

Phase Transition Enthalpy Measurements of Organic and Organometallic Compounds. Sublimation, Vaporization and Fusion Enthalpies From 1880 to 2010

William Acree, Jr.

Department of Chemistry, University of North Texas, Denton, Texas 76203

James S. Chickos^{a)}

Department of Chemistry and Biochemistry, University of Missouri—St. Louis, One University Boulevard, St. Louis, Missouri 63121

(Received 14 January 2010; accepted 15 January 2010; published online 4 October 2010)

A compendium of phase change enthalpies published within the period 1880–2010 is reported. Phase change enthalpies including fusion, vaporization, and sublimation are included for organic, organometallic, and a few inorganic compounds. This compendium is a combination of three previous series focusing on phase change enthalpies updated to 2009. Sufficient data are presently available for some compounds to permit thermodynamic cycles to be constructed, an important manner of evaluating the reliability of the measurements. Temperature adjustments of phase change enthalpies from the temperature of measurement to the standard reference temperature, $T=298.15$ K, are briefly discussed and a protocol for doing so is illustrated. © 2010 American Institute of Physics. [doi:10.1063/1.3309507]

Key words: Vaporization enthalpy; fusion enthalpy; sublimation enthalpy; compendium.

CONTENTS

1. Introduction.	1	7. Phase change enthalpies of C ₇ to C ₈ organic compounds.	135
2. Phase Change Enthalpies.	2	8. Phase change enthalpies of C ₉ to C ₁₀ organic compounds.	261
2.1. Estimation of heat capacities.	2	9. Phase change enthalpies of C ₁₁ to C ₁₄ organic compounds.	379
2.2. Vaporization enthalpies.	5	10. Phase change enthalpies of C ₁₅ to C ₂₀ organic compounds.	477
2.3. Sublimation enthalpies.	6	11. Phase change enthalpies of C ₂₁ to C ₁₉₂ organic compounds.	601
2.4. Fusion enthalpies.	6	12. Phase change enthalpies of organometallic and inorganic compounds.	689
2.5. Sample phase change enthalpy adjustment to $T=298.15$ K.	6		764
3. The Phase Change Enthalpy Compendium.	7		
4. References for Secs. 1–3.	8		
5. References.	872		

List of Tables

1. Group values for estimation of liquid [$\Gamma(l)$] and solid [$\Gamma(s)$] heat capacity at $T=298.15$ K (values in brackets are considered tentative values).	4
2. Some estimations of liquid and solid heat capacities ($\text{J mol}^{-1} \text{K}^{-1}$) at $T=298.15$ K.	5
3. Thermochemical cycles using Eq. (1).	7
4. Acronyms used in tables.	8
5. Phase change enthalpies of C ₁ to C ₄ organic compounds.	9
6. Phase change enthalpies of C ₅ to C ₆ organic compounds.	135

List of Figures

1. A hypothetical molecule illustrating the different carbon environments as defined in Table 1A.	2
2. A hypothetical molecule illustrating the different functional groups defined in Table 1B.	3

1. Introduction

Transition enthalpy measurements, that include measurements of sublimation, vaporization, and fusion of organic compounds, have been reported for well over 100 years. These properties find use in a number of disciplines that include chemical and environmental engineering, physics, and chemistry. The magnitude of these properties, when viewed

^{a)}Electronic mail: jsc@umsl.edu
© 2010 American Institute of Physics.

in perspective, provides valuable insight into the nature of both intra- and intermolecular interactions and thus are of interest to both experimentalists and theoreticians. While generally weaker than the chemical bond, the interactions responsible for these enthalpies are similar to those responsible for the self-assembly that occurs in everything from liquid crystals to biological systems.

The sheer number of measurements reported in the literature is testimony to the importance of these properties in providing an understanding of our physical world. One goal of this compendium is to bring together in one location measurements that often are scattered in the literature, sometimes in obscure places. Additionally, some of these measurements have significant errors associated with the property reported. These uncertainties are often not easily discernable, even to those familiar with the field. Despite significant improvement in instrumentation over the years, even recent measurements have errors significantly larger than reported. Provided all necessary transition enthalpies are available, an evaluation of the thermodynamic consistency of the values reported is possible. This compendium is an effort to compile these phase change enthalpies in a manner such that an assessment can be made as quantitatively as the data permits.

Equation (1) is the thermodynamic equality that relates phase change enthalpies, fusion, vaporization, and sublimation enthalpies but only if all enthalpies are referenced to a common temperature. Since experimental fusion enthalpies measured at ambient pressures are confined to the temperature T_{fus} , this equality is only applicable at the fusion temperature unless each enthalpy can be adjusted for temperature. Since $T=298.15$ K is the usual temperature of reference, adjustment of each of these enthalpies to $T=298.15$ K is preferable. A number of methods, some experimental and others computational, have been used to adjust both vaporization and sublimation enthalpies to $T=298.15$ K. A number of these have been summarized in previous compilations [1987CHI, 2003CHI/ACR2, 1999CHI/NIC]. The relationships outlined in this compendium for adjusting sublimation and vaporization enthalpies, those most familiar to the authors, have been in use for some time and have given acceptable results. Fusion enthalpies have been adjusted by combining the relationships used to adjust vaporization and sublimation enthalpies. The protocol is described below,

$$\Delta_{\text{sub}}H_m(T) = \Delta_{\text{vap}}H_m(T) + \Delta_{\text{fus}}H_m(T). \quad (1)$$

2. Phase Change Enthalpies

As noted above, phase change enthalpies are temperature dependent. Therefore, to adjust a phase change enthalpy for temperature requires the heat capacity of the two phases in question. For sublimation enthalpies, while the heat capacity of the solid, $C_{p,m}(s, 298.15 \text{ K})$, may be available, the heat capacity of the corresponding gas phase value, $C_{p,m}(g, 298.15 \text{ K})$, is generally not. Similarly for vaporization enthalpies, the experimental heat capacity of the liquid at T

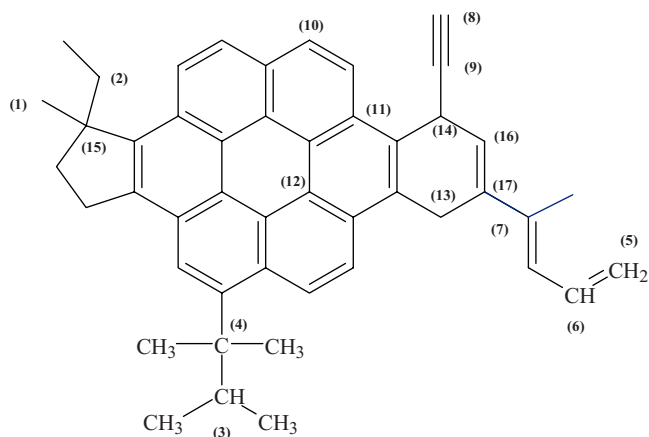


FIG. 1. (Color online) A hypothetical molecule illustrating the different carbon environments as defined in Table 1A.

$=298.15$ K, $C_{p,m}(l, 298.15 \text{ K})$, may be available, experimental data for the corresponding gas phase are usually lacking. For fusion enthalpies requiring both $C_{p,m}(s, 298.15 \text{ K})$ and $C_{p,m}(l, 298.15 \text{ K})$ for this adjustment, only one of these two properties is generally available at $T=298.15$ K. As a means of circumventing the lack of sufficient experimental data, empirical and theoretical relationships have been developed to adjust solid and liquid phase change enthalpies with temperature. Many of the phase change enthalpies reported in this compilation have been adjusted to $T=298.15$ K by the authors. The reader should consult the original literature to determine how this adjustment was made. In cases where the phase change enthalpy is reported only at the mean temperature of measurement, a few empirical relationships that can be used to adjust each respective phase change are discussed briefly below.

2.1. Estimation of heat capacities

A number of methods have been developed for the estimation of heat capacities [1990LYM/REE, 1908KOL/KUK, 1993CHI/HES]. The method employed in this article is a method developed by the authors and shown to give reasonably good temperature adjustments when used in combination with the equations to be described below [1998CHI, 1999SAB/XU, 2008ROU/TEM]. The method is based on group contributions. The group values (Γ) used in the estimation of $C_{p,m}(l, 298.15 \text{ K})$ and $C_{p,m}(s, 298.15 \text{ K})$ are provided in Table 1. For reference, Figs. 1 and 2 provide an example of a hypothetical molecule containing all of these groups. Each group, identified numerically in column 2 of Table 1, can be located in these figures. Some estimations using examples taken from the recent literature are given in Table 2. Note that some of the group values in Table 1 remain tentative and values for a few groups are not available for both condensed phases.

The calculations for *t*-butylbenzene using the group values of Table 1 are fairly straightforward and require no additional comments. Agreement with experiment for the solid is good. Agreement between estimated and experimental heat capaci-

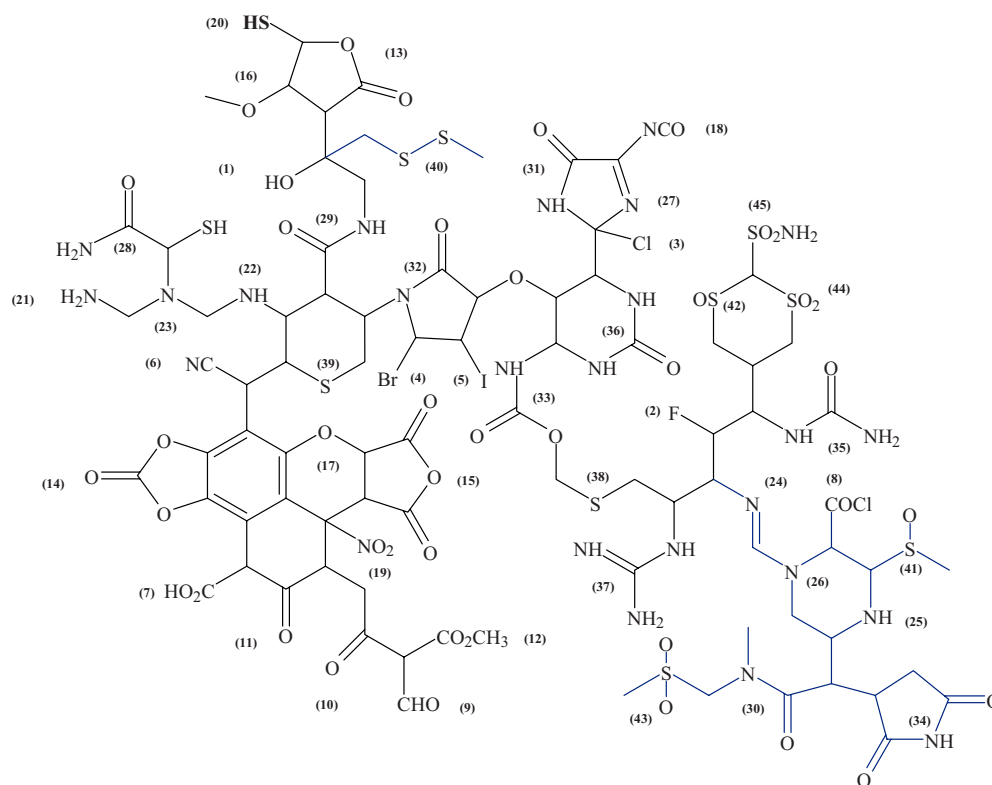


Fig. 2. (Color online) A hypothetical molecule illustrating the different functional groups defined in Table 1B.

ties for solid di-*tert*-butylbenzene also reported in the same article is not as good (estimated: 296.6; expt 347.8 J mol⁻¹ K⁻¹) [2009CHI/STE].

Estimations of the heat capacity of 5-aminouracil are not as obvious. This compound can be modeled as containing two cyclic secondary amides, a cyclic tertiary and cyclic quaternary sp² carbon atom and a primary sp³ nitrogen. Less intuitive models include modeling the ring as a cyclic urea and cyclic ketone or as a cyclic imide and cyclic secondary amine along with a cyclic tertiary and cyclic quaternary sp² carbon atom. All three of these estimations are illustrated in Table 2. All are in reasonable agreement with the experimental value [1907ZIE/SZT]. In this case, the less intuitive models give a slightly better agreement with experiment. Since group values for liquid cyclic ureas and cyclic imides are not available and cannot be used for the vaporization and fusion enthalpy temperature adjustments described below, the value for the more intuitive model should probably be used for fusion enthalpy adjustments. The heat capacity of 5-aminouracil can also be evaluated as its tautomer, 2,4-dihydroxy-5-aminopyrimidine. For comparative purposes, this estimation is also included in the table. In this case, the heat capacity of the solid is reproduced reasonably well but a larger discrepancy is observed between the predicted heat capacities of the liquid. In addition to 5-aminouracil, the heat capacities of a number of other solid uracils are reported in the same article [1907ZIE/SZT]: uracil (estimated: 124.6;

expt: 131.8); 6-aminouracil (estimated: 135; expt: 147.0), 6-amino-1-methyluracil (estimated: 177.9; expt: 166.2); 6-amino-1,3-dimethyluracil (estimated: 220.9; expt: 189 J mol⁻¹ K⁻¹). The estimated values were calculated using two cyclic amides to model the uracil ring. The values differ slightly from those reported by the authors of the article who used the same method but different groups to model the tertiary and quaternary sp² carbon groups [1907ZIE/SZT].

Calculations for 2,9-dimethyl-1,10-phenanthroline are fairly straightforward and in good agreement with the experimental heat capacity of the solid [2007BON/CAT]. The authors of this work also report the heat capacity of solid 1,10-phenanthroline as 115 J mol⁻¹ K⁻¹ at $T=298.15$ K. The estimated value for the solid at this temperature is 201.4 J mol⁻¹ K⁻¹ (calculations not shown). In this case the experimental value appears to be remarkably small when compared to the estimated value and to the value of naphthalene, 165 J mol⁻¹ K⁻¹ (300 K), used as a standard.

Other group additivity methods of varying complexity have been reported for the estimation of $C_{p,m}$ (1,298.15 K). References for these and an alternative method for calculating heat capacities can be found in the recent work of Kolska *et al.* [1908KOL/KUK].

TABLE 1. Group values for estimation of liquid [$\Gamma(l)$] and solid [$\Gamma(s)$] heat capacity at $T=298.15$ K (values in brackets are considered tentative values)

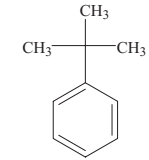
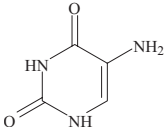
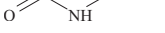

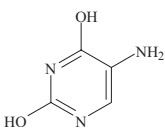
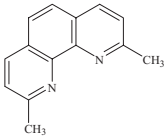
Group Atoms	$\Gamma(l)$ J mol ⁻¹ K ⁻¹	$\Gamma(s)$ J mol ⁻¹ K ⁻¹	
A. Hydrocarbon Groups			
primary sp ³ carbon	CH ₃ -	34.9	36.6
secondary sp ³ carbon	-CH ₂ -	31.9	26.9
tertiary sp ³ carbon	>CH-	22.4	9.0
quaternary sp ³ carbon	>C<	14.0	-5.0
secondary sp ² carbon	=CH ₂	25.8	[46.0]
tertiary sp ² carbon	=CH	27.8	21.4
quaternary sp ² carbon	=C<	21.7	[6.9]
tertiary sp carbon	≡CH	[34.3]	[37.1]
quaternary sp carbon	≡C-	28.9	[15.5]
tertiary aromatic sp ² carbon	=CH	21.8	17.5
quaternary aromatic sp ²	=C-	15.3	8.5
internal quaternary aromatic carbon ^a	=C-	16	[9.1]
cyclic secondary sp ³ carbon	-CH ₂ -	25.9	24.6
cyclic tertiary sp ³ carbon	>CH	20.6	11.7
cyclic quaternary sp ³ carbon	>C<	18	6.1
cyclic tertiary sp ²	=CH-	21.8	15.9
cyclic quaternary sp ²	=CR-	21.2	[4.7]
B. Functional Groups			
hydroxyl group (alcohols, phenols)	-OH	53.1	23.5
fluorine	-F	16.2	[24.8]
chlorine	-Cl	30.8	28.7
bromine	-Br	34.6	32.4
iodine	-I	39.1	[27.9]
nitrile	-C≡N	47.7	[42.3]
carboxylic acid	-(CO)OH	87.4	53.1
acid chloride	-(C=O)Cl	62.8	[60.2]
aldehyde	-(C=O)H	57.7	[84.5]
ketone	-(C=O)-	51.5	[28.0]
cyclic ketone	-(C=O)-	46.4	34.3
ester	-(C=O)O-	63.2	40.3
lactone	-(C=O)O-	[67.4]	[45.2]
cyclic carbonate	-O(C=O)O-	[92.0]	[68.2]
cyclic anhydride	-(C=O)O(C=O)-		[80.3]
ether	-O-	29.8	49.8
cyclic ether	-O-	24.6	9.7
isocyanate	O=C=N-	[58.2]	[52.7]
nitro group	-NO ₂	[58.6]	56.1
thiol	-SH	49.0	[51.9]
primary sp ³ nitrogen	-NH ₂	59.4	21.6
secondary sp ³ nitrogen	-NH-	[51.0]	[-0.3]
tertiary sp ³ nitrogen	-N<	22.0	[31.5]
tertiary sp ² nitrogen	=N-	[44.4]	10.7
cyclic secondary sp ³ nitrogen	-NH-	46.0	[23.9]
cyclic tertiary sp ³ nitrogen	-N<	[28.6]	1.2
cyclic tertiary sp ² nitrogen	=N-	20.7	13.9
primary amide	-(C=O)NH ₂	[41.0]	[54.4]
secondary amide	-(C=O)NH-	79.9	44.4
tertiary amide	-(C=O)N<	[82.4]	
cyclic secondary amide	-(C=O)NH-	[92.0]	46.4
cyclic tertiary amide	-(C=O)N-	[170]	[52.7]
Carbamate	-NH(C=O)O-		[76.1]
cyclic imide	-(C=O)NH(C=O)-		[74.1]
monsubstituted urea	-NH(C=O)NH ₂		[82.8]
cyclic urea	-NH(C=O)NH-		[63.6]
monsubstituted guanidine	-NH(C=NH)NH ₂		[59.4]

TABLE 1. Group values for estimation of liquid [$\Gamma(l)$] and solid [$\Gamma(s)$] heat capacity at $T=298.15$ K (values in brackets are considered tentative values)—Continued

	Group Atoms	$\Gamma(l)$ J mol ⁻¹ K ⁻¹	$\Gamma(s)$ J mol ⁻¹ K ⁻¹
sulfide	-S-	40.3	[116]
cyclic sulfide	-S-	33.8	18.2
disulfide	-S-S-	74.5	[41.0]
sulfoxide	-(S→O)-	[83.7]	[47.7]
cyclic sulfoxide	-(S→O)-		[31.0]
sulfone	-(S→O) ₂ -		[52.4]
cyclic sulfone	-(S→O) ₂ -		[38.7]
sulfonamide	-(S→O ₂)NH ₂		[104]
quaternary silicon	>Si<	30.9	32.4
tertiary aluminum	>Al-	[46.9]	
quaternary tin	>Sn<	58.6	[77.2]
quaternary germanium	>Ge<	[48.1]	[18.9]
phosphine oxide	-(P→O)<		[28.5]

^aAn internal quaternary carbon refers to internal carbon atoms as found in coronene or graphite.

TABLE 2. Some estimations of liquid and solid heat capacities (J mol⁻¹ K⁻¹) at $T=298.15$ K

	$3 \text{ CH}_3\text{-} + > \text{C} < + 5(\text{=CH})_{\text{arom}} + (\text{=C-})_{\text{arom}}$ $C_{p,m}(l, 298.15 \text{ K}) = 3(34.9) + 14.0 + 5(21.8) + 15.3 = 243$ (lit 241) [2009CHI/STE] $C_{p,m}(s, 298.15 \text{ K}) = 3(36.6) - 4.98 + 5(17.5) + 8.5 = 200.8$
	$2(-(\text{C=O})\text{NH-})_{\text{cyc}} + (\text{=CH-})_{\text{cyc}} + (\text{=CR-})_{\text{cyc}} + (-\text{NH}_2)$ $C_{p,m}(l, 298.15 \text{ K}) = 2(92) + 21.8 + 21.2 + 59.4 = 286.4$ $C_{p,m}(s, 298.15 \text{ K}) = 2(46.4) + 15.9 + 4.73 + 21.6 = 135.0$ (lit 145) [2007ZIE/SZT]
	$(-\text{NH}(\text{C=O})\text{NH-})_{\text{cyc}} + (-\text{C=O-})_{\text{cyc}} + (\text{=CH-})_{\text{cyc}} + (\text{=C-})_{\text{cyc}} + -\text{NH}_2$ $C_{p,m}(s, 298.15 \text{ K}) = 63.6 + 34.3 + 15.9 + 4.73 + 21.6 = 140.1$
	$(-\text{C=O})\text{NH}(\text{C=O-})_{\text{cyc}} + (-\text{NH-})_{\text{cyc}} + (\text{=CH-})_{\text{cyc}} + (\text{=C-})_{\text{cyc}} + -\text{NH}_2$ $C_{p,m}(s, 298.15 \text{ K}) = 74.1 + 23.9 + 15.9 + 4.73 + 21.6 = 140.2$
	$3(\text{=C-})_{\text{arom}} + (\text{=CH})_{\text{arom}} + 2(\text{=N-})_{\text{cyc}} + 2(-\text{OH}) + (-\text{NH}_2)$ $C_{p,m}(l, 298.15 \text{ K}) = 3(15.3) + (21.8) + 2(20.7) + 2(53.1) + 54.9 = 270.2$ $C_{p,m}(s, 298.15 \text{ K}) = 3(8.5) + 3(17.5) + 2(13.9) + 2(23.5) + 21.6 = 139.4$
	$6(\text{=C-})_{\text{arom}} + (\text{=CH})_{\text{arom}} + 2(\text{=N-})_{\text{cyc}} + 2(-\text{OH}) + (-\text{NH}_2)$ $C_{p,m}(l, 298.15 \text{ K}) = 6(21.8) + 6(15.3) + 2(20.7) + 2(34.9) = 333.8$ $C_{p,m}(s, 298.15 \text{ K}) = 6(17.5) + 6(8.5) + 2(13.9) + 2(36.6) = 248$ (lit 253) [2007BON/CAT]

2.2. Vaporization enthalpies

Equation (2) is an equation derived to model the differences in heat capacity between the liquid and gas phases [1993CHI/HOS]. It has been derived by correlating vaporization enthalpy differences measured at temperature T and generally at $T=298.15$ K with the heat capacity of the corresponding liquid at $T=298.15$ K for which reliable vaporization data are available. The vaporization enthalpy data used were obtained from the critical review and data compilation of Majer and Svoboda [1993CHI/HOS]. It has been found to provide satisfactory results for adjustments ranging

from approximately $T=500$ to 250 K [1998CHI, 1999SAB/XU, 2008ROU/TEM]. The adjustments have been generally been applied from the mean temperature of measurement, \bar{T} , to 298.15 K. Adjustments for temperatures above $T=500$ K should be viewed with caution,

$$\Delta_{\text{vap}} H_{\text{m}}^{\circ}(298.15 \text{ K}) / \text{kJ mol}^{-1} = \Delta_{\text{vap}} H_{\text{m}}^{\circ}(\bar{T}/\text{K}) + \{10.58 + 0.26[C_{p,m}(l, 298.15 \text{ K})] / \text{J mol}^{-1} \text{ K}^{-1}\} \{\bar{T}/\text{K} - 298.15\} / 1000. \quad (2)$$

The term $[C_{p,m}(l)](298.15 \text{ K}) / \text{J mol}^{-1} \text{ K}^{-1}$ refers to the mo-

lar heat capacity of the liquid phase at $T=298.15$ K; estimated heat capacities were used in deriving this relationship [1993CHI/HES]. Experimental values can also be used. An uncertainty of 16 J mol^{-1} has been associated with the bracketed term in Eq. (2). Some examples using Eq. (2) are given in Table 3.

2.3. Sublimation enthalpies

Sublimation enthalpies can also be adjusted to $T=298.15$ K from the mean temperature of measurement using a similar relationship, Eq. (3). This equation has also been derived by correlating sublimation enthalpy differences measured at temperature \bar{T} and generally at $T=298.15$ K with the estimated heat capacity of each corresponding crystalline compound at $T=298.15$ K [1993CHI/HOS]. Unlike vaporization enthalpy data, much less critically evaluated sublimation enthalpy data are available. As a consequence, a larger uncertainty is associated with this adjustment,

$$\begin{aligned} \Delta_{\text{sub}}H_{\text{m}}^{\circ}(298.15 \text{ K})/\text{kJ mol}^{-1} \\ = \Delta_{\text{sub}}H_{\text{m}}^{\circ}(\bar{T}/\text{K}) + \{0.75 \\ + 0.15[C_{p,m}(s,298.15 \text{ K})]/\text{J mol}^{-1} \text{ K}^{-1}\} \{\bar{T}/\text{K} \\ - 298.15\}/1000. \end{aligned} \quad (3)$$

The term $[C_{p,m}(s,298.15 \text{ K})]$ refers to the molar heat capacity of the solid phase at $T=298.15$ K. The relationship was derived using solid heat capacities estimated by the group additivity method described above [1993CHI/HES]. This equation has also been found to give satisfactory results for temperatures up to approximately $T=500$ K. Group values used in these estimations are also summarized in Table 1. As with heat capacities of the liquid state, experimental heat capacity values can be substituted for $C_{p,m}(s,298.15 \text{ K})$ if available. When using Eq. (3), an uncertainty equal to one-third the magnitude of the total temperature adjustment should be assumed. While this uncertainty, arbitrarily chosen, is significant, some compensation is afforded by the fact that temperature adjustments of sublimation enthalpies are generally much smaller than the corresponding adjustments for vaporization enthalpies. In cases where the experimental sublimation enthalpy is reported by the author at $T=298.15$ K, the reader should consult the original literature to determine how the temperature adjustment was achieved. A number of different methods have been used in the literature for this adjustment and it has been found that some methods provide more thermodynamically consistent results than others when using Eq. (1) [1998CHI].

2.4. Fusion enthalpies

Temperature adjustments for fusion enthalpies from T_{fus} to $T=298.15$ K can be achieved by noting that if Eq. (2) is subtracted from Eq. (3), the heat capacity of the gas phase common to both cancels and the remainder results in the difference in molar heat capacity between the liquid and solid phases, $\Delta C_p(\text{cr},1)$, Eq. (4). Equation (4) has been suc-

cessfully used in combination with Eqs. (1) and (2) to predict sublimation enthalpies at $T=298.15$ K [2008ROU/TEM, 2004BAS/CHI]. An uncertainty equal to one-third the magnitude of the total temperature adjustment has generally been assigned to this temperature adjustment,

$$\begin{aligned} \Delta_{\text{fus}}H_{\text{m}}^{\circ}(298.15 \text{ K})/\text{kJ mol}^{-1} \\ = \Delta_{\text{fus}}H_{\text{m}}(T_{\text{fus}}) + \Delta_{\text{trns}}H_{\text{m}}(T_{\text{trns}}) \\ + [0.15C_{p,m}(s,298.15 \text{ K}) - 0.26C_{p,m}(l,298.15 \text{ K}) \\ - 9.83][(T_{\text{fus}}/\text{K} - 298.15)/1000]. \end{aligned} \quad (4)$$

Many compounds do not exhibit solid-solid phase transitions at temperatures below fusion. For those compounds that do, the enthalpy of the transition also needs to be added to the fusion enthalpy when using Eq. (1) if the sublimation enthalpy was measured at temperatures below the transition temperature T_{trns} . If the sublimation enthalpy was measured above $T=T_{\text{trns}}$, the fusion enthalpy can be used directly in Eq. (1) to reproduce the sublimation enthalpy. If T_{trns} exceeds $T=298.15$ K, then the transition enthalpy will need to be added to the sublimation enthalpy to obtain $\Delta_{\text{sub}}H_{\text{m}}^{\circ}(298.15 \text{ K})$.

2.5. Sample phase change enthalpy adjustment to $T=298.15$ K

A few examples serve to illustrate the usefulness of Eqs. (1)–(4) when all three phase change enthalpies data for a particular substance are available. These are shown in Table 3. The first example, cyclohexanone, illustrates the use of these equations for temperature adjustments below ambient temperature. Cyclohexanone exhibits a solid-solid phase transition at a temperature below the temperature range used in the sublimation enthalpy measurement and hence is not included in Eq. (1). Since the heat capacity of the liquid generally exceeds that of the solid phase, adjustment to $T=298.15$ K in this case actually increases the fusion enthalpy from 1.3 to 3.0 kJ mol^{-1} . Three vaporization enthalpies have been reported, in good agreement with each other when adjusted to the reference temperature. The sublimation enthalpy, when adjusted to the reference temperature, is attenuated slightly, since the heat capacity of the solid generally exceeds that of the gas phase. The sublimation enthalpy, $48.2 \pm 0.3 \text{ kJ mol}^{-1}$, is within experimental error of the mean value calculated using Eq. (1), $49.4 \pm 0.8 \text{ kJ mol}^{-1}$. The uncertainty in the latter value is the mean uncertainty associated with each entry, and for one of the entries, the uncertainty associated with only the temperature adjustment.

Phenacetin illustrates a situation not uncommon with pharmaceuticals, the possible existence of polymorphism. The fusion enthalpy of phenacetin has been reported a number of times. Most reports are in reasonable agreement with each other except for one. In this case, the value is probably in error since the first and fourth fusion enthalpy entries, which are in disagreement, have been reported by the same research group. Ignoring the fourth value, an average value of $25.2 \pm 1.8 \text{ kJ mol}^{-1}$ is calculated, which when added to the

vaporization enthalpies results in two values, one of which appears to be in much better agreement with a single determination of the sublimation enthalpy. The uncertainty in $\Delta_{\text{fus}}H_m^\circ(298\text{ K})$ only reflects the uncertainty in the temperature adjustments.

Carbazole is another example of a substance that exhibits a solid-solid phase transition. In this case the fusion enthalpy at $T=298.15\text{ K}$ should include this transition since the subli-

mation enthalpy was measured at a temperature below the transition. A solid to liquid total phase change enthalpy of $15.6 \pm 3.5\text{ kJ mol}^{-1}$ is obtained which when added to the vaporization enthalpy results in sublimation enthalpy values of 97.9 ± 5.1 and $91.8 \pm 3.5\text{ kJ mol}^{-1}$. In this instance, only one of the sublimation enthalpy values calculated in this manner is consistent with the two sublimation enthalpies measured directly.

TABLE 3. Thermochemical cycles using Eq. (1)

Enthalpy	$\Delta H_m(T)$ kJ mol ⁻¹	\bar{T}/K	$C_{p,m}(l, 298\text{ K})$ J mol ⁻¹ K ⁻¹	$C_{p,m}(s, 298\text{ K})$ J mol ⁻¹ K ⁻¹	$\Delta C_{p,m}\Delta T$ kJ mol ⁻¹	$\Delta H_m(298\text{ K})$ kJ mol ⁻¹	$\Delta_{\text{fus}}H_m^\circ(298\text{ K})$ +		Reference ^a
							$\Delta_{\text{vap}}H_m^\circ(298\text{ K})$ kJ mol ⁻¹	$\Delta_{\text{sub}}H_m^\circ(298\text{ K})$ kJ mol ⁻¹	
C₆H₁₀O cyclohexanone									
$\Delta_{\text{trns}}H_m(T_{\text{trns}})$	8.66	220.8							[1980NAK/SUG]
$\Delta_{\text{fus}}H_m(T_{\text{fus}})$	1.33	245.2	175.9	157.3	1.7	3.0 ± 0.5			[1980NAK/SUG]
$\Delta_{\text{vap}}H_m^\circ(\bar{T}/\text{K})$	43.1	358	175.9	157.3	3.4 ± 1.0	46.5	49.5 ± 1.1		[2006TEO/BAR]
	46.6 ± 0.4	298	175.9	157.3	0	46.6 ± 0.4	49.6 ± 0.6		[1995CHI/HOS]
	44.0 ± 0.1	333	175.9	157.3	2.0 ± 0.6	46.0 ± 0.6	49.0 ± 0.8		[1993AUC/MON]
$\Delta_{\text{sub}}H_m^\circ(\bar{T}/\text{K})$	49.3	254	175.9	157.3	-1.07 ± 0.3	48.2 ± 0.3			[1948NIT/SEK2]
C₁₀H₁₃NO₂ 4'-ethoxyacetanilide (phenacetin)									
$\Delta_{\text{fus}}H_m(T_{\text{fus}})$	30 ± 1.0	409.6	329.2	281.3	-5.9 ± 1.8	24.1 ± 1.8			[2009VEC/TOM]
	28.8	408.3	329.2	281.3	-5.9 ± 1.8	22.9 ± 1.8			[2009PEN/ESC]
	34.1	407.4	329.2	281.3	-5.8 ± 1.7	28.3 ± 1.7			[2006WAS/HOL]
	21.4 ± 0.9	410.2	329.2	281.3	-6.0 ± 1.8	15.4 ± 2.0			[2004VEC/CAT]
	31.3	407.2	329.2	281.3	-5.8 ± 1.7	25.4 ± 1.7			[1990MAN/AHU]
					average	25.2 ± 1.8			
$\Delta_{\text{vap}}H_m^\circ(\bar{T}/\text{K})$	79.0 ± 1.0	459	329.2	281.3	15.5 ± 2.6	94.5 ± 2.6	119.6 ± 3.2		[2009VEC/TOM]
	82.6	478	329.2	281.3	17.3 ± 2.9	99.9 ± 2.9	125.0 ± 3.4		[1987STE/MAL]
$\Delta_{\text{sub}}H_m^\circ(\bar{T}/\text{K})$	115.5	349.5	329.2	281.3	2.2 ± 0.7	117.7 ± 0.7			[1972WIE]
C₁₂H₉N carbazole									
$\Delta_{\text{trns}}H_m(T_{\text{trns}})$	0.27	420							[1969ROB/SCO]
$\Delta_{\text{fus}}H_m(T_{\text{fus}})$	26.9	518.7	281.6	197.9	-11.8 ± 3.5	15.4 ± 3.5			[2000LIS/JAM]
	27.2	516	281.6	197.9	-11.6 ± 3.5	15.9 ± 3.5			[1996BUR/KOL]
					average	15.6 ± 3.5			
$\Delta_{\text{vap}}H_m^\circ(\bar{T}/\text{K})$	76.2	298	281.6	197.9	0	76.2	91.8 ± 3.5		[1996GOV/RUT]
	63.3	525	281.6	197.9	19.0 ± 3.6	82.3 ± 3.6	97.9 ± 5.1		[1983SIV/MAR]
$\Delta_{\text{sub}}H_m^\circ(\bar{T}/\text{K})$	101.2 ± 1.1	355	281.6	197.9	1.7 ± 0.5	102.9 ± 1.2			[1990JIM/ROU]
	97.7 ± 0.3	298	281.6	197.9	0	97.7 ± 0.3			[1987SAB/ANT]

^aReferences for fusion vaporization and sublimation can be found in Sec. 5.

3. The Phase Change Enthalpy Compendium

The phase change enthalpy data reported in this compendium have been reported over the time period of 1880–2010. The data are combination of three compendia published in recent years and updated to the present [1902CHI/ACR, 2003CHI/ACR, 2003CHI/ACR2, 1999CHI/ACR, 1909CHI/ACR]. Vaporization enthalpies and sublimation enthalpies have been measured for many years and numerous tech-

niques have been developed to do so. Generally, these techniques can be categorized into two groups, calorimetric techniques, in which the phase change enthalpy is measured directly, and techniques in which vapor pressure is measured directly or indirectly as a function of temperature. Providing on the experimental setup, calorimetric methods can also provide vapor pressures if an effusion cell is used. More recently, gas chromatographic techniques have also been developed to measure vaporization enthalpies. These techniques can provide both vaporization enthalpies and liquid

vapor pressures but since standards must be used, some care must be used in assessing their reliability. One technique, referred to as correlation-gas chromatography, has been shown to be successful provided standards are chosen with reliable values and appropriately related functional groups [2009LIP/HAN]. Fusion enthalpies have generally been measured by some form of calorimetry. These techniques are described by a number of acronyms used throughout the tables. A summary of these acronyms and their meanings can be found in Table 4.

In some cases, the vapor pressure–temperature data reported in the literature authors were analyzed by using the Clausius Clapeyron relationship, Eq. (5), where $C=0$, by the authors. The “Handbook of the Thermodynamics of Organic Compounds,” by Stephenson and Malanowski [1987STE/MAL], was a useful source of vaporization enthalpy information provided in the form of Antoine Constants. Unfortunately, references to the original literature data are not provided by these authors. Vaporization enthalpies and some sublimation enthalpies were calculated from the A, B, and C constants reported in this compendium using Eq. (6),

$$\log_{10} p \text{ (kPa)} = A - B/(C + T), \quad (5)$$

$$\Delta_{\text{vap}}H_m^{\circ}(T_m) = 2.303RB[\bar{T}/(\bar{T} + C)]^2. \quad (6)$$

Phase change enthalpies for organic compounds are reported in Tables 5–11. Organometallic compounds are reported in Table 12. Organometallic compounds are arranged alphabetically according to the metal.

In Tables 5–12, some enthalpy values are cited without a reference on the same line. The appropriate reference can be found with the next value with a reference (multiple enthalpy values were taken from the same source). In addition, in Tables 5–12, where a value is prefixed with a “U” (e.g., “U 66.0 ± 21.2”), this indicates unreliable data.

TABLE 4. Acronyms used in tables

A	calculated from the vapor pressure data reported by the method of least squares
AC	adiabatic calorimeter
B	calculated from the difference of the enthalpies of sublimation at temperature T and fusion at the melting point.
BG	Bourdon gauge
C	calorimetric determination
CATH	cathetometer
CGC	correlation-gas chromatography
CGC-DSC	combined correlation gas chromatography-differential scanning calorimetry
CR	Cryoscopy
DBM	dibutyl pthalate manometer
DM	diaphragm manometer
DSC	differential scanning calorimeter
DTA	differential thermal analysis
E	estimated value
EB	ebullimeter
EM	electronic manometer
EV	evaporation
F	fluorescence
GC	gas chromatography
GCC	gas chromatography-calorimetry
GS	gas saturation, transpiration
GSM	glass spring manometer
HG	Heise gauge
HSA	head space analysis
I	isoteniscope
IP	inclined piston manometry
KG	Knudsen gauge
LE	Langmuir evaporation
MDSC	Modulated differential scanning calorimetry
ME	mass effusion-Knudsen effusion
MEM	modified entrainment method
MG	McLeod Gauge
MM	mercury manometer
MS	mass spectrometry
OM	oil manometer
PG	pressure gauge
QCM	quartz crystal microbalance
QF	quartz fiber
QR	quartz resonator
RG	Rodebush gauge
S-F	sublimation-fusion
SG	spoon gauge
SRFG	spinning rotor friction gauge
STG	strain gauge
T	tensiometer
TCM	thermal conductivity manometer
TE	torsion effusion
TGA	thermal gravimetric analysis
TSGC	temperature scanning gas chromatography
U	unreliable
UV	ultraviolet spectroscopy
V	viscosity gauge
VG	viscosity gauge

4. References for Secs. 1–3

- 1972WIE H. G. Wiedemann, *Thermochim. Acta.* **3**, 355 (1972).
- 1985MAJ/SVO V. Majer and V. Svoboda, *Enthalpies of Vaporization of Organic Compounds: ACritical Review and Data Compilation*, IU-

	PAC Chemical Data Series No. 32 (Blackwell, Oxford, 1985).	1999SAB/XU	R. Sabbah, A. Xu-wu, J. S. Chickos, M. L. Plana Leitao, M. V. Roux, and L. A. Torres, <i>Thermochim. Acta</i> 331 , 93 (1999).
1987CHI	J. S. Chickos, in <i>Molecular Structure and Energetics</i> , edited by J. F. Liebman and A. Greenberg (VCH, New York, 1987), Vol. 2, Chap. 3.	2002CHI/ACR	J. S. Chickos and W. E. Acree, Jr., <i>J. Phys. Chem. Ref. Data</i> 31 , 537 (2002).
1987STE/MAL	R. M. Stephenson and S. Malanowski, <i>Handbook of the Thermodynamics of Organic Compounds</i> (Elsevier, New York, 1987).	2003CHI/ACR	J. S. Chickos and W. E. Acree, Jr., <i>Thermochim. Acta</i> 395 , 59 (2003).
1976WIE	H. G. Wiedemann, <i>Thermochim. Acta</i> 3 , 355 (1972).	2003CHI/ACR2	J. S. Chickos and W. E. Acree, Jr., <i>J. Phys. Chem. Ref. Data</i> 32 , 519 (2003).
1990LYM/REE	<i>Handbook of Chemical Property Estimation Methods</i> , edited by W. J. Lyman, W. F. Reehl, and D. H. Rosenblatt (American Chemical Society, Washington, D. C., 1990).	2004BAS/CHI	A. Bashir-Hashemi, J. S. Chickos, W. Hanshaw, H. Zhao, B. S. Farivar, and J. F. Liebman, <i>Thermochim. Acta</i> 424 , 91 (2004).
1993CHI/HES	J. S. Chickos, D. G. Hesse, and J. F. Liebman, <i>Struct. Chem.</i> 4 , 261 (1993).	2007ZIE/SZT	W. Zielenkiewicz and P. Szturner, <i>J. Chem. Eng. Data</i> 52 , 624 (2007).
1993CHI/HOS2	J. S. Chickos, S. Hosseini, D. G. Hesse, and J. F. Liebman, <i>Struct. Chem.</i> 4 , 271 (1993).	2007BON/CAT	M. G. Bonicelli, A. Catalani, G. Mariano, and S. Vecchio, <i>Thermochim. Acta</i> 466 , 69 (2007).
1998CHI	J. S. Chickos, <i>Thermochim. Acta</i> 313 , 19 (1998).	2008KOL/KUK	Z. Kolsa, J. Kukal, M. Zabransky, and V. Ruzicka, <i>Ind. Eng. Chem. Res.</i> 47 , 2075 (2008).
1999CHI/ACR	J. S. Chickos, W. E. Acree, Jr., and J. F. Liebman, <i>J. Phys. Chem. Ref. Data</i> 28 , 1535 (1999).	2008ROU/TEM	M. V. Roux, M. Temprado, J. S. Chickos, and Y. Nagano, <i>J. Phys. Chem. Ref. Data</i> 37 , 1855 (2008).
1999CHI/NIC	J. Chickos, G. Nichols, J. Wilson, M. Orf, P. Webb, and J. Wang, in <i>Energetics of Stable Molecules and Reactive Intermediates</i> , NATO Science Series C, Vol. 535, edited by M. da Piedade (Kluwer Academic, Boston: MA, 1999), pp 177–202	2009CHI/ACR	J. S. Chickos and W. E. Acree, Jr., <i>Thermochim. Acta</i> 495 , 5 (2009) and supplemental information.
		2009CHI/STE	R. D. Chirico and W. V. Steele, <i>J. Chem. Thermodyn.</i> 41 , 392 (2009).
		2009LIP/HAN	D. Lipkind, W. Hanshaw, and J. S. Chickos, <i>J. Chem. Eng. Data</i> 54 , 2930 (2009) and references cited.

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds

Molecular Formula	CAS Reg No	Compound		Method	Reference	
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)			T_m (K)
CBrClF ₂	[353-59-3]	bromochlorodifluoromethane				
	$\Delta_v H$	(268–324)	23.0	283	A	[1987STE/MAL]
	$\Delta_v H$	(194–287)	23.1	272	A	[1987STE/MAL]
	$\Delta_v H$	(321–403)	22.4	336	A	[1987STE/MAL]
	$\Delta_v H$	(403–427)	23.1	415	A	[1987STE/MAL]
	$\Delta_v H$	(178–283)	26.0	193		[1979KUD/KUD]
		(178–283)	18.7	268		[1960GLE, 1984BOU/FRI]
CBrCl ₃	[75-62-7]	bromotrichloromethane				
	$\Delta_{\text{trs}} H$		4.62	238.2		
	$\Delta_{\text{trs}} H$		0.53	259.4		
	$\Delta_{\text{fus}} H$		2.54	267.5	AC	[1991ACR, 1995OHT/YAM]
	$\Delta_v H$	(273–387)	35.0	288		[1979KUD/KUD]
		(294–443)	36.1	309	A	[1970DYK, 1987STE/MAL]
CBrF ₃	[75-63-8]	bromotrifluoromethane				
	$\Delta_v H$	(276–340)	17.8	291	A	[1987STE/MAL]
	$\Delta_v H$	(160–267)	17.7	252	A	[1987STE/MAL]
		(165–216)	19.1	180		[1979KUD/KUD]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
CBrFO	[753-56-0]	carbonic bromide fluoride				
	$\Delta_v H$	(197–256)	22.9	241	A	[1987STE/MAL]
CBrN	[506-68-3]	cyanogen bromide				
	$\Delta_{\text{sub}} H$	(273–308)	45.2 ± 4.2		MM	[1954LOR/WOO, 1970COX/PIL]
	$\Delta_{\text{sub}} H$	(256–308)	47.0		GS	[20BAX/BEZ]
	$\Delta_{\text{sub}} H$	(273–313)	45.9	288		[1954LOR/WOO, 1984BOU/FRI]
CBrN₃O₆	[560-95-2]	bromotrinitromethane				
	$\Delta_v H$	(318–335)	47.8	326	A	[1987STE/MAL, 1970CAR/ZIM]
CBr₂Cl₂	[594-18-3]	dibromodichloromethane				
	$\Delta_{\text{trs}} H$		5.43	258.8		
	$\Delta_{\text{fus}} H$		2.31	294.4		[1995OHT/YAM]
CBr₂F₂	[75-61-6]	dibromodifluoromethane				
	$\Delta_v H$	(247–297)	26.1	282	A	[1987STE/MAL, 1959MCD/SHR, 1979KUD/KUD, 1970DYK]
	$\Delta_v H$	(156–218)	18.6	203		[1948BAN/EME]
CBr₃F	[353-54-8]	tribromofluoromethane				
	$\Delta_v H$	(315–380)	34.4	330	A	[1987STE/MAL, 1948BAN/EME]
CBr₄	[558-13-4]	carbon tetrabromide				
	$\Delta_{\text{trs}} H$		5.94	320		
	$\Delta_{\text{fus}} H$		3.95	363.2		[1996DOM/HEA]
	$\Delta_{\text{sub}} H$ (<i>mono</i>)		54.5 ± 0.7	298	C	[1984BIC/MIN]
	$\Delta_{\text{sub}} H$ (<i>mono</i>)	(295–319)	54.4 ± 1.3	307	BG	[1959BRA/DRU]
	$\Delta_{\text{sub}} H$ (<i>cubic</i>)	(321–329)	49.4 ± 1.3	325	BG	[1959BRA/DRU]
	$\Delta_{\text{sub}} H$ (<i>cubic</i>)		48.3	320		[1955HAR/SWI]
	$\Delta_v H$	(375–463)	48.3	390		[1979KUD/KUD]
	$\Delta_v H$	(369–463)	48.2	384	A	[1987STE/MAL, 1947STU]
CClFO	[353-49-1]	carbonic chloride fluoride				
	$\Delta_v H$	(165–211)	22.7	196	A	[1987STE/MAL, 1964FIS/BUC]
	$\Delta_v H$	(157–227)	22.0	192		[1948EME/WOO]
CClF₂NO	[16847-30-6]	difluorocarbamoyl chloride				
	$\Delta_v H$	(189–234)	25.8	219	A	[1987STE/MAL]
CClF₃	[75-72-9]	chlorotrifluoromethane				
	$\Delta_v H$	(268–302)	16.0	283	A	[1987STE/MAL]
	$\Delta_v H$	(133–185)	17.0	170	A	[1987STE/MAL]
	$\Delta_v H$	(184–246)	15.7	231	A	[1987STE/MAL]
	$\Delta_v H$	(243–271)	15.7	257	A	[1987STE/MAL]
	$\Delta_v H$	(145–192)	16.8	177	A	[1987STE/MAL, 1979KUD/KUD]
	$\Delta_v H$	(124–191)	17.1	177	A	[1947STU]
$\Delta_v H$	(134–298)	NA			[1941RIE]	
CClF₃O	[22082-78-6]	trifluoromethyl hypochlorite				
	$\Delta_v H$	(160–226)	21.2	211	A	[1987STE/MAL]
	$\Delta_v H$	(142–219)	19.6	204	A	[1987STE/MAL]
CClF₃O₂	[32755-26-3]	peroxyhypochlorous acid, trifluoromethyl ester				
	$\Delta_v H$	(163–296)	23.4	281	A	[1987STE/MAL]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
CClF ₃ O ₃ S	[6069-31-4]	fluorosulfuric acid, chlorodifluoromethyl ester				
	$\Delta_{\text{v}}H$	(227–309)	32.1	243		[1999DYK/SVO]
	$\Delta_{\text{v}}H$	(228–310)	34.6	243	A	[1987STE/MAL, 1966DES/CAD]
CClF ₃ S	[421-17-0]	trifluoromethanesulfonyl chloride				
	$\Delta_{\text{v}}H$	(247–272)	24.5	260	A	[1987STE/MAL, 1999DYK/SVO]
CClF ₄ N	[13880-71-2]	difluoro(difluorochloromethyl)amine				
	$\Delta_{\text{v}}H$	(209–277)	26.6	262	A	[1987STE/MAL]
CClF ₄ NO ₂ S	[19419-95-5]	chloro(trifluoromethyl) sulfamoyl fluoride				
	$\Delta_{\text{v}}H$	(253–288)	28.8	273	A	[1987STE/MAL, 1999DYK/SVO]
CClF ₄ NO ₁₂ S ₄	[53684-03-0]	fluorosulfuric acid, bis[[fluorosulfonyl]oxy]amino]chloromethylene ester				
	$\Delta_{\text{v}}H$		42.6	424		[1975KIR/LAS]
CClF ₇ S	[42179-04-4]	chlorotetrafluoro (trifluoromethyl) sulfur				
	$\Delta_{\text{v}}H$	(293–353)	25.9	323		[1999DYK/SVO]
CCIN	[506-77-4]	cyanogen chloride				
	$\Delta_{\text{sub}}H$	(196–259)	35.7	228	A	[1947STU]
	$\Delta_{\text{v}}H$	(196–286)	32.2	271		[1947STU]
CCl ₂ FNO	[32751-02-3]	dichlorocarbamic fluoride				
	$\Delta_{\text{v}}H$		40.7			[1972DEM/SHR]
CCl ₂ F ₂	[75-71-8]	dichlorodifluoromethane				
	$\Delta_{\text{v}}H$	(282–345)	20.0	297	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$	(173–244)	21.4	229	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$	(173–240)	21.6	225	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$	(236–285)	20.4	270	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$	(341–385)	20.5	356	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$	(172–279)	22.9	187		[1979KUD/KUD]
	$\Delta_{\text{v}}H$	(154–243)	21.5	228		[1947STU]
	$\Delta_{\text{v}}H$		20.4	243		[1931BUF/FLE]
CCl ₂ F ₃ N	[24618-60-8]	N,N-difluoro-1,1-dichloro-1-fluoromethylamine				
	$\Delta_{\text{v}}H$	(209–277)	27.0	262	I	[1970ZAB/SHR]
CCl ₂ F ₃ N	[13880-73-4]	N,N-dichloro-1,1,1-trifluoromethylamine				
	$\Delta_{\text{v}}H$	(226–291)	25.8	276	A	[1987STE/MAL]
CCl ₂ F ₃ N	[33757-11-8]	N,1-dichloro-N,1,1-trifluoromethylamine				
	$\Delta_{\text{v}}H$	(226–291)	26.4	258		[1971SWI/ZAB]
CCl ₂ F ₃ NS	[10564-47-3]	(trifluoromethyl)imidodisulfurous dichloride				
	$\Delta_{\text{v}}H$	(284–344)	35.4	298		[1999DYK/SVO]
	$\Delta_{\text{v}}H$	(283–362)	33.7	298	A	[1987STE/MAL]
CCl ₂ F ₃ P	[421-58-9]	(trifluoromethyl)dichlorophosphine				
	$\Delta_{\text{v}}H$	(208–276)	29.2	260		[1964PET/BUR]
CCl ₂ F ₃ PS	[18799-78-5]	dichloro(trifluoromethylthio) phosphine				
	$\Delta_{\text{v}}H$	(293–363)	31.7	308	A	[1987STE/MAL, 1999DYK/SVO, 1960EME/PUG]
CCl ₂ O	[75-44-5]	phosgene				
	$\Delta_{\text{fus}}H$ (I)		5.74	145.4		
	$\Delta_{\text{fus}}H$ (II)		5.59	142.1		
	$\Delta_{\text{fus}}H$ (III)		4.73	139.2		[1960GIA/OTT]
	$\Delta_{\text{fus}}H$		5.73	145.3		[1948GIA/JON]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(280–341)	24.5	295	A	[1987STE/MAL]
	$\Delta_v H$	(240–281)	25.7	266	A	[1987STE/MAL]
	$\Delta_v H$	(338–410)	24.5	353	A	[1987STE/MAL]
	$\Delta_v H$	(406–455)	24.4	421	A	[1987STE/MAL]
	$\Delta_v H$	(215–248)	27.0	233		[1948GIA/JON]
	$\Delta_v H$	(180–273)	25.8	258		[1947STU]
CCl ₃ F	[75-69-4]	trichlorofluoromethane				
	$\Delta_{\text{fus}}H$		6.9	162.7		[1996DOM/HEA]
	$\Delta_v H$	(213–301)	28.5	228	A	[1987STE/MAL]
	$\Delta_v H$	(213–249)	28.2	234	A	[1987STE/MAL]
	$\Delta_v H$	(295–363)	25.6	310	A	[1987STE/MAL]
	$\Delta_v H$	(357–429)	24.7	372	A	[1987STE/MAL]
	$\Delta_v H$	(424–468)	25.1	439	A	[1987STE/MAL]
	$\Delta_v H$	(237–293)	27.3	251		[1979KUD/KUD]
	$\Delta_v H$	(237–293)	27.1	276		[1941OSB/GAR]
	$\Delta_v H$		25.2	290	C	[1941OSB/GAR]
	$\Delta_v H$	(244–334)	26.4	259		[1940BEN/MCH]
CCl ₃ F ₂ N	[24708-52-9]	N,N-difluoro-1,1,1-trichloromethylamine				
	$\Delta_v H$	(252–325)	33.4	267	I	[1987STE/MAL, 1970ZAB/SHR]
CCl ₃ F ₂ N	[33757-10-7]	N,1,1-trichloro-N,1-difluoromethylamine				
	$\Delta_v H$	(273–319)	27.8	296		[1971SWI/ZAB]
CCl ₃ F ₂ P	[1112-03-4]	difluoro(trichloromethyl) phosphine				
	$\Delta_{\text{sub}}H$	(264–283)	36.8	274		[1987STE/MAL]
	$\Delta_v H$	(289–313)	32.5	301	A	[1987STE/MAL]
CCl ₃ F ₄ P	[1184-80-1]	trichloromethyl tetrafluorophosphorane				
	$\Delta_v H$	(257–300)	10.4			[1965NIX]
CCl ₃ NO	[3711-49-7]	trichloronitrosomethane				
	$\Delta_v H$	(253–333)	32.4	268	A	[1987STE/MAL]
CCl ₃ NO ₂	[76-06-2]	trichloronitromethane				
	$\Delta_v H$	(273–333)	39.3	288	A	[1987STE/MAL]
	$\Delta_v H$	(301–449)	38.5	316	A	[1987STE/MAL, 1970DYK]
		(247–385)	40.0	262		[1947STU]
CCl ₄	[56-23-5]	carbon tetrachloride				
	$\Delta_{\text{ts}}H$		4.6	224.6		[1996DOM/HEA]
	$\Delta_{\text{fus}}H$		2.69	249		[1996DOM/HEA]
	$\Delta_{\text{sub}}H$		43.3	226	B	[1963BON]
	$\Delta_{\text{sub}}H$	(209–225)	38.8	217		[1960JON, 1948NIT/SEK]
	$\Delta_{\text{sub}}H$	(227–248)	37.9			[1948NIT/SEK]
	$\Delta_v H$	(349–416)	30.4	364	A	[1987STE/MAL]
	$\Delta_v H$	(412–497)	29.2	427	A	[1987STE/MAL]
	$\Delta_v H$	(494–555)	30.6	509	A	[1987STE/MAL]
	$\Delta_v H$		32.4	298	C	[1980MAJ/SVA]
	$\Delta_v H$	(262–349)	33.7	277	A, EB	[1987STE/MAL, 1972BOU/AIM]
	$\Delta_v H$	(293–351)	32.3	308		[1959HIL/MCD]
	$\Delta_v H$	(313–338)	31.7	325		[1953BAR/BRO]
CCl ₄ O ₂ S	[2547-61-7]	trichloromethanesulfonyl chloride				
	$\Delta_{\text{ts}}H$		7.1	227.4		

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{fus}}H$		7.46	418.5		[1994DOU/FEU]
CFIO	[1495-48-3] Δ_vH	carbonyl fluoride iodide (230–292)	26.1	277	A	[1987STE/MAL]
CFN	[1495-50-7] $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$ Δ_vH	cyanogen fluoride (147–191) (139–192) (133–203) (201–227)	28.9 24.4 29.3 22.4	176 166 168 214	A A	[1987STE/MAL, 1964FAW/LIP] [1947STU] [1931COS] [1987STE/MAL, 1964FAW/LIP, 1970DYK]
CFNO ₃ S	[1495-51-8] Δ_vH	sulfuryl fluoride isocyanate (294–335)	36.5	309	A	[1987STE/MAL, 1999DYK/SVO]
CFNO ₆ S ₂	[27931-74-4] Δ_vH	pyrosulfuryl fluoride isocyanate (330–405)	40.9	345	A	[1987STE/MAL, 1999DYK/SVO]
CFN ₃ O ₆	[1840-42-2] Δ_vH	fluorotrinitromethane (274–358)	34.2	289	A, T	[1987STE/MAL, 1966ZIM/ROB]
CF ₂ N ₂	[7127-18-6] Δ_vH	difluorocyanamide (179–198)	20.6	189		[1987STE/MAL, 1966MEY/FRA]
CF ₂ N ₂ OS	[19073-57-5] Δ_vH	cyanoimidodisulfuryl fluoride (262–354)	37.2	277	A	[1987STE/MAL, 1999DYK/SVO]
CF ₂ N ₂ O ₄	[1185-11-1] Δ_vH	difluorodinitromethane (283–310)	41.4	296	A	[1987STE/MAL, 1973PEP/LEB]
CF ₂ N ₂ S	[14453-41-9] Δ_vH	N-cyano-S,S-difluorosulfilimine (271–320)	44.1	286	A	[1987STE/MAL, 1999DYK/SVO]
CF ₂ O	[353-50-4] $\Delta_{\text{fus}}H$ $\Delta_{\text{sub}}H$ Δ_vH	carbonyl fluoride (130–159) (159–189)	6.7 23.2 20.0	161.9 145 174	 A	[1968PAC/REN] [1987STE/MAL, 1968PAC/REN] [1987STE/MAL]
CF ₂ O ₄ S	[7519-54-2] Δ_vH	fluoroformyl fluorosulfate (250–296)	27.3	281	A	[1987STE/MAL, 1999DYK/SVO]
CF ₂ S	[420-32-6] Δ_vH Δ_vH	thiocarbonyl fluoride (133–211) (178–211)	19.2 17.4	196 196	A A	[1987STE/MAL, 1970DYK] [1987STE/MAL, 1999DYK/SVO, 1962DOW]
CF ₃ I	[2314-97-8] Δ_vH	iodotrifluoromethane (188–296)	22.5	281	A	[1987STE/MAL, 1970DYK, 1948BAN/EME]
CF ₃ NO	[2368-32-3] Δ_vH	(difluoroamino) carbonyl fluoride (143–217)	21.6	202	A, MM	[1987STE/MAL, 1965FRA/SHR]
CF ₃ NO	[334-99-6] Δ_vH	trifluoronitrosomethane (141–174)	17.1	159	A	[1987STE/MAL]
CF ₃ NOS	[3855-41-2] Δ_vH	S,S-difluoro-N-(fluoroformyl)-sulfilimine (220–323)	37.3	235	A	[1987STE/MAL, 1999DYK/SVO]
CF ₃ NOS	[24892-54-4] Δ_vH	trifluoromethyl thionitrite (196–215)	25.8	205	T	[1969MAS]
CF ₃ NOS	[10564-49-5]	(N-sulfinyl)-trifluoromethylamine				

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(239–289)	27.0	274	A	[1987STE/MAL, 1999DYK/SVO]
CF ₃ NO ₂	[335-02-4]	trifluoronitromethane				
	$\Delta_v H$	(238–243)	21.6	240	A	[1987STE/MAL]
CF ₃ NO ₄	[50311-48-3]	(trifluoromethyl) peroxyxynitrate				
	$\Delta_v H$	(193–247)	24.8	232	A	[1987STE/MAL]
CF ₃ NO ₆ S ₂	[19252-48-3]	N-(fluoroformyl)-N,O-bis(fluorosulfonyl) hydroxylamine				
	$\Delta_v H$	(325–392)	36.3	340	A	[1987STE/MAL, 1999DYK/SVO]
CF ₄	[75-73-0]	carbon tetrafluoride				
	$\Delta_{\text{trs}}H$		1.71	76.27		
	$\Delta_{\text{fus}}H$		0.71	89.56		[1996DOM/HEA]
	$\Delta_{\text{sub}}H(\alpha)$	(76–90)	14.7	83		[1987STE/MAL, 70GEN/DUV]
	$\Delta_{\text{sub}}H(\beta)$	(70–76)	16.8	73		[1987STE/MAL, 1970GEN/DUV]
	$\Delta_{\text{sub}}H$	(86–89)	14.7	88		[1967SIM/KNO]
	$\Delta_{\text{sub}}H$		17.0	76		[1963BON]
	$\Delta_{\text{sub}}H$	(80–86)	14.0	83	A	[1933MEN/MOH]
	$\Delta_v H$	(195–227)	12.1	212	A	[1987STE/MAL]
	$\Delta_v H$	(89–163)	12.3	148	A	[1987STE/MAL]
	$\Delta_v H$	(160–197)	11.9	182	A	[1987STE/MAL]
	$\Delta_v H$	(116–146)	12.4	131		[1969SMI/PAC]
	$\Delta_v H$	(93–146)	12.8	131		[1933MEN/MOH, 1987STE/MAL]
CF ₄ N ₂ O	[815-10-1]	fluoro(trifluoromethyl) diimidoxide				
	$\Delta_v H$	(233–267)	27.7	252	A	[1987STE/MAL]
CF ₄ N ₂ O ₃ S ₂	[25523-80-2]	carbonylbis(imidosulfonyl fluoride)				
	$\Delta_v H$	(316–331)	41.3	323	A	[1987STE/MAL, 1999DYK/SVO]
CF ₄ O	[373-91-1]	hypofluorous acid trifluoromethyl ester				
	$\Delta_v H$	(153–194)	15.5	179	A	[1987STE/MAL, 1948KEL/CAD]
Note: The table in Ref. [1948KEL/CAD] gives the temperatures in °C; however, all of the equations and graphs in the article suggest that the temperature should be in Kelvin. We have assumed that the tabulated temperatures are in Kelvin; the results closely correspond to the entry in Ref. [1987STE/MAL].						
CF ₄ OS	[812-12-4]	trifluoromethyl sulfinyl fluoride				
	$\Delta_v H$	(204–271)	22.7	256	A, I	[1987STE/MAL, 1968RAT/SHR, 1970DYK, 1999DYK/SVO]
CF ₄ O ₂	[34511-13-2]	hydroperoxyfluoric acid trifluoromethyl ester				
	$\Delta_v H$	(156–203)	18.7	188	A	[1987STE/MAL]
CF ₄ O ₂ S	[335-05-7]	trifluoromethane sulfonyl fluoride				
	$\Delta_v H$	(226–249)	23.4	237	A	[1987STE/MAL, 1999DYK/SVO]
CF ₄ O ₃ S	[926-08-9]	trifluoromethyl fluorosulfonate				
	$\Delta_v H$	(194–269)	25.6	231		[1960VAN/CAD]
CF ₄ O ₄ S	[13990-10-8]	trifluoromethylperoxyfluorosulfonate				
	$\Delta_v H$	(233–286)	27.7	259		[1960VAN/CAD]
CF ₄ O ₅ S ₂	[21595-44-8]	fluorosulfonic acid trifluoromethane sulfonic acid anhydride				
	$\Delta_v H$	(308–338)	32.9	323	A	[1987STE/MAL, 1999DYK/SVO]
CF ₄ O ₆ S ₂	[6123-47-3]	trifluoromethyl fluorodisulfate				
	$\Delta_v H$	(292–351)	34.4	321		[1960VAN/CAD]
CF ₅ N	[335-01-3]	pentafluoromethyl amine				

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
	$\Delta_{\text{sub}}H$	(128–141)	18.6	135		[1987STE/MAL, 1951COA/HAR]
CF ₅ NO	[4217-93-0]	pentafluoromethoxyamine				
	$\Delta_{\text{v}}H$	(167–210)	18.5	195	A	[1987STE/MAL, 1965SHR/DUN]
CF ₅ OPS	[52752-66-6]	phosphorothionic difluoride, S-trifluoromethyl ester				
	$\Delta_{\text{v}}H$	(293–353)	23.1	323		[1999DYK/SVO]
CF ₅ OPS	[52752-66-6]	trifluoromethylthiophosphoryl difluoride				
	$\Delta_{\text{v}}H$		23.0			[1974SPE/SHR]
CF ₅ O ₃ P	[39125-42-3]	trifluoromethoxyphosphoryl difluoride				
	$\Delta_{\text{v}}H$	(225–264)	27.4	245		[1973BER/DES]
CF ₅ O ₃ P	[39125-42-3]	difluoroperoxyphosphoric acid trifluoromethyl ester				
	$\Delta_{\text{v}}H$	(241–280)	32.0	265	A	[1987STE/MAL, 1973BER/DES]
CF ₅ PS	[52752-65-5]	trifluoromethyl thiodifluorophosphine				
	$\Delta_{\text{v}}H$		24.3			[1974SPE/SHR]
CF ₆ N ₂ O ₂ S ₂	[20094-83-1]	N,N'-(difluoromethylene)bis imidosulfuryl fluoride				
	$\Delta_{\text{v}}H$	(283–308)	36.0	295		[1968GLE/VON]
CF ₆ N ₂ S ₂	[17686-45-2]	difluoromethane bis(S,S-difluorosulfilimine)				
	$\Delta_{\text{v}}H$	(230–313)	36.0	245	A	[1987STE/MAL, 1999DYK/SVO]
CF ₆ PS	[52752-65-5]	difluoro(trifluoromethylthio)phosphine				
	$\Delta_{\text{v}}H$	(293–353)	22.3	323		[1999DYK/SVO]
CF ₈ OS	[1873-23-0]	pentafluoro (trifluoromethoxy) sulfur				
	$\Delta_{\text{v}}H$	(217–262)	24.4	247	A	[1987STE/MAL, 1964DUN/CAD]
CF ₈ S	[373-80-8]	trifluoro(pentafluorothio)methane				
	$\Delta_{\text{v}}H$	(223–252)	20.2	253	I	[2001KUL/DES]
	$\Delta_{\text{v}}H$	(205–262)	23.8	247	A	[1987STE/MAL, 1999DYK/SVO]
CF ₉ NOS	[1840-45-5]	tetrafluoro(difluoroamino)(trifluoromethoxy) sulfur				
	$\Delta_{\text{v}}H$	(257–298)	28.7	272	A	[1987STE/MAL, 1964DUN/CAD2]
CF ₁₀ O ₅ S ₂	[60672-59-5]	[μ -(carono)diperoxato]decafluorodisulfur				
	$\Delta_{\text{v}}H$		38.1			[1976HOP/DES]
CIN	[506-78-5]	cyanogen iodide				
	$\Delta_{\text{sub}}H$	(337–426)	59.9	352	GSM	[1987STE/MAL, 1943KET/KRU]
	$\Delta_{\text{sub}}H$	(298–414)	58.6	356	A	[1947STU]
	$\Delta_{\text{sub}}H$	(337–426)	59.8 ± 0.4		GSM	[1943KET/KRU, 1970COX/PIL]
	$\Delta_{\text{sub}}H$	(278–374)	58.3	326		[1933YOS/STO]
	$\Delta_{\text{vap}}H$	(419–426)	40.0	423	A	[1987STE/MAL]
CN ₄ O ₈	[509-14-8]	tetranitromethane				
	$\Delta_{\text{sub}}H$	(255–286)	47.4	271		[1987STE/MAL, 1941SEK/NIT]
	$\Delta_{\text{v}}H$	(286–373)	43.1	301	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$	(313–373)	42.9	328	A	[1987STE/MAL, 1984BOU/FRI, 1952EDW]
	$\Delta_{\text{v}}H$	(273–313)	46.6	288		[1987STE/MAL, 1984BOU/FRI, 1949NIC]
CO	[630-08-0]	carbon monoxide				
	$\Delta_{\text{sub}}H$	(54–61)	7.6	58		[1987STE/MAL]
	$\Delta_{\text{sub}}H$	(51–68)	8.1	60	A	[1947STU]
	$\Delta_{\text{sub}}H$	(57–68)	7.9	62	A	[1931CRO/BIJ]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound					
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference	
		$\Delta_v H$	(68–108)	6.0	93	A	[1987STE/MAL]
		$\Delta_v H$	(69–83)	6.0	81		[1932CLA/GIA]
		$\Delta_v H$		6.0	81	C	[1932CLA/GIA]
COS	[463-58-1]	carbonyl sulfide					
		$\Delta_v H$	(161–284)	20.4	176		[1999DYK/SVO]
		$\Delta_v H$	(284–379)	18.3	299		[1999DYK/SVO]
		$\Delta_v H$	(140–224)	19.5	209	A	[1987STE/MAL]
		$\Delta_v H$		19.0 ± 0.1	214		[1939FRA/CLU]
		$\Delta_v H$	(162–224)	19.5	209		[1937KEM/GIA]
CO₂	[124-38-9]	carbon dioxide					
		$\Delta_{\text{sub}} H$	(198–216)	26.1	207	A	[1987STE/MAL]
		$\Delta_{\text{sub}} H$	(70–102)	27.2 ± 0.4		LE	[1974BRY/CAZ]
		$\Delta_{\text{sub}} H$	(179–198)	25.9	188		[1956AMB]
		$\Delta_{\text{sub}} H$	(139–195)	26.3	167	A	[1947STU]
		$\Delta_{\text{sub}} H$	(154–196)	25.2	195		[1937GIA/EGA]
		$\Delta_v H$	(273–304)	16.7	288	A	[1987STE/MAL]
		$\Delta_v H$	(216–273)	16.4	258	A	[1987STE/MAL]
		$\Delta_v H$	(267–303)	16.5	282		[1972BOU/AIM]
CS₂	[75-15-0]	carbon disulfide					
		$\Delta_v H$	(255–354)	28.7	270		[1999DYK/SVO]
		$\Delta_v H$	(354–552)	27.1	369		[1999DYK/SVO]
		$\Delta_v H$	(260–353)	28.5	275	A	[1987STE/MAL]
		$\Delta_v H$	(338–408)	27.4	353	A	[1987STE/MAL]
		$\Delta_v H$	(388–497)	27.0	403	A	[1987STE/MAL]
		$\Delta_v H$	(490–533)	28.7	505	A	[1987STE/MAL]
		$\Delta_v H$	(255–318)	28.7	270	EB	[1972BOU/AIM, 1987STE/MAL]
		$\Delta_v H$	(277–353)	28.1	292	EB	[1962WAD/SMI]
		$\Delta_v H$		28.1 ± 0.1	282	C	[1962WAD/SMI]
		$\Delta_v H$		27.5 ± 0.1	298	C	[1962WAD/SMI]
		$\Delta_v H$		26.7 ± 0.1	319	C	[1962WAD/SMI]
		$\Delta_v H$		27.7	298		[1961GOO/LAC]
		$\Delta_v H$	(303–358)	27.6	318		[1946THO]
CHBrF₂	[1511-62-2]	bromodifluoromethane					
		$\Delta_v H$	(194–259)	24.0	244	A	[1987STE/MAL]
		$\Delta_v H$	(194–288)	24.7	209		[1979KUD/KUD]
CHBr₃	[75-25-2]	tribromomethane					
		$\Delta_{\text{fus}} H$		11.09	281.5		[1987KAF/DOR]
		$\Delta_v H$		46.1 ± 0.1	298	C	[1972LAY/WAD]
		$\Delta_v H$	(320–412)	42.3	335	EB	[1972BOU/AIM, 1979KUD/KUD]
		$\Delta_v H$	(303–373)	44.0	318		[1941KIR/SIT, 1984BOU/FRI]
CHClF₂	[75-45-6]	chlorodifluoromethane					
		$\Delta_{\text{trs}} H$		0.07	59		
		$\Delta_{\text{fus}} H$		4.12	115.7		[1996DOM/HEA]
		$\Delta_v H$	(275–327)	20.0	290	A	[1987STE/MAL]
		$\Delta_v H$	(170–233)	21.3	218	A	[1987STE/MAL]
		$\Delta_v H$	(230–275)	20.4	260	A	[1987STE/MAL]
		$\Delta_v H$	(324–366)	20.1	339	A	[1987STE/MAL]
		$\Delta_v H$	(194–310)	21.8	209		[1979KUD/KUD]
		$\Delta_v H$	(229–236)	21.0	232		[1964KLE]
		$\Delta_v H$		20.2	232	C	[1957NEI/WHI]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
CHCl₂F	[75-43-4]	dichlorofluoromethane				
	$\Delta_{\text{v}}H$	(225–282)	26.1	267	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$	(279–344)	25.3	294	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$	(341–399)	24.2	356	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$	(397–450)	24.1	412	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$	(229–236)	U 20.9	233		[1964KLE]
	$\Delta_{\text{v}}H$	(181–282)	26.2	267		[1947STU]
$\Delta_{\text{v}}H$	(244–317)	36.7	259		[1940BEN/MCH]	
CHCl₂FO₃S	[42016-50-2]	fluorosulfuric acid, dichloromethyl ester				
	$\Delta_{\text{v}}H$	(275–293)	36.2	284	A	[1987STE/MAL, 1999DYK/SVO]
CHCl₃	[67-66-3]	chloroform				
	$\Delta_{\text{fus}}H$		8.8	209.6		[1991ACR]
	$\Delta_{\text{v}}H$	(306–427)	30.8	321		[1995CHE/WAN]
	$\Delta_{\text{v}}H$	(227–269)	31.8	254	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$	(333–416)	30.4	348	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$	(410–481)	28.9	425	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$	(479–523)	30.1	494	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$		31.1	298	C	[1980MAJ/SVA]
	$\Delta_{\text{v}}H$	(260–333)	32.5	275	EB	[1972BOU/AIM]
$\Delta_{\text{v}}H$	(215–334)	35.0	230		[1947STU]	
$\Delta_{\text{v}}H$	(308–333)	30.9	320		[1938SCT/RAY]	
CHFI₂	[1493-01-2]	diiodofluoromethane				
	$\Delta_{\text{v}}H$	(299–332)	32.9	314	A	[1987STE/MAL, 1979KUD/KUD, 1970DYK]
CHFN₂O₄	[7182-87-8]	fluorodinitromethane				
	$\Delta_{\text{v}}H$	(298–338)	43.6	313	A	[1987STE/MAL]
CHFO	[1493-02-3]	formyl fluoride				
	$\Delta_{\text{v}}H$	(178–235)	24.4	220	A	[1987STE/MAL, 1964FIS/BUC, 1970DYK]
CHF₂I	[1493-03-4]	difluoroiodomethane				
	$\Delta_{\text{v}}H$	(227–287)	26.0	272	A	[1987STE/MAL, 1979KUD/KUD, 1970DYK]
CHF₃	[75-46-7]	trifluoromethane				
	$\Delta_{\text{fus}}H$		4.06	118		[1996DOM/HEA]
	$\Delta_{\text{sub}}H$	(89–118)	25.6	103		[1987STE/MAL]
	$\Delta_{\text{v}}H$	(138–190)	18.1	175	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$	(198–298)	16.8	213	A	[1987STE/MAL]
$\Delta_{\text{v}}H$	(146–192)	18.0	177		[1962VAL/BRO]	
CHF₃O₂	[16156-36-8]	trifluoromethyl hydroperoxide				
	$\Delta_{\text{v}}H$	(248–285)	30.9	270	A	[1987STE/MAL]
CHF₃O₃S	[1493-13-6]	trifluoromethylsulfonic acid				
	$\Delta_{\text{v}}H$	(354–435)	47.7	369	A	[1987STE/MAL, 1999DYK/SVO]
CHF₃S	[1493-15-8]	trifluoromethanethiol				
	$\Delta_{\text{fus}}H$		4.93	116		[1996DOM/HEA]
	$\Delta_{\text{v}}H$	(167–236)	21.8	183		[1999DYK/SVO]
	$\Delta_{\text{v}}H$	(167–236)	21.0	221	A	[1987STE/MAL, 1999DYK/SVO]
CHF₇S	[420-67-7]	(difluoromethyl) sulfur pentafluoride				
	$\Delta_{\text{v}}H$	(221–292)	27.5	237		[1999DYK/SVO]
	$\Delta_{\text{v}}H$	(221–293)	25.6	278	A	[1987STE/MAL, 1999DYK/SVO]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
CHI ₃	[75-47-8]	iodoform				
	$\Delta_{\text{sub}}H$	(308–365)	69.9	323		[1943NIT/SEK]
CHN	[74-90-8]	hydrogen cyanide				
	$\Delta_{\text{sub}}H$	(236–259)	35.6	248		[1987STE/MAL]
	$\Delta_{\text{sub}}H$	(202–254)	37.6	228	A	[1947STU]
	Δ_vH	(259–299)	28.1	274	A	[1987STE/MAL]
	Δ_vH	(298–457)	27.8	313	A	[1987STE/MAL]
	Δ_vH	(257–315)	28.1	272		[1975IWA/DAT]
	Δ_vH	(259–294)	28.0	277		[1934LEW/SCH]
	Δ_vH	(257–319)	28.1	272		[1926SIN/HAR]
CDN	[3017-23-0]	deuterium cyanide				
	Δ_vH	(182–282)	26.2	267		[1947STU]
	Δ_vH	(265–293)	27.6	279		[1934LEW/SCH]
CHNO	[420-05-3]	cyanic acid				
	Δ_vH	(233–268)	30.7	253	A	[1987STE/MAL]
		(197–267)	NA			[1938LIN]
CHNS	[463-56-9]	thiocyanic acid				
	Δ_vH	(278–396)	28.0	293	A	[1987STE/MAL]
CHN ₃ O ₆	[517-25-9]	trinitromethane				
	$\Delta_{\text{sub}}H$		45.2 ± 2.1	298		[1999MIR/VOR]
	$\Delta_{\text{sub}}H$		54.8 ± 4.2			[1970BON/CAT]
	$\Delta_{\text{sub}}H$		46.7 ± 0.4			[1967MIR/LEB, 1970COX/PIL, 1977PED/RYL]
	Δ_vH	(290–317)	32.6	303	A	[1987STE/MAL, 1967MIR/LEB]
CH ₂ BrCl	[74-95-7]	bromochloromethane				
	Δ_vH	(226–341)	42.0	241	A	[1987STE/MAL]
	Δ_vH	(289–341)	33.5	304		[1959MCD/SHR, 1979KUD/KUD]
CH ₂ Br ₂	[74-95-3]	dibromomethane				
	Δ_vH	(273–373)	36.5	288	C	[1979KUD/KUD]
	Δ_vH		37.0 ± 0.1	298	A, E	[1972LAY/WAD]
	Δ_vH	(290–409)	37.2	305		[1987STE/MAL, 1956MAN, 1970DYK]
	Δ_vH	(238–371)	37.8	253		[1947STU]
CH ₂ ClF	[593-70-4]	chlorofluoromethane				
	Δ_vH	(140–264)	23.3	249	A	[1987STE/MAL, 1970DYK]
CH ₂ Cl ₂	[75-09-2]	dichloromethane				
	$\Delta_{\text{fus}}H$		6.16	178.2		[1996DOM/HEA]
	Δ_vH		30.6 ± 0.1	298	C	[1989AN/HU]
	Δ_vH	(311–383)	29.0	326	A	[1987STE/MAL]
	Δ_vH		28.8	298	C	[1980MAJ/SVA]
	Δ_vH	(264–311)	30.3	279	EB	[1972BOU/AIM]
	Δ_vH	(303–313)	29.2	308		[1960MUE/IGN]
	Δ_vH	(233–313)	30.2	248		[1948GAN/JUN]
	Δ_vH		NA		[1946DZU]	
	Δ_vH	(186–312)	29.4			[1927PER]
CH ₂ F ₂	[75-10-5]	difluoromethane				
	$\Delta_{\text{fus}}H$		4.36	136.4		[1996LUE/MAG]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(149–245)	20.6	230		[1987KAN/OI]
	$\Delta_v H$	(256–321)	19.9	271	A	[1987STE/MAL]
	$\Delta_v H$	(191–222)	21.2	207	A	[1987STE/MAL]
	$\Delta_v H$	(191–258)	20.3	243	A	[1987STE/MAL]
	$\Delta_v H$	(316–351)	20.3	331	A	[1987STE/MAL]
	$\Delta_v H$	(191–221)	21.2	206		[1968MAL/MEU]
	$\Delta_v H$	(191–242)	20.6	227		[1968MAL/MEU]
CH₂F₃NS	[1512-33-0]	1,1,1-trifluoromethanesulfenamide				
	$\Delta_v H$	(218–291)	34.1	276	A	[1987STE/MAL, 1999DYK/SVO, 1960EME/NAB]
CH₂I₂	[75-11-6]	diiodomethane				
	$\Delta_{\text{fus}} H$		12.05	279.2		[1987KAF/DOR]
	$\Delta_v H$		45.6	298	GC	[1994CAR/LAY]
	$\Delta_v H$		49.0	298	C	[1987FUC/CHA]
	$\Delta_v H$	(293–455)	48.8	307		[1979KUD/KUD]
	$\Delta_v H$	(356–505)	45.4	371	A	[1987STE/MAL, 1970DYK]
CH₂N₂	[420-04-2]	cyanamide				
	$\Delta_{\text{fus}} H$		8.76	317.2		[1991ACR]
	$\Delta_{\text{sub}} H$	(227–289)	75.9	290	TE,ME	[1983DEW/VAN]
	$\Delta_{\text{sub}} H$		75.2	298		[1983DEW/VAN]
CH₂N₄	[288-94-8]	tetrazole				
	$\Delta_{\text{fus}} H$		17.7	432.1		[1990KOZ/SIM3]
	$\Delta_{\text{sub}} H$		88.16	353	C	[1993KAB/KOZ]
	$\Delta_{\text{sub}} H$		87.8 ± 1.4	369	ME	[1993KAB/KOZ]
	$\Delta_{\text{sub}} H$	(333–404)	88.0 ± 1.6		ME	[1990KOZ/SIM]
	$\Delta_{\text{sub}} H$	(333–363)	97.5 ± 4.2	348	ME	[1951MCE/RIG, 1970COX/PIL]
CH₂O	[50-00-0]	formaldehyde				
	$\Delta_{\text{fus}} H$		7.53	155		[1998VAS/LEB]
	$\Delta_v H$	(184–251)	24.3	236	A	[1987STE/MAL]
	$\Delta_v H$	(173–251)	24.2	236		[1935SPE/WIL, 1987STE/MAL]
CH₂O₂	[64-18-6]	formic acid				
	$\Delta_{\text{sub}} H$	(268–281)	60.5	275		[1987STE/MAL]
	$\Delta_{\text{sub}} H$	(203–218)	62.1 ± 1	213	TE,ME	[1978CAL/CAL]
	$\Delta_{\text{sub}} H$	(265–268)	60.7	266		[1930COO, 1960JON]
	$\Delta_{\text{sub}} H$	(253–275)	60.1	264	A	[1947STU]
	$\Delta_v H$	(300–392)	35.2	315	EB	[1987AMB/GHI3]
	$\Delta_v H$	(283–384)	36.0	298	A	[1987STE/MAL]
	$\Delta_v H$ (monomer)		20.1 ± 0.1	298	C	[1970KON/WAD]
	$\Delta_v H$		46.3 ± 0.5	298	C	[1970KON/WAD]
	$\Delta_v H$	(310–374)	35.2	325		[1949DRE/SHR, 1949DRE/MAR]
	$\Delta_v H$		19.9	298		[1941STO/FIS]
	$\Delta_v H$		29.6	303		[1934CAM/CAM]
	$\Delta_v H$	(273–373)	20.3	315		[1930COO]
	$\Delta_v H$	(273–373)	20.9	338		[1930COO]
	$\Delta_v H$		20.4	315	C	[1930COO]
	$\Delta_v H$		21.1	338	C	[1930COO]
	$\Delta_v H$	(273–307)	36.8	288		[1994KAH]
	$\Delta_v H$	(295–374)	47.7	374		[1883KAH]
(CH₂O₂)₂	[14523-98-9]	formic acid dimer				

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
	$\Delta_{\text{sub}}H$	(203–218)	64.1 ± 1	213	TE,ME	[1978CAL/CAL]
CH₃Br	[74-83-9]	methyl bromide				
	$\Delta_{\text{trs}}H$		0.47	173.8		
	$\Delta_{\text{fus}}H$		5.98	179.5		[1996DOM/HEA, 1987KAF/DOR]
	$\Delta_{\text{v}}H$	(223–278)	25.8	238		[1979KUD/KUD]
	$\Delta_{\text{v}}H$	(201–296)	24.6	281	A, E	[1987STE/MAL, 1961LI/ROS]
	$\Delta_{\text{v}}H$	(203–277)	25.2	262		[1947BEE/JUN]
	$\Delta_{\text{v}}H$	(203–278)	25.3	263		[1938EGA/KEM]
CH₃Cl	[74-87-31]	methyl chloride				
	$\Delta_{\text{fus}}H$		6.42	174.5		[1996DOM/HEA]
	$\Delta_{\text{sub}}H$	(130–172)	31.6 ± 0.1	151		[1995BAH/DUP]
	$\Delta_{\text{sub}}H$		28.0		B	[1940MES/AST]
	$\Delta_{\text{v}}H$	(247–310)	22.0	262	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$	(368–416)	21.8	383	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$	(308–373)	21.0	323	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$	(198–278)	22.0	263		[1948GAN/JUN]
	$\Delta_{\text{v}}H$	(183–250)	22.7	235		[1947BEE/JUN]
	$\Delta_{\text{v}}H$	(191–249)	23.5	206		[1946THO]
	$\Delta_{\text{v}}H$	(192–249)	22.6	234		[1940MES/AST]
	$\Delta_{\text{v}}H$		20.1	293	C	[1926YAT]
CH₃ClFOP	[753-71-9]	methylphosphonyl chlorofluoride				
	$\Delta_{\text{fus}}H$		11.85	250.7	AC	[1964FUR/REI]
CH₃Cl₂P	[676-83-5]	dichloromethyl phosphine				
	$\Delta_{\text{v}}H$	(229–297)	35.5	282	A	[1987STE/MAL, 1963HOL/WAG]
CH₃Cl₂OP	[676-97-1]	methylphosphonic dichloride				
	$\Delta_{\text{fus}}H$		18.08	306.1		[1964FUR/REI]
	$\Delta_{\text{sub}}H$		62.3			[1970COX/PIL, 1955NEA/WIL]
CH₃F	[593-53-3]	methyl fluoride				
	$\Delta_{\text{v}}H$	(205–242)	16.9	227	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$	(240–288)	16.9	273	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$	(133–211)	17.9	172		[1983OI/SHU]
	$\Delta_{\text{v}}H$	(141–208)	17.1	193	A, E	[1987STE/MAL, 1961LI/ROS, 1970DYK]
	$\Delta_{\text{v}}H$	(165–217)	16.4	202		[1987STE/MAL, 1948MIC/WAS, 1984BOU/FRI]
	$\Delta_{\text{v}}H$	(170–197)	17.7	183	A	[1987STE/MAL, 19MOL/BAT, 1984BOU/FRI]
CH₃F₂N	[753-58-2]	N,N-difluoromethylamine				
	$\Delta_{\text{v}}H$	(203–257)	23.5	242	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$		22.9	257		[1960FRA]
CH₃F₂NS	[758-20-3]	methylimidodisulfurous difluoride				
	$\Delta_{\text{v}}H$	(194–258)	28.7	226		[1999DYK/SVO]
	$\Delta_{\text{v}}H$	(194–258)	28.6	243	A	[1987STE/MAL, 1999DYK/SVO]
CH₃F₂P	[753-59-3]	difluoromethyl phosphine				
	$\Delta_{\text{v}}H$	(174–236)	23.4	221	A	[1987STE/MAL]
CH₃F₂OP	[676-99-3]	methylphosphonyl difluoride				
	$\Delta_{\text{fus}}H$		11.88	236.3		[1964FUR/REI]
CH₃F₂OPS	[25237-37-0]	difluorothiophosphoric, S-methyl ester				
	$\Delta_{\text{v}}H$	(236–298)	31.2	251	A	[1987STE/MAL, 1999DYK/SVO]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
CH₃F₂PS₂	[21348-13-0] $\Delta_v H$	difluorodithiophosphoric acid, methyl ester (253–298)	39.0	268	A	[1987STE/MAL, 1999DYK/SVO]
CH₃F₄NP₂S₂	[25741-62-2] $\Delta_v H$	N,N-bis(difluorothiophosphoral) methylamine (273–325)	38.7	288	A	[1987STE/MAL, 1999DYK/SVO]
CH₃I	[74-88-4] $\Delta_{\text{fus}} H$	methyl iodide	9.12	206.8		[1982WRE/VIK]
	$\Delta_{\text{sub}} H$	(176–227)	40.2 ± 0.4	191	VG	[1982WRE/VIK]
	$\Delta_{\text{sub}} H$		U 69.9			[1943NIT/SEK, 1960JON]
	$\Delta_v H$	(228–337)	30.4	243	A	[1987STE/MAL]
	$\Delta_v H$	(315–502)	26.5	330	A	[1987STE/MAL]
	$\Delta_v H$	(208–227)	31.1	217		[1982WRE/VIK]
	$\Delta_v H$	(259–314)	29.2	274	EB	[1972BOU/AIM, 1979KUD/KUD]
	$\Delta_v H$	(218–315)	30.4	233		[1947STU]
	$\Delta_v H$	(273–307)	28.2	288		[1936EWE]
CH₃NO	[75-12-7] $\Delta_{\text{fus}} H$	formamide	7.98	275.7		[1996DOM/HEA]
	$\Delta_{\text{fus}} H$		8.67	275.6		[1983DEW/DEK]
	$\Delta_{\text{sub}} H$	(251–273)	72.4	264	TE,ME	[1983DEW/VAN]
	$\Delta_{\text{sub}} H$		71.7	298		[1983DEW/VAN]
	$\Delta_{\text{sub}} H$		71.7	276		[1979DAA/VAN]
	$\Delta_v H$	(293–377)	70.8	308	A	[1987STE/MAL]
	$\Delta_v H$	(415–466)	61.2	430	A	[1987STE/MAL]
	$\Delta_v H$		60.2	298	A	[1985BAR/CAS, 1985MAJ/SVO]
	$\Delta_v H$	(343–483)	64.0	358		[1947STU]
CH₃NOS	[4291-05-8] $\Delta_v H$	N-sulfinyl methanamine (252–277)	31.8	264	A	[1987STE/MAL, 1999DYK/SVO]
CH₃NO₂	[624-91-9] $\Delta_v H$	methyl nitrite (218–273)	22.1	258	A	[1987STE/MAL]
	$\Delta_v H$	(154–225)	26.2	190		[1982ROO]
	$\Delta_v H$		22.6 ± 0.2			[1958GRA/PRA]
CH₃NO₂	[75-52-2] $\Delta_{\text{fus}} H$	nitromethane	9.7	244.8		[1996DOM/HEA]
	$\Delta_v H$		38.5 ± 0.4	298		[1999MIR/VOR2]
	$\Delta_v H$	(313–353)	37.2	298	CGC	[1995CHI/HOS]
	$\Delta_v H$	(405–476)	35.2	420	A	[1987STE/MAL, 1967BER/WES]
	$\Delta_v H$	(328–410)	36.8	343	A	[1987STE/MAL, 1954MCC/SCO]
	$\Delta_v H$		37.2 ± 0.1	318	C	[1954MCC/SCO]
	$\Delta_v H$		36.3 ± 0.1	335	C	[1954MCC/SCO]
	$\Delta_v H$		35.2 ± 0.1	353	C	[1954MCC/SCO]
	$\Delta_v H$		34.0 ± 0.1	374	C	[1954MCC/SCO]
	$\Delta_v H$	(283–373)	38.0 ± 0.4	298	ZG	[1949HOL/DOR]
	$\Delta_v H$		38.3 ± 0.1	298	C	[1947JON/GIA]
CH₃NO₃	[598-58-3] $\Delta_{\text{fus}} H$	methyl nitrate	8.24	190.2		[1996DOM/HEA]
	$\Delta_v H$	(273–303)	34.8	288	A	[1987STE/MAL]
CH₃N₅	[4418-61-5] $\Delta_{\text{sub}} H$	5-aminotetrazole (383–443)	112.6 ± 1.2		ME	[1990KOZ/SIM]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
CH₄	[74-82-8]	methane				
	$\Delta_{\text{sub}}H$	(53–91)	9.7	72		[1987STE/MAL]
	$\Delta_{\text{sub}}H$	(54–90)	9.2	72		[1963BON, 1955ARM/BRI]
	$\Delta_{\text{sub}}H$	(79–89)	10.0	84		[1960JON]
	$\Delta_{\text{sub}}H$	(48–78)	9.7	63	A,MS	[1951TIC/LOS]
	$\Delta_{\text{sub}}H$	(67–88)	9.62	77	A	[1947STU]
	$\Delta_{\text{v}}H$	(90–120)	8.6	105	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$	(115–149)	8.4	134	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$	(148–189)	8.7	174	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$	(91–127)	8.6	112		[1972PRY/GOO, 1984BOU/FRI]
	$\Delta_{\text{v}}H$	(91–190)	8.5	175		[1972PRY/GOO]
	$\Delta_{\text{v}}H$		8.1	137		[1971WIL/ZWO]
	$\Delta_{\text{v}}H$	(100–190)	8.6	175		[1970AMB/COU]
	$\Delta_{\text{v}}H$		8.2	112	C	[1961HES/WHI]
	$\Delta_{\text{v}}H$		7.5	130	C	[1961HES/WHI]
	$\Delta_{\text{v}}H$		5.9	160	C	[1961HES/WHI]
	$\Delta_{\text{v}}H$		4.0	180	C	[1961HES/WHI]
	$\Delta_{\text{v}}H$	(109–189)	8.5	149		[1961HES/WHI]
$\Delta_{\text{v}}H$		8.5 ± 0.1	99		[1939FRA/CLU]	
$\Delta_{\text{v}}H$	(92–110)	8.6	101		[1921STO/HEN, 1984BOU/FRI]	
CH₄F₂NPS	[31411-30-0]	difluorothiophosphoric acid, N-methylamide				
	$\Delta_{\text{v}}H$	(273–325)	39.1	288	A	[1987STE/MAL, 1999DYK/SVO]
CH₄N₂	[12211-52-8]	ammonium cyanide				
	$\Delta_{\text{v}}H$	(222–305)	47.1	237		[1947STU]
CH₄N₂	[26981-93-1]	methyl diazene				
	$\Delta_{\text{v}}H$	(195–236)	27.5	221	A	[1987STE/MAL]
CH₄N₂O	[57-13-6]	urea				
	$\Delta_{\text{fus}}H$		14.6	407.2	DSC	[1999RAI/RAI]
	$\Delta_{\text{fus}}H$		14.6	406.7	DSC	[1998RAI/RAI]
	$\Delta_{\text{fus}}H$		13.6	405.2	DSC	[1998JAM/PAL]
	$\Delta_{\text{fus}}H$		15.03	407.9	DSC	[1995FER/DEL]
	$\Delta_{\text{fus}}H$		12.93	408.1		[1990KAB/MIR2]
	$\Delta_{\text{fus}}H$		13.9	405.8		[1986KOZ/DAL]
	$\Delta_{\text{sub}}H$	(358–402)	95.5 ± 0.3	298	GS	[2006EME/KAB]
	$\Delta_{\text{sub}}H$	(329–403)	94.6 ± 2.2	370	ME	[2003ZAI/KAB]
	$\Delta_{\text{sub}}H$	(329–403)	95.1 ± 2.2	350	ME	[2003ZAI/KAB]
	$\Delta_{\text{sub}}H$		94.6 ± 0.5	350	C	[2003ZAI/KAB]
	$\Delta_{\text{sub}}H$		90.9	381		[1987FER/DEL2]
	$\Delta_{\text{sub}}H$	(345–368)	87.7	357		[1987STE/MAL]
	$\Delta_{\text{sub}}H$	(338–362)	96.9	351	TE,ME	[1983DEW/VAN]
	$\Delta_{\text{sub}}H$		98.6	298		[1983DEW/VAN]
$\Delta_{\text{sub}}H$		95.4	361		[1978TRI/VOO]	
$\Delta_{\text{sub}}H$	(345–368)	87.9 ± 2.1	356		[1956SUZ/ONI, 1960JON, 1970COX/PIL]	
$\Delta_{\text{sub}}H$		88.2	357		[1953BRA/CLE2, 1983DEW/VAN]	
CH₄N₂S	[62-56-5]	thiourea				
	$\Delta_{\text{trs}}H$		0.026	169		
	$\Delta_{\text{trs}}H$		0.113	200	AC	[1993IGA/LOP]
	$\Delta_{\text{fus}}H$		15.64	444.7		[2000DEL/JOZ]
	$\Delta_{\text{trs}}H$		0.026	169		
	$\Delta_{\text{fus}}H$		12.6	452.2	AC	[1993IGA/LOP]
					[1994DOU/FUE]	

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	Enthalpy							
		$\Delta_{\text{fus}}H$			12.55	452.2		[1994KIM/LEE]
		$\Delta_{\text{sub}}H$			112.0 ± 2	298	ME	[2000DEL/JOZ]
		$\Delta_{\text{sub}}H$			109 ± 2.0	408	TE	[1994FER/MAR]
		$\Delta_{\text{sub}}H$			111 ± 3.0	298		[1994FER/MAR]
		$\Delta_{\text{sub}}H$		(378–396)	103.9 ± 0.3	387	ME	[1994TOR/HER]
		$\Delta_{\text{sub}}H$		(368–395)	106.6	384	TE,ME	[1983DEW/VAN]
		$\Delta_{\text{sub}}H$			107.6	298		[1983DEW/VAN]
		$\Delta_{\text{sub}}H$			112 ± 1.5	298	C	[1982TOR/SAB]
		$\Delta_{\text{sub}}H$			93.7 ± 10			[1975BAG/AND]
CH₄N₂O₂	[1111-78-0]		ammonium carbamate					
		Δ_vH		(247–331)	54.1	262		[1947STU]
CH₄N₄O₂	[556-88-7]		nitroguanidine					
		$\Delta_{\text{sub}}H$		(402–473)	142.7 ± 2.0	298	ME	[1978CUN/PAL]
CH₄N₄O₄	[14168-44-6]		N,N'-dinitro-diaminomethane					
		$\Delta_{\text{fus}}H$			35.85	371		[1987OYU/BR1]
CH₄O	[67-56-1]		methanol					
		$\Delta_{\text{us}}H$			0.59	161.1		
		$\Delta_{\text{fus}}H$			3.18	175.3		[1996DOM/HEA]
		Δ_vH		(298–333)	38.0	298		[2004NAS/ZIM]
		Δ_vH			34.3			[1999FAT]
		Δ_vH		(175–273)	39.2	258	A	[1987STE/MAL]
		Δ_vH		(338–487)	36.9	353	A	[1987STE/MAL]
		Δ_vH		(188–228)	43.7	213	A	[1987STE/MAL]
		Δ_vH		(224–290)	38.9	275	A	[1987STE/MAL]
		Δ_vH		(285–345)	38.3	300	A	[1987STE/MAL]
		Δ_vH		(335–376)	37.0	350	A	[1987STE/MAL]
		Δ_vH		(373–458)	36.1	388	A	[1987STE/MAL]
		Δ_vH		(453–513)	35.1	468	A	[1987STE/MAL]
		Δ_vH			32.7	373	C	[1986YER/WOR]
		Δ_vH			28.1	423	C	[1986YER/WOR]
		Δ_vH			20.6	473	C	[1986YER/WOR]
		Δ_vH			7.4	510	C	[1986YER/WOR]
		Δ_vH		(316–336)	37.5	331	EB	[1984CER/BOU]
		Δ_vH		(243–303)	37.8	298		[1983SCH/STR]
		Δ_vH		(288–337)	38.3	303		[1974GIB/CRE, 1984BOU/FRI]
		Δ_vH		(337–383)	37.0	352		[1973WIL/ZWO]
		Δ_vH			37.4 ± 0.1	298	C	[1973SVO/VES]
		Δ_vH			36.7 ± 0.1	313	C	[1973SVO/VES]
		Δ_vH			36.2 ± 0.1	323	C	[1973SVO/VES]
		Δ_vH			35.6 ± 0.1	333	C	[1973SVO/VES]
		Δ_vH			35.3 ± 0.1	338	C	[1973SVO/VES]
		Δ_vH			34.7 ± 0.1	343	C	[1973SVO/VES]
		Δ_vH			35.2 ± 0.1	338	C	[1973COU/LEE]
		Δ_vH			35.6 ± 0.1	331	C	[1973COU/LEE]
		Δ_vH			36.2 ± 0.1	321	C	[1973COU/LEE]
		Δ_vH			37.0 ± 0.1	306	C	[1973COU/LEE]
		Δ_vH		(275–336)	38.7	290	EB	[1972BOU/AIM, 1987STE/MAL]
		Δ_vH			37.43 ± 0.02	298	C	[1971POL/BEN]
		Δ_vH		(288–357)	38.3	303	EB	[1970AMB/SPR]
		Δ_vH		(353–483)	36.3	368		[1967HIR/SUD]
		Δ_vH			37.3 ± 0.1	298	C	[1966WAD]
		Δ_vH			37.7 ± 0.1	298	C	[1963MCC/LAI]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	Δ_vH	(278–323)	38.4	293		[1960KLY/MIS]
CH ₄ O ₂	[3031-73-0]	methyl hydroperoxide				
	Δ_vH	(253–313)	37.7	268	A	[1987STE/MAL, 1951EGE/EMT]
CH ₄ O ₃ S	[75-75-2]	methanesulfonic acid				
	Δ_vH	(395–440)	73.9	410	A	[1987STE/MAL, 1999DYK/SVO]
CH ₄ S	[74-93-1]	methyl mercaptan				
	$\Delta_{\text{trs}}H$		2.2	137.6		
	$\Delta_{\text{fus}}H$		5.9	150.2		[1996DOM/HEA]
	Δ_vH	(208–298)	27.2	223		[1999DYK/SVO]
	Δ_vH	(267–359)	25.2	359	A	[1987STE/MAL]
	Δ_vH	(221–283)	25.7	268	A	[1987STE/MAL]
	Δ_vH	(345–424)	23.7	360	A	[1987STE/MAL]
	Δ_vH	(414–470)	24.2	429	A	[1987STE/MAL]
	Δ_vH		23.8	298		[1971WIL/ZWO]
	Δ_vH	(222–279)	25.8	264		[1987STE/MAL, 1942RUS/OSB]
CH ₅ N	[74-89-5]	methylamine				
	$\Delta_{\text{fus}}H$		6.13	179.7		[1996DOM/HEA]
	Δ_vH	(319–381)	24.8	334	A	[1987STE/MAL]
	Δ_vH	(373–430)	23.5	388	A	[1987STE/MAL]
	Δ_vH	(263–329)	26.1	278	A	[1987STE/MAL]
	Δ_vH	(223–273)	27.2	258	A	[1987STE/MAL, 1970DYK]
	Δ_vH	(190–267)	27.4	252		[1937AST/SIL, 1984BOU/FRI]
CH ₅ NO	[593-77-1]	N-methylhydroxylamine				
	$\Delta_{\text{sub}}H$	(273–308)	56.6	288		[1987STE/MAL, 1957BIS/PAR]
	Δ_vH	(293–338)	49.7	308	A	[1987STE/MAL, 1970DYK]
	Δ_vH	(313–338)	49.3	325	A	[1987STE/MAL, 1957BIS/PAR, 1984BOU/FRI]
CH ₅ NO	[67-62-9]	O-methylhydroxylamine				
	Δ_vH	(228–322)	36.9	243	A	[1987STE/MAL]
	Δ_vH	(210–321)	38.0	225		[1957BIS/PAR, 1984BOU/FRI]
CH ₅ N ₃ O	[758-19-0]	1-methyl-1-nitrosohydrazine				
	$\Delta_{\text{sub}}H$		79.5 ± 0.4	298		[1998LEB/CHI]
CH ₅ N ₃ S	[79-19-6]	thiosemicarbazide				
	$\Delta_{\text{sub}}H$		125.8 ± 1.5	298	C	[1982TOR/SAB]
CH ₅ O ₃ P	[993-13-5]	methylphosphonic acid				
	$\Delta_{\text{sub}}H$		48.1 ± 4.2			[1955NEA/WIL, 1970COX/PIL]
CH ₆ ClN	[593-51-1]	methylamine hydrochloride				
	Δ_vH	(518–593)	114.5	533	A	[1987STE/MAL]
CH ₆ N ₂	[60-33-4]	methylhydrazine				
	$\Delta_{\text{fus}}H$		10.42	220.8		[1996DOM/HEA]
	Δ_vH	(274–299)	41.8	286	A	[1987STE/MAL, 1951AST/FIN]
CH ₆ N ₄ S	[2231-57-4]	thiocarbohydrazide				
	$\Delta_{\text{sub}}H$		152.1 ± 3.0	298	C	[1982TOR/SAB]
C ₂ BrCl ₂ F ₃ O ₄	[38217-36-6]	perchloric acid, 1,2,2-trifluoro-1-chloro-2-bromoethyl ester				
	Δ_vH	(273–294)	42.5	283	A	[1987STE/MAL, 1973SCH/PIL]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound					
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference	
C₂BrCl₃O	[34069-94-8] $\Delta_{\text{v}}H$	trichloroacetyl bromide (265–416)	42.6	280	A	[1987STE/MAL, 1947STU]	
C₂BrCl₅	[79504-02-2] $\Delta_{\text{sub}}H$	bromopentachloroethane (383–433)	44.4	398		[1987STE/MAL, 1949HIG/END]	
C₂BrF₃	[598-73-2] $\Delta_{\text{v}}H$	bromotrifluoroethylene (260–340)	25.0	275	A	[1987STE/MAL]	
C₂BrF₅O₃S	[757-02-8] $\Delta_{\text{v}}H$	2-bromotetrafluoroethyl fluorosulfate (273–298)	33.2	285		[1963GIL/CAD]	
C₂BrF₉S	[63011-81-4] $\Delta_{\text{v}}H$	pentafluoro(1-bromo-1,2,2,2-tetrafluoroethyl) sulfur (294–330)	30.7	309	A	[1987STE/MAL, 1999DYK/SVO]	
C₂Br₂ClF₃	[354-51-8] $\Delta_{\text{v}}H$	2-chloro-1,2-dibromo-1,1,2-trifluoroethane (343–428)	31.4	358	A	[1987STE/MAL]	
	$\Delta_{\text{v}}H$		35.0 ± 0.1	298	C	[1981MAJ/SVO]	
	$\Delta_{\text{v}}H$		34.2 ± 0.1	313	C	[1981MAJ/SVO]	
	$\Delta_{\text{v}}H$		33.5 ± 0.1	328	C	[1981MAJ/SVO]	
	$\Delta_{\text{v}}H$		32.6 ± 0.1	343	C	[1981MAJ/SVO]	
	$\Delta_{\text{v}}H$		31.6 ± 0.1	358	C	[1981MAJ/SVO]	
C₂Br₂Cl₄	[630-25-1] $\Delta_{\text{sub}}H$	1,2-dibromotetrachloroethane (383–453)	52.5	398		[1987STE/MAL, 1949HIG/END]	
	$\Delta_{\text{sub}}H$		(323–423)	56.7	373	A	[1935CAR/DIC]
C₂Br₂F₄	[124-73-2] $\Delta_{\text{fus}}H$	1,2-dibromotetrafluoroethane	7.04	162.8		[1991ACR]	
	$\Delta_{\text{v}}H$		(283–357)	28.5	298	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$		(354–443)	26.9	369	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$		(440–488)	27.1	455	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$			28.4 ± 0.1	298	C	[1981MAJ/SVO]
	$\Delta_{\text{v}}H$			27.5 ± 0.1	313	C	[1981MAJ/SVO]
	$\Delta_{\text{v}}H$			26.5 ± 0.1	328	C	[1981MAJ/SVO]
	$\Delta_{\text{v}}H$		(246–295)	30.0	280		[1987STE/MAL, 1970DYK]
C₂Br₄	[79-28-7] $\Delta_{\text{sub}}H$	tetrabromoethylene (221–310)	44.2	236		[1987STE/MAL]	
C₂ClFN₂	[30915-40-3] $\Delta_{\text{v}}H$	<i>cis</i> chloro(fluoroimino)acetonitrile (254–320)	31.7	269	A	[1987STE/MAL, 1971ZAB/SHR]	
C₂ClFN₂	[30915-39-0] $\Delta_{\text{v}}H$	<i>trans</i> chloro(fluoroimino)acetonitrile (257–320)	32.7	272	A	[1987STE/MAL, 1971ZAB/SHR]	
C₂ClF₂NO₂	[42016-33-1] $\Delta_{\text{v}}H$	chloro(fluorocarbonyl)carbamic fluoride (257–320)	36.8	376		[1973SPR/WRI]	
C₂ClF₃	[79-38-9] $\Delta_{\text{fus}}H$	chlorotrifluoroethylene	5.55	115		[1996DOM/HEA]	
	$\Delta_{\text{v}}H$		(206–262)	21.8	247	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$		(298–379)	20.2	313	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$		(206–263)	21.7	248		[1951MCC/PER]
	$\Delta_{\text{v}}H$		(195–250)	21.9	235		[1933HEI/MUR]
C₂ClF₃O₂	[23213-83-4] $\Delta_{\text{v}}H$	chloroformic acid, trifluoromethyl ester (195–273)	24.1	258	A	[1987STE/MAL]	

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
C ₂ ClF ₃ O ₄ S	[6069-32-5] $\Delta_{\text{v}}H$	difluorochloroacetic acid, fluorosulfuric acid anhydride (265–352)	39.8	280	A	[1987STE/MAL, 1966DES/CAD, 1999DYK/SVO]
C ₂ ClF ₄ NO	[42016-31-9] $\Delta_{\text{v}}H$	chloro(trifluoromethyl)carbamic fluoride	28.9	310		[1973SPR/WRI]
C ₂ ClF ₄ NO ₄ S	[42016-34-2] $\Delta_{\text{v}}H$	fluorosulfuric acid, chloro(trifluoromethyl)carbamic acid anhydride	28.5	398		[1973SPR/WRI]
C ₂ ClF ₅	[76-15-3] $\Delta_{\text{trs}}H$	chloropentafluoroethane	2.63	80.2		
	$\Delta_{\text{fus}}H$		1.88	173.7		[1996DOM/HEA]
	$\Delta_{\text{v}}H$	(262–317)	19.7	277	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$	(234–265)	20.1	250	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$	(312–353)	19.7	327	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$	(178–234)	20.9	219		[1966MEA/ROS]
	$\Delta_{\text{v}}H$	(176–235)	20.9	220	A	[1987STE/MAL, 1955AST/WIL]
$\Delta_{\text{v}}H$		19.4 ± 0.1	234	C	[1955AST/WIL]	
C ₂ ClF ₅ O	[22675-67-8] $\Delta_{\text{v}}H$	hypochlorous acid, pentafluoroethyl ester (193–248)	25.0	233	A	[1987STE/MAL, 1973DEM/SHR]
C ₂ ClF ₅ OS	[39937-08-1] $\Delta_{\text{v}}H$	pentafluoroethanesulfinyl chloride (273–338)	32.7	288	A	[1987STE/MAL, 1964RAT/SHR, 1999DYK/SVO]
C ₂ ClF ₅ O ₃ S	[649-61-6] $\Delta_{\text{v}}H$	2-chlorotetrafluoroethyl fluorosulfate (248–330)	32.9	289		[1963GIL/CAD]
C ₂ ClF ₅ O ₆ S ₂	[1957-17-1] $\Delta_{\text{v}}H$	1,2,2-trifluoro-1-chloro-1,2-ethanediol bis(fluorosulfate) (308–406)	53.2	323	A	[1987STE/MAL, 1999DYK/SVO]
C ₂ ClF ₆ NOS	[74366-11-3] $\Delta_{\text{v}}H$	(pentafluoroethyl)imidodisulfurous chloride fluoride	35.6	326	I	[1980ABE/SHR]
C ₂ ClF ₆ P	[650-52-2] $\Delta_{\text{v}}H$	bis(trifluoromethyl) chlorophosphine (193–273)	27.8	258		[1964PET/BUR, 1984BOU/FRI]
C ₂ ClF ₆ PS ₂	[660-05-9] $\Delta_{\text{v}}H$	chloro bis(trifluoromethylthio)phosphine (293–373)	33.0	333		[1960EME/PUG]
C ₂ ClF ₉ NP	[13105-57-2] $\Delta_{\text{v}}H$	[bis(trifluoromethyl)amino]trifluorochlorophosphorous(V) (223–273)	26.4	248		[1966RIN/ONE]
C ₂ ClF ₉ S	[646-63-9] $\Delta_{\text{v}}H$	2-chlorotetrafluoroethylsulfur pentafluoride	28.3	320		[1961CAS/RAY, 1999DYK/SVO]
C ₂ Cl ₂ F ₂	[598-88-9] $\Delta_{\text{v}}H$	1,2-dichloro-1,2-difluoroethylene (191–294)	27.9	279	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$	(240–294)	27.2	279		[1933BOO/BUR]
C ₂ Cl ₂ F ₂ N ₂	[30913-21-4] $\Delta_{\text{v}}H$	dichloro(difluoroamino)acetonitrile (238–341)	26.8	253	A	[1987STE/MAL, 1971ZAB/SHR]
C ₂ Cl ₂ F ₂ O	[354-18-7] $\Delta_{\text{v}}H$	fluorodichloroacetyl fluoride (208–273)	21.8	258	A	[1987STE/MAL]
C ₂ Cl ₂ F ₃ NO	[32751-03-4] $\Delta_{\text{v}}H$	N,N'-dichloro-2,2,2-trifluoroacetamide	40.9			[1972DEM/SHR]
C ₂ Cl ₂ F ₃ NO	[354-71-2] $\Delta_{\text{v}}H$	1,2-dichlorotrifluoro-1-nitrosoethane (307–310)	29.0	308	I	[1960GRI/HAZ]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
C ₂ Cl ₂ F ₃ NOS	[24433-67-8] $\Delta_{\text{v}}H$	S,S-dichloro-N-(trifluoroacetyl) sulfilimine (306–333)	44.2	319	A	[1987STE/MAL, 1969GLE/VON, 1999DYK/SVO]
C ₂ Cl ₂ F ₃ NO ₂ S	[51587-33-8] $\Delta_{\text{v}}H$	(trifluoromethyl)sulfonyl carbonimidic dichloride (312–405)	44.1	327	A	[1987STE/MAL, 1999DYK/SVO]
C ₂ Cl ₂ F ₄	[374-07-2] $\Delta_{\text{v}}H$	1,1-dichloro-1,2,2,2-tetrafluoroethane (231–373)	23.5	246	A	[1987STE/MAL, 1970DYK]
	$\Delta_{\text{v}}H$		23.2	233	BG	[1955MEA/STA]
	$\Delta_{\text{v}}H$		22.5	273	BG	[1955MEA/STA]
	$\Delta_{\text{v}}H$		20.8	313	BG	[1955MEA/STA]
	$\Delta_{\text{v}}H$		17.7	353	BG	[1955MEA/STA]
C ₂ Cl ₂ F ₄	[76-14-2] $\Delta_{\text{trs}}H$	1,2-dichloro-1,1,2,2-tetrafluoroethane (277–391)	1.21	109.3		
	$\Delta_{\text{trs}}H$		2.63	134.6		
	$\Delta_{\text{fus}}H$		1.51	180.6		[1996DOM/HEA]
	$\Delta_{\text{v}}H$		24.3	292	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$		25.1	262	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$		25.3	261		[1947STU]
C ₂ Cl ₂ F ₄ O ₄	[38126-28-2] $\Delta_{\text{v}}H$	perchloric acid, 1,1,2,2-tetrafluoro-2-chloroethyl ester (249–294)	32.6	279	A	[1987STE/MAL, 1973SCH/PIL]
C ₂ Cl ₂ F ₅ NS	[10564-48-4] $\Delta_{\text{v}}H$	S,S-dichloro-N-(pentafluoroethyl) sulfilimine (297–375)	37.4	312	A	[1987STE/MAL, 1999DYK/SVO]
C ₂ Cl ₂ F ₈ NP	[13105-58-3] $\Delta_{\text{v}}H$	[bis(trifluoromethyl)amino]difluorodichlorophosphorus(V) (262–305)	32.9	293		[1966EME/ONA]
C ₂ Cl ₃ F ₃	[354-58-5] $\Delta_{\text{v}}H$	1,1,1-trichloro-2,2,2-trifluoroethane (297–319)	28.1 ± 0.1	298	C	[1980MAJ/SVO]
	$\Delta_{\text{v}}H$		27.2 ± 0.1	313	C	[1980MAJ/SVO]
	$\Delta_{\text{v}}H$		26.3 ± 0.1	328	C	[1980MAJ/SVO]
	$\Delta_{\text{v}}H$		28.9	308		[1980MAJ/SVO]
	$\Delta_{\text{v}}H$		29.2	298	A	[1987STE/MAL, 1963HIR/HIL]
C ₂ Cl ₃ F ₃	[76-13-1] $\Delta_{\text{trs}}H$	1,1,2-trichloro-1,2,2-trifluoroethane (205–233)	0.83	82.5		
	$\Delta_{\text{fus}}H$		2.47	236.9		[1991ACR]
	$\Delta_{\text{sub}}H$		32.9	219	A	[1947STU]
	$\Delta_{\text{v}}H$		28.3	288	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$		30.9	253	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$		26.9	375	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$		28.8	307	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$		28.4 ± 0.1	298	C	[1980MAJ/SVO]
	$\Delta_{\text{v}}H$		27.5 ± 0.1	313	C	[1980MAJ/SVO]
	$\Delta_{\text{v}}H$		26.6 ± 0.1	328	C	[1980MAJ/SVO]
	$\Delta_{\text{v}}H$		28.2 ± 0.4	298		[1974VAR/BUL]
	$\Delta_{\text{v}}H$		28.2	288		[1963HIR/HIL]
	$\Delta_{\text{v}}H$		30.8	263		[1940BEN/MCH]
	$\Delta_{\text{v}}H$		NA			[1939REI]
	C ₂ Cl ₃ F ₃		[na] $\Delta_{\text{v}}H$	trichlorotrifluoroethane (248–352)	30.5	263
C ₂ Cl ₃ F ₃ O ₄	[38126-27-1]	perchloric acid, 1,2,2-trifluoro-1,2-dichloroethyl ester				

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
	$\Delta_{\text{v}}H$	(273–296)	26.9	284	A	[1987STE/MAL]
C ₂ Cl ₃ N	[545-06-2]	trichloroacetoneitrile				
	$\Delta_{\text{v}}H$	(289–357)	35.1	304	A	[1987STE/MAL, 1970DYK]
	$\Delta_{\text{v}}H$	(289–356)	34.7	304		[1954DAV/JEN]
C ₂ Cl ₄	[127-18-4]	tetrachloroethylene				
	$\Delta_{\text{trs}}H$		0.82	210		
	$\Delta_{\text{fus}}H$		10.88	250.8		[1996DOM/HEA]
	$\Delta_{\text{v}}H$	(307–393)	38.4	322		[1995DEJ/BUR]
	$\Delta_{\text{v}}H$	(310–393)	38.7	325	A	[1987STE/MAL, 1972BOU/AIM]
	$\Delta_{\text{v}}H$	(300–380)	38.9	315		[1970POL/MUR, 1984BOU/FRI]
	$\Delta_{\text{v}}H$	(333–373)	37.6	348		[1967FRI/GAL]
	$\Delta_{\text{v}}H$		39.7 ± 0.1	298	C	[1980MAJ/SVA]
C ₂ Cl ₄ F ₂	[76-12-0]	1,2-difluoro-1,1,2,2-tetrachloroethane				
	$\Delta_{\text{trs}}H$		0.79	130		
	$\Delta_{\text{fus}}H$		3.7	299.7		[1991ACR, 1978KIS/SUG]
	$\Delta_{\text{sub}}H$	(235–293)	36.4	278		[1987STE/MAL, 1947STU]
	$\Delta_{\text{sub}}H$	(237–293)	38.2	265	A	[1947STU]
	$\Delta_{\text{v}}H$	(313–361)	34.8 ± 0.4	298		[2007VAR/DRU, 1976VAR/BUL]
	$\Delta_{\text{v}}H$		34.6 ± 0.1	308	C	[1992SVO/KUB2]
	$\Delta_{\text{v}}H$		34.1 ± 0.1	315	C	[1992SVO/KUB2]
	$\Delta_{\text{v}}H$		33.6 ± 0.1	323	C	[1992SVO/KUB2]
	$\Delta_{\text{v}}H$		33.1 ± 0.1	330	C	[1992SVO/KUB2]
	$\Delta_{\text{v}}H$		32.6 ± 0.1	338	C	[1992SVO/KUB2]
	$\Delta_{\text{v}}H$	(301–365)	36.6	316	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$	(235–293)	36.4	278	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$	(312–362)	34.0	327	A	[1987STE/MAL]
$\Delta_{\text{v}}H$	(283–364)	32.7	298		[1933HOR/GEI]	
C ₂ Cl ₄ F ₂ O ₃ S	[na]	2-fluorotetrachloroethyl fluorosulfate				
	$\Delta_{\text{v}}H$	(311–437)	42.0	329		[1963GIL/CAD]
C ₂ Cl ₄ F ₂ O ₄	[38126-29-3]	perchloric acid, 1,2-difluoro-1,2,2-trichloroethyl ester				
	$\Delta_{\text{v}}H$	(273–294)	30.2	283	A	[1987STE/MAL, 1973SCH/PIL]
C ₂ Cl ₄ F ₄ N ₂	[35695-53-5]	1,2-bis(dichloroamino) tetrafluoroethane				
	$\Delta_{\text{v}}H$		43.1			[1972DEM/SHR]
C ₂ Cl ₄ F ₆ OS	[762-90-3]	pentafluoro(2-fluoro-1,1,2,2-tetrachloroethoxy) sulfur				
	$\Delta_{\text{v}}H$	(314–418)	42.8	329	A	[1987STE/MAL, 1962WIL/CAD]
C ₂ Cl ₄ O	[16650-10-5]	tetrachloroethylene oxide				
	$\Delta_{\text{v}}H$	(308–348)	36.9	323	A	[1987STE/MAL]
C ₂ Cl ₄ O	[76-02-8]	trichloroacetyl chloride				
	$\Delta_{\text{v}}H$	(305–393)	38.3	320	A	[1987STE/MAL, 1959MCD/SHR, 1970DYK]
C ₂ Cl ₆	[67-72-1]	hexachloroethane				
	$\Delta_{\text{trs}}H$		2.57	318		
	$\Delta_{\text{trs}}H$		8.22	345		
	$\Delta_{\text{fus}}H$		9.75	458		[1996DOM/HEA]
	$\Delta_{\text{sub}}H$	(317–345)	58.9	331		[1987STE/MAL]
	$\Delta_{\text{sub}}H$	(306–459)	48.8	382	A	[1947STU]
	$\Delta_{\text{sub}}H$ (mp 186.6)	(286–447)	59.1 ± 0.7	367		[1947IVI/DAI, 1960JON, 1970COX/PIL]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{sub}}H$ (<i>cubic</i>)	(286–447)	51.0	367		[1947IVI/DAI, 1960JON]
	$\Delta_{\text{sub}}H$		NA		GSM	[1941NIT/SEK]
	$\Delta_{\text{sub}}H$	(335–453)	50.5			[1935LEE]
	$\Delta_{\text{sub}}H$	(288–333)	59.0	310	GS,A	[1930NEL]
	Δ_vH	(460–513)	40.3	475	A	[1987STE/MAL]
	Δ_vH	(345–460)	51.2	360	A	[1987STE/MAL, 1970DYK]
	Δ_vH	(305–458)	53.7	320		[1947STU]
C ₂ FNO ₂	[15435-14-0]	fluorocarbonyl isocyanate				
	Δ_vH	(228–264)	33.5	249	A	[1987STE/MAL, 1967GLE/BIE]
C ₂ F ₂ N ₂ O	[32837-63-1]	difluorocarboncyamidic amide				
	Δ_vH		29.7	383		[1973WRI/SHR]
C ₂ F ₂ N ₂ O ₂	[32837-64-2]	difluorocarbonisocyanitic amide				
	Δ_vH		33.9	327		[1973WRI/SHR]
C ₂ F ₂ N ₄ O ₈	[20165-39-3]	1,2-difluoro-1,1,2,2-tetranitroethane				
	Δ_vH	(297–323)	62.8	310	A	[1987STE/MAL, 1973PEP/LEB]
C ₂ F ₂ O ₂	[359-40-0]	oxalyl fluoride				
	$\Delta_{\text{sub}}H$	(234–260)	16.7	247		[1987STE/MAL]
	Δ_vH	(264–272)	29.7	268		[1987STE/MAL]
C ₂ F ₂ O ₄	[692-74-0]	bis(fluorocarbonyl)peroxide				
	Δ_vH	(226–266)	30.6	251		[1962VON/AYM, 1984BOU/FRI]
C ₂ F ₃ N	[353-85-5]	trifluoroacetonitrile				
	$\Delta_{\text{fus}}H$		4.97	128.7		[1996DOM/HEA]
	Δ_vH	(151–206)	19.3	191	A	[1987STE/MAL]
	Δ_vH	(141–203)	19.2	188	A	[1987STE/MAL]
	Δ_vH	(197–241)	18.5	226	A	[1987STE/MAL]
	Δ_vH	(336–282)	17.4	309	A	[1987STE/MAL]
	Δ_vH	(272–311)	17.4	287	A	[1987STE/MAL]
	Δ_vH	(142–206)	19.2	191		[1961PAC/BOB]
C ₂ F ₃ NO	[460-49-1]	trifluoromethyl isocyanate				
	Δ_vH	(195–228)	22.5	213	A	[1987STE/MAL]
C ₂ F ₃ NO	[2713-04-4]	trifluoronitrosoethylene				
	Δ_vH	(247–250)	25.7	248	A,I	[1987STE/MAL, 1960GRI/HAZ]
C ₂ F ₃ NOS	[61951-27-7]	trifluoromethanesulfinyl cyanide				
	Δ_vH		40.2	352	I	[1977BUR/SHR]
C ₂ F ₃ NOS	[691-03-2]	trifluoroemethylsulphenyl isocyanate				
	Δ_vH	(231–293)	27.9	278	A	[1987STE/MAL, 1999DYK/SVO, 1963EME/HAA]
C ₂ F ₃ NO ₂ S	[26454-68-2]	2,2,2-trifluoro-N-sulfinylacetamide				
	Δ_vH	(267–302)	36.4	282	A	[1987STE/MAL, 1999DYK/SVO]
C ₂ F ₃ NO ₂ S ₂	[51587-30-5]	trifluoromethanesulfonyl isothiocyanate				
	Δ_vH	(297–385)	41.0	312	A	[1987STE/MAL, 1999DYK/SVO]
C ₂ F ₃ NO ₃ S	[30227-06-6]	trifluoromethanesulfonyl isocyanate				
	Δ_vH	(275–345)	36.9	290	A	[1987STE/MAL, 1999DYK/SVO]
C ₂ F ₃ NS	[690-24-4]	thiocyanic acid, trifluoromethyl ester				
	Δ_vH	(226–294)	32.6	279	A	[1987STE/MAL, 1999DYK/SVO, 1963EME/HAA]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
C ₂ F ₃ N ₃ O ₆	[20165-38-2] $\Delta_{\text{v}}H$	1,1,2-trifluoro-1,2,2-trinitroethane (313–353)	57.7	328	A	[1987STE/MAL, 1973PEP/LEB]
C ₂ F ₄	[116-14-3] $\Delta_{\text{fus}}H$	tetrafluoroethylene	7.71	142		[1996DOM/HEA]
	$\Delta_{\text{v}}H$	(197–273)	16.8	258	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$	(273–306)	16.6	288	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$	(142–208)	18.6	193	A	[1987STE/MAL, 1953FUR/MCC, 1984BOU/FRI]
C ₂ F ₄ N ₂	[5131-88-4] $\Delta_{\text{v}}H$	tetrafluoroaminoacetic, nitrile (193–238)	23.9	223	A	[1987STE/MAL]
C ₂ F ₄ N ₂ O ₃	[679-08-3] $\Delta_{\text{v}}H$	1,1,2,2-tetrafluoro-1-nitro-2-nitrosoethane (233–293)	28.8	278	A	[1987STE/MAL]
C ₂ F ₄ N ₂ O ₄	[356-16-1] $\Delta_{\text{v}}H$	1,1,2,2-tetrafluoro-1,2-dinitroethane (303–343)	67.8	323		[1973PEP/LEB]
	$\Delta_{\text{v}}H$	(259–333)	34.7	274	A, I	[1987STE/MAL, 1957FRA/SAN]
C ₂ F ₄ N ₂ O ₆ S ₂	[19252-50-7] $\Delta_{\text{v}}H$	1,2-bis(fluoroformyl)-1,2-bis(fluorosulfonyl)hydrazine (273–296)	49.8	284	A	[1987STE/MAL, 1999DYK/SVO]
C ₂ F ₄ O	[354-34-7] $\Delta_{\text{v}}H$	trifluoroacetyl fluoride (161–215)	20.9	200	A	[1987STE/MAL, 1972PAC/HOD]
C ₂ F ₄ O ₂ S	[684-106] $\Delta_{\text{v}}H$	trifluoroethylene sulfonyl fluoride (270–313)	27.0	285	A	[1987STE/MAL, 1999DYK/SVO]
C ₂ F ₄ O ₃	[16118-40-4] $\Delta_{\text{v}}H$	fluoroperoxyformic acid, trifluoromethyl ester (194–249)	27.3	234	A	[1987STE/MAL]
C ₂ F ₄ O ₄ S	[5762-53-8] $\Delta_{\text{v}}H$	trifluoroacetyl fluorosulfate (262–321)	34.3	277	A	[1987STE/MAL, 1966DEL/SHR]
	$\Delta_{\text{v}}H$	(250–320)	34.9	265		[1966MEA/ROS]
C ₂ F ₄ S ₂	[1717-50-6] $\Delta_{\text{v}}H$	tetrafluoro-1-3-dithietane	29.2			[1973ABE/SHR]
C ₂ F ₅ I	[354-64-3] $\Delta_{\text{v}}H$	pentafluoroiodoethane (248–283)	20.8	268	A	[1987STE/MAL]
C ₂ F ₅ NO	[32822-49-4] $\Delta_{\text{v}}H$	pentafluoroacetamide	23.8	252	HG	[1971DEM/SHR]
C ₂ F ₅ NO	[354-72-3] $\Delta_{\text{v}}H$	pentafluoronitrosoethane (193–227)	20.9	212	A	[1987STE/MAL, 1956BAR/HAS]
C ₂ F ₅ NOS	[32837-66-4] $\Delta_{\text{v}}H$	carbamothioic acid, difluoro-S-(trifluoromethyl) ester (193–227)	23.0	315		[1973WRI/SHR]
C ₂ F ₅ NOS	[24433-65-6] $\Delta_{\text{v}}H$	S,S-difluoro-N-(trifluoroacetyl) sulfilimine (240–282)	34.4	267	A	[1987STE/MAL, 1969GLE/VON, 1999DYK/SVO]
C ₂ F ₅ NOS	[28103-61-9] $\Delta_{\text{v}}H$	1,1,1-trifluoro-N-(fluoroformyl)methanesulfinimidyl fluoride (276–323)	38.9	291	A	[1987STE/MAL, 1999DYK/SVO]
C ₂ F ₅ NOS	[10564-50-8] $\Delta_{\text{v}}H$	1,1,2,2,2-pentafluoro-N-sulfinyl ethylamine (245–303)	29.0	260	A	[1987STE/MAL, 1999DYK/SVO]
C ₂ F ₅ NO ₄ S	[19252-49-4] $\Delta_{\text{v}}H$	(fluorosulfonyl)(trifluoromethoxy)carbamoyl fluoride (277–290)	30.3	283		[1999DYK/SVO]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₂ F ₅ N ₃ O ₃	[755-68-0] Δ_vH	fluoro(1,1,2,2-tetrafluoro-2-nitroethyl)-2-diimide oxide (257–350)	38.0	272	A	[1987STE/MAL]
C ₂ F ₆	[76-16-4] $\Delta_{\text{trs}}H$	hexafluoroethane	3.74	104		
	$\Delta_{\text{fus}}H$		2.69	173.1		[1996DOM/HEA]
	$\Delta_{\text{sub}}H$		26.0	103		[1963BON, 1948PAC/AST]
	Δ_vH	(172–200)	17.3	186	A	[1987STE/MAL]
	Δ_vH	(180–196)	17.1	188		[1948PAC/AST]
C ₂ F ₆ IN	[5764-87-4] Δ_vH	N-iodo- <i>bis</i> (trifluoromethyl)amine (261–318)	28.5	276	A	[1987STE/MAL]
	C ₂ F ₆ IP	[359-64-8] Δ_vH	<i>bis</i> (trifluoromethyl) phosphinous iodide (273–320)	33.2	288	
C ₂ F ₆ N ₂	[372-63-4] Δ_vH	hexafluoroazomethane (205–242)	22.9	227	A	[1987STE/MAL]
	C ₂ F ₆ N ₂ O	[371-56-2] Δ_vH	hexafluoroazoxymethane (274–281)	27.2	277	A
C ₂ F ₆ N ₂ O ₂	[359-75-1] Δ_vH	1,1,1-trifluoro-N-(nitrosooxy)-N-(trifluoromethyl)methanamine (245–285)	26.8	270	A	[1987STE/MAL]
	C ₂ F ₆ N ₂ O ₂	[367-54-4] Δ_vH	N-nitroso-O,N- <i>bis</i> (trifluoromethyl)hydroxylamine (272–283)	25.4	277	A
C ₂ F ₆ OS	[30341-37-8] Δ_vH	<i>bis</i> (trifluoromethyl)sulfoxide (248–303)	27.9	263	A	[1987STE/MAL, 1999DYK/SVO]
	C ₂ F ₆ OS	[20621-31-2] Δ_vH	pentafluoroethyl sulfinyl fluoride (234–293)	28.5	278	A, I
C ₂ F ₆ OS ₂	[63548-94-7] Δ_vH	S-trifluoromethyl-(trifluoromethyl)thiosulfinate (293–353)	30.7	333		[1999DYK/SVO]
	C ₂ F ₆ OS ₂	[na] Δ_vH	methanesulfonothioic acid, trifluoro-S-(trifluoromethyl) ester	27.8	329	I
C ₂ F ₆ O ₂ S	[354-87-0] Δ_vH	perfluoroethyl fluorosulfate (250–300)	28.8	275		[1963GIL/CAD]
	C ₂ F ₆ O ₃	[1718-18-9] Δ_vH	<i>bis</i> (trifluoromethyl) trioxide (193–248)	24.3	233	A
C ₂ F ₆ O ₃ S		[3582-05-6] Δ_vH	trifluoromethanesulfonic acid, trifluoromethyl ester (238–294)	29.4	252	
	Δ_vH	(238–294)	27.6	279	A	[1987STE/MAL, 1965NOF/CAD]
C ₂ F ₆ O ₄ S	[1479-52-3] Δ_vH	<i>bis</i> (trifluoromethyl) sulfate (219–304)	28.7	262		[1960VAN/CAD]
	C ₂ F ₆ O ₅ S	[41765-14-4] Δ_vH	peroxysulfuric acid, <i>bis</i> (trifluoromethyl) ester (253–319)	32.0	268	
C ₂ F ₆ O ₆ S ₂		[1479-53-4] Δ_vH	tetrafluoroethylene glycol, <i>bis</i> (fluorosulfate) (295–378)	43.7	310	A
	C ₂ F ₆ O ₇ S ₂	[na] Δ_vH	<i>bis</i> (trifluoromethyl) disulfate (328–357)	38.3	342	

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₂ F ₆ S	[371-78-8]	<i>bis</i> (trifluoromethyl) sulfide				
	$\Delta_v H$		23.6			[1952BRA/EME]
C ₂ F ₆ S ₂	[372-64-5]	<i>bis</i> (trifluoromethyl) disulfide				
	$\Delta_v H$		28.8			[1952BRA/EME]
C ₂ F ₇ N	[359-62-6]	perfluorodimethylamine				
	$\Delta_v H$	(199–230)	21.4	215	A	[1987STE/MAL]
	$\Delta_v H$	(203–233)	18.6	218	A	[1987STE/MAL, 1949THO/EME]
C ₂ F ₇ N	[354-80-3]	perfluoroethylamine				
	$\Delta_v H$	(171–236)	20.8	221	A	[1987STE/MAL, 1970DYK]
C ₂ F ₇ NOS	[59617-28-6]	(pentafluoroethyl)imidodisulfuryl fluoride				
	$\Delta_v H$		30.7			[1976STA/MEW]
C ₂ F ₇ NO ₃ S	[4188-34-5]	fluorosulfuric acid, 1,1,2,2-tetrafluoro-2-(difluoroamino)ethyl ester				
	$\Delta_v H$	(276–326)	31.1	291	A	[1987STE/MAL, 1999DYK/SVO]
C ₂ F ₇ NO ₁₂ S ₄	[53684-02-9]	fluorosulfuric acid, 1-[<i>bis</i> [(fluorosulfonyl)oxo]amino]-2,2,2-trifluoroethylidene ester				
	$\Delta_v H$		43.4	418		[1975KIR/LAS]
C ₂ F ₈ NOP	[13105-59-4]	[<i>bis</i> (difluoromethyl)amino] difluorophosphine oxide				
	$\Delta_v H$	(233–278)	30.4	255		[1966EME/ONA]
C ₂ F ₈ NOP	[36544-19-1]	phosphorous <i>bis</i> (trifluoromethyl)nitroxide difluoride				
	$\Delta_v H$		28.0	288		[1973WAN/SHR]
C ₂ F ₈ OS	[33716-15-3]	difluoro-oxo- <i>bis</i> -(trifluoromethyl)sulfur				
	$\Delta_v H$	(239–299)	22.4	254	A	[1987STE/MAL, 1999DYK/SVO]
	$\Delta_v H$		28.4			[1971SAU/SHR]
C ₂ F ₈ OS	[82390-51-0]	pentafluoro(trifluoroacetyl) sulfur				
	$\Delta_v H$	(162–290)	26.6	177		[1999DYK/SVO]
C ₂ F ₈ O ₃ S	[60672-61-9]	pentafluoro (trifluoroethaneperoxoato) sulfur				
	$\Delta_v H$		28.0			[1976HOP/DES]
C ₂ F ₈ S	[1186-51-2]	trifluorovinyl sulfur pentafluoride				
	$\Delta_v H$		25.1	292		[1961CAS/RAY]
C ₂ F ₈ S	[30341-38-9]	difluoro <i>bis</i> (trifluoromethyl) sulfur				
	$\Delta_v H$		28.8			[1971SAU/SHR]
C ₂ F ₁₀ OS	[1580-07-0]	pentafluoro(pentafluoroethoxy) sulfur				
	$\Delta_v H$	(245–287)	27.6	272	A	[1987STE/MAL, 1962WIL/CAD]
C ₂ F ₁₀ O ₂ S	[2004-38-8]	tetrafluoro- <i>bis</i> (trifluoromethoxy) sulfur				
	$\Delta_v H$	(246–302)	29.9	261	A	[1987STE/MAL, 1964DUN/CAD]
C ₂ F ₁₀ O ₃ S	[41938-43-6]	(trifluoromethoxy)[(trifluoromethyl)dioxy] sulfur tetrafluoride				
	$\Delta_v H$	(255–317)	32.5	270		[1999DYK/SVO]
C ₂ F ₁₀ O ₃ S ₂	[68010-32-2]	pentafluoro[1,2,2,2-tetrafluoro-1-[(fluorosulfonyl)oxy]ethyl] sulfur				
	$\Delta_v H$		34.8			[1978DEM/FOX]
C ₂ F ₁₀ S	[42179-02-2]	<i>trans</i> tetrafluoro- <i>bis</i> (trifluoromethyl) sulfur				
	$\Delta_v H$	(233–293)	23.3	278	A	[1987STE/MAL, 1999DYK/SVO]
C ₂ F ₁₁ NS	[13888-13-6]	[<i>bis</i> (trifluoromethyl)amino] sulfur pentafluoride				
	$\Delta_v H$	(233–306)	29.3	248	A	[1987STE/MAL, 1966DOB, 1999DYK/SVO]
C ₂ F ₁₂ S ₂	[42060-66-2]	perfluoro-1,3-dithietane octafluoride				

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
	$\Delta_{\text{v}}H$		35.6			[1973ABE/SHR]
C ₂ N ₂	[460-19-5]	cyanogen				
	$\Delta_{\text{fus}}H$		8.11	245.3		[1996DOM/HEA]
	$\Delta_{\text{sub}}H$	(202–239)	33.0	224		[1987STE/MAL]
	$\Delta_{\text{sub}}H$	(177–230)	33.6	204	A	[1947STU]
	$\Delta_{\text{sub}}H$	(202–245)	34.4		CATH	[1939RUE/GIA]
	$\Delta_{\text{sub}}H$	(198–240)	32.4	224		[1975GRO, 1925PER/BAR]
	$\Delta_{\text{sub}}H$		NA			[1916TER]
	$\Delta_{\text{v}}H$	(240–253)	24.5	246	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$	(246–273)	23.9	257		[1925PER/BAR]
$\Delta_{\text{v}}H$	(246–273)	23.5	267		[1925PER/BAR]	
C ₂ N ₆ O ₁₂	[918-37-6]	hexanitroethane				
	$\Delta_{\text{sub}}H$		70.7	298		[1999MIR/VOR]
	$\Delta_{\text{sub}}H$	(293–343)	30.4	308		[1987STE/MAL, 1963NOB/REE]
	$\Delta_{\text{sub}}H$	(293–313)	70.7 ± 1.7	303	ME	[1969MIR/LEB, 1977PED/RYL]
	$\Delta_{\text{sub}}H$	(293–343)	70.7 ± 1.7			[1968PEP/MIR]
C ₂ HBr	[593-61-3]	bromoacetylene				
	$\Delta_{\text{v}}H$	(214–273)	25.6	273	A	[1987STE/MAL]
C ₂ HBrClF ₃	[151-67-7]	2-bromo-2-chloro-1,1,1-trifluoroethane				
	$\Delta_{\text{fus}}H$		4.84	154.7		[1996DOM/HEA]
	$\Delta_{\text{v}}H$	(302–323)	29.8 ± 0.4	298		[2007VAR/DRU, 1985PAS/VAR]
	$\Delta_{\text{v}}H$	(298–323)	30.0	310	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$		29.6 ± 0.3	298		[1981PAP/ERA]
	$\Delta_{\text{v}}H$		29.6 ± 0.1	298	C	[1980MAJ/SVO]
	$\Delta_{\text{v}}H$		28.7 ± 0.1	313	C	[1980MAJ/SVO]
	$\Delta_{\text{v}}H$		27.8 ± 0.1	328	C	[1980MAJ/SVO]
	$\Delta_{\text{v}}H$		26.8 ± 0.1	343	C	[1980MAJ/SVO]
	$\Delta_{\text{v}}H$	(227–318)	34.3	242		[1965KUD/SAV]
$\Delta_{\text{v}}H$	(222–329)	33.2	237	A	[1987STE/MAL, 1963BOT/SEI, 1970DYK]	
C ₂ HBrClF ₃	[354-06-3]	1-bromo-2-chloro-1,1,2-trifluoroethane				
	$\Delta_{\text{fus}}H$		4.38	146.2		[1996DOM/HEA]
	$\Delta_{\text{v}}H$		30.0 ± 0.1	298	C	[1981MAJ/SVO]
	$\Delta_{\text{v}}H$		29.0 ± 0.1	313	C	[1981MAJ/SVO]
	$\Delta_{\text{v}}H$		28.1 ± 0.1	328	C	[1981MAJ/SVO]
$\Delta_{\text{v}}H$		27.2 ± 0.1	343	C	[1981MAJ/SVO]	
C ₂ HBrF ₈ S	[82390-50-9]	(1-bromo-2,2,2-trifluoroethyl)sulfur pentafluoride				
	$\Delta_{\text{v}}H$		32.0			[1982DEM/FOX]
C ₂ HBr ₂ FO ₂	[353-99-1]	dibromofluoroacetic acid				
	$\Delta_{\text{v}}H$	(403–468)	60.2	468	A	[1987STE/MAL]
C ₂ HBr ₃ O	[115-17-3]	tribromoacetaldehyde				
	$\Delta_{\text{v}}H$	(291–447)	47.8	306	A	[1987STE/MAL, 1947STU]
C ₂ HCl	[593-63-5]	chloroacetylene				
	$\Delta_{\text{v}}H$	(204–238)	21.8	223	A	[1987STE/MAL]
C ₂ HCIF ₂	[359-10-4]	1,1-difluoro-2-chloroethene				
	$\Delta_{\text{v}}H$		23.5	233	BG	[1955MEA/STA]
	$\Delta_{\text{v}}H$		21.4	273	BG	[1955MEA/STA]
$\Delta_{\text{v}}H$		18.5	313	BG	[1955MEA/STA]	

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$		14.1	353	BG	[1955MEA/STA]
C₂HCIF₆OS	[20407-78-7]	<i>trans</i> [(2-chloro-2-fluorovinyl)oxy] sulfur pentafluoride				
	$\Delta_v H$		36.8			[1968PLA/WIL]
C₂HCIF₆OS	[20407-79-8]	<i>cis</i> [(2-chloro-2-fluorovinyl)oxy] sulfur pentafluoride				
	$\Delta_v H$		34.3			[1968PLA/WIL]
C₂HCIF₈OS	[20334-47-8]	(2-chloro-1,2,2-trifluoroethoxy) sulfur pentafluoride				
	$\Delta_v H$		33.3			[1968PLA/WIL]
C₂HCIF₁₂O₂S	[20563-90-0]	[(2-chloro-2,2-difluoroethylidene)dioxy]bis(pentafluoro)sulfur				
	$\Delta_v H$		39.0			[1968PLA/WIL]
C₂HCIF₈S	[22756-13-4]	(1,1,2-trifluoro-2-chloroethyl) sulfur pentafluoride				
	$\Delta_v H$	(279–323)	30.2	294	A	[1987STE/MAL, 1999DYK/SVO]
C₂HCl₂F₃	[306-83-2]	1,1,1-trifluoro-2,2-dichloroethane				
	$\Delta_{\text{fus}} H$		5.51	145.7		[2002VAR/DRU]
	$\Delta_v H$		26.6 ± 0.3	298		[2002VAR/DRU]
	$\Delta_v H$	(243–448)	28.7	258	MM	[1992OGU/YAM]
C₂HCl₂F₃	[354-23-4]	1,1,2-trifluoro-1,2-dichloroethane				
	$\Delta_{\text{fus}} H$		7.08	135.7		[1999DRU/VAR]
	$\Delta_v H$		26.8 ± 0.3	298		[2002VAR/DRU]
C₂HCl₃	[79-01-6]	trichloroethylene				
	$\Delta_{\text{fus}} H$		8.45	188.5		[1996DOM/HEA]
	$\Delta_v H$	(297–360)	34.2	313		[1995AUC/GON]
	$\Delta_v H$		34.5 ± 0.1	298	C	[1980MAJ/SVA]
	$\Delta_v H$	(280–428)	34.6	295		[1987STE/MAL, 1970DYK]
	$\Delta_v H$	(290–359)	36.2	305		[1944MCD]
	$\Delta_v H$	(298–360)	35.6	313		[1912HER/RAT, 1984BOU/FRI]
C₂HCl₃F₂	[354-21-2]	1,1-difluoro-1,2,2-trichloroethane				
	$\Delta_v H$	(297–345)	32.8 ± 0.4	298		[2007VAR/DRU, 1976VAR/BUL]
	$\Delta_v H$		32.9 ± 0.1	298	C	[1997VAR/PAS]
C₂HCl₃F₂	[354-15-4]	1,2-difluoro-1,1,2-trichloroethane				
	$\Delta_v H$	(289–346)	33.0 ± 0.4	298		[2007VAR/DRU, 1997VAR/PAS]
	$\Delta_v H$		33.1 ± 0.1	298	C	[1997VAR/PAS]
C₂HCl₃F₂O₃S	[42087-88-7]	fluorosulfuric acid, 2-fluoro-1,1,2-trichloroethyl ester				
	$\Delta_v H$	(317–353)	36.6	332	A	[1987STE/MAL, 1999DYK/SVO]
C₂HCl₃O	[75-87-6]	trichloroacetaldehyde				
	$\Delta_v H$	(235–371)	36.6	250	A	[1987STE/MAL, 1947STU]
C₂HCl₃O₂	[76-03-9]	trichloroacetic acid				
	$\Delta_{\text{fus}} H$		5.88	330.7		[1991ACR]
	$\Delta_v H$	(326–473)	65.0	341	A	[1987STE/MAL, 1970DYK]
	$\Delta_v H$	(385–470)	57.2	400		[1959MCD/SHR]
C₂HCl₄FS	[2822-06-2]	(dichloromethyl)(fluorodichloromethyl) sulfide				
	$\Delta_v H$	(322–352)	46.5	337	A	[1987STE/MAL, 1999DYK/SVO]
C₂HCl₅	[76-01-7]	pentachloroethane				
	$\Delta_v H$	(274–434)	40.9	289	A	[1987STE/MAL]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(298–435)	45.5	313		[1930NEL]
C ₂ HF ₃ O ₂	[76-05-1]	trifluoroacetic acid				
	$\Delta_v H$	(285–345)	35.9	300	A	[1987STE/MAL, 1962KRE, 1970DYK, 1984BOU/FRI]
C ₂ HF ₅	[354-33-6]	pentafluoroethane				
	$\Delta_{\text{fus}} H$		2.25	172.6		[1996LUE/MAG]
	$\Delta_v H$		22.8	175	C	[1999WEB]
	$\Delta_v H$		21.9	190	C	[1999WEB]
	$\Delta_v H$		20.9	205	C	[1999WEB]
	$\Delta_v H$		20.3	215	C	[1999WEB]
	C ₂ HF ₅ O	[3822-68-2]	pentafluorodimethyl ether			
$\Delta_v H$		(216–238)	22.3	239	I	[2001KUL/DES]
$\Delta_v H$		(229–331)	19.3	260	EB	[1996WEB/DEF2]
$\Delta_v H$		(229–331)	17.6	280	EB	[1996WEB/DEF2]
$\Delta_v H$		(229–331)	15.6	300	EB	[1996WEB/DEF2]
$\Delta_v H$		(240–313)	20.4	255	A	[1992SAL/WAN]
C ₂ HF ₆ NOS	[34556-22-4]	S,S-bis(trifluoromethyl)sulfoximine				
	$\Delta_v H$		35.1	346	I	[1972SAU/SHR]
C ₂ HF ₆ NS ₂	[na]	bis(trifluoromethane) sulphenimide				
	$\Delta_v H$	(243–293)	36.5	268		[1960EME/NAB]
C ₂ HF ₆ OP	[359-65-9]	phosphinous acid, bis(trifluoromethyl) ester				
	$\Delta_{\text{sub}} H$	(233–251)	46.6	242		[1987STE/MAL, 1962GRI/BUR]
C ₂ HF ₆ OPS	[35814-49-4]	bis(trifluoromethyl) thiophosphinic acid				
	$\Delta_v H$	(283–324)	38.3	298	A	[1987STE/MAL]
C ₂ HF ₆ PS	[1486-19-7]	bis(trifluoromethyl)(mercapto)phosphine				
	$\Delta_v H$	(269–304)	30.7	286		[1964CAV/EME]
C ₂ HF ₆ PS ₂	[1486-19-7]	bis(trifluoromethyl) thiophosphinous acid				
	$\Delta_v H$	(217–280)	32.9	265	A	[1987STE/MAL, 1999DYK/SVO]
C ₂ HF ₆ PS ₂	[18799-75-2]	phosphinodithioic acid, bis(trifluoromethyl) ester				
	$\Delta_{\text{sub}} H$	(273–287)	41.9	280	A	[1987STE/MAL]
C ₂ HF ₇ S	[58636-78-5]	(2,2-difluoroethenyl) pentafluorosulfur				
	$\Delta_v H$		27.7			[1978DEM/FOX]
C ₂ HF ₉ S	[63011-80-3]	pentafluoro (1,2,2,2-tetrafluoroethyl) sulfur				
	$\Delta_v H$		28.0			[1978DEM/FOX]
C ₂ H ₂	[74-86-2]	acetylene				
	$\Delta_{\text{trs}} H$		2.54	142.7		
	$\Delta_{\text{fus}} H$		3.76	192.4		[1976MIS/RIE]
	$\Delta_{\text{sub}} H$	(98–145)	23.5	130	A	[1987STE/MAL]
	$\Delta_{\text{sub}} H$	(133–191)	21.8	162		[1960JON]
	$\Delta_{\text{sub}} H$	(151–193)	25.2	193		[1956AMB]
	$\Delta_{\text{sub}} H$	(130–189)	22.7	160	A	[1947STU]
	$\Delta_{\text{sub}} H$	(89–169)	22.1	129	A	[1943BUR]
	$\Delta_v H$	(258–308)	16.3	273	A	[1987STE/MAL]
	$\Delta_v H$	(192–308)	16.7	207	A	[1987STE/MAL]
	$\Delta_v H$	(192–225)	16.7	210	A	[1987STE/MAL]
$\Delta_v H$		17.0	214		[1971WIL/ZWO]	

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(215–308)	16.4	230		[1964AMB/TOW]
	$\Delta_v H$	(193–207)	16.8	200		[1956AMB, 1984BOU/FRI]
C ₂ H ₂ Br ₂	[590-11-4]	<i>cis</i> 1,2-dibromoethylene				
	$\Delta_v H$	(299–351)	40.6	314	A	[1987STE/MAL, 1950NOY/NOY, 1970DYK]
C ₂ H ₂ Br ₂	[590-12-5]	<i>trans</i> 1,2-dibromoethylene				
	$\Delta_v H$	(277–344)	35.2	310	A	[1987STE/MAL, 1970DYK]
	$\Delta_v H$	(277–343)	42.9	292		[1950NOY/NOY, 1984BOU/FRI]
C ₂ H ₂ Br ₂ Cl ₂	[75-81-0]	1,2-dibromo-1,1-dichloroethane				
	$\Delta_v H$	(354–519)	45.9	369	A	[1987STE/MAL, 1970DYK]
C ₂ H ₂ Br ₂ Cl ₂	[683-68-1]	1,2-dibromo-1,2-dichloroethane				
	$\Delta_v H$	(320–379)	45.9	335	A	[1987STE/MAL]
C ₂ H ₂ Br ₂ F ₂	[75-82-1]	1,2-dibromo-1,1-difluoroethane				
	$\Delta_{\text{fus}} H$		8.3	206.3		[1991ACR]
C ₂ H ₂ Br ₄	[630-16-0]	1,1,1,2-tetrabromoethane				
	$\Delta_v H$	(331–473)	61.5	346	A	[1987STE/MAL, 1947STU]
C ₂ H ₂ Br ₄	[79-27-6]	1,1,2,2-tetrabromoethane				
	$\Delta_v H$	(413–573)	56.9	428	A	[1987STE/MAL, 1970DYK]
C ₂ H ₂ ClFO	[359-14-8]	chloroacetyl fluoride				
	$\Delta_v H$	(273–333)	38.0	288	A, GS	[1987STE/MAL, 1948RED/CHA4, 1970DYK]
C ₂ H ₂ ClFO	[359-06-8]	fluoroacetyl chloride				
	$\Delta_v H$	(273–333)	36.7	288	A, GS	[1987STE/MAL, 1948RED/CHA4, 1970DYK]
C ₂ H ₂ ClF ₃ O ₂ S	[57169-80-9]	chlorosulfurous acid, 2,2,2-trifluoroethyl ester				
	$\Delta_v H$		36.0			[1975DEM/KOV2]
C ₂ H ₂ ClF ₇ S	[68010-35-5]	(2-chloro-2,2-difluoroethyl)pentafluorosulfur				
	$\Delta_v H$		32.9			[1978DEM/FOX]
C ₂ H ₂ Cl ₂	[75-35-4]	1,1-dichloroethylene				
	$\Delta_{\text{fus}} H$		6.51	150.9		[1996DOM/HEA]
	$\Delta_v H$	(245–305)	28.4	260	A	[1987STE/MAL, 1959HIL/MCD, 1970DYK]
C ₂ H ₂ Cl ₂	[156-59-2]	<i>cis</i> 1,2-dichloroethylene				
	$\Delta_v H$	(332–495)	29.3	347	A	[1987STE/MAL]
	$\Delta_v H$	(273–334)	31.5	288	A	[1987STE/MAL, 1970DYK]
	$\Delta_v H$	(292–335)	31.6	307		[1951FLO/ALP]
	$\Delta_v H$	(273–356)	31.8	288		[1947KET/VAN]
C ₂ H ₂ Cl ₂	[156-60-5]	<i>trans</i> 1,2-dichloroethylene				
	$\Delta_v H$	(321–473)	29.0	336	A	[1987STE/MAL]
	$\Delta_v H$	(273–319)	30.1	288		[1983MAC]
	$\Delta_v H$	(263–323)	30.4	278	A	[1987STE/MAL, 1970DYK]
	$\Delta_v H$	(235–358)	31.4	250		[1947KET/VAN]
C ₂ H ₂ Cl ₂ F ₂	[1649-08-7]	1,2-dichloro-1,1-difluoroethane				
	$\Delta_v H$	(323–493)	27.8	338	A	[1987STE/MAL]
C ₂ H ₂ Cl ₂ F ₂	[1842-05-3]	1,2-difluoro-2,2-dichloroethane				
	$\Delta_{\text{fus}} H$		8.19			[1996DOM/HEA]
C ₂ H ₂ Cl ₂ F ₆ OS	[20334-44-5]	(1,2-dichloro-2-fluoroethoxy)pentafluoro sulfur				
	$\Delta_v H$		38.8			[1968PLA/WIL]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C₂H₂Cl₂F₆OS	[20334-45-6]	(2,2-dichloro-2-fluoroethoxy)pentafluoro sulfur				
	$\Delta_v H$		38.5			[1968PLA/WIL]
C₂H₂Cl₂O	[79-04-9]	chloroacetyl chloride				
	$\Delta_v H$	(253–379)	45.0	268	A	[1987STE/MAL, 1970DYK]
	$\Delta_v H$	(301–380)	40.8	316		[1959MCD/SHR]
	$\Delta_v H$	(290–373)	44.1	305		[1935KIR/POP]
C₂H₂Cl₂O₂	[79-43-6]	dichloroacetic acid				
	$\Delta_{\text{fus}} H$		12.34	286.5		[1996DOM/HEA]
	$\Delta_v H$	(317–468)	55.7	332	A	[1987STE/MAL, 1947STU]
C₂H₂Cl₄	[630-20-6]	1,1,1,2-tetrachloroethane				
	$\Delta_v H$		45.7 ± 0.1	298	C	[1980MAJ/SVA]
	$\Delta_v H$	(316–447)	40.1	331	A	[1987STE/MAL, 1970DYK]
	$\Delta_v H$	(332–403)	39.2	347		[1949DRE/SHR, 1949DRE/MAR]
C₂H₂Cl₄	[79-34-5]	1,1,2,2-tetrachloroethane				
	$\Delta_{\text{trs}} H$		0.54	207.3		
	$\Delta_{\text{fus}} H$		9.17	230.8		[1996DOM/HEA]
	$\Delta_v H$	(343–418)	42.5	358	EB	[2006TEO/BAR]
	$\Delta_v H$	(377–419)	40.4	392	A	[1987STE/MAL]
	$\Delta_v H$	(371–419)	40.8	394		[1984CAS/FRA]
	$\Delta_v H$	(377–418)	40.1	398		[1978SUN/VIS]
	$\Delta_v H$		39	415		[1977RAO/VIU]
	$\Delta_v H$		45.8 ± 0.2	298	C	[1972LAY/WAD]
	$\Delta_v H$	(328–464)	41.9	343	A	[1987STE/MAL, 1970DYK]
	$\Delta_v H$	(298–403)	47.7	313		[1950MAT/SUM]
	$\Delta_v H$	(304–419)	45.7	319		[1930NEL]
C₂H₂Cl₄S	[51174-93-7]	<i>bis</i> (dichloromethyl) sulfide				
	$\Delta_v H$	(355–462)	47.6	370	A	[1987STE/MAL]
C₂H₂FN	[503-20-8]	fluoroacetonitrile				
	$\Delta_v H$	(273–333)	38.1	288	A, GS	[1987STE/MAL, 1948RED/CHA4, 1970DYK]
C₂H₂F₂	[75-38-7]	1,1-difluoroethene				
	$\Delta_v H$		13.2	233	BG	[1955MEA/STA]
	$\Delta_v H$		9.5	273	BG	[1955MEA/STA]
C₂H₂F₃NO	[354-38-1]	trifluoroacetamide				
	$\Delta_{\text{trs}} H$		5.58	336.9		
	$\Delta_{\text{trs}} H$		11.5	347.6		
	$\Delta_{\text{fus}} H$		1.34	388.9		[2000DI/TAN]
	$\Delta_{\text{sub}} H$	(288–329)	81.0	302	I	[1987STE/MAL, 1978BER/SPI]
	$\Delta_{\text{sub}} H$	(288–329)	77.7 ± 1.4	298	I	[1978BER/SPI]
C₂H₂F₄	[811-97-2]	1,1,1,2-tetrafluoroethane				
	$\Delta_{\text{trs}} H$		3.62	124.1		
	$\Delta_{\text{fus}} H$		2.01	169.4		[1999ZHE/KAT]
	$\Delta_v H$	(221–246)	23.7	249	I	[2001KUL/DES]
	$\Delta_v H$		26.4	180		[1998BLA/KLI]
	$\Delta_v H$		25.0	200		[1998BLA/KLI]
	$\Delta_v H$		23.8	220		[1998BLA/KLI]
	$\Delta_v H$		22.7	240		[1998BLA/KLI]
	$\Delta_v H$	(279–363)	22.0	294		[1992ZHU/WU]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	Enthalpy							
C ₂ H ₂ F ₄ O ₂ S	[na]	fluorosulfurous acid, 2,2,2-trifluoroethyl ester						
	$\Delta_v H$				33.6			[1975DEM/KOV2]
C ₂ H ₂ F ₆ P ₂	[462-57-7]	1,2-bis-(trifluoromethyl) diphosphine						
	$\Delta_v H$			(233–292)	33.8	277	A, SG	[1987STE/MAL, 1958MAH/BUR]
C ₂ H ₂ F ₈ S	[65227-29-4]	pentafluoro (2,2,2-trifluoroethyl) sulfur						
	$\Delta_v H$				29.3			[1978DEM/FOX]
C ₂ H ₂ I ₂	[590-26-1]	<i>cis</i> 1,2-diiodoethylene						
	$\Delta_v H$			(302–425)	46.5	317	A	[1987STE/MAL, 1970DYK]
	$\Delta_v H$			(302–424)	47.3	317		[1950NOY/NOY]
C ₂ H ₂ I ₂	[590-27-2]	<i>trans</i> 1,2-diiodoethylene						
	$\Delta_{\text{sub}} H$			(253–265)	40.7	258	ME	[1933BRO/FRA, 1960JON, 1987STE/MAL]
	$\Delta_v H$			(350–403)	42.3	365	A	[1987STE/MAL, 1970DYK]
	$\Delta_v H$			(350–403)	43.8	365		[1950NOY/NOY]
C ₂ H ₂ O	[463-51-4]	ketene						
	$\Delta_v H$			(159–224)	20.4 ± 0.1	209	A, MM	[1987STE/MAL, 1969RUE]
C ₂ H ₂ O ₄	[144-62-7]	oxalic acid (anhydrous)						
	$\Delta_{\text{sub}} H(\alpha)$			(303–328)	93.4	316		[1987STE/MAL]
	$\Delta_{\text{sub}} H(\beta)$			(310–325)	93.3	318		[1987STE/MAL]
	$\Delta_{\text{sub}} H(\alpha)$				98.5			[1983DEW/BOW]
	$\Delta_{\text{sub}} H(\beta)$				92.5			[1983DEW/BOW]
	$\Delta_{\text{sub}} H(\alpha)$			(303–328)	93.7 ± 1.3	298	TE	[1975DEK/VAN]
	$\Delta_{\text{sub}} H$			(311–325)	97.9 ± 2.2	318		[1953BRA/COT, 1960JON]
	$\Delta_{\text{sub}} H$			(311–323)	93.3	317		[1953BRA/COT, 1960JON]
	$\Delta_{\text{sub}} H$			(292–320)	61.8	306	A	[1947GRA]
$\Delta_{\text{sub}} H$			(333–378)	90.6		GS	[1926NOY/WEB]	
C ₂ H ₃ Br	[593-60-2]	vinyl bromide						
	$\Delta_v H$			(224–319)	27.3	239	A	[1987STE/MAL, 1970DYK]
	$\Delta_v H$			(186–289)	24.8	274		[1937GUY/SCH, 1984BOU/FRI]
	$\Delta_v H$			(207–285)	26.9	270		[1934MEH, 1984BOU/FRI]
C ₂ H ₃ BrO	[506-96-7]	acetyl bromide						
	$\Delta_v H$			(289–334)	29.5	304	A	[1987STE/MAL]
	$\Delta_v H$			(275–333)	31.4	290		[1969DEV/ONE]
C ₂ H ₃ BrO ₂	[79-08-3]	bromoacetic acid						
	$\Delta_{\text{fus}} H$				13.9	319.2	DSC	[2001LAG/DIO]
	$\Delta_v H$			(327–481)	57.2	342	A	[1987STE/MAL]
C ₂ H ₃ BrO ₂					83.5 ± 3.0	298	ME	[2001LAG/DIO]
	$\Delta_{\text{sub}} H$							
C ₂ H ₃ Br ₃	[78-74-0]	1,1,2-tribromoethane						
	$\Delta_{\text{fus}} H$				9.11	244		[1991ACR]
	$\Delta_v H$			(368–511)	50.5	383	A	[1987STE/MAL, 1970DYK]
	$\Delta_v H$			(305–461)	52.9	321		[1947STU]
C ₂ H ₃ Cl	[75-01-4]	vinyl chloride						
	$\Delta_{\text{fus}} H$				4.92	119.3		[1996DOM/HEA]
	$\Delta_v H$			(243–288)	22.7	265		[1967DAN/GOL]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
	$\Delta_{\text{v}}H$	(213–273)	22.9	258		[1967HAC/MAT]
	$\Delta_{\text{v}}H$	(209–260)	23.3	245	A	[1987STE/MAL, 1959MCD/SHR, 1970DYK]
C₂H₃ClF₂	[75-68-3]	1-chloro-1,1-difluoroethane				
	$\Delta_{\text{fus}}H$		2.69	142.4		[1996DOM/HEA]
	$\Delta_{\text{v}}H$	(225–285)	24.2	240	EB	[1993SIL/WEB]
	$\Delta_{\text{v}}H$	(248–390)	22.7	263	A	[1987STE/MAL, 1970DYK]
	$\Delta_{\text{v}}H$		24.0	233	BG	[1955MEA/STA]
	$\Delta_{\text{v}}H$		21.9	273	BG	[1955MEA/STA]
	$\Delta_{\text{v}}H$		19.2	313	BG	[1955MEA/STA]
	$\Delta_{\text{v}}H$		15.4	353	BG	[1955MEA/STA]
C₂H₃ClF₃N	[16276-45-2]	N-chloro-N,1,1-trifluoroethanamine				
	$\Delta_{\text{v}}H$	(220–294)	30.8	279	BG	[1987STE/MAL, 1967LUS]
C₂H₃ClF₃P	[4669-76-5]	chloromethyl(trifluoromethyl)phosphine				
	$\Delta_{\text{v}}H$	(236–294)	30.9	279		[1987STE/MAL]
C₂H₃ClO	[75-36-5]	acetyl chloride				
	$\Delta_{\text{v}}H$	(273–323)	24.5	288	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$	(267–324)	31.5	282	A	[1987STE/MAL, 1959MCD/SHR, 1970DYK]
C₂H₃ClO₂	[79-11-8]	chloroacetic acid				
	$\Delta_{\text{fus}}H$		16.3	334.8	DSC	[2001LAG/DIO]
	$\Delta_{\text{fus}}H(\alpha)$		16.3	334.3		[1996DOM/HEA, 1991ACR]
	$\Delta_{\text{fus}}H(\beta)$		13.93	329.2		[1996DOM/HEA, 1991ACR]
	$\Delta_{\text{sub}}H$		82.2 ± 0.9	298	ME	[2001LAG/DIO]
	$\Delta_{\text{sub}}H$		75.3 ± 4.2			[1928STE/JOH, 1949DRE/MAR, 1970COX/PIL]
	$\Delta_{\text{v}}H$	(336–463)	61.1	351	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$	(377–463)	56.8	392	A	[1987STE/MAL, 1959MCD/SHR, 1970DYK]
	$\Delta_{\text{v}}H$	(396–460)	55.7	411		[1949DRE/SHR, 1949DRE/MAR]
C₂H₃Cl₂F	[1717-00-6]	1,1-dichloro-1-fluoroethane				
	$\Delta_{\text{v}}H$	(250–450)	28.7	265		[1997DUA/HWA]
	$\Delta_{\text{v}}H$	(270–312)	27.8	285	EB	[1992WEB]
C₂H₃Cl₃	[71-55-6]	1,1,1-trichloroethane				
	$\Delta_{\text{trs}}H$		0.21	205		
	$\Delta_{\text{trs}}H$		7.45	223.6		
	$\Delta_{\text{fus}}H$		1.88	240.1		[1973AND/COU, 1996DOM/HEA]
	$\Delta_{\text{v}}H$	(295–372)	32.3	310	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$	(349–408)	30.5	364	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$	(399–487)	29.4	414	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$	(479–545)	29.5	494	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$		32.5 ± 0.1	298	C	[1980MAJ/SVA]
	$\Delta_{\text{v}}H$		32.4	344		[1977RAO/VIU]
	$\Delta_{\text{v}}H$	(196–298)	37.6	211		[1973AMB/SPR]
	$\Delta_{\text{v}}H$		32.5 ± 0.1	298	C	[1972LAY/WAD, 1971MAN/RIN]
	$\Delta_{\text{v}}H$	(268–290)	33.4	279		[1944RUB/LEV]
	$\Delta_{\text{v}}H$		33.4 ± 0.1	284	C	[1944RUB/LEV]
C₂H₃Cl₃	[79-00-5]	1,1,2-trichloroethane				
	$\Delta_{\text{fus}}H$		11.38	237.1		[1996DOM/HEA]
	$\Delta_{\text{v}}H$	(316–384)	40.1 ± 0.6	298		[2007VAR/DRU]
	$\Delta_{\text{v}}H$	(316–384)	38.6	331	A	[1987STE/MAL]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$		40.2 ± 0.1	298	C	[1980MAJ/SVA]
	$\Delta_v H$		40.3 ± 0.1	298	C	[1972LAY/WAD]
	$\Delta_v H$	(302–428)	38.2	317		[1987STE/MAL, 1970DYK]
	$\Delta_v H$	(323–386)	38.3	338		[1949DRE/SHR, 1949DRE/MAR]
C₂H₃Cl₃O	[115-20-8]	2,2,2-trichloroethanol				
	$\Delta_{\text{fus}} H$		10.05	290.6		[1997JEN/SAN]
C₂H₃Cl₃O₂	[302-17-0]	chloral hydrate				
	$\Delta_{\text{sub}} H$	(263–319)	50.9	291	A	[1947STU]
	$\Delta_v H$	(300–348)	38.4	324	EB	[1994WIB/MOR]
	$\Delta_v H$	(325–370)	49.6	340	A	[1987STE/MAL]
	$\Delta_v H$	(263–369)	51.5	278		[1947STU]
C₂H₃F	[75-02-5]	vinyl fluoride				
	$\Delta_v H$	(124–201)	16.6	186	A	[1987STE/MAL, 1947STU]
C₂H₃FN₂O₅	[17003-75-7]	2-fluoro-2,2-dinitroethanol				
	$\Delta_{\text{sub}} H$		55.6 ± 2.1			[1977PED/RYL, 1968BAR/CAR]
	$\Delta_v H$	(313–373)	55.7	343		[1968BAR/CAR]
C₂H₃FO	[557-99-3]	acetyl fluoride				
	$\Delta_v H$	(195–281)	14.3	266	A	[1987STE/MAL]
C₂H₃FO₂	[78948-09-1]	acetyl hypofluorite				
	$\Delta_v H$	(209–253)	35.6 ± 2.4	231		[1985APP/MEN]
C₂H₃FO₂	[144-49-0]	fluoroacetic acid				
	$\Delta_v H$	(293–443)	52.3	308	A	[1987STE/MAL, 1970DYK]
	$\Delta_v H$	(293–443)	53.6	308	T	[1955JAS/MIL]
C₂H₃F₃	[420-46-2]	1,1,1-trifluoroethane				
	$\Delta_{\text{fus}} H$		6.19	161.9		[1991ACR]
	$\Delta_v H$	(236–280)	18.1	240	EB	[1996WEB/DEF]
	$\Delta_v H$	(236–280)	17.5	250	EB	[1996WEB/DEF]
	$\Delta_v H$	(236–280)	16.7	260	EB	[1996WEB/DEF]
	$\Delta_v H$	(236–280)	15.9	270	EB	[1996WEB/DEF]
	$\Delta_v H$		18.9	233	BG	[1955MEA/STA]
	$\Delta_v H$		16.4	273	BG	[1955MEA/STA]
	$\Delta_v H$		13.8	303	BG	[1955MEA/STA]
	$\Delta_v H$		8.7	333	BG	[1955MEA/STA]
	$\Delta_v H$	(174–226)	20.5	211		[1944RUS/GOL]
	$\Delta_v H$		19.2 ± 0.1	224	C	[1944RUS/GOL]
C₂H₃F₃N₂	[690-21-1]	1,1,1-trifluoroazomethane				
	$\Delta_v H$	(240–273)	26.4	258	A	[1987STE/MAL]
C₂H₃F₃O	[421-14-7]	trifluoromethyl methyl ether				
	$\Delta_v H$	(233–313)	22.5	248	A	[1992SAL/WAN]
C₂H₃F₃O	[75-89-8]	2,2,2-trifluoroethanol				
	$\Delta_v H$	(276–302)	45.9	289	A	[1987STE/MAL]
	$\Delta_v H$	(298–328)	44.0	313	A	[1987STE/MAL, 1970DYK]
	$\Delta_v H$	(298–328)	41.5	313	MM	[1973ROC/SYM]
	$\Delta_v H$	(273–298)	44.5	285	MM	[1967MEE/GOL]
C₂H₃F₃O₂S	[30957-42-7]	methanesulfinic acid, trifluoromethyl ester				
	$\Delta_v H$		31.8	346		[1971SAU/SHR2]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
	Enthalpy							
C ₂ H ₃ F ₅ O ₃ S	[60672-60-8]	(ethaneperoxoato) pentafluoro sulfur		(217–377)	36.2	297		[1999DYK/SVO, 1976HOP/DES]
	$\Delta_{\text{v}}H$							
C ₂ H ₃ F ₅ S	[865-54-3]	vinylsulfur pentafluoride			28.5	314		[1961CAS/RAY2]
	$\Delta_{\text{v}}H$							
C ₂ H ₃ IO	[507-02-8]	acetyl iodide		(276–302)	37.1	289	A	[1987STE/MAL, 1969DEV/ONE]
	$\Delta_{\text{v}}H$							
C ₂ H ₃ IO ₂	[64-69-7]	iodoacetic acid			15.5	355.1	DSC	[2001LAG/DIO]
	$\Delta_{\text{fus}}H$							
C ₂ H ₃ IO ₂					86.5 ± 1.0	298	ME	[2001LAG/DIO]
	$\Delta_{\text{sub}}H$							
C ₂ H ₃ N	[75-05-8]	acetonitrile						
	$\Delta_{\text{trs}}H$				0.9	216.9		
	$\Delta_{\text{fus}}H$				8.17	229.3		[1996DOM/HEA]
	$\Delta_{\text{v}}H$	(302–353)	33	298	EB		[2004ANT/GAL]	
	$\Delta_{\text{v}}H$		33	298			[1983AN/MAN]	
	$\Delta_{\text{v}}H$	(288–362)	33.8	303			[1974DOJ/HEI]	
	$\Delta_{\text{v}}H$	(314–355)	33.3	329	A, EB		[1987STE/MAL, 1971MEY/REN]	
	$\Delta_{\text{v}}H$	(299–343)	34.8	315	BG		[1971HAL/BAL]	
C ₂ H ₃ NO	[624-83-9]	methyl isocyanate		(265–308)	29.9	280	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$			(253–310)	31.7	268	A	[1987STE/MAL]
C ₂ H ₃ NO ₃	[471-47-6]	oxalic acid, monoamide			108.9 ± 2.1	298	ME	[1988NUN/BAR]
	$\Delta_{\text{sub}}H$			(355–363)	107.9	359	ME	[1953BRA/CLE2, 1960JON, 1987STE/MAL]
C ₂ H ₃ NO ₅	[2278-22-0]	acetyl nitro peroxide		(277–330)	34.6	292	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$							
C ₂ H ₃ NS	[556-64-9]	methyl thiocyanate		(259–406)	40.7	274	A	[1987STE/MAL, 1947STU, 1999DYK/SVO]
	$\Delta_{\text{v}}H$							
C ₂ H ₃ NS	[556-61-6]	methyl isothiocyanate		(238–293)	31.5	266	A	[1947STU]
	$\Delta_{\text{sub}}H$			(309–392)	37.4	324	A	[1987STE/MAL, 1999DYK/SVO]
	$\Delta_{\text{v}}H$			(283–323)	37.3	298		[1935BAU/BUR, 1984BOU/FRI]
C ₂ H ₃ N ₃	[288-88-0]	1,2,4-triazole						
	$\Delta_{\text{fus}}H$				16.1	393.5		[1996DOM/HEA]
	$\Delta_{\text{fus}}H$				10.76 ± 0.08	393.3		[1999SAB/PER]
	$\Delta_{\text{sub}}H$				80.7 ± 0.5	298	C	[1999SAB/PER]
	$\Delta_{\text{sub}}H$				84.0 ± 0.7	298	ME	[1989JIM/ROU]
C ₂ H ₃ N ₃ O ₆	[595-86-8]	1,1,1-trinitroethane						
	$\Delta_{\text{trs}}H$				4.6	311.7		
	$\Delta_{\text{fus}}H$				11.72	329.2	DSC	[1969ROS, 1969ROS/HOL]
C ₂ H ₃ N ₃ O ₇	[918-54-7]	2,2,2-trinitroethanol						
	$\Delta_{\text{sub}}H$				72.0 ± 8.8	298		[1999MIR/VOR]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{trs}}H$		17.99	312.5		
	$\Delta_{\text{fus}}H$		2.72	344.9		[1969ROS/HOL]
C ₂ H ₄	[74-85-1]	ethylene				
	$\Delta_{\text{fus}}H$		3.35	104		[1996DOM/HEA]
	$\Delta_{\text{sub}}H$	(79–104)	18.4	91.5	A,MS	[1987STE/MAL, 1951TIC/LOS]
	$\Delta_{\text{sub}}H$	(77–103)	15.0			[1982MEN]
	Δ_vH	(252–282)	14.0	267	A	[1987STE/MAL]
	Δ_vH	(170–273)	13.7	258	A	[1987STE/MAL]
	Δ_vH	(120–170)	14.4	155	A	[1987STE/MAL]
	Δ_vH	(169–211)	13.7	196	A	[1987STE/MAL]
	Δ_vH	(209–254)	13.6	239	A	[1987STE/MAL]
	Δ_vH	(120–182)	14.1	167	A	[1987STE/MAL, 1970DYK]
	Δ_vH	(150–190)	14.0	175		[1950MIC/WAS]
	Δ_vH	(148–174)	14.3	161		[1940LAM/ROP]
Δ_vH	(124–171)	14.4	156		[1937EGA/KEM]	
C ₂ H ₄ BrCl	[593-96-4]	1-bromo-1-chloroethane				
	Δ_vH	(290–356)	33.1	305	A	[1987STE/MAL]
	Δ_vH	(237–356)	46.7	252		[1947STU]
C ₂ H ₄ BrCl	[107-04-0]	1-bromo-2-chloroethane				
	$\Delta_{\text{trs}}H$		3.1	182		
	$\Delta_{\text{fus}}H$		9.62	256.4		[1996DOM/HEA]
	Δ_vH		37.6 ± 0.1	308	C	[1992SVO/KUB2]
	Δ_vH		37.3 ± 0.1	315	C	[1992SVO/KUB2]
	Δ_vH		36.9 ± 0.1	323	C	[1992SVO/KUB2]
	Δ_vH		36.6 ± 0.1	330	C	[1992SVO/KUB2]
	Δ_vH	(244–379)	36.4 ± 0.1	338	C	[1992SVO/KUB2]
Δ_vH		39.5			[1947STU]	
C ₂ H ₄ Br ₂	[557-91-5]	1,1-dibromoethane				
	Δ_vH	(301–421)	39.6	316	E	[1987STE/MAL, 1956MAN, 1970DYK]
C ₂ H ₄ Br ₂	[106-93-4]	1,2-dibromoethane				
	$\Delta_{\text{trs}}H$		1.94	249.5		[1996DOM/HEA]
	$\Delta_{\text{fus}}H$		10.94	283		
	$\Delta_{\text{sub}}H$	(229–248)	54.8	239		[1948NIT/SEK]
	$\Delta_{\text{sub}}H$	(251–281)	49.8	258	A	[1948NIT/SEK, 1947STU, 1987STE/MAL]
	Δ_vH	(331–426)	41.7 ± 0.8	298		[2007VAR/DRU]
	Δ_vH	(283–323)	42.4	298		[1994GRA/PER]
	Δ_vH	(325–403)	41.7 ± 0.8	298		[1993VAR/PUC]
	Δ_vH		41.7 ± 0.1	308	C	[1992SVO/KUB2]
	Δ_vH		41.7 ± 0.1	315	C	[1992SVO/KUB2]
	Δ_vH		41.6 ± 0.1	323	C	[1992SVO/KUB2]
	Δ_vH		41.5 ± 0.1	330	C	[1992SVO/KUB2]
	Δ_vH		41.4 ± 0.1	338	C	[1992SVO/KUB2]
	Δ_vH	(283–317)	41.8	298	A	[1987STE/MAL]
	Δ_vH	(316–488)	40.0	331	A	[1987STE/MAL]
	Δ_vH	(404–578)	37.4	419	A	[1987STE/MAL]
	Δ_vH	(285–298)	49.6	291	MM,A	[1957CAL]
	Δ_vH	(325–404)	39.6	340		[1949DRE/SHR, 1949DRE/MAR]
	Δ_vH	(283–317)	41.8	300		[1948NIT/SEK]
	Δ_vH	(246–404)	31.1	261		[1947STU]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound					
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference	
C ₂ H ₄ ClF	[762-50-5]	1-chloro-2-fluoroethane					
	$\Delta_v H$	(288–327)	32.1	303	A	[1987STE/MAL]	
C ₂ H ₄ ClN ₃	[53422-48-3]	1-chloro-2-azidoethane					
	$\Delta_v H$	(273–333)	43.8	288	A	[1987STE/MAL]	
C ₂ H ₄ Cl ₂	[75-34-3]	1,1-dichloroethane					
	$\Delta_{\text{fus}} H$		7.87	176.2		[1991ACR]	
	$\Delta_v H$	(326–345)	33.5	336		[1987GAR/TRE]	
	$\Delta_v H$	(323–535)	29.2	338	A	[1987STE/MAL]	
	$\Delta_v H$	(363–535)	28.2	378	A	[1987STE/MAL]	
	$\Delta_v H$		30.6 ± 0.1	298	C	[1972LAY/WAD]	
	$\Delta_v H$	(234–290)	31.9	275		[1956LI/PIT]	
	$\Delta_v H$	(213–330)	34.4	228		[1947STU]	
C ₂ H ₄ Cl ₂	[107-06-2]	1,2-dichloroethane					
	$\Delta_{\text{fus}} H$		8.83	237.2		[1996DOM/HEA]	
	$\Delta_v H$	(299–356)	35.2 ± 0.4	298		[2007VAR/DRU]	
	$\Delta_v H$		34.4	298	GC	[1994CAR/LAY]	
	$\Delta_v H$		35.1 ± 0.1	298	C	[1989AN/HU]	
	$\Delta_v H$	(356–558)	31.1	371	A	[1987STE/MAL]	
	$\Delta_v H$	(279–374)	34.8	294	A	[1987STE/MAL]	
	$\Delta_v H$	(368–524)	31.1	383	A	[1987STE/MAL]	
	$\Delta_v H$	(523–561)	40.8	538	A	[1987STE/MAL]	
	$\Delta_v H$	(301–357)	34.7	316		[1982GUT/KNA]	
	$\Delta_v H$		35.2 ± 0.1	298	C	[1980MAJ/SVA]	
	$\Delta_v H$	(279–434)	34.8	294		[1987STE/MAL, 1970DYK]	
C ₂ H ₄ Cl ₂ S	[3592-44-7]	<i>bis</i> (chloromethyl) sulfide					
	$\Delta_v H$	(320–430)	45.1	335	A	[1987STE/MAL, 1999DYK/SVO]	
	C ₂ H ₄ D ₂ O ₂	[na]	dihydroxyethane-d2				
		$\Delta_{\text{fus}} H$		9.75	258.8		[1967NIK/RAB]
	C ₂ H ₄ FNO ₃	[763-97-3]	2-fluoroethyl nitrate				
		$\Delta_v H$	(273–333)	38.3	288	GS	[1987STE/MAL, 1948RED/CHA4, 1970DYK]
	C ₂ H ₄ F ₂	[75-37-6]	1,1-difluoroethane				
		$\Delta_{\text{fus}} H$		1.57	154.6		[1998MAG]
		$\Delta_v H$	(218–248)	22.7	249	I	[2001KUL/DES]
		$\Delta_v H$	(303–333)	22.1	318		[1999LIM/PAR]
$\Delta_v H$		(219–273)	23.3	234	EB	[1993SIL/WEB]	
$\Delta_v H$		(250–386)	21.8	265	A	[1987STE/MAL]	
$\Delta_v H$		(193–275)	22.1	260	A, E	[1987STE/MAL, 1970DYK, 1956MAN]	
$\Delta_v H$			21.8	233	BG	[1955MEA/STA]	
$\Delta_v H$			20.4	273	BG	[1955MEA/STA]	
$\Delta_v H$			17.8	313	BG	[1955MEA/STA]	
C ₂ H ₄ F ₃ NS	[62067-12-3]	1,1,1-trifluoro-N-methyl methanesulfenamide					
	$\Delta_v H$	(223–294)	33.6	279	A	[1987STE/MAL, 1960EME/NAB]	
	C ₂ H ₄ F ₃ OP	[6395-71-7]	(trifluoromethyl)phosphinous acid, methyl ester				

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
	$\Delta_{\text{v}}H$	(194–291)	29.4	276	A	[1987STE/MAL]
C ₂ H ₄ F ₃ OP	[26348-89-0]	methyl(trifluoromethyl)phosphine oxide				
	$\Delta_{\text{v}}H$	(305–322)	50.7	313		[1970BUR/KAN]
C ₂ H ₄ F ₆ OS	[na]	pentafluoro(2-fluoroethoxy) sulfur				
	$\Delta_{\text{v}}H$	(290–364)	39.3	305	A	[1987STE/MAL, 1962WIL/CAD, 1999DYK/SVO]
C ₂ H ₄ I ₂	[624-73-7]	1,2-diiodoethane				
	$\Delta_{\text{sub}}H$		65.7 ± 4.1			[1954ABR/DAV, 1970COX/PIL]
	$\Delta_{\text{v}}H$		49.8	298	GC	[1994CAR/LAY]
	$\Delta_{\text{v}}H$	(371–526)	47.7	386	A	[1987STE/MAL, 1970DYK]
C ₂ H ₄ N ₂ O ₂	[628-36-4]	diformylhydrazine				
	$\Delta_{\text{sub}}H$	(340–373)	205.1 ± 0.7	356	ME	[1980LEB/KAL]
	$\Delta_{\text{sub}}H$	(370–403)	100.8			[1956SUZ/ONI, 1960JON]
C ₂ H ₄ N ₂ O ₂	[471-46-5]	oxamide				
	$\Delta_{\text{trs}}H$		1.88	356.2		
	$\Delta_{\text{trs}}H$		3.56	455.5		
	$\Delta_{\text{trs}}H$		6.24	494.3	DSC	[2006BAD/DEL]
		Note: decomposed prior to melting				
	$\Delta_{\text{sub}}H$		117.3 ± 1.2	298	TE,ME	[1988NUN/BAR]
	$\Delta_{\text{sub}}H$	(370–398)	115.8	387	TE,ME	[1983DEW/VAN]
	$\Delta_{\text{sub}}H$	(353–369)	113	361	ME	[1953BRA/CLE, 1960JON, 1970COX/PIL]
C ₂ H ₄ N ₂ O ₄	[600-40-8]	1,1-dinitroethane				
	$\Delta_{\text{v}}H$	(303–363)	51.0	318	A	[1987STE/MAL]
C ₂ H ₄ N ₂ O ₆	[628-96-6]	ethylene glycol dinitrate				
	$\Delta_{\text{v}}H$	(283–535)	70.5	298	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$	(343–465)	55.3	358	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$	(240–298)	68.3	255		[1987STE/MAL, 1977PEL]
	$\Delta_{\text{v}}H$	(278–390)	62.3 ± 0.4			[1941BEL]
	$\Delta_{\text{v}}H$	(293–323)	68.6 ± 0.4			[1938BRA]
C ₂ H ₄ N ₂ S ₂	[628-96-6]	dithiooxamide				
	$\Delta_{\text{sub}}H$		103.8	298	TE,ME	[1988NUN/BAR]
	$\Delta_{\text{sub}}H$	(345–372)	105.1	361	TE,ME	[1983DEW/VAN]
	$\Delta_{\text{sub}}H$	(360–378)	105.4	369	ME	[1953BRA/CLE2, 1960JON, 1987STE/MAL]
C ₂ H ₄ N ₄	[461-58-5]	dicyandiamide				
	$\Delta_{\text{trs}}H$		2.98	269.5		
	$\Delta_{\text{fus}}H$		22.96	487.6		[1997ZHA/TAN]
	$\Delta_{\text{sub}}H$	(420–450)	128.7	436	TE,ME	[1983DEW/VAN]
C ₂ H ₄ N ₄	[16681-77-9]	1-methyltetrazole				
	$\Delta_{\text{fus}}H$		15.7	315		[1990KOZ/SIM3]
	$\Delta_{\text{sub}}H$	(282–311)	86.7 ± 1.9		ME	[1990KOZ/SIM]
C ₂ H ₄ N ₄	[16681-78-0]	2-methyltetrazole				
	$\Delta_{\text{fus}}H$		12.37	286		[1990KOZ/SIM3]
C ₂ H ₄ N ₄	[4076-36-2]	5-methyltetrazole				
	$\Delta_{\text{fus}}H$		16.0	418		[1990KOZ/SIM3]
	$\Delta_{\text{sub}}H$	(323–418)	93.8 ± 0.5		ME	[1990KOZ/SIM]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound					
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference	
C ₂ H ₄ N ₄	[61-82-5]	1 <i>H</i> -1,2,4-triazol-3-amine					
	$\Delta_{\text{fus}}H$		21.93	428.3		[1990DON/DRE]	
C ₂ H ₄ O	[75-07-0]	acetaldehyde					
	$\Delta_{\text{trs}}H$		2.31	149.8			
	$\Delta_{\text{fus}}H$		1.72	242.9		[1996DOM/HEA]	
	$\Delta_{\text{v}}H$	(293–377)	26.0	308	A	[1987STE/MAL]	
	$\Delta_{\text{v}}H$	(293–345)	26.3	308		[1977KIM/KIM]	
	$\Delta_{\text{v}}H$	(272–294)	27.6	283	A	[1987STE/MAL, 1970DYK]	
	$\Delta_{\text{v}}H$	(293–377)	26.9	298	EB	[1963BUL/SER, 2003VER/KRA2]	
C ₂ H ₄ O	[75-21-8]	ethylene oxide (oxirane)					
	$\Delta_{\text{fus}}H$		5.17	160.7		[1996DOM/HEA]	
	$\Delta_{\text{v}}H$	(283–385)	25.9	298	A	[1987STE/MAL]	
	$\Delta_{\text{v}}H$	(239–284)	26.8	269	A	[1987STE/MAL, 1959MCD/SHR, 1970DYK]	
	$\Delta_{\text{v}}H$	(223–284)	26.8	269	A	[1987STE/MAL, 1949GIA/GOR]	
C ₂ H ₄ O		(268–313)	26.9	290		[1937MOO/KAN]	
	C ₂ D ₄ O	[6552-57-4]	ethylene oxide-d ₄ oxide				
	$\Delta_{\text{v}}H$	(230–273)	27.6	258		[1952LEI/MOR, 1984BOU/FRI]	
	C ₂ H ₄ OS	[507-09-5]	thioacetic acid				
$\Delta_{\text{v}}H$	(307–360)	35.2	333		[1999DYK/SVO]		
C ₂ H ₄ O ₂	[64-19-7]	acetic acid					
	$\Delta_{\text{fus}}H$		11.72	298.7		[1996DOM/HEA, 1982MAR/AND]	
	$\Delta_{\text{fus}}H$		10.83	289.8		[1911LOU/DUP]	
	$\Delta_{\text{fus}}H$		11.52	283.7		[1910MEY]	
	$\Delta_{\text{sub}}H$	(213–230)	67.3 ± 1	223	TE,ME	[1978CAL/CAL]	
	$\Delta_{\text{sub}}H$	(213–230)	70 ± 1	213	TE,ME	[1978CAL/CAL]	
	$\Delta_{\text{v}}H$	(320–395)	40.9	335		[2001VER/VAZ]	
	$\Delta_{\text{v}}H$	(345–383)	39.1	360	EB	[2001MUN/KRA]	
	$\Delta_{\text{v}}H$	(303–378)	50.3	298	CGC	[2000VER2]	
	$\Delta_{\text{v}}H$	(391–550)	37.9	406	A	[1987STE/MAL]	
	$\Delta_{\text{v}}H$	(290–396)	42.0	305	A	[1987STE/MAL]	
	$\Delta_{\text{v}}H$	(391–447)	38.7	406	A	[1987STE/MAL]	
	$\Delta_{\text{v}}H$	(437–535)	38.1	452	A	[1987STE/MAL]	
	$\Delta_{\text{v}}H$	(525–593)	38.8	540	A	[1987STE/MAL]	
	$\Delta_{\text{v}}H$		43.0	308		[1983TAM/DRA]	
	$\Delta_{\text{v}}H$	(289–392)	41.6	304	A	[1987STE/MAL, 1970DYK]	
	$\Delta_{\text{v}}H$ (monomer)		23.3 ± 0.1	298	C	[1970KON/WAD]	
$\Delta_{\text{v}}H$		51.6 ± 1.6	298	C	[1970KON/WAD]		
$\Delta_{\text{v}}H$	(325–391)	40.3	340		[1959MCD/SHR]		
$\Delta_{\text{v}}H$	(303–399)	41.6	318	MM	[1954POR/RIT]		
(C ₂ H ₄ O ₂) ₂	[na]	acetic acid dimer					
	$\Delta_{\text{sub}}H$	(213–230)	70.2 ± 1	213	TE	[1978CAL/CAL]	
	$\Delta_{\text{sub}}H$	(213–230)	68.9 ± 1	213	ME	[1978CAL/CAL]	
C ₂ H ₄ O ₂	[107-31-3]	methyl formate					
	$\Delta_{\text{v}}H$	(279–305)	29.6	292	A	[1987STE/MAL]	
	$\Delta_{\text{v}}H$	(305–443)	28.4	320	A	[1987STE/MAL]	
	$\Delta_{\text{v}}H$		28.7 ± 0.1	293	C	[1976CIH/HYN]	
$\Delta_{\text{v}}H$		27.9 ± 0.1	305	C	[1976CIH/HYN]		

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$		27.4 ± 0.1	313	C	[1976CIH/HYN]
	$\Delta_v H$	(261–305)	30.1	283	BG	[1971HAL/BAL]
	$\Delta_v H$	(294–304)	U 52.7	299		[1928NEL, 1984BOU/FRI]
C₂H₄O₂	[141-46-8]	hydroxyacetaldehyde				
	$\Delta_v H$	(273–304)	70 ± 5	288		[2010PET/REY]
C₂H₄O₂S	[68-11-1]	mercaptoacetic acid				
	$\Delta_v H$	(333–427)	56.8	348	A	[1987STE/MAL, 1999DYK/SVO]
C₂H₄O₃	[289-14-5]	ethylene ozonide				
	$\Delta_v H$	(273–289)	34.8	281	A	[1987STE/MAL, 1956GAR/SCH]
C₂H₄O₃	[79-14-1]	hydroxyacetic acid				
	$\Delta_v H$	(350–375)	51.8	362	A	[1987STE/MAL]
C₂H₄O₃	[79-21-0]	peroxyacetic acid				
	$\Delta_v H$	(273–383)	44.2	288	A	[1987STE/MAL, 1970DYK]
C₂H₄O₃	[7456-87-3]	methyl hydrogen carbonate				
	$\Delta_{\text{sub}} H$	(204–237)	18.2 ± 1.6	220		[1973BEH/GAT]
C₂H₄S	[420-12-2]	ethylene sulfide				
	$\Delta_v H$	(291–361)	30.5	306	A	[1987STE/MAL, 1952GUT/SCO2, 1999DYK/SVO]
	$\Delta_v H$		30.3	298		[1971WIL/ZWO]
C₂H₅Br	[74-96-4]	ethyl bromide				
	$\Delta_v H$	(334–504)	26.9	349	A	[1987STE/MAL]
	$\Delta_v H$	(326–454)	26.6	341	A	[1987STE/MAL]
	$\Delta_v H$	(452–503)	31.0	467	A	[1987STE/MAL]
	$\Delta_v H$		27.6 ± 0.1	305	C	[1977SVO/MAJ]
	$\Delta_v H$		27.0 ± 0.1	312	C	[1977SVO/MAJ]
	$\Delta_v H$		26.2 ± 0.1	323	C	[1977SVO/MAJ]
	$\Delta_v H$	(225–333)	30.6	240	E	[1987STE/MAL, 1961LI/ROS, 1970DYK]
	$\Delta_v H$	(301–348)	27.9	316		[1930ZMA, 1984BOU/FRI]
C₂H₅BrO	[540-51-2]	2-bromoethanol				
	$\Delta_v H$		54.1 ± 0.4	298	C	[2007BER/MIN]
C₂H₅Cl	[75-00-3]	ethyl chloride				
	$\Delta_{\text{fus}} H$		4.45	134.8		[1991ACR]
	$\Delta_v H$	(285–344)	25.1	300	A	[1987STE/MAL]
	$\Delta_v H$	(334–413)	24.4	349	A	[1987STE/MAL]
	$\Delta_v H$	(403–460)	24.4	418	A	[1987STE/MAL]
	$\Delta_v H$	(207–305)	27.8	222	A, E	[1987STE/MAL, 1961LI/ROS, 1970DYK]
	$\Delta_v H$	(218–285)	25.9	270		[1948GOR/GIA]
	$\Delta_v H$		24.8		C	[1926YAT]
C₂H₅ClO	[107-07-3]	2-chloroethanol				
	$\Delta_v H$		48.3 ± 0.4	298	C	[2007BER/MIN]
	$\Delta_v H$	(328–401)	43.3	343	A	[1987STE/MAL]
	$\Delta_v H$	(323–363)	46.9	338		[1973GOT/MEN]
	$\Delta_v H$	(363–403)	39.1	378		[1973GOT/MEN]
	$\Delta_v H$	(269–401)	45.7	284		[1947STU]
C₂H₅ClO	[107-30-2]	methyl(chloromethyl) ether				
	$\Delta_v H$	(290–332)	32.2	305	A	[1987STE/MAL]
C₂H₅ClO₂S	[594-44-5]	ethane sulfonyl chloride				

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	Enthalpy							
	$\Delta_v H$	(349–449)		47.7	364			[1999DYK/SVO]
	$\Delta_v H$	(233–263)		56.4	248		A	[1987STE/MAL, 1999DYK/SVO]
C₂H₅Cl₂P	[1498-40-4]	dichloroethyl phosphine						
	$\Delta_v H$	(313–385)		36.8	328		A	[1987STE/MAL]
C₂H₅Cl₂OP	[1066-50-8]	ethylphosphonic acid dichloride						
	$\Delta_v H$			42.7 ± 4.2				[1956NEA/WIL, 1982PIL/SKI]
C₂H₅F	[353-36-6]	ethyl fluoride						
	$\Delta_v H$	(200–235)		20.7	236		I	[2001KUL/DES]
	$\Delta_v H$	(275–353)		20.2	290		A	[1987STE/MAL]
	$\Delta_v H$	(235–280)		20.5	265		A	[1987STE/MAL]
	$\Delta_v H$	(343–375)		20.7	358		A	[1987STE/MAL]
	$\Delta_v H$	(170–255)		U 4.2	240			[1975IWA/DAT]
	$\Delta_v H$	(173–251)		20.8	236		E	[1987STE/MAL, 1961LI/ROS, 1970DYK]
	$\Delta_v H$	(156–241)		22.0	226			[1947STU]
C₂H₅FO	[371-62-0]	2-fluoroethanol						
	$\Delta_v H$	(273–333)		44.1	288		GS	[1987STE/MAL, 1948RED/CHA4, 1970DYK]
C₂H₅FO₃S	[371-69-7]	ethyl fluorosulfonate						
	$\Delta_v H$	(273–333)		38.5	288		GS	[1987STE/MAL, 1948RED/CHA4, 1970DYK]
C₂H₅F₂N	[758-18-9]	N,N-difluoroethylamine						
	$\Delta_v H$	(241–259)		27.3	250		A	[1987STE/MAL]
	$\Delta_v H$			25.7	288			[1960FRA]
C₂H₅F₃NP	[4669-74-3]	methyl(trifluoromethyl) phosphinic acid amide						
	$\Delta_v H$	(238–294)		36.8	279			[1987STE/MAL]
C₂H₅I	[75-03-6]	ethyl iodide						
	$\Delta_v H$	(313–353)		31.7	298		CGC	[1995CHI/HOS]
	$\Delta_v H$			31.9 ± 0.1	298		C	[1968WAD]
	$\Delta_v H$	(249–369)		33.6	264		E	[1987STE/MAL, 1961LI/ROS]
	$\Delta_v H$	(219–345)		34.7	234			[1947STU]
	$\Delta_v H$	(254–293)		32	278			[1944MIL2]
	$\Delta_v H$	(303–333)		31.7	318			[1929SMY/ENG]
C₂H₅IO	[151-56-4]	2-iodoethanol						
	$\Delta_v H$			57.0 ± 0.2	288		C	[2007BER/MIN]
C₂H₅N	[151-56-4]	aziridine						
	$\Delta_v H$	(274–303)		34.9	288		A	[1987STE/MAL]
C₂H₅NO	[107-29-9]	acetaldehyde oxime						
	$\Delta_v H$	(288–388)		48	303		A	[1987STE/MAL]
C₂H₅NO	[60-35-5]	acetamide						
	$\Delta_{\text{fus}} H$			15.6	353			[1996DOM/HEA]
	$\Delta_{\text{sub}} H$			78.5 ± 0.3				[1998PRI/HAW]
	$\Delta_{\text{sub}} H$	(273–293)		77.8	284		TE,ME	[1983DEW/VAN]
	$\Delta_{\text{sub}} H$			77.2	298			[1983DEW/VAN]
	$\Delta_{\text{sub}} H$			78.7 ± 0.3				[1975BAR/PIL, 1977PED/RYL]
	$\Delta_{\text{sub}} H$			80.3 ± 1	298			[1971MOR2]
	$\Delta_{\text{sub}} H$			80.3 ± 1.3	298		C	[1965WAD]
	$\Delta_{\text{sub}} H$	(298–349)		77.4 ± 0.4	323		GS	[1959DAV/JON2, 1987STE/MAL]
	$\Delta_{\text{sub}} H$	(293–306)		U 57.2	300			[1952AIH]
	$\Delta_v H$	(381–492)		63.8	396		A	[1987STE/MAL]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(338–495)	60.9	353		[1947STU]
C ₂ H ₅ NO	[123-39-7]	N-methylformamide				
	$\Delta_{\text{trs}}H$		1.23	228.1		
	$\Delta_{\text{fus}}H$		10.44	270.6		[1999AHL/LOH]
	$\Delta_v H$	(340–440)	53.8	355		[1996USH/SED]
	$\Delta_v H$	(340–440)	54.4 ± 1.3	298		[1996USH/SED]
	$\Delta_v H$	(310–391)	54.5	325	A	[1987STE/MAL]
	$\Delta_v H$		56.2	298	A	[1985BAR/CAS, 1986VAR]
C ₂ H ₅ NO ₂	[109-95-5]	ethyl nitrite				
	$\Delta_v H$	(252–276)	25.7	264	A	[1987STE/MAL, 1937THO/DAI]
	$\Delta_v H$		27.8			[1934GOO]
C ₂ H ₅ NO ₂	[598-55-0]	methyl carbamate				
	$\Delta_{\text{fus}}H$		16.7	328.6		[1996DOM/HEA]
	$\Delta_{\text{sub}}H$	(287–305)	74.5 ± 0.8	296	GS	[1959DAV/JON]
C ₂ H ₅ NO ₂	[79-24-3]	nitroethane				
	$\Delta_{\text{fus}}H$		9.85	183.7		[1996DOM/HEA]
	$\Delta_v H$	(324–388)	38.6	339	EB	[1987STE/MAL, 1956TOO, 1970DYK]
	$\Delta_v H$	(252–387)	41.3	267		[1947STU]
C ₂ H ₅ NO ₂	[56-40-6]	glycine				
	$\Delta_{\text{sub}}H$	(408–431)	136.5 ± 2	419	TE,ME	[1979DEK/VOO]
	$\Delta_{\text{sub}}H$	(325–425)	U 96.2 ± 4	375	LE	[1977GAF/PIE]
	$\Delta_{\text{sub}}H$	(413–450)	138.1 ± 4.6	298	C	[1977NAG/SAB]
	$\Delta_{\text{sub}}H$	(453–471)	136.4 ± 4.0	462	ME	[1965SVE/CLY, 1970COX/PIL, 1964CLY/SVE, 1989CHI/GRO]
C ₂ H ₅ NO ₃	[625-58-1]	ethyl nitrate				
	$\Delta_{\text{fus}}H$		8.53	178.6		[1996DOM/HEA]
	$\Delta_v H$	(273–361)	37	288	A	[1987STE/MAL, 1970DYK]
	$\Delta_v H$	(273–343)	37.3	288		[1957GRA/PRA]
	$\Delta_v H$	(273–333)	37	288		[1956GRA/PRA]
C ₂ H ₅ NS	[62-55-5]	thioacetamide				
	$\Delta_{\text{fus}}H$		18.36	385.7		[1990DON/DRE]
	$\Delta_{\text{sub}}H$		83.3 ± 0.3	298	C	[1982INA/MUR, 1985MUR/SAK]
C ₂ H ₅ N ₃	[871-31-8]	azidoethane				
	$\Delta_v H$	(296–320)	31.5	308	A	[1987STE/MAL, 1970DYK]
	$\Delta_v H$	(253–298)	28.9	268	A	[1987STE/MAL, 1964GEI/KON, 1984BOU/FRI]
C ₂ H ₅ N ₃ O	[1517-05-1]	2-azidoethanol				
$\Delta_v H$		33.9 ± 1.3			[1997KOR/API]	
C ₂ H ₅ N ₃ O ₂	[na]	bis(nitrosomethyl)amine				
$\Delta_v H$	(276–426)	43.5	291		[1987STE/MAL, 1947STU]	

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₂ H ₅ N ₅	[5422-44-6]	1-methyl-5-aminotetrazole				
	$\Delta_{\text{sub}}H$	(379–438)	116.4 ± 1.7		ME	[1990KOZ/SIM]
C ₂ H ₅ N ₅	[6154-04-7]	2-methyl-5-aminotetrazole				
	$\Delta_{\text{sub}}H$	(310–373)	90.6 ± 1.1		ME	[1990KOZ/SIM]
C ₂ H ₆	[74-84-0]	ethane				
	$\Delta_{\text{fus}}H$		2.79	89.5		[1996DOM/HEA]
	$\Delta_{\text{trs}}H$		2.28	89.8		
	$\Delta_{\text{fus}}H$		0.58	90.3		[1976ATA/CHI]
	$\Delta_{\text{sub}}H$	(80–90)	22.6	85		[1972REG]
	$\Delta_{\text{sub}}H$		20.5	90	B	[1963BON]
	Δ_vH	(273–305)	15.3	288	A	[1987STE/MAL]
	Δ_vH	(154–185)	15.7	170	A	[1987STE/MAL]
	Δ_vH	(95–129)	17.7	114	A	[1987STE/MAL]
	Δ_vH	(185–229)	14.9	214	A	[1987STE/MAL]
	Δ_vH	(228–274)	14.9	259	A	[1987STE/MAL]
	Δ_vH	(91–144)	17.1	129		[1973CAR/KOB]
	Δ_vH		14.7	210		[1971WIL/ZWO]
	Δ_vH		14.7	184		[1937WIT/KEM]
Δ_vH	(136–200)	15.3	185		[1926LOO/WAL]	
C ₂ H ₆ BrF ₄ NS	[63324-17-4]	bromotetrafluoro(N-methylmethanaminato) sulfur				
	Δ_vH		38.1	372	I	[1977KIT/SHR2]
C ₂ H ₆ ClF ₄ NS	[63324-16-3]	chlorotetrafluoro(N-methylmethanaminato) sulfur				
	Δ_vH		36	359	I	[1977KIT/SHR2]
C ₂ H ₆ ClP	[811-62-1]	chlorodimethyl phosphine				
	$\Delta_{\text{sub}}H$	(233–268)	55.5	253		[1987STE/MAL, 1958BUR/SLO]
		(273–306)	32.9	288	A	[1987STE/MAL, 1958BUR/SLO]
C ₂ H ₆ Cl ₂ NP	[683-85-2]	(dimethylamino)dichlorophosphine				
	Δ_vH		40.8 ± 0.7	298	STG	[1995ALM/FIN2]
C ₂ H ₆ ClO ₃ P	[16672-87-0]	2-chloroethylphosphonic acid				
	$\Delta_{\text{fus}}H$		14.79	347.9	DSC	[1990DON/DRE]
C ₂ H ₆ FN	[14722-43-1]	fluorodimethylamine				
	Δ_vH	(249–273)	29.9	261	A	[1987STE/MAL]
C ₂ H ₆ FO ₃ P	[5954-50-7]	dimethylfluorophosphate				
	Δ_vH	(273–333)	44.4	288	A, GS	[1987STE/MAL, 1948RED/CHA4]
C ₂ H ₆ F ₂ NP	[814-97-1]	(dimethylamino)difluorophosphine				
	Δ_vH	(263–313)	29.3	288	I	[1964CAV]
C ₂ H ₆ F ₃ NOS	[22519-52-4]	(dimethylaminato)trifluoroxyo sulfur				
	Δ_vH	(313–357)	44.9	335		[1968GLE/VON2]
C ₂ H ₆ F ₃ NS	[3880-03-3]	(dimethylamino) sulfur trifluoride				
	Δ_vH	(296–327)	40.5	311	A	[1987STE/MAL, 1999DYK/SVO]
C ₂ H ₆ F ₄ NP	[2353-98-2]	(dimethylamino) tetrafluorophosphorane				
	Δ_vH		37.1			[1966BRO/FRA]
C ₂ H ₆ N ₂	[503-28-6]	azomethane				
	Δ_vH	(195–273)	26.4	258	A	[1987STE/MAL]
	Δ_vH	(209–236)	25.3	222	A	[1987STE/MAL]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₂ H ₆ N ₂	[na]	methylammonium cyanide				
	$\Delta_v H$	(251–295)	49.1	280	A	[1987STE/MAL, 1973DIE/MAR]
C ₂ H ₆ N ₂ O	[598-50-5]	N-methylurea				
	$\Delta_{\text{fus}}H$		12.5	372	DSC	[2005HAS/TAJ]
	$\Delta_{\text{fus}}H$		16.6	375	DSC	[1995FER/DEL]
	$\Delta_{\text{fus}}H$		14.06	378.1		[1990KAB/MIR2]
	$\Delta_{\text{sub}}H$	(331–365)	95.5 ± 0.5	298	GS	[2006EME/KAB]
	$\Delta_{\text{sub}}H$	(322–371)	96.9 ± 1.2	347	ME	[2003ZAI/KAB]
	$\Delta_{\text{sub}}H$	(322–371)	96.8 ± 1.2	350	ME	[2003ZAI/KAB]
	$\Delta_{\text{sub}}H$		94.4 ± 0.8	350	C	[2003ZAI/KAB]
	$\Delta_{\text{sub}}H$		94.4 ± 0.84	343	C	[1993KOZ/KAB]
	$\Delta_{\text{sub}}H$		97.1 ± 0.4	298		[1993KOZ/KAB]
	$\Delta_{\text{sub}}H$		94.9 ± 0.6	337	C	[1990KAB/MIR]
	$\Delta_{\text{sub}}H$		93.3 ± 1.2	355	TE	[1990PIA/FER, 1987FER/DEL2]
	$\Delta_{\text{sub}}H$		87.3	348		[1987FER/DEL2]
	$\Delta_{\text{sub}}H$		99.3 ± 0.7			[1986KRA/KOZ2]
				78.2	E	[1982AIR/CHA]
C ₂ H ₆ N ₂ O ₂	[4164-28-7]	N-methyl-N-nitromethanamine				
	$\Delta_{\text{sub}}H$	(315–324)	69.9	319	DBM	[1952BRA/COT, 1977PED/RYL]
C ₂ H ₆ N ₂ O ₂	[4164-28-7]	N-nitro-N-methylaminomethane				
	$\Delta_{\text{fus}}H$		37.66	327		[1987OYU/BR1]
C ₂ H ₆ N ₂ O ₂	[4164-28-7]	2-nitro-2-azapropene				
	$\Delta_{\text{fus}}H$		17.26	331.5	DSC	[1997ZEM]
C ₂ H ₆ N ₂ S	[598-52-7]	N-methylthiourea				
	$\Delta_{\text{fus}}H$		19.46	392.4		[2000DEL/JOZ]
	$\Delta_{\text{sub}}H$		112.9 ± 3	298	ME	[2000DEL/JOZ]
			111 ± 3.0	381	TE	[1994FER/MAR]
C ₂ H ₆ N ₂ S	[13849-02-0]	sulfur diimide, dimethyl				
	$\Delta_v H$	(248–298)	37.2	263	A	[1987STE/MAL, 1999DYK/SVO]
C ₂ H ₆ N ₄ O ₄	[505-71-5]	N,N'-dinitroethanediamine				
	$\Delta_{\text{fus}}H$		29.5	450		[1987OYU/BR1]
C ₂ H ₆ O	[115-10-6]	dimethyl ether				
	$\Delta_{\text{fus}}H$		4.94	131.7		[1996DOM/HEA]
	$\Delta_v H$	(183–265)	22.6	250	A	[1987STE/MAL]
	$\Delta_v H$	(180–249)	22.8	234	A	[1987STE/MAL]
	$\Delta_v H$	(293–360)	21.2	308	A	[1987STE/MAL]
	$\Delta_v H$	(349–400)	21.1	364	A	[1987STE/MAL]
	$\Delta_v H$	(241–303)	22.2	256	A	[1987STE/MAL]
	$\Delta_v H$	(171–248)	18.5	298		[1976AMB/ELL]
	$\Delta_v H$	(171–248)	21.4	248		[1976AMB/ELL]
	$\Delta_v H$	(195–248)	22.7	233		[1941KEN/SAG]
	$\Delta_v H$		21.5 ± 0.1	248	C	[1941KEN/SAG]
C ₂ H ₆ O	[64-17-5]	ethanol				
	$\Delta_{\text{trs}}H$		3.14	111.4		
	$\Delta_{\text{fus}}H$		4.64	158.8		[1996DOM/HEA]
	$\Delta_v H$	(311–351)	41.7	326		[2010MEJ/SEG]
	$\Delta_v H$	(298–348)	41.7	298		[2004NAS/ZIM]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	Enthalpy							
		$\Delta_v H$			38.9			[1999FAT]
		$\Delta_v H$	(323–357)		39.3	338		[1999AUC/LOR]
		$\Delta_v H$	(309–343)		40.7	321	EB	[1995DIO/SAN]
		$\Delta_v H$	(309–343)		42.4	298	EB	[1995DIO/SAN]
		$\Delta_v H$	(342–357)		40.5	357		[1990ORT/SUS]
		$\Delta_v H$			35.2	393	C	[1989VIN/WOR]
		$\Delta_v H$			30.6	423	C	[1989VIN/WOR]
		$\Delta_v H$			25.7	453	C	[1989VIN/WOR]
		$\Delta_v H$			21.8	473	C	[1989VIN/WOR]
		$\Delta_v H$			17.3	493	C	[1989VIN/WOR]
		$\Delta_v H$			14.2	503	C	[1989VIN/WOR]
		$\Delta_v H$			40.9	320	C	[1988DON/LIN]
		$\Delta_v H$			40.4	328	C	[1988DON/LIN]
		$\Delta_v H$			40.2	335	C	[1988DON/LIN]
		$\Delta_v H$			39.4	344	C	[1988DON/LIN]
		$\Delta_v H$			38.8	351	C	[1988DON/LIN]
		$\Delta_v H$	(320–359)		41.3	335	A	[1987STE/MAL]
		$\Delta_v H$	(210–271)		45.6	256	A	[1987STE/MAL]
		$\Delta_v H$	(193–223)		44	208	A	[1987STE/MAL]
		$\Delta_v H$	(320–359)		41.3	335	A	[1987STE/MAL]
		$\Delta_v H$	(349–374)		40.1	361	A	[1987STE/MAL]
		$\Delta_v H$	(370–464)		39.1	385	A	[1987STE/MAL]
		$\Delta_v H$	(459–514)		36.1	474	A	[1987STE/MAL]
		$\Delta_v H$	(292–353)		42.5	307	A	[1987STE/MAL]
		$\Delta_v H$	(243–303)		42.3	298		[1983SCH/STR]
		$\Delta_v H$	(271–373)		42.9	286		[1973WIL/ZWO]
		$\Delta_v H$			42.26 ± 0.02	298	C	[1971POL/BEN]
		$\Delta_v H$			41.0 ± 0.1	320	C	[1970COU/FEN]
		$\Delta_v H$			40.0 ± 0.1	335	C	[1970COU/FEN]
		$\Delta_v H$			38.7 ± 0.1	351	C	[1970COU/FEN]
		$\Delta_v H$	(293–366)		42.5	308	A, EB	[1987STE/MAL, 1970AMB/SPR]
		$\Delta_v H$	(288–348)		42.4	303		[1967VAN/SOC]
		$\Delta_v H$			42.3 ± 0.1	298	C	[1966WAD]
		$\Delta_v H$			42.2 ± 0.1	298	C	[1963MCC/LAI]
		$\Delta_v H$	(298–351)		42.2	313		[1949KRE/WIE]
		$\Delta_v H$			40.0	351		[1934OGU/ANJ]
		$\Delta_v H$	(286–351)		54.1	301		[1883KAH]
C₂H₆OS	[67-68-5]		dimethyl sulfoxide					
		$\Delta_{\text{fus}} H$			14.37	291.7		[1996DOM/HEA]
		$\Delta_v H$	(377–483)		48.6	392		[1999DYK/SVO]
		$\Delta_v H$	(353–383)		48.1	368	TGA	[1987ALN/ALS]
		$\Delta_v H$	(305–464)		51.7	320	A	[1987STE/MAL]
		$\Delta_v H$	(298–318)		52.3	308		[1974SAS/KON]
		$\Delta_v H$	(325–442)		50.6	340	MM	[1972JAK/VAN, 1984BOU/FRI]
		$\Delta_v H$	(303–423)		52.1	318		[1972NIS/HAK]
		$\Delta_v H$	(293–323)		52.5	308		[1969MES]
		$\Delta_v H$	(293–323)		52.9 ± 0.4	298	RG	[1948DOU]
C₂H₆OS	[60-24-2]		2-mercaptoethanol					
		$\Delta_v H$	(293–440)		54.1	308	A	[1987STE/MAL, 1970DYK, 1999DYK/SVO]
C₂H₆O₂	[107-21-1]		ethylene glycol					
		$\Delta_v H$	(307–384)		62.4 ± 4.0	345		[2010PET/REY]
		$\Delta_v H$	(308–336)		65.4 ± 0.3	298	GS	[2005VAS/VER]
		$\Delta_v H$	(278–328)		66.0 ± 0.2	298	GS	[2004VER2]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		Method	Reference
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)		
	$\Delta_v H$	(260–350)	66.2	298	[2003DER/MIC, 2004VER2]
	$\Delta_v H$	(264–353)	63.4 ± 0.1	298	[1999MOK/PRC, 2004VER2]
	$\Delta_v H$		65.6 ± 0.3	298	C [1988KNA/SAB, 1990KNA/SAB2]
	$\Delta_v H$	(363–408)	57.4	385	TGA [1987ALN/ALS]
	$\Delta_v H$	(323–473)	65.2	338	A [1987STE/MAL]
	$\Delta_v H$	(363–418)	62.5	378	A [1987STE/MAL]
	$\Delta_v H$	(283–373)	66.1 ± 0.3	298	GS [1981HAL/COG, 2004VER2]
	$\Delta_v H$	(335–420)	68.2 ± 0.8	298	EB [1981JOO/ARL, 2004VER2]
	$\Delta_v H$	(374–495)	63.7 ± 0.1	298	EB [1981AMB/HAL, 2005VAS/VER]
	$\Delta_v H$	(333–443)	63.6	298	[1972GAR/HUS, 2005VAS/VER]
	$\Delta_v H$	(323–473)	64.0	338	[1952JON/TAM, 1972GAR/HUS]
	$\Delta_v H$	(333–443)	63.8	298	DFM [1937GAL/HIB, 2004VER2, 1972GAR/HUS]
	$\Delta_v H$	(363–403)	61.1	383	[1935SCH/STA]
	$\Delta_v H$	(403–470)	57.3	436	[1935SCH/STA]
	$\Delta_v H$	(395–459)	61.1	410	[1901DEF]
C₂H₆O₂	[3031-74-1]	ethyl hydroperoxide			
	$\Delta_v H$	(253–363)	64.0	268	A [1987STE/MAL, 1951EGE/EMT, 1970DYK]
C₂H₆O₂	[na]	dihydroxyethane			
	$\Delta_{\text{fus}} H$		9.96	260.6	[1996DOM/HEA]
C₂H₆O₂S	[67-71-0]	dimethyl sulfone			
	$\Delta_{\text{fus}} H$		18.28	382	[1996DOM/HEA]
	$\Delta_{\text{sub}} H$		77 ± 2.9		[1970COX/PIL, UR/MAC]
	$\Delta_v H$	(387–523)	56.0	404	A [1987STE/MAL, 1970DYK, 1999DYK/SVO]
C₂H₆O₄	[17088-73-2]	<i>bis</i> -hydroxymethyl peroxide			
	$\Delta_{\text{sub}} H$		94.1 ± 4.2		ME [1953JEN/STY, 1970COX/PIL]
C₂H₆O₄S	[77-78-1]	dimethyl sulfate			
	$\Delta_v H$	(340–470)	46.7	355	A [1987STE/MAL, 1999DYK/SVO]
C₂H₆S	[75-18-3]	dimethyl sulfide			
	$\Delta_{\text{fus}} H$		7.98	174.9	[1996DOM/HEA]
	$\Delta_v H$		28.5 ± 0.1		[1997BAE]
	$\Delta_v H$		27.9 ± 0.6	298	C [1989VOR/KLY]
	$\Delta_v H$	(268–319)	28.9	283	A [1987STE/MAL]
	$\Delta_v H$	(307–379)	27.7	322	A [1987STE/MAL]
	$\Delta_v H$	(372–453)	26.6	387	A [1987STE/MAL]
	$\Delta_v H$	(447–503)	26.7	462	A [1987STE/MAL]
	$\Delta_v H$		27.5	298	[1981SHI/SAI]
	$\Delta_v H$		27.7	298	[1971WIL/ZWO]
	$\Delta_v H$		28.8 ± 0.1	276	C [1957MCC/HUB]
	$\Delta_v H$		27.9 ± 0.1	292	C [1957MCC/HUB]
	$\Delta_v H$		27.0 ± 0.1	310	C [1957MCC/HUB]
	$\Delta_v H$	(287–318)	28.2	302	EB [1952WHI/BER]
	$\Delta_v H$	(251–293)	28.9	278	[1942OSB/DOE]
	$\Delta_v H$		28.9	310	[1935THO/LIN]
C₂H₆S	[75-08-1]	ethyl mercaptan (ethanethiol)			
	$\Delta_{\text{fus}} H$		4.97	195.3	[1996DOM/HEA]
	$\Delta_v H$	(273–313)	28.4	288	A [1987STE/MAL]
	$\Delta_v H$	(303–375)	27.5	318	A [1987STE/MAL]
	$\Delta_v H$	(265–448)	26.3	380	A [1987STE/MAL]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(442–499)	26.6	457	A	[1987STE/MAL]
	$\Delta_v H$		27.3	298		[1971WIL/ZWO]
	$\Delta_v H$	(273–339)	28.4	288		[1966OSB/DOU]
	$\Delta_v H$	(273–339)	28.4	288	A, EB	[1987STE/MAL, 1952MCC/SCO, 1966OSB/DOU]
	$\Delta_v H$		28.7	306		[1935THO/LIN]
C ₂ H ₆ S ₂	[540-63-6]	1,2-ethanedithiol				
	$\Delta_v H$		44.7	298		[1962MAN/SUN]
C ₂ H ₆ S ₂	[624-92-0]	dimethyl disulfide				
	$\Delta_{\text{fus}} H$		9.19	188.4		[1996DOM/HEA]
	$\Delta_v H$		38.5 ± 0.6	298	C	[1989VOR/KLY]
	$\Delta_v H$	(297–402)	37.8	312	A	[1987STE/MAL]
	$\Delta_v H$		37.8 ± 0.1	298	C	[1985KUS]
	$\Delta_v H$		37.8	298		[1981SHI/SAI]
	$\Delta_v H$		38.4	298		[1971WIL/ZWO]
	$\Delta_v H$		36.0 ± 0.1	341	C	[1958HUB/DOU]
	$\Delta_v H$		34.9 ± 0.1	360	C	[1958HUB/DOU]
	$\Delta_v H$		33.7 ± 0.1	383	C	[1958HUB/DOU]
	$\Delta_v H$	(321–388)	36.7	336	EB	[1952WHI/BER]
	$\Delta_v H$	(334–401)	36.2	349		[1950SCO/FIN]
	$\Delta_v H$	(288–333)	38.2	303		[1950SCO/FIN]
C ₂ H ₇ N	[124-40-3]	dimethylamine				
	$\Delta_{\text{fus}} H$		5.94	181		[1996DOM/HEA]
	$\Delta_v H$	(277–360)	27.0	292	A	[1987STE/MAL]
	$\Delta_v H$	(358–438)	23.8	373	A	[1987STE/MAL]
	$\Delta_v H$	(202–279)	28.4	264	A	[1987STE/MAL, 1939AST/EID, 1984BOU/FRI]
C ₂ H ₇ N	[75-04-7]	ethyl amine				
	$\Delta_v H$	(213–297)	29.0	282	A	[1987STE/MAL]
	$\Delta_v H$	(290–449)	27.2	305	A	[1987STE/MAL]
	$\Delta_v H$	(291–387)	27.6	306	A	[1987STE/MAL]
	$\Delta_v H$	(377–456)	25.9	392	A	[1987STE/MAL]
	$\Delta_v H$	(275–288)	29.1	281		[1962BIT/KAU2]
	$\Delta_v H$	(190–290)	28.9	275		[1947STU]
C ₂ H ₇ NO	[5725-96-2]	N,N-dimethylhydroxyl amine				
	$\Delta_v H$	(290–363)	45.7	305	A	[1987STE/MAL, 1957BIS/PAR, 1984BOU/FRI]
C ₂ H ₇ NO	[1117-97-1]	N,O-dimethylhydroxyl amine				
	$\Delta_v H$	(228–316)	34.3	243	A	[1987STE/MAL, 1957BIS/PAR, 1984BOU/FRI]
C ₂ H ₇ NO	[141-43-5]	2-aminoethanol				
	$\Delta_v H$	(283–363)	60.6	298		[2009BEL/RAZ]
	$\Delta_v H$	(357–435)	55.9	372	EB	[2008KIM/SVE]
	$\Delta_v H$	(279–324)	59.6 ± 0.3	298	GS	[2005KAP/SLO]
	$\Delta_v H$	(358–440)	59.4	298	EB	[1999TOC/AKI, 2005KAP/SLO]
	$\Delta_v H$	(352–613)	59.0	298	EB	[1987DAU/JAL, 2005KAP/SLO]
	$\Delta_v H$	(310–444)	61.7	325	A	[1987STE/MAL]
	$\Delta_v H$	(298–308)	56.1	298		[1982TOU/OKA, 2005KAP/SLO]
	$\Delta_v H$	(293–298)	54.8	298		[1974GUS/REN, 2005KAP/SLO]
	$\Delta_v H$	(325–443)	57.7	298		[1969DAN/MAT, 2005KAP/SLO]
	$\Delta_v H$	(379–443)	54.7	394		[1959MCD/SHR]
	$\Delta_v H$	(273–301)	U 50.8	287	A, GS	[1957BES/KOC]
	$\Delta_v H$	(338–443)	58.9	353		[1950MAT/SUM, 1984BOU/FRI]
	$\Delta_v H$	(303–373)	57.4	298	EB	[1947LEI/SHO, 2005KAP/SLO]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	Δ_vH	(341–453)	57.9	298		[1935WIL, 2005KAP/SLO]
C ₂ H ₇ O ₃ P	[868-85-9]	dimethyl phosphite				
	Δ_vH	(353–443)	46.7	368		[2010FAN/WAN]
	Δ_vH	(243–443)	48.3	243	GS	[2009BUT/BUC]
	Δ_vH	(243–443)	47.9	263	GS	[2009BUT/BUC]
	Δ_vH	(243–443)	46.7	298	GS	[2009BUT/BUC]
	Δ_vH	(243–443)	46.4	313	GS	[2009BUT/BUC]
	Δ_vH	(243–443)	46.0	333	GS	[2009BUT/BUC]
	Δ_vH	(243–443)	45.4	373	GS	[2009BUT/BUC]
	Δ_vH		39.5			[1993OVC/SOB]
	Δ_vH	(346–456)	38.7	361	A	[1987STE/MAL, 1970DYK]
C ₂ H ₇ O ₃ P	[15845-66-6]	ethylphosphonic acid				
	$\Delta_{\text{sub}}H$		50.6 ± 4.2			[1955NEA/WIL, 1970COX/PIL]
C ₂ H ₈ CIN	[506-59-2]	dimethylammonium chloride				
	Δ_vH	(429–533)	95.6	444	A	[1987STE/MAL]
	Δ_vH	(533–569)	143.9	548	A	[1987STE/MAL]
C ₂ H ₈ CIN	[557-66-4]	ethylammonium chloride				
	Δ_vH	(382–480)	34.3	397	A	[1987STE/MAL]
C ₂ H ₈ NOPS ₂	[10265-92-6]	O,S-dimethyl phosphoroamidothioate				
	$\Delta_{\text{fus}}H$		13.34	316.8	DSC	[1990DON/DRE]
C ₂ H ₈ N ₂	[57-14-7]	1,1-dimethylhydrazine				
	$\Delta_{\text{fus}}H$		10.07	216		[1996DOM/HEA]
	Δ_vH	(267–303)	34.1	284		[2000BOU/YE]
	Δ_vH	(238–292)	36.5	277	A	[1987STE/MAL, 1953AST/WOO, 1984BOU/FRI]
C ₂ H ₈ N ₂	[540-73-8]	1,2-dimethylhydrazine				
	$\Delta_{\text{fus}}H$		13.64	264.3		[1996DOM/HEA]
	Δ_vH	(274–297)	41.0	286	A	[1987STE/MAL, 1951AST/JAN, 1984BOU/FRI]
C ₂ H ₈ N ₂	[107-15-3]	ethylenediamine				
	$\Delta_{\text{fus}}H$		21.08	284.1	DSC	[1997LEE/CHA]
	$\Delta_{\text{trs}}H$		0.49	189		
	$\Delta_{\text{fus}}H$		22.58	284.3	AC	[1991ACR, 1994LEE/LIE, 1975MES/FIN]
	$\Delta_{\text{sub}}H$	(242–278)	65.6	263	IP	[1987STE/MAL, 1975MES/FIN]
	Δ_vH	(294–325)	41.6	298	TGA	[1988AFZ/BUT, 2010EFI/EME]
	Δ_vH	(303–391)	43.9	318	A	[1987STE/MAL]
	Δ_vH	(284–419)	45.9	299	A, IP	[1987STE/MAL, 1975MES/FIN]
	Δ_vH		45.0 ± 0.1	298	C	[1969WAD]
	Δ_vH	(314–388)	54.4 ± 1.0	298		[1967SIV/MAT, 2010EFI/EME]
	Δ_vH		46.0 ± 0.2	298	IP	[1965DOU/OSB, 1970GOO/MOR]
	Δ_vH	(299–390)	45.6	314		[1934HIE/WOE, 1984BOU/FRI]
C ₂ H ₈ N ₂	[na]	diaminoethane				
	$\Delta_{\text{trs}}H$		0.49	189		
	$\Delta_{\text{fus}}H$		22.58	284.2		[1991ACR, 1994LEE/LIE]
C ₂ H ₈ N ₆ O ₂	[216489-95-1]	1,1'-(1,2-ethanediyl)bis(1-nitrosohydrazine)				
	$\Delta_{\text{sub}}H$		172.4 ± 1.3	298		[1998LEB/CHI]
C ₃ BrClF ₆ O ₄	[38126-26-0]	perchloric acid, 1,1,2,3,3,3-hexafluoro-2-bromopropyl ester				
	Δ_vH	(273–293)	38.1	283	A	[1987STE/MAL, 1973SCH/PIL]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₃ BrF ₅ O	[6129-62-0] $\Delta_v H$	2-bromo-2,3,3,3-tetrafluoropropionyl fluoride (224–282)	30.2	267	A	[1987STE/MAL]
C ₃ BrF ₆ NO	[na] $\Delta_v H$	N,N-bis(trifluoromethyl) carbamoyl bromide (233–293)	30.7	278	A	[1987STE/MAL]
C ₃ BrF ₉ N ₂	[na] $\Delta_v H$	N-bromo-tris(trifluoromethyl)hydrazine (283–333)	36.8	308		[1966BRO/FRA]
C ₃ BrF ₁₀ NS	[62977-73-5] $\Delta_v H$	bromotrifluoro[1,1,1,2,3,3,3-heptafluoro-2-propanaminoato(2-)] sulfur	35.1	394	I	[1977KIT/SHR2]
C ₃ Br ₂ F ₆ O	[2356-57-2] $\Delta_v H$	(trifluoromethyl)(1,2-dibromo-1,2,2-trifluoroethyl) ether (299–335)	34.6	314	A	[1987STE/MAL, 1968HAS/TIP]
C ₃ Br ₃ F ₆ NO	[29528-78-7] $\Delta_v H$	1,1,1,1',1',1'-hexafluoro-N-(tribromomethoxy)dimethylamine (297–338)	28.9	312	A	[1987STE/MAL]
C ₃ ClF ₄ NO ₂	[42016-32-0] $\Delta_v H$	chloro(trifluoroacetyl)carbamic fluoride	39.3	371		[1973SPR/WRI]
C ₃ ClF ₅ O	[79-53-8] $\Delta_v H$	chloropentafluoroacetone (232–303)	27.3	247	A	[1987STE/MAL, 1964MUR, 1984BOU/FRI]
C ₃ ClF ₅ O	[28627-00-1] $\Delta_v H$	2-chloro-2,3,3,3-tetrafluoropropionyl fluoride (195–273)	23.9	258	A	[1987STE/MAL]
C ₃ ClF ₆ NO ₂	[15496-01-2] $\Delta_v H$	O-(chloroformyl)-N,N-bis(trifluoromethyl)hydroxylamine (227–286)	34.5	271	A	[1987STE/MAL]
C ₃ ClF ₆ NS	[na] $\Delta_v H$	chloro(hexafluoroisopropylideneimino) sulfur	37.7	368	I	[1972MET/SHR]
C ₃ ClF ₇ O	[22675-68-9] $\Delta_v H$ $\Delta_v H$	heptafluoroisopropyl hypochlorite (196–287) (194–273)	26.7 22.7	272 258	A A	[1987STE/MAL] [1987STE/MAL]
C ₃ ClF ₈ N	[33757-13-0] $\Delta_v H$	N-chloro-N-1,2,2,2-pentafluoro-1-(trifluoromethyl)ethylamine (240–311)	28.8	255	A	[1987STE/MAL, 1971SWI/ZAB]
C ₃ ClF ₈ NOS	[74366-14-6] $\Delta_v H$	(heptafluoropropyl)imidisulfuryl chloride fluoride	26.7	346	I	[1980ABE/SHR]
C ₃ ClF ₁₀ NS	[62977-71-3] $\Delta_v H$	chlorotrifluoro[1,1,1,2,3,3,3-heptafluoro-2-propanaminoato(2-)] sulfur	33.5	391	I	[1977KIT/SHR2]
C ₃ Cl ₂ F ₅ N	[na] $\Delta_v H$	2,2-difluoro-1,2-dichloro-N-(trifluoromethyl)ethylideneimine (283–318)	31.2	298	A	[1987STE/MAL]
C ₃ Cl ₂ F ₆	[661-97-2] $\Delta_v H$ $\Delta_v H$ $\Delta_v H$	1,2-dichlorohexafluoropropane (296–307)	28.1 26.9 ± 0.1 25.9 ± 0.1	301 298 313	 C C	 [1980MAJ/SVO] [1980MAJ/SVO] [1980MAJ/SVO]
C ₃ Cl ₂ F ₆ N ₂	[na] $\Delta_v H$	bis(trifluoromethyl)aminocarbonylamine chloride (267–339)	35.0	303		[1966DOB/EME]
C ₃ Cl ₂ F ₆ O	[22675-69-0] $\Delta_v H$	hypochlorous acid, 2-chloro-1,1,2,3,3,3-hexafluoropropyl ester (273–293)	29.6	283	A	[1987STE/MAL]
C ₃ Cl ₂ F ₇ N	[32751-04-5] $\Delta_v H$	N,N-dichloro-1,2,2,2-tetrafluoro-1-(trifluoromethyl)ethylamine (299–344)	32.7	314	A	[1987STE/MAL, 1971SWI/ZAB]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound					
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference	
C ₃ Cl ₂ F ₇ NS	[26454-66-0] $\Delta_v H$	S,S-dichloro-N-[tetrafluoro-1-(trifluoromethyl)ethyl]sulfilimine (313–347)	39.3	328	A	[1987STE/MAL, 1970VON/GLE]	
C ₃ Cl ₂ F ₇ NS	[na] $\Delta_v H$	C ₃ F ₇ N=SCl ₂	25.9	361	I	[1980ABE/SHR]	
C ₃ Cl ₂ F ₇ P	[662-55-5] $\Delta_v H$	dichloro(heptafluoropropyl)phosphine (273–348)	33.5	310		[1959EME/SMI]	
C ₃ Cl ₃ F ₅ O	[37136-24-6] $\Delta_v H$	chlorodifluoromethyl 2,2-dichloro-1,1,2-trifluoroethyl ether (302–350)	33.4	317	A	[1987STE/MAL]	
	$\Delta_v H$		33.8 ± 0.5	298	EB	[1976AMM/BUL]	
C ₃ Cl ₅ F ₃ O	[428-73-9] $\Delta_v H$	trichloromethyl 2,2-dichloro-1,1,2-trifluoroethyl ether (341–423)	42.2	356	A	[1987STE/MAL]	
	$\Delta_v H$		45.7 ± 0.7	298	EB	[1976AMM/BUL]	
C ₃ Cl ₆	[1888-71-7] $\Delta_v H$	hexachloropropylene (366–510)	54.8 ± 0.4	298	EB	[1997STE/CHI4]	
	$\Delta_v H$		(382–540)	49.3	397	A	[1987STE/MAL, 1970DYK]
C ₃ Cl ₆	[2065-35-2] $\Delta_{\text{fus}} H$	hexachlorocyclopropane	18.6	376		[1996DOM/HEA]	
C ₃ F ₃ N ₂ P	[58310-46-6] $\Delta_v H$	dicyano(trifluoromethyl)phosphine (291–334)	45.6	306	A	[1987STE/MAL]	
C ₃ F ₃ N ₃	[675-14-9] $\Delta_v H$	2,4,6-trifluoro-1,3,5-triazine (277–344)	38.8	292	A	[1987STE/MAL]	
C ₃ F ₄	[20174-11-2] $\Delta_v H$	tetrafluoropropyne (179–218)	18.8	203	A	[1987STE/MAL]	
C ₃ F ₄ O ₂ S ₂	[58936-60-0] $\Delta_v H$	ethane(dithioperoxoic)acid, fluoro-oxo-trifluoromethyl ester	34.9	385	I	[1976BUR/SHR]	
C ₃ F ₅ N	[3291-42-7] $\Delta_v H$	2,2-difluoro-3-(trifluoromethyl)-2H-azirine (193–249)	24.0	234	A	[1987STE/MAL]	
C ₃ F ₅ N	[3291-41-6] $\Delta_v H$	2,3-difluoro-2-(trifluoromethyl)-2H-azirine (193–249)	24.3	220	A	[1987STE/MAL]	
C ₃ F ₆	[116-15-4] $\Delta_v H$	perfluoropropene (233–293)	21.9	278	A	[1987STE/MAL, 1952WHI, 1970DYK, 1984BOU/FRI]	
C ₃ F ₉ N ₂ OS	[34556-28-0] $\Delta_v H$	N-cyano-S,S-bis(trifluoromethyl)sulfoximine	30.8	382	I	[1972SAU/SHR]	
C ₃ F ₆ O	[425-82-1] $\Delta_v H$	hexafluoroacetone (232–313)	22.3	247	A	[1992SAL/WAN]	
C ₃ F ₆ O	[116-16-5] $\Delta_{\text{fus}} H$	perfluoroacetone	8.38	147.7		[1996DOM/HEA]	
	$\Delta_v H$		(195–246)	23.6	231	A	[1987STE/MAL, 1967PLA/PAC, 1984BOU/FRI]
	$\Delta_v H$		(240–357)	22.3	253		[1964MUR, 1984BOU/FRI]
	$\Delta_v H$		(213–245)	23.1	229		[1955MOR/AYS]
C ₃ F ₆ O	[1187-93-5] $\Delta_v H$	trifluoromethyl trifluorovinyl ether (248–338)	20.3	298		[2007LIU/LIU]	
	$\Delta_v H$		(208–241)	22.9	226	A	[1987STE/MAL]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₃ F ₆ O	[5930-63-2] $\Delta_v H$	pentafluoropropionyl hypofluorite (214–248)	25.8	233	A	[1987STE/MAL]
C ₃ F ₆ O ₂	[21297-65-4] $\Delta_v H$	1,3-perfluorodioxolane (234–367)	22.3	249	A	[1992SAL/WAN]
C ₃ F ₆ O ₄ S	[51689-98-6] $\Delta_v H$	pentafluoropropionic fluorosulfuric acid anhydride (252–335)	39.2	267	A	[1987STE/MAL, 1966DES/CAD]
C ₃ F ₆ O ₇ S ₂	[6378-48-9] $\Delta_v H$	hydroacrylic acid, tetrafluoroanhydride with fluorosulfuric acid, fluorosulfate (308–403)	49.4	323		[1999DYK/SVO]
C ₃ F ₇ I ₂ P	[678-07-9] $\Delta_v H$	diiodo(heptafluoropropyl)phosphine (313–393)	39.6	353		[1959EME/SMI]
C ₃ F ₇ N	[428-71-7] $\Delta_v H$	perfluoro-(2-ethyl-1,2-oxazetidine) 25.6				[1961BAR/HAS]
C ₃ F ₇ N	[760-43-0] $\Delta_v H$	perfluoro(ethylidene)methylamine 21.4				[1961BAR/HAS]
C ₃ F ₇ NO	[32822-50-7] $\Delta_v H$	heptafluoropropionamide 27.2	279		HG	[1971DEM/SHR]
C ₃ F ₇ NOS	[26454-67-1] $\Delta_v H$	1,1,1,2,3,3,3-heptafluoro-N-sulfinyl-2-propanamine (252–280)	34.1	266	A	[1987STE/MAL, 1999DYK/SVO, 1970VON/GLE]
C ₃ F ₇ NOS	[74366-13-5] $\Delta_v H$	1,1,2,2,3,3,3-heptafluoro-N-sulfinyl-1-propanamine 26.3	325			[1980ABE/SHR]
C ₃ F ₇ NS	[62067-06-5] $\Delta_v H$	2,2,2-trifluoro-N-[(trifluoromethyl)thio]ethanimidoyl fluoride 28.0	305		I	[1977BUR/SHR2]
C ₃ F ₇ NO ₂	[423-33-6] $\Delta_v H$	perfluoro-1-nitropropane (247–296)	28.5	281	A	[1987STE/MAL]
C ₃ F ₈	[76-19-7] $\Delta_{\text{fus}}H$ $\Delta_{\text{fus}}H$ $\Delta_v H$ $\Delta_v H$	perfluoropropane 3.56 0.48 (193–237) (213–259)	99.4 125.5 21.6 20.9		A	[1996DOM/HEA] [1987STE/MAL, 1967PAC/PLA] [1963BRO]
C ₃ F ₈ N ₂ O ₂	[32837-67-5] $\Delta_v H$	N-[(difluoroamino)carbonyl]oxy]-1,1,1-trifluoro-N-(trifluoromethyl)-methanamine 31.4	310			[1973WRI/SHR]
C ₃ F ₈ OS	[33622-17-2] $\Delta_v H$	pentafluoroethyl trifluoromethyl sulfur 32.5				[1971SAU/SHR]
C ₃ F ₈ S	[33547-10-3] $\Delta_v H$	pentafluoroethyl trifluoromethyl sulfide 28.8				[1971SAU/SHR]
C ₃ F ₉ N	[432-03-1] $\Delta_v H$	perfluorotrimethylamine (193–263)	23.9	248	A	[1987STE/MAL]
C ₃ F ₉ NO	[671-63-6] $\Delta_v H$	1,1,1-trifluoro-N-(trifluoromethoxy)-N-(trifluoromethyl)-methanamine (226–268)	27.0	253	A	[1987STE/MAL]
C ₃ F ₉ NOS	[59617-29-7] $\Delta_v H$	[1,2,2,2-tetrafluoro-1-(trifluoromethyl)ethyl] imidosulfonyl fluoride 28.7				[1976STA/MEW]
C ₃ F ₉ NOS ₂	[34556-26-8] $\Delta_v H$	S,S-bis(trifluoromethyl)-N-[(trifluoromethyl)thio]sulfoximine 31.2	360		I	[1972SAU/SHR]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₃ F ₉ NO ₂ S ₂	[34556-27-9] $\Delta_v H$	S,S-bis(trifluoromethyl)-N-[(trifluoromethyl)sulfinyl]sulfoximine	37.2	388	I	[1972SAU/SHR]
C ₃ F ₉ NO ₂ S ₃	[29749-02-8] $\Delta_v H$	1,1,1-trifluoro-N,N-bis(trifluoromethyl)thio]methanesulfonamide (288–403)	43.5	303	A	[1987STE/MAL, 1999DYK/SVO]
C ₃ F ₉ N ₃ O	[10405-30-8] $\Delta_v H$ $\Delta_v H$	nitrosotris(trifluoromethyl) hydrazine (279–300)	29.5	289	A	[1987STE/MAL]
			33.5	263		[1966DOB/EME]
C ₃ F ₉ N ₃ O ₂	[10405-31-9] $\Delta_v H$	nitrotris(trifluoromethyl) hydrazine (293–321)	31.6	307	A	[1987STE/MAL]
C ₃ F ₉ P	[432-04-2] $\Delta_v H$	tris(trifluoromethyl)phosphine (248–285)	24.7	270	A	[1987STE/MAL]
C ₃ F ₉ PS	[671-64-7] $\Delta_v H$	bis(trifluoromethyl)trifluoromethylthiophosphine (242–293)	32.5	267		[1962EME/PAC]
C ₃ F ₉ PS	[2025-08-3] $\Delta_v H$	tris(trifluoromethyl)phosphine sulfide (282–308)	29.1	295		[1964CAV/EME2]
C ₃ F ₉ PS ₂	[36121-49-0] $\Delta_v H$	(trifluoromethyl)dithiophosphite acid, bis(trifluoromethyl) ester (273–296)	37.9	284	A	[1987STE/MAL, 1999DYK/SVO]
C ₃ F ₉ P ₃ S ₅	[26349-17-7] $\Delta_{\text{sub}} H$	2,4,5-tris(trifluoromethyl)-1,3,2,4,5-dithiatriphospholane-2,4,5-trisulfide (363–373)	96.6	368		[1970BUR/PARC]
C ₃ F ₁₀ OS	[33564-24-8] $\Delta_v H$	difluoro(oxo(trifluoromethyl)(pentafluoroethyl) sulfur (291–324)	30.6	306	A	[1987STE/MAL, 1999DYK/SVO]
C ₃ F ₁₀ O ₃ S	[60672-62-0] $\Delta_v H$	pentafluoro (pentafluoropropaneperoxoato)sulfur	34.4			[1976HOP/DES]
C ₃ F ₁₀ S	[68010-33-3] $\Delta_v H$	[2,2-difluoro-(1-trifluoromethyl)ethenyl] pentafluoro sulfur	30.0			[1978DEM/FOX]
C ₃ F ₁₀ S	[31222-06-7] $\Delta_v H$	difluoro(pentafluoroethyl)(trifluoromethyl) sulfur	29.2			[1971SAU/SHR]
C ₃ F ₁₁ NO ₃ S ₂	[65844-08-8] $\Delta_v H$	trifluoro(trifluorosulfato-O)[1,1,1,2,3,3,3-heptafluoro-2-propanaminoato(2-)]sulfur	33.5	391	I	[1977KIT/SHR2]
C ₃ F ₁₂ O ₃ S ₂	[68010-30-0] $\Delta_v H$	pentafluoro [2,2,2-trifluoro-1-(fluorosulfonyl)oxo]-1-(trifluoromethyl)-ethyl] sulfur	37.2			[1978DEM/FOX]
C ₃ N ₃ P	[1116-01-4] $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$	tricyanophosphine (293–323)	78.3	308	ME	[1987STE/MAL, 1976DAV/FIN]
			75.3 ± 2.9	298		[1995ALM/FIN, 1976DAV/FIN]
C ₃ HCIF ₆ O ₂ S	[57169-81-0] $\Delta_v H$	chlorosulfurous acid, 2,2,2-trifluoro-1-(trifluoromethyl)ethyl ester	36.7			[1975KIR/LAS]
C ₃ HCIF ₁₀ S	[68010-36-6] $\Delta_v H$	[1-(chlorodifluoromethyl)-2,2,2-trifluoroethyl] pentafluorosulfur	31.2			[1978DEM/FOX]
C ₃ HCl ₇	[594-89-8] $\Delta_v H$	1,1,1,2,2,3,3-heptachloropropane (413–473)	34.8	428	A	[1987STE/MAL, 1949HIG/END, 1970DYK]
C ₃ HF ₃	[661-54-1] $\Delta_v H$	3,3,3-trifluoropropyne (138–213)	21.5	198	A	[1987STE/MAL]
C ₃ HF ₆ N	[3291-64-3]	2,2,3-trifluoro-3-fluoromethylaziridine				

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(268–298)	30.2	283	A	[1987STE/MAL]
C ₃ HF ₇	[431-89-0]	1,1,1,2,3,3,3-heptafluoropropane				
	$\Delta_v H$		18.9	298		[2007ECK/HUA]
	$\Delta_v H$	(278–308)	22.7	293		[2002BOB/ART]
	$\Delta_v H$	(293–353)	22.6	308		[2002VAL/COQ]
	$\Delta_v H$	(237–370)	22.3	250		[1992SAL/WAN]
	$\Delta_v H$	(237–370)	14.5	300		[1992SAL/WAN]
	$\Delta_v H$	(237–370)	12.5	325		[1992SAL/WAN]
C ₃ HF ₇ O	[2356-61-8]	trifluoromethyl 1 <i>H</i> -pentafluoroethyl ether				
	$\Delta_v H$	(232–313)	27.3	247	A	[1992SAL/WAN]
C ₃ HF ₇ O ₂ S	[52225-56-6]	fluorosulfurous acid, 2,2,2-trifluoro-1-(trifluoromethyl)ethyl ester				
	$\Delta_v H$		33.8			[1975DEM/KOV2]
C ₃ HF ₈ NOS	[34556-23-5]	S-(pentafluoroethyl)-S-(trifluoromethyl)sulfoximine				
	$\Delta_v H$		36.3	358	I	[1972SAU/SHR]
C ₃ HF ₉ N ₂	[13105-67-4]	<i>tris</i> (trifluoromethyl)hydrazine				
	$\Delta_v H$	(238–307)	29.9	273		[1966DOB/EME]
C ₃ HF ₁₁ S	[68010-34-4]	pentafluoro [2,2,2-trifluoro-1-(1-trifluoromethyl)ethyl]sulfur				
	$\Delta_v H$		30.1			[1978DEM/FOX]
C ₃ HN	[68010-34-4]	cyanoacetylene				
	$\Delta_{\text{fus}} H$		14.1		Sub-Vap	[1963DAN/FLU]
	$\Delta_{\text{sub}} H$	(247–279)	42.3	264		[1987STE/MAL, 1963DAN/FLU]
	$\Delta_v H$	(279–315)	28.1	294	A	[1987STE/MAL, 1963DAN/FLU]
C ₃ H ₂ ClF ₅	[460-92-4]	3-chloro-1,1,1,3,3-pentafluoropropane				
	$\Delta_{\text{fus}} H$		10.47	165.4		[1996DOM/HEA]
C ₃ H ₂ ClF ₅ O	[13838-16-9]	1-chloro-1,2,2-trifluoro-2-(difluoromethoxy)ethane				
	$\Delta_v H$	(274–351)	33.8	289		[1988AMB/GHI2]
	$\Delta_v H$	(290–329)	32.9	305	A	[1987STE/MAL]
	$\Delta_v H$		32.6 ± 0.1	298	C	[1984UCH/MAJ]
	$\Delta_v H$		31.3 ± 0.1	313	C	[1984UCH/MAJ]
	$\Delta_v H$		30.2 ± 0.1	328	C	[1984UCH/MAJ]
C ₃ H ₂ ClF ₅ O	[26675-46-7]	2-chloro-1,1,1-trifluoro-2-(difluoromethoxy)ethane				
	$\Delta_v H$	(280–344)	31.7	295		[1988AMB/GHI2]
	$\Delta_v H$	(283–312)	31.9	297	A	[1987STE/MAL]
C ₃ H ₂ Cl ₂ F ₄	[64712-27-2]	3,3-dichloro-1,1,1,3-tetrafluoropropane				
	$\Delta_v H$	(297–333)	31.7	312	A	[1987STE/MAL]
C ₃ H ₂ Cl ₂ F ₄ O	[37031-38-2]	2-chloro-1,1,2-trifluoroethyl chlorofluoromethyl ether				
	$\Delta_v H$		37.5 ± 0.1	298	C	[1984MAJ/UCH]
	$\Delta_v H$		36.4 ± 0.1	313	C	[1984MAJ/UCH]
	$\Delta_v H$		35.3 ± 0.1	328	C	[1984MAJ/UCH]
	$\Delta_v H$		34.1 ± 0.1	343	C	[1984MAJ/UCH]
	$\Delta_v H$		32.9 ± 0.1	353	C	[1984MAJ/UCH]
C ₃ H ₂ Cl ₃ F ₃	[7125-84-0]	1,1,1-trichloro-3,3,3-trifluoropropane				
	$\Delta_{\text{fus}} H$		14.07	232.7		[1991ACR]
	$\Delta_v H$		36.8 ± 0.1	298	C	[2007VAR/DRU]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(320–365)	35.2	335		[1987STE/MAL]
C ₃ H ₂ Cl ₄	[60320-18-5]	1,1,2,3-tetrachloropropylene				
	$\Delta_v H$	(347–416)	42.9	362	A	[1987STE/MAL]
C ₃ H ₂ Cl ₂ F ₄	[64712-27-2]	1,1,1,3-tetrafluoro-3,3-dichloropropane				
	$\Delta_v H$	(297–333)	31.9 ± 0.5	298		[2007VAR/DRU]
C ₃ H ₂ D ₅ N	[153557-95-0]	1,1,1,3-tetrafluoro-3,3-dichloropropane				
	$\Delta_v H$	(283–330)	31.7	298		[1993WOL/KIM]
C ₃ H ₂ FNOS	[459-71-2]	fluoracetyl isothiocyanate				
	$\Delta_v H$	(273–353)	49.3	288	A	[1987STE/MAL, 1970DYK, 1999DYK/SVO]
C ₃ H ₂ F ₆	[690-39-1]	1,1,1,3,3,3-hexafluoropropane				
	$\Delta_v H$	(283–323)	24.5	303	A	[2000BOB/CAM]
C ₃ H ₂ F ₆ N ₂ S	[na]	amino (hexafluoroisopropylideneimino) sulfur				
	$\Delta_v H$		37.7	388	I	[1972MET/SHR]
C ₃ H ₂ F ₆ N ₂ S	[62067-09-8]	2,2,2-trifluoro-N-[(trifluoromethyl)thio]ethanimidamide				
	$\Delta_v H$	(322–390)	39.8	337	A, I	[1987STE/MAL, 1977BUR/SHR2, 1999DYK/SVO]
C ₃ H ₂ F ₆ O	[57041-67-5]	2-(difluoromethoxy)-1,1,1,2-tetrafluoroethane				
	$\Delta_v H$	(274–311)	24.9	293	I	[1996SUS/SMI]
C ₃ H ₂ F ₆ O	[920-66-1]	1,1,1,3,3,3-hexafluoro-2-propanol				
	$\Delta_v H$	(294–330)	40.2	309	A, MM	[1987STE/MAL, 1973ROC/SYM]
	$\Delta_v H$	(294–330)	41.6	298	MM	[1973ROC/SYM]
	$\Delta_v H$	(273–296)	47.3	284		[1967VAN/SOC]
C ₃ H ₂ F ₆ O ₂	[30957-44-9]	bis-(difluoromethoxy)difluoromethane				
	$\Delta_{\text{fus}} H$		7.2	153		[1999MAR/BAS]
	$\Delta_v H$	(243–308)	31.5 ± 0.4			[1999MAR/BAS]
C ₃ H ₂ F ₆ O ₂ S	[30957-44-9]	trifluoromethanesulfinic acid, 2,2,2-trifluoromethyl ester				
	$\Delta_v H$		36.8	363		[1971SAU/SHR2]
C ₃ H ₂ F ₈ N ₂ S	[2433-66-1]	S,S-difluoro-N-[1-amino-2,2,2-trifluoro-1-(trifluoromethyl)ethyl]sulfilimine				
	$\Delta_v H$	(295–313)	38.7	304	A	[1987STE/MAL, 1969GLE/VON, 1999DYK/SVO]
C ₃ H ₂ N ₂	[109-77-3]	malononitrile				
	$\Delta_{\text{trs}} H$		0.43	260.9		
	$\Delta_{\text{fus}} H$		10.7	305	DSC	[2007BAD/BLA]
	$\Delta_{\text{fus}} H$		10.8	305		[1996DOM/HEA]
	$\Delta_{\text{sub}} H$	(278–299)	78.2 ± 1.0	298		[1990BEC/DOG]
	$\Delta_{\text{sub}} H$		79.1 ± 8		ME	[1967BOY/GUH, 1970COX/PIL]
C ₃ H ₂ N ₂ O ₃	[120-89-8]	imidazolidine-2,4,5-trione				
	$\Delta_{\text{trs}} H$		2.1	392.3		
	$\Delta_{\text{fus}} H$		NA			[2008RIB/RIB2]
	$\Delta_{\text{sub}} H$	(381–406)	114.6 ± 0.6	393.7	ME	[2008RIB/RIB2]
	$\Delta_{\text{sub}} H$	(381–406)	119.4 ± 0.6	298	ME	[2008RIB/RIB2]
C ₃ H ₂ OS ₂	[2314-40-1]	1,3-dithiol-2-one				
	$\Delta_{\text{sub}} H$		73.6 ± 0.8	298		[1973RAU/GEI, 1977PED/RYL]
C ₃ H ₂ OS ₃	[930-35-8]	1,3-dithiole-2-thione				
	$\Delta_{\text{sub}} H$		75.4 ± 0.4	298		[1973RAU/GEI, [1977PED/RYL]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₃ H ₂ O ₃	[872-36-6]	vinylene carbonate				
	$\Delta_v H$	(308–350)	46.9	323	A	[1987STE/MAL]
	$\Delta_v H$	(308–400)	41.3		MM	[1971CHO/JON]
C ₃ H ₃ Cl	[7747-84-4]	1-chloro-1-propyne				
	$\Delta_v H$	(200–289)	28.3	274	A	[1987STE/MAL]
C ₃ H ₃ Cl ₂ F ₃	[460-69-5]	1,1-dichloro-3,3,3-trifluoropropane				
	$\Delta_{\text{trs}} H$		0.2	167.7		
	$\Delta_{\text{fus}} H$		10.13	182.2		[1996DOM/HEA]
	$\Delta_v H$		34.1 ± 0.1	298	C	[2007VAR/DRU]
	$\Delta_v H$	(301–342)	33.7	316	A	[1987STE/MAL]
C ₃ H ₃ Cl ₂ F ₃ O	[428-92-2]	2-chloro-1,1,2-trifluoroethyl chloromethyl ether				
	$\Delta_v H$		42.4 ± 0.1	298	C	[1984UCH/MAJ]
	$\Delta_v H$		41.2 ± 0.1	313	C	[1984UCH/MAJ]
	$\Delta_v H$		40.1 ± 0.1	328	C	[1984UCH/MAJ]
	$\Delta_v H$		39.0 ± 0.1	343	C	[1984UCH/MAJ]
	$\Delta_v H$		37.8 ± 0.1	358	C	[1984UCH/MAJ]
C ₃ H ₃ Cl ₃ O ₂	[598-99-2]	methyl trichloroacetate				
	$\Delta_v H$		48.3 ± 0.1	298	C	[1972LAY/WAD]
C ₃ H ₃ Cl ₅	[16714-68-4]	1,1,2,2,3-pentachloropropane				
	$\Delta_v H$	(365–447)	46.3	380	A	[1987STE/MAL, 1970DYK]
C ₃ H ₃ F ₃	[677-21-4]	3,3,3-trifluoro-1-propene				
	$\Delta_v H$	(283–363)	22.0	298	A	[1987STE/MAL]
C ₃ H ₃ F ₄ I	[1737-76-4]	1,1,1,2-tetrafluoro-3-iodopropane				
	$\Delta_v H$	(295–356)	28.4	310	A	[1987STE/MAL]
C ₃ H ₃ F ₄ I	[460-74-2]	1,1,1,3-tetrafluoro-3-iodopropane				
	$\Delta_v H$	(301–356)	31.2	316	A	[1987STE/MAL]
C ₃ H ₃ F ₄ NO ₂	[na]	methoxy (trifluoromethyl)carbamic fluoride				
	$\Delta_v H$		27.8			[1979SEK/DES]
C ₃ H ₃ F ₅	[679-86-7]	1,1,2,2,3-pentafluoropropane				
	$\Delta_v H$	(258–353)	30.2	273	A	[2002DIN/PAS]
C ₃ H ₃ F ₅	[1814-88-6]	1,1,1,2,2-pentafluoropropane				
	$\Delta_v H$	(232–283)	22.9	268	A	[1987STE/MAL, 1970DYK]
	$\Delta_v H$	(233–379)	23.0	248		[1967SHA]
C ₃ H ₃ F ₅ O	[422-05-9]	2,2,3,3,3-pentafluoro-1-propanol				
	$\Delta_v H$	(273–297)	47.0	285	A, MM	[1987STE/MAL, 1967MEE/GOL, 1984BOU/FRI]
	$\Delta_v H$		44.4	298	MM	[1973ROC/SYM, 1967MEE/GOL]
	$\Delta_v H$		41.3	298		[1967MUR/KIV]
C ₃ H ₃ F ₅ O	[37031-31-5]	1,1,2,2-tetrafluoro-1-(fluoromethoxy)ethane				
	$\Delta_v H$	(288–317)	33.5	303	I	[2002MUR/YAM]
C ₃ H ₃ F ₆ NOS	[34556-25-7]	N-methyl-S,S-bis(trifluoromethyl)sulfoximine				
	$\Delta_v H$		30.7	338	I	[1972SAU/SHR]
C ₃ H ₃ F ₆ NS	[13105-12-9]	N,N-bis(trifluoromethyl)methanesulfenamide				
	$\Delta_v H$	(269–309)	31.1	284	A,T	[1987STE/MAL, 1966EME/TAT]
C ₃ H ₃ F ₆ O ₂ P	[25439-11-6]	bis(trifluoromethyl)phosphinic acid, methyl ester				
	$\Delta_v H$	(258–313)	40.5	273	A	[1987STE/MAL]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₃ H ₃ F ₆ PS	[1486-18-6] Δ_vH	<i>bis</i> (trifluoromethyl) methylthiophosphine (273–321)	36.9	297	T	[1964CAV/EME]
C ₃ H ₃ F ₆ PS ₂	[18799-79-6] Δ_vH	<i>bis</i> (trifluoromethyl)dithiophosphinic acid, methyl ester (273–344)	41.5	288	A	[1987STE/MAL, 1999DYK/SVO]
C ₃ H ₃ N	[107-13-1] $\Delta_{\text{trs}}H$ $\Delta_{\text{fus}}H$ Δ_vH Δ_vH Δ_vH Δ_vH Δ_vH	acrylonitrile (257–352) (283–343) (222–351) (293–343) (273–353)	1.19 6.23 33.6 31.6 35.5 32.9 32.6	162.5 189.6 272 298 237 308	 A A	[1996DOM/HEA] [1987STE/MAL] [1987STE/MAL] [1964SEV/SOK] [1964GUB/FER] [1945DAV/WIE]
C ₃ H ₃ NO	[288-42-6] Δ_vH Δ_vH	oxazole (293–344)	34.6 32.5 ± 0.1	308 298	A C	[1987STE/MAL] [1978MCC/HAM]
C ₃ H ₃ NO	[288-14-2] Δ_vH Δ_vH	isoxazole (314–404)	37.2 ± 0.2 36.5 ± 0.1	298 298	EB C	[1996STE/CHI3] [1978MCC/HAM]
C ₃ H ₃ NO ₂	[17640-15-2] Δ_vH	cyanofornic acid, methyl ester (273–333)	39.3	288	A	[1987STE/MAL, 1970DYK]
C ₃ H ₃ NS	[288-47-1] $\Delta_{\text{fus}}H$ Δ_vH Δ_vH	thiazole (333–393) (336–391)	9.58 39.7 38.9	239.4 348 351	 A A	[1968GOU/WES2, 1966MEY/MET] [1987STE/MAL] [1987STE/MAL, 1969SOU/GOU2]
C ₃ H ₃ N ₃	[290-87-9] $\Delta_{\text{trs}}H$ $\Delta_{\text{fus}}H$ $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$ Δ_vH	1,3,5-triazine (212–229) (242–264) (342–373)	0.07 14.56 58.2 54.2 ± 0.2 56.5 ± 2.1 U 43.1 38.8 ± 1.9	197.7 353.4 222 298 253 298	 TE,ME ME CGC	[1991ACR] [1983DEW/VAN] [1982BYS] [1982INI/LOP] [1968MAS/RAE] [2009LIP/CHI2, 2009LIP/CHI]
C ₃ H ₃ N ₃ O ₂	[461-89-2] $\Delta_{\text{sub}}H$	6-azauracil 		141	LE	[1974YAN/VER]
C ₃ H ₃ N ₃ O ₃	[108-80-5] $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$	cyanuric acid (440–473)	131 133	458 298	ME,TE	[1983DEW/VAN] [1983DEW/VAN]
C ₃ H ₃ N ₅ O ₁₀	[62626-83-9] $\Delta_{\text{sub}}H$	1,1,1,2,2-pentanitropropane 	77.4 ± 1.3	298		[1999MIR/VOR]
C ₃ H ₄	[463-49-0] Δ_vH Δ_vH Δ_vH Δ_vH Δ_vH Δ_vH	allene (136–274) (193–246) (153–238) (138–189) (203–236) (200–260)	22.6 19.9 21.3 22.9 20.9 21.5	259 231 223 174 220 245	A A MM	[1987STE/MAL] [1987STE/MAL] [1947STU] [1940LAM/ROP, 1984BOU/FRI] [1930LIV/HEI] [1921MAA/WRI, 1984BOU/FRI]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₃ H ₄	[74-99-7]	1-propyne				
	$\Delta_v H$	(183–257)	23.0	242	A	[1987STE/MAL]
	$\Delta_v H$	(257–402)	20.8	272	A	[1987STE/MAL]
	$\Delta_v H$	(303–361)	21.2	318	A	[1987STE/MAL]
	$\Delta_v H$	(359–402)	21.9	374	A	[1987STE/MAL]
	$\Delta_v H$	(249–306)	23.2	264	A	[1987STE/MAL]
	$\Delta_v H$		22.1	275		[1971WIL/ZWO]
	$\Delta_v H$	(162–255)	23.9	240		[1967VAN]
	$\Delta_v H$	(323–400)	21.6	338		[1962VOH/KAN]
		(194–250)	23.4	235		[1933BOO/BUR, 1984BOU/FRI]
		(200–260)	21.4	230		[1921MAA/WRI]
C ₃ H ₄ Br ₂	[513-31-5]	2,3-dibromopropylene				
	$\Delta_v H$	(267–415)	43.1	282	A	[1987STE/MAL, 1947STU]
C ₃ H ₄ Br ₄	[54268-02-9]	1,2,2,3-tetrabromopropane				
	$\Delta_v H$	(418–580)	57.7	433	A	[1987STE/MAL, 1970DYK]
C ₃ H ₄ ClFO ₃	[462-27-1]	carbonochloridic acid, 2-fluoroethyl ester				
	$\Delta_v H$	(273–333)	46.6	288	GS	[1987STE/MAL, 1948RED/CHA4, 1970DYK]
C ₃ H ₄ ClF ₃	[460-35-5]	1-chloro-3,3,3-trifluoropropane				
	$\Delta_{\text{trs}} H$		4.49	169.8		
	$\Delta_{\text{fus}} H$		5.31	179.4		[1996DOM/HEA]
	$\Delta_v H$	(297–315)	29.9	306	A	[1987STE/MAL]
	$\Delta_v H$	(301–341)	33.7	316		[1972VAR/DRU]
C ₃ H ₄ ClF ₃ O	[425-87-6]	2-chloro-1,1,2-trifluoroethyl methyl ether				
	$\Delta_v H$		34.4 ± 0.1	298	C	[1984MAJ/UCH]
	$\Delta_v H$		33.4 ± 0.1	313	C	[1984MAJ/UCH]
	$\Delta_v H$		31.1 ± 0.1	343	C	[1984MAJ/UCH]
C ₃ H ₄ ClF ₃ O ₂ S	[61915-99-9]	trifluoromethanesulfinic acid, 2-chloroethyl ester				
	$\Delta_v H$	(320–403)	40.5	335	I	[1987STE/MAL, 1977BUR/SHR, 1999DYK/SVO]
C ₃ H ₄ Cl ₂ F ₂ O	[76-38-0]	2,2-dichloro-1,1-difluoro-1-methoxyethane				
	$\Delta_v H$	(279–378)	40.3	294	A	[1987STE/MAL]
C ₃ H ₄ Cl ₂ O	[513-88-2]	1,1-dichloroacetone				
	$\Delta_v H$	(292–382)	35.8	307	A	[1987STE/MAL, 1970SMI/THO]
C ₃ H ₄ Cl ₂ O	[534-07-6]	1,3-dichloroacetone				
	$\Delta_v H$	(348–445)	49.6	363	A	[1987STE/MAL, 1970SMI/THO]
C ₃ H ₄ Cl ₂ O ₂	[116-54-1]	methyl dichloroacetate				
	$\Delta_v H$		47.7 ± 0.1	298	C	[1972LAY/WAD]
	$\Delta_v H$	(331–481)	44.9	346	A	[1987STE/MAL, 1970DYK]
	$\Delta_v H$	(276–416)	47.2	291		[1947STU]
C ₃ H ₄ Cl ₄	[812-03-3]	1,1,1,2-tetrachloropropane				
	$\Delta_v H$	(331–469)	42.3	346	A	[1987STE/MAL, 1970DYK]
C ₃ H ₄ Cl ₄	[1070-78-6]	1,1,1,3-tetrachloropropane				
	$\Delta_{\text{trs}} H$		2.2	219.9		
	$\Delta_{\text{fus}} H$		10.49	237.7		[1996DOM/HEA]
	$\Delta_v H$	(300–377)	57.8	315	A	[1987STE/MAL]
C ₃ H ₄ Cl ₄	[13116-53-5]	1,2,2,3-tetrachloropropane				
	$\Delta_v H$	(346–415)	42.8	361	A	[1987STE/MAL]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₃ H ₄ F ₂ O ₂	[432-53-4] Δ_vH	methyl difluoroacetate (273–333)	41.9	288	GS	[1987STE/MAL, 1948RED/CHA4, 1970DYK]
C ₃ H ₄ F ₄ O	[32778-16-8] Δ_vH	2-difluoromethoxy-1,1-difluoroethane (288–328)	32.9	303	I	[2002MUR/YAM]
C ₃ H ₄ F ₄ O	[76-37-9] Δ_vH	2,2,3,3-tetrafluoro-1-propanol (303–380)	47.9	318	A	[1987STE/MAL]
	Δ_vH	(298–333)	50.3	313	MM	[1973ROC/SYM]
	Δ_vH	(298–333)	53.6	298	MM	[1973ROC/SYM]
C ₃ H ₄ N ₂	[288-32-4] $\Delta_{\text{fus}}H$	imidazole	12.8	361.9		[1996DOM/HEA, 1983DEW/DEK]
	$\Delta_{\text{sub}}H$	(292–309)	83.1 ± 0.2	300	ME	[1987JIM/ROU]
	$\Delta_{\text{sub}}H$		83.1 ± 0.2	298	ME	[1986JIM/ROU]
	$\Delta_{\text{sub}}H$	(288–310)	80.8	301	ME,TE	[1983DEW/VAN]
	$\Delta_{\text{sub}}H$		74.5 ± 0.4	298	C	[1980SAB2]
	$\Delta_{\text{sub}}H$		85.3	298		[1961ZIM/GEI]
C ₃ H ₄ N ₂	[288-13-1] $\Delta_{\text{fus}}H$	pyrazole	14.2	343.2		[1996DOM/HEA]
	$\Delta_{\text{sub}}H$	(268–287)	74.3 ± 0.4	275	ME	[1987JIM/ROU]
	$\Delta_{\text{sub}}H$		74.0 ± 0.4	298		[1987JIM/ROU, 1986JIM/ROU]
	$\Delta_{\text{sub}}H$	(253–273)	72.7	265	TE,ME	[1983DEW/VAN]
	$\Delta_{\text{sub}}H$		69.2 ± 0.3	298	C	[1980SAB2]
	$\Delta_{\text{sub}}H$		71.8			[1979DAA/VAN]
	$\Delta_{\text{sub}}H$		67.7			[1961ZIM/GEI]
C ₃ H ₄ N ₂ O	[107-91-5] $\Delta_{\text{trs}}H$	2-cyanoacetamide	1.2	346.5		
	$\Delta_{\text{fus}}H$		21.7	387.3		[1996DOM/HEA]
	$\Delta_{\text{sub}}H$	(325–348)	99.7	336	TE,ME	[1983DEW/VAN]
C ₃ H ₄ N ₂ O ₄	[4122-45-6] $\Delta_{\text{sub}}H$	3-nitro-2-isoxazoline-2-oxide	71.1 ± 8.4			[1977PED/RYL, 1969MIR/LEB]
C ₃ H ₄ N ₄ O ₂	[107945-73-3] $\Delta_{\text{sub}}H$	2-methyl-4-nitro-1,2,3-triazole (288–314)	74.6 ± 3.1	301	ME	[2008MAT/IVA]
	$\Delta_{\text{sub}}H$	(288–314)	74.7 ± 3.1	298	ME	[2008MAT/IVA]
C ₃ H ₄ N ₄ O ₆	[97645-24-4] $\Delta_{\text{fus}}H$	1,3,3-trinitroazetidine	29	373.8		[2003SUC/RAJ]
	$\Delta_{\text{fus}}H$		30.31	375.5		[1996ZHA/HU, 1997ZHA/HU]
	$\Delta_{\text{sub}}H$		95.3	373.8	Fus+Vap	[2003SUC/RAJ]
	Δ_vH	(373–413)	66.8	393		[2003SUC/RAJ]
C ₃ H ₄ O	[107-02-8] Δ_vH	acrolein (250–306)	32.3	265	A	[1987STE/MAL]
	Δ_vH	(304–325)	30.9	314		[1979MAR/SAC]
	Δ_vH	(208–326)	33.5	223	A	[1987STE/MAL, 1947STU]
C ₃ H ₄ O	[107-19-7] Δ_vH	propargyl alcohol (2-propyn-1-ol) (293–387)	42.0	308	A	[1987STE/MAL]
C ₃ H ₄ OS ₂	[2080-58-2] $\Delta_{\text{sub}}H$	1,3-dithiolan-2-one	80.3 ± 0.4			[1973RAU/GEI, 1977PED/RYL]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₃ H ₄ O ₂	[79-10-7]	acrylic acid				
	$\Delta_{\text{fus}}H$		11.16	285.5		[1991ACR]
	Δ_vH		53.1 ± 4.2	298	C	[1996VAN/YU]
	Δ_vH		57.3	298		[1980VIL/PER]
	Δ_vH	(341–414)	45.3	356	A	[1987STE/MAL, 1973LIN/WIC]
		(293–343)	32.7	308		[1964GUB/FER]
C ₃ H ₄ O ₂	[57-57-8]	β -propiolactone (2-oxetanone)				
	$\Delta_{\text{fus}}H$		9.41	239.9		[1996DOM/HEA]
	Δ_vH	(324–435)	46.4	339	A	[1987STE/MAL]
			47.0 ± 0.1	298	C	[1966BOR/NAK]
C ₃ H ₄ O ₂ S	[7285-32-7]	thiete sulfone 2 <i>H</i> -thiete-1,1-dioxide)				
$\Delta_{\text{sub}}H$			83.7 ± 2.5		B	[1969MAC/MCN, 1969MAC/MCN2, 1977PED/RYL]
C ₃ H ₄ O ₃	[96-49-1]	ethylene carbonate				
	$\Delta_{\text{fus}}H$		13.02	311.2	DSC	[2004DIN]
	$\Delta_{\text{fus}}H$		13.3	309.5		[1973VAS/KOR]
	$\Delta_{\text{sub}}H$	(273–297)	68.7	285	A	[1987STE/MAL]
	$\Delta_{\text{sub}}H$		78.5 ± 4.2			[1971CHO/JON, 1977PED/RYL]
	$\Delta_{\text{sub}}H$		73.2 ± 2.5			[1970COX/PIL, 1958HOY/PEP]
	Δ_vH	(310–369)	60.8 ± 0.1	298	GS	[2008VER/TOK]
	Δ_vH	(451.6–505)	64.0 ± 0.1	298	EB	[2004CHE/CLE, 2008VER/TOK]
	Δ_vH	(381–437)	59.6	396	A	[1987STE/MAL]
	Δ_vH	(368–449)	60.3	383	EB	[1982HON/WAK]
	Δ_vH	(368–449)	56.3	423	EB	[1982HON/WAK]
	Δ_vH	(368–433)	55.0	433	EB	[1982HON/WAK]
	Δ_vH	(382–437)	63.4 ± 0.3	298	EB	[1975PET/SAN, 2008VER/TOK]
			62.4	298	EB	[1958PEP, 2008VER/TOK]
C ₃ H ₄ O ₃	[127-17-3]	pyruvic acid				
Δ_vH	(294–438)	51.4	309	A	[1987STE/MAL, 1947STU]	
C ₃ H ₄ O ₄	[141-82-2]	malonic acid				
	$\Delta_{\text{fus}}H$		23.1	407.5		[2004HAN/BEY]
	$\Delta_{\text{sub}}H$	(339–357)	108.9 ± 0.7	348	ME	[1999RIB/MON]
	$\Delta_{\text{sub}}H$		111.4 ± 0.7	298		[1999RIB/MON]
	$\Delta_{\text{sub}}H$	(291–320)	72.7	306	A	[1987STE/MAL, 1947GRA]
			105.1 ± 0.8		C	[1983ALT/PIL]
C ₃ H ₄ O ₅	[80-69-3]	tartronic acid				
$\Delta_{\text{sub}}H$		116.4 ± 0.3			C	[1983ALT/PIL]
C ₃ H ₄ S ₃	[822-38-8]	1,3-dithiolan-2-thione				
$\Delta_{\text{sub}}H$	(294–303)	81.8 ± 0.8	298			[1967GEI/SCH, 1970COX/PIL]
C ₃ H ₄ S ₃	[822-38-8]	trithiocarbonic acid, cyclic ethylene ester				
Δ_vH	(294–303)	82.9	298			[1999DYK/SVO]
C ₃ H ₅ Br	[106-95-6]	allyl bromide				
	Δ_vH	(297–338)	32.2	312	A, EB	[1987STE/MAL, 1977SVO/MAJ]
	Δ_vH		31.7 ± 0.1	318	C	[1977SVO/MAJ]
	Δ_vH		31.0 ± 0.1	330	C	[1977SVO/MAJ]
			30.4 ± 0.1	341	C	[1977SVO/MAJ]
C ₃ H ₅ Br	[590-14-7]	<i>cis</i> 1-bromopropylene				

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
	$\Delta_{\text{v}}H$	(257–366)	32.0	272	A	[1987STE/MAL, 1970DYK]
C ₃ H ₅ Br	[590-14-7]	<i>trans</i> 1-bromopropylene				
	$\Delta_{\text{v}}H$	(262–372)	32.5	277	A	[1987STE/MAL, 1970DYK]
C ₃ H ₅ Br ₃	[96-11-7]	1,2,3-tribromopropane				
	$\Delta_{\text{fus}}H$		23.78	289.4		[1991ACR]
	$\Delta_{\text{v}}H$	(390–595)	50.8	405	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$	(400–478)	50.2	415		[1949DRE/SHR, 1949DRE/MAR, 1984BOU/FRI]
C ₃ H ₅ Cl	[107-05-1]	allyl chloride				
	$\Delta_{\text{v}}H$	(203–318)	33.1	218	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$	(286–317)	30.0	301		[1944IOF/YAM, 1984BOU/FRI]
C ₃ H ₅ Cl	[590-21-6]	1-chloropropene				
	$\Delta_{\text{v}}H$	(191–310)	29.5	206		[1947STU]
C ₃ H ₅ Cl	[16136-84-8]	<i>cis</i> 1-chloropropene				
	$\Delta_{\text{v}}H$	(276–332)	27.9	291		[2001HOR/GAR]
	$\Delta_{\text{v}}H$	(237–338)	29.2	252	A	[1987STE/MAL, 1970DYK]
C ₃ H ₅ Cl	[16136-85-9]	<i>trans</i> 1-chloropropene				
	$\Delta_{\text{v}}H$	(277–340)	28.5	292		[2001HOR/GAR]
	$\Delta_{\text{v}}H$	(241–343)	29.7	256	A	[1987STE/MAL, 1970DYK]
C ₃ H ₅ Cl	[557-98-2]	2-chloropropene				
	$\Delta_{\text{v}}H$	(229–327)	28.0	244	A	[1987STE/MAL, 1970DYK]
C ₃ H ₅ Cl	[107-05-1]	3-chloro-1-propene				
	$\Delta_{\text{v}}H$	(276–320)	29.9	298		[1960DJK]
C ₃ H ₅ ClO	[106-89-8]	epichlorohydrin				
	$\Delta_{\text{v}}H$	(256–391)	42.9	272		[1947STU]
C ₃ H ₅ ClO	[78-95-5]	chloroacetone				
	$\Delta_{\text{v}}H$	(316–392)	40.1	331	A	[1987STE/MAL]
C ₃ H ₅ ClO ₂	[96-34-4]	methyl chloroacetate				
	$\Delta_{\text{v}}H$		46.7 ± 0.1	298	C	[1972LAY/WAD]
	$\Delta_{\text{v}}H$	(318–402)	45.5	333	A	[1987STE/MAL, 1967GOE/SCH, 1984BOU/FRI]
	$\Delta_{\text{v}}H$	(298–403)	46.7	313		[1928NEL2, 1984BOU/FRI]
C ₃ H ₅ ClO ₂	[541-41-3]	ethylchloroformate				
	$\Delta_{\text{v}}H$	(281–286)	38.7 ± 0.2	283	BG	[1980DAV/FIN]
	$\Delta_{\text{v}}H$	(281–286)	37.8 ± 0.2	298	BG	[1980DAV/FIN]
C ₃ H ₅ ClO ₂	[27617-66-1]	(S) 2-chloropropionic acid				
	$\Delta_{\text{v}}H$	(287–328)	63.4	308	GS	[2002LAG/DIO]
	$\Delta_{\text{v}}H$	(287–328)	64.9 ± 0.5	298	GS	[2002LAG/DIO]
C ₃ H ₅ Cl ₃	[7789-89-1]	1,1,1-trichloropropane				
	$\Delta_{\text{v}}H$	(244–382)	38.8	259	A	[1987STE/MAL, 1947STU]
C ₃ H ₅ Cl ₃	[20395-25-9]	1,1,3-trichloropropane				
	$\Delta_{\text{v}}H$	(328–464)	41.8	343	A	[1987STE/MAL, 1970DYK]
C ₃ H ₅ Cl ₃	[96-18-4]	1,2,3-trichloropropane				
	$\Delta_{\text{v}}H$		47.8 ± 0.1	298	C	[1989AN/HU]
	$\Delta_{\text{v}}H$	(337–477)	43	352	A	[1987STE/MAL, 1970DYK]
	$\Delta_{\text{v}}H$	(361–429)	43	376		[1959URB]
	$\Delta_{\text{v}}H$	(282–431)	46.8	297		[1947STU]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₃ H ₅ FO	[503-09-3] $\Delta_v H$	1,2-epoxy-3-fluoropropane (273–333)	39.9	288	A, GS	[1987STE/MAL, 1948RED/CHA4, 1970DYK]
C ₃ H ₅ FO ₂	[453-18-9] $\Delta_v H$	methyl fluoroacetate (273–333)	42.7	288	A, GS	[1987STE/MAL, 1948RED/CHA4, 1970DYK]
C ₃ H ₅ F ₃ O	[428-66-0] $\Delta_v H$	1,1,2-trifluoro-1-methoxyethane (276–317)	31.1	291	I	[2002MUR/YAM]
C ₃ H ₅ F ₃ O	[374-01-6] $\Delta_v H$ $\Delta_v H$	1,1,1-trifluoro-2-propanol (292–333) (294–333)	44.2 44.8	307 298	A, MM MM	[1987STE/MAL, 1973ROC/SYM, 1984BOU/FRI] [1973ROC/SYM]
C ₃ H ₅ F ₃ O ₂ S	[30957-43-8] $\Delta_v H$	trifluoromethylsulfonic acid, trifluoromethyl ester 37.2	37.2	370		[1971SAU/SHR2]
C ₃ H ₅ F ₃ S ₂	[691-05-4] $\Delta_v H$	ethyl(trifluoromethyl) disulfide (253–303)	33.8	268	A	[1987STE/MAL, 1999DYK/SVO]
C ₃ H ₅ N	[107-12-0] $\Delta_{\text{trs}}H$ $\Delta_{\text{fus}}H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$	propionitrile (310–353) (288–371) (308–363) (189–295) (308–370) (294–394)	17.07 5.03 37.1 ± 0.3 36.1 36.7 36.5 35.9 36.7 ± 0.3	177 180.4 298 303 326 280 323 298	 EB A BG MM	 [1996DOM/HEA] [2004ANT/GAL, 2005EME/VER] [1987STE/MAL] [1971HAL/BAL] [1956MIL, 1984BOU/FRI] [1949DRE/SHR, 1949DRE/MAR, 1984BOU/FRI] [1933HEI, 2005EME/VER]
C ₃ H ₅ NO	[79-06-1] $\Delta_{\text{fus}}H$ $\Delta_{\text{sub}}H$ $\Delta_v H$ $\Delta_v H$	acrylamide (303–358) (357–413) (373–413)	15.33 81.8 61.5 76.5	358 330 372 388	 A A	 [1996DOM/HEA] [1957CAR/DAV] [1987STE/MAL] [1987STE/MAL]
C ₃ H ₅ NO	[1738-36-9] $\Delta_v H$	methoxyacetone (285–316)	41.7 ± 0.6	298	GS	[1995VER/BEC]
C ₃ H ₅ NO	[109-78-4] $\Delta_v H$ $\Delta_v H$	2-cyanoethanol (306–361) (331–494)	62.3 ± 0.5 53.4	298 346	GS A	[2007ROU/NOT] [1987STE/MAL]
C ₃ H ₅ NO	[5314-33-0] $\Delta_v H$	2-propenal oxime (303–381)	42.2	318	A	[1987STE/MAL]
C ₃ H ₅ NO	[930-21-2] $\Delta_{\text{sub}}H$	2-azetidinone 77.4 ± 0.3	77.4 ± 0.3	298	ME	[1996ROU/JIM2]
C ₃ H ₅ NOS	[5840-81-3] $\Delta_{\text{fus}}H$ $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$	2-mercapto-2-oxazoline 15.9 (325–354) (325–354)	15.9 104.3 ± 4.4 105.0 ± 4.4	370 340 298	DSC ME ME	[2008TEM/ROU3] [2009ROU/TEM] [2009ROU/TEM]
C ₃ H ₅ NO ₂	[3156-70-5] $\Delta_v H$ $\Delta_v H$	1-nitropropylene (301–373) (273–333)	37.1 44.1	337 288	 A	[1984BOS/TUR] [1987STE/MAL, 1970DYK]
C ₃ H ₅ NO ₂	[4749-28-4]	2-nitropropylene				

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(273–333)	38.2	288	A	[1987STE/MAL, 1970DYK]
C ₃ H ₅ NO ₂	[497-25-6]	dimethylene urethane (2-oxazolidinone)				
	$\Delta_{\text{fus}} H$		17.3	360		[2004SMI/MAR]
C ₃ H ₅ NS	[542-85-8]	ethyl isothiocyanate				
	$\Delta_v H$	(283–404)	40.2	298	A	[1987STE/MAL]
	$\Delta_v H$	(283–323)	39.8	298		[1935BAU/BUR, 1984BOU/FRI]
C ₃ H ₅ NS	[542-90-5]	ethyl thiocyanate				
	$\Delta_v H$	(358–422)	44.2	373	A	[1987STE/MAL, 1999DYK/SVO]
C ₃ H ₅ NS ₂	[96-53-7]	2-mercapto-2-thiazoline				
	$\Delta_{\text{fus}} H$		16.8	377	DSC	[2008TEM/ROU3]
	$\Delta_{\text{sub}} H$	(333–351)	99.8 ± 3.4	342	ME	[2009ROU/TEM]
	$\Delta_{\text{sub}} H$	(333–351)	100.5 ± 3.4	298	ME	[2009ROU/TEM]
C ₃ H ₅ N ₃ O ₉	[55-63-0]	glycerol trinitrate				
	$\Delta_{\text{fus}} H$		21.87	285.5		[1991ACR]
	$\Delta_v H$	(293–373)	104.5	308	A	[1987STE/MAL]
	$\Delta_v H$	(400–524)	58.6	415		[1947STU]
C ₃ H ₅ P	[114596-02-0]	2-propynylphosphine				
	$\Delta_v H$	(228–273)	36.8	250		[1988SHA/DIE]
C ₃ H ₆	[75-19-4]	cyclopropane				
	$\Delta_{\text{fus}} H$		5.44	145.6		[1996DOM/HEA]
	$\Delta_{\text{sub}} H$		29.2	145	B	[1963BON]
	$\Delta_{\text{sub}} H$	(115–141)	28.2	128	A,MS	[1951TIC/LOS]
	$\Delta_v H$	(195–225)	21.8	210		[1997CAL/FIL]
	$\Delta_v H$	(358–398)	20.4	373	A	[1987STE/MAL]
	$\Delta_v H$	(297–359)	19.9	312	A	[1987STE/MAL]
	$\Delta_v H$	(188–239)	20.3	224	A	[1987STE/MAL]
	$\Delta_v H$	(239–298)	19.9	254	A	[1987STE/MAL]
	$\Delta_v H$		17.02	298		[1970LIN/SIL]
	$\Delta_v H$	(183–241)	21.1	226		[1946RUE/POW, 1984BOU/FRI]
C ₃ H ₆	[115-07-1]	propylene				
	$\Delta_{\text{fus}} H$		2.93	88.2		[1996DOM/HEA]
	$\Delta_v H$	(297–363)	18.7	312	A	[1987STE/MAL]
	$\Delta_v H$	(104–161)	22.2	146	A	[1987STE/MAL]
	$\Delta_v H$	(228–271)	18.7	256	A	[1987STE/MAL]
	$\Delta_v H$	(270–327)	18.5	285	A	[1987STE/MAL]
	$\Delta_v H$	(325–363)	18.8	340	A	[1987STE/MAL]
	$\Delta_v H$	(161–242)	19.2	227	A	[1987STE/MAL, 1970DYK]
	$\Delta_v H$	(298–423)	18.7	360		[1953MIC/WAS]
	$\Delta_v H$	(166–226)	19.6	211		[1939POW/GIA]
$\Delta_v H$	(236–283)	19.3	268		[1921MAA/WRI, 1984BOU/FRI]	
C ₃ H ₆ BrCl	[109-70-6]	1-bromo-3-chloropropane				
	$\Delta_v H$	(326–488)	42.0	341	A	[1987STE/MAL, 1970DYK]
C ₃ H ₆ BrNO	[7119-91-7]	2-bromo-2-nitrosopropane				
	$\Delta_v H$	(239–356)	41.0	254	A	[1987STE/MAL, 1970DYK]
C ₃ H ₆ Br ₂	[598-17-4]	1,1-dibromopropane				

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(322–449)	42.5	337	A, E	[1987STE/MAL, 1956MAN, 1970DYK]
C ₃ H ₆ Br ₂	[78-75-1]	1,2-dibromopropane				
	$\Delta_v H$	(312–403)	41.4	327	A	[1987STE/MAL]
	$\Delta_v H$	(310–400)	42.2	298		[1991BAS/SVO, 1975PIS/ROZ2]
	$\Delta_v H$		42.3 ± 0.7	298	EB	[1975PIS/ROZ]
	$\Delta_v H$	(329–456)	44.6	344	A	[1987STE/MAL, 1970DYK]
	$\Delta_v H$	(266–415)	42.8	281		[1947STU]
C ₃ H ₆ Br ₂	[109-64-8]	1,3-dibromopropane				
	$\Delta_{\text{fus}} H$		14.64	238.6		[1996DOM/HEA]
	$\Delta_v H$		47.6	298	GC	[1994CAR/LAY]
	$\Delta_v H$		47.3 ± 0.1	308	C	[1992SVO/KUB2]
	$\Delta_v H$		46.7 ± 0.1	315	C	[1992SVO/KUB2]
	$\Delta_v H$		46.1 ± 0.1	323	C	[1992SVO/KUB2]
	$\Delta_v H$		45.5 ± 0.1	330	C	[1992SVO/KUB2]
	$\Delta_v H$		44.8 ± 0.1	338	C	[1992SVO/KUB2]
	$\Delta_v H$	(307–437)	46.6	322	A	[1987STE/MAL]
	$\Delta_v H$	(351–487)	47.8	366	A	[1987STE/MAL, 1970DYK]
$\Delta_v H$	(283–440)	45.3	298		[1947STU]	
C ₃ H ₆ Br ₂ O	[96-13-9]	2,3-dibromo-1-propanol				
	$\Delta_v H$	(330–492)	57.3	345	A	[1987STE/MAL, 1947STU]
C ₃ H ₆ ClNO ₂	[594-71-8]	2-chloro-2-nitropropane				
	$\Delta_{\text{trs}} H$		9.54	213.8		
	$\Delta_{\text{fus}} H$		1.34	261.6		[1996DOM/HEA]
C ₃ H ₆ Cl ₂	[78-99-9]	1,1-dichloropropane				
	$\Delta_v H$		35.2 ± 0.4	298	C	[2007VAR/DRU]
	$\Delta_v H$	(310–360)	35.2	298		[1967HAC/MAT, 1991BAS/SVO]
	$\Delta_v H$	(282–399)	35.5	297	A, E	[1987STE/MAL, 1956MAN, 1970DYK]
C ₃ H ₆ Cl ₂	[78-87-5]	1,2-dichloropropane				
	$\Delta_{\text{fus}} H$		6.4	172.7		[1991ACR]
	$\Delta_v H$	(303–368)	36.3 ± 0.5	298		[2007VAR/DRU]
	$\Delta_v H$		36.2 ± 0.1	298	C	[2007VAR/DRU]
	$\Delta_v H$	(294–406)	38.4 ± 0.3	298	EB	[1997STE/CHI3]
	$\Delta_v H$	(300–370)	36.3	298		[1991BAS/SVO]
	$\Delta_v H$		36.1 ± 0.1	298	C	[1989AN/HU]
	$\Delta_v H$	(239–373)	39.4	254	A	[1987STE/MAL]
	$\Delta_v H$	(321–369)	34.7	336		[1949DRE/SHR, 1949DRE/MAR]
	$\Delta_v H$	(288–373)	34.3	303		[1933NEL/YOU]
C ₃ H ₆ Cl ₂	[142-28-9]	1,3-dichloropropane				
	$\Delta_v H$		41.0	298	GC	[1994CAR/LAY]
	$\Delta_v H$	(330–400)	41.0	298		[1987VAR/LOS, 1991BAS/SVO]
	$\Delta_v H$		40.6 ± 0.1	298	C	[1989AN/HU]
	$\Delta_v H$	(307–435)	39	322	A	[1987STE/MAL, 1970DYK]
C ₃ H ₆ Cl ₂	[594-20-7]	2,2-dichloropropane				
	$\Delta_{\text{trs}} H$		0.01	171.6		
	$\Delta_{\text{trs}} H$		0.73	188.2		
	$\Delta_{\text{fus}} H$		10.0	239.6		[1999KOB/OGU]
	$\Delta_{\text{trs}} H$		5.98	188		
	$\Delta_{\text{fus}} H$		2.34	239.3		[1996DOM/HEA]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
	$\Delta_{\text{v}}H$	(295–340)	32.1	298	A	[1987VAR/LOS, 1991BAS/SVO]
	$\Delta_{\text{v}}H$	(267–378)	33.2	282	A	[1987STE/MAL, 1970DYK]
C₃H₆Cl₂O	[616-23-9]	2,3-dichloro-1-propanol				
	$\Delta_{\text{v}}H$	(384–419)	48.5	399	A	[1987STE/MAL]
C₃H₆Cl₂O	[96-23-1]	1,3-dichloro-2-propanol				
	$\Delta_{\text{v}}H$	(301–448)	50.4	316	A	[1987STE/MAL, 1947STU]
C₃H₆F₂	[430-81-5]	1,1-difluoropropane				
	$\Delta_{\text{v}}H$	(219–311)	27.2	234	A, E	[1987STE/MAL, 1956MAN, 1970DYK]
C₃H₆F₂	[420-45-1]	2,2-difluoropropane				
	$\Delta_{\text{v}}H$	(211–302)	25.6	226	A	[1987STE/MAL, 1970DYK]
C₃H₆F₂O	[461-57-4]	1,1-difluoro-2-methoxyethane				
	$\Delta_{\text{v}}H$	(288–322)	31.8	303	I	[2002MUR/YAM]
C₃H₆F₃NS	[62067-13-4]	N,N-dimethyl-trifluoromethanesulfenamide				
	$\Delta_{\text{v}}H$	(223–295)	30.2	259		[1960EME/NAB]
C₃H₆F₃NS	[62067-13-4]	dimethyl(trifluoromethylthio)amine				
	$\Delta_{\text{v}}H$	(273–329)	31.1	288	A	[1987STE/MAL, 1999DYK/SVO]
C₃H₆F₃OP	[26348-84-5]	methyl(trifluoromethyl)phosphinous acid, methyl ester				
	$\Delta_{\text{v}}H$	(232–285)	33.9	258		[1970BUR/KAN]
C₃H₆F₃OP	[26348-91-4]	dimethyl(trifluoromethyl)phosphine oxide				
	$\Delta_{\text{v}}H$	(347–360)	52.4	358		[1970BUR/KAN]
C₃H₆F₃O₂P	[684-56-0]	(trifluoromethyl)phosphonic acid, dimethyl ester				
	$\Delta_{\text{v}}H$	(237–318)	37.4	252	A	[1987STE/MAL, 1961BUR/GRI]
C₃H₆F₃PS	[26348-86-7]	methyl(trifluoromethyl)phosphinothious acid, methyl ester				
	$\Delta_{\text{v}}H$	(273–313)	38.4	293		[1970BUR/KAN]
C₃H₆F₃PS	[26348-92-5]	dimethyl(trifluoromethyl)phosphine sulfide				
	$\Delta_{\text{sub}}H$	(300–320)	68.0	310		[1970BUR/KAN]
	$\Delta_{\text{v}}H$	(323–357)	47.2	340		[1970BUR/KAN]
C₃H₆I₂	[627-31-6]	1,3-diiodopropane				
	$\Delta_{\text{v}}H$		54.1	298	GC	[1994CAR/LAY]
C₃H₆N₂O	[120-93-4]	2-imadazolidinone				
	$\Delta_{\text{trs}}H$		3.6	344.6		
	$\Delta_{\text{fus}}H$		11.5	401.2		[2008RIB/RIB]
	$\Delta_{\text{trs}}H$		3.27	344		
	$\Delta_{\text{fus}}H$		5.11	397.3	DSC	[1984WEI/LEF]
	$\Delta_{\text{sub}}H$	(327–349)	96.6 ± 0.8	298	ME	[2008RIB/RIB]
	$\Delta_{\text{sub}}H$		83.7	298		[1999DEF/DEO]
C₃H₆N₂O₂	[591-07-1]	acetylurea				
	$\Delta_{\text{sub}}H$	(360–407)	102.4 ± 0.7	383		[1988IMA/MUR]
	$\Delta_{\text{sub}}H$		103.1 ± 0.7	298		[1988IMA/MUR]
	$\Delta_{\text{sub}}H$		103.1 ± 0.7	298	C	[1985MUR/SAK]
C₃H₆N₂O₂	[108-13-4]	malonamide				
	$\Delta_{\text{trs}}H$		1.89	418.9		
	$\Delta_{\text{fus}}H$		29.85	444.2	DSC	[2006BAD/DEL]
	$\Delta_{\text{trs}}H$		1.9	393		

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	Enthalpy							
		$\Delta_{\text{fus}}H$			35.8	443		[1996DOM/HEA]
		$\Delta_{\text{sub}}H$			126.4 ± 0.5		C	[1989IMA/TAK]
C ₃ H ₆ N ₂ O ₄	[601-76-3]		1,1-dinitropropane	(323–383)	57.9	338	A	[1987STE/MAL]
C ₃ H ₆ N ₂ O ₄	[595-49-3]		2,2-dinitropropane					
		$\Delta_{\text{trs}}H$			11.28	267.7		
		$\Delta_{\text{trs}}H$			1.87	259.7		
		$\Delta_{\text{fus}}H$			2.64	324.5		[1996DOM/HEA]
		Δ_vH		(363–553)	46.3	378	A	[1987STE/MAL]
C ₃ H ₆ N ₂ O ₅	[918-52-5]		2,2-dinitro-1-propanol					
		$\Delta_{\text{trs}}H$			15.06	281.7		
		$\Delta_{\text{fus}}H$			2.85	366.7		[1969ROS/HOL]
C ₃ H ₆ N ₂ O ₆	[6423-43-4]		1,2-propanediol dinitrate	(288–328)	63.8	303	A	[1987STE/MAL, 1970DYK]
C ₃ H ₆ N ₂ O ₆	[3457-90-7]		1,3-propanediol dinitrate	(293–313)	74.3 ± 4.6	303	A, GS	[1987STE/MAL, 1957KEM/GOL]
C ₃ H ₆ N ₂ O ₆	[2736-80-3]		2,2-dinitro-1,3-propanediol					
		$\Delta_{\text{trs}}H$			21.34	341.2		[1969ROS/HOL]
			Note: decomposes before melting.					
C ₃ H ₆ N ₄	[5144-11-6]		1,5-dimethyltetrazole					
		$\Delta_{\text{fus}}H$			14.7	349		[1990KOZ/SIM3]
		$\Delta_{\text{sub}}H$		(303–343)	86.2 ± 1.0		ME	[1990KOZ/SIM]
C ₃ H ₆ N ₄	[4135-93-7]		2,5-dimethyltetrazole					
		$\Delta_{\text{fus}}H$			13.5	256.4		[1990KOZ/SIM3]
C ₃ H ₆ N ₄ O ₄	[5754-91-6]		1,3-dinitro-1,3-diazacyclopentane					
		$\Delta_{\text{fus}}H$			25.08	410		[1996DOM/HEA]
C ₃ H ₆ N ₆	[108-78-1]		2,4,6-triamino-s-triazine (melamine)					
		$\Delta_{\text{sub}}H$		(417–614)	121.3 ± 4.2	515	GS	[1960HIR/STE, 1970COX/PIL]
		$\Delta_{\text{sub}}H$		(417–447)	123.3	432	A	[1987STE/MAL]
C ₃ H ₆ N ₆ O ₃	[13980-04-6]		1,3,5-trinitroso-1,3,5-triazacyclohexane					
		$\Delta_{\text{trs}}H$			17.78	367		
		$\Delta_{\text{fus}}H$			3.77	376		[1996DOM/HEA]
		$\Delta_{\text{sub}}H$		(343–447)	134.3 ± 0.7	298	ME	[1978CUN/PAL]
		$\Delta_{\text{sub}}H$		(383–411)	112.1		ME	[1974PEP/MAT]
		$\Delta_{\text{sub}}H$			112.1			[1953EDW, 1960JON]
C ₃ H ₆ N ₆ O ₅	[5755-27-1]		1,3-dinitro-5-nitroso-1,3,5-triazacyclohexane					
		$\Delta_{\text{fus}}H$			25.97	446		[1987OYU/BR1]
C ₃ H ₆ N ₆ O ₆	[121-82-4]		hexahydro-1,3,5-trinitro-1,3,5-triazine					
		$\Delta_{\text{fus}}H$			37.66	478.2		[1996DOM/HEA]
		$\Delta_{\text{sub}}H$			134.3	298		[1978CUN/PAL]
		$\Delta_{\text{sub}}H$		(325–360)	112.5 ± 0.8		ME	[1974PEP/MAT]
		$\Delta_{\text{sub}}H$		(329–371)	130.1	350		[1969ROS/DIC]
		Δ_vH		(503–523)	84.4	513	A	[1987STE/MAL]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₃ H ₆ O	[107-18-6]	2-propen-1-ol				
	$\Delta_v H$	(311–355)	46.1	298	EB	[2004LUB/MAL]
	$\Delta_v H$	(323–373)	47.3	298	CGC	[1995CHI/HOS]
C ₃ H ₆ O	[503-30-0]	oxetane				
	$\Delta_v H$		29.8	298	C	[1981HOS/SCO]
C ₃ H ₆ O	[67-64-1]	acetone				
	$\Delta_{\text{fus}} H$		5.72	176.6		[1996DOM/HEA]
	$\Delta_v H$	(298–318)	32.1	308		[2008SON/RAM]
	$\Delta_v H$	(329–488)	29.9	344	A	[1987STE/MAL]
	$\Delta_v H$	(178–243)	32.9	228	A	[1987STE/MAL]
	$\Delta_v H$	(203–269)	33.8	254	A	[1987STE/MAL]
	$\Delta_v H$	(323–379)	30.6	338	A	[1987STE/MAL]
	$\Delta_v H$	(374–464)	29.5	389	A	[1987STE/MAL]
	$\Delta_v H$	(457–508)	29.7	472	A	[1987STE/MAL]
	$\Delta_v H$		26.1	373	C	[1986DMI/KAC]
	$\Delta_v H$		21.7	423	C	[1986DMI/KAC]
	$\Delta_v H$		15.3	473	C	[1986DMI/KAC]
	$\Delta_v H$		9.2	498	C	[1986DMI/KAC]
	$\Delta_v H$	(285–329)	31.9	300	EB	[1986BAL/GNA]
	$\Delta_v H$	(305–333)	31.8	319		[1984CAS/FRA3]
	$\Delta_v H$	(259–351)	32.8	274	A	[1987STE/MAL, 1974AMB/SPR2, 1975AMB/ELL]
	$\Delta_v H$		31.3	298		[1975AMB/ELL]
	$\Delta_v H$	(261–328)	32.7	276	A, EB	[1987STE/MAL, 1972BOU/AIM]
	$\Delta_v H$	(278–293)	32.6	285		[1963SOK/ZHI]
	$\Delta_v H$	(310–329)	31.1	319		[1957BRO/SMI]
$\Delta_v H$	(204–339)	35	253	MG	[1926FEL/DUR]	
$\Delta_v H$	(204–339)	32.1	293	MG	[1926FEL/DUR]	
$\Delta_v H$	(204–339)	30.7	313	MG	[1926FEL/DUR]	
C ₃ H ₆ O	[107-18-6]	allyl alcohol				
	$\Delta_v H$	(310–340)	44.6	325		[2002LUB/BAN]
	$\Delta_v H$	(253–370)	46.7	268	A	[1987STE/MAL]
	$\Delta_v H$				NA	[1936EWE]
	$\Delta_v H$	(283–313)	44.8	298		[1935BAU/BUR]
C ₃ H ₆ O	[107-25-5]	methyl vinyl ether				
	$\Delta_v H$	(278–412)	23.4	293	A	[1987STE/MAL]
C ₃ H ₆ O	[503-30-3]	trimethylene oxide				
	$\Delta_{\text{fus}} H$	(85–270)	6.27	173.2		[1985HAN]
C ₃ H ₆ O	[123-38-6]	propanal				
	$\Delta_{\text{fus}} H$		8.59	171.3		[1996DOM/HEA]
	$\Delta_v H$	(263–373)	31.5	278		[1977KIM/KIM]
	$\Delta_v H$	(286–321)	30.5	301		[1974AMB/SPR]
	$\Delta_v H$		28.3	321		[1972COU/LEE]
	$\Delta_v H$		29.4	303		[1972COU/LEE]
	$\Delta_v H$		30.3	286		[1972COU/LEE]
	$\Delta_v H$		29.6	298		[1972COU/LEE]
	$\Delta_v H$	(290–322)	30.3	305	A	[1987STE/MAL, 1970DYK]
	$\Delta_v H$		29.7 ± 0.4	298	EB	[1967BUC/COX, 2003VER/KRA2]
	$\Delta_v H$		30.0	298	EB	[1962TJE2, 2003VER/KRA2]
$\Delta_v H$	(250–330)	31.9	265	EB	[1987STE/MAL, 1951SMI/BON]	
C ₃ H ₆ O	[75-56-9]	propylene oxide				

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	Enthalpy							
		$\Delta_{\text{fus}}H$			6.57	161.3		[1996DOM/HEA]
		Δ_vH	(225–308)		31.6	240	A	[1987STE/MAL, 1970DYK]
		Δ_vH	(292–345)		28.5	307		[1966BOT/ADL]
		Δ_vH			27.9	298	C	[1962SIN/HIL]
		Δ_vH	(249–308)		30.1	264		[1959MCD/SHR]
		Δ_vH	(285–322)		28.2	303		[1937MOO/KAN]
		Δ_vH	(243–306)		32.9	273		[1935KIR/POP]
C₃H₆O₂	[646-06-0]		1,3-dioxolane					
		$\Delta_{\text{us}}H$			2.68	142.4		
		$\Delta_{\text{fus}}H$			6.57	175.9		[1996DOM/HEA]
		Δ_vH	(305–347)		34.6	326	A	[1989WU/SAN]
		Δ_vH	(280–323)		35.8	295	A	[1987STE/MAL]
		Δ_vH	(321–357)		33.7	339		[1982CAS/FRA]
		Δ_vH	(306–346)		33.7	326		[1980FRA/CAS]
		Δ_vH	(280–355)		34.1	296		[1968CHE/TUR, 1984BOU/FRI]
		Δ_vH			35.6 ± 0.4			[1959FLE/MOR]
C₃H₆O₂	[109-94-4]		ethyl formate					
		Δ_vH	(300–326)		31.4	313		[1993FAR/WIC]
		Δ_vH	(327–498)		29.9	342	A	[1987STE/MAL]
		Δ_vH			31.6 ± 0.1	304	C	[1976CIH/HYN]
		Δ_vH			30.9 ± 0.1	313	C	[1976CIH/HYN]
		Δ_vH			29.8 ± 0.1	328	C	[1976CIH/HYN]
		Δ_vH	(213–336)		35.8	228	A	[1987STE/MAL, 1970DYK]
C₃H₆O₂	[79-20-9]		methyl acetate					
		$\Delta_{\text{fus}}H$			7.49	174.9		[1992OKA/OGU]
		Δ_vH	(260–351)		34.1	275	A	[1987STE/MAL]
		Δ_vH			32.3 ± 0.1	298	C	[1980SVO/UCH]
		Δ_vH			29.5 ± 0.1	343	C	[1980SVO/UCH]
		Δ_vH	(308–338)		31.8	323	DTA	[1980MEY/AWE]
		Δ_vH			32.6 ± 0.1	298	C	[1979SUN/SVE2]
		Δ_vH			32.2 ± 0.1	304	C	[1977SVO/VES]
		Δ_vH			31.6 ± 0.1	313	C	[1977SVO/VES]
		Δ_vH			30.5 ± 0.1	328	C	[1977SVO/VES]
		Δ_vH			30.3 ± 0.1	331	C	[1977SVO/VES]
		Δ_vH			32.5	295		[1976CON/COU]
		Δ_vH			30.2	330		[1976CON/COU]
		Δ_vH	(273–318)		34.5	296	BG	[1971HAL/BAL]
		Δ_vH	(274–329)		33.4	289	A	[1987STE/MAL, 1965MER/POL, 1970DYK]
C₃H₆O₂	[79-09-4]		propionic acid					
		$\Delta_{\text{fus}}H$			10.66	252.7		[1996DOM/HEA, 1982MAR/AND]
		$\Delta_{\text{sub}}H$	(225–238)		74.1 ± 1	233	TE	[1978CAL/CAL]
		$\Delta_{\text{sub}}H$			73.2 ± 1	233	ME	[1978CAL/CAL]
		Δ_vH	(303–378)		54.4	298	CGC	[2000VER]
		Δ_vH	(353–393)		54.9	298	CGC	[1995CHI/HOS]
		Δ_vH	(343–419)		47.0	358	A	[1987STE/MAL]
		Δ_vH	(414–511)		60.6	429	A	[1987STE/MAL]
		Δ_vH	(345–401)		46.4	360	A	[1987STE/MAL]
		Δ_vH			56.0	303		[1983TAM/DRA]
		Δ_vH (monomer)			31.1 ± 0.1	298	C	[1970KON/WAD]
		Δ_vH			55 ± 2	298	C	[1970KON/WAD]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(328–437)	48.3	343		[1981AMB/ELL]
(C ₃ H ₆ O ₂) ₂	[32574-16-6]	propionic acid dimer				
	$\Delta_{\text{sub}} H$	(225–238)	81.3 ± 1	233	TE	[1978CAL/CAL]
	$\Delta_{\text{sub}} H$		79.4 ± 1.0	233	ME	[1978CAL/CAL]
C ₃ H ₆ O ₂	[116-09-6]	hydroxyacetone				
	$\Delta_v H$	(296–356)	42 ± 3	326		[2010PET/REY]
C ₃ H ₆ O ₂ S	[107-96-0]	β -thiolactic acid				
	$\Delta_{\text{fus}} H$		16.97	291.9		[1996DOM/HEA]
C ₃ H ₆ O ₃	[625-45-6]	2-methoxyacetic acid				
	$\Delta_v H$	(325–477)	54.5	340	A	[1987STE/MAL, 1947STU]
C ₃ H ₆ O ₃	[96-35-5]	methyl glycolate				
	$\Delta_{\text{fus}} H$		11.4	272.8		[2000JAR/MAR]
	$\Delta_v H$	(326–381)	52.5 ± 6.3	298	EB	[1996STE/CHI2]
	$\Delta_v H$	(282–425)	47.4	297	A	[1987STE/MAL, 1947KET/VAN]
	[616-38-6]	dimethylcarbonate				
	$\Delta_{\text{trs}} H$		NA	220.1		
	$\Delta_{\text{fus}} H$		11.58	278.2		[2004DIN]
	$\Delta_v H$	(274–304)	38.0 ± 0.2	298	GS	[2008KOZ/EME]
	$\Delta_v H$	(326–411)	36.4	341		[2002ROD/CAN]
	$\Delta_v H$	(311–397)	37.7 ± 0.2	298	EB	[1997STE/CHI4, 1997STE/CHI2]
C ₃ H ₆ O ₃	[4212-43-5]	peroxypropionic acid				
	$\Delta_v H$	(273–393)	43.2	288	A	[1987STE/MAL, 1951EGE/EMT, 1970DYK]
C ₃ H ₆ O ₃	[38787-96-1]	propylene ozonide				
	$\Delta_v H$	(261–296)	36.9	281	A	[1987STE/MAL, 1956GAR/SCH]
C ₃ H ₆ O ₃	[110-88-3]	1,3,5-trioxane				
	$\Delta_{\text{fus}} H$		15.1	333.4		[1996DOM/HEA, 1991ACR]
	$\Delta_{\text{sub}} H$	(212–231)	57.9	223	TE,ME	[1983DEW/VAN]
	$\Delta_{\text{sub}} H$		55.6	298		[1983DEW/VAN]
	$\Delta_{\text{sub}} H$		56.5	298	C	[1975BOG/BER]
	$\Delta_{\text{sub}} H$		56.2 ± 0.2	298	C	[1969MAN/MOR, 1977PED/RYL]
	$\Delta_v H$	(329–386)	40.0	344	A	[1987STE/MAL, 1965SER/BYK]
C ₃ H ₆ O ₃	[50-21-5]	<i>(dl)</i> lactic acid				
	$\Delta_{\text{fus}} H$		11.34	289.9		[1996DOM/HEA]
C ₃ H ₆ S	[1072-43-1]	2-methylthiirane				
	$\Delta_v H$	(272–423)	34.6	287	A	[1987STE/MAL, 1970DYK, 1999DYK/SVO]
C ₃ H ₆ S	[287-27-4]	thiacyclobutane (thietane)				
	$\Delta_{\text{trs}} H$		0.67	176.7		
	$\Delta_{\text{fus}} H$		8.24	199.9		[1996DOM/HEA]
	$\Delta_v H$	(275–393)	36.5	290		[1999DYK/SVO]
	$\Delta_v H$		35.8	298		[1971WIL/ZWO]
	$\Delta_v H$	(321–404)	34.6	336	A, EB	[1987STE/MAL, 1953SCO/FIN, 1966OSB/DOU]
C ₃ H ₆ S ₃	[291-21-4]	1,3,5-trithiane				
	$\Delta_{\text{fus}} H$		32.2	488.4		[2002VAN/VAN2]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{sub}}H$		93.2 ± 0.2	298	ME	[2001ROU/JIM]
	$\Delta_{\text{sub}}H$	(320–339)	91.5	331	TE,ME	[1983DEW/VAN]
	$\Delta_{\text{sub}}H$		93.9	298		[1983DEW/VAN]
C₃H₇Br	[106-94-5]	1-bromopropane				
	Δ_vH	(301–344)	31.8	316	A, EB	[1987STE/MAL, 1977SVO/MAJ]
	Δ_vH		31.1 ± 0.1	322	C	[1977SVO/MAJ]
	Δ_vH		30.5 ± 0.1	332	C	[1977SVO/MAJ]
	Δ_vH		30.1 ± 0.1	339	C	[1977SVO/MAJ]
	Δ_vH		29.3 ± 0.1	352	C	[1977SVO/MAJ]
	Δ_vH		31.9 ± 0.1	298	C	[1966WAD]
	Δ_vH	(250–368)	34.1	265	A, E	[1987STE/MAL, 1961LI/ROS, 1970DYK]
	Δ_vH	(220–344)	35.5	235		[1947STU]
	Δ_vH	(273–303)	32.6	288		[1906REX, 1984BOU/FRI]
C₃H₇Br	[75-26-3]	2-bromopropane				
	$\Delta_{\text{fus}}H$		6.55	184.1		[1996DOM/HEA]
	Δ_vH	(323–363)	30.6	298	CGC	[1995CHI/HOS]
	Δ_vH	(236–328)	32.1	251	A	[1987STE/MAL]
	Δ_vH	(299–332)	30.1	314	EB	[1987STE/MAL, 1977SVO/MAJ]
	Δ_vH		29.8 ± 0.1	305	C	[1977SVO/MAJ]
	Δ_vH		29.2 ± 0.1	318	C	[1977SVO/MAJ]
	Δ_vH		28.5 ± 0.1	330	C	[1977SVO/MAJ]
	Δ_vH		28.0 ± 0.1	338	C	[1977SVO/MAJ]
	Δ_vH		30.2 ± 0.1	298	C	[1966WAD]
	Δ_vH	(211–333)	33.4	226		[1947STU]
	Δ_vH	(273–303)	30.9	288		[1906REX, 1984BOU/FRI]
C₃H₇Cl	[540-54-5]	1-chloropropane				
	Δ_vH	(250–320)	29.0	298		[1984BOU/FRI, 1991BAS/SVO]
	Δ_vH		28.5 ± 0.2	298	C	[1977MAN/SEL]
	Δ_vH	(248–320)	31.0	263	A	[1987STE/MAL, 1969KEM/KRE, 1970DYK]
	Δ_vH	(205–319)	33.1	219		[1947STU]
C₃H₇Cl	[75-29-6]	2-chloropropane				
	$\Delta_{\text{fus}}H$		7.39	156		[1991ACR]
	Δ_vH	(239–310)	30.2	254	A	[1987STE/MAL]
	Δ_vH	(271–312)	27.6	298		[1960DJK]
	Δ_vH	(194–309)	30.6	209		[1947STU]
	Δ_vH	(273–303)	27.3	288		[1906REX, 1984BOU/FRI]
C₃H₇ClO	[127-00-4]	1-chloro-2-propanol				
	Δ_vH	(308–399)	45.0 ± 2.2	340	EB	[2002STE/CHI]
	Δ_vH	(308–399)	42.2 ± 1.9	380	EB	[2002STE/CHI]
C₃H₇ClO	[78-89-7]	2-chloro-1-propanol				
	Δ_vH	(316–399)	45.0	331	A	[1987STE/MAL]
C₃H₇ClO₂	[96-24-2]	3-chloro-1,2-propanediol				
	Δ_vH	(343–409)	66.6	358		[1996GIL/WIL]
C₃H₇ClO₂S	[10147-36-1]	1-propanesulfonyl chloride				
	Δ_vH	(273–362)	52.3	288		[1999DYK/SVO]
	Δ_vH	(362–464)	49.9	377		[1999DYK/SVO]
	Δ_vH	(243–273)	60.1	258	A	[1987STE/MAL, 1999DYK/SVO]
C₃H₇ClS	[na]	methyl(2-chloroethyl) sulfide				

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(293–333)	42.4	308	A, GS	[1987STE/MAL, 1948RED/CHA, 1970DYK]
C ₃ H ₇ F	[460-13-9]	1-fluoropropane				
	$\Delta_v H$	(196–289)	24.0	274	A, E	[1987STE/MAL, 1961LI/ROS, 1970DYK]
C ₃ H ₇ F	[420-26-8]	2-fluoropropane				
	$\Delta_v H$	(190–264)	23.7	249	A	[1987STE/MAL]
C ₃ H ₇ I	[107-08-4]	1-iodopropane				
	$\Delta_v H$	(171–271)	37.8	256	A	[1987STE/MAL]
	$\Delta_v H$		36.3 ± 0.1	298	C	[1968WAD]
	$\Delta_v H$	(271–402)	36.8	286	A, E	[1987STE/MAL, 1961LI/ROS, 1970DYK]
$\Delta_v H$	(237–375)	37.0	252		[1947STU]	
C ₃ H ₇ I	[75-30-9]	2-iodopropane				
	$\Delta_v H$	(313–353)	34.0	298	CGC	[1995CHI/HOS]
	$\Delta_v H$	(173–262)	36.7	247	A	[1987STE/MAL]
	$\Delta_v H$		34.1 ± 0.1	298	C	[1968WAD]
$\Delta_v H$	(230–363)	36.3	244		[1947STU]	
C ₃ H ₇ N	[765-30-0]	cyclopropylamine				
	$\Delta_{\text{fus}} H$		13.18	237.8		[1991ACR]
$\Delta_v H$		31.3 ± 0.4	298	EB	[1971GOO/MOO]	
C ₃ D ₇ N	[153557-96-1]	perdeuterocyclopropylamine				
	$\Delta_v H$	(283–336)	32.0	298		[1993WOL/KIM]
C ₃ H ₇ N	[107-11-9]	allylamine				
	$\Delta_v H$	(273–303)	33.0	288	A	[1987STE/MAL]
	$\Delta_v H$	(273–324)	32.6	288	A	[1987STE/MAL]
C ₃ H ₇ N	[503-29-7]	azetidine				
	$\Delta_v H$	(273–303)	32.6	288	A	[1987STE/MAL]
C ₃ H ₇ NO	[127-06-0]	acetone oxime				
	$\Delta_{\text{sub}} H$	(313–333)	59.6	323	I	[1987STE/MAL, 1975MES/BAE]
$\Delta_v H$	(338–352)	51.4	345	A	[1987STE/MAL]	
C ₃ H ₇ NO	[79-05-0]	propionamide				
	$\Delta_{\text{fus}} H$		12.9	352.6		[2008ABA/BAD]
	$\Delta_{\text{fus}} H$		12.9	352.6		[2000BRU/DEL]
	$\Delta_{\text{sub}} H$	(283–343)	75 ± 4.0	298	TE	[2000BRU/DEL]
	$\Delta_{\text{sub}} H$		79.2 ± 0.3			[1975BAR/PIL, 1977PED/RYL]
	$\Delta_{\text{sub}} H$		73.3			[1960THO]
	$\Delta_{\text{sub}} H$	(318–346)	79.1 ± 0.4		GS	[1959DAV/JON2]
	$\Delta_v H$	(375–476)	63.9	390	EB	[2004HOR/FIS]
$\Delta_v H$	(338–486)	60.3	353		[1947STU]	
C ₃ H ₇ NO	[68-12-2]	N,N-dimethylformamide				
	$\Delta_{\text{fus}} H$		8.95	212.9	AC	[2007SMI/TSV]
	$\Delta_{\text{fus}} H$		8.95	212.9		[1996DOM/HEA]
	$\Delta_v H$	(463–513)	46.7 ± 0.5	298	CGC	[2009PAN/ANT]
	$\Delta_v H$	(346–425)	43.1	361		[2005MUN/MON]
	$\Delta_v H$	(377–426)	41.8	392		[1997BLA/BEL]
	$\Delta_v H$	(338–425)	43.6	353		[1995MAR/GAB]
	$\Delta_v H$	(301–426)	49.2	316	A	[1987STE/MAL]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$		46.9	298	A	[1985BAR/CAS, 1985MAJ/SVO]
	$\Delta_v H$	(318–423)	42.5	370		[1979BLU/BAE]
	$\Delta_v H$	(331–425)	56.7	346		[1974MYA/SCH, 1984BOU/FRI]
	$\Delta_v H$	(303–363)	46.7	318		[1968GOP/RIZ]
C₃H₇NO	[627-45-2]	N-ethylformamide				
	$\Delta_v H$		58.4	298	A	[1985BAR/CAS, 1985MAJ/SVO]
C₃H₇NO	[79-16-3]	N-methylacetamide				
	$\Delta_{\text{fus}} H$		10.11	303.7		[1999AHL/LOH]
	$\Delta_{\text{fus}} H$		9.73	303.8		[1969KRE/WOO]
	$\Delta_{\text{sub}} H$		70.8 ± 2.0	298		[1996ROU/JIM2]
	$\Delta_{\text{sub}} H$		69.87 ± 0.31	298		[1984STA/WAD]
	$\Delta_{\text{sub}} H$	(288–303)	54.0			[1952AIH, 1960JON]
	$\Delta_v H$	(363–414)	55.5	378		[1995SCH/PUS]
	$\Delta_v H$	(353–428)	62	368		[1993AUC/MON]
	$\Delta_v H$	(333–443)	59.6	348	A	[1987STE/MAL]
	$\Delta_v H$	(353–479)	53.5	368	A	[1987STE/MAL]
	$\Delta_v H$		NA			[1968GOP/RIZ]
C₃H₇NO	[627-39-4]	propionaldehyde oxime				
	$\Delta_v H$	(313–339)	51.2	326	A	[1987STE/MAL]
C₃H₇NO₂	[51-79-6]	ethyl carbamate				
	$\Delta_{\text{fus}} H$		15.23	321.9		[1991ACR]
	$\Delta_{\text{sub}} H$	(256–273)	77.7	265	TE,ME	[1983DEW/VAN]
	$\Delta_{\text{sub}} H$		76.3	298		[1983DEW/VAN]
	$\Delta_{\text{sub}} H$		71.9	322		[1976BAR/BOU]
	$\Delta_{\text{sub}} H$	(292–307)	89.1 ± 0.8	299	GS	[1959DAV/JON]
	$\Delta_v H$	(323–373)	U 25.8	338		[2004AHM/GIE]
	$\Delta_v H$	(338–457)	56.6	353	A	[1987STE/MAL, 1947STU]
C₃H₇NO₂	[541-42-4]	isopropyl nitrite				
	$\Delta_v H$	(253–268)	26.0	260	A	[1987STE/MAL, 1937THO/DAI]
C₃H₇NO₂	[108-03-2]	1-nitropropane				
	$\Delta_v H$	(313–353)	43.9	298	CGC	[1995CHI/HOS]
	$\Delta_v H$	(293–405)	42.6	308	A, EB	[1987STE/MAL, 1956TOO, 1970DYK]
	$\Delta_v H$	(331–404)	40.6	346		[1949DRE/SHR, 1949DRE/MAR]
C₃H₇NO₂	[79-46-9]	2-nitropropane				
	$\Delta_v H$	(313–353)	43.9	298	CGC	[1995CHI/HOS]
	$\Delta_v H$	(284–394)	40.9	299	A, EB	[1987STE/MAL, 1947STU, 1956TOO, 1970DYK]
C₃H₇NO₂	[543-67-9]	propyl nitrite				
	$\Delta_v H$	(253–268)	28.3	260	A	[1987STE/MAL, 1937THO/DAI]
C₃H₇NO₂	[56-41-7]	L-(<i>d</i>)-alanine				
	$\Delta_{\text{sub}} H$		132.8 ± 1	414	TE,ME	[1979DEK/VOO]
	$\Delta_{\text{sub}} H$	(413–450)	132.4 ± 1.3	433	C	[1977NAG/SAB]
	$\Delta_{\text{sub}} H$		144.8 ± 4.2	298		[1977NAG/SAB]
C₃H₇NO₂	[338-69-2]	D-(<i>l</i>)-alanine				
	$\Delta_{\text{sub}} H$	(407–426)	132.8	417	A	[1987STE/MAL]
	$\Delta_{\text{sub}} H$	(342–442)	U 105 ± 8	392	LE	[1977GAF/PIE]
	$\Delta_{\text{sub}} H$	(453–469)	138.3 ± 8	461	ME	[1965SVE/CLY, 1970COX/PIL, 1964CLY/SVE, 1989CHI/GRO]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound					
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference	
C ₃ H ₇ NO ₂	[107-95-9]	β -alanine					
	$\Delta_{\text{sub}}H$	(384–402)	133.1 ± 0.7	393	C	[1983SKO/SAB]	
	$\Delta_{\text{sub}}H$		134 ± 2	298	C	[1983SKO/SAB]	
C ₃ H ₇ NO ₂	[107-97-1]	sarcosine (N-methylglycine)					
	$\Delta_{\text{sub}}H$	(380–413)	146 ± 1	298	C	[1978SAB/LAF]	
C ₃ H ₇ NO ₂ S	[52-90-4]	L-cysteine					
	$\Delta_{\text{sub}}H$	(337–437)	U 96.2 ± 4.2	387	LE	[1977GAF/PIE]	
C ₃ H ₇ NO ₃	[1712-64-7]	isopropyl nitrate					
	$\Delta_{\text{fus}}H$		10.1	190.9		[1988LUS/RUB]	
	$\Delta_{\text{v}}H$		35.3 ± 0.6		DSC	[1999JON/FEN]	
C ₃ H ₇ NO ₃	[627-13-4]	propyl nitrate					
	$\Delta_{\text{v}}H$	(273–343)	41.7	288	A	[1987STE/MAL, 1957GRA/PRA, 1970DYK]	
C ₃ H ₇ NO ₃	[302-84-1]	(<i>dl</i>)-serine					
	$\Delta_{\text{sub}}H$	(354–454)	U 83.7 ± 4	404	LE	[1977GAF/PIE]	
C ₃ H ₇ N ₃	[22293-25-0]	1-azidopropane					
	$\Delta_{\text{v}}H$	(253–298)	31.1	268	A	[1987STE/MAL, 1964GEI/KON, 1984BOU/FRI]	
C ₃ H ₇ N ₃	[691-57-6]	2-azidopropane					
	$\Delta_{\text{v}}H$	(253–298)	33.2	268	A	[1987STE/MAL, 1964GEI/KON, 1984BOU/FRI]	
C ₃ H ₇ P	[81637-99-2]	2-propenylphosphine					
	$\Delta_{\text{v}}H$	(210–273)	32.7	241		[1988SHA/DIE]	
C ₃ H ₈	[74-98-6]	propane					
	$\Delta_{\text{fus}}H$		3.51	85.5	AC	[2009PER/OCH]	
	$\Delta_{\text{fus}}H$		3.52	85.5		[1991ACR]	
	$\Delta_{\text{sub}}H$		28.5	86	B	[1963BON]	
	$\Delta_{\text{v}}H$	(278–332)	18.8	293	A	[1987STE/MAL]	
	$\Delta_{\text{v}}H$	(165–248)	19.5	233	A	[1987STE/MAL]	
	$\Delta_{\text{v}}H$	(104–165)	22.1	150	A	[1987STE/MAL]	
	$\Delta_{\text{v}}H$	(231–281)	19.0	266	A	[1987STE/MAL]	
	$\Delta_{\text{v}}H$	(329–369)	19.2	344	A	[1987STE/MAL]	
	$\Delta_{\text{v}}H$	(312–367)	18.9	327		[1980MAJ/SVA]	
C ₃ H ₈ N ₂	[na]	dimethyl ammonium cyanide					
	$\Delta_{\text{v}}H$	(251–295)	49.0	280		[1987STE/MAL, 1973DIE/MAR]	
	C ₃ H ₈ N ₂ O	[625-52-5]	N-ethylurea				
		$\Delta_{\text{fus}}H$		9.6	356.7	DSC	[2005HAS/TAJ]
		$\Delta_{\text{trs}}H$		1.0	294.6		
		$\Delta_{\text{fus}}H$		14.65	368.9	DSC	[1995FER/DEL]
		$\Delta_{\text{fus}}H$		13.9	NA	DSC	[1995STR/ARG]
		$\Delta_{\text{fus}}H$		14.39	365.1		[1990KAB/MIR2]
		$\Delta_{\text{sub}}H$	(323–364)	98.1 ± 1.1	344	ME	[2003ZAI/KAB]
		$\Delta_{\text{sub}}H$	(323–364)	97.8 ± 1.1	350	ME	[2003ZAI/KAB]
$\Delta_{\text{sub}}H$			96.4 ± 1.1	350	C	[2003ZAI/KAB]	
$\Delta_{\text{sub}}H$		(333–365)	91.8 ± 1.2	354	TE	[1990PIA/FER, 1987FER/DEL2]	

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
	$\Delta_{\text{sub}}H$		100.3 ± 0.7			[1986KRA/KOZ2]
C ₃ H ₈ N ₂ O	[598-94-7]	1,1-dimethylurea				
	$\Delta_{\text{fus}}H$		29.11	454		[1991ACR]
	$\Delta_{\text{sub}}H$	(346–398)	93.5 ± 0.3	298	GS	[2006EME/KAB]
	$\Delta_{\text{sub}}H$	(323–372)	94.7 ± 1.4	348	ME	[2003ZAI/KAB]
	$\Delta_{\text{sub}}H$	(323–372)	94.7 ± 1.4	350	ME	[2003ZAI/KAB]
	$\Delta_{\text{sub}}H$		93.3 ± 0.5	350	C	[2003ZAI/KAB]
	$\Delta_{\text{sub}}H$	(326–369)	92.5 ± 1.3	357	TE	[1990PIA/FER, 1987FER/DEL2]
	$\Delta_{\text{sub}}H$		99.1 ± 0.4			[1986KRA/KOZ2]
C ₃ H ₈ N ₂ O	[96-31-1]	1,3-dimethylurea				
	$\Delta_{\text{fus}}H$		13.0	379.5		
	$\Delta_{\text{trs}}H$		0.08	301.2		
	$\Delta_{\text{trs}}H$		0.32	161.3		[1995KAB/KOZ2, 1990KAB/MIR2]
	$\Delta_{\text{sub}}H$	(313–357)	89.3 ± 0.4	298	GS	[2006EME/KAB]
	$\Delta_{\text{sub}}H$	(317–377)	87.6 ± 1.0	347	ME	[2003ZAI/KAB]
	$\Delta_{\text{sub}}H$	(317–377)	87.5 ± 1.0	350	ME	[2003ZAI/KAB]
	$\Delta_{\text{sub}}H$		86.6 ± 0.5	350	C	[2003ZAI/KAB]
	$\Delta_{\text{sub}}H$	(316–373)	87.2 ± 0.6	353	TE	[1990PIA/FER, 1987FER/DEL2]
C ₃ H ₈ N ₂ O ₂	[4114-31-2]	ethyl carbazate				
	$\Delta_{\text{fus}}H$	(78–371)	20.0	318.9	AC	[2001DI/SUN2]
C ₃ H ₈ N ₂ S	[534-13-4]	1,3-dimethylthiourea				
	$\Delta_{\text{fus}}H$		13.71	337		[2000DEL/JOZ]
	$\Delta_{\text{fus}}H$		12.7	336.9		[1994FER/MAR]
	$\Delta_{\text{sub}}H$		111.8 ± 3	298	B	[2000DEL/JOZ]
	$\Delta_{\text{sub}}H$		107.3 ± 4.0	298	B	[1994TER/PIA]
	$\Delta_{\text{sub}}H$		108 ± 3.0	361	B	[1994FER/MAR]
	$\Delta_{\text{v}}H$	(342–375)	93 ± 4.0	359	ME, TE	[1994TER/PIA]
C ₃ H ₈ N ₂ S	[625-53-6]	1-ethylthiourea				
	$\Delta_{\text{fus}}H$		21.34	380.8	DSC	[2000DEL/JOZ]
	$\Delta_{\text{sub}}H$	(360–380)	116.1 ± 2.0	370	ME, TE	[2007FER/BAD]
	$\Delta_{\text{sub}}H$	(360–380)	118.8 ± 2.1	298	ME, TE	[2007FER/BAD]
	$\Delta_{\text{sub}}H$		118.8 ± 5	298	ME	[2000DEL/JOZ]
C ₃ H ₈ N ₄ O ₄	[13232-00-3]	2,4-dinitro-2,4-diazapentane				
	$\Delta_{\text{fus}}H$		16.36	330.2	DSC	[1997ZEM]
C ₃ H ₈ O	[540-67-0]	methyl ethyl ether				
	$\Delta_{\text{v}}H$	(281–433)	30.1	296	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$	(216–299)	37.0	231	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$	(281–438)	37.1	296	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$	(278–281)	NA			[1967SHA]
	$\Delta_{\text{v}}H$	(182–280)	26.3	265		[1947STU]
C ₃ H ₈ O	[71-23-8]	1-propanol				
	$\Delta_{\text{fus}}H$		5.4	148.7		[2004VAN/VAN]
	$\Delta_{\text{fus}}H$		5.37	148.8		[1968COU/LEE]
	$\Delta_{\text{v}}H$	(298–363)	45.7	298		[2004NAS/ZIM]
	$\Delta_{\text{v}}H$	(310–356)	47.8	298	EB	[2004LUB/MAL]
	$\Delta_{\text{v}}H$		41.2	371		[2000WOR/VIN]
	$\Delta_{\text{v}}H$		35.2	423		[2000WOR/VIN]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	Enthalpy							
		$\Delta_v H$			29.4	453		[2000WOR/VIN]
		$\Delta_v H$			21.0	498		[2000WOR/VIN]
		$\Delta_v H$			11.4	528		[2000WOR/VIN]
		$\Delta_v H$	(323–373)		49.2	298	CGC	[1995CHI/HOS]
		$\Delta_v H$	(303–370)		47.0	318		[1995AUC/GON]
		$\Delta_v H$	(360–377)		42.9	375		[1990ORT/SUS]
		$\Delta_v H$	(200–228)		48.0	214	A	[1987STE/MAL]
		$\Delta_v H$	(356–376)		43.5	366	A	[1987STE/MAL]
		$\Delta_v H$	(369–407)		42.3	384	A	[1987STE/MAL]
		$\Delta_v H$	(401–482)		40.1	416	A	[1987STE/MAL]
		$\Delta_v H$	(478–507)		36.5	492	A	[1987STE/MAL]
		$\Delta_v H$	(243–303)		46.3	298		[1983SCH/STR]
		$\Delta_v H$	(275–373)		49.3	290		[1973WIL/ZWO]
		$\Delta_v H$			46.4 ± 0.1	313	C	[1973SVO/VES]
		$\Delta_v H$			45.7 ± 0.1	323	C	[1973SVO/VES]
		$\Delta_v H$			44.9 ± 0.1	333	C	[1973SVO/VES]
		$\Delta_v H$			44.0 ± 0.1	343	C	[1973SVO/VES]
		$\Delta_v H$			43.2 ± 0.1	353	C	[1973SVO/VES]
		$\Delta_v H$			42.4 ± 0.1	363	C	[1973SVO/VES]
		$\Delta_v H$			47.49 ± 0.02	298	C	[1971POL/BEN]
		$\Delta_v H$	(333–377)		44.7	348	EB	[1970AMB/SPR, 1987STE/MAL]
		$\Delta_v H$	(292–370)		46.9	307	DTA	[1969KEM/KRE]
		$\Delta_v H$	(288–348)		46.7	303		[1967VAN/SOC]
		$\Delta_v H$			47.3 ± 0.1	298	C	[1966WAD]
		$\Delta_v H$	(338–378)		44.3	353	EB	[1963BID/COL]
		$\Delta_v H$			46.6	298	C	[1963MCC/LAI]
		$\Delta_v H$	(405–537)		40.7	420		[1963AMB/TOW]
		$\Delta_v H$	(343–385)		44.1	358		[1961MAT/MCK]
		$\Delta_v H$			43.9 ± 0.1	343	C	[1961MAT/MCK]
		$\Delta_v H$			42.3 ± 0.1	360	C	[1961MAT/MCK]
		$\Delta_v H$			41.2 ± 0.1	370	C	[1961MAT/MCK]
		$\Delta_v H$			40.3 ± 0.1	378	C	[1961MAT/MCK]
		$\Delta_v H$			39.7 ± 0.1	384	C	[1961MAT/MCK]
		$\Delta_v H$	(321–367)		45.5			[1959ARO/KAS]
		$\Delta_v H$			43.2	354		[1957WIL/HAR]
C₃H₈O	[67-63-0]		2-propanol					
		$\Delta_{\text{fus}} H$			5.41	185.2		[1996DOM/HEA]
		$\Delta_v H$	(298–353)		44.0	298		[2004NAS/ZIM]
		$\Delta_v H$	(322–355)		43.2	337		[2002SEG/GAL]
		$\Delta_v H$			39.8	355		[2000WOR/VIN2]
		$\Delta_v H$			29.7	423		[2000WOR/VIN2]
		$\Delta_v H$			23.7	453		[2000WOR/VIN2]
		$\Delta_v H$			16.5	483		[2000WOR/VIN2]
		$\Delta_v H$			10.5	503		[2000WOR/VIN2]
		$\Delta_v H$			40.4			[1999FAT]
		$\Delta_v H$	(300–355)		44.8	315		[1995AUC/GON]
		$\Delta_v H$	(195–228)		50.3	213	A	[1987STE/MAL]
		$\Delta_v H$	(347–368)		42.0	355	A	[1987STE/MAL]
		$\Delta_v H$	(350–383)		41.3	365	A	[1987STE/MAL]
		$\Delta_v H$	(379–461)		39.2	394	A	[1987STE/MAL]
		$\Delta_v H$	(453–508)		35.3	468	A	[1987STE/MAL]
		$\Delta_v H$	(273–374)		45.7	288		[1973WIL/ZWO]
		$\Delta_v H$	(325–362)		43.1	340	A, EB	[1987STE/MAL, 1970AMB/SPR]
		$\Delta_v H$			45.34 ± 0.02	298	C	[1971POL/BEN]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(288–348)	45.5	303		[1967VAN/SOC]
	$\Delta_v H$		45.2 ± 0.1	298	C	[1966WAD]
	$\Delta_v H$		42.7 ± 0.1	330	C	[1964BER/LAR]
	$\Delta_v H$		41.0 ± 0.1	346	C	[1964BER/LAR]
	$\Delta_v H$		39.8 ± 0.1	355	C	[1964BER/LAR]
	$\Delta_v H$		38.9 ± 0.1	363	C	[1964BER/LAR]
	$\Delta_v H$	(329–363)	42.8	344	EB	[1963BID/COL]
	$\Delta_v H$		44.0	298	C	[1963MCC/LAI]
	$\Delta_v H$	(395–508)	39.1	410		[1963AMB/TOW]
	$\Delta_v H$		43.2	324	C	[1963HAL/COX]
	$\Delta_v H$		41.7	339	C	[1963HAL/COX]
	$\Delta_v H$		39.8	355	C	[1963HAL/COX]
	$\Delta_v H$	(354–420)	41.1	369		[1955FOZ/MOR]
	$\Delta_v H$	(273–363)	44.4	298		[1928PAR/BAR]
C ₃ H ₈ OS ₂	[59-52-9]	2,3-dimercaptopropanol				
	$\Delta_v H$	(353–413)	61.2	382		[1999DYK/SVO, 1987STE/MAL]
C ₃ H ₈ O ₂	[109-86-4]	2-methoxyethanol				
	$\Delta_v H$	(346–397)	43.0	361		[2010MAR/LOR]
	$\Delta_v H$	(333–423)	42.8	348	A	[1987STE/MAL]
	$\Delta_v H$		45.2 ± 0.2	298	C	[1971KUS/WAD]
	$\Delta_v H$	(329–396)	42.9	344		[1956PIC/FRI]
C ₃ H ₈ O ₂	[109-87-5]	dimethoxymethane				
	$\Delta_{\text{fus}} H$		8.33	168		[1996DOM/HEA]
	$\Delta_v H$	(273–357)	29.6	298		[2001ALB/HAH]
	$\Delta_v H$	(273–316)	31.2	288	A	[1987STE/MAL]
	$\Delta_v H$	(273–318)	29.8	288	A	[1987STE/MAL]
	$\Delta_v H$	(296–314)	30.3	305		[1976BRA/PES]
	$\Delta_v H$	(273–308)	30.1	288		[1949NIC/LAF]
C ₃ H ₈ O ₂	[57-55-6]	1,2-propanediol (propylene glycol)				
	$\Delta_v H$	(284–331)	67.5 ± 0.5	298	GS	[2009VER/KOZ]
	$\Delta_v H$	(284–331)	64.5 ± 0.2	298	GS	[2004VER2]
		Note: Author later recalculated this earlier value in a later paper [2009VER/KOZ]. The recalculated value was reported to be 68.5 ± 0.5				
	$\Delta_v H$	(293–423)	76	298	EB	[2004CHY/FRA2]
	$\Delta_v H$	(366–396)	62.2	298	TGA	[2002TAT/DOL]
	$\Delta_v H$	(365–496)	60.0 ± 0.3	380	EB	[2002STE/CHI3]
	$\Delta_v H$	(365–496)	56.2 ± 0.2	420	EB	[2002STE/CHI3]
	$\Delta_v H$	(365–496)	52.0 ± 0.3	460	EB	[2002STE/CHI3]
	$\Delta_v H$	(365–496)	47.5 ± 0.6	500	EB	[2002STE/CHI3]
	$\Delta_v H$	(348–453)	63.6 ± 0.3	298	EB	[1990KNA/SAB3, 2004VER2]
	$\Delta_v H$		U71.2 ± 0.1	298	C	[1990KNA/SAB3, 2004VER2]
	$\Delta_v H$	(373–408)	66.5	413	TGA	[1987ALN/ALS]
	$\Delta_v H$		U 51.7	298	I	[1971SUN/EIS]
	$\Delta_v H$	(359–461)	64.7	298	EB	[1966THO/MEA, 2004VER2]
	$\Delta_v H$	(318–461)	58.6	333	A	[1987STE/MAL, 1947STU]
	$\Delta_v H$	(353–403)	58.2	378		[1935SCH/STA]
	$\Delta_v H$	(403–460)	56	431		[1935SCH/STA]
C ₃ H ₈ O ₂	[4254-14-2]	(R)-1,2-propanediol (propylene glycol)				
	$\Delta_{\text{fus}} H$		8.4	240		[1995JAB/LET]
C ₃ H ₈ O ₂	[4254-15-3]	(S)-1,2-propanediol (propylene glycol)				
	$\Delta_{\text{fus}} H$		8.4	240		[1995JAB/LET]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(294–338)	69.2 ± 0.3	298	GS	[2009VER/KOZ]
C ₃ H ₈ O ₂	[504-63-2]	1,3-propanediol				
	$\Delta_{\text{fus}} H$		11.4	249	DSC	[1998JAB/LET]
	$\Delta_v H$	(314–460)	66.5	298	EB	[2008RIB/SAN4]
	$\Delta_v H$	(293–342)	70.5 ± 0.2	298	GS	[2007VER]
	$\Delta_v H$	(480–716)	69.1 ± 0.2	298		[2002WIL/VON, 2007VER]
	$\Delta_v H$	(413–458)	70.6 ± 0.5	298	EB	[1996OLS, 2007VER]
	$\Delta_v H$	(402–488)	70.1 ± 0.3	298	EB	[1981MAR/SAC, 2007VER]
	$\Delta_v H$	(367–489)	71.4	298	EB	[1966THO/MEA, 2007VER]
	$\Delta_v H$		72.4 ± 0.3	298	C	[1988KNA/SAB, 1990KNA/SAB2]
	$\Delta_v H$	(332–448)	57.2	347	A	[1987STE/MAL, 1947STU]
	$\Delta_v H$	(373–488)	65.2	298	EB	[1937GAL/HIB, 2007VER]
	$\Delta_v H$	(383–433)	63.3	408		[1935SCH/STA]
	$\Delta_v H$	(433–488)	60.4	460		[1935SCH/STA]
C ₃ H ₈ O ₂ S	[594-43-4]	ethyl methyl sulfone				
	$\Delta_{\text{fus}} H$		11.3	307.7		[1961BUS/IVI]
	$\Delta_{\text{sub}} H$		77.8 ± 2.9			[UR/MAC, 1970COX/PIL]
C ₃ H ₈ O ₃	[56-81-5]	glycerol				
	$\Delta_{\text{fus}} H$		18.28	293		[1991ACR]
	$\Delta_v H$		91.7 ± 0.9	298	C	[1988BAS/NIL]
	$\Delta_v H$	(469–563)	78.5	484	A	[1987STE/MAL]
	$\Delta_v H$	(291–341)	86.8	316	ME	[1977CAM/SCHL]
	$\Delta_v H$		67.5	343	GC	[1977NOV/NOV]
	$\Delta_v H$		66.8	353	GC	[1977NOV/NOV]
	$\Delta_v H$		66.2	363	GC	[1977NOV/NOV]
	$\Delta_v H$		65.5	373	GC	[1977NOV/NOV]
	$\Delta_v H$		64.8	383	GC	[1977NOV/NOV]
	$\Delta_v H$	(278–323)	71.5	300		[1972MCF/SOM]
	$\Delta_v H$	(293–343)	85.8	308	ME	[1987STE/MAL, 1962ROS/HEI, 1970DYK]
$\Delta_v H$	(456–553)	86	471		[1886RIC]	
C ₃ H ₈ S	[624-89-5]	ethyl methyl sulfide				
	$\Delta_{\text{fus}} H$		9.76	167.2		[1996DOM/HEA]
	$\Delta_v H$	(253–363)	33.7	268		[1999DYK/SVO]
	$\Delta_v H$		31.5	298		[1981SHI/SAI]
	$\Delta_v H$		32.3	287		[1960MAC/MAY]
	$\Delta_v H$	(296–373)	31.8	311	A, EB	[1987STE/MAL, 1951SCO/FIN]
	$\Delta_v H$		31.8	298		[1971WIL/ZWO, 1966OSB/DOU, 1954HUB/WAD]
$\Delta_v H$		30.3	338		[1935THO/LIN]	
C ₃ H ₈ S	[107-03-9]	1-propanethiol				
	$\Delta_{\text{trs}} H$		3.97	142.1		
	$\Delta_{\text{fus}} H$		5.48	160		[1996DOM/HEA]
	$\Delta_v H$	(254–364)	33.7	269		[1999DYK/SVO]
	$\Delta_v H$		31.9	298		[1971WIL/ZWO]
	$\Delta_v H$	(297–375)	31.8	312	A, EB	[1987STE/MAL, 1956PEN/SCO, 1966OSB/DOU, 1954HUB/WAD]
	$\Delta_v H$		31.6 ± 0.1	303	C	[1956PEN/SCO]
	$\Delta_v H$		30.7 ± 0.1	320	C	[1956PEN/SCO]
	$\Delta_v H$		29.5 ± 0.1	341	C	[1956PEN/SCO]
	$\Delta_v H$	(284–340)	31.5	312		[1933TAY/LAY]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₃ H ₈ S	[75-33-2]	2-propanethiol				
	$\Delta_{\text{trs}}H$		0.05	112.5		
	$\Delta_{\text{fus}}H$		5.73	142.6		[1996DOM/HEA]
	Δ_vH	(242–348)	31.9	257		[1999DYK/SVO]
	Δ_vH	(283–358)	29.5	298		[1971WIL/ZWO]
			30.1	298	A, EB	[1987STE/MAL, 1954MCC/SCO, 1966OSB/DOU, 1954HUB/WAD]
C ₃ H ₈ S ₂	[109-80-8]	1,3-propanedithiol				
	Δ_vH	(338–446)	50.9	353		[1999DYK/SVO]
	Δ_vH	(377–446)	41.6	398	A	[1987STE/MAL]
			49.7	298		[1962MAN/SUN]
C ₃ H ₉ N	[75-31-0]	isopropylamine				
	$\Delta_{\text{fus}}H$		7.33	178		[1991ACR]
	Δ_vH		28.4	298		[1979MAJ/SVO2]
	Δ_vH		27.2	313		[1979MAJ/SVO2]
		(277–334)	29.7	292	A, EB, IP	[1987STE/MAL, 1968OSB/DOU, 1970DYK]
C ₃ H ₉ N	[107-10-8]	propylamine				
	$\Delta_{\text{fus}}H$		10.97	188.4		[1991ACR]
	Δ_vH		31.3	298		[1979MAJ/SVO2]
	Δ_vH		30.1	313		[1979MAJ/SVO2]
	Δ_vH		28.9	328		[1979MAJ/SVO2]
		(296–350)	31.3	311	A, EB, IP	[1987STE/MAL, 1968OSB/DOU, 1970DYK]
C ₃ H ₉ N	[75-50-3]	trimethylamine				
	$\Delta_{\text{fus}}H$		6.54	156.1		[1991ACR]
	Δ_vH	(333–403)	23.0	368		[1950DAY/FEL]
	Δ_vH	(273–313)	24.1	288		[1945SWI/HOC]
	Δ_vH	(193–276)	24.6	261	A	[1987STE/MAL, 1944AST/SAG]
			24.5	250	C	[1944AST/SAG]
C ₃ H ₉ NO	[109-83-1]	2-(methylamino)ethanol				
	Δ_vH	(275–320)	57.8 ± 0.2	298	GS	[2005KAP/SLO]
	Δ_vH	(269–401)	57.9	298		[1998NOL/VAL, 2005KAP/SLO]
	Δ_vH	(340–461)	57.0 ± 0.5	298	EB	[1997STE/CHI3]
	Δ_vH	(351–410)	57.6	298	EB	[1987SMI/TER, 2005KAP/SLO]
	Δ_vH	(298–308)	57.5	298		[1982TOU/OKA, 2005KAP/SLO]
C ₃ H ₉ NO	[78-96-6]	1-amino-2-propanol				
	Δ_vH	(306–431)	51.6	321	A	[1987STE/MAL]
C ₃ H ₉ NO	[156-87-6]	3-amino-1-propanol				
	$\Delta_{\text{fus}}H$		16.9	284.1		[2003CAC/BAU]
C ₃ H ₉ NO	[5669-39-6]	N-methoxy dimethyl amine (trimethylhydroxylamine)				
	Δ_vH	(194–297)	28	282	A	[1987STE/MAL, 1957BIS/PAR]
C ₃ H ₉ NO	[109-85-3]	2-methoxyethyl amine				
	Δ_vH	(278–318)	38.8	293	A	[1987STE/MAL]
C ₃ H ₉ NO ₂ S	[177634-55-8]	trimethyl amine. sulfur dioxide complex				
	$\Delta_{\text{sub}}H$	(292–349)	60.6	307		[1987STE/MAL, 1943BUR2]
C ₃ H ₉ OP	[676-96-0]	trimethyl amine. sulfur dioxide complex				
	$\Delta_{\text{sub}}H$		50.2 ± 4.2		E	[1982PIL/SKI, 1960CLA/FOW]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₃ H ₉ O ₃ P	[756-79-6]	methylphosphonic acid, dimethyl ester				
	$\Delta_v H$	(258–454)	54.9	273	GS	[2009BUT/BUC]
	$\Delta_v H$	(258–454)	52.8	298	GS	[2009BUT/BUC]
	$\Delta_v H$	(258–454)	51.8	313	GS	[2009BUT/BUC]
	$\Delta_v H$	(258–454)	50.6	333	GS	[2009BUT/BUC]
	$\Delta_v H$	(258–454)	49.5	353	GS	[2009BUT/BUC]
C ₃ H ₉ O ₃ P	[121-45-9]	trimethyl phosphite				
	$\Delta_v H$	(302–342)	42.5	317	EB	[1990DUT/KAH]
C ₃ H ₉ O ₄ P	[512-56-1]	trimethyl phosphate				
	$\Delta_v H$	(408–438)	47.5	298	CGC	[2007PAN/ANT2]
C ₃ H ₉ P	[594-09-2]	trimethyl phosphine				
	$\Delta_v H$	(248–310)	28.9	263	A	[1987STE/MAL]
C ₃ H ₉ PS	[2404-55-9]	trimethylphosphine sulfide				
	$\Delta_{\text{sub}} H$	(366–394)	70.3	380		[1966BUR]
C ₃ H ₁₀ N ₂	[109-76-2]	1,3-diaminopropane				
	$\Delta_{\text{trs}} H$		10.53	260.6		
	$\Delta_{\text{fus}} H$		12.19	262.4	DSC	[2002DAL/DEL]
C ₃ H ₁₀ N ₂	[78-90-0]	<i>dl</i> 1,2-propanediamine				
	$\Delta_{\text{trs}} H$		0.07	222		
	$\Delta_{\text{fus}} H$		18.42	236.5		[1996DOM/HEA]
C ₃ H ₁₀ N ₂	[1741-01-1]	trimethylhydrazine				
	$\Delta_v H$	(257–287)	34.6	272		[1955AST/ZOL]
	$\Delta_v H$		33.4 ± 0.1	292	C	[1955AST/ZOL]
C ₃ N ₂ O	[1115-12-4]	carbonyl cyanide				
	$\Delta_v H$	(250–291)	37.5	276	A	[1987STE/MAL, 1948GLE/HAU]
C ₃ O ₂	[504-64-3]	carbon suboxide				
	$\Delta_v H$	(161–249)	26.2	234	A	[1987STE/MAL, 1965MCD/KIL]
C ₃ S ₂	[627-34-9]	carbon subsulfide				
	$\Delta_v H$	(287–383)	45.1	302	A	[1987STE/MAL]
C ₄ BrClF ₉ N	[na]	1,1,2-trifluoro-2-chloro-2-bromo-N,N-bis(trifluoromethyl)ethylamine				
	$\Delta_v H$	(329–364)	33.1	344	A	[1987STE/MAL, 1965HAS/TIP]
C ₄ BrCl ₂ F ₈ N	[4905-98-0]	2-bromo-1,2-dichloro-1,2-difluoro-N,N-bis(trifluoromethyl)ethamine				
	$\Delta_v H$	(358–394)	36.5	376		[1965HAS/TIP]
C ₄ BrF ₆ N	[22130-38-7]	2-bromo-N,N-bis(trifluoromethyl)ethynylamine				
	$\Delta_v H$	(311–329)	30.4	320	A	[1987STE/MAL, 1969FRE/TIP]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₄ BrF ₈ N	[17725-57-4] $\Delta_v H$	N,N-bis(trifluoromethyl)-2,2-difluoro-1-bromovinylamine (293–320)	31.2	320	A	[1987STE/MAL, 1968HAS/TIP]
C ₄ BrF ₉ O ₃ S	[14609-27-9] $\Delta_v H$	2-bromo-3-fluorosulfatooctafluorobutane (313–372)	43.5	342		[1966EAR/HIL]
C ₄ BrF ₁₀ N	[2261-32-7] $\Delta_v H$	1,1,2,2-tetrafluoro-2-bromo-N,N-bis(trifluoromethyl)ethylamine (289–329)	30.4	304	A	[1987STE/MAL, 1965HAS/TIP]
C ₄ Br ₂ Cl ₂ F ₆	[375-42-8] $\Delta_v H$ $\Delta_v H$	1,4-dibromo-2,3-dichloro-1,1,2,3,4,4-hexafluorobutane	47.7 ± 0.1	308	C	[1992SVO/KUB2]
			46.9 ± 0.1	315	C	[1992SVO/KUB2]
C ₄ Br ₂ F ₉ N	[17725-58-5] $\Delta_v H$	1,2-dibromo-1,2,2-trifluoro-N,N-bis(trifluoromethyl)ethylamine (326–366)	34.3	341	A	[1987STE/MAL, 1968HAS/TIP]
C ₄ ClF ₈ N	[14003-64-6] $\Delta_v H$	2-chloro-1,2-difluoro-N,N-bis(trifluoromethyl)vinylamine (273–312)	29.1	288	A	[1987STE/MAL, 1968HAS/TIP]
C ₄ ClF ₁₀ N	[54566-79-9] $\Delta_v H$	N-chloro-1,1,2,2,2-pentafluoro-N-(pentafluoroethyl)ethanamine	27.2	325		[1975PET/SHR2]
C ₄ ClF ₁₀ N	[53684-04-1] $\Delta_v H$	N-chloro-1,1,1,2,3,3,3-heptafluoro-N-(trifluoromethyl)-2-propanamine	28.9	325		[1975KIR/LAS]
C ₄ ClF ₁₂ NS	[62609-69-2] $\Delta_v H$	chlorodifluoro[1,1,1,2,3,3,3-heptafluoro-2-propanaminato(2-)]-(trifluoromethyl) sulfur	37.7	402		[1977KIT/SHR2]
C ₄ Cl ₂ F ₆	[20972-44-5] $\Delta_v H$	1,4-dichloro-hexafluoro-2-butene (279–330)	34	294	A	[1987STE/MAL]
C ₄ Cl ₂ F ₆	[2418-22-6] $\Delta_v H$	<i>cis</i> 2,3-dichloro-hexafluoro-2-butene (298–341)	32.5	313	A	[1987STE/MAL]
C ₄ Cl ₂ F ₆	[2418-21-5] $\Delta_v H$	<i>trans</i> 2,3-dichloro-hexafluoro-2-butene (298–340)	32.2	313	A	[1987STE/MAL]
C ₄ Cl ₂ F ₇ N	[89033-96-5] $\Delta_v H$	2,3,4,4-tetrafluoro-2,3-dichloro-(trifluoromethyl)azetidene (273–333)	32.6	288	A	[1987STE/MAL, 1965BAN/BAR]
C ₄ Cl ₂ F ₇ N	[4776-86-7] $\Delta_v H$	2,3-dichlorotetrafluoropropylidene-trifluoromethylamine (283–343)	27.0	313		[1965BAN/BAR]
C ₄ Cl ₃ F ₇	[335-44-4] $\Delta_v H$ $\Delta_v H$ $\Delta_v H$	2,3,3-trichloroheptafluorobutane (302–446)	33.3			[1959YEN/REE]
			35.6	298		[1959YEN/REE]
			36.4	317	MM, A	[1987STE/MAL, 56CAP/JAC]
C ₄ Cl ₄ F ₄	[457-20-0] $\Delta_v H$	1,2,3,4-tetrachlorotetrafluoro-1-butene (362–414)	39.4	377	A	[1987STE/MAL]
C ₄ Cl ₄ F ₆ O	[61136-57-0] $\Delta_v H$ $\Delta_v H$	trichloromethyl 2-chloro-1,1,2,3,3,3-hexafluoropropyl ether (325–403)	40.3	340	A	[1987STE/MAL]
			42.8 ± 0.7	298	EB	[1976AMM/BUL]
C ₄ Cl ₄ N ₂	[1780-40-1] $\Delta_{\text{sub}} H$	2,4,5,6-tetrachloropyrimidine	83.0 ± 1.0	298	C	[2007RIB/AMA]
C ₄ Cl ₆	[87-68-3] $\Delta_v H$ $\Delta_v H$	perchloro-1,3-butadiene (343–484) (343–473)	58.6	358	A	[1987STE/MAL]
			60.4	358		[1971GEL/SIM, 1984BOU/FRI]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₄ Cl ₆ O ₃	[4124-31-6] $\Delta_v H$	trichloroacetic anhydride (329–496)	56.0	344	A	[1987STE/MAL, 1947STU]
C ₄ F ₆	[685-63-2] $\Delta_v H$	hexafluoro-1,3-butadiene (273–343)	25.9	288		[2002BOB/FED]
C ₄ F ₆ N ₂ S	[71148-78-2] $\Delta_{\text{fus}} H$	3,4-bis(trifluoromethyl)-1,2,5-thiadiazole 11.5	284			[2002BRO/DU]
C ₄ F ₆ N ₂ S	[na] $\Delta_{\text{sub}} H$	bis(trifluoromethyl)-1,3,2-dithiazol-2-yl (253–283)	49.0 ± 1.5	268	PG	[2000BRO/DU]
C ₄ F ₆ O ₃	[407-25-0] $\Delta_v H$	trifluoroacetic anhydride (271–312)	34.7	286	A	[1987STE/MAL, 1962KRE, 1971DYK]
C ₄ F ₇ NO	[4222-29-1] $\Delta_v H$	4,4-difluoro-3-(difluoromethylene)-2-(trifluoromethyl)-1,2-oxazetidine (238–283)	31.1	268	A,I	[1987STE/MAL, 1960GRI/HAZ]
C ₄ F ₇ NO	[4777-13-3] $\Delta_v H$	3,6-dihydro-2,2,3,3,5,6,6-heptafluoro 2 <i>H</i> -1,4-oxazine (249–293)	27.3	278	A	[1987STE/MAL]
C ₄ F ₇ NO ₃ S	[26404-53-5] $\Delta_v H$	fluorosulfuric ester 3,3,3-trifluoro-2-(trifluoromethyl)lactonitrile (262–320)	31.2	277	A	[1987STE/MAL]
C ₄ F ₈	[357-26-6] $\Delta_v H$ $\Delta_v H$	perfluoro-1-butene (203–279) (250–293)	28.9 U 14.4	264 265	A	[1987STE/MAL, 1971DYK] [1947FOW/HAM, 1984BOU/FRI]
C ₄ F ₈	[115-25-3] $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$	perfluorocyclobutane (289–348) (343–388) (233–388) (234–269) (233–274)	23.5 23.2 25 25 24.9	304 358 248 254 259	A A A	[1987STE/MAL] [1987STE/MAL] [1967KLE/PET] [1962MAR, 1984BOU/FRI] [1987STE/MAL, 1954FUR/MCC]
C ₄ F ₈ N ₂ O ₂ S	[66918-60-3] $\Delta_v H$	N,N'-bis(trifluoroacetyl)sulfur difluorodiimide (328–383)	43.5	355		[1978STA/MEW]
C ₄ F ₈ N ₂ O ₃	[382-38-7] $\Delta_v H$	perfluoro-2-(tetrafluoro-2-nitroethyl)-1,2-oxazetidine (273–343)	31	288	A	[1987STE/MAL]
C ₄ F ₈ OS	[42060-62-8] $\Delta_v H$	perfluorotetramethylene sulfoxide 37.1				[1973ABE/SHR]
C ₄ F ₈ O ₂ S	[42060-64-0] $\Delta_v H$	perfluorotetramethylene sulfone 31.1				[1973ABE/SHR]
C ₄ F ₈ O ₄ S	[6069-35-8] $\Delta_v H$	heptafluorobutyric acid and fluorosulfuric acid anhydride (268–352)	44.8	283	A	[1987STE/MAL, 1966DES/CAD]
C ₄ F ₈ S	[706-76-3] $\Delta_{\text{trs}} H$ $\Delta_{\text{fus}} H$ $\Delta_v H$	octafluorotetramethylene sulfide 10.88 2.09 26.9	146 266.7			[1961VAN] [1973ABE/SHR]
C ₄ F ₈ S ₂	[710-65-6] $\Delta_v H$	perfluoro-1,4-dithiane 33				[1973ABE/SHR]
C ₄ F ₉ N	[453-22-5] $\Delta_v H$	1,1,1-trifluoro-N-[2,2,2-trifluoro-1-(trifluoromethyl)ethylidene]-methanamine 22.2	288			[1975KIR/LAS]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₄ F ₉ N	[13821-49-3] $\Delta_v H$	perfluoro[N,N-dimethyl(vinylamine)] (257–280)	27.5	268	A	[1987STE/MAL, 1968HAS/TIP]
C ₄ F ₉ N	[680-23-9] $\Delta_v H$	perfluoro[N-methyl(propylidineamine)] (245–280)	26.6	265	A	[1987STE/MAL, 1968HAS/TIP]
C ₄ F ₉ N	[378-00-7] $\Delta_v H$	perfluoro[N-propyl(methylenamine)] (250–291)	28.3	276	A	[1987STE/MAL, 1956BAR/HAS]
C ₄ F ₉ NO	[32822-51-8] $\Delta_v H$	nonafluorobutyramide 29.7	306	HG	[1971DEM/SHR]	
C ₄ F ₉ NO	[na] $\Delta_v H$	2,2,4,4,5,5-hexafluoro-3-(trifluoromethyl)oxazolidine (253–293)	27.4	278	A	[1987STE/MAL]
C ₄ F ₉ NO	[714-52-3] $\Delta_v H$	perfluoro[2,4-bis(trifluoromethyl)-1,2-oxazetidine] (266–289)	25.9	278	A	[1987STE/MAL, 1961BAR/HAS]
C ₄ F ₉ NOS	[31340-35-9] $\Delta_v H$	1,1,1-trifluoro-N-[2,2,2-trifluoro-1-(trifluoromethyl)ethylidene]-methanesulfinamide 36.4	361	I	[1972SWI/BAB]	
C ₄ F ₉ NO ₂	[15496-02-3] $\Delta_v H$	O-(trifluoroacetyl)-N,N-bis(trifluoromethyl)hydroxylamine (234–296)	30.5	281	A	[1987STE/MAL]
C ₄ F ₉ NO ₂ S	[34556-29-1] $\Delta_v H$	N-(trifluoroacetyl)-S,S-bis(trifluoromethyl)sulfoximine 35.1	363	I	[1972SAU/SHR]	
C ₄ F ₉ NO ₃	[55064-78-3] $\Delta_v H$	2,2,2-trifluoro-1,1-bis(trifluoromethyl) nitrate 33.5			[1975WAL/DES]	
C ₄ F ₉ NS	[31340-34-8] $\Delta_v H$	1,1,1-trifluoro-N-[2,2,2-trifluoro-1-(trifluoromethyl)ethylidene]-methanesulfenamide 31.4	324	I	[1972SWI/BAB]	
C ₄ F ₁₀	[355-25-9] $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$	perfluorobutane (272–327) (233–273) (323–386) (233–383) (233–383) (233–260)	24.2 24.2 23.1 21 17.1 25.8	287 258 338 293 333 247	A A A	[1987STE/MAL] [1987STE/MAL] [1987STE/MAL] [1958BRO/MEA] [1958BRO/MEA] [1952SIM/HAU, 1984BOU/FRI]
C ₄ F ₁₀ OS	[33622-18-3] $\Delta_v H$	heptafluoropropyl trifluoromethyl sulfoxide 33.6			[1971SAU/SHR]	
C ₄ F ₁₀ OS	[33622-19-4] $\Delta_v H$	bis(pentafluoroethyl) sulfoxide 35.1			[1971SAU/SHR]	
C ₄ F ₁₀ O ₃ S	[5762-52-7] $\Delta_v H$	fluorosulfuric acid, perfluoro(1-methylpropyl) ester (294–342)	33.8	309	A	[1987STE/MAL]
C ₄ F ₁₀ O ₄ S	[55064-77-2] $\Delta_v H$	fluoroperoxymonosulfuric acid, 2,2,2-trifluoro-1,1-bis(trifluoromethyl)-ethyl ester 37.6			[1975WAL/DES]	
C ₄ F ₁₀ O ₆ S ₂	[2261-44-1] $\Delta_v H$ $\Delta_v H$	1,1,1,2,3,4,4,4-octafluoro-2,3-bis(fluorosulfato)butane (316–393) (392–411)	30.1 27.1	331 401	A A	[1987STE/MAL, 1964RAT/SHR] [1987STE/MAL, 1964RAT/SHR]
C ₄ F ₁₀ S	[42060-60-6] $\Delta_v H$	octafluoroetramethylene sulfur difluoride 41.5			[1973ABE/SHR]	
C ₄ F ₁₀ S	[33547-11-4] $\Delta_v H$	heptafluoropropyl trifluoromethyl sulfide 27.7			[1971SAU/SHR]	

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₄ F ₁₁ NOS	[62609-62-5] $\Delta_v H$	difluoro(1,1,1,3,3,3-hexafluoro-2-propaniminato)oxo(trifluoromethyl) sulfur 35.1	396	I	[1977KIT/SHR]	
C ₄ F ₁₁ NS	[37826-43-0] $\Delta_v H$	fluoro(trifluoromethyl)[2,2,2,1-tetrafluoro-1-(trifluoromethyl)ethyl]-imino sulfur (300–333)	32.4	315	A, I	[1987STE/MAL, 1972SWI/SHR]
C ₄ F ₁₂ N ₂ O	[10405-32-0] $\Delta_v H$	perfluoro(2,3-dimethyl)-4-oxo-diazepentane (276–308)	32	291	A	[1987STE/MAL, 1971DYK, 1966HAS/TIP]
C ₄ F ₁₂ N ₂ O	[6141-72-6] $\Delta_v H$	perfluoro(2,4-dimethyl)-3-oxo-diazepentane (288–318)	30.1	303	A	[1987STE/MAL, 1971DYK, 1966HAS/TIP]
C ₄ F ₁₂ N ₂ S	[4101-59-1] $\Delta_v H$	difluorobis[1,1,2,2,2-pentafluorothananaminato]sulfur 37.0				[1976STA/MEW]
C ₄ F ₁₂ OS	[33564-25-9] $\Delta_v H$	difluorobis[1,1,2,2,2-pentafluorothananaminato]sulfur (284–341)	33.8	299		[1999DYK/SVO]
C ₄ F ₁₂ O ₂ S	[63465-11-2] $\Delta_v H$	bis(trifluoromethyl)bis(trifluoromethoxy) sulfur (273–325)	29.3	288	A, I	[1987STE/MAL, 1977KIT/SHR3, 1978KIT/SHR]
C ₄ F ₁₂ O ₃ S	[66632-46-0] $\Delta_v H$	oxobis(trifluoromethyl)bis(trifluoromethoxy) sulfur (273–335)	33.4	288	A, I	[1987STE/MAL, 1978KIT/SHR]
C ₄ F ₁₂ P ₂ S	[1486-20-0] $\Delta_v H$	di[bis(trifluoromethyl)phosphino] sulfide (273–335)	42.2	304	T	[1964CAV/EME]
C ₄ F ₁₂ P ₄	[393-02-2] $\Delta_{\text{sub}} H$	1,2,3,4-tetrakis(trifluoromethyl)tetraphosphetane (292–339)	65.3	307	A, SG	[1987STE/MAL, 1958MAH/BUR]
	$\Delta_v H$	(313–375)	43.2	328	A, SG	[1987STE/MAL, 1958MAH/BUR]
C ₄ F ₁₂ S	[33622-15-0] $\Delta_v H$	difluorobis(pentafluoroethyl) sulfur (284–341)	34.0	299	A	[1987STE/MAL]
	$\Delta_v H$		32.2			[1971SAU/SHR]
C ₄ F ₁₂ S	[31206-31-2] $\Delta_v H$	difluoro(heptafluoropropyl) (trifluoromethyl) sulfur 32.8				[1971SAU/SHR]
C ₄ F ₁₃ NOS	[65844-09-9] $\Delta_v H$	trifluoro[1,1,1,2,3,3,3-heptafluoro-2-propanaminoato(2-)]-(trifluoromethanolato) sulfur 33.9	389	I	[1978KIT/SHR]	
C ₄ F ₁₅ N ₂ O ₂ P	[36544-20-4] $\Delta_v H$	phosphorous bis[bis(trifluoromethyl)nitroxide] difluoride (303–370)	37.6	336		[1973WAN/SHR]
C ₄ F ₁₆ S ₂	[4556-31-4] $\Delta_v H$	hexadecafluoro-octahydro-1,4-dithiane (323–408)	40.5	365		[1999DYK/SVO, 1973ABE/SHR]
C ₄ N ₄	[1071-98-3] $\Delta_{\text{sub}} H$	dicyanoacetylene (263–273)	44.3	268	I	[1957SAG, 1975GRO, 1987STE/MAL]
	$\Delta_v H$	(295–350)	27.3	310	A	[1987STE/MAL]
C ₄ HBrF ₇ N	[25273-49-8] $\Delta_v H$	1-bromo-2-fluoro-N,N-bis(trifluoromethyl)vinylamine (321–342)	29.8	331	A	[1987STE/MAL]
C ₄ HBrF ₉ N	[4905-99-0] $\Delta_v H$	2-bromo-1,1,2-trifluoro-N,N-bis(trifluoromethyl)ethylamine (308–342)	31.9	323	A	[1987STE/MAL, 1965HAS/TIP]
C ₄ HBrF ₉ N	[4905-96-8] $\Delta_v H$	2-bromo-1,2,2-trifluoro-N,N-bis(trifluoromethyl)ethylamine (301–332)	33.8	316	A	[1987STE/MAL, 1968HAS/TIP]
C ₄ HBr ₂ F ₆ N	[22298-34-6]	<i>trans</i> 1,2-dibromo-N,N-bis(trifluoromethyl)vinylamine				

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(355–382)	33.4	369	A	[1987STE/MAL, 1969FRE/TIP]
C ₄ HCl ₂ F ₅ O ₂	[375-07-5]	3,4-dichloro-2,2,3,4,4-pentafluorobutyric acid				
	$\Delta_v H$	(373–456)	54.8	388	A	[1987STE/MAL, 1957BAR/SEF]
C ₄ HCl ₃ N ₂	[3764-01-0]	2,4,6-trichloropyrimidine				
	$\Delta_v H$		55.6 ± 0.6	298	C	[2007RIB/AMA]
C ₄ HF ₅	[7096-51-7]	3,3,4,4,4-pentafluoro-1-butyne				
	$\Delta_v H$	(203–261)	23.6	246	A	[1987STE/MAL]
C ₄ HF ₆ N	[13747-21-2]	N,N-bis(trifluoromethyl)ethynylamine				
	$\Delta_v H$	(229–271)	26.0	256	A	[1987STE/MAL, 1968FRE/TIP]
C ₄ HF ₆ N ₃	[709-62-6]	3,5-bis(trifluoromethyl)-1,2,4-triazole				
	$\Delta_{\text{sub}} H$	(271–283)	75.6 ± 0.8	277	ME	[1994TIP/JIM]
	$\Delta_{\text{sub}} H$	(271–283)	74.7 ± 0.8	298	ME	[1994TIP/JIM]
C ₄ HF ₇ O ₂	[375-22-4]	perfluorobutyric acid				
	$\Delta_v H$	(310–426)	50.1 ± 0.2	320	EB	[2002STE/CHI5]
	$\Delta_v H$	(310–426)	45.9 ± 0.2	360	EB	[2002STE/CHI5]
	$\Delta_v H$	(310–426)	41.0 ± 0.5	400	EB	[2002STE/CHI5]
	$\Delta_v H$	(329–493)	47.8	344	A	[1987STE/MAL]
	$\Delta_v H$	(353–393)	47.3	368	A	[1987STE/MAL]
C ₄ HF ₈ N	[14003-49-7]	N,N-bis(trifluoromethyl)-1,2-difluorovinylamine				
	$\Delta_v H$	(276–296)	28.8	286	A	[1987STE/MAL, 1968HAS/TIP]
C ₄ HF ₈ N	[13747-23-4]	N,N-bis(trifluoromethyl)-2,2-difluorovinylamine				
	$\Delta_v H$	(274–291)	27.7	282	A	[1987STE/MAL, 1968HAS/TIP]
C ₄ HF ₈ NO	[13580-54-6]	2,2,3,3,5,5,6,6-octafluoromorpholine				
	$\Delta_v H$	(273–323)	32.7	288	A	[1987STE/MAL, 67BAN/HAS]
C ₄ HF ₈ NOS	[77589-47-0]	2,2,3,3,4,4,5,5-octafluoro-1,1,2,3,4,5-hexahydro-1-iminothiophene-1-oxide				
	$\Delta_v H$		28.0	397		[1981ABE/SHR]
C ₄ HF ₉ N ₂ OS	[62609-65-8]	1,1,1-trifluoro-N-[2,2,2-trifluoro-1-(trifluoromethyl)ethylidene]-methanesulfonimidamide				
	$\Delta_v H$		37.2	388	I	[1977KIT/SHR]
C ₄ HF ₉ O ₂ S	[52225-50-0]	trifluoromethanesulfinic acid, 2,2,2-trifluoro-1-(trifluoromethyl)ethyl ester				
	$\Delta_v H$		39.3	362	HG	[1974MAJ/SHR]
C ₄ HF ₁₀ N	[54566-81-3]	1,1,1,2,3,3,3-heptafluoro-N-(pentafluoroethyl)ethanamine				
	$\Delta_v H$		26.4	306		[1975PET/SHR2]
C ₄ HF ₁₀ N	[53684-05-2]	1,1,1,2,3,3,3-heptafluoro-N-(trifluoromethyl)-2-propanamine				
	$\Delta_v H$		28.1	309		[1975KIR/LAS]
C ₄ HF ₁₀ NOS	[34556-24-6]	S,S-bis(pentafluoroethyl)sulfoximine				
	$\Delta_v H$		35.6	366	I	[1972SAU/SHR]
C ₄ H ₂	[460-12-8]	1,3-butadiyne				
	$\Delta_{\text{sub}} H$	(190–232)	36.2	211	A	[1947STU]
	$\Delta_{\text{sub}} H$	(188–234)	36.3			[1933TAN]
	$\Delta_v H$	(237–283)	26.1	268	A	[1987STE/MAL, 1971DYK]
	$\Delta_v H$	(191–282)	26.4	267		[1947STU]
	$\Delta_v H$	(188–234)	33.4	219		[1933TAN, 1984BOU/FRI]
	$\Delta_v H$	(195–273)	25.4	258		[1926STR, 1984BOU/FRI]
C ₄ H ₂ BrF ₆ N	[19451-87-7]	1-bromo-N,N-bis(trifluoromethyl)vinylamine				

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(288–327)	32.8	303	A	[1987STE/MAL, 1968FRE/TIP]
C ₄ H ₂ BrF ₆ N	[19483-21-7]	<i>cis</i> 2-bromo-N,N-bis(trifluoromethyl)vinylamine				
	$\Delta_v H$	(314–346)	29.7	329	A	[1987STE/MAL, 1968FRE/TIP]
C ₄ H ₂ BrF ₆ N	[19483-20-6]	<i>trans</i> 2-bromo-N,N-bis(trifluoromethyl)vinylamine				
	$\Delta_v H$	(314–341)	30.0	327	A	[1987STE/MAL, 1968FRE/TIP]
C ₄ H ₂ BrF ₈ N	[6857-63-2]	2-bromo-1,2-difluoro-N,N-bis(trifluoromethyl)ethylamine				
	$\Delta_v H$	(323–348)	32.4	328	A	[1987STE/MAL, 1965SHAS/TIP]
C ₄ H ₂ BrF ₈ N	[5003-73-6]	2-bromo-2,2-difluoro-N,N-bis(trifluoromethyl)ethylamine				
	$\Delta_v H$	(313–348)	33.6	328	A	[1987STE/MAL, 1965SHAS/TIP]
C ₄ H ₂ Br ₂ S	[3141-26-2]	3,4-dibromothiophene				
	$\Delta_v H$	(333–374)	32.1	348	A, I	[1987STE/MAL, 1971EON/POM, 1999DYK/SVO]
C ₄ H ₂ Cl ₂ N ₂	[4774-14-5]	2,6-dichloropyrazine				
	$\Delta_{\text{sub}} H$		69.9 ± 2.0	298	C	[2004MOR/MIR]
C ₄ H ₂ Cl ₂ N ₂	[3934-20-1]	2,4-dichloropyrimidine				
	$\Delta_{\text{sub}} H$		76.5 ± 2.0	298	C	[2007RIB/AMA]
C ₄ H ₂ Cl ₂ O ₂	[627-53-4]	<i>trans</i> fumaroyl chloride				
	$\Delta_v H$	(288–433)	45.6	303	A	[1987STE/MAL, 1947STU]
C ₄ H ₂ Cl ₂ S	[3172-52-9]	2,5-dichlorothiophene				
	$\Delta_{\text{fus}} H$		11.87	232.7		[2006FUJ/MAT]
	$\Delta_v H$	(323–425)	49.9	338		[1999DYK/SVO]
	$\Delta_v H$	(323–425)	36.2	338		[1999DYK/SVO]
	$\Delta_v H$	(323–425)	40.7	338		[1981DIT/SKO]
	$\Delta_v H$	(333–374)	33.7	348	A, I	[1987STE/MAL, 1971EON/POM, 1999DYK/SVO]
C ₄ H ₂ F ₄	[407-70-5]	1,1,4,4-tetrafluoro-1,3-butadiene				
	$\Delta_v H$	(239–271)	22.4	256	A	[1987STE/MAL]
C ₄ H ₂ F ₈	[662-35-1]	1,1,1,2,2,3,3,4-octafluorobutane				
	$\Delta_v H$	(260–278)	28.9	269	EB	[1997DEF/CAR]
C ₄ H ₂ F ₆ O ₂	[407-38-5]	trifluoroacetic acid, 2,2,2-trifluoroethyl ester				
	$\Delta_v H$		31.8	330	HG	[1973MAJ/SHR]
C ₄ H ₂ F ₇ S	[25273-51-2]	<i>cis</i> 2-fluoro-N,N-bis(trifluoromethyl)vinylamine				
	$\Delta_v H$	(289–311)	29.1	300	A	[1987STE/MAL]
C ₄ H ₂ F ₇ S	[25211-47-6]	<i>trans</i> 2-fluoro-N,N-bis(trifluoromethyl)vinylamine				
	$\Delta_v H$	(273–295)	28.5	284	A	[1987STE/MAL]
C ₄ H ₂ F ₈	[662-35-1]	1,1,1,2,2,3,3,4-octafluorobutane				
	$\Delta_v H$	(260–278)	28.9	269	EB	[1997DEF/CAR]
C ₄ H ₂ F ₈ O	[26103-08-2]	2-difluoromethoxy-1,1,1,3,3,3-hexafluoropropane				
	$\Delta_v H$	(283–315)	31.1	298	I	[2002MUR/YAM]
C ₄ H ₂ F ₈ O ₂	[188690-78-0]	1,2-bis(difluoromethoxy)-1,1,2,2-tetrafluoroethane				
	$\Delta_{\text{fus}} H$		11.8	195		[1999MAR/BAS]
	$\Delta_v H$	(253–331)	36.1 ± 0.2			[1999MAR/BAS]
C ₄ H ₂ F ₈ O ₃	[249932-25-0]	oxybis[(difluoromethoxy)difluoromethane]				
	$\Delta_{\text{fus}} H$		3.1	153		[1999MAR/BAS]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
	$\Delta_{\text{v}}H$	(253–341)	35.2 ± 0.2			[1999MAR/BAS]
C ₄ H ₂ N ₂	[764-42-1]	fumaronitrile				
	$\Delta_{\text{sub}}H$	(250–269)	69.6	260	TE,ME	[1983DEW/VAN]
	$\Delta_{\text{sub}}H$		68.6	298		[1983DEW/VAN]
	$\Delta_{\text{sub}}H$	(245–281)	72 ± 0.8	263	ME	[1967BOY/GUH, 1970COX/PIL]
C ₄ H ₂ N ₂ O ₄ S	[5347-12-6]	2,4-dinitrothiophene				
	$\Delta_{\text{v}}H$	(388–523)	59.7	403	A	[1987STE/MAL, 1971DYK, 1999DYK/SVO]
C ₄ H ₂ N ₂ O ₄ S	[59434-05-8]	2,5-dinitrothiophene				
	$\Delta_{\text{v}}H$	(388–523)	59.6	403		[1999DYK/SVO]
C ₄ H ₂ N ₂ S	[1452-15-9]	4-cyanothiazole				
	$\Delta_{\text{sub}}H$		73.9 ± 0.4	298	C	[1966MAN/SUN, 1970COX/PIL]
C ₄ H ₂ O ₃	[108-31-6]	maleic anhydride				
	$\Delta_{\text{fus}}H$		12.26	325.7		[1996DOM/HEA]
	$\Delta_{\text{sub}}H$	(308–326)	85.4	317		[1987STE/MAL]
	$\Delta_{\text{sub}}H$		68.8	258	TE,ME	[1983DEW/VAN]
	$\Delta_{\text{sub}}H$		70	298		[1978VIL/PER]
	$\Delta_{\text{sub}}H$	(308–325)	71.5 ± 5.0			[1949WIN/KUL, 1970COX/PIL]
	$\Delta_{\text{v}}H$	(336–475)	49.1	351	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$	(326–350)	54.8			[1949WIN/KUL]
C ₄ H ₂ O ₄	[142-45-0]	butyndioic acid				
	$\Delta_{\text{sub}}H$		NA			[1972LEB/KAT]
C ₄ H ₂ O ₄	[2892-51-5]	3,4-dihydroxy-3-cyclobutene-1,2-dione				
	$\Delta_{\text{sub}}H$	(469–499)	152.5	486	ME,TE	[1983DEW/VAN]
	$\Delta_{\text{sub}}H$		154.3	298		[1983DEW/VAN]
	$\Delta_{\text{sub}}H$		83.7 ± 16.7	298	E	[1971SEL2, 1977PED/RYL]
C ₄ H ₃ BrF ₇ N	[25237-12-1]	2-bromo-2-fluoro-N,N-bis(trifluoromethyl)ethylamine				
	$\Delta_{\text{v}}H$	(329–255)	30.9	342	A	[1987STE/MAL]
C ₄ H ₃ BrN ₂ O ₂	[51-20-7]	5-bromouracil				
	$\Delta_{\text{sub}}H$	(405–414)	148.1		ME	[2002SZT/KAM]
	$\Delta_{\text{sub}}H$		151.4 ± 2.5	298	ME	[2002SZT/KAM, 2005ZIE/SZT]
	$\Delta_{\text{sub}}H$		128.4		LE	[1974YAN/VER]
C ₄ H ₃ BrS	[1003-09-4]	2-bromothiophene				
	$\Delta_{\text{fus}}H$		0.01	55.3		
	$\Delta_{\text{fus}}H$		7.9	203.9		[1996DOM/HEA]
	$\Delta_{\text{v}}H$	(333–373)	27.9	348	A, I	[1987STE/MAL, 1971EON/POM, 1999DYK/SVO]
C ₄ H ₃ BrS	[872-31-1]	3-bromothiophene				
	$\Delta_{\text{v}}H$	(333–373)	28.9	348	A, I	[1987STE/MAL, 1971EON/POM, 1999DYK/SVO]
C ₄ H ₃ ClF ₆ O ₂ S	[57169-82-1]	chlorosulfurous acid, 2,2,2-trifluoro-1-methyl-1-(trifluoromethyl)ethyl ester				
	$\Delta_{\text{v}}H$		39.7			[1975DEM/KOV2]
C ₄ H ₃ ClN ₂	[14508-49-7]	2-chloropyrazine				
	$\Delta_{\text{v}}H$		45.1 ± 1.5	298	C	[2004MOR/MIR]
C ₄ H ₃ ClN ₂	[1722-12-9]	2-chloropyrimidine				
	$\Delta_{\text{sub}}H$		70.1 ± 1.3	298	C	[2007RIB/AMA]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
C ₄ H ₃ ClN ₂ O ₂	[1820-81-1]	5-chlorouracil				
	$\Delta_{\text{sub}}H$	(402–412)	145.5		ME	[2002SZT/KAM]
	$\Delta_{\text{sub}}H$		148.3 ± 2.4	298	ME	[2002SZT/KAM, 2005ZIE/SZT]
C ₄ H ₃ ClN ₂ O ₂	[4270-27-3]	6-chlorouracil				
	$\Delta_{\text{sub}}H$	(381–392)	134.3		ME	[2002SZT/KAM]
	$\Delta_{\text{sub}}H$		135.2 ± 2.0	298	ME	[2002SZT/KAM, 2005ZIE/SZT]
C ₄ H ₃ ClS	[96-43-5]	2-chlorothiophene				
	$\Delta_{\text{fus}}H$		8.97	201.3		[1993FUJ/OGU]
	$\Delta_{\text{v}}H$	(313–401)	34.7	328		[1999DYK/SVO]
	$\Delta_{\text{v}}H$	(320–401)	36.9	335		[1999DYK/SVO, 1981DIT/SKO]
	$\Delta_{\text{v}}H$	(333–374)	34.4	348	A, I	[1987STE/MAL, 1971EON/POM, 1999DYK/SVO]
C ₄ H ₃ ClS	[17249-80-8]	3-chlorothiophene				
	$\Delta_{\text{fus}}H$	(13–300)	9.39	214.2	AC	[2004FUJ/TOD]
C ₄ H ₃ Cl ₃ OS	[na]	2,3,3-trichloro-2-propenethioic acid, O-methyl ester				
	$\Delta_{\text{v}}H$	(383–423)	64.8		GC	[1980PIT/KIS]
C ₄ H ₃ FN ₂ O ₂	[51-21-8]	5-fluorouracil				
	$\Delta_{\text{sub}}H$	(394–401)	129.9		ME	[2002SZT/KAM]
	$\Delta_{\text{sub}}H$		133.2 ± 2.1	298	ME	[2002SZT/KAM, 2005ZIE/SZT]
	$\Delta_{\text{sub}}H$	(421–483)	150 ± 2		TE	[2002BRU/POR]
C ₄ H ₃ F ₅ OS	[35709-11-6]	trifluoroacetic acid, S-(2,2-difluoroethyl) ester				
	$\Delta_{\text{v}}H$	(282–322)	39.3	297	A	[1987STE/MAL, 1999DYK/SVO]
C ₄ H ₃ F ₅ O ₃	[2195-84-8]	α -(trifluoromethoxy)- α , α -difluoromethyl acetate				
	$\Delta_{\text{fus}}H$		8.51	167.4		[1996DOM/HEA]
C ₄ H ₃ F ₆ NO ₂	[22743-78-8]	N,N-bis(trifluoromethyl)acetamide-N-oxide				
	$\Delta_{\text{v}}H$	(268–336)	40.6	283	A	[1987STE/MAL, 1968NAS/BAB]
C ₄ H ₃ F ₆ O ₂ P	[2022-79-9]	bis(trifluoromethyl)acetoxyposphine				
	$\Delta_{\text{v}}H$	(273–313)	41	288		[1964PET/BUR, 1984BOU/FRI]
C ₄ H ₃ F ₇ O	[375-01-9]	2,2,3,3,4,4,4-heptafluoro-1-butanol				
	$\Delta_{\text{v}}H$	(273–298)	43.6	286	A, MM	[1987STE/MAL, 1971DYK, 1967MEE/GOL]
C ₄ H ₃ F ₇ O	[171182-95-9]	1-(2,2-difluoroethoxy)-1,1,2,2,2-pentafluoroethane				
	$\Delta_{\text{v}}H$	(288–318)	31.5	303	I	[2002MUR/YAM]
C ₄ H ₃ F ₇ O	[306-78-0]	1,1,2,2-tetrafluoro-1-(2,2,2-trifluoroethoxy)ethane				
	$\Delta_{\text{v}}H$	(283–329)	34	298	I	[2002MUR/YAM]
C ₄ H ₃ F ₇ O	[1683-81-4]	1,1,2,2-tetrafluoro-3-(trifluoromethoxy)propane				
	$\Delta_{\text{v}}H$	(288–319)	31.3	303	I	[2002MUR/YAM]
C ₄ H ₃ F ₇ O	[28523-86-6]	1,1,1,3,3,3-hexafluoro-2-fluoromethoxypropane				
	$\Delta_{\text{v}}H$	(288–331)	34.1	303	I	[2002MUR/YAM]
C ₄ H ₃ F ₇ O	[56860-81-2]	3-(difluoromethoxy)-1,1,1,2,2-pentafluoropropane				
	$\Delta_{\text{v}}H$	(283–319)	31.2	298	I	[2002MUR/YAM]
C ₄ H ₃ F ₇ O ₂ S	[57169-83-2]	fluorosulfurous acid, 2,2,2-trifluoro-1-methyl-1-(trifluoromethyl)ethyl ester				
	$\Delta_{\text{v}}H$		36.4			[1975DEM/KOV2]
C ₄ H ₃ IN ₂ O ₂	[696-07-1]	5-iodouracil				
	$\Delta_{\text{sub}}H$	(381–392)	127		ME	[2002SZT/KAM]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₄ H ₃ IS	[3437-95-4] Δ_vH	2-iodothiophene (333–374)	29.0	348	A, I	[1987STE/MAL, 1971EON/POM, 1999DYK/SVO]
C ₄ H ₃ NO ₂ S	[609-40-5] Δ_vH	2-nitrothiophene (378–443)	48.6	393		[1999DYK/SVO]
	Δ_vH	(321–498)	50.4	336	A	[1987STE/MAL]
C ₄ H ₃ NO ₃	[609-39-2] $\Delta_{\text{sub}}H$	2-nitrofuran	75.3 ± 2.1			[1980BAL/LEB, 1986PED/NAY]
C ₄ H ₄	[689-97-4] Δ_vH	1-butene-3-yne (180–278)	26.0	236	A	[1987STE/MAL, 1947STU]
C ₄ H ₄ BrF ₆ N	[1683-83-6] Δ_vH	2-bromo-N,N-bis(trifluoromethyl)ethylamine (323–356)	31.0	338	A	[1987STE/MAL, 1965HAS/TIP]
C ₄ H ₄ BrN ₃ O	[2240-25-7] $\Delta_{\text{sub}}H$	5-bromocytosine (403–468)	148.4 ± 1.5	435		[1975STEP/YAN]
C ₄ H ₄ Cl ₂	[3574-40-1] Δ_vH	1,2-dichloro-1,3-butadiene (260–308)	33.3	275	A	[1987STE/MAL]
C ₄ H ₄ Cl ₂	[1653-19-6] Δ_vH	2,3-dichloro-1,3-butadiene (299–368)	33.8	314	A	[1987STE/MAL]
C ₄ H ₄ Cl ₂ O ₂	[543-20-4] Δ_vH	succinyl chloride (312–466)	54.7	327	A	[1987STE/MAL, 1947STU]
C ₄ H ₄ Cl ₂ O ₃	[541-88-8] Δ_vH	chloroacetic acid anhydride (340–490)	61.8	355	A	[1987STE/MAL, 1947STU]
C ₄ H ₄ Cl ₄ O ₂ S	[3737-41-5] $\Delta_{\text{sub}}H$	3,3,4,4-tetrachlorotetrahydrothiophene-1,1-dioxide (303–348)	88.7	325	ME	[1978DEP]
	$\Delta_{\text{sub}}H$	(303–348)	88.6	318	A	[1987STE/MAL, 1999DYK/SVO]
C ₄ H ₄ F ₃ NO ₃	[383-70-0] $\Delta_{\text{sub}}H$	N-(trifluoroacetyl)aminoacetic acid (273–393)	98.8	288	A	[1987STE/MAL, 1960WEY/KLI]
C ₄ H ₄ F ₄ OS	[35709-10-5] Δ_vH	trifluorothioacetic acid, S-(2-fluoroethyl) ester (282–322)	41.4	297	A	[1987STE/MAL, 1999DYK/SVO]
C ₄ H ₄ F ₆ N ₂ S	[62067-10-1] Δ_vH	2,2,2-trifluoro-N-methyl-N'-[(trifluoromethyl)thio]ethanimidamide (339–387)	34.9	354	A, I	[1987STE/MAL, 1977BUR/SHR2, 1999DYK/SVO]
C ₄ H ₄ F ₆ O	[35042-99-0] Δ_vH	3-difluoromethoxy-1,1,2,2-tetrafluoropropane (283–349)	35.9	298	I	[2002MUR/YAM]
C ₄ H ₄ F ₆ O	[13171-18-1] Δ_vH	1,1,1,3,3,3-hexafluoro-2-methoxypropane (283–324)	32.6	298	I	[2002MUR/YAM]
C ₄ H ₄ F ₆ O	[25449-61-0] Δ_vH	1,1,1-trifluoro-2-(1,1,2-trifluoroethoxy)ethane (283–338)	35.4	298	I	[2002MUR/YAM]
C ₄ H ₄ F ₆ O	[333-36-8] Δ_vH	1,1,1-trifluoro-2-(2,2,2-trifluoroethoxy)ethane (283–337)	35	298	I	[2002MUR/YAM]
C ₄ H ₄ F ₆ O	[50807-77-7] Δ_vH	1-(1,1-difluoroethoxy)-1,1,2,2-tetrafluoroethane (288–352)	38.1	303	I	[2002MUR/YAM]
C ₄ H ₄ F ₆ O	[382-34-3] Δ_vH	1,1,1,2,3,3-hexafluoro-3-methoxypropane (288–327)	32.4	303	I	[2002MUR/YAM]
C ₄ H ₄ F ₆ O	[160620-20-2]	1,1,2,2,3,3-hexafluoro-1-methoxypropane				

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(288–341)	34.5	303	I	[2002MUR/YAM]
C ₄ H ₄ F ₆ O ₂ S	[52225-48-6]	trifluoromethanesulfinic acid, 2,2,2-trifluoro-1-methylethyl ester				
	$\Delta_v H$		36.8	375	HG	[1974MAJ/SHR]
C ₄ H ₄ IN ₃ O	[1122-44-7]	5-iodocytosine (4-amino-5-iodopyrimidinone)				
	$\Delta_{\text{sub}} H$	(413–463)	144 ± 1.5	438		[1975TEP/YAN]
C ₄ H ₄ N ₂	[290-37-9]	pyrazine				
	$\Delta_{\text{trs}} H$	(5–380)	1.09	300.5		
	$\Delta_{\text{trs}} H$	(5–380)	0.05	309.8		
	$\Delta_{\text{fus}} H$	(5–380)	14.78	325.4	AC	[2003CHI/KNI]
	$\Delta_{\text{fus}} H$		12.95	328.2		[1978BOU/LEC]
	$\Delta_{\text{sub}} H$	(288–317)	56.2	303		[1995SAK/UEO]
	$\Delta_{\text{sub}} H$		56.3 ± 0.5		C	[1962TJE, 1970COX/PIL]
	$\Delta_v H$	(342–373)	40.5 ± 1.7	298	CGC	[2009LIP/CHI2, 2009LIP/CHI]
	$\Delta_v H$	(354–426)	38.8 ± 0.1	340	EB	[2002STE/CHI3]
	$\Delta_v H$	(354–426)	36.5 ± 0.2	380	EB	[2002STE/CHI3]
	$\Delta_v H$	(354–426)	34.1 ± 0.4	420	EB	[2002STE/CHI3]
	$\Delta_v H$	(332–373)	37.9	352		[1995SAK/UEO]
C ₄ H ₄ N ₂	[289-95-2]	pyrimidine				
	$\Delta_v H$	(342–373)	41.0 ± 1.9	298	CGC	[2009LIP/CHI2, 2009LIP/CHI]
	$\Delta_v H$		49.9 ± 0.6	298	C	[1977NAB/SAB]
$\Delta_v H$		49.8 ± 0.3	298	C	[1962TJE]	
C ₄ H ₄ N ₂	[289-80-5]	pyridazine				
	$\Delta_v H$		53.5 ± 0.4	298	C	[1962TJE]
C ₄ H ₄ N ₂	[110-61-2]	succinonitrile				
	$\Delta_{\text{fus}} H$		3.7	329.7	DSC	[2009RAI/RED]
	$\Delta_{\text{trs}} H$		6.09	233.6		
	$\Delta_{\text{fus}} H$		3.75	330.3	DSC	[2007BAD/BLA]
	$\Delta_{\text{trs}} H$		6.2	233.3		
	$\Delta_{\text{fus}} H$	(5–350)	3.7	331.2	AC	[1996DOM/HEA, 1963WUL/WES]
$\Delta_{\text{sub}} H$	(279–298)	70 ± 0.3	289		[1960WOO/MUR, 1977PED/RYL, 1971RAP/WES, 1969STU/WES]	
C ₄ H ₄ N ₂ OS	[141-90-2]	2-thiouracil				
	$\Delta_{\text{sub}} H$		129.3		LE	[1974YAN/VER]
C ₄ H ₄ N ₂ OS	[591-28-6]	4-thiouracil				
	$\Delta_{\text{sub}} H$		125.5		LE	[1974YAN/VER]
C ₄ H ₄ N ₂ O ₂	[2423-84-9]	pyrazine 1,4-dioxide				
	$\Delta_{\text{sub}} H$		116.9 ± 0.8	298	C	[1997ACR/POW]
C ₄ H ₄ N ₂ O ₂	[66-22-8]	uracil				
	$\Delta_{\text{sub}} H$	(315–435)	125.3 ± 0.2	425	QR,ME	[2006DEB/MED]
	$\Delta_{\text{sub}} H$	(399–411)	101.3		ME	[2002SZT/KAM]
	$\Delta_{\text{sub}} H$	(394–494)	127.0 ± 2.0	439	TE	[2000BRU/PIA]
	$\Delta_{\text{sub}} H$	(452–587)	130.6 ± 4.0	519	ME, TE	[1980BAR/BEN]
	$\Delta_{\text{sub}} H$	(452–587)	131 ± 5	298	TE,GS	[1980BAR/BEN]
	$\Delta_{\text{sub}} H$	(378–428)	120.5 ± 1.3	403	QR	[1980TEP/YAN]
	$\Delta_{\text{sub}} H$		121.7	425	MS	[1979YAN/TEP]
	$\Delta_{\text{sub}} H$	(500–545)	133.9 ± 8	523	HSA	[1978NOW/SZC]
$\Delta_{\text{sub}} H$		126.5 ± 2.2	440	C	[1977NAB/SAB]	

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
	$\Delta_{\text{sub}}H$	(393–458)	120.5 ± 5.2	426	LE	[1975YAN/TEP, 1974YAN/VER]
	$\Delta_{\text{sub}}H$		115.5 ± 2.1		ME	[1972ROM/SUK, 2000BRU/PIA]
	$\Delta_{\text{sub}}H$		U 83.7			[1965CLA/PES]
C ₄ H ₄ N ₂ O ₂ S	[504-17-6]	thiobarbituric acid				
	$\Delta_{\text{sub}}H$	(400–461)	110 ± 4.0	430	TE	[1999BRU/PIA]
C ₄ H ₄ N ₂ O ₃	[67-52-7]	barbituric acid				
	$\Delta_{\text{trs}}H$		1.3	516		
	$\Delta_{\text{fus}}H$		20.87	526.4		[2008ROU/TEM2]
	$\Delta_{\text{sub}}H$	(294–438)	111.3 ± 0.3		GS	[1999ZIE/PER]
	$\Delta_{\text{sub}}H$	(392–493)	113 ± 4.0	442	TE	[1999BRU/PIA]
	$\Delta_{\text{sub}}H$	(404–479)	123.3 ± 1.7	440	ME	[1990SOL/KAB]
C ₄ H ₄ N ₂ S ₂	[2001-93-6]	2,4-dithiouracil				
	$\Delta_{\text{sub}}H$	(393–443)	119.7 ± 2.4	418		[1975STEP/YAN]
C ₄ H ₄ N ₄ O ₂	[5424-94-2]	N-nitro- <i>bis</i> (N,N-cyanomethyl) amine				
	$\Delta_{\text{fus}}H$		38.66	367		[1987OYU/BR1]
C ₄ H ₄ N ₄ O ₇	[57449-43-1]	furazandimethanol dinitrate				
	$\Delta_{\text{v}}H$	(399–433)	58.7	414	A	[1987STE/MAL, 1975PEP/MAT]
C ₄ H ₄ N ₄ O ₈	[57449-44-2]	furazandimethanol dinitrate, 2-oxide				
	$\Delta_{\text{v}}H$	(413–453)	64.3	428	A	[1987STE/MAL, 1975PEP/MAT]
C ₄ H ₄ N ₆	[1123-54-2]	8-azaadenine				
	$\Delta_{\text{sub}}H$	(418–463)	128.4 ± 1.3	440		[1975STEP/YAN]
C ₄ H ₄ N ₈ O ₁₃	[34882-73-0]	<i>bis</i> -(2,2,2-trinitroethyl)-N-nitrosoamine				
	$\Delta_{\text{sub}}H$	(333–354)	97.9 ± 0.8	343	ME	[1973DEK/OON]
C ₄ H ₄ N ₈ O ₁₄	[19836-28-3]	<i>bis</i> -(2,2,2-trinitroethyl)-N-nitroamine				
	$\Delta_{\text{sub}}H$	(340–356)	117.6 ± 0.8	348	ME	[1973DEK/OON]
C ₄ H ₄ O	[110-00-9]	furan				
	$\Delta_{\text{trs}}H$		2.05	150		
	$\Delta_{\text{fus}}H$		3.8	187.6		[1996DOM/HEA]
	$\Delta_{\text{sub}}H$		NA			[1953MIL/PAO]
	$\Delta_{\text{v}}H$	(238–356)	30.2	253	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$	(277–323)	28.2	298		[1970MOI/ANT]
	$\Delta_{\text{v}}H$	(275–334)	28.6	290		[1952GUT/SCO, 1984BOU/FRI]
C ₄ H ₄ OS	[3354-32-3]	2-(5 <i>H</i>)-thiophenone				
	$\Delta_{\text{v}}H$		56.0 ± 1.2	298	C	[2010RIB/SAN3]
C ₄ H ₄ O ₂	[674-82-8]	diketene				
	$\Delta_{\text{v}}H$	(297–388)	42.9	312	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$		42.9 ± 0.1	298	C	[1968MAN/NAK]
C ₄ H ₄ O ₂	[33689-28-0]	cyclobutane-1,2-dione				
	$\Delta_{\text{sub}}H$	(251–289)	69.1 ± 3.5	270	HSA	[UR/CHI]
	$\Delta_{\text{sub}}H$	(295–335)	54.8	315		[1985CAO/BAC]
C ₄ H ₄ O ₂	[15506-53-3]	cyclobutane-1,3-dione				
	$\Delta_{\text{sub}}H$	(274–322)	73.6 ± 3.7	298	HSA	[1978CHI/SHE]
C ₄ H ₄ O ₂	[497-23-4]	2-(5 <i>H</i>)-furanone				
	$\Delta_{\text{v}}H$		55.6 ± 1.3	298	C	[2010RIB/SAN3]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₄ H ₄ O ₃	[108-30-5]	succinic anhydride				
	$\Delta_{\text{sub}}H$	(298–320)	80.5 ± 1.6	309	ME	[1990MEN/PIL]
	$\Delta_{\text{sub}}H$		80.7 ± 1.6	298	C	[1990MEN/PIL]
	$\Delta_{\text{sub}}H$	(290–311)	82.2	302	ME,TE	[1983DEW/VAN]
	Δ_vH	(401–534)	57.3	416	A	[1987STE/MAL]
C ₄ H ₄ O ₄	[110-16-7]	<i>cis</i> -butenedioic acid (maleic acid)				
	$\Delta_{\text{fus}}H$		26.9	411.9	DSC	[2010GUO/SAD]
	$\Delta_{\text{sub}}H$	(348–389)	105.4 ± 1.7	368	ME	[1974ARS]
	$\Delta_{\text{sub}}H$	(357–367)	110 ± 2.5			[1938WOL/WEG, 1960JON, 1970COX/PIL]
	$\Delta_{\text{sub}}H$	(356–371)	109 ± 4.2			[1934WOL/TRI]
C ₄ H ₄ O ₄	[110-17-8]	<i>trans</i> -butenedioic acid				
	$\Delta_{\text{sub}}H$	(371–391)	123.6 ± 2.0	381	TE,ME	[1977DEK/VAN]
	$\Delta_{\text{sub}}H$		136 ± 6.3	365	QF	[1938WOL/WEG, 1935TRI, 1960JON]
	$\Delta_{\text{sub}}H$	(358–371)	134 ± 4.2			[1934WOL/TRI]
C ₄ H ₄ O ₄	[4480-83-5]	diglycolic anhydride				
	$\Delta_{\text{sub}}H$	(382–303)	84.2 ± 1.1	294	ME, TE	[1983DEW/VAN]
C ₄ H ₄ O ₄	[502-97-6]	1,4-dioxane-2,5-dione				
	$\Delta_{\text{trs}}H$		1.81	312.1		
	$\Delta_{\text{fus}}H$		14.8	356.2	AC	[1996DOM/HEA, 1988LEB/KAL]
	Δ_vH	(376–513)	50.4	391	A	[1987STE/MAL, 1947STU]
C ₄ H ₄ O ₄	[3524-70-7]	ethylene oxalate				
	$\Delta_{\text{fus}}H$		13.4	415		[1996DOM/HEA]
C ₄ H ₄ S	[110-02-1]	thiophene				
	$\Delta_{\text{trs}}H$		1.21	171.1		
	$\Delta_{\text{fus}}H$		4.97	235.2		[1996DOM/HEA, 1985FIG/SZW]
	$\Delta_{\text{sub}}H$	(195–228)	46.8	213		[1987STE/MAL, 1956MIL]
	$\Delta_{\text{sub}}H$	(192–213)	49	203		[1944MIL]
	Δ_vH	(267–381)	35.8	282		[1999DYK/SVO]
	Δ_vH	(333–373)	34.8	348	I	[1971EON/POM]
	Δ_vH		34.6	298		[1971WIL/ZWO]
	Δ_vH	(300–366)	34.1	315	EB	[1952WHI/BER]
	Δ_vH	(311–393)	33.7	326		[1949WAD/KNO]
	Δ_vH		33.6 ± 0.1	319	C	[1949WAD/KNO]
Δ_vH		32.7 ± 0.1	336	C	[1949WAD/KNO]	
Δ_vH		31.5 ± 0.1	357	C	[1949WAD/KNO]	
Δ_vH	(344–363)	32.6	353		[1945FAW/RAS]	
Δ_vH	(228–289)	35	270		[1944MIL]	
C ₄ H ₅ Cl	[126-99-8]	2-chloro-1,3-butadiene				
	Δ_vH	(243–263)	29.6	253	A	[1987STE/MAL]
	Δ_vH	(279–333)	29.6	294	A	[1987STE/MAL, 1971DYK]
	Δ_vH	(293–333)	30.9	308		[1964GUB/FER, 1984BOU/FRI]
C ₄ H ₅ ClO	[920-46-7]	methacryloyl chloride				
	Δ_vH	(313–372)	36.1	328	A	[1987STE/MAL]
C ₄ H ₅ ClO ₂	[na]	Z-3-chloro-2-butenic acid				
	$\Delta_{\text{fus}}H$		20.71	366.8		[1996DOM/HEA]
C ₄ H ₅ ClO ₂	[na]	E-3-chloro-2-butenic acid				

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	Enthalpy							
		$\Delta_{\text{fus}}H$			13.81	333.7		[1996DOM/HEA]
C ₄ H ₅ ClO ₃	[4755-77-5]	Δ_vH	ethyl chloroglyoxylate	(268–408)	44.9	283		[1947STU]
C ₄ H ₅ Cl ₃ O ₂	[515-84-4]	Δ_vH	ethyl trichloroacetate	(293–440)	49	308	A	[1987STE/MAL]
		Δ_vH			51.0 ± 0.1	298	C	[1972LAY/WAD]
		Δ_vH		(317–368)	47.5	332		[1959USA/DEM, 1984BOU/FRI]
C ₄ H ₅ Cl ₅	[2431-52-9]	Δ_vH	1,2,2,3,4-pentachlorobutane	(368–498)	62.6	383	A	[1987STE/MAL, 1968CIH/VOJ]
C ₄ H ₅ F ₂ I	[692-26-2]	Δ_vH	1,1-difluoro-4-iodo-1-butene	(318–342)	40.6	330	A	[1987STE/MAL]
C ₄ H ₅ F ₃ O	[406-90-6]	Δ_vH	vinyl 2,2,2-trifluoroethyl ether	(293–317)	32.0	305	A	[1987STE/MAL]
C ₄ H ₅ F ₃ OS	[383-64-2]	Δ_vH	trifluorothioacetic acid, S-ethyl ester	(273–313)	42.0	288	A	[1987STE/MAL]
C ₄ H ₅ F ₃ O ₂	[383-63-1]	Δ_vH	trifluoroacetic acid, ethyl ester		34.7	335	HG	[1973MAJ/SHR]
C ₄ H ₅ F ₅	[406-58-6]	Δ_vH	1,1,1,3,3-pentafluorobutane	(303–358)	29.2	318		[2002MAR/OLI]
C ₄ H ₅ F ₅ O	[378-16-5]	Δ_vH	1,1,1,2,2-pentafluoro-3-methoxypropane	(283–321)	31.6	298	I	[2002MUR/YAM]
C ₄ H ₅ F ₅ O	[69948-24-9]	Δ_vH	1-(difluoromethoxy)-1,1,2-trifluoroethane	(283–316)	31.7	298	I	[2002MUR/YAM]
C ₄ H ₅ F ₆ OP	[1692-49-5]	Δ_vH	ethyl bis(trifluoromethyl)phosphinite	(248–328)	33.2	288		[1959EME/SMI]
C ₄ H ₅ N	[109-75-1]	Δ_vH	3-butenitrile	(293–417)	40.3	308	A	[1987STE/MAL]
		Δ_vH			40.0	298		[1969KON/PRO]
		Δ_vH		(254–392)	41.6	268		[1947STU]
C ₄ H ₅ N	[627-26-9]	Δ_vH	(E) 2-butenitrile		40.0	298		[1969KON/PRO]
C ₄ H ₅ N	[1190-76-7]	Δ_vH	(Z) 2-butenitrile		38.9	298		[1969KON/PRO]
C ₄ H ₅ N	[1190-76-7]	Δ_vH	<i>cis</i> crotonitrile	(297–405)	37.1	312	A	[1987STE/MAL]
		Δ_vH		(244–381)	39.0	259		[1947STU]
C ₄ H ₅ N	[627-26-7]	Δ_vH	<i>trans</i> crotonitrile	(292–420)	39.7	307	A	[1987STE/MAL]
		Δ_vH		(254–395)	40.5	268		[1947STU]
C ₄ H ₅ N	[126-98-7]	Δ_vH	methacrylonitrile	(273–373)	36.5	288	A	[1987STE/MAL]
		Δ_vH		(229–363)	35.4	243		[1947STU]
C ₄ H ₅ N	[5500-21-0]	Δ_vH	cyclopropylcyanide		41.9 ± 0.1	298	C	[1982FUC/HAL]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(310–391)	39.4	325	BG	[1971HAL/BAL]
	$\Delta_v H$	(310–391)	39.8 ± 0.4	298	BG	[1971HAL/BAL]
C₄H₅N	[109-97-7]	pyrrole				
	$\Delta_{\text{fus}}H$		7.91	249.7		[1996DOM/HEA]
	$\Delta_{\text{sub}}H$		NA			[1941MIL]
	$\Delta_v H$	(285–329)	42.5	300		[1992KIM/SZY]
	$\Delta_v H$	(313–373)	41.9	328	I	[1971EON/POM]
	$\Delta_v H$	(338–440)	42.5	353	A, EB, IP	[1987STE/MAL, 1968OSB/DOU, 1967SCO/BER]
	$\Delta_v H$		45.2	298		[1967SCO/BER]
	$\Delta_v H$		41.8			[1961ZIM/GEI]
	$\Delta_v H$	(333–373)	41.9	348		[1947STU]
C₄DH₄N	[10162-82-0]	N-deutero pyrrole				
	$\Delta_v H$	(285–329)	42.9	300		[1992KIM/SZY]
C₄H₅NO	[30842-90-1]	3-methylisoxazole				
	$\Delta_v H$		39.8 ± 0.2	298	C	[1978HAM/BEN]
C₄H₅NO	[5765-44-6]	5-methylisoxazole				
	$\Delta_v H$		39.7 ± 0.2	298	C	[1978HAM/BEN]
C₄H₅NO₂	[105-34-0]	methyl cyanoacetate				
	$\Delta_v H$	(292–322)	66.2 ± 0.9	298	GS	[1995VER/BEC]
	$\Delta_v H$	(385–573)	54.9	400	A	[1987STE/MAL, 1971DYK]
C₄H₅NO₂	[123-56-8]	succinimide				
	$\Delta_{\text{fus}}H$		17.0	400		[1996DOM/HEA]
	$\Delta_{\text{sub}}H$	(317–340)	83.1 ± 1.5	329	ME	[1990MEN/PIL]
	$\Delta_{\text{sub}}H$		83.6 ± 1.5	298		[1990MEN/PIL]
	$\Delta_{\text{sub}}H$		88.0		B	[1989STE/CHI2]
	$\Delta_v H$	(416–561)	66.9	431	A	[1987STE/MAL]
	$\Delta_v H$	(388–560)	73.5	403		[1947STU]
C₄H₅NS	[57-06-7]	allyl isothiocyanate				
	$\Delta_v H$	(277–323)	47.6		GC	[1997LIM/TUN]
	$\Delta_v H$	(370–430)	56.8	385	A	[1987STE/MAL]
	$\Delta_v H$	(283–323)	42.1	298		[1935BAU/BUR, 1984BOU/FRI, 1999DYK/SVO]
C₄H₅NS	[3581-87-1]	2-methylthiazole				
	$\Delta_{\text{fus}}H$		12.16	248.6		[1968GOU/WES, 1969SOU/GOU]
	$\Delta_v H$	(353–402)	39.4	368	A	[1987STE/MAL, 1999DYK/SVO]
	$\Delta_v H$	(342–404)	40.0	357	A	[1987STE/MAL, 1969SOU/GOU]
C₄H₅NS	[693-95-8]	4-methylthiazole				
	$\Delta_{\text{fus}}H$		8.9	229.1		[1966MEY/MET]
	$\Delta_v H$	(346–408)	40.8	361	A	[1987STE/MAL]
	$\Delta_v H$		43.8 ± 0.2	298	C	[1966MAN/SUN]
C₄H₅NS	[3581-89-3]	5-methylthiazole				
	$\Delta_{\text{fus}}H$		7.65	232.8		[1966MEY/MET]
C₄H₅N₃O	[71-30-7]	cytosine				
	$\Delta_{\text{sub}}H$	(320–410)	167.7 ± 0.5	365	QR,ME	[2006DEB/MED]
	$\Delta_{\text{sub}}H$	(505–525)	151.7 ± 0.7		GS	[1998ZIE/WSZ]
	$\Delta_{\text{sub}}H$	(423–483)	147.2 ± 2.6	453	ME	[1984BUR/MOR]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{sub}}H$		155.0 ± 3.0	298		[1984BUR/MOR]
	$\Delta_{\text{sub}}H$		167 ± 10	298	TE	[1980FER/BEN]
	$\Delta_{\text{sub}}H$	(450–470)	176 ± 10	298	C	[1980SAB]
	$\Delta_{\text{sub}}H$		NA			[1977SAB/NAB]
	$\Delta_{\text{sub}}H$		150.6		ME	[1974YAN/VER, 1975YAN/TEP]
C ₄ H ₅ N ₃ O ₂	[932-52-5]	5-aminouracil				
	$\Delta_{\text{sub}}H$		145.6		LE	[1974YAN/VER]
C ₄ H ₅ N ₃ O ₂	[932-53-6]	6-azathymine				
	$\Delta_{\text{sub}}H$	(358–403)	112.5 ± 2.3	380		[1974MAN3]
C ₄ H ₅ N ₃ S	[333-49-3]	2-thiocytosine				
	$\Delta_{\text{sub}}H$	(408–458)		158 ± 1.6	433	[1975STEP/YAN]
C ₄ H ₅ N ₇ O ₁₂	[34880-53-0]	2,2,2-trinitro-N-(2,2,2-trinitroethyl)ethanamine				
	$\Delta_{\text{sub}}H$	(338–349)	80.8 ± 0.4		ME, A	[1973DEK/OON, 1977PED/RYL, 1987STE/MAL]
C ₄ H ₆	[590-19-2]	1,2-butadiene				
	$\Delta_{\text{fus}}H$		6.95	136.9		[1996DOM/HEA]
	Δ_vH	(243–291)	25.3	276	A	[1987STE/MAL]
	Δ_vH	(204–243)	26.4	228	A	[1987STE/MAL]
	Δ_vH		23.9	298		[1971WIL/ZWO]
	Δ_vH	(184–291)	25.2	276		[1947STU]
	Δ_vH		24.6 ± 0.1	273	C	[1947AST/SZA]
C ₄ H ₆	[106-99-0]	1,3-butadiene				
	$\Delta_{\text{fus}}H$		7.98	164.2		[1991ACR]
	Δ_vH	(270–318)	23	285	A	[1987STE/MAL]
	Δ_vH	(193–213)	25.7	203	A	[1987STE/MAL]
	Δ_vH	(213–276)	23.6	261	A	[1987STE/MAL]
	Δ_vH	(315–382)	22.4	330	A	[1987STE/MAL]
	Δ_vH	(380–425)	22.9	395	A	[1987STE/MAL]
	Δ_vH		21.1	298		[1971WIL/ZWO, 1945PRO/ROS]
	Δ_vH	(198–271)	23.7	256		[1984BOU/FRI, 1933HEI2]
	Δ_vH	(191–249)	24.7	235		[1932VAU, 1984BOU/FRI]
C ₄ H ₆	[107-00-6]	1-butyne				
	$\Delta_{\text{fus}}H$		6.03	147.4		[1996DOM/HEA]
	Δ_vH	(205–289)	26	274	A	[1987STE/MAL]
	Δ_vH		23.7	298		[1971WIL/ZWO]
	Δ_vH		25.8 ± 0.1	263	C	[1950AST/MAS]
	Δ_vH		24.5 ± 0.1	281	C	[1950AST/MAS]
C ₄ H ₆	[503-17-3]	2-butyne				
	$\Delta_{\text{fus}}H$		9.25	240.9		[1991ACR]
	$\Delta_{\text{sub}}H$	(200–239)	37.4	220	A	[1947STU]
	Δ_vH	(240–308)	29	255	A	[1987STE/MAL]
	Δ_vH		26.7	298		[1971WIL/ZWO]
	Δ_vH		26.9 ± 0.1	291	C	[1941YOU/GAR]
C ₄ H ₆	[822-35-2]	cyclobutene				
	Δ_vH	(206–275)	24.7	260	A	[1987STE/MAL]
	Δ_vH	(196–275)	24.6	260		[1941HEI, 1984BOU/FRI]
C ₄ H ₆ ClFO ₂	[1537-62-8]	2-chloroethyl fluoroacetate				

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
	$\Delta_{\text{v}}H$	(273–333)	56.4	288	A, GS	[1987STE/MAL, 1948RED/CHA4, 1971DYK]
C₄H₆ClF₃O	[310-71-4]	2-chloro-1,1,2-trifluoroethyl ethyl ether				
	$\Delta_{\text{v}}H$		37.5 ± 0.1	298	C	[1984UCH/MAJ]
	$\Delta_{\text{v}}H$		36.5 ± 0.1	313	C	[1984UCH/MAJ]
	$\Delta_{\text{v}}H$		35.3 ± 0.1	328	C	[1984UCH/MAJ]
	$\Delta_{\text{v}}H$		34.2 ± 0.1	343	C	[1984UCH/MAJ]
	$\Delta_{\text{v}}H$		32.9 ± 0.1	358	C	[1984UCH/MAJ]
C₄H₆Cl₂	[760-23-6]	3,4-dichloro-1-butene				
	$\Delta_{\text{v}}H$	(320–396)	38	335	A	[1987STE/MAL]
C₄H₆Cl₂	[7415-31-8]	<i>trans</i> 1,3-dichloro-2-butene				
	$\Delta_{\text{v}}H$	(306–401)	39.3	321	A	[1987STE/MAL]
C₄H₆Cl₂	[110-57-6]	<i>trans</i> 1,4-dichloro-2-butene				
	$\Delta_{\text{v}}H$	(340–379)	45.6	355	A	[1987STE/MAL]
C₄H₆Cl₂O₂	[3848-12-2]	2-chloroethyl chloroacetate				
	$\Delta_{\text{v}}H$	(319–478)	53.3	334	A	[1987STE/MAL, 1947STU]
C₄H₆Cl₂O₂	[535-15-9]	ethyl dichloroacetate				
	$\Delta_{\text{v}}H$		50.6 ± 0.1	298	C	[1972LAY/WAD]
	$\Delta_{\text{v}}H$	(283–430)	46.2	298	A	[1987STE/MAL, 1947STU]
C₄H₆Cl₄	[13138-51-7]	1,2,3,3-tetrachlorobutane				
	$\Delta_{\text{v}}H$	(349–464)	54.2	364	A	[1987STE/MAL, 1968CIH/VOJ]
C₄H₆FN	[407-83-0]	4-fluorobutyronitrile				
	$\Delta_{\text{v}}H$	(273–333)	45.2	288	A, GS	[1987STE/MAL, 1948RED/CHA4, 1971DYK]
C₄H₆F₂O₂	[459-99-4]	2-fluoroethyl fluoroacetate				
	$\Delta_{\text{v}}H$	(273–333)	55.1	288	A, GS	[1987STE/MAL, 1948RED/CHA4, 1971DYK]
C₄H₆F₃I	[540-87-4]	1,1,1-trifluoro-3-iodobutane				
	$\Delta_{\text{v}}H$	(304–321)	32.4	312	A	[1987STE/MAL]
C₄H₆F₃I	[26653-47-4]	1,1,1-trifluoro-3-iodo-2-methylpropane				
	$\Delta_{\text{v}}H$	(298–368)	30.4	313	A	[1987STE/MAL]
C₄H₆F₃NO₃	[72316-38-2]	carbamic acid, methoxy(trifluoromethyl)-, methyl ester				
	$\Delta_{\text{v}}H$		39			[1979SEK/DES]
C₄H₆F₄O	[512-51-6]	1-ethoxy-1,1,2,2-tetrafluoroethane				
	$\Delta_{\text{v}}H$	(283–330)	33	298	I	[2002MUR/YAM]
C₄H₆F₄O	[60598-17-6]	1,1,2,2-tetrafluoro-3-methoxypropane				
	$\Delta_{\text{v}}H$	(293–347)	35.2	308	I	[2002MUR/YAM]
C₄H₆F₄O₂	[73287-23-7]	1,2-dimethoxytetrafluoroethane				
	$\Delta_{\text{fus}}H$		10.75	252	DSC	[2005MAR/AVA]
	$\Delta_{\text{v}}H$	(270–356)	33.7			[2005MAR/AVA]
C₄H₆F₆N₂O	[30295-33-1]	1,1-dimethyl-2,2-bis(trifluoromethyl)hydrazine-2-oxide				
	$\Delta_{\text{v}}H$	(287–356)	36.4	302	A	[1987STE/MAL]
C₄H₆F₆P₂S	[26348-88-9]	methyl(trifluoromethyl)phosphinothious acid, anhydrosulfide				
	$\Delta_{\text{v}}H$	(316–342)	46.7	329		[1970BUR/KAN]
C₄H₆N₂	[693-98-1]	2-methylimidazole				
	$\Delta_{\text{sub}}H$	(301–318)	88.2 ± 0.7	309	ME	[1992JIM/ROU]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{sub}}H$		88.4 ± 0.7	298		
C ₄ H ₆ N ₂	[1453-58-3]	3-methylpyrazole				
	Δ_vH		65.9 ± 2.0	298	C	[2008RIB/FIG]
C ₄ H ₆ N ₂ O	[1072-67-9]	3-amino-5-methylisoxazole				
	$\Delta_{\text{sub}}H$		81.6 ± 2.5			[1973HAM/AYE, 1977PED/RYL]
C ₄ H ₆ N ₂ O	[4975-21-7]	dimethylfurazan				
	Δ_vH	(353–427)	51.1	368	A	[1987STE/MAL, 1971MAT/PEP]
C ₄ H ₆ N ₂ O ₂	[106-57-0]	2,5-piperazinedione				
	$\Delta_{\text{sub}}H$	(413–450)	103.8	428	A	[1987STE/MAL]
C ₄ H ₆ N ₂ O ₂	[2518-42-5]	dimethylfurazan-2-oxide				
	Δ_vH	(353–493)	57.0	368	A	[1987STE/MAL, 1971MAT/PEP]
C ₄ H ₆ N ₄ O	[56-06-4]	2,4-diamino-6-hydroxypyrimidine				
	$\Delta_{\text{sub}}H$	(423–471)	147.6 ± 0.2		GS	[1999ZIE/PER]
C ₄ H ₆ N ₄ O ₃ S ₂	[59-66-5]	2-acetamido-1,3,4-thiadiazole-5-sulfonamide				
	$\Delta_{\text{trs}}H$		1.2	506.2	DSC	
	$\Delta_{\text{fus}}H$		28.6	532.2	DSC	[2009BAR/GAM]
C ₄ H ₆ N ₄ O ₆	[14760-99-7]	2,5-dinitro-2,5-diazahehexane-3,4-dione				
	$\Delta_{\text{fus}}H$		23.4	397.1	DSC	[1997ZEM]
C ₄ H ₆ N ₄ O ₈	[3759-60-2]	1,1,3,3-tetranitrobutane				
	$\Delta_{\text{sub}}H$		87.9 ± 0.8	298		[1999MIR/VOR]
C ₄ H ₆ N ₄ O ₈	[20919-97-5]	2,2,3,3-tetranitrobutane				
	$\Delta_{\text{sub}}H$		78.2 ± 0.8	298		[1999MIR/VOR]
C ₄ H ₆ N ₄ O ₈	[20919-96-4]	1,1,1,4-tetranitrobutane				
	$\Delta_{\text{sub}}H$		99.6	298		[1999MIR/VOR]
C ₄ H ₆ N ₄ O ₈	[42216-58-0]	1,1,1,3-tetranitro-2-methylpropane				
	$\Delta_{\text{sub}}H$		91.2	298		[1999MIR/VOR]
	Δ_vH	(304–327)	75.7	316	A	[1987STE/MAL]
C ₄ H ₆ N ₄ O ₁₁	[20820-44-4]	2-nitro-2-hydroxymethyl-1,3-propanedioltrinitrate				
	Δ_vH	(313–353)	72.9	328	A	[1987STE/MAL]
C ₄ H ₆ N ₆ O ₈	[81360-42-1]	1,3,5,5-tetranitro-1,3-diazacyclohexane				
	$\Delta_{\text{fus}}H$		29.37	430		[1987OYU/BRI, 1995SKA/GOL]
C ₄ H ₆ O	[123-73-9]	<i>trans</i> crotonaldehyde				
	Δ_vH	(314–411)	36.6 ± 0.1	320	EB	[2002STE/CHI2]
	Δ_vH	(314–411)	34.5 ± 0.2	360	EB	[2002STE/CHI2]
	Δ_vH	(314–411)	32.1 ± 0.5	400	EB	[2002STE/CHI2]
C ₄ H ₆ O	[4170-30-3]	crotonaldehyde				
	Δ_vH		37.3 ± 0.4	298	C	[1996VAN/YU]
	Δ_vH		38.8	325	EB	[1994WIB/MOR]
	Δ_vH	(288–376)	35.1 ± 0.5	332		[1988BAG/GUR]
	Δ_vH	(304–377)	37.3	319		[1979MAR/SAC]
	Δ_vH	(306–376)	36.8	321	A	[1987STE/MAL, 1973WAR/SKU, 1984BOU/FRI]
C ₄ H ₆ O	[1191-95-3]	cyclobutanone				
	$\Delta_{\text{fus}}H$		10.8	220.5		[1998GON/SZW]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(301–344)	37.7	322	EB	[1994WIB/MOR]
	$\Delta_v H$	(283–313)	38.4	298	A	[1987STE/MAL]
	$\Delta_v H$	(317–380)	36.3	332	A, EB	[1987STE/MAL, 1976MEY/HOT]
	$\Delta_v H$		38.2 ± 0.4	298		[1972WOL]
	$\Delta_v H$	(249–298)	38.5	273		[1942BEN/KIS, 1984BOU/FRI]
C₄H₆O	[109-93-3]	divinyl ether				
	$\Delta_v H$	(253–323)	29.2	268	A	[1987STE/MAL]
	$\Delta_v H$	(253–323)	26.1	301	I	[1933MIL/MEN]
C₄H₆O	[78-94-4]	methyl vinyl ketone				
	$\Delta_v H$	(279–355)	32.9	294	A	[1987STE/MAL]
	$\Delta_v H$	(300–355)	33.6	315	A	[1987STE/MAL]
C₄H₆O	[1191-99-7]	2,3-dihydrofuran				
	$\Delta_v H$	(302–260)	30.8 ± 0.1	300	EB	[2002STE/CHI5]
	$\Delta_v H$	(302–360)	28.6 ± 0.3	340	EB	[2002STE/CHI5]
C₄H₆O	[927-74-2]	3-butyn-1-ol				
	$\Delta_v H$	(343–393)	51.7 ± 0.9	298	CGC	[2005VAL/QUI]
C₄H₆OS	[1115-15-7]	divinyl sulfoxide				
	$\Delta_v H$		51.2 ± 0.9	298	C	[1989VOR/KLY]
C₄H₆O₂	[1759-53-1]	cyclopropane carboxylic acid				
	$\Delta_v H$	(357–473)	58.9 ± 0.3	340	EB	[2002STE/CHI5]
	$\Delta_v H$	(357–473)	55.7 ± 0.2	380	EB	[2002STE/CHI5]
	$\Delta_v H$	(357–473)	52.4 ± 0.2	420	EB	[2002STE/CHI5]
	$\Delta_v H$	(357–473)	48.8 ± 0.4	460	EB	[2002STE/CHI5]
C₄H₆O₂	[431-03-8]	2,3-butanedione (biacetyl)				
	$\Delta_v H$	(273–348)	38.5	288	A, I	[1987STE/MAL, 1972NEE/HAL]
	$\Delta_v H$	(273–293)	39.6 ± 0.2	283		[1954NIC/SZA]
C₄H₆O₂	[503-64-0]	<i>cis</i> 2-butenic acid				
	$\Delta_v H$	(306–445)	55.8	321	A	[1987STE/MAL, 1947STU]
C₄H₆O₂	[107-93-7]	<i>trans</i> 2-butenic acid				
	$\Delta_v H$	(353–458)	56.7	368	A	[1987STE/MAL, 1947STU]
C₄H₆O₂	[110-65-6]	2-butyne-1,4-diol				
	$\Delta_v H$	(418–520)	69.0	433	A	[1987STE/MAL, 1966BRA/SEM, 1971DYK]
C₄H₆O₂	[96-48-0]	γ -butyrolactone				
	$\Delta_{\text{fus}} H$		9.57	230		[1991ACR]
	$\Delta_v H$	(293–333)	47.6 ± 4.0	298		[1997KLE, 2008EME/KOZ]
	$\Delta_v H$	(378–406)	49.5 ± 0.1	392	EB	[1991WIB/WAL]
	$\Delta_v H$	(378–406)	55.2 ± 1.3	298	EB	[1991WIB/WAL]
	$\Delta_v H$	(345–370)	51.8 ± 0.6	357	MM	[1991WIB/WAL]
	$\Delta_v H$	(345–370)	55.6 ± 1.4	298	MM	[1991WIB/WAL]
	$\Delta_v H$	(392–555)	48.9 ± 0.3	298		[1990RAM/KUD, 2008EME/KOZ]
	$\Delta_v H$		54.4 ± 0.4	298	C	[1990LEI/PIL2]
	$\Delta_v H$	(361–522)	54.9 ± 0.2	298	EB	[1989STE/CHI2, 2008EME/KOZ]
	$\Delta_v H$	(357–435)	51.5	298	EB	[1988ISM/GAB, 2008EME/KOZ]
	$\Delta_v H$	(392–474)	48.2	407	A	[1987STE/MAL]
	$\Delta_v H$	(273–478)	54.6 ± 0.2	298	GS	[1980JAR/AFA, 2008EME/KOZ]
	$\Delta_v H$	(413–478)	53.1 ± 0.2	298	EB	[1980JAR/AFA, 2008EME/KOZ]
	$\Delta_v H$		48.0 ± 0.4	298	EB	[1980YEV/LEB, 2008EME/KOZ]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₄ H ₆ O ₂	[79-41-4]	2-methyl-2-propenoic acid				
	$\Delta_v H$		47.5 ± 0.4	298	C	[1996VAN/YU]
	$\Delta_v H$	(321–435)	53.9	336	A	[1987STE/MAL]
	$\Delta_v H$	(298–434)	51.6	313	A	[1987STE/MAL, 1947STU]
C ₄ H ₆ O ₂	[96-33-3]	methyl acrylate				
	$\Delta_{\text{fus}} H$		9.73	197.5		[1996DOM/HEA]
	$\Delta_v H$	(316–354)	34.2	331	A	[1987STE/MAL]
	$\Delta_v H$	(299–337)	28.8	314	BG	[1971HAL/BAL]
	$\Delta_v H$	(229–353)	38	244		[1947STU]
C ₄ H ₆ O ₂	[108-05-4]	vinyl acetate				
	$\Delta_{\text{fus}} H$		8.46	180.6		[1997KUL/LEB]
	$\Delta_v H$	(294–346)	34.4	309	A	[1987STE/MAL, 1971DYK, 1963CAP/FRI, 1984BOU/FRI]
	$\Delta_v H$	(340–355)	31.4	348		[1965SWA/VAN]
C ₄ H ₆ O ₂	[na]	α -methylacrylic acid				
	$\Delta_{\text{fus}} H$		8.06	287.5		[1996DOM/HEA]
C ₄ H ₆ O ₂	[na]	<i>cis</i> -crotonic acid				
	$\Delta_{\text{fus}} H$		12.57	344.4		[1991ACR]
C ₄ H ₆ O ₂ S	[3232-39-1]	diacetyl sulfide				
	$\Delta_v H$	(325–355)	54.2	340		[1999DYK/SVO]
	$\Delta_v H$	(325–355)	50.9	340	A	[1987STE/MAL]
C ₄ H ₆ O ₂ S	[77-77-0]	divinyl sulfone				
	$\Delta_v H$		56.4 ± 0.9	298	C	[1989VOR/KLY]
	$\Delta_v H$		56.5 ± 0.8	298		[1969MAC/MCN]
C ₄ H ₆ O ₃	[108-24-7]	acetic anhydride				
	$\Delta_v H$	(349–429)	43.3	364	EB	[1987AMB/GHI]
	$\Delta_v H$	(413–526)	47.6	428	A	[1987STE/MAL]
	$\Delta_v H$	(320–413)	45.5	335	A	[1987STE/MAL, 1971DYK]
	$\Delta_v H$	(336–412)	44.2	351		[1959MCD/SHR]
C ₄ H ₆ O ₃	[108-32-7]	propylene carbonate				
	$\Delta_{\text{fus}} H$		8.96	220.3	DSC	[2004DIN]
	$\Delta_{\text{fus}} H$	(13–300)	8.01	218.7	AC	[1994FUJ/OGU]
	$\Delta_{\text{fus}} H$		9.62	218.2		[1974VAS/KOR]
	$\Delta_v H$	(298–345)	61.5 ± 0.3	298	GS	[2008VER/TOK]
	$\Delta_v H$	(294–473)	71.3	298		[2005NAS/NEU]
	$\Delta_v H$	(460–513)	61.3 ± 0.1	298	E	[2004CHE/CLE, 2008VER/TOK]
	$\Delta_v H$	(668–762)	71.2 ± 0.6	298		[2002WIL/VON, 2008VER/TOK]
	$\Delta_v H$	(412–466)	54.4	427	A	[1987STE/MAL]
	$\Delta_v H$	(368–462)	57.8	383	EB	[1982HON/WAK]
	$\Delta_v H$	(368–462)	55.2	423	EB	[1982HON/WAK]
	$\Delta_v H$	(368–462)	53.0	443	EB	[1982HON/WAK]
	$\Delta_v H$	(293–353)	55.2	323		[1972GRA/SAL]
$\Delta_v H$	(323–370)	33.8	338	A, MM	[1987STE/MAL, 1971CHO/JON]	
C ₄ H ₆ O ₃	[3041-16-5]	<i>p</i> -dioxanone				
	$\Delta_{\text{fus}} H$		16.14	301.7		[1995LEB/BYK]
C ₄ H ₆ O ₄	[553-90-2]	dimethyl oxalate				
	$\Delta_{\text{fus}} H$		21.07	327.6		[1991ACR]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
	Enthalpy							
	$\Delta_{\text{sub}}H$				74.6 ± 0.7	298	C	[1996CHI/SAB]
	$\Delta_{\text{sub}}H$	(268–298)			75.6 ± 1.6	283	HSA	[1996CHI/SAB]
	$\Delta_{\text{sub}}H$				75.3 ± 1.6	298		[1996CHI/SAB]
	$\Delta_{\text{sub}}H$				74.9 ± 0.6		B	[1996CHI/SAB]
	$\Delta_{\text{sub}}H$				47.4 ± 0.5		BG	[1976ANT/CAR2, 1975ANT/CAR]
	$\Delta_{\text{v}}H$	(330–365)			54.7 ± 0.3	298	GS	[2006VER/KOZ]
	$\Delta_{\text{v}}H$	(328–443)			52.5	298		[2004MA/LIU, 2006VER/KOZ]
	$\Delta_{\text{v}}H$	(347–485)			44.7	416	HG, EB	[1988ASK/DAU]
	$\Delta_{\text{v}}H$	(293–437)			48.8	308	A	[1987STE/MAL, 1947STU]
C₄H₆O₄	[110-15-6]	succinic acid						
	$\Delta_{\text{fus}}H$				34	455.2		[2005ROU/TEM]
	$\Delta_{\text{fus}}H$				32.95	457		[1991ACR]
	$\Delta_{\text{sub}}H$	(318–358)			128 ± 2		TPD	[2007CAP/LOV]
	$\Delta_{\text{sub}}H$	(280–302)			119.5		TPTD	[2005CHA/ZIE]
	$\Delta_{\text{sub}}H$	(356–376)			120.5	368	TE,ME	[1983DEW/VAN]
	$\Delta_{\text{sub}}H$				123.1	298		[1983DEW/VAN]
	$\Delta_{\text{sub}}H$	(372–401)			118.1 ± 3.3	386	ME	[1970COX/PIL, 1960DAV/THO]
	$\Delta_{\text{sub}}H$				120.3 ± 4.4	298		[1970COX/PIL, 1960DAV/THO]
	$\Delta_{\text{sub}}H$				121.8 ± 3.3	298		[1960DAV/THO, 1999RIB/MON]
	$\Delta_{\text{sub}}H$	(292–320)			73.6	306	A	[1947GRA]
	$\Delta_{\text{v}}H$	(424–503)			94.4	298	CGC	[2005ROU/TEM]
C₄H₆O₄	[516-05-2]	methylmalonic acid						
	$\Delta_{\text{sub}}H$				117.4 ± 1.9	298	ME	[2000RIB/MON]
	$\Delta_{\text{sub}}H$				113.2 ± 0.4		C	[1983ALT/PIL]
C₄H₆O₄	[931-40-8]	4-hydroxymethyl)-1,3-dioxolan-2-one						
	$\Delta_{\text{v}}H$	(330–398)			85.4 ± 0.4	298	GS	[2008VER/TOK]
	$\Delta_{\text{v}}H$	(430–455)			87.8 ± 0.4	298	EB	[2002WIL/VON, 2008VER/TOK]
C₄H₆O₅	[na]	<i>(dl)</i> malic acid I						
	$\Delta_{\text{fus}}H$ (I)				33.52	402		[1996DOM/HEA]
	$\Delta_{\text{fus}}H$ (II)				30.17	396		[1996DOM/HEA]
C₄H₆O₅	[na]	<i>(d)</i> malic acid						
	$\Delta_{\text{fus}}H$				23.01	376		[1976LEC/COL]
C₄H₆O₆	[147-73-9]	<i>meso</i> tartaric acid						
	$\Delta_{\text{sub}}H$				156.9			[1983DEW/BOW]
C₄H₆O₆	[na]	<i>(d)</i> -tartaric acid						
	$\Delta_{\text{fus}}H$				32.3	445.1		[1998MUR/BET]
C₄H₆S	[5954-75-6]	2-vinylthiirane						
	$\Delta_{\text{v}}H$	(273–335)			38.7	288	A	[1987STE/MAL, 1999DYK/SVO]
C₄H₆S	[627-51-0]	divinyl sulfide						
	$\Delta_{\text{v}}H$				38.3 ± 0.7	298	C	[1989VOR/KLY]
C₄H₆S₃	[1748-15-8]	1,3-dithian-2-thione						
	$\Delta_{\text{sub}}H$	(321–348)			88.6	335		[1967GEI/SCH]
	$\Delta_{\text{sub}}H$				91.4 ± 2.5	298		[1967GEI/SCH, 1970COX/PIL]
C₄H₇Br	[31849-78-2]	<i>cis</i> 1-bromo-1-butene						
	$\Delta_{\text{v}}H$	(280–397)			35.1	295	A	[1987STE/MAL, 1971DYK]
	$\Delta_{\text{v}}H$	(229–359)			36.5	244		[1947STU]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
C₄H₇Br	[32620-08-9] $\Delta_{\text{v}}H$	<i>trans</i> 1-bromo-1-butene (234–368)	36.1	249	A	[1987STE/MAL, 1947STU]
C₄H₇Br	[23074-36-4] $\Delta_{\text{v}}H$ $\Delta_{\text{v}}H$	2-bromo-1-butene (276–391) (226–354)	34.5 36.1	291 241	A	[1987STE/MAL, 1971DYK] [1947STU]
C₄H₇Br	[3017-71-8] $\Delta_{\text{v}}H$	<i>cis</i> 2-bromo-2-butene (234–367)	36.5	249	A	[1987STE/MAL, 1947STU]
C₄H₇Br	[3017-68-3] $\Delta_{\text{v}}H$	<i>trans</i> 2-bromo-2-butene (228–359)	35.7	243	A	[1987STE/MAL, 1947STU]
C₄H₇BrO	[816-40-0] $\Delta_{\text{v}}H$ $\Delta_{\text{v}}H$	1-bromo-2-butanone (322–428) (279–420)	49.9 47.7	337 294	A	[1987STE/MAL] [1947STU]
C₄H₇BrO	[814-75-5] $\Delta_{\text{v}}H$	3-bromo-2-butanone (306–409)	46.4	321	A	[1987STE/MAL]
C₄H₇BrO	[2736-37-0] $\Delta_{\text{v}}H$	isobutyryl bromide (286–436)	45.7	301	A	[1987STE/MAL, 1947STU]
C₄H₇Br₃	[62127-48-4] $\Delta_{\text{v}}H$	1,3-dibromo-2-(bromomethyl)propane (475–660)	66.1	490	A	[1987STE/MAL]
C₄H₇Br₃	[3675-68-1] $\Delta_{\text{v}}H$	1,1,2-tribromobutane (361–490)	49.4	376	A	[1987STE/MAL]
C₄H₇Br₃	[3675-69-2] $\Delta_{\text{v}}H$ $\Delta_{\text{v}}H$	1,2,2-tribromobutane (356–487) (314–486)	48.4 50.7	371 329	A	[1987STE/MAL] [1947STU]
C₄H₇Br₃	[632-05-3] $\Delta_{\text{v}}H$ $\Delta_{\text{v}}H$	1,2,3-tribromobutane (394–546) (318–489)	54.1 51.3	409 333	A	[1987STE/MAL, 1971DYK] [1947STU]
C₄H₇Br₃	[38300-67-3] $\Delta_{\text{v}}H$	1,2,4-tribromobutane (390–541)	53.5	405	A	[1987STE/MAL, 1971DYK]
C₄H₇Br₃	[62127-47-3] $\Delta_{\text{v}}H$	2,2,3-tribromobutane (311–480)	51.7	326	A	[1987STE/MAL, 1947STU]
C₄H₇Cl	[513-37-1] $\Delta_{\text{v}}H$	1-chloro-2-methyl-1-propene (285–343)	33.2	300	A	[1987STE/MAL]
C₄H₇Cl	[563-47-3] $\Delta_{\text{v}}H$	3-chloro-2-methyl-1-propene (285–348)	33.3	300	A	[1987STE/MAL]
C₄H₇ClO	[616-27-3] $\Delta_{\text{v}}H$	1-chloro-2-butanone (307–411)	49.2	322	A	[1987STE/MAL, 1971DYK]
C₄H₇ClO	[4091-39-8] $\Delta_{\text{v}}H$	3-chloro-2-butanone (313–389)	38.8	328	A	[1987STE/MAL]
C₄H₇ClO	[40605-42-3] $\Delta_{\text{v}}H$	3-chloro-2-butene-1-ol (345–437)	50	360	A	[1987STE/MAL]
C₄H₇ClO₂	[109-61-5] $\Delta_{\text{v}}H$ $\Delta_{\text{v}}H$	propyl chloroformate (293–303)	40.7 ± 0.4 40.7 ± 0.4	298 298	C	[1990DAV/FIN] [1990DAV/FIN]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₄ H ₇ ClO ₂	[105-39-5]	ethyl chloroacetate				
	$\Delta_v H$	(274–418)	45	289	A	[1987STE/MAL]
	$\Delta_v H$		49.5 ± 0.1	298	C	[1972LAY/WAD]
		(298–418)	48.5	313		[1928NEL2, 1984BOU/FRI]
C ₄ H ₇ ClS	[760996-44-9]	2-butene-3-chloro-1-thiol				
	$\Delta_v H$	(341–397)	48.2	356	A	[1987STE/MAL]
C ₄ H ₇ Cl ₂ O ₄ P	[62-73-7]	dimethyl-(2,2-dichlorovinyl) phosphate				
	$\Delta_v H$	(283–387)	68	298	A	[1987STE/MAL]
C ₄ H ₇ Cl ₃	[18338-40-4]	1,2,3-trichlorobutane				
	$\Delta_v H$	(273–442)	41.3	288	A	[1987STE/MAL, 1947STU]
C ₄ H ₇ FOS	[462-31-7]	2-fluoroethyl thioacetate				
	$\Delta_v H$	(273–333)	44.7	288	A, GS	[1987STE/MAL, 1948RED/CHA4]
C ₄ H ₇ FO ₂	[459-72-3]	ethyl fluoroacetate				
	$\Delta_v H$	(273–333)	41.9	288	A, GS	[1987STE/MAL, 1948RED/CHA4, 1971DYK]
C ₄ H ₇ F ₃	[460-34-4]	1,1,1-trifluorobutane				
	$\Delta_v H$	(226–320)	28.1	241	A	[1987STE/MAL, 1971DYK]
C ₄ H ₇ IO ₂	[623-48-3]	ethyl iodoacetate				
	$\Delta_v H$	(301–362)	52.1	316	A	[1987STE/MAL, 1947GOU/HOL]
C ₄ H ₇ N	[78-82-0]	isobutyronitrile				
	$\Delta_v H$	(324–354)	35.9	339		[1979SUK/VLA]
	$\Delta_v H$	(303–352)	37.5	321	BG	[1971HAL/BAL]
C ₄ H ₇ N	[109-74-0]	butyronitrile				
	$\Delta_v H$		39.2 ± 0.1	298	C	[1982FUC/HAL]
	$\Delta_v H$	(303–493)	38.8	318	EB	[1971MEY/REN]
	$\Delta_v H$	(332–401)	37.7	347	A, EB	[1987STE/MAL, 1947STU, 1973MEY/HOT]
	$\Delta_v H$		37.0	298	EB	[1959EVA/SKI, 2005EME/VER]
	$\Delta_v H$	(294–415)	40.5 ± 0.2	298	MM	[1933HEI, 2005EME/VER]
C ₄ H ₇ NO	[75-86-5]	acetone cyanohydrin				
	$\Delta_v H$	(355–393)	106.5	370	A	[1987STE/MAL]
C ₄ H ₇ NO	[4476-02-2]	2-hydroxybutyronitrile				
	$\Delta_v H$	(314–452)	57.9	329	A	[1987STE/MAL, 1947STU]
C ₄ H ₇ NO	[1120-64-5]	2-methyl-2-oxazoline				
	$\Delta_v H$		39.1 ± 0.3	298	C	[1976HAM/THO]
C ₄ H ₇ NO	[79-39-0]	methacrylamide				
	$\Delta_{\text{fus}} H$		15.0	385.1		[1996DOM/HEA]
	$\Delta_v H$	(390–418)	86.3	404	A	[1987STE/MAL]
C ₄ H ₇ NO	[62957-60-2]	ethoxyacetonitrile				
	$\Delta_v H$	(273–313)	46.5 ± 0.3	298	GS	[1995VER/BEC]
C ₄ H ₇ NO	[33695-59-9]	3-methoxypropionitrile				
	$\Delta_v H$	(328–438)	47.6	343	A	[1987STE/MAL]
C ₄ H ₇ NO	[616-45-5]	2-pyrrolidone				
	$\Delta_{\text{fus}} H$		13.92	299		[1996DOM/HEA]
	$\Delta_v H$	(391–484)	73.6 ± 1.3	298	EB, BG	[1998MOR/KOP]
	$\Delta_v H$		41.7 ± 0.6			[1995VIE/CDE]
	$\Delta_v H$	(395–518)	60	410	A	[1987STE/MAL]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₄ H ₇ NO	[31110-30-2]	<i>cis</i> 2-butenic acid amide				
	$\Delta_{\text{sub}}H$	(353–387)	68.0	368	A	[1987STE/MAL]
C ₄ H ₇ NO	[625-37-6]	<i>trans</i> 2-butenic acid amide				
	$\Delta_{\text{sub}}H$	(363–413)	80.0	378	A	[1987STE/MAL]
C ₄ H ₇ NOS	[17374-18-4]	tetrahydro-2 <i>H</i> -1,3-oxazine-2-thione				
	$\Delta_{\text{fus}}H$		18.4	400.2	DSC	[2008TEM/ROU3]
C ₄ H ₇ NO ₂	[625-77-4]	diacetamide				
	$\Delta_{\text{sub}}H$		73.2 ± 0.8	298	C	[1965WAD, 1971MOR]
	Δ_vH	(368–496)	59.7	383	A	[1987STE/MAL]
	Δ_vH	(343–496)	64.6	358		[1947STU]
C ₄ H ₇ NO ₂	[2783-12-2]	2-nitro-1-butene				
	Δ_vH	(273–333)	44.0	288	A	[1987STE/MAL, 1971DYK]
C ₄ H ₇ NO ₂	[22677-21-0]	(R)-4-hydroxy-2-pyrrolidone				
	$\Delta_{\text{fus}}H$		23.41	392	DSC	[2004WAN/WIE]
C ₄ H ₇ NO ₂	[68108-18-9]	(-) 4-hydroxy-2-pyrrolidone				
	$\Delta_{\text{fus}}H$		28.49	429.8	DSC	[1999LI/ZEL]
C ₄ H ₇ NO ₂	[25747-41-5]	(+) 4-hydroxy-2-pyrrolidone				
	$\Delta_{\text{fus}}H$		26.74	394.8	DSC	[1999LI/ZEL]
C ₄ H ₇ NO ₃	[543-24-8]	N-acetylglycine				
	$\Delta_{\text{sub}}H$		127.0 ± 1.0	389	TE,ME	[1979DEK/VOO]
C ₄ H ₇ NO ₄	[56-84-8]	L-aspartic acid				
	$\Delta_{\text{sub}}H$	(370–470)	U 96 ± 4.2	420	LE	[1977GAF/PIE]
C ₄ H ₇ N ₃ O ₉	[6859-60-5]	1,2,4-butanetriol trinitrate				
	Δ_vH	(293–313)	60.0 ± 11.3	303	A, GS	[1987STE/MAL, 1957MCC/DOU]
C ₄ H ₈	[106-98-9]	1-butene				
	$\Delta_{\text{fus}}H$		3.85	87.8		[1996DOM/HEA]
	Δ_vH	(200–274)	23.3	259	A	[1987STE/MAL]
	Δ_vH	(126–192)	28.3	177	A	[1987STE/MAL]
	Δ_vH	(267–345)	22.8	282	A	[1987STE/MAL]
	Δ_vH	(342–411)	22.0	357	A	[1987STE/MAL]
	Δ_vH	(267–411)	22.5	282	A	[1987STE/MAL]
	Δ_vH		20.1	298		[1971WIL/ZWO]
	Δ_vH		25.3	202		[1946AST/FIN]
	Δ_vH		24.5	219		[1946AST/FIN]
	Δ_vH		23.3	242		[1946AST/FIN]
	Δ_vH		21.9	267		[1946AST/FIN]
	Δ_vH	(216–273)	23.2	258		[1940LAM/ROP, 1984BOU/FRI]
C ₄ H ₈	[590-18-1]	<i>cis</i> 2-butene				
	$\Delta_{\text{fus}}H$		7.31	134.3		[1996DOM/HEA]
	Δ_vH	(221–290)	24.4	275	A	[1987STE/MAL]
	Δ_vH	(276–325)	24.0	291	A	[1987STE/MAL]
	Δ_vH	(324–386)	23.6	339	A	[1987STE/MAL]
	Δ_vH	(383–431)	23.6	398	A	[1987STE/MAL]
	Δ_vH		22.1	298		[1971WIL/ZWO]
	Δ_vH		22.5	246	C	[1944SCO/FER]
Δ_vH	(195–267)	25.3	252		[1942BEN, 1984BOU/FRI]	

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound					
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference	
C ₄ H ₈	[624-64-6]	<i>trans</i> 2-butene					
	$\Delta_{\text{fus}}H$		9.76	167.6		[1996DOM/HEA]	
	Δ_vH	(205–287)	23.9	272	A	[1987STE/MAL]	
	Δ_vH	(273–315)	23.6	288	A	[1987STE/MAL]	
	Δ_vH	(313–385)	23.3	328	A	[1987STE/MAL]	
	Δ_vH	(382–428)	23.2	397	A	[1987STE/MAL]	
	Δ_vH		21.3	298		[1971WIL/ZWO]	
	Δ_vH		22.8 ± 0.1	274	C	[1945GUT/PIT]	
C ₄ H ₈	[287-23-0]	cyclobutane					
	$\Delta_{\text{trs}}H$		5.71	145.7			
	$\Delta_{\text{fus}}H$		1.09	182.4		[1996DOM/HEA]	
	$\Delta_{\text{sub}}H$		36.4	145	B	[1963BON]	
	Δ_vH	(198–287)	25.2	272	A	[1987STE/MAL]	
	Δ_vH	(217–285)	25.2	270		[1953RAT/GWI, 1984BOU/FRI]	
	C ₄ H ₈	[594-11-6]	cyclobutane				
		Δ_vH	(177–278)	24.8	263	A	[1987STE/MAL, 1947STU]
C ₄ H ₈	[115-11-7]	2-methylpropene					
	$\Delta_{\text{fus}}H$		5.92	132.4		[1996DOM/HEA]	
	Δ_vH	(212–279)	23.1	264	A	[1987STE/MAL]	
	Δ_vH	(266–313)	22.7	281	A	[1987STE/MAL]	
	Δ_vH	(310–376)	22.2	325	A	[1987STE/MAL]	
	Δ_vH	(371–418)	22.3	386	A	[1987STE/MAL]	
	Δ_vH		20.6	298		[1971WIL/ZWO]	
C ₄ H ₈ BrClO	[51070-66-7]	2-bromoethyl 2-chloroethyl ether					
	Δ_vH	(309–469)	53.3	324	A	[1987STE/MAL, 1947STU]	
	C ₄ H ₈ Br ₂	[62168-25-6]	1,1-dibromobutane				
		Δ_vH	(342–477)	45.8	357	A, EST	[1987STE/MAL, 1956MAN, 1971DYK]
	C ₄ H ₈ Br ₂	[533-98-2]	1,2-dibromobutane				
		Δ_vH	(338–425)	43.5	353	A	[1987STE/MAL]
		Δ_vH	(330–425)	45.9	298		[1975PIS/ROZ2, 1975PIS/ROZ]
Δ_vH			45.6 ± 0.7	298	EB	[1975PIS/ROZ]	
Δ_vH		(281–439)	42.8	296	A	[1987STE/MAL, 1947STU]	
Δ_vH		(273–333)	45.1	300		[1941LIS]	
C ₄ H ₈ Br ₂	[107-80-2]	1,3-dibromobutane					
	Δ_vH	(351–450)	44.7	366	A	[1987STE/MAL]	
C ₄ H ₈ Br ₂	[110-52-1]	1,4-dibromobutane					
	Δ_vH		52.6	298	GC	[1994CAR/LAY]	
	Δ_vH	(375–520)	51.4	390	A	[1987STE/MAL, 1971DYK]	
C ₄ H ₈ Br ₂	[5780-13-2]	<i>meso</i> 2,3-dibromobutane					
	Δ_vH	(274–431)	41.7	289	A	[1987STE/MAL, 1947STU]	
	C ₄ H ₈ Br ₂	[598-71-0]	<i>threo</i> 2,3-dibromobutane				
Δ_vH		(278–434)	40.9	293	A	[1987STE/MAL]	

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₄ H ₈ Br ₂	[594-34-3]	1,2-dibromo-2-methylpropane				
	$\Delta_v H$		43.3 ± 0.1		C	[1974SUN/WUL]
		(244–422)	33.3	259	A	[1987STE/MAL, 1947STU]
C ₄ H ₈ Br ₂	[28148-04-1]	1,3-dibromo-2-methylpropane				
	$\Delta_v H$	(287–448)	45.1	302	A	[1987STE/MAL, 1947STU]
C ₄ H ₈ Br ₂ O	[5414-19-7]	<i>bis</i> (2-bromoethyl) ether				
	$\Delta_v H$	(320–486)	55.1	335	A	[1987STE/MAL, 1947STU]
C ₄ H ₈ Br ₂ O ₂	[na]	<i>(dl)</i> 2,3-dibromo-1,4-butanediol				
	$\Delta_{\text{fus}} H$		29.29	363.2		[1981CHI/GAR]
C ₄ H ₈ Br ₂ O ₂	[na]	<i>(d)</i> 2,3-dibromo-1,4-butanediol				
	$\Delta_{\text{fus}} H$		33.89	388.2		[1981CHI/GAR]
C ₄ H ₈ Cl ₂	[541-33-3]	1,1-dichlorobutane				
	$\Delta_v H$	(310–390)	39.5	298		[1991BAS/SVO]
	$\Delta_v H$	(304–386)	38.7	319	A	[1987STE/MAL]
	$\Delta_v H$		39.4 ± 0.6	298	EB	[1977PIS/ROZ]
	$\Delta_v H$	(303–428)	38.8	318	EST	[1987STE/MAL, 1956MAN]
C ₄ H ₈ Cl ₂	[616-21-7]	1,2-dichlorobutane				
	$\Delta_v H$		40.1 ± 0.1	298	C	[1992HE/AN]
	$\Delta_v H$		40.2 ± 0.1	298	C	[1989AN/HU]
	$\Delta_v H$	(312–394)	39.0	327	A	[1987STE/MAL]
	$\Delta_v H$	(310–390)	40.4	298		[1982ROO, 1991BAS/SVO]
	$\Delta_v H$		40.1 ± 0.6	298	EB	[1975PIS/ROZ2]
	$\Delta_v H$	(249–397)	38.1	264	A	[1987STE/MAL, 1947STU]
C ₄ H ₈ Cl ₂	[1190-22-3]	1,3-dichlorobutane				
	$\Delta_v H$		42.2 ± 0.1	298	C	[1992HE/AN]
	$\Delta_v H$	(320–400)	42.3	298		[1991BAS/SVO]
	$\Delta_v H$		42.3 ± 1.8	298	C	[1990AN/HE]
	$\Delta_v H$	(318–407)	40.5	333	A	[1987STE/MAL]
C ₄ H ₈ Cl ₂	[110-56-2]	1,4-dichlorobutane				
	$\Delta_v H$		46.7	298	GC	[1994CAR/LAY]
	$\Delta_v H$		46.5 ± 0.1	298	C	[1992HE/AN]
	$\Delta_v H$	(325–425)	46.4	298		[1991BAS/SVO]
	$\Delta_v H$		46.4 ± 0.1	298	C	[1990AN/HE]
	$\Delta_v H$		46.4 ± 0.1	298	C	[1989AN/HU]
	$\Delta_v H$	(336–425)	43.4	351	A	[1987STE/MAL, 1971DYK]
C ₄ H ₈ Cl ₂	[4279-22-5]	2,2-dichlorobutane				
	$\Delta_v H$		36.3 ± 0.1	298	C	[1992HE/AN]
	$\Delta_v H$	(300–370)	36.7	298		[1991BAS/SVO]
	$\Delta_v H$	(293–376)	36.4	308	A	[1987STE/MAL]
	$\Delta_v H$		33.7 ± 0.6	298	EB	[1977PIS/ROZ]
C ₄ H ₈ Cl ₂	[na]	<i>meso</i> 2,3-dichlorobutane				
	$\Delta_v H$		38.4	298	C	[1992HE/AN, 1993HE/AN]
C ₄ H ₈ Cl ₂	[na]	<i>(dl)</i> 2,3-dichlorobutane				
	$\Delta_v H$		39.7	298	C	[1993HE/AN]
C ₄ H ₈ Cl ₂	[7581-97-7]	2,3-dichlorobutane				
	$\Delta_v H$	(247–389)	39.6	262	A	[1987STE/MAL]
C ₄ H ₈ Cl ₂	[598-76-5]	1,1-dichloro-2-methylpropane				

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(242–379)	38.7	257	A	[1987STE/MAL]
C ₄ H ₈ Cl ₂	[594-37-6]	1,2-dichloro-2-methylpropane				
	$\Delta_v H$	(247–381)	40.4	262	A	[1987STE/MAL, 1947STU]
C ₄ H ₈ Cl ₂	[616-19-3]	1,3-dichloro-2-methylpropane				
	$\Delta_v H$	(270–408)	45.1	285	A	[1987STE/MAL, 1947STU]
C ₄ H ₈ Cl ₂ O	[111-44-4]	<i>bis</i> (2-chloroethyl) ether				
	$\Delta_{\text{fus}} H$		8.39	226.5		[1996DOM/HEA]
	$\Delta_v H$	(297–452)	49.8	312	A	[1987STE/MAL, 1947STU]
C ₄ H ₈ Cl ₂ S	[505-60-2]	<i>bis</i> (2-chloroethyl) sulfide				
	$\Delta_{\text{sub}} H$	(248–293)	80.9	271		[2006BUC/BUE]
	$\Delta_{\text{sub}} H$	(263–287)	77.2	275		[1987STE/MAL]
	$\Delta_{\text{sub}} H$		84.5		B	[1963BON, 1947BAL/DEN]
	$\Delta_v H$	(288–358)	59.6	303	A, MM	[1987STE/MAL, 1947BAL/DEN, 1984BOU/FRI, 1948BEN/FRA]
	$\Delta_v H$	(353–393)	50.3	373		[1943HOL/MEL]
C ₄ H ₈ Cl ₂ S ₃	[19149-77-0]	<i>bis</i> (2-chloroethyl) trisulfide				
	$\Delta_v H$	(293–333)	68.2	308	A, GS	[1987STE/MAL, 1948RED/CHA, 1999DYK/SVO]
C ₄ H ₈ Cl ₃ O ₄ P	[52-68-6]	(1-hydroxy-2,2,2-trichloroethyl)phosphonic acid dimethyl ester				
	$\Delta_{\text{fus}} H$		20.37	351		
	$\Delta_{\text{fus}} H$		22.4	357		
	$\Delta_{\text{fus}} H$		25.0	384		[1996DOM/HEA]
	$\Delta_{\text{sub}} H$	(293–357)	107		308	[1987STE/MAL]
C ₄ H ₈ F ₂	[353-81-1]	1,1-difluorobutane				
	$\Delta_v H$	(246–347)	31	261	A, EST	[1987STE/MAL, 1956MAN, 1971DYK]
C ₄ H ₈ F ₂	[353-81-1]	2,2-difluorobutane				
	$\Delta_v H$	(238–336)	30	253	A	[1987STE/MAL, 1971DYK]
C ₄ H ₈ F ₂ O	[184899-81-8]	1,1,1,2,2,3,3-heptafluoro-3-(fluoromethoxy)propane				
	$\Delta_v H$	(283–316)	31	298	I	[2002MUR/YAM]
C ₄ H ₈ F ₂ O ₄ S	[381-46-4]	<i>bis</i> (2-fluoroethyl) sulfate				
	$\Delta_v H$	(273–333)	63.9	288	A, GS	[1987STE/MAL, 1948RED/CHA4, 1999DYK/SVO]
C ₄ H ₈ I ₂	[628-21-7]	1,4-diiodobutane				
	$\Delta_v H$		59	298	GC	[1994CAR/LAY]
C ₄ H ₈ N ₂	[926-64-7]	(dimethylamino)acetonitrile				
	$\Delta_v H$	(277–307)	45.4 ± 0.6		GS	[1997WEL/VER]
C ₄ H ₈ N ₂	[1606-49-1]	1,4,5,6-tetrahydropyrimidine				
	$\Delta_v H$	(330–395)	75.6 ± 2.0	298	IP	[1996STE/CHI3]
C ₄ H ₈ N ₂ O	[1852-17-1]	tetrahydro-2-pyrimidone				
	$\Delta_{\text{sub}} H$	(363–385)	113.4 ± 0.7	298	ME	[2008RIB/RIB]
	$\Delta_{\text{sub}} H$		89.3	298		[1999DEF/DEO]
C ₄ H ₈ N ₂ O ₂	[3148-73-0]	1,2-diacetylhydrazine				
	$\Delta_{\text{sub}} H$	(347–358)	103.1 ± 1.7	352.5	A	[1987STE/MAL, 1959TAK/SHI]
C ₄ H ₈ N ₂ O ₂	[95-45-4]	dimethylglyoxime				
	$\Delta_{\text{sub}} H$	(331–352)	96.8	341.5	A	[1987STE/MAL]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
	$\Delta_{\text{sub}}H$		97.1 ± 2.1			[1956SEK/SUZ, 1970COX/PIL, 1960JON]
C ₄ H ₈ N ₂ O ₂	[2620-63-5]	N-acetylglycine amide				
	$\Delta_{\text{fus}}H$		25.6	408.2		[1996DOM/HEA]
	$\Delta_{\text{sub}}H$		123.5 ± 1.7	376	C	[1999DEL/BAR]
	$\Delta_{\text{sub}}H$		126.3 ± 2.3			[1999DEL/BAR]
	$\Delta_{\text{sub}}H$	(378–406)	140.2 ± 2.3	298	C	[1995DEL/SAB]
	$\Delta_{\text{sub}}H$		135 ± 3	392	TE	[1988FER/DEL, 1986BAR/FER]
C ₄ H ₈ N ₂ O ₂	[110-14-5]	succinamide				
	$\Delta_{\text{fus}}H$		6.08	485.9	DSC	[2006BAD/DEL]
C ₄ H ₈ N ₂ O ₂	[59-82-2]	N-nitrosomorpholine				
	$\Delta_{\text{vap}}H$		50.2	343		[1988SOL/SIT]
C ₄ H ₈ N ₂ O ₆	[6423-44-5]	1,3-butanediol dinitrate				
	$\Delta_{\text{v}}H$	(293–313)	71.4 ± 7.1	303	A, GS	[1987STE/MAL, 1957KEM/GOL]
C ₄ H ₈ N ₂ O ₆	[3457-91-8]	1,4-butanediol dinitrate				
	$\Delta_{\text{v}}H$	(293–313)	57.4 ± 0.8	303	A, GS	[1987STE/MAL, 1957KEM/GOL]
C ₄ H ₈ N ₂ O ₇	[693-21-0]	diethyleneglycol dinitrate				
	$\Delta_{\text{fus}}H$		25.4	276.5		[2000URY/KUP]
	$\Delta_{\text{v}}H$	(293–333)	94.3	308	A	[1987STE/MAL]
C ₄ H ₈ N ₄ O ₂	[140-79-4]	1,4-dinitrosopiperazine				
	$\Delta_{\text{sub}}H$	(325–360)	101.3 ± 8	343		[1974PEP/MAT, 1977PED/RYL]
C ₄ H ₈ N ₄ O ₄	[4164-37-8]	1,4-dinitropiperazine				
	$\Delta_{\text{fus}}H$		200.8	489.2		[2001OXL/SMI]
	$\Delta_{\text{fus}}H$		33.93	489.6	DSC	[1997ZEM]
	$\Delta_{\text{sub}}H$	(325–360)	111.3 ± 8	343		[1974PEP/MAT, 1977PED/RYL]
		Note: Enthalpy of fusion is abnormally high, compound may be decomposing				
C ₄ H ₈ N ₄ O ₄	[5754-89-2]	1,3-dinitro-1,3-diazacyclohexane				
	$\Delta_{\text{trs}}H$		15.8	343		
	$\Delta_{\text{fus}}H$		2.97	354		[1991PIC/RYL]
C ₄ H ₈ N ₆ O ₅	[5800-63-5]	1,5-dinitro-3-nitroso-1,3,5-triazacycloheptane				
	$\Delta_{\text{trs}}H$		25.7	404		
	$\Delta_{\text{fus}}H$		2.9	440		[1991PIC/RYL]
C ₄ H ₈ N ₆ O ₆	[5790-78-3]	1,3,5-trinitro-1,3,5-triazacycloheptane				
	$\Delta_{\text{fus}}H$		27.74	435.9	DSC	[1997ZEM]
C ₄ H ₈ N ₈ O ₈	[2691-41-0]	1,3,5,7-tetranitro-1,3,5,7-tetrazacyclooctane				
	$\Delta_{\text{fus}}H(\delta)$		69.87	553.2		[1996DOM/HEA]
	$\Delta_{\text{sub}}H$	(461–487)	161.9	474		[1976TAY/CRO]
	$\Delta_{\text{sub}}H(\delta)$	(415–479)	161 ± 0.3	447		[1978CUN/PAL]
	$\Delta_{\text{sub}}H(\beta)$	(371–403)	175.2	385		[1969ROS/DIC]
C ₄ H ₈ N ₁₂ O ₆	[62209-57-8]	1,7-diazido-2,4,6-trinitro-2,4,6-triazahexane				
	$\Delta_{\text{fus}}H$		40.17	406		[1987OYU/BRI]
C ₄ H ₈ O	[2919-23-5]	cyclobutanol				
	$\Delta_{\text{fus}}H$		8.53	228.4		[1982DWO/FUC]
C ₄ H ₈ O	[78-93-3]	2-butanone				
	$\Delta_{\text{fus}}H$		8.44	186.5		[1991ACR]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(294–342)	34.6	309	A	[1987STE/MAL]
	$\Delta_v H$	(353–403)	32.5	368	A	[1987STE/MAL]
	$\Delta_v H$	(397–479)	31.6	412	A	[1987STE/MAL]
	$\Delta_v H$	(473–537)	31.1	488	A	[1987STE/MAL]
	$\Delta_v H$		34.8 ± 0.1	298	C	[1983UCH/MAJ]
	$\Delta_v H$		34.5 ± 0.1	298	C	[1979SUN/SVE2]
	$\Delta_v H$	(258–362)	35.6	273		[1978CAV/CHA]
	$\Delta_v H$		34.7			[1975AMB/ELL]
	$\Delta_v H$	(315–363)	33.9	330	A, EB, GS	[1987STE/MAL, 1975AMB/ELL, 1965COL/COU]
	$\Delta_v H$		33.8	315	C	[1973GEI/QUI]
	$\Delta_v H$		33.8 ± 0.1	314	C	[1961NIC/KOB]
	$\Delta_v H$		32.3 ± 0.1	338	C	[1961NIC/KOB]
	$\Delta_v H$		31.3 ± 0.1	352	C	[1961NIC/KOB]
	$\Delta_v H$		30.5 ± 0.1	363	C	[1961NIC/KOB]
	$\Delta_v H$		30.0 ± 0.1	370	C	[1961NIC/KOB]
	$\Delta_v H$	(314–370)	33.9	329		[1947STU]
C₄H₈O	[513-42-8]	2-methyl-2-propen-1-ol				
	$\Delta_v H$	(323–373)	51.9	298	CGC	[1995CHI/HOS]
C₄H₈O	[627-27-0]	3-buten-1-ol				
	$\Delta_v H$	(343–393)	50.8 ± 0.0	298	CGC	[2005VAL/QUI]
	$\Delta_v H$		50.9 ± 0.1	313	C	[1996ULB/KLU]
	$\Delta_v H$		48.8 ± 0.1	328	C	[1996ULB/KLU]
	$\Delta_v H$		46.7 ± 0.1	343	C	[1996ULB/KLU]
C₄H₈O	[6118-14-5]	<i>(dl)</i> 3-buten-2-ol				
	$\Delta_v H$	(304–370)	39.2	319	A	[1987STE/MAL]
C₄H₈O	[123-72-8]	butyraldehyde				
	$\Delta_{\text{fus}} H$		11.09	176.8		[1996DOM/HEA, 1989VAS/LEB]
	$\Delta_v H$	(313–353)	33.2	298	CGC	[1995CHI/HOS]
	$\Delta_v H$	(293–349)	34.2	308	A	[1987STE/MAL]
	$\Delta_v H$		33.7 ± 0.4	298	EB	[1967BUC/COX, 2003VER/KRA2]
	$\Delta_v H$	(330–348)	32.9	339	EB	[1963WOJ]
	$\Delta_v H$	(304–347)	33.3	319		[1959SEP/PAU, 1984BOU/FRI]
	$\Delta_v H$	(258–353)	33.9	306		[1938KUC]
C₄H₈O	[106-88-7]	<i>(dl)</i> 1,2-epoxybutane				
	$\Delta_v H$	(254–347)	24.7	269	A	[1987STE/MAL]
C₄H₈O	[558-30-5]	1,2-epoxy-2-methylpropane (2,2-dimethyloxirane)				
	$\Delta_v H$	(204–329)	30.6	219	A	[1987STE/MAL, 1947STU]
C₄H₈O	[109-92-2]	ethyl vinyl ether				
	$\Delta_v H$	(223–309)	29.5	238	A	[1987STE/MAL]
C₄H₈O	[78-84-2]	isobutyraldehyde				
	$\Delta_v H$	(313–353)	32.3	298	CGC	[1995CHI/HOS]
	$\Delta_v H$	(313–324)	31.4	318		[1984ENG/SAN]
	$\Delta_v H$	(309–337)	31.8	324		[1976BRA/PES]
	$\Delta_v H$	(333–347)	33.4	340	EB	[1963WOJ]
	$\Delta_v H$	(283–337)	32.8	298	A	[1987STE/MAL, 1959SEP/PAU, 1964SER/TIM]
C₄H₈O	[116-11-0]	2-methoxy-1-propene				
	$\Delta_v H$	(281–309)	28.3 ± 0.1	295		[1988BAG/GUR]
C₄H₈O	[4188-68-5]	<i>cis</i> methyl propenyl ether				

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound					
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference	
		Δ_vH	(293–318)	30.6	305	A	[1987STE/MAL]
C₄H₈O	[4188-69-6]	<i>trans</i> methyl propenyl ether					
		Δ_vH	(293–322)	29.5	307	A	[1987STE/MAL]
C₄H₈O	[109-99-9]	tetrahydrofuran					
		$\Delta_{\text{fus}}H$		8.54	164.8		[1991ACR]
		Δ_vH	(290–339)	32.3	305		[2001LOR/AUC]
		Δ_vH	(273–339)	33.1	288	A	[1987STE/MAL]
		Δ_vH	(399–479)	29	414	A	[1987STE/MAL]
		Δ_vH	(467–541)	29.6	482	A	[1987STE/MAL]
		Δ_vH		32	298	C	[1981HOS/SCO]
		Δ_vH	(235–340)	32.5 ± 0.2	288		[1976BOR/CHU]
		Δ_vH	(302–339)	30.8	320		[1975RIV]
		Δ_vH	(273–308)	32.8	288		[1970KOI/OUN, 1984BOU/FRI]
		Δ_vH	(296–373)	31.9	311		[1970SCO, 1984BOU/FRI]
		Δ_vH	(224–360)	32.9	298		[1970MOI/ANT]
		Δ_vH	(293–341)	U 26.9	308		[1959BIS/FIN]
		Δ_vH	(293–313)	31.8	313		[1948KLA/MOH, 1958CAS/FLE3]
C₄H₈OS	[1600-44-8]	tetrahydrothiophene 1-oxide					
		$\Delta_{\text{trs}}H$		5.85	223.9		
		$\Delta_{\text{fus}}H$		0.51	231.8	DSC	[1990HAI/GIL]
C₄H₈OS	[15980-15-1]	1,4-oxathiane					
		Δ_vH	(342–411)	42.1	378		[1999DYK/SVO]
		Δ_vH	(342–411)	44.8	357	A	[1987STE/MAL]
C₄H₈OS	[625-60-5]	S-ethyl thiolacetate					
		Δ_vH		40.0 ± 0.2	298	C	[1966WAD]
C₄H₈OS₂	[16487-10-8]	1,3-dithiane sulfoxide					
		$\Delta_{\text{fus}}H$		22.6	361.9	DSC	[2003ROU/TEM2]
C₄H₈O₂	[497-26-7]	2-methyl-1,3-dioxolane					
		Δ_vH	(270–308)	43.0 ± 0.6		GS	[1998VER/PEN, 2002VER]
C₄H₈O₂	[6117-80-2]	<i>cis</i> 2-butene-1,4-diol					
		Δ_vH	(373–508)	74.7	388	A	[1987STE/MAL]
C₄H₈O₂	[922-69-0]	1,1-dimethoxyethene					
		Δ_vH	(303–362)	39.6	333		[1995GUT/LIU]
C₄H₈O₂	[107-92-6]	butanoic acid					
		$\Delta_{\text{fus}}H$		11.07	264.7		[1991ACR]
		$\Delta_{\text{trs}}H$		1.04	NA		
		$\Delta_{\text{fus}}H$		11.59	268		[1982MAR/AND]
		$\Delta_{\text{sub}}H$	(238–255)	76.0 ± 1.5	248	TE,ME	[1978CAL/CAL]
		Δ_vH	(384–435)	52.5	399		[2004CLI/RAM]
		Δ_vH	(391–429)	50.3	406	EB	[2001MUN/KRA]
		Δ_vH	(303–378)	58.5	298	CGC	[2000VER2]
		Δ_vH	(278–308)	58.5 ± 0.3	293	GS	[2000VER2]
		Δ_vH	(278–308)	58.2 ± 0.3	298	GS	[2000VER2]
		Δ_vH	(353–393)	60.7	298	CGC	[1995CHI/HOS]
		Δ_vH	(437–592)	47.7	452	A	[1987STE/MAL]
		Δ_vH	(301–358)	51.1	316	A	[1987STE/MAL]
		Δ_vH	(355–453)	53.2	370	A	[1987STE/MAL, 1971DYK]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	Enthalpy							
		$\Delta_v H$ (monomer)			40.5 ± 0.1	298	C	[1970KON/WAD]
		$\Delta_v H$			58 ± 4	298	C	[1970KON/WAD]
		$\Delta_v H$	(363–436)		52	378		[1949DRE/SHR, 1949DRE/MAR, 1984BOU/FRI]
(C ₄ H ₈ O ₂) ₂	[19496-06-1]	butanoic acid dimer						
		$\Delta_{\text{sub}} H$	(238–255)		85 ± 1.5	248	TE,ME	[1978CAL/CAL]
C ₄ H ₈ O ₂	[505-22-6]	1,3-dioxane						
		$\Delta_v H$			39.1 ± 0.1	298	C	[1982BYS/MAN]
		$\Delta_v H$			35.6 ± 0.4			[1959FLE/MOR]
C ₄ H ₈ O ₂	[123-91-1]	1,4-dioxane						
		$\Delta_{\text{trs}} H$			2.35	272.9		
		$\Delta_{\text{fus}} H$			12.84	284.1		[1996DOM/HEA]
		$\Delta_{\text{sub}} H$	(237–272)		35.6	255	A	[1947STU]
		$\Delta_v H$	(285–375)		38	300	A	[1987STE/MAL]
		$\Delta_v H$	(329–372)		36.5	350		[1984CAS/FRA]
		$\Delta_v H$			38.6 ± 0.1	298	C	[1982BYS/MAN]
		$\Delta_v H$	(293–398)		37.3	308		[1963VIN/MAR, 1984BOU/FRI]
		$\Delta_v H$	(283–353)		37	318		[1936HOV/SCH]
C ₄ H ₈ O ₂	[141-78-6]	ethyl acetate						
		$\Delta_{\text{fus}} H$			10.48	189.3		[1991ACR]
		$\Delta_v H$	(300–390)		34.1	315		[1997HER/ORT]
		$\Delta_v H$	(313–353)		35.0	298	CGC	[1995CHI/HOS]
		$\Delta_v H$	(271–373)		36.7	286		[1981AMB/ELL, 1984BOU/FRI]
		$\Delta_v H$			35.6 ± 0.1	298	C	[1980SVO/UCH]
		$\Delta_v H$			34.6 ± 0.1	313	C	[1980SVO/UCH]
		$\Delta_v H$			31.4 ± 0.1	343	C	[1980SVO/UCH]
		$\Delta_v H$			33.8 ± 0.1	326	C	[1977SVO/VES]
		$\Delta_v H$			33.4 ± 0.1	331	C	[1977SVO/VES]
		$\Delta_v H$			32.4 ± 0.1	344	C	[1977SVO/VES]
		$\Delta_v H$			31.9 ± 0.1	351	C	[1977SVO/VES]
		$\Delta_v H$			31.0 ± 0.1	363	C	[1977SVO/VES]
		$\Delta_v H$			34.0	320		[1976CON/COU]
		$\Delta_v H$			31.9	350		[1976CON/COU]
		$\Delta_v H$			35.1 ± 0.2	298	C	[1966WAD]
		$\Delta_v H$	(288–351)		35.7	303	A	[1987STE/MAL, 1965MER/POL, 1971DYK]
C ₄ H ₈ O ₂	[513-86-0]	3-hydroxy-2-butanone (acetoin)						
		$\Delta_v H$	(363–393)		48.7 ± 0.4	298	CGC	[2005TEM/CHI]
		$\Delta_v H$	(273–418)		38.4	288	A	[1987STE/MAL]
C ₄ H ₈ O ₂	[79-31-2]	2-methylpropanoic acid						
		$\Delta_v H$	(375–426)		50.5	390		[2004CLI/RAM]
		$\Delta_v H$	(303–378)		56.3	298	CGC	[2000VER2]
		$\Delta_v H$	(278–308)		55.8 ± 0.3	293	GS	[2000VER2]
		$\Delta_v H$	(278–308)		55.5 ± 0.3	298	GS	[2000VER2]
		$\Delta_v H$	(344–445)		51.6	359	EB	[1987AMB/GHI3]
		$\Delta_v H$	(288–428)		50.9	303	A	[1987STE/MAL]
		$\Delta_v H$	(428–562)		45.4	443	A	[1987STE/MAL]
		$\Delta_v H$	(228–243)		53.4 ± 3	398	TE	[1979DEK/OON]
		$\Delta_v H$ (monomer)			35.5 ± 0.1	298	C	[1970KON/WAD]
		$\Delta_v H$			53 ± 4	298	C	[1970KON/WAD]
C ₄ H ₈ O ₂	[625-55-8]	isopropyl formate						

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	Δ_vH	(221–342)	34.5	236	A	[1987STE/MAL, 1947STU]
C ₄ H ₈ O ₂	[922-67-8]	methyl propionate				
	Δ_vH	(313–363)	28.9	298	CGC	[1995CHI/HOS]
	Δ_vH		35.6 ± 0.4	298	GC	[1987AZA]
	Δ_vH	(231–353)	39.1	246	A	[1987STE/MAL]
	Δ_vH	(353–486)	32.8	368	A	[1987STE/MAL]
	Δ_vH		35.9 ± 0.1	298	C	[1980SVO/UCH]
	Δ_vH		34.9 ± 0.1	313	C	[1980SVO/UCH]
	Δ_vH		36.3 ± 0.3	298	GCC	[1980FUC/PEA]
	Δ_vH		35.8 ± 0.1	298	C	[1979SUN/SVE2]
	Δ_vH		34.2 ± 0.1	326	C	[1977SVO/VES]
	Δ_vH		33.8 ± 0.1	331	C	[1977SVO/VES]
	Δ_vH		32.8 ± 0.1	344	C	[1977SVO/VES]
	Δ_vH		32.1 ± 0.1	355	C	[1977SVO/VES]
Δ_vH		31.5 ± 0.1	363	C	[1977SVO/VES]	
Δ_vH	(293–353)	35.9	308	A	[1987STE/MAL, 1965MER/POL]	
C ₄ H ₈ O ₂	[110-74-7]	propyl formate				
	Δ_vH	(302–353)	35.3	317		[1993FAR/WIC]
	Δ_vH	(354–518)	32.7	369	A	[1987STE/MAL]
	Δ_vH	(230–355)	36.8	245	A	[1987STE/MAL]
	Δ_vH		37.5 ± 0.1	298	C	[1980SVO/UCH]
	Δ_vH		36.5 ± 0.1	313	C	[1980SVO/UCH]
	Δ_vH		35.8 ± 0.1	326	C	[1976CIH/HYN]
	Δ_vH		35.4 ± 0.1	331	C	[1976CIH/HYN]
	Δ_vH		34.4 ± 0.1	344	C	[1976CIH/HYN]
	Δ_vH		33.8 ± 0.1	351	C	[1976CIH/HYN]
	Δ_vH		33.5 ± 0.1	355	C	[1976CIH/HYN]
	Δ_vH		32.9 ± 0.1	363	C	[1976CIH/HYN]
Δ_vH	(299–355)	35.6	314		[1928NEL, 1984BOU/FRI]	
C ₄ H ₈ O ₂ S	[16215-14-8]	allyl methyl sulfone				
	Δ_vH	(405–450)	68.2	420	A	[1987STE/MAL, 1999DYK/SVO]
C ₄ H ₈ O ₂ S	[126-33-0]	tetrahydrothiophene-1,1-dioxide (sulfolane)				
	$\Delta_{\text{trs}}H$		7.86	288.6		
	$\Delta_{\text{fus}}H$		1.37	301.7		[1999AHL/LOH]
	Δ_vH		69.1 ± 1.4	298	C	[2004MOR/MAT]
	Δ_vH	(423–529)	59.0	438		[1999DYK/SVO]
	Δ_vH	(364–529)	53.7	379		[1999DYK/SVO]
	Δ_vH	(424–542)	67.8 ± 0.8	298	EB	[1997STE/CHI3]
	Δ_vH	(373–453)	58.2	413	TGA	[1987ALN/ALS]
	Δ_vH	(303–328)	31.0	315	A	[1987STE/MAL]
	Δ_vH	(413–558)	58.7	428	A	[1987STE/MAL]
Δ_vH	(360–400)	54.5	380		[1984SHC/KAP]	
C ₄ H ₈ O ₂ S	[126-33-0]	tetramethylene sulfone				
	$\Delta_{\text{trs}}H$		5.35	288.6		
	$\Delta_{\text{fus}}H$		1.43	301.6		[1996DOM/MOO]
C ₄ H ₈ O ₂ S ₂	[55337-75-2]	1,3-dithiane sulfone				
	$\Delta_{\text{fus}}H$		23.47	413.8	DSC	[2004ROU/TEM3]
	$\Delta_{\text{fus}}H$		22.0	414	DSC	[2003ROU/TEM2]
	$\Delta_{\text{sub}}H$	(342–358)	102.3 ± 0.9	350	ME	[2004ROU/TEM3]
	$\Delta_{\text{sub}}H$	(342–358)	103.6 ± 0.9	298	ME	[2004ROU/TEM3]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
C ₄ H ₈ O ₂ S ₂	[139408-38-1]	1,4-dithiane sulfone				
	$\Delta_{\text{fus}}H$		26.0	474.2	DSC	[2006TEM/ROU]
	$\Delta_{\text{sub}}H$	(340–354)	99.9 ± 1.2	298	ME	[2006ROU/TEM]
	$\Delta_{\text{sub}}H$	(340–354)	98.7 ± 1.2	347	ME	[2006ROU/TEM]
C ₄ H ₈ O ₃	[627-03-2]	ethoxyacetic acid				
	$\Delta_{\text{v}}H$	(280–310)	69.1	295	A	[1987STE/MAL]
C ₄ H ₈ O ₃	[19693-75-5]	2-methoxy-1,3-dioxolane				
	$\Delta_{\text{v}}H$	(278–308)	46.4 ± 0.8	298	GS	[2002VER]
	$\Delta_{\text{v}}H$	(278–308)	46.8 ± 0.8		GS	[1995RAK/VER2]
C ₄ H ₈ O ₃	[623-50-7]	ethyl glycolate				
	$\Delta_{\text{v}}H$	(287–432)	47.1	302	A	[1987STE/MAL, 1947STU]
C ₄ H ₈ O ₃	[594-61-6]	2-hydroxyisobutyric acid				
	$\Delta_{\text{v}}H$	(371–485)	67.5	386	A	[1987STE/MAL]
C ₄ H ₈ O ₃	[6149-41-3]	methyl 3-hydroxypropionate				
	$\Delta_{\text{v}}H$	(330–343)	60	336	A	[1987STE/MAL]
C ₄ H ₈ O ₃	[6290-49-9]	methoxyacetic acid, methyl ester				
	$\Delta_{\text{v}}H$	(285–310)	39.3	297	A	[1987STE/MAL]
C ₄ H ₈ O ₃	[547-64-8]	<i>(dl)</i> methyl lactate				
	$\Delta_{\text{v}}H$	(313–418)	44.7	328	A	[1987STE/MAL]
C ₄ H ₈ O ₃	[542-59-6]	ethylene glycol monoacetate				
	$\Delta_{\text{v}}H$	(301–346)	63.9 ± 0.3	298	GS	[2009VER/EME2]
	$\Delta_{\text{v}}H$	(363–448)	55.1	378	EB	[2007SCH/DOE]
C ₄ H ₈ O ₃	[13122-71-9]	peroxybutyric acid				
	$\Delta_{\text{v}}H$	(273–393)	45.5	288	A	[1987STE/MAL, 1971DYK]
C ₄ H ₈ O ₃	[623-53-0]	ethyl methyl carbonate				
	$\Delta_{\text{fus}}H$		11.24	219.4	DSC	[2004DIN]
C ₄ H ₈ O ₃ S	[109577-03-9]	1,3-oxathiane sulfone				
	$\Delta_{\text{fus}}H$		15.2	352.9	DSC	[2006TEM/ROU]
	$\Delta_{\text{sub}}H$	(307–324)	92.1 ± 0.7	298	ME	[2007ROU/TEM2]
	$\Delta_{\text{sub}}H$	(307–324)	91.7 ± 0.7	316	ME	[2007ROU/TEM2]
C ₄ H ₈ O ₃ S	[107-61-9]	1,4-oxathiane sulfone				
	$\Delta_{\text{trs}}H + \Delta_{\text{fus}}H$		20.2	403.3	DSC	[2006TEM/ROU]
	$\Delta_{\text{sub}}H$	(307–322)	92.0 ± 1.0	298	ME	[2007ROU/TEM2]
	$\Delta_{\text{sub}}H$	(307–322)	91.6 ± 1.0	314	ME	[2007ROU/TEM2]
C ₄ H ₈ O ₄	[293-30-1]	1,3,5,7-tetroxane				
	$\Delta_{\text{fus}}H$		22.59	385		[1996DOM/HEA]
	$\Delta_{\text{sub}}H$		79.6 ± 0.2	298	C	[1977PED/RYL, 1969MAN/MOR]
	$\Delta_{\text{sub}}H$		79.5		C	[1975BOG/BER]
C ₄ H ₈ S	[3772-13-2]	2,2-dimethylthiirane				
	$\Delta_{\text{v}}H$	(273–473)	37	288	A	[1987STE/MAL, 1971DYK, 1999DYK/SVO]
C ₄ H ₈ S	[3195-86-6]	2-ethylthiirane				
	$\Delta_{\text{v}}H$	(298–450)	39.7	313	A	[1987STE/MAL, 1971DYK]
C ₄ H ₈ S	[110-01-0]	tetrahydrothiophene				

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	Enthalpy							
		$\Delta_{\text{fus}}H$			7.35	177		[1985DEA]
		Δ_vH			38.8	298		[1971WIL/ZWO]
		Δ_vH	(331–401)		37.7	346	EB	[1952WHI/BER]
		Δ_vH	(343–434)		37.1	358	A, EB	[1987STE/MAL, 1952HUB/FIN, 1966OSB/DOU]
C₄H₈S₂	[505-23-7]		1,3-dithiane					
		$\Delta_{\text{trs}}H$			0.8	316.4		
		$\Delta_{\text{fus}}H$			14.4	327.2		[1996DOM/HEA]
		$\Delta_{\text{sub}}H$	(266–279)		62.9 ± 0.7	298	ME	[1999ROU/DAV]
		$\Delta_{\text{sub}}H$			69.9 ± 0.4	298	GC	[1989AZA]
		$\Delta_{\text{sub}}H$	(250–271)		72.6	263	TE,ME	[1983DEW/VAN]
		$\Delta_{\text{sub}}H$			52.3 ± 0.8	298	C	[1971MOR]
		Δ_vH			66.9 ± 0.4		GC	[1989AZA]
C₄H₈S₂	[505-29-3]		1,4-dithiane					
		$\Delta_{\text{fus}}H$			21.6	384.6		[1996DOM/HEA]
		$\Delta_{\text{sub}}H$			63	298		[1999DAV/FLO]
		$\Delta_{\text{sub}}H$			68.9	298		[1989AZA]
		$\Delta_{\text{sub}}H$	(253–276)		72.4	268	E	[1983DEW/VAN]
		Δ_vH	(389–437)		48.7	404		[1999DYK/SVO]
		Δ_vH			68.9 ± 0.5		GC	[1989AZA]
		Δ_vH	(388–437)		47.9	403	A	[1987STE/MAL]
C₄H₉Br	[109-65-9]		1-bromobutane					
		$\Delta_{\text{fus}}H$			9.23	160.4		[1996DOM/HEA]
		Δ_vH	(323–363)		36.4	298	CGC	[1995CHI/HOS]
		Δ_vH	(340–370)		36.4	298		[1991BAS/SVO]
		Δ_vH	(338–373)		34.6	353	A, EB	[1987STE/MAL, 1977SVO/MAJ]
		Δ_vH			35.6 ± 0.1	322	C	[1977SVO/MAJ]
		Δ_vH			34.9 ± 0.1	332	C	[1977SVO/MAJ]
		Δ_vH			34.5 ± 0.1	339	C	[1977SVO/MAJ]
		Δ_vH			33.7 ± 0.1	352	C	[1977SVO/MAJ]
		Δ_vH			33.0 ± 0.1	366	C	[1977SVO/MAJ]
		Δ_vH			36.6 ± 0.1	298	C	[1968WAD]
		Δ_vH			36.7 ± 0.1	298	C	[1966WAD]
		Δ_vH	(273–400)		37.5	288	A, EST	[1987STE/MAL, 1961LI/ROS, 1971DYK]
		Δ_vH	(293–343)		33.5	308		[1929SMY/ENG, 1984BOU/FRI]
C₄H₉Br	[78-76-2]		2-bromobutane					
		$\Delta_{\text{fus}}H$			6.88	160.3		[1991ACR]
		Δ_vH	(281–403)		33.9	296	A	[1987STE/MAL, 1971DYK]
		Δ_vH			34.5 ± 0.1	298	C	[1968WAD]
		Δ_vH			34.8 ± 0.1	298	C	[1966WAD]
C₄H₉Br	[78-77-3]		1-bromo-2-methylpropane					
		Δ_vH	(305–363)		34.1	320	A, EB	[1987STE/MAL, 1977SVO/MAJ]
		Δ_vH			33.1 ± 0.1	330	C	[1977SVO/MAJ]
		Δ_vH			32.6 ± 0.1	341	C	[1977SVO/MAJ]
		Δ_vH			32.0 ± 0.1	353	C	[1977SVO/MAJ]
		Δ_vH			31.4 ± 0.1	366	C	[1977SVO/MAJ]
		Δ_vH	(281–404)		34	296	A	[1987STE/MAL, 1971DYK]
		Δ_vH			34.9 ± 0.1	298	C	[1968WAD]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₄ H ₉ Br	[507-19-7]	2-bromo-2-methylpropane				
	$\Delta_{\text{trs}}H$		5.65	208.6		
	$\Delta_{\text{trs}}H$		1.05	231.5		
	$\Delta_{\text{fus}}H$		1.97	256.1		[1996DOM/HEA]
	$\Delta_{\text{trs}}H$		5.85	209.3		
	$\Delta_{\text{trs}}H$		0.96	231.8		
	$\Delta_{\text{fus}}H$		NA	NA	DSC	[1986WEN/SCH]
	Δ_vH	(248–346)	31.4	263	A	[1987STE/MAL]
	Δ_vH	(270–345)	31	298		[1984BOU/FRI, 1991BAS/SVO]
	Δ_vH	(298–323)	31.5	313		[1969CAL/VAL]
	Δ_vH		31.8 ± 0.1	298	C	[1968WAD]
Δ_vH	(273–346)	31.2	288		[1951BRY/HOW, 1984BOU/FRI]	
C ₄ H ₉ BrO	[2482-57-7]	1-bromo-2-butanol				
	Δ_vH	(296–418)	58.4	311	A	[1987STE/MAL, 1947STU]
C ₄ H ₉ Cl	[109-69-3]	1-chlorobutane				
	Δ_vH	(260–350)	33.5	298		[1984BOU/FRI, 1991BAS/SVO]
	Δ_vH		33.5 ± 0.1	298	C	[1981TEK/MAJ]
	Δ_vH		32.7 ± 0.1	313	C	[1981TEK/MAJ]
	Δ_vH		31.8 ± 0.1	328	C	[1981TEK/MAJ]
	Δ_vH		30.9 ± 0.1	343	C	[1981TEK/MAJ]
	Δ_vH		30.0 ± 0.1	358	C	[1981TEK/MAJ]
	Δ_vH		29.4 ± 0.1	358	C	[1981TEK/MAJ]
	Δ_vH	(256–352)	35.6	271	DTA	[1969KEM/KRE]
	Δ_vH		33.5 ± 0.1	298	C	[1968WAD]
	Δ_vH	(257–389)	35	272	A, EST	[1987STE/MAL, 1961LI/ROS, 1971DYK]
Δ_vH	(293–343)	37.2	308		[1929SMY/ENG, 1984BOU/FRI]	
C ₄ H ₉ Cl	[78-86-4]	2-chlorobutane				
	Δ_vH	(315–341)	30.9	328	EB	[1996DAH/WIC]
	Δ_vH	(266–377)	33.1	281	A	[1987STE/MAL]
	Δ_vH		31.5 ± 0.1	298	C	[1981TEK/MAJ]
	Δ_vH		30.7 ± 0.1	313	C	[1981TEK/MAJ]
	Δ_vH		29.9 ± 0.1	328	C	[1981TEK/MAJ]
	Δ_vH		29.1 ± 0.1	343	C	[1981TEK/MAJ]
	Δ_vH		28.2 ± 0.1	358	C	[1981TEK/MAJ]
	Δ_vH		31.6 ± 0.1	298	C	[1968WAD]
	Δ_vH	(273–312)	31.8	288		[1928ROL, 1984BOU/FRI]
C ₄ H ₉ Cl	[513-36-0]	1-chloro-2-methylpropane				
	Δ_vH	(219–342)	36.1	234	A	[1987STE/MAL, 1947STU]
	Δ_vH		31.7 ± 0.1	298	C	[1968WAD]
C ₄ H ₉ Cl	[507-20-0]	2-chloro-2-methylpropane				
	$\Delta_{\text{trs}}H$		2.08	183.6		
	$\Delta_{\text{trs}}H$		5.75	219.8		
	$\Delta_{\text{fus}}H$		1.89	247.8	DSC	[2000TAM/LOP]
	$\Delta_{\text{trs}}H$		1.87	182.9		
	$\Delta_{\text{trs}}H$		5.88	219.3		
	$\Delta_{\text{fus}}H$		1.97	248.1		[1985DEA]
	$\Delta_{\text{trs}}H$		1.86	183.1		
	$\Delta_{\text{trs}}H$		5.66	219.4		
	$\Delta_{\text{fus}}H$		1.99	248.4		[1972URB/JAN]
	Δ_vH	(313–353)	28.6	298	CGC	[1995CHI/HOS]
Δ_vH	(253–358)	32.3	268	A	[1987STE/MAL, 1971DYK]	

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(295–323)	27.8	309	A	[1987STE/MAL, 1969CAL/VAL]
	$\Delta_v H$	(295–323)	27	310		[1969CAL/VAL, 1984BOU/FRI]
	$\Delta_v H$		29.0 ± 0.1	298	C	[1968WAD]
	$\Delta_v H$	(254–324)	29.1	269		[1947STU]
C₄H₉ClO₂	[628-89-7]	2-(2-chloroethoxy)ethanol				
	$\Delta_v H$	(326–469)	59.8	341	A	[1987STE/MAL, 1947STU]
C₄H₉ClO₂S	[2386-60-9]	butyl sulfonyl chloride				
	$\Delta_v H$	(283–373)	55.7	298		[1999DYK/SVO]
	$\Delta_v H$	(373–474)	52.9	388		[1999DYK/SVO]
	$\Delta_v H$	(253–283)	60.2	268	A	[1987STE/MAL, 1999DYK/SVO]
C₄H₉ClS	[693-07-2]	ethyl (2-chloroethyl) sulfide				
	$\Delta_v H$	(293–333)	44.4	308	A, GS	[1987STE/MAL, 1948RED/CHA, 1971DYK]
C₄H₉F	[2366-52-1]	1-fluorobutane				
	$\Delta_v H$	(222–326)	30.1	237	A, EST	[1987STE/MAL, 1961LI/ROS, 1971DYK]
C₄H₉F	[359-01-3]	2-fluorobutane				
	$\Delta_v H$	(233–329)	29.2	248	A	[1987STE/MAL, 1971DYK]
C₄H₉F	[353-61-7]	2-fluoro-2-methylpropane				
	$\Delta_v H$	(222–315)	27.6	237	A	[1987STE/MAL, 1971DYK]
C₄H₉FO	[372-93-0]	4-fluoro-1-butanol				
	$\Delta_v H$	(323–343)	64	333	A	[1987STE/MAL]
C₄H₉I	[542-69-8]	1-iodobutane				
	$\Delta_v H$	(313–353)	40.3	298	CGC	[1995CHI/HOS]
	$\Delta_v H$	(313–353)	39.7	298	CGC	[1995CHI/HOS]
	$\Delta_v H$		40.6 ± 0.1	298	C	[1968WAD]
	$\Delta_v H$	(292–431)	39.9	307	A, EST	[1987STE/MAL, 1961LI/ROS, 1971DYK]
C₄H₉I	[513-48-4]	2-iodobutane				
	$\Delta_v H$	(313–353)	37.9	298	CGC	[1995CHI/HOS]
	$\Delta_v H$	(313–353)	38.8	298	CGC	[1995CHI/HOS]
	$\Delta_v H$		38.5 ± 0.1	298	C	[1968WAD]
C₄H₉I	[513-38-2]	1-iodo-2-methylpropane				
	$\Delta_v H$	(256–393)	41.1	271	A	[1987STE/MAL, 1947STU]
	$\Delta_v H$		38.8 ± 0.1	298	C	[1968WAD]
C₄H₉I	[558-17-8]	2-iodo-2-methylpropane				
	$\Delta_{\text{sub}} H$	(202–223)	49.8	212.5	MG	[1987STE/MAL, 1944MIL2]
	$\Delta_v H$	(313–353)	37.0	298	CGC	[1995CHI/HOS]
	$\Delta_v H$	(236–294)	34.8	279	A	[1987STE/MAL, 1971DYK]
	$\Delta_v H$		35.4 ± 0.1	298	C	[1968WAD]
C₄H₉N	[123-75-1]	pyrrolidine				
	$\Delta_{\text{fus}} H$		0.54	207.1		
	$\Delta_{\text{fus}} H$		8.58	215.3		[1996DOM/HEA]
	$\Delta_v H$	(273–313)	38.4	288	A	[1987STE/MAL]
	$\Delta_v H$	(316–394)	35.8	331	EB, IP	[1987STE/MAL, 1959MCC/DOU, 1968OSB/DOU]
	$\Delta_v H$		35.8 ± 0.1	322	C	[1959MCC/DOU]
	$\Delta_v H$		34.5 ± 0.1	340	C	[1959MCC/DOU]
	$\Delta_v H$		33.0 ± 0.1	360	C	[1959MCC/DOU]
	$\Delta_v H$	(294–360)	37.3	309		[1959HIL/SIN, 1984BOU/FRI]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₄ H ₉ NO	[96-29-7]	2-butanone oxime				
	$\Delta_v H$	(283–329)	58.6 ± 0.2	306	GS	[2009VER/EME4]
	$\Delta_v H$	(283–329)	59.1 ± 0.2	298	GS	[2009VER/EME4]
	$\Delta_v H$	(308–425)	53.7	323	A	[1987STE/MAL]
	$\Delta_v H$	(318–343)	55.5	330	A	[1987STE/MAL]
	$\Delta_v H$	(313–333)	57.2	323		[1975MES/BAE, 2009VER/EME4]
$\Delta_v H$	(313–333)	57.7	298		[1975MES/BAE, 2009VER/EME4]	
C ₄ H ₉ NO	[625-50-3]	N-ethylacetamide				
	$\Delta_v H$	(361–423)	55.7	376		[1995SCH/PUS]
			64.9 ± 0.2	298	C	[1984STA/WAD]
C ₄ H ₉ NO	[110-69-0]	butyraldehyde oxime				
	$\Delta_v H$	(313–343)	55.8	328	A	[1987STE/MAL]
C ₄ H ₉ NO	[541-35-5]	butyramide				
	$\Delta_{\text{fus}} H$		19.2	387.3		[2008ABA/BAD]
	$\Delta_{\text{fus}} H$		19.2	387.3		[2000BRU/DEL]
	$\Delta_{\text{sub}} H$	(288–354)	82 ± 4.0	298	TE	[2000BRU/DEL]
	$\Delta_{\text{sub}} H$	(298–347)	82 ± 4.0	298	TE	[2000BRU/DEL]
	$\Delta_{\text{sub}} H$		86.4 ± 0.4			[1975BAR/PIL, 1977PED/RYL]
	$\Delta_{\text{sub}} H$	(292–304)	85.4 ± 1.7	298	ME	[1973LEB/KAT2, 1977PED/RYL]
	$\Delta_{\text{sub}} H$	(353–373)	87	363		[1960JON]
	$\Delta_{\text{sub}} H$	(336–382)	86.4 ± 0.4	359	GS	[1959DAV/JON2]
	$\Delta_{\text{sub}} H$	(298–341)	87.0 ± 0.8	320	ME	[1959DAV/JON2]
			79.9		[1960THO]	
$\Delta_v H$	(397–504)	64	412	A	[1987STE/MAL]	
C ₄ H ₉ NO	[563-83-7]	2-methylpropanamide				
	$\Delta_{\text{fus}} H$		19.2	400.1		[2008ABA/BAD]
	$\Delta_{\text{sub}} H$		82			[2000BRU/DEL]
	$\Delta_{\text{sub}} H$	(285–302)	86.1 ± 0.2	294	ME	[1989ABB/JIM]
			86.0 ± 0.2	298		[1989ABB/JIM]
C ₄ H ₉ NO	[127-19-5]	N,N-dimethylacetamide				
	$\Delta_{\text{fus}} H$		10.2	254.2		[2007SMI/TSV]
	$\Delta_{\text{fus}} H$		8.2	253.2		[2000LIS/JAM]
	$\Delta_{\text{fus}} H$		10.42	251.4		[1999AHL/LOH]
	$\Delta_v H$	(463–513)	50.7 ± 0.7	298	CGC	[2009PAN/ANT]
	$\Delta_v H$	(298–423)	45.8	298		[2005NAS/NEU]
	$\Delta_v H$	(371–423)	45.1	386	A	[1987STE/MAL]
	$\Delta_v H$		50.2	298	A	[1985BAR/CAS, 1985MAJ/SVO]
	$\Delta_v H$	(297–438)	67.9	312		[1974MYA/SCH, 1984BOU/FRI]
	$\Delta_v H$		43.7	298	I	[1971SUN/EIS]
$\Delta_v H$	(303–363)	45.2	318	A	[1987STE/MAL, 1968GOP/RIZ]	
C ₄ H ₉ NO	[1187-58-2]	N-methylpropionamide				
	$\Delta_v H$	(307–371)	64.0 ± 0.2	339	GS	[2009VER/EME4]
	$\Delta_v H$	(307–371)	66.6 ± 0.2	298	GS	[2009VER/EME4]
	$\Delta_v H$	(368–473)	66.9 ± 1.3	298	EB,BG	[1998MOR/KOP]
	$\Delta_v H$	(368–473)	64.0 ± 0.3	420	EB,BG	[1998MOR/KOP]
	$\Delta_v H$	(361–414)	54.2	376		[1995SCH/PUS]
	$\Delta_v H$		64.9 ± 0.3	298	C	[1984STA/WAD]
	$\Delta_v H$	(303–363)	54.4	318	A	[1987STE/MAL, 1968GOP/RIZ]
$\Delta_v H$	(381–480)	56.6 ± 0.2	431	EB	[1983VAS, 2009VER/EME4]	

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	Δ_vH	(381–480)	63.9 ± 0.2	431	EB	[1983VAS, 2009VER/EME4]
C ₄ H ₉ NO	[110-91-8]	morpholine				
	Δ_vH		45.3 ± 0.5	298	DSC	[2005ROJ/GIN]
	Δ_vH	(274–303)	45.6 ± 0.4	288	GS	[1998VER2]
	Δ_vH	(274–303)	45.0 ± 0.4	298	GS	[1998VER2]
	Δ_vH	(346–401)	40.8	361		[1991WU/LOC]
	Δ_vH	(313–343)	44.3	328	TGA	[1987ALN/ALS]
	Δ_vH	(273–318)	45.3	288	A	[1987STE/MAL, 1975CAB/CON]
	Δ_vH	(317–443)	42.3	332	A	[1987STE/MAL]
C ₄ H ₉ NO ₂	[2835-81-6]	<i>(dl)</i> 2-aminobutyric acid				
	$\Delta_{\text{sub}}H$		132 ± 2	409	TE,ME	[1979DEK/VOO]
	$\Delta_{\text{sub}}H$	(400–418)	132	409	A	[1987STE/MAL]
C ₄ H ₉ NO ₂	[1492-24-6]	S 2-aminobutyric acid				
	$\Delta_{\text{sub}}H$		162.8 ± 0.8	455	ME	[1965SVE/CLY, 1964CLY/SVE]
	$\Delta_{\text{sub}}H$	(449–462)	162.5	455	A	[1987STE/MAL]
C ₄ H ₉ NO ₂	[62-57-7]	2-aminoisobutyric acid				
	$\Delta_{\text{sub}}H$	(439–462)	125.8	450.5	A	[1987STE/MAL]
	$\Delta_{\text{sub}}H$	(403–424)	134.2	413.5	A	[1987STE/MAL]
	$\Delta_{\text{sub}}H$		129.5 ± 0.4	455	ME	[1965SVE/CLY, 1964CLY/SVE]
C ₄ H ₉ NO ₂	[56-12-2]	4-aminobutanoic acid				
	$\Delta_{\text{sub}}H$	(460–475)	139 ± 4			[2009LEG/BAC]
	$\Delta_{\text{sub}}H$	(384–407)	138.9 ± 0.6	395	C	[1983SKO/SAB]
	$\Delta_{\text{sub}}H$		140 ± 2	395	C	[1983SKO/SAB]
	Δ_vH	(493–500)	87 ± 2			[2009LEG/BAC]
C ₄ H ₉ NO ₂	[924-43-6]	sec-butyl nitrite				
	Δ_vH	(267–287)	29.6	277	A	[1987STE/MAL, 1937THO/DAI]
C ₄ H ₉ NO ₂	[540-80-7]	<i>tert</i> -butyl nitrite				
	Δ_vH	(267–337)	30.8	282	A	[1987STE/MAL, 1937THO/DAI]
C ₄ H ₉ NO ₂	[na]	lactic acid N-methyl amide				
	Δ_vH	(359–415)	72.7	374	A	[1987STE/MAL]
C ₄ H ₉ NO ₂	[105-40-8]	N-methyl carbamic acid, ethyl ester				
	Δ_vH	(299–443)	51.7	314	A	[1987STE/MAL, 1947STU]
C ₄ H ₉ NO ₂	[625-74-1]	2-methyl-1-nitropropane				
	Δ_vH	(347–415)	41.1	362	A, EB	[1987STE/MAL, 1956TOO, 1971DYK]
C ₄ H ₉ NO ₂	[594-70-7]	2-methyl-2-nitropropane				
	$\Delta_{\text{trs}}H$		4.2	215.3		
	$\Delta_{\text{trs}}H$		4.7	260.1		
	$\Delta_{\text{fus}}H$		2.6	299.2		[1997REU/BUS]
	Δ_vH	(334–401)	39.1	349	EB	[1987STE/MAL, 1956TOO, 1971DYK]
C ₄ H ₉ NO ₂	[627-05-4]	1-nitrobutane				
	Δ_vH	(313–353)	47.0	298	CGC	[1995CHI/HOS]
	Δ_vH	(357–426)	42.7	372	A, EB	[1987STE/MAL, 1956TOO, 1971DYK]
C ₄ H ₉ NO ₂	[600-24-8]	<i>(dl)</i> 2-nitrobutane				
	Δ_vH	(345–413)	40.3	360	A, EB	[1987STE/MAL, 1956TOO, 1971DYK]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound					
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference	
C ₄ H ₉ NO ₂	[627-12-3] $\Delta_{\text{v}}H$	propyl carbamate (325–468)	61.6	340	A	[1987STE/MAL, 1947STU]	
C ₄ H ₉ NO ₃	[928-45-0] $\Delta_{\text{v}}H$	butyl nitrate (273–343)	44.1	288	A	[1987STE/MAL, 1971DYK, 1957GRA/PRA]	
C ₄ H ₉ NO ₃	[543-29-3] $\Delta_{\text{v}}H$	isobutyl nitrate (273–343)	42.8	288	A	[1987STE/MAL, 1971DYK, 1957GRA/PRA]	
C ₄ H ₉ NO ₃	[80-68-2] $\Delta_{\text{sub}}H$	(<i>dl</i>)-threonine (341–441)	U 96 ± 8	391	LE	[1977GAF/PIE]	
C ₄ H ₉ NO ₃	[76-39-1] $\Delta_{\text{trs}}H$ $\Delta_{\text{fus}}H$ $\Delta_{\text{trs}}H$ $\Delta_{\text{fus}}H$ $\Delta_{\text{sub}}H$ (<i>cryst</i>) $\Delta_{\text{sub}}H$ (<i>plastic</i>) $\Delta_{\text{sub}}H$ (<i>plastic</i>) $\Delta_{\text{sub}}H$ (<i>cryst</i>)	2-methyl-2-nitro-1-propanol	14.64	311.5			
			3.17	363.9		[1999SAL/LOP]	
			17.2	310			
			3.74	361		[1996DOM/HEA]	
			(293–309)	78 ± 1			[1995FON/MUN]
			(319–333)	64 ± 2			[1995FON/MUN]
				59.5 ± 3.0	319	C	[1994FON/MUN]
				73.2 ± 3.7	311	C	[1994FON/MUN]
C ₄ H ₉ NO ₄	[77-49-6] $\Delta_{\text{trs}}H$ $\Delta_{\text{fus}}H$ $\Delta_{\text{sub}}H$ (<i>cryst</i>) $\Delta_{\text{sub}}H$ (<i>plastic</i>) $\Delta_{\text{sub}}H$ (<i>plastic</i>) $\Delta_{\text{sub}}H$ (<i>cryst</i>)	2-methyl-2-nitro-1,3-propanediol	25.72	352			
			3.84	424		[1996DOM/HEA, 1994LOP/VAN]	
			(330–349)	98 ± 4			[1995FON/MUN]
			(361–382)	74 ± 5			[1995FON/MUN]
				79.3 ± 4.0	368	C	[1994FON/MUN]
				102.0 ± 5.1	339	C	[1994FON/MUN]
C ₄ H ₉ NO ₅	[126-11-4] $\Delta_{\text{sub}}H$ (<i>cryst</i>) $\Delta_{\text{sub}}H$ (<i>plastic</i>) $\Delta_{\text{sub}}H$ (<i>plastic</i>)	2-hydroxymethyl-2-nitro-1,3-propanediol	107 ± 10			[1995FON/MUN]	
			76 ± 8			[1995FON/MUN]	
			77.3 ± 3.9	368	C	[1994FON/MUN]	
C ₄ H ₉ N ₃ O ₂	[na] $\Delta_{\text{v}}H$	<i>bis</i> (nitrosoethyl)amine (291–450)	46.4	306	A	[1987STE/MAL]	
C ₄ H ₉ N ₃ O ₂	[216489-98-4] $\Delta_{\text{sub}}H$	1-[2-(ethenoxy)ethyl]-1-nitrosohydrazine	112.1 ± 1.9	298		[1998LEB/CHI]	
C ₄ H ₉ O ₂ PS ₂	[695-68-1] $\Delta_{\text{v}}H$	2-mercapto-4,5-dimethyl-1,3,2-dioxaphospholane-2-sulfide	66	298		[2008SAG/SAF]	
C ₄ H ₉ P	[62778-93-2] $\Delta_{\text{v}}H$	allymethylphosphine (242–291)	34.4	276	A	[1987STE/MAL]	
C ₄ H ₉ P	[114596-01-9] $\Delta_{\text{v}}H$	3-butenylphosphine (252–295)	34.5	273		[1988SHA/DIE]	
C ₄ H ₉ P	[3466-00-0] $\Delta_{\text{v}}H$	phospholane (257–347)	37.4	272	A	[1987STE/MAL]	
C ₄ H ₁₀	[106-97-8] $\Delta_{\text{trs}}H$ $\Delta_{\text{fus}}H$ $\Delta_{\text{sub}}H$ $\Delta_{\text{v}}H$	butane	2.05	107.6			
			4.66	134.9		[1996DOM/HEA]	
			35.9	107	B	[1966GEI/QUI]	
			(300–315)	22.9	308		[1997SAK/HOR]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
	$\Delta_{\text{v}}H$	(195–292)	23.4	277	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$	(273–321)	23.2	288	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$	(316–383)	22.6	331	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$	(375–425)	22.8	390	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$	(135–213)	27	198	A	[1987STE/MAL, 1973CAR/KOB]
	$\Delta_{\text{v}}H$		22.4	298		[1971WIL/ZWO]
	$\Delta_{\text{v}}H$	(206–279)	23.1	264		[1945WAC/LIN, 1984BOU/FRI]
	$\Delta_{\text{v}}H$	(195–273)	23.9	258		[1940AST/MES, 1984BOU/FRI]
C₄H₁₀	[75-28-5]	2-methylpropane (isobutane)				
	$\Delta_{\text{fus}}H$		4.49	113.7		[2009PER/MAG]
	$\Delta_{\text{fus}}H$		4.56	113.7		[1996DOM/HEA]
	$\Delta_{\text{v}}H$	(303–333)	21.5	318		[1999LIM/PAR]
	$\Delta_{\text{v}}H$	(186–280)	22.4	265	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$	(121–187)	26.9	172	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$	(263–306)	21.9	278	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$	(301–366)	21.4	316	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$	(361–408)	21.6	376	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$	(277–344)	21.6	292		[1976STE/POL, 1984BOU/FRI]
	$\Delta_{\text{v}}H$		21.3	286		[1971WIL/ZWO]
	$\Delta_{\text{v}}H$	(188–262)	22.6	247		[1940AST/KEN, 1984BOU/FRI]
C₄H₁₀F₃NOS	[26458-94-6]	(diethylamino)trifluorooxosulfur				
	$\Delta_{\text{v}}H$	(329–354)	49.5	341	A	[1987STE/MAL, 1999DYK/SVO]
C₄H₁₀F₃NS	[38078-09-0]	(N-ethylethaneaminato)trifluoro sulfur				
	$\Delta_{\text{v}}H$	(318–340)	45.2	329	A	[1987STE/MAL, 1999DYK/SVO]
C₄H₁₀N₂	[275-02-5]	piperazine				
	$\Delta_{\text{fus}}H$		26.7	384.6		[1997STE/CHI4]
	$\Delta_{\text{fus}}H$		22.1	381.8	DSC	[1997LEE/CHA]
	$\Delta_{\text{sub}}H$		72.1	298		[1998VER2]
	$\Delta_{\text{sub}}H$		65.2	385	B	[1997STE/CHI4]
	$\Delta_{\text{sub}}H$	(279–321)	73.1	294		[1987STE/MAL]
	$\Delta_{\text{v}}H$	(417–460)	50.1 ± 1.9	298	EB	[1997STE/CHI4]
C₄H₁₀N₂	[na]	trimethylammonium cyanide				
	$\Delta_{\text{sub}}H$	(219–236)	45.0	227.5		[1987STE/MAL]
C₄H₁₀N₂O	[927-67-3]	N-propylurea				
	$\Delta_{\text{trs}}H$		2.4	291.3		
	$\Delta_{\text{fus}}H$		11.9	370.2	DSC	[2005HAS/TAJ]
	$\Delta_{\text{trs}}H$		3.0	289.6		
	$\Delta_{\text{fus}}H$		14.92	383	DSC	[1995FER/DEL]
	$\Delta_{\text{fus}}H$		14.63	381		[1991ACR]
	$\Delta_{\text{sub}}H$	(333–357)	101.4 ± 0.6	298	GS	[2006EME/KAB]
	$\Delta_{\text{sub}}H$	(332–373)	90.7 ± 1.0	366		[1990PIA/FER, 1987FER/DEL2]
C₄H₁₀N₂O	[691-60-1]	N-isopropylurea				
	$\Delta_{\text{trs}}H$		1.41	280.8		
	$\Delta_{\text{trs}}H$		2.31	375.5		
	$\Delta_{\text{fus}}H$		17.5	429		[1990KAB/MIR2]
	$\Delta_{\text{sub}}H$	(333–372)	96.7 ± 1.6	353	ME	[2003ZAI/KAB]
	$\Delta_{\text{sub}}H$	(333–372)	96.8 ± 1.6	350	ME	[2003ZAI/KAB]
	$\Delta_{\text{sub}}H$		97.2 ± 0.6	350	C	[2003ZAI/KAB]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	Enthalpy							
		$\Delta_{\text{sub}}H$		100.6 ± 1.3		389		[1990PIA/FER]
C ₄ H ₁₀ N ₂ O	[632-14-4]		1,1,3-trimethylurea					
		$\Delta_{\text{fus}}H$		14.3		344.4		[1991ACR]
		Δ_vH	(345–375)	89.7 ± 1.2		360		[1990PIA/FER]
C ₄ H ₁₀ N ₂ O ₂	[7119-92-8]		diethylnitramine					
		Δ_vH	(338–378)	49.7		358		[1958CAS/FLE]
C ₄ H ₁₀ N ₄ O ₄	[4164-34-5]		N,N'-dimethyl-N,N'-dinitro-1,2-ethanediamine					
		$\Delta_{\text{fus}}H$		60.32		410		[1987OYU/BR1]
C ₄ H ₁₀ N ₆ O ₆	[13126-25-5]		2,4,6-trinitro-2,4,6-triazaheptane					
		$\Delta_{\text{fus}}H$		34.01		442.4	DSC	[1997ZEM]
C ₄ H ₁₀ O	[71-36-3]		1-butanol					
		$\Delta_{\text{fus}}H$		9.28		183.9		[1991ACR]
		Δ_vH	(298–363)	48.4		298		[2004NAS/ZIM]
		Δ_vH	(357–389)	46.0		372	EB	[2001MUN/KRA]
		Δ_vH		38.2		423		[2000WOR/FEN]
		Δ_vH		29.6		473		[2000WOR/FEN]
		Δ_vH		20.8		523		[2000WOR/FEN]
		Δ_vH		44.1				[1999FAT]
		Δ_vH	(323–373)	52.5		298	CGC	[1995CHI/HOS]
		Δ_vH	(315–390)	49.9		330		[1995DEJ/BUR]
		Δ_vH	(364–403)	45.3		379		[1993SUS/ORT2]
		Δ_vH	(283–323)	55.2		298		[1992GRA/SAN]
		Δ_vH	(376–399)	45.3		387	A	[1987STE/MAL]
		Δ_vH	(323–413)	50.1		338	A	[1987STE/MAL]
		Δ_vH	(413–550)	41.9		428	A	[1987STE/MAL]
		Δ_vH	(209–251)	51.6		236	A	[1987STE/MAL]
		Δ_vH	(376–397)	45.4		386	A	[1987STE/MAL]
		Δ_vH	(391–429)	43.8		406	A	[1987STE/MAL]
		Δ_vH	(415–501)	41.9		430	A	[1987STE/MAL]
		Δ_vH	(497–563)	37.4		512	A	[1987STE/MAL]
		Δ_vH	(243–303)	51.7		298		[1983SCH/STR]
		Δ_vH	(329–391)	49.0		344		[1982SAC/PES]
		Δ_vH		52.1		298	C	[1982FUC/PEA]
		Δ_vH	(288–404)	55.0		303		[1973WIL/ZWO]
		Δ_vH		49.5 ± 0.1		333	C	[1973SVO/VES]
		Δ_vH		48.6 ± 0.1		343	C	[1973SVO/VES]
		Δ_vH		47.5 ± 0.1		353	C	[1973SVO/VES]
		Δ_vH		46.4 ± 0.1		363	C	[1973SVO/VES]
		Δ_vH		52.34 ± 0.02		298	C	[1971POL/BEN]
		Δ_vH	(351–397)	47.2		366	EB	[1987STE/MAL, 1970AMB/SPR]
		Δ_vH	(295–391)	53.0		310	DTA	[1969KEM/KRE]
		Δ_vH		52.3 ± 0.1		298	C	[1966WAD]
		Δ_vH		47.2 ± 0.1		356	C	[1965COU/HAL]
	Δ_vH		45.4 ± 0.1		381	C	[1965COU/HAL]	
	Δ_vH		43.1 ± 0.1		391	C	[1965COU/HAL]	
	Δ_vH	(419–563)	42.1		434		[1963AMB/TOW]	
	Δ_vH	(362–398)	46.6		377	EB	[1963BID/COL]	
	Δ_vH		51.0 ± 0.1		298	C	[1963MCC/LAI]	
	Δ_vH	(337–390)	48.3		352		[1959BRO/SMI, 1984BOU/FRI]	
	Δ_vH	(314–390)	48.3		352		[1998KAH, 1984BOU/FRI]	

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₄ H ₁₀ O	[78-92-2]	2-butanol				
	$\Delta_{\text{fus}}H$		5.97	184.7		[1971AND/CON]
	Δ_vH	(315–371)	48.8	330	EB	[2009GIE/KOS]
	Δ_vH	(320–379)	46.2	335		[2009MAR/LLA]
	Δ_vH	(298–563)	46.2	298		[2004NAS/ZIM]
	Δ_vH	(306–373)	47.7	321		[1995DEJ/BUR]
	Δ_vH	(303–403)	49.3	318	A	[1987STE/MAL]
	Δ_vH	(359–381)	43.2	370	A	[1987STE/MAL]
	Δ_vH	(372–524)	47.9	387	A	[1987STE/MAL]
	Δ_vH	(210–303)	57.5	225	A	[1987STE/MAL]
	Δ_vH	(359–380)	43.2	369	A	[1987STE/MAL]
	Δ_vH	(368–404)	42	383	A	[1987STE/MAL]
	Δ_vH	(395–485)	39.6	410	A	[1987STE/MAL]
	Δ_vH	(476–536)	35	491	A	[1987STE/MAL]
	Δ_vH	(307–373)	47.8	322		[1982SAC/PES]
	Δ_vH	(293–380)	53.2	308		[1978CAV/CHA]
	Δ_vH	(319–372)	44.1	334		[1975BRA/AND]
	Δ_vH	(280–314)	50.2	295		[1975CAB/CON2]
	Δ_vH	(298–393)	48.1	313		[1973WIL/ZWO]
	Δ_vH		49.74 ± 0.02	298	C	[1971POL/BEN]
	Δ_vH	(323–373)	46.3	338		[1969BRO/FOC, 1984BOU/FRI]
	Δ_vH		49.7 ± 0.1	298	C	[1966WAD]
	Δ_vH	422–538)	38.4	437		[1963AMB/TOW]
	Δ_vH	(345–381)	44.1	360	EB	[1963BID/COL]
	Δ_vH		48.5	298	C	[1963MCC/LAI]
	Δ_vH	(340–379)	44.7	355	EB	[1987STE/MAL, 1962BER/MCK, 1970AMB/SPR]
	Δ_vH		45.3 ± 0.1	340	C	[1962BER/MCK]
Δ_vH		43.3 ± 0.1	355	C	[1962BER/MCK]	
Δ_vH		41.9 ± 0.1	365	C	[1962BER/MCK]	
Δ_vH		40.8 ± 0.1	372	C	[1962BER/MCK]	
C ₄ H ₁₀ O	[na]	(+) 2-butanol				
	$\Delta_{\text{fus}}H$		6.0	177.4		[1971AND/CON]
C ₄ H ₁₀ O	[78-83-1]	2-methyl-1-propanol				
	$\Delta_{\text{fus}}H$		6.32	171.2		[1968COU/LEE]
	Δ_vH	(350–400)	48.8	298		[1999ORT/HER]
	Δ_vH	(350–400)	45.4	365	EB	[1993SUS/ORT]
	Δ_vH	(313–411)	49.5	328	A	[1987STE/MAL]
	Δ_vH	(381–524)	46.0	396	A	[1987STE/MAL]
	Δ_vH	(202–243)	55.0	228	A	[1987STE/MAL]
	Δ_vH	(369–389)	44.2	379	A	[1987STE/MAL]
	Δ_vH	(383–416)	42.6	398	A	[1987STE/MAL]
	Δ_vH	(401–493)	41.1	416	A	[1987STE/MAL]
	Δ_vH	(483–548)	36.2	498	A	[1987STE/MAL]
	Δ_vH		50.8 ± 0.1	298	C	[1984MAJ/SVO]
	Δ_vH		49.7 ± 0.1	313	C	[1984MAJ/SVO]
	Δ_vH		48.3 ± 0.1	328	C	[1984MAJ/SVO]
	Δ_vH		45.0 ± 0.1	358	C	[1984MAJ/SVO]
	Δ_vH	(320–382)	48.1	335		[1982SAC/PES]
	Δ_vH	(293–388)	52.6	308		[1973WIL/ZWO]
	Δ_vH		50.79 ± 0.02	298	C	[1971POL/BEN]
	Δ_vH		46.2 ± 0.1	347	C	[1970COU/FEN]
	Δ_vH		44.2 ± 0.1	363	C	[1970COU/FEN]
	Δ_vH		41.9 ± 0.1	381	C	[1970COU/FEN]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	Δ_vH	(342–389)	46.2	357	A, EB	[1987STE/MAL, 1970AMB/SPR]
	Δ_vH	(333–381)	47.0	348		[1969BRO/FOC, 1984BOU/FRI]
	Δ_vH		50.8 ± 0.1	298	C	[1966WAD]
	Δ_vH	(423–548)	40.1	438		[1963AMB/TOW]
	Δ_vH	(353–388)	45.2	368	EB	[1963BID/COL]
	Δ_vH		49.8	298	C	[1963MCC/LAI]
C₄H₁₀O	[75-65-0]	2-methyl-2-propanol				
	$\Delta_{\text{trs}}H$		0.83	286.1		
	$\Delta_{\text{trs}}H$		0.49	294.5		
	$\Delta_{\text{fus}}H$		6.7	299		[1996DOM/HEA]
	$\Delta_{\text{sub}}H$	(253–298)	51.3	275	A	[1947STU]
	Δ_vH	(306–355)	47.4	298	EB	[2007MAL]
	Δ_vH	(323–368)	42.7	338		[2003ORT/ESP]
	Δ_vH	(321–359)	43.4	336		[1999AUC/LOR]
	Δ_vH	(323–373)	45.4	298	CGC	[1995CHI/HOS]
	Δ_vH	(299–375)	46.2	314	A	[1987STE/MAL]
	Δ_vH	(347–363)	41.4	355	A	[1987STE/MAL]
	Δ_vH	(356–480)	43.2	371	A	[1987STE/MAL]
	Δ_vH	(347–363)	41.4	355	A	[1987STE/MAL]
	Δ_vH	(357–461)	39.8	372	A	[1987STE/MAL]
	Δ_vH	(453–506)	33.6	468	A	[1987STE/MAL]
	Δ_vH		46.2 ± 0.1	303	C	[1984MAJ/SVO]
	Δ_vH		44.9 ± 0.1	313	C	[1984MAJ/SVO]
	Δ_vH		43.0 ± 0.1	328	C	[1984MAJ/SVO]
	Δ_vH		41.0 ± 0.1	343	C	[1984MAJ/SVO]
	Δ_vH		37.2 ± 0.1	368	C	[1984MAJ/SVO]
	Δ_vH	(306–357)	44.7	321		[1982SAC/PES]
	Δ_vH	(293–376)	46.5	308		[1973WIL/ZWO]
	Δ_vH		46.94 ± 0.02	298	C	[1971POL/BEN]
	Δ_vH	(313–355)	44.2	328		[1969BRO/FOC, 1984BOU/FRI]
	Δ_vH		46.6 ± 0.1	298	C	[1966WAD]
	Δ_vH	(333–363)	42.1	348	EB	[1963BEN/MCK]
	Δ_vH		42.5 ± 0.1	330	C	[1963BEN/MCK]
	Δ_vH		41.3 ± 0.1	340	C	[1963BEN/MCK]
	Δ_vH		40.4 ± 0.1	346	C	[1963BEN/MCK]
	Δ_vH		40.0 ± 0.1	349	C	[1963BEN/MCK]
	Δ_vH		39.0 ± 0.1	356	C	[1963BEN/MCK]
	Δ_vH	(329–363)	42.6	344	EB	[1987STE/MAL, 1970AMB/SPR, 1963BEN/MCK]
	Δ_vH		44.9	298	C	[1963MCC/LAI]
	Δ_vH	(373–506)	38.7	388		[1963AMB/TOW]
	Δ_vH	(293–363)	44.7	323		[1928PAR/BAR]
C₄H₁₀O	[60-29-7]	diethyl ether				
	$\Delta_{\text{fus}}H$		7.19	156.9		[1971COU/LEE]
	Δ_vH	(286–329)	28.1	301	A	[1987STE/MAL]
	Δ_vH	(307–457)	26.9	322	A	[1987STE/MAL]
	Δ_vH	(305–360)	27.5	320	A	[1987STE/MAL]
	Δ_vH	(351–420)	26.6	366	A	[1987STE/MAL]
	Δ_vH	(417–467)	26.7	432	A	[1987STE/MAL]
	Δ_vH		27.1 ± 0.1	298	C	[1980MAJ/WAN]
	Δ_vH	(250–329)	27.2	298		[1976AMB/ELL]
	Δ_vH	(250–329)	29.5	265	A	[1987STE/MAL, 1972AMB/SPR, 1976AMB/ELL]
	Δ_vH	(213–293)	28.4	278		[1922TAY/SMI]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₄ H ₁₀ O	[598-53-8]	isopropyl methyl ether				
	$\Delta_{\text{fus}}H$		5.85	127.3		[1996DOM/HEA]
	Δ_vH	(250–325)	28.8	265	A	[1987STE/MAL]
	Δ_vH		26.4 ± 0.1	298	C	[1980MAJ/WAN]
	Δ_vH	(260–325)	28.4	275	A	[1987STE/MAL, 1976AMB/ELL]
			26.4	298		[1976AMB/ELL]
C ₄ H ₁₀ O	[557-17-5]	methyl propyl ether				
	$\Delta_{\text{fus}}H$		7.67	134		[1996DOM/HEA]
	Δ_vH	(325–407)	27.2	340	A	[1987STE/MAL]
	Δ_vH	(401–476)	26.7	416	A	[1987STE/MAL]
	Δ_vH	(273–321)	30.7	288	A	[1987STE/MAL]
	Δ_vH		27.6 ± 0.1	298	C	[1980MAJ/WAN]
	Δ_vH	(253–328)	29.7	268	A	[1987STE/MAL, 1976AMB/ELL]
	Δ_vH		27.5	298		[1976AMB/ELL]
	Δ_vH		27.9 ± 0.2	298	C	[1975FEN/HAR]
	(273–312)	29.7	288		[1910BIN, 1984BOU/FRI]	
C ₄ H ₁₀ OS	[70-29-1]	diethyl sulfoxide				
	Δ_vH	(298–318)	58.7 ± 3.3	308		[2005MAR/ZAT]
C ₄ H ₁₀ O ₂	[26171-83-5]	(±) 1,2-butanediol				
	Δ_vH	(283–332)	70.4 ± 0.3	298	GS	[2004VER2]
	Δ_vH		73.3 ± 0.4	298	C	[2003EUS/LOP]
	Δ_vH	(372–506)	71.6 ± 0.8	298	EB	[1996STE/CHI]
	Δ_vH	(372–506)	51.46 ± 0.4	360	EB	[1996STE/CHI]
	Δ_vH	(372–506)	48.9 ± 0.4	400	EB	[1996STE/CHI]
	Δ_vH	(372–506)	46.3 ± 0.4	440	EB	[1996STE/CHI]
	Δ_vH	(372–506)	43.6 ± 0.4	480	EB	[1996STE/CHI]
			40.7 ± 0.5	520	EB	[1996STE/CHI]
C ₄ H ₁₀ O ₂	[107-88-0]	(±) 1,3-butanediol				
	Δ_vH	(288–332)	72.6 ± 0.3	298	GS	[2007VER]
	Δ_vH		72.8 ± 0.6	298	C	[2003EUS/LOP]
	Δ_vH	(365–518)	74.5 ± 1.0	298	EB	[1996STE/CHI]
	Δ_vH	(365–518)	72.3 ± 0.8	320	EB	[1996STE/CHI]
	Δ_vH	(365–518)	68.3 ± 0.7	360	EB	[1996STE/CHI]
	Δ_vH	(365–518)	64.1 ± 0.6	400	EB	[1996STE/CHI]
	Δ_vH	(365–518)	59.5 ± 0.5	440	EB	[1996STE/CHI]
	Δ_vH	(365–518)	54.4 ± 0.6	480	EB	[1996STE/CHI]
	Δ_vH	(362–483)	67.6	377	A	[1987STE/MAL]
	Δ_vH	(373–423)	59.7	398		[1935SCH/STA]
	(423–480)	58.1	451		[1935SCH/STA]	
C ₄ H ₁₀ O ₂	[na]	(R) 1,3-butanediol				
	Δ_vH		72.3 ± 0.7	298	C	[2003EUS/LOP]
C ₄ H ₁₀ O ₂	[110-63-4]	1,4-butanediol				
	$\Delta_{\text{fus}}H$		18.7	293.6		[1996DOM/HEA]
	Δ_vH	(330–363)	79.0 ± 0.9	298	GS	[2005VAS/VER]
	Δ_vH		78.3 ± 0.3	298	C	[2003EUS/LOP]
	Δ_vH		79.3 ± 0.5	298	C	[1988KNA/SAB, 1990KNA/SAB2]
	Δ_vH	(380–510)	72	395	A	[1987STE/MAL]
	Δ_vH	(416–501)	76.1 ± 0.5	298	EB	[1984PAL/CHO, 2005VAS/VER]
	(419–490)	76.6 ± 1.7	298		[1972GAR/HUS, 2003EUS/LOP]	

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₄ H ₁₀ O ₂	[513-85-9]	<i>(dl)</i> 2,3-butanediol				
	$\Delta_v H$	(348–457)	62.5	363	A	[1987STE/MAL]
	$\Delta_v H$	(317–455)	58.4	332		[1947STU]
	$\Delta_v H$	(353–403)	57.9	378		[1935SCH/STA]
	$\Delta_v H$	(303–456)	55.7	380		[1935SCH/STA]
C ₄ H ₁₀ O ₂	[na]	<i>meso</i> 2,3-butanediol				
	$\Delta_{\text{fus}} H$		10.8	306.6		[2003EUS/LOP]
	$\Delta_v H$		66.6 ± 0.4	298	C	[2003EUS/LOP]
	$\Delta_v H$	(413–453)	54.6	433		[1946KNO/SCH]
C ₄ H ₁₀ O ₂	[na]	<i>levo</i> 2,3-butanediol				
	$\Delta_v H$	(413–453)	52.6	433		[1946KNO/SCH]
C ₄ H ₁₀ O ₂	[na]	(S,S) 2,3-butanediol				
	$\Delta_v H$		63.2 ± 0.7	298	C	[2003EUS/LOP]
C ₄ H ₁₀ O ₂	[2163-42-0]	2-methyl-1,3-propanediol				
	$\Delta_v H$	(297–375)	73.6 ± 0.2	298	GS	[2007VER]
	$\Delta_v H$	(488–708)	71.3 ± 0.5	298		[2002WIL/VON, 2007VER]
C ₄ H ₁₀ O ₂	[628-37-5]	diethylperoxide				
	$\Delta_v H$	(253–333)	29.0	268	A	[1987STE/MAL, 1951EGE/EMT, 1971DYK]
C ₄ H ₁₀ O ₂	[534-15-6]	1,1-dimethoxyethane				
	$\Delta_v H$		36.4 ± 0.1	298	C	[1970KUS/WAD]
	$\Delta_v H$	(273–333)	33.4	288	A	[1987STE/MAL, 1949NIC/LAF, 1971DYK]
C ₄ H ₁₀ O ₂	[110-71-4]	1,2-dimethoxyethane				
	$\Delta_v H$	(304–358)	34.5		EB	[2009LI/FAN]
	$\Delta_v H$	(305–392)	36.8 ± 0.2	298	EB	[1996STE/CHI2]
	$\Delta_v H$	(238–298)	39.4	253	A	[1987STE/MAL]
	$\Delta_v H$	(238–363)	39.1	253	A	[1987STE/MAL]
	$\Delta_v H$	(225–366)	33.9	240		[1947STU]
C ₄ H ₁₀ O ₂	[110-80-5]	2-ethoxyethanol				
	$\Delta_v H$	(310–385)	47.4	325	EB	[2001CHY/FRA]
	$\Delta_v H$	(313–353)	49.4	298	EB	[1999ANT/FRA]
	$\Delta_v H$	(313–363)	50.0	298	EB	[1999ANT/FRA]
	$\Delta_v H$	(323–353)	45.9	338	TGA	[1987ALN/ALS]
	$\Delta_v H$		48.2 ± 0.1	298	C	[1971KUS/WAD]
	$\Delta_v H$		49.2	298	I	[1971SUN/EIS]
	$\Delta_v H$	(336–408)	44.7	351	A	[1987STE/MAL, 1956PIC/FRI]
C ₄ H ₁₀ O ₂	[107-98-2]	1-methoxy-2-propanol				
	$\Delta_v H$	(331–373)	46.2	298	EB	[2004ANT/GAL]
	$\Delta_v H$	(347–378)	46.4	298	EB	[2004CHY/FRA2]
C ₄ H ₁₀ O ₂ S	[111-48-8]	<i>bis</i> (2-hydroxyethyl) sulfide				
	$\Delta_v H$	(368–483)	27.1	383	A	[1987STE/MAL]
	$\Delta_v H$	(315–558)	28.3	330		[1947STU]
C ₄ H ₁₀ O ₂ S	[597-35-3]	diethyl sulfone				
	$\Delta_{\text{sub}} H$		86.2 ± 2.5			[UR/MAC, 1970COX/PIL]
C ₄ H ₁₀ O ₂ S ₂	[na]	<i>meso</i> 1,2- <i>bis</i> (methylsufinyl)ethane				
	$\Delta_{\text{fus}} H$		34.31	446.7		[2001CAL/MEL]
C ₄ H ₁₀ O ₂ S ₂	[na]	racemic 1,2- <i>bis</i> (methylsufinyl)ethane				
	$\Delta_{\text{fus}} H$		22.18	405.4		[2001CAL/MEL]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
		Note: The enthalpy and entropy of fusion values given in the paper are not consistent.				
C ₄ H ₁₀ O ₃	[111-46-6]	diethylene glycol				
	$\Delta_v H$	(410–539)	66.9 ± 0.3	420	EB	[2002STE/CHI2]
	$\Delta_v H$	(410–539)	63.1 ± 0.3	460	EB	[2002STE/CHI2]
	$\Delta_v H$	(410–539)	59.2 ± 0.3	500	EB	[2002STE/CHI2]
	$\Delta_v H$	(410–539)	55.1 ± 0.5	540	EB	[2002STE/CHI2]
	$\Delta_v H$	(373–453)	66.5	413	TGA	[1987ALN/ALS]
	$\Delta_v H$	(364–518)	59.8	379	A	[1987STE/MAL]
	$\Delta_v H$	(412–513)	66.8	427		[1981AMB/HAL, 1984BOU/FRI]
		(403–513)	69.2	418		[1927RIN, 1984BOU/FRI]
C ₄ H ₁₀ O ₃	[4435-50-1]	1,2,3-butanetriol				
	$\Delta_v H$	(375–537)	68.1	390		[1947STU]
C ₄ H ₁₀ O ₃	[149-73-5]	orthoformic acid trimethyl ester				
	$\Delta_v H$	(273–358)	39	288	A	[1987STE/MAL]
	$\Delta_v H$		38.1 ± 0.8	298		[1971PIH/TUO]
C ₄ H ₁₀ O ₃ S	[623-81-4]	diethyl sulfite				
	$\Delta_v H$		44.7			[1975DEM/KOV]
	$\Delta_v H$	(283–431)	44.5	298	A	[1987STE/MAL, 1947STU, 1999DYK/SVO]
C ₄ H ₁₀ O ₄	[149-32-6]	<i>meso</i> erythritol				
	$\Delta_{\text{fus}} H$		38.9	391.2		[2005LOP/TOM]
	$\Delta_{\text{fus}} H$		40.3	392.2		[2002JON/COO]
	$\Delta_{\text{sub}} H$		140	298	Vap+Fus	[2005LOP/TOM]
	$\Delta_{\text{sub}} H$		157	298	B	[1990BAR/DEL]
	$\Delta_{\text{sub}} H$		135.1 ± 2.2			[1950EDW, 1960JON, 1970COX/PIL]
	$\Delta_v H$		97 ± 1	398	C	[2005LOP/TOM]
	$\Delta_v H$	(397–428)	113.6 ± 1.1	412	TE	[1990BAR/DEL]
		(394–401)	93.3	397	A	[1987STE/MAL]
C ₄ H ₁₀ O ₄	[2319-57-5]	<i>(l)</i> -threitol				
	$\Delta_{\text{fus}} H$		29.1	361.8		[2005LOP/TOM]
	$\Delta_{\text{sub}} H$		123	298	Vap+Fus	[2005LOP/TOM]
	$\Delta_v H$		86 ± 1	398	C	[2005LOP/TOM]
C ₄ H ₁₀ O ₄ S	[64-67-5]	diethyl sulfate				
	$\Delta_v H$	(413–484)	50.1	428	A	[1987STE/MAL]
	$\Delta_v H$	(320–482)	54.9	335		[1947STU, 1999DYK/SVO]
C ₄ H ₁₀ S	[109-79-5]	1-butanethiol				
	$\Delta_{\text{fus}} H$		10.46	157.5		[1996DOM/HEA]
	$\Delta_v H$		36.5	298		[1971WIL/ZWO]
	$\Delta_v H$	(323–409)	35	338	A, EB	[1987STE/MAL, 1957SCO/FIN, 1966OSB/DOU]
	$\Delta_v H$		34.7 ± 0.1	330	C	[1957SCO/FIN]
	$\Delta_v H$		33.6 ± 0.1	350	C	[1957SCO/FIN]
			32.2 ± 0.1	371	C	[1957SCO/FIN]
C ₄ H ₁₀ S	[513-53-1]	<i>(dl)</i> 2-butanethiol				
	$\Delta_{\text{fus}} H$		6.48	133		[1996DOM/HEA]
	$\Delta_v H$		34.1	298		[1971WIL/ZWO]
	$\Delta_v H$	(310–395)	33.2	325	A, EB	[1987STE/MAL, 1958MCC/FIN, 1966OSB/DOU]
			32.9 ± 0.1	318	C	[1958MCC/FIN]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
		Δ_vH	32.3 ± 0.1	329	C	[1958MCC/FIN]
		Δ_vH	31.8 ± 0.1	337	C	[1958MCC/FIN]
		Δ_vH	30.6 ± 0.1	358	C	[1958MCC/FIN]
C₄H₁₀S	[513-44-0]	2-methyl-1-propanethiol				
		$\Delta_{\text{fus}}H$	4.98	128.3		[1996DOM/HEA]
		Δ_vH	34.6	298		[1971WIL/ZWO]
		Δ_vH	(314–399) 33.6	329	A, EB	[1987STE/MAL, 1958SCO/MCC, 1966OSB/DOU]
		Δ_vH	33.3 ± 0.1	321	C	[1958SCO/MCC]
		Δ_vH	32.3 ± 0.1	340	C	[1958SCO/MCC]
		Δ_vH	31.0 ± 0.1	361	C	[1958SCO/MCC]
C₄H₁₀S	[75-66-1]	<i>tert</i> -butyl mercaptan				
		$\Delta_{\text{trs}}H$	4.07	151.6		
		$\Delta_{\text{trs}}H$	0.65	157		
		$\Delta_{\text{trs}}H$	0.97	199.4		
		$\Delta_{\text{fus}}H$	2.48	274.4		[1996DOM/HEA]
		Δ_vH	(275–293) 30.1	284		[1998STO/NG]
		Δ_vH	30.8	298		[1971WIL/ZWO]
		Δ_vH	(293–373) 30.9	308	A, EB	[1987STE/MAL, 1953MCC/SCO, 1966OSB/DOU]
C₄H₁₀S	[352-93-2]	diethyl sulfide				
		$\Delta_{\text{fus}}H$	11.92	169.2		[1996DOM/HEA]
		Δ_vH	(293–361) 34.9	327		[2008BAE]
		Δ_vH	35.8 ± 0.7	298	C	[1989VOR/KLY]
		Δ_vH	35.5	298		[1981SHI/SAI]
		Δ_vH	35.8	298		[1971WIL/ZWO]
		Δ_vH	(318–396) 34.4	333	A, EB	[1987STE/MAL, 1952SCO/FIN, 1966OSB/DOU]
		Δ_vH	(309–371) 34.8	324	EB	[1952WHI/BER]
		Δ_vH	(233–361) 37.5	248		[1947STU]
		Δ_vH	33.5	364		[1935THO/LIN]
C₄H₁₀S	[1551-21-9]	methyl isopropyl sulfide				
		$\Delta_{\text{fus}}H$	9.36	171.7		[1996DOM/HEA]
		Δ_vH	34.1	298		[1971WIL/ZWO]
		Δ_vH	33.0 ± 0.1	318	C	[1955MCC/FIN]
		Δ_vH	32.0 ± 0.1	336	C	[1955MCC/FIN]
		Δ_vH	30.7 ± 0.1	358	C	[1955MCC/FIN]
		Δ_vH	(298–368) 33.8	313	A, EB	[1987STE/MAL, 1952WHI/BER]
C₄H₁₀S	[3877-15-4]	methyl propyl sulfide				
		$\Delta_{\text{fus}}H$	9.91	160.2		[1996DOM/HEA]
		Δ_vH	36.2	298		[1971WIL/ZWO]
		Δ_vH	(308–374) 35.3	323	A, EB	[1987STE/MAL, 1952WHI/BER]
		Δ_vH	34.5 ± 0.1	328	C	[1957SCO/FIN]
		Δ_vH	33.4 ± 0.1	347	C	[1957SCO/FIN]
		Δ_vH	32.1 ± 0.1	369	C	[1957SCO/FIN]
C₄H₁₀S₂	[1191-08-8]	1,4-butanedithiol				
		Δ_vH	(347–469) 50.9	362	A	[1987STE/MAL, 1999DYK/SVO]
		Δ_vH	55.3	298		[1962MAN/SUN]
		Δ_vH	54.9	298		[1962MAN/SUN]
C₄H₁₀S₂	[110-81-6]	diethyl disulfide				
		$\Delta_{\text{fus}}H$	9.4	171.6		[1996DOM/HEA]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(383–423)	44.8	298	CGC	[1995CHI/HOS]
	$\Delta_v H$		45.4 ± 0.8	298	C	[1989VOR/KLY]
	$\Delta_v H$	(287–434)	45.7	302	A	[1987STE/MAL]
	$\Delta_v H$		45.2 ± 0.1	298	C	[1985KUS]
	$\Delta_v H$		45.2	298		[1981SHI/SAI]
	$\Delta_v H$		45.6	298		[1971WIL/ZWO]
	$\Delta_v H$	(373–431)	40.9	388	EB	[1987STE/MAL, 1966OSB/DOU, 1952SCO/FIN2]
	$\Delta_v H$	(359–433)	41.5	374	EB	[1952WHI/BER]
C₄H₁₁N	[109-73-9]	butyl amine				
	$\Delta_v H$	(298–343)	35.2	313	I	[2000BEL/BEL]
	$\Delta_v H$	(283–373)	36.0	298		[1995WOL/LAN]
	$\Delta_v H$	(323–373)	35.6	298	CGC	[1995CHI/HOS]
	$\Delta_v H$	(313–350)	34.7	328	A	[1987STE/MAL]
	$\Delta_v H$		35.7 ± 0.2	298	C	[1985KUS]
	$\Delta_v H$	(296–349)	35.5	311	EB	[1979MAJ/SVO2]
	$\Delta_v H$		35.7 ± 0.1	298	C	[1979MAJ/SVO2]
	$\Delta_v H$		34.7 ± 0.1	313	C	[1979MAJ/SVO2]
	$\Delta_v H$		33.5 ± 0.1	323	C	[1979MAJ/SVO2]
	$\Delta_v H$		32.4 ± 0.1	343	C	[1979MAJ/SVO2]
	$\Delta_v H$		31.1 ± 0.1	358	C	[1979MAJ/SVO2]
	$\Delta_v H$		35.7 ± 0.1	298	C	[1969WAD]
C₄H₁₁N	[13952-84-6]	(<i>dl</i>) 2-aminobutane				
	$\Delta_v H$	(264–371)	34.1	279	A	[1987STE/MAL, 1971DYK]
C₄H₁₁N	[13952-84-6]	sec-butylamine				
	$\Delta_v H$	(300–335)	32.4	315	EB	[1979MAJ/SVO2]
	$\Delta_v H$		32.7 ± 0.1	298	C	[1979MAJ/SVO2]
	$\Delta_v H$		31.6 ± 0.1	313	C	[1979MAJ/SVO2]
	$\Delta_v H$		30.5 ± 0.1	328	C	[1979MAJ/SVO2]
	$\Delta_v H$		29.4 ± 0.1	343	C	[1979MAJ/SVO2]
	$\Delta_v H$		32.6 ± 0.1	298	C	[1969WAD]
C₄H₁₁N	[78-81-9]	isobutylamine				
	$\Delta_v H$	(248–347)	37.6	263	A	[1987STE/MAL]
	$\Delta_v H$		33.9 ± 0.1	298	C	[1979MAJ/SVO2]
	$\Delta_v H$		32.7 ± 0.1	313	C	[1979MAJ/SVO2]
	$\Delta_v H$		31.6 ± 0.1	328	C	[1979MAJ/SVO2]
	$\Delta_v H$	(297–340)	33.5	313	EB	[1979MAJ/SVO2]
	$\Delta_v H$		33.8 ± 0.1	298	C	[1969WAD]
	$\Delta_v H$		33.9 ± 0.2	298	IP	[1965DOU/OSB, 1970GOO/MOR]
C₄H₁₁N	[75-64-9]	<i>tert</i> -butylamine				
	$\Delta_{\text{us}}H$		0.11	91.3		
	$\Delta_{\text{us}}H$		6.05	202.3		
	$\Delta_{\text{fus}}H$		0.88	206.2		[1996DOM/HEA]
	$\Delta_v H$	(283–343)	30.5	298		[1995WOL/LAN]
	$\Delta_v H$		29.6 ± 0.1	298	C	[1969WAD]
	$\Delta_v H$	(292–349)	30.1	307	A,EB,IP	[1987STE/MAL, 1968OSB/DOU]
C₄H₁₁N	[109-89-7]	diethylamine				
	$\Delta_v H$	(302–328)	31.2	315	A	[1987STE/MAL]
	$\Delta_v H$	(325–437)	30.4	340	A	[1987STE/MAL]
	$\Delta_v H$	(431–496)	28.4	446	A	[1987STE/MAL]
	$\Delta_v H$		31.3 ± 0.1	298	C	[1979MAJ/SVO2]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$		30.2 ± 0.1	313	C	[1979MAJ/SVO2]
	$\Delta_v H$		29.1 ± 0.1	328	C	[1979MAJ/SVO2]
	$\Delta_v H$		28.0 ± 0.1	343	C	[1979MAJ/SVO2]
	$\Delta_v H$		31.2 ± 0.1	298	C	[1969WAD]
	$\Delta_v H$	(273–333)	32.7 ± 0.2	298	I	[1969FRA/WAT]
	$\Delta_v H$	(292–313)	31.8	307		[1965KIL/BIT, 1984BOU/FRI]
	$\Delta_v H$	(304–323)	31.5	319		[1962BIT/KAU, 1984BOU/FRI]
C₄H₁₁N	[4747-21-1]	N-methyl isopropyl amine				
	$\Delta_v H$		30.7 ± 0.1	298	C	[1979PET/MAJ]
	$\Delta_v H$		29.5 ± 0.1	313	C	[1979PET/MAJ]
	$\Delta_v H$		27.1 ± 0.1	343	C	[1979PET/MAJ]
	$\Delta_v H$	(293–319)	30.9	306	EB	[1979PET/MAJ]
C₄H₁₁NO	[108-01-0]	2-(dimethylamino)ethanol				
	$\Delta_v H$	(278–316)	46.5 ± 0.4	298	GS	[2005KAP/SLO]
	$\Delta_v H$	(350–387)	43.2	365	A	[1987STE/MAL]
	$\Delta_v H$	(323–408)	42.7	338	A	[1987STE/MAL]
	$\Delta_v H$	(298–308)	47.9	298		[1982TOU/OKA, 2005KAP/SLO]
	$\Delta_v H$	(333–423)	47.6	298	EB	[1970QUI/HOF, 2005KAP/SLO]
C₄H₁₁NO	[5332-73-0]	3-methoxypropylamine				
	$\Delta_v H$	(278–390)	44.5	293	A	[1987STE/MAL]
C₄H₁₁NO₂	[111-42-2]	2,2'-iminodiethanol				
	$\Delta_v H$	(463–582)	69	478		[1959MCD/SHR]
C₄H₁₁NO₂	[111-42-2]	diethanolamine				
	$\Delta_{\text{sub}}H$		105.9 ± 2.0	298	C	[1982MIN/SAB]
	$\Delta_v H$	(423–542)	74.4	438	A	[1987STE/MAL]
	$\Delta_v H$	(376–454)	77.0	391		[1969DAN/MAT, 1984BOU/FRI]
	$\Delta_v H$	(466–514)	70.6	481		[1959MCD/SHR, 1984BOU/FRI]
C₄H₁₁NO₂	[115-69-5]	2-amino-2-methyl-1,3-propanediol				
	$\Delta_{\text{trs}}H$		23.55	356.7		
	$\Delta_{\text{fus}}H$		2.76	384.1	DSC	[2006DIV/CHE]
	$\Delta_{\text{trs}}H$		25.21	352		
	$\Delta_{\text{fus}}H$		2.99	384		[1996DOM/HEA, 1994LOP/VAN]
	$\Delta_{\text{trs}}H$	(283–393)	5.0	352.9		
	$\Delta_{\text{trs}}H$	(283–393)	18.46	353.7		
	$\Delta_{\text{fus}}H$	(283–393)	2.78	384.1	AC	[1990ZHA/YAN]
	$\Delta_{\text{sub}}H$ (cryst)	(330–346)	110 ± 6			[1995FON/MUN]
	$\Delta_{\text{sub}}H$ (plastic)	(354–372)	81 ± 8			[1995FON/MUN]
	$\Delta_{\text{sub}}H$ (plastic)		86.5 ± 4.3	368	C	[1994FON/MUN]
	$\Delta_{\text{sub}}H$ (cryst)		114.5 ± 5.7	339	C	[1994FON/MUN]
C₄H₁₁NO₂S	[6338-68-7]	N,N-dimethylethanesulfonamide				
	$\Delta_v H$	(384–517)	54.3	399	A	[1987STE/MAL]
C₄H₁₁NO₃	[77-86-1]	2-amino-2-hydroxymethylpropane-1,3-diol				
	$\Delta_{\text{trs}}H$		33.48	409.2		
	$\Delta_{\text{fus}}H$		3.1	444.6	DSC	[2006DIV/CHE]
	$\Delta_{\text{trs}}H$		33.42	407.5		
	$\Delta_{\text{fus}}H$		2.41	443.6		[1990YIN/LIN, 1994LOP/VAN]
C₄H₁₁O₂PS₂	[298-06-6]	O,O-diethyl phosphorodithioate				
	$\Delta_v H$		67.7	298		[2008SAG/SAF]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₄ H ₁₁ O ₃ P	[762-04-9]	diethylphosphite				
	$\Delta_v H$		49.3	298		[2008SAG/SAF]
		(338–471)	38.1	353	A	[1987STE/MAL]
C ₄ H ₁₁ O ₃ P	[na]	dimethyl ethylphosphonate				
	$\Delta_v H$	(333–410)	70.1	348		[1987STE/MAL, 1955KOS, 1984BOU/FRI]
C ₄ H ₁₂ ClN	[3858-78-4]	butylammonium chloride				
	$\Delta_v H$	(489–508)	62.1	498	A	[1987STE/MAL]
C ₄ H ₁₂ ClN	[660-68-4]	diethylamine hydrochloride				
	$\Delta_v H$	(513–558)	177.6	528	A	[1987STE/MAL]
C ₄ H ₁₂ ClN ₂ P	[na]	bis(dimethylamino)chlorophosphine				
	$\Delta_v H$		45.9 ± 1.2	298	STG	[1995ALM/FIN2]
C ₄ H ₁₂ FN ₂ OP	[115-26-4]	bis(dimethylamido)fluorophosphate				
	$\Delta_v H$	(312–350)	50.4	327	A	[1987STE/MAL]
C ₄ H ₁₂ NP	[683-84-1]	dimethyl(dimethylamino)phosphine				
	$\Delta_v H$	(264–372)	36.8	279	A	[1987STE/MAL]
C ₄ H ₁₂ N ₂	[4426-48-6]	(dl) 1,2-butanediamine				
	$\Delta_v H$	(251–293)	50.2	278		[1987STE/MAL, 1975MES/FIN]
	$\Delta_v H$	(251–293)	46.9	298	IP	[1975MES/FIN]
	$\Delta_v H$		46.3 ± 0.2	298	IP	[1965DOU/OSB, 1970GOO/MOO]
C ₄ H ₁₂ N ₂	[110-60-1]	butane-1,4-diamine				
	$\Delta_{\text{fus}} H$		28.06	295.1	DSC	[2002DAL/DEL]
C ₄ H ₁₂ N ₂	[811-93-8]	2-methyl-1,2-propanediamine				
	$\Delta_{\text{trs}} H$		15.46	237.5		
	$\Delta_{\text{fus}} H$		2.23	256.1		[1996DOM/HEA, 1975MES/FIN]
	$\Delta_v H$	(256–293)	47.2	278	IP	[1987STE/MAL, 1975MES/FIN]
	$\Delta_v H$	(256–293)	43.5 ± 0.2	298	IP	[1975MES/FIN]
	$\Delta_v H$		43.6 ± 0.2	298	IP	[1965DOU/OSB, 1970GOO/MOR]
C ₄ H ₁₂ N ₂	[6415-12-9]	tetramethylhydrazine				
	$\Delta_v H$	(290–346)	32.9	305	T	[1987STE/MAL, 1957AYL]
C ₄ H ₁₂ N ₂	[6291-84-5]	3-(methylamino)propylamine				
	$\Delta_v H$	(327–413)	45.9	342	EB	[2008KIM/SVE]
C ₄ H ₁₂ N ₂ O	[111-41-1]	N-(2-hydroxyethyl)ethylenediamine				
	$\Delta_v H$	(383–517)	62.8	398	A	[1987STE/MAL]
C ₄ H ₁₂ N ₂ OS	[3768-60-3]	tetramethyl sulfurous diamide				
	$\Delta_v H$	(320–351)	41.9	335	A	[1987STE/MAL, 1999DYK/SVO]
C ₄ H ₁₂ N ₂ O ₂ S	[3768-63-6]	N,N,N',N'-tetramethylsulfamide				
	$\Delta_v H$	(358–495)	53.2	373	A	[1987STE/MAL, 1999DYK/SVO]
C ₄ H ₁₂ N ₂ S	[2129-20-6]	tetramethylsulfoxylic diamide				
	$\Delta_v H$	(301–326)	40.4	313	A	[1987STE/MAL]
C ₄ H ₁₃ NP ₂	[98023-09-7]	bis(dimethylphosphino)amine				
	$\Delta_{\text{sub}} H$	(300–310)	61.7	305		[1953WAG/BUR]
C ₄ H ₁₃ N ₃	[111-40-0]	2,2'-diaminodiethylamine				
	$\Delta_v H$	(371–521)	63.4 ± 0.7	298	EB	[1999RIB/MAT2]
	$\Delta_v H$	(371–441)	54.8	386	A	[1987STE/MAL]

TABLE 5. Phase change enthalpies of C₁ to C₄ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
	$\Delta_{\text{fus}}H$		NA		DSC	[2006BAD/DEL]
		Note: compound decomposed on melting				

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₅ BrF ₁₂ N	[4908-96-7] $\Delta_v H$	1,1,2,3,3,3-hexafluoro-2-bromo-N,N-bis(trifluoromethyl)propylamine (324–351)	30.2	337	A	[1987STE/MAL, 1965HAS/TIP]
C ₅ ClF ₅	[30221-57-9] $\Delta_v H$	1-chloro-2,3,4,5,5-pentafluoro-1,3-cyclopentadiene (273–303)	31.0	288	A	[1987STE/MAL]
C ₅ ClF ₅	[30221-56-8] $\Delta_v H$	5-chloro-1,2,3,4,5-pentafluoro-1,3-cyclopentadiene (283–323)	28.7	298	A	[1987STE/MAL]
C ₅ ClF ₁₀ N	[54120-14-8] $\Delta_v H$	2,2,2-trifluoro-N-[1,2,2,2-tetrafluoro-1-(trifluoromethyl)ethylidene]-ethanimidoyl chloride	28.6	361		[1975PET/SHR2]
C ₅ ClF ₁₂ N	[54566-78-8] $\Delta_v H$	N-chloro-1,1,2,3,3,3-heptafluoro-N-(pentafluoroethyl)-2-propanamine	28.6	346		[1975PET/SHR2]
C ₅ Cl ₂ F ₆	[706-79-6] $\Delta_v H$ $\Delta_v H$	1,2-dichlorohexafluorocyclopentene	33.0 36.5	298		[1959YEN/REE] [1959YEN/REE]
C ₅ Cl ₂ F ₉ N	[54566-77-7] $\Delta_v H$	1,1-dichloro-2,2,2-trifluoro-N-[2,2,2-trifluoro-1-(trifluoromethyl)-ethylidene]ethanamine	31.2	361		[1975PET/SHR2]
C ₅ Cl ₅ F ₇ O	[61196-11-0] $\Delta_v H$ $\Delta_v H$	(1,1,2-trifluoro-2,2-dichloroethyl)(2,2,3,3-tetrafluoro-1,1,3-trichloropropyl) ether (362–449)	45.3 50.7 ± 0.8	377 298	A EB	[1987STE/MAL] [1976AMM/BUL]
C ₅ Cl ₆	[77-47-4] $\Delta_{\text{sub}} H$ $\Delta_v H$ $\Delta_v H$	hexachlorocyclopentadiene (335–512)	73.6 53.7 67.4	283 350	B A	[1963BON, 1958UNG/MCB] [1987STE/MAL] [1977LYU/SMO]
C ₅ Cl ₈	[706-78-5] $\Delta_v H$	octachlorocyclopentene	83.4			[1977LYU/SMO]
C ₅ F ₅ N	[700-16-3] $\Delta_v H$	perfluoropyridine (273–363)	36.3	288	A	[1987STE/MAL, 1961BAN/GIN, 1972DYK]
C ₅ F ₈	[21972-01-1] $\Delta_v H$	perfluoro-1,2-pentadiene (262–276)	26.1	269	A	[1987STE/MAL]
C ₅ F ₉ N	[19451-91-3] $\Delta_v H$	3,3,3-trifluoro-N,N-bis(trifluoromethyl)-1-propylamine (277–293)	24.9	285	A	[1987STE/MAL]
C ₅ F ₉ N	[714-37-4] $\Delta_v H$	2,3,4,5-tetrahydro-nafluoropyridine (249–310)	29.3	264	A	[1987STE/MAL]
C ₅ F ₉ NO	[52225-57-7] $\Delta_v H$	2,2,2-trifluoro-N-[2,2,2-trifluoro-1-(trifluoromethyl)ethylidene]-acetamide	32.1	319		[1974PET/SHR]
C ₅ F ₉ NO	[4827-67-2] $\Delta_v H$	3,3,4,5,6,6-hexafluoro-3,6-dihydro-2-trifluoromethyl-2H-1,2-oxazine (263–323)	31.4	278	A	[1987STE/MAL, 1965BAN/BAR]
C ₅ F ₉ NO ₃ S	[34805-64-6] $\Delta_v H$	nonafluoro-1-butanesulfonyl isocyanate (309–401)	48.2	324	A	[1987STE/MAL]
C ₅ F ₁₀	[376-77-2] $\Delta_{\text{us}} H$ $\Delta_{\text{fus}} H$	perfluorocyclopentane	4.95 2.99	118.2 238.5		[1951BUR/CAD]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
		$\Delta_{\text{sub}}H$	(229–281)	32.1	266	[1987STE/MAL, 1967CRO/TAY]
		$\Delta_{\text{sub}}H$		38.2	115	[1963BON, 1951BUR/CAD, 1956BAR/CAD]
		Δ_vH	(285–297)	27	291	A [1987STE/MAL]
		Δ_vH	(290–330)	25.6	298	[1984BOU/FRI, 1991BAS/SVO]
		Δ_vH	(290–329)	26.3	298	[1956BAR/CAD]
C ₅ F ₁₀ N ₂ O ₂	[32822-52-9]	decafluoroglutaramide				
		Δ_vH		35.6	368	HG [1971DEM/SHR]
C ₅ F ₁₀ N ₂ O ₂	[1840-07-9]	1-nitrodecafluoropiperadine				
		Δ_vH	(283–343)	29.6	298	A [1987STE/MAL, 1964BAN/CHE]
C ₅ F ₁₀ O ₂	[55064-79-4]	carbonofluoric acid, 2,2,2-trifluoro-1,1-bis(trifluoromethyl) ethyl ether				
		Δ_vH	(275–305)	32.2	290	A [1987STE/MAL, 1975WAL/DES2]
C ₅ F ₁₀ O ₃ S	[2993-14-8]	perfluorocyclopentyl fluorosulfate				
		Δ_vH	(255–360)	36.6	307	[1963GIL/CAD]
C ₅ F ₁₀ O ₆ S ₂	[741-20-8]	octafluorocyclopentanediol bis(fluorosulfate)				
		Δ_vH	(334–423)	49.5	349	[1972DYK, 1987STE/MAL, 1999DYK/SVO]
C ₅ F ₁₁ N	[836-77-1]	perfluoropiperidine				
		$\Delta_{\text{trs}}H$		6.63	161	
		$\Delta_{\text{trs}}H$		1.84	171.9	
		$\Delta_{\text{fus}}H$		2.82	274.1	[1996DOM/HEA]
		Δ_vH	(302–355)	30	317	A [1987STE/MAL, 1963GOO/TOD, 1972DYK]
C ₅ F ₁₁ N	[2344-10-7]	octafluoro-1-(trifluoromethyl)pyrrolidine				
		Δ_vH	(249–306)	29.4	264	A [1987STE/MAL]
C ₅ F ₁₁ NO	[52225-65-7]	N,2,2,2-tetrafluoro-N-[1,2,2,2-tetrafluoro-1-(trifluoromethyl)ethyl]-acetamide				
		Δ_vH		32.6	332	[1974PET/SHR]
C ₅ F ₁₂	[594-21-2]	perfluoro-2-methylbutane				
		Δ_vH	(290–340)	26.3	298	[1984BOU/FRI, 1991BAS/SVO]
		Δ_vH	(228–308)	31.0	243	A [1987STE/MAL, 1967CRO/TAY]
		Δ_vH	(290–337)	27.4	298	[1956BAR/CAD]
C ₅ F ₁₂	[678-26-2]	perfluoropentane				
		$\Delta_{\text{fus}}H$		6.8	147.8	[1951BUR/CAD]
		$\Delta_{\text{sub}}H$		43.7	145	[1963BON, 1951BUR/CAD, 1956BAR/CAD]
		Δ_vH	(280–340)	26.6	298	[1984BOU/FRI, 1991BAS/SVO]
		Δ_vH	(221–303)	31.1	236	A [1987STE/MAL, 1967CRO/TAY]
		Δ_vH	(288–338)	27.5	298	[1956BAR/CAD]
C ₅ F ₁₂ N ₂	[53684-06-3]	[2,2,2-trifluoro-1,1-bis(trifluoromethyl)ethyl](trifluoromethyl)-diazene				
		Δ_vH		23.7	309	[1975KIR/LAS]
C ₅ F ₁₂ O ₂	[20822-11-1]	bis(pentafluoroethoxy)difluoromethane				
		Δ_vH	(246–299)	32.7	261	A [1987STE/MAL]
C ₅ F ₁₂ O ₂ S	[52225-54-4]	trifluoromethanesulfinic acid, 2,2,2-trifluoro-1,1-bis(trifluoromethyl) ethyl ether				
		Δ_vH		37.7	355	HG [1974MAJ/SHR]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₅ F ₁₂ O ₄ S	[60672-63-1] $\Delta_v H$		39.2			[1976HOP/DES]
C ₅ F ₁₃ N	[1481-55-6] $\Delta_v H$ $\Delta_v H$		(298–319) 30.2 29.4 ± 0.4	308 298	A	[1987STE/MAL] [1977VAR/AMM2]
C ₅ F ₁₃ N	[758-48-5] $\Delta_{\text{fus}} H$		7.16	149.7		[1996DOM/HEA]
C ₅ F ₁₃ NS	[37826-44-1] $\Delta_v H$		(314–360) 31.3		A	[1987STE/MAL, 1999DYK/SVO]
C ₅ F ₁₄ N ₂ O	[17636-89-4] $\Delta_v H$		(302–311) 34.7		A	[1987STE/MAL]
C ₅ F ₁₄ N ₂ O	[17636-88-3] $\Delta_v H$		(282–323) 33.7	297	A	[1987STE/MAL]
C ₅ F ₁₄ OS	[736-59-4] $\Delta_v H$		(300–361) 36.1	315	A	[1987STE/MAL, 1999DYK/SVO]
C ₅ F ₁₅ N	[758-48-5] $\Delta_v H$		29.4 ± 0.4	298		[1977VAR/AMM]
C ₅ F ₁₅ NS	[65844-10-2] $\Delta_v H$		32.2	375	I	[1978KIT/SHR]
C ₅ F ₁₅ P ₅	[745-23-3] $\Delta_v H$		(319–435) 51.8	334	A,SG	[1987STE/MAL, 1958MAH/BUR]
C ₅ N ₄	[24331-09-7] $\Delta_{\text{sub}} H$		61.1 ± 8.8	298	C	[1973BAR/MOR2]
C ₅ O ₂	[51799-36-1] $\Delta_v H$		(186–273) 4.6	258	A	[1987STE/MAL, 1937KLE/WAG]
C ₅ HCIF ₈ O ₂	[52225-55-5] $\Delta_v H$		37.2	338	HG	[1974MAJ/SHR]
C ₅ HF ₁₀ NO	[52225-63-5] $\Delta_v H$		42.3	367		[1974PET/SHR]
C ₅ HF ₉	[376-65-8] $\Delta_v H$ $\Delta_v H$		(289–348) 29.6 (289–348) 29.4	304 298	A	[1987STE/MAL, 1956BAR/CAD] [1956BAR/CAD]
C ₅ HF ₉ IN	[20257-34-5] $\Delta_v H$		(343–366) 31.3	354	A	[1987STE/MAL, 1968FRE/TIP]
C ₅ HF ₉ IN	[20257-35-6] $\Delta_v H$		(345–368) 35	356	A	[1987STE/MAL, 1968FRE/TIP]
C ₅ HF ₉ O ₂	[42031-15-2] $\Delta_v H$		28.5	321	HG	[1973MAJ/SHR]
C ₅ HF ₁₀ N	[559-31-9] $\Delta_v H$		(273–313) 32.7	288	A	[1987STE/MAL, 1964BAN/CHE]
C ₅ HF ₁₂ N	[54566-80-2] $\Delta_v H$		29.8	325		[1975PET/SHR2]
C ₅ HN ₃	[997-76-2]					ethylenetricarbonitrile

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(313–343)	66.0	328	A, MG	[1987STE/MAL, 1963BOY]
C ₅ H ₂ BrF ₈ N	[19451-93-5]	2-bromo-3,3-difluoro-N,N-bis(trifluoromethyl)allylamine				
	$\Delta_v H$	(336–367)	33.8	351	A	[1987STE/MAL, 1968HAS/TIP]
C ₅ H ₂ Cl ₃ N	[16063-70-0]	2,3,5-trichloropyridine				
	$\Delta_{\text{sub}} H$		74.4 ± 1.5	298	C	[2005GOM/AMA]
C ₅ H ₂ Cl ₃ NO	[6515-38-4]	3,5,6-trichloro-2-pyridinol				
	$\Delta_{\text{fus}} H$		25.79	448.1		[1991ACR]
	$\Delta_v H$	(373–403)	63.0		GC	[2007GOE/MCC]
C ₅ H ₂ F ₆ N ₂	[14704-41-7]	3,5-bis(trifluoromethyl)pyrazole				
	$\Delta_{\text{sub}} H$		69.0 ± 0.6	266	ME	[1991ELG/YRA]
C ₅ H ₂ F ₆ O ₂	[1522-22-1]	1,1,1,5,5,5-hexafluoropentan-2,4-dione				
	$\Delta_v H$	(273–330)	33.1	301	GS	[1998GEO/YOU]
	$\Delta_v H$		30.6 ± 0.1	298		[1997RIB/GON, 1975IRV/RIB, 1978RIB/IRV]
C ₅ H ₂ F ₉ N	[25273-42-1]	<i>trans</i> 3,3,3-trifluoro-N,N-bis(trifluoromethyl)propenylamine				
	$\Delta_v H$	(287–319)	28.2	302	A	[1987STE/MAL, 1968HAS/TIP]
C ₅ H ₂ F ₉ NOS	[62067-07-6]	2,2,2-trifluoro-N-[(trifluoromethyl)thio]ethanimidic acid, 2,2,2-trifluoroethyl ester				
	$\Delta_v H$		35.8	373	I	[1977BUR/SHR2]
C ₅ H ₂ F ₉ NS	[57682-29-8]	2,2,2-trifluoro-N-[2,2,2-trifluoro-1-(trifluoromethyl)ethyl]ethanethioamide				
	$\Delta_v H$		36.9			[1975PET/SHR3]
C ₅ H ₂ F ₁₀	[138495-42-8]	1,1,1,2,2,3,4,5,5,5-decafluoropentane				
	$\Delta_v H$	(289–326)	33.4	298		[2001LOR/AUC]
C ₅ H ₂ F ₁₀	[na]	<i>threo</i> 1,1,1,2,2,3,4,5,5,5-decafluoropentane				
	$\Delta_v H$	(293–328)	40.8	298	EB	[2004KAO/SIE]
C ₅ H ₂ F ₁₀	[na]	<i>erythro</i> 1,1,1,2,2,3,4,5,5,5-decafluoropentane				
	$\Delta_v H$	(293–328)	37.9	298	EB	[2004KAO/SIE]
C ₅ H ₂ F ₁₀ O	[142469-08-7]	1,1,1,2,2,3,3-heptafluoro-3-(2,2,2-trifluoroethoxy)propane				
	$\Delta_v H$	(288–325)	31.5	303	I	[2002MUR/YAM]
C ₅ H ₂ F ₁₀ O	[347148-74-7]	1,1,1,2,4,4,4-heptafluoro-2-(trifluoromethoxy)butane				
	$\Delta_v H$	(288–323)	31.8	303	I	[2002MUR/YAM]
C ₅ H ₂ F ₁₀ O	[155653-44-4]	1,1,1,2,2-pentafluoro-3-(pentafluoroethoxy)propane				
	$\Delta_v H$	(288–320)	31.2	303	I	[2002MUR/YAM]
C ₅ H ₂ F ₁₀ O ₃	[188690-77-9]	1-(difluoromethoxy)-2-[(difluoromethoxy)difluoromethoxy]-1,1,2,2-tetrafluoroethane				
	$\Delta_v H$	(263–357)	36.5 ± 0.7			[1999MAR/BAS]
C ₅ H ₂ N ₄ O ₆	[78013-51-1]	2,4,6-trinitropyridine				
	$\Delta_{\text{fus}} H$		22.0	436.2		[1988LIC/RIT]
	$\Delta_{\text{sub}} H$	(335–357)	101.7 ± 2.9			[1995LEB/CHI]
C ₅ H ₂ N ₄ O ₇	[25242-76-6]	2,4,6-trinitropyridine N-oxide				
	$\Delta_{\text{sub}} H$	(377–403)	106.3 ± 2.9			[1995LEB/CHI]
C ₅ H ₃ BrF ₉ N	[19451-92-4]	2-bromo-3,3,3-trifluoro-N,N-bis(trifluoromethyl)propylamine				
	$\Delta_v H$	(342–365)	34.2	353	A	[1987STE/MAL, 1968FRE/TIP]
C ₅ H ₃ Br ₂ N	[624-28-2]	2,5-dibromopyridine				

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
		$\Delta_{\text{sub}}H$	82.1 ± 2.2	298	C	[1997RIB/MAT]
C ₅ H ₃ Br ₂ N	[626-05-1]	2,6-dibromopyridine				
		$\Delta_{\text{sub}}H$	85.6 ± 3.0	298	C	[1997RIB/MAT]
C ₅ H ₃ Cl ₂ N	[2402-77-9]	2,3-dichloropyridine				
		$\Delta_{\text{sub}}H$	73.5 ± 3.1	298	C	[1997RIB/MAT]
C ₅ H ₃ Cl ₂ N	[16110-09-1]	2,5-dichloropyridine				
		$\Delta_{\text{sub}}H$	67.1 ± 2.0	298	C	[1997RIB/MAT]
C ₅ H ₃ Cl ₂ N	[2402-78-0]	2,6-dichloropyridine				
		$\Delta_{\text{sub}}H$	72.0 ± 1.6	298	C	[1997RIB/MAT]
C ₅ H ₃ Cl ₂ N	[2457-47-8]	3,5-dichloropyridine				
		$\Delta_{\text{sub}}H$	67.3 ± 1.9	298	C	[1997RIB/MAT]
C ₅ H ₃ F ₃ N ₂ O ₂	[54-20-6]	5-(trifluoromethyl)uracil				
		$\Delta_{\text{sub}}H$	(373–392) 108.5 ± 0.9	382	ME	[2004ZIE/SZT]
		$\Delta_{\text{sub}}H$	(373–392) 110.8 ± 0.9	298	ME	[2004ZIE/SZT]
C ₅ H ₃ F ₆ N	[25237-11-0]	N,N-bis(trifluoromethyl)-1-propynylamine				
		Δ_vH	(295–312) 31.1	303	A	[1987STE/MAL]
C ₅ H ₃ F ₇ O ₂	[356-24-1]	methyl perfluorobutyrate				
		$\Delta_{\text{fus}}H$	11.77	191.4		[1996DOM/HEA]
		Δ_vH	34.5			[1977DIT/KOL, 1978KOL/DIT]
C ₅ H ₃ F ₈ NOS	[77589-48-1]	2,2,3,3,4,4,5,5-octafluoro-1,1,2,3,4,5-hexahydro-1-(methylimino)-thiophene-1-oxide				
		Δ_vH	33.9	330		[1981ABE/SHR]
C ₅ H ₃ F ₉ N ₂ OS	[62609-63-6]	1,1,1-trifluoro-N'-methyl-N-[2,2,2-trifluoro-1-(trifluoromethyl)-ethylidene]methanesulfonimidamide				
		Δ_vH	32.6	417	I	[1977KIT/SHR]
C ₅ H ₃ F ₉ O	[176310-27-3]	1,1,2,2-tetrafluoro-3-(pentafluoroethoxy)propane				
		Δ_vH	(288–336) 34.0	303	I	[2002MUR/YAM]
C ₅ H ₃ F ₉ O	[176310-28-4]	1-(2,2-difluoroethoxy)-1,1,2,2,3,3,3-heptafluoropropane				
		Δ_vH	(288–340) 34.8	303	I	[2002MUR/YAM]
C ₅ H ₃ F ₉ O	[50807-74-4]	1,1,1,2,2-pentafluoro-3-(1,1,2,2-tetrafluoroethoxy)propane				
		Δ_vH	(293–343) 35.6	308	I	[2002MUR/YAM]
C ₅ H ₃ F ₉ O	[439152-54-2]	1,1,1,2,4,4-hexafluoro-2-(trifluoromethoxy)butane				
		Δ_vH	(283–332) 33.8	298	I	[2002MUR/YAM]
C ₅ H ₃ F ₉ O	[66670-22-2]	1,1,1,3,3,3-hexafluoro-2-methoxy-2-(trifluoromethyl)propane				
		Δ_vH	(288–326) 31.3	303	I	[2002MUR/YAM]
C ₅ H ₃ F ₉ O	[993-95-3]	1,1,1,2,3,3-hexafluoro-3-(2,2,2-trifluoroethoxy)propane				
		Δ_vH	(293–346) 36.1	308	I	[2002MUR/YAM]
C ₅ H ₃ F ₉ O	[69948-43-2]	1,1,1,2,3,3-hexafluoro-4-(trifluoromethoxy)butane				
		Δ_vH	(288–338) 34.0	303	I	[2002MUR/YAM]
C ₅ H ₃ F ₉ O ₂ S	[52225-51-1]	trifluoromethanesulfinic acid, 2,2,2-trifluoro-1-methyl-1-(trifluoromethyl)ethyl ester				
		Δ_vH	34.3	385	HG	[1974MAJ/SHR]
C ₅ H ₃ NO	[617-90-3]	2-furancarbonitrile				
		Δ_vH	44.8 ± 0.4	298	C	[2009RIB/AMA]
C ₅ H ₃ NO ₃	[698-63-5]	5-nitro-2-furancarboxaldehyde				

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{sub}}H$		75.3 ± 2.1			[1980BAL/LEB, 1986PED/NAY]
C ₅ H ₃ NS	[1003-31-2] Δ_vH	2-thiophenecarbonitrile	49.5 ± 1.1	298	C	[2008RIB/SAN]
C ₅ H ₃ NS	[1641-09-4] Δ_vH	3-thiophenecarbonitrile	51.6 ± 1.9	298	C	[2008RIB/SAN]
C ₅ H ₃ NS	[10359-20-3] $\Delta_{\text{fus}}H$	2,2-dicyanopropionitrile	18.7	367.2		[1994RAK/VER]
	$\Delta_{\text{sub}}H$	(293–333)	73.9 ± 0.5	313	T	[1994RAK/VER]
	Δ_vH	(293–333)	55.2		B	[1994RAK/VER]
C ₅ H ₃ N ₃	[19847-12-2] Δ_vH	pyrazinecarbonitrile	58.7 ± 1.2	298	C	[2005RIB/MIR]
C ₅ H ₄ BrF ₆ N	[25273-47-6] Δ_vH	<i>cis</i> 2-bromo-N,N-bis(trifluoromethyl)propenylamine (346–367)	35.3	356	A	[1987STE/MAL]
C ₅ H ₄ BrF ₆ N	[25273-48-7] Δ_vH	<i>trans</i> 2-bromo-N,N-bis(trifluoromethyl)propenylamine (336–360)	33.3	348	A	[1987STE/MAL]
C ₅ H ₄ BrN	[109-04-6] Δ_vH	2-bromopyridine	54.4 ± 1.3	298	C	[1997RIB/MAT]
C ₅ H ₄ BrN	[626-55-1] Δ_vH Δ_vH	3-bromopyridine (289–447)	52.1 ± 1.3 47.4	298 304	C A	[1997RIB/MAT] [1987STE/MAL, 1947STU]
C ₅ H ₄ ClN	[109-09-1] Δ_vH Δ_vH	2-chloropyridine (286–444)	51.0 ± 1.2 53	298 301	C A	[1997RIB/MAT2] [1987STE/MAL, 1947STU]
C ₅ H ₄ ClN	[626-60-8] Δ_vH	3-chloropyridine	47.9 ± 1.1	298	C	[1997RIB/MAT2]
C ₅ H ₄ F ₄ N ₄ O ₁₀	[58715-08-5] Δ_vH	<i>bis</i> (2-fluoro-2,2-dinitroethyl)difluoroformal (323–357)	72.7	340		[1997MIN/BEH]
C ₅ H ₄ F ₇ I	[1513-88-8] Δ_vH	1,1,1,2,2,3,3-heptafluoro-5-iodopentane (317–386)	38.7	332	A	[1987STE/MAL]
C ₅ H ₄ F ₈ O	[16627-68-2] Δ_vH	1,1,2,2-tetrafluoro-3-(1,1,2,2-tetrafluoroethoxy)propane (293–366)	40.2	308	I	[2002MUR/YAM]
C ₅ H ₄ F ₈ O	[382-26-3] Δ_vH	1,1,1,3,3-pentafluoro-3-methoxy-2-trifluoromethylpropane (288–343)	34.5	303	I	[2002MUR/YAM]
C ₅ H ₄ F ₉ N	[19451-89-9] Δ_vH	3,3,3-trifluoro-N,N-bis(trifluoromethyl)propylamine (290–333)	31	305	A	[1987STE/MAL, 1968FRE/TIP]
C ₅ H ₄ N ₂	[37580-43-1] Δ_vH	<i>cis</i> 2-methyl-2-butenedinitrile (395–467)	58.5	410	A	[1987STE/MAL, 1972DYK]
C ₅ H ₄ N ₂	[37580-44-2] Δ_vH	<i>trans</i> 2-methyl-2-butenedinitrile (339–411)	47.9	354	A	[1987STE/MAL, 1972DYK]
C ₅ H ₄ N ₂ O ₃	[1124-33-0] $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$	4-nitropyridine-N-oxide (311–335)	108.9 ± 0.3 89.1 ± 2.5	298	C	[1995ACR/TUC] [1995LEB/CHI]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₅ H ₄ N ₄	[275-02-5]	1,2,4-triazolo[1,5-a]pyrimidine				
	$\Delta_{\text{fus}}H$		19.4	419.5		[1997STE/CHI4]
	$\Delta_{\text{sub}}H$		86.9	419		[1997STE/CHI4]
	Δ_vH	(370–523)	82.5 ± 13.1	298	EB	[1997STE/CHI4]
	Δ_vH	(370–523)	63.5 ± 2.2	480	EB	[1997STE/CHI4]
C ₅ H ₄ N ₄	[120-73-0]	purine				
	$\Delta_{\text{sub}}H$		NA			[1974TEP/SUK]
C ₅ H ₄ N ₄ O	[68-94-0]	hypoxanthine				
	$\Delta_{\text{sub}}H$	(423–473)	158.1 ± 1.6	448		[1975TEP/YAN]
C ₅ H ₄ N ₄ S	[6112-76-1]	6-mercaptopurine				
	$\Delta_{\text{sub}}H$	(413–458)	148.5 ± 1.5	435		[1975TEP/YAN]
C ₅ H ₄ OS	[98-03-3]	2-thiophenecarboxyaldehyde				
	Δ_vH		54.9 ± 1.1	298	C	[2008RIB/SAN2]
C ₅ H ₄ OS	[498-62-4]	3-thiophenecarboxyaldehyde				
	Δ_vH		52.6 ± 1.2	298	C	[2008RIB/SAN2]
C ₅ H ₄ O ₂	[98-01-1]	2-furfuraldehyde				
	$\Delta_{\text{fus}}H$		14.37	235.1		[1996DOM/HEA]
	Δ_vH	(277–323)	50.7 ± 0.2	298	GS	[2007EME/DAB]
	Δ_vH	(357–435)	44.7	372	A	[1987STE/MAL]
	Δ_vH	(366–394)	50.7 ± 0.2	298	EB	[1987HAU/WU, 2007EME/DAB]
	Δ_vH	(329–433)	48.2	344		[1950MAT/SUM, 1984BOU/FRI]
	Δ_vH	(365–443)	47.6	380		[1926EVA/AYL, 1984BOU/FRI]
C ₅ H ₄ O ₂	[498-60-2]	3-furandaldehyde				
	Δ_vH		48.1 ± 0.5	298	C	[2009RIB/AMA]
C ₅ H ₄ O ₂ S	[527-72-0]	2-thiophene carboxylic acid				
	$\Delta_{\text{fus}}H$		21	400.9	DSC	[2003ROU/TEM]
	$\Delta_{\text{sub}}H$	(315–323)	97.1	319	E	[1953BRA/CAR, 1960JON]
C ₅ H ₄ O ₂ S	[88-13-1]	3-thiophenecarboxylic acid				
	$\Delta_{\text{fus}}H$		18.3	412.9	DSC	[2003ROU/TEM]
C ₅ H ₄ O ₃	[616-02-4]	citraconic anhydride				
	Δ_vH	(320–487)	53.3	335	A	[1987STE/MAL, 1947STU]
C ₅ H ₄ O ₃	[88-14-2]	2-furancarboxylic acid				
	$\Delta_{\text{fus}}H$		22.6	402.5	DSC	[2004ROU/TEM2]
	$\Delta_{\text{sub}}H$	(285–304)	88.4 ± 1.5	298	ME	[2003ROU/TEM3]
C ₅ H ₄ O ₃	[88-14-2]	2-furancarboxylic acid				
	$\Delta_{\text{sub}}H$	(317–328)	108.4 ± 2.2		ME	[1953BRA/CAR, 1960JON, 1970COX/PIL]
C ₅ H ₄ O ₃	[488-93-7]	3-furancarboxylic acid				
	$\Delta_{\text{fus}}H$		21.3	394.8	DSC	[2004ROU/TEM2]
C ₅ H ₄ O ₃	[488-93-7]	3-furancarboxylic acid				
	$\Delta_{\text{sub}}H$	(283–298)	87.1 ± 0.5	298	ME	[2003ROU/TEM3]
C ₅ H ₅ ClN ₂ O ₂	[31737-09-4]	1-methyl-6-chlorouracil				

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{sub}}H$	(417–465)	108.8 ± 8		HSA	[1978NOW/SZC]
C ₅ H ₅ ClN ₂ O ₂	[4318-56-3]	3-methyl-6-chlorouracil				
	$\Delta_{\text{sub}}H$	(444–493)	104.6 ± 6		HSA	[1978NOW/SZC]
C ₅ H ₅ Cl ₃ OS	[76619-92-6]	2,3,3-trichloro-2-propenethioic acid, O-ethyl ester				
	Δ_vH	(383–423)	66.9		GC	[1980PIT/KIS]
C ₅ H ₅ FN ₂ O ₂	[155-16-8]	1-methyl-5-fluorouracil				
	$\Delta_{\text{sub}}H$	(381–423)	116 ± 2		TE	[2002BRU/POR]
	$\Delta_{\text{sub}}H$	(480–515)	125.5 ± 8		HSA	[1978NOW/SZC]
C ₅ H ₅ FN ₂ O ₂	[4840-69-1]	3-methyl-5-fluorouracil				
	$\Delta_{\text{sub}}H$	(465–487)	79.5 ± 17		HSA	[1978NOW/SZC]
C ₅ H ₅ F ₃ N ₂	[10010-93-2]	3(5)-trifluoromethyl-5(3)-methylpyrazole				
	$\Delta_{\text{sub}}H$		78.2 ± 0.8	297	ME	[1991ELG/YRA]
C ₅ H ₅ F ₃ O ₂	[367-57-7]	1,1,1-trifluoropentane-2,4-dione				
	Δ_vH		37.2 ± 0.2	298		[1997RIB/GON, 1975IRV/RIB, 1978RIB/IRV]
C ₅ H ₅ F ₃ O ₂	[7291-30-7]	trifluoromethyl (2-hydroxy-1-propenyl)ketone				
	$\Delta_{\text{fus}}H$		8.45	232.4		[1996DOM/HEA]
C ₅ H ₅ F ₆ NO	[22743-77-7]	N,N-bis(trifluoromethyl)allylamine-N-oxide				
	Δ_vH	(254–328)	33.1	269	A	[1987STE/MAL]
C ₅ H ₅ F ₆ NO	[22130-39-8]	1-methoxy-N,N-bis(trifluoromethyl)vinylamine				
	Δ_vH	(321–343)	32.4	332	A	[1987STE/MAL, 1969FRE/TIP]
C ₅ H ₅ F ₆ NO	[22298-35-7]	<i>cis</i> 2-methoxy-N,N-bis(trifluoromethyl)vinylamine				
	Δ_vH	(341–362)	32.5	351	A	[1987STE/MAL, 1969FRE/TIP]
C ₅ H ₅ F ₆ NO ₂	[22743-66-4]	N,N-bis(trifluoromethyl)propionamide-N-oxide				
	Δ_vH	(278–361)	42.1	293	A	[1987STE/MAL, 1968NAS/BAB]
C ₅ H ₅ F ₇ O	[200501-98-0]	1,1,1,2-tetrafluoro-2-(trifluoromethoxy)butane				
	Δ_vH	(283–319)	30.3	298	I	[2002MUR/YAM]
C ₅ H ₅ F ₇ O	[22052-86-4]	1-ethoxy-1,1,2,2,3,3,3-heptafluoropropane				
	Δ_vH	(288–323)	31.0	303	I	[2002MUR/YAM]
C ₅ H ₅ F ₇ O	[376-98-7]	1,1,1,2,2,3,3-heptafluoro-4-methoxybutane				
	Δ_vH	(293–344)	34.6	308	I	[2002MUR/YAM]
C ₅ H ₅ N	[2180-69-0]	<i>cis</i> 2,4-pentadienenitrile (<i>cis</i> 1-cyano-1,3-butadiene)				
	Δ_vH	(318–383)	41.4	333	A	[1987STE/MAL, 1972DYK]
	Δ_vH		40.7	348		[1954WIS]
	Δ_vH		38.3	408		[1954WIS]
C ₅ H ₅ N	[16955-35-4]	bicyclo[1.1.0]butane-1-carbonitrile				
	Δ_vH	(307–349)	48.0	319	BG	[1971HAL/BAL]
C ₅ H ₅ N	[110-86-1]	pyridine				
	$\Delta_{\text{fus}}H$		8.28	231.5		[1996DOM/HEA]
	Δ_vH	(342–373)	40.5 ± 1.1	298	CGC	[2009LIP/CHI2]
	Δ_vH	(289–358)	39.3	324		[1997UKR/SOL]
	Δ_vH		40.16 ± 0.06	298		[1996CHI/STE]
	Δ_vH	(323–373)	40.4	298	CGC	[1995CHI/HOS]
	Δ_vH	(346–362)	37.6	354		[1994BLA/BEL]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		T _m (K)	Method	Reference	
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)				
		$\Delta_v H$	(295–388)	39.9	310	EB	[1990LEN]
		$\Delta_v H$	(296–353)	39.7	311	A	[1987STE/MAL]
		$\Delta_v H$	(348–434)	37.3	363	A	[1987STE/MAL]
		$\Delta_v H$	(431–558)	35.0	446	A	[1987STE/MAL]
		$\Delta_v H$	(552–620)	34.0	567	A	[1987STE/MAL]
		$\Delta_v H$	(298–333)	39.6	313	C	[1986MIC/JOS]
		$\Delta_v H$		40.2	298	C	[1984MAJ/SVO2]
		$\Delta_v H$		39.4	313	C	[1984MAJ/SVO2]
		$\Delta_v H$		38.5	328	C	[1984MAJ/SVO2]
		$\Delta_v H$		37.7	343	C	[1984MAJ/SVO2]
		$\Delta_v H$		36.3	368		[1984MAJ/SVO2]
		$\Delta_v H$	(340–426)	37.6	355	EB	[1987STE/MAL, 1957MCC/DOU]
		$\Delta_v H$		37.5 ± 0.1	346	C	[1957MCC/DOU]
		$\Delta_v H$		36.4 ± 0.1	366	C	[1957MCC/DOU]
		$\Delta_v H$		35.1 ± 0.1	388	C	[1957MCC/DOU]
		$\Delta_v H$	(320–388)	38.4	335	MG	[1953HER/MAR]
		$\Delta_v H$	(258–389)	44.4	273		[1931VAN/MAN]
C ₅ H ₅ NO	[142-08-5]		2-hydroxypyridine				
		$\Delta_{\text{sub}} H$		86.6 ± 1.3	298	C	[1982SUR/SAI, 1986PED/NAY]
C ₅ H ₅ NO	[109-00-2]		3-hydroxypyridine				
		$\Delta_{\text{sub}} H$		88.3 ± 1.3	298	C	[1982SUR/SAI, 1986PED/NAY]
C ₅ H ₅ NO	[626-64-2]		4-hydroxypyridine				
		$\Delta_{\text{sub}} H$		118.6 ± 5.2	298	C	[1992RIB/MAT]
		$\Delta_{\text{sub}} H$		103.8 ± 1.7	298	C	[1982SUR/SAI, 1986PED/NAY]
C ₅ H ₅ NO	[694-59-7]		pyridine N-oxide				
		$\Delta_{\text{sub}} H$		79.3 ± 1.0	298		[1988SHA/PIL]
C ₅ H ₅ NO ₂	[13161-30-3]		2-hydroxypyridine N-oxide				
		$\Delta_{\text{sub}} H$		89.4 ± 0.9	298	C	[2004RIB/MAT]
C ₅ H ₅ NO ₂	[6602-28-4]		3-hydroxypyridine N-oxide				
		$\Delta_{\text{sub}} H$	(345–392)	121.8 ± 4.4	298	ME	[1998RIB/MAT]
C ₅ H ₅ NO ₂	[634-97-9]		pyrrole-2-carboxylic acid				
		$\Delta_{\text{sub}} H$	(331–353)	98.6 ± 0.9	342	ME	[2009SAN/RIB]
		$\Delta_{\text{sub}} H$	(331–353)	100.8 ± 0.9	298	ME	[2009SAN/RIB]
		$\Delta_{\text{sub}} H$	(350–354)	126.8	352	ME	[1953BRA/CAR, 1960JON]
C ₅ H ₅ NO ₂	[930-88-1]		N-methylmaleimide				
		$\Delta_{\text{sub}} H$	(276–289)	75.3 ± 0.5	282	ME	[1997ROU/JIM]
		$\Delta_{\text{sub}} H$		73.3 ± 0.5	298		[1997ROU/JIM]
C ₅ H ₅ NO ₂	[137-05-3]		2-cyanoacrylic acid, methyl ester				
		$\Delta_v H$	(258–283)	57.8	270	A	[1987STE/MAL, 1969WOO/ADI, 1972DYK]
C ₅ H ₅ NO ₂	[16867-04-2]		2,3-dihydroxypyridine				
		$\Delta_{\text{sub}} H$		109.1 ± 4.3	298	C	[2006MOR/MIR]
C ₅ H ₅ N ₃ O	[98-96-4]		pyrazine carboxamide				
		$\Delta_{\text{fus}} H$		30.28	463		[1960NEG/MIK2]
		$\Delta_{\text{sub}} H$	(353–383)	87.9	368	ME	[1987STE/MAL, 1960NEG/MIK, 1959HAR]
C ₅ H ₅ N ₃ O ₂	[4214-76-0]		2-amino-5-nitropyridine				

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{fus}}H$	(80–395)	29.2	461.4	AC	[2007SHI/TAN2]
C ₅ H ₅ N ₅	[73-24-5]	adenine				
	$\Delta_{\text{sub}}H$		140.4		ME	[2000ZIE]
	$\Delta_{\text{sub}}H$	(448–473)	109.2	460.5	A	[1987STE/MAL]
	$\Delta_{\text{sub}}H$	(403–439)	127.2 ± 1.9		ME	[1984ZIE/ZIE]
	$\Delta_{\text{sub}}H$		126.3		LE	[1975YAN/TEP, 1974YAN/VER]
	$\Delta_{\text{sub}}H$		108.7 ± 8		ME	[1965CLA/PES, 1970COX/PIL]
C ₅ H ₅ N ₅ O	[73-40-5]	guanine				
	$\Delta_{\text{sub}}H$	(325–405)	168.3 ± 0.6	365	QR,ME	[2006DEB/MED]
	$\Delta_{\text{sub}}H$		186.2		LE	[1975YAN/TEP, 1974YAN/VER]
C ₅ H ₅ N ₇ O ₁₄	[20919-99-7]	1,1,1,3,5,5,5-heptanitropentane				
	$\Delta_{\text{sub}}H$		111.7	298		[1999MIR/VOR]
C ₅ H ₆	[542-92-7]	1,3-cyclopentadiene				
	$\Delta_{\text{fus}}H$		8.01	176.6		[1996DOM/HEA]
	Δ_vH	(271–314)	28.2	286		[1967LES/OGO, 1984BOU/FRI]
	Δ_vH	(291–314)	28.1	302	A,MM	[1987STE/MAL, 1965HUL/REI]
	Δ_vH	(291–314)	28.4 ± 0.3	298	MM	[1965HUL/REI]
	Δ_vH	(273–287)	29.7	298		[1965HUL/REI, 1933BAR/BUR]
C ₅ H ₆	[6746-94-7]	ethynylcyclopropane				
	Δ_vH	(290–320)	31.1	305	A	[1987STE/MAL]
C ₅ H ₆	[78-80-8]	isopropenylacetylene				
	Δ_vH		27.2			[1977LEB/RYA]
C ₅ H ₆ ClN	[32366-08-8]	4-chloro-3-pentenenitrile				
	Δ_vH	(349–433)	63.9	364	A	[1987STE/MAL]
C ₅ H ₆ Cl ₂ O ₂	[2873-74-7]	glutaryl chloride				
	Δ_vH	(329–490)	55.9	344	A	[1987STE/MAL, 1947STU]
C ₅ H ₆ F ₂ N ₄ O ₁₀	[17003-79-1]	bis(2-fluoro-2,2-dinitroethyl)formal				
	Δ_vH	(323–365)	85.1	344		[1997MIN/BEH]
C ₅ H ₆ F ₃ NO ₃	[383-72-2]	glycine, N-(trifluoroacetyl) methyl ester				
	$\Delta_{\text{sub}}H$	(293–463)	57.3	308		[1987STE/MAL, 1960WEY/KLI]
C ₅ H ₆ F ₆ N ₂ S	[62067-11-2]	2,2,2-trifluoro-N,N-dimethyl-N'-[(trifluoromethyl)thio]ethanimidamide				
	Δ_vH		40.4	400	I	[1977BUR/SHR2]
C ₅ H ₆ F ₆ O	[58705-93-4]	1,1,1,2,3,3-hexafluoro-4-methoxybutane				
	Δ_vH	(293–360)	37.0	308	I	[2002MUR/YAM]
C ₅ H ₆ F ₆ N ₂ S	[38005-19-5]	dimethylamino(hexafluoroisopropylideneimino) sulfur				
	Δ_vH		39.7	383	I	[1972MET/SHR]
C ₅ H ₆ F ₆ O ₂ S	[52225-49-7]	trifluoromethanesulfinic acid, 2,2,2-trifluoro-1,1-dimethylethyl ester				
	Δ_vH		35.6	388	HG	[1974MAJ/SHR]
C ₅ H ₆ F ₆ O ₃ S ₂	[61915-97-7]	3,3-bis[(trifluoromethyl)sulfonyl]-1-propanol				
	Δ_vH	(333–418)	32.8	348	A,I	[1987STE/MAL, 1977BUR/SHR, 1999DYK/SVO]
C ₅ H ₆ N ₂	[7321-55-3]	dimethylmalononitrile				
	$\Delta_{\text{trs}}H$		9.87	302.6		
	$\Delta_{\text{fus}}H$		4.05	307.5		[1996DOM/HEA]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		T _m (K)	Method	Reference	
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)				
		$\Delta_{\text{sub}}H$	62.0 ± 0.7	298		[1990BEC/DOG]	
		$\Delta_{\text{v}}H$	(322–413)	47.5	337	A	[1987STE/MAL, 1967RIB/WES]
C ₅ H ₆ N ₂	[109-08-0]	2-methylpyrazine					
		$\Delta_{\text{v}}H$	(342–373)	43.7 ± 1.9	298	CGC	[2009LIP/CHI2]
		$\Delta_{\text{v}}H$	(288–392)	42.4	340		[1995SAK/UEO]
C ₅ H ₆ N ₂	[1632-76-4]	3-methylpyridazine					
		$\Delta_{\text{v}}H$	(342–373)	49.7 ± 2.8	298	CGC	[2009LIP/CHI2]
C ₅ H ₆ N ₂	[3438-46-8]	4-methylpyrimidine					
		$\Delta_{\text{v}}H$	(342–373)	44.2 ± 2.4	298	CGC	[2009LIP/CHI2]
C ₅ H ₆ N ₂	[544-13-8]	glutaronitrile					
		$\Delta_{\text{fus}}H$		12.03	242	DSC	[2007BAD/BLA]
		$\Delta_{\text{fus}}H$		12.59	244.2		[1996DOM/HEA]
		$\Delta_{\text{v}}H$	(364–560)	60.1	379	A	[1987STE/MAL]
		$\Delta_{\text{v}}H$	(277–303)	66.8	290	A	[1987STE/MAL, 1972DYK, 1960WOO/MUR]
C ₅ H ₆ N ₂	[504-29-0]	2-aminopyridine					
		$\Delta_{\text{fus}}H$		15.3	331.5		[1998SAB/DAS]
		$\Delta_{\text{sub}}H$		76.5 ± 0.4	298	C	[1998SAB/DAS]
		$\Delta_{\text{sub}}H$		38.6 ± 1.9	298	DSC	[1985BRO/INI]
		$\Delta_{\text{sub}}H$		78.7 ± 0.8	298	C	[1984BIC/PIL]
C ₅ H ₆ N ₂	[462-08-8]	3-aminopyridine					
		$\Delta_{\text{fus}}H$		14.4	335.5		[1998SAB/DAS]
		$\Delta_{\text{sub}}H$		80.7 ± 0.3	298	C	[1998SAB/DAS]
		$\Delta_{\text{sub}}H$		84.0 ± 1.4	298	C	[1984BIC/PIL]
C ₅ H ₆ N ₂	[504-24-5]	4-aminopyridine					
		$\Delta_{\text{fus}}H$		20.07	429.9	DSC	[1990DON/DRE]
		$\Delta_{\text{sub}}H$		87.1 ± 0.4	298	C	[1998SAB/DAS]
		$\Delta_{\text{sub}}H$		53.8 ± 0.8	298	DSC	[1985BRO/INI]
		$\Delta_{\text{sub}}H$		88.1 ± 1.1	298	C	[1984BIC/PIL]
C ₅ H ₆ N ₂ OS	[2361-27-5]	2-thiophenecarboxylic acid hydrazide					
		$\Delta_{\text{sub}}H$	(339–361)	110.7 ± 0.5	350	ME	[2008RIB/AMA3]
		$\Delta_{\text{sub}}H$	(339–361)	113.3 ± 0.5	298	ME	[2008RIB/AMA3]
C ₅ H ₆ N ₂ O ₂	[615-77-0]	1-methyluracil					
		$\Delta_{\text{sub}}H$	(343–428)	121.7 ± 4.0	439	TE	[2000BRU/PIA]
		$\Delta_{\text{sub}}H$	(378–418)	112.5 ± 2.6	398	QR	[1980TEP/YAN]
		$\Delta_{\text{sub}}H$	(435–480)	104.6 ± 8	457	HSA	[1978NOW/SZC]
C ₅ H ₆ N ₂ O ₂	[608-34-4]	3-methyluracil					
		$\Delta_{\text{sub}}H$	(344–419)	118.8 ± 3.0	382	TE	[2000BRU/PIA]
		$\Delta_{\text{sub}}H$	(438–498)	75.3 ± 8	463	HSA	[1978NOW/SZC]
C ₅ H ₆ N ₂ O ₂	[65-71-4]	5-methyluracil (thymine)					
		$\Delta_{\text{fus}}H$		17.51	321.3		[1996DOM/HEA]
		$\Delta_{\text{sub}}H$	(383–438)	125.7 ± 3.6	411	ME	[1984BUR/MOR]
		$\Delta_{\text{sub}}H$		131.3 ± 4.0	298		[1984BUR/MOR]
		$\Delta_{\text{sub}}H$	(378–428)	124.4 ± 1.3	403	QR	[1980TEP/YAN]
		$\Delta_{\text{sub}}H$		138 ± 10	298	TE	[1980FER/BEN]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
		$\Delta_{\text{sub}}H$	134.1 ± 4.2	298	C	[1977NAB/SAB]
		$\Delta_{\text{sub}}H$	124.3		LE	[1975YAN/TEP, 1974YAN/VER]
C ₅ H ₆ N ₂ O ₂	[626-48-2]	6-methyluracil				
		$\Delta_{\text{sub}}H$	(426–503) 131	298		[1980FER/BEN2]
C ₅ H ₆ N ₂ O ₂	[3326-71-4]	2-furancarboxylic acid hydrazide				
		$\Delta_{\text{sub}}H$	(309–325) 98.1 ± 0.7	317.2	ME	[2008RIB/AMA3]
		$\Delta_{\text{sub}}H$	(309–325) 99.0 ± 0.7	298	ME	[2008RIB/AMA3]
C ₅ H ₆ O	[534-22-5]	2-methylfuran				
		$\Delta_{\text{fus}}H$	8.55	181.9		[1965CAR/WES]
		Δ_vH	(289–337) 32.4	304		[2002LOR/AUC]
		Δ_vH	(251–338) 34.4	266	A	[1987STE/MAL]
		Δ_vH	(309–339) 31.5	324		[1986KRE/PRA]
		Δ_vH	(288–303) 32.5	295		[1972DYK]
		Δ_vH	(333–373) 30.9	348		[1971EON/POM, 1984BOU/FRI]
		Δ_vH	(215–360) 32.2	298		[1970MOI/ANT]
C ₅ H ₆ O ₂	[591-12-8]	5-methyl-2(3 <i>H</i>)-furanone				
		Δ_vH	(324–442) 40.3	339	A	[1987STE/MAL]
C ₅ H ₆ O ₂	[591-11-7]	(<i>dl</i>) 5-methyl-2(5 <i>H</i>)-furanone				
		Δ_vH	(356–481) 48.2	371	A	[1987STE/MAL]
C ₅ H ₆ O ₂	[98-00-0]	furfuryl alcohol				
		$\Delta_{\text{fus}}H$	13.1	258.6		[1996DOM/HEA]
		Δ_vH	(304–443) 53.6	319	A	[1987STE/MAL, 1947KET/VAN]
C ₅ H ₆ O ₂	[15441-65-3]	5-hydroxy-3-pentyn-2-one				
		Δ_vH	(273–333) 64.4	288	A	[1987STE/MAL, 1972DYK]
C ₅ H ₆ O ₃	[108-55-4]	glutaric anhydride				
		$\Delta_{\text{sub}}H$	(298–320) 85.9 ± 1.6	309	ME	[1990MEN/PIL]
		$\Delta_{\text{sub}}H$	86.1 ± 1.6	298		[1990MEN/PIL]
		Δ_vH	(373–560) 60.9	388	A	[1987STE/MAL, 1947STU]
C ₅ H ₆ O ₃	[4100-80-5]	(<i>dl</i>) methylsuccinic anhydride				
		Δ_vH	(342–521) 59.3	357	A	[1987STE/MAL, 1947STU]
C ₅ H ₆ O ₅	[328-50-7]	α -ketoglutaric acid				
		$\Delta_{\text{fus}}H$	28.59	388.7	DSC	[2005CON/CHI]
		$\Delta_{\text{sub}}H$	(269–285) 100		TPTD	[2005CHA/ZIE]
		Note: Values based on TPTD method are not consistent with values determined by other experimental methods				
C ₅ H ₆ O ₅	[542-05-2]	3-oxopentanedioic acid				
		$\Delta_{\text{sub}}H$	(310–322) 160.2		TPTD	[2005CHA/ZIE]
		Note: Values based on TPTD method are not consistent with values determined by other experimental methods				
C ₅ H ₆ S	[554-14-3]	2-methylthiophene				
		$\Delta_{\text{fus}}H$	9.47	207.8		[1956PEN/FIN]
		Δ_vH	(333–373) 36.8	348	I	[1971EON/POM, 1984BOU/FRI]
		Δ_vH	38.7	298		[1971WIL/ZWO]
		Δ_vH	(324–391) 37.2	339	A,EB	[1987STE/MAL, 1952WHI/BER, 1999DYK/SVO]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₅ H ₆ S	[616-44-4]	3-methylthiophene				
	$\Delta_{\text{fus}}H$		10.54	204.2		[1985DEA]
	Δ_vH	(333–388)	37.3	348		[2009SAP/UUS2]
	Δ_vH	(326–398)	36.8	357		[1999DYK/SVO]
	Δ_vH	(333–373)	37.4	348	I	[1971EON/POM, 1984BOU/FRI]
	Δ_vH	(327–399)	39.5	298		[1971WIL/ZWO]
C ₅ H ₇ ClO ₃	[54166-91-5]	acetic acid, chlorooxo, propyl ester				
	Δ_vH	(282–396)	52.7	297	A	[1987STE/MAL, 1947STU]
C ₅ H ₇ Cl ₃ O ₂	[17831-70-8]	3-chloro-2,2-bis(chloromethyl)propionic acid				
	$\Delta_{\text{fus}}H$		20.9	383.9		[1999GOT/BUH]
C ₅ H ₇ FO ₂	[406-23-5]	allyl fluoroacetate				
	Δ_vH	(273–333)	48.9	288	A,GS	[1987STE/MAL, 1948RED/CHA4]
	Δ_vH					[1972DYK]
C ₅ H ₇ N	[4426-11-3]	cyclobutanecarbonitrile				
	Δ_vH		44.3	298	C	[1983FUC/HAL]
	Δ_vH	(328–402)	39.6	347	BG	[1971HAL/BAL]
	Δ_vH		40.0 ± 0.4	298	BG	[1971HAL/BAL]
C ₅ H ₇ N	[1647-11-6]	2-ethylacrylonitrile				
	Δ_vH	(244–387)	37.1	259	A	[1987STE/MAL, 1947STU]
C ₅ H ₇ N	[20068-02-4]	angelic acid, nitrile				
	Δ_vH	(265–413)	42.8	280	A	[1987STE/MAL, 1947STU]
C ₅ H ₇ N	[96-54-8]	1-methylpyrrole				
	$\Delta_{\text{fus}}H$		7.82	216.9		[1996DOM/HEA]
	$\Delta_{\text{sub}}H$		NA			[1941MIL]
	Δ_vH	(333–373)	38	343	I	[1971EON/POM]
	Δ_vH	(321–423)	39	336	A,EB,IP	[1987STE/MAL, 1968OSB/DOU, 1972DYK]
C ₅ H ₇ N	[25899-50-7]	(Z) 2-pentenitrile				
	Δ_vH		43.2	298		[1969KON/PRO]
C ₅ H ₇ N	[16529-66-1]	(E) 3-pentenitrile				
	Δ_vH		44.8	298		[1969KON/PRO]
C ₅ H ₇ N	[26294-98-4]	(E) 2-pentenitrile				
	Δ_vH		44.9	298		[1969KON/PRO]
C ₅ H ₇ N	[30574-97-1]	tiglic acid, nitrile				
	Δ_vH	(247–395)	37.4	262	A	[1987STE/MAL, 1947STU]
C ₅ H ₇ NO	[927-56-0]	4-oxo-pentanenitrile				
	Δ_vH	(293–473)	52.3	308	A	[1987STE/MAL]
C ₅ H ₇ NO ₂	[105-56-6]	ethyl cyanoacetate				
	$\Delta_{\text{fus}}H$		11.78	246.8		[1996DOM/HEA]
	Δ_vH	(340–479)	66.9	355	A	[1987STE/MAL, 1947STU]
C ₅ H ₇ NO ₂	[1121-89-7]	glutarimide				
	$\Delta_{\text{sub}}H$	(317–340)	93.6 ± 1.6	329	ME	[1990MEN/PIL]
	$\Delta_{\text{sub}}H$		94.1 ± 1.6	298		[1990MEN/PIL]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₅ H ₇ NO ₂	[1121-07-9]	N-methylsuccinimide				
	$\Delta_{\text{sub}}H$	(280–298)	80.6 ± 0.3	289	ME	[1997ROU/JIM]
	$\Delta_{\text{sub}}H$		80.1 ± 0.3	298		[1997ROU/JIM]
C ₅ H ₇ NO ₃	[149-87-1]	(<i>dl</i>)-5-oxoproline				
	$\Delta_{\text{sub}}H$	(394–416)	133.2 ± 1	405	TE,ME	[1979DEK/VOO]
C ₅ H ₇ NS	[3386-97-8]	isothiocyanic acid, 3-butenyl ester				
	Δ_vH	(342–443)	45.2	357	A	[1987STE/MAL, 1999DYK/SVO]
C ₅ H ₇ NS	[541-58-2]	2,4-dimethylthiazole				
	$\Delta_{\text{fus}}H$		2.9	222.9		[1966MEY/MET]
	Δ_vH	(357–421)	42.0	372	A	[1987STE/MAL]
C ₅ H ₇ N ₃ O	[1122-47-0]	1-methylcytosine				
	$\Delta_{\text{sub}}H$	(455–487)	141.2 ± 0.6		GS	[1998ZIE/WSZ]
	$\Delta_{\text{sub}}H$	(423–443)	141.8 ± 8.8	433	ME	[1984BUR/MOR]
	$\Delta_{\text{sub}}H$		149.1 ± 9.0	298		[1984BUR/MOR]
C ₅ H ₇ N ₃ O	[4776-08-3]	3-methylcytosine				
	$\Delta_{\text{sub}}H$	(487–526)	150.6		HAS	[1965CLA/PES]
C ₅ H ₇ N ₃ O	[1122-04-9]	3,5-dimethyl-4-nitrosopyrazole				
	$\Delta_{\text{sub}}H$		102.9 ± 3.0	298	C	[2001RIB/FER]
C ₅ H ₇ N ₃ O ₂	[20541-50-8]	1-methyl-N-hydroxycytosine				
	$\Delta_{\text{sub}}H$		126.7 ± 1.5			[1998ZIE/WSZ]
C ₅ H ₇ N ₃ O ₅	[179894-08-7]	N-acetyl-3,3-dinitroazetidine				
	$\Delta_{\text{fus}}H$		25.65	386.9		[1999GRI/SZE]
C ₅ H ₈	[185-94-4]	bicyclo[2.1.0]pentane				
	Δ_vH		28.0 ± 0.5	298	EB	[1998KOL/PIM, 1996VAR/PAS]
	Δ_vH	(296–315)	28.6	305	A	[1987STE/MAL]
C ₅ H ₈	[157-40-4]	spiropentane				
	$\Delta_{\text{fus}}H$		6.43	166.1		[1996DOM/HEA]
	Δ_vH	(276–344)	28.6	291	A	[1987STE/MAL, 1950SCO/FIN]
	Δ_vH		28.3 ± 0.1	283	C	[1950SCO/FIN2]
	Δ_vH		27.5 ± 0.1	298	C	[1950SCO/FIN2]
	Δ_vH		26.7 ± 0.1	312	C	[1950SCO/FIN2]
C ₅ H ₈	[693-86-7]	vinylcyclopropane				
	Δ_vH	(289–310)	28.9	299	A	[1987STE/MAL]
C ₅ H ₈	[142-29-0]	cyclopentene				
	$\Delta_{\text{trs}}H$		0.48	87.07		
	$\Delta_{\text{fus}}H$		3.36	138.1		[1996DOM/HEA]
	Δ_vH	(249–318)	29.9	264	A	[1987STE/MAL]
	Δ_vH	(289–318)	24.8	299	MM	[1950FOR/CAM]
	Δ_vH	(230–293)	28.4	300		[1941LIS]
C ₅ H ₈	[1120-56-5]	methylenecyclobutane				
	$\Delta_{\text{fus}}H$		5.86	138.5		[1996DOM/HEA]
	Δ_vH	(290–316)	26.1	303	A	[1987STE/MAL]
	Δ_vH	(292–306)	29.1	299	A	[1987STE/MAL, 1978LEB/TSV, 1978LEB/TSV2]
	Δ_vH		27.7 ± 0.4	298	EB	[1974GOO/MOO]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound					
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference	
C ₅ H ₈	[598-25-4]	3-methyl-1,2-butadiene					
	$\Delta_{\text{fus}}H$		7.95	159.5		[1996DOM/HEA]	
	Δ_vH	(227–253)	31	240	A	[1987STE/MAL]	
	Δ_vH	(252–323)	29.9	267	A	[1987STE/MAL]	
	Δ_vH		28	298		[1971WIL/ZWO]	
	Δ_vH	(213–242)	31.6	230	IP	[1969OSB/DOU]	
C ₅ H ₈	[78-79-5]	2-methyl-1,3-butadiene					
	$\Delta_{\text{fus}}H$		4.92	127.3		[1996DOM/HEA]	
	Δ_vH	(221–254)	29.4	239	A	[1987STE/MAL]	
	Δ_vH	(254–316)	28.3	269	A	[1987STE/MAL]	
	Δ_vH		26.4	298		[1971WIL/ZWO]	
	Δ_vH	(216–235)	31.5	225	IP	[1969OSB/DOU]	
C ₅ H ₈	[598-23-2]	3-methyl-1-butyne					
	Δ_vH	(218–320)	30.2	233	A	[1987STE/MAL]	
			25.8	298		[1971WIL/ZWO]	
	C ₅ H ₈	[591-95-7]	1,2-pentadiene				
		$\Delta_{\text{fus}}H$		7.56	135.9		[1996DOM/HEA]
		Δ_vH	(231–249)	31.6	240	A	[1987STE/MAL]
Δ_vH		(249–331)	30.6	264	A	[1987STE/MAL]	
Δ_vH			28.7	298		[1971WIL/ZWO]	
Δ_vH		(213–245)	32.2	231	IP	[1969OSB/DOU]	
C ₅ H ₈	[1574-41-0]	<i>cis</i> 1,3-pentadiene					
	$\Delta_{\text{fus}}H$		5.64	132.4		[1996DOM/HEA]	
	Δ_vH	(255–326)	30.1	270	A	[1987STE/MAL]	
	Δ_vH	(230–255)	31.2	242	A	[1987STE/MAL]	
	Δ_vH		28.3	298		[1971WIL/ZWO]	
	Δ_vH	(213–242)	31.9	230	IP	[1969OSB/DOU]	
C ₅ H ₈	[2004-41-0]	<i>trans</i> 1,3-pentadiene					
	$\Delta_{\text{fus}}H$		7.14	185.7		[1996DOM/HEA]	
	Δ_vH	(228–256)	30.7	242	A	[1987STE/MAL]	
	Δ_vH	(256–324)	29.5	271	A	[1987STE/MAL]	
	Δ_vH		27.8	298		[1971WIL/ZWO]	
	Δ_vH	(213–242)	31.3	230	IP	[1969OSB/DOU]	
C ₅ H ₈	[591-93-5]	1,4-pentadiene					
	$\Delta_{\text{fus}}H$		6.14	124.3		[1996DOM/HEA]	
	Δ_vH	(216–236)	29.1	226	A	[1987STE/MAL]	
	Δ_vH	(236–307)	28.1	251	A	[1987STE/MAL]	
	Δ_vH		25.2	298		[1971WIL/ZWO]	
	Δ_vH	(213–230)	29.3	221	IP	[1969OSB/DOU]	
	(288–300)	26.5	293	MM	[1950FOR/CAM]		
	(194–255)	28.4	240		[1940LAM/ROP]		

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₅ H ₈	[591-96-8]	2,3-pentadiene				
	$\Delta_{\text{fus}}H$		6.13	147.5		[1996DOM/HEA]
	Δ_vH	(234–258)	32.3	246	A	[1987STE/MAL]
	Δ_vH	(258–330)	31.1	273	A	[1987STE/MAL]
	Δ_vH		29.5	298		[1971WIL/ZWO]
	Δ_vH	(213–247)	33.2	232	IP	[1969OSB/DOU]
C ₅ H ₈	[627-19-0]	1-pentyne				
	Δ_vH	(229–315)	31.8	244	A	[1987STE/MAL]
C ₅ H ₈	[627-21-4]	2-pentyne				
	Δ_vH	(240–329)	33.1	255	A	[1987STE/MAL]
C ₅ H ₈ Br ₂	[10230-26-9]	<i>trans</i> 1,2-dibromocyclopentane				
	Δ_vH	(273–332)	47.9	288	A	[1987STE/MAL, 1941LIS]
C ₅ H ₈ Br ₄	[3229-00-3]	penterythritol tetrabromide				
	$\Delta_{\text{fus}}H$		27.97	433.5		[1996DOM/HEA]
	$\Delta_{\text{sub}}H$	(384–434)	84.0	399	A	[1987STE/MAL]
	$\Delta_{\text{sub}}H$		NA		GSM	[1941NIT/SEK]
C ₅ H ₈ ClFO ₂	[541-86-6]	3-chloro-4-fluorobutyric acid, methyl ester				
	Δ_vH	(273–333)	54.5	288	GS	[1987STE/MAL, 1948RED/CHA4, 1972DYK]
C ₅ H ₈ ClF ₃ O	[330-17-0]	2-chloro-1,1,2-trifluoroethyl isopropyl ether				
	Δ_vH		39.2	298	C	[1984MAJ/UCH]
	Δ_vH		38.1	313	C	[1984MAJ/UCH]
C ₅ H ₈ ClF ₃ O	[380-43-8]	2-chloro-1,1,2-trifluoroethyl propyl ether				
	Δ_vH		41.0	298	C	[1984UCH/MAJ]
	Δ_vH		39.9	313	C	[1984UCH/MAJ]
	Δ_vH		38.7	328	C	[1984UCH/MAJ]
	Δ_vH		37.5	343	C	[1984UCH/MAJ]
C ₅ H ₈ Cl ₂ O	[78-71-7]	3,3- <i>bis</i> (chloromethyl)oxetane				
	$\Delta_{\text{fus}}H$		16.95	292.2		[1996DOM/HEA]
C ₅ H ₈ Cl ₄	[2467-10-9]	1,1,1,5-tetrachloropentane				
	Δ_vH	(340–432)	61.7	355	A	[1987STE/MAL]
C ₅ H ₈ F ₂ O ₃	[406-15-5]	<i>bis</i> (2-fluoroethyl) carbonate				
	Δ_vH	(273–333)	61.5	288	GS	[1987STE/MAL, 1948RED/CHA4, 1972DYK]
C ₅ H ₈ F ₄	[338-23-8]	pentaerythritol tetrafluoride				
	$\Delta_{\text{fus}}H$		13.21	249.4		
C ₅ H ₈ F ₄ N ₄ O ₂	[298228-65-6]	4,4- <i>bis</i> (difluoroamino)-1-nitropiperidine				
	$\Delta_{\text{fus}}H$		5.14	367.4		[1996DOM/HEA]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
			50.2	366.2		[2001OXL/SMI]
C ₅ H ₈ NO ₂	[19947-75-2]	5-amino-3,4-dimethylisoxazole				
	$\Delta_{\text{sub}}H$		87.9 ± 2.5			[1973HAM/MIT, 1977PED/RYL]
C ₅ H ₈ N ₂	[80-73-9]	1,3-dimethyl-2-imidazolidinone				
	Δ_vH	(355–498)	54.3	375	EB	[1987KNE/ZON]
	Δ_vH	(355–498)	48.5	450	EB	[1987KNE/ZON]
C ₅ H ₈ N ₂	[7098-07-9]	1-ethylimidazole				
	Δ_vH		66.0 ± 3.9	298	C	[1999RIB/RIB]
C ₅ H ₈ N ₂	[1072-62-4]	2-ethylimidazole				
	$\Delta_{\text{sub}}H$	(303–321)	89.2 ± 0.4	312	ME	[1992JIM/ROU]
	$\Delta_{\text{sub}}H$		89.6 ± 0.4	298	ME	[1992JIM/ROU]
C ₅ H ₈ N ₂	[2817-71-2]	1-ethylpyrazole				
	Δ_vH		53.3 ± 2.4	298	C	[1999RIB/RIB]
C ₅ H ₈ N ₂	[2721-32-6]	2,3-diazabicyclo[2.2.1]hept-2-ene				
	$\Delta_{\text{sub}}H$		43.9 ± 2.1			[1974ENG/WOO, 1977PED/RYL]
	$\Delta_{\text{sub}}H$		55.3 ± 0.6	298		[1976ENG/MEL]
C ₅ H ₈ N ₂	[67-51-6]	3,5-dimethylpyrazole				
	$\Delta_{\text{sub}}H$		83.4 ± 2.4	298	C	[2001RIB/FER]
	$\Delta_{\text{sub}}H$		83.3 ± 0.2	301	ME	[1991ELG/YRA]
C ₅ H ₈ N ₂ O ₂	[19947-75-2]	1,3-dimethyluracil				
	$\Delta_{\text{sub}}H$		96.9 ± 1.2		C	[1989IMA/TAK]
C ₅ H ₈ N ₄ O ₆	[298228-66-7]	1,4,4-trinitropiperidine				
	$\Delta_{\text{fus}}H$		100.4	389.2		[2001OXL/SMI]
C ₅ H ₈ N ₄ O ₁₂	[78-11-5]	pentaerythritol tetranitrate				
	$\Delta_{\text{sub}}H$	(356–382)	156.9 ± 0.8	369	TE	[2004LAU/HIL]
	$\Delta_{\text{sub}}H$		152.3		DSC	[1990HWA/YOS]
	$\Delta_{\text{sub}}H$	(328–405)	150.4 ± 1.3	298	ME	[1978CUN/PAL]
	$\Delta_{\text{sub}}H$		146 ± 12			[1978CUN/PAL, 1971DIN/STA]
	$\Delta_{\text{sub}}H$	U	121.3		ME	[1969CRI]
	$\Delta_{\text{sub}}H$	(370–411)	151.9 ± 2.1			[1953EDW, 1960JON, 1970COX/PIL]
C ₅ H ₈ O	[120-93-2]	cyclopentanone				
	$\Delta_{\text{fus}}H$		11.4	221.2		[1998GON/SZW]
	Δ_vH	(323–403)	41.5	338	EB	[2006TEO/BAR]
	Δ_vH		42.1 ± 0.2	298		[1991DIK/KAB]
	Δ_vH		43.2 ± 0.3		GC	[1989AZA]
	Δ_vH	(317–427)	40.6	332		[1987AMB/GHI2]
	Δ_vH	(293–404)	42.6	308	A	[1987STE/MAL]
	Δ_vH	(338–416)	39.6	353	A,EB	[1987STE/MAL, 1976MEY/HOT]
	Δ_vH		42.6 ± 0.4	298		[1972WOL]
	Δ_vH		42.7 ± 0.1	298	C	[1968PLA/WIL]
	Δ_vH	(273–299)	43.6	286		[191942BEN/KIS]
C ₅ H ₈ O	[922-63-4]	2-ethylacrolein				
	Δ_vH		36.8 ± 0.4	298	C	[1996VAN/YU]
C ₅ H ₈ O	[765-43-5]	cyclopropyl methyl ketone				
	Δ_vH	(361–387)	37.6	374	A	[1987STE/MAL]
	Δ_vH		39.4			[1984KOZ/TIM]
	Δ_vH		39.4 ± 0.1	298	C	[1983FUC/HAL]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₅ H ₈ O	[25512-65-6] $\Delta_v H$	dihydro-2H-pyran (273–288)	32.2	280	A	[1987STE/MAL, 1972DYK, 1958CAS/FLE3]
C ₅ H ₈ O	[497-03-0] $\Delta_v H$	<i>trans</i> 2-methyl-2-butenal (248–390)	39.2	263	A	[1987STE/MAL, 1947STU]
C ₅ H ₈ O	[814-78-8] $\Delta_v H$	3-methyl-3-buten-2-one (313–371)	26.2	328	A	[1987STE/MAL, 1972DYK]
C ₅ H ₈ O	[115-19-5] $\Delta_v H$ $\Delta_v H$ $\Delta_v H$	2-methyl-3-buten-2-ol (333–377) (294–380) (294–380)	41.0	353	A	[1999ZAR/CHA]
			43.9	337	A	[1987STE/MAL, 1972DYK]
			49.5	309		[1984BOU/FRI, 1950CON/ELV]
C ₅ H ₈ O	[1629-58-9] $\Delta_v H$	1-penten-3-one (303–376)	36.7	318	A	[1987STE/MAL]
C ₅ H ₈ OS	[1072-72-6] $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$	tetrahydro-4H-thiopyran-4-one	71.7 ± 1.7	317	I	[1972GEI/SAW]
			72.6 ± 1.7	298		[1972GEI/SAW, 1977PED/RYL]
C ₅ H ₈ O ₂	[111-30-8] $\Delta_v H$ $\Delta_v H$	glutaraldehyde (347–382) (327–436)	51.4	362		[1998OLS]
			56.2	342		[1998OLS]
C ₅ H ₈ O ₂	[2868-37-3] $\Delta_v H$ $\Delta_v H$	methyl cyclopropanecarboxylate (273–313)	42.6 ± 0.4		GS	[1998VER/KUM]
			41.3 ± 0.1	298	C	[1983FUC/HAL]
C ₅ H ₈ O ₂	[123-54-6] $\Delta_v H$ (diketone) $\Delta_v H$ (enol form) $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ (84% enol) $\Delta_v H$ (100% enol) $\Delta_v H$	acetylacetone (307–414) (295–313) (378–411) (288–378) (297–398)	51.2 ± 2.2	298	CGC	[2005TEM/ROU]
			50.8 ± 0.6	298	CGC	[2005TEM/ROU]
			39.2	322	EB	[1985RAV/RAO]
			40.6	304		[1981INO/ARA]
			35.2	393	A,I,EB	[1987STE/MAL, 1972NAK/TOY]
			42.7	303	A,EB	[1987STE/MAL]
			41.8 ± 0.2	298	C	[1970IRV/WAD]
			43.2	298	C	[1970IRV/WAD]
39.4	347		[1969MEL/MER]			
C ₅ H ₈ O ₂	[na] $\Delta_{\text{fus}} H$	acetylacetone enol	14.5	254.8		[1969MEL/MER]
C ₅ H ₈ O ₂	[600-14-6] $\Delta_{\text{fus}} H$	2,3-pentanedione	7.84	221.2	DSC	[2006DOM/MOR]
C ₅ H ₈ O ₂	[565-63-9] $\Delta_v H$	<i>cis</i> 2-methyl-2-butenic acid (361–458)	61.8	376	A	[1987STE/MAL]
C ₅ H ₈ O ₂	[3586-58-1] $\Delta_v H$	2-ethylpropenoic acid	52.1 ± 0.4	298	C	[1996VAN/YU]
C ₅ H ₈ O ₂	[3586-58-1] $\Delta_v H$	2-ethylacrylic acid (320–453)	62.2	335	A	[1987STE/MAL, 1947STU]
C ₅ H ₈ O ₂	[626-96-0] $\Delta_v H$	4-oxovaleraldehyde (levulinaldehyde) (301–460)	48.8	316	A	[1987STE/MAL, 1947STU]
C ₅ H ₈ O ₂	[541-47-9]	3-methylcrotonic acid				

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound					
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference	
		$\Delta_v H$	(363–473)	57.7	378	A	[1987STE/MAL]
C ₅ H ₈ O ₂	[140-88-5]	2-ethyl acrylate					
	$\Delta_v H$	(243–372)	41.4	258		[1947STU]	
C ₅ H ₈ O ₂	[140-88-5]	2-propenoic acid, ethyl ester					
	$\Delta_v H$		39.2			[1975VIL/PER]	
C ₅ H ₈ O ₂	[80-62-6]	methyl methacrylate					
	$\Delta_{\text{fus}} H$		12.24	225		[1996DOM/HEA]	
	$\Delta_{\text{sub}} H$	(194–223)	60.7	205		[1952BYW, 1960JON]	
	$\Delta_v H$	(295–386)	38.8 ± 0.1	300	EB	[2002STE/CHI4]	
	$\Delta_v H$	(295–386)	36.3 ± 0.2	340	EB	[2002STE/CHI4]	
	$\Delta_v H$	(295–386)	33.3 ± 0.4	380	EB	[2002STE/CHI4]	
	$\Delta_v H$	(293–373)	37.9	308	A	[1987STE/MAL]	
	$\Delta_v H$	(318–348)	37.7	333		[1984HUL/LU]	
	$\Delta_v H$	(305–373)	38	320		[1984BOU/FRI]	
	$\Delta_v H$		40.1			[1975VIL/PER]	
	$\Delta_v H$	(312–362)	39	327		[1956VON/JEN]	
C ₅ H ₈ O ₂	[591-80-0]	4-pentenoic acid					
	$\Delta_v H$	(289–324)	65.8 ± 0.4	298	GS	[2008EME/VER]	
C ₅ H ₈ O ₂	[80-59-1]	<i>trans</i> 2-methyl-2-butenoic acid					
	$\Delta_v H$	(350–453)	61.2	365	A	[1987STE/MAL]	
C ₅ H ₈ O ₂	[542-28-9]	tetrahydro-2 <i>H</i> -pyran-2-one (δ -valerolactone)					
	$\Delta_{\text{trs}} H$		0.46	118			
	$\Delta_{\text{trs}} H$		0.3	135			
	$\Delta_{\text{fus}} H$		10.53	263		[1991ACR]	
	$\Delta_v H$	(278–353)	58.2 ± 0.3	298	GS	[2007EME/KOZ]	
	$\Delta_v H$	(393–428)	52.4 ± 0.2	410	EB	[1991WIB/WAL]	
	$\Delta_v H$	(393–428)	60.2 ± 1.3	298	EB	[1991WIB/WAL]	
	$\Delta_v H$		58.0 ± 0.4	298	C	[1990LEI/PIL2, 1989BRO/CON]	
	$\Delta_v H$	(342–433)	48.6	387		[1930SCH/THO]	
C ₅ H ₈ O ₂	[29943-42-8]	2,3,5,6-tetrahydropyran-4-one					
	$\Delta_v H$		50.7 ± 0.3	298	C	[2009FRE/GOM2]	
C ₅ H ₈ O ₂	[108-29-2]	(<i>dl</i>) γ -valerolactone					
	$\Delta_v H$	(276–350)	53.9 ± 0.2	298	GS	[2008EME/KOZ, 2009EME/VER]	
	$\Delta_v H$		54.8 ± 0.4	298	C	[1990LEI/PIL2]	
	$\Delta_v H$	(310–480)	53.5	325	A	[1987STE/MAL, 1947STU]	
C ₅ H ₈ O ₂	[105-38-4]	vinyl propanoate					
	$\Delta_v H$	(321–368)	36.7	336		[2005RES/GON]	
C ₅ H ₈ O ₂ S	[6007-71-2]	2,5-dihydro-2-methyl-thiophene-1,1-dioxide					
	$\Delta_{\text{sub}} H$		60.7 ± 2.5			[1969MAC/MCN, 1969MAC/MCN2, 1977PED/RYL]	
C ₅ H ₈ O ₂ S	[1193-10-8]	2,5-dihydro-3-methyl-thiophene-1,1-dioxide					
	$\Delta_{\text{sub}} H$		64.0 ± 2.5			[1969MAC/MCN, 1969MAC/MCN2, 1977PED/RYL]	
C ₅ H ₈ O ₃	[123-76-2]	4-oxopentanoic acid					
	$\Delta_v H$	(375–519)	74.4	390	A	[1987STE/MAL, 1947STU]	

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₅ H ₈ O ₃	[105-45-3]	methyl acetoacetate				
	$\Delta_v H$	(289–446)	45.4	304	A	[1987STE/MAL]
C ₅ H ₈ O ₃	[123-76-2]	levulinic acid				
	$\Delta_{\text{fus}} H$		9.22	306.2		[1991ACR]
C ₅ H ₈ O ₃	[4437-85-8]	butylene carbonate				
	$\Delta_v H$	(289–344)	63.2 ± 0.3	298	GS	[2008VER/TOK]
	$\Delta_v H$	(397–523)	63.8 ± 0.1	298	E	[2004CHE/CLE, 2008VER/TOK]
C ₅ H ₈ O ₄	[108-59-8]	dimethyl malonate				
	$\Delta_{\text{sub}} H$		111.7 ± 2.1	298	ME	[2000RIB/MON]
	$\Delta_v H$	(278–314)	57.5 ± 0.3	298	GS	[2006VER/KOZ]
	$\Delta_v H$	(351–460)	52.9 ± 0.2	360	EB	[2002STE/CHI6]
	$\Delta_v H$	(351–460)	49.5 ± 0.2	400	EB	[2002STE/CHI6]
	$\Delta_v H$	(351–460)	46.1 ± 0.3	440	EB	[2002STE/CHI6]
	$\Delta_v H$	(278–308)	61.8 ± 0.8	293	GS	[1992VER/BEC]
	$\Delta_v H$	(374–620)	50.0	497	EB,HG	[1988ASK/DAU]
	$\Delta_v H$	(308–454)	53.7	323	A	[1987STE/MAL]
C ₅ H ₈ O ₄	[110-94-1]	glutaric acid				
	$\Delta_{\text{trs}} H$		2.34	349.2		
	$\Delta_{\text{fus}} H$		21.3	372.3	DSC	[2009HA/HAN]
	$\Delta_{\text{fus}} H$		20.7	370.9	DSC	[2009GOO/ROD]
	$\Delta_{\text{trs}} H$		2.3	340.5		
	$\Delta_{\text{fus}} H$		18.8	363.9	DSC	[2005ROU/TEM]
	$\Delta_{\text{trs}} H$		2.4	338		
	$\Delta_{\text{fus}} H$		23	371		[2002STE/CHI6]
	$\Delta_{\text{trs}} H$		2.46	348.5		
	$\Delta_{\text{fus}} H$		20.9	371		[1991ACR]
	$\Delta_{\text{sub}} H$	(313–349)	134 ± 4		TPD	[2007CAP/LOV]
	$\Delta_{\text{sub}} H$	(275–294)	132.3		TPTD	[2005CHA/ZIE]
	Note: Values based on TPTD method are not consistent with values determined by other experimental methods					
$\Delta_{\text{sub}} H$	(348–363)	117.0 ± 1.2	356	ME	[1999RIB/MON]	
$\Delta_{\text{sub}} H$	(348–363)	119.8 ± 1.2	298	ME	[1999RIB/MON]	
$\Delta_{\text{sub}} H$	(292–320)	U 52.6	306	A	[1947GRA]	
$\Delta_v H$	(424–503)	101.6	298	GS	[2005ROU/TEM]	
$\Delta_v H$	(428–576)	98.1	443	A	[1987STE/MAL, 1947STU]	
C ₅ H ₈ O ₄	[601-75-2]	ethylmalonic acid				
	$\Delta_{\text{sub}} H$		112.8 ± 2.2	298	ME	[2000RIB/MON]
			105.5 ± 0.5		C	[1983ALT/PIL]
C ₅ H ₈ O ₄	[628-51-3]	diacetoxymethane				
	$\Delta_v H$	(334–443)	50.6	349	A	[1987STE/MAL]
C ₅ H ₉ BrO	[815-48-5]	3-bromo-2-pentanone				
	$\Delta_v H$	(273–333)	45.2	288	A	[1987STE/MAL, 1972DYK]
C ₅ H ₉ Cl	[930-28-9]	cyclopentyl chloride				
	$\Delta_{\text{trs}} H$		7.63	169.4		
	$\Delta_{\text{fus}} H$		0.64	180		[1993DIK/KAB]
	$\Delta_v H$		38.8	298	C	[1993DIK/KAB]
		(322–387)	37.4	337	A,EB	[1987STE/MAL, 1970AND/BRA]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₅ H ₉ ClO ₂	[5396-24-7]	propyl chloroacetate				
	$\Delta_{\text{trs}}H$		0.36	110		
	$\Delta_{\text{trs}}H$		0.69	204		
	$\Delta_{\text{fus}}H$		13.0	240		[1990MID/KAT]
C ₅ H ₉ ClO ₂	[105-48-6]	isopropyl chloroacetate				
	Δ_vH	(308–425)	44.3	323		[1928NEL2, 1984BOU/FRI]
C ₅ H ₉ ClO ₂	[535-13-7]	2-chloropropionic acid, ethyl ester				
	Δ_vH	(279–420)	46.5	294	A	[1987STE/MAL, 1947STU]
C ₅ H ₉ ClO ₂	[623-71-2]	3-chloropropionic acid, ethyl ester				
	Δ_vH	(316–358)	56.0	331	A	[1987STE/MAL]
C ₅ H ₉ ClS	[19155-35-2]	(2-chloroethyl) allyl sulfide				
	Δ_vH	(293–333)	50.2	308	A,GS	[1987STE/MAL, 1949WAD/SMI, 1972DYK, 1999DYK/SVO]
C ₅ H ₉ Cl ₃ O	[1067-09-0]	2-chloromethyl-2-methyl-1,3-dichloropropane				
	$\Delta_{\text{trs}}H$		12	246.6		
	$\Delta_{\text{fus}}H$		2.5	291.3		[1996DOU/FUE]
C ₅ H ₉ Cl ₃ O	[813-99-0]	3-chloro-2,2-bis(chloromethyl)-1-propanol				
	Δ_vH	(404–450)	79.6	419	A	[1987STE/MAL]
C ₅ H ₉ FOS	[63732-24-1]	4-fluorothiobutyric acid, methyl ester				
	Δ_vH	(273–333)	52.4	288	A,GS	[1987STE/MAL, 1948RED/CHA4, 1972DYK, 1999DYK/SVO]
C ₅ H ₉ FO ₂	[406-20-2]	4-fluorobutyric acid, methyl ester				
	Δ_vH	(273–333)	47.3	288	A,GS	[1987STE/MAL, 1948RED/CHA4, 1972DYK]
C ₅ H ₉ FO ₂	[406-06-4]	isopropyl fluoroacetate				
	Δ_vH	(273–333)	44.3	288	A,GS	[1987STE/MAL, 1948RED/CHA4]
C ₅ H ₉ FO ₃	[25309-12-0]	3-fluoro-2-hydroxybutyric acid, methyl ester				
	Δ_vH	(273–333)	62.3	288	GS	[1987STE/MAL, 1948RED/CHA4, 1972DYK]
C ₅ H ₉ N	[630-18-2]	pivalonitrile				
	$\Delta_{\text{trs}}H$		0.23	213		
	$\Delta_{\text{trs}}H$		1.91	232.7		
	$\Delta_{\text{fus}}H$		9.29	292.1		[1996DOM/HEA]
	Δ_vH	(299–365)	37.0	318	BG	[1971HAL/BAL]
	Δ_vH	(313–371)	36.5	328	A,I	[1987STE/MAL, 1967WES/RIB]
C ₅ H ₉ N	[18936-17-9]	2-methylbutyronitrile				
	Δ_vH	(274–313)	42.5 ± 0.3		GS	[1994RAK/VER]
	Δ_vH	(313–418)	42.3	328	A	[1987STE/MAL]
	Δ_vH	(342–414)	44.2	298	EB	[1969KON/PRO]
C ₅ H ₉ N			44.2			[1949DRE/SHR, 1949DRE/MAR, 2005EME/VER]
	Δ_vH	(394–439)	44.1 ± 0.2	298	MM	[1933HEL, 2005EME/VER]
C ₅ H ₉ NO	[872-50-4]	N-methyl-2-pyrrolidone				

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{fus}}H$		18.1	248.5		[2000LIS/JAM]
	Δ_vH	(380–475)	49.5	395		[2007PAL/ORAZ]
	Δ_vH	(352–378)	61.9	298	EB	[2004CHY/FRA2]
	Δ_vH	(330–373)	53.1	345	GS	[1996LIN/WIC]
	Δ_vH	(340–476)	53.4	350	EB	[1987KNE/ZON]
	Δ_vH	(340–476)	47.7	425	EB	[1987KNE/ZON]
	Δ_vH	(361–477)	49.2	376	A	[1987STE/MAL, 1972DYK]
	Δ_vH	(291–299)	55.3	295	A	[1987STE/MAL]
	Δ_vH	(333–473)	49.3	403		[1979BLU/BAE]
C ₅ H ₉ NO	[111-36-4]	butyl isocyanate				
	Δ_vH	(293–388)	38.5	308		[2004AHM/GIE]
	Δ_vH	(273–389)	46.8	288	A	[1987STE/MAL, 1974ZHU/MON]
C ₅ H ₉ NO	[1873-29-6]	isobutyl isocyanate				
	Δ_vH	(273–376)	44.2	288	A	[1987STE/MAL, 1974ZHU/MON]
C ₅ H ₉ NO	[3887-02-3]	N-methyl methacrylamide				
	Δ_vH	(355–489)	60.9	370	A	[1987STE/MAL]
C ₅ H ₉ NO	[15856-96-9]	<i>cis</i> 2-pentenoic acid amide				
	$\Delta_{\text{sub}}H$	(323–333)	106.5	328	A	[1987STE/MAL]
	Δ_vH	(343–384)	74.8	358	A	[1987STE/MAL]
C ₅ H ₉ NO	[15856-96-9]	<i>trans</i> 2-pentenoic acid amide				
	$\Delta_{\text{sub}}H$	(353–383)	57.9	368	A	[1987STE/MAL]
C ₅ H ₉ NO	[76474-09-4]	α -methoxyisobutyronitrile				
	Δ_vH	(261–285)	37.4 ± 0.8	298	GS	[1995VER/BEC]
C ₅ H ₉ NO	[14631-45-9]	2-ethoxypropanenitrile				
	Δ_vH	(348–445)	46.7	363	A,EB	[1987STE/MAL, 1976RAO/CHI]
C ₅ H ₉ NO	[675-20-7]	2-piperidone				
	$\Delta_{\text{fus}}H$		10.5	311.9		[1990DOM/HEA]
	Δ_vH	(293–312)	75.5	302	A	[1987STE/MAL]
C ₅ H ₉ NO	[10431-98-8]	2-ethyl-2-oxazoline				
	Δ_vH		44.2 ± 0.4	298	C	[1976HAM/THO]
	$\Delta_{\text{sub}}H$	(293–312)	74.5	303		[1953AIH, 1960JON, 1960AIH2]
C ₅ H ₉ NO ₂	[147-85-3]	L-(<i>l</i>)-proline				
	$\Delta_{\text{sub}}H$	(396–416)	127.4 ± 1	406	TE,ME	[1979DEK/VOO]
	$\Delta_{\text{sub}}H$	(380–420)	149 ± 4	400	C	[1978SAB/LAF]
	$\Delta_{\text{sub}}H$	(323–423)	U 50 ± 8	373	LE	[1977GAF/PIE]
C ₅ H ₉ NO ₂	[4394-85-8]	N-formylmorpholine				
	Δ_vH		56.9	416		[1989PAR/GME]
	Δ_vH	(375–423)	52.7	399	TGA	[1987ALN/ALS]
C ₅ H ₉ NO ₃	[51-35-4]	<i>trans</i> 4-hydroxy-L-proline				
	$\Delta_{\text{sub}}H$	(461–481)	162.6 ± 2	471	TE,ME	[1979DEK/VOO]
C ₅ H ₉ NO ₄	[56-86-0]	(<i>l</i>)-glutamic acid				
	$\Delta_{\text{sub}}H$	(353–453)	U 121 ± 34	403	LE	[1977GAF/PIE]
C ₅ H ₉ N ₃ O ₇	[26459-85-8]	2-ethoxy-1,1,1-trinitropropane				
	Δ_vH	(293–310)	57.7	301	A	[1987STE/MAL]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₅ H ₉ N ₃ O ₉	[3032-55-1]	2-hydroxymethyl-2-methyl-1,3-propanediol trinitrate				
	$\Delta_v H$	(299–345)	88.1	314	A	[1987STE/MAL]
C ₅ H ₉ N ₃ O ₉	[98071-55-7]	1,2,5-pentanetriol trinitrate				
	$\Delta_v H$	(293–313)	41.7 ± 2.1	303	A,GS	[1987STE/MAL, 1957KEM/GOL]
C ₅ H ₁₀	[1630-94-0]	1,1-dimethylcyclopropane				
	$\Delta_v H$		25.1 ± 0.8	298	EB	[1974GOO/MOO]
C ₅ H ₁₀	[287-92-3]	cyclopentane				
	$\Delta_{\text{trs}}H$		4.9	122		
	$\Delta_{\text{trs}}H$		0.34	138		
	$\Delta_{\text{fus}}H$		0.6	179.7		[1996DOM/HEA]
	$\Delta_{\text{sub}}H$		42.6	122	B	[1963BON]
	$\Delta_v H$	(280–331)	29.2	295	A	[1987STE/MAL]
	$\Delta_v H$	(322–384)	28.0	337	A	[1987STE/MAL]
	$\Delta_v H$	(381–455)	27.2	396	A	[1987STE/MAL]
	$\Delta_v H$	(452–511)	27.5	467	A	[1987STE/MAL]
	$\Delta_v H$		28.5	298		[1971WIL/ZWO]
	$\Delta_v H$		28.5 ± 0.1	298	C	[1959MCC/PEN]
	$\Delta_v H$		27.9 ± 0.1	310	C	[1959MCC/PEN]
	$\Delta_v H$		27.3 ± 0.1	322	C	[1959MCC/PEN]
	$\Delta_v H$		27.4	323		[1946SPI/PIT]
	$\Delta_v H$	(289–323)	29.0	304	MM	[1945WIL/TAY]
$\Delta_v H$		29.2	298	C	[1943AST/FIN]	
C ₅ H ₁₀	[109-67-1]	1-pentene				
	$\Delta_{\text{fus}}H$		5.81	107.9		[1991ACR, 1990MES/TOD]
	$\Delta_v H$	(218–311)	29.1	233	A	[1987STE/MAL]
	$\Delta_v H$	(286–304)	26.7	295	MM	[1950FOR/CAM]
	$\Delta_v H$		25.5	298		[1971WIL/ZWO]
	$\Delta_v H$	(273–334)	26.9	288		[1949SCO/WAD]
	$\Delta_v H$		26.2 ± 0.1	284	C	[1949SCO/WAD]
	$\Delta_v H$		25.5 ± 0.1	298	C	[1949SCO/WAD]
	$\Delta_v H$		25.2 ± 0.1	303	C	[1949SCO/WAD]
	$\Delta_v H$	(273–308)	26.3	290		[1948DAY/NIC]
$\Delta_v H$	(313–368)	25.7	341		[1948DAY/NIC]	
C ₅ H ₁₀	[627-20-3]	<i>cis</i> 2-pentene				
	$\Delta_{\text{fus}}H$		7.11	121.8		[1991ACR]
	$\Delta_v H$	(234–318)	29.8	249	A	[1987STE/MAL]
	$\Delta_v H$		26.8	298		[1971WIL/ZWO]
$\Delta_v H$	(274–341)	28.1	289	EB	[1950SCO/WAD]	
C ₅ H ₁₀	[646-04-8]	<i>trans</i> 2-pentene				
	$\Delta_{\text{fus}}H$		8.35	133		[1991ACR]
	$\Delta_v H$	(251–341)	28.8	266	A	[1987STE/MAL]
	$\Delta_v H$		26.7	298		[1971WIL/ZWO]
$\Delta_v H$	(274–341)	28	289	EB	[1950SCO/WAD]	
C ₅ H ₁₀	[563-46-2]	2-methyl-1-butene				
	$\Delta_{\text{fus}}H$		5.36	104.7		[1996DOM/HEA]
	$\Delta_v H$	(240–336)	28.5	255	A	[1987STE/MAL]
$\Delta_v H$		25.9	298		[1971WIL/ZWO]	

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound					
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference	
		$\Delta_v H$	(274–336)	27.3	289		[1949SCO/WAD]
		$\Delta_v H$		25.9 ± 0.1	298	C	[1949SCO/WAD]
		$\Delta_v H$		25.5 ± 0.1	304	C	[1949SCO/WAD]
C ₅ H ₁₀	[563-45-1]	3-methyl-1-butene					
		$\Delta_{\text{fus}} H$		5.36	104.7		[1996DOM/HEA]
		$\Delta_v H$	(237–324)	26.3	252	A	[1987STE/MAL]
		$\Delta_v H$		23.9	298		[1971WIL/ZWO]
		$\Delta_v H$	(273–324)	25.4	288	EB	[1950SCO/WAD]
C ₅ H ₁₀	[515-35-9]	2-methyl-2-butene					
		$\Delta_{\text{fus}} H$		7.59	139.4		[1996DOM/HEA]
		$\Delta_v H$	(271–343)	28.4	286	A	[1987STE/MAL]
		$\Delta_v H$		27.1	298		[1971WIL/ZWO]
		$\Delta_v H$	(276–344)	28.3	291		[1949SCO/WAD]
		$\Delta_v H$		27.5 ± 0.1	290	C	[1949SCO/WAD]
		$\Delta_v H$		27.1 ± 0.1	298	C	[1949SCO/WAD]
	$\Delta_v H$		26.3 ± 0.1	312	C	[1949SCO/WAD]	
C ₅ H ₁₀	[698-61-8]	methylcyclobutane					
		$\Delta_{\text{fus}} H$		5.76	138.6		[1991ACR]
C ₅ H ₁₀ Br ₂	[13320-56-4]	1,1-dibromopentane					
		$\Delta_v H$	(360–501)	48.8	375	A,EST	[1987STE/MAL, 1956MAN, 1972DYK]
C ₅ H ₁₀ Br ₂	[3234-49-9]	1,2-dibromopentane					
		$\Delta_v H$	(348–465)	46.5	363	A	[1987STE/MAL]
		$\Delta_v H$	(350–450)	49	298		[1975PIS/ROZ, 1991BAS/SVO]
		$\Delta_v H$		49.2 ± 0.8	298	EB	[1975PIS/ROZ]
		$\Delta_v H$	(292–448)	48.8	307	A	[1987STE/MAL, 1947STU]
C ₅ H ₁₀ Br ₂	[626-87-9]	1,4-dibromopentane					
		$\Delta_v H$	(377–524)	51.8	392	A	[1987STE/MAL, 1972DYK]
C ₅ H ₁₀ Br ₂	[111-24-0]	1,5-dibromopentane					
		$\Delta_v H$	(396–549)	54.4	411	A	[1987STE/MAL, 1972DYK]
C ₅ H ₁₀ Br ₂ O ₂	[3296-90-0]	2,2-bis(bromomethyl)-1,3-propanediol					
		$\Delta_{\text{fus}} H$		30.1	387.3		[1996DOU/FUE]
C ₅ H ₁₀ ClNO	[2895-21-8]	2-chloro-N-isopropylacetamide					
		$\Delta_{\text{fus}} H$		26.05	351.3	DSC	[1990DON/DRE]
C ₅ H ₁₀ Cl ₂	[820-55-3]	1,1-dichloropentane					
		$\Delta_v H$	(340–410)	44.3	298	A	[1987VAR/LOS2, 1991BAS/SVO]
		$\Delta_v H$	(325–457)	42.0	340	A,EST	[1987STE/MAL, 1956MAN, 1972DYK]
C ₅ H ₁₀ Cl ₂	[1674-33-5]	1,2-dichloropentane					
		$\Delta_v H$	(330–420)	44.4	298		[1991BAS/SVO]
		$\Delta_v H$	(332–418)	41.9	347	A	[1987STE/MAL]
		$\Delta_v H$		43.8 ± 0.7	298	EB	[1975PIS/ROZ2]
C ₅ H ₁₀ Cl ₂	[626-92-6]	1,4-dichloropentane					
		$\Delta_v H$	(350–440)	48.9	298		[1991BAS/SVO]
		$\Delta_v H$	(348–443)	45.0	363	A	[1987STE/MAL]
		$\Delta_v H$		48.1 ± 0.8	298	EB	[1975PIS/ROZ2]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		T _m (K)	Method	Reference
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)			
C ₅ H ₁₀ Cl ₂	[628-76-2]	1,5-dichloropentane				
	$\Delta_v H$	(360–450)	52.2	298		[1991BAS/SVO]
	$\Delta_v H$	(362–453)	47.2	377	A	[1987STE/MAL]
	$\Delta_v H$		51.3 ± 0.8	298	EB	[1975PIS/ROZ2]
C ₅ H ₁₀ Cl ₂	[29559-55-5]	1,3-dichloro-2,2-dimethylpropane				
	$\Delta_{\text{trs}} H$		0.6	193.8		
	$\Delta_{\text{trs}} H$		3.6	198.4		
	$\Delta_{\text{fus}} H$		1.6	262.2		[99GOT/BUH]
C ₅ H ₁₀ Cl ₂ O	[52250-75-6]	(2-chloroethyl)-(2-chloroisopropyl) ether				
	$\Delta_v H$	(297–453)	49.7	312	A	[1987STE/MAL, 1947STU]
C ₅ H ₁₀ Cl ₂ O	[42434-29-7]	(2-chloroethyl)-(2-chloropropyl) ether				
	$\Delta_v H$	(302–467)	49.3	317	A	[1987STE/MAL, 1947STU]
C ₅ H ₁₀ Cl ₂ O ₂	[111-91-1]	<i>bis</i> (2-chloroethoxy) methane				
	$\Delta_v H$	(326–488)	54.2	341	A	[1987STE/MAL, 1947STU]
C ₅ H ₁₀ F ₂	[62127-40-6]	1,1-difluoropentane				
	$\Delta_v H$	(268–378)	34.4	283	A,EST	[1987STE/MAL, 1956MAN, 1972DYK]
C ₅ H ₁₀ F ₂	[371-65-3]	2,2-difluoropentane				
	$\Delta_v H$	(262–367)	33.7	277	A	[1987STE/MAL, 1972DYK]
C ₅ H ₁₀ F ₂	[358-03-2]	3,3-difluoropentane				
	$\Delta_v H$	(262–368)	33.8	277	A	[1987STE/MAL, 1972DYK]
C ₅ H ₁₀ F ₂ O ₂	[373-40-0]	<i>bis</i> (2-fluoroethoxy) methane				
	$\Delta_v H$	(273–333)	52.3	288	A,GS	[1987STE/MAL, 1948RED/CHA4, 1972DYK]
C ₅ H ₁₀ N ₂	[1738-25-6]	3-(dimethylamino)propionitrile				
	$\Delta_v H$		42.2 ± 0.1			[1992PAP/PIM]
	$\Delta_v H$	(330–445)	45.9	345	A	[1987STE/MAL]
	$\Delta_v H$	(331–407)	52.4	346	A	[1987STE/MAL]
	$\Delta_v H$	(290–317)	44.1 ± 0.2			[1984LEB/GUT2]
	$\Delta_v H$		47.3			[1977VAS/KOT]
C ₅ H ₁₀ N ₂ O	[100-75-4]	1-nitrosopiperidine				
	$\Delta_v H$	(333–383)	47.7	348	A	[1987STE/MAL]
C ₅ H ₁₀ N ₂ O ₂	[7606-79-3]	N-acetyl glycine, N-methylamide				
	$\Delta_{\text{sub}} H$	(348–363)	97.8	355.5	A	[1987STE/MAL, 1955AIH]
C ₅ H ₁₀ N ₂ O ₂	[15962-47-7]	N-acetyl-L-alanine amide				
	$\Delta_{\text{fus}} H$		21.7	431		[1996DOM/HEA]
	$\Delta_{\text{sub}} H$		115.0 ± 1.2	376	C	[1999DEL/BAR]
	$\Delta_{\text{sub}} H$		118.1 ± 1.6	298		[1999DEL/BAR]
	$\Delta_{\text{sub}} H$	(366–410)	115 ± 3	388	TE	[1988FER/DEL, 1986BAR/FER]
C ₅ H ₁₀ N ₂ O ₂	[95048-77-4]	N-acetylsarcosinamide				
	$\Delta_{\text{fus}} H$		27.4	412.7		[1997PUL/DES]
C ₅ H ₁₀ N ₂ O ₂	[3424-60-0]	glutaramide				
	$\Delta_{\text{fus}} H$		38.4	453.9	DSC	[2006BAD/DEL]
C ₅ H ₁₀ N ₂ O ₂ S	[16752-77-5]	5-methyl N-(methylcarbamoyloxy)thioacetimidate				
	$\Delta_{\text{fus}} H$		21.73	352.7	DSC	[1990DON/DRE]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₅ H ₁₀ N ₂ O ₃	[na] $\Delta_{\text{fus}}H$ (decomp)	alanylglycine	56.6	508		[1996DOM/HEA]
C ₅ H ₁₀ N ₂ O ₆	[3457-92-9] Δ_vH	1,5-pentanediol dinitrate (293–313)	78.9 ± 5.9	303	A,GS	[1987STE/MAL, 1957KEM/GOL, 1972DYK]
C ₅ H ₁₀ N ₂ O ₆	[25385-63-1] Δ_vH	2,4-pentanediol dinitrate (293–313)	60.6 ± 5.9	303	A,GS	[1987STE/MAL, 1957KEM/GOL, 1972DYK]
C ₅ H ₁₀ N ₂ O ₆	[67727-92-8] Δ_vH	1-(methoxymethoxy)-2,2-dinitropropane (293–333)	71.3	308	A	[1987STE/MAL]
C ₅ H ₁₀ N ₄ O ₄	[5754-90-5] $\Delta_{\text{trs}}H$ $\Delta_{\text{fus}}H$	1,3-dinitro-1,3-diazacycloheptane	21.8 2.8	369 374		[1991PIC/RYL]
C ₅ H ₁₀ O	[557-31-3] Δ_vH	allyl ethyl ether (244–401)	34.6	259	A	[1987STE/MAL]
C ₅ H ₁₀ O	[616-25-1] Δ_vH Δ_vH Δ_vH	1-penten-3-ol	49.9 ± 0.1 48.4 ± 0.1 46.8 ± 0.1	313 328 343	C C C	[1996ULB/KLU] [1996ULB/KLU] [1996ULB/KLU]
C ₅ H ₁₀ O	[556-82-1] Δ_vH	3,3-dimethyl-2-propen-1-ol (348–372)	48.7	360	EB	[1989WAN/YIN]
C ₅ H ₁₀ O	[115-18-4] Δ_vH	2-methyl-3-buten-2-ol (290–372)	43.1 ± 0.1	331		[1988BAG/GUR]
C ₅ H ₁₀ O	[763-32-6] Δ_vH	3-buten-3-methyl-1-ol (338–409)	55.6	353	A	[1987STE/MAL]
C ₅ H ₁₀ O	[na] Δ_vH	(<i>dl</i>) 3-buten-3-methyl-2-ol (358–379)	41.0	368	A	[1987STE/MAL]
C ₅ H ₁₀ O	[96-41-3] $\Delta_{\text{trs}}H$ $\Delta_{\text{fus}}H$ Δ_vH Δ_vH Δ_vH Δ_vH Δ_vH Δ_vH Δ_vH	cyclopentanol	3.71 1.54 57.1 52.7 56.1 56.4 57.1 57.5 ± 0.2 57.5 ± 0.3	202.8 257.4 298 361 298 298 294 298 298	CGC A,EB A A C C	[1996DOM/HEA] [1995CHI/HOS] [1987AMB/GHI2] [1987STE/MAL] [1987STE/MAL] [1975CAB/CON2] [1968PLA/WIL] [1966WAD]
C ₅ H ₁₀ O	[142-68-7] Δ_vH Δ_vH Δ_vH Δ_vH Δ_vH	tetrahydropyran (286–361) (335–412) (273–362) (273–288)	36 38.2 ± 1.1 33.2 35.0 35.0	301 298 350 288 281	DSC A	[2006ROD/GIN] [2005ROJ/GIN] [2000ROD/ART] [1987STE/MAL] [1972DYK, 1958CAS/FLE]
C ₅ H ₁₀ O	[96-47-9] Δ_vH Δ_vH	2-methyltetrahydrofuran (283–353)	34.0 33.7	298 298	A	[1987STE/MAL] [1970MOI/ANT]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₅ H ₁₀ O	[563-80-4]	3-methyl-2-butanone				
	$\Delta_{\text{fus}}H$		9.34	180		[1996DOM/HEA]
	Δ_vH	(311–369)	35.5	326	A	[1987STE/MAL]
	Δ_vH	(363–415)	33.8	378	A	[1987STE/MAL]
	Δ_vH	(405–500)	32.6	420	A	[1987STE/MAL]
	Δ_vH		36.8	298	C	[1983UCH/MAJ]
	Δ_vH	(328–377)	35.0	343	A	[1987STE/MAL, 1975AMB/ELL]
	Δ_vH		36.9	298		[1975AMB/ELL]
	Δ_vH		35.0 ± 0.1	327	C	[1967HAL/LEE]
			33.8 ± 0.1	346	C	[1967HAL/LEE]
			32.3 ± 0.1	367	C	[1967HAL/LEE]
C ₅ H ₁₀ O	[107-87-9]	2-pentanone				
	$\Delta_{\text{trs}}H$		2.09	110		[1991ACR]
	$\Delta_{\text{fus}}H$		10.63	196.3		
	Δ_vH	(336–422)	36.1	351	A	[1987STE/MAL]
	Δ_vH	(416–501)	33.7	431	A	[1987STE/MAL]
	Δ_vH	(487–561)	33.3	502	A	[1987STE/MAL]
	Δ_vH		38.4	298	C	[1983UCH/MAJ]
	Δ_vH		38.3 ± 0.3	298	GCC	[1979SAL/PEA]
	Δ_vH		38.4	298		[1975AMB/ELL]
	Δ_vH	(268–373)	39.5	283	EB	[1966MEY/WAG]
	Δ_vH	(329–385)	36.5	344	A,GS,EB	[1987STE/MAL, 1975AMB/ELL, 1965COL/COU, 1972DYK]
	Δ_vH		36.1 ± 0.1	335	C	[1961NIC/KOB]
	Δ_vH		34.4 ± 0.1	360	C	[1961NIC/KOB]
	Δ_vH		33.4 ± 0.1	375	C	[1961NIC/KOB]
Δ_vH		32.8 ± 0.1	386	C	[1961NIC/KOB]	
Δ_vH		32.2 ± 0.1	394	C	[1961NIC/KOB]	
C ₅ H ₁₀ O	[96-22-0]	3-pentanone				
	$\Delta_{\text{trs}}H$		0.11	118.5		
	$\Delta_{\text{trs}}H$		0.01	180		
	$\Delta_{\text{fus}}H$		11.59	234.2		[1991ACR]
	Δ_vH	(290–375)	35.9 ± 0.2	332		[1988BAG/GUR]
	Δ_vH	(329–426)	36.6	344	A	[1987STE/MAL]
	Δ_vH	(421–502)	33.7	436	A	[1987STE/MAL]
	Δ_vH	(494–561)	33.3	509	A	[1987STE/MAL]
	Δ_vH		38.5	298	C	[1983UCH/MAJ]
	Δ_vH		38.7 ± 0.3	298	GCC	[1979SAL/PEA]
	Δ_vH		38.6	298		[1975AMB/ELL]
	Δ_vH		36.1 ± 0.1	335	C	[1967HAL/LEE]
	Δ_vH		34.9 ± 0.1	354	C	[1967HAL/LEE]
	Δ_vH		33.5 ± 0.1	375	C	[1967HAL/LEE]
	Δ_vH	(329–384)	36.6	344	A,GS,EB	[1987STE/MAL, 1975AMB/ELL]
Δ_vH					[1965COL/COU, 1972DYK]	
Δ_vH	(283–323)	36.9	303		[1937RIN/SAY]	
C ₅ H ₁₀ O	[110-62-3]	pentanal				
	$\Delta_{\text{fus}}H$		15.0	151.6		[1998VAS/LEB]
	Δ_vH	(307–343)	38.6	298	EB	[2002ANT/FRA, 2003VER/KRA2]
	Δ_vH	(313–353)	38.3	298	CGC	[1995CHI/HOS]
	Δ_vH	(290–385)	U 50.0	305	A	[1987STE/MAL]
Δ_vH		38.1 ± 0.1	298		[1981DYA/KOR]	

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₅ H ₁₀ O	$\Delta_v H$	(305–377)	37.3	320		[1979MAR/SAC]
	[630-19-3]	pivaldehyde				
	$\Delta_{\text{trs}} H$		0.5	158.5		
	$\Delta_{\text{fus}} H$		4.81	183.9		
C ₅ H ₁₀ O	$\Delta_v H$	(308–336)	34.2	322		[1988WHI/PER]
	[6921-35-3]	3,3-dimethyloxetane				
	$\Delta_v H$		33.9 ± 0.3	298	C	[1989VAR/SOM]
C ₅ H ₁₀ OS	[2307-10-0]	S-propyl thiolacetate				
	$\Delta_v H$		44.1 ± 0.2	298	C	[1966WAD]
C ₅ H ₁₀ OS	[926-73-8]	S-isopropyl thiolacetate				
	$\Delta_v H$		42.3 ± 0.2	298	C	[1966WAD]
C ₅ H ₁₀ OS	[6607-53-0]	1-(methylthio)-2-(vinylloxy)ethane				
	$\Delta_v H$	(316–347)	47.5	331	A	[1987STE/MAL, 1999DYK/SVO]
C ₅ H ₁₀ O ₂	[97-99-4]	tetrahydrofurfuryl alcohol				
	$\Delta_v H$	(393–453)	46.2	408	A	[1987STE/MAL]
	$\Delta_v H$	(333–443)	46.5	388		[1979BLU/BAE]
C ₅ H ₁₀ O ₂	[2916-31-6]	2,2-dimethyl-1,3-dioxolane				
	$\Delta_v H$	(278–318)	41.1 ± 0.2		GS	[1998VER/PEN, 2002VER]
C ₅ H ₁₀ O ₂	[1120-97-4]	4-methyl-1,3-dioxane				
	$\Delta_v H$	(273–313)	43.7 ± 0.3		GS	[1998VER/PEN, 2002VER]
C ₅ H ₁₀ O ₂	[50741-70-3]	1-methoxy-2-butanone				
	$\Delta_v H$	(297–408)	44.9	312	A	[1987STE/MAL, 1934RIG/FEL, 1972DYK]
C ₅ H ₁₀ O ₂	[115-22-0]	3-hydroxy-3-methyl-2-butanone				
	$\Delta_v H$	(317–419)	41.1	332	A	[1987STE/MAL, 1972DYK, 1950CON/ELV]
C ₅ H ₁₀ O ₂	[3393-64-4]	4-hydroxy-3-methyl-2-butanone				
	$\Delta_v H$	(375–528)	58.3	390	A	[1987STE/MAL, 1972DYK]
	$\Delta_v H$	(317–458)	59.0	332		[1947STU]
C ₅ H ₁₀ O ₂	[592-84-7]	butyl formate				
	$\Delta_v H$	(313–359)	40.1	298	EB	[2004CHY/FRA]
	$\Delta_v H$	(295–380)	37.9	310	A	[1987STE/MAL]
	$\Delta_v H$		41.3 ± 0.1	298	C	[1980SVO/UCH]
	$\Delta_v H$		40.1 ± 0.1	313	C	[1980SVO/UCH]
	$\Delta_v H$		39.0 ± 0.1	328	C	[1980SVO/UCH]
	$\Delta_v H$		38.7 ± 0.1	346	C	[1976CIH/HYN]
	$\Delta_v H$		38.1 ± 0.1	355	C	[1976CIH/HYN]
C ₅ H ₁₀ O ₂	[589-40-2]	sec butyl formate				
	$\Delta_v H$	(238–367)	37.7	253	A	[1987STE/MAL]
C ₅ H ₁₀ O ₂	[105-37-3]	ethyl propionate				
	$\Delta_v H$	(315–420)	36.7	330		[1997HER/ORT]
	$\Delta_v H$	(372–538)	34.4	387	A	[1987STE/MAL]
	$\Delta_v H$		39.3 ± 0.1	298	C	[1980SVO/UCH]
	$\Delta_v H$		38.2 ± 0.1	313	C	[1980SVO/UCH]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		T _m (K)	Method	Reference	
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)				
		$\Delta_v H$	36.6 ± 0.1	336	C	[1977SVO/VES]	
		$\Delta_v H$	36.0 ± 0.1	344	C	[1977SVO/VES]	
		$\Delta_v H$	35.5 ± 0.1	351	C	[1977SVO/VES]	
		$\Delta_v H$	34.5 ± 0.1	363	C	[1977SVO/VES]	
		$\Delta_v H$	39.1 ± 0.1	298	C	[1972MAN]	
		$\Delta_v H$	(306–372)	38.2	321	A	[1987STE/MAL, 1965MER/POL, 1972DYK]
C ₅ H ₁₀ O ₂	[542-55-2]	isobutyl formate					
		$\Delta_v H$	(371–507)	36.6	386	A	[1987STE/MAL]
		$\Delta_v H$	(240–372)	38.6	255	A	[1987STE/MAL, 1947STU]
C ₅ H ₁₀ O ₂	[108-21-4]	isopropyl acetate					
		$\Delta_v H$	(313–353)	37.0	298	CGC	[1995CHI/HOS]
		$\Delta_v H$		35.6	323	C	[1973GEI/QUI]
		$\Delta_v H$		37.2 ± 0.2	298	C	[1966WAD]
		$\Delta_v H$	(235–362)	38.8	250	A	[1987STE/MAL, 1947STU]
		$\Delta_v H$	(273–363)	36.3	288	A	[1929HAG/WEI]
C ₅ H ₁₀ O ₂	[623-42-7]	methyl butyrate					
		$\Delta_v H$	(274–303)	40.3 ± 0.5	298	GS	[2008VER/EME]
		$\Delta_v H$	(317–360)	38.4	332		[2002SWI/MAL]
		$\Delta_v H$		36.9	350		[2002VAN/VAN]
		$\Delta_v H$		41.1 ± 0.2	284		[2002VAN/VAN]
		$\Delta_v H$		40.6 ± 0.2	298		[2002VAN/VAN]
		$\Delta_v H$	(317–360)	40.6 ± 0.1	298	EB	[2002CON/WIC]
		$\Delta_v H$	(333–378)	39.3	298	CGC	[1999VER/HEI]
		$\Delta_v H$	(349–384)	36.4	364		[1990ORT/SUS]
		$\Delta_v H$		39.0 ± 0.4	298	GC	[1987AZA]
		$\Delta_v H$	(375–545)	34.2	390	A	[1987STE/MAL]
		$\Delta_v H$	(345–383)	40.4	298	EB	[1984WIS/TAM, 2008VER/EME]
		$\Delta_v H$		40.1 ± 0.4	298	C	[1981GAT/STR]
		$\Delta_v H$		39.8 ± 0.3	298	GCC	[1980FUC/PEA]
		$\Delta_v H$		39.3 ± 0.2	298	C	[1979SUN/SVE2]
		$\Delta_v H$	(246–375)	42.8	261	A	[1987STE/MAL, 1947STU]
C ₅ H ₁₀ O ₂	[547-63-7]	methyl isobutyrate					
		$\Delta_v H$	(366–533)	33.7	381	A	[1987STE/MAL]
		$\Delta_v H$		37.3	298		[UR/FUC, 1985MAJ/SVO]
		$\Delta_v H$	(239–366)	40.1	254	A	[1987STE/MAL, 1947STU]
C ₅ H ₁₀ O ₂	[109-60-4]	propyl acetate					
		$\Delta_v H$	(313–363)	37.7	298	CGC	[1995CHI/HOS]
		$\Delta_v H$	(333–372)	37.0	348		[1993FAR/WIC]
		$\Delta_v H$	(374–542)	34.8	389	A	[1987STE/MAL]
		$\Delta_v H$	(322–383)	38.1	327	DTA	[1980MEY/AWE]
		$\Delta_v H$		39.8 ± 0.1	298	C	[1980SVO/UCH]
		$\Delta_v H$		38.6 ± 0.1	313	C	[1980SVO/UCH]
		$\Delta_v H$		35.3 ± 0.1	343	C	[1980SVO/UCH]
		$\Delta_v H$		36.9 ± 0.1	336	C	[1977SVO/VES]
		$\Delta_v H$		36.4 ± 0.1	344	C	[1977SVO/VES]
		$\Delta_v H$		35.8 ± 0.1	351	C	[1977SVO/VES]
		$\Delta_v H$		34.8 ± 0.1	363	C	[1977SVO/VES]
		$\Delta_v H$		36.9	335		[1976CON/COU]
		$\Delta_v H$		33.9	375		[1976CON/COU]
		$\Delta_v H$		36.7	335	C	[1973GEI/QUI]
		$\Delta_v H$		39.1 ± 0.2	298	C	[1966WAD]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound					
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference	
C ₅ H ₁₀ O ₂	$\Delta_v H$	(312–374)	38.2	327	A	[1987STE/MAL, 1965MER/POL, 1972DYK]	
	[75-98-9]	trimethylacetic acid (pivalic acid)					
	$\Delta_{\text{fus}} H$		2.3	309.1		[2002STE/CHI4]	
	$\Delta_{\text{trs}} H$		8.18	278.3			
	$\Delta_{\text{fus}} H$		2.27	309.1		[1990SIN/GLI]	
	$\Delta_{\text{sub}} H$		62.3	291	GS	[2000VER2]	
	$\Delta_v H$	(344–472)	57.6 ± 0.2	320	EB	[2002STE/CHI4]	
	$\Delta_v H$	(344–472)	54.4 ± 0.2	360	EB	[2002STE/CHI4]	
	$\Delta_v H$	(344–472)	50.9 ± 0.2	400	EB	[2002STE/CHI4]	
C ₅ H ₁₀ O ₂	[109-52-4]	valeric acid (pentanoic acid)					
	$\Delta_{\text{fus}} H$		14.16	239.5		[1996DOM/HEA]	
	$\Delta_v H$	(283–313)	63.0 ± 9.5	298	GS	[2000VER2]	
	$\Delta_v H$	(353–393)	65.9	298	CGC	[1995CHI/HOS]	
	$\Delta_v H$	(373–465)	57.9	388	EB	[1987AMB/GHI3]	
	$\Delta_v H$	(375–523)	58.0	390	A	[1987STE/MAL]	
	$\Delta_v H$	(243–266)	62.4 ± 3	298	TE	[1979DEK/OON]	
C ₅ H ₁₀ O ₂	[503-74-2]	3-methylbutanoic acid (isovaleric acid)					
	$\Delta_v H$	(396–448)	53.8	411		[2004CLI/RAM]	
	$\Delta_v H$	(293–323)	60.7 ± 0.3	308	GS	[2000VER2]	
	$\Delta_v H$	(293–323)	61.2 ± 0.3	298	GS	[2000VER2]	
	$\Delta_v H$	(364–464)	55.8	379	A,EB	[1987AMB/GHI3]	
	$\Delta_v H$	(307–448)	56.6	322	A	[1987STE/MAL]	
	$\Delta_v H$	(243–259)	57.5 ± 3	298	TE	[1979DEK/OON]	
	$\Delta_v H$ (monomer)		46.9 ± 0.2	298	C	[1970KON/WAD]	
C ₅ H ₁₀ O ₃	[105-58-8]	diethyl carbonate					
	$\Delta_{\text{fus}} H$		9.24	198.2	DSC	[2004DIN]	
	$\Delta_v H$	(344–398)	42.3	359	EB	[2009XIN/FAN]	
	$\Delta_v H$	(273–315)	44.4 ± 0.2	298	GS	[2008KOZ/EME]	
	$\Delta_v H$	(352–403)	39.7	367		[2002ROD/CAN2]	
	$\Delta_v H$	(308–400)	40.9	323	A	[1987STE/MAL]	
	$\Delta_v H$		43.6 ± 0.2	298	C	[1973COU/LEE]	
	$\Delta_v H$	(308–368)	39.1		MM	[1971CHO/JON]	
C ₅ H ₁₀ O ₃	[110-49-6]	ethylene glycol methyl ethyl acetate					
	$\Delta_v H$		50.3 ± 0.1	298	C	[1970KUS/WAD]	
	$\Delta_v H$	(343–417)	44.3	358	A	[1987STE/MAL, 1957DYK/SEP, 1972DYK]	
	C ₅ H ₁₀ O ₃	[97-64-3]	(dl) ethyl lactate				
		$\Delta_v H$	(308–426)	49.2	323	A	[1987STE/MAL]
		$\Delta_v H$	(324–427)	51.3	339	A	[1987STE/MAL]
	C ₅ H ₁₀ O ₃	[623-72-3]	3-hydroxypropionic acid, ethyl ester				
$\Delta_v H$		(338–356)	62.2	347	A	[1987STE/MAL]	
C ₅ H ₁₀ O ₃	[3852-09-3]	3-methoxypropionic acid, methyl ester					
$\Delta_v H$	(350–438)	43.4	370	A	[1987STE/MAL]		

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₅ H ₁₀ O ₃	[1779-19-7]	1,3,6-trioxacyclooctane				
	$\Delta_v H$		48.8 ± 0.2	298	C	[1982BYS/MAN]
C ₅ H ₁₀ O ₄	[106-61-6]	glycerol, 1-monoacetate				
	$\Delta_v H$	(385–458)	74.0	400	A	[1987STE/MAL, 1972DYK]
C ₅ H ₁₀ O ₄	[4767-03-7]	2,2-bis-hydroxymethylpropanoic acid				
	$\Delta_{\text{trs}} H$		38.5	426		
	$\Delta_{\text{fus}} H$		3.59	468		[1996DOM/HEA]
C ₅ H ₁₀ O ₅	[16528-92-0]	1,3,5,7,9-pentoxecane				
	$\Delta_{\text{fus}} H$		21.88	334	DSC	[1996DOM/HEA]
	$\Delta_{\text{sub}} H$		87.9 ± 0.5	298	C	[1974MAN2]
C ₅ H ₁₀ O ₅	[58-86-6]	<i>(d)</i> -xylose				
	$\Delta_{\text{fus}} H$		31.7	416.2		[2002JON/COO]
	$\Delta_{\text{sub}} H$	(370–395)	158.0 ± 3.1	382	ME	[1999OJA/SUU]
C ₅ H ₁₀ S	[5296-62-8]	allyl ethyl sulfide				
	$\Delta_v H$	(300–327)	38.9	313	A,EB	[1987STE/MAL, 1962MAC/MAY3, 1999DYK/SVO]
	$\Delta_v H$	(300–327)	39.3	298		[1962MAC/MAY3]
C ₅ H ₁₀ S	[1679-07-8]	cyclopentanethiol				
	$\Delta_{\text{fus}} H$		7.83	155.4		[1996DOM/HEA]
	$\Delta_v H$	(354–446)	38.2	369		[1999DYK/SVO]
	$\Delta_v H$	(348–446)	38.4	363	A,EB	[1987STE/MAL, 1961BER/SCO, 1966OSB/DOU]
	$\Delta_v H$		37.9 ± 0.1	361	C	[1961BER/SCO]
	$\Delta_v H$		36.7 ± 0.1	381	C	[1961BER/SCO]
	$\Delta_v H$		35.3 ± 0.1	405	C	[1961BER/SCO]
C ₅ H ₁₀ S	[1795-09-1]	2-methyltetrahydrothiophene				
	$\Delta_{\text{fus}} H$		8.87	172.4		[1974MES/FIN, 1996DOM/HEA]
	$\Delta_v H$	(303–433)	40.6	318		[1999DYK/SVO]
	$\Delta_v H$		41.8	298		[1971WIL/ZWO]
	$\Delta_v H$		41.3 ± 0.1	298		[1972GOO, 1966OSB/DOU]
	$\Delta_v H$	(335–447)	39	350	A,EB	[1987STE/MAL, 1966OSB/DOU]
C ₅ H ₁₀ S	[4740-00-5]	3-methyltetrahydrothiophene				
	$\Delta_{\text{fus}} H$		10.37	192		[1974MES/FIN, 1996DOM/HEA]
	$\Delta_v H$	(307–439)	41.3	322		[1999DYK/SVO]
	$\Delta_v H$		42.7	298		[1971WIL/ZWO]
	$\Delta_v H$		42.1 ± 0.1	298		[1972GOO, 1966OSB/DOU]
	$\Delta_v H$	(340–453)	39.6	355	A,EB	[1987STE/MAL, 1966OSB/DOU]
C ₅ H ₁₀ S	[1613-51-0]	pentamethylene sulfide				
	$\Delta_{\text{trs}} H$		1.1	201.4		
	$\Delta_{\text{trs}} H$		7.77	240		
	$\Delta_{\text{fus}} H$		2.45	292.3		[1996DOM/HEA]
	$\Delta_v H$	(310–443)	41.4	325		[1999DYK/SVO]
	$\Delta_v H$	(338–393)	37.2	345	EB	[1984PAL/CHO]
			42.8	298	C	[1971WIL/ZWO]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
			39.7	351		[1954MCC/FIN]
			39.5	362	A,EB	[1987STE/MAL, 1952WHI/BER]
C ₅ H ₁₁ Br	[110-53-2]	1-bromopentane				
			14.37	185.1		[1996DOM/HEA]
			40.9	298	CGC	[1995CHI/HOS]
			41.4 ± 0.1	298	C	[1968WAD]
			41.1 ± 0.1	298	C	[1966WAD]
			41.0	308	A,EST	[1987STE/MAL, 1961LI/ROS, 1972DYK]
C ₅ H ₁₁ Br	[107-81-3]	(<i>dl</i>) sec-pentylbromide, 2-bromopentane				
			37.5	318	A	[1987STE/MAL, 1972DYK]
C ₅ H ₁₁ Br	[107-81-3]	2-bromopentane				
			38.5	298	CGC	[1995CHI/HOS]
C ₅ H ₁₁ Br	[1809-10-5]	3-bromopentane				
			8.4	167.3		[1995TAK/YAM]
			37.7	319	A	[1987STE/MAL, 1972DYK]
C ₅ H ₁₁ Br	[630-17-1]	1-bromo-2,2-dimethylpropane				
			35.6	308	A	[1987STE/MAL, 1972DYK]
C ₅ H ₁₁ Br	[10422-35-2]	1-bromo-2-methylbutane				
			37.9	321	A	[1987STE/MAL, 1972DYK]
C ₅ H ₁₁ Br	[107-82-4]	1-bromo-3-methylbutane				
			37.9	321	A	[1987STE/MAL, 1972DYK]
			41	268		[1947STU]
C ₅ H ₁₁ Br	[507-36-8]	2-bromo-2-methylbutane				
			36.4	310	A	[1987STE/MAL, 1972DYK]
C ₅ H ₁₁ Br	[18295-25-5]	2-bromo-3-methylbutane				
			37.2	316	A	[1987STE/MAL, 1972DYK]
C ₅ H ₁₁ Cl	[543-59-9]	1-chloropentane				
			38.8	298	CGC	[1995CHI/HOS]
			38.2	298	C	[1981TEK/MAJ]
			37.3	313	C	[1981TEK/MAJ]
			36.5	328	C	[1981TEK/MAJ]
			35.6	343	C	[1981TEK/MAJ]
			34.6	358	C	[1981TEK/MAJ]
			34.0	363	C	[1981TEK/MAJ]
			38.2 ± 0.1	298	C	[1968WAD]
			38.7	292	A,EST	[1987STE/MAL, 1961LI/ROS]
						[1972DYK]
C ₅ H ₁₁ Cl	[625-29-6]	2-chloropentane				
			36.0	298	C	[1981TEK/MAJ]
			35.2	313	C	[1981TEK/MAJ]
			34.4	328	C	[1981TEK/MAJ]
			33.5	358	C	[1981TEK/MAJ]
			31.9	368	C	[1981TEK/MAJ]
			36.2	304	A	[1987STE/MAL, 1972DYK]
C ₅ H ₁₁ Cl	[616-20-6]	3-chloropentane				

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
	$\Delta_{\text{v}}H$	(289–410)	36.5	304	A	[1987STE/MAL, 1972DYK]
C ₅ H ₁₁ Cl	[753-89-9]	1-chloro-2,2-dimethylpropane				
	$\Delta_{\text{v}}H$	(279–395)	34.9	294	A	[1987STE/MAL, 1972DYK]
C ₅ H ₁₁ Cl	[616-13-7]	<i>(dl)</i> 1-chloro-2-methylbutane				
	$\Delta_{\text{v}}H$	(300–374)	35.4	315	A	[1987STE/MAL, 1972DYK]
C ₅ H ₁₁ Cl	[594-36-5]	2-chloro-2-methylbutane				
	$\Delta_{\text{v}}H$	(280–396)	35.0	295	A	[1987STE/MAL, 1972DYK]
C ₅ H ₁₁ Cl	[631-65-2]	2-chloro-3-methylbutane				
	$\Delta_{\text{v}}H$	(285–405)	35.9	300	A	[1987STE/MAL, 1972DYK]
C ₅ H ₁₁ Cl	[107-84-6]	1-chloro-3-methylbutane				
	$\Delta_{\text{v}}H$	(313–353)	38.1	298	CGC	[1995CHI/HOS]
	$\Delta_{\text{v}}H$		36.2	298	C	[1981TEK/MAJ]
	$\Delta_{\text{v}}H$		35.4	313	C	[1981TEK/MAJ]
	$\Delta_{\text{v}}H$		34.6	328	C	[1981TEK/MAJ]
	$\Delta_{\text{v}}H$		33.7	343	C	[1981TEK/MAJ]
	$\Delta_{\text{v}}H$		32.8	358	C	[1981TEK/MAJ]
	$\Delta_{\text{v}}H$		32.3	368	C	[1981TEK/MAJ]
C ₅ H ₁₁ ClO ₂ S	[6303-18-0]	1-pentanesulfonyl chloride				
	$\Delta_{\text{v}}H$	(293–387)	58.5	308		[1999DYK/SVO]
	$\Delta_{\text{v}}H$	(387–492)	55.1	402		[1999DYK/SVO]
	$\Delta_{\text{v}}H$	(263–293)	60.5	278	A	[1987STE/MAL, 1999DYK/SVO]
C ₅ H ₁₁ Cl ₂ N	[51-75-2]	N-methyl- <i>bis</i> (2-chloroethyl)amine				
	$\Delta_{\text{v}}H$	(273–333)	54.6	288	A	[1987STE/MAL]
C ₅ H ₁₁ F	[592-50-7]	1-fluoropentane				
	$\Delta_{\text{v}}H$	(245–373)	33.7	260	EST	[1987STE/MAL, 1961LI/ROS, 1972DYK]
C ₅ H ₁₁ F	[10086-64-3]	1-fluoro-2-methylbutane				
	$\Delta_{\text{v}}H$	(287–329)	30.7	302	A	[1987STE/MAL, 1972DYK]
C ₅ H ₁₁ F	[661-53-0]	2-fluoro-2-methylbutane				
	$\Delta_{\text{v}}H$	(249–341)	31.8	264	A	[1987STE/MAL, 1972DYK]
C ₅ H ₁₁ I	[628-17-1]	1-iodopentane				
	$\Delta_{\text{v}}H$	(313–353)	44.4	298	CGC	[1995CHI/HOS]
	$\Delta_{\text{v}}H$		45.3 ± 0.1	298	C	[1968WAD]
	$\Delta_{\text{v}}H$	(312–473)	43.1	327	A,EST	[1987STE/MAL, 1961LI/ROS, 1972DYK]
C ₅ H ₁₁ I	[616-14-8]	1-iodo-2-methylbutane				
	$\Delta_{\text{v}}H$	(339–406)	39.8	354	A	[1987STE/MAL, 1972DYK]
C ₅ H ₁₁ I	[541-28-6]	1-iodo-3-methylbutane				
	$\Delta_{\text{v}}H$	(313–353)	42.2	298	CGC	[1995CHI/HOS]
	$\Delta_{\text{v}}H$	(270–422)	43.5	285	A	[1987STE/MAL, 1947STU]
C ₅ H ₁₁ I	[594-38-7]	2-iodo-2-methylbutane				
	$\Delta_{\text{v}}H$	(308–398)	40.4	323	A	[1987STE/MAL, 1972DYK]
C ₅ H ₁₁ N	[1003-03-8]	cyclopentylamine				
	$\Delta_{\text{trs}}H$		0.48	184.5		
	$\Delta_{\text{fus}}H$		8.31	190.4		[1996DOM/HEA]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(317–419)	38.3	332	EB	[1987STE/MAL, 1975AMB/CON]
	$\Delta_v H$	(317–419)	40.2 ± 0.4	298	EB	[1975GOO/MES]
C ₅ H ₁₁ N	[120-94-5]	1-methylpyrrolidine				
	$\Delta_v H$	(270–298)	35.0 ± 0.7	284	GS	[1998VER6]
	$\Delta_v H$	(270–298)	34.2 ± 0.7	298	GS	[1998VER6]
	$\Delta_v H$	(273–315)	33.7	288	A	[1987STE/MAL]
C ₅ H ₁₁ N	[110-89-4]	piperidine				
	$\Delta_{\text{fus}} H$		14.85	262.1		[1996DOM/HEA]
	$\Delta_v H$		36.6	338		[1988HOS/ARC]
	$\Delta_v H$	(315–417)	35.3	357		[1988HOS/ARC]
C ₅ H ₁₁ NO	$\Delta_v H$	(315–417)	37.6	330	A,EB,IP	[1987STE/MAL, 1968OSB/DOU]
	[617-84-5]	N,N-diethylformamide				
	$\Delta_v H$		50.3	298		[1985BAR/CAS, 1985MAJ/SVO]
C ₅ H ₁₁ NO	$\Delta_v H$	(303–363)	48.9	318	A	[1987STE/MAL, 1968GOP/RIZ]
	[1118-69-0]	N-isopropylacetamide				
C ₅ H ₁₁ NO	$\Delta_v H$		66.4 ± 0.3	298	C	[1984STA/WAD]
	[5331-48-6]	N-propylacetamide				
C ₅ H ₁₁ NO	$\Delta_v H$		69.8 ± 0.2	298	C	[1984STA/WAD]
	[626-97-1]	pentanamide				
C ₅ H ₁₁ NO	$\Delta_{\text{trs}} H$		1.9	211.8		
	$\Delta_{\text{trs}} H$		1.2	365		
	$\Delta_{\text{fus}} H$		17.9	377.2		[2008ABA/BAD]
	$\Delta_{\text{sub}} H$	(333–374)	89.3 ± 0.4		GS	[1959DAV/JON2, 1970COX/PIL]
	$\Delta_{\text{sub}} H$	(353–373)	89.1			[1960JON]
C ₅ H ₁₁ NO	[759-10-9]	2,2-dimethylpropanamide				
	$\Delta_{\text{fus}} H$		24.1	425.4		[2008ABA/BAD]
	$\Delta_{\text{sub}} H$	(298–359)	89 ± 2.0	298	TE	[2000BRU/DEL]
	$\Delta_{\text{sub}} H$	(288–306)	86.6 ± 0.4	298		[1989ABB/JIM]
C ₅ H ₁₁ NO	[758-96-3]	N,N-dimethylpropionamide				
	$\Delta_v H$	(326–424)	53.5	341	A	[1987STE/MAL]
	$\Delta_v H$		52.9			[1977VAS/KOT]
C ₅ H ₁₁ NO	[2675-88-9]	N-methyl-2-methylpropionamide				
	$\Delta_v H$		67.1 ± 0.2	298	C	[1984STA/WAD]
C ₅ H ₁₁ NO	[na]	methyl 2-(N,N-dimethylamino)propanoate				
	$\Delta_v H$	(278–308)	43.9 ± 0.4	293	GS	[1992VER/BECT]
C ₅ H ₁₁ NO	[109-02-4]	N-methylmorpholine				
	$\Delta_v H$	(273–353)	39.5	298		[2009BEL/RAZ]
	$\Delta_v H$	(273–353)	38.9	313		[2009RAZ/HAJ]
	$\Delta_v H$	(273–353)	39.8	298		[2009RAZ/HAJ]
	$\Delta_v H$		38.2 ± 1.1	298	DSC	[2005ROJ/GIN]
	$\Delta_v H$	(274–304)	40.2 ± 0.3	288	GS	[1998VER2]
	$\Delta_v H$	(274–304)	39.6 ± 0.3	298	GS	[1998VER2]
	$\Delta_v H$	(323–363)	33.6	343	TGA	[1987ALN/ALS]
	$\Delta_v H$	(297–389)	38.4	312	A	[1987STE/MAL]
	$\Delta_v H$	(276–390)	40.0	291	A	[1987STE/MAL]
$\Delta_v H$	(276–319)	39.4 ± 0.1	298		[1975CAB/CON]	

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₅ H ₁₁ NO	[15364-56-4] $\Delta_v H$	1-(dimethylamino)-2-propanone (298–338)	43.6 ± 0.3	298	GS	[1994WEL/VER]
C ₅ H ₁₁ NO	[1188-11-0] $\Delta_v H$	3-pentanone oxime (318–425)	55.8	333	A	[1987STE/MAL]
C ₅ H ₁₁ NO ₂	[628-05-7] $\Delta_v H$	1-nitropentane (278–318)	50.3 ± 0.2	298	GS	[1997VER3]
C ₅ H ₁₁ NO ₂	[543-28-2] $\Delta_v H$	isobutyl carbamate (356–479)	58.8	371	A	[1987STE/MAL, 1947STU]
C ₅ H ₁₁ NO ₂	[31502-31-5] $\Delta_v H$	N,N-dimethyl lactamide (351–417)	73.7	366	A	[1987STE/MAL]
C ₅ H ₁₁ NO ₂	[760-78-1] $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$	(<i>dl</i>) 2-aminopentanoic acid (DL-norvaline) (439–461)	120 121.1 ± 0.4	450 455	ME	[1987STE/MAL] [1965SVE/CLY, 1964CLY/SVE]
C ₅ H ₁₁ NO ₂	[592-35-8] $\Delta_{\text{sub}} H$	butyl carbamate (292–316)	94.1 ± 8		GS	[1959DAV/JON]
C ₅ H ₁₁ NO ₂	[465-58-7] $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$	(<i>dl</i>)-2-amino-2-methyl-butanoic acid (439–469)	134.2 ± 1 125.8 ± 0.4	413 454	TE,ME	[1979DEK/VOO] [1965SVE/CLY, 1964CLY/SVE]
C ₅ H ₁₁ NO ₂	[516-06-3] $\Delta_{\text{sub}} H$	DL-valine (320–420)	U 79.5 ± 8	370	LE	[1977GAF/PIE]
C ₅ H ₁₁ NO ₂	[72-18-4] $\Delta_{\text{sub}} H$	L-valine	162.8 ± 8	455	ME	[1965SVE/CLY, 1964CLY/SVE, 1989CHI/GRO]
C ₅ H ₁₁ NO ₂	[660-88-8] $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$	5-aminopentanoic acid (384–394)	141.8 ± 0.5 144 ± 3	389 289	C C	[1983SKO/SAB] [1983SKO/SAB]
C ₅ H ₁₁ NO ₂	[7529-22-8] $\Delta_{\text{fus}} H$	N-methylmorpholine-N-oxide	18.8	457.4		[1981NAV/HAU]
C ₅ H ₁₁ NO ₂ S	[59-51-8] $\Delta_{\text{sub}} H$	DL-methionine (363–463)	U 134 ± 8	413	LE	[1977GAF/PIE]
C ₅ H ₁₁ NO ₂ S	[63-68-3] $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$	L-(<i>d</i>)-methionine (463–485)	164 ± 4 125	298 474	C A	[1981SAB/MIN] [1987STE/MAL]
C ₅ H ₁₁ NO ₂ S	[348-67-4] $\Delta_{\text{sub}} H$	D-(<i>l</i>)-methionine	125.1 ± 0.8	455	ME	[1965SVE/CLY, 1964CLY/SVE]
C ₅ H ₁₁ NO ₃	[543-87-3] $\Delta_v H$	isopentyl nitrate (278–421)	47.0	293	A	[1987STE/MAL, 1947STU]
C ₅ H ₁₁ NO ₃ S	[14357-44-9] $\Delta_{\text{fus}} H$	2-methyl-2-(methylsulfonyl)propanal oxime	27.12	382	DSC	[1990DON/DRE]
C ₅ H ₁₁ O ₂ PS ₂	[77240-15-4] $\Delta_v H$	2-mercapto-4,6-dimethyl-1,3,2-dioxaphosphorinane-2-sulfide	72.3			[2008SAG/SAF]
C ₅ H ₁₁ P	[4743-40-2] $\Delta_{\text{sub}} H$	phosphorinane (250–291)	43.3	276	T	[1987STE/MAL, 1966MOR/TAM]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound					
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference	
C ₅ H ₁₂	$\Delta_v H$	(294–345)	39.9	309	A,T	[1987STE/MAL, 1966MOR/TAM]	
	[109-66-0]	pentane					
	$\Delta_{\text{fus}} H$		8.4	143.5		[1991ACR]	
	$\Delta_{\text{sub}} H$		42.0	143	B	[1963BON]	
	$\Delta_v H$	(308–423)	26.7	323		[2002PFO/RIE]	
	$\Delta_v H$		26.4	298		[1994RUZ/MAJ]	
	$\Delta_v H$	(223–352)	29.8	238	A	[1987STE/MAL]	
	$\Delta_v H$	(143–223)	32.3	208	A	[1987STE/MAL]	
	$\Delta_v H$	(350–422)	26.1	365	A	[1987STE/MAL]	
	$\Delta_v H$	(418–470)	26.2	433	A	[1987STE/MAL]	
	$\Delta_v H$		26.6 ± 0.1	298	C	[1982FUC/PEA]	
	$\Delta_v H$		26.4	298	C	[1981HOS/SCO2]	
	$\Delta_v H$		25.5	310		[1977DAS/REE]	
	$\Delta_v H$		23	350		[1977DAS/REE]	
	$\Delta_v H$		19.7	390		[1977DAS/REE]	
	$\Delta_v H$		15.1	430		[1977DAS/REE]	
	$\Delta_v H$	(216–296)	8.5	460		[1977DAS/REE]	
	$\Delta_v H$		26.2	298		[1975HOE/HOE]	
	$\Delta_v H$	(269–341)	27.9	284	EB	[1987STE/MAL, 1974OSB/DOU]	
	$\Delta_v H$		26.4	298		[1971WIL/ZWO]	
$\Delta_v H$		26.4	298	C	[1947OSB/GIN]		
$\Delta_v H$	(286–310)	27.4	298	MM	[1945WIL/TAY]		
$\Delta_v H$		26.2	298		[1940MES/KEN]		
C ₅ DH ₁₁	[55620-30-9]	1-deuteropentane					
$\Delta_v H$		(223–303)	26.2	298		[1975HOE/HOE]	
C ₅ DH ₁₁	[55620-31-0]	3-deuteropentane					
$\Delta_v H$		(213–294)	26.3	298		[1975HOE/HOE]	
C ₅ D ₁₂	[2031-90-5]	pentane-d ₁₂					
$\Delta_v H$		(205–298)	26.0	298		[1975HOE/HOE]	
C ₅ H ₁₂	[463-82-1]	2,2-dimethylpropane (neopentane)					
	$\Delta_{\text{trs}} H$		2.58	140			
	$\Delta_{\text{fus}} H$		3.26	256.5		[1996DOM/HEA]	
	$\Delta_{\text{sub}} H$	(223–256)	28.2	241		[1987STE/MAL]	
	$\Delta_{\text{sub}} H$		33.2			[1933WHI/FLE, 1936AST/MES]	
	$\Delta_{\text{sub}} H$	(171–249)	23.9	210	A	[1947STU]	
	$\Delta_v H$	(268–313)	24.0	283	A	[1987STE/MAL]	
	$\Delta_v H$	(312–385)	23.1	327	A	[1987STE/MAL]	
	$\Delta_v H$	(382–433)	23.1	397	A	[1987STE/MAL]	
	$\Delta_v H$		21.8	298	C	[1981HOS/SCO2]	
	$\Delta_v H$		22.2	290		[1977DAS/REE2]	
	$\Delta_v H$		19.5	330		[1977DAS/REE2]	
	$\Delta_v H$		16.2	370		[1977DAS/REE2]	
	$\Delta_v H$		11.1	410		[1977DAS/REE2]	
	$\Delta_v H$	(257–293)	24.3	272		[1975HOE/PAR, 1984BOU/FRI]	
	$\Delta_v H$	(343–433)	22.8	358		[1973DAW/SIL, 1984BOU/FRI]	
	$\Delta_v H$		21.85	298		[1971WIL/ZWO]	
	$\Delta_v H$		22.8 ± 0.1	283		[1936AST/MES]	
	C ₅ H ₁₂	[78-78-4]	2-methylbutane (isopentane)				
		$\Delta_{\text{fus}} H$		5.13	113.4		[1996DOM/HEA]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound					
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference	
		$\Delta_v H$	(255–323)	26.9	270		[1991EWI/GOO]
		$\Delta_v H$	(216–323)	28.5	231	A	[1987STE/MAL]
		$\Delta_v H$	(300–460)	25.2	315	A	[1987STE/MAL]
		$\Delta_v H$	(320–391)	25.2	335	A	[1987STE/MAL]
		$\Delta_v H$	(385–416)	24.8	400	A	[1987STE/MAL]
		$\Delta_v H$	(412–460)	25.3	427	A	[1987STE/MAL]
		$\Delta_v H$		24.4	310		[1977DAS/REE3]
		$\Delta_v H$		21.5	350		[1977DAS/REE3]
		$\Delta_v H$		18.0	390		[1977DAS/REE3]
		$\Delta_v H$		12.9	430		[1977DAS/REE3]
		$\Delta_v H$		24.8	298		[1971WIL/ZWO]
		$\Delta_v H$	(190–300)	30.2	205		[1947STU]
		$\Delta_v H$	(289–301)	26.2	295	MM	[1945WIL/TAY]
		$\Delta_v H$		25.0	298	C	[1942SCH/AST]
C₅H₁₂ClF₃N₂OS	[63265-73-6]	chlorobis(N-methylmethanaminato)oxo(trifluoromethyl)sulfur					
		$\Delta_v H$		40.2	477	I	[1977KIT/SHR2]
C₅H₁₂ClF₃N₂S	[63265-71-4]	chlorobis(N-methylmethanaminato)(trifluoromethyl) sulfur					
		$\Delta_v H$		38.1	368	I	[1977KIT/SHR2]
C₅H₁₂NO₃PS₂	[60-51-5]	phosphorodithioic acid, O,O-dimethyl-S-[2-(methylamino)-2-oxoethyl]ester					
		$\Delta_{\text{fus}}H$		20.49	321	DSC	[1990DON/DRE]
		$\Delta_v H$	(283–390)	95.0	298	A	[1987STE/MAL]
C₅H₁₂N₂	[4426-46-4]	methyl butyldiazene					
		$\Delta_v H$		36.4 ± 0.2	298	C	[1978ENG/MON]
C₅H₁₂N₂	[109-01-3]	N-methylpiperazine					
		$\Delta_v H$	(274–319)	46.7	289	A	[1987STE/MAL]
C₅H₁₂N₂O	[632-22-4]	1,1,3,3-tetramethylurea					
		$\Delta_{\text{fus}}H$		14.0	272.1		[2001JAM/DOB]
		$\Delta_{\text{fus}}H$		13.4	272.2		[1995KAB/KOZ2, 1996DOM/HEA]
		$\Delta_v H$	(320–450)	41.7	450	A,EB	[1987KNE/ZON]
		$\Delta_v H$	(320–450)	52.2	325	A,EB	[1987KNE/ZON]
C₅H₁₂N₂O	[634-95-7]	1,1-diethylurea					
		$\Delta_{\text{trs}}H$		2.07	197.3		
		$\Delta_{\text{fus}}H$		16.78	342.3		[1991ACR, 1995KAB/KOZ2, 1990KAB/MIR2]
		$\Delta_{\text{sub}}H$	(312–339)	95.7 ± 0.7	298	GS	[2006EME/KAB]
		$\Delta_{\text{sub}}H$	(305–347)	95.5 ± 0.8	324	ME	[2003ZAI/KAB]
		$\Delta_{\text{sub}}H$	(305–347)	94.9 ± 0.8	350	ME	[2003ZAI/KAB]
		$\Delta_{\text{sub}}H$		94.7 ± 0.2	350	C	[2003ZAI/KAB]
C₅H₁₂N₂O	[623-76-7]	1,3-diethylurea					
		$\Delta_{\text{trs}}H$		1.87	339.4		
		$\Delta_{\text{fus}}H$		12.46	383.4		[1991ACR, 1995KAB/KOZ2, 1990KAB/MIR2]
		$\Delta_{\text{sub}}H$	(343–379)	95.4 ± 0.3	298	GS	[2006EME/KAB]
		$\Delta_{\text{sub}}H$	(323–384)	91.8 ± 0.9	358	ME	[2003ZAI/KAB]
		$\Delta_{\text{sub}}H$	(323–384)	92.3 ± 0.9	350	ME	[2003ZAI/KAB]
		$\Delta_{\text{sub}}H$		95.6 ± 0.6	350	C	[2003ZAI/KAB]
		$\Delta_{\text{sub}}H$	(321–379)	96.8 ± 0.9	361	TE	[1990PIA/FER, 1987FER/DEL2]
		$\Delta_{\text{sub}}H$	(384–590)	NA		ME	[1986KRA/KOZ]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₅ H ₁₂ N ₂ O	[592-31-4]	N-butylurea				
	$\Delta_{\text{trs}}H$		6.3	310.5		
	$\Delta_{\text{trs}}H$		0.7	362.2		
	$\Delta_{\text{fus}}H$		10.8	365.4	DSC	[2005HAS/TAJ]
	$\Delta_{\text{trs}}H$		7.0	315		
	$\Delta_{\text{trs}}H$		1.0	346		
	$\Delta_{\text{fus}}H$		15.7	370	DSC	[1995FER/DEL]
	$\Delta_{\text{trs}}H$		7.02	313.1		
	$\Delta_{\text{trs}}H$		0.88	344.9		
	$\Delta_{\text{fus}}H$		14.55	369.3		[1991ACR]
	$\Delta_{\text{sub}}H$	(346–367)	105.8 ± 0.7	298	GS	[2006EME/KAB]
	$\Delta_{\text{sub}}H$	(339–364)	102.7 ± 2.8	354	ME	[2003ZAI/KAB]
	$\Delta_{\text{sub}}H$	(339–364)	103.0 ± 2.8	350	ME	[2003ZAI/KAB]
$\Delta_{\text{sub}}H$		101.1 ± 0.4	350	C	[2003ZAI/KAB]	
$\Delta_{\text{sub}}H$		99 ± 4			[1987FIO/FER]	
C ₅ H ₁₂ N ₂ O	[592-17-6]	N-isobutylurea				
	$\Delta_{\text{sub}}H$		101.1 ± 1.1	377	TE	[1990PIA/FER]
C ₅ H ₁₂ N ₂ O	[689-11-2]	N-sec-butylurea				
	$\Delta_{\text{sub}}H$	(345–394)	101.9 ± 0.5	298	GS	[2006EME/KAB]
	$\Delta_{\text{sub}}H$	(338–372)	104.3 ± 0.8	355	ME	[2003ZAI/KAB]
	$\Delta_{\text{sub}}H$	(338–372)	104.5 ± 0.8	350	ME	[2003ZAI/KAB]
$\Delta_{\text{sub}}H$		102.4 ± 0.5	350	C	[2009RIB/FER7]	
C ₅ H ₁₂ N ₂ O	[1118-12-3]	N-tert-butylurea				
	$\Delta_{\text{trs}}H$		0.1	249		
	$\Delta_{\text{fus}}H$		33.13	449.8		[1991ACR]
	$\Delta_{\text{sub}}H$	(335–397)	98.2 ± 0.4	298	GS	[2006EME/KAB]
	$\Delta_{\text{sub}}H$	(333–372)	97.6 ± 0.8	353	ME	[2003ZAI/KAB]
	$\Delta_{\text{sub}}H$	(333–372)	97.7 ± 0.8	350	ME	[2003ZAI/KAB]
	$\Delta_{\text{sub}}H$		94.4 ± 0.9	350	C	[2003ZAI/KAB]
$\Delta_{\text{sub}}H$		101.6 ± 0.7	379	TE	[1990PIA/FER]	
C ₅ H ₁₂ N ₂ O ₂	[52330-07-1]	N-methyl-N-nitrobutanamine				
$\Delta_{\text{fus}}H$			37.56	331		[1987OYU/BRI]
C ₅ H ₁₂ N ₂ S	[105-55-5]	1,3-diethylthiourea				
	$\Delta_{\text{fus}}H$		17.14	350.5	DSC	[2000DEL/JOZ]
	$\Delta_{\text{sub}}H$		121.7 ± 3	298	B	[2000DEL/JOZ]
	$\Delta_{\text{sub}}H$		120.2 ± 3.0	298	B	[1994TER/PIA]
Δ_vH	(351–384)	101 ± 3.0	368	ME,TE	[1994TER/PIA]	
C ₅ H ₁₂ N ₂ S	[2782-91-4]	tetramethylthiourea				
	$\Delta_{\text{fus}}H$		22.14	350.4	DSC	[2000DEL/JOZ]
	$\Delta_{\text{sub}}H$		84.5 ± 3	298	ME	[2000DEL/JOZ]
	$\Delta_{\text{sub}}H$		83 ± 3.0	333	TE	[1994FER/MAR]
	$\Delta_{\text{sub}}H$		84.0	298		[1994FER/MAR]
	$\Delta_{\text{sub}}H$		83.0 ± 0.5	298	C	[1985MUR/SAK]
$\Delta_{\text{sub}}H$		83.0 ± 0.2	298	C	[1982INA/MUR]	
C ₅ H ₁₂ O	[628-28-4]	1-methoxybutane				
	$\Delta_{\text{fus}}H$		10.85	157.5		[1996DOM/HEA]
Δ_vH	(293–367)		32.5	308	A	[1987STE/MAL]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		T _m (K)	Method	Reference	
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)				
		$\Delta_v H$		32.4	298	C	[1980MAJ/WAN]
		$\Delta_v H$	(265–367)	32.4	298		[1976AMB/ELL]
		$\Delta_v H$	(265–367)	29.6	343		[1976AMB/ELL]
		$\Delta_v H$		32.5 ± 0.1	298	C	[1975FEN/HAR]
		$\Delta_v H$	(296–342)	32.4	311	EB	[1969CID/POL]
C₅H₁₂O	[628-32-0]	1-ethoxypropane					
		$\Delta_{\text{fus}} H$		8.39	145.7		[1996DOM/HEA]
		$\Delta_v H$	(264–359)	33.0	279	A	[1987STE/MAL, 1976AMB/ELL]
		$\Delta_v H$		31.4	298	C	[1980MAJ/WAN]
		$\Delta_v H$	(264–359)	31.4	298		[1976AMB/ELL]
		$\Delta_v H$	(264–359)	29	336		[1976AMB/ELL]
		$\Delta_v H$		31.4 ± 0.1	298	C	[1975FEN/HAR]
		$\Delta_v H$	(293–335)	31.6	308		[1969CID/POL]
C₅H₁₂O	[625-54-7]	ethyl isopropyl ether					
		$\Delta_v H$		30.0	298	C	[1980MAJ/WAN]
C₅H₁₂O	[1634-04-4]	methyl <i>tert</i> -butyl ether					
		$\Delta_{\text{fus}} H$		7.6	164.6		[1996DOM/HEA]
		$\Delta_v H$	(300–328)	29.9	314		[2002SEG/GAL]
		$\Delta_v H$	(315–365)	29.6	330		[1998AUC/LOR]
		$\Delta_v H$	(298–322)	30.0	310		[1995BEL/AIT]
		$\Delta_v H$	(300–411)	31.2	315	EB	[1994KRA/GME]
		$\Delta_v H$	(287–326)	30.4	302		[1991WU/PIV]
		$\Delta_v H$		29.8	298	C	[1980MAJ/WAN]
		$\Delta_v H$	(287–351)	30.2	302	A	[1987STE/MAL, 1976AMB/ELL]
		$\Delta_v H$		29.6	298		[1976AMB/ELL]
		$\Delta_v H$		27.9	328		[1976AMB/ELL]
		$\Delta_v H$		30.4 ± 0.1	298	C	[1975FEN/HAR]
C₅H₁₂O	[71-41-0]	1-pentanol					
		$\Delta_{\text{fus}} H$		10.51	195.6		[2004VAN/VAN]
		$\Delta_{\text{fus}} H$		10.5	195.6		[1996DOM/HEA]
		$\Delta_v H$	(318–403)	55.4	298		[2006NAS/NEU]
		$\Delta_v H$		44.4	411		[2000WOR/JAM]
		$\Delta_v H$		40.1	448		[2000WOR/JAM]
		$\Delta_v H$		36.1	473		[2000WOR/JAM]
		$\Delta_v H$		31.7	498		[2000WOR/JAM]
		$\Delta_v H$		26.4	523		[2000WOR/JAM]
		$\Delta_v H$		22.0	548		[2000WOR/JAM]
		$\Delta_v H$		14.1	573		[2000WOR/JAM]
		$\Delta_v H$		7.1	586		[2000WOR/JAM]
		$\Delta_v H$		43.5			[1999FAT]
		$\Delta_v H$	(323–373)	57.8	298	CGC	[1995CHI/HOS]
		$\Delta_v H$	(323–373)	57.4	298	CGC	[1995CHI/HOS]
		$\Delta_v H$	(335–410)	51.5	350		[1994AUC/BUR]
		$\Delta_v H$	(388–420)	47.2	403	A	[1987STE/MAL]
		$\Delta_v H$	(326–411)	54.3	341	A	[1987STE/MAL]
		$\Delta_v H$	(408–441)	45.4	423	A	[1987STE/MAL]
		$\Delta_v H$		55.7 ± 0.2	313	C	[1985MAJ/SVO2]
		$\Delta_v H$		54.4 ± 0.2	328	C	[1985MAJ/SVO2]
		$\Delta_v H$		53.0 ± 0.2	343	C	[1985MAJ/SVO2]
		$\Delta_v H$		51.2 ± 0.2	358	C	[1985MAJ/SVO2]
		$\Delta_v H$	(343–303)	55.4	298		[1983SCH/STR]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
		(310–411)	55.0	325		[1973WIL/ZWO]
			50.5 ± 0.1	362	C	[1970COU/FEN]
			49.2 ± 0.1	374	C	[1970COU/FEN]
			47.0 ± 0.1	392	C	[1970COU/FEN]
			44.4 ± 0.1	411	C	[1970COU/FEN]
		(347–429)	51.6	362	EB	[1987STE/MAL, 1970AMB/SPR]
		(307–411)	56.2	322	DTA	[1969KEM/KRE]
			56.9 ± 0.2	298	C	[1966WAD]
			57.7 ± 1.1	298	EB	[1960GRE, 2001KUL/VER]
C₅H₁₂O	[6032-29-7]	2-pentanol				
			8.48	200		[1997LOH/JOH]
		(324–391)	51.2	339	EB	[2009GIE/KOS]
		(323–373)	53.6	298	CGC	[1995CHI/HOS]
		(274–393)	58.9	289	A	[1987STE/MAL]
			54.2 ± 0.2	298	C	[1985MAJ/SVO2]
			52.7 ± 0.2	313	C	[1985MAJ/SVO2]
			50.9 ± 0.2	328	C	[1985MAJ/SVO2]
			49.0 ± 0.2	343	C	[1985MAJ/SVO2]
			46.9 ± 0.1	358	C	[1985MAJ/SVO2]
			45.4 ± 0.1	368	C	[1985MAJ/SVO2]
		(322–393)	50.3	337		[1984SAC/MAR]
		(298–393)	54.0	313		[1973WIL/ZWO]
			53.0	298	C	[1963MCC/LAI]
		(298–383)	53.7	313		[1935BUT/RAM, 1984BOU/FRI]
C₅H₁₂O	[584-02-1]	3-pentanol				
			9.08	204.2		[1997LOH/JOH]
			53.2 ± 0.1	298	EB	[1988PES/SHV, 2001KUL/VER]
		(245–390)	59.9	260	A	[1987STE/MAL]
		(317–389)	49.6	332		[1984SAC/MAR]
		(279–318)	53.6	294		[1975CAB/CON2]
		(294–389)	50.2	319		[1973WIL/ZWO]
			52.9	298	C	[1963MCC/LAI]
C₅H₁₂O	[137-32-6]	2-methyl-1-butanol				
		(330–405)	51.2	345		[1994AUC/BUR]
		(338–402)	49.8	353	A	[1987STE/MAL]
		(317–403)	53.9	332	A	[1987STE/MAL]
		(249–319)	58.5	264	A	[1987STE/MAL, 1979THO/MEA]
		(307–403)	56.1	322		[1973WIL/ZWO]
			54.1	298	C	[1963MCC/LAI]
		(302–410)	43.4	317		[1957EAS/HAR, 1984BOU/FRI]
		(298–393)	56.7	313		[1984BOU/FRI, 1935BUT/RAM]
C₅H₁₂O	[75-85-4]	2-methyl-2-butanol				
			0.93	146.4		
			1.54	149.9		
			0.66	214.4		
			2.24	262.7	AC	[2008TON/TAN]
		(84–301)	0.9	145.8		
		(84–301)	2.0	264.7	AC	[2007STR/RUZ2]
			1.96	146		
			0.17	213		
			4.46	264		[1996DOM/HEA, 1933PAR/HUF]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		T _m (K)	Method	Reference	
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)				
		$\Delta_v H$	(303–373)	51.2	318	EB	[2009GIE/KOS]
		$\Delta_v H$	(274–306)	51.5 ± 0.3	298	GS	[2001KUL/VER]
		$\Delta_v H$	(323–373)	50.5	298	CGC	[1995CHI/HOS]
		$\Delta_v H$	(308–375)	47.3	323		[1994AUC/BUR]
		$\Delta_v H$		50.2 ± 0.3	298	EB	[1988PES/SHV, 2001KUL/VER]
		$\Delta_v H$	(280–375)	49.0	295	A	[1987STE/MAL]
		$\Delta_v H$	(323–376)	45.8	338	A	[1987STE/MAL]
		$\Delta_v H$		50.1 ± 0.2	298	C	[1985MAJ/SVO2]
		$\Delta_v H$		48.4 ± 0.2	313	C	[1985MAJ/SVO2]
		$\Delta_v H$		46.4 ± 0.2	328	C	[1985MAJ/SVO2]
		$\Delta_v H$		44.2 ± 0.1	343	C	[1985MAJ/SVO2]
		$\Delta_v H$		42.0 ± 0.1	358	C	[1985MAJ/SVO2]
		$\Delta_v H$		40.3 ± 0.1	368	C	[1985MAJ/SVO2]
		$\Delta_v H$	(298–375)	52.8	313		[1973WIL/ZWO]
		$\Delta_v H$		49.2	298	C	[1963MCC/LAI]
		$\Delta_v H$	(298–364)	48.5	313		[1935BUT/RAM]
C₅H₁₂O	[123-51-3]		3-methyl-1-butanol				
		$\Delta_v H$	(348–404)	49.8	363		[2008LLA/MON]
		$\Delta_v H$	(323–373)	55.3	298	CGC	[1995CHI/HOS]
		$\Delta_v H$	(325–385)	47.2	340		[1994AUC/BUR]
		$\Delta_v H$	(303–412)	57.1	318		[1987STE/MAL]
		$\Delta_v H$		55.2 ± 0.2	303	C	[1985MAJ/SVO2]
		$\Delta_v H$		54.2 ± 0.2	313	C	[1985MAJ/SVO2]
		$\Delta_v H$		52.9 ± 0.2	328	C	[1985MAJ/SVO2]
		$\Delta_v H$		51.4 ± 0.2	343	C	[1985MAJ/SVO2]
		$\Delta_v H$		49.7 ± 0.2	358	C	[1985MAJ/SVO2]
		$\Delta_v H$	(298–426)	56.5	313		[1973WIL/ZWO]
		$\Delta_v H$		54.3	298	C	[1963MCC/LAI]
C₅H₁₂O	[598-75-4]		3-methyl-2-butanol				
		$\Delta_v H$	(280–301)	51.6 ± 0.3	298	GS	[2001KUL/VER]
		$\Delta_v H$	(280–375)	49.0	295	A	[1987STE/MAL]
		$\Delta_v H$	(298–384)	52.7	313		[1973WIL/ZWO]
		$\Delta_v H$		51.7	298	C	[1963MCC/LAI]
C₅H₁₂O	[598-75-4]		(dl) 3-methyl-2-butanol				
		$\Delta_v H$	(293–385)	46.4	308	A	[1987STE/MAL]
C₅H₁₂O	[75-84-3]		2,2-dimethyl-1-propanol				
		$\Delta_{\text{trs}}H$	(10–370)	4.1	233.3		
		$\Delta_{\text{fus}}H$	(10–370)	2.9	328.2	AC	[2007STR/RUZ2]
		$\Delta_{\text{fus}}H$		3.87	329.8		[2003CEN/RUZ]
		$\Delta_{\text{trs}}H$		4.14	235.4		
		$\Delta_{\text{fus}}H$		3.73	329.8		[1999SAL/LOP]
		$\Delta_{\text{trs}}H$		4.6	242.1		
		$\Delta_{\text{fus}}H$		3.5	328.1	DSC	[1996GRA]
		$\Delta_{\text{trs}}H$		4.46	242		
		$\Delta_{\text{fus}}H$		4.06	325		[1970MER/BRE]
		$\Delta_v H$	(274–312)	51.8 ± 0.3	298	GS,B	[2001KUL/VER]
		$\Delta_v H$	(330–387)	47.5	345	A	[1987STE/MAL]
C₅H₁₂O₂	[5137-45-1]		1-ethoxy-2-methoxyethane				
		$\Delta_v H$		39.8 ± 0.1	298	C	[1970KUS/WAD]
C₅H₁₂O₂	[77-76-9]		2,2-dimethoxypropane				
		$\Delta_v H$	(272–301)	37.6 ± 0.4	298	GS	[2002VER]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound					
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference	
		$\Delta_v H$	(272–301)	38.2 ± 0.4		GS	[1998VER/PEN]
		$\Delta_v H$	(299–348)	35.3	324	EB	[1994WIB/MOR]
		$\Delta_v H$	(292–357)	33.4 ± 0.2	325		[1988BAG/GUR]
C ₅ H ₁₂ O ₂	[109-59-1]	2-isopropoxyethanol					
		$\Delta_v H$		50.1 ± 0.1	298	C	[1971MOR]
		$\Delta_v H$	(341–413)	45.1	356	A	[1987STE/MAL, 1957DYK/SEP, 1972DYK]
C ₅ H ₁₂ O ₂	[2807-30-9]	2-propoxyethanol					
		$\Delta_v H$		52.1 ± 0.1	298	C	[1971KUS/WAD]
		$\Delta_v H$	(350–422)	46.3	365	A	[1987STE/MAL, 1957DYK/SEP, 1972DYK]
C ₅ H ₁₂ O ₂	[462-95-3]	formaldehyde diethyl acetal (diethoxymethane)					
		$\Delta_v H$	(273–361)	36.1	288	A	[1987STE/MAL]
		$\Delta_v H$		35.7 ± 0.2	298	C	[1969MAN]
C ₅ H ₁₂ O ₂	[684-84-4]	2-methyl-1,3-butanediol					
		$\Delta_v H$	(399–561)	62.4	414	A	[1987STE/MAL]
C ₅ H ₁₂ O ₂	[2568-33-4]	3-methyl-1,3-butanediol					
		$\Delta_v H$	(346–475)	60.3	361	A	[1987STE/MAL]
C ₅ H ₁₂ O ₂	[126-30-7]	2,2-dimethyl-1,3-propanediol					
		$\Delta_{\text{trs}}H$		12.33	315.1		
		$\Delta_{\text{fus}}H$		4.55	401.2	DSC	[2009SIN/MUR]
		$\Delta_{\text{trs}}H$	(78–410)	14.78	314.3		
		$\Delta_{\text{fus}}H$	(78–410)	7.52	402.4	AC	[2007TON/TAN]
		$\Delta_{\text{trs}}H$		12.24	315.1		
		$\Delta_{\text{fus}}H$		4.23	401.6	DSC	[2006DIV/CHE]
		$\Delta_{\text{trs}}H$	(15–340)	0.18	60.4		
		$\Delta_{\text{trs}}H$	(15–340)	12.5	314.5	AC	[2001KAM/SUE]
		$\Delta_{\text{trs}}H$		0.18	60.4		
		$\Delta_{\text{trs}}H$		12.43	314.4		
		$\Delta_{\text{fus}}H$		4.34	402.8	DSC	[1999SAL/LOP]
		$\Delta_{\text{trs}}H$		13.8	315.2		
		$\Delta_{\text{trs}}H$		12.52	314.5	AC	[1999SUG]
		$\Delta_{\text{trs}}H$		12.8	315.2		
		$\Delta_{\text{fus}}H$		4.3	402.5	DSC	[1996GRA]
		$\Delta_{\text{fus}}H$		4.6	403.2		[1973FRA/KRZ, 1994LOP/VAN]
		$\Delta_{\text{sub}}H$ (cryst)	(294–311)	85 ± 2			[1995FON/MUN]
		$\Delta_{\text{sub}}H$ (plastic)	(319–333)	75 ± 2			[1995FON/MUN]
		$\Delta_{\text{sub}}H$ (plastic)		75.5 ± 3.8	368	C	[1994FON/MUN2, 1994FON/MUN]
		$\Delta_{\text{sub}}H$ (cryst)		87.6 ± 4.4	350	C	[1994FON/MUN2, 1994FON/MUN]
		$\Delta_v H$	(400–480)	79.4	415	A	[1987STE/MAL]
C ₅ H ₁₂ O ₂	[5343-92-0]	1,2-pentanediol					
		$\Delta_v H$	(289–345)	74.6 ± 0.3	298	GS	[2004VER2]
C ₅ H ₁₂ O ₂	[111-29-5]	1,5-pentanediol					
		$\Delta_{\text{fus}}H$		15.72	248		[1996DOM/HEA]
		$\Delta_v H$		86.8 ± 0.5	298	C	[1988KNA/SAB, 1990KNA/SAB2]
		$\Delta_v H$	(391–479)	78.6	406	A	[1987STE/MAL]
C ₅ H ₁₂ O ₂	[625-69-4]	2,4-pentanediol					
		$\Delta_v H$	(297–347)	72.5 ± 0.3	298	GS	[2007VER]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound					
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference	
C ₅ H ₁₂ O ₂ S	[14094-12-3]	<i>tert</i> -butyl methyl sulfone					
	$\Delta_{\text{fus}}H$		24.69	357.6		[1961BUS/IVI]	
			$\Delta_{\text{sub}}H$	82.4 ± 2.5		[UR/MAC, 1970COX/PIL]	
C ₅ H ₁₂ O ₃	[17742-78-8]	<i>tert</i> -butyldioxyethanol					
	Δ_vH		59.6 ± 2.4			[1983VAN/KAC]	
C ₅ H ₁₂ O ₃	[111-77-3]	diethylene glycol, methyl ether					
	Δ_vH	(385–466)	51.9	400	A	[1987STE/MAL, 1957DYK/SEP, 1972DYK]	
C ₅ H ₁₂ O ₃	[14642-48-9]	2,3,4-pentanetriol					
	Δ_vH	(428–600)	78.9	443		[1947STU]	
C ₅ H ₁₂ O ₃	[77-85-0]	2-hydroxymethyl-2-methyl-1,3-propanediol					
	$\Delta_{\text{trs}}H$		20.94	356.7			
	$\Delta_{\text{fus}}H$		4.72	474.4		[1999SAL/LOP]	
	$\Delta_{\text{trs}}H$		23.17	354			
	$\Delta_{\text{fus}}H$		5.38	470		[1996DOM/HEA]	
	$\Delta_{\text{trs}}H$		21.1	357.4			
	$\Delta_{\text{fus}}H$		4.8	472.4	DSC	[1996GRA]	
	$\Delta_{\text{trs}}H$	(18–375)	21.2	358.2	AC		
	$\Delta_{\text{fus}}H$	(353–483)	4.7	473.7	DSC	[1990SUE/MAT]	
	$\Delta_{\text{sub}}H$ (plastic)		84.2 ± 4.2	319	C	[1994FON/MUN]	
$\Delta_{\text{sub}}H$ (cryst)		109.2 ± 5.5	311	C	[1994FON/MUN]		
C ₅ H ₁₂ O ₄	[1850-14-2]	tetramethoxymethane					
	Δ_vH	(304–387)	41.2	319		[1980THO/SMI]	
C ₅ H ₁₂ O ₄	[115-77-5]	pentaerythritol					
	$\Delta_{\text{trs}}H$		43.93	460.4			
	$\Delta_{\text{fus}}H$		7.11	538.7		[1996DOM/HEA]	
	$\Delta_{\text{trs}}H$		41.2	459.7			
	$\Delta_{\text{fus}}H$		5.3	532.3	DSC	[1996GRA]	
	$\Delta_{\text{trs}}H$		40.5	458.3		[1990BAR/DEL]	
	$\Delta_{\text{fus}}H$		4.6	513.2	DSC	[1990BAR/DEL]	
	$\Delta_{\text{sub}}H$ (cryst)	(441–460)	134 ± 7			[1995FON/MUN]	
	$\Delta_{\text{sub}}H$ (plastic)	(465–477)	96 ± 9			[1995FON/MUN]	
	$\Delta_{\text{sub}}H$		131.3 ± 6.6	403	C	[1994FON/MUN]	
	$\Delta_{\text{sub}}H$	(418–455)	161 ± 1.0	437	TE	[1990BAR/DEL]	
	$\Delta_{\text{sub}}H$		163.0	298		[1990BAR/DEL]	
	$\Delta_{\text{sub}}H$ (tetragonal)	(397–410)	131.4		ME	[1951NIT/SEK2, 1960JON]	
$\Delta_{\text{sub}}H$	(379–408)	143.9 ± 0.8		ME	[1953BRA/COT, 1960JON]		
C ₅ H ₁₂ O ₅	[488-81-3]	1,2,3,4,5-pentahydroxypentane (adonitol)					
	$\Delta_{\text{fus}}H$		36.42	369.1	AC	[2010TON/YU]	
	$\Delta_{\text{fus}}H$		35.5	375.0	DSC	[2003CAR/DES]	
	$\Delta_{\text{fus}}H$		37.6	374.7	DSC	[1990BAR/DEL, 1996DOM/HEA]	
	$\Delta_{\text{sub}}H$		161.0	298	B	[1990BAR/DEL]	
	Δ_vH	(418–465)	111.1 ± 1.5	443	TE	[1990BAR/DEL]	
C ₅ H ₁₂ O ₅	[488-82-4]	1,2,3,4,5-pentahydroxypentane (D-arabinitol)					
	$\Delta_{\text{fus}}H$		38.8	376.0	DSC	[2003CAR/DES]	
	$\Delta_{\text{fus}}H$		38.9	379.4	DSC	[1990BAR/DEL, 1990DOM/HEA]	
			$\Delta_{\text{sub}}H$	160.0	298	B	[1990BAR/DEL]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound					
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference	
		$\Delta_v H$	(414–461)	110.1 ± 1.5	440	TE	[1990BAR/DEL]
C ₅ H ₁₂ O ₅	[7643-75-6]	1,2,3,4,5-pentahydroxypentane (L-arabinitol)					
	$\Delta_{\text{fus}} H$		43.2	374.0		DSC	[2003CAR/DES]
C ₅ H ₁₂ O ₅	[87-99-0]	xylitol					
	$\Delta_{\text{fus}} H$		33.26	369			[2007TON/TAN2]
	$\Delta_{\text{fus}} H$		37.7	368.0		DSC	[2003CAR/DES]
	$\Delta_{\text{fus}} H$		37.4	365.7		DSC	[1996DOM/HEA, 1990BAR/DEL]
	$\Delta_{\text{sub}} H$		161	298		B	[1990BAR/DEL]
		$\Delta_v H$	(406–460)	111.1 ± 0.8	433	TE	[1990BAR/DEL]
C ₅ H ₁₂ S	[10359-64-5]	3-methyl-2-thiapentane					
	$\Delta_v H$	(288–418)	38.5	303			[1999DYK/SVO]
C ₅ H ₁₂ S	[5008-69-5]	4-methyl-2-thiapentane					
	$\Delta_v H$	(288–411)	36.9	303			[1999DYK/SVO]
C ₅ H ₁₂ S	[628-29-5]	butyl methyl sulfide					
	$\Delta_{\text{fus}} H$		12.45	175.6			[1985DEA]
	$\Delta_v H$	(297–423)	40.4	312			[1999DYK/SVO]
	$\Delta_v H$	(301–330)	35.3	315		A	[1987STE/MAL]
	$\Delta_v H$		40.5	298			[1981SHI/SAI]
	$\Delta_v H$		41.0	298			[1971WIL/ZWO]
	$\Delta_v H$		40.9 ± 0.8	298		GC	[1964GUB/FER]
	$\Delta_v H$	(296–325)	38.1	313		EB	[1962MAC/MAY2]
	$\Delta_v H$	(343–436)	38.0	358		A,EB	[1987STE/MAL, 1961MCC/FIN, 1966OSB/DOU]
C ₅ H ₁₂ S	[6163-64-0]	methyl <i>tert</i> -butyl sulfide					
	$\Delta_{\text{fus}} H$		8.41	190.8			[1996DOM/HEA]
	$\Delta_v H$	(245–352)	34.2	298			[2004SAW/MOK]
	$\Delta_v H$	(276–397)	36.5	291			[1999DYK/SVO]
	$\Delta_v H$		35.9	298			[1971WIL/ZWO]
	$\Delta_v H$	(305–411)	35.1	320		A,EB	[1987STE/MAL, 1962SCO/GOO, 1966OSB/DOU]
C ₅ H ₁₂ S	[5145-99-3]	ethyl isopropyl sulfide					
	$\Delta_v H$	(284–406)	38.1	299			[1999DYK/SVO]
	$\Delta_v H$		37.8	298			[1981SHI/SAI]
	$\Delta_v H$		38.5	298			[1971WIL/ZWO]
	$\Delta_v H$		37.9 ± 0.8	298		GC	[1964MAC/MCC]
	$\Delta_v H$	(296–325)	38.1	313		EB	[1962MAC/MAY2]
	$\Delta_v H$	(319–391)	36.3	334		A,EB	[1987STE/MAL, 1952WHI/BER]
C ₅ H ₁₂ S	[4110-50-3]	ethyl propyl sulfide					
	$\Delta_{\text{fus}} H$		10.58	156.1			[1996DOM/HEA]
	$\Delta_v H$	(293–418)	39.8	308			[1999DYK/SVO]
	$\Delta_v H$		40	298			[1981SHI/SAI]
	$\Delta_v H$		39.5	298		C	[1981HOS/SCO]
	$\Delta_v H$	(331–398)	37.8	346		A,EB	[1987STE/MAL, 1952WHI/BER]
C ₅ H ₁₂ S	[110-66-7]	1-pentanethiol					
	$\Delta_{\text{fus}} H$		17.53	197.5			[1996DOM/HEA]
		$\Delta_v H$	(300–426)	40.6	315		[1999DYK/SVO]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		T _m (K)	Method	Reference
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)			
		$\Delta_{\text{v}}H$	41.1	298		[1971WIL/ZWO]
		$\Delta_{\text{v}}H$	37.1 ± 0.1	356	C	[1965FIN/HOS]
		$\Delta_{\text{v}}H$	36.4 ± 0.1	376	C	[1965FIN/HOS]
		$\Delta_{\text{v}}H$	34.9 ± 0.1	400	C	[1965FIN/HOS]
		$\Delta_{\text{v}}H$	(347–440)	38.1	A,EB	[1987STE/MAL, 1952FIN/SCO, 1966OSB/DOU]
C₅H₁₂S	[2084-19-7]	2-pentanethiol				
		$\Delta_{\text{v}}H$	(287–412)	38.4		[1999DYK/SVO]
		$\Delta_{\text{v}}H$	(347–435)	37.8	A	[1987STE/MAL]
C₅H₁₂S	[616-31-9]	3-pentanethiol				
		$\Delta_{\text{v}}H$	(288–413)	38.3		[1999DYK/SVO]
C₅H₁₂S	[1878-18-8]	2-methyl-1-butaneethiol				
		$\Delta_{\text{v}}H$	(293–418)	39.2		[1999DYK/SVO]
		$\Delta_{\text{v}}H$		39.9 ± 0.1		[1972GOO, 1966OSB/DOU]
		$\Delta_{\text{v}}H$		39.7		[1971WIL/ZWO]
		$\Delta_{\text{v}}H$	(324–432)	37.6	A,EB	[1987STE/MAL, 1966OSB/DOU]
C₅H₁₂S	[541-31-1]	3-methyl-1-butaneethiol				
		$\Delta_{\text{fus}}H$		7.41		[1996DOM/HEA]
		$\Delta_{\text{v}}H$	(292–418)	39.3		[1999DYK/SVO]
		$\Delta_{\text{v}}H$		39.7		[1971WIL/ZWO]
		$\Delta_{\text{v}}H$		39.9 ± 0.1		[1972GOO, 1966OSB/DOU]
		$\Delta_{\text{v}}H$	(323–431)	37.7	A,EB	[1987STE/MAL, 1966OSB/DOU]
C₅H₁₂S	[1679-09-0]	2-methyl-2-butaneethiol				
		$\Delta_{\text{v}}H$	(276–398)	36.3		[1999DYK/SVO]
		$\Delta_{\text{v}}H$		35.6		[1971WIL/ZWO]
		$\Delta_{\text{v}}H$	(320–411)	34.3	A,EB	[1987STE/MAL, 1962SCO/DOU]
		$\Delta_{\text{v}}H$		33.8 ± 0.1	C	[1962SCO/DOU]
		$\Delta_{\text{v}}H$		32.7 ± 0.1	C	[1962SCO/DOU]
		$\Delta_{\text{v}}H$		31.4 ± 0.1	C	[1962SCO/DOU]
C₅H₁₂S	[2084-18-6]	3-methyl-2-butaneethiol				
		$\Delta_{\text{trs}}H$		7.06		
		$\Delta_{\text{fus}}H$	(12–370)	0.61		[1996DOM/HEA, 1974MES/FIN]
		$\Delta_{\text{v}}H$	(285–409)	37.7		[1999DYK/SVO]
		$\Delta_{\text{v}}H$		37.5 ± 0.1		[1972GOO, 1966OSB/DOU]
		$\Delta_{\text{v}}H$		37.7		[1971WIL/ZWO]
		$\Delta_{\text{v}}H$	(315–422)	36.2	A,EB	[1987STE/MAL, 1966OSB/DOU]
C₅H₁₂S	[1679-08-9]	2,2-dimethyl-1-propaneethiol				
		$\Delta_{\text{v}}H$	(280–403)	36.9		[1999DYK/SVO]
		$\Delta_{\text{v}}H$		36.4 ± 0.1		[1972GOO, 1966OSB/DOU]
		$\Delta_{\text{v}}H$		36.8		[1971WIL/ZWO]
		$\Delta_{\text{v}}H$	(292–416)	36.2	A,EB	[1987STE/MAL, 1966OSB/DOU]
		$\Delta_{\text{v}}H$	(213–415)	42.1	EB,IP	[1966OSB/DOU]
C₅H₁₂S₂	[928-98-3]	1,5-pentanedithiol				
		$\Delta_{\text{v}}H$	(363–491)	51.6	A	[1987STE/MAL, 1999DYK/SVO]
		$\Delta_{\text{v}}H$		59.3		[1962MAN/SUN]
C₅H₁₂S₂	[5395-75-5]	3,5-dithiaheptane				
		$\Delta_{\text{v}}H$		50.8 ± 0.2	C	[1974MAN4]
C₅H₁₂S₂	[53966-36-2]	ethyl isopropyl disulfide				

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound						
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference		
		$\Delta_v H$	(369–426)	42.5	384		[1999DYK/SVO]	
		$\Delta_v H$	(363–427)	42.9	378	A,EB	[1987STE/MAL, 1952WHI/BER]	
C ₅ H ₁₂ S ₂	[30453-31-7]	ethyl propyl disulfide						
		$\Delta_v H$	(373–414)	44.0	388	A,EB	[1987STE/MAL, 1952WHI/BER, 1999DYK/SVO]	
C ₅ H ₁₂ S ₄	[6156-25-8]	tetra(methylthia)methane						
		$\Delta_{\text{trs}} H$	(13–360)	7.09	296.2			
		$\Delta_{\text{trs}} H$	(13–360)	7.29	318.8			
		$\Delta_{\text{fus}} H$	(13–360)	3.31	338.9	AC	[1998SOR/KIM]	
		$\Delta_{\text{trs}} H$		6.11	296.4			
		$\Delta_{\text{trs}} H$		7.61	318.7			
		$\Delta_{\text{fus}} H$		4.14	338.7		[1996DOM/HEA]	
C ₅ H ₁₃ N	[19961-27-4]	N-ethylisopropylamine						
		$\Delta_v H$		33.1 ± 0.1	298	C	[1979PET/MAJ]	
		$\Delta_v H$		32.1 ± 0.1	313	C	[1979PET/MAJ]	
		$\Delta_v H$		31.0 ± 0.1	328	C	[1979PET/MAJ]	
		$\Delta_v H$		28.8 ± 0.1	358	C	[1979PET/MAJ]	
		$\Delta_v H$	(303–342)	33.4	318	EB	[1979PET/MAJ]	
C ₅ H ₁₃ N	[616-39-7]	N,N-diethylmethylamine						
		$\Delta_v H$	(283–339)	31.8	298	A	[1987STE/MAL]	
C ₅ H ₁₃ N	[110-68-9]	N-methylbutylamine						
		$\Delta_v H$	(283–313)	38.1	298	A	[1987STE/MAL]	
C ₅ H ₁₃ N	[14610-37-8]	<i>tert</i> -butylmethylamine						
		$\Delta_v H$	(270–288)	32.3 ± 1.4	297		[1997VER]	
C ₅ H ₁₃ N	[110-58-7]	pentylamine						
		$\Delta_v H$	(322–378)	40.9	298	EB	[2004ANT/GAL]	
		$\Delta_v H$	(298–417)	39.0	313	A	[1987STE/MAL, 1972DYK]	
		$\Delta_v H$		40.1 ± 0.1	298	C	[1969WAD]	
C ₅ H ₁₃ NO	[110-73-6]	2-(ethylamino)ethanol						
		$\Delta_v H$	(282–321)	61.0 ± 0.4	298	GS	[2005KAP/SLO]	
C ₅ H ₁₃ NO ₂	[105-59-9]	N-methyl diethanolamine						
		$\Delta_v H$	(409–435)	71.5	422	EB	[2008KIM/SVE]	
		$\Delta_v H$	(390–520)	73.0	405	A	[1987STE/MAL]	
C ₅ H ₁₃ NO ₂ S	[2374-61-0]	N,N-diethyl methanesulfonamide						
		$\Delta_v H$	(384–528)	52.1	399	A	[1987STE/MAL]	
C ₅ H ₁₃ NS	[na]	N-methyl- <i>tert</i> -butylsulfenamide						
		$\Delta_v H$	(329–397)	41.9	364		[1999DYK/SVO]	
C ₅ H ₁₃ N ₃	[80-70-6]	1,1,3,3-tetramethylguanidine						
		$\Delta_v H$		46.9			[1967AND/HAM]	
C ₅ H ₁₃ O ₃ P	[683-08-9]	diethyl methylphosphonate						
		$\Delta_v H$	(253–465)	60.6	253	GS	[2009BUT/BUC]	
		$\Delta_v H$	(253–465)	57.2	283	GS	[2009BUT/BUC]	
		$\Delta_v H$	(253–465)	55.9	298	GS	[2009BUT/BUC]	
		$\Delta_v H$	(253–465)	54.7	313	GS	[2009BUT/BUC]	
		$\Delta_v H$	(253–465)	53.4	333	GS	[2009BUT/BUC]	
		$\Delta_v H$	(253–465)	51.2	373	GS	[2009BUT/BUC]	
		$\Delta_v H$	(343–402)	51.8	358	A	[1987STE/MAL, 1972DYK]	

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$		56.5 ± 4.2			[1956NEA/WIL, 1982PIL/SKI]
C ₅ H ₁₄ NP	[na]	trimethylphosphine-N-ethylimine				
	$\Delta_v H$		61.5 ± 4.2			[1960CLA/FOW, 1982PIL/SKI]
C ₅ H ₁₄ N ₂	[111-33-1]	N,N-dimethyl-1,3-propanediamine				
	$\Delta_{\text{fus}} H$		12.38	194.4		[1996DOM/HEA]
	$\Delta_v H$	(303–366)	45.7	318	A	[1987STE/MAL]
	$\Delta_v H$	(303–408)	42.0	318	A	[1987STE/MAL]
C ₅ H ₁₄ N ₂	[51-80-9]	<i>bis</i> (dimethylamino)methane				
	$\Delta_v H$	(273–348)	32.3	310		[1965AYL/PET]
	$\Delta_v H$		52.7			[1977VAS/KOT]
C ₅ H ₁₄ N ₂	[462-94-2]	pentane-1,5-diamine				
	$\Delta_{\text{fus}} H$		29.82	285	DSC	[2002DAL/DEL]
C ₅ H ₁₄ N ₂	[7328-91-8]	2,2-dimethyl-1,3-diaminopropane				
	$\Delta_{\text{trs}} H$		14.7	194.2		
	$\Delta_{\text{fus}} H$		1.7	301.7		[1996STR/BRA]
C ₆ BrF ₅	[344-04-7]	bromopentafluorobenzene				
	$\Delta_v H$	(400–522)	38.2	415	A	[1987STE/MAL, 1972DYK]
	$\Delta_v H$	(283–348)	43.1 ± 0.2	298		[1977KRE/PRI]
	$\Delta_v H$	(414–522)	38.0	429	EB	[1969WOO/ADI]
C ₆ BrF ₁₅ N ₂ S	[62977-74-6]	<i>bis</i> [1,2,2,2-tetrafluoro-1-(trifluoromethyl)ethyl]diimidodisulfurylbromide fluoride				
	$\Delta_v H$		41.0	476	I	[1977KIT/SHR2]
C ₆ Br ₆	[87-82-1]	hexabromobenzene				
	$\Delta_{\text{fus}} H$		24.6	598.8		[2004KUR/MAE2]
C ₆ ClF ₁₅ N ₂ S	[62977-72-4]	<i>bis</i> [1,2,2,2-tetrafluoro-1-(trifluoromethyl)ethyl]diimidodisulfurylchloride fluoride				
	$\Delta_v H$		37.2	458	I	[1977KIT/SHR2]
C ₆ ClF ₅	[344-07-0]	chloropentafluorobenzene				
	$\Delta_{\text{trs}} H$		3.64	191		
	$\Delta_{\text{trs}} H$		0.98	245		
	$\Delta_{\text{fus}} H$		8.36	257.5		[1996DOM/HEA, 1968AND/COU]
	$\Delta_v H$	(290–550)	41.3	298		[1991BAS/SVO]
	$\Delta_v H$	(348–402)	37.7	363	A	[1987STE/MAL]
	$\Delta_v H$	(307–417)	40.0	322	A	[1987STE/MAL, 1968AMB]
	$\Delta_v H$		37.7 ± 0.1	349		[1968AND/COU]
	$\Delta_v H$		36.4 ± 0.1	369		[1968AND/COU]
	$\Delta_v H$		34.8 ± 0.1	391		[1968AND/COU]
	$\Delta_v H$	(403–547)	35.2	418	EB	[1966EVA/TIL]
C ₆ ClF ₁₃ N ₂	[33757-14-1]	1-chloro-1', 2, 2, 2, 2', 2', 2-heptafluoro-1, 1'- <i>bis</i> (trifluoromethyl)azoethane				
	$\Delta_v H$	(297–355)	33.3	312	A	[1987STE/MAL, 1971SWI/ZAB]
C ₆ ClF ₁₄ P	[756-17-2]	<i>bis</i> (heptafluoropropyl) chlorophosphine				
	$\Delta_v H$	(283–373)	37.5	328		[1959EME/SMI]
C ₆ Cl ₂ F ₁₂ N ₂ S	[na]	<i>bis</i> (2-chlorohexafluoroisopropylimino) sulfur				
	$\Delta_v H$		43.5	404	I	[1972MET/SHR]
C ₆ Cl ₃ F ₃	[319-88-0]	1,3,5-trichloro-2,4,6-trifluorobenzene				
	$\Delta_{\text{fus}} H$		19.83	335		[1991ACR]
	$\Delta_v H$	(364–496)	49.2	379	A	[1987STE/MAL]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(364–550)	53.8	298		[1984BOU/FRI, 1991BAS/SVO]
C ₆ Cl ₃ F ₁₄ P	[na]	trichloro bis(heptafluoropropyl)phosphorane				
	$\Delta_v H$	(323–393)	40.1	358		[1959EME/SMI]
C ₆ Cl ₃ N ₃ O ₆	[2631-68-7]	1,3,5-trichloro-2,4,6-trinitrobenzene				
	$\Delta_v H$	(503–543)	68.9	518	A	[1987STE/MAL, 1968MAK]
	$\Delta_v H$	(503–543)	43.2	518		[1972DYK]
C ₆ Cl ₄ O ₂	[118-75-2]	2,3,5,6-tetrachloro-1,4-benzoquinone (chloranil)				
	$\Delta_{\text{fus}} H$		30.87	567.2		[1991ACR]
	$\Delta_{\text{sub}} H$	(333–356)	98.7 ± 8.3		QF	[1927COO/COO, 1960JON, 1970COX/PIL]
	$\Delta_v H$	(343–435)	88.5	358		[1947STU]
C ₆ Cl ₅ NO ₂	[82-68-8]	pentachloronitrobenzene				
	$\Delta_{\text{fus}} H$		18.41	418		[1991ACR]
	$\Delta_{\text{sub}} H$		96.3 ± 2.1	298	C	[2009RIB/FER6]
	$\Delta_{\text{sub}} H$	(317–339)	93.0 ± 0.4	328	ME	[2009RIB/FER6]
	$\Delta_{\text{sub}} H$	(317–339)	94.5 ± 0.4	298	ME	[2009RIB/FER6]
C ₆ Cl ₆	[118-74-1]	hexachlorobenzene				
	$\Delta_{\text{fus}} H$		25.2	502	DTA	[1991SAB/AN2]
	$\Delta_{\text{fus}} H$		23.85	505		[1991ACR]
	$\Delta_{\text{sub}} H$	(358–403)	96.8 ± 0.5	298	GS	[2007VER/EME]
	$\Delta_{\text{sub}} H$	(258–313)	105			[1994LIU/DIC]
	$\Delta_{\text{sub}} H$	(253–303)	77.4 ± 0.8	278	GS	[1994WAN/SHU]
	$\Delta_{\text{sub}} H$		89.6 ± 0.2	337	C	[1991SAB/AN2]
	$\Delta_{\text{sub}} H$		90.5 ± 0.2	298	C	[1991SAB/AN2]
	$\Delta_{\text{sub}} H$	(461–506)	85.5			[1989LUB/JAN]
	$\Delta_{\text{sub}} H$	(387–502)	62.7	402	A	[1987STE/MAL]
	$\Delta_{\text{sub}} H$	(314–373)	94.7	344	GS	[1986ROR/SAR, 1997DEL]
	$\Delta_{\text{sub}} H$	(288–318)	101.3	303	GS	[1980FAR/YAN]
	$\Delta_{\text{sub}} H$	(312–337)	79.5 ± 1.2			[1977STE2]
	$\Delta_{\text{sub}} H$	(369–397)	92 ± 8.2		RG	[1949SEA/HOP2, 1970COX/PIL]
	$\Delta_v H$		74.4 ± 0.7	298	GS	[2001PUR/CHI]
	$\Delta_v H$	(413–453)	76.8	298	GC	[1994SPI/LUI]
	$\Delta_v H$	(258–313)	81.3		GC	[1994LIU/DIC]
	$\Delta_v H$	(343–453)	68.6	398	GC	[1990HIN/BID2]
	$\Delta_v H$	(502–589)	68.7	517	A	[1987STE/MAL]
$\Delta_v H$	(387–582)	60.5	402		[1947STU]	
C ₆ D ₁₀ O	[51209-49-5]	cyclohexanone-d ₁₀				
	$\Delta_{\text{trs}} H$		7.1	216.8		
	$\Delta_{\text{trs}} H$		0.4	219.3		
	$\Delta_{\text{fus}} H$		1.19	241.5		[1997BUS/HAM]
C ₆ F ₅ NO ₂	[880-76-4]	pentafluoronitrobenzene				
	$\Delta_{\text{fus}} H$		11.81	250.5		[1996DOM/HEA]
C ₆ F ₆	[392-56-3]	hexafluorobenzene				
	$\Delta_{\text{fus}} H$		11.59	278.3		[1996DOM/HEA]
	$\Delta_{\text{sub}} H$	(215–278)	49.2	263	A	[1987STE/MAL, 1965DOU/OSB]
	$\Delta_{\text{sub}} H$	(238–268)	49.8	253	IP,A	[1979SCO/OSB]
	$\Delta_{\text{sub}} H$		46.0	316	B	[1965COU/GRE]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(288–333)	36.1 ± 0.1	298		[2005DIA/GON]
	$\Delta_v H$	(318–376)	34.4	333	EB	[1990AMB/EWI]
	$\Delta_v H$	(403–516)	31.8	425		[1988DAV/EWI]
	$\Delta_v H$	(278–354)	36.5	293	A	[1987STE/MAL]
	$\Delta_v H$	(348–389)	33.2	363	A	[1987STE/MAL]
	$\Delta_v H$	(384–462)	32.2	399	A	[1987STE/MAL]
	$\Delta_v H$	(458–517)	31.8	473	A	[1987STE/MAL]
	$\Delta_v H$	(290–510)	35.6	298		[1982INV, 1991BAS/SVO]
	$\Delta_v H$		35.7	298	C	[1981HOS/SCO]
	$\Delta_v H$	(293–323)	35.7	308		[1980PAT/TOM]
	$\Delta_v H$	(281–335)	36.4 ± 0.1	298		[1972KRE/PRI]
	$\Delta_v H$	(278–321)	36.2	292	MM	[1969FIN]
	$\Delta_v H$	(363–516)	32.2	378	EB	[1966EVA/TIL]
	$\Delta_v H$	(275–387)	36.5	293		[1965DOU/OSB]
	$\Delta_v H$	(293–356)	35.1	308		[1964PAT/PRO]
	$\Delta_v H$	(293–358)	35.5	308		[1964PAT/PRO, 1984BOU/FRI]
C₆F₇NOS	[20094-84-2]	N-(pentafluorophenyl)imidodisulfonyl fluoride				
	$\Delta_v H$	(309–355)	45.3	332		[1968GLE/VON]
C₆F₇OP	[59646-78-5]	pentafluorophenoxydifluorophosphine				
	$\Delta_v H$	(310–363)	42.4	325		[1976FAL/DES]
C₆F₇O₂P	[59617-42-4]	pentafluorophenoxyphosphoryl difluoride				
	$\Delta_v H$	(323–367)	46.4	338		[1976FAL/DES]
C₆F₈	[5680-05-7]	perfluoro(2-methyl-3-methylenecyclobutene)				
	$\Delta_v H$	(243–306)	31.0	258	A,I	[1987STE/MAL, 1966BAN/BAR]
C₆F₁₀	[355-75-9]	perfluorocyclohexene				
	$\Delta_v H$	(277–319)	31.0	298		[1979PRI/SAP]
C₆F₁₁NO	[52225-58-8]	2,2,3,3,3-pentafluoro-N-[2,2,2-trifluoro-1-(trifluoromethyl)ethylidene]propanamide				
	$\Delta_v H$		32.7	338		[1974PET/SHR]
C₆F₁₁NO₂S	[77589-41-4]	2,2,3,3,4,4,5,5-octafluoro-1,1,2,3,4,5-hexahydro-1-[(trifluoroacetyl)imino]thiophene-1-oxide				
	$\Delta_v H$		33.9	383		[1981ABE/SHR2]
C₆F₁₂	[1805-22-7]	perfluoromethylcyclopentane				
	$\Delta_v H$		30.68	298	EB	[1998EWI/SAN]
C₆F₁₂	[355-68-0]	perfluorocyclohexane				
	$\Delta_{\text{sub}} H$	(252–326)	36.4	267	A	[1987STE/MAL, 1967CRO/TAY]
	$\Delta_{\text{sub}} H$	(293–333)	36.2	313		[1957ROW/THA]
	$\Delta_v H$	(373–457)	28.0	388		[1988DAV/EWI]
	$\Delta_v H$	(350–451)	28.1	365	A	[1987STE/MAL]
	$\Delta_v H$	(274–322)	36.0	298		[1979PRI/SAP]
	$\Delta_v H$	(336–394)	29.6	351		[1957MCC/DOU, 1984BOU/FRI]
C₆F₁₂	[2994-71-0]	perfluoro(1,2-dimethylcyclobutane)				
	$\Delta_v H$	(242–318)	32.1	257	A	[1987STE/MAL]
C₆F₁₂N₂	[19451-96-8]	N,N,N,N-tetrakis(trifluoromethyl)-1,2-ethynylendiamine				
	$\Delta_v H$	(305–328)	32.1	316	A	[1987STE/MAL, 1968HAS/TIP]
C₆F₁₂N₂OS	[34619-84-6]	1,1,1,3,3,3-hexafluoro-2-isocyanato-N-[2,2,2-trifluoro-1-(trifluoromethyl)ethylidene]-2-propanamine				
	$\Delta_v H$		39.3	375	I	[1972SWI/BAB]
C₆F₁₂N₂O₂S	[62609-66-9]	1,1,1-trifluoro-N'-(trifluoroacetyl)-N-[2,2,2-trifluoro-1-(trifluoromethyl)ethylidene]methanesulfonimidamide				
	$\Delta_v H$		32.6	404	I	[1977KIT/SHR]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₆ F ₁₂ N ₂ S	[31340-33-7]	<i>bis</i> [2,2,2-trifluoro-1-(trifluoromethyl)ethylidene]sulfoxylic diamide				
	$\Delta_v H$		40.6	391	I	[1972SWI/BAB]
C ₆ F ₁₂ N ₂ S ₂	[38005-16-2]	<i>bis</i> (hexafluoroisopropylideneimino) disulfide				
	$\Delta_v H$		46	417	I	[1972MET/SHR]
C ₆ F ₁₂ O	[788-40-9]	perfluoro(methoxycyclopentane)				
	$\Delta_v H$	(246–330)	38.6	261	A	[1987STE/MAL, 1972DYK]
	$\Delta_v H$		30.3	330		[1957POR/CAD]
C ₆ F ₁₂ O ₂	[24165-10-4]	trifluoroacetic acid, 2,2,2-trifluoro-1,1- <i>bis</i> (trifluoromethyl) ethyl ester				
	$\Delta_v H$	(264–298)	34.3	279	A	[1987STE/MAL, 1975WAL/DES2]
	$\Delta_v H$		33.1	329	HG	[1973MAJ/SHR]
C ₆ F ₁₂ O ₄	[55100-93-1]	carbonoperoxoic acid, O-[2,2,2-trifluoro-1,1- <i>bis</i> (trifluoromethyl)ethyl-O-(trifluoromethyl) ester ester				
	$\Delta_v H$	(273–315)	33.5	288	A	[1987STE/MAL, 1975WAL/DES2]
C ₆ F ₁₃ NS	[53120-07-9]	2,2,2-trifluoro-N-[1,2,2,2-tetrafluoro-1-(trifluoromethyl)ethyl]ethanimidothioic acid, trifluoromethyl ester				
	$\Delta_v H$		35.3	360		[1975PET/SHR]
C ₆ F ₁₄	[355-42-0]	perfluorohexane				
	$\Delta_{\text{trs}}H$		0.97	103		
	$\Delta_{\text{fus}}H$		6.84	185		[1996DOM/HEA, 1986STA]
	$\Delta_v H$	(289–333)	32.5 ± 0.1	298		[2005DIA/GON]
	$\Delta_v H$	(261–334)	34.4	276	A	[1987STE/MAL]
	$\Delta_v H$	(285–340)	31.4	298		[1984BOU/FRI, 1991BAS/SVO]
	$\Delta_v H$	(433–449)	33.4	441	A	[1987STE/MAL, 1978MOU]
	$\Delta_v H$	(303–330)	31.5	316		[1958DUN/MUR, 1984BOU/FRI]
C ₆ F ₁₄	[335-04-4]	perfluoro-2-methylpentane				
	$\Delta_v H$	(280–340)	31.4	298		[1984BOU/FRI, 1991BAS/SVO]
	$\Delta_v H$	(253–329)	34.5	268	A	[1987STE/MAL, 1967CRO/TAY, 1984BOU/FRI]
	$\Delta_v H$	(277–341)	32.5	292		[1952STI/CAD, 1984BOU/FRI]
C ₆ F ₁₄	[865-71-4]	perfluoro-3-methylpentane				
	$\Delta_v H$	(282–333)	30.8	297	A	[1987STE/MAL]
C ₆ F ₁₄	[354-96-1]	perfluoro-2,3-dimethylbutane				
	$\Delta_v H$	(260–340)	31.6	298		[1984BOU/FRI, 1991BAS/SVO]
	$\Delta_v H$	(262–333)	33.0	277	A	[1987STE/MAL, 1967CRO/TAY, 1984BOU/FRI]
C ₆ F ₁₄ IP	[756-18-3]	<i>bis</i> (heptafluoropropyl) iodophosphine				
	$\Delta_v H$	(273–353)	41.6	313		[1959EME/SMI]
C ₆ F ₁₄ N ₂ S	[34451-12-2]	<i>bis</i> [1,2,2,2-tetrafluoro-1-(trifluoromethyl)ethyl] sulfur diimide				
	$\Delta_v H$	(325–378)	38.5	340	A	[1987STE/MAL, 1972SWI/SHR]
C ₆ F ₁₄ O	[356-62-7]	perfluorodipropyl ether				
	$\Delta_v H$	(306–327)	31.2 ± 0.4	298	EB	[1989VAR/PAS]
C ₆ F ₁₅ N	[359-70-6]	perfluorotriethylamine				
	$\Delta_{\text{trs}}H$		1.56	146.4		
	$\Delta_{\text{fus}}H$		5.56	156.2		[1996DOM/HEA]
	$\Delta_v H$	(297–343)	34.0 ± 0.4	298	EB	[1995VAR/DRO]
	$\Delta_v H$		34.2 ± 0.1	298	C	[1995VAR/DRO]
		(320–334)	32.8	327	A	[1987STE/MAL]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(317–349)	32.9	332	A	[1987STE/MAL]
C ₆ F ₁₅ NO	[54566-82-4]	1,1,1,2,3,3,3,-heptafluoro-N-(pentafluoroethyl)-N-(trifluoromethyl)-2-propanamine				
	$\Delta_v H$		27.1	338		[1975PET/SHR2]
C ₆ F ₁₅ O ₄ S ₂	[63441-15-6]	2,2,4,4-tetrafluoro-1,1,3,3-tetrahydro-1,1,3,3-tetrakis(trifluoromethoxy)-1,3-dithietane				
	$\Delta_v H$		37.2	404	I	[1977KIT/SHR3]
C ₆ F ₁₆ N ₂ S	[59617-31-1]	<i>bis</i> [1,2,2,2-tetrafluoro-1-(trifluoromethyl)ethyl]diimidodisulfuryl fluoride				
	$\Delta_v H$		35.8			[1976STA/MEW]
C ₆ F ₁₆ S	[1423-18-3]	difluoro <i>bis</i> [1,2,2,2-tetrafluoro-1-(trifluoromethyl)ethyl] sulfur				
	$\Delta_v H$	(273–383)	36.6	328	A	[1987STE/MAL, 1999DYK/SVO]
C ₆ F ₁₈ NP ₃	[na]	nitriolo <i>tris</i> [<i>bis</i> (trifluoromethyl)phosphine]				
	$\Delta_{\text{sub}} H$	(273–309)	68.4	291		[1965BUR/HEN]
C ₆ F ₂₀ N ₃ O ₃ P	[na]	phosphorous <i>tris</i> [<i>bis</i> (trifluoromethyl)nitroxide] difluoride				
	$\Delta_v H$		39.3	421		[1973WAN/SHR]
C ₆ N ₂	[16419-78-6]	dicyanobutadiyne				
	$\Delta_{\text{sub}} H$	(294–335)	34.4	309	A	[1987STE/MAL]
	$\Delta_{\text{sub}} H$	(295–335)	35.9	315	I	[1957SAG]
	$\Delta_v H$	(341–369)	30.2	355	A	[1987STE/MAL, 1957SAG]
C ₆ N ₄	[670-54-2]	tetracyanoethylene				
	$\Delta_{\text{fus}} H$		24.92	472.2		[1991RAD/RAD]
	$\Delta_{\text{sub}} H$	(333–371)	81.4	348	A	[1987STE/MAL]
	$\Delta_{\text{sub}} H$	(290–312)	84.3	302	TE,ME	[1983DEW/VAN]
	$\Delta_{\text{sub}} H$		81.2 ± 5.9	350		[1963BOY, 1970COX/PIL]
$\Delta_{\text{sub}} H$		78.0		GS	[1958LOO/DOW]	
C ₆ N ₆ O ₃	[na]	benzotrifurazan				
	$\Delta_{\text{sub}} H$	(303–333)	95.8 ± 3.8			[1999MAT/PEP]
C ₆ N ₆ O ₆	[na]	benzotrifuroxan				
	$\Delta_{\text{sub}} H$	(363–433)	172.0 ± 2.5			[1999MAT/PEP]
C ₆ N ₈ O ₈	[19451-95-7]	4,4''-dinitro-3,3':4',3''-ter-1,2,5-oxadiazole-2'-oxide				
	$\Delta_{\text{fus}} H$		93.83	381.9		[2004GEN/PEI]
C ₆ HBrF ₁₂ N ₂	[19451-95-7]	<i>trans</i> 1-bromo-N,N,N',N'-tetrakis(trifluoromethyl)vinylenediamine				
	$\Delta_v H$	(348–371)	32.3	359	A	[1987STE/MAL, 1968FRE/TIP]
C ₆ HBr ₅ O	[608-71-9]	pentabromophenol				
	$\Delta_{\text{fus}} H$		27.6	469.8		[2004KUR/MAE]
	$\Delta_{\text{trs}} H$		11.29	441.5		
	$\Delta_{\text{fus}} H$		19.14	502		[1995WOJ/TOU]
C ₆ HCIF ₁₁ NO	[52225-62-4]	N-[1-chloro-2,2,2-trifluoro-1-(trifluoromethyl)ethyl]-2,2,3,3,3-pentafluoropropanamide				
	$\Delta_v H$		40.8	381		[1974PET/SHR]
C ₆ HCl ₂ N ₃ O ₆	[1630-09-7]	1,3-dichloro-2,4,6-trinitrobenzene				
	$\Delta_v H$	(504–563)	46.9	519	A	[1987STE/MAL, 1972DYK]
	$\Delta_v H$	(504–533)	80.4			[1968MAK]
C ₆ HCl ₃ F ₈ O ₂	[2106-54-9]	octafluoro-3,5,6-trichlorohexanoic acid				
	$\Delta_v H$	(373–505)	64.2	388	A	[1987STE/MAL, 1972DYK, 1957BAR/SEF]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound					
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference	
C ₆ HCl ₃ O ₂	[634-85-5]	trichloro-1,4-benzoquinone					
	$\Delta_{\text{sub}}H$	(301–327)	88.7 ± 8.3		QF	[1927COO/COO, 1960JON, 1970COX/PIL]	
C ₆ HCl ₄ NO ₂	[117-18-0]	1,2,4,5-tetrachloro-3-nitrobenzene					
	$\Delta_{\text{fus}}H$		19.46	373.3	DSC	[1991ACR, 1990DON/DRE]	
			$\Delta_{\text{sub}}H$	91.3 ± 2.5	298	C	[2009RIB/FER6]
C ₆ HCl ₅	[608-93-5]	pentachlorobenzene					
	$\Delta_{\text{fus}}H$		20.6	357.7		[1991ACR]	
	$\Delta_{\text{sub}}H$		87.1 ± 0.4	298	C	[1991SAB/AN2]	
	Δ_vH		66.0	357		[1999ROH/RUZ]	
	Δ_vH	(413–453)	67.7	298	GC	[1994SPI/LUI]	
		(371–549)	62.1	386	A	[1987STE/MAL, 1947STU]	
C ₆ HCl ₅ O	[87-86-5]	pentachlorophenol					
	$\Delta_{\text{fus}}H$		17.15	462.5		[1991ACR, 1995WOJ/TOU]	
	$\Delta_{\text{sub}}H$	(348–403)	91.6 ± 0.5	298	GS	[2007VER/EME]	
	$\Delta_{\text{sub}}H$		67.4 ± 2.1			[UR/STU, 1970COX/PIL]	
		(463–507)	69.0	478	A	[1987STE/MAL, 1972DYK]	
C ₆ HF ₅	[363-72-4]	pentafluorobenzene					
	$\Delta_{\text{fus}}H$		10.88	225.7		[1991ACR]	
	Δ_vH	(358–397)	33.5	373	A	[1987STE/MAL]	
	Δ_vH	(393–479)	32.6	408	A	[1987STE/MAL]	
	Δ_vH	(473–531)	32.2	488	A	[1987STE/MAL]	
	Δ_vH	(290–510)	36.2	298		[1982INV, 1991BAS/SVO]	
	Δ_vH	(322–368)	34.8	337	A	[1987STE/MAL, 1968AMB]	
	Δ_vH	(373–530)	32.0	388	EB	[1966EVA/TIL]	
		(298–358)	35.7	313		[1964PAT/PRO, 1984BOU/FRI]	
C ₆ HF ₅ O	[771-61-9]	pentafluorophenol					
	$\Delta_{\text{trs}}H$		1.16	287			
	$\Delta_{\text{fus}}H$		16.41	310.6		[1968AND/COU]	
	$\Delta_{\text{sub}}H$	(273–299)	67.4 ± 1.7		GS	[1969COX/GUN, 1970COX/PIL]	
	Δ_vH	(323–455)	52.2 ± 0.4	298	EB	[1997STE/CHI2]	
		(378–428)	44.2	393	A	[1987STE/MAL, 1968AMB]	
C ₆ HF ₁₂ NO	[52225-64-6]	2,2,3,3,3-pentafluoro-N-[1,2,2,-tetrafluoro-1-(trifluoromethyl)ethyl]propanamide					
Δ_vH			41.3	368		[1974PET/SHR]	
C ₆ HF ₁₂ NOS	[62067-08-7]	2,2,2-trifluoro-N-[(trifluoromethyl)thio]ethanimidic acid, 2,2,2-trifluoro-1-(trifluoromethyl)ethyl ether					
Δ_vH			33.6	364	I	[1977BUR/SHR2]	
C ₆ H ₂ BrCl ₃ O	[na]	3-bromo-2,4,6-trichlorophenol					
	Δ_vH	(385–579)	67.1	400	A	[1987STE/MAL, 1947STU]	
C ₆ H ₂ Br ₄	[636-28-2]	1,2,4,5-tetrabromobenzene					
	$\Delta_{\text{fus}}H$		24.4	454.5		[2004KUR/MAE2]	
	$\Delta_{\text{trs}}H$		0.34	306.8			
	$\Delta_{\text{fus}}H$		27.88	453.1		[1996DOM/HEA]	
C ₆ H ₂ ClN ₃ O ₆	[88-88-0]	1-chloro-2,4,6-trinitrobenzene					
	$\Delta_{\text{sub}}H$		103.8		DSC	[1990HWA/YOS]	
			103.0			[1950NIT/SEK]	

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$		85.4		DSC	[1990HWA/YOS]
	$\Delta_v H$	(473–543)	63.1	488	A	[1987STE/MAL, 1968MAK]
C ₆ H ₂ Cl ₂ O ₂	[697-91-6]	2,6-dichloro-1,4-benzoquinone				
	$\Delta_{\text{sub}} H$	(274–315)	69.9 ± 8.3		QF	[1927COO/COO, 1960JON, 1970COX/PIL]
C ₆ H ₂ Cl ₃ F	[36556-33-9]	1-fluoro-2,4,6-trichlorobenzene				
	$\Delta_v H$	(344–489)	41.1	359	A	[1987STE/MAL]
C ₆ H ₂ Cl ₃ NO ₂	[89-69-0]	2,4,5-trichloro-1-nitrobenzene				
	$\Delta_v H$	(427–560)	56.7	442	A	[1987STE/MAL]
C ₆ H ₂ Cl ₃ NO ₂	[18708-70-8]	2,4,6-trichloro-1-nitrobenzene				
	$\Delta_{\text{sub}} H$		84.3 ± 1.9	298	C	[2009RIB/FER6]
	$\Delta_{\text{sub}} H$	(287–303)	86.9 ± 1.1	295	ME	[2009RIB/FER6]
	$\Delta_{\text{sub}} H$	(287–303)	86.7 ± 1.1	298	ME	[2009RIB/FER6]
C ₆ H ₂ Cl ₄	[634-66-2]	1,2,3,4-tetrachlorobenzene				
	$\Delta_{\text{fus}} H$		17.0	319.7	DTA	[1991SAB/AN2]
	$\Delta_{\text{fus}} H$		17.0	320		[1991ACR]
	$\Delta_{\text{sub}} H$		78.8 ± 0.2	298	C	[1991SAB/AN2]
	$\Delta_v H$	(413–453)	60.1	298	GC	[1994SPI/LUI]
	$\Delta_v H$	(341–527)	56.7	356	A	[1987STE/MAL, 1947STU]
C ₆ H ₂ Cl ₄	[634-90-2]	1,2,3,5-tetrachlorobenzene				
	$\Delta_{\text{fus}} H$		19.0	323.9		[1991ACR]
	$\Delta_{\text{sub}} H$		79.6 ± 0.3	298	C	[1991SAB/AN2]
	$\Delta_v H$	(413–453)	60.7	298	GC	[1994SPI/LUI]
C ₆ H ₂ Cl ₄	[95-94-3]	1,2,4,5-tetrachlorobenzene				
	$\Delta_{\text{fus}} H$		24.4	410.1	DSC	[2002RAI/PAN]
	$\Delta_{\text{fus}} H$		24.9	412.6	DTA	[1991SAB/AN2]
	$\Delta_{\text{fus}} H$		24.1	412.2		[1991ACR]
C ₆ H ₂ Cl ₄	[95-94-3]	1,2,4,5-tetrachlorobenzene				
	$\Delta_{\text{sub}} H$		83.2 ± 0.3	298	C	[1991SAB/AN2]
	$\Delta_v H$	(413–453)	60.7	298	GC	[1994SPI/LUI]
	$\Delta_v H$	(419–518)	52.0	434	A	[1987STE/MAL, 1947STU]
C ₆ H ₂ Cl ₄ O	[58-90-2]	2,3,4,6-tetrachlorophenol				
	$\Delta_v H$	(373–548)	64.8	388	A	[1987STE/MAL, 1947STU]
C ₆ H ₂ Cl ₄ O ₂	[1198-55-6]	3,4,5,6-tetrachloro-1,2-benzenediol				
	$\Delta_v H$	(293–323)	77.9	308	CGC	[1999LEI/WAN2]
C ₆ H ₂ Cl ₄ O ₂	[87-87-6]	tetrachlorohydroquinone				
	$\Delta_{\text{sub}} H$	(298–359)	89.0	313		[1987STE/MAL]
	$\Delta_{\text{sub}} H$	(333–356)	88.7		QF	[1927COO/COO, 1960JON]
C ₆ H ₂ Cl ₅ N	[527-20-8]	pentachloroaniline				
	$\Delta_{\text{fus}} H$		18.7	505.8		[1991ACR]
C ₆ H ₂ F ₄	[551-62-2]	1,2,3,4-tetrafluorobenzene				
	$\Delta_{\text{fus}} H$		10.93	233.3		[1973AND/MAR]
	$\Delta_v H$	(300–390)	37.5	298		[1984BOU/FRI, 1991BAS/SVO]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₆ H ₂ F ₄	$\Delta_v H$	(300–392)	36.8	315	A	[1987STE/MAL, 1975AMB/ELL2, 1984BOU/FRI]
	$\Delta_v H$	(279–323)	37.0	294	MM	[1987STE/MAL, 1969FIN]
	[2367-82-0]	1,2,3,5-tetrafluorobenzene				
	$\Delta_{\text{fus}} H$		10.67	226.9		[1973AND/MAR]
	$\Delta_v H$	(385–416)	32.4	400	A	[1987STE/MAL]
	$\Delta_v H$	(290–380)	36.0	298		[1984BOU/FRI, 1991BAS/SVO]
C ₆ H ₂ F ₄	$\Delta_v H$	(287–382)	36.0	302	A	[1987STE/MAL, 1975AMB/ELL2, 1984BOU/FRI]
	$\Delta_v H$	(279–323)	36.0	294	MM	[1987STE/MAL, 1969FIN]
	[327-54-8]	1,2,4,5-tetrafluorobenzene				
	$\Delta_{\text{fus}} H$		15.05	277		[1973AND/MAR]
	$\Delta_v H$	(290–390)	37.2	298		[1984BOU/FRI, 1991BAS/SVO]
C ₆ H ₂ F ₄	$\Delta_v H$	(390–488)	33.1	405	A	[1987STE/MAL]
	$\Delta_v H$	(488–543)	32.6	503	A	[1987STE/MAL]
	$\Delta_v H$	(293–390)	36.8	308	A	[1987STE/MAL, 1975AMB/ELL2, 1984BOU/FRI]
	[771-60-8]	pentafluoroaniline				
	$\Delta_{\text{fus}} H$		14.27	306.8		[1996DOM/HEA]
C ₆ H ₂ F ₁₂ O	[176310-30-8]	1,1,1,2,2,3,3,3-heptafluoro-3-(2,2,3,3,3-pentafluoropropoxy)propane				
	$\Delta_v H$	(288–344)	34.8	303	I	[2002MUR/YAM]
C ₆ H ₂ F ₁₂ O ₃	[205367-61-9]	1,1'-oxybis[2-(difluoromethoxy)-1,1,2,2-tetrafluoroethane]				
	$\Delta_v H$	(268–283)	38.5 ± 0.8			[1999MAR/BAS]
C ₆ H ₂ F ₁₂ O ₃ S	[53517-89-9]	<i>bis</i> (1,1,1,3,3,3-hexafluoro-2-propanol) sulfite				
	$\Delta_v H$		42.4			[1975DEM/KOV]
C ₆ H ₂ F ₁₂ O ₄	[249932-26-1]	1,1,3,3,5,5,7,7,8,8,10,10-dodecafluoro-12,4,6,9-tetraoxadecane				
	$\Delta_v H$	(263–381)	42.3 ± 0.4			[1999MAR/BAS]
C ₆ H ₂ F ₁₄ NP	[na]	amino <i>bis</i> (heptafluoropropyl)phosphine				
	$\Delta_v H$	(293–393)	38.7	343		[1959EME/SMI]
C ₆ H ₂ N ₄	[13481-25-9]	2,3-dicyanopyrazine				
	$\Delta_{\text{fus}} H$		19.8	405.1	DSC	[2006MIR/MOR]
	$\Delta_{\text{sub}} H$		89.1 ± 2.7	298	C	[2006MIR/MOR]
C ₆ H ₂ N ₄ O ₆	[5128-28-9]	4,6-dinitrobenzofurazan 1-oxide				
	$\Delta_{\text{fus}} H$		20.73	452.7		[1983RED/MUR]
C ₆ H ₃ BrCl ₂ O	[45524-77-0]	2-bromo-4,6-dichlorophenol				
	$\Delta_v H$	(357–541)	58.6	372	A	[1987STE/MAL, 1947STU]
C ₆ H ₃ BrCl ₂ O	[1940-42-7]	4-bromo-2,5-dichlorophenol				
	$\Delta_{\text{fus}} H$		22.11	343.4	DSC	[1990DON/DRE]
C ₆ H ₃ Br ₃	[615-54-3]	1,2,4-tribromobenzene				
	$\Delta_{\text{fus}} H$		17.9	317		[2004KUR/MAE2]
C ₆ H ₃ Br ₃	[626-39-1]	1,3,5-tribromobenzene				
	$\Delta_{\text{fus}} H$		21.72	395		[2005VAN/VAN]
C ₆ H ₃ Br ₃ NO ₂	[3460-18-2]	2,5-dibromonitrobenzene				

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{sub}}H$	(302–322)	96.8 ± 0.4	312	ME	[2009RIB/FER8]
	$\Delta_{\text{sub}}H$	(302–322)	97.0 ± 0.4	298	ME	[2009RIB/FER8]
C₆H₃Br₃O	[118-79-6]	2,4,6-tribromophenol				
	$\Delta_{\text{fus}}H$		20.9	367.5		[2004KUR/MAE]
	$\Delta_{\text{fus}}H$		18.52	366.2		[1991ACR]
	$\Delta_{\text{sub}}H$		97.6 ± 1.1			[1987ALL/FIN]
C₆H₃ClN₂O₂	[17348-69-5]	5-chlorobenzofurazan-1-oxide				
	$\Delta_{\text{sub}}H$		81.2 ± 1.8	298	C	[1996ACR/BOT]
C₆H₃ClN₂O₄	[97-00-7]	1-chloro-2,4-dinitrobenzene				
	$\Delta_{\text{fus}}H$		20.17	325.2		[1932KUB]
	Δ_vH	(430–590)	80.5	445	A	[1987STE/MAL]
C₆H₃ClN₂O₄	[606-21-3]	2,6-dinitrochlorobenzene				
	$\Delta_{\text{fus}}H$		18.95	361.2		[1932KUB]
C₆H₃ClO₂	[695-99-8]	chlorobenzoquinone				
	$\Delta_{\text{sub}}H$	(264–289)	69.0 ± 8.3	276	QF	[1927COO/COO, 1960JON, 1970COX/PIL]
C₆H₃Cl₂NO₂	[611-06-3]	2,4-dichloro-1-nitrobenzene				
	$\Delta_{\text{sub}}H$		87.8 ± 1.7	298	C	[2009RIB/FER9]
C₆H₃Cl₂NO₂	[89-61-2]	2,5-dichloro-1-nitrobenzene				
	$\Delta_{\text{sub}}H$		87.4 ± 2.4	298	C	[2009RIB/FER9]
C₆H₃Cl₂NO₂	[99-54-7]	3,4-dichloro-1-nitrobenzene				
	$\Delta_{\text{fus}}H$		17.95	314.1	DSC	[2003VER/SCH]
	$\Delta_{\text{fus}}H$		17.6	316		[1981MAS/OLE]
	$\Delta_{\text{sub}}H$		85.8 ± 2.5	298	C	[2009RIB/FER9]
	$\Delta_{\text{sub}}H$	(283–311)	83.1 ± 0.6	298	GS	[2003VER/SCH]
	Δ_vH	(316–346)	65.2 ± 0.2	298	GS	[2003VER/SCH]
	Δ_vH	(417–515)	55.5	432	A	[1987STE/MAL]
C₆H₃Cl₂NO₂	[618-62-2]	3,5-dichloro-1-nitrobenzene				
	$\Delta_{\text{sub}}H$		83.2 ± 1.5	298	C	[2009RIB/FER9]
C₆H₃Cl₃	[120-82-1]	1,2,4-trichlorobenzene				
	$\Delta_{\text{sub}}H$	(279–298)	62.3	289	RG	[1949SEA/HOP, 1960JON]
	Δ_vH		55.8	290		[1999ROH/RUZ]
	Δ_vH	(391–490)	49.5	406	EB	[1998ROH/RUZ]
	Δ_vH	(413–453)	57.6	298	GC	[1994SPI/LUI]
	Δ_vH		55.5 ± 0.1	298	C	[1987YAN/GU]
	Δ_vH	(279–298)	47.0	288	RG	[1949SEA/HOP]
	Δ_vH	(311–486)	49.3	326		[1947STU]
C₆H₃Cl₃	[87-61-6]	1,2,3-trichlorobenzene				
	$\Delta_{\text{fus}}H$		20.5	326.9		[1991ACR]
	$\Delta_{\text{fus}}H$		17.25	322.9	DSC	[1990DON/DRE]
	$\Delta_{\text{sub}}H$	(258–313)	72.7			[1994LIU/DIC]
	$\Delta_{\text{sub}}H$		75.1 ± 0.75	298		[1985YAN/GU, 1987YAN/GU]
	$\Delta_{\text{sub}}H$	(289–303)	65.7	296	RG	[1949SEA/HOP, 1960JON]
	Δ_vH		54.5	325		[1999ROH/RUZ]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound					
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference	
		$\Delta_v H$	(413–453)	57.2	298	GC	[1994SPI/LUI]
		$\Delta_v H$	(258–313)	54.3		GC	[1994LIU/DIC]
		$\Delta_v H$	(293–383)	53.5	308	A	[1987STE/MAL]
		$\Delta_v H$	(313–492)	47.4	328	A	[1987STE/MAL, 1947STU]
C₆H₃Cl₃	[108-70-3]	1,3,5-trichlorobenzene					
		$\Delta_{\text{fus}} H$		17.56	335.9		[2005VAN/VAN]
		$\Delta_{\text{fus}} H$		18.2	336.7		[1991ACR]
		$\Delta_{\text{sub}} H$		72.7 ± 0.5	298		[1985YAN/GU, 1987YAN/GU]
		$\Delta_{\text{sub}} H$	(282–301)	56.5	291	RG	[1949SEA/HOP, 1960JON]
		$\Delta_v H$	(338–415)	50.3 ± 0.1	375	DM	[2001BLO/VAN]
		$\Delta_v H$		51.7	337		[1999ROH/RUZ]
		$\Delta_v H$	(413–453)	59	298	GC	[1994SPI/LUI]
		$\Delta_v H$	(336–482)	48.8	351	A	[1987STE/MAL, 1947STU]
C₆H₃Cl₃O	[95-95-4]	2,4,5-trichlorophenol					
		$\Delta_{\text{fus}} H$		21.59	340.3	DSC	[1990DON/DRE]
		$\Delta_v H$	(345–525)	54.5	360	A	[1987STE/MAL, 1947STU]
C₆H₃Cl₃O	[88-06-2]	2,4,6-trichlorophenol					
		$\Delta_{\text{sub}} H$	(299–340)	82.3 ± 0.3	298	GS	[2007FRE/OLI]
		$\Delta_v H$	(343–375)	62.5	359	GS	[2007VER/EME]
		$\Delta_v H$	(343–375)	67.2 ± 0.3	298	GS	[2007VER/EME]
		$\Delta_v H$	(344–463)	58.2	404		[1995MOK/PAU, 2007VER/EME]
		$\Delta_v H$	(344–463)	66.1 ± 0.4	298		[1995MOK/PAU, 2007VER/EME]
		$\Delta_v H$	(349–519)	58.8	364	A	[1987STE/MAL, 1947STU]
C₆H₃Cl₃O₂	[56961-20-7]	3,4,5-trichloro-1,2-benzenediol					
		$\Delta_v H$	(293–323)	79.3	308	CGC	[1999LEI/WAN2]
C₆H₃Cl₃O₂	[608-94-6]	trichlorohydroquinone					
		$\Delta_{\text{sub}} H$	(298–336)	101.5	313	A	[1987STE/MAL]
		$\Delta_{\text{sub}} H$	(314–335)	101.3	324	QF	[1927COO/COO, 1960JON]
C₆H₃Cl₄N	[3481-20-7]	2,3,5,6-tetrachloroaniline					
		$\Delta_{\text{sub}} H$		86.0 ± 2.0	298	C	[2007RIB/AMA3]
C₆H₃Cl₄N	[69045-78-9]	2-chloro-5-(trichloromethyl)pyridine					
		$\Delta_{\text{fus}} H$	(80–345)	14.5	324.7	AC	[2004KON/TAN]
C₆H₃Cl₄N	[1929-82-4]	2-chloro-6-(trichloromethyl)pyridine					
		$\Delta_{\text{fus}} H$	(13–316)	20.3	337.8	AC	[1996DOM/HEA]
C₆H₃F₃	[372-38-3]	1,3,5-trifluorobenzene					
		$\Delta_v H$	(280–320)	33.9	298		[1984BOU/FRI, 1991BAS/SVO]
		$\Delta_v H$	(279–350)	34.5	294	A,MM	[1987STE/MAL, 1969FIN, 1972DYK]
C₆H₃F₄N	[363-73-5]	2,3,4,6-tetrafluoroaniline					
		$\Delta_v H$		50.4 ± 0.6	298	C	[2007RIB/FER]
C₆H₃F₉O₂	[42031-16-3]	trifluoroacetic acid, 2,2,2-trifluoro-1-methyl-1-(trifluoromethyl)ethyl ester					
		$\Delta_v H$		33.5	338	HG	[1973MAJ/SHR]
C₆H₃F₉O₂	[24165-09-1]	acetic acid, 2,2,2-trifluoro-1,1-bis(trifluoromethyl)ethyl ester					
		$\Delta_v H$	(273–328)	40.1	288	A	[1987STE/MAL, 1975WAL/DES2]
C₆H₃F₁₀NS	[54120-08-0]	2,2,2-trifluoro-N-[1,2,2,2-tetrafluoro-1-(trifluoromethyl)ethyl]ethanimidothioic acid, methyl ester					

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
		$\Delta_v H$	31.6	383		[1975PET/SHR]
C ₆ H ₃ F ₁₁ O	[176310-29-5]	1,1,1,2,2,3,3-heptafluoro-3-(2,2,3,3-tetrafluoropropoxy)propane				
	$\Delta_v H$	(288–357)	37.2	303	I	[2002MUR/YAM]
C ₆ H ₃ F ₁₁ O	[181214-74-4]	1,1,1,2,2,3,3,4,4,5,5-undecafluoro-5-methoxypentane				
	$\Delta_v H$	(288–358)	36.6	303	I	[2002MUR/YAM]
C ₆ H ₃ F ₁₁ O	[203783-56-6]	1,1,1,2,3,3,4,4-octafluoro-4-methoxy-2-(trifluoromethyl)butane				
	$\Delta_v H$	(288–357)	36		I	[2002MUR/YAM]
C ₆ H ₃ F ₁₁ O	[290-28-8]	1,1,1,2,3,3-hexafluoro-3-(2,2,3,3,3-pentafluoropropoxy)propane				
	$\Delta_v H$	(293–360)	38.4	308	I	[2002MUR/YAM]
C ₆ H ₃ N ₃ O ₄	[18771-85-2]	4-nitrobenzofurazan-1-oxide				
	$\Delta_{\text{sub}} H$		97.3 ± 1.6	298	C	[1996ACR/BOT]
C ₆ H ₃ N ₃ O ₆	[603-13-4]	1,2,3-trinitrobenzene				
	$\Delta_v H$	(523–573)	60.3	538	A	[1987STE/MAL, 1968MAK, 1972DYK]
C ₆ H ₃ N ₃ O ₆	[610-31-1]	1,2,4-trinitrobenzene				
	$\Delta_v H$	(523–573)	82.6	538	A	[1987STE/MAL, 1968MAK, 1972DYK]
C ₆ H ₃ N ₃ O ₆	[99-35-4]	1,3,5-trinitrobenzene				
	$\Delta_{\text{fus}} H$		1.9	370		
	$\Delta_{\text{fus}} H$		14.81	380.3		[1996DOM/HEA]
	$\Delta_{\text{sub}} H$	(313–395)	107.3 ± 0.6	298	ME	[1978CUN/PAL]
	$\Delta_{\text{sub}} H$		99.6 ± 2.1			[1950NIT/SEK, 1970COX/PIL]
C ₆ H ₃ N ₃ O ₇	[88-89-1]	2,4,6-trinitrophenol (picric acid)				
	$\Delta_{\text{fus}} H$		17.1	394.1		[1996DOM/HEA]
	$\Delta_{\text{sub}} H$		106.3		DSC	[1990HWA/YOS]
	$\Delta_{\text{sub}} H$	(314–406)	105.1 ± 1.6	298	ME	[1978CUN/PAL]
	$\Delta_v H$		87.9		DSC	[1990HWA/YOS]
		(468–598)	106.4	483	A	[1987STE/MAL]
		Note: The value of 106.4 kJ/mole from [1987STE/MAL] is likely an enthalpy of sublimation				
C ₆ H ₃ N ₃ O ₈	[82-71-3]	2,4,6-trinitroresorcinol (styphnic acid)				
	$\Delta_{\text{fus}} H$		33.5	454.8		[1996DOM/HEA]
	$\Delta_{\text{sub}} H$		120.1		DSC	[1990HWA/YOS]
	$\Delta_{\text{sub}} H$	(325–436)	120.8 ± 1.1	298	ME	[1978CUN/PAL]
			92.9		DSC	[1990HWA/YOS]
C ₆ H ₄ BrCl	[694-80-4]	1-bromo-2-chlorobenzene				
	$\Delta_{\text{fus}} H$		12.37	260.6		[1996DOM/HEA]
C ₆ H ₄ BrCl	[108-37-2]	1-bromo-3-chlorobenzene				
	$\Delta_{\text{fus}} H$		12.29	252		[1996DOM/HEA]
	$\Delta_v H$	(252–469)	52.2	267	A	[1987STE/MAL, 1972DYK]
C ₆ H ₄ BrCl	[106-39-8]	1-bromo-4-chlorobenzene				
	$\Delta_{\text{fus}} H$	(6–350)	18.4	338	AC	[2000TOZ/AKU]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
		$\Delta_{\text{fus}}H$	18.76	337.8		[1996DOM/HEA]
		$\Delta_{\text{sub}}H$	69.34 ± 0.11	298	DM	[2000OON/VAN]
		$\Delta_{\text{sub}}H$	(250–335) 69.3 ± 0.4	298	TE,ME,DM	[1998OON/VAN]
		$\Delta_{\text{sub}}H$	69.1 ± 0.2	298		[1998OON/VAN]
		$\Delta_{\text{sub}}H$	(294–337) 67.9 ± 0.8	316		[1961WAL/SMI]
		Δ_vH	(333–470) 49.1	348	A	[1987STE/MAL]
		Δ_vH	(305–470) 49.7	320		[1947STU]
C₆H₄BrI	[583-55-1]	1-bromo-2-iodobenzene				
		$\Delta_{\text{fus}}H$	14.42	294.2		[1991ACR]
C₆H₄BrI	[591-18-4]	1-bromo-3-iodobenzene				
		$\Delta_{\text{fus}}H$	12.16	282.5		[1991ACR]
C₆H₄BrI	[589-87-7]	1-bromo-4-iodobenzene				
		$\Delta_{\text{fus}}H$	19.38	363.5		[2001VAN/OON]
		$\Delta_{\text{fus}}H$	19.13	363.3		[1991ACR]
		$\Delta_{\text{sub}}H$	78.53 ± 0.16	298	DM	[2000OON/VAN]
		$\Delta_{\text{sub}}H$	(279–355) 78.5 ± 0.4	298	ME,TE,DM	[1998OON/VAN]
C₆H₄BrNO₂	[577-19-5]	2-bromo-1-nitrobenzene				
		$\Delta_{\text{sub}}H$	(275–295) 85.5 ± 0.3	285	ME	[2010RIB/FER4]
		$\Delta_{\text{sub}}H$	(275–295) 85.2 ± 0.3	298	ME	[2010RIB/FER4]
C₆H₄BrNO₂	[585-79-5]	3-bromo-1-nitrobenzene				
		$\Delta_{\text{sub}}H$	(280–295) 87.0 ± 0.5	287	ME	[2010RIB/FER4]
		$\Delta_{\text{sub}}H$	(280–295) 86.8 ± 0.5	298	ME	[2010RIB/FER4]
C₆H₄BrNO₂	[586-78-7]	4-bromo-1-nitrobenzene				
		$\Delta_{\text{sub}}H$	(289–309) 86.6 ± 0.6	299	ME	[2010RIB/FER4]
		$\Delta_{\text{sub}}H$	(289–309) 86.6 ± 0.6	298	ME	[2010RIB/FER4]
		$\Delta_{\text{sub}}H$	(293–303) 88.3	303	ME	[1987STE/MAL, 1925SWA/MAC]
C₆H₄Br₂	[583-53-9]	1,2-dibromobenzene				
		$\Delta_{\text{fus}}H$	12.61	275		[1991ACR]
		Δ_vH	(388–568) 50.1	403	A	[1987STE/MAL, 1972DYK]
C₆H₄Br₂	[108-36-1]	1,3-dibromobenzene				
		$\Delta_{\text{fus}}H$	13.21	266.3		[1991ACR]
		Δ_vH	(417–500) 48.3	432	A	[1987STE/MAL]
C₆H₄Br₂	[106-37-6]	1,4-dibromobenzene				
		$\Delta_{\text{fus}}H$	18.6	357.7		
		$\Delta_{\text{fus}}H$	20.39	360.5		[2004KUR/MAE2, 2005VAN/VAN]
		$\Delta_{\text{fus}}H$	20.04	360.1		[1991ACR]
		$\Delta_{\text{sub}}H$	74.23 ± 0.11	298	ME	[2000OON/VAN]
		$\Delta_{\text{sub}}H$	(298–354) 73.2	313		[1987STE/MAL]
		$\Delta_{\text{sub}}H$	(278–353) 73.3 ± 0.4	326		[1961WAL/SMI]
		$\Delta_{\text{sub}}H$	(228–347) 73.8	288		[1959STE/GRE]
		$\Delta_{\text{sub}}H$	(248–303) 59.8	298	ME,GS	[1940ZIB, 1960JON]
		Δ_vH	(373–493) 49.9	388	A	[1987STE/MAL, 1972DYK]
C₆H₄Br₂O	[615-58-7]	2,4-dibromophenol				
		$\Delta_{\text{fus}}H$	14.64	313		[1991ACR]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₆ H ₄ Br ₃ N	[147-82-0]	2,4,6-tribromoaniline				
	$\Delta_{\text{fus}}H$		25.75	393	DSC	[2006RIB/FER]
	$\Delta_{\text{sub}}H$		96.7 ± 1.7	298	C	[2006RIB/FER]
			101.1 ± 1.1			[1987ALL/FIN]
C ₆ H ₄ ClF	[625-98-9]	1-chloro-3-fluorobenzene				
	Δ_vH	(273–403)	37.4	288	A	[1987STE/MAL]
C ₆ H ₄ ClF	[352-33-0]	1-chloro-4-fluorobenzene				
	$\Delta_{\text{fus}}H$		13.9	245		[2004CER/PER]
C ₆ H ₄ ClF	[352-33-0]	1-chloro-4-iodobenzene				
	$\Delta_{\text{fus}}H$		16.1	326.7		[2000VAN/OON]
	$\Delta_{\text{sub}}H$		71.86 ± 0.21	298	DM	[2000OON/VAN]
	$\Delta_{\text{sub}}H$	(259–320)	71.9 ± 0.4	298	ME,TE,DM	[1998OON/VAN]
	$\Delta_{\text{sub}}H$	(303–323)	61.1 ± 0.6			[1953EWA, 1960JON]
		(333–500)	56.5	348	A	[1987STE/MAL]
C ₆ H ₄ ClNO ₂	[88-73-3]	1-chloro-2-nitrobenzene				
	$\Delta_{\text{fus}}H$		18.21	305.8		[2003VER/SCH]
	$\Delta_{\text{fus}}H$		18.11	305.8		[2007STR/RUZ]
	$\Delta_{\text{fus}}H$		19.08	308.2		[1981MAS/OLE]
	$\Delta_{\text{sub}}H$		80.9 ± 1.5	298	C	[2009RIB/FER7]
	$\Delta_{\text{sub}}H$	(278–305)	80.8 ± 0.3	298	GS	[2003VER/SCH]
	Δ_vH	(307–334)	60.4 ± 0.3	298	GS	[2003VER/SCH]
		(420–516)	52.1	435	EB	[1984PUT/IVA]
C ₆ H ₄ ClNO ₂	[121-73-3]	1-chloro-3-nitrobenzene				
	$\Delta_{\text{fus}}H$		18.65	318		[2003VER/SCH]
	$\Delta_{\text{fus}}H$		19.52	316.9		[2007STR/RUZ]
	$\Delta_{\text{fus}}H$		19.37	317.6		[1991ACR]
	$\Delta_{\text{sub}}H$		82.5 ± 1.5	298	C	[2009RIB/FER7]
	$\Delta_{\text{sub}}H$	(281–314)	81.3 ± 0.3	298	GS	[2003VER/SCH]
	$\Delta_{\text{sub}}H$	(275–286)	74.7 ± 1.7			[1935TRI, 1938WOL/WEG]
	Δ_vH	(319–364)	60.2 ± 0.2	298	GS	[2003VER/SCH]
		(414–506)	51.5	429	EB	[1984PUT/IVA]
C ₆ H ₄ ClNO ₂	[100-00-5]	1-chloro-4-nitrobenzene				
	$\Delta_{\text{fus}}H$		15.03	355.1		[2007STR/RUZ]
	$\Delta_{\text{fus}}H$		16.17	356.1	DSC	[2003VER/SCH]
	$\Delta_{\text{fus}}H$		11.85	354.6		[1996DOM/HEA]
	$\Delta_{\text{fus}}H$		14.1	357		[1981MAS/OLE]
	$\Delta_{\text{sub}}H$	(303–339)	74.7 ± 0.1	298	GS	[2003VER/SCH]
	$\Delta_{\text{sub}}H$	(283–303)	83.2	293	ME	[1987STE/MAL, 1925SWA/MAC]
		(385–515)	51.3	400	A	[1987STE/MAL]
C ₆ H ₄ ClNO ₃	[619-08-9]	2-chloro-4-nitrophenol				
	$\Delta_{\text{fus}}H$		20.88	380.7		[2007MOR/MIR]
			99.0 ± 2.1	298	C	[2007MOR/MIR]
C ₆ H ₄ ClNO ₃	[89-64-5]	4-chloro-2-nitrophenol				
	$\Delta_{\text{fus}}H$		22.69	360.3		[2007MOR/MIR]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		T _m (K)	Method	Reference
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)			
C ₆ H ₄ ClNO ₃	$\Delta_{\text{sub}}H$		87.6 ± 0.9	298	C	[2007MOR/MIR]
	[610-78-6]	4-chloro-3-nitrophenol				
	$\Delta_{\text{fus}}H$		25.97	399.4		[2007MOR/MIR]
C ₆ H ₄ ClNO ₃	$\Delta_{\text{sub}}H$		111.0 ± 3.3	298	C	[2007MOR/MIR]
	[54127-63-8]	5-chloro-6-hydroxynicotinic acid				
	$\Delta_{\text{sub}}H$	(457–487)	149.1 ± 2.6	472	DSC	[2009SAN/FIG]
C ₆ H ₄ ClN ₃ O ₄	$\Delta_{\text{sub}}H$	(457–487)	151.3 ± 2.8	298	DSC	[2009SAN/FIG]
	[3531-19-9]	2-chloro-4,6-dinitroaniline				
	$\Delta_{\text{sub}}H$	(358–380)	114.2 ± 0.5	369	ME	[2010RIB/RIB]
C ₆ H ₄ ClN ₃ O ₄	$\Delta_{\text{sub}}H$	(358–380)	115.0 ± 0.9	298	ME	[2010RIB/RIB]
	[5388-62-5]	4-chloro-2,6-dinitroaniline				
	$\Delta_{\text{sub}}H$	(335–359)	104.7 ± 0.4	347	ME	[2010RIB/RIB]
C ₆ H ₄ Cl ₂	$\Delta_{\text{sub}}H$	(335–359)	105.2 ± 0.7	298	ME	[2010RIB/RIB]
	[95-50-1]	1,2-dichlorobenzene				
	$\Delta_{\text{fus}}H$		12.4	255.9	DSC	[2009WEI/JIN, 2008WEI]
	$\Delta_{\text{fus}}H$		12.93	256.5		[1991ACR]
	Δ_vH		51.2	256		[1999ROH/RUZ]
	Δ_vH	(363–454)	44.5	376	EB	[1998ROH/RUZ]
	Δ_vH	(256–287)	50.8	271		[1996POL/GUE]
	Δ_vH	(413–453)	50.9	298	GC	[1994SPI/LUI]
	Δ_vH	(258–313)	51.2		GC	[1994LIU/DIC]
	Δ_vH		48.5 ± 0.1	298	C	[1989ZN/ZHE]
	Δ_vH	(373–453)	44.0	388	A	[1987STE/MAL]
	Δ_vH	(360–450)	49.9	298		[1984BOU/FRI, 1991BAS/SVO]
	Δ_vH	(301–343)	50.0	322	GS	[1982GRA/FOS]
C ₆ H ₄ Cl ₂	[541-73-1]	1,3-dichlorobenzene				
	$\Delta_{\text{fus}}H$		12.6	248.3	DSC	[2009WEI/JIN, 2008WEI]
	$\Delta_{\text{fus}}H$		12.64	248.4		[1991ACR]
	Δ_vH		50.4	248		[1999ROH/RUZ]
	Δ_vH	(357–448)	44.1	372	EB	[1998ROH/RUZ]
	Δ_vH	(250–274)	50.0	262		[1996POL/GUE]
	Δ_vH	(413–453)	53.9	298	GC	[1994SPI/LUI]
	Δ_vH	(360–450)	47.0	298		[1984BOU/FRI, 1991BAS/SVO]
	Δ_vH	(348–513)	44.7	363	A	[1987STE/MAL, 1972DYK]
C ₆ H ₄ Cl ₂	[106-46-7]	1,4-dichlorobenzene				
	$\Delta_{\text{fus}}H$		18.19	326.2	DSC	[2009WEI/JIN, 2008WEI]
	$\Delta_{\text{trs}}H$		1.24	275		
	$\Delta_{\text{trs}}H$		0.18	306		
	$\Delta_{\text{fus}}H$	(5–380)	17.91	326.2	AC	[2005VAN/VAN]
	$\Delta_{\text{fus}}H$		18.16	326		[1991ACR]
	$\Delta_{\text{sub}}H$		64.75 ± 0.15	298	DM	[2000OON/VAN]
	$\Delta_{\text{sub}}H$	(258–313)	53.1			[1994LIU/DIC]
	$\Delta_{\text{sub}}H$		65.2 ± 2.0	298	C	[1989AN/ZHE]
	$\Delta_{\text{sub}}H$	(303–423)	65.4	313	GS	[1985ROR]
	$\Delta_{\text{sub}}H$		65.7			[1981DEK/VAN]
	$\Delta_{\text{sub}}H$	(293–311)	64.8 ± 0.8	303		[1961WAL/SMI, 1970COX/PIL]
	$\Delta_{\text{sub}}H$	(311–325)	63 ± 0.4	318		[1961WAL/SMI]
$\Delta_{\text{sub}}H$	(248–303)	56.9	275	ME	[1940DAR/VER, 1960JON]	

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		T _m (K)	Method	Reference	
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)				
		$\Delta_v H$	46.4	326		[1999ROH/RUZ]	
		$\Delta_v H$	(358–448)	44.2	EB	[1998ROH/RUZ]	
		$\Delta_v H$	(413–453)	54.8	GC	[1994SPI/LUI]	
		$\Delta_v H$	(258–313)	U 35.0	GC	[1994LIU/DIC]	
		$\Delta_v H$	(341–448)	45.0	A	[1987STE/MAL]	
		$\Delta_v H$	(370–450)	47.8		[1984BOU/FRI, 1991BAS/SVO]	
C₆H₄Cl₂N₂O₂	[6627-34-5]	2,5-dichloro-4-nitrobenzenamine					
		$\Delta_{\text{sub}} H$	(352–374)	113.2 ± 0.6	363	ME	[2009RIB/RIB]
		$\Delta_{\text{sub}} H$	(352–374)	114.3 ± 0.9	298	ME	[2009RIB/RIB]
C₆H₄Cl₂N₂O₂	[99-30-9]	2,6-dichloro-4-nitroaniline					
		$\Delta_{\text{fus}} H$		29.48	467.2	DSC	[1991ACR, 1990DON/DRE]
		$\Delta_{\text{sub}} H$	(344–366)	108.2 ± 0.6	355	ME	[2009RIB/RIB]
		$\Delta_{\text{sub}} H$	(344–366)	109.2 ± 0.9	298	ME	[2009RIB/RIB]
C₆H₄Cl₂N₂O₂	[6641-64-1]	4,5-dichloro-2-nitroaniline					
		$\Delta_{\text{sub}} H$	(351–367)	108.4 ± 0.7	359	ME	[2009RIB/RIB2]
		$\Delta_{\text{sub}} H$	(351–367)	109.4 ± 0.9	298	ME	[2009RIB/RIB2]
C₆H₄Cl₂O	[576-24-9]	2,3-dichlorophenol					
		$\Delta_{\text{fus}} H$		21.36	330		[1991ACR]
		$\Delta_{\text{sub}} H$	(294–327)	76.9 ± 0.4	298	GS	[2007VER/EME]
		$\Delta_{\text{sub}} H$		71.7 ± 2.2	298	C	[1994RIB/FER2]
		$\Delta_v H$	(331–358)	57.3 ± 0.2	298	GS	[2007VER/EME]
C₆H₄Cl₂O	[128-83-2]	2,4-dichlorophenol					
		$\Delta_{\text{fus}} H$		20.09	318		[1996DOM/HEA]
		$\Delta_{\text{sub}} H$	(278–315)	78.0 ± 0.3	298	GS	[2007VER/EME]
		$\Delta_{\text{sub}} H$		70.1 ± 1.1	298	C	[1994RIB/FER2]
		$\Delta_v H$	(317–344)	56.6	331	GS	[2007VER/EME]
		$\Delta_v H$	(317–344)	59.0 ± 0.4	298	GS	[2007VER/EME]
		$\Delta_v H$	(323–443)	52.3	383		[1995MOK/PAU, 2007VER/EME]
		$\Delta_v H$	(323–443)	58.1 ± 0.3	298		[1995MOK/PAU, 2007VER/EME]
		$\Delta_v H$	(326–483)	60.8	341	A	[1987STE/MAL, 1947STU, 1975ARR/MEL]
C₆H₄Cl₂O	[583-78-8]	2,5-dichlorophenol					
		$\Delta_{\text{fus}} H$		22.43	331		[1996DOM/HEA]
		$\Delta_{\text{sub}} H$	(294–327)	77.3 ± 0.1	298	GS	[2007VER/EME]
		$\Delta_{\text{sub}} H$		73.6 ± 2.1	298	C	[1994RIB/FER2]
		$\Delta_v H$	(333–361)	53.1	347	GS	[2007VER/EME]
		$\Delta_v H$	(333–361)	56.7 ± 0.1	298	GS	[2007VER/EME]
C₆H₄Cl₂O	[87-65-0]	2,6-dichlorophenol					
		$\Delta_{\text{fus}} H$		22.14	340		[1996DOM/HEA]
		$\Delta_{\text{sub}} H$	(299–331)	79.3 ± 0.2	298	GS	[2007VER/EME]
		$\Delta_{\text{sub}} H$		75.8 ± 1.1	298	C	[1994RIB/FER2]
		$\Delta_v H$	(341–371)	55.4	356	GS	[2007VER/EME]
		$\Delta_v H$	(341–371)	59.6 ± 0.3	298	GS	[2007VER/EME]
		$\Delta_v H$	(343–457)	51.6	400		[1995MOK/PAU, 2007VER/EME]
		$\Delta_v H$	(343–457)	58.5 ± 0.5	298		[1995MOK/PAU, 2007VER/EME]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₆ H ₄ Cl ₂ O	$\Delta_v H$	(333–493)	57.9	348	A	[1987STE/MAL, 1947STU]
	[95-77-2]	3,4-dichlorophenol				
	$\Delta_{\text{fus}} H$		20.93	341		[1996DOM/HEA]
	$\Delta_{\text{sub}} H$	(291–337)	89.8 ± 0.4	298	GS	[2007VER/EME]
	$\Delta_{\text{sub}} H$		81.3 ± 2.3	298	C	[1994RIB/FER2]
C ₆ H ₄ Cl ₂ O	$\Delta_v H$	(341–368)	66.7	355	GS	[2007VER/EME]
	$\Delta_v H$	(341–368)	70.8 ± 0.2	298	GS	[2007VER/EME]
	[591-35-5]	3,5-dichlorophenol				
	$\Delta_{\text{fus}} H$		20.51	341		[1996DOM/HEA]
	$\Delta_{\text{sub}} H$		82.8 ± 1.1	298	C	[1994RIB/FER2]
C ₆ H ₄ Cl ₂ O ₂	[3428-24-8]	4,5-dichloro-1,2-benzenediol				
	$\Delta_v H$	(293–323)	70.5	308	CGC	[1999LEI/WAN2]
C ₆ H ₄ Cl ₂ O ₂	[20103-10-0]	2,6-dichlorohydroquinone				
	$\Delta_{\text{sub}} H$	(324–345)	92.0 ± 8.3		QF	[1927COO/COO, 1960JON, 1970COX/PIL]
C ₆ H ₄ Cl ₂ O ₃	[na]	vinyl mucochlorate				
	$\Delta_v H$	(273–333)	63.9	288	A	[1987STE/MAL]
C ₆ H ₄ Cl ₃ N	[634-67-3]	2,3,4-trichloroaniline				
	$\Delta_{\text{sub}} H$		92.4 ± 1.7	298	C	[2002RIB/AMA]
C ₆ H ₄ Cl ₃ N	[636-30-6]	2,4,5-trichloroaniline				
	$\Delta_{\text{sub}} H$		86.3 ± 2.5	298	C	[2002RIB/AMA]
C ₆ H ₄ Cl ₃ N	[634-93-5]	2,4,6-trichloroaniline				
	$\Delta_{\text{sub}} H$		85.3 ± 1.9	298	C	[2002RIB/AMA]
	$\Delta_v H$	(407–535)	92.9	422	A	[1987STE/MAL, 1947STU]
		Note: Enthalpy of vaporization is likely in error				
C ₆ H ₄ Cl ₃ N	[634-91-3]	3,4,5-trichloroaniline				
	$\Delta_{\text{sub}} H$		92.9 ± 3.3	298	C	[2002RIB/AMA]
C ₆ H ₄ F ₂	[367-11-3]	1,2-difluorobenzene				
	$\Delta_{\text{fus}} H$		11.05	226		[1996DOM/HEA]
	$\Delta_v H$	(300–400)	36.2	298		[1984BOU/FRI, 1991BAS/SVO]
	$\Delta_v H$	(304–403)	35.5	319	EB	[1963SCO/MES, 1984BOU/FRI]
	$\Delta_v H$		34.6 ± 0.1	327	C	[1963SCO/MES]
	$\Delta_v H$		33.5 ± 0.1	345	C	[1963SCO/MES]
C ₆ H ₄ F ₂	[372-18-9]	1,2-difluorobenzene				
	$\Delta_{\text{trs}} H$		0.83	186.8		
	$\Delta_{\text{fus}} H$		8.58	204		[1996DOM/HEA]
	$\Delta_v H$	(310–400)	34.6	298		[1980OSB/SCO, 1991BAS/SVO]
C ₆ H ₄ F ₂	[540-36-3]	1,4-difluorobenzene				
	$\Delta_v H$	(300–400)	35.8	298		[1980OSB/SCO, 1991BAS/SVO]
C ₆ H ₄ F ₂ O	[6418-38-8]	2,3-difluorophenol				
	$\Delta_{\text{sub}} H$		68.2 ± 1.5	298	C	[2010RIB/FER2]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₆ H ₄ F ₂ O	[367-27-1] $\Delta_v H$	2,4-difluorophenol	58.8 ± 0.9	298	C	[2010RIB/FER2]
C ₆ H ₄ F ₂ O	[2713-31-7] $\Delta_{\text{sub}} H$	2,5-difluorophenol	68.0 ± 1.4	298	C	[2010RIB/FER2]
C ₆ H ₄ F ₂ O	[28177-48-2] $\Delta_{\text{sub}} H$	2,6-difluorophenol	77.8 ± 2.0	298	C	[2010RIB/FER2]
C ₆ H ₄ F ₂ O	[2713-33-9] $\Delta_{\text{sub}} H$	3,4-difluorophenol	72.9 ± 1.5	298	C	[2010RIB/FER2]
C ₆ H ₄ F ₂ O	[2713-34-0] $\Delta_{\text{sub}} H$	3,5-difluorophenol	72.8 ± 1.5	298	C	[2010RIB/FER2]
C ₆ H ₄ F ₃ N	[3862-73-5] $\Delta_v H$	2,3,4-trifluoroaniline	53.7 ± 0.5	298	C	[2007RIB/FER]
C ₆ H ₄ F ₃ N	[67815-56-9] $\Delta_v H$	2,3,6-trifluoroaniline	50.1 ± 0.5	298	C	[2007RIB/FER]
C ₆ H ₄ F ₁₀ O	[65064-78-0] $\Delta_v H$	1,1,1,2,3,3-hexafluoro-3-(2,2,3,3-tetrafluoropropoxy)propane (293–379)	42.3	308	I	[2002MUR/YAM]
C ₆ H ₄ INO ₂	[609-73-4] $\Delta_v H$	2-iodo-1-nitrobenzene (433–563)	59.9	448	A	[1987STE/MAL]
C ₆ H ₄ INO ₂	[645-00-1] $\Delta_{\text{sub}} H$	3-iodo-1-nitrobenzene (295–306)	83.2 ± 1.2	300		[1935TRI, 1938WOL/WEG, 1960JON]
C ₆ H ₄ I ₂	[615-42-9] $\Delta_{\text{fus}} H$	1,2-diiodobenzene	14.01	296.6		[1991ACR]
C ₆ H ₄ I ₂	[626-00-6] $\Delta_{\text{fus}} H$	1,3-diiodobenzene	15.93	307.4		[1991ACR]
C ₆ H ₄ I ₂	[624-38-4] $\Delta_{\text{trs}} H$ $\Delta_{\text{fus}} H$ $\Delta_{\text{fus}} H$	1,4-diiodobenzene	0.22 22.3 22.37	320 402.4 402		[2001VAN/OON] [1991ACR]
	$\Delta_{\text{sub}} H$	(372–401)	63.4	386.5	A	[1987STE/MAL]
	$\Delta_v H$	(402–560)	52.6	417	A	[1987STE/MAL]
C ₆ H ₄ N ₂	[100-54-9] $\Delta_v H$	nicotinic acid nitrile (453–479)	45	466	A	[1987STE/MAL, 1972DYK]
C ₆ H ₄ N ₂	[100-70-9] $\Delta_{\text{sub}} H$	2-cyanopyridine	70.7 ± 1.2	298	C	[1984BIC/PIL]
C ₆ H ₄ N ₂	[100-54-9] $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$	3-cyanopyridine	72.1 ± 1.8 79	298	C DSC	[1984BIC/PIL] [1989SHI/SHI]
C ₆ H ₄ N ₂	[100-48-1] $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$	4-cyanopyridine	73.2 ± 0.6 75.6	298	C DSC	[1984BIC/PIL] [1989SHI/SHI]
C ₆ H ₄ N ₂ O	[14906-64-0] $\Delta_{\text{sub}} H$	3-cyanopyridine N-oxide (345–392)	101.9 ± 2.0	298	ME	[1998RIB/MAT]
C ₆ H ₄ N ₂ O	[14906-59-3]	4-cyanopyridine N-oxide				

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound					
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference	
		$\Delta_{\text{sub}}H$	(345–392)	104.4 ± 4.3	298	ME	[1998RIB/MAT]
C ₆ H ₄ N ₂ O	[273-09-6]		benzofurazan				
		$\Delta_{\text{sub}}H$		64.4 ± 1.6	298	C	[1990LEI/PIL]
		$\Delta_{\text{sub}}H$		64.9	298		[1980ARS]
C ₆ H ₄ N ₂ O ₂	[480-96-6]		benzofurazan N-oxide				
		$\Delta_{\text{sub}}H$		79.6 ± 1.7	298	C	[1990LEI/PIL]
C ₆ H ₄ N ₂ O ₃ -C ₆ H ₄ N ₂ O ₃	[56079-22-2]		1-nitro-2-nitrosobenzene (dimer)				
		$\Delta_{\text{sub}}H$	(323–343)	95.5	333	A	[1987STE/MAL, 1974PEP/LEB]
C ₆ H ₄ N ₂ O ₄	[528-29-0]		1,2-dinitrobenzene				
		$\Delta_{\text{fus}}H$		22.84	396.1		[1996DOM/HEA]
		$\Delta_{\text{sub}}H$	(323–353)	95.5 ± 0.9	298	GS	[1997VER3]
		$\Delta_{\text{sub}}H$	(323–353)	93.1 ± 0.9	338	GS	[1997VER3]
		$\Delta_{\text{sub}}H$	(343–387)	82.9	358		[1987STE/MAL]
		$\Delta_{\text{sub}}H$	(343–397)	87.9 ± 2.1	298	TE	[1976FER/PIA]
		$\Delta_{\text{sub}}H$	(328–338)	86.6 ± 1.2	309		[1935TRI, 1938WOL/WEG, 1960JON]
		Δ_vH	(454–593)	60.0	469	A	[1987STE/MAL, 1972DYK]
C ₆ H ₄ N ₂ O ₄	[99-65-0]		1,3-dinitrobenzene				
		$\Delta_{\text{fus}}H$		19.68	360.4		[2002MUS/RAZ]
		$\Delta_{\text{fus}}H$		17.36	363.2		[1991ACR]
		$\Delta_{\text{sub}}H$	(335–356)	76.1	345.5		[1987STE/MAL]
		$\Delta_{\text{sub}}H$	(332–383)	87.0 ± 0.8	298	TE	[1976FER/PIA]
		$\Delta_{\text{sub}}H$	(315–329)	81.1 ± 1.7	323		[1935TRI, 1938WOL/WEG, 1960JON]
		$\Delta_{\text{sub}}H$		81.2 ± 1.7			[1950NIT/SEK3, 1970COX/PIL]
	Δ_vH	(336–379)	96.7	351	A	[1987STE/MAL]	
C ₆ H ₄ N ₂ O ₄	[100-25-4]		1,4-dinitrobenzene				
		$\Delta_{\text{fus}}H$		17.58	446		[2002MUS/RAZ]
		$\Delta_{\text{fus}}H$		28.12	446.7		[1991ACR]
		$\Delta_{\text{sub}}H$		94.3 ± 0.7	298		[1997VER3]
		$\Delta_{\text{sub}}H$	(339–398)	96.2 ± 2.5	298	TE	[1976FER/PIA]
		$\Delta_{\text{sub}}H$	(345–368)	89.1 ± 1.7	343		[1935TRI, 1938WOL/WEG, 1960JON]
	Δ_vH	(445–572)	60.3	460	A	[1987STE/MAL, 1972DYK]	
C ₆ H ₄ N ₂ O ₅	[66-56-8]		2,3-dinitrophenol				
		$\Delta_{\text{fus}}H$		22.67	419		[2002MUS/RAZ]
		$\Delta_{\text{fus}}H$		26.24	417		[1996DOM/HEA]
		$\Delta_{\text{sub}}H$	(303–343)	96.6	323		[1958HOY/PEP]
C ₆ H ₄ N ₂ O ₅	[51-28-5]		2,4-dinitrophenol				
		$\Delta_{\text{fus}}H$		26.19	383.2		[2002MUS/RAZ]
		$\Delta_{\text{fus}}H$		24.17	388		[1996DOM/HEA]
		$\Delta_{\text{sub}}H$	(293–333)	104.6 ± 4.2	313		[1958HOY/PEP, 1970COX/PIL]
C ₆ H ₄ N ₂ O ₅	[329-71-5]		2,5-dinitrophenol				
		$\Delta_{\text{fus}}H$		23.73	381		[1996DOM/HEA]
	$\Delta_{\text{sub}}H$	(278–333)	93.4	306		[1958HOY/PEP]	
C ₆ H ₄ N ₂ O ₅	[573-56-8]		2,6-dinitrophenol				

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
			22.91	329		[2002MUS/RAZ]
			19.58	336		[1996DOM/HEA]
		(293–333)	112.1 ± 4.2	313		[1958HOY/PEP, 1970COX/PIL]
C₆H₄N₂O₅	[577-71-9]	3,4-dinitrophenol				
			25.37	407		[1996DOM/HEA]
		(328–383)	123.5	383		[1958HOY/PEP]
C₆H₄N₄O₂	[1516-60-5]	4-nitrophenyl azide				
			17.1	345.4	DSC	[1997FIN/GAR]
C₆H₄N₄O₆	[489-98-5]	2,4,6-trinitroaniline				
			124.7		DSC	[1990HWA/YOS]
		(328–371)	115.9	343	LE	[1987STE/MAL, 1969ROS/DIC]
		(326–449)	125.3 ± 0.8	298	ME	[1978CUN/PAL]
			95.8		DSC	[1990HWA/YOS]
C₆H₄N₂S	[273-13-2]	2,1,3-benzothiadiazole				
			70.73 ± 0.2	298	C	[1998SAB/KUA]
C₆H₄O₂	[106-51-4]	1,4-benzoquinone				
			18.40 ± 0.1	385.1	DSC	[2004ROJ/FOR]
			18.35 ± 0.3	385.7	HFC	[2004ROJ/FOR]
			18.45	388		[1991ACR]
			66.7 ± 1.6	298	DSC	[2004ROJ/FOR]
			68.0 ± 0.5	262	ME,TE	[1981DEK/SMI]
			62.8 ± 3.3			[1956MAG, 1977PED/RYL]
			68.5 ± 0.6			[1953SEK/SUZ]
		(260–278)	62.8	269	QF	[1927COO/COO]
		(388–402)	47.8	395	A	[1987STE/MAL]
(C₆H₄O₂)–(C₆H₆O₂)	[106-34-3]	quinhydrone (quinone-hydroquinone)				
			88.6 ± 1	313	ME,TE	[1981DEK/SMI]
			U 181.2			[1953SEK/SUZ, 1960JON]
			NA			[1951NIT/SEK]
C₆H₄O₅	[3238-40-2]	furan-2,5-dicarboxylic acid				
		(378–402)	121.3	391	TE,ME	[1983SPE/CLI]
C₆H₄S₄	[31366-25-3]	tetrathiafulvene				
			61.0		TGA	[1995YAS/TAK]
			95.3 ± 1	345	TE,ME	[1980DEK/GOV]
		(341–361)	92 ± 6.3	351	HSA	[1979SAN/EPS]
		(331–355)	95.3	343		[1999DYK/SVO]
(C₆H₄S₄)–(C₁₂H₄N₄)	[40210-84-2]	(tetrathiofulvalene)-(7,7,8,8-tetracyanoquinodimethane) (TTF-TCNQ)				
			130 ± 2	410	TE,ME	[1980DEK/GOV]
C₆H₅Br	[108-86-1]	bromobenzene				
			10.7	242.4		[1996DOM/HEA]
		(330–430)	44.8	298		[1984BOU/FRI, 1991BAS/SVO]
			44.0	293	C	[1975MAS/SCO]
		(333–463)	42.3	348	A	[1987STE/MAL, 1972DYK]
			44.5 ± 0.1	298	C	[1968WAD]
		(329–427)	42.4	344		[1955DRE/MAR]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₆ H ₅ BrO	[95-56-7]	2-bromophenol				
	$\Delta_v H$		55.5 ± 1.3	298	C	[2009RIB/FER5]
	$\Delta_v H$		50.2			[1986BAL/GNA]
C ₆ H ₅ BrO	[591-20-8]	3-bromophenol				
	$\Delta_v H$	(410–510)	73.5	425	A	[1987STE/MAL]
	$\Delta_v H$		55.2			[1986BAL/GNA]
C ₆ H ₅ BrO	[106-41-2]	4-bromophenol				
	$\Delta_{\text{fus}} H$		17.6	338.2		[2004KUR/MAE]
	$\Delta_{\text{fus}} H$		16.57	336		[1996DOM/HEA]
	$\Delta_{\text{sub}} H$		83.1 ± 1.6	298	C	[2009RIB/FER5]
	$\Delta_{\text{sub}} H$	(260–302)	87.3 ± 0.4	298	ME	[1971PAR/ROC]
	$\Delta_v H$	(390–511)	58.8	405	A	[1987STE/MAL]
	$\Delta_v H$		58.6			[1986BAL/GNA]
C ₆ H ₅ BrS	[6320-02-1]	2-bromobenzenethiol				
	$\Delta_v H$		50.6			[1986BAL/GNA]
C ₆ H ₅ BrS	[6320-01-0]	3-bromobenzenethiol				
	$\Delta_v H$		51.1			[1986BAL/GNA]
C ₆ H ₅ BrS	[106-53-6]	4-bromobenzenethiol				
	$\Delta_v H$		52.3			[1986BAL/GNA]
C ₆ H ₅ Br ₂ N	[615-57-6]	2,4-dibromoaniline				
	$\Delta_{\text{fus}} H$		21.37	351.4	DSC	[2006RIB/FER]
	$\Delta_{\text{sub}} H$		88.0 ± 1.5	298	C	[2006RIB/FER]
C ₆ H ₅ Br ₂ N	[3638-73-1]	2,5-dibromoaniline				
	$\Delta_{\text{fus}} H$		20.47	328.1	DSC	[2006RIB/FER]
	$\Delta_{\text{sub}} H$		85.7 ± 1.9	298	C	[2006RIB/FER]
C ₆ H ₅ Br ₂ N	[608-30-0]	2,6-dibromoaniline				
	$\Delta_{\text{fus}} H$		21.79	355.5	DSC	[2006RIB/FER]
	$\Delta_{\text{sub}} H$		80.7 ± 1.4	298	C	[2006RIB/FER]
C ₆ H ₅ Cl	[108-90-7]	chlorobenzene				
	$\Delta_{\text{fus}} H$		9.55	227.9		[1996DOM/HEA]
	$\Delta_v H$	(313–353)	40.3	298	CGC	[1995CHI/HOS]
	$\Delta_v H$	(413–453)	43.9	298	GC	[1994SPI/LUI]
	$\Delta_v H$	(258–313)	48.1		GC	[1994LIU/DIC]
	$\Delta_v H$		40.6 ± 0.3		GC	[1989AZA]
	$\Delta_v H$	(405–597)	35.4	420	A	[1987STE/MAL]
	$\Delta_v H$	(335–405)	41.0	298		[1984BOU/FRI, 1991BAS/SVO]
	$\Delta_v H$		41.0 ± 0.1	298	C	[1968WAD]
	$\Delta_v H$	(333–405)	38.8	348	A	[1987STE/MAL, 1952BRO, 1984BOU/FRI]
$\Delta_v H$	(253–303)	37.3	278	ME	[1940ZIB]	
C ₆ H ₅ ClN ₂ O ₂	[121-87-9]	2-chloro-4-nitroaniline				
	$\Delta_{\text{sub}} H$		101.8 ± 1.8	298	C	[2003RIB/LIM]
	$\Delta_{\text{sub}} H$	(335–351)	100.3 ± 1.5	343	ME	[2003RIB/LIM]
	$\Delta_{\text{sub}} H$	(335–351)	102.6 ± 1.5	298	ME	[2003RIB/LIM]
C ₆ H ₅ ClN ₂ O ₂	[6283-25-6]	2-chloro-5-nitroaniline				

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₆ H ₅ ClO	$\Delta_{\text{sub}}H$		100.3 ± 2.2	298	C	[2003RIB/LIM]
	$\Delta_{\text{sub}}H$		99.3 ± 1.6	333	ME	[2003RIB/LIM]
	$\Delta_{\text{sub}}H$		101.0 ± 1.6	298	ME	[2003RIB/LIM]
	[95-57-8]	2-chlorophenol				
	$\Delta_{\text{trs}}H$		0.09	276		
	$\Delta_{\text{fus}}H$		12.52	283		[1991ACR]
	Δ_vH	(288–321)	51.9	305	GS	[2007VER/EME]
	Δ_vH	(288–321)	52.3 ± 0.2	298	GS	[2007VER/EME]
	Δ_vH	(337–447)	47.0	352		[1995GAB/MAR]
	Δ_vH		45.2			[1966GOO/DEP]
Δ_vH	(354–448)	47.2	369	A	[1987STE/MAL]	
Δ_vH	(333–449)	50.1	348	A	[1987STE/MAL, 1974KIV/NAD]	
Δ_vH	(285–447)	45.2	300		[1947STU]	
C ₆ H ₅ ClO	[108-43-0]	3-chlorophenol				
	$\Delta_{\text{fus}}H$		14.91	305.8		[1991ACR]
	$\Delta_{\text{sub}}H$	(275–306)	76.9 ± 0.3	298	GS	[2007VER/EME]
	$\Delta_{\text{sub}}H$		53.1			[1938WOL/WEG, 1960JON, 1970COX/PIL]
	Δ_vH	(308–335)	61.9	322	GS	[2007VER/EME]
	Δ_vH	(308–335)	63.5 ± 0.3	298	GS	[2007VER/EME]
	Δ_vH		52.3			[1986BAL/GNA]
	Δ_vH	(317–487)	53.1	332	A	[1987STE/MAL, 1947STU]
C ₆ H ₅ ClO	[106-48-9]	4-chlorophenol				
	$\Delta_{\text{fus}}H$		14.07	315.9		[1991ACR]
	$\Delta_{\text{sub}}H$	(283–313)	77.1 ± 0.2	298	GS	[2007VER/EME]
	$\Delta_{\text{sub}}H$	(252–293)	60.8	278	A	[1987STE/MAL]
	$\Delta_{\text{sub}}H$		51.9			[1938WOL/WEG, 1960JON, 1970COX/PIL]
	Δ_vH	(318–351)	61.9	335	GS	[2007VER/EME]
	Δ_vH	(318–351)	64.4 ± 0.3	298	GS	[2007VER/EME]
	Δ_vH	(373–493)	60.6	388	A	[1987STE/MAL]
	Δ_vH		54.0			[1986BAL/GNA]
	Δ_vH	(323–493)	52.8	338		[1947STU]
C ₆ H ₅ ClO ₂	[2138-22-9]	4-chloro-1,2-benzenediol				
	Δ_vH	(293–323)	70.2	308	CGC	[1999LEI/WAN2]
C ₆ H ₅ ClO ₂	[615-67-8]	chlorohydroquinone				
	$\Delta_{\text{sub}}H$	(306–334)	102.9 ± 8.3	320	QF	[1927COO/COO, 1960JON, 1970COX/PIL]
C ₆ H ₅ ClO ₂ S	[98-09-9]	benzenesulfonyl chloride				
	Δ_vH	(339–524)	54.4	354		[1999DYK/SVO]
	Δ_vH	(338–525)	57.2	353	A	[1987STE/MAL, 1947STU]
C ₆ H ₅ ClO ₃ S	[99-66-8]	4-chlorobenzene sulfonic acid				
	$\Delta_{\text{fus}}H$		10.6	333.2	DSC	[1995MAC/JOY]
C ₆ H ₅ ClS	[6320-03-2]	2-chlorobenzenethiol				
	Δ_vH		47.7			[1986BAL/GNA]
C ₆ H ₅ ClS	[2037-31-2]	3-chlorobenzenethiol				
	Δ_vH		48.5			[1986BAL/GNA]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₆ H ₅ ClS	[106-54-7] $\Delta_v H$	4-chlorobenzenethiol	48.5			[1986BAL/GNA]
C ₆ H ₅ Cl ₂ N	[608-27-5] $\Delta_{\text{sub}} H$	2,3-dichloroaniline	82.2 ± 1.0	298	C	[2006RIB/AMA]
C ₆ H ₅ Cl ₂ N	[554-00-7] $\Delta_{\text{sub}} H$	2,4-dichloroaniline	84.7 ± 1.3	298	C	[2006RIB/AMA]
C ₆ H ₅ Cl ₂ N	[95-82-9] $\Delta_{\text{sub}} H$	2,5-dichloroaniline	83.4 ± 1.3	298	C	[2006RIB/AMA]
C ₆ H ₅ Cl ₂ N	[608-31-1] $\Delta_{\text{sub}} H$	2,6-dichloroaniline	74.2 ± 0.9	298	C	[2006RIB/AMA]
C ₆ H ₅ Cl ₂ N	[95-76-1] $\Delta_{\text{fus}} H$	3,4-dichloroaniline	21.69	344.5		[2003VER/SCH]
	$\Delta_v H$	(420–545)	58.6	435	A	[1987STE/MAL]
C ₆ H ₅ Cl ₂ OP	[3426-89-9] $\Delta_v H$	phenyl dichlorophosphite (363–480)	52.7	378		[2008SHA/WU]
C ₆ H ₅ Cl ₂ O ₂ P	[770-12-7] $\Delta_v H$	phenyl dichlorophosphate (339–513)	63.6	354	A	[1987STE/MAL, 1947STU]
C ₆ H ₅ F	[462-06-6] $\Delta_{\text{fus}} H$	fluorobenzene	11.31	230.9		[1996DOM/HEA]
	$\Delta_v H$	(358–530)	31.9	373	A	[1987STE/MAL]
	$\Delta_v H$	(373–419)	31.8	388	A	[1987STE/MAL]
	$\Delta_v H$	(414–501)	31.0	429	A	[1987STE/MAL]
	$\Delta_v H$	(497–561)	30.9	512	A	[1987STE/MAL]
	$\Delta_v H$	(255–360)	34.5	298		[1984BOU/FRI, 1991BAS/SVO]
	$\Delta_v H$	(312–394)	33.6	327	EB	[1987STE/MAL, 1956SCO/MCC2]
	$\Delta_v H$		33.5 ± 0.1	318	C	[1956SCO/MCC2]
	$\Delta_v H$		32.4 ± 0.1	337	C	[1956SCO/MCC2]
	$\Delta_v H$		31.2 ± 0.1	358	C	[1956SCO/MCC2]
$\Delta_v H$		29.7 ± 0.1	382	C	[1956SCO/MCC2]	
C ₆ H ₅ FO	[367-12-4] $\Delta_v H$	2-fluorophenol	52.3 ± 0.8	298	C	[2009RIB/FER]
C ₆ H ₅ FO	[372-20-3] $\Delta_v H$	3-fluorophenol	60.1 ± 0.9	298	C	[2009RIB/FER]
	$\Delta_v H$	(373–451)	50.3	388	A	[1987STE/MAL]
C ₆ H ₅ FO	[371-41-5] $\Delta_{\text{sub}} H$	4-fluorophenol	73.9 ± 1.4	298	C	[2009RIB/FER]
	$\Delta_v H$	(360–460)	48.8	375	A	[1987STE/MAL]
C ₆ H ₅ F ₂ N	[4519-40-8] $\Delta_v H$	2,3-difluoroaniline	49.3 ± 0.5	298	C	[2007RIB/FER]
C ₆ H ₅ F ₂ N	[367-25-9] $\Delta_v H$	2,4-difluoroaniline	52.1 ± 0.5	298	C	[2007RIB/FER]
C ₆ H ₅ F ₂ N	[367-30-6] $\Delta_v H$	2,5-difluoroaniline	52.5 ± 0.5	298	C	[2007RIB/FER]
C ₆ H ₅ F ₂ N	[5509-65-9]	2,6-difluoroaniline				

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$		47.5 ± 0.5	298	C	[2007RIB/FER]
C ₆ H ₅ F ₂ N	[3863-11-4]	3,4-difluoroaniline				
	$\Delta_v H$		53.3 ± 0.5	298	C	[2007RIB/FER]
C ₆ H ₅ F ₈ NOS	[77984-30-6]	1-(ethylimino)-2,2,3,3,4,4,5,5-octafluoro-1,1,2,3,4,5-hexahydrothiophene-1-oxide				
	$\Delta_v H$		31.4	333		[1981ABE/SHR]
C ₆ H ₅ F ₉ O	[163702-05-4]	1-ethoxy-1,1,2,2,3,3,4,4,4-nonafluorobutane				
	$\Delta_v H$	(293–350)	34.2	308	I	[2002MUR/YAM]
C ₆ H ₅ I	[591-50-4]	iodobenzene				
	$\Delta_{\text{fus}} H$		9.75	241.8		[1996DOM/HEA]
	$\Delta_v H$	(243–255)	43.1			[1960JON]
	$\Delta_v H$	(248–303)	40.0	275	ME	[1940ZIB]
	$\Delta_v H$	(313–353)	47.4	298	CGC	[1995CHI/HOS]
	$\Delta_v H$	(462–679)	41.1	477	A	[1987STE/MAL]
	$\Delta_v H$	(320–460)	48.9	298		[1984BOU/FRI, 1991BAS/SVO]
	$\Delta_v H$	(273–358)	51.4	288	A	[1987STE/MAL, 1972DYK]
C ₆ H ₅ NO	[586-96-9]	nitrosobenzene (dimer)				
	$\Delta_{\text{sub}} H$	(297–339)	85.1	312	A	[1987STE/MAL, 1974PEP/LEB]
	$\Delta_{\text{sub}} H$		80.8			[1930DRU/FLA]
C ₆ H ₅ NO ₂	[98-95-3]	nitrobenzene				
	$\Delta_{\text{fus}} H$		12.12	278.8		[1996DOM/HEA]
	$\Delta_v H$	(313–353)	54.5	298	CGC	[1995CHI/HOS]
	$\Delta_v H$	(288–318)	54.3	303		[1985ZAR]
	$\Delta_v H$	(291–305)	56.1 ± 1.7	298	ME	[1971LEB/KAT]
	$\Delta_v H$		55.0	298		[1971KUS/WAD2]
	$\Delta_v H$	(279–296)	54.7	287	A	[1987STE/MAL, 1972DYK, 1960LYN/WIL]
	$\Delta_v H$	(283–303)	52.5	293	ME	[1958SKL/MAR]
C ₆ H ₅ NO ₂	[98-98-6]	2-pyridinecarboxylic acid (picolinic acid)				
	$\Delta_{\text{fus}} H$		30.0	411		[1996DOM/HEA]
	$\Delta_{\text{sub}} H$		91.0 ± 0.5	329	C	[1999SAB/IDE]
	$\Delta_{\text{sub}} H$		92.7 ± 0.5	298		[1999SAB/IDE]
	$\Delta_{\text{sub}} H$	(345–392)	98.0 ± 2.3	298	ME	[1998RIB/MAT]
C ₆ H ₅ NO ₂	[59-67-6]	3-pyridinecarboxylic acid (nicotinic acid)				
	$\Delta_{\text{fus}} H$		97.1	509.3		[2004WAN/WAN]
		Note: This value is considerably larger than values below—likely in error.				
	$\Delta_{\text{trs}} H$		0.81	451.4		
	$\Delta_{\text{fus}} H$		27.57	509.1		[2004WAN/TAN3]
	$\Delta_{\text{trs}} H$		0.78	452		
	$\Delta_{\text{fus}} H$		26.7	510		[1993ELM/CHA]
	$\Delta_{\text{sub}} H$	(473–498)	89.3		TG, DTA	[2002MEN/DOL]
	$\Delta_{\text{sub}} H$	(352–360)	123.9 ± 3.7	298	ME	[2000RIB/GON]
	$\Delta_{\text{sub}} H$		101.1 ± 0.6	362	C	[1999SAB/IDE]
$\Delta_{\text{sub}} H$		105.2 ± 0.6	298		[1999SAB/IDE]	
$\Delta_{\text{sub}} H$		123.4 ± 1.2	298	C	[1984BIC/PIL]	

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₆ H ₅ NO ₂	[55-22-1]	4-pyridinecarboxylic acid (isonicotinic acid)				
	$\Delta_{\text{fus}}H$		135	593		[1996DOM/HEA]
		Note: Enthalpy of fusion is much too large, compound likely decomposed				
	$\Delta_{\text{sub}}H$		107.7 ± 0.7	362	C	[1999SAB/IDE]
	$\Delta_{\text{sub}}H$		111.3 ± 0.6	298		[1999SAB/IDE]
C ₆ H ₅ NO ₃	[88-75-5]	2-nitrophenol				
	$\Delta_{\text{fus}}H$		17.05	316.3		[2002MUS/RAZ]
	$\Delta_{\text{fus}}H$		17.45	318.2		[1991ACR, 1994SAB/GOU]
	$\Delta_{\text{sub}}H$		73.3	298	C	[1994SAB/GOU]
	$\Delta_{\text{sub}}H$	(273–292)	54.8	282.5		[1987STE/MAL]
	$\Delta_{\text{sub}}H$	(298–310)	73.2 ± 1.3			[1935TRI, 1938WOL/WEG, 1960JON]
	Δ_vH	(319–346)	58.4 ± 0.5	298	GS	[2007HEI/KAP]
	Δ_vH	(366–490)	55.9	381	A	[1987STE/MAL]
	Δ_vH	(324–347)	U 43.3	298	ME	[1958SKL/MAR, 2007HEI/KAP]
	Δ_vH	(322–357)	54.4	337	A	[1947STU]
C ₆ H ₅ NO ₃	[554-84-7]	3-nitrophenol				
	$\Delta_{\text{fus}}H$		18.06	369		[2002MUS/RAZ]
	$\Delta_{\text{fus}}H$		19.2	370		[1982POE/FAN]
	$\Delta_{\text{fus}}H$		19.19	371.2		[1991ACR, 1994SAB/GOU]
	$\Delta_{\text{sub}}H$		91.2 ± 0.5	298	C	[1994SAB/GOU]
	$\Delta_{\text{sub}}H$		98.5 ± 0.6	321	ME	[1992RIB/REI]
	$\Delta_{\text{sub}}H$		100.2 ± 0.6	298		[1992RIB/REI]
	$\Delta_{\text{sub}}H$	(305–334)	76.2	319.5	A	[1987STE/MAL]
C ₆ H ₅ NO ₃	[100-02-7]	4-nitrophenol				
	$\Delta_{\text{fus}}H$		11.0	386.4		[2002MUS/RAZ]
	$\Delta_{\text{fus}}H$		18.25	388.2		[1996DOM/HEA, 1994SAB/GOU]
	$\Delta_{\text{sub}}H$		92.4	298	C	[1994SAB/GOU]
	$\Delta_{\text{sub}}H$	(305–352)	98.8 ± 1	298	ME	[1971PAR/ROC]
C ₆ H ₅ NO ₃	[824-40-8]	pyridine-2-carboxylic acid N-oxide				
	$\Delta_{\text{sub}}H$	(345–392)	94.4 ± 4.0	298	ME	[1998RIB/MAT]
	C ₆ H ₅ NO ₃	[2398-81-4]	pyridine-3-carboxylic acid N-oxide			
$\Delta_{\text{sub}}H$			152.3 ± 1.9	298	ME	[1995ACR/TUC, 1995ABB/JIM]
C ₆ H ₅ NO ₃	[13602-12-5]	pyridine-4-carboxylic acid N-oxide				
	$\Delta_{\text{sub}}H$	(345–392)	136.1 ± 1.2	298	ME	[1998RIB/MAT]
C ₆ H ₅ NO ₃	[84522-17-8]	(2-furyl)oxoacetamide				
	$\Delta_{\text{fus}}H$ (I)		22	373.4		
	$\Delta_{\text{fus}}H$ (II)		17.8	367.2	DSC	[2008BAR/BER]
C ₆ H ₅ NO ₃	[609-71-2]	2-hydroxynicotinic acid				
	$\Delta_{\text{sub}}H$	(433–461)	125.4 ± 5.0	447	ME	[2009SAN/FIG]
	$\Delta_{\text{sub}}H$	(433–461)	128.3 ± 5.1	298	ME	[2009SAN/FIG]
C ₆ H ₅ NO ₃	[609-70-1]	4-hydroxynicotinic acid				
	$\Delta_{\text{sub}}H$	(425–456)	144.6 ± 3.6	441	ME	[2009SAN/FIG]
	$\Delta_{\text{sub}}H$	(425–456)	148.1 ± 3.7	298	ME	[2009SAN/FIG]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₆ H ₅ NO ₃	[27828-71-3]	5-hydroxynicotinic acid				
	$\Delta_{\text{sub}}H$	(436–458)	147.1 ± 7.0	447	ME	[2009SAN/FIG]
		(436–458)	149.8 ± 7.1	298	ME	[2009SAN/FIG]
C ₆ H ₅ NO ₃	[5006-66-6]	6-hydroxynicotinic acid				
	$\Delta_{\text{sub}}H$	(461–488)	143.0 ± 4.5	475	ME	[2009SAN/FIG]
		(461–488)	146.4 ± 4.6	298	ME	[2009SAN/FIG]
C ₆ H ₅ NO ₄	[601-89-8]	2-nitro-1,3-dihydroxybenzene				
	$\Delta_{\text{sub}}H$	(253–293)	74.5	273		[1958HOY/PEP]
C ₆ H ₅ NO ₄	[3316-09-4]	4-nitrocatechol				
	$\Delta_{\text{sub}}H$		121.1 ± 1.4		C	[1986RIB/RIB]
C ₆ H ₅ NO ₅	[na]	methyl 5-nitro-2-furancarboxylate				
	$\Delta_{\text{sub}}H$		104.2 ± 2.1			[1980BAL/LEB, 1986PED/NAY]
C ₆ H ₅ NS	[20893-30-5]	2-thiopheneacetonitrile				
	Δ_vH		60.5 ± 1.3	298	C	[2008RIB/SAN]
C ₆ H ₅ NS	[13781-53-8]	3-thiopheneacetonitrile				
	Δ_vH		61.1 ± 1.3	298	C	[2008RIB/SAN]
C ₆ H ₅ NS	[55406-13-8]	3-methyl-2-thiophenecarbonitrile				
	Δ_vH		54.4 ± 1.2	298	C	[2008RIB/SAN]
C ₆ H ₅ N ₃	[622-37-3]	phenyl azide				
	Δ_vH	(348–368)	45.2	358	A	[1987STE/MAL, 1972DYK]
C ₆ H ₅ N ₃	[95-14-7]	1- <i>H</i> -benzotriazole				
	$\Delta_{\text{fus}}H$		7.7	369.9		[1999SAB/PER]
	$\Delta_{\text{sub}}H$		98.2 ± 0.7	298	C	[1999SAB/PER]
	$\Delta_{\text{sub}}H$		99.0 ± 0.5	298	ME	[1989JIM/ROU]
	$\Delta_{\text{sub}}H$		97.9			[1961ZIM/GEI]
C ₆ H ₅ N ₅ O ₆	[28930-29-2]	1,3-diamino-2,4,6-trinitrobenzene				
	$\Delta_{\text{sub}}H$		146.9		DSC	[1990HWA/YOS]
	$\Delta_{\text{sub}}H$	(335–382)	140	350	LE	[1987STE/MAL, 1969ROS/DIC]
	$\Delta_{\text{sub}}H$		143.5	298		[1978CUN/PAL]
			110.9		DSC	[1990HWA/YOS]
C ₆ H ₆	[71-43-2]	benzene				
	$\Delta_{\text{fus}}H$		9.87	278.7	C	[1996DOM/HEA, 1926AND/LYN, 1942ZIE/AND]
	$\Delta_{\text{sub}}H$	(258–273)	41.7			[1994LIU/DIC]
	$\Delta_{\text{sub}}H$	(223–279)	45.2	264	A	[1987STE/MAL, 1976HA/MOR]
	$\Delta_{\text{sub}}H$		45.1	278		[1984HES/WIS]
	$\Delta_{\text{sub}}H$	(183–197)	44.4	298	TE,ME	[1980DEK]
	$\Delta_{\text{sub}}H$		53.9 ± 0.8	193		[1977DEK/VAN]
	$\Delta_{\text{sub}}H$		49.4 ± 0.4	193		[1977DEK/VAN]
	$\Delta_{\text{sub}}H$	(221–268)	45.6	279	MM	[1974JAC]
	$\Delta_{\text{sub}}H$		44.1	261		[1960JON]
	$\Delta_{\text{sub}}H$		43.1	229		[1960JON]
	$\Delta_{\text{sub}}H$		44.6	279		[1956MIL]
	$\Delta_{\text{sub}}H$	(263–270)	46.6	282	A	[1947STU]
	$\Delta_{\text{sub}}H$		44.6	273		[1936DEB, 1974JAC]
	$\Delta_{\text{sub}}H$	(184–200)	U 33.2	192		[1933DEI]
$\Delta_{\text{sub}}H$	(214–238)	43.3	226	A	[1913MUN]	

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound					
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference	
		$\Delta_v H$	(305–345)	33.2	320		[2002LUB/BAN]
		$\Delta_v H$	(258–313)	35.6		GC	[1994LIU/DIC]
		$\Delta_v H$	(296–377)	33.5	311	EB	[1990AMB/EWI]
		$\Delta_v H$		33.9 ± 0.2		GC	[1989AZA]
		$\Delta_v H$		33.4	307	C	[1988DON/LIN]
		$\Delta_v H$		33.1	314	C	[1988DON/LIN]
		$\Delta_v H$		32.4	324	C	[1988DON/LIN]
		$\Delta_v H$		31.9	332	C	[1988DON/LIN]
		$\Delta_v H$		31.4	344	C	[1988DON/LIN]
		$\Delta_v H$		30.6	353	C	[1988DON/LIN]
		$\Delta_v H$	(279–377)	34.4	294	A	[1987STE/MAL]
		$\Delta_v H$	(353–422)	31.5	368	A	[1987STE/MAL]
		$\Delta_v H$	(420–502)	30.2	435	A	[1987STE/MAL]
		$\Delta_v H$	(501–562)	30.3	516	A	[1987STE/MAL]
		$\Delta_v H$		30.8	352		[1983NAT/VIS]
		$\Delta_v H$		30.5	361		[1983NAT/VIS]
		$\Delta_v H$		30.2	366		[1983NAT/VIS]
		$\Delta_v H$	(313–373)	35.3	343		[1983TSO/WIL]
		$\Delta_v H$		31	350		[1977RAO/VIU]
		$\Delta_v H$		33.8 ± 0.1	298	C	[1973SVO/VES]
		$\Delta_v H$		33.0 ± 0.1	313	C	[1973SVO/VES]
		$\Delta_v H$		32.2 ± 0.1	328	C	[1973SVO/VES]
		$\Delta_v H$		31.8 ± 0.1	333	C	[1973SVO/VES]
		$\Delta_v H$		31.4 ± 0.1	343	C	[1973SVO/VES]
		$\Delta_v H$		30.9 ± 0.1	353	C	[1973SVO/VES]
		$\Delta_v H$		32.6 ± 0.4	313	DSC	[1971MIT/IMA]
		$\Delta_v H$		32.5 ± 0.5	328	DSC	[1971MIT/IMA]
		$\Delta_v H$		33.9	298		[1971WIL/ZWO]
		$\Delta_v H$		31.6 ± 0.4	345	DSC	[1971MIT/IMA]
		$\Delta_v H$		34.1	293		[49YAR/FED]
		$\Delta_v H$	(284–354)	34.1	299		[1949FOR/NOR]
		$\Delta_v H$		33.8	298	C	[1947OSB/GIN]
		$\Delta_v H$	(282–354)	34.1	297		[1946THO]
		$\Delta_v H$		31.2	294		[1946SCO/BRI]
		$\Delta_v H$	(288–354)	34.1	303	MM	[1945WIL/TAY]
		$\Delta_v H$	(298–373)	33.4	313	EB	[1941SMI]
		$\Delta_v H$	(273–348)	34.5	288		[1940STU/SAY]
		$\Delta_v H$		34	298		[1927NAG]
C₆D₆	[1076-43-3]	benzene-d ₆					
		$\Delta_{\text{fus}}H$		9.79	279.9		[1942ZIE/AND]
		$\Delta_v H$	(283–352)	34.2	298		[1953DAV/SCH]
C₆H₆	[821-08-9]	1,5-hexadien-3-yne					
		$\Delta_v H$	(223–357)	40.4	238	A	[1987STE/MAL]
C₆H₆	[10420-90-3]	1,3-hexadien-5-yne					
		$\Delta_v H$	(223–303)	44.0	238	A	[1987STE/MAL]
C₆H₆	[2809-69-0]	2,4-hexadiyne					
		$\Delta_{\text{trs}}H$		1.0	118		[1982BAL/MRA]
		$\Delta_{\text{sub}}H$	(282–333)	47 ± 2	307	MM	[1982BAL/MRA]
		$\Delta_v H$	(364–408)	42.5	298	EB	[1986MEY/MEY]
C₆H₆BrN	[615-36-1]	2-bromoaniline					

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
			16.14	304.1	DSC	[2006RIB/FER]
			20.04	305		[1983KHA/KHE]
			75.0 ± 1.4	298	C	[2006RIB/FER]
C ₆ H ₆ BrN	[591-19-5]	3-bromoaniline				
			14.68	291		[1983KHA/KHE]
			63.4 ± 1.5	298	C	[2006RIB/FER]
C ₆ H ₆ BrN	[106-40-1]	4-bromoaniline				
			16.75	336	DSC	[2006RIB/FER]
			13.36	336		[1983KHA/KHE]
C ₆ H ₆ ClN			79.4 ± 1.7	298	C	[2006RIB/FER]
	[95-51-2]	2-chloroaniline				
			12.38	269.2		[2007STR/RUZ]
C ₆ H ₆ ClN			8.81	271		[1983KHA/KHE]
			56.4 ± 1.6	298	C	[2005RIB/GOM]
		(288–327)	57.1 ± 0.5	298	GS	[2003VER/SCH]
		(397–482)	50.7	412	A	[1987STE/MAL]
		(287–336)	58.2 ± 1.4	311	TE,ME	[1985PIA/SCA]
		(294–330)	57.1 ± 1.0	312	TE,ME	[1985PIA/SCA]
C ₆ H ₆ ClN	[108-42-9]	3-chloroaniline				
			12	263		[1983KHA/KHE]
			61.1 ± 2.8	298	C	[2005RIB/GOM]
		(291–340)	60.2 ± 0.1	298	GS	[2003VER/SCH]
		(398–573)	53.6	413	A	[1987STE/MAL, 1972DYK]
		(292–346)	60.3 ± 0.6	319	TE,ME	[1985PIA/SCA]
		(304–342)	61.0 ± 0.8	323	TE,ME	[1985PIA/SCA]
C ₆ H ₆ ClN	[106-47-8]	4-chloroaniline				
			21.06	343.5		
			20.47	342.8		[2003VER/SCH, 2007STR/RUZ]
			16.9	344		[1983KHA/KHE]
		(291–337)	80.5 ± 0.3	298	GS	[2003VER/SCH]
		(283–303)	90.7	293	ME	[1987STE/MAL, 1925SWA/MAC]
		(346–374)	62.3 ± 0.5	298	GS	[2003VER/SCH]
C ₆ H ₆ Cl ₄	[41992-55-6]	α -3,4,5,6-tetrachlorocyclohexene				
		(353–399)	58.0	368	A	[1987STE/MAL]
C ₆ H ₆ Cl ₆	[319-84-6]	α -hexachlorocyclohexane				
		(313–363)	95.7	328	A	[1987STE/MAL, 1960JON]
		(324–344)	92.9	334	TE	[1947BAL]
C ₆ H ₆ Cl ₆			68.5	398	GC	[1990HIN/BID2]
	[319-85-7]	β -hexachlorocyclohexane (mp 314 °C)				
		(506–551)	103.7			[1989LUB/JAN]
C ₆ H ₆ Cl ₆			107	328	A	[1987STE/MAL, 1960JON]
		(368–390)	102.9	379	TE	[1947BAL]
C ₆ H ₆ Cl ₆	[58-89-9]	γ -hexachlorocyclohexane				
		(310–384)	92.4 ± 4.0	298	ME,TE	[1998GIU/BRU]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{sub}}H$	(292–326)	97.7 ± 0.6	308	ME	[1996BOE/MAR]
	$\Delta_{\text{sub}}H$	(243–303)	106.6 ± 0.9	273	GS	[1994WAN/SHU]
	$\Delta_{\text{sub}}H$		90.1 ± 0.7	338	C	[1991SAB/AN]
	$\Delta_{\text{sub}}H$		90.8 ± 0.7	298	C	[1991SAB/AN]
	$\Delta_{\text{sub}}H$	(313–363)	99.2	328	A	[1987STE/MAL, 1960JON]
	$\Delta_{\text{sub}}H$	(293–313)	88.9	303	GS	[1983SPE/CLI, 1970SPE/CLI]
	$\Delta_{\text{sub}}H$	(293–313)	101.2	303		[1970SPE/CLI]
	$\Delta_{\text{sub}}H$	(313–343)	89.7	328		[1960SCH/LEG]
	$\Delta_{\text{sub}}H$	(333–365)	115.5		TE	[1947BAL]
	Δ_vH	(343–453)	70.5	398	GC	[1990HIN/BID2]
C₆H₆Cl₆	[319-86-8]	δ -hexachlorocyclohexane (mp 142 °C)				
	$\Delta_{\text{sub}}H$	(313–363)	97.3	328	A	[1987STE/MAL, 1960JON]
	$\Delta_{\text{sub}}H$	(328–358)	97.5			[1947BAL]
C₆H₆Cl₆	[na]	1 α , 2 α , 3 β , 4 α , 5 α , 6 β -hexachlorocyclohexane				
	$\Delta_{\text{fus}}H$		22.13	386.8	DSC	[1990DON/DRE]
C₆H₆Cl₆	[na]	1 α , 2 α , 3 β , 4 α , 5 α , 6 β -hexachlorocyclohexane (lindane)				
	$\Delta_{\text{fus}}H$		15.9	388.9	DSC	[1969PLA.GLA]
C₆H₆FN	[348-54-9]	2-fluoroaniline				
	Δ_vH		52.0 ± 0.6	298	C	[2007RIB/FER]
C₆H₆FN	[372-19-0]	3-fluoroaniline				
	Δ_vH		54.7 ± 0.6	298	C	[2007RIB/FER]
C₆H₆FN	[371-40-4]	4-fluoroaniline				
	Δ_vH		54.8 ± 0.8	298	C	[2007RIB/FER]
C₆H₆F₈O	[77527-96-9]	1,1,2,2,3,3,4,4-octafluoro-5-methoxypentane				
	Δ_vH	(293–396)	44.8	308	I	[2002MUR/YAM]
C₆H₆F₈O₂	[355-74-8]	2,2,3,3,4,4,5,5-octafluoro-1,6-hexanediol				
	$\Delta_{\text{sub}}H$		89.2 ± 8.4			[1974COX, 1977PED/RYL]
C₆H₆F₈O₃	[485399-46-0]	1, 1'-oxybis[1,1,2,2-tetrafluoro-2-methoxyethane]				
	Δ_vH	(280–370)	33.4			[2005MAR/AVA]
C₆H₆F₉N₃S	[63265-76-9]	N-[N, N'-dimethyl-S-(trifluoromethyl)sulfonodiimidoyl]-1,1,1,3,3,3-hexafluoro-2-propanimine				
	Δ_vH		32.6	426	I	[1977KIT/SHR2]
C₆H₆IN	[615-43-0]	2-iodoaniline				
	$\Delta_{\text{fus}}H$		19.38	329.6		[2006RIB/FER2]
	$\Delta_{\text{fus}}H$		13.95	333		[1983KHA/KHE]
	$\Delta_{\text{sub}}H$		81.3 ± 1.4	298	C	[2006RIB/FER2]
C₆H₆IN	[626-01-7]	3-iodoaniline				
	$\Delta_{\text{fus}}H$		14.5	298		[1983KHA/KHE]
	Δ_vH		67.5 ± 1.4	298	C	[2006RIB/FER2]
C₆H₆IN	[540-37-4]	4-iodoaniline				
	$\Delta_{\text{fus}}H$		16.94	336		[2006RIB/FER2]
	$\Delta_{\text{fus}}H$		15.1	334		[1983KHA/KHE]
	$\Delta_{\text{sub}}H$		84.8 ± 1.4	298	C	[2006RIB/FER2]
C₆H₆N₂	[1119-85-3]	3-hexenedinitrile				
	Δ_vH	(353–448)	49.4	368	A	[1987STE/MAL]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	Enthalpy							
C ₆ H ₆ N ₂ O	[1452-77-3]	2-pyridinecarboxamide			16.82	381		[1960NEG/MIK2]
	$\Delta_{\text{fus}}H$							
	$\Delta_{\text{sub}}H$			93.1 ± 3.3	298		C	[2001RIB/GON]
				(323–373)	93.1	338	ME	[1987STE/MAL, 1960NEG/MIK, 1959HAR]
C ₆ H ₆ N ₂ O	[98-92-0]	3-pyridinecarboxamide (nicotinamide)			23.8	403.8	DSC	[2009GOO/ROD]
	$\Delta_{\text{fus}}H$				25.5	401.6	DSC	[2008NIC/BEL]
	$\Delta_{\text{fus}}H$				26.94	402		[1960NEG/MIK2]
	$\Delta_{\text{sub}}H$			121.2 ± 3.3	298		C	[2001RIB/GON]
	$\Delta_{\text{sub}}H$			(363–393)	111.8	378	ME	[1987STE/MAL, 1960NEG/MIK, 1959HAR]
C ₆ H ₆ N ₂ O	[1453-82-3]	4-pyridinecarboxamide			26.81	431		[1960NEG/MIK2]
	$\Delta_{\text{fus}}H$							
	$\Delta_{\text{sub}}H$			116.1 ± 1.5	298		C	[2001RIB/GON]
				(383–412)	99.9	397.5	ME	[1987STE/MAL, 1960NEG/MIK, 1959HAR]
C ₆ H ₆ N ₂ O	[na]	2-pyridinealdoxime			19.98	388		[2007SHI/TAN3]
	$\Delta_{\text{fus}}H$							
C ₆ H ₆ N ₂ O ₂	[88-74-4]	2-nitroaniline			16.11	344.4		[1996DOM/HEA]
	$\Delta_{\text{fus}}H$							
	$\Delta_{\text{sub}}H$			(313–342)	89.0 ± 0.7	298	GS	[1997VER3]
	$\Delta_{\text{sub}}H$			(313–342)	87.2 ± 0.7	328	GS	[1997VER3]
	$\Delta_{\text{sub}}H$				90 ± 3.0		ME, TE	[1985FER/PIA]
	$\Delta_{\text{sub}}H$				82.4 ± 2	313		[1938WOL/WEG, 1960JON, 1935TRI]
	$\Delta_{\text{sub}}H$				90 ± 4.2			[1958HOY/PEP, 1970COX/PIL]
	$\Delta_{\text{sub}}H$			(310–319)	79.9 ± 1.7			[1934WOL/TRI]
				(423–553)	59.3	438	A	[1987STE/MAL]
				(377–558)	64.8	392		[1947STU]
C ₆ H ₆ N ₂ O ₂	[99-09-2]	3-nitroaniline			23.68	387.2		[1996DOM/HEA]
	$\Delta_{\text{fus}}H$							
	$\Delta_{\text{sub}}H$				108.3 ± 3		ME, TE	[1985FER/PIA]
	$\Delta_{\text{sub}}H$			(320–384)	93.6 ± 0.7	351	ME	[1973MAL/GIG2]
	$\Delta_{\text{sub}}H$			(320–384)	94.6 ± 0.3	351	C	[1973MAL/GIG2]
	$\Delta_{\text{sub}}H$				96.5 ± 0.3	298	C	[1973MAL/GIG2]
	$\Delta_{\text{sub}}H$			(288–343)	97.6	316	ME	[1958HOY/PEP, 1970COX/PIL]
	$\Delta_{\text{sub}}H$			(332–341)	88.3 ± 1.7		TE	[1934WOL/TRI]
	$\Delta_{\text{sub}}H$			(332–341)	88.7 ± 2.5			[1938WOL/WEG, 1960JON, 1935TRI]
				(443–578)	64.9	458	A	[1985FER/PIA]
C ₆ H ₆ N ₂ O ₂	[100-01-6]	4-nitroaniline			21.09	420.2		[1996DOM/HEA]
	$\Delta_{\text{fus}}H$							
	$\Delta_{\text{sub}}H$				101.4 ± 1.3	298	ME	[1990RIB/RIB]
	$\Delta_{\text{sub}}H$				101.5 ± 1.7	298	TE	[1990RIB/RIB]
	$\Delta_{\text{sub}}H$				94.6		GS	[1987SHI/OHK, 1991HOR]
	$\Delta_{\text{sub}}H$				107 ± 3		ME, TE	[1985FER/PIA]
	$\Delta_{\text{sub}}H$				100.4 ± 2.1	298	ME	[1977FRA, 1990RIB/RIB]
					100.9 ± .6	298	ME	[1973MAL/GIG2]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		T _m (K)	Method	Reference	
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)				
		$\Delta_{\text{sub}}H$		101.3 ± 0.7	298	C	[1973MAL/GIG2]
		$\Delta_{\text{sub}}H$	(303–363)	109.3	333	ME	[1958HOY/PEP, 1970COX/PIL]
		$\Delta_{\text{sub}}H$		99.3 ± 1.7	298	ME	[1956MAJ]
		$\Delta_{\text{sub}}H$	(346–366)	97.5 ± 1.7	356	ME	[1956MAJ]
		$\Delta_{\text{sub}}H$		100.7 ± 2.5	298	TE	[1938WOL/WEG]
		$\Delta_{\text{sub}}H$		98.7 ± 2.5	361	TE	[1938WOL/WEG, 1960JON]
		$\Delta_{\text{sub}}H$	(357–367)	103.3 ± 1.7	362		[1934WOL/TRI]
		Δ_vH	(473–538)	77.9	488	A	[1987STE/MAL]
		Δ_vH	(415–609)	70.0	430		[1947STU]
C ₆ H ₆ N ₂ O ₂	[1986-81-8]		3-pyridinecarboxamide N-oxide				
		$\Delta_{\text{sub}}H$		119.2 ± 2.3	298	ME	[2001RIB/GON]
C ₆ H ₆ N ₂ O ₂	[38557-82-3]		4-pyridinecarboxamide N-oxide				
		$\Delta_{\text{sub}}H$		125.3 ± 1.8	298	ME	[2001RIB/GON]
C ₆ H ₆ N ₂ O ₂	[5521-55-1]		2-methyl-5-pyrazine carboxylic acid				
		$\Delta_{\text{sub}}H$		100.9 ± 1.5	298	C	[1997ACR/POW]
C ₆ H ₆ N ₂ O ₂	[1445-69-8]		phthalhydrazide				
		$\Delta_{\text{sub}}H$	(428–450)	132.7 ± 0.7	439	ME	[2008RIB/CAB2]
		$\Delta_{\text{sub}}H$	(428–450)	139.8 ± 0.7	298	ME	[2008RIB/CAB2]
C ₆ H ₆ N ₂ O ₃	[1074-98-2]		3-methyl-4-nitropyridine N-oxide				
		$\Delta_{\text{sub}}H$	(345–392)	106.7 ± 2.0	298	ME	[1998RIB/MAT]
C ₆ H ₆ N ₄ O	[1006-08-02]		7-methylhypoxanthine				
		$\Delta_{\text{sub}}H$		100.4 ± 13			[1978NOW/SZC]
C ₆ H ₆ N ₄ O	[875-31-0]		9-methylhypoxanthine				
		$\Delta_{\text{sub}}H$	(500–552)	84.0		HS	[1965CLA/PES]
C ₆ H ₆ N ₄ O ₄	[na]		(2,4-dinitrophenyl)hydrazine				
		$\Delta_{\text{fus}}H$		18.89	474.1		[2002MUS/RAZ]
C ₆ H ₆ N ₆ O ₆	[3058-38-6]		2,4,6-trinitro-1,3,5-benzenetriamine				
		$\Delta_{\text{sub}}H$		182.4		DCA	[1990HWA/YOS]
		$\Delta_{\text{sub}}H$		168	423		[2008RAI/BHA, 1979GAR/LAW]
		$\Delta_{\text{sub}}H$	(402–451)	168.2	417	LE	[1987STE/MAL, 1969ROS/DIC]
C ₆ H ₆ N ₆ O ₁₄	[866-65-9]		2,2,2-trinitroethyl 4,4,4-trinitrobutyrate				
		$\Delta_{\text{trs}}H$		25.94	362.7		
		$\Delta_{\text{fus}}H$		6.69	366.5	DSC	[1971ROS/HOL]
C ₆ H ₆ N ₁₂ O ₁₂	[135285-90-4]		hexanitrohexaazaisowurtzitane				
		$\Delta_{\text{trs}}H$		1.97	428.2		
		$\Delta_{\text{trs}}H$		5.91	435.2		
		$\Delta_{\text{fus}}H$		NA			
		$\Delta_{\text{trs}}H$		7.25	442.2		
		$\Delta_{\text{fus}}H$		NA			[2005TUR/VAC, 1998LOB/BOH]
C ₆ H ₆ O	[108-95-2]		phenol				
		$\Delta_{\text{fus}}H$		11.51	314		[1972INO/LIA, 1996DOM/HEA]
		$\Delta_{\text{sub}}H$	(263–298)	65.3 ± 3.3	280	HSA	[1975CHI]
		$\Delta_{\text{sub}}H$	(230–273)	69.7 ± 0.9	298	ME	[1971PAR/ROC]
		$\Delta_{\text{sub}}H$	(282–313)	68.7 ± 0.5		GS	[1960AND/BID, 1970COX/PIL]
		$\Delta_{\text{sub}}H$	(283–303)	68.2	293	ME	[1958SKL/MAR]
		$\Delta_{\text{sub}}H$	(270–313)	68.1	292		[1948NIT/SEK2]
		$\Delta_{\text{sub}}H$	(278–305)	67.8		TE	[1947BAL, 1960JON]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound						
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference		
		$\Delta_v H$	(363–391)	53.2	378	EB	[2001CHY/FRA]	
		$\Delta_v H$	(393–433)	58.8	298	CGC	[1995CHI/HOS]	
		$\Delta_v H$	(455–655)	49.5	470	A	[1987STE/MAL]	
		$\Delta_v H$	(314–395)	57.4	329	A	[1987STE/MAL]	
		$\Delta_v H$	(387–456)	50.9	402	A	[1987STE/MAL]	
		$\Delta_v H$	(449–526)	46.8	464	A	[1987STE/MAL]	
		$\Delta_v H$	(520–625)	43.8	535	A	[1987STE/MAL]	
		$\Delta_v H$		51.1			[1986BAL/GNA]	
		$\Delta_v H$	(383–473)	51.3	398	EB,GS	[1987STE/MAL, 1960AND/BID, 1972DYK]	
		$\Delta_v H$	(380–455)	51.4	395		[1949DRE/SHR, 1949DRE/MAR, 1984BOU/FRI]	
		$\Delta_v H$	(414–454)	48.1	434		[1939GOL/MAR]	
C₆H₆O	[1192-62-7]	2-acetylfuran						
		$\Delta_{\text{fus}}H$		15.5	301.6		[2009FLO/CAM]	
		$\Delta_v H$		53.0 ± 0.6	298	C	[2009RIB/AMA]	
C₆H₆OS	[88-15-3]	2-acetylthiophene						
		$\Delta_v H$		58.8 ± 1.2	298	C	[2007ROU/TEM]	
C₆H₆OS	[1468-83-3]	3-acetylthiophene						
		$\Delta_{\text{fus}}H$		18.9	333.6		[2006TEM/ROU]	
C₆H₆OS	[1379-70-4]	5-methyl-2-thiophenecarboxaldehyde						
		$\Delta_v H$		57.7 ± 1.3	298	C	[2008RIB/SAN2]	
C₆H₆OS	[5834-16-2]	3-methyl-2-thiophenecarboxaldehyde						
		$\Delta_v H$		56.2 ± 1.2	298	C	[2008RIB/SAN2]	
C₆H₆O₂	[120-80-9]	1,2-dihydroxybenzene (catechol)						
		$\Delta_{\text{fus}}H$		22.87	377.6		[2008VER/KOZ]	
		$\Delta_{\text{fus}}H$		18.55	377.6		[2000VER/SCH]	
		$\Delta_{\text{fus}}H$		22.54	377.7	DSC	[1997LEE/CHA]	
		$\Delta_{\text{fus}}H$		22.01	376.9		[1989BRE/LIC]	
		$\Delta_{\text{fus}}H$		22.76	377.5	C	[1926AND/LYN]	
		$\Delta_{\text{sub}}H$	(295–310)	80.0 ± 0.5	302		[2006CHE/OJA]	
		$\Delta_{\text{sub}}H$		87.5 ± 0.3	298	C	[1991SAB/BUL]	
		$\Delta_{\text{sub}}H$		86.6 ± 1.6	298	C	[1984CAR]	
		$\Delta_{\text{sub}}H$		80.8			[1938WOL/WEG, 1960JON, 1935TRI]	
		$\Delta_v H$	(378–389)	71.9 ± 0.8	298	GS	[2008VER/KOZ]	
		$\Delta_v H$	(395–519)	63.1	410	A	[1987STE/MAL]	
		$\Delta_v H$	(378–439)	61.2	393	GC	[1975KUN/LIL]	
C₆H₆O₂	[108-46-3]	1,3-dihydroxybenzene (resorcinol)						
		$\Delta_{\text{trs}}H$		1.2	366.8			
		$\Delta_{\text{fus}}H$		18.9	382.6		[1989BRE/LIC]	
		$\Delta_{\text{fus}}H$		21.3	382.9	C	[1926AND/LYN]	
		$\Delta_{\text{sub}}H$		85.3 ± 0.5	334	C	[1991SAB/BUL]	
		$\Delta_{\text{sub}}H$		87.5 ± 0.5	298	C	[1991SAB/BUL]	
		$\Delta_{\text{sub}}H$	(328–379)	92.3	353	GS	[1983BEN/BIE]	
		$\Delta_{\text{sub}}H$	(324–335)	93.3 ± 21			[1968DES/WIL]	
		$\Delta_{\text{sub}}H$	(283–323)	93.4	303		[1958HOY/PEP]	
		$\Delta_{\text{sub}}H$		95.4 ± 1.7			[1938WOL/WEG, 1960JON, 1935TRI]	
		$\Delta_v H$		78.4 ± 1.3	298		[2008VER/KOZ]	

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(419–550)	74.3	434	A	[1987STE/MAL]
	$\Delta_v H$	(392–463)	74.3	407	GC	[1987STE/MAL, 1975KUN/LIL]
C ₆ H ₆ O ₂	[123-31-9]	1,4-dihydroxybenzene (hydroquinone)				
	$\Delta_{\text{fus}}H$		27.23	445.1		[2008VER/KOZ]
	$\Delta_{\text{fus}}H$		27.2	NA	DSC	[1999VER7]
	$\Delta_{\text{fus}}H$		26.48	453		[1989BRE/LIC]
	$\Delta_{\text{fus}}H$		27.11	445.1	C	[1926AND/LYN]
	$\Delta_{\text{sub}}H$	(325–339)	100.6 ± 1.3	332		[2006CHE/OJA]
	$\Delta_{\text{sub}}H$		94.1 ± 0.5	298	C	[1991SAB/BUL]
	$\Delta_{\text{sub}}H$		93.7 ± 0.5	334	C	[1991SAB/BUL]
	$\Delta_{\text{sub}}H$	(341–400)	101.3		GS	[1983BEN/BIE]
	$\Delta_{\text{sub}}H$		103.9 ± 1	342	ME,TE	[1981DEK/SMI]
	$\Delta_{\text{sub}}H$	(298–346)	103.8	313		[1956MAG]
	$\Delta_{\text{sub}}H$		90.1 ± 0.8			[1953SEK/SUZ]
	$\Delta_{\text{sub}}H$	(326–345)	103.8		QF	[1927COO/COO]
	$\Delta_v H$		84.4 ± 0.7	298		[2008VER/KOZ]
$\Delta_v H$	(448–559)	70.5	463	A	[1987STE/MAL]	
(C ₆ H ₆ O ₂)–(C ₁₀ H ₈ O ₂)	[60706-28-7]	1,4-hydroquinone-1,4-naphthoquinone				
$\Delta_{\text{sub}}H$		98.7 ± 1	324	TE,ME	[1981DEK/SMI]	
C ₆ H ₆ O ₂ S	[1918-77-0]	2-thiopheneacetic acid				
	$\Delta_{\text{fus}}H$		14.0	337.4	DSC	[2006TEM/ROU]
$\Delta_{\text{sub}}H$	(292–307)	97.5 ± 1.4	298	ME	[2008TEM/ROU2]	
C ₆ H ₆ O ₂ S	[6964-21-2]	3-thiopheneacetic acid				
	$\Delta_{\text{fus}}H$		18.8	353.3	DSC	[2006TEM/ROU]
$\Delta_{\text{sub}}H$	(294–312)	100.9 ± 1.9	298	ME	[2008TEM/ROU2]	
C ₆ H ₆ O ₂ S	[5380-42-7]	methyl 2-thiophenecarboxylate				
	$\Delta_v H$		57.6 ± 1.2	298	C	[2009RIB/SAN2]
C ₆ H ₆ O ₂ S	[23806-24-8]	methyl 2-thiophenecarboxylate				
	$\Delta_{\text{sub}}H$	(312–334)	96.7 ± 0.4	323.2	ME	[2008RIB/SAN5]
$\Delta_{\text{sub}}H$	(312–334)	98.0 ± 0.4	298	ME	[2008RIB/SAN5]	
C ₆ H ₆ O ₂ S	[1918-79-2]	5-methyl-2-thiophenecarboxylic acid				
	$\Delta_{\text{sub}}H$	(316–338)	100.4 ± 0.3	327.2	ME	[2008RIB/SAN5]
$\Delta_{\text{sub}}H$	(316–338)	101.9 ± 0.3	298	ME	[2008RIB/SAN5]	
C ₆ H ₆ O ₃	[87-66-1]	1,2,3-trihydroxybenzene				
	$\Delta_{\text{fus}}H$		25.9	405.6		[2004VER/SCH]
	$\Delta_{\text{fus}}H$		18.55	407.2		[1992RAI/GEO]
	$\Delta_{\text{sub}}H$	(326–370)	104	298	GS	[2004VER/SCH]
	$\Delta_{\text{sub}}H$		116.9 ± 0.6	298	C	[1986RIB/RIB]
	$\Delta_{\text{sub}}H$	(377–398)	89.1	387		[1934HIR]
$\Delta_v H$	(425–582)	69.5	440	A	[1987STE/MAL, 1955VON/GEB]	
C ₆ H ₆ O ₃	[533-73-3]	1,2,4-trihydroxybenzene				
	$\Delta_{\text{fus}}H$		28.8	413.2		[2004VER/SCH]
	$\Delta_{\text{sub}}H$	(341–382)	124.2 ± 0.6	298	GS	[2004VER/SCH]
$\Delta_{\text{sub}}H$		119.8 ± 1.6	298	C	[1986RIB/RIB]	
C ₆ H ₆ O ₃	[108-73-6]	1,3,5-trihydroxybenzene				

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound					
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference	
		$\Delta_{\text{fus}}H$	34.5	491.8		[2004VER/SCH]	
		$\Delta_{\text{sub}}H$	(355–382)	135.5 ± 1.3	298	GS	[2004VER/SCH]
		$\Delta_{\text{sub}}H$		131.7 ± 1.0	298	C	[1986RIB/RIB]
		$\Delta_{\text{sub}}H$	(383–406)	127.9		TE,ME	[1983DEW/BOW]
C₆H₆O₃	[67-47-0]	5-hydroxymethylfurfural					
		$\Delta_{\text{fus}}H$	19.8	308.5	DSC	[2009VER/EME]	
		Δ_vH	(314–368)	83.4 ± 0.2	298	GS	[2009VER/EME]
C₆H₆O₄	[na]	butynedioic acid, dimethyl ester					
		Δ_vH	(273–460)	56.3	288	A	[1987STE/MAL, 1972DYK]
C₆H₆S	[108-98-5]	benzenethiol (thiophenol)					
		$\Delta_{\text{fus}}H$	11.48	258.2		[1996DOM/HEA]	
		Δ_vH	(333–471)	45.9	348		[1999DYK/SVO]
		Δ_vH		43.5			[1986BAL/GNA]
		Δ_vH	(385–486)	43.1	400	A,EB	[1987STE/MAL, 1966OSB/DOU, 1956SCO/MCC]
		Δ_vH		43.8 ± 0.1	375	C	[1956SCO/MCC]
		Δ_vH		42.6 ± 0.1	395	C	[1956SCO/MCC]
		Δ_vH		41.8 ± 0.1	407	C	[1956SCO/MCC]
		Δ_vH		41.3 ± 0.1	417	C	[1956SCO/MCC]
		Δ_vH	(324–440)	44.3	339		[1955VON/GEB, 1984BOU/FRI]
C₆H₇Cl₂N	[137-04-2]	2-chloroaniline hydrochloride					
		$\Delta_{\text{sub}}H$	(373–473)	77.6	388	A	[1987STE/MAL, 1975KON/SEL]
C₆H₇Cl₂N	[141-85-5]	3-chloroaniline hydrochloride					
		$\Delta_{\text{sub}}H$	(383–473)	71.3	398	A	[1987STE/MAL, 1975KON/SEL]
C₆H₇Cl₂N	[20265-96-7]	4-chloroaniline hydrochloride					
		$\Delta_{\text{sub}}H$	(373–483)	77.8	388	A	[1987STE/MAL, 1975KON/SEL]
C₆H₇Cl₃OS	[76619-93-7]	2,3,3-trichloro-2-propenethioic acid, O-propyl ester					
		Δ_vH	(383–433)	69.4		GC	[1980PIT/KIS]
C₆H₇FN₂O₂	[na]	1,3-dimethyl-5-fluorouracil					
		$\Delta_{\text{sub}}H$	(338–373)	119 ± 4		TE	[2002BRU/POR]
C₆H₇F₃N₂O₄	[400-58-8]	N-[N-(trifluoroacetyl)glycyl]glycine					
		$\Delta_{\text{sub}}H$	(273–423)	67.0	288	A	[1987STE/MAL, 1960WEY/KLI]
C₆H₇N	[15760-35-7]	3-methylenecyclobutanecarbonitrile					
		Δ_vH	(348–435)	45.9	366	BG	[1971HAL/BAL]
C₆H₇N	[31357-71-8]	bicyclo[2.1.0]pentane-1-carbonitrile					
		Δ_vH	(332–390)	41.8	343	BG	[1971HAL/BAL]
C₆H₇N	[62-53-3]	aniline					
		$\Delta_{\text{fus}}H$		10.54	267.1		[1972AHM/EAD2, 1996DOM/HEA]
		$\Delta_{\text{fus}}H$		10.92	267.3		[1942ZIE/AND]
		Δ_vH	(350–499)	51.0 ± 0.2	360	EB	[2002STE/CHI]
		Δ_vH	(350–499)	48.0 ± 0.2	400	EB	[2002STE/CHI]
		Δ_vH	(350–499)	45.2 ± 0.2	440	EB	[2002STE/CHI]
		Δ_vH	(350–499)	42.2 ± 0.4	480	EB	[2002STE/CHI]
		Δ_vH	(421–591)	45.8	444		[1992LEE/CHE]
		Δ_vH	(273–338)	52.2	288	A	[1987STE/MAL]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound					
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference	
		$\Delta_v H$	(304–485)	53.6	319	A	[1987STE/MAL]
		$\Delta_v H$	(373–458)	48.6	388	A	[1987STE/MAL]
		$\Delta_v H$	(455–523)	46.3	470	A	[1987STE/MAL]
		$\Delta_v H$	(313–386)	51.4	350		[1979MAH/SMI]
		$\Delta_v H$	(288–298)	52.9	293		[1968RAV/DAN]
		$\Delta_v H$	(304–457)	54.0	319		[1962HAT/DOU]
		$\Delta_v H$		53.0	333	C	[1962HAT/DOU]
C ₆ H ₇ N	[109-06-8]	2-methylpyridine					
		$\Delta_{\text{fus}}H$		9.72	206.5		[1996DOM/HEA]
		$\Delta_v H$	(308–441)	41.2 ± 0.1	320	EB	[1999CHI/KNI]
		$\Delta_v H$	(308–441)	38.8 ± 0.1	360	EB	[1999CHI/KNI]
		$\Delta_v H$	(308–441)	36.4 ± 0.1	400	EB	[1999CHI/KNI]
		$\Delta_v H$	(308–441)	33.7 ± 0.3	440	EB	[1999CHI/KNI]
		$\Delta_v H$	(323–373)	43.6	298	CGC	[1995CHI/HOS]
		$\Delta_v H$	(292–403)	42.0	307	EB	[1990LEN]
		$\Delta_v H$	(209–245)	46.9	230	A	[1987STE/MAL]
		$\Delta_v H$	(429–537)	36.5	444	A	[1987STE/MAL]
		$\Delta_v H$	(521–621)	35.4	536	A	[1987STE/MAL]
		$\Delta_v H$		42.5 ± 0.1	298	C	[1984MAJ/SVO2]
		$\Delta_v H$		41.6 ± 0.1	313	C	[1984MAJ/SVO2]
		$\Delta_v H$		40.7 ± 0.1	328	C	[1984MAJ/SVO2]
		$\Delta_v H$		39.8 ± 0.1	343	C	[1984MAJ/SVO2]
		$\Delta_v H$		38.3 ± 0.1	368	C	[1984MAJ/SVO2]
		$\Delta_v H$	(352–445)	39.1	367	EB,IP	[1987STE/MAL, 1968OSB/DOU]
		$\Delta_v H$	(352–442)	39.1	367	EB	[1987STE/MAL, 1963SCO/HUB]
		$\Delta_v H$		38.8 ± 0.1	359	C	[1963SCO/HUB]
		$\Delta_v H$		37.7 ± 0.1	379	C	[1963SCO/HUB]
		$\Delta_v H$		36.2 ± 0.1	402	C	[1963SCO/HUB]
		$\Delta_v H$	(337–403)	39.8	352	MG	[1953HER/MAR]
C ₆ H ₇ N	[108-99-6]	3-methylpyridine					
		$\Delta_{\text{fus}}H$		14.18	255		[1996DOM/HEA]
		$\Delta_{\text{sub}}H$	(225–255)	62.2	240		[1987STE/MAL]
		$\Delta_v H$	(342–373)	44.5 ± 2.0	298	CGC	[2009LIP/CHI2]
		$\Delta_v H$	(314–457)	43.2 ± 0.1	320	EB	[1999CHI/KNI]
		$\Delta_v H$	(314–457)	40.9 ± 0.1	360	EB	[1999CHI/KNI]
		$\Delta_v H$	(314–457)	38.6 ± 0.1	400	EB	[1999CHI/KNI]
		$\Delta_v H$	(314–457)	36.1 ± 0.2	440	EB	[1999CHI/KNI]
		$\Delta_v H$	(374–458)	40.1	389	A	[1987STE/MAL]
		$\Delta_v H$	(450–570)	37.7	465	A	[1987STE/MAL]
		$\Delta_v H$	(561–645)	36.8	576	A	[1987STE/MAL]
		$\Delta_v H$		44.6 ± 0.1	298	C	[1984MAJ/SVO2]
		$\Delta_v H$		43.6 ± 0.1	313	C	[1984MAJ/SVO2]
		$\Delta_v H$		42.7 ± 0.1	328	C	[1984MAJ/SVO2]
		$\Delta_v H$		42.0 ± 0.1	343	C	[1984MAJ/SVO2]
		$\Delta_v H$		40.4 ± 0.1	368	C	[1984MAJ/SVO2]
		$\Delta_v H$	(347–458)	41.3	362	EB,IP	[1987STE/MAL, 1968OSB/DOU]
		$\Delta_v H$	(347–458)	41.3	362	EB	[1987STE/MAL, 1963SCO/GOO]
		$\Delta_v H$		40.2 ± 0.1	372	C	[1963SCO/GOO]
		$\Delta_v H$		38.9 ± 0.1	393	C	[1963SCO/GOO]
		$\Delta_v H$		37.4 ± 0.1	417	C	[1963SCO/GOO]
		$\Delta_v H$	(354–418)	41.0	369	MG	[1953HER/MAR]
C ₆ H ₇ N	[108-89-4]	4-methylpyridine					

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound					
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference	
		$\Delta_{\text{sub}}H$	(213–239)	62.7	226	A	[1987STE/MAL]
		Δ_vH	(328–459)	43.4 ± 0.1	320	EB	[1999CHI/KN1]
		Δ_vH	(328–459)	41.1 ± 0.1	360	EB	[1999CHI/KN1]
		Δ_vH	(328–459)	38.8 ± 0.1	400	EB	[1999CHI/KN1]
		Δ_vH	(328–459)	36.2 ± 0.2	440	EB	[1999CHI/KN1]
		Δ_vH	(323–373)	44.7	298	CGC	[1995CHI/HOS]
		Δ_vH	(348–460)	41.4	363	A	[1987STE/MAL]
		Δ_vH	(348–347)	42.1	347	A	[1987STE/MAL]
		Δ_vH	(381–460)	40.0	396	A	[1987STE/MAL]
		Δ_vH	(452–573)	37.9	467	A	[1987STE/MAL]
		Δ_vH	(564–646)	37.2	579	A	[1987STE/MAL]
		Δ_vH		44.9 ± 0.1	298	C	[1984MAJ/SVO2]
		Δ_vH		43.9 ± 0.1	313	C	[1984MAJ/SVO2]
		Δ_vH		42.9 ± 0.1	328	C	[1984MAJ/SVO2]
		Δ_vH		42.1 ± 0.1	343	C	[1984MAJ/SVO2]
		Δ_vH		44.8 ± 0.1	298	C	[1981HOS/SCO]
		Δ_vH	(348–459)	41.4	363	EB,IP	[1987STE/MAL, 1968OSB/DOU]
		Δ_vH	(350–418)	41.3	365	MG	[1953HER/MAR]
C₆H₇N	[26555-56-5]		2-cyclopentene-1-carbonitrile				
		Δ_vH		44.9 ± 0.1	298	C	[1970PRO/KRE]
(C₆H₇N)–(SO₂)	[na]		aniline-sulfur dioxide complex				
		$\Delta_{\text{sub}}H$	(277–323)	82.1	300		[1931HIL]
C₆H₇NO	[95-55-6]		2-aminophenol				
		$\Delta_{\text{fus}}H$		31.4	443.2		[2001ROT/GLA]
		$\Delta_{\text{fus}}H$		21.72	447.6		[2003HUA, 2005HUA/TAN]
		$\Delta_{\text{fus}}H$		34.0	447.4		[1996DOM/HEA]
		$\Delta_{\text{sub}}H$		93.5 ± 0.8	332	C	[1996SAB/GOU]
		$\Delta_{\text{sub}}H$		95.3 ± 0.7	337	C	[1996SAB/GOU]
		$\Delta_{\text{sub}}H$		96.9 ± 0.6	298	C	[1996SAB/GOU]
		$\Delta_{\text{sub}}H$		103.9 ± 0.9	298	C	[1986NUN/BAR]
C₆H₇NO	[591-27-5]		3-aminophenol				
		$\Delta_{\text{fus}}H$		23.9	390.7		[2001ROT/GLA]
		$\Delta_{\text{fus}}H$		21.95	396.8		[2003HUA, 2005HUA/TAN]
		$\Delta_{\text{fus}}H$		22.98	399		[1991RAI/GEO]
		$\Delta_{\text{sub}}H$		98.8 ± 0.9	335	C	[1996SAB/GOU]
		$\Delta_{\text{sub}}H$		101.6 ± 0.9	298	C	[1996SAB/GOU]
		$\Delta_{\text{sub}}H$		104.7 ± 1.2	298	C	[1986NUN/BAR]
C₆H₇NO	[123-30-8]		4-aminophenol				
		$\Delta_{\text{fus}}H$		23.8	455.2		[2001ROT/GLA]
		$\Delta_{\text{fus}}H$		31.2	459.5		[1996DOM/HEA, 1989BRE/LIC]
		$\Delta_{\text{fus}}H$		26.0	462.5		[1996DOM/HEA, 1996SAB/GOU]
		$\Delta_{\text{sub}}H$		101.1 ± 0.7	335	C	[1996SAB/GOU]
		$\Delta_{\text{sub}}H$		103.6 ± 0.7	298	C	[1996SAB/GOU]
		$\Delta_{\text{sub}}H$	(423–459)	111.0	438		[1987STE/MAL]
		$\Delta_{\text{sub}}H$		109.1 ± 1.4	298	C	[1986NUN/BAR]
		$\Delta_{\text{sub}}H$	(403–430)	92.1	417	I	[1954DUN, 1960JON]
C₆H₇NO	[1121-25-1]		2-methyl-3-hydroxypyridine				
		$\Delta_{\text{sub}}H$		89.3 ± 1.3	298	C	[1982SUR/SAI, 1986PED/NAY]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		T _m (K)	Method	Reference
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)			
C ₆ H ₇ NO	[18617-86-6] $\Delta_{\text{sub}}H$	2-methyl-4-hydroxypyridine	113.0 ± 1.3	298		[1982SUR/SAI, 1986PED/NAY]
C ₆ H ₇ NO	[1121-78-4] $\Delta_{\text{sub}}H$	2-methyl-5-hydroxypyridine	96.2 ± 2.1	298	C	[1982SUR/SAI, 1986PED/NAY]
C ₆ H ₇ NO	[3279-76-3] $\Delta_{\text{sub}}H$	2-methyl-6-hydroxypyridine	92.0 ± 1.3	298	C	[1982SUR/SAI, 1986PED/NAY]
C ₆ H ₇ NO	[931-19-1] $\Delta_{\text{sub}}H$	2-methylpyridine N-oxide	92.9 ± 1.9	298	C	[2010CAB/MON]
	$\Delta_{\text{sub}}H$		78.2 ± 2.2	298	C	[1995ACR/TUC]
C ₆ H ₇ NO	[1003-73-2] $\Delta_{\text{sub}}H$	3-methylpyridine N-oxide	82.2 ± 2.4	298	C	[1995ACR/TUC]
C ₆ H ₇ NO	[1003-67-4] $\Delta_{\text{sub}}H$	4-methylpyridine N-oxide	(345–392) 85.3 ± 2.6	298	ME	[1998RIB/MAT]
	$\Delta_{\text{sub}}H$		(316–341) 79.1 ± 1.3			[1995LEB/CHI]
C ₆ H ₇ NO	[1628-89-3] Δ_vH	2-methoxypyridine	(304–338) 40.5	319	A	[1987STE/MAL]
C ₆ H ₇ NO	[694-85-9] Δ_vH	1-methyl-2(1H)-pyridone	(353–399) 60.2	368	A	[1987STE/MAL]
C ₆ H ₇ NO	[586-95-8] $\Delta_{\text{fus}}H$	4-pyridinemethanol	11.78	325.2		[2005WAN/TAN]
C ₆ H ₇ NO	[1072-83-9] $\Delta_{\text{fus}}H$	2-acetylpyrrole	14.08	363		[2009FLO/CAM]
	$\Delta_{\text{sub}}H$		(277–293) 81.3 ± 1.0	285	ME	[2009SAN/GOM]
	$\Delta_{\text{sub}}H$		(277–293) 81.2 ± 1.0	298	ME	[2009SAN/GOM]
C ₆ H ₇ NO	[1072-82-8] $\Delta_{\text{sub}}H$	3-acetylpyrrole	(316–338) 93.2 ± 0.5	327	ME	[2009SAN/GOM]
	$\Delta_{\text{sub}}H$		(316–338) 94.7 ± 0.5	298	ME	[2009SAN/GOM]
C ₆ H ₇ NO ₂	[na] $\Delta_{\text{fus}}H$	<i>n</i> -ethyl- α -cyanoacrylate	12.86	243.2		[1991BYK/KIP]
C ₆ H ₇ NO ₂	[6973-60-0] $\Delta_{\text{sub}}H$	1-methyl-2-pyrrolecarboxylic acid	(305–327) 95.3 ± 0.7	316	ME	[2009SAN/RIB]
	$\Delta_{\text{sub}}H$		(305–327) 96.2 ± 0.7	298	ME	[2009SAN/RIB]
C ₆ H ₇ NO ₂ S	[98-10-2] $\Delta_{\text{fus}}H$	benzenesulfonamide	25.17	425.8		[2005MAT/MIR]
	$\Delta_{\text{sub}}H$		115.3 ± 1.7	298	C	[2005MAT/MIR]
C ₆ H ₇ NO ₃ S	[121-57-3] $\Delta_{\text{sub}}H$	sulfanilic acid (4-aminobenzene sulfonic acid)	66.9			[1938WOL/WEG, 1960JON]
C ₆ H ₇ NS	[22581-72-2] $\Delta_{\text{sub}}H$	4-(methylthio)pyridine	(347–383) 75.3 ± 3.8	365	B	[1974BEA/MUE]
	Δ_vH		(346–383) 55.8	361	A	[1987STE/MAL]
C ₆ H ₇ NS	[6887-59-8] $\Delta_{\text{sub}}H$	1-methyl-4-thiopyridone	(440–465) 188.3 ± 9.2	452	B	[1974BEA/MUE]
C ₆ H ₇ N ₃ O ₂	[3034-19-3]	2-hydrazino-1-nitrobenzene				

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
		$\Delta_{\text{sub}}H$	124.7		DSC	[1990HWA/YOS]
		Δ_vH	99.6		DSC	[1990HWA/YOS]
C ₆ H ₇ N ₅	[5142-22-3]	1-methyladenine				
		$\Delta_{\text{sub}}H$	138.2		ME	[2000ZIE]
C ₆ H ₇ N ₅	[1445-08-5]	2-methyladenine				
		$\Delta_{\text{sub}}H$	121.7		ME	[2000ZIE]
C ₆ H ₇ N ₅	[5142-23-4]	3-methyladenine				
		$\Delta_{\text{sub}}H$	117.5		ME	[2000ZIE]
		$\Delta_{\text{sub}}H$	83.7 ± 9		HSA	[1978NOW/SZC]
C ₆ H ₇ N ₅	[443-72-1]	N-methyladenine				
		$\Delta_{\text{sub}}H$	(395–425) 123.4 ± 21		ME	[1984ZIE/ZIE]
C ₆ H ₇ N ₅	[22387-37-7]	8-methyladenine				
		$\Delta_{\text{sub}}H$	103.2		ME	[2000ZIE]
C ₆ H ₇ N ₅	[700-00-5]	9-methyladenine				
		$\Delta_{\text{sub}}H$	(381–411) 121.3 ± 4.6		ME	[1984ZIE/ZIE]
		$\Delta_{\text{sub}}H$	(413–458) 121.7	428	HSA	[1987STE/MAL, 1965CLA/PES]
		$\Delta_{\text{sub}}H$	92 ± 8		HSA	[1978NOW/SZC]
C ₆ H ₈	[na]	<i>cis, anti, cis</i> -tricyclo[3.1.0.0 ^{2,4}]hexane				
		Δ_vH	(273–329) 30.6	293		[1979LET/ORC]
		Δ_vH	(273–329) 29.7	313		[1979LET/ORC]
C ₆ H ₈	[592-57-4]	1,3-cyclohexadiene				
		$\Delta_{\text{fus}}H$	4.2	161		[1996DOM/HEA]
		Δ_vH	(307–364) 32.6	322	A,EB	[1987STE/MAL, 1973MEY/HOT]
		Δ_vH	(304–322) 32.4	308	MM	[1974LET/MAR]
C ₆ H ₈	[628-41-1]	1,4-cyclohexadiene				
		$\Delta_{\text{trs}}H$	0.82	192		
		$\Delta_{\text{fus}}H$	5.72	224		[1996DOM/HEA]
		Δ_vH	(304–360) 34.0	319	A	[1987STE/MAL]
		Δ_vH	(304–322) 33.9	308	MM	[1974LET/MAR]
C ₆ H ₈	[2612-46-6]	<i>cis</i> 1,3,5-hexatriene				
		Δ_vH	(306–323) 33.3	314	A,MM	[1987STE/MAL, 1974LET/MAR]
C ₆ H ₈ CIN	[142-04-1]	aniline hydrochloride				
		$\Delta_{\text{sub}}H$	(383–471) 87.5	398	A	[1987STE/MAL, 1975KON/SEL]
C ₆ H ₈ CIN	[14401-92-4]	3-methylpyridine hydrochloride				
		Δ_vH	(420–471) 68.7	435	A	[1987STE/MAL]
C ₆ H ₈ CIN	[14401-93-5]	4-methylpyridine hydrochloride				
		Δ_vH	(437–473) 64.7	452	A	[1987STE/MAL]
C ₆ H ₈ CIO	[na]	chloroethyl methacrylate				
		$\Delta_{\text{fus}}H$	17.0	235.1		[1996DOM/HEA]
C ₆ H ₈ Cl ₂ O ₄	[6941-69-1]	ethylene glycol, <i>bis</i> chloroacetate				
		Δ_vH	(385–557) 73.9	400	A	[1987STE/MAL, 1947STU]
C ₆ H ₈ F ₈ N ₄	[na]	1,1,4,4-tetrakis(difluoroamino)cyclohexane				
		$\Delta_{\text{fus}}H$	46.02	382.2		[2001OXL/SMI]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound					
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference	
C ₆ H ₈ N ₂	[5910-89-4]	2,3-dimethylpyrazine					
	$\Delta_v H$		52.6 ± 1.7	298	C	[2003MOR/MIR]	
C ₆ H ₈ N ₂	[123-32-0]	2,5-dimethylpyrazine					
	$\Delta_v H$	(342–373)	47.2 ± 2.2	298	CGC	[2009LIP/CHI2]	
	$\Delta_v H$	(303–411)	44.5	357		[1995SAK/UEO]	
C ₆ H ₈ N ₂	[111-69-3]	adiponitrile					
	$\Delta_{\text{fus}} H$		18.0	275	DSC	[2007BAD/BLA]	
C ₆ H ₈ N ₂	[4597-87-9]	2-methylaminopyridine					
		$\Delta_v H$	(308–323)	49.0	316	A	[1987STE/MAL]
C ₆ H ₈ N ₂	[18364-47-1]	3-methylaminopyridine					
		$\Delta_v H$	(313–343)	57.2	326	A	[1987STE/MAL]
C ₆ H ₈ N ₂	[1121-58-0]	4-methylaminopyridine					
		$\Delta_v H$	(313–343)	54.1	328	A	[1987STE/MAL]
C ₆ H ₈ N ₂	[95-54-5]	1,2-diaminobenzene					
		$\Delta_{\text{fus}} H$		23.1	373.9		[1997LEE/CHA, 1989BRE/LIC]
		$\Delta_{\text{sub}} H$		85.5 ± 0.3	298	C	[1997SAB/PER]
C ₆ H ₈ N ₂	[108-45-2]	1,3-diaminobenzene					
		$\Delta_{\text{fus}} H$		15.4	335.5		[1996DOM/HEA]
		$\Delta_{\text{sub}} H$		90.4 ± 0.4	298	C	[1997SAB/PER]
		$\Delta_v H$	(372–559)	63.7	387	A	[1987STE/MAL, 1947STU]
C ₆ H ₈ N ₂	[106-50-3]	1,4-diaminobenzene					
		$\Delta_{\text{fus}} H$		21.7	412.3		[1996DOM/HEA]
		$\Delta_{\text{sub}} H$		92.2 ± 0.2	298	C	[1997SAB/PER]
C ₆ H ₈ N ₂	[100-63-0]	phenyl hydrazine					
		$\Delta_{\text{fus}} H$		16.43	292.8		[1996DOM/HEA]
		$\Delta_v H$	(413–518)	57.3	428	A	[1987STE/MAL, 1972DYK]
		$\Delta_v H$	(345–517)	59.2	360		[1947STU]
C ₆ H ₈ N ₂	[13925-00-3]	ethylpyrazine					
		$\Delta_v H$		48.8 ± 1.9	298	C	[2003MOR/MIR]
C ₆ H ₈ N ₂ O ₂	[874-14-6]	1,3-dimethyluracil					
		$\Delta_{\text{fus}} H$		14.6	398		[1996DOM/HEA]
		$\Delta_{\text{sub}} H$	(311–367)	115.8 ± 3.0	338	TE	[2000BRU/PIA]
		$\Delta_{\text{sub}} H$		96.4 ± 1.4	298	C	[1985MUR/SAK]
		$\Delta_{\text{sub}} H$	(313–363)	101.7 ± 2.1	338	QR	[1980TEP/YAN]
		$\Delta_{\text{sub}} H$	(400–454)	46 ± 4.2	426	HAS	[1978NOW/SZC]
C ₆ H ₈ N ₂ O ₂	[4160-72-9]	1-methylthymine					
		$\Delta_{\text{sub}} H$	(378–428)	124.4 ± 1.3	398	QR	[1980TEP/YAN]
C ₆ H ₈ N ₂ O ₂	[na]	N-acetylglycine amide					
		$\Delta_{\text{fus}} H$		25.6	408.2		[1988FER/DEL]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₆ H ₈ N ₂ O ₂	[4538-37-8] $\Delta_{\text{fus}}H$	1,4-diisocyanatobutane (8-360)	20.76	231.2	AC	[2005SMI/KAN2]
C ₆ H ₈ N ₂ O ₂ S	[na] $\Delta_{\text{trs}}H$ $\Delta_{\text{fus}}H$ $\Delta_{\text{fus}}H$	<i>p</i> -aminobenzene sulphonamide	1.63 24.02 23.0	407 439.3 438.7		[1996CIO/MEL] [1996CIO/MEL] [1985OHM/LIP]
C ₆ H ₈ N ₂ O ₂ S	[63-74-1] $\Delta_{\text{fus}}H$	4-aminobenzenesulfonamide (sulfanilamide)	23.3	435.4		[2002MAR/GOM]
C ₆ H ₈ N ₂ O ₈	[na] $\Delta_{\text{fus}}H$	1,4:3,6-dianhydromannitol dinitrate (isomannide dinitrate)	20.5	337.2		[1998HAT/SUZ]
C ₆ H ₈ N ₂ O ₈	[na] $\Delta_{\text{fus}}H$	1,4:3,6-dianhydro-(<i>d</i>)-glucitol dinitrate (isosorbide dinitrate)	27.63	341.7		[1998HAT/SUZ]
C ₆ H ₈ N ₂ O ₈	[na] $\Delta_{\text{fus}}H$	1,4:3,6-dianhydroiditol dinitrate (isoidide dinitrate)	12.81	325.9		[1998HAT/SUZ]
C ₆ H ₈ N ₄ O ₂	[4164-33-4] $\Delta_{\text{fus}}H$	<i>bis</i> (2-cyanoethyl)- <i>N</i> -nitroamine	44.99	327		[1987OYU/BR1]
C ₆ H ₈ N ₄ O ₈	[146028-82-2] $\Delta_{\text{fus}}H$	1,1,4,4-tetranitrocyclohexane	108.8	489.2		[2001OXL/SMI]
		Note: Experimental enthalpy is abnormally large—compound may be decomposing				
C ₆ H ₈ N ₆ O ₈	[na] $\Delta_{\text{fus}}H$	1,3-dinitro-3(1',3')-dinitroazetidid-3'-yl)azetidide	25.52	387.5		[1998MCK/FLO]
C ₆ H ₈ O	[930-68-7] Δ_vH Δ_vH	2-cyclohexen-1-one (351-445) (335-481)	45.0 49.5 ± 0.4	366 298	EB EB	[2006PAL/ORA] [1997STE/CHI3]
C ₆ H ₈ O	[625-36-3] Δ_vH Δ_vH	2,5-dimethylfuran (271-308) (271-308)	32.3 ± 0.3 31.8 ± 0.3	290 298	GS GS	[1998VER/WEL] [1998VER/WEL]
C ₆ H ₈ O ₂	[4935-01-7] Δ_vH	methyl bicyclo[1.1.0]butane-1-carboxylate (299-377)	37.3	318	BG	[1971HAL/BAL]
C ₆ H ₈ O ₂	[504-02-9] $\Delta_{\text{sub}}H$	1,3-cyclohexanedione	89.8 ± 1.1	298	C	[1993PIL/PAR]
C ₆ H ₈ O ₂	[637-88-7] $\Delta_{\text{trs}}H$ $\Delta_{\text{trs}}H$ $\Delta_{\text{fus}}H$ $\Delta_{\text{trs}}H$ $\Delta_{\text{trs}}H$ $\Delta_{\text{fus}}H$ $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$	1,4-cyclohexanedione	6.20 1.18 11.26 6.15 0.96 10.04 75.0 ± 1.0 84.4 84.2	319.9 338.8 351.5 322.2 339.2 348.2 298 289 298		[1983DEW/DEK] [1972ALV/BOR] [1993PIL/PAR] [1983DEW/VAN] [1983DEW/VAN]
C ₆ H ₈ O ₃	[17347-61-4] Δ_vH	2,2-dimethylsuccinic acid anhydride (334-493)	57.3	349	A	[1987STE/MAL, 1947STU]
C ₆ H ₈ O ₃	[31468-33-4] Δ_vH	2-methylglutaric acid anhydride (366-556)	60.7	381	A	[1987STE/MAL, 1947STU]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₆ H ₈ O ₄	[624-49-7]	dimethyl fumarate				
	$\Delta_{\text{fus}}H$		35.15	375		[1996DOM/HEA]
	$\Delta_{\text{sub}}H$		NA			[1972LEB/KAT]
	$\Delta_{\text{sub}}H$		84.5 ± 1.7			[1934WOL/TRI]
C ₆ H ₈ O ₄	[624-48-6]	dimethyl maleate				
	$\Delta_{\text{fus}}H$		14.64	254		[1996DOM/HEA]
	$\Delta_{\text{sub}}H$		44.8			[1938WOL/WEG, 1960JON, 1935TRI]
	$\Delta_{\text{sub}}H$	(317–341)	41.8 ± 4.2			[1934WOL/TRI]
C ₆ H ₈ O ₄	[5445-51-2]	cyclobutane-1,1-dicarboxylic acid				
	$\Delta_{\text{sub}}H$		112.2 ± 0.7		C	[1983ALT/PIL]
	[3396-14-3]	cyclobutane-1,2-dicarboxylic acid				
	$\Delta_{\text{sub}}H$		120.0 ± 0.9		C	[1983ALT/PIL]
C ₆ H ₈ O ₄	[95-96-5]	(<i>dl</i>) 3,6-dimethyl-1,4-dioxane-2,5-dione				
	$\Delta_{\text{fus}}H$		24.7	397.5		[1996DOM/HEA]
C ₆ H ₈ O ₄	[4511-42-6]	(<i>l</i>) 3,6-dimethyl-1,4-dioxane-2,5-dione				
	$\Delta_{\text{fus}}H$		16.94	366.6		[1999LEB/KUL]
C ₆ H ₈ O ₅	[3184-35-8]	2-oxohexanedioic acid				
	$\Delta_{\text{sub}}H$	(281–301)	127.0		TPTD	[2005CHA/ZIE]
Note: Values based on TPTD method are not consistent with values determined by other experimental methods						
C ₆ H ₈ O ₅	[689-31-6]	3-oxohexanedioic acid				
	$\Delta_{\text{sub}}H$	(307–329)	151		TPTD	[2005CHA/ZIE]
Note: Values based on TPTD method are not consistent with values determined by other experimental methods						
C ₆ H ₈ O ₆	[na]	(<i>l</i>)-ascorbic acid				
	$\Delta_{\text{fus}}H$		37.04	466.2		[1998MUR/BET]
C ₆ H ₈ S	[632-16-6]	2,3-dimethylthiophene				
	Δ_vH	(353–473)	39.4	368	A	[1987STE/MAL, 1972DYK]
C ₆ H ₈ S	[638-00-6]	2,4-dimethylthiophene				
	Δ_vH	(323–493)	41.4	338	A	[1987STE/MAL, 1972DYK, 1999DYK/SVO]
C ₆ H ₈ S	[638-02-8]	2,5-dimethylthiophene				
	$\Delta_{\text{fus}}H$		8.91	210.6		[1996DOM/HEA]
	Δ_vH		40.2 ± 0.9	298	C	[2008RIB/SAN3]
	Δ_vH	(333–374)	39.7	348	IA	[1987STE/MAL, 1971EON/POM, 1999DYK/SVO]
C ₆ H ₈ S	[632-15-5]	3,4-dimethylthiophene				
	Δ_vH	(328–478)	41.1	343	A	[1987STE/MAL, 1972DYK, 1999DYK/SVO]
C ₆ H ₈ S	[872-55-9]	2-ethylthiophene				
	Δ_vH		39.7 ± 0.9	298	C	[2007RIB/SAN]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(333–374)	39.7	348	IA	[1987STE/MAL, 1971EON/POM, 1999DYK/SVO]
C ₆ H ₈ S	[1795-01-3] $\Delta_v H$	3-ethylthiophene (318–473)	40.7	333	A	[1987STE/MAL, 1972DYK]
C ₆ H ₉ F ₃ O ₂	[367-64-6] $\Delta_v H$	butyl trifluoroacetate (343–377)	37.8	358	A,EB	[1987STE/MAL, 1969SHE/LAN]
C ₆ H ₉ N	[4254-02-8] $\Delta_v H$	cyclopentanecarbonitrile (340–418)	48.1 ± 0.1	298	C	[1983FUC/HAL]
	$\Delta_v H$		43.4 ± 0.1	298		[1973KON]
	$\Delta_v H$		40.9	359	BG	[1971HAL/BAL]
	$\Delta_v H$		43.5 ± 0.1	298	C	[1970PRO/KRE]
C ₆ H ₉ N	[625-82-1] $\Delta_{\text{fus}} H$	2,4-dimethylpyrrole	9.6	268.5		[1994CHI/HOS2]
C ₆ H ₉ N	[625-84-3] $\Delta_{\text{fus}} H$	2,5-dimethylpyrrole	9.3	280.9		[1996DOM/HEA]
	$\Delta_v H$	(373-43)	49.5	388	A,IP,EB	[1987STE/MAL, 1968OSB/DOU]
C ₆ H ₉ NO	[88-12-0] $\Delta_{\text{fus}} H$	N-vinylpyrrolidone	15.28	286.2		[1997KUL/LEB2]
C ₆ H ₉ NO ₂	[1572-99-2] $\Delta_v H$	ethyl 2-cyanopropionate (283–323)	58.6 ± 0.3	298	GS	[1995VER/BEC]
C ₆ H ₉ NO ₆	[na] $\Delta_{\text{fus}} H$	isomannide mononitrate	20.64	344.8		[1998HAT/SUZ]
C ₆ H ₉ NO ₆	[na] $\Delta_{\text{fus}} H$	isosorbide-2-mononitrate	26.38	328		[1998HAT/SUZ]
C ₆ H ₉ NO ₆	[na] $\Delta_{\text{fus}} H$	isosorbide-5-mononitrate	22.36	364		[1998HAT/SUZ]
C ₆ H ₉ NS	[13623-11-5] $\Delta_{\text{fus}} H$	2,4,5-trimethylthiazole	9.0	240.7		[1966MEY/MET]
C ₆ H ₉ N ₃ O	[7171-70-2] $\Delta_{\text{sub}} H$	1,3,5-trimethyl-4-nitrosopyrazole	88.0 ± 2.0	298	C	[2001RIB/FER]
C ₆ H ₉ N ₃ O	[17634-60-5] $\Delta_{\text{sub}} H$	1,5-dimethylcytosine (390–437)	132.8 ± 0.6		GS	[1998ZIE/WSZ]
C ₆ H ₉ N ₃ O	[6220-49-1] $\Delta_{\text{sub}} H$	1,N-dimethylcytosine (401–426)	122.2 ± 0.3		GS	[1998ZIE/WSZ]
C ₆ H ₉ N ₃ O ₂	[71-00-1] $\Delta_{\text{sub}} H$	L-histidine (392–492)	142 ± 8	442	LE	[1977GAF/PIE]
C ₆ H ₉ N ₃ O ₂	[20555-80-0] $\Delta_{\text{sub}} H$	1-methyl-N4-methoxycytosine (316–325) (320–357)	107.6 ± 0.3		ME	[1999ZIE/PER]
	$\Delta_{\text{sub}} H$		106.9 ± 0.4		GS	[1999ZIE/PER]
	$\Delta_{\text{sub}} H$		106.4 ± 0.8			[1998ZIE/WSZ]
C ₆ H ₉ N ₃ O ₂	[6220-53-7] $\Delta_{\text{sub}} H$	1,5-dimethyl-N-hydroxycytosine (357–394)	115.2 ± 0.6		GS	[1998ZIE/WSZ]
C ₆ H ₉ N ₃ O ₂	[36315-01-2]	2-amino-4,6-dimethoxypyrimidine				

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{fus}}H$		29.85	371		[2003SUN/SON]
C ₆ H ₉ N ₃ O ₃	[877-89-4]	2,4,6-trimethoxy-1,3,5-triazine				
	$\Delta_{\text{fus}}H$ (β -form)		18.1	404		[2004FRI/KAP]
	$\Delta_{\text{fus}}H$ (γ -form)		11.4	409		[2004FRI/KAP]
	$\Delta_{\text{trs}}H$	(298–523)	3.9	340.2	DSC	
	$\Delta_{\text{fus}}H$	(298–523)	18.1	395.2	DSC	[2000HAN/BOT]
C ₆ H ₉ N ₃ O ₃	[na]	6-methoxy-3,5-dimethyl-tetrahydrotriazine-2,4-dione				
	$\Delta_{\text{fus}}H$		12.7	363.2		[2000HAN/BOT, 2004FRI/KAP]
		Note: Compound rearranges shortly after melting.				
C ₆ H ₉ N ₃ O ₃	[877-89-4]	trimethyl isocyanurate				
	$\Delta_{\text{sub}}H$	(330–346)	86.6 ± 1.3	338	C	[1988IMA/MUR]
	$\Delta_{\text{sub}}H$		88.2 ± 1.3	298	C	[1988IMA/MUR]
	$\Delta_{\text{sub}}H$		88.2 ± 1.3	298	C	[1989IMA/TAK, 1985MUR/SAK]
C ₆ H ₉ N ₃ O ₃	[877-89-4]	trimethyl cyanurate				
	$\Delta_{\text{sub}}H$		90.3 ± 1.0	298	C	[1989IMA/TAK, 1985MUR/SAK]
C ₆ H ₉ P	[3746-01-8]	trivinylphosphine				
	Δ_vH	(289–334)	33.7	304		[1957MAI/SEY, 1984BOU/FRI]
C ₆ H ₁₀	[285-58-5]	<i>cis</i> bicyclo[3.1.0]hexane				
	Δ_vH	(273–300)	33.7	286	A	[1987STE/MAL]
	Δ_vH		33.5 ± 0.4	298		[1970CHA/MCN]
C ₆ H ₁₀	[5685-46-1]	bicyclopropyl				
	Δ_vH		31.7 ± 0.5	298	C	[2007PAS/KUZ]
C ₆ H ₁₀	[3664-56-0]	1,2,2-trimethylcyclopropene				
	Δ_vH		27.9 ± 1.4	298	C	[2007PAS/KUZ]
	Δ_vH		26.8 ± 1.7	298		[1986PIM/DOM, 2007PAS/KUZ]
C ₆ H ₁₀	[110-83-8]	cyclohexene				
	$\Delta_{\text{trs}}H$		4.23	138.7		
	$\Delta_{\text{fus}}H$		3.28	169.7		[1996DOM/HEA]
	Δ_vH	(315–356)	32.8	330		[2009MAR/AUC]
	Δ_vH	(310–356)	32.9	325		[2004STE/SUN]
	Δ_vH	(312–356)	32.6	327		[2001SEG/LAM]
	Δ_vH	(285–357)	33.5 ± 0.5	298	EB	[1996STE/CHI3]
	Δ_vH	(309–365)	32.7	324	A,EB	[1987STE/MAL, 1973MEY/HOT]
	Δ_vH	(305–322)	33.1	308	MM	[1974LET/MAR]
	Δ_vH		32.7 ± 0.1	313	C	[1973SVO/VES]
	Δ_vH		32.2 ± 0.1	323	C	[1973SVO/VES]
	Δ_vH		31.7 ± 0.1	333	C	[1973SVO/VES]
	Δ_vH		31.2 ± 0.1	343	C	[1973SVO/VES]
	Δ_vH		30.7 ± 0.1	353	C	[1973SVO/VES]
	Δ_vH	(285–357)	33.7	300	MM	[1950FOR/CAM]
	Δ_vH	(229–292)	32.6	300		[1941LIS]
C ₆ H ₁₀	[na]	1-methylcyclopentene				
	Δ_vH		32.6 ± 0.2	298	GCC	[1979FUC/PEA]
	Δ_vH	(268–403)	33.4	283	A	[1987STE/MAL, 1972DYK]
C ₆ H ₁₀	[1120-62-3]	3-methylcyclopentene				
	Δ_vH		31.0 ± 0.2	298	GCC	[1979FUC/PEA]
	Δ_vH	(263–392)	32.1	278	A	[1987STE/MAL, 1972DYK]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		T _m (K)	Method	Reference
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)			
C ₆ H ₁₀	[1759-81-5] $\Delta_{\text{v}}H$	4-methylcyclopentene (271–403)	33.2	286	A	[1987STE/MAL, 1972DYK]
C ₆ H ₁₀	[1489-61-8] $\Delta_{\text{v}}H$	1,3-dimethylcyclobutene (269–296)	31.3	282	A	[1987STE/MAL]
C ₆ H ₁₀	[513-81-5] $\Delta_{\text{v}}H$	2,3-dimethyl-1,3-butadiene (273–342)	32.2	288	A	[1987STE/MAL, 1955CUM/MCL]
C ₆ H ₁₀	[592-48-3] $\Delta_{\text{v}}H$	<i>trans</i> 1,3-hexadiene (299–319)	32.1	309	A,MM	[1987STE/MAL, 1974LET/MAR]
C ₆ H ₁₀	[7319-00-8] $\Delta_{\text{v}}H$	<i>trans</i> 1,4-hexadiene (304–323)	30.2	313	A,MM	[1987STE/MAL, 1974LET/MAR]
C ₆ H ₁₀	[592-42-7] $\Delta_{\text{v}}H$ $\Delta_{\text{v}}H$ $\Delta_{\text{v}}H$	1,5-hexadiene (299–333) (300–319) (273–333)	29.4	314	A	[1987STE/MAL]
			28.6	308		[1974LET/MAR]
			30.5	288	A	[1987STE/MAL, 1955CUM/MCL, 1972DYK]
C ₆ H ₁₀	[5194-51-4] $\Delta_{\text{v}}H$ $\Delta_{\text{v}}H$	<i>trans trans</i> 2,4-hexadiene (304–354) (305–323)	33.2	319	A	[1987STE/MAL]
			33.2	308	MM	[1974LET/MAR]
C ₆ H ₁₀	[693-02-7] $\Delta_{\text{v}}H$ $\Delta_{\text{v}}H$ $\Delta_{\text{v}}H$	1-hexyne (250–290) (237–287) (265–391)	33.5	270	MM	[1981CHI/HYM]
			34.2	262	HSA	[1981CHI/HYM]
			33.4	280	A	[1987STE/MAL, 1972DYK]
C ₆ H ₁₀	[264-35-2] $\Delta_{\text{v}}H$ $\Delta_{\text{v}}H$	2-hexyne (283–313) (283–313)	35.8	298		[2007BOU/BEL]
			35.9	298		[2006BOU/BEL]
C ₆ H ₁₀	[764-35-2] $\Delta_{\text{v}}H$ $\Delta_{\text{v}}H$	3-hexyne (253–354) (253–298)	30.5	268	A	[1987STE/MAL]
			31.6	275	T	[1965RON/HAR]
C ₆ H ₁₀ Br ₂	[7429-37-0] $\Delta_{\text{v}}H$	<i>trans</i> 1,2-dibromocyclohexane (350–416)	53.3	365	A	[1987STE/MAL]
C ₆ H ₁₀ ClFO ₂	[na] $\Delta_{\text{v}}H$	3-fluorobutyric acid, 2-chloroethyl ester (273–333)	60.4	288	A	[1987STE/MAL, 1972DYK]
C ₆ H ₁₀ ClF ₃ O	[358-36-1] $\Delta_{\text{v}}H$ $\Delta_{\text{v}}H$ $\Delta_{\text{v}}H$ $\Delta_{\text{v}}H$	2-chloro-1,1,2-trifluoroethyl butyl ether	45.1 ± 0.1	298	C	[1984MAJ/UCH]
			43.9 ± 0.1	313	C	[1984MAJ/UCH]
			42.8 ± 0.1	328	C	[1984MAJ/UCH]
			41.6 ± 0.1	343	C	[1984MAJ/UCH]
C ₆ H ₁₀ Cl ₂	[2108-92-1] $\Delta_{\text{trs}}H$ $\Delta_{\text{fus}}H$ $\Delta_{\text{v}}H$	1,1-dichlorocyclohexane (335–444)	9.16	225		
			1.47	236		[1999KAB/KOZ]
			43.5	350	A	[1987STE/MAL]
C ₆ H ₁₀ Cl ₂	[10498-35-8] $\Delta_{\text{v}}H$	<i>cis</i> 1,2-dichlorocyclohexane (364–480)	45.8	379	A	[1987STE/MAL]
C ₆ H ₁₀ Cl ₂	[822-86-6] $\Delta_{\text{v}}H$	<i>trans</i> 1,2-dichlorocyclohexane (344–462)	45.8	359	A	[1987STE/MAL]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₆ H ₁₀ Cl ₂	[19398-57-3] $\Delta_v H$	1,4-dichlorocyclohexane (353–406)	47.8	368	A	[1987STE/MAL]
C ₆ H ₁₀ Cl ₂ O ₂	[37079-08-6] $\Delta_v H$	isobutyl dichloroacetate (301–456)	51.4	316	A	[1987STE/MAL, 1947STU]
C ₆ H ₁₀ F ₂ O ₂	[na] $\Delta_v H$	3-fluorobutyric acid, 2-fluoroethyl ester (273–333)	54.8	288	A	[1987STE/MAL, 1972DYK]
C ₆ H ₁₀ N ₂ O	[na] $\Delta_{\text{trs}}H$ $\Delta_{\text{trs}}H$ $\Delta_{\text{fus}}H$	2,3-diazabicyclo[2.2.2]oct-2-ene N-oxide	5.02 8.05 3.84	359.3 399.3 438		[1980BYS]
C ₆ H ₁₀ N ₂ O ₂	[7491-74-1] $\Delta_{\text{fus}}H$ (I) $\Delta_{\text{fus}}H$ (II)	2-oxo-1-pyrrolidineacetamide (piracetam)	25.59 29.85	426 412	DSC	[1996CEO/AGA]
C ₆ H ₁₀ N ₆ O ₉	[28464-26-8] $\Delta_{\text{sub}}H$	N-(2,2-dinitropropyl)-2,2-dinitro-N-nitroso-1-propanamine (323–336)	110.9 ± 8		ME	[1973PEP/GAF, 1977PED/RYL]
C ₆ H ₁₀ N ₆ O ₁₀	[28464-24-6] $\Delta_{\text{sub}}H$	N-(2,2-dinitropropyl)-2,2-dinitro- N-nitro-1-propanamine (398–423)	99.2 ± 0.8		ME	[1973PEP/GAF]
C ₆ H ₁₀ O	[279-49-2] $\Delta_{\text{trs}}H$ $\Delta_{\text{trs}}H$ $\Delta_{\text{fus}}H$	7-oxabicyclo[2.2.1]heptane	4.86 0.98 0.71	180.5 218.5 244	DSC	[1998PAR/GIL]
C ₆ H ₁₀ O	[1462-03-9] $\Delta_{\text{sub}}H$ (cryst III) $\Delta_{\text{sub}}H$ (cryst III) $\Delta_{\text{sub}}H$ (cryst III)	1-methylcyclopentanol (253–281)	73.7 ± 0.4 67.0 ± 0.2 67.4 ± 0.2	267 298 291	ME C	[1997BLO/KAB] [1997BLO/KAB] [1997BLO/KAB]
C ₆ H ₁₀ O	[na] $\Delta_v H$ $\Delta_v H$	cyclopentenyl methyl ether (274–313) (274–313)	42.3 ± 0.8 42.1 ± 0.8	294 298	GS GS	[1998VER/WEL] [1998VER/WEL]
C ₆ H ₁₀ O	[12655-16-2] $\Delta_v H$	2,3-dihydro-4-methyl-2H-pyran (304–392)	38.1	319	A	[1987STE/MAL, 1968KAC/NEM, 1984BOU/FRI]
C ₆ H ₁₀ O	[35656-02-1] $\Delta_v H$	methylenetetrahydro-2H-pyran (339–382)	36.8	354	A	[1987STE/MAL]
C ₆ H ₁₀ O	[108-94-1] $\Delta_{\text{trs}}H$ $\Delta_{\text{fus}}H$ $\Delta_{\text{sub}}H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$	cyclohexanone (243–265) (343–427) (343–383) (318–428)	8.66 1.33 49.3 43.1 46.6 ± 0.4 44.0 44.4 ± 0.1 44.0 ± 0.1 43.4 ± 0.1 43.1 ± 0.1 42.2 ± 0.1 41.8 ± 0.1 41.4 ± 0.1	220.8 245.2 254 358 298 333 308 313 323 328 338 343 348	EB CGC C C C C C C C C	[1980NAK/SUG] [1948NIT/SEK2] [2006TEO/BAR] [1995CHI/HOS] [1993AUC/MON] [1992SVO/KUB] [1992SVO/KUB] [1992SVO/KUB] [1992SVO/KUB] [1992SVO/KUB] [1992SVO/KUB] [1992SVO/KUB] [1992SVO/KUB]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		T _m (K)	Method	Reference
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)			
		$\Delta_v H$		42.3 ± 0.2		GC [1989AZA]
		$\Delta_v H$	(345–458)	42.2	360	EB [1987AMB/GHI2]
		$\Delta_v H$	(395–426)	40.4	410	[1984CAS/FRA2]
		$\Delta_v H$	(362–439)	41.5	377	A,EB [1987STE/MAL, 1973MEY/HOT]
		$\Delta_v H$		44.9 ± 0.6	298	[1972WOL]
		$\Delta_v H$		45.1 ± 0.1	298	C [1968PLA/WIL]
		$\Delta_v H$	(273–298)	40.3	286	[1938RAD/ALE]
C₆H₁₀O	[109-49-9]		5-hexen-2-one			
		$\Delta_v H$	(317–440)	42.1 ± 0.1	320	EB [2002STE/CHI5]
		$\Delta_v H$	(317–440)	39.4 ± 0.2	360	EB [2002STE/CHI5]
		$\Delta_v H$	(317–440)	36.6 ± 0.3	400	EB [2002STE/CHI5]
		$\Delta_v H$	(317–440)	33.5 ± 0.6	440	EB [2002STE/CHI5]
		$\Delta_v H$	(449–561)	34.6	464	A [1987STE/MAL]
C₆H₁₀O	[141-79-7]		mesityl oxide			
		$\Delta_v H$		35.2	401	[1998LOU, 1997STE/CHI]
			Note: May be a mixture of 2-methyl-1-penten-4-one and 4-methyl-3-penten-2-one			
C₆H₁₀O	[3744-02-3]		2-methyl-1-penten-4-one			
		$\Delta_v H$	(389–461)	36.9	404	A [1987STE/MAL]
		$\Delta_v H$	(286–461)	41.9	298	[1975AMB/ELL]
		$\Delta_v H$	(306–398)	41.1	321	MM [1987STE/MAL, 1947STR/MON, 1972DYK]
C₆H₁₀O	[141-79-7]		4-methyl-3-penten-2-one			
		$\Delta_v H$	(303–442)	42.7 ± 0.3	298	EB [1997STE/CHI]
		$\Delta_v H$	(303–442)	41.4 ± 0.3	320	EB [1997STE/CHI]
		$\Delta_v H$	(303–442)	39.1 ± 0.3	360	EB [1997STE/CHI]
		$\Delta_v H$	(303–442)	36.5 ± 0.3	400	EB [1997STE/CHI]
		$\Delta_v H$	(303–442)	33.5 ± 0.6	440	EB [1997STE/CHI]
		$\Delta_v H$	(343–383)	44.8	298	CGC [1995CHI/HOS]
		$\Delta_v H$	(399–471)	37.8	414	A [1987STE/MAL]
		$\Delta_v H$	(292–471)	43.3	298	[1975MES/FIN]
		$\Delta_v H$	(313–405)	41.5	328	MM [1987STE/MAL, 1947STR/MON, 1972DYK]
C₆H₁₀O	[286-20-4]		cyclohexene oxide (1,2-epoxycyclohexane)			
		$\Delta_{\text{trs}}H$	(10–300)	9.54	193.1	
		$\Delta_{\text{fus}}H$	(10–300)	1.06	238.1	[1980NAK/SUG]
C₆H₁₀O₂	[765-85-5]		methyl cyclobutanecarboxylate			
		$\Delta_v H$		44.2 ± 0.2		GS [1998VER/KUM]
		$\Delta_v H$		44.7 ± 0.1	298	C [1983FUC/HAL]
		$\Delta_v H$	(319–378)	41.4	340	BG [1971HAL/BAL]
C₆H₁₀O₂	[na]		cyclopropanecarboxylic acid ethyl ester			
		$\Delta_v H$	(278–308)	44.0 ± 0.5		GS [1998VER/KUM]
C₆H₁₀O₂	[106-92-3]		allyl glycidyl ether			
		$\Delta_v H$	(323–420)	47.0	338	A [1987STE/MAL]
C₆H₁₀O₂	[123-20-6]		butyric acid, vinyl ester			
		$\Delta_v H$	(365–387)	39.3	376	A [1987STE/MAL]
C₆H₁₀O₂	[1072-96-4]		4-vinyl-1,3-dioxane			
		$\Delta_v H$	(306–416)	54.5	321	A [1987STE/MAL]
C₆H₁₀O₂	[502-44-3]		ϵ -caprolactone			
		$\Delta_{\text{fus}}H$		13.82	272	[1991ACR]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound						
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference		
		$\Delta_v H$	(283–343)	U 38.2	298		[2008BIA/CEZ]	
		$\Delta_v H$	(395–436)	54.0 ± 0.2	415	EB	[1991WIB/WAL]	
		$\Delta_v H$	(395–436)	62.0 ± 1.3	298	EB	[1991WIB/WAL]	
C ₆ H ₁₀ O ₂	[823-22-3]	δ -hexanolactone						
		$\Delta_v H$	(283–343)	58.1	298		[2008BIA/CEZ]	
		$\Delta_v H$	(283–353)	60.9 ± 0.1	298	GS	[2007EME/KOZ]	
C ₆ H ₁₀ O ₂	[695-06-7]	γ -caprolactone						
		$\Delta_v H$	(283–353)	57.2 ± 0.3	298	GS	[2008EME/KOZ, 2009EME/VER]	
		$\Delta_v H$	(243–298)	55.3 ± 0.6	298		[2004COV/MOK, 2008EME/KOZ]	
C ₆ H ₁₀ O ₂	[924-50-5]	methyl 3-methylbut-2-enoate						
		$\Delta_v H$	(274–304)	46.9 ± 0.2	298	GS	[2008EME/TOK]	
C ₆ H ₁₀ O ₂	[10544-63-5]	ethyl crotonate						
		$\Delta_v H$	(329–420)	47.1	344	A	[1987STE/MAL]	
C ₆ H ₁₀ O ₂	[97-63-2]	ethyl methacrylate						
		$\Delta_v H$	(285–390)	38.3	300	A	[1987STE/MAL]	
C ₆ H ₁₀ O ₂	[3123-97-5]	5,5-dimethyldihydro-2(3H)-furanone						
		$\Delta_v H$	(311–480)	52.7	326	A	[1987STE/MAL]	
C ₆ H ₁₀ O ₂	[925-60-0]	propyl acrylate						
		$\Delta_v H$	(287–395)	37.9	302	A	[1987STE/MAL]	
C ₆ H ₁₀ O ₂	[110-13-4]	2,5-hexanedione						
		$\Delta_v H$	(386–474)	50.1	401	A	[1987STE/MAL]	
C ₆ H ₁₀ O ₃	[na]	<i>cis/trans</i> 2,5-dimethoxy-2,5-dihydrofuran						
		$\Delta_v H$		44.2 ± 0.3	298	CGC	[2000NIC/ORF]	
C ₆ H ₁₀ O ₃	[na]	cyclohexene ozonide						
		$\Delta_v H$	(276–311)	74.2	291	A	[1987STE/MAL]	
		$\Delta_v H$	(353–403)	58.6	378		[1977BOL/MAK]	
C ₆ H ₁₀ O ₃	[141-97-9]	ethyl acetoacetate						
		$\Delta_v H$	(301–454)	52.5	316	A	[1987STE/MAL]	
C ₆ H ₁₀ O ₃	[624-45-3]	methyl levulinate						
		$\Delta_v H$	(312–471)	50.4	327	A	[1987STE/MAL, 1947STU]	
		$\Delta_v H$		51.1	410		[1931SCH/COW]	
C ₆ H ₁₀ O ₃	[123-62-6]	propionic anhydride						
		$\Delta_v H$	(293–440)	48.2	308	A	[1987STE/MAL]	
		$\Delta_v H$	(341–440)	52.2	356		[1883KAH]	
C ₆ H ₁₀ O ₃	[141-97-9]	ethyl 3-oxobutanoate						
		$\Delta_v H$		54.2 ± 1.0	298	C	[1995RIB/FER]	
		$\Delta_v H$		55.0			[1975VIL/PER]	
C ₆ H ₁₀ O ₃	[766-32-5]	4-methyl-2,6,7-trioxabicyclo[2.2.2]octane						
		$\Delta_{\text{fus}} H$		10.4	369.2		[1995RAK/VER2]	
		$\Delta_{\text{sub}} H$		67.4	298		[1995RAK/VER2]	
C ₆ H ₁₀ O ₃	[3592-12-9]	2,2-dimethyltrimethylene carbonate						
		$\Delta_{\text{trs}} H$		10.3	324.1			
		$\Delta_{\text{fus}} H$		5.62	387.2		[1995LEB/KUL2]	
C ₆ H ₁₀ O ₄	[75096-35-4]	<i>cis</i> -1,3,5,7-tetraoxadecalin or [54933-94-7]						

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	Enthalpy							
		$\Delta_{\text{fus}}H$			28.62	450.2		[1998LIN/BEC]
		$\Delta_{\text{sub}}H$			94.9	298		[1998LIN/BEC]
C₆H₁₀O₄	[75096-35-4]		<i>trans</i> -1,3,5,7-tetraoxadecalin or					[54933-94-7]
		$\Delta_{\text{fus}}H$			23.14	374.5		[1998LIN/BEC]
		$\Delta_{\text{sub}}H$			81.5	298		[1998LIN/BEC]
C₆H₁₀O₄	[542-10-9]		1,1-diacetoxyethane					
		Δ_vH	(343–438)		49.7	358	A	[1987STE/MAL]
C₆H₁₀O₄	[6284-75-9]		2-acetoxypropionic acid, methyl ester					
		Δ_vH	(337–445)		52.9	352	A	[1987STE/MAL]
C₆H₁₀O₄	[38003-42-8]		3-acetoxypropionic acid, methyl ester					
		Δ_vH	(343–358)		68.0	350	A	[1987STE/MAL]
C₆H₁₀O₄	[95-92-1]		diethyl oxalate					
		Δ_vH	(343–457)		53.9	358	A	[1987STE/MAL]
		Δ_vH	(320–459)		62.3	335	A	[1987STE/MAL, 1947STU]
C₆H₁₀O₄	[106-65-0]		dimethyl succinate					
		Δ_vH	(286–340)		61.0 ± 0.3	298	GS	[2006VER/KOZ]
		Δ_vH	(342–468)		61.7 ± 0.4	298		[1992KAT, 2006VER/KOZ]
		Δ_vH	(340–470)		49.3	364	A	[1987STE/MAL]
		Δ_vH	(367–460)		60.9 ± 0.4	298	EB	[1987DAU/JAL, 2006VER/KOZ]
		Δ_vH	(398–468)		62.4	298	EB	[1963VLA/GRA, 2006VER/KOZ]
C₆H₁₀O₄	[111-55-7]		ethylene glycol diacetate					
		Δ_vH	(291–334)		61.4 ± 0.2	298	GS	[2009VER/EME2]
		Δ_vH	(311–464)		55.2	326	A	[1987STE/MAL]
		Δ_vH			61.4 ± 0.2	298	C	[1986NIL/WAD]
		Δ_vH			61.0 ± 0.1	298	C	[1970KUS/WAD]
		Δ_vH	(373–463)		57.6	388		[1926TAY/RIN, 1984BOU/FRI]
C₆H₁₀O₄	[609-02-9]		dimethyl methylmalonate					
		Δ_vH	(278–308)		57.8 ± 0.8	293	GS	[1992VER/BEC]
C₆H₁₀O₄	[124-04-9]		adipic acid					
		$\Delta_{\text{fus}}H$			33.7	419		[2005ROU/TEM]
		$\Delta_{\text{fus}}H$			34.85	426.4		[1991ACR]
		$\Delta_{\text{sub}}H$	(353–373)		124.7 ± 20		ME	[2009TAU/SIT]
		$\Delta_{\text{sub}}H$	(328–368)		145 ± 4		TPD	[2007CAP/LOV]
		$\Delta_{\text{sub}}H$	(285–307)		146.2		TPTD	[2005CHA/ZIE]
		$\Delta_{\text{sub}}H$			NA			[2001ALB]
		$\Delta_{\text{sub}}H$	(295–318)		140		TPTD	[2001CHA/TOB]
								Note: Values based on TPTD method are not consistent with values determined by other experimental methods
		$\Delta_{\text{sub}}H$			133.6 ± 1.3	298	ME	[1999RIB/MON, 1960DAV/THO]
		$\Delta_{\text{sub}}H$	(359–406)		129.3 ± 2.5	383	ME	[1950NIT/SEK2, 1960JON, 1970COX/PIL]
		$\Delta_{\text{sub}}H$	(292–320)		U 37.2	306	A	[1947GRA]
		Δ_vH	(424–503)		105.2	298	CGC	[2005ROU/TEM]
		Δ_vH	(432–611)		92.0	447	A	[1987STE/MAL, 1947STU]
C₆H₁₀O₅	[498-07-7]		(<i>l</i>) glucosane					
		Δ_vH	(468–528)		92.2	483	A	[1987STE/MAL, 1964ENS/DUR, 1972DYK]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
C ₆ H ₁₀ O ₅	[498-07-7]	1,6-anhydro- β -(<i>d</i>)-glucose				
	$\Delta_{\text{sub}}H$	(344–386)	125.1 ± 1.0	365	ME	[1999OJA/SUU]
	$\Delta_{\text{sub}}H$	(386–405)	100.3 ± 5.9	395	ME	[1999OJA/SUU]
C ₆ H ₁₀ O ₅	[na]	1,6-anhydro- β -(<i>d</i>)-gulopyranose				
	$\Delta_{\text{fus}}H$		24.5	404		[1970SHA/MCG]
C ₆ H ₁₀ O ₅	[617-55-0]	<i>(l)</i> malic acid, dimethyl ester				
	$\Delta_{\text{v}}H$	(348–516)	58.7	363	A	[1987STE/MAL, 1947STU]
C ₆ H ₁₀ O ₅	[na]	methyl[1-(methoxycarbonyl)ethyl]carbonate				
	$\Delta_{\text{v}}H$	(358–483)	55.9	373	A	[1987STE/MAL]
C ₆ H ₁₀ O ₆	[608-68-4]	<i>(d)</i> dimethyl tartrate				
	$\Delta_{\text{fus}}H$		17.36	322.2	DTA	[1981CHI/GAR, 1991CHI/BRA]
	$\Delta_{\text{sub}}H$	(310–320)	77.4 ± 8	315	HSA	[1981CHI/GAR]
	$\Delta_{\text{sub}}H$	(308–317)	U 113	312		[1954CRO/JON, 1977PED/RYL]
	$\Delta_{\text{sub}}H$		88.3			[1938WOL/WEG, 1960JON]
	$\Delta_{\text{sub}}H$		85.8			[1937DUN/WOL]
	$\Delta_{\text{v}}H$	(322–365)	76.4	337	A,ME	[1987STE/MAL, 1954CRO/JON]
	$\Delta_{\text{v}}H$	(375–553)	66.0	390		[1947STU]
C ₆ H ₁₀ O ₆	[609-69-5]	<i>(dl)</i> dimethyl tartrate				
	$\Delta_{\text{fus}}H$		26.94	360.2	DTA	[1981CHI/GAR, 1991CHI/BRA]
	$\Delta_{\text{sub}}H$	(314–339)	112 ± 5.6	326	HSA	[1981CHI/GAR]
	$\Delta_{\text{sub}}H$	(315–358)	113.8	336	ME	[1954CRO/JON, 1977PED/RYL]
	$\Delta_{\text{sub}}H$		U 95.0			[1938WOL/WEG, 1960JON]
	$\Delta_{\text{sub}}H$		U 92.5			[1937DUN/WOL]
	$\Delta_{\text{v}}H$	(373–555)	62.5	388	A	[1987STE/MAL, 1947STU]
C ₆ H ₁₀ O ₆	[na]	<i>meso</i> -dimethyl tartrate				
	$\Delta_{\text{sub}}H$		98.3			[1938WOL/WEG, 1960JON]
	$\Delta_{\text{sub}}H$		95.8			[1937DUN/WOL]
C ₆ H ₁₀ O ₆	[na]	<i>(d)</i> -galactono-1,4-lactone				
	$\Delta_{\text{fus}}H$		35.77	410.3		[2004FLO/AMA]
C ₆ H ₁₀ O ₆	[na]	<i>(l)</i> -galactono-1,4-lactone				
	$\Delta_{\text{fus}}H$		35.98	409.8		[2004FLO/AMA]
C ₆ H ₁₀ O ₆	[na]	<i>(d)</i> -gulono-1,4-lactone				
	$\Delta_{\text{fus}}H$		40.13	459.3		[2004FLO/AMA]
C ₆ H ₁₀ O ₆	[na]	<i>(l)</i> -gulono-1,4-lactone				
	$\Delta_{\text{fus}}H$		41.5	459		[2004FLO/AMA]
C ₆ H ₁₀ O ₆	[na]	<i>(l)</i> -mannono-1,4-lactone				
	$\Delta_{\text{fus}}H$		36.13	426.5		[2004FLO/AMA]
C ₆ H ₁₀ S	[592-88-1]	diallyl sulfide				
	$\Delta_{\text{v}}H$	(263–411)	46.6	278		[1999DYK/SVO]
	$\Delta_{\text{v}}H$	(263–412)	43.2	278	A	[1987STE/MAL, 1947STU]
C ₆ H ₁₁ Br	[108-85-0]	bromocyclohexane				
	$\Delta_{\text{fus}}H$		10.79	216.9		[1995KOB/OGU]
		(347–439)	42.8	362		[1997ART/LAF]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C₆H₁₁BrO₂	[600-00-0]	ethyl 2-bromo-2-methylpropionate				
	$\Delta_v H$	(283–437)	45.4	298	A	[1987STE/MAL, 1947STU]
C₆H₁₁Cl	[542-18-7]	chlorocyclohexane				
	$\Delta_{\text{trs}}H$		7.88	221.1		
	$\Delta_{\text{fus}}H$		1.67	228.7	DSC	[2008SIN/MUR]
	$\Delta_{\text{trs}}H$		0.05	120		
	$\Delta_{\text{trs}}H$		8.01	220.4		
	$\Delta_{\text{fus}}H$	(5–304)	2.04	229.3	AC	[1994DIK/KAB]
	$\Delta_v H$	(313–353)	41.8	298	CGC	[1995CHI/HOS]
	$\Delta_v H$		40.7 ± 0.1	298	C	[1995XUW/DAJ]
	$\Delta_v H$		42.9 ± 0.6	298	C	[1994DIK/KAB]
	$\Delta_v H$	(350–416)	39.3	365	A	[1987STE/MAL, 1969AND/BRA]
C₆H₁₁Cl	[6196-85-6]	1-chloro-1-methylcyclopentane				
	$\Delta_v H$		39.7 ± 0.1	297	C	[1997BLO/KAB]
C₆H₁₁ClO	[2736-40-5]	diethylacetyl chloride				
	$\Delta_v H$	(313–412)	39.4	328	A	[1987STE/MAL]
C₆H₁₁ClO	[2177-22-2]	3-ethyl-3-(chloromethyl)oxetane				
	$\Delta_v H$		49.7 ± 0.2	298	C	[1971RIN/SUN]
C₆H₁₁ClO₂	[17696-64-9]	chloroacetic acid, sec-butyl ester				
	$\Delta_v H$	(290–441)	49.6	305	A	[1987STE/MAL]
C₆H₁₁ClO₂	[13361-38-8]	chloroacetic acid, isobutyl ester				
	$\Delta_v H$	(293–323)	43.9	308	A	[1987STE/MAL]
C₆H₁₁F	[372-46-3]	fluorocyclohexane				
	$\Delta_{\text{trs}}H$		7.82	186.7		
	$\Delta_{\text{fus}}H$		2.58	285.3		[1986GON/SZW]
	$\Delta_v H$	(271–301)	37.5 ± 0.3	298	GS	[1997SCH/VER]
	$\Delta_v H$	(316–373)	35.0	331	A	[1987STE/MAL]
C₆H₁₁FO₂	[1578-57-0]	2-fluorohexanoic acid				
	$\Delta_v H$	(387–411)	80.9	399	A	[1987STE/MAL]
C₆H₁₁FO₅	[na]	2-deoxy-2-fluoro-(<i>d</i>)-glucopyranose				
	$\Delta_{\text{fus}}H$		38.2	427.2		[1996SCH]
C₆H₁₁FO₅	[na]	6-deoxy-6-fluoro-(<i>d</i>)-glucopyranose				
	$\Delta_{\text{fus}}H$		27.2	412.2		[1996SCH]
C₆H₁₁FO₅	[na]	3-deoxy-3-fluoro-(<i>d</i>)-glucopyranose				
	$\Delta_{\text{fus}}H$		18.3	378.2		[1996SCH]
C₆H₁₁I	[626-62-0]	iodocyclohexane				
	$\Delta_v H$	(313–353)	48.3	298	CGC	[1995CHI/HOS]
	$\Delta_v H$	(358–408)	43.0	383	A,I	[1987STE/MAL, 1956BRE/UBB]
C₆H₁₁N	[628-73-9]	hexanenitrile				
	$\Delta_v H$	(371–442)	43.3	386	A,EB	[1987STE/MAL, 1973MEY/HOT]
	$\Delta_v H$	(344–441)	44.6	359	EB	[1971MEY/REN]
	$\Delta_v H$		47.9 ± 0.1	298	C	[1970HOW/WAD]
	$\Delta_v H$	(365–437)	49.1	298	EB	[1949DRE/SHR, 1949DRE/MAR, 2005EME/VER]
	$\Delta_v H$	(293–452)	47.7 ± 0.1	298	MM	[1933HEL, 2005EME/VER]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₆ H ₁₁ N	[542-54-1]	4-methylvaleronitrile				
	$\Delta_v H$	(332–430)	35.7	347	A	[1987STE/MAL]
C ₆ H ₁₁ NO	[100-64-1]	cyclohexanone oxime				
	$\Delta_{\text{fus}} H$		12.7	362.5		[2002STE/CHI6]
	$\Delta_{\text{fus}} H$		12.45	362.2		[2008ZAI/PAU]
	$\Delta_{\text{trs}} H$		0.01	240.8		
	$\Delta_{\text{trs}} H$		0.09	273.4		
	$\Delta_{\text{fus}} H$		12.7	362.6		[1992KOZ/KAB]
	$\Delta_{\text{sub}} H$		74.0 ± 0.3	354	C	[1992KOZ/KAB]
	$\Delta_{\text{sub}} H$	(288–348)	79.9 ± 0.7	317	ME	[1992KOZ/KAB]
	$\Delta_{\text{sub}} H$		76.5 ± 1.0	378		[2002STE/CHI6]
	$\Delta_{\text{sub}} H$		79.0 ± 2.0	298		[2002STE/CHI6]
	$\Delta_v H$		58.7 ± 0.6	368	C	[1992KOZ/KAB]
	$\Delta_v H$	(371–446)	59.5 ± 0.5			[1992KOZ/KAB]
	$\Delta_v H$	(370–385)	63.1 ± 1.0	298		[2002STE/CHI6]
C ₆ H ₁₁ NO	[105-60-2]	ϵ -caprolactam				
	$\Delta_{\text{fus}} H$		16.16	342.2		[2007SHE/ZAI]
	$\Delta_{\text{fus}} H$		16.2	342.3	DSC	[2002STE/CHI3]
	$\Delta_{\text{fus}} H$		16.1	343.3		[1992KAB/KOZ]
	$\Delta_{\text{sub}} H$	(293–338)	86.3	316	ME	[2006ZAI/PAU]
	$\Delta_{\text{sub}} H$	(302–339)	86.9	320	GS	[2006ZAI/PAU]
	$\Delta_{\text{sub}} H$	(330–340)	89.3 ± 0.8	335	ME	[1992KAB/KOZ]
	$\Delta_{\text{sub}} H$		86.3 ± 0.2	338	C	[1992KAB/KOZ]
	$\Delta_{\text{sub}} H$		87.3 ± 0.2	298		[1992KAB/KOZ]
	$\Delta_{\text{sub}} H$	(258–308)	77.5	273	A	[1987STE/MAL]
	$\Delta_{\text{sub}} H$	(294–314)	83.3 ± 0.8			[1953AIH, 1960JON, 1970COX/PIL, 1960AIH2]
	$\Delta_v H$	(350–568)	69.2 ± 0.3	360	EB	[2002STE/CHI3]
	$\Delta_v H$	(350–568)	65.7 ± 0.3	400	EB	[2002STE/CHI3]
	$\Delta_v H$	(350–568)	62.3 ± 0.2	440	EB	[2002STE/CHI3]
	$\Delta_v H$	(350–568)	59.0 ± 0.2	480	EB	[2002STE/CHI3]
	$\Delta_v H$	(350–568)	55.7 ± 0.3	520	EB	[2002STE/CHI3]
	$\Delta_v H$	(350–568)	52.4 ± 0.5	560	EB	[2002STE/CHI3]
	$\Delta_v H$	(373–543)	62.3	388	A	[1987STE/MAL]
C ₆ H ₁₁ NO	[820-99-5]	<i>cis</i> 2-hexenoic acid amide				
	$\Delta_{\text{sub}} H$	(323–333)	80.0	328	A	[1987STE/MAL]
	$\Delta_v H$	(343–383)	61.7	358	A	[1987STE/MAL]
C ₆ H ₁₁ NO	[19841-69-3]	<i>trans</i> 2-hexenoic acid amide				
	$\Delta_{\text{sub}} H$	(353–393)	55.8	368	A	[1987STE/MAL]
C ₆ H ₁₁ NO	[931-20-4]	1-methyl-2-piperidone				
	$\Delta_v H$		60.3 ± 0.9	298	C	[2006RIB/CAB]
	$\Delta_v H$	(341–385)	55.4	356	A	[1987STE/MAL]
C ₆ H ₁₁ NO	[1445-73-4]	1-methyl-4-piperidone				
	$\Delta_v H$		54.2 ± 1.0	298	C	[2006RIB/CAB]
C ₆ H ₁₁ NO	[5693-62-9]	2,3,4,5-tetrahydro-6-methoxypyridine				
	$\Delta_v H$	(292–338)	42.8	307	A	[1987STE/MAL]
C ₆ H ₁₁ NO ₂	[1122-60-7]	nitrocyclohexane				

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(298–318)	54.7 ± 0.6	298	GS	[1997VER3]
C ₆ H ₁₁ NO ₂	[52-52-8]	1-aminocyclopentanecarboxylic acid				
	$\Delta_{\text{sub}} H$		123.4 ± 4	455	ME	[1965SVE/CLY, 1964CLY/SVE]
	$\Delta_{\text{sub}} H$	(443–468)	123.3	455	A	[1987STE/MAL]
C ₆ H ₁₁ NO ₂	[na]	lactic acid N-allyl amide				
	$\Delta_v H$	(359–419)	78.2	374	A	[1987STE/MAL]
C ₆ H ₁₁ NO ₂	[na]	5,5-dimethylperhydro-1,3-oxazine-2-one				
	$\Delta_{\text{fus}} H$		28.5	399		[1996LEB/SMI]
C ₆ H ₁₁ NO ₃	[1906-82-7]	ethyl acetamidoacetate				
	$\Delta_v H$	(383–466)	69.4	398	A	[1987STE/MAL, 1972DYK]
C ₆ H ₁₁ NO ₃	[1596-84-5]	N-dimethylaminosuccinamic acid				
	$\Delta_{\text{fus}} H$		36.97	431.4	DSC	[1990DON/DRE]
C ₆ H ₁₁ N ₂ O ₃ PS ₂	[950-37-8]	S-2,3-dihydro-5-methoxy-2-oxo-1,3,4-thiadiazol-3-ylmethyl O,O-dimethyl phosphorodithioate				
	$\Delta_{\text{fus}} H$		28.54	315.1	DSC	[1990DON/DRE]
C ₆ H ₁₁ NS	[13070-07-0]	2-piperidinethione				
	$\Delta_{\text{sub}} H$	(363–370)	81.2 ± 2.9	366	B	[1974BEA/MUE]
	$\Delta_v H$	(363–370)	63.3	366	A	[1987STE/MAL]
C ₆ H ₁₁ NS	[19766-29-1]	2,3,4,5-tetrahydro-(methylthio)pyridine				
	$\Delta_v H$	(313–351)	52.6	328	A	[1987STE/MAL]
C ₆ H ₁₁ N ₃ O ₆	[62154-78-3]	2,3,3-trinitro-2-methylpentane				
	$\Delta_{\text{sub}} H$		90.8	298		[1999MIR/VOR]
C ₆ H ₁₁ N ₅ O ₈	[1924-47-6]	N-(2,2-dinitropropyl)-2,2-dinitro-1-propanamine				
	$\Delta_{\text{sub}} H$		105.4 ± 4.2			[1973DEK/OON, 1977PED/RYL]
C ₆ H ₁₂	[4806-61-5]	ethylcyclobutane				
	$\Delta_v H$		31.2 ± 0.2	298	C	[1983FUC/HAL]
	$\Delta_v H$		32.6 ± 0.8	298	EB	[1974GOO/MOO]
C ₆ H ₁₂	[110-82-7]	cyclohexane				
	$\Delta_{\text{trs}} H$		6.74	186.1		
	$\Delta_{\text{fus}} H$		2.68	279.8		[1996DOM/HEA]
	$\Delta_{\text{trs}} H$		6.73	186		[1984DOM/EVA]
	$\Delta_{\text{sub}} H$	(223–280)	27.6	265	A	[1987STE/MAL]
	$\Delta_{\text{sub}} H$		36.4 ± 0.7		B	[1974JAC]
	$\Delta_{\text{sub}} H$		46.6	186	B	[1963BON]
	$\Delta_{\text{sub}} H$	(268–278)	37.2	273		[1960JON]
	$\Delta_{\text{sub}} H$	(228–268)	37.7	248	A	[1947STU]
	$\Delta_{\text{sub}} H$	(269–279)	36.5	274	A	[1934ROT/NAG]
	$\Delta_v H$	(296–353)	33.1	315	EB	[2009GIE/KOS]
	$\Delta_v H$	(300–345)	32.7	315		[2002LUB/BAN]
	$\Delta_v H$	(360–470)	32.2	375		[1993LEE/HOL]
	$\Delta_v H$	(313–336)	31.9	324	EB	[1995DIO/SAN]
	$\Delta_v H$	(313–336)	33.1	298	EB	[1995DIO/SAN]
	$\Delta_v H$		32.3	314	C	[1988DON/LIN]
	$\Delta_v H$		31.1	332	C	[1988DON/LIN]
$\Delta_v H$		30.3	345	C	[1988DON/LIN]	
$\Delta_v H$		30.0	355	C	[1988DON/LIN]	
$\Delta_v H$	(353–414)	30.9	368	A	[1987STE/MAL]	

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	Enthalpy							
		$\Delta_v H$		(412–491)	29.6	427	A	[1987STE/MAL]
		$\Delta_v H$		(489–553)	29.6	504	A	[1987STE/MAL]
		$\Delta_v H$			33.0 ± 0.1	298	C	[1982FUR/SAK]
		$\Delta_v H$			33.0	298		[1981SHI/SAI]
		$\Delta_v H$			33.0 ± 0.1	298	C	[1979MAJ/SVO]
		$\Delta_v H$			32.3 ± 0.1	313	C	[1979MAJ/SVO]
		$\Delta_v H$			31.2 ± 0.1	333	C	[1979MAJ/SVO]
		$\Delta_v H$			31.0 ± 0.1	338	C	[1979MAJ/SVO]
		$\Delta_v H$			30.4 ± 0.1	348	C	[1979MAJ/SVO]
		$\Delta_v H$			30.1 ± 0.1	353	C	[1979MAJ/SVO]
		$\Delta_v H$			32.2 ± 0.1	313	C	[1973SVO/VES]
		$\Delta_v H$			31.9 ± 0.1	323	C	[1973SVO/VES]
		$\Delta_v H$			31.1 ± 0.1	333	C	[1973SVO/VES]
		$\Delta_v H$			30.6 ± 0.1	343	C	[1973SVO/VES]
		$\Delta_v H$			30.1 ± 0.1	354	C	[1973SVO/VES]
		$\Delta_v H$			32.9 ± 0.3	298	ME	[1972SAB/CHA]
		$\Delta_v H$			32.9	298		[1971MOR]
		$\Delta_v H$			33.0	298		[1971WIL/ZWO]
		$\Delta_v H$		(303–343)	32.5	318		[1968GAW/SWI]
		$\Delta_v H$		(298–348)	32.9	313		[1967CRU/CUT]
		$\Delta_v H$		(316–354)	32.8	331		[1965MAR/SUS]
		$\Delta_v H$			33.0 ± 0.1	298	C	[1960WAD]
		$\Delta_v H$			31.4 ± 0.1	324	C	[1951MCC/PER]
		$\Delta_v H$			30.4 ± 0.1	346	C	[1951MCC/PER]
		$\Delta_v H$			33.0	298	C	[1947OSB/GIN]
		$\Delta_v H$			30.1	354		[1946SPI/PIT]
		$\Delta_v H$		(293–355)	32.9	308	A,MM	[1987STE/MAL, 1945WIL/TAY]
		$\Delta_v H$			33.3 ± 0.1	298	C	[1943AST/SZA]
		$\Delta_v H$			33.5	298		[1927NAG]
C₆D₁₂	[1735-17-7]		cyclohexane-d ₁₂					
		$\Delta_v H$		(283–353)	33.1	298		[1953DAV/SCH]
C₆H₁₂	[96-37-7]		methylcyclopentane					
		$\Delta_{\text{fus}} H$			6.93	130.7		[1996DOM/HEA]
		$\Delta_v H$		(300–345)	31.4	315		[2010SAP/UUS]
		$\Delta_v H$			31.6	298		[1971WIL/ZWO]
		$\Delta_v H$			31.3 ± 0.1	304	C	[1959MCC/PEN]
		$\Delta_v H$			30.2 ± 0.1	326	C	[1959MCC/PEN]
		$\Delta_v H$			29.1 ± 0.1	345	C	[1959MCC/PEN]
		$\Delta_v H$			31.6 ± 0.1	298	C	[1947OSB/GIN]
		$\Delta_v H$		(288–346)	31.9	303	A,MM	[1987STE/MAL, 1945WIL/TAY]
C₆H₁₂	[592-41-6]		1-hexene					
		$\Delta_{\text{fus}} H$			9.35	133.4		[1996DOM/HEA]
		$\Delta_v H$		(298–336)	30.4	313		[2009MAR/AUC]
		$\Delta_v H$		(300–337)	30.6	315		[2001SEG/LAM]
		$\Delta_v H$		(273–343)	31.6	288	A	[1987STE/MAL]
		$\Delta_v H$			30.6	298		[1971WIL/ZWO]
		$\Delta_v H$		(289–337)	30.6	298		[1956CAM/ROS]
		$\Delta_v H$		(289–337)	31.0	304	MM	[1950FOR/CAM]
C₆H₁₂	[7688-21-3]		<i>cis</i> 2-hexene					
		$\Delta_{\text{fus}} H$			8.88	132		[1990MES/TOD]
		$\Delta_v H$		(278–343)	32.2	293	A	[1987STE/MAL]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$		31.5	298		[1971WIL/ZWO]
	$\Delta_v H$	(298–342)	31.5	298		[1956CAM/ROS]
C ₆ H ₁₂	[4050-45-7]	<i>trans</i> 2-hexene				
	$\Delta_v H$	(283–342)	32.2	298	A	[1987STE/MAL]
	$\Delta_v H$		31.6	298		[1971WIL/ZWO]
	$\Delta_v H$	(292–341)	31.5	298		[1956CAM/ROS]
C ₆ H ₁₂	[7642-09-3]	<i>cis</i> 3-hexene				
	$\Delta_v H$	(276–348)	32.1	291	A	[1987STE/MAL]
	$\Delta_v H$		31.3	298		[1971WIL/ZWO]
C ₆ H ₁₂	[13269-52-8]	<i>trans</i> 3-hexene				
	$\Delta_v H$	(278–341)	32.3	293	A	[1987STE/MAL]
	$\Delta_v H$		31.6	298		[1971WIL/ZWO]
C ₆ H ₁₂	[763-29-1]	2-methyl-1-pentene				
	$\Delta_v H$	(272–341)	31.6	287	A	[1987STE/MAL]
	$\Delta_v H$		30.5	298		[1971WIL/ZWO]
C ₆ H ₁₂	[760-20-3]	3-methyl-1-pentene				
	$\Delta_v H$	(265–333)	30	280	A	[1987STE/MAL]
	$\Delta_v H$		28.6	298		[1971WIL/ZWO]
C ₆ H ₁₂	[691-37-2]	4-methyl-1-pentene				
	$\Delta_{\text{fus}} H$		4.93	118.9		[1994LEB/SMI3]
	$\Delta_v H$	(310–360)	28.6 ± 0.2	298	EB	[1997STE/CHI]
C ₆ H ₁₂	$\Delta_v H$	(310–360)	27.4 ± 0.3	320	EB	[1997STE/CHI]
	$\Delta_v H$	(310–360)	26.2 ± 0.4	340	EB	[1997STE/CHI]
	$\Delta_v H$	(310–360)	24.9 ± 0.5	360	EB	[1997STE/CHI]
	$\Delta_v H$	(265–333)	30.1	280	A	[1987STE/MAL]
	$\Delta_v H$		28.7	298		[1971WIL/ZWO]
	$\Delta_v H$	(287–328)	28.7	298		[1956PEN/SCO]
	$\Delta_v H$		28.7	298		[1956PEN/SCO]
C ₆ H ₁₂	[625-27-4]	2-methyl-2-pentene				
	$\Delta_v H$	(277–346)	32.4	292	A	[1987STE/MAL]
	$\Delta_v H$		31.6	298		[1971WIL/ZWO]
C ₆ H ₁₂	[922-62-3]	<i>cis</i> 3-methyl-2-pentene				
	$\Delta_v H$	(277–347)	32.2	292	A	[1987STE/MAL]
	$\Delta_v H$		31.3	298		[1971WIL/ZWO]
C ₆ H ₁₂	[616-12-6]	<i>trans</i> 3-methyl-2-pentene				
	$\Delta_v H$	(280–349)	32.8	295	A	[1987STE/MAL]
	$\Delta_v H$		32.1	298		[1971WIL/ZWO]
C ₆ H ₁₂	[691-38-3]	<i>cis</i> 4-methyl-2-pentene				
	$\Delta_v H$	(267–330)	30.8	282	A	[1987STE/MAL]
	$\Delta_v H$		29.5	298		[1971WIL/ZWO]
C ₆ H ₁₂	[674-76-0]	<i>trans</i> 4-methyl-2-pentene				
	$\Delta_v H$	(300–330)	29.5	298		[1956CAM/ROS]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound						
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference		
C ₆ H ₁₂		$\Delta_v H$	(269–337)	31.2	284	A	[1987STE/MAL]	
		$\Delta_v H$		30.0	298		[1971WIL/ZWO]	
		$\Delta_v H$	(291–332)	30.0	298		[1956CAM/ROS]	
C ₆ H ₁₂	[563-78-0]	2,3-dimethyl-1-butene						
		$\Delta_v H$	(267–335)	30.5	282	A	[1987STE/MAL]	
		$\Delta_v H$		29.2	298		[1971WIL/ZWO]	
C ₆ H ₁₂		$\Delta_v H$	(289–329)	29.2	298		[1956CAM/ROS]	
	[558-37-2]	3,3-dimethyl-1-butene						
		$\Delta_{\text{trs}} H$		4.35	124.9			
		$\Delta_{\text{fus}} H$		1.09	158.4		[1996DOM/HEA]	
		$\Delta_v H$	(254–316)	28.6	269	A	[1987STE/MAL]	
C ₆ H ₁₂		$\Delta_v H$		26.6	298		[1971WIL/ZWO]	
		$\Delta_v H$	(264–314)	27.4	298		[1971BAG/MAL]	
		$\Delta_v H$	(281–315)	26.6	298		[1956CAM/ROS]	
	[563-79-1]	2,3-dimethyl-2-butene						
		$\Delta_{\text{trs}} H$		3.53	196.8			
		$\Delta_{\text{fus}} H$		6.44	198.9		[1996DOM/HEA]	
		$\Delta_v H$	(313–346)	32.1	328		[2004UUS/POK]	
		$\Delta_v H$	(289–347)	32.6	298		[1956CAM/ROS]	
C ₆ H ₁₂		$\Delta_v H$		32.5	298		[1971WIL/ZWO]	
		$\Delta_v H$	(289–347)	32.7	304		[1971BAG/MAL]	
		$\Delta_v H$	(282–348)	33.1	297	A	[1987STE/MAL, 1955CUM/MCL]	
		$\Delta_v H$		32.9 ± 0.1	292	C	[1955SCO/FIN]	
		$\Delta_v H$		32.0 ± 0.1	308	C	[1955SCO/FIN]	
		$\Delta_v H$		30.9 ± 0.1	326	C	[1955SCO/FIN]	
		$\Delta_v H$		29.7 ± 0.1	346	C	[1955SCO/FIN]	
	C ₆ H ₁₂	[760-21-4]	2-ethyl-1-butene					
		$\Delta_v H$	(289–338)	31.0	298		[1956CAM/ROS]	
C ₆ H ₁₂ Br ₂	[58133-26-9]	1,1-dibromohexane						
		$\Delta_v H$	(378–526)	51.6	393	A,EST	[1987STE/MAL, 1956MAN, 1972DYK]	
C ₆ H ₁₂ Br ₂	[624-20-4]	1,2-dibromohexane						
		$\Delta_v H$	(363–450)	56.5 ± 2.0	298		[1993VAR/PUC]	
C ₆ H ₁₂ ClNO	[3240-94-6]							
		$\Delta_v H$	(273–333)	53.8	288	A	[1987STE/MAL, 1972DYK]	
C ₆ H ₁₂ Cl ₂	[62017-16-7]	1,1-dichlorohexane						
		$\Delta_v H$	(330–440)	48.7	298		[1987VAR/LOS2, 1991BAS/SVO]	
		$\Delta_v H$	(345–484)	45.1	360	A,EST	[1987STE/MAL, 1956MAN, 1972DYK]	
C ₆ H ₁₂ Cl ₂	[2162-92-7]	<i>(dl)</i> 1,2-dichlorohexane						
		$\Delta_v H$	(350–440)	48.8	298		[1991BAS/SVO]	
		$\Delta_v H$	(352–442)	44.9	367	A	[1987STE/MAL]	
C ₆ H ₁₂ Cl ₂		$\Delta_v H$		47.9 ± 0.7	298	EB	[1975PIS/ROZ2]	
	[2163-00-0]	1,6-dichlorohexane						
		$\Delta_v H$	(380–480)	56.3	298		[1988VAR/LOS, 1991BAS/SVO]	
C ₆ H ₁₂ Cl ₂ O	[108-60-1]	<i>bis</i> (2-chloro-1-methylethyl) ether						
		$\Delta_v H$	(302–456)	53.6	317	A	[1987STE/MAL, 1947STU]	

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₆ H ₁₂ Cl ₂ O ₂	[14689-97-5] $\Delta_v H$	<i>bis</i> (2-chloroethyl)acetaldehyde acetal (329–486)	59.4	344	A	[1987STE/MAL, 1947STU]
C ₆ H ₁₂ Cl ₃ N	[555-77-1] $\Delta_v H$	<i>bis</i> (2-chloroethyl)acetaldehyde acetal (273–333)	65.0	288	A,GS	[1987STE/MAL, 1948RED/CHA3, 1972DYK]
C ₆ H ₁₂ Cl ₃ O ₄ P	[na] $\Delta_v H$	tris(2-chloroethyl)phosphate (293–445)	36.7	308	A	[1987STE/MAL, 1972DYK]
C ₆ H ₁₂ F ₂	[62127-41-7] $\Delta_v H$	1,1-difluorohexane (290–407)	37.7	305	A,EST	[1987STE/MAL, 1956MAN, 1972DYK]
C ₆ H ₁₂ F ₃ OP	[na] $\Delta_v H$	methyl (trifluoromethyl)phosphinous acid, <i>tert</i> -butyl ester (273–329)	39.7	296		[1970BUR/KAN]
C ₆ H ₁₂ F ₃ PS	[26348-87-8] $\Delta_v H$	methyl (trifluoromethyl)phosphinothious acid, <i>tert</i> -butyl ester (296–337)	43.2	312		[1970BUR/KAN]
C ₆ H ₁₂ F ₄ N ₂	[16096-76-7] $\Delta_v H$	N,N,N',N'-tetrafluoro-2-methyl-1,2-pentanediamine (253–293)	42.8	278	A,IP	[1987STE/MAL, 1963GOO/DOU, 1962GOO/DOU]
C ₆ H ₁₂ N ₂	[3010-02-4] $\Delta_v H$	(diethylamino)acetonitrile (283–318)	49.9 ± 0.3		GS	[1997WEL/VER]
C ₆ H ₁₂ N ₂	[280-57-9] $\Delta_{\text{trs}}H$ $\Delta_{\text{fus}}H$ $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$	1,4-diazabicyclo[2.2.2]octane (324–351) (353–369) (323–373)	10.54 7.45 61.9 ± 3.3 52.3 ± 3.3 54.4	351.1 433 338 361 348		[1996DOM/HEA] [1960WAD/KIS, 1970COX/PIL] [1960WAD/KIS, 1970COX/PIL] [1963BON]
C ₆ H ₁₂ N ₂	[na] $\Delta_{\text{sub}}H$	3,3,4,4-tetramethyl- Δ 1-1,2-diazetidine 62.3 ± 1.0	298		C	[1978MON/ENG]
C ₆ H ₁₂ N ₂ O	[7226-23-5] $\Delta_v H$	1,3-dimethyl-3,4,5,6-tetrahydro-2(1 <i>H</i>) pyrimidinone (370–520)	58.0	400	EB	[1987KNE/ZON]
C ₆ H ₁₂ N ₂ O	[18503-52-1] $\Delta_{\text{trs}}H$ $\Delta_{\text{fus}}H$	1,4-diazabicyclo[2.2.2]octane N-oxide 3.4 0.45	418 493		DSC	[1990MIH/BAS, 1987MIH/BAS]
C ₆ H ₁₂ N ₂ O	[2158-03-4] $\Delta_{\text{sub}}H$	1-piperidinecarboxamide 100.2 ± 1.2	298		C	[2007RIB/CAB2]
C ₆ H ₁₂ N ₂ O	[4138-26-5] $\Delta_{\text{sub}}H$	3-piperidinecarboxamide 112.5 ± 1.3	298		C	[2007RIB/CAB2]
C ₆ H ₁₂ N ₂ O	[39546-32-2] $\Delta_{\text{sub}}H$	4-piperidinecarboxamide 123.6 ± 1.3	298		C	[2007RIB/CAB2]
C ₆ H ₁₂ N ₂ OS	[62528-85-2] $\Delta_{\text{fus}}H$ $\Delta_v H$	tetramethyl monothiooxamide 17.0 59.0	350.2 508		DSC TGA,DSC	[2003CLO/JAN] [2003CLO/JAN]
C ₆ H ₁₂ N ₂ O ₂	[608-14-6] $\Delta_{\text{fus}}H$ $\Delta_v H$	N,N,N',N'-tetramethyloxamide 18.0 52.5	352.2 460			[2003CLO/JAN] [2003CLO/JAN]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₆ H ₁₂ N ₂ O ₂	[628-94-4]	adipamide				
	$\Delta_{\text{trs}}H$		1.59	458.4		
	$\Delta_{\text{fus}}H$		52.72	499.1	DSC	[2006BAD/DEL]
C ₆ H ₁₂ N ₂ O ₃	[na]	β -alanyl- β -alanine				
	$\Delta_{\text{fus}}H$		58.3	480.1		[1996DOM/HEA]
C ₆ H ₁₂ N ₂ O ₃	[na]	α -alanyl- α -alanine (<i>dl</i>)				
	$\Delta_{\text{fus}}H$		33.2	483.2		[1996DOM/HEA]
C ₆ H ₁₂ N ₂ O ₄	[3964-18-9]	2,3-dinitro-2,3-dimethylbutane				
	$\Delta_{\text{trs}}H$		1.0	322		
	$\Delta_{\text{trs}}H$		18.0	389		
	$\Delta_{\text{fus}}H$		8.8	473		[2002JON/LIG]
	$\Delta_{\text{sub}}H$		74 ± 5		TGA	[2002JON/LIG]
	$\Delta_{\text{sub}}H$		79.5 ± 0.8	298		[1999MIR/VOR]
	$\Delta_{\text{sub}}H$	(303–330)	85 ± 2		ME	[1994SMI/MAT, 2002JON/LIG]
	$\Delta_{\text{sub}}H$	(253–323)	94		GC	[1991ELI, 2002JON/LIG]
C ₆ H ₁₂ N ₂ O ₆	[99115-63-6]	2,5-hexanediol dinitrate				
	$\Delta_{\text{sub}}H$	(293–313)	119	303	A	[1987STE/MAL, 1957KEM/GOL, 1972DYK]
	Δ_vH	(293–313)	54.4	303	B,GS	[1957KEM/GOL, 1972DYK]
C ₆ H ₁₂ N ₂ O ₈	[111-22-8]	triethylene glycol dinitrate				
	Δ_vH	(303–348)	88.3	318	A	[1987STE/MAL, 1972DYK, 1963WOO/ADI]
C ₆ H ₁₂ N ₂ S ₂	[35840-78-9]	tetramethyl dithiooxamide				
	$\Delta_{\text{fus}}H$		21.0	409.2	DSC	[2003CLO/JAN]
	Δ_vH		60.5	533	TGA,DSC	[2003CLO/JAN]
C ₆ H ₁₂ N ₄	[100-97-0]	1,3,5,7-tetraazatricyclo[3.3.1.1 ^{3,7}]decane				
	$\Delta_{\text{sub}}H$	(339–378)	77.7 ± 0.4	359	GS	[2002VER2]
	$\Delta_{\text{sub}}H$	(339–378)	79.6 ± 0.4	298	GS	[2002VER2]
	$\Delta_{\text{sub}}H$	(298–453)	76.8	313	A	[1987STE/MAL]
	$\Delta_{\text{sub}}H$	(302–328)	78.8	316	TE,ME	[1983DEW/VAN]
	$\Delta_{\text{sub}}H$		74.9 ± 2.9	298		[1960WAD/KIS, 1970MAN/RAP]
	$\Delta_{\text{sub}}H$	(281–298)	74.1 ± 0.8	289	TE	[1960BUD]
	$\Delta_{\text{sub}}H$		75.3			[1958KLI/STR]
C ₆ H ₁₂ O	[100-38-9]	<i>(dl)</i> 2,5-dimethyltetrahydrofuran				
	Δ_vH	(278–370)	35.4	293	A	[1987STE/MAL]
C ₆ H ₁₂ O	[1436-34-6]	1,2-epoxyhexane				
	Δ_vH	(300–390)	43.1	315	A	[1987STE/MAL, 1969VOJ/CIH, 1984BOU/FRI]
C ₆ H ₁₂ O	[1192-22-9]	2-methyl-2,3-epoxypentane				
	Δ_vH	(306–369)	40.6	321	A	[1987STE/MAL]
C ₆ H ₁₂ O	[6140-80-3]	allyl isopropyl ether				
	Δ_vH	(253–415)	36.1	268	A	[1987STE/MAL]
	Δ_vH	(229–353)	36.8	244	A	[1987STE/MAL, 1947STU]
C ₆ H ₁₂ O	[1471-03-0]	allyl propyl ether				
	Δ_vH	(261–428)	37.5	276	A	[1987STE/MAL]
	Δ_vH	(234–364)	36.4	249	A	[1987STE/MAL, 1947STU]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₆ H ₁₂ O	[111-34-2]	butyl vinyl ether				
	$\Delta_v H$	(311–403)	36.7 ± 0.2	298	EB	[1996STE/CHI2]
	$\Delta_v H$	(311–403)	35.2 ± 0.2	320	EB	[1996STE/CHI2]
	$\Delta_v H$	(311–403)	32.5 ± 0.2	360	EB	[1996STE/CHI2]
	$\Delta_v H$	(311–403)	29.6 ± 0.2	400	EB	[1996STE/CHI2]
	$\Delta_v H$	(353–393)	36.5	298	CGC	[1995CHI/HOS]
		(269–368)	36.1	284	A	[1987STE/MAL]
C ₆ H ₁₂ O	[109-53-5]	isobutyl vinyl ether				
$\Delta_v H$		(266–357)	37.4	281	A	[1987STE/MAL]
C ₆ H ₁₂ O	[108-93-0]	cyclohexanol				
	$\Delta_{\text{trs}} H$		8.7	245.2		
	$\Delta_{\text{fus}} H$		1.73	298.2	DSC	[2009SIN/MUR]
	$\Delta_{\text{trs}} H$		8.21	263.5		
	$\Delta_{\text{fus}} H$		1.7	297		[1984PIN/POS, 1968ADA/SUG, 1996DOM/HEA]
	$\Delta_{\text{sub}} H$	(272–298)	60.7	285	A	[1987STE/MAL, 1948NIT/SEK2]
	$\Delta_v H$	(322–433)	60.1	337		[2004STE/SUN]
	$\Delta_v H$	(390–430)	49.8	405		[2002SWI/MAL]
	$\Delta_v H$		62.0 ± 0.3	298	C	[1999COS/EUS]
	$\Delta_v H$	(288–328)	61.2 ± 0.6	308	GS	[1998VER5]
	$\Delta_v H$	(288–328)	61.8 ± 0.6	298	GS	[1998VER5]
	$\Delta_v H$	(341–471)	63.5 ± 0.7	298	EB	[1997STE/CHI3]
	$\Delta_v H$	(323–373)	61.3	298	CGC	[1995CHI/HOS]
	$\Delta_v H$	(350–456)	55.0	365	EB	[1987AMB/GHI2]
	$\Delta_v H$	(318–434)	59.9	333	A	[1987STE/MAL]
	$\Delta_v H$	(300–434)	62.7	315	A	[1987STE/MAL]
	$\Delta_v H$	(404–432)	49.3	418		[1984CAS/FRA2]
	$\Delta_v H$	(303–373)	58.4	318		[1984SIP/WIE]
	$\Delta_v H$	(299–319)	60.4	309		[1975CAB/CON2]
			62.0 ± 0.9	298		[1975CAB/CON2]
$\Delta_v H$		62.0 ± 0.2	298	C	[1968PLA/WIL]	
$\Delta_v H$		62.0 ± 0.3	298		[1966WAD]	
$\Delta_v H$		62.0 ± 0.2	298	C	[1962SEL/SUN]	
$\Delta_v H$	(367–433)	52.6	382		[1960NOV/MAT2, 1960NOV/MAT]	
$\Delta_v H$	(307–422)	54.8	322		[1946THO]	
C ₆ H ₁₂ O	[1462-03-9]	1-methylcyclopentanol				
	$\Delta_{\text{fus}} H$		8.41	310.2		[1985WIB/WAS]
	$\Delta_v H$	(354–407)	45.7	369	A	[1987STE/MAL]
C ₆ H ₁₂ O	[821-41-0]	5-hexen-1-ol				
	$\Delta_v H$		60.2 ± 0.1	298	C	[1996ULB/KLU]
	$\Delta_v H$		58.0 ± 0.1	343	C	[1996ULB/KLU]
			55.7 ± 0.1	358	C	[1996ULB/KLU]
C ₆ H ₁₂ O	[565-61-7]	(dl) 3-methyl-2-pentanone				
	$\Delta_v H$	(286–400)	39.8	301	A	[1987STE/MAL]
	$\Delta_v H$	(283–395)	41.5	298	A	[1987STE/MAL]
	$\Delta_v H$	(385–455)	36.5	400	A	[1987STE/MAL]
		(283–457)	41.2	298		[1975AMB/ELL]
C ₆ H ₁₂ O	[591-78-6]	2-hexanone				
	$\Delta_{\text{fus}} H$		14.9	217.7		[1996DOM/HEA]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	Enthalpy							
		$\Delta_v H$	(359–401)	39.0	374	EB	[2002SII/KIR]	
		$\Delta_v H$		43.1 ± 0.1	298	C	[1992SVO/KUB]	
		$\Delta_v H$		42.5 ± 0.1	308	C	[1992SVO/KUB]	
		$\Delta_v H$		41.6 ± 0.1	323	C	[1992SVO/KUB]	
		$\Delta_v H$		40.7 ± 0.1	338	C	[1992SVO/KUB]	
		$\Delta_v H$		40.1 ± 0.1	348	C	[1992SVO/KUB]	
		$\Delta_v H$		39.5 ± 0.1	358	C	[1992SVO/KUB]	
		$\Delta_v H$	(293–411)	40.8	308	A	[1987STE/MAL]	
		$\Delta_v H$	(279–423)	43.8	294	A	[1987STE/MAL]	
		$\Delta_v H$	(310–427)	41.5	325	A	[1987STE/MAL]	
		$\Delta_v H$	(421–523)	36.7	436	A	[1987STE/MAL]	
		$\Delta_v H$	(513–587)	36.1	528	A	[1987STE/MAL]	
		$\Delta_v H$		43.1 ± 0.1	298	C	[1983UCH/MAJ]	
		$\Delta_v H$		43.0 ± 0.3	298	GCC	[1979SAL/PEA]	
		$\Delta_v H$	(307–482)	42.9	298		[1975AMB/ELL]	
		$\Delta_v H$		42.2 ± 0.1	298	C	[1970HAR/HEA]	
		$\Delta_v H$	(280–400)	53.8	295		[1947STU]	
C₆H₁₂O	[589-38-8]		3-hexanone					
		$\Delta_{\text{trs}} H$		0.68	145			
		$\Delta_{\text{fus}} H$		13.47	217.7		[1996DOM/HEA]	
		$\Delta_v H$	(408–517)	36.5	423	A	[1987STE/MAL]	
		$\Delta_v H$	(511–583)	35.4	526	A	[1987STE/MAL]	
		$\Delta_v H$		40.6 ± 0.1	298	C	[1983UCH/MAJ]	
		$\Delta_v H$		42.3 ± 0.3	298	GCC	[1979SAL/PEA]	
		$\Delta_v H$	(348–413)	38.9	363	A	[1987STE/MAL, 1975AMB/ELL]	
		$\Delta_v H$		42.3	298		[1975AMB/ELL]	
		$\Delta_v H$	(292–406)	42.2	307	A	[1987STE/MAL, 1972DYK]	
		$\Delta_v H$		41.9 ± 0.2	298	C	[1970HAR/HEA]	
		$\Delta_v H$		38.4 ± 0.1	354	C	[1967HAL/LEE]	
		$\Delta_v H$		37.0 ± 0.1	374	C	[1967HAL/LEE]	
		$\Delta_v H$		35.4 ± 0.1	396	C	[1967HAL/LEE]	
		$\Delta_v H$	(349–406)	38.8	364	GS,EB	[1965COL/COU]	
C₆H₁₂O	[75-97-8]		3,3-dimethyl-2-butanone (pinacolone)					
		$\Delta_{\text{fus}} H$		11.34	221.7		[1996DOM/HEA]	
		$\Delta_v H$		37.8 ± 0.1	308	C	[1992SVO/KUB]	
		$\Delta_v H$		37.5 ± 0.1	313	C	[1992SVO/KUB]	
		$\Delta_v H$		36.9 ± 0.1	323	C	[1992SVO/KUB]	
		$\Delta_v H$		36.7 ± 0.1	328	C	[1992SVO/KUB]	
		$\Delta_v H$		35.8 ± 0.1	338	C	[1992SVO/KUB]	
		$\Delta_v H$		35.4 ± 0.1	343	C	[1992SVO/KUB]	
		$\Delta_v H$		35.0 ± 0.1	348	C	[1992SVO/KUB]	
		$\Delta_v H$	(311–381)	36.9	326	A	[1987STE/MAL]	
		$\Delta_v H$	(363–400)	34.9	378	A	[1987STE/MAL]	
		$\Delta_v H$	(396–509)	33.8	411	A	[1987STE/MAL]	
		$\Delta_v H$	(491–567)	33.1	506	A	[1987STE/MAL]	
		$\Delta_v H$	(289–402)	38.3	304	A	[1987STE/MAL, 1975AMB/ELL]	
		$\Delta_v H$		38.3	298		[1975AMB/ELL]	
		$\Delta_v H$		36.1	338	C	[1973GEI/QUI]	
		$\Delta_v H$		37.9 ± 0.1	298	C	[1970HAR/HEA]	
C₆H₁₂O	[108-10-1]		4-methyl-2-pentanone					
		$\Delta_v H$	(321–397)	38.7	336		[2009MAR/LLA]	
		$\Delta_v H$		40.1 ± 0.1	308	C	[1992SVO/KUB]	

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	Enthalpy							
		$\Delta_v H$			39.0 ± 0.1	323	C	[1992SVO/KUB]
		$\Delta_v H$			38.0 ± 0.1	338	C	[1992SVO/KUB]
		$\Delta_v H$			37.4 ± 0.1	348	C	[1992SVO/KUB]
		$\Delta_v H$	(309–416)		39.2	324		[1988AMB/GHI3]
		$\Delta_v H$	(281–400)		42.5	296	A	[1987STE/MAL]
		$\Delta_v H$	(349–389)		37.0	365	EB	[1985RED/RAO]
		$\Delta_v H$			42.5 ± 0.1	298	C	[1983UCH/MAJ]
		$\Delta_v H$	(282–456)		41.0	298		[1975AMB/ELL]
		$\Delta_v H$			37.6	347	C	[1973GEI/QUI]
		$\Delta_v H$	(294–390)		41.2	309	A	[1987STE/MAL, 1952FUG/BOW]
C₆H₁₂O	[565-69-5]		2-methyl-3-pentanone					
		$\Delta_v H$	(300–387)		43.4	315	A	[1987STE/MAL]
		$\Delta_v H$	(280–387)		41.0	295	A	[1987STE/MAL]
		$\Delta_v H$	(377–450)		36.2	392	A	[1987STE/MAL]
		$\Delta_v H$	(280–452)		40.5	298		[1975AMB/ELL]
		$\Delta_v H$			39.8 ± 0.2	298	C	[1970SEL2]
C₆H₁₂O	[66-25-1]		hexanal					
		$\Delta_{\text{fus}}H$			13.3	214.9		
		$\Delta_{\text{trs}}H$ (liq anomaly)			0.34	243.2		[1993LEB/VAS, 1991YAS/BYK]
		$\Delta_v H$	(322–402)		40.8	337	EB	[2006PAL/ORA]
		$\Delta_v H$	(287–309)		42.5 ± 0.4	298	GS	[2003VER/KRA2]
		$\Delta_v H$			42.3 ± 0.1	298		[1981DYA/KOR]
		$\Delta_v H$	(315–402)		41.0	330		[1979MAR/SAC]
C₆H₁₂OS	[926-47-2]		S-butyl thiolacetate					
		$\Delta_v H$			48.1 ± 0.2	298	C	[1966WAD]
C₆H₁₂OS	[999-90-6]		S-tert-butyl thiolacetate					
		$\Delta_v H$			42.9 ± 0.2	298	C	[1966WAD]
C₆H₁₂O₂	[1792-81-0]		cis 1,2-cyclohexanediol					
		$\Delta_{\text{fus}}H$			20.27	373.2	DSC	[2002ZHO/PEN, 2003ZHO/ZHA]
		$\Delta_{\text{trs}}H$			19.9	360.4		
		$\Delta_{\text{fus}}H$			3.3	371.6		[2008MAR/EUS]
		$\Delta_{\text{trs}}H$			19.89	360.4		
		$\Delta_{\text{fus}}H$			3.32	371.6		[1995MAR/COS]
		$\Delta_{\text{sub}}H$			89.0			[1999COS/EUS]
		$\Delta_{\text{sub}}H$ (cryst. I)			70 ± 3.0	366	C	[1995MAR/COS]
		$\Delta_{\text{sub}}H$ (cryst. III)			88.0 ± 1.9	343	C	[1995MAR/COS]
		$\Delta_{\text{sub}}H$	(289–320)		43.7	304	ME	[1987STE/MAL, 1940ZIB]
C₆H₁₂O₂	[1460-57-7]		trans 1,2-cyclohexanediol					
		$\Delta_{\text{fus}}H$			16.37	375.7	DSC	[2002ZHO/PEN, 2003ZHO/ZHA]
		$\Delta_{\text{fus}}H$			21.0	382.6		[2008MAR/EUS]
		$\Delta_{\text{fus}}H$			18.51	372.3		[1995MAR/COS]
		$\Delta_{\text{sub}}H$			85.9 ± 1.4	343	C	[1995MAR/COS]
		$\Delta_{\text{sub}}H$	(289–320)		42.5	304	ME	[1987STE/MAL, 1940ZIB]
C₆H₁₂O₂	[126-39-6]		2-ethyl-2-methyl-1,3-dioxolane					
		$\Delta_v H$	(274–313)		44.8 ± 0.3	298	GS	[2002VER]
		$\Delta_v H$	(274–313)		43.1 ± 0.3		GS	[1998VER/PEN]
C₆H₁₂O₂	[3390-13-4]		2-propyl-1,3-dioxolane					
		$\Delta_v H$	(278–313)		45.3 ± 0.3	298	GS	[1998VER/PEN, 2002VER]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₆ H ₁₂ O ₂	[1121-61-5] $\Delta_v H$	4-ethyl-1,3-dioxane (362–412)	39.3	377	A	[1987STE/MAL]
C ₆ H ₁₂ O ₂	[766-20-1] $\Delta_v H$	2,4-dimethyl-1,3-dioxane (274–313)	44.9 ± 0.6	298	GS	[2002VER]
C ₆ H ₁₂ O ₂	[141-79-7] $\Delta_{\text{fus}} H$	2,2-dimethyl-1,3-dioxane	12.1	229.6		[1975BOR]
C ₆ H ₁₂ O ₂	[766-15-4] $\Delta_v H$ $\Delta_v H$	4,4-dimethyl-1,3-dioxane (333–407) (363–406)	37.1 38.8	348 378	A	[1987STE/MAL, 1968KAC/NEM] [1969LES/MOR]
C ₆ H ₁₂ O ₂	[2391-24-4] $\Delta_v H$	<i>cis</i> 4,5-dimethyl-1,3-dioxane (353–410)	48.5	368	A	[1987STE/MAL]
C ₆ H ₁₂ O ₂	[1121-20-6] $\Delta_v H$	<i>trans</i> 4,5-dimethyl-1,3-dioxane (353–408)	39.1	368	A	[1987STE/MAL]
C ₆ H ₁₂ O ₂	[100-72-1] $\Delta_v H$	2-hydroxymethyltetrahydropyran (344–460)	49.0	359	A	[1987STE/MAL]
C ₆ H ₁₂ O ₂	[6581-66-4] $\Delta_v H$	2-methoxytetrahydropyran	39.3 ± 1.2	298	DSC	[2005ROJ/GIN]
C ₆ H ₁₂ O ₂	[4415-90-1] $\Delta_v H$ $\Delta_v H$ $\Delta_v H$	1,1-dimethoxycyclobutane (274–313) (273–313) (301–348)	42.0 ± 0.3 42.3 ± 0.3 42.6	298 325	GS GS EB	[2002VER] [1998VER/PEN] [1994WIB/MOR]
C ₆ H ₁₂ O ₂	[4016-14-2] $\Delta_v H$	[(1-methylethoxy)methyl]oxirane	43.5 ± 2.1			[1987VAN/KAC]
C ₆ H ₁₂ O ₂	[3126-95-2] $\Delta_v H$	(propoxymethyl)oxirane	48.5 ± 0.4			[1987VAN/KAC]
C ₆ H ₁₂ O ₂	[72380-56-4] $\Delta_v H$	1,1-dimethoxy-3-butene (305–334)	42.0	320	EB	[1994WIB/MOR]
C ₆ H ₁₂ O ₂	[123-86-4] $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$	butyl acetate (313–363) (313–353)	42.4 42.7 43.1 41.0 ± 0.5 43.0 ± 0.1 41.7 ± 0.1 40.6 ± 0.1 39.4 ± 0.1 41.3 43.6 ± 0.5 43.7 ± 0.2 40.5 43.6 ± 0.2 40.8	298 298 298 298 313 328 343 358 341 298 298 356 298 347	GC CGC CGC GC C C C C DTA GCC GCC A,EB C A	[1997KOU/HOS] [1995CHI/HOS] [1995CHI/HOS] [1987AZA] [1980SVO/UCH] [1980SVO/UCH] [1980SVO/UCH] [1980SVO/UCH] [1980MEY/AWE] [1980FUC/PEA] [1980FUC/PEA] [1987STE/MAL, 1969SHE/LAN] [1966WAD] [1987STE/MAL, 1964KLI/FRI, 1984BOU/FRI] [1961SCH/BOT]
C ₆ H ₁₂ O ₂	[123-42-2] $\Delta_v H$	diacetone alcohol (301–388)	47.5	316	A,I	[1987STE/MAL, 1952FUG/BOW]
C ₆ H ₁₂ O ₂	[123-42-2]	4-hydroxy-4-methyl-2-pentanone				

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound					
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference	
		$\Delta_v H$	52.1			[1975VIL/PER]	
		$\Delta_v H$	(295–441)	51.0	310		[1947STU]
C ₆ H ₁₂ O ₂	[105-54-4]	ethyl butyrate					
		$\Delta_v H$	(330–435)	39.4	345		[1997HER/ORT]
		$\Delta_v H$	(332–393)	40.2	347		[1993FAR/WIC]
		$\Delta_v H$	(310–336)	42.1 ± 0.1	323	EB	[1991WIB/WAL]
		$\Delta_v H$	(310–336)	43.7 ± 1.3	298	EB	[1991WIB/WAL]
		$\Delta_v H$	(263–404)	48.3	278	A	[1987STE/MAL]
		$\Delta_v H$		42.0 ± 0.1	298	C	[1966WAD]
		$\Delta_v H$	(254–394)	41.8	270		[1947STU]
C ₆ H ₁₂ O ₂	[97-62-1]	ethyl isobutyrate					
		$\Delta_v H$	(383–483)	36.0	398	A	[1987STE/MAL]
		$\Delta_v H$		39.8 ± 0.1	298	C	[1966WAD]
	$\Delta_v H$	(249–393)	44.1	264	A	[1987STE/MAL, 1947STU]	
C ₆ H ₁₂ O ₂	[110-19-0]	isobutyl acetate					
		$\Delta_v H$	(308–391)	39.9	323		[2005MON/MUN]
		$\Delta_v H$	(325–393)	39.2	340		[1996BUR/MON]
	$\Delta_v H$	(252–391)	39.8	267	A	[1987STE/MAL, 1947STU]	
C ₆ H ₁₂ O ₂	[540-88-5]	<i>tert</i> -butyl acetate					
		$\Delta_v H$	(308–372)	36.7	323		[2005MON/MUN]
	$\Delta_v H$		38.0 ± 0.2	298	C	[1966WAD, 1996VER/BEC]	
C ₆ H ₁₂ O ₂	[598-98-1]	methyl 2,2-dimethylpropanoate					
		$\Delta_v H$	(313–363)	37.7	298	CGC	[1995CHI/HOS]
		$\Delta_v H$		39.0 ± 0.5	298	GC	[1987AZA]
		$\Delta_v H$		38.8	298		[UR/FUC, 1985MAJ/SVO]
		$\Delta_v H$		39.7 ± 0.3	298	GCC	[1980FUC/PEA]
	$\Delta_v H$	(299–356)	35.2	319	BG	[1971HAL/BAL]	
C ₆ H ₁₂ O ₂	[110-45-2]	isopentyl formate					
		$\Delta_v H$	(255–397)	38.9	270	A	[1987STE/MAL]
C ₆ H ₁₂ O ₂	[556-24-1]	methyl isovalerate					
		$\Delta_v H$	(254–390)	41.2	269	A	[1987STE/MAL, 1947STU]
C ₆ H ₁₂ O ₂	[624-24-8]	methyl valerate					
		$\Delta_v H$	(275–311)	43.7 ± 0.3	298	GS	[2008VER/EME]
		$\Delta_v H$	(350–415)	44.1 ± 0.1		EB	[2007CAM/MOL]
		$\Delta_v H$	(281–547)	44.4	296		[2006CAM/MAR, 2006CAM]
		$\Delta_v H$	(364–417)	39.2	379		[2003ORT/ESP2]
		$\Delta_v H$		41.3	350		[2002VAN/VAN]
		$\Delta_v H$		43.7 ± 0.2	298		[2002VAN/VAN]
		$\Delta_v H$	(297–411)	42.5	312	A	[1987STE/MAL]
		$\Delta_v H$		43.3 ± 0.5	298	GC	[1987AZA]
		$\Delta_v H$		44.3 ± 0.5	298	C	[1981GAT/STR]
		$\Delta_v H$		46.1 ± 0.3	298	GCC	[1980FUC/PEA]
		$\Delta_v H$		44.1 ± 0.1	298	GCC	[1980FUC/PEA]
	$\Delta_v H$		43.1 ± 0.1	298	C	[1977MAN/SEL]	
C ₆ H ₁₂ O ₂	[106-36-5]	propyl propionate					
		$\Delta_v H$	(378–406)	37.6	392		[1994ORT/GAL]
		$\Delta_v H$	(336–394)	39.9	351		[1993FAR/WIC]
		$\Delta_v H$	(259–396)	43.1	274	A	[1987STE/MAL, 1947STU]
		$\Delta_v H$		42.1 ± 0.1	313	C	[1980SVO/UCH]
		$\Delta_v H$		41.1 ± 0.1	328	C	[1980SVO/UCH]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$		40.0 ± 0.1	343	C	[1980SVO/UCH]
	$\Delta_v H$		38.8 ± 0.1	358	C	[1980SVO/UCH]
C ₆ H ₁₂ O ₂	[142-62-1]	hexanoic acid				
	$\Delta_v H$	(297–328)	68.4 ± 0.9	313	GS	[2000VER2]
	$\Delta_v H$	(297–328)	69.2 ± 0.9	298	GS	[2000VER2]
	$\Delta_v H$	(353–393)	71.3	298	CGC	[1995CHI/HOS]
	$\Delta_v H$		70.9	271		[1982DEK/SCH]
	$\Delta_v H$	(270–280)	73.2 ± 2.0	298	TE	[1979DEK/OON]
	$\Delta_v H$	(335–487)	65.9	350	A	[1987STE/MAL, 1972DYK]
	$\Delta_v H$	(371–452)	66.6	386		[1957ROS/ACC, 1984BOU/FRI]
	$\Delta_v H$		64.6	367	I	[1943CRA]
C ₆ H ₁₂ O ₂	[88-09-5]	2-ethyl butyric acid				
	$\Delta_v H$	(373–466)	58.2	388	A	[1987STE/MAL]
C ₆ H ₁₂ O ₂	[646-07-1]	4-methylvaleric acid				
	$\Delta_v H$	(339–481)	91.7	354	A	[1987STE/MAL]
C ₆ H ₁₂ O ₂	[595-37-9]	2,2-dimethylbutanoic acid				
	$\Delta_v H$	(364–498)	59.4 ± 0.3	370	EB	[2002STE/CHI]
	$\Delta_v H$	(364–498)	54.6 ± 0.3	410	EB	[2002STE/CHI]
	$\Delta_v H$	(364–498)	50.0 ± 0.4	450	EB	[2002STE/CHI]
	$\Delta_v H$	(364–498)	46.0 ± 0.7	490	EB	[2002STE/CHI]
C ₆ H ₁₂ O ₂	[1070-83-3]	3,3-dimethylbutanoic acid				
	$\Delta_v H$	(283–325)	63.6 ± 0.9	304	GS	[2000VER2]
	$\Delta_v H$	(283–325)	64.0 ± 0.9	298	GS	[2000VER2]
C ₆ H ₁₂ O ₃	[na]	1-hexene ozonide				
	$\Delta_v H$	(353–373)	43.9	363	MM	[1977BOL/MAK]
C ₆ H ₁₂ O ₃	[37160-61-5]	sec-butyl glycolate				
	$\Delta_v H$	(301–451)	52.3	316	A	[1987STE/MAL, 1947STU]
C ₆ H ₁₂ O ₃	[123-63-7]	2,4,6-trimethyl-1,3,5-trioxane				
	$\Delta_{\text{trs}} H$		0.26	142.7		
	$\Delta_{\text{trs}} H$		0.77	147.5		
	$\Delta_{\text{fus}} H$		13.52	285.7		[1996DOM/HEA]
	$\Delta_v H$	(323–396)	41.5	338	A	[1987STE/MAL]
	$\Delta_v H$		41.4 ± 0.4			[1959FLE/MOR]
C ₆ H ₁₂ O ₃	[na]	glycerol 1-monoallyl ether				
	$\Delta_v H$	(323–383)	74.7	338	A	[1987STE/MAL, 1972DYK]
C ₆ H ₁₂ O ₃	[817-95-8]	2-ethoxyacetic acid, ethyl ester				
	$\Delta_v H$	(330–430)	46.1	345	A	[1987STE/MAL]
C ₆ H ₁₂ O ₃	[14144-33-3]	3-ethoxypropionic acid, methyl ester				
	$\Delta_v H$	(320–432)	44.3	335	A	[1987STE/MAL, 1972DYK]
C ₆ H ₁₂ O ₃	[111-15-9]	2-ethoxyethanol acetate				
	$\Delta_v H$	(322–430)	50.9	337	A	[1987STE/MAL]
	$\Delta_v H$		52.7 ± 0.1	298	C	[1970KUS/WAD]
	$\Delta_v H$	(330–468)	52.6 ± 0.4	298	EB	[1966BOT/ADL]
C ₆ H ₁₂ O ₃	[na]	3-hydroxypropionic acid, propyl ester				
	$\Delta_v H$	(350–375)	60.9	362	A	[1987STE/MAL]
C ₆ H ₁₂ O ₃	[10606-42-5]	3-methoxypropionic acid, ethyl ester				

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
	$\Delta_{\text{v}}H$	(313–432)	44.6	328	A	[1987STE/MAL]
C ₆ H ₁₂ O ₃	[616-09-1]	propyl lactate				
	$\Delta_{\text{v}}H$	(334–442)	52.1	349	A	[1987STE/MAL]
C ₆ H ₁₂ O ₃	[617-51-6]	isopropyl lactate				
	$\Delta_{\text{v}}H$	(356–430)	44.5	371		[2005PEN/MUR]
C ₆ H ₁₂ O ₃	[54078-53-4]	ethoxymethyl propionate				
	$\Delta_{\text{v}}H$		49.9 ± 0.1	298	C	[1974MAN]
C ₆ H ₁₂ O ₃	[5405-41-4]	ethyl 3-hydroxybutyrate				
	$\Delta_{\text{v}}H$	(363–393)	55.9 ± 0.6	298	CGC	[2005TEM/CHI]
C ₆ H ₁₂ O ₄	[624-47-5]	(<i>dl</i>) glycerol 1-propionate				
	$\Delta_{\text{v}}H$	(388–456)	75.8	403	A	[1987STE/MAL, 1972DYK]
C ₆ H ₁₂ O ₅	[na]	1-deoxy-(<i>d</i>)-glucopyranose				
	$\Delta_{\text{fus}}H$		27.4	403.2		[1996SCH]
C ₆ H ₁₂ O ₅	[na]	2-deoxy-(<i>d</i>)-glucopyranose				
	$\Delta_{\text{fus}}H$		34.5	398.7		[1996SCH]
C ₆ H ₁₂ O ₅	[na]	3-deoxy-(<i>d</i>)-glucopyranose				
	$\Delta_{\text{fus}}H$		32.6	387.2		[1996SCH]
C ₆ H ₁₂ O ₅	[na]	6-deoxy-(<i>d</i>)-glucopyranose				
	$\Delta_{\text{fus}}H$		22.7	409.2		[1996SCH]
C ₆ H ₁₂ O ₆	[87-89-8]	<i>myo</i> -inositol				
	$\Delta_{\text{fus}}H$		47.9	496.9		[1996DOM/HEA]
	$\Delta_{\text{sub}}H$	(438–458)	174.0 ± 2.6	448		[2006CHE/OJA]
	$\Delta_{\text{sub}}H$		181			[1999COS/EUS]
	$\Delta_{\text{sub}}H$		154.7 ± 1.4	477	TE	[1990BAR/DEL]
	$\Delta_{\text{sub}}H$		161	298		[1990BAR/DEL]
	$\Delta_{\text{sub}}H$		178	298	B	[1990BAR/DEL]
	$\Delta_{\text{sub}}H$	(454–472)	168			[1983DEW/BOW]
C ₆ H ₁₂ O ₆	$\Delta_{\text{v}}H$	(497–524)	119.0 ± 1.4	519	TE	[1990BAR/DEL]
	[na]	α -(<i>d</i>)-glucose				
	$\Delta_{\text{fus}}H$		34.3	423.2		[1996SCH]
C ₆ H ₁₂ O ₆	$\Delta_{\text{fus}}H$		31.42	414		[1996DOM/HEA]
	[na]	(<i>d</i>)-mannopyranose				
C ₆ H ₁₂ O ₆	$\Delta_{\text{fus}}H$		24.7	391.2		[1996SCH]
	[59-23-4]	(<i>d</i>)-galactose				
C ₆ H ₁₂ O ₆	$\Delta_{\text{fus}}H$		43.8	436.2		[2002JON/COO]
	[7133-36-0]	cyclopentyl methyl sulfide				
C ₆ H ₁₂ S	$\Delta_{\text{trs}}H$		0.9	165		
	$\Delta_{\text{fus}}H$		9.2	169.9		[1974MES/FIN]
	$\Delta_{\text{v}}H$		45.1 ± 0.1	298		[1972GOO, 1966OSB/DOU]
	$\Delta_{\text{v}}H$	(354–473)	41.7	369	A,EB	[1987STE/MAL, 1966OSB/DOU]
C ₆ H ₁₂ S	[5161-13-7]	<i>cis</i> 2,5-dimethyltetrahydrothiophene				
	$\Delta_{\text{v}}H$	(311–444)	41.7	326		[1999DYK/SVO]
	$\Delta_{\text{v}}H$	(349–427)	39.7	364	A,EB	[1987STE/MAL, 1952WHI/BER]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound					
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference	
C₆H₁₂S	[5161-14-8] $\Delta_{\text{v}}H$	<i>trans</i> 2,5-dimethyltetrahydrothiophene (348–396)	39.3	363	EB	[1987STE/MAL, 1952WHI/BER, 1999DYK/SVO]	
C₆H₁₂S	[1551-32-2] $\Delta_{\text{v}}H$	2-ethyltetrahydrothiophene (333–488)	42.6	348	A	[1987STE/MAL, 1972DYK, 1999DYK/SVO]	
C₆H₁₂S	[62184-67-2] $\Delta_{\text{v}}H$	3-ethyltetrahydrothiophene (343–503)	43.1	358	A	[1987STE/MAL, 1972DYK, 1999DYK/SVO]	
C₆H₁₂S	[5161-16-0] $\Delta_{\text{v}}H$ $\Delta_{\text{v}}H$	2-methyltetrahydro-2 <i>H</i> -thiopyrane (317–455) (356–438)	42.1	332	A,EB	[1999DYK/SVO]	
			40.2	371		[1987STE/MAL, 1952WHI/BER]	
C₆H₁₂S	[5258-50-4] $\Delta_{\text{v}}H$ $\Delta_{\text{v}}H$	3-methyltetrahydro-2 <i>H</i> -thiopyrane (321–460) (361–435)	42.5	336	A,EB	[1999DYK/SVO]	
			40.7	376		[1987STE/MAL, 1952WHI/BER]	
C₆H₁₂S	[5161-17-1] $\Delta_{\text{v}}H$ $\Delta_{\text{v}}H$	4-methyltetrahydro-2 <i>H</i> -thiopyrane (321–461) (361–441)	42.8	336	A,EB	[1999DYK/SVO]	
			40.8	376		[1987STE/MAL, 1952WHI/BER]	
C₆H₁₂S	[1569-69-3] $\Delta_{\text{fus}}H$ $\Delta_{\text{v}}H$ $\Delta_{\text{v}}H$ $\Delta_{\text{v}}H$	cyclohexanethiol	10.0	189.6	A,EB	[1996DOM/HEA]	
			44.9	298		C	[1981HOS/SCO]
			44.6 ± 0.1	298			[1972GOO, 1966OSB/DOU]
			41.2	370			[1987STE/MAL, 1966OSB/DOU, 1999DYK/SVO]
C₆H₁₂S₃	[6573-11-1] $\Delta_{\text{fus}}H$	1,4,7-trithiacyclononane	29.0	354.2	DSC	[2002ROC/GRI]	
C₆H₁₃Br	[111-25-1] $\Delta_{\text{fus}}H$ $\Delta_{\text{v}}H$ $\Delta_{\text{v}}H$ $\Delta_{\text{v}}H$	1-bromohexane (323–363) (333–456)	18.05	188.1	A,EST	[1996DOM/HEA]	
			45.5	298		CGC	[1995CHI/HOS]
			46.1 ± 0.1	298		C	[1968WAD]
			45.6 ± 0.1	298		C	[1966WAD]
			43.2	348			[1987STE/MAL, 1961LI/ROS, 1972DYK]
C₆H₁₃Br	[3377-86-4] $\Delta_{\text{v}}H$	(<i>dl</i>) 2-bromohexane (303–416)	43.8	318	A	[1987STE/MAL]	
C₆H₁₃Br	[30310-22-6] $\Delta_{\text{v}}H$	2-bromo-4-methylpentane (315–448)	29.3	330	A	[1987STE/MAL, 1972DYK]	
C₆H₁₃Br	[26356-06-9] $\Delta_{\text{v}}H$	2-bromo-3,3-dimethylbutane (315–449)	39.5	330	A	[1987STE/MAL, 1972DYK]	
C₆H₁₃Cl	[544-10-5] $\Delta_{\text{v}}H$ $\Delta_{\text{v}}H$ $\Delta_{\text{v}}H$ $\Delta_{\text{v}}H$ $\Delta_{\text{v}}H$ $\Delta_{\text{v}}H$	1-chlorohexane (319–376) (290–410)	41.1	334	C	[1988PAU/KRU]	
			42.0	298		[1984BOU/FRI, 1991BAS/SVO]	
			42.8 ± 0.1	298		[1981TEK/MAJ]	
			40.5 ± 0.1	328		[1981TEK/MAJ]	
			40.0 ± 0.1	343		[1981TEK/MAJ]	
			39.0 ± 0.1	358		[1981TEK/MAJ]	
			38.4 ± 0.1	368		[1981TEK/MAJ]	

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		T _m (K)	Method	Reference
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)			
		$\Delta_v H$	(288–409)	43.5	303	A,DTA [1987STE/MAL, 1969KEM/KRE, 1972DYK]
		$\Delta_v H$		42.8 ± 0.1	298	C [1968WAD]
C ₆ H ₁₃ Cl	[638-28-8]		(dl) 2-chlorohexane			
		$\Delta_v H$	(300–399)	40.9	315	A [1987STE/MAL]
C ₆ H ₁₃ Cl	[594-57-0]		2-chloro-2,3-dimethylbutane			
		$\Delta_v H$	(301–426)	38.0	316	A [1987STE/MAL, 1972DYK]
C ₆ H ₁₃ Cl	[5750-00-5]		(dl) 2-chloro-3,3-dimethylbutane			
		$\Delta_v H$	(300–425)	38.0	315	A [1987STE/MAL, 1972DYK]
C ₆ H ₁₃ ClO ₂ S	[14532-24-2]		1-hexanesulfonyl chloride			
		$\Delta_v H$	(273–304)	60.7	288	[1999DYK/SVO, 1963QUI/NOW]
		$\Delta_v H$	(303–400)	61.7	318	[1999DYK/SVO]
		$\Delta_v H$	(400–507)	57.2	415	[1999DYK/SVO]
C ₆ H ₁₃ Cl ₂ N	[13426-57-8]		N-ethyl-bis(2-chloroethyl)amine			
		$\Delta_v H$	(273–333)	54.9	288	A,GS [1987STE/MAL, 1948RED/CHA3, 1972DYK]
C ₆ H ₁₃ F	[373-14-8]		1-fluorohexane			
		$\Delta_v H$	(273–388)	36.9	288	A,EST [1987STE/MAL, 1961LI/ROS, 1972DYK]
C ₆ H ₁₃ F	[52688-75-2]		3-fluorohexane			
		$\Delta_v H$	(281–393)	36.8	296	A [1987STE/MAL, 1972DYK]
C ₆ H ₁₃ I	[638-45-9]		1-iodohexane			
		$\Delta_v H$		49.8 ± 0.1	298	C [1968WAD]
		$\Delta_v H$	(331–485)	46.2	346	A,EST [1987STE/MAL, 1961LI/ROS, 1972DYK]
C ₆ H ₁₃ N	[108-91-8]		cyclohexylamine			
		$\Delta_{\text{trs}}H$		1.0	258.2	
		$\Delta_{\text{fus}}H$		16.5	255.1	[1999HAM/WUR]
		$\Delta_{\text{fus}}H$		14.92	255.4	[1939VAN, 1999KAB/KOZ]
		$\Delta_v H$	(363–407)	40.6	378	[1987STE/MAL]
		$\Delta_v H$		42.7 ± 0.1	313	C [1979MAJ/SVO2]
		$\Delta_v H$		40.7 ± 0.1	343	C [1979MAJ/SVO2]
		$\Delta_v H$		39.6 ± 0.1	358	C [1979MAJ/SVO2]
		$\Delta_v H$		42.8 ± 0.1	298	C [1975BER/OLO]
		$\Delta_v H$	(333–408)	40.8	348	A [1987STE/MAL, 1972DYK]
		$\Delta_v H$	(334–401)	40.8	349	[1960NOV/MAT2, 1984BOU/FRI, 1960NOV/MAT]
C ₆ H ₁₃ N	[111-49-9]		hexahydro-1H-azepine			
		$\Delta_v H$	(348–423)	37.7	363	A [1987STE/MAL]
		$\Delta_v H$	(312–411)	40.4	327	A [1987STE/MAL, 1972DYK]
C ₆ H ₁₃ N	[626-67-5]		N-methylpiperidine			
		$\Delta_v H$		36.8 ± 0.6	298	C [2006RIB/CAB5]
C ₆ H ₁₃ N	[109-05-7]		(dl) 2-methylpiperidine			
		$\Delta_{\text{fus}}H$		18.58	269.4	[1996DOM/HEA]
		$\Delta_v H$	(323–431)	38.2	338	EB,IP [1987STE/MAL, 1968OSB/DOU]
C ₆ H ₁₃ N	[626-56-2]		3-methylpiperidine			
		$\Delta_v H$		44.4 ± 0.7	298	C [2006RIB/CAB5]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound							
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference			
C ₆ H ₁₃ N	[626-58-4] $\Delta_v H$	4-methylpiperidine	40.6 ± 0.9	298	C	[2006RIB/CAB5]			
C ₆ H ₁₃ N	[626-67-5] $\Delta_v H$	N-methylpiperidine	36.8 ± 0.6	298	C	[2006RIB/CAB5]			
	$\Delta_v H$		(298–343)	36.5			313	[1995BEL/AIT]	
	$\Delta_v H$		(273–380)	37.3	288	A	[1987STE/MAL]		
	$\Delta_v H$			36.7 ± 0.1	298		[1979BER/ANG, 1998EWI/SAN]		
C ₆ H ₁₃ NO	[100-74-3] $\Delta_v H$	N-ethylmorpholine	(274–313)	42.3 ± 0.3	294	GS	[1998VER2]		
	$\Delta_v H$		(274–313)	42.1 ± 0.3	298	GS	[1998VER2]		
C ₆ H ₁₃ NO	[na] $\Delta_v H$	methyl 2-(N,N-dimethylamino)propanoate	(278–306)	46.1 ± 1.1	290	GS	[1992VER/BEC]		
C ₆ H ₁₃ NO	[na] $\Delta_v H$	ethyl 2-(N,N-dimethylamino)ethanoate	(278–308)	47.6 ± 0.8	293	GS	[1992VER/BEC]		
C ₆ H ₁₃ NO	[127-19-5] $\Delta_v H$	N,N-diethylacetamide	(463–513)	53.7 ± 0.4	298	CGC	[2009PAN/ANT]		
	$\Delta_v H$			54.1	298	A	[1985BAR/CAS, 1985MAJ/SVO]		
C ₆ H ₁₃ NO	[1119-49-9] $\Delta_v H$	N-butylacetamide		75.0 ± 0.3	298	C	[1984STA/WAD]		
C ₆ H ₁₃ NO	[760-79-2] $\Delta_v H$	N,N-dimethyl butyramide	(251–432)	50.8	366	A	[1987STE/MAL]		
	$\Delta_v H$			55.2			[1977VAS/KOT]		
C ₆ H ₁₃ NO	[762-84-5] $\Delta_{\text{sub}} H$	<i>tert</i> -butylacetamide	(278–295)	78.3 ± 0.3	287	ME	[1983ZIE/ZIE]		
	$\Delta_{\text{sub}} H$			77.9 ± 0.4	298		[1983ZIE/ZIE]		
C ₆ H ₁₃ NO	[628-02-4] $\Delta_{\text{trs}} H$	hexanamide		7.9	305.1	DSC	[2008ABA/BAD] [1973LEB/KAT2]		
	$\Delta_{\text{fus}} H$			16.7	373				
	$\Delta_{\text{fus}} H$			25.1	374				
	$\Delta_{\text{sub}} H$		(301–371)	85 ± 4.0	298			TE	[2000BRU/DEL]
	$\Delta_{\text{sub}} H$		(293–303)	98.7 ± 1.7	298				[1973LEB/KAT2, 1977PED/RYL]
	$\Delta_{\text{sub}} H$		(338–368)	95.1 ± 4	353			GS	[1959DAV/JON2, 1970COX/PIL, 1987STE/MAL]
C ₆ H ₁₃ NO	[3554-74-3] $\Delta_v H$	1-methyl-3-piperidinol		73.0 ± 0.6	298	C	[2006RIB/CAB2]		
C ₆ H ₁₃ NO	[106-52-5] $\Delta_v H$	1-methyl-4-piperidinol		80.8 ± 0.4	298	C	[2006RIB/CAB2]		
C ₆ H ₁₃ NO	[3433-37-2] $\Delta_{\text{sub}} H$	2-piperidinemethanol		93.0 ± 0.5	298	C	[2006RIB/CAB3]		
C ₆ H ₁₃ NO	[4606-65-9] $\Delta_{\text{sub}} H$	3-piperidinemethanol		95.9 ± 1.4	298	C	[2006RIB/CAB3]		
C ₆ H ₁₃ NO	[6457-49-4] $\Delta_{\text{sub}} H$	4-piperidinemethanol		98.3 ± 0.7	298	C	[2006RIB/CAB3]		
C ₆ H ₁₃ NO ₂	[na] $\Delta_v H$	N-isopropyl lactamide	(369–407)	69.9	384	A	[1987STE/MAL]		

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₆ H ₁₃ NO ₂	[na]	N-propyl lactamide				
	$\Delta_v H$	(373–423)	74.0	388	A	[1987STE/MAL]
C ₆ H ₁₃ NO ₂	[616-06-8]	<i>dl</i> -2-aminohexanoic acid				
	$\Delta_{\text{sub}} H$	(435–469)	114.5 ± 0.4	450	ME	[1965SVE/CLY, 1964CLY/SVE, 1987STE/MAL]
C ₆ H ₁₃ NO ₂	[3107-04-8]	2-amino-3-methylpentanoic acid				
	$\Delta_{\text{sub}} H$		120.1 ± 0.8	455	ME	[1965SVE/CLY, 1964CLY/SVE]
C ₆ H ₁₃ NO ₂	[328-38-1]	<i>L</i> -(<i>d</i>)-2-amino-4-methylpentanoic acid (<i>L</i> -(<i>d</i>)-leucine)				
	$\Delta_{\text{sub}} H$	(323–423)	U 83.7 ± 4	373	LE	[1977GAF/PIE]
C ₆ H ₁₃ NO ₂	[61-90-5]	<i>D</i> -(<i>l</i>)-leucine				
	$\Delta_{\text{sub}} H$	(401–517)	148.7 ± 6.5		TGA	[2009LAH/RAU]
	$\Delta_{\text{sub}} H$	(446–464)	150.6 ± 0.8	455	ME	[1965SVE/CLY, 1970COX/PIL, 1964CLY/SVE]
C ₆ H ₁₃ NO ₂	[60-32-2]	6-aminohexanoic acid				
	$\Delta_{\text{sub}} H$	(388–407)	153.3 ± 0.8	398	C	[1983SKO/SAB]
	$\Delta_{\text{sub}} H$		155 ± 3	298	C	[1983SKO/SAB]
C ₆ H ₁₄	[110-54-3]	hexane				
	$\Delta_{\text{fus}} H$		13.08	177.8		[1996DOM/HEA]
	$\Delta_{\text{sub}} H$		50.8	178	B	[1963BON]
	$\Delta_v H$		31.4 ± 0.2	298	C	[2007PAS/KUZ]
	$\Delta_v H$		31.5 ± 0.1	298	C	[1996VAR/PAS]
	$\Delta_v H$		31.5	298		[1994RUZ/MAJ]
	$\Delta_v H$	(283–323)	32.1	298		[1992GRA/SAN]
	$\Delta_v H$		31.3 ± 0.3		GC	[1989AZA]
	$\Delta_v H$	(238–298)	34.9	253	A	[1987STE/MAL]
	$\Delta_v H$	(189–259)	35.7	244	A	[1987STE/MAL]
	$\Delta_v H$	(298–343)	31.5	313	A	[1987STE/MAL]
	$\Delta_v H$	(341–377)	30.1	356	A	[1987STE/MAL]
	$\Delta_v H$	(374–451)	29.3	389	A	[1987STE/MAL]
	$\Delta_v H$	(445–508)	29.4	460	A	[1987STE/MAL]
	$\Delta_v H$		26.6	373	C	[1985WOR/YER]
	$\Delta_v H$		22.5	423	C	[1985WOR/YER]
	$\Delta_v H$		15.7	473	C	[1985WOR/YER]
	$\Delta_v H$		8.9	498	C	[1985WOR/YER]
	$\Delta_v H$		31.6	298		[UR/FUC, 1985MAJ/SVO]
	$\Delta_v H$	(298–338)	30.9	313		[1984MIC/JOS]
	$\Delta_v H$		31.6 ± 0.1	298	C	[1979MAJ/SVO]
	$\Delta_v H$		30.7 ± 0.1	313	C	[1979MAJ/SVO]
	$\Delta_v H$		29.5 ± 0.1	333	C	[1979MAJ/SVO]
	$\Delta_v H$		28.2 ± 0.1	353	C	[1979MAJ/SVO]
	$\Delta_v H$	(300–321)	31.6	310		[1974LET/MAR, 1984BOU/FRI]
	$\Delta_v H$	(178–265)	32.5	250		[1973CAR/KOB]
	$\Delta_v H$		31.55	298		[1971WIL/ZWO]
	$\Delta_v H$		30.9 ± 0.1	309	C	[1947WAD/DOU]
	$\Delta_v H$		29.8 ± 0.1	328	C	[1947WAD/DOU]
	$\Delta_v H$		31.5 ± 0.1	298	C	[1947OSB/GIN]
	$\Delta_v H$	(286–343)	32.0	301	A,MM	[1987STE/MAL, 1945WIL/TAY]
	$\Delta_v H$		31.0 ± 0.2	298	C	[1943LEM/FEL]
	$\Delta_v H$		30.5 ± 0.2	313	C	[1943LEM/FEL]
	$\Delta_v H$		29.0 ± 0.2	333	C	[1943LEM/FEL]
	$\Delta_v H$		28.2 ± 0.2	353	C	[1943LEM/FEL]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₆ H ₁₄	[107-83-5]	2-methylpentane				
	$\Delta_{\text{fus}}H$		6.27	119.6		[1996DOM/HEA]
	Δ_vH	(290–333)	30.2	305		[2010SAP/UUS]
	Δ_vH	(301–333)	30.0	316		[2002POK/UUS]
	Δ_vH	(310–359)	29.7	325		[1998AUC/LOR]
	Δ_vH	(293–335)	30.5	308	A	[1987STE/MAL]
	Δ_vH		29.9	298		[1971WIL/ZWO]
	Δ_vH		29.9 ± 0.1	298	C	[1949WAD/SMI]
	Δ_vH		28.7 ± 0.1	318	C	[1949WAD/SMI]
	Δ_vH		27.8 ± 0.1	333	C	[1949WAD/SMI]
	Δ_vH		29.9 ± 0.1	298	C	[1947OSB/GIN]
	Δ_vH	(286–334)	30.4	301	MM	[1945WIL/TAY]
	Δ_vH		29.8 ± 0.2	293	C	[1943LEM/FEL]
	Δ_vH		29.0 ± 0.2	313	C	[1943LEM/FEL]
Δ_vH		27.6 ± 0.2	333	C	[1943LEM/FEL]	
Δ_vH		26.9 ± 0.2	353	C	[1943LEM/FEL]	
C ₆ H ₁₄	[96-14-0]	3-methylpentane				
	$\Delta_{\text{fus}}H$		5.31	110.3		[1996DOM/HEA]
	Δ_vH	(316–361)	29.9	331		[1999LOR/AUC]
	Δ_vH	(293–338)	30.5	308	A	[1987STE/MAL]
	Δ_vH		30.3 ± 0.1	298	C	[1979MAJ/SVO]
	Δ_vH		29.5 ± 0.1	313	C	[1979MAJ/SVO]
	Δ_vH		28.3 ± 0.1	333	C	[1979MAJ/SVO]
	Δ_vH		27.0 ± 0.1	353	C	[1979MAJ/SVO]
	Δ_vH		30.3	298		[1971WIL/ZWO]
	Δ_vH		30.0 ± 0.1	303	C	[1949WAD/SMI]
	Δ_vH		28.8 ± 0.1	324	C	[1949WAD/SMI]
	Δ_vH		28.1 ± 0.1	336	C	[1949WAD/SMI]
	Δ_vH		30.3 ± 0.1	298	C	[1947OSB/GIN]
	Δ_vH	(288–337)	30.2	303	MM	[1945WIL/TAY]
C ₆ H ₁₄	[79-29-8]	2,3-dimethylbutane				
	$\Delta_{\text{trs}}H$		6.43	136.1		
	$\Delta_{\text{trs}}H$		2.37	107		
	$\Delta_{\text{fus}}H$		0.79	145.2		[1996DOM/HEA]
	Δ_vH		29.1	298		[1971WIL/ZWO]
	Δ_vH		29.2 ± 0.1	296	C	[1949WAD/SMI]
	Δ_vH		28.9 ± 0.1	303	C	[1949WAD/SMI]
	Δ_vH		28.3 ± 0.1	313	C	[1949WAD/SMI]
	Δ_vH		27.3 ± 0.1	331	C	[1949WAD/SMI]
	Δ_vH	(287–332)	29.6	302	MM	[1945WIL/TAY]
	Δ_vH		29.2 ± 0.1	293	C	[1943LEM/FEL]
	Δ_vH		28.2 ± 0.1	313	C	[1943LEM/FEL]
	Δ_vH		27.0 ± 0.1	333	C	[1943LEM/FEL]
	Δ_vH		26.1 ± 0.1	353	C	[1943LEM/FEL]
C ₆ H ₁₄	[75-83-2]	2,2-dimethylbutane				
	$\Delta_{\text{trs}}H$		5.4	126.8		
	$\Delta_{\text{trs}}H$		0.28	140.8		
	$\Delta_{\text{fus}}H$		0.58	174.3		[1996DOM/HEA]
	Δ_vH		27.7	298		[1971WIL/ZWO]
	Δ_vH	(273–318)	28.7	288		[1949NIC/LAF, 1984BOU/FRI]
	Δ_vH		27.8 ± 0.1	296	C	[1947WAD/DOU]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		T _m (K)	Method	Reference	
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)				
		$\Delta_v H$	26.3 ± 0.1	323	C	[1947WAD/DOU]	
		$\Delta_v H$	(211–289)	29.2		[1946KIL/PIT]	
		$\Delta_v H$	(288–323)	28.3	MM	[1945WIL/TAY]	
C₆H₁₄FO₃P	[55-91-4]	fluorophosphoric acid, diisopropyl ester					
		$\Delta_v H$	(273–348)	29.4	A	[1987STE/MAL]	
C₆H₁₄N₂	[821-67-0]	dipropyldiazene					
		$\Delta_v H$	39.9 ± 0.4	298	C	[1976ENG/MEL]	
		$\Delta_v H$	(295–305)	39.5	UV	[1974ENG/WOO]	
		$\Delta_v H$	41.1			[1968BAC/NOV, 1974ENG/WOO]	
C₆H₁₄N₂	[3880-49-7]	diisopropyldiazene					
		$\Delta_v H$	35.9 ± 0.4	298	C	[1976ENG/MEL]	
		$\Delta_v H$	(296–308)	36.1	UV	[1974ENG/WOO]	
		$\Delta_v H$	37.7		I	[1974ENG/WOO]	
		$\Delta_v H$	34.9			[1968GEI/HOF, 1974ENG/WOO]	
C₆H₁₄N₂	[3114-70-3]	1,4-diaminocyclohexane					
		$\Delta_v H$	(383–473)	48.2	A	[1987STE/MAL]	
C₆H₁₄N₂	[1436-59-5]	<i>cis</i> 1,2-cyclohexanediamine					
		$\Delta_v H$	62.2 ± 1.0	298	ME	[2007TOM/ROS]	
C₆H₁₄N₂	[2615-25-0]	<i>trans</i> -cyclohexyl-1,4-diamine					
		$\Delta_{\text{fus}} H$	27.0	342.1	DSC	[2007TOM/ROS]	
		$\Delta_{\text{sub}} H$	105.0 ± 0.8	298	C	[2007TOM/ROS]	
C₆H₁₄N₂	[106-58-1]	1,4-dimethylpiperazine					
		$\Delta_v H$	(268–311)	41.2 ± 0.4	298	GS	[2010EFI/EME]
		$\Delta_v H$	(270–309)	44.3 ± 0.3	289	GS	[1998VER2]
		$\Delta_v H$	(270–309)	43.8 ± 0.3	298	GS	[1998VER2]
		$\Delta_v H$	(276–319)	41.6	291	A	[1987STE/MAL, 1975CAB/CON]
C₆H₁₄N₂	[106-55-8]	2,5-dimethylpiperazine					
		$\Delta_v H$	(437–609)	48.4	452	A	[1987STE/MAL]
C₆H₁₄N₂	[7423-00-9]	propylhydrazone acetone					
		$\Delta_v H$	(288–318)	44.0	300		[1980LEB/NAZ]
C₆H₁₄N₂	[7423-01-0]	isopropylhydrazone acetone					
		$\Delta_v H$	(288–323)	44.6	303		[1980LEB/NAZ]
C₆H₁₄N₂O	[103-76-4]	N-(hydroxyethyl)piperazine					
		$\Delta_v H$	(308–343)	77.3 ± 0.7	326	GS	[2002VER2]
		$\Delta_v H$	(308–343)	78.8 ± 0.7	298	GS	[2002VER2]
C₆H₁₄N₂O	[17697-55-1]	dipropyldiazene N-oxide					
		$\Delta_v H$	51.7 ± 0.1	298	C	[1981BYS]	
C₆H₁₄N₂O	[38869-91-9]	1-pentyl urea					
		$\Delta_{\text{trs}} H$	2.5	355.1			
		$\Delta_{\text{fus}} H$	21.0	375.2	DSC	[2005HAS/TAJ]	
C₆H₁₄N₂O₂	[56-87-1]	<i>(l)</i> -lysine					
		$\Delta_{\text{sub}} H$	(397–497)	U 88 ± 8	447	LE	[1977GAF/PIE]
C₆H₁₄N₄O₂	[74-79-3]	<i>(l)</i> -arginine					
		$\Delta_{\text{sub}} H$	(441–541)	U 134 ± 8	491	LE	[1977GAF/PIE]
C₆H₁₄N₈O₈	[13405-40-8]	2,4,7,9-tetranitro-2,4,7,9-tetraazadecane					

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound					
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference	
		$\Delta_{\text{fus}}H$	68.09	488.1	DSC	[1997ZEM]	
C ₆ H ₁₄ O	[62881-9]	butyl ethyl ether					
	Δ_vH		36.3 ± 0.1	298	C	[1980MAJ/WAN]	
	Δ_vH	(311–365)	36.5	298		[1976AMB/ELL]	
	Δ_vH	(311–365)	32.1	365		[1976AMB/ELL]	
		Δ_vH	(311–365)	35.2	326	A,EB	[1987STE/MAL, 1976AMB/ELL, 1969CID/POL, 1972DYK]
C ₆ H ₁₄ O	[994-05-8]	<i>tert</i> -amyl methyl ether					
	Δ_vH		35.3 ± 0.4	298		[UR/VER, 2002VER, 2003VER/KRA]	
	Δ_vH	(309–396)	36.6 ± 0.1	320	EB	[2002STE/CHI2]	
	Δ_vH	(309–396)	34.5 ± 0.2	360	EB	[2002STE/CHI2]	
	Δ_vH	(309–396)	32.1 ± 0.5	400	EB	[2002STE/CHI2]	
	Δ_vH		34.8	298	EB	[1999HEI/FIS, 2003VER/KRA]	
	Δ_vH	(314–362)	33.4	329		[1998AUC/LOR]	
	Δ_vH	(283–308)	35.7 ± 1.0	295	GS	[1998VER/WEL]	
	Δ_vH	(283–308)	35.5 ± 1.0	298	GS	[1998VER/WEL]	
	Δ_vH		35.2	298	EB	[1996TOG/TOG, 2003VER/KRA]	
	Δ_vH		35.0	298	EB	[1994ANT/SAN, 2003VER/KRA]	
	Δ_vH	(306–359)	33.8	321	EB	[1994KRA/GME]	
	Δ_vH		35.8	298	C	[1991ROZ/SAF]	
	Δ_vH	(294–359)	33.5	298	EB	[1991ROZ/SAF, 1984CER/BOU, 2003VER/KRA]	
		Δ_vH	(294–359)	34.3	309	EB	[1984CER/BOU]
		Δ_vH	(309–358)	33.7	324	EB	[1984PAL/CHO]
C ₆ H ₁₄ O	[637-92-3]	<i>tert</i> -butyl ethyl ether					
	Δ_vH	(303–345)	33.1 ± 0.4	298	EB	[2007EFI/PAS]	
	Δ_vH	(313–346)	32.1	328		[2007SAP/ZAY]	
	Δ_vH	(313–345)	31.9	328		[2004KIM/KES]	
	Δ_vH	(307–346)	32.1	322		[2000REI/CAR]	
	Δ_vH	(306–345)	32.2	321	EB	[1994KRA/GME]	
	Δ_vH	(284–346)	33.5	299	A	[1987STE/MAL]	
	Δ_vH	(248–350)	35.3	263	A	[1987STE/MAL]	
		Δ_vH	(340–407)	31.2	355	A	[1987STE/MAL]
C ₆ H ₁₄ O	[111-43-3]	dipropyl ether					
	$\Delta_{\text{trs}}H$		2.3	149.4			
	$\Delta_{\text{fus}}H$		10.77	158.4		[1996DOM/HEA]	
	Δ_vH	(308–338)	34.8	323	EB	[2002ANT/FRA]	
	Δ_vH	(385–467)	32.2	400	A	[1987STE/MAL]	
	Δ_vH	(465–530)	32.4	480	A	[1987STE/MAL]	
	Δ_vH		35.7 ± 0.1	298	C	[1980MAJ/WAN]	
	Δ_vH	(292–389)	35.6	307	A	[1987STE/MAL, 1976AMB/ELL]	
	Δ_vH		31.4	363		[1976AMB/ELL]	
	Δ_vH	(312–371)	34.6	327	A,EB	[1987STE/MAL, 1973MEY/HOT]	
		Δ_vH	(300–362)	35.1	315	EB	[1969CID/POL]
		Δ_vH	(340–379)	34.5	360		[1968LAP/NIS]
C ₆ H ₁₄ O	[108-20-3]	diisopropyl ether					
	$\Delta_{\text{fus}}H$		12.05	187.8		[1974AND/COU]	
	Δ_vH	(285–365)	32.7 ± 0.5	298	EB	[2007EFI/PAS]	
	Δ_vH	(278–323)	33.0	293		[1999GAR/AND]	
	Δ_vH	(307–349)	31.1	322		[1999MON/DEL]	
		Δ_vH	(360–440)	29.9	375	A	[1987STE/MAL]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(436–500)	29.5	451	A	[1987STE/MAL]
	$\Delta_v H$		32.1 ± 0.1	298	C	[1980MAJ/WAN]
	$\Delta_v H$	(284–365)	32.6	299	A	[1987STE/MAL, 1976AMB/ELL]
	$\Delta_v H$	(284–365)	29.2	341		[1976AMB/ELL]
	$\Delta_v H$	(296–342)	32.1	311	A,EB	[1987STE/MAL, 1969CID/POL]
	$\Delta_v H$	(321–350)	30.1	336		[1965NIS/LAP2, 1972DYK]
	$\Delta_v H$	(273–333)	33.2	288		[1949NIC/LAF, 1984BOU/FRI]
C ₆ H ₁₄ O	[111-27-3]	1-hexanol				
	$\Delta_{\text{fus}} H$		15.48	225.8		[1996DOM/HEA]
	$\Delta_v H$	(344–384)	59.7	359	EB	[2009GIE/KOS]
	$\Delta_v H$	(328–423)	59.9	298		[2006NAS/NEU]
	$\Delta_v H$	(265–363)	61.7 ± 0.3	298	GS	[2005ROG/PIS]
	$\Delta_v H$	(370–416)	51.4	385	EB	[2004TAN/LI]
	$\Delta_v H$	(265–328)	62.0	288	GS	[2001KUL/VER2]
	$\Delta_v H$	(265–328)	61.1	298	GS	[2001KUL/VER2]
	$\Delta_v H$	(268–333)	61.9 ± 0.2	301	GS	[1998VER5]
	$\Delta_v H$	(268–333)	62.1 ± 0.2	298	GS	[1998VER5]
	$\Delta_v H$	(373–423)	61.5	298	CGC	[1995CHI/HOS]
	$\Delta_v H$	(323–373)	61.6	298	CGC	[1995CHI/HOS]
	$\Delta_v H$	(253–338)	61.2	296		[1992NGU/KAS]
	$\Delta_v H$	(298–343)	57.7	313	A	[1987STE/MAL]
	$\Delta_v H$	(380–417)	47.9	395	EB	[1985RED/RAO]
	$\Delta_v H$		58.5 ± 0.2	328	C	[1985MAJ/SVO3]
	$\Delta_v H$		57.6 ± 0.2	343	C	[1985MAJ/SVO3]
	$\Delta_v H$		55.2 ± 0.2	358	C	[1985MAJ/SVO3]
	$\Delta_v H$		53.8 ± 0.2	368	C	[1985MAJ/SVO3]
	$\Delta_v H$	(243–303)	59.1	298		[1983SCH/STR]
	$\Delta_v H$		60.8 ± 0.2	298	C	[1977MAN/SEL]
	$\Delta_v H$	(308–430)	57.9	323		[1973WIL/ZWO]
	$\Delta_v H$	(325–431)	58.5	340	DTA	[1987STE/MAL, 1969KEM/KRE]
$\Delta_v H$					[1972DYK]	
$\Delta_v H$		61.6 ± 0.2	298	C	[1966WAD]	
$\Delta_v H$	(334–381)	56.0	349		[1961ROS/SUP]	
$\Delta_v H$	(308–428)	U 55.8	323	I	[1938HOV/LAN]	
$\Delta_v H$	(333–425)	57.9	348		[1935BUT/RAM, 1984BOU/FRI]	
C ₆ H ₁₄ O	[626-93-7]	(dl) 2-hexanol				
	$\Delta_v H$	(274–309)	57.0 ± 0.2	298	GS	[2005ROG/PIS]
	$\Delta_v H$	(274–309)	58.3 ± 0.3	298	GS	[2001KUL/VER]
	$\Delta_v H$	(224–323)	61.8	239		[1999NGU/BER]
	$\Delta_v H$	(360–415)	48.7	375	A	[1987STE/MAL]
	$\Delta_v H$		56.8 ± 0.2	313	C	[1985MAJ/SVO2]
	$\Delta_v H$		55.0 ± 0.2	328	C	[1985MAJ/SVO2]
	$\Delta_v H$		53.0 ± 0.2	343	C	[1985MAJ/SVO2]
	$\Delta_v H$		50.7 ± 0.2	358	C	[1985MAJ/SVO2]
	$\Delta_v H$		49.2 ± 0.2	368	C	[1985MAJ/SVO2]
	$\Delta_v H$	(337–413)	52.4	352		[1984SAC/MAR]
	$\Delta_v H$	(351–412)	47.8	366	A	[1987STE/MAL, 1975BRA/AND]
	$\Delta_v H$	(301–415)	53.1	316		[1973WIL/ZWO]
	$\Delta_v H$	(298–413)	49.7	356	I	[1938HOV/LAN]
C ₆ H ₁₄ O	[623-37-0]	(dl) 3-hexanol				
	$\Delta_v H$	(278–311)	58.6 ± 0.4	298	GS	[2001KUL/VER]
	$\Delta_v H$	(244–318)	U 50.7	259		[1999NGU/BER]
$\Delta_v H$	(354–410)	46.1	369	A	[1987STE/MAL]	

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound					
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference	
		$\Delta_v H$	(280–320)	57.5	295	A	[1987STE/MAL]
		$\Delta_v H$	(333–409)	51.5	348		[1984SAC/MAR]
		$\Delta_v H$	(280–316)	57.4	295		[1975CAB/CON2]
		$\Delta_v H$	(298–408)	46.4	353	I	[1938HOV/LAN]
C₆H₁₄O	[105-30-6]	<i>(dl)</i> 2-methyl-1-pentanol					
		$\Delta_v H$	(275–313)	59.4 ± 0.3	298	GS	[2001KUL/VER]
		$\Delta_v H$	(367–423)	49.3	382	A	[1987STE/MAL]
		$\Delta_v H$	(261–294)	64.9	279	A	[1987STE/MAL, 1979THO/MEA]
		$\Delta_v H$		57.4 ± 0.2	328	C	[1985MAJ/SVO2]
		$\Delta_v H$		55.7 ± 0.2	343	C	[1985MAJ/SVO2]
		$\Delta_v H$		53.9 ± 0.2	358	C	[1985MAJ/SVO2]
		$\Delta_v H$		52.7 ± 0.2	368	C	[1985MAJ/SVO2]
		$\Delta_v H$	(298–423)	54.2	313		[1973WIL/ZWO]
		$\Delta_v H$	(298–413)	50.2	356	I	[1938HOV/LAN]
C₆H₁₄O	[589-35-5]	<i>(dl)</i> 3-methyl-1-pentanol					
		$\Delta_v H$	(280–316)	61.7 ± 0.3	298	GS	[2001KUL/VER]
		$\Delta_v H$	(328–427)	54.8	343	A	[1987STE/MAL]
		$\Delta_v H$	(298–427)	59.7	313		[1973WIL/ZWO]
		$\Delta_v H$	(298–423)	47.2	360	I	[1940HOV/LAN2]
C₆H₁₄O	[626-89-1]	4-methyl-1-pentanol					
		$\Delta_v H$	(357–427)	53.0	372	A	[1987STE/MAL]
		$\Delta_v H$	(371–427)	51.1	386	A	[1987STE/MAL]
		$\Delta_v H$	(298–427)	63.9	313		[1973WIL/ZWO]
		$\Delta_v H$	(298–423)	46.5	360	I	[1940HOV/LAN2]
C₆H₁₄O	[590-36-3]	2-methyl-2-pentanol					
		$\Delta_v H$	(341–396)	44.2	356	A	[1987STE/MAL]
		$\Delta_v H$	(330–397)	48.9	345	A	[1987STE/MAL]
		$\Delta_v H$		54.7 ± 0.2	298	C	[1985MAJ/SVO2]
		$\Delta_v H$		52.8 ± 0.2	313	C	[1985MAJ/SVO2]
		$\Delta_v H$		50.7 ± 0.2	328	C	[1985MAJ/SVO2]
		$\Delta_v H$		48.5 ± 0.2	343	C	[1985MAJ/SVO2]
		$\Delta_v H$		46.1 ± 0.2	358	C	[1985MAJ/SVO2]
		$\Delta_v H$		44.4 ± 0.2	368	C	[1985MAJ/SVO2]
		$\Delta_v H$	(288–396)	58.3	303		[1973WIL/ZWO]
		$\Delta_v H$	(268–394)	49.1	283		[1947STU]
		$\Delta_v H$	(288–396)	51.3	303	I	[1933HOR/LAN]
C₆H₁₄O	[565-60-5]	<i>(dl)</i> 3-methyl-2-pentanol					
		$\Delta_v H$	(275–310)	58.2 ± 0.3	298	GS	[2001KUL/VER]
		$\Delta_v H$	(314–409)	54.4	329	A	[1987STE/MAL]
		$\Delta_v H$	(255–295)	60.4	280	A	[1987STE/MAL, 1979THO/MEA]
		$\Delta_v H$	(296–408)	54.8	311		[1973WIL/ZWO]
C₆H₁₄O	[108-11-2]	<i>(dl)</i> 4-methyl-2-pentanol					
		$\Delta_v H$	(274–301)	57.3 ± 0.3	298	GS	[2001KUL/VER]
		$\Delta_v H$	(240–295)	59.6	280	A	[1987STE/MAL]
		$\Delta_v H$	(293–406)	49.6	308		[1973WIL/ZWO]
		$\Delta_v H$	(353–404)	47.3	368	A,EB	[1987STE/MAL, 1970AND/BRA]
		$\Delta_v H$	(298–403)	45.6	350	I	[1938HOV/LAN]
C₆H₁₄O	[565-67-3]	<i>(dl)</i> 2-methyl-3-pentanol					
		$\Delta_v H$	(275–307)	56.0 ± 0.5	298	GS	[2001KUL/VER]
		$\Delta_v H$	(307–401)	52.2	322	A	[1987STE/MAL]
		$\Delta_v H$	(342–400)	45.4	357	A	[1987STE/MAL, 1975BRA/AND]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(298–401)	52.0	313		[1973WIL/ZWO]
	$\Delta_v H$	(298–399)	44.4	349	I	[1940HOV/LAN]
C ₆ H ₁₄ O	[77-74-7]	3-methyl-3-pentanol				
	$\Delta_v H$	(275–301)	55.7 ± 0.3	298	GS	[2001KUL/VER]
	$\Delta_v H$	(322–397)	40.1	337	A	[1987STE/MAL]
	$\Delta_v H$	(338–396)	46.4	353		[1973WIL/ZWO]
	$\Delta_v H$		56.7 ± 0.8	298		[1991WIB/HAO]
	$\Delta_v H$	(298–393)	42.1	346	I	[1940HOV/LAN]
C ₆ H ₁₄ O	[1185-33-7]	2,2-dimethyl-1-butanol				
	$\Delta_v H$	(356–415)	47.2	371	A	[1987STE/MAL]
	$\Delta_v H$	(298–415)	53.7	313		[1973WIL/ZWO]
C ₆ H ₁₄ O		(298–408)	52.1	313	I	[1940HOV/LAN3]
	[19550-30-2]	(dl) 2,3-dimethyl-1-butanol				
	$\Delta_v H$	(324–431)	51.4	339	A	[1987STE/MAL]
C ₆ H ₁₄ O		(373–422)	49.6	388		[1973WIL/ZWO]
	[624-95-3]	3,3-dimethyl-1-butanol				
C ₆ H ₁₄ O	$\Delta_{\text{fus}} H$		9.54	235.7		[2004MAS/NAK]
	$\Delta_v H$	(276–312)	58.0 ± 0.2	298	GS	[2001KUL/VER]
	$\Delta_v H$		58.6 ± 0.1	328	C	[1996ULB/KLU]
	$\Delta_v H$		55.4 ± 0.1	343	C	[1996ULB/KLU]
	$\Delta_v H$		52.4 ± 0.1	358	C	[1996ULB/KLU]
	$\Delta_v H$	(319–424)	50.8	334	A	[1987STE/MAL]
	$\Delta_v H$	(353–417)	49.4	368		[1973WIL/ZWO]
	C ₆ H ₁₄ O	[594-60-5]	2,3-dimethyl-2-butanol			
$\Delta_v H$		(303–340)	54.0 ± 0.8	298		[1991WIB/HAO]
$\Delta_v H$		(299–400)	48.8	314	A	[1987STE/MAL]
$\Delta_v H$		(298–393)	49.1	313		[1973WIL/ZWO]
C ₆ H ₁₄ O	[464-07-3]	(dl) 3,3-dimethyl-2-butanol				
	$\Delta_v H$	(280–315)	53.8 ± 0.3	298	GS	[2001KUL/VER]
	$\Delta_v H$	(302–401)	48.3	317	A	[1987STE/MAL]
C ₆ H ₁₄ O		(338–393)	46.8	353		[1973WIL/ZWO]
	[97-95-0]	2-ethyl-1-butanol				
	$\Delta_v H$	(275–313)	60.3 ± 0.3	298	GS	[2001KUL/VER]
C ₆ H ₁₄ O		(321–426)	53.1	336	A	[1987STE/MAL]
	$\Delta_v H$	(262–295)	65.4	280	A	[1987STE/MAL, 1979THO/MEA]
	$\Delta_v H$	(298–426)	59.6	313		[1973WIL/ZWO]
	$\Delta_v H$	(298–418)	U 45.5	313	I	[1940HOV/LAN3]
	C ₆ H ₁₄ OS	[na]	2-methyl-2-propanesulfonic acid, ethyl ester			
$\Delta_v H$		(337–343)	U 14.0	340	A	[1987STE/MAL]
C ₆ H ₁₄ O ₂	[111-76-2]	2-butoxyethanol				
	$\Delta_{\text{fus}} H$		11.8	199.5		[2000ATA/KAW]
	$\Delta_v H$	(363–382)	51.2	373	MM	[1999ESC/SAN]
	$\Delta_v H$	(336–443)	49.5	351	A	[1987STE/MAL, 1972DYK]
C ₆ H ₁₄ O ₂		(336–443)	52.6	351		[1957DYK/SEP, 1984BOU/FRI]
	[4461-87-4]	1,1-dimethoxybutane				
C ₆ H ₁₄ O ₂	$\Delta_v H$	(304–329)	41.2	317	EB	[1994WIB/MOR]
	[3453-99-4]	2,2-dimethoxybutane				

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{fus}}H$		9.32	174		[2003TEO/WIL]
C ₆ H ₁₄ O ₂	[105-57-7]	1,1-diethoxyethane				
	$\Delta_{\text{fus}}H$		10.95	167		[2003TEO/WIL]
	Δ_vH	(275–308)	39.6 ± 0.3	298	GS	[1998VER/PEN, 2002VER]
	Δ_vH	(281–384)	41.6	296	A	[1987STE/MAL, 1972DYK]
	Δ_vH	(273–343)	39.8	288		[1949NIC/LAF, 1984BOU/FRI]
	Δ_vH	(239–392)	36.2	255		[1947STU]
C ₆ H ₁₄ O ₂	[629-14-1]	1,2-diethoxyethane				
	Δ_vH	(339–382)	39.3	361		[1987TRE/LU]
	Δ_vH	(239–393)	37.9	254	A	[1987STE/MAL]
	Δ_vH		43.2 ± 0.1	298	C	[1970KUS/WAD]
C ₆ H ₁₄ O ₂	[77078-18-3]	1-methoxy-2-propoxyethane				
	Δ_vH		43.7 ± 0.1	298	C	[1970KUS/WAD]
C ₆ H ₁₄ O ₂	[4439-24-1]	2-isobutoxyethanol				
	Δ_vH	(344–432)	48.1	359	A	[1987STE/MAL, 1972DYK, 1957DYK/SEP, 1984BOU/FRI]
C ₆ H ₁₄ O ₂	[7580-85-0]	2- <i>tert</i> -butoxyethanol				
	$\Delta_{\text{fus}}H$		11.4	223.1		[2000ATA/KAW]
C ₆ H ₁₄ O ₂	[6920-22-5]	1,2-hexanediol				
	Δ_vH	(294–348)	78.7 ± 0.3	298	GS	[2004VER2]
C ₆ H ₁₄ O ₂	[629-11-8]	1,6-hexanediol				
	$\Delta_{\text{fus}}H$		25.5	316		[2005SMI/KAN]
	$\Delta_{\text{fus}}H$		25.52	340.6		[1996DOM/HEA]
	$\Delta_{\text{sub}}H$		112.0 ± 0.4	298	C	[1990KNA/SAB]
	$\Delta_{\text{sub}}H$		108.8			[1972GAR/HUS, 1977PED/RYL]
	Δ_vH	(355–559)	98.5 ± 1.8	298	EB,IP	[1996STE/CHI2]
	Δ_vH	(355–559)	87.8 ± 1.1	360	EB,IP	[1996STE/CHI2]
	Δ_vH	(355–559)	80.8 ± 0.9	400	EB,IP	[1996STE/CHI2]
	Δ_vH	(355–559)	73.9 ± 0.7	440	EB,IP	[1996STE/CHI2]
	Δ_vH	(355–559)	67.0 ± 0.6	480	EB,IP	[1996STE/CHI2]
	Δ_vH		87.0	342		[1993PIA/FER, 2006UMN/KWE]
	Δ_vH		90.9 ± 4.1	298		[1993PIA/FER, 2006UMN/KWE]
	Δ_vH		90.7 ± 1.1	298		[1990KNA/SAB]
	Δ_vH		83.3 ± 1.7			[1972GAR/HUS, 1977PED/RYL]
C ₆ H ₁₄ O ₂	[4457-71-0]	3-methyl-1,5-pentanediol				
	Δ_vH	(402–485)	76.9	417	A	[1987STE/MAL]
C ₆ H ₁₄ O ₂	[107-41-5]	(<i>dl</i>) 2-methyl-2,4-pentanediol				
	Δ_vH	(285–329)	68.9 ± 0.4	298	GS	[2007VER]
	Δ_vH	(370–547)	68.6 ± 0.4	298	EB	[1990DAU/HUT, 2007VER]
	Δ_vH	(373–473)	58.1	388	A	[1987STE/MAL]
C ₆ H ₁₄ O ₂	[76-09-5]	2,3-dimethyl-2,3-butanediol				
	$\Delta_{\text{fus}}H$		14.7	316.2	DSC	[1983PRI/WOO]
	Δ_vH	(346–448)	59.1	361	A	[1987STE/MAL]
C ₆ H ₁₄ O ₂ S	[34008-94-1]	<i>tert</i> -butyl ethyl sulfone				
	$\Delta_{\text{sub}}H$		86.6 ± 2.5			[UR/MAC, 1970COX/PIL]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₆ H ₁₄ O ₃	[111-96-6]	diethylene glycol, dimethyl ether				
	$\Delta_{\text{fus}}H$		17.78	209.1		[1996DOM/HEA]
	Δ_vH	(371–434)	45.4		EB	[2009LI/FAN]
	Δ_vH	(286–433)	48.0 ± 0.6	298	CGC	[2000NIC/ORF]
C ₆ H ₁₄ O ₃	[5648-29-3]	3,5,7-trioxanonane				
	Δ_vH		44.7 ± 0.2	298	C	[1969MAN]
C ₆ H ₁₄ O ₃	[15476-85-4]	<i>tert</i> -butyl 2-hydroxyethyl peroxide				
	Δ_vH		66.4 ± 1.9			[1983VAN/KAC]
C ₆ H ₁₄ O ₃	[111-90-0]	diethylene glycol, monoethyl ether				
	Δ_vH	(318–475)	52.1	333	A	[1987STE/MAL, 1947STU]
C ₆ H ₁₄ O ₃	[25265-71-8]	dipropylene glycol				
	Δ_vH	(423–505)	61.2	438	A	[1987STE/MAL]
C ₆ H ₁₄ O ₃	[77-99-6]	2-ethyl-2-hydroxymethyl-1,3-propanediol				
	$\Delta_{\text{trs}}H$		16.36	327.8		
	$\Delta_{\text{fus}}H$		0.9	332.7	DSC	[2002CHA/MAN]
	$\Delta_{\text{fus}}H$	(270–354)	21.45	333.4	AC	[1989ZHA/YAN]
C ₆ H ₁₄ O ₃	[106-69-4]	1,2,6-trihydroxyhexane				
	Δ_vH	(393–433)	97.2	408	A	[1987STE/MAL]
C ₆ H ₁₄ O ₄	[2517-44-4]	1,1,2,2-tetramethoxyethane				
	Δ_vH	(351–432)	42.9	366	A	[1987STE/MAL]
C ₆ H ₁₄ O ₄	[112-27-6]	triethylene glycol				
	Δ_vH	(442–562)	72.2 ± 0.3	440	EB	[2002STE/CHI3]
	Δ_vH	(442–562)	68.5 ± 0.3	480	EB	[2002STE/CHI3]
	Δ_vH	(442–562)	64.6 ± 0.3	520	EB	[2002STE/CHI3]
	Δ_vH	(442–562)	60.8 ± 0.5	560	EB	[2002STE/CHI3]
	Δ_vH	(278–323)	77.0	300		[1972MCF/SOM]
	Δ_vH	(288–303)	67.7	295	A	[1987STE/MAL, 1955ISH/MAT]
	Δ_vH	(293–303)	60.5	298		[1950WIS/PUC]
C ₆ H ₁₄ O ₆	[na]	dulcitol				
	$\Delta_{\text{fus}}H$		65.1	460.3		[1996DOM/HEA]
	Δ_vH	(464–496)	133.8 ± 1.4	482	TE	[1990BAR/DEL]
C ₆ H ₁₄ O ₆	[69-65-8]	<i>(d)</i> -mannitol				
	$\Delta_{\text{fus}}H$		54.69	437.3	DSC	[2010TON/LIU]
	$\Delta_{\text{fus}}H$		56.1	439.1	DSC	[1996DOM/HEA, 1990BAR/DEL]
	$\Delta_{\text{sub}}H$		202	298	B	[1990BAR/DEL]
C ₆ H ₁₄ O ₆	[50-70-4]	<i>(d)</i> -sorbitol				
	$\Delta_{\text{fus}}H$		30.2	366.5	DSC	[1996DOM/HEA, 1990BAR/DEL]
	$\Delta_{\text{sub}}H$		186	298	B	[1990BAR/DEL]
C ₆ H ₁₄ O ₆	[na]					
	Δ_vH	(461–497)	132.4 ± 2.0	477	TE	[1990BAR/DEL]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound						
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference		
C ₆ H ₁₄ O ₆	[na]	sorbitol						
	$\Delta_{\text{fus}}H$	(80–390)	30.35	369.2	AC	[2008TON/TAN2]		
C ₆ H ₁₄ O ₆	[50-70-4]	sorbitol						
	$\Delta_{\text{fus}}H(\alpha)$		29.8	359.1	DSC	[2009NEZ/AER]		
	$\Delta_{\text{fus}}H(\beta)$		31.7	371.2	DSC	[2009NEZ/AER]		
C ₆ H ₁₄ O ₆	[608-66-2]	(<i>d</i>)-galactitol	205	298	B	[1990BAR/DEL]		
C ₆ H ₁₄ O ₆	[na]	(<i>l</i>)-iditol	30.9	352.8		[1993SIN/CAR]		
C ₆ H ₁₄ S	[1741-83-9]	methyl pentyl sulfide						
	Δ_vH	(321–349)	44.2	336		[1999DYK/SVO]		
	Δ_vH		45.2	298		[1981SHI/SAI]		
	Δ_vH		44.6 ± 0.8	298	GC	[1964MAC/MCC]		
C ₆ H ₁₄ S	[638-46-0]	butyl ethyl sulfide	$\Delta_{\text{fus}}H$	12.39	178.1		[1985DEA]	
			Δ_vH	(314–445)	43.7	319		[1999DYK/SVO]
			Δ_vH		44.5	298		[1981SHI/SAI]
			Δ_vH		44.9	298		[1971WIL/ZWO]
			Δ_vH		44.6 ± 0.8	298	GC	[1964MAC/MCC]
			Δ_vH	(316–348)	43.5	333	EB	[1962MAC/MAY2]
			Δ_vH	(354–424)	40.7	369	A,EB	[1987STE/MAL, 1952WHI/BER]
C ₆ H ₁₄ S	[5008-72-0]	sec-butyl ethyl sulfide	Δ_vH	(304–434)	41.2	319		[1999DYK/SVO]
			Δ_vH	(345–409)	39.0	360	A,EB	[1987STE/MAL, 1952WHI/BER]
C ₆ H ₁₄ S	[14290-92-7]	<i>tert</i> -butyl ethyl sulfide	Δ_vH	(293–420)	39.2	308		[1999DYK/SVO]
			Δ_vH		39.3	298		[1971WIL/ZWO]
			Δ_vH	(332–400)	37.1	347	A,EB	[1987STE/MAL, 1952WHI/BER]
C ₆ H ₁₄ S	[625-80-9]	diisopropyl sulfide	$\Delta_{\text{fus}}H$		10.42	195.1		[1996DOM/HEA]
			Δ_vH	(293–420)	39.4	308		[1999DYK/SVO]
			Δ_vH		39.6 ± 0.1	298		[1972GOO, 1966OSB/DOU]
			Δ_vH	(324–433)	37.7	339	A,EB	[1987STE/MAL, 1966OSB/DOU]
			Δ_vH		39.6 ± 0.8	298	GC	[1964MAC/MCC]
			Δ_vH	(303–328)	38.5	318	EB	[1962MAC/MAY2]
			Δ_vH	(330–400)	37.4	345	EB	[1952WHI/BER]
C ₆ H ₁₄ S	[111-47-7]	dipropyl sulfide	$\Delta_{\text{fus}}H$		12.13	170.4		[1996DOM/HEA]
			Δ_vH	(313–411)	42.9	328		[1999DYK/SVO]
			Δ_vH		44.2	298		[1981SHI/SAI]
			Δ_vH		44.5	298		[1971WIL/ZWO]
			Δ_vH		39.5	298		[1971WIL/ZWO]
			Δ_vH		44.7 ± 0.8	298	GC	[1964MAC/MCC]
			Δ_vH	(353–427)	40.6	368	A,EB	[1987STE/MAL, 1952WHI/BER]
C ₆ H ₁₄ S	[5008-73-1]	isopropyl propyl sulfide						
Δ_vH	(303–432)	41.1	318		[1999DYK/SVO]			

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$		41.8	298		[1981SHI/SAI]
	$\Delta_v H$	(343–416)	39.0	358	A,EB	[1987STE/MAL, 1952WHI/BER]
C ₆ H ₁₄ S	[1613-45-2]	ethyl isobutyl sulfide				
	$\Delta_v H$	(305–401)	41.3	320		[1999DYK/SVO]
	$\Delta_v H$	(345–414)	39.2	360	A,EB	[1987STE/MAL, 1952WHI/BER]
C ₆ H ₁₄ S	[111-31-9]	1-hexanethiol				
	$\Delta_{\text{fus}} H$		18.03	192.6		[1996DOM/HEA]
	$\Delta_v H$	(320–454)	43.9	335		[1999DYK/SVO]
	$\Delta_v H$		44.8 ± 0.2	298		[1966GOO/DEP, 1966OSB/DOU]
	$\Delta_v H$	(352–468)	42.4	367	A,EB	[1987STE/MAL, 1966OSB/DOU]
C ₆ H ₁₄ S	[1679-06-7]	2-hexanethiol				
	$\Delta_v H$	(310–440)	42.7	325		[1999DYK/SVO]
	$\Delta_v H$	(328–423)	41.4	343	A	[1987STE/MAL]
C ₆ H ₁₄ S	[1639-01-6]	2,3-dimethyl-2-butanethiol				
	$\Delta_v H$	(285–318)	39.3	300		[1999DYK/SVO]
	$\Delta_v H$	(318–441)	37.8	333		[1999DYK/SVO]
	$\Delta_v H$		39.3 ± 0.1	298		[1972GOO, 1966OSB/DOU]
	$\Delta_v H$	(328–441)	37.4	343	A,EB	[1987STE/MAL, 1966OSB/DOU]
C ₆ H ₁₄ S	[1633-97-2]	2-methyl-2-pentanethiol				
	$\Delta_v H$		40.0 ± 0.1	298		[1972GOO, 1966OSB/DOU]
	$\Delta_v H$	(327–439)	38.0	342	A,EB	[1987STE/MAL, 1966OSB/DOU, 1999DYK/SVO]
C ₆ H ₁₄ S ₂	[4253-89-8]	diisopropyl disulfide				
	$\Delta_v H$	(383–423)	49.3	298	CGC	[1995CHI/HOS]
	$\Delta_v H$		39.6	298		[1981SHI/SAI]
	$\Delta_v H$	(377–447)	43.8	392	A,EB	[1987STE/MAL, 1952WHI/BER, 1999DYK/SVO]
C ₆ H ₁₄ S ₂	[629-19-6]	dipropyl disulfide				
	$\Delta_{\text{fus}} H$		13.81	187.7		[1996DOM/HEA]
	$\Delta_v H$	(354–499)	47.8	369		[1999DYK/SVO]
	$\Delta_v H$		53.8 ± 0.1	298	C	[1985KUS]
	$\Delta_v H$		53.8	298		[1981SHI/SAI]
	$\Delta_v H$	(389–447)	47.0	404	A,EB	[1987STE/MAL, 1958HUB/DOU, 1966OSB/DOU]
	$\Delta_v H$	(395–456)	46.6	410	EB	[1952WHI/BER]
C ₆ H ₁₄ S ₂	[4151-69-3]	ethyl (1,1-dimethylethyl) disulfide				
	$\Delta_v H$	(373–461)	43.4	388	A,EB	[1987STE/MAL, 1952WHI/BER, 1999DYK/SVO]
C ₆ H ₁₄ S ₂	[33672-51-4]	isopropyl propyl disulfide				
	$\Delta_v H$	(383–433)	45.4	398	A	[1987STE/MAL, 1999DYK/SVO]
C ₆ H ₁₄ S ₂	[1191-43-1]	1,6-hexanedithiol				
	$\Delta_v H$	(379–511)	55.7	394	A	[1987STE/MAL, 1999DYK/SVO]
C ₆ H ₁₄ S ₂	[5395-75-5]	3,6-dithiaoctane				
	$\Delta_v H$		59.5 ± 0.1	298	C	[1974MAN4]
C ₆ H ₁₄ S ₃	[na]	trithiodiethylene glycol, dimethyl ether				
	$\Delta_v H$	(391–418)	103.7	404	A	[1987STE/MAL]

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₆ H ₁₄ S ₃	[37460-04-1]	2,5,8-trithianonane				
	$\Delta_v H$	(391–533)	116.4	406		[1999DYK/SVO]
C ₆ H ₁₅ N	[111-26-2]	hexylamine				
	$\Delta_v H$	(323–373)	45.0	298	CGC	[1995CHI/HOS]
	$\Delta_v H$	(303–406)	42.2	318	A	[1987STE/MAL]
	$\Delta_v H$		45.1 ± 0.1	298	C	[1969WAD]
C ₆ H ₁₅ N	[21035-44-9]	<i>(dl)</i> sec-butyl ethyl amine				
	$\Delta_v H$	(283–372)	37.9	298		[1987STE/MAL, 1972DYK]
C ₆ H ₁₅ N	[108-18-9]	diisopropylamine				
	$\Delta_v H$	(260–412)	35.4	275	A	[1987STE/MAL]
	$\Delta_v H$	(273–367)	35.6	288	A	[1987STE/MAL]
	$\Delta_v H$		34.6 ± 0.1	298	C	[1979PET/MAJ]
	$\Delta_v H$		33.7 ± 0.1	313	C	[1979PET/MAJ]
	$\Delta_v H$		32.6 ± 0.1	328	C	[1979PET/MAJ]
	$\Delta_v H$		31.5 ± 0.1	343	C	[1979PET/MAJ]
	$\Delta_v H$		30.2 ± 0.1	358	C	[1979PET/MAJ]
	$\Delta_v H$	(300–356)	34.4	315	EB	[1979PET/MAJ]
	$\Delta_v H$	(291–305)	34.6	298		[1971LEB/KAT2]
	$\Delta_v H$		34.5 ± 0.1	298	C	[1969WAD]
$\Delta_v H$	(273–333)	33.8 ± 0.2	298	I	[1969FRA/WAT]	
C ₆ H ₁₅ N	[21968-17-2]	N-isopropyl propylamine				
	$\Delta_v H$		37.3 ± 0.1	298	C	[1979PET/MAJ]
	$\Delta_v H$		36.2 ± 0.1	313	C	[1979PET/MAJ]
	$\Delta_v H$		35.2 ± 0.1	328	C	[1979PET/MAJ]
	$\Delta_v H$		34.1 ± 0.1	343	C	[1979PET/MAJ]
	$\Delta_v H$		33.0 ± 0.1	358	C	[1979PET/MAJ]
	$\Delta_v H$	(312–369)	36.2	327	EB	[1979PET/MAJ]
C ₆ H ₁₅ N	[13360-63-9]	N-butylethylamine				
	$\Delta_v H$		40.2 ± 0.1	298	C	[1979PET/MAJ]
	$\Delta_v H$		39.1 ± 0.1	313	C	[1979PET/MAJ]
	$\Delta_v H$		38.0 ± 0.1	328	C	[1979PET/MAJ]
	$\Delta_v H$		36.9 ± 0.1	343	C	[1979PET/MAJ]
	$\Delta_v H$		35.8 ± 0.1	358	C	[1979PET/MAJ]
	$\Delta_v H$	(313–375)	39.9	328	EB	[1979PET/MAJ]
$\Delta_v H$	(283–382)	41.4	298	A	[1987STE/MAL, 1972DYK]	
C ₆ H ₁₅ N	[142-84-7]	dipropylamine				
	$\Delta_v H$	(321–382)	40.0	336		[2000RES/GON]
	$\Delta_v H$	(302–422)	39.8	317	A	[1987STE/MAL, 1972DYK]
	$\Delta_v H$	(291–305)	41.5	298		[1971LEB/KAT2]
	$\Delta_v H$		40.0 ± 0.1	298	C	[1969WAD]
	$\Delta_v H$	(273–333)	40.2 ± 0.3	298	I	[1969FRA/WAT]
C ₆ H ₁₅ N	[121-44-8]	triethylamine				
	$\Delta_v H$	(273–353)	33.4 ± 0.2	313		[2009MOK/RAZ]
	$\Delta_v H$	(273–353)	35.4 ± 0.2	298		[2009MOK/RAZ]
	$\Delta_v H$	(310–362)	33.9	325	EB	[2006WAN/FAN]
	$\Delta_v H$	(231–319)	35.2 ± 0.9	275		[2001BAE]
	$\Delta_v H$	(302–338)	34.1	317	EB	[1990DUT/KAH]
	$\Delta_v H$	(298–324)	34.6	311		[1987STE/MAL]
	$\Delta_v H$	(283–363)	35.5	298		[1987STE/MAL]
	$\Delta_v H$		34.8 ± 0.2	298	C	[1979MAJ/SVO2]
$\Delta_v H$		33.9 ± 0.1	313	C	[1979MAJ/SVO2]	

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		T _m (K)	Method	Reference
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)			
		$\Delta_v H$	33.0 ± 0.2	328	C	[1979MAJ/SVO2]
		$\Delta_v H$	32.2 ± 0.1	343	C	[1979MAJ/SVO2]
		$\Delta_v H$	31.3 ± 0.2	358	C	[1979MAJ/SVO2]
		$\Delta_v H$	(303–361) 34.8	318	EB	[1979MAJ/SVO2]
		$\Delta_v H$	(283–313) 35.1	298		[1975CHU/DRU]
		$\Delta_v H$	(324–357) 33.3	339		[1971BAY/LET]
		$\Delta_v H$	34.9 ± 0.1	298	C	[1969WAD]
		$\Delta_v H$	(298–363) U 25.3	313	I	[1953COP/EVE]
		$\Delta_v H$	(285–337) 33.0	311		[1936THO/LIN]
C₆H₁₅N	[918-02-5]		N,N-dimethyl <i>tert</i> -butyl amine			
		$\Delta_v H$	(283–318) 34.8	298	A	[1987STE/MAL]
C₆H₁₅NO	[5888-29-9]		N-(methoxymethyl)diethylamine			
		$\Delta_v H$	(293–318) 38.0	305	A	[1987STE/MAL]
C₆H₁₅NO	[100-37-8]		N,N-diethylethanolamine			
		$\Delta_v H$	(278–318) 52.5 ± 0.2	298	GS	[2005KAP/SLO]
		$\Delta_v H$	(332–475) 48.5 ± 0.2	340	EB	[2002STE/CHI5]
		$\Delta_v H$	(332–475) 45.0 ± 0.2	380	EB	[2002STE/CHI5]
		$\Delta_v H$	(332–475) 41.6 ± 0.4	420	EB	[2002STE/CHI5]
		$\Delta_v H$	(332–475) 37.8 ± 0.7	460	EB	[2002STE/CHI5]
		$\Delta_v H$	(328–433) 48.5	343	A	[1987STE/MAL]
		$\Delta_v H$	(283–318) 58.5 ± 1.3	298		[1977LEB/NAZ, 2005KAP/SLO]
C₆H₁₅NO₂	[110-97-4]		diisopropanolamine			
		$\Delta_v H$	(390–521) 68.0	405	A	[1987STE/MAL, 1972DYK]
C₆H₁₅NO₂	[1704-62-7]		2-[2-(dimethylamino)ethoxy]ethanol			
		$\Delta_v H$	(412–452) 54.4	427	A	[1987STE/MAL]
C₆H₁₅NO₂S	[33718-39-7]		N,N-diethyl ethanesulfonamide			
		$\Delta_v H$	(392–526) 55.4	407	A	[1987STE/MAL]
C₆H₁₅NO₃	[102-71-6]		triethanolamine			
		$\Delta_v H$	(523–579) 79.3	538	A	[1987STE/MAL, 1959MCD/SHR, 1984BOU/FRI]
C₆H₁₅NS	[na]		N,N-dimethyl-S- <i>tert</i> -butylthiohydroxylamine			
		$\Delta_v H$	(328–334) 28.3	331	A	[1987STE/MAL, 1999DYK/SVO]
C₆H₁₅N₃	[108-74-7]		1,3,5-trimethylhexahydro-s-triazine			
		$\Delta_v H$	(284–328) 50.8 ± 0.8	306	GS	[2002VER2]
		$\Delta_v H$	(284–328) 51.2 ± 0.8	298	GS	[2002VER2]
C₆H₁₅N₃	[140-31-8]		1-(2-aminoethyl)piperazine			
		$\Delta_v H$	(296–338) 68.7 ± 0.3	298	GS	[2010EFI/EME]
C₆H₁₅O₂PS₃	[640-15-3]		O,O-dimethyl-S-[2-(ethylthio)ethyl]dithiophosphate			
		$\Delta_v H$	(283–394) 76.8	298	A	[1987STE/MAL, 1999DYK/SVO]
C₆H₁₅O₃P	[1809-21-8]		phosphonic acid, dipropyl ester			
		$\Delta_v H$	(318–467) 38.1	333	A	[1987STE/MAL, 1972DYK]
C₆H₁₅O₃P	[122-52-1]		triethylphosphite			
		$\Delta_v H$	53.0	298		[2008SAG/SAF]
C₆H₁₅O₃PS	[126-68-1]		O,O,O-triethylthiophosphate			
		$\Delta_v H$	(305–335) 87.5	320	A	[1987STE/MAL, 1999DYK/SVO]
C₆H₁₅O₃PS	[1186-09-0]		O,O,S-triethylthiophosphate			

TABLE 6. Phase change enthalpies of C₅ to C₆ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(312–352)	76.3	327	A	[1987STE/MAL, 1999DYK/SVO]
C ₆ H ₁₅ O ₃ PS ₂	[867-27-6]	phosphorothioic acid, O-[2-(ethylthio)ethyl]-O,O-dimethyl ester				
	$\Delta_v H$	(283–379)	71	298	A	[1987STE/MAL, 1999DYK/SVO]
C ₆ H ₁₅ O ₃ PS ₂	[919-86-8]	phosphorothioic acid, S-[2-(ethylthio)ethyl]-O,O-dimethyl ester				
	$\Delta_v H$	(283–407)	78.8	298	A	[1987STE/MAL, 1999DYK/SVO]
C ₆ H ₁₅ O ₄ P	[78-40-0]	triethylphosphate				
	$\Delta_v H$	(413–453)	55.7	298	CGC	[2007PAN/ANT2]
	$\Delta_v H$	(312–484)	46.3	327	A	[1987STE/MAL, 1947STU]
C ₆ H ₁₅ P	[554-70-1]	triethylphosphine				
	$\Delta_{\text{fus}} H$		10.73	188.2		[1999SHE/KAM]
	$\Delta_v H$	(291–402)	38.3	306	A	[1987STE/MAL, 1972DYK]
C ₆ H ₁₆ FN ₂ OP	[371-86-8]	N,N'-diisopropyl phosphorodiamidic fluoride				
	$\Delta_v H$	(278–398)	58.1	293	A	[1987STE/MAL]
C ₆ H ₁₆ N ₂	[124-09-4]	1,6-hexanediamine				
	$\Delta_{\text{fus}} H$		39.38	311.6	DSC	[2006KHI/DAH2]
	$\Delta_{\text{fus}} H$		40.21	312.3	DSC	[2002DAL/DEL]
	$\Delta_v H$	(348–474)	49.3	363	A	[1987STE/MAL]
	$\Delta_v H$	(338–473)	51.3	353	A	[1987STE/MAL]
C ₆ H ₁₆ N ₂	[110-18-9]	1,2-bis(dimethylamino)ethane				
	$\Delta_v H$	(295–365)	39.8	330		[2009RAZ/HAJ]
	$\Delta_v H$	(295–365)	42.2	298		[2009RAZ/HAJ]
C ₆ H ₁₆ N ₂ O ₂	[4439-20-7]	N,N-bis(2-hydroxyethyl)ethylenediamine				
	$\Delta_{\text{fus}} H$		49.7	373.2		[1997STE/CHI4]
	$\Delta_{\text{sub}} H$		142.7	373	B	[1997STE/CHI4]
	$\Delta_v H$	(399–500)	106.4 ± 6.4	298	EB,IP	[1997STE/CHI2, 1997STE/CHI4]
	$\Delta_v H$	(399–500)	91.2 ± 0.2	400	EB,IP	[1997STE/CHI2, 1997STE/CHI4]
	$\Delta_v H$	(399–500)	87.7 ± 0.2	440	EB,IP	[1997STE/CHI2, 1997STE/CHI4]
C ₆ H ₁₆ N ₂ O ₂			84.8 ± 0.2	480	EB,IP	[1997STE/CHI2, 1997STE/CHI4]
	[929-59-9]	1,2-bis(2-aminoethoxy)ethane				
	$\Delta_v H$	(293–353)	56.2	323		[2009RAZ/HAJ]
	$\Delta_v H$	(293–353)	58.8	298		[2009RAZ/HAJ]
C ₆ H ₁₆ N ₂ O ₂	[3129-93-9]	diisopropyl ammonium nitrite				
	$\Delta_{\text{sub}} H$	(288–299)	39.0	293.5	A	[1987STE/MAL, 1965MAR]
C ₆ H ₁₈ N ₃ P	[1608-26-0]	tris(dimethylamino)phosphine				
	$\Delta_v H$		41.5 ± 0.6	298	STG	[1995ALM/FIN2]
	$\Delta_v H$	(298–333)	63.2	313		[1984MIC/JOS]
C ₆ H ₁₈ N ₄	[112-24-3]	triethylenetetramine				
	$\Delta_v H$	(338–373)	84.7 ± 0.3	298	GS	[2010EFI/EME]
	$\Delta_v H$	(294–325)	75.6	298	TGA	[1988AFZ/BUT, 2010EFI/EME]
	$\Delta_v H$	(431–550)	59.8	446	A	[1987STE/MAL, 1972DYK]
	$\Delta_v H$	(431–492)	71.0 ± 2.6	298		[1967SIV/MAT, 2010EFI/EME]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₇ ClF ₁₇ N ₂ S	[na]	CF ₃ SCI[=NCF(CF ₃) ₂] ₂				
	$\Delta_v H$		38.9	467	I	[1977KIT/SHR2]
		Note: This is the structure and molecular formula given in paper. A search of Chemical Abstracts by molecular formula failed to turn up any hits with this formula.				
C ₇ D ₈	[2037-26-5]	perdeuterotoluene				
	$\Delta_{\text{us}}H$		3.51	136	DSC	[1972AHM/EAD]
	$\Delta_{\text{fus}}H$		6.02	178	DSC	
C ₇ F ₆ O ₂	[59483-82-8]	carbonofluoridic acid pentafluorophenyl ester				
	$\Delta_v H$		42.3			[1976FAL/DES2]
C ₇ F ₈	[434-64-0]	perfluorotoluene				
	$\Delta_{\text{fus}}H$		13.2	207.7		
	$\Delta_{\text{fus}}H$		11.49	207		[1996DOM/HEA]
	$\Delta_v H$	(288–334)	40.5 ± 0.2	298		[2005DIA/GON]
	$\Delta_v H$	(291–378)	40.0	306		[1999DYK/SVO]
	$\Delta_v H$	(285–376)	40.9	300	A	[1987STE/MAL]
	$\Delta_v H$	(290–400)	40.4	298		[1984BOU/FRI, 1991BAS/SVO]
	$\Delta_v H$	(285–334)	41.6 ± 0.2	298		[1973KRE/PRI]
C ₇ F ₁₀	[14451-74-2]	3,3-difluoro-1,2-bis(trifluoromethyl)-4-(difluoroethylene)cyclobutene				
	$\Delta_v H$	(272–316)	31.5	287	A	[1987STE/MAL, 1999DYK/SVO]
C ₇ F ₁₂ O ₂ S ₄	[58936-62-2]	pentanebis(dithioperoxyic) acid, hexafluoro-bis(trifluoromethyl) ester				
	$\Delta_v H$		33.6	370	I	[1976BUR/SHR]
C ₇ F ₁₂ O ₆	[32751-20-5]	hexafluoroperoxyglutaric acid, bis(trifluoromethyl) ester				
	$\Delta_v H$	(200–390)	47.3	215	A	[1987STE/MAL, 1999DYK/SVO]
C ₇ F ₁₄	[355-02-2]	perfluoromethylcyclohexane				
	$\Delta_{\text{sub}}H$		51.6	234	B	[1963BON, 1957ROW/THA]
	$\Delta_v H$	(305–414)	33.1	320		[1999DYK/SVO]
	$\Delta_v H$	(413–488)	30.2	428		[1999DYK/SVO]
	$\Delta_v H$	(306–384)	34.0	298		[1984BOU/FRI, 1991BAS/SVO]
	$\Delta_v H$		34.1 ± 0.3	298		[1981VAR/BUL]
	$\Delta_v H$	(305–385)	33.4	320	A	[1987STE/MAL, 1970DYK/VAN, 1973KKY/REP]
	$\Delta_v H$	(306–384)	33.3	321		[1959GOO/DOU, 1984BOU/FRI]
	$\Delta_v H$	(298–353)	33.8	313		[1957ROW/THA, 1984BOU/FRI]
$\Delta_v H$	(272–349)	33.3	310		[1956GLE/REE, 1970DYK/VAN]	
C ₇ F ₁₅ NS	[77984-26-0]	2,2,3,3,4,4,5,5-octafluoro-1,1,2,3,4,5-hexahydro-1-[[1,2,2,2-tetrafluoro-1-(trifluoromethyl)ethyl]imino]thiophene				
	$\Delta_v H$		33.9	371		[1981ABE/SHR]
C ₇ F ₁₆	[333-57-9]	perfluoroheptane				
	$\Delta_{\text{us}}H$		6.67	180.4		
	$\Delta_{\text{fus}}H$		6.95	221.9		[1996DOM/HEA, 1986STA]
	$\Delta_{\text{sub}}H$		57.7		B	[1963BON, 1951OLI/GRI]
	$\Delta_v H$	(363–474)	32.6	378		[1999DYK/SVO]
	$\Delta_v H$	(304–390)	36.3 ± 0.3	298	EB	[1997STE/CHI3]
	$\Delta_v H$	(290–355)	35.9	298		[1984BOU/FRI, 1991BAS/SVO]
	$\Delta_v H$		33.1			[1959YEN/REE]
	$\Delta_v H$	(293–355)	34.9	324		[1956GLE/REE]
$\Delta_v H$	(271–379)	37.7	286	A	[1987STE/MAL, 1951OLI/GRI, 1970DYK/VAN]	

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₇ F ₁₆ N ₂ OS	[62609-64-7]	1,1,1-trifluoro-N'-[1,2,2,2-tetrafluoro-1-(trifluoromethyl)ethyl]-N-[2,2,2-trifluoro-1-(trifluoromethyl)ethylidene]methanesulfonimidamide	$\Delta_v H$ 33.5	451		[1977KIT/SHR]
C ₇ F ₁₇ N	[338-81-8]	perfluoro-N,N-diethylpropylamine	$\Delta_v H$ (283–366)	39.2	325	[1999DYK/SVO]
C ₇ HF ₅ O ₂	[602-94-8]	pentafluorobenzoic acid	$\Delta_{\text{sub}} H$ (335–359)	91.6 ± 4.2		GS [1969COX/GUN, 1970COX/PIL]
C ₇ HF ₁₃ O ₂	[375-85-9]	tridecafluoroheptanoic acid	$\Delta_v H$ (359–485)	61.4 ± 0.3	370	EB [2002STE/CHI]
			$\Delta_v H$ (359–485)	55.5 ± 0.3	410	EB [2002STE/CHI]
			$\Delta_v H$ (359–485)	48.7 ± 0.7	450	EB [2002STE/CHI]
C ₇ HF ₁₅	[375-83-7]	1,1,1,2,2,3,3,4,4,5,5,6,6,7,7-pentafluoroheptane	$\Delta_v H$ (365–369)	30.7	367	[1966CAR/STE]
			$\Delta_v H$ (292–370)	37.1	307	A [1987STE/MAL, 1953KAR/SAY, 1973KKY/REP, 1999DYK/SVO]
C ₇ H ₂ F ₁₃ NO	[54181-88-3]	(E) 1,1,1,2,3,3,3-heptafluoro-N-[2,2,2-trifluoro-1-(2,2,2-trifluoroethoxy)ethylidene]-2-propanamine	$\Delta_v H$ 35.7	369		[1975PET/SHR]
C ₇ H ₃ Br ₂ NO	[1689-84-5]	3,5-dibromo-4-hydroxybenzotrile	$\Delta_{\text{fus}} H$ 32.03	464	DSC	[1990DON/DRE]
C ₇ H ₃ ClF ₃ NO ₂	[777-37-7]	1-(trifluoromethyl)-2-chloro-5-nitrobenzene	$\Delta_v H$ (364–508)	58.1	379	A [1987STE/MAL, 1953KAR/SAY, 1973KKY/REP, 1999DYK/SVO]
C ₇ H ₃ ClF ₃ NO ₂	[121-17-5]	1-(trifluoromethyl)-4-chloro-3-nitrobenzene	$\Delta_v H$ (358–495)	57.6	373	A [1987STE/MAL, 1973KKY/REP, 1999DYK/SVO]
C ₇ H ₃ Cl ₂ F ₃	[328-84-7]	1-(trifluoromethyl)-3,4-dichlorobenzene	$\Delta_v H$ (353–453)	44.1	368	A [1987STE/MAL]
			$\Delta_v H$ (284–446)	41.8	299	[1947STU]
C ₇ H ₃ Cl ₂ N	[1194-65-6]	2,6-dichlorobenzotrile	$\Delta_{\text{fus}} H$	24.56	421.2	DSC [2000ROD/VEC]
			$\Delta_{\text{fus}} H$	26.17	417.2	DSC [1991ACR, 1990DON/DRE]
			$\Delta_{\text{fus}} H$	25.94	416.7	DSC [1972PLA]
C ₇ H ₃ Cl ₂ NO	[102-36-3]	3,4-dichlorophenylisocyanate	$\Delta_v H$ (373–473)	47.4	388	A [1987STE/MAL]
C ₇ H ₃ Cl ₃ O ₂	[50-31-7]	2,3,6-trichlorobenzoic acid	$\Delta_{\text{fus}} H$ 23.85	402.7		[1991ACR]
C ₇ H ₃ Cl ₅	[13014-24-9]	1-(trichloromethyl)-3,4-dichlorobenzene	$\Delta_v H$ (438–663)	59.3	453	A [1987STE/MAL, 1970DYK/VAN, 1973KKY/REP, 1999DYK/SVO]
C ₇ H ₃ F ₅	[771-56-2]	2,3,4,5,6-pentafluorotoluene	$\Delta_{\text{ms}} H$	0.7	70.3	
			$\Delta_{\text{fus}} H$	13.28	243.7	[1996DOM/HEA, 1968COU/HAL]
			$\Delta_v H$ (403–523)	36.1	418	[1999DYK/SVO]
			$\Delta_v H$ (493–564)	34.9	508	[1999DYK/SVO]
			$\Delta_v H$ (310–410)	41.2	298	[1984BOU/FRI, 1991BAS/SVO]
		$\Delta_v H$ (312–416)	39.9	327	A [1987STE/MAL, 1968AMB, 1973KKY/REP, 1999DYK/SVO]	

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₇ H ₃ I ₂ NO	[1689-83-4] $\Delta_{\text{fus}}H$	4-hydroxy-3,5-diiodobenzonitrile	33.63	482.9	DSC	[1990DON/DRE]
C ₇ H ₃ I ₃ O ₂	[88-82-4] $\Delta_{\text{fus}}H$	2,3,5-triiodobenzoic acid	32.23	503.8		[1991ACR]
C ₇ H ₄ ClF ₃	[88-16-4] $\Delta_{\text{fus}}H$	1-(trifluoromethyl)-2-chlorobenzene	11.6	264	DSC	[1972AHM/EAD]
	Δ_vH	(310–426)	44.6	325	A	[1987STE/MAL, 1951POT/SAY, 1970DYK/VAN, 1973KKY/REP]
C ₇ H ₄ ClF ₃	[98-15-7] Δ_vH	1-(trifluoromethyl)-3-chlorobenzene	43.0	317	A	[1987STE/MAL, 1951POT/SAY, 1970DYK/VAN, 1973KKY/REP]
C ₇ H ₄ ClF ₃	[98-56-6] Δ_vH	1-(trifluoromethyl)-4-chlorobenzene	42.2	317	A	[1987STE/MAL, 1951POT/SAY, 1970DYK/VAN, 1973KKY/REP]
C ₇ H ₄ CIN	[873-32-5] Δ_vH	2-chlorobenzonitrile	53.5	393	EB	[1994AIM2]
C ₇ H ₄ CIN	[623-03-0] Δ_vH	4-chlorobenzonitrile	51.9	404	EB	[1994AIM2]
C ₇ H ₄ CINO	[2909-38-8] Δ_vH	3-chlorophenyl isocyanate	53.1	359	A	[1987STE/MAL, 1964GOL/GOR]
C ₇ H ₄ CINO	[104-12-1] Δ_vH	4-chlorophenyl isocyanate	48.9	378	A	[1987STE/MAL]
	Δ_vH	(323–433)	44.3	338		[1967KON/ZHU]
C ₇ H ₄ CINO ₃	[121-90-4] Δ_vH	3-nitrobenzoyl chloride	62.4	443	A	[1987STE/MAL, 1999DYK/SVO]
C ₇ H ₄ Cl ₂ O	[609-65-4] Δ_vH	2-chlorobenzoyl chloride	53.4	384	A	[1987STE/MAL, 1999DYK/SVO]
C ₇ H ₄ Cl ₂ O	[618-46-2] Δ_vH	3-chlorobenzoyl chloride	49.4	382	A	[1987STE/MAL, 1999DYK/SVO]
C ₇ H ₄ Cl ₂ O	[122-01-0] Δ_vH	4-chlorobenzoyl chloride	55.7	381	A	[1987STE/MAL, 1999DYK/SVO]
C ₇ H ₄ Cl ₂ O	[874-42-0] $\Delta_{\text{fus}}H$	2,4-dichlorobenzaldehyde	20.47	347.2		[2004WAN/TAN4]
C ₇ H ₄ Cl ₂ O ₂	[51-36-5] $\Delta_{\text{fus}}H$	3,5-dichlorobenzoic acid	22.97	459.3		[1991ACR]
C ₇ H ₄ Cl ₃ NO ₃	[55335-06-3] $\Delta_{\text{fus}}H$	3,5,6-trichloro-2-pyridinyloxyacetic acid	31.17	423.3	DSC	[1990DON/DRE]
C ₇ H ₄ Cl ₄	[2136-89-2] Δ_vH	1-(trichloromethyl)-2-chlorobenzene	55.0	438	A	[1987STE/MAL, 1970DYK/VAN]
C ₇ H ₄ Cl ₄	[1006-31-1] Δ_vH	2,3,5,6-tetrachlorotoluene	52.6	414	A	[1987STE/MAL, 1973FEL/SAV, 1999DYK/SVO]
C ₇ H ₄ F ₃ NO ₂	[98-46-4] Δ_vH	1-(trifluoromethyl)-3-nitrobenzene	53.8	356	A	[1987STE/MAL, 1953KAR/SAY, 1999DYK/SVO]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₇ H ₄ F ₄	[5230-78-4]	1,2,4,5-tetrafluoro-3-methylbenzene				
	$\Delta_{\text{trs}}H$		3.64	218	DSC	
	$\Delta_{\text{fus}}H$		5.84	233	DSC	[1972AHM/EAD]
C ₇ H ₄ F ₄	[392-85-8]	1-(trifluoromethyl)-2-fluorobenzene				
	$\Delta_{\text{fus}}H$		10.7	222	DSC	[1972AHM/EAD]
	Δ_vH	(310–410)	38.1	298		[1984BOU/FRI, 1991BAS/SVO]
C ₇ H ₄ F ₄	[401-80-9]	1-(trifluoromethyl)-3-fluorobenzene				
	Δ_vH	(313–410)	36.8	328	A	[1987STE/MAL, 1970DYK/VAN]
C ₇ H ₄ F ₄	[402-44-8]	1-(trifluoromethyl)-4-fluorobenzene				
	Δ_vH	(286–381)	35.8	301	A	[1987STE/MAL, 1970DYK/VAN]
C ₇ H ₄ F ₁₂ O	[335-99-9]	2,2,3,3,4,4,5,5,6,6,7,7-dodecafluoro-1-heptanol				
	Δ_vH	(355–446)	53.4	370	A	[1987STE/MAL, 1999DYK/SVO]
C ₇ H ₄ N ₂ O ₂	[612-24-8]	2-nitrobenzotrile				
	$\Delta_{\text{trs}}H$		1.57	338.1		
	$\Delta_{\text{fus}}H$		15.72	382.7	DSC	[2002JIM/ROU2]
	$\Delta_{\text{sub}}H$	(297–311)	87.9 ± 1.4	304	ME	[2003ROU/JIM]
	$\Delta_{\text{sub}}H$	(297–311)	88.1 ± 1.4	298	ME	[2003ROU/JIM]
C ₇ H ₄ N ₂ O ₂	[619-24-9]	3-nitrobenzotrile				
	$\Delta_{\text{fus}}H$		20.49	389.7	DSC	[2002JIM/ROU2]
	$\Delta_{\text{sub}}H$	(306–324)	92.2 ± 0.3	316	ME	[2003RIB/SAN]
	$\Delta_{\text{sub}}H$	(306–324)	92.8 ± 0.3	298	ME	[2003ROU/JIM]
C ₇ H ₄ N ₂ O ₂	[619-72-7]	4-nitrobenzotrile				
	$\Delta_{\text{trs}}H$		0.45	349		
	$\Delta_{\text{trs}}H$		1.01	386		
	$\Delta_{\text{fus}}H$		17.73	420.6	DSC	[2002JIM/ROU2]
	$\Delta_{\text{sub}}H$	(305–322)	90.5 ± 1.3	313	ME	[2003ROU/JIM]
	$\Delta_{\text{sub}}H$	(305–322)	91.1 ± 1.3	298	ME	[2003ROU/JIM]
C ₇ H ₄ N ₂ O ₅	[528-75-6]	2,4-dinitrobenzaldehyde				
	$\Delta_{\text{fus}}H$	(78–368)	21.18	344.9	AC	[2005WAN/TAN3]
C ₇ H ₄ N ₂ O ₆	[610-30-0]	2,4-dinitrobenzoic acid				
	$\Delta_{\text{fus}}H$		30.6	455.8	DSC	[2009VEC/BRU]
	$\Delta_{\text{sub}}H$	(364–402)	132 ± 2	383	TE	[2009VEC/BRU]
	$\Delta_{\text{sub}}H$	(364–402)	135 ± 2	298	TE	[2009VEC/BRU]
	Δ_vH	(503–544)	92 ± 3	522	TGA	[2009VEC/BRU]
	Δ_vH	(494–539)	91 ± 1	517	TGA	[2009VEC/BRU]
C ₇ H ₄ N ₂ O ₆	[528-45-0]	3,4-dinitrobenzoic acid				
	$\Delta_{\text{fus}}H$		24.6	438.2	DSC	[2009VEC/BRU]
	$\Delta_{\text{sub}}H$	(366–399)	126 ± 2	386	TE	[2009VEC/BRU]
	$\Delta_{\text{sub}}H$	(366–399)	129 ± 2	298	TE	[2009VEC/BRU]
	Δ_vH	(518–540)	91 ± 3	529	TGA	[2009VEC/BRU]
	Δ_vH	(506–536)	91 ± 1	521	TGA	[2009VEC/BRU]
C ₇ H ₄ N ₂ O ₆	[99-34-3]	3,5-dinitrobenzoic acid				
	$\Delta_{\text{fus}}H$		22.8	480.4		[1971LEB/RYA]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound					
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference	
C ₇ H ₄ S ₃	[3354-42-5]	4,5-benzo-1,2-dithiole-3-thione					
		$\Delta_{\text{sub}}H$	(350–361)	102.6 ± 0.4	355		[1972GEI/RAU]
		$\Delta_{\text{sub}}H$		107 ± 0.4	298		[1972GEI/RAU]
C ₇ H ₄ S ₃	[934-36-1]	4,5-benzo-1,3-dithiole-2-thione					
		$\Delta_{\text{sub}}H$		118.8 ± 0.4	298		[1973RAU/GEI, 1977PED/RYL]
C ₇ H ₅ BrO	[618-32-6]	benzoyl bromide					
		Δ_vH	(320–492)	52.3	335	A	[1987STE/MAL, 1947STU]
C ₇ H ₅ BrO	[1122-91-4]	4-bromobenzaldehyde					
		$\Delta_{\text{fus}}H$		22.6	334.2		[2008FAV/FRE]
C ₇ H ₅ BrO ₂	[88-65-3]	2-bromobenzoic acid					
		$\Delta_{\text{fus}}H$		24.83	421.6	DSC	[2005RIB/FON]
		$\Delta_{\text{sub}}H$	(328–347)	106.8 ± 0.4	338	ME	[2005RIB/FON]
		$\Delta_{\text{sub}}H$	(328–347)	108.5 ± 0.6	298	ME	[2005RIB/FON]
		$\Delta_{\text{sub}}H$		95.9 ± 0.4	298	C	[1994TAN/SAB]
		$\Delta_{\text{sub}}H$		110.9 ± 1.1	298	C	[1987FER/PIL]
C ₇ H ₅ BrO ₂	[585-76-2]	3-bromobenzoic acid					
		$\Delta_{\text{fus}}H$		24.91	430.1	DSC	[2005RIB/FON]
		$\Delta_{\text{sub}}H$	(328–347)	104.2 ± 0.5	338	ME	[2005RIB/FON]
		$\Delta_{\text{sub}}H$	(328–347)	105.9 ± 0.7	298	ME	[2005RIB/FON]
		$\Delta_{\text{sub}}H$		99.2 ± 0.2	298	C	[1994TAN/SAB]
		$\Delta_{\text{sub}}H$		105.0 ± 1.1	298	C	[1987FER/PIL]
C ₇ H ₅ BrO ₂	[586-76-5]	4-bromobenzoic acid					
		$\Delta_{\text{fus}}H$		30.87	526.3	DSC	[2005RIB/FON]
		$\Delta_{\text{sub}}H$	(349–366)	107.4 ± 0.5	358	ME	[2005RIB/FON]
		$\Delta_{\text{sub}}H$	(349–366)	110.1 ± 0.8	358	ME	[2005RIB/FON]
		$\Delta_{\text{sub}}H$		103.1 ± 0.6	298	C	[1994TAN/SAB]
		$\Delta_{\text{sub}}H$		107.6 ± 1.1	298	C	[1987FER/PIL]
C ₇ H ₅ ClO	[98-88-4]	benzoyl chloride					
		Δ_vH	(305–470)	49.6	320	A	[1987STE/MAL, 1947STU, 1999DYK/SVO]
C ₇ H ₅ ClO	[89-98-5]	2-chlorobenzaldehyde					
		Δ_vH	(382–563)	49.8	397	A	[1987STE/MAL, 1999DYK/SVO]
C ₇ H ₅ ClO ₂	[118-91-2]	2-chlorobenzoic acid					
		$\Delta_{\text{fus}}H$		25.25	414	DSC	[2005RIB/FON]
		$\Delta_{\text{fus}}H$		25.73	413.4		[1991ACR]
		$\Delta_{\text{fus}}H$		26.3	414		[1991SAB/HIR]
		$\Delta_{\text{sub}}H$	(320–339)	105.0 ± 0.4	330	ME	[2005RIB/FON]
		$\Delta_{\text{sub}}H$	(320–339)	106.3 ± 0.5	298	ME	[2005RIB/FON]
		$\Delta_{\text{sub}}H$		100.9 ± 0.5	298	C	[1995SAB/AGU]
		$\Delta_{\text{sub}}H$		116.2 ± 0.6		DSC	[1983HOL]
		$\Delta_{\text{sub}}H$		79.5 ± 3.3			[1938WOL/WEG, 1960JON, 1970COX/PIL]
C ₇ H ₅ ClO ₂	[535-80-8]	3-chlorobenzoic acid					
		$\Delta_{\text{fus}}H$		23.67	427.9	DSC	[2005RIB/FON]
		$\Delta_{\text{fus}}H$		23.85	427.4		[1991ACR]
		$\Delta_{\text{fus}}H$		22.0	427.8		[1991SAB/HIR]
		$\Delta_{\text{sub}}H$	(320–340)	101.2 ± 0.4	330	ME	[2005RIB/FON]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{sub}}H$	(320–340)	102.5 ± 0.5	298	ME	[2004RIB/SAN3]
	$\Delta_{\text{sub}}H$		101.4 ± 0.4	298	C	[1995SAB/AGU]
	$\Delta_{\text{sub}}H$		99.6	413	C	[1975ADE/BRO]
	$\Delta_{\text{sub}}H$		105.8	298	C	[1975ADE/BRO]
	$\Delta_{\text{sub}}H$		80.8 ± 3.3			[1938WOL/WEG, 1960JON, 1970COX/PIL]
C₇H₅ClO₂	[74-11-3]	4-chlorobenzoic acid				
	$\Delta_{\text{fus}}H$		30.91	512.5	DSC	[2005RIB/FON]
	$\Delta_{\text{fus}}H$	(80–580)	13.5	512.3	AC	[2002TAN/SUN]
		Note: This value is considerably lower than the other three reported enthalpies of fusion. This value is likely in error.				
	$\Delta_{\text{fus}}H$		32.26	512.9		[1991ACR]
	$\Delta_{\text{fus}}H$		34.26	513.5		[1991SAB/HIR]
	$\Delta_{\text{sub}}H$	(333–356)	103.3 ± 0.5	344	ME	[2005RIB/FON]
	$\Delta_{\text{sub}}H$	(333–356)	105.2 ± 0.7	298	ME	[2005RIB/FON]
	$\Delta_{\text{sub}}H$		102.5 ± 0.4	298	C	[1995SAB/AGU]
	$\Delta_{\text{sub}}H$		101.9	413	C	[1975ADE/BRO]
	$\Delta_{\text{sub}}H$		107.9	298	C	[1975ADE/BRO]
	$\Delta_{\text{sub}}H$		87.9 ± 3.3			[1938WOL/WEG, 1960JON, 1970COX/PIL]
C₇H₅Cl₂N	[622-44-6]	phenylcarbonimidic dichloride				
	Δ_vH	(273–333)	54.0	288	A	[1987STE/MAL, 1973KKY/REP, 1999DYK/SVO]
C₇H₅Cl₂NO₂	[133-90-4]	3-amino-2,5-dichlorobenzoic acid				
	$\Delta_{\text{fus}}H$		37.42	475.6		[1991ACR]
C₇H₅Cl₃	[94-99-5]	1-(chloromethyl)-2,4-dichlorobenzene				
	Δ_vH	(413–578)	54.6	428	A	[1987STE/MAL, 1970DYK/VAN, 1973KKY/REP]
C₇H₅Cl₃	[98-07-7]	(trichloromethyl)benzene				
	$\Delta_{\text{fus}}H$		13.95	236		[1996DOM/HEA]
	$\Delta_{\text{fus}}H$		10.6	270	DSC	[1972AHM/EAD]
		Note: There is a large discrepancy in the two melting points. The 270 K value is correct				
	Δ_vH		57.6			[1995PAP/PIM]
	Δ_vH	(318–487)	52.0	333	A	[1987STE/MAL, 1947STU]
C₇H₅Cl₃	[2077-46-5]	2,3,6-trichlorotoluene				
	Δ_vH	(384–509)	62.2	399	A	[1987STE/MAL, 1973FEL/SAV]
C₇H₅Cl₃N₂O₂	[na]	methyl 4-amino-3,5,6-trichloro-2-picolinate				
	$\Delta_{\text{fus}}H$		26.78	394.3	DSC	[1969PLA/GLA]
C₇H₅FN₂O₄	[17003-70-2]	(fluorodinitromethyl)benzene				
	Δ_vH	(328–363)	52.8	343	A	[1987STE/MAL]
C₇H₅FO₂	[445-29-4]	2-fluorobenzoic acid				
	$\Delta_{\text{sub}}H$	(309–323)	93.9 ± 0.5	316	ME	[2000MON/HIL]
	$\Delta_{\text{sub}}H$		94.4 ± 0.8	298		[2000MON/HIL]
C₇H₅FO₂	[455-38-9]	3-fluorobenzoic acid				
	$\Delta_{\text{sub}}H$	(303–317)	93.3 ± 0.5	310	ME	[2000MON/HIL]
	$\Delta_{\text{sub}}H$		93.6 ± 0.6	298		[2000MON/HIL]
C₇H₅FO₂	[456-22-4]	4-fluorobenzoic acid				
	$\Delta_{\text{fus}}H$		20.9	451.2		[2000KAN/SAM]
	$\Delta_{\text{sub}}H$	(358–382)	91.2 ± 1.3	370	GS	[1969COX/GUN, 1970COX/PIL, 1987STE/MAL]
	$\Delta_{\text{sub}}H$		93.1 ± 3.8	298		[1969COX/GUN, 2000MON/HIL]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₇ H ₅ F ₃	[98-08-8]	(trifluoromethyl)benzene				
	$\Delta_{\text{fus}}H$		13.77	244		[1996DOM/HEA]
	$\Delta_{\text{fus}}H$		11.99	242	DSC	[1972AHM/EAD]
	$\Delta_{\text{sub}}H$	(222–233)	54.4	227	MG	[1948SEA/HOP]
	Δ_vH	(328–413)	35.6	343		[1999DYK/SVO]
	Δ_vH	(468–532)	31.6	483		[1999DYK/SVO]
	Δ_vH	(323–384)	35.9	338	I	[1992JAD/FRA]
	Δ_vH	(460–530)	32.4	475		[1985MOU]
	Δ_vH	(330–410)	37.1	298		[1984BOU/FRI, 1991BAS/SVO]
	Δ_vH		35.4 ± 0.1	334	C	[1959SCO/DOU]
	Δ_vH		34.1 ± 0.1	353	C	[1959SCO/DOU]
	Δ_vH		32.6 ± 0.1	375	C	[1959SCO/DOU]
	Δ_vH	(328–413)	35.7	343	A,EB	[1987STE/MAL, 1951POT/SAY, 1970DYK/VAN, 1959SCO/DOU]
Δ_vH	(241–375)	39.1	256		[1947STU]	
Δ_vH	(275–353)	38.5	290		[1946FIE/SAY]	
C ₇ H ₅ F ₄ NO ₂	[27827-91-4]	1,1,3-trihydrotetrafluoropropyl α -cyanoacrylate				
	$\Delta_{\text{trs}}H$		0.3	154		
	$\Delta_{\text{fus}}H$		19.95	287.4		[1995LEB/BYK2]
C ₇ H ₅ F ₁₀ NS	[na]	2,2,2-trifluoro-N-[1,2,2,2-tetrafluoro-1-(trifluoromethyl)ethyl]ethanimidithioic acid, ethyl ester				
	Δ_vH		30.1	394		[1975PET/SHR]
C ₇ H ₅ F ₁₁ O	[181214-75-5]	1-ethoxy-1,1,2,2,3,3,4,4,5,5,5-undecafluoropentane				
	Δ_vH	(288–373)	39.0	303	I	[2002MUR/YAM]
C ₇ H ₅ F ₁₁ O	[203783-57-7]	1-ethoxy-1,1,2,2,3,4,4,4-octafluoro-3-(trifluoromethyl)butane				
	Δ_vH	(288–373)	38.3	303	I	[2002MUR/YAM]
C ₇ H ₅ IO ₂	[88-67-5]	2-iodobenzoic acid				
	$\Delta_{\text{fus}}H$		21.38	435.1		[1994TAN/SAB2]
	$\Delta_{\text{sub}}H$	(345–359)	111.4 ± 0.8	352	ME	[2000MON/HIL]
	$\Delta_{\text{sub}}H$		112.8 ± 2.0	298		[2000MON/HIL]
	$\Delta_{\text{sub}}H$		92.6 ± 0.2	298	C	[1994ZHI/SAB, 1995SAB/AGU, 1994TAN/SAB2]
$\Delta_{\text{sub}}H$		103.0 ± 0.4	298	DSC	[1983HOL]	
C ₇ H ₅ IO ₂	[618-51-9]	3-iodobenzoic acid				
	$\Delta_{\text{fus}}H$		28.7	460.4		[1994TAN/SAB2]
	$\Delta_{\text{sub}}H$	(347–363)	109.6 ± 0.5	355	ME	[2000MON/HIL]
	$\Delta_{\text{sub}}H$		111.1 ± 1.9	298		[2000MON/HIL]
$\Delta_{\text{sub}}H$		96.4 ± 0.3	298	C	[1994ZHI/SAB, 1995SAB/AGU, 1994TAN/SAB2]	
C ₇ H ₅ IO ₂	[619-58-9]	4-iodobenzoic acid				
	$\Delta_{\text{fus}}H$		35.24	543.8		[1994TAN/SAB2]
	$\Delta_{\text{sub}}H$	(363–379)	111.0 ± 0.4	372	ME	[2000MON/HIL]
	$\Delta_{\text{sub}}H$		112.9 ± 2.5	298		[2000MON/HIL]
$\Delta_{\text{sub}}H$		99.3 ± 0.4	298	C	[1994ZHI/SAB, 1995SAB/AGU, 1994TAN/SAB2]	
C ₇ H ₅ N	[100-47-0]	benzotrile				
	$\Delta_{\text{fus}}H$		10.98	260.3		[1985LEB/BYK]
	Δ_vH	(301–464)	49.1	316	A	[1987STE/MAL, 1947STU]
C ₇ H ₅ N	[931-54-4]	phenyl isocyanide				
	Δ_vH	(285–438)	46.2	300	A	[1987STE/MAL, 1947STU]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₇ H ₅ NO	[273-53-0]	benzoxazole				
	$\Delta_{\text{fus}}H$		0.02	247		
	$\Delta_{\text{fus}}H$		16.78	302.5		[1996DOM/HEA]
	$\Delta_{\text{sub}}H$		69.5 ± 0.4	298	C	[1998SAB/HEV]
	Δ_vH		51.2	320	EB	[1992STE/CHI2]
	Δ_vH		48.6	360	EB	[1992STE/CHI2]
	Δ_vH		46.1	400	EB	[1992STE/CHI2]
	Δ_vH		43.5	440	EB	[1992STE/CHI2]
C ₇ H ₅ NO	[103-71-9]	phenyl isocyanate				
	Δ_vH	(329–445)	46.5 ± 0.3	298	EB	[1996STE/CHI3]
C ₇ H ₅ NO						
	Δ_vH	(283–439)	45.0	298	A	[1987STE/MAL, 1947STU]
C ₇ H ₅ NO	[7187-01-1]	2-furanacrylonitrile				
	Δ_vH		65.2 ± 0.6	298	C	[2009RIB/AMA2]
C ₇ H ₅ NO	[271-58-9]	benz[a]isoxazole (anthranil)				
	Δ_vH		55.3 ± 0.3	298	C	[2004MAT/MIR3]
C ₇ H ₅ NOS	[2382-96-9]	2-mercaptobenzoxazole				
	$\Delta_{\text{fus}}H$		25.0	470	DSC	[2008TEM/ROU3]
	$\Delta_{\text{fus}}H$		22.6 ± 0.9	468.3	DSC	[2008MEN/FLO]
C ₇ H ₅ NO ₂	[59-49-4]	2-benzoxazolinone				
	$\Delta_{\text{sub}}H$		97.6 ± 2.2	298	C	[2006MOR/MIR2]
C ₇ H ₅ NO ₃	[552-89-6]	2-nitrobenzaldehyde				
	Δ_vH	(390–547)	58.7	405	A	[1987STE/MAL]
	Δ_vH	(359–547)	59.5	373		[1947STU]
C ₇ H ₅ NO ₃	[99-61-6]	3-nitrobenzaldehyde				
	Δ_vH	(401–552)	62.0	416	A	[1987STE/MAL]
C ₇ H ₅ NO ₃ S	[81-07-2]	1,1-dioxo-1,2-benzisothiazol-3(2H)-one (saccharin)				
	$\Delta_{\text{fus}}H$		32.1	502.9	DSC	[2009GOO/ROD]
	$\Delta_{\text{fus}}H$		26.77	502.7	DSC	[2008BAS/BOS]
	$\Delta_{\text{fus}}H$		29.89	500.7	DSC	[2005MAT/MIR]
	$\Delta_{\text{sub}}H$		112.6 ± 4.2	298	C	[2005MAT/MIR]
C ₇ H ₅ NO ₄	[552-16-9]	2-nitrobenzoic acid				
	$\Delta_{\text{fus}}H$		27.99	419		[1996DOM/HEA]
	$\Delta_{\text{sub}}H$	(346–356)	115.8 ± 0.5	356	ME	[1999RIB/MAT]
	$\Delta_{\text{sub}}H$		118.7 ± 0.5	298	ME	[1999RIB/MAT]
C ₇ H ₅ NO ₄	[121-92-6]	3-nitrobenzoic acid				
	$\Delta_{\text{fus}}H$		19.33	414.3		[1996DOM/HEA]
	$\Delta_{\text{sub}}H$	(347–361)	107.2 ± 0.4	354	ME	[1999RIB/MAT]
	$\Delta_{\text{sub}}H$		110.0 ± 0.4	298	ME	[1999RIB/MAT]
C ₇ H ₅ NO ₄	[62-23-7]	4-nitrobenzoic acid				
	$\Delta_{\text{fus}}H$		36.9	512.4		[1996DOM/HEA]
	$\Delta_{\text{sub}}H$	(367–381)	115.4 ± 0.6	374	ME	[1999RIB/MAT]
	$\Delta_{\text{sub}}H$		119.7 ± 0.6	298	ME	[1999RIB/MAT]
C ₇ H ₅ NO ₄	[100-26-5]	pyridine-2,5-dicarboxylic acid				
	$\Delta_{\text{sub}}H$		163.6 ± 2.7	298	C	[2005MAT/MOR]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₇ H ₅ NO ₄	[499-83-2]	pyridine-2,6-dicarboxylic acid				
	$\Delta_{\text{sub}}H$		137.1 ± 5.7	298	C	[2005MAT/MOR]
C ₇ H ₅ NO ₄	[1874-22-2]	3-(5-nitro-2-furyl)-2-propenal				
	$\Delta_{\text{sub}}H$		97.9 ± 2.1			[1980BAL/LEB, 1986PED/NAY]
C ₇ H ₅ NO ₄	[2620-44-2]	5-nitro-1,3-benzodioxole				
	$\Delta_{\text{fus}}H$		28.2	420.2	DSC	[2007MAT/SOU]
	$\Delta_{\text{sub}}H$		97.4 ± 2.2	298	C	[2007MAT/SOU]
C ₇ H ₅ NS	[95-16-9]	benzothiazole				
	$\Delta_{\text{fus}}H$		11.95	275.5	DTA	[1998SAB/HEV]
	$\Delta_{\text{fus}}H$		12.8	275.6		[1996DOM/HEA]
	$\Delta_{\text{sub}}H$		72.9 ± 0.6	298	B	[1998SAB/HEV]
	Δ_vH		58.7	320	EB	[1992STE/CHI2]
	Δ_vH		56.0	360	EB	[1992STE/CHI2]
	Δ_vH		53.5	400	EB	[1992STE/CHI2]
	Δ_vH		50.9	440	EB	[1992STE/CHI2]
	Δ_vH		48.4	480	EB	[1992STE/CHI2]
	Δ_vH		45.7	520	EB	[1992STE/CHI2]
C ₇ H ₅ NS	[103-72-0]	phenyl isothiocyanate				
	Δ_vH	(320–492)	52.6	335	A	[1987STE/MAL, 1947STU]
C ₇ H ₅ NS ₂	[149-30-4]	2-mercaptobenzothiazole				
	$\Delta_{\text{fus}}H$		22.3 ± 0.2	455.9	DSC	[2008TEM/ROU3]
	$\Delta_{\text{fus}}H$		20.56	453.5	DSC	[2008MEN/FLO]
C ₇ H ₅ N ₃ O ₆	[610-25-3]	2,4,5-trinitrotoluene				
	$\Delta_{\text{fus}}H$		24.7	376.2		[1996DOM/HEA]
C ₇ H ₅ N ₃ O ₆	[118-96-7]	2,4,6-trinitrotoluene				
	$\Delta_{\text{fus}}H$		23.43	352.2		[1993ACR]
	$\Delta_{\text{sub}}H$		104.2		DSC	[1990HWA/YOS]
	$\Delta_{\text{sub}}H$	(293–353)	112.4	308	A	[1987STE/MAL]
	$\Delta_{\text{sub}}H$	(301–349)	113.2 ± 1.5	298	ME	[1979KUD/KUD2]
	$\Delta_{\text{sub}}H$	(297–330)	99.2 ± 2		GS	[1976PEL, 1977PEL]
	$\Delta_{\text{sub}}H$		104.6 ± 1.7	298	ME	[1971LEN/VEL]
	$\Delta_{\text{sub}}H$	(327–349)	103.3 ± 2.5	338		[1970LEN/VEL]
	$\Delta_{\text{sub}}H$		U 112-132		TGA	[1970MAY/VEN, 1978CUN/PAL]
	$\Delta_{\text{sub}}H$	(323–353)	118.4 ± 4.2		ME	[1950EDW, 1960JON, 1970COX/PIL]
	$\Delta_{\text{sub}}H$		102.2			[1950NIT/SEK]
	Δ_vH		80.8		DSC	[1990HWA/YOS]
	Δ_vH	(353–523)	93.7	368	A	[1987STE/MAL]
	Δ_vH		87.0 ± 1.9	298	ME	[1978CUN/PAL]
C ₇ H ₅ N ₃ O ₇	[606-35-9]	2,4,6-trinitroanisole				
	$\Delta_{\text{sub}}H$	(334–342)	132.4	338	A	[1987STE/MAL]
	$\Delta_{\text{sub}}H$		133.1 ± 2.1			[1950NIT/SEK, 1970COX/PIL]
	Δ_vH	(342–363)	91.9	352	A	[1987STE/MAL]
C ₇ H ₅ N ₃ O ₇	[602-99-3]	3-hydroxy-2,4,6-trinitrotoluene				
	$\Delta_{\text{sub}}H$	(310–365)	111.2 ± 2.1	298		[1978CUN/PAL]
	$\Delta_{\text{sub}}H$	(325–350)	103.3	337		[1970LEN/VEL]
	$\Delta_{\text{sub}}H$		104.6	298		[1970LEN/VEL]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₇ H ₅ N ₅ O ₈	[479-45-8]	2,4,6-N-tetranitro-N-methylaniline				
	$\Delta_{\text{fus}}H$		25.86	402.6		[1996DOM/HEA]
	$\Delta_{\text{sub}}H$		133.1		DSC	[1990HWA/YOS]
	$\Delta_{\text{sub}}H$	(335–416)	133.8 ± 1.6	298	ME	[1978CUN/PAL]
C ₇ H ₆ ClF	[443-83-4]	1-chloro-3-fluoro-2-methylbenzene				
	$\Delta_{\text{fus}}H$		12.6	246	DSC	[1972AHM/EAD]
	Δ_vH		108.4		DSC	[1990HWA/YOS]
C ₇ H ₆ ClNO ₂	[612-23-7]	1-(chloromethyl)-2-nitrobenzene				
	$\Delta_{\text{sub}}H$		96.2 ± 3.5		ME	[2005HOS/NAG]
C ₇ H ₆ Cl ₂	[98-87-3]	(dichloromethyl)benzene				
	Δ_vH	(308–487)	49.5	323	A	[1987STE/MAL, 1947STU]
C ₇ H ₆ Cl ₂	[95-73-8]	2,4-dichlorotoluene				
	Δ_vH	(346–475)	50.6	361	A	[1987STE/MAL, 1973FEL/SAV, 1999DYK/SVO]
C ₇ H ₆ Cl ₂	[118-69-4]	2,6-dichlorotoluene				
	$\Delta_{\text{fus}}H$		10.7	272	DSC	[1972AHM/EAD]
C ₇ H ₆ Cl ₂	[95-75-0]	3,4-dichlorotoluene				
	Δ_vH	(378–543)	49.4	393	A	[1987STE/MAL, 1970DYK/VAN, 1999DYK/SVO]
C ₇ H ₆ Cl ₂ O	[1984-59-4]	2,3-dichloroanisole				
	$\Delta_{\text{fus}}H$		22.04	304.1	DSC	[2008RIB/FER]
	$\Delta_{\text{sub}}H$		83.6 ± 1.5	298	C	[2008RIB/FER]
C ₇ H ₆ Cl ₂ O	[33719-74-3]	3,5-dichloroanisole				
	$\Delta_{\text{fus}}H$		23.68	310.5	DSC	[2008RIB/FER]
	$\Delta_{\text{sub}}H$		79.0 ± 1.5	298	C	[2008RIB/FER]
C ₇ H ₆ Cl ₃ NO ₂	[77765-42-5]	2,2,4-trichloro-5-(dimethylamino)-4-cyclopentene-1,3-dione				
	Δ_vH	(453–483)	70.9	468	GC	[1980SHA/SAD]
C ₇ H ₆ F ₃ N	[98-16-8]	1-(trifluoromethyl)-3-aminobenzene				
	Δ_vH	(334–464)	53.1	349	A	[1987STE/MAL, 1953KAR/SAY, 1999DYK/SVO]
C ₇ H ₆ F ₃ NS	[na]	N-(trifluoromethyl)thioaniline				
	Δ_vH	(333–413)	47.0	348	A	[1987STE/MAL]
C ₇ H ₆ F ₆ O ₄	[na]	dimethylperfluoroglutarate				
	Δ_vH		52.3	298	EB	[1976KOL/SLA]
C ₇ H ₆ F ₈ O ₃	[na]	<i>bis</i> -(tetrafluoropropyl)carbonate				
	$\Delta_{\text{fus}}H$		41.05	253.4		[1996DOM/HEA]
C ₇ H ₆ INO ₂	[6277-17-4]	2-iodo-3-nitrotoluene				
	$\Delta_{\text{fus}}H$	(79–373)	20.68	339.3	AC	[2000DI/LI]
C ₇ H ₆ N ₂	[51-17-2]	benzimidazole				
	$\Delta_{\text{fus}}H$		0.71	384.4		
	$\Delta_{\text{fus}}H$		20.47	445.5		[2002DOM/KOZ]
	$\Delta_{\text{fus}}H$		19.25	443.2		[1984DOM/EVA]
	$\Delta_{\text{sub}}H$		90.2 ± 0.6	363	C	[1998SAB/HEV2]
	$\Delta_{\text{sub}}H$		94.3 ± 0.6	298		[1998SAB/HEV2]
	$\Delta_{\text{sub}}H$	(340–359)	101.8 ± 0.4	350	ME	[1987JIM/ROU]
	$\Delta_{\text{sub}}H$		102.2 ± 0.4	298	ME	[1987JIM/ROU]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{sub}}H$		98.9 ± 0.4	298		[1986JIM/ROU]
C ₇ H ₆ N ₂	[271-44-3]	indazole				
	$\Delta_{\text{sub}}H$	(308–317)	90.9 ± 0.2	318	ME	[1987JIM/ROU]
	$\Delta_{\text{sub}}H$		91.1 ± 0.2	298		[1987JIM/ROU, 1986JIM/ROU]
	$\Delta_{\text{sub}}H$		87.7 ± 0.9			[1985SKI/PIL]
	$\Delta_{\text{sub}}H$		97.1			[1961ZIM/GEI]
C ₇ H ₆ N ₂ O	[615-16-7]	1,3-dihydro-2 <i>H</i> -benzimidazol-2-one				
	$\Delta_{\text{sub}}H$		126.4 ± 2.4	298	C	[2006MOR/MIR2]
C ₇ H ₆ N ₂ O	[7364-25-2]	1,2-dihydro-3 <i>H</i> -indazol-3-one				
	$\Delta_{\text{sub}}H$		127.6 ± 1.5	298	C	[2006MOR/MIR2]
C ₇ H ₆ N ₂ O ₂	[4413-48-3]	5-methoxybenzofurazan				
	$\Delta_{\text{sub}}H$		89.2 ± 0.7	298	C	[1996ACR/BOT]
C ₇ H ₆ N ₂ O ₂	[19164-41-1]	5-methylbenzofurazan-1-oxide				
	$\Delta_{\text{sub}}H$		92.2 ± 1.2	298	C	[1996ACR/BOT]
C ₇ H ₆ N ₂ O ₃	[7791-49-3]	5-methoxybenzofurazan-1-oxide				
	$\Delta_{\text{sub}}H$		96.0 ± 1.6	298	C	[1996ACR/BOT]
C ₇ H ₆ N ₂ O ₃	[6635-41-2]	2-nitrobenzaloxime				
	$\Delta_{\text{sub}}H$ (<i>anti</i>)		U 26.4 ± 1.7		MS	[1983MAJ/AZZ]
	$\Delta_{\text{sub}}H$ (<i>syn</i>)		U 40.2 ± 1.7		MS	[1983MAJ/AZZ]
C ₇ H ₆ N ₂ O ₃	[3431-62-7]	3-nitrobenzaloxime				
	$\Delta_{\text{sub}}H$ (<i>anti</i>)		U 41.0 ± 1.7		MS	[1983MAJ/AZZ]
	$\Delta_{\text{sub}}H$ (<i>syn</i>)		U 42.7 ± 1.7		MS	[1983MAJ/AZZ]
C ₇ H ₆ N ₂ O ₃	[1129-37-9]	4-nitrobenzaloxime				
	$\Delta_{\text{sub}}H$ (<i>anti</i>)		U 56.4 ± 1.7		MS	[1983MAJ/AZZ]
C ₇ H ₆ N ₂ O ₄	[25321-14-6]	1,1-dinitrophenylmethane				
	$\Delta_{\text{sub}}H$	(312–323)	76.1 ± 0.8		ME	[1972PEP/MAT]
C ₇ H ₆ N ₂ O ₄	[602-01-7]	2,3-dinitrotoluene				
	$\Delta_{\text{fus}}H$		17.57	329.8		[1993ACR]
	$\Delta_{\text{sub}}H$	(270–315)	97.0 ± 2.1		ME	[2008FRE/KEB]
C ₇ H ₆ N ₂ O ₄	[121-14-2]	2,4-dinitrotoluene				
	$\Delta_{\text{fus}}H$		20.12	343.3		[1991ACR]
	$\Delta_{\text{sub}}H$	(270–315)	94.7 ± 2.3		ME	[2008FRE/KEB]
	$\Delta_{\text{sub}}H$		94.2 ± 2.5		GS	[2008FRE/KEB, 2001RIT]
	$\Delta_{\text{sub}}H$	(332–342)	98.3 ± 2.5	337	ME	[1977PED/RYL, 1970LEN/VEL]
	$\Delta_{\text{sub}}H$		99.6 ± 2.5	298		[1977PED/RYL, 1970LEN/VEL]
	$\Delta_{\text{sub}}H$	(277–344)	95.8 ± 1.25	310	GS	[1976PEL, 1977PEL]
	$\Delta_{\text{sub}}H$		99.6 ± 1.3		ME	[1970COX/PIL, 1971LEN/VEL]
	Δ_vH	(344–572)	76.9	359	A	[1987STE/MAL]
	Δ_vH	(473–572)	58.2	488		[1987STE/MAL, 1968MAK, 1973KKY/REP]
Δ_vH	(354–439)	70.2			[1977PEL, 1958MOL]	
C ₇ H ₆ N ₂ O ₄	[606-20-2]	2,6-dinitrotoluene				
	$\Delta_{\text{fus}}H$		23.85	327.5		[1993ACR]
	$\Delta_{\text{sub}}H$	(275–325)	99.6 ± 2.3		ME	[2008FRE/KEB]
$\Delta_{\text{sub}}H$	(277–323)	98.3 ± 0.8	300	GS	[1976PEL, 1977PEL]	

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(330–533)	77.8	345	A	[1987STE/MAL]
	$\Delta_v H$	(423–553)	56.9	438	A	[1987STE/MAL, 1968MAK]
	$\Delta_v H$	(344–427)	68.7			[1977PEL, 1958MOL]
C ₇ H ₆ N ₂ O ₄	[610-39-9]	3,4-dinitrotoluene				
	$\Delta_{\text{fus}}H$		18.83	329.5		[1993ACR]
	$\Delta_{\text{sub}}H$	(270–315)	99.6 ± 1.9		ME	[2008FRE/KEB]
C ₇ H ₆ N ₂ O ₄	[618-85-9]	3,5-dinitrotoluene				
	$\Delta_v H$	(493–543)	62.6	508	A	[1987STE/MAL, 1968MAK]
C ₇ H ₆ N ₂ O ₄	[611-38-1]	(dinitromethyl)benzene				
	$\Delta_{\text{sub}}H$	(312–323)	76.1	317.5	A	[1987STE/MAL]
C ₇ H ₆ N ₂ O ₅	[497-56-3]	3,5-dinitro- <i>o</i> -cresol				
	$\Delta_{\text{sub}}H$	(290–324)	103.3		TE	[1947BAL, 1960JON]
C ₇ H ₆ N ₂ O ₅	[534-52-1]	2-methyl-4,6-dinitrophenol				
	$\Delta_{\text{fus}}H$		19.41	359.3		[1991ACR]
C ₇ H ₆ N ₂ S	[583-39-1]	2-mercaptobenzimidazole				
	$\Delta_{\text{fus}}H$		24.11	589.4	DSC	[2008MEN/FLO]
C ₇ H ₆ O	[100-52-7]	benzaldehyde				
	$\Delta_{\text{fus}}H$		9.33	216		[1996DOM/HEA]
	$\Delta_v H$	(278–313)	49.0 ± 0.7	298	GS	[2007EME/DAB]
	$\Delta_v H$	(313–353)	49.1	298	CGC	[1995CHI/HOS]
	$\Delta_v H$	(348–452)	49.5	363	A	[1987STE/MAL]
	$\Delta_v H$	(409–481)	43.8	424	A	[1987STE/MAL]
	$\Delta_v H$	(311–376)	48.6	326	A	[1987STE/MAL]
	$\Delta_v H$	(370–475)	45.5	385	A	[1987STE/MAL]
	$\Delta_v H$	(465–541)	41.9	480	A	[1987STE/MAL]
	$\Delta_v H$	(529–599)	40.6	544	A	[1987STE/MAL]
	$\Delta_v H$	(311–404)	50.3	298	EB	[1975AMB/CON]
	$\Delta_v H$	(311–404)	42.5	452	EB	[1975AMB/CON]
	$\Delta_v H$	(273–373)	47.0	288	A,BG	[1987STE/MAL, 1973DEM/LEH]
	$\Delta_v H$	(299–452)	54.4	314		[1947STU]
C ₇ H ₆ O	[539-80-0]	2,4,6-cycloheptatrienone (tropone)				
	$\Delta_v H$	(273–323)	54.2	288	A	[1987STE/MAL]
C ₇ H ₆ O ₂	[65-85-0]	benzoic acid				
	$\Delta_{\text{fus}}H$		16.99	396.9	DSC	[2009BR12]
	$\Delta_{\text{fus}}H$		17.3	394.4	DSC	[2003SHA/KAN, 2004SHA/JAM]
	$\Delta_{\text{fus}}H$		17.1	395.4	DSC	[2002ROY/RIG]
	$\Delta_{\text{fus}}H$		17.99	395.5		[1984DOM/EVA]
	$\Delta_{\text{sub}}H$	(340–410)	90.9 ± 2.0		TG-TS	[2009SEL/RAG]
	$\Delta_{\text{sub}}H$		93.3 ± 1.2	298	QCM	[2008FRE/KEB]
	$\Delta_{\text{sub}}H$	(299–317)	90.0 ± 0.5	307	ME	[2006RIB/MON]
	$\Delta_{\text{sub}}H$	(299–317)	90.4 ± 0.5	298	ME	[2006RIB/MON]
	$\Delta_{\text{sub}}H$		91.7 ± 3.4		ME	[2005HOS/NAG]
	$\Delta_{\text{sub}}H$		88.3 ± 0.5	298	C	[2001KIY/MIN]
	$\Delta_{\text{sub}}H$	(323–394)	90.5 ± 0.3		GS	[1999ZIE/PER]
	$\Delta_{\text{sub}}H$		89 ± 6		TGA	[1999PRI/BAS]
	$\Delta_{\text{sub}}H$		87.5 ± 0.4			[1998PRI/HAW]
	$\Delta_{\text{sub}}H$	(313–343)	86.7		TGA	[1997ELD]
	$\Delta_{\text{sub}}H$	(307–314)	88.7 ± 0.9	311	ME	[1990RIB/MON]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
		$\Delta_{\text{sub}}H$	89.3 ± 0.9	298		[1990RIB/MON]
		$\Delta_{\text{sub}}H$	87.5 ± 0.3	335	C	[1988TOR/BAR]
		$\Delta_{\text{sub}}H$	89.2 ± 1.0	298		[1988TOR/BAR]
		$\Delta_{\text{sub}}H$	95.1 ± 1.8	294		[1985KAI/HAD]
		$\Delta_{\text{sub}}H$	(293–319) 90.8 ± 0.6	306	QR	[1985GLU/ARK]
		$\Delta_{\text{sub}}H$	(368–428) 87.8		GS	[1985MAT/KUW2]
		$\Delta_{\text{sub}}H$	89.5 ± 0.4		DSC	[1983HOL]
		$\Delta_{\text{sub}}H$	(320–370) 89.1 ± 0.2		C	[1982MUR/SAK]
		$\Delta_{\text{sub}}H$	(316–391) 89.5 ± .05	353	DM	[1982DEK/BLO]
		$\Delta_{\text{sub}}H$	(293–313) 90.6 ± 2		ME	[1982COL/JIM]
		$\Delta_{\text{sub}}H$	93.45 ± 1		GS	[1981BRO/MCE]
		$\Delta_{\text{sub}}H$	(328–398) U 133.5 ± 4.5		C	[1980MUR/CAV]
		$\Delta_{\text{sub}}H$	(344–395) 85 ± 2	369	SG	[1980SAC/HIL]
		$\Delta_{\text{sub}}H$	(281–323) 88.3 ± 2.9		LE	[1978NOW/SZC]
		$\Delta_{\text{sub}}H$	88.5 ± 0.8		C	[1976MIR/LEB]
		$\Delta_{\text{sub}}H$	(294–331) 92.5 ± 4		ME	[1975VAN/DEK]
		$\Delta_{\text{sub}}H$	(293–318) 88.5 ± 1.6		TE	[1975DEK/VAN]
		$\Delta_{\text{sub}}H$	(273–318) 92.9 ± 0.2	296	ME	[1974ARS]
		$\Delta_{\text{sub}}H$	(293–311) 88.1 ± 0.2		TCM	[1973DEK/OON]
		$\Delta_{\text{sub}}H$	(338–383) 89.0 ± 0.4		ME	[1973MAL/GIG]
		$\Delta_{\text{sub}}H$	(338–383) 89.3 ± 0.4		C	[1973MAL/GIG]
		$\Delta_{\text{sub}}H$	(290–315) 86.6 ± 1.3		ME,C	[1972WIE]
		$\Delta_{\text{sub}}H$	(293–308) 90. ± 0.3		ME	[1972COL/MON]
		$\Delta_{\text{sub}}H$	89.5 ± 0.2	298	C	[1972MOR, 1971BEE/LIN]
		$\Delta_{\text{sub}}H$	(299–329) 89.1	314		[1971ASH]
		$\Delta_{\text{sub}}H$	(290–315) 86.6 ± 1.7	303	ME	[1970WIE/WAU, 1999ZIE/PER]
		$\Delta_{\text{sub}}H$	(324–392) 90.4 ± 0.8	367	HSA	[1970MEL/MER]
		$\Delta_{\text{sub}}H$	89.7 ± 0.6	298	C	[1969CHA/STE]
		$\Delta_{\text{sub}}H$	(348–378) 88.9 ± 0.5	363	GS	[1968MER]
		$\Delta_{\text{sub}}H$	(291–307) 90.9	299	ME	[1965DAV/KYB]
		$\Delta_{\text{sub}}H$	(243–387) 91.5 ± 0.5	298	GS	[1954DAV/JON, 1970COX/PIL, 1960JON]
		$\Delta_{\text{sub}}H$	84.2 ± 0.8	318	TE	[1938WOL/WEG]
		$\Delta_{\text{sub}}H$	(333–389) 85.8	383	T	[1934HIR]
		$\Delta_{\text{sub}}H$	(377–394) 84.5 ± 0.5	364	I	[1927KLO/WOO]
		Δ_vH	(401–416) 63.3 ± 0.6			[2003PEN/RIB]
		Δ_vH	(353–393) 78.9	298	CGC	[1995CHI/HOS]
		Δ_vH	(368–428) 67.8		GS	[1985MAT/KUW2]
		Δ_vH	(405–523) 66.3	420	A	[1987STE/MAL]
		Δ_vH	65.4	428	I	[1943CRA]
		Δ_vH	(401–520) 67.7	416	MM,A	[1927KLO/WOO]
C₇H₆O₂	[90-02-8]	2-hydroxybenzaldehyde				
	$\Delta_{\text{fus}}H$		13.3	278.7	DSC	[2008BER/MIN]
		Note: Authors noted in the paper that their melting point temperature differed significantly from published literature values				
		Δ_vH	53.3 ± 0.3	298	C	[2008BER/MIN]
		Δ_vH	50.4 ± 1.3	298	C	[2007RIB/ARA]
		Δ_vH	(383–470) 30.6	398	A	[1987STE/MAL]
		Δ_vH	47.7			[1986BAL/GNA]
		Δ_vH	(306–470) 49.6	321		[1947STU]
C₇H₆O₂	[100-83-4]	3-hydroxybenzaldehyde				
	$\Delta_{\text{sub}}H$	(312–330) 99.7 ± 0.6		321	ME	[2010RIB/GON]
	$\Delta_{\text{sub}}H$	(312–330) 100.1 ± 0.6		298	ME	[2010RIB/GON]
C₇H₆O₂	[123-08-0]	4-hydroxybenzaldehyde				

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
		$\Delta_{\text{fus}}H$	20.3	390.8	DSC	[2008BER/MIN]
		$\Delta_{\text{fus}}H$	21.6	390.8		[2008TEM/ROU]
		$\Delta_{\text{sub}}H$	(324–341) 101.8 ± 0.5	333	ME	[2010RIB/GON]
		$\Delta_{\text{sub}}H$	(324–341) 102.5 ± 0.5	298	ME	[2010RIB/GON]
		$\Delta_{\text{sub}}H$	99.7 ± 0.4	298	C	[2008BER/MIN]
		$\Delta_{\text{sub}}H$	(303–336) 98.2 ± 1.3	298		[1987STE/MAL, 1971PAR/ROC]
		$\Delta_{\text{sub}}H$	(312–336) 91.2	324		[1960AIH]
		Δ_vH	(394–583) 72.3	409	A	[1987STE/MAL, 1947STU]
C ₇ H ₆ O ₂	[1864-94-4]	phenylformate				
		Δ_vH	(287–305) 52.9 ± 0.6	298	BG	[1976ANT/CAR, 1975ANT/CAR]
C ₇ H ₆ O ₂	[274-09-9]	1,3-benzodioxole				
		Δ_vH	41.4			[1958CAS/FLE2]
C ₇ H ₆ O ₂	[533-75-5]	tropolone				
		$\Delta_{\text{sub}}H$	(273–333) 84.1 ± .4		ME	[1971JAC/HUN]
		$\Delta_{\text{sub}}H$	83.7 ± 0.8	298		[1951NIC, 1970COX/PIL]
C ₇ H ₆ O ₂	[623-30-3]	3-(2-furyl)-2-propenal				
		$\Delta_{\text{sub}}H$	76.1 ± 2.1			[1980BAL/LEB, 1986PED/NAY]
C ₇ H ₆ O ₃	[539-47-9]	2-furanacrylic acid				
		$\Delta_{\text{sub}}H$	(322–338) 101.7 ± 0.5	298	ME	[2009RIB/AMA2]
		$\Delta_{\text{sub}}H$	103.0 ± 0.7	298	C	[2009RIB/AMA2]
C ₇ H ₆ O ₃	[81311-95-7]	3-furanacrylic acid				
		$\Delta_{\text{sub}}H$	(327–343) 105.0 ± 0.5	298	ME	[2009RIB/AMA2]
		$\Delta_{\text{sub}}H$	104.9 ± 1.1	298	C	[2009RIB/AMA2]
C ₇ H ₆ O ₃	[69-72-7]	2-hydroxybenzoic acid (salicylic acid)				
		$\Delta_{\text{fus}}H$	27.1	434.1	DSC	[2009GOO/ROD]
		$\Delta_{\text{fus}}H$	23.05	432.5	DSC	[2009PEN/ESC]
		$\Delta_{\text{fus}}H$	26.1	432.4		
		$\Delta_{\text{fus}}H$	24.45	431.1		[2005PIN/DIO, 2008MOT/QUE]
		$\Delta_{\text{fus}}H$	24.6	NA	DSC	[2003SHA/KAN]
		$\Delta_{\text{fus}}H$	24.6	431.8		[1996DOM/HEA, 1993SAB/LE]
		$\Delta_{\text{sub}}H$	94.4 ± 0.4	298	C	[2005PIN/DIO]
		$\Delta_{\text{sub}}H$	95.1 ± 0.5	333	C	[1993SAB/LE]
		$\Delta_{\text{sub}}H$	96.3 ± 0.5	298		[1993SAB/LE]
		$\Delta_{\text{sub}}H$	(307–324) 95.7 ± 0.8	315	ME	[1980COL/JIM, 1981COL/JIM]
		$\Delta_{\text{sub}}H$	(312–332) 94.9 ± 0.4	322	TE	[1977DEK/VAN]
		$\Delta_{\text{sub}}H$	(312–332) 93.22 ± 0.8	322	ME	[1977DEK/VAN]
		$\Delta_{\text{sub}}H$	(298–328) 99.2 ± 2	313	ME	[1974ARS]
		$\Delta_{\text{sub}}H$	(368–408) 94.8 ± 0.4			[1973MAL/GIG]
		$\Delta_{\text{sub}}H$	(368–408) 95.1 ± 0.4		GS	[1954DAV/JON, 1970COX/PIL, 1960JON]
		Δ_vH	66.7			[2002CHA/DOL]
		Δ_vH	(445–504) 79.4	460	A	[1987STE/MAL]
C ₇ H ₆ O ₃	[99-06-9]	3-hydroxybenzoic acid				
		$\Delta_{\text{fus}}H$	36.5	476.4		[2005PIN/DIO]
		$\Delta_{\text{fus}}H$	26.2	475.1		[1996DOM/HEA, 1993SAB/LE]
		$\Delta_{\text{sub}}H$	118.3 ± 1.1	298	C	[2005PIN/DIO]
		$\Delta_{\text{sub}}H$	123.5 ± 0.74	363	C	[1993SAB/LE]
		$\Delta_{\text{sub}}H$	125.0 ± 0.74	298		[1993SAB/LE]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound					
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference	
		$\Delta_v H$	80.1		TGA	[2002CHA/DOL]	
C ₇ H ₆ O ₃	[99-96-7]	4-hydroxybenzoic acid					
		$\Delta_{\text{fus}} H$	31.4	487.2		[2002GRA/RAS]	
		$\Delta_{\text{fus}} H$	30.85	489		[2006NOR/RAS2]	
		$\Delta_{\text{fus}} H$	32.0	488	DSC	[1992HEA/SIN]	
		$\Delta_{\text{fus}} H$	30.9	488.1		[1979ARM/JAM]	
		$\Delta_{\text{sub}} H$	117.0 ± 0.5	298	C	[2005PIN/DIO]	
		$\Delta_{\text{sub}} H$	112.4 ± 0.7	363	C	[1993SAB/LE]	
		$\Delta_{\text{sub}} H$	114.1 ± 0.7	298		[1993SAB/LE]	
		(398–433)	116.3		GS	[1954DAV/JON, 1960JON]	
C ₇ H ₆ O ₃	[533-31-3]	5-hydroxy-1,3-benzodioxole (sesamol)					
		$\Delta_{\text{fus}} H$	16.96	337.7		[2004MAT/MON]	
		$\Delta_{\text{sub}} H$	(293–309)	92.1 ± 0.6	301	ME	[2004MAT/MON]
		(293–309)	92.2 ± 0.6	298	ME	[2004MAT/MON]	
C ₇ H ₆ O ₃ S	[4066-41-5]	5-acetyl-2-thiophenecarboxylic acid					
		$\Delta_{\text{sub}} H$	(364–387)	119.6 ± 0.6	375.2	ME	[2008RIB/SAN5]
		$\Delta_{\text{sub}} H$	(364–387)	123.5 ± 0.6	298	ME	[2008RIB/SAN5]
C ₇ H ₆ O ₄	[303-38-8]	2,3-dihydroxybenzoic acid					
		$\Delta_{\text{fus}} H$	31.9	476.6	DSC	[2010MON/GON]	
		$\Delta_{\text{sub}} H$	(345–363)	109.1 ± 0.8	354	ME	[2010MON/GON]
		$\Delta_{\text{sub}} H$	(345–363)	110.7 ± 0.8	298	ME	[2010MON/GON]
			116 ± 4		TGA	[1999PRI/BAS]	
C ₇ H ₆ O ₄	[89-86-1]	2,4-dihydroxybenzoic acid					
		$\Delta_{\text{sub}} H$	(376–392)	124.0 ± 0.8	384	ME	[2010MON/GON]
		$\Delta_{\text{sub}} H$	(376–392)	126.4 ± 0.8	298	ME	[2010MON/GON]
			126 ± 6		TGA	[1999PRI/BAS]	
C ₇ H ₆ O ₄	[490-79-9]	2,5-dihydroxybenzoic acid					
		$\Delta_{\text{fus}} H$	20.8	476.2	DSC	[2010MON/GON]	
		$\Delta_{\text{sub}} H$	(372–389)	128.1 ± 1.4	380	ME	[2010MON/GON]
		$\Delta_{\text{sub}} H$	(372–389)	130.4 ± 1.3	298	ME	[2010MON/GON]
		$\Delta_{\text{sub}} H$	(362–379)	117.9 ± 1.4	370		[2006CHE/OJA]
			109 ± 3		TGA	[1999PRI/BAS]	
C ₇ H ₆ O ₄	[303-07-1]	2,6-dihydroxybenzoic acid					
		$\Delta_{\text{sub}} H$	(347–365)	107.5 ± 1.0	356	ME	[2010MON/GON]
		$\Delta_{\text{sub}} H$	(347–365)	109.1 ± 1.0	298	ME	[2010MON/GON]
			111 ± 7		TGA	[1999PRI/BAS]	
C ₇ H ₆ O ₄	[99-50-3]	3,4-dihydroxybenzoic acid					
		$\Delta_{\text{fus}} H$	31.2	472.3	DSC	[2009QUE/MOT]	
		$\Delta_{\text{sub}} H$	(387–403)	132.3 ± 1.2	395	ME	[2010MON/GON]
		$\Delta_{\text{sub}} H$	(387–403)	135.1 ± 1.2	298	ME	[2010MON/GON]
			153 ± 9		TGA	[1999PRI/BAS]	
C ₇ H ₆ O ₄	[99-10-5]	3,5-dihydroxybenzoic acid					
		$\Delta_{\text{fus}} H$	38.3	508.3	DSC	[2010MON/GON]	
		$\Delta_{\text{sub}} H$	(345–363)	139.8 ± 1.8	416	ME	[2010MON/GON]
		$\Delta_{\text{sub}} H$	(345–363)	143.2 ± 1.8	298	ME	[2010MON/GON]
			135 ± 6		TGA	[1999PRI/BAS]	

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₇ H ₆ O ₅	[149-91-7]	3,4,5-trihydroxybenzoic acid				
	$\Delta_{\text{sub}}H$	(391–421)	75.1	406		[1934HIR]
C ₇ H ₇ Br	[100-39-0]	benzylbromide				
	$\Delta_{\text{fus}}H$		13.2	271.8		[1976ASH]
	Δ_vH	(284–306)	53.3 ± 0.7	298	GS	[2002KRA/VAS]
	Δ_vH		53.7	298	CGC	[2002KRA/VAS]
	Δ_vH	(340–409)	48.1	355	I,A	[1976ASH, 1987STE/MAL]
	Δ_vH		50.5 ± 0.5	298		[1976ASH]
C ₇ H ₇ Br	[95-46-5]	2-bromotoluene				
	Δ_vH	(322–455)	47.2	337		[1999DYK/SVO]
	Δ_vH	(353–518)	45.3	368	A	[1987STE/MAL, 1970DYK/VAN, 1973KKY/REP]
	Δ_vH	(297–455)	52.6	312		[1947STU]
	Δ_vH	(273–348)	48.8	288		[1940STU/SAY]
C ₇ H ₇ Br	[591-17-3]	3-bromotoluene				
	Δ_vH	(351–457)	47.7	366		[1999DYK/SVO]
	Δ_vH	(287–457)	48.3	302	A	[1987STE/MAL, 1947STU]
C ₇ H ₇ Br	[108-38-7]	4-bromotoluene				
	$\Delta_{\text{fus}}H$		15.13	301.2		[1996VAN/ALV]
	Δ_vH	(320–458)	47.1	335		[1999DYK/SVO]
	Δ_vH	(273–472)	55.3	288		[1999DYK/SVO]
C ₇ H ₇ BrO	[578-57-4]	2-bromoanisole				
	Δ_vH		61.8 ± 1.3	298	C	[2009RIB/FER3]
	Δ_vH		52.3			[1986BAL/GNA]
	[2398-37-0]	3-bromoanisole				
Δ_vH		58.0 ± 1.2	298	C	[2009RIB/FER3]	
Δ_vH		50.2			[1986BAL/GNA]	
C ₇ H ₇ BrO	[104-92-7]	4-bromoanisole				
	Δ_vH		58.3 ± 1.2	298	C	[2009RIB/FER3]
	Δ_vH		50.6			[1986BAL/GNA]
C ₇ H ₇ BrO	[104-92-7]	4-bromoanisole				
	Δ_vH	(318–496)	48.9	333		[1947STU]
C ₇ H ₇ BrS	[19614-16-5]	2-bromothioanisole				
	Δ_vH		56.5			[1986BAL/GNA]
C ₇ H ₇ BrS	[33733-73-2]	3-bromothioanisole				
	Δ_vH		54.4			[1986BAL/GNA]
C ₇ H ₇ BrS	[104-95-0]	4-bromothioanisole				
	Δ_vH		55.7			[1986BAL/GNA]
C ₇ H ₇ Cl	[100-44-7]	benzyl chloride				
	$\Delta_{\text{fus}}H$		8.74	230	DSC	[1972AHM/EAD]
	Δ_vH	(276–309)	50.1 ± 0.3	298	GS	[2002KRA/VAS]
	Δ_vH		49.9	298	CGC	[2002KRA/VAS]
	Δ_vH	(320–390)	48.6	335	A,I	[1987STE/MAL, 1976ASH]
	Δ_vH		50.1 ± 0.5	298		[1976ASH, 1999DYK/SVO]
Δ_vH	(295–453)	48.6	310	A	[1987STE/MAL, 1947STU]	

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₇ H ₇ Cl	[95-49-8]	2-chlorotoluene				
	$\Delta_{\text{fus}}H$		10.3	237	DSC	[1972AHM/EAD]
	Δ_vH	(370–432)	41.6	385		[1999DYK/SVO]
	Δ_vH	(345–430)	45.3	298		[1984BOU/FRI, 1991BAS/SVO]
	Δ_vH	(345–430)	42.5	361		[1984BOU/FRI]
	Δ_vH	(338–493)	42.8	353	A	[1987STE/MAL, 1973KKY/REP, 1970DYK/VAN]
	Δ_vH	(278–432)	44.8	293		[1947STU]
C ₇ H ₇ Cl	[108-41-8]	3-chlorotoluene				
	Δ_vH	(373–435)	41.9	388		[1999DYK/SVO]
	Δ_vH	(277–436)	43.7	292	A	[1987STE/MAL, 1947STU]
C ₇ H ₇ Cl	[106-43-4]	4-chlorotoluene				
	$\Delta_{\text{fus}}H$		13.55	280.7		[1996VAN/ALV]
	Δ_vH	(362–435)	41.8	375		[1999DYK/SVO]
	Δ_vH	(304–436)	41.7	319	A	[1987STE/MAL]
	Δ_vH	(340–430)	46.0	298		[1984BOU/FRI, 1991BAS/SVO]
	Δ_vH	(338–433)	43.5	353		[1984BOU/FRI]
C ₇ H ₇ ClN ₂ O	[5814-05-1]	2-chlorobenzoic acid hydrazide				
	$\Delta_{\text{fus}}H$		25.6	392.2	DSC	[2003CHI/ACR]
C ₇ H ₇ ClN ₂ O	[536-40-3]	4-chlorobenzoic acid hydrazide				
	$\Delta_{\text{fus}}H$		32.9	437.2	DSC	[2003CHI/ACR]
C ₇ H ₇ ClN ₂ O ₂	[23042-32-2]	N-methyl-N-(4-chlorophenyl)nitramine				
	$\Delta_{\text{fus}}H$		19.5	323.4		[2002DAS/ZAL]
C ₇ H ₇ ClN ₂ S	[5344-82-1]	1-(<i>o</i> -chlorophenyl)thiourea				
	$\Delta_{\text{fus}}H$		22.29	413.5	DSC	[1990DON/DRE]
C ₇ H ₇ ClO	[766-51-8]	2-chloroanisole				
	Δ_vH		55.0 ± 0.8	298	C	[2008RIB/FER2]
	Δ_vH		49.4			[1986BAL/GNA]
	Δ_vH	(388–460)	48.3	403	A	[1987STE/MAL, 1973KKY/REP, 1999DYK/SVO]
C ₇ H ₇ ClO	[2845-89-8]	3-chloroanisole				
	Δ_vH		53.6 ± 0.8	298	C	[2008RIB/FER2]
	Δ_vH		48.1			[1986BAL/GNA]
C ₇ H ₇ ClO	[623-12-1]	4-chloroanisole				
	Δ_vH		54.8 ± 0.8	298	C	[2008RIB/FER2]
	Δ_vH		47.7			[1986BAL/GNA]
C ₇ H ₇ ClO	[17733-22-1]	2-chlorothioanisole				
	Δ_vH		53.6			[1986BAL/GNA]
C ₇ H ₇ ClO	[4867-37-2]	3-chlorothioanisole				
	Δ_vH		51.9			[1986BAL/GNA]
C ₇ H ₇ ClO	[123-09-1]	4-chlorothioanisole				
	Δ_vH		53.1			[1986BAL/GNA]
C ₇ H ₇ Cl ₃ NO ₃ PS	[5598-13-0]	O,O-dimethyl-O-3,5,6-trichloro-2-pyridyl phosphorothioate (chlorpyrifos methyl)				
	Δ_vH	(373–403)	73.0		GC	[2007GOE/MCC]
C ₇ H ₇ F	[350-50-5]	benzyl fluoride				

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(278–318)	46.2 ± 0.3	298	GS	[2002KRA/VAS]
	$\Delta_v H$		46.5	298	CGC	[2002KRA/VAS]
	$\Delta_v H$	(278–318)	46.3 ± 0.3	298	GS	[1997SCH/VER]
	$\Delta_v H$	(297–410)	43.7	312	A	[1987STE/MAL]
	$\Delta_v H$	(298–356)	44.3	312	I	[1976ASH]
	$\Delta_v H$		44.5 ± 0.4	298		[1976ASH]
C₇H₇F	[95-52-3]	2-fluorotoluene				
	$\Delta_{\text{fus}}H$		9.8	210.7		[1996DOM/HEA]
	$\Delta_{\text{fus}}H$		9.8	204	DSC	[1972AHM/EAD]
	$\Delta_v H$	(248–388)	42.0	263		[1999DYK/SVO]
	$\Delta_v H$	(452–531)	31.5	465		[1999DYK/SVO]
	$\Delta_v H$	(453–530)	32.3	468		[1984MOU]
	$\Delta_v H$	(308–348)	38.0	323		[1974MOZ/KOL, 1984BOU/FRI]
	$\Delta_v H$	(295–388)	38.7	310	A	[1987STE/MAL, 1951POT/SAY]
	$\Delta_v H$	(248–387)	40.5	264		[1947STU]
C₇H₇F	[352-70-5]	3-fluorotoluene				
	$\Delta_{\text{fus}}H$		8.3	184		[1996DOM/HEA]
	$\Delta_{\text{fus}}H$		6.31	185	DSC	[1972AHM/EAD]
	$\Delta_v H$	(250–390)	41.6	265		[1999DYK/SVO]
	$\Delta_v H$	(293–390)	39.2	308	A	[1987STE/MAL, 1951POT/SAY]
	$\Delta_v H$	(250–389)	40.7	266		[1947STU]
C₇H₇F	[352-32-9]	4-fluorotoluene				
	$\Delta_{\text{fus}}H$		9.35	216.5		[1996DOM/HEA]
	$\Delta_{\text{fus}}H$		7.65	213	DSC	[1972AHM/EAD]
	$\Delta_v H$	(340–430)	39.5	298		[1984BOU/FRI, 1991BAS/SVO]
	$\Delta_v H$	(340–429)	37.0	355	A	[1987STE/MAL, 1951POT/SAY, 1999DYK/SVO]
C₇H₇FO	[321-28-8]	2-fluoroanisole				
	$\Delta_v H$		52.1 ± 1.1	298	C	[2009RIB/FER2]
C₇H₇FO	[456-49-5]	3-fluoroanisole				
	$\Delta_v H$		48.1 ± 1.1	298	C	[2009RIB/FER2]
C₇H₇FO	[459-60-9]	4-fluoroanisole				
	$\Delta_v H$		48.7 ± 1.2	298	C	[2009RIB/FER2]
C₇H₇F₂N	[23162-99-4]	N,N-difluorobenzylamine				
	$\Delta_v H$	(313–333)	77.8	323	A	[1987STE/MAL]
C₇H₇F₉O	[72372-80-6]	1,1,1,2,2,3,3,4,4-nonfluoro-4-propoxybutane				
	$\Delta_v H$	(288–369)	37.9	303	I	[2002MUR/YAM]
C₇H₇I	[620-05-3]	benzyl iodide				
	$\Delta_{\text{fus}}H$		13.2	299.5		[1976ASH]
	$\Delta_v H$	(301–337)	57.4 ± 0.3	298	GS	[2002KRA/VAS]
	$\Delta_v H$		57.7	298	CGC	[2002KRA/VAS]
	$\Delta_v H$	(360–400)	46.8	375	I,A	[1987STE/MAL, 1976ASH]
	$\Delta_v H$		50.6 ± 1.4	298		[1976ASH]
C₇H₇I	[615-37-2]	2-iodotoluene				
	$\Delta_v H$	(310–484)	49.7	325	A	[1987STE/MAL, 1947STU]
C₇H₇I	[624-31-7]	4-iodotoluene				
	$\Delta_{\text{fus}}H$		14.96	306.7		[1996VAN/ALV]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₇ H ₇ IO	[696-62-8]	4-iodoanisole				
	$\Delta_v H$	(401–520)	54.4	416	A	[1987STE/MAL, 1999DYK/SVO]
	$\Delta_v H$	(401–479)	53.1 ± 0.4	440	I	[1956BRE/UBB]
C ₇ H ₇ NO	[6264-93-3]	2-aminotropone				
	$\Delta_{\text{sub}} H$	(273–333)	71.13 ± 0.4		ME	[1971JAC/HUN]
C ₇ H ₇ NO	[55-21-0]	benzamide				
	$\Delta_{\text{fus}} H$		19.15	402.1	DSC	[2009BRI]
	$\Delta_{\text{fus}} H$		23.14	NA	DSC	[2008SIN/DAS2]
	$\Delta_{\text{fus}} H$		18.49	402.3		[1991ACR]
	$\Delta_{\text{sub}} H$	(325–342)	96.9	333.5	A	[1987STE/MAL, 1960AIH2]
	$\Delta_{\text{sub}} H$	(323–349)	101.7 ± 1	298	C	[1982TOR/SAB2]
C ₇ H ₇ NO	[103-70-8]	formanilide				
	$\Delta_{\text{sub}} H$	(298–318)	77.8	308	A	[1987STE/MAL, 1960AIH2]
C ₇ H ₇ NO	[1122-62-9]	2-acetylpyridine				
	$\Delta_v H$		60.5 ± 0.3	298	C	[2007FRE/OLI]
C ₇ H ₇ NO	[350-03-8]	3-acetylpyridine				
	$\Delta_v H$		66.1 ± 0.8	298	C	[2007FRE/OLI]
C ₇ H ₇ NO	[1122-54-9]	4-acetylpyridine				
	$\Delta_v H$		66.5 ± 0.9	298	C	[2007FRE/OLI]
C ₇ H ₇ NO ₂	[622-42-4]	(nitromethyl)benzene				
	$\Delta_v H$	(363–413)	53.8	378	A	[1987STE/MAL]
C ₇ H ₇ NO ₂	[88-72-2]	2-nitrotoluene				
	$\Delta_v H$	(283–313)	59.6 ± 1.6	298	GS	[2010WID/BRU]
	$\Delta_v H$	(274–323)	59.0 ± 0.3	299	GS	[2000VER/HEI]
	$\Delta_v H$		59.1 ± 0.3	298		[2000VER/HEI]
	$\Delta_v H$	(388–448)	52.0	403	EB	[1994AIM]
	$\Delta_v H$	(402–496)	51.0	417	A	[1987STE/MAL]
	$\Delta_v H$	(387–493)	52.2	402		[1938LEV/SHT, 1994AIM]
C ₇ H ₇ NO ₂	[99-08-1]	3-nitrotoluene				
	$\Delta_v H$	(293–313)	56.6 ± 2.5	303	GS	[2010WID/BRU]
	$\Delta_v H$	(397–452)	52.8	413	EB	[1994AIM]
	$\Delta_v H$	(353–505)	49.8	368	A	[1987STE/MAL]
C ₇ H ₇ NO ₂	[99-99-0]	4-nitrotoluene				
	$\Delta_{\text{fus}} H$		16.81	324.8		[1991ACR]
	$\Delta_{\text{sub}} H$	(283–313)	74.8 ± 1.0	298	GS	[2010WID/BRU]
	$\Delta_{\text{sub}} H$		79.1 ± 2.5	298	ME	[1971LEN/VEL]
	$\Delta_{\text{sub}} H$	(298–310)	79.1	298		[1970LEN/VEL]
	$\Delta_v H$	(407–457)	52.8	422	EB	[1994AIM]
	$\Delta_v H$	(423–512)	49.8	438	A	[1987STE/MAL]
	$\Delta_v H$	(387–493)	54.2	402		[1938LEV/SHT, 1994AIM]
	C ₇ H ₇ NO ₂	[118-92-3]	2-aminobenzoic acid (I)			
$\Delta_{\text{fus}} H$			20.5	417.8		[1991ACR]
$\Delta_{\text{sub}} H$			111.6 ± 1.7	298		[1972ARN/JON]
C ₇ H ₇ NO ₂	[na]	2-aminobenzoic acid (II)				
	$\Delta_{\text{sub}} H$		100 ± 1	338	TE,ME	[1979DEK/VOO]
	$\Delta_{\text{sub}} H$		99.6 ± 0.5	378	C	[1974SAB/CHA]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₇ H ₇ NO ₂	$\Delta_{\text{sub}}H$		104.9 ± 1	298	C	[1974SAB/CHA]
	[99-05-8]	3-aminobenzoic acid				
	$\Delta_{\text{fus}}H$ (I)		35.5	445.2	DSC	[2010SVA/NOR]
	$\Delta_{\text{fus}}H$ (II)		26.7	451.2		[2010SVA/NOR]
	$\Delta_{\text{fus}}H$		33.7	445.7		[2001ROT/GLA]
	$\Delta_{\text{fus}}H$		21.84	452.9		[1991ACR]
	$\Delta_{\text{sub}}H$		122 ± 1	374.8	TE	[1979DEK/VOO]
$\Delta_{\text{sub}}H$	(367–389)	122.3 ± 3		C	[1974SAB/CHA]	
$\Delta_{\text{sub}}H$		128 ± 3.2	298	C	[1974SAB/CHA, 1977NAB/SAB]	
C ₇ H ₇ NO ₂	[150-13-0]	4-aminobenzoic acid				
	$\Delta_{\text{fus}}H$		2.06	355.2	DSC	
	$\Delta_{\text{fus}}H$		22.62	458.7		[2004GRA/RAS]
	$\Delta_{\text{fus}}H$		24.5	459.2		[2001ROT/GLA]
	$\Delta_{\text{fus}}H$		20.92	461.4		[1991ACR]
	$\Delta_{\text{sub}}H$		112.3 ± 1	373	TE	[1979DEK/VOO]
	$\Delta_{\text{sub}}H$	(367–389)	114 ± 3.5		C	[1974SAB/CHA]
$\Delta_{\text{sub}}H$		116 ± 3.7	298	C	[1974SAB/CHA, 1977NAB/SAB]	
$\Delta_{\text{sub}}H$		U 142			[1938WOL/WEG, 1960JON]	
C ₇ H ₇ NO ₂	[94-67-7]	2-hydroxybenzaloxime				
	$\Delta_{\text{sub}}H$ (<i>mp</i> 330 K)	(423–513)	96.7 ± 9.4	468	DSC	[1984BUR/MOR]
	$\Delta_{\text{sub}}H$		105.2 ± 10	298		[1984BUR/MOR]
	$\Delta_{\text{sub}}H$ (<i>anti</i>)		U 51 ± 1.7		MS	[1983MAJ/AZZ]
$\Delta_{\text{sub}}H$ (<i>syn</i>)		U 65.3 ± 1.7		MS	[1983MAJ/AZZ]	
C ₇ H ₇ NO ₂	[22241-18-5]	3-hydroxybenzaloxime				
	$\Delta_{\text{sub}}H$ (<i>anti</i>)		U 52.7 ± 1.7		MS	[1983MAJ/AZZ]
$\Delta_{\text{sub}}H$ (<i>syn</i>)		U 57.3 ± 1.7		MS	[1983MAJ/AZZ]	
C ₇ H ₇ NO ₂	[699-06-9]	4-hydroxybenzaloxime				
	$\Delta_{\text{sub}}H$ (<i>anti</i>)		U 54.4 ± 1.7		MS	[1983MAJ/AZZ]
C ₇ H ₇ NO ₂	[65-45-2]	2-hydroxybenzamide (salicylamide)				
	$\Delta_{\text{fus}}H$		27.1	414.9	DSC	[2008BER/MIN]
	$\Delta_{\text{fus}}H$		29.0	411.9		[2006NOR/RAS]
	$\Delta_{\text{sub}}H$		101.9 ± 0.4	298	C	[2008BER/MIN]
$\Delta_{\text{sub}}H$		99.3 ± 1.3	298	C	[2007RIB/ARA]	
C ₇ H ₇ NO ₂	[619-57-8]	4-hydroxybenzamide				
	$\Delta_{\text{fus}}H$		25.4	433.8	DSC	[2008BER/MIN]
	$\Delta_{\text{fus}}H$		25.2	433.2	DSC	[2007PER/HAN]
	$\Delta_{\text{sub}}H$		129.7 ± 1.9	298	C	[2008BER/MIN]
	$\Delta_{\text{sub}}H$	(360–420)	115.6 ± 0.6	390	GS	[2007PER/HAN]
$\Delta_{\text{sub}}H$	(360–420)	117.8 ± 0.6	298	GS	[2007PER/HAN]	
C ₇ H ₇ NO ₂	[622-42-8]	phenyl carbamate				
	$\Delta_{\text{fus}}H$ (I)		15.42	414.9	DSC	
$\Delta_{\text{fus}}H$ (II)		22.12	417.6	DSC	[2008WIS/BER]	
C ₇ H ₇ NO ₂	[2459-07-6]	methyl picolinate				
	Δ_vH		67.0 ± 1.8	298	C	[2007RIB/FRE]
Δ_vH	(273–340)	64.1 ± 0.1	298		[2007RIB/FRE]	
C ₇ H ₇ NO ₂	[93-60-7]	methyl nicotinate				
	Δ_vH	(298–324)	61.2 ± 0.2	298		[2007RIB/FRE]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₇ H ₇ NO ₂	[2459-09-8]	methyl isonicotinate				
	$\Delta_v H$		65.4 ± 1.4	298	C	[2007RIB/FRE]
	$\Delta_v H$	(298–320)	59.4 ± 0.1	298		[2007RIB/FRE]
C ₇ H ₇ NO ₃	[5399-68-8]	2,4-dihydroxybenzaloxime				
	$\Delta_{\text{sub}}H$ (<i>anti</i>)		U 76.2 ± 1.7		MS	[1983MAJ/AZZ]
	$\Delta_{\text{sub}}H$ (<i>syn</i>)		U 93.7 ± 1.7		MS	[1983MAJ/AZZ]
C ₇ H ₇ NO ₃	[91-23-6]	2-nitroanisole				
	$\Delta_v H$	(424–545)	58.6	439	A	[1987STE/MAL]
C ₇ H ₇ NO ₃	[2581-34-2]	4-nitro-5-methylphenol				
	$\Delta_{\text{fus}}H$		27.4	401		[1991ACR]
C ₇ H ₇ NO ₃	[700-38-9]	2-nitro-5-methylphenol				
	$\Delta_{\text{fus}}H$		20.79	302.8		[1991ACR]
	$\Delta_v H$	(331–358)	62.8 ± 0.5	298	GS	[2007HEI/KAP]
C ₇ H ₇ NO ₃	[2581-34-2]	3-methyl-4-nitrophenol				
	$\Delta_v H$		85.8	298	B	[2007HEI/KAP]
C ₇ H ₇ NO ₃	[554-84-7]	2-methyl-5-nitrophenol				
	$\Delta_v H$		85.9	298	B	[2007HEI/KAP]
C ₇ H ₇ NO ₃	[65-49-6]	4-aminosalicylic acid				
	$\Delta_{\text{fus}}H$		47.9	406.2		[2001ROT/GLA]
C ₇ H ₇ NO ₃	[89-57-6]	5-aminosalicylic acid				
	$\Delta_{\text{fus}}H$		67.2	543.2		[2001ROT/GLA]
C ₇ H ₇ NO ₃	[619-73-8]	4-nitrobenzyl alcohol				
	$\Delta_{\text{fus}}H$	(78–396)	20.97	336.4	AC	[2009MEN/TAN]
C ₇ H ₇ NO ₄	[3251-56-7]	2-methoxy-4-nitrophenol				
	$\Delta_{\text{fus}}H$		21.69	374.4		[2004MIR/MOR]
	$\Delta_{\text{sub}}H$		99.4 ± 2.0	298	C	[2004MIR/MOR]
C ₇ H ₇ NO ₄	[636-93-1]	2-methoxy-5-nitrophenol				
	$\Delta_{\text{fus}}H$		21.43	377.6		[2004MIR/MOR]
	$\Delta_{\text{sub}}H$		106.2 ± 2.2	208	C	[2004MIR/MOR]
C ₇ H ₇ NO ₄	[1568-70-3]	4-methoxy-2-nitrophenol				
	$\Delta_{\text{fus}}H$		22.42	352.3		[2004MIR/MOR]
	$\Delta_{\text{sub}}H$		90.8 ± 1.7	298	C	[2004MIR/MOR]
C ₇ H ₇ NS	[2227-79-4]	thiobenzamide				
	$\Delta_{\text{sub}}H$		103.4 ± 2.2	298	C	[1989RIB/SOU]
	$\Delta_{\text{sub}}H$		97.2 ± 0.6	298	C	[1982SAB/TOR]
C ₇ H ₇ N ₃	[622-79-7]	(azidomethyl)benzene				
	$\Delta_v H$	(333–363)	48.0	348	A	[1987STE/MAL]
C ₇ H ₇ N ₃ O ₂	[na]	N-acetyl-pyrazinamide				
	$\Delta_{\text{fus}}H$		23.6	366.7		[1991LIU/GUO]
C ₇ H ₇ N ₃ O ₄	[16698-03-6]	N-methyl-N-(4-nitrophenyl)nitramine				
	$\Delta_{\text{fus}}H$		25.0	416.9		[2002DAS/ZAL]
C ₇ H ₇ N ₃ O ₄	[55739-03-2]	N-methyl-N-(3-nitrophenyl)nitramine				

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound					
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference	
		$\Delta_{\text{fus}}H$	25.3	350.5		[2002DAS/ZAL]	
C ₇ H ₈	[121-46-0]	bicyclo[2.2.1]hepta-2,5-diene					
		$\Delta_{\text{trs}}H$	8.2	202			
		$\Delta_{\text{fus}}H$	1.91	255.6		[2004BYK/SMI]	
		$\Delta_{\text{trs}}H$	8.93	202			
		$\Delta_{\text{fus}}H$	NA			[1974CLA/MCK]	
		Δ_vH	34.8 ± 0.1	298	C	[1993AN/XIE]	
		Δ_vH	(300–364)	33.6	315	A	[1987STE/MAL]
		Δ_vH	34.7 ± 0.1	298	C	[1985KUS]	
		Δ_vH	33.8 ± 0.9	298		[1978STE2]	
		Δ_vH	(300–353)	32.9 ± 0.8	298	BG	[1973HAL/SMI]
C ₇ H ₈	[544-25-2]	1,3,5-cycloheptatriene					
		$\Delta_{\text{trs}}H$	2.35	154			
		$\Delta_{\text{fus}}H$	1.16	198		[1996DOM/HEA]	
		Δ_vH	(273–338)	40.8	288	A	[1987STE/MAL, 1973KKY/REP]
		Δ_vH	(273–416)	39.4	288	A,EB	[1987STE/MAL, 1956FIN/SCO]
		Δ_vH	38.7 ± 0.2	298		[1956FIN/SCO]	
C ₇ H ₈	[278-06-8]	tetracyclo[3.2.0.0 ^{2,7} .0 ^{4,6}]heptane (quadricyclane)					
		$\Delta_{\text{trs}}H$	7.2	180			
		$\Delta_{\text{fus}}H$	1.09	228		[1996DOM/HEA]	
		Δ_vH	37.9 ± 0.1	298	C	[1993AN/XIE]	
		Δ_vH	37.9 ± 0.1	298	C	[1985KUS]	
		Δ_vH	37.0 ± 0.8	298		[1978STE2]	
		Δ_vH	(302–372)	37.3 ± 0.8	317	BG	[1987STE/MAL, 1973HAL/SMI]
C ₇ H ₈	[108-88-3]	toluene					
		$\Delta_{\text{fus}}H$	6.61	178		[1996DOM/HEA, 1931SOU/AND]	
		$\Delta_{\text{sub}}H$	43.1	298	B	[1970LEN/VEL]	
		Δ_vH	(331–496)	35.7	346		[1993LEE/HOL]
		Δ_vH	(210–279)	40.6	264	A	[1987STE/MAL]
		Δ_vH	(383–445)	34.4	398	A	[1987STE/MAL]
		Δ_vH	(440–531)	33.2	455	A	[1987STE/MAL]
		Δ_vH	(530–592)	33.3	545	A	[1987STE/MAL]
		Δ_vH	(273–295)	38.9	284	A	[1987STE/MAL]
		Δ_vH	33.5 ± 0.1	380	C	[1985NAT/VIS]	
		Δ_vH	32.1 ± 0.1	403	C	[1985NAT/VIS]	
		Δ_vH	29.4 ± 0.1	441	C	[1985NAT/VIS]	
		Δ_vH	27.1 ± 0.1	470	C	[1985NAT/VIS]	
		Δ_vH	24.0 ± 0.1	505	C	[1985NAT/VIS]	
		Δ_vH	35.4	333		[1984EUB/CED]	
		Δ_vH	33.4	373		[1984EUB/CED]	
		Δ_vH	31.4	413		[1984EUB/CED]	
		Δ_vH	28.4	453		[1984EUB/CED]	
		Δ_vH	24.0	493		[1984EUB/CED]	
		Δ_vH	(343–383)	35.4	360		[1975RIV]
		Δ_vH		38.0	298		[1971WIL/ZWO]
		Δ_vH	(303–343)	37.3	318		[1968GAW/SWI2]
		Δ_vH	(288–348)	36.9	303		[1967VAN/SOC]
	Δ_vH	(210–293)	37.8	278		[1956MIL]	
	Δ_vH	(308–386)	37.0	323		[1987STE/MAL, 1949FOR/NOR]	
	Δ_vH		38.0	298	C	[1947OSB/GIN]	

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(286–362)	37.8	301		[1946THO]
	$\Delta_v H$	(308–384)	37.0	323	MM	[1945WIL/TAY]
	$\Delta_v H$	(273–323)	38.8	288		[1943PIT/SCO]
C ₇ H ₈ ClN ₃ O ₄ S ₂	[58-93-5]	6-chloro-3,4-dihydro-2H-1,2,4-benzothiadiazine-7-sulfonamide-1,1-dioxide (hydrochlorthiazide)				
	$\Delta_{\text{fus}}H$		33.6	540.8	DSC	[2006WAS/HOL]
	$\Delta_{\text{fus}}H$		30.96	547.2		[2000HAN/PAR]
C ₇ H ₈ FN	[452-80-2]	2-fluoro-4-methylaniline				
	$\Delta_v H$		56.6 ± 0.6	298	C	[2007RIB/FER2]
C ₇ H ₈ FN	[452-84-6]	2-fluoro-5-methylaniline				
	$\Delta_v H$		56.9 ± 0.5	298	C	[2007RIB/FER2]
C ₇ H ₈ FN	[443-86-7]	3-fluoro-2-methylaniline				
	$\Delta_v H$		57.8 ± 0.6	298	C	[2007RIB/FER2]
C ₇ H ₈ FN	[452-71-1]	4-fluoro-2-methylaniline				
	$\Delta_v H$		59.8 ± 0.8	298	C	[2007RIB/FER2]
C ₇ H ₈ N ₂	[33496-46-7]	1-amino-7-imino-1,3,5-cycloheptatriene				
	$\Delta_{\text{sub}}H$	(273–333)	49.4 ± 0.4		ME	[1971JAC/HUN]
C ₇ H ₈ N ₂ O	[64-10-8]	monophenylurea				
	$\Delta_{\text{fus}}H$		23.68	420.6		[1991ACR]
	$\Delta_{\text{sub}}H$	(392–412)	136 ± 6	406	TE	[1987FER/DEL]
C ₇ H ₈ N ₂ O	[3398-07-0]	2-aminobenzaldoxime				
	$\Delta_{\text{sub}}H$ (anti)		U 33.9 ± 1.7		MS	[1983MAJ/AZZ]
	$\Delta_{\text{sub}}H$ (syn)		U 63.6 ± 1.7		MS	[1983MAJ/AZZ]
C ₇ H ₈ N ₂ O	[5231-96-9]	(2-pyridyl)acetamide				
	$\Delta_{\text{fus}}H$		16.0	343		[1979GON/CHA]
	$\Delta_{\text{sub}}H$		103.8	298	B,E	[1979GON/CHA]
C ₇ H ₈ N ₂ O	[613-94-5]	benzoic acid hydrazide				
	$\Delta_{\text{fus}}H$		25.7	388.2	DSC	[2003CHI/ACR]
C ₇ H ₈ N ₄ O	[31010-51-2]	9-ethylhypoxanthine				
	$\Delta_{\text{sub}}H$		108.8 ± 13		HSA	[1978NOW/SZC]
	$\Delta_{\text{sub}}H$		U 83.7		HSA	[1965CLA/PES]
C ₇ H ₈ N ₄ O	[20535-82-4]	1,9-dimethylhypoxanthine				
	$\Delta_{\text{sub}}H$		75.3 ± 13		HSA	[1978NOW/SZC]
C ₇ H ₈ N ₄ O ₂	[58-55-9]	1,3-dimethylxanthine (theophylline)				
	$\Delta_{\text{fus}}H$ (form I)		28.02	547.9	DSC	[2010SZT/LEG, 2009SZT]
	$\Delta_{\text{fus}}H$ (form II)		30.1	543.9	DSC	[2010SZT/LEG, 2009SZT]
	$\Delta_{\text{fus}}H$		19	546.8	DSC	[2009GOO/ROD]
	$\Delta_{\text{fus}}H$		28.2	544		[1989GON/KRA]
	$\Delta_{\text{fus}}H$		28.2	542.3		[1989SUZ/SHI]
	$\Delta_{\text{fus}}H$		31.2	543.7		[1983FOK/VAN]
	$\Delta_{\text{sub}}H$ (form I)	(413–453)	132.0 ± 0.3	433	T	[1999EMM/PIC]
	$\Delta_{\text{sub}}H$ (form I)		142	298		[1999EMM/PIC]
	$\Delta_{\text{sub}}H$ (form II)	(413–453)	134.2 ± 0.3	433	T	[1999EMM/PIC]
	$\Delta_{\text{sub}}H$ (form II)		144	298		[1999EMM/PIC]
	$\Delta_{\text{sub}}H$		126	421	ME,TE	[1983FOK/VAN]
	$\Delta_{\text{sub}}H$		135	298		[1983FOK/VAN, 1999EMM/PIC]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₇ H ₈ O	[100-66-3]	anisole				
	$\Delta_{\text{fus}}H$		11.66	237		[1996DOM/HEA, 1994LEE/LIE]
	$\Delta_{\text{fus}}H$		12.89	236		[1996DOM/HEA]
	Δ_vH	(278–312)	46.6 ± 0.2	298	GS	[2005VAS/VER]
	Δ_vH	(363–463)	44.3	298	GC	[2005HOS/GRY]
	Δ_vH	(353–393)	45.3	298	CGC	[1995CHI/HOS]
	Δ_vH	(382–429)	41.8	397		[1993REI/SAN]
	Δ_vH		41.0			[1986BAL/GNA]
	Δ_vH	(382–437)	41.9	397	A	[1987STE/MAL, 1976AMB/ELL]
	Δ_vH	(382–437)	46.9	298		[1976AMB/ELL]
	Δ_vH	(282–437)	39.0	426		[1976AMB/ELL]
	Δ_vH		46.8 ± 0.2	298	C	[1975FEN/HAR]
	Δ_vH	(363–383)	38.2 ± 0.4	298		[1972LEB/KAT2, 2005VAS/VER]
	Δ_vH		42.9 ± 0.1	367	C	[1967HAL/LEE]
	Δ_vH		42.0 ± 0.1	382	C	[1967HAL/LEE]
	Δ_vH		40.5 ± 0.1	402	C	[1967HAL/LEE]
	Δ_vH		38.9 ± 0.1	427	C	[1967HAL/LEE]
Δ_vH		39.4	298		[1957MCC/DOU, 2005VAS/VER]	
Δ_vH	(382–437)	41.9	397		[1955VON/GEB, 1965COL/COU]	
C ₇ H ₈ O	[100-51-6]	benzyl alcohol				
	$\Delta_{\text{fus}}H$		8.79	257.6		[1991ACR]
	Δ_vH	(282–323)	65.5 ± 0.4	298	GS	[2005VAS/VER]
	Δ_vH	(277–381)	64.8 ± 0.6	298	GS	[1999VER4]
	Δ_vH	(323–373)	69.5	298	CGC	[1995CHI/HOS]
	Δ_vH		60.5			[1995PAP/PIM]
	Δ_vH	(404–507)	62.5 ± 0.3	298	EB	[1990AMB/GHI, 2005VAS/VER]
	Δ_vH	(303–333)	66.2	318	GS	[1982GRA/FOS]
	Δ_vH	(385–573)	54.6	400	A	[1987STE/MAL, 1973KKY/REP]
	Δ_vH	(293–313)	61.5	303	A,ME	[1987STE/MAL, 1957SER/VOI, 1973KKY/REP]
	Δ_vH	(396–478)	62.1 ± 0.3	298	EB	[1949DRE/SHR, 1949DRE/MAR, 2005VAS/VER]
	Δ_vH	(312–348)	63.0 ± 2.2	298	EB	[1937GAR/BRE, 2005VAS/VER]
Δ_vH		60.3 ± 0.4	298	EB	[1926MAT, 2005VAS/VER]	
C ₇ H ₈ O	[95-48-7]	2-hydroxytoluene				
	$\Delta_{\text{fus}}H$		14.8	305.4	DSC	[2007RIC/BER]
	$\Delta_{\text{fus}}H$		15.9	304.1	DSC	[1998JAM/PAL]
	$\Delta_{\text{fus}}H$		15.82	304.2		[1996DOM/HEA]
	$\Delta_{\text{fus}}H$		14.8	304.1		[1990MEV/LIC]
	$\Delta_{\text{sub}}H$		73.7 ± 0.5	298	C	[2007RIC/BER]
	$\Delta_{\text{sub}}H$	(273–303)	74.8	288	A	[1987STE/MAL]
	$\Delta_{\text{sub}}H$	(273–303)	76.0 ± 0.8	288		[1960AND/BID, 1970COX/PIL]
	Δ_vH	(304–409)	58.5	319	A	[1987STE/MAL]
	Δ_vH	(399–470)	50.1	414	A	[1987STE/MAL]
	Δ_vH	(463–526)	46.2	478	A	[1987STE/MAL]
	Δ_vH	(517–630)	44.0	532	A	[1987STE/MAL]
	Δ_vH		50.2			[1986BAL/GNA]
	Δ_vH	(383–473)	51.3	398	GS,EB	[1987STE/MAL, 1960AND/BID, 1973KKY/REP]
	Δ_vH	(415–462)	48.2	438		[1939GOL/MAR]
C ₇ H ₈ O	[108-39-4]	3-hydroxytoluene				
	$\Delta_{\text{fus}}H$		8.9	282.3	DSC	[2007RIC/BER]
	$\Delta_{\text{fus}}H$		10.67	285.3	DSC	[1998JAM/PAL]
	$\Delta_{\text{fus}}H$		10.71	285.4		[1996DOM/HEA]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound					
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference	
		$\Delta_{\text{fus}}H$	9.1	280.8		[1990MEV/LIC]	
		$\Delta_{\text{sub}}H$	(273–285)	56.1	279	A	[1987STE/MAL]
		$\Delta_{\text{sub}}H$	(284–313)	61.7 ± 1.0		GS	[1960AND/BID]
		Δ_vH		65.0 ± 0.7	298	C	[2007RIC/BER]
		Δ_vH	(393–433)	62.5	298	CGC	[1995CHI/HOS]
		Δ_vH	(284–313)	61.7	298	A	[1987STE/MAL]
		Δ_vH	(285–416)	63.1	300	A	[1987STE/MAL]
		Δ_vH	(410–477)	52.7	425	A	[1987STE/MAL]
		Δ_vH	(471–531)	47.6	486	A	[1987STE/MAL]
		Δ_vH	(523–633)	43.8	538	A	[1987STE/MAL]
		Δ_vH	(383–473)	55.0	398	GS,EB	[1987STE/MAL, 1960AND/BID]
		Δ_vH	(388–429)	60.6	409	GS	[1980NAS/HWA, 1973KKY/REP]
		Δ_vH		61.7	298		[1958BID/MAR]
		Δ_vH	(359–473)	58.8	374		[1955VON/GEB, 1984BOU/FRI]
		Δ_vH	(422–474)	50.7	448		[1939GOL/MAR]
C₇H₈O	[106-44-5]	4-hydroxytoluene					
		$\Delta_{\text{fus}}H$	12.6	308.8	DSC	[2007RIC/BER]	
		$\Delta_{\text{fus}}H$	8.58	307.6	DSC	[1998JAM/PAL]	
		$\Delta_{\text{fus}}H$	12.72	307.9		[1996DOM/HEA]	
		$\Delta_{\text{fus}}H$	11.8	307.4		[1990MEV/LIC]	
		$\Delta_{\text{sub}}H$	73.1 ± 0.6	298	C	[2007RIC/BER]	
		$\Delta_{\text{sub}}H$	(273–307)	73.9 ± 1.5	290		[1960AND/BID, 1970COX/PIL]
		Δ_vH	(308–393)	62.0	323	A	[1987STE/MAL]
		Δ_vH	(385–477)	55.4	400	A	[1987STE/MAL]
		Δ_vH	(463–533)	49.2	478	A	[1987STE/MAL]
		Δ_vH	(523–635)	46.0	538	A	[1987STE/MAL]
		Δ_vH		54.0			[1986BAL/GNA]
		Δ_vH	(383–473)	55.6	398	A,GS,EB	[1987STE/MAL, 1960AND/BID, 1973KKY/REP]
		Δ_vH	(419–474)	51.3	446		[1939GOL/MAR]
C₇H₈OS	[106-53-6]	4-methoxybenzenethiol					
		Δ_vH	52.3			[1986BAL/GNA]	
C₇H₈OS	[13679-73-7]	2-acetyl-4-methylthiophene					
		Δ_vH	63.0 ± 2.6	298	C	[2008RIB/SAN4]	
C₇H₈OS	[13679-72-6]	2-acetyl-3-methylthiophene					
		Δ_vH	57.1 ± 2.4	298	C	[2008RIB/SAN4]	
C₇H₈OS	[13679-74-8]	2-acetyl-5-methylthiophene					
		Δ_vH	62.0 ± 2.6	298	C	[2008RIB/SAN4]	
C₇H₈OS	[36880-33-8]	5-ethyl-2-thiophenecarboxaldehyde					
		Δ_vH	62.2 ± 1.3	298	C	[2008RIB/SAN2]	
C₇H₈O₂	[488-17-5]	3-methyl-1,2-dihydroxybenzene					
		$\Delta_{\text{sub}}H$	93.2 ± 1.0	298	C	[1984CAR]	
C₇H₈O₂	[496-73-1]	2,4-dihydroxytoluene					
		$\Delta_{\text{fus}}H$	27.6	404.2		[1999VER7]	
		$\Delta_{\text{sub}}H$	(317–333)	106.8 ± 0.9	325	ME	[2009RIB/FER4]
		$\Delta_{\text{sub}}H$	(317–333)	107.3 ± 3.0	298	ME	[2009RIB/FER4]
		Δ_vH	(391–459)	72.2	406	A,GC	[1987STE/MAL, 1975KUN/LIL]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₇ H ₈ O ₂	[608-25-3]	2,6-dihydroxytoluene				
	$\Delta_{\text{sub}}H$	(309–329)	98.8 ± 0.3	319	ME	[2009RIB/FER4]
	$\Delta_{\text{sub}}H$	(309–329)	99.2 ± 2.3	298	ME	[2009RIB/FER4]
	Δ_vH	(398–434)	66.9	413	A,GC	[1987STE/MAL, 1975KUN/LIL]
C ₇ H ₈ O ₂	[452-86-8]	3,4-dihydroxytoluene				
	$\Delta_{\text{sub}}H$		94.9 ± 1.0	298	C	[1984CAR]
	Δ_vH	(387–415)	90.0	401	A	[1987STE/MAL]
C ₇ H ₈ O ₂	[504-15-4]	3,5-dihydroxytoluene				
	$\Delta_{\text{sub}}H$	(322–338)	102.3 ± 0.7	330	ME	[2009RIB/FER4]
	$\Delta_{\text{sub}}H$	(322–338)	102.9 ± 3.5	298	ME	[2009RIB/FER4]
	Δ_vH	(402–468)	76.6	417	A,GC	[1987STE/MAL, 1975KUN/LIL]
C ₇ H ₈ O ₂	[95-71-6]	2-methyl-1,4-dihydroxybenzene				
	$\Delta_{\text{fus}}H$		27.6	404.2	DSC	[1999VER7]
	$\Delta_{\text{sub}}H$	(325–341)	107.8 ± 1.1	333	ME	[2009RIB/FER4]
	$\Delta_{\text{sub}}H$	(325–341)	108.4 ± 3.9	298	ME	[2009RIB/FER4]
	$\Delta_{\text{sub}}H$	(333–368)	97.2 ± 1.4	351	GS	[1999VER7]
	$\Delta_{\text{sub}}H$	(333–368)	100.4 ± 1.4	298	GS	[1999VER7]
C ₇ H ₈ O ₂	[90-05-1]	2-methoxyphenol				
	Δ_vH		62.6 ± 0.5	298	C	[2003MAT/MIR]
	Δ_vH	(378–479)	52.7	393	A	[1987STE/MAL, 1973KKY/REP]
	Δ_vH	(355–478)	52.7	370		[1955VON/GEB]
C ₇ H ₈ O ₂	[150-19-6]	3-methoxyphenol				
	Δ_vH		75.9 ± 1.2	298	C	[2003MAT/MIR]
	Δ_vH	(413–518)	64.8	428	A	[1987STE/MAL, 1973KKY/REP]
C ₇ H ₈ O ₂	[150-76-5]	4-methoxyphenol				
	$\Delta_{\text{fus}}H$		18.3	328.4		[1997LEE/CHA]
	$\Delta_{\text{sub}}H$		94.4 ± 1.2	298	C	[2003MAT/MIR]
	$\Delta_{\text{sub}}H$	(278–300)	88.7	289		[1987STE/MAL, 1960AIH]
	Δ_vH		58.6			[1986BAL/GNA]
	Δ_vH	(418–518)	61.4	433	A	[1987STE/MAL, 1973KKY/REP]
C ₇ H ₈ O ₂	[63233-31-8]	2,3-dimethyl-2H-pyran-2-one				
	Δ_vH	(352–518)	64.9	367	A	[1987STE/MAL, 1947STU]
C ₇ H ₈ O ₂	[90-01-7]	2-hydroxybenzyl alcohol				
	$\Delta_{\text{fus}}H$		21.5	358.3		[2008PIN/DIO]
C ₇ H ₈ O ₂ S	[52911-98-5]	6-methyl-4-methoxy-2H-pyran-2-thione				
	$\Delta_{\text{sub}}H$	(402–415)	130.5 ± 5.9	408	B	[1974BEA/MUE]
	Δ_vH	(401–415)	108.9	408	A	[1987STE/MAL, 1999DYK/SVO, 1974BEA/MUE]
C ₇ H ₈ O ₂ S	[52911-99-6]	2-methyl-6-(methylthio)-4H-pyran-4-one				
	$\Delta_{\text{sub}}H$	(388–433)	87.4 ± 3.8	410	B	[1974BEA/MUE]
	Δ_vH	(387–432)	62.7	402	A	[1987STE/MAL, 1999DYK/SVO, 1974BEA/MUE]
C ₇ H ₈ O ₂ S	[3112-85-4]	methyl phenyl sulfone				
	$\Delta_{\text{sub}}H$		92 ± 2.9			[UR/MAC, 1970COX/PIL]
C ₇ H ₈ O ₂ S	[19432-68-9]	methyl 2-thiopheneacetate				

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$		61.9 ± 1.4	298	C	[2007ROU/TEM3]
C ₇ H ₈ O ₂ S	[58414-52-1]	methyl 3-thiopheneacetate				
	$\Delta_v H$		60.9 ± 1.3	298	C	[2007ROU/TEM3]
C ₇ H ₈ O ₂ S	[2810-04-0]	ethyl 2-thiophenecarboxylate				
	$\Delta_v H$		56.6 ± 1.3	298	C	[2009RIB/SAN2]
C ₇ H ₈ O ₃	[614-99-3]	2-furancarboxylic acid, ethyl ester				
	$\Delta_v H$	(354–389)	51.2	369	A	[1987STE/MAL]
	$\Delta_v H$	(310–468)	52.6	325		[1947STU]
C ₇ H ₈ O ₃	[4225-42-7]	3-methoxy-6-methyl-4H-pyran-4-one				
	$\Delta_v H$	(370–384)	72.8	377	A	[1987STE/MAL]
C ₇ H ₈ O ₃	[672-89-9]	4-methoxy-6-methyl-2H-pyran-2-one				
	$\Delta_v H$	(385–434)	57.4	400	A	[1987STE/MAL]
C ₇ H ₈ O ₃	[934-00-9]	3-methoxycatechol				
	$\Delta_v H$		91.7 ± 0.8	298		[1986RIB/RIB]
C ₇ H ₈ S	[100-53-8]	benzenemethanethiol				
	$\Delta_v H$	(394–436)	47.5	409		[1999DYK/SVO]
	$\Delta_v H$		56.6 ± 0.1	298		[1972GOO]
C ₇ H ₈ S	[137-06-4]	2-methylbenzenethiol				
	$\Delta_v H$	(351–498)	48.1	366		[1999DYK/SVO]
	$\Delta_v H$	(370–470)	46.6	394	A	[1987STE/MAL]
	$\Delta_v H$		46.0			[1986BAL/GNA]
C ₇ H ₈ S	[108-40-7]	3-methylbenzenethiol				
	$\Delta_v H$	(353–498)	48.7	368		[1999DYK/SVO]
	$\Delta_v H$	(380–471)	47.1	395	A	[1987STE/MAL]
C ₇ H ₈ S	[106-45-6]	4-methylbenzenethiol				
	$\Delta_v H$	(351–499)	48.1	366		[1999DYK/SVO]
	$\Delta_v H$	(379–471)	46.5	394	A	[1987STE/MAL]
	$\Delta_v H$		46.4			[1986BAL/GNA]
C ₇ H ₈ S	[100-68-5]	methyl phenyl sulfide				
	$\Delta_{\text{fus}}H$		14.85	256.4		[1974MES/FIN]
	$\Delta_v H$		47.7			[1986BAL/GNA]
	$\Delta_v H$		54.3 ± 0.1	298		[1972GOO, 1966OSB/DOU]
	$\Delta_v H$	(389–475)	47.5	404	A,EB	[1987STE/MAL, 1966OSB/DOU, 1999DYK/SVO]
	$\Delta_v H$	(323–353)	50.6 ± 2.1	298		[1962MAC/MAY]
C ₇ H ₈ S ₃	[698-42-0]	4,5-tetramethylene-1,3-dithiole-2-thione				
	$\Delta_{\text{sub}}H$	(340–352)	98.3	346		[1967GEI/SCH, 1970COX/PIL]
	$\Delta_{\text{sub}}H$		102.1 ± 2.9			[1967GEI/SCH, 1970COX/PIL]
C ₇ H ₈ S ₃	[14085-34-8]	4,5-tetramethylene-1,2-dithiole-3-thione				
	$\Delta_{\text{sub}}H$	(335–350)	101.6	342		[1972GEI/RAU]
	$\Delta_{\text{sub}}H$		105.3	298		[1972GEI/RAU]
C ₇ H ₉ Cl ₃ NO ₃ PS	[5598-13-0]	O,O-dimethyl-O-(3,5,6-trichloro-2-pyridyl)phosphorothioate				
	$\Delta_{\text{fus}}H$		25.92	318.7	DSC	[1990DON/DRE]
C ₇ H ₉ Cl ₃ OS	[79886-21-8]	2,3,3-trichloro-2-propenethioic acid, O-butyl ester				
	$\Delta_v H$	(383–433)			GC	[1980PIT/KIS]
C ₇ H ₉ F ₃ N ₂ O ₄	[433-33-0]	glycine, N-[N-(trifluoroacetyl)glycyl]methyl ester				

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
		$\Delta_{\text{sub}}H$	(323–419)	127.9	338	[1987STE/MAL, 1960WEY/KLI]
		Δ_vH	(420–443)	93.8	431	A [1987STE/MAL, 1973KKY/REP]
C ₇ H ₉ F ₅ O ₂	[680-28-4]	pentafluoropropionic acid, butyl ester				
		Δ_vH	(354–389)	38.6	369	A,EB [1987STE/MAL, 1969SHE/LAN, 1999DYK/SVO]
C ₇ H ₉ N	[100-46-9]	benzylamine				
		Δ_vH	(293–362)	52.7 ± 0.3	328	[2009MOK/RAZ]
		Δ_vH	(293–362)	54.6 ± 0.3	298	[2009MOK/RAZ]
		Δ_vH	(302–458)	51.8	317	A [1987STE/MAL, 1977CAR/LAY, 1947BEE/JUN]
C ₇ H ₉ N	[583-61-9]	2,3-dimethylpyridine				
		$\Delta_{\text{fus}}H$		13.48	258.6	[1994CHI/HOS]
		Δ_vH	(283–313)	52.0 ± 0.6	298	GS [1999VER4]
		Δ_vH	(328–476)	45.2	340	EB [1995STE/CHI2]
		Δ_vH	(328–476)	42.7	380	EB [1995STE/CHI2]
		Δ_vH	(328–476)	40.2	420	EB [1995STE/CHI2]
		Δ_vH	(328–476)	37.4	460	EB [1995STE/CHI2]
		Δ_vH	(323–373)	47.6	298	CGC [1995CHI/HOS]
		Δ_vH		46.9	313	C [1985MAJ/SVO2]
		Δ_vH		45.0	343	C [1985MAJ/SVO2]
		Δ_vH		43.5	368	C [1985MAJ/SVO2]
		Δ_vH	(372–436)	43.0	387	A [1987STE/MAL, 1973KKY/REP]
C ₇ H ₉ N	[108-47-4]	2,4-dimethylpyridine				
		$\Delta_{\text{fus}}H$		8.82	209.4	[1994CHI/HOS]
		Δ_vH	(323–373)	47.5	298	CGC [1995CHI/HOS]
		Δ_vH	(288–373)	45.5	330	[1995SAK/UEO]
		Δ_vH	(331–473)	44.8	340	EB [1995STE/CHI2]
		Δ_vH	(331–473)	42.3	380	EB [1995STE/CHI2]
		Δ_vH	(331–473)	39.8	420	EB [1995STE/CHI2]
		Δ_vH	(331–473)	37.0	460	EB [1995STE/CHI2]
		Δ_vH	(298–431)	47.1	313	EB [1990LEN]
		Δ_vH	(267–358)	47.5	282	MM [1986WIS/LEN]
		Δ_vH		46.5	313	C [1985MAJ/SVO2]
		Δ_vH		44.6	343	C [1985MAJ/SVO2]
		Δ_vH		43.9	368	C [1985MAJ/SVO2]
		Δ_vH	(349–433)	43.5	364	A [1987STE/MAL, 1973KKY/REP]
C ₇ H ₉ N	[589-93-5]	2,5-dimethylpyridine				
		$\Delta_{\text{fus}}H$		14.64	259.1	[1994CHI/HOS]
		Δ_vH	(330–471)	44.4	340	EB [1995STE/CHI2]
		Δ_vH	(330–471)	41.9	380	EB [1995STE/CHI2]
		Δ_vH	(330–471)	39.4	420	EB [1995STE/CHI2]
		Δ_vH	(330–471)	36.5	460	EB [1995STE/CHI2]
		Δ_vH	(358–431)	42.8	373	A,MG [1987STE/MAL, 1953HER/MAR, 1973KKY/REP]
C ₇ H ₉ N	[108-48-5]	2,6-dimethylpyridine				
		$\Delta_{\text{fus}}H$		13.04	267.1	[1994CHI/HOS]
		Δ_vH	(342–373)	45.9 ± 2.4	298	CGC [2009LIP/CHI2]
		Δ_vH	(263–353)	45.3	298	[2005BEN/AIT]
		Δ_vH	(323–373)	46.4	298	CGC [1995CHI/HOS]
		Δ_vH	(288–373)	43.7	330	[1995SAK/UEO]
		Δ_vH	(315–457)	43.9	320	EB [1995STE/CHI2]
		Δ_vH	(315–457)	41.4	360	EB [1995STE/CHI2]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound					
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference	
		$\Delta_v H$	(315–457)	38.8	400	EB	[1995STE/CHI2]
		$\Delta_v H$	(315–457)	36.0	440	EB	[1995STE/CHI2]
		$\Delta_v H$	(295–417)	45.0	310	EB	[1990LEN]
		$\Delta_v H$	(267–358)	46.1	282	MM	[1986WIS/LEN]
		$\Delta_v H$		44.4	313	C	[1985MAJ/SVO2]
		$\Delta_v H$		42.5	343	C	[1985MAJ/SVO2]
		$\Delta_v H$		40.8	368	C	[1985MAJ/SVO2]
		$\Delta_v H$	(352–418)	41.6	367	A, MG	[1987STE/MAL, 1953HER/MAR]
C ₇ H ₉ N	[583-58-4]	3,4-dimethylpyridine					
		$\Delta_{\text{fus}} H$		14.7	262.7		[1994CHI/HOS]
		$\Delta_v H$	(341–495)	46.6	360	EB	[1995STE/CHI2]
		$\Delta_v H$	(341–495)	44.2	400	EB	[1995STE/CHI2]
		$\Delta_v H$	(341–495)	41.7	440	EB	[1995STE/CHI2]
		$\Delta_v H$	(341–495)	39.0	480	EB	[1995STE/CHI2]
		$\Delta_v H$	(288–422)	47.6	355		[1995SAK/UEO]
		$\Delta_v H$		48.8	328	C	[1985MAJ/SVO2]
		$\Delta_v H$		47.6	343	C	[1985MAJ/SVO2]
		$\Delta_v H$		45.9	368	C	[1985MAJ/SVO2]
		$\Delta_v H$	(385–454)	44.8	400	A	[1987STE/MAL, 1973KKY/REP]
C ₇ H ₉ N	[591-22-0]	3,5-dimethylpyridine					
		$\Delta_{\text{fus}} H$		13.11	266.9		[1994CHI/HOS]
		$\Delta_v H$	(273–353)	48.5	298		[2005BEN/AIT]
		$\Delta_v H$	(323–373)	48.7	298	CGC	[1995CHI/HOS]
		$\Delta_v H$	(288–392)	47.0	340		[1995SAK/UEO]
		$\Delta_v H$	(335–487)	46.7	340	EB	[1995STE/CHI2]
		$\Delta_v H$	(335–487)	44.3	380	EB	[1995STE/CHI2]
		$\Delta_v H$	(335–487)	41.8	420	EB	[1995STE/CHI2]
		$\Delta_v H$	(335–487)	39.2	460	EB	[1995STE/CHI2]
		$\Delta_v H$	(273–358)	49.1	288	MM	[1986WIS/LEN]
		$\Delta_v H$		49.6	313	C	[1985MAJ/SVO2]
		$\Delta_v H$		46.5	343	C	[1985MAJ/SVO2]
		$\Delta_v H$		44.8	368	C	[1985MAJ/SVO2]
		$\Delta_v H$	(373–446)	44.3	388	A	[1987STE/MAL, 1973KKY/REP]
C ₇ H ₉ N	[536-78-7]	3-ethylpyridine					
		$\Delta_v H$	(334–373)	44.6	349	A	[1987STE/MAL, 1973KKY/REP]
C ₇ H ₉ N	[100-71-0]	2-ethylpyridine					
		$\Delta_v H$		44.7 ± 0.8	298	C	[2003MOR/MIR]
		$\Delta_v H$	(323–373)	43.7	338	A	[1987STE/MAL, 1973KKY/REP]
C ₇ H ₉ N	[536-75-4]	4-ethylpyridine					
		$\Delta_v H$		46.3 ± 0.7	298	C	[2003MOR/MIR]
		$\Delta_v H$	(333–372)	45.3	348	A	[1987STE/MAL, 1973KKY/REP]
C ₇ H ₉ N	[100-61-8]	N-methylaniline					
		$\Delta_v H$	(309–469)	53.6	324	A	[1987STE/MAL]
C ₇ H ₉ N	[95-53-4]	<i>o</i> -toluidine					
		$\Delta_{\text{fus}} H$		11.66	287.6		[1994STE/CHI]
		$\Delta_v H$	(282–313)	57.3 ± 0.2	298	GS	[2005EME/VER2]
		$\Delta_v H$	(290–517)	57.8	300	EB, IP	[1994STE/CHI]
		$\Delta_v H$	(290–517)	54.5	340	EB, IP	[1994STE/CHI]
		$\Delta_v H$	(290–517)	51.5	380	EB, IP	[1994STE/CHI]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound					
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference	
		$\Delta_v H$	(290–517)	48.6	420	EB,IP	[1994STE/CHI]
		$\Delta_v H$	(290–517)	45.7	460	EB,IP	[1994STE/CHI]
		$\Delta_v H$	(290–517)	42.7	500	EB,IP	[1994STE/CHI]
		$\Delta_v H$		62.7 ± 0.5	298		[1990CHA/GAD, 2005EME/VER2]
		$\Delta_v H$	(473–690)	63.1	298		[1957GLA/RUL, 2005EME/VER2]
		$\Delta_v H$	(391–474)	50.0	406	A	[1987STE/MAL, 1949DRE/SHR, 1984BOU/FRI]
		$\Delta_v H$	(313–473)	56.2	298	EB	[1927BER/MAY, 2005EME/VER2]
C ₇ H ₉ N	[108-44-1]	<i>m</i> -toluidine					
		$\Delta_{\text{fus}}H$		8.8	241.7		[1991ACR]
		$\Delta_v H$	(282–313)	58.3 ± 0.4	298	GS	[2005EME/VER2]
		$\Delta_v H$		62.7 ± 0.5	298		[1990CHA/GAD, 2005EME/VER2]
		$\Delta_v H$	(420–439)	59.6 ± 0.3	298	EB	[1990CAB/BEL, 2005EME/VER2]
		$\Delta_v H$	(476–704)	64.1	298		[1957GLA/RUL, 2005EME/VER2]
		$\Delta_v H$	(394–477)	51.1	409	A	[1987STE/MAL, 1949DRE/SHR, 1984BOU/FRI]
		$\Delta_v H$	(313–473)	56.4	298	EB	[1927BER/MAY, 2005EME/VER2]
C ₇ H ₉ N	[106-49-0]	<i>p</i> -toluidine					
		$\Delta_{\text{fus}}H$		17.19	317		[2001CEN/LIP]
		$\Delta_{\text{fus}}H$		17.89	316.5		[1991ACR]
		$\Delta_{\text{fus}}H$		17.3	316.6		[1990MEV/LIC]
		$\Delta_{\text{fus}}H$		18.9	316.9		[1963RAS/NIG]
		$\Delta_{\text{fus}}H$		17.28	315.6		[1889EYK]
		$\Delta_{\text{sub}}H$	(284–313)	76.2 ± 0.3	298	GS	[2005EME/VER2]
		$\Delta_{\text{sub}}H$		78.8 ± 0.5	298		[1990CHA/GAD]
		$\Delta_v H$	(319–345)	57.8 ± 0.3	298	GS	[2005EME/VER2]
		$\Delta_v H$	(393–474)	51.1	408	A	[1987STE/MAL]
		$\Delta_v H$	(474–641)	62.1	298		[1957GLA/RUL, 2005EME/VER2]
		$\Delta_v H$	(315–473)	54.9	330		[1947STU]
		$\Delta_v H$	(313–473)	55.9	298	EB	[1927BER/MAY, 2005EME/VER2]
C ₇ H ₉ N	[1855-63-6]	1-cyclohexene-1-carbonitrile					
		$\Delta_v H$		53.6 ± 0.1	298	C	[1970PRO/KRE]
C ₇ H ₉ N	[31357-72-9]	bicyclo[3.1.0]hexane-1-carbonitrile					
		$\Delta_v H$	(366–444)	U43.2	382	BG	[1971HAL/BAL]
C ₇ H ₉ NO	[90-04-0]	2-methoxyaniline					
		$\Delta_v H$	(334–492)	57.5	349	A	[1987STE/MAL, 1947STU]
C ₇ H ₉ NO	[104-94-9]	4-methoxyaniline					
		$\Delta_{\text{fus}}H$		18.53	NA	DSC	[2008SIN/DAS]
C ₇ H ₉ NO	[3718-65-8]	3,5-dimethylpyridine N-oxide					
		$\Delta_{\text{sub}}H$		100.9 ± 2.3	298	C	[2010CAB/MON]
C ₇ H ₉ N ₅	[87578-82-3]	8,9-dimethyladenine					
		$\Delta_{\text{sub}}H$	(369–374)	105.8 ± 0.8	361	ME	[1987KAM/ZIE]
C ₇ H ₉ N ₅	[76470-20-7]	2,9-dimethyladenine					
		$\Delta_{\text{sub}}H$	(359–364)	123.5	371		[1992KAM]
C ₇ H ₉ N ₅	[938-55-6]	N,N-dimethyladenine					
		$\Delta_{\text{sub}}H$	(379–409)	115.5 ± 2.1		ME	[1984ZIE/ZIE]
C ₇ H ₉ N ₅	[2009-52-1]	N,9-dimethyladenine					
		$\Delta_{\text{sub}}H$	(336–369)	115.5 ± 1.7		ME	[1984ZIE/ZIE]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₇ H ₉ N ₅ O ₁₂	[34001-49-5]	2,2,2-trinitroethyl-4,4-dinitropentanoate				
	$\Delta_{\text{us}}H$		20.08	363.8		
	$\Delta_{\text{fus}}H$		6.69	366.7	DSC	[1971ROS/HOL]
C ₇ H ₉ N ₅ O ₁₂	[2555-56-8]	2,2-dinitropropyl-4,4,4-trinitrobutyrate				
	$\Delta_{\text{us}}H$		25.94	284.2		
	$\Delta_{\text{fus}}H$		6.69	368.2	DSC	[1971ROS/HOL]
C ₇ H ₁₀	[498-66-8]	bicyclo[2.2.1]hept-2-ene (norbornene)				
	$\Delta_{\text{us}}H$		4.37	130.3		
	$\Delta_{\text{fus}}H$		3.48	319.5		[1992LEB/SMI, 1992SMI/LEB]
	$\Delta_{\text{us}}H$		4.37	130		[1992LEB/SMI]
	$\Delta_{\text{sub}}H$		37.8 ± 0.14	298	C	[1982JOC/DEK]
	$\Delta_{\text{sub}}H$		37.7 ± 0.9	298	BG	[1978STE2]
	$\Delta_{\text{sub}}H$		38.7 ± 0.5	298	C	[1976KOZ/BYC]
	$\Delta_{\text{sub}}H$		33.6 ± 0.08			[1973HAL/SMI]
	Δ_vH	(338–406)	35.1 ± 0.2	298	EB	[1996STE/CHI2]
	Δ_vH	(338–406)	33.0 ± 0.2	340	EB	[1996STE/CHI2]
	Δ_vH	(338–406)	30.8 ± 0.3	380	EB	[1996STE/CHI2]
C ₇ H ₁₀	[16554-83-9]	bicyclo[4.1.0]hept-3-ene				
	Δ_vH	(333–384)	36.7	348	A	[1987STE/MAL]
	Δ_vH		38.4 ± 0.6	298	EB	[1974VAR/DRU]
C ₇ H ₁₀	[279-19-6]	tricyclo[2.2.1.0 ^{2,6}]heptane				
	$\Delta_{\text{sub}}H$		38.7 ± 0.7	298	BG	[1978STE2]
	$\Delta_{\text{sub}}H$		39.2 ± 1.1	298	C	[1976KOZ/BYC]
	Δ_vH		38.5	298		[2008OSM/CAT]
C ₇ H ₁₀	[187-26-8]	tricyclo[4.1.0.0 ^{2,4}]heptane				
	Δ_vH		36.5 ± 0.5	298	EB	[1974VAR/DRU]
C ₇ H ₁₀	[na]	tricyclo[4.1.0.0 ^{2,6}]heptane				
	Δ_vH	(322–373)	35.3	337	A	[1987STE/MAL]
C ₇ H ₁₀	[33475-22-8]	dispiro[2.0.2.1]heptane				
	Δ_vH		35.1 ± 0.5	298		[2008OSM/CAT]
C ₇ H ₁₀ ClN ₃ O ₃	[16773-42-5]	1-(2-hydroxy-3-chloropropyl)-2-methyl-5-nitroimidazole (ornidazole)				
	$\Delta_{\text{fus}}H$		21.38	358.6		[2004WAN/TAN2]
C ₇ H ₁₀ N ₂	[14667-55-1]	2,3,5-trimethylpyrazine				
	Δ_vH		53.9 ± 1.6	298	C	[1996RIB/MOR]
C ₇ H ₁₀ N ₂	[538-08-9]	diallycyanamide				
	Δ_vH	(369–495)	52.3	384	A	[1987STE/MAL]
C ₇ H ₁₀ N ₂	[646-20-8]	1,5-dicyanopentane				
	Δ_vH	(306–331)	74.5	318	A	[1987STE/MAL]
C ₇ H ₁₀ N ₂	[95-80-7]	2,4-diaminotoluene				
	Δ_vH	(379–553)	67.7	394	A	[1987STE/MAL, 1947STU]
C ₇ H ₁₀ N ₂	[539-44-6]	4-tolyhydrazine				
	Δ_vH	(355–515)	65.4	370	A	[1987STE/MAL, 1947STU]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₇ H ₁₀ N ₂	[4210-60-0] $\Delta_{\text{sub}}H$	α - <i>tert</i> -butylmalononitrile (293–323)	59.8 ± 0.7	298		[1990BEC/DOG]
C ₇ H ₁₀ N ₂	[646-20-8] $\Delta_{\text{fus}}H$	pimelonitrile	15.0	241.7	DSC	[2007BAD/BLA]
C ₇ H ₁₀ N ₂	[1122-58-3] $\Delta_{\text{fus}}H$	4-N,N-dimethylaminopyridine (80–402)	21.63	387.1	AC	[2007SHI/TAN]
C ₇ H ₁₀ N ₂ O	[na] $\Delta_{\text{us}}H$ $\Delta_{\text{fus}}H$	6,7-diazatricyclo[3.2.2.0 ^{2,4}]non-6-ene-N-oxide	15.8 2.6	372.6 411.4		[1980BYS]
C ₇ H ₁₀ N ₂ O ₂	[4401-71-2] $\Delta_{\text{sub}}H$	1,3-dimethylthymine (313–363)	109.2 ± 2.1	338	QR	[1980TEP/YAN]
C ₇ H ₁₀ N ₂ O ₂	[4401-71-2] $\Delta_{\text{fus}}H$ $\Delta_{\text{sub}}H$	1,3,5-trimethyluracil (321–331)	16.1 103.5 ± 1.5	428.7 326		[1996KAM/ZIE] [1996KAM/ZIE]
C ₇ H ₁₀ N ₂ O ₂	[13509-52-9] $\Delta_{\text{fus}}H$ $\Delta_{\text{sub}}H$	1,3,6-trimethyluracil (300–340)	21.2 106.7 ± 2.5	384.5 320		[1996DOM/HEA] [1980TEP/YAN]
C ₇ H ₁₀ N ₂ O ₂	[na] $\Delta_{\text{fus}}H$	N-acetyl-(<i>l</i>)-alanine amide	21.7	431		[1988FER/DEL]
C ₇ H ₁₀ O	[17356-19-3] Δ_vH	1-ethynyl-1-cyclopentanol (323–373)	62.1	298	CGC	[1995CHI/HOS]
C ₇ H ₁₀ O	[10218-02-7] $\Delta_{\text{sub}}H$ Δ_vH	7-norbornanone (300–340) (322–348)	47.3 ± 2.2 47.9	298 335	BG EB	[1978STE] [1994WIB/MOR]
C ₇ H ₁₀ O	[497-38-1] $\Delta_{\text{fus}}H$ $\Delta_{\text{sub}}H$ Δ_vH Δ_vH Δ_vH	2-norbornanone (300–340) (343–383) (343–383)	3.39 49.0 ± 1.7 50.0 51.5 49.6	368.7 298 298 298 298		[1993ACR] [1978STE] [2002VAN/PAR] [1995CHI/HOS] [1995CHI/HOS]
C ₇ H ₁₀ O ₂	[20583-46-4] Δ_vH	5-methyl-5-hexene-2,4-dione (323–363)	26.4	338	A	[1987STE/MAL, 1973KKY/REP]
C ₇ H ₁₀ O ₂	[4350-84-9] $\Delta_{\text{sub}}H$	2-oxabicyclo[2.2.2]octan-3-one	69.6 ± 21			[1980AND/PIL]
C ₇ H ₁₀ O ₃	[815-68-9] Δ_vH	3-acetyl-2,4-pentanedione (369–477)	54.9	384	A	[1987STE/MAL]
C ₇ H ₁₀ O ₃	[106-91-2] Δ_vH Δ_vH	glycidyl methacrylate	61.2 ± 0.4 60.6 ± 0.9	298 298	A C	[1987VAN/KAC] [1986YER/WOR2]
C ₇ H ₁₀ O ₃	[281-32-3] $\Delta_{\text{sub}}H$	2,4,10-trioxaadamantane	74.4 ± 0.4	298	C	[1974MAN2]
C ₇ H ₁₀ O ₃	[na]	3,3-dimethylpentanedioic anhydride				

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound					
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference	
		$\Delta_{\text{fus}}H$	17.99	396.2		[1974BOR]	
C ₇ H ₁₀ O ₃	[35046-67-4]	trimethylsuccinic anhydride					
		$\Delta_{\text{sub}}H$	74.1 ± 4.2			[1954JOR, 1970COX/PIL]	
		Δ_vH	(326–504)	52.9	341	A	[1987STE/MAL, 1947STU]
C ₇ H ₁₀ O ₄	[617-54-9]	dimethyl citraconate					
		Δ_vH	(324–484)	55.8	339	A	[1987STE/MAL, 1947STU]
C ₇ H ₁₀ O ₄	[617-52-7]	dimethyl itaconate					
		Δ_vH	(342–481)	67.0	357	A	[1987STE/MAL, 1947STU]
C ₇ H ₁₀ O ₄	[617-53-8]	dimethyl mesaconate					
		Δ_vH	(319–479)	55.2	334	A	[1987STE/MAL, 1947STU]
C ₇ H ₁₀ O ₆	[na]	tris(methoxycarbonyl)methane					
		$\Delta_{\text{fus}}H$	18.2	301.2		[1995RAK/VER]	
		Δ_vH	(308–348)	74.4 ± 0.6		GS	[1995RAK/VER]
C ₇ H ₁₀ S	[1551-27-5]	2-propylthiophene					
		Δ_vH	43.7 ± 1.0	298		C	[2007RIB/SAN]
		Δ_vH	(243–303)	46.0	273		[1981EDW/PRA, 1999DYK/SVO]
C ₇ H ₁₀ S	[4095-22-1]	2-isopropylthiophene					
		Δ_vH	(352–468)	41.5	367		[1999DYK/SVO]
C ₇ H ₁₀ S ₃	[2164-87-6]	4,5-tetramethylene-1,3-dithiolan-2-thione					
		$\Delta_{\text{sub}}H$	(353–369)	99.0	360		[1967GEI/SCH, 1970COX/PIL]
		$\Delta_{\text{sub}}H$		103.9 ± 2.9	298		[1967GEI/SCH, 1970COX/PIL]
C ₇ H ₁₁ BrO ₂	[26918-14-9]	4-bromo-3-methylcrotonic acid, ethyl ester					
		Δ_vH	(346–381)	43.1	361	A	[1987STE/MAL]
C ₇ H ₁₁ ClO ₅	[na]	(2-chloroethyl)[(1-methoxycarbonyl)ethyl] carbonate					
		Δ_vH	(365–525)	66.8	380	A	[1987STE/MAL, 1999DYK/SVO]
C ₇ H ₁₁ Cl ₃ O ₂	[57392-56-0]	trichloroacetic acid, neopentyl ester					
		Δ_vH	(378–473)	57.7	393	A	[1987STE/MAL, 1999DYK/SVO]
C ₇ H ₁₁ N	[766-05-2]	cyclohexanecarbonitrile					
		$\Delta_{\text{us}}H$	7.43	215			
		$\Delta_{\text{fus}}H$	3.64	285.1		[1996DOM/HEA]	
		Δ_vH	(333–427)	39.4	351	BG	[1971HAL/BAL]
		Δ_vH		51.9 ± 0.1	298	C	[1970PRO/KRE]
C ₇ H ₁₁ N	[931-53-3]	isocyanocyclohexane					
		$\Delta_{\text{fus}}H$	3.98	277.7		DSC	[2008SIN/MUR2]
		$\Delta_{\text{us}}H$	6.18	192.6			
		$\Delta_{\text{fus}}H$	4.23	279.6		[1996DOM/HEA]	
C ₇ H ₁₁ NO ₂	[na]	2-methyl-2-acetoxybutyronitrile					
		Δ_vH	(315–469)	58.1	330	A	[1987STE/MAL, 1947STU]
C ₇ H ₁₁ NO ₂	[na]	5-oxo-2-pyrrolidinecarboxylic acid, ethyl ester					
		Δ_vH	(418–511)	73.7	433	A	[1987STE/MAL]
C ₇ H ₁₁ N ₃ O	[2228-27-5]	1,N,N-trimethylcytosine					
		$\Delta_{\text{sub}}H$	110.9 ± 1.7			[1998ZIE/WSZ]	
C ₇ H ₁₁ N ₃ O	[25307-94-2]	1,5,N-trimethylcytosine					

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₇ H ₁₁ N ₃ O ₂	$\Delta_{\text{sub}}H$	(396–431)	108.0 ± 2.0		GS	[1998ZIE/WSZ]
	[na] $\Delta_{\text{sub}}H$	1,5-dimethyl-N-methoxycytosine (327–365)	95.6 ± 0.7		GS	[1998ZIE/WSZ]
C ₇ H ₁₁ N ₅ O ₁₀	[242800-94-8] $\Delta_{\text{sub}}H$	1,1,1,4,4-pentanitro-2,2-dimethylpentane	103.8	298		[1999MIR/VOR]
C ₇ H ₁₂	[279-23-2] $\Delta_{\text{fus}}H$	bicyclo[2.2.1]heptane (norbornane)	4.45	360.8	DSC	[2004VER/EME]
	$\Delta_{\text{sub}}H$	(278–308)	40.3 ± 0.4	293	GS	[2004VER/EME]
	$\Delta_{\text{sub}}H$	(278–308)	40.1 ± 0.4	298	GS	[2004VER/EME]
	$\Delta_{\text{sub}}H$		40.0 ± 0.1	298	C	[1987AN/ZHU]
	$\Delta_{\text{sub}}H$		40.3 ± 0.32	298	C	[1982JOC/DEK]
	$\Delta_{\text{sub}}H$		40.4 ± 0.8			[1978STE2]
	$\Delta_{\text{sub}}H$	(284–326)	40.0 ± 0.8	305	TSGC	[1975CLA/KNO]
	$\Delta_{\text{sub}}H$		39.33 ± 0.13			[1973HAL/SMI]
C ₇ H ₁₂	[286-08-8] Δ_vH	<i>cis</i> bicyclo[4.1.0]heptane	40.6 ± 0.2	298		[2008OSM/CAT]
	Δ_vH	(298–385)	38.0 ± 0.8	313	A	[1987STE/MAL, 1970CHA/MCN]
	[286-08-8] Δ_vH	(<i>dl</i>) bicyclo[4.1.0]heptane (333–385)	36.5	348	A	[1987STE/MAL]
C ₇ H ₁₂	[4625-24-5] Δ_vH	1-methylbicyclo[3.1.0]hexane (312–362)	34.0	327	A	[1987STE/MAL]
C ₇ H ₁₂	[628-92-2] $\Delta_{\text{trs}}H$	cycloheptene	5.28	154		
	$\Delta_{\text{trs}}H$		0.71	210		
	$\Delta_{\text{fus}}H$		0.97	217		[1996DOM/HEA, 1994LEB/SMI]
	Δ_vH	(251–313)	38.5	266	A	[1987STE/MAL, 1941LIS]
	Δ_vH	(251–312)	36.7	300		[1941LIS]
C ₇ H ₁₂	[765-47-9] Δ_vH	1,2-dimethylcyclopentene (294–431)	36.4	309	A	[1987STE/MAL, 1973KKY/REP]
C ₇ H ₁₂	[62184-82-1] Δ_vH	(<i>dl</i>) 1,3-dimethylcyclopentene (283–410)	35.0	298	A	[1987STE/MAL, 1973KKY/REP]
C ₇ H ₁₂	[57426-81-0] Δ_vH	(<i>dl</i>) 1,4-dimethylcyclopentene (273–413)	35.1	288	A	[1987STE/MAL, 1973KKY/REP]
C ₇ H ₁₂	[16491-15-9] Δ_vH	(<i>dl</i>) 1,5-dimethylcyclopentene (273–423)	37.1	288	A	[1987STE/MAL, 1973KKY/REP]
C ₇ H ₁₂	[na] Δ_vH	1-ethylidenecyclopentane	38.6 ± 0.2	298	GCC	[1979FUC/PEA]
C ₇ H ₁₂	[2146-38-5] Δ_vH	1-ethylcyclopentene	38.5 ± 0.3	298	GCC	[1979FUC/PEA]
	Δ_vH	(293–433)	36.5	308	A	[1987STE/MAL, 1973KKY/REP]
	[694-35-9] Δ_vH	3-ethylcyclopentene (288–435)	36.5	303	A	[1987STE/MAL, 1973KKY/REP]
C ₇ H ₁₂	[3742-38-9] Δ_vH	4-ethylcyclopentene (288–435)	36.5	303	A	[1987STE/MAL, 1973KKY/REP]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₇ H ₁₂	[591-49-1]	1-methyl-1-cyclohexene				
	$\Delta_v H$	(275–313)	37.7 ± 0.2	294	GS	[2000VER/WAN]
	$\Delta_v H$	(275–313)	37.5 ± 0.2	298	GS	[2000VER/WAN]
	$\Delta_v H$	(333–384)	35.7	348	A	[1987STE/MAL, 1970EIS/ORA, 1984BOU/FRI]
C ₇ H ₁₂	[591-48-0]	<i>(dl)</i> 3-methyl-1-cyclohexene				
	$\Delta_v H$	(335–376)	34.8	350	A	[1987STE/MAL, 1970EIS/ORA, 1984BOU/FRI]
C ₇ H ₁₂	[591-47-9]	<i>(dl)</i> 4-methyl-1-cyclohexene				
	$\Delta_{\text{fus}} H$		6.63	153.6		[1994LEB/SMI]
	$\Delta_v H$	(275–296)	37.0 ± 0.6	286	GS	[2000VER/WAN]
	$\Delta_v H$	(275–296)	36.3 ± 0.6	298	GS	[2000VER/WAN]
C ₇ H ₁₂	[1192-37-6]	methylenecyclohexane				
	$\Delta_v H$		36.1 ± 0.3	298	GCC	[1979FUC/PEA]
	$\Delta_v H$	(331–387)	34.4	346	A,EB	[1987STE/MAL, 1973MEY/HOT]
C ₇ H ₁₂	[628-71-7]	1-heptyne				
	$\Delta_v H$	(336–373)	37.9	351	A	[1987STE/MAL, 1970EIS/ORA, 1984BOU/FRI]
C ₇ H ₁₂	[1119-65-9]	2-heptyne				
	$\Delta_v H$	(346–385)	38.6	361	A	[1987STE/MAL, 1970EIS/ORA, 1984BOU/FRI]
C ₇ H ₁₂	[2586-89-2]	3-heptyne				
	$\Delta_v H$	(343–380)	39.1	358	A	[1987STE/MAL, 1970EIS/ORA, 1984BOU/FRI]
C ₇ H ₁₂ Br ₂	[29974-68-3]	1,2-dibromocycloheptane				
	$\Delta_v H$	(292–353)	50.3	307	A	[1987STE/MAL, 1941LIS, 1973KKY/REP]
C ₇ H ₁₂ ClNO	[13654-91-6]	6-chlorohexylisocyanate				
	$\Delta_v H$	(363–453)	52.5	378	A	[1987STE/MAL, 1968ZHU/KON, 1973KKY/REP]
C ₇ H ₁₂ CIN ₅	[122-34-9]	2-chloro-4,6-bis(ethylamino)-s-triazine (Simazin)				
	$\Delta_{\text{fus}} H$		47.35	502.5	DSC	[1990DON/DRE]
	$\Delta_{\text{sub}} H$	(323–403)	130.8	338	GS-GC	[1987STE/MAL, 1964FRI/SAM]
C ₇ H ₁₂ Cl ₂ O ₂	[na]	dichloroacetic acid, neopentyl ester				
	$\Delta_v H$	(368–463)	57.4	383	A	[1987STE/MAL, 1999DYK/SVO]
C ₇ H ₁₂ Cl ₂ S	[na]	(2-chloroethyl)(2-chlorocyclopentyl) sulfide				
	$\Delta_v H$	(273–333)	65.9	303	A,GS	[1987STE/MAL, 1948RED/CHA, 1999DYK/SVO]
C ₇ H ₁₂ Cl ₄	[3922-36-9]	1,1,1,7-tetrachloroheptane				
	$\Delta_v H$	(342–455)	71.7	357		[1999DYK/SVO]
	$\Delta_v H$	(370–454)	69.9	385	A	[1987STE/MAL]
C ₇ H ₁₂ N ₂	[3010-03-5]	1-piperidinoacetonitrile				
	$\Delta_{\text{fus}} H$		17.57	293.2		[1997WEL/VER]
	$\Delta_v H$	(303–338)	56.0 ± 0.5		GS	[1997WEL/VER]
C ₇ H ₁₂ N ₄ O ₁₀	[5917-61-3]	<i>bis</i> (2,4-dinitropropyl)formal				
	$\Delta_v H$	(333–383)	84.8 ± 0.9	358		[2007RAU/BEH]
C ₇ H ₁₂ O	[497-37-0]	exo-norborneol				
	$\Delta_v H$		52.5	298	GC	[2002VAN/PAR]
C ₇ H ₁₂ O	[931-57-7]	1-methoxycyclohexene				
	$\Delta_v H$	(274–313)	44.0 ± 0.2	294	GS	[1998VER/WEL]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₇ H ₁₂ O	$\Delta_v H$	(274–313)	43.7 ± 0.2	298	GS	[1998VER/WEL]
	[502-42-1]	cycloheptanone				
	$\Delta_{\text{fus}}H$		12.4	227		
	$\Delta_{\text{fus}}H$		0.43	232.6		
	$\Delta_{\text{fus}}H$		1.39	259.3		[1998GON/SZW]
	$\Delta_v H$	(343–383)	50.6	298	CGC	[1995CHI/HOS]
	$\Delta_v H$	(343–383)	51.9	298	CGC	[1995CHI/HOS]
	$\Delta_v H$	(343–383)	50.7	298	CGC	[1995CHI/HOS]
	$\Delta_v H$	(313–453)	48.5	328	A	[1987STE/MAL]
C ₇ H ₁₂ O	$\Delta_v H$	(373–465)	44.8	388	A,EB	[1987STE/MAL, 1976MEY/HOT]
	$\Delta_v H$		49.5 ± 0.6	298		[1972WOL]
C ₇ H ₁₂ O	[583-60-8]	2-methylcyclohexanone				
	$\Delta_v H$	(339–437)	44.0	298	EB	[2006PAL/ORO]
	$\Delta_v H$	(331–437)	45.0	346		[1993BRU/MON]
C ₇ H ₁₂ O	[591-24-2]	3-methylcyclohexanone				
	$\Delta_v H$	(334–441)	44.9	349		[1993AUC/MON]
C ₇ H ₁₂ O	[589-92-4]	4-methylcyclohexanone				
	$\Delta_v H$	(339–444)	45.3	354		[1993AUC/MON]
C ₇ H ₁₂ O ₂	[na]	cyclobutanecarboxylic acid ethyl ester				
	$\Delta_v H$	(274–308)	44.9 ± 0.4		GS	[1998VER/KUM]
C ₇ H ₁₂ O ₂	[176-32-9]	1,4-dioxaspiro[4.4]nonane				
	$\Delta_v H$	(278–313)	47.6 ± 0.5	298	GS	[1998VER/PEN, 2002VER]
C ₇ H ₁₂ O ₂	[141-32-2]	butyl acrylate				
	$\Delta_{\text{fus}}H$		17.31	209.5		[1996DOM/HEA]
	$\Delta_v H$	(318–419)	47.3 ± 0.3	298	EB	[1996STE/CHI2]
	$\Delta_v H$	(318–419)	45.7 ± 0.3	320	EB	[1996STE/CHI2]
	$\Delta_v H$	(318–419)	42.8 ± 0.3	360	EB	[1996STE/CHI2]
	$\Delta_v H$	(318–419)	40.0 ± 0.3	400	EB	[1996STE/CHI2]
	$\Delta_v H$	(272–421)	44.8	287	A	[1987STE/MAL, 1947STU]
C ₇ H ₁₂ O ₂	[106-63-8]	isobutyl acrylate				
	$\Delta_v H$	(330–410)	43.8	345	A	[1987STE/MAL]
C ₇ H ₁₂ O ₂	[539-87-7]	heptanolactone				
	$\Delta_v H$	(368–390)	48.2 ± 0.3	379	MM	[1991WIB/WAL]
	$\Delta_v H$	(369–390)	53.3 ± 1.3	298	MM	[1991WIB/WAL]
C ₇ H ₁₂ O ₂	[105-21-5]	γ -heptalactone				
	$\Delta_v H$	(298–363)	62.3 ± 0.3	298	GS	[2008EME/KOZ]
C ₇ H ₁₂ O ₂	[2210-28-8]	propyl methacrylate				
	$\Delta_v H$	(304–413)	41.6	319	A	[1987STE/MAL]
C ₇ H ₁₂ O ₂	[98-89-5]	cyclohexanecarboxylic acid				
	$\Delta_{\text{fus}}H$		9.2	301.9		[2008DOM/MOR]
C ₇ H ₁₂ O ₂	[4351-54-6]	cyclohexyl formate				
	$\Delta_{\text{fus}}H$		10.49	201.3		[1999KAB/KOZ]
	$\Delta_v H$	(307–434)	47.1	322		[2005STE/SUN]
	$\Delta_v H$		49.3 ± 0.2	298	C	[2004PAU/ZAI, 2003ZAI/VER]
	$\Delta_v H$	(243–273)	52.0 ± 1.3	298	ME	[2003ZAI/VER]
	$\Delta_v H$	(243–273)	49.5 ± 1.2	298	ME	[2003ZAI/VER]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound					
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference	
C ₇ H ₁₂ O ₂	[638-10-8]	ethyl 3-methylbut-2-enoate					
	$\Delta_v H$	(274–319)	49.3 ± 0.2	298	GS	[2008EME/TOK]	
C ₇ H ₁₂ O ₃	[2461-40-7]	glycidyl butyrate					
	$\Delta_v H$		58.7 ± 0.4	298		[1987VAN/KAC]	
	$\Delta_v H$		58.0 ± 0.4	298	C	[1986YER/WOR2]	
C ₇ H ₁₂ O ₃	[10235-71-9]	2-acetoxy-2-methyl-3-butanone					
	$\Delta_v H$	(337–368)	54.8	352	A	[1987STE/MAL]	
C ₇ H ₁₂ O ₃	[539-88-8]	ethyl levulinate					
	$\Delta_v H$	(320–480)	58.3	335	A	[1987STE/MAL, 1947STU]	
	$\Delta_v H$		51.6	420		[1931SCH/COW]	
C ₇ H ₁₂ O ₃	[27761-61-1]	1,4-dimethyl-2,6,7-trioxabicyclo[2.2.2]octane					
	$\Delta_{\text{fus}} H$		18.0	370.2		[1995RAK/VER2]	
	$\Delta_{\text{sub}} H$		74.9	298		[1995RAK/VER2]	
C ₇ H ₁₂ O ₄	[2985-28-6]	2-acetoxypropionic acid, ethyl ester					
	$\Delta_v H$	(313–454)	57.9	328	A	[1987STE/MAL]	
C ₇ H ₁₂ O ₄	[40326-37-2]	3-acetoxypropionic acid, ethyl ester					
	$\Delta_v H$	(350–367)	72.1	358	A	[1987STE/MAL, 1973KKY/REP]	
C ₇ H ₁₂ O ₄	[623-84-7]	1,2-propylene glycol diacetate					
	$\Delta_v H$	(318–367)	54.9	323		[2001HOR/GAR]	
C ₇ H ₁₂ O ₄	[105-53-3]	diethyl malonate					
	$\Delta_v H$	(288–318)	64.7 ± 0.2	293	GS	[1992VER/BEC]	
			Note: Steele and coworkers in reference [2002STE/CHI6] refer to a personal communication with one of the authors of [1992VER/BEC]—stating that it was established that the compound studied was not diethyl malonate.				
	$\Delta_v H$	(293–318)	63.3	305	A	[1987STE/MAL]	
	$\Delta_v H$	(384–468)	59.9	399	A	[1987STE/MAL]	
	$\Delta_v H$	(313–472)	51.2	328	A	[1987STE/MAL, 1947STU]	
	C ₇ H ₁₂ O ₄	[6065-54-9]	dimethyl dimethylmalonate				
		$\Delta_v H$	(278–307)	55.6 ± 0.8	293	GS	[1992VER/BEC]
	C ₇ H ₁₂ O ₄	[1119-40-0]	glutaric acid, dimethyl ester				
		$\Delta_v H$	(283–348)	65.7 ± 0.4	298	GS	[2006VER/KOZ]
$\Delta_v H$		(366–483)	54.7	381	A	[1987STE/MAL]	
$\Delta_v H$		(388–483)	66.1	298	EB	[1963VLA/GRA, 2006VER/KOZ]	
C ₇ H ₁₂ O ₄	[627-91-8]	methyl adipate					
	$\Delta_v H$	(453–503)	82.9	468	A	[1987STE/MAL]	
C ₇ H ₁₂ O ₄	[111-16-0]	heptanedioic acid (pimelic acid)					
	$\Delta_{\text{us}} H$		1.5	337.7			
	$\Delta_{\text{fus}} H$		23.7	368.2		[2005ROU/TEM]	
	$\Delta_{\text{fus}} H$		27.62	377.5		[1993ACR]	
	$\Delta_{\text{sub}} H$	(328–363)	153 ± 4		TPD	[2007CAP/LOV]	
	$\Delta_{\text{sub}} H$	(283–300)	124		TPTD	[2005CHA/ZIE]	
	$\Delta_{\text{sub}} H$	(318–336)	80.8		TPTD	[2005CHA/ZIE]	
	$\Delta_{\text{sub}} H$	(288–308)	178		TPTD	[2001CHA/TOB]	
		Note: Values based on the TPTD method are not consistent with values determined by other experimental methods					
	$\Delta_{\text{sub}} H$	(358–371)	136.61.0	365	ME	[1999RIB/MON]	
	$\Delta_{\text{sub}} H$		139.9 ± 1.0	298		[1999RIB/MON]	

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(424–503)	112.0 ± 0.8	298	CGC	[2005ROU/TEM]
	$\Delta_v H$	(436–615)	88.6	451	A	[1987STE/MAL, 1947STU]
C ₇ H ₁₂ O ₄	[126-54-5]	2,4,8,10-tetraoxaspiro[5.5]undecane				
	$\Delta_v H$		56.0			[1959FLE/MOR]
C ₇ H ₁₂ O ₄	[534-59-8]	butylmalonic acid				
	$\Delta_{\text{sub}} H$		124.6 ± 2.3	298	ME	[2000RIB/MON]
C ₇ H ₁₂ O ₄ S ₂	[na]	<i>(dl)</i> methylenebisthiopropionic acid				
	$\Delta_{\text{fus}} H$		39.33	429		[1976LEC/COL]
C ₇ H ₁₂ O ₄ S ₂	[na]	<i>(d)</i> methylenebisthiopropionic acid				
	$\Delta_{\text{fus}} H$		22.59	355		[1976LEC/COL]
C ₇ H ₁₂ O ₅	[na]	ethyl[(1-methoxycarbonyl)ethyl]carbonate				
	$\Delta_v H$	(343–473)	60.0	358	A	[1987STE/MAL]
C ₇ H ₁₂ O ₅	[na]	2-(lactyloxy)propionic acid, methyl ester				
	$\Delta_v H$	(317–384)	72.0	332	A	[1987STE/MAL]
C ₇ H ₁₃ Cl	[932-78-2]	1-chloro-1-methylcyclohexane				
	$\Delta_{\text{us}} H$		9.38	214.4		
	$\Delta_{\text{fus}} H$		1.63	234.5		[1998KAB/BLO]
C ₇ H ₁₃ ClO	[2528-61-2]	heptanoyl chloride				
	$\Delta_v H$	(307–418)	63.7	322	A	[1987STE/MAL, 1947STU]
C ₇ H ₁₃ ClO ₂	[na]	chloroacetic acid, neopentyl ester				
	$\Delta_v H$	(378–448)	55.6	393	A	[1987STE/MAL, 1999DYK/SVO]
C ₇ H ₁₃ F ₃ O ₃	[2339-51-7]	<i>tris</i> (2-fluoroethyl)orthoformate				
	$\Delta_v H$	(273–333)	59.7	288	A	[1987STE/MAL, 1999DYK/SVO]
C ₇ H ₁₃ N	[na]	2,2-dimethylpentanenitrile				
	$\Delta_v H$	(274–303)	46.9 ± 0.4		GS	[1994RAK/VER]
C ₇ H ₁₃ N	[629-08-3]	heptanonitrile				
	$\Delta_v H$	(280–307)	51.9 ± 0.3	298	GS	[2005EME/VER]
	$\Delta_v H$	(313–473)	46.0	328	A	[1987STE/MAL]
	$\Delta_v H$		51.9 ± 0.8	298	EB	[1973LEB/KAT, 2005EME/VER]
	$\Delta_v H$	(294–457)	46.4	309		[1947STU]
	$\Delta_v H$	(313–473)	49.1	298	EB	[1941RAL/SEL, 2005EME/VER]
	$\Delta_v H$	(314–472)	51.3 ± 0.3	298	MM	[1933HEI, 2005EME/VER]
C ₇ H ₁₃ N	[100-76-5]	1-azabicyclooctane				
	$\Delta_{\text{us}} H$		5.23	196		
	$\Delta_{\text{fus}} H$		5.86	430		[1996DOM/HEA]
	$\Delta_{\text{sub}} H$		50.8 ± 0.4			[1971WON/WES, 1977PED/RYL]
	$\Delta_{\text{sub}} H$	(273–362)	50.8 ± 0.2	298		[1948BRO/SUI, 1970COX/PIL, 1960JON]
C ₇ H ₁₃ NO	[2556-73-2]	N-methylcaprolactam				
	$\Delta_v H$	(340–400)	49.4	370		[1984SHC/KAP]
C ₇ H ₁₃ NO	[na]	2-butoxypropionitrile				
	$\Delta_v H$	(373–423)	46.7	388	A	[1987STE/MAL, 1973KKY/REP]
C ₇ H ₁₃ NO	[162047-91-8]	2-methoxy-3,3-dimethylbutanenitrile				
	$\Delta_v H$	(295–324)	58.8 ± 1.1	298	GS	[1995VER/BEC]
C ₇ H ₁₃ NO	[162047-90-7]	2-methoxy-2-methylpentanenitrile				

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(278–308)	48.5 ± 0.6	298	GS	[1995VER/BEC]
C ₇ H ₁₃ NO	[na]	<i>trans</i> -6-heptenoic acid amide				
	$\Delta_{\text{sub}} H$	(362–393)	97.2	377	A	[1987STE/MAL]
C ₇ H ₁₃ NO	[673-66-5]	ζ -enantholactam				
	$\Delta_{\text{fus}} H$		13.78	310.3		[1996DOM/HEA]
C ₇ H ₁₃ NO	[3612-18-8]	1-ethyl-4-piperidone				
	$\Delta_v H$		56.7 ± 0.6	298	C	[2006RIB/CAB]
C ₇ H ₁₃ NO ₂	[na]	lactic acid N-(methallyl) amide				
	$\Delta_v H$	(360–428)	81.8	375	A	[1987STE/MAL]
C ₇ H ₁₃ NO ₂	[na]	N-lactylmorpholine				
	$\Delta_v H$	(371–423)	62.7	386	A	[1987STE/MAL]
C ₇ H ₁₃ NO ₃	[5143-72-6]	<i>dl</i> N-acetylalanine ethyl ester				
	$\Delta_v H$	(372–460)	65.2	387	A	[1987STE/MAL, 1973KKY/REP]
C ₇ H ₁₃ N ₃ O ₃ S	[23135-22-0]	N,N-dimethyl-2-methylcarbomoyloxymino-2-(methylthio)acetamide				
	$\Delta_{\text{fus}} H$		30.17	372.2	DSC	[1990DON/DRE]
C ₇ H ₁₃ O ₆ P	[7786-34-7]	mevinphos				
	$\Delta_v H$	(293–383)	68.1	308	A	[1987STE/MAL]
C ₇ H ₁₄	[291-64-5]	cycloheptane				
	$\Delta_{\text{trs}} H$		4.98	134.8		
	$\Delta_{\text{trs}} H$		0.29	198.2		
	$\Delta_{\text{trs}} H$		0.45	212.4		
	$\Delta_{\text{fus}} H$		1.88	265.1		[1996DOM/HEA]
	$\Delta_{\text{sub}} H$		53.5	134		[1963BON]
	$\Delta_v H$	(282–333)	38.6	297	A	[1987STE/MAL]
	$\Delta_v H$	(476–604)	31.7	491	A	[1987STE/MAL]
	$\Delta_v H$	(333–398)	36.4	348	A,EB	[1987STE/MAL, 1976MEY/HOT]
	$\Delta_v H$	(283–323)	38.5	298		[1975ANA/GRO]
	$\Delta_v H$	(341–433)	36.1	356	A,EB	[1987STE/MAL, 1956FIN/SCO]
	$\Delta_v H$		38.5 ± 0.2	298		[1956FIN/SCO]
C ₇ H ₁₄	[108-87-2]	methylcyclohexane				
	$\Delta_{\text{fus}} H$		6.69	146.6		[1996DOM/HEA]
	$\Delta_v H$	(325–374)	33.8	340		[2010SAP/UUS]
	$\Delta_v H$	(295–333)	36.2	310		[1991WU/PIV]
	$\Delta_v H$		35.1 ± 0.4	298	GC	[1987AZA]
	$\Delta_v H$	(373–511)	32.3	388	A	[1987STE/MAL]
	$\Delta_v H$	(501–573)	31.2	516	A	[1987STE/MAL]
	$\Delta_v H$		32.2	353		[1984EUB/CED]
	$\Delta_v H$		29.9	393		[1984EUB/CED]
	$\Delta_v H$		26.9	433		[1984EUB/CED]
	$\Delta_v H$		23.4	473		[1984EUB/CED]
	$\Delta_v H$		35.4 ± 0.1	298	C	[1979MAJ/SVO]
	$\Delta_v H$		34.6 ± 0.1	313	C	[1979MAJ/SVO]
	$\Delta_v H$		33.5 ± 0.1	333	C	[1979MAJ/SVO]
	$\Delta_v H$		32.5 ± 0.1	353	C	[1979MAJ/SVO]
	$\Delta_v H$		35.4	298	GCC	[1978FUC/PEA]
	$\Delta_v H$		35.3	298		[1975KUS/SAI]
	$\Delta_v H$	(308–368)	34.6	323	A	[1987STE/MAL, 1970VAL/KIL, 1984BOU/FRI]
	$\Delta_v H$		31.8	374		[1946SPI/PIT]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound					
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference	
		$\Delta_v H$	(299–375)	34.9	314	MM	[1945WIL/TAY]
		$\Delta_v H$	(273–348)	36.1	288		[1940STU/SAY]
C ₇ H ₁₄	[1638-26-2]	1,1-dimethylcyclopentane					
		$\Delta_{\text{fus}} H$		6.49	146.8		
		$\Delta_{\text{fus}} H$		1.09	203.7		[1996DOM/HEA]
		$\Delta_v H$	(284–363)	34.0	299	A	[1987STE/MAL]
		$\Delta_v H$		33.8	298		[1971WIL/ZWO]
		$\Delta_v H$	(289–362)	33.8	304		[1949FOR/NOR]
C ₇ H ₁₄	[1192-18-3]	<i>cis</i> 1,2-dimethylcyclopentane					
		$\Delta_{\text{fus}} H$		6.65	141.5		
		$\Delta_{\text{fus}} H$		1.66	219.4		[1996DOM/HEA]
		$\Delta_v H$	(293–375)	35.5	308	A	[1987STE/MAL]
		$\Delta_v H$		35.8	298		[1971WIL/ZWO]
		$\Delta_v H$	(298–373)	35.2	313		[1949FOR/NOR]
C ₇ H ₁₄	[822-50-4]	<i>(dl)</i> <i>trans</i> 1,2-dimethylcyclopentane					
		$\Delta_v H$	(295–367)	34.2	310	A	[1987STE/MAL]
		$\Delta_v H$		34.6	298		[1971WIL/ZWO]
		$\Delta_v H$	(299–366)	34.0	314		[1949FOR/NOR]
C ₇ H ₁₄	[2532-58-3]	<i>cis</i> 1,3-dimethylcyclopentane					
		$\Delta_v H$	(295–366)	34.2	310	A	[1987STE/MAL]
		$\Delta_v H$		34.3	298		[1971WIL/ZWO]
		$\Delta_v H$		32.8 ± 0.1	323	C	[1959MCC/PEN]
		$\Delta_v H$		31.7 ± 0.1	342	C	[1959MCC/PEN]
		$\Delta_v H$	(299–366)	30.4 ± 0.1	364	C	[1959MCC/PEN]
C ₇ H ₁₄	[1759-58-6]	<i>(dl)</i> <i>trans</i> 1,3-dimethylcyclopentane					
		$\Delta_{\text{fus}} H$		7.41	139.5		[1996DOM/HEA]
		$\Delta_v H$	(295–367)	34.0	310	A	[1987STE/MAL]
		$\Delta_v H$		34.5	298		[1971WIL/ZWO]
C ₇ H ₁₄	[1640-89-7]	ethylcyclopentane					
		$\Delta_{\text{fus}} H$		6.86	134.7		[1996DOM/HEA]
		$\Delta_v H$	(308–387)	35.5	323	A	[1987STE/MAL]
		$\Delta_v H$	(386–507)	32.9	401	A	[1987STE/MAL]
		$\Delta_v H$	(499–569)	31.9	514	A	[1987STE/MAL]
		$\Delta_v H$		35.6 ± 0.1	313	C	[1981SVO/CHA]
		$\Delta_v H$		34.8 ± 0.1	328	C	[1981SVO/CHA]
		$\Delta_v H$		33.9 ± 0.1	343	C	[1981SVO/CHA]
		$\Delta_v H$		33.0 ± 0.1	358	C	[1981SVO/CHA]
		$\Delta_v H$		32.5 ± 0.1	368	C	[1981SVO/CHA]
		$\Delta_v H$		36.5	298		[1971WIL/ZWO]
		$\Delta_v H$	(302–377)	35.7	317		[1949FOR/NOR]
C ₇ H ₁₄	[592-76-7]	1-heptene					
		$\Delta_{\text{fus}} H$		12.64	154.3		[1996DOM/HEA]
		$\Delta_v H$	(311–368)	34.6	326	A	[1987STE/MAL]
		$\Delta_v H$	(327–367)	33.9	342		[1970EIS/ORA, 1984BOU/FRI]
		$\Delta_v H$		35.7	298		[1971WIL/ZWO]
		$\Delta_v H$	(295–318)	35.3	310	MM	[1950FOR/CAM]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(255–312)	35.9	300		[194ILIS]
	$\Delta_v H$	(273–362)	34.5	288		[1936BEN/CUT]
C₇H₁₄	[6443-92-1]	<i>cis</i> 2-heptene				
	$\Delta_v H$	(276–304)	39.0 ± 0.3	290	GS	[2000VER/WAN]
	$\Delta_v H$	(276–304)	38.6	298	GS	[2000VER/WAN]
	$\Delta_v H$	(315–372)	35.3	330	A	[1987STE/MAL]
	$\Delta_v H$	(332–371)	34.6	347		[1970EIS/ORA, 1984BOU/FRI]
	$\Delta_v H$		36.0	298		[1971WIL/ZWO]
C₇H₁₄	[14686-13-6]	<i>trans</i> 2-heptene				
	$\Delta_v H$	(314–373)	35.3	329	A	[1987STE/MAL]
	$\Delta_v H$		36.0	298		[1971WIL/ZWO]
	$\Delta_v H$	(331–370)	34.6	346		[1970EIS/ORA]
C₇H₁₄	[7642-10-6]	<i>cis</i> 3-heptene				
	$\Delta_v H$	(312–369)	35.0	327	A	[1987STE/MAL]
	$\Delta_v H$		35.6	298		[1971WIL/ZWO]
C₇H₁₄	[14686-14-7]	<i>trans</i> 3-heptene				
	$\Delta_v H$	(312–368)	34.6	327	A	[1987STE/MAL]
	$\Delta_v H$		35.6	298		[1971WIL/ZWO]
C₇H₁₄	[6094-02-6]	2-methyl-1-hexene				
	$\Delta_v H$	(318–390)	33.9	333	A	[1987STE/MAL, 1973KKY/REP]
	$\Delta_v H$		35.1	298		[1971WIL/ZWO]
C₇H₁₄	[3404-61-3]	(<i>dl</i>) 3-methyl-1-hexene				
	$\Delta_v H$	(311–381)	33.4	326	A	[1987STE/MAL, 1973KKY/REP]
	$\Delta_v H$		34.3	298		[1971WIL/ZWO]
C₇H₁₄	[3769-23-1]	(<i>dl</i>) 4-methyl-1-hexene				
	$\Delta_v H$	(313–384)	33.6	328	A	[1987STE/MAL, 1973KKY/REP]
	$\Delta_v H$		34.7	298		[1971WIL/ZWO]
C₇H₁₄	[3524-73-0]	5-methyl-1-hexene				
	$\Delta_v H$	(313–393)	33.5	328	A	[1987STE/MAL, 1973KKY/REP]
	$\Delta_v H$		34.3	298		[1971WIL/ZWO]
C₇H₁₄	[2738-19-4]	2-methyl-2-hexene				
	$\Delta_v H$	(322–394)	34.0	337	A	[1987STE/MAL, 1973KKY/REP]
	$\Delta_v H$		35.6	298		[1971WIL/ZWO]
C₇H₁₄	[10574-36-4]	<i>cis</i> 3-methyl-2-hexene				
	$\Delta_v H$	(322–396)	34.2	337	A	[1987STE/MAL, 1973KKY/REP]
	$\Delta_v H$		35.6	298		[1971WIL/ZWO]
C₇H₁₄	[20710-38-7]	<i>trans</i> 3-methyl-2-hexene				
	$\Delta_v H$	(321–394)	34.1	336	A	[1987STE/MAL, 1973KKY/REP]
	$\Delta_v H$		35.6	298		[1971WIL/ZWO]
C₇H₁₄	[3683-19-0]	<i>cis</i> 4-methyl-2-hexene				
	$\Delta_v H$	(313–384)	33.5	328	A	[1987STE/MAL, 1973KKY/REP]
	$\Delta_v H$		34.7	298		[1971WIL/ZWO]
C₇H₁₄	[3683-22-5]	<i>trans</i> 4-methyl-2-hexene				
	$\Delta_v H$	(314–385)	33.6	329	A	[1987STE/MAL, 1973KKY/REP]
	$\Delta_v H$		34.7	298		[1971WIL/ZWO]
C₇H₁₄	[13151-17-2]	<i>cis</i> 5-methyl-2-hexene				
	$\Delta_v H$	(354–372)	32.6	363	A	[1987STE/MAL, 1973KKY/REP]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
		$\Delta_v H$	34.7	298		[1971WIL/ZWO]
C ₇ H ₁₄	[3683-22-5]	<i>trans</i> 5-methyl-2-hexene				
	$\Delta_v H$	(315–386)	33.6	330	A	[1987STE/MAL, 1973KKY/REP]
	$\Delta_v H$		34.7	298		[1971WIL/ZWO]
C ₇ H ₁₄	[15840-60-5]	<i>cis</i> 2-methyl-3-hexene				
	$\Delta_v H$	(262–383)	36.1	277	A	[1987STE/MAL, 1973KKY/REP]
	$\Delta_v H$		34.3	298		[1971WIL/ZWO]
C ₇ H ₁₄	[692-24-0]	<i>trans</i> 2-methyl-3-hexene				
	$\Delta_v H$	(313–383)	33.5	328	A	[1987STE/MAL, 1973KKY/REP]
	$\Delta_v H$		34.3	298		[1971WIL/ZWO]
C ₇ H ₁₄	[4914-89-0]	<i>cis</i> 3-methyl-3-hexene				
	$\Delta_v H$	(307–375)	35.4	322	A	[1987STE/MAL]
	$\Delta_v H$		36.4	298		[1971WIL/ZWO]
	$\Delta_v H$	(302–368)	35.7	317	MM	[1960CAM/ROS]
C ₇ H ₁₄	[3899-36-3]	<i>trans</i> 3-methyl-3-hexene				
	$\Delta_v H$	(310–368)	34.8	325	A	[1987STE/MAL]
	$\Delta_v H$		35.8	298		[1971WIL/ZWO]
	$\Delta_v H$	(300–367)	35.3	315	MM	[1960CAM/ROS]
C ₇ H ₁₄	[3404-72-6]	<i>(dl)</i> 2,3-dimethylpent-1-ene				
	$\Delta_v H$	(311–382)	33.4	326	A	[1987STE/MAL, 1973KKY/REP]
C ₇ H ₁₄	[na]	2,3-dimethylpent-2-ene				
	$\Delta_v H$		34.3	298		[1971WIL/ZWO]
C ₇ H ₁₄	[2213-32-3]	2,4-dimethylpent-1-ene				
	$\Delta_v H$	(311–361)	32.3	326	A	[1987STE/MAL]
	$\Delta_v H$		33.1	298		[1971WIL/ZWO]
	$\Delta_v H$	(289–355)	33.2	304	MM	[1960CAM/ROS]
C ₇ H ₁₄	[3404-73-7]	3,3-dimethylpent-1-ene				
	$\Delta_v H$	(306–374)	33.0	321	A	[1987STE/MAL, 1973KKY/REP]
	$\Delta_v H$		33.5	298		[1971WIL/ZWO]
C ₇ H ₁₄	[7385-78-6]	<i>(dl)</i> 3,4-dimethylpent-1-ene				
	$\Delta_v H$	(309–378)	33.2	324	A	[1987STE/MAL, 1973KKY/REP]
	$\Delta_v H$		33.9	298		[1971WIL/ZWO]
C ₇ H ₁₄	[762-62-9]	4,4-dimethylpent-1-ene				
	$\Delta_v H$	(299–347)	31.0	314	A	[1987STE/MAL]
	$\Delta_v H$		31.2	298		[1971WIL/ZWO]
	$\Delta_v H$	(290–346)	31.0	315	MM	[1960CAM/ROS]
C ₇ H ₁₄	[10574-37-5]	2,3-dimethylpent-2-ene				
	$\Delta_v H$	(322–396)	34.2	337	A	[1987STE/MAL, 1973KKY/REP]
	$\Delta_v H$		35.6	298		[1971WIL/ZWO]
C ₇ H ₁₄	[625-65-0]	2,4-dimethylpent-2-ene				
	$\Delta_v H$	(276–297)	35.2 ± 1.5	286	GS	[2000VER/WAN]
	$\Delta_v H$	(276–297)	34.5 ± 1.5	298	GS	[2000VER/WAN]
	$\Delta_v H$	(286–363)	34.5	301	A	[1987STE/MAL]
	$\Delta_v H$		34.3	298		[1971WIL/ZWO]
	$\Delta_v H$	(292–357)	34.2	307	MM	[1960CAM/ROS]
C ₇ H ₁₄	[4914-91-4]	<i>cis</i> 3,4-dimethylpent-2-ene				
	$\Delta_v H$	(316–387)	33.7	331	A	[1987STE/MAL, 1973KKY/REP]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$		34.7	298		[1971WIL/ZWO]
C ₇ H ₁₄	[4914-92-5]	<i>trans</i> 3,4-dimethylpent-2-ene				
	$\Delta_v H$	(317–390)	33.9	332	A	[1987STE/MAL, 1973KKY/REP]
	$\Delta_v H$		35.1	298		[1971WIL/ZWO]
C ₇ H ₁₄	[762-63-0]	<i>cis</i> 4,4-dimethylpent-2-ene				
	$\Delta_v H$	(303–355)	32.2	318	A	[1987STE/MAL]
	$\Delta_v H$		32.6	298		[1971WIL/ZWO]
C ₇ H ₁₄	[690-08-4]	<i>trans</i> 4,4-dimethylpent-2-ene				
	$\Delta_v H$	(295–352)	32.8	310	A	[1987STE/MAL]
	$\Delta_v H$		32.8	298		[1971WIL/ZWO]
C ₇ H ₁₄	[690-08-4]	<i>trans</i> 4,4-dimethylpent-2-ene				
	$\Delta_v H$	(289–350)	33.0	304	MM	[1960CAM/ROS]
	$\Delta_v H$		33.0	304		[1960CAM/ROS]
C ₇ H ₁₄	[7357-93-9]	2-ethyl-3-methyl-1-butene				
	$\Delta_v H$	(303–381)	33.8	318	A	[1987STE/MAL]
	$\Delta_v H$		34.3	298		[1971WIL/ZWO]
C ₇ H ₁₄	[7357-93-9]	2-ethyl-3-methyl-1-butene				
	$\Delta_v H$	(290–360)	34.4	305	MM	[1960CAM/ROS]
	$\Delta_v H$		34.4	305		[1960CAM/ROS]
C ₇ H ₁₄	[3404-71-5]	2-ethyl-1-pentene				
	$\Delta_v H$	(267–392)	36.6	282	A	[1987STE/MAL, 1973KKY/REP]
	$\Delta_v H$		35.1	298		[1971WIL/ZWO]
C ₇ H ₁₄	[4038-04-4]	3-ethyl-1-pentene				
	$\Delta_v H$	(311–382)	33.4	326	A	[1987STE/MAL, 1973KKY/REP]
	$\Delta_v H$		34.3	298		[1971WIL/ZWO]
C ₇ H ₁₄	[816-79-5]	3-ethyl-2-pentene				
	$\Delta_v H$	(321–395)	34.1	336	A	[1987STE/MAL, 1973KKY/REP]
	$\Delta_v H$		35.6	298		[1971WIL/ZWO]
C ₇ H ₁₄	[594-56-9]	2,3,3-trimethyl-1-butene				
	$\Delta_v H$	(288–353)	32.4	303	A	[1987STE/MAL]
	$\Delta_v H$		34.3	298		[1971WIL/ZWO]
C ₇ H ₁₄	[594-56-9]	2,3,3-trimethyl-1-butene				
	$\Delta_v H$	(288–351)	32.1	303	MM	[1960CAM/ROS]
	$\Delta_v H$		32.1	303		[1960CAM/ROS]
C ₇ H ₁₄ Br ₂	[59104-79-9]	1,1-dibromoheptane				
	$\Delta_v H$	(395–548)	54.4	410	A,E	[1987STE/MAL, 1956MAN, 1970DYK/VAN]
C ₇ H ₁₄ Br ₂	[42474-21-5]	(<i>dl</i>) 1,2-dibromoheptane				
	$\Delta_v H$	(295–553)	52.9	310	A	[1987STE/MAL, 1973KKY/REP, 1999DYK/SVO]
	$\Delta_v H$	(295–355)	54.4	300		[1941LIS]
C ₇ H ₁₄ Cl ₂	[821-25-0]	1,1-dichloroheptane				
	$\Delta_v H$	(375–460)	53.5	298		[1987VAR/LOS2, 1991BAS/SVO]
	$\Delta_v H$	(364–510)	48.4	379	A,E	[1987STE/MAL, 1956MAN, 1970DYK/VAN]
C ₇ H ₁₄ Cl ₂	[10575-87-8]	1,2-dichloroheptane				
	$\Delta_v H$	(353–466)	49.0	368		[1999DYK/SVO]
	$\Delta_v H$	(350–470)	53.2	298		[1982VAR/PUC, 1991BAS/SVO]
C ₇ H ₁₄ Cl ₂	[821-76-1]	1,7-dichloroheptane				
	$\Delta_v H$	(406–491)	52.3	421		[1999DYK/SVO]
	$\Delta_v H$	(410–490)	61.2	298		[1988VAR/LOS, 1991BAS/SVO]
C ₇ H ₁₄ F ₂	[407-96-5]	1,1-difluoroheptane				
	$\Delta_v H$	(311–424)	41.1	326	A,E	[1987STE/MAL, 1956MAN, 1970DYK/VAN]
C ₇ H ₁₄ NO ₅ P	[6923-22-4]	dimethyl (E)-1-methyl-2-methylcarbamoylvinyl phosphate				

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
		$\Delta_{\text{fus}}H$	22.36	326.9	DSC	[1990DON/DRE]
C ₇ H ₁₄ N ₂	[5351-04-2]	3-(diethylamino)propionitrile				
		Δ_vH	(338–470) 53.7	353	A	[1987STE/MAL]
C ₇ H ₁₄ N ₂	[na]	2-(diethylamino)propionitrile				
		Δ_vH	(278–315) 50.8 ± 0.3		GS	[1997WEL/VER]
C ₇ H ₁₄ N ₂	[2721-31-5]	3,3,5,5-tetramethyl-1-pyrazoline				
		$\Delta_{\text{sub}}H$	61.6 ± 0.2	298		[1976ENG/MEL]
C ₇ H ₁₄ N ₂ O ₂	[37933-88-3]	N-acetyl L-valinamide				
		$\Delta_{\text{fus}}H$	39.1	509		[1997SAN/ROC]
		$\Delta_{\text{sub}}H$	129.8 ± 1.9	376	C	[1999DEL/BAR]
		$\Delta_{\text{sub}}H$	133.1 ± 2.2	298		[1999DEL/BAR]
		$\Delta_{\text{sub}}H$	(391–425) 126 ± 2.0	418		[1990PUL/MAT]
C ₇ H ₁₄ N ₂ O ₂	[1740-56-3]	pimelamide				
		$\Delta_{\text{fus}}H$	44.56	446.8	DSC	[2006BAD/DEL]
C ₇ H ₁₄ N ₂ O ₂ S	[116-06-3]	2-methyl-2(methylthio)propanal, O-[(methylamino)-carbonyl]oxime				
		$\Delta_{\text{fus}}H$	22.71	374	DSC	[1990DON/DRE]
		$\Delta_{\text{sub}}H$	(298–323) 80.0	310	ME	[1987STE/MAL, 1976DEP]
C ₇ H ₁₄ O	[5063-65-0]	1,2-epoxyheptane				
		Δ_vH	(305–414) 45.5	320	A	[1987STE/MAL, 1970VOJ/CIH]
C ₇ H ₁₄ O	[502-41-0]	cycloheptanol				
		$\Delta_{\text{us}}H$	0.45	227.9		
		$\Delta_{\text{us}}H$	0.78	250.4		
		$\Delta_{\text{fus}}H$	1.51	278.3		[2003RUT/SAL]
		$\Delta_{\text{us}}H$	2.93	172.2		
		$\Delta_{\text{us}}H$	0.55	227.3		
		$\Delta_{\text{us}}H$	0.88	258.4		
		$\Delta_{\text{fus}}H$	1.6	280.3		[1996DOM/HEA]
		Δ_vH	(284–323) 64.7	299	A	[1987STE/MAL]
		Δ_vH	(284–321) 67.4	299		[1975CAB/CON2]
C ₇ H ₁₄ O	[590-67-0]	1-methylcyclohexanol				
		$\Delta_{\text{fus}}H$	14.32	299.4		[1998KAB/BLO]
		$\Delta_{\text{fus}}H$	10.87	299.2		[1985WIB/WAS]
		$\Delta_{\text{sub}}H$	75.9 ± 0.4	291	C	[1998KAB/BLO]
		Δ_vH	(340–430) 49.1	355	A	[1987STE/MAL]
C ₇ H ₁₄ O	[583-59-5]	2-methylcyclohexanol				
		Δ_vH	(361–439) 51.7	376	EB	[2007PAL/ORO]
		Δ_vH	(323–373) 63.3	298	CGC	[1995CHI/HOS]
C ₇ H ₁₄ O	[7443-70-1]	<i>cis</i> 2-methylcyclohexanol				
		Δ_vH	61.8			[1975VIL/PER]
C ₇ H ₁₄ O	[na]	3-methylcyclohexanol				
		Δ_vH	(323–373) 65.5	298	CGC	[1995CHI/HOS]
C ₇ H ₁₄ O	[24965-90-0]	(<i>dl</i>) <i>cis</i> 3-methylcyclohexanol				
		Δ_vH	(340–450) 54.3	355	A	[1987STE/MAL]
C ₇ H ₁₄ O	[23068-71-5]	(<i>dl</i>) <i>trans</i> 3-methylcyclohexanol				

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(350–450)	50	365	A	[1987STE/MAL]
C ₇ H ₁₄ O	[na]	4-methylcyclohexanol				
	$\Delta_v H$	(323–373)	65.9	298	CGC	[1995CHI/HOS]
C ₇ H ₁₄ O	[7731-28-4]	<i>cis</i> 4-methylcyclohexanol				
	$\Delta_v H$	(340–450)	49.9	355	A	[1987STE/MAL]
C ₇ H ₁₄ O	[7731-29-5]	<i>trans</i> 4-methylcyclohexanol				
	$\Delta_v H$	(340–350)	52.1	355	A	[1987STE/MAL]
C ₇ H ₁₄ O	[1462-96-0]	1-ethyl-1-cyclopentanol				
	$\Delta_v H$	(347–426)	58.4	362	A	[1987STE/MAL, 1973KKY/REP]
C ₇ H ₁₄ O	[110-43-0]	2-heptanone				
	$\Delta_{\text{fus}} H$		19.71	237.7		[1996FIE/JOH]
	$\Delta_v H$	(343–383)	46.1	298	CGC	[1995CHI/HOS]
	$\Delta_v H$	(343–383)	48.5	298	CGC	[1995CHI/HOS]
	$\Delta_v H$	(303–424)	47.5	318	A	[1987STE/MAL]
	$\Delta_v H$	(449–480)	39.1	464	A	[1987STE/MAL]
	$\Delta_v H$		47.4 ± 0.3	298	GCC	[1979SAL/PEA]
	$\Delta_v H$		47.2 ± 0.1	298	C	[1979SUN/SVE2]
	$\Delta_v H$	(327–457)	44.7	342		[1987STE/MAL, 1975AMB/ELL]
	$\Delta_v H$		48.0	298		[1975AMB/ELL]
C ₇ H ₁₄ O	[123-19-3]	3-heptanone				
	$\Delta_{\text{fus}} H$		17.53	236		[1996FIE/JOH]
C ₇ H ₁₄ O	[123-19-3]	4-heptanone				
	$\Delta_{\text{fus}} H$		16.16	240.2		[1996FIE/JOH]
	$\Delta_v H$	(343–383)	47.8	298	CGC	[1995CHI/HOS]
	$\Delta_v H$		46.2 ± 0.4	298	GCC	[1979SAL/PEA]
	$\Delta_v H$	(304–490)	45.5	319	A	[1987STE/MAL, 1975AMB/ELL]
	$\Delta_v H$		46.7	298		[1975AMB/ELL]
	$\Delta_v H$	(296–417)	57.5	311	A	[1987STE/MAL, 1947STU]
	$\Delta_v H$	(283–323)	40.7	303		[1937RIN/SAY]
C ₇ H ₁₄ O	[7379-12-6]	2-methyl-3-hexanone				
	$\Delta_v H$	(296–406)	41.3	311	A	[1987STE/MAL]
C ₇ H ₁₄ O	[564-04-5]	2,2-dimethyl-3-pentanone				
	$\Delta_v H$		42.3 ± 0.1	298	C	[1970SEL2]
	$\Delta_v H$		42.3 ± 0.1	298	C	[1966WAD]
C ₇ H ₁₄ O	[565-80-0]	2,4-dimethyl-3-pentanone				
	$\Delta_{\text{fus}} H$		11.2	204.8		[1996DOM/HEA]
	$\Delta_v H$	(321–399)	39.4	336	A	[1987STE/MAL, 1973KKY/REP]
	$\Delta_v H$		41.6 ± 0.1	298	C	[1970SEL2]
	$\Delta_v H$		41.5 ± 0.1	298	C	[1966WAD]
C ₇ H ₁₄ O	[111-71-7]	heptanal				
	$\Delta_{\text{fus}} H$		22.89	229.3		[1980DYA/VAS]
	$\Delta_v H$	(313–353)	48.7	298	CGC	[1995CHI/HOS]
	$\Delta_v H$		47.3 ± 0.1	298		[1981DYA/KOR]
	$\Delta_v H$		48.0 ± 1.3	298	EB	[1960NIC, 2003VER/KRA2]
	$\Delta_v H$	(285–428)	62.0	300		[1947STU]
C ₇ H ₁₄ O	[19269-28-4]	3-methylhexanal				

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(314–417)	42.8	329	EB	[1987MIL/FEN]
C ₇ H ₁₄ O	[19353-21-0]	3,4-dimethylpentanal				
	$\Delta_v H$	(319–417)	42.4	334	EB	[1987MIL/FEN]
C ₇ H ₁₄ O ₂	[931-94-2]	1,1-dimethoxycyclopentane				
	$\Delta_v H$	(278–318)	44.5 ± 0.3	298	GS	[1998VER/PEN, 2002VER]
	$\Delta_v H$	(307–343)	46.1	325	EB	[1994WIB/MOR]
C ₇ H ₁₄ O ₂	[106-70-7]	methyl hexanoate				
	$\Delta_v H$	(281–331)	48.4 ± 0.2	298	GS	[2008VER/EME]
	$\Delta_v H$		45.2	350		[2002VAN/VAN]
	$\Delta_v H$		46.4 ± 0.1	325		[2002VAN/VAN]
	$\Delta_v H$		47.7 ± 0.1	298		[2002VAN/VAN]
	$\Delta_v H$	(313–363)	47.9	298	CGC	[1995CHI/HOS]
	$\Delta_v H$	(313–353)	48.2	298	CGC	[1995CHI/HOS]
	$\Delta_v H$		47.8 ± 0.5	298	GC	[1987AZA]
	$\Delta_v H$		48.7 ± 0.3	298	GCC	[1980FUC/PEA]
	$\Delta_v H$		48.0 ± 0.1	298	C	[1977MAN/SEL]
	$\Delta_v H$	(315–383)	45.3	330	A	[1987STE/MAL, 1963ROS/SCH]
C ₇ H ₁₄ O ₂	[590-01-2]	butyl propionate				
	$\Delta_v H$	(305–417)	49.1	320	A	[1987STE/MAL]
	$\Delta_v H$	(305–365)	47.4	320		[1959USA/DEM2, 1984BOU/FRI]
C ₇ H ₁₄ O ₂	[3938-95-2]	ethyl pivalate				
	$\Delta_v H$	(308–429)	39.8 ± 0.1	320	EB	[2002STE/CHI4]
	$\Delta_v H$	(308–429)	36.9 ± 0.2	360	EB	[2002STE/CHI4]
	$\Delta_v H$	(308–429)	33.8 ± 0.6	400	EB	[2002STE/CHI4]
			41.3 ± 0.1	298	C	[1966WAD]
C ₇ H ₁₄ O ₂	[108-64-5]	ethyl isovalerate				
	$\Delta_v H$	(301–418)	42.8	316	A	[1987STE/MAL]
	$\Delta_v H$	(267–407)	44.5	282		[1947STU]
C ₇ H ₁₄ O ₂	[540-42-1]	isobutyl propionate				
	$\Delta_v H$	(271–410)	44.9	286	A	[1987STE/MAL, 1947STU]
C ₇ H ₁₄ O ₂	[123-92-2]	isopentyl acetate				
	$\Delta_v H$	(230–435)	46.4	300		[1999DIA/GUE]
	$\Delta_v H$	(278–305)	46.8 ± 0.2	292	GS	[1999VER/HEI]
	$\Delta_v H$	(278–305)	46.4 ± 0.2	298	GS	[1999VER/HEI]
	$\Delta_v H$	(308–424)	44.3	323	A	[1987STE/MAL]
	$\Delta_v H$	(313–368)	45.1	328		[1959TER/BRI, 1984BOU/FRI]
C ₇ H ₁₄ O ₂	[617-50-5]	isopropyl isobutyrate				
	$\Delta_v H$	(257–394)	43.3	272	A	[1987STE/MAL, 1947STU]
C ₇ H ₁₄ O ₂	[107-70-0]	4-methoxy-4-methyl-2-pentanone				
	$\Delta_v H$	(343–423)	45.0	358	A	[1987STE/MAL]
C ₇ H ₁₄ O ₂	[10250-48-3]	methyl <i>tert</i> -butylacetate				
	$\Delta_v H$	(274–313)	43.9 ± 0.2	298	GS	[2008VER/EME]
	$\Delta_v H$	(274–313)	44.4 ± 0.2	298	GS	[1996VER/BEC]
C ₇ H ₁₄ O ₂	[926-41-0]	neopentyl acetate				
	$\Delta_v H$	(301–400)	49.1	316	A	[1987STE/MAL]
C ₇ H ₁₄ O ₂	[625-16-1]	<i>tert</i> -pentyl acetate				
	$\Delta_v H$	(274–308)	40.3	298	GS	[2008VER/EME]
	$\Delta_v H$		42.8	209	CGC	[2000MAT/MIR]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(274–308)	42.8 ± 0.3	298	GS	[1996VER/BEC]
C ₇ H ₁₄ O ₂	[628-63-7]	pentyl acetate				
	$\Delta_v H$	(321–462)	48.6 ± 0.4	298	EB	[1996STE/CHI]
	$\Delta_v H$	(329–423)	43.2	344	A	[1987STE/MAL]
C ₇ H ₁₄ O ₂	[105-66-8]	propyl butyrate				
	$\Delta_v H$	(390–430)	39.6	405		[1995ORT/GAL]
	$\Delta_v H$	(355–416)	42.0	370		[1993FAR/WIC]
	$\Delta_v H$	(271–416)	44.3	286	A	[1987STE/MAL, 1947STU]
C ₇ H ₁₄ O ₂	[644-49-5]	propyl isobutyrate				
	$\Delta_v H$	(267–407)	50.5	282	A	[1987STE/MAL, 1947STU]
C ₇ H ₁₄ O ₂	[na]	ethyl 2-methylbutanoate				
	$\Delta_v H$	(288–308)	64.7	298	GS	[1992VER/BEC]
C ₇ H ₁₄ O ₂	[2426-08-6]	butyl glycidyl ether				
	$\Delta_v H$		53.3 ± 0.4			[1987VAN/KAC]
C ₇ H ₁₄ O ₂	[7665-72-7]	<i>tert</i> -butyl glycidyl ether				
	$\Delta_v H$		50.2 ± 0.4			[1987VAN/KAC]
C ₇ H ₁₄ O ₂	[na]	2,2,4-trimethyl-1,3-dioxane				
	$\Delta_v H$		41.9 ± 1.2	298		[1967PIH/HEI]
C ₇ H ₁₄ O ₂	[4352-98-1]	2-methyl-2-propyl-1,3-dioxolane				
	$\Delta_v H$	(278–313)	46.3 ± 0.3	298	GS	[1998VER/PEN, 2002VER]
C ₇ H ₁₄ O ₂	[4405-16-7]	2-methyl-2-isopropyl-1,3-dioxolane				
	$\Delta_v H$	(274–303)	43.9 ± 0.2	298	GS	[2002VER]
	$\Delta_v H$	(273–303)	44.6 ± 0.2	298	GS	[1998VER/PEN]
C ₇ H ₁₄ O ₂	[3814-55-9]	[(1-methylpropoxy)methyl]oxirane				
	$\Delta_v H$		45.2 ± 1.8			[1987VAN/KAC]
C ₇ H ₁₄ O ₂	[111-14-8]	heptanoic acid				
	$\Delta_{\text{us}}H$		2.04	224.8		
	$\Delta_{\text{fus}}H$		15.44	265.8		[1996DOM/HEA, 1991LAB/WES]
	$\Delta_v H$	(413–453)	69.0	428		[2004CLI/RAM]
	$\Delta_v H$	(283–328)	72.5 ± 0.8	306	GS	[2000VER2]
	$\Delta_v H$	(283–328)	72.9 ± 0.8	298	GS	[2000VER2]
	$\Delta_v H$	(353–393)	75.7	298	CGC	[1995CHI/HOS]
	$\Delta_v H$		76.0	266		[1982DEK/SCH]
	$\Delta_v H$	(271–291)	72.0 ± 1.5	298	TE	[1979DEK/OON]
	$\Delta_v H$	(351–495)	68.3	366	A	[1987STE/MAL, 1947STU]
C ₇ H ₁₄ O ₃	[33415-52-0]	<i>tert</i> -butylperoxymethyloxirane				
	$\Delta_v H$		53.9 ± 0.4			[1987VAN/KAC]
C ₇ H ₁₄ O ₃	[20706-25-6]	2-propoxyethylacetate				
	$\Delta_v H$		55.6 ± 0.1	298	C	[1970KUS/WAD]
C ₇ H ₁₄ O ₃	[138-22-7]	(<i>dl</i>) butyl lactate				
	$\Delta_v H$	(391–460)	49.9	406		[2005PEN/MUR]
	$\Delta_v H$	(339–456)	58.7	354	A	[1987STE/MAL]
C ₇ H ₁₄ O ₃	[763-69-9]	3-ethoxypropionic acid, ethyl ester				
	$\Delta_v H$	(312–446)	45.5	327	A	[1987STE/MAL]
C ₇ H ₁₄ O ₃	[na]	1-heptene ozonide				

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(353–373)	44.4	363	A	[1987STE/MAL, 1977BOL/MAK]
C ₇ H ₁₄ O ₃	[na]	4-(2-hydroxyethyl)-4-methyl-1,3-dioxane				
	$\Delta_v H$	(329–455)	51.7	344	A	[1987STE/MAL]
C ₇ H ₁₄ O ₃	[na]	3-hydroxypropionic acid, butyl ester				
	$\Delta_v H$	(361–382)	60.3	371	A	[1987STE/MAL, 1973KKY/REP]
C ₇ H ₁₄ O ₃	[5349-56-4]	3-methoxypropionic acid, propyl ester				
	$\Delta_v H$	(323–433)	47.0	338	A	[1987STE/MAL]
C ₇ H ₁₄ O ₃	[14144-39-9]	3-propoxypropionic acid, methyl ester				
	$\Delta_v H$	(323–453)	46.6	338	A	[1987STE/MAL]
C ₇ H ₁₄ O ₃	[557-25-5]	(<i>dl</i>) butyric acid, 2,3-dihydroxypropyl ester				
	$\Delta_v H$	(392–449)	80.4	407	A	[1987STE/MAL]
C ₇ H ₁₄ O ₃	[14620-87-2]	2-butoxypropionic acid				
	$\Delta_v H$	(373–473)	52.8	388	A	[1987STE/MAL, 1973KKY/REP]
C ₇ H ₁₄ O ₃	[623-96-1]	dipropyl carbonate				
	$\Delta_v H$	(274–318)	53.2 ± 0.3	298	GS	[2008KOZ/EME]
C ₇ H ₁₄ O ₆	[na]	1-methoxy- α -(<i>d</i>)-glucopyranoside				
	$\Delta_{\text{fus}} H$		35.11	436	DSC	[2000AGU/GUA]
	$\Delta_{\text{fus}} H$		37.6	424.2		[1996SCH]
C ₇ H ₁₄ O ₆	[na]	3-methoxy- α -(<i>d</i>)-glucopyranoside				
	$\Delta_{\text{fus}} H$		41.3	425.6		[1996SCH]
C ₇ H ₁₄ O ₆	[na]	methyl α -(<i>d</i>)-mannopyranoside				
	$\Delta_{\text{fus}} H$		44.7	455.2		[1996SCH]
C ₇ H ₁₄ O ₆	[na]	methyl- α -(<i>d</i>)-glucopyranoside				
	$\Delta_{\text{fus}} H$		35.11	436		[2000AGU/GUA]
C ₇ H ₁₄ O ₆	[na]	methyl- β -(<i>d</i>)-glucopyranoside				
	$\Delta_{\text{fus}} H$		27.03	384.9		[2000AGU/GUA]
C ₇ H ₁₄ O ₆	[na]	methyl- α -(<i>d</i>)-galactopyranoside				
	$\Delta_{\text{fus}} H$		27.95	397.6		[2000AGU/GUA]
C ₇ H ₁₄ O ₆	[na]	methyl- β -(<i>d</i>)-galactopyranoside				
	$\Delta_{\text{fus}} H$		33.18	450.9		[2000AGU/GUA]
C ₇ H ₁₄ S	[37850-75-2]	allyl <i>tert</i> -butyl sulfide				
	$\Delta_v H$	(319–339)	41.9	332		[1999DYK/SVO]
	$\Delta_v H$	(319–339)	43.1	329	A,EB	[1987STE/MAL, 1962MAC/MAY3]
	$\Delta_v H$	(319–339)	44.8	298	EB	[1962MAC/MAY3]
C ₇ H ₁₅ Br	[629-04-9]	1-bromoheptane				
	$\Delta_{\text{fus}} H$		21.76	214.4		[1950CRO/SMY]
	$\Delta_v H$	(341–481)	47.0	356		[1999DYK/SVO]
	$\Delta_v H$	(323–363)	50.2	298	CGC	[1995CHI/HOS]
	$\Delta_v H$		50.8 ± 0.1	298	C	[1968WAD]
	$\Delta_v H$		50.4 ± 0.2	298	C	[1966WAD]
	$\Delta_v H$	(333–483)	47.5	348	A,E	[1987STE/MAL, 1961LI/ROS]
C ₇ H ₁₅ Br	[1974-04-5]	(<i>dl</i>) 2-bromoheptane				
	$\Delta_v H$	(333–440)	45.0	348	A	[1987STE/MAL, 1999DYK/SVO]
C ₇ H ₁₅ Cl	[629-06-1]	1-chloroheptane				

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(326–462)	45.1	341		[1999DYK/SVO]
	$\Delta_v H$	(313–353)	47.9	298	CGC	[1995CHI/HOS]
	$\Delta_v H$	(300–430)	47.0	298		[1984BOU/FRI, 1991BAS/SVO]
	$\Delta_v H$	(307–434)	46.9	322	A,DTA	[1987STE/MAL, 1969KEM/KRE]
	$\Delta_v H$		47.7 ± 0.1	298	C	[1968WAD]
C₇H₁₅Cl	[1001-89-4]	<i>(dl)</i> 2-chloroheptane				
	$\Delta_v H$	(313–424)	44.8	328	A	[1987STE/MAL, 1999DYK/SVO]
C₇H₁₅Cl₂N	[52802-03-6]	N-methyl- <i>bis</i> (2-chloropropyl)amine				
	$\Delta_v H$	(273–333)	54.6	288	A,GS	[1987STE/MAL, 1948RED/CHA3, 1973KKY/REP]
C₇H₁₅Cl₂N	[621-68-1]	N-propyl- <i>bis</i> (2-chloroethyl)amine				
	$\Delta_v H$	(273–369)	56.8	288	A,GS	[1987STE/MAL, 1948RED/CHA3, 1973KKY/REP]
C₇H₁₅Cl₂N₂O₂P	[50-18-0]	2- <i>bis</i> (2-chloroethyl)amino]tetrahydro-2 <i>H</i> -1,3,2-oxazophosphorine-2-oxide				
	$\Delta_{\text{fus}}H$		33.13	322.6	DSC	[1990DON/DRE]
C₇H₁₅F	[661-11-0]	1-fluoroheptane				
	$\Delta_v H$	(294–416)	40.3	309		[1999VER2]
	$\Delta_v H$	(287–417)	40.8	302	A,E	[1987STE/MAL, 1961LI/ROS]
C₇H₁₅I	[4282-40-0]	1-iodoheptane				
	$\Delta_v H$	(373–513)	55.0	298		[2006BOL/NER, 1961LI/ROS]
	$\Delta_v H$	(373–513)	47.8	388	A,E	[1987STE/MAL, 1961LI/ROS, 1970DYK/VAN]
C₇H₁₅N	[766-09-6]	N-ethylpiperidine				
	$\Delta_v H$	(274–313)	41.1 ± 0.6	294	GS	[1998VER6]
	$\Delta_v H$	(274–313)	40.8 ± 0.6	298	GS	[1998VER6]
C₇H₁₅N	[766-80-6]	1-ethylpiperidine				
	$\Delta_v H$		39.4 ± 0.7	298	C	[2006RIB/CAB6]
C₇H₁₅N	[1484-80-6]	2-ethylpiperidine				
	$\Delta_v H$		42.2 ± 0.9	298	C	[2006RIB/CAB6]
C₇H₁₅N	[766-17-6]	<i>cis</i> 2,6-dimethylpiperidine				
	$\Delta_v H$	(295–365)	39.7 ± 0.1	330		[2009MOK/RAZ]
	$\Delta_v H$	(295–365)	41.3 ± 0.1	298		[2009MOK/RAZ]
	$\Delta_v H$		42.4 ± 0.6	298	C	[2006RIB/CAB5]
Note: Authors indicate that the compound is <i>cis</i> 2,6-dimethylpiperidine; however they give the CAS registry number of 504-03-0 (which does not specify <i>cis</i> or <i>trans</i> isomer).						
C₇H₁₅N	[35794-11-7]	3,5-dimethylpiperidine				
	$\Delta_v H$		49.1 ± 0.6	298	C	[2006RIB/CAB5]
C₇H₁₅N	[1121-92-2]	octahydroazocine				
	$\Delta_v H$	(273–313)	46.5	288	A	[1987STE/MAL]
C₇H₁₅NO	[3040-44-6]	1-piperidineethanol				
	$\Delta_v H$		64.2 ± 0.8	298	C	[2006RIB/CAB4]
C₇H₁₅NO	[1484-84-0]	2-piperidineethanol				
	$\Delta_v H$		75.2 ± 0.5	298	C	[2006RIB/CAB4]
C₇H₁₅NO	[20845-34-5]	1-methyl-2-piperidinemethanol				
	$\Delta_v H$		68.2 ± 0.7	298	C	[2006RIB/CAB4]
C₇H₁₅NO	[13444-24-1]	1-ethyl-3-piperidinol				
	$\Delta_v H$		75.7 ± 0.4	298	C	[2006RIB/CAB2]
C₇H₁₅NO	[na]	N,N-dimethyl- <i>tert</i> -butylcarboxamide				

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$		55.1 ± 0.4	298	ME	[1995ABB/JIM, 1993ABB/JIM]
C ₇ H ₁₅ NO	[1620-14-0]	1-(diethylamino)-2-propanone				
	$\Delta_v H$	(278–318)	47.7 ± 0.3	298	GS	[1994WEL/VER]
C ₇ H ₁₅ NO	[628-62-6]	heptanamide (enanthamide)				
	$\Delta_{\text{sub}} H$	(345–365)	99.6		A	[1954SER/VOI, 1960JON, 1987STE/MAL]
C ₇ H ₁₅ NO	[na]	methyl 2-(N,N-dimethylamino)-2-methylpropanoate				
	$\Delta_v H$	(278–308)	49.2 ± 1.0	293	GS	[1992VER/BEC]
C ₇ H ₁₅ NO ₂	[30220-58-7]	lactic acid N-butylamide				
	$\Delta_v H$	(365–433)	77.4	380	A	[1987STE/MAL]
C ₇ H ₁₅ NO ₂	[na]	lactic acid N-sec-butylamide				
	$\Delta_v H$	(368–418)	74.6	383	A	[1987STE/MAL]
C ₇ H ₁₅ NO ₂	[na]	lactic acid N-isobutylamide				
	$\Delta_v H$	(388–418)	73.5	403	A	[1987STE/MAL]
C ₇ H ₁₅ NO ₂	[2666-93-5]	(<i>d</i>) leucine methyl ester				
	$\Delta_v H$	(320–353)	39.4	366	A	[1987STE/MAL]
C ₇ H ₁₅ NO ₂	[2114-20-7]	hexyl carbamate				
	$\Delta_{\text{sub}} H$	(291–314)	96.2 ± 0.8		GS	[1959DAV/JON]
C ₇ H ₁₅ NO ₄	[69567-10-8]	N-methyl-5-amino-1,5-dideoxy-(<i>d</i>)-glycopyranose				
	$\Delta_{\text{fus}} H$		27.5	425.7		[1994BLU/PRA]
C ₇ H ₁₆	[142-85-5]	heptane				
	$\Delta_{\text{fus}} H$		14.04	182.6		[1996DOM/HEA]
	$\Delta_{\text{sub}} H$		57.9	183	B	[1963BON]
	$\Delta_v H$	(330–371)	34.7	345		[2002SEG/WIS]
	$\Delta_v H$	(328–393)	35.2	343		[2001ORT/GON]
	$\Delta_v H$		36.6	298		[1994RUZ/MAJ]
	$\Delta_v H$	(298–363)	36.1	313		[1984SIP/WIE]
	$\Delta_v H$	(298–338)	36.1	313		[1984MIC/JOS]
	$\Delta_v H$		36.6 ± 0.1	298	C	[1979MAJ/SVO]
	$\Delta_v H$		35.6 ± 0.1	313	C	[1979MAJ/SVO]
	$\Delta_v H$		34.4 ± 0.1	333	C	[1979MAJ/SVO]
	$\Delta_v H$		33.1 ± 0.1	353	C	[1979MAJ/SVO]
	$\Delta_v H$	(278–353)	36.9	298		[1979JAK/TZI]
	$\Delta_v H$		36.55	298		[1971WIL/ZWO]
	$\Delta_v H$	(288–348)	36.4	303		[1967VAN/SOC]
	$\Delta_v H$	(297–375)	36.1	312	A	[1987STE/MAL, 1949FOR/NOR]
	$\Delta_v H$		34.5 ± 0.1	331	C	[1947WAD/TOD]
	$\Delta_v H$		33.2 ± 0.1	350	C	[1947WAD/TOD]
	$\Delta_v H$	(313–398)	35.4	328		[1946THO]
	$\Delta_v H$	(299–372)	36.0	314	MM	[1945WIL/TAY]
	$\Delta_v H$		32.0	371	C	[1940PIT]
	$\Delta_v H$	(310–397)	35.5	325	EB	[1940SMI]
C ₇ H ₁₆	[591-76-4]	2-methylhexane				
	$\Delta_{\text{fus}} H$		9.18	154.9		[1996DOM/HEA]
	$\Delta_v H$	(296–365)	34.6	311	A	[1987STE/MAL]
	$\Delta_v H$		34.9 ± 0.1	298	C	[1979MAJ/SVO]
	$\Delta_v H$		33.9 ± 0.1	313	C	[1979MAJ/SVO]
	$\Delta_v H$		32.7 ± 0.1	333	C	[1979MAJ/SVO]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	Enthalpy							
		$\Delta_v H$			31.3 ± 0.1	353	C	[1979MAJ/SVO]
		$\Delta_v H$			34.8	298		[1971WIL/ZWO]
		$\Delta_v H$	(273–318)		34.8	298		[1961HUF/GRO]
		$\Delta_v H$	(291–364)		34.8	306		[1949FOR/NOR]
C ₇ H ₁₆	[589-34-4]		(dl) 3-methylhexane					
		$\Delta_v H$	(289–366)		35.1	304	A	[1987STE/MAL]
		$\Delta_v H$			35.1 ± 0.1	298	C	[1979MAJ/SVO]
		$\Delta_v H$			34.2 ± 0.1	313	C	[1979MAJ/SVO]
		$\Delta_v H$			32.9 ± 0.1	333	C	[1979MAJ/SVO]
		$\Delta_v H$			31.7 ± 0.1	353	C	[1979MAJ/SVO]
		$\Delta_v H$			35.1	298		[1971WIL/ZWO]
		$\Delta_v H$	(293–366)		34.9	308		[1949FOR/NOR]
C ₇ H ₁₆	[590-35-2]		2,2-dimethylpentane					
		$\Delta_{\text{fus}} H$			5.86	148.1		[1991ACR]
		$\Delta_v H$			32.4 ± 0.1	298	C	[1998SVO/HYN]
		$\Delta_v H$			31.8 ± 0.1	308	C	[1998SVO/HYN]
		$\Delta_v H$			31.4 ± 0.1	315	C	[1998SVO/HYN]
		$\Delta_v H$			31.0 ± 0.1	323	C	[1998SVO/HYN]
		$\Delta_v H$			30.5 ± 0.1	330	C	[1998SVO/HYN]
		$\Delta_v H$			30.1 ± 0.1	338	C	[1998SVO/HYN]
		$\Delta_v H$			29.4 ± 0.1	348	C	[1998SVO/HYN]
		$\Delta_v H$			28.8 ± 0.1	358	C	[1998SVO/HYN]
		$\Delta_v H$			28.1 ± 0.1	368	C	[1998SVO/HYN]
		$\Delta_v H$	(277–354)		33.2	292	A	[1987STE/MAL]
		$\Delta_v H$	(353–483)		30.1	368	A	[1987STE/MAL]
		$\Delta_v H$	(285–353)		32.8	300		[1949FOR/NOR]
		$\Delta_v H$			32.4 ± 0.1	298	C	[1947OSB/GIN]
		$\Delta_v H$			32.2 ± 0.1	298	C	[1947OSB/GIN]
	$\Delta_v H$	(288–353)		32.6	303	MM	[1945WIL/TAY]	
C ₇ H ₁₆	[565-59-3]		(dl) 2,3-dimethylpentane					
		$\Delta_v H$	(309–371)		33.0	324		[1999MON/DEL]
		$\Delta_v H$	(208–286)		35.9	271	A	[1987STE/MAL]
		$\Delta_v H$			34.3 ± 0.1	298	C	[1979MAJ/SVO]
		$\Delta_v H$			33.4 ± 0.1	313	C	[1979MAJ/SVO]
		$\Delta_v H$			32.2 ± 0.1	333	C	[1979MAJ/SVO]
		$\Delta_v H$			31.1 ± 0.1	353	C	[1979MAJ/SVO]
		$\Delta_v H$	(286–365)		34.5	301	A	[1987STE/MAL, 1973KKY/REP]
		$\Delta_v H$	(291–364)		34.4	306		[1949FOR/NOR]
	$\Delta_v H$			34.2 ± 0.1	298	C	[1947OSB/GIN]	
C ₇ H ₁₆	[108-08-7]		2,4-dimethylpentane					
		$\Delta_{\text{fus}} H$			6.85	154		[1996DOM/HEA]
		$\Delta_v H$			32.7 ± 0.1	298	C	[1998SVO/HYN]
		$\Delta_v H$			32.3 ± 0.1	308	C	[1998SVO/HYN]
		$\Delta_v H$			31.9 ± 0.1	315	C	[1998SVO/HYN]
		$\Delta_v H$			31.5 ± 0.1	323	C	[1998SVO/HYN]
		$\Delta_v H$			31.0 ± 0.1	330	C	[1998SVO/HYN]
		$\Delta_v H$			30.6 ± 0.1	338	C	[1998SVO/HYN]
		$\Delta_v H$			30.0 ± 0.1	348	C	[1998SVO/HYN]
		$\Delta_v H$	(284–355)		33.3	299	A	[1987STE/MAL, 1973KKY/REP]
		$\Delta_v H$	(287–354)		33.2	302		[1949FOR/NOR]
	$\Delta_v H$			32.9 ± 0.1	298	C	[1947OSB/GIN]	

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₇ H ₁₆	[562-49-2]	3,3-dimethylpentane				
	$\Delta_{\text{fus}}H$		7.07	138.2		[1996DOM/HEA]
	Δ_vH	(213–281)	34.8	266		[1987STE/MAL]
	Δ_vH	(280–360)	33.6	295		[1987STE/MAL]
	Δ_vH		33.0 ± 0.1	298	C	[1981HOS/SCO2]
	Δ_vH		33.0	298		[1971WIL/ZWO]
	Δ_vH	(287–360)	33.2	302		[1949FOR/NOR]
	Δ_vH		33.0 ± 0.1	298	C	[1947OSB/GIN]
	(285–360)	33.3	300	MM	[1987STE/MAL, 1945WIL/TAY]	
C ₇ H ₁₆	[617-78-7]	3-ethylpentane				
	$\Delta_{\text{fus}}H$		9.55	154.6		[1996DOM/HEA]
	Δ_vH		35.1 ± 0.1	298	C	[1998SVO/HYN]
	Δ_vH		34.5 ± 0.1	308	C	[1998SVO/HYN]
	Δ_vH		34.1 ± 0.1	315	C	[1998SVO/HYN]
	Δ_vH		33.7 ± 0.1	232	C	[1998SVO/HYN]
	Δ_vH		33.3 ± 0.1	330	C	[1998SVO/HYN]
	Δ_vH		32.7 ± 0.1	338	C	[1998SVO/HYN]
	Δ_vH		32.2 ± 0.1	348	C	[1998SVO/HYN]
	Δ_vH	(291–368)	35.2	306	A	[1987STE/MAL]
	Δ_vH		35.2	298		[1971WIL/ZWO]
	Δ_vH		35.2 ± 0.1	298	C	[1947OSB/GIN]
	Δ_vH	(294–367)	35.0	309		[1945WIL/TAY]
Δ_vH	(308–391)	34.4	323	EB	[1941SMI, 1984BOU/FRI]	
C ₇ H ₁₆	[464-06-2]	2,2,3-trimethylbutane				
	$\Delta_{\text{us}}H$		2.38	121		
	$\Delta_{\text{fus}}H$		2.2	247.7		[1996DOM/HEA]
	Δ_vH	(284–355)	32.4	299	A	[1987STE/MAL]
	Δ_vH	(353–483)	29.9	368	A	[1987STE/MAL]
	Δ_vH		32	298		[1971WIL/ZWO]
	Δ_vH	(286–355)	32.3	301		[1949FOR/NOR]
	Δ_vH		31.2 ± 0.1	314	C	[1947WAD/TOD]
Δ_vH	(296–378)	31.9	311	EB	[1941SMI]	
C ₇ H ₁₆ N ₂ O	[2158-11-4]	1-hexyl urea				
	$\Delta_{\text{fus}}H$		22.4	383	DSC	[2005HAS/TAJ]
	$\Delta_{\text{fus}}H$		26.5	380.2		[1999WEL/DRU]
C ₇ H ₁₆ N ₂ S	[26536-60-7]	1,3-propylthiourea				
	$\Delta_{\text{fus}}H$		22.9	342.6	DSC	[2000DEL/JOZ]
	$\Delta_{\text{sub}}H$		134.9 ± 3	298	B	[2000DEL/JOZ]
	$\Delta_{\text{sub}}H$		132.5 ± 3.0	298	C	[1994TER/PIA]
	Δ_vH	(346–394)	107 ± 3.0	370	ME,TE	[1994TER/PIA]
C ₇ H ₁₆ O	[919-94-8]	<i>tert</i> -amyl ethyl ether				
	Δ_vH		39.2 ± 0.4	298	GS	[UR/VER, 2002VER, 2003VER/KRA]
	Δ_vH		38.2 ± 0.2	298	C	[2002VAR/PAS]
	Δ_vH	(318–374)	35.7	333	EB	[2002VAR/PAS]
	Δ_vH	(320–374)	35.6	335	EB	[1994KRA/GME]
C ₇ H ₁₆ O	[29072-93-3]	propyl <i>tert</i> -butyl ether				
	$\Delta_{\text{fus}}H$		9.87	179.6		[2001VAR/DRU, 2004DOR/YAN]
	Δ_vH		38.3	298	CGC	[UR/VER, 2002VER, 2003VER/KRA]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound					
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference	
		$\Delta_v H$	(315–370)	37.2 ± 0.6	298	EB	[2002VAR/PAS2]
		$\Delta_v H$		36.6 ± 0.2	298	C	[2002VAR/PAS2, 2004DOR/YAN]
C ₇ H ₁₆ O	[17348-59-3]	isopropyl <i>tert</i> -butyl ether					
		$\Delta_{\text{fus}} H$		8.46	184.8		[2001VAR/DRU, 2004DOR/YAN]
		$\Delta_v H$		36.2	298	CGC	[UR/VER, 2002VER, 2003VER/KRA]
		$\Delta_v H$	(305–360)	34.4 ± 0.6	298	EB	[2002VAR/PAS2]
		$\Delta_v H$		34.5 ± 0.2	298	C	[2002VAR/PAS2, 2004DOR/YAN]
		$\Delta_v H$	(307–360)	34.0	322	EB	[1994KRA/GME]
C ₇ H ₁₆ O	[111-70-6]	1-heptanol					
		$\Delta_{\text{fus}} H$		18.35	239.9		[2003VAN/GAB]
		$\Delta_{\text{fus}} H$		18.16	240.4		[1996DOM/HEA]
		$\Delta_v H$	(348–443)	67.1	298		[2006NAS/NEU]
		$\Delta_v H$	(283–323)	66.9 ± 0.4	298	GS	[2005ROG/PIS]
		$\Delta_v H$	(323–373)	66.5	298	CGC	[1995CHI/HOS]
		$\Delta_v H$	(373–423)	66.4	298	CGC	[1995CHI/HOS]
		$\Delta_v H$	(258–363)	65.2	310		[1992NGU/KAS]
		$\Delta_v H$	(335–450)	62.5	350	A	[1987STE/MAL]
		$\Delta_v H$		66.8 ± 0.2	298	C	[1977MAN/SEL]
		$\Delta_v H$	(333–449)	65.2	348		[1973WIL/ZWO]
		$\Delta_v H$	(336–450)	62.6	351	DTA	[1987STE/MAL, 1969KEM/KRE]
		$\Delta_v H$	(333–425)	62.9	348		[1935BUT/RAM, 1984BOU/FRI]
C ₇ H ₁₆ O	[543-49-7]	<i>(dl)</i> 2-heptanol					
		$\Delta_v H$	(275–312)	62.1 ± 0.4	298	GS	[2007VER/SCH]
		$\Delta_v H$	(244–338)	66.1	259		[1999NGU/BER]
		$\Delta_v H$	(351–433)	54.4	366		[1984SAC/MAR]
		$\Delta_v H$	(357–431)	51.6	372	A	[1987STE/MAL, 1975BRA/AND]
		$\Delta_v H$	(323–433)	59.8	338		[1973WIL/ZWO]
C ₇ H ₁₆ O	[589-82-2]	<i>(dl)</i> 3-heptanol					
		$\Delta_v H$	(244–333)	67.0	259		[1999NGU/BER]
		$\Delta_v H$	(325–430)	60.3	340	A	[1987STE/MAL]
		$\Delta_v H$	(263–295)	64.7	280	A	[1987STE/MAL, 1979THO/MEA]
		$\Delta_v H$	(349–430)	53.1	364		[1984SAC/MAR]
		$\Delta_v H$	(328–429)	59.2	343		[1973WIL/ZWO]
C ₇ H ₁₆ O	[589-55-9]	4-heptanol					
		$\Delta_v H$		62.4 ± 0.3	298		[2007VER/SCH]
		$\Delta_v H$	(320–428)	58.2	335	A	[1987STE/MAL]
		$\Delta_v H$	(349–428)	53.1	364		[1984SAC/MAR]
		$\Delta_v H$	(282–320)	63.1	297	A	[1987STE/MAL, 1975CAB/CON2]
		$\Delta_v H$	(320–428)	56.9	335		[1973WIL/ZWO]
C ₇ H ₁₆ O	[624-22-6]	2-methyl-1-hexanol					
	$\Delta_v H$	(343–438)	53.5	390		[1973WIL/ZWO]	
C ₇ H ₁₆ O	[13231-81-7]	3-methyl-1-hexanol					
	$\Delta_v H$	(353–445)	57.4	399		[1973WIL/ZWO]	
C ₇ H ₁₆ O	[818-49-5]	4-methyl-1-hexanol					
	$\Delta_v H$	(348–448)	62.6	363		[1973WIL/ZWO]	
C ₇ H ₁₆ O	[625-23-0]	2-methyl-2-hexanol					
		$\Delta_v H$	(274–306)	58.6 ± 0.4	298	GS	[2005ROG/PIS]
		$\Delta_v H$	(311–415)	54.5	326	A	[1987STE/MAL, 1973WIL/ZWO]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₇ H ₁₆ O	[627-59-8] $\Delta_v H$	5-methyl-2-hexanol (348–428)	49.4	388		[1973WIL/ZWO]
C ₇ H ₁₆ O	[617-29-8] $\Delta_v H$	2-methyl-3-hexanol (323–420)	55.7	338		[1973WIL/ZWO]
C ₇ H ₁₆ O	[597-96-6] $\Delta_v H$	3-methyl-3-hexanol (323–416)	53.6	338		[1973WIL/ZWO]
C ₇ H ₁₆ O	[623-55-2] $\Delta_v H$	5-methyl-3-hexanol (275–311)	59.8 ± 0.3	298	GS	[2005ROG/PIS]
C ₇ H ₁₆ O	[6570-87-2] $\Delta_v H$	3,4-dimethyl-1-pentanol (393–438)	50.3	388		[1973WIL/ZWO]
C ₇ H ₁₆ O	[625-06-9] $\Delta_v H$	2,4-dimethyl-2-pentanol (328–408)	49.7	343		[1973WIL/ZWO]
C ₇ H ₁₆ O	[3970-62-5] $\Delta_v H$	2,2-dimethyl-3-pentanol (318–411)	51.4	333		[1973WIL/ZWO]
C ₇ H ₁₆ O	[595-41-5] $\Delta_v H$	2,3-dimethyl-3-pentanol (318–413)	53.2	333		[1973WIL/ZWO]
C ₇ H ₁₆ O	[600-36-2] $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$	2,4-dimethyl-3-pentanol (307–412)	51.8 48.8 45.7 53.6	328 343 358 322	C C C A	[1996WEB/DEF2] [1996WEB/DEF2] [1996WEB/DEF2] [1987STE/MAL, 1973WIL/ZWO]
C ₇ H ₁₆ O	[597-49-9] $\Delta_v H$ $\Delta_v H$ $\Delta_v H$	3-ethyl-3-pentanol (275–311) (317–408) (308–416)	57.3 ± 0.2 51.3 55.2	298 332 323	GS A	[2005ROG/PIS] [1987STE/MAL, 1973KKY/REP] [1973WIL/ZWO]
C ₇ H ₁₆ O	[18371-13-6] $\Delta_v H$	2-methyl-2-ethyl-1-butanol (358–428)	55.7	373		[1973WIL/ZWO]
C ₇ H ₁₆ O	[594-83-2] $\Delta_v H$	2,3,3-trimethyl-2-butanol (298–363)	48.7	313	MM	[1985WIE/SIP]
C ₇ H ₁₆ O ₂	[13343-98-1] $\Delta_v H$	1-butoxy-2-methoxyethane 47.8 ± 0.1	298		C	[1970KUS/WAD]
C ₇ H ₁₆ O ₂	[18854-31-4] $\Delta_v H$	1-propoxy-2-ethoxyethane 46.8 ± 0.1	298		C	[1970KUS/WAD]
C ₇ H ₁₆ O ₂	[3459-83-4] $\Delta_v H$	1,3-diethoxypropane 45.9 ± 0.2	298		C	[1972MAN2]
C ₇ H ₁₆ O ₂	[126-84-1] $\Delta_v H$ $\Delta_v H$ $\Delta_v H$	2,2-diethoxypropane (273–308) (273–308) (286–304)	43.2 ± 0.4 43.9 U28.2	298 298 295	GS GS A,I	[2002VER] [1998VER/PEN] [1987STE/MAL, 1962STE/DOR]
C ₇ H ₁₆ O ₂	[57018-52-7] $\Delta_v H$	1- <i>tert</i> -butoxy-2-propanol (346–420)	45.4	361	EB	[2001BER/WIC]
C ₇ H ₁₆ O ₂	[141-73-1] $\Delta_v H$	4-methyl-4-methoxy-2-pentanol (343–423)	46.6	358	A	[1987STE/MAL, 1973KKY/REP]
C ₇ H ₁₆ O ₂	[629-30-1]	1,7-heptanediol				

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		T _m (K)	Method	Reference
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)			
		$\Delta_{\text{fus}}H$	21.3	295.2		[1991ACR]
		Δ_vH	92.4	341		[1994PIA/FER, 2006UMN/KWE]
		Δ_vH	96.5 ± 3.2	298		[1994PIA/FER, 2006UMN/KWE]
		Δ_vH	93.8	323		[1990KNA/SAB, 2006UMN/KWE]
		Δ_vH	96.2 ± 1.2	298		[1990KNA/SAB, 2006UMN/KWE]
C ₇ H ₁₆ O ₂	[115-76-4]	2,2-diethyl-1,3-propanediol				
		Δ_vH	(343–380) 80.2 ± 0.2	298		[2007VER]
C ₇ H ₁₆ O ₃	[6881-94-3]	diethylene glycol monopropyl ether				
		Δ_vH	(369–404) 65.3	384	A	[1987STE/MAL]
C ₇ H ₁₆ O ₃	[6881-94-3]	2-(2-propoxyethoxy)ethanol				
		Δ_vH	(378–495) 65.7 ± 0.8	298	EB	[1996STE/CHI]
C ₇ H ₁₆ O ₃	[122-51-0]	triethoxymethane (triethyl orthoformate)				
		Δ_vH	(293–323) 49.0	308	A	[1987STE/MAL]
		Δ_vH	47.8 ± 0.1	298	C	[1985MAR/MAN]
		Δ_vH	46.0 ± 0.8	298		[1971PIH/TUO]
		Δ_vH	(278–419) 47.2	293	A	[1987STE/MAL, 1947STU]
C ₇ H ₁₆ O ₃	[4431-82-7]	3,5,7,9-tetraoxaundecane				
		Δ_vH	53.6 ± 0.7	298	C	[1969MAN]
C ₇ H ₁₆ O ₃	[51452-08-5]	<i>tert</i> -pentylperoxyethanol				
		Δ_vH	70.1 ± 2.5			[1983VAN/KAC]
C ₇ H ₁₆ O ₄	[38578-50-6]	3- <i>tert</i> -butyldioxy-1,2-propanediol				
		Δ_vH	88.0 ± 2.6			[1983VAN/KAC]
C ₇ H ₁₆ S	[1639-09-4]	1-heptanethiol				
		$\Delta_{\text{fus}}H$	25.4	229.9		[1996DOM/HEA]
		Δ_vH	(273–345) 49.5	288		[1999DYK/SVO]
		Δ_vH	50.6 ± 0.2	298		[1966GOO/DEP, 1966OSB/DOU]
		Δ_vH	(373–472) 45.0	388	A,EB	[1987STE/MAL, 1965DOU/OSB, 1966OSB/DOU]
C ₇ H ₁₆ S	[628-00-2]	2-heptanethiol				
		Δ_vH	(343–437) 44.1	358		[1999DYK/SVO]
		Δ_vH	(343–471) 47.2	360		[1999DYK/SVO]
		Δ_vH	(341–443) 44.2	356	A	[1987STE/MAL, 1973KKY/REP]
C ₇ H ₁₆ S ₂	[62224-02-6]	1,7-heptanedithiol				
		Δ_vH	(392–526) 59.0	407	A	[1987STE/MAL, 1973KKY/REP, 1999DYK/SVO]
C ₇ H ₁₇ N	[na]	<i>tert</i> -butylisopropylamine				
		Δ_vH	(275–299) 35.7 ± 1.0	287		[1997VER]
C ₇ H ₁₇ N	[39099-23-5]	N-butyl isopropylamine				
		Δ_vH	42.1 ± 0.1	298	C	[1979PET/MAJ]
		Δ_vH	40.9 ± 0.1	313	C	[1979PET/MAJ]
		Δ_vH	39.9 ± 0.1	328	C	[1979PET/MAJ]
		Δ_vH	38.7 ± 0.1	343	C	[1979PET/MAJ]
		Δ_vH	37.6 ± 0.1	358	C	[1979PET/MAJ]
		Δ_vH	(325–395) 40.0	340	C	[1979PET/MAJ]
C ₇ H ₁₇ N	[111-68-2]	heptylamine				
		Δ_vH	(323–373) 49.9	298	CGC	[1995CHI/HOS]
		Δ_vH	(326–430) 46.5	341	A	[1987STE/MAL]
		Δ_vH	50.0 ± 0.1	298	C	[1969WAD]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₇ H ₁₇ NO	[7352-03-6] $\Delta_v H$	N-(ethoxymethyl)diethylamine (285–400)	39.6	300	A	[1987STE/MAL]
C ₇ H ₁₇ N ₃	[934-98-5] $\Delta_v H$	1-(2-aminoethyl)-4-methylpiperazine (298–332)	64.0 ± 0.2	298	GS	[2010EFI/EME]
C ₇ H ₁₇ O ₂ PS ₃	[298-02-2] $\Delta_v H$	O,O-diethyl-S-[(ethylthio)methyl]dithiophosphate (283–387)	70.8	298	A	[1987STE/MAL, 1973KKY/REP, 1999DYK/SVO]
C ₇ H ₁₇ O ₃ P	[1445-75-6] $\Delta_v H$	diisopropyl methylphosphonate (253–468)	62.1	253	GS	[2009BUT/BUC]
	$\Delta_v H$	(253–468)	58.9	283	GS	[2009BUT/BUC]
	$\Delta_v H$	(253–468)	57.6	298	GS	[2009BUT/BUC]
	$\Delta_v H$	(253–468)	56.5	313	GS	[2009BUT/BUC]
	$\Delta_v H$	(253–468)	54.2	353	GS	[2009BUT/BUC]
	$\Delta_v H$	(253–468)	52.4	393	GS	[2009BUT/BUC]
C ₇ H ₁₈ N ₂	[646-19-5] $\Delta_{\text{fus}} H$	1,7-heptanediamine	36.95	298.5	DSC	[2002DAL/DEL]
	$\Delta_v H$	(273–313)	46.5	288	A	[1987STE/MAL]
C ₇ H ₁₈ N ₂	[104-78-9] $\Delta_v H$	N,N-diethyl-1,3-propanediamine (329–443)	46.4	344	A	[1987STE/MAL]
C ₇ H ₁₈ N ₂ O	[5966-51-8] $\Delta_v H$	1,3-bis(dimethylamino)-2-propanol (355–450)	50.3	370	A	[1987STE/MAL]
C ₇ H ₁₈ N ₃	[67727-91-7] $\Delta_v H$	N,N-diethyl-2-(1-methylhydrazino)ethanamine (283–313)	61.8	298	A	[1987STE/MAL]
C ₇ H ₂₀ N ₄	[4741-99-5] $\Delta_v H$	1,4,8,11-tetraazaundecane (332–348)	98.3 ± 1.3	340	TE	[1983CLA/COR]
C ₈ BrF ₁₇	[423-55-2] $\Delta_{\text{us}} H$	1-bromoperfluorooctane	1.6	146.4		
	$\Delta_{\text{fus}} H$		12.13	278.9		[1997VAR/DRU]
	$\Delta_v H$	(288–332)	45.6 ± 0.4	298		[2005DIA/GON]
C ₈ Cl ₄ N ₂	[1897-45-6] $\Delta_{\text{us}} H$	2,4,5,6-tetrachloro-1,3-benzenedicarbonitrile	4.03	423.7		
	$\Delta_{\text{fus}} H$		NA			[2005RON/GU]
	$\Delta_{\text{fus}} H$		30	526.2	DSC	[1990DON/DRE]
	$\Delta_{\text{sub}} H$	(363–418)	109.1	378	ME,GS	[1987STE/MAL, 1980DEP]
	$\Delta_v H$	(373–403)	67.0		GC	[2007GOE/MCC]
C ₈ F ₈ O ₂	[14533-84-7] $\Delta_v H$	trifluoroacetic acid, pentafluorophenyl ester	42.1			[1976HOP/DES]
C ₈ F ₈ O ₄	[59483-83-9] $\Delta_v H$	carbonoperoxoic acid, O-(pentafluorophenyl) O,O-(trifluoromethyl) ester	51.8			[1976FAL/DES2]
C ₈ F ₁₆	[335-21-7] $\Delta_v H$	perfluoroethylcyclohexane (308–512)	37.2	323		[1999DYK/SVO]
	$\Delta_v H$	(310–400)	38.6	298		[1984BOU/FRI, 1991BAS/SVO]
	$\Delta_v H$		38.7 ± 0.4	298		[1981VAR/BUL]
	$\Delta_v H$	(311–411)	37.1	326	A	[1987STE/MAL, 1970DYK/VAN, 1959GOO/DOU, 1999DYK/SVO]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₈ F ₁₆	[335-27-3] $\Delta_v H$	perfluoro-1,3-dimethylcyclohexane (308–375)	37.4	323		[1999DYK/SVO]
C ₈ F ₁₆	[na] $\Delta_v H$	<i>cis/trans</i> perfluoro-1,3-dimethylcyclohexane	38.6 ± 0.1	298	C	[1996VAR/DRU]
C ₈ F ₁₆ N ₂	[57682-63-0] $\Delta_v H$	2,2,2-trifluoro-N'-[1,2,2-tetrafluoro-1-(trifluoromethyl)ethyl]-N-[2,2,2-trifluoro-1-(trifluoromethyl)ethylidene]-ethanimidamide	32.8			[1975PET/SHR3]
C ₈ F ₁₆ O	[na] $\Delta_v H$	perfluoro-2-butyltetrahydrofuran (383–433)	34.7	408	EST	[1960YAR/KAY]
C ₈ F ₁₈	[307-34-6] $\Delta_{\text{ms}}H$ $\Delta_{\text{fus}}H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$	perfluorooctane (289–333) (437–503) (309–378) (310–379)	3.14 9.58 39.9 32 41.2 ± 0.8 41.1 ± 0.1 39.5	176.5 254.2 298 452 298 298 325		[1986STA] [2004DIA/CAC] [1999DYK/SVO] [1981VAR/BUL] [1981VAR/BUL] [1987STE/MAL, 1962KRE, 1970DYK/VAN]
C ₈ F ₁₈ N ₂ OS	[66632-47-1] $\Delta_v H$	<i>bis</i> (1,1,1,3,3,3-hexafluoro-2-propanamino)oxobis(trifluoromethyl)sulfur (273–333)	39.6	288	A	[1987STE/MAL, 1978KIT/SHR, 1999DYK/SVO]
C ₈ F ₁₈ N ₂ S	[37826-45-2] $\Delta_v H$	<i>S,S-bis</i> (trifluoromethyl)-N-[2,2,2-trifluoro-1-(trifluoromethyl)-1-[(2,2,2-trifluoro-1-trifluoromethyl)ethylidene]amino]ethyl sulfilimine (329–373)	41.1	344	A	[1987STE/MAL, 1972SWI/SHR]
C ₈ F ₁₈ O	[308-48-5] $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$	<i>bis</i> (nonafluorobutyl) ether (315–374) (343–375) (288–313) (374–413)	40.3 ± 0.8 40.7 ± 0.1 36.6 42.2 56.3	298 298 358 300 389	EB C A A A	[1989VAR/PAS] [1989VAR/PAS] [1987STE/MAL] [1987STE/MAL] [1987STE/MAL, 1999DYK/SVO]
C ₈ F ₁₈ O ₂	[na] $\Delta_v H$	dodecafluoro-1,6- <i>bis</i> (trifluoromethoxy)hexane (293–353)	33.6	323		[1999DYK/SVO]
C ₈ F ₁₈ O ₃ S	[53517-90-1] $\Delta_v H$	<i>bis</i> (1,1,1,3,3,3-hexafluoro-2-(trifluoromethyl)-2-propanol) sulfite (293–353)	38.7			[1975DEM/KOV]
C ₈ F ₂₀ N ₂ S	[65844-11-3] $\Delta_v H$	difluoro[1,1,1,3,3,3-hexafluoro-N-[2,2,2-trifluoro-1-(trifluoromethyl)-ethylidene-2,2-propanediamino(2-)-N]- <i>bis</i> (trifluoromethyl) sulfur (293–353)	39.3	390	I	[1978KIT/SHR]
C ₈ HCl ₄ F ₁₁ O ₂	[2923-68-4] $\Delta_v H$	3,5,7,8-tetrachloro-2,2,3,4,4,5,6,6,7,8,8-undecafluorooctanoic acid (373–553)	70.6	388	A	[1987STE/MAL, 1957BAR/SEF, 1999DYK/SVO]
C ₈ HF ₁₅ O ₂	[335-67-1] $\Delta_{\text{sub}}H$	perfluorooctanoic acid (298–318)	88.9		GS	[2008BAR/BOT]
C ₈ HF ₁₆ NO	[54181-87-2] $\Delta_v H$	1,1,1,2,3,3,3-heptafluoro-N-[2,2,2-trifluoro-1-[2,2,2-trifluoro-1-(trifluoromethyl)ethoxy]ethylidene]-2-propanamine (298–318)	36	364		[1975PET/SHR]
C ₈ HF ₁₇	[335-65-9] $\Delta_v H$	1,1,1,2,2,3,3,4,4,5,5,6,6,7,7,8,8-heptafluorooctane (288–332)	43.4 ± 0.2	298		[2005DIA/GON]
C ₈ H ₂ Cl ₄ N ₂	[25983-14-6] $\Delta_{\text{fus}}H$	2,3,6,7-tetrachloroquinoxaline (298–318)	29.6	446		[2000MON/HIL2]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{sub}}H$		105.9 ± 2.2	298	ME	[2004MOR/MIR]
	$\Delta_{\text{sub}}H$	(347–361)	106.2 ± 0.3	354	ME	[2000MON/HIL2]
	$\Delta_{\text{sub}}H$		108.2 ± 1.9	298	ME	[2000MON/HIL2]
C₈H₂F₁₆	[307-99-3]	1,1,2,2,3,3,4,4,5,5,6,6,7,7,8,8-hexadecafluorooctane				
	Δ_vH	(298–323)	41.1	310	A	[1987STE/MAL, 1999DYK/SVO]
C₈H₃ClF₆	[327-76-4]	4-chloro-1,3-bis(trifluoromethyl)benzene				
	Δ_vH	(275–353)	48.0	290		[1987STE/MAL, 1946FIE/SAY, 1970DYK/VAN, 1999DYK/SVO]
C₈H₃ClF₆	[328-72-3]	5-chloro-1,3-bis(trifluoromethyl)benzene				
	Δ_vH	(275–353)	46.2	290	A	[1987STE/MAL, 1946FIE/SAY, 1970DYK/VAN, 1999DYK/SVO]
C₈H₃Cl₄F₃	[328-82-5]	1,1,1-trifluoro-2,2-dichloro-2-(3,4-dichlorophenyl)ethane				
	Δ_vH	(417–461)	56.9	432	A	[1987STE/MAL, 1999DYK/SVO]
C₈H₃F₅O₂	[19220-93-0]	acidic acid, pentafluorophenyl ester				
	Δ_vH	(283–322)	48.1	298	A	[1987STE/MAL, 1999DYK/SVO]
C₈H₃F₁₅O	[307-30-2]	2,2,3,3,4,4,5,5,6,6,7,7,8,8,8-pentadecafluoro-1-octanol				
	Δ_vH	(350–437)	53.3	365	A	[1987STE/MAL, 1999DYK/SVO]
C₈H₃NO₅	[641-70-3]	3-nitrophthalic anhydride				
	$\Delta_{\text{fus}}H$		18.4	436.2	DSC	[1977CAS/VEC]
C₈H₃NO₅	[5466-84-2]	4-nitrophthalic anhydride				
	$\Delta_{\text{fus}}H$		17.14	388.2	DSC	[1977CAS/VEC]
C₈H₄ClF₃O	[321-31-3]	trifluoromethyl 3-chlorophenyl ketone				
	Δ_vH	(366–405)	52.7	386	A	[1987STE/MAL, 1999DYK/SVO]
C₈H₄Cl₂N₂	[2213-63-0]	2,3-dichloroquinoxaline				
	$\Delta_{\text{fus}}H$		24.36	424.4	DSC	[2000MON/HIL2]
	$\Delta_{\text{sub}}H$		91.8 ± 1.1	298	ME	[2004MOR/MIR]
	$\Delta_{\text{sub}}H$	(313–329)	92.4 ± 0.4	321	ME	[2000MON/HIL2]
	$\Delta_{\text{sub}}H$		93.1 ± 0.9	298	ME	[2000MON/HIL2]
C₈H₄Cl₂O₂	[99-63-8]	isophthaloyl chloride				
	Δ_vH	(443–550)	61.5	458	A	[1987STE/MAL, 1999DYK/SVO]
C₈H₄Cl₂O₂	[88-95-9]	phthaloyl chloride				
	Δ_vH	(391–549)	58.0	406	A	[1987STE/MAL, 1947STU, 1999DYK/SVO]
C₈H₄Cl₂O₂	[100-20-9]	terephthaloyl chloride				
	$\Delta_{\text{us}}H$		2.34	337.3		
	$\Delta_{\text{fus}}H$		21.1	356.1		[1996DOM/HEA]
	Δ_vH	(454–473)	56.2	463	A	[1987STE/MAL, 1999DYK/SVO]
C₈H₄Cl₃F₃	[309-12-6]	1,1,1-trifluoro-2,2-dichloro-(3-chlorophenyl)ethane				
	Δ_vH	(387–475)	49.6	431		[1999DYK/SVO]
	Δ_vH	(387–474)	50.4	402	A	[1987STE/MAL]
C₈H₄F₆	[402-31-3]	1,3-bis(trifluoromethyl)benzene				
	Δ_vH	(275–353)	42.4	290	A	[1987STE/MAL, 1951POT/SAY, 1970DYK/VAN, 1999DYK/SVO]
C₈H₄F₆	[433-19-2]	1,4-bis(trifluoromethyl)benzene				
	Δ_vH	(287–390)	41.8	302	A	[1987STE/MAL, 1951POT/SAY, 1999DYK/SVO]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound					
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference	
C ₈ H ₄ N ₂	[91-15-6]	1,2-dicyanobenzene					
	$\Delta_{\text{fus}}H$		20.0	414.1		[1982KAR/SHV]	
C ₈ H ₄ N ₂	[626-17-5]	1,3-dicyanobenzene					
	$\Delta_{\text{sub}}H$		86.9 ± 1.5	298	GS	[1980SAT/SAK]	
C ₈ H ₄ N ₂	[623-26-7]	1,4-dicyanobenzene					
	$\Delta_{\text{sub}}H$		89.7 ± 1.8	298	ME	[1992ACR/TUC]	
C ₈ H ₄ N ₂ O ₂	[3729-34-8]	1,4-dicyanobenzene di-N-oxide					
	$\Delta_{\text{sub}}H$		73.0 ± 2.0	298	ME	[1992ACR/TUC]	
C ₈ H ₄ O ₂	[6383-11-5]	benzocyclobutenedione					
	$\Delta_{\text{sub}}H$	(304–367)	U 89.5	336		[1989ROR/RUT]	
C ₈ H ₄ O ₃	[85-44-9]	phthalic anhydride					
	$\Delta_{\text{fus}}H$		23.09	403.3	DSC	[1991ACR, 1990DON/DRE]	
	$\Delta_{\text{fus}}H$		22.1	404.5	DSC	[1979DAS/DHA]	
	$\Delta_{\text{sub}}H$	(313–383)	87.9	348	A	[1987STE/MAL, 1972AMI/VAK]	
	$\Delta_{\text{sub}}H$	(333–403)	84.4 ± 1.2	388	GS	[1979DAS/DHA]	
	$\Delta_{\text{sub}}H$		81 ± 1		C	[1971BEE/LIN]	
	$\Delta_{\text{sub}}H$	(303–333)	88.4 ± 1.2	318		[1946CRO/FEE, 1970COX/PIL, 1960JON]	
	Δ_vH	(407–558)	52.1	422	A	[1987STE/MAL]	
	Δ_vH	(411–450)	63.9 ± 2.5	422	GS	[1979DAS/DHA]	
	Δ_vH		63.1			[1952GOT]	
C ₈ H ₅ Br ₃	[24162-65-0]	2,4,5-tribromostyrene					
	$\Delta_{\text{fus}}H$		25.1	340.3		[1993OIS/HOR]	
	C ₈ H ₅ Cl ₂ F ₃	[309-10-4]	1,1,1-trifluoro-2,2-dichloro-2-phenylethane				
		Δ_vH	(365–446)	47.2	380	A	[1987STE/MAL, 1999DYK/SVO]
	C ₈ H ₅ Cl ₂ N	[40626-45-7]	α,α -dichlorophenylacetonitrile				
		Δ_vH	(329–497)	57.2	344	A	[1987STE/MAL, 1947STU]
	C ₈ H ₅ Cl ₃ O ₂	[85-34-7]	2,3,6-trichlorophenylacetic acid				
		$\Delta_{\text{fus}}H$		22.43	432.3	DSC	[1991ACR, 1990DON/DRE]
	C ₈ H ₅ Cl ₃ O ₃	[93-76-5]	(2,4,5-trichlorophenoxy)acetic acid				
		$\Delta_{\text{fus}}H$		38.0	428.7	DSC	[1991ACR, 1990DON/DRE]
C ₈ H ₅ Cl ₅	[606-07-5]	pentachloroethylbenzene					
	Δ_vH	(369–572)	58.8	384	A	[1987STE/MAL, 1947STU, 1999DYK/SVO]	
C ₈ H ₅ F ₃ OS ₂	[4552-64-1]	1,1,1-trifluoro-4-(2-thienyl)-4-mercapto-3-buten-2-one					
	$\Delta_{\text{sub}}H$		95.1 ± 3.7	298	C	[1997RIB/SAN]	
C ₈ H ₅ F ₃ O ₂ S	[15788-02-0]	1,1,1-trifluoro-4-(2-thienyl)-4-hydroxy-3-buten-2-one					
	$\Delta_{\text{sub}}H$		86.2 ± 0.6	298	C	[1997RIB/SAN]	
			86.2 ± 0.6	298	ME	[1992RIB/MON]	
C ₈ H ₅ F ₃ O ₃	[326-90-9]	4,4,4-trifluoro-1-(2-furanyl)-butane-1,3-dione					
	$\Delta_{\text{sub}}H$		70 ± 10	298		[1997RIB/GON]	
C ₈ H ₅ F ₅ O	[434-45-7]	2,2,2-trifluoroacetophenone					

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(342–425)	43.1	357	A	[1987STE/MAL, 1999DYK/SVO]
C₈H₅F₁₄OP	[na]	ethyl bis(heptafluoropropyl)phosphinite				
	$\Delta_v H$	(303–393)	41.9	348		[1959EME/SMI]
C₈H₅NO	[613-90-1]	benzoylnitrile				
	$\Delta_{\text{sub}} H$	(292–304)	78.7 ± 4.2	298		[1969LEB/DNE, 1977PED/RYL, 1987STE/MAL]
	$\Delta_v H$	(318–481)	52.0	333	A	[1987STE/MAL, 1947STU]
C₈H₅NO₂	[85-41-6]	phthalimide				
	$\Delta_{\text{fus}} H$		28.6	507.2	DSC	[1978CAN]
	$\Delta_{\text{sub}} H$	(347–365)	104.0 ± 0.4	356	ME	[2006RIB/SAN]
	$\Delta_{\text{sub}} H$	(347–365)	106.3 ± 1.3	298	ME	[2006RIB/SAN]
	$\Delta_{\text{sub}} H$	(378–418)	82.8	393	RG	[1987STE/MAL, 1956KLO]
C₈H₅NO₂	[91-56-5]	1 <i>H</i> -indole-2,3-dione (isatin)				
	$\Delta_{\text{fus}} H$		27.82	475.7	DSC	[2003MAR/AVI]
	$\Delta_{\text{sub}} H$		118.8 ± 5.1	298	C	[2003MAT/MIR2]
C₈H₅NO₂	[4421-09-4]	5-cyano-1,3-benzodioxole				
	$\Delta_{\text{fus}} H$		20.79	366.3	DSC	[2007MAT/SOU]
	$\Delta_{\text{sub}} H$		90.9 ± 0.9	298	C	[2007MAT/SOU]
C₈H₅NO₂	[3839-22-3]	2-cyanobenzoic acid				
	$\Delta_{\text{sub}} H$		114.6 ± 1.3	298	C	[2008RIB/AMA2]
C₈H₅NO₂	[1877-72-1]	3-cyanobenzoic acid				
	$\Delta_{\text{sub}} H$		116.6 ± 0.9	298	C	[2008RIB/AMA2]
C₈H₅NO₂	[619-65-8]	4-cyanobenzoic acid				
	$\Delta_{\text{sub}} H$		112.8 ± 0.4	298		[2008RIB/AMA2]
C₈H₅NO₃	[118-48-9]	isatoic anhydride				
	$\Delta_{\text{sub}} H$		115.6 ± 2.8	298	C	[2004MAT/MIR]
	$\Delta_{\text{sub}} H$		82.7 ± 2.8	298	C	[2003MAT/MIR2]
Note: in [2004MAT/MIR] the authors state that an error was made in [2003MAT/MIR2] in converting the enthalpy of sublimation measured at a higher temperature back to 298 K.						
C₈H₅NO₃	[2037-95-8]	2 <i>H</i> -1,3-benzoxazine-2,4(3 <i>H</i>)-dione				
	$\Delta_{\text{fus}} H$		28.63	500.5	DSC	[2004MAT/MIR]
	$\Delta_{\text{sub}} H$		114.2 ± 2.7	298	C	[2004MAT/MIR]
C₈H₅N₃	[27032-01-5]	pyridinium dicyanomethylide				
	$\Delta_{\text{sub}} H$	(403–433)	125.4	418	A	[1987STE/MAL]
	$\Delta_{\text{sub}} H$	(403–406)	125.5 ± 1.3		ME	[1967BOY/GUH, 1970COX/PIL]
C₈H₆	[536-74-3]	phenylacetylene				
	$\Delta_{\text{fus}} H$		9.46	228		[1982LEB/BYK]
	$\Delta_v H$	(313–416)	42.6 ± 0.1	320	EB	[2002STE/CHI3]
	$\Delta_v H$	(313–416)	40.4 ± 0.1	360	EB	[2002STE/CHI3]
	$\Delta_v H$	(313–416)	38.0 ± 0.2	400	EB	[2002STE/CHI3]
	$\Delta_v H$	(265–291)	43.9	278	MM	[1981CHI/HYM]
	$\Delta_v H$	(270–292)	45.2	281	HSA	[1981CHI/HYM]
C₈H₆BrN	[5798-79-8]	<i>(dl)</i> α-bromophenylacetonitrile				
	$\Delta_v H$	(293–515)	64.7	308	A	[1987STE/MAL]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₈ H ₆ ClN	[612-13-5] $\Delta_{\text{fus}}H$	2-(chloromethyl)benzotrile (300–372)	20.53	344.2	AC	[2007XU/ZEN]
C ₈ H ₆ ClNO ₃	[22751-23-1] $\Delta_{\text{sub}}H$	2-nitrobenzeneacetyl chloride (296–327)	103.6	311	TE	[1987STE/MAL, 1947BAL, 1960JON]
C ₈ H ₆ ClNO ₃	[99-47-8] $\Delta_{\text{sub}}H$	3-nitrobenzeneacetyl chloride (299–343)	109.1	314	TE	[1987STE/MAL, 1947BAL, 1960JON]
C ₈ H ₆ ClNS ₂	[28908-00-1] $\Delta_{\text{fus}}H$	2-(chloromethylthio)benzothiazole (80–350)	17.02	351.1	AC	[2005WAN/TAN2]
C ₈ H ₆ Cl ₂	[2123-28-6] Δ_vH Δ_vH	2,3-dichlorostyrene (334–508) (334–508)	55.4 54.3	349 349	 A	 [1999DYK/SVO] [1987STE/MAL, 1947STU]
C ₈ H ₆ Cl ₂	[2123-27-5] Δ_vH	2,4-dichlorostyrene (327–498)	55.0	342	A	[1987STE/MAL, 1970DYK/VAN]
C ₈ H ₆ Cl ₂	[1123-84-8] Δ_vH	2,5-dichlorostyrene (328–500)	54.3	343	A	[1987STE/MAL, 1947STU, 1970DYK/VAN]
C ₈ H ₆ Cl ₂	[28469-92-3] Δ_vH Δ_vH	2,6-dichlorostyrene (321–490)	53.8 ± 1.5 50.4	298 336	GS A	[2001PUR/CHI] [1987STE/MAL, 1947STU, 1970DYK/VAN]
C ₈ H ₆ Cl ₂	[2039-83-0] Δ_vH	3,4-dichlorostyrene (330–503)	53.3	345	A	[1987STE/MAL, 1947STU, 1970DYK/VAN]
C ₈ H ₆ Cl ₂	[2155-42-2] Δ_vH	3,5-dichlorostyrene (326–498)	55.1	341	A	[1987STE/MAL, 1970DYK/VAN]
C ₈ H ₆ Cl ₂ O	[63024-77-1] Δ_vH	3-(chloromethyl)benzoyl chloride (424–464)	54.7	439	A	[1987STE/MAL, 1999DYK/SVO]
C ₈ H ₆ Cl ₂ O	[876-08-4] Δ_vH	4-(chloromethyl)benzoyl chloride (440–466)	68.3	453	A	[1987STE/MAL, 1999DYK/SVO]
C ₈ H ₆ Cl ₂ O ₃	[94-75-7] $\Delta_{\text{fus}}H$ $\Delta_{\text{fus}}H$ $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$	(2,4-dichlorophenoxy)acetic acid (346–387) (346–387)	32 35.33 115 ± 6 123 ± 2 125 ± 3	416.2 412.5 298 361 298	DSC DSC DSC TE TE	[2005VEC/BRU] [1990DON/DRE] [2005VEC/BRU] [2005VEC/BRU] [2005VEC/BRU]
C ₈ H ₆ Cl ₂ O ₃	[1918-00-9] $\Delta_{\text{fus}}H$	3,6-dichloro-2-methoxybenzoic acid	22.9	386.7	DSC	[1991ACR, 1990DON/DRE]
C ₈ H ₆ Cl ₂ O ₄	[7600-50-2] $\Delta_{\text{fus}}H$	3,6-dichloro-5-hydroxy-2-methoxybenzoic acid	28.98	409.8	DSC	[1991ACR, 1990DON/DRE]
C ₈ H ₆ Cl ₄	[877-08-7] Δ_vH	2,3,4,6-tetrachloro-1-ethylbenzene (350–543)	53.6	365	A	[1987STE/MAL, 1947STU]
C ₈ H ₆ Cl ₄	[877-08-7] $\Delta_{\text{fus}}H$ Δ_vH	3,4,5,6-tetrachloro-1,2-dimethylbenzene (367–547)	21.46 63.6	359.2 382	 A	 [1991ACR] [1987STE/MAL, 1970DYK/VAN]
C ₈ H ₆ Cl ₄	[877-10-1] $\Delta_{\text{fus}}H$	tetrachloro- <i>p</i> -xylene	22.59	368.2		[1991ACR]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₈ H ₆ F ₁₂ O ₃ S	[53602-64-5] $\Delta_v H$	bis(1,1,1,3,3,3-hexafluoro-2-methyl-2-propanol) sulfite	46.5			[1975DEM/KOV]
C ₈ H ₆ F ₁₂ O ₄	[485399-48-2] $\Delta_{\text{fus}} H$ $\Delta_{\text{fus}} H$ $\Delta_v H$	3,3,4,4,6,6,7,7,9,9,10,10-dodecafluoro-2,5,8,11-tetraoxododecane	22.07 1.14 36.7	239 251		DSC [2005MAR/AVA] [2005MAR/AVA]
C ₈ H ₆ N ₂	[253-52-1] $\Delta_{\text{fus}} H$ $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$	phthalazine	13.32 82.3 ± 2.3 81.1 ± 0.4 96.7	364.5 298 298		[1993SAB/PEM] C [1995RIB/MAT4] C [1998SAB/TAB, 1993SAB/PEM] ME [1972MIL]
C ₈ H ₆ N ₂	[91-19-0] $\Delta_{\text{fus}} H$ $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$	quinoxaline	11.8 66.6 ± 2.0 69.4 ± 0.6	305.7 298 298		[1993SAB/PEM] C [1995RIB/MAT4] C [1993SAB/PEM]
C ₈ H ₆ N ₂	[253-82-7] $\Delta_{\text{fus}} H$ $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$ $\Delta_v H$	quinazoline	16.95 77.6 ± 0.5 76.6 ± 1.4 58.9 ± 2.0	320.9 298 298		[1993SAB/PEM] C [1998SAB/TAB, 1993SAB/PEM] C [1995RIB/MAT4] CGC [2009LIP/HAN]
C ₈ H ₆ N ₂ O	[1196-57-2] $\Delta_{\text{fus}} H$ $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$	2-hydroxyquinoxaline	32.5 116.1 ± 0.6 118.5 ± 3.1 125.8 ± 4.0	542.5 391 298 298		DSC [2000MON/HIL2] ME [2000MON/HIL2] ME [2000MON/HIL2] C [2000RIB/MAT]
C ₈ H ₆ N ₂ O	[119-39-1] $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$	1-(2H)-phthalazinone	(348–368) 104.4 ± 0.5 (348–368) 107.4 ± 0.5	358 298		ME [2008RIB/CAB2] ME [2008RIB/CAB2]
C ₈ H ₆ N ₂ OS ₂	[2439-01-2] $\Delta_{\text{fus}} H$	6-methyl-1,3-dithiolo[4,5-b]quinoxalin-2-one	29.92	443.2		DSC [1990DON/DRE]
C ₈ H ₆ N ₂ O ₂	[15804-19-0] $\Delta_{\text{sub}} H$	2,3-dihydroxyquinoxaline	156.3 ± 5.5	298		C [2000RIB/MAT]
C ₈ H ₆ N ₂ O ₂	[2433-66-7] $\Delta_{\text{sub}} H$	quinoxaline-1,4-dioxide	112.0 ± 1.9	298		C [1997ACR/POW]
C ₈ H ₆ N ₂ O ₂	[2518-24-3] $\Delta_{\text{sub}} H$	3-aminophthalimide	(386–459) 108.3	401		A [1987STE/MAL, 1956KLO]
C ₈ H ₆ N ₂ O ₂	[3676-85-5] $\Delta_{\text{sub}} H$	4-aminophthalimide	(444–498) 135.3	459		A [1987STE/MAL]
C ₈ H ₆ N ₂ O ₂	[6146-52-7] $\Delta_{\text{sub}} H$	5-nitroindole	(353–375) 110.5 ± 1.0	298		ME [2009RIB/CAB]
C ₈ H ₆ N ₄	[na] $\Delta_{\text{sub}} H$	monobenzo-1,3 α ,4,6 α -tetraazapentalene	(323–373) 74.9 ± 2.9	348		[1967CHI/SIM]
C ₈ H ₆ N ₄	[na]	monobenzo-1,3 α ,6,6 α -tetraazapentalene				

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{sub}}H$	(323–383)	63.6 ± 2.9	350		[1967CHI/SIM]
C ₈ H ₆ O	[271-89-6]	2,3-benzofuran				
	Δ_vH	(323–403)	46.2	338	A	[1987STE/MAL]
C ₈ H ₆ O	[5101-44-0]	2-ethynylphenol				
	Δ_vH	(300–373)	33.5	315	A	[1987STE/MAL]
C ₈ H ₆ O ₂	[1074-12-0]	phenyl glyoxal				
	Δ_vH	(348–467)	59.7	363	A	[1987STE/MAL, 1947STU]
C ₈ H ₆ O ₂	[87-41-2]	phthalide				
	Δ_vH	(368–563)	59.3	383	A	[1987STE/MAL, 1947STU]
C ₈ H ₆ O ₃	[120-57-0]	piperonal				
	$\Delta_{\text{sub}}H$	(293–353)	90.8	323		[1953SER/VOI, 1960JON, 1987STE/MAL]
	Δ_vH	(310–353)	65.7	331		[1953SER/VOI]
	Δ_vH	(360–536)	60.6	375	A	[1987STE/MAL, 1947STU]
C ₈ H ₆ O ₃	[619-66-9]	4-formylbenzoic acid				
	$\Delta_{\text{fus}}H$		21.3	452.2		[2004CHE/MA]
C ₈ H ₆ O ₄	[88-99-3]	phthalic acid				
	$\Delta_{\text{fus}}H$		36.5	463.5		[1999SAB/PER2]
	$\Delta_{\text{sub}}H$		129.8 ± 0.6	298	C	[1999SAB/PER2]
C ₈ H ₆ O ₄	[121-91-5]	isophthalic acid				
	$\Delta_{\text{fus}}H$		43.2	617.4	DTA	[1999SAB/PER2]
	$\Delta_{\text{sub}}H$	(424–476)	134.6 ± 1.6	450	ME	[2000KOZ/MAK]
	$\Delta_{\text{sub}}H$		142.0 ± 0.7	298	C	[1999SAB/PER2]
	$\Delta_{\text{sub}}H$	(493–563)	114.2	508	A	[1987STE/MAL, 1962KRA/BER]
	$\Delta_{\text{sub}}H$	(493–563)	106.7 ± 2.2	523	GS	[1962KRA/BER, 1970COX/PIL]
C ₈ H ₆ O ₄	[100-21-0]	terephthalic acid				
	$\Delta_{\text{sub}}H$	(442–500)	142.2 ± 1.5	471	ME	[2000KOZ/MAK]
	$\Delta_{\text{sub}}H$		146.6 ± 0.5	298	C	[1999SAB/PER2]
	$\Delta_{\text{sub}}H$	(568–675)	139.3 ± 3.8		DTA	[1968LUC/LEW]
	$\Delta_{\text{sub}}H$	(523–633)	139.2	538	A	[1987STE/MAL, 1962KRA/BER]
	$\Delta_{\text{sub}}H$	(523–633)	131	573	GS	[1962KRA/BER]
	$\Delta_{\text{sub}}H$	(392–425)	U 98.24 ± 2.5	408		[1934HIR, 1970COX/PIL]
C ₈ H ₆ O ₄	[94-53-1]	1,3-benzodioxole-5-carboxylic acid (piperonylic acid)				
	$\Delta_{\text{fus}}H$		30.5	501.6		[2004MAT/MON]
	$\Delta_{\text{sub}}H$	(363–377)	113.6 ± 1.1	370	ME	[2004MAT/MON]
	$\Delta_{\text{sub}}H$	(363–377)	117.2 ± 1.8	298	ME	[2004MAT/MON]
C ₈ H ₆ S	[95-15-8]	benzo[b]thiophene				
	$\Delta_{\text{fus}}H$		11.84	304.5		[1996DOM/HEA]
	$\Delta_{\text{sub}}H$		65.7 ± 0.2	298	C	[1979SAB]
	Δ_vH	(349–424)	52.1	364		[1999DYK/SVO]
	Δ_vH	(424–498)	47.9	439		[1999DYK/SVO]
	Δ_vH	(498–631)	45.0	513		[1999DYK/SVO]
	Δ_vH	(310–542)	54.3	320		[1991CHI/KN12]
	Δ_vH	(310–542)	52.0	360		[1991CHI/KN12]
	Δ_vH	(310–542)	49.7	400		[1991CHI/KN12]
	Δ_vH	(310–542)	46.2	460		[1991CHI/KN12]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(310–542)	43.8	500		[1991CHI/KNI2]
	$\Delta_v H$	(310–542)	41.2	540		[1991CHI/KNI2]
	$\Delta_v H$		47.2	425		[1981WIE/KOB]
	$\Delta_v H$		42.8	505		[1981WIE/KOB]
	$\Delta_v H$		36.1	605		[1981WIE/KOB]
	$\Delta_v H$	(306–346)	53.8	326		[1981EDW/PRA, 1999DYK/SVO]
C₈H₆S₂	[492-97-7]	2,2'-bithiophene				
	$\Delta_{\text{fus}}H$		16.5	304.2		[2006TEM/ROU]
	$\Delta_{\text{sub}}H$	(275–291)	86.0 ± 0.4	283	ME	[2009RIB/SAN]
	$\Delta_{\text{sub}}H$	(275–291)	85.2 ± 0.4	298	ME	[2009RIB/SAN]
C₈H₆S₂	[3172-56-3]	3,3'-biothiophene				
	$\Delta_{\text{sub}}H$	(301–317)	88.6 ± 0.3	309	ME	[2009RIB/SAN]
	$\Delta_{\text{sub}}H$	(301–317)	89.2 ± 0.3	298	ME	[2009RIB/SAN]
C₈H₇Br	[2039-88-5]	2-bromostyrene				
	$\Delta_v H$	(378–543)	48.7	393	A	[1987STE/MAL, 1970DYK/VAN]
C₈H₇Br	[2039-82-9]	4-bromostyrene				
	$\Delta_v H$	(393–420)	48.5	406		[1999DYK/SVO]
	$\Delta_v H$	(383–543)	49.9	398	A	[1987STE/MAL, 1970DYK/VAN]
C₈H₇Cl	[2039-87-4]	2-chlorostyrene				
	$\Delta_v H$	(363–523)	46.0	378	A	[1987STE/MAL, 1970DYK/VAN]
C₈H₇Cl	[2039-85-2]	3-chlorostyrene				
	$\Delta_v H$	(298–463)	46.1	313	A	[1987STE/MAL, 1947STU, 1970DYK/VAN]
C₈H₇Cl	[1073-67-2]	4-chlorostyrene				
	$\Delta_v H$	(363–523)	48.1	378	A	[1987STE/MAL, 1947STU, 1970DYK/VAN]
C₈H₇ClN₂O₂	[14722-82-8]	N-(2-chlorophenyl)-2-(hydroxyimino)acetamide				
	$\Delta_{\text{fus}}H$		29.7	432.7	DTA	[1982CUE/SOL]
C₈H₇ClN₂O₂S	[364-98-7]	7-chloro-3-methyl-2H-1,2,4-benzothiadiazine 1,1-dioxide (diazoxide)				
	$\Delta_{\text{fus}}H$		34.1	600.4	DSC	[2006WAS/HOL]
C₈H₇ClO	[532-27-4]	2-chloroacetophenone				
	$\Delta_{\text{sub}}H$	(278–323)	90.7	293	TE	[1987STE/MAL, 1947BAL, 1960JON]
C₈H₇ClO	[99-91-2]	4'-chloroacetophenone				
	$\Delta_v H$	(404–623)	54.0	419	A	[1987STE/MAL, 1999DYK/SVO]
	$\Delta_v H$	(395–485)	50.7	410		[1949DRE/SHR, 1949DRE/MAR, 1984BOU/FRI]
C₈H₇ClO	[103-80-0]	phenylacetyl chloride				
	$\Delta_v H$	(321–483)	56.5	336	A	[1987STE/MAL, 1947STU, 1999DYK/SVO]
C₈H₇ClO₂	[501-53-1]	benzyl chloroformate				
	$\Delta_v H$	(293–303)	38.5 ± 0.1	298		[1990DAV/FIN]
C₈H₇ClO₂	[2444-36-2]	(2-chlorophenyl)acetic acid				
	$\Delta_{\text{fus}}H$		24.33	367.4	DSC	[2008RIB/FER3]
C₈H₇ClO₂	[1878-65-5]	(3-chlorophenyl)acetic acid				
	$\Delta_{\text{fus}}H$		22.6	349.8	DSC	[2008RIB/FER3]
C₈H₇ClO₂	[1878-66-6]	(4-chlorophenyl)acetic acid				
	$\Delta_{\text{fus}}H$		23.57	377.9	DSC	[2008RIB/FER3]
C₈H₇ClO₃	[na]	<i>(d)</i> <i>o</i> -chloromandelic acid				

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{fus}}H$		24.69	392.5		[1991CHI/BRA]
C ₈ H ₇ ClO ₃	[10421-85-9]	(<i>dl</i>) <i>o</i> -chloromandelic acid				
	$\Delta_{\text{fus}}H$		20.08	358.5		[1991CHI/BRA]
C ₈ H ₇ ClO ₃	[na]	2-chloromandelic acid				
	$\Delta_{\text{fus}}H$		23.1	363.5	DSC	[2009HE/ZHU]
C ₈ H ₇ ClO ₃	[na]	(<i>R</i>) 2-chloromandelic acid				
	$\Delta_{\text{fus}}H$		24.9	391.3	DSC	[2009HE/ZHU]
C ₈ H ₇ ClO ₃	[492-86-4]	(<i>dl</i>) <i>p</i> -chloromandelic acid				
	$\Delta_{\text{fus}}H$		27.2	394		[1991CHI/BRA]
C ₈ H ₇ ClO ₃	[na]	(<i>d</i>) <i>p</i> -chloromandelic acid				
	$\Delta_{\text{fus}}H$		23.01	394		[1991CHI/BRA]
C ₈ H ₇ ClO ₃	[122-88-3]	4-chlorophenoxyacetic acid				
	$\Delta_{\text{fus}}H$		36.27	429.6	DSC	[1990DON/DRE]
C ₈ H ₇ ClO ₃	[19463-48-0]	5-chloro-4-hydroxy-3-methoxybenzaldehyde				
	$\Delta_{\text{fus}}H$		39.4	442.6	DSC	[2000LAR/LER]
C ₈ H ₇ Cl ₂ NO	[1918-18-9]	methyl-3,4-dichlorophenylcarbamate				
	$\Delta_{\text{fus}}H$		23.19	381.4	DSC	[1990DON/DRE]
C ₈ H ₇ Cl ₃ O ₃	[2539-26-6]	3,4,5-trichloro-2,6-dimethoxyphenol				
	Δ_vH	(293–323)	77.4	308	CGC	[1999LEI/WAN2]
C ₈ H ₇ FO	[450-95-3]	2-fluoroacetophenone				
	Δ_vH	(273–333)	62	288	A,GS	[1987STE/MAL, 1948RED/CHA4, 1999DYK/SVO]
C ₈ H ₇ FO ₃	[395-05-1]	(<i>dl</i>) <i>m</i> -fluoromandelic acid				
	$\Delta_{\text{fus}}H$		24.69	370		[1991CHI/BRA, 1994LAR/MAR]
C ₈ H ₇ FO ₃	[na]	(<i>d</i>) <i>m</i> -fluoromandelic acid				
	$\Delta_{\text{fus}}H$		24.27	394		[1991CHI/BRA]
C ₈ H ₇ FO ₃	[389-31-1]	(<i>dl</i>) <i>o</i> -fluoromandelic acid				
	$\Delta_{\text{fus}}H$		30.12	390		[1991CHI/BRA, 1994LAR/MAR]
C ₈ H ₇ FO ₃	[na]	(<i>d</i>) <i>o</i> -fluoromandelic acid				
	$\Delta_{\text{fus}}H$		20.92	363		[1991CHI/BRA]
C ₈ H ₇ FO ₃	[395-33-5]	(<i>dl</i>) <i>p</i> -fluoromandelic acid				
	$\Delta_{\text{fus}}H$		29.29	403		[1991CHI/BRA, 1994LAR/MAR]
C ₈ H ₇ FO ₃	[na]	(<i>d</i>) <i>p</i> -fluoromandelic acid				
	$\Delta_{\text{fus}}H$		30.54	426		[1991CHI/BRA]
C ₈ H ₇ F ₃	[21249-93-4]	1,1,1-trifluoro-2-phenylethane				
	Δ_vH	(273–313)	46.1 ± 0.3	298	GS	[1997SCH/VER]
C ₈ H ₇ N	[140-29-4]	benzylcyanide				
	Δ_vH	(283–328)	60.1 ± 0.7	306	GS	[2000VER]
	Δ_vH	(283–328)	60.5 ± 0.7	298	GS	[2000VER]
	Δ_vH	(333–507)	54.8	348	A	[1987STE/MAL, 1947STU]
C ₈ H ₇ N	[620-22-4]	2-tolunitrile				
	Δ_vH	(309–479)	50.8	324	A	[1987STE/MAL, 1947STU]
C ₈ H ₇ N	[104-85-8]	4-tolunitrile				
	Δ_vH	(315–491)	48.0	330	A	[1987STE/MAL, 1947STU]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound					
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference	
C ₈ H ₇ N	[10468-64-1]	2-tolylisocyanide					
	$\Delta_v H$	(298–457)	48.5	313	A	[1987STE/MAL, 1947STU]	
C ₈ H ₇ N	[120-72-9]	indole					
	$\Delta_{\text{sub}}H$	(275–291)	78.4 ± 1.4	283	ME	[2008RIB/CAB3]	
	$\Delta_{\text{sub}}H$	(275–291)	77.6 ± 1.1	298	ME	[2008RIB/CAB3]	
	$\Delta_{\text{sub}}H$	(291–319)	75.0	305	A	[1987STE/MAL]	
	$\Delta_{\text{sub}}H$	(275–303)	77.8 ± 1.6	289	ME	[1974ARS]	
	$\Delta_{\text{sub}}H$	(283–301)	70.0	292		[1955AIH2]	
	$\Delta_{\text{sub}}H$	(283–328)	74.9	305		[1954SER/VOI, 1960JON]	
$\Delta_v H$		57.3		GC	[1996GOV/RUT]		
C ₈ H ₇ NO	[3173-56-6]	benzyl isocyanate					
	$\Delta_v H$	(333–393)	42.3	348	A	[1987STE/MAL]	
C ₈ H ₇ NO ₂	[5466-88-6]	2 <i>H</i> -1,4-benzoxazin-3(4 <i>H</i>)-one					
	$\Delta_{\text{fus}}H$		22.8	445.6	DSC	[2006MAT/MIR]	
			$\Delta_{\text{sub}}H$	106.4 ± 3.0	298	C	[2006MAT/MIR]
C ₈ H ₇ NO ₃	[577-59-3]	2'-nitroacetophenone					
	$\Delta_v H$	(293–333)	103.6	308	A	[1987STE/MAL]	
C ₈ H ₇ NO ₃	[121-89-1]	3'-nitroacetophenone					
	$\Delta_{\text{sub}}H$	(293–343)	110	308	A	[1987STE/MAL]	
C ₈ H ₇ NO ₄	[16498-20-7]	2,3-dihydro-6-nitro-1,4-benzodioxin					
	$\Delta_{\text{fus}}H$		24.27	394	DSC	[2008MAT/SOU2]	
			$\Delta_{\text{sub}}H$	100.6 ± 1.2	298	C	[2008MAT/SOU2]
C ₈ H ₇ NO ₄	[606-27-9]	2-nitrobenzoic acid, methyl ester					
	$\Delta_v H$	(423–453)	56.1	438	A	[1987STE/MAL]	
C ₈ H ₇ NO ₄	[610-69-5]	(2-nitrophenyl) acetate					
	$\Delta_v H$	(373–526)	71.1	388	A	[1987STE/MAL, 1947STU]	
C ₈ H ₇ NO ₄	[1975-50-4]	2-methyl-3-nitrobenzoic acid					
	$\Delta_{\text{sub}}H$		119.5 ± 2.3	298	ME	[2001MON/HIL3]	
C ₈ H ₇ NO ₄	[13506-76-8]	2-methyl-6-nitrobenzoic acid					
	$\Delta_{\text{sub}}H$		120.0 ± 2.2	298	ME	[2001MON/HIL3]	
C ₈ H ₇ NO ₄	[5437-38-7]	3-methyl-2-nitrobenzoic acid					
	$\Delta_{\text{sub}}H$		124.4 ± 2.7	298	ME	[2001MON/HIL3]	
C ₈ H ₇ NO ₄	[3113-71-1]	3-methyl-4-nitrobenzoic acid					
	$\Delta_{\text{sub}}H$		119.3 ± 2.5	298	ME	[2001MON/HIL3]	
C ₈ H ₇ NO ₄	[96-98-0]	4-methyl-3-nitrobenzoic acid					
	$\Delta_{\text{sub}}H$		118.6 ± 2.5	298	ME	[2001MON/HIL3]	
C ₈ H ₇ NO ₄	[3113-72-2]	2-methyl-3-nitrobenzoic acid					
	$\Delta_{\text{sub}}H$		118.7 ± 2.2	298	ME	[2001MON/HIL3]	
C ₈ H ₇ NO ₅	[4920-80-3]	3-methoxy-2-nitrobenzoic acid					
	$\Delta_{\text{sub}}H$	(398–410)	136.6 ± 1.3	404	ME	[1999RIB/MAT]	
			$\Delta_{\text{sub}}H$	141.9 ± 1.3	298	ME	[1999RIB/MAT]
C ₈ H ₇ NO ₅	[89-41-8]	4-methoxy-3-nitrobenzoic acid					
	$\Delta_{\text{sub}}H$	(387–401)	126.5 ± 0.8	394	ME	[1999RIB/MAT]	
			$\Delta_{\text{sub}}H$	131.2 ± 0.8	298	ME	[1999RIB/MAT]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₈ H ₇ NO ₅	[5081-36-7]	3-methoxy-4-nitrobenzoic acid				
	$\Delta_{\text{sub}}H$	(388–402)	126.1 ± 1.1	395	ME	[1999RIB/MAT]
	$\Delta_{\text{sub}}H$		131.0 ± 1.1	298	ME	[1999RIB/MAT]
C ₈ H ₇ NS	[622-78-6]	benzyl isothiocyanate				
	Δ_vH	(352–516)	62.2	367	A	[1987STE/MAL, 1947STU]
C ₈ H ₇ NS	[120-75-2]	2-methylbenzothiazole				
	Δ_vH	(343–499)	61.3	358	A	[1987STE/MAL, 1947STU, 1999DYK/SVO]
C ₈ H ₇ N ₃ O ₂	[1660-15-7]	3,6-diaminophthalimide				
	$\Delta_{\text{sub}}H$	(461–508)	98.5	476	A	[1987STE/MAL, 1956KLO]
C ₈ H ₇ N ₃ O ₆	[38677-56-4]	2,2,2-trinitro-1-phenylethane				
	$\Delta_{\text{sub}}H$	(293–308)	84.1 ± 0.4	301	ME	[1972PEP/MAT, 1977PED/RYL, 1987STE/MAL]
C ₈ H ₇ N ₃ O ₆	[632-92-8]	3-methyl-2,4,6-trinitrotoluene				
	$\Delta_{\text{fus}}H$		38.49	455.4		[1919BEL/SAW]
	$\Delta_{\text{sub}}H$		122.6		DSC	[1990HWA/YOS]
	$\Delta_{\text{sub}}H$	(319–411)	129.8 ± 1.1	365	ME	[1987STE/MAL, 1978CUN/PAL]
	Δ_vH		87.9		DSC	[1990HWA/YOS]
C ₈ H ₇ N ₃ O ₇	[4732-14-3]	2,4,6-trinitrophenetole				
	$\Delta_{\text{sub}}H$	(352–364)	79.0	358	A	[1987STE/MAL]
	$\Delta_{\text{sub}}H$		120.5 ± 2.1			[1950NIT/SEK3, 1970COX/PIL]
	Δ_vH	(342–351)	120.5	346	A	[1987STE/MAL]
C ₈ H ₇ N ₅ O ₈	[43072-20-4]	2,4,6,-trinitro-N-(nitromethyl)- <i>m</i> -toluidine				
	$\Delta_{\text{fus}}H$		19.33	375.6	DSC	[1996DOM/HEA, 1973KRI/LIC]
C ₈ H ₇ N ₅ O ₈	[6052-13-7]	2,4,6-N-tetranitro-N-ethylaniline				
	$\Delta_{\text{fus}}H$		23.51	369	DSC	[1996DOM/HEA, 1973KRI/LIC]
C ₈ H ₈	[277-10-1]	cubane				
	$\Delta_{\text{us}}H$		5.94	394		
	$\Delta_{\text{fus}}H$		8.7	404.9	AC	[1992WHI/WAS]
	$\Delta_{\text{sub}}H$		55.2 ± 2.0	298	AC+CGC	[2004BAS/CHI]
	$\Delta_{\text{sub}}H$	(239–262)	80.3 ± 1.6	298	ME	[1966KYB/CAR, 1970COX/PIL, 1987STE/MAL, 2003DIK/FRE]
		Note: in Ref. [2003DIK/FRE] the authors state that the value of 80.3 kJ/mole from [1966KYB/CAR] pertains to the average temperature and not to 298 K. The authors give a value of 79.1 ± 1.7 for the 298 K value				
	Δ_vH		44.6 ± 0.8	298	CGC	[2004BAS/CHI]
C ₈ H ₈	[629-20-9]	cyclooctatetraene				
	$\Delta_{\text{fus}}H$		11.25	268.5		[1996DOM/HEA]
	$\Delta_{\text{sub}}H$		54.4		B	[1949SCO/GRO]
	Δ_vH	(273–348)	43.9	288	A	[1987STE/MAL]
	Δ_vH		43.1	298		[1949SCO/GRO]
C ₈ H ₈	[500-24-3]	bicyclo[2.2.2]octa-2,5,7-triene				
	Δ_vH		42.9 ± 0.1	298	C	[1985KUS]
C ₈ H ₈	[16607-77-5]	1,5,7-octatriene-3-yene				
	Δ_vH	(313–429)	35.1	328	A	[1987STE/MAL]
C ₈ H ₈	[100-42-5]	styrene				

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{fus}}H$		10.96	242.3		[1996DOM/HEA]
	Δ_vH	(245–334)	42.5	260	A	[1987STE/MAL]
	Δ_vH	(334–419)	41.5	349	A	[1987STE/MAL]
	Δ_vH	(306–333)	42.5	319		[1959CHA/VAN]
	Δ_vH	(303–417)	43.1	318		[1955DRE/MAR]
	Δ_vH	(285–333)	43.9	298		[1946PIT/GUT]
	Δ_vH		43.5 ± 0.4	298		[1946PIT/GUT]
	Δ_vH	(306–389)	40.2	348		[1942BUR]
	Δ_vH	(245–357)	43.2	298		[1939PAT/SCH]
C₈H₈	[116316-76-8]	1-cyclopropyl-1,3-pentadiene				
	Δ_vH		51.9 ± 0.1	298	C	[2007PAS/KUZ]
C₈H₈BrCl₂O₃PS	[2104-96-3]	O-(4-bromo-2,5-dichlorophenyl) O,O-dimethylphosphorothioate				
	$\Delta_{\text{fus}}H$		31.15	325.3	DSC	[1990DON/DRE]
C₈H₈BrNO	[103-88-8]	4-bromoacetanilide				
	$\Delta_{\text{fus}}H$		26.0	440.3		[2009BAR/ESP]
	$\Delta_{\text{fus}}H$		25.8	441.2		[2004VEC/CAT]
	$\Delta_{\text{sub}}H$		110 ± 4	298	Fus + Vap	[2009VEC/TOM]
	Δ_vH		78 ± 2	480	TGA	[2009VEC/TOM]
	Δ_vH		77 ± 1	460	TGA	[2009VEC/TOM]
C₈H₈Br₂	[93-52-7]	(1,2-dibromoethyl)benzene				
	Δ_vH	(359–527)	64.9	374	A	[1987STE/MAL, 1947STU, 1970DYK/VAN]
C₈H₈Br₂	[91-13-4]	α,α' -dibromo- <i>o</i> -xylene				
	$\Delta_{\text{fus}}H$		26.78	368.2		[1991ACR]
C₈H₈Br₂	[626-15-3]	α,α' -dibromo- <i>m</i> -xylene				
	$\Delta_{\text{fus}}H$		23.69	350.2		[1991ACR]
C₈H₈ClNO₂	[3942-54-9]	N-methyl-2-chlorophenylcarbamic acid ester				
	$\Delta_{\text{fus}}H$		21.81	362.7	DSC	[1990DON/DRE]
C₈H₈Cl₂	[1124-05-6]	2,5-dichloro-1,4-dimethylbenzene				
	Δ_vH	(393–573)	52.7	408	A	[1987STE/MAL, 1970DYK/VAN, 1999DYK/SVO]
C₈H₈Cl₂	[54484-61-6]	2,5-dichloro-1,4-dimethylbenzene				
	Δ_vH	(319–495)	48.9	334	A	[1987STE/MAL, 1947STU, 1970DYK/VAN]
C₈H₈Cl₂	[54484-63-8]	2,5-dichloro-1,4-dimethylbenzene				
	Δ_vH	(311–490)	46.0	326	A	[1987STE/MAL, 1947STU, 1970DYK/VAN]
C₈H₈Cl₂	[6623-59-2]	3,4-dichloro-1-ethylbenzene				
	Δ_vH	(320–500)	49.3	335	A	[1987STE/MAL, 1947STU, 1970DYK/VAN]
C₈H₈Cl₂	[93-52-7]	1,4-bis(chloromethyl)benzene				
	Δ_vH	(412–504)	50.8	427		[1999DYK/SVO]
C₈H₈Cl₂	[612-12-4]	α,α' -dichloro- <i>o</i> -xylene				
	$\Delta_{\text{fus}}H$		21.26	328.2		[1991ACR]
C₈H₈Cl₂	[626-16-4]	α,α' -dichloro- <i>m</i> -xylene				
	$\Delta_{\text{fus}}H$		19.51	307.2		[1991ACR]
C₈H₈Cl₂	[623-25-6]	α,α' -dichloro- <i>p</i> -xylene				
	$\Delta_{\text{fus}}H$		23.97	373.2		[1991ACR]
C₈H₈Cl₂O₂	[120-67-2]	2-(2,4-dichlorophenoxy)ethanol				

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(484–560)	65.1	499	A	[1959MCD/SHR, 1984BOU/FRI] [1999DYK/SVO, 1987STE/MAL]
C ₈ H ₈ Cl ₂ O ₂	[2675-77-6] $\Delta_{\text{fus}} H$	1,4-dichloro-2,5-dimethoxybenzene	27.56	403.9	DSC	[1991ACR, 1990DON/DRE]
C ₈ H ₈ Cl ₂ O ₃	[na] $\Delta_v H$	3,5-dichloro-2,6-dimethoxyphenol	70.4	308	CGC	[1999LEI/WAN2]
C ₈ H ₈ Cl ₂ O ₃	[6597-78-0] $\Delta_{\text{fus}} H$	methyl 3,6-dichloro-2-methoxybenzoate	18.49	304.6	DSC	[1990DON/DRE]
C ₈ H ₈ Cl ₃ O ₃ PS	[299-84-3] $\Delta_{\text{fus}} H$	O,O-dimethyl-O-(2,4,5-trichlorophenyl)thiophosphate	18.94	313	DSC	[1990DON/DRE]
	$\Delta_v H$	(298–373)	56.8	313	A	[1987STE/MAL]
C ₈ H ₈ NO ₃	[6705-03-9] $\Delta_{\text{fus}} H$	2-amino-5-methoxybenzoic acid	22.88	425	DSC	[2010MON/ALM]
	$\Delta_{\text{sub}} H$	(353–375)	116.9 ± 0.8	364	ME	[2010MON/ALM]
	$\Delta_{\text{sub}} H$	(353–375)	119.3 ± 0.8	298	ME	[2010MON/ALM]
C ₈ H ₈ NO ₃	[2840-26-8] $\Delta_{\text{fus}} H$	3-amino-4-methoxybenzoic acid	25.34	477.9	DSC	[2010MON/ALM]
	$\Delta_{\text{sub}} H$	(380–399)	127.4 ± 0.8	389	ME	[2010MON/ALM]
	$\Delta_{\text{sub}} H$	(380–399)	130.7 ± 0.8	298	ME	[2010MON/ALM]
C ₈ H ₈ NO ₃	[74165-74-5] $\Delta_{\text{fus}} H$	3-amino-5-methoxybenzoic acid	22.4	456.9	DSC	[2010MON/ALM]
	$\Delta_{\text{sub}} H$	(380–400)	132.8 ± 1.0	390	ME	[2010MON/ALM]
	$\Delta_{\text{sub}} H$	(380–400)	136.1 ± 1.0	298	ME	[2010MON/ALM]
C ₈ H ₈ NO ₃	[2486-69-3] $\Delta_{\text{fus}} H$	4-amino-3-methoxybenzoic acid	25.27	462.4	DSC	[2010MON/ALM]
	$\Delta_{\text{sub}} H$	(373–395)	128.9 ± 1.2	384	ME	[2010MON/ALM]
	$\Delta_{\text{sub}} H$	(373–395)	132.0 ± 1.2	298	ME	[2010MON/ALM]
C ₈ H ₈ N ₂	[615-15-6] $\Delta_{\text{us}} H$ $\Delta_{\text{fus}} H$	2-methylbenzimidazole	0.59 20.49	383.9 451.4	DSC	[2002DOM/KOZ]
C ₈ H ₈ N ₂ O ₂	[32692-19-6] $\Delta_{\text{sub}} H$	5-nitroindoline	109.8 ± 0.8	298	ME	[2009RIB/CAB]
C ₈ H ₈ N ₂ O ₂	[122-96-3] $\Delta_v H$	1,4-bis(2-hydroxyethyl)piperazine	67.8 ± 5.3			[1998ABD/MEI]
C ₈ H ₈ N ₂ O ₂	[88-96-0] $\Delta_{\text{sub}} H$	1,2-benzenedicarboxamide	57.3 ± 4.2		ME	[1972HAM/WIT, 1977PED/RYL]
C ₈ H ₈ N ₂ O ₂	[1740-57-4] $\Delta_{\text{sub}} H$	1,3-benzenedicarboxamide	54.4 ± 4.2		ME	[1971HAM/WIT, 1977PED/RYL]
C ₈ H ₈ N ₂ O ₂	[3010-82-0] $\Delta_{\text{sub}} H$	1,4-benzenedicarboxamide	57.3 ± 4.2			[1972HAM/WIT]
C ₈ H ₈ N ₂ O ₂	[1769-41-1] $\Delta_{\text{fus}} H$	2-(hydroxyimino)-N-phenylacetamide	10.4	453.1	DTA	[1982CUE/SOL]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₈ H ₈ N ₂ O ₃	[552-32-9]	2'-nitroacetanilide				
	$\Delta_v H$	(473–593)	44.0	488	A	[1987STE/MAL]
C ₈ H ₈ O	[98-86-2]	acetophenone				
	$\Delta_{\text{fus}} H$		1.36	289.7		[2006SED/MAI]
	$\Delta_{\text{fus}} H$		16.65	292.7		[1911LOU/DUP, 2006SED/MAI]
	$\Delta_v H$	(360–520)	55.4 ± 0.4	298	EB	[1996STE/CHI]
	$\Delta_v H$	(360–520)	52.6 ± 0.4	340	EB	[1996STE/CHI]
	$\Delta_v H$	(360–520)	50.1 ± 0.3	380	EB	[1996STE/CHI]
	$\Delta_v H$	(360–520)	47.5 ± 0.3	420	EB	[1996STE/CHI]
	$\Delta_v H$	(360–520)	45.0 ± 0.4	460	EB	[1996STE/CHI]
	$\Delta_v H$	(360–520)	42.2 ± 0.4	500	EB	[1996STE/CHI]
	$\Delta_v H$	(343–383)	53.4	298	CGC	[1995CHI/HOS]
	$\Delta_v H$	(343–383)	52.7	298	CGC	[1995CHI/HOS]
	$\Delta_v H$	(343–383)	57.9	298	CGC	[1995CHI/HOS]
	$\Delta_v H$	(375–603)	49.7	390	A	[1987STE/MAL]
	$\Delta_v H$	(383–437)	41.9	398	GS,EB	[1965COL/COU]
$\Delta_v H$	(310–476)	51.2	325		[1947STU]	
C ₈ H ₈ O	[496-14-0]	2,5-dihydrobenzo-3,4-furan				
	$\Delta_v H$	(285–510)	53.7 ± 0.4	298	EB	[1996STE/CHI3]
C ₈ H ₈ O	[122-78-1]	phenylacetaldehyde				
	$\Delta_v H$	(293–343)	59.8 ± 0.3	298	GS	[2007EME/DAB]
	$\Delta_v H$	(283–333)	54.5	298	A	[1987STE/MAL]
C ₈ H ₈ O ₂	[493-09-4]	1,4-benzodioxan				
	$\Delta_v H$		67.4 ± 1.7	298	C	[2008MAT/SOU2]
	$\Delta_v H$	(400–486)	50.4	415	A	[1987STE/MAL]
	$\Delta_v H$		50.4			[1958CAS/FLE2]
C ₈ H ₈ O ₂	[104-57-4]	benzyl formate				
	$\Delta_v H$	(298–357)	51.6	313	A	[1987STE/MAL]
C ₈ H ₈ O ₂	[118-93-4]	2'-hydroxyacetophenone				
	$\Delta_{\text{fus}} H$		13.0	278.5	DSC	[2008BER/MIN]
	$\Delta_v H$		58.3 ± 0.3	298	C	[2008BER/MIN]
	$\Delta_v H$	(369–491)	58.3	384	A	[1987STE/MAL]
	$\Delta_v H$		50.2			[1986BAL/GNA]
C ₈ H ₈ O ₂	[121-71-1]	3'-hydroxyacetophenone				
	$\Delta_{\text{fus}} H$		23.4	366.7	DSC	[2005CHE/TAN]
C ₈ H ₈ O ₂	[99-93-4]	4'-hydroxyacetophenone				
	$\Delta_{\text{fus}} H$		18.08	382.8		[2008BER/PIE]
	$\Delta_{\text{fus}} H$		17.0	381.3	DSC	[2005CHE/TAN]
	$\Delta_{\text{sub}} H$	(320–349)	95.7	335	A	[1987STE/MAL, 1960AIH]
C ₈ H ₈ O ₂	[123-11-5]	4-methoxybenzaldehyde				
	$\Delta_v H$	(348–521)	58.4	363	A,EB	[1985SCH/BRU]
	$\Delta_v H$	(283–323)	60.4	298	A	[1987STE/MAL, 1955SER/VOI]
	$\Delta_v H$	(346–521)	57.1	361	A	[1987STE/MAL, 1947STU]
C ₈ H ₈ O ₂	[93-58-3]	methyl benzoate				
	$\Delta_{\text{fus}} H$	(5–320)	14.8	260.8	AC	[2002BLO/PAU]
	$\Delta_{\text{fus}} H$		14.83	260.8		[1998MAK/KAB]
	$\Delta_{\text{fus}} H$		13.9	261		[1978DOZ/FUJ]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(358–517)	51.1 ± 0.2	360	EB	[2002STE/CHI2]
	$\Delta_v H$	(358–517)	48.5 ± 0.2	400	EB	[2002STE/CHI2]
	$\Delta_v H$	(358–517)	45.8 ± 0.2	440	EB	[2002STE/CHI2]
	$\Delta_v H$	(358–517)	43.0 ± 0.4	480	EB	[2002STE/CHI2]
	$\Delta_v H$		57.2 ± 0.1	303	C	[1998MAK/KAB]
	$\Delta_v H$	(313–353)	53.4	298	CGC	[1995CHI/HOS]
	$\Delta_v H$	(313–363)	53.8	298	CGC	[1995CHI/HOS]
	$\Delta_v H$	(433–473)	54.7	298	CGC	[1995CHI/HOS]
	$\Delta_v H$	(334–428)	50.7	379	BG	[1988KAT2]
	$\Delta_v H$	(334–428)	48.3	410	BG	[1988KAT2]
	$\Delta_v H$	(283–323)	53.9	298	A	[1987STE/MAL]
	$\Delta_v H$	(373–533)	49.7	388	A	[1987STE/MAL]
	$\Delta_v H$		55.6 ± 0.1	298	C	[1972COL/LAY]
	$\Delta_v H$	(341–433)	52.8	363	BG	[1971HAL/BAL]
C₈H₈O₂	[118-90-1]	2-methylbenzoic acid				
	$\Delta_{\text{fus}}H$		20.17	376.9		[1991ACR]
	$\Delta_{\text{sub}}H$	(297–337)	95.9 ± 0.1	298	ME	[1986COL/JIM]
	$\Delta_{\text{sub}}H$		137.7 ± 0.5		DSC	[1983HOL]
C₈H₈O₂	[99-04-7]	3-methylbenzoic acid				
	$\Delta_{\text{fus}}H$		15.73	381.9		[1991ACR]
	$\Delta_{\text{sub}}H$	(303–323)	97.0 ± 0.3	298	ME	[1986COL/JIM]
	$\Delta_v H$	(473–533)	62.8	503	A	[1987STE/MAL, 1970MUL/GAL]
C₈H₈O₂	[99-94-5]	4-methylbenzoic acid				
	$\Delta_{\text{fus}}H$		22.72	452.8		[1991ACR]
	$\Delta_{\text{sub}}H$	(320–337)	98.6 ± 0.6	298	ME	[2004MON/ALM]
	$\Delta_{\text{sub}}H$	(318–337)	98.8 ± 0.3	298	ME	[1986COL/JIM]
C₈H₈O₂	[122-79-2]	phenyl acetate				
	$\Delta_v H$	(313–363)	53.3	298	CGC	[1995CHI/HOS]
	$\Delta_v H$	(433–473)	53.6	298	CGC	[1995CHI/HOS]
	$\Delta_v H$	(313–353)	53.1	298	CGC	[1995CHI/HOS]
	$\Delta_v H$	(311–469)	51.7	326	A	[1987STE/MAL, 1947STU]
C₈H₈O₂	[103-82-2]	phenylacetic acid				
	$\Delta_{\text{fus}}H$		32.0	NA	DSC	[2003SHA/KAN]
	$\Delta_{\text{fus}}H$		15.2	349.2		[2002GRA/RAS]
	$\Delta_{\text{fus}}H$		16.5	350.8		[2001MON/HIL]
	$\Delta_{\text{fus}}H$		14.49	349.9	DSC	[1991ACR]
	$\Delta_{\text{sub}}H$	(307–339)	93.5 ± 0.3	298	GS	[2004ROU/TEM]
	$\Delta_{\text{sub}}H$	(305–321)	98.6 ± 0.4	313	ME	[2001MON/HIL]
	$\Delta_{\text{sub}}H$	(305–321)	99.0 ± 0.6	298	ME	[2001MON/HIL]
	$\Delta_v H$	(353–392)	79.1 ± 0.3	298	GS	[2004ROU/TEM]
	$\Delta_v H$	(370–539)	65.0	385	A	[1987STE/MAL]
C₈H₈O₂	[137-18-8]	2,5-dimethyl-1,4-benzoquinone				
	$\Delta_{\text{sub}}H$	(273–293)	77.0	283	QF	[1927COO/COO, 1960JON, 1987STE/MAL]
C₈H₈O₂	[7145-99-5]	5-methyl-1,3-benzodioxole				
	$\Delta_v H$		54.9 ± 1.2	298	C	[2007MAT/SOU]

Note: There is a large discrepancy between the value of 32.0 kJ/mole and the other reported literature values. Reference [2003SHA/KAN] quotes a literature value of 32.0 kJ/mole; however, the authors do not provide the source of the cited literature value.

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₈ H ₈ O ₂ S	[5535-48-8]	phenyl vinyl sulfone				
	$\Delta_{\text{fus}}H$		11.72	343.4		[1969MAC/MCN]
	$\Delta_{\text{sub}}H$		82 ± 2.5		B	[1969MAC/MCN, 1969MAC/MCN2, 1977PED/RYL]
C ₈ H ₈ O ₃	[935-79-5]	<i>cis</i> 4-cyclohexene-1,2-dicarboxylic acid anhydride				
	Δ_vH	(325–525)	53.1 ± 0.1			[1984NUR/MEK]
C ₈ H ₈ O ₃	[na]	1-cyclohexene-1,2-dicarboxylic acid anhydride				
	$\Delta_{\text{fus}}H$	(80–360)	11.88	343.5	AC	[2004LU/TAN]
C ₈ H ₈ O ₃	[25326-19-6]	5,6-dioxycarbonyl[2.2.1]bicyclohept-2-ene				
	$\Delta_{\text{trs}}H$		0.9	323.6		
	$\Delta_{\text{fus}}H$		8.7	342.4		
			3.6	388.4	DSC	[1987CUR/ASR]
C ₈ H ₈ O ₃	[99-76-3]	4-hydroxybenzoic acid, methyl ester				
	$\Delta_{\text{fus}}H$		25.3	399.2		[1999GIO/BET]
	$\Delta_{\text{fus}}H$		24.31	398.5		[1990MAN/AHU]
	$\Delta_{\text{sub}}H$	(303–327)	98.8 ± 0.8	298	GS	[2005PER/ROD]
	$\Delta_{\text{sub}}H$		77.1		TGA	[2002CHA/DOL]
	Δ_vH		83.1	298	CGC	[2005TEM/ROU]
		(446–517)	81.5	461	A	[1987STE/MAL]
C ₈ H ₈ O ₃	[119-36-8]	methyl salicylate				
	Δ_vH	(333–433)	U56.2	298	GC	[2005HOS/GRY]
	Δ_vH		52.3		TG,DTA	[2001CHE/HUA]
	Δ_vH	(327–497)	59.9	342	A	[1987STE/MAL]
	Δ_vH	(329–496)	58.7	344	A	[1987STE/MAL]
	Δ_vH	(288–333)	56.9	303	A	[1987STE/MAL]
C ₈ H ₈ O ₃	[148-53-8]	2-hydroxy-3-methoxybenzaldehyde				
	$\Delta_{\text{sub}}H$	(282–303)	54.1	292.5	A	[1987STE/MAL]
C ₈ H ₈ O ₃	[121-33-5]	4-hydroxy-3-methoxybenzaldehyde (vanillin)				
	$\Delta_{\text{fus}}H$		22.35	NA	DSC	[2008SIN/DAS]
	$\Delta_{\text{fus}}H$		22.4	355.4		[2008TEM/ROU]
	$\Delta_{\text{sub}}H$	(293–353)	88.7	323		[1953SER/VOI, 1960JON]
	Δ_vH	(353–463)	66.9	298	GC	[2005HOS/GRY]
	Δ_vH	(380–558)	66.9	395	A	[1987STE/MAL, 1947STU]
C ₈ H ₈ O ₃	[579-75-9]	2-methoxybenzoic acid				
	$\Delta_{\text{fus}}H$		24.0	374.6	DSC	[2008PER/VOL]
	$\Delta_{\text{sub}}H$	(309–363)	110.7 ± 0.8	298	GS	[2008PER/VOL]
	$\Delta_{\text{sub}}H$	(318–353)	101.2	333	A	[1987STE/MAL]
	$\Delta_{\text{sub}}H$		104.7 ± 0.3	298	ME	[1978COL/JIM]
	$\Delta_{\text{sub}}H$	(353–368)	90.8 ± 0.4	360	GS	[1973MAL/GIG, 1987STE/MAL]
	$\Delta_{\text{sub}}H$	(353–368)	90.9	360	GS	[1954DAV/JON, 1960JON]
	Δ_vH		91.8	298	Sub-Fus	[2008PER/VOL]
C ₈ H ₈ O ₃	[586-38-9]	3-methoxybenzoic acid				
	$\Delta_{\text{fus}}H$		24.9	378.7	DSC	[2008PER/VOL]
	$\Delta_{\text{sub}}H$	(310–349)	114.7 ± 0.8	298	GS	[2008PER/VOL]
	$\Delta_{\text{sub}}H$		107.5 ± .4	298	ME	[1987STE/MAL, 1978COL/JIM]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$		95	298	Sub-Fus	[2008PER/VOL]
C ₈ H ₈ O ₃	[100-09-4]	4-methoxybenzoic acid				
	$\Delta_{\text{fus}} H$		29.9	455.6	DSC	[2008PER/VOL]
	$\Delta_{\text{fus}} H$		28.4	457.8		[1991ACR]
	$\Delta_{\text{sub}} H$	(316–369)	111.6 ± 0.6	298	GS	[2008PER/VOL]
	$\Delta_{\text{sub}} H$		109.8 ± 0.6	298	ME	[1987STE/MAL, 1978COL/JIM]
	$\Delta_v H$		92	298	Sub-Fus	[2008PER/VOL]
C ₈ H ₈ O ₃	[na]	(dl) mandelic acid				
	$\Delta_{\text{fus}} H$		26.1	392.1	MDSC	[2004PRO/RAS]
	$\Delta_{\text{fus}} H$		25.52	392		[1991CHI/BRA]
C ₈ H ₈ O ₃	[na]	(d) mandelic acid				
	$\Delta_{\text{fus}} H$		26.2	404.1	MDSC	[2004PRO/RAS]
	$\Delta_{\text{fus}} H$		26.36	406		[1991CHI/BRA]
C ₈ H ₈ O ₃	[156-38-7]	4-hydroxyphenylacetic acid				
	$\Delta_{\text{fus}} H$		28.0	422.9		[2002GRA/RAS]
	$\Delta_{\text{fus}} H$		28.4	423.6		[1991ACR]
C ₈ H ₈ O ₃	[495-76-1]	1,3-benzodioxole-5-methanol (piperonyl alcohol)				
	$\Delta_{\text{fus}} H$		18.05	327.1		[2004MAT/MON]
	$\Delta_{\text{sub}} H$	(305–319)	103.0 ± 0.6	312	ME	[2004MAT/MON]
	$\Delta_{\text{sub}} H$	(305–319)	103.7 ± 0.7	298	ME	[2004MAT/MON]
C ₈ H ₈ O ₄	[520-45-6]	2-acetyl-5-hydroxy-3-oxo-4-hexenoic acid- δ -lactone (dehydroacetic acid)				
	$\Delta_v H$	(364–542)	62.1	379	A	[1987STE/MAL, 1947STU]
C ₈ H ₈ S	[4565-32-6]	2,3-dihydrobenzo[b]thiophene				
	$\Delta_{\text{fus}} H$		14.84	269.8		[2003STE/CHI]
	$\Delta_v H$	(345–557)	59.1 ± 0.2	298	IP,EB	[2003STE/CHI]
	$\Delta_v H$	(345–557)	56.3 ± 0.2	340	IP,EB	[2003STE/CHI]
	$\Delta_v H$	(345–557)	53.8 ± 0.2	380	IP,EB	[2003STE/CHI]
	$\Delta_v H$	(345–557)	51.4 ± 0.2	420	IP,EB	[2003STE/CHI]
	$\Delta_v H$	(345–557)	49.0 ± 0.2	460	IP,EB	[2003STE/CHI]
	$\Delta_v H$	(345–557)	46.4 ± 0.3	500	IP,EB	[2003STE/CHI]
C ₈ H ₉ Br	[553-94-6]	1-bromo-2,5-dimethylbenzene				
	$\Delta_v H$	(310–480)	50.9	325		[1999DYK/SVO, 1947STU]
	$\Delta_v H$	(310–480)	53.6	325	A	[1987STE/MAL, 1970DYK/VAN]
C ₈ H ₉ Br	[585-71-7]	(1-bromoethyl)benzene				
	$\Delta_v H$	(298–333)	56.4 ± 0.3	298	GS	[2002KRA/VAS]
	$\Delta_v H$		52.4	298	CGC	[2002KRA/VAS]
C ₈ H ₉ Br	[103-63-9]	(2-bromoethyl)benzene				
	$\Delta_v H$	(348–401)	51.5	363	A	[1987STE/MAL, 1999DYK/SVO]
C ₈ H ₉ Br	[1973-22-4]	1-bromo-2-ethylbenzene				
	$\Delta_v H$	(368–523)	48.1	383	A	[1987STE/MAL, 1970DYK/VAN, 1999DYK/SVO]
C ₈ H ₉ Br	[1585-07-5]	1-bromo-4-ethylbenzene				
	$\Delta_v H$	(347–479)	46.2	362		[1999DYK/SVO]
	$\Delta_v H$	(378–533)	49.4	393	A	[1987STE/MAL, 1970DYK/VAN]
	$\Delta_v H$	(303–479)	52.0	318		[1947STU]
C ₈ H ₉ Cl	[627-65-1]	(dl) (1-chloroethyl)benzene				

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(281–319)	52.8 ± 0.2	298	GS	[2002KRA/VAS]
	$\Delta_v H$		52.4	298	CGC	[2002KRA/VAS]
	$\Delta_v H$	(336–372)	51.4	351		[1999DYK/SVO]
	$\Delta_v H$	(342–378)	47.0	357	A	[1987STE/MAL]
C₈H₉Cl	[622-24-2]	(2-chloroethyl)benzene				
	$\Delta_v H$	(356–480)	53.1	368		[1999DYK/SVO]
	$\Delta_v H$	(356–380)	51.7	368	A	[1987STE/MAL]
C₈H₉Cl	[89-96-3]	1-chloro-2-ethylbenzene				
	$\Delta_v H$	(353–503)	46.1	368	A	[1987STE/MAL, 1970DYK/VAN]
	$\Delta_v H$	(290–450)	47.2	305		[1947STU]
C₈H₉Cl	[620-16-6]	1-chloro-3-ethylbenzene				
	$\Delta_v H$	(348–457)	46.4	363		[1999DYK/SVO]
	$\Delta_v H$	(358–508)	46.8	373	A	[1987STE/MAL, 1970DYK/VAN]
	$\Delta_v H$	(291–454)	46.4	307		[1947STU]
C₈H₉Cl	[622-98-0]	1-chloro-4-ethylbenzene				
	$\Delta_v H$	(350–458)	45.8	365		[1999DYK/SVO]
	$\Delta_v H$	(358–508)	46.8	373	A	[1987STE/MAL, 1970DYK/VAN]
	$\Delta_v H$	(381–457)	45.5	396		[1947STU]
C₈H₉Cl	[104-82-5]	1-(chloromethyl)-4-methylbenzene				
	$\Delta_v H$	(376–457)	44.9	391	A	[1987STE/MAL, 1999DYK/SVO]
C₈H₉ClNO₃PS	[500-28-7]	O,O-(dimethyl)-O-(3-chloro-4-nitrophenyl)thiophosphate				
	$\Delta_v H$	(283–409)	92	346	A	[1987STE/MAL, 1999DYK/SVO]
C₈H₉ClNO₃PS	[2463-84-5]	O-(2-chloro-4-nitrophenyl) O,O-dimethyl phosphorothioate				
	$\Delta_{\text{fus}} H$		29.08	323.9	DSC	[1990DON/DRE]
C₈H₉ClN₂O	[52943-21-2]	2-chloro-N,N-dimethylnicotinamide				
	$\Delta_{\text{fus}} H$	(82–380)	21.39	342.2	AC	[2005SUN/LIU3]
C₈H₉ClO	[614-72-2]	1-chloro-2-ethoxybenzene				
	$\Delta_v H$	(318–481)	52.4	333	A	[1987STE/MAL, 1947STU]
C₈H₉ClO	[1875-88-3]	4-chlorophenethyl alcohol				
	$\Delta_v H$	(426–673)	59.3	411	A	[1987STE/MAL, 1947STU]
C₈H₉ClO	[622-61-7]	4-chloro-1-ethoxybenzene				
	$\Delta_v H$	(395–485)	49.5	410	A	[1987STE/MAL, 1947STU]
C₈H₉ClO₂	[7477-64-7]	ethylene glycol, 4-chlorophenyl ether				
	$\Delta_v H$	(410–554)	68.5	425	A	[1987STE/MAL, 1999DYK/SVO]
C₈H₉ClO₃	[na]	3-chloro-2,6-dimethoxyphenol				
	$\Delta_v H$	(293–323)	68.6	308	CGC	[1999LEI/WAN2]
C₈H₉Cl₃O₄	[na]	2-acetyl-4,4,4-trichloro-3-oxobutyric acid, ethyl ester				
	$\Delta_v H$	(374–409)	53.1	389	A	[1987STE/MAL, 1999DYK/SVO]
C₈H₉I	[10604-60-1]	(1-iodoethyl)benzene				
	$\Delta_v H$	(303–340)	59.9 ± 0.4	298	GS	[2002KRA/VAS]
C₈H₉N	[140-76-1]	2-methyl-5-vinylpyridine				
	$\Delta_v H$	(342–457)	55.2	357	A	[1987STE/MAL]
	$\Delta_v H$	(342–457)	54.5	357		[1961FRO/LOG, 1984BOU/FRI]
C₈H₉N	[na]	N-methylbenzaldehyde-imine				
	$\Delta_v H$	(283–318)	51.1 ± 0.2	301	GS	[1997VER/MOR]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(283–318)	51.2 ± 0.2	298	GS	[1997VER/MOR]
C ₈ H ₉ N	[496-15-1]	indoline				
	$\Delta_v H$		61.9 ± 1.7	298	C	[2008RIB/CAB3]
C ₈ H ₉ NO	[103-84-4]	acetanilide (N-phenylacetamide)				
	$\Delta_{\text{fus}} H$		21.4	386.9	DSC	[2005CHE/TAN]
	$\Delta_{\text{fus}} H$		18.3	387.2		[2004VEC/CAT]
	$\Delta_{\text{fus}} H$		21.67	387.5		[1996DOM/HEA]
	$\Delta_{\text{sub}} H$	(303–324)	80.6	313.5	A	[1987STE/MAL, 1955AIH3]
	$\Delta_{\text{sub}} H$	(317–336)	87.2	326.5	A	[1987STE/MAL, 1960AIH2]
	$\Delta_v H$	(473–577)	64.8	488	A	[1987STE/MAL]
	$\Delta_v H$	(387–577)	66.3	402		[1947STU]
C ₈ H ₉ NO	[41977-54-2]	<i>anti</i> 3-methylbenzaloxime				
	$\Delta_{\text{sub}} H$		U 31 ± 1.7		MS	[1983MAJ/AZZ]
C ₈ H ₉ NO	[3235-02-7]	<i>anti</i> 4-methylbenzaloxime				
	$\Delta_{\text{sub}} H$		U 36 ± 1.7		MS	[1983MAJ/AZZ]
C ₈ H ₉ NO	[103-81-1]	2-phenylacetamide				
	$\Delta_{\text{sub}} H$	(329–352)	96.4	340.5	A	[1987STE/MAL, 1960AIH2]
C ₈ H ₉ NO	[99-03-6]	3-aminoacetophenone				
	$\Delta_{\text{fus}} H$		29.0	371.2		[1971LEB/RYA]
C ₈ H ₉ NO	[99-92-3]	4-aminoacetophenone				
	$\Delta_{\text{fus}} H$		19.6	378.2	DSC	[2005CHE/TAN]
	$\Delta_{\text{fus}} H$		38.0	379.2		[1971LEB/RYA]
	$\Delta_{\text{sub}} H$	(314–338)	92.7	326	A	[1987STE/MAL, 1960AIH2]
C ₈ H ₉ NO	[613-93-4]	N-methylbenzamide				
	$\Delta_{\text{sub}} H$	(297–321)	75.0	309	A	[1987STE/MAL, 1955AIH]
	$\Delta_{\text{sub}} H$	(307–329)	85.7	318	A	[1987STE/MAL, 1960AIH2]
C ₈ H ₉ NO ₂	[134-20-3]	anthranilic acid, methyl ester				
	$\Delta_v H$	(299–333)	62.3	314	A,ME	[1987STE/MAL, 1954SER/VOI]
C ₈ H ₉ NO ₂	[81-20-9]	2-nitro-1,3-dimethylbenzene				
	$\Delta_v H$	(284–323)	57.2 ± 0.8	303	GS	[2000VER/HEI]
	$\Delta_v H$		57.5 ± 0.8	298		[2000VER/HEI]
	$\Delta_v H$	(373–498)	49.7	388	A	[1987STE/MAL]
C ₈ H ₉ NO ₂	[89-87-2]	4-nitro-1,3-dimethylbenzene				
	$\Delta_v H$	(368–518)	56.7	383	A	[1987STE/MAL]
	$\Delta_v H$	(338–517)	57.3	353		[1947STU]
C ₈ H ₉ NO ₂	[83-41-0]	1,2-dimethyl-3-nitrobenzene				
	$\Delta_v H$	(383–518)	59.4	398		[1984BOU/FRI]
C ₈ H ₉ NO ₂	[99-51-4]	1,2-dimethyl-4-nitrobenzene				
	$\Delta_v H$	(399–536)	63.6	414		[1984BOU/FRI]
C ₈ H ₉ NO ₂	[612-22-6]	2-nitro-1-ethylbenzene				
	$\Delta_v H$	(284–323)	62.7 ± 0.4	303	GS	[2000VER/HEI]
	$\Delta_v H$		63.0 ± 0.4	298		[2000VER/HEI]
	$\Delta_v H$	(353–422)	56.3	368	A	[1987STE/MAL]
C ₈ H ₉ NO ₂	[100-12-9]	4-nitro-1-ethylbenzene				

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(353–433)	59.4	368	A	[1987STE/MAL]
C ₈ H ₉ NO ₂	[103-01-5]	N-phenylglycine				
	$\Delta_{\text{sub}}H$		114.1 ± 1.0	365.4	C	[1980SAB/SKO]
	$\Delta_{\text{sub}}H$		128.0 ± 2.0	298		[1980SAB/SKO]
C ₈ H ₉ NO ₂	[875-74-1]	<i>(d)</i> α -phenylglycine				
	$\Delta_{\text{sub}}H$		148.9 ± 2.2	443	C	[1980SAB/SKO]
	$\Delta_{\text{sub}}H$		165.0 ± 6.0	298	C	[1980SAB/SKO]
C ₈ H ₉ NO ₂	[29577-53-5]	2-methoxybenzaldoxime				
	$\Delta_{\text{sub}}H$ (<i>anti</i>)		U 20.1 ± 1.7		MS	[1983MAJ/AZZ]
	$\Delta_{\text{sub}}H$ (<i>syn</i>)		U 32.6 ± 1.7		MS	[1983MAJ/AZZ]
C ₈ H ₉ NO ₂	[5235-04-9]	4-methoxybenzaldoxime				
	$\Delta_{\text{sub}}H$		U 67.3 ± 1.7		MS	[1983MAJ/AZZ]
C ₈ H ₉ NO ₂	[134-20-3]	methyl 2-aminobenzoate				
	$\Delta_{\text{sub}}H$	(287–298)	78.4	292.5	ME	[1987STE/MAL, 1954SER/VOI, 1960JON]
C ₈ H ₉ NO ₂	[619-45-4]	methyl 4-aminobenzoate				
	$\Delta_{\text{fus}}H$		22.55	385.1		[1990MAN/AHU]
C ₈ H ₉ NO ₂	[614-80-2]	<i>o</i> -hydroxyacetanilide				
	$\Delta_{\text{fus}}H$		21.25	364.5		[1996DOM/HEA]
C ₈ H ₉ NO ₂	[103-90-2]	<i>p</i> -hydroxyacetanilide (acetaminophen)				
	$\Delta_{\text{fus}}H$		27.6	443.2	DSC	[2009MOT/CAR]
	$\Delta_{\text{fus}}H$		27.0	440.3	DSC	[2009VEC/TOM]
	$\Delta_{\text{fus}}H$		NA	149.9		
	$\Delta_{\text{fus}}H$		26.49	441.9	AC,DSC	[2006XU/SUN]
	$\Delta_{\text{fus}}H$		26.2	443	DSC	[2004ROM/BUS]
	$\Delta_{\text{fus}}H$		26.02	441.2		[1990MAN/AHU]
	$\Delta_{\text{sub}}H$		138 ± 3	298	Fus+Vap	[2009VEC/TOM]
	$\Delta_v H$		103 ± 3	521	TGA	[2009VEC/TOM]
	$\Delta_v H$		99 ± 1	494	TGA	[2009VEC/TOM]
	C ₈ H ₉ NO ₂	[2603-10-3]	methyl N-phenylcarbamate			
$\Delta_{\text{fus}}H$			14.56	325		[1971PRI, 1996DOM/HEA]
C ₈ H ₉ NO ₂	[4389-45-1]	2-amino-3-methylbenzoic acid				
	$\Delta_{\text{fus}}H$		27.3	447.4	DSC	[2001MON/HIL2]
	$\Delta_{\text{sub}}H$	(343–357)	107.3 ± 1.8	298	ME	[2001MON/HIL2]
	$\Delta_{\text{sub}}H$	(343–357)	105.8 ± 0.8	350	ME	[2001MON/HIL2]
C ₈ H ₉ NO ₂	[2941-78-8]	2-amino-5-methylbenzoic acid				
	$\Delta_{\text{fus}}H$		27.63	450	DSC	[2001MON/HIL2]
	$\Delta_{\text{sub}}H$	(345–361)	110.6 ± 1.9	298	ME	[2001MON/HIL2]
	$\Delta_{\text{sub}}H$	(345–361)	108.9 ± 0.5	353	ME	[2001MON/HIL2]
C ₈ H ₉ NO ₂	[4389-50-8]	2-amino-6-methylbenzoic acid				
	$\Delta_{\text{fus}}H$		27.49	398.7	DSC	[2001MON/HIL2]
	$\Delta_{\text{sub}}H$	(339–355)	116.1 ± 2.0	298	ME	[2001MON/HIL2]
	$\Delta_{\text{sub}}H$	(339–355)	114.7 ± 1.2	347	ME	[2001MON/HIL2]
C ₈ H ₉ NO ₂	[52130-17-3]	3-amino-2-methylbenzoic acid				
	$\Delta_{\text{fus}}H$		38.47	458.8	DSC	[2001MON/HIL2]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{sub}}H$	(367–381)	127.8 ± 2.6	298	ME	[2001MON/HIL2]
	$\Delta_{\text{sub}}H$	(367–381)	115.6 ± 0.8	374	ME	[2001MON/HIL2]
C₈H₉NO₂	[2458-12-0]	3-amino-4-methylbenzoic acid				
	$\Delta_{\text{fus}}H$		26.89	438.8	DSC	[2001MON/HIL2]
	$\Delta_{\text{sub}}H$	(363–377)	119.4 ± 2.5	298	ME	[2001MON/HIL2]
	$\Delta_{\text{sub}}H$	(363–377)	117.3 ± 0.9	370	ME	[2001MON/HIL2]
C₈H₉NO₂	[2486-70-6]	4-amino-3-methylbenzoic acid				
	$\Delta_{\text{fus}}H$		21.77	439.4	DSC	[2001MON/HIL2]
	$\Delta_{\text{sub}}H$	(367–383)	122.0 ± 2.6	298	ME	[2001MON/HIL2]
	$\Delta_{\text{sub}}H$	(367–383)	119.8 ± 0.7	375	ME	[2001MON/HIL2]
C₈H₉NO₂	[1197-55-3]	4-aminophenylacetic acid				
	$\Delta_{\text{fus}}H$		42.7	468.2	DSC	[2002GRA/RAS]
C₈H₉NO₂S₂	[949171-63-5]	N-theonylthiocarbamic-O-ethyl ester				
	$\Delta_{\text{fus}}H$		21.9	345.9	DSC	[2007RIB/MON]
	$\Delta_{\text{sub}}H$		143.2 ± 3.1	298	C	[2007RIB/MON]
C₈H₉NO₇	[22401-53-2]	methyl 5-nitro-2-acetoxy-2,5-dihydro-2-furancarboxylate				
	$\Delta_{\text{sub}}H$		89.1 ± 2.1			[1980BAL/LEB, 1986PED/NAY]
C₈H₉N₃	[62679-52-1]	2,2-dicyanohexanenitrile				
	Δ_vH	(288–323)	61.0 ± 0.2		GS	[1994RAK/VER]
C₈H₉N₃O₂	[na]	N-(1-oxopropyl)pyrazinecarboxamide				
	$\Delta_{\text{fus}}H$		23.5	366.7		[1991ZHA/HUA]
C₈H₉O₃PS	[3811-49-2]	2-methoxy-4 <i>H</i> -1,3,2-benzodioxaphosphorin 2-sulfide				
	$\Delta_{\text{fus}}H$		16.92	327.9	DSC	[1990DON/DRE]
C₈H₁₀	[95-47-6]	1,2-dimethylbenzene				
	$\Delta_{\text{fus}}H$		13.6	247.8		[1996DOM/HEA]
	$\Delta_{\text{sub}}H$		60.1	248	B	[1986HES/LIC]
	Δ_vH	(373–423)	42.9	298	CGC	[1995CHI/HOS]
	Δ_vH	(333–419)	41.1	348	A	[1987STE/MAL]
	Δ_vH	(416–473)	38.0	431	A	[1987STE/MAL]
	Δ_vH	(471–571)	36.7	486	A	[1987STE/MAL]
	Δ_vH	(567–630)	36.7	582	A	[1987STE/MAL]
	Δ_vH	(386–416)	39.8	401		[1982CAS/FRA]
	Δ_vH		43.4	298		[1971WIL/ZWO]
	Δ_vH		43.4 ± 0.1	298	C	[1947OSB/GIN]
	Δ_vH	(337–419)	40.8	352	MM	[1945WIL/TAY, 1949FOR/NOR]
	Δ_vH	(273–323)	45.0	288		[1943PIT/SCO, 1984BOU/FRI]
C₈D₁₀	[56004-61-6]	o-xylene-d10				
	Δ_vH		43.0	298	CGC	[2008ZHA/UNH]
C₈D₁₀	[41051-88-1]	p-xylene-d10				
	Δ_vH		42.4	298	CGC	[2008ZHA/UNH]
C₈H₁₀	[108-38-3]	1,3-dimethylbenzene				
	$\Delta_{\text{fus}}H$		11.59	225.3		[1996DOM/HEA]
	Δ_vH	(360–410)	39.2	375		[2002SWI/MAL]
	Δ_vH	(327–412)	40.7	342		[1989PAR/GME]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound					
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference	
		$\Delta_v H$	(267–301)	44.7	282	A	[1987STE/MAL]
		$\Delta_v H$	(412–462)	37.5	427	A	[1987STE/MAL]
		$\Delta_v H$	(461–554)	36.4	476	A	[1987STE/MAL]
		$\Delta_v H$	(550–617)	36.2	565	A	[1987STE/MAL]
		$\Delta_v H$	(380–411)	38.7	395		[1983MAC]
		$\Delta_v H$		42.7	298		[1971WIL/ZWO]
		$\Delta_v H$		42.7 ± 0.1	298	C	[1947OSB/GIN]
		$\Delta_v H$	(331–415)	40.4	346	MM	[1987STE/MAL, 1945WIL/TAY, 1949FOR/NOR]
		$\Delta_v H$	(273–333)	43.2	288		[1943PIT/SCO, 1984BOU/FRI]
C₈H₁₀	[106-42-3]	1,4-dimethylbenzene					
		$\Delta_{\text{fus}} H$		17.11	286.3		[1996DOM/HEA]
		$\Delta_{\text{sub}} H$	(247–286)	59.4	271		[1987STE/MAL, 1974OSB/DOU]
		$\Delta_{\text{sub}} H$		60.8	286	B	[1986HES/LIC]
		$\Delta_v H$	(373–423)	42.3	298	CGC	[1995CHI/HOS]
		$\Delta_v H$	(293–323)	43.0 ± 0.1	298		[1990SMI]
		$\Delta_v H$		42.3 ± 0.01	298		[1988MES/FIN]
		$\Delta_v H$		40.3	353		[1988HOS/ARC]
		$\Delta_v H$	(411–463)	37.3	426	A	[1987STE/MAL]
		$\Delta_v H$	(460–553)	36.1	475	A	[1987STE/MAL]
		$\Delta_v H$	(551–616)	36.2	566	A	[1987STE/MAL]
		$\Delta_v H$		36.0 ± 0.1	411	C	[1985NAT/VIS]
		$\Delta_v H$		34.5 ± 0.1	436	C	[1985NAT/VIS]
		$\Delta_v H$		30.5 ± 0.1	484	C	[1985NAT/VIS]
		$\Delta_v H$		24.7 ± 0.1	540	C	[1985NAT/VIS]
		$\Delta_v H$	(380–410)	37.3	395		[1982CAS/FRA]
		$\Delta_v H$		42.3 ± 0.1	298	C	[1981HOS/SCO3]
		$\Delta_v H$		42.6	298		[1974AMB/ELL]
		$\Delta_v H$	(286–453)	42.4	301	IP,EB	[1987STE/MAL, 1974OSB/DOU]
		$\Delta_v H$		42.4	298		[1971WIL/ZWO]
		$\Delta_v H$	(303–343)	41.6	318		[1968GAW/SWI2]
		$\Delta_v H$		42.4 ± 0.1	298	C	[1947OSB/GIN]
		$\Delta_v H$	(332–413)	40.1	347	MM	[1945WIL/TAY, 1949FOR/NOR]
C₈H₁₀	[100-41-4]	ethylbenzene					
		$\Delta_{\text{fus}} H$		9.16	178.2		[1996DOM/HEA]
		$\Delta_v H$	(298–420)	41.8	313	A	[1987STE/MAL]
		$\Delta_v H$	(409–459)	37.0	424	A	[1987STE/MAL]
		$\Delta_v H$	(457–554)	35.8	472	A	[1987STE/MAL]
		$\Delta_v H$	(320–400)	40.6	335		[1986PAU/KRU]
		$\Delta_v H$	(549–617)	35.5	564	A	[1987STE/MAL]
		$\Delta_v H$		40.5 ± 0.1	328	C	[1982SVO/CHA]
		$\Delta_v H$		39.5 ± 0.1	343	C	[1982SVO/CHA]
		$\Delta_v H$		38.6 ± 0.1	358	C	[1982SVO/CHA]
		$\Delta_v H$		42.4 ± 0.1	298	C	[1981HOS/SCO3]
		$\Delta_v H$		42.3	298		[1971WIL/ZWO]
		$\Delta_v H$		42.2 ± 0.1	298	C	[1947OSB/GIN]
		$\Delta_v H$	(330–410)	40.0	345	MM	[1945WIL/TAY, 1949FOR/NOR]
C₈H₁₀	[na]	1,2-bicyclopropylacetylene					
		$\Delta_v H$		47.6 ± 0.2	298	C	[2007PAS/KUZ]
C₈H₁₀Cl₂O₂	[55701-05-8]	3-(2,2-dichloroethenyl)-2,2-dimethylcyclopropanecarboxylic acid					
		$\Delta_{\text{fus}} H$		16.32	331.5		[2007XUE/WAN]
C₈H₁₀F₃NO₃	[715-58-2]	N-trifluoroacetyl- <i>l</i> -proline, methyl ester					

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound					
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference	
		$\Delta_v H$	(303–523)	57.9	318	A	[1987STE/MAL, 1999DYK/SVO]
C ₈ H ₁₀ F ₃ NO ₅	[81084-01-7]	N-trifluoroacetyl- <i>l</i> -aspartic acid, dimethyl ester					
	$\Delta_v H$	(303–423)	58.2	318	A	[1987STE/MAL, 1999DYK/SVO]	
C ₈ H ₁₀ NO ₅ PS	[298-00-0]	O,O-dimethyl-O-(4-nitrophenyl) thiophosphate					
	$\Delta_{\text{fus}} H$		20.07	308.2		[1991ACR]	
	$\Delta_{\text{sub}} H$	(298–308)	125.1	303	GS,A	[1984KIM/WOO]	
	$\Delta_{\text{sub}} H$	(278–288)	108.7	283	GS	[1983SPE/CLI, 1979SPE/SHO]	
	$\Delta_v H$	(293–427)	88.9	308	A	[1987STE/MAL]	
		$\Delta_v H$		87.0	GS	[1979SPE/SHO]	
C ₈ H ₁₀ N ₂ O	[138-89-6]	4-N,N-dimethylaminonitrosobenzene					
	$\Delta_{\text{sub}} H$	(323–334)	82.0 ± 1.7	298	ME	[1994ACR/TUC]	
C ₈ H ₁₀ N ₂ O ₂	[619-31-8]	3-nitro-N,N-dimethylaniline					
	$\Delta_{\text{sub}} H$		92.7 ± 0.3	298	C	[1985MUR/SAK]	
	$\Delta_v H$	(427–558)	52.3	442	A,GS,EB	[1987STE/MAL, 1960AND/BID]	
	$\Delta_v H$	(357–492)	48.2	372		[1955VON/GEB]	
C ₈ H ₁₀ N ₂ O ₂	[100-23-2]	N,N-dimethyl-4-nitroaniline					
	$\Delta_{\text{sub}} H$		102.7 ± 1.1	298	C	[1985MUR/SAK]	
	$\Delta_{\text{sub}} H$	(344–366)	98.7 ± 1.7	355	ME	[1987STE/MAL, 1956MAJ]	
	$\Delta_{\text{sub}} H$	(372–393)	101.3 ± 2.0	298	ME	[1994ACR/TUC]	
C ₈ H ₁₀ N ₂ O ₃	[22809-78-3]	N-methyl-N-(4-methoxyphenyl)nitramine					
	$\Delta_{\text{fus}} H$		22.7	342.6		[2002DAS/ZAL]	
C ₈ H ₁₀ N ₂ O ₃ S	[144-80-9]	N-[(4-aminophenyl)sulfonyl]acetamide (sulfacetamide)					
	$\Delta_{\text{fus}} H$		29.8	455.2		[2002MAR/GOM]	
C ₈ H ₁₀ N ₂ O ₄ S	[156461-84-6]	N-methyl-N-(3-methylsulfonylphenyl)nitramine					
	$\Delta_{\text{fus}} H$		26.1	377.8		[2002DAS/ZAL]	
C ₈ H ₁₀ N ₂ O ₄ S	[23042-38-8]	N-methyl-N-(4-methylsulfonylphenyl)nitramine					
	$\Delta_{\text{fus}} H$		19.2	438.1		[2002DAS/ZAL]	
C ₈ H ₁₀ N ₄ O ₂	[58-08-2]	caffeine (1,3,7-trimethylxanthine)					
	$\Delta_{\text{fus}} H$		24.8	507.7	DSC	[2010GUO/SAD]	
	$\Delta_{\text{us}} H$		3.43	420.9			
	$\Delta_{\text{fus}} H$		19.86	509.5	DSC	[2007DON/LI]	
	$\Delta_{\text{us}} H$		4.02	428.2			
	$\Delta_{\text{fus}} H$		21.9	510.2	DSC	[2006PIN/DIO]	
	$\Delta_{\text{us}} H$		3.9	428	DSC	[1905DES/COR]	
	$\Delta_{\text{fus}} H$		19.38	510.2	DSC	[2005KLO/BRO]	
	$\Delta_{\text{us}} H$		0.94	426			
	$\Delta_{\text{fus}} H$		23.43	512			
	$\Delta_{\text{fus}} H$		18.3	510		[1990DOM/HEA, 1985OHM/LIP]	
	$\Delta_{\text{us}} H$		1.71	420.8			
	$\Delta_{\text{fus}} H$		20.95	508.3	DSC	[1984WEI/LEF]	
	$\Delta_{\text{sub}} H$ (form I)	(413–463)	104.8 ± 0.2	438	T	[1999EMM/PIC]	
	$\Delta_{\text{sub}} H$ (form I)	(413–463)	115	298	T	[1999EMM/PIC]	
	$\Delta_{\text{sub}} H$ (form II)	(413–463)	113.6 ± 0.2	369	T	[1999EMM/PIC]	
	$\Delta_{\text{sub}} H$ (form II)	(413–463)	119	298	T	[1999EMM/PIC]	
$\Delta_{\text{sub}} H$	(315–364)	112.6 ± 2.4		ME	[1998BOL/WIE]		
$\Delta_{\text{sub}} H$		105.1 ± 0.7		ME	[1985KAM/ZIE]		
$\Delta_{\text{sub}} H$	(373–473)	103.6	423	UV	[84EBE/FRA]		

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{sub}}H$ (<i>form I</i>)	(446–509)	100.0 ± 0.6	478	MM	[1979BOT/CAM]
	$\Delta_{\text{sub}}H$ (<i>form I</i>)		110	298		[1979BOT/CAM, 1999EMM/PIC]
	$\Delta_{\text{sub}}H$ (<i>form II</i>)	(446–509)	110.7 ± 0.7	362	MM	[1979BOT/CAM]
	$\Delta_{\text{sub}}H$ (<i>form II</i>)		114	298		[1979BOT/CAM, 1999EMM/PIC]
	Δ_vH	(634–743)	64.9 ± 2.4		DSC	[1998BOL/WIE]
C ₈ H ₁₀ O	[526-75-0]	2,3-dimethylphenol				
	$\Delta_{\text{fus}}H$		20.29	345.8	DSC	[1998JAM/PAL]
	$\Delta_{\text{fus}}H$		21.02	346		[1991ACR]
	$\Delta_{\text{sub}}H$	(283–323)	84. ± 1.0		GS	[1960AND/BID, 1970COX/PIL, 1987STE/MAL]
	Δ_vH	(433–492)	52.1	448	A,GS,EB	[1987STE/MAL, 1960AND/BID]
C ₈ H ₁₀ O	[105-67-9]	2,4-dimethylphenol				
	$\Delta_{\text{fus}}H$		12.76	297.3	DSC	[1998JAM/PAL]
	Δ_vH	(282–318)	65.9 ± 0.2		GS	[1960AND/BID, 1970COX/PIL]
	Δ_vH	(393–433)	64.6	298	CGC	[1995CHI/HOS]
	Δ_vH	(282–318)	65.9	297	A	[1987STE/MAL]
	Δ_vH	(282–318)	65.9 ± 0.2		GS	[1960AND/BID, 1970COX/PIL]
	Δ_vH	(429–486)	51.8	444	A,GS,EB	[1987STE/MAL, 1960AND/BID]
C ₈ H ₁₀ O	[95-87-4]	2,5-dimethylphenol				
	$\Delta_{\text{fus}}H$		13.81	348.1	DSC	[1998JAM/PAL]
	$\Delta_{\text{fus}}H$		23.38	348		[1991ACR]
	$\Delta_{\text{sub}}H$	(282–323)	85.0 ± 0.25		GS	[1960AND/BID, 1970COX/PIL, 1987STE/MAL]
	Δ_vH	(427–485)	51.7	442	A,GS,EB	[1987STE/MAL, 1960AND/BID]
C ₈ H ₁₀ O	[576-26-1]	2,6-dimethylphenol				
	$\Delta_{\text{fus}}H$		18.83	318.6	DSC	[1998JAM/PAL]
	$\Delta_{\text{fus}}H$		18.9	318.9		[1991ACR]
	$\Delta_{\text{sub}}H$		75.6	298		[1971MOR]
	$\Delta_{\text{sub}}H$		75.1	298		[1968MOR]
	$\Delta_{\text{sub}}H$	(277–313)	75.6 ± 0.17		GS	[1960AND/BID, 1970COX/PIL]
	Δ_vH		75.6	298		[1971MOR]
	Δ_vH		75.1	298		[1968MOR]
	Δ_vH	(417–476)	48.5	432	A,GS,EB	[1987STE/MAL, 1960AND/BID]
						Note: Author of [1968MOR] refers to the determined value as the enthalpy of vaporization even though the compound is a solid
C ₈ H ₁₀ O	[96-65-8]	3,4-dimethylphenol				
	$\Delta_{\text{fus}}H$		19.04	338.5	DSC	[1998JAM/PAL]
	$\Delta_{\text{fus}}H$		18.13	334		[1991ACR]
	$\Delta_{\text{sub}}H$		85.1	298		[1971MOR]
	$\Delta_{\text{sub}}H$		85.0	298		[1968MOR]
	$\Delta_{\text{sub}}H$	(282–323)	85.7 ± 0.1		GS	[1960AND/BID, 1970COX/PIL, 1987STE/MAL]
	Δ_vH		85.1	298		[1971MOR]
	Δ_vH		85.0	298		[1968MOR]
	Δ_vH	(444–502)	54.9	459	A,GS,EB	[1987STE/MAL, 1960AND/BID]
						Note: Author of [1968MOR] refers to the determined value as the enthalpy of vaporization even though the compound is a solid
C ₈ H ₁₀ O	[108-68-9]	3,5-dimethylphenol				

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
		$\Delta_{\text{fus}}H$	18.0	336.8		[1991ACR]
		$\Delta_{\text{sub}}H$	(282–323)	82.8 ± 0.3		GS [1960AND/BID, 1970LEN/VEL, 1987STE/MAL]
		$\Delta_{\text{sub}}H$		82 (sub)	298	[1971MOR]
		Δ_vH	(427–497)	55.3	442	A,GS,EB [1987STE/MAL, 1960AND/BID]
C ₈ H ₁₀ O	[538-86-3]	benzyl methyl ether				
		Δ_vH	(274–314)	51.4 ± 0.3	298	GS [2002KRA/VAS]
C ₈ H ₁₀ O	[103-73-1]	ethoxybenzene				
		Δ_vH	(390–454)	44.5	405	A [1987STE/MAL, 1976AMB/ELL]
		Δ_vH	(390–454)	50.7	298	[1976AMB/ELL]
		Δ_vH	(390–454)	40.7	443	[1976AMB/ELL]
		Δ_vH		51.0 ± 0.1	298	C [1975FEN/HAR]
		Δ_vH	(400–454)	44.0	415	[1965COL/COU, 1984BOU/FRI]
C ₈ H ₁₀ O	[90-00-6]	2-ethylphenol				
		$\Delta_{\text{sub}}H$	(278–317)	80.3 ± 0.5		GS [1963BID/HAN, 1970COX/PIL, 1987STE/MAL]
		Δ_vH	(393–433)	64.5	298	CGC [1995CHI/HOS]
		Δ_vH	(423–491)	50.5	438	A,GS,EB [1987STE/MAL, 1963BID/HAN]
		Δ_vH	(277–318)	63.5	292	A,GS,EB [1987STE/MAL, 1963BID/HAN]
		Δ_vH	(359–480)	51.6	374	[1955VON/GEB]
		Δ_vH	(321–492)	51.4	348	[1953STA/MUL]
		Δ_vH	(321–492)	49.5	373	[1953STA/MUL]
		Δ_vH	(321–492)	48.6	398	[1953STA/MUL]
		Δ_vH	(321–492)	47.0	423	[1953STA/MUL]
		Δ_vH	(321–492)	43.1	473	[1953STA/MUL]
C ₈ H ₁₀ O	[620-17-7]	3-ethylphenol				
		Δ_vH	(445–503)	53.1	460	A,GS,EB [1987STE/MAL, 1963BID/HAN]
		Δ_vH	(277–323)	68.1	292	A,GS,EB [1987STE/MAL, 1963BID/HAN]
		Δ_vH	(334–501)	58.3	348	[1953STA/MUL]
		Δ_vH	(334–501)	56.5	373	[1953STA/MUL]
		Δ_vH	(334–501)	55.2	398	[1953STA/MUL]
		Δ_vH	(334–501)	53.7	423	[1953STA/MUL]
		Δ_vH	(334–501)	48.8	473	[1953STA/MUL]
C ₈ H ₁₀ O	[123-07-9]	4-ethylphenol				
		Δ_vH	(444–503)	53.2	459	A,GS,EB [1987STE/MAL, 1963BID/HAN]
		Δ_vH	(337–503)	56.5	348	[1953STA/MUL]
		Δ_vH	(337–503)	54.7	373	[1953STA/MUL]
		Δ_vH	(337–503)	53.8	398	[1953STA/MUL]
		Δ_vH	(337–503)	51.3	423	[1953STA/MUL]
		Δ_vH	(337–503)	47.6	473	[1953STA/MUL]
C ₈ H ₁₀ O	[589-18-4]	4-methylbenzyl alcohol				
		$\Delta_{\text{us}}H$		0.73	179	
		$\Delta_{\text{us}}H$		0.21	210	
		$\Delta_{\text{fus}}H$	(10–350)	20.17	331.9	AC [2005SAI/IKE]
		Δ_vH	(338–376)	64.2	353	A [1987STE/MAL]
C ₈ H ₁₀ O	[13323-81-4]	<i>(dl)</i> 1-phenylethanol				
		Δ_vH	(353–480)	53.5	368	A [1987STE/MAL]
C ₈ H ₁₀ O	[60-12-8]	2-phenylethanol				
		Δ_vH	(288–363)	66.7	298	GS [2007EME/DAB]
		Δ_vH	(313–413)	U54.55	298	GC [2005HOS/GRY]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(394–613)	55.1	409	A	[1987STE/MAL]
	$\Delta_v H$	(305–363)	73.4 ± 1.5	298	GC	[1981SHC/RUD, 2007EME/DAB]
	$\Delta_v H$	(284–287)	69.7 ± 1.6	298	ME	[1958SER/VOI, 2007EME/DAB]
	$\Delta_v H$	(283–318)	68.4	298	A,ME	[1987STE/MAL, 1954SER/VOI]
	$\Delta_v H$	(406–492)	64.9 ± 0.3	298	EB	[1949DRE/SHR, 1949DRE/MAR, 2007EME/DAB]
	$\Delta_v H$	(331–492)	68	298	EB	[1947STU, 2007EME/DAB]
C ₈ H ₁₀ O	[578-58-5]	2-methylanisole				
	$\Delta_v H$		45.2			[1986BAL/GNA]
C ₈ H ₁₀ O	[104-93-8]	4-methylanisole				
	$\Delta_v H$		46.0			[1986BAL/GNA]
C ₈ H ₁₀ O	[98-85-1]	α -methyl benzyl alcohol				
	$\Delta_v H$	(358–398)	75.2	298	EB	[2004CHY/FRA]
C ₈ H ₁₀ OS	[1879-16-9]	4-methoxythioanisole				
	$\Delta_v H$		53.6			[1986BAL/GNA]
C ₈ H ₁₀ OS	[2530-10-4]	3-acetyl-2,5-dimethylthiophene				
	$\Delta_v H$		61.3 ± 1.3	298	C	[2008RIB/SAN3]
C ₈ H ₁₀ O ₂	[na]	ethyl <i>trans</i> b-(2-furyl)acrylate				
	$\Delta_v H$	(428–500)	56.8	464		[1956FRO/LOE]
C ₈ H ₁₀ O ₂	[105-13-5]	4-methoxybenzyl alcohol				
	$\Delta_v H$	(394–424)	95.6	409	A	[1987STE/MAL]
	$\Delta_v H$	(354–453)	71.7	369	EB	[1985SCH/BRU]
C ₈ H ₁₀ O ₂	[488-87-9]	1,3-dihydroxy-2,5-dimethylbenzene				
	$\Delta_v H$	(393–459)	74.7	408	A,GC	[1987STE/MAL, 1975KUN/LIL]
C ₈ H ₁₀ O ₂	[527-55-9]	1,3-dihydroxy-4,5-dimethylbenzene				
	$\Delta_v H$	(424–453)	67.5	438	A,GC	[1987STE/MAL, 1975KUN/LIL]
C ₈ H ₁₀ O ₂	[615-89-4]	1,3-dihydroxy-4,6-dimethylbenzene				
	$\Delta_v H$	(388–466)	74.7	403	A,GC	[1975KUN/LIL]
C ₈ H ₁₀ O ₂	[615-90-7]	1,4-dihydroxy-2,5-dimethylbenzene				
	$\Delta_v H$	(331–361)	101.1	346	A	[1987STE/MAL]
C ₈ H ₁₀ O ₂	[4299-72-3]	1,3-dihydroxy-5-ethylbenzene				
	$\Delta_v H$	(408–479)	81.3	423	A,GC	[1987STE/MAL, 1975KUN/LIL]
C ₈ H ₁₀ O ₂	[615-90-7]	2,5-dimethylhydroquinone				
	$\Delta_{\text{sub}} H$	(332–361)	100.8		QF	[1927COO/COO, 1960JON]
C ₈ H ₁₀ O ₂	[91-16-7]	1,2-dimethoxybenzene				
	$\Delta_v H$	(373–468)	52.7	388		[2002SU, 2004LEE/SU]
	$\Delta_v H$		68.1 ± 1.4	298	C	[2000MAT/MIR]
	$\Delta_v H$		66.9			[1958CAS/FLE2]
C ₈ H ₁₀ O ₂	[151-10-0]	1,3-dimethoxybenzene				
	$\Delta_v H$	(358–423)	60.8	373	A,GC	[1987STE/MAL, 1975KUN/LIL]
	$\Delta_v H$		61.5 ± 1.4	298	C	[2000MAT/MIR]
C ₈ H ₁₀ O ₂	[150-78-7]	1,4-dimethoxybenzene				
	$\Delta_{\text{sub}} H$		84.1 ± 2.3	298	C	[2000MAT/MIR]
	$\Delta_v H$	(298–357)	62.1	313	A	[1987STE/MAL]
	$\Delta_v H$		51.5			[1986BAL/GNA]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₈ H ₁₀ O ₂	[122-99-6] $\Delta_v H$	2-phenoxyethanol (351–519)	66.0	366	A	[1987STE/MAL, 1947STU]
C ₈ H ₁₀ O ₂	[2896-67-5] $\Delta_v H$	3-methoxy-4-hydroxytoluene (356–495)	53.2	371	A	[1987STE/MAL]
C ₈ H ₁₀ O ₂	[135-02-4] $\Delta_v H$	2-methoxybenzaldehyde	55.2			[1986BAL/GNA]
C ₈ H ₁₀ O ₂	[501-94-0] $\Delta_{\text{fus}} H$	4-hydroxybenzeneethanol	25.9	364	DSC	[2009QUE/MOT]
C ₈ H ₁₀ O ₂ S	[3112-90-1] $\Delta_{\text{fus}} H$	benzyl methyl sulfone	25.52	400.5		[1961BUS/IVI]
	$\Delta_v H$	(455–529)	64.9	470	A	[1987STE/MAL, 1999DYK/SVO]
C ₈ H ₁₀ O ₂ S	[57382-97-5] $\Delta_v H$	ethyl 2-thiopheneacetate	61.8 ± 1.3	298	C	[2009RIB/SAN2]
C ₈ H ₁₀ O ₂ S	[37784-63-7] $\Delta_v H$	ethyl 3-thiopheneacetate	63.2 ± 1.3	298	C	[2009RIB/SAN2]
C ₈ H ₁₀ O ₃	[13149-00-3] $\Delta_{\text{us}} H$ $\Delta_{\text{fus}} H$	<i>cis</i> cyclohexane-1,2-dicarboxylic acid anhydride	5.59 0.85	304 310.5		[1983GEI/NUR]
	$\Delta_v H$	(325–525)	48.8 ± 0.1			[1984NUR/MEK]
C ₈ H ₁₀ O ₃	[85-42-7] $\Delta_{\text{fus}} H$	1,2-cyclohexanedicarboxylic anhydride (80–390)	14.71	303.8	AC	[2008LU/GAO]
C ₈ H ₁₀ O ₃	[5150-42-5] $\Delta_v H$	2,3-dimethoxyphenol	76.5 ± 0.5	298	C	[2003MAT/MIR]
C ₈ H ₁₀ O ₃	[91-10-1] $\Delta_{\text{sub}} H$	2,6-dimethoxyphenol	98.4 ± 1.1	298	C	[2003MAT/MIR]
C ₈ H ₁₀ O ₃	[500-99-2] $\Delta_{\text{sub}} H$	3,5-dimethoxyphenol	101.1 ± 2.3	298	C	[2003MAT/MIR]
C ₈ H ₁₀ O ₄	[na] $\Delta_{\text{us}} H$ $\Delta_{\text{fus}} H$	<i>trans,trans</i> -2,6-octadiene-1,8-dioic acid	11.04 27.77	439 541		[1969COR/FRA]
C ₈ H ₁₀ O ₄	[na] $\Delta_{\text{fus}} H$	<i>trans,cis</i> -2,6-octadiene-1,8-dioic acid	22.78	380		[1969COR/FRA]
C ₈ H ₁₀ O ₆	[59743-08-7] $\Delta_v H$	dioxobutanedioic acid, diethyl ester (343–507)	59.3	358	A	[1987STE/MAL, 1947STU]
C ₈ H ₁₀ S	[766-92-7] $\Delta_v H$ $\Delta_v H$ $\Delta_v H$	benzyl methyl sulfide (336–368) (336–368) (323–358)	51.8 50.8 55.2 ± 2.1	351 351 298	A	[1999DYK/SVO] [1987STE/MAL] [1962MAC/MAY]
C ₈ H ₁₀ S	[622-38-8] $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$	ethyl phenyl sulfide (255–451) (338–367) (338–477) (323–358)	56.4 50.9 51.7 53.6 ± 2.1	298 353 353	A	[2004SAW/MOK] [1999DYK/SVO] [1987STE/MAL] [1962MAC/MAY]
C ₈ H ₁₀ S	[14092-00-3]	2-(methylthio)toluene				

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$		50.2			[1986BAL/GNA]
C ₈ H ₁₀ S	[623-13-2]	4-(methylthio)toluene				
	$\Delta_v H$		50.2			[1986BAL/GNA]
C ₈ H ₁₁ Cl ₃ OS	[76619-94-8]	2,3,3-trichloro-2-propenethioic acid, O-pentyl ester				
	$\Delta_v H$	(413–455)	74.1		GC	[1980PIT/KIS]
C ₈ H ₁₁ F ₃ O ₂	[1549-45-7]	trifluoroacetic acid, cyclohexyl ester				
	$\Delta_v H$	(345–420)	43	360	A,EB	[1987STE/MAL, 1969SHE/LAN, 1999DYK/SVO]
C ₈ H ₁₁ N	[121-69-7]	N,N-dimethylaniline				
	$\Delta_{\text{fus}} H$		11.56	275.6		[1972AHM/EAD2]
	$\Delta_v H$	(284–323)	53.7 ± 0.5	304		[1997VER]
	$\Delta_v H$	(363–418)	49.2	378	A	[1987STE/MAL]
	$\Delta_v H$		52.8 ± 0.1	298	C	[1982FUR/SAK]
	$\Delta_v H$	(302–467)	47.6	317	A	[1987STE/MAL, 1947STU]
C ₈ H ₁₁ N	[95-68-1]	2,4-dimethylaniline				
	$\Delta_v H$	(295–339)	61.3 ± 0.6	317		[1997VER]
	$\Delta_v H$	(383–485)	55.5	398	A	[1987STE/MAL]
	$\Delta_v H$	(326–485)	56.9	341		[1947STU]
C ₈ H ₁₁ N	[95-78-3]	2,5-dimethylaniline				
	$\Delta_{\text{fus}} H$		13.7	279		[1972AHM/EAD2]
	$\Delta_v H$	(295–339)	61.7 ± 0.7	317		[1997VER]
C ₈ H ₁₁ N	[87-62-7]	2,6-dimethylaniline				
	$\Delta_v H$	(286–326)	59.2 ± 0.3	306		[2000VER3]
	$\Delta_v H$		59.6 ± 0.3	298		[2000VER3]
	$\Delta_v H$	(373–490)	48.5	388	A	[1987STE/MAL]
	$\Delta_v H$	(317–491)	50.7	332		[1947STU]
C ₈ H ₁₁ N	[103-69-5]	N-ethylaniline				
	$\Delta_v H$	(279–318)	58.3 ± 0.6	298		[1997VER]
	$\Delta_v H$	(311–477)	52.2	326	A	[1987STE/MAL]
C ₈ H ₁₁ N	[578-54-1]	2-ethylaniline				
	$\Delta_v H$	(283–323)	60.3 ± 0.9	304.3	GS	[2000VER3]
	$\Delta_v H$		60.6 ± 0.9	298		[2000VER3]
C ₈ H ₁₁ N	[589-16-2]	4-ethylaniline				
	$\Delta_v H$	(393–491)	53.1	408	A	[1987STE/MAL]
	$\Delta_v H$	(325–490)	54.6	340		[1947STU]
C ₈ H ₁₁ N	[104-90-5]	5-ethyl-2-methylpyridine				
	$\Delta_v H$	(348–451)	45.4	363	A	[1987STE/MAL]
	$\Delta_v H$	(253–276)	51.6	264	GS	[1980VAN/PRA]
C ₈ H ₁₁ N	[98-84-0]	α -methyl benzylamine				
	$\Delta_v H$	(283–318)	54.7 ± 0.3	301	GS	[1999VER4]
	$\Delta_v H$	(283–318)	54.9 ± 0.3	298	GS	[1999VER4]
C ₈ H ₁₁ N	[618-36-0]	<i>dl</i> α -methyl benzylamine				
	$\Delta_v H$	(292–318)	36.7	305	A	[1987STE/MAL]
	$\Delta_v H$		54.5 ± 0.1	298	C	[1987ATI/SAI]
C ₈ H ₁₁ N	[3886-69-9]	(+) α -methylbenzylamine				
	$\Delta_v H$		54.1 ± 0.1	298	C	[1987ATI/SAI]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₈ H ₁₁ N	[2627-86-3] $\Delta_v H$	(-) α -methylbenzylamine	54.6 ± 0.1	298	C	[1987ATI/SAI]
C ₈ H ₁₁ N	[104-84-7] $\Delta_v H$	4-methylbenzylamine (353–466)	54.4	368	A	[1987STE/MAL]
C ₈ H ₁₁ N	[64-04-0] $\Delta_v H$ $\Delta_v H$	2-phenylethylamine (277–351) (277–351)	55.7 ± 0.2 56.8 ± 0.2	313 298		[2009MOK/RAZ] [2009MOK/RAZ]
C ₈ H ₁₁ N	[695-98-7] $\Delta_v H$	2,3,5-trimethylpyridine (293–426)	44.0	359		[1995SAK/UEO]
C ₈ H ₁₁ N	[1462-84-6] $\Delta_v H$ $\Delta_v H$ $\Delta_v H$	2,3,6-trimethylpyridine	48.5 47.5 45.7	328 343 368	C C C	[1985MAJ/SVO2] [1985MAJ/SVO2] [1985MAJ/SVO2]
C ₈ H ₁₁ N	[108-75-8] $\Delta_{\text{fus}} H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$	2,4,6-trimethylpyridine	9.54 50.4 ± 2.9 50.2 46.5 51.2 50.3 ± 0.2 48.3 47.2 45.5	229 298 298 363 313 298 328 343 368		[1996DOM/HEA] CGC [2009LIP/CHI2] CGC [1995CHI/HOS] [1995SAK/UEO] EB [1990LEN] [1985MAJ/SVO2] C [1985MAJ/SVO2] C [1985MAJ/SVO2] C [1985MAJ/SVO2]
C ₈ H ₁₁ N	[622-39-9] $\Delta_v H$	2-propylpyridine (338–445)	46.6	353	A	[1987STE/MAL]
C ₈ H ₁₁ N	[4673-31-8] $\Delta_v H$	3-propylpyridine (350–450)	49.9	365	A	[1987STE/MAL]
C ₈ H ₁₁ N	[1122-81-2] $\Delta_v H$	4-propylpyridine (354–465)	47.8	369	A	[1987STE/MAL]
C ₈ H ₁₁ N	[103434-09-9] $\Delta_{\text{sub}} H$	1-norbornylisocyanide	60.6 ± 0.5	298		[1987MEI/DOG]
C ₈ H ₁₁ N	[3211-90-3] $\Delta_{\text{us}} H$ $\Delta_{\text{fus}} H$	<i>exo</i> -2-cyanobicyclo[2.2.1]heptane	7.95 2.93	237.7 298.8		[1996DOM/HEA]
C ₈ H ₁₁ N	[3211-87-8] $\Delta_{\text{us}} H$ $\Delta_{\text{fus}} H$	<i>endo</i> -2-cyanobicyclo[2.2.1]heptane	2.25 2.96	177.3 331.2		[1996DOM/HEA]
C ₈ H ₁₁ NO	[122-98-5] $\Delta_v H$	2-anilinoethanol (377–553)	69.9	392	A	[1987STE/MAL, 1947STU]
C ₈ H ₁₁ NO	[94-70-2] $\Delta_v H$	2-ethoxyaniline (373–458)	57.3	388	A	[1987STE/MAL]
C ₈ H ₁₁ NO	[156-43-4] $\Delta_v H$	4-ethoxyaniline (421–523)	61.2	436	A	[1987STE/MAL]
C ₈ H ₁₁ NO ₂	[4355-17-3] $\Delta_{\text{us}} H$ $\Delta_{\text{fus}} H$	3-azabicyclo[3.3.1]nonane-2,4-dione	16.3 3.3	408.6 463.6		[2007HUL/JOH]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₈ H ₁₁ N ₃ O ₄	[87473-90-3]	methyl N-(4,6-dimethoxypyrimidin-2-yl)carbamate				
	$\Delta_{\text{fus}}H$	(80–380)	26.29	357.2	AC	[2004XIN/TAN]
C ₈ H ₁₁ N ₅	[116988-56-8]	8-ethyl-9-methyladenine				
	$\Delta_{\text{sub}}H$		127.1 ± 0.7			[1994ZIE/ZIE]
	$\Delta_{\text{sub}}H$	(365–370)	115.2 ± 1.0	368	ME	[1987KAM/ZIE]
C ₈ H ₁₁ N ₅	[139909-51-6]	6,8,9-trimethyladenine				
	$\Delta_{\text{fus}}H$		23.1	438		[1994ZIE/ZIE]
	$\Delta_{\text{sub}}H$	(334–342)	98.6 ± 0.2	338	ME	[1994ZIE/ZIE]
C ₈ H ₁₁ N ₅	[3013-82-9]	N,N,9-trimethyladenine				
	$\Delta_{\text{sub}}H$	(319–349)	101.7 ± 2.1		ME	[1984ZIE/ZIE]
C ₈ H ₁₁ N ₅ O ₂	[na]	2-amino-9-[(2-hydroxyethoxy)methyl]-9H-purine				
	$\Delta_{\text{fus}}H$		42.2	462.2		[1995KRI/VES]
C ₈ H ₁₁ N ₅ O ₃	[59277-89-3]	2-amino-9-[(2-hydroxyethoxy)methyl]-1,9-dihydro-6H-purin-6-one				
	$\Delta_{\text{fus}}H$		30.44	528.2		[1995KRI/VES]
C ₈ H ₁₂	[13027-75-3]	<i>anti</i> -tricyclo[4.2.0.0 ^{2,5}]octane				
	Δ_vH		41.8 ± 1.7	298		[2008OSM/CAT]
C ₈ H ₁₂	[28636-10-4]	<i>syn</i> -tricyclo[4.2.0.0 ^{2,5}]octane				
	Δ_vH		41.8 ± 1.7	298		[2008OSM/CAT]
C ₈ H ₁₂	[250-21-5]	tricyclo[3.3.0.0 ^{2,6}]octane				
	Δ_vH	(273–343)	40.0	273		[1988LET/SEW]
	Δ_vH	(273–343)	39.1	298		[1988LET/SEW]
	Δ_vH	(273–343)	38.3	323		[1988LET/SEW]
C ₈ H ₁₂	[na]	cyclooctadiene (mixed isomers)				
	Δ_vH	(290–474)	34.6	305	A	[1987STE/MAL]
C ₈ H ₁₂	[1552-12-1]	<i>cis</i> cis 1,5-cyclooctadiene				
	Δ_vH		43.4 ± 0.1	298	C	[1996VAR/PAS]
C ₈ H ₁₂	[10092-71-4]	1,5-cyclooctadiene				
	$\Delta_{\text{us}}H$		0.38	194.4		
	$\Delta_{\text{fus}}H$		9.83	204		[1996DOM/HEA, 1975LEB/TSV]
	Δ_vH	(348–386)	40.9	363	A	[1987STE/MAL]
C ₈ H ₁₂	[6553-48-6]	<i>dl</i> <i>trans</i> 1,2-divinylcyclobutane				
	Δ_vH	(319–371)	38.9 ± 0.5	298	EB	[1996VAR/PAS]
	Δ_vH	(350–385)	39.1	365	A	[1987STE/MAL]
	Δ_vH		42.3	298		[1973RAU/GEY]
	Δ_vH		39.0 ± 0.5	367		[1973RAU/GEY]
C ₈ H ₁₂	[100-40-3]	<i>dl</i> 4-vinyl-1-cyclohexene				
	Δ_vH	(292–405)	40.1	307	A	[1987STE/MAL]
C ₈ H ₁₂	[931-64-6]	bicyclo[2.2.2]octane				
	$\Delta_{\text{us}}H$		0.19	110.5		
	$\Delta_{\text{us}}H$		5.65	176.5		
	$\Delta_{\text{fus}}H$		5.4	389.8		[1977WON/WES]
	$\Delta_{\text{sub}}H$		43.8 ± 0.4		C	[1970WES/WON, 1977PED/RYL, 1971WON/WES]
C ₈ H ₁₂	[21426-37-9]	dispiro[2.0.2.2]octane				
	Δ_vH	(280–369)	38.3		DSC	[1995BEC/RUC]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₈ H ₁₂	[25399-32-0] $\Delta_v H$	dispiro[2.1.2.1]octane (278–362)	32.2		DSC	[1995BEC/RUC]
C ₈ H ₁₂ BrN ₅ O ₃	[81475-44-7] $\Delta_{\text{fus}} H$	8-bromo-9-[(2-hydroxyethoxy)methyl]guanine	36.44	452.9		[1999ZIE/GOL]
C ₈ H ₁₂ Cl ₂ O ₅	[na] $\Delta_v H$	diethylene glycol bis(chloroacetate) (421–586)	87.3	436		[1987STE/MAL, 1947STU, 1999DYK/SVO]
C ₈ H ₁₂ NO ₅ PS ₂	[115-93-5] $\Delta_{\text{fus}} H$	O,O-dimethyl O-(4-aminosulfonylphenyl)phosphorodithioate	26.21	344.2	DSC	[1990DON/DRE]
C ₈ H ₁₂ N ₂	[42046-61-7] $\Delta_v H$	pentylmalodinitrile (298–328)	66.9 ± 0.4		GS	[1990BEC/DOG]
C ₈ H ₁₂ N ₂	[629-40-3] $\Delta_{\text{fus}} H$ $\Delta_v H$	suberic acid dinitrile (suberonitrile) (303–339)	21.97 77.3	268.9 318	DSC A	[2007BAD/BLA] [1987STE/MAL]
C ₈ H ₁₂ N ₂	[3333-52-6] $\Delta_{\text{fus}} H$ $\Delta_{\text{fus}} H$ $\Delta_{\text{sub}} H$	tetramethylsuccinonitrile	18.1 7.15 81.2 ± 1.7	345 442		[1996DOM/HEA] [1973LEB/KAT, 1977PED/RYL]
C ₈ H ₁₂ N ₂	[1124-11-4] $\Delta_{\text{sub}} H$	tetramethylpyrazine	94.6 ± 4.0	298	C	[1996RIB/MOR]
C ₈ H ₁₂ N ₂ O	[15029-30-8] $\Delta_{\text{sub}} H$	1-(cyanoacetyl)piperidine	103.5 ± 1.9	298	C	[2008RIB/CAB]
C ₈ H ₁₂ N ₂ O ₂	[31703-08-9] $\Delta_{\text{fus}} H$ $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$	1,3-dimethyl-5-ethyluracil (312–321) (300–316) (319–340)	19.4 98.7 ± 1.7 99.3 ± 0.2 110 ± 1.2	354.4 316 308 330		[1996KAM/ZIE] [1996KAM/ZIE] [1983COL/JIM] [1983COL/JIM]
C ₈ H ₁₂ N ₂ O ₂	[na] $\Delta_{\text{fus}} H$	1,6-hexamethylene diisocyanate	18.64	206.1		[1996DOM/HEA]
C ₈ H ₁₂ N ₂ O ₃	[57-44-3] $\Delta_{\text{fus}} H$	5,5-diethyl-2,4,6(1 <i>H</i> ,3 <i>H</i> ,5 <i>H</i>)-pyrimidinetrione (barbital)	24.98	462.6		[1986CHU/DEM]
C ₈ H ₁₂ N ₄ O ₁₀	[2555-54-6] $\Delta_{\text{fus}} H$ $\Delta_{\text{fus}} H$	2,2-dinitropropyl-4,4-dinitropentanoate	23.01 6.28	330.6 370.8		[1971ROS/HOL]
C ₈ H ₁₂ N ₄ O ₁₀	[34001-51-9] $\Delta_{\text{fus}} H$ $\Delta_{\text{fus}} H$	2-methyl-2-nitropropyl-4,4,4-trinitrobutyrate	24.69 5.27	346.1 349.4		[1971ROS/HOL]
C ₈ H ₁₂ O	[59348-18-4] $\Delta_v H$	1-methylnorcamphor	47.6			[1984KOZ/TIM]
C ₈ H ₁₂ O	[7040-43-9] $\Delta_v H$ $\Delta_v H$	2- <i>tert</i> -butylfuran (270–308) (270–308)	38.7 ± 0.4 38.1 ± 0.4	289 298	GS GS	[1998VER/WEL] [1998VER/WEL]
C ₈ H ₁₂ OS	[20387-67-1] $\Delta_{\text{fus}} H$ $\Delta_{\text{fus}} H$	2-oxa-6-thiaadamantane	4.11 8.12	224 557		[1978AND/CAR]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₈ H ₁₂ O ₂	[10279-96-6]	2,6-dioxaadamantane				
	$\Delta_{\text{fus}}H$		5.84	276		
	$\Delta_{\text{fus}}H$		3.78	444		[1978AND/CAR]
C ₈ H ₁₂ O ₂	[562-46-9]	4,4-dimethyl-1,3-cyclohexanedione				
	$\Delta_{\text{sub}}H$		99.2 ± 2.1	298	ME	[1993PIL/PAR]
C ₈ H ₁₂ O ₂	[126-81-8]	5,5-dimethyl-1,3-cyclohexanedione				
	$\Delta_{\text{sub}}H$		99.8 ± 1.1	298	ME	[1993PIL/PAR]
C ₈ H ₁₂ O ₂	[933-52-8]	2,2,4,4-tetramethyl-1,3-cyclobutanedione				
	$\Delta_{\text{sub}}H$		70.3 ± 3.5		HSA	[1975CHI]
	$\Delta_{\text{sub}}H$		72.2 ± 0.6			[1971SEL2]
	$\Delta_{\text{sub}}H$		72.4 ± 0.6		C	[1971MOR]
C ₈ H ₁₂ O ₂	[1489-74-3]	1,5-cyclooctanedione				
	$\Delta_{\text{fus}}H$		11.92	341.2		[1972ALV/BOR]
C ₈ H ₁₂ O ₄	[623-91-6]	diethyl fumarate				
	Δ_vH	(326–492)	53.2	341	A	[1987STE/MAL, 1947STU]
C ₈ H ₁₂ O ₄	[141-05-9]	diethyl maleate				
	Δ_vH	(330–498)	55.2	345	A	[1987STE/MAL, 1947STU]
C ₈ H ₁₂ O ₄ S	[925-47-3]	thiodiacetic acid, diethyl ether				
	Δ_vH	(385–448)	77.7	400		[1999DYK/SVO]
C ₈ H ₁₂ S ₆	[6327-74-8]	1,3,5,7-tetramethyl-2,4,6,8,9,10-hexathiaadamantane				
	$\Delta_{\text{fus}}H$		23.7	501.4	DSC	[2002BOU/SAI]
C ₈ H ₁₃ ClN ₂ O ₂	[4902-51-2]	5-chloro-3-(1,1-dimethylethyl)-6-methyl-2,4(1 <i>H</i> ,3 <i>H</i>)-pyrimidinedione				
	$\Delta_{\text{fus}}H$		12.51	448	DSC	[1990DON/DRE]
C ₈ H ₁₃ N ₂ O ₃	[57-44-3]	5,5'-diethylbarbituric acid				
	$\Delta_{\text{fus}}H$		0.7	413.3		
	$\Delta_{\text{fus}}H$		24.8	462	DSC	[2009RIB/RIB3]
	$\Delta_{\text{sub}}H$	(355–377)	113.9 ± 0.6	366	ME	[2009RIB/RIB3]
	$\Delta_{\text{sub}}H$	(355–377)	117.3 ± 0.6	298	ME	[2009RIB/RIB3]
C ₈ H ₁₄	[280-33-1]	bicyclo[2.2.2]octane				
	$\Delta_{\text{fus}}H$		4.6	164.3		
	$\Delta_{\text{fus}}H$		8.37	447.5		[1991ACR, 1970WON/WES]
	$\Delta_{\text{fus}}H$		4.6	164		[1984DOM/EVA]
	$\Delta_{\text{sub}}H$	(323–363)	46.3 ± 0.8		BG	[1971BOY/SAN, 1977PED/RYL, 1987STE/MAL]
	$\Delta_{\text{sub}}H$		47.7 ± 0.8	298		[1971BOY/SAN, 1977PED/RYL]
	$\Delta_{\text{sub}}H$		48.0 ± 2		C	[1970WES/WON, 1971BOY/SAN, 1977PED/RYL]
C ₈ H ₁₄	[1755-05-1]	<i>cis</i> bicyclo[3.3.0]octane				
	Δ_vH	(298–318)	42.0	308	A	[1987STE/MAL]
	Δ_vH		41.5 ± 0.4	318		[1970CHA/MCN]
	Δ_vH		43.1 ± 0.8	298		[1970CHA/MCN]
C ₈ H ₁₄	[5597-89-7]	<i>trans</i> bicyclo[3.3.0]octane				
	Δ_vH	(298–320)	41.4	309	A	[1987STE/MAL]
	Δ_vH		41.3 ± 0.4	320		[1970CHA/MCN]
	Δ_vH		42.7 ± 0.8	298		[1970CHA/MCN]
C ₈ H ₁₄	[28282-35-1]	<i>cis</i> bicyclo[4.2.0]octane				
	Δ_vH	(298–347)	40.7	313	A	[1987STE/MAL]
	Δ_vH		39.5 ± 0.4	347		[1970CHA/MCN]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
		$\Delta_v H$	42.7 ± 1.2	298		[1970CHA/MCN]
C ₈ H ₁₄	[16526-90-2]	<i>cis</i> bicyclo[5.1.0]octane				
		$\Delta_v H$	(297–322) 43.6 ± 0.8	309	A	[1987STE/MAL, 1970CHA/MCN]
C ₈ H ₁₄	[286-43-1]	bicyclo[5.1.0]octane				
		$\Delta_v H$	43.5 ± 0.8	298		[2008OSM/CAT]
C ₈ H ₁₄	[931-88-4]	cyclooctene				
		$\Delta_{\text{us}}H$	9.8	190.1		
		$\Delta_{\text{fus}}H$	1.81	259.2		[1994LEB/SMI]
		$\Delta_v H$	(273–411) 42.0	288	A	[1987STE/MAL]
		$\Delta_v H$	(273–333) 41.6	300		[1941LIS]
C ₈ H ₁₄	[695-12-5]	vinylcyclohexane				
		$\Delta_v H$	39.7 ± 0.2	298	GCC	[1979FUC/PEA]
C ₈ H ₁₄	[3524-75-2]	allylcyclopentane				
		$\Delta_v H$	40.4 ± 0.2	298	GCC	[1979FUC/PEA]
C ₈ H ₁₄	[627-58-7]	2,5-dimethyl-1,5-hexadiene				
		$\Delta_v H$	(330–388) 38.8	345	A	[1987STE/MAL]
C ₈ H ₁₄	[24253-25-6]	3,3-dimethyl-1,5-hexadiene				
		$\Delta_v H$	(293–371) 35.2	308	A	[1987STE/MAL]
C ₈ H ₁₄	[na]	3,4-dimethylhexadiene				
		$\Delta_{\text{sub}}H$	53.1			[1956SEK/SUZ, 1960JON]
C ₈ H ₁₄	[1453-24-3]	1-ethylcyclohexene				
		$\Delta_v H$	(353–412) 39.1	368	A	[1987STE/MAL]
		$\Delta_v H$	(332–411) 40.1	347	MM	[1960CAM/ROS]
C ₈ H ₁₄	[2439-79-4]	1-methylbicyclo[4.1.0]heptane				
		$\Delta_v H$	(340–394) 37.2	355	A	[1987STE/MAL]
C ₈ H ₁₄	[629-05-0]	1-octyne				
		$\Delta_v H$	42.3 ± 0.1	298	C	[1983HAL/STE]
		$\Delta_v H$	(357–400) 38.5	372	A	[1987STE/MAL, 1970EIS/ORA, 1984BOU/FRI]
C ₈ H ₁₄	[2809-67-8]	2-octyne				
		$\Delta_v H$	44.5 ± 0.1	298	C	[1983HAL/STE]
		$\Delta_v H$	(368–412) 39.9	383	A	[1987STE/MAL, 1970EIS/ORA, 1984BOU/FRI]
C ₈ H ₁₄	[15232-76-5]	3-octyne				
		$\Delta_v H$	43.9	298		[UR/FUC, 1985MAJ/SVO]
		$\Delta_v H$	(363–406) 39.7	378	A	[1987STE/MAL, 1970EIS/ORA, 1984BOU/FRI]
C ₈ H ₁₄	[1942-45-6]	4-octyne				
		$\Delta_v H$	42.7 ± 0.1	298	C	[1983HAL/STE]
		$\Delta_v H$	(362–405) 39.6	377	A	[1987STE/MAL, 1970EIS/ORA, 1984BOU/FRI]
C ₈ H ₁₄	[765-90-2]	<i>endo</i> -2-methylbicyclo[2.2.1]heptane				
		$\Delta_{\text{us}}H$	4.73	152.4		
		$\Delta_{\text{fus}}H$	1.62	278.3		[1996DOM/HEA]
C ₈ H ₁₄	[872-78-6]	<i>exo</i> 2-methylbicyclo[2.2.1]heptanes				
		$\Delta_{\text{fus}}H$	8.37	164.1		[1996DOM/HEA]
C ₈ H ₁₄ Br ₂	[29974-69-4]	1,2-dibromocyclooctane				
		$\Delta_v H$	(292–354) 50.3	307	A	[1987STE/MAL, 1941LIS]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₈ H ₁₄ ClN ₅	[1912-24-9]	2-chloro-4-ethylamino-6-isopropylamino-1,3,5-triazine (atrazine)				
	$\Delta_{\text{sub}}H$	(324–354)	114.6	339	GS	[1982GRA/FOS]
	$\Delta_{\text{sub}}H$	(323–403)	113.8	338	GS-GC	[1964FRI/SAM, 1987STE/MAL]
C ₈ H ₁₄ Cl ₂ S	[16660-53-0]	(2-chlorocyclohexyl)(2-chloroethyl) sulfide				
	Δ_vH	(293–333)	62.5	308	A,GS	[1987STE/MAL, 1948RED/CHA, 1999DYK/SVO]
C ₈ H ₁₄ N ₂	[62842-38-0]	2-piperidinopropionitrile				
	Δ_vH	(283–318)	57.6 ± 0.3		GS	[1997WEL/VER]
C ₈ H ₁₄ N ₂	[49570-30-1]	1,4-dimethyl-2,3-diazabicyclo[2.2.2]octane				
	$\Delta_{\text{sub}}H$		72.0 ± 0.5	298	C	[1976ENG/MEL]
C ₈ H ₁₄ N ₂ O ₂	[19701-85-0]	α -acetylproline N-methylamide				
	$\Delta_{\text{sub}}H$	(308–318)	69.1	313	A	[1987STE/MAL, 1955AIH]
C ₈ H ₁₄ N ₂ O ₂	[na]	β -acetylproline N-methylamide				
	$\Delta_{\text{sub}}H$	(319–335)	60.7	327	A	[1987STE/MAL, 1955AIH]
C ₈ H ₁₄ N ₄ OS	[21087-64-9]	4-amino-6-(1,1-dimethylethyl)-3-(methylthio)1,2,4-triazin-5(4H)-one				
	$\Delta_{\text{fus}}H$		18.0	399.4	DSC	[1991ACR, 1990DON/DRE]
C ₈ H ₁₄ N ₅ Cl	[1912-24-9]	6-chloro-N-ethyl-N'(isopropyl)-1,3,5-triazine-2,4-diamine				
	$\Delta_{\text{fus}}H$		38.15	449.7	DSC	[1990DON/DRE]
C ₈ H ₁₄ N ₆ O ₁₀	[na]	1,7-diacetoxy-2,4,6-trinitro-2,4,6-triazaheptane				
	$\Delta_{\text{fus}}H$		38.49	422.5		[1996DOM/HEA]
C ₈ H ₁₄ O	[502-49-8]	cyclooctanone				
	Δ_vH	(343–383)	54.4	298	CGC	[1995CHI/HOS]
	Δ_vH	(343–383)	53.6	298	CGC	[1995CHI/HOS]
	Δ_vH	(343–383)	54.2	298	CGC	[1995CHI/HOS]
	Δ_vH	(323–403)	47.3	338	A	[1987STE/MAL]
	Δ_vH	(394–484)	46.8	409	A,EB	[1987STE/MAL, 1976MEY/HOT]
	Δ_vH		48.5 ± 0.6	298		[1972WOL]
C ₈ H ₁₄ O	[66266-68-2]	2-ethyl-2-hexenal				
	Δ_vH	(326–448)	48.4	341	A	[1987STE/MAL, 1961DYK/SEP]
C ₈ H ₁₄ O	[28419-86-5]	2-ethyl-4-methyl-2-pentenal				
	Δ_vH	(311–436)	46.7	326	A	[1987STE/MAL, 1961DYK/SEP]
C ₈ H ₁₄ O	[110-93-0]	6-methyl-5-hepten-2-one				
	Δ_vH	(364–393)	45.9	379		[1989WAN/YIN]
	Δ_vH	(328–451)	44.7 ± 0.2	390		[1988BAG/GUR]
C ₈ H ₁₄ O	[1193-70-0]	(dl) 2-propylcyclopentanone				
	Δ_vH	(332–457)	46.0	347	A	[1987STE/MAL]
C ₈ H ₁₄ O	[283-27-2]	3-oxabicyclo[3.2.2]nonane				
	$\Delta_{\text{us}}H$		7.02	208.5		
	$\Delta_{\text{fus}}H$		6.75	448.4		[1996DOM/HEA]
	$\Delta_{\text{sub}}H$		53.1 ± 0.5			[1971WON/WES, 1977PED/RYL]
C ₈ H ₁₄ O ₂	[5292-21-7]	cyclohexylacetic acid				
	$\Delta_{\text{fus}}H$		13.8	302.6		[2008DOM/MOR]
C ₈ H ₁₄ O ₂	[5698-29-3]	octanolactone				
	Δ_vH	(345–380)	48.9 ± 0.2	362	MM	[1991WIB/WAL]
	Δ_vH	(345–380)	52.8 ± 1.3	298	MM	[1991WIB/WAL]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₈ H ₁₄ O ₂	[698-76-0] $\Delta_v H$	d-octanolactone (288–353)	67.0 ± 0.2	298	GS	[2007EME/KOZ]
C ₈ H ₁₄ O ₂	[na] $\Delta_v H$	acrylic acid, neopentyl ester (301–325)	45.7	313	A	[1987STE/MAL]
C ₈ H ₁₄ O ₂	[3891-33-6] $\Delta_v H$	1,4-butanediol divinyl ether (335–440)	49.0	350	A	[1987STE/MAL]
C ₈ H ₁₄ O ₂	[97-88-1] $\Delta_{\text{fus}} H$	butyl methacrylate	15.55	197.8		[1995LEB/KUL]
	$\Delta_v H$	(343–373)	47.4	358	A	[1987STE/MAL]
	$\Delta_v H$	(344–437)	45.1	359	A	[1987STE/MAL]
C ₈ D ₁₄ O ₂	[na] $\Delta_{\text{fus}} H$	perdeuteriobutylmethacrylate	16.04	198.1		[1995LEB/KUL]
C ₈ H ₁₄ O ₂	[na] $\Delta_v H$	cyclopentanecarboxylic acid ethyl ester (275–308)	51.2 ± 0.6		GS	[1996VIT/CHA]
C ₈ H ₁₄ O ₂	[177-10-6] $\Delta_v H$	1,4-dioxaspiro[4.5]decane (278–308)	50.6 ± 0.6	298	GS	[1998VER/PEN, 2002VER]
C ₈ H ₁₄ O ₂	[622-45-7] $\Delta_{\text{us}} H$ $\Delta_{\text{fus}} H$	cyclohexyl acetate	5.23	221.8		[1999KAB/KOZ]
	$\Delta_v H$		52.3 ± 0.2	298	C	[2004PAU/ZAI, 2003ZAI/VERI]
	$\Delta_v H$	(253–283)	56.5 ± 0.5	298	ME	[2003ZAI/VER]
	$\Delta_v H$	(253–283)	52.6 ± 0.5	298	ME	[2003ZAI/VER]
	$\Delta_v H$	(274–318)	52.3 ± 0.8	298	GS	[2003ZAI/VER]
	$\Delta_v H$	(333–378)	51.7	298	CGC	[1999VER/HEI]
	$\Delta_v H$	(278–318)	51.7 ± 0.2	298	GS	[1996VER/BEC]
	$\Delta_v H$	(368–446)	46.7	383	A,EB	[1987STE/MAL, 1969SHE/LAN]
C ₈ H ₁₄ O ₂	[585-07-9] $\Delta_v H$	methylacrylic acid, <i>tert</i> -butyl ester (313–410)	42.9	328	A	[1987STE/MAL]
C ₈ H ₁₄ O ₂	[2998-23-4] $\Delta_v H$	pentyl acrylate (325–440)	44.9	340	A	[1987STE/MAL]
C ₈ H ₁₄ O ₂	[142-30-2] $\Delta_v H$	2,5-dimethyl-3-hexyne-2,5-diol	82.8 ± 1.0	298	CGC	[2006UMN/KWE]
C ₈ H ₁₄ O ₂	[56922-71-5] $\Delta_v H$	propyl 3-methylbut-2-enoate (278–311)	53.0 ± 0.2	298	GS	[2008EME/TOK]
C ₈ H ₁₄ O ₂	[25859-51-2] $\Delta_v H$	isopropyl 3-methylbut-2-enoate (279–313)	50.0 ± 0.2	298	GS	[2008EME/TOK]
C ₈ H ₁₄ O ₃	[106-31-0] $\Delta_v H$	butyric anhydride (349–470)	49.1	364	A	[1987STE/MAL]
C ₈ H ₁₄ O ₃	[764-99-8] $\Delta_v H$	diethylene glycol divinyl ether (336–470)	50.0	351	A	[1987STE/MAL]
C ₈ H ₁₄ O ₃	[607-97-6] $\Delta_v H$	2-ethylacetoacetic acid, ethyl ester (313–471)	53.3	328	A	[1987STE/MAL, 1947STU]
C ₈ H ₁₄ O ₃	[21884-26-4] $\Delta_v H$	isopropyl levulinate (321–481)	56.6	336	A	[1987STE/MAL, 1947STU]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound					
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference	
		$\Delta_v H$	52.0	422		[1931SCH/COW]	
C ₈ H ₁₄ O ₃	[645-67-0]	propyl levulinate					
	$\Delta_v H$	(332–495)	56.3	347	A	[1987STE/MAL]	
	$\Delta_v H$		54.0	436		[1931SCH/COW]	
C ₈ H ₁₄ O ₄	[na]	2-acetoxypropionic acid, propyl ester					
	$\Delta_v H$	(318–469)	59.5	333	A	[1987STE/MAL]	
C ₈ H ₁₄ O ₄	[na]	3-acetoxypropionic acid, propyl ester					
	$\Delta_v H$	(361–373)	74.7	367	A	[1987STE/MAL]	
C ₈ H ₁₄ O ₄	[123-25-1]	diethyl succinate					
	$\Delta_v H$	(327–490)	56.5	342	A	[1987STE/MAL, 1947STU]	
C ₈ H ₁₄ O ₄	[615-81-6]	diisopropyl oxalate					
	$\Delta_v H$	(418–501)	57.8	433	A	[1987STE/MAL]	
	$\Delta_v H$	(316–467)	57.6	331	A	[1987STE/MAL, 1947STU]	
C ₈ H ₁₄ O ₄	[627-93-0]	dimethyl adipate					
	$\Delta_v H$	(353–443)	74.6	368		[2007LEE/LAI]	
	$\Delta_v H$	(294–373)	69.0 ± 0.2	298	GS	[2006VER/KOZ]	
	$\Delta_v H$	(293–344)	67.1 ± 0.3	298	GS	[2006VAS/VER, 2006VER/KOZ]	
	$\Delta_v H$	(293–323)	U 55.9 ± 2.0	298	TE	[1997CHE/LIA, 2006VER/KOZ]	
	$\Delta_v H$	(382–500)	58.8	397	A	[1987STE/MAL]	
	$\Delta_v H$	(428–498)	73.4	298		[1963VLA/GRA, 2006VER/KOZ]	
C ₈ H ₁₄ O ₄	[615-98-5]	dipropyl oxalate					
	$\Delta_v H$	(326–487)	57.8	341	A	[1987STE/MAL, 1947STU]	
C ₈ H ₁₄ O ₄	[123-80-8]	ethylene dipropionate					
	$\Delta_v H$		67.6 ± 0.5	298	C	[1986NIL/WAD]	
C ₈ H ₁₄ O ₄	[609-08-5]	2-methylmalonic acid, diethyl ester					
	$\Delta_v H$	(312–475)	52.5	327	A	[1987STE/MAL, 1947STU]	
C ₈ H ₁₄ O ₄	[505-48-6]	octanedioic acid (suberic acid)					
	$\Delta_{\text{us}}H$		2.0	335.9			
	$\Delta_{\text{us}}H$		9.1	403.6			
	$\Delta_{\text{fus}}H$		30.7	413.2		[2005ROU/TEM]	
	$\Delta_{\text{fus}}H$		28.82	415.3		[1991ACR]	
	$\Delta_{\text{sub}}H$	(348–378)	168 ± 7		TPD	[2007CAP/LOV]	
	$\Delta_{\text{sub}}H$	(310–320)	148		TPTD	[2001CHA/TOB]	
	$\Delta_{\text{sub}}H$		147.8 ± 3.8	298		[1999RIB/MON, 1960DAV/THO]	
	$\Delta_{\text{sub}}H$	(379–407)	143.1 ± 3.8	393	M	[1960DAV/THO, 1970COX/PIL, 1987STE/MAL]	
	$\Delta_v H$	(424–503)	116.7 ± 0.8	298	CGC	[2005ROU/TEM]	
	$\Delta_v H$	(445–619)	91.4	460	A	[1987STE/MAL, 1947STU]	
	C ₈ H ₁₄ O ₄	[360-51-3]	tetramethysuccinic acid				
		$\Delta_{\text{us}}H$		13.43	383		
		$\Delta_{\text{fus}}H$		6.47	464		[1996DOM/HEA]
C ₈ H ₁₄ O ₄ S	[925-47-3]	thiodiacetic acid, diethyl ester					
	$\Delta_v H$	(384–448)	77.3	399	A	[1987STE/MAL]	
C ₈ H ₁₄ O ₅	[na]	isopropyl[1-(methoxycarbonyl)ethyl] carbonate					
	$\Delta_v H$	(330–493)	55.5	345	A	[1987STE/MAL]	
C ₈ H ₁₄ O ₅	[na]	2-(lactyloxy)propionic acid, ethyl ester					
	$\Delta_v H$	(321–389)	72.8	336	A	[1987STE/MAL]	

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₈ H ₁₄ O ₅	[7554-12-3] $\Delta_v H$	malic acid, diethyl ester (353–527)	59.6	368	A	[1987STE/MAL, 1947STU]
C ₈ H ₁₄ O ₅	[na] $\Delta_v H$	propyl[1-(methoxycarbonyl)ethyl] carbonate (373–495)	58.0	388	A	[1987STE/MAL]
C ₈ H ₁₄ O ₆	[13811-71-7] $\Delta_v H$	(<i>d</i>) diethyl tartrate (375–553)	65.9	390		[1947STU]
C ₈ H ₁₄ O ₆	[87-91-2] $\Delta_v H$	(<i>dl</i>) diethyl tartrate (375–553)	67.3	390	A	[1987STE/MAL, 1947STU]
C ₈ H ₁₄ O ₆	$\Delta_v H$	(<i>d</i>)-dimethoxysuccinic acid dimethyl ester 53.1				[1937DUN/WOL]
C ₈ H ₁₄ O ₆	$\Delta_v H$	(<i>dl</i>)-dimethoxysuccinic acid dimethyl ester 57.7				[1937DUN/WOL]
C ₈ H ₁₄ O ₆	$\Delta_v H$	<i>meso</i> dimethoxysuccinic acid dimethyl ester 74.1				[1937DUN/WOL]
C ₈ H ₁₄ O ₆ S	[29771-87-7] $\Delta_v H$	sulfonyldiacetic acid, diethyl ester (421–494)	88.2	426		[1999DYK/SVO]
	$\Delta_v H$	(421–494)	87.6	436	A	[1987STE/MAL]
C ₈ H ₁₄ O ₆ S	[na] $\Delta_{\text{fus}} H$	dimethyl 3,3'-sulfonyldipropionate 41.1		390.3		[1994WAN/KUO]
C ₈ H ₁₅ Br	[1647-26-3] $\Delta_v H$	(2-bromoethyl)cyclohexane (311–486)	54.2	326	A	[1987STE/MAL, 1947STU, 1970DYK/VAN]
C ₈ H ₁₅ ClO	[111-64-8] $\Delta_v H$	octanoyl chloride (343–373)	74.5	358	A	[1987STE/MAL, 1999DYK/SVO]
C ₈ H ₁₅ ClO	[na] $\Delta_v H$	5-methylheptanoyl chloride (338–373)	66.3	353	A	[1987STE/MAL, 1999DYK/SVO]
C ₈ H ₁₅ Cl ₃ O ₄	[na] $\Delta_v H$	trichlorohydrine pentaerythritol (404–449)	80.4	419		[1965LUT/KOL]
C ₈ H ₁₅ N	[283-24-9] $\Delta_{\text{us}} H$	3-azabicyclo[3.2.2]nonane 14.55		297.8		
	$\Delta_{\text{fus}} H$	6.92		466.6		[1996DOM/HEA]
	$\Delta_{\text{sub}} H$	57.8 ± 1.3		298	C	[1970WES/WON]
	$\Delta_v H$	(303–443)	52.2	318	A	[1987STE/MAL]
C ₈ H ₁₅ N	[124-12-9] $\Delta_v H$	octanenitrile (283–310)	55.7 ± 0.2	298	GS	[2005EME/VER]
	$\Delta_v H$	(373–480)	50.0	388	A	[1987STE/MAL]
	$\Delta_v H$	56.8 ± 0.3		298	C	[1977STRI/SUN]
	$\Delta_v H$	(374–420)	49.8	389	EB	[1971MEY/REN]
	$\Delta_v H$	(420–479)	48.0	435	EB	[1971MEY/REN]
	$\Delta_v H$	(316–477)	56.7	331		[1947STU]
	$\Delta_v H$	(322–460)	53.5	298	EB	[1941RAL/SEL, 2005EME/VER]
	$\Delta_v H$	(294–477)	57.2 ± 0.3	298	MM	[1933HEI, 2005EME/VER]
C ₈ H ₁₅ NO	[4747-81-3] $\Delta_v H$	heptyl isocyanate (326–461)	47.5	341	A	[1987STE/MAL]
C ₈ H ₁₅ NO	[6554-73-0]	methacrylic acid <i>N-tert</i> -butylamide				

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(340–467)	49.6	355	A	[1987STE/MAL]
C ₈ H ₁₅ NO	[na]	<i>trans</i> 2-octenoic acid amide				
	$\Delta_{\text{sub}} H$	(373–393)	73.5	383	A	[1987STE/MAL]
C ₈ H ₁₅ NO ₂	[2867-47-2]	methacrylic acid, 2-(dimethylamino)ethyl ester				
	$\Delta_v H$	(372–460)	48.8	387	A	[1987STE/MAL]
C ₈ H ₁₅ NO ₂	[na]	1-lactopiperidine				
	$\Delta_v H$	(346–408)	62.1	361	A	[1987STE/MAL]
C ₈ H ₁₅ NO ₂	[1563-86-6]	N-acetyl-N-butylacetamide				
	$\Delta_v H$		64.4 ± 0.4	298	C	[1965WAD]
C ₈ H ₁₅ NO ₂	[na]	dimethylaminoethyl methacrylate				
	$\Delta_{\text{fus}} H$		16.85	237.7		[1996DOM/HEA]
C ₈ H ₁₅ NO ₃	[5411-58-5]	N,N-diethyloxamic acid, ethyl ester				
	$\Delta_v H$	(349–525)	60.5	364	A	[1987STE/MAL, 1947STU]
C ₈ H ₁₅ N ₅ O	[673-04-1]	2-methoxy-4,6- <i>bis</i> (ethylamino)-1,3,5-triazine				
	$\Delta_{\text{sub}} H$	(323–403)	98.2	338	GS-GC	[1987STE/MAL, 1964FRI/SAM]
C ₈ H ₁₅ N ₅ S	[1014-70-6]	2-methylthio-4,6- <i>bis</i> (ethylamino)-1,3,5-triazine (simetryn)				
	$\Delta_{\text{fus}} H$		24.0	353.2		[2007VEC/BRU]
	$\Delta_{\text{sub}} H$	(323–355)	101.3	338	GS-GC	[1987STE/MAL, 1964FRI/SAM]
	$\Delta_v H$		88 ± 4	461	DSC	[2007VEC/BRU]
	$\Delta_v H$		120 ± 6	298	DSC	[2007VEC/BRU]
	$\Delta_v H$		83.7 ± 1.3	453	TGA	[2007VEC/BRU]
	$\Delta_v H$		115 ± 4	298	DSC	[2007VEC/BRU]
C ₈ H ₁₅ N ₅ S	[1014-69-3]	2-methylthio-4-methylamino-6-isopropyl-1,3,5-triazine				
	$\Delta_{\text{sub}} H$	(323–357)	101.5	338	GS-GC	[1987STE/MAL, 1964FRI/SAM]
C ₈ H ₁₅ N ₇ O ₂ S ₃	[76824-35-6]	3-[[[2-[(aminoiminomethyl)amino]-4-thiazolyl]methyl]thio]-N-(aminosulfonyl)propanimidamide (famotidine)				
	$\Delta_{\text{fus}} H$ (I)		49.7	447		
	$\Delta_{\text{fus}} H$ (II)		48.6	438.6	DSC	[2007LU/WAN]
	$\Delta_{\text{fus}} H$ (I)		45.81	444.4		
	$\Delta_{\text{fus}} H$ (II)		43.92	436.6		[2002ROU/DAV]
	$\Delta_{\text{sub}} H$		207		TGA	[1997ELD]
C ₈ H ₁₆	[16747-50-5]	1-ethyl-1-methylcyclopentane				
	$\Delta_v H$		38.8	298	C	[1981HOS/SCO3]
C ₈ H ₁₆	[292-64-8]	cyclooctane				
	$\Delta_{\text{fus}} H$		6.32	166.5		
	$\Delta_{\text{fus}} H$		0.48	183.8		
	$\Delta_{\text{fus}} H$		2.41	288		[1991ACR]
	$\Delta_{\text{sub}} H$		58.7	166	B	[1963BON]
	$\Delta_v H$	(358–413)	40.3	373		[1991WU/LOC]
	$\Delta_v H$		43.1 ± 0.2		GC	[1989AZA]
	$\Delta_v H$	(289–369)	43.3	304	A	[1987STE/MAL]
	$\Delta_v H$	(373–434)	39.3	388	EB	[1976MEY/HOT]
	$\Delta_v H$	(291–323)	43.1	306		[1975ANA/GRO]
	$\Delta_v H$		43.3 ± 0.2	298		[1956FIN/SCO]
	$\Delta_v H$	(369–467)	39.4	384	A,EB	[1987STE/MAL, 1956FIN/SCO]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₈ H ₁₆	[590-66-9]	1,1-dimethylcyclohexane				
	$\Delta_{\text{fus}}H$		5.98	153.2		
	$\Delta_{\text{fus}}H$		2.01	239.8		[1996DOM/HEA]
	Δ_vH	(271–303)	39.6 ± 0.1	287	GS	[1995CHI/HES]
	Δ_vH		38.8 ± 0.1	298		[1995CHI/HES]
	Δ_vH		37.9	298		[1975KUS/SAI]
	Δ_vH	(313–395)	37.8	298		[1971WIL/ZWO]
			36.6	328	A	[1987STE/MAL, 1949FOR/NOR]
C ₈ H ₁₆	[2207-01-4]	<i>cis</i> 1,2-dimethylcyclohexane				
	$\Delta_{\text{fus}}H$		8.26	172.5		
	$\Delta_{\text{fus}}H$		1.64	223.3		[1996DOM/HEA]
	Δ_vH		39.4	298		[1975KUS/SAI]
	Δ_vH		39.7	298		[1971WIL/ZWO]
	Δ_vH		35.5 ± 0.1	370	C	[1951MCC/PER]
	Δ_vH		34.5 ± 0.1	387	C	[1951MCC/PER]
	Δ_vH	(322–405)	39.7 ± 0.1	298	C	[1947OSB/GIN]
			38.0	337	A,MM	[1987STE/MAL, 1945WIL/TAY]
C ₈ H ₁₆	[6876-23-9]	<i>(dl)</i> <i>trans</i> 1,2-dimethylcyclohexane				
	$\Delta_{\text{fus}}H$		10.5	185		[1996DOM/HEA]
	Δ_vH		38.3	298		[1975KUS/SAI]
	Δ_vH		38.4	298		[1971WIL/ZWO]
	Δ_vH		34.4 ± 0.1	373	C	[1951MCC/PER]
	Δ_vH		33.5 ± 0.1	387	C	[1951MCC/PER]
	Δ_vH	(316–399)	38.4 ± 0.1	298	C	[1947OSB/GIN]
			37.0	331	A,MM	[1987STE/MAL, 1945WIL/TAY]
C ₈ H ₁₆	[638-04-0]	<i>cis</i> 1,3-dimethylcyclohexane				
	$\Delta_{\text{fus}}H$		10.82	197.6		[1996DOM/HEA]
	Δ_vH	(318–396)	36.8	333	A	[1987STE/MAL]
	Δ_vH		38.1	298		[1975KUS/SAI]
	Δ_vH		38.2	298		[1971WIL/ZWO]
	Δ_vH		34.9 ± 0.1	363	C	[1951MCC/PER]
	Δ_vH		33.3 ± 0.1	385	C	[1951MCC/PER]
	Δ_vH	(316–398)	38.2 ± 0.1	298	C	[1947OSB/GIN]
			37.7	331	MM	[1945WIL/TAY]
C ₈ H ₁₆	[2207-03-6]	<i>(dl)</i> <i>trans</i> 1,3-dimethylcyclohexane				
	$\Delta_{\text{fus}}H$		9.87	183.1		[1996DOM/HEA]
	Δ_vH	(314–400)	37.9	329	A	[1987STE/MAL]
	Δ_vH		39.1	298		[1975KUS/SAI]
	Δ_vH		39.2	298		[1971WIL/ZWO]
	Δ_vH	(314–394)	39.2 ± 0.1	298	C	[1947OSB/GIN]
			37.4	329	MM	[1945WIL/TAY]
C ₈ H ₁₆	[624-29-3]	<i>cis</i> 1,4-dimethylcyclohexane				
	$\Delta_{\text{fus}}H$		9.29	185.7		[1996DOM/HEA]
	Δ_vH		39	298		[1975KUS/SAI]
	Δ_vH		39.00	298		[1971WIL/ZWO]
	Δ_vH	(317–400)	39.0 ± 0.1	298	C	[1947OSB/GIN]
			37.6	332	A,MM	[1987STE/MAL, 1945WIL/TAY]
C ₈ H ₁₆	[2207-04-7]	<i>(dl)</i> <i>trans</i> 1,4-dimethylcyclohexane				

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound					
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference	
		$\Delta_{\text{fus}}H$	12.34	236.2		[1996DOM/HEA]	
		Δ_vH	37.6	298		[1975KUS/SAI]	
		Δ_vH	37.9	298		[1971WIL/ZWO]	
		Δ_vH	35.6 ± 0.1	341	C	[1951MCC/PER]	
		Δ_vH	34.6 ± 0.1	357	C	[1951MCC/PER]	
		Δ_vH	33.5 ± 0.1	377	C	[1951MCC/PER]	
		Δ_vH	39.9 ± 0.1	298	C	[1947OSB/GIN]	
		Δ_vH	(313–395)	36.7	328	A,MM	[1987STE/MAL, 1945WIL/TAY]
C₈H₁₆	[1678-91-7]	ethylcyclohexane					
		$\Delta_{\text{fus}}H$	8.5	161.5		[2006MAN/CUT]	
		$\Delta_{\text{fus}}H$	8.33	161.4		[1996DOM/HEA]	
		Δ_vH	39.2 ± 0.4	298	GC	[1987AZA]	
		Δ_vH	39.8 ± 0.1	313	C	[1981SVO/CHA]	
		Δ_vH	38.9 ± 0.1	328	C	[1981SVO/CHA]	
		Δ_vH	37.9 ± 0.1	343	C	[1981SVO/CHA]	
		Δ_vH	37.0 ± 0.1	358	C	[1981SVO/CHA]	
		Δ_vH	36.3 ± 0.1	368	C	[1981SVO/CHA]	
		Δ_vH	40.0 ± 0.4	298	GCC	[1978FUC/PEA]	
		Δ_vH	40.4	298		[1975KUS/SAI]	
		Δ_vH	40.5	298		[1971WIL/ZWO]	
		Δ_vH	40.5 ± 0.1	298	C	[1947OSB/GIN]	
		Δ_vH	(323–407)	38.6	338	A,MM	[1987STE/MAL, 1945WIL/TAY]
C₈H₁₆	[2040-96-2]	propylcyclopentane					
		$\Delta_{\text{fus}}H$	10.04	155.8		[1996DOM/HEA]	
		Δ_vH	41.1	298		[1971WIL/ZWO]	
		Δ_vH	41.1 ± 0.1	298	C	[1947OSB/GIN]	
		Δ_vH	(323–406)	39.2	338	A,MM	[1987STE/MAL, 1945WIL/TAY]
C₈H₁₆	[3875-51-2]	isopropylcyclopentane					
		Δ_vH	37.9	298		[1971WIL/ZWO]	
		Δ_vH	39.4 ± 0.1	298	C	[1947OSB/GIN]	
		Δ_vH	(320–403)	37.9	335	A,MM	[1987STE/MAL, 1945WIL/TAY]
C₈H₁₆	[4259-00-1]	1,1,2-trimethylcyclopentane					
		Δ_vH	36.3	324	A,MM	[1987STE/MAL, 1949FOR/NOR]	
C₈H₁₆	[4516-69-2]	1,1,3-trimethylcyclopentane					
		Δ_vH	35.4	316	A,MM	[1987STE/MAL, 1949FOR/NOR]	
C₈H₁₆	[16747-50-5]	1-ethyl-1-methylcyclopentane					
		Δ_vH	(331–397)	36.7	346	A	[1987STE/MAL]
		Δ_vH	(238–288)	40.2	273	IP	[1987STE/MAL, 1974OSB/DOU]
		Δ_vH		38.9	298		[1971WIL/ZWO]
		Δ_vH	(316–396)	37.3	331		[1949FOR/NOR]
C₈H₁₆	[930-89-2]	<i>(dl)</i> <i>cis</i> 1-ethyl-2-methylcyclopentane					
		Δ_vH	(238–304)	42.5	253	A	[1987STE/MAL]
		Δ_vH	(303–403)	39.3	318	A	[1987STE/MAL]
		Δ_vH	(238–288)	41.6	273	IP	[1974OSB/DOU]
		Δ_vH		40.2	298		[1971WIL/ZWO]
		Δ_vH	(322–402)	38.3	337		[1949FOR/NOR]
C₈H₁₆	[na]	<i>trans</i> 1-ethyl-2-methylcyclopentane					
		Δ_vH		39.3	298		[1971WIL/ZWO]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₈ H ₁₆	[na]	<i>cis</i> 1-ethyl-3-methylcyclopentane				
	$\Delta_v H$		39.3	298		[1941KIR/SIT]
C ₈ H ₁₆	[na]	<i>trans</i> 1-ethyl-3-methylcyclopentane				
	$\Delta_v H$		38.9	298		[1941KIR/SIT]
C ₈ H ₁₆	[4259-00-1]	1,1,2-trimethylcyclopentane				
	$\Delta_v H$		37.2	298		[1971WIL/ZWO]
C ₈ H ₁₆	[4516-69-2]	1,1,3-trimethylcyclopentane				
	$\Delta_v H$		36.0	298		[1971WIL/ZWO]
C ₈ H ₁₆	[na]	<i>cis, cis</i> 1,2,3-trimethylcyclopentane				
	$\Delta_v H$		38.9	298		[1971WIL/ZWO]
C ₈ H ₁₆	[na]	<i>cis, cis</i> 1,2,4-trimethylcyclopentane				
	$\Delta_v H$		38.1	298		[1971WIL/ZWO]
C ₈ H ₁₆	[na]	<i>cis</i> 1,2- <i>trans</i> -3-trimethylcyclopentane				
	$\Delta_v H$		38.1	298		[1971WIL/ZWO]
C ₈ H ₁₆	[4850-28-6]	<i>cis</i> 1,2- <i>trans</i> -4-trimethylcyclopentane (311–392)				
	$\Delta_v H$		36.8	326	A,MM	[1987STE/MAL, 1949FOR/NOR]
C ₈ H ₁₆	[13398-35-1]	<i>trans</i> 1,2- <i>cis</i> -4-trimethylcyclopentane				
	$\Delta_v H$		36.8	298		[1971WIL/ZWO]
	$\Delta_v H$	(305–385)	36.0	320	A,MM	[1987STE/MAL, 1949FOR/NOR]
C ₈ H ₁₆	[111-66-0]	1-octene				
	$\Delta_{\text{fus}} H$		15.31	171.5		[1996DOM/HEA]
	$\Delta_v H$	(373–423)	40.6	298	CGC	[1995CHI/HOS]
	$\Delta_v H$		39.5 ± 0.1	313	C	[1982SVO/CHA]
	$\Delta_v H$		38.6 ± 0.1	328	C	[1982SVO/CHA]
	$\Delta_v H$		37.6 ± 0.1	343	C	[1982SVO/CHA]
	$\Delta_v H$		36.6 ± 0.1	358	C	[1982SVO/CHA]
	$\Delta_v H$		35.8 ± 0.1	368	C	[1982SVO/CHA]
	$\Delta_v H$	(263–291)	40.2	277	MM	[1981CHI/HYM]
	$\Delta_v H$	(260–291)	41.2	275	HSA	[1981CHI/HYM]
	$\Delta_v H$		40.3 ± 0.2	298	C	[1977MAN/SEL]
	$\Delta_v H$		38	298		[1971WIL/ZWO]
$\Delta_v H$	(317–395)	38.8	332	A,MM	[1987STE/MAL, 1950FOR/CAM]	
C ₈ H ₁₆	[7642-04-8]	<i>cis</i> 2-octene				
	$\Delta_v H$		40.2	298		[1971WIL/ZWO]
	$\Delta_v H$	(356–400)	37.8	371	A	[1987STE/MAL]
C ₈ H ₁₆	[13389-42-9]	<i>trans</i> 2-octene				
	$\Delta_v H$	(356–399)	37.9	371	A	[1987STE/MAL]
	$\Delta_v H$		40.2	298		[1971WIL/ZWO]
C ₈ H ₁₆	[14850-22-7]	<i>cis</i> 3-octene				
	$\Delta_v H$		39.7	298		[1971WIL/ZWO]
C ₈ H ₁₆	[14919-01-8]	<i>trans</i> 3-octene				
	$\Delta_v H$	(354–396)	37.6	369	A	[1987STE/MAL]
	$\Delta_v H$		40.2	298		[1971WIL/ZWO]
C ₈ H ₁₆	[7642-15-1]	<i>cis</i> 4-octene				
	$\Delta_v H$	(353–395)	37.2	368	A	[1987STE/MAL]
$\Delta_v H$		39.7	298		[1971WIL/ZWO]	

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound						
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference		
C ₈ H ₁₆	[14850-23-8]	<i>trans</i> 4-octene						
	$\Delta_v H$	(276–308)	43.2 ± 0.3	292	GS	[2000VER/WAN]		
	$\Delta_v H$	(276–308)	42.9 ± 0.3	298	GS	[2000VER/WAN]		
	$\Delta_v H$	(353–396)	37.4	368	A	[1987STE/MAL]		
C ₈ H ₁₆	[15870-10-7]	2-methyl-1-heptene	$\Delta_v H$	39.3	298	[1971WIL/ZWO]		
C ₈ H ₁₆	[4810-09-7]	3-methyl-1-heptene	$\Delta_v H$	38.5	298	[1971WIL/ZWO]		
C ₈ H ₁₆	[13151-05-8]	4-methyl-1-heptene	$\Delta_v H$	38.9	298	[1971WIL/ZWO]		
C ₈ H ₁₆	[13151-04-7]	5-methyl-1-heptene	$\Delta_v H$	38.9	298	[1971WIL/ZWO]		
C ₈ H ₁₆	[5026-76-6]	6-methyl-1-heptene	$\Delta_v H$	38.9	298	[1971WIL/ZWO]		
C ₈ H ₁₆	[627-97-4]	2-methyl-2-heptene	$\Delta_v H$	39.7	298	[1971WIL/ZWO]		
			$\Delta_v H$	(257–396)	41.2	272	A	[1987STE/MAL, 1947STU]
C ₈ H ₁₆	[22768-19-0]	3-methyl- <i>cis</i> -2-heptene	$\Delta_v H$	39.7	298	[1971WIL/ZWO]		
C ₈ H ₁₆	[22768-20-3]	3-methyl- <i>trans</i> -2-heptene	$\Delta_v H$	39.7	298	[1971WIL/ZWO]		
C ₈ H ₁₆	[na]	4-methyl- <i>cis</i> -2-heptene	$\Delta_v H$	38.9	298	[1971WIL/ZWO]		
C ₈ H ₁₆	[na]	4-methyl- <i>trans</i> -2-heptene	$\Delta_v H$	38.9	298	[1971WIL/ZWO]		
C ₈ H ₁₆	[24608-84-2]	5-methyl- <i>cis</i> -2-heptene	$\Delta_v H$	39.3	298	[1971WIL/ZWO]		
C ₈ H ₁₆	[24608-85-3]	5-methyl- <i>trans</i> -2-heptene	$\Delta_v H$	39.3	298	[1971WIL/ZWO]		
C ₈ H ₁₆	[na]	6-methyl- <i>cis</i> -2-heptene	$\Delta_v H$	39.3	298	[1971WIL/ZWO]		
C ₈ H ₁₆	[51065-65-7]	6-methyl- <i>trans</i> -2-heptene	$\Delta_v H$	39.3	298	[1971WIL/ZWO]		
C ₈ H ₁₆	[20488-34-0]	2-methyl- <i>cis</i> -3-heptene	$\Delta_v H$	38.9	298	[1971WIL/ZWO]		
C ₈ H ₁₆	[692-96-6]	2-methyl- <i>trans</i> -3-heptene	$\Delta_v H$	38.9	298	[1971WIL/ZWO]		
C ₈ H ₁₆	[22768-17-8]	3-methyl- <i>cis</i> -3-heptene	$\Delta_v H$	39.7	298	[1971WIL/ZWO]		
C ₈ H ₁₆	[22768-18-9]	3-methyl- <i>trans</i> -3-heptene	$\Delta_v H$	39.7	298	[1971WIL/ZWO]		
C ₈ H ₁₆	[14255-24-4]	4-methyl- <i>cis</i> -3-heptene	$\Delta_v H$	39.7	298	[1971WIL/ZWO]		

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₈ H ₁₆	[13714-85-7] $\Delta_v H$	4-methyl- <i>trans</i> -3-heptene	39.7	298		[1971WIL/ZWO]
C ₈ H ₁₆	[50422-80-5] $\Delta_v H$	5-methyl- <i>cis</i> -3-heptene	38.9	298		[1971WIL/ZWO]
C ₈ H ₁₆	[53510-18-2] $\Delta_v H$	5-methyl- <i>trans</i> -3-heptene	38.9	298		[1971WIL/ZWO]
C ₈ H ₁₆	[66225-19-2] $\Delta_v H$	6-methyl- <i>cis</i> -3-heptene	38.9	298		[1971WIL/ZWO]
C ₈ H ₁₆	[66225-20-5] $\Delta_v H$	6-methyl- <i>trans</i> -3-heptene	38.9	298		[1971WIL/ZWO]
C ₈ H ₁₆	[1632-16-2] $\Delta_v H$	2-ethyl-1-hexene	39.7	298		[1971WIL/ZWO]
C ₈ H ₁₆	[3404-58-8] $\Delta_v H$	3-ethyl-1-hexene	38.5	298		[1971WIL/ZWO]
C ₈ H ₁₆	[16746-85-3] $\Delta_v H$	4-ethyl-1-hexene	38.9	298		[1971WIL/ZWO]
C ₈ H ₁₆	[16746-86-4] $\Delta_v H$	2,3-dimethyl-1-hexene	38.5	298		[1971WIL/ZWO]
C ₈ H ₁₆	[16746-87-5] $\Delta_v H$	2,4-dimethyl-1-hexene	38.5	298		[1971WIL/ZWO]
C ₈ H ₁₆	[6795-92-4] $\Delta_v H$	2,5-dimethyl-1-hexene	38.9	298		[1971WIL/ZWO]
C ₈ H ₁₆	[3404-77-1] $\Delta_v H$	3,3-dimethyl-1-hexene	36.0	298		[1971WIL/ZWO]
C ₈ H ₁₆	[16745-94-1] $\Delta_v H$	3,4-dimethyl-1-hexene	38.9	298		[1971WIL/ZWO]
C ₈ H ₁₆	[7423-69-0] $\Delta_v H$	3,5-dimethyl-1-hexene	38.1	298		[1971WIL/ZWO]
C ₈ H ₁₆	[1647-08-1] $\Delta_v H$	4,4-dimethyl-1-hexene	31.0	298		[1971WIL/ZWO]
C ₈ H ₁₆	[16106-59-5] $\Delta_v H$	4,5-dimethyl-1-hexene	38.5	298		[1971WIL/ZWO]
C ₈ H ₁₆	[7116-86-1] $\Delta_v H$	5,5-dimethyl-1-hexene	37.7	298		[1971WIL/ZWO]
C ₈ H ₁₆	[36880-72-5] $\Delta_v H$	3-ethyl- <i>cis</i> -2-hexene	39.7	298		[1971WIL/ZWO]
C ₈ H ₁₆	[19781-63-6] $\Delta_v H$	3-ethyl- <i>trans</i> -2-hexene	39.7	298		[1971WIL/ZWO]
C ₈ H ₁₆	[54616-49-8] $\Delta_v H$	4-ethyl- <i>cis</i> -2-hexene	38.9	298		[1971WIL/ZWO]
C ₈ H ₁₆	[19781-63-6] $\Delta_v H$	4-ethyl- <i>trans</i> -2-hexene	38.9	298		[1971WIL/ZWO]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₈ H ₁₆	[7145-20-2] $\Delta_v H$	2,3-dimethyl-2-hexene	39.7	298		[1971WIL/ZWO]
C ₈ H ₁₆	[14255-23-3] $\Delta_v H$	2,4-dimethyl-2-hexene	38.5	298		[1971WIL/ZWO]
C ₈ H ₁₆	[3404-78-2] $\Delta_v H$	2,5-dimethyl-2-hexene	38.9	298		[1971WIL/ZWO]
C ₈ H ₁₆	[19550-81-3] $\Delta_v H$	3,4-dimethyl- <i>cis</i> -2-hexene	39.3	298		[1971WIL/ZWO]
C ₈ H ₁₆	[19550-82-4] $\Delta_v H$	3,4-dimethyl- <i>trans</i> -2-hexene	39.3	298		[1971WIL/ZWO]
C ₈ H ₁₆	[66225-31-8] $\Delta_v H$	3,5-dimethyl- <i>cis</i> -2-hexene	38.9	298		[1971WIL/ZWO]
C ₈ H ₁₆	[66225-12-5] $\Delta_v H$	3,5-dimethyl- <i>trans</i> -2-hexene	38.9	298		[1971WIL/ZWO]
C ₈ H ₁₆	[66225-13-6] $\Delta_v H$	4,4-dimethyl- <i>cis</i> -2-hexene	38.1	298		[1971WIL/ZWO]
C ₈ H ₁₆	[19550-83-5] $\Delta_v H$	4,4-dimethyl- <i>trans</i> -2-hexene	38.1	298		[1971WIL/ZWO]
C ₈ H ₁₆	[na] $\Delta_v H$	4,5-dimethyl- <i>cis</i> -2-hexene	38.5	298		[1971WIL/ZWO]
C ₈ H ₁₆	[66225-14-7] $\Delta_v H$	4,5-dimethyl- <i>trans</i> -2-hexene	38.5	298		[1971WIL/ZWO]
C ₈ H ₁₆	[39761-61-0] $\Delta_v H$	5,5-dimethyl- <i>cis</i> -2-hexene	38.1	298		[1971WIL/ZWO]
C ₈ H ₁₆	[39782-43-9] $\Delta_v H$	5,5-dimethyl- <i>trans</i> -2-hexene	38.1	298		[1971WIL/ZWO]
C ₈ H ₁₆	[16789-51-8] $\Delta_v H$	3-ethyl-3-hexene	39.3	298		[1971WIL/ZWO]
C ₈ H ₁₆	[690-92-6] $\Delta_v H$ $\Delta_v H$ $\Delta_v H$	<i>cis</i> 2,2-dimethyl-3-hexene (319–380) 35.3 37.2 (305–379) 36.1	334 298 320	A MM	[1987STE/MAL] [1971WIL/ZWO] [1960CAM/ROS]	
C ₈ H ₁₆	[690-93-7] $\Delta_v H$ $\Delta_v H$ $\Delta_v H$	<i>trans</i> 2,2-dimethyl-3-hexene (306–379) 36.1 37.2 (303–374) 36.3	321 298 318	A MM	[1987STE/MAL] [1971WIL/ZWO] [1960CAM/ROS]	
C ₈ H ₁₆	[59643-75-3] $\Delta_v H$	<i>cis</i> 2,3-dimethyl-3-hexene	38.9	298		[1971WIL/ZWO]
C ₈ H ₁₆	[66225-30-7] $\Delta_v H$	<i>trans</i> 2,3-dimethyl-3-hexene	38.9	298		[1971WIL/ZWO]
C ₈ H ₁₆	[37549-89-6] $\Delta_v H$	<i>cis</i> 2,4-dimethyl-3-hexene	38.5	298		[1971WIL/ZWO]
C ₈ H ₁₆	[61847-78-7] $\Delta_v H$	<i>trans</i> 2,4-dimethyl-3-hexene	38.5	298		[1971WIL/ZWO]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₈ H ₁₆	[10557-44-5] $\Delta_v H$	<i>cis</i> 2,5-dimethyl-3-hexene	37.2	298		[1971WIL/ZWO]
C ₈ H ₁₆	[692-70-6] $\Delta_v H$	<i>trans</i> 2,5-dimethyl-3-hexene	37.5	298		[1971WIL/ZWO]
C ₈ H ₁₆	[19550-87-9] $\Delta_v H$	<i>cis</i> 3,4-dimethyl-3-hexene	39.7	298		[1971WIL/ZWO]
C ₈ H ₁₆	[19550-88-0] $\Delta_v H$	<i>trans</i> 3,4-dimethyl-3-hexene	39.7	298		[1971WIL/ZWO]
C ₈ H ₁₆	[15918-08-8] $\Delta_v H$	2-propyl-1-pentene	39.3	298		[1971WIL/ZWO]
C ₈ H ₁₆	[61847-79-8] $\Delta_v H$	2-isopropyl-1-pentene	38.8	298		[1971WIL/ZWO]
C ₈ H ₁₆	[3404-67-9] $\Delta_v H$ $\Delta_v H$ $\Delta_v H$	2-ethyl-3-methyl-1-pentene (307–389)	36.4 38.9	322 298	A	[1987STE/MAL] [1971WIL/ZWO]
		(308–383)	36.4	323	MM	[1960CAM/ROS]
C ₈ H ₁₆	[3404-80-6] $\Delta_v H$	2-ethyl-4-methyl-1-pentene	38.5	298		[1971WIL/ZWO]
C ₈ H ₁₆	[19780-66-6] $\Delta_v H$	3-ethyl-2-methyl-1-pentene	37.7	298		[1971WIL/ZWO]
C ₈ H ₁₆	[6196-60-7] $\Delta_v H$	3-ethyl-3-methyl-1-pentene	38.9	298		[1971WIL/ZWO]
C ₈ H ₁₆	[61847-80-1] $\Delta_v H$	3-ethyl-4-methyl-1-pentene	38.5	298		[1971WIL/ZWO]
C ₈ H ₁₆	[560-23-6] $\Delta_v H$	2,3,3-trimethyl-1-pentene	38.5	298		[1971WIL/ZWO]
C ₈ H ₁₆	[565-76-4] $\Delta_v H$	2,3,4-trimethyl-1-pentene	38.5	298		[1971WIL/ZWO]
C ₈ H ₁₆	[107-39-1] $\Delta_{\text{fus}} H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$	2,4,4-trimethyl-1-pentene	8.79	178.9		[1996DOM/HEA]
		(306–356)	35.7	298	EB	[2007MAL]
		(343–381)	33.5	358	A	[1987STE/MAL]
			35.7	298		[1971WIL/ZWO]
		(301–375)	35.1	316	MM	[1960CAM/ROS]
C ₈ H ₁₆	[564-03-4] $\Delta_v H$	3,3,4-trimethyl-1-pentene	38.1	298		[1971WIL/ZWO]
C ₈ H ₁₆	[19780-67-7] $\Delta_v H$	2-methyl-3-ethyl-2-pentene	39.3	298		[1971WIL/ZWO]
C ₈ H ₁₆	[42067-48-1] $\Delta_v H$	4-methyl-3-ethyl- <i>cis</i> -2-pentene	39.3	298		[1971WIL/ZWO]
C ₈ H ₁₆	[42067-49-2] $\Delta_v H$	4-methyl-3-ethyl- <i>trans</i> -2-pentene	38.9	298		[1971WIL/ZWO]
C ₈ H ₁₆	[565-77-5] $\Delta_v H$	2,3,4-trimethyl-2-pentene	39.3	298		[1971WIL/ZWO]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₈ H ₁₆	[107-40-4]	2,4,4-trimethyl-2-pentene				
	$\Delta_{\text{fus}}H$		6.78	166		[1996DOM/HEA]
	Δ_vH	(319–380)	35.7	334	A	[1987STE/MAL]
	Δ_vH	(305–378)	39.3	298		[1971WIL/ZWO]
C ₈ H ₁₆	[39761-64-3]	3,4,4-trimethyl- <i>cis</i> -2-pentene				
	Δ_vH		38.9	298		[1971WIL/ZWO]
C ₈ H ₁₆	[39761-64-3]	3,4,4-trimethyl- <i>trans</i> -2-pentene				
	Δ_vH		38.9	298		[1971WIL/ZWO]
C ₈ H ₁₆	[na]	3-methyl-2-isopropyl-1-butene				
	Δ_vH		38.1	298		[1971WIL/ZWO]
C ₈ H ₁₆	[18231-53-3]	3,3-dimethyl-2-ethyl-1-butene				
	Δ_vH		38.5	298		[1971WIL/ZWO]
C ₈ H ₁₆ Br ₂	[62168-26-7]	1,1-dibromooctane				
	Δ_vH	(412–571)	57.1	427	A,EST	[1987STE/MAL, 1956MAN, 1970DYK/VAN]
C ₈ H ₁₆ Cl ₂	[20395-24-8]	1,1-dichlorooctane				
	Δ_vH	(380–480)	57.7	298	A	[1987VAR/LOS2, 1991BAS/SVO]
	Δ_vH	(382–533)	51.4	397	A,EST	[1987STE/MAL, 1956MAN, 1970DYK/VAN, 1999DYK/SVO]
C ₈ H ₁₆ Cl ₂	[21948-46-9]	1,2-dichlorooctane				
	Δ_vH	(370–490)	52.0	385		[1982VAR/PUC, 1999DYK/SVO]
	Δ_vH	(370–490)	57.6	298		[1982VAR/PUC, 1992LEE/CHE]
C ₈ H ₁₆ Cl ₂	[2162-99-4]	1,8-dichlorooctane				
	Δ_vH	(410–510)	55.9	426		[1999DYK/SVO]
	Δ_vH	(410–510)	65.6	298		[1988VAR/LOS, 1991BAS/SVO]
C ₈ H ₁₆ Cl ₂	[2162-99-4]	<i>erythro</i> -4,5-dichlorooctane				
	Δ_vH	(351–480)	47.3	415		[1999DYK/SVO]
C ₈ H ₁₆ F ₂	[61350-03-6]	1,1-difluorooctane				
	Δ_vH	(329–459)	44.2	344	A,EST	[1987STE/MAL, 1956MAN, 1970DYK/VAN, 1999DYK/SVO]
C ₈ H ₁₆ NO ₂	[na]	ethyl 2-(<i>N,N</i> -dimethylamino)-2-methylpropanoate				
	Δ_vH	(278–313)	51.6 ± 0.5	298	GS	[1996VER/ZUF]
C ₈ H ₁₆ N ₂	[na]	methyl ethyl ketazine				
	Δ_vH	(439–524)	40.0			[1993FER/MOR]
C ₈ H ₁₆ N ₂	[na]	1,1,4,4-tetramethyltetramethylenediazine				
	Δ_vH		50.1 ± 0.4	298	C	[1976ENG/MEL]
C ₈ H ₁₆ N ₂	[na]	2-diethylamino-2-methylpropionitrile				
	Δ_vH		56.3 ± 1.1		GS	[1997WEL/VER]
C ₈ H ₁₆ N ₂ O ₂	[28529-34-2]	N-acetyl (<i>l</i>)-leucine amide				
	$\Delta_{\text{sub}}H$		115.6 ± 1.4	376	C	[1999DEL/BAR]
	$\Delta_{\text{sub}}H$		119.8 ± 1.5	298		[1999DEL/BAR]
C ₈ H ₁₆ N ₂ O ₂	[16624-68-3]	N-acetyl (<i>d</i>)-leucine amide				
	$\Delta_{\text{fus}}H$		20.2	404		[1996DOM/HEA]
	$\Delta_{\text{sub}}H$		114.8 ± 0.3	393	C	[1999DEL/BAR]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{sub}}H$		120.4 ± 0.4	298		[1999DEL/BAR]
	$\Delta_{\text{sub}}H$	(374–401)	101 ± 3	388	TE	[1988FER/DEL, 1986BAR/FER]
C ₈ H ₁₆ N ₂ O ₂	[56711-06-9]	N-acetyl (<i>l</i>)-isoleucine amide				
	$\Delta_{\text{fus}}H$		41.8	529.6		[1997PUL/DES]
	$\Delta_{\text{sub}}H$		142.7 ± 0.2	390	C	[1999DEL/BAR]
	$\Delta_{\text{sub}}H$		147.4 ± 0.3	298		[1999DEL/BAR]
C ₈ H ₁₆ N ₂ O ₂	[3891-73-4]	suberamide				
	$\Delta_{\text{us}}H$		5.45	431.3		
	$\Delta_{\text{fus}}H$		58.4	493.2	DSC	[2006BAD/DEL]
C ₈ H ₁₆ N ₆	[na]	1-(methylamino)-3,5-bis(dimethylamino)-s-triazine				
	$\Delta_{\text{fus}}H$		22.34	378.8		[1989BRA/RYT]
C ₈ H ₁₆ N ₆ O	[na]	1-(hydroxylamino)-3,5-bis(dimethylamino)-s-triazine				
	$\Delta_{\text{fus}}H$		30.67	381.5		[1989BRA/RYT]
C ₈ H ₁₆ O	[124-13-0]	octanal				
	$\Delta_{\text{fus}}H$		25.86	288.2		[1980DYA/VAS]
	Δ_vH	(277–310)	51.0 ± 0.3	298	GS	[2003VER/KRA2]
	Δ_vH	(313–353)	53.8	298	CGC	[1995CHI/HOS]
	Δ_vH	(293–438)	43.4	308	A	[1987STE/MAL]
	Δ_vH		51.3 ± 0.2	298		[1981DYA/KOR]
C ₈ H ₁₆ O	[18641-70-8]	2,4-dimethyl-3-hexanone				
	Δ_vH	(350–418)	42.5	365	A	[1987STE/MAL]
C ₈ H ₁₆ O	[696-71-9]	cyclooctanol				
	$\Delta_{\text{fus}}H$		2.02	291.2	DSC	[2008SIN/MUR]
	$\Delta_{\text{us}}H$		2.05	264.1		
	$\Delta_{\text{fus}}H$		1.97	297.1		[2003RUT/SAL]
	$\Delta_{\text{us}}H$		2.12	261.3		
	$\Delta_{\text{fus}}H$		2.06	295		[1995SCI/MAY]
Note: Authors did not report enthalpic data for all transitions						
C ₈ H ₁₆ O	[1940-18-7]	1-ethylcyclohexanol				
	Δ_vH	(324–440)	46.9	339	A	[1987STE/MAL]
C ₈ H ₁₆ O	[13019-20-0]	2-methyl-3-heptanone				
	Δ_vH	(350–428)	43.5	365	A	[1987STE/MAL]
C ₈ H ₁₆ O	[51500-48-2]	6-methyl-3-hepten-2-ol				
	Δ_vH	(314–449)	59.7	329	A	[1987STE/MAL, 1947STU]
C ₈ H ₁₆ O	[4630-06-2]	<i>(dl)</i> 6-methyl-5-hepten-2-ol				
	Δ_vH	(314–448)	57	329	A	[1987STE/MAL, 1947STU]
C ₈ H ₁₆ O	[111-13-7]	2-octanone				
	$\Delta_{\text{fus}}H$		24.42	252.9		[1996DOM/HEA]
	Δ_vH	(343–383)	52.6	298	CGC	[1995CHI/HOS]
	Δ_vH	(317–446)	49.8	332	A	[1987STE/MAL]
	Δ_vH		52.0 ± 0.3	298	GCC	[1979SAL/PEA]
	Δ_vH	(324–520)	49.1	339	A	[1987STE/MAL, 1975AMB/ELL]
	Δ_vH	(324–520)	51.8	298		[1975AMB/ELL]
	Δ_vH	(296–446)	50.6	311		[1947STU]
C ₈ H ₁₆ O	[106-68-3]	3-octanone				
	Δ_vH	(293–348)	43.8	308	A	[1987STE/MAL]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₈ H ₁₆ O	[589-63-9]	4-octanone				
	$\Delta_v H$	(288–433)	36.4	303	A	[1987STE/MAL]
C ₈ H ₁₆ O	[1604-02-0]	1-propylcyclopentanol				
	$\Delta_v H$	(344–447)	64.2	359	A	[1987STE/MAL]
C ₈ H ₁₆ O	[5857-36-2]	2,2,4-trimethyl-3-pentanone				
	$\Delta_v H$	(287–408)	55.7	302	A	[1987STE/MAL, 1947STU]
	$\Delta_v H$		43.3 ± 0.2	298	C	[1970SEL2]
	$\Delta_v H$		43.3 ± 0.1	298	C	[1966WAD]
C ₈ H ₁₆ O ₂	[na]	3-butoxy-2-butanone				
	$\Delta_v H$	(323–398)	36.7	338	A	[1987STE/MAL]
C ₈ H ₁₆ O ₂	[na]	<i>trans</i> 2,2,4,6-tetramethyl-1,3-dioxane				
	$\Delta_v H$		41.9 ± 1.2	298		[1967PIH/HEI]
C ₈ H ₁₆ O ₂	[na]	<i>cis</i> 2,2,4,6-tetramethyl-1,3-dioxane				
	$\Delta_v H$		42.3 ± 1.2	298		[1967PIH/HEI]
C ₈ H ₁₆ O ₂	[na]	2,2,6,6-tetramethyl-1,3-dioxane				
	$\Delta_{\text{fus}} H$		10.9	250.6		[1975BOR]
C ₈ H ₁₆ O ₂	[933-40-4]	1,1-dimethoxycyclohexane				
	$\Delta_v H$	(278–308)	48.6 ± 0.2	298	GS	[2002VER]
	$\Delta_v H$	(278–308)	49.0 ± 0.2		GS	[1998VER/PEN]
	$\Delta_v H$	(315–347)	52.4	331	EB	[1994WIB/MOR]
C ₈ H ₁₆ O ₂	[124-07-2]	octanoic acid (caprylic acid)				
	$\Delta_{\text{fus}} H$		21.38	289.7		[1996DOM/HEA]
	$\Delta_{\text{sub}} H$		113.3 ± 6	298	TPD	[2008CAP/LOV]
	$\Delta_v H$	(297–343)	79.8 ± 0.6	320	GS	[2000VER2]
	$\Delta_v H$	(297–434)	81.0 ± 0.6	298	GS	[2000VER2]
	$\Delta_v H$	(353–393)	81.2	298	CGC	[1995CHI/HOS]
	$\Delta_v H$	(417–514)	66.6	432	A,EB	[1987AMB/GHI3]
	$\Delta_v H$	(296–331)	85.3	311	A	[1987STE/MAL]
	$\Delta_v H$	(360–512)	74.4	375	A	[1987STE/MAL]
	$\Delta_v H$		80.0	290		[1982DEK/SCH]
	$\Delta_v H$	(291–303)	82.9 ± 1.0	298	TE	[1979DEK/OON]
	$\Delta_v H$		70.0	407	I	[1943CRA]
C ₈ H ₁₆ O ₂	[123-66-0]	ethyl hexanoate				
	$\Delta_v H$	(345–374)	47.4 ± 0.3	359	EB	[1991WIB/WAL]
	$\Delta_v H$	(345–379)	51.5 ± 1.3	298	EB	[1991WIB/WAL]
	$\Delta_v H$	(396–449)	51.8	311	A	[1987STE/MAL]
	$\Delta_v H$	(300–376)	48.6	315	A	[1987STE/MAL]
C ₈ H ₁₆ O ₂	[149-57-5]	(<i>dl</i>) 2-ethylhexanoic acid				
	$\Delta_v H$	(397–514)	76.3 ± 0.9	298	EB	[1997STE/CHI3]
	$\Delta_v H$	(403–500)	61.8	418	A	[1987STE/MAL]
	$\Delta_v H$		75.6 ± 0.5	298	C	[1976STR]
C ₈ H ₁₆ O ₂	[142-92-7]	hexyl acetate				
	$\Delta_{\text{fus}} H$		19.83	212.1		[1996DOM/HEA]
	$\Delta_v H$	(274–309)	51.9 ± 0.3	298	GS	[2006KRA/VER]
	$\Delta_v H$		52.1	298	GC	[1997KOU/HOS]
	$\Delta_v H$	(303–444)	50.9	318		[1995ARC/BLA]
	$\Delta_v H$	(304–381)	48.9	319	A	[1987STE/MAL]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(378–459)	46.2	387	DTA	[1980MEY/AWE]
C ₈ H ₁₆ O ₂	[109-21-7]	butyl butanoate				
	$\Delta_{\text{fus}} H$		14.93	181.7		[1996DOM/HEA]
C ₈ H ₁₆ O ₂	[539-90-2]	isobutyl butyrate				
	$\Delta_v H$	(277–430)	41.7	292	A	[1987STE/MAL, 1947STU]
C ₈ H ₁₆ O ₂	[97-85-8]	isobutyl isobutyrate				
	$\Delta_v H$	(274–319)	48.5	298	GS	[2008VER/EME]
	$\Delta_v H$	(278–313)	44.5 ± 0.1	298	GS	[1996VER/BEC]
	$\Delta_v H$	(277–421)	46.9	292	A	[1987STE/MAL, 1947STU]
C ₈ H ₁₆ O ₂	[624-54-4]	isopentyl propionate				
	$\Delta_v H$	(281–434)	44.1	296	A	[1987STE/MAL]
C ₈ H ₁₆ O ₂	[106-73-0]	methyl heptanoate				
	$\Delta_v H$	(421–444)	46.3	433		[2009POS/MAR]
	$\Delta_v H$	(278–310)	53.2 ± 0.2	298	GS	[2008VER/EME]
	$\Delta_v H$		49.1	350		[2002VAN/VAN]
	$\Delta_v H$		50.2 ± 0.1	326		[2002VAN/VAN]
	$\Delta_v H$		51.8 ± 0.1	298		[2002VAN/VAN]
	$\Delta_v H$	(313–363)	53.4	298	CGC	[1995CHI/HOS]
	$\Delta_v H$	(433–473)	53.7	298	CGC	[1995CHI/HOS]
	$\Delta_v H$	(313–353)	53.5	298	CGC	[1995CHI/HOS]
	$\Delta_v H$		49.7 ± 0.5	298	GC	[1987AZA]
	$\Delta_v H$		53.1 ± 0.4	298	GCC	[1980FUC/PEA]
	$\Delta_v H$		53.1 ± 0.1	298	C	[1980FUC/PEA]
	$\Delta_v H$		51.6 ± 0.5	298	C	[1977MAN/SEL]
	$\Delta_v H$	(332–402)	49.0	347	A,EST	[1987STE/MAL, 1963ROS/SCH]
C ₈ H ₁₆ O ₂	[25415-67-2]	4-methylvaleric acid, ethyl ester				
	$\Delta_v H$	(284–434)	45.4	299	A	[1987STE/MAL, 1947STU]
C ₈ H ₁₆ O ₂	[557-00-6]	propyl isovalerate				
	$\Delta_v H$	(281–429)	44.3	296	A	[1987STE/MAL, 1947STU]
C ₈ H ₁₆ O ₂	[123-66-0]	ethyl hexanoate				
	$\Delta_v H$	(279–309)	50.8 ± 0.4	294	GS	[1999VER/HEI]
	$\Delta_v H$	(279–309)	50.6 ± 0.4	298	GS	[1999VER/HEI]
C ₈ H ₁₆ O ₂	[34949-22-9]	<i>tert</i> -pentyl propionate				
	$\Delta_v H$	(274–310)	45.3 ± 0.1	298	GS	[2008VER/EME]
	$\Delta_v H$	(333–378)	45.7	298	CGC	[1999VER/HEI]
C ₈ H ₁₆ O ₂	[34859-98-8]	1,1-dimethylbutyl acetate				
	$\Delta_v H$	(333–378)	45.6	298	CGC	[1999VER/HEI]
C ₈ H ₁₆ O ₂	[15965-97-6]	[(3-methylbutoxy)methyl]oxirane				
	$\Delta_v H$		55.8 ± 1.9			[1987VAN/KAC]
C ₈ H ₁₆ O ₃	[112-07-2]	2-butoxyethyl acetate				
	$\Delta_v H$		59.5 ± 0.1	298	C	[1970KUS/WAD]
C ₈ H ₁₆ O ₃	[4126-55-0]	2-butoxypropionic acid, methyl ester				
	$\Delta_v H$	(348–417)	51.9	363	A	[1987STE/MAL]
C ₈ H ₁₆ O ₃	[14144-34-4]	3-butoxypropionic acid, methyl ester				
	$\Delta_v H$	(311–469)	51.1	326	A	[1987STE/MAL]
C ₈ H ₁₆ O ₃	[na]	3-ethoxypropionic acid, propyl ester				

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(343–461)	48.6	358	A	[1987STE/MAL, 1954FRI/PIC]
C ₈ H ₁₆ O ₃	[112-07-2] $\Delta_v H$	ethylene glycol monobutyl ether acetate (293–465)	51.9	308	A	[1987STE/MAL]
C ₈ H ₁₆ O ₃	[816-50-2] $\Delta_v H$	2-hydroxyisobutyric acid, butyl ester (384–458)	47.7	399	A	[1987STE/MAL]
C ₈ H ₁₆ O ₃	[4195-88-4] $\Delta_v H$	3-methoxypropionic acid, butyl ester (311–469)	50.9	326	A	[1987STE/MAL]
C ₈ H ₁₆ O ₃	[6382-06-5] $\Delta_v H$	pentyl lactate (288–469)	73.9	303	A	[1987STE/MAL]
C ₈ H ₁₆ O ₃	[112-15-2] $\Delta_v H$	diethylene glycol monoethyl ether acetate (293–491)	51.7	308	A	[1987STE/MAL]
C ₈ H ₁₆ O ₃	[2305-25-1] $\Delta_v H$	ethyl 3-hydroxyhexanoate (363–393)	61.9 ± 0.6	298	CGC	[2005TEM/CHI]
C ₈ H ₁₆ O ₄	[294-93-9] $\Delta_{\text{fus}} H$	1,4,7,10-tetraoxacyclododecane (12-crown-4)	22.46	290.7		[1998DOM]
	$\Delta_v H$		65.7 ± 3.7	298	CGC	[2000NIC/ORF]
	$\Delta_v H$		65.6 ± 0.4	298	C	[1982BYS/MAN]
C ₈ H ₁₆ O ₄	[2780-59-8] $\Delta_v H$	3,6-diethyl-3,6-dimethyl-1,2,4,5-tetraoxacyclohexane (403–473)	45.09	298	CGC	[2007CAN/EYL]
C ₈ H ₁₆ S ₄	[25423-56-7] $\Delta_{\text{fus}} H$	1,4,7,10-tetrathiacyclododecane	31.0	499.2	DSC	[2002ROC/GRI]
C ₈ H ₁₇ Br	[111-83-1] $\Delta_{\text{fus}} H$	1-bromooctane	24.69	218.2		[1950CRO/SMY]
	$\Delta_v H$	(323–363)	55.1	298	CGC	[1995CHI/HOS]
	$\Delta_v H$		55.8 ± 0.1	298	C	[1976STR3, 1977MAN/SEL]
	$\Delta_v H$	(373–475)	49.3	388	A,EST	[1987STE/MAL, 1961LI/ROS, 1970DYK/VAN]
C ₈ H ₁₇ Br	[555-35-7] $\Delta_v H$	(dl) 2-bromooctane (343–463)	48.4	358	A	[1987STE/MAL, 1999DYK/SVO]
C ₈ H ₁₇ Cl	[111-85-3] $\Delta_v H$	1-chlorooctane (330–460)	51.4	298		[1984BOU/FRI, 1991BAS/SVO]
	$\Delta_v H$	(327–457)	50.3	342	A,DTA	[1987STE/MAL, 1969KEM/KRE]
	$\Delta_v H$		52.4 ± 0.1	298	C	[1968WAD]
C ₈ H ₁₇ Cl	[628-61-5] $\Delta_v H$	(dl) 2-chlorooctane (330–446)	47.8	345	A	[1987STE/MAL, 1999DYK/SVO]
C ₈ H ₁₇ Cl	[123-04-6] $\Delta_v H$	(3-chloromethyl)heptane (371–443)	44.2	386	A	[1987STE/MAL, 1999DYK/SVO]
C ₈ H ₁₇ ClO ₄	[5197-66-0] $\Delta_v H$	triethylene glycol mono(2-chloroethyl) ether (383–555)	68.6	398	A	[1987STE/MAL, 1947STU]
C ₈ H ₁₇ Cl ₂ N	[42520-97-8] $\Delta_v H$	N-butyl bis(2-chloroethyl)amine (273–380)	60.7	288	A,GS	[1987STE/MAL, 1948RED/CHA3, 1999DYK/SVO]
C ₈ H ₁₇ Cl ₂ N	[na] $\Delta_v H$	N-sec-butyl bis(2-chloromethyl)amine (273–373)	59.5	288	GS	[1987STE/MAL, 1948RED/CHA3, 1999DYK/SVO]
C ₈ H ₁₇ Cl ₂ N	[10125-86-7]	N-tert-butyl bis(2-chloromethyl)amine				

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(273–345)	58.4	288	A,GS	[1987STE/MAL, 1948RED/CHA3, 1999DYK/SVO]
C ₈ H ₁₇ Cl ₂ N	[87289-70-1]	N-isobutyl bis(2-chloromethylamine)				
	$\Delta_v H$	(273–345)	60.3	288	A,GS	[1987STE/MAL, 1948RED/CHA3, 1999DYK/SVO]
C ₈ H ₁₇ F	[463-11-6]	1-fluorooctane				
	$\Delta_v H$		49.7	298		[UR/FUC, 1985MAJ/SVO]
	$\Delta_v H$	(307–446)	44.1	322	A,EST	[1987STE/MAL, 1961LI/ROS, 1970DYK/VAN]
C ₈ H ₁₇ I	[629-27-6]	1-iodooctane				
	$\Delta_v H$	(391–554)	59.7	298	A,EST	[1987STE/MAL, 1961LI/ROS, 1970DYK/VAN, 2006BOL/NER]
	$\Delta_v H$	(391–554)	50.7	406	A,EST	[1987STE/MAL, 1961LI/ROS, 1970DYK/VAN]
C ₈ H ₁₇ NO	[629-01-6]	octanamide				
	$\Delta_{\text{us}}H$		1.91	194.4		
	$\Delta_{\text{us}}H$		0.97	304.5		
	$\Delta_{\text{fus}}H$		27.6	377	DSC	[2008ABA/BAD]
	$\Delta_{\text{sub}}H$	(325–374)	110.5 ± 2.9		GS,ME	[1959DAV/JON2, 1987STE/MAL]
C ₈ H ₁₇ NO	[na]	N-propylpiperidine				
	$\Delta_v H$	(275–314)	45.2 ± 0.4	294	GS	[1998VER6]
	$\Delta_v H$	(275–314)	44.9 ± 0.4	298	GS	[1998VER6]
C ₈ H ₁₇ NO	[1114-76-7]	butyric acid N,N-diethylamide				
	$\Delta_v H$	(298–373)	38.7	313	A	[1987STE/MAL]
C ₈ H ₁₇ NO	[929-55-5]	caprylaldehyde oxime				
	$\Delta_v H$	(313–400)	71.3	328	A	[1987STE/MAL]
C ₈ H ₁₇ NO	[7207-49-0]	2-octanone oxime				
	$\Delta_v H$	(293–487)	67.5	308	A	[1987STE/MAL]
C ₈ H ₁₇ NO	[7207-50-3]	3-octanone oxime				
	$\Delta_v H$	(293–400)	67.2	308	A	[1987STE/MAL]
C ₈ H ₁₇ NO	[7207-51-4]	4-octanone oxime				
	$\Delta_v H$	(293–400)	68.8	308	A	[1987STE/MAL]
C ₈ H ₁₇ NO ₂	[na]	2,4,4-trimethyl-2-nitropentane				
	$\Delta_v H$	(288–324)	54.2 ± 0.8	306	GS	[1997VER3]
	$\Delta_v H$	(288–324)	54.7 ± 0.8	298	GS	[1997VER3]
C ₈ H ₁₇ NO ₂	[na]	lactic acid N-isopentylamide				
	$\Delta_v H$	(386–433)	77.9	401	A	[1987STE/MAL]
C ₈ H ₁₇ NO ₂	[na]	lactic acid N-pentylamide				
	$\Delta_v H$	(373–448)	81.8	388	A	[1987STE/MAL]
C ₈ H ₁₇ NO ₂	[2743-60-4]	(l) leucine ethyl ester				
	$\Delta_v H$	(333–449)	43.5	348	A	[1987STE/MAL]
C ₈ H ₁₇ NO ₂	[na]	ethyl 2-(N,N-dimethylamino)-2-methylpropionate				
	$\Delta_v H$		55.6 ± 0.4	283	DSC	[1993SCH/BEC]
C ₈ H ₁₇ NO ₂	[na]	(1-methylheptyl)nitrite				
	$\Delta_v H$	(303–338)	44.9	318	A	[1987STE/MAL]
C ₈ H ₁₇ NO ₂	[1002-57-9]	8-aminooctanoic acid				
	$\Delta_{\text{sub}}H$	(391–402)	166.2 ± 0.9	397	C	[1983SKO/SAB]
	$\Delta_{\text{sub}}H$		170 ± 4	298	C	[1983SKO/SAB]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₈ H ₁₇ NO ₄	[72458-42-5]	N-ethyl-5-amino-1,5-dideoxy-(<i>d</i>)-glycopyranose				
	$\Delta_{\text{fus}}H$		26.3	429.1		[1994BLU/PRA]
C ₈ H ₁₈	[590-73-8]	2,2-dimethylhexane				
	Δ_vH		37.3	298		[1971WIL/ZWO]
	Δ_vH	(243–380)	39.7	258		[1947STU]
	Δ_vH		37.3 ± 0.1	298	C	[1947OSB/GIN]
	Δ_vH	(302–381)	36.6	317	A,MM	[1987STE/MAL, 1945WIL/TAY]
C ₈ H ₁₈	[584-94-1]	2,3-dimethylhexane				
	Δ_vH		38.8	298		[1971WIL/ZWO]
	Δ_vH	(250–388)	41.4	265		[1947STU]
	Δ_vH		38.8 ± 0.1	298	C	[1947OSB/GIN]
	Δ_vH	(310–390)	37.6	325	A,MM	[1987STE/MAL, 1945WIL/TAY]
C ₈ H ₁₈	[589-43-5]	2,4-dimethylhexane				
	Δ_vH		37.8	298		[1971WIL/ZWO]
	Δ_vH	(246–382)	41	261		[1947STU]
	Δ_vH		37.8 ± 0.1	298	C	[1947OSB/GIN]
	Δ_vH	(305–385)	36.9	320	A,MM	[1987STE/MAL, 1945WIL/TAY]
C ₈ H ₁₈	[592-13-2]	2,5-dimethylhexane				
	Δ_vH		37.9	298		[1971WIL/ZWO]
	Δ_vH	(246–382)	41.1	261		[1947STU]
	Δ_vH		37.9 ± 0.1	298	C	[1947OSB/GIN]
	Δ_vH	(307–383)	36.9	322	A,MM	[1987STE/MAL, 1945WIL/TAY]
C ₈ H ₁₈	[563-16-6]	3,3-dimethylhexane				
	Δ_vH	(247–385)	41.2	262		[1947STU]
	Δ_vH		37.5 ± 0.1	298	C	[1947OSB/GIN]
	Δ_vH	(308–386)	36.6	323	A,MM	[1987STE/MAL, 1945WIL/TAY]
C ₈ H ₁₈	[583-48-2]	3,4-dimethylhexane				
	Δ_vH	(251–390)	41.3	266		[1947STU]
	Δ_vH		39.0 ± 0.1	298	C	[1947OSB/GIN]
	Δ_vH	(313–392)	37.7	328	A,MM	[1987STE/MAL, 1945WIL/TAY]
C ₈ H ₁₈	[619-99-8]	3-ethylhexane				
	Δ_vH		39.7	298		[1971WIL/ZWO]
	Δ_vH	(251–391)	42.4	268		[1947STU]
	Δ_vH		39.6 ± 0.1	298	C	[1947OSB/GIN]
	Δ_vH	(314–393)	38.2	329	A,MM	[1987STE/MAL, 1945WIL/TAY]
C ₈ H ₁₈	[609-26-7]	3-ethyl-2-methylpentane				
	Δ_vH		38.5	298		[1971WIL/ZWO]
	Δ_vH		38.5 ± 0.1	298	C	[1947OSB/GIN]
	Δ_vH	(311–390)	37.4	326	A,MM	[1987STE/MAL, 1945WIL/TAY]
C ₈ H ₁₈	[1067-08-9]	3-ethyl-3-methylpentane				
	Δ_vH		38.0	298		[1971WIL/ZWO]
	Δ_vH	(249–391)	40.2	264		[1947STU]
	Δ_vH		38.0 ± 0.1	298	C	[1947OSB/GIN]
	Δ_vH	(312–393)	36.9	327	A,MM	[1987STE/MAL, 1945WIL/TAY]
C ₈ H ₁₈	[592-27-8]	2-methylheptane				
	$\Delta_{\text{fus}}H$		11.92	164.2		[1996DOM/HEA]
	Δ_vH	(285–392)	39.8	300	A	[1987STE/MAL]
	Δ_vH		39.7 ± 0.1	298	C	[1979MAJ/SVO]
	Δ_vH		38.7 ± 0.1	313	C	[1979MAJ/SVO]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$		37.3 ± 0.1	333	C	[1979MAJ/SVO]
	$\Delta_v H$		36.0 ± 0.1	353	C	[1979MAJ/SVO]
	$\Delta_v H$	(233–283)	41.6	268	IP	[1987STE/MAL, 1974OSB/DOU]
	$\Delta_v H$		39.7	298		[1971WIL/ZWO]
	$\Delta_v H$		39.8 ± 0.1	298	C	[1947OSB/GIN]
	$\Delta_v H$	(315–391)	38.1	330	MM	[1945WIL/TAY]
C ₈ H ₁₈	[589-81-1]	3-methylheptane				
	$\Delta_{\text{fus}} H$		11.67	152.6		[1996DOM/HEA]
	$\Delta_v H$		39.8 ± 0.2	298	C	[1987AN/HU]
	$\Delta_v H$	(286–393)	40.1	301	A	[1987STE/MAL]
	$\Delta_v H$	(238–286)	41.6	271	IP	[1987STE/MAL, 1974OSB/DOU]
	$\Delta_v H$		39.8	298		[1971WIL/ZWO]
	$\Delta_v H$		39.8 ± 0.1	298	C	[1947OSB/GIN]
C ₈ H ₁₈	[589-53-7]	4-methylheptane				
	$\Delta_{\text{fus}} H$		10.84	152.2		[1991ACR]
	$\Delta_v H$		39.7 ± 0.1	298	C	[1979MAJ/SVO]
	$\Delta_v H$		38.7 ± 0.1	313	C	[1979MAJ/SVO]
	$\Delta_v H$		37.4 ± 0.1	333	C	[1979MAJ/SVO]
	$\Delta_v H$		36.1 ± 0.1	353	C	[1979MAJ/SVO]
	$\Delta_v H$	(253–391)	39.7	298		[1971WIL/ZWO]
C ₈ H ₁₈	[111-65-9]	octane				
	$\Delta_{\text{fus}} H$		21.8	216.6	DSC	[2004MON/RAJ]
	$\Delta_{\text{fus}} H$		20.74	216.4		[1996DOM/HEA]
	$\Delta_{\text{sub}} H$		68.1	216	B	[1963BON]
	$\Delta_v H$	(323–563)	39.4	338	EB	[2003EWI/OCH]
	$\Delta_v H$		41.6	298		[1994RUZ/MAJ]
	$\Delta_v H$	(297–400)	41.0	312	A	[1987STE/MAL]
	$\Delta_v H$	(216–278)	44.4	263	A	[1987STE/MAL]
	$\Delta_v H$	(396–432)	36.3	411	A	[1987STE/MAL]
	$\Delta_v H$	(428–510)	35.5	443	A	[1987STE/MAL]
	$\Delta_v H$	(506–569)	34.9	521	A	[1987STE/MAL]
	$\Delta_v H$	(295–402)	41.2	310		[1986PAU/KRU]
	$\Delta_v H$	(298–333)	41.9	313		[1984MIC/JOS]
	$\Delta_v H$		41.5 ± 0.1	298	C	[1981HOS/SCO2]
$\Delta_v H$		41.5 ± 0.1	298	C	[1979MAJ/SVO]	
$\Delta_v H$		40.5 ± 0.1	313	C	[1979MAJ/SVO]	
$\Delta_v H$		39.1 ± 0.1	333	C	[1979MAJ/SVO]	
$\Delta_v H$		37.8 ± 0.1	353	C	[1979MAJ/SVO]	
$\Delta_v H$	(217–297)	43.0	282		[1973CAR/KOB]	
$\Delta_v H$		41.5	298		[1971WIL/ZWO]	
$\Delta_v H$		38.0 ± 0.1	311	C	[1960MCK/SAG]	
$\Delta_v H$		36.7 ± 0.1	328	C	[1960MCK/SAG]	
$\Delta_v H$		35.4 ± 0.1	344	C	[1960MCK/SAG]	
$\Delta_v H$		41.5 ± 0.1	298	C	[1947OSB/GIN]	
$\Delta_v H$	(326–400)	39.2	341	MM	[1945WIL/TAY]	
C ₈ D ₁₈	[17252-77-6]	octane-d ₁₈				
	$\Delta_v H$		41.4	298	CGC	[2008ZHA/UNH]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₈ H ₁₈	[560-21-4]	<i>(dl)</i> 2,2,3-trimethylpentane				
	$\Delta_v H$		37.7 ± 0.1	298	C	[1998SVO/HYN]
	$\Delta_v H$		37.1 ± 0.1	308	C	[1998SVO/HYN]
	$\Delta_v H$		36.6 ± 0.1	315	C	[1998SVO/HYN]
	$\Delta_v H$		36.0 ± 0.1	323	C	[1998SVO/HYN]
	$\Delta_v H$		35.5 ± 0.1	330	C	[1998SVO/HYN]
	$\Delta_v H$		35.1 ± 0.1	338	C	[1998SVO/HYN]
	$\Delta_v H$		34.8 ± 0.1	348	C	[1998SVO/HYN]
	$\Delta_v H$		34.1 ± 0.1	358	C	[1998SVO/HYN]
	$\Delta_v H$		33.5 ± 0.1	368	C	[1998SVO/HYN]
	$\Delta_v H$		36.9	298		[1971WIL/ZWO]
	$\Delta_v H$		36.9 ± 0.1	298	C	[1947OSB/GIN]
$\Delta_v H$	(306–384)	36.1	321	A,MM	[1987STE/MAL, 1945WIL/TAY]	
C ₈ H ₁₈	[540-84-1]	2,2,4-trimethylpentane				
	$\Delta_{\text{fus}} H$		9.04	165.3		[1996DOM/HEA]
	$\Delta_v H$	(373–423)	34.9	298	CGC	[1995CHI/HOS]
	$\Delta_v H$	(289–333)	36.1	304		[1991WU/PIV]
	$\Delta_v H$	(423–523)	31.6	438	A	[1987STE/MAL]
	$\Delta_v H$	(372–416)	32.2	387	A	[1987STE/MAL]
	$\Delta_v H$	(413–494)	31.5	428	A	[1987STE/MAL]
	$\Delta_v H$	(490–544)	31.4	505	A	[1987STE/MAL]
	$\Delta_v H$		35.2 ± 0.1	298	C	[1982SVO/CHA]
	$\Delta_v H$		34.4 ± 0.1	313	C	[1982SVO/CHA]
	$\Delta_v H$		33.4 ± 0.1	328	C	[1982SVO/CHA]
	$\Delta_v H$		32.6 ± 0.1	343	C	[1982SVO/CHA]
	$\Delta_v H$		31.7 ± 0.1	358	C	[1982SVO/CHA]
	$\Delta_v H$		31.0 ± 0.1	368	C	[1982SVO/CHA]
	$\Delta_v H$		35.1 ± 0.1	298	C	[1979MAJ/SVO]
	$\Delta_v H$		34.3 ± 0.1	313	C	[1979MAJ/SVO]
	$\Delta_v H$		33.2 ± 0.1	333	C	[1979MAJ/SVO]
	$\Delta_v H$		32.0 ± 0.1	353	C	[1979MAJ/SVO]
	$\Delta_v H$		35.1	298		[1971WIL/ZWO]
	$\Delta_v H$	(194–299)	40.7	209	A	[1987STE/MAL, 1956MIL]
$\Delta_v H$		35.1 ± 0.1	298	C	[1947OSB/GIN]	
$\Delta_v H$	(297–374)	34.8	312	A,MM	[1987STE/MAL, 1945WIL/TAY]	
$\Delta_v H$		31.0	371	C	[1940PIT]	
$\Delta_v H$	(318–399)	33.9	333	EB	[1940SMI]	
C ₈ H ₁₈	[560-21-4]	2,3,3-trimethylpentane				
	$\Delta_v H$		37.6 ± 0.1	298	C	[1998SVO/HYN]
	$\Delta_v H$		36.9 ± 0.1	308	C	[1998SVO/HYN]
	$\Delta_v H$		36.5 ± 0.1	315	C	[1998SVO/HYN]
	$\Delta_v H$		36.0 ± 0.1	323	C	[1998SVO/HYN]
	$\Delta_v H$		35.5 ± 0.1	330	C	[1998SVO/HYN]
	$\Delta_v H$		35.1 ± 0.1	338	C	[1998SVO/HYN]
	$\Delta_v H$		34.4 ± 0.1	348	C	[1998SVO/HYN]
	$\Delta_v H$		33.9 ± 0.1	358	C	[1998SVO/HYN]
	$\Delta_v H$		33.3 ± 0.1	368	C	[1998SVO/HYN]
	$\Delta_v H$		37.2	298		[1971WIL/ZWO]
	$\Delta_v H$		36.9 ± 0.1	298	C	[1947OSB/GIN]
	$\Delta_v H$		37.2 ± 0.1	298	C	[1947OSB/GIN]
	$\Delta_v H$	(308–390)	36.4	323	A,MM	[1987STE/MAL, 1945WIL/TAY]
C ₈ H ₁₈	[565-75-3]	2,3,4-trimethylpentane				
	$\Delta_{\text{fus}} H$		9.27	163.6		[1996DOM/HEA]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(288–400)	37.7	303	A	[1987STE/MAL]
	$\Delta_v H$	(223–289)	39.1	274	A	[1987STE/MAL]
	$\Delta_v H$		37.7 ± 0.1	298	C	[1981HOS/SCO2]
	$\Delta_v H$	(223–426)	41.3	238	IP,EB	[1974OSB/DOU]
	$\Delta_v H$	(223–278)	39.8	263	IP	[1974OSB/DOU]
	$\Delta_v H$		37.7	298		[1971WIL/ZWO]
	$\Delta_v H$		37.7 ± 0.1	298	C	[1947OSB/GIN]
	$\Delta_v H$	(310–388)	36.7	325	MM	[1945WIL/TAY]
C₈H₁₈	[594-82-1]	2,2,3,3-tetramethylbutane				
	$\Delta_{\text{us}}H$		2.0	153		[1984DOM/EVA]
	$\Delta_{\text{us}}H$		2.0	152.5		
	$\Delta_{\text{fus}}H$		7.54	373.9		[1996DOM/HEA]
	$\Delta_{\text{sub}}H$	(286–377)	43.6	301		[1987STE/MAL]
	$\Delta_{\text{sub}}H$	(273–338)	43.4 ± 0.2	298		[1952SCO/DOU, 1970COX/PIL]
	$\Delta_{\text{sub}}H$		42.9 ± 0.9	298	C	[1947OSB/GIN]
	$\Delta_{\text{sub}}H$	(263–279)	56.2		A, MG	[1931LIN]
	$\Delta_v H$	(377–390)	333	383	A	[1987STE/MAL]
C₈H₁₈N₂	[2159-75-3]	dibutyldiazene				
	$\Delta_v H$		49.3 ± 0.2	298	C	[1978ENG/MON]
C₈H₁₈N₂	[927-83-3]	di- <i>tert</i> -butyldiazene				
	$\Delta_{\text{us}}H$		4.89	242.6		
	$\Delta_{\text{fus}}H$		10.28	258.6		[1980BYS]
	$\Delta_v H$		39.1 ± 0.3	298	C	[1976ENG/MEL]
	$\Delta_v H$	(294–305)	39.6	299	UV	[1974ENG/WOO]
	$\Delta_v H$		32.7		I	[1974ENG/WOO]
C₈H₁₈N₂	[na]	butylhydrazone butyraldehyde				
	$\Delta_v H$	(298–323)	55.6	310		[1980LEB/NAZ]
C₈H₁₈N₂	[na]	isobutylhydrazone isobutyraldehyde				
	$\Delta_v H$	(288–313)	57.2	300		[1980LEB/NAZ]
C₈H₁₈N₂O	[16649-52-8]	dibutyldiazene N-oxide				
	$\Delta_{\text{us}}H$		8.34	268		
	$\Delta_{\text{fus}}H$		11.52	288.4		[1980BYS]
	$\Delta_v H$		45.9 ± 0.1	298	C	[1981BYS]
C₈H₁₈N₂O	[42955-46-4]	1-heptyl urea				
	$\Delta_{\text{fus}}H$		29.0	386.1	DSC	[2005HAS/TAJ]
	$\Delta_{\text{us}}H$		1.4	379.2		
	$\Delta_{\text{fus}}H$		26.3	382.2		[1999WEL/DRU]
C₈H₁₈N₂O₂	[122-96-3]	1,4- <i>bis</i> -(2-hydroxyethyl)piperazine				
	$\Delta_{\text{fus}}H$		25.9	405		[1996DOM/HEA]
	$\Delta_{\text{sub}}H$	(354–396)	128.0 ± 1.0	375	GS	[2002VER2]
	$\Delta_{\text{sub}}H$	(354–396)	130.5 ± 1.0	298	GS	[2002VER2]
	$\Delta_{\text{sub}}H$	(334–356)	104.1			[1984LEB/GUT]
C₈H₁₈N₄O₄	[35823-10-0]	N,N'-dimethyl-N,N'-dinitro-1,6-hexanediamine				
	$\Delta_{\text{fus}}H$		61.68	331		[1987OYU/BRI]
C₈H₁₈O	[629-32-3]	heptyl methyl ether				
	$\Delta_v H$		46.9	298		[UR/FUC, 1985MAJ/SVO]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₈ H ₁₈ O	[1000-63-1]	butyl <i>tert</i> -butyl ether				
	$\Delta_v H$		43.2	298		[UR/VER, 2002VER]
	$\Delta_v H$		41.6 ± 0.2	298	C	[2002VAR/AIT]
	$\Delta_v H$		42.3 ± 0.3	298	C	[1991SHA/MIS]
	$\Delta_v H$	(293–397)	41.7	308	A	[1987STE/MAL]
		(356–397)	38.3	371	EB	[1987STE/MAL, 1969SHE/ALN]
C ₈ H ₁₈ O	[33021-02-2]	<i>tert</i> -butyl isobutyl ether				
	$\Delta_{\text{fus}} H$		8.65	162.3		[2006DRU/DOR]
	$\Delta_v H$	(349–386)	39.1 ± 0.5	298	EB	[2007EFI/PAS]
	$\Delta_v H$		41.2 ± 0.3	298		[UR/VER, 2002VER]
	$\Delta_v H$		39.2 ± 0.3	298	C	[2002VAR/AIT]
	$\Delta_v H$	(273–308)	40.9 ± 0.3	298	GS	[1996VER/BEC]
			40.1 ± 0.1	298	C	[1991SHA/MIS]
C ₈ H ₁₈ O	[17071-47-5]	butyl isobutyl ether				
	$\Delta_v H$	(328–406)	40.3	343	A	[1987STE/MAL]
C ₈ H ₁₈ O	[32970-45-9]	sec-butyl <i>tert</i> -butyl ether				
	$\Delta_v H$		41.3	298		[2002VER, 2003VER/KRA]
C ₈ H ₁₈ O	[32970-45-9]	1-methyl-1- <i>tert</i> -butoxypropane				
	$\Delta_v H$		40.3 ± 0.2	298	C	[1991SHA/MIS]
C ₈ H ₁₈ O	[142-96-1]	dibutyl ether				
	$\Delta_v H$	(339–415)	40.9	354	A	[1987STE/MAL]
	$\Delta_v H$	(336–415)	41.7	351	A	[1987STE/MAL]
	$\Delta_v H$		44.7 ± 0.1	298	C	[1982FUC/PEA]
	$\Delta_v H$		45.0 ± 0.1	298	C	[1980MAJ/WAN]
	$\Delta_v H$	(362–414)	44.4	298		[1976AMB/ELL]
	$\Delta_v H$	(362–414)	36.4	413		[1976AMB/ELL]
	$\Delta_v H$	(362–413)	40.6	377	EB	[1969CID/POL]
		(386–440)	39.4	413		[1965NIS/LAP]
C ₈ H ₁₈ O	[6163-66-2]	di- <i>tert</i> -butyl ether				
	$\Delta_v H$	(290–386)	37.7 ± 0.3	298	EB	[1996STE/CHI2]
	$\Delta_v H$	(289–382)	37.3	304	A	[1987STE/MAL, 1976AMB/ELL]
	$\Delta_v H$	(289–382)	37.2	298		[1976AMB/ELL]
	$\Delta_v H$	(289–382)	31.6	380		[1976AMB/ELL]
	$\Delta_v H$		37.6 ± 0.1	298	C	[1975FEN/HAR]
		(277–382)	38.7	292	A	[1987STE/MAL, 1961SMU/BON]
C ₈ H ₁₈ O	[628-55-7]	diisobutyl ether				
	$\Delta_{\text{fus}} H$	(8–373)	1.8	170.7		
	$\Delta_{\text{fus}} H$	(8–383)	11.33	190.4	AC	[2009EFI/DRU]
	$\Delta_v H$	(331–395)	41.2 ± 0.7	298	EB	[2009EFI/DRU]
		(320–396)	38.9	335	A	[1987STE/MAL]
C ₈ H ₁₈ O	[74058-13-2]	propyl <i>tert</i> -amyl ether				
	$\Delta_v H$		43.8 ± 0.7	298		[2002VER, 2003VER/KRA]
C ₈ H ₁₈ O	[3249-46-5]	isopropyl <i>tert</i> -amyl ether				
	$\Delta_v H$		41.6	298		[2002VER, 2003VER/KRA]
C ₈ H ₁₈ O	[104-76-7]	2-ethyl-1-hexanol				
	$\Delta_v H$	(293–331)	68.5 ± 0.2	298	GS	[2005ROG/PIS]
	$\Delta_v H$	(373–398)	52.7	388		[1973LIN/WIC]
		(347–457)	60.2	362	A	[1987STE/MAL, 1961DYK/SEP]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		T _m (K)	Method	Reference
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)			
C ₈ H ₁₈ O	[597-76-2] $\Delta_v H$	3-ethyl-3-hexanol (331–433)	49.2	345		[1973WIL/ZWO]
C ₈ H ₁₈ O	[10137-88-9] $\Delta_v H$	2-ethyl-4-methyl-1-pentanol (343–450)	58.9	358	A	[1987STE/MAL, 1961DYK/SEP, 1973WIL/ZWO]
C ₈ H ₁₈ O	[106-67-2] $\Delta_v H$	2-methyl-1-heptanol (350–449)	53.3	365	A	[1987STE/MAL, 1973WIL/ZWO]
C ₈ H ₁₈ O	[1070-32-2] $\Delta_v H$	3-methyl-1-heptanol (360–459)	53.4	375	A	[1987STE/MAL, 1973WIL/ZWO]
C ₈ H ₁₈ O	[817-91-4] $\Delta_v H$ $\Delta_v H$	4-methyl-1-heptanol (357–456) (354–456)	55.9 56.7	372 369	A	[1987STE/MAL] [1973WIL/ZWO]
C ₈ H ₁₈ O	[7212-53-5] $\Delta_v H$	(dl) 5-methyl-1-heptanol (364–460)	57.6	379	A	[1987STE/MAL, 1973WIL/ZWO]
C ₈ H ₁₈ O	[1653-40-3] $\Delta_v H$	6-methyl-1-heptanol (368–610)	61.0	383	A	[1987STE/MAL, 1973WIL/ZWO]
C ₈ H ₁₈ O	[625-25-2] $\Delta_v H$ $\Delta_v H$ $\Delta_v H$	2-methyl-2-heptanol (275–314) (343–430) (339–429)	62.9 ± 0.2 53.1 55	298 358 354	GS A	[2005ROG/PIS] [1987STE/MAL] [1973WIL/ZWO]
C ₈ H ₁₈ O	[31367-46-1] $\Delta_v H$	3-methyl-2-heptanol (341–440)	48.0	356	A	[1987STE/MAL, 1973WIL/ZWO]
C ₈ H ₁₈ O	[56298-90-9] $\Delta_v H$	4-methyl-2-heptanol (351–445)	54.2	366	A	[1987STE/MAL, 1973WIL/ZWO]
C ₈ H ₁₈ O	[54630-50-1] $\Delta_v H$ $\Delta_v H$	5-methyl-2-heptanol (348–445) (348–445)	51.9 47.2	363 363	A	[1987STE/MAL] [1973WIL/ZWO]
C ₈ H ₁₈ O	[4730-22-7] $\Delta_v H$	(dl) 6-methyl-2-heptanol (354–445)	55.2	369	A	[1987STE/MAL, 1973WIL/ZWO]
C ₈ H ₁₈ O	[18720-62-2] $\Delta_v H$	(dl) 2-methyl-3-heptanol (349–441)	54.8	364	A	[1987STE/MAL, 1973WIL/ZWO]
C ₈ H ₁₈ O	[5582-82-1] $\Delta_v H$ $\Delta_v H$	3-methyl-3-heptanol (344–433) (338–433)	54.1 54.7	359 353	A	[1987STE/MAL] [1973WIL/ZWO]
C ₈ H ₁₈ O	[14979-39-6] $\Delta_v H$	4-methyl-3-heptanol (330–429)	43.9	345	A	[1987STE/MAL, 1973WIL/ZWO]
C ₈ H ₁₈ O	[18720-65-5] $\Delta_v H$	5-methyl-3-heptanol (330–427)	46.5	345	A	[1987STE/MAL, 1973WIL/ZWO]
C ₈ H ₁₈ O	[18720-66-6] $\Delta_v H$	(dl) 6-methyl-3-heptanol (333–432)	47.6	348	A	[1987STE/MAL]
C ₈ H ₁₈ O	[21570-35-4] $\Delta_v H$ $\Delta_v H$	2-methyl-4-heptanol (348–440) (345–437)	54.8 56.3	363 360	A	[1987STE/MAL] [1973WIL/ZWO]
C ₈ H ₁₈ O	[1838-73-9] $\Delta_v H$	(dl) 3-methyl-4-heptanol (340–438)	48.0	355	A	[1987STE/MAL, 1973WIL/ZWO]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₈ H ₁₈ O	[598-01-6]	4-methyl-4-heptanol				
	$\Delta_v H$	(344–434)	54.4	359	A	[1987STE/MAL]
	$\Delta_v H$	(331–434)	54.8	345		[1973WIL/ZWO]
C ₈ H ₁₈ O	[19550-07-3]	2,5-dimethyl-3-hexanol				
	$\Delta_v H$	(337–431)	55	352		[1973WIL/ZWO]
C ₈ H ₁₈ O	[111-87-5]	1-octanol				
	$\Delta_{\text{fus}} H$		25.24	258.4		[2003VAN/GAB]
	$\Delta_{\text{sub}} H$		100.4	298		[1965DAV/KYB]
	$\Delta_v H$	(358–463)	69.3	298		[2006NAS/NEU]
	$\Delta_v H$	(282–321)	69.6	303	GS	[2001KUL/VER2]
	$\Delta_v H$	(282–321)	70.1	298	GS	[2001KUL/VER2]
	$\Delta_v H$	(373–423)	71.6	298	CGC	[1995CHI/HOS]
	$\Delta_v H$	(273–363)	68.7	318		[1992NGU/KAS]
	$\Delta_v H$	(328–400)	67.3	343	A	[1987STE/MAL]
	$\Delta_v H$	(430–474)	52.5	445	A	[1987STE/MAL]
	$\Delta_v H$	(397–479)	56.6	412	A	[1987STE/MAL]
	$\Delta_v H$	(475–555)	47.8	490	A	[1987STE/MAL]
	$\Delta_v H$		71.0 ± 0.4	298	C	[1977MAN/SEL]
	$\Delta_v H$	(343–468)	67.5	358		[1973WIL/ZWO]
	$\Delta_v H$	(386–480)	58.3	401	EB	[1987STE/MAL, 1970AMB/SPR]
	$\Delta_v H$	(352–468)	65.0	367	DTA	[1969KEM/KRE]
	$\Delta_v H$	(293–353)	70.4	308		[1966GEI/FRU]
$\Delta_v H$	(267–282)	64.0	274	A,ME	[1987STE/MAL, 1965DAV/KYB]	
$\Delta_v H$	(365–427)	61.6	380		[1958ROS/PAP]	
C ₈ H ₁₈ O	[123-96-6]	<i>(dl)</i> 2-octanol				
	$\Delta_v H$	(284–329)	67.9 ± 0.3	298	GS	[2007VER/SCH]
	$\Delta_v H$	(253–353)	70.7	268		[1999NGU/BER]
	$\Delta_v H$	(333–453)	60.7	348	A	[1987STE/MAL]
	$\Delta_v H$	(367–453)	56.1	382		[1984SAC/MAR]
$\Delta_v H$	(345–453)	60.0	360		[1973WIL/ZWO]	
C ₈ H ₁₈ O	[20296-29-1]	<i>(dl)</i> 3-octanol				
	$\Delta_v H$	(288–324)	67.9 ± 0.3	298	GS	[2007VER/SCH]
	$\Delta_v H$	(253–348)	71.6	268		[1999NGU/BER]
	$\Delta_v H$	(313–450)	64.1	328	A	[1987STE/MAL]
	$\Delta_v H$	(366–450)	54.5	381		[1984SAC/MAR]
$\Delta_v H$	(349–449)	58.8	364		[1973WIL/ZWO]	
C ₈ H ₁₈ O	[589-62-8]	<i>(dl)</i> 4-octanol				
	$\Delta_v H$	(288–322)	67.2 ± 0.5	298	GS	[2007VER/SCH]
	$\Delta_v H$	(343–450)	57.3	358	A	[1987STE/MAL]
	$\Delta_v H$	(364–449)	54.8	379		[1984SAC/MAR]
$\Delta_v H$	(341–449)	62.1	356		[1973WIL/ZWO]	
C ₈ H ₁₈ O	[123-44-4]	<i>(dl)</i> 2,4,4-trimethyl-1-pentanol				
	$\Delta_v H$		60.6 ± 0.1	328	C	[1996ULB/KLU]
	$\Delta_v H$		58.6 ± 0.1	343	C	[1996ULB/KLU]
	$\Delta_v H$		56.5 ± 0.1	358	C	[1996ULB/KLU]
	$\Delta_v H$	(352–446)	54.2	367	A	[1987STE/MAL]
$\Delta_v H$	(333–441)	54.7	348		[1973WIL/ZWO]	
C ₈ H ₁₈ O	[7294-05-5]	2,2,3-trimethyl-3-pentanol				
	$\Delta_v H$	(318–426)	47.3	333		[1973WIL/ZWO]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₈ H ₁₈ O	[5162-48-1] $\Delta_v H$	2,2,4-trimethyl-3-pentanol (328–428)	57.1	343		[1973WIL/ZWO]
C ₈ H ₁₈ O ₂	[110-05-4] $\Delta_v H$	di- <i>tert</i> -butyl peroxide (308–358)	37.0	333		[1995DIO/MIN]
	$\Delta_v H$	(308–358)	38.9	298		[1995DIO/MIN]
	$\Delta_v H$		36.6 ± 0.6	298	C	[1990VAN/PAV]
	$\Delta_v H$	(246–311)	32.0	261	A	[1987STE/MAL, 1978IND/STO]
	$\Delta_v H$	(273–384)	31.0	288	A	[1987STE/MAL, 1951EGE/EMT]
	$\Delta_v H$		40.2	298		[1951VAU, 1948RAL/RUS]
C ₈ H ₁₈ O ₂	[18854-56-3] $\Delta_v H$	1,2-dipropoxyethane (234–453)	U28.2	249	A	[1987STE/MAL]
	$\Delta_v H$		50.6 ± 0.1	298	C	[1970KUS/WAD]
C ₈ H ₁₈ O ₂	[4413-13-2] $\Delta_v H$	1-butoxy-2-ethoxyethane 50.9 ± 0.1		298	C	[1970KUS/WAD]
C ₈ H ₁₈ O ₂	[4468-93-3] $\Delta_v H$	ethylene glycol mono(2-ethylbutyl) ether (357–470)	53.4	372	A	[1987STE/MAL]
C ₈ H ₁₈ O ₂	[112-25-4] $\Delta_v H$	ethylene glycol monoheptyl ether (363–483)	54.6	378	A	[1987STE/MAL]
C ₈ H ₁₈ O ₂	[na] $\Delta_v H$	3-hydroxymethyl-4-heptanol (375–518)	61.6	390	A	[1987STE/MAL]
C ₈ H ₁₈ O ₂	[629-41-4] $\Delta_{\text{fus}} H$	1,8-octanediol	36.1	332.8		[1991ACR]
	$\Delta_{\text{sub}} H$		139.3 ± 0.9	298	C	[1990KNA/SAB]
	$\Delta_v H$		101	356		[1993PIA/FER, 2006UMN/KWE]
	$\Delta_v H$		107.0 ± 2.2	298		[1993PIA/FER, 2006UMN/KWE]
	$\Delta_v H$		105.4 ± 1.8	298		[1990KNA/SAB, 2006UMN/KWE]
C ₈ H ₁₈ O ₂	[144-19-4] $\Delta_{\text{fus}} H$	2,2,4-trimethyl-1,3-pentanediol	24.2	328.3		[2002STE/CHI]
	$\Delta_v H$	(396–489)	66.6 ± 2.1	400	EB	[2002STE/CHI]
	$\Delta_v H$	(396–489)	60.3 ± 1.7	440	EB	[2002STE/CHI]
	$\Delta_v H$	(396–489)	55.0 ± 1.6	480	EB	[2002STE/CHI]
	$\Delta_v H$	(413–502)	58.5	428	A	[1987STE/MAL]
C ₈ H ₁₈ O ₂	[94-96-2] $\Delta_v H$	2-ethyl-1,3-hexanediol (331–413)	79.5	298		[1974BLA/LEV, 2007VER]
C ₈ H ₁₈ O ₂	[110-03-2] $\Delta_v H$	2,5-dimethyl-2,5-hexanediol	85.2 ± 3.5	298	CGC	[2006UMN/KWE]
C ₈ H ₁₈ O ₂ S	[598-04-9] $\Delta_{\text{sub}} H$	di- <i>n</i> -butyl sulfone	100.4 ± 2.5			[UR/MAC, 1970COX/PIL]
C ₈ H ₁₈ O ₂ S	[1886-75-7] $\Delta_{\text{sub}} H$	di- <i>tert</i> -butyl sulfone	94.1 ± 2.9			[UR/MAC, 1970COX/PIL]
C ₈ H ₁₈ O ₃	[1538-75-6] $\Delta_v H$	trimethylacetic acid anhydride (355–513)	50.7 ± 0.2	360	EB	[2002STE/CHI4]
	$\Delta_v H$	(355–513)	47.4 ± 0.2	400	EB	[2002STE/CHI4]
	$\Delta_v H$	(355–513)	44.0 ± 0.4	440	EB	[2002STE/CHI4]
	$\Delta_v H$	(355–513)	40.3 ± 0.7	480	EB	[2002STE/CHI4]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₈ H ₁₈ O ₃	[112-36-7]	diethylene glycol diethyl ether				
	$\Delta_v H$		56.4 ± 1.4	298	CGC	[2000NIC/ORF]
		(330–461)	48.3	345	A	[1987STE/MAL]
C ₈ H ₁₈ O ₃	[112-34-5]	diethylene glycol monobutyl ether				
	$\Delta_v H$	(415–505)	55.7	430	A	[1987STE/MAL]
C ₈ H ₁₈ O ₄	[112-49-2]	1,2- <i>bis</i> (2-methoxyethoxy)ethane (triglyme)				
	$\Delta_v H$		63.7 ± 3.3	298	CGC	[2000NIC/ORF]
C ₈ H ₁₈ O ₄	[112-49-2]	2,5,8,11-tetraoxadodecane				
	$\Delta_{\text{fus}} H$		23.71	229.3		[1996DOM/HEA]
C ₈ H ₁₈ O ₄	[na]	1,2,7,8-tetrahydroxyoctane				
	$\Delta_{\text{fus}} H$		36.7	352.2		[1991HEN/TSC, 1994HEN/DIE]
C ₈ H ₁₈ O ₄ S ₂	[76-20-0]	2,2-butanediol <i>bis</i> (ethylsulfonate)				
	$\Delta_v H$	(443–493)	75.7	458	A	[1987STE/MAL, 1999DYK/SVO]
C ₈ H ₁₈ O ₅	[112-60-7]	tetraethylene glycol				
	$\Delta_v H$	(426–581)	92.2	441	A	[1987STE/MAL, 1947STU]
C ₈ H ₁₈ S	[544-40-1]	dibutyl sulfide				
	$\Delta_{\text{fus}} H$		19.41	198.1		[1996DOM/HEA]
	$\Delta_v H$	(255–422)	44.8	339		[2004SAW/MOK]
	$\Delta_v H$	(283–390)	40.3	298		[1999DYK/SVO]
	$\Delta_v H$		53	298		[1981SHI/SAI]
	$\Delta_v H$		54.2 ± 0.8	298	GC	[1964MAC/MCC]
	$\Delta_v H$	(390–470)	46.5	405	A,EB	[1987STE/MAL, 1952WHI/BER]
C ₈ H ₁₈ S	[626-26-6]	di- <i>tert</i> -butyl sulfide				
	$\Delta_v H$	(264–329)	44.9	279		[1999DYK/SVO]
	$\Delta_v H$	(329–470)	41.4	344		[1999DYK/SVO]
	$\Delta_v H$	(390–470)	46.4	405		[1999DYK/SVO]
	$\Delta_v H$	(278–308)	44.8	293		[1998STO/NG]
	$\Delta_v H$	(324–420)	42.4	339	A	[1987STE/MAL]
	$\Delta_v H$		43.8	298		[1981SHI/SAI]
	$\Delta_v H$		43.8 ± 0.1	298		[1972GOO]
	$\Delta_v H$		49.3 ± 0.8	298	GC	[1964MAC/MCC]
	(325–350)	42.3	333	EB	[1962MAC/MAY2]	
C ₈ H ₁₈ S	[592-65-4]	diisobutyl sulfide				
	$\Delta_v H$	(325–346)	46.4	335	A	[1987STE/MAL, 1999DYK/SVO]
	$\Delta_v H$		48.7	298		[1981SHI/SAI]
	$\Delta_v H$		48.5 ± 0.8	298	GC	[1964MAC/MCC]
		(326–346)	43.1	336	EB	[1962MAC/MAY2]
C ₈ H ₁₈ S	[626-26-6]	di- <i>sec</i> -butyl sulfide				
	$\Delta_v H$	(255–422)	47.0	298		[2004SAW/MOK]
C ₈ H ₁₈ S	[111-88-6]	1-octanethiol				
	$\Delta_{\text{fus}} H$		24.27	224		[1996DOM/HEA, 1985DEA]
		(372–473)	49.6	387	A	[1987STE/MAL, 1999DYK/SVO, 1932ELL/REI]
C ₈ H ₁₈ S	[3001-66-9]	2-octanethiol				
	$\Delta_v H$	(347–489)	49.0	362		[1999DYK/SVO]
C ₈ H ₁₈ S	[10435-81-1]	<i>dl</i> 2-octanethiol				
	$\Delta_v H$	(361–460)	48.0	376	A	[1987STE/MAL, 1999DYK/SVO, 1932ELL/REI]

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₈ H ₁₈ S ₂	[629-45-8]	dibutyl disulfide				
	$\Delta_v H$	(383–423)	64.1	298	CGC	[1995CHI/HOS]
	$\Delta_v H$		62.3 ± 0.2	298	C	[1985KUS]
C ₈ H ₁₈ S ₂	[na]	diisobutyl disulfide				
	$\Delta_v H$		57.2	298		[1981SHI/SAI]
C ₈ H ₁₈ S ₂	[1518-72-5]	2,7-dimethyl-4,5-dithiaoctane				
	$\Delta_v H$		57.2 ± 0.1	298	C	[1985KUS]
C ₈ H ₁₈ S ₂	[110-06-5]	2,2,5,5-tetramethyl-3,4-dithiahexane				
	$\Delta_v H$	(383–423)	53.8	298	CGC	[1995CHI/HOS]
	$\Delta_v H$		52.5 ± 0.2	298	C	[1985KUS]
C ₈ H ₁₈ S ₂	[na]					
	$\Delta_v H$		52.5	298		[1981SHI/SAI]
C ₈ H ₁₈ S ₂	[1191-62-4]	1,8-octanedithiol				
	$\Delta_v H$	(405–543)	60.9	420	A	[1987STE/MAL, 1943HAL/REI, 1999DYK/SVO]
C ₈ H ₁₉ N	[20810-06-4]	N-butyl isobutylamine				
	$\Delta_v H$	(313–423)	41.2	328	A	[1987STE/MAL]
C ₈ H ₁₉ N	[111-92-2]	N,N-dibutylamine				
	$\Delta_v H$	(343–479)	46.0	358	A	[1987STE/MAL]
	$\Delta_v H$		46.0 ± 0.1	343	C	[1979PET/MAJ]
	$\Delta_v H$		44.8 ± 0.1	358	C	[1979PET/MAJ]
	$\Delta_v H$	(291–305)	48.1	298		[1971LEB/KAT2]
	$\Delta_v H$		49.4 ± 0.1	298	C	[1969WAD]
C ₈ H ₁₉ N	[110-93-3]	N,N-diisobutylamine				
	$\Delta_v H$	(291–305)	39.3	298		[1971LEB/KAT2]
	$\Delta_v H$	(273–333)	43.1 ± 0.3	298	I	[1969FRA/WAT]
C ₈ H ₁₉ N	[na]					
	$\Delta_v H$	(268–413)	43.8	283	A	[1987STE/MAL, 1947STU]
C ₈ H ₁₉ N	[626-23-3]	N,N-di-sec-butylamine				
	$\Delta_v H$	(273–333)	41.3 ± 0.3	298	I	[1969FRA/WAT]
C ₈ H ₁₉ N	[104-75-6]	2-ethylhexylamine				
	$\Delta_v H$	(341–447)	44.8	356	A	[1987STE/MAL]
C ₈ H ₁₉ N	[111-86-4]	octylamine				
	$\Delta_v H$	(343–494)	54.8 ± 0.5	298	EB	[1996STE/CHI3]
	$\Delta_v H$	(323–373)	54.6	298	CGC	[1995CHI/HOS]
C ₈ H ₁₉ N	[na]					
	$\Delta_v H$	(308–453)	50.8	323	A	[1987STE/MAL]
C ₈ H ₁₉ O ₂ PS ₃	[298-04-4]	O,O-diethyl-S-[2-(ethylthio)ethyl] dithiophosphate				
	$\Delta_v H$	(283–401)	76.7	298	A	[1987STE/MAL, 1999DYK/SVO]
C ₈ H ₁₉ O ₂ PS ₃	[2253-44-3]	O,O'-dibutyl dithiophosphate				
	$\Delta_v H$		81.8	298		[2008SAG/SAF]
C ₈ H ₁₉ O ₃ P	[1809-19-4]	dibutyl phosphite				
	$\Delta_v H$	(298–438)	37.8	313	A	[1987STE/MAL]
C ₈ H ₁₉ O ₃ P	[na]	diisopropyl ethylphosphonate				
	$\Delta_v H$		60.7 ± 4.2			[1956NEA/WIL, 1982PIL/SKI]
C ₈ H ₁₉ O ₃ PS ₂	[298-03-3]	O,O-diethyl-O-[2-(ethylthio)ethyl] thiophosphate				
	$\Delta_v H$	(283–411)	78.7	298	A	[1987STE/MAL, 1999DYK/SVO]
C ₈ H ₁₉ O ₃ PS ₂	[126-75-0]	O,O-diethyl-S-[2-(ethylthio)ethyl] thiophosphate				

TABLE 7. Phase change enthalpies of C₇ to C₈ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(283–401)	76.4	298	A	[1987STE/MAL, 1999DYK/SVO]
C ₈ H ₂₀ ClN	[6287-40-7]	dibutylammonium chloride				
	$\Delta_v H$	(553–563)	116.7	558	A	[1987STE/MAL, 1999DYK/SVO]
C ₈ H ₂₀ N ₂	[373-44-4]	octane-1,8-diamine				
	$\Delta_{\text{fus}} H$		50.51	324.9	DSC	[2006KHI/DAH2]
	$\Delta_{\text{fus}} H$		50.98	324.8	DSC	[2002DAL/DEL]
C ₈ H ₂₀ N ₂	[4267-00-9]	tetraethylhydrazine				
	$\Delta_v H$	(308–368)	33.4	323	A	[1987STE/MAL]
C ₈ H ₂₀ N ₂	[97-84-7]	N,N,N',N'-tetramethyl-1,3-butanediamine				
	$\Delta_v H$	(273–363)	49.2	288		[2002DAH/MOK]
	$\Delta_v H$	(335–439)	42.7	350	A	[1987STE/MAL]
C ₈ H ₂₀ N ₂ O ₂ S	[2832-49-7]	N,N,N',N'-tetraethylsulfamide				
	$\Delta_v H$	(407–528)	59.1	422	A	[1987STE/MAL]
C ₈ H ₂₀ N ₂ O ₃	[na]	<i>tris</i> (2-hydroxyethyl)ethylenediamine				
	$\Delta_v H$	(373–472)	90.0		GS	[1998ABD/MEI]
C ₈ H ₂₀ O ₅ P ₂ S ₂	[3689-24-5]	dithiopyrophosphoric acid, tetraethyl ester				
	$\Delta_v H$	(293–409)	80.6	308	A	[1987STE/MAL, 1999DYK/SVO]
C ₈ H ₂₀ O ₇ P ₂	[107-49-3]	pyrophosphoric acid, tetraethyl ester				
	$\Delta_v H$	(283–411)	82.2	298	A	[1987STE/MAL]
C ₈ H ₂₃ N ₅	[112-57-2]	tetraethylene pentamine				
	$\Delta_v H$	(464–615)	71.3	478	A	[1987STE/MAL]
C ₈ H ₂₄ N ₄ O ₃ P ₂	[152-16-9]	pyrophosphoric acid <i>tetrakis</i> (dimethylamide)				
	$\Delta_v H$	(273–415)	65.5	288	A	[1987STE/MAL]

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₉ F ₁₆	[75240-06-1]	<i>trans</i> -perfluorohydrindane				
	$\Delta_v H$		45.2 ± 0.1	298	C	[1996VAR/DRU]
C ₉ F ₁₆	[75262-87-2]	<i>cis</i> -perfluorobicyclo[4.3.0]nonane				
	$\Delta_{\text{us}} H$	(5–350)	8.76	200.6		
	$\Delta_{\text{us}} H$	(5–350)	1.27	245.6		
	$\Delta_{\text{fus}} H$	(5–350)	2.72	291.3	AC	[1998VAR/DRU]
C ₉ F ₁₆	[75240-06-1]	<i>trans</i> -perfluorobicyclo[4.3.0]nonane				
	$\Delta_{\text{us}} H$	(5–350)	8.91	236.6		
	$\Delta_{\text{fus}} H$	(5–350)	2.63	248.1	AC	[1998VAR/DRU]
C ₉ F ₁₇ NO ₃ S	[34834-20-3]	perfluoro-1-octanesulfonylisocyanate				
	$\Delta_v H$	(324–470)	67.7	339	A	[1987STE/MAL, 1999DYK/SVO]
C ₉ F ₁₈	[374-59-4]	perfluoro(propyl)cyclohexane				
	$\Delta_v H$	(321–396)	40.4	336		[1999DYK/SVO]
	$\Delta_v H$		43.1 ± 0.1	298	C	[1996VAR/DRU]
	$\Delta_v H$		43.1 ± 0.5	298	EB	[1981VAR/BUL]
	$\Delta_v H$		43.1 ± 0.1	298	C	[1981VAR/BUL]
C ₉ F ₁₈	[423-02-9]	perfluoro(isopropyl)cyclohexane				
	$\Delta_v H$		46.7 ± 0.1	298	C	[1996VAR/DRU]
C ₉ F ₁₈ N ₂	[34451-14-4]	1,1,1,3,3,3-hexafluoro-N,N'-bis[2,2,2-trifluoro-1-(trifluoromethyl)-ethylidene]-2,2-propanediamine				
	$\Delta_v H$	(314–381)	35.5	329	A	[1987STE/MAL, 1972SWI/SHR]
C ₉ F ₁₈ O ₃	[40719-69-5]	carbonic acid, bis[1,1,1,3,3,3-hexafluoro-2-(trifluoromethyl)-2-propyl ester				
	$\Delta_v H$	(316–358)	39.7	331	A	[1987STE/MAL, 1975WAL/DES2]
C ₉ F ₁₉ NO	[54120-06-8]	2,2,2-trifluoro-N-[1,2,2,2-tetrafluoro-1-(trifluoromethyl)ethyl]ethanimidic acid, 1,2,2,2-tetrafluoro-1-(trifluoromethyl)ethyl ester				
	$\Delta_v H$		37.6	385		[1975PET/SHR]
C ₉ F ₂₀	[375-96-2]	perfluorononane				
	$\Delta_v H$	(288–333)	45.3 ± 0.2	298		[2005DIA/GON]
	$\Delta_v H$	(387–524)	32.8	402	A	[1987STE/MAL, 1967BER/WES, 1999DYK/SVO]
C ₉ F ₂₁ N	[514-03-4]	perfluoro-N-methyl-N,N-dibutylamine				
	$\Delta_v H$	(339–407)	48.8 ± 0.8	298	EB	[1995VAR/DRO]
	$\Delta_v H$		48.2 ± 0.1	298	C	[1995VAR/DRO]
C ₉ F ₂₁ N	[338-83-3]	<i>tris</i> (heptafluoropropyl)amine				
	$\Delta_v H$		46.6 ± 0.3	298	C	[1995VAR/DRO]
	$\Delta_v H$	(329–403)	46.9 ± 0.7	298	EB	[1995VAR/DRO]
	$\Delta_v H$	(333–403)	40.6	348	A	[1987STE/MAL]
C ₉ H ₂ Cl ₆ O ₃	[7365-74-4]	4,5,6,7,8,8-hexachloro-3a,4,7,7a-tetrahydro-4,7-methanoisobenzofuran-1,3-dione				
	$\Delta_{\text{us}} H$		10.64	385.4		
	$\Delta_{\text{fus}} H$		2.67	506	DSC	[1984WEI/LEF]
C ₉ H ₄ Cl ₃ NO ₂ S	[133-07-3]	2-[(trichloromethyl)thio]-1 <i>H</i> -isoindole-1,3(2 <i>H</i>)-dione				
	$\Delta_{\text{fus}} H$		35.49	454.2	DSC	[1991ACR, 1990DON/DRE]
C ₉ H ₄ Cl ₄ O ₄	[887-54-7]	methyl tetrachloroterephthalic acid ester				
	$\Delta_{\text{fus}} H$		16.89	444.3	DSC	[1990DON/DRE]
C ₉ H ₄ Cl ₈ O	[76341-69-0]	1,3,4,5,6,7,8,8-octachloro-2,3,3a,4,7,7a-hexahydro-4,7-methanoisobenzofuran				
	$\Delta_{\text{fus}} H$		25.94	395.4	DSC	[1969PLA/GLA]
C ₉ H ₄ O ₅	[552-30-7]	trimellitic acid anhydride				
	$\Delta_{\text{fus}} H$		10.46	385		[1996DOM/HEA]

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(558–596)	65.6	573	A	[1987STE/MAL]
C ₉ H ₅ BrClNO	[7640-33-7]	7-bromo-5-chloro-8-hydroxyquinoline				
	$\Delta_{\text{sub}} H$	(353–368)	110.1 ± 0.8	361	ME	[1992RIB/MON2]
	$\Delta_{\text{sub}} H$	(353–368)	113.2 ± 0.8	298	ME	[1992RIB/MON2]
C ₉ H ₅ Br ₂ NO	[521-74-4]	5,7-dibromo-8-hydroxyquinoline				
	$\Delta_{\text{sub}} H$	(365–380)	113.6 ± 1.3	372	ME	[1992RIB/MON2]
	$\Delta_{\text{sub}} H$	(365–380)	117.3 ± 1.3	298	ME	[1992RIB/MON2]
	$\Delta_{\text{sub}} H$	(363–393)	94.1			[1963HOR/WEN]
C ₉ H ₅ ClINO	[130-26-7]	5-chloro-7-iodo-8-hydroxyquinoline				
	$\Delta_{\text{sub}} H$	(359–378)	111.3 ± 0.4	368	ME	[1992RIB/MON2]
	$\Delta_{\text{sub}} H$	(359–378)	114.8 ± 0.4	298	ME	[1992RIB/MON2]
C ₉ H ₅ ClINO	[35048-13-6]	5-iodo-7-chloro-8-hydroxyquinoline				[1963HOR/WEN]
	$\Delta_{\text{sub}} H$	(383–414)	131			
C ₉ H ₅ Cl ₂ N ₂ O ₂	[15166-26-4]	5-chloro-2,4-diisocyanato-1-methylbenzene				
	$\Delta_v H$	(373–433)	66.7	388	A	[1987STE/MAL]
	$\Delta_v H$	(373–433)	60.2 ± 0.2	403		[1972STR/NOV]
C ₉ H ₅ Cl ₂ N	[86-98-6]	4,7-dichloroquinoline				
	$\Delta_{\text{sub}} H$		89.5 ± 2.3	298	C	[2006RIB/MAT]
C ₉ H ₅ Cl ₂ NO	[773-76-2]	5,7-dichloro-8-hydroxyquinoline				
	$\Delta_{\text{sub}} H$	(351–366)	106.3 ± 0.7	358	ME	[1992RIB/MON2]
	$\Delta_{\text{sub}} H$	(351–366)	109.3 ± 0.7	298	ME	[1992RIB/MON2]
	$\Delta_{\text{sub}} H$	(363–393)	92.9			[1963HOR/WEN]
C ₉ H ₅ Cl ₃ N ₄	[101-05-3]	4,6-dichloro-N-(2-chlorophenyl)-1,3,5-triazin-2-amine				
	$\Delta_{\text{fus}} H$		31.48	431	DSC	[1990DON/DRE]
C ₉ H ₅ I ₂ NO	[83-73-8]	5,7-diiodo-8-hydroxyquinoline				
	$\Delta_{\text{sub}} H$	(389–404)	121.9 ± 0.8	396	ME	[1992RIB/MON2]
	$\Delta_{\text{sub}} H$	(389–404)	126.8 ± 0.8	298	ME	[1992RIB/MON2]
	$\Delta_{\text{sub}} H$	(403–423)	110.9			[1963HOR/WEN]
C ₉ H ₆ BrN	[5332-24-1]	3-bromoquinoline				
	$\Delta_v H$		70.7 ± 2.3	298	C	[2008RIB/AMA]
C ₉ H ₆ ClN	[612-62-4]	2-chloroquinoline				
	$\Delta_{\text{sub}} H$		84.3 ± 2.6	298	C	[2006RIB/MAT]
C ₉ H ₆ ClN	[611-35-8]	4-chloroquinoline				
	$\Delta_{\text{sub}} H$		78.6 ± 1.7	298	C	[2006RIB/MAT]
C ₉ H ₆ ClN	[612-57-7]	6-chloroquinoline				
	$\Delta_{\text{sub}} H$		80.8 ± 1.9	298	C	[2006RIB/MAT]
C ₉ H ₆ ClNO	[130-16-5]	5-chloro-8-hydroxyquinoline				
	$\Delta_{\text{sub}} H$	(317–327)	97.5 ± 0.9	322	ME	[1992RIB/MON2]
	$\Delta_{\text{sub}} H$	(317–327)	98.7 ± 0.9	298	ME	[1992RIB/MON2]
C ₉ H ₆ ClNO ₂	[17564-64-6]	N-chloromethylphthalimide				
	$\Delta_{\text{sub}} H$		103.5 ± 1.1	298	C	[2007RIB/SAN3]
	$\Delta_{\text{sub}} H$	(323–343)	103.6 ± 0.9	298	ME	[2007RIB/SAN3]
C ₉ H ₆ Cl ₂ N ₂ O ₃	[20354-26-1]	2-(3,4-dichlorophenyl)-4-methyl-1,2,4-oxadiazolidine-3,5-dione				
	$\Delta_{\text{fus}} H$		29.5	396.3	DSC	[1990DON/DRE]
C ₉ H ₆ Cl ₂ O ₃	[17812-11-2]	2,3-dichloro-5-norbornene-2,3-dicarboxylic anhydride				
	$\Delta_{\text{trs}} H$		17.94	339.1		
	$\Delta_{\text{fus}} H$		5.36	457.6	DSC	[1984WEI/LEF]

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₉ H ₆ Cl ₆ O ₃ S	[959-98-8]	endosulfan I				
	$\Delta_{\text{fus}}H$		10.0	380	DSC	[1990DON/DRE]
	Δ_vH	(343–453)	80.4	398	GC	[1990HIN/BID2]
C ₉ H ₆ Cl ₆ O ₃ S	[33213-65-9]	endosulfan II				
	Δ_vH	(343–453)	82.4	398	GC	[1990HIN/BID2]
C ₉ H ₆ Cl ₆ O ₃ S	[115-29-7]	6,7,8,9,10,10-hexachloro-1,5,5a,6,9,9a-hexahydro-6,9-methano-2,4,3-benzodioxathiepin-3-oxide (endosulfan)				
	$\Delta_{\text{fus}}H$		16.52	368	DSC	[2000ROD/VEC]
C ₉ H ₆ Cl ₆ O ₄ S	[1031-07-8]	endosulfan sulfate				
	Δ_vH	(343–453)	85.6	398	GC	[1990HIN/BID2]
C ₉ H ₆ Cl ₆ O ₄ S	[1031-07-6]	6,7,8,9,10,10-hexachloro-6,9-methano-2,4,3-benzodioxathiapin-3,3-dioxide				
	$\Delta_{\text{fus}}H$		21.66	419.7	DSC	[1990DON/DRE]
C ₉ H ₆ INO	[13207-63-1]	5-iodo-8-hydroxyquinoline				
	$\Delta_{\text{sub}}H$	(363–393)	118.8		ME	[1963HOR/WEN]
C ₉ H ₆ N ₂ O ₂	[584-84-9]	2,4-toluene diisocyanate				
	Δ_vH	(373–530)	59.7	388	A	[1987STE/MAL]
	Δ_vH	(393–530)	59.5	408	A	[1987STE/MAL]
	Δ_vH	(373–530)	61.3	388	I	[1975FRE/ADA]
	Δ_vH	(373–433)	57.7 ± 0.2	403		[1972STR/NOV]
C ₉ H ₆ N ₂ O ₂	[91-08-7]	2,6-toluene diisocyanate				
	Δ_vH	(373–463)	60.4	388	A	[1987STE/MAL]
C ₉ H ₆ N ₂ O ₂	[607-34-1]	5-nitroquinoline				
	$\Delta_{\text{sub}}H$	(310–324)	93.2 ± 0.7	317	ME	[1997RIB/MAT5]
	$\Delta_{\text{sub}}H$	(310–324)	94.2 ± 0.7	298	ME	[1997RIB/MAT5]
C ₉ H ₆ N ₂ O ₂	[613-50-3]	6-nitroquinoline				
	$\Delta_{\text{sub}}H$	(336–350)	101.5 ± 1.0	343	ME	[1997RIB/MAT5]
	$\Delta_{\text{sub}}H$	(336–350)	103.8 ± 1.0	298	ME	[1997RIB/MAT5]
C ₉ H ₆ N ₂ O ₂	[607-35-2]	8-nitroquinoline				
	$\Delta_{\text{sub}}H$	(338–352)	104.3 ± 0.9	345	ME	[1997RIB/MAT5]
	$\Delta_{\text{sub}}H$	(338–352)	106.7 ± 0.9	298	ME	[1997RIB/MAT5]
C ₉ H ₆ N ₂ O ₃	[4008-48-4]	5-nitro-8-hydroxyquinoline				
	$\Delta_{\text{fus}}H$		19.61	455.2	DSC	[2010GAO/LIN]
	$\Delta_{\text{fus}}H$		24.7	453.2		[2001ZOR/COS]
	$\Delta_{\text{sub}}H$	(413–453)	81.66		TGA	[2010GAO/LIN]
	$\Delta_{\text{sub}}H$	(413–453)	86.14	298	TGA	[2010GAO/LIN]
						Note: The authors of [2010GAO/LIN] did note that their experimental value differed significantly from the earlier published literature values.
$\Delta_{\text{sub}}H$	(352–362)	114.1 ± 2.2	298	ME	[1989RIB/MON]	
$\Delta_{\text{sub}}H$		111.2 ± 3.0	298	C	[1989RIB/MON]	
C ₉ H ₆ N ₄ O ₂	[23190-84-3]	3-amino-2-quinoxalinecarbonitrile 1,4-dioxide				
	$\Delta_{\text{sub}}H$		139.7 ± 3.7	298	ME	[2004RIB/GOM]
C ₉ H ₆ O ₂	[91-64-5]	coumarin				
	$\Delta_{\text{fus}}H$		18.63	342.3	DSC	[2009MAT/SOU4]
	$\Delta_{\text{fus}}H$		19.14	342.1		[1996DOM/HEA]
	$\Delta_{\text{sub}}H$		95.4 ± 2.6	298	C	[2009MAT/SOU4]
	$\Delta_{\text{sub}}H$		83.1	298	C	[1991ELW/SAB, 1992SAB/WAT]
	$\Delta_{\text{sub}}H$	(293–353)	86.2	323	ME	[1953SER/VOI, 1960JON, 1987STE/MAL]
Δ_vH	(379–463)	63.2	394	A	[1987STE/MAL, 1947STU]	

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₉ H ₆ O ₂	[491-38-3]	chromone				
	$\Delta_{\text{fus}}H$		15.44	329.9	DSC	[2009MAT/SOU4]
	$\Delta_{\text{fus}}H$		17.31	330.3		[1991ACR, 1996DOM/HEA]
	$\Delta_{\text{sub}}H$		86.5 ± 1.1	298	C	[2009MAT/SOU4]
C ₉ H ₆ O ₂	[606-23-5]	1,3-indandione				
	$\Delta_{\text{fus}}H$		21.8	401.5	DSC	[2007MAT/MIR]
C ₉ H ₆ O ₆	[554-95-0]	1,3,5-benzenetricarboxylic acid (553–593)	159.4	573	GS	[1987STE/MAL, 1962KRA/BER]
	$\Delta_{\text{sub}}H$					
C ₉ H ₆ S ₃	[3445-76-9]	5-phenyl-1,2-dithiole-3-thione (363–373)	117.4 ± 0.4			[1972GEI/RAU]
	$\Delta_{\text{sub}}H$		123.3 ± 0.4	298		[1972GEI/RAU]
C ₉ H ₆ S ₃	[3445-76-9]	5-phenyldithiolethione				
	$\Delta_{\text{fus}}H$		26.27	398	DSC	[1999DOL/LEC]
C ₉ H ₇ BrO ₂	[14473-91-7]	3-bromo- <i>trans</i> -cinnamic acid				
	$\Delta_{\text{fus}}H$		31.06	443.2	DSC	[2001AHN/HAR]
C ₉ H ₇ Cl ₃ O ₃	[93-72-1]	2-(2,4,5-trichlorophenoxy)propanoic acid				
	$\Delta_{\text{fus}}H$		39.58	450.6	DSC	[1990DON/DRE]
C ₉ H ₇ Cl ₃ O ₃	[1928-37-6]	2,4,5-trichlorophenoxyacetic acid, methyl ester				
	$\Delta_{\text{fus}}H$		30.46	361.9	DSC	[1990DON/DRE]
	Δ_vH	(444–573)	76.9	459	A	[1987STE/MAL, 1999DYK/SVO]
C ₉ H ₇ F ₃ O ₂	[1736-09-0]	trifluoroacetic acid, 3-tolyl ester (363–439)	47.4	378	A,EB	[1987STE/MAL, 1969SHE/LAN, 1999DYK/SVO]
	Δ_vH					
C ₉ H ₇ F ₃ O ₂	[1813-29-2]	trifluoroacetic acid, 4-tolyl ester (365–442)	47.8	380	A,EB	[1987STE/MAL, 1969SHE/LAN, 1999DYK/SVO]
	Δ_vH					
C ₉ H ₇ N	[119-65-3]	isoquinoline				
	$\Delta_{\text{fus}}H$		13.54	299.6		[1996DOM/HEA]
	Δ_vH		53.3		GC	[1996GOV/RUT]
	Δ_vH	(313–566)	58.9 ± 0.1	320	IP,EB	[1988STE/ARC]
	Δ_vH	(313–566)	56.4 ± 0.1	360	IP,EB	[1988STE/ARC]
	Δ_vH	(313–566)	54.1 ± 0.1	400	IP,EB	[1988STE/ARC]
	Δ_vH	(313–566)	51.7 ± 0.1	440	IP,EB	[1988STE/ARC]
	Δ_vH	(313–566)	49.4 ± 0.2	480	IP,EB	[1988STE/ARC]
	Δ_vH	(313–566)	47.0 ± 0.3	520	IP,EB	[1988STE/ARC]
Δ_vH	(439–517)	51.0	454	A,EB	[1987STE/MAL, 1961MAL]	
C ₉ H ₇ N	[91-22-5]	quinoline				
	$\Delta_{\text{ms}}H$		0.07	220		
	$\Delta_{\text{fus}}H$		10.66	258.4		[1996DOM/HEA]
	Δ_vH		53.3		GC	[1996GOV/RUT]
	Δ_vH	(573–668)	46.9	588	DSC	[1996BAC/GRZ]
	Δ_vH	(504–616)	46.5	519		[1992LEE/CHE]
	Δ_vH	(298–559)	57.9 ± 0.1	320	IP,EB	[1988STE/ARC]
	Δ_vH	(298–559)	55.5 ± 0.1	360	IP,EB	[1988STE/ARC]
	Δ_vH	(298–559)	53.1 ± 0.1	400	IP,EB	[1988STE/ARC]
	Δ_vH	(298–559)	50.7 ± 0.1	440	IP,EB	[1988STE/ARC]

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(298–559)	48.4 ± 0.2	480	IP,EB	[1988STE/ARC]
	$\Delta_v H$	(298–559)	46.0 ± 0.3	520	IP,EB	[1988STE/ARC]
	$\Delta_v H$	(473–548)	U65.4	488		[1987KLA/MOH]
	$\Delta_v H$	(463–794)	46.1	478	A	[1987STE/MAL]
	$\Delta_v H$	(286–309)	58.1	298	GS	[1980VAN/PRA]
	$\Delta_v H$	(433–511)	49.2	448	EB	[1987STE/MAL, 1961MAL]
C₉H₇NO	[59-31-4]	2-hydroxyquinoline				
	$\Delta_{\text{sub}}H$	(375–390)	115.2 ± 0.6	383	ME	[1990RIB/MAT]
	$\Delta_{\text{sub}}H$	(375–390)	119.4 ± 0.6	298	ME	[1990RIB/MAT]
C₉H₇NO	[611-36-9]	4-hydroxyquinoline				
	$\Delta_{\text{sub}}H$	(415–433)	128.8 ± 1.1	424	ME	[1990RIB/RIB]
	$\Delta_{\text{sub}}H$	(415–433)	135.1 ± 1.1	298	ME	[1990RIB/RIB]
C₉H₇NO	[148-24-3]	8-hydroxyquinoline				
	$\Delta_{\text{fus}}H$		40.3	345.7	AC	[2008WAN/TAN]
	$\Delta_{\text{fus}}H$		22.1	346.8		[2001ZOR/COS]
	$\Delta_{\text{sub}}H$	(293–303)	89.5 ± 0.9	298	ME	[1989RIB/MON]
	$\Delta_{\text{sub}}H$		89.0 ± 1.4	298	C	[1989RIB/MON]
	$\Delta_{\text{sub}}H$	(308–328)	108.8 ± 1.7		ME	[1963HOR/WEN, 1970COX/PIL, 1987STE/MAL]
C₉H₇NO	[491-30-5]	1-hydroxyisoquinoline				
	$\Delta_{\text{sub}}H$		113.6 ± 2.2	298	C	[2005RIB/MAT]
C₉H₇NO	[2439-04-5]	5-hydroxyisoquinoline				
	$\Delta_{\text{sub}}H$		109.6 ± 2.1	298	C	[2005RIB/MAT]
C₉H₇NO	[614-16-4]	Ω -cyanoacetophenone				
	$\Delta_{\text{sub}}H$	(318–333)	99.8	325.5		[1987STE/MAL]
	$\Delta_{\text{sub}}H$		92.5 ± 4.2		ME	[1969LEB/DNE, 1977PED/RYL]
C₉H₇NO₂	[5154-02-9]	1,5-dihydroxyisoquinoline				
	$\Delta_{\text{sub}}H$		123.6 ± 2.2		C	[2005RIB/FON]
C₉H₇NO₂	[550-44-7]	N-methylphthalimide				
	$\Delta_{\text{sub}}H$	(298–316)	91.1 ± 0.5	307	ME	[1997ROU/JIM]
	$\Delta_{\text{sub}}H$	(298–316)	91.1 ± 0.5	298	ME	[1997ROU/JIM]
C₉H₇NO₂	[2058-74-4]	1-methyl-1 <i>H</i> -indole-2,3-dione (N-methylisatin)				
	$\Delta_{\text{fus}}H$		19.5	403.3	DSC	[2003MAT/MIR2]
	$\Delta_{\text{sub}}H$		105.6 ± 3.3	298	C	[2003MAT/MIR2]
C₉H₇N₃O₂	[35975-00-9]	5-amino-6-nitroquinoline				
	$\Delta_{\text{sub}}H$	(400–424)	130.7 ± 0.8	412	ME	[1998RIB/CAR]
	$\Delta_{\text{sub}}H$	(400–424)	136.4 ± 0.8	298	ME	[1998RIB/CAR]
C₉H₇N₃S	[41814-78-2]	5-methyl-1,2,4-triazolo[3,4- <i>b</i>]benzothiazole				
	$\Delta_{\text{fus}}H$		24.07	460.2	DSC	[1990DON/DRE]
C₉H₈	[95-13-6]	indene				
	$\Delta_{\text{fus}}H$		10.2	271.7		[1996DOM/HEA]
	$\Delta_v H$	(369–457)	45.3	384	A	[1987STE/MAL]
	$\Delta_v H$	(289–455)	43.6	304	A	[1987STE/MAL, 1947STU]
	$\Delta_v H$	(329–454)	43.9	392		[1942BUR]
C₉H₈Cl₂O₃	[6597-78-0]	methyl 3,6-dichloro-2-methoxybenzoate				
	$\Delta_{\text{fus}}H$		18.49	304.6		[1991ACR]
C₉H₈Cl₂O₃	[120-36-5]	2-(2,4-dichlorophenoxy)propanoic acid				

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{fus}}H$		32.0	391.3	DSC	[2005VEC/BRU]
	$\Delta_{\text{fus}}H$		30.43	389.2		[1991ACR]
	$\Delta_{\text{sub}}H$		116 ± 6	298	DSC	[2005VEC/BRU]
	$\Delta_{\text{sub}}H$	(343–375)	128 ± 2	359	TE	[2005VEC/BRU]
	$\Delta_{\text{sub}}H$	(343–375)	130 ± 3	298	TE	[2005VEC/BRU]
C₉H₈Cl₂O₃	[1928-38-7]	2,4-dichlorophenoxyacetic acid, methyl ester				
	$\Delta_{\text{fus}}H$		20.0	313.4	DSC	[2005VEC/BRU]
	$\Delta_{\text{fus}}H$		25.1	315.4	DSC	[1969PLA/GLA]
	Δ_vH	(403–548)	68.0	418	A	[1987STE/MAL, 1999DYK/SVO]
C₉H₈Cl₂O₄	[76330-06-8]	2,6-dichlorosyringaldehyde				
	Δ_vH	(293–323)	82.2	308	CGC	[1999LEI/WAN2]
C₉H₈Cl₃NO₃	[75907-45-8]	2,2,4-trichloro-5-(4-morpholinyl)-4-cyclopentene-1,3-dione				
	Δ_vH	(453–483)	79.6	468	GC	[1980SHA/SAD]
C₉H₈N₂	[1126-00-7]	1-phenylpyrazole				
	Δ_vH		70.2 ± 3.4	298	C	[2000RIB/RIB2]
C₉H₈N₂	[7164-98-9]	1-phenylimidazole				
	Δ_vH		84.6 ± 3.7	298	C	[2000RIB/RIB2]
C₉H₈N₂	[670-96-2]	2-phenylimidazole				
	$\Delta_{\text{fus}}H$		17.81	420	DSC	[2007SIF/AIT]
C₉H₈N₂	[580-17-6]	3-aminoquinoline				
	$\Delta_{\text{sub}}H$	(329–345)	101.1 ± 0.9	337	ME	[1993RIB/MAT]
	$\Delta_{\text{sub}}H$	(329–345)	103.1 ± 0.9	298	ME	[1993RIB/MAT]
	$\Delta_{\text{sub}}H$		104.8 ± 4.8	298	C	[1993RIB/MAT]
C₉H₈N₂	[611-34-7]	5-aminoquinoline				
	$\Delta_{\text{sub}}H$	(329–349)	102.9 ± 0.7	339	ME	[1993RIB/MAT]
	$\Delta_{\text{sub}}H$	(329–349)	105.0 ± 0.7	298	ME	[1993RIB/MAT]
	$\Delta_{\text{sub}}H$		103.3 ± 3.4	298	C	[1993RIB/MAT]
C₉H₈N₂	[580-15-4]	6-aminoquinoline				
	$\Delta_{\text{sub}}H$	(333–349)	103.6 ± 1.0	341	ME	[1993RIB/MAT]
	$\Delta_{\text{sub}}H$	(333–349)	105.7 ± 1.0	298	ME	[1993RIB/MAT]
C₉H₈N₂	[578-66-5]	8-aminoquinoline				
	$\Delta_{\text{sub}}H$	(296–314)	93.0 ± 0.5	305	ME	[1993RIB/MAT]
	$\Delta_{\text{sub}}H$	(296–314)	93.33 ± 0.5	298	ME	[1993RIB/MAT]
C₉H₈N₂O	[14003-34-0]	2-methyl-3-hydroxyquinoxaline				
	$\Delta_{\text{fus}}H$		33.4	522.9	DSC	[2000MON/HIL2]
	$\Delta_{\text{sub}}H$	(375–391)	117.2 ± 0.4	383	ME	[2000MON/HIL2]
	$\Delta_{\text{sub}}H$	(375–391)	119.7 ± 2.8	298	ME	[2000MON/HIL2]
	$\Delta_{\text{sub}}H$		123.0 ± 4.4	298	C	[2000RIB/MAT]
C₉H₈N₂O₂	[6639-86-7]	2-methylquinoxaline-1,4-dioxide				
	$\Delta_{\text{sub}}H$		107.0 ± 6.2	298	C	[1997ACR/POW]
C₉H₈N₂O₂	[5972-09-8]	3-methylaminophthalimide				
	$\Delta_{\text{sub}}H$	(402–450)	104.9	417	RG	[1987STE/MAL, 1956KLO]
C₉H₈O	[83-33-0]	1-indanone				
	$\Delta_{\text{fus}}H$		17.6	314.1	DSC	[2007MAT/MIR]
	$\Delta_{\text{fus}}H$		17.78	312.9	DSC	[1998VER4]
	$\Delta_{\text{sub}}H$		78.7 ± 2.8	298	C	[2007MAT/MIR]

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
			83.5 ± 0.7	298	GS	[1998VER4]
			60.3 ± 0.4		GS	[1998GUD/TOR]
C ₉ H ₈ O	[615-13-4]	2-indanone	16.89	330	DSC	[2007MAT/MIR]
			78.3 ± 1.1	298	C	[2007MAT/MIR]
C ₉ H ₈ O	[104-55-2]	3-phenyl-2-propenal (cinnamaldehyde)	62.4	298	GC	[2002VAN/PAR]
			51.7	444	TGA	[2002HAZ/DOL]
			58.2	364	A	[1987STE/MAL, 1947STU]
			72.7	363	A	[1987STE/MAL]
C ₉ H ₈ O ₂	[140-10-3]	<i>trans</i> cinnamic acid	22.21	406.1	DSC	[2008MOT/QUE]
			22.6	404.8		[2004SHA/JAM]
			22.63	406.2		[1991ACR]
			107.1 ± 0.8	298	ME	[1999MON/HIL]
			73.9	445	A	[1987STE/MAL]
C ₉ H ₈ O ₂	[102-94-3]	allocinnamic acid (<i>cis</i> cinnamic acid)	16.95	341.2		[1991ACR]
C ₉ H ₈ O ₂	[39869-70-0]	7,7-dimethoxynorborane	49.0	339	EB	[1994WIB/MOR]
C ₉ H ₈ O ₂	[119-84-6]	3,4-dihydrocoumarin	69.9 ± 0.5	298	C	[2009MAT/SOU2]
C ₉ H ₈ O ₂	[491-37-2]	chromanone	16.7	312.3	DSC	[2009MAT/SOU]
			84.6 ± 1.3	298	C	[2009MAT/SOU]
C ₉ H ₈ O ₂	[4385-35-7]	3-isochromanone	18.3	355.9	DSC	[2009MAT/SOU]
			97.3 ± 1.4	298	C	[2009MAT/SOU]
C ₉ H ₈ O ₂ S	[2525-42-0]	phenyl propadienyl sulfone	105.4 ± 2.5			[1969MAC/STE, 1970COX/PIL]
C ₉ H ₈ O ₂ S	[2525-41-9]	phenyl prop-1-ynyl sulfone	95.4 ± 2.5		B	[1969MAC/STE, 1970COX/PIL]
C ₉ H ₈ O ₂ S	[2525-40-8]	phenyl prop-2-ynyl sulfone	$105. \pm 2.5$		B	[1969MAC/STE, 1970COX/PIL]
C ₉ H ₈ O ₃	[129-64-6]	<i>endo</i> -5-norbornene-2,3-dicarboxylic anhydride	15.73	367.2		[1967PIN/WIL]
			3.71	437.2		
			97 ± 4.2	298	MG	[1973ROG/QUA, 1977PED/RYL]
C ₉ H ₈ O ₃	[2746-19-2]	<i>exo</i> -5-norbornene-2,3-dicarboxylic anhydride	21.77	416.2	DSC	[1967PIN/WIL]
C ₉ H ₈ O ₃	[3162-29-6]	5-oxoethyl-1,3-benzodioxole	26.23	358.9	DSC	[2007MAT/SOU]
			104.4 ± 2.2	298	C	[2007MAT/SOU]

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₉ H ₈ O ₃	[29668-44-8]	2,3-dihydro-1,4-benzodioxin-6-carboxaldehyde				
	$\Delta_{\text{fus}}H$		19.44	324.4	DSC	[2008MAT/SOU2]
			98.2 ± 1.4	298	C	[2008MAT/SOU2]
C ₉ H ₈ O ₄	[331-39-5]	3,4-dihydroxycinnamic acid (caffeic acid)				
	$\Delta_{\text{sub}}H$	(409–424)	170.2 ± 4.6	411		[2006CHE/OJA]
C ₉ H ₈ O ₄	[1679-64-7]	monomethyl terephthalate				
	$\Delta_{\text{fus}}H$		37.68	92.5	DSC	[2005MON/SOU]
	$\Delta_{\text{sub}}H$	(363–381)	121.0 ± 0.5	372.3	ME	[2005MON/SOU]
	$\Delta_{\text{sub}}H$	(363–381)	124.1 ± 1.0	298	ME	[2005MON/SOU]
	$\Delta_{\text{sub}}H$	(433–493)	72.1	448		[1987STE/MAL]
	$\Delta_{\text{sub}}H$	(433–493)	82.8	473	GS	[1962KRA/BER]
			130.4		C	[1998MAK/KAB]
C ₉ H ₈ O ₄	[4376-18-5]	monomethyl phthalate				
	$\Delta_{\text{fus}}H$		21.63	357.5	DSC	[2005MON/SOU]
	$\Delta_{\text{sub}}H$	(335–355)	115.9 ± 0.6	345.3	ME	[2005MON/SOU]
			117.9 ± 0.8	298	ME	[2005MON/SOU]
C ₉ H ₈ O ₄	[1877-71-0]	monomethyl isophthalate				
	$\Delta_{\text{fus}}H$		36.5	466.7	DSC	[2005MON/SOU]
	$\Delta_{\text{sub}}H$	(359–379)	122.6 ± 0.7	369.2	ME	[2005MON/SOU]
			125.6 ± 1.0	298	ME	[2005MON/SOU]
C ₉ H ₈ O ₄	[50-78-2]	2-acetoxybenzoic acid (aspirin)				
	$\Delta_{\text{fus}}H$		29.17	409.2	DSC	[2004XU/SUN2]
	$\Delta_{\text{fus}}H$		31.01	412.7	DSC	[2001PER/BAU, 2003PER/BAU]
			29.8	414		[2000KIR]
C ₉ H ₈ O ₄	[2345-34-8]	4-acetoxybenzoic acid				
	$\Delta_{\text{fus}}H$		26.35	467.2		[1997YUA/ZHA]
C ₉ H ₈ O ₄	[2861-28-1]	1,3-benzodioxole-5-acetic acid (homopiperonylic acid)				
	$\Delta_{\text{fus}}H$		24.94	401.7		[2004MAT/MON]
	$\Delta_{\text{sub}}H$	(346–364)	120.1 ± 0.8	355	ME	[2004MAT/MON]
			122.9 ± 1.4	298	ME	[2004MAT/MON]
C ₉ H ₈ O ₄	[3663-80-7]	<i>(dl)</i> 1,4-benzodioxan-2-carboxylic acid				
	$\Delta_{\text{sub}}H$		117.8 ± 2.1	298	C	[2008MAT/MIR2]
C ₉ H ₉ BrO ₃	[na]	<i>(dl)</i> 2-(<i>p</i> -bromophenoxy)propanoic acid				
	$\Delta_{\text{fus}}H$		31.8	385		[1991CHI/BRA]
C ₉ H ₉ BrO ₃	[na]	<i>(d)</i> 2-(<i>p</i> -bromophenoxy)propanoic acid				
	$\Delta_{\text{fus}}H$		27.61	380		[1991CHI/BRA]
C ₉ H ₉ BrO ₃	[40620-67-5]	<i>(dl)</i> 3-(<i>m</i> -bromophenyl)-3-hydroxypropanoic acid				
	$\Delta_{\text{fus}}H$		26.78	349		[1991CHI/BRA]
C ₉ H ₉ BrO ₃	[40620-57-3]	<i>(d)</i> 3-(<i>m</i> -bromophenyl)-3-hydroxypropanoic acid				
	$\Delta_{\text{fus}}H$		23.85	350		[1991CHI/BRA]
C ₉ H ₉ BrO ₃	[40620-68-6]	<i>(dl)</i> 3-(<i>p</i> -bromophenyl)-3-hydroxypropanoic acid				
	$\Delta_{\text{fus}}H$		28.87	371		[1991CHI/BRA]
C ₉ H ₉ BrO ₃	[40620-58-4]	<i>(d)</i> 3-(<i>p</i> -bromophenyl)-3-hydroxypropanoic acid				
	$\Delta_{\text{fus}}H$		35.56	398		[1991CHI/BRA]
C ₉ H ₉ ClO ₃	[na]	<i>(dl)</i> 2-(<i>o</i> -chlorophenoxy)propanoic acid				

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		T _m (K)	Method	Reference
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)			
	$\Delta_{\text{fus}}H$		32.22	388		[1991CHI/BRA]
C ₉ H ₉ ClO ₃	[na]	(<i>d</i>) 2-(<i>o</i> -chlorophenoxy)propanoic acid				
	$\Delta_{\text{fus}}H$		26.78	369		[1991CHI/BRA]
C ₉ H ₉ ClO ₃	[101-10-0]	(<i>dl</i>) 2-(<i>m</i> -chlorophenoxy)propanoic acid				
	$\Delta_{\text{fus}}H$		33.05	386		[1991CHI/BRA]
C ₉ H ₉ ClO ₃	[na]	(<i>d</i>) 2-(<i>m</i> -chlorophenoxy)propanoic acid				
	$\Delta_{\text{fus}}H$		29.71	367.5		[1991CHI/BRA]
C ₉ H ₉ ClO ₃	[na]	(<i>dl</i>) 3-(<i>p</i> -chlorophenyl)-3-hydroxypropanoic acid				
	$\Delta_{\text{fus}}H$		29.71	357		[1991CHI/BRA]
C ₉ H ₉ ClO ₃	[40620-55-1]	(<i>d</i>) 3-(<i>p</i> -chlorophenyl)-3-hydroxypropanoic acid				
	$\Delta_{\text{fus}}H$		28.03	385		[1991CHI/BRA]
C ₉ H ₉ ClO ₃	[40620-64-2]	(<i>dl</i>) 3-(<i>m</i> -chlorophenyl)-3-hydroxypropanoic acid				
	$\Delta_{\text{fus}}H$		23.85	340		[1991CHI/BRA]
C ₉ H ₉ ClO ₃	[40620-54-0]	(<i>d</i>) 3-(<i>m</i> -chlorophenyl)-3-hydroxypropanoic acid				
	$\Delta_{\text{fus}}H$		28.03	368		[1991CHI/BRA]
C ₉ H ₉ ClO ₃	[94-74-6]	(4-chloro- <i>o</i> -tolylxy)acetic acid				
	$\Delta_{\text{fus}}H$		29.98	392.9	DSC	[1991ACR, 1990DON/DRE]
C ₉ H ₉ ClO ₄	[76341-69-0]	2-chlorosyringaldehyde (293–323)	77.7	308	CGC	[1999LEI/WAN2]
C ₉ H ₉ Cl ₂ NO	[709-98-8]	3',4'-dichloropropionanilide				
	$\Delta_{\text{fus}}H$		18.26	363.7	DSC	[1991ACR, 1990DON/DRE]
C ₉ H ₉ FO ₃	[40620-61-9]	(<i>dl</i>) 3-(<i>m</i> -fluorophenyl)-3-hydroxypropanoic acid				
	$\Delta_{\text{fus}}H$		20.5	290		[1991CHI/BRA]
C ₉ H ₉ FO ₃	[40620-51-7]	(<i>d</i>) 3-(<i>m</i> -fluorophenyl)-3-hydroxypropanoic acid				
	$\Delta_{\text{fus}}H$		24.27	311		[1991CHI/BRA]
C ₉ H ₉ FO ₃	[na]	(<i>dl</i>) 3-(<i>o</i> -fluorophenyl)-3-hydroxypropanoic acid				
	$\Delta_{\text{fus}}H$		27.2	342		[1991CHI/BRA]
C ₉ H ₉ FO ₃	[40620-50-6]	(<i>d</i>) 3-(<i>o</i> -fluorophenyl)-3-hydroxypropanoic acid				
	$\Delta_{\text{fus}}H$		22.59	348		[1991CHI/BRA]
C ₉ H ₉ FO ₃	[40620-62-0]	(<i>dl</i>) 3-(<i>p</i> -fluorophenyl)-3-hydroxypropanoic acid				
	$\Delta_{\text{fus}}H$		27.61	362		[1991CHI/BRA]
C ₉ H ₉ FO ₃	[na]	(<i>d</i>) 3-(<i>p</i> -fluorophenyl)-3-hydroxypropanoic acid				
	$\Delta_{\text{fus}}H$		30.96	381		[1991CHI/BRA]
C ₉ H ₉ F ₆ NO ₅	[1548-45-4]	(<i>l</i>) N,O-bis(trifluoroacetal)-threonine methyl ester (323–413)	72.5	338	A	[1987STE/MAL, 1999DYK/SVO]
C ₉ H ₉ N	[603-76-9]	1-methylindole				
	Δ_vH		62.2 ± 1.6	298	C	[2009RIB/CAB2]
C ₉ H ₉ N	[95-20-5]	2-methylindole				
	$\Delta_{\text{fus}}H$		15.72	329.4		[1997PEY/LET]
	$\Delta_{\text{sub}}H$		88.7 ± 2.4	298	C	[2009RIB/CAB2]
C ₉ H ₉ N	[83-34-1]	3-methylindole (skatole)				
	$\Delta_{\text{sub}}H$		90.4 ± 1.9	298	C	[2009RIB/CAB2]
	$\Delta_{\text{sub}}H$	(288–333)	83.3	303	A	[1987STE/MAL]

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(368–540)	64.5	383	A	[1987STE/MAL, 1947STU]
C ₉ H ₉ N	[1823-91-2]	α -methylbenzylcyanide				
	$\Delta_v H$	(284–318)	60.8 ± 0.7	301	GS	[2000VER]
	$\Delta_v H$	(284–318)	60.9 ± 0.7	298	GS	[2000VER]
C ₉ H ₉ N	[21789-36-6]	2,6-dimethylbenzotrile				
	$\Delta_{\text{sub}} H$		83.9 ± 2.8	298	C	[1991ACR/TUC]
C ₉ H ₉ NO ₂	[122-85-0]	4-acetomidobenzaldehyde				
	$\Delta_{\text{sub}} H$	(328–346)	99.0	337	A	[1987STE/MAL, 1960AIH2]
C ₉ H ₉ NO ₄	[618-98-4]	3-nitrobenzoic acid, ethyl ester				
	$\Delta_v H$	(381–571)	65.1	396	A	[1987STE/MAL, 1947STU]
C ₉ H ₉ NO ₄	[5251-93-4]	[(benzoylamino)oxy] acetic acid				
	$\Delta_{\text{fus}} H$		31.46	416.9	DSC	[1991ACR, 1990DON/DRE]
C ₉ H ₉ NO ₄	[5453-67-8]	dimethyl pyridine-2,6-dicarboxylate				
	$\Delta_{\text{sub}} H$		113.5 ± 3.8	298	C	[2005MAT/MOR]
C ₉ H ₉ NO ₅	[na]	(dl) 2-(p-nitrophenoxy)propanoic acid				
	$\Delta_{\text{fus}} H$		32.22	411.4		[1991CHI/BRA]
C ₉ H ₉ NO ₅	[na]	(d) 2-(p-nitrophenoxy)propanoic acid				
	$\Delta_{\text{fus}} H$		20.92	362		[1991CHI/BRA]
C ₉ H ₉ N ₃ O ₂ S ₂	[72-14-0]	4-amino-N-2-thiazolylbenzenesulfonamide (sulfathiazole)				
	$\Delta_{\text{fus}} H$		30.3	473	DSC	[2003MAR/AVI, 2002MAR/GOM]
C ₉ H ₉ N ₃ O ₆	[602-96-0]	2,4,6-trinitromesitylene				
	$\Delta_{\text{sub}} H$	(319–397)	103.6 ± 1.2		ME	[1987STE/MAL, 1978CUN/PAL]
C ₉ H ₁₀	[496-11-7]	indane				
	$\Delta_{\text{fus}} H$		8.6	221.8		[1996DOM/HEA]
	$\Delta_v H$	(374–466)	44.0	389	A	[1987STE/MAL]
	$\Delta_v H$		49.0	298	C	[1981HOS/SCO3]
	$\Delta_v H$	(355–482)	45.0	370		[1976AMB/SPR]
C ₉ H ₁₀	[611-15-4]	2-methylstyrene				
	$\Delta_v H$	(305–385)	47.9	320	A	[1987STE/MAL, 1953CLE/WIS]
C ₉ H ₁₀	[100-80-1]	3-methylstyrene				
	$\Delta_v H$	(314–385)	47.5	329	A	[1987STE/MAL, 1953CLE/WIS]
C ₉ H ₁₀	[622-97-9]	4-methylstyrene				
	$\Delta_v H$	(304–390)	47.6	319	A	[1987STE/MAL, 1953CLE/WIS]
C ₉ H ₁₀	[98-83-9]	α -methylstyrene				
	$\Delta_{\text{fus}} H$		11.92	250.8		[1996DOM/HEA]
	$\Delta_v H$	(274–314)	49.2 ± 0.3	294	GS	[1999VER6]
	$\Delta_v H$	(274–314)	48.9 ± 0.3	298	GS	[1999VER6]
	$\Delta_v H$	(331–467)	48.6 ± 0.4	298	EB	[1997STE/CHI2]
	$\Delta_v H$	(331–467)	45.9 ± 0.3	340	EB	[1997STE/CHI2]
	$\Delta_v H$	(331–467)	43.3 ± 0.3	380	EB	[1997STE/CHI2]
	$\Delta_v H$	(331–467)	40.6 ± 0.3	420	EB	[1997STE/CHI2]
	$\Delta_v H$	(331–467)	37.7 ± 0.5	460	EB	[1997STE/CHI2]
	$\Delta_v H$	(343–493)	44.3	358	A	[1987STE/MAL]
	$\Delta_v H$	(353–413)	44.8	368	A	[1987STE/MAL]
	C ₉ H ₁₀	[766-90-5]	<i>cis</i> β -methylstyrene			
$\Delta_v H$		(348–498)	44.8	363	A	[1987STE/MAL]

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₉ H ₁₀	[873-66-5] $\Delta_v H$	<i>trans</i> β -methylstyrene (291–452)	46.4	306	A	[1987STE/MAL, 1947STU]
C ₉ H ₁₀	[300-57-2] $\Delta_v H$ $\Delta_v H$	allylbenzene (274–313) (274–313)	46.5 ± 0.2	294	GS	[1999VER6]
			46.3 ± 0.2	298	GS	[1999VER6]
C ₉ H ₁₀ BrClN ₂ O ₂	[13360-45-7] $\Delta_{\text{fus}} H$	3-(4-bromo-3-chlorophenyl)-1-methoxy-1-methylurea	26.54	369.8	DSC	[1990DON/DRE]
C ₉ H ₁₀ ClNO ₃	[194085-75-1] $\Delta_{\text{fus}} H$	(1S)-1-(2-chlorophenyl)-1,2-ethanediol, 2-carbamate, (carisbamate)	33.0	409.6	DSC	[2009WU/MEH]
C ₉ H ₁₀ Cl ₂ N ₂ O	[330-54-1] $\Delta_{\text{fus}} H$ $\Delta_{\text{fus}} H$ $\Delta_{\text{fus}} H$ $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$	3-(3,4-dichlorophenyl)-1,1-dimethylurea (diuron)	25.28	435.1	DSC	[2000ROD/VEC]
			30.47	430.5	DSC	[1990DON/DRE]
			33.89	429.7	DSC	[1969PLA/GLA]
			119 ± 0.6	393	C	[1997PFE/SAB]
			133.9 ± 0.7	298	C	[1997PFE/SAB]
C ₉ H ₁₀ Cl ₂ N ₂ O ₂	[330-55-2] $\Delta_{\text{fus}} H$	N'-(3,4-dichlorophenyl)-N-methoxy-N-methylurea (linuron)	26.56	365.8	DSC	[1991ACR, 1990DON/DRE]
C ₉ H ₁₀ F ₂	[146377-62-0] $\Delta_v H$	1,1-difluoro-3-phenylpropane (278–318)	53.3 ± 0.4	298	GS	[1997SCH/VER]
C ₉ H ₁₀ N ₂	[1075-76-9] $\Delta_{\text{us}} H$ $\Delta_{\text{fus}} H$	N-(2-cyanoethyl)aniline (83–353) (83–353)	0.98	310.6		
			19.4	323.3	AC	[2005TIA/TAN]
C ₉ H ₁₀ N ₂	[53406-41-0] $\Delta_{\text{fus}} H$	N-amino-2-methylindole	25.55	384.6		[1997PEY/LET]
C ₉ H ₁₀ N ₂ O	[92-43-3] $\Delta_{\text{sub}} H$	1-phenyl-3-pyrazolidinone (327–348)	84.3	337.5	A	[1987STE/MAL, 1960AIH2]
C ₉ H ₁₀ N ₂ O ₃	[612-45-3] $\Delta_{\text{fus}} H$ (white cryst) $\Delta_{\text{fus}} H$ (amber cryst) $\Delta_{\text{fus}} H$ (yellow cryst)	N-(4-methyl-2-nitrophenyl)acetamide	24.25	366.2		
			20.97	356.7		
			22.37	364.2	DSC	[2001HE/STO]
C ₉ H ₁₀ N ₂ O ₃	[6335-41-7] $\Delta_{\text{fus}} H$	2-(hydroxyimino)-N-(4-methoxyphenyl)acetamide	8.3	457.6	DTA	[1982CUE/SOL]
C ₉ H ₁₀ N ₂ O ₃	[6335-42-8] $\Delta_{\text{fus}} H$	2-(hydroxyimino)-N-(2-methoxyphenyl)acetamide	27.8	424.6	DTA	[1982CUE/SOL]
C ₉ H ₁₀ O	[1746-13-0] $\Delta_v H$	allyl phenyl ether (349–456)	49.4	364	A	[1987STE/MAL]
C ₉ H ₁₀ O	[104-54-1] $\Delta_{\text{fus}} H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$	cinnamyl alcohol	15.73	308.2		[1991CHI/BRA]
			68.1 ± 0.1	310	TG,DTA	[2002SOR/DOL]
			79.8	319	A	[1987STE/MAL]
			56.2	388	A	[1987STE/MAL]
C ₉ H ₁₀ O	[15764-16-6] $\Delta_v H$	2,4-dimethylbenzaldehyde (358–489)	57.4	373	A	[1987STE/MAL]
C ₉ H ₁₀ O	[1470-94-6] $\Delta_v H$	5-hydroxyindane (393–524)	55.4	408	A	[1987STE/MAL]
C ₉ H ₁₀ O	[122-00-9]	4'-methylacetophenone				

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(288–333)	59.6	303	A	[1987STE/MAL]
C ₉ H ₁₀ O	[93-53-8]	2-phenylpropionaldehyde				
	$\Delta_v H$	(364–517)	52.3 ± 0.2	360	EB	[2002STE/CHI5]
	$\Delta_v H$	(364–517)	49.4 ± 0.2	400	EB	[2002STE/CHI5]
	$\Delta_v H$	(364–517)	46.6 ± 0.3	440	EB	[2002STE/CHI5]
	$\Delta_v H$	(364–517)	43.4 ± 0.5	480	EB	[2002STE/CHI5]
C ₉ H ₁₀ O	[104-53-0]	3-phenylpropionaldehyde				
	$\Delta_v H$	(330–363)	67.5	345	A	[1987STE/MAL]
C ₉ H ₁₀ O	[103-79-7]	benzyl methyl ketone				
	$\Delta_v H$	(343–383)	56.1	298	CGC	[1995CHI/HOS]
	$\Delta_v H$	(343–383)	55.0	298	CGC	[1995CHI/HOS]
C ₉ H ₁₀ O	[93-55-0]	ethyl phenyl ketone (propiofenone)				
	$\Delta_v H$	(388–623)	52.1	403	A	[1987STE/MAL]
	$\Delta_v H$	(391–454)	44.4	406	EB,GS	[1965COL/COU]
C ₉ H ₁₀ O	[612-15-7]	2-vinylanisole				
	$\Delta_v H$	(314–467)	56.7	329	A	[1987STE/MAL, 1947STU]
C ₉ H ₁₀ O	[626-20-0]	3-vinylanisole				
	$\Delta_v H$	(316–471)	55.9	331	A	[1987STE/MAL, 1947STU]
C ₉ H ₁₀ O	[637-69-4]	4-vinylanisole				
	$\Delta_v H$	(318–478)	54.9	333	A	[1987STE/MAL, 1947STU]
C ₉ H ₁₀ O	[4407-36-7]	<i>(E)</i> -3-phenyl-2-propen-1-ol				
	$\Delta_{\text{sub}} H$	(288–307)	109.6	297.5	A	[1987STE/MAL]
	$\Delta_{\text{sub}} H$		69.5		ME	[1954SER/VOI]
C ₉ H ₁₀ O	[493-08-3]	chroman				
	$\Delta_{\text{fus}} H$		16.26	269.8		[1996DOM/HEA]
	$\Delta_{\text{sub}} H$	(293–535)	56.7 ± 0.1	298	IPM,EB	[1990CHI/ARC]
	$\Delta_{\text{sub}} H$	(293–535)	55.2 ± 0.1	320	IPM,EB	[1990CHI/ARC]
	$\Delta_{\text{sub}} H$	(293–535)	52.7 ± 0.1	360	IPM,EB	[1990CHI/ARC]
	$\Delta_{\text{sub}} H$	(293–535)	50.2 ± 0.1	400	IPM,EB	[1990CHI/ARC]
	$\Delta_{\text{sub}} H$	(293–535)	48.9 ± 0.1	440	IPM,EB	[1990CHI/ARC]
	$\Delta_{\text{sub}} H$	(293–535)	45.1 ± 0.3	480	IPM,EB	[1990CHI/ARC]
C ₉ H ₁₀ O	[493-05-0]	isochroman				
	$\Delta_{\text{fus}} H$		16.75	277.5		[1996DOM/HEA]
	$\Delta_{\text{sub}} H$	(295–536)	57.1 ± 0.1	298	IPM,EB	[1990CHI/ARC]
	$\Delta_{\text{sub}} H$	(295–536)	55.6 ± 0.1	320	IPM,EB	[1990CHI/ARC]
	$\Delta_{\text{sub}} H$	(295–536)	52.9 ± 0.1	360	IPM,EB	[1990CHI/ARC]
	$\Delta_{\text{sub}} H$	(295–536)	50.3 ± 0.1	400	IPM,EB	[1990CHI/ARC]
	$\Delta_{\text{sub}} H$	(295–536)	48.7 ± 0.1	440	IPM,EB	[1990CHI/ARC]
	$\Delta_{\text{sub}} H$	(295–536)	45.3 ± 0.3	480	IPM,EB	[1990CHI/ARC]
C ₉ H ₁₀ O ₂	[89-71-4]	methyl <i>o</i> -toluate				
	$\Delta_{\text{fus}} H$	(5–320)	12.47	228.8	AC	[2002BLO/PAU]
	$\Delta_{\text{fus}} H$		12.5	228.8	AC	[1998MAK/KAB]
	$\Delta_v H$		57.3 ± 0.2	293	C	[1998MAK/KAB]
C ₉ H ₁₀ O ₂	[99-36-5]	methyl <i>m</i> -toluate				
	$\Delta_{\text{fus}} H$	(5–320)	17.14	270.6	AC	[2002BLO/PAU]
	$\Delta_{\text{fus}} H$		21.15	269.9	AC	[1998MAK/KAB]

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$		60.3 ± 0.2	296	C	[1998MAK/KAB]
	$\Delta_v H$		53.5	388		[1974MUR/TUD]
C ₉ H ₁₀ O ₂	[99-75-2]	methyl <i>p</i> -toluate				
	$\Delta_{\text{fus}} H$	(5–320)	20.78	306.5	AC	[2002BLO/PAU]
	$\Delta_{\text{fus}} H$		20.77	306.5	AC	[1998MAK/KAB]
	$\Delta_{\text{sub}} H$		83.3 ± 0.3	298	C	[1998MAK/KAB]
C ₉ H ₁₀ O ₂	$\Delta_{\text{sub}} H$	(403–493)	51.6	418		[1998SEM/WIL]
	[122-46-3]	acetic acid, 3-tolyl ester				
C ₉ H ₁₀ O ₂	$\Delta_v H$	(385–480)	55.7	400	A,EB	[1987STE/MAL, 1969SHE/LAN]
	[140-39-6]	acetic acid, 4-tolyl ester				
C ₉ H ₁₀ O ₂	$\Delta_v H$	(385–480)	55.9	400	A,EB	[1987STE/MAL, 1969SHE/LAN]
	[579-74-8]	2-acetylanisole				
C ₉ H ₁₀ O ₂	$\Delta_v H$		56.5			[1986BAL/GNA]
	[100-06-1]	4-acetylanisole				
C ₉ H ₁₀ O ₂	$\Delta_{\text{sub}} H$	(276–300)	77.7		V	[1959AIH]
	$\Delta_{\text{sub}} H$	(283–333)	93.7	308	A	[1954SER/VOI, 1960JON, 1987STE/MAL]
C ₉ H ₁₀ O ₂	$\Delta_v H$	(311–334)	66.5	322	A,ME	[1987STE/MAL, 1954SER/VOI]
	[7216-18-4]	3,4-dihydro-2 <i>H</i> -1,5-benzodioxepin				
C ₉ H ₁₀ O ₂	$\Delta_v H$		55.6			[1958CAS/FLE2]
	[140-11-4]	benzyl acetate				
C ₉ H ₁₀ O ₂	$\Delta_v H$	(283–490)	55.5	298	A	[1987STE/MAL]
	$\Delta_v H$	(283–328)	60.4	305	ME	[1954SER/VOI]
C ₉ H ₁₀ O ₂	[93-89-0]	ethylbenzoate				
	$\Delta_v H$	(283–332)	61.1 ± 0.3	298	GS	[2006VAS/VER]
	$\Delta_v H$	(369–531)	52.5 ± 0.2	380	EB	[2002STE/CHI2]
	$\Delta_v H$	(369–531)	49.6 ± 0.2	420	EB	[2002STE/CHI2]
	$\Delta_v H$	(369–531)	46.7 ± 0.3	460	EB	[2002STE/CHI2]
	$\Delta_v H$	(369–531)	43.6 ± 0.5	500	EB	[2002STE/CHI2]
	$\Delta_v H$	(344–440)	57.0	356	BG	[1988KAT2]
	$\Delta_v H$	(344–440)	50.5	419	BG	[1988KAT2]
	$\Delta_v H$	(288–333)	55.9	303	A	[1987STE/MAL]
	$\Delta_v H$	(358–487)	50.4	373	A	[1987STE/MAL]
	$\Delta_v H$	(317–486)	51.9	332		[1947STU]
	C ₉ H ₁₀ O ₂	[501-52-0]	3-phenylpropionic acid (hydrocinnamic acid)			
$\Delta_{\text{fus}} H$			15.61	321.2	DSC	[2001MON/HIL4]
$\Delta_{\text{fus}} H$			17.68	321.2		[1991ACR]
$\Delta_{\text{sub}} H$		(305–315)	102.0 ± 0.7	310	ME	[2001MON/HIL4]
$\Delta_{\text{sub}} H$		(305–315)	102.4 ± 0.8	298	ME	[2001MON/HIL4]
C ₉ H ₁₀ O ₂	$\Delta_v H$	(375–553)	67.0	390	A	[1987STE/MAL, 1947STU]
	[99-36-5]	3-methylbenzoic acid, methyl ester				
C ₉ H ₁₀ O ₂	$\Delta_v H$	(359–500)	54.8	374	A	[1987STE/MAL]
	[122-60-1]	(phenoxy)methyl)oxirane				
C ₉ H ₁₀ O ₂	$\Delta_v H$	(400–532)	69.9 ± 0.7	298	EB	[1997STE/CHI]
	$\Delta_v H$	(400–532)	60.3 ± 0.5	400	EB	[1997STE/CHI]
	$\Delta_v H$	(400–532)	56.7 ± 0.4	440	EB	[1997STE/CHI]
	$\Delta_v H$	(400–532)	53.1 ± 0.4	480	EB	[1997STE/CHI]
	$\Delta_v H$	(400–532)	51.3 ± 0.5	500	EB	[1997STE/CHI]
	$\Delta_v H$	(400–532)	49.4 ± 0.6	520	EB	[1997STE/CHI]

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(343–373)	65.6 ± 0.1			[1976KUZ/MIR]
C ₉ H ₁₀ O ₂	[101-41-7]	methyl phenylacetate				
	$\Delta_v H$	(333–433)	57.4	298	GC	[2005HOS/GRY]
C ₉ H ₁₀ O ₂	[122-46-3]	3-methylphenyl acetate				
	$\Delta_{\text{sub}} H$	(274–317)	60.7	295	TE	[1947BAL, 1960JON]
C ₉ H ₁₀ O ₂	[936-51-6]	2-phenyl-1,3-dioxolane				
	$\Delta_v H$	(285–333)	62.6 ± 0.7	298	GS	[2002VER]
	$\Delta_v H$	(298–333)	62.1 ± 0.3	316	GS	[1995VER/DOG]
C ₉ H ₁₀ O ₂	[612-19-1]	2-ethylbenzoic acid				
	$\Delta_{\text{sub}} H$	(298–313)	100.5	305.5	ME	[1987STE/MAL, 1976COL/JIM]
	$\Delta_{\text{sub}} H$		101.1 ± 0.4	298	ME	[1984COL/JIM]
	$\Delta_{\text{sub}} H$	(298–313)	100.7 ± 2.5	298	ME	[1976COL/JIM]
C ₉ H ₁₀ O ₂	[619-20-5]	3-ethylbenzoic acid				
	$\Delta_{\text{sub}} H$	(300–318)	99.1	309	ME	[1987STE/MAL, 1976COL/JIM]
	$\Delta_{\text{sub}} H$		99.7 ± 0.4	298	ME	[1984COL/JIM]
	$\Delta_{\text{sub}} H$	(300–318)	99.11 ± 2.5	298	ME	[1976COL/JIM]
C ₉ H ₁₀ O ₂	[619-64-7]	4-ethylbenzoic acid				
	$\Delta_{\text{fus}} H$		14.06	386.2		[1991CHI/BRA]
	$\Delta_{\text{sub}} H$	(321–335)	101.2 ± 0.8	298	ME	[2004MON/ALM]
	$\Delta_{\text{sub}} H$	(310–329)	98.2	319.5	ME	[1987STE/MAL, 1976COL/JIM]
	$\Delta_{\text{sub}} H$		98.9 ± 0.2	298	ME	[1984COL/JIM]
	$\Delta_{\text{sub}} H$	(311–330)	97.6 ± 0.2	321	ME	[1984COL/JIM]
	$\Delta_{\text{sub}} H$	(320–329)	97.5 ± 2.5	298	ME	[1976COL/JIM]
C ₉ H ₁₀ O ₂	[603-79-2]	2,3-dimethylbenzoic acid				
	$\Delta_{\text{fus}} H$		18.3	417.6		[1996BEL/UFN]
	$\Delta_{\text{sub}} H$	(316–337)	102.3 ± 0.4	326	ME	[1984COL/JIM2]
	$\Delta_{\text{sub}} H$	(316–337)	104.6 ± 0.4	298	ME	[1984COL/JIM2]
C ₉ H ₁₀ O ₂	[611-01-8]	2,4-dimethylbenzoic acid				
	$\Delta_{\text{sub}} H$	(312–331)	102.7 ± 0.3	321	ME	[1984COL/JIM2]
	$\Delta_{\text{sub}} H$	(312–331)	103.5 ± 0.3	298	ME	[1984COL/JIM2]
C ₉ H ₁₀ O ₂	[610-72-0]	2,5-dimethylbenzoic acid				
	$\Delta_{\text{sub}} H$	(315–334)	103.6 ± 0.6	324	ME	[1984COL/JIM2]
	$\Delta_{\text{sub}} H$	(315–334)	105.0 ± 0.6	298	ME	[1984COL/JIM2]
C ₉ H ₁₀ O ₂	[632-46-2]	2,6-dimethylbenzoic acid				
	$\Delta_{\text{sub}} H$	(309–324)	98.2 ± 0.2	317	ME	[1984COL/JIM2]
	$\Delta_{\text{sub}} H$	(309–324)	99.1 ± 0.2	298	ME	[1984COL/JIM2]
C ₉ H ₁₀ O ₂	[632-46-2]	3,4-dimethylbenzoic acid				
	$\Delta_{\text{sub}} H$	(325–347)	104.5 ± 0.3	336	ME	[1984COL/JIM2]
	$\Delta_{\text{sub}} H$	(325–347)	106.4 ± 0.3	298	ME	[1984COL/JIM2]
C ₉ H ₁₀ O ₂	[499-06-9]	3,5-dimethylbenzoic acid				
	$\Delta_{\text{fus}} H$		22.6	442.9		[1996BEL/UFN]
	$\Delta_{\text{sub}} H$	(322–341)	100.8 ± 0.3	332	ME	[1984COL/JIM2]
	$\Delta_{\text{sub}} H$	(322–341)	102.3 ± 0.3	298	ME	[1984COL/JIM2]
C ₉ H ₁₀ O ₂	[122-60-1]	phenyl glycidyl ether				
	$\Delta_{\text{fus}} H$		17.32	279.8		[1988LEB/BYK]
C ₉ H ₁₀ O ₂	[935-92-2]	2,3,5-trimethyl-1,4-benzoquinone				
	$\Delta_v H$	(393–450)	49.9	408	EB	[2004TAN/LI]

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₉ H ₁₀ O ₂ S	[5535-52-4]	p-tolyl vinyl sulfone				
	$\Delta_{\text{fus}}H$		10.88	340.4		[1969MAC/MCN]
	$\Delta_{\text{sub}}H$		82.4 ± 2.5		B	[1969MAC/MCN, 1969MAC/MCN2, 1977PED/RYL]
C ₉ H ₁₀ O ₃	[118-61-6]	ethyl salicylate				
	Δ_vH	(431–461)	53.4			[1982WAY]
	Δ_vH	(288–333)	59.2	303	A	[1987STE/MAL]
	Δ_vH	(334–505)	55.2	349	A	[1987STE/MAL]
C ₉ H ₁₀ O ₃	[623-20-1]	2-furanacrylic acid, ethyl ester				
	Δ_vH	(428–500)	56.8	443	A	[1987STE/MAL]
C ₉ H ₁₀ O ₃	[121-98-2]	methyl 4-methoxybenzoate				
	Δ_vH	(382–472)	61.1	397	EB	[1985SCH/BRU]
C ₉ H ₁₀ O ₃	[120-47-8]	ethyl 4-hydroxybenzoate				
	$\Delta_{\text{fus}}H$		27.9	389.2	DSC	[2008WAS/HOL]
	$\Delta_{\text{fus}}H$		32.49	388.9	DSC	[2008NIC/BEL]
	$\Delta_{\text{fus}}H$		26.4	389		[1999GIO/BET]
	$\Delta_{\text{fus}}H$	(313–326)	100.9 ± 0.7	298	GS	[2005PER/ROD]
	Δ_vH		75.0		TGA	[2002CHA/DOL]
	Δ_vH		72.6		TGA	[2001CHA/DOL]
C ₉ H ₁₀ O ₃	[35438-32-5]	<i>cis,cis</i> 3-methyl-4-cyclohexene-1,2-dicarboxylic acid anhydride				
	Δ_vH	(325–525)	49.5 ± 1.0			[1984NUR/MEK]
C ₉ H ₁₀ O ₃	[3425-89-6]	4-methyl-4-cyclohexene-1,2-dicarboxylic anhydride				
	$\Delta_{\text{fus}}H$	(80–361)	17.67	335.5	AC	[2005LU/TAN]
C ₉ H ₁₀ O ₃	[121-32-4]	3-ethoxy-4-hydroxybenzaldehyde (ethyl vanillin)				
	$\Delta_{\text{fus}}H$		23.1	349.8	DSC	[2008TEM/ROU]
	$\Delta_{\text{sub}}H$	(296–338)	101.5	311		[1987STE/MAL, 1957LIT, 1960JON]
C ₉ H ₁₀ O ₃	[120-14-9]	3,4-dimethoxybenzaldehyde				
	$\Delta_{\text{fus}}H$		20.3	317	DSC	[2008TEM/ROU]
C ₉ H ₁₀ O ₃	[na]	(<i>dl</i>) 3-phenyl-3-hydroxypropanoic acid				
	$\Delta_{\text{fus}}H$		29.71	366		[1991CHI/BRA]
C ₉ H ₁₀ O ₃	[2768-42-5]	(<i>d</i>) 3-phenyl-3-hydroxypropanoic acid				
	$\Delta_{\text{fus}}H$		32.64	391		[1991CHI/BRA]
C ₉ H ₁₀ O ₃	[940-31-8]	(<i>dl</i>) 2-phenoxypropionic acid				
	$\Delta_{\text{fus}}H$		33.05	388		[1991CHI/BRA]
C ₉ H ₁₀ O ₃	[na]	(<i>d</i>) 2-phenoxypropionic acid				
	$\Delta_{\text{fus}}H$		22.59	359		[1991CHI/BRA]
C ₉ H ₁₀ O ₃	[104-01-8]	4-methoxyphenylacetic acid				
	$\Delta_{\text{fus}}H$		21.8	358.1		[1991ACR]
C ₉ H ₁₀ O ₃	[na]	4-hydroxyphenylpropionic acid				
	$\Delta_{\text{fus}}H$		28.9	402.5		[1991ACR]
C ₉ H ₁₀ O ₃	[619-86-3]	4-ethoxybenzoic acid				
	$\Delta_{\text{fus}}H$		29.4	472.8		[1991ACR]
	$\Delta_{\text{sub}}H$		123.3 ± 0.9	298		[2010RIB/FER3]
C ₉ H ₁₀ O ₃	[3663-82-9]	1,4-benzodioxan-2-hydroxymethyl				
	$\Delta_{\text{fus}}H$		28.78	362.4	DSC	[2008MAT/SOU]

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{sub}}H$		106.9 ± 0.8	298	C	[2008MAT/MIR2]
C ₉ H ₁₀ O ₃	[53282-12-5]	ethyl <i>trans</i> β-(2-furyl)acrylate (428–500)	56.8	464		[1956FRO/LOE]
C ₉ H ₁₀ O ₄	[1521-38-6]	2,3-dimethoxybenzoic acid (336–356)	115.1 ± 0.3	346	ME	[1985COL/JIM]
	$\Delta_{\text{sub}}H$	(336–356)	116.6 ± 0.3	298	ME	[1985COL/JIM]
C ₉ H ₁₀ O ₄	[91-52-1]	2,4-dimethoxybenzoic acid (346–367)	120.5 ± 0.4	357	ME	[1985COL/JIM]
	$\Delta_{\text{sub}}H$	(346–367)	123.4 ± 0.4	298	ME	[1985COL/JIM]
C ₉ H ₁₀ O ₄	[1466-76-8]	2,6-dimethoxybenzoic acid (335–378)	118.4 ± 0.4	367	ME	[1985COL/JIM]
	$\Delta_{\text{sub}}H$	(335–378)	121.7 ± 0.4	298	ME	[1985COL/JIM]
C ₉ H ₁₀ O ₄	[93-07-2]	3,4-dimethoxybenzoic acid (359–378)	126.1 ± 0.6	369	ME	[1985COL/JIM]
	$\Delta_{\text{sub}}H$	(359–378)	129.8 ± 0.6	298	ME	[1985COL/JIM]
C ₉ H ₁₀ O ₄	[2785-98-0]	2,5-dimethoxybenzoic acid (324–342)	113.3 ± 0.7	333	ME	[1996JIM/ROU]
	$\Delta_{\text{sub}}H$	(324–342)	116.1 ± 0.7	298	ME	[1996JIM/ROU]
C ₉ H ₁₀ O ₄	[1132-21-4]	3,5-dimethoxybenzoic acid (356–376)	124.5 ± 0.6	369	ME	[1985COL/JIM]
	$\Delta_{\text{sub}}H$	(356–376)	127.1 ± 0.6	298	ME	[1985COL/JIM]
C ₉ H ₁₀ O ₄	[na]	(±) bicyclo[2.2.1]hept-5-ene- <i>trans</i> -2,3-dicarboxylic acid	29.8	458.2		[1971PIN/TON]
	$\Delta_{\text{fus}}H$					
C ₉ H ₁₀ O ₄	[32216-02-7]	(+) bicyclo[2.2.1]hept-5-ene- <i>trans</i> -2,3-dicarboxylic acid	22.5	449.2		[1971PIN/TON]
	$\Delta_{\text{fus}}H$					
C ₉ H ₁₀ O ₄	[na]	(<i>dl</i>) erythro phenylglyceric acid	31.38	395		[1991CHI/BRA]
	$\Delta_{\text{fus}}H$					
C ₉ H ₁₀ O ₄	[na]	(<i>d</i>) erythro phenylglyceric acid	23.43	371.5		[1991CHI/BRA]
	$\Delta_{\text{fus}}H$					
C ₉ H ₁₀ O ₅	[613-75-2]	2-(diacetoxymethyl)furan	109.6 ± 2.5			[1980BAL/LEB, 1986PED/NAY]
	$\Delta_{\text{sub}}H$					
C ₉ H ₁₀ O ₅	[530-57-4]	3,5-dimethoxy-4-hydroxybenzoic acid (syringic acid)	33.7	480.3	DSC	[2009QUE/MOT]
	$\Delta_{\text{fus}}H$					
C ₉ H ₁₁ Br	[7073-94-1]	1-bromo-2-isopropylbenzene (404–484)	48.4	419		[1999DYK/SVO]
	Δ_vH	(378–528)	49.8	393	A	[1987STE/MAL, 1970DYK/VAN]
C ₉ H ₁₁ Br	[586-61-8]	1-bromo-4-isopropylbenzene (362–493)	51.1	377		[1999DYK/SVO]
	Δ_vH	(388–528)	50.4	403	A	[1987STE/MAL, 1970DYK/VAN]
C ₉ H ₁₁ Br	[3575-19-7]	cumyl bromide	58.0	298	CGC	[2002KRA/VAS]
	Δ_vH					
C ₉ H ₁₁ BrN ₂ O	[3060-89-7]	N'-(4-bromophenyl)-N-methoxy-N-methylurea	24.44	368.3	DSC	[1991ACR, 1990DON/DRE]
	$\Delta_{\text{fus}}H$					
C ₉ H ₁₁ BrO ₂	[109417-60-9]	1-bromo-2-(2-methoxyethoxy)benzene (302–368)	64.5 ± 0.3	298	GS	[2006DAB/SPO]
	Δ_vH					
C ₉ H ₁₁ BrO ₃	[63834-58-2]	(racemic) 3-(2-bromophenoxy)propane-1,2-diol				

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		T _m (K)	Method	Reference
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)			
		$\Delta_{\text{fus}}H$	33.4	353.5		[2006ZAK/LAZ]
C ₉ H ₁₁ BrO ₃	[386702-67-6]	(R) 3-(2-bromophenoxy)propane-1,2-diol				
		$\Delta_{\text{fus}}H$	38.4	374.3		[2006ZAK/LAZ]
C ₉ H ₁₁ Cl	[2077-13-6]	1-chloro-2-isopropylbenzene				
		Δ_vH	(341–465) 48.1	356		[1999DYK/SVO]
		Δ_vH	(363–508) 47.7	378	A	[1987STE/MAL, 1970DYK/VAN]
C ₉ H ₁₁ Cl	[2621-46-7]	1-chloro-4-isopropylbenzene				
		Δ_vH	(307–472) 51.4	322		[1999DYK/SVO]
		Δ_vH	(368–513) 48.5	383	A	[1987STE/MAL, 1970DYK/VAN]
C ₉ H ₁₁ Cl	[934-53-2]	cumyl chloride				
		Δ_vH	54.7	298	CGC	[2002KRA/VAS]
C ₉ H ₁₁ ClN ₂ O	[150-68-5]	3-(4-chlorophenyl)-1,1-dimethylurea (monuron)				
		$\Delta_{\text{fus}}H$	29.3	447.6		[2004KON/TAN2]
		$\Delta_{\text{fus}}H$	29.46	447.6		[1991ACR]
		$\Delta_{\text{sub}}H$	(303–379) 114.6 ± 4.9	341	ME,C	[1987STE/MAL, 1972WIE]
C ₉ H ₁₁ ClN ₂ O ₂	[1746-81-2]	N'-(4-chlorophenyl)-N-methoxy-N-methylurea				
		$\Delta_{\text{fus}}H$	22.54	353.4	DSC	[1991ACR, 1990DON/DRE]
C ₉ H ₁₁ ClO ₂	[67146-43-4]	propylene glycol mono(4-chlorophenyl) ether				
		Δ_vH	(417–542) 64.9	432	A	[1987STE/MAL, 1999DYK/SVO]
C ₉ H ₁₁ ClO ₃	[93-65-2]	2-(4-chloro-2-methylphenoxy)propanoic acid				
		$\Delta_{\text{fus}}H$	26.43	366.2	DSC	[1990DON/DRE]
C ₉ H ₁₁ ClO ₃	[5112-21-0]	(racemic) 3-(2-chlorophenoxy)propane-1,2-diol				
		$\Delta_{\text{fus}}H$	29.0	344.9		[2006ZAK/LAZ]
C ₉ H ₁₁ ClO ₃	[153547-60-5]	(R) 3-(2-chlorophenoxy)propane-1,2-diol				
		$\Delta_{\text{fus}}H$	38.2	363.3		[2006ZAK/LAZ]
C ₉ H ₁₁ ClS	[4322-51-8]	benzyl (2-chloroethyl) sulfide				
		Δ_vH	(293–333) 52.3	308	A,GS	[1987STE/MAL, 1948RED/CHA, 1999DYK/SVO]
C ₉ H ₁₁ Cl ₃ NO ₃ PS	[330-55-2]	O,O-diethyl-O-(3,5,6-trichloro-2-pyridyl)phosphorothioate				
		$\Delta_{\text{fus}}H$	24.53	315	DSC	[1991ACR, 1990DON/DRE]
C ₉ H ₁₁ Cl ₃ NO ₄ P	[5598-15-2]	3,5,6-trichloro-2-pyridyl diethylphosphate (chlorpyrifos oxon)				
		$\Delta_{\text{sub}}H$	(373–403) 79		GC	[2007GOE/MCC]
C ₉ H ₁₁ FO ₃	[399-28-0]	(racemic) 3-(2-fluorophenoxy)propane-1,2-diol				
		$\Delta_{\text{fus}}H$	20.5	318.2		[2006ZAK/LAZ]
C ₉ H ₁₁ FO ₃	[912556-93-5]	(R) 3-(2-fluorophenoxy)propane-1,2-diol				
		$\Delta_{\text{fus}}H$	26.0	333.7		[2006ZAK/LAZ]
C ₉ H ₁₁ F ₅ O ₂	[24262-73-5]	pentafluoropropionic acid, cyclohexyl ester				
		Δ_vH	(335–428) 46.4	350	A,EB	[1987STE/MAL, 1969SHE/LAN]
C ₉ H ₁₁ I	[54290-22-1]	cumyl iodide				
		Δ_vH	63.3	298	CGC	[2002KRA/VAS]
C ₉ H ₁₁ IO ₃	[55169-06-7]	(racemic) 3-(2-iodophenoxy)propane-1,2-diol				
		$\Delta_{\text{fus}}H$	34.2	362.5		[2006ZAK/LAZ]
C ₉ H ₁₁ IO ₃	[912556-94-6]	(R) 3-(2-iodophenoxy)propane-1,2-diol				
		$\Delta_{\text{fus}}H$	37.4	383.5		[2006ZAK/LAZ]

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₉ H ₁₁ N	[635-46-1]	1,2,3,4-tetrahydroquinoline				
	$\Delta_{\text{fus}}H$		11.81	290		[1991ACR]
	Δ_vH	(323–572)	65.3 ± 0.2	298	IP,EB	[1989STE/CHI3]
	Δ_vH	(323–572)	62.1 ± 0.1	340	IP,EB	[1989STE/CHI3]
	Δ_vH	(323–572)	59.2 ± 0.1	380	IP,EB	[1989STE/CHI3]
	Δ_vH	(323–572)	56.3 ± 0.1	420	IP,EB	[1989STE/CHI3]
	Δ_vH	(323–572)	53.5 ± 0.2	460	IP,EB	[1989STE/CHI3]
	Δ_vH	(323–572)	50.8 ± 0.3	500	IP,EB	[1989STE/CHI3]
Δ_vH	(323–572)	47.9 ± 0.4	540	IP,EB	[1989STE/CHI3]	
C ₉ H ₁₁ N	[10500-57-9]	5,6,7,8-tetrahydroquinoline				
	$\Delta_{\text{fus}}H$		9.08	222.7		[1991ACR]
	Δ_vH	(303–544)	57.6 ± 0.2	298	IP,EB	[1989STE/CHI3]
	Δ_vH	(303–544)	56.1 ± 0.1	320	IP,EB	[1989STE/CHI3]
	Δ_vH	(303–544)	53.6 ± 0.1	360	IP,EB	[1989STE/CHI3]
	Δ_vH	(303–544)	51.1 ± 0.1	400	IP,EB	[1989STE/CHI3]
	Δ_vH	(303–544)	48.7 ± 0.3	440	IP,EB	[1989STE/CHI3]
	Δ_vH	(303–544)	46.2 ± 0.4	480	IP,EB	[1989STE/CHI3]
Δ_vH	(303–544)	43.5 ± 0.5	520	IP,EB	[1989STE/CHI3]	
C ₉ H ₁₁ NO	[579-10-2]	N-methylacetanilide				
	Δ_vH	(383–519)	60.1	398	A	[1987STE/MAL]
Δ_vH	(377–526)	56.7	392		[1947STU]	
C ₉ H ₁₁ NO	[120-66-1]	N-(2-methylphenyl)acetamide				
	$\Delta_{\text{sub}}H$	(315–340)	96.8	327.5		[1987STE/MAL, 1960AIH2]
C ₉ H ₁₁ NO	[103-89-9]	N-(4-methylphenyl)acetamide				
	$\Delta_{\text{sub}}H$	(331–350)	99.0	341		[1960AIH2]
C ₉ H ₁₁ NO	[611-74-5]	N,N-dimethylbenzamide				
	$\Delta_{\text{sub}}H$	(289–305)	89.7 ± 0.3	298		[1995ABB/JIM]
	$\Delta_{\text{sub}}H$		94.8 ± 2.0	298	C	[1989RIB/SOU]
C ₉ H ₁₁ NO	[120-66-1]	2-(acetylamino)toluene				
	$\Delta_{\text{fus}}H$		21.7	382.7	DSC	[2003HUA, 2005HUA/TAN]
C ₉ H ₁₁ NO	[103-89-9]	4-(acetylamino)toluene				
	$\Delta_{\text{fus}}H$		28.93	424	DSC	[2003HUA, 2005HUA/TAN]
C ₉ H ₁₁ NO	[100-10-7]	4-(N,N-dimethylamino)benzaldehyde				
	$\Delta_{\text{fus}}H$		19.07	346.2		[1999MEN/LIA]
C ₉ H ₁₁ NO ₂	[6526-72-3]	1-nitro-2-isopropylbenzene				
	Δ_vH	(278–323)	65.5 ± 0.7	301	GS	[2000VER/HEI]
	Δ_vH		65.6 ± 0.7	298		[2000VER/HEI]
C ₉ H ₁₁ NO ₂	[87-25-2]	ethyl 2-aminobenzoate (ethyl anthranilate)				
	Δ_vH	(433–593)	59.6	448	A	[1987STE/MAL]
C ₉ H ₁₁ NO ₂	[94-09-7]	ethyl 4-aminobenzoate (benzocaine)				
	$\Delta_{\text{fus}}H$		24.6	362.6	DSC	[2008WAS/HOL]
	$\Delta_{\text{fus}}H$		21.16	NA	DSC	[1995YAM/KIT]
	$\Delta_{\text{fus}}H$		23.56	362.8		[1991ACR]
C ₉ H ₁₁ NO ₂	[101-99-5]	ethyl carbanilate				
	Δ_vH	(380–510)	84.2	395	A	[1987STE/MAL, 1947STU]
C ₉ H ₁₁ NO ₂	[603-71-4]	2,4,6-trimethylnitrobenzene				
	$\Delta_{\text{sub}}H$		78.6 ± 1.0	298	C	[1993ACR/TUC2, 1993ACR/TUC2]
C ₉ H ₁₁ NO ₂	[63-91-2]	L-(l)-phenylalanine				

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{sub}}H$	(342–442)	U 90 ± 6.3	392	LE	[1977GAF/PIE]
	$\Delta_{\text{sub}}H$		154 ± 8	455	ME	[1965SVE/CLY, 1970COX/PIL, 1987STE/MAL, 1964CLY/SVE]
C ₉ H ₁₁ NO ₂	[101-88-5]	ethyl phenyl carbamate				
	$\Delta_{\text{fus}}H$		16.27	326		[1971PRI]
C ₉ H ₁₁ NO ₂	[51-66-1]	<i>p</i> -methoxyacetanilide				
	$\Delta_{\text{fus}}H$		27.82	400.3		[1990MAN/AHU]
C ₉ H ₁₁ NO ₂ S ₂	[949171-64-6]	N-theonylthiocarbamic-O-propyl ester				
	$\Delta_{\text{fus}}H$		26.1	370.1	DSC	[2007RIB/MON]
	$\Delta_{\text{sub}}H$		136.5 ± 1.8	298	C	[2007RIB/MON]
C ₉ H ₁₁ NO ₃	[60-18-4]	<i>(l)</i> -tyrosine				
	$\Delta_{\text{sub}}H$	(412–512)	101 ± 8	462	LE	[1977GAF/PIE]
C ₉ H ₁₂	[38451-18-2]	<i>cis</i> bicyclo[4.3.0]nona-3,7-diene				
	Δ_vH	(356–429)	41.8	371	A	[1987STE/MAL]
C ₉ H ₁₂	[28304-66-7]	<i>(Z)</i> 5-ethylidene-2-norbornene				
	Δ_vH	(315–462)	44.3 ± 0.3	298	EB	[1997STE/CHI]
	Δ_vH	(315–462)	43.0 ± 0.3	320	EB	[1997STE/CHI]
	Δ_vH	(315–462)	40.5 ± 0.3	360	EB	[1997STE/CHI]
	Δ_vH	(315–462)	38.0 ± 0.3	400	EB	[1997STE/CHI]
	Δ_vH	(315–462)	35.2 ± 0.5	440	EB	[1997STE/CHI]
C ₉ H ₁₂	[3048-64-4]	5-ethylidene-2-norbornene				
	Δ_vH	(314–420)	42.3 ± 0.3	298	EB	[1996STE/CHI2]
C ₉ H ₁₂	[28304-67-8]	<i>trans</i> 5-ethylidene-2-norbornene				
	Δ_vH	(346–416)	41.2	361	A	[1987STE/MAL]
C ₉ H ₁₂	[611-14-3]	2-ethyltoluene				
	Δ_vH		46.9	298		[1994RUZ/ZAB]
	Δ_vH		47.7	298		[1971WIL/ZWO]
	Δ_vH	(353–443)	43.6	368	A	[1987STE/MAL, 1949FOR/NOR]
C ₉ H ₁₂	[620-14-4]	3-ethyltoluene				
	Δ_vH		46.6	298		[1994RUZ/ZAB]
	Δ_vH		46.9	298		[1971WIL/ZWO]
	Δ_vH	(348–438)	43.4	363	A	[1987STE/MAL, 1949FOR/NOR]
C ₉ H ₁₂	[622-96-8]	4-ethyltoluene				
	Δ_vH		46.5	298		[1994RUZ/ZAB]
	Δ_vH		46.6	298		[1971WIL/ZWO]
	Δ_vH	(349–442)	43.2	364	A	[1987STE/MAL, 1949FOR/NOR]
C ₉ H ₁₂	[98-82-8]	isopropylbenzene				
	$\Delta_{\text{fus}}H$		7.32	177.1		[1973KIS/SUG]
	Δ_vH		45.1	298		[1994RUZ/ZAB]
	Δ_vH	(349–426)	41.2	364		[1989CEP/GON]
	Δ_vH	(339–433)	42.1	354	A	[1987STE/MAL]
	Δ_vH		45.1 ± 0.1	298	C	[1982FUC/HAL]
	Δ_vH		44.0	298		[1975KUS/SAI]
	Δ_vH		45.1	298		[1971WIL/ZWO]
	Δ_vH		45.1	298	C	[1947OSB/GIN]
	Δ_vH	(343–426)	41.9	358	MM	[1949FOR/NOR, 1945WIL/TAY]
C ₉ H ₁₂	[103-65-1]	propylbenzene				
	$\Delta_{\text{fus}}H$		9.27	173.6		[1991ACR]

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$		46.2	298		[1994RUZ/ZAB]
	$\Delta_v H$	(340–391)	43.8	355		[1986PAU/KRU]
	$\Delta_v H$		45.0	298		[1975KUS/SAI]
	$\Delta_v H$		46.2	298		[1971WIL/ZWO]
	$\Delta_v H$		46.2	298	C	[1947OSB/GIN]
	$\Delta_v H$	(348–433)	42.7	363	A,MM	[1987STE/MAL, 1949FOR/NOR, 1945WIL/TAY]
C₉H₁₂	[3048-65-5]	3a,4,7,7a-tetrahydro-1 <i>H</i> -indene				
	$\Delta_v H$	(338–440)	42.3	353	A	[1987STE/MAL]
C₉H₁₂	[526-73-8]	1,2,3-trimethylbenzene				
	$\Delta_{\text{us}} H$		0.66	218.7		[1996DOM/HEA]
	$\Delta_{\text{us}} H$		1.33	230.3		
	$\Delta_{\text{fus}} H$		8.18	247.8		[1996DOM/HEA]
	$\Delta_v H$		49.0	298		[1994RUZ/ZAB]
	$\Delta_v H$		48.8	298		[1974KUS/SAI]
	$\Delta_v H$	(363–456)	44.8	378	A	[1987STE/MAL, 1949FOR/NOR]
	$\Delta_v H$		49.1	298		[1971WIL/ZWO]
	$\Delta_v H$	(259–270)	42.5	265	RG	[1948HOP/SEA]
	$\Delta_v H$		49.1	298	C	[1947OSB/GIN]
C₉H₁₂	[95-63-6]	1,2,4-trimethylbenzene				
	$\Delta_{\text{fus}} H$		12.65	228.6		
	$\Delta_{\text{fus}} H$		13.19	229.3		[1996DOM/HEA]
	$\Delta_v H$		48.0	298		[1994RUZ/ZAB]
	$\Delta_v H$		47.2	298		[1974KUS/SAI]
	$\Delta_v H$		47.9	298		[1971WIL/ZWO]
	$\Delta_v H$	(357–450)	44.1	372	A	[1987STE/MAL, 1949FOR/NOR]
	$\Delta_v H$	(257–267)	46.5	262	RG	[1948HOP/SEA]
	$\Delta_v H$		47.9	298	C	[1947OSB/GIN]
C₉H₁₂	[108-67-8]	1,3,5-trimethylbenzene				
	$\Delta_{\text{us}} H$		0.33	91.3		
	$\Delta_{\text{us}} H$		0.07	188.5		[2000YAM/TAN]
	$\Delta_{\text{fus}} H$		9.51	228.4		[1991RAD/RAD]
	$\Delta_{\text{fus}} H$		9.51	228.4		[1996DOM/HEA, 1991RAD/RAD]
	$\Delta_v H$		47.6	298		[1994RUZ/ZAB]
	$\Delta_v H$	(296–342)	46.2 ± 1.3	319	MM	[1991WIB/WAL]
	$\Delta_v H$	(296–342)	47.5 ± 2.1	298	MM	[1991WIB/WAL]
	$\Delta_v H$	(348–424)	43.5	363		[1989PAR/GME]
	$\Delta_v H$	(249–356)	49.7	264	A	[1987STE/MAL]
	$\Delta_v H$		47.5 ± 0.1	298	C	[1987AN/HU]
	$\Delta_v H$	(273–299)	47.7	286	MM	[1981CHI/HYM]
	$\Delta_v H$	(354–445)	43.9	369	A	[1987STE/MAL, 1949FOR/NOR]
	$\Delta_v H$		47.5	298		[1971WIL/ZWO]
	$\Delta_v H$	(255–268)	51.1	262	RG	[1948HOP/SEA]
	$\Delta_v H$		47.5	298	C	[1947OSB/GIN]
C₉H₁₂	[3048-64-4]	5-vinyl-2-norbornene				
	$\Delta_v H$	(301–410)	42.0	316	A	[1987STE/MAL]
	$\Delta_v H$	(354–409)	48.9	369	A	[1987STE/MAL]
C₉H₁₂	[31561-59-8]	trispiro[2.0.2.0.2.0]nonane ([3] rotane)				
	$\Delta_{\text{fus}} H$		14.14	312.1	DSC	[1995BEC/RUC]
	$\Delta_v H$	(273–308)	59.2 ± 0.5		GS	[1995BEC/RUC]
C₉H₁₂	[50874-24-3]	trispiro[2.0.0.2.1.1]nonane ([3] triangulane)				
	$\Delta_v H$	(275–314)	46.3 ± 0.5		GS	[1995BEC/RUC]

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₉ H ₁₂ ClN ₅	[22936-86-3] $\Delta_{\text{fus}}H$	6-chloro-N-cyclopropyl-N'-(1-methylethyl)-1,3,5-triazine-2,4-diamine	28.76	441.6	DSC	[1990DON/DRE]
C ₉ H ₁₂ F ₃ N ₃ O ₅	[651-18-3] $\Delta_{\text{sub}}H$	N-[N-(N-[trifluoroacetyl]glycyl)glycyl]glycine methyl ester	(343–433) 133.4	358		[1987STE/MAL, 1960WEY/KLI]
C ₉ H ₁₂ NO ₅ PS	[122-14-5] Δ_vH	O,O-dimethyl-O-(3-methyl-4-nitrophenyl)thiophosphate	(293–382) 78.0	308	A	[1987STE/MAL]
C ₉ H ₁₂ N ₂	[103-02-6] Δ_vH	phenylhydrazone acetone	(413–436) 74.6	424	A	[1987STE/MAL]
C ₉ H ₁₂ N ₂	[1502-10-9] $\Delta_{\text{sub}}H$	N-methyl-7-(methylimino)-1,3,5-cycloheptatrienylamine	49.4 ± 4			[1971JAC/HUN, 1977PED/RYL]
C ₉ H ₁₂ N ₂	[31529-46-1] $\Delta_{\text{fus}}H$	N-amino-2-methylindoline	24.45	318.2		[1997PEY/LET]
C ₉ H ₁₂ N ₂ O	[101-42-8] $\Delta_{\text{fus}}H$	1,1-dimethyl-3-phenylurea	22.81	404.8	DSC	[1991ACR, 1990DON/DRE]
C ₉ H ₁₂ N ₂ O ₂	[13142-86-4] $\Delta_{\text{sub}}H$	3-ethoxyphenylurea	75.3 ± 8.3			[1954TSU/KAT, 1970COX/PIL]
C ₉ H ₁₂ N ₂ O ₂	[150-69-6] $\Delta_{\text{sub}}H$	4-ethoxyphenylurea (dulcin)	83.7 ± 8.3			[1954TSU/KAT, 1970COX/PIL]
C ₉ H ₁₂ N ₂ O ₂ S ₂	[2651-16-3] $\Delta_{\text{fus}}H$	S-methyl-N'-tosylisothiourea	31.2	401.2	DSC	[1992REI/HAN]
C ₉ H ₁₂ N ₂ S	[14222-60-7] $\Delta_{\text{fus}}H$	2-propyl-4-pyridinecarbothioamide	23.21	414.1		[2007ZHA/TAN]
C ₉ H ₁₂ N ₄ O ₂	[5770-28-5] $\Delta_{\text{fus}}H$	8-ethyltheophylline	37.2	545.3	DSC	[1989GON/KRA]
C ₉ H ₁₂ O	[4013-34-7] Δ_vH Δ_vH	(1-methoxyethyl)benzene	(298–313) 49.2 ± 0.4 (298–313) 49.1 ± 0.4	296 298	GS GS	[2001VER/HEI] [2001VER/HEI]
C ₉ H ₁₂ O	[539-30-0] Δ_vH Δ_vH	benzyl ethyl ether	(278–314) 53.5 ± 0.4 (299–460) 48.0	298 314	GS A	[2002KRA/VAS] [1987STE/MAL, 1947STU]
C ₉ H ₁₂ O	[14804-32-1] Δ_vH	2-ethylanisole	(302–460) 49.8	317	A	[1987STE/MAL, 1947STU]
C ₉ H ₁₂ O	[10568-38-4] Δ_vH	3-ethylanisole	(306–470) 49.3	321	A	[1987STE/MAL, 1947STU]
C ₉ H ₁₂ O	[1515-95-3] Δ_vH	4-ethylanisole	(306–470) 51.9	321	A	[1987STE/MAL, 1947STU]
C ₉ H ₁₂ O	[698-71-5] Δ_vH Δ_vH	5-ethyl-3-methylphenol	(468–521) 55.0 (385–506) 58.5	483 58.5	A,GS,EB	[1987STE/MAL, 1964HAN/HAR] [1955VON/GEB]
C ₉ H ₁₂ O	[88-69-7] Δ_vH Δ_vH Δ_vH Δ_vH	2-isopropylphenol	(375–493) 63.5 (370–489) 55.1 (375–493) 56.1 (335–501) 57.3	390 385 390 350	EB A	[1990NES/NAZ] [1987STE/MAL] [1986TSV/NAZ] [1947STU]
C ₉ H ₁₂ O	[618-45-1] Δ_vH	3-isopropylphenol	(377–497) 64.3	392	A	[1987STE/MAL]

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound					
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference	
C ₉ H ₁₂ O	[99-89-8]	4-isopropylphenol					
		$\Delta_v H$	(391–507)	63.7	406	EB	[1990NES/NAZ]
			(380–496)	63.1	395	A	[1987STE/MAL]
C ₉ H ₁₂ O	[2741-16-4]	isopropyl phenyl ether					
		$\Delta_v H$	(345–448)	49.5	360	A	[1987STE/MAL, 1965HEI/SUR, 1984BOU/FRI]
C ₉ H ₁₂ O	[122-97-4]	3-phenyl-1-propanol					
		$\Delta_v H$	(284–328)	62.8	299	A	[1987STE/MAL]
			(347–508)	62.6	362		[1947STU]
C ₉ H ₁₂ O	[617-94-7]	2-phenyl-2-propanol					
		$\Delta_v H$	(391–423)	52.9	406	A	[1987STE/MAL]
C ₉ H ₁₂ O	[622-85-5]	phenyl propyl ether					
		$\Delta_v H$	(374–463)	46.5	389	A	[1987STE/MAL]
C ₉ H ₁₂ O	[644-35-9]	2-propylphenol					
		$\Delta_v H$	(377–495)	56.9	392	A	[1987STE/MAL]
		$\Delta_v H$	(381–504)	59.9	398		[1953STA/MUL]
		$\Delta_v H$	(381–504)	57.2	423		[1953STA/MUL]
		$\Delta_v H$	(381–504)	53.0	473		[1953STA/MUL]
C ₉ H ₁₂ O	[621-27-2]	3-propylphenol					
		$\Delta_v H$	(408–538)	60.2	423	A	[1987STE/MAL]
		$\Delta_v H$	(386–512)	59.9	398		[1953STA/MUL]
		$\Delta_v H$	(386–512)	57.2	423		[1953STA/MUL]
		$\Delta_v H$	(386–512)	53.0	473		[1953STA/MUL]
C ₉ H ₁₂ O	[645-56-7]	4-propylphenol					
		$\Delta_v H$	(383–508)	56.7	398	A	[1987STE/MAL]
		$\Delta_v H$	(347–517)	61.3	348		[1953STA/MUL]
		$\Delta_v H$	(347–517)	59.5	373		[1953STA/MUL]
		$\Delta_v H$	(347–517)	58.4	398		[1953STA/MUL]
		$\Delta_v H$	(347–517)	56.2	423		[1953STA/MUL]
		$\Delta_v H$	(347–517)	51.5	473		[1953STA/MUL]
C ₉ H ₁₂ O	[697-82-5]	2,3,5-trimethylphenol					
		$\Delta_v H$	(459–521)	53.9	474	A,GS,EB	[1987STE/MAL, 1964HAN/HAR]
		$\Delta_v H$	(379–506)	55.1	394		[1955VON/GEB]
C ₉ H ₁₂ O	[2416-94-6]	2,3,6-trimethylphenol					
		$\Delta_{\text{fus}}H$		22.05	331.2		[1999VER]
		$\Delta_{\text{sub}}H$		86.7 ± 0.6	298	GS	[1999VER]
C ₉ H ₁₂ O	[496-78-6]	2,4,5-trimethylphenol					
		$\Delta_v H$	(379–505)	56.5	394	A	[1987STE/MAL, 1955VON/GEB]
C ₉ H ₁₂ O	[527-60-6]	2,4,6-trimethylphenol					
		$\Delta_{\text{sub}}H$		82.8 ± 0.3	298	GS	[1999VER]
		$\Delta_{\text{sub}}H$		95.0	298	C	[1971BER/GIR, 1999VER]
		$\Delta_v H$	(367–494)	53.2	382	A	[1987STE/MAL, 1955VON/GEB]
C ₉ H ₁₂ O	[2416-94-6]	2,5,6-trimethylphenol					
		$\Delta_v H$	(359–503)	51.1 ± 0.2	431		[1988BAG/GUR]
C ₉ H ₁₂ O	[527-54-8]	3,4,5-trimethylphenol					
		$\Delta_v H$	(396–521)	61.1	411	A	[1987STE/MAL]
C ₉ H ₁₂ O	[617-94-7]	α, α -dimethylbenzyl alcohol					
		$\Delta_{\text{sub}}H$	(276–302)	82.8 ± 0.7	289	GS	[1999VER4]

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
			82.3 ± 0.7	298		[1999VER4]
			63.4 ± 0.5	325	GS	[1999VER4]
			65.0 ± 0.5	298	GS	[1999VER4]
C ₉ H ₁₂ O ₂	[700-13-0]	trimethylhydroquinone				
		(450–501)	45.5 ± 0.3	475		[1988BAG/GUR]
C ₉ H ₁₂ O ₂	[80-15-9]	cumene hydroperoxide				
		(283–333)	69.9	298	A	[1987STE/MAL]
		(347–390)	74.0	362	A	[1987STE/MAL]
C ₉ H ₁₂ O ₂	[na]	1,3-dihydroxy-5-methyl-2-ethylbenzene				
		(388–453)	77.1	403	A,GC	[1987STE/MAL, 1975KUN/LIL]
C ₉ H ₁₂ O ₂	[4179-19-5]	3,5-dimethoxytoluene				
		(374–520)	59.5	389	A	[1987STE/MAL]
C ₉ H ₁₂ O ₂	[622-08-2]	ethylene glycol monobenzyl ether				
		(453–530)	58.6	468	A	[1987STE/MAL]
C ₉ H ₁₂ O ₂	[770-35-4]	propylene glycol 1-phenyl ether				
		(389–509)	59.5	404	A	[1987STE/MAL]
C ₉ H ₁₂ O ₂	[na]	isopropyl catechol (isomer not specified)				
		(393–453)	65.3	423		[1965GAK/BAB]
C ₉ H ₁₂ O ₂	[1125-88-8]	benzaldehyde dimethyl acetal				
		(278–318)	60.9 ± 0.5	298	GS	[2002VER]
		(283–318)	56.5 ± 0.7	300	GS	[1995VER/DOG]
C ₉ H ₁₂ O ₂	[2138-48-9]	3-isopropyl-1,2-dihydroxybenzene				
			97.8 ± 1.7	298	C	[1984RIB/RIB2]
C ₉ H ₁₂ O ₃	[634-36-6]	1,2,3-trimethoxybenzene				
			113.4 ± 0.3	375	C	[2000MAT/MIR]
			98.0 ± 0.3	298	C	[2000MAT/MIR]
C ₉ H ₁₂ O ₃	[621-23-8]	1,3,5-trimethoxybenzene				
			116.0 ± 1.9	375	C	[2000MAT/MIR]
			100.6 ± 1.9	298	C	[2000MAT/MIR]
			68.2 ± 2.0	298	CGC	[2000NIC/ORF]
C ₉ H ₁₂ O ₃	[538-43-2]	(racemic) 3-phenoxypropane-1,2-diol				
			28.0	331.7	DSC	[2008BRE/BRE, 2006ZAK/LAZ]
C ₉ H ₁₂ O ₃	[82430-38-4]	(R)-3-phenoxypropane-1,2-diol				
			31.8	341.5	DSC	[2008BRE/BRE, 2006ZAK/LAZ]
C ₉ H ₁₂ O ₃	[5662-95-3]	3,3-tetramethyleneglutaric acid anhydride				
			15.3	338.4	DSC	[2008MAT/MIR]
			96.4 ± 1.1	298	C	[2008MAT/MIR]
C ₉ H ₁₂ O ₄	[642-71-7]	3,4,5-trimethoxyphenol				
			31.94	420.2		[2008MAT/MIR2]
C ₉ H ₁₂ S	[6263-62-3]	benzyl ethyl sulfide				
		(346–370)	56.0	358		[1999DYK/SVO]
		(345–500)	54.8	360	A	[1987STE/MAL]
			56.9 ± 2.1	298		[1962MAC/MAY]
C ₉ H ₁₂ S	[20760-06-9]	2-ethylthioanisole				
		(481–511)	44.3	496		[1999DYK/SVO]

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound					
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference	
C ₉ H ₁₂ S	[34786-24-8] $\Delta_v H$	ethyl <i>m</i> -tolyl sulfide (472–502)	43.5	487		[1999DYK/SVO]	
C ₉ H ₁₂ S	[622.63-9] $\Delta_v H$	ethyl <i>p</i> -tolyl sulfide (473–503)	43.6	488		[1999DYK/SVO]	
C ₉ H ₁₂ S	[3019-20-3] $\Delta_v H$	(isopropylthio)benzene (461–491)	U23.6	476		[1999DYK/SVO]	
C ₉ H ₁₂ S	[874-79-3] $\Delta_v H$	(propylthio)benzene (473–503)	44.3	488		[1999DYK/SVO]	
C ₉ H ₁₃ BrN ₂ O ₂	[314-40-9] $\Delta_{\text{fus}} H$	5-bromo-6-methyl-3-(1-methylpropyl)-2,4-(1 <i>H</i> ,3 <i>H</i>)-pyrimidinedione	22.02	428.3	DSC	[1991ACR, 1990DON/DRE]	
C ₉ H ₁₃ ClN ₆	[21725-46-2] $\Delta_{\text{fus}} H$	2-[[4-chloro-6-(ethylamino)-1,3,5-triazin-2-yl]amino]-2-methylpropanenitrile	41.96	437.9	DSC	[1991ACR, 1990DON/DRE]	
C ₉ H ₁₃ ClN ₆	[21725-46-2] $\Delta_{\text{sub}} H$	2-[(4-chloro-6-ethylamino- <i>s</i> -triazin-2-yl)amino]-2-methylpropionitrile (cyanazine) (339–365)	90.7	352	GS	[1982GRA/FOS]	
C ₉ H ₁₃ Cl ₃ NO ₄ P	[5598-15-2] $\Delta_{\text{fus}} H$	O,O-diethyl-O-(3,5,6-trichloro-2-pyridyl) phosphate	15.61	312.5	DSC	[1990DON/DRE]	
C ₉ H ₁₃ Cl ₃ OS	[76619-95-9] $\Delta_v H$	2,3,3-trichloro-2-propenethioic acid, O-hexyl ester (433–483)	69.5		GC	[1980PIT/KIS]	
C ₉ H ₁₃ N	[na] $\Delta_v H$	α, α -dimethylbenzylamine (283–323)	56.4 ± 0.7	303	GS	[1999VER4]	
	$\Delta_v H$		56.7 ± 0.7	298	GS	[1999VER4]	
C ₉ H ₁₃ N	[103-83-3] $\Delta_v H$	N,N-dimethylbenzylamine (288–328)	48.9 ± 0.4	308	GS	[1999VER4]	
			$\Delta_v H$	49.5 ± 0.4	298	GS	[1999VER4]
			$\Delta_v H$	50.1 ± 0.9	298	C	[1996MIR/ORL]
C ₉ H ₁₃ N	[609-72-3] $\Delta_v H$	N,N-dimethyl-2-toluidine (301–458)	52.4	316	A	[1987STE/MAL, 1947STU]	
C ₉ H ₁₃ N	[121-72-2] $\Delta_v H$	N,N-dimethyl-3-toluidine	58.2 ± 6.9	298	CGC	[1996RIB/RIB]	
C ₉ H ₁₃ N	[99-97-8] $\Delta_v H$	N,N-dimethyl-4-toluidine (323–483)	60.7	338	A	[1987STE/MAL, 1947STU]	
C ₉ H ₁₃ N	[102-27-2] $\Delta_v H$	N-ethyl-3-toluidine	60.0 ± 3.0	298	CGC	[1996RIB/RIB]	
C ₉ H ₁₃ N	[643-28-7] $\Delta_v H$	2-isopropylaniline (286–326)	61.3 ± 0.9	306	GS	[2000VER3]	
			$\Delta_v H$	61.8 ± 0.9	298	GS	[2000VER3]
C ₉ H ₁₃ N	[99-88-7] $\Delta_v H$	4-isopropylaniline (333–500)	57.5	348	A	[1987STE/MAL, 1947STU]	
C ₉ H ₁₃ N	[300-62-9] $\Delta_v H$	1-phenyl-2-propylamine (333–353)	53.4	343	A	[1987STE/MAL]	
C ₉ H ₁₃ N	[88-05-1] $\Delta_v H$	2,4,6-trimethylaniline (341–510)	64.1	356	A	[1987STE/MAL, 1947STU]	
C ₉ H ₁₃ N	[3978-81-2] $\Delta_v H$	4- <i>tert</i> -butylpyridine	54.4 ± 1.3	298	C	[2008FRI/ACR]	
C ₉ H ₁₃ NO	[492-41-1]	(–) 2-amino-1-phenyl-1-propanol (norephedrine)					

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{fus}}H$		15.87	324.4	DSC	[1999LI/ZEL]
C ₉ H ₁₃ NO	[14838-15-4]	(±) 2-amino-1-phenyl-1-propanol (norephedrine)	$\Delta_{\text{fus}}H$ 26.11	374.3	DSC	[1999LI/ZEL]
C ₉ H ₁₃ NO	[104-63-2]	N-benzylethanolamine	Δ_vH (293–363) 71.7 Δ_vH (293–363) 74.5	328 298		[2009RAZ/HAJ] [2009RAZ/HAJ]
C ₉ H ₁₃ NO ₂	[1075-89-4]	3,3-tetramethyleneglutarimide	$\Delta_{\text{fus}}H$ 24.2	426.6	DSC	[2008MAT/MIR]
	$\Delta_{\text{sub}}H$		106.8 ± 2.0	298	C	[2008MAT/MIR]
C ₉ H ₁₃ N ₅	[139909-52-7]	6,9-dimethyl-8-ethyladenine	$\Delta_{\text{fus}}H$ 29.8	436.8		[1994ZIE/ZIE]
	$\Delta_{\text{sub}}H$	(345–351)	94.1 ± 0.1	348	ME	[1994ZIE/ZIE]
C ₉ H ₁₃ N ₅	[117954-97-9]	8-propyl-9-methyladenine	$\Delta_{\text{sub}}H$ (364–370) 124.2 ± 0.8	367	ME	[1987KAM/ZIE]
C ₉ H ₁₃ N ₅ O ₄	[82410-32-0]	9-[(1,3-dihydroxy-2-propoxy)methyl]guanine	$\Delta_{\text{fus}}H$ 37.88	509.2		[1999ZIE/GOL]
C ₉ H ₁₄	[na]	1-ethyltricyclo[2,2,1,0 ^{2,6}]heptane	Δ_vH 42.0 ± 0.1	298	C	[1996VAR/PAS]
C ₉ H ₁₄	[2972-20-5]	2-methylenebicyclo[2.2.2]octane	Δ_vH 45.2			[1974KOZ/BYC]
C ₉ H ₁₄	[4893-13-4]	2-methylbicyclo[2.2.2]oct-2-ene	Δ_vH (363–402) 40.2 Δ_vH 43.5 ± 0.4	378 298	A EB	[1987STE/MAL] [1974VAR/DRU, 1974KOZ/BYC]
C ₉ H ₁₄	[2146-39-6]	2-vinylbicyclo[2.2.1]heptane	Δ_vH (350–385) 38.6	365	A	[1987STE/MAL]
C ₉ H ₁₄	[7124-86-9]	bicyclo[3.2.2]non-6-ene	$\Delta_{\text{sub}}H$ 48 ± 1.0	298	C	[1982JOC/DEK2]
C ₉ H ₁₄	[6671-66-5]	bicyclo[3.3.1]non-2-ene	$\Delta_{\text{sub}}H$ 48.2 ± 0.4	298	C	[1982JOC/DEK2]
C ₉ H ₁₄	[1456-33-0]	bicyclo[4.2.1]non-3-ene	$\Delta_{\text{sub}}H$ 49.7 ± 0.8	298	C	[1982JOC/DEK2]
C ₉ H ₁₄ ClN ₅	[139-40-2]	2-chloro-4,6-bis(isopropylamino)-1,3,5-triazine	$\Delta_{\text{fus}}H$ 41.87	490.3	DSC	[1991ACR, 1990DON/DRE]
C ₉ H ₁₄ F ₃ NO ₃	[1115-39-5]	N-trifluoroacetyl-L-leucine, methyl ester	Δ_vH (273–463) 55.9	288	A	[1987STE/MAL, 1999DYK/SVO]
C ₉ H ₁₄ N ₂	[1675-69-0]	azelaic acid dinitrile	Δ_vH (308–341) 80.4	323	A	[1987STE/MAL]
C ₉ H ₁₄ N ₂	[1675-69-0]	azelanitrile	$\Delta_{\text{fus}}H$ 18.68	251.1	DSC	[2007BAD/BLA]
C ₉ H ₁₄ N ₂ O ₂	[82413-39-6]	1,3-dimethyl-5-propyluracil	$\Delta_{\text{fus}}H$ 26.3	355		[1996KAM/ZIE]
	$\Delta_{\text{sub}}H$	(317–327)	111.0 ± 1.6	322	ME	[1996KAM/ZIE]
C ₉ H ₁₄ N ₂ O ₂	[175412-48-3]	1,3-dimethyl-5-isopropyluracil				

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		T _m (K)	Method	Reference	
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)				
		$\Delta_{\text{fus}}H$	22.4	354.7		[1996KAM/ZIE]	
		$\Delta_{\text{sub}}H$	(316–328)	102.9 ± 1.6	322	ME	[1996KAM/ZIE]
C ₉ H ₁₄ N ₂ O ₂	[21472-93-5]	1,3-diethylthymine					
		$\Delta_{\text{sub}}H$	89.8 ± 0.4	298	C	[1980SAB/KOM]	
		$\Delta_{\text{sub}}H$	(307–325)	95.0 ± 2.1	317	QR	[1980TEP/YAN]
C ₉ H ₁₄ O	[281-24-3]	2-oxaadamantane					
		$\Delta_{\text{fus}}H$	8.12	567	DSC	[1978AND/CAR]	
C ₉ H ₁₄ O	[17931-55-4]	bicyclo[3.3.1]nonan-9-one					
		$\Delta_{\text{fus}}H$	13.99	299	DSC	[1998PAR/GIL2]	
		$\Delta_{\text{fus}}H$	14.11	300.5	AC	[1991WHI/PER]	
C ₉ H ₁₄ O	[5689-04-3]	<i>cis</i> 2-hexahydroindanone					
		Δ_vH	57.5	298		[1971SEL3]	
C ₉ H ₁₄ O	[16484-17-6]	<i>trans</i> 2-hexahydroindanone					
		Δ_vH	56.1	298		[1971SEL3]	
C ₉ H ₁₄ O	[20030-30-2]	2,5,6-trimethyl-2-cyclohexen-1-one					
		Δ_vH	(371–478)	45.5 ± 0.3	425		[1988BAG/GUR]
C ₉ H ₁₄ O	[78-59-1]	3,3,5-trimethyl-2-cyclohex-1-one (isophorone)					
		Δ_vH	(311–489)	48.6	326	A	[1987STE/MAL, 1947STU]
C ₉ H ₁₄ O	[504-20-1]	2,6-dimethyl-2,5-heptadien-4-one (phorone)					
		Δ_vH	(315–471)	54.1	330	A	[1987STE/MAL, 1947STU]
C ₉ H ₁₄ O ₂	[na]	bicyclo[2.2.1]heptan-7-one ethylene ketal					
		Δ_vH	(283–318)	53.8 ± 0.2		GS	[1998VER/PEN, 2002VER]
C ₉ H ₁₄ O ₂	[111-12-6]	methyl 2-octynoate					
		Δ_vH	(283–312)	64.5	297	A,ME	[1987STE/MAL, 1955SER/VOI]
C ₉ H ₁₄ O ₄	[691-83-8]	diethyl citraconate					
		Δ_vH	(332–504)	54.9	347	A	[1987STE/MAL, 1947STU]
C ₉ H ₁₄ O ₄	[2409-52-1]	diethyl itaconate					
		Δ_vH	(324–501)	51.0	339	A	[1987STE/MAL, 1947STU]
C ₉ H ₁₄ O ₄	[2418-31-7]	diethyl mesaconate					
		Δ_vH	(335–502)	55.9	350	A	[1987STE/MAL, 1947STU]
C ₉ H ₁₄ O ₄	[1559-02-0]	1,1-cyclopropanedicarboxylic acid diethyl ester					
		Δ_vH	(288–318)	63.9 ± 0.5		GS	[1998VER/KUM]
C ₉ H ₁₄ O ₄	[16713-66-9]	3,3-tetramethyleneglutaric acid					
		$\Delta_{\text{fus}}H$	32.1	452.9	DSC	[2008MAT/MIR]	
		$\Delta_{\text{sub}}H$	126.9 ± 2.4	298	C	[2008MAT/MIR]	
C ₉ H ₁₄ O ₅	[570-08-1]	diethyl acetylmalonate					
		Δ_vH	(363–510)	54.0	378	A	[1987STE/MAL]
C ₉ H ₁₄ O ₅	[na]	ethyl[(1-allyloxycarbonyl)ethyl] carbonate					
		Δ_vH	(342–496)	61.3	357	A	[1987STE/MAL]
C ₉ H ₁₄ O ₅	[na]	2-lactyloxypropionic acid, allyl ester					
		Δ_vH	(331–401)	75.1	346	A	[1987STE/MAL]
C ₉ H ₁₄ O ₅	[57822-06-7]	3-oxononanedioic acid					
		$\Delta_{\text{sub}}H$	(312–330)	118		TPTD	[2005CHA/ZIE]

Note: Values based on TPTD method are not consistent with values determined by other experimental methods

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₉ H ₁₄ O ₆	[102-76-1]	glycerol triacetate				
	$\Delta_{\text{fus}}H$		25.8	275.3		[1996DOM/HEA]
	Δ_vH	(320–361)	81.9 ± 0.3	298	GS	[2009VER/EME2]
	Δ_vH	(440–590)	83.8 ± 0.9	298	EB	[1990DAU/HUT, 2009VER/EME2]
	Δ_vH	(284–319)	82.0	299	A	[1987STE/MAL, 1963WOO/ADI]
	Δ_vH		85.7 ± 0.3	298	C	[1986NIL/WAD]
C ₉ H ₁₄ O ₇	[1587-20-8]	trimethyl citrate				
	Δ_vH	(379–560)	617.4	394	A	[1987STE/MAL, 1947STU]
C ₉ H ₁₄ S	[281-25-4]	2-thiaadamantane				
	$\Delta_{\text{ms}}H$		1.95	211		
	$\Delta_{\text{fus}}H$		8.1	597	DSC	[1978AND/CAR]
C ₉ H ₁₄ S	[4861-58-9]	2-pentylthiophene				
	Δ_vH		52.0 ± 1.2	298	C	[2007RIB/SAN]
C ₉ H ₁₅ Cl ₃ O ₂	[na]	3-chloro-2,2-bis(chloromethyl)propyl butyrate				
	Δ_vH	(426–482)	73.6	441	A	[1987STE/MAL, 1999DYK/SVO]
C ₉ H ₁₅ NOS	[59300-33-3]	carbamothioic acid, (1-methylethyl)-2-propynyl-S-ethyl ester				
	Δ_vH	(298–313)	72.8	305	A	[1987STE/MAL, 1999DYK/SVO]
C ₉ H ₁₅ NOS	[59300-32-2]	carbamothioic acid, propyl-2-propynyl-S-ethyl ester				
	Δ_vH	(298–313)	64.6	305	A	[1987STE/MAL]
C ₉ H ₁₅ NO ₃ S	[62571-86-2]	1-[(2S)-3-mercapto-2-methyl-1-oxopropyl]-(<i>l</i>)-proline				
	$\Delta_{\text{fus}}H$		20.38	379.5	DSC	[2008STU/ROR]
C ₉ H ₁₅ N ₃ O ₃	[82859-98-1]	N-acetylglycyl-(<i>l</i>)-prolinamide				
	$\Delta_{\text{ms}}H$		5.6	450.6		
	$\Delta_{\text{fus}}H$		27.0	457.8		[1996PUL/BAR]
C ₉ H ₁₅ N ₃ O ₃	[52186-41-1]	N-acetyl-(<i>l</i>)-prolyl-glycinamide				
	$\Delta_{\text{fus}}H$		32.2	434.1	DSC	[1992BAR/GIA]
C ₉ H ₁₅ N ₃ O ₈	[34001-52-0]	neopentyl-4,4,4-trinitrobutyrate				
	$\Delta_{\text{fus}}H$		22.59	333.5	DSC	[1971ROS/HOL]
C ₉ H ₁₆	[3452-09-3]	1-nonyne				
	Δ_vH	(320–464)	45.6 ± 0.2	320	EB	[2002STE/CHI4]
	Δ_vH	(320–464)	42.7 ± 0.2	360	EB	[2002STE/CHI4]
	Δ_vH	(320–464)	39.7 ± 0.3	400	EB	[2002STE/CHI4]
	Δ_vH	(320–464)	36.4 ± 0.5	440	EB	[2002STE/CHI4]
C ₉ H ₁₆	[39124-79-3]	<i>trans</i> bicyclo[6.1.0]nonane				
	Δ_vH		42.7 ± 0.6		EB	[1978COR/PER]
C ₉ H ₁₆	[13757-43-2]	<i>cis</i> bicyclo[6.1.0]nonane				
	Δ_vH		49.8 ± 0.8		EB	[1978COR/PER]
	Δ_vH	(297–360)	50.4 ± 0.8	312	A	[1987STE/MAL, 1970CHA/MCN]
C ₉ H ₁₆	[20454-81-3]	1,4-dimethylbicyclo[2.2.1]heptane				
	Δ_vH	(328–393)	36.8	343	A	[1987STE/MAL, 1970VAR/BEL, 1984BOU/FRI]
C ₉ H ₁₆	[20558-16-1]	<i>trans</i> 2,3-dimethylbicyclo[2.2.1]heptane				
	Δ_vH	(345–411)	39.3	360	A	[1987STE/MAL, 1970VAR/BEL, 1984BOU/FRI]
C ₉ H ₁₆	[2146-41-0]	2-ethylbicyclo[2.2.1]heptane				
	Δ_vH	(349–396)	44.4	364	A	[1987STE/MAL]

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₉ H ₁₆	[4551-51-3]	<i>cis</i> hexahydroindan				
	$\Delta_{\text{us}}H$		8.26	182.3		
	$\Delta_{\text{us}}H$		0.39	184.5		
	$\Delta_{\text{fus}}H$		1.4	236.5		[1972FIN/MCC]
	Δ_vH	(263–293)	47.1	278	A	[1987STE/MAL]
	Δ_vH	(290–366)	45.9	305	A	[1987STE/MAL]
	Δ_vH	(363–463)	41.9	378	A	[1987STE/MAL]
C ₉ H ₁₆	[3296-50-2]	<i>trans</i> hexahydroindan				
	$\Delta_{\text{fus}}H$		10.9	213.9		[1972FIN/MCC]
	Δ_vH	(281–362)	45.1	296	A	[1987STE/MAL]
	Δ_vH	(356–457)	41.1	371	A	[1987STE/MAL]
	Δ_vH	(262–283)	45.9	272	A	[1987STE/MAL]
	Δ_vH	(358–479)	41.0	373	A	[1987STE/MAL]
	Δ_vH	(345–435)	41.6	360	GS	[1955CAM/ROS]
C ₉ H ₁₆	[2114-42-3]	allylcyclohexane				
	Δ_vH		44.0 ± 0.2	298	GCC	[1979FUC/PEA]
C ₉ H ₁₆	[1003-64-1]	ethylidenecyclohexane				
	Δ_vH		42.0 ± 0.2	298	GCC	[1979FUC/PEA]
C ₉ H ₁₆	[175-93-9]	spiro[4.4]nonane				
	Δ_vH	(278–313)	44.5 ± 0.6	298	GS	[2002VER]
C ₉ H ₁₆	[260-65-9]	bicyclo[3.3.1]nonane				
	$\Delta_{\text{sub}}H$		50.6 ± 2	298	TSGC	[1977PAR/STE]
C ₉ H ₁₆ ClN ₅	[139-40-2]	2-chloro-4,6-bis(isopropylamino)-1,3,5-triazine				
	$\Delta_{\text{sub}}H$	(323–403)	125.1	338	GS-GC	[1987STE/MAL, 1964FRI/SAM]
C ₉ H ₁₆ ClN ₅	[5915-41-3]	6-chloro-N-(1,1-dimethylethyl)-N'-ethyl-1,3,5-triazine-2,4-diamine				
	$\Delta_{\text{fus}}H$		33.57	448.6	DSC	[1990DON/DRE]
C ₉ H ₁₆ Cl ₄	[1561-48-4]	1,1,1,9-tetrachlorononane				
	Δ_vH	(303–434)	78.0	318		[1999DYK/SVO]
	Δ_vH	(298–338)	89.0	313	A	[1987STE/MAL]
C ₉ H ₁₆ NO ₂	[2896-70-0]	2,2,6,6-tetramethyl-4-oxopiperidine-1-oxyl				
	$\Delta_{\text{sub}}H$		83.3 ± 1.7		ME	[1965KAL/ROZ, 1970COX/PIL, 1987STE/MAL]
C ₉ H ₁₆ N ₂	[2273-41-8]	2-methyl-2-piperidinopropionitrile				
	$\Delta_{\text{fus}}H$		21.59	316.2		[1997WEL/VER]
	$\Delta_{\text{sub}}H$		80.3 ± 0.5	298		[1997WEL/VER]
C ₉ H ₁₆ N ₄ OS	[34014-18-1]	N-[5-(1,1-dimethylethyl)-1,3,4-thiadiazol-2-yl]-N,N'-dimethylurea				
	$\Delta_{\text{fus}}H$		29.48	435.3	DSC	[1991ACR, 1990DON/DRE]
	$\Delta_{\text{sub}}H$		1.6	298		[1998GON/SZW]
C ₉ H ₁₆ O	[3350-30-9]	cyclononanone				
	$\Delta_{\text{us}}H$		14.7	247		
	$\Delta_{\text{fus}}H$		1.6	298		[1998GON/SZW]
	Δ_vH	(333–413)	51.4	348	A	[1987STE/MAL]
C ₉ H ₁₆ O	[2890-62-2]	1-(1-methyl-3-cyclohexen-3-yl)ethanol				
	Δ_vH	(358–410)	54.6	373	A	[1987STE/MAL]
	Δ_vH		53.1 ± 0.6	298		[1972WOL]

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₉ H ₁₆ O	[2890-62-2] $\Delta_v H$	methyl (1-methylcyclohexyl) ketone (374–414)	46.1	389	A	[1987STE/MAL]
C ₉ H ₁₆ O	[18829-56-6] $\Delta_v H$	<i>trans</i> 2-nonenal (363–398)	56.1	378	A	[1987STE/MAL]
C ₉ H ₁₆ O	[873-94-9] $\Delta_v H$	(<i>dl</i>) 3,5,5-trimethylcyclohexanone (423–463)	39.3	438	A	[1987STE/MAL]
C ₉ H ₁₆ O	[1000-30-2] $\Delta_v H$	2,5,5-trimethyl-4-hexene-1-al (293–353)	57.0	308	A	[1987STE/MAL]
C ₉ H ₁₆ OS	[22842-41-7] $\Delta_v H$	tetrahydro-2,2,6,6-tetramethyl-4 <i>H</i> -thiopyran-4-one (300–360)	34.7	315	A	[1987STE/MAL, 1972GEI/SAW, 1999DYK/SVO]
C ₉ H ₁₆ O ₂	[6222-35-1] $\Delta_v H$	cyclohexyl propanoate (253–293)	55.9 ± 0.1	298	C	[2004PAU/ZAI, 2003ZAI/VER]
	$\Delta_v H$		59.4 ± 0.8	298	ME	[2003ZAI/VER]
	$\Delta_v H$		56.4 ± 0.5	298	ME	[2003ZAI/VER]
	$\Delta_v H$		54.3 ± 0.4	298	GS	[2003ZAI/VER]
	$\Delta_v H$		56.8	298	CGC	[1999VER/HEI]
C ₉ H ₁₆ O ₂	[16737-30-7] $\Delta_v H$	1-methylcyclohexyl acetate (333–378)	52.4	298	CGC	[1999VER/HEI]
C ₉ H ₁₆ O ₂	[66922-08-5] $\Delta_v H$	3-methylcyclohexyl acetate (333–378)	53.6	298	CGC	[1999VER/HEI]
C ₉ H ₁₆ O ₂	[22597-23-5] $\Delta_v H$	4-methylcyclohexyl acetate (333–378)	54.1	298	CGC	[1999VER/HEI]
C ₉ H ₁₆ O ₂	[39869-70-0] $\Delta_v H$	bicyclo[2.2.1]heptan-7-one dimethyl ketal (283–318)	50.2 ± 0.2		GS	[1998VER/PEN, 2002VER]
C ₉ H ₁₆ O ₂	[5726-19-2] $\Delta_v H$	acetic acid, 2-methylcyclohexyl ester, mixed isomers (337–457)	49.0	353	A	[1987STE/MAL]
C ₉ H ₁₆ O ₂	[61732-95-4] $\Delta_v H$	2-butyl-4,7-dihydro-1,3-dioxepine (318–453)	50.9	333	A	[1987STE/MAL]
C ₉ H ₁₆ O ₂	[2499-95-8] $\Delta_v H$	hexyl acrylate (342–461)	48.2	357	A	[1987STE/MAL]
C ₉ H ₁₆ O ₂	[na] $\Delta_v H$	methacrylic acid, neopentyl ester (313–338)	40.5	325	A	[1987STE/MAL]
C ₉ H ₁₆ O ₂	[6008-27-1] $\Delta_v H$	oxo-2-cyclodecanone (nonanolactone) (352–381)	54.5 ± 0.2	366	MM	[1991WIB/WAL]
	$\Delta_v H$		59.0 ± 1.3	298	MM	[1991WIB/WAL]
	$\Delta_v H$		60.9	348	A	[1987STE/MAL]
C ₉ H ₁₆ O ₂	[104-61-0] $\Delta_v H$	γ -nonanolactone (296–363)	70.3 ± 0.2	298	GS	[2008EME/KOZ, 2009EME/VER]
C ₉ H ₁₆ O ₂	[3301-94-8] $\Delta_v H$	δ -nonanolactone (293–348)	70.7 ± 0.4	298	GS	[2007EME/KOZ]
C ₉ H ₁₆ O ₂	[18362-64-6] $\Delta_v H$	2,6-dimethyl-3,5-heptanedione 56.1		298		[1978RIB/IRV]
C ₉ H ₁₆ O ₂	[2849-98-1] $\Delta_v H$	pentyl methacrylate (339–456)	47.6	354	A	[1987STE/MAL]
C ₉ H ₁₆ O ₂	[54056-51-8]	butyl 3-methylbut-2-enoate				

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(279–323)	56.6 ± 0.3	298	GS	[2008EME/TOK]
C ₉ H ₁₆ O ₂	[30434-54-9]	isobutyl 3-methylbut-2-enoate				
	$\Delta_v H$	(280–323)	54.7 ± 0.2	298	GS	[2008EME/TOK]
C ₉ H ₁₆ O ₃	[2052-15-5]	butyl levulinate				
	$\Delta_v H$	(338–511)	55.5	373	A	[1987STE/MAL]
	$\Delta_v H$		56.0	452		[1931SCH/COW]
C ₉ H ₁₆ O ₃	[2052-15-5]	sec-butyl levulinate				
	$\Delta_v H$	(393–499)	51.0	408	A	[1987STE/MAL]
C ₉ H ₁₆ O ₃	[3757-32-2]	isobutyl levulinate				
	$\Delta_v H$	(338–503)	61.5	353	A	[1987STE/MAL, 1947STU]
	$\Delta_v H$		54.7	444		[1931SCH/COW]
C ₉ H ₁₆ O ₃	[na]	4-methyl-1-propyl-2,6,7-trioxabicyclo[2.2.2]octane				
	$\Delta_{\text{fus}} H$		16.1	311.2		[1995RAK/VER2]
C ₉ H ₁₆ O ₄	[na]	2-acetoxypropionic acid, butyl ester				
	$\Delta_v H$	(325–485)	63.2	340	A	[1987STE/MAL]
C ₉ H ₁₆ O ₄	[40326-38-3]	3-acetoxypropionic acid, butyl ester				
	$\Delta_v H$	(373–391)	75.4	382	A	[1987STE/MAL]
C ₉ H ₁₆ O ₄	[818-38-2]	diethyl glutarate				
	$\Delta_v H$	(338–510)	55.7	353	A	[1987STE/MAL, 1947STU]
C ₉ H ₁₆ O ₄	[133-13-1]	ethylmalonic acid, diethyl ester				
	$\Delta_v H$	(323–485)	55.3	338	A	[1987STE/MAL, 1947STU]
C ₉ H ₁₆ O ₄	[123-99-9]	nonanedioic acid (azelaic acid)				
	$\Delta_{\text{fus}} H$		35.3	375.6	DSC	[2009CHE/XIA]
		Note: Authors explicitly state in the manuscript that no solid-solid phase transition was observed				
	$\Delta_{\text{trs}} H$		0.01	330.6		
	$\Delta_{\text{trs}} H$		0.7	339.8		
	$\Delta_{\text{fus}} H$		29.7	372.4	DSC	[2005ROU/TEM]
	$\Delta_{\text{fus}} H$		32.67	380		[1991ACR]
	$\Delta_{\text{sub}} H$	(348–373)	178 ± 5		TPD	[2007CAP/LOV]
	$\Delta_{\text{sub}} H$	(294–311)	138		TPTD	[2005CHA/ZIE]
		Note: Values based on TPTD method are not consistent with values determined by other experimental methods				
	$\Delta_{\text{sub}} H$	(367–377)	156.2 ± 0.5	372	ME	[1999RIB/MON]
	$\Delta_{\text{sub}} H$	(367–377)	159.9 ± 1.0	298	ME	[1999RIB/MON]
	$\Delta_v H$	(434–503)	119.7 ± 0.8	298	CGC	[2005ROU/TEM]
	$\Delta_v H$	(451–630)	89.3	466	A	[1987STE/MAL, 1947STU]
C ₉ H ₁₆ O ₄	[1732-08-7]	dimethyl pimelate				
	$\Delta_v H$	(291–353)	73.5 ± 0.3	298	GS	[2006VER/KOZ]
C ₉ H ₁₆ O ₅	[na]	butyl[1-(methoxycarbonyl)ethyl] carbonate				
	$\Delta_v H$	(349–510)	61.7	364	A	[1987STE/MAL]
C ₉ H ₁₆ O ₅	[na]	isobutyl[1-(methoxycarbonyl)ethyl] carbonate				
	$\Delta_v H$	(340–501)	59.1	355	A	[1987STE/MAL]
C ₉ H ₁₆ O ₅	[na]	2-lactoylpropionic acid, propyl ester				
	$\Delta_v H$	(327–397)	73.5	342	A	[1987STE/MAL]
C ₉ H ₁₆ O ₅	[na]	methyl[1-(butoxycarbonyl)ethyl] carbonate				
	$\Delta_v H$	(311–503)	60.2	326	A	[1987STE/MAL]
C ₉ H ₁₇ N	[767-92-0]	<i>trans</i> (R,S)-decahydroquinoline				

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
		$\Delta_{\text{fus}}H$	25.72	321.4		[1994STE/CHI]
		Δ_vH	(325–525) 50.4	340	EB,IP	[1994STE/CHI]
		Δ_vH	(325–525) 47.6	380	EB,IP	[1994STE/CHI]
		Δ_vH	(325–525) 45.0	420	EB,IP	[1994STE/CHI]
		Δ_vH	(325–525) 42.3	460	EB,IP	[1994STE/CHI]
		Δ_vH	(325–525) 39.5	500	EB,IP	[1994STE/CHI]
C₉H₁₇N	[2243-27-8]	octyl cyanide				
		Δ_vH	(285–323) 62.0 ± 0.3	298	GS	[2005EME/VER]
		Δ_vH	(328–503) 56.8	343	A	[1987STE/MAL]
		Δ_vH	(314–480) 58.0	298	EB	[1941RAL/SEL, 2005EME/VER]
C₉H₁₇NO	[2896-70-0]	2,2,6,6-tetramethyl-4-oxopiperidine				
		$\Delta_{\text{sub}}H$	60.8 ± 2.7		ME	[1966LEB/ROS, 1970COX/PIL]
C₉H₁₇NO	[14952-05-7]	<i>trans</i> 2-nonenic acid amide				
		$\Delta_{\text{sub}}H$	(383–393) 111.9	388	A	[1987STE/MAL]
C₉H₁₇NO₂	[3637-11-4]	2,2,6,6-tetramethyl-1-hydroxy-4-oxopiperidine				
		$\Delta_{\text{sub}}H$	(288–328) 80.0	303	A	[1987STE/MAL]
		$\Delta_{\text{sub}}H$	80.1 ± 4.6		ME	[1965KAL/ROZ, 1970COX/PIL]
C₉H₁₇NO₃	[56430-36-5]	(<i>dl</i>) N-acetylvaline ethyl ester				
		Δ_vH	(382–466) 67.7	397	A	[1987STE/MAL]
C₉H₁₇NO₃S	[33280-93-2]	(<i>dl</i>) N-acetylmethionine ethyl ester				
		Δ_vH	(432–519) 81.6	447	A	[1987STE/MAL, 1999DYK/SVO]
C₉H₁₇N₅O	[1610-17-9]	2-methoxy-4-ethylamino-6-isopropylamino-1,3,5-triazine				
		$\Delta_{\text{sub}}H$	(323–403) 94.4	338	GS-GC	[1987STE/MAL, 1964FRI/SAM]
C₉H₁₇N₅S	[834-12-8]	2-methylthio-4-ethylamino-6-isopropylamino-1,3,5-triazine (ametryn)				
		$\Delta_{\text{fus}}H$	26.0	359.1	DSC	[2007VEC/BRU]
		$\Delta_{\text{sub}}H$	(323–403) 100.9	338	GS-GC	[1987STE/MAL, 1964FRI/SAM]
		Δ_vH	91 ± 4	466	DSC	[2007VEC/BRU]
		Δ_vH	125 ± 6	298	DSC	[2007VEC/BRU]
		Δ_vH	84.9 ± 1.3	453	TGA	[2007VEC/BRU]
		Δ_vH	118 ± 4	298	TGA	[2007VEC/BRU]
C₉H₁₈	[2040-95-1]	butylcyclopentane				
		$\Delta_{\text{fus}}H$	11.31	165.2		[1996DOM/HEA]
		Δ_vH	(413–432) 39.4	422	A	[1987STE/MAL]
		Δ_vH	43.8 ± 0.1	328	C	[1981SVO/CHA]
		Δ_vH	42.7 ± 0.1	343	C	[1981SVO/CHA]
		Δ_vH	41.6 ± 0.1	358	C	[1981SVO/CHA]
		Δ_vH	40.9 ± 0.1	368	C	[1981SVO/CHA]
		Δ_vH	46.0	298	298	[1971WIL/ZWO]
C₉H₁₈	[19489-10-2]	<i>cis</i> 1-ethyl-3-methylcyclohexane				
		Δ_vH	(373–465) 39.0	388	A	[1987STE/MAL]
C₉H₁₈	[696-29-7]	isopropylcyclohexane				
		Δ_vH	(295–431) 44.1	310	A	[1987STE/MAL]
		Δ_vH	(344–429) 41.1	359		[1949FOR/NOR]
C₉H₁₈	[1678-92-8]	propylcyclohexane				
		$\Delta_{\text{fus}}H$	10.6	178		[2006MAN/CUT]
		$\Delta_{\text{fus}}H$	10.37	178.3		[1991ACR]
		Δ_vH	42.8 ± 0.5	298	GC	[1987AZA]

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$		44.7 ± 0.4	298	GCC	[1978FUC/PEA]
	$\Delta_v H$		45.2	298		[1971WIL/ZWO]
	$\Delta_v H$		45.1	298		[1965FIN/MES]
	$\Delta_v H$		45.2	298	C	[1947OSB/GIN]
	$\Delta_v H$	(346–431)	41.7	361	A,MM	[1987STE/MAL, 1947STU, 1949FOR/NOR]
C₉H₁₈	[3073-66-3]	1,1,3-trimethylcyclohexane				
	$\Delta_v H$	(348–411)	37.7	363	A	[1987STE/MAL]
	$\Delta_v H$	(327–410)	38.6	342		[1962PAS/THO, 1984BOU/FRI]
	$\Delta_v H$	(328–411)	38.4	343		[1949FOR/NOR]
C₉H₁₈	[7094-27-1]	1,1,4-trimethylcyclohexane				
	$\Delta_v H$		45.6 ± 0.2	298		[1995LUK/KOZ2, 1995LUK/KOZ]
C₉H₁₈	[1795-27-3]	<i>cis</i> 1,3,5-trimethylcyclohexane				
	$\Delta_v H$	(318–410)	38.3	333	A	[1987STE/MAL]
C₉H₁₈	[3074-78-0]	2,6-dimethyl-1-heptene				
	$\Delta_v H$	(273–306)	46.3 ± 0.5	290	GS	[2000VER/WAN]
	$\Delta_v H$	(273–306)	45.9 ± 0.5	298	GS	[2000VER/WAN]
C₉H₁₈	[124-11-8]	1-nonene				
	$\Delta_{\text{fus}} H$		19.97	191.6		[1990MES/TOD]
	$\Delta_v H$	(278–318)	44.7 ± 0.2	298	GS	[2000VER/WAN]
	$\Delta_v H$		45.5	298		[1971WIL/ZWO]
	$\Delta_v H$	(339–423)	42.0	354	A,MM	[1987STE/MAL, 1950FOR/CAM]
C₉H₁₈	[6434-77-1]	<i>cis</i> 2-nonene				
	$\Delta_v H$	(379–424)	40.7	394	A	[1987STE/MAL]
C₉H₁₈	[6434-78-2]	<i>trans</i> 2-nonene				
	$\Delta_v H$	(379–422)	40.8	394	A	[1987STE/MAL]
C₉H₁₈	[20237-46-1]	<i>cis</i> 3-nonene				
	$\Delta_v H$	(376–422)	40.3	391	A	[1987STE/MAL]
C₉H₁₈	[20063-92-7]	<i>trans</i> 3-nonene				
	$\Delta_v H$	(377–421)	40.6	392	A	[1987STE/MAL]
C₉H₁₈	[10405-84-2]	<i>cis</i> 4-nonene				
	$\Delta_v H$	(376–421)	40.1	391	A	[1987STE/MAL]
C₉H₁₈	[10405-85-3]	<i>trans</i> 4-nonene				
	$\Delta_v H$	(376–420)	40.4	391	A	[1987STE/MAL]
C₉H₁₈Br₂	[62168-27-8]	1,1-dibromononane				
	$\Delta_v H$	(427–591)	59.5	442	A,EST	[1987STE/MAL, 1956MAN, 1970DYK/VAN, 1999DYK/SVO]
C₉H₁₈Cl₂	[821-88-5]	1,1-dichlorononane				
	$\Delta_v H$	(420–490)	62.3	298		[1987VAR/LOS2, 1991BAS/SVO]
	$\Delta_v H$	(398–556)	54.0	413	A,EST	[1987STE/MAL, 1956MAN, 1970DYK/VAN, 1999DYK/SVO]
C₉H₁₈Cl₂	[56375-96-3]	1,2-dichlorononane				
	$\Delta_v H$	(430–510)	52.1	443		[1999DYK/SVO]
	$\Delta_v H$	(430–510)	62.1	298		[1986VAR, 1991BAS/SVO]
C₉H₁₈F₂	[62127-42-8]	1,1-difluorononane				
	$\Delta_v H$	(347–482)	47.2	362	A,EST	[1987STE/MAL, 1956MAN, 1970DYK/VAN, 1999DYK/SVO]

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound					
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference	
C ₉ H ₁₈ F ₂	[145853-37-8]	2,2-difluorononane $\Delta_v H$	(279–313) 46.7 ± 0.2	298	GS	[1997SCH/VER]	
C ₉ H ₁₈ NO ₂	[na]	propyl 2-(N,N-dimethylamino)-2-methylpropanoate $\Delta_v H$	(282–318) 54.0 ± 0.5	298	GS	[1996VER/ZUF]	
C ₉ H ₁₈ NO ₂	[na]	ethyl 2-(N,N-diethylamino)2-propanoate $\Delta_v H$	(283–313) 54.9 ± 0.6	298	GS	[1996VER/ZUF]	
C ₉ H ₁₈ NO ₂	[2226-96-2]	2,2,6,6-tetramethyl-4-hydroxypiperidine-1-oxyl $\Delta_{\text{sub}} H$	(293–318) 101.5 ± 5.2	306	ME	[1966LEB/ROS, 1970COX/PIL]	
C ₉ H ₁₈ N ₂	[19340-91-9]	2-(diethylamino)pentanenitrile $\Delta_v H$	(283–318) 57.4 ± 0.4	298	GS	[1997WEL/VER]	
			(283–326) 58.8				
C ₉ H ₁₈ N ₂ OS	[na]	N,N-diethyl-N'-isobutanoylthiourea $\Delta_{\text{sub}} H$	363 120.8 ± 2.5	298	C	[2001RIB/RIB]	
C ₉ H ₁₈ N ₂ O ₂	[1842-72-4]	azelamide $\Delta_{\text{fus}} H$	55.0	450.4	DSC	[2006BAD/DEL]	
C ₉ H ₁₈ N ₂ O ₂ S	[39196-18-4]	3,3-dimethyl-1-(methylthio)-2-butanone O-[(methylamino)carbonyl]oxime $\Delta_{\text{fus}} H$	19.83	330.2	DSC	[1991ACR, 1990DON/DRE]	
			$\Delta_{\text{sub}} H$	(298–328) 93.5 ± 6	308	ME	[1987STE/MAL, 1976DEP]
C ₉ H ₁₈ N ₆	[645-05-6]	1,3,5-tris(dimethylamino)-s-triazine $\Delta_{\text{fus}} H$	23.01	444.4	DSC	[1991ACR]	
C ₉ H ₁₈ N ₆	[16268-64-7]	1-(ethylamino)-3,5-bis(dimethylamino)-s-triazine $\Delta_{\text{fus}} H$	16.74	333	DSC	[1991ACR]	
C ₉ H ₁₈ O	[1462-97-1]	1-butylcyclopentanol $\Delta_v H$	(359–466) 63.5	374	A	[1987STE/MAL]	
C ₉ H ₁₈ O	[815-24-7]	2,2,4,4-tetramethyl-3-pentanone $\Delta_v H$	45.5 ± 0.4	298	C	[1977PEA/FUC]	
			45.4 ± 0.1	298	C	[1970SEL2]	
			45.4 ± 0.1	298	C	[1966WAD]	
C ₉ H ₁₈ O	[108-83-8]	2,6-dimethyl-4-heptanone $\Delta_v H$	49.8 ± 0.1	308	C	[1992SVO/KUB]	
			49.3 ± 0.1	313	C	[1992SVO/KUB]	
			48.4 ± 0.1	323	C	[1992SVO/KUB]	
			47.9 ± 0.1	328	C	[1992SVO/KUB]	
			47.1 ± 0.1	338	C	[1992SVO/KUB]	
			46.6 ± 0.1	343	C	[1992SVO/KUB]	
			46.1 ± 0.1	348	C	[1992SVO/KUB]	
			45.2 ± 0.1	358	C	[1992SVO/KUB]	
			51.0	298		[1975AMB/ELL]	
			50.9 ± 0.1	298	C	[1970SEL2]	
			46.8	351	A,MM	[1987STE/MAL, 1947STR/GAB]	
C ₉ H ₁₈ O	[na]	1-(1-methylcyclohexyl)ethanol $\Delta_v H$	(358–408) 55.5	373	A	[1987STE/MAL]	
C ₉ H ₁₈ O	[124-19-6]	nonanal $\Delta_{\text{fus}} H$	29.6	253.9		[1980DYA/VAS]	
			$\Delta_v H$	(276–309) 55.3 ± 0.3	298	GS	[2003VER/KRA2]
			$\Delta_v H$	(323–343) 55.2	298	CGC	[1996KOU/HOS, 2000OVA/KOU]
			$\Delta_v H$	(313–353) 58.9	298	CGC	[1995CHI/HOS]
			$\Delta_v H$	(306–458) 51.1	321	A	[1987STE/MAL]

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$		56.3 ± 0.2	298		[1981DYA/KOR]
C ₉ H ₁₈ O	[925-78-0]	2-nonanone				
	$\Delta_v H$	(285–454)	55.6	300		[1999DIA/GUE]
	$\Delta_v H$	(335–468)	52.6	350	A	[1987STE/MAL]
	$\Delta_v H$		56.6 ± 0.6	298	GCC	[1979SAL/PEA]
	$\Delta_v H$		56.4 ± 0.1	298	C	[1977SEL]
	$\Delta_v H$	(342–545)	56.4	298		[1975AMB/ELL]
	$\Delta_v H$	(335–437)	52.7	348		[1966MEY/WAG]
C ₉ H ₁₈ O	[502-56-7]	5-nonanone				
	$\Delta_{\text{fus}} H$		24.94	269.3		
	$\Delta_{\text{fus}} H$		11.27	451.8		[1991ACR]
	$\Delta_v H$	(443–486)	44.7	458	A	[1987STE/MAL]
	$\Delta_v H$		54.9 ± 0.4	298	GCC	[1979SAL/PEA]
	$\Delta_v H$	(357–468)	49.7	372	A	[1987STE/MAL, 1975AMB/ELL]
	$\Delta_v H$		55.0	298		[1975AMB/ELL]
	$\Delta_v H$		53.3 ± 0.1	298	C	[1970HAR/HEA]
	$\Delta_v H$	(283–323)	40.2	298	A	[1987STE/MAL, 1937RIN/SAY]
C ₉ H ₁₈ O	[116-02-9]	3,3,5-trimethylcyclohexanol				
	$\Delta_v H$	(343–473)	61.8	358	A	[1987STE/MAL]
C ₉ H ₁₈ O	[53965-16-5]	2,2,5-trimethyl-4-hexene-1-ol				
	$\Delta_v H$	(323–373)	61.5	338	A	[1987STE/MAL]
C ₉ H ₁₈ O ₂	[858858-08-9]	2-butoxy-3-pentanone				
	$\Delta_v H$	(333–398)	39.8	348	A	[1987STE/MAL, 1933HEN/MUR]
C ₉ H ₁₈ O ₂	[22432-66-2]	2-butyl-1,3-dioxepane				
	$\Delta_v H$	(325–358)	57.4	340	A	[1987STE/MAL]
C ₉ H ₁₈ O ₂	[3274-29-1]	2-ethylheptanoic acid				
	$\Delta_v H$	(386–475)	63.4	401	A,EB	[1987STE/MAL, 1960TRE/MIL]
C ₉ H ₁₈ O ₂	[4352-95-8]	2-methyl-2-pentyl-1,3-dioxolane				
	$\Delta_v H$	(278–318)	54.0 ± 0.3	298	GS	[1998VER/PEN, 2002VER]
C ₉ H ₁₈ O ₂	[4421-10-7]	2,2-diisopropyl-1,3-dioxolane				
	$\Delta_v H$	(278–318)	49.9 ± 0.3	293	GS	[1998VER/PEN, 2002VER]
C ₉ H ₁₈ O ₂	[1708-34-5]	2-hexyl-1,3-dioxolane				
	$\Delta_v H$	(325–353)	55.0	339	A	[1987STE/MAL]
C ₉ H ₁₈ O ₂	[64198-22-7]	methyl 2,4,4-trimethylpentanoate				
	$\Delta_v H$	(278–318)	48.4 ± 0.2	298	GS	[1996VER/BEC]
C ₉ H ₁₈ O ₂	[5129-37-3]	butyl pivalate				
	$\Delta_v H$	(274–313)	49.5 ± 0.2	298	GS	[2008VER/EME]
	$\Delta_v H$	(274–313)	50.4 ± 0.3	298	GS	[1996VER/BEC]
C ₉ H ₁₈ O ₂	[589-59-3]	isobutyl isovalerate				
	$\Delta_v H$	(289–442)	47.3	304	A	[1987STE/MAL, 1947STU]
C ₉ H ₁₈ O ₂	[106-27-4]	isopentyl butyrate				
	$\Delta_v H$	(294–452)	47.4	309	A	[1987STE/MAL, 1947STU]
C ₉ H ₁₈ O ₂	[2050-01-3]	isopentyl isobutyrate				
	$\Delta_v H$	(287–442)	47.4	302	A	[1987STE/MAL, 1947STU]
C ₉ H ₁₈ O ₂	[2311-46-8]	isopropyl caproate				
	$\Delta_v H$	(307–383)	51.6	322	A	[1987STE/MAL]

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound					
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference	
C ₉ H ₁₈ O ₂	[111-11-5]	methyl octanoate (methyl caprylate)					
	$\Delta_v H$		53.3	350		[2002VAN/VAN]	
	$\Delta_v H$		52.6 ± 0.1	363		[2002VAN/VAN]	
	$\Delta_v H$		56.9 ± 0.1	298		[2002VAN/VAN]	
	$\Delta_v H$		54.7 ± 0.6	298	GC	[1987AZA]	
	$\Delta_v H$		57.3 ± 0.4	298	GCC	[1980FUC/PEA]	
	$\Delta_v H$		57.9 ± 0.4	298	C	[1977PEA/FUC]	
	$\Delta_v H$		56.4 ± 0.5	298	C	[1977MAN/SEL]	
	$\Delta_v H$	(347–470)	52.4	362	A,EST	[1987STE/MAL, 1963ROS/SCH]	
C ₉ H ₁₈ O ₂	[112-05-0]	nonanoic acid					
	$\Delta_{\text{us}} H$		5.61	268			
	$\Delta_{\text{fus}} H$		20.31	285.5		[1991ACR]	
	$\Delta_v H$	(381–528)	76.9	396	A	[1987STE/MAL]	
	$\Delta_v H$	(292–313)	85.3 ± 2.0	304	ME,TE	[1982DEK/SCH]	
	$\Delta_v H$	(293–303)	82.4 ± 0.4	298		[1968BAC/NOV]	
	$\Delta_v H$	(387–483)	64.2		EB	[1960TRE/MIL]	
	C ₉ H ₁₈ O ₂	[626-77-7]	propyl caproate				
		$\Delta_v H$	(315–394)	52.8	330	A	[1987STE/MAL]
$\Delta_v H$		(315–394)	52.1	330		[1961ROS/SUP, 1984BOU/FRI]	
C ₉ H ₁₈ O ₂	[112-06-1]	heptyl acetate					
	$\Delta_v H$	(274–306)	57.1 ± 0.2	298	GS	[2006KRA/VER]	
	$\Delta_v H$		56.9	298	GC	[1997KOU/HOS]	
C ₉ H ₁₈ O ₂	[2050-00-2]	<i>tert</i> -amyl butyrate					
	$\Delta_v H$	(274.5-309)	50.3 ± 0.2	298	GS	[2008VER/EME]	
	$\Delta_v H$	(333–378)	48.9	298	CGC	[1999VER/HEI]	
C ₉ H ₁₈ O ₂	[194784-95-5]	<i>tert</i> -amyl isobutyrate					
	$\Delta_v H$	(333–378)	47.8	298	CGC	[1999VER/HEI]	
	C ₉ H ₁₈ O ₂	[245658-26-8]	2-methyl-2-pentanol propanoate				
$\Delta_v H$		(333–378)	49.8	298	CGC	[1999VER/HEI]	
C ₉ H ₁₈ O ₂	[245658-31-5]	2,3-dimethyl-2-butanol 2-propanoate					
	$\Delta_v H$	(333–378)	49.9	298	CGC	[1999VER/HEI]	
C ₉ H ₁₈ O ₂	[15706-73-7]	butyl 2-methylbutanoate					
	$\Delta_v H$	(274–319)	54.4 ± 0.3	298	GS	[2008VER/EME]	
	$\Delta_v H$	(278–313)	50.6 ± 0.5	298	GS	[1996VER/BEC]	
C ₉ H ₁₈ O ₃	[109857-47-8]	2-butoxypropionic acid, ethyl ester					
$\Delta_v H$	(348–438)	80.3	363	A	[1987STE/MAL, 1933HEN/MUR]		
C ₉ H ₁₈ O ₃	[14144-35-5]	3-ethoxypropionic acid, butyl ester					
$\Delta_v H$	(346–479)	51.8	361	A	[1987STE/MAL]		
C ₉ H ₁₈ O ₃	[na]	3-hydroxypropionic acid, hexyl ester					
$\Delta_v H$	(408–432)	69.6	420	A	[1987STE/MAL]		
C ₉ H ₁₈ O ₃	[20279-51-0]	lactic acid, hexyl ester					
$\Delta_v H$	(307–494)	67.4	322	A	[1987STE/MAL]		
C ₉ H ₁₈ O ₃	[10500-16-0]	3-methoxypropionic acid, pentyl ester					
$\Delta_v H$	(322–485)	53.3	337	A	[1987STE/MAL]		
C ₉ H ₁₈ O ₃	[14144-41-3]	3-propoxypropionic acid, propyl ester					

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(317–484)	50.9	332	A	[1987STE/MAL]
C ₉ H ₁₈ O ₃	[542-52-9]	dibutyl carbonate				
	$\Delta_v H$	(287–329)	62.9 ± 0.4	298	GS	[2008KOZ/EME]
C ₉ H ₁₈ O ₃	[34619-03-9]	di- <i>tert</i> -butylcarbonate				
	$\Delta_{\text{sub}} H$		65.4 ± 0.2	298	C	[1985KUS]
C ₉ H ₁₈ S ₄	[25423-58-9]	1,5,9-trithiacyclododecane				
	$\Delta_{\text{fus}} H$		12	349.2		
	$\Delta_{\text{fus}} H$		19.4	373.2	DSC	[2002ROC/GRI]
C ₉ H ₁₉ Br	[693-58-3]	1-bromononane				
	$\Delta_{\text{fus}} H$		30.12	243.2		[1950CRO/SMY]
	$\Delta_v H$	(376–525)	53.1	391		[1999DYK/SVO]
	$\Delta_v H$	(391–549)	52.2	406	A,EST	[1987STE/MAL, 1961LI/ROS, 1970DYK/VAN]
C ₉ H ₁₉ Cl	[2473-01-0]	1-chlorononane				
	$\Delta_v H$	(363–509)	51.5	378		[1999DYK/SVO]
	$\Delta_v H$	(340–480)	55.9	298		[1984BOU/FRI, 1991BAS/SVO]
	$\Delta_v H$	(342–478)	53.4	357	A,DTA	[1987STE/MAL, 1969KEM/KRE]
C ₉ H ₁₉ F	[463-18-3]	1-fluorononane				
	$\Delta_v H$	(278–313)	50.8 ± 0.9	298	GS	[1994STE/CHI]
	$\Delta_v H$	(333–473)	46.8	348	A,EST	[1987STE/MAL, 1961LI/ROS, 1970DYK/VAN, 1999DYK/SVO]
C ₉ H ₁₉ I	[4282-42-2]	1-iodononane				
	$\Delta_v H$		64.5	298		[2006BOL/NER, 1961LI/ROS]
	$\Delta_v H$	(391–551)	54.6	406		[1999DYK/SVO]
	$\Delta_v H$	(408–577)	53.5	423	A	[1987STE/MAL, 1970DYK/VAN]
	$\Delta_v H$	(343–493)	64.3	358		[1947STU]
C ₉ H ₁₉ N	[768-66-1]	2,2,6,6-tetramethylpiperidine				
	$\Delta_v H$	(288–313)	44.5 ± 0.5	300		[1997VER]
C ₉ H ₁₉ N	[4945-48-6]	N-butylpiperidine				
	$\Delta_v H$	(275–313)	49.2 ± 0.2	294	GS	[1998VER6]
	$\Delta_v H$	(275–313)	48.9 ± 0.2	298	GS	[1998VER6]
C ₉ H ₁₉ N	[13173-21-2]	N,N-diethyl-4-pentenylamine				
$\Delta_v H$	(338–430)	41.5	353	A	[1987STE/MAL]	
C ₉ H ₁₉ NO	[103-00-4]	1-(cyclohexylamino)-2-propanol				
$\Delta_v H$	(423–512)	56.6	438	A	[1987STE/MAL, 1984BOU/FRI, 1959MCD/SHR]	
C ₉ H ₁₉ NO	[1120-07-6]	nonanamide				
	$\Delta_{\text{sub}} H$	(353–370)	114.6 ± 3.3		ME	[1959DAV/JON2, 1987STE/MAL]
C ₉ H ₁₉ NO ₂	[35601-84-4]	heptylcarbamic acid, methyl ester				
$\Delta_v H$	(368–408)	109.8	383	A	[1987STE/MAL]	
C ₉ H ₁₉ NO ₂	[3637-10-3]	2,2,6,6-tetramethyl-1,4-dihydroxypiperidine				
$\Delta_{\text{sub}} H$	(318–348)	100.4 ± 0.6	328	ME	[1966LEB/ROS, 1970COX/PIL]	
C ₉ H ₂₀	[111-84-2]	nonane				
	$\Delta_{\text{us}} H$		6.2	218.2		
	$\Delta_{\text{fus}} H$		15.0	219.5	DSC	[2004MON/RAJ]
	$\Delta_{\text{us}} H$		6.28	217.2		
	$\Delta_{\text{fus}} H$		15.48	219.7		[1991ACR, 1996DOM/HEA]

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{sub}}H$		74.6	219	B	[1963BON]
	Δ_vH		46.5 ± 0.2	298	C	[2007PAS/KUZ]
	Δ_vH		46.7	299	C	[1996VIT/CHA]
	Δ_vH		46	314	C	[1996VIT/CHA]
	Δ_vH		46.6 ± 0.2	298	C	[1996VAR/PAS]
	Δ_vH		46.6	298		[1994RUZ/MAJ]
	Δ_vH	(322–413)	43.9	337		[1986PAU/KRU]
	Δ_vH		44.3	328	C	[1984MAJ/SVO3]
	Δ_vH		43.2	343	C	[1984MAJ/SVO3]
	Δ_vH		42.1	358	C	[1984MAJ/SVO3]
	Δ_vH		46.4	298		[1971WIL/ZWO]
	Δ_vH	(219–308)	48.3	234	A	[1987STE/MAL, 1973CAR/KOB]
	Δ_vH		46.4	298	C	[1947OSB/GIN]
	Δ_vH	(344–426)	42.7	359	A,MM	[1987STE/MAL, 1945WIL/TAY, 1949FOR/NOR]
C₉H₂₀	[3221-61-2]	2-methyloctane				
	$\Delta_{\text{fus}}H$		17.99	192.8		[1998HEL/OWE]
	Δ_vH	(305–417)	43.2	320	A	[1987STE/MAL]
	Δ_vH		44.9	298		[1971WIL/ZWO, 1961LAB/GRE]
C₉H₂₀	[2216-33-3]	3-methyloctane				
	$\Delta_{\text{fus}}H$		16.99	165.6		[1998HEL/OWE]
	Δ_vH		44.9	298		[1971WIL/ZWO, 1961LAB/GRE]
C₉H₂₀	[2216-34-4]	4-methyloctane				
	Δ_vH		44.5	298		[1971WIL/ZWO, 1961LAB/GRE]
C₉H₂₀	[15869-80-4]	3-ethylheptane				
	Δ_vH		44.5	298		[1971WIL/ZWO, 1961LAB/GRE]
C₉H₂₀	[2216-32-2]	4-ethylheptane				
	Δ_vH		44.1	298		[1961LAB/GRE]
C₉H₂₀	[1071-26-7]	2,2-dimethylheptane				
	Δ_vH		42.3	298		[1971WIL/ZWO, 1961LAB/GRE]
C₉H₂₀	[3074-71-3]	2,3-dimethylheptane				
	Δ_vH		43.6	298		[1971WIL/ZWO, 1961LAB/GRE]
C₉H₂₀	[2213-23-2]	2,4-dimethylheptane				
	Δ_vH		42.9	298		[1971WIL/ZWO, 1961LAB/GRE]
C₉H₂₀	[2216-30-0]	2,5-dimethylheptane				
	Δ_vH		43.3	298		[1971WIL/ZWO, 1961LAB/GRE]
C₉H₂₀	[1072-05-5]	2,6-dimethylheptane				
	Δ_vH		43.3	298		[1971WIL/ZWO, 1961LAB/GRE]
C₉H₂₀	[4032-86-4]	3,3-dimethylheptane				
	Δ_vH		42.6	298		[1971WIL/ZWO, 1961LAB/GRE]
C₉H₂₀	[922-28-1]	3,4-dimethylheptane				
	Δ_vH		43.6	298		[1971WIL/ZWO, 1961LAB/GRE]
C₉H₂₀	[926-82-9]	3,5-dimethylheptane				
	Δ_vH		43.3	298		[1971WIL/ZWO]
C₉H₂₀	[1068-19-5]	4,4-dimethylheptane				
	Δ_vH		42.2	298		[1971WIL/ZWO, 1961LAB/GRE]

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₉ H ₂₀	[1067-20-5]	3,3-diethylpentane				
	$\Delta_{\text{us}}H$		0.48	208.3		
	$\Delta_{\text{us}}H$		0.81	210.4		
	$\Delta_{\text{fus}}H$		10.09	240.1		[1996DOM/HEA]
	Δ_vH		42.6 ± 0.3	298	GCC	[1979FUC/PEA]
	Δ_vH	(335–426)	39.8	350	A	[1971WIL/ZWO, 1961LAB/GRE] [1987STE/MAL, 1949FOR/NOR]
C ₉ H ₂₀	[16789-46-1]	2-methyl-3-ethylhexane				
	Δ_vH		43.2	298		[1961LAB/GRE]
C ₉ H ₂₀	[3074-75-7]	2-methyl-4-ethylhexane				
	Δ_vH		42.9	298		[1961LAB/GRE]
C ₉ H ₂₀	[3074-76-8]	3-methyl-3-ethylhexane				
	Δ_vH		42.9	298		[1961LAB/GRE]
C ₉ H ₂₀	[na]	3-methyl-4-ethylhexane				
	Δ_vH		43.6	298		[1961LAB/GRE]
C ₉ H ₂₀	[16747-25-4]	2,2,3-trimethylhexane				
	Δ_vH	(238–303)	42.2	288	IP	[1974OSB/DOU]
	Δ_vH		41.7	298		[1971WIL/ZWO, 1961LAB/GRE]
C ₉ H ₂₀	[16747-26-5]	2,2,4-trimethylhexane				
	Δ_vH	(288–410)	39.5	303	A	[1987STE/MAL]
	Δ_vH	(238–393)	41.0	278	A	[1987STE/MAL]
	Δ_vH	(238–303)	40.5	288	IP	[1974OSB/DOU]
	Δ_vH		40.7	298		[1971WIL/ZWO]
C ₉ H ₂₀	[3522-94-9]	2,2,5-trimethylhexane				
	Δ_vH	(288–399)	40.1	303	A	[1987STE/MAL]
	Δ_vH	(238–303)	41.1	288	A,IP	[1987STE/MAL, 1974OSB/DOU]
	Δ_vH		40.2	298		[1971WIL/ZWO]
	Δ_vH	(319–398)	38.5	334		[1949FOR/NOR]
	Δ_vH		40.2	298	C	[1947OSB/GIN]
C ₉ H ₂₀	[16747-28-7]	2,3,3-trimethylhexane				
	Δ_vH	(238–303)	44.2	253	A	[1987STE/MAL]
	Δ_vH	(288–422)	39.4	303	A	[1987STE/MAL]
	Δ_vH		42.1	298		[1971WIL/ZWO, 1961LAB/GRE]
C ₉ H ₂₀	[921-47-1]	2,3,4-trimethylhexane				
	Δ_vH		42.7	298		[1971WIL/ZWO, 1961LAB/GRE]
C ₉ H ₂₀	[1069-53-0]	2,3,5-trimethylhexane				
	Δ_vH		41.4	298		[1971WIL/ZWO]
	Δ_vH		41.4	298	C	[1947OSB/GIN]
C ₉ H ₂₀	[16747-30-0]	2,4,4-trimethylhexane				
	Δ_vH		41.1	298		[1971WIL/ZWO, 1961LAB/GRE]
	Δ_vH	(323–406)	38.5	338	A	[1987STE/MAL, 1949FOR/NOR]
C ₉ H ₂₀	[16747-31-2]	3,3,4-trimethylhexane				
	Δ_vH		42.2	298		[1971WIL/ZWO, 1961LAB/GRE]
C ₉ H ₂₀	[16747-32-3]	2,2-dimethyl-3-ethylpentane				
	Δ_vH		41.7	298		[1971WIL/ZWO, 1961LAB/GRE]
C ₉ H ₂₀	[16747-33-4]	2,3-dimethyl-3-ethylpentane				
	Δ_vH		42.7	298		[1971WIL/ZWO, 1961LAB/GRE]
C ₉ H ₂₀	[1068-87-7]	2,4-dimethyl-3-ethylpentane				

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$		42.3	298		[1971WIL/ZWO, 1961LAB/GRE]
C ₉ H ₂₀	[7154-79-2]	2,2,3,3-tetramethylpentane				
	$\Delta_{\text{trs}}H$		7.33	174.5		
	$\Delta_{\text{fus}}H$		2.33	263.4		[1996DOM/HEA]
C ₉ H ₂₀	$\Delta_v H$		41.2	298		[1971WIL/ZWO, 1961LAB/GRE]
	$\Delta_v H$	(328–415)	39.2	343	A	[1987STE/MAL, 1949FOR/NOR]
C ₉ H ₂₀	[1186-53-4]	2,2,3,4-tetramethylpentane				
	$\Delta_v H$		40.8	298		[1971WIL/ZWO, 1961LAB/GRE]
	$\Delta_v H$	(325–413)	38.4	340	A	[1987STE/MAL, 1949FOR/NOR]
C ₉ H ₂₀	[1070-87-7]	2,2,4,4-tetramethylpentane				
	$\Delta_{\text{fus}}H$		9.75	206.7		[1996DOM/HEA]
	$\Delta_v H$		38.5 ± 0.1	298	C	[1982FUC/PEA]
	$\Delta_v H$		38.5 ± 0.3	298	GCC	[1979FUC/PEA]
	$\Delta_v H$		38.2	298		[1971WIL/ZWO, 1961LAB/GRE]
	$\Delta_v H$	(313–397)	37.2	328	A	[1987STE/MAL, 1949FOR/NOR]
	$\Delta_v H$	(331–375)	36.5	346	EB	[1941SMI, 1984BOU/FRI]
	$\Delta_v H$	(375–422)	34.8	390		[1941SMI]
C ₉ H ₂₀	[16747-38-9]	2,3,3,4-tetramethylpentane				
	$\Delta_v H$		41.8	298		[1971WIL/ZWO, 1961LAB/GRE]
	$\Delta_v H$	(331–416)	39.3	346	A	[1987STE/MAL, 1949FOR/NOR]
C ₉ H ₂₀ ClF ₃ N ₂ S	[63265-72-5]	chlorobis(N-ethylethanaminato)(trifluoromethyl) sulfur				
	$\Delta_v H$		39.8	479	I	[1977KIT/SHR2]
C ₉ H ₂₀ ClF ₃ N ₂ OS	[63265-74-7]	chlorobis(N-ethylethanaminato)oxo(trifluoromethyl) sulfur				
	$\Delta_v H$		44.4	486	I	[1977KIT/SHR2]
C ₉ H ₂₀ N ₂ O	[1792-17-2]	1,3-dibutylurea				
	$\Delta_{\text{trs}}H$		11.1	311.5		
	$\Delta_{\text{fus}}H$		14.87	349.6		[1996DOM/HEA]
	$\Delta_{\text{sub}}H$	(323–372)	91.9 ± 0.9	348	ME	[2003ZAI/KAB]
	$\Delta_{\text{sub}}H$	(323–372)	91.9 ± 0.9	350	ME	[2003ZAI/KAB]
	$\Delta_{\text{sub}}H$		90.0 ± 1.0	350	C	[2003ZAI/KAB]
	$\Delta_v H$	(379–413)	101.1 ± 1.6	396		[1990PIA/FER]
C ₉ H ₂₀ N ₂ O	[2158-10-3]	1-octyl urea				
	$\Delta_{\text{trs}}H$		11.8	353.2		
	$\Delta_{\text{fus}}H$		24.4	374.6	DSC	[2005HAS/TAJ]
	$\Delta_{\text{trs}}H$		11.5	350.2		
	$\Delta_{\text{fus}}H$		24.6	372.2		[1999WEL/DRU]
C ₉ H ₂₀ N ₂ O	[1187-03-7]	1,1,3,3-tetraethylurea				
	$\Delta_{\text{fus}}H$		20.55	253		[1990KOZ/SIM2, 1996DOM/HEA, 1995KAB/KOZ2]
C ₉ H ₂₀ N ₂ S	[109-46-6]	1,3-butylthiourea				
	$\Delta_{\text{fus}}H$		28.34	338	DSC	[2000DEL/JOZ]
	$\Delta_{\text{sub}}H$		141.0 ± 2	298	B	[2000DEL/JOZ]
	$\Delta_{\text{sub}}H$		137 ± 3.0	298	C	[1994TER/PIA]
	$\Delta_v H$	(368–403)	105 ± 2.0	386	ME,TE	[1994TER/PIA]
C ₉ H ₂₀ N ₂ S ₂	[1518-58-7]	diethylammonium diethyldithiocarbamate				
	$\Delta_{\text{sub}}H$		209.9 ± 3.0		C	[1988RIB/REI]
	$\Delta_{\text{sub}}H$		111.8 ± 3.0			[1979CAV/HIL]

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
Note: Authors of [1988RIB/REI] state compound decomposes on sublimation						
C ₉ H ₂₀ O	[143-08-8]	1-nonanol				
	$\Delta_v H$	(284–353)	74.7 ± 0.3	298	GS	[2005ROG/PIS]
	$\Delta_v H$		72.2	298	CGC	[2000OVA/KOU]
	$\Delta_v H$	(373–423)	76.7	298	CGC	[1995CHI/HOS]
	$\Delta_v H$	(273–323)	77.4	298		[1992NGU/KAS]
	$\Delta_v H$	(368–500)	65.0	383	A	[1987STE/MAL]
	$\Delta_v H$	(381–495)	62.9	396	A	[1987STE/MAL]
	$\Delta_v H$		76.9 ± 0.8	298	C	[1977MAN/SEL]
	$\Delta_v H$	(425–494)	59.7	440	EB	[1976HON/SIN]
C ₉ H ₂₀ O	[628-99-9]	2-nonanol				
	$\Delta_v H$	(286–324)	72.9 ± 0.6	298	GS	[2007VER/SCH]
C ₉ H ₂₀ O						
	$\Delta_v H$	(253–353)	79.6	268		[1999NGU/BER]
	$\Delta_v H$	(364–471)	55.5	379		[1973WIL/ZWO]
C ₉ H ₂₀ O	[624-51-1]	3-nonanol				
	$\Delta_v H$	(263–363)	70.9 ± 0.3	298	GS	[2007VER/SCH]
	$\Delta_v H$	(263–363)	75.5	278		[1999NGU/BER]
C ₉ H ₂₀ O						
	$\Delta_v H$	(366–468)	57.1	381		[1973WIL/ZWO]
C ₉ H ₂₀ O	[5932-79-6]	4-nonanol				
	$\Delta_v H$	(285–324)	71.5 ± 0.3	298	GS	[2007VER/SCH]
C ₉ H ₂₀ O	[623-93-8]	5-nonanol				
	$\Delta_v H$	(289–334)	71.4 ± 0.4	298	GS	[2007VER/SCH]
C ₉ H ₂₀ O	[628-44-4]	2-methyl-2-octanol				
	$\Delta_v H$	(338–451)	64.6	353		[1973WIL/ZWO]
C ₉ H ₂₀ O	[26533-34-6]	2-methyl-3-octanol				
	$\Delta_v H$	(388–453)	49.5	403		[1973WIL/ZWO]
C ₉ H ₂₀ O	[5340-36-3]	3-methyl-3-octanol				
	$\Delta_v H$	(353–388)	53.2	368		[1973WIL/ZWO]
C ₉ H ₂₀ O	[na]	2,2-dimethyl-4-heptanol				
	$\Delta_v H$	(320–445)	50.2	335		[1973WIL/ZWO]
C ₉ H ₂₀ O	[108-82-7]	2,6-dimethyl-4-heptanol				
	$\Delta_v H$	(278–321)	65.2 ± 0.3	298	GS	[2005ROG/PIS]
	$\Delta_v H$	(363–453)	54.5	378		[1973WIL/ZWO]
	$\Delta_v H$	(374–452)	52.8	389	A,MM	[1987STE/MAL, 1947STR/GAB]
C ₉ H ₂₀ O	[5340-41-0]	2,2,3-trimethyl-3-hexanol				
	$\Delta_v H$	(343–441)	55.1	358		[1973WIL/ZWO]
C ₉ H ₂₀ O	[3452-97-9]	3,5,5-trimethyl-1-hexanol				
	$\Delta_v H$	(288–324)	67.9 ± 0.4	298	GS	[2005ROG/PIS]
C ₉ H ₂₀ O	[3970-59-0]	2,4-dimethyl-3-ethyl-3-pentanol				
	$\Delta_v H$	(369–451)	50.0	384		[1973WIL/ZWO]
C ₉ H ₂₀ O	[29772-39-2]	2,2,3,4-tetramethyl-3-pentanol				
	$\Delta_v H$	(329–448)	60.8	344		[1973WIL/ZWO]
C ₉ H ₂₀ O	[14609-79-1]	di- <i>tert</i> -butylmethanol				
	$\Delta_v H$		62.7 ± 0.9	298		[1998VER3]
C ₉ H ₂₀ O	[14609-79-1]	2,2,4,4-tetramethylpentan-3-ol				
	$\Delta_{\text{vs}} H$		1.9	263		

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
		$\Delta_{\text{fus}}H$	7.3	322		[1996DOM/HEA]
C ₉ H ₂₀ O	[3249-47-6]	butyl 1,1-dimethylpropyl ether Δ_vH	(278–308) 46.1 ± 0.3	298	GS	[1996VER/BEC]
C ₉ H ₂₀ O	[62108-41-2]	2-methoxy-2,4,4-trimethylpentane Δ_vH	(381–418) 38.5	396		[2001UUS/POK]
C ₉ H ₂₀ O	[62108-41-2]	methyl <i>tert</i> -octyl ether Δ_vH	45.3	298		[2002VER, 2003VER/KRA]
		Δ_vH	45.33	298	EB	[2001UUS/POK, 2003VER/KRA]
C ₉ H ₂₀ O	[10100-95-5]	pentyl <i>tert</i> -butyl ether Δ_vH	48.3	298		[2002VER, 2003VER/KRA]
		Δ_vH	(319–365) 43.7	334	EB	[1990ROZ/BAR]
		Δ_vH	(319–365) 46.9 ± 1.0	298	EB	[1990ROZ/BAR]
C ₉ H ₂₀ O	[10100-95-5]	isobutyl <i>tert</i> -amyl ether Δ_vH	46.3	298		[2002VER, 2003VER/KRA]
C ₉ H ₂₀ O	[na]	sec-butyl <i>tert</i> -amyl ether Δ_vH	46.8	298		[2002VER, 2003VER/KRA]
C ₉ H ₂₀ O	[3249-47-6]	butyl <i>tert</i> -amyl ether Δ_vH	48.3	298		[2002VER, 2003VER/KRA]
C ₉ H ₂₀ O ₂	[22419-28-9]	2,6,6-trimethyl-5-oxa-2-heptanol Δ_vH	(329–454) 53.3	344		[1968KAC/NEM, 1984BOU/FRI]
C ₉ H ₂₀ O ₂	[2568-90-3]	dibutyoxymethane Δ_vH	(366–452) 47.9	381	EB	[2000PAL/SZA]
		Δ_vH	(363–452) 48.1	298	EB	[2000PAL/SZA]
C ₉ H ₂₀ O ₂	[18854-58-5]	1-butyoxy-2-propoxyethane Δ_vH	54.7 ± 0.1	298	C	[1970KUS/WAD]
C ₉ H ₂₀ O ₂	[115-84-4]	2-butyl-2-ethyl-1,3-propanediol $\Delta_{\text{fus}}H$	20.8	317.3		[2002STE/CHI]
		Δ_vH	(424–523) 74.3 ± 0.3	420	EB	[2002STE/CHI]
		Δ_vH	(424–523) 67.2 ± 0.3	460	EB	[2002STE/CHI]
		Δ_vH	(424–523) 61.4 ± 0.6	500	EB	[2002STE/CHI]
C ₉ H ₂₀ O ₂	[22419-28-9]	4- <i>tert</i> -butoxy-2-methyl-2-butanol Δ_vH	(367–483) 61.5	382	A	[1987STE/MAL]
C ₉ H ₂₀ O ₂	[3937-56-2]	1,9-nonanediol $\Delta_{\text{fus}}H$	36.4	319.6		[1991ACR]
		$\Delta_{\text{sub}}H$	148.7			[1990KNA/SAB]
		Δ_vH	104.4	360		[1994PIA/FER, 2006UMN/KWE]
		Δ_vH	111.4 ± 7.0	298		[1994PIA/FER, 2006UMN/KWE]
		Δ_vH	110	323		[1990KNA/SAB, 2006UMN/KWE]
		Δ_vH	112.8 ± 2.1	298		[1990KNA/SAB, 2006UMN/KWE]
C ₉ H ₂₀ O ₂	[3089-24-5]	2,2,4-trimethyl-1,6-hexanediol Δ_vH	(419–541) 68.0	434	A	[1987STE/MAL]
C ₉ H ₂₀ O ₂ S	[54581-77-0]	3-(hexylthio)-1,2-propanediol $\Delta_{\text{fus}}H$	48.5	290.8	DSC	[1993ACR, 1990VAN/KEL]
C ₉ H ₂₀ O ₂ S ₂	[na]	<i>meso</i> 1,3-bis(propylsufinyl)propane $\Delta_{\text{fus}}H$	40.58	411.3		[2001CAL/MEL]

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₉ H ₂₀ O ₂ S ₂	[na] $\Delta_{\text{fus}}H$	racemic 1,3- <i>bis</i> (propylsufinyl)propane	35.15	387.8		[2001CAL/MEL]
C ₉ H ₂₀ O ₃	[na] Δ_vH	dipropylene glycol isopropyl ether (319–479)	55.0	334	A	[1987STE/MAL, 1947STU]
C ₉ H ₂₀ O ₃	[10305-38-1] $\Delta_{\text{fus}}H$	3-(hexyloxy)-1,2-propanediol	10.2	272.9	DSC	[1990VAN/KEL]
C ₉ H ₂₀ O ₄	[na] Δ_vH	tripropylene glycol (369–541)	63.3	384	A	[1987STE/MAL]
C ₉ H ₂₀ O ₄	[4161-32-4] $\Delta_{\text{fus}}H$	3,3'-[1,3-propanediyl bis (oxy)] <i>bis</i> -1-propanol	21.15	263.1	DSC	[1991BED/BOO]
C ₉ H ₂₀ O ₄	[78-09-1] Δ_vH	tetraethoxymethane	52.9 ± 0.2	298	C	[1985MAR/MAN]
C ₉ H ₂₀ S	[1455-21-6] $\Delta_{\text{fus}}H$	1-nonanethiol	33.5	267.7		[1985DEA]
	Δ_vH	(390–494)	52.6	405	A	[1987STE/MAL, 1999DYK/SVO, 1932ELL/REI]
C ₉ H ₂₀ S	[13281-11-3] Δ_vH	2-nonanethiol (379–482)	50.3	394		[1999DYK/SVO, 1932ELL/REI]
C ₉ H ₂₀ S ₂	[3489-28-9] Δ_vH	1,9-nonanedithiol (418–557)	63.6	433	A	[1987STE/MAL, 1999DYK/SVO, 1943HAL/REI]
C ₉ H ₂₁ N	[2439-54-5] Δ_vH	N-methyl octylamine (365–508)	49.2	380	A	[1987STE/MAL]
C ₉ H ₂₁ N	[112-20-9] Δ_vH	nonylamine (377–478)	50.7	392	A	[1987STE/MAL]
C ₉ H ₂₁ N	[102-69-2] Δ_vH	tripropylamine (341–475)	45.6	356	A	[1987STE/MAL]
	Δ_vH		46.2 ± 0.1	298	C	[1969WAD]
C ₉ H ₂₁ NO ₃	[122-20-3] Δ_vH	triisopropanolamine (428–573)	73.7	443	A	[1987STE/MAL]
C ₉ H ₂₁ O ₄ P	[513-08-6] Δ_vH	tripropylphosphate (394–525)	56.7	409	A	[1987STE/MAL]
C ₉ H ₂₁ P	[2234-97-1] Δ_vH	tripropylphosphine (324–368)	39.4 ± 0.2	346		[2001BAE]
C ₉ H ₂₂ ClN ₂ PS	[58023-20-4] Δ_vH	P-(chloromethyl)-N,N'- <i>bis</i> (1-methylpropyl)phosphorothioic diamide (333–368)	66.8	348	A	[1987STE/MAL, 1999DYK/SVO]
C ₉ H ₂₂ N ₂	[646-24-2] $\Delta_{\text{us}}H$ $\Delta_{\text{fus}}H$	nonane-1,9-diamine	7.77 36.24	301.7 308.1	DSC	[2002DAL/DEL]
C ₁₀ Cl ₈	[2234-13-1] Δ_vH	octachloronaphthalene (323–423)	96.1	373	GC	[1999LEI/WAN]
C ₁₀ Cl ₁₂	[2385-85-5] Δ_vH	mirex (343–453)	90.3	398	GC	[1990HIN/BID2]
C ₁₀ F ₈	[313-72-4] $\Delta_{\text{us}}H$	octafluoronaphthalene	2.12	283.6		

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound						
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference		
			$\Delta_{\text{fus}}H$	17.55	358.8		[1999MIC/NEG]	
		(293–323)	$\Delta_{\text{sub}}H$	79.4 ± 2.5	308	A	[1987STE/MAL, 1974RAD/KAT]	
C ₁₀ F ₁₆	[54939-04-7]	perfluorobicyclo[4.4.0]dec-1(6)-ene						
			$\Delta_{\text{us}}H$	0.75	200			
			$\Delta_{\text{us}}H$	1.12	234			
			$\Delta_{\text{fus}}H$	10.47	263		[1996DOM/HEA]	
			Δ_vH	45.3 ± 0.1	298	C	[1996VAR/DRU]	
C ₁₀ F ₁₈	[60433-11-6]	perfluoro(<i>cis</i> -decahydronaphthalene)						
			$\Delta_{\text{us}}H$	4.24	232.5			
			$\Delta_{\text{fus}}H$	10.3	266.7		[1996DOM/HEA]	
		(313–415)	Δ_vH	43.9	328		[1999DYK/SVO]	
			Δ_vH	46.2 ± 0.1	298	C	[1996VAR/DRU]	
			Δ_vH	46.7 ± 0.6	298	EB	[1981VAR/BUL]	
			Δ_vH	46.2 ± 0.1	298	C	[1981VAR/BUL]	
C ₁₀ F ₁₈	[60433-12-7]	perfluoro(<i>trans</i> -decahydronaphthalene)						
			$\Delta_{\text{fus}}H$	17.96	294.6		[1996DOM/HEA]	
		(315–417)	Δ_vH	43.3	330		[1999DYK/SVO]	
			Δ_vH	45.4 ± 0.1	298	C	[1996VAR/DRU]	
			Δ_vH	45.9 ± 0.6	298	EB	[1981VAR/BUL]	
			Δ_vH	45.4 ± 0.1	298	C	[1981VAR/BUL]	
C ₁₀ F ₁₈	[306-94-5]	perfluorodecalin						
		(288–333)	Δ_vH	41.5 ± 0.5	298		[2005DIA/GON]	
C ₁₀ F ₂₀	[35328-43-9]	perfluoro-1-decene						
		(315–399)	Δ_vH	42.3	330		[1999DYK/SVO]	
			Δ_vH	45.2 ± 0.6	298	EB	[1981VAR/BUL]	
C ₁₀ F ₂₀	[116667-53-9]	perfluoro(1-methyl-4-isopropyl)cyclohexane						
		(339–418)	Δ_vH	42.7	354		[1999DYK/SVO]	
			Δ_vH	46.7 ± 0.5	298	EB	[1981VAR/BUL]	
C ₁₀ F ₂₀	[na]	perfluoro(isobutyl)cyclohexane						
		(327–415)	Δ_vH	43.4	342		[1999DYK/SVO]	
			Δ_vH	46.3 ± 0.6	298	EB	[1981VAR/BUL]	
			Δ_vH	46.7 ± 0.1	298	C	[1981VAR/BUL]	
C ₁₀ F ₂₀ N ₂ S	[77984-27-1]	2,2,3,3,4,4,5,5-octafluoro-1,1,2,3,4,5-hexahydro-1-[[2,2,2-trifluoro-1-(trifluoromethyl)-1-[[2,2,2-trifluoro-1-(trifluoromethyl)ethylidene]amino]ethyl]imino]thiophene						
			Δ_vH	29.3	389		[1981ABE/SHR]	
C ₁₀ F ₂₂	[307-45-9]	perfluorodecane						
		(404–543)	Δ_vH	34	420	A	[1987STE/MAL, 1967ERM/SKR, 1999DYK/SVO]	
C ₁₀ F ₂₂ O	[464-36-8]	<i>bis</i> (undecafluoropentyl)ether						
		(337–411)	Δ_vH	49.9 ± 1.5	298	EB	[1989VAR/PAS]	
			Δ_vH	49.5 ± 0.1	298	C	[1989VAR/PAS]	
		(288–313)	Δ_vH	51.5	300	A	[1987STE/MAL, 1999DYK/SVO]	
			Δ_vH	47.3 ± 0.8	298	EB	[1976KOL/SLA]	
C ₁₀ F ₂₂ O ₈	[927699-29-4]	perfluoro-2,4,6,8,11,13,15,17-octaoxy-n-octadecane						
		(363–437)	Δ_vH	61.9 ± 1.2	298	EB	[2006DRU/KRO]	
C ₁₀ HCl ₅ F ₁₄ O ₂	[335-74-0]	2,2,3,4,4,5,6,6,7,8,8,9,10,10-tetradecafluoro-3,5,7,9,10-pentachlorodecanoic acid						
		(373–578)	Δ_vH	80.6	388	A	[1987STE/MAL, 1957BAR/SEF, 1999DYK/SVO]	

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₀ HCl ₇	[58863-14-2]	1,2,3,4,5,6,7-heptachloronaphthalene				
	$\Delta_{\text{fus}}H$		8.54	454.5		
	$\Delta_{\text{fus}}H$		8.75	456	DSC	[2006LAH/PAA]
	Δ_vH	(323–423)	90.6	373	GC	[1999LEI/WAN]
C ₁₀ H ₂ Cl ₆	[90948-28-0]	1,2,4,5,6,8-hexachloronaphthalene				
	$\Delta_{\text{fus}}H$		28.89	449.4	DSC	[2006LAH/PAA]
	Δ_vH	(323–423)	85.3	373	GC	[1999LEI/WAN]
C ₁₀ H ₂ Cl ₆	[103426-94-4]	1,2,3,5,7,8-hexachloronaphthalene				
	Δ_vH	(323–423)	85.0	373	GC	[1999LEI/WAN]
C ₁₀ H ₂ Cl ₆	[103426-97-7]	1,2,3,5,6,7-hexachloronaphthalene				
	Δ_vH	(323–423)	84.5	373	GC	[1999LEI/WAN]
C ₁₀ H ₂ Cl ₆	[103426-96-6]	1,2,3,4,6,7-hexachloronaphthalene				
	Δ_vH	(323–423)	84.5	373	GC	[1999LEI/WAN]
C ₁₀ H ₂ O ₆	[89-32-7]	1,2,4,5-benzenetetracarboxylic dianhydride (pyromellitic acid dianhydride)				
	$\Delta_{\text{fus}}H$		15.82	557.2		[1996DOM/HEA]
	$\Delta_{\text{trs}}H$		3.38	505.9		
	$\Delta_{\text{fus}}H$		32.39	558.9	DSC	[1984WEI/LEF]
	$\Delta_{\text{sub}}H$		122.3 ± 2.4	298	C	[2007MAT/MIR2]
	$\Delta_{\text{sub}}H$		100.4	559		[1975BAG/AND]
	$\Delta_{\text{sub}}H$		88.4			[1967MUL/MUK]
	Δ_vH	(641–665)	79.6	576	A	[1987STE/MAL]
C ₁₀ H ₃ Cl ₅	[150224-24-1]	1,2,3,5,8-pentachloronaphthalene				
	$\Delta_{\text{trs}}H$		14.39	387.8		
	$\Delta_{\text{trs}}H$		0.92	443.4		
	$\Delta_{\text{fus}}H$		8.44	453.3	DSC	[2006LAH/PAA]
	Δ_vH	(323–423)	80.5	373	GC	[1999LEI/WAN]
C ₁₀ H ₃ Cl ₅	[53555-65-0]	1,2,3,5,7-pentachloronaphthalene				
	$\Delta_{\text{fus}}H$		27.14	444.9	DSC	[2006LAH/PAA]
	Δ_vH	(323–423)	78.2	373	GC	[1999LEI/WAN]
C ₁₀ H ₃ Cl ₅	[67922-26-3]	1,2,3,4,6-pentachloronaphthalene				
	$\Delta_{\text{trs}}H$		15.64	404.2		
	$\Delta_{\text{fus}}H$		8.41	412.7	DSC	[2006LAH/PAA]
	Δ_vH	(323–423)	78.9	373	GC	[1999LEI/WAN]
C ₁₀ H ₃ Cl ₅	[150224-16-1]	1,2,3,6,7-pentachloronaphthalene				
	$\Delta_{\text{fus}}H$		22.94	416.6	DSC	[2006LAH/PAA]
C ₁₀ H ₃ Cl ₅	[150205-21-3]	1,2,3,7,8-pentachloronaphthalene				
	$\Delta_{\text{fus}}H$		17.55	381.6	DSC	[2006LAH/PAA]
C ₁₀ H ₃ Cl ₅	[150224-17-2]	1,2,4,6,7-pentachloronaphthalene				
	$\Delta_{\text{fus}}H$		22.58	404.3	DSC	[2006LAH/PAA]
C ₁₀ H ₃ Cl ₅	[150224-22-9]	1,2,4,6,8-pentachloronaphthalene				
	$\Delta_{\text{fus}}H$		19.75	429	DSC	[2006LAH/PAA]
C ₁₀ H ₄ Cl ₂ O ₂	[117-80-6]	2,3-dichloro-1,4-naphthalenedione				
	$\Delta_{\text{fus}}H$		28.53	469	DSC	[1991ACR, 1990DON/DRE]
C ₁₀ H ₄ Cl ₄	[67922-21-8]	1,2,4,7-tetrachloronaphthalene				

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound					
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference	
C ₁₀ H ₄ Cl ₄		$\Delta_v H$	(323–423)	72.1	373	GC	[1999LEI/WAN]
	[53555-63-8]	1,2,3,5-tetrachloronaphthalene					
	$\Delta_{\text{fus}} H$		27.14	453	DSC	[2006LAH/PAA]	
C ₁₀ H ₄ Cl ₄		$\Delta_v H$	(323–423)	73.4	373	GC	[1999LEI/WAN]
	[20020-02-4]	1,2,3,4-tetrachloronaphthalene					
	$\Delta_{\text{us}} H$		10.53	440.5			
	$\Delta_{\text{us}} H$		1.33	454.3			
C ₁₀ H ₄ Cl ₄		$\Delta_{\text{fus}} H$		11.54	470.8	DSC	[2006LAH/PAA]
		$\Delta_v H$	(323–423)	73.2	373	GC	[1999LEI/WAN]
	[149864-82-4]	1,2,7,8-tetrachloronaphthalene					
C ₁₀ H ₄ Cl ₄		$\Delta_{\text{fus}} H$		14.62	401.3	DSC	[2006LAH/PAA]
	[27304-13-8]	2,3,4,5,6,6a,7,7-octachloro-1a,1b,5,5a,6,6a-hexahydro-2,5-methano-2H-indeno-[1,2-b]oxirene (oxychlordane)					
C ₁₀ H ₄ Cl ₈ O		$\Delta_v H$	(373–403)	75		GC	[2007GOE/MCC]
	[50402-52-3]	1,2,3-trichloronaphthalene					
C ₁₀ H ₅ Cl ₃		$\Delta_{\text{fus}} H$		18.44	354.7	DSC	[2006LAH/PAA]
		$\Delta_v H$	(323–423)	68	373	GC	[1999LEI/WAN]
	[76-44-8]	1,4,5,6,7,8,8-heptachloro-3a,4,7,7a-tetrahydro-4,7-endomethanoindene (heptachlor)					
C ₁₀ H ₅ Cl ₇		$\Delta_{\text{fus}} H$		20.72	362.2	TGA,DSC	[2000ROD/VEC]
		$\Delta_{\text{us}} H$		23.4	358.2		
		$\Delta_{\text{fus}} H$		2.09	371	DSC	[1995KSI/NAG]
		$\Delta_v H$	(343–453)	76.5	398	GC	[1990HIN/BID2]
C ₁₀ H ₅ Cl ₇ O	[1024-57-3]	1,4,5,6,7,8,8-heptachloro-2,3-epoxy-3a,4,7,7a-tetrahydro-4,7-endo-methanoindan					
		$\Delta_{\text{us}} H$		18.9	385.2		
		$\Delta_{\text{fus}} H$		2.85	434.9	DSC	[1995KSI/NAG]
		$\Delta_v H$	(373–423)	75		GC	[2007GOE/MCC]
C ₁₀ H ₅ Cl ₉	[5103-73-1]	<i>cis</i> nonachlor					
		$\Delta_v H$	(343–453)	83.8	398	GC	[1990HIN/BID2]
C ₁₀ H ₅ Cl ₉	[39765-80-5]	<i>trans</i> nonachlor					
		$\Delta_v H$	(343–453)	85.5	398	GC	[1990HIN/BID2]
C ₁₀ H ₆ BrNO ₂	[13380-67-1]	1-(4-bromophenyl)-1H-pyrrole-2,5-dione					
		$\Delta_{\text{sub}} H$	(350–370)	105.9 ± 0.7		C	[1998KIS/KAS]
C ₁₀ H ₆ Br ₂	[83-53-4]	1,4-dibromonaphthalene					
		$\Delta_{\text{sub}} H$	(297–322)	90.8 ± 1.7		ME	[2008GOL/SUU2]
C ₁₀ H ₆ Cl ₂	[2050-69-3]	1,2-dichloronaphthalene					
		$\Delta_v H$	(323–423)	60.7	373	GC	[1999LEI/WAN]
C ₁₀ H ₆ Cl ₂	[1825-31-6]	1,4-dichloronaphthalene					
		$\Delta_v H$	(323–423)	58.7	373	GC	[1999LEI/WAN]
C ₁₀ H ₆ Cl ₄ O ₄	[1861-32-1]	dimethyltetrachloroterephthalane (chlorthal)					
		$\Delta_{\text{sub}} H$	(348–433)	104.9 ± 1.4	390	ME,GS	[1981DEP]
C ₁₀ H ₆ Cl ₄ O ₄	[1861-32-1]	dimethyl-2,3,5,6-tetrachloro-1,4-benzenedicarboxylate					
		$\Delta_{\text{fus}} H$		30.23	431.7		[1991ACR]
C ₁₀ H ₆ Cl ₈	[5103-71-9]	<i>cis</i> chlordane					
		$\Delta_v H$	(323–409)	83	338		[1999DYK/SVO]
		$\Delta_v H$	(343–453)	82	398	GC	[1990HIN/BID2]

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₀ H ₆ Cl ₈	[5103-74-2]	<i>trans</i> chlordane				
	$\Delta_v H$	(373–409)	81.7	388		[1999DYK/SVO]
	$\Delta_v H$	(343–453)	80.7	398	GC	[1990HIN/BID2]
C ₁₀ H ₆ Cl ₈	[5103-71-9]	1,2,4,5,6,7,8,8-octachloro-2,3,3a,4,7,7a-hexahydro-4,7-ethano-1 <i>H</i> -indene				
	$\Delta_{\text{fus}} H$		23.15	379.9	DSC	[1990DON/DRE]
C ₁₀ H ₆ F ₃ NO ₂	[53518-15-3]	4-trifluoromethyl-7-aminocoumarin				
	$\Delta_{\text{fus}} H$		31.7	494.8	DSC	[1991ZHA/HUA]
C ₁₀ H ₆ N ₂	[1436-43-7]	2-cyanoquinoline				
	$\Delta_{\text{sub}} H$		89.3 ± 3.3	298	C	[1995RIB/MAT2]
	$\Delta_{\text{sub}} H$	(312–326)	93.4 ± 0.7	319	ME	[1995RIB/MAT2]
	$\Delta_{\text{sub}} H$		94.4 ± 0.7	298		[1995RIB/MAT2]
C ₁₀ H ₆ N ₂	[34846-64-5]	3-cyanoquinoline				
	$\Delta_{\text{sub}} H$		91.3 ± 1.8	298	C	[1995RIB/MAT2]
	$\Delta_{\text{sub}} H$	(312–326)	93.4 ± 0.7	319	ME	[1995RIB/MAT2]
	$\Delta_{\text{sub}} H$		93.2 ± 0.8	298		[1995RIB/MAT2]
C ₁₀ H ₆ N ₂ O ₄	[7300-93-8]	1-(3-nitrophenyl)-1 <i>H</i> -pyrrole-2,5-dione				
	$\Delta_{\text{sub}} H$	(350–370)	115.7 ± 0.9		C	[1998KIS/KAS]
C ₁₀ H ₆ N ₂ O ₄	[4338-06-1]	1-(4-nitrophenyl)-1 <i>H</i> -pyrrole-2,5-dione				
	$\Delta_{\text{sub}} H$	(350–370)	117.3 ± 1.2		C	[1998KIS/KAS]
C ₁₀ H ₆ N ₂ O ₆	[605-71-0]	1,5-dinitronaphthalene				
	$\Delta_v H$	(506–642)	74.7	521	A	[1987STE/MAL]
C ₁₀ H ₆ N ₂ O ₆	[602-38-0]	1,8-dinitronaphthalene				
	$\Delta_v H$	(553–715)	78.5	568	A	[1987STE/MAL]
C ₁₀ H ₆ OS ₂	[49833-12-7]	naphthalene 1,8-disulfide S-oxide				
	$\Delta_{\text{trs}} H$		3.2	363		
	$\Delta_{\text{fus}} H$		23.3	421.2	DSC	[1975CUC]
C ₁₀ H ₆ O ₂	[130-15-4]	1,4-naphthoquinone				
	$\Delta_{\text{sub}} H$		91.0 ± 0.8	298	C	[1989RIB/RIB]
	$\Delta_{\text{sub}} H$		90.7 ± 2	313	TE,ME	[1981DEK/SMI]
	$\Delta_{\text{sub}} H$		72.4 ± 3.8			[1956MAG, 1970COX/PIL]
(C ₁₀ H ₆ O ₂)– (C ₁₀ H ₈ O ₂)	[21414-85-7]	(1,4-naphthoquinone)-(1,4-naphthohydroquinone)				
	$\Delta_{\text{sub}} H$		102.3 ± 2	342.4	ME,TE	[1981DEK/SMI]
2(C ₁₀ H ₆ O ₂)– (C ₁₀ H ₈ O ₂)	[66653-77-8]	2(1,4-naphthoquinone)-(1,4-naphthohydroquinone)				
	$\Delta_{\text{sub}} H$		88.7 ± 3	328.5	ME,TE	[1981DEK/SMI]
C ₁₀ H ₆ S ₂	[209-22-3]	naphthalene 1,8-disulfide				
	$\Delta_{\text{fus}} H$		13.0	394.8	DSC	[1975CUC2]
C ₁₀ H ₇ Br	[90-11-9]	1-bromonaphthalene				
	$\Delta_{\text{fus}} H$		15.16	271.4		[1991ACR]
	$\Delta_v H$	(303–336)	63.9 ± 0.4	298	GS	[2003VER]
	$\Delta_v H$	(357–555)	58.5	372		[1987STE/MAL]
	$\Delta_v H$	(295–359)	56 ± 6	329	ME	[1980URB/GIG]
	$\Delta_v H$	(469–559)	45.8	484	A,EB	[1987STE/MAL, 1976HON/SIN, 1999DYK/SVO]
C ₁₀ H ₇ Br	[580-13-2]	2-bromonaphthalene				
	$\Delta_{\text{trs}} H$		5.76	319		
	$\Delta_{\text{fus}} H$		14.4	329		[1996DOM/HEA]
	$\Delta_{\text{sub}} H$		81.2 ± 1.0	298	C	[1993RIB/FER]

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound					
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference	
		$\Delta_{\text{sub}}H$	(275–378)	64 ± 5	298	TE,ME	[1981FER/PIA]
		Δ_vH	(330–360)	66.1 ± 0.4	298	GS	[2003VER]
		Δ_vH	(330–378)	42.5	354		[1999DYK/SVO]
		Δ_vH	(322–359)	40.4	340	ME,TE	[1981FER/PIA]
C₁₀H₇Cl	[90-13-1]	1-chloronaphthalene					
		$\Delta_{\text{fus}}H$		12.9	270.7		[1991ACR]
		Δ_vH		64.7	298		[2006BOL/NER2]
		Δ_vH	(289–332)	62.0 ± 0.4	298	GS	[2003VER]
		Δ_vH		64.0 ± 0.3	298	GS	[2001PUR/CHI]
		Δ_vH	(323–423)	58.6	373	GC	[1999LEI/WAN]
		Δ_vH	(353–553)	59.6	368	A	[1987STE/MAL, 1947STU]
		Δ_vH	(400–435)	57.8	415	A	[1987STE/MAL]
C₁₀H₇Cl	[91-58-7]	2-chloronaphthalene					
		$\Delta_{\text{fus}}H$		14.7	332		[1991ACR]
		Δ_vH	(332–362)	62.3 ± 1.1	298	GS	[2003VER]
		Δ_vH	(400–435)	57.9	417		[1999DYK/SVO]
		Δ_vH	(323–423)	58.5	373	GC	[1999LEI/WAN]
C₁₀H₇Cl₅O	[58138-08-2]	2-(3,5-dichlorophenyl)-2-(2,2,2-trichloroethyl)oxirane					
		$\Delta_{\text{fus}}H$		18.54	313.2	DSC	[1990DON/DRE]
C₁₀H₇Cl₇	[2589-15-3]	1,4,5,6,7,8,8-heptachloro-3a,4,7,7a-tetrahydro-4,7-methanoindan					
		$\Delta_{\text{sub}}H$	(333–353)	83.8	343		[1987STE/MAL, 1974BES/CHE]
		Δ_vH	(333–353)	83.8	343		[1999DYK/SVO]
C₁₀H₇F₃O₂	[326-06-7]	benzoyltrifluoroacetone					
		$\Delta_{\text{sub}}H$		87.1 ± 0.9	298	ME	[1992RIB/MON]
C₁₀H₇F₅O₂	[24277-51-0]	pentafluoropropionic acid, 3-tolyl ester					
		Δ_vH	(371–446)	48.5	386	A,EB	[1987STE/MAL, 1969SHE/LAN, 1999DYK/SVO]
C₁₀H₇F₅O₂	[24271-52-1]	pentafluoropropionic acid, 4-tolyl ester					
		Δ_vH	(371–448)	48.3	386	A,EB	[1987STE/MAL, 1969SHE/LAN, 1999DYK/SVO]
C₁₀H₇I	[90-14-2]	1-iodonaphthalene					
		$\Delta_{\text{fus}}H$		15.91	280		[1991ACR]
		Δ_vH	(303–347)	69.9 ± 0.3	298	GS	[2003VER]
		Δ_vH	(321–428)	78.9	336		[1999DYK/SVO]
C₁₀H₇I	[612-55-5]	2-iodonaphthalene					
		$\Delta_{\text{fus}}H$		16.04	327.6		[1991ACR]
		$\Delta_{\text{sub}}H$		90.8			[1956SMI]
C₁₀H₇NO₂	[86-57-7]	1-nitronaphthalene					
		$\Delta_{\text{fus}}H$		17.3	328.9	DSC	[2010KES/AUC]
		$\Delta_{\text{fus}}H$		18.43	329.9		[1991ACR]
		$\Delta_{\text{sub}}H$	(305–321)	94.4 ± 0.4	313	ME	[2006RIB/AMA3]
		$\Delta_{\text{sub}}H$	(305–321)	95.1 ± 0.4	298	ME	[2006RIB/AMA3]
		$\Delta_{\text{sub}}H$	(309–326)	U68.5 ± 1.9	318		[1987STE/MAL, 1974RAD/KAT]
		$\Delta_{\text{sub}}H$	(325–332)	106.9	328.5		[1987STE/MAL]
		$\Delta_{\text{sub}}H$		107.1 ± 2.1			[1950NIT/SEK, 1970COX/PIL]
		Δ_vH	(332–580)	66.4	347	A	[1987STE/MAL]

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₀ H ₇ NO ₂	[581-89-5]	2-nitronaphthalene	14.5	348.2		[2010KES/AUC]
C ₁₀ H ₇ NO ₂	[131-91-9]	1-nitroso-2-naphthol	86.6 ± 4.2		ME	[1968HAM/FAG, 1977PED/RYL]
C ₁₀ H ₇ NO ₂	[132-53-6]	2-nitroso-1-naphthol	56.5 ± 4.2		ME	[1968HAM/FAG, 1977PED/RYL]
C ₁₀ H ₇ NO ₂	[605-60-7]	4-nitroso-1-naphthol	87.4 ± 4.2		ME	[1968HAM/FAG, 1977PED/RYL]
C ₁₀ H ₇ NO ₂	[941-69-5]	1-phenyl-1 <i>H</i> -pyrrole-2,5-dione (350–370)	98.1 ± 1		C	[1998KIS/KAS]
C ₁₀ H ₇ N ₃ S	[148-79-8]	2-(4-thiazolyl)-1 <i>H</i> -benzimidazole (thiabenzazole)	35.2	573.2	DSC	[2010MUE/ESC]
C ₁₀ H ₈	[275-51-4]	azulene	17.53	373.5	DSC	[1998CHI/HES]
	$\Delta_{\text{sub}}H$	(283–326)	78.4 ± 1.3	303	HSA	[1998CHI/HES]
	$\Delta_{\text{sub}}H$		72.7	298	CGC-DSC	[1998CHI/HES]
	$\Delta_{\text{sub}}H$	(290–372)	82.8	305	S	[1987STE/MAL]
	$\Delta_{\text{sub}}H$		82.9	298	H	[1987STE/MAL, 1993CHI/HOS]
	$\Delta_{\text{sub}}H$	(253–293)	75.8	273		[1958HOY/PEP]
	$\Delta_{\text{sub}}H$		75.3	298	H	[1987STE/MAL, 1993CHI/HOS]
	$\Delta_{\text{sub}}H$		76.8 ± 0.2		C	[1972MOR]
	$\Delta_{\text{sub}}H$	(293–323)	95.4 ± 0.4	298	ME	[1962BAU/GUN, 1970COX/PIL]
	$\Delta_{\text{sub}}H$		67.6			[1947HEI/WIE]
	Δ_vH		52.8	298	CGC	[1998CHI/HES]
	Δ_vH	(369–515)	53.0	384	A	[1987STE/MAL]
	Δ_vH	(442–534)	51.2	457	EB	[1977MEY/GEN]
	Δ_vH	(373–423)	55.5	373		[1962BAU/GUN]
C ₁₀ H ₈	[91-20-3]	naphthalene	19.1	353.5	DSC	[2008SHA/GUP]
	$\Delta_{\text{fus}}H$		16.44	353.8	DSC	[2007HAF/MAH]
	$\Delta_{\text{fus}}H$		19.55	354.7	DSC	[2006KHI/DAH2]
	$\Delta_{\text{fus}}H$		19.55	354.7	DSC	[2006KHI/DAH]
	$\Delta_{\text{fus}}H$		19.1	Not given	DSC	[2003SHA/KAN]
	$\Delta_{\text{fus}}H$	(5–440)	19.0	353.4	AC	[2002CHI/KNI]
	$\Delta_{\text{fus}}H$		19.1	353.4		[1991ACR]
	$\Delta_{\text{sub}}H$	(267–303)	88.0 ± 2.5		ME	[1998BOL/WIE]
	$\Delta_{\text{sub}}H$		70.4	298	CGC-DSC	[1998CHI/HES]
	$\Delta_{\text{sub}}H$	(313–353)	71.7	333	GS	[1995NAS/LEN]
	$\Delta_{\text{sub}}H$	(243–273)	73.7 ± 1.0	258	GS	[1994WAN/SHU]
	$\Delta_{\text{sub}}H$	(337–352)	78.2 ± 1		GC	[1988KHU]
	$\Delta_{\text{sub}}H$		70.9 ± 0.4	323	DSC	[1988TOR/BAR]
	$\Delta_{\text{sub}}H$		72.3 ± 0.4	298	DSC	[1988TOR/BAR]
	$\Delta_{\text{sub}}H$	(299–331)	73.4	315	GS	[1986SAT/INO]
	$\Delta_{\text{sub}}H$	(333–393)	69.9		GS	[1985MAT/KUW2]
	$\Delta_{\text{sub}}H$	(293–331)	72.3 ± 0.8		QR	[1985GLU/ARK]
	$\Delta_{\text{sub}}H$	(283–323)	75.8 ± 1.1	303	GS	[1983SON/ZOL]
	$\Delta_{\text{sub}}H$		72.6 ± 0.4		DSC	[1983HOL]
	$\Delta_{\text{sub}}H$		72.6 ± 0.1	298	TE,ME,DM	[1983VAN/JAC, 1981DEK/KUI]
	$\Delta_{\text{sub}}H$	(302–352)	72.8	327	GS	[1982GRA/FOS]
	$\Delta_{\text{sub}}H$	(271–285)	72.8 ± 0.3		ME	[1982COL/JIM]
	$\Delta_{\text{sub}}H$		72.4 ± 0.7	298	C	[1982MUR/SAK]
$\Delta_{\text{sub}}H$	(274–353)	72.5 ± 0.1		DM	[1981DEK/KUI]	
$\Delta_{\text{sub}}H$	(328–398)	76.0 ± 2.0		DSC	[1980MUR/CAV]	

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{sub}}H$	(253–273)	72.6 ± 0.6		TE	[1980DEK]
	$\Delta_{\text{sub}}H$	(280–305)	71.3	293	GS	[1979MAC/PRA]
	$\Delta_{\text{sub}}H$	(253–273)	74.77 ± 0.4		TE	[1977DEK/VAN]
	$\Delta_{\text{sub}}H$	(253–273)	73.9 ± 0.2		ME	[1977DEK/VAN]
	$\Delta_{\text{sub}}H$	(303–329)	74.35 ± 1.7		TSGC	[1975MCE/SAN]
	$\Delta_{\text{sub}}H$		72.3 ± 0.4		C	[1976FER/PIA]
	$\Delta_{\text{sub}}H$	(263–343)	72.5 ± 0.3		DM	[1975AMB/LAW]
	$\Delta_{\text{sub}}H$		67.8 ± 3.5	280	HSA	[1975CHI]
	$\Delta_{\text{sub}}H$		72.5	298	GS	[1974SIN]
	$\Delta_{\text{sub}}H$		72.7 ± 1.7			[1974RAD/KAT]
	$\Delta_{\text{sub}}H$	(281–290)	64 ± .5		LE	[1973MCE/SAN]
	$\Delta_{\text{sub}}H$		72.1 ± 0.25	298	C	[1972MOR]
	$\Delta_{\text{sub}}H$		73.0 ± 0.3	298	C	[1972IRV]
	$\Delta_{\text{sub}}H$	(283–323)	72.7		ME	[1971RAD]
	$\Delta_{\text{sub}}H$		66.5			[1968KAR/RAB]
	$\Delta_{\text{sub}}H$	(230–260)	72.7 ± 0.3		KG	[1963MIL, 1970COX/PIL]
	$\Delta_{\text{sub}}H$	(276–283)	66.3		V	[1959AIH]
	$\Delta_{\text{sub}}H$	(283–303)	65.8	293	Effusion	[1958SKL/MAR]
	$\Delta_{\text{sub}}H$	(253–283)	69.2	268		[1958HOY/PEP]
	$\Delta_{\text{sub}}H$	(273–311)	72.1	292		[1957SHE/BRY]
	$\Delta_{\text{sub}}H$	(279–294)	72.4			[1953BRA/CLE2, 1960JON, 1954SEA/HOP]
	$\Delta_{\text{sub}}H$		64.0	298	ME	[1951INO]
	$\Delta_{\text{sub}}H$	(288–306)	65.7	297	Effusion	[1940ZIB]
	$\Delta_{\text{sub}}H$		66.5 ± 1.7	298	QF	[1938WOL/WEG]
	$\Delta_{\text{sub}}H$	(237–276)	76.6			[1926AND]
	$\Delta_{\text{sub}}H$	(283–303)	82.0	293	ME	[1925SWA/MAC]
	Δ_vH		54.6	298	CGC	[2008ZHA/UNH]
	Δ_vH	(333–403)	60.3 ± 1.1	298	GC	[2006HAF/PAR]
	Δ_vH	(323–473)	56.1	398	GC	[2002LEI/CHA]
	Δ_vH	(491–747)	47.6 ± 1.8		DSC	[1998BOL/WIE]
	Δ_vH		53.4	298	CGC	[1998CHI/HES]
	Δ_vH	(460–647)	45.4	475	DSC	[1996BAC/GRZ]
	Δ_vH	(403–453)	56.6	298	CGC	[1995CHI/HOS]
	Δ_vH		48.7 ± 0.3	400	EB	[1993CHI/KNI]
	Δ_vH		46.4	440	EB	[1993CHI/KNI]
	Δ_vH		44.0	480	EB	[1993CHI/KNI]
	Δ_vH		41.5	520	EB	[1993CHI/KNI]
	Δ_vH	(513–613)	44.4	528		[1993LEE/HOL]
	Δ_vH	(418–613)	47.9	423	EB	[1990AMB/EWI]
	Δ_vH	(352–500)	50.6	367	A	[1987STE/MAL]
	Δ_vH	(491–565)	44.8	506	A	[1987STE/MAL]
	Δ_vH	(563–663)	43.2	578	A	[1987STE/MAL]
	Δ_vH	(661–750)	43.3	676	A	[1987STE/MAL]
	Δ_vH		51.5		GS	[1985MAT/KUW2]
	Δ_vH	(441–727)	44.7	466		[1981WIL/JOH]
	Δ_vH	(353–388)	50.3 ± 0.2	370		[1981DEK/KUI]
	Δ_vH	(354–453)	50.7	369		[1968FOW/TRU]
	Δ_vH	(399–491)	49.0	414		[1955CAM/ROS]
	Δ_vH		51.5			[1952GOT]
	Δ_vH		46.4	441	C	[1951BAR/MCC]
	Δ_vH		48.3	379	I	[1943CRA]
	Δ_vH	(373–473)	47.2	423	I	[1923MOR/MUR]
	Δ_vH	(360–494)	47.7	427	I	[1922NEL/SEN]
C₁₀D₈	[1146-65-2]	naphthalene-d ₈				
	$\Delta_{\text{sub}}H$	(282–323)	70.6 ± 0.5	303	GS	[1983SON/ZOL]
	Δ_vH		54.7	298	CGC	[2008ZHA/UNH]

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₀ H ₈ BrNO ₂	[574-98-1] $\Delta_{\text{sub}}H$	N-(2-bromoethyl)phthalimide	108.7 ± 1.0	298	C	[2007RIB/SAN3]
C ₁₀ H ₈ Br ₂ N ₂	[3138-86-1] $\Delta_{\text{fus}}H$	2,3- <i>bis</i> -(bromomethyl)quinoxaline	32.43	423.6	DSC	[2000MON/HIL2]
	$\Delta_{\text{sub}}H$	(351–365)	111.7 ± 0.5	358	ME	[2000MON/HIL2]
	$\Delta_{\text{sub}}H$	(351–365)	114.0 ± 2.0	298	ME	[2000MON/HIL2]
C ₁₀ H ₈ ClNO ₂	[6270-06-0] $\Delta_{\text{sub}}H$	N-(2-chloroethyl)phthalimide	98.4 ± 1.9	298	C	[2007RIB/SAN3]
C ₁₀ H ₈ ClN ₃ O	[1698-60-8] $\Delta_{\text{fus}}H$	5-amino-4-chloro-2-phenyl-3(2 <i>H</i>)-pyridazinone	26.75	479.2	DSC	[1991ACR, 1990DON/DRE]
C ₁₀ H ₈ ClN ₃ O ₂	[5707-69-7] $\Delta_{\text{fus}}H$	4-(2-chlorophenylhydrazone)-3-methyl-5-isoxazolone	28.04	440.4	DSC	[1990DON/DRE]
C ₁₀ H ₈ Cl ₂ O ₆	[24648-18-8] $\Delta_{\text{fus}}H$	3,6-dichloro-2,5-dihydroxyterephthalate	1.7	380		
	$\Delta_{\text{fus}}H$ (white)		41.0	455		[1990RIC/YAN]
	$\Delta_{\text{fus}}H$ (white)		44.6	461	DSC	[1972BYR/CUR]
	$\Delta_{\text{fus}}H$ (yellow--white)		2.6	403	DSC	[1972BYR/CUR]
C ₁₀ H ₈ NO ₂	[87-51-4] $\Delta_{\text{sub}}H$	indole-3-acetic acid	64.0 ± 1.5	368	ME	[1988TOR/BAR]
		(313–423)				
C ₁₀ H ₈ N ₂	[366-18-7] $\Delta_{\text{fus}}H$	2,2'-bipyridine	20.4	345	DSC	[2009LIP/HAN]
	$\Delta_{\text{sub}}H$		81.8 ± 2.3	298	C	[1995RIB/MOR]
	$\Delta_{\text{sub}}H$		75.0 ± 5.0	298	B	[1996CHA/EMM]
	$\Delta_{\text{sub}}H$		81.9 ± 0.3			[1985SKI/PIL]
	Δ_vH		67.0 ± 2.3	298	CGC	[2009LIP/CHI, 2009LIP/HAN]
C ₁₀ H ₈ N ₂	[581-47-5] $\Delta_{\text{fus}}H$	2,4'-bipyridine	17.4	332.8	DSC	[2009LIP/HAN]
	$\Delta_{\text{sub}}H$		87.9 ± 1.7	298	C	[1995RIB/MOR]
	Δ_vH		70.9 ± 1.6	298	CGC	[2009LIP/CHI, 2009LIP/HAN]
C ₁₀ H ₈ N ₂	[553-26-4] $\Delta_{\text{fus}}H$	4,4'-bipyridine	16.1	377.5	DSC	[2009LIP/HAN]
	$\Delta_{\text{sub}}H$		106.3 ± 2.8	298	C	[1995RIB/MOR]
	Δ_vH		71.1 ± 2.6	298	CGC	[2009LIP/CHI, 2009LIP/HAN]
C ₁₀ H ₈ N ₂	[3438-48-0] $\Delta_{\text{fus}}H$	4-phenylpyrimidine	18.8	334.1	DSC	[2009LIP/HAN]
	Δ_vH		68.8 ± 2.5	298	CGC	[2009LIP/HAN]
C ₁₀ H ₈ N ₂ O ₂	[3634-83-1] Δ_vH	1,3- <i>bis</i> -(isocyanatomethyl)benzene	46.7	418	A	[1987STE/MAL]
		(403–473)				
C ₁₀ H ₈ N ₂ O ₂	[1014-98-8] Δ_vH	1,4- <i>bis</i> -(isocyanatomethyl)benzene	56.9	418	A	[1987STE/MAL]
		(403–473)				
C ₁₀ H ₈ N ₂ O ₂	[64711-83-7] Δ_vH	benzene, ethyldiisocyanato (mixed isomers)	60.7	378	A	[1987STE/MAL, 1977ZHU/MEL]
		(363–473)				
C ₁₀ H ₈ N ₂ O ₂	[881-07-2]	8-nitroquinaldine				

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound					
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference	
		$\Delta_{\text{sub}}H$	(346–360)	108.3 ± 0.8	353	ME	[1997RIB/MAT5]
		$\Delta_{\text{sub}}H$	(346–360)	111.0 ± 0.8	298	ME	[1997RIB/MAT5]
C ₁₀ H ₈ N ₂ O ₃	[6118-65-6]	3-acetamidophthalimide					
		$\Delta_{\text{sub}}H$	(428–468)	108.5	443	A	[1987STE/MAL, 1956KLO]
C ₁₀ H ₈ O	[90-15-3]	1-naphthol					
		$\Delta_{\text{fus}}H$		23.3	368.7	DSC	[2002RAI/PAN]
		$\Delta_{\text{fus}}H$		24.4	369.7	DSC	[1998RAI/RAI]
		$\Delta_{\text{fus}}H$		23.01	369		[1991ACR]
		$\Delta_{\text{fus}}H$		23.47	368.2	C	[1926AND/LYN]
		$\Delta_{\text{sub}}H$	(296–313)	91.2 ± 0.4		ME	[1974COL/ROU2]
		$\Delta_{\text{sub}}H$	(279–328)	89.1 ± 1.7	304	ME	[1974ARS]
		$\Delta_{\text{sub}}H$	(298–312)	93.3	305	A	[1987STE/MAL, 1960AIH]
		$\Delta_{\text{sub}}H$	(314–324)	84.3	319	A	[1987STE/MAL, 1960AIH]
		Note: In Ref. [1960AIH] the author mentions that there may be a small phase transition at 39.4 °C as evidenced in the log P versus 1/T graph					
		$\Delta_{\text{sub}}H$		91.5 ± 3.8	298	B	[1926AND/LYN, 1970COX/PIL, 1927MAY/BER]
		Δ_vH	(399–556)	58.5	414	A	[1987STE/MAL]
		Δ_vH	(423–563)	60.8	473		[1927MAY/BER]
C ₁₀ H ₈ O	[135-19-3]	2-naphthol					
		$\Delta_{\text{fus}}H$	(363–413)	20.9	392.5	DSC	[2003ROJ/ORO]
		$\Delta_{\text{fus}}H$		18.79	393.6	C	[1991ACR, 1926AND/LYN]
		$\Delta_{\text{sub}}H$		85.5 ± 1.2	298	DSC	[2003ROJ/ORO]
		$\Delta_{\text{sub}}H$	(305–323)	94.2 ± 0.5		ME	[1974COL/ROU2]
		$\Delta_{\text{sub}}H$	(277–324)	87.4 ± 2.5	300	ME	[1974ARS]
		$\Delta_{\text{sub}}H$	(283–323)	78.7 ± 0.8	298	A	[1968KAR/RAB, 1977PED/RYL, 1987STE/MAL]
		$\Delta_{\text{sub}}H$	(298–312)	97.8	305	A	[1987STE/MAL, 1960AIH]
		$\Delta_{\text{sub}}H$	(314–332)	87.8	323	A	[1987STE/MAL, 1960AIH]
		Note: In reference [1960AIH] the author mentions that there may be a small Phase transition at 39.1 °C as evidenced in the log P versus 1/T graph					
		$\Delta_{\text{sub}}H$		83.0 ± 3.8	298	B	[1926AND/LYN, 1927MAY/BER, 1970COX/PIL]
		Δ_vH	(393–433)	76.2	298	CGC	[1995CHI/HOS]
		Δ_vH	(401–561)	59.7	416	A	[1987STE/MAL]
		Δ_vH	(417–561)	59.7	432		[1955VON/GEB]
		Δ_vH	(423–563)	61.8	473		[1927MAY/BER]
C ₁₀ H ₈ O	[4759-11-9]	1,6-oxido[10]annulene					
		$\Delta_{\text{sub}}H$		80.4 ± 8.4		B	[1969BRE/HAG, 1977PED/RYL]
C ₁₀ H ₈ OS ₃	[532-11-6]	5-(4-methoxyphenyl)-3H-1,2-dithiole-3-thione					
		$\Delta_{\text{fus}}H$		24.39	382.2	DSC	[1999DOL/LEC]
C ₁₀ H ₈ O ₂	[571-60-8]	1,4-naphthohydroquinone					
		$\Delta_{\text{sub}}H$		119 ± 1	381	ME,TE	[1981DEK/SMI]
C ₁₀ H ₈ O ₂	[574-00-5]	1,2-dihydroxynaphthalene					
		$\Delta_{\text{sub}}H$		109.3 ± 0.9	298	C	[1988RIB/RIB]
C ₁₀ H ₈ O ₂	[132-86-5]	1,3-dihydroxynaphthalene					
		$\Delta_{\text{sub}}H$		116.0 ± 1.1	298	C	[1988RIB/RIB]
C ₁₀ H ₈ O ₂	[92-44-4]	2,3-dihydroxynaphthalene					
		$\Delta_{\text{sub}}H$		109.6 ± 1.0	298	C	[1988RIB/RIB]
		$\Delta_{\text{sub}}H$	(341–359)	109.4 ± 0.5	350	ME	[1979COL/JIM2]

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₀ H ₈ O ₃	[90-33-5] $\Delta_{\text{fus}}H$	4-methyl-7-hydroxycoumarin	29.14	460.7		[1996DOM/HEA]
C ₁₀ H ₉ Br	[2007844-51-7] $\Delta_{\text{fus}}H$	bromobullvalene	14.2	317.2	DSC	[1998LUS/OLI]
C ₁₀ H ₉ Cl	[27576-94-9] $\Delta_{\text{fus}}H$	chlorobullvalene	13.5	287.2	DSC	[1998LUS/OLI]
C ₁₀ H ₉ Cl ₂ NO	[2164-09-2] $\Delta_{\text{fus}}H$	N-(3,4-dichlorophenyl)-2-methyl-2-propenamide	32.04	395.5	DSC	[1991ACR, 1990DON/DRE]
C ₁₀ H ₉ Cl ₃ O ₃	[1928-39-8] Δ_vH	(2,4,5-trichlorophenoxy)acetic acid, ethyl ester (444–573)	76.4	459	A	[1987STE/MAL, 1999DYK/SVO]
C ₁₀ H ₉ Cl ₃ O ₃	[1928-37-6] $\Delta_{\text{fus}}H$	methyl 2-(2,4,5-trichlorophenoxy)propionate	31.95	360.6		[1991ACR]
C ₁₀ H ₉ Cl ₃ O ₃	[93-80-1] $\Delta_{\text{fus}}H$	4-(2,4,5-trichlorophenoxy)butanoic acid	30.28	386.7	DSC	[1991ACR, 1990DON/DRE]
C ₁₀ H ₉ Cl ₄ NO ₂ S	[2425-06-1] $\Delta_{\text{fus}}H$ $\Delta_{\text{fus}}H$	N-[(1,1,2,2-tetrachloroethyl)thio]-4-cyclohexene-1,2-dicarboximide	40.22 43.1	432.7 432	DSC DSC	[1990DON/DRE] [1969PLA/GLA]
C ₁₀ H ₉ I	[207844-52-8] $\Delta_{\text{fus}}H$	iodobullvalene	15.5	376.2	DSC	[1998LUS/OLI]
C ₁₀ H ₉ IO ₂	[122200-58-2] $\Delta_{\text{fus}}H$	methyl-4-iodocubanecarboxylate	22.3	395		[2010GRI/TSA]
C ₁₀ H ₉ N	[91-63-4] $\Delta_{\text{fus}}H$	2-methylquinoline (quinaldine)	12.52	270.5	AC,DSC	[2005CHI/STE]
	Δ_vH	(319–553)	62.6 ± 0.1	298	IP,EB	[2005CHI/STE]
	Δ_vH	(319–553)	61.0 ± 0.1	320	IP,EB	[2005CHI/STE]
	Δ_vH	(319–553)	58.2 ± 0.1	360	IP,EB	[2005CHI/STE]
	Δ_vH	(319–553)	55.6 ± 0.1	400	IP,EB	[2005CHI/STE]
	Δ_vH	(319–553)	53.0 ± 0.1	440	IP,EB	[2005CHI/STE]
	Δ_vH	(319–553)	50.4 ± 0.1	480	IP,EB	[2005CHI/STE]
	Δ_vH	(319–553)	47.7 ± 0.2	520	IP,EB	[2005CHI/STE]
	Δ_vH		66.1 ± 1.9	298	C	[1995RIB/MAT]
	Δ_vH	(281–313)	61.2	297	GS	[1980VAN/PRA]
	Δ_vH	(443–521)	54.7	548	A,EB	[1987STE/MAL, 1961MAL2, 1961MAL]
C ₁₀ H ₉ N	[612-58-8] Δ_vH	3-methylquinoline (443–528)	55.8	458	A	[1987STE/MAL, 1961MAL2]
C ₁₀ H ₉ N	[491-35-0] Δ_vH Δ_vH	4-methylquinoline (lepidine) (463–539)	67.6 ± 1.8 58.2	298 478	C A,EB	[1995RIB/MAT] [1987STE/MAL, 1961MAL2, 1961MAL]
C ₁₀ H ₉ N	[91-62-3] Δ_vH Δ_vH	6-methylquinoline (453–540)	67.7 ± 1.8 56.1	298 468	C A	[1995RIB/MAT] [1987STE/MAL]
C ₁₀ H ₉ N	[612-60-2] Δ_vH	7-methylquinoline (493–532)	56.7	508	A,EB	[1987STE/MAL, 1961MAL]
C ₁₀ H ₉ N	[611-32-5] $\Delta_{\text{fus}}H$ Δ_vH Δ_vH	8-methylquinoline (324–553) (324–553)	10.73 62.1 ± 0.1 59.2 ± 0.1	246.9 298 340	AC,DSC IP,EB IP,EB	[2005CHI/STE] [2005CHI/STE] [2005CHI/STE]

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(324–553)	56.6 ± 0.1	380	IP,EB	[2005CHI/STE]
	$\Delta_v H$	(324–553)	54.0 ± 0.1	420	IP,EB	[2005CHI/STE]
	$\Delta_v H$	(324–553)	51.3 ± 0.1	460	IP,EB	[2005CHI/STE]
	$\Delta_v H$	(324–553)	49.0 ± 0.2	500	IP,EB	[2005CHI/STE]
	$\Delta_v H$	(324–553)	46.2 ± 0.3	540	IP,EB	[2005CHI/STE]
	$\Delta_v H$		65.7 ± 1.9	298	C	[1995RIB/MAT]
	$\Delta_v H$	(493–523)	52.2	508	A,EB	[1987STE/MAL, 1961MAL]
C₁₀H₉N	[134-32-7]	1-naphthylamine				
	$\Delta_{\text{fus}}H$		15.53	323.2		[1991ACR]
	$\Delta_{\text{sub}}H$	(290–320)	88.1 ± 0.4	298	GS	[2007VER/GEO]
	$\Delta_{\text{sub}}H$		90.0 ± 4.2		TE	[1947BAL, 1970COX/PIL]
	$\Delta_v H$	(323–353)	73.3 ± 0.4	298	GS	[2007VER/GEO]
	$\Delta_v H$	(377–574)	63.6	392	A	[1987STE/MAL, 1947STU]
C₁₀H₉N	[91-59-8]	2-naphthylamine				
	$\Delta_{\text{fus}}H$		23.33	386.2		[1991ACR]
	$\Delta_{\text{sub}}H$	(283–323)	73.9	298	A	[1987STE/MAL]
	$\Delta_{\text{sub}}H$		74.1 ± 1.7			[1968KAR/RAB, 1977PED/RYL]
	$\Delta_{\text{sub}}H$		88.3 ± 4.2			[1947STU, 1970COX/PIL]
	$\Delta_v H$	(388–579)	63.5	403	A	[1987STE/MAL, 1947STU]
C₁₀H₉NO	[5263-87-6]	6-methoxyquinoline				
	$\Delta_v H$		78.1 ± 2.3	298	C	[2003RIB/SAN]
C₁₀H₉NO	[5343-98-6]	β -cyanopropiophenone				
	$\Delta_{\text{sub}}H$		101.7 ± 4.2		ME	[1969LEB/DNE, 1977PED/RYL]
	$\Delta_{\text{sub}}H$	(318–333)	108.5	325.5	A	[1987STE/MAL]
C₁₀H₉NO	[826-81-3]	2-methyl-8-hydroxyquinoline				
	$\Delta_{\text{sub}}H$	(296–307)	90.4 ± 0.7		ME	[1989RIB/MON]
	$\Delta_{\text{sub}}H$		87.2 ± 1.9		C	[1989RIB/MON]
	$\Delta_{\text{sub}}H$	(308–333)	87.9		ME	[1987STE/MAL, 1963HOR/WEN]
C₁₀H₉NO	[18615-86-6]	2-methyl-4-hydroxyquinoline				
	$\Delta_{\text{sub}}H$	(424–442)	132.2 ± 1.0	433	ME	[1990RIB/MAT]
	$\Delta_{\text{sub}}H$		139.0 ± 1.0	298		[1990RIB/MAT]
C₁₀H₉NO	[607-66-9]	4-methyl-2-hydroxyquinoline				
	$\Delta_{\text{sub}}H$	(391–405)	123.1 ± 1.6	398	ME	[1990RIB/MAT]
	$\Delta_{\text{sub}}H$		128.1 ± 1.6	298		[1990RIB/MAT]
C₁₀H₉NO₂	[87-51-4]	indole-3-acetic acid				
	$\Delta_{\text{sub}}H$	(313–423)	64.0 ± 1.4U	368	ME	[1988GAL/GON]
C₁₀H₉NO₂	[26093-31-2]	4-methyl-7-aminocoumarin				
	$\Delta_{\text{fus}}H$		32.09	499.9		[1996DOM/HEA]
C₁₀H₉NO₂	[5022-29-7]	N-ethylphthalimide				
	$\Delta_{\text{sub}}H$		90.9 ± 1.2	298	C	[2006RIB/SAN]
C₁₀H₉NO₂	[6563-13-9]	6-methoxyquinoline N-oxide				
	$\Delta_{\text{sub}}H$		117.9 ± 1.0	298	C	[2003RIB/SAN]
C₁₀H₁₀	[5603-34-9]	pentacyclo[4.4.0.0 ^{2,5} .0 ^{3,8} .0 ^{4,7}]dec-9-ene (basketene)				
	$\Delta_{\text{fus}}H$		2.72	331.8	DSC	[2002VER/KUM]
	$\Delta_{\text{sub}}H$	(274–313)	55.4 ± 0.5	294	GS	[2002VER/KUM]
	$\Delta_{\text{sub}}H$	(274–313)	55.3 ± 0.5	298	GS	[2002VER/KUM]

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₀ H ₁₀	[21604-76-2]	tricyclo[4.2.2.0 ^{2,5}]dec-3,7,9-triene				
	$\Delta_{\text{fus}}H$		1.46	293.7	DSC	[2002VER/KUM]
	Δ_vH	(296–326)	47.2 ± 0.4	311	GS	[2002VER/KUM]
		(296–326)	47.9 ± 0.4	298	GS	[2002VER/KUM]
C ₁₀ H ₁₀	[26934-61-2]	2,2a,2b,3,5a,5b-hexahydro-1,2,3-metheno-1H-cycloprop[cd]indene (snoutene)				
	$\Delta_{\text{fus}}H$		7.87	334	DSC	[2002VER/KUM]
	$\Delta_{\text{sub}}H$	(274–313)	58.9 ± 0.4	294	GS	[2002VER/KUM]
		(274–313)	58.7 ± 0.4	298	GS	[2002VER/KUM]
C ₁₀ H ₁₀	[108-57-6]	1,3-divinylbenzene				
Δ_vH		(305–453)	48.3	320	A	[1987STE/MAL, 1947STU]
C ₁₀ H ₁₀	[77-73-6]	dicyclopentadiene				
Δ_vH		(307–440)	42.4	322		[1947STU]
C ₁₀ H ₁₀	[447-53-0]	1,2-dihydronaphthalene				
	$\Delta_{\text{fus}}H$	(5–444)	10.53	264.4	AC	[2008CHI/STE]
	Note: Authors report two solid/solid phase transitions having negligible enthalpy of transition					
	Δ_vH		54.8 ± 0.1	298	IP,EB	[2008CHI/STE]
	Δ_vH		53.4 ± 0.1	320	IP,EB	[2008CHI/STE]
	Δ_vH		50.8 ± 0.1	360	IP,EB	[2008CHI/STE]
	Δ_vH		48.4 ± 0.1	400	IP,EB	[2008CHI/STE]
	Δ_vH		45.9 ± 0.1	440	IP,EB	[2008CHI/STE]
	Δ_vH	(274–319)	51.9 ± 0.4	296	GS	[1999VER6]
	(274–319)	51.9 ± 0.4	298	GS	[1999VER6]	
C ₁₀ H ₁₀	[612-17-9]	1,4-dihydronaphthalene				
	$\Delta_{\text{fus}}H$		9.35	298.1	DSC	[1999VER6]
	$\Delta_{\text{sub}}H$		63.6 ± 1.6	298		[1999VER6]
	Δ_vH	(300–333)	53.2 ± 0.4	296	GS	[1999VER6]
		(300–333)	54.2 ± 0.4	298	GS	[1999VER6]
C ₁₀ H ₁₀	[5187-81-5]	diisopropenyldiacetylene				
Δ_vH			50.2			[1977LEB/RYA]
C ₁₀ H ₁₀	[1005-51-2]	bullvalene				
	$\Delta_{\text{fus}}H$		15.25	366.5		[1996DOM/HEA]
			71.8	298	C	[1981MAN/SUN]
C ₁₀ H ₁₀ Cl ₂ O ₃	[533-23-3]	(2,4-dichlorophenoxy)acetic acid, ethyl ester				
Δ_vH		(444–573)	72.6	459	A	[1987STE/MAL, 1999DYK/SVO]
C ₁₀ H ₁₀ Cl ₂ O ₃	[94-82-6]	4-(2,4-dichlorophenoxy)butyric acid				
	$\Delta_{\text{fus}}H$		35.0	395.5	DSC	[2005VEC/BRU]
	$\Delta_{\text{fus}}H$		38.42	391.4		[1991ACR]
	$\Delta_{\text{sub}}H$		124 ± 6	298	DSC	[2005VEC/BRU]
	$\Delta_{\text{sub}}H$	(356–391)	146 ± 1	374	TE	[2005VEC/BRU]
	$\Delta_{\text{sub}}H$	(356–391)	149 ± 2	298	TE	[2005VEC/BRU]
C ₁₀ H ₁₀ N ₂	[2243-62-1]	1,5-diaminonaphthalene				
	$\Delta_{\text{sub}}H$	(367–389)	118.5 ± 0.9	378	ME	[2010RIB/FER]
	$\Delta_{\text{sub}}H$	(367–389)	122.5 ± 0.9	298	ME	[2010RIB/FER]
		(345–371)	120.2 ± 0.7	298	GS	[2007VER/GEO]
C ₁₀ H ₁₀ N ₂	[479-27-6]	1,8-diaminonaphthalene				
$\Delta_{\text{fus}}H$			16.15	339.8	DSC	[2007VER/GEO]

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{sub}}H$	(314–338)	97.6 ± 0.7	326	ME	[2010RIB/FER]
	$\Delta_{\text{sub}}H$	(314–338)	99.0 ± 0.7	298	ME	[2010RIB/FER]
	$\Delta_{\text{sub}}H$	(304–335)	94.1 ± 0.4	298	GS	[2007VER/GEO]
	Δ_vH	(339–379)	79.6 ± 0.3	298	GS	[2007VER/GEO]
C ₁₀ H ₁₀ N ₂	[10199-67-4]	1-benzylpyrazole				
	Δ_vH		73.8 ± 2.0	298	C	[1999MO/YAN]
C ₁₀ H ₁₀ N ₂	[2379-55-7]	2,3-dimethylquinoxaline				
	$\Delta_{\text{fus}}H$		22.35	379.5	DSC	[2000MON/HIL2]
	$\Delta_{\text{sub}}H$	(294–308)	87.7 ± 0.4	301	ME	[2000MON/HIL2]
	$\Delta_{\text{sub}}H$		87.8 ± 0.4	298	ME	[2000MON/HIL2]
	$\Delta_{\text{sub}}H$		85.8 ± 1.8	298	C	[1996RIB/MOR]
C ₁₀ H ₁₀ N ₂	[6628-04-2]	4-aminoquinaldine				
	$\Delta_{\text{sub}}H$	(352–373)	112.1 ± 0.8	363	ME	[1998RIB/CAR]
	$\Delta_{\text{sub}}H$		115.3 ± 0.8	298		[1998RIB/CAR]
C ₁₀ H ₁₀ N ₂	[4238-71-5]	1-benzylimidazole				
	$\Delta_{\text{sub}}H$		102.1 ± 0.4	298	ME	[1999MO/YAN]
C ₁₀ H ₁₀ N ₂ O ₂	[6118-66-7]	3-dimethylaminophthalimide				
	$\Delta_{\text{sub}}H$	(392–431)	90.9	407	RG	[1987STE/MAL, 1956KLO]
C ₁₀ H ₁₀ N ₂ O ₂	[5432-74-6]	2,3-dimethylquinoxaline 1,4-dioxide				
	$\Delta_{\text{sub}}H$		124.4 ± 2.7	298	C	[2004RIB/GOM2]
C ₁₀ H ₁₀ N ₂ O ₅	[143248-63-9]	2,3-dihydro-3-[2-(nitrooxy)ethyl]-4H-1,3-benzoxazin-4-one				
	$\Delta_{\text{fus}}H$		23.0	326.3	DSC	[1996FON/ROS]
C ₁₀ H ₁₀ N ₄ O ₂ S	[68-35-9]	4-amino-N-(2-pyrimidinyl)benzene sulfonamide (sulfadiazine)				
	$\Delta_{\text{fus}}H$		44.3	532.7	DSC	[2003MAR/AVI, 2002MAR/GOM]
	$\Delta_{\text{fus}}H$		44.3	520.4		[1985MAR/WU]
	$\Delta_{\text{fus}}H$		31.2	538.7	DTA	[1971SUN/EIS]
C ₁₀ H ₁₀ O	[101-39-3]	2-methyl-3-phenyl-2-propenal				
	Δ_vH	(401–556)	59.3 ± 0.2	400	EB	[2002STE/CHI4]
	Δ_vH	(401–556)	56.3 ± 0.2	440	EB	[2002STE/CHI4]
	Δ_vH	(401–556)	53.4 ± 0.2	480	EB	[2002STE/CHI4]
	Δ_vH	(401–556)	50.5 ± 0.4	520	EB	[2002STE/CHI4]
	Δ_vH	(401–556)	47.7 ± 0.6	560	EB	[2002STE/CHI4]
	Δ_vH	(343–393)	71.5	358	A	[1987STE/MAL]
C ₁₀ H ₁₀ O	[122-57-6]	4-phenyl-3-buten-2-one				
	Δ_vH	(354–534)	58.5	369	A	[1987STE/MAL, 1947STU]
C ₁₀ H ₁₀ O	[529-34-0]	1-tetralone				
	Δ_vH	(284–324)	65.0 ± 0.3	298	GS	[1998VER4]
	Δ_vH	(388–535)	61.5	403	A	[1987STE/MAL]
C ₁₀ H ₁₀ O ₂	[103-26-4]	<i>trans</i> cinnamic acid, methyl ester				
	$\Delta_{\text{fus}}H$		33.1	309		[2002STE/CHI4]
	Δ_vH	(409–557)	59.9 ± 0.2	420	EB	[2002STE/CHI4]
	Δ_vH	(409–557)	56.9 ± 0.2	460	EB	[2002STE/CHI4]
	Δ_vH	(409–557)	53.8 ± 0.3	500	EB	[2002STE/CHI4]
	Δ_vH	(409–557)	50.5 ± 0.5	540	EB	[2002STE/CHI4]
	Δ_vH	(288–333)	62.4	303	A	[1987STE/MAL]
	Δ_vH	(350–536)	58.3	365	A	[1987STE/MAL, 1947STU]
C ₁₀ H ₁₀ O ₂	[6781-42-6]	1,3-diacetylbenzene				
	Δ_vH	(323–418)	43.2	338	A	[1987STE/MAL]

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₀ H ₁₀ O ₂	[1009-61-6] $\Delta_v H$	1,4-diacetylbenzene (388–431)	82.2	403	A	[1987STE/MAL]
C ₁₀ H ₁₀ O ₂	[120-58-1] $\Delta_v H$	isosafole (393–531)	59.4	408	A	[1987STE/MAL]
C ₁₀ H ₁₀ O ₂	[1199-77-5] $\Delta_v H$	α -methylcinnamic acid (398–561)	78.5	413	A	[1987STE/MAL, 1947STU]
C ₁₀ H ₁₀ O ₂	[1963-36-6] $\Delta_{\text{fus}} H$	4-methoxycinnamaldehyde	19.0	332.7		[2008TEM/ROU]
C ₁₀ H ₁₀ O ₂	[94-59-7] $\Delta_v H$	safrole (336–506)	54.6	351	A	[1987STE/MAL, 1947STU]
C ₁₀ H ₁₀ O ₂	[15844-05-0] $\Delta_v H$	4-carboxypentacyclo[4.3.0.0. ^{2,5} 0 ^{4,7}]nonane	82.0		C	[1984BEC/RUC]
C ₁₀ H ₁₀ O ₂	[93-91-4] $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$	1-phenyl-1,3-butanedione (278–300)	91.0 ± 0.6 83.7	298 289	ME V	[1992RIB/MON] [1987STE/MAL, 1959AIH]
C ₁₀ H ₁₀ O ₂	[28315-93-7] $\Delta_{\text{fus}} H$ $\Delta_{\text{sub}} H$	5-hydroxy- α -tetralone	33.67 118.5 ± 1.5	480.1 298	DSC C	[2009MAT/SOU3] [2009MAT/SOU3]
C ₁₀ H ₁₀ O ₂	[3470-50-6] $\Delta_{\text{fus}} H$ $\Delta_{\text{fus}} H$ $\Delta_{\text{sub}} H$	6-hydroxy- α -tetralone	13.02 18.98 117.5 ± 1.4	387.5 425.9 298	DSC C	[2009MAT/SOU3] [2009MAT/SOU3]
C ₁₀ H ₁₀ O ₂ S	[16192-08-8] $\Delta_{\text{sub}} H$	p-tolyl propadienyl sulfone	113 ± 2.5		B	[1969MAC/STE, 1970COX/PIL]
C ₁₀ H ₁₀ O ₂ S	[14027-53-3] $\Delta_{\text{sub}} H$	p-tolyl prop-1-ynyl sulfone	103.3 ± 2.5		B	[1969MAC/STE, 1970COX/PIL]
C ₁₀ H ₁₀ O ₂ S	[16192-07-7] $\Delta_{\text{sub}} H$	p-tolyl prop-2-ynyl sulfone	107.5 ± 2.5		B	[1969MAC/STE, 1970COX/PIL]
C ₁₀ H ₁₀ O ₃	[14737-91-8] $\Delta_{\text{sub}} H$	<i>cis</i> -2-methoxycinnamic acid (339–352)	121.7 ± 0.6	298	ME	[1999MON/HIL]
C ₁₀ H ₁₀ O ₃	[3943-97-3] $\Delta_{\text{fus}} H$	methyl 4-hydroxycinnamate	30.22	410.7	DSC	[2010PAN/SAR]
C ₁₀ H ₁₀ O ₃	[1011-54-7] $\Delta_{\text{sub}} H$	<i>trans</i> -2-methoxycinnamic acid (368–382)	128.8 ± 0.6	298	ME	[1999MON/HIL]
C ₁₀ H ₁₀ O ₃	[6099-03-2] $\Delta_{\text{fus}} H$	2-methoxycinnamic acid	32.54	458.7	DSC	[1994HUA/CHE]
C ₁₀ H ₁₀ O ₃	[17570-26-2] $\Delta_{\text{sub}} H$	<i>trans</i> -3-methoxycinnamic acid (353–367)	124.0 ± 0.9	298	ME	[1999MON/HIL]
C ₁₀ H ₁₀ O ₃	[6099-04-3] $\Delta_{\text{fus}} H$	3-methoxycinnamic acid	22.58	390.5	DSC	[1994HUA/CHE]
C ₁₀ H ₁₀ O ₃	[943-89-5] $\Delta_{\text{sub}} H$	<i>trans</i> -4-methoxycinnamic acid (369–383)	134.0 ± 1.0	298	ME	[1999MON/HIL]
C ₁₀ H ₁₀ O ₃	[830-09-1] $\Delta_{\text{us}} H$	4-methoxycinnamic acid	24.75	446.4		

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound					
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference	
		$\Delta_{\text{fus}}H$	2.49	461.9	DSC	[1994HUA/CHE]	
C ₁₀ H ₁₀ O ₃	[2879-20-1]	6-acetylbenzodioxan					
		$\Delta_{\text{fus}}H$	23.49	356	DSC	[2008MAT/SOU2]	
		$\Delta_{\text{sub}}H$	102.5 ± 1.1	298	C	[2008MAT/SOU2]	
C ₁₀ H ₁₀ O ₃	[24327-08-0]	<i>endo</i> -bicyclo[2.2.2]oct-5-ene-2,3-dicarboxylic anhydride					
		$\Delta_{\text{trs}}H$	18.19	382.9			
		$\Delta_{\text{fus}}H$	4.54	419.2	DSC	[1984WEI/LEF]	
C ₁₀ H ₁₀ O ₄	[635-67-6]	1,2-diacetoxybenzene (371–551)	62.9	386	A	[1987STE/MAL]	
C ₁₀ H ₁₀ O ₄	[1459-93-4]	dimethyl isophthalate					
		$\Delta_{\text{fus}}H$	25.3	341.2		[1993ACR]	
		$\Delta_{\text{sub}}H$	(295–309)	100.7 ± 0.2	302	ME	[1998ROU/JIM]
		$\Delta_{\text{sub}}H$		100.9 ± 0.2	298		[1998ROU/JIM]
		$\Delta_{\text{sub}}H$		100.7	298	C	[1998MAK/KAB]
		Δ_vH	(350–607)	77.2 ± 0.8	298	EB,IP	[1997STE/CHI2]
		Δ_vH	(393–550)	60.5	408	A,GS	[1987STE/MAL, 1963VOI]
C ₁₀ H ₁₀ O ₄	[131-11-3]	dimethyl phthalate					
		$\Delta_{\text{fus}}H$	16.95	274.2		[1998MAK/KAB]	
		$\Delta_{\text{fus}}H$	16.95	274.2		[1993ACR]	
		Δ_vH	(466–518)	61.5	481	EB	[1999ROH/MUS]
		Δ_vH		69.4 ± 0.1	365	C	[1998MAK/KAB]
		Δ_vH		72.5 ± 0.6	344	C	[1998MAK/KAB]
		Δ_vH		74.5 ± 0.3	326	C	[1998MAK/KAB]
		Δ_vH	(304–371)	78.7	319	A	[1987STE/MAL]
		Δ_vH	(371–547)	63.7	386	A	[1987STE/MAL]
		Δ_vH	(377–440)	68.6	409		[1969DAV/MAK]
C ₁₀ H ₁₀ O ₄	[120-61-6]	dimethyl terephthalate					
		$\Delta_{\text{fus}}H$	32.09	413.8		[1993ACR]	
		$\Delta_{\text{sub}}H$	(311–330)	103.8 ± 0.3	321	ME	[1998ROU/JIM]
		$\Delta_{\text{sub}}H$		104.6 ± 0.3	298		[1998ROU/JIM]
		$\Delta_{\text{sub}}H$	(373–413)	94.4	388		[1987STE/MAL]
		$\Delta_{\text{sub}}H$	(373–413)	88.3	393	GS	[1962KRA/BER]
		$\Delta_{\text{sub}}H$		105.3		C	[1998MAK/KAB]
		Δ_vH	(413–523)	62	428	A	[1987STE/MAL]
C ₁₀ H ₁₀ O ₄	[1135-24-6]	4-hydroxy-3-methoxycinnamic acid (ferulic acid)					
		$\Delta_{\text{fus}}H$	17.89	435.3	DSC	[1994HUA/CHE]	
		$\Delta_{\text{sub}}H$	(369–390)	132.4 ± 1.3	379		[2006CHE/OJA]
C ₁₀ H ₁₀ O ₄	[635-51-8]	(RS)-phenylsuccinic acid					
		$\Delta_{\text{fus}}H$	37.37	440.1	DSC	[2006PRO/RAS]	
C ₁₀ H ₁₀ O ₄	[4036-30-0]	(S)-phenylsuccinic acid					
		$\Delta_{\text{fus}}H$	41.84	446.9	DSC	[2006PRO/RAS]	
C ₁₀ H ₁₀ O ₄	[1135-24-6]	3-(4-hydroxy-3-methoxyphenyl)-2-propenoic acid (ferulic acid)					
		$\Delta_{\text{fus}}H$	33.34	444.6		[2008MOT/QUE]	
C ₁₀ H ₁₁ ClN ₂ O ₄	[310412-18-1]	ethyl (2-chloromethyl-2,3-dihydro-5 <i>H</i> -oxazolo[3,2- <i>a</i>]-pyrimidin-5-one)-6-carboxylate					
		$\Delta_{\text{trs}}H$	5.38	379.7			
		$\Delta_{\text{fus}}H$	10.77	413.4	DSC	[2000CHA/SOS]	

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₀ H ₁₁ ClO ₃	[na] $\Delta_{\text{fus}}H$	(dl) 2-(2-chloro-3-methylphenoxy)propionic acid	30.54	391.5		[1976LEC/COL]
C ₁₀ H ₁₁ ClO ₃	[na] $\Delta_{\text{fus}}H$	(d) 2-(2-chloro-3-methylphenoxy)propionic acid	22.18	359.5		[1976LEC/COL]
C ₁₀ H ₁₁ F ₃ N ₂ O	[2164-17-2] $\Delta_{\text{fus}}H$	N,N-dimethyl-N'-[3-(trifluoromethyl)-phenyl]urea	29.82	434.1	DSC	[1991ACR, 1990DON/DRE]
C ₁₀ H ₁₁ F ₃ N ₂ O ₃ S	[47000-92-0] $\Delta_{\text{fus}}H$	N-[4-methyl-3-[[trifluoromethyl)sulfonyl]amino]phenyl]acetamide	40.47	455.7	DSC	[1990DON/DRE]
C ₁₀ H ₁₁ N	[1195-98-8] Δ_vH Δ_vH	α,α -dimethylbenzylcyanide (284–323) (284–323)	60.3 ± 0.6 60.6 ± 0.6	303 298	GS GS	[2000VER] [2000VER]
C ₁₀ H ₁₁ N	[769-68-6] Δ_vH	α -ethylbenzylcyanide (283–313)	64.3 ± 0.6	298	GS	[2000VER]
C ₁₀ H ₁₁ N	[2571-52-0] $\Delta_{\text{sub}}H$	2,4,6-trimethylbenzotrile	82.9 ± 1.6	298	C	[1991ACR/TUC]
C ₁₀ H ₁₁ NO	[2904-59-8] $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$	2,4,6-trimethylbenzotrile N-oxide	87.5 ± 0.5 87.9 ± 1.9 84.7 ± 1.8 77.5 ± 3.7	314 298 319 298	C C ME C	[1993ACR/SEV] [1993ACR/SEV] [1992ACR/SIM] [1991ACR/TUC]
C ₁₀ H ₁₁ NO	[1128-85-4] $\Delta_{\text{sub}}H$	3-amino-1-phenyl-but-2-enone	109.4 ± 2.1	298	C	[1993RIB/RIB]
C ₁₀ H ₁₁ NO ₂	[1563-87-7] $\Delta_{\text{sub}}H$	N-phenyldiacetamide	90.0 ± 0.8	298	C	[1965WAD, 1970COX/PIL]
C ₁₀ H ₁₁ NO ₂ S	[6958-78-7] $\Delta_{\text{sub}}H$	N-benzoylthiocarbamic O-ethyl ester	112.2 ± 1.3	298	C	[2004RIB/SAN2]
C ₁₀ H ₁₁ NO ₃	[2571-54-2] $\Delta_{\text{sub}}H$	2,4,6-trimethoxybenzotrile	112.6 ± 2.0	298	C	[1991ACR/TUC]
C ₁₀ H ₁₁ NO ₃	[34295-85-7] $\Delta_{\text{fus}}H$	N-salicylidene- β -alanine	28.5	408		[1996DOM/HEA]
C ₁₀ H ₁₁ NO ₃	[2623-33-8] $\Delta_{\text{fus}}H$	N-[4-(acetyloxy)phenyl] acetamide	30.97	427.5		[1990BHA/LAL]
C ₁₀ H ₁₁ NO ₄	[2904-59-8] $\Delta_{\text{sub}}H$	2,4,6-trimethoxybenzotrile N-oxide	91.9 ± 1.9	298	C	[1991ACR/TUC]
C ₁₀ H ₁₁ N ₃ O	[5809-38-1] $\Delta_{\text{sub}}H$	3,5-dimethyl-1-phenyl-4-nitrosopyrazole	89.7 ± 2.0	298	C	[2001RIB/FER]
C ₁₀ H ₁₁ N ₃ O ₂	[10495-38-2] $\Delta_{\text{sub}}H$	3-dimethylamino-6-aminophthalimide (434–459)	108.8	446.5		[1987STE/MAL, 1956KLO]
C ₁₀ H ₁₁ N ₃ O ₃ S	[723-46-6] $\Delta_{\text{fus}}H$ $\Delta_{\text{fus}}H$	4-amino-N-(5-methyl-3-isoxazolyl)benzene sulfonamide (sulfamethoxazole)	33.8 35.26	440.7 440	DSC	[2003MAR/AVI, 2002MAR/GOM] [1998ISS/ELA]
C ₁₀ H ₁₁ N ₅ O ₃	[134287-59-5] $\Delta_{\text{fus}}H$	3,9-dihydro-3-[(2-hydroxyethoxy)methyl]-9-oxo-5H-imidazo[1,2-a]pyrine	43.33	520.6	DSC	[1999ZIE/GOL]
C ₁₀ H ₁₂	[1755-01-7] $\Delta_{\text{fus}}H$ $\Delta_{\text{fus}}H$	<i>endo</i> dicyclopentadiene	9.66 2.22	216 304.8		[1996DOM/HEA]

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound						
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference		
			8.04	216.1				
			$\Delta_{\text{fus}}H$	1.79	304.7		[1997SMI/LEB]	
			Δ_vH	(350–446)	43.6	365	A	[1987STE/MAL]
C ₁₀ H ₁₂	[933-60-8]	<i>exo</i> -dicyclopentadiene						
			$\Delta_{\text{fus}}H$	7.11	189.8		[1997SMI/LEB]	
C ₁₀ H ₁₂	[2234-20-0]	2,4-dimethylstyrene						
		(307–453)	Δ_vH	50.0	322	A	[1987STE/MAL, 1947STU]	
C ₁₀ H ₁₂	[2039-89-6]	2,5-dimethylstyrene						
		(302–453)	Δ_vH	48.1	317	A	[1987STE/MAL, 1947STU]	
C ₁₀ H ₁₂	[2039-93-2]	α -ethylstyrene						
		(274–313)	Δ_vH	52.0 ± 0.3	294	GS	[1999VER/EBE]	
		(274–313)	Δ_vH	51.8 ± 0.3	298	GS	[1999VER/EBE]	
C ₁₀ H ₁₂	[7564-63-8]	1-ethyl-2-vinylbenzene						
		(363–413)	Δ_vH	46.3	378	A	[1987STE/MAL]	
C ₁₀ H ₁₂	[7525-62-4]	1-ethyl-3-vinylbenzene						
		(343–453)	Δ_vH	49.6	358	A	[1987STE/MAL]	
C ₁₀ H ₁₂	[3454-07-7]	1-ethyl-4-vinylbenzene						
		(341–448)	Δ_vH	48.4	356	A	[1987STE/MAL]	
C ₁₀ H ₁₂	[119-64-2]	1,2,3,4-tetrahydronaphthalene (tetralin)						
			$\Delta_{\text{fus}}H$	12.45	237.4		[1996DOM/HEA]	
		(465–580)	Δ_vH	44.1	480		[1992LEE/DEM]	
		(311–481)	Δ_vH	51.1	326	A	[1987STE/MAL]	
			Δ_vH	41.3 ± 0.1	498	C	[1985NAT/VIS]	
			Δ_vH	37.6 ± 0.1	552	C	[1985NAT/VIS]	
			Δ_vH	35.7 ± 0.1	567	C	[1985NAT/VIS]	
			Δ_vH	33.9 ± 0.1	585	C	[1985NAT/VIS]	
			Δ_vH	32.0 ± 0.1	604	C	[1985NAT/VIS]	
		(331–437)	Δ_vH	52.1	346		[1984KAT/HAR]	
		(367–479)	Δ_vH	48.6	382		[1947STU]	
C ₁₀ H ₁₂	[3451-55-6]	cyclodeca-1,2,6,7-tetraene						
			$\Delta_{\text{sub}}H$	73.0 ± 0.4	298	C	[1991LUK/TIM]	
C ₁₀ H ₁₂ ClNO ₂	[2950-21-0]	isopropyl-3-chlorophenylcarbamate						
			$\Delta_{\text{fus}}H$	17.75	313.9		[1991ACR]	
C ₁₀ H ₁₂ ClN ₃ O ₂	[15271-41-7]	5-chloro-6-[[[(methylamino)carbonyl]oxy]imino]bicyclo[2.2.1]-heptane-2-carbonitrile						
			$\Delta_{\text{fus}}H$	26.07	431.6	DSC	[1990DON/DRE]	
C ₁₀ H ₁₂ Cl ₄ NOPS	[21844-03-1]	P-chloromethyl-N-(1-methylethyl)amidothiophosphonic acid, O-(2,4,6-trichlorophenyl) ester						
		(323–363)	Δ_vH	78.3	338	A	[1987STE/MAL]	
C ₁₀ H ₁₂ N ₂	[827-36-1]	2-dimethylamino-2-phenylacetone nitrile						
		(298–338)	Δ_vH	62.8 ± 0.4		GS	[1997WEL/VER]	
C ₁₀ H ₁₂ N ₂	[5465-29-2]	2-propylbenzimidazole						
			$\Delta_{\text{us}}H$	1.5	385.8			
			$\Delta_{\text{fus}}H$	NA	431.6	DSC	[2004RIB/RIB]	
		(344–364)	$\Delta_{\text{sub}}H$	109.4 ± 1.2	298	ME	[2004RIB/RIB]	
C ₁₀ H ₁₂ N ₂	[5851-43-4]	2-isopropylbenzimidazole						
			$\Delta_{\text{us}}H$	1.0	410.4			
			$\Delta_{\text{fus}}H$	NA	507.9	DSC	[2004RIB/RIB]	

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{sub}}H$		109.9 ± 2.7	298	C	[2004RIB/RIB]
C ₁₀ H ₁₂ N ₂ O ₂	[30764-27-3] $\Delta_{\text{sub}}H$	acetylglycine anilide (362–365)	122.1	363.5	A	[1987STE/MAL, 1955AIH]
C ₁₀ H ₁₂ N ₂ O ₃	[52-43-7] $\Delta_{\text{fus}}H$	allobarbital	32.31	442.6		[1986CHU/DEM]
C ₁₀ H ₁₂ N ₂ O ₃	[38423-62-0] $\Delta_{\text{fus}}H$ $\Delta_{\text{fus}}H$	2-ethoxyisonitrosoacetanilide	23.0 27.8	405 434.6	DTA	[1996DOM/HEA] [1982CUE/SOL]
C ₁₀ H ₁₂ N ₂ O ₃	[17122-74-6] $\Delta_{\text{fus}}H$ $\Delta_{\text{fus}}H$	4-ethoxyisonitrosoacetanilide	7.6 7.6	490 491.2	DTA	[1996DOM/HEA] [1982CUE/SOL]
C ₁₀ H ₁₂ N ₂ O ₃ S	[25057-89-0] $\Delta_{\text{fus}}H$	3-(1-methylethyl)-(1 <i>H</i>)-2,1,3-benzothiadiazin-4(3 <i>H</i>)-one 2,2-dioxide	21.77	412.5	DSC	[1990DON/DRE]
C ₁₀ H ₁₂ N ₂ O ₄	[84592-41-6] $\Delta_{\text{fus}}H$	(2-hydroxyethyl)[3-[(hydroxyimino)methyl]phenyl carbamic acid	28.9	508.3	DTA	[1982CUE/SOL]
C ₁₀ H ₁₂ N ₂ O ₄	[3056-17-5] $\Delta_{\text{fus}}H$	2,3'-didehydro-3'-deoxythymidine (stavudine)	26.91	443.2		[2000GAN/BOG]
C ₁₀ H ₁₂ N ₂ O ₄ S	[138517-12-1] $\Delta_{\text{fus}}H$	(4-nitrophenyl)-2-(methylthio)ethyl carbamate	31.27	349.4	DSC	[1993TIE/FRA]
C ₁₀ H ₁₂ N ₂ O ₅	[88-85-7] $\Delta_{\text{fus}}H$	2-sec-butyl-4,6-dinitrophenol	21.81	313.7	DSC	[1990DON/DRE]
C ₁₀ H ₁₂ N ₂ S	[na] $\Delta_{\text{fus}}H$	N-allyl-N-phenylthiourea	27.61	375		[1996DOM/HEA]
C ₁₀ H ₁₂ N ₃ O ₃ PS ₂	[86-50-0] $\Delta_{\text{fus}}H$	S-(3,4-dihydro-4-oxobenzo[d][1,2,3]-triazin-3-ylmethyl) O,O-dimethylphosphorodithioate	27.76	345.3	DSC	[1991ACR, 1990DON/DRE]
C ₁₀ H ₁₂ O	[104-46-1] Δ_vH	anethole	61.9	298	GC	[2002VAN/PAR]
C ₁₀ H ₁₂ O	[25679-28-1] Δ_vH	<i>cis</i> anethole (333–363)	68.7	348	A	[1987STE/MAL]
C ₁₀ H ₁₂ O	[4180-23-8] Δ_vH	<i>trans</i> anethole (333–363)	78.3	348	A	[1987STE/MAL]
C ₁₀ H ₁₂ O	[140-67-0] Δ_vH	estragole (325–488)	56.3	340	A	[1987STE/MAL]
C ₁₀ H ₁₂ O	[2142-64-5] Δ_vH Δ_vH	2'-ethylacetophenone (363–397) (293–423)	52.8 U23.7	378 368	A	[1987STE/MAL] [1968KHO/BYK]
C ₁₀ H ₁₂ O	[937-30-4] Δ_vH Δ_vH	4'-ethylacetophenone (294–368) (293–423)	42.2 39.8	309 368	A	[1987STE/MAL] [1968KHO/BYK]
C ₁₀ H ₁₂ O	[122-03-2] Δ_vH	4-isopropylbenzaldehyde (cuminal) (331–505)	55.3	346	A	[1987STE/MAL, 1947STU]
C ₁₀ H ₁₂ O	[5445-77-2] Δ_vH	2-methyl-3-phenylpropanal (333–373)	59.1	348	A	[1987STE/MAL]
C ₁₀ H ₁₂ O	[5337-93-9] Δ_vH	4'-methylpropiophenone (332–512)	52.6	347	A	[1987STE/MAL, 1947STU]

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₀ H ₁₂ O	[5459-40-5] $\Delta_v H$	4-vinylphenetole (337–498)	59.2	352	A	[1987STE/MAL, 1947STU]
C ₁₀ H ₁₂ O	[1712-69-2] $\Delta_{\text{fus}} H$	4-methoxy- α -methylstyrene	19.07	309.2		[1999VER6]
	$\Delta_{\text{sub}} H$		81.2 ± 0.4	298		[1999VER6]
	$\Delta_v H$	(308–343)	60.6 ± 0.3	326	GS	[1999VER6]
	$\Delta_v H$	(308–343)	62.1 ± 0.3	298	GS	[1999VER6]
C ₁₀ H ₁₂ O ₂	[3674-77-9] $\Delta_{\text{sub}} H$	2-phenyl-2-methyl-1,3-dioxolane (293–324)	81.9 ± 0.5	308	T	[1995VER/DOG]
C ₁₀ H ₁₂ O ₂	[103-45-7] $\Delta_v H$	acetic acid, phenethyl ester (283–318)	67.4	298	A	[1987STE/MAL]
	$\Delta_v H$	(422–506)	52.2	437	A	[1987STE/MAL]
C ₁₀ H ₁₂ O ₂	[31508-44-8] $\Delta_v H$	methyl 2-phenylpropionate (284–318)	61.8 ± 0.7	301	GS	[1999VER8]
	$\Delta_v H$	(284–318)	62.0 ± 0.7	298	GS	[1999VER8]
C ₁₀ H ₁₂ O ₂	[97-53-0] $\Delta_v H$	4-allyl-2-methoxyphenol (eugenol)	66.3	298	GC	[2002VAN/PAR]
	$\Delta_v H$	(395–527)	57.7	410	A	[1987STE/MAL]
	$\Delta_v H$	(285–333)	66.1	300	ME	[1987STE/MAL, 1959SCO/DOU]
	$\Delta_v H$	(351–526)	60.3	366		[1947STU]
C ₁₀ H ₁₂ O ₂	[122-63-4] $\Delta_v H$	benzyl propionate (298–378)	59.0	313	A	[1987STE/MAL]
C ₁₀ H ₁₂ O ₂	[501-19-9] $\Delta_v H$	5-allyl-2-methoxyphenol (345–527)	61.4	371	A	[1987STE/MAL]
C ₁₀ H ₁₂ O ₂	[97-54-1] $\Delta_v H$	2-methoxy-4-(1-propenyl)phenol (isoeugenol) (359–540)	60.7	374		[1957DYK/SEP]
C ₁₀ H ₁₂ O ₂	[5912-86-7] $\Delta_v H$	<i>cis</i> isoeugenol (373–403)	69.7	388	A	[1987STE/MAL]
C ₁₀ H ₁₂ O ₂	[5932-68-3] $\Delta_v H$	<i>trans</i> isoeugenol (363–420)	69.1	378	A	[1987STE/MAL]
C ₁₀ H ₁₂ O ₂	[na] $\Delta_{\text{fus}} H$	acetophenone ethylene glycol ketal	25.2	333.6		[1995VER/DOG]
C ₁₀ H ₁₂ O ₂	[16108-50-2] $\Delta_{\text{fus}} H$	2-acetyl-3,5-dimethylphenol	1.36	333.2	DTA	[1989SAL/ABA]
		Note: Reported enthalpy of fusion is too small, and the published enthalpy and entropy of fusion data are internally inconsistent				
C ₁₀ H ₁₂ O ₂	[101-97-3] $\Delta_v H$	phenylacetic acid, ethyl ester (333–433)	60.7	298	GC	[2005HOS/GRY]
	$\Delta_v H$	(288–328)	63.9 ± 0.4	308	GS	[1999VER8]
	$\Delta_v H$	(288–328)	64.5 ± 0.4	298	GS	[1999VER8]
	$\Delta_v H$	(393–500)	54	408	A	[1987STE/MAL]
C ₁₀ H ₁₂ O ₂	[2315-68-6] $\Delta_v H$	propylbenzoate (359–458)	60.2	379	BG	[1988KAT2]
	$\Delta_v H$	(359–458)	52.7	440	BG	[1988KAT2]
	$\Delta_v H$	(327–504)	53.8	342	A	[1987STE/MAL, 1947STU]
C ₁₀ H ₁₂ O ₂	[2930-05-4] $\Delta_v H$	[(phenylmethoxy)methyl]oxirane	71.0 ± 0.4			[1987VAN/KAC]

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₀ H ₁₂ O ₂	[2529-36-4]	2,3,6-trimethylbenzoic acid				
	$\Delta_{\text{sub}}H$	(314–336)	104.4 ± 0.2	298	ME	[1987COL/JIM2]
	$\Delta_{\text{sub}}H$	(314–336)	103.6 ± 0.2	325	ME	[1987COL/JIM2]
C ₁₀ H ₁₂ O ₂	[480-63-7]	2,4,6-trimethylbenzoic acid				
	$\Delta_{\text{sub}}H$	(316–340)	103.6 ± 0.3	298	ME	[1987COL/JIM2]
	$\Delta_{\text{sub}}H$	(316–340)	102.5 ± 0.3	328	ME	[1987COL/JIM2]
C ₁₀ H ₁₂ O ₂	[1076-47-7]	2,3,4-trimethylbenzoic acid				
	$\Delta_{\text{sub}}H$	(329–351)	109.3 ± 0.3	298	ME	[1987COL/JIM2]
	$\Delta_{\text{sub}}H$	(329–351)	108.2 ± 0.3	340	ME	[1987COL/JIM2]
C ₁₀ H ₁₂ O ₂	[2437-66-3]	2,3,5-trimethylbenzoic acid				
	$\Delta_{\text{sub}}H$	(320–338)	106.7 ± 0.3	298	ME	[1987COL/JIM2]
	$\Delta_{\text{sub}}H$	(320–338)	105.7 ± 0.3	329	ME	[1987COL/JIM2]
C ₁₀ H ₁₂ O ₂	[528-90-5]	2,4,5-trimethylbenzoic acid				
	$\Delta_{\text{sub}}H$	(324–346)	109.6 ± 0.5	298	ME	[1987COL/JIM2]
	$\Delta_{\text{sub}}H$	(324–346)	108.3 ± 0.5	335	ME	[1987COL/JIM2]
C ₁₀ H ₁₂ O ₂	[1076-88-6]	3,4,5-trimethylbenzoic acid				
	$\Delta_{\text{sub}}H$	(340–359)	111.0 ± 0.5	298	ME	[1987COL/JIM2]
	$\Delta_{\text{sub}}H$	(340–359)	109.3 ± 0.5	350	ME	[1987COL/JIM2]
C ₁₀ H ₁₂ O ₂	[2438-05-3]	4-propylbenzoic acid				
	$\Delta_{\text{fus}}H$		3.4	301		
	$\Delta_{\text{fus}}H$		23.3	422		[1985PRI/PUC]
	$\Delta_{\text{sub}}H$	(331–347)	109.1 ± 0.8	298	ME	[2004MON/ALM]
C ₁₀ H ₁₂ O ₂	[2438-04-2]	2-isopropylbenzoic acid				
	$\Delta_{\text{sub}}H$	(300–320)	100.2 ± 0.4	310	ME	[1987COL/JIM]
	$\Delta_{\text{sub}}H$		101.0 ± 0.4	298		[1987COL/JIM]
C ₁₀ H ₁₂ O ₂	[5651-47-8]	3-isopropylbenzoic acid				
	$\Delta_{\text{sub}}H$	(300–316)	103.3 ± 0.3	308	ME	[1987COL/JIM]
	$\Delta_{\text{sub}}H$		104.1 ± 0.3	298		[1987COL/JIM]
C ₁₀ H ₁₂ O ₂	[536-66-3]	4-isopropylbenzoic acid				
	$\Delta_{\text{sub}}H$	(316–334)	99.0 ± 0.3	310	ME	[1987COL/JIM]
	$\Delta_{\text{sub}}H$		99.0 ± 0.3	298		[1987COL/JIM]
C ₁₀ H ₁₂ O ₂	[1821-12-1]	4-phenylbutyric acid				
	$\Delta_{\text{fus}}H$		19.5	324.2	DSC	[2001MON/HIL]
	$\Delta_{\text{sub}}H$	(309–323)	112.4 ± 0.8	316	ME	[2001MON/HIL]
C ₁₀ H ₁₂ O ₂	[527-17-3]	2,3,5,6-tetramethyl-1,4-benzoquinone				
	$\Delta_{\text{fus}}H$		18.39 ± 0.1	384.1	DSC	[2004ROJ/FOR]
	$\Delta_{\text{fus}}H$		18.54 ± 0.1	384.8	HFC	[2004ROJ/FOR]
C ₁₀ H ₁₂ O ₂	[527-17-3]	2,3,5,6-tetramethyl-1,4-benzoquinone				
	$\Delta_{\text{sub}}H$		93.2 ± 1.2	298	DSC	[2004ROJ/FOR]
	[490-91-5]	2-isopropyl-5-methyl-1,4-benzoquinone (thymoquinone)				
$\Delta_{\text{fus}}H$		18.16	323.2		[2003PAG/BEN]	
C ₁₀ H ₁₂ O ₂ S	[32228-15-2]	p-tolyl trans-prop-1-enyl sulfone				
$\Delta_{\text{sub}}H$			83.7 ± 2.1		B	[1969MAC/STE, 1970COX/PIL]
C ₁₀ H ₁₂ O ₂ S	[3112-87-6]	p-tolyl prop-2-enyl sulfone				
$\Delta_{\text{sub}}H$			95.8 ± 2.9		B	[1969MAC/STE, 1970COX/PIL]
C ₁₀ H ₁₂ O ₂ S	[67605-02-1]	p-tolyl isopropenyl sulfone				

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
			88.7 ± 2.5		B	[1969MAC/MCN, 1969MAC/MCN2, 1977PED/RYL]
C ₁₀ H ₁₂ O ₃	[6192-44-5] $\Delta_v H$	acetic acid, (2-phenoxyethyl) ester (355–533)	56.8	370	A	[1987STE/MAL, 1947STU]
C ₁₀ H ₁₂ O ₃	[94-13-3] $\Delta_{\text{fus}} H$ $\Delta_{\text{fus}} H$	propyl 4-hydroxybenzoate	27.2 27.99	369.3 369.2		[1999GIO/BET] [1990MAN/AHU]
	$\Delta_{\text{sub}} H$	(319–333)	123.7 ± 0.6	298	GS	[2005PER/ROD]
	$\Delta_v H$		79.1		TGA	[2002CHA/DOL]
	$\Delta_v H$		76.5		TGA	[2001CHA/DOL]
C ₁₀ H ₁₂ O ₃	[3759-31-7] $\Delta_{\text{fus}} H$	(dl) 3-hydroxy-3-phenylbutyric acid	19.66	330		[1991CHI/BRA]
C ₁₀ H ₁₂ O ₃	[na] $\Delta_{\text{fus}} H$	(d) 3-hydroxy-3-phenylbutyric acid	22.59	357		[1991CHI/BRA]
C ₁₀ H ₁₂ O ₃	[4919-33-9] $\Delta_{\text{fus}} H$	4-ethoxyphenylacetic acid	23.0	360.2		[1991ACR]
C ₁₀ H ₁₂ O ₃	[6342-77-4] $\Delta_{\text{fus}} H$	3-(2-methoxyphenyl)propionic acid	25.33	360.5	DSC	[2001MON/HIL4]
	$\Delta_{\text{sub}} H$	(331–347)	116.0 ± 0.4	339	ME	[2001MON/HIL4]
	$\Delta_{\text{sub}} H$	(331–347)	117.8 ± 1.4	298	ME	[2001MON/HIL4]
C ₁₀ H ₁₂ O ₃	[129-29-9] $\Delta_{\text{fus}} H$ $\Delta_{\text{fus}} H$	3-(4-methoxyphenyl)propionic acid	29.57 28.5	376.1 376.9	DSC	[2001MON/HIL4] [1991ACR]
	$\Delta_{\text{sub}} H$	(341–357)	122.3 ± 0.3	349	ME	[2001MON/HIL4]
	$\Delta_{\text{sub}} H$	(341–357)	124.5 ± 1.7	298	ME	[2001MON/HIL4]
C ₁₀ H ₁₂ O ₃	[5438-19-7] $\Delta_{\text{us}} H$ (liq cryst) $\Delta_{\text{us}} H$ (liq cryst) $\Delta_{\text{us}} H$ (liq cryst-liq)	4-propoxybenzoic acid	7.95 16.74 2.51	394.2 419.9 426.7		[1967HER]
C ₁₀ H ₁₂ O ₄	[999-21-3] $\Delta_v H$	maleic acid, diallyl ester (392–426)	77.7	407	A	[1987STE/MAL]
C ₁₀ H ₁₂ O ₄	[20765-04-2] $\Delta_{\text{fus}} H$	2,5-diethoxy-1,4-benzoquinone	28.7	459.3	DSC	[1996KEE/VAN]
C ₁₀ H ₁₂ O ₅	[490-64-2] $\Delta_{\text{fus}} H$	2,4,5-trimethoxybenzoic acid	31.15	417.9	DSC	[2003HUA, 2005HUA/TAN]
C ₁₀ H ₁₂ O ₅	[118-41-2] $\Delta_{\text{fus}} H$	3,4,5-trimethoxybenzoic acid	29.9	444.5	DSC	[2003HUA, 2005HUA/TAN]
	$\Delta_{\text{sub}} H$	(354–372)	127.9 ± 0.8	363	ME	[2001ROU/JIM2]
	$\Delta_{\text{sub}} H$		131.2 ± 0.8	298	ME	[2001ROU/JIM2]
C ₁₀ H ₁₃ Br	[2437-76-5] $\Delta_v H$	2-bromo-4-isopropyltoluene (400–510)	50.2	415	A	[1987STE/MAL, 1970DYK/VAN, 1999DYK/SVO]
C ₁₀ H ₁₃ Br	[4478-10-8] $\Delta_v H$	3-bromo-4-isopropyltoluene (400–510)	48.3	415	A	[1987STE/MAL, 1970DYK/VAN, 1999DYK/SVO]

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₀ H ₁₃ BrO	[54514-31-7] Δ_vH	2-bromophenyl isobutyl ether (293–323)	64.0 ± 0.2	298	GS	[2005STR/SPO]
C ₁₀ H ₁₃ BrO	[223564-75-8] Δ_vH	3-bromophenyl isobutyl ether (290–323)	65.3 ± 0.2	298	GS	[2005STR/SPO]
C ₁₀ H ₁₃ BrO	[30752-23-9] Δ_vH	4-bromophenyl isobutyl ether (293–333)	66.8 ± 0.6	298	GS	[2005STR/SPO]
C ₁₀ H ₁₃ Cl	[4395-79-3] Δ_vH	2-chloro-4-isopropyltoluene (400–490)	49.3	415	A	[1987STE/MAL, 1970DYK/VAN, 1999DYK/SVO]
C ₁₀ H ₁₃ Cl	[15146-00-6] Δ_vH	3-chloro-4-isopropyltoluene (400–490)	47.1	415	A	[1987STE/MAL, 1970DYK/VAN]
C ₁₀ H ₁₃ ClN ₂ O ₂	[19937-59-8] $\Delta_{\text{fus}}H$	N'-(3-chloro-4-methoxyphenyl)-N,N-dimethylurea	27.48	399.2	DSC	[1990DON/DRE]
C ₁₀ H ₁₃ ClN ₂ O ₃ S	[94-20-2] $\Delta_{\text{fus}}H$	1-(4-chlorobenzenesulfonyl)-3-propylurea	25.7	401	DSC	[2006WAS/HOL, 2008WAS/HOL]
C ₁₀ H ₁₃ ClN ₆	[32889-48-8] $\Delta_{\text{fus}}H$	2-((4-chloro-6-(cyclopropylamino)-1,3,5-triazin-2-yl)amino)-2-methylpropanenitrile	22.51	438.5	DSC	[1990DON/DRE]
C ₁₀ H ₁₃ ClO	[4446-91-7] Δ_vH	2-chloroethyl α -methylbenzyl ether (335–508)	54.8	350	A	[1987STE/MAL, 1947STU, 1999DYK/SVO]
C ₁₀ H ₁₃ ClO ₃	[58498-77-4] Δ_vH	diethylene glycol 4-chlorophenyl ether (450–523)	75.9	465	A	[1987STE/MAL, 1999DYK/SVO]
C ₁₀ H ₁₃ ClO ₃	[94-81-5] $\Delta_{\text{fus}}H$	4-(4-chloro-2-methylphenoxy)butanoic acid	32.02	373.4	DSC	[1990DON/DRE]
C ₁₀ H ₁₃ Cl ₂ O ₂ P	[18351-36-5] Δ_vH	4-tert-butyl phenyl dichlorophosphate (369–572)	59.6	384		[1947STU]
C ₁₀ H ₁₃ Cl ₃ NOPS	[18361-88-1] Δ_vH	P-chloromethyl-N-(1-methylethyl)amidothiophosphonic acid, O-(2,4-dichlorophenyl) ester (323–368)	93.1	345	A	[1987STE/MAL, 1999DYK/SVO]
C ₁₀ H ₁₃ NO	[6935-65-5] Δ_vH	N,N-dimethyl- <i>m</i> -toluamide (374–405)	29.9	390		[1969DAV/MAK2]
C ₁₀ H ₁₃ NO ₂	[62-44-2] $\Delta_{\text{fus}}H$	4'-ethoxyacetanilide (phenacetin)	30	409.6	DSC	[2009VEC/TOM]
	$\Delta_{\text{fus}}H$		28.75	408.3	DSC	[2009PEN/ESC]
	$\Delta_{\text{fus}}H$		34.1	407.4	DSC	[2006WAS/HOL, 2008WAS/HOL]
	$\Delta_{\text{fus}}H$		21.4	410.2	DSC	[2004VEC/CAT]
	$\Delta_{\text{fus}}H$		31.25	407.2	DSC	[1990MAN/AHU]
	$\Delta_{\text{sub}}H$		120 ± 3	298	Vap+Fus	[2009VEC/TOM]
	$\Delta_{\text{sub}}H$	(312–387)	115.5 ± 2.4		C,ME	[1972WIE, 1987STE/MAL]
	Δ_vH		79 ± 1	459	TGA	[2009VEC/TOM]
	Δ_vH		82 ± 1	476	TGA	[2009VEC/TOM]
Δ_vH	(463–533)	82.6	478	A	[1987STE/MAL]	
C ₁₀ H ₁₃ NO ₂	[943-15-7] Δ_vH	2-nitro-4-isopropyltoluene (370–415)	67.7	385	A	[1987STE/MAL]
C ₁₀ H ₁₃ NO ₂	[35480-94-5] Δ_vH	3-nitro-4-isopropyltoluene (330–430)	54	345	A	[1987STE/MAL]
C ₁₀ H ₁₃ NO ₂	[1886-57-3]	2-nitro-1- <i>tert</i> -butylbenzene				

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(278–323)	64.8 ± 0.6	301	GS	[2000VER/HEI]
	$\Delta_v H$	(278–323)	65.0 ± 0.6	298	GS	[2000VER/HEI]
C ₁₀ H ₁₃ NO ₂	[94-12-2]	propyl 4-aminobenzoate				
	$\Delta_{\text{fus}} H$		20.54	347.1		[1991ACR]
C ₁₀ H ₁₃ NO ₂	[1202-25-1]	methyl <i>p</i> -N,N-dimethylaminobenzoate				
	$\Delta_{\text{fus}} H$		26.07	371.8		[1991ACR]
C ₁₀ H ₁₃ NO ₂	[5532-90-1]	propyl N-phenylcarbamate				
	$\Delta_{\text{fus}} H$		21.08	331		[1971PRI]
C ₁₀ H ₁₃ NO ₂	[122-42-9]	isopropyl phenylcarbamate				
	$\Delta_{\text{fus}} H$		19.37	359.5		[1991ACR]
C ₁₀ H ₁₃ NO ₂	[2425-10-7]	3,4-dimethylphenyl methylcarbamate				
	$\Delta_{\text{fus}} H$		24.97	350.8		[1991ACR]
C ₁₀ H ₁₃ NO ₂ S ₂	[949171-65-7]	N-theonylthiocarbamic-O-butyl ester				
	$\Delta_{\text{fus}} H$		23.89	364.3	DSC	[2007RIB/MON]
	$\Delta_{\text{sub}} H$		147.5 ± 1.9	298	C	[2007RIB/MON]
C ₁₀ H ₁₃ NO ₄	[6988-21-2]	2-(1,3-dioxolan-2-yl)phenyl methylcarbamate				
	$\Delta_{\text{fus}} H$		23.82	387.2	DSC	[1990DON/DRE]
C ₁₀ H ₁₃ N ₅ O ₃	[na]	2-acetylamino-9-[(2-hydroxyethoxy)methyl]-9 <i>H</i> -purine				
	$\Delta_{\text{fus}} H$		54.92	454.2		[1995KRI/VES]
C ₁₀ H ₁₃ N ₅ O ₃	[na]	9-[(2-acetoxyethoxy)methyl]-2-amino-9 <i>H</i> -purine				
	$\Delta_{\text{fus}} H$		42.69	408.2		[1995KRI/VES]
C ₁₀ H ₁₃ N ₅ O ₄	[110104-37-5]	2-acetylamino-9-[(2-hydroxyethoxy)methyl]-1,9-dihydro-6 <i>H</i> -purin-6-one				
	$\Delta_{\text{fus}} H$		53.83	490.2	DSC	[1995KRI/VES]
C ₁₀ H ₁₃ N ₅ O ₄	[102728-64-3]	2-amino-9-[(2-acetoxyethoxy)methyl]-1,9-dihydro-6 <i>H</i> -purin-6-one				
	$\Delta_{\text{fus}} H$		49.9	515.2	DSC	[1995KRI/VES]
C ₁₀ H ₁₃ N ₅ O ₄	[30516-87-1]	3'-azido-2,3'-dideoxythymidine (zidovudine)				
	$\Delta_{\text{fus}} H$		33.03	359.8	DSC	[2003ARA/STO]
C ₁₀ H ₁₄	[104-51-8]	butylbenzene				
	$\Delta_{\text{fus}} H$		11.22	185.3		[1996DOM/HEA]
	$\Delta_v H$	(343–501)	47.4 ± 0.2	350	EB	[2002STE/CHI, 2006VER]
	$\Delta_v H$	(343–501)	43.5 ± 0.2	410	EB	[2002STE/CHI]
	$\Delta_v H$	(343–501)	40.6 ± 0.4	450	EB	[2002STE/CHI]
	$\Delta_v H$	(343–501)	37.5 ± 0.7	490	EB	[2002STE/CHI]
	$\Delta_v H$		50.8	298		[1994RUZ/ZAB]
	$\Delta_v H$	(243–403)	53.5	258		[1993KAS/MOK]
	$\Delta_v H$		48.0 ± 0.1	343	C	[1982SVO/CHA]
	$\Delta_v H$		46.8 ± 0.1	358	C	[1982SVO/CHA]
	$\Delta_v H$		46.0 ± 0.1	368	C	[1982SVO/CHA]
	$\Delta_v H$		50.1	298		[1971WIL/ZWO]
	$\Delta_v H$	(374–454)	45.2	389		[1965LIN/FRI, 1984BOU/FRI]
	$\Delta_v H$	(369–463)	45.7	384	A	[1987STE/MAL, 1949FOR/NOR]
C ₁₀ H ₁₄	[135-98-8]	<i>(dl)</i> sec-butylbenzene				
	$\Delta_v H$	(335–491)	45.7 ± 0.2	340	EB	[2002STE/CHI]
	$\Delta_v H$	(335–491)	43.2 ± 0.2	380	EB	[2002STE/CHI]
	$\Delta_v H$	(335–491)	40.6 ± 0.3	420	EB	[2002STE/CHI]
	$\Delta_v H$	(335–491)	37.8 ± 0.5	460	EB	[2002STE/CHI]
	$\Delta_v H$		48.1	298		[1994RUZ/ZAB]
	$\Delta_v H$	(243–373)	50.6	258		[1993KAS/MOK]

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(384–448)	42.8	399	A	[1987STE/MAL]
	$\Delta_v H$		49.5	298		[1971WIL/ZWO]
	$\Delta_v H$	(368–448)	44.0	375		[1949FOR/NOR]
C₁₀H₁₄	[98-06-6]	<i>tert</i> -butylbenzene				
	$\Delta_{\text{fus}}H$		8.4	215.3	DSC,AC	[2009CHI/STE]
	$\Delta_{\text{fus}}H$		8.41	215		[1996DOM/HEA]
	$\Delta_v H$	(332–486)	45.3 ± 0.2	340	EB	[2002STE/CHI]
	$\Delta_v H$	(332–486)	42.6 ± 0.2	380	EB	[2002STE/CHI]
	$\Delta_v H$	(332–486)	39.9 ± 0.3	420	EB	[2002STE/CHI]
	$\Delta_v H$	(332–486)	37.0 ± 0.5	460	EB	[2002STE/CHI]
	$\Delta_v H$	(278–308)	47.8 ± 0.4	293	GS	[1998VER]
	$\Delta_v H$		47.5 ± 0.4	298		[1998VER]
	$\Delta_v H$		47.6	298		[1994RUZ/ZAB]
	$\Delta_v H$	(368–444)	43.1	383	A	[1987STE/MAL]
	$\Delta_v H$		47.7	298		[1974KUS/SAI]
	$\Delta_v H$		49.1	298		[1971WIL/ZWO]
	$\Delta_v H$	(357–443)	43.7	372		[1949FOR/NOR]
C₁₀H₁₄	[135-01-3]	1,2-diethylbenzene				
	$\Delta_v H$		52.8	298		[1971WIL/ZWO]
	$\Delta_v H$	(369–464)	46.0	384	A	[1987STE/MAL, 1949FOR/NOR]
C₁₀H₁₄	[141-93-5]	1,3-diethylbenzene				
	$\Delta_v H$		52.5	298		[1971WIL/ZWO]
	$\Delta_v H$	(368–457)	45.8	383	A	[1987STE/MAL, 1949FOR/NOR]
C₁₀H₁₄	[105-05-5]	1,4-diethylbenzene				
	$\Delta_v H$		52.5	298		[1971WIL/ZWO]
	$\Delta_v H$	(369–464)	45.8	384	A	[1987STE/MAL, 1949FOR/NOR]
C₁₀H₁₄	[933-98-2]	1,2-dimethyl-3-ethylbenzene				
	$\Delta_v H$	(344–497)	49.7	359	A	[1987STE/MAL]
	$\Delta_v H$		54.9	298		[1971WIL/ZWO]
C₁₀H₁₄	[934-80-5]	1,2-dimethyl-4-ethylbenzene				
	$\Delta_v H$	(340–493)	48.9	355	A	[1987STE/MAL]
	$\Delta_v H$		53.9	298		[1971WIL/ZWO]
C₁₀H₁₄	[2870-04-0]	1,3-dimethyl-2-ethylbenzene				
	$\Delta_v H$	(341–493)	48.8	356	A	[1987STE/MAL]
	$\Delta_v H$		53.9	298		[1971WIL/ZWO]
	$\Delta_v H$	(299–461)	48.6	314		[1947STU]
C₁₀H₁₄	[874-41-9]	1,3-dimethyl-4-ethylbenzene				
	$\Delta_v H$	(339–492)	48.5	354	A	[1987STE/MAL]
	$\Delta_v H$		53.3	298		[1971WIL/ZWO]
C₁₀H₁₄	[934-74-7]	1,3-dimethyl-5-ethylbenzene				
	$\Delta_v H$	(336–487)	48.0	351	A	[1987STE/MAL]
	$\Delta_v H$		52.4	298		[1971WIL/ZWO]
	$\Delta_v H$	(295–456)	47.5	310		[1947STU]
C₁₀H₁₄	[1758-88-9]	1,4-dimethyl-2-ethylbenzene				
	$\Delta_v H$	(338–490)	48.0	353	A	[1987STE/MAL]
	$\Delta_v H$		52.6	298		[1971WIL/ZWO]
	$\Delta_v H$	(299–440)	48.7	313		[1947STU]
C₁₀H₁₄	[538-93-2]	isobutylbenzene				
	$\Delta_v H$		48.0	298		[1994RUZ/ZAB]
	$\Delta_v H$	(373–447)	43.2	388	A	[1987STE/MAL]
	$\Delta_v H$		49.5	298		[1971WIL/ZWO]

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₀ H ₁₄	$\Delta_v H$	(360–447)	43.8	375		[1949FOR/NOR]
	[527-84-4] $\Delta_v H$	2-isopropyltoluene (354–453)	44.4	369	A	[1987STE/MAL, 1959MCD/SHR, 1984BOU/FRI]
	$\Delta_v H$		50.6	298		[1971WIL/ZWO]
C ₁₀ H ₁₄	[535-77-3] $\Delta_v H$	3-isopropyltoluene (351–450)	44.7	366	A	[1987STE/MAL, 1959MCD/SHR, 1984BOU/FRI]
	$\Delta_v H$		50.0	298		[1971WIL/ZWO]
C ₁₀ H ₁₄	[99-87-6] $\Delta_{\text{fus}} H$	4-isopropyltoluene	9.67	204.2		[1996DOM/HEA]
	$\Delta_v H$	(333–443)	49.2	298	GC	[2005HOS/GRY]
	$\Delta_v H$		48.9	298		[1994RUZ/ZAB]
	$\Delta_v H$	(380–452)	44.0	395	A	[1987STE/MAL, 1959MCD/SHR, 1984BOU/FRI]
C ₁₀ H ₁₄	$\Delta_v H$		50.3	298		[1971WIL/ZWO]
	[1074-17-5] $\Delta_v H$	2-propyltoluene (337–488)	48.0	352	A	[1987STE/MAL]
	$\Delta_v H$		52.7	298		[1971WIL/ZWO]
	C ₁₀ H ₁₄	[1074-43-7] $\Delta_v H$	3-propyltoluene (334–485)	47.8	349	A
$\Delta_v H$			52.1	298		[1971WIL/ZWO]
C ₁₀ H ₁₄	[1074-55-1] $\Delta_v H$	4-propyltoluene (335–487)	47.6	350	A	[1987STE/MAL]
	$\Delta_v H$		51.9	298		[1971WIL/ZWO]
C ₁₀ H ₁₄	[488-23-3] $\Delta_{\text{fus}} H$	1,2,3,4-tetramethylbenzene	11.23	265.4		[1996DOM/HEA]
	$\Delta_v H$		54.0	298		[1994RUZ/ZAB]
	$\Delta_v H$		52.6 ± 0.2	298	C	[1994SAB/TAB, 1990YAW/YAN]
	$\Delta_v H$	(352–509)	50.7	367	A	[1987STE/MAL]
	$\Delta_v H$		57.2	298		[1971WIL/ZWO]
	$\Delta_v H$	(316–477)	55.7	331		[1947STU]
C ₁₀ H ₁₄	[527-53-7] $\Delta_{\text{fus}} H$	1,2,3,5-tetramethylbenzene	12.93	248.6		[1996DOM/HEA]
	$\Delta_v H$		53.2	298		[1994RUZ/ZAB]
	$\Delta_v H$		52.0 ± 0.2	298	C	[1994SAB/TAB]
	$\Delta_v H$	(348–502)	50.0	363	A	[1987STE/MAL]
	$\Delta_v H$		55.8	298		[1971WIL/ZWO]
	$\Delta_v H$	(314–471)	58.9	329		[1947STU]
C ₁₀ H ₁₄	[95-93-2] $\Delta_{\text{fus}} H$	1,2,4,5-tetramethylbenzene	20.88	352.4		[1996DOM/HEA]
	$\Delta_{\text{sub}} H$		71.7 ± 0.3	298	C	[1994SAB/TAB]
	$\Delta_{\text{sub}} H$	(263–277)	74.6 ± 0.3	298	ME	[1989COL/JIM]
	$\Delta_{\text{sub}} H$	(318–348)	71.3	333	A	[1947BAL/DEN]
	$\Delta_{\text{sub}} H$		72.4	298	H	[1947BAL/DEN, 1993CHI/HOS]
	$\Delta_v H$	(363–381)	47.7 ± 0.3	375	DM	[2001BLO/VAN]
	$\Delta_v H$	(353–500)	49.4	368	A	[1987STE/MAL]
	[na] $\Delta_v H$	spirocyclopropane-1,6-tricyclo[3.2.1.0 ^{2,4}]octane	47.8 ± 0.1	298	C	[1998KOL/PIM, 1996VAR/PAS]

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₀ H ₁₄ Cl ₂ NO ₂ PS	[299-85-4] $\Delta_{\text{fus}}H$	O-(2,4-dichlorophenyl) O-methyl-(1-methylethyl) phosphoramidothioate	29.25	321.5	DSC	[1990DON/DRE]
C ₁₀ H ₁₄ NO ₅ PS	[56-38-2] $\Delta_{\text{fus}}H$	O,O-diethyl O-4-nitrophenyl phosphorothioate (parathion)	15.72	278.1	DSC	[1991ACR, 1990DON/DRE]
	$\Delta_{\text{sub}}H$	(298–318)	100.6	308		[1979SPE/SHO, 1983SPE/CLI]
	Δ_vH	(293–433)	93.4 (sub)	308	A	[1987STE/MAL, 1999DYK/SVO]
C ₁₀ H ₁₄ NO ₅ PS	[3270-86-8] Δ_vH	phosphorothioic acid, O,O'-diethyl-S-(4-nitrophenyl)	(313–366) 75.9	328	A	[1987STE/MAL, 1999DYK/SVO]
C ₁₀ H ₁₄ NO ₅ PS	[597-88-6] Δ_vH	phosphorothioic acid, O,S-diethyl-O'-(4-nitrophenyl)	(332–364) 75.1	347	A	[1987STE/MAL, 1999DYK/SVO]
C ₁₀ H ₁₄ NO ₆ P	[311-45-5] Δ_vH	O,O-diethyl-O-(4-nitrophenyl) phosphate	(273–422) 87.9	288	A	[1987STE/MAL]
C ₁₀ H ₁₄ N ₂	[22083-74-5] Δ_vH	(<i>dl</i>) nicotine	(406–520) 53.3	421	A	[1987STE/MAL]
	[54-11-5] Δ_vH Δ_vH	(<i>l</i>) nicotine	 63.9 ± 2.1 46.1	298 448	CGC	[2009LIP/HAN] [1934GOR]
C ₁₀ H ₁₄ N ₂	[494-52-0] Δ_vH	3-(2S)-2-piperidinylpyridine	(373–523) 49.5	448		[1934GOR]
C ₁₀ H ₁₄ N ₂ O	[120-22-9] $\Delta_{\text{sub}}H$	4-diethylaminonitrosobenzene	107.9 ± 3.7	298	C	[1998RIB/MAT2]
C ₁₀ H ₁₄ N ₂ O ₂ S	[na] $\Delta_{\text{sub}}H$	N,N-diethyl-N'-furoylthiourea	132.0 ± 3.5	298	C	[2002RIB/RIB]
C ₁₀ H ₁₄ N ₂ O ₄	[88381-75-3] $\Delta_{\text{us}}H$	2,2-dinitroadamantane	16.76	362.2		
	$\Delta_{\text{fus}}H$		5.06	491.2		[1990FRI/DOG]
	$\Delta_{\text{sub}}H$		(278–317) 96.4 ± 1.4	298	T	[1990FRI/DOG]
C ₁₀ H ₁₄ N ₄ O ₂	[2850-41-1] $\Delta_{\text{fus}}H$	8-propyltheophylline	33.3	534.3	DSC	[1991ACR, 1989GON/KRA]
C ₁₀ H ₁₄ N ₄ O ₂	[2850-40-0] $\Delta_{\text{fus}}H$	8-isopropyltheophylline	34.4	569.3	DSC	[1991ACR, 1989GON/KRA]
C ₁₀ H ₁₄ N ₄ O ₃	[603-00-9] $\Delta_{\text{fus}}H$ (I)	(R,S)-3,7-dihydro-7-(2-hydroxypropyl)-1,3-dimethyl-1 <i>H</i> -pyrine-2,6-dione (proxiphylline)	24.7	408.2		
	$\Delta_{\text{fus}}H$ (II)		20.8	389.2		[2000GRI/AUE]
C ₁₀ H ₁₄ N ₄ O ₄	[479-18-5] $\Delta_{\text{fus}}H$	7-(2,3-dihydroxypropyl)-3,7-dihydro-1,3-dimethyl-1 <i>H</i> -purine-2,6-dione (dyphylline)	39.3	435.7		[1999GRI/AUE]
C ₁₀ H ₁₄ O	[3180-09-4] Δ_vH	2-butylphenol	(403–533) 55.1	418	A	[1987STE/MAL, 1975ARR/MEL]
	Δ_vH		(382–520) 52.9	398		[1953STA/MUL]
	Δ_vH		(382–520) 51.0	423		[1953STA/MUL]
	Δ_vH		(382–520) 47.0	473		[1953STA/MUL]
C ₁₀ H ₁₄ O	[89-72-5] Δ_vH	2-sec-butylphenol	(451–513) 52.1	466	A,GS,EB	[1987STE/MAL, 1964HAN/HAR]
C ₁₀ H ₁₄ O	[88-18-6] Δ_vH	2- <i>tert</i> -butylphenol	(289–329) 62.6 ± 0.2	309	GS	[1999VER2]

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$		63.2 ± 0.2	298		[1999VER2]
	$\Delta_v H$	(409–467)	74.1	424	EB	[1990NES/NAZ]
	$\Delta_v H$	(409–465)	52.9	424		[1986TSV/NAZ]
	$\Delta_v H$	(353–498)	54.9	368	A	[1987STE/MAL]
	$\Delta_v H$	(330–507)	55.6	348		[1953STA/MUL]
	$\Delta_v H$	(330–507)	53.9	373		[1953STA/MUL]
	$\Delta_v H$	(330–507)	51.0	423		[1953STA/MUL]
	$\Delta_v H$	(330–507)	47.0	473		[1953STA/MUL]
C₁₀H₁₄O	[4074-43-5]	3-butylphenol				
	$\Delta_v H$	(396–533)	62.5	411	A	[1987STE/MAL]
	$\Delta_v H$	(396–533)	56.6	398		[1953STA/MUL]
	$\Delta_v H$	(396–533)	54.4	423		[1953STA/MUL]
	$\Delta_v H$	(396–533)	49.9	473		[1953STA/MUL]
C₁₀H₁₄O	[585-34-2]	3- <i>tert</i> -butylphenol				
	$\Delta_{\text{sub}} H$		88.9 ± 0.5	298	C	[1999RIB/MAT2]
	$\Delta_{\text{sub}} H$	(278–319)	86.0 ± 0.5	298	GS	[1999VER2]
	$\Delta_{\text{sub}} H$	(266–299)	70.7	281	A	[1987STE/MAL]
	$\Delta_v H$	(320–348)	69.1 ± 0.8	334	GS	[1999VER2]
	$\Delta_v H$	(320–348)	71.3 ± 0.8	298	GS	[1999VER2]
	$\Delta_v H$	(391–524)	62.4	406	A	[1987STE/MAL]
	$\Delta_v H$	(391–524)	56.6	398		[1953STA/MUL]
	$\Delta_v H$	(391–524)	54.4	423		[1953STA/MUL]
	$\Delta_v H$	(391–524)	49.9	473		[1953STA/MUL]
C₁₀H₁₄O	[1638-22-8]	4-butylphenol				
	$\Delta_v H$	(395–653)	61.7	410	A	[1987STE/MAL]
	$\Delta_v H$	(357–529)	57.6	373		[1953STA/MUL]
	$\Delta_v H$	(357–529)	56.6	398		[1953STA/MUL]
	$\Delta_v H$	(357–529)	54.4	423		[1953STA/MUL]
	$\Delta_v H$	(357–529)	49.9	473		[1953STA/MUL]
C₁₀H₁₄O	[99-71-8]	4- <i>sec</i> -butylphenol				
	$\Delta_v H$	(344–516)	59.0	359	A	[1987STE/MAL, 1947STU]
C₁₀H₁₄O	[98-54-4]	4- <i>tert</i> -butylphenol				
	$\Delta_{\text{fus}} H$		14.52	373.2		[1972INO/LIA]
	$\Delta_{\text{sub}} H$		89.4 ± 2.5	298	C	[1999RIB/MAT2]
	$\Delta_{\text{sub}} H$	(293–334)	85.0 ± 0.5	313	GS	[1999VER2]
	$\Delta_{\text{sub}} H$	(293–334)	85.9 ± 0.5	298	GS	[1999VER2]
	$\Delta_{\text{sub}} H$	(280–304)	84.3	292	A	[1987STE/MAL, 1960AIH]
	$\Delta_v H$		67.9 ± 1.0	298	C	[1999RIB/MAT2]
	$\Delta_v H$	(471–525)	54.3	486	A,GS,EB	[1987STE/MAL, 1947STU, 1964HAN/HAR]
	$\Delta_v H$	(346–523)	59.6	348		[1953STA/MUL]
	$\Delta_v H$	(346–523)	57.6	373		[1953STA/MUL]
	$\Delta_v H$	(346–523)	56.6	398		[1953STA/MUL]
	$\Delta_v H$	(346–523)	54.4	423		[1953STA/MUL]
	$\Delta_v H$	(346–523)	49.9	473		[1953STA/MUL]
C₁₀H₁₄O	[1126-79-0]	butyl phenyl ether				
	$\Delta_v H$	(391–483)	48.9	406	A	[1987STE/MAL, 1949DRE/SHR, 1984BOU/FRI]
C₁₀H₁₄O	[4371-48-6]	3-isopropyl-2-methylphenol				
	$\Delta_v H$	(365–516)	60.2	380	EB	[1969LAM/PER]
C₁₀H₁₄O	[1740-97-2]	4-isopropyl-2-methylphenol				
	$\Delta_v H$	(382–503)	59.8	397	EB	[1969LAM/PER]

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₀ H ₁₄ O	[499-75-2]	5-isopropyl-2-methylphenol (carvacrol)				
	$\Delta_v H$		68.2	298	GC	[2002VAN/PAR]
	$\Delta_v H$	(387–512)	59.4	402	EB	[1969LAM/PER]
	$\Delta_v H$	(343–510)	56.5	358	A	[1987STE/MAL, 1947STU]
C ₁₀ H ₁₄ O	[3228-04-4]	6-isopropyl-2-methylphenol				
	$\Delta_v H$	(371–499)	54.5	386	EB	[1969LAM/PER]
C ₁₀ H ₁₄ O	[1197-34-8]	3,5-diethylphenol				
	$\Delta_v H$	(387–521)	54.3	402	A	[1987STE/MAL, 1955VON/GEB]
C ₁₀ H ₁₄ O	[4167-74-2]	4-isobutylphenol				
	$\Delta_v H$	(345–510)	58.1	360	A	[1987STE/MAL, 1947STU]
C ₁₀ H ₁₄ O	[527-35-5]	2,3,5,6-tetramethylphenol				
	$\Delta_v H$	(381–522)	51.2	396	A	[1987STE/MAL, 1955VON/GEB]
C ₁₀ H ₁₄ O	[89-83-8]	2-isopropyl-5-methylphenol (thymol)				
	$\Delta_{\text{fus}} H$		22.01	324.2		[1991CHI/BRA]
	$\Delta_{\text{sub}} H$	(273–295)	75.1	284	A	[1987STE/MAL, 1960AIH]
	$\Delta_{\text{sub}} H$	(293–323)	89.1 ± 4.5	303	HSA	[1975CHI]
	$\Delta_{\text{sub}} H$	(229–312)	U 69.0	270	TGA	[1971ASH]
	$\Delta_{\text{sub}} H$	(273–313)	91.2 ± 4.1		TE	[1970COX/PIL, 1960JON, 1947BAL]
	$\Delta_{\text{sub}} H$	(283–323)	91.5	298		[1957SHE/BRY, 1987STE/MAL]
	$\Delta_v H$	(333–433)	70.9	298	GC	[2005HOS/GRY]
	$\Delta_v H$		68.7	298	GC	[2002VAN/PAR]
	$\Delta_v H$	(393–433)	70.5	298	CGC	[1995CHI/HOS]
	$\Delta_v H$	(381–514)	58.4	396	A	[1987STE/MAL]
	$\Delta_v H$	(339–514)	63.2	373		[1953STA/MUL]
	$\Delta_v H$	(339–514)	58.4	398		[1953STA/MUL]
	$\Delta_v H$	(339–514)	55.2	423		[1953STA/MUL]
	$\Delta_v H$	(339–514)	52.8	448		[1953STA/MUL]
	$\Delta_v H$	(339–514)	51.5	473		[1953STA/MUL]
$\Delta_v H$	(337–505)	54.9	352		[1947STU]	
C ₁₀ H ₁₄ O	[22327-39-5]	<i>(dl)</i> carvone				
	$\Delta_v H$	(330–501)	55	345	A	[1987STE/MAL, 1947STU]
C ₁₀ H ₁₄ O	[na]	<i>(l)</i> -carvone				
	$\Delta_{\text{fus}} H$		11.55	247.7		[1996GAL/BOU]
C ₁₀ H ₁₄ O	[2244-16-8]	<i>(+)</i> -carvone				
	$\Delta_v H$	(323–433)	58.2	298	GC	[2005HOS/GRY]
C ₁₀ H ₁₄ O	[18486-69-6]	<i>(-)</i> -myrtenal				
	$\Delta_v H$	(323–423)	55.1	298	GC	[2005HOS/GRY]
C ₁₀ H ₁₄ O	[1585-06-4]	4-ethylphenetole				
	$\Delta_v H$	(321–481)	54.3	336	A	[1987STE/MAL, 1947STU]
C ₁₀ H ₁₄ O	[22545-12-6]	2-(2-ethylphenyl)ethanol				
	$\Delta_v H$	(420–653)	59.5	435	A	[1987STE/MAL]
C ₁₀ H ₁₄ O	[22545-13-7]	2-(4-ethylphenyl)ethanol				
	$\Delta_v H$	(420–653)	59.1	435	A	[1987STE/MAL]
C ₁₀ H ₁₄ O	[536-60-7]	4-isopropylbenzyl alcohol				
	$\Delta_v H$	(347–520)	59.7	362	A	[1987STE/MAL, 1947STU]
C ₁₀ H ₁₄ O	[7384-80-7]	2-methyl-3-phenyl-1-propanol				
	$\Delta_v H$	(343–393)	71.9	358	A	[1987STE/MAL]

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₀ H ₁₄ O	[3299-05-6]	(1-ethoxyethyl)benzene				
	$\Delta_v H$	(286–318)	52.4 ± 0.2	302	GS	[2001VER/HEI]
	$\Delta_v H$	(286–318)	52.6 ± 0.2	298	GS	[2001VER/HEI]
C ₁₀ H ₁₄ O	[700-58-3]	2-adamantanone				
	$\Delta_{\text{trs}} H$	(5–310)	7.63	216.4	AC	[2006BAR/BLO]
	$\Delta_{\text{fus}} H$		11.77	557.5	DSC	[2006BAR/BLO]
	$\Delta_{\text{sub}} H$		66.4 ± 0.3	298	C	[2006BAR/BLO]
	$\Delta_{\text{sub}} H$	(280–333)	66.3 ± 0.8	298	ME	[2006BAR/BLO]
	$\Delta_{\text{sub}} H$		76.1 ± 1.5	298		[2002MIR/LEB, 2006BAR/BLO]
	$\Delta_{\text{sub}} H$		80.3 ± 2.5	298	BG	[1978ARO/STE]
C ₁₀ H ₁₄ O	[935-67-1]	(1-methoxy-1-methylethyl)-benzene				
	$\Delta_v H$	(278–313)	53.0 ± 0.5	296	GS	[2001HEI/VER]
	$\Delta_v H$	(278–313)	52.9 ± 0.5	298	GS	[2001HEI/VER]
C ₁₀ H ₁₄ O ₂	[4026-05-5]	1,2-dihydroxy-3- <i>tert</i> -butylbenzene				
	$\Delta_v H$	(334–384)	70.1 ± 0.8	359	GS	[2000VER/SCH]
	$\Delta_v H$	(334–384)	73.5 ± 0.8	298	GS	[2000VER/SCH]
C ₁₀ H ₁₄ O ₂	[98-29-3]	1,2-dihydroxy-4- <i>tert</i> -butylbenzene				
	$\Delta_{\text{fus}} H$		15.1	330.4		[2000VER/SCH]
	$\Delta_{\text{sub}} H$		98.7 ± 0.9	313	GS	[2000VER/SCH]
	$\Delta_{\text{sub}} H$		99.2 ± 0.9	298	GS	[2000VER/SCH]
	$\Delta_{\text{sub}} H$		99.3 ± 1.4	298	C	[1984CAR]
C ₁₀ H ₁₄ O ₂	[439-516]	(439–516)	96.5 ± 2.8	298	EB	[1997STE/CHI2]
	[1948-33-0]	2- <i>tert</i> -butyl-1,4-dihydroxybenzene				
	$\Delta_{\text{fus}} H$		27.74	350.9		[1999VER7]
	$\Delta_{\text{sub}} H$	(333–368)	101.2 ± 1.3	351	GS	[1999VER7]
C ₁₀ H ₁₄ O ₂	[13331-20-9]	1,3-dihydroxy-2-butylbenzene				
		$\Delta_v H$	(413–469)	75.3	428	A,GC
C ₁₀ H ₁₄ O ₂	[2785-87-7]	2-methoxy-4-propylphenol				
	$\Delta_v H$	(373–413)	78.0	388	A	[1987STE/MAL]
C ₁₀ H ₁₄ O ₂	[na]	<i>tert</i> -butylcatechol (isomer not specified)				
	$\Delta_v H$	(421–466)	58.2	443		[1965GAK/BAB]
C ₁₀ H ₁₄ O ₂	[490-06-2]	6-methyl-3-isopropyl-1,2-dihydroxybenzene				
	$\Delta_{\text{sub}} H$		96.6 ± 0.9	298	C	[1984CAR]
C ₁₀ H ₁₄ O ₂	[18523-34-7]	1,1-dimethoxy-2-phenylcyclopropane				
	$\Delta_v H$	(278–313)	63.7 ± 0.6	298	GS	[2002VER]
C ₁₀ H ₁₄ O ₂	[4316-35-2]	acetophenone dimethyl ketal				
	$\Delta_v H$	(268–303)	54.0 ± 0.8	298	GS	[2002VER]
	$\Delta_v H$	(268–303)	55.0 ± 1.3	286	GS	[1995VER/DOG]
C ₁₀ H ₁₄ O ₂	[na]	(1S)-(+)-camphorquinone				
	$\Delta_{\text{fus}} H$		6.07	473.2		[1992ELS/PRA]
		Note: Fusion enthalpy seems low, compound may have an unmeasured phase transition at a lower temperature]				
C ₁₀ H ₁₄ O ₃	[707-07-3]	trimethyl orthobenzoate				
	$\Delta_v H$	(294–333)	59.9 ± 0.4	298	GS	[2002VER]
	$\Delta_v H$	(294–333)	58.6 ± 0.4		GS	[1995RAK/VER2]

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₀ H ₁₄ O ₃	[595-31-3]	(d)-camphoric anhydride				
	$\Delta_{\text{trs}}H$		19.5	404.4		
	$\Delta_{\text{fus}}H$		5.65	493.6	DSC	[1984WEI/LEF]
	$\Delta_{\text{trs}}H$		29	406		
C ₁₀ H ₁₄ O ₃	[76-32-4]	(dl)-camphoric anhydride				
	$\Delta_{\text{trs}}H$		17.31	374.5		
	$\Delta_{\text{fus}}H$		5.72	493.9	DSC	[1984WEI/LEF]
	$\Delta_{\text{trs}}H$		24.0	375		
C ₁₀ H ₁₄ O ₃	[na]	(racemic) 3-(2-methylphenoxy)propane-1,2-diol				
	$\Delta_{\text{fus}}H$		32.2	343.8	DSC	[2008BRE/BRE]
C ₁₀ H ₁₄ O ₃	[52153-44-3]	(S)- 3-(2-methylphenoxy)propane-1,2-diol				
	$\Delta_{\text{fus}}H$		34.4	364.2	DSC	[2008BRE/BRE]
C ₁₀ H ₁₄ O ₅	[na]	allyl[(1-allyloxycarbonyl)ethyl] carbonate				
	Δ_vH	(353–503)	61.9	368	A	[1987STE/MAL]
C ₁₀ H ₁₄ O ₈	[na]	(dl) dimethyl diacetyltartrate				
	$\Delta_{\text{fus}}H$		25.94	355.2		[1991CHI/BRA]
C ₁₀ H ₁₄ O ₈	[na]	(d) dimethyl diacetyltartrate				
	$\Delta_{\text{fus}}H$		29.29	377.2		[1991CHI/BRA]
C ₁₀ H ₁₅ Br	[768-90-1]	1-bromoadamantane				
	$\Delta_{\text{trs}}H$	(6–320)	0.001	31	AC	
	$\Delta_{\text{trs}}H$	(6–320)	1.39	282.3	AC	
	$\Delta_{\text{trs}}H$	(290–430)	7.42	309.9	DSC	
	$\Delta_{\text{fus}}H$	(290–430)	3.97	391.8	DSC	[2005BAZ/BLO]
	$\Delta_{\text{trs}}H$		0.88	279		
	$\Delta_{\text{trs}}H$		6.93	310.5		
	$\Delta_{\text{fus}}H$		3.83	396.5		[1977CLA/KNO]
	$\Delta_{\text{sub}}H$ (form II)	(288–310)	71.6 ± 1.1	299	ME	[2005BAZ/KAB]
C ₁₀ H ₁₅ Cl	[935-56-8]	1-chloroadamantane				
	$\Delta_{\text{trs}}H$		6.01	244.2		
	$\Delta_{\text{fus}}H$		4.87	442.5		[1977CLA/KNO]
C ₁₀ H ₁₅ Cl	[7346-41-0]	2-chloroadamantane				
	$\Delta_{\text{trs}}H$		0.47	227		
	$\Delta_{\text{trs}}H$		8.3	242	DSC	[1988PAR/KAW]
	$\Delta_{\text{sub}}H$		61.5 ± 0.8	298	ME	[2002ABB/CAS]
C ₁₀ H ₁₅ Cl ₃ OS	[76633-71-1]	2,3,3-trichloro-2-propenethioic acid, O-heptyl ester				
	Δ_vH	(433–483)	72.7		GC	[1980PIT/KIS]
C ₁₀ H ₁₅ F	[768-92-3]	1-fluoroadamantane				
	$\Delta_{\text{fus}}H$		1.5	221.6		[1991KAW/GIL]
C ₁₀ H ₁₅ I	[768-93-4]	1-iodoadamantane				
	$\Delta_{\text{trs}}H$		2.14	211		
	$\Delta_{\text{fus}}H$		10.22	347		[1977CLA/KNO]
C ₁₀ H ₁₅ N	[6310-21-0]	2-tert-butylaniline				
	Δ_vH	(279–318)	62.7 ± 0.4	298	GS	[2000VER3]

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₀ H ₁₅ N	[579-66-8]	2,6-diethylaniline				
	$\Delta_v H$	(284–328)	69.5 ± 0.6	306		[2000VER3]
			65.9 ± 0.6	298		[2000VER3]
C ₁₀ H ₁₅ N	[2051-53-8]	5-isopropyl-2-methylaniline				
	$\Delta_v H$	(360–386)	72.0	373	A	[1987STE/MAL]
C ₁₀ H ₁₅ N	[537-46-2]	N- α -dimethylphenethylamine				
	$\Delta_v H$	(270–304)	52.8	285	A	[1987STE/MAL]
C ₁₀ H ₁₅ N	[1126-78-9]	N-butylaniline				
	$\Delta_v H$	(413–643)	55.6	428	A	[1987STE/MAL]
C ₁₀ H ₁₅ N	[91-66-7]	N,N-diethylaniline				
	$\Delta_v H$	(343–493)	54.5	358	A	[1987STE/MAL]
C ₁₀ H ₁₅ NO	[na]	2-(dimethylamino)-1-phenylethanone				
	$\Delta_v H$	(293–333)	69.7 ± 0.5	298	GS	[1994WEL/VER]
C ₁₀ H ₁₅ NO	[103-62-8]	4-(butylamino)phenol				
	$\Delta_v H$	(464–511)	71.2	478	A	[1987STE/MAL]
C ₁₀ H ₁₅ NO	[55658-55-4]	(<i>dl</i>)-carvoxime				
	$\Delta_{\text{fus}} H$		17.03	365.1	DTA	[1981CHI/GAR]
	$\Delta_{\text{sub}} H$	(324–343)	101.6 ± 5	334	HAS	[1981CHI/GAR]
C ₁₀ H ₁₅ NO	[80124-30-7]	(<i>d</i>)-carvoxime				
	$\Delta_{\text{fus}} H$		22.72	346.5	DTA	[1981CHI/GAR]
	$\Delta_{\text{sub}} H$	(324–343)	90.8 ± 4.5	334	HAS	[1981CHI/GAR]
C ₁₀ H ₁₅ NO	[90-82-4]	(+) <i>threo</i> -2-(methylamino)-1-phenyl-1-propanol (pseudoephedrine)				
	$\Delta_{\text{fus}} H$		31.95	392.4	DSC	[1999LI/ZEL]
C ₁₀ H ₁₅ NO	[4125-58-0]	(+) <i>threo</i> -2-(methylamino)-1-phenyl-1-propanol (pseudoephedrine)				
	$\Delta_{\text{fus}} H$		34.1	391.1	DSC	[1999LI/ZEL]
C ₁₀ H ₁₅ NO	[299-42-3]	(–) 2-(methylamino)-1-phenyl-1-propanol (ephedrine)				
	$\Delta_{\text{fus}} H$		17.33	312.9	DSC	[1999LI/ZEL]
C ₁₀ H ₁₅ NO	[90-81-3]	(+) 2-(methylamino)-1-phenyl-1-propanol (ephedrine)				
	$\Delta_{\text{fus}} H$		29.09	350.7	DSC	[1999LI/ZEL]
C ₁₀ H ₁₅ NO ₂	[120-07-0]	N,N-bis(2-hydroxyethyl)aniline				
	$\Delta_v H$	(418–611)	77.6	433	A	[1987STE/MAL, 1947STU]
C ₁₀ H ₁₅ NO ₂	[7575-82-8]	1-nitroadamantane				
	$\Delta_{\text{fus}} H$		4.18	435.2		[1990FRI/DOG]
	$\Delta_{\text{sub}} H$	(321–357)	63.6 ± 1.0	339	T	[1990FRI/DOG]
		Note: Entropy seems low, compound may have lower temperature phase transitions.				
C ₁₀ H ₁₅ NO ₂	[54654-31-7]	2-nitroadamantane				
	$\Delta_{\text{fus}} H$		4.23	452.2		[1990FRI/DOG]
	$\Delta_{\text{sub}} H$	(331–368)	58.0 ± 2.3	350	T	[1990FRI/DOG]
		Note: Entropy seems low, compound may have lower temperature phase transitions.				
C ₁₀ H ₁₅ N ₅	[153495-35-3]	6,9-dimethyl-8-propyladenine				
	$\Delta_{\text{fus}} H$		30.2	411.9		[1994ZIE/ZIE]
		(345–349)	129.0 ± 0.1	347	ME	[1994ZIE/ZIE]
C ₁₀ H ₁₅ N ₅	[117954-98-0]	8-butyl-9-methyladenine				

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{sub}}H$	(363–368)	135.1 ± 1.2	366	ME	[1987KAM/ZIE7]
C ₁₀ H ₁₅ N ₅ O ₃	[39809-25-1] $\Delta_{\text{fus}}H$	9-(4-hydroxy-3-hydroxymethylbut-1-yl)guanine	35.74	551.2		[2004AHM/BAR]
C ₁₀ H ₁₅ O ₃ PS ₂	[55-38-9] $\Delta_{\text{v}}H$	O,O-dimethyl-O-[3-methyl-4-(methylthio)phenyl]thiophosphate	75.6	308	A	[1987STE/MAL, 1999DYK/SVO]
C ₁₀ H ₁₆	[53130-19-1] $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$	tricyclo[4.3.1.0 ^{3,8}]decane	64.9 ± 1.8 65.6	323 298	TSGC H	[1975CLA/KNO] [1975CLA/KNO, 1993CHI/HOS]
C ₁₀ H ₁₆	[6004-38-2] $\Delta_{\text{trs}}H$ $\Delta_{\text{fus}}H$ $\Delta_{\text{fus}}H$	tricyclo[5.2.1.0 ^{2,6}]decane	2.57 3.07 2.95	204.3 345.3 352	AC	[2003KON/TAN] [1996DOM/HEA]
	$\Delta_{\text{sub}}H$	(359–443)	52.9 ± 1.3	298	BG	[1971BOY/SAN, 1977PED/RYL]
	$\Delta_{\text{v}}H$	(394–457)	46.0	409	EB	[2009XIN/FAN]
C ₁₀ H ₁₆	[17760-91-7] $\Delta_{\text{sub}}H$	tricyclo[5.2.1.0 _{4,10}]decane (hexahydrotriquinacene)	56.6 ± 1.3	307	TSGC	[1979CLA/KNO]
C ₁₀ H ₁₆	[283-50-1] $\Delta_{\text{sub}}H$	bicyclo[3.3.2]decane	58.2 ± 2	298		[1977PAR/STE]
C ₁₀ H ₁₆	[281-23-2] $\Delta_{\text{trs}}H$ $\Delta_{\text{fus}}H$ $\Delta_{\text{trs}}H$ $\Delta_{\text{fus}}H$ $\Delta_{\text{trs}}H$ $\Delta_{\text{fus}}H$	adamantane	3.21 10.9 3.38 10.9 3.38 13.8	208 541 208.6 541.2 208.6 543.2	AC DSC AC AC AC	[2006VAN/VAN] [1998CHI/HES] [1960CHA/WES] [1996DOM/HEA, 1966PIR, 1993HAK] [1960CHA/WES] [2000KAB/BLO]
	$\Delta_{\text{sub}}H$		59.1	298		[2000MOK/RUZ]
	$\Delta_{\text{sub}}H$		58.3	308		[2000KAB/BLO]
	$\Delta_{\text{sub}}H$		52.6	298	CGC-DSC	[1998CHI/HES]
	$\Delta_{\text{sub}}H$	(278–368)	59.7	293	A	[1987STE/MAL]
	$\Delta_{\text{sub}}H$	(328–373)	55.3	343	A	[1987STE/MAL]
	$\Delta_{\text{sub}}H$	(343–483)	54.3	358	A	[1987STE/MAL, 1968FLO]
	$\Delta_{\text{sub}}H$		58.45	298	C	[1982JOC/DEK]
	$\Delta_{\text{sub}}H$	(278–443)	59.5	300		[1975LEE/SLU]
	$\Delta_{\text{sub}}H$	(310–336)	59.7 ± 0.8	326	TSGC	[1975CLA/KNO]
	$\Delta_{\text{sub}}H$		58.6	298	H	[1975CLA/KNO, 1993CHI/HOS]
	$\Delta_{\text{sub}}H$	(310–336)	59.3 ± 0.2	326	BG	[1971BOY/SAN]
	$\Delta_{\text{sub}}H$		60.5 ± 1.3	298	H	[1971BOY/SAN, 1993CHI/HOS]
	$\Delta_{\text{sub}}H$	(312–366)	53.6	332	I	[1971WU/HSU]
	$\Delta_{\text{sub}}H$		54.8	298	H	[1971WU/HSU, 1993CHI/HOS]
	$\Delta_{\text{sub}}H$		59.3 ± 0.16	298	C	[1970MAN/RAP]
	$\Delta_{\text{sub}}H$		59.5	298		[1970VON/WIL, 1971BUT/CAR]
	$\Delta_{\text{sub}}H$	(313–353)	58.6 ± 0.6	333	DBM	[1967BRA/SZI]
	$\Delta_{\text{sub}}H$		59.6	298	H	[1967BRA/SZI, 1993CHI/HOS]
	$\Delta_{\text{sub}}H$		62.3	298		[1967BRA/SZI]
	$\Delta_{\text{v}}H$		48.2	298	GC	[2002VAN/PAR]
	$\Delta_{\text{v}}H$	(403–453)	51.7	298	CGC	[1995CHI/HOS]
C ₁₀ H ₁₆	[79-92-5] $\Delta_{\text{sub}}H$	(dl) 2,2-dimethyl-3-methylenebicyclo[2.2.1]heptane (camphene)	46.8		C	[1977KOZ/BYC]
	$\Delta_{\text{v}}H$	(320–434)	44.0	335	A	[1987STE/MAL, 1947STU]

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₀ H ₁₆	[4497-92-1] $\Delta_v H$	(<i>d</i>) 3,7,7-trimethylbicyclo[4.1.0]hept-2-ene (2-carene) (293–450)	45.5	308	A	[1987STE/MAL, 1954BUK/MAJ]
C ₁₀ H ₁₆	[498-15-7] $\Delta_v H$	(<i>d</i>) 3,7,7-trimethylbicyclo[4.1.0]hept-3-ene (3-carene) (359–443)	42.8	374	A	[1987STE/MAL]
C ₁₀ H ₁₆	[na] $\Delta_v H$	(+) limonene (373–423)	49.6	298	CGC	[1995CHI/HOS]
C ₁₀ H ₁₆	[na] $\Delta_v H$	(S)-(–) limonene (320–451)	47.4	335		[1996ROD/BER]
C ₁₀ H ₁₆	[5989-27-5] $\Delta_{\text{fus}} H$	(<i>d</i>) limonene	11.38	199.2		[1996GAL/VAN, 1996GAL/VAN]
	$\Delta_v H$	(326–445)	49.5	298		[2009CLA/GOM]
	$\Delta_v H$	(313–413)	49.6	298	GC	[2005HOS/GRY]
	$\Delta_v H$		49.9	298	GC	[2002VAN/PAR]
	$\Delta_v H$	(339–495)	46.1 ± 0.2	350	EB	[2002STE/CHI2]
	$\Delta_v H$	(339–495)	43.5 ± 0.2	390	EB	[2002STE/CHI2]
	$\Delta_v H$	(339–495)	40.9 ± 0.3	430	EB	[2002STE/CHI2]
	$\Delta_v H$	(339–495)	37.9 ± 0.6	470	EB	[2002STE/CHI2]
	$\Delta_v H$	(250–434)	49.2	300		[1999DIA/GUE]
	$\Delta_v H$		48.9 ± 0.1	298	C	[1987ATI/SAI]
	$\Delta_v H$	(287–448)	44.5	302	A	[1987STE/MAL]
	$\Delta_v H$	(288–323)	47.7	303	A	[1987STE/MAL]
C ₁₀ H ₁₆	[5989-54-8] $\Delta_v H$	(<i>l</i>) limonene (325–450)	47.0	340		[1993NAD/BER]
	$\Delta_v H$		49.0 ± 0.1	298	C	[1987ATI/SAI]
	$\Delta_v H$	(303–363)	45.5	318	A	[1987STE/MAL, 1954BUK/MAJ]
C ₁₀ H ₁₆	[138-86-3] $\Delta_v H$	(<i>dl</i>) limonene (287–448)	45.9	302	A	[1987STE/MAL]
C ₁₀ H ₁₆	[na] $\Delta_v H$	limonene (353–405)	39.4	379	TGA	[2002HAZ/DOL]
C ₁₀ H ₁₆	[na] $\Delta_v H$	β -myrcene (303–363)	47.0	318		[1954BUK/MAJ]
C ₁₀ H ₁₆	[123-35-3] $\Delta_v H$	7-methyl-3-methylene-1,6-octadiene (myrcene)	50.6	298	GC	[2002VAN/PAR]
	$\Delta_v H$	(287–445)	45.7	302	A	[1987STE/MAL, 1947STU]
C ₁₀ H ₁₆	[99-83-2] $\Delta_v H$	5-isopropyl-2-methyl-1,3-cyclohexadiene (293–448)	47.7	308	A	[1987STE/MAL]
C ₁₀ H ₁₆	[555-10-2] $\Delta_v H$	3-isopropyl-6-methylenecyclohexene (303–363)	47.7	318	A	[1987STE/MAL, 1954BUK/MAJ]
C ₁₀ H ₁₆	[na] $\Delta_v H$	α -pinene (320–429)	42.5	335		[1996ROD/BER]
	$\Delta_v H$	(365–430)	40.2	380		[1993REI/SAN]
	$\Delta_v H$		44.6 ± 0.1	298	C	[1987AN/HU]
C ₁₀ H ₁₆	[80-56-8] $\Delta_v H$	(<i>d</i>) α -pinene (308–427)	45.4	298		[2009CLA/GOM]
	$\Delta_v H$	(292–433)	45.0	307	A	[1987STE/MAL]
	$\Delta_v H$	(293–363)	43.4	308		[1954BUK/MAJ]
C ₁₀ H ₁₆	[127-91-3] $\Delta_v H$	β -pinene (290–439)	46.0	305		[1996ROD/BER]
	$\Delta_v H$	(364–439)	41.6	379		[1993REI/SAN]

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$		45.8 ± 0.1	298	C	[1987AN/HU]
C ₁₀ H ₁₆	[na]	(l) β -pinene				
	$\Delta_v H$	(291–441)	46.1	306	A	[1987STE/MAL]
	$\Delta_v H$	(293–363)	44.9	308		[1954BUK/MAJ]
C ₁₀ H ₁₆	[586-62-9]	terpinolene				
	$\Delta_v H$	(313–363)	50.8	328		[1954BUK/MAJ]
	$\Delta_v H$	(305–458)	50.5	320	A	[1987STE/MAL, 1947STU]
C ₁₀ H ₁₆	[79-92-5]	camphene				
	$\Delta_v H$	(313–413)	44.7	298	GC	[2005HOS/GRY]
C ₁₀ H ₁₆	[99-85-4]	γ -terpinene				
	$\Delta_v H$	(313–413)	51.4	298	GC	[2005HOS/GRY]
C ₁₀ H ₁₆	[4221-98-1]	sabinene				
	$\Delta_v H$	(313–413)	46.9	298	GC	[2005HOS/GRY]
C ₁₀ H ₁₆	[4221-98-1]	(-)- α -phellandrene				
	$\Delta_v H$	(313–423)	48.3	298	GC	[2005HOS/GRY]
C ₁₀ H ₁₆	[4497-92-1]	(+) 2-carene				
	$\Delta_v H$	(313–413)	48.5	298	GC	[2005HOS/GRY]
C ₁₀ H ₁₆	[498-15-7]	(+) 3-carene				
	$\Delta_v H$	(313–413)	48.5	298	GC	[2005HOS/GRY]
C ₁₀ H ₁₆	[2867-05-2]	α -thujene				
	$\Delta_v H$	(313–413)	44.8	298	GC	[2005HOS/GRY]
C ₁₀ H ₁₆	[6004-38-2]	tetrahydrodicyclopentadiene				
	$\Delta_v H$	(358–465)	43.5	373	A	[1987STE/MAL]
C ₁₀ H ₁₆	[2825-82-3]	<i>exo</i> tetrahydrodicyclopentadiene				
	$\Delta_{\text{trs}}H$		3.18	162.1		
	$\Delta_{\text{fus}}H$		1.2	183.2	DSC	[2002CHI/HIL]
C ₁₀ H ₁₆	[2825-83-4]	<i>endo</i> tetrahydrodicyclopentadiene				
	$\Delta_{\text{trs}}H$		10.7	213.8		
	$\Delta_{\text{fus}}H$		3.48	356.8	DSC	[2002CHI/HIL]
C ₁₀ H ₁₆ ClO ₆	[na]	lactic acid, O-ethoxycarbonyl, 2-(2-chloroethoxy)ethyl ester				
	$\Delta_v H$	(406–523)	83.8 (sub)	421	A	[1987STE/MAL]
C ₁₀ H ₁₆ Cl ₃ NOS	[2303-17-5]	carbamothioic acid, <i>bis</i> (isopropyl), S-(2,3,3-trichloroallyl) ester				
	$\Delta_{\text{fus}}H$		27.11	306.4	DSC	[1991ACR, 1990DON/DRE]
	$\Delta_v H$	(293–318)	84.3	305	A	[1987STE/MAL]
C ₁₀ H ₁₆ NOS	[2303-17-5]	S-2,3,3-trichloroallyl N,N-diisopropylthiocarbonate (triallate)				
	$\Delta_{\text{sub}}H$	(293–318)	84.0			[1983SPE/CLI, 1978GRO/SPE]
C ₁₀ H ₁₆ NO ₄ PS	[52-85-7]	O-[4-(dimethylamino)sulfonyl]phenyl] O,O-dimethylphosphorothionate				
	$\Delta_{\text{fus}}H$		26.5	326.8	DSC	[1990DON/DRE]
C ₁₀ H ₁₆ N ₂	[1871-96-1]	sebaconitrile				
	$\Delta_{\text{fus}}H$		28.2	281.2	DSC	[2007BAD/BLA]
	$\Delta_v H$	(303–343)	83.7	318	A	[1987STE/MAL]
C ₁₀ H ₁₆ N ₂	[33089-74-6]	N'-(2,4-dimethylphenyl)-N-methylformamidine				
	$\Delta_v H$		89.2	303		[1998ZHA/MO]
C ₁₀ H ₁₆ N ₂	[85688-96-6]	methyl(1,1,1-trimethylpropyl)propanedinitrile				

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound					
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference	
		$\Delta_{\text{sub}}H$	62.0 ± 0.7	298		[1990BEC/DOG]	
C ₁₀ H ₁₆ N ₂	[85688-95-5]	(1,1-dimethylpropyl)ethylpropanedinitrile					
	$\Delta_{\text{fus}}H$		19.25	307.5		[1990BEC/DOG]	
	$\Delta_{\text{sub}}H$		76.2 ± 0.8	298		[1990BEC/DOG]	
C ₁₀ H ₁₆ N ₂	[na]	<i>meso</i> 2,3-diethyl-2,3-dimethylsuccinonitrile					
	$\Delta_{\text{fus}}H$		26.78	370.2		[1983BAR/BEC]	
C ₁₀ H ₁₆ N ₂ O ₂	[82413-40-9]	1,3-dimethyl-5-butyluracil					
	$\Delta_{\text{fus}}H$		22	312.1		[1996KAM/ZIE]	
		$\Delta_{\text{sub}}H$	(306–311)	106.3 ± 1.3	309	ME	[1996KAM/ZIE]
C ₁₀ H ₁₆ N ₂ O ₂	[77-28-1]	5-butyl-5-ethylbarbituric acid (butobarbital)					
	$\Delta_{\text{fus}}H$ (I)		13.6	394.2			
	$\Delta_{\text{fus}}H$ (II)		14.7	392.7			
	$\Delta_{\text{fus}}H$ (III)		17.3	396.2	DSC	[1989CHA/DEM]	
C ₁₀ H ₁₆ N ₄ O ₂ S	[55511-98-3]	3-(5-(1,1-dimethylethyl)-1,3,4-thiadiazol-2-yl)-4-hydroxy-1-methyl-2-imidazolidinone					
	$\Delta_{\text{fus}}H$		25.46	408.9	DSC	[1990DON/DRE]	
C ₁₀ H ₁₆ N ₆ S	[51481-61-9]	N-cyano-N'-methyl-N''-[2-[(5-methyl-1 <i>H</i> -imidazol-4-yl)-thio]ethyl]guanidine (cimetidine)					
	$\Delta_{\text{fus}}H$ (I)		41.0	413.8			
	$\Delta_{\text{fus}}H$ (II)		39.7	413.5	DSC	[1999BAU/MAR]	
	$\Delta_{\text{fus}}H$ (A)		38.2	412.2	DSC		
	$\Delta_{\text{fus}}H$ (B)		34.8	413.7	DSC	[1996BAU]	
		Note: Cimetidine is reported to have four different crystalline forms					
C ₁₀ H ₁₆ O	[29171-20-8]	3,7-dimethyl-6-octen-1-yn-3-ol (dehydrolinalool)					
	Δ_vH	(406–471)	52.1	421	EB	[2001ZHU/LI]	
	Δ_vH	(359–381)	U			[1999ZAR/CHA]	
	Δ_vH	(369–445)	50.4 ± 0.1	407		[1988BAG/GUR]	
C ₁₀ H ₁₆ O	[na]	camphor					
	Δ_vH	(343–383)	54.4	298	CGC	[1995CHI/HOS]	
	Δ_vH	(343–383)	54.5	298	CGC	[1995CHI/HOS]	
	Δ_vH	(343–383)	55.2	298	CGC	[1995CHI/HOS]	
C ₁₀ H ₁₆ O	[464-49-3]	(<i>d</i>)-camphor					
	$\Delta_{\text{fus}}H$		16.0	242			
	$\Delta_{\text{us}}H$		0.23	374			
	$\Delta_{\text{fus}}H$		5.3	452	DSC	[1979MJO]	
C ₁₀ H ₁₆ O	[76-22-2]	(<i>dl</i>)-camphor					
	$\Delta_{\text{sub}}H$		51.8 ± 0.8			[1977STE]	
	$\Delta_{\text{sub}}H$	(273–293)	51.5 ± 2.6	283	HSA	[1975CHI]	
	$\Delta_{\text{sub}}H$	(273–298)	U 65.8			[1960JON, 1940ZIB]	
	$\Delta_{\text{sub}}H$		50.7			[1960JON, 1937DEW]	
	$\Delta_{\text{sub}}H$	(273–453)	53.6	363		[1960JON]	
	$\Delta_{\text{sub}}H$	(285–318)	54.7	301		[1957SHE/BRY]	
C ₁₀ H ₁₆ O	[464-49-3]	(+) camphor					
	Δ_vH		55.3	298	GC	[2002VAN/PAR]	
C ₁₀ H ₁₆ O	[1686-14-2]	α -pinene oxide					
	Δ_vH		53.6	298	GC	[2002VAN/PAR]	
C ₁₀ H ₁₆ O	[13854-85-8]	(<i>d</i>) 3-bornanone					
	$\Delta_{\text{sub}}H$	(273–408)	54.2	288	A	[1987STE/MAL]	
	$\Delta_{\text{sub}}H$	(323–339)	55.0	331	A	[1987STE/MAL]	
		$\Delta_{\text{sub}}H$	(408–451)	49.8	423	A	[1987STE/MAL]

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(452–488)	44.6	467	A	[1987STE/MAL]
C ₁₀ H ₁₆ O	[4695-62-9]	(d) 1,3,3-trimethylbicyclo[2.2.1]heptan-2-one (fenchone)				
	$\Delta_v H$	(365–384)	47.0	374		[2002BAT]
	$\Delta_v H$		51.7 ± 0.1	298	C	[1987ATI/SAI]
	$\Delta_v H$		51.4 ± 0.1	298	C	[1987ATI/SAI]
	$\Delta_v H$		51.1 ± 0.1	298	C	[1985KUS]
	$\Delta_v H$	(301–464)	48.9	316	A	[1987STE/MAL, 1947STU]
C ₁₀ H ₁₆ O	[na]	(l) 1,3,3-trimethylbicyclo[2.2.1]heptan-2-one (fenchone)				
	$\Delta_v H$		51.1 ± 0.1	298	C	[1987ATI/SAI]
	$\Delta_v H$		51.3 ± 0.1	298	C	[1987ATI/SAI]
	$\Delta_v H$		51.4 ± 0.1	298	C	[1985KUS]
C ₁₀ H ₁₆ O	[89-82-7]	pulegone				
	$\Delta_v H$	(353–453)	58.0	298	GC	[2005HOS/GRY]
	$\Delta_v H$		62.0	298	GC	[2002VAN/PAR]
	$\Delta_v H$	(331–494)	U99.8	346	A	[1987STE/MAL, 1947STU]
C ₁₀ H ₁₆ O	[471-15-8]	(d) 1-isopropyl-4-methylbicyclo[3.1.0]hexan-3-one (thujone)				
	$\Delta_v H$	(311–474)	51.8	326	A	[1987STE/MAL, 1947STU]
C ₁₀ H ₁₆ O	[499-74-1]	(dl) 3-isopropyl-6-methyl-2-cyclohexene-1-one				
	$\Delta_v H$	(364–507)	56.9	379	A	[1987STE/MAL]
C ₁₀ H ₁₆ O	[43205-82-9]	5-isopropyl-2-methyl-2-cyclohexen-1-one				
	$\Delta_v H$	(361–503)	56.8	376	A	[1987STE/MAL]
C ₁₀ H ₁₆ O	[na]	3,4-epoxycarane (α -isomer)				
	$\Delta_v H$		49.4			[1977ALE/KOZ]
C ₁₀ H ₁₆ O	[na]	3,4-epoxycarane (β -isomer)				
	$\Delta_v H$		50.2			[1977ALE/KOZ]
C ₁₀ H ₁₆ O	[4584-09-2]	(dl) dihydrocarvone				
	$\Delta_v H$	(319–496)	51.2	334	A	[1987STE/MAL, 1947STU]
C ₁₀ H ₁₆ O	[5392-40-5]	citral				
	$\Delta_v H$	(372–473)	55.9	298		[2009CLA/GOM]
C ₁₀ H ₁₆ O	[141-27-5]	geranial (<i>trans</i> -citral)				
	$\Delta_v H$	(343–453)	62.5	298	GC	[2005HOS/GRY]
	$\Delta_v H$	(283–333)	61	298	A	[1987STE/MAL]
	$\Delta_v H$	(373–501)	54.9	388	A	[1987STE/MAL]
C ₁₀ H ₁₆ O	[13040-03-4]	(+)– <i>cis</i> -verbenol				
	$\Delta_v H$	(323–433)	54.9	298	GC	[2005HOS/GRY]
C ₁₀ H ₁₆ O	[22339-08-8]	(+)– <i>trans</i> -verbenol				
	$\Delta_v H$	(323–433)	55.0	298	GC	[2005HOS/GRY]
C ₁₀ H ₁₆ O	[106-26-3]	neral				
	$\Delta_v H$	(343–453)	60.2	298	GC	[2005HOS/GRY]
C ₁₀ H ₁₆ O	[768-95-6]	1-adamantanol				
	$\Delta_{\text{us}}H$		11.29	357.1		
	$\Delta_{\text{fus}}H$		12.36	552.9		[2003CHA/BLO]
	$\Delta_{\text{us}}H$		2.5	369.2		[1989SAL/ABA2]
	$\Delta_{\text{sub}}H$		86.8 ± 0.2	298	C	[2003CHA/BLO2]
	$\Delta_{\text{sub}}H$	(288–323)	86.7 ± 0.2	298	ME	[2003CHA/BLO2]
	$\Delta_{\text{sub}}H$		86.6 ± 2.5	298	BG	[1978ARO/STE]
	$\Delta_v H$		60.8	298	GC	[2002VAN/PAR]

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₀ H ₁₆ O	[700-57-2]	adamantan-2-ol				
	$\Delta_{\text{us}}H$		0.16	238.4		
	$\Delta_{\text{us}}H$		2.3	322.3		
	$\Delta_{\text{us}}H$		7.98	389.3		
	$\Delta_{\text{fus}}H$		11.94	567.3		[2003CHA/BLO]
	$\Delta_{\text{us}}H$		0.3	325.2		
	$\Delta_{\text{us}}H$		3.74	391.2		[1989SAL/ABA2]
	$\Delta_{\text{sub}}H$	(303–318)	88.1 ± 1.6	298	ME	[2003CHA/BLO2]
$\Delta_{\text{sub}}H$		88.7 ± 2.5	298	BG	[1978ARO/STE]	
C ₁₀ H ₁₆ O	[20379-99-1]	<i>trans</i> octahydro-3a-methyl-2 <i>H</i> -inden-2-one				
	Δ_vH		58.3 ± 0.2	298	C	[1970SEL]
C ₁₀ H ₁₆ O	[13351-29-6]	<i>cis</i> -8-methyl-2-hydrindanone				
	$\Delta_{\text{sub}}H$		60.9 ± 0.2	298	C	[1970SEL, 1977PED/RYL]
C ₁₀ H ₁₆ O ₂	[2704-78-1]	3-acetyl-2,2-dimethylcyclobutaneacetaldehyde (pinonaldehyde)				
	Δ_vH	(283–308)	75.5 ± 5.6		ME	[1997HAL/WAN]
C ₁₀ H ₁₆ O ₂	[26946-56-5]	2,2-dimethyl-3-(2-oxopropyl)cyclopropaneacetaldehyde (caronaldehyde)				
	Δ_vH	(283–308)	77.4 ± 6.9		ME	[1997HAL/WAN]
C ₁₀ H ₁₆ O ₂	[490-03-9]	diosphenol				
	Δ_vH	(339–505)	56.2	354	A	[1987STE/MAL, 1947STU]
C ₁₀ H ₁₆ O ₂	[512-77-6]	1-methyl-3-(1-methylethyl)-cyclopentanecarboxylic acid (fencholic acid)				
	Δ_vH	(374–537)	77.5	389		[1947STU]
C ₁₀ H ₁₆ O ₂	[na]	(2,3,3-trimethyl-3-cyclopentadienyl)acetic acid				
	Δ_vH	(370–529)	71.3	385	A	[1987STE/MAL, 1947STU]
C ₁₀ H ₁₆ O ₂	[38734-05-3]	1,6-cyclodecanedione				
	$\Delta_{\text{fus}}H$		29.58	372.2		[1972ALV/BOR]
C ₁₀ H ₁₆ O ₂	[10453-89-1]	2,2-dimethyl-3-(2-methyl-1-propen-1-yl)cyclopropanecarboxylic acid (chrysanthemic acid)				
	$\Delta_{\text{fus}}H$		14.51	390.7		[2004XUE/WAN]
C ₁₀ H ₁₆ O ₄	[na]	1,4-cyclohexanedione <i>bis</i> ethylene ketal				
	$\Delta_{\text{fus}}H$		25.77	353.2		[1972ALV/BOR]
C ₁₀ H ₁₆ O ₄	[na]	1,1-cyclobutanedicarboxylic acid diethyl ester				
	Δ_vH	(288–318)	65.8 ± 0.4	GS		[1998VER/KUM]
C ₁₀ H ₁₆ O ₆	[na]	lactic acid, O-ethoxycarbonyl, tetrafurfuryl ester				
	Δ_vH	(390–523)	71.2	405	A	[1987STE/MAL]
C ₁₀ H ₁₆ O ₆	[6279-86-3]	<i>tris</i> -(carboethoxy)methane				
	Δ_vH	(297–338)	79.1 ± 0.7	298	GS	[1992VER/BEC]
C ₁₀ H ₁₆ S	[53402-10-1]	(1 <i>R</i>) (-)-thiocamphor				
	$\Delta_{\text{sub}}H$	(262–282)	62.2 ± 0.9	272	ME	[1999ROU/JIM]
	$\Delta_{\text{sub}}H$		61.7 ± 0.9	298		[1999ROU/JIM]
	Δ_vH		55.5	298	GC	[2002VAN/PAR]
C ₁₀ H ₁₆ S	[18794-77-9]	2-hexylthiophene				
	Δ_vH		56.4 ± 1.3	298	C	[2007RIB/SAN]
C ₁₀ H ₁₆ S	[1693-86-3]	3-hexylthiophene				
	Δ_vH		58.5 ± 1.3	298	C	[2007RIB/SAN]
C ₁₀ H ₁₆ S ₄	[7000-79-5]	1,3,5,7-tetramethyl-2,4,6,8-tetrathiaadamantane				
	$\Delta_{\text{sub}}H$		117.1 ± 4.1	298	TE	[78HEA/HEF]

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₀ H ₁₇ N	[na]	1-cyclohexylimino-2-butene	58.3			[1993OVC/SOB]
C ₁₀ H ₁₇ N	[768-94-5]	1-aminoadamantane				
	$\Delta_{\text{us}}H$	(5–370)	1.72	241.4	AC	
	$\Delta_{\text{us}}H$	(5–370)	5.31	284.6	AC	[2008BAZ/BLO]
	$\Delta_{\text{fus}}H$		NA		AC	
C ₁₀ H ₁₇ NO	[2792-42-9]	(<i>d</i>) camphor oxime				
	$\Delta_{\text{us}}H$		14.48	384.5		
	$\Delta_{\text{fus}}H$		2.1	393.3	DSC	[1984WEI/LEF]
	$\Delta_{\text{us}}H$		13.3	383		
C ₁₀ H ₁₇ NO	[na]	(<i>dl</i>) camphor oxime				
	$\Delta_{\text{us}}H$		3	375		
	$\Delta_{\text{us}}H$		11.2	380		
	$\Delta_{\text{fus}}H$		1.2	388	DSC	[1979MJO]
C ₁₀ H ₁₇ NOS	[59300-35-5]	carbamothioic acid, N-butyl-N-(2-propynyl), S-ethyl ester				
$\Delta_{\text{sub}}H$	(298–313)	82.1	305.5	ME	[1987STE/MAL, 1976DEP]	
C ₁₀ H ₁₇ NOS	[59300-36-6]	carbamothioic acid, N,N-dipropyl S-(2-propynyl) ester				
$\Delta_{\text{sub}}H$	(298–313)	92.4	305.5	ME	[1987STE/MAL, 1976DEP]	
C ₁₀ H ₁₇ NOS	[59300-34-4]	carbamothioic acid, N-2-methylpropyl-N-(2-propynyl), S-ethyl ester				
$\Delta_{\text{sub}}H$	(298–313)	74.0	305.5	ME	[1987STE/MAL, 1976DEP]	
C ₁₀ H ₁₇ NO ₃	[na]	2-(2-cyanoethoxy)propionic acid, butyl ester				
Δ_vH	(328–382)	61.7	343	A	[1987STE/MAL]	
C ₁₀ H ₁₇ NO ₅	[1069-39-2]	(<i>l</i>) N-acetylaspartic acid, diethyl ester				
Δ_vH	(418–508)	76.0	433	A	[1987STE/MAL]	
C ₁₀ H ₁₇ N ₅ O	[1610-18-0]	(<i>l</i>) N-acetylaspartic acid, diethyl ester				
$\Delta_{\text{fus}}H$		21.18	363.5	DSC	[1991ACR, 1990DON/DRE]	
C ₁₀ H ₁₈	[176-63-6]	spiro[4.5]decane				
	Δ_vH		54.8	298	C	[1975SUB/ZWO]
	Δ_vH	(348–389)	44.0	363		[1965NAR]
C ₁₀ H ₁₈	[16189-46-1]	<i>cis</i> bicyclo[5.3.0]decane				
	Δ_vH	(298–377)	49.8	313	A	[1987STE/MAL]
	Δ_vH		46.9 ± 0.8	377		[1970CHA/MCN]
	Δ_vH		53.6 ± 1.2	298		[1970CHA/MCN]
C ₁₀ H ₁₈	[1636-39-1]	bicyclopentyl				
	$\Delta_{\text{us}}H$		0.26	171.5		
	$\Delta_{\text{fus}}H$		13.4	237.8		[2004CHI/STE]
	Δ_vH	(383–509)	50.4 ± 0.1	298	EB	[2005CHI/STE]
	Δ_vH	(384–509)	44.1 ± 0.1	400	EB	[2004CHI/STE]
	Δ_vH	(384–509)	41.6 ± 0.1	440	EB	[2004CHI/STE]
	Δ_vH	(384–509)	38.9 ± 0.2	480	EB	[2004CHI/STE]
Δ_vH	(350–393)	43.2	365	A	[1987STE/MAL]	
C ₁₀ H ₁₈	[18968-24-6]	<i>cis</i> carane				
Δ_vH	(362–445)	42.8	377	A	[1987STE/MAL]	
C ₁₀ H ₁₈	[493-01-6]	<i>cis</i> decahydronaphthalene				
$\Delta_{\text{fus}}H$		14.43	242.8		[1996DOM/HEA]	

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound					
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference	
		$\Delta_{\text{sub}}H$	64.8	230	B	[1963BON]	
		$\Delta_{\text{sub}}H$	62.5	298	H	[1963BON, 1993CHI/HOS]	
		Δ_vH	(371–473)	45.5	386	A,GS	[1987STE/MAL, 1955CAM/ROS]
C₁₀H₁₈	[493-02-7]	<i>trans</i> decahydronaphthalene					
		$\Delta_{\text{us}}H$	2.13	216.1			
		$\Delta_{\text{fus}}H$	9.49	230.2		[1996DOM/HEA]	
		$\Delta_{\text{sub}}H$	66.2	241	B	[1963BON]	
		$\Delta_{\text{sub}}H$	64.3	298	H	[1963BON, 1993CHI/HOS]	
		Δ_vH	(363–461)	44.2	378	A,GS	[1987STE/MAL, 1955CAM/ROS]
C₁₀H₁₈	[1942-46-7]	5-decyne					
		Δ_vH	(477–487)	45.5	366	A	[1987STE/MAL]
C₁₀H₁₈	[4755-33-3]	<i>cis</i> 2,6,6-trimethylbicyclo[3.1.1]heptane (<i>cis</i> -pinane)					
		Δ_vH	(378–441)	41.8	393		[2003WAN/LI]
C₁₀H₁₈Cl₄	[205646-11-3]	1,2,9,10-tetrachlorodecane					
		Δ_vH	75.4			[1998DRO/TOM]	
C₁₀H₁₈N₆O₂	[na]	1-(sarcosino)-3,5-bis(dimethylamino)-s-triazine					
		$\Delta_{\text{fus}}H$	29.83	431		[1989BRA/RYT]	
C₁₀H₁₈O	[470-82-6]	1,8-epoxy- <i>p</i> -menthane					
		Δ_vH	(264–303)	35.6	279	A	[1987STE/MAL]
C₁₀H₁₈O	[6627-72-1]	<i>(dl)</i> borneol					
		$\Delta_{\text{sub}}H$	(350–475)	69.3	365	A	[1987STE/MAL]
		Δ_vH	(477–487)	50.9	482	A	[1987STE/MAL]
C₁₀H₁₈O	[470-67-7]	1,4-cineole					
		Δ_vH	(288–449)	46.1	303	A	[1987STE/MAL]
C₁₀H₁₈O	[470-82-6]	1,8-cineole					
		Δ_vH	(313–413)	49.0	298	GC	[2005HOS/GRY]
		Δ_vH		53.2	298	GC	[2002VAN/PAR]
		Δ_vH	(353–403)	41.1	378	TGA	[2002HAZ/DOL]
C₁₀H₁₈O	[619-01-2]	<i>(d)</i> dihydrocarveol					
		Δ_vH	(336–498)	58.2	351	A	[1987STE/MAL]
C₁₀H₁₈O	[2217-01-8]	<i>(dl)</i> fenchyl alcohol					
		Δ_vH	(318–474)	89.1	333	A	[1987STE/MAL]
C₁₀H₁₈O	[106-24-1]	geraniol					
		Δ_vH	(288–333)	62.9	303	A	[1987STE/MAL]
		Δ_vH	(342–503)	59.1	357	A	[1987STE/MAL]
C₁₀H₁₈O	[7786-67-6]	<i>(d)</i> isopulegol					
		Δ_vH	(335–485)	49.8	350	A	[1987STE/MAL]
C₁₀H₁₈O	[126-90-9]	<i>(d)</i> linalool					
		Δ_vH	(273–321)	65.4	297		[1999DIA/GUE]
		Δ_vH	(313–471)	52.4	328	A	[1987STE/MAL]
C₁₀H₁₈O	[78-70-6]	linalool					
		Δ_vH	(352–468)	65.0	298		[2009CLA/GOM]
		Δ_vH	(333–433)	55.3	298	GC	[2005HOS/GRY]
		Δ_vH	(368–428)	51.4	399	TGA	[2002HAZ/DOL]
		Δ_vH	(409–465)	50.3	424	EB	[2002DEN/LI]

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₀ H ₁₈ O	[106-25-2] $\Delta_v H$	<i>cis</i> 3,7-dimethyl-2,6-octadien-1-ol (nerol) (334–499)	55.4	349	A	[1987STE/MAL]
C ₁₀ H ₁₈ O	[98-55-5] $\Delta_{\text{sub}} H$	(<i>dl</i>) α -terpineol (287–308)	80.1	297.5	A	[1987STE/MAL]
	$\Delta_v H$	(325–491)	54.0	340	A	[1987STE/MAL]
C ₁₀ H ₁₈ O	[7785-53-7] $\Delta_v H$	(+) α -terpineol	60.7	298	GC	[2002VAN/PAR]
C ₁₀ H ₁₈ O	[10482-56-1] $\Delta_{\text{sub}} H$	α -terpineol (283–328)	80.3	305	ME	[1954SER/VOI, 1960JON]
C ₁₀ H ₁₈ O	[562-74-3] $\Delta_v H$	terpinen-4-ol (323–433)	55.5	298	GC	[2005HOS/GRY]
C ₁₀ H ₁₈ O	[124-76-5] $\Delta_{\text{sub}} H$	(<i>dl</i>)-isoborneol (373–457)	41.1	388	A	[1987STE/MAL, 1936GRE]
C ₁₀ H ₁₈ O	[na] $\Delta_v H$	1-(1-methylcyclohex-3-enyl)-1-propanol (397–422)	53.6	409	A	[1987STE/MAL]
C ₁₀ H ₁₈ O	[1502-06-3] $\Delta_{\text{fus}} H$	cyclodecanone	24.3	294.9		[1998GON/SZW]
	$\Delta_v H$	(353–423)	55.2	368	A	[1987STE/MAL]
	$\Delta_v H$		58.4 ± 0.6	298		[1972WOL]
C ₁₀ H ₁₈ O	[na] $\Delta_v H$	ethyl (1-methylcyclohexyl) ketone (388–431)	45.2	403	A	[1987STE/MAL]
C ₁₀ H ₁₈ O	[10458-14-7] $\Delta_v H$	2-isopropyl-5-methylcyclohexanone (menthone) (372–397)	50.1	385		[2002BAT]
	$\Delta_v H$	(350–483)	51.2	365	A	[1987STE/MAL]
C ₁₀ H ₁₈ O	[2385-77-5] $\Delta_v H$	(<i>d</i>) citronellal (288–333)	54.9	303	A	[1987STE/MAL]
	$\Delta_v H$	(317–480)	53.2	332	A	[1987STE/MAL, 1947STU]
C ₁₀ H ₁₈ O	[69891-94-7] $\Delta_v H$	(<i>Z</i>) 3-decenal (323–343)	59.2	298	CGC	[1996KOU/HOS, 2000OVA/KOU]
C ₁₀ H ₁₈ O	[68676-85-7] $\Delta_v H$	(<i>E</i>) 3-decenal (323–343)	59.8	298	CGC	[1996KOU/HOS, 2000OVA/KOU]
C ₁₀ H ₁₈ O	[21662-09-9] $\Delta_v H$	(<i>Z</i>) 4-decenal (323–343)	59.3	298	CGC	[1996KOU/HOS, 2000OVA/KOU]
C ₁₀ H ₁₈ O	[65405-70-1] $\Delta_v H$	(<i>E</i>) 4-decenal (323–343)	60	298	CGC	[1996KOU/HOS, 2000OVA/KOU]
C ₁₀ H ₁₈ O	[21662-08-8] $\Delta_v H$	(<i>Z</i>) 5-decenal (323–343)	58.5	298	CGC	[1996KOU/HOS, 2000OVA/KOU]
C ₁₀ H ₁₈ O	[21662-11-3] $\Delta_v H$	(<i>E</i>) 5-decenal (323–343)	59.2	298	CGC	[1996KOU/HOS, 2000OVA/KOU]
C ₁₀ H ₁₈ O	[147159-48-6] $\Delta_v H$	(<i>Z</i>) 6-decenal (323–343)	59.3	298	CGC	[1996KOU/HOS, 2000OVA/KOU]
C ₁₀ H ₁₈ O	[147159-48-6] $\Delta_v H$	(<i>E</i>) 6-decenal (323–343)	59.5	298	CGC	[1996KOU/HOS, 2000OVA/KOU]
C ₁₀ H ₁₈ O	[21661-97-2]	(<i>Z</i>) 7-decenal				

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(323–343)	59.9	298	CGC	[1996KOU/HOS, 2000OVA/KOU]
C ₁₀ H ₁₈ O	[21662-10-2]	(E) 7-decenal				
	$\Delta_v H$	(323–343)	59.8	298	CGC	[1996KOU/HOS, 2000OVA/KOU]
C ₁₀ H ₁₈ O	[174155-46-5]	(Z) 8-decenal				
	$\Delta_v H$	(323–343)	60.5	298	CGC	[1996KOU/HOS, 2000OVA/KOU]
C ₁₀ H ₁₈ O	[174155-47-6]	(E) 8-decenal				
	$\Delta_v H$	(323–343)	60.2	298	CGC	[1996KOU/HOS, 2000OVA/KOU]
C ₁₀ H ₁₈ O ₂	[502-47-6]	3,7-dimethyl-6-octenoic acid				
	$\Delta_v H$	(372–530)	68.7	387	A	[1987STE/MAL, 1947STU]
C ₁₀ H ₁₈ O ₂	[na]	8,8-dimethyl-6,10-dioxaspiro[4.5]decane				
	$\Delta_v H$	(283–313)	53.7 ± 0.5	298	GS	[1998VER/PEN, 2002VER]
C ₁₀ H ₁₈ O ₂	[7333-23-5]	2,2,6-dimethyl-3,5-heptanedione				
	$\Delta_v H$		57.7	298		[1976BUR/SHR]
C ₁₀ H ₁₈ O ₂	[na]	δ -decanolactone				
	$\Delta_v H$	(365–387)	57.7 ± 0.8	376	MM	[1991WIB/WAL]
	$\Delta_v H$	(365–387)	63.0 ± 1.5	298	MM	[1991WIB/WAL]
C ₁₀ H ₁₈ O ₂	[706-14-9]	γ -decanolactone				
	$\Delta_v H$	(298–365)	75.6 ± 0.3	298	GS	[2008EME/KOZ]
C ₁₀ H ₁₈ O ₂	[705-86-2]	δ -decanolactone				
	$\Delta_v H$	(309–358)	74.2 ± 0.3	298	GS	[2007EME/KOZ]
C ₁₀ H ₁₈ O ₂	[2499-58-3]	heptyl acrylate				
	$\Delta_v H$	(359–481)	51.1	374	A	[1987STE/MAL]
C ₁₀ H ₁₈ O ₂	[142-09-6]	hexyl methacrylate				
	$\Delta_v H$	(354–475)	50.5	369	A	[1987STE/MAL]
C ₁₀ H ₁₈ O ₂	[512-77-6]	1-methyl-3-isopropylcyclopentane carboxylic acid				
	$\Delta_v H$	(374–538)	91.6	389	A	[1987STE/MAL]
C ₁₀ H ₁₈ O ₂	[1551-44-6]	cyclohexyl butyrate				
	$\Delta_v H$		60.0 ± 0.2	298	C	[2004PAU/ZAI]
	$\Delta_v H$		60.1 ± 0.2	298	C	[2003ZAI/VER]
	$\Delta_v H$	(273–310)	64.1 ± 0.6	298	ME	[2003ZAI/VER]
	$\Delta_v H$	(273–310)	59.8 ± 0.6	298	ME	[2003ZAI/VER]
	$\Delta_v H$	(278–313)	58.4 ± 0.7	298	GS	[2003ZAI/VER]
	$\Delta_v H$	(333–378)	60.0	298	CGC	[1999VER/HEI]
	$\Delta_v H$	(283–313)	60.0 ± 0.6	298	GS	[1996VER/BEC]
C ₁₀ H ₁₈ O ₂	[1129-47-1]	cyclohexyl isobutyrate				
	$\Delta_v H$	(333–378)	57.7	298	CGC	[1999VER/HEI]
C ₁₀ H ₁₈ O ₂	[na]	1-methylcyclohexyl propanoate				
	$\Delta_v H$	(333–378)	55.8	298	CGC	[1999VER/HEI]
C ₁₀ H ₁₈ O ₂	[na]	3-methylcyclohexyl propanoate				
	$\Delta_v H$	(333–378)	58.3	298	CGC	[1999VER/HEI]
C ₁₀ H ₁₈ O ₂	[na]	4-methylcyclohexyl propanoate				
	$\Delta_v H$	(333–378)	58.9	298	CGC	[1999VER/HEI]
C ₁₀ H ₁₈ O ₂	[56922-76-2]	pentyl 3-methylbut-2-enoate				
	$\Delta_v H$		61.8 ± 0.4	298	GS	[2008EME/TOK]
C ₁₀ H ₁₈ O ₂	[71697-84-2]	(–) 5-hydroxy- $\alpha,\alpha,4$ -trimethyl-3-cyclohexene-1-methanol (<i>trans</i> -sobrerol)				
	$\Delta_{\text{fus}} H$		34.69	423.6	DSC	[1999LI/ZEL]

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₀ H ₁₈ O ₂	[42370-41-2] $\Delta_{\text{fus}}H$	(+) 5-hydroxy- $\alpha,\alpha,4$ -trimethyl-3-cyclohexene-1-methanol (<i>trans</i> -sobrerol)	34.39	404.9	DSC	[1999LI/ZEL]
C ₁₀ H ₁₈ O ₂	[na] $\Delta_{\text{fus}}H$	(-) 5-hydroxy- $\alpha,\alpha,4$ -trimethyl-3-cyclohexene-1-methanol (<i>cis</i> -sobrerol)	23.18	382.9	DSC	[1999LI/ZEL]
C ₁₀ H ₁₈ O ₂	[54164-89-5] $\Delta_{\text{fus}}H$	(+) 5-hydroxy- $\alpha,\alpha,4$ -trimethyl-3-cyclohexene-1-methanol (<i>cis</i> -sobrerol)	25.86	378.9	DSC	[1999LI/ZEL]
C ₁₀ H ₁₈ O ₃	[na] Δ_vH	3-hydroxy-2,3-dimethyl-4-hexenoic acid, ethyl ester (362–387)	57.4	374	A	[1987STE/MAL]
C ₁₀ H ₁₈ O ₃	[na] Δ_vH Δ_vH	isopentyl levulinate (403–521)	59.4 56.3	418 461	A	[1987STE/MAL] [1931SCH/COW]
C ₁₀ H ₁₈ O ₃	[na] Δ_vH	1-ethylpropyl levulinate (397–513)	58.6	412	A	[1987STE/MAL]
C ₁₀ H ₁₈ O ₃	[na] Δ_vH	1-methylbutyl levulinate (397–513)	57.2	412	A	[1987STE/MAL]
C ₁₀ H ₁₈ O ₃	[na] Δ_vH	2-methylbutyl levulinate (391–473)	56.5	406	A	[1987STE/MAL]
C ₁₀ H ₁₈ O ₃	[20279-49-6] Δ_vH Δ_vH	pentyl levulinate (354–527)	66.3 56.2	369 466	A	[1987STE/MAL, 1947STU] [1931SCH/COW]
C ₁₀ H ₁₈ O ₃	[na] $\Delta_{\text{fus}}H$	1- <i>tert</i> -butyl-4-methyl-2,6,7-trioxabicyclo[2.2.2]octane	16.4	375.2		[1995RAK/VER2]
C ₁₀ H ₁₈ O ₄	[na] Δ_vH	pentyl 2-acetoxypropionate (312–501)	68.5	327	A	[1987STE/MAL]
C ₁₀ H ₁₈ O ₄	[141-28-6] Δ_vH	diethyl adipate (347–513)	57.5	362	A	[1987STE/MAL, 1947STU]
C ₁₀ H ₁₈ O ₄	[2050-61-5] Δ_vH	diisobutyl oxalate (336–503)	55.5	351	A	[1987STE/MAL, 1947STU]
C ₁₀ H ₁₈ O ₄	[925-15-5] Δ_vH	dipropyl succinate (350–524)	59.4	365	A	[1987STE/MAL, 1947STU]
C ₁₀ H ₁₈ O ₄	[2049-70-9] Δ_vH	diethyl ethylmethylmalonate (317–481)	53.2	332	A	[1987STE/MAL]
C ₁₀ H ₁₈ O ₄	[1732-09-8] Δ_vH	dimethyl suberate (293–352)	78.1 ± 0.2	298	GS	[2006VER/KOZ]
C ₁₀ H ₁₈ O ₄	[105-72-6] Δ_vH	ethylene glycol dibutyrate	73.2 ± 0.6	298	C	[1986NIL/WAD]
C ₁₀ H ₁₈ O ₄	[111-20-6] $\Delta_{\text{fus}}H$ $\Delta_{\text{fus}}H$ $\Delta_{\text{fus}}H$ $\Delta_{\text{fus}}H$ $\Delta_{\text{fus}}H$	sebacic acid	46.9 45.3 0.4 46.6 40.8	405.7 405.6 370.3 403.9 404	DSC	[2008VEN/BAY] [2008XIA/ZHA] [2005ROU/TEM] [1996DOM/HEA]
	$\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$	(353–385) (302–320)	181 ± 8 146.5		TPD TPTD	[2007CAP/LOV] [2005CHA/ZIE]
	$\Delta_{\text{sub}}H$	Note: Values based on TPTD method are not consistent with values determined by other experimental methods	165.3 ± 2.9			[1999RIB/MON, 1960DAV/THO]

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{sub}}H$	(375–403)	160.7 ± 2.5	389	ME	[1960DAV/THO, 1970COX/PIL, 1987STE/MAL]
	Δ_vH	(424–503)	124.8	298	CGC	[2005ROU/TEM]
	Δ_vH	(456–625)	85.9	471		[1947STU]
C ₁₀ H ₁₈ O ₅	[na]	ethyl[1-(butoxycarbonyl)ethyl] carbonate				
	Δ_vH	(324–473)	70.2	339	A	[1987STE/MAL]
C ₁₀ H ₁₈ O ₅	[na]	2-lactoyloxypropionic acid, butyl ester				
	Δ_vH	(336–407)	74.8	351	A	[1987STE/MAL]
C ₁₀ H ₁₈ O ₅	[na]	2-lactoyloxypropionic acid, sec-butyl ester				
	Δ_vH	(329–399)	74.3	344	A	[1987STE/MAL]
C ₁₀ H ₁₈ O ₅	[500790-23-8]	pentyl[1-(ethoxycarbonyl)methyl] carbonate				
	Δ_vH	(383–503)	68.2	398	A	[1987STE/MAL]
C ₁₀ H ₁₈ O ₅	[na]	pentyl[1-(methoxycarbonyl)ethyl] carbonate				
	Δ_vH	(360–524)	63.7	375	A	[1987STE/MAL]
C ₁₀ H ₁₈ O ₆	[62961-64-2]	<i>(d)</i> diisopropyl tartrate				
	Δ_vH	(376–548)	65.7	391	A	[1987STE/MAL, 1947STU]
C ₁₀ H ₁₈ O ₆	[2217-14-3]	<i>(d)</i> dipropyl tartrate				
	Δ_vH	(388–576)	71.8	403	A	[1987STE/MAL, 1947STU]
C ₁₀ H ₁₈ O ₆ S	[7355-12-6]	diethyl 3,3'-sulfonyldipropionate				
	$\Delta_{\text{fus}}H$		38.0	359.7		[1994WAN/KUO]
C ₁₀ H ₁₉ ClNO ₅ P	[13171-21-6]	phosphamidon				
	Δ_vH	(293–388)	90.1	308	A	[1987STE/MAL, 1999DYK/SVO]
C ₁₀ H ₁₉ Cl ₂ N	[4261-59-0]	N,N-bis(2-chloroethyl)cyclohexylamine				
	Δ_vH	(273–333)	62.4	288	A,GS	[1987STE/MAL, 1948RED/CHA3, 1999DYK/SVO]
C ₁₀ H ₁₉ N	[7335-04-8]	N-cyclopentylpiperidine				
	Δ_vH	(283–318)	54.9 ± 0.3	301	GS	[1998VER6]
	Δ_vH	(283–318)	55.1 ± 0.3	298	GS	[1998VER6]
C ₁₀ H ₁₉ N	[1975-78-6]	decanenitrile (caprinitrile)				
	Δ_vH	(295–326)	66.3 ± 0.4	298	GS	[2005EME/VER]
	Δ_vH	(381–519)	58.0	396	A	[1987STE/MAL]
	Δ_vH		66.8 ± 0.4	298	C	[1977STRI/SUN]
	Δ_vH	(381–431)	57.8	396	EB	[1971MEY/REN]
	Δ_vH	(431–518)	54.4	446	EB	[1971MEY/REN]
C ₁₀ H ₁₉ NO ₃	[na]	<i>(l)</i> N-acetylisoleucine, ethyl ester				
	Δ_vH	(391–476)	69.1	406	A	[1987STE/MAL]
C ₁₀ H ₁₉ NO ₃	[na]	<i>(l)</i> N-acetylleucine, ethyl ester				
	Δ_vH	(396–476)	74.8	411	A	[1987STE/MAL]
C ₁₀ H ₁₉ N ₅ O	[1610-18-0]	2-methoxy-4,6-bis(isopropylamino)-1,3,5-triazine				
	$\Delta_{\text{sub}}H$	(323–365)	92.2	338	GS-GC	[1987STE/MAL, 1964FRI/SAM]
C ₁₀ H ₁₉ N ₅ S	[7287-19-6]	2-methylthio-4,6-bis(isopropylamino)-1,3,5-triazine				
	$\Delta_{\text{sub}}H$	(323–393)	100	338	GS-GC	[1987STE/MAL, 1964FRI/SAM]
C ₁₀ H ₁₉ N ₅ S	[886-50-0]	2-methylmercapto-4-ethylamino-6- <i>tert</i> -butylamino-1,3,5-triazine (terbutryn)				
	$\Delta_{\text{fus}}H$		21	376.1		[2007VEC/BRU]
	$\Delta_{\text{fus}}H$		21.42	375.9	DSC	[1990DON/DRE]
	Δ_vH		87 ± 5	467	DSC	[2007VEC/BRU]

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$		115 ± 7	298	DSC	[2007VEC/BRU]
	$\Delta_v H$		83.2 ± 1.0	452	TGA	[2007VEC/BRU]
	$\Delta_v H$		111 ± 4	298	TGA	[2007VEC/BRU]
C₁₀H₁₉O₆PS₂	[121-75-5]	(<i>dl</i>) malathion (283–419)	71.1	298	A	[1987STE/MAL]
C₁₀H₁₉O₇PS	[na]	O,O-dimethyl-S-[1,2- <i>bis</i> (ethoxycarbonyl)ethyl]thiophosphate (283–406)	93.4	298	A	[1987STE/MAL]
C₁₀H₂₀	[2158-55-6]	1,1,4-trimethylcycloheptane 45.5 ± 0.2		298	C	[1996VAR/PAS]
	$\Delta_v H$					Note: Text in Ref. [1996VAR/PAS] states 1,1,4-trimethylcycloheptane; however, the molecular structure of 1,1,4-trimethylcyclohexane is given in the paper.
C₁₀H₂₀	[293-96-9]	cyclodecane				
	$\Delta_{\text{fus}} H$		18.95	282.7		[2005HUA/SIM]
	$\Delta_v H$	(404–489)	45.1	419	A,EB	[1987STE/MAL, 1976MEY/HOT]
	$\Delta_v H$	(343–386)	48.2	358	EB	[1987STE/MAL, 1976MEY/HOT]
C₁₀H₂₀	[1678-93-9]	butylcyclohexane				
	$\Delta_{\text{fus}} H$		14.2	198		[2006MAN/CUT]
	$\Delta_{\text{fus}} H$		14.14	198.4		[1996DOM/HEA]
	$\Delta_v H$	(274–313)	47.4 ± 0.2	294	GS	[1995CHI/HES]
	$\Delta_v H$		47.0 ± 0.2	298		[1995CHI/HES]
	$\Delta_v H$		48.9 ± 0.5	298	GC	[1987AZA]
	$\Delta_v H$		49.4 ± 0.4	298	GCC	[1978FUC/PEA]
	$\Delta_v H$		49.4	298		[1975KUS/SAI]
	$\Delta_v H$		50	298		[1971WIL/ZWO]
	$\Delta_v H$		50.0	298		[1965FIN/MES]
	$\Delta_v H$	(367–457)	44.9	382	A	[1987STE/MAL, 1949FOR/NOR]
C₁₀H₂₀	[7058-01-7]	sec-butylcyclohexane (369–455)	44.1	384	A	[1987STE/MAL, 1949FOR/NOR]
C₁₀H₂₀	[3178-22-1]	<i>tert</i> -butylcyclohexane				
	$\Delta_v H$		45.0 ± 0.1	328	C	[1981SVO/CHA]
	$\Delta_v H$		44.0 ± 0.1	343	C	[1981SVO/CHA]
	$\Delta_v H$		43.0 ± 0.1	358	C	[1981SVO/CHA]
	$\Delta_v H$		42.4 ± 0.1	368	C	[1981SVO/CHA]
	$\Delta_v H$	(355–446)	42.9	370	A	[1987STE/MAL, 1949FOR/NOR]
C₁₀H₂₀	[1678-98-4]	isobutylcyclohexane				
	$\Delta_v H$		47.5	298		[1975KUS/SAI]
	$\Delta_v H$	(355–446)	43.5	370	A	[1987STE/MAL, 1949FOR/NOR]
C₁₀H₂₀	[99-82-1]	1-isopropyl-4-methylcyclohexane (382–443)	43.6	297	A	[1987STE/MAL]
C₁₀H₂₀	[3741-00-2]	n-pentylcyclopentane	51.0	298		[1971WIL/ZWO]
C₁₀H₂₀	[872-05-9]	1-decene				
	$\Delta_{\text{fus}} H$		7.95	198.3		
	$\Delta_{\text{fus}} H$		13.81	206.9		[1996DOM/HEA]
	$\Delta_v H$	(383–445)	43.8	398	A	[1987STE/MAL]
	$\Delta_v H$		50.4 ± 0.2	298	C	[1977MAN/SEL]
	$\Delta_v H$		50.5	298		[1971WIL/ZWO]
	$\Delta_v H$	(360–445)	45.1	375		[1950FOR/CAM]
C₁₀H₂₀	[20348-51-0]	<i>cis</i> 2-decene				

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(401–447)	43.6	416	A	[1987STE/MAL]
C ₁₀ H ₂₀	[20063-97-2]	<i>trans</i> 2-decene				
	$\Delta_v H$	(401–447)	43.7	416	A	[1987STE/MAL]
C ₁₀ H ₂₀	[19398-86-8]	<i>cis</i> 3-decene				
	$\Delta_v H$	(398–444)	43.1	413	A	[1987STE/MAL]
C ₁₀ H ₂₀	[19150-21-1]	<i>trans</i> 3-decene				
	$\Delta_v H$	(398–445)	43.4	413	A	[1987STE/MAL]
C ₁₀ H ₂₀	[19398-88-0]	<i>cis</i> 4-decene				
	$\Delta_v H$	(397–444)	43.0	412	A	[1987STE/MAL]
C ₁₀ H ₂₀	[19398-89-1]	<i>trans</i> 4-decene				
	$\Delta_v H$	(398–444)	43.2	413	A	[1987STE/MAL]
C ₁₀ H ₂₀	[7433-78-5]	<i>cis</i> 5-decene				
	$\Delta_v H$	(397–443)	42.9	412	A	[1987STE/MAL]
C ₁₀ H ₂₀	[7433-56-9]	<i>trans</i> 5-decene				
	$\Delta_v H$	(398–444)	42.3	413	A	[1987STE/MAL]
C ₁₀ H ₂₀	[4485-13-6]	4-propyl-3-heptene				
	$\Delta_v H$	(333–371)	43.7	348	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₁₀ H ₂₀	[na]	<i>trans</i> 2,2,4,4-tetramethyl-3-hexene				
	$\Delta_v H$		42.0 ± 0.2	298	GCC	[1979BOT/CAM]
C ₁₀ H ₂₀	[22808-06-6]	2,2,5,5-tetramethylhex-3-ene				
	$\Delta_{\text{ms}} H$		1.21	235.8		
	$\Delta_{\text{ws}} H$		4.33	243.5		
	$\Delta_{\text{fus}} H$		10.25	268.9	DSC	[1980BYS]
C ₁₀ H ₂₀ Br ₂	[59104-80-2]	1,1-dibromodecane				
	$\Delta_v H$	(442–610)	62.2	457	A, EST	[1987STE/MAL, 1956MAN, 1970DYK/VAN, 1999DYK/SVO]
C ₁₀ H ₂₀ Br ₂	[28467-71-2]	1,2-dibromodecane				
	$\Delta_v H$	(368–524)	67.0	383	A	[1987STE/MAL, 1947STU, 1970DYK/VAN]
C ₁₀ H ₂₀ Cl ₂	[3162-62-7]	1,1-dichlorodecane				
	$\Delta_v H$	(415–577)	56.9	430	A, EST	[1987STE/MAL, 1956MAN, 1970DYK/VAN]
C ₁₀ H ₂₀ Cl ₂	[2162-98-3]	1,10-dichlorodecane				
	$\Delta_v H$	(441–520)	61.1	456		[1999DYK/SVO]
	$\Delta_v H$		67.3			[1998DRO/TOM]
	$\Delta_v H$	(440–540)	73.1	298		[1991BAS/SVO]
C ₁₀ H ₂₀ F ₂	[62127-43-9]	1,1-difluorodecane				
	$\Delta_v H$	(364–504)	50.2	379	A, EST	[1987STE/MAL, 1956MAN, 1970DYK/VAN, 1999DYK/SVO]
C ₁₀ H ₂₀ NO ₂	[na]	ethyl 2-(N,N-diethylamino)butanoate				
	$\Delta_v H$	(283–313)	57.3 ± 0.2	298	GS	[1996VER/ZUF]
C ₁₀ H ₂₀ N ₂ OS	[398995-30-7]	N,N-diethyl-N'-isovaleroylthiourea				
	$\Delta_{\text{sub}} H$	363	121.5 ± 3.2	298	C	[2001RIB/RIB]
C ₁₀ H ₂₀ N ₂ OS	[398995-31-8]	N,N-diethyl-N'-pivaloylthiourea				
	$\Delta_{\text{sub}} H$	366	114.9 ± 2.7	298	C	[2001RIB/RIB]
C ₁₀ H ₂₀ N ₂ O ₂	[na]	tetraethyloxamide				

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
		$\Delta_{\text{fus}}H$	17	310.2		[2003CLO/JAN]
		Δ_vH	63	464	TGA,DSC	[2003CLO/JAN]
C ₁₀ H ₂₀ N ₂ O ₂	[1740-54-1]	sebacamide				
		$\Delta_{\text{fus}}H$	68.7	484.3	DSC	[2006BAD/DEL]
C ₁₀ H ₂₀ N ₆	[16268-73-8]	1-(ethylmethylamino)-3,5-bis(dimethylamino)-s-triazine				
		$\Delta_{\text{fus}}H$	21.3	384	DSC	[1991ACR]
C ₁₀ H ₂₀ O	[106-22-9]	citronellol				
		$\Delta_{\text{sub}}H$	(283–333)	66.1	308	[1954SER/VOI, 1960JON]
		Δ_vH	(343–453)	63.5	298	GC [2005HOS/GRY]
C ₁₀ H ₂₀ O	[na]	bis(3-methyl-2-butenyl) ether				
		Δ_vH	(383–413)	47.8	398	[1989WAN/YIN]
C ₁₀ H ₂₀ O	[103-44-6]	(2-ethylhexyl) vinyl ether				
		Δ_vH	(330–451)	44.7	345	A [1987STE/MAL]
C ₁₀ H ₂₀ O	[5445-30-7]	1-butylcyclohexanol				
		Δ_vH	(362–481)	55.7	377	A [1987STE/MAL]
C ₁₀ H ₂₀ O	[106-22-9]	3,7-dimethyl-6-octene-1-ol				
		Δ_vH	(293–333)	72.6	308	A [1987STE/MAL]
		Δ_vH	(373–500)	65.9	388	A [1987STE/MAL]
C ₁₀ H ₂₀ O	[2216-51-5]	(<i>l</i>) menthol				
		$\Delta_{\text{fus}}H$	11.88	316.2	DTA	[1981CHI/GAR]
		$\Delta_{\text{sub}}H$	(279–299)	95.8 ± 4.8	289	HSA [1981CHI/GAR]
		Δ_vH	(372–488)	59.1	387	A [1987STE/MAL]
		Δ_vH	(329–485)	58.2	344	[1947STU]
C ₁₀ H ₂₀ O	[89-78-1]	(<i>dl</i>)-menthol				
		$\Delta_{\text{fus}}H$	10.25	301.2	DTA	[1981CHI/GAR]
		$\Delta_{\text{sub}}H$	(279–299)	78.6 ± 4	289	HSA [1981CHI/GAR]
C ₁₀ H ₂₀ O	[2216-51-5]	(–)-menthol				
		Δ_vH	(323–433)	56.6	298	GC [2005HOS/GRY]
C ₁₀ H ₂₀ O	[1502-05-2]	cyclodecanol				
		$\Delta_{\text{sub}}H$	(287–292)	100.5 ± 0.5	288	TM [1955ENG]
C ₁₀ H ₂₀ O	[na]	1-(1-methylcyclohexyl)-1-propanol				
		Δ_vH	(396–420)	55.4	408	A [1987STE/MAL]
C ₁₀ H ₂₀ O	[27331-02-8]	2-(1-methylcyclohexyl)-2-propanol				
		Δ_vH	(393–418)	53.0	405	A [1987STE/MAL]
C ₁₀ H ₂₀ O	[10340-22-4]	(<i>Z</i>) 3-decen-1-ol				
		Δ_vH	(323–363)	78.7	298	CGC [2000OVA/KOU, 1994KOU/HOS]
C ₁₀ H ₂₀ O	[10339-60-3]	(<i>E</i>) 3-decen-1-ol				
		Δ_vH	(323–363)	78.8	298	CGC [2000OVA/KOU, 1994KOU/HOS]
C ₁₀ H ₂₀ O	[57074-37-0]	(<i>Z</i>) 4-decen-1-ol				
		Δ_vH	(323–363)	79.6	298	CGC [2000OVA/KOU, 1994KOU/HOS]
C ₁₀ H ₂₀ O	[10339-62-5]	(<i>E</i>) 4-decen-1-ol				
		Δ_vH	(323–363)	80.3	298	CGC [2000OVA/KOU, 1994KOU/HOS]

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₀ H ₂₀ O	[51652-47-2] $\Delta_v H$	(Z) 5-decen-1-ol (323–363)	80.3	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₀ H ₂₀ O	[56578-18-8] $\Delta_v H$	(E) 5-decen-1-ol (323–363)	80.6	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₀ H ₂₀ O	[68760-59-8] $\Delta_v H$	(Z) 6-decen-1-ol (323–363)	80.3	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₀ H ₂₀ O	[38421-92-0] $\Delta_v H$	(E) 6-decen-1-ol (323–363)	80.6	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₀ H ₂₀ O	[16504-66-8] $\Delta_v H$	(Z) 7-decen-1-ol (323–363)	80.8	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₀ H ₂₀ O	[52957-12-7] $\Delta_v H$	(E) 7-decen-1-ol (323–363)	81.1	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₀ H ₂₀ O	[83799-67-1] $\Delta_v H$	(Z) 8-decen-1-ol (323–363)	81.6	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₀ H ₂₀ O	[83799-68-2] $\Delta_v H$	(E) 8-decen-1-ol (323–363)	81.5	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₀ H ₂₀ O	[693-54-9] $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$	2-decanone (317–484) (357–560) 60.9 ± 0.5 (358–568)	51.7 55.1 60.9 ± 0.5 44.6	332 372 298 487	A A GCC	[1987STE/MAL, 1947STU] [1987STE/MAL] [1979SAL/PEA] [1975AMB/ELL]
C ₁₀ H ₂₀ O	[868-91-7] $\Delta_v H$	2,2,5,5-tetramethyl-3-hexanone	48.8 ± 0.2	298	C	[1970SEL2]
C ₁₀ H ₂₀ O	[112-31-2] $\Delta_{\text{fus}} H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$	decanal (280–316) (308–353) (288–333) (293–358) 60.4 ± 0.3 (324–482)	30.6 59.5 ± 0.4 60.5 57.3 57.3 60.4 ± 0.3 56.3	268.2 298 298 303 308 298 339	GS CGC A A	[1980DYA/VAS] [2003VER/KRA2] [1996KOU/HOS, 2000OVA/KOU] [1987STE/MAL] [1987STE/MAL] [198IDYA/KOR] [1987STE/MAL, 1947STU]
C ₁₀ H ₂₀ O ₂	[4359-57-3] $\Delta_v H$	2-heptyl-1,3-dioxolane (318–453)	62.0	333	A	[1987STE/MAL]
C ₁₀ H ₂₀ O ₂	[4359-47-1] $\Delta_v H$	2-(1-ethylpentyl)-1,3-dioxolane (333–453)	55.3	348	A	[1987STE/MAL]
C ₁₀ H ₂₀ O ₂	[2244-85-1] $\Delta_v H$	4-hexyl-1,3-dioxane (318–453)	56.9	333	A	[1987STE/MAL]
C ₁₀ H ₂₀ O ₂	[61827-60-9] $\Delta_v H$	3-pentyl-4-hydroxytetrahydropyran (383–453)	72.6	398	A	[1987STE/MAL]
C ₁₀ H ₂₀ O ₂	[859773-58-3] $\Delta_v H$	2-butoxy-3-hexanone (333–418)	39.5	348	A	[1987STE/MAL, 1933HEN/MUR]
C ₁₀ H ₂₀ O ₂	[107-75-5] $\Delta_v H$	hydroxycitronellal (283–333)	75.3	298	A,ME	[1987STE/MAL, 1955SER/VOI]
C ₁₀ H ₂₀ O ₂	[112-14-1] $\Delta_v H$ $\Delta_v H$	octyl acetate (274–309)	60.7 ± 0.4 61.7	298 298	GS GC	[2006KRA/VER] [1997DEF/CAR]

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(334–417)	54.9	349	A	[1987STE/MAL]
	$\Delta_v H$	(345–472)	47.8	360	A	[1987STE/MAL]
C ₁₀ H ₂₀ O ₂	[106-32-1]	ethyl octanoate				
	$\Delta_v H$	(382–412)	52.5 ± 0.2	397	EB	[1991WIB/WAL]
	$\Delta_v H$	(382–412)	59.5 ± 1.3	298	EB	[1991WIB/WAL]
	$\Delta_v H$	(330–480)	53.2	345	A	[1987STE/MAL]
C ₁₀ H ₂₀ O ₂	[103-09-3]	2-ethylhexylacetate				
	$\Delta_v H$	(333–472)	50.1	348	A	[1987STE/MAL]
C ₁₀ H ₂₀ O ₂	[659-70-1]	isopentyl isovalerate				
	$\Delta_v H$	(341–479)	46.4	356	A	[1987STE/MAL]
	$\Delta_v H$	(300–467)	47.2	315		[1947STU]
C ₁₀ H ₂₀ O ₂	[5340-26-1]	neopentyl pivalate				
	$\Delta_v H$	(280–310)	49.1 ± 0.5	295	GS	[1999VER/HEI]
	$\Delta_v H$	(280–310)	48.9 ± 0.5	298	GS	[1999VER/HEI]
C ₁₀ H ₂₀ O ₂	[89397-96-6]	2,2-dimethylpropanoic acid, 1,1-dimethylpropyl ester				
	$\Delta_v H$	(333–378)	48.0	298	CGC	[1999VER/HEI]
C ₁₀ H ₂₀ O ₂	[245658-27-9]	2-methylpropanoic acid, 1,1-dimethylbutyl ester				
	$\Delta_v H$	(333–378)	51.4	298	CGC	[1999VER/HEI]
C ₁₀ H ₂₀ O ₂	[245658-32-6]	2-methylpropanoic acid, 1,1,2-trimethylpropyl ester				
	$\Delta_v H$	(333–378)	51.7	298	CGC	[1999VER/HEI]
C ₁₀ H ₂₀ O ₂	[na]	butanoic acid, 1,1,2-trimethylpropyl ester				
	$\Delta_v H$	(333–378)	53.8	298	CGC	[1999VER/HEI]
C ₁₀ H ₂₀ O ₂	[245658-37-1]	2,4-dimethyl-2-pentanol, propanoate				
	$\Delta_v H$	(333–378)	51.7	298	CGC	[1999VER/HEI]
C ₁₀ H ₂₀ O ₂	[1731-84-6]	methyl nonanoate				
	$\Delta_v H$		57.4	350		[2002VAN/VAN]
	$\Delta_v H$		56.7 ± 0.3	364		[2002VAN/VAN]
	$\Delta_v H$		61.6 ± 0.4	298		[2002VAN/VAN]
	$\Delta_v H$		57.7 ± 0.7	298	GC	[1987AZA]
	$\Delta_v H$		62.0 ± 0.5	298	GCC	[1980FUC/PEA]
	$\Delta_v H$		62.0 ± 0.4	298	C	[1977MAN/SEL]
	$\Delta_v H$	(364–439)	55.6	379	A,EST	[1987STE/MAL, 1963ROS/SCH]
C ₁₀ H ₂₀ O ₂	[334-48-5]	decanoic acid (capric acid)				
	$\Delta_{\text{fus}} H$		28.3	303.8	DSC	[2007MOR/COR]
	$\Delta_{\text{fus}} H$		27.82	304.5		[1996DOM/HEA]
	$\Delta_{\text{sub}} H$		129.6 ± 5	298	TPD	[2008CAP/LOV]
	$\Delta_{\text{sub}} H$	(293–303)	118.8 ± 2.2	298	ME	[1968BAC/NOV, 1970COX/PIL, 1987STE/MAL]
	$\Delta_{\text{sub}} H$	(290–301)	117.1 ± 1.7	295	ME	[1961DAV/MAL]
	$\Delta_v H$	(398–543)	76.4	413	A	[1987STE/MAL]
	$\Delta_v H$	(305–323)	88.6	314	ME,TE	[1982DEK/SCH]
	$\Delta_v H$		71.4	418	I	[1943CRA]
C ₁₀ H ₂₀ O ₃	[869190-73-8]	propyl 3-butoxypropionate				
	$\Delta_v H$	(373–473)	44.2	388	A	[1987STE/MAL, 1933HEN/MUR]
C ₁₀ H ₂₀ O ₃	[14144-36-6]	pentyl 3-ethoxypropionate				
	$\Delta_v H$	(374–498)	54.1	389	A	[1987STE/MAL]
C ₁₀ H ₂₀ O ₃	[7419-97-8]	methyl 3-hexyloxypropionate				
	$\Delta_v H$	(373–473)	55.1	388	A	[1987STE/MAL]

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₀ H ₂₀ O ₃	[14156-10-6]	peroxydecanoic acid				
	$\Delta_{\text{sub}}H$	(293–303)	117.1 ± 0.8	298	ME	[1980SWA/KWA]
C ₁₀ H ₂₀ O ₄	[124-17-4]	diethylene glycol monobutyl ether acetate				
	Δ_vH	(393–520)	57.7	408	A	[1987STE/MAL]
C ₁₀ H ₂₀ O ₄	[33785-99-8]	3,3,6,6-tetraethyl-1,2,4,5-tetraoxacyclohexane				
	Δ_vH	(403–473)	50.1	298	CGC	[2007CAN/EYL]
C ₁₀ H ₂₀ N ₆ O	[52298-71-2]	1-(2-hydroxyethylmethylamino)-3,5-bis(dimethylamino)-s-triazine				
	$\Delta_{\text{fus}}H$		17.32	373.3		[1989BRA/RYT]
C ₁₀ H ₂₀ O ₅	[33100-27-5]	1,4,7,10,13-pentaoxacyclopentadecane (15-crown-5)				
	Δ_vH		75.7 ± 1.7	298	CGC	[2000NIC/ORF]
	Δ_vH		79.6 ± 0.3	298	C	[1982BYS/MAN]
C ₁₀ H ₂₀ S ₄	[24194-61-4]	1,4,8,11-tetrathiacyclotetradecane				
	$\Delta_{\text{trs}}H$		2.7	345.2		
	$\Delta_{\text{fus}}H$		33.0	393.2	DSC	[2002ROC/GRI]
C ₁₀ H ₂₀ S ₅	[36338-04-2]	1,4,7,10,13-pentathiacyclopentadecane				
	$\Delta_{\text{trs}}H$		11.0	318.2		
	$\Delta_{\text{trs}}H$		4.2	340.2		
	$\Delta_{\text{fus}}H$		17.0	391.2	DSC	[2002ROC/GRI]
C ₁₀ H ₂₁ Br	[112-29-8]	1-bromodecane				
	Δ_vH	(391–545)	56.1	406		[1999DYK/SVO]
	Δ_vH	(383–570)	56.6	398	A,EST	[1987STE/MAL, 1961LI/ROS]
C ₁₀ H ₂₁ Cl	[1002-69-3]	1-chlorodecane				
	Δ_vH		64.0 ± 0.2	298	GS	[2001PUR/CHI]
	Δ_vH	(379–530)	54.4	394		[1999DYK/SVO]
	Δ_vH	(359–499)	56.2	374	A,DTA	[1987STE/MAL, 1969KEM/KRE]
C ₁₀ H ₂₁ F	[334-56-5]	1-fluorodecane				
	Δ_vH	(342–503)	50.4	357	A	[1987STE/MAL, 1961LI/ROS]
C ₁₀ H ₂₁ I	[2050-77-3]	1-iododecane				
	Δ_vH	(397–598)	69.8	298	A,EST	[1987STE/MAL, 1961LI/ROS, 2006BOL/NER]
	Δ_vH	(407–571)	57.4	422		[1999DYK/SVO]
	Δ_vH	(397–598)	58.1	412	A,EST	[1987STE/MAL, 1961LI/ROS]
C ₁₀ H ₂₁ N	[101-40-6]	N,α-dimethylcyclohexanethylamine				
	Δ_vH	(270–300)	50.2	285	A	[1987STE/MAL]
C ₁₀ H ₂₁ NO	[6282-97-9]	N,N-diethylhexanamide				
	Δ_vH	(373–443)	47.7	388	A	[1987STE/MAL]
C ₁₀ H ₂₁ NO	[2319-29-1]	decanamide				
	$\Delta_{\text{trs}}H$		1.05	218.7		
	$\Delta_{\text{trs}}H$		18.8	366.6		
	$\Delta_{\text{fus}}H$		15.1	370.6	DSC	[2008ABA/BAD]
	$\Delta_{\text{sub}}H$	(353–370)	125.9 ± 1.3	361.5	ME	[1959DAV/JON2]
C ₁₀ H ₂₂	[124-18-5]	decane				
	$\Delta_{\text{fus}}H$		27.6	243	DSC	[2004MAR/KAI]
	$\Delta_{\text{fus}}H$		28.7	243.5		[1996DOM/HEA]
	$\Delta_{\text{sub}}H$		80.3	298	B	[1980SWA/KWA]
	$\Delta_{\text{sub}}H$		84.8	243	B	[1963BON]
	$\Delta_{\text{sub}}H$		82.4	298	H	[1963BON, 1993CHI/HOS]

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(324–402)	48.3	339	GC	[2007MOK/RAZ]
	$\Delta_v H$	(337–376)	46.6	352		[2002BAT]
	$\Delta_v H$		51.1 ± 3.9	298	CGC	[2000NIC/ORF]
	$\Delta_v H$		51.5	299	C	[1996VIT/CHA]
	$\Delta_v H$		50.5	314	C	[1996VIT/CHA]
	$\Delta_v H$		50.1	324	C	[1996VIT/CHA]
	$\Delta_v H$		49.2	334	C	[1996VIT/CHA]
	$\Delta_v H$	(403–453)	50.9	298	CGC	[1995CHI/HOS]
	$\Delta_v H$	(423–473)	51.5	298	CGC	[1995CHI/HOS]
	$\Delta_v H$		51.4	298		[1994RUZ/MAJ]
	$\Delta_v H$	(409–584)	42.5	424		[1992LEE/DEM]
	$\Delta_v H$	(268–490)	48.1	340	EB,IP	[1989CHI/NGU]
	$\Delta_v H$	(268–490)	51.4	298	EB,IP	[1989CHI/NGU]
	$\Delta_v H$	(252–383)	53.8	267	A	[1987STE/MAL]
	$\Delta_v H$	(447–526)	41.7	462	A	[1987STE/MAL]
	$\Delta_v H$	(524–617)	38.6	539	A	[1987STE/MAL]
	$\Delta_v H$	(298–347)	50.3	313	GS	[1986ALL/JOS]
	$\Delta_v H$	(308–351)	49.8 ± 1.7			[1984BEC/RUC]
	$\Delta_v H$		51.4 ± 0.1	298	C	[1982FUR/SAK]
	$\Delta_v H$	(243–310)	55.9	258		[1973CAR/KOB]
	$\Delta_v H$		51.4	298		[1971WIL/ZWO]
	$\Delta_v H$	(373–443)	45.3	388		[1987STE/MAL, 1970VAR/BEL]
	$\Delta_v H$		51.4	298	C	[1947OSB/GIN]
	$\Delta_v H$	(368–440)	45.5	383	MM	[1945WIL/TAY]
C₁₀H₂₂	[871-83-0]	2-methylnonane				
	$\Delta_{\text{fus}}H$		17.49	198.8		[1996DOM/HEA]
	$\Delta_v H$	(324–441)	46.4 ± 0.2	339	A	[1987STE/MAL]
	$\Delta_v H$		47.3 ± 0.2	328	C	[1984MAJ/SVO3]
	$\Delta_v H$		46.2 ± 0.2	343	C	[1984MAJ/SVO3]
	$\Delta_v H$		45.0 ± 0.2	358	C	[1984MAJ/SVO3]
	$\Delta_v H$		51.0	298		[1971WIL/ZWO]
C₁₀D₂₂	[16416-29-8]	decane-d ₂₂				
	$\Delta_v H$		51.8	298	CGC	[2008ZHA/UNH]
C₁₀H₂₂	[5911-04-6]	3-methylnonane				
	$\Delta_{\text{fus}}H$		18.7	188.5		[1996DOM/HEA]
	$\Delta_v H$		47.3 ± 0.2	328	C	[1984MAJ/SVO3]
	$\Delta_v H$		46.2 ± 0.2	343	C	[1984MAJ/SVO3]
	$\Delta_v H$		45.1 ± 0.2	358	C	[1984MAJ/SVO3]
	$\Delta_v H$		50.2	298		[1971WIL/ZWO]
C₁₀H₂₂	[17301-94-9]	4-methylnonane				
	$\Delta_{\text{fus}}H$		15.19	174.7		[1996DOM/HEA]
	$\Delta_v H$		49.5	298		[1961LAB/GRE]
C₁₀H₂₂	[15869-85-9]	5-methylnonane				
	$\Delta_{\text{fus}}H$		16.65	186.7		[1996DOM/HEA]
	$\Delta_v H$		47.0 ± 0.2	328	C	[1984MAJ/SVO3]
	$\Delta_v H$		45.9 ± 0.2	343	C	[1984MAJ/SVO3]
	$\Delta_v H$		44.6 ± 0.2	358	C	[1984MAJ/SVO3]
	$\Delta_v H$		49.8	298		[1971WIL/ZWO]
C₁₀H₂₂	[5881-17-4]	3-ethyloctane				
	$\Delta_v H$		49.0	298		[1971WIL/ZWO]
C₁₀H₂₂	[15869-86-0]	4-ethyloctane				
	$\Delta_v H$		48.1	298		[1971WIL/ZWO]

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₀ H ₂₂	[3178-29-8]	4-propylheptane				
	Δ_vH	(331–430)	48.5	298		[1971WIL/ZWO]
C ₁₀ H ₂₂	[52896-87-4]	4-isopropylheptane				
	Δ_vH		47.3	298	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₁₀ H ₂₂	[15869-87-1]	2,2-dimethyloctane	49.0	298		[1971WIL/ZWO]
C ₁₀ H ₂₂	[7146-60-3]	2,3-dimethyloctane	48.1	298		[1971WIL/ZWO]
C ₁₀ H ₂₂	[4032-94-4]	2,4-dimethyloctane				
	Δ_vH		44.9 ± 0.2	328	C	[1984MAJ/SVO2]
	Δ_vH		43.6 ± 0.2	343	C	[1984MAJ/SVO2]
	Δ_vH		42.4 ± 0.2	358	C	[1984MAJ/SVO2]
C ₁₀ H ₂₂	[15869-89-3]	2,5-dimethyloctane	49.0	298		[1971WIL/ZWO]
C ₁₀ H ₂₂	[2051-30-1]	2,6-dimethyloctane	49.3	298		[1971WIL/ZWO]
C ₁₀ H ₂₂	[1072-16-8]	2,7-dimethyloctane				
	Δ_vH	(279–433)	47.7	298	A	[1971WIL/ZWO]
C ₁₀ H ₂₂	[4110-44-5]	3,3-dimethyloctane	48.5	298		[1971WIL/ZWO]
C ₁₀ H ₂₂	[15869-92-8]	3,4-dimethyloctane	48.1	298		[1971WIL/ZWO]
C ₁₀ H ₂₂	[15869-93-9]	3,5-dimethyloctane	48.5	298		[1971WIL/ZWO]
C ₁₀ H ₂₂	[15869-94-0]	3,6-dimethyloctane	47.3	298		[1971WIL/ZWO]
C ₁₀ H ₂₂	[15869-95-1]	4,4-dimethyloctane	48.1	298		[1971WIL/ZWO]
C ₁₀ H ₂₂	[15869-96-2]	4,5-dimethyloctane	48.5	298		[1971WIL/ZWO]
C ₁₀ H ₂₂	[14676-29-0]	2-methyl-3-ethylheptane	48.1	298		[1971WIL/ZWO]
C ₁₀ H ₂₂	[52896-88-5]	2-methyl-4-ethylheptane	47.3	298		[1971WIL/ZWO]
C ₁₀ H ₂₂	[13475-78-0]	2-methyl-5-ethylheptane	48.1	298		[1971WIL/ZWO]
C ₁₀ H ₂₂	[17302-01-1]	3-methyl-3-ethylheptane	47.7	298		[1971WIL/ZWO]
C ₁₀ H ₂₂	[52896-89-6]	3-methyl-4-ethylheptane	47.7	298		[1971WIL/ZWO]
C ₁₀ H ₂₂	[52896-90-0]	3-methyl-5-ethylheptane	47.7	298		[1971WIL/ZWO]

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₀ H ₂₂	[52896-91-0] $\Delta_v H$	4-methyl-3-ethylheptane	48.1	298		[1971WIL/ZWO]
C ₁₀ H ₂₂	[17302-04-4] $\Delta_v H$	4-methyl-4-ethylheptane	47.2	298		[1961LAB/GRE]
C ₁₀ H ₂₂	[52896-92-1] $\Delta_v H$	2,2,3-trimethylheptane	46.9	298		[1971WIL/ZWO]
C ₁₀ H ₂₂	[14720-74-2] $\Delta_v H$	2,2,4-trimethylheptane	45.6	298		[1971WIL/ZWO]
C ₁₀ H ₂₂	[20291-95-6] $\Delta_v H$	2,2,5-trimethylheptane	46.0	298		[1971WIL/ZWO]
C ₁₀ H ₂₂	[1190-83-6] $\Delta_v H$	2,2,6-trimethylheptane	46.4	298		[1971WIL/ZWO]
C ₁₀ H ₂₂	[52896-93-2] $\Delta_v H$	2,3,3-trimethylheptane	46.9	298		[1971WIL/ZWO]
C ₁₀ H ₂₂	[52896-95-4] $\Delta_v H$	2,3,4-trimethylheptane	47.3	298		[1971WIL/ZWO]
C ₁₀ H ₂₂	[20278-85-7] $\Delta_v H$	2,3,5-trimethylheptane	47.3	298		[1971WIL/ZWO]
C ₁₀ H ₂₂	[4032-93-3] $\Delta_v H$	2,3,6-trimethylheptane	47.3	298		[1971WIL/ZWO]
C ₁₀ H ₂₂	[4032-92-2] $\Delta_v H$	2,4,4-trimethylheptane	45.2	298		[1971WIL/ZWO]
C ₁₀ H ₂₂	[20278-84-6] $\Delta_v H$	2,4,5-trimethylheptane	46.9	298		[1971WIL/ZWO]
C ₁₀ H ₂₂	[2613-61-8] $\Delta_v H$	2,4,6-trimethylheptane	46.4	298		[1971WIL/ZWO]
C ₁₀ H ₂₂	[1189-99-7] $\Delta_v H$	2,5,5-trimethylheptane	46.0	298		[1971WIL/ZWO]
C ₁₀ H ₂₂	[20278-88-0] $\Delta_v H$	3,3,4-trimethylheptane	46.9	298		[1971WIL/ZWO]
C ₁₀ H ₂₂	[7154-80-5] $\Delta_v H$	3,3,5-trimethylheptane	46.0	298		[1971WIL/ZWO]
C ₁₀ H ₂₂	[20278-88-0] $\Delta_v H$	3,4,4-trimethylheptane	46.4	298		[1971WIL/ZWO]
C ₁₀ H ₂₂	[20278-89-1] $\Delta_v H$	3,4,5-trimethylheptane	47.3	298		[1971WIL/ZWO]
C ₁₀ H ₂₂	[62016-13-1] $\Delta_v H$	2-methyl-3-isopropylhexane	46.4	298		[1971WIL/ZWO]
C ₁₀ H ₂₂	[17302-02-2] $\Delta_v H$	3,3-diethylhexane	47.3	298		[1971WIL/ZWO]
C ₁₀ H ₂₂	[19398-77-7] $\Delta_v H$	3,4-diethylhexane	47.7	298		[1971WIL/ZWO]
C ₁₀ H ₂₂	[20291-91-2] $\Delta_v H$	2,2-dimethyl-3-ethylhexane	46.0	298		[1971WIL/ZWO]

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₀ H ₂₂	[52896-99-8] $\Delta_v H$	2,2-dimethyl-4-ethylhexane	45.2	298		[1971WIL/ZWO]
C ₁₀ H ₂₂	[52897-00-4] $\Delta_v H$	2,3-dimethyl-3-ethylhexane	46.9	298		[1971WIL/ZWO]
C ₁₀ H ₂₂	[52897-01-5] $\Delta_v H$	2,3-dimethyl-4-ethylhexane	46.9	298		[1971WIL/ZWO]
C ₁₀ H ₂₂	[7220-26-0] $\Delta_v H$	2,4-dimethyl-3-ethylhexane	46.9	298		[1971WIL/ZWO]
C ₁₀ H ₂₂	[52897-03-7] $\Delta_v H$	2,4-dimethyl-4-ethylhexane	46.4	298		[1971WIL/ZWO]
C ₁₀ H ₂₂	[52897-04-8] $\Delta_v H$	2,5-dimethyl-3-ethylhexane	46.4	298		[1971WIL/ZWO]
C ₁₀ H ₂₂	[52897-05-9] $\Delta_v H$	3,3-dimethyl-4-ethylhexane	46.4	298		[1971WIL/ZWO]
C ₁₀ H ₂₂	[52897-06-0] $\Delta_v H$	3,4-dimethyl-3-ethylhexane	46.4	298		[1971WIL/ZWO]
C ₁₀ H ₂₂	[13475-81-5] $\Delta_v H$	2,2,3,3-tetramethylhexane	45.2	298		[1971WIL/ZWO]
C ₁₀ H ₂₂	[52897-08-2] $\Delta_v H$	2,2,3,4-tetramethylhexane	45.6	298		[1971WIL/ZWO]
C ₁₀ H ₂₂	[52897-09-3] $\Delta_v H$	2,2,3,5-tetramethylhexane	45.2	298		[1971WIL/ZWO]
C ₁₀ H ₂₂	[51750-65-3] $\Delta_v H$	2,2,4,4-tetramethylhexane	43.5	298		[1971WIL/ZWO]
C ₁₀ H ₂₂	[16747-42-5] $\Delta_v H$	2,2,4,5-tetramethylhexane	44.4	298		[1971WIL/ZWO]
C ₁₀ H ₂₂	[1071-81-4] $\Delta_v H$	2,2,5,5-tetramethylhexane	43.5	298		[1971WIL/ZWO]
C ₁₀ H ₂₂	[52897-10-6] $\Delta_v H$	2,3,3,4-tetramethylhexane	46.4	298		[1971WIL/ZWO]
C ₁₀ H ₂₂	[52897-11-7] $\Delta_v H$	2,3,3,5-tetramethylhexane	45.2	298		[1971WIL/ZWO]
C ₁₀ H ₂₂	[52897-12-8] $\Delta_v H$	2,3,4,4-tetramethylhexane	46.0	298		[1971WIL/ZWO]
C ₁₀ H ₂₂	[52897-15-1] $\Delta_v H$	2,3,4,5-tetramethylhexane	46.0	298		[1971WIL/ZWO]
C ₁₀ H ₂₂	[5171-84-6] $\Delta_v H$	3,3,4,4-tetramethylhexane	42.3	298		[1971WIL/ZWO]
C ₁₀ H ₂₂	[13475-79-1] $\Delta_v H$	2,4-dimethyl-3-isopropylpentane	45.6	298		[1971WIL/ZWO]
C ₁₀ H ₂₂	[52897-16-2] $\Delta_v H$	2-methyl-3,3-diethylpentane	47.3	298		[1971WIL/ZWO]
C ₁₀ H ₂₂	[52897-17-3] $\Delta_v H$	2,2,3-trimethyl-3-ethylpentane	46.0	298		[1971WIL/ZWO]

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		T_m (K)	Method	Reference
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)			
C ₁₀ H ₂₂	[52897-18-4] $\Delta_v H$	2,2,4-trimethyl-3-ethylpentane	44.8	298		[1971WIL/ZWO]
C ₁₀ H ₂₂	[52897-19-5] $\Delta_v H$	2,3,4-trimethyl-3-ethylpentane	46.4	298		[1971WIL/ZWO]
C ₁₀ H ₂₂	[16747-44-7] $\Delta_v H$	2,2,3,3,4-pentamethylpentane	45.2	298		[1971WIL/ZWO]
C ₁₀ H ₂₂	[16747-45-8] $\Delta_v H$	2,2,3,4,4-pentamethylpentane	43.5	298		[1971WIL/ZWO]
C ₁₀ H ₂₂ N ₂ O	[28141-55-1] $\Delta_{\text{fus}} H$	1-nonyl urea	38.9	380.3	DSC	[2005HAS/TAJ]
C ₁₀ H ₂₂ O	[69775-79-7] $\Delta_v H$	hexyl <i>tert</i> -butyl ether	53.2	298	CGC	[2002VER, 2003VER/KRA]
C ₁₀ H ₂₂ O	[na] $\Delta_v H$	pentyl <i>tert</i> -amyl ether	53.5	298	CGC	[2002VER, 2003VER/KRA]
C ₁₀ H ₂₂ O	[na] $\Delta_v H$	ethyl <i>tert</i> -octyl ether	45.3 ± 0.3	298	GS	[2002VER, 2003VER/KRA]
C ₁₀ H ₂₂ O	[693-65-2] $\Delta_v H$ $\Delta_v H$	dipentyl ether (373–460) (423–480)	46.2 45.6	388 451	A	[1987STE/MAL] [1968LAP/NIS]
C ₁₀ H ₂₂ O	[54459-71-1] $\Delta_v H$	butyl hexyl ether	53.2 ± 0.1	298	C	[1985KUS]
C ₁₀ H ₂₂ O	[544-01-4] $\Delta_v H$ $\Delta_v H$ $\Delta_v H$	diisopentyl ether (353–393) (417–470) (291–447)	51.4 41.4 47.6	298 443 306	CGC A	[1995CHI/HOS] [1968LAP/NIS] [1987STE/MAL, 1947STU]
C ₁₀ H ₂₂ O	[112-30-1] $\Delta_{\text{fus}} H$ $\Delta_{\text{fus}} H$	1-decanol	33.67 37.66	280 280.1		[2003VAN/GAB] [1997DOM/GON]
	$\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$	(264–273)	115.5 ± 6.3 112.5 ± 6.3	268 298	ME	[1965KAR/KYB, 1987STE/MAL] [1965KAR/KYB]
	$\Delta_v H$	(281–327)	79.5	309	GS	[2001KUL/VER2]
	$\Delta_v H$	(281–327)	80.9	298	GS	[2001KUL/VER2]
	$\Delta_v H$	(278–378)	81.1	293		[1999NGU/BER]
	$\Delta_v H$	(373–423)	81.7	298	CGC	[1995CHI/HOS]
	$\Delta_v H$	(353–393)	79.3	298	CGC	[1994KOU/HOS, 2000OVA/KOU]
	$\Delta_v H$	(283–388)	75.4	336		[1992NGU/KAS]
	$\Delta_v H$	(349–410)	71.6	364	A	[1987STE/MAL]
	$\Delta_v H$	(405–528)	62.6	420	A	[1987STE/MAL]
	$\Delta_v H$	(474–529)	53.9	489	A	[1987STE/MAL]
	$\Delta_v H$		78.2 ± 0.8	323	C	[1979SEV]
	$\Delta_v H$		81.5 ± 0.8	298	C	[1979SEV]
	$\Delta_v H$		81.5 ± 0.8	298	C	[1977MAN/SEL]
	$\Delta_v H$	(298–325)	77.6	313		[1973WIL/ZWO]
	$\Delta_v H$	(400–529)	63.5	415	A,EB	[1987STE/MAL, 1970AMB/SPR]
	$\Delta_v H$	(378–504)	69.5	393	DTA	[1969KEM/KRE]
	$\Delta_v H$	(298–325)	77.6	311	ME	[1965DAV/KYB]
	$\Delta_v H$	(364–461)	69.6	379		[1958ROS/PAP]
C ₁₀ H ₂₂ O	[106-21-8] $\Delta_v H$	(<i>dl</i>) 3,7-dimethyl-1-octanol (341–467)	79.1	356	A	[1987STE/MAL]

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₀ H ₂₂ O ₂	[112-48-1]	ethylene glycol dibutyl ether				
	$\Delta_v H$	(356–476)	55.9	371	A	[1987STE/MAL]
			58.8 ± 0.1	298	C	[1970KUS/WAD]
C ₁₀ H ₂₂ O ₂	[5669-09-0]	ethylene glycol diisobutyl ether				
	$\Delta_v H$	(336–456)	46.1	351	A	[1987STE/MAL]
C ₁₀ H ₂₂ O ₂	[1559-35-9]	ethylene glycol mono(2-ethylhexyl) ether				
	$\Delta_v H$	(381–502)	56.5	396	A	[1987STE/MAL]
C ₁₀ H ₂₂ O ₂	[871-22-7]	acetaldehyde dibutyl ether				
	$\Delta_v H$	(303–464)	47.3	318	A	[1987STE/MAL]
C ₁₀ H ₂₂ O ₂	[6931-71-1]	3,4-diethyl-3,4-hexanediol				
	$\Delta_v H$	(405–507)	54.7	420	A,EB	[1987STE/MAL, 1979BAL/FRI]
C ₁₀ H ₂₂ O ₂	[na]	3-ethyl-3-hydroxymethyl-2-heptanol				
	$\Delta_v H$	(338–500)	63.4	353	A	[1987STE/MAL]
C ₁₀ H ₂₂ O ₂	[112-47-0]	1,10-decanediol				
	$\Delta_{\text{fus}} H$		41.7	345.5		[1991ACR]
	$\Delta_{\text{sub}} H$		155.8 ± 0.9	298	C	[1990KNA/SAB]
	$\Delta_v H$		126.6 ± 4.2	298		[1994STE/CHI2, 2006UMN/KWE]
	$\Delta_v H$		112.4	364		[1993PIA/FER, 2006UMN/KWE]
	$\Delta_v H$		120.4 ± 4.9	298		[1993PIA/FER, 2006UMN/KWE]
C ₁₀ H ₂₂ O ₂ S	[126835-71-0]	3-(heptylthio)-1,2-propanediol				
	$\Delta_{\text{us}} H$		27.3	289.5		
			1.7	292.5	DSC	[1993ACR]
C ₁₀ H ₂₂ O ₃	[112-59-4]	diethylene glycol monoheptyl ether				
	$\Delta_v H$	(403–423)	86.5	298	EB	[2004CHY/FRA2]
			62.7	421	A	[1987STE/MAL]
C ₁₀ H ₂₂ O ₃	[24083-03-2]	dipropylene glycol monobutyl ether				
	$\Delta_v H$	(337–500)	63.2	352	A	[1987STE/MAL, 1947STU]
C ₁₀ H ₂₂ O ₃	[127748-41-8]	3-(heptyloxy)-1,2-propanediol				
	$\Delta_{\text{fus}} H$		28.8	288	DSC	[1993ACR]
C ₁₀ H ₂₂ O ₄	[20324-33-8]	tripropylene glycol monomethyl ether				
	$\Delta_v H$	(308–515)	58.7	323	A	[1987STE/MAL]
C ₁₀ H ₂₂ O ₅	[143-24-8]	tetraethylene glycol dimethyl ether (tetraglyme)				
	$\Delta_v H$		76.9 ± 2.6	298	CGC	[2000NIC/ORF]
			58	434	A	[1987STE/MAL]
C ₁₀ H ₂₂ S	[3698-94-0]	1-ethylthiooctane				
	$\Delta_v H$	(384–545)	63.9 ± 0.6	298	EB	[1996STE/CHI]
C ₁₀ H ₂₂ S	[143-10-2]	1-decanethiol				
	$\Delta_{\text{fus}} H$		33.3	247.9		[1996DOM/HEA]
	$\Delta_v H$	(390–544)	56.4	405		[1999DYK/SVO]
	$\Delta_v H$	(283–293)	58.6	288	A	[1987STE/MAL]
	$\Delta_v H$	(413–534)	54.6	428	A	[1987STE/MAL]
			65.5 ± 0.5	298	C	[1977MAN/SEL]
C ₁₀ H ₂₂ S	[13402-60-3]	2-decanethiol				
	$\Delta_v H$	(380–534)	54.6	395		[1999DYK/SVO]
C ₁₀ H ₂₂ S	[544-02-5]	diisopentyl sulfide				

TABLE 8. Phase change enthalpies of C₉ to C₁₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(339–366)	57.9	352	A	[1987STE/MAL, 1999DYK/SVO]
	$\Delta_v H$	(340–365)	56.9	352	C	[1962MAC/MAY2]
C ₁₀ H ₂₂ S	[872-10-6]	dipentyl sulfide				
	$\Delta_v H$	(346–365)	U66.3	356		[1999DYK/SVO]
	$\Delta_v H$	(346–366)	58.7	356	A	[1987STE/MAL]
	$\Delta_v H$	(346–366)	57.5	358	EB	[1962MAC/MAY2]
C ₁₀ H ₂₂ S ₂	[112-51-6]	dipentyl disulfide				
	$\Delta_v H$	(410–571)	59.8	425		[1999DYK/SVO]
	$\Delta_v H$		71.1 ± 0.2	298	C	[1985KUS]
C ₁₀ H ₂₂ S ₂	[1191-67-9]	1,10-decanedithiol				
	$\Delta_v H$	(434–571)	72.3	449	A	[1987STE/MAL, 1999DYK/SVO, 1943HAL/REI]
C ₁₀ H ₂₃ N	[2016-57-1]	decylamine				
	$\Delta_v H$	(410–506)	52.4	425	A,EST	[1987STE/MAL, 1956MAN2]
C ₁₀ H ₂₃ N	[7378-99-6]	N,N-methyloctylamine				
	$\Delta_v H$	(284–323)	54.0 ± 0.5	303		[1997VER]
	$\Delta_v H$	(371–517)	50.2	386	A	[1987STE/MAL]
C ₁₀ H ₂₃ N	[2050-92-2]	dipentylamine				
	$\Delta_v H$	(379–527)	51.2	394	A	[1987STE/MAL]
C ₁₀ H ₂₃ NO ₂	[126835-62-9]	3-(heptylamino)-1,2-propanediol				
	$\Delta_{\text{fus}}H$		28.8	324.9	DSC	[1993ACR]
C ₁₀ H ₂₃ N ₃	[67752-90-3]	[2-(dimethylamino)ethyl]methylhydrazone-2-propanone				
	$\Delta_v H$	(288–315)	62.3	301	A	[1987STE/MAL, 1980LEB/NAZ]
C ₁₀ H ₂₄ NO ₃ PS	[78-53-5]	O,O-diethyl-S-[2-(diethylamino)ethyl]thiophosphate				
	$\Delta_v H$	(358–407)	94.5	373	A	[1987STE/MAL, 1999DYK/SVO]
C ₁₀ H ₂₄ N ₂	[646-25-3]	decane-1,10-diamine				
	$\Delta_{\text{fus}}H$		57.81	332.9		[2002DAL/DEL]
C ₁₀ H ₂₄ N ₄	[996-70-3]	tetrakis(dimethylamino)ethylene				
	$\Delta_v H$	(358–485)	53.9 ± 0.5	298	EB	[1997STE/CHI4]
C ₁₀ H ₂₄ N ₄	[295-37-4]	1,4,8,11-tetraazacyclotetradecane				
	$\Delta_{\text{sub}}H$	(352–372)	133.9 ± 2.5	362	TE	[1983CLA/COR]
C ₁₀ H ₂₇ N ₅ O ₆	[114606-56-3]	8-[[[(1R)-1-(3,4-dimethoxyphenyl)-2-hydroxyethyl]amino]-3,7-dihydro-7-(2-methoxyethyl)-1,3-dimethyl-1H-pyridine-2,6-dione				
	$\Delta_{\text{fus}}H$ (I)		40.31	384.2		
	$\Delta_{\text{fus}}H$ (II)		39.88	401.2		
	$\Delta_{\text{fus}}H$ (III)		38.58	391.2		[1999GIR/PIE]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₁ F ₂₁ N ₃	[57731-09-6] $\Delta_v H$	2,2,2-trifluoro-N-[2,2,2-trifluoro-1-(trifluoromethyl)ethylidene]-N'-[2,2,2-trifluoro-1-(trifluoromethyl)-1-[[2,2,2-trifluoro-1-(trifluoromethyl)ethylidene]amino]ethyl] ethanimidamide	39.8			[1975PET/SHR3]
C ₁₁ F ₂₂	[75169-50-5] $\Delta_v H$	perfluoro(1-methyl-4- <i>tert</i> -butylcyclohexane) (mix <i>cis</i> + <i>trans</i>)	45.8	360		[1999DYK/SVO]
C ₁₁ F ₂₂	[na] $\Delta_v H$	perfluoro(1-methyl-4- <i>tert</i> -butylcyclohexane) (isomer not specified)	54.1 ± 0.5	298	EB	[1981VAR/BUL]
C ₁₁ F ₂₄ O ₂	[678-38-6] $\Delta_v H$	octadecafluoro-1,9- <i>bis</i> (trifluoromethoxy)nonane	43.0	323		[1999DYK/SVO]
C ₁₁ H ₄ Cl ₅ NO ₂	[77765-41-4] $\Delta_v H$	2,2,4-trichloro-5-[(3,4-dichlorophenyl)amino]-4-cyclopentene-1,3-dione	87.5	468	GC	[1980SHA/SAD]
C ₁₁ H ₅ BrCl ₃ NO ₂	[73373-59-8] $\Delta_v H$	2,2,4-trichloro-5-[(2-bromophenyl)amino]-4-cyclopentene-1,3-dione	67.5	468	GC	[1980SHA/SAD]
C ₁₁ H ₅ BrCl ₃ NO ₂	[73373-60-1] $\Delta_v H$	2,2,4-trichloro-5-[(3-bromophenyl)amino]-4-cyclopentene-1,3-dione	78.1	468	GC	[1980SHA/SAD]
C ₁₁ H ₅ BrCl ₃ NO ₂	[73373-61-2] $\Delta_v H$	2,2,4-trichloro-5-[(4-bromophenyl)amino]-4-cyclopentene-1,3-dione	82.9	468	GC	[1980SHA/SAD]
C ₁₁ H ₅ Cl ₄ NO ₂	[73373-63-4] $\Delta_v H$	2,2,4-trichloro-5-[(4-chlorophenyl)amino]-4-cyclopentene-1,3-dione	86.2	468	GC	[1980SHA/SAD]
C ₁₁ H ₆ N ₄	[6343-21-1] $\Delta_{\text{sub}} H$	bicyclo[2.2.1]hept-5-ene-2,2,3,3-tetracarbonitrile	117.2 ± 5.4	408	MG	[1972ROG2, 1977PED/RYL]
C ₁₁ H ₇ BrO ₂	[20717-79-7] $\Delta_{\text{sub}} H$	1-bromo-2-naphthoic acid	109.0 ± 2.7		ME	[2008GOL/SUU]
C ₁₁ H ₇ N ₂	[6023-46-7] $\Delta_{\text{fus}} H$ $\Delta_{\text{sub}} H$	2,2-dicyano-3-phenylpropionitrile	29.29 96.2 ± 0.4	411.2 353		[1994RAK/VER] T [1994RAK/VER]
C ₁₁ H ₇ N ₃	[6023-46-7] $\Delta_v H$	2,2-dicyano-1-phenylpropionitrile	66.9		B	[1994RAK/VER]
C ₁₁ H ₇ N ₃ O ₂ S	[186792-85-8] $\Delta_{\text{fus}} H$ (<i>red cryst</i>) $\Delta_{\text{fus}} H$ (<i>orange</i>)	2-[(2-nitrophenyl)amino]-3-thiophenecarbonitrile	30.4 30.4	407.8 407.8	DSC DSC	[2006LI/STO]
C ₁₁ H ₈ F ₃ N ₃ O ₇	[185852-05-5] $\Delta_{\text{fus}} H$	2,3-dihydro-6-nitro-3-[2-(nitrooxy)ethyl]-7-(trifluoromethyl)-4 <i>H</i> -1,3-benzoxazin-4-one	28.9	384.7	DSC	[1996FON/ROS]
C ₁₁ H ₈ N ₂	[244-63-3] $\Delta_{\text{fus}} H$	9 <i>H</i> -pyrido[3,4- <i>b</i>]indole (norharmaline)	25.5	471.5	DSC	[1996BUR/DAG]
C ₁₁ H ₈ N ₄	[13358-02-6] $\Delta_{\text{sub}} H$	3-methyl-1,1,2,2-tetracyanocyclohex-4-ene	82 ± 2.1	350	MG	[1971ROG, 1977PED/RYL]
C ₁₁ H ₈ O ₂	[86-55-5] $\Delta_{\text{fus}} H$ $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$	1-naphthoic acid	19.89 117.6 ± 0.4 110.4 ± 0.2 113.6	435.2		[1991ACR] DSC [1983HOL] ME [1974COL/ROU, 1977PED/RYL, 1987STE/MAL] C [1974SAB/GIL]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(457–573)	97.2	472	A	[1987STE/MAL]
C ₁₁ H ₈ O ₂	[93-09-4]	2-naphthoic acid				
	$\Delta_{\text{fus}} H$		23.54	460.2		[1991ACR]
	$\Delta_{\text{sub}} H$		119.5 ± 0.6		DSC	[1983HOL]
	$\Delta_{\text{sub}} H$	(347–363)	113.6 ± 0.8	365	ME	[1974COL/ROU, 1977PED/RYL, 1987STE/MAL]
	$\Delta_{\text{sub}} H$		117.2	298	C	[1974SAB/GIL]
	$\Delta_v H$	(463–582)	98.9	478	A	[1987STE/MAL]
C ₁₁ H ₉ Cl ₂ NO ₂	[101-27-9]	4-chlorobut-2-ynyl 3-chlorophenylcarbamate				
	$\Delta_{\text{fus}} H$		26.91	344.1	DSC	[1990DON/DRE]
C ₁₁ H ₉ Cl	[86-52-2]	1-(chloromethyl)naphthalene				
	$\Delta_v H$	(423–565)	59.8	494		[1999DYK/SVO]
	$\Delta_v H$	(407–447)	U 90.2	422	A	[1987STE/MAL]
C ₁₁ H ₉ N	[1009-89-5]	2-phenylpyridine				
	$\Delta_v H$		68.4 ± 1.9	298	CGC	[2009LIP/CHI, 2009LIP/HAN]
	$\Delta_v H$		68.7 ± 4.6	298	CGC	[2000RIB/MAT2]
	$\Delta_v H$		63.2		GC	[1996GOV/RUT]
C ₁₁ H ₉ N	[1008-88-4]	3-phenylpyridine				
	$\Delta_v H$		68.4 ± 1.6	298	CGC	[2009LIP/CHI, 2009LIP/HAN]
	$\Delta_v H$		64.5 ± 4.5	298	CGC	[2000RIB/MAT2]
C ₁₁ H ₉ N	[939-23-1]	4-phenylpyridine				
	$\Delta_{\text{fus}} H$		19.95	346.9	DSC	[2000RIB/MAT2]
	$\Delta_{\text{sub}} H$		81.4 ± 1.6	298		[2000RIB/MAT2]
	$\Delta_v H$		63.3		GC	[1996GOV/RUT]
C ₁₁ H ₉ NO ₂	[1631-28-3]	1-(4-methylphenyl)-1H-pyrrole-2,5-dione				
	$\Delta_{\text{sub}} H$	(350–370)	104.6 ± 0.8		C	[1998KIS/KAS]
C ₁₁ H ₉ NO ₃	[1081-17-0]	1-(4-methoxyphenyl)-1H-pyrrole-2,5-dione				
	$\Delta_{\text{sub}} H$	(350–370)	121.1 ± 0.8		C	[1998KIS/KAS]
C ₁₁ H ₁₀	[92-12-0]	1-methylnaphthalene				
	$\Delta_{\text{us}} H$		4.98	240.7		
	$\Delta_{\text{fus}} H$		6.95	242.7		[1996DOM/HEA]
	$\Delta_v H$	(343–423)	65.1 ± 1.1	298	GC	[2006HAF/PAR]
	$\Delta_v H$	(294–324)	60.1 ± 0.8	298	GS	[2003VER]
	$\Delta_v H$	(323–473)	62.4	298	GC	[2002LEI/CHA]
	$\Delta_v H$	(485–595)	50.0	500		[1992LEE/DEM]
	$\Delta_v H$	(259–388)	63.3	274		[1988SAS/JOS]
	$\Delta_v H$	(424–536)	49.6	455		[1981WIE/KOB]
	$\Delta_v H$	(424–536)	45.9	525		[1981WIE/KOB]
	$\Delta_v H$	(278–313)	57.5	293	A, GS	[1987STE/MAL, 1979MAC/PRA]
	$\Delta_v H$		57.3 ± 0.4	298	C	[1974SAB/CHA2]
	$\Delta_v H$	(415–526)	52.3	430	A, GS	[1987STE/MAL, 1955CAM/ROS]
C ₁₁ H ₁₀	[91-57-6]	2-methylnaphthalene				
	$\Delta_{\text{us}} H$		5.61	288.5		
	$\Delta_{\text{fus}} H$		12.13	307.7		[1996DOM/HEA]
	$\Delta_{\text{sub}} H$		65.7 ± 0.85		C	[1974SAB/CHA2]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{sub}}H$		61.7 ± 1.7			[1968KAR/RAB, 1977PED/RYL]
	Δ_vH	(424–535)	48.4	465		[1981WIE/KOB]
	Δ_vH	(424–535)	46.4	505		[1981WIE/KOB]
	Δ_vH	(423–515)	51.2	438	A, GS	[1987STE/MAL, 1955CAM/ROS]
C ₁₁ H ₁₀ BrNO ₂	[5460-29-7]	N-(3-bromopropyl)phthalimide				
	$\Delta_{\text{sub}}H$		116.0 ± 1.0	298	C	[2007RIB/SAN3]
C ₁₁ H ₁₀ N ₂ O	[72583-92-7]	2-(2-benzofuryl) Δ -2-imidazoline				
	$\Delta_{\text{fus}}H$ (I)		25.95	412.7		
	$\Delta_{\text{fus}}H$ (II)		28.53	420.3	DSC	[2001LEG/BAZ]
C ₁₁ H ₁₀ N ₂ O ₃	[13297-17-1]	2-methyl-3-acetylquinoxaline-1,4-dioxide				
	$\Delta_{\text{sub}}H$		117.0 ± 2.4	298	ME	[1997ACR/POW]
C ₁₁ H ₁₀ N ₂ O ₃	[40016-70-4]	2-methyl-3-carboxymethoxyquinoxaline-1,4-dioxide				
	$\Delta_{\text{sub}}H$		118.3 ± 2.6	298	C	[1997ACR/POW]
C ₁₁ H ₁₀ N ₂ O ₃	[61522-53-0]	3-(methoxycarbonyl)-2-methoxyquinoxaline-1-oxide				
	$\Delta_{\text{sub}}H$		118.1 ± 3.3	298	C	[2009GOM/MON]
C ₁₁ H ₁₀ O	[4780-79-4]	1-naphthalenemethanol				
	$\Delta_{\text{sub}}H$		102.3 ± 1.9	298	C	[2007MAT/MOR]
C ₁₁ H ₁₀ O	[1592-38-7]	2-naphthalenemethanol				
	$\Delta_{\text{sub}}H$		106.0 ± 2.1	298	C	[2007MAT/MOR]
C ₁₁ H ₁₀ O ₂	[2958-72-7]	pentacyclo[5.4.0 ^{2,6} 0 ^{3,10} 0 ^{5,9}]undecane-8,11-dione				
	$\Delta_{\text{us}}H$		0.32	309.8		
	$\Delta_{\text{us}}H$		9.61	345.3		
	$\Delta_{\text{fus}}H$		5.23	516.8	DSC	[1999JIM/ROU]
	$\Delta_{\text{us}}H$		32.14	365.9		
	$\Delta_{\text{fus}}H$		3.94	516.8	DSC	[1984WEI/LEF]
		Note: There is a large difference in the experimental enthalpies and transition temperature reported by the two research groups for the solid/solid transition around 355 K				
	$\Delta_{\text{sub}}H$		92.6 ± 1.0	298	ME	[1999JIM/ROU]
C ₁₁ H ₁₀ O ₂	[711-79-5]	2-acetyl-1-naphthol				
	$\Delta_{\text{fus}}H$		22.52	371.8		[1991ACR]
C ₁₁ H ₁₀ O ₂	[574-19-6]	1-acetyl-2-naphthol				
	$\Delta_{\text{fus}}H$		21.34	337		[1991ACR]
C ₁₁ H ₁₀ O ₄	[na]	<i>p</i> -methacryloyloxybenzoic acid				
	$\Delta_{\text{fus}}H$		34.0	455		[1996DOM/HEA]
C ₁₁ H ₁₁ Cl ₃ O ₃	[1928-40-1]	2,4,5-trichlorophenoxyacetic acid, propyl ester				
	Δ_vH	(444–573)	83.2	459	A	[1987STE/MAL, 1999DYK/SVO]
C ₁₁ H ₁₁ Cl ₃ O ₃	[na]	methyl 2-(2,4,5-trichlorophenoxy)butyrate				
	$\Delta_{\text{fus}}H$		28.87	316.5	DSC	[1969PLA/GLA]
C ₁₁ H ₁₁ N	[1198-37-4]	2,4-dimethylquinoline				
	Δ_vH	(458–543)	56.3	473	A	[1987STE/MAL]
C ₁₁ H ₁₁ N	[877-43-0]	2,6-dimethylquinoline				
	$\Delta_{\text{fus}}H$		20.4	330.8	AC,DSC	[2007CHI/JOH]
	$\Delta_{\text{sub}}H$		84.5 ± 1.5	298	C	[1995RIB/MAT3]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(337–591)	64.0 ± 0.1	340	IP,EB	[2007CHI/JOH]
	$\Delta_v H$	(337–591)	61.1 ± 0.1	380	IP,EB	[2007CHI/JOH]
	$\Delta_v H$	(337–591)	58.4 ± 0.1	420	IP,EB	[2007CHI/JOH]
	$\Delta_v H$	(337–591)	55.7 ± 0.1	460	IPEB	[2007CHI/JOH]
	$\Delta_v H$	(337–591)	53.0 ± 0.1	500	IP,EB	[2007CHI/JOH]
	$\Delta_v H$	(337–591)	50.0 ± 0.2	540	IP,EB	[2007CHI/JOH]
	$\Delta_v H$	(337–591)	46.8 ± 0.4	580	IP,EB	[2007CHI/JOH]
	$\Delta_v H$	(337–591)	67.1 ± 0.2	298	IP,EB	[2007CHI/JOH]
	$\Delta_v H$	(461–541)	55.7	476	A	[1987STE/MAL]
C ₁₁ H ₁₁ N	[93-37-8] $\Delta_{\text{sub}} H$	2,7-dimethylquinoline	87.5 ± 1.5	298	C	[1995RIB/MAT3]
C ₁₁ H ₁₁ NO ₂	[5323-50-2] $\Delta_{\text{sub}} H$	N-propylphthalimide	98.2 ± 1.4	298	C	[2006RIB/SAN]
C ₁₁ H ₁₁ N ₃ O ₂ S	[144-83-2] $\Delta_{\text{fus}} H$	4-amino-N-2-pyridinylbenzenesulfonamide (sulfapyridine)	40.47	462.7	DSC	[2003MAR/AVI, 2002MAR/GOM]
C ₁₁ H ₁₁ N ₃ O ₈	[53848-90-1] $\Delta_{\text{fus}} H$ $\Delta_{\text{fus}} H$	butyl 2,4,6-trinitrobenzoate	2.5 28.13	360 395.2	DSC	[1974WAR/WIL]
C ₁₁ H ₁₂ BrN ₅ O ₃	[244272-55-7] $\Delta_{\text{fus}} H$	2-bromo-6-methyl-3,9-dihydro-3-[(2-hydroxyethoxy)methyl]-9-oxo-5H-imidazol[1,2-a]pyrine	50.44	474.3	DSC	[1999ZIE/GOL]
C ₁₁ H ₁₂ Cl ₂ O ₃	[94-11-1] $\Delta_v H$	2,4-dichlorophenoxyacetic acid, isopropyl ester	(460–573) 69.5	475	A	[1987STE/MAL, 1999DYK/SVO]
C ₁₁ H ₁₂ Cl ₂ O ₃	[1928-61-1] $\Delta_v H$	2,4-dichlorophenoxyacetic acid, propyl ester	(444–573) 77.3	459	A	[1987STE/MAL, 1999DYK/SVO]
C ₁₁ H ₁₂ Cl ₂ O ₃	[na] $\Delta_{\text{fus}} H$	methyl 4-(2,4-dichlorophenoxy)butyrate	32.64	309.6	DSC	[1969PLA/GLA]
C ₁₁ H ₁₂ Cl ₂ O ₃	[18625-12-2] $\Delta_{\text{fus}} H$	4-(2,4-dichlorophenoxy)butanoic acid, methyl ester	22	309.7	DSC	[2005VEC/BRU]
C ₁₁ H ₁₂ Cl ₂ O ₄	[28191-20-0] $\Delta_v H$	2,4-dichlorophenoxyacetic acid, 3-hydroxypropyl ester	(463–483) 72.1	473	A	[1987STE/MAL, 1999DYK/SVO]
C ₁₁ H ₁₂ I ₃ NO ₂	[na] $\Delta_{\text{fus}} H$	(–) 3-[3-(amino-2,4,6-triiodophenyl)-2-ethylpropanoic acid (iopanoic acid)]	25.98	438.8	DSC	[1999LI/ZEL]
C ₁₁ H ₁₂ I ₃ NO ₂	[96-83-3] $\Delta_{\text{fus}} H$	(+) 3-[3-(amino-2,4,6-triiodophenyl)-2-ethylpropanoic acid (iopanoic acid)]	27.7	427	DSC	[1999LI/ZEL]
C ₁₁ H ₁₂ NO ₃ PS	[732-11-6] $\Delta_{\text{fus}} H$	O,O-dimethyl S-phthalimidomethyl phosphorodithioate	26.96	343.2	DSC	[1990DON/DRE]
C ₁₁ H ₁₂ N ₂ O	[60-80-0] $\Delta_{\text{fus}} H$	1,2-dihydro-1,5-dimethyl-2-phenyl-3H-pyrazol-3-one (antipyrine)	24.52	385.8		[1985OHM/LIP]
C ₁₁ H ₁₂ N ₂ OS ₂	[102-77-2] $\Delta_{\text{fus}} H$ (I) $\Delta_{\text{fus}} H$ (II)	2-(4-morpholiniothio)benzothiazole	12.97 17.99	345 357		[1978GUZ/LAR]
C ₁₁ H ₁₂ N ₂ O ₂	[73-22-3] $\Delta_{\text{sub}} H$	(l)-tryptophane	(340–440) 87.9 ± 8U	390	LE	[1977GAF/PIE]
C ₁₁ H ₁₂ N ₂ O ₃	[20771-72-6] $\Delta_{\text{sub}} H$	4-[(4-nitrophenyl)amino]pent-3-ene-2-one	121.9 ± 3.9	298	C	[1993RIB/RIB]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound					
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference	
C ₁₁ H ₁₂ N ₂ O ₅	[143248-64-0] $\Delta_{\text{fus}}H$	2,3-dihydro-6-methyl-3-[2-(nitrooxy)ethyl]-4H-1,3-benzoxazin-4-one	27.1	351.2	DSC	[1996FON/ROS]	
C ₁₁ H ₁₂ N ₄ O ₂ S	[127-79-7] $\Delta_{\text{fus}}H$	2-(4-aminobenzenesulfonamido)-4-methylpyrimidine (sulfamerazine)	41.3	508.5	DSC	[2003MAR/AVI, 2002MAR/GOM]	
	$\Delta_{\text{fus}}H$		31.6	515.2	DTA	[1971SUN/EIS]	
C ₁₁ H ₁₂ N ₄ O ₃ S	[80-35-3] $\Delta_{\text{fus}}H$	4-amino-N-(6-methoxy-3-pyridazinyl)benzenesulfonamide (sulphamethoxypyridazine)	22.3	453.4		[1995BUS/ESC]	
C ₁₁ H ₁₂ O	[na] Δ_vH	2-ethylidene-3-phenylpropanal (333–374)	73.6	348	A	[1987STE/MAL]	
C ₁₁ H ₁₂ O ₂	[2495-37-6] Δ_vH	benzyl methacrylate (347–431)	70.5	362	A	[1987STE/MAL]	
C ₁₁ H ₁₂ O ₂	[103-36-6] Δ_vH	ethyl cinnamate (453–544)	57.8	468	A	[1987STE/MAL]	
C ₁₁ H ₁₂ O ₂	[5331-64-6] Δ_vH	1-phenyl-1,3-pentanedione (371–550)	64.6	386	A	[1987STE/MAL]	
C ₁₁ H ₁₂ O ₂	[39522-76-4] $\Delta_{\text{fus}}H$	1-phenyl-4,7-dioxaspiro[2.4]heptane	22.6	303.1	DSC	[1998VER/PEN]	
	$\Delta_{\text{sub}}H$		91.8 ± 0.8	298		[1998VER/PEN]	
	Δ_vH		(307–333)	71.3 ± 0.7	298	GS	[2002VER]
	Δ_vH		(288–302)	69.6 ± 0.7	298	GS	[1998VER/PEN]
C ₁₁ H ₁₂ O ₂	[40317-63-3] Δ_vH	4-carboxymethylpentacyclo[4.3.0.0.2.5 ⁰ 4.7]nonane (303–343)	80.0 ± 1.7	333		[1984BEC/RUC]	
C ₁₁ H ₁₂ O ₂	[33892-75-0] $\Delta_{\text{fus}}H$	3,4-dihydro-5-methoxy-1(2H)-naphthalenone	22.2	362.5	DSC	[2009MAT/SOU2]	
	$\Delta_{\text{sub}}H$		97.9 ± 0.4	298	C	[2009MAT/SOU2]	
C ₁₁ H ₁₂ O ₂	[1078-19-9] $\Delta_{\text{fus}}H$	3,4-dihydro-6-methoxy-1(2H)-naphthalenone	22.8	351.3	DSC	[2009MAT/SOU2]	
	$\Delta_{\text{sub}}H$		104.7 ± 0.9	298	C	[2009MAT/SOU2]	
C ₁₁ H ₁₂ O ₂	[6836-19-7] $\Delta_{\text{fus}}H$	3,4-dihydro-7-methoxy-1(2H)-naphthalenone	23.2	334.8	DSC	[2009MAT/SOU2]	
	$\Delta_{\text{sub}}H$		103.1 ± 0.9	298	C	[2009MAT/SOU2]	
C ₁₁ H ₁₂ O ₃	[94-02-0] Δ_vH	benzoylactic acid, ethyl ester (380–538)	72.1	395	A	[1987STE/MAL]	
C ₁₁ H ₁₂ O ₃	[607-91-0] Δ_vH	myristicin (368–553)	61.2	383	A	[1987STE/MAL]	
C ₁₁ H ₁₂ O ₃	[na] Δ_vH	2-piperonylpropanal (373–423)	74.5	388	A	[1987STE/MAL]	
C ₁₁ H ₁₂ O ₄	[2309-07-1] $\Delta_{\text{fus}}H$	methyl 4'-hydroxy-3'-methoxycinnamate (methyl ferulate)	25.84	335.7	DSC	[2010PAN/SAR]	
C ₁₁ H ₁₂ O ₄	[7345-82-6] $\Delta_{\text{sub}}H$	<i>trans</i> -2,3-dimethoxycinnamic acid (380–392)	141.0 ± 0.9	298	ME	[1999MON/HIL]	

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₁ H ₁₂ O ₄	[16909-09-4] $\Delta_{\text{sub}}H$	<i>trans</i> -2,4-dimethoxycinnamic acid (391–404)	149.2 ± 1.3	298	ME	[1999MON/HIL]
C ₁₁ H ₁₂ O ₄	[10538-51-9] $\Delta_{\text{sub}}H$	<i>trans</i> -2,5-dimethoxycinnamic acid (376–391)	138.8 ± 1.1	298	ME	[1999MON/HIL]
C ₁₁ H ₁₂ O ₄	[2316-26-9] $\Delta_{\text{sub}}H$	<i>trans</i> -3,4-dimethoxycinnamic acid (390–404)	149.9 ± 0.8	298	ME	[1999MON/HIL]
C ₁₁ H ₁₂ O ₄	[16909-11-8] $\Delta_{\text{sub}}H$	<i>trans</i> -3,5-dimethoxycinnamic acid (385–397)	141.4 ± 0.5	298	ME	[1999MON/HIL]
C ₁₁ H ₁₃ ClF ₃ N ₃ O ₄ S ₃	[346-18-9] $\Delta_{\text{fus}}H$	6-chloro-3,4-dihydro-2-methyl-3-[[2,2,2-trifluoroethyl]-thio]methyl-2 <i>H</i> -1,2,4-benzothiadiazine-5-sulfonamide-1,1-dioxide (polythiazide)	42.67	493.2		[2000HAN/PAR]
C ₁₁ H ₁₃ ClO ₃	[94-81-5] $\Delta_{\text{fus}}H$	4-(4-chloro-2-methylphenoxy)butanoic acid	32.02	373.5	DSC	[1990DON/DRE]
C ₁₁ H ₁₃ Cl ₃	[61468-36-8] Δ_vH	4- <i>tert</i> -butyl-2,3,6-trichlorotoluene (423–570)	62.7	438	A	[1987STE/MAL, 1973FEL/SAV, 1999DYK/SVO]
C ₁₁ H ₁₃ F ₃ N ₂ O ₃ S	[53780-34-0] $\Delta_{\text{fus}}H$	5'-(trifluoromethanesulphonamide)acet-2',4-xylylide	37.66	457.3	DSC	[1990DON/DRE]
C ₁₁ H ₁₃ F ₃ N ₄ O ₄	[2091-05-2] $\Delta_{\text{fus}}H$	N(3),N(3)-diethyl-2,4-dinitro-6-(trifluoromethyl)-1,3-benzenediamine	29.13	372.1	DSC	[1990DON/DRE]
C ₁₁ H ₁₃ NO	[14091-93-1] $\Delta_{\text{sub}}H$	(<i>E</i>)-3-(methylamino)-1-phenyl-but-2-en-1-one	99.2 ± 4.2	298	C	[1993RIB/RIB]
C ₁₁ H ₁₃ NO	[7294-89-5] $\Delta_{\text{sub}}H$	4-phenylaminopent-3-ene-2-one	89.9 ± 3.8	298	C	[1993RIB/RIB]
C ₁₁ H ₁₃ NO ₄	[22781-23-3] $\Delta_{\text{fus}}H$	2,3-isopropylidenedioxyphenyl-N-methylcarbamate	29.45	402.6	DSC	[1990DON/DRE]
C ₁₁ H ₁₃ N ₃ O ₃ S	[127-69-5] $\Delta_{\text{fus}}H$	3,4-dimethylisoxazol 5-sulphanylamide	8.41	448.2	DSC	[1996CIO/MEL]
C ₁₁ H ₁₃ N ₅ O ₃	[114199-19-8] $\Delta_{\text{fus}}H$	6-methyl-3,9-dihydro-3-[(2-hydroxyethoxy)methyl]-9-oxo-5 <i>H</i> -imidazol[1,2- <i>a</i>]pyrine	36.06	465.4	DSC	[1999ZIE/GOL]
C ₁₁ H ₁₄	[4912-92-9] $\Delta_{\text{fus}}H$	1,1-dimethylindane	11.99	227.4		[1996DOM/HEA]
	Δ_vH	(313–348)	50.1	328	A,IP,EB	[1987STE/MAL, 1978OSB/SCO]
	Δ_vH	(313–467)	50.5	328	A,IP,EB	[1987STE/MAL]
	Δ_vH	(387–467)	45.9	402	A,IP,EB	[1987STE/MAL, 1978OSB/SCO]
	Δ_vH	(313–467)	51.9 ± 0.3	298	IP,EB	[1978OSB/SCO]
C ₁₁ H ₁₄	[1685-82-1] $\Delta_{\text{fus}}H$	4,6-dimethylindane	12.88	256.5		[1996DOM/HEA]
	Δ_vH	(313–467)	56.9	328	A,IP,EB	[1987STE/MAL, 1978OSB/SCO]
	Δ_vH	(313–363)	56.4	328	A,IP,EB	[1987STE/MAL, 1978OSB/SCO]
	Δ_vH	(415–467)	50.3	430	A,IP,EB	[1987STE/MAL, 1978OSB/SCO]
	Δ_vH	(313–467)	57.9 ± 0.4	298	IP,EB	[1978OSB/SCO]
C ₁₁ H ₁₄	[6682-71-9] $\Delta_{\text{fus}}H$	4,7-dimethylindane	13.52	272.7		[1996DOM/HEA]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(313–470)	54.7	328	A,IP,EB	[1987STE/MAL, 1978OSB/SCO]
	$\Delta_v H$	(313–363)	56.9	328	A,IP,EB	[1987STE/MAL, 1978OSB/SCO]
	$\Delta_v H$	(417–470)	50.6	432	A,IP,EB	[1987STE/MAL, 1978OSB/SCO]
	$\Delta_v H$	(313–470)	58.3 ± 0.4	298	IP, EB	[1978OSB/SCO]
C ₁₁ H ₁₄	[2055-40-5]	4-isopropylstyrene				
	$\Delta_v H$	(408–478)	48.5	423	A	[1987STE/MAL]
C ₁₁ H ₁₄	[17498-71-4]	α -isopropylstyrene				
	$\Delta_v H$	(278–318)	53.3 ± 0.3	298	GS	[1999VER/EBE]
C ₁₁ H ₁₄	[2809-64-5]	5-methyl-1,2,3,4-tetrahydronaphthalene				
	$\Delta_v H$	(416–508)	53.4	431	A	[1987STE/MAL]
C ₁₁ H ₁₄	[1680-51-9]	6-methyl-1,2,3,4-tetrahydronaphthalene				
	$\Delta_v H$	(411–502)	53.7	426	A	[1987STE/MAL]
C ₁₁ H ₁₄	[3937-24-4]	2,4,5-trimethylstyrene				
	$\Delta_v H$	(352–490)	56.4	367	A	[1987STE/MAL, 1949BUC/COL]
C ₁₁ H ₁₄	[769-25-5]	2,4,6-trimethylstyrene				
	$\Delta_v H$	(362–483)	50.9	377	A	[1987STE/MAL, 1949BUC/COL]
C ₁₁ H ₁₄	[4421-32-3]	pentacyclo[5.4.0 ^{2,6} 0 ^{3,10} 0 ^{5,9}]undecane				
	$\Delta_{\text{fus}} H$	(5–320)	4.86	164.4		
	$\Delta_{\text{fus}} H$	(5–320)	6.38	475.8	AC	[1995KAB/KOZ]
	$\Delta_{\text{sub}} H$		54.7 ± 0.9	337	C	[1995KAB/KOZ]
	$\Delta_{\text{sub}} H$	(273–323)	54.9 ± 1.1	298	ME	[1995KAB/KOZ]
C ₁₁ H ₁₄ ClNO	[1918-16-7]	2-chloro-N-isopropyl N-phenylacetamide				
	$\Delta_{\text{fus}} H$		26.05	351.4	DSC	[1990DON/DRE]
C ₁₁ H ₁₄ Cl ₂	[61468-35-7]	4- <i>tert</i> -butyl-2,5-dichlorotoluene				
	$\Delta_v H$	(395–538)	57.0	410	A	[1987STE/MAL, 1973FEL/SAV, 1999DYK/SVO]
C ₁₁ H ₁₄ N ₂ O ₂	[718-36-5]	4-nitrobenzylidene <i>tert</i> -butylamine				
	$\Delta_{\text{sub}} H$		91.1 ± 3.1	298	C	[1989ACR/KIR]
C ₁₁ H ₁₄ N ₂ O ₂	[128478-71-7]	2-cyano-2-nitroadamantane				
	$\Delta_{\text{fus}} H$		4.98	470.2		[1990FRI/DOG]
		Note: Enthalpy seems low, compound may have lower temperature phase transitions.				
	$\Delta_{\text{sub}} H$	(307–368)	70.0 ± 1.9	338	T	[1990FRI/DOG]
C ₁₁ H ₁₄ N ₂ O ₃	[3585-88-4]	4-nitrobenzylidene <i>tert</i> -butylamine N-oxide				
	$\Delta_{\text{sub}} H$		116.5 ± 3.1	298	C	[1989ACR/KIR]
C ₁₁ H ₁₄ N ₂ O ₄	[204189-06-0]	3-nitro-3-(4-nitrophenyl)pentane				
	$\Delta_{\text{fus}} H$		20.29	na	DSC	[1997VER3]
	$\Delta_{\text{sub}} H$		110.8 ± 0.8	298	Fus+Vap	[1997VER3]
	$\Delta_v H$	(321–358)	88.0 ± 0.8	340	GS	[1997VER3]
	$\Delta_v H$	(321–358)	90.5 ± 0.8	298	GS	[1997VER3]
C ₁₁ H ₁₄ O	[938-16-9]	<i>tert</i> -butyl phenyl ketone				
	$\Delta_v H$	(330–493)	55.5	345	A	[1987STE/MAL, 1947STU]
C ₁₁ H ₁₄ O	[na]	2-ethyl-3-phenylpropanal				
	$\Delta_v H$	(343–388)	64.6	358	A	[1987STE/MAL]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₁ H ₁₄ O	[582-62-7] $\Delta_v H$	isobutyl phenyl ketone (331–501)	55.7	346	A	[1987STE/MAL, 1947STU]
C ₁₁ H ₁₄ O	[na] $\Delta_v H$	2,3,5-trimethylacetophenone (352–557)	57.9	367	A	[1987STE/MAL, 1947STU]
C ₁₁ H ₁₄ O ₂	[18523-34-7] $\Delta_v H$	1,1-dimethoxy-2-phenylcyclopropane (278–313)	63.9 ± 0.6		GS	[1998VER/PEN]
C ₁₁ H ₁₄ O ₂	[122-72-5] $\Delta_v H$ $\Delta_v H$	3-acetoxy-1-phenylpropane (293–333) (392–516)	74.3 56.8	306 402	A	[1987STE/MAL] [1986CIH/VOJ]
C ₁₁ H ₁₄ O ₂	[210009-92-7] $\Delta_{\text{fus}} H$	2-acetyl-3,5-dimethylanisole	0.99	323.2	DTA	[1989SAL/ABA]
		Note: Reported enthalpy of fusion is too small, and the published enthalpy and entropy of fusion data are internally inconsistent				
C ₁₁ H ₁₄ O ₂	[120-50-3] $\Delta_v H$ $\Delta_v H$ $\Delta_v H$	butyl benzoate (374–474) (374–474) (343–405)	63.2 55.7 59.1	394 452 358	BG BG A	[1988KAT2] [1988KAT2] [1987STE/MAL]
C ₁₁ H ₁₄ O ₂	[93-16-3] $\Delta_v H$	1,2-dimethoxy-4-(1-propenyl)benzene (358–521)	61.9	373	A	[1987STE/MAL, 1947STU]
C ₁₁ H ₁₄ O ₂	[136-60-7] $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$	isobutyl benzoate (370–467) (370–467) (291–300) (338–510)	60.4 54.4 58.1 57.1	393 449 295 353	BG BG A A	[1988KAT2] [1988KAT2] [1987STE/MAL] [1987STE/MAL, 1947STU]
C ₁₁ H ₁₄ O ₂	[2510-99-8] $\Delta_v H$ $\Delta_v H$	ethyl 2-phenylpropionate (293–329) (293–329)	63.2 ± 0.3 64.0 ± 0.3	311 298	GS GS	[1999VER8] [1999VER8]
C ₁₁ H ₁₄ O ₂	[53917-01-4] $\Delta_v H$	1-(4-methoxyphenyl)-2-butanone (373–443)	62.6	388	A	[1987STE/MAL]
C ₁₁ H ₁₄ O ₂	[2270-20-4] $\Delta_{\text{fus}} H$ $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$	5-phenylvaleric acid (315–327) (315–327)	23.4 118.5 ± 0.8 119.4 ± 1.1	332 321 298	DSC ME ME	[2001MON/HIL] [2001MON/HIL] [2001MON/HIL]
C ₁₁ H ₁₄ O ₂	[1077-58-3] $\Delta_{\text{sub}} H$	2- <i>tert</i> -butylbenzoic acid (306–322)	99.8 ± 0.4	315	ME	[1979COL/JIM]
C ₁₁ H ₁₄ O ₂	[7498-54-6] $\Delta_{\text{sub}} H$	3- <i>tert</i> -butylbenzoic acid (318–335)	103. ± 0.5	327	ME	[1979COL/JIM]
C ₁₁ H ₁₄ O ₂	[98-73-7] $\Delta_{\text{fus}} H$ $\Delta_{\text{sub}} H$	4- <i>tert</i> -butylbenzoic acid (325–343)	17.91 103.8 ± 0.4	440 334	DSC ME	[1993ACR] [1979COL/JIM]
C ₁₁ H ₁₄ O ₂	[20651-71-2] $\Delta_{\text{sub}} H$	4-butylbenzoic acid (333–349)	110.5 ± 0.7	298	ME	[2004MON/ALM]
C ₁₁ H ₁₄ O ₂	[2529-39-7] $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$	2,3,4,5-tetramethylbenzoic acid (337–360) (337–360)	113.4 ± 0.6 115.9 ± 0.6	348 298	ME ME	[1988COL/JIM] [1988COL/JIM]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₁ H ₁₄ O ₂	[2408-38-0]	2,3,4,6-tetramethylbenzoic acid				
	$\Delta_{\text{sub}}H$	(330–351)	106.9 ± 0.5	341	ME	[1988COL/JIM]
	$\Delta_{\text{sub}}H$	(330–351)	109.7 ± 0.5	298	ME	[1988COL/JIM]
C ₁₁ H ₁₄ O ₂	[2604-45-7]	2,3,5,6-tetramethylbenzoic acid				
	$\Delta_{\text{sub}}H$	(330–351)	104.6 ± 0.8	341	ME	[1988COL/JIM]
	$\Delta_{\text{sub}}H$	(330–351)	106.1 ± 0.8	298	ME	[1988COL/JIM]
C ₁₁ H ₁₄ O ₂	[3854-90-8]	3,5-diethylbenzoic acid				
	$\Delta_{\text{sub}}H$	(325–343)	104.1 ± 4.2	334	A	[1974ROU/TUR, 1977PED/RYL, 1987STE/MAL]
C ₁₁ H ₁₄ O ₂ S	[111895-49-9]	<i>p</i> -tolyl but-1-enyl sulfone				
	$\Delta_{\text{sub}}H$		106.3 ± 2.5		B	[1969MAC/MCN, 1969MAC/MCN2, 1970COX/PIL]
C ₁₁ H ₁₄ O ₂ S	[24931-66-6]	<i>p</i> -tolyl but-2-enyl sulfone				
	$\Delta_{\text{sub}}H$		107.5 ± 2.5		B	[1969MAC/STE, 1970COX/PIL]
C ₁₁ H ₁₄ O ₂ S	[17482-19-8]	<i>p</i> -tolyl but-3-enyl sulfone				
	$\Delta_{\text{sub}}H$		113.4 ± 2.9		B	[1969MAC/STE, 1970COX/PIL]
C ₁₁ H ₁₄ O ₂ S	[16192-03-3]	<i>p</i> -tolyl-isobutenyl sulfone				
	$\Delta_{\text{sub}}H$		102.1 ± 2.5		B	[1969MAC/MCN, 1969MAC/MCN2, 1977PED/RYL]
C ₁₁ H ₁₄ O ₂ S	[16192-04-4]	<i>p</i> -tolyl 2-methylprop-2-enyl sulfone				
	$\Delta_{\text{sub}}H$		106.7 ± 2.9			[1969MAC/STE, 1970COX/PIL]
C ₁₁ H ₁₄ O ₃	[94-26-8]	butyl 4-hydroxybenzoate				
	$\Delta_{\text{fus}}H$		26.6	341.8		[1999GIO/BET]
	$\Delta_{\text{sub}}H$	(320–333)	108.4 ± 0.8	298	GS	[2005PER/ROD]
	Δ_vH		76.9		TGA	[2002CHA/DOL]
	Δ_vH		72.2		TGA	[2001CHA/DOL]
C ₁₁ H ₁₄ O ₃	[na]	<i>(dl)</i> 3-phenyl-3-hydroxy-2,2-dimethylpropanoic acid				
	$\Delta_{\text{fus}}H$		37.24	407		[1991CHI/BRA]
C ₁₁ H ₁₄ O ₃	[na]	<i>(d)</i> 3-phenyl-3-hydroxy-2,2-dimethylpropanoic acid				
	$\Delta_{\text{fus}}H$		39.75	431		[1991CHI/BRA]
C ₁₁ H ₁₄ O ₃	[4521-28-2]	(4-methoxyphenyl)-4-butyric acid				
	$\Delta_{\text{fus}}H$		25.3	330.9		[1979ARM/JAM]
C ₁₁ H ₁₄ O ₃	[na]	<i>(dl)</i> 3-hydroxy-3-phenylvaleric acid				
	$\Delta_{\text{fus}}H$		35.15	394		[1991CHI/BRA]
C ₁₁ H ₁₄ O ₃	[na]	<i>(d)</i> 3-hydroxy-3-phenylvaleric acid				
	$\Delta_{\text{fus}}H$		30.96	379		[1991CHI/BRA]
C ₁₁ H ₁₄ O ₃	[na]	2-piperonylpropanol				
	Δ_vH	(373–443)	84.8	388	A	[1987STE/MAL]
C ₁₁ H ₁₄ O ₃	[1498-96-0]	4- <i>n</i> -butoxybenzoic acid				
	$\Delta_{\text{fus}}H$ (<i>solid-to-liq</i> <i>cryst</i>)		18.83	420.7		
	$\Delta_{\text{fus}}H$ (<i>liq</i> <i>cryst-to-liq</i>)		2.93	432.2		[1967HER]
	$\Delta_{\text{sub}}H$		129.0 ± 0.8	298		[2010RIB/FER3]
C ₁₁ H ₁₄ O ₃	[6627-89-0]	<i>tert</i> -butyl phenyl carbonate				
	Δ_vH	(294–348)	67.6 ± 0.6	298	GS	[2008VER/EME2]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₁ H ₁₄ O ₄	[2107-70-2]	3-(3,4-dimethoxyphenyl)propionic acid				
	$\Delta_{\text{fus}}H$		32.38	370.9	DSC	[2001MON/HIL4]
	$\Delta_{\text{sub}}H$	(352–366)	140.3 ± 0.8	359	ME	[2001MON/HIL4]
	$\Delta_{\text{sub}}H$	(352–366)	143.6 ± 2.2	298	ME	[2001MON/HIL4]
C ₁₁ H ₁₅ BrO	[99857-52-0]	4-methyl-2-bromophenyl isobutyl ether				
	Δ_vH	(293–328)	71.0 ± 0.3	298	GS	[2005STR/SPO]
C ₁₁ H ₁₅ BrO ₃	[929259-36-9]	1-bromo-2-[2-(2-methoxyethoxy)ethoxy]benzene				
	Δ_vH	(310–373)	83.1 ± 0.3	298	GS	[2006DAB/SPO]
C ₁₁ H ₁₅ Cl	[42597-10-4]	4- <i>tert</i> -butyl-2-chlorotoluene				
	Δ_vH	(372–503)	54.0	387	A	[1987STE/MAL, 1973FEL/SAV, 1999DYK/SVO]
C ₁₁ H ₁₅ N	[42525-65-2]	2-phenylethylazetidide				
	Δ_vH	(302–333)	62.2	317	A	[1987STE/MAL, 1976KIP/TSV]
C ₁₁ H ₁₅ N	[4096-20-2]	N-phenylpiperidine				
	Δ_vH	(284–323)	64.0 ± 0.4	303	GS	[1998VER6]
	Δ_vH	(284–323)	64.3 ± 0.4	298	GS	[1998VER6]
C ₁₁ H ₁₅ N	[23074-42-2]	1-adamantyl-1-carbonitrile				
	$\Delta_{\text{us}}H$		5.06	279.4	DSC	[2008SIN/MUR2]
	$\Delta_{\text{us}}H$		5.5	280		
	$\Delta_{\text{fus}}H$		15.0	458		[1984FOU/AMO]
	$\Delta_{\text{sub}}H$	(294–312)	67.1 ± 0.8	303	ME	[1992ABB/JIM]
	$\Delta_{\text{sub}}H$	(294–312)	67.2 ± 0.8	298	ME	[1992ABB/JIM]
C ₁₁ H ₁₅ NO	[3376-24-7]	benzylidene <i>tert</i> -butylamine N-oxide				
	$\Delta_{\text{sub}}H$		86.8 ± 0.9	298	C	[1989ACR/KIR]
C ₁₁ H ₁₅ NO	[1696-17-9]	N,N-diethylbenzamide				
	Δ_vH	(373–403)	56.5	388	A	[1987STE/MAL]
	Δ_vH	(374–405)	53.2	389		[1969DAV/MAK]
C ₁₁ H ₁₅ NO	[na]	(4R,5R)-3,4-dimethyl-5-phenyl-1,3-oxazolidine				
	Δ_vH	(293–303)	50.0 ± 1.3	298		[1998GUD/TOR]
C ₁₁ H ₁₅ NO	[na]	(4S,5R)-3,4-dimethyl-5-phenyl-1,3-oxazolidine				
	Δ_vH	(293–303)	52.4 ± 0.9	298		[1998GUD/TOR]
C ₁₁ H ₁₅ NO	[15351-09-1]	2-(dimethylamino)-1-phenyl-1-propanone				
	Δ_vH	(293–333)	64.8 ± 1.2	298	GS	[1994WEL/VER]
C ₁₁ H ₁₅ NO ₂	[na]	4- <i>trans</i> -cyanocyclohexyl (E) 2-butenate				
	$\Delta_{\text{fus}}H$		24.4	366.2		[1995KEL/SCH]
C ₁₁ H ₁₅ NO ₂	[94-25-7]	butyl 4-aminobenzoate				
	$\Delta_{\text{fus}}H$		23.9	330.6	DSC	[2005SCH]
	$\Delta_{\text{fus}}H$		20.46	331.1		[1991ACR]
C ₁₁ H ₁₅ NO ₂	[2631-40-5]	2-(1-methylethyl)phenyl methylcarbamate				
	$\Delta_{\text{fus}}H$		26.14	369.3	DSC	[1990DON/DRE]
C ₁₁ H ₁₅ NO ₂	[94-14-4]	4-aminobenzoic acid, 2-methylpropyl ester				
	$\Delta_{\text{fus}}H$		10.7	327.8	DSC	[2005SCH]
C ₁₁ H ₁₅ NO ₂ S	[2032-65-7]	4-methylthio-3,5-xylyl methylcarbamate				
	$\Delta_{\text{fus}}H$		30.36	393.8	DSC	[1991ACR, 1990DON/DRE]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₁ H ₁₅ NO ₂ S ₂	[949171-66-8] $\Delta_{\text{fus}}H$	N-theonylthiocarbamic-O-pentyl ester				
			24.59	354.3	DSC	[2007RIB/MON]
	$\Delta_{\text{sub}}H$		165.6 ± 2.1	298	C	[2007RIB/MON]
C ₁₁ H ₁₅ NO ₃	[75587-96-1] $\Delta_{\text{fus}}H$	1,2-dihydro-6-neopentyl-2-oxonicotinic acid				
			19.33	469.2	DSC	[1986SHA/BRI]
C ₁₁ H ₁₅ NS	[1696-17-9] $\Delta_{\text{sub}}H$	N,N-diethylbenzamide				
			91.4 ± 3.2	298	C	[1989RIB/SOU]
C ₁₁ H ₁₅ N ₃ O ₂	[140670-55-9] $\Delta_{\text{fus}}H$	N-caproyl-pyrazinamide				
			35.95	351.7		[1991LIU/GUO]
C ₁₁ H ₁₆	[1777-44-2] Δ_vH	tetracyclo[6.2.1.0 ^{2,7} .0 ^{3,5}]undecane				
			55.3 ± 0.3	298	C	[1996VAR/PAS]
C ₁₁ H ₁₆	[538-68-1] Δ_vH	pentylbenzene				
		(284–323)	55.1 ± 0.4	298	GS	[2006VER]
	Δ_vH		55.3	298		[1994RUZ/ZAB]
	Δ_vH		55.1	298		[1971WIL/ZWO]
C ₁₁ H ₁₆	[2049-94-7] Δ_vH	isopentylbenzene				
		(302–466)	53.0	298	EB	[1947STU, 2006VER]
C ₁₁ H ₁₆	[2049-95-8] Δ_vH	<i>tert</i> -pentylbenzene				
		(294–318)	52.3 ± 0.3	298	GS	[2009VER/EME3]
C ₁₁ H ₁₆	[2719-52-0] Δ_vH	<i>dl</i> 2-phenylpentane				
		(302–466)	50.3	317	A	[1987STE/MAL]
C ₁₁ H ₁₆	[1075-38-3] Δ_vH	1- <i>tert</i> -butyl-3-methylbenzene				
		(274–318)	51.1 ± 0.3	298	GS	[2008VER/KOZ2]
	Δ_vH	(279–314)	51.4 ± 0.6	296	GS	[1998VER]
	Δ_vH		51.3 ± 0.6	298		[1998VER]
C ₁₁ H ₁₆	[98-51-1] Δ_vH	1- <i>tert</i> -butyl-4-methylbenzene				
		(279–323)	52.2 ± 0.1	298	GS	[2008VER/KOZ2]
	Δ_vH	(279–314)	52.3 ± 0.5	296	GS	[1998VER]
	Δ_vH		52.2 ± 0.6	298		[1998VER]
C ₁₁ H ₁₆	[98-15-1] Δ_vH	4- <i>tert</i> -butyltoluene				
		(342–465)	49.1	357	A	[1987STE/MAL, 1973FEL/SAV]
C ₁₁ H ₁₆	[2050-24-0] Δ_vH	3,5-diethyltoluene				
		(307–474)	49.6	322	A	[1987STE/MAL, 1947STU]
C ₁₁ H ₁₆	[4920-99-4] Δ_vH	1-ethyl-3-isopropylbenzene				
		(301–466)	48.8	316	A	[1987STE/MAL, 1947STU]
C ₁₁ H ₁₆	[4218-48-8] Δ_vH	1-ethyl-4-isopropylbenzene				
		(304–469)	49.4	319	A	[1987STE/MAL, 1947STU]
C ₁₁ H ₁₆	[3982-67-0] Δ_vH	2-ethyl-1,3,5-trimethylbenzene				
		(312–481)	52.6	327	A	[1987STE/MAL]
C ₁₁ H ₁₆	[18262-85-6] Δ_vH	3-ethyl-1,2,4-trimethylbenzene				
		(347–488)	61.3	362	A	[1987STE/MAL]
C ₁₁ H ₁₆	[17851-27-3] Δ_vH	5-ethyl-1,2,4-trimethylbenzene				
		(317–481)	56.4	332	A	[1987STE/MAL, 1947STU]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₁ H ₁₆	[700-12-9]	pentamethylbenzene				
	$\Delta_{\text{us}}H$		1.98	296.8		
	$\Delta_{\text{fus}}H$		10.67	328.2		[1996DOM/HEA]
	$\Delta_{\text{sub}}H$		71.6 ± 0.1	298	C	[1994SAB/TAB]
	$\Delta_{\text{sub}}H$	(296–313)	77.4 ± 0.4	298	ME	[1989COL/JIM]
	Δ_vH	(338–503)	57.8	353	A	[1987STE/MAL]
C ₁₁ H ₁₆ N ₂ O ₂	[82413-41-0]	1,3-dimethyl-5,6-pentamethyleneuracil				
	$\Delta_{\text{sub}}H$	(335–358)	111.9 ± 0.2	346	ME	[1983COL/JIM]
	$\Delta_{\text{sub}}H$	(323–338)	108.8 ± 5	330	QR	[1980TEP/YAN, 1983COL/JIM]
	$\Delta_{\text{sub}}H$	(340–370)	113.4 ± 1.3	355	MS	[1980TEP/YAN, 1983COL/JIM]
C ₁₁ H ₁₆ N ₂ O ₂	[156461-80-2]	N-methyl-N-(4- <i>tert</i> -butylphenyl)nitramine				
$\Delta_{\text{fus}}H$			23.4	351.2		[2002DAS/ZAL]
C ₁₁ H ₁₆ N ₂ O ₂ S ₂	[19475-21-9]	N-ethyl-S-methyl-N'-tosylisothiourea				
$\Delta_{\text{fus}}H$			35.8	390.2	DSC	[1992REI/HAN]
C ₁₁ H ₁₆ N ₂ O ₂ S ₂	[21017-78-7]	N-methyl-S-ethyl-N'-tosylisothiourea				
$\Delta_{\text{fus}}H$			26.5	414.2	DSC	[1992REI/HAN]
C ₁₁ H ₁₆ N ₄ O ₂	[35873-40-6]	8-butyltheophylline				
$\Delta_{\text{fus}}H$			32.3	509.2	DSC	[1989GON/KRA]
C ₁₁ H ₁₆ N ₄ O ₂	[15030-44-1]	8- <i>tert</i> -butyltheophylline				
$\Delta_{\text{fus}}H$			48.2	402.3	DSC	[1989GON/KRA]
C ₁₁ H ₁₆ N ₄ O ₄	[24613-06-7]	(–) 4,4'-(1-methyl-1,2-ethanediyl)bis-2,6-piperazinedione (dexrazoxane)				
$\Delta_{\text{fus}}H$			37.82	467.6	DSC	[1999LI/ZEL]
C ₁₁ H ₁₆ N ₄ O ₄	[21416-67-1]	(+) 4,4'-(1-methyl-1,2-ethanediyl)bis-2,6-piperazinedione (dexrazoxane)				
$\Delta_{\text{fus}}H$			44.98	507.4	DSC	[1999LI/ZEL]
C ₁₁ H ₁₆ O	[51528-17-7]	2- <i>sec</i> -butyl-4-methylphenol				
	Δ_vH	(413–548)	58.4	428	A	[1987STE/MAL]
	Δ_vH	(383–523)	59.0	373		[1953STA/MUL]
	Δ_vH	(383–523)	58.0	398		[1953STA/MUL]
	Δ_vH	(383–523)	55.8	423		[1953STA/MUL]
	Δ_vH	(383–523)	51.4	473		[1953STA/MUL]
C ₁₁ H ₁₆ O	[2409-55-4]	2- <i>tert</i> -butyl-4-methylphenol				
	$\Delta_{\text{sub}}H$	(288–318)	82.6 ± 0.5	303	GS	[1999VER2]
	$\Delta_{\text{sub}}H$	(288–318)	82.9 ± 0.5	298	GS	[1999VER2]
	$\Delta_{\text{sub}}H$	(274–294)	77.4	284	A	[1987STE/MAL, 1960AIH]
	Δ_vH	(327–358)	63.0 ± 0.3	343	GS	[1999VER2]
	Δ_vH	(327–358)	65.7 ± 0.3	298	GS	[1999VER2]
	Δ_vH	(385–517)	58.9	400	A	[1987STE/MAL]
	Δ_vH	(343–507)	57.7	348		[1953STA/MUL]
	Δ_vH	(343–507)	55.7	373		[1953STA/MUL]
	Δ_vH	(343–507)	52.6	423		[1953STA/MUL]
	Δ_vH	(343–507)	48.5	473		[1953STA/MUL]
C ₁₁ H ₁₆ O	[88-60-8]	2- <i>tert</i> -butyl-5-methylphenol				
	$\Delta_{\text{sub}}H$	(277–294)	80.4 ± 1.3	287	GS	[1999VER2]
	$\Delta_{\text{sub}}H$	(277–294)	79.7 ± 1.3	298	GS	[1999VER2]
	Δ_vH	(296–343)	65.9 ± 0.3	320	GS	[1999VER2]
	Δ_vH	(296–343)	67.2 ± 0.3	298	GS	[1999VER2]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(378–490)	59.8	393	A	[1987STE/MAL]
	$\Delta_v H$	(383–518)	53.0	398	A	[1987STE/MAL]
C ₁₁ H ₁₆ O	[2219-82-1]	2- <i>tert</i> -butyl-6-methylphenol				
	$\Delta_{\text{fus}} H$		17.32	302.5	DSC	[1999VER]
	$\Delta_v H$	(308–343)	62.2 ± 0.5	326	GS	[1999VER]
	$\Delta_v H$	(308–343)	63.8 ± 0.5	298	GS	[1999VER]
	$\Delta_v H$	(375–505)	55.2	390	A	[1987STE/MAL]
C ₁₁ H ₁₆ O	[98-27-1]	4- <i>tert</i> -butyl-2-methylphenol				
	$\Delta_v H$	(291–333)	71.3 ± 0.6	312	GS	[1999VER2]
	$\Delta_v H$	(291–333)	72.1 ± 0.6	298	GS	[1999VER2]
	$\Delta_v H$	(347–520)	61.5	362	A	[1987STE/MAL]
	$\Delta_v H$	(275–297)	75.7	286	A	[1987STE/MAL, 1960AIH]
	$\Delta_v H$	(347–532)	55.7	348		[1953STA/MUL]
	$\Delta_v H$	(347–532)	53.9	373		[1953STA/MUL]
	$\Delta_v H$	(347–532)	53.2	398		[1953STA/MUL]
	$\Delta_v H$	(347–532)	50.9	423		[1953STA/MUL]
	$\Delta_v H$	(347–532)	46.7	473		[1953STA/MUL]
C ₁₁ H ₁₆ O	[3968-87-4]	2-ethyl-3-phenyl-1-propanol				
	$\Delta_v H$	(348–393)	70.9	363	A	[1987STE/MAL]
C ₁₁ H ₁₆ O	[87-26-3]	2-(2-pentyl)phenol				
	$\Delta_v H$	(397–501)	74.4	413	EB	[1990NES/NAZ]
	$\Delta_v H$	(397–501)	59.6	412		[1993KAS/MOK]
C ₁₁ H ₁₆ O	[14938-35-3]	4-pentylphenol				
	$\Delta_v H$	(423–563)	60.9	438	A	[1987STE/MAL]
C ₁₁ H ₁₆ O	[80-46-6]	4- <i>tert</i> -pentylphenol				
	$\Delta_{\text{sub}} H$	(293–333)	87.4 ± 0.5	313	GS	[1999VER2]
	$\Delta_{\text{sub}} H$	(293–333)	88.3 ± 0.5	298	GS	[1999VER2]
	$\Delta_v H$	(297–333)	64.2 ± 0.2	329	GS	[1999VER2]
	$\Delta_v H$	(297–333)	65.3 ± 0.2	298	GS	[1999VER2]
	$\Delta_v H$	(385–548)	58.2	400	A	[1987STE/MAL]
C ₁₁ H ₁₆ O	[10521-91-2]	5-phenyl-1-pentanol				
	$\Delta_v H$	(373–430)	58.2	388	A	[1987STE/MAL]
C ₁₁ H ₁₆ O	[91967-71-4]	(1-propoxyethyl)benzene				
	$\Delta_v H$	(288–321)	56.4 ± 0.2	305	GS	[2001VER/HEI]
	$\Delta_v H$	(288–321)	56.7 ± 0.2	298	GS	[2001VER/HEI]
C ₁₁ H ₁₆ O	[65757-61-1]	(1-isopropoxyethyl)benzene				
	$\Delta_v H$	(278–313)	55.4 ± 0.3	298	GS	[2002KRA/VAS, 2002VER/HEI]
C ₁₁ H ₁₆ O	[1712-74-9]	ethyl cumyl ether				
	$\Delta_v H$	(278–313)	54.8 ± 0.5	296	GS	[2001VER/HEI2]
	$\Delta_v H$	(278–313)	54.7 ± 0.5	298	GS	[2001VER/HEI2]
C ₁₁ H ₁₆ O	[31108-34-6]	1-(2,4,6-trimethylphenyl)ethanol				
	$\Delta_{\text{sub}} H$	(282–313)	U 5.7	297	A	[1987STE/MAL]
C ₁₁ H ₁₆ O ₂	[121-00-6]	2- <i>tert</i> -butyl-4-methoxyphenol				
	$\Delta_v H$	(403–463)	54.4	418	A	[1987STE/MAL]
C ₁₁ H ₁₆ O ₂	[533-24-4]	1,3-dihydroxy-4-pentylbenzene				
	$\Delta_v H$	(423–488)	84.9	438	A,GC	[1987STE/MAL, 1975KUN/LIL]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₁ H ₁₆ O ₂	[774-48-1] $\Delta_v H$	phenyldiethoxymethane (283–329)	62.8 ± 0.6	298	GS	[2002VER]
C ₁₁ H ₁₆ O ₂	[25310-92-3] $\Delta_v H$	1,1-dimethoxy-1-phenylpropane (288–328)	58.9 ± 0.3	298	GS	[2002VER]
	$\Delta_v H$	(288–328)	57.9 ± 0.3		GS	[1998VER/PEN]
C ₁₁ H ₁₆ O ₂	[na] $\Delta_v H$	<i>tert</i> -pentylcatechol (isomer not specified) (398–473)	58.2	436		[1965GAK/BAB]
C ₁₁ H ₁₆ O ₂	[828-51-3] $\Delta_{\text{fus}} H$	1-adamantanecarboxylic acid	2.25	524.2	DSC	[1986HAR/GIL]
C ₁₁ H ₁₆ O ₃	[7149-82-8] $\Delta_{\text{fus}} H$	(racemic) 3-(2-ethylphenoxy)-propane-1,2-diol	34.8	324.1	DSC	[2008BRE/BRE]
C ₁₁ H ₁₆ O ₃	[1092799-92-2] $\Delta_{\text{fus}} H$	(S)-3-(2-ethylphenoxy)-propane-1,2-diol	35.0	342.1	DSC	[2008BRE/BRE]
C ₁₁ H ₁₆ O ₅	[na] $\Delta_v H$	ethylcamphoric acid anhydride (391–571)	70.8	406	A	[1987STE/MAL, 1947STU]
C ₁₁ H ₁₆ O ₅	[na] $\Delta_v H$	(1-methylallyl)[1-(allyloxycarbonyl)ethyl]carbonate (368–508)	60.2	383	A	[1987STE/MAL]
C ₁₁ H ₁₇ Cl ₃ OS	[76619-96-0] $\Delta_v H$	2,3,3-trichloro-2-propenethioic acid, O-octyl ester (443–483)	74.2		GC	[1980PIT/KIS]
C ₁₁ H ₁₇ NO	[552-79-4] $\Delta_{\text{fus}} H$	(–) 2-dimethylamino-1-phenyl-1-propanol (methylephedrine) (79–399)	21.8	358.6	AC	[2008DI/WAN]
	$\Delta_{\text{fus}} H$		30.56	361.2	DSC	[1999LI/ZEL]
C ₁₁ H ₁₇ NO	[1201-56-5] $\Delta_{\text{fus}} H$	(+) 2-dimethylamino-1-phenyl-1-propanol (methylephedrine)	26.6	336	DSC	[1999LI/ZEL]
C ₁₁ H ₁₇ NO	[5511-18-2] $\Delta_{\text{sub}} H$	1-adamantyl carboxamide (336–354)	105.9 ± 0.5	345	ME	[1989ABB/JIM]
	$\Delta_{\text{sub}} H$		108.0 ± 0.5	298	ME	[1989ABB/JIM]
C ₁₁ H ₁₇ N ₅	[153495-36-4] $\Delta_{\text{fus}} H$	6,9-dimethyl-8-butyladenine	36.0	409.2		[1994ZIE/ZIE]
	$\Delta_{\text{sub}} H$	(348–354)	106.0 ± 0.1	351	ME	[1994ZIE/ZIE]
C ₁₁ H ₁₈	[768-91-2] $\Delta_{\text{us}} H$	1-methyladamantane	1.91	169.5		
	$\Delta_{\text{us}} H$		1.47	211.5		
	$\Delta_{\text{fus}} H$		3.71	392		[1977CLA/KNO]
	$\Delta_{\text{sub}} H$	(300–342)	67.8 ± 1.3	298	BG	[1977STE/WAT]
C ₁₁ H ₁₈	[700-56-1] $\Delta_{\text{sub}} H$	2-methyladamantane (310–330)	67.5 ± 2.1	320		[1975CLA/KNO]
	$\Delta_{\text{sub}} H$	(300–340)	68.2 ± 1.3	298		[1977STE/WAT]
C ₁₁ H ₁₈ N ₂	[71172-36-6] $\Delta_{\text{fus}} H$	undecanedinitrile	26.0	266.1	DSC	[2007BAD/BLA]
C ₁₁ H ₁₈ O	[26533-38-0] $\Delta_v H$	6-methyl-3-isopropenyl-5-hepten-2-one (390–420)	49.7	405		[1989WAN/YIN]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₁ H ₁₈ O	[702-98-7]	2-methyl-2-adamantanol				
	$\Delta_{\text{sub}}H$	(298–334)	91.3 ± 0.8	298	ME	[2003CHA/BLO2]
	$\Delta_{\text{sub}}H$		91.4 ± 0.3	298	C	[2003CHA/BLO2]
C ₁₁ H ₁₈ O ₂	[7492-41-3]	borneol formate				
	Δ_vH	(320–487)	52.7	335	A	[1987STE/MAL, 1947STU]
C ₁₁ H ₁₈ O ₂	[2142-94-1]	3,7-dimethyl- <i>cis</i> -2,6-octadienyl formate				
	Δ_vH	(330–498)	58.1	345	A	[1987STE/MAL]
C ₁₁ H ₁₈ O ₂	[105-86-2]	3,7-dimethyl- <i>trans</i> -2,6-octadienyl formate				
	Δ_vH	(334–503)	57.1	349	A	[1987STE/MAL, 1947STU]
C ₁₁ H ₁₈ O ₂	[1200-67-5]	isoborneol formate				
	Δ_vH	(383–441)	53.5	398	A	[1987STE/MAL]
C ₁₁ H ₁₈ O ₄	[4167-77-5]	1,1-cyclopentanedicarboxylic acid diethyl ester				
	Δ_vH	(293–323)	66.8 ± 0.4		GS	[1998VER/KUM]
C ₁₁ H ₁₈ O ₅	[na]	4-oxononanedioic acid, dimethyl ester				
	Δ_vH	(394–559)	72.7	409	A	[1987STE/MAL]
C ₁₁ H ₁₈ O ₆	[na]	1,1,1- <i>tris</i> (ethoxycarbonyl)methane				
	Δ_vH	(298–338)	74.1 ± 0.4		GS	[1995RAK/VER]
C ₁₁ H ₁₈ O ₆	[na]	1,1,1- <i>tris</i> (methoxycarbonyl)pentane				
	Δ_vH	(298–338)	81.0 ± 0.4		GS	[1995RAK/VER]
C ₁₁ H ₁₉ NO ₂	[62391-95-1]	ethyl <i>bis</i> (isopropyl)cyanoacetate				
	Δ_vH	(284–319)	65.0 ± 0.9	298	GS	[1995VER/BEC]
C ₁₁ H ₁₉ NO ₃	[114-26-1]	2-isopropoxyphenyl N-methylcarbamate				
	$\Delta_{\text{fus}}H$		22.96	362.7	DSC	[1991ACR, 1990DON/DRE]
C ₁₁ H ₁₉ NS	[27149-31-1]	2,4-di- <i>tert</i> -butylthiazole				
	$\Delta_{\text{fus}}H$		10.5	258.2		[1966MEY/MET]
C ₁₁ H ₁₉ N ₃ O	[23947-60-6]	5-butyl-2-ethylamino-6-methylpyrimidin-4-ol				
	$\Delta_{\text{fus}}H$		20.32	432.5	DSC	[1991ACR, 1990DON/DRE]
C ₁₁ H ₁₉ N ₅ S	[4147-51-7]	6-ethylthio-N,N'- <i>bis</i> (1-methylethyl)-1,3,5-triazine-2,4-diamine				
	$\Delta_{\text{fus}}H$		23.94	377.7	DSC	[1991ACR, 1990DON/DRE]
C ₁₁ H ₁₉ O ₅	[1446-19-1]	N-acetyl-(<i>l</i>)-glutamic acid, diethyl ester				
	Δ_vH	(403–503)	67.2	418	A	[1987STE/MAL]
C ₁₁ H ₂₀	[180-43-8]	spiro[5.5]undecane				
	Δ_vH		56.1	298	C	[1975SUB/ZWO]
C ₁₁ H ₂₀	[1606-08-2]	cyclopentylcyclohexane				
	Δ_vH	(383–488)	47.9	398	A	[1987STE/MAL]
C ₁₁ H ₂₀	[180-43-8]	bicyclo[3.3.3]undecane				
	$\Delta_{\text{sub}}H$		63.6 ± 0.8	298	C	[1975PAR/STE, 1977PED/RYL]
C ₁₁ H ₂₀ Cl ₄	[3922-34-7]	1,1,1,11-tetrachloroundecane				
	Δ_vH	(303–353)	92.5	318	A	[1987STE/MAL, 1999DYK/SVO]
C ₁₁ H ₂₀ Cl ₄	[210049-49-3]	1,2,10,11-tetrachloroundecane				
	Δ_vH		78.7			[1998DRO/TOM]
C ₁₁ H ₂₀ N ₆	[13452-85-2]	1-pyrrolidinyl-3,5- <i>bis</i> (dimethylamino)-s-triazine				
	$\Delta_{\text{fus}}H$		25.61	403.1	DSC	[1991ACR]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₁ H ₂₀ N ₆ O	[16269-02-6] $\Delta_{\text{fus}}H$	1-morpholinyl-3,5- <i>bis</i> (dimethylamino)- <i>s</i> -triazine				
			24.69	397.4	DSC	[1991ACR]
C ₁₁ H ₂₀ N ₆ S	[41492-69-7] $\Delta_{\text{fus}}H$	1-(thiomorpholinyl)-3,5- <i>bis</i> (dimethylamino)- <i>s</i> -triazine				
			29.08	391.2	DSC	[1991ACR]
C ₁₁ H ₂₀ O	[878-13-7] Δ_vH Δ_vH Δ_vH	cycloundecanone				
		(363–433)	60.3	378	A	[1987STE/MAL]
		(448–501)	51.8	463	A, EB	[1987STE/MAL, 1976MEY/HOT]
			64.3 ± 0.6	298		[1972WOL]
C ₁₁ H ₂₀ O ₂	[1118-71-4] Δ_vH	2,2,6,6-tetramethyl-3,5-heptanedione				
			59.5	298		[1978RIB/IRV]
C ₁₁ H ₂₀ O ₂	[103-11-7] Δ_vH	<i>(dl)</i> 2-ethylhexyl acrylate				
		(323–489)	55.3	338	A	[1987STE/MAL, 1947STU]
C ₁₁ H ₂₀ O ₂	[na] Δ_vH	formic acid, 3- <i>para</i> -menthol ester				
		(320–492)	52.0	335	A	[1987STE/MAL, 1947STU]
C ₁₁ H ₂₀ O ₂	[1551-43-5] Δ_vH Δ_vH Δ_vH Δ_vH	cyclohexyl valerate				
			63.7 ± 0.1	298	C	[2004PAU/ZAI, 2003ZAI/VER]
		(273–318)	67.2 ± 0.8	298	ME	[2003ZAI/VER]
		(273–318)	63.9 ± 0.4	298	ME	[2003ZAI/VER]
		(293–332)	62.4 ± 0.7	298	GS	[2003ZAI/VER]
C ₁₁ H ₂₀ O ₂	[na] Δ_vH	1-methylcyclohexyl isobutyrate				
		(333–378)	57.2	298	CGC	[1999VER/HEI]
C ₁₁ H ₂₀ O ₂	[na] Δ_vH	3-methylcyclohexyl isobutyrate				
		(333–378)	59.3	298	CGC	[1999VER/HEI]
C ₁₁ H ₂₀ O ₂	[na] Δ_vH	4-methylcyclohexyl isobutyrate				
		(333–378)	59.7	298	CGC	[1999VER/HEI]
C ₁₁ H ₂₀ O ₂	[29878-49-7] Δ_vH	cyclohexyl pivalate				
		(333–378)	59.0	298	CGC	[1999VER/HEI]
C ₁₁ H ₂₀ O ₂	[61732-96-5] Δ_vH	2-hexyl-4,7-dihydro-1,3-dioxepin				
		(333–453)	66.0	348	A	[1987STE/MAL]
C ₁₁ H ₂₀ O ₂	[2499-59-4] Δ_vH	octyl acrylate				
		(331–500)	56.2	346	A	[1987STE/MAL, 1947STU]
C ₁₁ H ₂₀ O ₂	[1725-03-7] $\Delta_{\text{fus}}H$ $\Delta_{\text{fus}}H$ Δ_vH Δ_vH Δ_vH	oxa-2-cyclododecanone (undecanolactone)				
			3.36	250.2		
			12.61	275.3		[1996DOM/HEA]
		(365–387)	57.7 ± 0.8	376	MM	[1991WIB/WAL]
		(365–387)	66.2 ± 1.3	298	MM	[1991WIB/WAL]
		(353–413)	70.5	368	A	[1987STE/MAL]
C ₁₁ H ₂₀ O ₂	[112-38-9] Δ_vH	10-undecenoic acid				
		(387–548)	70.6	402	A	[1987STE/MAL, 1947STU]
C ₁₁ H ₂₀ O ₂	[707-29-9] Δ_vH	3,3-dimethyl-1,5-dioxaspiro[5.5]undecane				
		(283–323)	59.0 ± 0.6		GS	[1998VER/PEN, 2002VER]
C ₁₁ H ₂₀ O ₃	[24431-34-3] Δ_vH Δ_vH	hexyl levulinate				
		(363–540)	66.6	378	A	[1987STE/MAL, 1947STU]
			59.1	479		[1931SCH/COW]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₁ H ₂₀ O ₄	[77008-66-3] $\Delta_v H$	(<i>dl</i>) hexyl 2-acetoxypropionate (322–517)	70.3	337	A	[1987STE/MAL]
C ₁₁ H ₂₀ O ₄	[1732-10-1] $\Delta_v H$	azelaic acid, dimethyl ester (413–540)	63.6	428	A	[1987STE/MAL]
C ₁₁ H ₂₀ O ₄	[77-25-8] $\Delta_v H$	diethyl diethylmalonate (386–491)	68.5	401	A	[1987STE/MAL]
C ₁₁ H ₂₀ O ₄	[1852-04-6] $\Delta_{\text{fus}} H$ $\Delta_{\text{fus}} H$ $\Delta_{\text{fus}} H$ $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$ $\Delta_v H$	undecanedioic acid (295–313)	1.6 41.2 39.65 141.5	355.3 380.1 385	DSC TPTD	[2005ROU/TEM] [1996DOM/HEA] [2005CHA/ZIE]
Note: Values based on TPTD method are not consistent with values determined by other experimental methods						
		(371–381)	158.6 ± 1.9	376	ME	[1999RIB/MON]
		(371–381)	162.5 ± 1.9	298	ME	[1999RIB/MON]
		(424–503)	128.2 ± 2.3	298	CGC	[2005ROU/TEM]
C ₁₁ H ₂₀ O ₄	[1732-10-1] $\Delta_v H$	dimethyl azelate (298–373)	82.3 ± 0.4	298	GS	[2006VER/KOZ]
C ₁₁ H ₂₀ O ₅	[na] $\Delta_v H$	hexyl[1-(methoxycarbonyl)ethyl]carbonate (371–538)	65.9	386	A	[1987STE/MAL]
C ₁₁ H ₂₀ O ₅	[na] $\Delta_v H$	propyl[1-(butoxycarbonyl)ethyl]carbonate (330–463)	66.4	345	A	[1987STE/MAL]
C ₁₁ H ₂₁ N	[80606-32-2] $\Delta_v H$	2-butyl-2-methylhexanenitrile (298–388)	59.8 ± 0.4		GS	[1994RAK/VER]
C ₁₁ H ₂₁ N	[2244-07-1] $\Delta_v H$ $\Delta_v H$ $\Delta_v H$	undecanonitrile (290–340) (355–534)	71.8 ± 0.3 63.7	298 370	GS A	[2005EME/VER] [1987STE/MAL]
			71.1 ± 0.1	298	C	[1977STRI/SUN]
C ₁₁ H ₂₁ N	[3319-01-5] $\Delta_v H$ $\Delta_v H$	N-cyclohexylpiperidine (288–328)	59.9 ± 0.6	308	GS	[1998VER6]
		(288–328)	60.5 ± 0.6	298	GS	[1998VER6]
C ₁₁ H ₂₁ NO	[15770-38-4] $\Delta_v H$	N-hexanoylpiperidone (383–433)	66.3	398	A	[1987STE/MAL]
C ₁₁ H ₂₁ N ₅ S	[4147-51-7] $\Delta_{\text{fus}} H$	6-(ethylthio)-N,N'-bis(1-methylethyl)-1,3,5-triazine-2,4-diamine	23.94	377.7	DSC	[1990DON/DRE]
C ₁₁ H ₂₁ N ₇	[125867-94-9] $\Delta_{\text{fus}} H$	1-(piperiziny)-3,5-bis(dimethylamino)-s-triazine	23.01	382	DSC	[1991ACR]
C ₁₁ H ₂₂	[4292-92-6] $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$	pentylcyclohexane	52.9 ± 0.5 54.1 ± 0.3 53.9 55.0	298 298 298 298	GCC	[1987AZA] [1978FUC/PEA] [1975KUS/SAI] [1971WIL/ZWO]
C ₁₁ H ₂₂	[4457-00-5] $\Delta_v H$	hexylcyclopentane	55.9	298		[1971WIL/ZWO]
C ₁₁ H ₂₂	[821-95-4]	1-undecene				

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{fus}}H$		9.2	217.3		
	$\Delta_{\text{fus}}H$		16.99	224		[1996DOM/HEA]
	Δ_vH	(283–312)	54.3 ± 0.3	298	GS	[2000VER/WAN]
	Δ_vH		55.4	298		[1971WIL/ZWO]
	Δ_vH	(378–473)	48.2	393	A	[1987STE/MAL, 1950FOR/CAM]
C ₁₁ H ₂₂	[821-96-5]	<i>cis</i> 2-undecene				
	Δ_vH	(333–393)	53.2	348	A	[1987STE/MAL]
C ₁₁ H ₂₂	[693-61-8]	<i>trans</i> 2-undecene				
	Δ_vH	(333–393)	53.0	348	A	[1987STE/MAL]
C ₁₁ H ₂₂	[821-97-6]	<i>cis</i> 3-undecene				
	Δ_vH	(333–393)	52.3	348	A	[1987STE/MAL]
C ₁₁ H ₂₂	[1002-68-2]	<i>trans</i> 3-undecene				
	Δ_vH	(333–393)	52.0	348	A	[1987STE/MAL]
C ₁₁ H ₂₂	[821-98-7]	<i>cis</i> 4-undecene				
	Δ_vH	(333–393)	51.6	348	A	[1987STE/MAL]
C ₁₁ H ₂₂	[693-62-9]	<i>trans</i> 4-undecene				
	Δ_vH	(333–393)	52.1	348	A	[1987STE/MAL]
C ₁₁ H ₂₂	[764-96-5]	<i>cis</i> 5-undecene				
	Δ_vH	(333–393)	51.4	348	A	[1987STE/MAL]
C ₁₁ H ₂₂	[764-97-6]	<i>trans</i> 5-undecene				
	Δ_vH	(333–393)	51.8	348	A	[1987STE/MAL]
C ₁₁ H ₂₂	[na]	3-methyl-3-propyl-1-heptene				
	Δ_vH	(263–293)	52.8 ± 1.0	278	HSA	[1995CHI/HES]
	Δ_vH	(263–293)	50.9	298	HSA	[1995CHI/HES]
	Δ_vH		51.5	298	CGC	[1995CHI/HES]
C ₁₁ H ₂₂ Cl ₂	[822-01-5]	1,1-dichloroundecane				
	Δ_vH	(430–500)	59.5	445		[1999DYK/SVO, 1987VAR/LOS2]
	Δ_vH	(430–500)	71.7	298		[1987VAR/LOS2, 1991BAS/SVO]
C ₁₁ H ₂₂ N ₂	[880-09-1]	bis(piperidino)methane				
	Δ_vH	(283–322)	61.9 ± 0.9	303	GS	[2002VER2]
	Δ_vH	(283–322)	62.2 ± 0.9	298	GS	[2002VER2]
C ₁₁ H ₂₂ N ₂ O ₂	[73154-82-2]	undecandiamide				
	$\Delta_{\text{fus}}H$		64.4	451.2	DSC	[2006BAD/DEL]
C ₁₁ H ₂₂ O	[878-13-7]	cycloundecanone				
	$\Delta_{\text{fus}}H$		23.0	287.7		[1998GON/SZW]
C ₁₁ H ₂₂ O	[36633-49-5]	1-hexylcyclopentanol				
	Δ_vH	(387–509)	59.2	402	A	[1987STE/MAL]
C ₁₁ H ₂₂ O	[na]	cyclohexyl <i>tert</i> -amyl ether				
	Δ_vH		54.3 ± 0.2	298		[2002VER]
C ₁₁ H ₂₂ O	[112-12-9]	2-undecanone				
	$\Delta_{\text{fus}}H$		28.78	290.5		[1993VIL/HAM]
	Δ_vH	(461–538)	51.5	476	A	[1987STE/MAL]
	Δ_vH		69.7 ± 0.5	298	GCC	[1979SAL/PEA]
	Δ_vH		67.0 ± 0.4	298	C	[1979SUN/SVE2]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(393–523)	56.2	408	A	[1987STE/MAL, 1975AMB/ELL]
	$\Delta_v H$		46.4	506		[1975AMB/ELL]
	$\Delta_v H$	(335–433)	61.6	350	A, EB	[1987STE/MAL, 1966MEY/WAG]
	$\Delta_v H$	(341–497)	61.9	356		[1947STU]
C₁₁H₂₂O	[927-49-1]	6-undecanone				
	$\Delta_v H$	(343–383)	59	298	CGC	[1995CHI/HOS]
	$\Delta_v H$	(343–383)	61.8	298	CGC	[1995CHI/HOS]
	$\Delta_v H$	(388–543)	55.3	403	A	[1987STE/MAL]
	$\Delta_v H$	(461–513)	50.4	476	A	[1987STE/MAL]
	$\Delta_v H$		63.5 ± 0.5	298	GCC	[1979SAL/PEA]
	$\Delta_v H$	(383–514)	45.8	500		[1975STR/SUN]
C₁₁H₂₂O	[4436-99-1]	2,2,6,6-tetramethyl-4-heptanone				
	$\Delta_v H$		52.9 ± 0.2	298	C	[1971SEL]
C₁₁H₂₂O	[112-44-7]	undecanal				
	$\Delta_v H$	(293–329)	64.6 ± 0.5	298	GS	[2003VER/KRA2]
	$\Delta_v H$	(323–343)	69.3	298	CGC	[1996KOU/HOS, 2000OVA/KOU]
	$\Delta_v H$	(288–400)	60.2	303	A	[1987STE/MAL]
C₁₁H₂₂O₂	[5870-93-9]	heptyl butyrate				
	$\Delta_v H$	(384–498)	58.7	399	A	[1987STE/MAL]
C₁₁H₂₂O₂	[6454-22-4]	4,5-dimethyl-2-hexyl-1,3-dioxolane				
	$\Delta_v H$	(333–453)	65.6	348	A	[1987STE/MAL]
C₁₁H₂₂O₂	[2244-84-0]	4-heptyl-1,3-dioxane				
	$\Delta_v H$	(353–453)	64.4	368	A	[1987STE/MAL]
C₁₁H₂₂O₂	[41277-7502]	3-hexyl-4-hydroxytetrahydro-2H-pyran				
	$\Delta_v H$	(383–453)	73.6	398	A	[1987STE/MAL]
C₁₁H₂₂O₂	[5458-59-3]	isopropyl caprylate				
	$\Delta_v H$	(338–420)	57.5	353	A	[1987STE/MAL]
	$\Delta_v H$	(338–419)	58.3	353		[1948BON/ATH, 1984BOU/FRI]
C₁₁H₂₂O₂	[110-42-9]	methyl decanoate (methyl caprate)				
	$\Delta_v H$		62.0	350		[2002VAN/VAN]
	$\Delta_v H$		62.9 ± 0.1	337		[2002VAN/VAN]
	$\Delta_v H$		66.1 ± 0.2	298		[2002VAN/VAN]
	$\Delta_v H$	(373–433)	66.9	298	GC	[1997KRO/VEL]
	$\Delta_v H$	(453–543)	49.9	498	GC	[1993HUS/SAR]
	$\Delta_v H$		66.3 ± 0.5	298	GCC	[1980FUC/PEA]
	$\Delta_v H$		66.8 ± 0.6	298	C	[1977MAN/SEL]
	$\Delta_v H$	(379–500)	57.1	394	A, E	[1987STE/MAL, 1963ROS/SCH]
	$\Delta_v H$	(324–370)	63	339	MG,OM	[1952SCO/MAC]
C₁₁H₂₂O₂	[5432-30-4]	2-octyl-1,3-dioxolane				
	$\Delta_v H$	(333–453)	60.3	348	A	[1987STE/MAL]
C₁₁H₂₂O₂	[624-13-5]	propyl caprylate				
	$\Delta_v H$	(343–500)	58.8	358	A	[1987STE/MAL]
	$\Delta_v H$	(343–426)	58.2	358		[1948BON/ATH, 1984BOU/FRI]
C₁₁H₂₂O₂	[143-13-5]	nonyl acetate				
	$\Delta_v H$	(277–309)	66.2 ± 0.2	298	GS	[2006KRA/VER]
	$\Delta_v H$		66.8	298		[1997DEF/CAR]
	$\Delta_v H$	(313–358)	67	298	GC	[1997KOU/HOS, 2000OVA/KOU]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound					
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference	
C ₁₁ H ₂₂ O ₂	[245658-29-1] $\Delta_v H$	2,2-dimethylpropanoic acid, 1,1-dimethylbutyl ester (333–378)	52.3	298	CGC	[1999VER/HEI]	
C ₁₁ H ₂₂ O ₂	[245658-35-9] $\Delta_v H$	2,2-dimethylpropanoic acid, 1,1,2-trimethylpropyl ester (333–378)	52.8	298	CGC	[1999VER/HEI]	
C ₁₁ H ₂₂ O ₂	[245658-24-6] $\Delta_v H$	3,3-dimethylbutanoic acid, 1,1-dimethylpropyl ester (333–378)	53.2	298	CGC	[1999VER/HEI]	
C ₁₁ H ₂₂ O ₂	[245658-38-2] $\Delta_v H$	2-methylpropanoic acid, 1,1,3-trimethylbutyl ester (333–378)	53.4	298	CGC	[1999VER/HEI]	
C ₁₁ H ₂₂ O ₂	[na] $\Delta_v H$	2,6-dimethyl-2-heptanol acetate (333–378)	56.4	298	CGC	[1999VER/HEI]	
C ₁₁ H ₂₂ O ₂	[112-37-8] $\Delta_{\text{trs}} H$	undecanoic acid	8.13	290			
	$\Delta_{\text{fus}} H$		25.98	301.6		[1996DOM/HEA]	
	$\Delta_{\text{sub}} H$	(303–308)	121.3 ± 1.3	298	ME	[1968BAC/NOV, 1970COX/PIL]	
	$\Delta_v H$	(393–557)	81.3	408	A	[1987STE/MAL]	
	$\Delta_v H$	(310–332)	90.7 ± 2.0	323	ME, TE	[1982DEK/SCH]	
	$\Delta_v H$	(303–308)	97.9 ± 6.3	305		[1968BAC/NOV]	
C ₁₁ H ₂₂ O ₃	[38611-89-1] $\Delta_v H$	butyl 2-butoxypropionate (373–398)	40.8	385	A	[1987STE/MAL, 1933HEN/MUR]	
C ₁₁ H ₂₂ O ₃	[14144-48-0] $\Delta_v H$	butyl 3-butoxypropionate (343–493)	57.6	358	A	[1987STE/MAL]	
C ₁₁ H ₂₂ O ₃	[14144-37-7] $\Delta_v H$	hexyl 3-ethoxypropionate (373–514)	56.7	388	A	[1987STE/MAL]	
C ₁₁ H ₂₂ O ₃	[51191-33-4] $\Delta_v H$	octyl lactate (328–528)	71.5	343	A	[1987STE/MAL]	
C ₁₁ H ₂₂ O ₃	[676-08-4] $\Delta_{\text{sub}} H$	peroxyundecanoic acid (293–303)	125.9 ± 3.4	298	ME	[1980SWA/KWA]	
C ₁₁ H ₂₂ O ₃	[3669-80-5] $\Delta_{\text{sub}} H$	11-hydroxyundecanoic acid (307–321)	105		TPTD	[2005CHA/ZIE]	
C ₁₁ H ₂₃ Br	[693-67-4] $\Delta_{\text{fus}} H$	1-bromoundecane	33.47	263.3		[1950CRO/SMY]	
	$\Delta_v H$		(407–564)	58.8	422		[1999DYK/SVO]
	$\Delta_v H$		(398–591)	59.5	413	A, E	[1987STE/MAL, 1961LI/ROS, 1970DYK/VAN]
C ₁₁ H ₂₃ Cl	[2473-03-2] $\Delta_v H$	1-chloroundecane	70.2	298		[2006BOL/NER2]	
	$\Delta_v H$		(370–520)	65.9	298		[1984BOU/FRI, 1991BAS/SVO]
	$\Delta_v H$		(374–519)	59.4	389	A, DTA	[1987STE/MAL, 1969KEM/KRE]
C ₁₁ H ₂₃ F	[506-05-8] $\Delta_v H$	1-fluoroundecane (373–523)	52.3	388	A, E	[1987STE/MAL, 1961LI/ROS, 1970DYK/VAN]	
C ₁₁ H ₂₃ I	[4282-44-4] $\Delta_v H$	1-iodoundecane (412–618)	74.8	298	A,E	[1987STE/MAL, 1961LI/ROS, 1970DYK/VAN, 2006BOL/NER]	

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound		T _m (K)	Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)			
	$\Delta_v H$	(422–589)	60.1	437		[1999DYK/SVO]
	$\Delta_v H$	(412–618)	60.9	427	A, E	[1987STE/MAL, 1961LI/ROS, 1970DYK/VAN]
C₁₁H₂₃NO	[6225-08-7] $\Delta_v H$	N,N-dimethyl nonamide (411–509)	69.3	426	A	[1987STE/MAL]
C₁₁H₂₃NO	[23220-25-9] $\Delta_{\text{sub}} H$	N-methyl decanamide (303–325)	102.8 ± 0.8	314	GS	[1959DAV/JON, 1987STE/MAL]
C₁₁H₂₃NO₂	[6288-16-0] $\Delta_v H$	N,N-dibutyl lactamide (393–418)	88.3	405	A	[1987STE/MAL]
C₁₁H₂₃NO₂	[6280-23-5] $\Delta_v H$	N-octyl lactamide (428–468)	96.3	443	A	[1987STE/MAL]
C₁₁H₂₄	[1120-21-4] $\Delta_{\text{fus}} H$	undecane	0.1	236.3		
	$\Delta_{\text{fus}} H$		7.0	237.4		
	$\Delta_{\text{fus}} H$		22.5	247.6	DSC	[2004MON/RAJ]
	$\Delta_{\text{us}} H$		6.86	236.6		
	$\Delta_{\text{fus}} H$		22.18	247.6		[1996DOM/HEA]
	$\Delta_{\text{sub}} H$		91.5	236	B	[1963BON]
	$\Delta_v H$		56.4 ± 0.4	298	C	[2007PAS/KUZ]
	$\Delta_v H$		56.6 ± 0.6	298	C	[2006RIB/CAB2]
	$\Delta_v H$		56.2	299	C	[1996VIT/CHA]
	$\Delta_v H$		55.4	314	C	[1996VIT/CHA]
	$\Delta_v H$		54.5	324	C	[1996VIT/CHA]
	$\Delta_v H$		54.0	334	C	[1996VIT/CHA]
	$\Delta_v H$		53.1	344	C	[1996VIT/CHA]
	$\Delta_v H$		56.6	298		[1994RUZ/MAJ]
	$\Delta_v H$	(278–470)	60.0	293	A	[1987STE/MAL]
	$\Delta_v H$		56.3	298		[1971WIL/ZWO]
	$\Delta_v H$	(378–470)	49.1	393		[1955CAM/ROS]
C₁₁H₂₄	[6975-98-0] $\Delta_{\text{fus}} H$	2-methyldecane	25.06	224.3		[1996DOM/HEA]
	$\Delta_v H$	(273–353)	55.5	288	A	[1987STE/MAL]
	$\Delta_v H$	(379–463)	47.4	394	A	[1987STE/MAL]
	$\Delta_v H$		51.9	328	C	[1984MAJ/SVO3]
	$\Delta_v H$		50.6	343	C	[1984MAJ/SVO3]
	$\Delta_v H$		49.5	358	C	[1984MAJ/SVO3]
	$\Delta_v H$	(273–293)	55.4	283	IP	[1974OSB/DOU]
C₁₁H₂₄	[13151-34-3] $\Delta_v H$	3-methyldecane (340–464)	46.5	355	A	[1987STE/MAL]
C₁₁H₂₄	[2847-72-5] $\Delta_v H$	4-methyldecane (339–460)	46.6	354	A	[1987STE/MAL]
	$\Delta_v H$		50.4	343	C	[1984MAJ/SVO3]
	$\Delta_v H$		49.2	358	C	[1984MAJ/SVO3]
	$\Delta_v H$		48.5	368	C	[1984MAJ/SVO3]
C₁₁H₂₄	[13151-35-4] $\Delta_v H$	5-methyldecane (334–452)	46.0	349	A	[1987STE/MAL]
C₁₁H₂₄	[2884-06-2]	2,3-dimethylnonane				

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(336–460)	45.1	351	A	[1987STE/MAL]
C ₁₁ H ₂₄	[17302-24-8]	2,4-dimethylnonane				
	$\Delta_v H$	(334–452)	46.8	349	A	[1987STE/MAL]
C ₁₁ H ₂₄	[62016-37-9]	2,4,6-trimethyloctane				
	$\Delta_v H$	(325–442)	44.9	340	A	[1987STE/MAL]
C ₁₁ H ₂₄	[62016-38-0]	2,4,7-trimethyloctane				
	$\Delta_v H$		47.6	328	C	[1984MAJ/SVO3]
	$\Delta_v H$		46.4	343	C	[1984MAJ/SVO3]
	$\Delta_v H$		45.3	358	C	[1984MAJ/SVO3]
C ₁₁ H ₂₄ N ₂ O	[17450-44-1]	1-decyl urea				
	$\Delta_{\text{us}} H$		1.3	294.4		
	$\Delta_{\text{fus}} H$		38.3	385.3	DSC	[2005HAS/TAJ]
C ₁₁ H ₂₄ O	[7289-52-3]	decyl methyl ether				
	$\Delta_{\text{fus}} H$		31.71	243.5		[1996DOM/HEA]
	$\Delta_v H$	(341–429)	56.9	356	A	[1987STE/MAL]
	$\Delta_v H$	(341–471)	57.0	356	A	[1987STE/MAL, 1976AMB/ELL]
	$\Delta_v H$	(341–471)	62.6	298		[1976AMB/ELL]
	$\Delta_v H$	(341–471)	45.5	489		[1976AMB/ELL]
	$\Delta_v H$		62.3 ± 0.3	298	C	[1975FEN/HAR]
C ₁₁ H ₂₄ O	[16979-32-1]	ethyl nonyl ether				
	$\Delta_v H$		60.3 ± 0.1	298	C	[1985KUS]
C ₁₁ H ₂₄ O	[29379-41-7]	propyl octyl ether				
	$\Delta_v H$		58.8 ± 0.1	298	C	[1985KUS]
C ₁₁ H ₂₄ O	[71112-90-8]	butyl heptyl ether				
	$\Delta_v H$		58.2 ± 0.1	298	C	[1985KUS]
C ₁₁ H ₂₄ O	[78972-97-1]	heptyl <i>tert</i> -butyl ether				
	$\Delta_v H$		56.6	298	CGC	[UR/VER, 2002VER, 2003VER/KRA]
C ₁₁ H ₂₄ O	[na]	hexyl <i>tert</i> -amyl ether				
	$\Delta_v H$		58.6	298	CGC	[UR/VER, 2002VER, 2003VER/KRA]
C ₁₁ H ₂₄ O	[na]	propyl <i>tert</i> -octyl ether				
	$\Delta_v H$		50.1 ± 0.3	298	CGC	[UR/VER, 2002VER, 2003VER/KRA]
C ₁₁ H ₂₄ O	[112-42-5]	1-undecanol				
	$\Delta_v H$		85.8 ± 2.1	298	CGC	[2006NIC/KWE]
	$\Delta_v H$	(313–354)	79.5	336	GS	[2001KUL/VER2]
	$\Delta_v H$	(313–354)	84.7	298	GS	[2001KUL/VER2]
	$\Delta_v H$	(373–423)	86.8	298	CGC	[1995CHI/HOS]
	$\Delta_v H$	(353–393)	85.6	298	CGC	[1994KOU/HOS, 2000OVA/KOU]
	$\Delta_v H$	(293–342)	83.6	318		[1992NGU/KAS]
	$\Delta_v H$	(283–393)	83.5	298		[1999NGU/BER]
	$\Delta_v H$	(393–523)	68.7	408	A	[1987STE/MAL]
	$\Delta_v H$	(393–534)	68.5	408	A	[1987STE/MAL]
	$\Delta_v H$	(393–516)	72.3	408		[1973WIL/ZWO]
	C ₁₁ H ₂₄ O	[1653-30-1]	2-undecanol			
$\Delta_v H$		(344–505)	61.4	359		[1947STU]
C ₁₁ H ₂₄ O	[57233-26-8]	2,5-dimethyl-3-isopropyl-3-hexanol				
	$\Delta_v H$	(321–458)	57.2	336		[1973WIL/ZWO]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₁ H ₂₄ O	[5457-41-0] $\Delta_v H$	2,2,4-trimethyl-3-isopropyl-3-pentanol (321–458)	67.1	336		[1973WIL/ZWO]
C ₁₁ H ₂₄ O ₂ S	[54581-75-8] $\Delta_{\text{fus}} H$	3-(octylthio)-1,2-propanediol	39.8	306.5	DSC	[1993ACR]
C ₁₁ H ₂₄ O ₃	[10438-94-5] $\Delta_{\text{fus}} H$	3-(octyloxy)-1,2-propanediol	33.4	296.1	DSC	[1993ACR]
C ₁₁ H ₂₄ O ₄	[75899-69-3] $\Delta_v H$	tripropylene glycol, monoethyl ether (317–521)	60.0	332	A	[1987STE/MAL]
C ₁₁ H ₂₄ S	[5332-52-5] $\Delta_v H$	1-undecanethiol (405–563)	59.3	420		[1999DYK/SVO]
C ₁₁ H ₂₄ S ₂	[63476-06-2] $\Delta_v H$	1,11-undecanedithiol (444–582)	75.1	459	A	[1987STE/MAL, 1943HAL/REI, 1999DYK/SVO]
C ₁₁ H ₂₅ N	[7307-55-3] $\Delta_v H$	undecylamine (428–527)	55.1	443	A, E	[1987STE/MAL, 1956MAN2]
C ₁₁ H ₂₅ NO ₂	[929-31-7] $\Delta_{\text{fus}} H$	3-(octylamino)-1,2-propanediol	45.1	335.9	DSC	[1993ACR]
C ₁₁ H ₂₆ NO ₂ PS	[50782-69-9] $\Delta_v H$ $\Delta_v H$	methylthiophosphonic acid, O-ethyl-S-[2-(N,N-diisopropylamino)ethyl] ester (261–385) (280–315)	77.9 101	323 295	GC A	[2001RIT] [1987STE/MAL, 1999DYK/SVO, 1974FRO]
C ₁₁ H ₂₆ NO ₂ PS	[159939-87-4] $\Delta_v H$	P-methylphosphonothioic acid, S-[2(diethylamino)ethyl] O-(2-methylpropyl) ester (263–385)	76.6	324	GC	[2001RIT]
C ₁₁ H ₂₆ N ₂	[822-08-2] $\Delta_{\text{fus}} H$	undecane-1,11-diamine	48.08	313.6	DSC	[2002DAL/DEL]
C ₁₂ Cl ₈ O	[39001-02-2] $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$	octachlorodibenzofuran (438–473) (373–474)	141.7 ± 1.8 149.4	455 423	ME T	[2004LI/SHI] [1989ROR, 1986ROR]
C ₁₂ Cl ₈ O ₂	[3268-87-9] $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$	octachlorodibenzo[b,e][1,4]dioxin (463–493) (393–573)	145.7 ± 4.0 149.8	478 483	ME T	[2004LI/SHI] [1989ROR, 1986ROR]
C ₁₂ Cl ₁₀	[2051-24-3] $\Delta_{\text{fus}} H$ $\Delta_{\text{fus}} H$ $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$ $\Delta_v H$ $\Delta_v H$	decachlorobiphenyl (338–358) (324–363) (343–393) (343–453)	39.34 41.2 U 93.6 121.8 103.4 103.4	577.7 580.3 348 343 368 398	DSC DSC ME GS GC GC	[1991ACR] [1990DON/DRE] [1997GOO] [1984BUR/ARM] [1994FAL/BID] [1990HIN/BID2]
C ₁₂ F ₁₀	[434-90-2] $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$ $\Delta_v H$	decafluorobiphenyl (297–323) (453–608)	87.8 85.3 ± 2.3 49.9	310 468	A DSC	[1987STE/MAL] [1974RAD/KAT] [1996BAC/GRZ]
C ₁₂ F ₁₈	[23174-55-2] $\Delta_v H$	hexakis(trifluoromethyl)bicyclo[2.2.0]hexa-2,5-diene (293–343)	41.4	308	A	[1987STE/MAL, 1970BAR/HAS, 1999DYK/SVO]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₂ F ₁₈	[22736-20-5]	<i>hexakis</i> (trifluoromethyl)tetracyclo[2.2.0.0 ^{2,6} .0 ^{3,5}]hexane				
	$\Delta_{\text{sub}}H$	(293–306)	49.2	299.5	A	[1987STE/MAL]
	Δ_vH	(313–353)	33.1	328	A	[1987STE/MAL, 1970BAR/HAS, 1999DYK/SVO]
C ₁₂ F ₁₈	[22186-64-7]	<i>hexakis</i> (trifluoromethyl)tricyclo[3.1.0.0 ^{2,6}]hex-3-ene				
	Δ_vH	(293–353)	38.6	308	A	[1987STE/MAL, 1970BAR/HAS, 1999DYK/SVO]
C ₁₂ F ₂₃ N	[86630-50-4]	perfluoro-N-(4-methylcyclohexyl)piperidine				
	$\Delta_{\text{fus}}H$		8.32	293.3	AC	[2007DRU/EFI]
	$\Delta_{\text{fus}}H$		8.6	293.7	DSC	[2007DRU/EFI]
	Δ_vH		56.6	298	C	[2007DRU/EFI]
C ₁₂ F ₂₆	[307-59-5]	perfluorododecane				
	$\Delta_{\text{fus}}H$		6.9	170.2		
	$\Delta_{\text{fus}}H$		38.16	348.5	DSC	[1986STA]
C ₁₂ F ₂₆ O ₁₀	[927699-30-7]	perfluoro-2,4,6,8,10,13,15,17,19,21-decaoxy-n-docosane				
	Δ_vH	(397–468)	74.6 ± 2.9	298	EB	[2006DRU/KRO]
C ₁₂ F ₂₇ N	[311-89-7]	perfluorotributylamine				
	Δ_vH		60.3 ± 0.1	298	C	[1995VAR/DRO]
	Δ_vH	(298–450)	57.4	313	A	[1987STE/MAL]
	Δ_vH	(371–544)	51.1	386	A	[1987STE/MAL]
	Δ_vH		60.4 ± 1.2	298		[1977VAR/AMM2, 1977VAR/AMM]
C ₁₂ HCl ₇ O ₂	[58200-70-7]	1,2,3,4,6,7,9-heptachlorodibenzo[b,e] [1,4]dioxin				
	$\Delta_{\text{sub}}H$	(418–453)	144.2 ± 0.3	435	ME	[2004LI/SHI]
C ₁₂ HCl ₉	[52663-77-1]	2,2',3,3',4,5,5',6,6'-nonachlorobiphenyl				
	$\Delta_{\text{fus}}H$		22.6	455.8		[1991ACR]
C ₁₂ HF ₂₅	[66563-68-6]	1,1,1,2,2,3,3,4,4,5,5,6,6,7,7,8,8,9,9,10,10,11,11,12,12-pentacosofluorododecane				
	$\Delta_{\text{fus}}H$		21	345		[1988HOP/PUG]
	$\Delta_{\text{fus}}H$		23	344.5	DSC	[1986RUS/RAB]
C ₁₂ H ₂ Cl ₆ O ₂	[58200-68-3]	1,2,3,4,6,9-hexachlorodibenzo[b,e] [1,4]dioxin				
	$\Delta_{\text{sub}}H$	(418–438)	128.5 ± 1.5	428	ME	[2004LI/SHI]
C ₁₂ H ₂ Cl ₈	[2136-99-4]	2,2',3,3',5,5',6,6'-octachlorobiphenyl				
	$\Delta_{\text{fus}}H$		22.8	433.8		[1991ACR]
	$\Delta_{\text{sub}}H$	(302–334)	101.7	318	GS	[1984BUR/ARM]
	Δ_vH	(343–393)	92.9	368	GC	[1994FAL/BID]
	Δ_vH	(343–453)	92.9	398	GC	[1990HIN/BID2]
C ₁₂ H ₃ Br ₇ O	[327185-13-7]	2',3,3',4,4',5,6-heptabromodiphenyl ether				
	Δ_vH	(363–473)	115.8	418	GC	[2001WON/LEI]
	Δ_vH	(403–475)	121.2		CGC	[2001TIT/TOM]
C ₁₂ H ₃ Cl ₅ O ₂	[58802-08-7]	1,2,4,7,8-pentachlorodibenzo[b,e] [1,4]dioxin				
	$\Delta_{\text{sub}}H$	(403–428)	125.3 ± 2.3	415	ME	[2004LI/SHI]
C ₁₂ H ₃ Cl ₇	[52663-68-0]	2,2',3,4',5,5',6-heptachlorobiphenyl				
	Δ_vH	(343–393)	94.0	368	GC	[1994FAL/BID]
C ₁₂ H ₃ Cl ₇	[35065-29-3]	2,2',3,4,4',5,5'-heptachlorobiphenyl				
	Δ_vH	(343–393)	96.5	268	GC	[1994FAL/BID]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound		T _m (K)	Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)			
C ₁₂ H ₃ Cl ₇	[52663-71-5]	2,2',3,3',4,4',6-heptachlorobiphenyl		298	CGC	[2001PUR/CHI]
	$\Delta_v H$	(343–393)	109.1 95.9	368	GC	[1994FAL/BID]
C ₁₂ H ₃ Cl ₇	[35065-30-6]	2,2',3,3',4,4',5-heptachlorobiphenyl		368	GC	[1994FAL/BID]
	$\Delta_v H$	(343–393)	98.4			
C ₁₂ H ₃ Cl ₇	[52663-67-9]	2,2',3,3',5,5',6-heptachlorobiphenyl		395.4		[1991ACR]
	$\Delta_{\text{fus}} H$		20.3			
C ₁₂ H ₃ Cl ₇ O	[67517-48-0]	1,2,3,4,8-pentachlorodibenzofuran		400	ME	[2004LI/SHI]
	$\Delta_{\text{sub}} H$	(388–413)	125.2 ± 2.0			
C ₁₂ H ₄ Br ₆ O	[182677-30-4]	2,2',3,4,4',5'-hexabromodiphenyl ether			CGC	[2001TIT/TOM]
	$\Delta_v H$	(403–475)	114.1			
C ₁₂ H ₄ Br ₆ O	[68631-49-2]	2,2',4,4',5,5'-hexabromodiphenyl ether		418	GC	[2001WON/LEI]
	$\Delta_v H$	(363–473)	107.6			
C ₁₂ H ₄ Cl ₂ F ₆ N ₄ OS	[120068-37-3]	1-(2,6-dichloro-4-trifluoromethylphenyl)-3-cyano-5-amino-4-(trifluoromethylsulfinyl)pyrazole (fipronil)			GC	[2007GOE/MCC]
	$\Delta_v H$	(373–423)	85.0			
C ₁₂ H ₄ Cl ₄ O	[24478-72-6]	1,2,3,4-tetrachlorodibenzofuran		363	T	[1989ROR, 1986ROR]
	$\Delta_{\text{sub}} H$	(333–393)	118.5			
C ₁₂ H ₄ Cl ₄ O	[51207-31-9]	2,3,7,8-tetrachlorodibenzofuran		323	T	[1989ROR, 1986ROR]
	$\Delta_{\text{sub}} H$	(303–344)	124			
C ₁₂ H ₄ Cl ₄ O ₂	[30746-58-8]	1,2,3,4-tetrachlorodibenzo[b,e][1,4]dioxin		390	ME	[2004LI/SHI]
	$\Delta_{\text{sub}} H$	(378–403)	111.3 ± 1.4			
C ₁₂ H ₄ Cl ₄ O ₂	[40581-90-6]	1,2,6,7-tetrachlorodibenzo[b,e][1,4]dioxin		403	ME	[2004LI/SHI]
	$\Delta_{\text{sub}} H$	(393–413)	120.4 ± 3.3			
C ₁₂ H ₄ Cl ₄ O ₂	[33423-92-6]	1,3,6,8-tetrachlorodibenzo[b,e][1,4]dioxin		393	ME	[2004LI/SHI]
	$\Delta_{\text{sub}} H$	(378–408)	118.6 ± 3.2			
C ₁₂ H ₄ Cl ₄ O ₂	[62470-53-5]	1,3,7,9-tetrachlorodibenzo[b,e][1,4]dioxin		395	ME	[2004LI/SHI]
	$\Delta_{\text{sub}} H$	(383–408)	123.6 ± 1.5			
C ₁₂ H ₄ Cl ₄ O ₂	[1746-01-6]	2,3,7,8-tetrachlorodibenzo[b,e][1,4]dioxin		578.2		[1986ROR2]
	$\Delta_{\text{fus}} H$		38.9	578		
	$\Delta_{\text{sub}} H$		124			[1985SCH/HIL]
C ₁₂ H ₄ Cl ₆	[38380-08-4]	2,3,3',4,4',5-hexachlorobiphenyl		298	CGC	[2001PUR/CHI]
	$\Delta_v H$	(343–393)	112.6 ± 0.4 94.8	368	GC	[1994FAL/BID]
	$\Delta_v H$					
C ₁₂ H ₄ Cl ₆	[35065-27-1]	2,2',4,4',5,5'-hexachlorobiphenyl		298	CGC	[2001PUR/CHI]
	$\Delta_v H$	(343–393)	103.5 ± 0.1 91.4	368	GC	[1994FAL/BID]
	$\Delta_v H$					
C ₁₂ H ₄ Cl ₆	[33976-03-2]	2,2',4,4',6,6'-hexachlorobiphenyl		386.7		[1991ACR]
	$\Delta_{\text{fus}} H$		17.5	283	GS	[1994WAN/SHU]
	$\Delta_{\text{sub}} H$	(263–303)	103.4 ± 2.3			
C ₁₂ H ₄ Cl ₆	[38380-04-0]	2,2',3,4',5',6-hexachlorobiphenyl		368	GC	[1994FAL/BID]
	$\Delta_v H$	(343–393)	89.8			
C ₁₂ H ₄ Cl ₆	[35065-28-2]	2,2',3,4,4',5'-hexachlorobiphenyl		368	GC	[1994FAL/BID]
	$\Delta_v H$	(343–393)	91.9			

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₂ H ₄ Cl ₆	[38380-07-3] $\Delta_v H$	2,2',3,3',4,4'-hexachlorobiphenyl (343–393)	93.5	368	GC	[1994FAL/BID]
C ₁₂ H ₄ Cl ₆	[35694-04-3] $\Delta_{\text{fus}} H$	2,2',3,3',5,5'-hexachlorobiphenyl	29.2	424.9		[1991ACR]
C ₁₂ H ₄ Cl ₆	[38411-22-2] $\Delta_{\text{fus}} H$	2,2',3,3',6,6'-hexachlorobiphenyl	21.1	385.2		[1991ACR]
C ₁₂ H ₄ N ₄	[1518-16-7] $\Delta_{\text{sub}} H$	7,7,8,8-tetracyanoquinodimethane	79		TGA	[1995YAS/TAK]
	$\Delta_{\text{sub}} H$	(452–553)	108 ± 2	500	T	[1984KER/OPP]
	$\Delta_{\text{sub}} H$	(382–464)	122 ± 2	423	ME	[1984KER/OPP]
	$\Delta_{\text{sub}} H$		126.1 ± 1	413	ME,TE	[1980DEK/GOV]
	$\Delta_{\text{sub}} H$	(433–499)	104.8 ± 10	448		[1980SWA/KWA, 1987STE/MAL]
	$\Delta_{\text{sub}} H$		105 ± 9.2	465	MG	[1963BOY, 1970COX/PIL]
C ₁₂ H ₅ Br ₅ O	[na] $\Delta_v H$	2,2',3,4,4'-pentabromodiphenyl ether (403–475)	111		CGC	[2001TIT/TOM]
C ₁₂ H ₅ Br ₅ O	[327185-11-5] $\Delta_v H$	2,2',3,3',4-pentabromodiphenyl ether (363–473)	99.1	418	GC	[2001WON/LEI]
C ₁₂ H ₅ Br ₅ O	[60348-60-9] $\Delta_v H$	2,2',4,4',5-pentabromodiphenyl ether (363–473)	100.3	418	GC	[2001WON/LEI]
	$\Delta_v H$	(405–475)	104.8		CGC	[2001TIT/TOM]
C ₁₂ H ₅ Br ₅ O	[189084-66-0] $\Delta_v H$	2,2',4,4',6-pentabromodiphenyl ether (363–473)	101.8	418	GC	[2001WON/LEI]
C ₁₂ H ₅ Cl ₃ O	[58802-14-5] $\Delta_{\text{sub}} H$	2,4,6-dibenzofuran (338–373)	108.8 ± 2.2	355	ME	[2004LI/SHI]
C ₁₂ H ₅ Cl ₃ O ₂	[54536-17-3] $\Delta_{\text{sub}} H$	1,2,3-trichlorodibenzo[b,e] [1,4]dioxin (363–388)	117.1 ± 3.7	375	ME	[2004LI/SHI]
C ₁₂ H ₅ Cl ₃ O ₂	[39227-58-2] $\Delta_{\text{sub}} H$	1,2,4-trichlorodibenzo[b,e] [1,4]dioxin (348–383)	121.0 ± 1.8	365	ME	[2004LI/SHI]
	$\Delta_{\text{sub}} H$	(310–374)	118.8	342	T	[1989ROR, 1986ROR]
C ₁₂ H ₅ Cl ₃ O ₂	[67028-17-5] $\Delta_{\text{fus}} H$	1,3,7-trichlorodibenzo[b,e] [1,4]dioxin	30.8	421.7		[1986ROR2]
	$\Delta_{\text{sub}} H$	(310–373)	116.2	342	T	[1989ROR, 1986ROR]
C ₁₂ H ₅ Cl ₃ O ₂	[82306-65-8] $\Delta_{\text{sub}} H$	1,7,8-trichlorodibenzo[b,e] [1,4]dioxin (358–388)	113.5 ± 3.3	373	ME	[2004LI/SHI]
C ₁₂ H ₅ Cl ₅	[31508-00-6] $\Delta_v H$	2,3',4,4',5-pentachlorobiphenyl (343–393)	89.3	368	GC	[1994FAL/BID]
C ₁₂ H ₅ Cl ₅	[32598-14-4] $\Delta_v H$	2,3,3',4,4'-pentachlorobiphenyl (343–393)	91.1	368	GC	[1994FAL/BID]
C ₁₂ H ₅ Cl ₅	[37680-73-2] $\Delta_{\text{fus}} H$	2,2',4,5,5'-pentachlorobiphenyl	18.8	350.1		[1991ACR]
	$\Delta_{\text{sub}} H$	(303–313)	92.7	308	GS	[1981WES/SIM]
	$\Delta_v H$	(343–393)	86.4	368	GC	[1994FAL/BID]
	$\Delta_v H$	(343–453)	83.7	398	GC	[1990HIN/BID2]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound		T _m (K)	Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)			
C ₁₂ H ₅ Cl ₅	[38379-99-6] $\Delta_v H$	2,2',3,5',6-pentachlorobiphenyl	92.3 ± 0.6	298	CGC	[2001PUR/CHI]
C ₁₂ H ₅ Cl ₅	[73575-54-9] $\Delta_v H$	2,2',3,6,6'-pentachlorobiphenyl	89.6 ± 0.2	298	CGC	[2001PUR/CHI]
C ₁₂ H ₅ Cl ₅	[60145-21-3] $\Delta_v H$	2,2',4,5',6-pentachlorobiphenyl	91.6 ± 0.5	298	CGC	[2001PUR/CHI]
C ₁₂ H ₅ Cl ₅	[38380-02-8] $\Delta_v H$	2,2',3,4,5'-pentachlorobiphenyl (343–393)	87.3	368	GC	[1994FAL/BID]
C ₁₂ H ₅ Cl ₅	[38380-01-7] $\Delta_v H$	2,2',4,4',5-pentachlorobiphenyl (343–393)	86.8	368	GC	[1994FAL/BID]
C ₁₂ H ₅ Cl ₅	[18259-05-7] $\Delta_{\text{fus}} H$	2,3,4,5,6-pentachlorobiphenyl	21.8	397.6		[1991ACR]
C ₁₂ H ₆ Br ₄ O	[5436-43-1] $\Delta_v H$ $\Delta_v H$	2,2',4,4'-tetrabromodiphenyl ether (363–473) (403–475)	92.0 103.1	418	GC CGC	[2001WON/LEI] [2001PUR/CHI]
C ₁₂ H ₆ Br ₄ O	[189084-61-5] $\Delta_v H$	2,3',4,4'-tetrabromodiphenyl ether (363–473)	93.5	418	GC	[2001WON/LEI]
C ₁₂ H ₆ Br ₄ O	[327185-09-1] $\Delta_v H$	2,3',4,6-tetrabromodiphenyl ether (363–473)	91.1	418	GC	[2001WON/LEI]
C ₁₂ H ₆ Br ₄ O	[189084-63-7] $\Delta_v H$	2,4,4',6-tetrabromodiphenyl ether (363–473)	90.1	418	GC	[2001WON/LEI]
C ₁₂ H ₆ Br ₄ O	[93703-48-1] $\Delta_v H$	3,3',4,4'-tetrabromodiphenyl ether (363–473)	95.3	418	GC	[2001WON/LEI]
C ₁₂ H ₆ Cl ₂ O	[5409-83-6] $\Delta_{\text{sub}} H$	2,8-dichlorodibenzofuran (348–383)	110.3 ± 1.2	360	ME	[2004LI/SHI]
C ₁₂ H ₆ Cl ₂ O	[74919-40-4] $\Delta_{\text{fus}} H$ $\Delta_{\text{sub}} H$	3,6-dichlorodibenzofuran (305–374)	32.4 110.9	461.2 340		[1986ROR2] [1989ROR, 1986ROR]
C ₁₂ H ₆ Cl ₂ O ₂	[38178-38-0] $\Delta_{\text{sub}} H$	1,6-dichlorodibenzo[b,e][1,4]dioxin (348–383)	113.6 ± 2.3	365	ME	[2004LI/SHI]
C ₁₂ H ₆ Cl ₂ O ₂	[29446-15-9] $\Delta_{\text{fus}} H$ $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$	2,3-dichlorodibenzo[b,e][1,4]dioxin (338–378)	27.1 106.2 ± 1.1 108.6 ± 1.0 107.2 ± 0.8 108.6 ± 1.0 106.2	431.6 358 298 358 298 340		[1999KOL/DOR] [2004LI/SHI] [1999KOL/DOR] [1998PAP/KOL] [1998PAP/KOL] [1989ROR, 1986ROR]
C ₁₂ H ₆ Cl ₂ O ₂	[33857-26-0] $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$	2,7-dichlorodibenzo[b,e][1,4]dioxin (358–393) (314–374)	113.8 ± 2.0 105.5	375 344	ME T	[2004LI/SHI] [1989ROR, 1986ROR]
C ₁₂ H ₆ Cl ₂ O ₂	[38964-22-6] $\Delta_{\text{sub}} H$	2,8-dichlorodibenzo[b,e][1,4]dioxin (305–363)	109	334	T	[1989ROR, 1986ROR]
C ₁₂ H ₆ Cl ₄	[38444-93-8]	2,2',3,3'-tetrachlorobiphenyl				

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(343–398)	81.8	368	GC	[1994FAL/BID]
C ₁₂ H ₆ Cl ₄	[35693-99-3]	2,2',5,5'-tetrachlorobiphenyl				
	$\Delta_{\text{sub}} H$	(323–353)	102.0 ± 0.5	338	ME	[2005NAK/SHI]
	$\Delta_{\text{sub}} H$	(303–312)	94.6	308	GS	[1981WES/SIM]
	$\Delta_v H$	(343–398)	80.8	368	GC	[1994FAL/BID]
C ₁₂ H ₆ Cl ₄		(343–453)	79	398	GC	[1990HIN/BID2]
	[33284-53-6]	2,3,4,5-tetrachlorobiphenyl				
	$\Delta_{\text{fus}} H$		25.2	363.9		[1991ACR]
C ₁₂ H ₆ Cl ₄		(253–393)	88.7 ± 1.2	273	GS	[1994WAN/SHU]
	[32598-10-0]	2,3',4,4'-tetrachlorobiphenyl				
	$\Delta_{\text{sub}} H$	(348–373)	105.9 ± 2.5	353	ME	[2005NAK/SHI]
C ₁₂ H ₆ Cl ₄		(343–398)	83.3	368	GC	[1994FAL/BID]
	[32598-11-1]	2,3',4',5-tetrachlorobiphenyl				
C ₁₂ H ₆ Cl ₄		(343–398)	84.8	368	GC	[1994FAL/BID]
	[41464-40-8]	2,2',4,5'-tetrachlorobiphenyl				
C ₁₂ H ₆ Cl ₄			23.4	339.1		[1991ACR]
	$\Delta_v H$		87.4 ± 0.8	298	CGC	[2001PUR/CHI]
	[41464-41-9]	2,2',5,6'-tetrachlorobiphenyl				
C ₁₂ H ₆ Cl ₄			84.9 ± 0.6	298	CGC	[2001PUR/CHI]
	$\Delta_v H$	(343–398)	78.8	368	GC	[1994FAL/BID]
C ₁₂ H ₆ Cl ₄	[32598-13-3]	3,3',4,4'-tetrachlorobiphenyl				
	$\Delta_{\text{sub}} H$	(383–403)	121.6 ± 1.3	393	ME	[2005NAK/SHI]
C ₁₂ H ₆ Cl ₄ O ₂ S		(343–393)	87.2	368	GC	[1994FAL/BID]
	[116-29-0]	1,2,4-trichloro-5-((4-chlorophenyl)sulfonyl)benzene				
C ₁₂ H ₆ O ₃			28.94	419.9	DSC	[1991ACR]
	[81-84-5]	1-8-naphthalic anhydride (protect)				
C ₁₂ H ₇ Br ₃ O			23.32	542.3	DSC	[1990DON/DRE]
	[41318-75-6]	2,4,4'-tribromodiphenyl ether				
C ₁₂ H ₇ Br ₃ O		(403–475)	94.1		CGC	[2001TIT/TOM]
	[147217-81-0]	3,4,4'-tribromodiphenyl ether				
C ₁₂ H ₇ Br ₃ O		(363–473)	86.7	418	GC	[2001WON/LEI]
	[155999-95-4]	2,4,6-tribromodiphenyl ether				
C ₁₂ H ₇ Br ₃ O		(363–473)	85.1	418	GC	[2001WON/LEI]
	[189084-60-4]	2,4',6-tribromodiphenyl ether				
C ₁₂ H ₇ Br ₃ O		(363–473)	83.3	418	GC	[2001WON/LEI]
	[147217-78-5]	2',3,4-tribromodiphenyl ether				
C ₁₂ H ₇ Br ₃ O		(363–473)	81.0	418	GC	[2001WON/LEI]
	[147217-80-9]	3,3',4-tribromodiphenyl ether				
C ₁₂ H ₇ ClO ₂		(363–473)	86.4	418	GC	[2001WON/LEI]
	[39227-53-7]	1-chlorodibenzo[b,e][1,4]dioxin				
	$\Delta_{\text{fus}} H$		23.2	378.2		[1986ROR2]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₂ H ₇ ClO ₂	$\Delta_{\text{sub}}H$	(308–343)	100.5 ± 0.8	325	ME	[2004LI/SHI]
	$\Delta_{\text{sub}}H$		95.2 ± 1.1	298	C	[1999KOL/DOR]
	$\Delta_{\text{sub}}H$		95.2			[1998PAP/LUK]
	$\Delta_{\text{sub}}H$	(303–338)	98.6	321	T	[1989ROR, 1986ROR]
	[39227-54-8]	2-chlorodibenzo[b,e] [1,4]dioxin				
	$\Delta_{\text{fus}}H$		23.1	362.2		[1986ROR2]
	$\Delta_{\text{sub}}H$	(308–343)	98.1 ± 1.1	298	ME	[2004LI/SHI]
	$\Delta_{\text{sub}}H$		97.2	298	C	[1999KOL/DOR]
C ₁₂ H ₇ Cl ₂ NO ₃	[1836-75-5]	2,4-dichlorophenyl 4-nitrophenyl ether				
	$\Delta_{\text{fus}}H$		22.96	342	DSC	[1990DON/DRE]
	Δ_vH	(328–403)	90.4	343	A	[1987STE/MAL]
C ₁₂ H ₇ Cl ₃	[15862-07-4]	2,4,5-trichlorobiphenyl				
	$\Delta_{\text{fus}}H$		22.8	349.5		[1991ACR]
	Δ_vH	(343–393)	76.6	368	GC	[1994FAL/BID]
C ₁₂ H ₇ Cl ₃	[35693-92-6]	2,4,6-trichlorobiphenyl				
	$\Delta_{\text{fus}}H$		16.5	334.3		[1991ACR]
	Δ_vH	(343–393)	74.4	368	GC	[1994FAL/BID]
C ₁₂ H ₇ Cl ₃	[16606-02-3]	2,4',5-trichlorobiphenyl				
	Δ_vH	(343–398)	77.7	368	GC	[1994FAL/BID]
C ₁₂ H ₇ Cl ₃	[36780-65-2]	2,2',5-trichlorobiphenyl				
	$\Delta_{\text{sub}}H$	(303–313)	93.7 ± 6.2	308	ME	[2005NAK/SHI]
	Δ_vH		80.2 ± 0.9	298	CGC	[2001PUR/CHI]
C ₁₂ H ₇ Cl ₃	[7012-37-5]	2,4,4'-trichlorobiphenyl				
	$\Delta_{\text{sub}}H$	(313–328)	96.7 ± 3.4	320	ME	[2005NAK/SHI]
C ₁₂ H ₇ Cl ₃	[38444-86-9]	2',3,4-trichlorobiphenyl				
	$\Delta_{\text{sub}}H$	(313–328)	98.2 ± 5.5	320	ME	[2005NAK/SHI]
C ₁₂ H ₈	[208-96-8]	acenaphthylene				
	$\Delta_{\text{trs}}H$		1.4	116.6		
	$\Delta_{\text{fus}}H$		6.95	362.6		
	$\Delta_{\text{fus}}H$		10.96	362		[1996DOM/HEA, 1994CHE/WES]
	$\Delta_{\text{sub}}H$		70.0	298	CGC-DSC	[1998CHI/HES]
	$\Delta_{\text{sub}}H$	(313–453)	77.2	383	GS	[1995NAS/LEN]
	$\Delta_{\text{sub}}H$	(238–323)	73.2 ± 0.5	303	GS	[1983SON/ZOL]
	$\Delta_{\text{sub}}H$		73.0 ± 0.4	298	C	[1972MOR]
	$\Delta_{\text{sub}}H$	(286–318)	71.1 ± 1.3		A	[1970COX/PIL, 1987STE/MAL, 1965BOY/CHR]
	Δ_vH		64.6 ± 5.8	298	CGC	[2008ROU/TEM]
	Δ_vH		69.1 ± 2.2	298	GC	[2006HAF/PAR]
C ₁₂ H ₈	[259-79-0]	biphenylene				
	$\Delta_{\text{sub}}H$	(313–453)	82.7	383	GS	[1995NAS/LEN]
	$\Delta_{\text{sub}}H$	(309–336)	U 104.5	319		[1989ROR/RUT]
	$\Delta_{\text{sub}}H$		87.3 ± 0.3	298	B	[1980OSB/SCO]
	$\Delta_{\text{sub}}H$		83.8 ± .3		C	[1972MOR]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{sub}}H$	(371–381)	U 128.9 ± 2	376	A	[1955CAS/SPR, 1970COX/PIL, 1987STE/MAL]
C ₁₂ H ₈ Br ₂	[na] $\Delta_{\text{fus}}H$	(dl) 1,2-dibromoacenaphthene	25.1	397		[1976LEC/COL]
C ₁₂ H ₈ Br ₂	[na] $\Delta_{\text{fus}}H$	(d) 1,2-dibromoacenaphthene	26.36	416		[1976LEC/COL]
C ₁₂ H ₈ Br ₂	[92-86-4] $\Delta_{\text{fus}}H$	4,4'-dibromobiphenyl	28.38	440.7	DSC	[2009RAI/RED]
C ₁₂ H ₈ Br ₂ O	[171977-44-9] Δ_vH	2,4-dibromodiphenyl ether (363–473)	75.4	418	GC	[2001WON/LEI]
C ₁₂ H ₈ Br ₂ O	[189084-59-1] Δ_vH	3,4-dibromodiphenyl ether (363–473)	77.4	418	GC	[2001WON/LEI]
C ₁₂ H ₈ Br ₂ O	[83694-71-7] Δ_vH	3,4'-dibromodiphenyl ether (363–473)	77.4	418	GC	[2001WON/LEI]
C ₁₂ H ₈ Br ₂ O	[2050-47-7] Δ_vH	4,4'-dibromodiphenyl ether (363–473)	78.0	418	GC	[2001WON/LEI]
C ₁₂ H ₈ Br ₂ O	[147217-71-8] Δ_vH	2,4'-dibromodiphenyl ether (363–473)	76.4	418	GC	[2001WON/LEI]
C ₁₂ H ₈ Br ₂ O	[51930-04-2] Δ_vH	2,6-dibromodiphenyl ether (363–473)	73.1	418	GC	[2001PUR/CHI]
C ₁₂ H ₈ Cl ₂	[13029-08-8] $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$	2,2'-dichlorobiphenyl (303–323) (310–328) (310–328)	87.8 ± 1.2 96.1 96.2 ± 4.2	313 314 298	ME ME ME	[2005NAK/SHI] [1964SMI/GOR] [1964SMI/GOR, 1970COX/PIL, 1987STE/MAL]
C ₁₂ H ₈ Cl ₂	[33284-50-3] Δ_vH Δ_vH	2,4-dichlorobiphenyl (343–393)	75.3 ± 1.5 73.5	298 368	CGC GC	[2001PUR/CHI] [1994FAL/BID]
C ₁₂ H ₈ Cl ₂	[34883-39-1] Δ_vH Δ_vH	2,5-dichlorobiphenyl (343–393)	76.8 ± 0.4 73.9	298 368	CGC GC	[2001PUR/CHI] [1994FAL/BID]
C ₁₂ H ₈ Cl ₂	[33146-45-1] $\Delta_{\text{fus}}H$	2,6-dichlorobiphenyl	12.6	307.9		[1991ACR]
C ₁₂ H ₈ Cl ₂	[2050-67-1] Δ_vH Δ_vH	3,3'-dichlorobiphenyl (343–393)	81.0 ± 0.2 75.4	298 368	CGC GC	[2001PUR/CHI] [1994FAL/BID]
C ₁₂ H ₈ Cl ₂	[2050-68-2] $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$ Δ_vH Δ_vH	4,4'-dichlorobiphenyl (263–303) (303–360) (303–360) (343–393)	95.3 ± 1.3 103.7 103.8 ± 4.2 81.4 ± 0.3 76.0	283 331 298 298 368	GS ME ME CGC GC	[1994WAN/SHU] [1964SMI/GOR, 1987STE/MAL] [1964SMI/GOR, 1970COX/PIL] [2001PUR/CHI] [1994FAL/BID]
C ₁₂ H ₈ Cl ₂	[na] $\Delta_{\text{fus}}H$	(dl) 1,2-dichloroacenaphthene	20.5	339		[1976LEC/COL]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₂ H ₈ Cl ₂	[na]	(d) 1,2-dichloroacenaphthene				
	$\Delta_{\text{fus}}H$		21.34	375		[1976LEC/COL]
C ₁₂ H ₈ Cl ₂ O ₂ S	[80-07-9]	4,4'-dichlorodiphenylsulfone				
	$\Delta_{\text{fus}}H$		24.4	422		[1996DOM/HEA]
		(463–573)	59.7	478		[1999DYK/SVO]
C ₁₂ H ₈ Cl ₂ O ₃ S	[80-33-1]	4-chlorophenyl 4-chlorobenzenesulfonate				
	$\Delta_{\text{fus}}H$		23.63	360	DSC	[1991ACR, 1990DON/DRE]
C ₁₂ H ₈ Cl ₃ NO ₂	[77765-39-0]	2,2,4-trichloro-5-[(2-methylphenyl)amino]-4-cyclopentene-1,3-dione				
	Δ_vH	(453–483)	85	468	GC	[1980SHA/SAD]
C ₁₂ H ₈ Cl ₃ NO ₃	[77765-40-3]	2,2,4-trichloro-5-[(2-methoxyphenyl)amino]-4-cyclopentene-1,3-dione				
	Δ_vH	(453–483)	84.6	468	GC	[1980SHA/SAD]
C ₁₂ H ₈ Cl ₃ NO ₃	[73373-64-5]	2,2,4-trichloro-5-[(3-methoxyphenyl)amino]-4-cyclopentene-1,3-dione				
	Δ_vH	(453–483)	63.1	468	GC	[1980SHA/SAD]
C ₁₂ H ₈ Cl ₆	[309-00-2]	1,2,3,4,10,10-hexachloro-1,4,4a,5,8,8a-hexahydro-endo-exo-1,4:5,8-dimethylnaphthalene (aldrin)				
	$\Delta_{\text{us}}H$		16.59	383.7		
	$\Delta_{\text{fus}}H$		4.15	562.4	DSC	[1995KSI/NAG]
	$\Delta_{\text{sub}}H$	(309–343)	91.8	326	GS	[1982GRA/FOS]
		(343–453)	75.1	398	GC	[1990HIN/BID2]
C ₁₂ H ₈ Cl ₆ O	[60-57-1]	1,2,3,4,10,10-hexachloro-6,7-epoxy-1,4,4a,5,6,7,8,8a-octahydro-1,4-endo-exo-5,8-dimethanonaphthalene (dieldrin)				
	$\Delta_{\text{us}}H$		19.33	405.6		
	$\Delta_{\text{fus}}H$		3.04	452.9	DSC	[1995KSI/NAG]
	$\Delta_{\text{sub}}H$	(308–348)	93.8	328	GS	[1982GRA/FOS]
	$\Delta_{\text{sub}}H$	(293–313)	98.7	303	GS	[1969SPE/CLI]
		(343–453)	82.5	398	GC	[1990HIN/BID2]
C ₁₂ H ₈ F ₂	[388-82-9]	2,2'-difluorobiphenyl				
	$\Delta_{\text{sub}}H$	(301–319)	95.1	310		[1987STE/MAL, 1964SMI/GOR]
	$\Delta_{\text{sub}}H$	(301–318)	95 ± 4.2	298	ME	[1964SMI/GOR, 1970COX/PIL]
C ₁₂ H ₈ F ₂	[2050-68-2]	4,4'-difluorobiphenyl				
	$\Delta_{\text{sub}}H$	(294–318)	91.4	306	ME	[1964SMI/GOR]
	$\Delta_{\text{sub}}H$	(294–318)	91.2 ± 4.2	298	ME	[1964SMI/GOR, 1970COX/PIL]
C ₁₂ H ₈ N ₂	[230-46-6]	1,7-phenanthroline				
	Δ_vH		79.4 ± 4.7	298	CGC	[2009LIP/CHI, 2009LIP/HAN]
C ₁₂ H ₈ N ₂	[66-71-7]	1,10-phenanthroline				
	$\Delta_{\text{fus}}H$		15.5	391.7	AC	[2010CHI/STE]
	$\Delta_{\text{fus}}H$		11.8	391.1	DSC	[2007BON/CAT]
	$\Delta_{\text{sub}}H$		98.3		ME	[1972MIL]
	Δ_vH		77.7 ± 0.1	520	EB	[2010CHI/STE]
	Δ_vH		74.9 ± 0.2	560	EB	[2010CHI/STE]
			72.1 ± 0.2	600	EB	[2010CHI/STE]
C ₁₂ H ₈ N ₂	[230-07-9]	4,7-phenanthroline				
	Δ_vH		80.8 ± 4.7	298	CGC	[2009LIP/CHI, 2009LIP/HAN]
C ₁₂ H ₈ N ₂	[92-82-0]	phenazine				

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{fus}}H$		24.92	447.9	AC	[2010CHI/STE2]
	$\Delta_{\text{fus}}H$		20.92	450.2	DSC	[1975MCE/SAN]
	$\Delta_{\text{sub}}H$		95.9 ± 0.4	298		[2010CHI/STE2]
	$\Delta_{\text{sub}}H$		94.3 ± 0.4	354		[2010CHI/STE2]
	$\Delta_{\text{sub}}H$		92.7 ± 0.4	354		[1991SAB/WAT]
	$\Delta_{\text{sub}}H$		97.0 ± 0.4	298		[1991SAB/WAT]
	$\Delta_{\text{sub}}H$		91.8 ± 2.1	298	C	[1990LEI/PIL]
	$\Delta_{\text{sub}}H$	(280–318)	92.4	295		[1987STE/MAL]
	$\Delta_{\text{sub}}H$		99.9 ± 2.5		ME,GS	[1980ARS]
	$\Delta_{\text{sub}}H$	(303–328)	90.4 ± 2.5	298	TE	[1975DEK/VAN]
	$\Delta_{\text{sub}}H$	(303–323)	90.0 ± 1.5	298	TCM	[UR/DEK, 1975DEK/VAN]
	$\Delta_{\text{sub}}H$	(281–293)	90.4 ± 1.7		LE	[1975MCE/SAN]
	$\Delta_{\text{sub}}H$		U 81.5			[1946ALB/WIL]
	Δ_vH		66.1 ± 0.1	450		[2010CHI/STE2]
	Δ_vH		65.5 ± 0.1	460		[2010CHI/STE2]
	Δ_vH		65.0 ± 0.1	470		[2010CHI/STE2]
C ₁₂ H ₈ N ₂	[230-17-1]	benzo[c]cinnoline				
	$\Delta_{\text{fus}}H$		20.92	432.2	DTA	[1977SCH/PET]
	$\Delta_{\text{sub}}H$	(320–360)	101.7 ± .2	340	ME	[1977SCH/PET, 1987STE/MAL]
	$\Delta_{\text{sub}}H$		113		ME	[1972MIL]
C ₁₂ H ₈ N ₂ O	[304-81-4]	phenazine-N-oxide				
	$\Delta_{\text{sub}}H$		100. ± 1.3	298	C	[1990LEI/PIL]
C ₁₂ H ₈ N ₂ O ₄	[1528-74-1]	4,4'-dinitrobiphenyl				
	$\Delta_{\text{sub}}H$	(441–428)	104.6 ± 1.8	420	ME	[1953SEK/SUZ, 1960JON]
C ₁₂ H ₈ N ₂ O ₅	[101-63-3]	4,4'-dinitrodiphenyl ether				
	$\Delta_{\text{fus}}H$		10.29	418.2		[1993ACR]
C ₁₂ H ₈ N ₄	[1017-93-2]	bicyclo[2.2.2]oct-5-ene-2,2,3,3-tetracarbonitrile				
	$\Delta_{\text{fus}}H$		18.91	476.7		
	$\Delta_{\text{fus}}H$		4.54	533.2	DSC	[1984WEI/LEF]
	$\Delta_{\text{sub}}H$		111.7 ± 5.4	433		[1972ROG2, 1977PED/RYL]
C ₁₂ H ₈ N ₄	[7120-73-2]	dibenzo-1,3a,4,6a-tetraazapentalene				
	$\Delta_{\text{sub}}H$	(363–433)	70.3 ± 1.7	400		[1967CHI/SIM]
C ₁₂ H ₈ N ₄	[2055-55-2]	dibenzo-1,3a,6,6a-tetraazapentalene				
	$\Delta_{\text{sub}}H$	(363–443)	42.3 ± 3.4	403		[1967CHI/SIM]
C ₁₂ H ₈ O	[132-64-9]	dibenzofuran				
	$\Delta_{\text{fus}}H$		18.6	354.7	DSC	[2010KES/AUC]
	$\Delta_{\text{fus}}H$		19.41	355.1		[2007HAF/MAH]
	$\Delta_{\text{fus}}H$		20.5	355.8	DSC	[2000LIS/JAM]
	$\Delta_{\text{fus}}H$		17.6	355.2	DSC	[2000MAH/SOL]
	$\Delta_{\text{fus}}H$		18.6	355.7		[1990CHI/GAM]
	$\Delta_{\text{sub}}H$	(295–318)	82.1 ± 1.5	307	ME	[2004LI/SHI]
	$\Delta_{\text{sub}}H$	(293–353)	82.0 ± 0.2	298	GS	[2003VER2]
	$\Delta_{\text{sub}}H$		84.4 ± 0.7	298		[1990CHI/GAM]
	$\Delta_{\text{sub}}H$		76.5 ± 0.2	298		[1987SAB/ANT]
	$\Delta_{\text{sub}}H$	(304–343)	85.6	324	T	[1989ROR, 1986ROR]
	$\Delta_{\text{sub}}H$	(303–343)	79.1	323		[1986HAN/ECK]
	$\Delta_{\text{sub}}H$	(299–346)	76.7	323	GS	[1986SAT/INO]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{sub}}H$		88.7 ± 2.1			[1958CAS/FLE3]
	Δ_vH	(323–473)	66.2	398	GC	[2002LEI/CHA]
	Δ_vH	(403–559)	55.1	418	A	[1987STE/MAL]
	Δ_vH	(403–418)	66.2	410		[1958CAS/FLE3]
C₁₂H₈OS	[262-20-4]	phenoxathiin				
	$\Delta_{\text{fus}}H$		19.43	329.6	DSC	[2008MON/SAN]
	$\Delta_{\text{fus}}H$		20.27	328.8		[1993STE/CHI]
	Δ_vH	(318–373)	77.3 ± 0.1	298	ME	[2008MON/SAN]
	Δ_vH	(365–640)	68.7	400	EB,IP	[1993STE/CHI]
	Δ_vH	(365–640)	66	440	EB,IP	[1993STE/CHI]
	Δ_vH	(365–640)	63.4	480	EB,IP	[1993STE/CHI]
	Δ_vH	(365–640)	60.8	520	EB,IP	[1993STE/CHI]
	Δ_vH	(365–640)	58	560	EB,IP	[1993STE/CHI]
	Δ_vH	(365–640)	55.1	600	EB,IP	[1993STE/CHI]
C₁₂H₈OS₂	[49833-13-8]	diphenylene-2,2'-disulfide S-oxide				
	$\Delta_{\text{fus}}H$		17.99	407	DSC	[1996DOM/HEA, 1975CUC]
C₁₂H₈O₂	[262-12-4]	dibenzo[b,e][1,4]dioxin				
	$\Delta_{\text{fus}}H$		23.2	395.7		[1986ROR2]
	$\Delta_{\text{sub}}H$	(303–333)	93.6 ± 1.2	318	ME	[2004LI/SHI]
	$\Delta_{\text{sub}}H$		91.5 ± 0.8	298	C	[2002PIM/MEL]
	$\Delta_{\text{sub}}H$		89.6 ± 0.7	298	C	[1999KOL/DOR]
	$\Delta_{\text{sub}}H$		89.6 ± 0.7	318	C	[1997LUK/KOL]
	$\Delta_{\text{sub}}H$	(303–333)	92.3	318	T	[1989ROR, 1986ROR]
C₁₂H₈O₂S	[1016-05-3]	dibenzothiophene sulfone				
	$\Delta_{\text{fus}}H$		27.17	509.2	DSC	[2007RAM/ROJ]
	$\Delta_{\text{fus}}H$		23.72	507.8	DSC	[UR/MCC]
C₁₂H₈S	[132-64-0]	dibenzothiophene				
	$\Delta_{\text{fus}}H$		21.6	371.8	DSC	[2000LIS/JAM]
	$\Delta_{\text{fus}}H$		21.58	371		[1996DOM/HEA]
	$\Delta_{\text{fus}}H$		21.6	373.2		[1983ORO/MRA]
	$\Delta_{\text{sub}}H$		93.2 ± 0.5	298	C	[2009FRE/GOM]
	$\Delta_{\text{sub}}H$		85.1 ± 0.4	298	C	[1987SAB/ANT, 1979SAB]
	$\Delta_{\text{sub}}H$	(303–348)	91.2	325	T	[1986HAN/ECK]
	$\Delta_{\text{sub}}H$	(333–363)	90.7	348	GS	[1981EDW/PRA]
	$\Delta_{\text{sub}}H$		97.5	298		[1975AUB/MAY, 2009FRE/GOM]
	Δ_vH	(413–473)	78.3 ± 1.1	298	GC	[2006HAF/PAR]
	Δ_vH	(373–424)	65.6	388		[1999DYK/SVO]
	Δ_vH	(424–608)	63.4	439		[1999DYK/SVO]
	Δ_vH		69.5 ± 0.3	380		[1995STE/CHI]
	Δ_vH		66.8 ± 0.3	420		[1995STE/CHI]
	Δ_vH		64.3 ± 0.3	460		[1995STE/CHI]
	Δ_vH		61.8 ± 0.3	500		[1995STE/CHI]
	Δ_vH		59.3 ± 0.3	540		[1995STE/CHI]
	Δ_vH		56.8 ± 0.3	580		[1995STE/CHI]
	Δ_vH		54.0 ± 0.3	620		[1995STE/CHI]
	Δ_vH	(375–662)	68.0 ± 0.1	400	EB,IP	[1991CHI/KN1]
	Δ_vH	(375–662)	64.9 ± 0.1	450	EB,IP	[1991CHI/KN1]
	Δ_vH	(375–662)	61.8 ± 0.1	500	EB,IP	[1991CHI/KN1]
	Δ_vH	(375–662)	58.7 ± 0.1	550	EB,IP	[1991CHI/KN1]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(375–662)	55.4 ± 0.3	600	EB,IP	[1991CHI/KN1]
	$\Delta_v H$	(375–662)	51.8 ± 0.4	650	EB,IP	[1991CHI/KN1]
	$\Delta_v H$	(385–574)	60.1	400	A	[1987STE/MAL]
	$\Delta_v H$		56.9	590	C	[1984MRA/KEW]
	$\Delta_v H$		55.3	610	C	[1984MRA/KEW]
	$\Delta_v H$		53.6	630	C	[1984MRA/KEW]
	$\Delta_v H$	(373–403)	69.4	385	GS	[1981EDW/PRA]
C ₁₂ H ₈ S ₂	[92-85-3]	thianthrene				
	$\Delta_{\text{fus}}H$		27.55	429.6		[1993STE/CHI]
	$\Delta_{\text{sub}}H$		103.6 ± 0.4	350	IP	[1993STE/CHI]
	$\Delta_{\text{sub}}H$	(338–368)	98.6 ± 0.5	353		[1989SAB/ELW]
	$\Delta_{\text{sub}}H$		99.4 ± 0.6	298		[1989SAB/ELW]
	$\Delta_{\text{sub}}H$	(358–428)	98.0	393	GS	[1981EDW/PRA]
	$\Delta_{\text{sub}}H$	(338–368)	97.5 ± 6.3	353	HSA	[1979SAN/EPS]
	$\Delta_v H$	(429–460)	71.2	444		[1999DYK/SVO]
	$\Delta_v H$	(460–539)	68.4	475		[1999DYK/SVO]
	$\Delta_v H$	(395–639)	72.7	440	EB,IP	[1993STE/CHI]
	$\Delta_v H$	(395–639)	69.9	480	EB,IP	[1993STE/CHI]
	$\Delta_v H$	(395–639)	67.2	520	EB,IP	[1993STE/CHI]
	$\Delta_v H$	(395–639)	64.5	560	EB,IP	[1993STE/CHI]
	$\Delta_v H$	(395–639)	61.7	600	EB,IP	[1993STE/CHI]
	$\Delta_v H$	(430–593)	69.1	465		[1983SIV/KOB]
	$\Delta_v H$	(430–593)	68.7	515		[1983SIV/KOB]
	$\Delta_v H$	(428–448)	71.1	438	GS	[1981EDW/PRA]
C ₁₂ H ₈ S ₂	[230-26-2]	dibenzo[c,e] [1,2]dithiin				
	$\Delta_{\text{fus}}H$		19.3	386.2	DSC	[1975CUC2]
C ₁₂ H ₉ Br	[92-66-0]	4-bromobiphenyl				
	$\Delta_v H$	(371–583)	62.2	386	A	[1987STE/MAL, 1947STU]
C ₁₂ H ₉ Br	[2051-98-1]	5-bromoacenaphthene				
	$\Delta_{\text{sub}}H$	(295–321)	87.4 ± 2.6		ME	[2008GOL/SUU2]
C ₁₂ H ₉ BrO	[7025-06-1]	5-bromoacenaphthene				
	$\Delta_v H$	(363–473)	63.7	418	GC	[2001WON/LEI]
C ₁₂ H ₉ BrO	[6876-00-2]	3-bromodiphenyl ether				
	$\Delta_v H$	(363–473)	65.4	418	GC	[2001WON/LEI]
C ₁₂ H ₉ BrO	[101-55-3]	4-bromodiphenyl ether				
	$\Delta_v H$	(463–673)	64.6	478	A	[1987STE/MAL]
C ₁₂ H ₉ BrO	[92-03-5]	2-bromo-4-phenylphenol				
	$\Delta_v H$	(373–584)	57.8	388	A	[1987STE/MAL, 1947STU]
C ₁₂ H ₉ Cl	[2051-60-7]	2-chlorobiphenyl				
	$\Delta_{\text{fus}}H$		14.54	304.9		[1991ACR]
	$\Delta_v H$		72.1 ± 2.0	298	CGC	[2001PUR/CHI]
	$\Delta_v H$	(343–393)	64.4	368	GC	[1994FAL/BID]
	$\Delta_v H$	(409–540)	57.8	424	A	[1987STE/MAL]
	$\Delta_v H$	(306–350)	74.5	328	ME	[1983FER/PIA]
	$\Delta_v H$	(410–540)	55.8	424	QM	[1975GEI/DZH]
	$\Delta_v H$	(362–541)	61.1	377	A	[1987STE/MAL, 1947STU]
C ₁₂ H ₉ Cl	[2051-61-8]	3-chlorobiphenyl				

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$		74.3 ± 1.1	298	CGC	[2001PUR/CHI]
	$\Delta_v H$	(343–393)	66.6	368	GC	[1994FAL/BID]
	$\Delta_v H$	(310–359)	66.2	335	ME	[1983FER/PIA]
	$\Delta_v H$	(341–402)	69.2	372	TE	[1983FER/PIA]
	$\Delta_v H$	(452–536)	66.0	494	QM	[1975GEI/DZH]
C₁₂H₉Cl	[2051-62-9]	4-chlorobiphenyl				
	$\Delta_{\text{fus}}H$		13.32	348.6		[1991ACR]
	$\Delta_{\text{sub}}H$	(253–303)	86.0 ± 0.9	278	GS	[1994WAN/SHU]
	$\Delta_{\text{sub}}H$	(306–346)	73.7 ± 0.7	326	TE,ME	[1983FER/PIA]
	$\Delta_v H$		71.6 ± 0.7	298	CGC	[2001PUR/CHI]
	$\Delta_v H$	(343–393)	66.8	368	GC	[1994FAL/BID]
	$\Delta_v H$	(451–536)	65.9	466	A	[1987STE/MAL]
	$\Delta_v H$	(348–409)	67.8	378	TE	[1983FER/PIA]
	$\Delta_v H$	(369–566)	59.0	384	A	[1987STE/MAL, 1947STU]
C₁₂H₉ClN₂	[na]	4-chloroazobenzene				
	$\Delta_{\text{fus}}H$		27.2	361.2		[1988BAU/PER]
C₁₂H₉ClO	[666747-18-8]	2-chloro-3-phenylphenol				
	$\Delta_v H$	(391–591)	65	406	A	[1987STE/MAL, 1947STU]
C₁₂H₉ClO	[85-97-2]	2-chloro-6-phenylphenol				
	$\Delta_v H$	(393–590)	67.6	408	A	[1987STE/MAL, 1947STU]
C₁₂H₉ClO₂S	[80-38-6]	4-chlorophenylbenzenesulfonate				
	$\Delta_{\text{fus}}H$		21.44	332.2	DSC	[1990DON/DRE]
C₁₂H₉Cl₂NO₂S	[92589-22-5]	N-(2,3-dichlorophenyl)benzene sulfonamide				
	$\Delta_{\text{fus}}H$		27.2	387.2	DSC	[2007PER/STR]
C₁₂H₉F₃N₂O₂	[75706-12-6]	5-methyl-N-[4-(trifluoromethyl)phenyl]-4-isoxazolcarboxamide (leflunomide)				
	$\Delta_{\text{fus}}H$		32.43	438.2		[2006VEG/PET]
C₁₂H₉N	[86-74-8]	carbazole				
	$\Delta_{\text{fus}}H$		26.9	518.7	DSC	[2000LIS/JAM]
	$\Delta_{\text{fus}}H$		27.2	516		[1996DOM/HEA]
	$\Delta_{\text{fus}}H$		0.27	420		[1969ROB/SCO]
	$\Delta_{\text{sub}}H$	(346–364)	101.2 ± 1.1	355	ME	[1990JIM/ROU]
	$\Delta_{\text{sub}}H$		103.3 ± 1.1	298	ME	[1990JIM/ROU]
	$\Delta_{\text{sub}}H$		97.7 ± 0.3	298	C	[1987SAB/ANT]
	$\Delta_{\text{sub}}H$		108.8			[1961ZIM/GEI, 1990JIM/ROU]
	$\Delta_{\text{sub}}H$		84.5 ± 0.8			[1970COX/PIL, 1955AIH3]
	$\Delta_v H$		76.2		GC	[1996GOV/RUT]
	$\Delta_v H$		63.3	525		[1983SIV/MAR]
	$\Delta_v H$		61.8	565		[1983SIV/MAR]
	$\Delta_v H$		60.8	605		[1983SIV/MAR]
	$\Delta_v H$	(525–631)	65.7	540	A	[1987STE/MAL, 1923SEN/NEL, 1984BOU/FRI]
	$\Delta_v H$	(517–624)	66	532		[1923MOR/MUR, 1984BOU/FRI]
C₁₂H₉NO	[91-02-1]	2-benzoylpyridine				
	$\Delta_{\text{fus}}H$	(80–340)	20.91	316.5	AC	[2006WAN/TAN]
C₁₂H₉NO	[135-67-1]	10H-phenoxazine				
	$\Delta_{\text{sub}}H$		96.1 ± 0.3	298	C	[1992SAB/ELW2]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₂ H ₉ NS	[92-84-2]	phenothiazine				
	$\Delta_{\text{fus}}H$		28.4	457.2	DSC	[2007GUP/SIN]
	$\Delta_{\text{fus}}H$		25.7	458.4		[1992SAB/ELW2]
	$\Delta_{\text{fus}}H$		26.92	458.2		[1991ACR]
	$\Delta_{\text{sub}}H$		114.5 ± 0.4	298	C	[1992SAB/ELW2]
		(336–395)	86.0	351	A	[1987STE/MAL, 1942NEL/SMI]
C ₁₂ H ₉ N ₃ O ₂	[2491-52-3]	4-nitroazobenzene				
	$\Delta_{\text{sub}}H$		110		GS	[1987SHI/OHK, 1991HOR]
C ₁₂ H ₉ N ₃ O ₂ S	[138564-59-7]	5-methyl-[(2-nitrophenyl)amino]-3-thiophene carbonitrile				
	$\Delta_{\text{fus}}H$ (<i>yellow prism</i>)		27.2	383		
	$\Delta_{\text{fus}}H$ (<i>orange needle</i>)		25.1	388		
	$\Delta_{\text{fus}}H$ (<i>orange prism</i>)		25.5	385.9		
	$\Delta_{\text{fus}}H$ (<i>red prism</i>)		26.0	379.4	DSC	[2000YU/STE]
C ₁₂ H ₉ N ₃ O ₃	[1435-60-5]	4-hydroxy-4'-nitroazobenzene				
	$\Delta_{\text{sub}}H$		140.1		GS	[1987SHI/OHK, 1991HOR]
	$\Delta_{\text{sub}}H$	(417–444)	143.8	430.5	A	[1987STE/MAL]
	$\Delta_{\text{sub}}H$		146 ± 2.5		TE,ME	[1970KOJ]
	$\Delta_{\text{sub}}H$		136.8			[1968TSU/KOJ, 1988BAU/PER]
C ₁₂ H ₉ N ₃ O ₄	[961-68-2]	N-(2,4-dinitrophenyl)-N-phenylamine				
	$\Delta_{\text{sub}}H$	(402–420)	147.6	411		[1987STE/MAL]
	$\Delta_{\text{sub}}H$		149 ± 3.0		TE,ME	[1970KOJ]
	$\Delta_{\text{sub}}H$		131.8			[1968TSU/KOJ, 1988BAU/PER]
C ₁₂ H ₉ N ₃ O ₅	[119-15-3]	N-(2,4-dinitrophenyl)-N-(4-hydroxyphenyl)amine				
	$\Delta_{\text{sub}}H$	(440–470)	155.6 ± 4.2	455	TE,ME	[1970KOJ, 1987STE/MAL]
	$\Delta_{\text{sub}}H$		154			[1968TSU/KOJ, 1988BAU/PER]
C ₁₂ H ₉ N ₄ O ₄	[961-68-2]	N-(4-aminophenyl)-N-(2,4-dinitrophenyl)amine				
	$\Delta_{\text{sub}}H$	(437–460)	156.6	448.5	A	[1987STE/MAL]
	$\Delta_{\text{sub}}H$		154 ± 2.9		TE,ME	[1970KOJ]
	$\Delta_{\text{sub}}H$		139.3			[1968TSU/KOJ, 1988BAU/PER]
C ₁₂ H ₁₀	[83-32-9]	acenaphthene				
	$\Delta_{\text{fus}}H$		21.0	367	DSC	[2008SHA/GUP]
	$\Delta_{\text{fus}}H$		20.5	na	DSC	[2003SHA/KAN]
	$\Delta_{\text{fus}}H$		21.46	366.6		[1996DOM/HEA]
	$\Delta_{\text{sub}}H$		84.6	298	CGC-DSC	[1998CHI/HES]
	$\Delta_{\text{sub}}H$	(313–453)	83.2	383	GS	[1995NAS/LEN]
	$\Delta_{\text{sub}}H$	(293–342)	77.0	318	GS	[1986SAT/INO]
	$\Delta_{\text{sub}}H$	(283–323)	86.8 ± 0.9	303	GS	[1983SON/ZOL]
	$\Delta_{\text{sub}}H$		83.4 ± 1.0	298		[1975OSB/DOU, 1977FIN/MES]
	$\Delta_{\text{sub}}H$		82.4	366	B	[1975OSB/DOU]
	$\Delta_{\text{sub}}H$		84.7 ± 2.7		ME	[1974RAD/KAT]
	$\Delta_{\text{sub}}H$	(290–340)	86.2 ± 0.8		ME	[1965BOY/CHR, 1970COX/PIL]
	$\Delta_{\text{sub}}H$	(291–310)	82.1 ± 0.4	300	V	[1959AIH, 1987STE/MAL]
	$\Delta_{\text{sub}}H$	(258–308)	81.6			[1958HOY/PEP]
	Δ_vH		68.0	298	CGC	[2008ZHA/UNH]
	Δ_vH	(363–423)	70.5 ± 1.1	298	GC	[2006HAF/PAR]
	Δ_vH		61.1	366		[1998RUU/MOK, 2008HAN/NUT]
Δ_vH		66.2	298		[1998RUU/MOK, 2008HAN/NUT]	
Δ_vH	(323–473)	63.9	398	GC	[2002LEI/CHA]	
Δ_vH		66.2	298	CGC	[1998CHI/HES]	

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$		60.6	378		[1995MOK/GUE, 2008HAN/NUT]
	$\Delta_v H$		66.5 ± 1.3	298		[1995MOK/GUE, 2008HAN/NUT]
	$\Delta_v H$	(368–552)	54	403	A	[1987STE/MAL]
	$\Delta_v H$	(368–413)	60.3	383	A	[1987STE/MAL, 1975OSB/DOU, 1984BOU/FRI]
	$\Delta_v H$		61.3	395	I	[1943CRA]
	$\Delta_v H$	(413–561)	54.3	466	I	[1923MOR/MUR]
	$\Delta_v H$	(420–561)	55.4	435		[1923MOR/MUR, 1984BOU/FRI]
C₁₂H₁₀	[92-52-4]	biphenyl				
	$\Delta_{\text{fus}} H$		19.7	342.3	DSC	[2006KHI/DAH]
	$\Delta_{\text{fus}} H$		19.27	344.34	DSC	[2004BEN/KHI]
	$\Delta_{\text{fus}} H$		18.66	341.5		[1996DOM/HEA]
	$\Delta_{\text{sub}} H$		82.9	298	CGC-DSC	[1998CHI/HES]
	$\Delta_{\text{sub}} H$	(313–453)	81.8	383	GS	[1995NAS/LEN]
	$\Delta_{\text{sub}} H$	(283–338)	83.4	311	EM	[1989SAS/NGU]
	$\Delta_{\text{sub}} H$	(303–333)	U 113.3	318		[1989ROR/RUT]
	$\Delta_{\text{sub}} H$		81.5 ± 0.2	298		[1989CHI/KRI]
	$\Delta_{\text{sub}} H$		77.9 ± 0.3	298	C	[1979SAB2]
	$\Delta_{\text{sub}} H$		81.8 ± 0.2	298	C	[1978MON/ROS]
	$\Delta_{\text{sub}} H$	(306–332)	80.4 ± 1.6	319	TSGC	[1975CLA/KNO]
	$\Delta_{\text{sub}} H$	(273–313)	76.0 ± 4.0		HSA	[1975CHI]
	$\Delta_{\text{sub}} H$		83.6 ± 2.5			[1974RAD/KAT]
	$\Delta_{\text{sub}} H$	(298–318)	75.2		ME	[1974PRI/POU]
	$\Delta_{\text{sub}} H$		81.8 ± 0.4	298	C	[1972MOR]
	$\Delta_{\text{sub}} H$	(279–299)	75.8 ± 0.6	289		[1955AIH3]
	$\Delta_{\text{sub}} H$		81.6 ± 2			[1953BRA/CLE2, 1970COX/PIL, 1960JON]
	$\Delta_{\text{sub}} H$	(287–307)	75.1 ± 1.7	297		[1953SEK/SUZ]
	$\Delta_{\text{sub}} H$	(288–314)	81.6 ± 1.7	301		[1953BRA/CLE]
	$\Delta_{\text{sub}} H$	(278–307)	72.8 + 3	302	ME	[1951BRI]
	$\Delta_{\text{sub}} H$		68.6 ± 0.8	292	QF	[1938WOL/WEG]
	$\Delta_v H$		65.0	298	CGC	[2008ZHA/UNH]
	$\Delta_v H$	(323–473)	62.5	298	GC	[2002LEI/CHA]
	$\Delta_v H$		64.5 ± 2.2	298	GS	[2001PUR/CHI]
	$\Delta_v H$		66.2	298	CGC	[1998CHI/HES]
	$\Delta_v H$	(495–688)	51.2	510	DSC	[1996BAC/GRZ]
	$\Delta_v H$	(403–453)	66.0	298	CGC	[1995CHI/HOS]
	$\Delta_v H$	(348–453)	59.6	363	GS	[1989SAK/IWA]
	$\Delta_v H$	(350–578)	64.9	298	EB	[1989CHI/KRI]
	$\Delta_v H$	(350–578)	57.4	400	EB	[1989CHI/KRI]
	$\Delta_v H$	(350–578)	60.3	360	EB	[1989CHI/KRI]
	$\Delta_v H$	(350–578)	50.4	500	EB	[1989CHI/KRI]
	$\Delta_v H$	(333–393)	60.4	363		[1989SAS/NGU]
	$\Delta_v H$	(390–563)	57.3	405	A	[1987STE/MAL]
	$\Delta_v H$	(396–437)	54.9	417	GS	[1980NAS/HWA]
	$\Delta_v H$	(528–766)	48.0	647		[1957GLA/RUL]
	$\Delta_v H$	(342–544)	59.4	357		[1930CUN, 1984BOU/FRI]
C₁₂D₁₀	[1486-01-7]	biphenyl-d ₁₀				
	$\Delta_v H$		64.9	298	CGC	[2008ZHA/UNH]
C₁₂D₁₀	[15067-26-2]	acenaphthene-d ₁₀				
	$\Delta_v H$		67.2	298	CGC	[2008ZHA/UNH]
C₁₂H₁₀CINO₂S	[21226-30-2]	N-(2-chlorophenyl)benzene sulfonamide				
	$\Delta_{\text{fus}} H$		33.5	398.2	DSC	[2007PER/STR]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₂ H ₁₀ ClNO ₂ S	[4750-28-1] $\Delta_{\text{fus}}H$	N-(4-chlorophenyl)benzene sulfonamide				
			25.8	394.6	DSC	[2007PER/STR]
C ₁₂ H ₁₀ ClN ₃ S	[27429-35-2] $\Delta_{\text{fus}}H$	N-2-pyridyl-N'-(2-chlorophenyl) thiourea				
			28.3	429.7	DSC	[2002KEL/SZC]
C ₁₂ H ₁₀ ClN ₃ S	[53385-84-5] $\Delta_{\text{fus}}H$	N-2-pyridyl-N'-(4-chlorophenyl) thiourea				
			34.3	462.2	DSC	[2002SZC/KEL]
C ₁₂ H ₁₀ Cl ₂ N ₂ O ₂ S	[943757-10-6] $\Delta_{\text{fus}}H$	4-amino-N-(2,3-chlorophenyl)benzene sulfonamide				
			40.9	454.3	DSC	[2008PER/STR]
	$\Delta_{\text{sub}}H$	(345–391)	137.5 ± 0.7	368	GS	[2008PER/STR]
	$\Delta_{\text{sub}}H$	(345–391)	141.1 ± 0.7	298	GS	[2008PER/STR]
	Δ_vH		114.3	298	S-F	[2008PER/STR]
C ₁₂ H ₁₀ Cl ₂ N ₂ O ₂ S	[439118-58-8] $\Delta_{\text{fus}}H$	4-amino-N-(2,5-chlorophenyl)benzene sulfonamide				
			41.3	445.9	DSC	[2008PER/STR]
	$\Delta_{\text{sub}}H$	(379–417)	151.3 ± 1.6	398	GS	[2008PER/STR]
	$\Delta_{\text{sub}}H$	(379–417)	155.4 ± 1.6	298	GS	[2008PER/STR]
	Δ_vH		127.8	298	S-F	[2008PER/STR]
C ₁₂ H ₁₀ Cl ₂ N ₂ O ₂ S	[34392-63-7] $\Delta_{\text{fus}}H$	4-amino-N-(3,4-chlorophenyl)benzene sulfonamide				
			51.5	497.9	DSC	[2008PER/STR]
	$\Delta_{\text{sub}}H$	(418–448)	161.4 ± 3.6	433	GS	[2008PER/STR]
	$\Delta_{\text{sub}}H$	(418–448)	167.5 ± 3.6	298	GS	[2008PER/STR]
	Δ_vH		136.7	298	S-F	[2008PER/STR]
C ₁₂ H ₁₀ F ₃ NO ₂	[52840-38-7] $\Delta_{\text{fus}}H$	4-(trifluoromethyl)-7-(N-ethylamino)coumarin				
			30.47	432.5	DSC	[1991ZHA/HUA]
C ₁₂ H ₁₀ F ₃ NO ₂	[53518-14-2] $\Delta_{\text{fus}}H$	4-(trifluoromethyl)-7-(N,N-dimethylamino)coumarin				
			26.25	420.5	DSC	[1991ZHA/HUA]
C ₁₂ H ₁₀ N ₂	[1080-16-6] $\Delta_{\text{sub}}H$	<i>cis</i> -azobenzene				
		(273–323)	92.9	288		[1987STE/MAL]
	$\Delta_{\text{sub}}H$	(298–357)	92.9 ± 0.12	328	ME	[1977SCH/PET]
	$\Delta_{\text{sub}}H$	(303–333)	U 74.9	318	ME	[1950BRI/CAR, 1960JON]
C ₁₂ H ₁₀ N ₂	[17082-12-1] $\Delta_{\text{fus}}H$	<i>trans</i> -azobenzene				
			22.53	341.1		[1996DOM/HEA]
	$\Delta_{\text{fus}}H$		22.65	341.9	DTA	[1977SCH/PET]
	$\Delta_{\text{fus}}H$		22.1		CR	[1977SCH/PET]
	$\Delta_{\text{sub}}H$		94.1 ± 0.8	298	B	[1996STE/CHI2]
	$\Delta_{\text{sub}}H$	(298–302)	93.6 ± 1.9	298	ME	[1992DIA/MIN]
	$\Delta_{\text{sub}}H$	(298–341)	92.1 ± 0.9	319	TE,ME	[1984BOU/OON]
	$\Delta_{\text{sub}}H$	(299–317)	96.9 ± 0.8	308	TE	[1977DEK/VAN]
	$\Delta_{\text{sub}}H$	(299–317)	94.9 ± 0.8	308	ME	[1977DEK/VAN]
	$\Delta_{\text{sub}}H$	(298–347)	93.8 ± 1.2	323	ME	[1977SCH/PET]
	$\Delta_{\text{sub}}H$	(303–333)	U 74.9	318	A	[1950BRI/CAR, 1960JON, 1987STE/MAL]
	Δ_vH	(436–626)	72.8 ± 0.7	298	EB	[1996STE/CHI2]
	Δ_vH	(376–566)	62.3	391	A	[1987STE/MAL, 1947STU]
C ₁₂ H ₁₀ N ₂	[486-84-0] $\Delta_{\text{fus}}H$	1-methyl-9 <i>H</i> -pyrido[3,4- <i>b</i>]indole (harmaline)				
			27.2	509.9	DSC	[1996BUR/DAG]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₂ H ₁₀ N ₂ O	[21650-65-7]	<i>trans</i> -diphenyldiazene N-oxide				
	$\Delta_{\text{fus}}H$		17.93	309.2		[1991ACR]
	$\Delta_{\text{sub}}H$		98.6 ± 0.9	298	C	[1986KIR/ACR]
C ₁₂ H ₁₀ N ₂ O	[na]	4-hydroxyazobenzene				
	$\Delta_{\text{fus}}H$		32.99	425.2		[1988BAU/PER]
C ₁₂ H ₁₀ N ₂ O ₂	[119-75-5]	N-(2-nitrophenyl)-N-phenylamine				
	$\Delta_{\text{sub}}H$	(335–346)	100.9	340.5	A	[1987STE/MAL]
	$\Delta_{\text{sub}}H$		101.9 ± 1.7		TE,ME	[1970KOJ]
	$\Delta_{\text{sub}}H$		108.4			[1968TSU/KOJ, 1988BAU/PER]
C ₁₂ H ₁₀ N ₂ O ₂	[836-30-6]	N-(4-nitrophenyl)-N-phenylamine				
	$\Delta_{\text{sub}}H$	(382–403)	130.6	392.5	A	[1987STE/MAL]
	$\Delta_{\text{sub}}H$		126.2 ± 1.6		TE,ME	[1970KOJ]
	$\Delta_{\text{sub}}H$		120.9			[1968TSU/KOJ, 1988BAU/PER]
C ₁₂ H ₁₀ N ₄	[69155-29-9]	4,5-dimethyl-1,1,1,2,2-tetracyanocyclohex-4-ene				
	$\Delta_{\text{sub}}H$		107.9 ± 4.2	378		[1972ROG2, 1977PED/RYL]
C ₁₂ H ₁₀ N ₄ O ₂	[730-40-5]	4-amino-4'-nitroazobenzene				
	$\Delta_{\text{fus}}H$		31.88	488.2		[1988BAU/PER]
	$\Delta_{\text{sub}}H$	(403–465)	123	434	GS	[1989NIS/AND]
	$\Delta_{\text{sub}}H$		140.1		GS	[1987SHI/OHK, 1991HOR]
	$\Delta_{\text{sub}}H$		127.6		UV	[1984KAR/ROD, 1984KAR/KRU]
	$\Delta_{\text{sub}}H$		136.4		ME	[1980NIG/DEP, 1991HOR]
	$\Delta_{\text{sub}}H$		140.4 ± 1.2		TE,ME	[1970KOJ]
	$\Delta_{\text{sub}}H$		134.3		ME	[1968TSU/KOJ, 1988BAU/PER]
	$\Delta_{\text{sub}}H$	(404–424)	137.7 ± 0.8	414	TE	[1967GRE/JON, 1987STE/MAL]
	$\Delta_{\text{sub}}H$	(404–423)	136.4 ± 5.0	413	ME	[1967GRE/JON, 1966JON/KRA]
C ₁₂ H ₁₀ O	[941-98-0]	1-acetylnaphthalene				
	Δ_vH	(388–569)	65.4	403	A	[1987STE/MAL]
C ₁₂ H ₁₀ O	[93-08-3]	2-acetylnaphthalene				
	$\Delta_{\text{sub}}H$	(295–316)	87.9 ± 0.4	305	V	[1959AIH, 1987STE/MAL]
	Δ_vH	(393–574)	74.1	408	A	[1987STE/MAL]
C ₁₂ H ₁₀ O	[101-84-8]	diphenyl ether				
	$\Delta_{\text{fus}}H$		17.21	300		[1996DOM/HEA]
	$\Delta_{\text{fus}}H$		16.51	300.4	DSC	[1992BAB/WHA]
	$\Delta_{\text{sub}}H$		82 ± 2.1		E	[1958CAS/FLE3, 1970COX/PIL]
	Δ_vH	(353–393)	67.1	298	CGC	[1995CHI/HOS]
	Δ_vH	(477–544)	65.0	298		[1976AMB/ELL]
	Δ_vH	(477–544)	48.2	531		[1976AMB/ELL]
	Δ_vH	(477–544)	53.0	492	GS,EB	[1987STE/MAL, 1976AMB/ELL]
	Δ_vH		66.1 ± 0.4	298	C	[1972MOR, 1965COL/COU]
	Δ_vH	(313–333)	64.2	323	A	[1987STE/MAL, 1948BEN/FRA]
C ₁₂ H ₁₀ O	[90-43-7]	2-hydroxybiphenyl				
	$\Delta_{\text{fus}}H$		15.48	333.7	DSC	[1998VER5]
	$\Delta_{\text{fus}}H$		16.21	330.6		[1973GEI/DZH]
	$\Delta_{\text{sub}}H$	(301–328)	87.6 ± 0.9	314	GS	[1998VER5]
	$\Delta_{\text{sub}}H$		88.5 ± 0.9	298	GS	[1998VER5]
	$\Delta_{\text{sub}}H$	(292–314)	82.9	303		[1987STE/MAL, 1960AIH]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₂ H ₁₀ O	$\Delta_v H$	(434–547)	94.2	449	A	[1987STE/MAL]
	[92-69-3]	4-hydroxybiphenyl				
	$\Delta_{\text{fus}} H$		31.59	443.1	DSC	[1998VER5]
	$\Delta_{\text{sub}} H$	(333–368)	106.6 ± 1.0	351	GS	[1998VER5]
	$\Delta_{\text{sub}} H$		109.8 ± 1.0	298	GS	[1998VER5]
C ₁₂ H ₁₀ O ₂	$\Delta_{\text{sub}} H$	(327–348)	97.0	337.5	A	[1987STE/MAL, 1960AIH]
	$\Delta_v H$	(450–581)	72.3	465	A	[1987STE/MAL]
	[1806-29-7]	2,2'-dihydroxybiphenyl				
	$\Delta_{\text{fus}} H$		25.36	386.7	DSC	[1998VER5]
	$\Delta_{\text{sub}} H$	(334–363)	111.4 ± 1.2	349	GS	[1998VER5]
C ₁₂ H ₁₀ O ₂	$\Delta_{\text{sub}} H$	(334–363)	114.4 ± 1.2	298	GS	[1998VER5]
	$\Delta_v H$	(444–598)	61.7	459	A	[1987STE/MAL]
	[92-88-6]	4,4'-dihydroxybiphenyl				
	$\Delta_{\text{fus}} H$		43.05	560.7	DSC	[1998VER5]
	$\Delta_{\text{sub}} H$	(354–388)	138.6 ± 2.0	371	GS	[1998VER5]
C ₁₂ H ₁₀ O ₂	$\Delta_{\text{sub}} H$	(354–388)	143.0 ± 2.0	298	GS	[1998VER5]
	[713-68-8]	3-phenoxyphenol				
	$\Delta_v H$	(416–494)	69.5	431	A	[1987STE/MAL]
C ₁₂ H ₁₀ O ₂	[830-81-9]	α -naphthyl acetate				
	$\Delta_{\text{fus}} H$		20.21	319.2		[1981BYS]
	$\Delta_{\text{sub}} H$	(286–317)	95.1 ± 0.6	298	GS	[2003VER/ROU]
C ₁₂ H ₁₀ O ₂	[1523-11-1]	β -naphthyl acetate				
	$\Delta_{\text{fus}} H$		20.05	342.2		[1981BYS]
	$\Delta_{\text{sub}} H$	(313–341)	96.3 ± 0.6	298	GS	[2003VER/ROU]
C ₁₂ H ₁₀ O ₂	[86-87-3]	1-naphthaleneacetic acid				
	$\Delta_{\text{fus}} H$		22.26	405.3		[1991ACR]
	$\Delta_{\text{sub}} H$	(343–372)	112.3 ± 0.9	298	GS	[2004ROU/TEM]
C ₁₂ H ₁₀ O ₂	[581-96-4]	2-naphthaleneacetic acid				
	$\Delta_{\text{sub}} H$	(343–372)	124.6 ± 1.0	298	GS	[2004ROU/TEM]
C ₁₂ H ₁₀ O ₂	[na]	2-carbomethoxynaphthalene				
	$\Delta_{\text{fus}} H$		27.1	350.2		[1978DOZ/FUJ]
C ₁₂ H ₁₀ O ₂ S	[127-63-9]	diphenyl sulfone				
	$\Delta_{\text{fus}} H$		21.78	398.2	DSC	[UR/MCC, 2000DEF/VAN]
	$\Delta_{\text{sub}} H$		106.3 ± 2.9			[UR/MAC, 1970COX/PIL]
C ₁₂ H ₁₀ O ₂ S	[2664-63-3]	4,4'-thiodiphenol				
	$\Delta_{\text{fus}} H$		31.04	432.9		[2001LI/HE]
C ₁₂ H ₁₀ O ₄	[106-34-3]	quinhydrone				
	$\Delta_{\text{sub}} H$	(317–334)	89.1	325.5	A	[1987STE/MAL]
C ₁₂ H ₁₀ O ₄ S ₂	[10409-06-0]	diphenyl disulfone				
	$\Delta_{\text{sub}} H$		161.9 ± 3.3		E	[1964MAC/OHA]
	$\Delta_v H$		149.0 ± 2.9	298	E	[1964MAC/OHA]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₂ H ₁₀ S	[139-66-2]	diphenyl sulfide				
	$\Delta_{\text{fus}}H$	(5–440)	13.98	258	AC	[1995STE/CHI]
	$\Delta_{\text{sub}}H$		95. ± 3.0		E	[1962MAC/MAY3, 1970COX/PIL]
	Δ_vH	(369–566)	60.5	384		[1999DYK/SVO]
	Δ_vH	(345–611)	67.3 ± 0.3	360	EB,IP	[1995STE/CHI]
	Δ_vH	(345–611)	64.3 ± 0.3	400	EB,IP	[1995STE/CHI]
	Δ_vH	(345–611)	61.3 ± 0.3	440	EB,IP	[1995STE/CHI]
	Δ_vH	(345–611)	58.3 ± 0.3	480	EB,IP	[1995STE/CHI]
	Δ_vH	(345–611)	55.3 ± 0.3	520	EB,IP	[1995STE/CHI]
Δ_vH	(345–611)	52.0 ± 0.3	560	EB,IP	[1995STE/CHI]	
Δ_vH	(369–566)	58.2	384	A	[1987STE/MAL, 1949KRE/WIE]	
C ₁₂ H ₁₀ S ₂	[882-33-7]	diphenyl disulfide				
	Δ_vH	(405–583)	72.4	420		[1999DYK/SVO]
	Δ_vH		78.7 ± 2.9	298		[1962MAC/MAY3]
C ₁₂ H ₁₁ ClN ₂ O ₂ S	[16803-92-2]	4-amino-N-(4-chlorophenyl)benzene sulfonamide				
	$\Delta_{\text{fus}}H$		37.3	467.9	DSC	[2008PER/STR]
	$\Delta_{\text{sub}}H$	(400–432)	129.2 ± 1.2	416	GS	[2008PER/STR]
	$\Delta_{\text{sub}}H$	(400–432)	134.1 ± 1.2	298	GS	[2008PER/STR]
C ₁₂ H ₁₁ Cl ₂ NO	[23950-58-5]	3,5-dichloro-N-(1,1-dimethyl-2-propynyl)benzamide				
	$\Delta_{\text{fus}}H$		28.68	428.4	DSC	[1990DON/DRE]
C ₁₂ H ₁₁ N	[90-41-5]	2-aminobiphenyl				
	$\Delta_{\text{fus}}H$		13.99	322.3		[1996DOM/HEA]
	Δ_vH	(340–623)	68.6	400	EB,IP	[1991STE/CHI]
	Δ_vH	(340–623)	65.1	440	EB,IP	[1991STE/CHI]
	Δ_vH	(340–623)	61.8	480	EB,IP	[1991STE/CHI]
	Δ_vH	(340–623)	58.5	520	EB,IP	[1991STE/CHI]
	Δ_vH	(340–623)	55.2	560	EB,IP	[1991STE/CHI]
C ₁₂ H ₁₁ N	[na]	diphenylamine				
	$\Delta_{\text{fus}}H$		19.9	326.1	DSC	[2009SUR/TER]
	$\Delta_{\text{fus}}H$		17.86	326.2		[1991ACR]
	$\Delta_{\text{sub}}H$	(303–319)	110.0 ± 1.0	311	GS	[2009SUR/TER]
	$\Delta_{\text{sub}}H$	(303–319)	110	298	GS	[2009SUR/TER]
	$\Delta_{\text{sub}}H$		96.7 ± 2.5		TE,ME	[1970KOJ]
	$\Delta_{\text{sub}}H$		99.2			[1968TSU/KOJ, 1988BAU/PER]
	$\Delta_{\text{sub}}H$	(298–323)	96.7 ± 2.5	310	QF	[1953AIH, 1970COX/PIL]
	Δ_vH	(381–575)	64.1	396	A	[1987STE/MAL, 1947STU]
Δ_vH	(573–673)	54.2	588	A	[1987STE/MAL]	
C ₁₂ H ₁₁ N	[101-82-6]	2-benzylpyridine				
Δ_vH		69.9 ± 2.8	298	CGC	[2009LIP/CHI, 2009LIP/HAN]	
C ₁₂ H ₁₁ NO	[575-36-0]	N-acetyl-1-naphthylamine				
$\Delta_{\text{sub}}H$	(337–360)	94.1	348.5	A	[1987STE/MAL, 1960AIH2]	
C ₁₂ H ₁₁ NO	[86-86-2]	1-naphthaleneacetamide				
$\Delta_{\text{fus}}H$		32.82	455.5	DSC	[1990DON/DRE]	

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₂ H ₁₁ NO ₂	[63-25-2]	1-naphthyl methylcarbamate				
	$\Delta_{\text{fus}}H$		24.51	416.3	DSC	[1990DON/DRE]
C ₁₂ H ₁₁ N ₃	[60-09-3]	4-aminoazobenzene				
	$\Delta_{\text{fus}}H$		21.7	398.2		[1988BAU/PER]
	$\Delta_{\text{sub}}H$		106.3		GS	[1987SHI/OHK, 1991HOR]
	$\Delta_{\text{sub}}H$		109.4			[1984KAR/KRU]
C ₁₂ H ₁₁ N ₃ O ₂ S	[1900019-63-7]	5-nitro-2-thiophenecarboxaldehyde-4-methylphenylhydrazone				
	$\Delta_{\text{fus}}H$ (<i>red greenish plates</i>)		5.23	425		
	$\Delta_{\text{fus}}H$ (<i>orange red prisms</i>)		15.15	429.2		
	$\Delta_{\text{fus}}H$ (<i>black needles</i>)	(356–373)	110.9 ± 1.7	364	ME	[1956MAJ, 1987STE/MAL]
C ₁₂ H ₁₁ N ₃ S	[886-60-2]	N-2-pyridyl-N'-phenylthiourea				
	$\Delta_{\text{fus}}H$		41.0	na		[2002VAL/HER]
C ₁₂ H ₁₁ O ₂ P	[1707-03-5]	P,P-diphenylphosphinic acid				
	$\Delta_{\text{fus}}H$		21.91	466.1	DSC	[2008ZHA/WAN]
C ₁₂ H ₁₂	[571-58-4]	1,4-dimethylnaphthalene				
	$\Delta_{\text{fus}}H$		10.6	279.2	DSC	[2007CHE/KIM]
	$\Delta_{\text{fus}}H$		10.6	279.9		[1991ACR]
C ₁₂ H ₁₂	[571-61-7]	1,5-dimethylnaphthalene				
	$\Delta_{\text{fus}}H$		20	355.2	DSC	[2007CHE/KIM]
	Δ_vH	(323–473)	64.1	398	GC	[2002LEI/CHA]
C ₁₂ H ₁₂	[575-43-9]	1,6-dimethylnaphthalene				
	$\Delta_{\text{fus}}H$		8.5	257	DSC	[2007CHE/KIM]
	Δ_vH	(323–473)	63.6	398	GC	[2002LEI/CHA]
C ₁₂ H ₁₂	[569-41-5]	1,8-dimethylnaphthalene				
	$\Delta_{\text{fus}}H$		18.53	338.2	DSC	[2007CHE/KIM]
	$\Delta_{\text{fus}}H$		15.77	336.3		[1991ACR]
	$\Delta_{\text{sub}}H$	(328–336)	77.9	332	A	[1987STE/MAL]
	$\Delta_{\text{sub}}H$		79.6	336	B	[1975OSB/DOU, 1979COL/JIM2]
	$\Delta_{\text{sub}}H$		82.7 ± 0.3	298	C	[1974MAN3, 1977PED/RYL]
	Δ_vH	(338–413)	62.8	353	A	[1987STE/MAL, 1975OSB/DOU, 1984BOU/FRI]
	Δ_vH	(338–413)	64.8	336	IP	[1977FIN/MES]
	Δ_vH	(338–413)	62.2	360	IP	[1977FIN/MES]
	Δ_vH	(338–413)	60.7	380	IP	[1977FIN/MES]
C ₁₂ H ₁₂	[581-40-8]	2,3-dimethylnaphthalene				
	$\Delta_{\text{fus}}H$		23.97	377.2	DSC	[2007CHE/KIM]
	$\Delta_{\text{fus}}H$		15.9	378		[1991ACR]
	$\Delta_{\text{sub}}H$	(333–373)	82.8	348	A	[1987STE/MAL]
	$\Delta_{\text{sub}}H$	(287–300)	82.2 ± 0.4	294	ME	[1979COL/JIM2]
	$\Delta_{\text{sub}}H$		81.0		B	[1978ARO/STE]
	$\Delta_{\text{sub}}H$	(278–301)	79.9 ± 0.4	290	V	[1959AIH, 1987STE/MAL]
	Δ_vH		60.9 ± 0.7	380		[1988MES/FIN]
	Δ_vH	(378–408)	60.0	393	A	[1987STE/MAL]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₂ H ₁₂	[581-42-0]	2,6-dimethylnaphthalene				
	$\Delta_{\text{fus}}H$		25.3	385.2	DSC	[2007CHE/KIM]
	$\Delta_{\text{fus}}H$		25.06	383.3		[1991ACR]
	$\Delta_{\text{sub}}H$	(350–383)	84.4 ± 1.9	366		[1977FIN/MES, 1975OSB/DOU, 1987STE/MAL]
	$\Delta_{\text{sub}}H$		82.5	383	B	[1975OSB/DOU]
	$\Delta_{\text{sub}}H$	(279–304)	84.1	291	V	[1959AIH, 1987STE/MAL]
	Δ_vH	(384–418)	57.4	383	IP	[1977FIN/MES]
	Δ_vH	(384–418)	56.6	400	IP	[1977FIN/MES]
	Δ_vH	(384–418)	55.7	420	IP	[1977FIN/MES]
		(384–418)	57.3	399	A	[1987STE/MAL, 1975OSB/DOU, 1984BOU/FRI]
C ₁₂ H ₁₂	[582-16-1]	2,7-dimethylnaphthalene				
	$\Delta_{\text{fus}}H$		22.2	370.2	DSC	[2007CHE/KIM]
	$\Delta_{\text{fus}}H$		23.35	368.8		[1991ACR]
	$\Delta_{\text{sub}}H$	(340–369)	83.8 ± 1	345		[1977FIN/MES, 1975OSB/DOU]
	$\Delta_{\text{sub}}H$		83.2	369	B	[1975OSB/DOU]
	$\Delta_{\text{sub}}H$	(333–368)	84.6	348	A	[1987STE/MAL]
	Δ_vH		57.3	400		[1993CHI/KNI]
	Δ_vH		54.8	440		[1993CHI/KNI]
	Δ_vH		52.2	480		[1993CHI/KNI]
	Δ_vH		49.5	520		[1993CHI/KNI]
	Δ_vH		46.6	560		[1993CHI/KNI]
	Δ_vH	(369–398)	59.5	368.8	IP	[1977FIN/MES]
	Δ_vH	(369–398)	58.6	380	IP	[1977FIN/MES]
	Δ_vH	(369–398)	58.1	390	IP	[1977FIN/MES]
Δ_vH	(369–400)	58.5	384	A	[1987STE/MAL, 1975OSB/DOU, 1984BOU/FRI]	
C ₁₂ H ₁₂	[1127-76-0]	1-ethylnaphthalene				
	Δ_vH	(393–565)	57.3	408	A, GS	[1987STE/MAL, 1979MAC/PRA]
C ₁₂ H ₁₂	[939-27-5]	2-ethylnaphthalene				
	Δ_vH	(323–473)	64.7	398	GC	[2002LEI/CHA]
	Δ_vH	(269–398)	69.3	284		[1988SAS/JOS]
	Δ_vH	(286–319)	61.9	301	A	[1987STE/MAL]
		(393–565)	56.7	408	A	[1987STE/MAL]
C ₁₂ H ₁₂ ClN ₅	[na]	2-amino-4-(<i>p</i> -chloranilino)-6-isopropenyl-s-triazine				
	$\Delta_{\text{fus}}H$ (I)		23.85	415.2		
	$\Delta_{\text{fus}}H$ (II)		20.5	403.2		[1986KUN/YUK]
C ₁₂ H ₁₂ N ₂	[530-50-7]	1,1-diphenylhydrazine				
	Δ_vH	(399–596)	68.8	68.8	A	[1987STE/MAL, 1947STU]
C ₁₂ H ₁₂ N ₂	[na]	hydrazobenzene (1,2-diphenylhydrazine)				
	$\Delta_{\text{fus}}H$		17.65	407.2		[1991ACR]
C ₁₂ H ₁₂ N ₂	[1134-35-6]	4,4'-dimethyl-2,2'-bipyridyl				
	$\Delta_{\text{sub}}H$		99.7 ± 2.3	298	C	[1997RIB/MAT4]
C ₁₂ H ₁₂ N ₂	[92-87-5]	benzidine				
	$\Delta_{\text{fus}}H$		19.1	400.2		[1992RAI/GEO]
C ₁₂ H ₁₂ N ₂ O	[101-80-4]	4,4'-diaminodiphenyl oxide				

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{fus}}H$		7.74	465.4		[1996DOM/HEA]
	$\Delta_{\text{sub}}H$		62.8			[1975BAG/AND]
C ₁₂ H ₁₂ N ₂ O ₂	[6953-81-7] $\Delta_{\text{sub}}H$	1-(4-dimethylaminophenyl)-1 <i>H</i> -pyrrole-2,5-dione (350–370)	122.6 ± 0.9		C	[1998KIS/KAS]
C ₁₂ H ₁₂ N ₂ O ₃	[389-08-2] $\Delta_{\text{fus}}H$ $\Delta_{\text{fus}}H$	1-ethyl-1,4-dihydro-7-methyl-4-oxo-1,8-naphthyridine-3-carboxylic acid (nalidixic acid)	35.92 35.92	501.9 501.9	DSC	[2004ROM/BUS2] [1998BUS/ROM]
C ₁₂ H ₁₂ N ₂ O ₃	[94098-94-9] $\Delta_{\text{sub}}H$	3-(methoxycarbonyl)-2-methylquinoxaline-1-oxide	129.2 ± 4.1	298	C	[2009GOM/MON]
C ₁₂ H ₁₂ N ₂ O ₃	[50-06-6] $\Delta_{\text{fus}}H$ (I) $\Delta_{\text{fus}}H$ (II)	5-ethyl-5-phenylpyridine-2,4,6(1 <i>H</i> ,3 <i>H</i> ,5 <i>H</i>)trione (phenobarbital)	28.0 27.9	449 447	DSC DSC	[2010ZEN/GEL] [2010ZEN/GEL]
C ₁₂ H ₁₂ N ₂ O ₄	[13297-18-2] $\Delta_{\text{sub}}H$	3-methyl-2-quinoxalinecarboxylic acid-1,4-dioxide, ethyl ester	133.4 ± 2.1	298	C	[2004RIB/GOM2]
C ₁₂ H ₁₂ O ₄	[29412-62-2] $\Delta_{\text{fus}}H$ $\Delta_{\text{fus}}H$ $\Delta_{\text{sub}}H$ Δ_vH	1,4-dimethylcubane dicarboxylate	38.1 41.0 117.2 ± 3.9 88.5 ± 2.2	438.2 437.8 298 298	DSC DSC Vap+Fus CGC	[2005ROU/DAV] [1996DOM/HEA] [2005ROU/DAV] [2005ROU/DAV]
C ₁₂ H ₁₂ O ₄	[30296-80-1] $\Delta_{\text{fus}}H$ $\Delta_{\text{sub}}H$ Δ_vH	dimethyl 2,6-cuneanedicarboxylate	23.4 106.8 ± 3.0 89.7 ± 2.1	392.7 298 298	DSC Vap+Fus CGC	[2005ROU/DAV] [2005ROU/DAV] [2005ROU/DAV]
C ₁₂ H ₁₂ O ₆	[2672-57-3] $\Delta_{\text{fus}}H$ Δ_vH	1,2,3-benzenetricarboxylic acid, trimethyl ester	32.7 72.5	375.7 468	DSC A, GS	[1993ACR, 1978DOZ/FUJ] [1987STE/MAL, 1962KRA/BER]
C ₁₂ H ₁₂ O ₆	[28904-30-5] Δ_vH Δ_vH	1,2,4-benzenetricarboxylic acid, trimethyl ester	78.5 ± 0.4 61.1	399 458	C A, GS	[1998MAK/KAB] [1987STE/MAL, 1962KRA/BER]
C ₁₂ H ₁₂ O ₆	[2672-58-4] $\Delta_{\text{fus}}H$ $\Delta_{\text{fus}}H$ $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$ Δ_vH	1,3,5-benzenetricarboxylic acid, trimethyl ester	4.6 17.6 115.9 ± 0.4 118.9 ± 0.4 117.5 ± 0.8 75.4	408.2 419.4 359 298 298 458	DSC DSC ME	[1978DOZ/FUJ] [1995JIM/MEN] [1995JIM/MEN] [1967TUR2, 1995JIM/MEN] [1987STE/MAL]
C ₁₂ H ₁₂ S	[16587-33-0] $\Delta_{\text{fus}}H$ Δ_vH Δ_vH Δ_vH Δ_vH	1,2,3,4-tetrahydrodibenzothiophene	32.03 70.3 ± 0.3 67.3 ± 0.3 64.5 ± 0.2 61.8 ± 0.2	275 360 400 440 480		[2004STE/CHI2] [2004STE/CHI2] [2004STE/CHI2] [2004STE/CHI2] [2004STE/CHI2]

Note: Fusion enthalpy may not be reliable—authors reported a mass loss as the sample melted

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound					
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference	
	$\Delta_v H$	(360–600)	59.2 ± 0.3	520	IP,EB	[2004STE/CHI2]	
	$\Delta_v H$	(360–600)	56.5 ± 0.4	560	IP,EB	[2004STE/CHI2]	
	$\Delta_v H$	(360–600)	75.3 ± 0.7	298	IP,EB	[2004STE/CHI2]	
C ₁₂ H ₁₃ ClF ₃ N ₃ O ₄	[33245-39-5] $\Delta_{\text{fus}} H$	N-(2-chloroethyl)-2,6-dinitro-N-propyl-4-(trifluoromethyl)benzeneamine		23.08	318.4	DSC	[1990DON/DRE]
C ₁₂ H ₁₃ Cl ₃ O ₃	[93-79-8] $\Delta_v H$	2,4,5-trichlorophenoxyacetic acid, butyl ester		87.3	475	A	[1987STE/MAL]
C ₁₂ H ₁₃ N	[86-56-6] $\Delta_v H$	N,N-dimethyl-1-aminonaphthalene		66.9 ± 0.2	298	GS	[2007VER/GEO]
C ₁₂ H ₁₃ NO ₂	[na] $\Delta_{\text{fus}} H$	4-methyl-7-dimethylaminocoumarin		23.92	416.1		[1996DOM/HEA]
C ₁₂ H ₁₃ NO ₂ S	[5237-68-4] $\Delta_{\text{fus}} H$	5,6-dihydro-2-methyl-N-phenyl-1,4-oxathin-3-carboxanilide (carboxin)		28.19	365.3	AC	[2004WAN/TAN]
C ₁₂ H ₁₃ NO ₄ S	[5259-88-1] $\Delta_{\text{fus}} H$	2,3-dihydro-6-methyl-5-phenylcarbamoyl-1,4-oxathiin-4,4-dioxide		26.66	401.5	DSC	[1990DON/DRE]
C ₁₂ H ₁₃ N ₃	[53112-28-0] $\Delta_{\text{fus}} H$	4,6-dimethyl-N-phenyl-2-pyrimidinamine		21.23	370.8	AC	[2004SUN/SON]
C ₁₂ H ₁₄ Cl ₂	[79995-39-4] $\Delta_v H$	cyclohexyl-3,4-dichlorobenzene		64.7	398		[1981GUS/KAS]
C ₁₂ H ₁₄ Cl ₂ O ₃	[94-80-4] $\Delta_v H$	2,4-dichlorophenoxyacetic acid, butyl ester		76.3	459	A	[1987STE/MAL, 1999DYK/SVO]
C ₁₂ H ₁₄ Cl ₂ O ₃	[94-79-1] $\Delta_v H$	2,4-dichlorophenoxyacetic acid, sec-butyl ester		74.2	459	A	[1987STE/MAL, 1999DYK/SVO]
C ₁₂ H ₁₄ Cl ₂ O ₄	[74944-83-5] $\Delta_v H$	2,4-dichlorophenoxyacetic acid, 2-ethoxyethyl ester		63.5	458	A	[1987STE/MAL]
C ₁₂ H ₁₄ Cl ₂ O ₄	[36227-43-7] $\Delta_v H$	2,4-dichlorophenoxyacetic acid, 4-hydroxybutyl ester		72.1	458	A	[1987STE/MAL]
C ₁₂ H ₁₄ N ₂ O ₅	[131-89-5] $\Delta_{\text{fus}} H$	2-cyclohexyl-4,6-dinitrophenol		28.03	378.7	DSC	[1969PLA/GLA]
	$\Delta_v H$	(405–565)	88.6	420	A	[1987STE/MAL, 1947STU]	
C ₁₂ H ₁₄ N ₄ O	[2676-59-7] $\Delta_{\text{fus}} H$	3,3',4'4'-tetraaminodiphenyl ether		25.3	402.6		[1990DOM/HEA]
C ₁₂ H ₁₄ N ₄ O ₂ S	[na] $\Delta_{\text{fus}} H$	4-amino-N-[2,6-dimethyl-4-pyrimidinyl]benzene sulfonamide		45.11	515.6		[1982MAR/MIR]
C ₁₂ H ₁₄ N ₄ O ₂ S	[515-64-0] $\Delta_{\text{fus}} H$	2,4-dimethyl-6-sulfamylamidopyrimidine (sulfisomidine)		42.7	523.6	DTA	[1971SUN/EIS]
C ₁₂ H ₁₄ N ₄ O ₂ S	[55-68-1] $\Delta_{\text{fus}} H$	2-(4-aminobenzenesulfonamido)-4,6-dimethylpyrimidinesulfamethazine		39.2	469	DSC	[2003MAR/AVI, 2002MAR/GOM]
	$\Delta_{\text{fus}} H$		31.1	471.6	DTA	[1971SUN/EIS]	
C ₁₂ H ₁₄ O ₂	[946-38-3] $\Delta_v H$	ethyl <i>cis</i> -2-phenylcyclopropanecarboxylate		70.7 ± 0.6	298	C	[1998KOL/PIM]
C ₁₂ H ₁₄ O ₂	[946-39-4] $\Delta_{\text{sub}} H$	ethyl <i>trans</i> -2-phenylcyclopropanecarboxylate		96.9 ± 0.4	298	C	[1998KOL/PIM]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₂ H ₁₄ O ₃	[93-28-7] $\Delta_v H$	1-acetoxy-2-methoxy-4-allylbenzene (eugenol acetate) (374–555)	63.1	389	A	[1987STE/MAL, 1947STU]
C ₁₂ H ₁₄ O ₃	[na] $\Delta_{\text{fus}} H$	4-methyl-1-phenyl-2,6,7-trioxabicyclo[2.2.2]octane	20.9	410.2		[1995RAK/VER2]
C ₁₂ H ₁₄ O ₄	[523-80-8] $\Delta_v H$	4,7-dimethoxy-5-(2-propen-1-yl)-1,3-benzodioxole (apiol) (389–558)	70.6	404	A	[1987STE/MAL]
C ₁₂ H ₁₄ O ₄	[84-66-2] $\Delta_{\text{fus}} H$	diethyl phthalate	17.99	269.9		[1996DOM/HEA]
	$\Delta_v H$		82.1 ± 0.5	298	EB,ME	[2004ROH/RUZ]
	$\Delta_v H$		87.4	298	EB,ME	[2004ROH/RUZ]
	$\Delta_v H$		74.6	426	BG	[1988KAT]
	$\Delta_v H$	(345–453)	77.9	360	A	[1987STE/MAL]
	$\Delta_v H$	(421–570)	59.1	436	A	[1987STE/MAL]
	$\Delta_v H$	(307–333)	86.8	310	GS	[1982GRA/FOS]
	$\Delta_v H$		81.1 ± 0.8	298	GCC	[1980FUC/PEA]
	$\Delta_v H$	(381–567)	65.9	396		[1947STU]
C ₁₂ H ₁₄ O ₄	[636-09-9] $\Delta_{\text{fus}} H$	diethyl terephthalate	24.69	317.2		[1996DOM/HEA]
C ₁₂ H ₁₄ O ₄	[28153-24-4] $\Delta_{\text{sub}} H$	1,1-diacetoxy-1-phenylethane (308–338)	94.4 ± 2.2	318	GS	[1996VER/PEN]
C ₁₂ H ₁₄ O ₅	[20733-94-2] $\Delta_{\text{fus}} H$	methyl 4-hydroxy-3,5-dimethoxycinnamate (methyl sinapate)	29.85	361.8	DSC	[2010PAN/SAR]
C ₁₂ H ₁₅ ClNO ₄ S ₂	[2310-17-0] $\Delta_{\text{fus}} H$	S-6-chloro-2,3-dihydro-2-oxobenzoxazol-3-ylmethyl O,O-diethylphosphorodithioate	30.03	320	DSC	[1990DON/DRE]
C ₁₂ H ₁₅ N	[6247-00-3] $\Delta_v H$	N,N-diallyl aniline (421–513)	54.8	436	A	[1987STE/MAL]
C ₁₂ H ₁₅ NO	[4783-65-7] $\Delta_v H$	1-benzyl-2-piperidone	91.3 ± 1.0	298	C	[2006RIB/CAB]
C ₁₂ H ₁₅ NO	[3612-20-2] $\Delta_v H$	1-benzyl-4-piperidone	78.0 ± 0.8	298	C	[2006RIB/CAB]
C ₁₂ H ₁₅ NO ₂	[na] $\Delta_{\text{fus}} H$	phenylaminoethyl methacrylate	25.47	297.5		[1996DOM/HEA]
C ₁₂ H ₁₅ NO ₂ S	[21406-29-1] $\Delta_{\text{sub}} H$	N-benzoylthiocarbamic O-butyl ester	120.7 ± 1.8	298	C	[2004RIB/SAN2]
C ₁₂ H ₁₅ NO ₃	[1563-66-2] $\Delta_{\text{fus}} H$	2,3-dihydro-2,2-dimethylbenzofuran-7-yl methylcarbamate	30.33	426.2	DSC	[1990DON/DRE]
C ₁₂ H ₁₅ N ₂ O ₃ PS	[13593-03-8] $\Delta_{\text{fus}} H$	O,O-diethyl O-quinoxalin-2-yl phosphothioate	25.4	304.1	DSC	[1990DON/DRE]
C ₁₂ H ₁₅ N ₃ O ₂	[5972-07-6] $\Delta_{\text{sub}} H$	3,6-bis(dimethylamino)phthalimide (400–457)	105	415	A	[1987STE/MAL]
	$\Delta_{\text{sub}} H$		135.3		RG	[1958KLO]
C ₁₂ H ₁₅ N ₃ O ₆	[81-15-2] $\Delta_{\text{sub}} H$	2,4,6-trinitro-1,3-dimethyl-5- <i>tert</i> -butylbenzene (312–348)	100.4	327	A	[1987STE/MAL, 1956SER/VOI]
C ₁₂ H ₁₅ N ₅ O ₄	[na]	9-[(2-acetoxyethoxy)methyl]-2-acetylamino-9H-purine				

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{fus}}H$		42.33	407.2		[1995KRI/VES]
C ₁₂ H ₁₅ N ₅ O ₅	[75128-73-3] $\Delta_{\text{fus}}H$	9-[(2-acetoxyethoxy)methyl]-2-acetylamino-1,9-dihydro-6H-purin-6-one	47.37	477.2		[1995KRI/VES]
C ₁₂ H ₁₅ N ₃ O ₆	[81-15-2] $\Delta_{\text{fus}}H$	5- <i>tert</i> -butyl-2,4,6-trinitro-1,3-dimethylbenzene (musk xylene)	20.79	386.7	DSC	[2004QU/BAI]
C ₁₂ H ₁₆	[827-52-1] $\Delta_{\text{fus}}H$	cyclohexylbenzene	15.3	280.5		[1996DOM/HEA]
	Δ_vH	(344–462)	60.8 ± 0.2	298	MM	[1998MOK/RAU, 2006VER]
	Δ_vH	(283–462)	60.4	298		[1993KAS/MOK]
	Δ_vH	(333–343)	56.4	348		[1990SOH/OKA]
	Δ_vH	(421–513)	51.3	436	A	[1987STE/MAL]
	Δ_vH		59.9 ± 0.3	298	C	[1978MON/ROS]
C ₁₂ H ₁₆	[na] Δ_vH	dicyclohexadiene (377–505)	77.9	329	A	[1987STE/MAL]
C ₁₂ H ₁₆	[2715-29-9] Δ_vH	2,5-diethylstyrene (322–496)	52.2	337	A	[1987STE/MAL, 1947STU]
C ₁₂ H ₁₆	[5676-29-9] Δ_vH	α - <i>tert</i> -butylstyrene (298–318)	53.2 ± 0.1	298	GS	[1999VER/EBE]
C ₁₂ H ₁₆	[2388-14-9] Δ_vH	1-isopropenyl-4-isopropylbenzene (403–479)	50.9	418	A	[1987STE/MAL]
C ₁₂ H ₁₆	[24375-17-5] $\Delta_{\text{fus}}H$	tetraspiro[2.0.2.0.2.0.2.0]dodecane ([4] rotane)	21	394.9	DSC	[1995BEC/RUC]
	$\Delta_{\text{sub}}H$	(298–338)	74.9 ± 0.5		GS	[1995BEC/RUC]
C ₁₂ H ₁₆ Cl ₂ N ₂ O	[555-37-3] $\Delta_{\text{fus}}H$	N-butyl-N'-(3,4-dichlorophenyl)-N-methylurea	27.23	374.3	DSC	[1990DON/DRE]
C ₁₂ H ₁₆ NO ₂	[2631-37-0] $\Delta_{\text{fus}}H$	5-isopropyl- <i>m</i> -tolyl methylcarbamate	23.04	361.3	DSC	[1990DON/DRE]
C ₁₂ H ₁₆ N ₂ OS	[479578-80-8] $\Delta_{\text{fus}}H$	N-[(3-methoxyphenyl)methyl]-N'-2-propenylthiourea	15.43	313	DSC	[2002ABB/WHO]
C ₁₂ H ₁₆ N ₂ O ₂	[315-18-4] $\Delta_{\text{fus}}H$	4-dimethylamino-3,5-xylol methylcarbamate	18.37	361.7	DSC	[1990DON/DRE]
C ₁₂ H ₁₆ N ₂ O ₂	[na] $\Delta_{\text{sub}}H$	N-benzoyl-N',N'-diethylurea	132.2 ± 2.8	298	C	[2000RIB/RIB]
C ₁₂ H ₁₆ N ₂ O ₄	[na] $\Delta_{\text{fus}}H$	2,4-dinitro-1,3-dimethyl-5- <i>tert</i> -butylbenzene	16.68	340.4		[2004QU/BAI]
C ₁₂ H ₁₆ N ₂ O ₄	[90429-36-0] $\Delta_{\text{fus}}H$	pentyl N-(4-nitrophenyl) carbamate	25.98	363.8	DSC	[1993TIE/FRA]
C ₁₂ H ₁₆ N ₂ O ₅	[83-66-9] $\Delta_{\text{sub}}H$	1-methyl-4- <i>tert</i> -butyl-3-methoxy-2,6-dinitrobenzene (293–353)	102.9			[1953SER/VOI, 1960JON]
C ₁₂ H ₁₆ N ₃ O ₃ PS ₂	[2642-71-9] $\Delta_{\text{sub}}H$	azinphos-ethyl (326–420)	86.8	341	A	[1987STE/MAL]
C ₁₂ H ₁₆ N ₃ O ₃ PS ₂	[2642-71-9] $\Delta_{\text{fus}}H$	S-(3,4-dihydro-4-oxobenzod[d]-[1,2,3]-triazin-3-ylmethyl) O,O-diethylphosphorodithioate	25.22	322.2	DSC	[1990DON/DRE]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₂ H ₁₆ N ₃ O ₆ S	[4726-14-1] $\Delta_{\text{fus}}H$	4-methylsulphonyl-2,6-dinitro-N,N-dipropylaniline	28.05	424.3	DSC	[1990DON/DRE]
C ₁₂ H ₁₆ O ₂	[2049-96-9] Δ_vH	pentyl benzoate (395–492)	85.9	410	A	[1987STE/MAL]
C ₁₂ H ₁₆ O ₂	[94-46-2] Δ_vH	isopentyl benzoate (345–535)	51.6	360	A	[1987STE/MAL, 1947STU]
C ₁₂ H ₁₆ O ₂	[94-46-2] Δ_vH	ethyl 2-phenylbutyrate (404–489)	56.0	419	A	[1987STE/MAL]
C ₁₂ H ₁₆ O ₂	[na] $\Delta_{\text{fus}}H$	benzaldehyde 2,2-dimethylpropylene glycol acetal	18.6	307.6		[1995VER/DOG]
C ₁₂ H ₁₆ O ₂	[26311-45-5] $\Delta_{\text{trs}}H$ (liq cryst) $\Delta_{\text{trs}}H$ (liq cryst) $\Delta_{\text{trs}}H$ (liq cryst-to-liq) $\Delta_{\text{sub}}H$	4-pentylbenzoic acid (341–357)	2.6 9.9 1.5 118.2 ± 1.0	252 362 395 298	ME	[1985PRI/PUC] [2004MON/ALM]
C ₁₂ H ₁₆ O ₂	[2243-32-5] $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$	pentamethylbenzoic acid (347–363)	111.5 ± 1.7 113.4 ± 1.8	355 298	ME	[1988COL/JIM] [1988COL/JIM]
C ₁₂ H ₁₆ O ₃	[2050-08-0] Δ_vH	pentyl salicylate (402–540)	66.5	417	A	[1987STE/MAL]
C ₁₂ H ₁₆ O ₃	[87-20-7] Δ_vH	isopentyl salicylate (287–329)	73.0	302	A, ME	[1987STE/MAL, 1955SER/VOI]
C ₁₂ H ₁₆ O ₃	[15872-41-0] $\Delta_{\text{trs}}H$ (liq cryst) $\Delta_{\text{trs}}H$ (liq cryst-to-liq)	4-pentoxybenzoic acid	21.76 2.09	398.2 422.2		[1967HER]
C ₁₂ H ₁₆ O ₃	[63905-22-6] $\Delta_{\text{fus}}H$	(racemic) 3-(2-allylphenoxy)-propane-1,2-diol	27.8	314.9	DSC	[2008BRE/BRE]
C ₁₂ H ₁₆ O ₃	[476169-18-3] $\Delta_{\text{fus}}H$	(S)-3-(2-allylphenoxy)-propane-1,2-diol	28.8	331.2	DSC	[2008BRE/BRE]
C ₁₂ H ₁₆ O ₄	[14174-08-4] $\Delta_{\text{sub}}H$ Δ_vH	benzo-12-crown-4	104.3 ± 2.6 82.7 ± 2.3	298 298	CGC-DSC CGC	[2000NIC/ORF] [2000NIC/ORF]
C ₁₂ H ₁₆ O ₄	[25762-98-5] $\Delta_{\text{trs}}H$ $\Delta_{\text{fus}}H$	2,5-dipropoxy-1,4-benzoquinone	8.6 33.6	357 460.8	DSC	[1996KEE/VAN]
C ₁₂ H ₁₆ O ₆	[na] $\Delta_{\text{fus}}H$	α -phenoxy- α -(d)-glucopyranoside	39.0	429.2		[1996SCH]
C ₁₂ H ₁₇ N	[31252-42-3] Δ_vH	4-benzylpiperidine	74.2 ± 1.0	298	C	[2007RIB/CAB]
C ₁₂ H ₁₇ NO	[91-49-6] Δ_vH	N-butylacetanilide (443–653)	60.2	458	A	[1987STE/MAL]
C ₁₂ H ₁₇ NO	[2431-96-1] Δ_vH	N,N-diethyl-2-phenylacetamide (404–460)	82.8	419	A	[1987STE/MAL, 1969DAV/MAK]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₂ H ₁₇ NO	[52486-76-7] $\Delta_v H$	2-(dimethylamino)-2-methyl-1-phenylpropanone (298–338)	66.7 ± 0.4	298	GS	[1994WEL/VER]
C ₁₂ H ₁₇ NO	[4061-29-4] $\Delta_v H$	2-(diethylamino)-1-phenylethanone (293–338)	71.6 ± 0.9	298	GS	[1994WEL/VER]
C ₁₂ H ₁₇ NO ₂	[91563-76-7] $\Delta_{\text{fus}} H$	1-nitro-2,6-diisopropylbenzene	12.51	301.2	DSC	[2000VER/HEI]
	$\Delta_{\text{sub}} H$	(279–294)	81.0 ± 1.0	286	GS	[2000VER/HEI]
	$\Delta_{\text{sub}} H$	(279–294)	80.6 ± 1.0	298	GS	[2000VER/HEI]
	$\Delta_v H$	(308–343)	66.9 ± 0.6	326	GS	[2000VER/HEI]
C ₁₂ H ₁₇ NO ₂	[13110-37-7] $\Delta_{\text{fus}} H$	pentyl 4-aminobenzoate	23.93	325.1		[1991ACR]
C ₁₂ H ₁₇ NO ₂ S ₂	[949171-67-9] $\Delta_{\text{fus}} H$	N-theonylthiocarbamic-O-hexyl ester	22.48	346.4	DSC	[2007RIB/MON]
	$\Delta_{\text{sub}} H$		180.1 ± 3.0	298	C	[2007RIB/MON]
C ₁₂ H ₁₇ N ₃ O ₃	[105910-97-2] $\Delta_{\text{fus}} H$	1-pentyl-3-(4-nitrophenyl) urea	19.85	404.2	DSC	[1993TIE/FRA]
C ₁₂ H ₁₇ N ₃ S	[na] $\Delta_{\text{sub}} H$	N-(diethylaminothiocarbonyl)benzamideine	126.0 ± 1.5	298	C	[2004RIB/SAN]
C ₁₂ H ₁₈	[4904-61-4] $\Delta_v H$	1- <i>cis</i> -5- <i>trans</i> -9- <i>trans</i> -cyclododecatriene (344–387)	49.9	359	A	[1987STE/MAL]
	$\Delta_v H$	(400–423)	60.0	411	A	[1987STE/MAL]
	$\Delta_v H$	(426–503)	47.8	441	A	[1987STE/MAL]
C ₁₂ H ₁₈	[706-31-0] $\Delta_v H$	1- <i>trans</i> -5- <i>trans</i> -9- <i>cis</i> -cyclododecatriene (286–373)	68.0	301	A	[1987STE/MAL]
C ₁₂ H ₁₈	[1077-16-3] $\Delta_v H$	hexylbenzene	60.2	298		[1994RUZ/ZAB]
	$\Delta_v H$	(274–463)	61.6	289		[1993KAS/MOK]
	$\Delta_v H$		60.0	298		[1971WIL/ZWO]
C ₁₂ H ₁₈	[577-55-9] $\Delta_v H$	1,2-diisopropylbenzene (388–476)	48.9	403	A	[1987STE/MAL]
C ₁₂ H ₁₈	[99-62-7] $\Delta_v H$	1,3-diisopropylbenzene (283–318)	56.0 ± 0.8	301	GS	[1998VER7]
	$\Delta_v H$	(283–318)	56.2 ± 0.8	298	GS	[1998VER7]
	$\Delta_v H$	(387–477)	48.9	402	A	[1987STE/MAL]
C ₁₂ H ₁₈	[100-18-5] $\Delta_v H$	1,4-diisopropylbenzene (366–530)	50.7 ± 0.2	400	EB	[2002STE/CHI6]
	$\Delta_v H$	(366–530)	46.3 ± 0.3	440	EB	[2002STE/CHI6]
	$\Delta_v H$	(366–530)	43.0 ± 0.5	480	EB	[2002STE/CHI6]
	$\Delta_v H$	(366–530)	39.3 ± 0.9	520	EB	[2002STE/CHI6]
	$\Delta_v H$	(283–318)	56.3 ± 0.3	301	GS	[1998VER7]
	$\Delta_v H$	(283–318)	56.5 ± 0.3	298	GS	[1998VER7]
	$\Delta_v H$	(393–485)	47.6	408	A	[1987STE/MAL]
	$\Delta_v H$	(393–485)	48.9	408		[1959MCD/SHR, 1984BOU/FRI]
C ₁₂ H ₁₈	[98-19-1]	1,3-dimethyl-5- <i>tert</i> -butylbenzene				

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(284–318)	56.5 ± 0.6	301	GS	[1998VER]
	$\Delta_v H$		56.6 ± 0.6	298		[1998VER]
	$\Delta_v H$	(253–443)	59.8	268		[1993KAS/MOK]
C₁₂H₁₈	[87-85-4]	hexamethylbenzene				
	$\Delta_{\text{fus}} H$		1.76	383.7		
	$\Delta_{\text{fus}} H$		20.63	438.7		[1996DOM/HEA, 1932SPA/THO]
	$\Delta_{\text{sub}} H$		80		TGA	[1997GIL/BOT]
	$\Delta_{\text{sub}} H$		81.4 ± 0.1	298	C	[1994SAB/TAB]
	$\Delta_{\text{sub}} H$	(288–304)	85.0 ± 0.2	298	ME	[1989COL/JIM]
	$\Delta_{\text{sub}} H$		74.9 ± 0.6		DSC	[1984HOL]
	$\Delta_{\text{sub}} H$	(303–338)	85.2	320	A	[1976AMB/LAW]
	$\Delta_{\text{sub}} H$		86.1	298	H	[1976AMB/LAW, 1993CHI/HOS]
	$\Delta_{\text{sub}} H$	(314–364)	83.2	329	A	[1969OVE/STE]
	$\Delta_{\text{sub}} H$		74.7 ± 2		ME	[1965FRA/AST, 1970COX/PIL]
	$\Delta_{\text{sub}} H$		80.8			[1957VAN, 1960JON]
	$\Delta_{\text{sub}} H$		80.8			[1949NIT/SEK]
	$\Delta_v H$		68.6	298	CGC	[2008ZHA/UNH]
	$\Delta_v H$	(443–537)	56.8	458	A	[1987STE/MAL]
C₁₂D₁₈	[4342-40-9]	hexamethylbenzene-d ₁₈				
	$\Delta_v H$		68.2	298	CGC	[2008ZHA/UNH]
(C₁₂H₁₈) -(C₆H₃N₂ClO₄)	[57230-36-1]	(hexamethylbenzene)- (picryl chloride)				
	$\Delta_{\text{sub}} H$		93.7			[1949NIT/SEK]
C₁₂H₁₈	[877-44-1]	1,2,4-triethylbenzene				
	$\Delta_v H$	(319–491)	51.2	334	A	[1987STE/MAL, 1947STU]
C₁₂H₁₈	[102-25-0]	1,3,5-triethylbenzene				
	$\Delta_v H$	(371–534)	59.2 ± 0.3	298	EB	[1997STE/CHI2]
C₁₂H₁₈	[10222-95-4]	1,2,4-trimethyl-5-isopropylbenzene				
	$\Delta_v H$		64.9	298		[1975VIL/PER]
C₁₂H₁₈	[6902-73-4]	2-isopropenyl-1-methyl-1-vinyl-3-cyclohexane				
	$\Delta_v H$	(348–404)	47.8	363	A	[1987STE/MAL]
C₁₂H₁₈	[676-22-2]	<i>E,E,E</i> -1,5,9-cyclododecatriene				
	$\Delta_{\text{sub}} H$	(273–307)	75.2	288	A	[1987STE/MAL]
	$\Delta_{\text{sub}} H$		74.7 ± 0.8			[1973RAU/GEY, 1977PED/RYL]
C₁₂H₁₈ClNO	[41570-61-0]	2-chloro- α -[[1,1-dimethylethyl]amino]methyl]benzenemethanol (tulobuterol)				
	$\Delta_{\text{fus}} H$ (I)		27.1	364		
	$\Delta_{\text{fus}} H$ (II)		25.4	354	DSC	[2004CAI/BOU]
C₁₂H₁₈Cl₂NO₃	[42585-08-0]					
	$\Delta_v H$	(309–363)	62.6	324	A	[1987STE/MAL]
C₁₂H₁₈N₂O	[34123-59-6]	<i>N,N</i> -dimethyl- <i>N'</i> -[4-(1-methylethyl)phenyl]urea				
	$\Delta_{\text{fus}} H$		33.87	430.4	DSC	[1991ACR]
C₁₂H₁₈N₂O	[34123-59-6]	<i>N'</i> -(<i>p</i> -cumenyl)- <i>N,N</i> -dimethylurea (isoprotruron)				
	$\Delta_{\text{fus}} H$		21.33	427.4		[2003YU/TAN2]
C₁₂H₁₈N₂O₂	[315-18-4]	3,5-dimethyl-4-(dimethylamino)phenyl methylcarbamate				
	$\Delta_{\text{fus}} H$		18.37	361.7	DSC	[1991ACR, 1990DON/DRE]
C₁₂H₁₈N₂O₂S₂	[120563-92-0]	<i>N</i> -isopropyl- <i>S</i> -methyl- <i>N'</i> -tosylisothiourea				

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{fus}}H$		32.7	392.2	DSC	[1992REI/HAN]
C ₁₂ H ₁₈ N ₂ O ₂ S ₂	[145198-68-1] $\Delta_{\text{fus}}H$	N-ethyl-S-ethyl-N'-tosylisothiourea	30.3	390.2	DSC	[1992REI/HAN]
C ₁₂ H ₁₈ N ₂ O ₃	[76-73-3] $\Delta_{\text{fus}}H$	5-(1-methylbutyl)-5-(2-propen-1-yl)-2,4,6(1 <i>H</i> ,3 <i>H</i> ,5 <i>H</i>)-pyrimidinetrione (secobarbital)	22.9	371.8	DSC	[2008WAS/HOL]
C ₁₂ H ₁₈ N ₂ O ₃ S	[64-77-7] $\Delta_{\text{fus}}H$ $\Delta_{\text{fus}}H$	3-(p-tolyl-4-sulfonyl)-1-butyl urea (tolbutamide)	27.2 25.61	400.2 404.8	DSC	[1999KIM/HIR] [1982MAR/MIR]
C ₁₂ H ₁₈ N ₄ O ₂	[35873-41-7] $\Delta_{\text{fus}}H$	8-pentyltheophylline	35.1	498.4		[1991ACR]
C ₁₂ H ₁₈ N ₄ O ₆ S	[19044-88-3] $\Delta_{\text{fus}}H$	4-(N,N-dipropylamino)-3,5-dinitrobenzenesulphonamide	38.48	414.8	DSC	[1990DON/DRE]
C ₁₂ H ₁₈ O	[4157-77-1] Δ_vH	(1-butoxyethyl)benzene (278–318)	59.8 ± 0.3	298	GS	[2001VER/HEI]
C ₁₂ H ₁₈ O	[6857-85-1] Δ_vH	R,S (1-sec-butoxyethyl)benzene (296–332)	58.7 ± 0.5	298	GS	[2002KRA/VAS, 2002VER/HEI]
C ₁₂ H ₁₈ O	[8760-63-8] Δ_vH	S,S (1-sec-butoxyethyl)benzene (297–332)	59.1 ± 0.5	298	GS	[2002KRA/VAS, 2002BAE/SHI2]
C ₁₂ H ₁₈ O	[24142-77-6] Δ_vH Δ_vH	propyl cumyl ether (278–325) (278–325)	59.1 ± 0.2 59.3 ± 0.2	302 298	GS GS	[2001VER/HEI2] [2001VER/HEI2]
C ₁₂ H ₁₈ O	[6382-14-5] Δ_vH	benzyl pentyl ether (363–512)	50.8	378	A	[1987STE/MAL, 1969KRO]
C ₁₂ H ₁₈ O	[2934-05-6] Δ_vH	2,4-diisopropylphenol (395–528)	58.4	410	A	[1987STE/MAL]
C ₁₂ H ₁₈ O	[2078-54-8] $\Delta_{\text{fus}}H$ Δ_vH Δ_vH	2,6-diisopropylphenol (293–328)	14.64 67.9 ± 0.3 68.7 ± 0.3	292.8 310 298	 GS	[1975BER/PER] [1999VER] [1999VER]
C ₁₂ H ₁₈ O	[26886-05-5] $\Delta_{\text{fus}}H$	3,5-diisopropylphenol	12.13	326.3		[1975BER/PER]
C ₁₂ H ₁₈ O	[68189-19-5] Δ_vH	2,3-dimethyl-4- <i>tert</i> -butylphenol (418–523)	60.2	433	A	[1987STE/MAL]
C ₁₂ H ₁₈ O	[46170-85-8] Δ_vH	2,3-dimethyl-6- <i>tert</i> -butylphenol (412–525)	60.0	427	S	[1987STE/MAL]
C ₁₂ H ₁₈ O	[1879-09-0] Δ_vH Δ_vH Δ_vH Δ_vH Δ_vH Δ_vH Δ_vH	2,4-dimethyl-6- <i>tert</i> -butylphenol (304–333) (388–522) (344–535) (344–535) (344–535) (344–535) (344–535) (344–535)	67.2 ± 0.8 68.4 ± 0.8 58.4 54.4 52.7 51.7 49.7 45.4	318 298 403 348 373 398 423 473	GS A	[1999VER] [1999VER] [1987STE/MAL] [1953STA/MUL] [1953STA/MUL] [1953STA/MUL] [1953STA/MUL] [1953STA/MUL]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₂ H ₁₈ O	[17696-37-6]	2,5-dimethyl-4- <i>tert</i> -butylphenol				
	$\Delta_v H$	(408–538)	61.7	423	A	[1987STE/MAL]
	$\Delta_v H$	(361–548)	62.0	373		[1953STA/MUL]
	$\Delta_v H$	(361–548)	59.4	398		[1953STA/MUL]
	$\Delta_v H$	(361–548)	57.1	423		[1953STA/MUL]
$\Delta_v H$	(361–548)	52.8	473		[1953STA/MUL]	
C ₁₂ H ₁₈ O	[879-97-0]	2,6-dimethyl-4- <i>tert</i> -butylphenol				
	$\Delta_v H$	(392–522)	59.7	407	A	[1987STE/MAL]
	$\Delta_v H$	(347–530)	58.4	348		[1953STA/MUL]
	$\Delta_v H$	(347–530)	57.0	373		[1953STA/MUL]
	$\Delta_v H$	(347–530)	55.4	398		[1953STA/MUL]
	$\Delta_v H$	(347–530)	54.2	423		[1953STA/MUL]
$\Delta_v H$	(347–530)	49.3	473		[1953STA/MUL]	
C ₁₂ H ₁₈ O	[1445-23-4]	3,4-dimethyl-6- <i>tert</i> -butylphenol				
$\Delta_v H$	(413–532)	62.7	428	A	[1987STE/MAL]	
C ₁₂ H ₁₈ O	[63452-61-9]	2-ethyl-4- <i>tert</i> -butylphenol				
	$\Delta_v H$	(428–623)	61.6	443	A	[1987STE/MAL]
	$\Delta_v H$	(397–543)	55.4	398		[1953STA/MUL]
	$\Delta_v H$	(397–543)	54.2	423		[1953STA/MUL]
$\Delta_v H$	(397–543)	49.3	473		[1953STA/MUL]	
C ₁₂ H ₁₈ O	[63551-41-7]	2-ethyl-6- <i>tert</i> -butylphenol				
$\Delta_v H$	(393–443)	58.2	408	A	[1987STE/MAL]	
C ₁₂ H ₁₈ O	[4237-25-6]	3-ethyl-6- <i>tert</i> -butylphenol				
$\Delta_v H$	(415–530)	59.5	430	A	[1987STE/MAL]	
C ₁₂ H ₁₈ O	[96-70-8]	4-ethyl-2- <i>tert</i> -butylphenol				
	$\Delta_v H$	(394–523)	59.2	409	A	[1987STE/MAL]
	$\Delta_v H$	(349–533)	57.0	373		[1953STA/MUL]
	$\Delta_v H$	(349–533)	55.4	398		[1953STA/MUL]
	$\Delta_v H$	(349–533)	54.2	423		[1953STA/MUL]
$\Delta_v H$	(349–533)	49.3	473		[1953STA/MUL]	
C ₁₂ H ₁₈ O	[71745-63-6]	2-methyl-4- <i>tert</i> -pentylphenol				
	$\Delta_v H$	(443–653)	65.6	458	A	[1987STE/MAL]
	$\Delta_v H$	(409–561)	55.3	423		[1953STA/MUL]
$\Delta_v H$	(409–561)	50.7	473		[1953STA/MUL]	
C ₁₂ H ₁₈ O	[na]	3-methyl-4- <i>tert</i> -pentylphenol				
	$\Delta_v H$	(443–683)	65.1	458	A	[1987STE/MAL]
	$\Delta_v H$	(409–561)	55.3	423		[1953STA/MUL]
	$\Delta_v H$	(409–561)	50.7	473		[1953STA/MUL]
C ₁₂ H ₁₈ O	[34072-71-4]	4-methyl-2- <i>tert</i> -pentylphenol				
	$\Delta_v H$	(423–653)	61.4	438	A	[1987STE/MAL]
	$\Delta_v H$	(394–538)	58.1	398		[1953STA/MUL]
	$\Delta_v H$	(394–538)	55.3	423		[1953STA/MUL]
$\Delta_v H$	(394–538)	50.7	473		[1953STA/MUL]	
C ₁₂ H ₁₈ O	[1660-04-4]	1-adamantyl methyl ketone				
$\Delta_{\text{sub}} H$	(287–305)	84.2 ± 0.6	298	ME	[1992ABB/JIM2]	
C ₁₂ H ₁₈ O	[7273-98-5]	<i>exo</i> -4-hydroxy- <i>endo</i> - <i>endo</i> -tetracyclo[6.2.1.1. ^{3,6} .0 ^{2,7}]dodecane				
$\Delta_{\text{sub}} H$	(303–343)	79.0 ± 2.5	298	TSGC	[1980STE]	
C ₁₂ H ₁₈ O	[107133-43-7]	<i>exo</i> -4-hydroxy- <i>exo</i> - <i>endo</i> -tetracyclo[6.2.1.1. ^{3,6} .0 ^{2,7}]dodecane				

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{sub}}H$	(323–353)	74.3 ± 1.8		TSGC	[1980STE]
	$\Delta_{\text{sub}}H$		76.3 ± 2.0	298		[1980STE]
C ₁₂ H ₁₈ O	[74007-11-7]	<i>exo</i> -4-hydroxy- <i>exo</i> - <i>exo</i> -tetracyclo[6.2.1.1. ^{3,6} .0 ^{2,7}]dodecane				
	$\Delta_{\text{sub}}H$	(313–353)	73.9 ± 2		TSGC	[1980STE]
	$\Delta_{\text{sub}}H$		75.9 ± 2.2	298		[1980STE]
C ₁₂ H ₁₈ O	[na]	2-(1'-cyclohexenyl)cyclohexanone				
	$\Delta_{\text{fus}}H$		17.26	278.8		[1992MAR/KOZ]
C ₁₂ H ₁₈ O ₂	[5673-09-6]	1,3-dihydroxy-2-hexylbenzene				
	Δ_vH	(433–494)	76.8	448	A, GC	[1987STE/MAL, 1975KUN/LIL]
C ₁₂ H ₁₈ O ₂	[136-77-6]	1,3-dihydroxy-4-hexylbenzene				
	$\Delta_{\text{fus}}H$		19.04	341.5		[1991ACR]
	Δ_vH	(434–494)	88.1	449	A, GC	[1987STE/MAL, 1975KUN/LIL]
C ₁₂ H ₁₈ O ₂	[711-01-3]	1-adamantyl-1-carboxylic acid methyl ester				
	$\Delta_{\text{sub}}H$	(267–283)	84.3 ± 0.6	275	ME	[1992ABB/JIM]
	$\Delta_{\text{sub}}H$		82.4 ± 0.6	298		[1992ABB/JIM]
C ₁₂ H ₁₈ O ₂	[na]	<i>trans</i> - <i>syn</i> - <i>trans</i> decahydro-3-hydroxy-2-naphthalene acetic γ -lactone				
	$\Delta_{\text{sub}}H$	(240–310)	NA		ME	[1957SPI]
C ₁₂ H ₁₈ O ₂	[na]	<i>trans</i> - <i>anti</i> - <i>trans</i> decahydro-3-hydroxy-2-naphthalene acetic γ -lactone				
	$\Delta_{\text{sub}}H$	(240–310)	NA		ME	[1957SPI]
C ₁₂ H ₁₈ O ₃	[63911-78-6]	(racemic) 3-(2-propylphenoxy)-propane-1,2-diol				
	$\Delta_{\text{fus}}H$		29.5	326.5	DSC	[2008BRE/BRE]
C ₁₂ H ₁₈ O ₃	[1092799-99-9]	(R)-3-(2-propylphenoxy)-propane-1,2-diol				
	$\Delta_{\text{fus}}H$		31.9	340.5	DSC	[2008BRE/BRE]
C ₁₂ H ₁₈ O ₃	[204583-98-2]	(racemic) 3-(2-isopropylphenoxy)-propane-1,2-diol				
	$\Delta_{\text{fus}}H$		31.5	353.7	DSC	[2008BRE/BRE]
C ₁₂ H ₁₈ O ₃	[204584-38-3]	(R)-3-(2-isopropylphenoxy)-propane-1,2-diol				
	$\Delta_{\text{fus}}H$		30.6	345	DSC	[2008BRE/BRE]
C ₁₂ H ₁₈ O ₄	[532-34-3]	3,4-dihydro-2,2-dimethyl-4-oxo-2H-pyran-6-carboxylic acid, butyl ester				
	Δ_vH	(357–435)	64.7	372	A	[1987STE/MAL]
C ₁₂ H ₁₈ O ₆	[na]	triethyl aconitrate				
	Δ_vH	(423–540)	79.6	438	A	[1987STE/MAL]
C ₁₂ H ₁₈ O ₆	[na]	R,R,R-4,8,12-trimethyl-1,5,9-trioxacyclododeca-2,6,10-trione				
	$\Delta_{\text{fus}}H$		21.51	380.2		[1996LEB/BYK]
C ₁₂ H ₁₉ ClNO ₃ P	[299-86-5]	N-methyl O-methyl O-2-chloro-4- <i>tert</i> -butylphenylphosphoramidate				
	$\Delta_{\text{fus}}H$		21.98	332	DSC	[1990DON/DRE]
C ₁₂ H ₁₉ F ₃ N ₂ O ₄	[na]	N-[(N-trifluoroacetyl)valyl]alanine ethyl ester				
	$\Delta_{\text{sub}}H$	(323–424)	115.5	338	A	[1987STE/MAL, 1960WEY/KLI]
C ₁₂ H ₁₉ N	[24544-04-5]	2,6-diisopropylaniline				
	Δ_vH	(284–323)	69.2 ± 0.3	303		[2000VER3]
	Δ_vH		69.5 ± 0.3	298		[2000VER3]
C ₁₂ H ₁₉ N	[202925-84-6]	N-methyl-3-methyl-3-phenyl-2-butaneamine				
	Δ_vH	(283–330)	67.0 ± 0.8	307	GS	[1998VER/BEC]
	Δ_vH	(283–330)	67.5 ± 0.8	298	GS	[1998VER/BEC]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₂ H ₁₉ F ₃ N ₂ O ₄	[na] $\Delta_v H$	N-[N-(trifluoroacetyl)valyl]alanine ethyl ester (425–453) 86.4 439 A [1987STE/MAL, 1999DYK/SVO]				
C ₁₂ H ₂₀	[770-69-4] $\Delta_{\text{fus}} H$	1-ethyladamantane (8–373) 11.28 225.6 AC [2005VAR/DRU]				
	$\Delta_v H$		55.3 ± 1.1	298		[2000MEL/PIM]
	$\Delta_v H$	(383–492)	49.1	398	A	[1987STE/MAL]
C ₁₂ H ₂₀	[707-79-4] $\Delta_{\text{fus}} H$	1,3-dimethyladamantane (8–373) 9.31 223.4				
	$\Delta_{\text{fus}} H$	(8–373)	1.54	247.8	AC	[2005VAR/DRU]
	$\Delta_{\text{fus}} H$		7.65	221		
	$\Delta_{\text{fus}} H$		0.94	244	DSC	[1980ARN/SCH]
	$\Delta_{\text{fus}} H$		7.36	221		
	$\Delta_{\text{fus}} H$		0.92	245		[1977CLA/KNO]
	$\Delta_{\text{sub}} H$		67.8 ± 1.3	298	EB	[1977STE/WAT]
	$\Delta_v H$		49.2 ± 0.2	308	C	[2001VAN/PAS]
	$\Delta_v H$		49.7 ± 0.2	298	C	[2001VAN/PAS]
	$\Delta_v H$	(352–526)	49.4 ± 0.3	298	EB	[1996STE/CHI]
	$\Delta_v H$	(352–526)	45.9 ± 0.3	360	EB	[1996STE/CHI]
	$\Delta_v H$	(352–526)	43.7 ± 0.3	400	EB	[1996STE/CHI]
	$\Delta_v H$	(352–526)	41.5 ± 0.3	440	EB	[1996STE/CHI]
$\Delta_v H$	(352–526)	39.1 ± 0.3	480	EB	[1996STE/CHI]	
$\Delta_v H$	(352–526)	36.4 ± 0.3	520	EB	[1996STE/CHI]	
C ₁₂ H ₂₀	[19740-34-2] $\Delta_{\text{sub}} H$	2,2-dimethyladamantane (300–360) 73.6 ± 1.3 298 BG [1977STE/WAT]				
C ₁₂ H ₂₀ N ₂	[3867-15-0] $\Delta_{\text{fus}} H$	1-(1-piperidinyl)cyclohexanecarbonitrile 25.44 339.2 [1997WEL/VER]				
	$\Delta_{\text{sub}} H$		87.8 ± 0.6	298		[1997WEL/VER]
C ₁₂ H ₂₀ N ₂	[4543-66-2] $\Delta_{\text{fus}} H$	dodecanedinitrile 34.33 294.2 DSC [2007BAD/BLA]				
C ₁₂ H ₂₀ N ₂ O ₂	[6310-76-5] $\Delta_{\text{sub}} H$	N,N'-ethylenebis(4-aminopent-3-ene-2-one) (358–374) 128.2 ± 0.7 366 ME [1995RIB/RIB]				
	$\Delta_{\text{sub}} H$	(358–374)	131.6	298	ME	[1995RIB/RIB]
C ₁₂ H ₂₀ N ₂ O ₂	[63254-50-2] $\Delta_{\text{fus}} H$	(1R,2S,5R)-2-isopropyl-5-methylcyclohexyl diazoacetate 17.2 320.4 [2000DI/TAN2]				
C ₁₂ H ₂₀ N ₄ O ₂	[51235-04-2] $\Delta_{\text{fus}} H$	3-cyclohexyl-6-(dimethylamino)-1-methyl-1,3,5-triazine-2,4(1H,3H)-dione 20.36 389.6 DSC [1990DON/DRE]				
C ₁₂ H ₂₀ O	[4789-40-6] $\Delta_v H$	2,5-di- <i>tert</i> -butylfuran (274–323) 56.1 ± 1.1 298 GS [1998VER/WEL]				
C ₁₂ H ₂₀ O	[90-42-6] $\Delta_{\text{fus}} H$	2-cyclohexylcyclohexanone 18.0 277 [1992MAR/KOZ]				
C ₁₂ H ₂₀ O ₂	[76-49-3] $\Delta_v H$	bornyl acetate (319–496) 50.8 334 A [1987STE/MAL, 1947STU]				
C ₁₂ H ₂₀ O ₂	[105-87-3] $\Delta_v H$	geranyl acetate (346–516) 58.1 361 A [1987STE/MAL, 1947STU]				
C ₁₂ H ₂₀ O ₂	[125-12-2]	isobornyl acetate				

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(404–450)	56.1	419	A	[1987STE/MAL]
C ₁₂ H ₂₀ O ₂	[na]	bicyclo[2,2,1]heptane-7-one 2,2-dimethylpropylene acetal				
	$\Delta_v H$	(293–323)	60.5 ± 0.9	298	GS	[2002VER]
C ₁₂ H ₂₀ O ₂	[115-95-7]	3,7-dimethyl-1,6-octadien-3-ol acetate (linalyl acetate)				
	$\Delta_v H$	(281–490)	57.8	296	A	[1987STE/MAL]
	$\Delta_v H$	(328–493)	56.8	343		[1947STU]
C ₁₂ H ₂₀ O ₂	[80-26-2]	terpineol acetate				
	$\Delta_v H$	(310–424)	68.1	325	A	[1987STE/MAL]
C ₁₂ H ₂₀ O ₂	[217467-40-8]	bicyclo[2.2.1]heptane-7-one 2,2-dimethylpropylene ketal				
	$\Delta_{\text{fus}} H$		23.9	346.7		[1998VER/PEN]
	$\Delta_{\text{sub}} H$		84.0 ± 0.9	298		[1998VER/PEN]
C ₁₂ H ₂₀ O ₂	[10329-90-5]	1,7-cyclododecanedione				
	$\Delta_{\text{fus}} H$		15.77	405.2		[1972ALV/BOR]
C ₁₂ H ₂₀ O ₂	[28746-99-8]	2-(1'-hydroxycyclohexyl)cyclohexanone				
	$\Delta_{\text{fus}} H$	(5–310)	20.81	306.8	DSC	[2006SHE/KAB]
C ₁₂ H ₂₀ O ₃	[49540-29-6]	3,3,6,6-tetramethyloctanedioic anhydride				
	$\Delta_{\text{fus}} H$		18.83	344.2		[1974BOR]
C ₁₂ H ₂₀ O ₄	[na]	1,5-cyclooctanedione bis ethylene ketal				
	$\Delta_{\text{fus}} H$		18.03	296.2		[1972ALV/BOR]
C ₁₂ H ₂₀ O ₄	[105-76-0]	dibutyl maleate				
	$\Delta_v H$	(255–550)	41.1	270	A	[1987STE/MAL]
C ₁₂ H ₂₀ O ₅	[na]	2-ethoxycarbonylpropionic acid, cyclohexyl ester				
	$\Delta_v H$	(388–523)	67.6	403	A	[1987STE/MAL]
C ₁₂ H ₂₀ O ₆	[139-45-7]	tripropionin				
	$\Delta_v H$		91.4 ± 0.4	298	C	[1986NIL/WAD]
C ₁₂ H ₂₀ O ₇	[77-93-0]	triethyl citrate				
	$\Delta_v H$	(380–567)	68.2	395	A	[1987STE/MAL]
C ₁₂ H ₂₀ S	[880-36-4]	2-octylthiophene				
	$\Delta_v H$		65.4 ± 1.4	298	C	[2007RIB/SAN]
C ₁₂ H ₂₀ S	[65016-62-8]	3-octylthiophene				
	$\Delta_v H$		67.6 ± 1.5	298	C	[2007RIB/SAN]
C ₁₂ H ₂₁ O ₄ P	[na]	trimethylallyl phosphate				
	$\Delta_v H$	(367–597)	53.9	381		[1947STU]
C ₁₂ H ₂₁ N ₂ O ₃ PS	[333-41-5]	diazinon				
	$\Delta_v H$	(293–398)	87.4	308	A	[1987STE/MAL, 1999DYK/SVO]
C ₁₂ H ₂₂	[92-51-3]	cis bicyclohexyl				
	$\Delta_v H$	(331–511)	53.8	346	A	[1987STE/MAL]
C ₁₂ H ₂₂	[92-51-3]	bicyclohexyl				
	$\Delta_{\text{ms}} H$	(6–440)	3.7	267.4		
	$\Delta_{\text{ms}} H$	(6–440)	7.26	273		
	$\Delta_{\text{fus}} H$	(6–440)	6.86	276.8	AC	[1998CHI/COW]
	$\Delta_{\text{ms}} H$		1.54	256.1		
	$\Delta_{\text{ms}} H$		0.74	267.5		

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{fus}}H$		7.08	273.5		
	$\Delta_{\text{fus}}H$		6.78	277.2	DSC	[1996DOM/HEA, 1983ORO/MRA]
	Δ_vH		50.1	435		[1981WIE/KOB]
	Δ_vH		42.5	525		[1981WIE/KOB]
	Δ_vH		58.0 ± 0.2	298	C	[1978MON/ROS]
	Δ_vH		58.5 ± 0.6	298	C	[1978MAN, 1978MON/ROS]
C ₁₂ H ₂₂	[6975-99-1]	6-dodecyne				
	Δ_vH	(373–388)	60.9	380	A	[1987STE/MAL]
C ₁₂ H ₂₂	[na]	perhydroacenaphthylene				
	Δ_vH	(422–514)	49.6	437	EB	[2000ROH/CEN]
C ₁₂ H ₂₂ Cl ₄	[210115-98-3]	1,2,11,12-tetrachlorododecane				
	Δ_vH		81.9			[1998DRO/TOM]
C ₁₂ H ₂₂ N ₂ O ₂	[56403-09-9]	1,8-diaza-2,9-dioxocyclotetradecane				
	$\Delta_{\text{fus}}H$		13.6	517.4		
	$\Delta_{\text{fus}}H$		49.3	617.8	DSC	[1993SCH/KVA]
C ₁₂ H ₂₂ N ₆	[na]	1-(piperidinyl)-3,5-(dimethylamino)-s-triazine				
	$\Delta_{\text{fus}}H$		23.22	361.5		[1991ACR]
C ₁₂ H ₂₂ O	[58879-21-3]	<i>trans</i> 2-cyclohexylcyclohexanol				
	$\Delta_{\text{fus}}H$		14.52	325.8		[1997MAK/KAB]
	$\Delta_{\text{sub}}H$	(293–325)	98.6 ± 0.5	320	ME	[1997MAK/KAB]
	Δ_vH	(324–364)	83.2 ± 1.2	344	ME	[1997MAK/KAB]
C ₁₂ H ₂₂ O	[830-13-7]	cyclododecanone				
	$\Delta_{\text{fus}}H$		16.85	335.6		[1996ROU/JIM, 1998GON/SZW]
	$\Delta_{\text{sub}}H$		83.2 ± 0.3	298	ME	[1996ROU/JIM]
	Δ_vH	(373–443)	61	388	A	[1987STE/MAL]
	Δ_vH	(408–450)	57.9	423	A, EB	[1987STE/MAL, 1976MEY/HOT]
	Δ_vH	(458–556)	54.7	473	A, EB	[1987STE/MAL, 1976MEY/HOT]
	Δ_vH		65.5 ± 0.6	298		[1972WOL]
C ₁₂ H ₂₂ O	[81149-96-4]	(Z) 2-dodecenal				
	Δ_vH	(323–363)	72.5	298	CGC	[1996KOU/HOS, 2000OVA/KOU]
C ₁₂ H ₂₂ O	[20407-84-5]	(E) 2-dodecenal				
	Δ_vH	(323–363)	72.6	298	CGC	[1996KOU/HOS, 2000OVA/KOU]
C ₁₂ H ₂₂ O	[68141-15-1]	(Z) 3-dodecenal				
	Δ_vH	(323–363)	69.6	298	CGC	[1996KOU/HOS, 2000OVA/KOU]
C ₁₂ H ₂₂ O	[76595-72-7]	(E) 3-dodecenal				
	Δ_vH	(323–363)	70.2	298	CGC	[1996KOU/HOS, 2000OVA/KOU]
C ₁₂ H ₂₂ O	[21944-98-9]	(Z) 4-dodecenal				
	Δ_vH	(323–363)	69.4	298	CGC	[1996KOU/HOS, 2000OVA/KOU]
C ₁₂ H ₂₂ O	[174155-48-7]	(E) 4-dodecenal				
	Δ_vH	(323–363)	69.9	298	CGC	[1996KOU/HOS, 2000OVA/KOU]
C ₁₂ H ₂₂ O	[68820-33-7]	(Z) 5-dodecenal				
	Δ_vH	(323–363)	69.1	298	CGC	[1996KOU/HOS, 2000OVA/KOU]
C ₁₂ H ₂₂ O	[68820-34-8]	(E) 5-dodecenal				

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound		T _m (K)	Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)			
	$\Delta_v H$	(323–363)	69.6	298	CGC	[1996KOU/HOS, 2000OVA/KOU]
C ₁₂ H ₂₂ O	[12674-61-7] $\Delta_v H$	(Z) 6-dodecenal (323–363)	69.2	298	CGC	[1996KOU/HOS, 2000OVA/KOU]
C ₁₂ H ₂₂ O	[174155-49-8] $\Delta_v H$	(E) 6-dodecenal (323–363)	67.7	298	CGC	[1996KOU/HOS, 2000OVA/KOU]
C ₁₂ H ₂₂ O	[63851-40-1] $\Delta_v H$	(Z) 7-dodecenal (323–363)	69.4	298	CGC	[1996KOU/HOS, 2000OVA/KOU]
C ₁₂ H ₂₂ O	[82944-76-1] $\Delta_v H$	(E) 7-dodecenal (323–363)	69.6	298	CGC	[1996KOU/HOS, 2000OVA/KOU]
C ₁₂ H ₂₂ O	[139909-65-2] $\Delta_v H$	(Z) 8-dodecenal (323–363)	70.0	298	CGC	[1996KOU/HOS, 2000OVA/KOU]
C ₁₂ H ₂₂ O	[144298-64-6] $\Delta_v H$	(E) 8-dodecenal (323–363)	69.8	298	CGC	[1996KOU/HOS, 2000OVA/KOU]
C ₁₂ H ₂₂ O	[56219-03-5] $\Delta_v H$	(Z) 9-dodecenal (323–363)	70.1	298	CGC	[1996KOU/HOS, 2000OVA/KOU]
C ₁₂ H ₂₂ O	[155235-07-7] $\Delta_v H$	(E) 9-dodecenal (323–363)	70.4	298	CGC	[1996KOU/HOS, 2000OVA/KOU]
C ₁₂ H ₂₂ O	[81892-61-7] $\Delta_v H$	(Z) 10-dodecenal (323–363)	71.0	298	CGC	[1996KOU/HOS, 2000OVA/KOU]
C ₁₂ H ₂₂ O	[81892-62-8] $\Delta_v H$	(E) 10-dodecenal (323–363)	70.9	298	CGC	[1996KOU/HOS, 2000OVA/KOU]
C ₁₂ H ₂₂ O ₂	[947-05-7] $\Delta_v H$	dodecanolactone (377–403)	64.2 ± 1.1	390	MM	[1991WIB/WAL]
	$\Delta_v H$	(377–403)	70.5 ± 1.7	298	MM	[1991WIB/WAL]
C ₁₂ H ₂₂ O ₂	[na] $\Delta_v H$	acetic acid, 4- <i>tert</i> -butylcyclohexyl ester (285–318)	63.8	300	A, ME	[1987STE/MAL, 1958SER/VOI, 1957SER/VOI]
C ₁₂ H ₂₂ O ₂	[na] $\Delta_v H$	3,3-dimethylbutanoic acid, cyclohexyl ester (333–378)	62.1	298	CGC	[1999VER/HEI]
C ₁₂ H ₂₂ O ₂	[na] $\Delta_v H$	1-methylcyclohexyl pivalate (333–378)	57.9	298	CGC	[1999VER/HEI]
C ₁₂ H ₂₂ O ₂	[na] $\Delta_v H$	3-methylcyclohexyl pivalate (333–378)	60.5	298	CGC	[1999VER/HEI]
C ₁₂ H ₂₂ O ₂	[na] $\Delta_v H$	4-methylcyclohexyl pivalate (333–378)	60.9	298	CGC	[1999VER/HEI]
C ₁₂ H ₂₂ O ₂	[16409-45-3] $\Delta_v H$	(<i>d</i>) menthyl acetate (330–500)	55.3	345	A	[1987STE/MAL, 1947STU]
C ₁₂ H ₂₂ O ₂	[150-84-5] $\Delta_v H$	citronellyl acetate (347–490)	68.7	362	A	[1987STE/MAL, 1947STU]
C ₁₂ H ₂₂ O ₂	[61732-97-6] $\Delta_v H$	2-(1-ethylpentyl)-4,7-dihydro-1,3-dioxepin (333–453)	66.3	348	A	[1987STE/MAL]
C ₁₂ H ₂₂ O ₂	[2664-55-3] $\Delta_{\text{fus}} H$	nonyl acrylate	23.36	236.5		[1996DOM/HEA]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₂ H ₂₂ O ₂	[2157-01-9] $\Delta_{\text{fus}}H$	octyl methacrylate	24.09	230.3		[1990DOM/HEA]
	Δ_vH	(384–513)	55.6	399	A	[1987STE/MAL]
C ₁₂ H ₂₂ O ₂	[111-81-9] Δ_vH	methyl 10-undecenoate	59.2	412	A	[1987STE/MAL]
C ₁₂ H ₂₂ O ₂	[81634-99-3] Δ_vH	(Z) 3-decenyl acetate	69.5	298	GC	[1997KOU/HOS, 2000OVA/KOU]
	Δ_vH	(299–313)	72.0	306	GC	[1983OLS/JON]
C ₁₂ H ₂₂ O ₂	[81634-98-2] Δ_vH	(E) 3-decenyl acetate	70.1	298	GC	[1997KOU/HOS, 2000OVA/KOU]
C ₁₂ H ₂₂ O ₂	[67452-27-1] Δ_vH	(Z) 4-decenyl acetate	69.0	298	GC	[1997KOU/HOS, 2000OVA/KOU]
C ₁₂ H ₂₂ O ₂	[69222-16-8] Δ_vH	(E) 4-decenyl acetate	70.1	298	GC	[1997KOU/HOS, 2000OVA/KOU]
C ₁₂ H ₂₂ O ₂	[67446-07-5] Δ_vH	(Z) 5-decenyl acetate	69.7	298	GC	[1997KOU/HOS, 2000OVA/KOU]
	Δ_vH	(299–313)	72.0	306	GC	[1983OLS/JON]
C ₁₂ H ₂₂ O ₂	[38421-90-8] Δ_vH	(E) 5-decenyl acetate	70.6	298	GC	[1997KOU/HOS, 2000OVA/KOU]
C ₁₂ H ₂₂ O ₂	[68760-70-3] Δ_vH	(Z) 6-decenyl acetate	70.1	298	GC	[1997KOU/HOS, 2000OVA/KOU]
C ₁₂ H ₂₂ O ₂	[na] Δ_vH	(E) 6-decenyl acetate	70.6	298	GC	[1997KOU/HOS, 2000OVA/KOU]
	Δ_vH	(299–313)	72.0	306	GC	[1983OLS/JON]
C ₁₂ H ₂₂ O ₂	[13857-03-9] Δ_vH	(Z) 7-decenyl acetate	70.7	298	GC	[1997KOU/HOS, 2000OVA/KOU]
C ₁₂ H ₂₂ O ₂	[13857-04-0] Δ_vH	(E) 7-decenyl acetate	71.1	298	GC	[1997KOU/HOS, 2000OVA/KOU]
C ₁₂ H ₂₂ O ₂	[83808-51-9] Δ_vH	(Z) 8-decenyl acetate	71.5	298	GC	[1997KOU/HOS, 2000OVA/KOU]
C ₁₂ H ₂₂ O ₂	[83808-51-9] Δ_vH	(Z) 8-decenyl acetate	71.5	298	GC	[1997KOU/HOS, 2000OVA/KOU]
C ₁₂ H ₂₂ O ₃	[na] Δ_vH	heptyl levulinate	62.6	408	A	[1987STE/MAL]
	Δ_vH	(393–558)	60.0	496		[1931SCH/COW]
C ₁₂ H ₂₂ O ₃	[18871-14-2] Δ_vH	3-pentyl-4-acetoxytetrahydro-2H-pyran	65.8	398	A	[1987STE/MAL]
C ₁₂ H ₂₂ O ₄	[106-19-4] Δ_vH	dipropyl adipate	63.6	428	A	[1987STE/MAL]
C ₁₂ H ₂₂ O ₄	[141-03-7] $\Delta_{\text{fus}}H$	di- <i>n</i> -butyl succinate	29.21	244.1		[1996DOM/HEA]
C ₁₂ H ₂₂ O ₄	[5398-08-3] Δ_vH	isopentylmalonic acid, diethyl ester	64.1	392	A	[1987STE/MAL]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound							
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference			
C ₁₂ H ₂₂ O ₄	[22328-91-2] $\Delta_v H$	(1-methylbutyl)malonic acid, diethyl ester (395–516)	67.4	410	A	[1987STE/MAL]			
C ₁₂ H ₂₂ O ₄	[106-79-6] $\Delta_v H$	dimethyl sebacate (304–374)	86.4 ± 0.3	298	GS	[2006VER/KOZ]			
C ₁₂ H ₂₂ O ₄	[2051-00-5] $\Delta_v H$	diisopentyl oxalate (358–538)	58.6	373	A	[1987STE/MAL, 1947STU]			
C ₁₂ H ₂₂ O ₄	[693-23-2] $\Delta_{\text{fus}} H$	dodecanedioic acid	52.5	401.6	DSC	[2008VEN/BAY]			
	$\Delta_{\text{fus}} H$		49.8	400.3		[2005ROU/TEM]			
	$\Delta_{\text{fus}} H$		50.57	402.5		[1996DOM/HEA]			
	$\Delta_{\text{sub}} H$	(346–377)	169 ± 4		TPD	[2007CAP/LOV]			
	$\Delta_{\text{sub}} H$	(298–316)	156		TPTD	[2005CHA/ZIE]			
	$\Delta_{\text{sub}} H$	(375–296)	153.1 ± 2.9	386	ME	[1960DAV/THO, 1970COX/PIL]			
	$\Delta_v H$	(424–503)	130.0 ± 2.3	298	CGC	[2005ROU/TEM]			
C ₁₂ H ₂₂ O ₄ S	[4121-12-4] $\Delta_v H$	thiodiglycolic acid, diethyl ester (298–383)	75.7	313	A	[1987STE/MAL, 1999DYK/SVO]			
C ₁₂ H ₂₂ O ₅	[902261-31-8] $\Delta_v H$	butyl[1-(butoxycarbonyl)ethyl] carbonate (338–513)	68.1	353	A	[1987STE/MAL]			
C ₁₂ H ₂₂ O ₅	[na] $\Delta_v H$	pentyl[1-(ethoxycarbonyl)isopropyl] carbonate (368–513)	63.8	383	A	[1987STE/MAL]			
C ₁₂ H ₂₂ O ₆	[856371-29-4] $\Delta_v H$	lactic acid, O-ethoxycarbonyl, 2-butoxyethyl ester (383–521)	74.6	398	A	[1987STE/MAL]			
C ₁₂ H ₂₂ O ₆	[87-92-3] $\Delta_v H$	dibutyl tartrate (428–511)	79.8	443	A	[1987STE/MAL]			
C ₁₂ H ₂₂ O ₆	[4054-82-4] $\Delta_v H$	(<i>d</i>) diisobutyl tartrate (390–597)	64.6	405	A	[1987STE/MAL]			
C ₁₂ H ₂₂ O ₁₁	[528-50-7] $\Delta_{\text{sub}} H$	(<i>d</i>) cellobiose (474–488)	302 ± 44.0	481	ME	[1999OJA/SUU]			
C ₁₂ H ₂₂ O ₁₁	[14641-93-1] $\Delta_{\text{fus}} H$	α -lactose	75.2	496.2		[2000MAC/COU, 1983RAE/SCH]			
C ₁₂ H ₂₂ O ₁₁	[57-50-1] $\Delta_{\text{fus}} H$	sucrose	46.2	459		[1988SOP/KEA]			
C ₁₂ H ₂₂ S	[7133-46-2] $\Delta_{\text{us}} H$	dicyclohexyl sulfide	10.01	274.7					
	$\Delta_{\text{fus}} H$		5.68	284.2			[2004STE/CHI]		
	$\Delta_v H$		(335–523)	65.4 ± 0.2			340	IP,EB	[2004STE/CHI]
	$\Delta_v H$		(335–523)	62.5 ± 0.1			380	IP,EB	[2004STE/CHI]
	$\Delta_v H$		(335–523)	59.5 ± 0.1			420	IP,EB	[2004STE/CHI]
	$\Delta_v H$		(335–523)	56.6 ± 0.1			440	IP,EB	[2004STE/CHI]
	$\Delta_v H$		(335–523)	53.7 ± 0.1			480	IP,EB	[2004STE/CHI]
$\Delta_v H$	(421–523)	69.0 ± 0.7	298	EB	[1997STE/CHI4]				
C ₁₂ H ₂₃ N	[101-83-7] $\Delta_v H$	dicyclohexylamine (408–529)	54.0	423	A	[1987STE/MAL]			
C ₁₂ H ₂₃ N	[2437-25-4]	lauroitrile							

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(298–367)	74.9 ± 0.2	298	GS	[2005EME/VER]
	$\Delta_v H$		76.1 ± 0.1	298	C	[1977STRI/SUN]
	$\Delta_v H$	(393–462)	65.2	408	EB	[1971MEY/REN]
	$\Delta_v H$	(440–556)	60.7	455	A, EB	[1987STE/MAL, 1971MEY/REN, 1973MEY/HOT]
C ₁₂ H ₂₃ NO ₃	[14305-32-9] $\Delta_{\text{fus}}H$	N-decanoylglycine	42.2	387.6	DSC	[1986MIY/MAT]
C ₁₂ H ₂₃ N ₇	[5512-05-0] $\Delta_{\text{fus}}H$	1-(4'-methylpiperiziny)-3,5-bis(dimethylamino)-s-triazine	20.42	354.2	DSC	[1989BRA/RYT]
C ₁₂ H ₂₄	[294-62-2] $\Delta_{\text{ms}}H$ $\Delta_{\text{fus}}H$ $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$	cyclododecane	0.6 14.8 76.2 76.4 ± 1.7 63.0 62.8 52.6 49.8	199 333.8 298	CGC-DSC	[1987DRO/MOL] [1998CHI/HES] [1957VAN]
				298	CGC	[1998CHI/HES]
		(403–453)	62.8	298	CGC	[1995CHI/HOS]
		(386–441)	52.6	401	A, EB	[1987STE/MAL, 1976MEY/HOT]
		(440–529)	49.8	455	A, EB	[1987STE/MAL, 1976MEY/HOT]
C ₁₂ H ₂₄	[112-41-4] $\Delta_{\text{ms}}H$ $\Delta_{\text{fus}}H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$	1-dodecene	4.55 19.87 60.8 ± 0.3 60.3 51.1	212.9 237.9 298 298 411	C A	[1996DOM/HEA] [1976STR2, 1977MAN/SEL] [1971WIL/ZWO] [1987STE/MAL, 1950FOR/CAM]
C ₁₂ H ₂₄	[4292-75-5] $\Delta_v H$ $\Delta_v H$ $\Delta_v H$	hexylcyclohexane	55.9 ± 0.5 59.0 ± 0.5 59.9	298 298 298	GC GCC	[1987AZA] [1978FUC/PEA] [1971WIL/ZWO]
C ₁₂ H ₂₄	[5617-42-5] $\Delta_v H$	heptylcyclopentane	60.8	298		[1971WIL/ZWO]
C ₁₂ H ₂₄	[27656-49-1] $\Delta_v H$ $\Delta_v H$	<i>trans</i> 2,2,4,6,6-pentamethyl-3-heptene	65.6 ± 0.5 65.9 ± 0.3	305 298	GS GS	[2000VER/WAN] [2000VER/WAN]
C ₁₂ H ₂₄	[27656-50-4] $\Delta_v H$ $\Delta_v H$	<i>cis</i> 2,2,4,6,6-pentamethyl-3-heptene	63.0 ± 0.5 63.2 ± 0.5	303 298	GS GS	[2000VER/WAN] [2000VER/WAN]
C ₁₂ H ₂₄ Cl ₂	[3922-28-9] $\Delta_v H$	1,12-dichlorododecane	73.1			[1998DRO/TOM]
C ₁₂ H ₂₄ N ₂ O ₂	[10263-96-4] $\Delta_{\text{fus}}H$	N,N'-di- <i>n</i> -propyladipamide	36.11	452		[1984DOM/EVA]
C ₁₂ H ₂₄ N ₂ O ₂	[6224-99-3] $\Delta_{\text{ms}}H$ $\Delta_{\text{fus}}H$	dodecandiamide	5.09 73.7	422.8 466.1	DSC	[2006BAD/DEL]
C ₁₂ H ₂₄ N ₂ O ₂	[3129-91-7] $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$	dicyclohexyl ammonium nitrite	99.1 U 161.8	294	TE	[1987STE/MAL, 1965MAR] [1985TRU/KRA]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{sub}}H$	(308–339)	105.9	324		[1961ROZ/POL]
C ₁₂ H ₂₄ O	[1724-39-6]	cyclododecanol				
	Δ_vH	(405–468)	68.8	420	A	[1987STE/MAL]
	Δ_vH	(467–557)	57.1	482	A	[1987STE/MAL]
C ₁₂ H ₂₄ O	[112-54-9]	dodecanal				
	Δ_vH	(314–347)	68.3 ± 0.9	298	GS	[2003VER/KRA2]
	Δ_vH	(308–353)	70.2	298	CGC	[1996KOU/HOS, 2000OVA/KOU]
	Δ_vH	(350–530)	56.5	365	A	[1987STE/MAL, 1947STU]
C ₁₂ H ₂₄ O	[6175-49-1]	2-dodecanone				
	Δ_vH	(350–520)	61.1	365	A	[1987STE/MAL, 1947STU]
	Δ_vH		71.8 ± 0.6	298	C	[1977SEL]
	Δ_vH	(386–609)	60.8	401	A	[1987STE/MAL, 1975AMB/ELL]
	Δ_vH	(386–609)	48.1	524		[1975AMB/ELL]
C ₁₂ H ₂₄ O	[19321-39-2]	ethyl <i>p</i> -menthyl ether				
	Δ_vH	(366–414)	50.9	381	A	[1987STE/MAL]
C ₁₂ H ₂₄ O	[20999-39-7]	1-heptylcyclopentanol				
	Δ_vH	(395–524)	58.6	410	A	[1987STE/MAL]
C ₁₂ H ₂₄ O	[3964-63-4]	1-hexylcyclohexanol				
	Δ_vH	(380–491)	53.5	395	A	[1987STE/MAL]
C ₁₂ H ₂₄ O	[69064-36-4]	(Z) 2-dodecen-1-ol				
	Δ_vH	(333–373)	90.7	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₂ H ₂₄ O	[69064-37-5]	(E) 2-dodecen-1-ol				
	Δ_vH	(333–373)	91.0	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₂ H ₂₄ O	[32451-95-9]	(Z) 3-dodecen-1-ol				
	Δ_vH	(333–373)	89.3	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₂ H ₂₄ O	[68900-87-8]	(E) 3-dodecen-1-ol				
	Δ_vH	(333–373)	89.2	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₂ H ₂₄ O	[40642-37-3]	(Z) 4-dodecen-1-ol				
	Δ_vH	(333–373)	89.9	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₂ H ₂₄ O	[81745-38-2]	(E) 4-dodecen-1-ol				
	Δ_vH	(333–373)	90.6	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₂ H ₂₄ O	[40642-38-4]	(Z) 5-dodecen-1-ol				
	Δ_vH	(333–373)	90.2	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₂ H ₂₄ O	[62936-12-3]	(E) 5-dodecen-1-ol				
	Δ_vH	(333–373)	90.7	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₂ H ₂₄ O	[40642-39-5]	(Z) 6-dodecen-1-ol				
	Δ_vH	(333–373)	90.2	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₂ H ₂₄ O	[52957-14-9]	(E) 6-dodecen-1-ol				
	Δ_vH	(333–373)	90.7	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₂ H ₂₄ O	[20056-92-2]	(Z) 7-dodecen-1-ol				
	Δ_vH	(333–373)	90.5	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₂ H ₂₄ O	[16695-40-2]	(E) 7-dodecen-1-ol				
	Δ_vH	(333–373)	90.8	298	CGC	[2000OVA/KOU, 1994KOU/HOS]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound					
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference	
C ₁₂ H ₂₄ O	[40642-40-8] $\Delta_v H$	(Z) 8-dodecen-1-ol (333–373)	91.0	298	CGC	[2000OVA/KOU, 1994KOU/HOS]	
C ₁₂ H ₂₄ O	[42513-42-8] $\Delta_v H$	(E) 8-dodecen-1-ol (333–373)	91.0	298	CGC	[2000OVA/KOU, 1994KOU/HOS]	
C ₁₂ H ₂₄ O	[35148-18-6] $\Delta_v H$	(Z) 9-dodecen-1-ol (333–373)	91.1	298	CGC	[2000OVA/KOU, 1994KOU/HOS]	
C ₁₂ H ₂₄ O	[35237-62-8] $\Delta_v H$	(E) 9-dodecen-1-ol (333–373)	91.7	298	CGC	[2000OVA/KOU, 1994KOU/HOS]	
C ₁₂ H ₂₄ O	[35289-30-6] $\Delta_v H$	(Z) 10-dodecen-1-ol (333–373)	92.4	298	CGC	[2000OVA/KOU, 1994KOU/HOS]	
C ₁₂ H ₂₄ O	[35237-63-9] $\Delta_v H$	(E) 10-dodecen-1-ol (333–373)	91.9	298	CGC	[2000OVA/KOU, 1994KOU/HOS]	
C ₁₂ H ₂₄ O ₂	[110-38-3] $\Delta_{\text{fus}} H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$	ethyl decanoate (5–370)	32.29	253.6	AC	[2009ZAI/PAU]	
			69.9 ± 0.7	305	C	[2009ZAI/PAU]	
			70.5	298		[2009ZAI/PAU]	
			(404–440)	58.4 ± 0.1	422	MM	[1991WIB/WAL]
			(404–440)	67.4 ± 1.3	298	MM	[1991WIB/WAL]
			(359–515)	59.6	374	A	[1987STE/MAL]
C ₁₂ H ₂₄ O ₂	[112-17-4] $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$	decyl acetate (284–321)	70.2 ± 0.3	298	GS	[2006KRA/VER]	
			71.6	298	GC	[1997KOU/HOS, 2000OVA/KOU]	
			(363–515)	61.9	378	A	[1987STE/MAL]
			(299–313)	72	306	GC	[1983OLS/JON]
			(445–530)	56.3	460	DTA	[1980MEY/AWE]
C ₁₂ H ₂₄ O ₂	[61732-91-0] $\Delta_v H$	4,5-dimethyl-2-heptyl-1,3-dioxolane (333–453)	69.8	346	A	[1987STE/MAL]	
C ₁₂ H ₂₄ O ₂	[143-07-7] $\Delta_{\text{fus}} H$ $\Delta_{\text{fus}} H$ $\Delta_{\text{fus}} H$ $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$	dodecanoic acid	36.1	316.6	DSC	[2007MOR/COR]	
			34.7	317.9	DSC	[2007MIS/MIS]	
			36.65	316.9		[1996DOM/HEA]	
			147.2 ± 4	298	TPD	[2008CAP/LOV]	
			(293–303)	127.9	298		[1987STE/MAL]
			(293–308)	132.6	300	ME	[1968BAC/NOV]
			(296–314)	140.2 ± 3.3	304	ME	[1961DAV/MAL]
			(293–313)	117.2 ± 2.9	303	ME	[1957LIT]
			(393–573)	88.8	408	A	[1987STE/MAL]
			(321–341)	95.8	332	ME, TE	[1982FUC/HAL]
				81.3	437	I	[1943CRA]
C ₁₂ H ₂₄ O ₂	[61732-93-2] $\Delta_v H$	2-(1-ethylpentyl)-1,3-dioxepane (333–373)	68.1	348	A	[1987STE/MAL]	
C ₁₂ H ₂₄ O ₂	[61732-92-1] $\Delta_v H$	2-heptyl-1,3-dioxepane (328–373)	70.3	343	A	[1987STE/MAL]	
C ₁₂ H ₂₄ O ₂	[62159-06-2] $\Delta_v H$	3-heptyl-4-hydroxytetrahydro-2H-pyran (383–453)	77.6	398	A	[1987STE/MAL]	
C ₁₂ H ₂₄ O ₂	[23433-02-5]	4-octyl-1,3-dioxane					

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(353–453)	65.5	368	A	[1987STE/MAL]
C ₁₂ H ₂₄ O ₂	[1731-86-8]	methyl undecanoate				
	$\Delta_v H$		66.1	350		[2002VAN/VAN]
	$\Delta_v H$		67.0 ± 0.1	340		[2002VAN/VAN]
	$\Delta_v H$		70.8 ± 0.4	298		[2002VAN/VAN]
	$\Delta_v H$	(433–473)	70.6	298	CGC	[1995CHI/HOS]
	$\Delta_v H$		71.4 ± 0.3	298	C	[1977MAN/SEL]
	$\Delta_v H$	(393–473)	60.9	408	A, E	[1987STE/MAL, 1963ROS/SCH]
C ₁₂ H ₂₄ O ₂	[245658-36-0]	3,3-dimethylbutanoic acid, 1,1,2-trimethylpropyl ester				
	$\Delta_v H$	(333–378)	57.1	298	CGC	[1999VER/HEI]
C ₁₂ H ₂₄ O ₂	[245658-40-6]	2,2-dimethylpropanoic acid, 1,1,3-trimethylbutyl ester				
	$\Delta_v H$	(333–378)	54.2	298	CGC	[1999VER/HEI]
C ₁₂ H ₂₄ O ₂	[245658-43-9]	2,6-dimethyl-2-heptanol propanoate				
	$\Delta_v H$	(333–378)	59.4	298	CGC	[1999VER/HEI]
C ₁₂ H ₂₄ O ₃	[2388-12-7]	peroxydodecanoic acid				
	$\Delta_{\text{sub}} H$	(293–303)	131.4 ± 1.7	298	ME	[1980SWA/KWA]
C ₁₂ H ₂₄ O ₃	[na]	pentyl 2-butoxypropionate				
	$\Delta_v H$	(373–398)	47.3	385	A	[1987STE/MAL]
C ₁₂ H ₂₄ O ₃	[7419-98-9]	methyl 3-octyloxypropionate				
	$\Delta_v H$	(373–513)	59.8	388	A	[1987STE/MAL]
C ₁₂ H ₂₄ O ₄	[53759-20-9]	2,2,8,8-tetramethyl-1,3,7,9-tetraoxacyclododecane				
	$\Delta_{\text{fus}} H$		23.4	383		[1975BOR]
C ₁₂ H ₂₄ O ₄	[43091-26-5]	1,3,9,11-tetraoxacyclohexadecane				
	$\Delta_{\text{fus}} H$		35.56	332		[1973DAL/EKE]
C ₁₂ H ₂₄ O ₄	[20732-35-8]	3,6-dimethyl-3,6-di- <i>tert</i> -butyl-1,2,4,5-tetraoxacyclohexane				
	$\Delta_v H$	(403–473)	53.7	298	CGC	[2007CAN/EYL]
C ₁₂ H ₂₄ O ₆	[24748-23-0]	3,6,9-triethyl-3,6,9-trimethyl-1,2,4,5,7,8-hexaoxacyclononane				
	$\Delta_v H$	(403–473)	59.2	298	CGC	[2007CAN/EYL]
C ₁₂ H ₂₄ O ₆	[17455-13-9]	1,4,7,10,13,16-hexaoxacyclooctadecane (18-crown-6)				
	$\Delta_{\text{fus}} H$		34.0	312.2		[1972DAL/KRI]
	$\Delta_{\text{sub}} H$		119.1 ± 6.7	298	CGC-DSC	[2000NIC/ORF]
	$\Delta_{\text{sub}} H$		133.2 ± 0.3		C	[1990BRI/WAD]
	$\Delta_v H$		86.1 ± 6.7	298	CGC	[2000NIC/ORF]
C ₁₂ H ₂₄ O ₁₁	[585-88-6]	1,4-O- α -D-glucopyranosyl-D-glucitol (maltitol)				
	$\Delta_{\text{fus}} H$		55.07	420	DSC	[2001LEB/VAN, 2003LEB/VAN]
C ₁₂ H ₂₄ O ₁₁	[534-73-6]	α -(<i>d</i>)-glucopyranosyl-1,6-sorbitol				
	$\Delta_{\text{fus}} H$		56.4	439		[1996CAM/FIG]
C ₁₂ H ₂₄ O ₁₁	[20942-99-8]	α -(<i>d</i>)-glucopyranosyl-1,6-mannitol				
	$\Delta_{\text{fus}} H$		55.0	440.8		[1996CAM/FIG]
C ₁₂ H ₂₄ O ₁₁	[64519-82-0]	6-O- α -D-glucopyranosyl-D-arabino-hexitol (isomalt)				
	$\Delta_{\text{fus}} H$		44.3		DSC	[2002BOR/CES]
C ₁₂ H ₂₄ S ₄	[297181-32-9]	1,4,8,11-tetrathiacyclohexadecane				
	$\Delta_{\text{us}} H$ [solid-solid (<i>I</i>)]		32.0	328.2		

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound		T _m (K)	Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)			
	$\Delta_{\text{fus}}H$		5.2	333.2		[2002ROC/GRI]
	$\Delta_{\text{us}}H$ [solid-solid (II)]		27.0	328.2		
	$\Delta_{\text{fus}}H$		5.2	333.2	DSC	[2002ROC/GRI]
C₁₂H₂₅Br	[143-15-7]	1-bromododecane				
	Δ_vH		74.8 ± 0.4	298	C	[1976STR3, 1977MAN/SEL]
	Δ_vH	(411–610)	62.2	426	A, E	[1987STE/MAL, 1961LI/ROS, 1970DYK/VAN]
C₁₂H₂₅Cl	[112-52-7]	1-chlorododecane				
	Δ_vH		75.8	298		[2006BOL/NER2]
	Δ_vH		73.9 ± 1.4	298	GS	[2001PUR/CHI]
	Δ_vH	(390–520)	70.5	298		[1984BOU/FRI, 1991BAS/SVO]
	Δ_vH		71.9 ± 0.3	298	C	[1977MAN/SEL]
	Δ_vH		70.3 ± 0.5	298	C	[1975STR/SUN]
	Δ_vH	(389–519)	62.4	404	A, DTA	[1987STE/MAL, 1969KEM/KRE]
C₁₂H₂₅Cl	[2350-12-1]	(dl) 2-chlorododecane				
	Δ_vH	(283–328)	65.3	298	A	[1987STE/MAL, 1970DYK/VAN, 1962GEI/QUI]
C₁₂H₂₅Cl	[2350-12-1]	(dl) 3-chlorododecane				
	Δ_vH	(283–328)	65.9	298	A	[1987STE/MAL, 1970DYK/VAN, 1962GEI/QUI]
C₁₂H₂₅Cl	[2350-13-2]	(dl) 4-chlorododecane				
	Δ_vH	(283–328)	64.1	298	A	[1987STE/MAL, 1970DYK/VAN, 1962GEI/QUI]
C₁₂H₂₅Cl	[2350-14-3]	(dl) 5-chlorododecane				
	Δ_vH	(283–328)	65.9	298	A	[1987STE/MAL, 1970DYK/VAN, 1962GEI/QUI]
C₁₂H₂₅Cl	[26535-66-0]	6-chlorododecane				
	Δ_vH	(283–328)	65.5	298	A	[1987STE/MAL, 1970DYK/VAN, 1962GEI/QUI]
C₁₂H₂₅F	[334-68-9]	1-fluorododecane				
	Δ_vH	(288–328)	64.0 ± 0.2	298	GS	[1997SCH/VER]
	Δ_vH	(374–533)	56.2	389	A, E	[1987STE/MAL, 1961LI/ROS, 1970DYK/VAN]
C₁₂H₂₅I	[4292-19-7]	1-iodododecane				
	Δ_vH	(426–636)	79.9	298	A,E	[1987STE/MAL, 1961LI/ROS, 1970DYK/VAN, 2006BOL/NER]
	Δ_vH	(426–636)	63.5	441	A, E	[1987STE/MAL, 1961LI/ROS, 1970DYK/VAN]
C₁₂H₂₅NO	[996-97-4]	N,N-diethylcaprylamide				
	Δ_vH	(373–510)	71.2	388	A	[1987STE/MAL]
C₁₂H₂₅NO	[1120-16-7]	dodecanamide				
	$\Delta_{\text{us}}H$		9.7	321.1		
	$\Delta_{\text{fus}}H$		36.3	373.3	DSC	[2008ABA/BAD]
	$\Delta_{\text{sub}}H$	(349–368)	152.7 ± 0.8	358.5	ME	[1959DAV/JON2, 1987STE/MAL]
C₁₂H₂₆	[112-40-3]	dodecane				
	$\Delta_{\text{fus}}H$		35.7	263.1	DSC	[2004MON/RAJ]
	$\Delta_{\text{fus}}H$		36.82	263.6		[1996DOM/HEA]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{sub}}H$		100.2	298	B	[1972MOR3]
	$\Delta_{\text{sub}}H$		101.7	263	B	[1963BON]
	Δ_vH		62.1 ± 0.2	298	GS	[2001PUR/CHI]
	Δ_vH		60.3 ± 0.8	298	CGC	[2000NIC/ORF]
	Δ_vH		61.4	299	C	[1996VIT/CHA]
	Δ_vH		58.1	334	C	[1996VIT/CHA]
	Δ_vH		57.4	344	C	[1996VIT/CHA]
	Δ_vH	(373–423)	60.7	298	CGC	[1995CHI/HOS]
	Δ_vH	(363–413)	61.2	298	CGC	[1995CHI/HOS]
	Δ_vH	(423–473)	61.2	298	CGC	[1995CHI/HOS]
	Δ_vH		61.5	298		[1994RUZ/MAJ]
	Δ_vH	(263–371)	65.7	278		[1988SAS/JOS]
	Δ_vH	(278–400)	61.8	293	A	[1987STE/MAL]
	Δ_vH	(298–389)	61.1	313	GS	[1986ALL/JOS]
	Δ_vH		61.8 ± 0.5	298	C	[1976MEL/MAN]
	Δ_vH		61.2 ± 0.2	298	C	[1974MAN4]
	Δ_vH		60.4 ± 0.3	298	C	[1972MOR2]
	Δ_vH		61.3	298		[1971WIL/ZWO]
	Δ_vH	(400–492)	51.6	415	A, MM	[1987STE/MAL, 1945WIL/TAY]
C₁₂H₂₆	[7045-71-8]	2-methylundecane				
	Δ_vH	(356–484)	49.5	371	A	[1987STE/MAL]
C₁₂H₂₆	[1002-43-3]	(<i>dl</i>) 3-methylundecane				
	Δ_vH	(357–485)	48.8	372	A	[1987STE/MAL]
C₁₂H₂₆	[2980-69-0]	4-methylundecane				
	Δ_vH	(359–481)	51.6	374	A	[1987STE/MAL]
C₁₂H₂₆	[1632-70-8]	5-methylundecane				
	Δ_vH	(357–480)	50.3	372	A	[1987STE/MAL]
C₁₂H₂₆	[17312-44-6]	2,3-dimethyldecane				
	Δ_vH	(369–480)	50.0	384	A	[1987STE/MAL]
C₁₂H₂₆	[2801-84-5]	2,4-dimethyldecane				
	Δ_vH	(348–472)	47.5	363	A	[1987STE/MAL]
C₁₂H₂₆	[62184-10-5]	2,4,6-trimethylnonane				
	Δ_vH	(339–459)	46.4	354	A	[1987STE/MAL]
C₁₂H₂₆	[62199-46-6]	3,3,6,6-tetramethyloctane				
	Δ_vH	(347–463)	52.9	362	A	[1987STE/MAL]
C₁₂H₂₆	[13475-82-6]	2,2,4,6,6-pentamethylheptane				
	Δ_vH		49.0 ± 0.2	298	C	[1976MEL/MAN]
C₁₂H₂₆N₂O	[4128-38-5]	1-undecyl urea				
	$\Delta_{\text{fus}}H$		38.4	385.6	DSC	[2005HAS/TAJ]
C₁₂H₂₆O	[55962-01-1]	ethyl decyl ether				
	Δ_vH		65.9 ± 0.1	298	C	[1985KUS]
C₁₂H₂₆O	[112-58-3]	dihexyl ether				
	Δ_vH		63.6 ± 0.8	298	CGC	[2000NIC/ORF]
	Δ_vH	(353–393)	63.5	298	CGC	[1995CHI/HOS]
	Δ_vH	(372–510)	52.9	387	A	[1987STE/MAL]
	Δ_vH		64.1 ± 0.1	298	C	[1985KUS]
C₁₂H₂₆O	[51323-70-7]	octyl <i>tert</i> -butyl ether				

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound					
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference	
	$\Delta_v H$		61.4	298	CGC	[UR/VER, 2002VER, 2003VER/KRA]	
C ₁₂ H ₂₆ O	[na]	isobutyl <i>tert</i> -octyl ether					
	$\Delta_v H$		51.6	298	CGC	[UR/VER, 2002VER, 2003VER/KRA]	
C ₁₂ H ₂₆ O	[na]	butyl <i>tert</i> -octyl ether					
	$\Delta_v H$		52.9 ± 0.4	298	CGC	[UR/VER, 2002VER, 2003VER/KRA]	
C ₁₂ H ₂₆ O	[112-53-8]	1-dodecanol					
	$\Delta_{\text{fus}}H$		40.31	297.3		[2003VAN/VAN]	
	$\Delta_{\text{fus}}H$		40.17	300.2		[1993ACR]	
	$\Delta_{\text{sub}}H$	(285–294)	130.1 ± 1.2	290	ME	[1965DAV/KYB, 1987STE/MAL]	
	$\Delta_{\text{sub}}H$		129.3	298		[1965DAV/KYB]	
	$\Delta_v H$		90.8 ± 1.2	298	CGC	[2006NIC/KWE]	
	$\Delta_v H$	(303–348)	85.8	327	GS	[2001KUL/VER2]	
	$\Delta_v H$	(303–348)	90	298	GS	[2001KUL/VER2]	
	$\Delta_v H$	(373–423)	91.7	298	CGC	[1995CHI/HOS]	
	$\Delta_v H$	(353–393)	91.7	298	CGC	[1994KOU/HOS, 2000OVA/KOU]	
	$\Delta_v H$	(303–413)	80.5	358		[1992NGU/KAS]	
	$\Delta_v H$	(383–438)	73.8	398	A	[1987STE/MAL]	
	$\Delta_v H$	(505–550)	57.1	520	A	[1987STE/MAL]	
	$\Delta_v H$		84.7 ± 0.5	343	C	[1979SEV]	
	$\Delta_v H$		91.8 ± 0.6	298	C	[1979SEV]	
	$\Delta_v H$		92.0 ± 0.6	298	C	[1977MAN/SEL]	
	$\Delta_v H$	(297–363)	92.5	312		[1973WIL/ZWO]	
	$\Delta_v H$	(411–487)	67.6	426		[1973WIL/ZWO]	
	C ₁₂ H ₂₆ O	[10203-28-8]	2-dodecanol				
		$\Delta_v H$	(293–393)	87.0	308		[1999NGU/BER]
$\Delta_v H$		(293–343)	85.0	318	A, ME	[1987STE/MAL, 1962GEI/QUI2]	
C ₁₂ H ₂₆ O		[10203-30-2]	<i>(dl)</i> 3-dodecanol				
		$\Delta_v H$	(293–343)	78.3	318	A, ME	[1987STE/MAL, 1962GEI/QUI2]
C ₁₂ H ₂₆ O		[10203-32-4]	4-dodecanol				
		$\Delta_v H$	(293–343)	80.6	318	A, ME	[1987STE/MAL, 1962GEI/QUI2]
C ₁₂ H ₂₆ O		[10203-33-5]	5-dodecanol				
		$\Delta_v H$	(293–343)	79.4	318	A, ME	[1987STE/MAL, 1962GEI/QUI2]
C ₁₂ H ₂₆ O		[6836-38-0]	6-dodecanol				
	$\Delta_v H$	(293–343)	81.5	318	A, ME	[1987STE/MAL, 1962GEI/QUI2]	
C ₁₂ H ₂₆ O	[5457-42-1]	di- <i>tert</i> -butyl-isopropylmethanol					
	$\Delta_{\text{fus}}H$		2.09	314	DSC	[1998VER3]	
	Note: Compound likely has an unmeasured solid phase transition.						
	$\Delta_{\text{sub}}H$	(274–308)	59.3 ± 0.8	298	GS	[1998VER3]	
	$\Delta_{\text{sub}}H$	(274–308)	59.7 ± 0.8	291	GS	[1998VER3]	
	$\Delta_v H$	(317–348)	54.9 ± 0.8	333	GS	[1998VER5]	
	$\Delta_v H$	(317–348)	57.0 ± 0.8	298	GS	[1998VER5]	
	C ₁₂ H ₂₆ O ₂	[na]	<i>(dl)</i> 3,4-diethyl-3,4-dimethoxyhexane				

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(302–332)	59.8 ± 1.3	317	GS	[1990DOG/BEC]
C ₁₂ H ₂₆ O ₂	[5675-51-4] $\Delta_{\text{fus}} H$	1,12-dodecanediol	51.2	352		[2006UMN/KWE]
C ₁₂ H ₂₆ O ₃	[112-73-2] $\Delta_v H$ $\Delta_v H$	diethylene glycol dibutyl ether	73.8 ± 1.7	298	GC	[2000NIC/ORF]
		(293–528)	56.6	308	A	[1987STE/MAL]
C ₁₂ H ₂₆ O ₃	[113676-50-9] $\Delta_{\text{fus}} H$	3-(nonyloxy)-1,2-propanediol	29.5	297.2	DSC	[1993ACR]
C ₁₂ H ₂₆ O ₄	[41407-59-4] $\Delta_v H$	2,2-bis(tert-butylperoxy)butane	77.1	311	A	[1987STE/MAL]
C ₁₂ H ₂₆ O ₄	[na] $\Delta_v H$	tripropylene glycol monoisopropyl ether	56.9	370	A	[1987STE/MAL, 1947STU]
C ₁₂ H ₂₆ O ₄	[4161-33-5] $\Delta_{\text{fus}} H$	4,4'-[1,4-butanediyl bis(oxy)]bis-1-butanol	39.37	306.7	DSC	[1991BED/BOO]
C ₁₂ H ₂₆ S	[112-55-0] $\Delta_v H$	1-dodecanethiol	62.0	435		[1999DYK/SVO]
C ₁₂ H ₂₆ S	[6294-31-1] $\Delta_v H$	dihexylsulfide	72.4	310		[2004SAW/MOK]
C ₁₂ H ₂₆ S ₂	[33528-63-1] $\Delta_v H$	1,12-dodecanedithiol	77.8	469	A	[1987STE/MAL, 1943HAL/REI, 1999DYK/SVO]
C ₁₂ H ₂₆ S ₂	[10496-15-8] $\Delta_v H$	dihexyl disulfide	64.9	450		[1999DYK/SVO]
C ₁₂ H ₂₇ N	[124-22-1] $\Delta_v H$ $\Delta_v H$	dodecylamine	61.0	458	A, E	[1987STE/MAL, 1956MAN2]
		(443–545)	63.4	371		[1947STU]
C ₁₂ H ₂₇ N	[143-16-8] $\Delta_v H$	dihexylamine	55.1	423	A	[1987STE/MAL]
C ₁₂ H ₂₇ N	[1120-24-7] $\Delta_v H$	N,N-dimethyldecylamine	55.2	420	A	[1987STE/MAL]
C ₁₂ H ₂₇ N	[102-82-9] $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$	tributylamine	62.7 ± 1.3	298	CGC	[2009LIP/CHI, 2009LIP/HAN]
		(432–488)	49.9	447	EB	[2008GUA/YAN]
		(298–337)	64.4	313	A	[1987STE/MAL]
		(333–487)	48.1	348	A	[1987STE/MAL]
C ₁₂ H ₂₇ N	[1116-40-1] $\Delta_v H$	triisobutylamine	54.3	320	A	[1987STE/MAL, 1947STU]
C ₁₂ H ₂₇ NO ₂	[126835-64-1] $\Delta_{\text{fus}} H$	3-(nonylamino)-1,2-propanediol	53.2	343.2		[1993ACR]
C ₁₂ H ₂₇ O ₄ P	[126-73-8] $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$	tributyl phosphate	81.3	298	CGC	[2007PAN/ANT2]
		(443–483)	78.8	298	CGC	[2007PAN/ANT2]
		(423–463)	81.7	298	CGC	[2007PAN/ANT2]
		(453–493)	61.4	515	A	[1987STE/MAL]
C ₁₂ H ₂₇ O ₄ P	[126-71-6]	triisobutyl phosphate				

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(443–483)	73.0	298	CGC	[2007PAN/ANT2]
	$\Delta_v H$	(443–473)	76.3	298	CGC	[2007PAN/ANT2]
	$\Delta_v H$	(411–537)	62.8	426	A	[1987STE/MAL]
C₁₂H₂₇O₄P	[2528-45-2]	tri- <i>sec</i> -butyl phosphate				
	$\Delta_v H$	(413–453)	69.6	298	CGC	[2007PAN/ANT2]
	$\Delta_v H$	(443–483)	70.5	298	CGC	[2007PAN/ANT2]
C₁₂H₂₇P	[998-40-3]	tributyl phosphine				
	$\Delta_v H$	(353–428)	51.7 ± 0.5	390		[2001BAE]
C₁₂H₂₈N₂	[4843-89-4]	1,12-dodecanediamine				
	$\Delta_{\text{fus}} H$		67.1	341.8	DSC	[2006KHI/DAH2]
	$\Delta_{\text{fus}} H$		67.1	340.5	DSC	[2002DAL/DEL]
	$\Delta_v H$	(313–353)	110.1	328	A	[1987STE/MAL]
C₁₂H₂₈N₂	[60678-69-5]	tetrapropyl hydrazine				
	$\Delta_v H$	(362–423)	65.2	377	A	[1987STE/MAL]
C₁₂H₃₀N₃P	[2283-11-6]	<i>tris</i> (diethylamino)phosphine				
	$\Delta_v H$		60.7 ± 0.4			[1959FOL/MOR]
C₁₃H₄Cl₆O	[38178-99-3]	1,2,4,5,7,8-hexachloroxanthene				
	$\Delta_{\text{sub}} H$	(353–449)	147	401	T	[1986ROR]
C₁₃H₄N₄O₁₀	[185141-40-6]	2,3,5,7-tetranitroxanthone				
	$\Delta_{\text{fus}} H$		33.56	514	DSC	[1997IBR/FRA]
C₁₃H₄N₄O₁₀	[54849-77-3]	2,4,5,7-tetranitroxanthone				
	$\Delta_{\text{fus}} H$		32.2	593.9	DSC	[1997IBR/FRA]
C₁₃H₅N₃O₇	[129-79-3]	2,4,7-trinitrofluoren-9-one				
	$\Delta_{\text{fus}} H$		2.9	430.2		
	$\Delta_{\text{fus}} H$		23.5	449.2	DSC	[1980KRA/PIG]
C₁₃H₅N₃O₈	[185141-39-3]	1,2,7-trinitroxanthone				
	$\Delta_{\text{fus}} H$		11.89	554.9	DSC	[1997IBR/FRA]
		Note: Decomposes near melting point temperature.				
C₁₃H₅N₃O₈	[54849-76-2]	2,3,7-trinitroxanthone				
	$\Delta_{\text{fus}} H$		24.91	538.9	DSC	[1997IBR/FRA]
C₁₃H₅N₃O₈	[131032-92-3]	2,4,7-trinitroxanthone				
	$\Delta_{\text{fus}} H$		31.4	477.8	DSC	[1997IBR/FRA]
C₁₃H₆Cl₆O₂	[70-30-4]	2,2'-methylene <i>bis</i> (3,4,6-trichlorophenol)				
	$\Delta_{\text{fus}} H$		33.26	437.5	DSC	[1991ACR, 1990DON/DRE]
C₁₃H₆N₂O₆	[185141-35-9]	1,7-dinitroxanthone				
	$\Delta_{\text{fus}} H$		37.23	536.4	DSC	[1997IBR/FRA]
C₁₃H₆N₂O₆	[185141-37-1]	2,5-dinitroxanthone				
	$\Delta_{\text{fus}} H$		31.37	491.2	DSC	[1997IBR/FRA]
C₁₃H₆N₂O₆	[185141-38-2]	2,6-dinitroxanthone				
	$\Delta_{\text{fus}} H$		26.13	541	DSC	[1997IBR/FRA]
C₁₃H₆N₂O₆	[51792-18-8]	2,7-dinitroxanthone				
	$\Delta_{\text{fus}} H$		30.59	540	DSC	[1997IBR/FRA]
C₁₃H₇F₃N₂O₅	[15457-05-3]	2-nitro-1-(4-nitrophenoxy)-4-(trifluoromethyl)benzene				

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{fus}}H$		18.44	364.6	DSC	[1991ACR, 1990DON/DRE]
C ₁₃ H ₇ NO ₂	[46492-08-4] $\Delta_{\text{sub}}H$	benz[g]isoquinoline-5,10-dione (334–381)	108.1 ± 1.6	358	ME	[1998OJA/SUU]
C ₁₃ H ₇ NO ₄	[17607-01-1] $\Delta_{\text{fus}}H$	1-nitroxanthone	28.9	477.7	DSC	[1997IBR/FRA]
C ₁₃ H ₇ NO ₄	[20061-39-0] $\Delta_{\text{fus}}H$	2-nitroxanthone	26.75	477.9	DSC	[1997IBR/FRA]
C ₁₃ H ₇ NO ₄	[17607-10-2] $\Delta_{\text{fus}}H$	3-nitroxanthone	25.37	448	DSC	[1997IBR/FRA]
C ₁₃ H ₈ Br ₃ NO ₂	[87-10-5] $\Delta_{\text{fus}}H$	3,5-dibromo-N-(4-bromophenyl)-2-hydroxybenzamide	28.67	497.7	DSC	[1990DON/DRE]
C ₁₃ H ₈ Cl ₂ N ₂ O ₄	[50-65-7] $\Delta_{\text{fus}}H$ $\Delta_{\text{fus}}H$	5-chloro-N-(2-chloro-4-nitrophenyl)-2-hydroxybenzamide (niclosamide)	40.7 35.98	505.4 502.2	DSC	[2005YAN/DEV] [2004VAN/MAL]
C ₁₃ H ₈ Cl ₂ O	[90-98-2] $\Delta_{\text{fus}}H$ $\Delta_{\text{fus}}H$ $\Delta_{\text{fus}}H$ $\Delta_{\text{fus}}H$ $\Delta_{\text{fus}}H$ $\Delta_{\text{fus}}H$ $\Delta_{\text{fus}}H$	4,4'-dichlorobenzophenone (90–280) (90–280) (90–280) (10–298) (10–298) (10–298) (10–298)	0.04 0.05 NA 0.14 0.39 na 21.65	187 192 188.3 338.4	 AC AC DSC	 [2002DIA/LOP] [1999HUZ/SAI] [1990DON/DRE]
	$\Delta_{\text{fus}}H$ $\Delta_{\text{fus}}H$ $\Delta_{\text{fus}}H$ $\Delta_{\text{fus}}H$		0.15 0.25 NA 30.12	186.1 189.5 420	 DSC DSC	 [1987ECO/BER, 1999HUZ/SAI] [1991ACR, 1972PLA]
	$\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$	(349–367) (349–367)	114.5 ± 0.3 117.5 ± 0.3	358 298	ME ME	[2007RIB/AMA2] [2007RIB/AMA2]
C ₁₃ H ₈ F ₂ O ₃	[22494-42-4] $\Delta_{\text{fus}}H$ (I) $\Delta_{\text{fus}}H$ (II) $\Delta_{\text{fus}}H$ (III) $\Delta_{\text{sub}}H$	5-[2,4-difluorophenyl]salicylic acid (diflunisal) (349–414)	 35.9 35.8 35.9 119.3 ± 0.6	 486 485.5 486.4 	 DSC GS	 [2002PER/HAN] [2003PER/KUR]
C ₁₃ H ₈ N ₂ O ₂	[2538-68-3] $\Delta_{\text{fus}}H$	1-phenazinecarboxylic acid	35.44	514.5	DSC	[1997CIO/MEL]
C ₁₃ H ₈ N ₄	[19139-24-3] $\Delta_{\text{fus}}H$ $\Delta_{\text{fus}}H$ $\Delta_{\text{fus}}H$	8,8,9,9-tetracyanoquadracyclo[2.2.1.0 ^{3,5} .2]nonane	4.14 0.37 14.47	425.8 462.1 467.9	 DSC	 [1984WEI/LEF]
C ₁₃ H ₈ O	[548-39-0] $\Delta_{\text{sub}}H$	perinaphthenone (326–348)	97.2 ± 2.5	337	ME	[1998OJA/SUU]
C ₁₃ H ₈ O	[486-25-9] $\Delta_{\text{fus}}H$	9-fluorenone	14.85	353.3	DSC	[1998VER4]

Note: the entry for Ref. [1990DON/DRE] may likely be in error. The authors give the name of the compound as 4,4'-dichlorobenzophenone in the paper; however, they give the CAS Registry number of [85-29-0] which corresponds to 2,4'-dichlorobenzophenone. The observed melting point temperature of 338.4 K is more in line with the melting point temperature of 2,4'-dichlorobenzophenone

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound		T _m (K)	Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)			
	$\Delta_{\text{fus}}H$		18.12	356.4		[1991ACR]
	$\Delta_{\text{sub}}H$		93.9 ± 1.8	298	GS	[1998VER4]
	$\Delta_{\text{sub}}H$		87.6 ± 0.3	319	C	[1988SAB/ELW2]
	$\Delta_{\text{sub}}H$		88.4 ± 0.4	298	C	[1988SAB/ELW2]
	Δ_vH		60.9	435		[1983SIV/MAR]
	Δ_vH		59.8	475		[1983SIV/MAR]
	Δ_vH		59.1	525		[1983SIV/MAR]
	Δ_vH		58.6	565		[1983SIV/MAR]
	Δ_vH		57.9	595		[1983SIV/MAR]
C ₁₃ H ₈ OS	[492-22-8]	thioxanthone				
	$\Delta_{\text{fus}}H$		35.5	487.9		[1992SAB/ELW]
	$\Delta_{\text{sub}}H$		114.8 ± 0.4	298	C	[1992SAB/ELW]
C ₁₃ H ₈ O ₂	[90-47-1]	xanthone				
	$\Delta_{\text{fus}}H$		26.12	449.7		[1996DOM/HEA]
	$\Delta_{\text{sub}}H$		102.7 ± 2.3	298	C	[2009FRE/GOM2]
	$\Delta_{\text{sub}}H$		98.57 ± 0.4	298	C	[1988SAB/ELW]
C ₁₃ H ₈ O ₂	[5472-84-4]	3-hydroxy-1 <i>H</i> -phenalen-1-one (402–432)	151.5 ± 4.7	417	ME	[1998OJA/SUU]
C ₁₃ H ₉ ClO	[5162-03-8]	2-chlorobenzophenone				
	$\Delta_{\text{sub}}H$		100.2 ± 0.4	298	C	[2007RIB/AMA2]
C ₁₃ H ₉ ClO	[1016-78-0]	3-chlorobenzophenone (321–339)	108.8 ± 0.4	330	ME	[2007RIB/AMA2]
	$\Delta_{\text{sub}}H$	(321–339)	110.4 ± 0.4	298	ME	[2007RIB/AMA2]
C ₁₃ H ₉ ClO	[134-85-0]	4-chlorobenzophenone (320–338)	105.4 ± 0.3	329	ME	[2007RIB/AMA2]
	$\Delta_{\text{sub}}H$	(320–338)	108.2 ± 0.3	298	ME	[2007RIB/AMA2]
C ₁₃ H ₉ ClO ₂	[85-19-8]	5-chloro-2-hydroxybenzophenone (293–367)	91.9	308	A, UV	[1987STE/MAL, 1960SCH/HIR]
	Δ_vH	(367–493)	73.3	382	A, UV	[1987STE/MAL, 1960SCH/HIR]
C ₁₃ H ₉ Cl ₃ N ₂ O	[na]	benzoic acid, 2,4,6-trichlorophenyl hydrazide				
	$\Delta_{\text{fus}}H$		32.71	439.7	DSC	[1990DON/DRE]
C ₁₃ H ₉ Cl ₃ N ₂ O	[101-20-2]	3,4,4'-trichlorocarbanilide				
	$\Delta_{\text{us}}H$		6.1	428	DSC	[2010RIB/RIB2]
	$\Delta_{\text{sub}}H$		182.2 ± 1.7	298	C	[2010RIB/RIB2]
C ₁₃ H ₉ F ₃ N ₂ O ₂	[4394-00-7]	2-[3-(trifluoromethyl)anilino]nicotinic acid (niflumic acid)				
	$\Delta_{\text{fus}}H$		36.5	478	DSC	[2007PER/SUR2, 2009SUR/TER]
	$\Delta_{\text{fus}}H$		35.7	476.4	DSC	[2004ROM/BUS]
	$\Delta_{\text{fus}}H$		32.73	477.2		[1998BUS/PEN]
	$\Delta_{\text{fus}}H$		38.0	476		[1989PIN/GON]
	$\Delta_{\text{sub}}H$	(355–396)	127.8 ± 0.8	376	GS	[2007PER/SUR2, 2009SUR/TER]
	$\Delta_{\text{sub}}H$	(355–396)	130.2 ± 0.8	298	GS	[2007PER/SUR2, 2009SUR/TER]
	Δ_vH		107.5	298	S-F	[2007PER/SUR2]
C ₁₃ H ₉ N	[260-94-6]	acridine				

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{fus}}H$		18.58	383.2	DSC	[1975MCE/SAN]
	$\Delta_{\text{sub}}H$		86.0	430	TGA	[1998LEB/CHI]
	$\Delta_{\text{sub}}H$		89.5 ± 0.2	333	C	[1994SAB/TAB2]
	$\Delta_{\text{sub}}H$		91.7 ± 0.4	298	C	[1994SAB/TAB2]
	$\Delta_{\text{sub}}H$		94.5	298		[1989STE/CHI]
	$\Delta_{\text{sub}}H$	(280–328)	92.6	295		[1987STE/MAL]
	$\Delta_{\text{sub}}H$	(303–328)	90.8 ± 1.3	298	TE	[1975DEK/VAN]
	$\Delta_{\text{sub}}H$	(303–326)	93.3 ± 0.8	298	TCM	[UR/DEK, 1975DEK/VAN]
	$\Delta_{\text{sub}}H$	(281–298)	91.6 ± 2.5	290	LE	[1975MCE/SAN]
	$\Delta_{\text{sub}}H$	(306–345)	92.8 ± 1.3	298	ME	[UR/DEK, 1975DEK/VAN]
	$\Delta_{\text{sub}}H$		78.7		E	[1946ALB/WIL]
	Δ_vH		72.1		GC	[1996GOV/RUT]
	Δ_vH	(383–637)	71.5 ± 0.2	400	IPM,EB	[1989STE/CHI]
	Δ_vH	(383–637)	68.9 ± 0.1	440	IPM,EB	[1989STE/CHI]
	Δ_vH	(383–637)	66.4 ± 0.1	480	IPM,EB	[1989STE/CHI]
	Δ_vH	(383–637)	63.8 ± 0.1	520	IPM,EB	[1989STE/CHI]
	Δ_vH	(383–637)	61.3 ± 0.2	560	IPM,EB	[1989STE/CHI]
	Δ_vH	(423–621)	62.9	465		[1983SIV/KOB]
	Δ_vH	(423–621)	62.1	515		[1983SIV/KOB]
	Δ_vH	(423–621)	61.5	595		[1983SIV/KOB]
	Δ_vH	(402–619)	66.2	417	A	[1987STE/MAL, 1947STU]
C₁₃H₉N	[260-27-3]	3,4-benzoquinoline (phenanthridine)				
	$\Delta_{\text{fus}}H$		0.02	354		
	$\Delta_{\text{fus}}H$		22.83	379.7		[1996DOM/HEA]
	$\Delta_{\text{sub}}H$	(288–323)	100.1 ± 10.1	306	ME	[1998OJA/SUU]
	$\Delta_{\text{sub}}H$		98.6	298		[1989STE/CHI]
	$\Delta_{\text{sub}}H$	(288–323)	94.6 ± 4	308	ME	[1975MCE/INI, 1987STE/MAL]
	$\Delta_{\text{sub}}H$		107.5		ME	[1965DAV/KYB]
	Δ_vH	(383–473)	74.3 ± 0.1	380	IPM	[1989STE/CHI]
	Δ_vH	(383–473)	71.6 ± 0.1	420	IPM	[1989STE/CHI]
	Δ_vH	(383–473)	68.9 ± 0.1	460	IPM	[1989STE/CHI]
C₁₃H₉N	[85-02-9]	5,6-benzoquinoline				
	$\Delta_{\text{sub}}H$	(288–323)	83.1 ± 3.6	308	ME	[1975MCE/INI, 1987STE/MAL]
	$\Delta_{\text{sub}}H$		106.3		ME	[1972MIL]
C₁₃H₉N	[230-27-3]	7,8-benzoquinoline				
	$\Delta_{\text{fus}}H$		14.1	324.1		[1996DOM/HEA]
	$\Delta_{\text{sub}}H$		90.2 ± 2.0	298		[1989STE/CHI]
	$\Delta_{\text{sub}}H$	(293–323)	80.8 ± 2.5	308	ME	[1975MCE/INI, 1987STE/MAL]
	$\Delta_{\text{sub}}H$		100.4		ME	[1972MIL]
	Δ_vH		71.4		GC	[1996GOV/RUT]
	Δ_vH	(373–672)	71.7 ± 0.1	380	IPM,EB	[1989STE/CHI]
	Δ_vH	(373–672)	69.0 ± 0.1	420	IPM,EB	[1989STE/CHI]
	Δ_vH	(373–672)	66.5 ± 0.1	460	IPM,EB	[1989STE/CHI]
	Δ_vH	(373–672)	64.0 ± 0.1	500	IPM,EB	[1989STE/CHI]
	Δ_vH	(373–672)	61.5 ± 0.3	540	IPM,EB	[1989STE/CHI]
	Δ_vH	(373–672)	59.0 ± 0.3	580	IPM,EB	[1989STE/CHI]
C₁₃H₉NO	[598-95-0]	acridone				
	$\Delta_{\text{fus}}H$		32.5	640	DSC	[2003STO/KRZ]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{sub}}H$		133		DSC	[2003STO/KRZ]
	$\Delta_{\text{sub}}H$		136.2 ± 0.5	298	C	[1992SAB/ELW]
C ₁₃ H ₉ NO ₂	[2382-08-3]	N-methyl-1,8-naphthalimide				
	$\Delta_{\text{sub}}H$	(379–398)	107.4 ± 0.8	389	ME	[2000ROU/JIM]
	$\Delta_{\text{sub}}H$		109.7 ± 0.8	298	ME	[2000ROU/JIM]
C ₁₃ H ₉ NO ₂	[607-57-8]	2-nitrofluorene				
	$\Delta_{\text{sub}}H$	(349–384)	114.2 ± 3.0		ME	[2008GOL/SUU]
C ₁₃ H ₉ NO ₄	[75965-74-1]	2-nitro-7-methoxynaphtho[2,1b]furan				
	$\Delta_{\text{fus}}H$		28.7	460.4	DSC	[2010KES/AUC]
C ₁₃ H ₉ N ₂	[716-79-0]	2-phenylbenzimidazole				
	$\Delta_{\text{fus}}H$		22.18	572.2		[1971KAM/MIT]
C ₁₃ H ₁₀	[86-73-7]	fluorene				
	$\Delta_{\text{fus}}H$		19.1	387.7	DSC	[2000LIS/JAM]
	$\Delta_{\text{fus}}H$		19.58	387.9		[1996DOM/HEA]
	$\Delta_{\text{sub}}H$	(289–359)	86.1 ± 0.1	298	GS	[2004VER]
	$\Delta_{\text{sub}}H$		87.6	298	CGC-DSC	[1998CHI/HES]
	$\Delta_{\text{sub}}H$	(313–453)	84.9	383	GS	[1995NAS/LEN]
	$\Delta_{\text{sub}}H$	(323–363)	84.9 ± 0.4	343	GS	[1994RAK/VER2]
	$\Delta_{\text{sub}}H$		85.1 ± 0.4	298		[1994RAK/VER2]
	$\Delta_{\text{sub}}H$	(318–333)	87.0 ± 1.0	318	PG	[1988SAS/JOS]
	$\Delta_{\text{sub}}H$		80.2 ± 0.2	298	C	[1987SAB/ANT]
	$\Delta_{\text{sub}}H$	(348–388)	78.9	363	A	[1987STE/MAL]
	$\Delta_{\text{sub}}H$	(308–347)	83.2	328	GS	[1986SAT/INO]
	$\Delta_{\text{sub}}H$	(298–343)	92.2	320	T	[1986HAN/ECK]
	$\Delta_{\text{sub}}H$	(283–323)	88.4 ± 0.6	303	GS	[1983SON/ZOL]
	$\Delta_{\text{sub}}H$	(350–388)	83.1 ± 1.3			[1977FIN/MES, 1975OSB/DOU]
	$\Delta_{\text{sub}}H$		81.8	388	B	[1975OSB/DOU]
	$\Delta_{\text{sub}}H$	(286–300)	80.3 ± 0.8	293	TE	[1960BUD]
	$\Delta_{\text{sub}}H$	(306–323)	82.8	315		[1953BRA/CLE, 1987STE/MAL]
	$\Delta_{\text{sub}}H$	(306–322)	82.8			[1953BRA/CLE2, 1960JON]
	Δ_vH		72.4 ± 1.7	298	CGC	[2008HAN/NUT]
	Δ_vH	(373–423)	74.4 ± 1.2	298	GC	[2006HAF/PAR]
	Δ_vH	(323–473)	66.9	398	GC	[2002LEI/CHA]
	Δ_vH		72.3	298	CGC	[1998CHI/HES]
	Δ_vH	(403–453)	72.2	298	CGC	[1995CHI/HOS]
	Δ_vH	(323–363)	65.7	298	B	[1994RAK/VER2]
	Δ_vH	(383–427)	63.3	398		[1988SAS/JOS]
	Δ_vH	(402–568)	54.2	417	A	[1987STE/MAL]
	Δ_vH	(423–573)	56.6	498	I	[1923MOR/MUR]
C ₁₃ H ₁₀ BrCl ₂ O ₂ PS	[21609-90-5]	O-(4-bromo-2,5-dichlorophenyl) O-methyl phenylphosphonothioate				
	$\Delta_{\text{fus}}H$		31.35	345.6	DSC	[1990DON/DRE]
C ₁₃ H ₁₀ BrN ₃ O ₄	[192219-62-8]	2-cyano-6-nitro-1(2H)-quinolinecarboxylic acid, 2-bromoethyl ester				
	$\Delta_{\text{fus}}H$		19.79	419.2	DSC	[2005LIZ/ZAB]
C ₁₃ H ₁₀ ClN ₃ O ₄	[850836-65-6]	2-cyano-6-nitro-1(2H)-quinolinecarboxylic acid, 2-chloroethyl ester				
	$\Delta_{\text{fus}}H$		17.17	418.9	DSC	[2005LIZ/ZAB]
C ₁₃ H ₁₀ Cl ₂ S	[na]	<i>p</i> -chlorobenzyl <i>p</i> -chlorophenyl sulfide				
	$\Delta_{\text{fus}}H$		32.22	343.8	DSC	[1969PLA/GLA]
C ₁₃ H ₁₀ N ₂	[622-16-2]	N,N'-diphenylcarbodiimide				

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{fus}}H$		18.55	287.4		[1990DOM/HEA]
	Δ_vH	(500–599)	65.6	515	A, I	[1987STE/MAL, 1962JOH/MCE]
C ₁₃ H ₁₀ N ₂	[90-45-9]	9-aminoacridine				
	$\Delta_{\text{sub}}H$		115	520	TGA	[1998STO/KRZ]
C ₁₃ H ₁₀ N ₂	[716-79-0]	2-phenylbenzimidazole				
	$\Delta_{\text{sub}}H$		123.0 ± 1.7	298	C	[2005RIB/RIB]
C ₁₃ H ₁₀ N ₂ O ₂	[785-80-8]	N-phenyl 4-nitrobenzaldehyde imine				
	$\Delta_{\text{fus}}H$		24.56	347.2		[1997VER/MOR]
	$\Delta_{\text{sub}}H$		126 ± 1.3	298		[1997VER/MOR]
C ₁₃ H ₁₀ N ₂ O ₄	[50-35-1]	thalidomide				
	$\Delta_{\text{fus}}H$ (I)		39.97	546.7	DSC	[2007LAR/PER]
	$\Delta_{\text{fus}}H$ (II)		37.91	550.8	DSC	[2007LAR/PER]
	$\Delta_{\text{fus}}H$		36.02	548.2		[2002GOO/LAI]
C ₁₃ H ₁₀ N ₄	[7477-73-8]	1,5-diphenyltetrazole				
	$\Delta_{\text{sub}}H$	(348–363)	121.5 ± 4.2	355	ME	[1951MCE/RIG, 1970COX/PIL]
C ₁₃ H ₁₀ N ₄	[18038-45-7]	2,5-diphenyltetrazole				
	$\Delta_{\text{sub}}H$	(333–353)	119.7 ± 4.2	343	ME	[1951MCE/RIG, 1970COX/PIL]
C ₁₃ H ₁₀ N ₄ O	[14031-13-1]	1-phenazinecarboxylic acid hydrazide				
	$\Delta_{\text{fus}}H$		27.62	505	DSC	[1997CIO/MEL]
C ₁₃ H ₁₀ O	[119-61-9]	benzophenone				
	$\Delta_{\text{fus}}H$		18.47	321.3	AC	[2002HAN/HIK]
	$\Delta_{\text{fus}}H$	(5–440)	18.81	321.2	AC	[2002CHI/KN12]
	$\Delta_{\text{fus}}H$		18.19	324.2		[1996DOM/HEA]
	$\Delta_{\text{sub}}H$		93.1 ± 2.1	298	GS	[1998VER4]
	$\Delta_{\text{sub}}H$		94.7 ± 1	321	DM	[1983DEK/VAN]
	$\Delta_{\text{sub}}H$		92 ± 0.83	298	C	[1974SAB, 1983DEK/VAN]
	$\Delta_{\text{sub}}H$	(295–313)	95.0 ± 0.2	304	ME	[1980COL/JIM2]
	$\Delta_{\text{sub}}H$		84.4 ± 1.13	298	C	[1978SAB/LAF2]
	$\Delta_{\text{sub}}H$	(297–317)	93.9 ± 0.5	307	TE,ME	[1977DEK/VAN]
	$\Delta_{\text{sub}}H$	(293–318)	95.0 ± 1.5	305	TE	[1975DEK/VAN]
	$\Delta_{\text{sub}}H$	(294–318)	92.9 ± 0.8	306	ME	[1975VAN/DEK]
	$\Delta_{\text{sub}}H$	(278–311)	77.0 ± 2.5	298	ME	[1974ARS]
	$\Delta_{\text{sub}}H$	(298–318)	89.96	308	ME	[1987STE/MAL, 1974PRI/POU]
	$\Delta_{\text{sub}}H$	(295–304)	94.6 ± 0.8	298	TCM	[1973DEK/OON]
	$\Delta_{\text{sub}}H$		93.4 ± 0.3	298	C	[1972MOR3]
	$\Delta_{\text{sub}}H$	(293–319)	96.1	306		[1956SER/VOI]
	$\Delta_{\text{sub}}H$		91.2			[1950NIT/SEK]
	$\Delta_{\text{sub}}H$	(290–315)	78.2 ± 1.2	303		[1938WOL/WEG, 1934WOL/TR1]
	$\Delta_{\text{sub}}H$		95 ± 2.5	298	TE	[1932NEU/VOL, 1970COX/PIL, 1960JON]
	$\Delta_{\text{sub}}H$	(273–320)	91.2 ± 1.6	298	ME	[1925VOL/KIR]
	Δ_vH	(433–673)	65.1	448	A	[1987STE/MAL]
	Δ_vH	(473–579)	62.2	488		[1949DRE/SHR, 1949DRE/MAR, 1984BOU/FRI]
	Δ_vH	(530–575)	59.0	545		[1904JAQ/WAS, 1984BOU/FRI]
C ₁₃ H ₁₀ O	[92-83-1]	xanthene				
	$\Delta_{\text{fus}}H$		20.67	374.6	DSC	[2008MON/SAN]
	$\Delta_{\text{fus}}H$		15.87	374.3	DSC	[2000MAH/SOL]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{fus}}H$		19.2	373.7		[1991ACR]
	$\Delta_{\text{sub}}H$	(305–353)	92.5	329	T	[1986ROR]
	$\Delta_{\text{sub}}H$		112.1 ± 2.1			[1958CAS/FLE3, 1970COX/PIL]
		The authors refer to the compound as dibenzopyran but the melting temperature corresponds to xanthene				
	Δ_vH	(424–589)	64.5	435		[1984SIV/KOB]
	Δ_vH	(424–589)	61.1	475		[1984SIV/KOB]
	Δ_vH	(424–589)	59.2	515		[1984SIV/KOB]
	Δ_vH	(424–589)	56.7	555		[1984SIV/KOB]
	Δ_vH	(424–589)	54.4	585		[1984SIV/KOB]
	Δ_vH	(413–433)	88.7	423	A	[1987STE/MAL, 1958CAS/FLE3]
C ₁₃ H ₁₀ O	[1689-64-1]	9-hydroxyfluorene				
	Δ_vH		50.4	435		[1983SIV/MAR]
	Δ_vH		49.7	465		[1983SIV/MAR]
	Δ_vH		48.9	505		[1983SIV/MAR]
C ₁₃ H ₁₀ O ₂	[947-84-2]	2-biphenylcarboxylic acid				
	$\Delta_{\text{sub}}H$		121.3 ± 4.3	298	C	[2004MAT/MIR2]
C ₁₃ H ₁₀ O ₂	[92-92-2]	4-biphenylcarboxylic acid				
	$\Delta_{\text{sub}}H$		127.5 ± 4.1	298	C	[2004MAT/MIR2]
C ₁₃ H ₁₀ O ₂	[93-99-2]	phenyl benzoate				
	$\Delta_{\text{sub}}H$		99.0 ± 0.4	298		[1971CAR/FIN]
	$\Delta_{\text{sub}}H$		89.5 ± 4.2			[1971KIP/RAB, 1977PED/RYL]
	$\Delta_{\text{sub}}H$		96.2 ± 1.7			[1947STU, 1970COX/PIL]
	Δ_vH	(379–587)	62.4	394	A	[1987STE/MAL, 1947STU]
C ₁₃ H ₁₀ O ₂	[117-99-7]	(2-hydroxyphenyl)phenylmethanone				
	$\Delta_{\text{fus}}H$		0.67	308.2	DTA	[1989SAL/ABA]
		Note: Reported enthalpy of fusion is too small, and the published enthalpy and entropy of fusion data are internally inconsistent				
C ₁₃ H ₁₀ O ₃	[835-11-0]	2,2'-dihydroxybenzophenone				
	$\Delta_{\text{fus}}H$		20.07	334.5		[2005TOM/MIZ]
C ₁₃ H ₁₀ O ₃	[131-56-6]	2,4-dihydroxybenzophenone				
	$\Delta_{\text{sub}}H$	(312–353)	134	327	A	[1987STE/MAL]
	Δ_vH	(418–485)	87.1	433	A, UV	[1987STE/MAL, 1960SCH/HIR]
C ₁₃ H ₁₀ O ₃	[118-55-8]	phenyl salicylate				
	$\Delta_{\text{fus}}H$		19.2	315.1	DSC	[2010LAZ/RIE]
	$\Delta_{\text{fus}}H$		18.4 ± 0.5	312.7	DSC	[2006PER/CON]
	$\Delta_{\text{fus}}H$ (I)		16.5	304.2	DSC	[2004RAM/COR]
	$\Delta_{\text{fus}}H$ (II)		18.6	315.2	DSC	[2004RAM/COR]
	$\Delta_{\text{fus}}H$		19.16	315		[2002HAN/HIK]
	$\Delta_{\text{fus}}H$		18.98	314.2	DSC	[1995MUR/PAI]
	$\Delta_{\text{sub}}H$	(279–315)	109.1	294	A	[1987STE/MAL]
	$\Delta_{\text{sub}}H$		92 ± 4.2			[1947STU, 1970COX/PIL]
	Δ_vH	(423–587)	69.9	438	A, UV	[1987STE/MAL, 1960SCH/HIR]
C ₁₃ H ₁₀ O ₃	[102-09-0]	diphenyl carbonate				
	$\Delta_{\text{fus}}H$		23.43	355		[1971CAR/FIN]
	$\Delta_{\text{sub}}H$		90 ± 8.4	298	E	[1971CAR/FIN, 1977PED/RYL]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(355–381)	80.9 ± 0.6	298	GS	[2008VER/EME2]
C ₁₃ H ₁₀ O ₄	[1470-79-7]	2,4,4'-trihydroxybenzophenone				
	$\Delta_{\text{fus}} H$		31.3	482.6	DSC	[1999PRI/HAWN]
	$\Delta_{\text{sub}} H$		139		TGA	[1999PRI/HAWN]
C ₁₃ H ₁₀ O ₅	[131-55-5]	2,2',4,4'-tetrahydroxybenzophenone				
	$\Delta_{\text{fus}} H$		28.0	472	DSC	[1999PRI/HAWN]
	$\Delta_{\text{sub}} H$		178.5		B	[1999PRI/HAWN]
	$\Delta_{\text{sub}} H$	(363–471)	143.4	378	A	[1987STE/MAL]
C ₁₃ H ₁₀ S	[261-31-4]	thioxanthene				
	$\Delta_{\text{fus}} H$		26.1	401.8		[1991ACR]
	$\Delta_{\text{sub}} H$		101.7 ± 1.6	298	C	[2009FRE/MON]
	$\Delta_{\text{sub}} H$	(339–402)	98.4 ± 0.2	370	ME	[2009FRE/MON]
	$\Delta_{\text{sub}} H$	(339–402)	100.9 ± 0.2	298	ME	[2009FRE/MON]
	$\Delta_v H$	(383–447)	69.5 ± 0.2	415	ME	[2009FRE/MON]
	$\Delta_v H$	(383–447)	77.8 ± 2.6	298	ME	[2009FRE/MON]
C ₁₃ H ₁₀ S	[7372-88-5]	4-methyldibenzothiophene				
	$\Delta_{\text{sub}} H$		90.3 ± 0.7	298	C	[2010FRE/GOM]
C ₁₃ H ₁₁ BrO ₅	[111171-29-0]	8-(hydroxymethyl)-6-bromo-2-oxo-2H-1-benzopyran-3-carboxylic acid, ethyl ester				
	$\Delta_{\text{fus}} H$		28.19	434.1	DSC	[1992HUA/ZHO2]
C ₁₃ H ₁₁ Cl	[90-99-3]	chlorodiphenylmethane				
	$\Delta_v H$	(381–450)	70.4	396	A	[1987STE/MAL]
C ₁₃ H ₁₁ ClO ₅	[111171-28-9]	8-(hydroxymethyl)-6-chloro-2-oxo-2H-1-benzopyran-3-carboxylic acid, ethyl ester				
	$\Delta_{\text{fus}} H$		15.53	418.2		
	$\Delta_{\text{fus}} H$		22.47	424	DSC	[1992HUA/ZHO2]
C ₁₃ H ₁₁ F	[579-55-5]	fluorodiphenylmethane				
	$\Delta_v H$	(288–333)	69.8 ± 0.4	298	GS	[1997SCH/VER]
C ₁₃ H ₁₁ N	[1013-88-3]	benzophenone imine				
	$\Delta_v H$	(308–338)	74.2 ± 1.0	323	GS	[1997VER/MOR]
	$\Delta_v H$	(308–338)	75.7 ± 1.0	298	GS	[1997VER/MOR]
	$\Delta_v H$	(373–422)	62.3	388	A	[1987STE/MAL]
C ₁₃ H ₁₁ N	[538-51-2]	N-phenyl-benzaldehyde imine				
	$\Delta_{\text{fus}} H$		20.42	329.7		[1997VER/MOR]
	$\Delta_{\text{sub}} H$	(294–326)	97.4 ± 1.2	309	T	[1997VER/MOR]
	$\Delta_{\text{sub}} H$		98.1 ± 1.2	298	T	[1997VER/MOR]
	$\Delta_{\text{sub}} H$		93.7 ± 0.9	298	C	[1986KIR/ACR]
	$\Delta_{\text{sub}} H$		85.5 ± 2.1	293	EST	[1948COA/SUT]
C ₁₃ H ₁₁ N	[1484-12-4]	9-methylcarbazole				
	$\Delta_{\text{fus}} H$		17.15	362.5		[1996DOM/HEA]
	$\Delta_{\text{sub}} H$	(313–332)	95.0	322	ME	[1990JIM/ROU]
	$\Delta_{\text{sub}} H$		95.5	298		[1990JIM/ROU]
	$\Delta_v H$		79.5 ± 3.2	298	CGC	[2009LIP/CHI, 2009LIP/HAN]
	$\Delta_v H$	(373–673)	73.4	400	EB,IP	[1992STE/CHI]
	$\Delta_v H$	(373–673)	70.5	440	EB,IP	[1992STE/CHI]
	$\Delta_v H$	(373–673)	67.7	480	EB,IP	[1992STE/CHI]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(373–673)	65.0	520	EB,IP	[1992STE/CHI]
	$\Delta_v H$	(373–673)	62.1	560	EB,IP	[1992STE/CHI]
	$\Delta_v H$	(373–673)	59.1	600	EB,IP	[1992STE/CHI]
	$\Delta_v H$	(373–673)	55.9	640	EB,IP	[1992STE/CHI]
	$\Delta_v H$	(348–384)	74.9	366	GS	[1980VAN/PRA]
C ₁₃ H ₁₁ N	[5097-92-7]	<i>cis</i> 4-(2-phenylethylenyl)pyridine				
	$\Delta_{\text{fus}}H$		9.14	400.3	DSC	[2007LIU/LIU]
C ₁₃ H ₁₁ NO	[1137-98-8]	N-phenylmethylene benzenamine N-oxide				
	$\Delta_{\text{sub}}H$		115.0 ± 0.8	298	C	[1986KIR/ACR]
C ₁₃ H ₁₁ NO	[779-84-0]	2-hydroxybenzaldehyde N-phenylimine				
	$\Delta_{\text{sub}}H$	(288–325)	115.9	303	A	[1987STE/MAL]
	$\Delta_{\text{sub}}H$	(348–408)	129.9	378		[1958HOY/PEP]
C ₁₃ H ₁₁ NO	[1689-73-2]	4-hydroxybenzaldehyde N-phenylimine				
	$\Delta_{\text{sub}}H$	(348–408)	127.9	363	A	[1987STE/MAL]
	$\Delta_{\text{sub}}H$	(288–338)	116	313		[1958HOY/PEP]
C ₁₃ H ₁₁ NO	[93-98-1]	benzanilide				
	$\Delta_{\text{fus}}H$		32.4	436.3	DSC	[2006MAT/MIR2]
	$\Delta_{\text{fus}}H$		29.61	436.5		[1996DOM/HEA]
	$\Delta_{\text{sub}}H$		125.4 ± 2.3	298	C	[2006MAT/MIR2]
	$\Delta_{\text{sub}}H$	(352–369)	99.2	360.5	A	[1987STE/MAL, 1960AIH2]
C ₁₃ H ₁₁ NO ₂	[20357-59-9]	N-(2-hydroxyphenylmethylene) benzenamine N-oxide				
	$\Delta_{\text{sub}}H$		116.5 ± 1.4	298	C	[1986KIR/ACR]
C ₁₃ H ₁₁ NO ₂	[91-40-7]	N-phenylanthranilic acid				
	$\Delta_{\text{fus}}H$		39.7	458.2	DSC	[2009SUR/TER]
	$\Delta_{\text{sub}}H$	(353–411)	123.0 ± 1.3	382	GS	[2009SUR/TER]
	$\Delta_{\text{sub}}H$	(353–411)	126.0 ± 1.3	298	GS	[2009SUR/TER]
C ₁₃ H ₁₁ NO ₅	[14698-29-4]	1-ethyl-1,4-dihydro-6,7-methylenedioxy-4-oxo-3-quinoline-carboxylic acid (oxolinic acid)				
	$\Delta_{\text{fus}}H$		43.59	592.5	DSC	[2004ROM/BUS2]
C ₁₃ H ₁₁ N ₃ O	[2440-22-4]	2-(2'-hydroxy-5'-methylphenyl)benzotriazole				
	$\Delta_{\text{sub}}H$	(293–333)	125.2	308	A	[1987STE/MAL]
	$\Delta_v H$	(413–433)	79.1	423	ME	[1984SUR]
	$\Delta_v H$	(404–435)	70.6	419	A, UV	[1987STE/MAL, 1960SCH/HIR]
C ₁₃ H ₁₁ N ₃ O ₂ S	[na]	5-methyl-2-[(4-methyl-2-nitrophenyl)amino]-3-thiophene carbonitrile				
	$\Delta_{\text{fus}}H$		24.57	400.2		[2001HE/GRI]
C ₁₃ H ₁₁ N ₃ O ₄	[1979-00-7]	2-cyano-6-nitro-1(2 <i>H</i>)-quinolinecarboxylic acid, ethyl ester				
	$\Delta_{\text{fus}}H$		27.11	391.2	DSC	[2005LIZ/ZAB]
C ₁₃ H ₁₂	[643-93-6]	3-methylbiphenyl				
	$\Delta_v H$	(283–463)	69.6	298		[1993KAS/MOK]
C ₁₃ H ₁₂	[644-08-6]	4-methylbiphenyl				
	$\Delta_{\text{sub}}H$		80.2 ± 1.4	298	C	[1997RIB/MAT4]
C ₁₃ H ₁₂	[101-81-5]	diphenylmethane				
	$\Delta_{\text{fus}}H$		19.01	298.4		[2005CHI/STE2]
	$\Delta_{\text{fus}}H$		18.58	298.3	AC	[1996DOM/HEA]
	$\Delta_{\text{sub}}H$	(273–295)	88.5 ± 0.8	284	GS	[1999VER5]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{sub}}H$		87.6 ± 0.8	298		[1999VER5]
	$\Delta_{\text{sub}}H$	(273–298)	71.5	286	EM	[1989SAS/NGU]
	$\Delta_{\text{sub}}H$	(276–295)	83.3 ± 3.3	286	HSA	[1986CHI/ANN]
	$\Delta_{\text{sub}}H$		82.4 ± 8		V	[1959AIH, 1970COX/PIL]
	$\Delta_{\text{sub}}H$	(278–299)	64.0			[1951BRI, 1960JON]
	$\Delta_{\text{sub}}H$		72.0 ± 0.8	297		[1938WOL/WEG]
	Δ_vH		67.6 ± 0.2	298		[2005CHI/STE2, 2008HAN/NUT]
	Δ_vH	(343–393)	64.7 ± 0.2	298		[2006HAF/PAR]
	Δ_vH	(330–588)	64.1 ± 0.1	340	IP,EB	[2005CHI/STE2]
	Δ_vH	(330–588)	61.0 ± 0.1	380	IP,EB	[2005CHI/STE2]
	Δ_vH	(330–588)	57.9 ± 0.1	420	IP,EB	[2005CHI/STE2]
	Δ_vH	(330–588)	55.0 ± 0.1	460	IP,EB	[2005CHI/STE2]
	Δ_vH	(330–588)	52.0 ± 0.2	500	IP,EB	[2005CHI/STE2]
	Δ_vH	(330–588)	48.9 ± 0.3	540	IP,EB	[2005CHI/STE2]
	Δ_vH		65.7	298	GC	[2002VAN/PAR]
	Δ_vH	(303–343)	66.4 ± 0.5	323	GS	[1999VER5]
	Δ_vH	(303–343)	67.9 ± 0.5	298	GS	[1999VER5]
	Δ_vH	(353–433)	61.8	368		[1990SOH/OKA]
	Δ_vH	(303–402)	63.7	363		[1989SAS/NGU]
	Δ_vH	(295–383)	72.2	310	A	[1987STE/MAL]
	Δ_vH	(423–583)	56.7	438	A	[1987STE/MAL]
	Δ_vH		55.8	445		[1981WIE/KOB]
	Δ_vH		49.0	535		[1981WIE/KOB]
	Δ_vH		66.6 ± 0.1	298	C	[1972MOR]
	Δ_vH	(490–555)	54.2	505		[1915CRA, 1984BOU/FRI]
C ₁₃ H ₁₂ ClN ₂ O ₂	[556836-77-2] $\Delta_{\text{fus}}H$	4-chloro-2'-hydroxy-4'-methoxyazobenzene	33.7	390	DSC	[2003PAJ/ROS]
C ₁₃ H ₁₂ ClN ₃ S	[281212-47-3] $\Delta_{\text{fus}}H$	N-2-(3-picoly)-N'-(2-chlorophenyl) thiourea	11.2	400.2	DSC	[2002KEL/SZC]
C ₁₃ H ₁₂ ClN ₃ S	[457886-93-0] $\Delta_{\text{fus}}H$	N-2-(4-picoly)-N'-(2-chlorophenyl) thiourea	44.5	441.2	DSC	[2002KEL/SZC]
C ₁₃ H ₁₂ ClN ₃ S	[457886-96-3] $\Delta_{\text{fus}}H$	N-2-(5-picoly)-N'-(2-chlorophenyl) thiourea	24.2	460.2	DSC	[2002KEL/SZC]
C ₁₃ H ₁₂ ClN ₃ S	[457886-94-1] $\Delta_{\text{fus}}H$	N-2-(6-picoly)-N'-(2-chlorophenyl) thiourea	27.3	449.7	DSC	[2002KEL/SZC]
C ₁₃ H ₁₂ ClN ₃ S	[53385-87-8] $\Delta_{\text{fus}}H$	N-2-(3-picoly)-N'-(4-chlorophenyl) thiourea	16.8	391.2	DSC	[2002SZC/KEL]
C ₁₃ H ₁₂ ClN ₃ S	[53385-88-9] $\Delta_{\text{fus}}H$	N-2-(4-picoly)-N'-(4-chlorophenyl) thiourea	35.2	460.2	DSC	[2002SZC/KEL]
C ₁₃ H ₁₂ ClN ₃ S	[53385-89-0] $\Delta_{\text{fus}}H$	N-2-(5-picoly)-N'-(4-chlorophenyl) thiourea	51.1	473.7	DSC	[2002SZC/KEL]
C ₁₃ H ₁₂ ClN ₃ S	[53385-90-3] $\Delta_{\text{fus}}H$	N-2-(6-picoly)-N'-(4-chlorophenyl) thiourea	40.1	464.2	DSC	[2002SZC/KEL]
C ₁₃ H ₁₂ N ₂ O	[102-07-8] $\Delta_{\text{fus}}H$	1,3-diphenylurea	34.62	512.1		[1996DOM/HEA, 1991ACR]
	$\Delta_{\text{fus}}H$		37.7	na	DSC	[1995STR/ARG]
	$\Delta_{\text{sub}}H$	(445–484)	152 ± 6		TE	[1987FER/DEL]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₃ H ₁₂ N ₂ O	[442-51-3] $\Delta_{\text{fus}}H$	7-methoxy-1-methyl-9H-pyrido[3,4-b]indole (harmine)				
			48.8	536.6	DSC	[1996BUR/DAG]
C ₁₃ H ₁₂ N ₂ O ₂	[23042-34-4] $\Delta_{\text{fus}}H$	N-methyl-N-(4-biphenyl)nitramine				
			24.0	415.1		[2002DAS/ZAL]
C ₁₃ H ₁₂ N ₂ O ₂	[17954-23-3] $\Delta_{\text{fus}}H$	2-cyano-1(2H)-quinolinecarboxylic acid, ethyl ester				
			19.88	337	DSC	[2005LIZ/ZAB]
C ₁₃ H ₁₂ N ₂ O ₅ S	[51803-78-2] $\Delta_{\text{fus}}H$	N-(4-nitro-2-phenoxyphenyl)methanesulfonamide (nimesulide)				
			37.3	422.5	DSC	[2007MON/PER]
C ₁₃ H ₁₂ N ₄ O ₂	[na] $\Delta_{\text{sub}}H$	4'-nitro-2-methylaminoazobenzene				
			134.7		GS	[1987SHI/OHK, 1991HOR]
C ₁₃ H ₁₂ O	[946-80-5] Δ_vH	benzyl phenyl ether (368–560)				
			58.8	383	A	[1987STE/MAL, 1947STU]
C ₁₃ H ₁₂ O	[91-01-0] $\Delta_{\text{fus}}H$	diphenylmethanol				
			23.0	338.5		[1996DOM/HEA]
		$\Delta_{\text{sub}}H$ (301–335)	105.7 ± 0.7	298	GS	[1998VER3]
		$\Delta_{\text{sub}}H$ (301–335)	104.5 ± 0.7	318	GS	[1998VER3]
		Δ_vH (342–373)	79.4 ± 0.7	358	GS	[1998VER5]
		Δ_vH (342–373)	83.0 ± 0.7	298	GS	[1998VER5]
		Δ_vH (438–574)	65.4	453	A	[1987STE/MAL]
C ₁₃ H ₁₂ O	[2876-63-3] Δ_vH	ethyl 1-naphthyl ketone (397–579)				
			74.1	412	A	[1987STE/MAL, 1947STU]
C ₁₃ H ₁₂ O	[28994-41-4] $\Delta_{\text{fus}}H$ (I) $\Delta_{\text{fus}}H$ (II) $\Delta_{\text{fus}}H$	2-(phenylmethyl)phenol (2-benzylphenol)				
			21.8	326.2		
			17.0	288.2	DSC	[2004ROM/ROC]
			23.4	325.7	DSC	[1995MUR/PAI]
C ₁₃ H ₁₂ O	[101-53-1] $\Delta_{\text{sub}}H$	4-benzylphenol (313–335)				
			97.4	324	A	[1987STE/MAL, 1960AIH]
C ₁₃ H ₁₂ O	[2928-43-0] $\Delta_{\text{fus}}H$ $\Delta_{\text{fus}}H$	2-biphenylmethanol				
			19.7	324	DSC	[2007PIN/BER, 2006DIO/PIN]
			18.5	326.8	DSC	[2006BAR/DAV]
		$\Delta_{\text{sub}}H$	106.0 ± 1.1	315	ME	[2007PIN/BER]
		$\Delta_{\text{sub}}H$	107.1 ± 1.1	298	ME	[2007PIN/BER]
		Δ_vH	85.6 ± 0.6	326	C	[2007PIN/BER]
C ₁₃ H ₁₂ O	[3597-91-9] $\Delta_{\text{fus}}H$	4-biphenylmethanol				
			27.0	375.5	DSC	[2007PIN/BER, 2006DIO/PIN]
		$\Delta_{\text{sub}}H$	105.7 ± 1.8	349	C	[2007PIN/BER]
		$\Delta_{\text{sub}}H$	107.3 ± 1.8	298	C	[2007PIN/BER]
C ₁₃ H ₁₂ OS	[40932-63-6] $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$	3-acetyl-2-methyl-5-phenylthiophene				
			(321–339) 107.3 ± 0.4	330	ME	[2010RIB/SAN]
			(321–339) 108.9 ± 0.4	298	ME	[2010RIB/SAN]
C ₁₃ H ₁₂ S	[831-91-4] $\Delta_{\text{sub}}H$	phenyl benzyl sulfide				
			98.4 ± 1.4	298	C	[2006MUL/MOZ]
C ₁₃ H ₁₃ BrS	[148681-80-5]	2-propyl-5-(4-bromophenyl)thiophene				

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{fus}}H$		15.7	360.4	DSC	[1993BRE/DUN]
C ₁₃ H ₁₃ ClN ₂ O ₂ S	[34392-72-8]	4-amino-N-(5-chloro-2-methylphenyl)benzenesulfonamide				
	$\Delta_{\text{fus}}H$		36.8	422.7	DSC	[2009PER/TKA]
	$\Delta_{\text{sub}}H$		130 ± 1	298	GS	[2009PER/TKA]
	Δ_vH		104	298	S-F	[2009PER/TKA]
C ₁₃ H ₁₃ N	[552-82-9]	N-methyldiphenylamine				
	Δ_vH	(376–555)	65.2	391	A	[1987STE/MAL, 1947STU]
C ₁₃ H ₁₃ N	[103-32-2]	N-benzylaniline				
	$\Delta_{\text{fus}}H$		16.76	305.6		[1991ACR]
	$\Delta_{\text{sub}}H$	(293–312)	103.6 ± 1.6	303	T	[1997VER]
	$\Delta_{\text{sub}}H$		51.3			[1980AIH]
	Δ_vH	(316–343)	79.6 ± 1.1	330		[1997VER]
	Δ_vH		79.5			[1980AIH]
Note: The value reported in Ref. [1980AIH] for the enthalpy of sublimation is smaller than the value given for the enthalpy of vaporization. The author of [1980AIH] noted the anomalous behavior.						
C ₁₃ H ₁₃ NO	[3449-48-7]	1-keto-1,2,3,4-tetrahydro-6-methylcarbazole				
	$\Delta_{\text{fus}}H$		26.9	468.5		[2006COR/LOP]
C ₁₃ H ₁₃ NO	[na]	2-(4-methoxyphenyl)-5-methylpyridine				
	$\Delta_{\text{fus}}H$		20.0	328		[2000MOR/HAR]
C ₁₃ H ₁₃ NO ₂	[na]	(dl) 2-(1-naphthoxy)propionamide				
	$\Delta_{\text{fus}}H$		37.66	445		[1976LEC/COL]
C ₁₃ H ₁₃ NO ₂	[na]	(d) 2-(1-naphthoxy)propionamide				
	$\Delta_{\text{fus}}H$		38.07	475		[1976LEC/COL]
C ₁₃ H ₁₃ N ₃ O	[na]	N-(4-methylphenyl)-N'-(2-pyridyl) urea				
	$\Delta_{\text{fus}}H$		204.45	447		[2002LU/SON, 2004SON/TAN]
Note: Value is too large. The compound may have decomposed upon melting, or there is a decimal place error in the numerical value.						
C ₁₃ H ₁₃ N ₃ S	[53385-83-4]	N-2-(6-picoly)-N'-phenylthiourea				
	$\Delta_{\text{fus}}H$		43.5	460.7	DSC	[2002VAL/HER]
C ₁₃ H ₁₃ OP	[2129-89-7]	methyldiphenylphosphine oxide				
	$\Delta_{\text{fus}}H$		20.37	385.4	DSC	[2010HU/WAN]
C ₁₃ H ₁₄	[2245-38-7]	1,6,7-trimethylnaphthalene				
	Δ_vH	(323–473)	68.6	398	GC	[2002LEI/CHA]
C ₁₃ H ₁₄	[6158-45-8]	1-isopropylnaphthalene				
	Δ_vH	(402–541)	50.4	417	A	[1987STE/MAL]
C ₁₃ H ₁₄	[2027-17-0]	2-isopropylnaphthalene				
	Δ_vH	(402–541)	60.3	417	A	[1987STE/MAL]
C ₁₃ H ₁₄ N ₂	[1208-52-2]	2,4'-diaminodiphenylmethane				
	Δ_vH	(353–403)	111.5	368	A	[1987STE/MAL]
C ₁₃ H ₁₄ N ₂	[101-77-9]	4,4'-diaminodiphenylmethane				
	$\Delta_{\text{fus}}H$		19.69	362.7	DSC	[2006KHI/DAH]
	$\Delta_{\text{fus}}H$		9.23	363.7		[1996DOM/HEA]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	Δ_vH	(343–393)	109.3	358	A	[1987STE/MAL]
	Δ_vH	(486–545)	98.0	501	A	[1987STE/MAL]
	Δ_vH	(471–545)	100.6	502	A	[1966ZAL/STR]
C ₁₃ H ₁₄ N ₂	[6582-52-1]	2,2'-diaminodiphenylmethane				
	$\Delta_{\text{sub}}H$	(343–403)	111.3	358	A	[1987STE/MAL]
C ₁₃ H ₁₄ N ₂ O ₃ S	[19837-74-2]	4-amino-N-(4-methoxyphenyl)benzenesulfonamide				
	$\Delta_{\text{fus}}H$		38.6	467.4	DSC	[2009PER/TKA]
	$\Delta_{\text{sub}}H$		124 ± 1	298	GS	[2009PER/TKA]
	Δ_vH		99.4	298	S-F	[2009PER/TKA]
C ₁₃ H ₁₅ Cl ₂ N ₃	[56518-41-3]	5-[(4-bromo-3,5-dimethoxyphenyl)methyl]-2,4-pyrimidinediamine				
	$\Delta_{\text{fus}}H$		49.9	505.4		[2007CAI/BET]
C ₁₃ H ₁₅ Cl ₂ N ₃	[66246-88-6]	1-[2-(2,4-dichlorophenyl)pentyl]-1 <i>H</i> -1,2,4-triazole (penconazole)				
	$\Delta_{\text{fus}}H$		33.58	332.4		[2004SUN/SON2]
C ₁₃ H ₁₅ Cl ₃ O ₃	[120-39-8]	2,4,6-trichlorophenoxyacetic acid, pentyl ester				
	Δ_vH	(460–573)	78.8	475	A,GC	[1987STE/MAL, 1966JEN/SCH]
C ₁₃ H ₁₅ N	[17058-12-7]	1,2,3,4-tetrahydro-9-methylcarbazole				
	$\Delta_{\text{us}}H$		0.08	162		
	$\Delta_{\text{fus}}H$		14.67	323.8		[1992STE/CHI]
	Δ_vH	(370–655)	72.5	400	EB,IP	[1992STE/CHI]
	Δ_vH	(370–655)	69.6	440	EB,IP	[1992STE/CHI]
	Δ_vH	(370–655)	66.7	480	EB,IP	[1992STE/CHI]
	Δ_vH	(370–655)	63.8	520	EB,IP	[1992STE/CHI]
	Δ_vH	(370–655)	60.7	560	EB,IP	[1992STE/CHI]
	Δ_vH	(370–655)	57.4	600	EB,IP	[1992STE/CHI]
	Δ_vH	(370–655)	53.8	640	EB,IP	[1992STE/CHI]
C ₁₃ H ₁₅ NO	[na]	1-(1-isocyanato-1-methylethyl)-4-(1-methylethylbenzene)				
	Δ_vH	(298–463)	68.5	308	DTA, T, HSA	[1986ACH/HAS]
C ₁₃ H ₁₅ NO ₂	[24691-76-7]	3,4-dihydro-6-methyl-2 <i>H</i> -pyran-5-carboxanilide				
	$\Delta_{\text{fus}}H$		19.21	381.1	DSC	[1990DON/DRE]
C ₁₃ H ₁₅ NO ₂	[na]	ethyl 4,7-dihydro-4,7-ethano-2 <i>H</i> -isoindole-1-carboxylate				
	$\Delta_{\text{fus}}H$		45.4	401.8		[2000UNO/ITO]
C ₁₃ H ₁₅ N ₃ O ₂	[87-47-8]	3-methyl-1-phenyl-1 <i>H</i> -pyrazol-5-yl dimethylcarbamate				
	$\Delta_{\text{fus}}H$		21.39	324.3	DSC	[1990DON/DRE]
C ₁₃ H ₁₅ N ₃ O ₈	[53848-89-8]	hexyl 2,4,6-trinitrobenzoate				
	$\Delta_{\text{us}}H$		1.7	264		
	$\Delta_{\text{fus}}H$		32.96	402	DSC	[1974WAR/WIL]
C ₁₃ H ₁₆ ClNO	[6740-88-1]	(RS)-2-(2-chlorophenyl)-2-(methylamino)cyclohexanone				
	$\Delta_{\text{fus}}H$		26.6	365.7		[2009TAM/MIR]
C ₁₃ H ₁₆ Cl ₂ O ₃	[67821-07-2]	2,4-dichlorophenoxyacetic acid, isopentyl ester				
	Δ_vH	(460–573)	75.8	475	A,GC	[1987STE/MAL, 1966JEN/SCH]
C ₁₃ H ₁₆ Cl ₂ O ₃	[1917-96-6]					
	Δ_vH	(444–573)	73.6	459	A,GC	[1987STE/MAL, 1966JEN/SCH]
C ₁₃ H ₁₆ F ₃ N ₃ O ₄	[na]	2,6-dinitro-N,N-dipropyl-4-(trifluoromethyl)benzenamine				

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{fus}}H$		22.32	321.4		[1991ACR]
C ₁₃ H ₁₆ F ₃ N ₃ O ₄	[na] $\Delta_{\text{fus}}H$	N-butyl-N-ethyl-2,6-dinitro-4-trifluoromethylaniline	36.5	338.5		[1991ACR]
C ₁₃ H ₁₆ N ₂	[5766-79-0] $\Delta_{\text{fus}}H$	2-phenyl-1-piperidinoacetonitrile	19.71	335.2		[1997WEL/VER]
	Δ_vH	(338–378)	73.2 ± 0.4	298	GS	[1997WEL/VER]
C ₁₃ H ₁₆ N ₂ O ₃	[37000-08-1] $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$	hexahydro-1-(3-nitrobenzoyl)-1 <i>H</i> -azepine (compound is called hexamethyleneimine <i>m</i> -nitro-benzoate in paper) (310–321)	113 104.6	315		[1972ROZ/POL] ME [1970POL/PER, 1972ROZ/POL]
C ₁₃ H ₁₆ N ₂ O ₆	[159432-36-7] $\Delta_{\text{fus}}H$	2,3-dihydro-7-(1-methylethoxy)-3-[2-(nitrooxy)ethyl]-4 <i>H</i> -1,3-benzoxazin-4-one	32.1	344.7	DSC	[1996FON/ROS]
C ₁₃ H ₁₇ ClN ₂ O ₄	[138517-06-3] $\Delta_{\text{fus}}H$	(4-nitrophenyl)-6-chlorohexyl carbamate	30.31	360.8	DSC	[1993TIE/FRA]
C ₁₃ H ₁₇ NO	[3626-62-8] Δ_vH Δ_vH	1-(phenacyl)piperidine (381–446) (382–450)	51.4 47.2	396 416	A	[1987STE/MAL, 1969DAV/MAK] [1969DAV/MAK]
C ₁₃ H ₁₇ NO	[13290-48-7] Δ_vH	1-(<i>m</i> -toluoyl)piperidine (373–403)	53.8	388	A	[1987STE/MAL, 1969DAV/MAK]
C ₁₃ H ₁₇ NO ₃	[4134-09-2] Δ_vH	(<i>dl</i>) N-acetylphenylalanine, ether ester (438–528)	82.4	453	A	[1987STE/MAL]
C ₁₃ H ₁₇ NO ₃	[na] $\Delta_{\text{sub}}H$	morpholine cinnamate (298–349)	118.8	313	A	[1987STE/MAL]
C ₁₃ H ₁₈	[941-60-6] $\Delta_{\text{fus}}H$ Δ_vH Δ_vH Δ_vH Δ_vH	1,1,4,6-tetramethylindane (313–383) (313–469) (423–469) (313–469)	15.74 59.4 60.2 51.9 61.4 ± 0.5	273.6 328 328 439 298		[1991ACR] A,EB,IP [1987STE/MAL, 1978OSB/SCO] A,EB,IP [1987STE/MAL, 1978OSB/SCO] A,EB,IP [1987STE/MAL, 1978OSB/SCO] EB, IP [1978OSB/SCO]
C ₁₃ H ₁₈	[1078-04-2] $\Delta_{\text{fus}}H$ Δ_vH Δ_vH Δ_vH Δ_vH	1,1,4,7-tetramethylindane (313–388) (313–469) (431–469) (313–469)	11.28 59.6 60.4 52.0 61.4 ± 0.6	245.6 328 328 446 298		[1991ACR] A,EB,IP [1987STE/MAL, 1978OSB/SCO] A,EB,IP [1987STE/MAL, 1978OSB/SCO] A,EB,IP [1987STE/MAL, 1978OSB/SCO] EB, IP [1978OSB/SCO]
C ₁₃ H ₁₈ Br ₂ N ₂ O	[18683-91-5] $\Delta_{\text{fus}}H$ (I) $\Delta_{\text{fus}}H$ (II)	<i>trans</i> -4-[[2-amino-3,5-dibromophenyl)methyl]amino]cyclohexanol (ambroxol)	31.46 36.52	372.7 365.6		[2004CAI/FOP]
C ₁₃ H ₁₈ ClNO	[7287-36-7] $\Delta_{\text{fus}}H$	N-(4-chlorophenyl)-2,2-dimethylpentanamide	23.31	360.2	DSC	[1990DON/DRE]
C ₁₃ H ₁₈ ClNO	[2307-68-8] $\Delta_{\text{fus}}H$	N-(3-chloro-4-methylphenyl)-2-methylpentanamide	16.35	353.2	DSC	[1990DON/DRE]
C ₁₃ H ₁₈ N ₂ O ₂	[2164-08-1] $\Delta_{\text{fus}}H$	3-cyclohexyl-6,7-dihydro-1 <i>H</i> -cyclopentapyrimidine-2,4-(3 <i>H</i> ,5 <i>H</i>)-dione	42.31	584.3	DSC	[1990DON/DRE]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound					
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference	
C ₁₃ H ₁₈ N ₂ O ₂ S ₂	[na] $\Delta_{\text{fus}}H$	N-allyl-S-ethyl-N'-tosylisothiourea		28	335.2	DSC	[1992REI/HAN]
C ₁₃ H ₁₈ N ₂ O ₄	[87458-01-3] $\Delta_{\text{fus}}H$	hexyl N-(4-nitrophenyl) carbamate		32.74	376.7	DSC	[1993TIE/FRA]
C ₁₃ H ₁₈ N ₄ O ₃	[6493-05-6] $\Delta_{\text{fus}}H$	1-(5-oxohexyl)-3,7-dimethylxanthine (pentoxifylline)		36.6	376.8	DSC	[2009DOM/POB]
C ₁₃ H ₁₈ O	[5195-24-4] Δ_vH	4,4-dimethyl-1-phenyl-3-pentanone (405–520)		63.5	420	A	[1987STE/MAL]
C ₁₃ H ₁₈ O	[103-95-7] Δ_vH	<i>p</i> -isopropyl- α -methylhydrocinnamaldehyde (283–499)		72.6	298	A	[1987STE/MAL]
C ₁₃ H ₁₈ O	[1671-75-6] Δ_vH	1-phenyl-1-heptanone (373–550)		64.6	388	A	[1987STE/MAL, 1947STU]
C ₁₃ H ₁₈ O ₂	[41643-38-9] $\Delta_{\text{us}}H$ (liq cryst) $\Delta_{\text{us}}H$ (liq cryst-to-liq) $\Delta_{\text{sub}}H$	4-hexylbenzoic acid (347–363)		17.4 2.4 123.6 ± 1.0	371 380 298	ME	[1985PRI/PUC] [2004MON/ALM]
C ₁₃ H ₁₈ O ₂	[15687-27-1] $\Delta_{\text{fus}}H$ $\Delta_{\text{fus}}H$ $\Delta_{\text{fus}}H$ $\Delta_{\text{fus}}H$ $\Delta_{\text{fus}}H$	(±) α -methyl-4-(isobutyl)phenylacetic acid (ibuprofen)		27.94 26.6 25.5 26.65 25.7	347.6 346.4 347.2 348 350.9	DSC DSC DSC	[2010HON/HUA] [2006WAS/HOL] [2002GRA/RAS, 2004XU/SUN] [1999LI/ZEL]
C ₁₃ H ₁₈ O ₂	[51146-57-7] $\Delta_{\text{fus}}H$	(–) α -methyl-4-(isobutyl)phenylacetic acid (ibuprofen)		17.9	327.2	DSC	[1999LI/ZEL]
C ₁₃ H ₁₈ O ₂	[51146-56-6] $\Delta_{\text{fus}}H$ $\Delta_{\text{fus}}H$	(S) α -methyl-4-(2-methylpropyl)benzeneacetic acid (S)-ibuprofen		18.05 18.7	324.2 325.5	AC DSC	[2005XU/SUN] [1996BUR/KOL]
C ₁₃ H ₁₈ O ₂	[na] $\Delta_{\text{fus}}H$	S-ibuprofen		28.3	326.9	DSC	[2010CIL/ALB]
C ₁₃ H ₁₈ O ₂	[na] $\Delta_{\text{fus}}H$	RS-ibuprofen		39.5	350.4	DSC	[2010CIL/ALB]
C ₁₃ H ₁₈ O ₃	[200570-98-5] $\Delta_{\text{fus}}H$	3-hexyloxybenzoic acid		22.72	343		[2001LAI/LEE]
C ₁₃ H ₁₈ O ₅ S	[26225-79-6] $\Delta_{\text{fus}}H$	<i>(dl)</i> -2-ethoxy-2,3-dihydro-3,3-dimethyl-5-benzofuranylmethanesulfonate		26.25	344.1	DSC	[1990DON/DRE]
C ₁₃ H ₁₈ O ₇	[138-52-3] $\Delta_{\text{fus}}H$	2-(hydroxymethyl)phenyl- β - <i>(d)</i> -glucopyranoside, <i>(d)</i> -salicin		55.5	474.7	DSC	[2008PIN/DIO, 2008DIO/PIN]
C ₁₃ H ₁₉ BrO ₄	[929259-37-0] Δ_vH	1-bromo-2-[2-[2-(2-methoxyethoxy)ethoxy]ethoxy]benzene (333–370)		96.1 ± 0.4	298	GS	[2006DAB/SPO]
C ₁₃ H ₁₉ NO	[na] Δ_vH	3-phenylpropionic acid, N,N-diethylamide (353–439)		46.5	368	A	[1987STE/MAL]
C ₁₃ H ₁₉ NO	[141271-51-4] Δ_vH	(4R,5R)-2,2,3,4-tetramethyl-5-phenyl-1,3-oxazolidine (293–301)		61.6 ± 1.8	298		[1998GUD/TOR]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₃ H ₁₉ NO ₂	[3129-92-8] $\Delta_{\text{sub}}H$	cyclohexyl ammonium benzoate (289–298)	103.1	293.5	A	[1987STE/MAL, 1965MAR]
C ₁₃ H ₁₉ NO ₂	[7461-26-9] $\Delta_{\text{fus}}H$	hexyl N-phenylcarbamate	32.76	328		[1971PRI]
C ₁₃ H ₁₉ NO ₄	[73243-69-3] $\Delta_{\text{fus}}H$	N-phenylethyl-5-amino-1,5-dideoxy-(<i>d</i>)-glucopyranose	39.9	455.8		[1994BLU/PRA]
C ₁₃ H ₁₉ NO ₄ S	[57-66-9] $\Delta_{\text{fus}}H$ $\Delta_{\text{fus}}H$	4-[(dipropylamino)sulfonyl]benzoic acid (probenecid)	33.57	471	DSC	[2009PEN/ESC]
			40.9	472.1	DSC	[2006WAS/HOL]
C ₁₃ H ₁₉ N ₃ O ₃	[16577-64-3] $\Delta_{\text{fus}}H$	1-hexyl-3-(4-nitrophenyl) urea	25.47	384.4	DSC	[1993TIE/FRA]
C ₁₃ H ₁₉ N ₃ O ₄	[40487-42-1] $\Delta_{\text{fus}}H$	N-(1-ethylpropyl)-2,6-dinitro-3,4-xylylidine	25.19	327.5	DSC	[1991ACR, 1990DON/DRE]
C ₁₃ H ₂₀	[1078-71-3] Δ_vH Δ_vH Δ_vH	heptylbenzene (292–353) (423–527)	64.2 ± 0.2	298	GS	[2006VER]
			54.0	438	A	[1987STE/MAL]
			64.9	298		[1971WIL/ZWO]
C ₁₃ H ₂₀ N ₂ O ₂ S ₂	[81261-44-1] $\Delta_{\text{fus}}H$	N-isobutyl-S-methyl-N'-tosylisothiurea	29.4	363.2	DSC	[1992REI/HAN]
C ₁₃ H ₂₀ N ₂ O ₂ S ₂	[120563-91-9] $\Delta_{\text{fus}}H$	N- <i>tert</i> -butyl-S-methyl-N'-tosylisothiurea	30.4	394.2	DSC	[1992REI/HAN]
C ₁₃ H ₂₀ N ₂ O ₂ S ₂	[145198-70-5] $\Delta_{\text{fus}}H$	N-isopropyl-S-ethyl-N'-tosylisothiurea	29.1	392.2	DSC	[1992REI/HAN]
C ₁₃ H ₂₀ O	[na] Δ_vH	butyl cumyl ether (278–318)	63.8 ± 0.5	298	GS	[2001VER/HEI2]
C ₁₃ H ₂₀ O	[127-41-3] Δ_vH Δ_vH	α -ionone (352–523) (286–333)	62.0	367	A	[1987STE/MAL, 1947STU]
			67.5	301	A, ME	[1987STE/MAL, 1957SER/VOI]
C ₁₃ H ₂₀ O	[14901-07-6] Δ_vH	β -ionone (291–334)	69.0	306	A, ME	[1987STE/MAL, 1957SER/VOI]
C ₁₃ H ₂₀ O	[16647-05-5] Δ_vH	6,10-dimethyl-4,5,9-undecatrien-2-one (349–421)	63.6 ± 1.4	385		[1988BAG/GUR]
C ₁₃ H ₂₀ O	[141-10-6] Δ_vH	6,10-dimethyl-3,5,9-undecatrien-2-one (382–457)	67.6 ± 1.1	420		[1988BAG/GUR]
C ₁₃ H ₂₀ O	[79-77-6] Δ_vH	4-(2,6,6-trimethyl-1-cyclohexen-1-yl)-3-buten-2-one (373–442)	49.6 ± 1.1	408		[1988BAG/GUR]
C ₁₃ H ₂₀ O ₂	[500-67-4] Δ_vH	1,3-dihydroxy-5-heptylbenzene (443–504)	91.6	458	A, GC	[1987STE/MAL, 1975KUN/LIL]
C ₁₃ H ₂₀ O ₂	[41395-27-1] Δ_vH	1,3-dihydroxy-5-methyl-2-hexylbenzene (433–493)	82.8	448	A, GC	[1987STE/MAL, 1975KUN/LIL]
C ₁₃ H ₂₁ Cl ₃ OS	[76619-97-1] Δ_vH	2,3,3-trichloro-2-propenethioic acid, O-decyl ester (483–503)	79.9		GC	[1980PIT/KIS]
C ₁₃ H ₂₁ N	[29772-98-3]	N,N-dimethyl-3-methyl-3-phenyl-2-butaneamine				

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	Δ_vH	(283–330)	59.8 ± 0.7	307	GS	[1998VER/BEC]
	Δ_vH	(283–330)	60.3 ± 0.7	298	GS	[1998VER/BEC]
C ₁₃ H ₂₁ N	[29772-82-5]	N-methyl-2,3-dimethyl-3-phenyl-2-butaneamine				
	Δ_vH	(285–332)	71.9 ± 1.1	309	GS	[1998VER/BEC]
	Δ_vH	(285–332)	72.5 ± 1.1	298	GS	[1998VER/BEC]
C ₁₃ H ₂₁ N	[585-48-8]	2,6-di- <i>tert</i> -butylpyridine				
	Δ_vH		56.6 ± 1.2	298		[2008FRI/ACR]
C ₁₃ H ₂₁ NO	[90-84-6]	2-(diethylamino)-1-phenyl-1-propanone				
	Δ_vH	(293–333)	71.6 ± 1.0	298	GS	[1994WEL/VER]
C ₁₃ H ₂₁ NO	[1502-00-7]	N,N-dimethyl-1-adamantylcarboxamide				
	$\Delta_{\text{sub}}H$	(303–322)	96.9 ± 0.3	313	ME	[1993ABB/JIM, 1995ABB/JIM]
	$\Delta_{\text{sub}}H$		97.5 ± 0.3	298	ME	[1995ABB/JIM]
C ₁₃ H ₂₁ NO ₂	[3246-04-6]	N-(3-phenoxy-2-hydroxypropyl)butylamine				
	$\Delta_{\text{sub}}H$	(323–348)	133.9	335.5	A	[1987STE/MAL]
C ₁₃ H ₂₁ N ₂ O	[18530-56-8]	N,N-dimethyl-N'-(octahydro-4,7-methano-1 <i>H</i> -inden-5-yl)urea				
	$\Delta_{\text{fus}}H$		21.74	436.5	DSC	[1990DON/DRE]
C ₁₃ H ₂₂	[na]	2-allyl- <i>cis</i> -decahydronaphthalene				
	Δ_vH	(296–320)	89.9	308	A	[1987STE/MAL]
C ₁₃ H ₂₂	[na]	2-allyl- <i>trans</i> -decahydronaphthalene				
	Δ_vH	(296–320)	91.7	308	A	[1987STE/MAL]
C ₁₃ H ₂₂	[5744-03-6]	dodecahydrofluorene				
	Δ_vH	(332–525)	55.8	347	A	[1987STE/MAL]
C ₁₃ H ₂₂	[707-35-7]	1,3,5-trimethyladamantane				
	$\Delta_{\text{fus}}H$		8.19	234.4		
	$\Delta_{\text{fus}}H$		2.06	255.6		[2000DRU/VAR2]
	$\Delta_{\text{fus}}H$		6.3	228.2		
	$\Delta_{\text{fus}}H$		1.73	253.6		[1977CLA/KNO]
	$\Delta_{\text{sub}}H$	(300–360)	77.8 ± 1.3	298	BG	[1977STE/WAT]
	Δ_vH		51.7 ± 0.2	298		[2000DRU/VAR, 2000MEL/PIM]
C ₁₃ H ₂₂ Cl ₂ O ₄	[na]	2,2- <i>bis</i> (chloromethyl)-1,3-propanediol dibutyrate				
	Δ_vH	(454–572)	43.1	469	A	[1987STE/MAL]
C ₁₃ H ₂₂ O ₂	[78548-53-5]	bornyl propionate				
	Δ_vH	(337–508)	55.9	352	A	[1987STE/MAL, 1947STU]
C ₁₃ H ₂₂ O ₃	[4427-97-8]	dicyclohexyl carbonate				
	$\Delta_{\text{sub}}H$	(293–313)	66.5 ± 4.2	303	ME	[1971KIP/RAB, 1977PED/RYL]
C ₁₃ H ₂₂ O ₃	[49540-31-0]	3,3,7,7-tetramethylnonanedioic anhydride				
	$\Delta_{\text{fus}}H$		20.5	396.2		[1974BOR]
C ₁₃ H ₂₄ Cl ₄	[3922-33-6]	1,1,1,1,3-tetrachlorotridecane				
	Δ_vH	(320–370)	97.4	335	A	[1987STE/MAL]
C ₁₃ H ₂₄ O	[42023-59-6]	5-methyl-2-ethyl-2-butyl-4-hexenal				
	Δ_vH	(323–393)	69.1	338	A	[1987STE/MAL]
C ₁₃ H ₂₄ N ₆	[na]	1-(hexamethyleneimine)-3,5- <i>bis</i> (dimethylamino)- <i>s</i> -triazine				
	$\Delta_{\text{fus}}H$		16.32	335.8		[1989BRA/RYT]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₃ H ₂₄ O ₂	[2156-96-9] $\Delta_v H$	decyl acrylate (404–536)	59.6	419	A	[1987STE/MAL]
C ₁₃ H ₂₄ O ₂	[1725-04-8] $\Delta_{\text{m}} H$ $\Delta_{\text{fus}} H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$	oxa-2-cyclotetradecanone (tridecanolactone)	18.16 9.08 66.6 ± 1.1 72.9 ± 1.7 67.5	290.6 300.4 390 298 408	MM MM A	[1996DOM/HEA] [1991WIB/WAL] [1991WIB/WAL] [1987STE/MAL]
C ₁₃ H ₂₄ O ₂	[na] $\Delta_v H$	3,3-dimethylbutanoic acid, 1-methylcyclohexyl ester (333–378)	61.4	298	CGC	[1999VER/HEI]
C ₁₃ H ₂₄ O ₂	[na] $\Delta_v H$	3,3-dimethylbutanoic acid, 3-methylcyclohexyl ester (333–378)	63.5	298	CGC	[1999VER/HEI]
C ₁₃ H ₂₄ O ₂	[1027080-67-6] $\Delta_v H$	3,3-dimethylbutanoic acid, 4-methylcyclohexyl ester (333–378)	64.1	298	CGC	[1999VER/HEI]
C ₁₃ H ₂₄ O ₂	[692-86-4] $\Delta_v H$	ethyl 10-undecenoate (404–532)	77.4	419	A	[1987STE/MAL]
C ₁₃ H ₂₄ O ₃	[1898-91-1] $\Delta_v H$	1,4-dioxo-5-cyclopentadecanone (403–443)	69.6	418	A, GC	[1987STE/MAL, 1971VOI/SHC]
C ₁₃ H ₂₄ O ₃	[36575-54-9] $\Delta_v H$	1,6-dioxo-7-cyclopentadecanone (403–443)	75.7	418	A, GC	[1987STE/MAL, 1971VOI/SHC]
C ₁₃ H ₂₄ O ₃	[36575-53-8] $\Delta_v H$	1,8-dioxo-9-cyclopentadecanone (403–443)	66.5	418	A, GC	[1987STE/MAL, 1971VOI/SHC]
C ₁₃ H ₂₄ O ₃	[18871-17-5] $\Delta_v H$	3-hexyl-4-acetoxytetrahydro-2H-pyran (383–453)	72.1	398	A	[1987STE/MAL]
C ₁₃ H ₂₄ O ₃	[41780-57-8] $\Delta_v H$ $\Delta_v H$	octyl levulinate (413–565)	66.3 65.1	428 507	A	[1987STE/MAL] [1933COW/SCH]
C ₁₃ H ₂₄ O ₄	[na] $\Delta_v H$	octyl 3-acetoxypionate (420–440)	88.4	430	A	[1987STE/MAL]
C ₁₃ H ₂₄ O ₄	[72030-39-8] $\Delta_v H$	ethylisopentylmalonic acid, ethyl methyl ester (392–501)	73.1	407	A	[1987STE/MAL]
C ₁₃ H ₂₄ O ₄	[505-52-2] $\Delta_{\text{fus}} H$	1,13-tridecanedioic acid (brassylic acid)	49.4	386.3	DSC	[2005ROU/TEM]
C ₁₃ H ₂₄ O ₅	[5456-15-5] $\Delta_v H$	octyl[1-(methoxycarbonyl)ethyl] carbonate (391–566)	70.0	406	A	[1987STE/MAL]
C ₁₃ H ₂₄ O ₅	[na] $\Delta_v H$	pentyl[1-(butoxycarbonyl)ethyl] carbonate (348–513)	70.1	363	A	[1987STE/MAL]
C ₁₃ H ₂₅ N	[629-60-7] $\Delta_v H$ $\Delta_v H$	tridecanonitrile (301–363) (380–566)	80.3 ± 0.4 69.5	298 395	GS A	[2005EME/VER] [1987STE/MAL]
C ₁₃ H ₂₅ NO	[20299-83-6] $\Delta_v H$	1-octanoyl piperidine (373–443)	50.0	388	A	[1987STE/MAL]
C ₁₃ H ₂₆	[7367-38-6]	5-butyl-4-nonene				

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(310–361)	55.8	325	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₁₃ H ₂₆	[2437-56-1]	1-tridecene				
	$\Delta_v H$		65.3	298		[1971WIL/ZWO]
	$\Delta_v H$	(413–509)	53.9	428	A	[1987STE/MAL, 1955CAM/ROS]
C ₁₃ H ₂₆	[1795-20-6]	<i>n</i> -octylcyclopentane				
	$\Delta_v H$		65.8	298		[1971WIL/ZWO]
C ₁₃ H ₂₆	[5617-41-4]	<i>n</i> -heptylcyclohexane				
	$\Delta_{\text{fus}} H$		22.22	232.8		[1996DOM/HEA]
	$\Delta_v H$		64.9	298		[1971WIL/ZWO]
C ₁₃ H ₂₆	[295-02-3]	cyclotridecane				
	$\Delta_{\text{fus}} H$		0.9	285.6		
	$\Delta_{\text{fus}} H$		7.4	297.6		[1987DRO/MOL]
C ₁₃ H ₂₆ O	[53144-53-9]	5-methyl-2-ethyl-2-butyl-4-hexene-1-ol				
	$\Delta_v H$	(333–393)	76.9	348	A	[1987STE/MAL]
C ₁₃ H ₂₆ O	[30089-09-9]	1-octylcyclopentanol				
	$\Delta_v H$	(468–541)	60.9	483	A	[1987STE/MAL]
C ₁₃ H ₂₆ O	[10486-19-8]	tridecanal				
	$\Delta_v H$	(325–349)	73.3 ± 0.4	298	GS	[2003VER/KRA2]
C ₁₃ H ₂₆ O	[593-08-8]	2-tridecanone				
	$\Delta_v H$	(335–534)	69.6	350	A	[1987STE/MAL]
	$\Delta_v H$	(424–510)	61	439	A	[1987STE/MAL]
	$\Delta_v H$	(400–628)	49.6	541		[1975AMB/ELL]
	$\Delta_v H$	(335–431)	69.8	348	EB	[1966MEY/WAG]
	$\Delta_v H$	(360–535)	62.1	375		[1947STU]
C ₁₃ H ₂₆ O	[462-18-0]	7-tridecanone				
	$\Delta_{\text{sub}} H$	(287–293)	103.8	290	ME	[1938UBB]
	$\Delta_v H$	(395–600)	62.7	410	A	[1987STE/MAL]
	$\Delta_v H$	(396–623)	49.3	536		[1975AMB/ELL]
C ₁₃ H ₂₆ O	[64470-31-1]	(Z) 7-tridecen-1-ol				
	$\Delta_v H$	(343–383)	95.1	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₃ H ₂₆ O	[64437-28-1]	(E) 7-tridecen-1-ol				
	$\Delta_v H$	(343–383)	95.6	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₃ H ₂₆ O	[52957-10-5]	(Z) 9-tridecen-1-ol				
	$\Delta_v H$	(343–383)	95.8	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₃ H ₂₆ O	[52957-15-0]	(E) 9-tridecen-1-ol				
	$\Delta_v H$	(343–383)	96.4	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₃ H ₂₆ O	[34010-24-7]	(Z) 11-tridecen-1-ol				
	$\Delta_v H$	(343–383)	97.1	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₃ H ₂₆ O	[56195-34-7]	(E) 11-tridecen-1-ol				
	$\Delta_v H$	(343–383)	97.2	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₃ H ₂₆ O	[1604-34-8]	6,10-dimethyl-2-undecanone				
	$\Delta_v H$	(379–473)	59.3 ± 0.4	426		[1988BAG/GUR]
C ₁₃ H ₂₆ O ₂	[5452-11-9]	4,5-dimethyl-2-octyl-1,3-dioxolane				

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(333–453)	72.8	348	A	[1987STE/MAL]
C ₁₃ H ₂₆ O ₂	[61732-94-3] $\Delta_v H$	2-octyl-1,3-dioxepane (323–373)	61.2	338	A	[1987STE/MAL]
C ₁₃ H ₂₆ O ₂	[1731-81-3] $\Delta_v H$ $\Delta_v H$	undecyl acetate (289–329) (333–378)	75.1 ± 0.3 77.2	298 298	GS GC	[2006KRA/VER] [1997KOU/HOS, 2000OVA/KOU]
C ₁₃ H ₂₆ O ₂	[2311-59-3] $\Delta_v H$	isopropyl decanoate (363–451)	60.8	378	A	[1987STE/MAL]
C ₁₃ H ₂₆ O ₂	[627-90-7] $\Delta_{\text{fus}} H$	ethyl undecanoate	36.16	259.2	AC	[2005VAN/OON]
C ₁₃ H ₂₆ O ₂	[30673-60-0] $\Delta_v H$	propyl decanoate (369–459)	62.4	384	A	[1987STE/MAL]
C ₁₃ H ₂₆ O ₂	[111-82-0] $\Delta_{\text{sub}} H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$	methyl laurate (262–273)	121.8 ± 2.1	267	ME	[1965DAV/KYB, 1987STE/MAL]
			71.4	350		[2002VAN/VAN]
			70.7 ± 0.2	356		[2002VAN/VAN]
			76.6 ± 0.4	298		[2002VAN/VAN]
		(295–452)	74.9	310		[2001BUR/JOS]
		(393–463)	76.8	298	GC	[1997KRO/VEL]
		(453–543)	53.3	498	GC	[1993HUS/SAR]
		(287–333)	83.6	302	A	[1987STE/MAL]
			76.5 ± 0.7	298	C,GC	[1980FUC/PEA]
			77.2 ± 0.6	298	C	[1977MAN/SEL]
		(407–540)	63.6	422	A	[1987STE/MAL, 1963ROS/SCH]
		(336–409)	71.4	351	MG,OM	[1952SCO/MAC]
		(373–439)	62.3	388		[1944ALT/TRI]
C ₁₃ H ₂₆ O ₂	[245658-41-7] $\Delta_v H$	3,3-dimethylbutanoic acid, 1,1,3-trimethylbutyl ester (333–378)	58.4	298	CGC	[1999VER/HEI]
C ₁₃ H ₂₆ O ₂	[na] $\Delta_v H$	2,6-dimethyl-2-heptanol butanoate (333–378)	62.5	298	CGC	[1999VER/HEI]
C ₁₃ H ₂₆ O ₂	[245658-44-0] $\Delta_v H$	2-methylpropanoic acid, 1,1,5-trimethylhexyl ester (333–378)	60.8	298	CGC	[1999VER/HEI]
C ₁₃ H ₂₆ O ₂	[638-53-9] $\Delta_{\text{trs}} H$ $\Delta_{\text{trs}} H$ $\Delta_{\text{fus}} H$ $\Delta_{\text{trs}} H$ $\Delta_{\text{fus}} H$ $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$	tridecanoic acid	0.06 8.5 33.0 8.72 33.74	287.7 309.1 314.6 307.1 315	DSC	[2007GBA/NEG] [1996DOM/HEA]
		(271–282)	112.5		TPTD	[2005CHA/ZIE]
		(282–299)	170		TPTD	[2001CHA/TOB]
						Note: Experimental values based on the TPTD method are often inconsistent with values determined using other experimental methods
		(409–585)	90.1	424	A	[1987STE/MAL]
		(328–350)	100.4 ± 2.0	340	ME, TE	[1982DEK/SCH]
C ₁₃ H ₂₆ O ₃	[42175-34-8] $\Delta_v H$	decyl lactate (349–556)	76.6	364	A	[1987STE/MAL]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₃ H ₂₆ O ₃	[na] $\Delta_v H$	octyl 3-ethoxypropionate (398–543)	56.9	413	A	[1987STE/MAL]
C ₁₃ H ₂₆ O ₃	[14144-56-0] $\Delta_v H$	pentyl 3-pentyloxypropionate (378–498)	62.3	393	A	[1987STE/MAL]
C ₁₃ H ₂₆ O ₃	[40915-96-6] $\Delta_{\text{sub}} H$	peroxytridecanoic acid (293–303)	142.7 ± 5	298	ME	[1980SWA/KWA]
C ₁₃ H ₂₇ Br	[765-09-3] $\Delta_v H$	1-bromotridecane (425–628)	64.6	440	A, EST	[1987STE/MAL, 1961LI/ROS, 1970DYK/VAN]
C ₁₃ H ₂₇ Cl	[822-13-9] $\Delta_v H$ $\Delta_v H$	1-chlorotridecane	81.3	298		[2006BOL/NER2]
		(414–611)	63.0	429	A, EST	[1987STE/MAL, 1961LI/ROS, 1970DYK/VAN]
C ₁₃ H ₂₇ F	[1536-21-6] $\Delta_v H$	1-fluorotridecane (387–558)	58.9	402	A, EST	[1987STE/MAL, 1961LI/ROS, 1970DYK/VAN]
C ₁₃ H ₂₇ I	[35599-77-0] $\Delta_v H$ $\Delta_v H$	1-iodotridecane (440–655)	85.0	298	A, EST	[1987STE/MAL, 1961LI/ROS, 1970DYK/VAN, 2006BOL/NER]
		(440–655)	66.1	455	A, EST	[1987STE/MAL, 1961LI/ROS, 1970DYK/VAN]
C ₁₃ H ₂₇ NO	[27563-67-3] $\Delta_{\text{sub}} H$	N-methyl dodecanamide (323–337)	116.6 ± 0.8	330	GS	[1959DAV/JON, 1987STE/MAL]
C ₁₃ H ₂₇ NO ₂	[6280-24-6] $\Delta_v H$	N-decyl lactamide (413–483)	97.9	428	A	[1987STE/MAL]
C ₁₃ H ₂₇ NO ₂	[na] $\Delta_v H$	O-decyl lactamide (413–483)	95.0	428	A	[1987STE/MAL]
C ₁₃ H ₂₈	[629-50-5] $\Delta_{\text{us}} H$ $\Delta_{\text{fus}} H$ $\Delta_{\text{fus}} H$ $\Delta_{\text{fus}} H$ $\Delta_{\text{sub}} H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$	tridecane	7.7	255.2		
			28.9	267.7	DSC	[2004MON/RAJ]
			7.66	255		
			28.49	267.8		[1996DOM/HEA]
			91.4	298	B	[1972MOR3]
			65.3	309	C	[1996VIT/CHA]
			64.9	314	C	[1996VIT/CHA]
			64.2	324	C	[1996VIT/CHA]
			63.3	334	C	[1996VIT/CHA]
			62.4	344	C	[1996VIT/CHA]
			62.3	349	C	[1996VIT/CHA]
			66.7	298		[1994RUZ/MAJ]
			65.6	308	C	[1979SUN/SVE]
			64.6	318	C	[1979SUN/SVE]
			61.7	348	C	[1979SUN/SVE]
	66.5 ± 0.2	298	C	[1979SUN/SVE]		
	66.4 ± 0.3	298	C	[1972MOR2]		
	66.2	298		[1971WIL/ZWO]		
	(417–511)	54.5	432	A	[1987STE/MAL, 1955CAM/ROS]	
C ₁₃ H ₂₈	[1560-97-0]	2-methyldodecane				

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(373–503)	52.5	388	A	[1987STE/MAL]
C ₁₃ H ₂₈	[17312-57-1]	3-methyldodecane				
	$\Delta_v H$	(372–504)	51.4	387	A	[1987STE/MAL]
C ₁₃ H ₂₈	[6117-97-1]	4-methyldodecane				
	$\Delta_v H$	(372–501)	52.0	387	A	[1987STE/MAL]
C ₁₃ H ₂₈	[17453-93-9]	5-methyldodecane				
	$\Delta_v H$	(368–500)	50.6	383	A	[1987STE/MAL]
C ₁₃ H ₂₈	[17312-77-5]	2,3-dimethylundecane				
	$\Delta_v H$	(383–500)	53.2	398	A	[1987STE/MAL]
C ₁₃ H ₂₈	[17312-80-0]	2,4-dimethylundecane				
	$\Delta_v H$	(365–490)	52.1	380	A	[1987STE/MAL]
C ₁₃ H ₂₈	[62108-27-4]	2,4,6-trimethyldecane				
	$\Delta_v H$	(352–478)	48.7	367	A	[1987STE/MAL]
C ₁₃ H ₂₈	[na]	5-ethyl-5-methyldecane				
	$\Delta_v H$	(273–307)	61.4 ± 1.1	290	HSA	[1995CHI/HES]
	$\Delta_v H$		60.5 ± 1.1	298		[1995CHI/HES]
	$\Delta_v H$		61.4 ± 1.8	298	CGC	[1995CHI/HES]
C ₁₃ H ₂₈	[17312-63-9]	5-butylnonane				
	$\Delta_v H$	(298–365)	52.6	313	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₁₃ H ₂₈	[35660-96-9]	tri- <i>tert</i> -butylmethane				
	$\Delta_{\text{fus}}H$		3.53	358.2		[1986FLA/BEC]
	$\Delta_{\text{sub}}H$		55.4	298	CGC-DSC	[1998CHI/HES]
	$\Delta_{\text{sub}}H$	(265–319)	57.0 ± 0.4	288	T	[1997VER/NOL]
	$\Delta_{\text{sub}}H$	(273–306)	57.7 ± 2.8	290	HSA	[1995CHI/HES]
	$\Delta_{\text{sub}}H$		61.1 ± 1.3			[1995CHI/HES]
C ₁₃ H ₂₈ N ₂ O	[2158-09-0]	1-dodecyl urea				
	$\Delta_{\text{fus}}H$		1.3	275.4		
C ₁₃ H ₂₈ O	[508181-43-9]	pentyl <i>tert</i> -octyl ether				
	$\Delta_v H$	(278–303)	55.9 ± 0.3	298	GS	[UR/VER, 2002VER, 2003VER/KRA]
C ₁₃ H ₂₈ O	[112-70-9]	1-tridecanol				
	$\Delta_{\text{fus}}H$		44.78	304.7		[2003VAN/VAN]
	$\Delta_{\text{fus}}H$		45.1	304.6		
	$\Delta_{\text{fus}}H$		41.42	304.5		
	$\Delta_{\text{fus}}H$		23.3	303.5		
	$\Delta_{\text{fus}}H$		3.6	301.6		
	$\Delta_{\text{fus}}H$		22.09	305.8		
	$\Delta_{\text{fus}}H$		18.74	306.6		[1974MOS/MOU]
	$\Delta_v H$		94.7 ± 0.4	298	CGC	[2006NIC/KWE]
	$\Delta_v H$	(307–348)	91.1	327	GS	[2001KUL/VER2]
	$\Delta_v H$	(307–348)	95.8	298	GS	[2001KUL/VER2]
	$\Delta_v H$	(313–373)	87.4	343		[1992NGU/KAS]
$\Delta_v H$	(431–568)	69.2	446	A	[1987STE/MAL]	
C ₁₃ H ₂₈ O	[42930-67-6]	2,2-dimethyl-3- <i>tert</i> -butyl-3-heptanol				
	$\Delta_v H$	(379–513)	58.3	394		[1973WIL/ZWO]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₃ H ₂₈ O	[na] $\Delta_v H$	3,3,5,5-tetramethyl-4-ethyl-4-heptanol (393–526)	55.9	408		[1973WIL/ZWO]
C ₁₃ H ₂₈ O	[na] $\Delta_v H$	3,3,6-trimethyl-4-isopropyl-4-heptanol (381–512)	59.1	396		[1973WIL/ZWO]
C ₁₃ H ₂₈ O	[na] $\Delta_v H$	3,3,6-trimethyl-4-propyl-4-heptanol (383–513)	60.1	398		[1973WIL/ZWO]
C ₁₃ H ₂₈ O	[32579-70-7] $\Delta_v H$	2,2,5-trimethyl-3- <i>tert</i> -butyl-3-hexanol (377–513)	57.6	392		[1973WIL/ZWO]
C ₁₃ H ₂₈ O	[41902-42-5] $\Delta_{\text{fus}} H$ $\Delta_{\text{fus}} H$	tri- <i>tert</i> -butylmethanol	7.2 3.43	302 390		[1996DOM/HEA]
	$\Delta_{\text{sub}} H$ (<i>plastic</i>)	(278–318)	56.5 ± 1.0	298	TE	[1983MAS/STE]
	$\Delta_{\text{sub}} H$ (<i>cryst</i>)	(269–300)	63.2 ± 1.2	298	TE	[1983MAS/STE]
C ₁₃ H ₂₈ O ₂	[13362-25-2] $\Delta_{\text{fus}} H$ $\Delta_{\text{fus}} H$	1,13-tridecanediol	28.9 17.8	343 351	DSC	[1999OGA/NAK]
C ₁₃ H ₂₈ O ₂ S	[24724-30-9] $\Delta_{\text{fus}} H$ $\Delta_{\text{fus}} H$	3-(decylthio)-1,2-propanediol	17.3 17.3	291.9 311.9	DSC	[1993ACR]
C ₁₃ H ₂₈ O ₃	[10430-97-4] $\Delta_{\text{fus}} H$	3-(decyloxy)-1,2-propanediol	38.9	311	DSC	[1993ACR]
C ₁₃ H ₂₈ O ₄	[57499-93-1] $\Delta_v H$	tripropylene glycol, monobutyl ether (374–543)	67.1	389	A	[1987STE/MAL, 1947STU]
C ₁₃ H ₂₈ O ₅ S ₂	[123483-21-6] $\Delta_{\text{fus}} H$	(<i>l</i>)-arabinose dibutyl dithioacetal	41.5	380.4	DSC	[1989VAN/VAN]
C ₁₃ H ₂₈ S	[19484-26-5] $\Delta_v H$	1-tridecanethiol (433–598)	64.7	448		[1999DYK/SVO]
C ₁₃ H ₂₉ N	[2869-34-3] $\Delta_v H$	tridecylamine (458–562)	60.1	473	A, EST	[1987STE/MAL, 1956MAN2]
C ₁₃ H ₂₉ NO ₂	[1191-45-3] $\Delta_{\text{fus}} H$	3-(decylamino)-1,2-propanediol	54.8	346.6	DSC	[1993ACR]
C ₁₄ H ₅ F ₂₅	[89109-68-2] $\Delta_{\text{fus}} H$	1,1,1,2,2,3,3,4,4,5,5,6,6,7,7,8,8,9,9,10,10,11,11,12,12-pentacosofluorotetradecane	20.8	344.2	DSC	[1986RUS/RAB]
C ₁₄ H ₆ Cl ₂ N ₂ O ₄	[66121-41-3] $\Delta_{\text{sub}} H$	1-amino-4-nitro-5,8-dichloroanthraquinone	158.2			[1968TSU/KOJ, 1988BAU/PER]
C ₁₄ H ₆ N ₂ O ₆	[66121-37-7] $\Delta_{\text{sub}} H$	1,4-dinitroanthraquinone	131			[1968TSU/KOJ, 1988BAU/PER]
C ₁₄ H ₆ N ₆ O ₁₂	[20062-22-0] $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$	1,2- <i>bis</i> (2,4,6-trinitrophenyl)ethylene (434–479)	179.9 180.3	449	LE	[1987STE/MAL, 1969ROS/DIC] [1968MAR/ARM, 1966ROS]
C ₁₄ H ₇ ClF ₃ NO ₅	[50594-66-6] $\Delta_{\text{fus}} H$	5-[2-chloro-4-(trifluoromethyl)phenoxy]-2-nitrobenzoic acid	37.67	436.6	DSC	[1990DON/DRE]
C ₁₄ H ₇ ClO ₂	[131-09-9]	2-chloroanthraquinone				

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{fus}}H$		39.0	483		[1996DOM/HEA]
C ₁₄ H ₇ NO ₄	[82-34-8]	1-nitroanthraquinone				
	$\Delta_{\text{sub}}H$	(407–440)	139.7	422	A	[1987STE/MAL]
	$\Delta_{\text{sub}}H$		108.9 ± 2.1	396	C	[1982MUR/SAK]
	$\Delta_{\text{sub}}H$		137.9 ± 1.7		TE,ME	[1970KOJ]
	$\Delta_{\text{sub}}H$		115.5			[1968TSU/KOJ, 1988BAU/PER]
C ₁₄ H ₈	[187-78-0]	paracylene				
	$\Delta_{\text{sub}}H$		82.0	342	ME	[2002DIO/KIY]
	$\Delta_{\text{sub}}H$		83.2	298	ME	[2002DIO/KIY]
C ₁₄ H ₈ Br ₂	[3278-82-8]	1,5-dibromoanthracene				
	$\Delta_{\text{sub}}H$	(358–408)	116.7 ± 3.0		ME	[2008GOL/SUU2]
C ₁₄ H ₈ Br ₂	[523-27-3]	9,10-dibromoanthracene				
	$\Delta_{\text{sub}}H$	(359–391)	114.2 ± 2.8		ME	[2008GOL/SUU2]
C ₁₄ H ₈ Cl ₂	[605-48-1]	9,10-dichloroanthracene				
	$\Delta_{\text{sub}}H$	(316–376)	113.9 ± 4.5		ME	[2008GOL/SUU2]
C ₁₄ H ₈ Cl ₄	[3424-92-6]	1-chloro-2-(2,2-dichloro-1-(4-chlorophenylethynyl)benzene				
	$\Delta_{\text{fus}}H$		23.84	349.8	DSC	[1990DON/DRE]
C ₁₄ H ₈ Cl ₄	[72-55-9]	1,1-dichloro-2,2-bis(4-chlorophenyl)ethylene (<i>p,p'</i> -DDE)				
	$\Delta_{\text{fus}}H$		23.55	360.4	DSC	[1990DON/DRE]
	$\Delta_{\text{sub}}H$		74.2			[1995RUL/RAK, 1989LUB/JAN]
	Δ_vH	(343–453)	87.2	398	GC	[1990HIN/BID2]
C ₁₄ H ₈ Cl ₆	[3563-45-9]	1,1,1-trichloro-2-chloro-2,2-bis(4-chlorophenyl)ethane				
	$\Delta_{\text{sub}}H$		89.4			[1995RUL/RAK, 1989LUB/JAN]
C ₁₄ H ₈ O ₂	[84-65-1]	9,10-anthraquinone				
	$\Delta_{\text{fus}}H$		32.57	558		[1996DOM/HEA]
	$\Delta_{\text{sub}}H$		111.3		GS	[1987SHI/OHK, 1991HOR]
	$\Delta_{\text{sub}}H$	(373–453)	98.3	413	GS	[1977NIS/ISH, 1978NIS/ISH]
	$\Delta_{\text{sub}}H$		113. ± 0.8	298	C	[1973BAR/MAL]
	$\Delta_{\text{sub}}H$		107.5 ± 0.8	434	ME	[1973BAR/MAL]
	$\Delta_{\text{sub}}H$	(397–471)	107.9 ± 0.8		ME	[1973BAR/MAL]
	$\Delta_{\text{sub}}H$	(355–356)	U 105.9		TGA	[1971ASH]
	$\Delta_{\text{sub}}H$	(470–590)	127.0 ± 3.0		C	[1971BEE/LIN]
	$\Delta_{\text{sub}}H$		136.6 ± 3	298	C	[1971BEE/LIN]
	$\Delta_{\text{sub}}H$		116.1 ± 1.7		ME,TE	[1970KOJ]
	$\Delta_{\text{sub}}H$		115.1		ME	[1968TSU/KOJ, 1988BAU/PER]
	$\Delta_{\text{sub}}H$	(343–403)	126.4	373	ME	[1958HOY/PEP, 1987STE/MAL]
	$\Delta_{\text{sub}}H$		112.1	298		[1956MAG, 1970COX/PIL]
	$\Delta_{\text{sub}}H$		110.9	298		[1956BEY/NIC]
	$\Delta_{\text{sub}}H$		107.9	298		[1954JOR]
$\Delta_{\text{sub}}H$		104.6	367	ME	[1952INO/SHI]	
$\Delta_{\text{sub}}H$		108	298	ME	[1952INO/SHI]	
	Δ_vH	(559–660)	64.3	574	A	[1987STE/MAL]
C ₁₄ H ₈ O ₂	[84-11-7]	9,10-phenanthraquinone				
	$\Delta_{\text{sub}}H$		108.1	289	C	[1989RIB/RIB]
	$\Delta_{\text{sub}}H$		132	383		[1956MAG, 1970COX/PIL]
C ₁₄ H ₈ O ₃	[129-43-1]	1-hydroxy-9,10-anthraquinone				

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{sub}}H$		113.4		GS	[1987SHI/OHK, 1991HOR]
	$\Delta_{\text{sub}}H$	(333–383)	120.6	358		[1958HOY/PEP, 1987STE/MAL]
	$\Delta_{\text{sub}}H$		101.3 ± 0.4	407	HSA	[1956BEY/NIC]
C ₁₄ H ₈ O ₃	[605-32-3]	2-hydroxy-9,10-anthraquinone				
	$\Delta_{\text{sub}}H$		136.8		GS	[1987SHI/OHK, 1991HOR]
	$\Delta_{\text{sub}}H$	(393–453)	153.1	408	A	[1987STE/MAL]
C ₁₄ H ₈ O ₃	[74553-57-4]	9-hydroxy-1,4-anthraquinone				
	$\Delta_{\text{sub}}H$	(377–394)	108.2 ± 2.2	386	ME	[2002JIM/ROU]
	$\Delta_{\text{sub}}H$	(377–394)	109.5 ± 2.2	298	ME	[2002JIM/ROU]
C ₁₄ H ₈ O ₃	[6050-13-1]	2,2'-biphenyldicarboxylic anhydride				
	$\Delta_{\text{sub}}H$		120.7 ± 4.0	298	C	[2005MAT/MIR2]
	$\Delta_{\text{sub}}H$	(433–490)	91.4	448	A	[1987STE/MAL]
C ₁₄ H ₈ O ₄	[72-48-0]	1,2-dihydroxyanthraquinone				
	$\Delta_{\text{sub}}H$	(368–498)	123.8	383	A	[1987STE/MAL]
	$\Delta_{\text{sub}}H$		121.9 ± 0.5	469	C	[1973MAL/BAR]
	$\Delta_{\text{sub}}H$	(434–505)	121.5 ± 0.4	469	ME	[1973MAL/BAR]
	$\Delta_{\text{sub}}H$		123.9	403	ME	[1958HOY/PEP]
C ₁₄ H ₈ O ₄	[81-64-1]	1,4-dihydroxy-9,10-anthraquinone (quinizarin)				
	$\Delta_{\text{sub}}H$		115.3	363		[2003HIN/RAF]
	$\Delta_{\text{sub}}H$		114.6		GS	[1987SHI/OHK, 1991HOR]
	$\Delta_{\text{sub}}H$	(353–373)	102.4 ± 4.4	363		[1984KRI]
	$\Delta_{\text{sub}}H$	(473–553)	89.1	513	GS	[1977NIS/ISH, 1978NIS/ISH]
	$\Delta_{\text{sub}}H$	(394–463)	121.9 ± 0.8	429	ME	[1973MAL/BAR]
	$\Delta_{\text{sub}}H$		121.1 ± 4	429	C	[1973MAL/BAR]
	$\Delta_{\text{sub}}H$	(324–351)	U 94.5	338	TGA	[1971ASH]
	$\Delta_{\text{sub}}H$		123.5	376		[1958HOY/PEP, 1987STE/MAL]
	$\Delta_{\text{sub}}H$		103.5 ± 1.3	409	HSA	[1956BEY/NIC]
	Δ_vH	(469–633)	74	484	A	[1987STE/MAL, 1947STU]
C ₁₄ H ₈ O ₄	[117-12-4]	1,5-dihydroxyanthraquinone				
	$\Delta_{\text{sub}}H$		123.2 ± 7		ME	[1973BAR/MAL]
	$\Delta_{\text{sub}}H$	(363–433)	126.8	398	ME	[1958HOY/PEP, 1987STE/MAL]
	$\Delta_{\text{sub}}H$		111.3	456		[1956BEY/NIC]
	$\Delta_{\text{sub}}H$		117.6	298		[1956BEY/NIC]
C ₁₄ H ₈ O ₄	[117-10-2]	1,8-dihydroxyanthraquinone				
	$\Delta_{\text{sub}}H$		116.8		ME	[1973BAR/MAL]
	$\Delta_{\text{sub}}H$	(333–403)	123	368	ME	[1958HOY/PEP, 1987STE/MAL]
	$\Delta_{\text{sub}}H$	(335–356)	U 96.5	345	TGA	[1971ASH]
	$\Delta_{\text{sub}}H$		105.8 ± 8	404	HSA	[1956BEY/NIC]
	$\Delta_{\text{sub}}H$		109.6 ± 8	298		[1956BEY/NIC]
C ₁₄ H ₈ O ₄	[84-60-6]	2,6-dihydroxyanthraquinone				
	$\Delta_{\text{sub}}H$	(463–533)	173.8	498		[1958HOY/PEP, 1987STE/MAL]
C ₁₄ H ₈ O ₆	[81-60-7]	1,4,5,8-tetrahydroxyanthraquinone				
	$\Delta_{\text{sub}}H$	(403–473)	151.6	438		[1958HOY/PEP, 1987STE/MAL]
C ₁₄ H ₉ Br	[1564-64-3]	9-bromoanthracene				
	$\Delta_{\text{sub}}H$	(315–368)	100.5 ± 1.8		ME	[2008GOL/SUU2]
C ₁₄ H ₉ Cl	[4985-70-0]	1-chloroanthracene				
	$\Delta_{\text{fus}}H$		14.14	355.2		[1970GUA/SAR]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₄ H ₉ Cl	[17135-78-3] $\Delta_{\text{sub}}H$	2-chloroanthracene (331–371)	99.3 ± 2.7		ME	[2008RIB/SCH]
C ₁₄ H ₉ Cl	[716-53-0] $\Delta_{\text{fus}}H$	9-chloroanthracene	18.66	379.2		[1970GUA/SAR]
C ₁₄ H ₉ ClF ₂ N ₂ O ₂	[35367-38-5] $\Delta_{\text{fus}}H$	N-[(4-chlorophenylamino)carbonyl]-2,6-difluorobenzamide	55.99	499.5	DSC	[1990DON/DRE]
C ₁₄ H ₉ ClN ₂ O ₄	[12217-79-7] $\Delta_{\text{sub}}H$	1,5-diaminochloro-4,8-dihydroxyanthraquinone (C.I. disperse blue 56) (483–533)	93.3	498	A	[1987STE/MAL]
C ₁₄ H ₉ Cl ₂ NO ₃	[42576-02-3] $\Delta_{\text{fus}}H$	methyl 5-(2,4-dichlorophenoxy)-2-nitrobenzoate	26.31	358.3	DSC	[1991ACR, 1990DON/DRE]
C ₁₄ H ₉ Cl ₃	[na] $\Delta_{\text{fus}}H$	1-chloro-2,2-(bis-(4-chlorophenyl)ethylene	25.52	337.9	DSC	[1969PLA/GLA]
C ₁₄ H ₉ Cl ₅	[50-29-3] $\Delta_{\text{fus}}H$	1,1,1-trichloro-2,2-bis(4-chlorophenyl)ethane (<i>pp'</i> DDT)	26.28	382.1	DSC	[1991ACR, 1990DON/DRE]
	$\Delta_{\text{sub}}H$	(273–313)	120.2 ± 1.0	293	GS	[1994WAN/SHU]
	$\Delta_{\text{sub}}H$	(323–363)	115	338	A	[1987STE/MAL]
	$\Delta_{\text{sub}}H$	(293–353)	110	304	GS	[1980ROT]
	$\Delta_{\text{sub}}H$	(293–313)	117.8	303	GS	[1972SPE/CLI]
	$\Delta_{\text{sub}}H$	(323–363)	117.5	338	GS	[1956DIC, 1960JON]
	$\Delta_{\text{sub}}H$	(313–363)	84	338	GS	[1949KUH/MAS]
	$\Delta_{\text{sub}}H$	(339–373)	118	356	TE	[1947BAL]
	Δ_vH		106.1 ± 1.3	398	GS	[2001PUR/CHI]
Δ_vH	(343–453)	93.2	398	GC	[1990HIN/BID2]	
C ₁₄ H ₉ Cl ₅	[789-02-6] Δ_vH	1,1,1-trichloro-2-(4-chlorophenyl)-2-(2-chlorophenyl)ethane (<i>p, o'</i> -DDT) (343–453)	88.6	398	GC	[1990HIN/BID2]
C ₁₄ H ₉ Cl ₅	[789-02-6] $\Delta_{\text{fus}}H$	1-chloro-2-(2,2,2-trichloro-1-(4-chlorophenyl)ethyl)benzene	23.09	345.8	DSC	[1990DON/DRE]
C ₁₄ H ₉ Cl ₅	[na] Δ_vH	DDT (313–363)	338	83.7		[1949KUH/MAS]
C ₁₄ H ₉ Cl ₅ O	[10606-46-9] $\Delta_{\text{fus}}H$	2-chloro- α -(4-chlorophenyl)- α -(trichloromethyl)benzenemethanol	25.2	396.3		[1991ACR, 1990DON/DRE]
C ₁₄ H ₉ Cl ₅ O	[115-32-2] $\Delta_{\text{fus}}H$	4-chloro- α -(4-chlorophenyl)- α -(trichloromethyl)benzenemethanol	19.56	347.2	DSC	[1990DON/DRE]
C ₁₄ H ₉ F ₃ O ₂	[893-33-4] $\Delta_{\text{sub}}H$	4,4,4-trifluoro-1-(2-naphthyl)-butan-1,3-dione	108.7 ± 0.6	298	ME	[1997RIB/GON]
C ₁₄ H ₉ F ₁₇ O ₂	[1996-88-9] $\Delta_{\text{trs}}H$	perfluorooctylethylene methacrylate	5.0	210		
	$\Delta_{\text{fus}}H$		9.0	253		[1992HOP/FAU]
C ₁₄ H ₉ F ₂₁ O	[na] $\Delta_{\text{fus}}H$	ω -perfluorodecyl-1-butanol	21.3	360		[1991HOP/MOL]
C ₁₄ H ₉ NO ₂	[82-45-1] $\Delta_{\text{fus}}H$	1-aminoanthraquinone	28.78	524.2		[1988BAU/PER]
	$\Delta_{\text{sub}}H$		121.8		GS	[1987SHI/OHK, 1991HOR]
	$\Delta_{\text{sub}}H$		(413–443)	126.5	428	A

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{sub}}H$	(368–393)	116.3 ± 3.9	380		[1984KRI]
	$\Delta_{\text{sub}}H$	(473–553)	103.3	513	GS	[1977NIS/ISH, 1978NIS/ISH]
	$\Delta_{\text{sub}}H$	(361–386)	U 90.9	374	TGA	[1971ASH]
	$\Delta_{\text{sub}}H$		125.9 ± 2.5		TE,ME	[1970KOJ]
	$\Delta_{\text{sub}}H$		131			[1968TSU/KOJ, 1988BAU/PER]
	$\Delta_{\text{sub}}H$		113 ± 0.4	463	HSA	[1956BEY/NIC]
C ₁₄ H ₉ NO ₂	[117-79-3]	2-aminoanthraquinone				
	$\Delta_{\text{sub}}H$		136.8		GS	[1987SHI/OHK, 1991HOR]
	$\Delta_{\text{sub}}H$		143.5 ± 2.9		TE,ME	[1970KOJ]
	$\Delta_{\text{sub}}H$		162.3			[1968TSU/KOJ, 1988BAU/PER]
C ₁₄ H ₉ NO ₂	[602-60-8]	9-nitroanthracene				
	$\Delta_{\text{fus}}H$		20.1	420.4	DSC	[2010KES/AUC]
	$\Delta_{\text{sub}}H$	(361–377)	111.9 ± 0.6	369	ME	[2006RIB/AMA3]
	$\Delta_{\text{sub}}H$	(361–377)	115.4 ± 0.6	298	ME	[2006RIB/AMA3]
C ₁₄ H ₉ NO ₃	[116-85-8]	1-hydroxy-4-aminoanthraquinone				
	$\Delta_{\text{sub}}H$		127.2		GS	[1987SHI/OHK, 1991HOR]
	$\Delta_{\text{sub}}H$	(418–438)	131.3	428	A	[1987STE/MAL, 1980ROD/KRU]
	$\Delta_{\text{sub}}H$	(444–473)	144	458.5	A	[1987STE/MAL]
	$\Delta_{\text{sub}}H$		119.6			[1984KAR/KRU]
	$\Delta_{\text{sub}}H$		133.5 ± 2.1		TE,ME	[1970KOJ]
	$\Delta_{\text{sub}}H$		120.1			[1968TSU/KOJ, 1988BAU/PER]
C ₁₄ H ₉ N ₃ O ₄	[82-33-7]	1,4-diamino-5-nitroanthraquinone				
	$\Delta_{\text{sub}}H$	(473–553)	U 50.2	513	GS	[1977NIS/ISH, 1978NIS/ISH]
		<i>(not crystalline)</i>				
C ₁₄ H ₁₀	[120-12-7]	anthracene				
	$\Delta_{\text{fus}}H$	(463–503)	29.8	492	DSC	[2003ROJ/ORO]
	$\Delta_{\text{fus}}H$		31.5	491	DSC	[2003STO/KRZ]
	$\Delta_{\text{fus}}H$		28.8	489.4	DSC	[2000LIS/JAM]
	$\Delta_{\text{fus}}H$		29.37	488.9		[1996DOM/HEA]
	$\Delta_{\text{sub}}H$	(339–399)	97.6 ± 1.3	369	ME	[2009SID/SID]
	$\Delta_{\text{sub}}H$	(339–399)	98.2	298	ME	[2009SID/SID]
	$\Delta_{\text{sub}}H$	(320–355)	97.9 ± 0.6		ME	[2009OJA/CHE]
	$\Delta_{\text{sub}}H$	(320–350)	98.4 ± 0.7		ME	[2009OJA/CHE]
	$\Delta_{\text{sub}}H$	(320–354)	95.6 ± 1.2	337		[2006CHE/OJA]
	$\Delta_{\text{sub}}H$	(340–360)	98.8 ± 0.4	350	ME	[2006RIB/MON]
	$\Delta_{\text{sub}}H$	(340–360)	100.2 ± 0.4	298	ME	[2006RIB/MON]
	$\Delta_{\text{sub}}H$	(348–368)	102.5 ± 1.9	358	ME	[2004VER]
	$\Delta_{\text{sub}}H$		96.3 ± 0.7	298	DSC	[2003ROJ/ORO]
	$\Delta_{\text{sub}}H$		106		DSC	[2003STO/KRZ]
	$\Delta_{\text{sub}}H$	(423–488)	94.5		MEM	[1999EMM/PIC]
	$\Delta_{\text{sub}}H$	(338–353)	102.5		ME	[1998KLO/LAU]
	$\Delta_{\text{sub}}H$		99.4	298	CGC-DSC	[1998CHI/HES]
	$\Delta_{\text{sub}}H$	(318–363)	100.0 ± 2.8	341	ME	[1998OJA/SUU]
	$\Delta_{\text{sub}}H$	(343–448)	84.0 ± 3.0	298	TGA	[1997TES/PIK]
	$\Delta_{\text{sub}}H$	(313–453)	99.7	383	GS	[1995NAS/LEN]
	$\Delta_{\text{sub}}H$	(318–373)	98.7	346	GS	[1986ROR]
	$\Delta_{\text{sub}}H$	(313–363)	102.6	338	GS	[1986HAN/ECK]
	$\Delta_{\text{sub}}H$	(353–399)	94.3		GS	[1983BEN/BIE]
	$\Delta_{\text{sub}}H$	(283–323)	91.8 ± 0.9	303	GS	[1983SON/ZOL]
	$\Delta_{\text{sub}}H$	(323–353)	91.2	338	GS	[1982GRA/FOS]
	$\Delta_{\text{sub}}H$		97.4 ± 1.1		GS,C	[1981BRO/MCE]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{sub}}H$		97.8 ± 0.1		HSA	[1980DYG/STE]
	$\Delta_{\text{sub}}H$	(337–361)	104.5 ± 1.5	298	TE,ME	[1980DEK]
	$\Delta_{\text{sub}}H$	(358–393)	94.8	376	GS	[1979MAC/PRA]
	$\Delta_{\text{sub}}H$	(363–448)	98.8 ± 0.4		HSA	[1977DYG/STE]
	$\Delta_{\text{sub}}H$	(328–372)	97.2		ME	[1976TAY/CRO]
	$\Delta_{\text{sub}}H$		97.1		C	[1975ADE/BRO]
	$\Delta_{\text{sub}}H$	(323–353)	102.9 ± 4.8	298	TE	[1975DEK/VAN]
	$\Delta_{\text{sub}}H$	(283–323)	95.8 ± 6		LE	[1973MCE/SAN]
	$\Delta_{\text{sub}}H$	(353–432)	101.0 ± 0.5		ME	[1973MAL/GIG]
	$\Delta_{\text{sub}}H$		99.7	393	C	[1973MAL/GIG]
	$\Delta_{\text{sub}}H$	(290–358)	84.1		ME,C	[1972WIE, 1971BEE/LIN]
	$\Delta_{\text{sub}}H$	(342–359)	98.3 ± 2.1			[1964KEL/RIC, 1970COX/PIL]
	$\Delta_{\text{sub}}H$	(327–346)	90 ± 0.13	337	TE	[1960BUD]
	$\Delta_{\text{sub}}H$		100.8			[1958HOY/PEP, 1970COX/PIL]
	$\Delta_{\text{sub}}H$	(303–373)	103.4 ± 2.9			[1958HOY/PEP, 1970COX/PIL]
	$\Delta_{\text{sub}}H$		100.8 ± 4.2			[1958HOY/PEP, 1970COX/PIL]
	$\Delta_{\text{sub}}H$	(396–421)	97.5 ± 2		HSA	[1953STE]
	$\Delta_{\text{sub}}H$	(339–353)	102.1	346		[1953BRA/CLE]
	$\Delta_{\text{sub}}H$	(338–353)	102.1 ± 2.1			[1953BRA/CLE2, 1970COX/PIL]
	$\Delta_{\text{sub}}H$		92.0 ± 2.1	364	ME	[1952INO/SHI]
	$\Delta_{\text{sub}}H$		90.4	353	ME	[1951INO]
	$\Delta_{\text{sub}}H$		95.4			[1951NIT/SEK]
	$\Delta_{\text{sub}}H$		95.0			[1950NIT/SEK3]
	$\Delta_{\text{sub}}H$	(378–398)	97.3 ± 1.2		RG	[1949SEA/HOP2]
	$\Delta_{\text{sub}}H$		104.6 ± 4.2			[1949KLA, 1970COX/PIL]
	$\Delta_{\text{sub}}H$		93.3 ± 4.2	353		[1938WOL/WEG]
	Δ_vH		66.7	498		[2003ROJ/ORO, 2008HAN/NUT]
	Δ_vH		78.5	298	CGC	[2008ZHA/UNH]
	Δ_vH	(413–473)	79.5 ± 1.2	298	GC	[2006HAF/PAR]
	Δ_vH	(323–473)	72.4	398	GC	[2002LEI/CHA]
	Δ_vH		79.1	298	CGC	[2001PUR/CHI]
	Δ_vH		79.8	298	CGC	[1998CHI/HES]
	Δ_vH	(453–503)	79.6	298	CGC	[1995CHI/HOS]
	Δ_vH	(343–453)	69.7	398	GC	[1990HIN/BID2]
	Δ_vH	(504–615)	58.6	519	A	[1987STE/MAL]
	Δ_vH		62.1	500		[1979KUD/KUD2, 2008HAN/NUT]
	Δ_vH	(500–616)	59.2	558	I	[1923MOR/MUR]
	Δ_vH	(500–616)	60.3	515	I	[1923MOR/MUR, 1984BOU/FRI]
	Δ_vH	(496–614)	59.6	555	I	[1922NEL/SEN]
	Δ_vH	(496–614)	60.7	511	I	[1922NEL/SEN, 1984BOU/FRI]
C₁₄D₁₀	[1719-06-8]	anthracene-d ₁₀				
	Δ_vH		78.4	298	CGC	[2008ZHA/UNH]
C₁₄H₁₀	[85-01-8]	phenanthrene				
	$\Delta_{\text{fus}}H$		18.1	na	DSC	[2003SHA/KAN]
	$\Delta_{\text{fus}}H$	(353–383)	16.6	367.6	DSC	[2003ROJ/ORO]
	$\Delta_{\text{trs}}H$		0.22	347.5		
	$\Delta_{\text{fus}}H$		16.2	372.9	DSC	[2000LIS/JAM]
	$\Delta_{\text{fus}}H$		16.46	372.4		[1996DOM/HEA]
	$\Delta_{\text{sub}}H$	(313–333)	91.6 ± 0.4	323	ME	[2006RIB/MON]
	$\Delta_{\text{sub}}H$	(313–333)	92.5 ± 0.4	298	ME	[2006RIB/MON]
	$\Delta_{\text{sub}}H$		89.6 ± 0.8	298	DSC	[2003ROJ/ORO]
	$\Delta_{\text{sub}}H$		92 ± 1		LE	[1998PRI/HAW]
	$\Delta_{\text{sub}}H$		90.5	298	CGC-DSC	[1998CHI/HES]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{sub}}H$	(303–333)	95.0 ± 4.4	318	ME	[1998OJA/SUU]
	$\Delta_{\text{sub}}H$	(313–453)	88.9	383	GS	[1995NAS/LEN]
	$\Delta_{\text{sub}}H$		87.2 ± 1.1	350	DSC	[1988TOR/BAR]
	$\Delta_{\text{sub}}H$		90.9 ± 1.7	298	DSC	[1988TOR/BAR]
	$\Delta_{\text{sub}}H$	(323–348)	96.2	335	GS	[1986SAT/INO]
	$\Delta_{\text{sub}}H$	(317–362)	82 ± 2	340	TE	[1983FER/IMP]
	$\Delta_{\text{sub}}H$	(283–323)	95.0 ± 0.6	303	GS	[1983SON/ZOL]
	$\Delta_{\text{sub}}H$	(315–335)	92.5 ± 2	298	TE,ME	[1980DEK]
	$\Delta_{\text{sub}}H$	(325–364)	87.2	345	GS	[1979MAC/PRA]
	$\Delta_{\text{sub}}H$		87.2	372	B	[1975OSB/DOU]
	$\Delta_{\text{sub}}H$	(300–330)	87.4 ± 0.8	298	TE	[1975DEK/VAN]
	$\Delta_{\text{sub}}H$	(312–326)	86.6 ± 0.8	298	TCM	[UR/DEL, 1975DEK/VAN]
	$\Delta_{\text{sub}}H$		90.9 ± 0.4	298	C	[1972MOR, 1977PED/RYL]
	$\Delta_{\text{sub}}H$	(279–315)	84.1 ± 2.5	297	TE	[1960BUD]
	$\Delta_{\text{sub}}H$	(273–333)	95.9	303		[1958HOY/PEP, 1970COX/PIL]
	$\Delta_{\text{sub}}H$	(310–323)	86.6			[1953BRA/CLE2, 1970COX/PIL]
	$\Delta_{\text{sub}}H$					[1960JON]
	$\Delta_{\text{sub}}H$		90.7 ± 1.2	315	ME	[1952INO/SHI]
	$\Delta_{\text{sub}}H$		81.6	323	ME	[1951INO]
	$\Delta_{\text{sub}}H$		92.9			[1949KLA, 1970COX/PIL]
	$\Delta_{\text{sub}}H$		84.1 ± 0.8	313		[1938WOL/WEG]
	Δ_vH		78.7	298	CGC	[2008ZHA/UNH]
	Δ_vH	(413–483)	79.0 ± 1.2	298	GC	[2006HAF/PAR]
	Δ_vH	(323–473)	72.2	398	GC	[2002LEI/CHA]
	Δ_vH		78.7	298	CGC	[1998CHI/HES]
	Δ_vH		72.5		GC	[1996GOV/RUT]
	Δ_vH	(403–453)	78.5	298	CGC	[1995CHI/HOS]
	Δ_vH	(343–453)	71.2	398	GC	[1990HIN/BID2]
	Δ_vH	(391–613)	58.2	406	A	[1987STE/MAL]
	Δ_vH	(373–423)	69.6	388	A	[1987STE/MAL, 1975OSB/DOU]
	Δ_vH		71.2	372		[1977FIN/MES]
	Δ_vH		69.7	390		[1977FIN/MES]
	Δ_vH		67.5	420		[1977FIN/MES]
	Δ_vH	(476–620)	57.2	548	I	[1923MOR/MUR]
	Δ_vH	(476–620)	61.2	491	I	[1923MOR/MUR, 1984BOU/FRI]
	Δ_vH	(505–614)	59.3	560	I	[1922NEL/SEN]
	Δ_vH	(505–614)	61.2	520	I	[1922NEL/SEN, 1984BOU/FRI]
C₁₄D₁₀	[1517-22-2]	phenanthrene-d ₁₀				
	$\Delta_{\text{sub}}H$	(283–323)	92.2 ± 1.1	303	GS	[1983SON/ZOL]
	Δ_vH		78.6	298	CGC	[2008ZHA/UNH]
C₁₄H₁₀	[501-65-5]	diphenylacetylene				
	$\Delta_{\text{fus}}H$		21.5	335	DSC	[2002STE/CHI3]
	$\Delta_{\text{fus}}H$		20.5	334		[1996DOM/HEA]
	$\Delta_{\text{sub}}H$		95.3	298	CGC-DSC	[1998CHI/HES]
	$\Delta_{\text{sub}}H$	(298–316)	95.1 ± 1.1	298	ME	[1993DIO/MIN]
	$\Delta_{\text{sub}}H$	(299–321)	90.0 ± 4.5	310	HSA	[1986CHI/ANN]
	$\Delta_{\text{sub}}H$	(299–321)	88.7 ± 1.25	313		[1938WOL/WEG, 1938WEG, 1960JON]
	Δ_vH	(439–517)	63.8 ± 0.2	440	EB	[2002STE/CHI3]
	Δ_vH	(439–517)	60.9 ± 0.2	480	EB	[2002STE/CHI3]
	Δ_vH	(439–517)	58.1 ± 0.3	520	EB	[2002STE/CHI3]
C₁₄H₁₀Cl₂O₂	[83-05-6]	bis(4-chlorophenyl)acetic acid				

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{fus}}H$		31.66	440.2	DSC	[1991ACR, 1990DON/DRE]
C ₁₄ H ₁₀ Cl ₄	[72-54-8]	1,1-dichloro-2,2-bis(4-chlorophenyl)ethane p,p'-DDD				
	$\Delta_{\text{fus}}H$		27.31	382.1	DSC	[1991ACR, 1990DON/DRE]
	Δ_vH	(343–453)	88.5	398	GC	[1990HIN/BID2]
C ₁₄ H ₁₀ Cl ₄	[121107-48-0]	(2,2',4,6'-tetrachloro-5-methyldiphenyl)methane				
	Δ_vH		98.6	298	GC	[1996VAN/VAN]
C ₁₄ H ₁₀ Cl ₄	[121107-46-8]	(2,2',4,5'-tetrachloro-5-methyldiphenyl)methane				
	Δ_vH		101	298	GC	[1996VAN/VAN]
	Δ_vH		92.4		GC	[1996GOV/RUT]
C ₁₄ H ₁₀ Cl ₄	[121107-54-8]	(2,2',5,5'-tetrachloro-4-methyldiphenyl)methane				
	Δ_vH		101.2	298	GC	[1996VAN/VAN]
	Δ_vH		92.6		GC	[1996GOV/RUT]
C ₁₄ H ₁₀ Cl ₄	[121107-44-6]	(2,2',4,4'-tetrachloro-5-methyldiphenyl)methane				
	Δ_vH		101.3	298	GC	[1996VAN/VAN]
	Δ_vH		92.8		GC	[1996GOV/RUT]
C ₁₄ H ₁₀ Cl ₄	[121107-47-9]	(2,2',4,6'-tetrachloro-3-methyldiphenyl)methane				
	Δ_vH		100.1	298	GC	[1996VAN/VAN]
C ₁₄ H ₁₀ Cl ₄	[121107-83-5]	(2',3,4,6'-tetrachloro-6-methyldiphenyl)methane				
	Δ_vH		101.1	298	GC	[1996VAN/VAN]
C ₁₄ H ₁₀ Cl ₄	[121107-43-5]	(2,2',4,4'-tetrachloro-3-methyldiphenyl)methane				
	Δ_vH		101.8	298	GC	[1996VAN/VAN]
	Δ_vH		93.0		GC	[1996GOV/RUT]
C ₁₄ H ₁₀ Cl ₄	[121107-65-1]	(2,3',4,4'-tetrachloro-5-methyldiphenyl)methane				
	Δ_vH		103.8	298	GC	[1996VAN/VAN]
C ₁₄ H ₁₀ Cl ₄	[121107-77-5]	(2'3,4,4'-tetrachloro-6-methyldiphenyl)methane				
	Δ_vH		103	298	GC	[1996VAN/VAN]
	Δ_vH		94.2		GC	[1996GOV/RUT]
C ₁₄ H ₁₀ F ₃ NO ₂	[530-78-9]	2-[[3-(trifluoromethyl)phenyl]amino]benzoic acid (flufenamic acid)				
	$\Delta_{\text{fus}}H$		26.7	405	DSC	[2007PER/SUR2, 2009SUR/TER]
	$\Delta_{\text{fus}}H$		27.0	407.3	DSC	[2004ROM/BUS]
	$\Delta_{\text{sub}}H$	(339–376)	119.4 ± 0.7	358	GS	[2007PER/SUR2, 2009SUR/TER]
	$\Delta_{\text{sub}}H$	(339–376)	121.2 ± 0.7	298	GS	[2007PER/SUR2, 2009SUR/TER]
C ₁₄ H ₁₀ F ₄	[425-32-1]	1,1,2,2-tetrafluoro-1,2-diphenylethane				
	$\Delta_{\text{fus}}H$		28.83	399.2		[1997SCH/VER]
	$\Delta_{\text{sub}}H$		101.8	298		[1997SCH/VER]
C ₁₄ H ₁₀ N ₂ O ₂	[128-95-0]	1,4-diaminoanthraquinone				
	$\Delta_{\text{fus}}H$		24.2	484.2		[1988BAU/PER]
	$\Delta_{\text{sub}}H$		143		GS	[1987SHI/OHK, 1991HOR]
	$\Delta_{\text{sub}}H$	(448–474)	151.2	461		[1987STE/MAL, 1980ROD/KRU]
	$\Delta_{\text{sub}}H$		136			[1984KAR/KRU]
	$\Delta_{\text{sub}}H$	(378–403)	102.6 ± 9.7	390		[1984KRI]
	$\Delta_{\text{sub}}H$	(473–553)	123	513	GS	[1977NIS/ISH, 1978NIS/ISH]
$\Delta_{\text{sub}}H$		199.2 ± 2.5		TE,ME	[1970KOJ]	

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{sub}}H$		123.4			[1968TSU/KOJ, 1988BAU/PER]
	$\Delta_{\text{sub}}H$		138.1		GS	[1967DAT/KAN, 1991HOR]
C ₁₄ H ₁₀ N ₂ O ₂	[129-44-2]	1,5-diaminoanthraquinone (405–427)	118.5 ± 4.8	416		[1984KRI]
C ₁₄ H ₁₀ N ₂ O ₂	[4870-16-0]	N-anilinophthalimide				
	$\Delta_{\text{fus}}H$		1.62	401		
	$\Delta_{\text{fus}}H$		26.9	457		[1998BOT/ELL]
C ₁₄ H ₁₀ N ₂ O ₃	[58658-02-9]	10-methyl-2-nitroacridin-9(10H)-one				
	$\Delta_{\text{fus}}H$		37.6	561	DSC	[2003STO/KRZ]
C ₁₄ H ₁₀ O	[90-44-8]	anthrone				
	$\Delta_{\text{fus}}H$		26.8	429		[1996DOM/HEA]
		Note: Some decomposition upon melting				
	$\Delta_{\text{sub}}H$		106.1 ± 0.8	298	GS	[1998VER4]
	$\Delta_{\text{sub}}H$		103.3	298		[1991ELW/SAB, 1992SAB/WAT]
	$\Delta_{\text{sub}}H$		99.6	354	C	[1991ELW/SAB]
C ₁₄ H ₁₀ O	[30084-90-3]	2-fluorencarboxaldehyde (338–356)	100.0 ± 3.4		ME	[2008GOL/SUU]
C ₁₄ H ₁₀ O ₂	[134-81-6]	benzil				
	$\Delta_{\text{fus}}H$		22.88	368.1		[2005FAT/KAS]
	$\Delta_{\text{fus}}H$		23.2	na	DSC	[2003SHA/KAN]
	$\Delta_{\text{fus}}H$		23.8	369.2	DSC	[2001RAI/VAR]
	$\Delta_{\text{fus}}H$		0.04	84.0		
	$\Delta_{\text{fus}}H$		23.56	368		[1996DOM/HEA]
	$\Delta_{\text{fus}}H$	(60–100)	0.05	84.1	AC	[1977DWO/FUC]
	$\Delta_{\text{sub}}H$	(319–340)	98.4 ± 1.1	329		[1959AIH, 1970COX/PIL, 1987STE/MAL]
	$\Delta_{\text{sub}}H$		82.8			[1938WOL/WEG, 1938WEG, 1960JON]
	Δ_vH	(401–620)	69.2	416	A	[1987STE/MAL, 1947STU]
C ₁₄ H ₁₀ O ₂	[1989-33-9]	9-fluorencarboxylic acid (349–418)	110.1 ± 4.6		ME	[2008GOL/SUU]
C ₁₄ H ₁₀ O ₃	[93-97-0]	benzoic acid anhydride				
	$\Delta_{\text{fus}}H$		17.15	313.2		[1971CAR/FIN]
	$\Delta_{\text{sub}}H$		96.2 ± 4.2	298	B	[1971CAR/FIN, 1977PED/RYL]
	$\Delta_{\text{sub}}H$		96.7 ± 4.2			[1947STU, 1970COX/PIL]
	Δ_vH	(416–633)	69.1	431	A	[1987STE/MAL, 1947STU]
C ₁₄ H ₁₀ O ₄	[94-36-0]	benzoyl peroxide				
	$\Delta_{\text{fus}}H$		22.59	378		[1967FIN/GRA]
		Note: Large uncertainty in reported value. Compound may undergo some decomposition upon melting.				
	$\Delta_{\text{sub}}H$	(310–340)	97.9 ± 2.5	298	ME	[1975CAR/LAY]
	$\Delta_{\text{sub}}H$	(293–313)	89.7 ± 4.2	303	ME	[1971KIP/RAB, 1977PED/RYL]
C ₁₄ H ₁₀ O ₄	[3155-16-6]	diphenyl oxalate				
	$\Delta_{\text{fus}}H$		31.38	403		[1971CAR/FIN]
	$\Delta_{\text{sub}}H$		102.5 ± 8.4		B	[1971CAR/FIN, 1977PED/RYL]
C ₁₄ H ₁₀ O ₄	[482-05-3]	2,2'-biphenyldicarboxylic acid				
	$\Delta_{\text{sub}}H$		151.9 ± 3.5	298	C	[2004MAT/MIR2]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{sub}}H$	(433–493)	166.1	448	A	[1987STE/MAL]
C ₁₄ H ₁₀ O ₄	[787-70-2]	4,4'-biphenyldicarboxylic acid				
	$\Delta_{\text{sub}}H$		196.4 ± 7.1	298	C	[2004MAT/MIR2]
C ₁₄ H ₁₀ O ₄	[40498-13-3]	2,3-dihydro-1,4-dihydroxy-9,10-anthraquinone				
	$\Delta_{\text{sub}}H$		110.7	363		[2003HIN/RAF]
C ₁₄ H ₁₀ O ₅	[962-16-3]	O-phenyl-O,O-benzoyl peroxycarbonate				
	$\Delta_{\text{sub}}H$		97.9 ± 2.5			[1975CAR/LAY, 1977PED/RYL]
	$\Delta_{\text{sub}}H$		133.9 ± 4.2		E	[1971KIP/RAB, 1977PED/RYL]
C ₁₄ H ₁₀ O ₅	[552-94-3]	salicylsalicylic acid				
	$\Delta_{\text{fus}}H$		29.0	430.2	DSC	[2004RAM/DIO]
C ₁₄ H ₁₁ BrN ₂ S	[109768-69-6]	N-(4-bromophenyl)-4 <i>H</i> -3,1-benzothiazin-2-amine				
	$\Delta_{\text{fus}}H$		24.5	478.3	DSC	[2004GON/KOS]
C ₁₄ H ₁₁ ClN ₂ S	[461662-90-8]	N-(4-chlorophenyl)-4 <i>H</i> -3,1-benzothiazin-2-amine				
	$\Delta_{\text{fus}}H$		27.5	476.4	DSC	[2004GON/KOS]
C ₁₄ H ₁₁ Cl ₂ NO ₂	[32809-16-8]	3-(3,5-dichlorophenyl)-1,5-dimethyl-3-azabicyclo[3.1.0]hexanedione				
	$\Delta_{\text{fus}}H$		30.09	438.2	DSC	[1990DON/DRE]
C ₁₄ H ₁₁ Cl ₂ NO ₂	[15307-86-5]	2-[(2,6-dichlorophenyl)amino]benzoic acid (diclofenac acid)				
	$\Delta_{\text{fus}}H$		40.4	452.6	DSC	[2009SUR/TER]
	$\Delta_{\text{fus}}H$		39.4	454.2	DSC	[2007PAS/BET]
	$\Delta_{\text{sub}}H$	(323–355)	114.7 ± 1.3	339	GS	[2007PER/SUR, 2009SUR/TER]
	$\Delta_{\text{sub}}H$	(323–355)	115.6 ± 1.3	298	GS	[2007PER/SUR, 2009SUR/TER]
C ₁₄ H ₁₁ FO ₃	[3119-88-8]	2'-fluoro-2-hydroxy-4-methoxybenzophenone				
	$\Delta_{\text{sub}}H$	(307–318)	109.3	312.5	EV	[1987STE/MAL, 1966GRA/BUR]
C ₁₄ H ₁₁ FO ₃	[3506-35-2]	3'-fluoro-2-hydroxy-4-methoxybenzophenone				
	$\Delta_{\text{sub}}H$	(322–343)	U 17.3	332.5	EV	[1987STE/MAL, 1966GRA/BUR]
C ₁₄ H ₁₁ FO ₃	[3602-47-9]	4'-fluoro-2-hydroxy-4-methoxybenzophenone				
	$\Delta_{\text{sub}}H$	(322–343)	U 37.7	332.5	EV	[1987STE/MAL, 1966GRA/BUR]
C ₁₄ H ₁₁ F ₃	[68936-77-6]	1,1,2-trifluoro-1,2-diphenylethane				
	$\Delta_{\text{fus}}H$		28.37	354.2		[1997SCH/VER]
	$\Delta_{\text{sub}}H$		93.1	298		[1997SCH/VER]
C ₁₄ H ₁₁ F ₃	[384-94-1]	1,1,1-trifluoro-2,2-diphenylethane				
	Δ_vH	(286–328)	69.1 ± 0.9	298	GS	[1997SCH/VER]
C ₁₄ H ₁₁ IO ₃ S	[313057-05-5]	4-(2-propenyloxy)phenyl 5-iodo-2-thiophene carboxylate				
	$\Delta_{\text{fus}}H$		83.68	383.2	DSC	[2000WU/WAN]
C ₁₄ H ₁₁ NO	[574-39-0]	N-acetylcarbazole				
	$\Delta_{\text{fus}}H$		15.1	349.9		[2001JAM/DOB]
C ₁₄ H ₁₁ NO	[719-54-0]	10-methylacridin-9(10 <i>H</i>)-one				
	$\Delta_{\text{fus}}H$		29.7	479	DSC	[2003STO/KRZ]
	$\Delta_{\text{sub}}H$		105		DSC	[2003STO/KRZ]
C ₁₄ H ₁₁ NO ₃	[na]	N-salicylidene- <i>m</i> -aminobenzoic acid				
	$\Delta_{\text{fus}}H$		33.11	464		[1996DOM/HEA]
C ₁₄ H ₁₁ NS	[150993-53-6]	2-cyanophenyl benzyl sulfide				

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{sub}}H$		117.8 ± 2.1	298	C	[2006MUL/MOZ]
C ₁₄ H ₁₁ N ₃ O ₂	[6407-69-8]	1,4,5-triaminoanthraquinone				
	$\Delta_{\text{sub}}H$	(473–553)	U 70.3	513	GS	[1977NIS/ISH, 1978NIS/ISH]
C ₁₄ H ₁₁ N ₃ O ₄	[1979-14-3]	2-cyano-6-nitro-1(2 <i>H</i>)-quinolinecarboxylic acid, 2-propenyl ester				
	$\Delta_{\text{fus}}H$		26.87	383.7	DSC	[2005LIZ/ZAB]
C ₁₄ H ₁₁ N ₃ O ₄	[na]	2-cyano-6-nitro-1(2 <i>H</i>)-quinolinecarboxylic acid, 1-propenyl ester				
	$\Delta_{\text{fus}}H$		26.27	389.4	DSC	[2005LIZ/ZAB]
C ₁₄ H ₁₂	[1730-37-6]	1-methylfluorene				
	$\Delta_{\text{sub}}H$	(285–317)	91.2 ± 0.4	298	GS	[2004VER]
	Δ_vH		77.2 ± 3.6	298	CGC	[2008HAN/NUT]
	Δ_vH	(361–375)	78.7 ± 0.7	298	GS	[2004VER]
C ₁₄ H ₁₂		(323–473)	71.1	398	GC	[2002LEI/CHA]
	[2523-37-7]	9-methylfluorene				
	$\Delta_{\text{fus}}H$		16.32	319.2	DSC	[1994RAK/VER2]
	$\Delta_{\text{sub}}H$	(285–317)	83.7 ± 0.6	298	GS	[2004VER]
	$\Delta_{\text{sub}}H$	(318–358)	82.8 ± 0.3	338	B	[1994RAK/VER2]
	$\Delta_{\text{sub}}H$		82.8 ± 0.3	298		[1994RAK/VER2]
	Δ_vH	(320–353)	70.6 ± 0.3	298	GS	[2004VER]
C ₁₄ H ₁₂		(318–358)	71.3 ± 0.2	298	GS	[2004VER]
		(318–358)	66.5	298	B	[1994RAK/VER2]
	[613-31-0]	9,10-dihydroanthracene				
	$\Delta_{\text{sub}}H$	(313–453)	93.9	383	GS	[1995NAS/LEN]
	$\Delta_{\text{sub}}H$	(318–379)	92.4 ± 4		ME	[1975MAL/GIG, 1987STE/MAL]
	$\Delta_{\text{sub}}H$		94.2 ± 0.8	298	ME	[1975MAL/GIG]
C ₁₄ H ₁₂		(279–328)	93.3 ± 4	304		[1958HOY/PEP, 1970COX/PIL]
	$\Delta_{\text{sub}}H$		89.5	388		[1951MAG/HAR, 1960JON]
	[776-35-2]	9,10-dihydrophenanthrene				
	$\Delta_{\text{fus}}H$		12.8	306.5	AC	[1996DOM/HEA, 1979LEE/HOS]
	Δ_vH	(417–453)	64.0	432	A	[1987STE/MAL]
C ₁₄ H ₁₂		(353–418)	72.3 ± 0.6	340	IP	[1979LEE/HOS]
		(353–418)	76.6 ± 0.1	298	IP	[1979LEE/HOS]
	[530-48-3]	1,1-diphenylethylene				
	Δ_vH	(298–331)	70.2 ± 0.7	314	GS	[1999VER/EBE]
C ₁₄ H ₁₂		(298–331)	71.2 ± 0.7	298	GS	[1999VER/EBE]
	Δ_vH	(360–550)	59.3	375	A	[1987STE/MAL, 1947STU]
	[645-49-8]	<i>cis</i> 1,2-diphenylethylene (<i>cis</i> stilbene)				
C ₁₄ H ₁₂		(308–343)	70.5 ± 0.4	298	GS	[2009CAM/EME]
	Δ_vH	(373–428)	66.5	388	A	[1987STE/MAL]
	Δ_vH	(276–286)	50.3 ± 1.0	298		[1952BRA/PLE, 2009CAM/EME]
C ₁₄ H ₁₂	[103-30-0]	<i>trans</i> 1,2-diphenylethylene (<i>trans</i> stilbene)				
	$\Delta_{\text{fus}}H$		27.4	398.2		[1991ACR]
	$\Delta_{\text{fus}}H$		27.7	397.4	AC	[1985BOU/DEL]
	$\Delta_{\text{sub}}H$	(324–367)	102 ± 0.4	298	GS	[2009CAM/EME]
	$\Delta_{\text{sub}}H$		102	298	CGC-DSC	[1998CHI/HES]
	$\Delta_{\text{sub}}H$	(298–343)	99.6	313	A	[1987STE/MAL]
	$\Delta_{\text{sub}}H$		U 61.1		MS	[1983MAJ/AZZ]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{sub}}H$	(293–338)	103.8 ± 2.5	315		[1983KRA/BEC]
	$\Delta_{\text{sub}}H$		100.7 ± 0.4	298	SRFG	[1983VAN/JAC]
	$\Delta_{\text{sub}}H$	(310–340)	99.6 ± 1.7	298	TE	[1975DEK/VAN]
	$\Delta_{\text{sub}}H$		102.1 ± 0.6		TCM	[1973DEK/OON]
	$\Delta_{\text{sub}}H$		99.2 ± 0.4			[1972MOR3]
	$\Delta_{\text{sub}}H$	(303–315)	86.5 ± 0.1	309	TM	[1955ENG]
	Δ_vH		79.7	298	CGC	[1998CHI/HES]
	Δ_vH	(453–503)	79.8	298	CGC	[1995CHI/HOS]
	Δ_vH	(403–453)	79.6	298	CGC	[1995CHI/HOS]
	Δ_vH	(419–580)	65.5	434	A	[1987STE/MAL]
C₁₄H₁₂BrNOS	[127351-08-0]	3-bromo-N-(4-methoxyphenyl)benzenecarbothioamide				
	$\Delta_{\text{fus}}H$ (I)		28.5	376.7		
	$\Delta_{\text{fus}}H$ (II)		22.6	373.2		[2007BAS/AND]
C₁₄H₁₂ClN	[33442-36-3]	4-chlorobenzylidene-4'-methylaniline				
	$\Delta_{\text{fus}}H$		25.58	400.4	DSC	[1999GAL/COL]
C₁₄H₁₂ClNO₂	[13710-19-5]	2-[(3-chloro-2-methylphenyl)amino]benzoic acid (tolfenamic acid)				
	$\Delta_{\text{fus}}H$ (white crys)		41.0	484.2		
	$\Delta_{\text{fus}}H$ (yellow crys)		49.0	485.8	DSC	[2009SUR/SZT]
	$\Delta_{\text{fus}}H$		38.6	484.3	DSC	[2009SUR/TER]
	$\Delta_{\text{fus}}H$		41.2	485.3		[2007BER/WAS]
	$\Delta_{\text{sub}}H$	(346–373)	125.7 ± 0.8	360	GS	[2009SUR/TER]
	$\Delta_{\text{sub}}H$	(346–373)	128.4 ± 0.8	298	GS	[2009SUR/TER]
C₁₄H₁₂F₂	[350-62-9]	1,1-difluoro-1,2-diphenylethane				
	$\Delta_{\text{fus}}H$		24.35	339.2		[1997SCH/VER]
	$\Delta_{\text{sub}}H$		94.7 ± 0.9	298		[1997SCH/VER]
C₁₄H₁₂F₃NO₄S₂	[37924-13-3]	1,1,1-trifluoro-N-[2-methyl-4-(phenylsulphonyl)phenyl]methanesulfonamide				
	$\Delta_{\text{fus}}H$		31.79	418.4	DSC	[1990DON/DRE]
C₁₄H₁₂N₂	[22739-29-3]	N-methyl-9-acridinamine				
	$\Delta_{\text{sub}}H$		107	480	TGA	[1998STO/KRZ]
C₁₄H₁₂N₂	[5291-44-1]	10-methyl-9-acridinimine				
	$\Delta_{\text{sub}}H$		94	550	TGA	[1998STO/KRZ]
C₁₄H₁₂N₂	[588-68-1]	dibenzylideneazaine				
	$\Delta_{\text{sub}}H$		93.3 ± 2.1	293	EST	[1948COA/SUT]
C₁₄H₁₂N₂	[484-11-7]	2,9-dimethyl-1,10-phenanthroline				
	$\Delta_{\text{fus}}H$		17.6	435.9		[2007BON/CAT]
C₁₄H₁₂N₂	[621-72-7]	2-benzylbenzimidazole				
	$\Delta_{\text{sub}}H$	(393–412)	134.5 ± 0.5	403	ME	[2005RIB/RIB]
	$\Delta_{\text{sub}}H$	(393–412)	136.2 ± 0.5	298	ME	[2005RIB/RIB]
C₁₄H₁₂N₂O₂	[192998-96-2]	<i>cis</i> -5a,6,11a,12-tetrahydro[1,4]benzothiazino[3,2-b][1,4]-benzoxazine				
	$\Delta_{\text{sub}}H$	(383–392)	122	387	ME	[1997GUD/TOR]
	$\Delta_{\text{sub}}H$		129.0 ± 1.3	298		[1997GUD/TOR]
C₁₄H₁₂N₂O₂	[na]	4-nitro-4'-methylbenzylidene aniline				
	$\Delta_{\text{fus}}H$		27.3	402		[1997KER/LOC]
C₁₄H₁₂N₂O₄	[42472-93-5]	N-methylthalidomide				
	$\Delta_{\text{fus}}H$		18.12	432.2	DTA	[2002GOO/LAI]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₄ H ₁₂ N ₂ S ₂	[165454-33-1]	<i>cis</i> -5a,6,11a,12-tetrahydro[1,4]benzothiazino[3,2-b]-[1,4]-benzothiazine				
	$\Delta_{\text{sub}}H$	(383–392)	118	387	ME	[1997GUD/TOR]
	$\Delta_{\text{sub}}H$		123.3 ± 1.2	298		[1997GUD/TOR]
C ₁₄ H ₁₂ N ₄ O ₂	[2475-45-8]	1,4,5,8-tetraminoanthraquinone				
	$\Delta_{\text{sub}}H$	(473–553)	U 82	513	GS	[1977NIS/ISH, 1978NIS/ISH]
C ₁₄ H ₁₂ O	[451-40-1]	benzyl phenyl ketone				
	Δ_vH	(396–594)	68.1	411	A	[1987STE/MAL, 1947STU]
C ₁₄ H ₁₂ O	[131-58-8]	2-methylbenzophenone				
	Δ_vH	(435–580)	65.1	450	A	[1987STE/MAL]
C ₁₄ H ₁₂ O	[643-65-2]	3-methylbenzophenone				
	Δ_vH	(445–585)	68.4	460	A	[1987STE/MAL]
C ₁₄ H ₁₂ O	[134-84-9]	4-methylbenzophenone				
	Δ_vH	(450–492)	72.0	465	A	[1987STE/MAL]
C ₁₄ H ₁₂ O	[451-40-1]	desoxybenzoin				
	$\Delta_{\text{sub}}H$		99.3 ± 4.2			[1947STU, 1970COX/PIL]
C ₁₄ H ₁₂ O	[24324-17-2]	9-fluorenylmethanol				
	$\Delta_{\text{fus}}H$	(78–390)	26.27	376.6	AC	[2004DI/TAN]
C ₁₄ H ₁₂ O ₂	[579-44-2]	<i>dl</i> benzoin				
	Δ_vH		98.5 ± 12.5	298	CGC	[2006PER/CON]
	Δ_vH	(408–616)	69.0	423	A	[1987STE/MAL, 1947STU]
C ₁₄ H ₁₂ O ₂	[120-51-4]	benzyl benzoate				
	$\Delta_{\text{fus}}H$		20.44	293.1	DSC	[1990DON/DRE]
	Δ_vH	(497–602)	59.7	512	A, EB	[1987STE/MAL, 1976HON/SIN]
	Δ_vH	(297–353)	77.7	312	A, ME	[1987STE/MAL, 1957SER/VOI]
C ₁₄ H ₁₂ O ₂	[117-34-0]	diphenylacetic acid				
	$\Delta_{\text{fus}}H$		31.18	420.4		[2010CHA/LAY]
	$\Delta_{\text{fus}}H$		31.25	420.4		[1996DOM/HEA]
C ₁₄ H ₁₂ O ₂	[na]	<i>dl</i> 1,2-diphenyl-1,2-dihydroxyethane				
	$\Delta_{\text{fus}}H$		31.38	393		[1976LEC/COL]
C ₁₄ H ₁₂ O ₂	[na]	<i>d</i> 1,2-diphenyl-1,2-dihydroxyethane				
	$\Delta_{\text{fus}}H$		34.31	420.5		[1976LEC/COL]
C ₁₄ H ₁₂ O ₂	[2553-04-0]	(2-methoxyphenyl)phenylmethanone				
	$\Delta_{\text{fus}}H$		0.68	350.2	DTA	[1989SAL/ABA]
		Note: Reported enthalpy of fusion is too small, and the published enthalpy and entropy of fusion data are internally inconsistent				
C ₁₄ H ₁₂ O ₂ S	[16212-06-9]	E-(2-phenylethenyl)sulfonyl benzene (phenyl <i>trans</i> - β -styrylsulfone)				
	$\Delta_{\text{sub}}H$		105 ± 3.8		B	[1969MAC/MCN, 1969MAC/MCN2, 1977PED/RYL]
C ₁₄ H ₁₂ O ₃	[118-58-1]	benzyl salicylate				
	Δ_vH	(295–334)	78.7	310	A, ME	[1987STE/MAL, 1955SER/VOI]
C ₁₄ H ₁₂ O ₃	[131-57-7]	2-hydroxy-4-methoxybenzophenone				
	$\Delta_{\text{fus}}H$		21.77	336.7		[2008LAG/JIM]
	$\Delta_{\text{sub}}H$	(281–337)	118.9	296	A	[1987STE/MAL]
		(308–323)	U 39.7	315	EV	[1966GRA/BUR]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(337–413)	74.7	352	A, UV	[1987STE/MAL, 1960SCH/HIR]
C ₁₄ H ₁₂ O ₃	[na] $\Delta_{\text{fus}} H$	2-pivaloylindan-1,3-dione	25.99	381.5		[1991ACR]
C ₁₄ H ₁₂ O ₄	[131-53-3] $\Delta_{\text{fus}} H$	2,2'-dihydroxy-4-methoxybenzophenone	22.0	343	DSC	[1999PRI/HAWN]
	$\Delta_{\text{sub}} H$		103.8		B	[1999PRI/HAWN]
	$\Delta_{\text{sub}} H$	(303–342)	228	318	A	[1987STE/MAL]
	$\Delta_v H$	(342–481)	75.6	357	A, UV	[1987STE/MAL, 1960SCH/HIR]
C ₁₄ H ₁₂ O ₄	[131-53-3] $\Delta_{\text{fus}} H$	2,4-dihydroxy-4'-methoxybenzophenone	35.6	436.8	DSC	[1999PRI/HAWN]
	$\Delta_{\text{sub}} H$		138.3		B	[1999PRI/HAWN]
C ₁₄ H ₁₂ O ₄	[na] $\Delta_{\text{fus}} H$	1,2-dicarbomethoxynaphthalene	27.6	358.2	DSC	[1993ACR]
C ₁₄ H ₁₂ O ₄	[18713-38-7] $\Delta_{\text{fus}} H$	1,3-dicarbomethoxynaphthalene	30.5	378.7	DSC	[1993ACR]
C ₁₄ H ₁₂ O ₄	[7487-15-2] $\Delta_{\text{fus}} H$	1,4-dicarbomethoxynaphthalene	20.4	340.2	DSC	[1993ACR]
C ₁₄ H ₁₂ O ₄	[19458-95-8] $\Delta_{\text{fus}} H$	1,5-dicarbomethoxynaphthalene	26.4	392	DSC	[1993ACR]
C ₁₄ H ₁₂ O ₄	[16144-94-8] $\Delta_{\text{fus}} H$	1,6-dicarbomethoxynaphthalene	22.1	371.8	DSC	[1993ACR]
C ₁₄ H ₁₂ O ₄	[68267-12-9] $\Delta_{\text{fus}} H$	1,7-dicarbomethoxynaphthalene	20.0	363.2	DSC	[1993ACR]
C ₁₄ H ₁₂ O ₄	[13728-34-2] $\Delta_{\text{fus}} H$	2,3-dicarbomethoxynaphthalene	20.2	324.2	DSC	[1993ACR]
C ₁₄ H ₁₂ O ₄	[2549-47-5] $\Delta_{\text{fus}} H$	2,7-dicarbomethoxynaphthalene	26.6	410.2	DSC	[1993ACR]
C ₁₄ H ₁₂ O ₅	[82-02-0] $\Delta_{\text{fus}} H$ (I) $\Delta_{\text{fus}} H$ (II)	4,9-dimethoxy-7-methyl-5H-furo[3,2g][1]benzopyran-5-one (khellin)	27.9 32.32	423.5 426.5		[1979MAS/MAL]
C ₁₄ H ₁₃ N	[na] $\Delta_v H$ $\Delta_v H$	N-benzybenzaldehyde-imine	83.4 ± 1.2 85.0 ± 1.2	324 298	GS GS	[1997VER/MOR] [1997VER/MOR]
C ₁₄ H ₁₃ N	[86-28-2] $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$	N-ethylcarbazole	98.4 ± 0.3 99.1 ± 0.3	319 298	ME ME	[1990JIM/ROU] [1990JIM/ROU]
C ₁₄ H ₁₃ NO	[519-87-9] $\Delta_{\text{fus}} H$ $\Delta_{\text{sub}} H$	N,N-diphenylacetamide	23.4 122.7	374.4 358		[2001JAM/DOB] [1987STE/MAL]
C ₁₄ H ₁₃ NO ₂	[3585-93-1] $\Delta_{\text{sub}} H$	N-(4-methoxyphenylmethylene) benzenamine N-oxide	130.6 ± 1.2	298	C	[1986KIR/ACR]
C ₁₄ H ₁₃ N ₃ O ₄	[1979-08-5]	2-cyano-6-nitro-1(2H)-quinolinecarboxylic acid, propyl ester				

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{fus}}H$		24.99	372.2	DSC	[2005LIZ/ZAB]
C ₁₄ H ₁₃ N ₃ O ₄	[1979-12-1]	2-cyano-6-nitro-1(2 <i>H</i>)-quinolinecarboxylic acid, 1-methylethyl ester				
	$\Delta_{\text{fus}}H$		27.75	402	DSC	[2005LIZ/ZAB]
C ₁₄ H ₁₃ N ₃ O ₄ S	[71125-38-7]	4-hydroxy-2-methyl-N-(5-methyl-2-thiazolyl)-2 <i>H</i> -1,2-benzothiazine-3-carboxamide-1,1-dioxide (meloxicam)				
	$\Delta_{\text{fus}}H$		71.73	530	DSC	[2007BAB/SUB]
C ₁₄ H ₁₄	[620-83-7]	(4-methylphenyl)phenylmethane				
	Δ_vH	(293–333)	68.6 ± 0.3	313	GS	[1999VER5]
	Δ_vH	(293–333)	69.5 ± 0.3	298	GS	[1999VER5]
C ₁₄ H ₁₄	[605-39-0]	2,2'-dimethylbiphenyl				
	$\Delta_{\text{fus}}H$		2.28	293.1		[1996DOM/HEA]
C ₁₄ H ₁₄		(283–288)	65.7	285	ME	[1974PRI/POU, 1987STE/MAL]
	$\Delta_{\text{sub}}H$					
	Δ_vH	(288–308)	71.9	298	ME	[1974PRI/POU]
C ₁₄ H ₁₄		(288–308)	71.9	298	A	[1987STE/MAL, 1974PRI/POU]
	$\Delta_{\text{sub}}H$					
	Δ_vH					
C ₁₄ H ₁₄	[613-33-2]	4,4'-dimethylbiphenyl				
	$\Delta_{\text{sub}}H$		95.1 ± 2.0	298	C	[1997RIB/MAT4]
C ₁₄ H ₁₄	[1812-51-7]	2-ethylbiphenyl				
	$\Delta_{\text{fus}}H$		2.07	267.1		[1996DOM/HEA]
C ₁₄ H ₁₄	[612-00-0]	1,1-diphenylethane				
	Δ_vH	(293–328)	68.2 ± 0.6	313	GS	[1999VER5]
	Δ_vH	(293–328)	68.9 ± 0.6	298	GS	[1999VER5]
	Δ_vH	(348–405)	62.4	363	A	[1987STE/MAL]
C ₁₄ H ₁₄	[103-29-7]	1,2-diphenylethane				
	$\Delta_{\text{trs}}H$		2.25	273.2		
	$\Delta_{\text{fus}}H$		22.73	324.3		[1996DOM/HEA]
	$\Delta_{\text{sub}}H$	(293–323)	92.9	308	EM	[1989SAS/NGU]
	$\Delta_{\text{sub}}H$	(273–318)	91.2 ± 0.4	295		[1983KRA/BEC]
	$\Delta_{\text{sub}}H$		91.5 ± 0.7	298	B	[1980OSB/SCO]
	$\Delta_{\text{sub}}H$		91.4 ± 0.5	298	C	[1972MOR]
	$\Delta_{\text{sub}}H$	(286–307)	84.1 ± 0.4		V	[1959AIH, 1970COX/PIL]
	$\Delta_{\text{sub}}H$	(290–317)	72.4 ± 1.3	304	ME	[1951BRI]
	$\Delta_{\text{sub}}H$		73.2			[1938WOL/WEG, 1960JON, 1938WEG]
	Δ_vH	(323–473)	67.4	398	GC	[2002LEI/CHA]
	Δ_vH		66.2 ± 0.2	340		[1988MES/FIN]
Δ_vH	(333–413)	64.1	373		[1989SAS/NGU]	
Δ_vH	(359–557)	57	374	A	[1987STE/MAL, 1947STU]	
C ₁₄ H ₁₄	[2141-42-6]	1,2,3,4-tetrahydroanthracene				
	$\Delta_{\text{trs}}H$		19.16	373.3		
	$\Delta_{\text{fus}}H$		2.92	388		[1996DOM/HEA]
C ₁₄ H ₁₄	[1013-08-7]	1,2,3,4-tetrahydrophenanthrene				
	$\Delta_{\text{fus}}H$		11.17	302.6		
	$\Delta_{\text{trs}}H$		0.10	285		
	$\Delta_{\text{trs}}H$		1.77	298		[1994CHI/GAM]
C ₁₄ H ₁₄	[1857-75-6]	1,2- <i>cis</i> -dimethylacenaphthene				
	$\Delta_{\text{fus}}H$		22.59	325.2		[1974CAN/JAC]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₄ H ₁₄	[51921-69-8] $\Delta_{\text{fus}}H$	1,2- <i>trans</i> -dimethylacenaphthene	18.83	350.2		[1974CAN/JAC]
C ₁₄ H ₁₄	[56137-64-5] $\Delta_{\text{fus}}H$	1,3-dimethylacenaphthene	12.55	283.2		[1974CAN/JAC]
C ₁₄ H ₁₄	[56137-75-8] $\Delta_{\text{fus}}H$	1,4-dimethylacenaphthene	17.57	279.7		[1974CAN/JAC]
C ₁₄ H ₁₄	[56137-80-5] $\Delta_{\text{fus}}H$	1,5-dimethylacenaphthene	22.18	316.7		[1974CAN/JAC]
C ₁₄ H ₁₄	[56137-89-4] $\Delta_{\text{fus}}H$	1,7-dimethylacenaphthene	14.23	288.2		[1974CAN/JAC]
C ₁₄ H ₁₄	[56137-90-7] $\Delta_{\text{fus}}H$	1,8-dimethylacenaphthene	14.23	289.2		[1974CAN/JAC]
C ₁₄ H ₁₄	[56137-94-1] $\Delta_{\text{fus}}H$	3,4-dimethylacenaphthene	17.57	357.2		[1974CAN/JAC]
C ₁₄ H ₁₄	[56137-95-2] $\Delta_{\text{fus}}H$	3,8-dimethylacenaphthene	17.99	341.7		[1974CAN/JAC]
C ₁₄ H ₁₄	[56137-98-5] $\Delta_{\text{fus}}H$	4,7-dimethylacenaphthene	15.06	314.2		[1974CAN/JAC]
C ₁₄ H ₁₄	[56138-04-6] $\Delta_{\text{fus}}H$	5,6-dimethylacenaphthene	21.76	443.2		[1974CAN/JAC]
C ₁₄ H ₁₄ ClN ₂ O ₂	[457899-89-7] $\Delta_{\text{fus}}H$	4-chloro-2'-hydroxy-4'-ethoxyazobenzene	34.3	421	DSC	[2003PAJ/ROS]
C ₁₄ H ₁₄ ClN ₃ S	[436847-00-6] $\Delta_{\text{fus}}H$	N-2-(4,6-lutidyl)-N'-(2-chlorophenyl) thiourea	42.2	467.2	DSC	[2002KEL/SZC]
C ₁₄ H ₁₄ ClN ₃ S	[436847-02-8] $\Delta_{\text{fus}}H$	N-2-(4,6-lutidyl)-N'-(4-chlorophenyl) thiourea	66.5	499.7	DSC	[2002SZC/KEL]
C ₁₄ H ₁₄ Cl ₂ N ₂ O	[35554-44-0] $\Delta_{\text{fus}}H$	1-[2-(2,4-dichlorophenyl)-2-(propenyloxy)ethyl]-1 <i>H</i> -imidazole	30.5	322.6	DSC	[1990DON/DRE]
C ₁₄ H ₁₄ FN ₃	[150-74-3] $\Delta_{\text{sub}}H$	N,N-dimethyl-4-[(fluorophenyl)azo]benzenamine	91.2		UV	[1984KAR/ROD]
C ₁₄ H ₁₄ FN ₃ O ₂ S	[4644-89-7] $\Delta_{\text{sub}}H$	4-[[4-(dimethylamino)phenyl]azo]benzenesulfonyl fluoride	105.6		UV	[1984KAR/ROD]
C ₁₄ H ₁₄ F ₃ NO ₂	[41934-47-8] $\Delta_{\text{fus}}H$	4-trifluoromethyl-7-(N,N-diethylamino)coumarin	23.3	360	DSC	[1991ZHA/HUA]
C ₁₄ H ₁₄ NO ₃	[2643-00-7] $\Delta_{\text{sub}}H$	<i>bis</i> (4-methoxyphenyl)nitrogen oxide (328–363)	100.7	343	A	[1987STE/MAL, 1965KAL/ROZ]
C ₁₄ H ₁₄ NO ₄ PS	[2104-64-5] $\Delta_{\text{fus}}H$	O-ethyl O-(4-nitrophenyl)phenylphosphonothioate	25.05	308.2	DSC	[1990DON/DRE]
C ₁₄ H ₁₄ N ₂	[621-09-0] $\Delta_{\text{sub}}H$	(343–383)	122.6 ± 3.8	363	ME	[1958DUN/HAN]
C ₁₄ H ₁₄ N ₂ O	[1562-94-3] $\Delta_{\text{sub}}H$	<i>p</i> -azoxyanisole	134.8 ± 3.7	298	C	[1993ACR/TUC]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₄ H ₁₄ N ₂ O ₂	[na] $\Delta_{\text{sub}}H$	4-(2-hydroxyethoxy)azobenzene	120.9		GS	[1956MAJ2, 1991HOR]
C ₁₄ H ₁₄ N ₂ O ₂	[na] $\Delta_{\text{fus}}H$	N-methyl-N-nitro-4-(phenylmethyl)benzenamine	21.7	329.6		[2002DAS/ZAL]
C ₁₄ H ₁₄ N ₂ O ₃	[1562-94-3] Δ_vH	4,4'-dimethoxyazoxybenzene (395–418)	73.7	406	A	[1987STE/MAL]
C ₁₄ H ₁₄ N ₂ O ₃	[57721-89-8] $\Delta_{\text{fus}}H$	2-cyano-6-methoxy-1(2H)-quinolinecarboxylic acid, ethyl ester	22.37	359.1	DSC	[2005LIZ/ZAB]
C ₁₄ H ₁₄ N ₄ O ₂	[3837-55-6] $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$	3-nitro-4'-(N,N-dimethylamino)-azobenzene (388–412) (392–410)	133.9 ± 3.8 133.1 ± 3.8	400 401	ME TE	[1967GRE/JON] [1967GRE/JON, 1987STE/MAL]
C ₁₄ H ₁₄ N ₄ O ₂	[2491-74-9] $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$	4-nitro-4'-(N,N-dimethylamino)-azobenzene (413–425) (414–428)	134.3 ± 7.5 135.1 ± 0.9	419 421	ME TE	[1967GRE/JON, 1966JON/KRA] [1967GRE/JON, 1987STE/MAL]
C ₁₄ H ₁₄ O	[103-50-4] Δ_vH Δ_vH	dibenzyl ether (275–417) (413–461)	45.6 59.4	290 428	A A	[1987STE/MAL] [1987STE/MAL]
C ₁₄ H ₁₄ O	[59502-28-2] Δ_vH	isopropyl 2-naphthyl ketone (406–586)	75.9	421	A	[1987STE/MAL, 1947STU]
C ₁₄ H ₁₄ O	[52857-29-1] Δ_vH Δ_vH	2-(1-phenylethyl)phenol (443–521) (442–523)	82.8 72.8	458 482	A	[1987STE/MAL] [1939GOL/MAR]
C ₁₄ H ₁₄ O	[1988-89-2] Δ_vH Δ_vH	4-(1-phenylethyl)phenol (447–517) (447–523)	90.8 75.4	462 485	A	[1987STE/MAL] [1939GOL/MAR]
C ₁₄ H ₁₄ O	[599-67-7] $\Delta_{\text{fus}}H$ $\Delta_{\text{sub}}H$	1,1-diphenylethanol	26.49 105.0 ± 0.8	357.9 298		[1998VER3] [1998VER3]
C ₁₄ H ₁₄ OS	[26905-24-8] $\Delta_{\text{sub}}H$	4-methoxyphenyl benzyl sulfide	112.5 ± 2.7	298	C	[2006MUL/MOZ]
C ₁₄ H ₁₄ O ₂	[7501-02-2] Δ_vH	2-(2-biphenyloxy)ethanol (410–608)	71.9	425	A	[1987STE/MAL]
C ₁₄ H ₁₄ O ₂ S	[620-32-6] $\Delta_{\text{sub}}H$	dibenzyl sulfone	125.5 ± 2.9			[UR/MAC, 1970COX/PIL]
C ₁₄ H ₁₄ O ₂ S	[599-66-6] $\Delta_{\text{sub}}H$	di-p-tolyl sulfone	109.6 ± 2.9			[UR/MAC, 1970COX/PIL]
C ₁₄ H ₁₄ O ₃	[22204-53-1] $\Delta_{\text{fus}}H$ $\Delta_{\text{fus}}H$ $\Delta_{\text{fus}}H$ $\Delta_{\text{sub}}H$	6-methoxy- α -methyl-2-naphthaleneacetic acid (naproxen) 34.2 31.5 29.41 (341–397)	428.8 428.5 439.2 128.3 ± 0.5		DSC DSC DSC GS	[2006WAS/HOL] [1997NEA/BHA] [1993CON/VIA] [2004PER/KUR]
C ₁₄ H ₁₄ O ₅	[11171-30-3] $\Delta_{\text{us}}H$	8-(hydroxymethyl)-6-methyl-2-oxo-2H-1-benzopyran-3-carboxylic acid, ethyl ester	2.96	415.6		

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{fus}}H$		29.58	429.8	DSC	[1992HUA/ZHO2]
C ₁₄ H ₁₄ O ₆	[111171-31-4] $\Delta_{\text{fus}}H$	8-(hydroxymethyl)-6-methoxy-2-oxo-2H-1-benzopyran-3-carboxylic acid, ethyl ester	36.42	431.9	DSC	[1992HUA/ZHO2]
C ₁₄ H ₁₄ O ₈	[3451-02-3] $\Delta_{\text{fus}}H$	1,2,3,4-tetracarbomethoxybenzene	40.4	404.7	DSC	[1993ACR]
C ₁₄ H ₁₄ O ₈	[3034-97-7] $\Delta_{\text{fus}}H$	1,2,3,5-tetracarbomethoxybenzene	32.6	389.2	DSC	[1993ACR]
C ₁₄ H ₁₄ O ₈	[635-10-9] $\Delta_{\text{fus}}H$	1,2,4,5-tetramethoxycarbonylbenzene	35.7	416.7	DSC	[1993ACR]
	$\Delta_{\text{sub}}H$	(371–391)	140.4 ± 0.8	381	ME	[1995JIM/MEN]
	$\Delta_{\text{sub}}H$		143.3 ± 0.8	298		[1995JIM/MEN]
	$\Delta_{\text{sub}}H$		135.9 ± 1.3	298		[1967TUR2, 1995JIM/MEN]
C ₁₄ H ₁₄ S	[538-74-9] $\Delta_{\text{sub}}H$	dibenzyl sulfide	93.3 ± 5		E	[1962MAC/MAY3, 1970COX/PIL]
C ₁₄ H ₁₄ S ₂	[150-60-7] $\Delta_{\text{fus}}H$	benzyl disulfide	44.7	341.7		[2007WAN/TAN]
C ₁₄ H ₁₅ N	[103-49-1] Δ_vH	dibenzylamine (391–573)	70.5	406	A	[1987STE/MAL, 1947STU]
C ₁₄ H ₁₅ N	[606-99-5] Δ_vH	N,N-diphenyl-N-ethylamine (371–559)	63.2	386	A	[1987STE/MAL, 1947STU]
C ₁₄ H ₁₅ NO	[na] $\Delta_{\text{fus}}H$	2-(4-ethoxyphenyl)-5-methylpyridine	21.0	364		[2000MOR/HAR]
C ₁₄ H ₁₅ N ₃	[60-11-7] $\Delta_{\text{sub}}H$	4-(N,N-dimethylamino)azobenzene (346–354)	117.6 ± 1.7	350	ME	[1967GRE/JON]
	$\Delta_{\text{sub}}H$	(352–354)	115.9 ± 1.3	353	TE	[1967GRE/JON]
	$\Delta_{\text{sub}}H$		120.9 ± 1.7	373	ME	[1956MAJ, 1987STE/MAL]
C ₁₄ H ₁₅ N ₃	[25548-37-2] $\Delta_{\text{sub}}H$	(E) 4-(N,N-dimethylamino)azobenzene	132 ± 8	381	TE	[1985CAM/FER]
C ₁₄ H ₁₅ N ₃	[na] $\Delta_{\text{fus}}H$	N,N-dimethyl-4-phenylazoaniline	23.08	389.2		[1988BAU/PER]
C ₁₄ H ₁₅ N ₃	[na] $\Delta_{\text{sub}}H$	2,3'-dimethyl-4-aminoazobenzene	112.5		GS	[1987SHI/OHK, 1991HOR]
C ₁₄ H ₁₅ N ₃ S	[na] $\Delta_{\text{fus}}H$	N-2-(4,6-lutidyl)-N'-phenylthiourea	50.9	489.7	DSC	[2002VAL/HER]
C ₁₄ H ₁₅ N ₃ S	[71196-80-0] $\Delta_{\text{fus}}H$	N-2-(6-picolyl)-N'-2-tolylthiourea	44.1	468.7	DSC	[2002HER/ACK]
C ₁₄ H ₁₅ N ₃ S	[476443-76-2] $\Delta_{\text{fus}}H$	N-2-(6-picolyl)-N'-3-tolylthiourea	33.2	460.7	DSC	[2002HER/ACK]
C ₁₄ H ₁₅ N ₃ S	[71196-81-1] $\Delta_{\text{fus}}H$	N-2-(6-picolyl)-N'-4-tolylthiourea	47.2	492.2	DSC	[2002HER/ACK]
C ₁₄ H ₁₆	[2717-39-7] $\Delta_{\text{sub}}H$	1,4,5,8-tetramethylnaphthalene	99.8 ± 1.4	298	C	[1974MAN3, 1977PED/RYL]
C ₁₄ H ₁₆	[na]	heptacyclo[6.6.0 ^[2,6] .0 ^[3,13] .0 ^[4,11] .0 ^[5,9] .0 ^[8,1] .0 ^[10,14]] tetradecane				

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{fus}}H$		14.67	355		
	$\Delta_{\text{fus}}H$		5.57	440		[1994KAB/KOZ]
C ₁₄ H ₁₆ ClNO	[124771-55-7]	2-chloromethylbenzoxazole				
	Δ_vH	(348–424)	84.1	363		[2006HUO/ZEN]
C ₁₄ H ₁₆ ClN ₃ O	[67129-08-2]	2-chloro-N-(2,6-dimethylphenyl)-N-(1 <i>H</i> -pyrazol-1-ylmethyl)acetamide (metazachlor)				
	$\Delta_{\text{fus}}H$ (I)		19.7	356.2		
	$\Delta_{\text{fus}}H$ (II)		23	353.2		
	$\Delta_{\text{fus}}H$ (III)		26.6	349.2	DSC	[2004GRI/WEI]
C ₁₄ H ₁₆ ClN ₃ O ₂	[43121-43-3]	{1-(4-chlorophenoxy)-3,3-dimethyl-1-(1 <i>H</i> -1,2,4-triazol-1-yl)}butanone (triadimefon)				
	$\Delta_{\text{sub}}H$	(298–343)	111.1 ± 2.2	303	GS	[1997DAS/DAS]
C ₁₄ H ₁₆ ClN ₃ O ₂	[43121-43-3]	1-(4-chlorophenoxy)-3,3-dimethyl-(1 <i>H</i> ,1,2,4-triazol-1-yl)-2-butanone				
	$\Delta_{\text{fus}}H$		22.87	351.4	DSC	[1990DON/DRE]
C ₁₄ H ₁₆ F ₃ N ₃ O ₄	[26399-36-0]	N-(cyclopropylmethyl)-2,6-dinitro-N-propyl-4-(trifluoromethyl)benzenamine				
	$\Delta_{\text{fus}}H$		22.51	305.8	DSC	[1990DON/DRE]
C ₁₄ H ₁₆ N ₂ O ₂	[na]	1,3- <i>bis</i> (1-isocyanato-1-methylethyl)benzene				
	Δ_vH	(298–426)	65.2	361	HSA, T, DTA	[1986ACH/HAS]
C ₁₄ H ₁₆ N ₂ O ₂	[na]	1,4- <i>bis</i> (1-isocyanato-1-methylethyl)benzene				
	Δ_vH	(373–428)	74.0	400	HSA, T, DTA	[1986ACH/HAS]
C ₁₄ H ₁₆ N ₂ O ₂ S	[108929-67-5]	4-amino-N-(4-ethylphenyl)benzenesulfonamide				
	$\Delta_{\text{fus}}H$		36.3	436.2	DSC	[2009PER/TKA]
	$\Delta_{\text{sub}}H$		143.6 ± 0.9	298	GS	[2009PER/TKA]
	Δ_vH		118.8	298	S-F	[2009PER/TKA]
C ₁₄ H ₁₆ N ₂ O ₇	[152672-90-7]	2-methylpropanoic acid 2,3-dihydro-7-(1-methylethoxy)-3-[2-(nitrooxy)-ethyl]-4-oxo-2 <i>H</i> -1,3-benzoxazin-7-yl ester				
	$\Delta_{\text{fus}}H$		26	345.7	DSC	[1996FON/ROS]
C ₁₄ H ₁₆ O ₅	[20666-86-8]	benzoyl (3-cyclohexyloxy)carbonyl peroxide				
	$\Delta_{\text{sub}}H$	(293–313)	96.2 ± 4.2	303	ME	[1971KIP/RAB, 1977PED/RYL]
C ₁₄ H ₁₇ ClNO ₄ PS ₂	[10311-84-9]	S-[2-chloro-1-(1,3-dihydro-1,3-dioxo-2 <i>H</i> -isoindol-2-yl)ethyl] O,O-diethylphosphorodithioate				
	$\Delta_{\text{fus}}H$		25.27	340	DSC	[1990DON/DRE]
C ₁₄ H ₁₇ Cl ₂ N ₃ O	[79983-71-4]	α -butyl- α -(2,4-dichlorophenyl)-1 <i>H</i> -1,2,4-triazole-1-ethanol (\pm)-hexaconazole				
	$\Delta_{\text{sub}}H$	(318–358)	160.1	338	ME	[1997GOO]
C ₁₄ H ₁₇ Cl ₃ O ₃	[2630-13-9]	hexyl 2,4,5-trichlorophenoxyacetate				
	Δ_vH	(460–573)	85.3	475	A	[1987STE/MAL]
C ₁₄ H ₁₇ NO ₂	[na]	4-methyl-7-diethylaminocoumarin				
	$\Delta_{\text{fus}}H$		17.88	343.8		[1996DOM/HEA]
C ₁₄ H ₁₇ N ₅ O ₃	[51940-44-4]	8-ethyl-5,8-dihydro-5-oxo-2-(1-piperazinyl)pyrido[2,3- <i>d</i>]-pyrimidine-6-carboxylic acid (pipemidic acid)				
	$\Delta_{\text{fus}}H$		32.85	529.9		[2004ROM/BUS2]
C ₁₄ H ₁₈	[1079-71-6]	1,2,3,4,5,6,7,8-octahydroanthracene				
	$\Delta_{\text{fus}}H$		2.51	331.4		
	$\Delta_{\text{fus}}H$		18.34	345.4		[1996DOM/HEA]
	Δ_vH	(437–498)	45.6	452	A	[1987STE/MAL]
	Δ_vH	(348–433)	NA		IP	[1982GAM/CAL]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₄ H ₁₈	[5325-97-3] $\Delta_v H$	1,2,3,4,5,6,7,8-octahydrophenanthrene (402–570)	55.8	417	A	[1987STE/MAL]
C ₁₄ H ₁₈ ClN ₃ O ₂	[55219-65-3] $\Delta_{\text{fus}} H$	β -(4-chlorophenoxy)- α -(1,1-dimethylethyl)-1 <i>H</i> -1,2,4-triazole-1-ethanol	24.47	377.8	DSC	[1990DON/DRE]
C ₁₄ H ₁₈ Cl ₂ O ₃	[1917-95-9] $\Delta_v H$	hexyl 2,4-dichlorophenoxyacetate (444–573)	81.3	459	A,GC	[1987STE/MAL, 1966JEN/SCH]
C ₁₄ H ₁₈ Cl ₂ O ₃	[na] $\Delta_v H$	isohexyl 2,4-dichlorophenoxyacetate (460–573)	69.1	475	A,GC	[1987STE/MAL, 1999DYK/SVO, 1966JEN/SCH]
C ₁₄ H ₁₈ N ₂	[10075-69-1] $\Delta_{\text{sub}} H$	1,5-N,N,N',N'-tetramethyldiaminonaphthalene (318–356)	98.6 ± 0.4	298	GS	[2007VER/GEO]
C ₁₄ H ₁₈ N ₂	[20734-58-1] $\Delta_{\text{sub}} H$	1,8-N,N,N',N'-tetramethyldiaminonaphthalene (324–364)	94.7 ± 0.8	298	GS	[2007VER/GEO]
	$\Delta_v H$	(324–364)	76.7 ± 0.4	298	GS	[2007VER/GEO]
C ₁₄ H ₁₈ N ₂ O ₅	[81-14-1] $\Delta_{\text{fus}} H$	2,6-dimethyl-3,5-dinitro-4- <i>tert</i> -butylacetophenone	23.81	408.5		[2004QU/BAI]
	$\Delta_{\text{sub}} H$	(293–353)	107.9	323	ME	[1953SER/VOI, 1960JON]
C ₁₄ H ₁₈ N ₄ O ₃	[738-70-5] $\Delta_{\text{fus}} H$	5-[(3,4,5-trimethoxyphenyl)methyl]-2,4-pyrimidinediamine (trimethoprim)	49.8	472.9	DSC	[2006WAS/HOL]
	$\Delta_{\text{fus}} H$		53.65	474		[1998ISS/ELA]
C ₁₄ H ₁₈ O	[122-40-7] $\Delta_v H$	α -pentylcinnamaldehyde (282–333)	75.3	297	A, ME	[1987STE/MAL, 1955SER/VOI]
C ₁₄ H ₁₈ O	[30545-23-4] $\Delta_{\text{sub}} H$	diamantanone	103.1 ± .62	320	TSGC	[1980CLA/KNO]
C ₁₄ H ₁₈ O ₂	[180988-52-7] $\Delta_{\text{fus}} H$	6,6-dimethyl-1-phenyl-4,8-dioxaspiro[2.5]octane	27.2	351.2		[1998VER/PEN]
	$\Delta_{\text{sub}} H$		97.5 ± 0.3	298		[1998VER/PEN]
C ₁₄ H ₁₈ O ₃	[49763-96-4] $\Delta_{\text{fus}} H$	E(+) 4,4-dimethyl-1-(3,4-methylenedioxyphenyl)-1-penten-3-ol (stiripentol)	29.0	348.2		[1991CEO/DUG]
C ₁₄ H ₁₈ O ₄	[131-16-8] $\Delta_v H$	dipropyl phthalate (403–578)	73.2	418	A	[1987STE/MAL]
C ₁₄ H ₁₈ O ₄	[na] $\Delta_v H$	diisopropyl phthalate	74.8	430	BG	[1988KAT]
C ₁₄ H ₁₉ Cl ₂ NO ₂	[305-03-3] $\Delta_{\text{fus}} H$	4- <i>p</i> -[bis(2-chloroethyl)amino]benzene]butanoic acid	29.18	338.9	DSC	[1990DON/DRE]
C ₁₄ H ₁₉ NO	[na] $\Delta_{\text{fus}} H$	2-(dimethylamino)-1,2-diphenylethanone	22.38	334.2		[1994WEL/VER]
C ₁₄ H ₁₉ NO	[18494-61-6] $\Delta_v H$	hexahydro-1-(phenylacetyl)-1 <i>H</i> -azepine (370–418)	53.9	385	A	[1987STE/MAL, 1969DAV/MAK]
	$\Delta_v H$	(371–420)	49.4	396		[1969DAV/MAK]
C ₁₄ H ₁₉ NO ₂ S	[na] $\Delta_{\text{sub}} H$	N-benzoylthiocarbamic O-hexyl ester	139.7 ± 2.4	298	C	[2004RIB/SAN2]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound						
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference		
C ₁₄ H ₁₉ N ₂ O ₂	[70585-35-2]	(1RS,2SR)-1-(4-chlorophenoxy)-3,3-dimethyl-1-(1 <i>H</i> -1,2,4-triazol-1-yl)-butan-2-ol (<i>erythro</i> triadimenol)						
	$\Delta_{\text{fus}}H$ (I)		32	411.2				
	$\Delta_{\text{fus}}H$ (II)		33.1	406.2				
	$\Delta_{\text{fus}}H$ (III)		25.1	385.2	DSC	[2000BUR/VAN]		
C ₁₄ H ₁₉ N ₂ O ₂	[70585-37-4]	(1RR,2SS)-1-(4-chlorophenoxy)-3,3-dimethyl-1-(1 <i>H</i> -1,2,4-triazol-1-yl)-butan-2-ol (<i>threo</i> triadimenol)						
	$\Delta_{\text{fus}}H$		33.2	406.2	DSC	[2000BUR/VAN]		
C ₁₄ H ₁₉ N ₅ O ₃	[157891-99-1]	6- <i>tert</i> -butyl-3,9-dihydro-3-[(2-hydroxyethoxy)methyl]-9-oxo-5 <i>H</i> -imidazo[1,2- <i>a</i>]pyrine						
	$\Delta_{\text{fus}}H$		37.95	478.5	DSC	[1999ZIE/GOL]		
C ₁₄ H ₂₀	[4413-16-5]	1-cyclohexyl-1-phenylethane						
	Δ_vH	(359–400)	70.8	374	A, MG	[1987STE/MAL, 1955SCH/WHI]		
C ₁₄ H ₂₀	[1603-61-8]	1-cyclohexyl-2-phenylethane						
	Δ_vH	(372–406)	60.7	387	A, MG	[1987STE/MAL, 1955SCH/WHI]		
C ₁₄ H ₂₀	[2883-12-7]	1-cyclopentyl-3-phenylpropane						
	Δ_vH	(373–540)	61.3	388	A, MG	[1987STE/MAL, 1955SCH/WHI]		
C ₁₄ H ₂₀	[1540-80-3]	1,8-cyclotetradecadiyne						
	$\Delta_{\text{fus}}H$		22.6	370		[1974AUG/BOR]		
	$\Delta_{\text{sub}}H$	(315–364)	87.6 ± 1.0	338	HSA	[1998CHI/HES]		
	$\Delta_{\text{sub}}H$		94.3	298	CGC-DSC	[1998CHI/HES]		
C ₁₄ H ₂₀	[1540-80-3]	$\Delta_{\text{sub}}H$	(317–332)	166.0 ± 3.2	325	ME	[1964FRI/BAU, 1970COX/PIL]	
		C ₁₄ H ₂₀	[1079-71-6]	1,2,3,4,5,6,7,8-octahydroanthracene (octhracene)				
			$\Delta_{\text{sub}}H$	(438–499)	82.3 ± 1.2	298	BG	[1971BOY/SAN, 1977PED/RYL]
C ₁₄ H ₂₀	[2292-79-7]	diadamantane						
	$\Delta_{\text{fus}}H$		4.44	407.2				
	$\Delta_{\text{fus}}H$		8.95	440.4				
	$\Delta_{\text{fus}}H$		8.66	517.9		[1996DOM/HEA]		
	$\Delta_{\text{sub}}H$	(305–333)	96.0 ± 0.8	319	TSGC	[1975CLA/KNO]		
C ₁₄ H ₂₀ ClNO ₂	[15972-60-8]	2-chloro- <i>N</i> -(2,6-diethylphenyl)- <i>N</i> -(methoxymethyl)acetamide						
	$\Delta_{\text{fus}}H$		26.7	317.7	DSC	[2005SBI/VEC]		
	$\Delta_{\text{fus}}H$		25.31	315.9	DSC	[1990DON/DRE]		
	Δ_vH		85 ± 1	436	TGA	[2007VEC]		
C ₁₄ H ₂₀ Cl ₂	[na]	1,2-dichloro-3,4,5,6-tetraethylbenzene						
	Δ_vH	(378–575)	66.2	393	A	[1987STE/MAL, 1947STU, 1970DYK/VAN]		
C ₁₄ H ₂₀ Cl ₂	[na]	1,4-dichloro-2,3,5,6-tetraethylbenzene						
	Δ_vH	(364–570)	60.8	379	A	[1987STE/MAL, 1947STU, 1970DYK/VAN]		
C ₁₄ H ₂₀ N ₂ O	[27262-40-4]	<i>N</i> -(2,6-dimethylphenyl)-2-piperidinecarboxamide						
	$\Delta_{\text{fus}}H$		24.19	403.2	DSC	[1997NEM/ACS]		
C ₁₄ H ₂₀ N ₂ OS	[90473-84-0]	<i>N</i> -(diethylaminothiocarbonyl)benzimidazole ethyl ester						
	$\Delta_{\text{sub}}H$		135.6 ± 2.6	298	C	[2006RIB/SAN3]		
C ₁₄ H ₂₀ N ₂ O ₂	[na]	(–) 1-(1 <i>H</i> -indol-4-yloxy)-3-(isopropylamino)-2-propanol (pindolol)						
	$\Delta_{\text{fus}}H$		25.69	365.7		[1999LI/ZEL]		
C ₁₄ H ₂₀ N ₂ O ₂	[13523-86-9]	(±) 1-(1 <i>H</i> -indol-4-yloxy)-3-(isopropylamino)-2-propanol (pindolol)						
	$\Delta_{\text{fus}}H$		60.6	423.6	DSC	[2007PER/VOL]		
	$\Delta_{\text{fus}}H$		58	443.8		[2004NUN/EUS]		

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{fus}}H$		57.9	442.9		[1999LI/ZEL]
	$\Delta_{\text{sub}}H$	(355–427)	146.0 ± 1.2	298	GS	[2007PER/VOL]
C ₁₄ H ₂₀ N ₃ O ₅ PS	[13457-18-6] $\Delta_{\text{fus}}H$	O-6-ethoxycarbonyl-5-methylpyrazolo[1,5-a]pyrimidin-2-yl O,O-diethyl phosphorothioate	27.32	324.4	DSC	[1990DON/DRE]
C ₁₄ H ₂₀ O	[61812-55-3] Δ_vH	(1-cyclohexyloxyethyl)benzene	69.8 ± 0.5	298	GS	[2002KRA/VAS, 2002VER/HEI]
C ₁₄ H ₂₀ O	[30545-14-3] $\Delta_{\text{fus}}H$	diamantan-1-ol	18.0	395		
	$\Delta_{\text{fus}}H$		4.9	408		
	$\Delta_{\text{fus}}H$		9.6	573	DSC	[1974CLA/MCK]
	$\Delta_{\text{sub}}H$	(319–349)	118. ± 0.6	334		[1980CLA/KNO, 1975CLA/KNO]
C ₁₄ H ₂₀ O	[30545-24-5] $\Delta_{\text{sub}}H$	diamantan-3-ol	116.1 ± 4.4	338		[1980CLA/KNO, 1975CLA/KNO]
C ₁₄ H ₂₀ O	[30651-03-7] $\Delta_{\text{fus}}H$	diamantan-4-ol	9.77	448		
	$\Delta_{\text{fus}}H$		16.4	484	DSC	[1974CLA/MCK]
	$\Delta_{\text{sub}}H$	(322–353)	117.8 ± 0.2	337		[1980CLA/KNO, 1975CLA/KNO]
C ₁₄ H ₂₀ O ₂	[3383-21-9] $\Delta_{\text{fus}}H$	3,5-di- <i>tert</i> -butyl- <i>o</i> -benzoquinone	26.53	387.9		[2005FAT/KAS]
C ₁₄ H ₂₀ O ₂	[38350-87-7] $\Delta_{\text{sub}}H$	4-heptylbenzoic acid	130.0 ± 0.9	298	ME	[2004MON/ALM]
C ₁₄ H ₂₀ O ₃	[na] Δ_vH	2-(4- <i>tert</i> -butylphenoxy)ethyl acetate	78.8	406	A	[1987STE/MAL, 1947STU]
C ₁₄ H ₂₀ O ₄	[58608-07-4] $\Delta_{\text{fus}}H$	2,5-dibutoxy-1,4-benzoquinone	4.7	328.3		
	$\Delta_{\text{fus}}H$		2.3	364.5		
	$\Delta_{\text{fus}}H$		31.5	473.3	DSC	[1996KEE/VAN]
C ₁₄ H ₂₀ O ₅	[14098-44-3] $\Delta_{\text{sub}}H$	benzo-15-crown-5	128.1 ± 10.8	298	CGC-DSC	[2000NIC/ORF]
	Δ_vH		98.9 ± 1.3	298	CGC	[2000NIC/ORF]
C ₁₄ H ₂₁ F ₃ N ₂ O ₄	[na] $\Delta_{\text{sub}}H$	proline, 1-[N-(trifluoroacetyl)-(l)-leucyl]methyl ester	121.3	328	A	[1987STE/MAL, 1960WEY/KLI]
	Δ_vH	(366–453)	105.8	381	A	[1987STE/MAL]
C ₁₄ H ₂₁ NO	[121678-88-4] $\Delta_{\text{sub}}H$	4-isopropylbenzylidene <i>tert</i> -butylamine N-oxide	101.8 ± 4.1	298	C	[1989ACR/KIR]
C ₁₄ H ₂₁ N ₃ O ₄	[33629-47-9] $\Delta_{\text{fus}}H$	4-(1,1-dimethylethyl)-N-(1-methylpropyl)-2,6-dinitrobenzeneamine	20.84	338.8	DSC	[1990DON/DRE]
C ₁₄ H ₂₁ N ₃ S	[90473-92-0] $\Delta_{\text{sub}}H$	N-(diethylaminothiocarbonyl)-N-monoethylbenzamide	141.2 ± 1.2	298	C	[2006RIB/SAN3]
C ₁₄ H ₂₂	[1012-72-2] $\Delta_{\text{fus}}H$	1,4-di- <i>tert</i> -butylbenzene	14.4	350.7	AC, DSC	[2009CHI/STE]
	$\Delta_{\text{fus}}H$		8.2	350.8	AC, DSC	[2009CHI/STE]
	$\Delta_{\text{fus}}H$		22.48	341.5		[1997STE/CHI3]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{sub}}H$	(288–333)	82.1 ± 0.4	310	T	[1998VER]
	$\Delta_{\text{sub}}H$		82.8 ± 0.4	298		[1998VER]
	$\Delta_{\text{sub}}H$	(285–325)	82.8	305	ME	[1951HOP/SEA, 1987STE/MAL]
	Δ_vH	(319–559)	63.0 ± 0.1	298	EB,IP	[2009CHI/STE]
	Δ_vH	(319–559)	55.8 ± 0.1	360	EB,IP	[2009CHI/STE]
	Δ_vH	(319–559)	54.6 ± 0.1	400	EB,IP	[2009CHI/STE]
	Δ_vH	(319–559)	61.4 ± 0.1	440	EB,IP	[2009CHI/STE]
	Δ_vH	(319–559)	46.4 ± 0.2	480	EB,IP	[2009CHI/STE]
	Δ_vH	(319–559)	44.6 ± 0.3	520	EB,IP[[2009CHI/STE]
	Δ_vH	(354–382)	61.4 ± 0.3	298	GS	[2008VER/KOZZ]
	Δ_vH	(387–559)	63.0 ± 0.6	298	EB	[1997STE/CHI3]
C ₁₄ H ₂₂	[1014-60-4]	1,3-di- <i>tert</i> -butylbenzene				
	Δ_vH	(288–333)	58.9 ± 0.5	310	GS	[1998VER]
	Δ_vH		59.6 ± 0.5	298		[1998VER]
	Δ_vH	(346–374)	58.0	360	A	[1987STE/MAL]
C ₁₄ H ₂₂	[2189-60-8]	octylbenzene				
	$\Delta_{\text{fus}}H$		29.96	234.2		[1996DOM/HEA]
	Δ_vH	(293–462)	67.4	308		[1993KAS/MOK]
	Δ_vH	(368–400)	63.1	383	A	[1987STE/MAL]
	Δ_vH	(316–399)	66.2	336	GS	[1986ALL/JOS]
C ₁₄ H ₂₂	[777-22-0]	2-phenyloctane				
	Δ_vH	(361–392)	61.6	376	A	[1987STE/MAL]
	Δ_vH		70.0	298		[1971WIL/ZWO]
C ₁₄ H ₂₂	[642-32-0]	1,2,3,4-tetraethylbenzene				
	Δ_vH	(423–525)	62.6	438	A	[1987STE/MAL]
C ₁₄ H ₂₂	[38842-05-6]	1,2,3,5-tetraethylbenzene				
	Δ_vH	(413–521)	64.8	428	A	[1987STE/MAL]
C ₁₄ H ₂₂	[635-81-4]	1,2,4,5-tetraethylbenzene				
	Δ_vH	(338–521)	54.5	353	A	[1987STE/MAL]
C ₁₄ H ₂₂ N ₂ O	[137-58-6]	2-(diethylamino)-N-(2,6-dimethylphenyl)acetamide (lidocaine)				
	$\Delta_{\text{fus}}H$		16.4	340.7	DSC	[2010LAZ/RIE]
	$\Delta_{\text{fus}}H$		18.8	341	DSC	[2008WAS/HOL]
C ₁₄ H ₂₂ N ₂ O ₂ S	[na]	N,N-diisobutyl-N'-furoylthiourea				
	$\Delta_{\text{sub}}H$		141.7 ± 5.6	298	C	[2002RIB/RIB]
C ₁₄ H ₂₂ N ₂ O ₃	[56715-13-0]	(+) 4-[2'-hydroxy-3'-(isopropylamino)propoxy]-phenylacetamide (atenolol)				
	$\Delta_{\text{fus}}H$		36.77	420.3	DSC	[1999LI/ZEL]
C ₁₄ H ₂₂ N ₂ O ₃	[29122-68-7]	(+) 4-[2'-hydroxy-3'-(isopropylamino)propoxy]-phenylacetamide (atenolol)				
	$\Delta_{\text{fus}}H$		38.7	426.1	DSC	[2007PER/VOL]
	$\Delta_{\text{fus}}H$		35.66	423.4	DSC	[1999LI/ZEL]
	$\Delta_{\text{sub}}H$	(396–418)	140.0 ± 3.7	298	GS	[2007PER/VOL]
C ₁₄ H ₂₂ N ₄ O ₂	[35873-43-9]	8-heptyltheophylline				
	$\Delta_{\text{fus}}H$		33	472.7	DSC	[1991ACR, 1989GON/KRA]
C ₁₄ H ₂₂ N ₄ O ₆	[74734-25-1]	N,N'-bis(2-oxo-3-oxazolidin-3-ylcarbonyl)-1,6-hexandiamine				
	$\Delta_{\text{fus}}H$		8.9	400.8		[1990SHI/HAY]
C ₁₄ H ₂₂ N ₄ O ₆ S	[19044-94-1]	4-(dipropylamino)-N,N-dimethyl-3,5-dinitrobenzenesulfonamide				

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₄ H ₂₂ O	$\Delta_{\text{fus}}H$		32.57	413.6	DSC	[1990DON/DRE]
	[96-76-4]	2,4-di- <i>tert</i> -butylphenol				
	$\Delta_{\text{sub}}H$	(288–327)	86.1 ± 0.3	308	GS	[1999VER2]
	$\Delta_{\text{sub}}H$		86.7 ± 0.3	298		[1999VER2]
	$\Delta_{\text{sub}}H$		92.9 ± 2.8	298	C	[1999RIB/MAT2]
	Δ_vH	(333–368)	69.2 ± 0.5	350	GS	[1999VER2]
	Δ_vH		72.4 ± 0.5	298		[1999VER2]
C ₁₄ H ₂₂ O	[128-39-2]	2,6-di- <i>tert</i> -butylphenol				
	$\Delta_{\text{fus}}H$		16.57	310.7		[1972INO/LIA]
	$\Delta_{\text{sub}}H$		84.6 ± 0.5	298	GS	[1999VER]
	$\Delta_{\text{sub}}H$		81.5 ± 2.3	298	C	[1999RIB/MAT2]
	$\Delta_{\text{sub}}H$		U 110.9	298	C	[1971BER/GIR, 1999VER]
	Δ_vH	(313–368)	63.5 ± 0.2	341	GS	[1999VER]
	Δ_vH		66.0 ± 0.2	298		[1999VER]
C ₁₄ H ₂₂ O	Δ_vH	(386–530)	60.4	401	A	[1987STE/MAL]
	[1138-52-9]	3,5-di- <i>tert</i> -butylphenol				
	$\Delta_{\text{sub}}H$		97.7 ± 3.7	298	C	[2001RIB/MAT]
C ₁₄ H ₂₂ O	$\Delta_{\text{sub}}H$	(302–325)	68.2	313.5	A	[1987STE/MAL]
	[63264-81-3]	4-(1,1-diethylbutyl)phenol				
C ₁₄ H ₂₂ O	Δ_vH	(404–549)	69.5	419	A	[1987STE/MAL]
	[65152-07-0]	2,4-diisobutylphenol				
C ₁₄ H ₂₂ O	Δ_vH	(448–598)	65.0	463	A	[1987STE/MAL]
	[59048-99-6]	4-[(1,2-dimethyl-1-ethyl)butyl]phenol				
C ₁₄ H ₂₂ O	Δ_vH	(415–578)	64.7	430	A	[1987STE/MAL]
	[na]	4-[(1,3-dimethyl-1-ethyl)butyl]phenol				
C ₁₄ H ₂₂ O	Δ_vH	(409–571)	60.9	424	A	[1987STE/MAL]
	[na]	4-[(2,2-dimethyl-1-ethyl)butyl]phenol				
C ₁₄ H ₂₂ O	Δ_vH	(413–553)	67.0	428	A	[1987STE/MAL]
	[79-70-9]	β -irone				
C ₁₄ H ₂₂ O	Δ_vH	(288–333)	72.1	303	A	[1987STE/MAL]
	[127-51-5]	α -isomethylionone				
C ₁₄ H ₂₂ O	Δ_vH	(288–333)	69.5	303	A	[1987STE/MAL]
	[1988-35-8]	4-(1-methyl-1-ethyl)pentyl]phenol				
C ₁₄ H ₂₂ O	Δ_vH	(413–578)	62.8	428	A	[1987STE/MAL]
	[127-42-4]	α -methylionone				
C ₁₄ H ₂₂ O	Δ_vH	(288–333)	70.1	303	A	[1987STE/MAL]
	[127-43-5]	β -methylionone				
C ₁₄ H ₂₂ O	Δ_vH	(288–333)	70.3	303	A	[1987STE/MAL]
	[140-66-9]	4-(1,1,3,3-tetramethylbutyl)phenol				
	Δ_vH	(309–350)	68.8 ± 0.3	329	GS	[1999VER2]
	Δ_vH		70.7 ± 0.3	298	GS	[1999VER2]
	Δ_vH	(381–563)	72.4	396	A	[1987STE/MAL, 1959MCD/SHR, 1984BOU/FRI]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₄ H ₂₂ O	[124765-79-3]	4- <i>tert</i> -octylphenol				
		(297–351)	96.3 ± 0.9	324	GS	[1999VER2]
			97.9 ± 0.9	298	GS	[1999VER2]
C ₁₄ H ₂₂ O ₂	[1020-31-1]	3,5-di- <i>tert</i> -butyl-1,2-dihydroxybenzene				
			24.1	372.8		[2000VER/SCH]
			103.7 ± 0.5	330	GS	[2000VER/SCH]
			104.7 ± 0.5	298	GS	[2000VER/SCH]
		100.1 ± 0.6	298	C	[1984CAR]	
C ₁₄ H ₂₂ O ₂	[88-58-4]	2,5-di- <i>tert</i> -butyl-1,4-dihydroxybenzene				
			43.85	496.5		[1999VER7]
		(333–368)	108.8 ± 1.7	351	GS	[1999VER7]
		122.4 ± 1.7	298		[1999VER7]	
C ₁₄ H ₂₂ O ₄	[620-82-6]	dicyclohexyl oxalate				
		(333–360)	92.1 ± 0.7	298	GS	[2008LIP/KRA]
C ₁₄ H ₂₂ O ₆	[1561-49-5]	dicyclohexyl peroxydicarbonate				
			100.4 ± 4.2			[1971KIP/RAB, 1977STE/WAT]
		(293–313)	100.4 ± 8.3	303	ME	[1962RAB/TEL, 1970COX/PIL]
C ₁₄ H ₂₂ O ₁₁	[na]	diethyleneglycol, O,O-dicarboxylic acid, di[1-(methoxycarbonyl)-ethyl] ester				
		(403–493)	98.2	418	A	[1987STE/MAL]
C ₁₄ H ₂₃ N	[na]	N,N-dimethyl-2,3-dimethyl-3-phenyl-2-butanamine				
		(280–335)	65.8 ± 1.3	308	GS	[1998VER/BEC]
		(280–335)	66.4 ± 1.3	298	GS	[1998VER/BEC]
C ₁₄ H ₂₃ NO ₂	[na]	decyl- α -cyanoacrylate				
			41.8	294.5		[1993BYK/KIP]
C ₁₄ H ₂₄	[5743-97-5]	perhydrophenanthrene				
		(455–551)	55.7	470	EB	[2000ROH/CEN]
C ₁₄ H ₂₄	[28071-99-0]	<i>trans-anti-trans</i> perhydroanthracene				
		(269–313)	66.1	284		[1987STE/MAL]
		(275–313)	72.7 ± 3.3	294	ME	[1963MAR/FRI, 1970COX/PIL]
C ₁₄ H ₂₄	[1755-19-7]	<i>trans-syn-trans</i> perhydroanthracene				
		(293–335)	88.1	308	A	[1987STE/MAL]
		(335–393)	87.4 ± 2.4	365	ME	[1963MAR/FRI, 1970COX/PIL]
C ₁₄ H ₂₄	[1687-36-1]	1,3,5,7-tetramethyladamantane				
		(310–350)	83.7 ± 1.3	298	BG	[1977STE/WAT]
		(295–315)	81.1 ± 10.9	305	TSGC	[1975CLA/KNO]
C ₁₄ H ₂₄	[na]	<i>cis-anti-trans</i> -perhydrophenanthrene				
			11.16	313		[1996DOM/HEA]
C ₁₄ H ₂₄	[na]	<i>cis-syn-trans</i> -perhydrophenanthrene				
			10.48	273		[1996DOM/HEA]
C ₁₄ H ₂₄	[2108-89-6]	<i>trans-anti-trans</i> -perhydrophenanthrene				
			11.83	283		[1996DOM/HEA]
C ₁₄ H ₂₄	[1687-36-1]	1,3,5,7-tetramethyladamantane				
			0.23	183.3		
			9.82	337.2		[1977CLA/KNO]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₄ H ₂₄ NO ₄ PS ₃	[741-58-2] $\Delta_{\text{us}}H$	O,O-diisopropyl S-2-phenylsulfonylaminoethylphosphorodithioate				
			30.61	310.4	DSC	[1990DON/DRE]
C ₁₄ H ₂₄ N ₂	[101-96-2] Δ_vH	N,N'-di-sec-butyl-1,4-phenylenediamine				
		(370–507)	70.3	385	A	[1987STE/MAL]
C ₁₄ H ₂₄ N ₂	[7735-44-6] $\Delta_{\text{us}}H$ $\Delta_{\text{fus}}H$	tetradecanedinitrile				
			1.77	261.1	DSC	[2007BAD/BLA]
			40.17	309.6		
C ₁₄ H ₂₄ O	[53131-20-7] Δ_vH	2,2,5,9-tetramethyl-4,8-decanedienal				
		(353–416)	66.4	368	A	[1987STE/MAL]
C ₁₄ H ₂₄ O	[na] Δ_vH	borneol butyrate				
		(347–520)	59.6	362	A	[1987STE/MAL, 1947STU]
C ₁₄ H ₂₄ O	[24717-86-0] Δ_vH	<i>dl</i> borneol isobutyrate				
		(343–516)	58.8	358	A	[1987STE/MAL, 1947STU]
C ₁₄ H ₂₄ O	[na] Δ_vH	geraniol butyrate				
		(369–531)	68.6	384	A	[1987STE/MAL, 1947STU]
C ₁₄ H ₂₄ O	[na] Δ_vH	geraniol isobutyrate				
		(363–524)	67.8	378	A	[1987STE/MAL, 1947STU]
C ₁₄ H ₂₄ O ₂	[38300-49-1] $\Delta_{\text{fus}}H$	1,8-cyclotetradecanedione				
			27.53	417.2		[1972ALV/BOR]
C ₁₄ H ₂₄ O ₄	[na] $\Delta_{\text{fus}}H$	1,6-cyclodecanedione <i>bis</i> ethylene ketal				
			32.68	450.2		[1972ALV/BOR]
C ₁₄ H ₂₄ O ₆	[na] Δ_vH	1,1,1- <i>tris</i> (ethoxycarbonyl)pentane				
		(298–343)	81.4 ± 0.4		GS	[1995RAK/VER]
C ₁₄ H ₂₆	[2883-07-0] Δ_vH	1-cyclohexyl-3-cyclopentylpropane				
		(371–403)	64.5	386	A	[1987STE/MAL]
C ₁₄ H ₂₆	[2319-61-1] Δ_vH	1,1-dicyclohexylethane				
		(370–402)	62.1	385	A	[1987STE/MAL]
C ₁₄ H ₂₆	[3321-50-4] Δ_vH	1,2-dicyclohexylethane				
		(371–402)	65.4	386	A	[1987STE/MAL]
C ₁₄ H ₂₆ O	[53965-17-6] Δ_vH	<i>cis</i> 2,2,5,9-tetramethyl-4,8-decadiene-1-ol				
		(363–393)	94.0	378	A	[1987STE/MAL]
C ₁₄ H ₂₆ O	[53965-18-7] Δ_vH	<i>trans</i> 2,2,5,9-tetramethyl-4,8-decadiene-1-ol				
		(363–393)	86.3	378	A	[1987STE/MAL]
C ₁₄ H ₂₆ O	[3021-89-4] Δ_vH	2-pentyl-2-nonenal				
		(384–553)	65.7	399		[1987MIL/FEN2]
C ₁₄ H ₂₆ O	[142628-55-5] Δ_vH	(Z) 2-tetradecenal				
		(353–393)	82.5	298	CGC	[1996KOU/HOS, 2000OVA/KOU]
C ₁₄ H ₂₆ O	[51534-36-2] Δ_vH	(E) 2-tetradecenal				
		(353–393)	82.6	298	CGC	[1996KOU/HOS, 2000OVA/KOU]
C ₁₄ H ₂₆ O	[174155-51-2] Δ_vH	(Z) 3-tetradecenal				
		(353–393)	79.4	298	CGC	[1996KOU/HOS, 2000OVA/KOU]
C ₁₄ H ₂₆ O	[174155-50-1] Δ_vH	(E) 3-tetradecenal				
		(353–393)	80.1	298	CGC	[1996KOU/HOS, 2000OVA/KOU]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₄ H ₂₆ O	[115018-49-0] $\Delta_v H$	(Z) 4-tetradecenal (353–393)	79.2	298	CGC	[1996KOU/HOS, 2000OVA/KOU]
C ₁₄ H ₂₆ O	[115018-39-8] $\Delta_v H$	(E) 4-tetradecenal (353–393)	79.9	298	CGC	[1996KOU/HOS, 2000OVA/KOU]
C ₁₄ H ₂₆ O	[63851-42-3] $\Delta_v H$	(Z) 5-tetradecenal (353–393)	78.4	298	CGC	[1996KOU/HOS, 2000OVA/KOU]
C ₁₄ H ₂₆ O	[174155-52-3] $\Delta_v H$	(E) 5-tetradecenal (353–393)	79.1	298	CGC	[1996KOU/HOS, 2000OVA/KOU]
C ₁₄ H ₂₆ O	[174155-53-4] $\Delta_v H$	(Z) 6-tetradecenal (353–393)	78.5	298	CGC	[1996KOU/HOS, 2000OVA/KOU]
C ₁₄ H ₂₆ O	[174155-54-5] $\Delta_v H$	(E) 6-tetradecenal (353–393)	79.3	298	CGC	[1996KOU/HOS, 2000OVA/KOU]
C ₁₄ H ₂₆ O	[65128-96-3] $\Delta_v H$	(Z) 7-tetradecenal (353–393)	78.7	298	CGC	[1996KOU/HOS, 2000OVA/KOU]
C ₁₄ H ₂₆ O	[37011-96-4] $\Delta_v H$	(E) 7-tetradecenal (353–393)	79.2	298	CGC	[1996KOU/HOS, 2000OVA/KOU]
C ₁₄ H ₂₆ O	[169054-69-7] $\Delta_v H$	(Z) 8-tetradecenal (353–393)	78.8	298	CGC	[1996KOU/HOS, 2000OVA/KOU]
C ₁₄ H ₂₆ O	[174155-55-6] $\Delta_v H$	(E) 8-tetradecenal (353–393)	79.3	298	CGC	[1996KOU/HOS, 2000OVA/KOU]
C ₁₄ H ₂₆ O	[53939-27-8] $\Delta_v H$	(Z) 9-tetradecenal (353–393)	79.1	298	CGC	[1996KOU/HOS, 2000OVA/KOU]
C ₁₄ H ₂₆ O	[71377-13-4] $\Delta_v H$	(E) 9-tetradecenal (353–393)	79.5	298	CGC	[1996KOU/HOS, 2000OVA/KOU]
C ₁₄ H ₂₆ O	[144525-16-6] $\Delta_v H$	(Z) 10-tetradecenal (353–393)	79.6	298	CGC	[1996KOU/HOS, 2000OVA/KOU]
C ₁₄ H ₂₆ O	[148238-39-5] $\Delta_v H$	(E) 10-tetradecenal (353–393)	79.8	298	CGC	[1996KOU/HOS, 2000OVA/KOU]
C ₁₄ H ₂₆ O	[35237-64-0] $\Delta_v H$	(Z) 11-tetradecenal (353–393)	80.3	298	CGC	[1996KOU/HOS, 2000OVA/KOU]
C ₁₄ H ₂₆ O	[35746-21-5] $\Delta_v H$	(E) 11-tetradecenal (353–393)	80.5	298	CGC	[1996KOU/HOS, 2000OVA/KOU]
C ₁₄ H ₂₆ O	[174155-56-7] $\Delta_v H$	(Z) 12-tetradecenal (353–393)	80.8	298	CGC	[1996KOU/HOS, 2000OVA/KOU]
C ₁₄ H ₂₆ O	[124499-92-9] $\Delta_v H$	(E) 12-tetradecenal (353–393)	80.8	298	CGC	[1996KOU/HOS, 2000OVA/KOU]
C ₁₄ H ₂₆ O	[na] $\Delta_v H$	2-(1,2-dimethylpropyl)-5,6-dimethylheptenal (385–535)	60.0	400	EB	[1987MIL/FEN2]
C ₁₄ H ₂₆ O	[3021-89-4] $\Delta_v H$	2-pentyl-2-nonenal (385–553)	65.0	409	EB	[1987MIL/FEN2]
C ₁₄ H ₂₆ O	[295-17-0] $\Delta_{\text{sub}} H$	cyclotetradecanone	80.75			[1938WOL/WEG, 1960JON]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound		T _m (K)	Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)			
C ₁₄ H ₂₆ O	[37608-02-9] $\Delta_{\text{fus}}H$	4,4,8,8-tetramethylcyclodecanone		378.2		[1976BOR/DAL]
C ₁₄ H ₂₆ O ₂	[3179-47-3] $\Delta_{\text{fus}}H$	decyl methacrylate		250.7		[1996DOM/HEA]
	Δ_vH	(350–541)	62.7	365	A	[1987STE/MAL]
C ₁₄ H ₂₆ O ₂	[84801-15-0] Δ_vH	(Z) 2-dodecenyl acetate (333–378)		298	GC	[1997KOU/HOS, 2000OVA/KOU]
C ₁₄ H ₂₆ O ₂	[84801-16-1] Δ_vH	(E) 2-dodecenyl acetate (333–378)		298	GC	[1997KOU/HOS, 2000OVA/KOU]
C ₁₄ H ₂₆ O ₂	[38363-24-5] Δ_vH	(Z) 3-dodecenyl acetate (333–378)		298	GC	[1997KOU/HOS, 2000OVA/KOU]
C ₁₄ H ₂₆ O ₂	[56218-63-4] Δ_vH	(E) 3-dodecenyl acetate (333–378)		298	GC	[1997KOU/HOS, 2000OVA/KOU]
C ₁₄ H ₂₆ O ₂	[38363-25-6] Δ_vH	(Z) 4-dodecenyl acetate (333–378)		298	GC	[1997KOU/HOS, 2000OVA/KOU]
C ₁₄ H ₂₆ O ₂	[38363-26-7] Δ_vH	(E) 4-dodecenyl acetate (333–378)		298	GC	[1997KOU/HOS, 2000OVA/KOU]
C ₁₄ H ₂₆ O ₂	[16676-96-3] Δ_vH	(Z) 5-dodecenyl acetate (333–378)		298	GC	[1997KOU/HOS, 2000OVA/KOU]
C ₁₄ H ₂₆ O ₂	[16676-97-4] Δ_vH	(E) 5-dodecenyl acetate (333–378)		298	GC	[1997KOU/HOS, 2000OVA/KOU]
C ₁₄ H ₂₆ O ₂	[16974-12-2] Δ_vH	(Z) 6-dodecenyl acetate (333–378)		298	GC	[1997KOU/HOS, 2000OVA/KOU]
C ₁₄ H ₂₆ O ₂	[29868-16-4] Δ_vH	(E) 6-dodecenyl acetate (333–378)		298	GC	[1997KOU/HOS, 2000OVA/KOU]
C ₁₄ H ₂₆ O ₂	[14959-86-5] Δ_vH	(Z) 7-dodecenyl acetate (333–378)		298	GC	[1997KOU/HOS, 2000OVA/KOU]
	Δ_vH	(303–317)	77.5	310	GC	[1983OLS/JON]
C ₁₄ H ₂₆ O ₂	[16695-41-3] Δ_vH	(E) 7-dodecenyl acetate (333–378)		298	GC	[1997KOU/HOS, 2000OVA/KOU]
C ₁₄ H ₂₆ O ₂	[28079-04-1] Δ_vH	(Z) 8-dodecenyl acetate (333–378)		298	GC	[1997KOU/HOS, 2000OVA/KOU]
C ₁₄ H ₂₆ O ₂	[38363-29-0] Δ_vH	(E) 8-dodecenyl acetate (333–378)		298	GC	[1997KOU/HOS, 2000OVA/KOU]
C ₁₄ H ₂₆ O ₂	[16974-11-1] Δ_vH	(Z) 9-dodecenyl acetate (333–378)		298	GC	[1997KOU/HOS, 2000OVA/KOU]
C ₁₄ H ₂₆ O ₂	[35148-19-7] Δ_vH	(E) 9-dodecenyl acetate (333–378)		298	GC	[1997KOU/HOS, 2000OVA/KOU]
C ₁₄ H ₂₆ O ₂	[35148-20-0] Δ_vH	(Z) 10-dodecenyl acetate (333–378)		298	GC	[1997KOU/HOS, 2000OVA/KOU]
C ₁₄ H ₂₆ O ₂	[35153-09-4] Δ_vH	(E) 10-dodecenyl acetate (333–378)		298	GC	[1997KOU/HOS, 2000OVA/KOU]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₄ H ₂₆ O ₃	[5963-13-3] $\Delta_v H$	1,7-dioxa-8-cyclohexadecanone (403–453)	73.3	418	A	[1987STE/MAL]
C ₁₄ H ₂₆ O ₃	[23144-23-2] $\Delta_v H$	3-heptyl-4-acetoxytetrahydro-2H-pyran (383–453)	74.4	398	A	[1987STE/MAL]
C ₁₄ H ₂₆ O ₃	[872803-07-1] $\Delta_v H$ $\Delta_v H$	nonyl levulinate (423–571)	69.4 68.4	438 516	A	[1987STE/MAL] [1933COW/SCH]
C ₁₄ H ₂₆ O ₄	[821-38-5] $\Delta_{\text{fus}}H + \Delta_{\text{trs}}H$ $\Delta_{\text{fus}}H$ $\Delta_v H$	1,14-tetradecanedioic acid (424–503)	56.9 56.5 127.4 ± 2.3	398 397.3 298	DSC DSC CGC	[2006VEN/MET] [2005ROU/TEM] [2005ROU/TEM]
C ₁₄ H ₂₆ O ₄	[105-99-7] $\Delta_v H$	dibutyl adipate (435–563)	68.7	450	A	[1987STE/MAL]
C ₁₄ H ₂₆ O ₄	[na] $\Delta_v H$	diethyl isopentylmalonate (388–526)	75.3	403	A	[1987STE/MAL]
C ₁₄ H ₂₆ O ₄	[na] $\Delta_v H$	2-methylheptane-5,5-dicarboxylic acid, diethyl ester (394–427)	70.1	409	A	[1987STE/MAL]
C ₁₄ H ₂₆ O ₄	[110-40-7] $\Delta_v H$	diethyl decanedioate (398–579)	74.1	413	A	[1987STE/MAL, 1947STU]
C ₁₄ H ₂₆ O ₅	[na] $\Delta_v H$	ethyl[1-(1-octyloxycarbonyl)ethyl]carbonate (413–513)	74.0	428	A	[1987STE/MAL]
C ₁₄ H ₂₆ O ₅	[902261-33-0] $\Delta_v H$	hexyl[1-(1-butoxycarbonyl)ethyl]carbonate (357–501)	72.1	372	A	[1987STE/MAL]
C ₁₄ H ₂₆ O ₆ S	[5423-27-8] $\Delta_{\text{fus}}H$	dibutyl 3,3'-sulfonyldipropionate 31.4	344			[1994WAN/KUO]
C ₁₄ H ₂₇ N	[629-63-0] $\Delta_v H$ $\Delta_v H$ $\Delta_v H$	myristonitrile (327–369) (391–580)	84.2 ± 0.2 71.4 85.3 ± 0.5	298 406 298	GS A C	[2005EME/VER] [1987STE/MAL] [1977STRI/SUN]
C ₁₄ H ₂₇ NO ₃	[na] $\Delta_{\text{fus}}H$	N-dodecanoylglycine 48.4	393.1		DSC	[1986MIY/MAT]
C ₁₄ H ₂₇ NO ₃	[14379-35-2] $\Delta_{\text{trs}}H$ $\Delta_{\text{fus}}H$	N-octanoyl-(l)-leucine 7.6 29.3	357.1 398.1		DSC	[1986MIY/MAT]
C ₁₄ H ₂₇ NO ₃	[107396-11-2] $\Delta_{\text{trs}}H$ $\Delta_{\text{fus}}H$	N-octanoyl-(dl)-leucine 6.8 27.2	353.6 367.1		DSC	[1986MIY/MAT]
C ₁₄ H ₂₈	[295-17-0] $\Delta_{\text{fus}}H$ $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$ $\Delta_v H$	cyclotetradecane 28.7 (300–321) (295–307) (285–290) 62.3 ± 0.2	328 298 310 301 287 343		CGC-DSC HSA ME TM	[1970BOR/DAL] [1998CHI/HES] [1992CHI/HES] [1964FRI/BAU, 1970COX/PIL] [1955ENG] [1992CHI/HES]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound		T _m (K)	Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)			
	$\Delta_v H$		65.3 ± 0.2	298		[1992CHI/HES]
C ₁₄ H ₂₈	[na] $\Delta_v H$	3- <i>tert</i> -butyl-1-methyl-4-isopropylcyclohexane (329–505)	53.8	344	A	[1987STE/MAL]
C ₁₄ H ₂₈	[na] $\Delta_v H$	(1-methylheptyl)cyclohexane (364–397)	60.4	379	A	[1987STE/MAL]
C ₁₄ H ₂₈	[1795-15-9] $\Delta_v H$ $\Delta_v H$	octylcyclohexane (367–399)	62.7 69.8	382 298	A	[1987STE/MAL] [1971WIL/ZWO]
C ₁₄ H ₂₈	[2882-98-6] $\Delta_v H$	nonylcyclopentane	70.7	298		[1971WIL/ZWO]
C ₁₄ H ₂₈	[1120-36-1] $\Delta_v H$ $\Delta_v H$	1-tetradecene (430–527)	70.2 56.5	298 445	A	[1971WIL/ZWO] [1987STE/MAL, 1955CAM/ROS]
C ₁₄ H ₂₈	[54845-26-0] $\Delta_v H$	2,2,3,5,5,6,6-heptamethyl-3-heptene (303–355)	51.2	318	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₁₄ H ₂₈	[4789-35-9] $\Delta_{\text{fus}} H$	<i>trans</i> -1,4-di- <i>tert</i> -butylcyclohexane	17.15	363.2		[1968VAN/HOE]
C ₁₄ H ₂₈	[4789-34-8] $\Delta_{\text{fus}} H$	<i>cis</i> -1,4-di- <i>tert</i> -butylcyclohexane	8.79	293.2		[1968VAN/HOE]
C ₁₄ H ₂₈ N ₂ O ₂	[163678-36-2] $\Delta_{\text{fus}} H$ $\Delta_v H$	tetrapropyloxamide	21.0 67.0	317.2 489	TGA,DSC TGA,DSC	[2003CLO/JAN] [2003CLO/JAN]
C ₁₄ H ₂₈ N ₂ O ₂	[61382-93-2] $\Delta_{\text{fus}} H$	tetradecanediamide	77.45	469.3	DSC	[2006BAD/DEL]
C ₁₄ H ₂₈ O	[5770-04-7] $\Delta_v H$	1-octylcyclohexanol (373–403)	105.6	388	A	[1987STE/MAL]
C ₁₄ H ₂₈ O	[75039-85-9] $\Delta_v H$	(Z) 2-tetradecen-1-ol (353–393)	101.1	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₄ H ₂₈ O	[75039-86-0] $\Delta_v H$	(E) 2-tetradecen-1-ol (353–393)	101.5	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₄ H ₂₈ O	[68892-27-3] $\Delta_v H$	(Z) 3-tetradecen-1-ol (353–393)	99.8	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₄ H ₂₈ O	[68900-86-7] $\Delta_v H$	(E) 3-tetradecen-1-ol (353–393)	99.7	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₄ H ₂₈ O	[40642-41-9] $\Delta_v H$	(Z) 4-tetradecen-1-ol (353–393)	100.0	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₄ H ₂₈ O	[59101-24-5] $\Delta_v H$	(E) 4-tetradecen-1-ol (353–393)	100.7	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₄ H ₂₈ O	[40642-42-0] $\Delta_v H$	(Z) 5-tetradecen-1-ol (353–393)	100.3	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₄ H ₂₈ O	[62936-14-5] $\Delta_v H$	(E) 5-tetradecen-1-ol (353–393)	100.8	298	CGC	[2000OVA/KOU, 1994KOU/HOS]

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₄ H ₂₈ O	[68760-63-4] $\Delta_v H$	(Z) 6-tetradecen-1-ol (353–393)	100	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₄ H ₂₈ O	[68760-62-3] $\Delta_v H$	(E) 6-tetradecen-1-ol (353–393)	100.5	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₄ H ₂₈ O	[40642-43-1] $\Delta_v H$	(Z) 7-tetradecen-1-ol (353–393)	99.9	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₄ H ₂₈ O	[37011-95-3] $\Delta_v H$	(E) 7-tetradecen-1-ol (353–393)	100.5	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₄ H ₂₈ O	[64470-32-2] $\Delta_v H$	(Z) 8-tetradecen-1-ol (353–393)	100.3	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₄ H ₂₈ O	[64437-34-9] $\Delta_v H$	(E) 8-tetradecen-1-ol (353–393)	101.4	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₄ H ₂₈ O	[35153-15-2] $\Delta_v H$	(Z) 9-tetradecen-1-ol (353–393)	100.6	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₄ H ₂₈ O	[52957-16-1] $\Delta_v H$	(E) 9-tetradecen-1-ol (353–393)	101	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₄ H ₂₈ O	[57393-02-9] $\Delta_v H$	(Z) 10-tetradecen-1-ol (353–393)	101.1	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₄ H ₂₈ O	[64437-35-0] $\Delta_v H$	(E) 10-tetradecen-1-ol (353–393)	101.5	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₄ H ₂₈ O	[34010-15-6] $\Delta_v H$	(Z) 11-tetradecen-1-ol (353–393)	101.7	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₄ H ₂₈ O	[35153-18-5] $\Delta_v H$	(E) 11-tetradecen-1-ol (353–393)	101.8	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₄ H ₂₈ O	[70711-48-7] $\Delta_v H$	(Z) 12-tetradecen-1-ol (353–393)	102.5	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₄ H ₂₈ O	[70711-49-8] $\Delta_v H$	(E) 12-tetradecen-1-ol (353–393)	102.5	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₄ H ₂₈ O	[2345-27-9] $\Delta_{\text{fus}} H$	2-tetradecanone	49.12	306.7		[1996DOM/HEA]
	$\Delta_{\text{sub}} H$		130.9 ± 0.5	298	C	[1979SUN/SVE2]
	$\Delta_v H$	(411–560)	65.6	426	A	[1987STE/MAL]
	$\Delta_v H$	(549–643)	55.6	564	A	[1987STE/MAL]
	$\Delta_v H$		82.1 ± 0.6	298	S-F	[1979SUN/SVE2]
	$\Delta_v H$	(412–643)	51.6	556		[1975AMB/ELL]
C ₁₄ H ₂₈ O	$\Delta_v H$	(372–551)	64.4	387	A	[1987STE/MAL, 1947STU]
	[6137-34-4] $\Delta_v H$	7-tetradecanone (438–462)	66.9	450	A, ME	[1987STE/MAL, 1938UBB]
	[124-25-4] $\Delta_v H$	tetradecanal (334–370)	77.4 ± 0.4	298	GS	[2003VER/KRA2]
	$\Delta_v H$	(343–383)	80.2	298	CGC	[1996KOU/HOS, 2000OVA/KOU]
	$\Delta_v H$	(372–571)	63.4	387	A	[1987STE/MAL, 1947STU]
C ₁₄ H ₂₈ O ₂	[112-66-3]	dodecyl acetate				

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(289–333)	79.6 ± 0.3	298	GS	[2006KRA/VER]
	$\Delta_v H$	(333–378)	81.8	298	GC	[1997KOU/HOS, 2000OVA/KOU]
	$\Delta_v H$	(398–540)	70.5	413	A	[1987STE/MAL]
C ₁₄ H ₂₈ O ₂	[106-33-2]	ethyl dodecanoate				
	$\Delta_{\text{fus}}H$		9.31	271.5		[1996DOM/HEA]
	$\Delta_v H$	(423–483)	80.0	298	GC	[1997KRO/VEL]
	$\Delta_v H$	(386–435)	67.2	401	A	[1987STE/MAL]
C ₁₄ H ₂₈ O ₂	[1731-88-0]	methyl tridecanoate				
	$\Delta_v H$		74.0	350		[2002VAN/VAN]
	$\Delta_v H$		72.3 ± 0.1	368		[2002VAN/VAN]
	$\Delta_v H$		80.0 ± 0.5	298		[2002VAN/VAN]
	$\Delta_v H$		81.3 ± 0.7	298	GC, C	[1980FUC/PEA]
	$\Delta_v H$		82.7 ± 0.8	298	C	[1977MAN/SEL]
	$\Delta_v H$	(377–504)	72.6	392	A, EST	[1987STE/MAL, 1963ROS/SCH]
C ₁₄ H ₂₈ O ₂	[245658-44-0]	2,2-dimethylpropanoic acid, 1,1,5-trimethylhexyl ester				
	$\Delta_v H$	(333–378)	61.3	298	CGC	[1999VER/HEI]
C ₁₄ H ₂₈ O ₂	[544-63-8]	tetradecanoic acid				
	$\Delta_{\text{fus}}H$		45.75	326.2	DSC	[2010HON/HUA]
	$\Delta_{\text{fus}}H$		1.8	315		
	$\Delta_{\text{fus}}H$		6.4	325.3		
	$\Delta_{\text{fus}}H$		45.0	326.5	DSC	[2007MOR/COR]
	$\Delta_{\text{fus}}H$		40.1	326.6	DSC	[2007MIS/MIS]
	$\Delta_{\text{fus}}H$		45.1	327		[1996DOM/HEA]
	$\Delta_{\text{sub}}H$		168.6 ± 9	298	TPD	[2008CAP/LOV]
	$\Delta_{\text{sub}}H$	(272–288)		125.6	TPTD	[2005CHA/ZIE]
	$\Delta_{\text{sub}}H$	(282–305)		174	TPTD	[2001CHA/TOB]
	$\Delta_{\text{sub}}H$	(312–325)	139.7 ± 3.8	318	ME	[1961DAV/MAL, 1970COX/PIL]
	$\Delta_v H$	(383–459)	100.4	398	A	[1987STE/MAL]
	$\Delta_v H$	(423–599)	91.6	438	A	[1987STE/MAL]
	$\Delta_v H$	(339–358)	104.1 ± 2.0	349	ME, TE	[1982DEK/SCH]
	$\Delta_v H$		88.9	455	I	[1943CRA]
C ₁₄ H ₂₈ O ₃	[na]	decyl 3-methoxypropionate				
	$\Delta_v H$	(403–513)	68.9	418	A	[1987STE/MAL]
C ₁₄ H ₂₈ O ₃	[19816-73-0]	peroxytetradecanoic acid				
	$\Delta_{\text{sub}}H$	(293–303)	156.0 ± 4.1		ME	[1980SWA/KWA]
C ₁₄ H ₂₈ O ₄	[56444-61-2]	2,2,9,9-tetramethyl-1,3,8,10-tetraoxacyclotetradecane				
	$\Delta_{\text{fus}}H$		30.5	409.4		[1994KIM/LEE]
C ₁₄ H ₂₈ O ₄	[55208-76-9]	3,3,6,6-tetrapropyl-1,2,4,5-tetraoxacyclohexane				
	$\Delta_v H$	(403–473)	65.1	298	CGC	[2007CAN/EYL]
C ₁₄ H ₂₈ O ₆	[125590-73-0]	2-ethylhexyl α -(D)-glucoside				
	$\Delta_{\text{fus}}H$		33.47	341.2		
	$\Delta_{\text{fus}}H$		3.56	387.2	DSC	[1998NIL/SOE]
C ₁₄ H ₂₈ O ₆	[125590-74-1]	2-ethylhexyl β -(D)-glucoside				
	$\Delta_{\text{fus}}H$		10.88	330.2	DSC	[1998NIL/SOE]
C ₁₄ H ₂₉ Br	[112-71-0]	1-bromotetradecane				

Note: Experimental values based on the TPTD method are often inconsistent with values determined using other experimental methods

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(437–645)	67.1	452	A, EST	[1987STE/MAL, 1961LI/ROS, 1970DYK/VAN]
C ₁₄ H ₂₉ Cl	[2425-54-9]	1-chlorotetradecane				
	$\Delta_v H$		86.6	298		[2006BOL/NER2]
	$\Delta_v H$	(313–373)	80.2	313	GC	[1980JON/MAT]
	$\Delta_v H$	(313–373)	78.0	333	GC	[1980JON/MAT]
	$\Delta_v H$	(313–373)	74.4	353	GC	[1980JON/MAT]
	$\Delta_v H$	(313–373)	72.9	373	GC	[1980JON/MAT]
C ₁₄ H ₂₉ F	[593-33-9]	1-fluorotetradecane				
	$\Delta_v H$	(288–335)	73.5 ± 0.4	298	GS	[1997SCH/VER]
	$\Delta_v H$	(400–593)	61.4	415	A, EST	[1987STE/MAL, 1961LI/ROS, 1970DYK/VAN]
C ₁₄ H ₂₉ I	[19218-94-1]	1-iodotetradecane				
	$\Delta_v H$	(452–672)	90.0	298	A, EST	[1987STE/MAL, 1961LI/ROS, 1970DYK/VAN, 2006BOL/NER]
	$\Delta_v H$	(452–672)	68.6	467	A, EST	[1987STE/MAL, 1961LI/ROS, 1970DYK/VAN]
C ₁₄ H ₂₉ NO	[638-58-4]	tetradecanamide				
	$\Delta_{\text{sub}} H$	(248–375)	167.4 ± 2.5	352	ME	[1959DAV/JON2, 1987STE/MAL]
C ₁₄ H ₃₀	[629-54-4]	tetradecane				
	$\Delta_{\text{fus}} H$		42.7	278.3	DSC	[2004MON/RAJ]
	$\Delta_{\text{fus}} H$		45.07	279		[1996DOM/HEA]
	$\Delta_{\text{sub}} H$		117.6	298	B	[1972MOR3]
	$\Delta_v H$	(283–313)	71.6 ± 1.3	298	GS	[1909LEG/BAC]
	$\Delta_v H$		72.1	298	GS	[2001PUR/CHI]
	$\Delta_v H$		72.0 ± 2.4	298	CGC	[2000NIC/ORF]
	$\Delta_v H$		69.0	324	C	[1996VIT/CHA]
	$\Delta_v H$		68.6	329	C	[1996VIT/CHA]
	$\Delta_v H$		67.9	334	C	[1996VIT/CHA]
	$\Delta_v H$		66.8	344	C	[1996VIT/CHA]
	$\Delta_v H$		65.7	359	C	[1996VIT/CHA]
	$\Delta_v H$	(423–473)	71.2	298	CGC	[1995CHI/HOS]
	$\Delta_v H$	(363–413)	71.4	298	CGC	[1995CHI/HOS]
	$\Delta_v H$		71.7	298		[1994RUZ/MAJ]
	$\Delta_v H$	(313–433)	67.8	328	A	[1987STE/MAL]
	$\Delta_v H$	(343–395)	64.1	361	GS	[1986ALL/JOS]
	$\Delta_v H$		70.1	313	C	[1979SUN/SVE]
	$\Delta_v H$		68.9	328	C	[1979SUN/SVE]
	$\Delta_v H$		71.8 ± 0.6	298	C	[1979SUN/SVE]
	$\Delta_v H$		71.1 ± 0.4	298	C	[1972MOR2]
$\Delta_v H$		71.7	298		[1971WIL/ZWO]	
$\Delta_v H$	(432–529)	57.1	447	A	[1987STE/MAL, 1955CAM/ROS]	
$\Delta_v H$	(429–468)	57.8	449	ME	[1938UBB]	
C ₁₄ H ₃₀	[1560-96-9]	2-methyltridecane				
	$\Delta_v H$	(388–530)	56.3	403	A	[1987STE/MAL]
C ₁₄ H ₃₀	[6418-41-3]	3-methyltridecane				
	$\Delta_v H$	(389–521)	55.1	404	A	[1987STE/MAL]
C ₁₄ H ₃₀	[26730-12-1]	4-methyltridecane				

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound		T _m (K)	Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)			
	$\Delta_v H$	(386–520)	54.2	401	A	[1987STE/MAL]
C ₁₄ H ₃₀	[25117-31-1]	5-methyltridecane				
	$\Delta_v H$	(385–518)	53.8	400	A	[1987STE/MAL]
C ₁₄ H ₃₀	[26730-14-3]	7-methyltridecane				
	$\Delta_v H$	(357–389)	59.0	372	A	[1987STE/MAL]
C ₁₄ H ₃₀	[6117-98-2]	2,3-dimethyldodecane				
	$\Delta_v H$	(385–519)	53.4	400	A	[1987STE/MAL]
C ₁₄ H ₃₀	[6117-99-3]	2,4-dimethyldodecane				
	$\Delta_v H$	(379–509)	54.0	394	A	[1987STE/MAL]
C ₁₄ H ₃₀	[na]	2,4,6-trimethylundecane				
	$\Delta_v H$	(368–491)	53.2	383	A	[1987STE/MAL]
C ₁₄ H ₃₀	[5171-86-8]	2,2,3,4,6,6-heptamethylheptane				
	$\Delta_v H$	(313–366)	54.5	366	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₁₄ H ₃₀	[5171-86-8]	hexaethylethane (3,3,4,4-tetraethylhexane)				
	$\Delta_v H$	(298–307)	63.9 ± 1.2	298	GS	[1997VER/NOL]
	$\Delta_v H$	(283–302)	65.7 ± 1.2	292	GS	[1973BEC/RUC, 1995CHI/HES]
	$\Delta_v H$		65.0 ± 1.2	298		[1973BEC/RUC]
C ₁₄ H ₃₀	[65149-84-0]	2,2,3,3,4,4,5,5-octamethylhexane				
	$\Delta_v H$	(288–325)	56.9 ± 0.7	298	GS	[1997VER/NOL]
C ₁₄ H ₃₀ N ₂ O	[842173-55-1]	1-tridecyl urea				
	$\Delta_{\text{us}}H$		1.5	261.6		
	$\Delta_{\text{us}}H$		2.8	306.5		
	$\Delta_{\text{fus}}H$		46.0	384.6	DSC	[2005HAS/TAJ]
C ₁₄ H ₃₀ O	[629-64-1]	diheptyl ether				
	$\Delta_v H$	(360–547)	63.1	375	A	[1987STE/MAL]
C ₁₄ H ₃₀ O	[na]	4-methylpentyl <i>tert</i> -octyl ether				
	$\Delta_v H$		57.5	298	CGC	[UR/VER, 2002VER, 2003VER/KRA]
C ₁₄ H ₃₀ O	[na]	3-methylpentyl <i>tert</i> -octyl ether				
	$\Delta_v H$		58.0	298	CGC	[UR/VER, 2002VER, 2003VER/KRA]
C ₁₄ H ₃₀ O	[na]	3,3-dimethylbutyl <i>tert</i> -octyl ether				
	$\Delta_v H$		56.4	298	CGC	[UR/VER, 2002VER, 2003VER/KRA]
C ₁₄ H ₃₀ O	[508181-44-0]	hexyl <i>tert</i> -octyl ether				
	$\Delta_v H$	(296–326)	59.8 ± 0.6	298	GS	[2003VER/KRA]
	$\Delta_v H$		59.2	298		[UR/VER, 2002VER]
C ₁₄ H ₃₀ O	[112-72-1]	1-tetradecanol				
	$\Delta_{\text{fus}}H$		47.29	308.1	DSC	[2009ZEN/CAO]
	$\Delta_{\text{fus}}H + \Delta_{\text{us}}H$		47.01	311.2		
	$\Delta_{\text{us}}H$		25.1	310.8		
	$\Delta_{\text{us}}H$		1.8	306		
	$\Delta_{\text{us}}H$		23.81	311		
	$\Delta_{\text{fus}}H$		22.01	311.6		
	$\Delta_{\text{fus}}H + \Delta_{\text{us}}H$		49.37	311		[1974MOS/MOU]
	$\Delta_{\text{sub}}H$		126.0 ± 0.6			[1977MAN/SEL]
	$\Delta_{\text{sub}}H$	(293–307)	143.9	300	ME	[1965DAV/KYB]
$\Delta_v H$		98.9 ± 2.5	298	CGC	[2006NIC/KWE]	

TABLE 9. Phase change enthalpies of C₁₁ to C₁₄ organic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(312–346)	93.6	328	GS	[2001KUL/VER2]
	$\Delta_v H$	(312–346)	98.7	298	GS	[2001KUL/VER2]
	$\Delta_v H$	(333–438)	81.8	386		[1992NGU/KAS]
	$\Delta_v H$	(317–358)	109	332	A	[1987STE/MAL]
	$\Delta_v H$		102.2 ± 2.3	298	C	[1977MAN/SEL]
	$\Delta_v H$	(313–358)	106.4	328		[1973WIL/ZWO]
	$\Delta_v H$	(424–569)	76.6	439	A	[1987STE/MAL, 1969KEM/KRE]
	$\Delta_v H$	(313–326)	104.2	320	ME	[1965DAV/KYB]
C₁₄H₃₀O	[4706-81-4]	2-tetradecanol				
	$\Delta_v H$	(313–428)	95.7	328		[1999NGU/BER]
C₁₄H₃₀O₂	[4536-30-5]	2-(dodecyloxy)ethanol				
	$\Delta_v H$	(414–467)	71.5	429	A	[1987STE/MAL]
C₁₄H₃₀O₂	[19812-64-7]	1,14-tetradecanediol				
	$\Delta_{\text{fus}} H$		61.9	360.4	DSC	[1999OGA/NAK]
C₁₄H₃₀O₂S	[126835-75-4]	3-(undecylthio)-1,2-propanediol				
	$\Delta_{\text{us}} H$		2.5	280.2		
	$\Delta_{\text{us}} H$		4.9	289.1		
	$\Delta_{\text{us}} H$		4.6	295.2		
	$\Delta_{\text{fus}} H$		18.3	317.4		[1993ACR]
C₁₄H₃₀O₃	[10430-98-5]	3-(undecyloxy)-1,2-propanediol				
	$\Delta_{\text{fus}} H$		43.1	311.7	DSC	[1993ACR]
C₁₄H₃₀O₄S₂	[na]	2-deoxy-(D)-glucose dibutyl dithioacetal				
	$\Delta_{\text{fus}} H$		60.3	409.5	DSC	[1989VAN/VAN]
C₁₄H₃₀O₄S₂	[na]	(l)-rhamnose dibutyl dithioacetal				
	$\Delta_{\text{fus}} H$		37.9	389.9	DSC	[1989VAN/VAN]
		Note: Authors report that there are several transitions prior to melting.				
C₁₄H₃₀O₅S₂	[115395-52-3]	(D)-glucose dibutyl dithioacetal				
	$\Delta_{\text{fus}} H$		50.2	399	DSC	[1989VAN/VAN]
C₁₄H₃₀O₅S₂	[68747-93-3]	(D)-galactose dibutyl dithioacetal				
	$\Delta_{\text{fus}} H$		46.4	399.2	DSC	[1989VAN/VAN]
C₁₄H₃₀S	[2079-95-0]	1-tetradecanethiol				
	$\Delta_v H$	(446–614)	67.3	461		[1999DYK/SVO]
C₁₄H₃₀S₂	[10496-16-9]	diheptyl disulfide				
	$\Delta_v H$	(458–630)	69.8	473		[1999DYK/SVO]
C₁₄H₃₁N	[2470-68-0]	diheptylamine				
	$\Delta_v H$	(435–605)	60.0	450	A	[1987STE/MAL]
C₁₄H₃₁N	[112-18-5]	N,N-dimethyldodecylamine				
	$\Delta_v H$	(283–324)	69.5	299		[2004FUL/RUZ]
	$\Delta_v H$	(380–604)	64.4	395	A	[1987STE/MAL]
C₁₄H₃₁N	[2016-42-4]	tetradecylamine				
	$\Delta_v H$	(471–577)	62.4	486	A, EST	[1987STE/MAL, 1956MAN2]
C₁₄H₃₁NO₂	[126835-66-3]	3-(undecylamino)-1,2-propanediol				
	$\Delta_{\text{fus}} H$		58.2	348.8		[1993ACR]
C₁₄H₃₁O₂P	[na]	diheptylphosphinic acid				
	$\Delta_v H$	(482–664)	64.1	573		[1971NAK/SMI]

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₅ H ₈ Cl ₃ NO ₂	[77765-38-9] $\Delta_v H$	2,2,4-trichloro-5-(2-naphthalenylamino)-4-cyclopentene-1,3-dione (453–483)	91.4	468	GC	[1980SHA/SAD]
C ₁₅ H ₉ N	[1210-12-4] $\Delta_{\text{fus}} H$	9-cyanoanthracene	25.19	445.2		[1970GUA/SAR]
C ₁₅ H ₉ N ₃	[217-88-9] $\Delta_v H$	pyrido[2,3-f] [1,7]phenanthroline (648–707)	65.1	663	A	[1987STE/MAL, 1962JOH/MCE]
C ₁₅ H ₉ N ₃	[217-81-2] $\Delta_v H$	pyrido[3,2-f] [1,7]phenanthroline (648–706)	67.4	663	A, I	[1987STE/MAL, 1962JOH/MCE]
C ₁₅ H ₁₀	[203-64-5] $\Delta_v H$	4 <i>H</i> -cyclopenta[def]phenanthrene 83.4 ± 0.7		298	CGC	[2008HAN/NUT]
C ₁₅ H ₁₀ ClFN ₂ O	[2886-65-9] $\Delta_{\text{fus}} H$	7-chloro-1,3-dihydro-5-(2'-fluorophenyl)-2 <i>H</i> -1,4-benzodiazapin-2-one (desalkylflurazepam)	30.7	481.2	DSC	[2008WAS/HOL]
C ₁₅ H ₁₀ Cl ₂ N ₂ O ₂	[50264-69-2] $\Delta_{\text{fus}} H$	1-[(2,4-dichlorophenyl)methyl]-1- <i>H</i> -indazole-3-carboxylic acid	45.92	480.2	DSC	[1998PAL/WEH]
C ₁₅ H ₁₀ N ₂ O ₂	[2536-05-2] $\Delta_v H$	2,2'-diisocyanatodiphenylmethane (343–413)	90.1	358	A	[1987STE/MAL]
C ₁₅ H ₁₀ N ₂ O ₂	[5873-54-1] $\Delta_v H$	2,4'-diisocyanatodiphenylmethane (343–413)	89.3	358	A	[1987STE/MAL]
C ₁₅ H ₁₀ N ₂ O ₂	[101-68-8] $\Delta_{\text{fus}} H$	4,4'-diisocyanatodiphenylmethane	27.3	313.6		[1996DOM/HEA, 1977LEB/EVS]
	$\Delta_v H$	(343–413)	90.5	358	A	[1987STE/MAL]
	$\Delta_v H$	(442–530)	93.8	457	A	[1987STE/MAL]
	$\Delta_v H$	(442–530)	90.6	483	A	[1966ZAL/STR]
C ₁₅ H ₁₀ O	[886-38-4] $\Delta_{\text{sub}} H$	diphenylcyclopropenone (353–378)	119.7 ± 8	365	HSA	[1985STE/GAN]
	$\Delta_{\text{sub}} H$	(323–343)	141 ± 4	333	ME	[1976HOP/BOS, 1987STE/MAL]
C ₁₅ H ₁₀ O	[642-31-9] $\Delta_{\text{fus}} H$	9-anthraldehyde	17.61	377.2		[1970GUA/SAR]
	$\Delta_{\text{sub}} H$	(329–363)	100.6 ± 3.9		ME	[2008GOL/SUU]
C ₁₅ H ₁₀ O	[1139-82-8] $\Delta_{\text{fus}} H$	5,7-dihydro-6 <i>H</i> -dibenzo[a,c]cyclohepten-6-one	18.16	350.3	DSC	[1998VER4]
C ₁₅ H ₁₀ O ₂	[na] $\Delta_{\text{sub}} H$	α -benzoyloxypthalide (343–388)	U 125.3	366		[1989ROR/RUT]
C ₁₅ H ₁₀ O ₂	[613-08-1] $\Delta_{\text{sub}} H$	2-anthracenecarboxylic acid (401–421)	134.8 ± 3.4		ME	[2008GOL/SUU]
C ₁₅ H ₁₀ O ₂	[723-62-6] $\Delta_{\text{sub}} H$	9-anthracenecarboxylic acid (385–420)	120.1 ± 3.8		ME	[2008GOL/SUU]
C ₁₅ H ₁₀ O ₂	[525-82-6] $\Delta_{\text{fus}} H$	2-phenyl-4 <i>H</i> -1-benzopyran-4-one (flavone)	20.32	369.9	DSC	[2009SOU/MAT]
	$\Delta_{\text{sub}} H$		108.2 ± 1.7	298	C	[2009SOU/MAT]
C ₁₅ H ₁₀ O ₃	[82-39-3] $\Delta_{\text{sub}} H$	1-methoxy-9,10-anthraquinone	128		GS	[1987SHI/OHK, 1991HOR]
	$\Delta_{\text{sub}} H$		106.6	385	HSA	[1956BEY/NIC]

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		T _m (K)	Method	Reference
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)			
C ₁₅ H ₁₀ O ₃	[3274-20-2]	2-methoxy-9,10-anthraquinone				
	$\Delta_{\text{sub}}H$		124.7		GS	[1987SHI/OHK, 1991HOR]
			118.4 ± 0.4	419	HSA	[1956BEY/NIC]
C ₁₅ H ₁₀ O ₃	[60466-75-3]	9-methoxy-1,4-anthraquinone				
	$\Delta_{\text{sub}}H$	(363–386)	130.5 ± 2.3	375	ME	[2002JIM/ROU]
		(363–386)	131.5 ± 2.3	298	ME	[2002JIM/ROU]
C ₁₅ H ₁₀ O ₄	[480-40-0]	5,7-dihydroxy-2-phenyl-4 <i>H</i> -1-benzopyran-4-one (chrysin)				
	$\Delta_{\text{fus}}H$		39.2	558.2	DSC	[2007CHE/HUM]
C ₁₅ H ₁₀ O ₇	[117-39-5]	2-(3,4-dihydroxyphenyl)-3,5,7-trihydroxy-4 <i>H</i> -1-benzopyran-4-one (quercetin)				
	$\Delta_{\text{fus}}H$		41.5	595.2	DSC	[2007CHE/HUM]
C ₁₅ H ₁₁ ClF ₃ NO ₄	[42874-03-3]	2-chloro-1-(3-ethoxy-4-nitrophenoxy)-4-(trifluoromethyl)benzene				
	$\Delta_{\text{fus}}H$		30.07	358.8	DSC	[1991ACR, 1990DON/DRE]
C ₁₅ H ₁₁ ClN ₂ O	[1088-11-5]	7-chloro-1,3-dihydro-5-phenyl-2 <i>H</i> -1,4-benzodiazepin-2-one (nordazepam)				
	$\Delta_{\text{fus}}H$ (I)		24.45	494.5		
	$\Delta_{\text{fus}}H$ (II)		34	489.9		
	$\Delta_{\text{fus}}H$ (III)		27.4	489.2		
	$\Delta_{\text{fus}}H$ (IV)		33.62	487.4	TGA	[1996DOM/HEA, 1992CHA/MOU]
C ₁₅ H ₁₁ Cl ₂ N ₂ O ₂	[846-49-1]	7-chloro-5-(2-chlorophenyl)-1,3-dihydro-3-hydroxy-2 <i>H</i> -1,4-benzodiazepin-2-one ((±)lorazepam)				
	$\Delta_{\text{fus}}H$		75.2	453.2	DSC	[2008WAS/HOL]
	$\Delta_{\text{fus}}H$		92.57	446.5	DSC	[2001VER/AUG]
C ₁₅ H ₁₁ F ₃ O ₃	[3119-86-6]	2-hydroxy-2'-trifluoromethyl-4-methoxybenzophenone				
	$\Delta_{\text{sub}}H$	(323–363)	U 13.3	338	EV	[1987STE/MAL, 1966GRA/BUR]
C ₁₅ H ₁₁ F ₃ O ₃	[7396-89-6]	2-hydroxy-3'-trifluoromethyl-4-methoxybenzophenone				
	$\Delta_{\text{sub}}H$	(313–323)	103.8	318	EV	[1987STE/MAL, 1966GRA/BUR]
C ₁₅ H ₁₁ F ₃ O ₃	[7396-90-9]	2-hydroxy-4'-trifluoromethyl-4-methoxybenzophenone				
	$\Delta_{\text{sub}}H$	(313–333)	91.0	323	EV	[1987STE/MAL, 1966GRA/BUR]
C ₁₅ H ₁₁ N	[612-96-4]	2-phenylquinoline				
	$\Delta_{\text{sub}}H$	(337–351)	103.1 ± 0.8	344	ME	[1997RIB/MAT3]
	$\Delta_{\text{sub}}H$		105.4 ± 0.9	298	ME	[1997RIB/MAT3]
C ₁₅ H ₁₁ NO ₂	[82-38-2]	1-methylamino-9,10-anthraquinone				
	$\Delta_{\text{fus}}H$		28.81	443.2		[1991BAU/WEB]
	$\Delta_{\text{sub}}H$		112.6			[1984KAR/KRU]
	$\Delta_{\text{sub}}H$	(363–383)	115.9 ± 3.5	373		[1984KRI]
	$\Delta_{\text{sub}}H$	(384–405)	123.8 ± 3.3	395	ME	[1960BRA/BIR, 1987STE/MAL]
	$\Delta_{\text{sub}}H$		123.8		ME	[1964JON/SED, 1991HOR, 1966JON/KRA]
	$\Delta_{\text{sub}}H$		115.5 ± 0.4	461	HSA	[1956BEY/NIC]
	$\Delta_{\text{sub}}H$		114.7 ± 3	406	HSA	[1956BEY/NIC]
	Δ_vH	(433–493)	103.5	448	A	[1987STE/MAL]
C ₁₅ H ₁₁ NO ₂	[82-28-0]	1-amino-2-methyl-9,10-anthraquinone				
	$\Delta_{\text{sub}}H$	(360–388)	124.6 ± 7.3	374		[1984KRI]
C ₁₅ H ₁₁ NO ₃ S	[313057-09-9]	4-(2-propenyloxy)phenyl 5-cyano-2-thiophene carboxylate				
	$\Delta_{\text{fus}}H$		103.8	361.5	DSC	[2000WU/WAN]
C ₁₅ H ₁₁ NO ₄	[na]	1-amino-2-methoxy-4-hydroxy-9,10-anthraquinone				
	$\Delta_{\text{sub}}H$		132			[1984KAR/KRU]
C ₁₅ H ₁₁ N ₃ O ₂	[6407-80-3]	4-hydroxy-3-(phenylazo)-2(1 <i>H</i>)-quinolinone (Disperse Yellow 4)				

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		T _m (K)	Method	Reference
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)			
		$\Delta_{\text{sub}}H$		127.2		[1968TSU/KOJ, 1988BAU/PER]
C ₁₅ H ₁₂	[610-48-0]	1-methylanthracene				
		$\Delta_{\text{v}}H$	87.0 ± 1.0	298	CGC	[2008HAN/NUT]
C ₁₅ H ₁₂	[613-12-7]	2-methylanthracene				
		$\Delta_{\text{v}}H$	84.5 ± 2.7	298	CGC	[2008HAN/NUT]
		$\Delta_{\text{v}}H$	(413–473) 84.4 ± 1.2	298	GC	[2006HAF/PAR]
		$\Delta_{\text{v}}H$	(323–473) 76.1	398	GC	[2002LEI/CHA]
C ₁₅ H ₁₂	[779-02-0]	9-methylanthracene				
		$\Delta_{\text{sub}}H$	(329–345) 99.8 ± 1.0	337	ME	[2006RIB/AMA2]
		$\Delta_{\text{sub}}H$	(329–345) 101.8 ± 1.0	298	ME	[2006RIB/AMA2]
		$\Delta_{\text{sub}}H$	98.9		RG	[1958KLO]
		$\Delta_{\text{v}}H$	88.1 ± 1.0	298	CGC	[2008HAN/NUT]
		$\Delta_{\text{v}}H$	(354–402) 98.9	369	A	[1987STE/MAL]
		$\Delta_{\text{v}}H$	(423–587) 58.5	465		[1983SIV/KOB]
		$\Delta_{\text{v}}H$	(423–515) 58.1	515		[1983SIV/KOB]
		$\Delta_{\text{v}}H$	(423–515) 56.5	555		[1983SIV/KOB]
C ₁₅ H ₁₂	[832-69-9]	1-methylphenanthrene				
		$\Delta_{\text{v}}H$	84.5 ± 1.4	298	CGC	[2008HAN/NUT]
		$\Delta_{\text{v}}H$	(323–473) 76.3	398	GC	[2002LEI/CHA]
C ₁₅ H ₁₂	[832-64-4]	4-methylphenanthrene				
		$\Delta_{\text{trs}}H$	0.02	182		
		$\Delta_{\text{trs}}H$	0.03	295		
		$\Delta_{\text{fus}}H$	14.04	324.9		[1996DOM/HEA]
		$\Delta_{\text{v}}H$	(368–647) 74.4 ± 0.2	380	EB,IP	[1989CHI/HOS]
		$\Delta_{\text{v}}H$	(368–647) 71.8 ± 0.1	420	EB,IP	[1989CHI/HOS]
		$\Delta_{\text{v}}H$	(368–647) 69.2 ± 0.1	460	EB,IP	[1989CHI/HOS]
		$\Delta_{\text{v}}H$	(368–647) 66.7 ± 0.1	500	EB,IP	[1989CHI/HOS]
		$\Delta_{\text{v}}H$	(368–647) 64.2 ± 0.1	540	EB,IP	[1989CHI/HOS]
		$\Delta_{\text{v}}H$	(368–647) 61.6 ± 0.1	580	EB,IP	[1989CHI/HOS]
C ₁₅ H ₁₂	[4505-48-0]	2-phenylindene				
		$\Delta_{\text{v}}H$	84.3 ± 0.7	298	CGC	[2008HAN/NUT]
C ₁₅ H ₁₂ Br ₄ O ₂	[79-94-7]	2,2',6,6'-tetrabromo-4,4-isopropylidenediphenol				
		$\Delta_{\text{fus}}H$	29.1	451.5	DSC	[2008KUR/KAW]
		$\Delta_{\text{sub}}H$	153 ± 3		ME	[2008KUR/KAW]
C ₁₅ H ₁₂ ClN ₂ O ₂	[604-75-1]	7-chloro-1,3-dihydroxy-5-phenyl-2 <i>H</i> -1,4-benzodiazepin-2-one ((±)-oxazepam)				
		$\Delta_{\text{fus}}H$	86.4	478.8	DSC	[2008WAS/HOL]
		$\Delta_{\text{fus}}H$	84.11	467.5	DSC	[2001VER/AUG]
C ₁₅ H ₁₂ ClN ₅ O ₄	[na]	5-[(4-chloro-2-nitrophenylazo)-1-ethyl-1,2-dihydro-6-hydroxy-4-methyl-2-oxo-3-pyridinecarbonitrile				
		$\Delta_{\text{fus}}H$	35.16	500.2		[1991BAU/WEB]
C ₁₅ H ₁₂ N ₂	[668-94-0]	4,5-diphenylimidazole				
		$\Delta_{\text{fus}}H$	32.34	505	DSC	[2007SIF/AIT]
C ₁₅ H ₁₂ N ₂ O	[298-46-4]	5 <i>H</i> -dibenz[b,f]azepine-5-carboxamide (carbamazepine)				
		$\Delta_{\text{fus}}H$	25.6	465.3	DSC	[2009GOO/ROD]
		$\Delta_{\text{fus}}H$ (I)	25.52	466.7		
		$\Delta_{\text{fus}}H$ (II)	26.82	464.4		
		$\Delta_{\text{fus}}H$ (III)	24.89	464.7	DSC	[2003GRZ/LAN]

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
	Enthalpy							
C ₁₅ H ₁₂ N ₂ O ₂	[1220-94-6]		1-amino-4-(N-methylamino)anthra-9,10-quinone		140.6		GS	[1967DAT/KAN, 1991HOR]
C ₁₅ H ₁₂ N ₂ O ₂	[51-41-0]		5,5-diphenyl-2,4-imidazolidinedione (phenytoin)		40.1	568.8	DSC	[2006WAS/HOL, 2008WAS/HOL]
		$\Delta_{\text{fus}}H$			47.08	570.8	DSC	[2003NOK/BOL]
C ₁₅ H ₁₂ N ₂ O ₂	[52955-48-3]		N-(N'-methylanilino)phthalamide		3.6	374		
		$\Delta_{\text{fus}}H$			21.7	399	DSC	[1998BOT/ELL]
C ₁₅ H ₁₂ N ₂ O ₂	[28721-07-5]		10,11-dihydro-10-oxo-5H-dibenz[b,f]azepine-5-carboxamide (oxcarbazepine)		40.3	495.6	DSC	
		$\Delta_{\text{fus}}H$ (I)			33.3	491.4	DSC	
		$\Delta_{\text{fus}}H$ (III)			26.07	486.2	DSC	[2010LUT/MAT]
Note: All three polymorphic forms decomposed on melting.								
C ₁₅ H ₁₂ N ₂ O ₂	[57-41-0]		5,5-diphenylhydantoin		36.29	574		[1985OHM/LIP]
C ₁₅ H ₁₂ N ₂ O ₃	[2872-48-2]		1,4-diamino-2-methoxyanthra-9,10-quinone		35.29	515.2		[1988BAU/PER]
		$\Delta_{\text{sub}}H$			147.0			[1984KAR/KRU]
		$\Delta_{\text{sub}}H$			151.9		GS	[1967DAT/KAN, 1991HOR]
C ₁₅ H ₁₂ N ₄ O ₂	[340820-68-0]		4-phenyl-5-(2-pyridinyl)-4H-1,2,4-triazole-3-carboxylic acid, methyl ester		24.4	465.2		[2005SIK/MOD]
C ₁₅ H ₁₂ O	[1210-35-1]		dibenzosuberone		17.15	305.5	DSC	[1998VER4]
		$\Delta_{\text{sub}}H$			109.3	298		[1998VER4]
		$\Delta_{\text{v}}H$		(314–338)	90.0 ± 1.5		GS	[1998VER4]
C ₁₅ H ₁₂ O	[1139-82-8]		5,7-dihydro-6H-dibenzo[a,c]cyclohepten-6-one		95.6 ± 0.8	298		[1998VER4]
C ₁₅ H ₁₂ OS	[na]		monothiodibenzoylmethane		125.5 ± 4.9	298	C	[2004RIB/SAN3]
		$\Delta_{\text{sub}}H$						
C ₁₅ H ₁₂ O ₂	[120-46-7]		dibenzoylmethane		113.3 ± 4.8	298	C	[2004RIB/SAN3]
		$\Delta_{\text{sub}}H$			115.7 ± 0.9	298	ME	[1992RIB/MON]
C ₁₅ H ₁₂ O ₂	[120-46-7]		1,3-diphenyl-1,3-propanedione	(368–383)	60.1	375	A	[1987STE/MAL]
C ₁₅ H ₁₂ O ₂	[487-26-3]		2,3-dihydro-2-phenyl-4H-1-benzopyran-4-one (flavanone)		21.04	349.5	DSC	[2009SOU/MAT]
		$\Delta_{\text{sub}}H$			107.2 ± 2.3	298	C	[2009SOU/MAT]
C ₁₅ H ₁₂ O ₅	[480-41-1]		2,3-dihydro-5,7-dihydroxy-2-(4-hydroxyphenyl)-4H-1-benzopyran-4-one (naringenin)		39.8	523.2	DSC	[2007CHE/HUM]
C ₁₅ H ₁₃ ClN ₂ O ₅	[1562-85-2]		gallocyanine (C. I. Disperse Blue 95)	(433–493)	88.2	448	A	[1987STE/MAL]
C ₁₅ H ₁₃ ClN ₂ S	[688319-94-0]		N-(2-methyl-4-chlorophenyl)-4H-3,1-benzothiazin-2-amine		17.5	495.8	DSC	[2004GON/KOS]

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
	Enthalpy							
C ₁₅ H ₁₃ Cl ₂ NO ₂	[117-27-1]		1,1-(di-p-chlorophenyl)-2-nitropropane		21.39	354.3	DSC	[1990DON/DRE]
C ₁₅ H ₁₃ FO ₂	[5104-49-4]		2-fluoro- α -methyl[1,1'-biphenyl]-4-acetic acid (flurbiprofen)		27.9	386.7	DAC	[1999HEN/KUH]
		$\Delta_{\text{sub}}H$	(342–367)		108.4 ± 0.5		GS	[2003PER/KUR]
C ₁₅ H ₁₃ NO	[68347-34-2]		2,10-dimethylacridin-9(10H)-one		22.4	426	DSC	[2003STO/KRZ]
		$\Delta_{\text{sub}}H$			119		DSC	[2003STO/KRZ]
C ₁₅ H ₁₃ NO	[2207-41-2]		10-ethylacridin-9(10H)-one		27.5	434		[2003STO/KRZ]
		$\Delta_{\text{sub}}H$			117		DSC	[2003STO/KRZ]
C ₁₅ H ₁₃ NO ₂	[23825-32-3]		N-benzoyl-N-methylbenzamide					
		$\Delta_{\text{sub}}H$	(246–269)		116.8 ± 0.4	356	ME	[1997ROU/JIM]
		$\Delta_{\text{sub}}H$			120.1 ± 0.4	298		[1997ROU/JIM]
C ₁₅ H ₁₃ NO ₃	[74103-06-3]		5-benzoyl-2,3-dihydro-1H-pyrrolizine-1-carboxylic acid (ketorolac)					
		$\Delta_{\text{fus}}H$ (I)			28.62	431.2	DSC	
		$\Delta_{\text{fus}}H$ (II)			171.74	430.2	DSC	
		$\Delta_{\text{fus}}H$ (III)		25.42	426.2	DSC	[2004SOH/SEO]	Note: Value for (II) seems much too large in comparison with fusion enthalpies of the other two crystalline forms
C ₁₅ H ₁₃ N ₃ O ₄ S	[36322-90-4]		2H-1,2-benzothiazine-3-carboxamide-4-hydroxy-2-methyl-N-2-pyridinyl-1,1-dioxide (piroxicam)					
		$\Delta_{\text{fus}}H$			36.3	473.4	DSC	[2006WAS/HOL, 2008WAS/HOL]
		$\Delta_{\text{fus}}H$			35	473.9	DSC	[2006DRE/SHA]
		$\Delta_{\text{fus}}H$			35	474.5		[1998GIO/GAZ]
		$\Delta_{\text{fus}}H$			34.5	473	DSC	[1998BUS/PEN]
C ₁₅ H ₁₄ CIN	[113788-74-2]		4-chlorobenzylidene-4'-ethylaniline					
		$\Delta_{\text{fus}}H$			17.21	358.4	DSC	[1999GAL/COL]
C ₁₅ H ₁₄ Cl ₂ N ₄ O ₃	[6232-56-0]		4-(N-methyl-N-2-hydroxyethylamino)-4'-nitro-2',6'-dichloroazobenzene					
		$\Delta_{\text{sub}}H$			135.1			[1968TSU/KOJ, 1988BAU/PER]
C ₁₅ H ₁₄ Cl ₃ O ₂ PS	[57875-65-7]		(chloromethyl)thiophosphonic acid, O,O-bis(2-chloro-4-methylphenyl) ester					
		$\Delta_{\text{v}}H$	(343–365)		93.2	354	A	[1987STE/MAL, 1999DYK/SVO]
C ₁₅ H ₁₄ F ₃ N ₃	[6232-56-0]		N,N-dimethyl-4-[[4-(trifluoromethyl)phenyl]azo]benzenamine					
		$\Delta_{\text{sub}}H$			95.8		UV	[1984KAR/ROD]
C ₁₅ H ₁₄ F ₃ N ₃ O	[1494-75-3]		N,N-dimethyl-4-[[4-(trifluoromethoxy)phenyl]azo]benzenamine					
		$\Delta_{\text{sub}}H$			96.8		UV	[1984KAR/ROD]
C ₁₅ H ₁₄ F ₃ N ₃ S	[1494-77-5]		N,N-dimethyl-4-[[4-(trifluoromethylthio)phenyl]azo]benzenamine					
		$\Delta_{\text{sub}}H$			100.8		UV	[1984KAR/ROD]
C ₁₅ H ₁₄ N ₂	[3295-59-8]		N,N-dimethyl-9-acridinamine					
		$\Delta_{\text{sub}}H$			86.0	510	TGA	[1998STO/KRZ]
C ₁₅ H ₁₄ N ₂	[213623-43-9]		N-methyl-10-methylacridinimine					
		$\Delta_{\text{sub}}H$			72.0	480	TGA	[1998STO/KRZ]
C ₁₅ H ₁₄ N ₂ OS	[109768-68-5]		N-(4-methoxyphenyl)-4H-3,1-benzothiazin-2-amine					
		$\Delta_{\text{fus}}H$			16.1	436.2	DSC	[2004GON/KOS]

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₅ H ₁₄ N ₂ S	[109768-67-4]	N-(4-methylphenyl)-4 <i>H</i> -3,1-benzothiazin-2-amine				
	$\Delta_{\text{fus}}H$		19.9	448.4	DSC	[2004GON/KOS]
C ₁₅ H ₁₄ O	[102-04-5]	1,3-diphenylacetone				
	$\Delta_{\text{fus}}H$		20.2	307.2		[1993ACR]
	$\Delta_{\text{sub}}H$		89.1 ± 5			[1954SPR/WHI, 1977PED/RYL, 1970COX/PIL]
	Δ_vH	(398–604)	65.7	413	A	[1987STE/MAL, 1947STU]
C ₁₅ H ₁₄ O	[10435-68-4]	4,5,6-trimethylbenzoxalene				
	$\Delta_{\text{sub}}H$		139.7 ± 2.5			[1966GEI/QUI, 1970COX/PIL]
C ₁₅ H ₁₄ O	[1210-34-0]	5 <i>H</i> -10,11-dihydrodibenzo[<i>a,d</i>]cyclohexane-5-ol				
	$\Delta_{\text{fus}}H$		19.0	365.2	DSC	[2005PER/BAN]
C ₁₅ H ₁₄ O	[2571-39-3]	3,4-dimethylbenzophenone				
	$\Delta_{\text{sub}}H$		107.9 ± 0.8	298	C	[2008GOM/AMA]
C ₁₅ H ₁₄ O ₂	[4359-34-6]	2,2-diphenyl-1,3-dioxolane				
	$\Delta_{\text{fus}}H$		15.9	328.1		[1998VER/PEN]
	$\Delta_{\text{sub}}H$		99.7 ± 1.1	298		[1998VER/PEN]
	Δ_vH	(331–370)	84.6 ± 0.6	298	GS	[2002VER]
	Δ_vH	(331–370)	81.2 ± 0.6		GS	[1998VER/PEN]
C ₁₅ H ₁₄ O ₂	[7144-65-2]or [4698-96-8]	1-biphenyloxy-2,3-epoxypropane				
	Δ_vH	(408–613)	80.0	423	A	[1987STE/MAL]
C ₁₅ H ₁₄ O ₂	[2929-45-5]	(2-hydroxy-4,6-dimethylphenyl)phenylmethanone				
	$\Delta_{\text{fus}}H$		0.67	405.2	DTA	[1989SAL/ABA]
		Note: Reported enthalpy of fusion is too small, and the published enthalpy and entropy of fusion data are internally inconsistent				
C ₁₅ H ₁₄ O ₂	[na]	4,4'-dihydroxy- α -methylstilbene				
	$\Delta_{\text{fus}}H$		20.82	465.2	DSC	[2000PUN]
		Note: DSC thermogram showed an un-quantified transition between 373 and 393 K.				
C ₁₅ H ₁₄ O ₂ S	[54897-33-5]	(Z)-1-methyl-4-(2-phenylethenyl)sulfonyl benzene				
	$\Delta_{\text{sub}}H$		116.3 ± 3.8		B	[1969MAC/MCN, 1969MAC/MCN2, 1977PED/RYL]
C ₁₅ H ₁₄ O ₂ S	[16212-08-1]	(E)-1-methyl-4-(2-phenylethenyl)sulfonyl benzene				
	$\Delta_{\text{sub}}H$		108.4 ± 2.5		B	[1969MAC/MCN, 1969MAC/MCN2, 1977PED/RYL]
C ₁₅ H ₁₄ O ₃	[15889-70-0]	2-hydroxy-4-ethoxybenzophenone				
	Δ_vH	(373–433)	90.7	403	ME	[1984SUR]
C ₁₅ H ₁₄ O ₃	[6547-53-1]	4-(phenylmethoxy)benzeneacetic acid				
	$\Delta_{\text{fus}}H$		29.3	396.1	DSC	[2006KUR/PER]
	$\Delta_{\text{sub}}H$	(378–387)	107.3 ± 3.0	383	GS	[2006KUR/PER]
C ₁₅ H ₁₄ O ₃	[3459-92-5]	dibenzyl carbonate				
	Δ_vH	(342–373)	96.7 ± 0.7	298	GS	[2008KOZ/EME]
C ₁₅ H ₁₄ O ₄	[631-38-0]	2-hydroxy-4,4'-dimethoxybenzophenone				
	$\Delta_{\text{fus}}H$		37.6	390.4	DSC	[1999PRI/HAWN]
	$\Delta_{\text{sub}}H$		121.1		B	[1999PRI/HAWN]

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
C ₁₅ H ₁₄ O ₄ S	[313057-13-5]	4-(2-propenyloxy)phenyl 5-methoxy-2-thiophene carboxylate	66.94	336.9	DSC	[2000WU/WAN]
C ₁₅ H ₁₄ O ₅	[131-54-4]	2,2'-dihydroxy-4,4'-dimethoxybenzophenone	33.2	412.3	DSC	[1999PRI/HAWN]
			130.2		B	[1999PRI/HAWN]
		(406–497)	77.4	423	A, UV	[1987STE/MAL, 1960SCH/HIR]
C ₁₅ H ₁₅ Cl	[13389-70-3]	chloro-di-4-tolylmethane				
		(406–453)	75.2	421	A	[1987STE/MAL]
C ₁₅ H ₁₅ ClN ₂ O ₂	[1982-47-4]	3-[4-[4-chlorophenoxy]phenyl]-1,1-dimethylurea	34.87	425.8	DSC	[1991ACR, 1990DON/DRE]
C ₁₅ H ₁₅ ClO ₅	[111171-33-6]	8-(hydroxymethyl)-6-chloro-5,7-dimethyl-2-oxo-2H-1-benzopyran-3-carboxylic acid, ethyl ester	3.59	446.5		
			25.08	456.4	DSC	[1992HUA/ZHO2]
C ₁₅ H ₁₅ N	[1484-09-9]	N-isopropylcarbazole				
			0.64	137.5	DSC	[1986BER/COL]
			0.38	180		
			17.73	395.2	DSC	[1991ACR, 1990KAL/DRE]
C ₁₅ H ₁₅ NO	[954-21-2]	N-methyldiphenylacetamide				
			30.23	439.8	DSC	[1990DON/DRE]
C ₁₅ H ₁₅ NO ₂	[61-68-7]	2-[(2,3-dimethylphenyl)amino]benzoic acid (mefenamic acid)				
			38.7	503.5	DSC	[2009SUR/TER]
			18.1	463.2		
			38.25	503.6	DSC	[2004ROM/BUS]
			38.2	503.6		[1999ROM/ESC]
		(357–398)	132.7 ± 0.8	377	GS	[2009SUR/TER]
		(357–398)	136.3 ± 0.8	298	GS	[2009SUR/TER]
C ₁₅ H ₁₅ NO ₃	[24033-07-6]	2-methoxy-4-[[4-methoxyphenyl]imino]methyl]phenol				
			18.53	408	DSC	[2008SIN/DAS]
C ₁₅ H ₁₅ N ₃ O ₂	[2832-40-8]	N-[4-[(2-hydroxy-5-methylphenyl)azo]phenyl]acetamide (Disperse Yellow 3)				
		(403–465)	107	434	GS	[1989NIS/AND]
			140.6			[1968TSU/KOJ, 1988BAU/PER]
C ₁₅ H ₁₅ N ₃ O ₃	[1979-02-9]	6-(acetlamino)-2-cyano-1(2H)-quinolinecarboxylic acid, ethyl ester	35.02	441.2	DSC	[2005LIZ/ZAB]
C ₁₅ H ₁₅ N ₃ O ₄	[1979-19-8]	2-cyano-6-nitro-1(2H)-quinolinecarboxylic acid, butyl ester	25.16	359.1	DSC	[2005LIZ/ZAB]
C ₁₅ H ₁₅ N ₃ O ₄	[1979-23-4]	2-cyano-6-nitro-1(2H)-quinolinecarboxylic acid, 2-methylpropyl ester	28.26	388.8	DSC	[2005LIZ/ZAB]
C ₁₅ H ₁₆	[1335-47-3]	ditolylmethane				
		(573–673)	51.8	588		[1964MAN]
C ₁₅ H ₁₆	[1530-03-6]	1,1-diphenylpropane				
		(298–343)	71.4 ± 0.4	321	GS	[1999VER5]
		(298–343)	72.8 ± 0.4	298	GS	[1999VER5]
C ₁₅ H ₁₆	[1081-75-0]	1,3-diphenylpropane				
		(342–577)	61.5	357	A	[1987STE/MAL]

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
C ₁₅ H ₁₆ ClN ₂ O ₂	[556836-79-4] $\Delta_{\text{fus}}H$	4-chloro-2'-hydroxy-4'-propoxyazobenzene	29.8	371	DSC	[2003PAJ/ROS]
C ₁₅ H ₁₆ N ₂ O ₂	[12771-68-5] $\Delta_{\text{fus}}H$	α -cyclopropyl- α -(4-methoxyphenyl)-5-pyrimidinemethanol	26.63	383.1	DSC	[1990DON/DRE]
C ₁₅ H ₁₆ N ₂ O ₂	[na] $\Delta_{\text{fus}}H$	2-cyano-1(2 <i>H</i>)-quinolinecarboxylic acid, butyl ester	22.5	347.5	DSC	[2005LIZ/ZAB]
C ₁₅ H ₁₆ N ₂ O ₂ S	[6601-00-9] $\Delta_{\text{fus}}H$	N,N'-bis(3-methoxyphenyl)thiourea	43.83	405.2	DSC	[2002ABB/WOH]
C ₁₅ H ₁₆ N ₂ O ₃	[16460-28-9] $\Delta_{\text{fus}}H$	N,N'-bis(3-methoxyphenyl)urea	36.76	443.2	DSC	[2002ABB/WOH]
C ₁₅ H ₁₆ N ₂ O ₃	[1979-07-4] $\Delta_{\text{fus}}H$	2-cyano-6-methoxy-1(2 <i>H</i>)-quinolinecarboxylic acid, propyl ester	15.6	339.9	DSC	[2005LIZ/ZAB]
C ₁₅ H ₁₆ N ₄ O ₂	[4313-14-8] $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$ $\Delta_{\text{v}}H$	3-methyl-3'-nitro-4,N,N-dimethylaminoazobenzene (368–393) (370–388) (370–388)	101.7 ± 1.7 98.7 ± 2.5 98.6	381 379 379	ME TE A	[1967GRE/JON] [1967GRE/JON] [1987STE/MAL]
C ₁₅ H ₁₆ N ₄ O ₂	[92114-99-3] $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$	3-methyl-4'-nitro-4,N,N-dimethylaminoazobenzene (369–392) (371–390)	125.5 ± 1.3 126.4 ± 3.8	381 381	TE ME	[1967GRE/JON, 1987STE/MAL] [1967GRE/JON]
C ₁₅ H ₁₆ N ₄ O ₆	[74734-24-0] $\Delta_{\text{fus}}H$	2,4-bis(2-oxo-3-oxazolidin-3-ylcarbonylamino)toluene	5.2	479.5		[1990SHI/HAY]
C ₁₅ H ₁₆ O	[885-77-8] $\Delta_{\text{v}}H$	di-(4-tolyl)methanol (413–478)	81.7	428	A	[1987STE/MAL]
C ₁₅ H ₁₆ O	[na] $\Delta_{\text{v}}H$	1-isovaleronaphthone (409–593)	76.2	424	A	[1987STE/MAL, 1947STU]
C ₁₅ H ₁₆ O	[599-64-4] $\Delta_{\text{fus}}H$ $\Delta_{\text{fus}}H$	4-(1-methyl-1-phenylmethyl)phenol (p- α -cumylphenol)	22.8 21.68	346.2 346.4	DSC	[1998JAM/PAL] [1996DOM/HEA]
C ₁₅ H ₁₆ O ₂	[80-05-7] $\Delta_{\text{fus}}H$ $\Delta_{\text{v}}H$	4,4'-dihydroxydiphenyl-2,2-propane (bisphenol A) (466–634)	30.1 102.2	433 481	A	[1996DOM/HEA] [1987STE/MAL, 1947STU]
C ₁₅ H ₁₆ O ₂	[2235-01-0] $\Delta_{\text{fus}}H$ $\Delta_{\text{sub}}H$	dimethoxydiphenylmethane	27.8 103.9 ± 1.7	380 298	DSC	[1998VER/PEN] [1998VER/PEN]
C ₁₅ H ₁₆ S ₂	[14252-46-1] $\Delta_{\text{fus}}H$	2,2-bis(phenylthio)propane	24.4	329		[1997STE/CHI]
C ₁₅ H ₁₇ BrNO ₂	[1689-99-2] $\Delta_{\text{fus}}H$	3,5-dibromo-4-hydroxybenzoxonitrile octanoyl ester	26.49	318.3	DSC	[1990DON/DRE]
C ₁₅ H ₁₇ ClN ₄	[88671-89-0] $\Delta_{\text{fus}}H$	α -butyl- α -(4-chlorophenyl)-1 <i>H</i> -1,2,4-triazole-1-propanenitrile (myclobutanil)	30.93	348.8		[2005SUN/LIU2]
C ₁₅ H ₁₇ NO ₂	[16112-55-3] $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$	N-(2-hydroxy-3-phenoxypropyl)phenylamine (323–333)	113.9 113.8 ± 2.1	328	A	[1987STE/MAL] [1976KUZ/MIR]

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(343–373)	99.9	358	A	[1987STE/MAL]
		Note: There is a problem with these experimental values				
C ₁₅ H ₁₈	[86-89-5]	1-pentyl-naphthalene				
	$\Delta_v H$	(415–535)	62.7	430	A	[1987STE/MAL]
C ₁₅ H ₁₈ Cl ₂ N ₂ O ₃	[19666-30-9]	3-[2,4-dichloro-5-(1-methylethoxy)phenyl]-5-(1,1-dimethylethyl)-1,3,4-oxadiazol-2(3H)-one				
	$\Delta_{\text{fus}} H$		26.39	360.6	DSC	[1990DON/DRE]
C ₁₅ H ₁₈ N ₂	[101-72-4]	4-isopropylaminodiphenylamine				
	$\Delta_{\text{sub}} H$	(323–348)	120.7	335	GS	[1971FEL/KUZ]
C ₁₅ H ₁₈ N ₂ O ₆	[485-31-4]	2-sec-butyl-4,6-dinitrophenyl 3-methylcrotonate				
	$\Delta_{\text{fus}} H$		18.89	341.3	DSC	[1990DON/DRE]
C ₁₅ H ₁₈ O	[20490-22-6]	2,4,6-triallylphenol				
	$\Delta_v H$	(423–571)	61.0	438	A	[1987STE/MAL]
C ₁₅ H ₁₈ O	[5737-13-3]	4H-cyclopenta[def]phenanthren-4-one				
	$\Delta_{\text{fus}} H$		16.3	443.9	DSC	[2010KES/AUC]
C ₁₅ H ₁₉ Cl ₃ O ₃	[1928-41-2]	2,4,5-trichlorophenoxyacetic acid, heptyl ester				
	$\Delta_v H$	(460–573)	92.3	475	A,GC	[1987STE/MAL, 1966JEN/SCH]
C ₁₅ H ₁₉ N ₃ O ₈	[53848-88-7]	octyl 2,4,6-trinitrobenzoate				
	$\Delta_{\text{trs}} H$		2.07	312		
	$\Delta_{\text{fus}} H$		29.16	396.7	DSC	[1974WAR/WIL]
C ₁₅ H ₂₀ Cl ₂ O ₃	[1917-96-0]	2,4-dichlorophenoxyacetic acid, heptyl ester				
	$\Delta_v H$	(460–573)	88.3	475	A,GC	[1987STE/MAL, 1966JEN/SCH]
C ₁₅ H ₂₀ Cl ₂ O ₃	[1917-94-8]	2,4-dichlorophenoxyacetic acid, 1-propylbutyl ester				
	$\Delta_v H$	(460–573)	77.3	475	A,GC	[1987STE/MAL, 1966JEN/SCH]
C ₁₅ H ₂₀ Cl ₂ O ₄	[3966-11-8]	2,4-dichlorophenoxyacetic acid, (1-methyl-2-butoxy)ethyl ester				
	$\Delta_v H$	(443–573)	82.5	458	A	[1987STE/MAL]
C ₁₅ H ₂₀ N ₂ O ₄ S	[968-81-0]	4-acetyl-N-[(cyclohexylamino)carbonyl]benzene sulfonamide				
	$\Delta_{\text{fus}} H$		41.08	457		[1982MAR/MIR]
C ₁₅ H ₂₀ N ₄ O ₄	[na]	1,1'-(1,5-pentanediyloxy)bis thymine				
	$\Delta_{\text{fus}} H$		32.03	524		[2002ITA/KAM]
C ₁₅ H ₂₀ O ₂	[1407-13-3]	helenine, alantolactone				
	$\Delta_v H$	(430–548)	112.7	445	A	[1987STE/MAL]
C ₁₅ H ₂₁ NO	[13430-30-3]	2-methyl-1-phenyl-2-N-piperidinyl-1-propanone				
	$\Delta_{\text{fus}} H$		16.74	310.2		[1994BEC/RUE]
	$\Delta_{\text{sub}} H$		94.8 ± 1.3		B	[1994WEL/VER]
C ₁₅ H ₂₁ NO ₂	[57-42-1]	1-methyl-4-phenylpiperidine-4-carboxylic acid ethyl ester (meperidine)				
	$\Delta_{\text{fus}} H$		24.6	308.2	DSC	[1988ROY/FLY]
C ₁₅ H ₂₁ NO ₄	[57837-19-1]	methyl N-(2-methoxyacetyl)-n-(2,6-xylyl)-(dl)-alaninate				
	$\Delta_{\text{fus}} H$		26.46	345.5	DSC	[1990DON/DRE]
C ₁₅ H ₂₁ N ₃ O ₃ S	[21187-98-4]	N-(4-methylbenzenesulfonyl)-N'-[3-azabicyclo(3,3,0)oct-3-yl]urea (gliclazide)				
	$\Delta_{\text{fus}} H$		44.2	444.6	DSC	[2006WAS/HOL]
C ₁₅ H ₂₂	[26460-76-4]	1-methyldiamantane				
	$\Delta_{\text{sub}} H$	(310–333)	80.7 ± 0.4	321	TSGC	[1975CLA/KNO]

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
C ₁₅ H ₂₂	[38375-86-2] $\Delta_{\text{sub}}H$	3-methyldiamantane (305–327)	103.1 ± 1.0	316	TSGC	[1975CLA/KNO]
C ₁₅ H ₂₂	[30545-18-9] $\Delta_{\text{sub}}H$	4-methyldiamantane (310–333)	79.4 ± 1.25	321	TSGC	[1975CLA/KNO]
C ₁₅ H ₂₂ ClNO ₂	[51218-45-2] $\Delta_{\text{fus}}H$	2-chloro-N-(2-ethyl-6-methylphenyl)-N-(2-methoxy-1-methylethyl)acetamide	17.0	299	DSC	[2005SBI/VEC]
	$\Delta_{\text{v}}H$		70 ± 1	436	TGA	[2007VEC]
C ₁₅ H ₂₂ N ₂ O	[24358-84-7] $\Delta_{\text{fus}}H$	N-(2,6-dimethylphenyl)-1-methyl-2-piperidinecarboxamide	17.77	426.2	DSC	[1997NEM/ACS]
C ₁₅ H ₂₂ N ₂ O ₂	[5124-30-1] $\Delta_{\text{v}}H$	dicyclohexylmethane-4,4'-diisocyanate (326–404)	80.4	341	A	[1987STE/MAL]
C ₁₅ H ₂₂ N ₂ O ₄	[92700-71-5] $\Delta_{\text{fus}}H$	octyl N-(4-nitrophenyl) carbamate	38.85	383.6	DSC	[1993TIE/FRA]
C ₁₅ H ₂₂ N ₂ O ₅	[138517-11-0] $\Delta_{\text{fus}}H$	(4-nitrophenyl)-8-hydroxyoctyl carbamate	44.07	386.9	DSC	[1993TIE/FRA]
C ₁₅ H ₂₂ O ₂	[16225-26-6] $\Delta_{\text{sub}}H$	3,5-di- <i>tert</i> -butylbenzoic acid (339–357)	108.4 ± 4.2	348	ME	[1974ROU/TUR, 1987STE/MAL, 1977PED/RYL]
C ₁₅ H ₂₂ O ₂	[37942-07-7] $\Delta_{\text{sub}}H$	3,5-di- <i>tert</i> -butyl-2-hydroxybenzaldehyde (296–312)	95.7 ± 0.5	304	ME	[2010RIB/GON]
	$\Delta_{\text{sub}}H$		96.0 ± 0.5	298	ME	[2010RIB/GON]
C ₁₅ H ₂₂ O ₂	[3575-31-3] $\Delta_{\text{sub}}H$ (I)	4-octylbenzoic acid (357–365)	134.7 ± 1.5	298	ME	[2004MON/ALM]
	$\Delta_{\text{sub}}H$ (II)		135.4 ± 1.3	298	ME	[2004MON/ALM]
C ₁₅ H ₂₂ O ₃	[79785-45-8] $\Delta_{\text{fus}}H$	3-octyloxybenzoic acid	33.12	347.1		[2001LAI/LEE]
C ₁₅ H ₂₂ O ₃	[2493-84-7] $\Delta_{\text{sub}}H$	4-octyloxybenzoic acid	163.0 ± 1.2	298		[2010RIB/FER3]
C ₁₅ H ₂₂ O ₃	[19715-19-6] $\Delta_{\text{fus}}H$	3,5-di- <i>tert</i> -butylsalicylic acid	22.92	437.5	DSC	[2003YU/TAN]
	$\Delta_{\text{sub}}H$		83.9 ± 2.6		DSC	[2003YU/TAN]
C ₁₅ H ₂₂ O ₅	[63968-64-9] $\Delta_{\text{fus}}H$ (I)	octahydro-3,6,9-trimethyl-3,12-epoxy-12 <i>H</i> -pyrano[4,3- <i>j</i>]-1,2-benzodioxepin-10(3 <i>H</i>)-one (artemisinin)	22.8	428.2		
	$\Delta_{\text{fus}}H$ (II)		23.41	428.1	DSC	[1997CHA/YUE]
C ₁₅ H ₂₃ NO ₂	[na] $\Delta_{\text{fus}}H$	(+) 1-(<i>o</i> -allylphenoxy)-3-(isopropylamino)-2-propanol (alprenolol)	23.78	298.5	DSC	[1999LI/ZEL]
C ₁₅ H ₂₃ NO ₂	[13655-52-2] $\Delta_{\text{fus}}H$	(±) 1-(<i>o</i> -allylphenoxy)-3-(isopropylamino)-2-propanol (alprenolol)	35.61	331.2	DSC	[1999LI/ZEL]
C ₁₅ H ₂₃ N ₃ O ₂	[135742-55-1] $\Delta_{\text{fus}}H$	N-capryl-pyrazinamide	50.58	360.5		[1991LIU/GUO]
C ₁₅ H ₂₃ N ₃ O ₄ S	[na] $\Delta_{\text{fus}}H$	(–) N-1-(ethylpyrrolidin-2-ylmethyl)-2-methoxy-5-sulfamoylbenzamide (sulpiride)	42.01	459.5		[1999LI/ZEL]
C ₁₅ H ₂₃ N ₃ O ₄ S	[15676-16-1]	(+) N-1-(ethylpyrrolidin-2-ylmethyl)-2-methoxy-5-sulfamoylbenzamide (sulpiride)				

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		T _m (K)	Method	Reference
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)			
		$\Delta_{\text{fus}}H$		46.15	451	[1999LI/ZEL]
C ₁₅ H ₂₄	[1081-77-2]	nonylbenzene				
		$\Delta_{\text{v}}H$	(304–466)	74.1 ± 0.5	298	MM [1998MOK/RAU, 2006VER/KOZ]
		$\Delta_{\text{v}}H$	(316–415)	69.7	331	GS [1986ALL/JOS]
		$\Delta_{\text{v}}H$		74.8	298	[1971WIL/ZWO]
C ₁₅ H ₂₄	[717-74-8]	1,3,5-triisopropylbenzene				
		$\Delta_{\text{v}}H$	(283–323)	64.3 ± 0.3	303	GS [1998VER7]
		$\Delta_{\text{v}}H$	(283–323)	64.6 ± 0.6	298	GS [1998VER7]
		$\Delta_{\text{v}}H$	(282–388)	67.4	297	[1993KAS/MOK]
C ₁₅ H ₂₄	[na]	1,3-di- <i>tert</i> -butyl-5-methylbenzene				
		$\Delta_{\text{sub}}H$	(275–301)	82.4 ± 0.5	288	T [1998VER]
		$\Delta_{\text{sub}}H$	(275–301)	81.8 ± 0.5	298	T [1998VER]
		$\Delta_{\text{v}}H$	(309–338)	61.8 ± 0.9	310	GS [1998VER]
		$\Delta_{\text{v}}H$	(309–338)	63.3 ± 0.9	298	GS [1998VER]
C ₁₅ H ₂₄	[18794-84-8]	(E)- β -farnesene				
		$\Delta_{\text{v}}H$	(363–473)	72.5	298	GC [2005HOS/GRY]
C ₁₅ H ₂₄	[87-44-5]	β -caryophyllene				
		$\Delta_{\text{v}}H$	(363–463)	65.5	298	GC [2005HOS/GRY]
C ₁₅ H ₂₄ N ₂ O ₃	[490-98-2]	4-(butylamino)-2-hydroxybenzoic acid, 2-(dimethylamino)ethyl ester (salicaine)				
		$\Delta_{\text{fus}}H$		26.8	319.4	DSC [2006SCH/SCH]
C ₁₅ H ₂₄ O	[497-39-2]	2,4-di- <i>tert</i> -butyl-5-methylphenol				
		$\Delta_{\text{v}}H$	(376–555)	67.0	391	A [1987STE/MAL, 1947STU]
C ₁₅ H ₂₄ O	[616-55-7]	2,4-di- <i>tert</i> -butyl-6-methylphenol				
		$\Delta_{\text{v}}H$	(359–543)	59.8	374	A [1987STE/MAL]
C ₁₅ H ₂₄ O	[128-37-0]	2,6-di- <i>tert</i> -butyl-4-methylphenol				
		$\Delta_{\text{fus}}H$		19.85	341.7	DSC [1999VER]
		$\Delta_{\text{fus}}H$		23.85	343.7	DTA [1972INO/LIA]
		$\Delta_{\text{sub}}H$		91.9 ± 3.2	298	C [2001RIB/MAT]
		$\Delta_{\text{sub}}H$	(298–338)	86.8 ± 0.8	319	GS [1999VER]
		$\Delta_{\text{sub}}H$	(298–338)	88.0 ± 0.8	298	GS [1999VER]
		$\Delta_{\text{sub}}H$	(303–343)	87.8	318	GS [1987STE/MAL, 1971FEL/KUZ]
		$\Delta_{\text{sub}}H$		U 117.3	298	C [1971BER/GIR, 1999VER]
		$\Delta_{\text{v}}H$	(303–343)	87.8	318	A [1987STE/MAL]
		$\Delta_{\text{v}}H$	(358–536)	61.5	373	A [1987STE/MAL, 1947STU]
	C ₁₅ H ₂₄ O	[2219-84-3]	2-methyl-4-(1,1,3,3-tetramethylbutyl)phenol			
		$\Delta_{\text{v}}H$	(447–683)	67.1	462	A [1987STE/MAL]
C ₁₅ H ₂₄ O	[2219-84-3]	3-methyl-4-(1,1,3,3-tetramethylbutyl)phenol				
		$\Delta_{\text{v}}H$	(436–549)	65.5	451	A [1987STE/MAL]
C ₁₅ H ₂₄ O	[4979-46-8]	4-methyl-2-(1,1,3,3-tetramethylbutyl)phenol				
		$\Delta_{\text{v}}H$	(415–545)	65.0	430	A [1987STE/MAL]
C ₁₅ H ₂₄ O	[na]	4-(3',6'-dimethyl-3'-heptyl)phenol				
		$\Delta_{\text{v}}H$		89.4	298	ME [2001LAL/SCH]
C ₁₅ H ₂₄ O	[104-40-5]	4-nonylphenol				
		$\Delta_{\text{v}}H$	(487–595)	65.0	502	A, EB [1987STE/MAL, 1976HON/SIN]

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₅ H ₂₄ O	[115-71-9] $\Delta_v H$	α -santalol (293–450)	58.3	308	A	[1987STE/MAL]
C ₁₅ H ₂₄ O ₂	[1991-52-2] $\Delta_{\text{fus}} H$ $\Delta_v H$	2,5-di- <i>tert</i> -butyl-4-methoxyphenol (423–453)	26.9 64.4	374.4 438	A	[1972ALV/BOR] [1987STE/MAL]
C ₁₅ H ₂₄ O ₂	[6121-64-8] $\Delta_v H$	1,3-dimethoxy-5-heptylbenzene (419–488)	75.5	434	A, GC	[1987STE/MAL, 1975KUN/LIL]
C ₁₅ H ₂₄ O ₂	[41442-51-7] $\Delta_v H$	1,3-dimethoxy-5-methyl-2-hexylbenzene (410–475)	72.3	425	A, GC	[1987STE/MAL, 1975KUN/LIL]
C ₁₅ H ₂₄ O ₄	[1152-57-4] $\Delta_v H$	dicyclohexyl malonate (324–353)	93.7 ± 1.1	298	GS	[2008LIP/KRA]
C ₁₅ H ₂₄ O ₆	[64617-28-3] $\Delta_v H$	aconitic acid, tripropyl ester (359–500)	72.3	374	A	[1987STE/MAL]
C ₁₅ H ₂₆ O	[489-86-1] $\Delta_v H$	guaiol (373–561)	62.2	388	A	[1987STE/MAL]
C ₁₅ H ₂₆ O ₆	[na] $\Delta_v H$	camphorenic acid, triethyl ester (423–574)	69.0	438	A	[1987STE/MAL, 1947STU]
C ₁₅ H ₂₆ O ₆	[5333-54-0] $\Delta_v H$	tripropyl 1,2,3-propanetricarboxylate (360–460)	76.5	375	A	[1987STE/MAL]
C ₁₅ H ₂₆ O ₆	[60-01-5] $\Delta_v H$	glycerol tributyrate (318–364)	81.4	333	A	[1987STE/MAL]
C ₁₅ H ₂₆ O ₆	[60-01-5] $\Delta_v H$ $\Delta_v H$ $\Delta_v H$	tributyryn	83.5 84.9 ± 2.5 107.1 ± 1.0	308 298 298	TGA TGA C	[1990KIS/SHO] [1990KIS/SHO] [1986NIL/WAD]
C ₁₅ H ₂₈ Cl ₄	[3922-32-5] $\Delta_v H$	1,1,1,15-tetrachloropentadecane (340–392)	103.5	355	A	[1987STE/MAL]
C ₁₅ H ₂₈ O	[1604-35-9] $\Delta_v H$	3,7,11-trimethyl-1-dodecyn-3-ol (401–524)	43.2 ± 1.1	463		[1988BAG/GUR, 1986WHI]
C ₁₅ H ₂₈ O	[502-72-7] $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$	cyclopentadecanone (296–315)	86.0 ± 0.6 77.4	305	ME	[1997JIM/ROU] [1938WOL/WEG, 1960JON, 1970COX/PIL]
C ₁₅ H ₂₈ O ₂	[2156-97-0] $\Delta_v H$	dodecyl acrylate (432–573)	64.6	447	A	[1987STE/MAL]
C ₁₅ H ₂₈ O ₂	[106-02-5] $\Delta_{\text{sub}} H$ $\Delta_v H$ $\Delta_v H$	pentadecanolide (290–310) (363–443) (310–320)	81.3 78.2 74.2	300 378 315	ME A A, ME	[1987STE/MAL, 1960JON, 1954SER/VOI] [1987STE/MAL] [1987STE/MAL, 1954SER/VOI]
C ₁₅ H ₂₈ O ₂	[34270-22-9] $\Delta_v H$	(Z) 7-tridecenyl acetate (343–388)	84.3	298	GC	[1997KOU/HOS, 2000OVA/KOU]
C ₁₅ H ₂₈ O ₂	[56577-30-1] $\Delta_v H$	(E) 7-tridecenyl acetate (343–388)	84.8	298	GC	[1997KOU/HOS, 2000OVA/KOU]
C ₁₅ H ₂₈ O ₂	[35835-78-0]	(Z) 9-tridecenyl acetate				

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		T _m (K)	Method	Reference	
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)				
		$\Delta_{\text{v}}H$	(343–388)	85.1	298	GC	[1997KOU/HOS, 2000OVA/KOU]
C ₁₅ H ₂₈ O ₂	[52957-19-4]	(E) 9-tridecenyl acetate					
	$\Delta_{\text{v}}H$	(343–388)	85.5	298	GC	[1997KOU/HOS, 2000OVA/KOU]	
C ₁₅ H ₂₈ O ₂	[33951-95-0]	(Z) 11-tridecenyl acetate					
	$\Delta_{\text{v}}H$	(343–388)	86.4	298	GC	[1997KOU/HOS, 2000OVA/KOU]	
C ₁₅ H ₂₈ O ₂	[56195-36-9]	(E) 11-tridecenyl acetate					
	$\Delta_{\text{v}}H$	(343–388)	86.4	298	GC	[1997KOU/HOS, 2000OVA/KOU]	
C ₁₅ H ₂₈ O ₂	[56219-06-8]	methyl Z 9-tetradecenoate					
	$\Delta_{\text{v}}H$		87.1 ± 0.7	298	CGC	[2007LIP/KAP]	
C ₁₅ H ₂₈ O ₂	[106-02-5]	pentadecanolactone					
	$\Delta_{\text{trs}}H$		27.3	283			
	$\Delta_{\text{fus}}H$	(10–370)	6.99	308.5	AC	[1984DOM/EVA, 1981LEB/YEV]	
C ₁₅ H ₂₈ O ₃	[37826-51-0]	decyl levulinate					
	$\Delta_{\text{v}}H$	(423–580)	76.1	438	A	[1987STE/MAL]	
	$\Delta_{\text{v}}H$		72.0	524		[1933COW/SCH]	
C ₁₅ H ₂₈ O ₃	[6707-60-4]	1,6-dioxa-7-cycloheptadecanone					
	$\Delta_{\text{v}}H$	(403–463)	75.9	418	A	[1987STE/MAL]	
C ₁₅ H ₂₈ O ₅	[1085702-05-1]	decyl[1-(methoxycarbonyl)ethyl]carbonate					
	$\Delta_{\text{v}}H$	(411–592)	73.8	426	A	[1987STE/MAL]	
C ₁₅ H ₂₉ N	[2570-26-5]	pentadecanenitrile					
	$\Delta_{\text{v}}H$	(336–372)	88.1 ± 0.3	298	GS	[2005EME/VER]	
	$\Delta_{\text{v}}H$	(403–596)	75.5	418	A	[1987STE/MAL]	
C ₁₅ H ₂₉ NO ₃	[na]	2-[2-ethyl(hexanoyloxy)]propionic acid, butylamide					
	$\Delta_{\text{v}}H$	(378–433)	81.0	393	A	[1987STE/MAL]	
C ₁₅ H ₂₉ NO ₃	[22220-07-1]	N-decanoyl-(<i>l</i>)-valine					
	$\Delta_{\text{trs}}H$		21.3	378.1			
	$\Delta_{\text{fus}}H$		15.4	380.6	DSC	[1986MIY/MAT]	
C ₁₅ H ₂₉ NO ₃	[83871-16-3]	N-decanoyl-(<i>dl</i>)-valine					
	$\Delta_{\text{fus}}H$		63.1	358.1	DSC	[1986MIY/MAT]	
C ₁₅ H ₂₉ NO ₃	[na]	N-dodecanoyl-(<i>l</i>)-alanine					
	$\Delta_{\text{fus}}H$		37.6	356.1	DSC	[1986MIY/MAT]	
C ₁₅ H ₃₀	[1795-21-7]	decylcyclopentane					
	$\Delta_{\text{fus}}H$		33.14	251		[1996DOM/HEA]	
	$\Delta_{\text{v}}H$	(358–411)	71.1	373	A	[1987STE/MAL]	
	$\Delta_{\text{v}}H$		75.7	298		[1971WIL/ZWO]	
	$\Delta_{\text{v}}H$	(453–553)	59.7	468	A, MM	[1987STE/MAL, 1954CAM/FOR]	
C ₁₅ H ₃₀	[2883-02-5]	nonylcyclohexane					
	$\Delta_{\text{v}}H$		74.7	298		[1971WIL/ZWO]	
C ₁₅ H ₃₀	[13360-61-7]	1-pentadecene					
	$\Delta_{\text{v}}H$	(375–407)	65.2	390	A	[1987STE/MAL]	
	$\Delta_{\text{v}}H$	(423–658)	53.2	570		[1975AMB/ELL]	
	$\Delta_{\text{v}}H$		75.1	298		[1971WIL/ZWO]	
		(443–543)	59.3	458	A	[1987STE/MAL, 1955CAM/ROS]	
C ₁₅ H ₃₀	[295-48-7]	cyclopentadecane					

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
	Enthalpy							
		$\Delta_{\text{trs}}H$			8.5	210.1		
		$\Delta_{\text{fus}}H$			8.5	336.6		[1987DRO/MOL]
		$\Delta_{\text{sub}}H$			74.6 ± 0.4			[1957VAN, 1970COX/PIL]
C ₁₅ H ₃₀ N ₃ PS ₆	[17767-20-3]		phosphorus- <i>tris</i> (N,N-diethyldithiocarbamate)		143 ± 2	298		[1987AIR/DES]
C ₁₅ H ₃₀ O	[56218-94-1]		(Z) 9-pentadecen-1-ol	(363–403)	105.3	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₅ H ₃₀ O	[64437-40-7]		(E) 9-pentadecen-1-ol	(363–403)	105.9	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₅ H ₃₀ O	[64437-42-9]		(Z) 10-pentadecen-1-ol	(363–403)	105.9	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₅ H ₃₀ O	[64437-44-1]		(E) 10-pentadecen-1-ol	(363–403)	106.2	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₅ H ₃₀ O	[69282-63-9]		(Z) 11-pentadecen-1-ol	(363–403)	106.3	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₅ H ₃₀ O	[69222-14-6]		(E) 11-pentadecen-1-ol	(363–403)	106.5	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₅ H ₃₀ O	[158906-50-4]		(Z) 12-pentadecen-1-ol	(363–403)	106.7	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₅ H ₃₀ O	[69222-15-7]		(E) 12-pentadecen-1-ol	(363–403)	107	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₅ H ₃₀ O	[158906-51-5]		(Z) 13-pentadecen-1-ol	(363–403)	107.7	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₅ H ₃₀ O	[158906-52-6]		(E) 13-pentadecen-1-ol	(363–403)	107.7	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₅ H ₃₀ O	[2345-28-0]		2-pentadecanone		54.39	312.2		[1996DOM/HEA]
		$\Delta_{\text{sub}}H$			139.3 ± 1.6	298	C	[1979SUN/SVE2]
		$\Delta_{\text{v}}H$		(422–575)	67.8	437	A	[1987STE/MAL]
		$\Delta_{\text{v}}H$		(559–658)	57.9	574	A	[1987STE/MAL]
		$\Delta_{\text{v}}H$			85.4 ± 1.7	298	S-F	[1979SUN/SVE2]
C ₁₅ H ₃₀ O	[818-23-5]		8-pentadecanone		65.3	458	A	[1987STE/MAL]
		$\Delta_{\text{v}}H$		(443–568)	65.4	458	A	[1987STE/MAL, 1975AMB/ELL]
		$\Delta_{\text{v}}H$		(444–590)	53.0	567		[1975AMB/ELL]
		$\Delta_{\text{v}}H$		(438–462)	61.9	450	A, ME	[1987STE/MAL, 1938UBB]
C ₁₅ H ₃₀ O ₂	[124-10-7]		methyl tetradecanoate (methyl myristate)		50.21	291.6		[1993ACR]
		$\Delta_{\text{sub}}H$			137.7 ± 2.1	281	ME	[1965DAV/KYB]
		$\Delta_{\text{v}}H$			79.8	350		[2002VAN/VAN]
		$\Delta_{\text{v}}H$			76.0 ± 0.2	382		[2002VAN/VAN]
		$\Delta_{\text{v}}H$			85.9 ± 0.8	298		[2002VAN/VAN]
		$\Delta_{\text{v}}H$		(393–473)	86.6	298	GC	[1997KRO/VEL]

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		T _m (K)	Method	Reference	
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)				
		$\Delta_{\text{v}}H$	(453–543)	65.3	498	GC	[1993HUS/SAR]
		$\Delta_{\text{v}}H$		86.2 ± 1.0	298	GC, C	[1980FUC/PEA]
		$\Delta_{\text{v}}H$		87.0 ± 0.9	298	C	[1977MAN/SEL]
		$\Delta_{\text{v}}H$	(389–519)	75.6	404	A	[1987STE/MAL, 1963ROS/SCH]
		$\Delta_{\text{v}}H$	(364–417)	77.4	379	MG, OM	[1952SCO/MAC]
C ₁₅ H ₃₀ O ₂	[28267-29-0]	ethyl tridecanoate					
		$\Delta_{\text{fus}}H$		40.7	272.4	AC	[2005VAN/OON]
C ₁₅ H ₃₀ O ₂	[10233-13-3]	isopropyl dodecanoate					
		$\Delta_{\text{v}}H$	(305–452)	81.5	320		[2001BUR/JOS]
		$\Delta_{\text{v}}H$	(390–469)	66.1	405	A	[1987STE/MAL, 1948BON/ATH, 1984BOU/FRI]
C ₁₅ H ₃₀ O ₂	[3681-78-5]	propyl dodecanoate					
		$\Delta_{\text{v}}H$	(423–483)	84.7	298	GC	[1997KRO/VEL]
		$\Delta_{\text{v}}H$	(396–479)	66.9	411	A	[1987STE/MAL, 1948BON/ATH, 1984BOU/FRI]
C ₁₅ H ₃₀ O ₂	[1072-33-9]	tridecyl acetate					
		$\Delta_{\text{v}}H$	(313–358)	87.2	298	GC	[1997KOU/HOS, 2000OVA/KOU]
C ₁₅ H ₃₀ O ₂	[245658-47-3]	3,3-dimethylbutanoic acid, 1,1,5-trimethylhexyl ester					
		$\Delta_{\text{v}}H$	(333–378)	67.4	298	CGC	[1999VER/HEI]
C ₁₅ H ₃₀ O ₂	[1002-84-2]	pentadecanoic acid					
		$\Delta_{\text{trs}}H$		8.2	321.9		
		$\Delta_{\text{fus}}H$		40.4	325.5	DSC	[2007GBA/NEG, 2008GBA/NEG, 2009GBA/NEG]
		$\Delta_{\text{trs}}H$		8.12	318.7		
		$\Delta_{\text{fus}}H$		41.52	325.7		[1996DOM/HEA]
		$\Delta_{\text{fus}}H$		46.1	324.9		[1976BER/BER]
		$\Delta_{\text{sub}}H$	(275–293)	144.3		TPTD	[2005CHA/ZIE]
		$\Delta_{\text{sub}}H$	(283–305)	178		TPTD	[2001CHA/TOB]
		Note: Experimental values based on the TPTD method are often inconsistent with values determined using other experimental methods					
		$\Delta_{\text{v}}H$	(431–613)	94	446	A	[1987STE/MAL]
		$\Delta_{\text{v}}H$	(347–367)	108.5 ± 2.0	357	ME, TE	[1982DEK/SCH]
C ₁₅ H ₃₀ O ₃	[4617-33-8]	15-hydroxypentadecanoic acid					
		$\Delta_{\text{sub}}H$	(294–316)	103		TPTD	[2005CHA/ZIE]
		Note: Experimental values based on the TPTD method are often inconsistent with values determined using other experimental methods					
C ₁₅ H ₃₀ O ₃	[6283-92-7]	dodecyl lactate					
		$\Delta_{\text{v}}H$	(367–583)	80.5	382	A	[1987STE/MAL]
C ₁₅ H ₃₀ O ₃	[70160-09-7]	decyl 2-ethoxypropionate					
		$\Delta_{\text{v}}H$	(423–523)	69.8	438	A	[1987STE/MAL]
C ₁₅ H ₃₀ O ₆	[63364-38-5]	3,3,6,6,9,9-tetraethyl-1,2,4,5,7,8-hexaoxacyclonane					
		$\Delta_{\text{v}}H$	(403–473)	63.6	298	CGC	[2007CAN/EYL]
C ₁₅ H ₃₁ Br	[629-72-1]	1-bromopentadecane					
		$\Delta_{\text{v}}H$	(450–661)	69.5	465	A, E	[1987STE/MAL, 1961LI/ROS, 1970DYK/VAN]
C ₁₅ H ₃₁ Cl	[4862-03-7]	1-chloropentadecane					
		$\Delta_{\text{v}}H$		92.6	298		[2006BOL/NER2]
		$\Delta_{\text{v}}H$	(439–645)	55.4	454	A, E	[1987STE/MAL, 1961LI/ROS, 1970DYK/VAN]
C ₁₅ H ₃₁ F	[1555-17-5]	1-fluoropentadecane					

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		T _m (K)	Method	Reference
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)			
C ₁₅ H ₃₁ I		(413–593)	63.8	428	A, E	[1987STE/MAL, 1961LI/ROS, 1970DYK/VAN]
	[35599-78-1]	1-iodopentadecane				
	$\Delta_{\text{v}}H$	(464–673)	94.6	298	A,E	[1987STE/MAL, 1961LI/ROS, 1970DYK/VAN, 2006BOL/NER]
C ₁₅ H ₃₁ NO		(464–673)	70.6	479	A, E	[1987STE/MAL, 1961LI/ROS, 1970DYK/VAN]
	[7438-09-7]	N-methyl tetradecanamide				
C ₁₅ H ₃₁ NO ₂		(332–347)	130.4 ± 0.8	340	GS	[1959DAV/JON, 1987STE/MAL]
	[5468-40-6]	N,N-dihexyl lactamide				
C ₁₅ H ₃₁ NO ₂		(418–453)	79.4	433	A	[1987STE/MAL]
	[5422-41-3]	N-dodecyl lactamide				
C ₁₅ H ₃₂		(408–476)	103.9	423	A	[1987STE/MAL]
	[629-62-9]	pentadecane				
	$\Delta_{\text{trs}}H$		8.7	270.3		
	$\Delta_{\text{fus}}H$		34.2	282.7	DSC	[2004MON/RAJ]
	$\Delta_{\text{trs}}H$		9.17	270.9		
	$\Delta_{\text{fus}}H$		34.6	283.1		[1996DOM/HEA]
	$\Delta_{\text{sub}}H$		107.8	298	B	[1972MOR3]
	$\Delta_{\text{v}}H$		72.9	334	C	[1996VIT/CHA]
	$\Delta_{\text{v}}H$		71.8	344	C	[1996VIT/CHA]
	$\Delta_{\text{v}}H$	(453–503)	75.7	298	CGC	[1995CHI/HOS]
	$\Delta_{\text{v}}H$	(423–473)	76.2	298	CGC	[1995CHI/HOS]
	$\Delta_{\text{v}}H$	(363–413)	76.4	298	CGC	[1995CHI/HOS]
	$\Delta_{\text{v}}H$		76.8	298		[1994RUZ/MAJ]
	$\Delta_{\text{v}}H$	(366–409)	67.5	381	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$	(333–409)	66.4	350	GS	[1986ALL/JOS]
	$\Delta_{\text{v}}H$		75.4 ± 1.2	298	C	[1979SUN/SVE]
	$\Delta_{\text{v}}H$		70.8	353	C	[1979SUN/SVE]
	$\Delta_{\text{v}}H$		68.8	373	C	[1979SUN/SVE]
	$\Delta_{\text{v}}H$		72.2 ± 1.2	333	C	[1979SUN/SVE]
	$\Delta_{\text{v}}H$		76.2 ± 0.4	298	C	[1972MOR2]
$\Delta_{\text{v}}H$		76.2	298		[1971WIL/ZWO]	
$\Delta_{\text{v}}H$	(447–546)	59.6	462	A	[1987STE/MAL, 1955CAM/ROS]	
$\Delta_{\text{v}}H$	(430–464)	61.9	447	ME	[1938UBB]	
C ₁₅ H ₃₂	[1560-95-8]	2-methyltetradecane				
	$\Delta_{\text{v}}H$	(402–537)	58.8	417	A	[1987STE/MAL]
C ₁₅ H ₃₂	[18435-22-8]	3-methyltetradecane				
	$\Delta_{\text{v}}H$	(403–538)	58.4	418	A	[1987STE/MAL]
C ₁₅ H ₃₂	[25117-24-2]	4-methyltetradecane				
	$\Delta_{\text{v}}H$	(398–536)	55.9	413	A	[1987STE/MAL]
C ₁₅ H ₃₂	[25117-32-2]	5-methyltetradecane				
	$\Delta_{\text{v}}H$	(398–535)	56.1	413	A	[1987STE/MAL]
C ₁₅ H ₃₂	[18435-20-6]	2,3-dimethyltridecane				
	$\Delta_{\text{v}}H$	(399–537)	56.3	414	A	[1987STE/MAL]
C ₁₅ H ₃₂	[61868-05-1]	2,4-dimethyltridecane				
	$\Delta_{\text{v}}H$	(393–523)	57.9	408	A	[1987STE/MAL]
C ₁₅ H ₃₂	[na]	2,4,6-trimethyldodecane				
	$\Delta_{\text{v}}H$	(382–508)	55.8	397	A	[1987STE/MAL]

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference	
	Enthalpy								
C ₁₅ H ₃₂ N ₂ O	[32954-73-7]	1-tetradecyl urea							
	$\Delta_{\text{trs}}H$			1.0	227.1				
	$\Delta_{\text{trs}}H$			1.7	369.2				
	$\Delta_{\text{fus}}H$			50.9	387.4	DSC	[2005HAS/TAJ]		
C ₁₅ H ₃₂ O	[629-76-5]	1-pentadecanol							
	$\Delta_{\text{fus}}H$			29.6	316.4	DSC	[2004VEN/CAL]		
	$\Delta_{\text{fus}}H$			53.62	316.9		[2003VAN/VAN]		
	Note: The value of 53.62 includes both the enthalpy of fusion as well as the enthalpy of solid-to-solid transition that occurs at 315.4 K								
	$\Delta_{\text{trs}}H$			23.64	316				
	$\Delta_{\text{fus}}H$			54.73	316.6		[1974MOS/MOU]		
	Δ_vH			103.5 ± 3.3	298	CGC	[2006NIC/KWE]		
	Δ_vH		(319–358)	95.5	339	GS	[2001KUL/VER2]		
	Δ_vH		(319–358)	102.5	298	GS	[2001KUL/VER2]		
	Δ_vH		(353–393)	107.2	298	CGC	[1994KOU/HOS, 2000OVA/KOU]		
	Δ_vH		(343–393)	92.4	368		[1992NGU/KAS]		
	Δ_vH		(438–600)	75.0	453	A	[1987STE/MAL]		
Δ_vH		(453–584)	72.4	468	A	[1987STE/MAL]			
C ₁₅ H ₃₂ O ₂	[14722-40-8]	1,15-pentadecanediol							
	$\Delta_{\text{trs}}H$			35.1	349.4				
	$\Delta_{\text{fus}}H$			23.6	361.4	DSC	[1999OGA/NAK]		
C ₁₅ H ₃₂ O ₂ S	[18023-86-4]	3-(dodecylthio)-1,2-propanediol							
	$\Delta_{\text{trs}}H$			18.1	299				
	$\Delta_{\text{fus}}H$			20.3	325.5	DSC	[1993ACR]		
C ₁₅ H ₃₂ O ₃	[1561-07-5]	3-(dodecyloxy)-1,2-propanediol							
	$\Delta_{\text{fus}}H$			51.4	323	DSC	[1993ACR]		
C ₁₅ H ₃₂ O ₄	[4161-34-6]	5,5'-[1,5-pentanediy]bis(oxy)]bis-1-pentanol							
	$\Delta_{\text{fus}}H$			35.66	302.7	DSC	[1991BED/BOO]		
C ₁₅ H ₃₂ O ₅	[na]	tetrapropylene glycol monoisopropyl ether							
	Δ_vH		(389–566)	71.5	404	A	[1987STE/MAL, 1947STU]		
C ₁₅ H ₃₂ O ₅ S ₂	[na]	(l)-arabinose dipentyl dithioacetal							
	$\Delta_{\text{fus}}H$			37.3	368	DSC	[1989VAN/VAN]		
C ₁₅ H ₃₂ S	[25276-70-4]	1-pentadecanethiol							
	Δ_vH		(459–629)	69.8	474		[1999DYK/SVO]		
C ₁₅ H ₃₃ N	[2570-26-5]	1-aminopentadecane							
	Δ_vH		(400–594)	71.2	415	A, E	[1987STE/MAL, 1956MAN2]		
C ₁₅ H ₃₃ NO ₂	[821-91-0]	3-(dodecylamino)-1,2-propanediol							
	$\Delta_{\text{fus}}H$			62.1	351.9	DSC	[1993ACR]		
C ₁₅ H ₃₃ O ₄ P	[2528-38-3]	tripentyl phosphate							
	Δ_vH		(443–473)	92.3	298	CGC	[2007PAN/ANT2]		
	Δ_vH		(443–483)	90.7	298	CGC	[2007PAN/ANT2]		
C ₁₅ H ₃₃ O ₄ P	[919-62-0]	triisopentyl phosphate							
	Δ_vH		(453–493)	86.6	298	CGC	[2007PAN/ANT2]		
	Δ_vH		(483–513)	86.5	298	CGC	[2007PAN/ANT2]		
C ₁₅ H ₃₃ O ₄ P	[646621-37-1]	tri-sec-pentyl phosphate							
	Δ_vH		(463–493)	80.7	298	CGC	[2007PAN/ANT2]		
	Δ_vH		(453–493)	81.5	298	CGC	[2007PAN/ANT2]		

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
C ₁₅ H ₃₃ O ₄ P	[45241-53-0]	tri-(2-methylbutyl) phosphate				
	$\Delta_{\text{v}}H$	(463–493)	86.1	298	CGC	[2007PAN/ANT2]
	$\Delta_{\text{v}}H$	(493–523)	86.7	298	CGC	[2007PAN/ANT2]
C ₁₆ F ₃₄	[355-49-7]	<i>n</i> -perfluorohexadecane				
	$\Delta_{\text{trs}}H$		1.13	176.5		
	$\Delta_{\text{trs}}H$		3.01	177.7		
	$\Delta_{\text{trs}}H$		1.89	186.7		
	$\Delta_{\text{fus}}H$		61.09	402.2	DSC	[1986STA]
	$\Delta_{\text{sub}}H$	(288–303)	104.6	295	ME	[1951BRA/WAG, 1987STE/MAL]
C ₁₆ H ₆ Br ₄ N ₂ O ₂	[2475-31-2]	5,7-dibromo-2-(5,7-dibromo-1,3-dihydro-3-oxo-2 <i>H</i> -indol-2-ylidene)-1,2-dihydro-3 <i>H</i> -indol-3-one (C.I. Vat Blue 5)				
	$\Delta_{\text{sub}}H$	(519–634)	129	577	GS	[1986NIS/AND]
C ₁₆ H ₉ Br	[1714-29-0]	1-bromopyrene				
	$\Delta_{\text{sub}}H$	(321–368)	99.2 ± 4.4		ME	[2008GOL/SUU2]
C ₁₆ H ₉ BrN ₂ O ₂	[6492-73-5]	5-bromo-2-(1,3-dihydro-3-oxo-2 <i>H</i> -indol-2-ylidene)-1,2-dihydro-3 <i>H</i> -indol-3-one (C.I. Vat Blue 3)				
	$\Delta_{\text{sub}}H$	(519–634)	57.0	577	GS	[1986NIS/AND]
C ₁₆ H ₉ F ₂₅	[89109-69-3]	1,1,1,2,2,3,3,4,4,5,5,6,6,7,7,8,8,9,9,10,10,11,11,12,12-pentacosofluorohexadecane				
	$\Delta_{\text{trs}}H$		0.7	147		
	$\Delta_{\text{trs}}H$		1.4	314		
	$\Delta_{\text{fus}}H$		21.0	349	DSC	[1991HOP/MOL]
	$\Delta_{\text{trs}}H$		1.4	312.2		
	$\Delta_{\text{fus}}H$		20.9	349.2	DSC	[1986RUS/RAB]
C ₁₆ H ₉ NO ₂	[5522-43-0]	1-nitropyrene				
	$\Delta_{\text{fus}}H$		18.9	425.9	DSC	[2010KES/AUC]
	$\Delta_{\text{sub}}H$	(379–408)	125.2 ± 3.8		ME	[2008GOL/SUU]
C ₁₆ H ₉ NO ₂	[892-21-7]	3-nitrofluoranthene				
	$\Delta_{\text{fus}}H$		22.6	435.0	DSC	[2010KES/AUC]
C ₁₆ H ₁₀	[206-44-0]	fluoranthene				
	$\Delta_{\text{fus}}H$		18.74	383.4		[1996DOM/HEA]
	$\Delta_{\text{sub}}H$	(313–453)	98.3	383	GS	[1995NAS/LEN]
	$\Delta_{\text{sub}}H$	(283–323)	84.6 ± 0.9	303	GS	[1983SON/ZOL]
	$\Delta_{\text{sub}}H$		99.2 ± 0.8	298	C	[1972MOR, 1977PED/RYL]
	$\Delta_{\text{sub}}H$	(328–353)	102.1 ± 2	340	ME	[1965BOY/CHR, 1970COX/PIL]
	$\Delta_{\text{sub}}H$	(298–358)	102.6	328		[1958HOY/PEP]
	$\Delta_{\text{v}}H$	(423–493)	86.8 ± 1.3	298	GC	[2005RIB/GOM]
	$\Delta_{\text{v}}H$	(323–473)	79.3	398	GC	[2002LEI/CHA]
	$\Delta_{\text{v}}H$	(343–453)	77.4	398	GC	[1990HIN/BID2]
	$\Delta_{\text{v}}H$	(503–658)	62.2	518	A	[1987STE/MAL, 1955TSY]
C ₁₆ H ₁₀	[129-00-0]	pyrene				
	$\Delta_{\text{fus}}H$	(403–433)	16.7	422.4	DSC	[2003ROJ/ORO]
	$\Delta_{\text{trs}}H$		0.29	120.8		
	$\Delta_{\text{fus}}H$		17.36	423.8		[1996DOM/HEA]
	$\Delta_{\text{sub}}H$	(341–418)	103.3 ± 2.1	380	ME	[2009SID/SID]
	$\Delta_{\text{sub}}H$	(341–418)	104.5	298	ME	[2009SID/SID]
	$\Delta_{\text{sub}}H$		98.5 ± 1.0	298	DSC	[2003ROJ/ORO]
	$\Delta_{\text{sub}}H$	(308–398)	103.1 ± 6.5	353	ME	[1998OJA/SUU]
	$\Delta_{\text{sub}}H$	(313–453)	97.9	383	GS	[1995NAS/LEN]

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	Enthalpy							
		$\Delta_{\text{sub}}H$		(369–383)	100.3 ± 0.3	353	PG	[1988SAS/JOS]
		$\Delta_{\text{sub}}H$		(283–323)	91.2 ± 0.5	303	GS	[1983SON/ZOL]
		$\Delta_{\text{sub}}H$		(398–423)	100.2 ± 0.4	410	IP	[1980SMI/STE]
		$\Delta_{\text{sub}}H$			101.0 ± 0.5		C	[1974MAL/BAR]
		$\Delta_{\text{sub}}H$		(348–419)	100.8 ± 1.5		ME	[1974MAL/BAR]
		$\Delta_{\text{sub}}H$			95.7		ME	[1953BRA/CLE2, 1977PED/RYL, 1970COX/PIL]
		$\Delta_{\text{sub}}H$		(298–363)	100.5	330	ME	[1958HOY/PEP]
		$\Delta_{\text{sub}}H$		(345–358)	100.1 ± 1.7	351	ME	[1952INO/SHI]
		Δ_vH			92.4 ± 1.1	298	CGC	[2008HAN/NUT]
		Δ_vH		(423–493)	87.2 ± 1.3	298	GC	[2006TEO/BAR]
		Δ_vH		(343–453)	78.6	398	GC	[1990HIN/BID2]
		Δ_vH		(413–467)	76	428		[1988SAS/JOS]
		Δ_vH		(398–458)	76.4	440		[1980SMI/STE]
		Δ_vH		(513–668)	73	528	A	[1987STE/MAL, 1955TSY]
C ₁₆ H ₁₀ ClN ₃ O	[1978-95-7]		1-[(2-chloro-3-pyridinyl)carbonyl]-1,2-dihydro-2-quinolinecarbonitrile					
		$\Delta_{\text{fus}}H$			50.89	489.8	DSC	[2005LIZ/ZAB]
C ₁₆ H ₁₀ N ₂ O ₂	[482-89-3]		2-(1,3-dihydro-3-oxo-2 <i>H</i> -indol-2-ylidene)-1,2-dihydro-3 <i>H</i> -indol-3-one (C.I. Vat Blue 1)					
		$\Delta_{\text{sub}}H$		(519–634)	136	577	GS	[1986NIS/AND]
C ₁₆ H ₁₀ O	[5315-79-7]		1-hydroxypyrene					
		$\Delta_{\text{sub}}H$		(369–394)	129.0 ± 3.2	382	ME	[1998OJA/SUU]
C ₁₆ H ₁₀ O	[243-24-3]		2,3,5,6-dibenzoxalene (benz[b]indeno[1,2- <i>e</i>]pyran)					
		$\Delta_{\text{sub}}H$		(375–388)	125.9	381.5	A	[1987STE/MAL]
		$\Delta_{\text{sub}}H$			129.4 ± 1.3			[1966GEI/QUI, 1970COX/PIL]
C ₁₆ H ₁₀ O	[955-83-9]		2,5-diphenylfuran					
		$\Delta_{\text{sub}}H$			102	340	HSA	[1989SCH/PEN]
C ₁₆ H ₁₀ O	[205-39-0]		benzo[b]naphtho[1,2 <i>d</i>]furan					
		$\Delta_{\text{fus}}H$			13.7	315.9	DSC	[2010KES/AUC]
C ₁₆ H ₁₀ O	[239-30-5]		benzo[b]naphtho[2,1 <i>d</i>]furan					
		$\Delta_{\text{fus}}H$			20.9	373.7	DSC	[2010KES/AUC]
C ₁₆ H ₁₀ S	[239-35-0]		1,2-benzodiphenylene sulfide					
		$\Delta_{\text{sub}}H$		(325–373)	111.9 ± 1.2	349	ME	[1998OJA/SUU]
C ₁₆ H ₁₀ S	[205-43-6]		dibenzo[b]naphtho[1,2 <i>d</i>]thiophene					
		$\Delta_{\text{fus}}H$			19.0	375.5	DSC	[2010KES/AUC]
C ₁₆ H ₁₀ S ₄	[5632-29-1]		2,2',5',2''-quaterthiophene					
		$\Delta_{\text{sub}}H$		(383–413)	132.6		ME	[1998KLO/LAU]
		$\Delta_{\text{sub}}H$		(428–457)	145.6		ME	[1998KLO/LAU]
C ₁₆ H ₁₁ F ₃ O	[172424-69-0]		4-ethoxy-2',3',4'-trifluorodiphenylacetylene					
		$\Delta_{\text{fus}}H$			32.2	356.8	DSC	[1995HSU/TSA]
C ₁₆ H ₁₁ N ₃ O	[1978-94-6]		1-[(3-pyridinyl)carbonyl]-1,2-dihydro-2-quinolinecarbonitrile					
		$\Delta_{\text{fus}}H$			31.01	412.5	DSC	[2005LIZ/ZAB]
C ₁₆ H ₁₂	[6572-60-7]		[2.2]-paracyclophane-1,9-diene					
		$\Delta_{\text{fus}}H$			30.71	505.9	DSC	[2003DEM/KOZ]
		$\Delta_{\text{sub}}H$		(318–343)	92.0 ± 1.2	331	GS	[2003DEM/KOZ]
		$\Delta_{\text{sub}}H$		(318–343)	93.1 ± 1.2	298	GS	[2003DEM/KOZ]
C ₁₆ H ₁₂	[605-02-7]		1-phenylnaphthalene					

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		T _m (K)	Method	Reference
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)			
C ₁₆ H ₁₂		$\Delta_{\text{sub}}H$	(313–453)	88.6	383	GS [1995NAS/LEN]
		$\Delta_{\text{v}}H$		81.1 ± 1.8	298	C [2008ROU/LIM]
	[612-94-2]	2-phenylnaphthalene				
C ₁₆ H ₁₂ F ₂		$\Delta_{\text{fus}}H$		17.9	373.5	DSC [2008ROU/LIM]
		$\Delta_{\text{sub}}H$	(333–353)	106.6 ± 0.4	343	ME [2008ROU/LIM]
		$\Delta_{\text{sub}}H$	(333–353)	107.6 ± 0.6	298	ME [2008ROU/LIM]
C ₁₆ H ₁₂ F ₂ O	[145698-42-6]	4-ethyl-3',4'-difluorodiphenylacetylene				
		$\Delta_{\text{fus}}H$		16.6	301.2	DSC [1995HSU/TSA]
C ₁₆ H ₁₂ F ₂ O	[172424-66-7]	4-ethoxy-2',4'-difluorodiphenylacetylene				
		$\Delta_{\text{fus}}H$		27.0	343.4	DSC [1995HSU/TSA]
C ₁₆ H ₁₂ N ₂ O	[842-07-9]	2-hydroxy-1-phenylazonaphthalene				
		$\Delta_{\text{sub}}H$	(350–374)	116.7 ± 5.4	362	[1984KRI]
C ₁₆ H ₁₂ O ₂	[134852-10-1]	5-hydroxymethylene-5H-6,7-dihydrodibenzo[a,c]-cyclohepten-6-one				
		$\Delta_{\text{fus}}H$		16.9	357.7	DSC [2006PER/CON]
		$\Delta_{\text{v}}H$		116.1 ± 12.1	298	CGC [2006PER/CON]
C ₁₆ H ₁₂ S ₂	[16212-85-4]	3,6-diphenyl-1,2-dithiin				
		$\Delta_{\text{sub}}H$		174.5 ± 2.5	355	[1973GEI/SAW, 1977PED/RYL]
		$\Delta_{\text{sub}}H$		183.1 ± 2.5	298	[1973GEI/SAW, 1977PED/RYL]
C ₁₆ H ₁₂ S ₂	[92802-27-2]	2,6-diphenyl-1,4-dithiin				
		$\Delta_{\text{fus}}H$ (I)		20.9	336.6	
		$\Delta_{\text{fus}}H$ (II)		24.6	350.7	DSC [2004PIA/SUG]
C ₁₆ H ₁₃ ClN ₂ O	[439-14-5]	7-chloro-1,3-dihydro-1-methyl-5-phenyl-2H-1,4-benzodiazepin-2-one (diazepam)				
		$\Delta_{\text{fus}}H$		24.7	404.8	DSC [2006WAS/HOL]
		$\Delta_{\text{fus}}H$		25.49	403.6	DSC [2001VER/AUG]
C ₁₆ H ₁₃ ClN ₂ O ₂	[846-50-4]	7-chloro-1,3-dihydro-3-hydroxy-1-methyl-5-phenyl-2H-1,4-benzodiazepin-2-one (temazepam)				
		$\Delta_{\text{fus}}H$		27.4	432.6	[1998VAN/AUG]
		$\Delta_{\text{fus}}H$		25.58	432.5	DSC [1992RIC/MCC]
C ₁₆ H ₁₃ FO	[na]	4-ethoxy-4'-fluorodiphenylacetylene				
		$\Delta_{\text{fus}}H$		22.8	354.4	[1995HSU/TSA]
C ₁₆ H ₁₃ N	[90-32-2]	N-phenyl-1-naphthylamine				
		$\Delta_{\text{sub}}H$	(313–333)	96.5	323	GS [1987STE/MAL, 1971FEL/KUZ]
		$\Delta_{\text{v}}H$	(338–368)	89.6	353	A [1987STE/MAL]
C ₁₆ H ₁₃ N	[135-88-6]	N-phenyl-2-naphthylamine				
		$\Delta_{\text{sub}}H$	(333–363)	115.8	348	GS [1987STE/MAL, 1971FEL/KUZ]
		$\Delta_{\text{v}}H$	(383–520)	88.7	398	A [1987STE/MAL]
C ₁₆ H ₁₃ NO	[37170-96-0]	9-acetamidoanthracene				
		$\Delta_{\text{sub}}H$	(446–500)	134.8	461	RG [1958KLO, 1987STE/MAL]
C ₁₆ H ₁₃ NO	[93-45-8]	N-(4-hydroxyphenyl)-2-naphthylamine				
		$\Delta_{\text{sub}}H$	(373–408)	126.8	390	GS [1971FEL/KUZ]
C ₁₆ H ₁₃ NO ₂	[5960-55-4]	1-(dimethylamino)-9,10-anthraquinone				
		$\Delta_{\text{sub}}H$	(396–408)	U 3.6	402	A [1987STE/MAL]
C ₁₆ H ₁₃ NO ₂	[4465-58-1]	1-(2-hydroxyethylamino)-9,10-anthraquinone				

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
	$\Delta_{\text{sub}}H$	(403–417)	152.7	410	ME	[1960BRA/BIR, 1966JON/KRA]
C ₁₆ H ₁₃ NO ₃	[483362-77-2]	1-[(4-nitrophenyl)ethynyl]-4-ethoxybenzene				
	$\Delta_{\text{fus}}H$		26.02	388.1	DSC	[2002SPA/DZI]
C ₁₆ H ₁₃ NO ₅	[na]	1-amino-2-hydroxyethyl-4-hydroxy-9,10-anthraquinone				
	$\Delta_{\text{sub}}H$		135.2			[1984KAR/KRU]
C ₁₆ H ₁₃ NO ₇	[na]	2-acetoxybenzoic acid, 3'-(nitrooxymethyl)phenyl ester				
	$\Delta_{\text{fus}}H$ (I)		33.79	335.2		
	$\Delta_{\text{fus}}H$ (I)		26.83	328.5		[2004FOP/SAN]
C ₁₆ H ₁₄	[781-17-9]	4,5,9,10-tetrahydropyrene				
	$\Delta_{\text{trs}}H$		1.85	319.9		
	$\Delta_{\text{trs}}H$		0.13	385.1		
	$\Delta_{\text{fus}}H$		17.09	412.8		[1993CHI/KNI2]
	$\Delta_{\text{sub}}H$	(385–410)	90.4	400	IP	[1993CHI/KNI2]
	$\Delta_{\text{v}}H$		70.9	440	EB,IP	[1993CHI/KNI2]
	$\Delta_{\text{v}}H$		68.1	480	EB,IP	[1993CHI/KNI2]
	$\Delta_{\text{v}}H$		65.3	520	EB,IP	[1993CHI/KNI2]
	$\Delta_{\text{v}}H$		62.5	560	EB,IP	[1993CHI/KNI2]
	$\Delta_{\text{v}}H$		59.5	600	EB,IP	[1993CHI/KNI2]
	$\Delta_{\text{v}}H$		56.4	640	EB,IP	[1993CHI/KNI2]
C ₁₆ H ₁₄	[20279-21-4]	1,2,3,10b-tetrahydrofluoranthene				
	$\Delta_{\text{v}}H$	(400–469)	68.0	415	A	[1987STE/MAL]
C ₁₆ H ₁₄	[781-43-1]	9,10-dimethylanthracene				
	$\Delta_{\text{sub}}H$	(363–378)	109.4 ± 1.7	371	ME	[2006RIB/AMA2]
	$\Delta_{\text{sub}}H$	(363–378)	113.0 ± 1.7	298	ME	[2006RIB/AMA2]
	$\Delta_{\text{sub}}H$	(372–382)	114.6	377		[1987STE/MAL]
	$\Delta_{\text{sub}}H$	(381–434)	103.2	396	RG	[1958KLO, 1987STE/MAL]
	$\Delta_{\text{v}}H$		94.5 ± 0.2	298	CGC	[2008HAN/NUT]
C ₁₆ H ₁₄	[52251-71-5]	2-ethylanthracene				
	$\Delta_{\text{sub}}H$	(343–359)	104.9 ± 0.6	351	ME	[2006RIB/AMA2]
	$\Delta_{\text{sub}}H$	(343–359)	107.6 ± 0.6	298	ME	[2006RIB/AMA2]
	$\Delta_{\text{v}}H$		91.4 ± 1.1	298	CGC	[2008HAN/NUT]
C ₁₆ H ₁₄	[1576-69-8]	2,7-dimethylphenanthrene				
	$\Delta_{\text{sub}}H$		106.7 ± 0.8		ME	[1965KAR/KYB, 1970COX/PIL]
C ₁₆ H ₁₄	[3674-69-9]	4,5-dimethylphenanthrene				
	$\Delta_{\text{sub}}H$	(313–453)	85.7	383	GS	[1995NAS/LEN]
	$\Delta_{\text{sub}}H$		104.6 ± 1.3		ME	[1965KAR/KYB, 1970COX/PIL]
C ₁₆ H ₁₄	[604-83-1]	9,10-dimethylphenanthrene				
	$\Delta_{\text{sub}}H$		119.5 ± 1.3			[1966GEI/QUI, 1970COX/PIL]
C ₁₆ H ₁₄	[866-65-7]	1,4-diphenylbutadiene				
	$\Delta_{\text{sub}}H$		87.0		RG	[1958KLO]
C ₁₆ H ₁₄	[31297-12-8]	[2.2]-paracyclophane-1-ene				
	$\Delta_{\text{fus}}H$		17.61	469.9		[2003DEM/KOZ]
	$\Delta_{\text{sub}}H$	(318–343)	93.3 ± 1.1	331	GS	[2003DEM/KOZ]
	$\Delta_{\text{sub}}H$	(318–343)	94.4 ± 1.1	298	GS	[2003DEM/KOZ]

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
C ₁₆ H ₁₄ Cl ₂ O ₂	[4359-34-6] $\Delta_{\text{sub}}H$	1,1-dichloro-2,2-bis-(4-methoxyphenyl)ethylene	79.2			[1995RUL/RAK, 1989LUB/JAN]
C ₁₆ H ₁₄ Cl ₂ O ₃	[na] $\Delta_{\text{fus}}H$	ethyl 2-hydroxy-2,2-bis-(4-chlorophenyl)acetate	23.48	310.4		[1991ACR]
C ₁₆ H ₁₄ Cl ₂ O ₃	[51338-27-3] $\Delta_{\text{fus}}H$	methyl 2-(4-(2,4-dichlorophenoxy)phenoxy)propionate	27.08	314.4	DSC	[1990DON/DRE]
C ₁₆ H ₁₄ F ₄ N ₄ O ₂	[80135-84-8] $\Delta_{\text{sub}}H$	N-methyl-N-(2,2,3,3-tetrafluoropropyl)-4-[(4-nitrophenyl)azo]benzenamine	100.8		UV	[1984KAR/ROD]
C ₁₆ H ₁₄ N ₂	[19311-79-6] $\Delta_{\text{fus}}H$	1-methyl-3,5-diphenylpyrazole	17.46	332.9	AC	[2001DI/SUN]
C ₁₆ H ₁₄ N ₂ OS	[688319-93-9] $\Delta_{\text{fus}}H$	N-(4-acetylphenyl)-4 <i>H</i> -3,1-benzothiazin-2-amine	16.1	436.2	DSC	[2004GON/KOS]
C ₁₆ H ₁₄ N ₂ O ₂	[2475-44-7] $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$	1,4-bis(N-methylamino)anthra-9,10-quinone	151.8 ± 3.9	399		[1984KRI] [1967DAT/KAN, 1991HOR]
C ₁₆ H ₁₄ N ₂ O ₂	[65990-96-7] $\Delta_{\text{sub}}H$	2-methyl-3-(phenylmethyl)quinoxaline-1,4-dioxide	146.6 ± 3.2	298	C	[2004RIB/GOM2]
C ₁₆ H ₁₄ N ₂ O ₃ S	[181695-72-7] $\Delta_{\text{fus}}H$	4-(5-methyl-3-phenyl-4-isoxazolyl)benzenesulfonamide (valdecobix)	30.35	446.4		[2004AMB/MAH]
C ₁₆ H ₁₄ N ₄ O ₂	[340820-69-1] $\Delta_{\text{fus}}H$	4-(4-methylphenyl)-5-(2-pyridinyl)-4 <i>H</i> -1,2,4-triazole-3-carboxylic acid, methyl ester	38.2	423.4		[2005SIK/MOD]
C ₁₆ H ₁₄ O	[838-15-3] $\Delta_{\text{fus}}H$ $\Delta_{\text{sub}}H$ $\Delta_{\text{v}}H$	2,3:6,7-dibenzocycloocta-2,6-dien-1-one	17.2	366.6	DSC	[2003PER/CON]
			103.3 ± 3.2	298	Vap+Fus	[2003PER/CON]
			90.6 ± 2.0	298	CGC	[2003PER/CON]
C ₁₆ H ₁₄ O	[na] $\Delta_{\text{fus}}H$ $\Delta_{\text{sub}}H$ $\Delta_{\text{v}}H$	2,3:7,8-dibenzocycloocta-2,7-dien-1-one	27.8	420	DSC	[2003PER/CON]
			112.8 ± 4.1	298	Vap+Fus	[2003PER/CON]
			92.0 ± 2.9	298	CGC	[2003PER/CON]
C ₁₆ H ₁₄ O ₂	[103-41-3] $\Delta_{\text{v}}H$	benzyl cinnamate	89.4	461	A	[1987STE/MAL, 1947STU]
C ₁₆ H ₁₄ O ₂	[495-71-6] $\Delta_{\text{trs}}H$ $\Delta_{\text{fus}}H$	1,2-dibenzoylthane	0.22	187		
			38.99	418.6		[1996DOM/HEA]
C ₁₆ H ₁₄ O ₃	[22071-15-4] $\Delta_{\text{fus}}H$ $\Delta_{\text{fus}}H$ $\Delta_{\text{fus}}H$ $\Delta_{\text{sub}}H$	(+)- α -(3-benzoylphenyl)propionic acid ((+)-ketoprofen)	37.3	368	DSC	[2006WAS/HOL, 2008WAS/HOL]
			25.04	369		[1998MUR/BET2, 1999MUR/FAU]
			28.23	367.4		[1995ESP/BIS]
		(341–365)	110.1 ± 0.5		GS	[2003PER/KUR2]
C ₁₆ H ₁₄ O ₃	[36330-85-5] $\Delta_{\text{fus}}H$ $\Delta_{\text{fus}}H$	3-(4-biphenylcarbonyl)propionic acid (fenbufen)	41.1	462.9	DSC	[2008KUR/PER]
			46.2	459.3	DSC	[2006WAS/HOL]

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
	Enthalpy							
		$\Delta_{\text{sub}}H$		(378–420)	154.9 ± 0.8	298	GS	[2008KUR/PER]
C ₁₆ H ₁₄ O ₄	[5673-22-3]		1,2- <i>cis</i> -dicarbomethoxyacenaphthene		37.66	398.2		[1974CAN/JAC]
		$\Delta_{\text{fus}}H$						
C ₁₆ H ₁₄ O ₄	[51869-93-3]		1,2- <i>trans</i> -dicarbomethoxyacenaphthene		27.61	388.7		[1974CAN/JAC]
		$\Delta_{\text{fus}}H$						
C ₁₆ H ₁₄ O ₄	[56137-73-6]		1,3-dicarbomethoxyacenaphthene		23.01	371.2		[1974CAN/JAC]
		$\Delta_{\text{fus}}H$						
C ₁₆ H ₁₄ O ₄	[51870-00-9]		1,5-dicarbomethoxyacenaphthene		28.87	386.7		[1974CAN/JAC]
		$\Delta_{\text{fus}}H$						
C ₁₆ H ₁₄ O ₄	[4599-96-6]		5,6-dicarbomethoxyacenaphthene		34.73	450.2		[1974CAN/JAC]
		$\Delta_{\text{fus}}H$						
C ₁₆ H ₁₄ O ₆	[36063-02-2]		1,2,3-tricarbomethoxynaphthalene		23.7	362.7	DSC	[1993ACR]
		$\Delta_{\text{fus}}H$						
C ₁₆ H ₁₄ O ₆	[36063-03-3]		1,2,4-tricarbomethoxynaphthalene		32.1	393.7	DSC	[1993ACR]
		$\Delta_{\text{fus}}H$						
C ₁₆ H ₁₄ O ₆	[68267-11-8]		1,2,5-tricarbomethoxynaphthalene		25.5	363	DSC	[1993ACR]
		$\Delta_{\text{fus}}H$						
C ₁₆ H ₁₄ O ₆	[36063-04-4]		1,2,6-tricarbomethoxynaphthalene		35.9	416.7	DSC	[1993ACR]
		$\Delta_{\text{fus}}H$						
C ₁₆ H ₁₄ O ₆	[68257-10-7]		1,2,7-tricarbomethoxynaphthalene		36.1	427.2	DSC	[1993ACR]
		$\Delta_{\text{fus}}H$						
C ₁₆ H ₁₄ O ₆	[36440-23-0]		1,2,8-tricarbomethoxynaphthalene		24.8	366.7	DSC	[1993ACR]
		$\Delta_{\text{fus}}H$						
C ₁₆ H ₁₄ O ₆	[36440-28-5]		1,3,5-tricarbomethoxynaphthalene		25.9	402.7	DSC	[1993ACR]
		$\Delta_{\text{fus}}H$						
C ₁₆ H ₁₄ O ₆	[36440-29-6]		1,3,6-tricarbomethoxynaphthalene		37.4	469.7	DSC	[1993ACR]
		$\Delta_{\text{fus}}H$						
C ₁₆ H ₁₄ O ₆	[36440-30-9]		1,3,7-tricarbomethoxynaphthalene		37.2	446.7	DSC	[1993ACR]
		$\Delta_{\text{fus}}H$						
C ₁₆ H ₁₄ O ₆	[36440-24-1]		1,3,8-tricarbomethoxynaphthalene		27.7	388.2	DSC	[1993ACR]
		$\Delta_{\text{fus}}H$						
C ₁₆ H ₁₄ O ₆	[36440-25-2]		1,4,5-tricarbomethoxynaphthalene		26.5	402.2	DSC	[1993ACR]
		$\Delta_{\text{fus}}H$						
C ₁₆ H ₁₄ O ₆	[36063-05-5]		1,4,6-tricarbomethoxynaphthalene		30.2	409.2	DSC	[1993ACR]
		$\Delta_{\text{fus}}H$						
C ₁₆ H ₁₄ O ₆	[36440-26-3]		2,3,5-tricarbomethoxynaphthalene		41.0	401.7	DSC	[1993ACR]
		$\Delta_{\text{fus}}H$						
C ₁₆ H ₁₄ O ₆	[36440-27-4]		2,3,6-tricarbomethoxynaphthalene		34.4	399.2	DSC	[1993ACR]
		$\Delta_{\text{fus}}H$						
C ₁₆ H ₁₄ O ₆	[520-33-2]		2,3-dihydro-5,7-dihydroxy-2-(3-hydroxy-4-methoxyphenyl)-4 <i>H</i> -1-benzopyran-4-one (hesperitin)		35.9	499.2	DSC	[2007CHE/HUM]
		$\Delta_{\text{fus}}H$						
C ₁₆ H ₁₅ BrO	[556052-89-2]		4-bromo-4'-(3-butenyloxy)-1,1'-biphenyl					

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	Enthalpy							
C ₁₆ H ₁₅ Cl ₂ NO ₂		$\Delta_{\text{trs}}H$		13.3	324.2			
		$\Delta_{\text{fus}}H$		15.8	396.8	DSC	[2003WIL/VAN]	
C ₁₆ H ₁₅ Cl ₂ NO ₂	[117-26-0]		1,1- <i>bis</i> -(4-chlorophenyl)-2-nitrobutane					
		$\Delta_{\text{fus}}H$		15.41	330.3	DSC	[1990DON/DRE]	
C ₁₆ H ₁₅ Cl ₃ O ₂	[30667-90-3]		1-methoxy-2-(2,2,2-trichloro-1-(4-methoxyphenyl)ethyl)benzene					
		$\Delta_{\text{fus}}H$		22.45	347.6	DSC	[1990DON/DRE]	
C ₁₆ H ₁₅ Cl ₃ O ₂	[72-43-5]		1,1'-(2,2,2-trichloroethylidene- <i>bis</i> -(4-methoxy)benzene					
		$\Delta_{\text{fus}}H$		27.48	360.6	DSC	[1990DON/DRE]	
C ₁₆ H ₁₅ IO ₃ S	[313057-06-6]		4-(4-pentyloxy)phenyl 5-iodo-2-thiophene carboxylate					
		$\Delta_{\text{fus}}H$		68.2	332.7	DSC	[2000WU/WAN]	
C ₁₆ H ₁₅ N	[na]		4'-propylbiphenyl-4-carbonitrile					
		$\Delta_{\text{fus}}H$		22.7	338.8		[1996DOM/HEA]	
C ₁₆ H ₁₅ NO	[18594-93-9]		3-anilino-1-phenylbut-2-enone					
		$\Delta_{\text{sub}}H$		126.8 ± 3.0	298	C	[1993RIB/RIB]	
C ₁₆ H ₁₅ NO ₄	[483362-66-9]		2-(4-nitrophenyl)-1-(4-ethoxyphenyl)ethanone					
		$\Delta_{\text{fus}}H$		28.2	390.3	DSC	[2002SPA/DZI]	
C ₁₆ H ₁₅ N ₅ O ₃	[157892-00-7]		6-phenyl-3,9-dihydro-3-[(2-hydroxyethoxy)methyl]-9-oxo-5 <i>H</i> -imidazol[1,2- <i>a</i>]pyrrole					
		$\Delta_{\text{fus}}H$		63.78	485.8	DSC	[1999ZIE/GOL]	
C ₁₆ H ₁₆	[1732-13-4]		1,2,3,6,7,8-hexahydropyrene					
		$\Delta_{\text{trs}}H$		5.02	377			
		$\Delta_{\text{fus}}H$		18.09	407.7		[1993CHI/KNI2]	
		$\Delta_{\text{sub}}H$	(390–405)	92.3	398	IP	[1993CHI/KNI2]	
		Δ_vH		72.0	440	EB,IP	[1993CHI/KNI2]	
		Δ_vH		69.4	480	EB,IP	[1993CHI/KNI2]	
		Δ_vH		66.8	520	EB,IP	[1993CHI/KNI2]	
		Δ_vH		64.2	560	EB,IP	[1993CHI/KNI2]	
C ₁₆ H ₁₆	[1633-22-3]		[2.2]- <i>para</i> -cyclophane					
		$\Delta_{\text{fus}}H$		0.21	323.2		[1970AND/WES]	
		$\Delta_{\text{sub}}H$	(353–409)	96.4 ± 1.5		TSGC	[1980NIS/SAK]	
		$\Delta_{\text{sub}}H$		96.3 ± 4.2			[1973ROD/WES, 1977PED/RYL]	
C ₁₆ H ₁₆		$\Delta_{\text{sub}}H$	(343–383)	92.9 ± 0.84	363	ME	[1966BOY, 1987STE/MAL, 1970COX/PIL]	
	[2319-97-3]		[2.2]- <i>meta</i> -cyclophane					
		$\Delta_{\text{fus}}H$		21.42	404		[1969SHI/MCN]	
		$\Delta_{\text{sub}}H$	(308–332)	91.6 ± 1.7	320	ME	[1969SHI/MCN, 1977PED/RYL, 1987STE/MAL]	
C ₁₆ H ₁₆		$\Delta_{\text{sub}}H$		92.0 ± 2	298	ME	[1969SHI/MCN, 1977PED/RYL]	
	[5385-36-4]		[2.2]- <i>meta-para</i> -cyclophane					
		$\Delta_{\text{trs}}H$		0.98	315			
		$\Delta_{\text{fus}}H$		12.76	354		[1969SHI/MCN]	
C ₁₆ H ₁₆		$\Delta_{\text{sub}}H$	(311–328)	86.6	336	ME	[1969SHI/MCN, 1977PED/RYL, 1987STE/MAL]	
		$\Delta_{\text{sub}}H$		87.5 ± 0.9	298	ME	[1969SHI/MCN, 1977PED/RYL]	
	[2919-20-2]		1,1- <i>bis</i> -(4-methylphenyl)ethene					
	$\Delta_{\text{fus}}H$		23.31	334.1		[1999VER6]		

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	Enthalpy							
		$\Delta_{\text{sub}}H$		(309–332)	100.3 ± 1.4	320	GS	[1999VER6]
		$\Delta_{\text{sub}}H$			101.0 ± 1.4	298	GS	[1999VER6]
C ₁₆ H ₁₆ ClN	[113788-75-3]		4-chlorobenzylidene-4'-propylaniline		24.61	343.7	DSC	[1999GAL/COL]
C ₁₆ H ₁₆ N ₂ O ₂	[94-93-9]		N,N'-bis(salicylaldehyde)ethylenediimine		34.09	397.9	DSC	[2004RIB/GON]
		$\Delta_{\text{sub}}H$		(348–363)	141.3 ± 3.2	298	ME	[2004RIB/GON]
C ₁₆ H ₁₆ N ₂ O ₂	[2299-73-2]		2-[(4-methoxyphenyl)methylene]hydrazone-4-methoxybenzaldehyde (anisaldazine)		29.75	442		[1996DOM/HEA]
C ₁₆ H ₁₆ N ₂ O ₄	[13684-56-5]		ethyl [3-[[[(phenylamino)carbonyl]oxy]phenyl]carbamate		32.75	394.1	DSC	[1990DON/DRE]
C ₁₆ H ₁₆ N ₂ O ₄	[13684-63-4]		methyl 3- <i>m</i> -tolylcarbamoyloxyphenylcarbamate		39.62	423.8	DSC	[1990DON/DRE]
C ₁₆ H ₁₆ N ₂ O ₄	[54946-22-4]		N-propylthalidomide		27.28	409.2	DTA	[2002GOO/LAI]
C ₁₆ H ₁₆ O	[130935-82-9]		6-hydroxymethyl-5,6-dihydro-7 <i>H</i> -dibenzo[<i>a,c</i>]cycloheptane		25.9	405.7	DSC	[2005PER/BAN]
C ₁₆ H ₁₆ O	[29817-04-7]		5-hydroxymethyl-5,6-dihydro-7 <i>H</i> -dibenzo[<i>a,c</i>]cycloheptane		16	352.5	DSC	[2005PER/BAN]
C ₁₆ H ₁₆ O ₂	[29783-24-2]		<i>trans</i> -9,10- <i>bis</i> hydroxymethyl-9,10-dihydrophenanthrene		30.3	450.8	DSC	[2005PER/BAN]
C ₁₆ H ₁₆ O ₂	[29790-58-7]		<i>trans</i> -5-hydroxymethyl-5,6-dihydro-7 <i>H</i> -dibenzo[<i>a,c</i>]cycloheptan-6-ol		31.8	460.2	DSC	[2005PER/BAN]
C ₁₆ H ₁₆ O ₂	[na]		(<i>d</i>) 2-(<i>p</i> -methoxyphenyl)propiophenone		21.76	326		[1976LEC/COL]
C ₁₆ H ₁₆ O ₂	[na]		(<i>dl</i>) 2-(<i>p</i> -methoxyphenyl)propiophenone		26.36	353		[1976LEC/COL]
C ₁₆ H ₁₆ O ₂	[46863-20-1]		(2-hydroxyphenyl)-2,4,6-trimethylphenylmethanone		0.49	353.2	DTA	[1989SAL/ABA]
			Note: Reported enthalpy of fusion is too small, and the published enthalpy and entropy of fusion data are internally inconsistent					
C ₁₆ H ₁₆ O ₃	[24650-42-8]		2,2-dimethoxy-1,2-diphenylethanone		20.86	338.5		[1994SAN/DEF]
C ₁₆ H ₁₆ O ₃	[7074-00-2]		2-phenylisopropoxybenzoate	(293–313)	43.1 ± 4.2	303	ME	[1971KIP/RAB, 1977PED/RYL]
C ₁₆ H ₁₆ O ₁₀	[3327-06-8]		pentamethoxycarbonylbenzene		38.0	424.7	DSC	[1978DOZ/FUJ]
		$\Delta_{\text{sub}}H$		(389–413)	160.0 ± 0.8	401	ME	[1995JIM/MEN]
		$\Delta_{\text{sub}}H$			165.1 ± 0.8	298		[1995JIM/MEN]
		$\Delta_{\text{sub}}H$			165.1 ± 0.8	298		[1967TUR2, 1995JIM/MEN]
C ₁₆ H ₁₇ ClN ₄ O ₃	[3180-81-2]		4-(<i>N</i> -ethyl- <i>N</i> -2-hydroxyethylamino)-4'-nitro-2'-chloroazobenzene		142.7			[1968TSU/KOJ, 1988BAU/PER]
C ₁₆ H ₁₇ ClN ₄ O ₄	[na]		2,2'-[[3-chloro-4-[(4-nitrophenyl)azo]phenyl]imino]bis-ethanol					

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
	$\Delta_{\text{fus}}H$		29.78	463.2		[1988BAU/PER]
C ₁₆ H ₁₇ Cl ₂ N ₅ O ₄	[na]	1-[[2-chloro-4-[(2-chloro-4-nitrophenyl)azo]-5-(methylamino)phenyl]amino]-2-propanol N-oxide				
	$\Delta_{\text{fus}}H$		30.62	371.2		[1991BAU/WEB]
C ₁₆ H ₁₇ F	[193472-70-7]	2-fluoro-2-methyl-1,3-diphenylpropane				
	$\Delta_{\text{fus}}H$		29.7	332.7		[1997SCH/VER]
	$\Delta_{\text{sub}}H$		102.2 ± 1.1	298		[1997SCH/VER]
C ₁₆ H ₁₇ F ₁₅ O	[41049-15-4]	1,1,1,2,2,3,3,4,4,5,5,6,6,7,7-pentadecafluoro-8-hexadecanone				
	$\Delta_{\text{fus}}H$		34.2	285.8		[1992VIL/WEI]
C ₁₆ H ₁₇ NO	[15582-77-5]	1,2-diphenyl-2-N,N-dimethylamino-1-ethanone				
	$\Delta_{\text{sub}}H$		140.1 ± 1.9		B	[1994WEL/VER]
C ₁₆ H ₁₇ NO	[99081-88-6]	N-(4-isopropylphenylmethylene) benzenamine N-oxide				
	$\Delta_{\text{sub}}H$		127.2 ± 1.7	298	C	[1986KIR/ACR]
C ₁₆ H ₁₇ NO	[na]	N,N-dimethyl-2,2-diphenylacetamide				
	$\Delta_{\text{fus}}H$		25.43	407.1		[1993ACR]
C ₁₆ H ₁₇ N ₃ O ₃	[850836-66-7]	6-(acetylamino)-2-cyano-1(2 <i>H</i>)-quinolinecarboxylic acid, 1-methylethyl ester				
	$\Delta_{\text{fus}}H$		12.66	377.4	DSC	[2005LIZ/ZAB]
C ₁₆ H ₁₈	[na]	1-(2-tolyl)-2-(4-tolyl)ethane				
	$\Delta_{\text{v}}H$	(298–473)	85.6	313	A	[1987STE/MAL, 1963BES]
C ₁₆ H ₁₈	[719-79-9]	1,1-diphenylbutane				
	$\Delta_{\text{v}}H$	(298–342)	75.9 ± 0.6	320	GS	[1999VER5]
	$\Delta_{\text{v}}H$	(298–342)	77.2 ± 0.6	298	GS	[1999VER5]
C ₁₆ H ₁₈	[1520-44-1]	<i>dl</i> 1,3-diphenylbutane				
	$\Delta_{\text{sub}}H$	(288–303)	73.6	296	ME	[1974PRI/POU, 1987STE/MAL]
C ₁₆ H ₁₈	[5789-35-5]	2,3-diphenylbutane				
	$\Delta_{\text{sub}}H$	(293–348)	96.7	326		[1984BEC/RUC]
C ₁₆ H ₁₈	[1634-11-3]	2-methyl-1,1-diphenylpropane				
	$\Delta_{\text{v}}H$	(298–338)	72.0 ± 0.5	318	GS	[1999VER5]
	$\Delta_{\text{v}}H$	(298–338)	73.2 ± 0.5	298	GS	[1999VER5]
C ₁₆ H ₁₈	[530-45-0]	1,1- <i>bis</i> (4-methylphenyl)ethane				
	$\Delta_{\text{v}}H$	(298–338)	75.3 ± 0.6	318	GS	[1999VER5]
	$\Delta_{\text{v}}H$	(298–338)	76.5 ± 0.6	298	GS	[1999VER5]
C ₁₆ H ₁₈	[1625-92-9]	4- <i>tert</i> -butylbiphenyl				
	$\Delta_{\text{sub}}H$		98.1 ± 2.1	298	C	[2009MEL/PIM]
	$\Delta_{\text{v}}H$		80.0 ± 1.9	298	C	[2009MEL/PIM]
C ₁₆ H ₁₈ Cl ₄ O ₄	[3015-66-5]	dibutyl tetrachlorophthalate				
	$\Delta_{\text{v}}H$	(368–421)	99.7	383	A, T	[1987STE/MAL, 1949PER/WEB, 1999DYK/SVO]
C ₁₆ H ₁₈ FN ₃ O ₃	[70458-96-7]	1-ethyl-6-fluoro-1,4-dihydro-4-oxo-7-(1-piperazinyl)-3-quinolinecarboxylic acid (norfloxacin)				
	$\Delta_{\text{fus}}H$		32.42	492.6	DSC	[2009OLI/BER]
	$\Delta_{\text{fus}}H$		32.97	500.2	DSC	[1994YU/ZIP]
C ₁₆ H ₁₈ NO ₅	[3788-15-6]	<i>bis</i> (2,4-dimethoxyphenyl)nitrogen oxide				
	$\Delta_{\text{sub}}H$	(333–363)	144.1 ± 11.4	348	A	[1987STE/MAL, 1965KAL/ROZ]
C ₁₆ H ₁₈ N ₂ O	[na]	4- <i>n</i> -butyl-4'-hydroxyazobenzene				

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
	Enthalpy							
		$\Delta_{\text{fus}}H$			5.25	351.6		[1990JIN/KAN]
C ₁₆ H ₁₈ N ₂ OS	[373642-48-9]		N-[(3-methoxyphenyl)methyl]-N'-(phenylmethyl)thiourea		21.02	345	DSC	[2002ABB/WHO]
		$\Delta_{\text{fus}}H$						
C ₁₆ H ₁₈ N ₂ O ₂	[101225-69-8]		2,2',6,6'-tetramethylazobenzene-N,N-dioxide		107 ± 12	298	ME	[1993ACR/TUC2]
		$\Delta_{\text{sub}}H$						
C ₁₆ H ₁₈ N ₂ O ₃	[4792-83-0]		<i>p</i> -azoxyphenetole		126.2 ± 2.7	298	C	[1993ACR/TUC]
		$\Delta_{\text{sub}}H$						
C ₁₆ H ₁₈ N ₂ O ₃	[1979-22-3]		2-cyano-6-methoxy-1(2 <i>H</i>)-quinolinecarboxylic acid, 2-methylpropyl ester		24.45	346.4	DSC	[2005LIZ/ZAB]
		$\Delta_{\text{fus}}H$						
C ₁₆ H ₁₈ N ₄ O ₂	[3025-52-3]		4-(<i>N,N</i> -diethylamino)-4'-nitroazobenzene		146		GS	[1987SHI/OHK, 1991HOR]
		$\Delta_{\text{sub}}H$						
		$\Delta_{\text{sub}}H$	(422–441)	151.5 ± 4.2	431		ME	[1960BRA/BIR]
C ₁₆ H ₁₈ N ₄ O ₃	[2872-52-8]		4-(<i>N</i> -ethyl- <i>N</i> -2-hydroxyethylamino)-4'-nitroazobenzene		136.8		UV	[1984KAR/ROD, 1984KAR/KRU]
		$\Delta_{\text{sub}}H$						
		$\Delta_{\text{sub}}H$			189.5			[1968TSU/KOJ, 1988BAU/PER]
		$\Delta_{\text{sub}}H$	(420–433)	176.6 ± 1.3	426		ME	[1960BRA/BIR, 1966JON/KRA]
C ₁₆ H ₁₈ N ₄ O ₄	[na]		<i>N,N</i> -(2-hydroxyethyl)-4-(4-nitrophenyl)azoaniline		32.43	484.2		[1988BAU/PER]
		$\Delta_{\text{fus}}H$						
C ₁₆ H ₁₈ O	[93-96-9]		<i>bis</i> (α -methylbenzyl) ether		62.1	384	A	[1987STE/MAL, 1947STU]
		$\Delta_{\text{v}}H$	(369–554)					
C ₁₆ H ₁₉ BrO ₂	[164591-96-2]		4- <i>trans</i> -(4-bromophenyl)cyclohexyl (E)-2-butenate		28.4	388.2		[1995KEL/SCH]
		$\Delta_{\text{fus}}H$						
C ₁₆ H ₁₉ ClO ₂	[164591-95-1]		4- <i>trans</i> -(4-chlorophenyl)cyclohexyl (E)-2-butenate		30.2	386.2		[1995KEL/SCH]
		$\Delta_{\text{fus}}H$						
C ₁₆ H ₁₉ FO ₂	[164591-94-0]		4- <i>trans</i> -(4-fluorophenyl)cyclohexyl (E)-2-butenate		25.1	354.2		[1995KEL/SCH]
		$\Delta_{\text{fus}}H$						
C ₁₆ H ₁₉ NO	[na]		2-(4-butoxyphenyl)-5-methylpyridine		33.0	363		[2000MOR/HAR]
		$\Delta_{\text{fus}}H$						
C ₁₆ H ₁₉ N ₃	[2481-94-9]		4-(<i>N,N</i> -diethylamino)azobenzene		132.2		GS	[1987SHI/OHK, 1991HOR]
		$\Delta_{\text{sub}}H$						
		$\Delta_{\text{sub}}H$	(330–353)	91.4 ± 2.9	342			[1984KRI]
		$\Delta_{\text{sub}}H$		106.4			UV	[1984KAR/ROD]
		$\Delta_{\text{sub}}H$		106.3				[1984KAR/KRU]
C ₁₆ H ₁₉ N ₃ O ₂	[na]		<i>N,N</i> -(2-hydroxyethyl)-4-phenylazoaniline		29.96	407		[1988BAU/PER]
		$\Delta_{\text{fus}}H$						
C ₁₆ H ₂₀ N ₂	[19219-01-3]		tetracyclopropylsuccinonitrile		22.3	390		[1996DOM/HEA]
		$\Delta_{\text{fus}}H$						
C ₁₆ H ₂₀ N ₄ O ₃ S	[56211-40-6]		<i>N</i> -[[[(1-methylethyl)amino]carbonyl]-4-[(3'-methylphenyl)amino]-3-pyridinesulfonamide (torasemide)		37.2	434.7		
		$\Delta_{\text{fus}}H$ (I)						
		$\Delta_{\text{fus}}H$ (II)			29.0	430		[2002ROL/GST]
C ₁₆ H ₂₀ O ₂	[105443-43-4]		2-isopropyl-6-(1-hydroperoxy-1-methylethyl)naphthalene		24.9	335.2		[1998STE/ZAW]
		$\Delta_{\text{fus}}H$						
C ₁₆ H ₂₀ O ₃	[146683-17-2]		3-benzoyl-1,2,2-trimethylcyclopentanecarboxylic acid		20.35	387.6	DSC	[1992TER/PAU]
		$\Delta_{\text{fus}}H$						

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
C ₁₆ H ₂₀ O ₄	[96783-79-8] $\Delta_{\text{fus}}H$	2,6- <i>bis</i> -(1-hydroperoxy-1-methylethyl)naphthalene	38.3	394.2		[1998STE/ZAW]
C ₁₆ H ₂₀ O ₆ P ₂ S ₃	[3383-96-8] $\Delta_{\text{fus}}H$	O,O,O',O'-tetramethyl O,O'-thiodi- <i>p</i> -phenylene <i>bis</i> (phosphorothioate)	33.03	303.2	DSC	[1990DON/DRE]
C ₁₆ H ₂₁ Cl ₃ O ₃	[1928-47-8] $\Delta_{\text{v}}H$	2,4,5-trichlorophenoxyacetic acid, (2-ethylhexyl) ester	(460–575) 85.4	475	A,GC	[1987STE/MAL, 1966JEN/SCH]
C ₁₆ H ₂₁ Cl ₃ O ₃	[2630-15-1] $\Delta_{\text{v}}H$	2,4,5-trichlorophenoxyacetic acid, octyl ester	(460–575) 92.2	475	A,GC	[1987STE/MAL, 1966JEN/SCH]
C ₁₆ H ₂₁ N	[61203-99-4] $\Delta_{\text{trs}}H$ $\Delta_{\text{fus}}H$	4-(<i>trans</i> -4-propylcyclohexyl)benzotrile	(15–385) 20.4	316.3		
			(15–385) 1.1	319	AC	[1998ASA/SOR]
C ₁₆ H ₂₁ NO ₂	[4199-09-1] $\Delta_{\text{fus}}H$	(–) 1-(isopropylamino)-3-(1-naphthoxy)-2-propanol (propranolol)	36.25	344.7	DSC	[1999LI/ZEL]
C ₁₆ H ₂₁ NO ₂	[525-66-6] $\Delta_{\text{fus}}H$	(±) 1-(isopropylamino)-3-(1-naphthoxy)-2-propanol (propranolol)	43.45	365.5	DSC	[1999LI/ZEL]
C ₁₆ H ₂₂ ClNO ₃	[38727-55-8] $\Delta_{\text{fus}}H$	N-(chloroacetyl)-N-(2,6-diethylphenyl)glycine ethyl ester	23.84	318	DSC	[1990DON/DRE]
C ₁₆ H ₂₂ Cl ₂ O ₃	[1928-43-4] $\Delta_{\text{v}}H$	2,4-dichlorophenoxyacetic acid, (2-ethylhexyl) ester	(460–575) 83	475	A,GC	[1987STE/MAL, 1966JEN/SCH]
C ₁₆ H ₂₂ Cl ₂ O ₃	[1917-97-1] $\Delta_{\text{v}}H$	2,4-dichlorophenoxyacetic acid, (1-methylheptyl) ester	(460–575) 83	475	A,GC	[1987STE/MAL, 1966JEN/SCH]
C ₁₆ H ₂₂ Cl ₂ O ₃	[1928-44-5] $\Delta_{\text{v}}H$	2,4-dichlorophenoxyacetic acid, octyl ester	(460–573) 87.9	475	A,GC	[1987STE/MAL, 1966JEN/SCH]
C ₁₆ H ₂₂ N ₄ O ₄	[53808-87-0] $\Delta_{\text{fus}}H$	5-[[3,5-dimethoxy-4-(2-methoxyethoxy)phenyl]methyl]-2,4-pyrimidinediamine (tetroxoprim)	46.36	423.3	DSC	[2002CAI/BET]
C ₁₆ H ₂₂ O ₄	[84-74-2] $\Delta_{\text{v}}H$ $\Delta_{\text{v}}H$ $\Delta_{\text{v}}H$ $\Delta_{\text{v}}H$	dibutyl phthalate	80.4 94.0 76.1 91.7	462 329 483 300		[1988KAT] [1987STE/MAL] [1987STE/MAL] [1949BIR/BRA]
			(314–469)		A	
			(468–605)		A	
			(288–313)			
C ₁₆ H ₂₂ O ₄	[4489-61-6] $\Delta_{\text{v}}H$	di- <i>sec</i> -butyl phthalate	(313–373) 93.8	328	A, ME	[1987STE/MAL, 1948SMA/SMA]
C ₁₆ H ₂₂ O ₄	[1962-75-0] $\Delta_{\text{v}}H$	dibutyl terephthalate	(393–483) 86.2	408	A	[1987STE/MAL]
C ₁₆ H ₂₃ N	[199394-72-4] $\Delta_{\text{fus}}H$ $\Delta_{\text{sub}}H$	N-cyclohexyl-(2,4,6-trimethyl)benzaldehyde imine	25.61	339.4		[1997VER/MOR]
			104.9 ± 0.8	298	B	[1997VER/MOR]
C ₁₆ H ₂₄ N ₂	[na] $\Delta_{\text{fus}}H$	2-(4- <i>tert</i> -butylphenyl)-2-(diethylamino)acetonitrile	24.39	327.2		[1997WEL/VER]
C ₁₆ H ₂₄ N ₂ O	[98626-60-9] $\Delta_{\text{fus}}H$	N-(2,6-dimethylphenyl)-1-ethyl-2-piperidinecarboxamide	19.9	408.2	DSC	[1997NEM/ACS]
C ₁₆ H ₂₄ N ₂ OS	[862582-66-9] $\Delta_{\text{fus}}H$	2-[(diethylamino)thioxomethyl]-N,N-dimethylbenzamide	28.79	353.5	DSC	[2005ALT/COP]

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		T _m (K)	Method	Reference
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)			
C ₁₆ H ₂₄ N ₂ O ₂	[81994-74-3]	N-benzoyl-N',N'-diisobutylurea		298	C	[2000RIB/RIB]
	$\Delta_{\text{sub}}H$		137.5 ± 4.4			
C ₁₆ H ₂₄ N ₂ O ₄	[na]	nonyl N-(4-nitrophenyl)carbamate		378.6	DSC	[1993TIE/FRA]
	$\Delta_{\text{fus}}H$		37.0			
C ₁₆ H ₂₄ N ₂ S ₂	[862582-67-0]	N,N,N',N'-tetraethyl-1,2-benzenedicarbothiamide		388.4	DSC	[2005ALT/COP]
	$\Delta_{\text{fus}}H$		23.39			
C ₁₆ H ₂₄ N ₆	[125867-93-8]	1-(methylphenethylamino)-3,5-bis(dimethylamino)-s-triazine		334.2		[1991ACR]
	$\Delta_{\text{fus}}H$		20.04			
C ₁₆ H ₂₄ O ₄	[175848-65-4]	2,5-dipentoxy-1,4-benzoquinone		333.7		
	$\Delta_{\text{trs}}H$		9.0			
	$\Delta_{\text{fus}}H$		36.5	414.6	DSC	[1996KEE/VAN]
C ₁₆ H ₂₅ Cl	[412027-20-4]	chloro(pentaethyl)benzene		378	A	[1987STE/MAL, 1947STU]
	$\Delta_{\text{v}}H$	(363–558)	60.3			
C ₁₆ H ₂₅ NO ₂	[33689-71-3]	nonyl phenylcarbamate		327		[1971PRI]
	$\Delta_{\text{fus}}H$		28.07			
C ₁₆ H ₂₅ N ₃ S	[90473-97-5]	N-(diethylaminothiocarbonyl)-N',N'-diethylbenzamide		298	C	[2004RIB/SAN]
	$\Delta_{\text{sub}}H$		122.2 ± 2.0			
C ₁₆ H ₂₆	[104-72-3]	decylbenzene		298	GS	[2006VER]
	$\Delta_{\text{v}}H$	(318–363)	78.2 ± 0.3			
	$\Delta_{\text{v}}H$	(313–433)	78.0	328		[1993KAS/MOK]
	$\Delta_{\text{v}}H$	(371–427)	75.1	386	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$		79.8	298		[1971WIL/ZWO]
	$\Delta_{\text{v}}H$	(475–571)	61.6	490	A, IP	[1987STE/MAL, 1954CAM/FOR]
C ₁₆ H ₂₆	[605-01-6]	pentaethylbenzene		374	A	[1987STE/MAL, 1947STU]
	$\Delta_{\text{v}}H$	(359–550)	56.5			
C ₁₆ H ₂₆ O	[4130-42-1]	2,6-di- <i>tert</i> -butyl-4-ethylphenol		348		[1953STA/MUL]
	$\Delta_{\text{v}}H$	(362–557)	62.8			
	$\Delta_{\text{v}}H$	(362–557)	60.4	373		[1953STA/MUL]
	$\Delta_{\text{v}}H$	(362–557)	58.6	398		[1953STA/MUL]
	$\Delta_{\text{v}}H$	(362–557)	57.3	423		[1953STA/MUL]
	$\Delta_{\text{v}}H$	(362–557)	52.6	473		[1953STA/MUL]
C ₁₆ H ₂₆ O	[6287-47-4]	4,6-di- <i>tert</i> -butyl-2-ethylphenol		428	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$	(413–556)	61.9			
	$\Delta_{\text{v}}H$	(413–562)	57.3	423		[1953STA/MUL]
	$\Delta_{\text{v}}H$	(413–562)	52.6	473		[1953STA/MUL]
C ₁₆ H ₂₆ O	[70766-54-0]	2,4-di- <i>tert</i> -butyl-5,6-dimethylphenol		446	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$	(431–565)	69.1			
C ₁₆ H ₂₆ O	[19245-41-1]	2,4-di- <i>tert</i> -butyl-5-ethylphenol		399	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$	(384–563)	69.3			
C ₁₆ H ₂₆ O	[na]	2,4,5-triisopropylbenzyl alcohol		327	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$	(312–346)	113.1			
C ₁₆ H ₂₆ O ₄	[965-40-2]	dicyclohexyl succinate		298	GS	[2008LIP/KRA]
	$\Delta_{\text{v}}H$	(338–365)	98.0 ± 0.8			
C ₁₆ H ₂₆ O ₁₁	[na]	diethylene glycol dicarboxylic acid, di[1-(ethoxycarbonyl)ethyl] ester		433	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$	(418–503)	99.3			

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₆ H ₂₈	[3752-92-9]	tricyclopentylmethane				
	$\Delta_v H$	(273–351)	77.8	288	A	[1987STE/MAL, 1964MOR]
	$\Delta_v H$	(371–429)	71.4	386	A	[1987STE/MAL]
C ₁₆ H ₂₈	[283-68-1]	tricyclo[8.2.2.2 ^{4,7}]hexadecane				
	$\Delta_{\text{sub}} H$	(316–338)	91.6 ± 0.9	327	ME	[1969SHI/MCN, 1977PED/RYL]
	$\Delta_{\text{sub}} H$	(316–338)	85.2	327		[1987STE/MAL]
C ₁₆ H ₂₈ O ₂	[31067-25-1]	1,9-cyclohexadecanedione				
	$\Delta_{\text{trs}} H$		17.95	301.2		
	$\Delta_{\text{fus}} H$		8.03	351.2		[1972ALV/BOR]
C ₁₆ H ₂₈ O ₄	[na]	1,7-cyclododecanedione <i>bis</i> ethylene ketal				
	$\Delta_{\text{fus}} H$		36.94	478.2		[1972ALV/BOR]
C ₁₆ H ₃₀ N ₂	[19219-01-3]	tetracyclopropylsuccinonitrile				
	$\Delta_{\text{sub}} H$		110.2 ± 1.5			[1984BER/BEC2]
C ₁₆ H ₃₀ O	[541-91-3]	3-methylcyclopentadecanone				
	$\Delta_v H$	(391–601)	63.5	406	A	[1987STE/MAL]
C ₁₆ H ₃₀ O	[2550-52-8]	cyclohexadecanone				
	$\Delta_{\text{sub}} H$		82.0			[1938WOL/WEG, 1960JON]
C ₁₆ H ₃₀ O	[174155-58-9]	(Z) 3-hexadecenal				
	$\Delta_v H$	(373–413)	89.9	298	CGC	[1996KOU/HOS, 2000OVA/KOU]
C ₁₆ H ₃₀ O	[174155-57-8]	(E) 3-hexadecenal				
	$\Delta_v H$	(373–413)	89.6	298	CGC	[1996KOU/HOS, 2000OVA/KOU]
C ₁₆ H ₃₀ O	[88373-69-7]	(Z) 4-hexadecenal				
	$\Delta_v H$	(373–413)	88.7	298	CGC	[1996KOU/HOS, 2000OVA/KOU]
C ₁₆ H ₃₀ O	[174155-59-0]	(E) 4-hexadecenal				
	$\Delta_v H$	(373–413)	88.9	298	CGC	[1996KOU/HOS, 2000OVA/KOU]
C ₁₆ H ₃₀ O	[88373-68-6]	(Z) 5-hexadecenal				
	$\Delta_v H$	(373–413)	87.8	298	CGC	[1996KOU/HOS, 2000OVA/KOU]
C ₁₆ H ₃₀ O	[99142-11-7]	(E) 5-hexadecenal				
	$\Delta_v H$	(373–413)	88.6	298	CGC	[1996KOU/HOS, 2000OVA/KOU]
C ₁₆ H ₃₀ O	[88373-67-5]	(Z) 6-hexadecenal				
	$\Delta_v H$	(373–413)	87.9	298	CGC	[1996KOU/HOS, 2000OVA/KOU]
C ₁₆ H ₃₀ O	[103346-18-5]	(E) 6-hexadecenal				
	$\Delta_v H$	(373–413)	88.5	298	CGC	[1996KOU/HOS, 2000OVA/KOU]
C ₁₆ H ₃₀ O	[56797-40-1]	(Z) 7-hexadecenal				
	$\Delta_v H$	(373–413)	87.8	298	CGC	[1996KOU/HOS, 2000OVA/KOU]
C ₁₆ H ₃₀ O	[72698-27-2]	(E) 7-hexadecenal				
	$\Delta_v H$	(373–413)	88.6	298	CGC	[1996KOU/HOS, 2000OVA/KOU]
C ₁₆ H ₃₀ O	[66644-98-2]	(Z) 8-hexadecenal				
	$\Delta_v H$	(373–413)	87.7	298	CGC	[1996KOU/HOS, 2000OVA/KOU]
C ₁₆ H ₃₀ O	[72698-28-3]	(E) 8-hexadecenal				
	$\Delta_v H$	(373–413)	88.4	298	CGC	[1996KOU/HOS, 2000OVA/KOU]
C ₁₆ H ₃₀ O	[56219-04-6]	(Z) 9-hexadecenal				
	$\Delta_v H$	(373–413)	88.0	298	CGC	[1996KOU/HOS, 2000OVA/KOU]

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
	Enthalpy	Temp (K) Range					
C ₁₆ H ₃₀ O	[72698-29-4] $\Delta_{\text{v}}H$	(E) 9-hexadecenal (373–413)	88.6	298	CGC	[1996KOU/HOS, 2000OVA/KOU]	
C ₁₆ H ₃₀ O	[68279-24-3] $\Delta_{\text{v}}H$	(Z) 10-hexadecenal (373–413)	88.2	298	CGC	[1996KOU/HOS, 2000OVA/KOU]	
C ₁₆ H ₃₀ O	[72698-30-7] $\Delta_{\text{v}}H$	(E) 10-hexadecenal (373–413)	88.8	298	CGC	[1996KOU/HOS, 2000OVA/KOU]	
C ₁₆ H ₃₀ O	[53939-28-9] $\Delta_{\text{v}}H$	(Z) 11-hexadecenal (373–413)	88.5	298	CGC	[1996KOU/HOS, 2000OVA/KOU]	
C ₁₆ H ₃₀ O	[57491-33-5] $\Delta_{\text{v}}H$	(E) 11-hexadecenal (373–413)	89.2	298	CGC	[1996KOU/HOS, 2000OVA/KOU]	
C ₁₆ H ₃₀ O	[72698-31-8] $\Delta_{\text{v}}H$	(Z) 12-hexadecenal (373–413)	89.3	298	CGC	[1996KOU/HOS, 2000OVA/KOU]	
C ₁₆ H ₃₀ O	[72698-32-9] $\Delta_{\text{v}}H$	(E) 12-hexadecenal (373–413)	89.3	298	CGC	[1996KOU/HOS, 2000OVA/KOU]	
C ₁₆ H ₃₀ O	[71545-96-5] $\Delta_{\text{v}}H$	(Z) 13-hexadecenal (373–413)	89.7	298	CGC	[1996KOU/HOS, 2000OVA/KOU]	
C ₁₆ H ₃₀ O	[72698-33-0] $\Delta_{\text{v}}H$	(E) 13-hexadecenal (373–413)	90.0	298	CGC	[1996KOU/HOS, 2000OVA/KOU]	
C ₁₆ H ₃₀ O ₂	[142-90-5] $\Delta_{\text{v}}H$	dodecyl methacrylate (438–580)	64.9	453	A	[1987STE/MAL]	
C ₁₆ H ₃₀ O ₂	[109-29-5] $\Delta_{\text{v}}H$	oxa-2-cycloheptadecanone (403–463)	71.6	418	A	[1987STE/MAL]	
C ₁₆ H ₃₀ O ₂	[51309-20-7] $\Delta_{\text{v}}H$	(Z) 2-tetradecenyl acetate (353–398)	89.1	298	GC	[1997KOU/HOS, 2000OVA/KOU]	
C ₁₆ H ₃₀ O ₂	[51309-21-8] $\Delta_{\text{v}}H$	(E) 2-tetradecenyl acetate (353–398)	90.3	298	GC	[1997KOU/HOS, 2000OVA/KOU]	
C ₁₆ H ₃₀ O ₂	[54897-65-3] $\Delta_{\text{v}}H$	(Z) 3-tetradecenyl acetate (353–398)	88.5	298	GC	[1997KOU/HOS, 2000OVA/KOU]	
C ₁₆ H ₃₀ O ₂	[56221-90-0] $\Delta_{\text{v}}H$	(E) 3-tetradecenyl acetate (353–398)	89.2	298	GC	[1997KOU/HOS, 2000OVA/KOU]	
C ₁₆ H ₃₀ O ₂	[54897-66-4] $\Delta_{\text{v}}H$	(Z) 4-tetradecenyl acetate (353–398)	87.8	298	GC	[1997KOU/HOS, 2000OVA/KOU]	
C ₁₆ H ₃₀ O ₂	[56209-67-7] $\Delta_{\text{v}}H$	(E) 4-tetradecenyl acetate (353–398)	89.0	298	GC	[1997KOU/HOS, 2000OVA/KOU]	
C ₁₆ H ₃₀ O ₂	[35153-13-0] $\Delta_{\text{v}}H$	(Z) 5-tetradecenyl acetate (353–398)	88.3	298	GC	[1997KOU/HOS, 2000OVA/KOU]	
C ₁₆ H ₃₀ O ₂	[34010-13-4] $\Delta_{\text{v}}H$	(E) 5-tetradecenyl acetate (353–398)	89.1	298	GC	[1997KOU/HOS, 2000OVA/KOU]	
C ₁₆ H ₃₀ O ₂	[39650-11-8] $\Delta_{\text{v}}H$	(Z) 6-tetradecenyl acetate (353–398)	88.1	298	GC	[1997KOU/HOS, 2000OVA/KOU]	
C ₁₆ H ₃₀ O ₂	[39650-10-7] $\Delta_{\text{v}}H$	(E) 6-tetradecenyl acetate (353–398)	88.9	298	GC	[1997KOU/HOS, 2000OVA/KOU]	

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		T _m (K)	Method	Reference
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)			
C ₁₆ H ₃₀ O ₂	[16974-10-0]	(Z) 7-tetradecenyl acetate	88.4	298	GC	[1997KOU/HOS, 2000OVA/KOU]
	$\Delta_{\text{v}}H$	(353–398)				
C ₁₆ H ₃₀ O ₂	[28540-79-6]	(E) 7-tetradecenyl acetate	89.0	298	GC	[1997KOU/HOS, 2000OVA/KOU]
	$\Delta_{\text{v}}H$	(353–398)				
C ₁₆ H ₃₀ O ₂	[35835-80-4]	(Z) 8-tetradecenyl acetate	88.7	298	GC	[1997KOU/HOS, 2000OVA/KOU]
	$\Delta_{\text{v}}H$	(353–398)				
C ₁₆ H ₃₀ O ₂	[56218-64-5]	(E) 8-tetradecenyl acetate	89.2	298	GC	[1997KOU/HOS, 2000OVA/KOU]
	$\Delta_{\text{v}}H$	(353–398)				
C ₁₆ H ₃₀ O ₂	[16725-53-4]	(Z) 9-tetradecenyl acetate	89.1	298	GC	[1997KOU/HOS, 2000OVA/KOU]
	$\Delta_{\text{v}}H$	(353–398)				
	$\Delta_{\text{v}}H$	(303–317)	90.0	310	GC	[1983OLS/JON]
C ₁₆ H ₃₀ O ₂	[23192-82-7]	(E) 9-tetradecenyl acetate	89.6	298	GC	[1997KOU/HOS, 2000OVA/KOU]
	$\Delta_{\text{v}}H$	(353–398)				
C ₁₆ H ₃₀ O ₂	[35153-16-3]	(Z) 10-tetradecenyl acetate	89.6	298	GC	[1997KOU/HOS, 2000OVA/KOU]
	$\Delta_{\text{v}}H$	(353–398)				
C ₁₆ H ₃₀ O ₂	[35153-17-4]	(E) 10-tetradecenyl acetate	89.9	298	GC	[1997KOU/HOS, 2000OVA/KOU]
	$\Delta_{\text{v}}H$	(353–398)				
C ₁₆ H ₃₀ O ₂	[20711-10-8]	(Z) 11-tetradecenyl acetate	90.0	298	GC	[1997KOU/HOS, 2000OVA/KOU]
	$\Delta_{\text{v}}H$	(353–398)				
C ₁₆ H ₃₀ O ₂	[33189-72-9]	(E) 11-tetradecenyl acetate	90.4	298	GC	[1997KOU/HOS, 2000OVA/KOU]
	$\Delta_{\text{v}}H$	(353–398)				
C ₁₆ H ₃₀ O ₂	[35153-20-9]	(Z) 12-tetradecenyl acetate	90.9	298	GC	[1997KOU/HOS, 2000OVA/KOU]
	$\Delta_{\text{v}}H$	(353–398)				
C ₁₆ H ₃₀ O ₂	[35153-21-0]	(E) 12-tetradecenyl acetate	90.8	298	GC	[1997KOU/HOS, 2000OVA/KOU]
	$\Delta_{\text{v}}H$	(353–398)				
C ₁₆ H ₃₀ O ₂	[90176-51-5]	methyl Z 10-pentadecenoate	91.7	298	CGC	[2007LIP/KAP]
	$\Delta_{\text{v}}H$					
C ₁₆ H ₃₀ O ₂	[373-49-9]	<i>cis</i> -9-hexadecenoic acid (palmitoleic acid)				
	$\Delta_{\text{trs}}H$		7.5	254.8		
	$\Delta_{\text{fus}}H$		32.1	275.2		[1997SAT/YAN]
C ₁₆ H ₃₀ O ₃	[6720-22-5]	1,7-dioxa-8-cyclooctadecanone	73.3	418	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$	(403–463)				
C ₁₆ H ₃₀ O ₃	[36575-58-3]	1,9-dioxa-2-cyclooctadecanone	74.5	418	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$	(403–463)				
C ₁₆ H ₃₀ O ₄	[14027-78-2]	dipentyl adipate	74.7	464	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$	(449–575)				
C ₁₆ H ₃₀ O ₄	[505-54-4]	hexadecanedioic acid				
	$\Delta_{\text{fus}}H + \Delta_{\text{trs}}H$		60.1	396.4	DSC	[2006VEN/MET]
	$\Delta_{\text{fus}}H$		52.2	395.4	DSC	[2005ROU/TEM]
	$\Delta_{\text{sub}}H$	(377–398)	151.0 ± 3.3	388	ME	[1960DAV/THO, 1987STE/MAL]
	$\Delta_{\text{sub}}H$		155.4 ± 3.3	298		[1960DAV/THO, 1999RIB/MON]
C ₁₆ H ₃₀ O ₅	[5420-72-4]	octyl[1-(butoxycarbonyl)ethyl]carbonate	76.2	389	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$	(374–503)				

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		T _m (K)	Method	Reference
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)			
C ₁₆ H ₃₁ N	[629-79-8]	hexadecanenitrile (palmitonitrile)				
	$\Delta_{\text{v}}H$	(345–382)	93.3 ± 0.4	298		[2005EME/VER]
	$\Delta_{\text{v}}H$	(503–608)	70.1	518	A	[1987STE/MAL]
C ₁₆ H ₃₁ NO ₃	[14246-55-0]	N-tetradecanoylglycine				
	$\Delta_{\text{trs}}H$		6.8	379.6		
	$\Delta_{\text{fus}}H$		47.4	396.6	DSC	[1986MIY/MAT]
C ₁₆ H ₃₁ NO ₃	[14379-38-5]	N-decanoyl-(l)-leucine				
	$\Delta_{\text{trs}}H$		1.2	343.1		
	$\Delta_{\text{fus}}H$		27.5	383.1	DSC	[1986MIY/MAT]
C ₁₆ H ₃₁ NO ₃	[107396-12-3]	N-decanoyl-(dl)-leucine				
	$\Delta_{\text{fus}}H$		28.9	357.1	DSC	[1986MIY/MAT]
C ₁₆ H ₃₂	[15220-85-6]	tetraisobutylene				
	$\Delta_{\text{v}}H$	(381–440)	54.5	397		[1943STE]
C ₁₆ H ₃₂	[1795-16-0]	decylcyclohexane				
	$\Delta_{\text{fus}}H$		38.62	271.4		[1991ACR]
	$\Delta_{\text{v}}H$	(371–425)	76.7	386	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$		79.7	298		[1971WIL/ZWO]
	$\Delta_{\text{v}}H$	(469–571)	61.6	484	A, MM	[1987STE/MAL, 1954CAM/FOR]
C ₁₆ H ₃₂	[6785-23-5]	undecylcyclopentane				
	$\Delta_{\text{v}}H$		80.6	298		[1971WIL/ZWO]
C ₁₆ H ₃₂	[629-73-2]	1-hexadecene				
	$\Delta_{\text{trs}}H$		3.87	249.2		
	$\Delta_{\text{fus}}H$		30.21	277.5		[1990MES/TOD]
	$\Delta_{\text{v}}H$		80.3 ± 0.4	298	C	[1977MAN/SEL]
	$\Delta_{\text{v}}H$		80.3 ± 0.4	298	C	[1976STR2]
	$\Delta_{\text{v}}H$		80.1	298		[1971WIL/ZWO]
	$\Delta_{\text{v}}H$	(461–558)	61.5	476	A	[1987STE/MAL, 1954CAM/FOR]
C ₁₆ H ₃₂	[295-65-8]	cyclohexadecane				
	$\Delta_{\text{trs}}H$		18.83	271.2		
	$\Delta_{\text{trs}}H$		1.26	283.2		
	$\Delta_{\text{fus}}H$		4.18	332.2		[1975BJO/BOR2]
	$\Delta_{\text{sub}}H$		81.8 ± 0.4			[1957VAN, 1970COX/PIL]
C ₁₆ H ₃₂ O	[141694-91-9]	(Z) 3-hexadecen-1-ol				
	$\Delta_{\text{v}}H$	(373–413)	110.7	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₆ H ₃₂ O	[128999-42-8]	(E) 3-hexadecen-1-ol				
	$\Delta_{\text{v}}H$	(373–413)	110.8	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₆ H ₃₂ O	[145235-63-8]	(Z) 4-hexadecen-1-ol				
	$\Delta_{\text{v}}H$	(373–413)	110.6	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₆ H ₃₂ O	[59101-23-4]	(E) 4-hexadecen-1-ol				
	$\Delta_{\text{v}}H$	(373–413)	111.5	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₆ H ₃₂ O	[106463-48-3]	(Z) 5-hexadecen-1-ol				
	$\Delta_{\text{v}}H$	(373–413)	110.9	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₆ H ₃₂ O	[85388-16-5]	(E) 5-hexadecen-1-ol				
	$\Delta_{\text{v}}H$	(373–413)	111.4	298	CGC	[2000OVA/KOU, 1994KOU/HOS]

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
C ₁₆ H ₃₂ O	[40642-45-3] $\Delta_{\text{v}}H$	(Z) 6-hexadecen-1-ol (373–413)	110.5	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₆ H ₃₂ O	[34500-33-9] $\Delta_{\text{v}}H$	(E) 6-hexadecen-1-ol (373–413)	111	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₆ H ₃₂ O	[24880-48-6] $\Delta_{\text{v}}H$	(Z) 7-hexadecen-1-ol (373–413)	110.2	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₆ H ₃₂ O	[51824-10-3] $\Delta_{\text{v}}H$	(E) 7-hexadecen-1-ol (373–413)	111.4	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₆ H ₃₂ O	[64437-46-3] $\Delta_{\text{v}}H$	(Z) 8-hexadecen-1-ol (373–413)	110.4	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₆ H ₃₂ O	[64470-33-3] $\Delta_{\text{v}}H$	(E) 8-hexadecen-1-ol (373–413)	111.1	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₆ H ₃₂ O	[10378-01-5] $\Delta_{\text{v}}H$	(Z) 9-hexadecen-1-ol (373–413)	110.6	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₆ H ₃₂ O	[64437-47-4] $\Delta_{\text{v}}H$	(E) 9-hexadecen-1-ol (373–413)	111.3	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₆ H ₃₂ O	[64437-48-5] $\Delta_{\text{v}}H$	(Z) 10-hexadecen-1-ol (373–413)	111	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₆ H ₃₂ O	[54502-94-2] $\Delta_{\text{v}}H$	(E) 10-hexadecen-1-ol (373–413)	111.5	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₆ H ₃₂ O	[56683-54-6] $\Delta_{\text{v}}H$	(Z) 11-hexadecen-1-ol (373–413)	111.3	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₆ H ₃₂ O	[61301-56-2] $\Delta_{\text{v}}H$	(E) 11-hexadecen-1-ol (373–413)	111.8	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₆ H ₃₂ O	[72698-34-1] $\Delta_{\text{v}}H$	(Z) 12-hexadecen-1-ol (373–413)	111.8	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₆ H ₃₂ O	[72698-35-2] $\Delta_{\text{v}}H$	(E) 12-hexadecen-1-ol (373–413)	112.1	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₆ H ₃₂ O	[69282-65-1] $\Delta_{\text{v}}H$	(Z) 13-hexadecen-1-ol (373–413)	112.3	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₆ H ₃₂ O	[69282-66-2] $\Delta_{\text{v}}H$	(E) 13-hexadecen-1-ol (373–413)	112.6	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₆ H ₃₂ O	[18787-63-8] $\Delta_{\text{v}}H$	2-hexadecanone (382–580)	72.3	397	A	[1987STE/MAL]
C ₁₆ H ₃₂ O	[629-80-1] $\Delta_{\text{v}}H$ $\Delta_{\text{v}}H$	hexadecanal (343–383) (394–594)	89.7 67.6	298 409	CGC A	[1996KOU/HOS, 2000OVA/KOU] [1987STE/MAL, 1947STU]
C ₁₆ H ₃₂ O ₂	[7132-64-1] $\Delta_{\text{v}}H$ $\Delta_{\text{v}}H$ $\Delta_{\text{v}}H$ $\Delta_{\text{v}}H$ $\Delta_{\text{v}}H$ $\Delta_{\text{v}}H$	methyl pentadecanoate (433–473)	82.1 79.8 ± 0.2 89.3 ± 0.8 88.8 91.6 ± 0.9 93.5 ± 1.0	350 372 298 298 298 298	 CGC GC,C C	[2002SEG/GAL] [2002SEG/GAL] [2002SEG/GAL] [1995CHI/HOS] [1980FUC/PEA] [1977MAN/SEL]

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference	
	Enthalpy								
		$\Delta_v H$	(295–303)	87.9 ± 1.3	299			[1968BAC/NOV]	
		$\Delta_v H$	(400–527)	78.3	415	A, E		[1987STE/MAL, 1963ROS/SCH]	
C ₁₆ H ₃₂ O ₂	[124-06-1]	ethyl tetradecanoate							
		$\Delta_v H$	(407–568)	71.8	422	A		[1987STE/MAL]	
C ₁₆ H ₃₂ O ₂	[106-18-3]	butyl dodecanoate							
		$\Delta_v H$	(423–483)	89.2	298	GC		[1997KRO/VEL]	
		$\Delta_v H$	(343–383)	75.8	358	A		[1987STE/MAL]	
C ₁₆ H ₃₂ O ₂	[na]	isobutyl dodecanoate							
		$\Delta_v H$	(345–452)	80.0	360			[2001BUR/JOS]	
C ₁₆ H ₃₂ O ₂	[638-59-5]	tetradecyl acetate							
		$\Delta_v H$	(303–340)	89.9 ± 0.2	298	GS		[2006KRA/VER]	
		$\Delta_v H$	(353–398)	91.7	298	GC		[1997KOU/HOS, 2000OVA/KOU]	
		$\Delta_v H$	(411–462)	72.7	426	A		[1987STE/MAL]	
C ₁₆ H ₃₂ O ₂	[57-10-3]	hexadecanoic acid (palmitic acid)							
		$\Delta_{\text{trs}}H$		3.8	318.8				
		$\Delta_{\text{fus}}H$		53.9	335.4	DSC		[2009GBA/NEG]	
		$\Delta_{\text{fus}}H$		51.37	332.7	DSC		[2009ZEN/CAO]	
		$\Delta_{\text{trs}}H$		3.1	316.7				
		$\Delta_{\text{trs}}H$		4.9	317.5				
		$\Delta_{\text{fus}}H$		53.0	334.7	DSC		[2007MOR/COR]	
		$\Delta_{\text{fus}}H$		47.0	336.5	DSC		[2007MIS/MIS]	
		$\Delta_{\text{fus}}H$		54.81	335.7			[1996DOM/HEA]	
		$\Delta_{\text{sub}}H$		193.8 ± 11	298	TPD		[2008CAP/LOV]	
		$\Delta_{\text{sub}}H$	(273–303)	134		TPTD		[2005CHA/ZIE]	
		$\Delta_{\text{sub}}H$	(294–316)	154		TPTD		[2001CHA/TOB]	
		Note: Experimental values based on the TPTD method are often inconsistent with values determined using other experimental methods							
		$\Delta_{\text{sub}}H$	(320–333)	154.4 ± 4.2	326	ME			[1961DAV/MAL, 1970COX/PIL, 1987STE/MAL]
	$\Delta_v H$	(440–625)	97.5	455	A			[1987STE/MAL]	
	$\Delta_v H$	(347–374)	110.2 ± 2.0	364	ME, TE			[1982DEK/SCH]	
	$\Delta_v H$		90.1	475	I			[1943CRA]	
C ₁₆ H ₃₂ O ₃	[764-67-0]	2-hydroxyhexadecanoic acid							
		$\Delta_{\text{sub}}H$	(294–311)	121		TPTD		[2005CHA/ZIE]	
	Note: Experimental values based on the TPTD method are often inconsistent with values determined using other experimental methods								
C ₁₆ H ₃₂ O ₃	[506-13-8]	16-hydroxyhexadecanoic acid							
		$\Delta_{\text{sub}}H$	(316–329)	114		TPTD		[2005CHA/ZIE]	
	Note: Experimental values based on the TPTD method are often inconsistent with values determined using other experimental methods								
C ₁₆ H ₃₂ O ₄	[43091-27-6]	6,6,14,14-tetramethyl-1,3,9,11-tetraoxacyclohexadecane							
		$\Delta_{\text{fus}}H$		29.71	358.6			[1973DAL/EKE]	
C ₁₆ H ₃₂ O ₄	[43091-28-7]	2,2,10,10-tetramethyl-1,3,9,11-tetraoxacyclohexadecane							
		$\Delta_{\text{fus}}H$		25.94	371.3			[1973DAL/EKE]	
C ₁₆ H ₃₂ O ₈	[33089-37-1]	1,4,7,10,13,16,19,22-octaoxacyclotetracosane							
		$\Delta_{\text{fus}}H$		34.5	292.2			[1972DAL/KRI]	
C ₁₆ H ₃₃ Br	[112-82-3]	1-bromohexadecane							
		$\Delta_v H$		94.4 ± 1.5	298	C		[1996WEB/DEF2]	

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		T _m (K)	Method	Reference
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)			
C ₁₆ H ₃₃ Cl		(461–673)	71.9	476	A, E	[1987STE/MAL, 1961LI/ROS, 1970DYK/VAN]
	[4860-03-1]	1-chlorohexadecane				
	$\Delta_{\text{v}}H$		97.9	298		[2006BOL/NER2]
	$\Delta_{\text{v}}H$		96.4 ± 0.9	298	GS	[2001PUR/CHI]
	$\Delta_{\text{v}}H$	(439–600)	91.8 ± 1.1	298	C	[1977MAN/SEL]
C ₁₆ H ₃₃ F	[408-38-8]	1-fluorohexadecane				
	$\Delta_{\text{v}}H$	(425–608)	66.1	440	A, E	[1970DYK/VAN]
C ₁₆ H ₃₃ I	[544-77-4]	1-iodohexadecane				
	$\Delta_{\text{v}}H$	(475–673)	99.6	298	A,E	[1987STE/MAL, 1961LI/ROS, 1970DYK/VAN, 2006BOL/NER]
C ₁₆ H ₃₃ NO		(475–673)	73.0	490	A, E	[1987STE/MAL, 1961LI/ROS, 1970DYK/VAN]
	[629-54-9]	hexadecanamide				
	$\Delta_{\text{trs}}H$		10.4	355.5		
	$\Delta_{\text{fus}}H$		45.4	376	DSC	[2008ABA/BAD]
C ₁₆ H ₃₃ NO		(364–378)	181.6 ± 1.3	371	ME	[1959DAV/JON2, 1987STE/MAL]
	[74534-10-4]	N-hexyl decanamide				
	$\Delta_{\text{trs}}H$		6.0	301		
C ₁₆ H ₃₃ NO			31.0	311	DSC	[1980CAR/BUS]
	[6284-08-8]	N-butyl dodecanamide				
	$\Delta_{\text{fus}}H$		39.0	322.1	DSC	[1980CAR/BUS]
C ₁₆ H ₃₃ NO	[57303-23-8]	N,N-dibutyl octanamide				
	$\Delta_{\text{v}}H$	(463–513)	75.6 ± 0.7	298	CGC	[2009PAN/ANT]
C ₁₆ H ₃₄	[544-76-3]	hexadecane				
	$\Delta_{\text{fus}}H$		53.0	290.7	DSC	[2004MON/RAJ]
	$\Delta_{\text{fus}}H$		53.35	291.3		
	$\Delta_{\text{fus}}H$		51.46	291.1		[1996DOM/HEA]
	$\Delta_{\text{sub}}H$		135.1	298	B	[1972MOR3]
	$\Delta_{\text{sub}}H$		134.9	291	B	[1963BON]
	$\Delta_{\text{sub}}H$	(288–290)	83.4 ± 8		ME	[1949BRA/SHE]
	$\Delta_{\text{v}}H$		81.8 ± 1.3	298	CGC	[2000NIC/ORF]
	$\Delta_{\text{v}}H$	(453–503)	81.4	298	CGC	[1995CHI/HOS]
	$\Delta_{\text{v}}H$	(423–473)	81.4	298	CGC	[1995CHI/HOS]
	$\Delta_{\text{v}}H$	(363–413)	81.2	298	CGC	[1995CHI/HOS]
	$\Delta_{\text{v}}H$	(393–583)	68.5	408		[1994MOR/KOB]
	$\Delta_{\text{v}}H$		81.4	298		[1994RUZ/MAJ]
	$\Delta_{\text{v}}H$	(505–589)	59.8	520		[1992LEE/DEM]
	$\Delta_{\text{v}}H$	(323–423)	74.9	338	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$		66.9	343	GC	[1977NOV/NOV]
	$\Delta_{\text{v}}H$		66.2	353	GC	[1977NOV/NOV]
	$\Delta_{\text{v}}H$		65.6	363	GC	[1977NOV/NOV]
	$\Delta_{\text{v}}H$		64.9	373	GC	[1977NOV/NOV]
	$\Delta_{\text{v}}H$		64.2	383	GC	[1977NOV/NOV]
	$\Delta_{\text{v}}H$		81.4 ± 0.4	298	C	[1972MOR2]
	$\Delta_{\text{v}}H$		81.1	298		[1971WIL/ZWO]
	$\Delta_{\text{v}}H$	(467–563)	61.7	482	A, MM	[1987STE/MAL, 1954CAM/FOR]
$\Delta_{\text{v}}H$	(299–324)	93.4	311	ME	[1949PAR/MOO]	
$\Delta_{\text{v}}H$	(293–308)	80.2	300	ME	[1949BRA/SHE2]	
$\Delta_{\text{v}}H$	(442–469)	65.7	455	ME	[1938UBB]	

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		T _m (K)	Method	Reference
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)			
C ₁₆ H ₃₄	[1560-93-6]	2-methylpentadecane				
	$\Delta_{\text{v}}H$	(417–554)	62.0	432	A	[1987STE/MAL]
C ₁₆ H ₃₄	[2882-96-4]	3-methylpentadecane				
	$\Delta_{\text{v}}H$	(417–555)	61.0	432	A	[1987STE/MAL]
C ₁₆ H ₃₄	[2801-87-8]	4-methylpentadecane				
	$\Delta_{\text{v}}H$	(411–553)	57.8	426	A	[1987STE/MAL]
C ₁₆ H ₃₄	[25117-33-3]	5-methylpentadecane				
	$\Delta_{\text{v}}H$	(408–551)	57.3	423	A	[1987STE/MAL]
C ₁₆ H ₃₄	[6165-40-8]	7-methylpentadecane				
	$\Delta_{\text{v}}H$	(355–410)	66.3	370	A	[1987STE/MAL]
C ₁₆ H ₃₄	[18435-23-9]	2,3-dimethyltetradecane				
	$\Delta_{\text{v}}H$	(412–554)	57.4	427	A	[1987STE/MAL]
C ₁₆ H ₃₄	[61868-06-2]	2,4-dimethyltetradecane				
	$\Delta_{\text{v}}H$	(404–539)	60.6	419	A	[1987STE/MAL]
C ₁₆ H ₃₄	[na]	2,4,6-trimethyltridecane				
	$\Delta_{\text{v}}H$	(395–521)	59.1	410	A	[1987STE/MAL]
C ₁₆ H ₃₄	[4390-04-9]	2,2,4,4,6,8,8-heptamethylnonane				
	$\Delta_{\text{v}}H$	(423–545)	52.4	438		[1988AMB/GHI]
C ₁₆ H ₃₄	[78715-64-7]	3,3,6,6-tetraethyloctane				
	$\Delta_{\text{v}}H$	(301–330)	73.0 ± 1.9	308	HSA	[1995CHI/HES]
	$\Delta_{\text{v}}H$		74.3 ± 1.9	298		[1995CHI/HES]
	$\Delta_{\text{v}}H$		72.3 ± 1.8	298	CGC	[1995CHI/HES]
C ₁₆ H ₃₄ N ₂	[39198-34-0]	bis(1,1,3,3-tetramethylbutyl)diazene				
	$\Delta_{\text{v}}H$		66.5 ± 0.6	298	C	[1976ENG/MEL]
C ₁₆ H ₃₄ O	[36653-82-4]	1-hexadecanol				
	$\Delta_{\text{fus}}H$	(80–370)	57.7	322.2	AC	[2008XIN/TAN]
	$\Delta_{\text{fus}}H$		33.1	321.6	DSC	[2004VEN/CAL]
	$\Delta_{\text{trs}}H$		21.21	322.2		[1979KUC/SKU]
	$\Delta_{\text{fus}}H$		33.97	322.9		[1979KUC/SKU]
	$\Delta_{\text{fus}}H + \Delta_{\text{trs}}H$		58.41	322.2		[1974MOS/MOU]
	$\Delta_{\text{sub}}H$	(308–320)	167.4 ± 2.1	314	ME	[1965DAV/KYB, 1987STE/MAL]
	$\Delta_{\text{sub}}H$		169.5 ± 2.1	298		[1965DAV/KYB]
	$\Delta_{\text{v}}H$		107.7 ± 1.2	298	CGC	[2006NIC/KWE]
	$\Delta_{\text{v}}H$	(328–362)	100.4	347	GS	[2001KUL/VER2]
	$\Delta_{\text{v}}H$	(328–362)	108.8	298	GS	[2001KUL/VER2]
	$\Delta_{\text{v}}H$		112.5	298	CGC	[2000OVA/KOU]
	$\Delta_{\text{v}}H$	(343–463)	88.2	403		[1992NGU/KAS]
	$\Delta_{\text{v}}H$	(509–569)	68.9	524	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$	(415–487)	83.2	430	A	[1987STE/MAL, 1974AMB/ELL]
	$\Delta_{\text{v}}H$	(323–335)	109.4	329		[1987STE/MAL]
	$\Delta_{\text{v}}H$	(323–376)	112.3	338		[1973WIL/ZWO]
	$\Delta_{\text{v}}H$	(418–463)	78.8	423		[1973WIL/ZWO]
	$\Delta_{\text{v}}H$	(498–569)	70.0	513	A, EB	[1987STE/MAL, 1970AMB/SPR]
	$\Delta_{\text{v}}H$	(445–598)	77.3	460	DTA	[1969KEM/KRE]
	$\Delta_{\text{v}}H$	(323–335)	109.5	329	ME	[1965DAV/KYB]
C ₁₆ H ₃₄ O	[14852-31-4]	2-hexadecanol				
	$\Delta_{\text{v}}H$	(333–453)	102.2	348		[1999NGU/BER]

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		T _m (K)	Method	Reference
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)			
C ₁₆ H ₃₄ O ₂	[7735-42-4] $\Delta_{\text{fus}}H$	1,16-hexadecanediol	64.2	365.4	DSC	[1999OGA/NAK]
C ₁₆ H ₃₄ O ₂ S	[126835-77-6] $\Delta_{\text{trs}}H$ $\Delta_{\text{fus}}H$	3-(tridecylthio)-1,2-propanediol	11.3 22.7	296.9 330.6	DSC	[1993ACR]
C ₁₆ H ₃₄ O ₃	[10431-00-2] $\Delta_{\text{fus}}H$	3-(tridecyloxy)-1,2-propanediol	51.4	324.2	DSC	[1993ACR]
C ₁₆ H ₃₄ O ₃	[3055-93-4] $\Delta_{\text{v}}H$	2[2-(dodecyloxy)ethoxy]ethanol (448–489)	82.1	463	A	[1987STE/MAL]
C ₁₆ H ₃₄ O ₄ S ₂	[na] $\Delta_{\text{fus}}H$	2-deoxy-(D)-glucose dipentyl dithioacetal	63.1	393.3	DSC	[1989VAN/VAN]
C ₁₆ H ₃₄ O ₄ S ₂	[na] $\Delta_{\text{fus}}H$	(l)-rhamnose dipentyl dithioacetal	46.5	388.2	DSC	[1989VAN/VAN]
C ₁₆ H ₃₄ O ₅ S ₂	[115395-53-4] $\Delta_{\text{fus}}H$	(D)-glucose dipentyl dithioacetal	49.1	389	DSC	[1989VAN/VAN]
C ₁₆ H ₃₄ O ₅ S ₂	[123389-86-6] $\Delta_{\text{trs}}H$ $\Delta_{\text{fus}}H$	(D)-galactose dipentyl dithioacetal	2.7 41.1	384.6 392.1	DSC	[1989VAN/VAN]
C ₁₆ H ₃₄ S	[2917-26-2] $\Delta_{\text{v}}H$	1-hexadecanethiol (470–643)	72.4	485		[1999DYK/SVO]
C ₁₆ H ₃₄ S	[2690-08-6] $\Delta_{\text{v}}H$ $\Delta_{\text{v}}H$ $\Delta_{\text{v}}H$ $\Delta_{\text{v}}H$	dioctyl sulfide (335–442) (465–550) (465–550) (465–550)	71.6 95.0 ± 10.7 72.0 ± 0.6 67.8 ± 0.5	388 298 460 500	EB EB EB EB	[2004SAW/MOK] [1997STE/CHI4] [1997STE/CHI4] [1997STE/CHI4] [1997STE/CHI4]
C ₁₆ H ₃₄ S ₂	[822-27-5] $\Delta_{\text{v}}H$	diheptyl disulfide (479–656)	73.9	494		[1999DYK/SVO]
C ₁₆ H ₃₅ N	[1120-48-5] $\Delta_{\text{v}}H$	dioctylamine (448–597)	87.1 ± 1.3	298	EB	[1996STE/CHI3]
C ₁₆ H ₃₅ N	[143-27-1] $\Delta_{\text{v}}H$	hexadecylamine (498–609)	66.9	513	A	[1987STE/MAL]
C ₁₆ H ₃₅ N	[na] $\Delta_{\text{v}}H$	N,N-dimethyl-2-pentylnonylamine (401–552)	64.8	425	EB	[1987MIL/FEN2]
C ₁₆ H ₃₅ NO ₂	[126835-68-5] $\Delta_{\text{fus}}H$	3-(tridecylamino)-1,2-propanediol	68.7	354.9	DSC	[1993ACR]
C ₁₆ H ₃₆ N ₂	[60678-70-8] $\Delta_{\text{v}}H$	tetraethyl hydrazine (392–453)	51.1	407	A	[1987STE/MAL]
C ₁₇ H ₁₀ ClN ₃ O ₃	[1978-90-2] $\Delta_{\text{fus}}H$	1-(4-chlorobenzoyl)-1,2-dihydro-6-nitro-1-quinolinecarbonitrile	25.62	430.5	DSC	[2005LIZ/ZAB]
C ₁₇ H ₁₀ O	[82-05-3] $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$	benzanthrone (390–410) (390–410) (389–409) (389–409)	122.6 ± 0.6 125.6 ± 0.6 121.6 ± 0.6 126.6 ± 0.6	400 298 399 298	ME ME ME ME	[2006RIB/MON] [2006RIB/MON] [1999RIB/FER] [1999RIB/FER]

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
	$\Delta_{\text{sub}}H$	(373–393)	129.7 ± 2.1	298	QR	[1999RIB/FER]
	$\Delta_{\text{sub}}H$	(373–393)	125.5 ± 2.1	382	QR	[1999RIB/FER]
	$\Delta_{\text{sub}}H$	(353–388)	119.7 ± 5.4	370	ME	[1984BUR/MOR]
	$\Delta_{\text{sub}}H$	(353–388)	124.6 ± 6.0	298	ME	[1984BUR/MOR]
	$\Delta_{\text{sub}}H$		114.2 ± 0.8		QR	[1979YAN/TEP]
	$\Delta_{\text{sub}}H$		115.5	398		[1952INO/SHI, 1960JON]
	$\Delta_{\text{v}}H$	(498–673)	91.4	513	A	[1987STE/MAL, 1947STU]
C ₁₇ H ₁₀ O	[116232-62-3]	benzo[a]fluorenone				
	$\Delta_{\text{fus}}H$		15.6	406.9	DSC	[2010KES/AUC]
C ₁₇ H ₁₁ ClN ₂ O ₂	[1979-25-6]	2-cyano-1(2 <i>H</i>)-quinolinecarboxylic acid, 4-chlorophenyl ester				
	$\Delta_{\text{fus}}H$		37.19	424.2	DSC	[2005LIZ/ZAB]
C ₁₇ H ₁₁ N	[225-11-6]	benz[a]acridine				
	$\Delta_{\text{fus}}H$		21.9	402.8	DSC	[2010KES/AUC]
C ₁₇ H ₁₁ N	[225-51-4]	benz[c]acridine				
	$\Delta_{\text{fus}}H$		20.3	381.4	DSC	[2010KES/AUC]
C ₁₇ H ₁₂	[238-84-6]	1,2-benzofluorene				
	$\Delta_{\text{fus}}H$		3.8	399.9		
	$\Delta_{\text{fus}}H$		18.4	462.8		[1996DOM/HEA]
	$\Delta_{\text{sub}}H$	(313–453)	105.4	383	GS	[1995NAS/LEN]
	$\Delta_{\text{v}}H$	(323–473)	83.7	398	GC	[2002LEI/CHA]
C ₁₇ H ₁₂	[243-17-4]	2,3-benzofluorene				
	$\Delta_{\text{fus}}H$		23.4	489.7		[1996DOM/HEA]
	$\Delta_{\text{sub}}H$	(344–398)	119.3 ± 1.3	371	ME	[1998OJA/SUU]
	$\Delta_{\text{sub}}H$	(313–453)	111.2	383	GS	[1995NAS/LEN]
	$\Delta_{\text{v}}H$		97.5 ± 3.9	298	CGC	[2008HAN/NUT]
	$\Delta_{\text{v}}H$	(323–473)	84.7	398	GC	[2002LEI/CHA]
C ₁₇ H ₁₂	[2381-21-7]	1-methylpyrene				
	$\Delta_{\text{v}}H$	(423–493)	92.3 ± 1.3	298	GC	[2006HAF/PAR]
C ₁₇ H ₁₂ Cl ₂ N ₄	[28911-01-5]	8-chloro-6-(2-chlorophenyl)-1-methyl-4 <i>H</i> -[1,2,4]triazolo[4,3- <i>a</i>]-[1,4]benzodiazepine (triazolam)				
	$\Delta_{\text{fus}}H$		41.0	514.5	DSC	[2008WAS/HOL]
C ₁₇ H ₁₂ N ₂ O ₂	[40448-93-9]	2-cyano-1(2 <i>H</i>)-quinolinecarboxylic acid, phenyl ester				
	$\Delta_{\text{fus}}H$		31.52	399	DSC	[2005LIZ/ZAB]
C ₁₇ H ₁₂ O	[152464-07-8]	4-ethynyl-1-[(4-ethynylphenyl)methoxy]benzene				
	$\Delta_{\text{fus}}H$		21.2	371.2	DSC	[1994MEL/LIT]
C ₁₇ H ₁₂ O ₂	[24776-44-1]	4-benzoyl-1-naphthol				
	$\Delta_{\text{fus}}H$		28.64	440.6	DSC	[1991ACR]
C ₁₇ H ₁₂ O ₂	[6333-07-9]	1-benzoyl-2-naphthol				
	$\Delta_{\text{fus}}H$		31.35	414.1	DSC	[1991ACR]
C ₁₇ H ₁₂ O ₂	[21009-99-4]	2-benzoyl-1-naphthol				
	$\Delta_{\text{fus}}H$		20.18	343.9	DSC	[1991ACR]
C ₁₇ H ₁₂ O ₂	[607-55-6]	1-naphthyl benzoate				
	$\Delta_{\text{fus}}H$		16.98	329.2		[1981BYS]
C ₁₇ H ₁₂ O ₂	[93-44-7]	2-naphthyl benzoate				

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	Enthalpy							
		$\Delta_{\text{fus}}H$			26.23	381.2		[1981BYS]
C ₁₇ H ₁₃ ClN ₄	[28981-97-7]		8-chloro-1-methyl-6-phenyl-4 <i>H</i> -[1,2,4]triazolo[4,3- <i>a</i>] [1,4]-benzodiazepine (alprazolam)		32.0	501.8	DSC	[2008WAS/HOL]
		$\Delta_{\text{fus}}H$						
C ₁₇ H ₁₃ F ₃ O	[172424-70-3]		4- <i>n</i> -propoxy-2',3',4'-trifluorodiphenylacetylene		26.1	327.3	DSC	[1995HSU/TSA]
		$\Delta_{\text{fus}}H$						
C ₁₇ H ₁₃ N	[6626-64-8]		5-methyl-5 <i>H</i> -indeno[2,1- <i>b</i>]quinoline		131.8 ± 1.3			[1966GEI/QUI, 1970COX/PIL]
		$\Delta_{\text{sub}}H$						
		Δ_vH	(375–388)	122.2	381		A	[1966GEI/QUI, 1970COX/PIL]
C ₁₇ H ₁₄ F ₂	[145698-43-7]		4- <i>n</i> -propyl-3',4'-difluorodiphenylacetylene		20.2	311	DSC	[1995HSU/TSA]
		$\Delta_{\text{fus}}H$						
C ₁₇ H ₁₄ F ₂ O	[172424-67-8]		4- <i>n</i> -propoxy-2',4'-difluorodiphenylacetylene		25.2	326.9	DSC	[1995HSU/TSA]
		$\Delta_{\text{fus}}H$						
C ₁₇ H ₁₄ F ₃ N ₃ O ₂ S	[169590-42-5]		4-[5-(3-methylphenyl)-3-(trifluoromethyl)-1 <i>H</i> -pyrazol-1-yl]-benzenesulphonamide (celecoxib)		34.35	436		[2003CHA/GUP]
		$\Delta_{\text{fus}}H$						
C ₁₇ H ₁₄ N ₂ O ₂	[122955-6]		1-[(2-methoxyphenyl)azo]-2-hydroxynaphthalene		142.4 ± 2.2	381		[1984KRI]
		$\Delta_{\text{sub}}H$	(374–388)					
C ₁₇ H ₁₄ N ₂ O ₂	[na]		2,2- <i>bis</i> -(4-cyanatophenyl)propane		26.69	355.8		[1996DOM/HEA]
		$\Delta_{\text{fus}}H$						
C ₁₇ H ₁₄ N ₄ O ₃	[243445-12-5]		2[4,5-dihydro-5-oxo-4-phenyl-3-(2-pyridinyl)-1,2,4-triazine-6(<i>H</i>)-ylidene]acetic acid, methyl ester		21.6	420.6		[2005SIK/MOD]
		$\Delta_{\text{fus}}H$						
C ₁₇ H ₁₄ O	[24330-03-8]		2:3,6:7-dibenzobicyclo[3.2.2]nona-2,6-dien-4-one		10.9	383.2	DSC	[2006PER/CON]
		$\Delta_{\text{fus}}H$						
		Δ_vH		94.5 ± 2.2	298		CGC	[2006PER/CON]
C ₁₇ H ₁₄ O ₄ S	[162011-90-7]		3-phenyl-4-[4-(methylsulfonyl)phenyl]-2(<i>5H</i>)-furanone (rofecoxib)		11.98	482.1	DSC	[2008TUN/TAB]
		$\Delta_{\text{fus}}H$						
C ₁₇ H ₁₄ O ₅	[117-52-2]		3-[1-(2-furanyl)-3-oxobutyl]-4-hydroxy-2 <i>H</i> -1-benzopyran-2-one		33.88	391.8	DSC	[1990DON/DRE]
		$\Delta_{\text{fus}}H$						
C ₁₇ H ₁₅ F	[145698-32-4]		4- <i>n</i> -propyl-4'-fluorodiphenylacetylene		24.1	324	DSC	[1995HSU/TSA]
		$\Delta_{\text{fus}}H$						
C ₁₇ H ₁₅ FO	[145532-20-3]		4- <i>n</i> -propoxy-4'-fluorodiphenylacetylene		27.1	356.8	DSC	[1995HSU/TSA]
		$\Delta_{\text{fus}}H$						
C ₁₇ H ₁₅ NO ₂	[154924-24-0]		1-[(4-nitrophenyl)ethynyl]-4-propylbenzene		23.26	351.3	DSC	[2002SPA/DZI]
		$\Delta_{\text{fus}}H$						
C ₁₇ H ₁₅ NO ₃	[483362-78-3]		1-[(4-nitrophenyl)ethynyl]-4-propoxybenzene		31.42	377	DSC	[2002SPA/DZI]
		$\Delta_{\text{fus}}H$						
C ₁₇ H ₁₅ NO ₃	[31842-01-0]		4-(1,3-dihydro-1-oxo-2 <i>H</i> -isoindol-2-yl)- α -methylbenzeneacetic acid (\pm) indoprofen		40.3	484.6	DSC	[2006WAS/HOL, 2008WAS/HOL]
		$\Delta_{\text{fus}}H$						
C ₁₇ H ₁₅ NO ₃ S	[313057-10-2]		4-(4-pentenyl)oxyphenyl 5-cyano-2-thiophene carboxylate		72.8	337.6	DSC	[2000WU/WAN]
		$\Delta_{\text{fus}}H$						
C ₁₇ H ₁₆ Br ₂ O ₃	[na]		isopropyl 4,4'-dibromobenzilate		24.55	348.1		[1996DOM/HEA]
		$\Delta_{\text{fus}}H$						

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
C ₁₇ H ₁₆ ClN ₅ O ₃	[na] $\Delta_{\text{fus}}H$	3-[[4-[(2-chloro-4-nitrophenyl)azo]phenyl](2-hydroxyethyl)amino]propanenitrile	26.29	428.2		[1991BAU/WEB]
C ₁₇ H ₁₆ F ₄ N ₄ O ₂	[91488-84-5] $\Delta_{\text{sub}}H$	N-ethyl-N-(2,2,3,3-tetrafluoropropyl)-4-[4-nitrophenyl]azobenzenamine	103		UV	[1984KAR/ROD]
C ₁₇ H ₁₆ F ₄ N ₄ O ₄	[1543-74-4] $\Delta_{\text{sub}}H$	2-[[4-(4-nitrophenyl)azo]phenyl](2,2,3,3-tetrafluoropropyl)amino]ethanol	103		UV	[1984KAR/ROD]
C ₁₇ H ₁₆ N ₂ O ₄	[129555-39-1] $\Delta_{\text{fus}}H$	5-phenoxyethyl-3-phenylcarbamoyl-2-oxazolidone	12.9	415.9		[1990SHI/HAY]
C ₁₇ H ₁₆ OS	[37014-01-0] $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$	tetrahydro-2,6-diphenyl-4 <i>H</i> -thiopyran-4-one	136 144 ± 3	375 298	ME ME	[1972GEI/SAW] [1972GEI/SAW, 1977PED/RYL]
C ₁₇ H ₁₆ O ₄	[54334-63-3] $\Delta_{\text{sub}}H$	diphenylmethylene diacetate (348–388)	122.1 ± 1.2	368	GS	[1996VER/PEN]
C ₁₇ H ₁₇ ClO ₆	[126-07-8] $\Delta_{\text{fus}}H$ $\Delta_{\text{fus}}H$	([1 <i>S</i>]- <i>trans</i> -7-chloro-,2',4,6-trimethoxy-6'-methylspiro[benzofuran-2(3 <i>H</i>), 1'-(2)cyclohexene]-3,4'-dione (griseofulvin)	44.7 39.39	491.2 495.2	DSC	[2006WAS/HOL]2008WAS/HOL] [1983GRA/ABO]
C ₁₇ H ₁₇ Cl ₂ N ₅ O ₄	[na] $\Delta_{\text{fus}}H$	N-[4-chloro-2-[(2-chloro-4-nitrophenyl)azo]-5-[(2-hydroxypropyl)amino]phenyl] acetamide	38.87	471.2		[1991BAU/WEB]
C ₁₇ H ₁₇ NO ₄	[483362-67-0] $\Delta_{\text{fus}}H$	2-(4-nitrophenyl)-1-(4-propoxyphenyl)ethanone	31.97	372.4	DSC	[2002SPA/DZI]
C ₁₇ H ₁₇ N ₅ O ₂	[na] $\Delta_{\text{sub}}H$	4-nitro-4'-[N-2-cyanoethyl-N-ethylamino]azobenzene	147.3			[1984KAR/KRU]
C ₁₇ H ₁₇ N ₅ O ₄	[231629-80-4] $\Delta_{\text{fus}}H$	6-(4-methoxyphenyl)-3,9-dihydro-3-[(2-hydroxyethoxy)methyl]-9-oxo-5 <i>H</i> -imidazol[1,2- <i>a</i>]pyrine	48.41	507.3	DSC	[1999ZIE/GOL]
C ₁₇ H ₁₈ ClNO ₂ S	[178870-32-1] $\Delta_{\text{fus}}H$	N-[4-chloro-3-[(3-methyl-2-butenyl)oxy]phenyl]-2-methyl-3-furancarbothiamide	36.94	400.8	DSC	[2001DAM/BLA]
C ₁₇ H ₁₈ FNO ₂	[164591-98-4] $\Delta_{\text{fus}}H$	4- <i>trans</i> -(3-fluoro-4-cyanophenyl)cyclohexyl (E)-but-2-enoate	21.1	393.2		[1995KEL/SCH]
C ₁₇ H ₁₈ FN ₃ O ₃	[85721-33-1] $\Delta_{\text{fus}}H$	1-cyclopropyl-6-fluoro-1,4-dihydro-4-oxo-7-(1-piperazinyl)-3-quinolinecarboxylic acid (ciprofloxacin)	64.48	541.5	DSC	[1994YU/ZIP]
C ₁₇ H ₁₈ N ₂ O ₃ S	[479578-81-9] $\Delta_{\text{fus}}H$	4-methoxy-N-[[[(3-methoxyphenyl)methyl]amino]thioxomethyl]benzamide	31.02	389.2	DSC	[2002ABB/WOH]
C ₁₇ H ₁₈ O ₃	[15131-43-8] $\Delta_{\text{v}}H$	2-hydroxy-4-butoxybenzophenone (393–443)	92.7	418	ME	[1984SUR]
C ₁₇ H ₁₈ O ₃	[87-18-3] $\Delta_{\text{sub}}H$ $\Delta_{\text{v}}H$	salicylic acid, 4-(<i>t</i> -butylphenyl) ester (293–336) (336–438)	137.4 90.4	308 351	A A, UV	[1987STE/MAL] [1987STE/MAL, 1960SCH/HIR]
C ₁₇ H ₁₈ O ₄	[101595-31-7] $\Delta_{\text{fus}}H$ $\Delta_{\text{sub}}H$	2-hydroxy-4,4'-diethoxybenzophenone	34.7 134.9	373.6 298	DSC B	[1999PRI/HAWN] [1999PRI/HAWN]
C ₁₇ H ₁₈ O ₄ S	[313057-14-6] $\Delta_{\text{fus}}H$	4-(4-pentenyl)oxyphenyl 5-methoxy-2-thiophene carboxylate	74.48	333.7	DSC	[2000WU/WAN]

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₇ H ₁₉ F ₃ O ₃	[164591-97-3]	4- <i>trans</i> -(trifluoromethoxyphenyl)cyclohexyl (E)-but-2-enoate				
	$\Delta_{\text{fus}}H$		21.6	340.2		[1995KEL/SCH]
C ₁₇ H ₁₉ NO ₃	[57-27-2]	7,8-didehydro-4,5-epoxy-17-methylmorphinan-3,6-diol (morphine)				
	$\Delta_{\text{fus}}H$		28.87	528.2	DTA	[1988ROY/FLY]
C ₁₇ H ₁₉ NO ₄	[72490-01-8]	N-[2-(4-phenoxyphenoxy)ethyl]carbamic acid, ethyl ester (fenoxycarb)				
	$\Delta_{\text{fus}}H$		26.98	326.3		[2005SUN/LIU4]
C ₁₇ H ₁₉ N ₃ O ₃	[850836-67-8]	6-(acetylamino)-2-cyano-1(2 <i>H</i>)-quinolinecarboxylic acid, 2-methylpropyl ester				
	$\Delta_{\text{fus}}H$		19.83	404.6	DSC	[2005LIZ/ZAB]
C ₁₇ H ₁₉ N ₃ O ₃	[1979-20-1]	6-(acetylamino)-2-cyano-1(2 <i>H</i>)-quinolinecarboxylic acid, butyl ester				
	$\Delta_{\text{fus}}H$		36.44	436.1	DSC	[2005LIZ/ZAB]
C ₁₇ H ₂₀ N ₂ O ₂ S	[373643-58-1]	N,N'- <i>bis</i> [(3-methoxyphenyl)methyl]thiourea				
	$\Delta_{\text{fus}}H$		33.22	354.7	DSC	[2002ABB/WOH]
C ₁₇ H ₂₀ O ₂	[6397-77-9]	diethoxydiphenylmethane				
	$\Delta_{\text{fus}}H$		19.9	323.2		[1998VER/PEN]
	$\Delta_{\text{sub}}H$		97.1 ± 1.1	298		[1998VER/PEN]
C ₁₇ H ₂₁ ClO ₄	[104225-37-8]	3-(3-chloro-4-methoxybenzoyl)-1,2,2-trimethylcyclopentanecarboxylic acid				
	$\Delta_{\text{fus}}H$		29.1	440.2	DSC	[1992TER/PAU]
C ₁₇ H ₂₁ F ₁₅	[139277-00-2]	1,1,1,2,3,3,4,4,5,5,6,6-dodecafluoro-2-(trifluoromethyl)hexadecane				
	$\Delta_{\text{trs}}H$		3.0	220		
	$\Delta_{\text{fus}}H$		18.0	261	DSC	[1992HOP/MOL]
C ₁₇ H ₂₁ NO ₂	[15299-99-7]	N,N-diethyl-2-(1-naphthyl)propionamide				
	$\Delta_{\text{fus}}H$		24.57	345.3	DSC	[1990DON/DRE]
C ₁₇ H ₂₁ NO ₃	[509-60-4]	4,5-epoxy-3-hydroxy-17-methylmorphinan-6-one (hydromorphone)				
	$\Delta_{\text{fus}}H$		35.61	539.2	DSC	[1988ROY/FLY]
C ₁₇ H ₂₁ NO ₄	[50-36-2]	cocaine				
	$\Delta_{\text{sub}}H$		127.2		GS	[1996ZIE/EIK]
	$\Delta_{\text{sub}}H$	(294–314)	112.3 ± 2.8	304	GS	[1984LWA/ELI]
C ₁₇ H ₂₁ NO ₆	[146607-85-4]	3-(3-nitro-4-methoxybenzoyl)-1,2,2-trimethylcyclopentanecarboxylic acid				
	$\Delta_{\text{fus}}H$		32.36	426.9	DSC	[1992TER/PAU]
C ₁₇ H ₂₁ N ₃ O ₂	[na]	2,2'-[[[3-methyl-4-(phenylazo)phenyl]imino]bis-ethanol				
	$\Delta_{\text{fus}}H$		31.9	384.2		[1988BAU/PER]
C ₁₇ H ₂₂ N ₂ O ₆	[76035-96-6]	<i>l</i>)-menthyl 3,5-dinitrobenzoate				
	$\Delta_{\text{fus}}H$		34.4	427.2	DTA	[1981CHI/GAR]
C ₁₇ H ₂₂ N ₂ O ₆	[80124-31-8]	<i>dl</i>)-menthyl 3,5-dinitrobenzoate				
	$\Delta_{\text{fus}}H$		30.6	401.2	DTA	[1981CHI/GAR]
C ₁₇ H ₂₂ O ₃	[115969-40-9]	3-(4-methylbenzoyl)-1,2,2-trimethylcyclopentanecarboxylic acid				
	$\Delta_{\text{fus}}H$		30.07	468.2	DSC	[1993TER/BOU]
C ₁₇ H ₂₂ O ₃ S	[145918-70-3]	<i>p</i> -(1 <i>R</i> ,3 <i>S</i>)-3-thianisoyl-1,2,2-trimethylcyclopentanecarboxylic acid				
	$\Delta_{\text{fus}}H$		23.56	393.7	DSC	[1994TER/CAS, 1993RAM/BOU]
C ₁₇ H ₂₃ NO ₃	[172589-28-5]	3-[(hydroxyimino)phenylmethyl]-1,2,2-trimethylcyclopentanecarboxylic acid methyl ester				
	$\Delta_{\text{fus}}H$		33.8	422	DSC	[1995NUR/LEL]
C ₁₇ H ₂₃ NO ₃	[51-55-8]	α -(hydroxymethyl)-benzene acetic acid, (3- <i>endo</i>)-8-methyl-8-azabicyclo[3.2.1]oct-3-yl ester (atropine)				
	$\Delta_{\text{fus}}H$		35.5	388.5	DSC	[2009DOM/POB]

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
C ₁₇ H ₂₃ NO ₄	[146607-86-5]	3-(4-methoxy-3-aminobenzoyl)-1,2,2-trimethylcyclopentanecarboxylic acid	41.32	498.6	DSC	[1992TER/PAU]
C ₁₇ H ₂₄ O ₂	[6284-35-1]	menthyl benzoate	69.9	411	A	[1987STE/MAL, 1947STU]
C ₁₇ H ₂₆ N ₂ O	[84057-95-4]	N-(2,6-dimethylphenyl)-1-propyl-2-piperidinecarboxamide	44.5	414.2	DSC	[1997NEM/ACS]
C ₁₇ H ₂₆ O ₃	[na]	3-decyloxybenzoic acid	33.88	345.1		[2001LAI/LEE]
C ₁₇ H ₂₇ NO ₂	[93413-69-5]	1-[2-(dimethylamino)-1-(4-methoxyphenyl)ethyl]cyclohexanol (venlafaxine)	27.2	348.1	DSC	
			26.4	349.7	DSC	
			24.4	351.3	DSC	[2009VAN/WES]
C ₁₇ H ₂₇ N ₃ O ₃	[83963-52-4]	1-decyl-3-(4-nitrophenyl) urea	37.92	390.6	DSC	[1993TIE/FRA]
C ₁₇ H ₂₈	[6742-54-7]	undecylbenzene	82.4 ± 0.4	298	GS	[2006VER]
			66.7	465		[1999DYK/SVO]
			84.7	298		[1971WIL/ZWO]
C ₁₇ H ₂₈ N ₂ OS	[373642-33-2]	N-[(3-methoxyphenyl)methyl]-N'-octylthiourea	37.94	350.7	DSC	[2002ABB/WOH]
C ₁₇ H ₂₈ O	[56103-67-4]	4-methyl-2,6-di- <i>tert</i> -pentylphenol	65.9	453	A	[1987STE/MAL]
C ₁₇ H ₂₈ O ₂	[55095-35-7]	1,3-dimethoxy-2-nonylbenzene	79.2	458	A, GC	[1987STE/MAL, 1975KUN/LIL]
C ₁₇ H ₂₈ O ₄	[3220-58-4]	dicyclohexyl glutarate	101.1 ± 0.8	298	GS	[2008LIP/KRA]
C ₁₇ H ₃₂	[26186-00-5]	1-heptadecyne	62.7	453		[1999DYK/SVO]
C ₁₇ H ₃₂	[61847-96-9]	2-heptadecyne	63.7	461		[1999DYK/SVO]
C ₁₇ H ₃₂	[61886-63-3]	3-heptadecyne	62.5	453		[1999DYK/SVO]
C ₁₇ H ₃₂ Cl ₄	[93479-16-4]	1,1,1,1,17-tetrachloroheptadecane	108	366	A	[1987STE/MAL]
C ₁₇ H ₃₂ O	[3661-77-6]	cycloheptadecanone	75.7			[1938WOL/WEG, 1960JON, 1970COX/PIL]
C ₁₇ H ₃₂ O ₂	[5637-97-8]	oxa-2-cyclotetradecanone	73.5	418	A	[1987STE/MAL]
C ₁₇ H ₃₂ O ₂	[21643-42-5]	tetradecyl acrylate	69.4	473	A	[1987STE/MAL]
C ₁₇ H ₃₂ O ₂	[35835-77-9]	(Z) 9-pentadecenyl acetate	93.6	298	GC	[1997KOU/HOS, 2000OVA/KOU]
C ₁₇ H ₃₂ O ₂	[64437-41-8]	(E) 9-pentadecenyl acetate				

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		T _m (K)	Method	Reference	
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)				
		$\Delta_v H$	(363–408)	94.3	298	GC	[1997KOU/HOS, 2000OVA/KOU]
C ₁₇ H ₃₂ O ₂	[64437-43-0]		(Z) 10-pentadecenyl acetate				
		$\Delta_v H$	(363–408)	94.1	298	GC	[1997KOU/HOS, 2000OVA/KOU]
C ₁₇ H ₃₂ O ₂	[64437-45-2]		(E) 10-pentadecenyl acetate				
		$\Delta_v H$	(363–408)	94.6	298	GC	[1997KOU/HOS, 2000OVA/KOU]
C ₁₇ H ₃₂ O ₂	[35153-25-4]		(Z) 11-pentadecenyl acetate				
		$\Delta_v H$	(363–408)	94.6	298	GC	[1997KOU/HOS, 2000OVA/KOU]
C ₁₇ H ₃₂ O ₂	[40535-40-8]		(E) 11-pentadecenyl acetate				
		$\Delta_v H$	(363–408)	94.9	298	GC	[1997KOU/HOS, 2000OVA/KOU]
C ₁₇ H ₃₂ O ₂	[70711-45-4]		(Z) 12-pentadecenyl acetate				
		$\Delta_v H$	(363–408)	95.1	298	GC	[1997KOU/HOS, 2000OVA/KOU]
C ₁₇ H ₃₂ O ₂	[73304-17-3]		(E) 12-pentadecenyl acetate				
		$\Delta_v H$	(363–408)	94.5	298	GC	[1997KOU/HOS, 2000OVA/KOU]
C ₁₇ H ₃₂ O ₂	[70711-46-5]		(Z) 13-pentadecenyl acetate				
		$\Delta_v H$	(363–408)	95.9	298	GC	[1997KOU/HOS, 2000OVA/KOU]
C ₁₇ H ₃₂ O ₂	[na]		(E) 13-pentadecenyl acetate				
		$\Delta_v H$	(363–408)	95.9	298	GC	[1997KOU/HOS, 2000OVA/KOU]
C ₁₇ H ₃₂ O ₂	[1120-25-8]		methyl Z 9-hexadecenoate				
		$\Delta_v H$		96.4 ± 0.7	298	CGC	[2007LIP/KAP]
C ₁₇ H ₃₂ O ₂	[na]		methyl Z 10-heptadecenoate				
		$\Delta_v H$		100.8	298	CGC	[2007LIP/KAP]
C ₁₇ H ₃₂ O ₃	[1725-00-4]		1,8-dioxo-9-cyclononadecanone				
		$\Delta_v H$	(403–463)	77.0	418	A	[1987STE/MAL]
C ₁₇ H ₃₂ O ₄	[2917-73-9]		dibutyl nonadioate				
		$\Delta_v H$	(313–450)	88.4	328	A	[1987STE/MAL]
C ₁₇ H ₃₂ O ₅	[na]		nonyl[1-(butoxycarbonyl)ethyl]carbonate				
		$\Delta_v H$	(420–534)	73.8	435	A	[1987STE/MAL]
C ₁₇ H ₃₃ N	[5399-02-0]		heptadecanonitrile				
		$\Delta_v H$	(348–385)	98.9 ± 0.4	298	GS	[2005EME/VER]
		$\Delta_v H$	(425–620)	81.2	440	A	[1987STE/MAL]
C ₁₇ H ₃₃ NO ₃	[71448-29-8]		N-tetradecanyl-(<i>l</i>)-alanine				
		$\Delta_{\text{fus}} H$		52.3	367.1	DSC	[1986MIY/MAT]
C ₁₇ H ₃₃ NO ₃	[na]		N-dodecanoyl-(<i>l</i>)-valine				
		$\Delta_{\text{fus}} H$		33.1	380.1	DSC	[1986MIY/MAT]
C ₁₇ H ₃₃ NO ₃	[na]		N-dodecyl-(<i>dl</i>)-valine				
		$\Delta_{\text{fus}} H$		64.4	364.6	DSC	[1986MIY/MAT]
C ₁₇ H ₃₄	[5634-30-0]		dodecylcyclopentane				
		$\Delta_v H$	(450–619)	68.0	465		[1999DYK/SVO]
		$\Delta_v H$		85.5	298		[1971WIL/ZWO]
C ₁₇ H ₃₄	[54105-66-7]		undecylcyclohexane				
		$\Delta_v H$	(450–622)	67.0	465		[1999DYK/SVO]
		$\Delta_v H$		84.6	298		[1971WIL/ZWO]
C ₁₇ H ₃₄	[6765-39-5]		1-heptadecene				

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		T _m (K)	Method	Reference
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)			
		$\Delta_{\text{v}}H$	(598–746)	55.5	613	[1999DYK/SVO]
		$\Delta_{\text{v}}H$	(376–432)	72.3	391	A [1987STE/MAL]
		$\Delta_{\text{v}}H$		84.9	298	[1971WIL/ZWO]
C ₁₇ H ₃₄	[295-97-6]	cycloheptadecane				
		$\Delta_{\text{sub}}H$		66.1 ± 0.6		[1957VAN, 1970COX/PIL]
C ₁₇ H ₃₄ O	[2922-51-2]	2-heptadecanone				
		$\Delta_{\text{v}}H$	(402–593)	77.0	417	A [1987STE/MAL, 1947STU]
C ₁₇ H ₃₄ O	[6064-42-2]	7-heptadecanone				
		$\Delta_{\text{v}}H$		94.5 ± 1.8	298	CGC [2006PER/CON]
C ₁₇ H ₃₄ O	[540-08-9]	9-heptadecanone				
		$\Delta_{\text{fus}}H$		66.68	323.9	[1993VIL/HAM]
		$\Delta_{\text{v}}H$	(439–482)	78.3	454	A, ME [1987STE/MAL, 1938UBB]
C ₁₇ H ₃₄ O ₂	[112-39-0]	methyl hexadecanoate (methyl palmitate)				
		$\Delta_{\text{fus}}H$		58.1	302.2	
		$\Delta_{\text{fus}}H$		56.0	305.2	[2003NIK/MAR, 2004CHI/ZHA]
		$\Delta_{\text{fus}}H$		68.16	307.2	[1993ACR]
		$\Delta_{\text{sub}}H$	(291–301)	152.3 ± 2	296	ME [1965DAV/KYB, 1987STE/MAL]
		$\Delta_{\text{v}}H$		93.4	350	[2002VAN/VAN]
		$\Delta_{\text{v}}H$		83.3 ± 0.4	397	[2002VAN/VAN]
		$\Delta_{\text{v}}H$		96.8 ± 0.6	298	[2002VAN/VAN]
		$\Delta_{\text{v}}H$	(463–523)	96.4	298	GC [1997KRO/VEL]
		$\Delta_{\text{v}}H$	(433–473)	93.2	298	CGC [1995CHI/HOS]
		$\Delta_{\text{v}}H$	(453–543)	78.2	498	GC [1993HUS/SAR]
		$\Delta_{\text{v}}H$	(287–322)	U 69.6	302	A [1987STE/MAL]
		$\Delta_{\text{v}}H$	(411–543)	82.4	426	A [1987STE/MAL, 1963ROS/SCH]
		$\Delta_{\text{v}}H$	(378–445)	82.6	393	MG,OM [1952SCO/MAC]
		$\Delta_{\text{v}}H$	(422–475)	71.4	437	[1948BON/ATH]
C ₁₇ H ₃₄ O ₂	[110-27-0]	isopropyl tetradecanoate				
		$\Delta_{\text{v}}H$	(413–466)	70.2	428	A [1987STE/MAL, 1948BON/ATH, 1984BOU/FRI]
C ₁₇ H ₃₄ O ₂	[14303-70-9]	propyl tetradecanoate				
		$\Delta_{\text{v}}H$	(420–474)	71.3	435	A [1987STE/MAL, 1948BON/ATH, 1984BOU/FRI]
C ₁₇ H ₃₄ O ₂	[506-12-7]	heptadecanoic acid				
		$\Delta_{\text{trs}}H$		7.5	331.2	
		$\Delta_{\text{fus}}H$		46.5	333.5	DSC [2007GBA/NEG, 2008GBA/NEG]
		$\Delta_{\text{trs}}H$		7.44	329.2	
		$\Delta_{\text{fus}}H$		51.33	334.3	[1996DOM/HEA]
		$\Delta_{\text{fus}}H$		51.9	333	[1976BER/BER]
		$\Delta_{\text{sub}}H$	(280–302)	151		TPTD [2005CHA/ZIE]
		$\Delta_{\text{sub}}H$	(291–316)	168		TPTD [2001CHA/TOB]
Note: Experimental values based on the TPTD method are often inconsistent with values determined using other experimental methods						
		$\Delta_{\text{v}}H$	(449–637)	100.7	464	A [1987STE/MAL]
		$\Delta_{\text{v}}H$	(357–382)	112.7 ± 2.0	372	ME, TE [1982DEK/SCH]
C ₁₇ H ₃₄ O ₃	[1323-03-1]	tetradecyl lactate				
		$\Delta_{\text{v}}H$	(388–608)	86.4	403	A [1987STE/MAL]
C ₁₇ H ₃₅ Br	[3508-00-7]	1-bromoheptadecane				

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		T _m (K)	Method	Reference
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)			
C ₁₇ H ₃₅ Cl		$\Delta_{\text{v}}H$	(472–673)	71.6	487	A, E [1987STE/MAL, 1961LI/ROS, 1970DYK/VAN]
	[62016-75-5]		1-chloroheptadecane			
		$\Delta_{\text{v}}H$	(450–673)	103.6	298	[2006BOL/NER2]
C ₁₇ H ₃₅ F		$\Delta_{\text{v}}H$	(437–623)	73.2	465	A, E [1987STE/MAL, 1970DYK/VAN, 1961LI/ROS]
	[1545-17-1]		1-fluoroheptadecane			
C ₁₇ H ₃₅ I		$\Delta_{\text{v}}H$	(517–673)	68.4	452	A, E [1987STE/MAL, 1961LI/ROS, 1970DYK/VAN]
	[26825-83-2]		1-iodoheptadecane			
C ₁₇ H ₃₅ NO		$\Delta_{\text{v}}H$	(517–673)	104.7	298	A,E [1987STE/MAL, 1961LI/ROS, 1970DYK/VAN, 2006BOL/NER]
		$\Delta_{\text{v}}H$	(517–673)	73.0	532	A, E [1987STE/MAL, 1961LI/ROS, 1970DYK/VAN]
C ₁₇ H ₃₅ NO ₂		$\Delta_{\text{sub}}H$	(345–355)	144.5 ± 0.8	350	GS [1959DAV/JON, 1987STE/MAL]
	[7388-58-1]		N-methyl hexadecanamide			
C ₁₇ H ₃₅ NO ₂		$\Delta_{\text{v}}H$	(413–491)	107.5	428	A [1987STE/MAL]
	[96945-44-7]		N-tetradecyllactamide			
C ₁₇ H ₃₆		$\Delta_{\text{trs}}H$		10.8	284.2	
		$\Delta_{\text{fus}}H$		39.4	294.7	DSC [2004MON/RAJ]
		$\Delta_{\text{trs}}H$		10.96	284.3	
		$\Delta_{\text{fus}}H$		40.17	295.1	[1996DOM/HEA]
		$\Delta_{\text{sub}}H$		125.1	298	[1972MOR3]
		$\Delta_{\text{sub}}H$	(288–293)	131.3 ± 13	290	ME [1949BRA/SHE, 1960JON]
		$\Delta_{\text{v}}H$		86.5	298	[1994RUZ/MAJ]
		$\Delta_{\text{v}}H$	(289–320)	91.1	304	A [1987STE/MAL]
		$\Delta_{\text{v}}H$	(488–577)	62.9	503	A [1987STE/MAL]
		$\Delta_{\text{v}}H$		86.0 ± 0.8	298	C [1972MOR]
		$\Delta_{\text{v}}H$		86.2	298	[1971WIL/ZWO]
		$\Delta_{\text{v}}H$	(445–470)	71.6	457	ME [1938UBB]
	C ₁₇ H ₃₆		$\Delta_{\text{v}}H$	(428–569)	63.5	443
[1560-92-5]			2-methylhexadecane			
C ₁₇ H ₃₆		$\Delta_{\text{v}}H$	(428–567)	63.4	443	A [1987STE/MAL, 1959TER/BRI]
	[6418-43-5]		3-methylhexadecane			
C ₁₇ H ₃₆		$\Delta_{\text{v}}H$	(420–567)	58.7	435	A [1987STE/MAL, 1959TER/BRI]
	[25117-26-4]		4-methylhexadecane			
C ₁₇ H ₃₆		$\Delta_{\text{v}}H$	(422–566)	59.8	437	A [1987STE/MAL, 1959TER/BRI]
	[25117-34-4]		5-methylhexadecane			
C ₁₇ H ₃₆		$\Delta_{\text{v}}H$	(424–569)	60.6	439	A [1987STE/MAL, 1959TER/BRI]
	[2882-97-5]		2,3-dimethylpentadecane			
C ₁₇ H ₃₆		$\Delta_{\text{v}}H$	(419–546)	65.2	434	A [1987STE/MAL, 1959TER/BRI]
	[61868-07-3]		2,4-dimethylpentadecane			
C ₁₇ H ₃₆		$\Delta_{\text{v}}H$	(411–534)	64.5	426	A [1987STE/MAL]
	[101791-53-1]		2,4,6-trimethyltetradecane			
C ₁₇ H ₃₆		$\Delta_{\text{v}}H$		78.0 ± 1.8	298	CGC [1995CHI/HES]
	[93816-24-1]		4,4-dipropylundecane			
C ₁₇ H ₃₆ O		$\Delta_{\text{fus}}H$		63.06	327.3	[2003VAN/VAN]
	[1454-85-9]		1-heptadecanol			

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
	Enthalpy							
			Note: The enthalpy of fusion value includes the enthalpy of solid-to-solid transition that occurs at 323.2 K					
		$\Delta_{\text{fus}}H$			37.0	326.6	DSC	[2004VEN/CAL]
		$\Delta_{\text{trs}}H$			25.2	323.6		
		$\Delta_{\text{fus}}H$			37.0	326.6		[2002VEN/RAM]
		$\Delta_{\text{sub}}H$			169.5 ± 2.2			[1965DAV/KYB, 1970COX/PIL]
		$\Delta_{\text{v}}H$			112.5 ± 0.5	298	CGC	[2006NIC/KWE]
		$\Delta_{\text{v}}H$		(460–620)	78.3	475	A	[1987STE/MAL]
		$\Delta_{\text{v}}H$		(473–623)	75.9	488	A	[1987STE/MAL]
C ₁₇ H ₃₆ O	[103385-34-8]		4-heptadecanol					
		$\Delta_{\text{fus}}H + \Delta_{\text{trs}}H$			35.7	311.5	DSC	[2006NIC/KWE]
C ₁₇ H ₃₆ O	[112283-13-3]		6-heptadecanol					
		$\Delta_{\text{fus}}H + \Delta_{\text{trs}}H$			49.0	315.8	DSC	[2006NIC/KWE]
		$\Delta_{\text{v}}H$			108.6 ± 1.0	298	CGC	[2006NIC/KWE]
C ₁₇ H ₃₆ O	[93658-33-4]		7-heptadecanol					
		$\Delta_{\text{fus}}H + \Delta_{\text{trs}}H$			28.8	314.4	DSC	[2006NIC/KWE]
		$\Delta_{\text{v}}H$			108.2 ± 0.8	298	CGC	[2006NIC/KWE]
C ₁₇ H ₃₆ O	[624-08-8]		9-heptadecanol					
		$\Delta_{\text{fus}}H + \Delta_{\text{trs}}H$			43.2	330.2	DSC	[2006NIC/KWE]
		$\Delta_{\text{v}}H$			108.5 ± 0.4	298	CGC	[2006NIC/KWE]
C ₁₇ H ₃₆ O ₂	[66577-59-1]		1,17-heptadecanediol					
		$\Delta_{\text{trs}}H$			34.9	98.4		
		$\Delta_{\text{fus}}H$			30.8	367.3	DSC	[1999OGA/NAK]
C ₁₇ H ₃₆ O ₂ S	[79768-75-5]		3-(tetradecylthio)-1,2-propanediol					
		$\Delta_{\text{trs}}H$			16.3	302.5		
		$\Delta_{\text{fus}}H$			26.8	336.4	DSC	[1993ACR]
C ₁₇ H ₃₆ O ₃	[1561-06-4]		3-(tetradecyloxy)-1,2-propanediol					
		$\Delta_{\text{fus}}H$			62.1	331.3	DSC	[1993ACR]
C ₁₇ H ₃₆ S	[53193-22-9]		1-heptadecanethiol					
		$\Delta_{\text{v}}H$	(481–657)		74.6	496		[1999DYK/SVO]
C ₁₇ H ₃₇ N	[4200-95-7]		heptadecylamine					
		$\Delta_{\text{v}}H$	(522–636)		68.2	537	A	[1987STE/MAL, 1956MAN2]
C ₁₇ H ₃₇ NO ₂	[111953-19-6]		3-(tetradecylamino)-1,2-propanediol					
		$\Delta_{\text{fus}}H$			64.9	356.2	DSC	[1993ACR]
C ₁₈ H ₁₀	[65513-20-4]		benzo[3,4]cyclobuta[1,2-a]biphenylene ([3]phenylene)					
		$\Delta_{\text{sub}}H$			115.1 ± 0.8			[2000BEC/FAU]
C ₁₈ H ₁₀	[203-12-3]		benzo[ghi]fluoranthene					
		$\Delta_{\text{trs}}H$			5.35	402.8		
		$\Delta_{\text{trs}}H$			0.88	402.1		
		$\Delta_{\text{trs}}H$			0.44	352.7		
		$\Delta_{\text{fus}}H$			11.8	424		[1980SMI]
C ₁₈ H ₁₀ BrNO ₃	[10319-14-9]		2(4-bromo-3-hydroxy-2-quinolinyl)-1 <i>H</i> -indene-1,3(2 <i>H</i>)-dione (C. I. disperse yellow 64)					
		$\Delta_{\text{sub}}H$	(483–523)		130.6	498		[1987STE/MAL]

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₈ H ₁₀ Cl ₂ O ₂ S ₂	[2379-74-0]	6-chloro-2-(6-chloro-4-methyl-3-oxobenzo[b]thien-2(3H)-ylidene)-4-methyl-benzo[b]thiophen-3(2H)-one (C.I. Vat Red 1)				
	$\Delta_{\text{sub}}H$	(519–634)	148	577	GS	[1986NIS/AND]
C ₁₈ H ₁₀ Cl ₂ O ₂ S ₂	[5462-29-3]	5-chloro-2-(5-chloro-7-methyl-3-oxobenzo[b]thien-2(3H)-ylidene)-7-methyl-benzo[b]thiophen-3(2H)-one (C.I. Vat Violet 2)				
	$\Delta_{\text{sub}}H$	(519–634)	93.0	577	GS	[1986NIS/AND]
C ₁₈ H ₁₀ O ₂	[2498-66-0]	1,2-benzanthra-9,10-quinone				
	$\Delta_{\text{sub}}H$		82.8 ± 4.0			[1956MAG, 1970COX/PIL]
C ₁₈ H ₁₀ O ₂	[1090-13-7]	5,12-tetracenequinone				
	$\Delta_{\text{sub}}H$		108.8 ± 5.0			[1956MAG, 1970COX/PIL]
C ₁₈ H ₁₀ O ₄	[1785-52-0]	6,11-dihydroxy-5,12-naphthacenedione				
	$\Delta_{\text{sub}}H$	(426–446)	144.2 ± 1.4	436	ME	[1998OJA/SUU]
C ₁₈ H ₁₁ NO ₂	[7496-02-8]	6-nitrochrysene				
	$\Delta_{\text{fus}}H$		28.4	486.6	DSC	[2010KES/AUC]
C ₁₈ H ₁₁ NO ₃	[7576-65-0]	2-(3-hydroxy-2-quinolinylidene)-indeno-1,3-dione (Disperse yellow 54)				
	$\Delta_{\text{fus}}H$		30.89	539.2		[1991BAU/WEB]
	$\Delta_{\text{sub}}H$		125.2 ± 0.4		LE	[1998PRI/HAW]
	$\Delta_{\text{sub}}H$	(483–513)	139	498		[1973MCD]
C ₁₈ H ₁₂	[92-24-0]	naphthacene (tetracene)				
	$\Delta_{\text{sub}}H$	(399–430)	124.8 ± 2.6		ME	[2009OJA/CHE]
	$\Delta_{\text{sub}}H$	(386–472)	126.1 ± 9.0	429	ME	[1998OJA/SUU]
	$\Delta_{\text{sub}}H$	(313–453)	126.5	383	GS	[1995NAS/LEN]
	$\Delta_{\text{sub}}H$	(419–446)	143.7 ± 0.5	298	TE,M	[1980DEK]
	$\Delta_{\text{sub}}H$		124.7 ± 4	422	ME	[1967WAK/INO, 1977PED/RYL]
	$\Delta_{\text{sub}}H$					[1970COX/PIL]
	$\Delta_{\text{sub}}H$	(433–493)	128.8	473	HSA	[1965MOR]
	$\Delta_{\text{sub}}H$	(433–483)	132.6	468	HSA	[1964FIE/MAC]
	$\Delta_{\text{sub}}H$		117.2	459	ME	[1952INO/SHI, 1960JON]
	$\Delta_{\text{sub}}H$		U 92.0	384	ME	[1951INO]
	$\Delta_{\text{sub}}H$		124.3			[1951MAG/HAR, 1960JON]
	Δ_vH		106.2 ± 3.7	298	CGC	[2008HAN/NUT]
C ₁₈ H ₁₂	[56-55-3]	benz[a]anthracene				
	$\Delta_{\text{fus}}H$		20.1	433.5	DSC	[2010KES/AUC]
	$\Delta_{\text{fus}}H$		21.38	434.3		[1991ACR]
	$\Delta_{\text{sub}}H$	(313–453)	115.5	383	GS	[1995NAS/LEN]
	$\Delta_{\text{sub}}H$	(330–390)	113.4	345	ME	[1987STE/MAL, 1974MUR/POL]
	$\Delta_{\text{sub}}H$		104 ± 2	351	TE	[1983FER/IMP]
	$\Delta_{\text{sub}}H$	(283–323)	U 81.3 ± 2.5	303	GS	[1983SON/ZOL]
	$\Delta_{\text{sub}}H$	(373–396)	123.3 ± 3	298		[1980DEK]
	$\Delta_{\text{sub}}H$	(357–454)	120.5	405	ME	[1967WAK/INO]
	$\Delta_{\text{sub}}H$	(377–403)	104.6 ± 4.2	390	ME	[1964KEL/RIC, 1987STE/MAL]
	$\Delta_{\text{sub}}H$	(333–393)	119.7	363		[1958HOY/PEP]
	$\Delta_{\text{sub}}H$		U 109.2			[1951MAG/HAR, 1960JON]
	Δ_vH		105.8 ± 1.9	298	CGC	[2008HAN/NUT]
	Δ_vH	(463–525)	96.6 ± 1.4	298	GC	[2006HAF/PAR]
	Δ_vH	(343–453)	91.0	398	GC	[1990HIN/BID2]
C ₁₈ H ₁₂	[217-59-4]	triphenylene				

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	Enthalpy							
		$\Delta_{\text{fus}}H$			23.0	471.2	DSC	[2010KES/AUC]
		$\Delta_{\text{fus}}H$			24.74	471		[1996DOM/HEA]
		$\Delta_{\text{sub}}H$		(313–453)	114.5	383	GS	[1995NAS/LEN]
		$\Delta_{\text{sub}}H$		(381–406)	126.5 ± 4	298	TE,ME	[1980DEK]
		$\Delta_{\text{sub}}H$		(363–468)	107.6	378		[1987STE/MAL]
		$\Delta_{\text{sub}}H$		(338–398)	118 ± 4	368		[1958HOY/PEP, 1970COX/PIL]
		$\Delta_{\text{sub}}H$			107.1	425	ME	[1967WAK/INO]
		Δ_vH			106.1 ± 3.9	298	CGC	[2008HAN/NUT]
		Δ_vH		(323–473)	88.5	398	GC	[2002LEI/CHA]
		Δ_vH		(535–768)	67.7	550		[1999DYK/SVO]
C₁₈H₁₂	[218-01-9]		chrysene					
		$\Delta_{\text{fus}}H$			23.6	527	DSC	[2010KES/AUC]
		$\Delta_{\text{trs}}H$			3.22	512.2		
		$\Delta_{\text{fus}}H$			26.15	531.4	DSC	[1973CAS/VEC]
		$\Delta_{\text{sub}}H$		(313–453)	118.8	383	GS	[1995NAS/LEN]
		$\Delta_{\text{sub}}H$			131 ± 4	298	TE,ME	[1980DEK]
		$\Delta_{\text{sub}}H$			117.6 ± 4	400	ME	[1967WAK/INO, 1970COX/PIL]
		$\Delta_{\text{sub}}H$		(353–418)	121.4	385		[1958HOY/PEP]
		$\Delta_{\text{sub}}H$			117.6			[1951MAG/HAR, 1960JON]
		Δ_vH			106.2	298	CGC	[2008ZHA/UNH]
		Δ_vH		(463–513)	97.0 ± 1.4	298	GC	[2006HAF/PAR]
		Δ_vH		(323–473)	89.6	398	GC	[2002LEI/CHA]
C₁₈D₁₂	[1719-03-5]		chrysene - d ₁₂					
		Δ_vH			106	298	CGC	[2008ZHA/UNH]
C₁₈H₁₂	[195-19-7]		benzo[c]phenanthrene (3,4-benzophenanthrene)					
		$\Delta_{\text{fus}}H$			15.5	339.2	DSC	[2010KES/AUC]
		$\Delta_{\text{fus}}H$			16.32	334.7		[1991ACR]
		$\Delta_{\text{sub}}H$			106.3 ± 4.2			[1951MAG/HAR, 1970COX/PIL, 1967WAK/INO]
C₁₈H₁₂N₂	[119-91-5]		2,2'-biquinoline					
		$\Delta_{\text{sub}}H$		(393–411)	129.5 ± 0.8	402	ME	[1997RIB/MAT3]
		$\Delta_{\text{sub}}H$			134.7 ± 1.3	298		[1997RIB/MAT3]
		$\Delta_{\text{sub}}H$			96.6 ± 0.9			[1985SKI/PIL]
C₁₈H₁₂O	[10435-67-3]		2-phenylindeno[2,1-b]pyran					
		$\Delta_{\text{sub}}H$		(394–424)	132.8	409		[1987STE/MAL, 1966GEI/QUI]
C₁₈H₁₂O₃	[568-73-0]		1,6-dimethylphenanthro[1,2-b]furan-10,11-dione					
		$\Delta_{\text{fus}}H$			22.09	495.4	DSC	[1988HUA/TAN]
C₁₈H₁₃ClN₂O₃	[1979-26-7]		2-cyano-6-methoxy-1(2 <i>H</i>)-quinolinecarboxylic acid, 4-chlorophenyl ester					
		$\Delta_{\text{fus}}H$			22.35	374.2	DSC	[2005LIZ/ZAB]
C₁₈H₁₃FO	[145532-14-5]		4-ethoxy-4'-fluorodiphenylodiacetylene					
		$\Delta_{\text{fus}}H$			33.9	400.2	DSC	[1993JUA/CHE]
C₁₈H₁₃F₂₅	[89109-70-6]		1,1,1,2,2,3,3,4,4,5,5,6,6,7,7,8,8,9,9,10,10,11,11,12,12-pentacosofluorooctadecane					
		$\Delta_{\text{trs}}H$			3.3	317.2		
		$\Delta_{\text{fus}}H$			21.8	352.2	DSC	[1986RUS/RAB]
C₁₈H₁₃O₄P	[99208-50-1]		2-(6-oxido-6 <i>H</i> -dibenz[<i>c,e</i>][1,2]oxaphosphorin-6-yl)-1,4-dihydroxyphenylene					
		$\Delta_{\text{fus}}H$			41.7	524.1	DSC	[2008FAN/WAN]

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	Enthalpy	Temp (K) Range					
C ₁₈ H ₁₄	[84-15-1]	<i>o</i> -terphenyl					
	$\Delta_{\text{fus}}H$			16.9	327.8	DSC	[1997VER2]
	$\Delta_{\text{fus}}H$			17.2	328.4		[1995MUR/PAI]
	$\Delta_{\text{fus}}H$			17.2	329.4		[1972CHA/BES]
	$\Delta_{\text{sub}}H$	(312–328)		103.0 ± 0.4	298	ME	[2008RIB/SAN6]
	$\Delta_{\text{sub}}H$			97 ± 1	298	B	[1979KIM/TAK]
	Δ_vH	(335–368)		81.0 ± 0.4	352	GS	[1997VER2]
	Δ_vH	(335–368)		84.2 ± 0.4	298	GS	[1997VER2]
	Δ_vH	(576–786)		60.5	591	DSC	[1996BAC/GRZ]
	Δ_vH	(343–462)		77.6	403		[1989SAS/NGU]
Δ_vH	(462–650)		68.5	477	A	[1987STE/MAL]	
C ₁₈ H ₁₄	[92-06-8]	<i>m</i> -terphenyl					
	$\Delta_{\text{fus}}H$			31.0	361.2	DSC	[1997VER2]
	$\Delta_{\text{fus}}H$			22.59	360		[1971KAM/MIT]
	$\Delta_{\text{sub}}H$	(337–359)		118.6 ± 0.7	298	ME	[2008RIB/SAN6]
	$\Delta_{\text{sub}}H$	(329–353)		115.5 ± 1.6	341	T	[1997VER2]
	$\Delta_{\text{sub}}H$			118.1 ± 1.6	298		[1997VER2]
	$\Delta_{\text{sub}}H$			120 ± 1	298		[1979KIM/TAK]
	$\Delta_{\text{sub}}H$	(313–363)		119	338		[1958HOY/PEP]
	Δ_vH			97.2 ± 0.3	298	CGC	[2001PUR/CHI]
	Δ_vH	(462–691)		76.1	477	A	[1987STE/MAL]
C ₁₈ H ₁₄	[92-94-4]	<i>p</i> -terphenyl					
	$\Delta_{\text{fus}}H$			35.3	482.4	DSC	[1997VER2]
	$\Delta_{\text{trs}}H$			1.96	193.6		
	$\Delta_{\text{fus}}H$			35.5	486.3		[1988SAI/ATA, 1983CHA, 1991ACR]
	$\Delta_{\text{sub}}H$	(373–395)		125.6 ± 0.8	298	ME	[2008RIB/SAN6]
	$\Delta_{\text{sub}}H$	(353–383)		116.2 ± 2.4	368	T	[1997VER2]
	$\Delta_{\text{sub}}H$			120.4 ± 2.4	298		[1997VER2]
	$\Delta_{\text{sub}}H$			113 ± 2	298	B	[1979KIM/TAK]
	$\Delta_{\text{sub}}H$			118.4	397	ME	[1967WAK/INO]
	$\Delta_{\text{sub}}H$	(333–393)		120.6	363		[1958HOY/PEP]
C ₁₈ D ₁₄	[1718-51-0]	<i>p</i> -terphenyl - d ₁₄					
	Δ_vH			101.6	298	CGC	[2008ZHA/UNH]
	Δ_vH			99.5 ± 4.4	298	CGC	[2008HAN/NUT]
C ₁₈ H ₁₄	[959-02-4]	5,12-dihydrotetracene					
	$\Delta_{\text{sub}}H$	(338–398)		115.9 ± 4	368		[1958HOY/PEP, 1970COX/PIL]
$\Delta_{\text{sub}}H$			120.5			[1951MAG/HAR, 1960JON]	
C ₁₈ H ₁₄	[2175-90-8]	diphenylfulvene					
	$\Delta_{\text{sub}}H$			104.6 ± 8.3		E	[1957DAY/OES, 1970COX/PIL]
C ₁₈ H ₁₄ F ₄ N ₂ O ₄ S	[90357-06-5]	(±) N-[4-cyano-3-(trifluoromethyl)phenyl]-3-[(4-fluorophenyl)-sulfonyl]-2-hydroxy-2-methylpropanamide (bicalutamide)					
	$\Delta_{\text{fus}}H$			53.8	469.2	DSC	[2010AND/ABU]
	$\Delta_{\text{fus}}H$ (I)			47.77	465.2		
	$\Delta_{\text{fus}}H$ (II)			43.04	462.2		[2006VEG/POL]

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
C ₁₈ H ₁₄ N ₂ O ₂	[6334-31-2] $\Delta_{\text{fus}}H$	1-benzoyl-1,2-dihydro-6-methoxy-2-quinolinecarbonitrile	23.11	396	DSC	[2005LIZ/ZAB]
C ₁₈ H ₁₄ N ₂ O ₃	[1979-24-5] $\Delta_{\text{fus}}H$	2-cyano-6-methoxy-1(2 <i>H</i>)-quinolinecarboxylic acid, phenyl ester	22.88	384	DSC	[2005LIZ/ZAB]
C ₁₈ H ₁₄ N ₄ O ₂	[21811-64-3] $\Delta_{\text{v}}H$	1,4- <i>bis</i> [(4-hydroxyphenyl)azo]benzene (473–533)	68.0	488	A	[1987STE/MAL]
C ₁₈ H ₁₄ O	[2432-11-3] $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$	2,6-diphenylphenol (334–363)	116.1 ± 1.1	348	GS	[1998VER5]
			119.1 ± 1.1	298	GS	[1998VER5]
C ₁₈ H ₁₄ O ₃	[538-56-7] $\Delta_{\text{fus}}H$	cinnamic anhydride	32.77	321.2		[1991ACR]
C ₁₈ H ₁₄ O ₃	[87205-99-0] $\Delta_{\text{fus}}H$	1,2-dihydro-1,6-dimethylphenanthro[1,2- <i>b</i>]furan-10,11-dione	22.22	490.3	DSC	[1988HUA/TAN]
C ₁₈ H ₁₅ ClN ₂ O ₂ S	[202409-33-4] $\Delta_{\text{fus}}H$	5-chloro-6'-methyl-3-[4-(methylsulfonyl)phenyl]-2,3'-bipyridine (etoricoxib)	30.43	407.1	DSC	[2008TUN/TAB]
C ₁₈ H ₁₅ F ₃ O	[172424-71-4] $\Delta_{\text{fus}}H$	4-butoxy-2',3',4'-trifluorodiphenylacetylene	36.0	344.4	DSC	[1995HSU/TSA]
C ₁₈ H ₁₅ N	[603-34-9] $\Delta_{\text{fus}}H$ $\Delta_{\text{sub}}H$ $\Delta_{\text{v}}H$	triphenylamine (322–373) (473–640)	24.89	400.2		[1993ACR]
			87.9 ± 1.3	337	BG	[1978STE3, 1987STE/MAL]
			65.2	488	A	[1987STE/MAL, 1949FOR/BOW]
C ₁₈ H ₁₅ NO ₂	[3808-37-5] $\Delta_{\text{v}}H$	N-9-anthryldiacetamide (399–455)	106.3	414	A	[1987STE/MAL]
C ₁₈ H ₁₅ NO ₂	[3808-37-5] $\Delta_{\text{sub}}H$	9-diacetylaminoanthracene	106.4		RG	[1958KLO]
C ₁₈ H ₁₅ OP	[791-28-6] $\Delta_{\text{fus}}H$ $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$	triphenylphosphine oxide	24.22	431.9		[1991ACR]
			131 ± 2	399	ME,TE	[1989HUI/VAN]
			66 ± 6	298	B	[1978JOR/AIR]
C ₁₈ H ₁₅ O ₃ P	[13291-46-8] $\Delta_{\text{fus}}H$	(2,5-dihydroxyphenyl) diphenylphosphine oxide	37.26	487.8	DSC	[2010GUO/WAN]
C ₁₈ H ₁₅ O ₄ P	[115-86-6] $\Delta_{\text{fus}}H$ $\Delta_{\text{sub}}H$ $\Delta_{\text{v}}H$	triphenyl phosphate (548–683)	29.61	322.5		[1991ACR]
			114.4 ± 2.6	298	B	[1989KIR/DOM]
			81.4	563	LA	[1987STE/MAL, 1957DOB/KEL]
C ₁₈ H ₁₅ P	[603-35-0] $\Delta_{\text{fus}}H$ $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$ $\Delta_{\text{v}}H$ $\Delta_{\text{v}}H$	triphenylphosphine (483–660) (364–392)	19.69	354.4		[1991ACR]
			113.2 ± 3.0	298		[1988KIR/DOM]
			109.2 ± 1.1	350		[1984GRI/KON]
			96.2 ± 8.4	298		[1982PIL/SKI, 1960BED/MOR]
			71.2	498	A	[1987STE/MAL, 1949FOR/BOW]
91.4 ± 2	378	TE,ME	[1981DEK/HER]			

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₈ H ₁₅ PS	[3878-45-3]	triphenylphosphine sulfide				
	$\Delta_{\text{sub}}H$	(388–419)	136.8 ± 6.1	403	HSA	[1996KIR/CHI]
	$\Delta_{\text{sub}}H$		142.8 ± 6.8	298		[1996KIR/CHI]
C ₁₈ H ₁₆ F ₂	[109970-65-2]	4- <i>n</i> -butyl-3',4'-difluorodiphenylacetylene				
	$\Delta_{\text{fus}}H$		25.3	323.5	DSC	[1995HSU/TSA]
C ₁₈ H ₁₆ N ₂ O ₂	[na]	<i>meso</i> 2,3-dimethoxy-2,3-diphenylsuccinonitrile				
	$\Delta_{\text{fus}}H$		25.1	469.7		[1983ZAM/KAI]
C ₁₈ H ₁₆ N ₄ O ₃	[243445-13-8]	2[4,5-dihydro-4-(4-methylphenyl)-5-oxo-3-(2-pyridinyl)-1,2,4-triazine-6(1 <i>H</i>)-ylidene]acetic acid, methyl ester				
	$\Delta_{\text{fus}}H$		42.8	468.3		[2005SIK/MOD]
C ₁₈ H ₁₆ N ₄	[22119-35-3]	dihydrodibenzotetra-aza-annulene				
	$\Delta_{\text{sub}}H$	(443–583)	81.5 ± 6.4	513	T	[1983ZVE/MOT]
C ₁₈ H ₁₆ O ₂	[84-47-9]	2- <i>tert</i> -butyl-9,10-anthraquinone				
	Δ_vH	(483–523)	101.4	498	A	[1987STE/MAL]
	Δ_vH		97.7			[1977SAS/FAL]
C ₁₈ H ₁₆ O ₃	[114390-57-7]	1,8-diphenyl-2,3,5-trioxabicyclo[4.3.0]non-7-ene				
	$\Delta_{\text{fus}}H$		21.7	371.2	DSC	[1991JEF/JAB]
C ₁₈ H ₁₆ O ₈	[36063-07-7]	1,2,3,4-tetracarboxymethoxynaphthalene				
	$\Delta_{\text{fus}}H$		35.9	423.7	DSC	[1993ACR]
C ₁₈ H ₁₆ O ₈	[68267-09-4]	1,2,4,5-tetracarboxymethoxynaphthalene				
	$\Delta_{\text{fus}}H$		36.4	438.2	DSC	[1993ACR]
C ₁₈ H ₁₆ O ₈	[36063-08-8]	1,2,5,6-tetracarboxymethoxynaphthalene				
	$\Delta_{\text{fus}}H$		42.1	470.2	DSC	[1993ACR]
C ₁₈ H ₁₆ O ₈	[68267-08-3]	1,2,6,7-tetracarboxymethoxynaphthalene				
	$\Delta_{\text{fus}}H$		34.2	407.2	DSC	[1993ACR]
C ₁₈ H ₁₆ O ₈	[56110-97-5]	2,3,6,7-tetracarboxymethoxynaphthalene				
	$\Delta_{\text{fus}}H$		42.2	458.2	DSC	[1993ACR]
C ₁₈ H ₁₆ O ₈	[31996-10-8]	1,4,5,8-tetracarboxymethoxynaphthalene				
	$\Delta_{\text{fus}}H$		36.1	477.2	DSC	[1993ACR]
C ₁₈ H ₁₇ Cl ₂ NO ₃	[22212-55-1]	ethyl <i>N</i> -benzoyl- <i>N</i> -(3,4-dichlorophenyl)-(<i>dl</i>)-alaninate				
	$\Delta_{\text{fus}}H$		27.06	341.7	DSC	[1990DON/DRE]
C ₁₈ H ₁₇ F	[109970-63-0]	4- <i>n</i> -butyl-4'-fluorodiphenylacetylene				
	$\Delta_{\text{fus}}H$		18.5	329.9	DSC	[1995HSU/TSA]
C ₁₈ H ₁₇ FO	[130746-61-1]	4-butoxy-4'-fluorodiphenylacetylene				
	$\Delta_{\text{fus}}H$		25.4	346.7	DSC	[1995HSU/TSA]
C ₁₈ H ₁₇ NO ₃	[483362-79-4]	1-[(4-nitrophenyl)ethynyl]-4-butoxybenzene				
	$\Delta_{\text{fus}}H$		21.84	374.7	DSC	[2002SPA/DZI]
C ₁₈ H ₁₇ NO ₅	[53902-12-8]	2-[[3-(3,4-dimethoxyphenyl)-1-oxo-2-propenyl]amino]benzoic acid (tranilast)				
	$\Delta_{\text{fus}}H$		48.87	486.4		[2005VOG/COH]
C ₁₈ H ₁₈	[18801-00-8]	2-(<i>tert</i> -butyl)anthracene				
	Δ_vH	(323–473)	84.5	398	GC	[2002LEI/CHA]
C ₁₈ H ₁₈	[1498-69-7]	9-butylanthracene				
	$\Delta_{\text{sub}}H$	(293–313)	108.1	303	A	[1987STE/MAL, 1964MOR]

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
	Enthalpy							
		$\Delta_{\text{v}}H$	(422–492)	77.1	437		A	[1987STE/MAL]
		$\Delta_{\text{v}}H$	(328–373)	83.9	343		A	[1987STE/MAL, 1964MOR]
C ₁₈ H ₁₈	[483-65-8]	1-methyl-7-isopropylphenanthrene						
		$\Delta_{\text{fus}}H$		18.03	369			[1996DOM/HEA]
C ₁₈ H ₁₈		$\Delta_{\text{v}}H$	(539–678)	54.0	554		A	[1987STE/MAL]
	[7396-38-5]	2,4,5,7-tetramethylphenanthrene						
		$\Delta_{\text{sub}}H$		114.2 ± 1.7			ME	[1965MCD/KIL, 1970COX/PIL]
C ₁₈ H ₁₈	[7343-06-8]	3,4,5,6-tetramethylphenanthrene						
		$\Delta_{\text{sub}}H$		133.5 ± 3.8			ME	[1965MCD/KIL, 1970COX/PIL]
C ₁₈ H ₁₈ CIN ₃	[113-59-7]	3-(2-chloro-9 <i>H</i> -thioxanthen-9-ylidene)- <i>N,N</i> -dimethyl-1-propanamine (2-chloroprothixene)						
		$\Delta_{\text{fus}}H$		27.82	370.3			[1996DOM/HEA]
		$\Delta_{\text{fus}}H$		28.9	370.5		DTA	[1983MAS/CHA]
C ₁₈ H ₁₈ F ₃ N ₃ O ₃	[15168-05-9]	3-{3,5-dimethyl-4-[3-(3-methylisoxazol-5-yl)propoxy]phenyl}-5-trifluoromethyl[1,2,4]oxodiazole (pleconaril)						
		$\Delta_{\text{fus}}H$ (I)		29.3	336.5			
		$\Delta_{\text{fus}}H$ (II)		32.7	333.4		DSC	[2004COS/SCH]
C ₁₈ H ₁₈ N ₂ O ₂	[na]	<i>N,N'</i> -(2-hydroxyethyl)-1,4-diaminoanthraquinone						
		$\Delta_{\text{fus}}H$		32.34	521.2			[1988BAU/PER]
C ₁₈ H ₁₈ O ₂	[19672-37-8]	3-diphenylmethyl-2,4-pentanedione						
		$\Delta_{\text{fus}}H$		27.02	387.2			[1995NOL/VER]
		$\Delta_{\text{sub}}H$	(348–383)	112.8 ± 0.4	366		T	[1995NOL/VER]
C ₁₈ H ₁₈ O ₃	[na]	butyl 9-hydroxy-9 <i>H</i> -fluorene-9-carboxylate						
		$\Delta_{\text{fus}}H$		25.56	343.9			[1991ACR]
C ₁₈ H ₁₈ O ₄	[25062-95-7]	2,2'-diphenyl-bi-(1,3-dioxolan-2-yl)						
		$\Delta_{\text{fus}}H$		32.1	456.1			[1995VER/DOG]
		$\Delta_{\text{sub}}H$	(320–362)	132.8 ± 2.1	341		T	[1995VER/DOG]
C ₁₈ H ₁₈ O ₁₂	[6237-59-8]	hexamethoxycarbonylbenzene						
		$\Delta_{\text{fus}}H$		22.5	463.7		DSC	[1978DOZ/FUJ]
		$\Delta_{\text{sub}}H$	(403–422)	140.7 ± 1.1	413		ME	[1995JIM/MEN]
		$\Delta_{\text{sub}}H$		154.3 ± 1.2	298			[1995JIM/MEN]
C ₁₈ H ₁₉ BrO	[556052-88-1]	4-bromo-4'-(5-hexenyloxy)-1,1'-biphenyl						
		$\Delta_{\text{fus}}H$		13.8	308.2			
		$\Delta_{\text{fus}}H$		16.0	393.9		DSC	[2003WIL/VAN]
C ₁₈ H ₁₉ CIN ₄	[5786-21-0]	8-chloro-11-(4-methyl-1-piperazinyl)-5 <i>H</i> -dibenzo[<i>b,e</i>] [1,4]diazepine (clozapine)						
		$\Delta_{\text{fus}}H$		35.9	457.1		DSC	[2006WAS/HOL]
C ₁₈ H ₁₉ Cl ₂ NO ₄	[72509-76-3]	(±) ethyl methyl 1,4-dihydro-2,6-dimethyl-4-(2,3-dichlorophenyl)-3,5-pyridine dicarboxylate (felodipine)						
		$\Delta_{\text{fus}}H$ (I)		31.5	417.4			
		$\Delta_{\text{fus}}H$ (II)		26.7	405.7			[2001ROL/BUR]
		$\Delta_{\text{fus}}H$		34.8	412.3			[2007BER/WAS]
		$\Delta_{\text{fus}}H$		35.21	414.9		DSC	[1992SRC/KER]
C ₁₈ H ₁₉ Cl ₂ NO ₄	[119945-59-4]	(+) ethyl methyl 1,4-dihydro-2,6-dimethyl-4-(2,3-dichlorophenyl)-3,5-pyridine dicarboxylate (felodipine)						
		$\Delta_{\text{fus}}H$		25.4	415.7			[2001ROL/BUR]

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
	Enthalpy							
C ₁₈ H ₁₉ N	[na]	(S) 4-(2-methylbutyl)-4'-cyanobiphenyl						
	$\Delta_{\text{fus}}H$			10.7	276			[1999MAY/WIT]
		Note: Compound may have phase transitions at lower temperatures.						
C ₁₈ H ₁₉ NO ₄	[483362-68-1]	2-(4-nitrophenyl)-1-(4-butoxyphenyl)ethanone						
	$\Delta_{\text{fus}}H$			24.94	336.1		DSC	[2002SPA/DZI]
C ₁₈ H ₂₀	[2913-24-8]	[3.3]para-cyclophane						
	$\Delta_{\text{trs}}H$			7.36	332			
	$\Delta_{\text{trs}}H$			0.46	351			
	$\Delta_{\text{fus}}H$			11.76	377			[1969SHI/MCN]
	$\Delta_{\text{sub}}H$		(322–343)	103.3 ± 1	332		ME	[1969SHI/MCN, 1977PED/RYL]
			(321–343)	97.8	332		A	[1987STE/MAL]
C ₁₈ H ₂₀	[115181-05-0]	6-(4-biphenyl)-1-hexene						
	$\Delta_{\text{fus}}H$			15.1	274.5		DSC	[1989MAL/KAN]
C ₁₈ H ₂₀ BrN ₅ O ₅	[191355-38-1]	8-bromo-(R)-7-[2-hydroxy-3-(4-acetylamino)-fenoxypropyl]-1,3-dimethylxanthine						
	$\Delta_{\text{fus}}H$			37.8	492.2		DSC	[2000DAN/PRO]
C ₁₈ H ₂₀ Cl ₂	[72-56-0]	1,1'-(2,2-dichloroethylidene)bis(4-ethylbenzene)						
	$\Delta_{\text{fus}}H$			23.34	331.6		DSC	[1991ACR, 1990DON/DRE]
C ₁₈ H ₂₀ N ₂ O ₂	[3955-57-5]	N,N'-bis(salicylaldehyde)tetramethylenediimine						
	$\Delta_{\text{fus}}H$			35.54	362.8		DSC	[2004RIB/GON]
	$\Delta_{\text{sub}}H$		(349–361)	165.1 ± 3.1	298		ME	[2004RIB/GON]
C ₁₈ H ₂₀ N ₂ O ₄	[na]	diethyl 2,4,6,8-tetrahydro-4,8-ethanobenzo[1,2-c:4,5-c']dipyrrole-1,7-dicarboxylate						
	$\Delta_{\text{fus}}H$			18.2	439.3			[2000UNO/ITO]
C ₁₈ H ₂₀ N ₂ O ₄	[54946-24-6]	N-pentylthalidomide						
	$\Delta_{\text{fus}}H$			23.97	378.2		DTA	[2002GOO/LAI]
C ₁₈ H ₂₀ N ₂ O ₆	[na]	3-ethyl-5-methyl-1,4-dihydro-2,6-dimethyl-4-(3-nitrophenyl)-3,5-pyridinedicarboxylate ((RS)-nitrendipine)						
	$\Delta_{\text{fus}}H$			41.1	430.7			[1997BUR/ROL]
C ₁₈ H ₂₀ N ₆ O ₇	[191355-39-2]	8-nitro-(R)-7-[2-hydroxy-3-(4-acetylamino)-fenoxypropyl]-1,3-dimethylxanthine						
	$\Delta_{\text{fus}}H$			56.8	481.2		DSC	[2000DAN/PRO]
C ₁₈ H ₂₀ OS	[556052-90-5]	4'-(5-hexenyloxy)-[1,1'-biphenyl]-4-thiol						
	$\Delta_{\text{trs}}H$			11.6	358.4			
	$\Delta_{\text{fus}}H$			13.6	384.6		DSC	[2003WIL/VAN]
C ₁₈ H ₂₀ O ₂	[56-53-1]	diethylstilbestrol						
	$\Delta_{\text{fus}}H$			31.76	443.8		DSC	[1990DON/DRE]
C ₁₈ H ₂₀ O ₂	[100923-74-8]	(2-hydroxyl-4,6-dimethylphenyl)-2,4,6-trimethylphenylmethanone						
	$\Delta_{\text{fus}}H$			0.84	380.2		DTA	[1989SAL/ABA]
								Note: Reported enthalpy of fusion is too small, and the published enthalpy and entropy of fusion data are internally inconsistent
C ₁₈ H ₂₀ O ₄	[na]	2-hydroxy-4-butoxy-4'-methoxybenzophenone						
	$\Delta_{\text{fus}}H$			33.7	345.6		DSC	[1999PRI/HAWN]
	$\Delta_{\text{sub}}H$			126.3			B	[1999PRI/HAWN]
C ₁₈ H ₂₁ N	[na]	N-benzyl-pivalophenone imine						
	$\Delta_{\text{fus}}H$			27.86	339.6			[1997VER/MOR]
	$\Delta_{\text{sub}}H$			109.7 ± 3.3	298		B	[1997VER/MOR]

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
	Enthalpy							
C ₁₈ H ₂₁ NO	[97402-82-9]	(E)-N-(4-methoxybenzylidene)-4-butylaniline			na			[1989KRE/AZA]
C ₁₈ H ₂₁ NO ₃	[76-57-3]	7,8-didehydro-4,5-epoxy-3-methoxy-17-ethylmorphanan-6-ol (codeine)			23.81	430.3		[1995YAN/YIN]
					18.28	428.2	DTA	[1988ROY/FLY]
C ₁₈ H ₂₁ N ₃ S	[69186-17-0]	N-(diethylaminothiocarbonyl)-N'-phenylbenzamidine			159.4 ± 3.3	298	C	[2004RIB/SAN]
C ₁₈ H ₂₁ N ₅ O ₅	[191355-37-0]	(R)-7-[2-hydroxy-3-(4-acetylamino)-fenoxypropyl]-1,3-dimethylxanthine			65.9	477.2	DSC	[2000DAN/PRO]
C ₁₈ H ₂₂	[1087-49-6]	1,6-diphenylhexane			88.0	308	A	[1987STE/MAL, 1964MOR]
		(293–373)						
C ₁₈ H ₂₂	[na]	2,2-di(<i>p</i> -tolyl)butane			85.4	313		[1999DYK/SVO]
		(298–473)						
C ₁₈ H ₂₂	[na]	1- <i>p</i> -tolyl-(1- <i>p</i> -propylphenyl)ethane			85.4	313		[1999DYK/SVO]
		(298–473)						
C ₁₈ H ₂₂	[na]	1- <i>o</i> -tolyl- <i>p</i> -tolylbutane			85.4	313		[1999DYK/SVO]
		(298–473)						
C ₁₈ H ₂₂	[1889-67-4]	2,3-dimethyl-2,3-diphenylbutane			25.52	392	DSC	[1983KRA/BEC]
		(293–348)			96.7 ± 0.8	320		[1983KRA/BEC]
C ₁₈ H ₂₂ FNO ₃	[34809-52-2]	(R)-deoxyephedrinium (S)-4'-fluoromandelate			27.9	374.5	DSC	[2001VAL/SMI]
C ₁₈ H ₂₂ FNO ₃	[348098-53-3]	(R)-deoxyephedrinium (R)-4'-fluoromandelate			26.6	369.2	DSC	[2001VAL/SMI]
C ₁₈ H ₂₂ FNO ₄	[174966-70-2]	(1R,2S)-ephedrinium (R)-4'-fluoromandelate			37.5	438.3	DSC	[2001VAL/SMI]
C ₁₈ H ₂₂ FNO ₄	[174966-62-2]	(1R,2S)-ephedrinium (S)-4'-fluoromandelate			24.5	380.4	DSC	[2001VAL/SMI]
C ₁₈ H ₂₂ N ₂ O ₂	[100046-00-2]	2,2',4,4',6,6'-hexamethylazobenzene-N,N-dioxide			107 ± 12	298	ME	[1993ACR/TUC2]
C ₁₈ H ₂₂ N ₄	[133930-64-0]	2,3-dimethyl-2,3- <i>bis</i> (phenylazo)butane			21.09	342.3		[1993ENG/WAN]
					113.8 ± 1.8		B	[1993ENG/WAN]
C ₁₈ H ₂₂ N ₄	[na]	<i>trans, trans</i> -1,6-diphenyl-3,3,4,4-tetramethyl-1,2,5,6-tetraazahexane			92.8 ± 1.5	368	GS	[1993ENG/WAN]
		(348–388)						
C ₁₈ H ₂₂ O ₂	[41047-48-7]	(<i>dl</i>)-2,3-dimethoxy-2,3-diphenylbutane			114.2 ± 6.3	339		[1990DOG/BEC]
		(322–355)						
C ₁₈ H ₂₂ O ₂	[na]	(<i>dl</i>) anisylidenecamphor			26.36	371.5		[1976LEC/COL]
C ₁₈ H ₂₂ O ₂	[na]	(<i>d</i>) anisylidenecamphor			30.12	399.5		[1976LEC/COL]
C ₁₈ H ₂₂ O ₂	[80-43-3]	di- α -cumyl peroxide						

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
	Enthalpy							
		$\Delta_{\text{fus}}H$			28.14	312.4		[1996DOM/HEA]
C ₁₈ H ₂₂ O ₄	[39787-30-9]		1,2-diphenyl-1,1,2,2-tetramethoxyethane					
		$\Delta_{\text{fus}}H$			20.1	328.5		[1995VER/DOG]
		$\Delta_{\text{sub}}H$	(351–399)		77.6 ± 0.6	375	T	[1995VER/DOG]
C ₁₈ H ₂₂ O ₄	[11921-29-1]		4,4'-di-(2-methoxyethoxy)biphenyl					
		$\Delta_{\text{trs}}H$			17.53	409.5		
		$\Delta_{\text{fus}}H$			22.67	412.4	DSC	[1995BOW/HER]
C ₁₈ H ₂₃ FO ₂	[157396-75-3]		4- <i>trans</i> -(4-fluorophenylethyl)cyclohexyl (E)-butenoate					
		$\Delta_{\text{fus}}H$			25.0	335.2		[1995KEL/SCH]
C ₁₈ H ₂₄	[1610-22-6]		1,2,3,4,4a,7,8,9,10,12,12a-dodecahydrochrysene					
		$\Delta_{\text{sub}}H$	(293–313)		115.4	303	A	[1987STE/MAL, 1964MOR]
		$\Delta_{\text{v}}H$	(318–358)		84.2	333	A	[1987STE/MAL, 1947STU]
C ₁₈ H ₂₄	[83171-44-2]		(E)-9-(bicyclo[4.2.1]non-3-en-9-ylidene)bicyclo[4.2.1]non-3-ene					
		$\Delta_{\text{fus}}H$			9.1	441.7	DSC	[1984MAR/MEL]
C ₁₈ H ₂₄	[83171-45-3]		(Z)-9-(bicyclo[4.2.1]non-3-en-9-ylidene)bicyclo[4.2.1]non-3-ene					
		$\Delta_{\text{fus}}H$			27.96	440.8	DSC	[1984MAR/MEL]
C ₁₈ H ₂₄ N ₂ O ₄	[138516-98-0]		(4-nitrophenyl)-10-undecynyl carbamate					
		$\Delta_{\text{fus}}H$			53.18	385.4	DSC	[1993TIE/FRA]
C ₁₈ H ₂₄ N ₂ O ₆	[na]		3-[(hydroxyimino)(4-methoxy-3-nitrophenyl)methyl]-1,2,2-trimethylcyclopentanecarboxylic acid methyl ester					
		$\Delta_{\text{fus}}H$			31.99	433		[1995NUR/LEL]
C ₁₈ H ₂₄ O	[na]		(1R,1''R,6S,6''S,9S,9''S)-dispiro(bicyclo[4.2.1]non-3-ene-9,2'-oxirane-3',9''-bicyclo[4.2.1]non-3-ene)					
		$\Delta_{\text{trs}}H$			5.92	354.3		
		$\Delta_{\text{fus}}H$			8.19	373.8	DSC	[1984MAR/MEL]
C ₁₈ H ₂₄ O	[na]		(1R,1''R,6S,6''S,9R,9''S)-dispiro(bicyclo[4.2.1]non-3-ene-9,2'-oxirane-3',9''-bicyclo[4.2.1]non-3-ene)					
		$\Delta_{\text{fus}}H$			14.92	333.9	DSC	[1984MAR/MEL]
C ₁₈ H ₂₄ O ₃	[104225-40-3]		3-(4-ethylbenzoyl)-1,2,2-trimethylcyclopentanecarboxylic acid					
		$\Delta_{\text{fus}}H$			22.54	387.6	DSC	[1992TER/PAU]
C ₁₈ H ₂₄ O ₃	[104225-33-4]		3-(3,4-dimethylbenzoyl)-1,2,2-trimethylcyclopentanecarboxylic acid					
		$\Delta_{\text{fus}}H$			32.31	460.6	DSC	[1992TER/PAU]
C ₁₈ H ₂₄ O ₃	[104225-36-7]		3-(2,4-dimethylbenzoyl)-1,2,2-trimethylcyclopentanecarboxylic acid					
		$\Delta_{\text{fus}}H$			18.81	386.8	DSC	[1992TER/PAU]
C ₁₈ H ₂₄ O ₄	[57403-79-7]		3-(4-ethoxybenzoyl)-1,2,2-trimethylcyclopentanecarboxylic acid					
		$\Delta_{\text{fus}}H$			22.05	394.6	DSC	[1992TER/PAU]
C ₁₈ H ₂₄ O ₄	[84-64-0]		butylcyclohexylphthalate					
		$\Delta_{\text{v}}H$	(368–485)		94.3	383	A	[1987STE/MAL]
C ₁₈ H ₂₅ NO ₃	[104225-18-5]		3-[4-(dimethylamino)benzoyl]-1,2,2-trimethylcyclopentanecarboxylic acid					
		$\Delta_{\text{fus}}H$			25.03	445	DSC	[1992TER/PAU]
C ₁₈ H ₂₅ NO ₄	[na]		3-[(hydroxyimino)(4-methoxyphenyl)methyl]-1,2,2-trimethylcyclopentanecarboxylic acid methyl ester					
		$\Delta_{\text{fus}}H$			36.99	433		[1995NUR/LEL]
C ₁₈ H ₂₆ N ₂ O ₄	[138516-97-9]		(4-nitrophenyl)-10-undecenyl carbamate					
		$\Delta_{\text{fus}}H$			43.81	376.4	DSC	[1993TIE/FRA]

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
	Enthalpy							
C ₁₈ H ₂₆ O ₄	[606-50-5] $\Delta_{\text{v}}H$	diisopentyl phthalate		(390–610)	81.6	405	A	[1987STE/MAL]
C ₁₈ H ₂₆ O ₄	[131-18-0] $\Delta_{\text{v}}H$	dipentyl phthalate		(323–390)	87.3	338	T	[1949PER/WEB]
	$\Delta_{\text{v}}H$			(303–500)	99.4	318	A, ME	[1987STE/MAL, 1948SMA/SMA]
C ₁₈ H ₂₇ BrN ₂ O ₄	[138517-07-4] $\Delta_{\text{fus}}H$	(4-nitrophenyl)-11-bromoundecyl carbamate			58.56	395.6	DSC	[1993TIE/FRA]
C ₁₈ H ₂₇ IN ₂ O ₄	[138517-10-9] $\Delta_{\text{fus}}H$	(4-nitrophenyl)-11-iodoundecyl carbamate			63.34	399.6	DSC	[1993TIE/FRA]
C ₁₈ H ₂₈	[66553-12-6] $\Delta_{\text{v}}H$	1,2,3,4-tetrahydro-6-octylnaphthalene		(503–574)	103.3	538		[1999DYK/SVO]
C ₁₈ H ₂₈	[83171-46-4] $\Delta_{\text{trs}}H$	(E)-9-(bicyclo[4.2.1]nonan-9-ylidene)bicyclo[4.2.1]nonane			2.68	224.8		
	$\Delta_{\text{fus}}H$				9.32	412.4	DSC	[1984MAR/MEL]
C ₁₈ H ₂₈	[83171-47-5] $\Delta_{\text{trs}}H$	(Z)-9-(bicyclo[4.2.1]nonan-9-ylidene)bicyclo[4.2.1]nonane			10.12	344.4		
	$\Delta_{\text{fus}}H$				11.04	393.3	DSC	[1984MAR/MEL]
C ₁₈ H ₂₈ N ₂ O	[27262-45-9] $\Delta_{\text{fus}}H$	N-(2,6-dimethylphenyl)-1-butyl-2-piperidinecarboxamide			26.25	413.2	DSC	[1997NEM/ACS]
C ₁₈ H ₂₈ O	[na] $\Delta_{\text{trs}}H$	(1R,1''R,6S,6''S,9R,9''R)-dispiro(bicyclo[4.2.1]nonane-9,2'-oxirane-3',9''-bicyclo[4.2.1]nonane)			9.74	348.5		
	$\Delta_{\text{fus}}H$				13.13	444.1	DSC	[1984MAR/MEL]
C ₁₈ H ₂₈ O	[na] $\Delta_{\text{trs}}H$	(1R,1''R,6S,6''S,9R,9''S)-dispiro(bicyclo[4.2.1]nonane-9,2'-oxirane-3',9''-bicyclo[4.2.1]nonane)			0.72	274.9		
	$\Delta_{\text{fus}}H$				6.35	342.5	DSC	[1984MAR/MEL]
C ₁₈ H ₂₈ O ₄	[118476-22-5] $\Delta_{\text{trs}}H$	2,5-di- <i>n</i> -hexyloxy-1,4-benzoquinone			5.3	332.3		
	$\Delta_{\text{fus}}H$				38.9	412.1	DSC	[1996KEE/VAN]
C ₁₈ H ₂₈ S ₈	[106920-28-9] $\Delta_{\text{fus}}H$	2-[4,5- <i>bis</i> (propylthio)-1,3-dithiol-2-ylidene]-4,5- <i>bis</i> (propylthio)-1,3-dithiole			42.7	306.6	AC	[1997TAN/ATA]
C ₁₈ H ₂₉ NO	[24973-59-9] $\Delta_{\text{sub}}H$	2,4,6-tri- <i>tert</i> -butylnitrosobenzene			91.0 ± 3.2	298	C	[1995ACR/BOT]
C ₁₈ H ₂₉ NO ₂	[4074-25-3] $\Delta_{\text{fus}}H$	2,4,6-tri- <i>tert</i> -butylnitrobenzene			19.25	482.8	DSC	[2000VER/HEI]
	$\Delta_{\text{sub}}H$	(333–368)			94.8 ± 1.0	351	GS	[2000VER/HEI]
	$\Delta_{\text{sub}}H$	(333–368)			96.4 ± 1.0	298	GS	[2000VER/HEI]
	$\Delta_{\text{sub}}H$				81.4 ± 1.8	298	C	[1995ACR/BOT]
C ₁₈ H ₃₀	[123-01-3] $\Delta_{\text{v}}H$	dodecylbenzene			92	275		[2000MOK/RUZ]
	$\Delta_{\text{v}}H$	(333–453)			83.2	348		[1993KAS/MOK]
	$\Delta_{\text{v}}H$	(496–609)			67.4	511	A	[1987STE/MAL]
	$\Delta_{\text{v}}H$	(336–456)			80.6	356	GS	[1986ALL/JOS]
	$\Delta_{\text{v}}H$				89.6	298		[1971WIL/ZWO]
C ₁₈ H ₃₀	[2090-14-4] $\Delta_{\text{v}}H$	perhydrochrysene		(273–353)	82.4	288		[1964MOR]

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
C ₁₈ H ₃₀	[604-88-6]	hexaethylbenzene				
	$\Delta_{\text{sub}}H$	(327–352)	95.0 ± 4.0	340	HSA	[1986CHI/ANN]
	$\Delta_{\text{sub}}H$		U 41.3 ± 0.9		DSC	[1984HOL]
	$\Delta_{\text{v}}H$	(407–572)	62.6	422	A	[1987STE/MAL, 1947STU]
C ₁₈ H ₃₀	[635-11-0]	1,2,4,5-tetraisopropylbenzene				
	$\Delta_{\text{fus}}H$		19.6	393		[2002STE/CHI6]
	$\Delta_{\text{v}}H$	(410–575)	61.1 ± 0.3	420	EB	[2002STE/CHI6]
	$\Delta_{\text{v}}H$	(410–575)	56.8 ± 0.3	460	EB	[2002STE/CHI6]
	$\Delta_{\text{v}}H$	(410–575)	52.3 ± 0.5	500	EB	[2002STE/CHI6]
	$\Delta_{\text{v}}H$	(410–575)	47.5 ± 0.9	540	EB	[2002STE/CHI6]
C ₁₈ H ₃₀	[1460-02-2]	1,3,5-tri- <i>tert</i> -butylbenzene				
	$\Delta_{\text{sub}}H$	(298–341)	79.9 ± 0.3	319	T	[1998VER]
	$\Delta_{\text{sub}}H$		81.2 ± 0.3	298		[1998VER]
	$\Delta_{\text{sub}}H$	(273–315)	79.7 ± 0.4	294	ME	[1965DAV/KYB, 1987STE/MAL]
C ₁₈ H ₃₀	[2090-14-4]	perhydrochrysene				
	$\Delta_{\text{sub}}H$	(273–353)	82.3 (liq)	288	A	[1987STE/MAL, 1964MOR]
C ₁₈ H ₃₀	[123-01-3]	1-phenyldodecane				
	$\Delta_{\text{fus}}H$		43.1	274.6	DSC	[2000MOK/RUZ]
	$\Delta_{\text{sub}}H$		135.1	298		[2000MOK/RUZ]
C ₁₈ H ₃₀ N ₄ O ₂	[126235-05-0]	8-undecyltheophylline				
	$\Delta_{\text{fus}}H$		25.8	433.5	DSC	[1989GON/KRA, 1991ACR]
C ₁₈ H ₃₀ O	[732-26-3]	2,4,6-tri- <i>tert</i> -butylphenol				
	$\Delta_{\text{fus}}H$		19.46	405.2		[1991CHI/BRA]
	$\Delta_{\text{sub}}H$		87.5 ± 0.4	298	GS	[1999VER]
	$\Delta_{\text{sub}}H$	(295–339)	85.6 ± 0.4	317	ME	[1965DAV/KYB, 1987STE/MAL]
	$\Delta_{\text{sub}}H$		U 128.1	298	C	[1971BER/GIR, 1999VER]
	$\Delta_{\text{sub}}H$	(292–313)	83.9	302		[1960AIH]
	$\Delta_{\text{sub}}H$		84.2 ± 0.5	298	V	[1960AIH, 1999VER]
	$\Delta_{\text{v}}H$	(415–551)	63.2	430	A	[1987STE/MAL]
C ₁₈ H ₃₀ O ₂	[59968-12-6]	1,3-dimethoxy-4-decylbenzene				
	$\Delta_{\text{v}}H$	(443–493)	76.6	458	A, GC	[1987STE/MAL, 1975KUN/LIL]
C ₁₈ H ₃₀ O ₂	[41442-52-8]	1,3-dimethoxy-5-decylbenzene				
	$\Delta_{\text{v}}H$	(459–519)	78.4	474	A, GC	[1987STE/MAL, 1975KUN/LIL]
C ₁₈ H ₃₀ O ₄	[47189-08-2]	1,4- <i>bis</i> (1,1-diethoxyethyl)benzene				
	$\Delta_{\text{v}}H$	(329–347)	88.5	338	A	[1987STE/MAL]
C ₁₈ H ₃₀ O ₄	[47189-08-2]	4-diacetylbenzene diethyl ketal				
	$\Delta_{\text{trs}}H$		1.31	168.2		
	$\Delta_{\text{fus}}H$		23.5	326.2		[1996DOM/HEA]
	$\Delta_{\text{sub}}H$	(306–327)	112.5	316.5		[1978KAR/KAM, 1987STE/MAL]
C ₁₈ H ₃₀ O ₄	[849-30-4]	dicyclohexyl adipate				
	$\Delta_{\text{v}}H$	(338–369)	106.3 ± 1.5	298	GS	[2008LIP/KRA]
C ₁₈ H ₃₀ O ₆	[7568-58-3]	<i>trans</i> aconitic acid, tributyl ester				
	$\Delta_{\text{v}}H$	(385–483)	87.4	400	A	[1987STE/MAL]

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₈ H ₃₀ O ₆	[na] $\Delta_v H$	diethylene glycol dicarboxylic acid, di[1-(isopropoxycarbonyl)ethyl] ester (418–493)	97.6	433	A	[1987STE/MAL]
C ₁₈ H ₃₀ O ₆	[na] $\Delta_v H$	diethylene glycol dicarboxylic acid, di[1-(propoxycarbonyl)ethyl] ester (418–514)	101.5	433	A	[1987STE/MAL]
C ₁₈ H ₃₁ N	[961-38-6] $\Delta_{\text{fus}} H$	2,4,6-tri- <i>tert</i> -butylaniline	19.38	426.4		[2000VER3]
	$\Delta_{\text{sub}} H$		92.5 ± 1.1	298	GS	[2000VER3]
C ₁₈ H ₃₂	[55133-89-6] $\Delta_v H$	9-butyltetradecahydroanthracene (420–456)	72.8	435	A	[1987STE/MAL]
C ₁₈ H ₃₂	[2456-43-1] $\Delta_v H$	1,2-dicyclohexylcyclohexane (375–563)	72.8	390	A	[1987STE/MAL]
C ₁₈ H ₃₂ O	[1604-32-6] $\Delta_v H$	6,10,14-trimethyl-3,5-pentadecadien-2-one (404–560)	43.4 ± 0.5	482		[1988BAG/GUR]
C ₁₈ H ₃₂ O ₂	[506-21-8] $\Delta_{\text{fus}} H$	linoelaidic acid	47.7	303	TA	[1982JAL/ZOG]
C ₁₈ H ₃₂ O ₂	[19307-18-7] $\Delta_{\text{fus}} H$	4-octadecyonic acid	57.94	348	TA	[1982JAL/ZOG]
C ₁₈ H ₃₂ O ₂	[676-30-2] $\Delta_{\text{fus}} H$	5-octadecyonic acid	54.41	325	TA	[1982JAL/ZOG]
C ₁₈ H ₃₂ O ₂	[544-74-1] $\Delta_{\text{fus}} H$	6-octadecyonic acid	54.92	324	TA	[1982JAL/ZOG]
C ₁₈ H ₃₂ O ₂	[19220-35-0] $\Delta_{\text{fus}} H$	7-octadecyonic acid	53.61	322	TA	[1982JAL/ZOG]
C ₁₈ H ₃₂ O ₂	[19220-36-1] $\Delta_{\text{fus}} H$	8-octadecyonic acid	55.3	320	TA	[1982JAL/ZOG]
C ₁₈ H ₃₂ O ₂	[506-24-1] $\Delta_{\text{fus}} H$	9-octadecyonic acid	54.87	319	TA	[1982JAL/ZOG]
C ₁₈ H ₃₂ O ₂	[19220-39-4] $\Delta_{\text{fus}} H$	10-octadecyonic acid	52.32	319	TA	[1982JAL/ZOG]
C ₁₈ H ₃₂ O ₂	[19220-40-7] $\Delta_{\text{fus}} H$	11-octadecyonic acid	55.97	320	TA	[1982JAL/ZOG]
C ₁₈ H ₃₂ O ₂	[19220-41-8] $\Delta_{\text{fus}} H$	12-octadecyonic acid	49.79	320	TA	[1982JAL/ZOG]
C ₁₈ H ₃₂ O ₂	[19220-42-9] $\Delta_{\text{fus}} H$	13-octadecyonic acid	55.51	322	TA	[1982JAL/ZOG]
C ₁₈ H ₃₂ O ₂	[34494-26-3] $\Delta_{\text{fus}} H$	14-octadecyonic acid	52.74	337	TA	[1982JAL/ZOG]
C ₁₈ H ₃₂ O ₂	[34450-17-4] $\Delta_{\text{fus}} H$	16-octadecyonic acid	60.1	347	TA	[1982JAL/ZOG]
C ₁₈ H ₃₂ O ₂	[34450-18-5] $\Delta_{\text{fus}} H$	17-octadecyonic acid	54.2	340	TA	[1982JAL/ZOG]
C ₁₈ H ₃₂ O ₂	[112-79-8]	1,10-cyclooctadecanedione				

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
	Enthalpy							
		$\Delta_{\text{trs}}H$			11.84	359.2		
		$\Delta_{\text{fus}}H$			27.03	371.2		[1972ALV/BOR]
C ₁₈ H ₃₂ O ₄	[na]		1,8-cyclotetradecanedione bis ethylene ketal					
		$\Delta_{\text{fus}}H$			30.67	457.2		[1972ALV/BOR]
C ₁₈ H ₃₂ O ₆	[38094-11-0]		tributyl 1,2,3-propanetricarboxylate				A	[1987STE/MAL]
		$\Delta_{\text{v}}H$	(385–482)		87.8	400		
C ₁₈ H ₃₄	[629-89-0]		1-octadecyne					
		$\Delta_{\text{v}}H$	(450–623)		64.9	465		[1999DYK/SVO]
C ₁₈ H ₃₄	[61847-97-0]		2-octadecyne					
		$\Delta_{\text{v}}H$	(458–633)		65.7	473		[1999DYK/SVO]
C ₁₈ H ₃₄	[61886-64-4]		3-octadecyne					
		$\Delta_{\text{v}}H$	(449–622)		64.5	464		[1999DYK/SVO]
C ₁₈ H ₃₄	[95049-67-5]		1,4-dipentylbicyclo[2.2.2]octane					
		$\Delta_{\text{fus}}H$			20.4	261.5		[1999DOU/BOT]
C ₁₈ H ₃₄	[1610-23-7]		1,6-dicyclohexylhexane				A	[1987STE/MAL, 1964MOR]
		$\Delta_{\text{v}}H$	(288–373)		85.9	303		
C ₁₈ H ₃₄ O	[6907-37-5]		cyclooctadecanone					
		$\Delta_{\text{sub}}H$			77.4			[1938WOL/WEG, 1960JON]
C ₁₈ H ₃₄ O ₂	[593-39-5]		<i>cis</i> -6-octadecenoic acid					
		$\Delta_{\text{fus}}H$			59.9	302.3		[2007MOO/KOE]
		$\Delta_{\text{fus}}H$			47.5	303.7		[1996DOM/HEA]
C ₁₈ H ₃₄ O ₂	[112-80-1]		<i>cis</i> 9-octadecenoic acid					
		$\Delta_{\text{fus}}H$			39.6	286.5		[1996DOM/HEA]
		$\Delta_{\text{v}}H$	(441–633)		83.8	456	A	[1987STE/MAL]
C ₁₈ H ₃₄ O ₂	[112-79-8]		<i>trans</i> 9-octadecenoic acid (elaidic acid)					
		$\Delta_{\text{fus}}H$			61.55	317.6		[1996DOM/HEA]
		$\Delta_{\text{v}}H$	(444–635)		82.3	459	A	[1987STE/MAL, 1947STU]
C ₁₈ H ₃₄ O ₂	[2549-53-3]		tetradecyl methacrylate				A	[1987STE/MAL]
		$\Delta_{\text{v}}H$	(463–611)		69.1	478		
C ₁₈ H ₃₄ O ₂	[141694-86-2]		(Z) 3-hexadecenyl acetate				GC	[1997KOU/HOS, 2000OVA/KOU]
		$\Delta_{\text{v}}H$	(373–418)		98.5	298		
C ₁₈ H ₃₄ O ₂	[128984-60-1]		(E) 3-hexadecenyl acetate				GC	[1997KOU/HOS, 2000OVA/KOU]
		$\Delta_{\text{v}}H$	(373–418)		99.1	298		
C ₁₈ H ₃₄ O ₂	[65954-24-7]		(Z) 4-hexadecenyl acetate				GC	[1997KOU/HOS, 2000OVA/KOU]
		$\Delta_{\text{v}}H$	(373–418)		97.7	298		
C ₁₈ H ₃₄ O ₂	[155055-27-9]		(E) 4-hexadecenyl acetate				GC	[1997KOU/HOS, 2000OVA/KOU]
		$\Delta_{\text{v}}H$	(373–418)		98.9	298		
C ₁₈ H ₃₄ O ₂	[34010-18-9]		(Z) 5-hexadecenyl acetate				GC	[1997KOU/HOS, 2000OVA/KOU]
		$\Delta_{\text{v}}H$	(373–418)		98.0	298		
C ₁₈ H ₃₄ O ₂	[56218-65-6]		(E) 5-hexadecenyl acetate				GC	[1997KOU/HOS, 2000OVA/KOU]
		$\Delta_{\text{v}}H$	(373–418)		98.8	298		
C ₁₈ H ₃₄ O ₂	[34010-19-0]		(Z) 6-hexadecenyl acetate					

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		T _m (K)	Method	Reference	
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)				
		$\Delta_{\text{v}}H$	(373–418)	97.8	298	GC	[1997KOU/HOS, 2000OVA/KOU]
C ₁₈ H ₃₄ O ₂	[56218-66-7]	(E) 6-hexadecenyl acetate					
		$\Delta_{\text{v}}H$	(373–418)	98.6	298	GC	[1997KOU/HOS, 2000OVA/KOU]
C ₁₈ H ₃₄ O ₂	[23192-42-9]	(Z) 7-hexadecenyl acetate					
		$\Delta_{\text{v}}H$	(373–418)	97.8	298	GC	[1997KOU/HOS, 2000OVA/KOU]
C ₁₈ H ₃₄ O ₂	[23192-83-8]	(E) 7-hexadecenyl acetate					
		$\Delta_{\text{v}}H$	(373–418)	98.5	298	GC	[1997KOU/HOS, 2000OVA/KOU]
C ₁₈ H ₃₄ O ₂	[56218-67-8]	(Z) 8-hexadecenyl acetate					
		$\Delta_{\text{v}}H$	(373–418)	97.8	298	GC	[1997KOU/HOS, 2000OVA/KOU]
C ₁₈ H ₃₄ O ₂	[56218-68-9]	(E) 8-hexadecenyl acetate					
		$\Delta_{\text{v}}H$	(373–418)	98.6	298	GC	[1997KOU/HOS, 2000OVA/KOU]
C ₁₈ H ₃₄ O ₂	[34010-20-3]	(Z) 9-hexadecenyl acetate					
		$\Delta_{\text{v}}H$	(373–418)	98.2	298	GC	[1997KOU/HOS, 2000OVA/KOU]
C ₁₈ H ₃₄ O ₂	[56218-69-0]	(E) 9-hexadecenyl acetate					
		$\Delta_{\text{v}}H$	(373–418)	98.9	298	GC	[1997KOU/HOS, 2000OVA/KOU]
C ₁₈ H ₃₄ O ₂	[56218-70-3]	(Z) 10-hexadecenyl acetate					
		$\Delta_{\text{v}}H$	(373–418)	98.5	298	GC	[1997KOU/HOS, 2000OVA/KOU]
C ₁₈ H ₃₄ O ₂	[56218-71-4]	(E) 10-hexadecenyl acetate					
		$\Delta_{\text{v}}H$	(373–418)	99.1	298	GC	[1997KOU/HOS, 2000OVA/KOU]
C ₁₈ H ₃₄ O ₂	[34010-21-4]	(Z) 11-hexadecenyl acetate					
		$\Delta_{\text{v}}H$	(373–418)	98.9	298	GC	[1997KOU/HOS, 2000OVA/KOU]
C ₁₈ H ₃₄ O ₂	[56218-72-5]	(E) 11-hexadecenyl acetate					
		$\Delta_{\text{v}}H$	(373–418)	99.5	298	GC	[1997KOU/HOS, 2000OVA/KOU]
C ₁₈ H ₃₄ O ₂	[56218-73-6]	(Z) 12-hexadecenyl acetate					
		$\Delta_{\text{v}}H$	(373–418)	99.5	298	GC	[1997KOU/HOS, 2000OVA/KOU]
C ₁₈ H ₃₄ O ₂	[64789-90-8]	(E) 12-hexadecenyl acetate					
		$\Delta_{\text{v}}H$	(373–418)	99.8	298	GC	[1997KOU/HOS, 2000OVA/KOU]
C ₁₈ H ₃₄ O ₂	[56218-74-7]	(Z) 13-hexadecenyl acetate					
		$\Delta_{\text{v}}H$	(373–418)	100	298	GC	[1997KOU/HOS, 2000OVA/KOU]
C ₁₈ H ₃₄ O ₂	[69282-67-3]	(E) 13-hexadecenyl acetate					
		$\Delta_{\text{v}}H$	(373–418)	100.3	298	GC	[1997KOU/HOS, 2000OVA/KOU]
C ₁₈ H ₃₄ O ₂	[13161-77-8]	<i>trans</i> -3-octadecenoic acid					
		$\Delta_{\text{fus}}H$		57.15	334	TA	[1982JAL/ZOG]
C ₁₈ H ₃₄ O ₂	[34450-19-6]	<i>trans</i> -4-octadecenoic acid					
		$\Delta_{\text{fus}}H$		55.88	333	TA	[1982JAL/ZOG]
C ₁₈ H ₃₄ O ₂	[7056-85-1]	<i>trans</i> -5-octadecenoic acid					
		$\Delta_{\text{fus}}H$		45.11	319	TA	[1982JAL/ZOG]
C ₁₈ H ₃₄ O ₂	[593-40-8]	<i>trans</i> -6-octadecenoic acid					
		$\Delta_{\text{fus}}H$		60.15	326	TA	[1982JAL/ZOG]
C ₁₈ H ₃₄ O ₂	[5684-82-2]	<i>trans</i> -10-octadecenoic acid					
		$\Delta_{\text{fus}}H$		58.52	326	TA	[1982JAL/ZOG]
C ₁₈ H ₃₄ O ₂	[506-17-2]	<i>cis</i> -11-octadecenoic acid (asclepic acid)					

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	Enthalpy							
		$\Delta_{\text{trs}}H$			7.8	257.8		
		$\Delta_{\text{fus}}H$			39.8	287		[1997SAT/YAN]
C ₁₈ H ₃₄ O ₂	[693-72-1]		<i>trans</i> -11-octadecenoic acid					
		$\Delta_{\text{fus}}H$			58.49	317	TA	[1982JAL/ZOG]
C ₁₈ H ₃₄ O ₂	[13126-38-0]		<i>trans</i> -12-octadecenoic acid					
		$\Delta_{\text{fus}}H$			56.71	325	TA	[1982JAL/ZOG]
C ₁₈ H ₃₄ O ₂	[693-72-1]		<i>trans</i> -13-octadecenoic acid					
		$\Delta_{\text{fus}}H$			55.62	318	TA	[1982JAL/ZOG]
C ₁₈ H ₃₄ O ₂	[13126-42-6]		<i>trans</i> -14-octadecenoic acid					
		$\Delta_{\text{fus}}H$			57.06	327	TA	[1982JAL/ZOG]
C ₁₈ H ₃₄ O ₂	[13126-44-8]		<i>trans</i> -15-octadecenoic acid					
		$\Delta_{\text{fus}}H$			58.98	331	TA	[1982JAL/ZOG]
C ₁₈ H ₃₄ O ₄	[110-33-8]		dihexyl adipate					
		Δ_vH	(470–595)		80.4	485	A	[1987STE/MAL]
C ₁₈ H ₃₄ O ₄	[109-43-3]		dibutyl decanedioate					
		Δ_vH	(468–532)		106.4	483	EB	[2008ZHU/XU]
		Δ_vH	(400–532)		87.6	465		[2008ZHU/XU]
			Note: First value based on published Antoine constants determined from the authors' experimental data; second value is what the authors reported from combining their experimental data with published literature data					
		Δ_vH			88.1	327	TGA	[1990KIS/SHO]
		Δ_vH			91.8 ± 3.2	298	TGA	[1990KIS/SHO]
		Δ_vH	(401–520)		94.3	416	A	[1987STE/MAL]
C ₁₈ H ₃₄ O ₅	[1081524-92-6]		decyl[1-(butoxycarbonyl)ethyl]carbonate					
		Δ_vH	(391–503)		79.3	406	A	[1987STE/MAL]
C ₁₈ H ₃₄ O ₆	[95-08-9]		triethylene glycol, <i>bis</i> (2-ethylbutyrate)					
		Δ_vH	(313–528)		91.7	328	A	[1987STE/MAL]
C ₁₈ H ₃₅ N	[638-65-3]		stearonitrile					
		$\Delta_{\text{fus}}H$			56.5	315.5		[2006TEI/GON]
		Δ_vH	(478–631)		78.6	493	A	[1987STE/MAL]
C ₁₈ H ₃₅ NO ₃	[2441-41-0]		N-hexadecanoylglycine					
		$\Delta_{\text{trs}}H$			4.5	384.6		
		$\Delta_{\text{trs}}H$			5.6	366.1		
		$\Delta_{\text{fus}}H$			56.5	393.1	DSC	[1986MIY/MAT]
C ₁₈ H ₃₅ NO ₃	[14379-40-9]		N-dodecanoyl-(<i>l</i>)-leucine					
		$\Delta_{\text{fus}}H$			33.5	383.1	DSC	[1986MIY/MAT]
C ₁₈ H ₃₅ NO ₃	[97850-50-5]		N-dodecanoyl-(<i>dl</i>)-leucine					
		$\Delta_{\text{trs}}H$			28.9	341.1		
		$\Delta_{\text{fus}}H$			31.0	356.6	DSC	[1986MIY/MAT]
C ₁₈ H ₃₆	[6006-34-4]		tridecylcyclopentane					
		Δ_vH	(463–634)		70.9	478		[1999DYK/SVO]
		Δ_vH			90.5	298		[1971WIL/ZWO]
C ₁₈ H ₃₆	[1795-17-1]		dodecylcyclohexane					
		$\Delta_{\text{fus}}H$			45.84	258.8		[1996DOM/HEA]
		Δ_vH			88.9 ± 0.8	298	GCC	[1978FUC/PEA]

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
	Enthalpy							
		$\Delta_{\text{v}}H$			89.5	298		[1971WIL/ZWO]
		$\Delta_{\text{v}}H$		(299–324)	93.4	311	A, ME	[1987STE/MAL, 1949PAR/MOO]
C ₁₈ H ₃₆	[296-18-4]	cyclooctadecane						
		$\Delta_{\text{trs}}H$			29.29	298.2		
		$\Delta_{\text{fus}}H$			9.87	346.2		[1969BOR/DAL]
C ₁₈ H ₃₆	[42506-48-9]	1,1-dimethylcyclohexadecane						
		$\Delta_{\text{trs}}H$			1.26	216.2		
		$\Delta_{\text{fus}}H$			0.42	221.2		
		$\Delta_{\text{fus}}H$			14.23	290.2		[1975BJO/BOR2]
C ₁₈ H ₃₆	[112-88-9]	1-octadecene						
		$\Delta_{\text{v}}H$		(399–589)	76.4	414	A	[1987STE/MAL]
		$\Delta_{\text{v}}H$			90	298		[1971WIL/ZWO]
C ₁₈ H ₃₆	[24584-00-7]	<i>cis,trans</i> -1,3,5-tri- <i>tert</i> -butylcyclohexane						
		$\Delta_{\text{fus}}H$			17.99	338.2		[1968VAN/HOE]
C ₁₈ H ₃₆	[24583-99-1]	<i>cis,cis</i> -1,3,5-tri- <i>tert</i> -butylcyclohexane						
		$\Delta_{\text{fus}}H$			26.78	393.2		[1968VAN/HOE]
C ₁₈ H ₃₆ N ₂ O ₂	[21150-82-3]	N,N'-di- <i>n</i> -hexyladipamide						
		$\Delta_{\text{fus}}H$			40.79	432		[1996DOM/HEA]
C ₁₈ H ₃₆ O	[41207-35-6]	(Z) 3-octadecen-1-ol						
		$\Delta_{\text{v}}H$		(393–433)	120.5	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₈ H ₃₆ O	[41207-36-7]	(E) 3-octadecen-1-ol						
		$\Delta_{\text{v}}H$		(393–433)	120.0	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₈ H ₃₆ O	[143-28-2]	(Z) 9-octadecen-1-ol						
		$\Delta_{\text{v}}H$		(393–433)	119.3	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₈ H ₃₆ O	[506-42-3]	(E) 9-octadecen-1-ol						
		$\Delta_{\text{v}}H$		(393–433)	120.1	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₈ H ₃₆ O	[57716-88-8]	(Z) 11-octadecen-1-ol						
		$\Delta_{\text{v}}H$		(393–433)	119.6	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₈ H ₃₆ O	[62972-93-4]	(E) 11-octadecen-1-ol						
		$\Delta_{\text{v}}H$		(393–433)	120.4	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₈ H ₃₆ O	[69820-27-5]	(Z) 13-octadecen-1-ol						
		$\Delta_{\text{v}}H$		(393–433)	120.8	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₈ H ₃₆ O	[76836-10-7]	(E) 13-octadecen-1-ol						
		$\Delta_{\text{v}}H$		(393–433)	121.2	298	CGC	[2000OVA/KOU, 1994KOU/HOS]
C ₁₈ H ₃₆ O	[502-69-2]	6,10,14-trimethyl-2-pentadecanone						
		$\Delta_{\text{v}}H$		(402–500)	56.0 ± 0.6	451		[1988BAG/GUR]
C ₁₈ H ₃₆ O	[638-66-4]	octadecanal						
		$\Delta_{\text{v}}H$		(413–616)	75.7	428	A	[1987STE/MAL, 1947STU]
C ₁₈ H ₃₆ O ₂	[629-70-9]	hexadecyl acetate						
		$\Delta_{\text{v}}H$		(373–418)	102.3	298	GC	[1997KOU/HOS, 2000OVA/KOU]
		$\Delta_{\text{v}}H$		(431–469)	70.3	446	A	[1987STE/MAL]
C ₁₈ H ₃₆ O ₂	[628-97-7]	ethyl palmitate						
		$\Delta_{\text{fus}}H$			15.09	296.4		[1996DOM/HEA]
		$\Delta_{\text{fus}}H$			53.14	296		[1967OMA]

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference	
	Enthalpy								
		$\Delta_{\text{sub}}H$		(286–294)	150.8	290	ME	[1987STE/MAL, 1967OMA]	
		Δ_vH		(429–466)	73.9	444	A	[1987STE/MAL]	
		Δ_vH		(298–318)	100.7	308	ME	[1987STE/MAL, 1967OMA]	
C ₁₈ H ₃₆ O ₂	[1731-92-6]	methyl heptadecanoate							
		$\Delta_{\text{fus}}H$			48.1	304.2	DSC	[2004CHI/ZHA]	
		Δ_vH		(467–558)	100.8 ± 1.0	298	CGC	[2004CHI/ZHA]	
		Δ_vH			89.3	350		[2002VAN/VAN]	
		Δ_vH			89.0 ± 0.7	353		[2002VAN/VAN]	
		Δ_vH			97.0 ± 1.2	298		[2002VAN/VAN]	
		Δ_vH		(421–525)	84.4	436	A, EST	[1987STE/MAL, 1963ROS/SCH]	
C ₁₈ H ₃₆ O ₂	[55-11-4]	octadecanoic acid (stearic acid)							
		$\Delta_{\text{trs}}H$			5.4	324.4			
		$\Delta_{\text{trs}}H$			5.7	325.9			
		$\Delta_{\text{fus}}H$			63.2	342.8	DSC	[2007MOR/COR]	
		$\Delta_{\text{fus}}H$			60.4	338.3		[2007MOO/KOE]	
		$\Delta_{\text{fus}}H$			57.8	344.1	DSC	[2006TEI/GON]	
		$\Delta_{\text{fus}}H$			50.93	340.2	AC	[2000YU/MEN]	
		$\Delta_{\text{fus}}H$			61.21	342.5		[1996DOM/HEA]	
		$\Delta_{\text{sub}}H$			204.1 ± 9	298	TPD	[2008CAP/LOV]	
		$\Delta_{\text{sub}}H$		(291–309)	158.5		TPTD	[2005CHA/ZIE]	
		$\Delta_{\text{sub}}H$		(296–319)	158		TPTD	[2001CHA/TOB]	
		Note: Experimental values based on the TPTD method are often inconsistent with values determined using other experimental methods							
		$\Delta_{\text{sub}}H$		(331–340)	166.5 ± 4.2	336	ME	[1961DAV/MAL, 1970COX/PIL]	
		Δ_vH		(349–415)	124.3	364	A	[1987STE/MAL]	
	Δ_vH		(457–649)	100.6	472	A	[1987STE/MAL]		
	Δ_vH		(366–389)	118.9 ± 2.0	379	ME, TE	[1982DEK/SCH]		
	Δ_vH			79.8	515	I	[1943CRA]		
C ₁₈ H ₃₆ O ₄	[56444-62-3]	2,2,11,11-tetramethyl-1,3,10,12-tetraoxacyclooctadecane							
		$\Delta_{\text{fus}}H$			35.1	373		[1975BOR]	
C ₁₈ H ₃₇ Br	[112-89-0]	1-bromooctadecane							
		Δ_vH		(430–673)	81.0	445	A, EST	[1987STE/MAL, 1961LI/ROS, 1970DYK/VAN]	
C ₁₈ H ₃₇ Cl	[3386-33-2]	1-chlorooctadecane							
		Δ_vH			108.8	298		[2006BOL/NER2]	
		Δ_vH		(333–393)	96.9	333	GC	[1980JON/MAT]	
		Δ_vH		(333–393)	93.4	353	GC	[1980JON/MAT]	
		Δ_vH		(333–393)	88.4	373	GC	[1980JON/MAT]	
		Δ_vH		(333–393)	86.7	393	GC	[1980JON/MAT]	
	Δ_vH		(472–673)	74.2	487	A	[1987STE/MAL, 1970DYK/VAN]		
C ₁₈ H ₃₇ F	[1649-73-6]	1-fluorooctadecane							
		Δ_vH		(477–633)	68.2	492	A, EST	[1987STE/MAL, 1961LI/ROS, 1970DYK/VAN]	
C ₁₈ H ₃₇ I	[629-93-6]	1-iodooctadecane							
		Δ_vH		(496–673)	109.3	298	A, EST	[1987STE/MAL, 1961LI/ROS, 1970DYK/VAN, 2006BOL/NER]	
		Δ_vH		(496–673)	77.2	511	A, EST	[1987STE/MAL, 1961LI/ROS, 1970DYK/VAN]	
C ₁₈ H ₃₇ NO	[124-26-5]	octadecanamide							
		$\Delta_{\text{trs}}H$			2.2	298.7			
		$\Delta_{\text{fus}}H$			54.8	379.7	DSC	[2008ABA/BAD]	

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		T _m (K)	Method	Reference
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)			
		$\Delta_{\text{fus}}H$	59.91	377.2		[1993ACR]
		$\Delta_{\text{sub}}H$	(367–379)	195.8 ± 4.2	373	ME [1959DAV/JON2, 1987STE/MAL]
C₁₈H₃₇NO	[41328-72-7]	N-butyl tetradecanamide				
		$\Delta_{\text{fus}}H$	45.0	336.1	DSC	[1993ACR, 1980CAR/BUS]
C₁₈H₃₇NO	[146985-21-9]	N,N-dihexyl hexanamide				
		$\Delta_{\text{v}}H$	(463–513)	80.4 ± 1.6	298	CGC [2009PAN/ANT]
C₁₈H₃₈	[593-45-3]	octadecane				
		$\Delta_{\text{fus}}H$	60.1	301.1	DSC	[2004MON/RAJ]
		$\Delta_{\text{fus}}H$	61.5	301.3		[1996DOM/HEA]
		$\Delta_{\text{sub}}H$	152.7	298	C	[1972MOR3]
		$\Delta_{\text{sub}}H$	(288–298)	153.0 ± 5	293	ME [1949BRA/SHE, 1960JON, 1970COX/PIL]
		$\Delta_{\text{v}}H$	90.6 ± 1.0	298	CGC	[2002CHI/WEB]
		$\Delta_{\text{v}}H$	91.3 ± 2.9	298	GS	[2001PUR/CHI]
		$\Delta_{\text{v}}H$	91.4 ± 1.3	298	CGC	[2000NIC/ORF]
		$\Delta_{\text{v}}H$	(363–413)	91.8	298	CGC [1995CHI/HOS]
		$\Delta_{\text{v}}H$	(423–473)	91.8	298	CGC [1995CHI/HOS]
		$\Delta_{\text{v}}H$	(453–503)	92.8	298	CGC [1995CHI/HOS]
		$\Delta_{\text{v}}H$	(413–588)	74.4	428	[1994MOR/KOB]
		$\Delta_{\text{v}}H$	91.4	298		[1994RUZ/MAJ]
		$\Delta_{\text{v}}H$	(501–548)	64.8	516	A [1987STE/MAL]
		$\Delta_{\text{v}}H$	(335–439)	80.0	348	GS [1986ALL/JOS]
		$\Delta_{\text{v}}H$	(318–361)	84.3	333	A, GS [1987STE/MAL, 1979MAC/PRA]
		$\Delta_{\text{v}}H$	72.5	343	GC	[1977NOV/NOV]
		$\Delta_{\text{v}}H$	71.8	353	GC	[1977NOV/NOV]
		$\Delta_{\text{v}}H$	71.1	363	GC	[1977NOV/NOV]
		$\Delta_{\text{v}}H$	70.5	373	GC	[1977NOV/NOV]
		$\Delta_{\text{v}}H$	69.8	383	GC	[1977NOV/NOV]
		$\Delta_{\text{v}}H$	90.8	298		[1971WIL/ZWO]
		$\Delta_{\text{v}}H$	(447–474)	78.1	460	ME [1938UBB]
		$\Delta_{\text{v}}H$	(447–590)	69.4	462	[1882KRA, 1984BOU/FRI]
C₁₈H₃₈	[1560-89-0]	2-methylheptadecane				
		$\Delta_{\text{v}}H$	(442–581)	67.8	457	A [1987STE/MAL, 1959TER/BRI]
C₁₈H₃₈	[6418-44-6]	3-methylheptadecane				
		$\Delta_{\text{v}}H$	(441–583)	65.6	456	A [1987STE/MAL, 1959TER/BRI]
C₁₈H₃₈	[26429-11-8]	4-methylheptadecane				
		$\Delta_{\text{v}}H$	(429–580)	58.9	444	A [1987STE/MAL, 1959TER/BRI]
C₁₈H₃₈	[26730-95-0]	5-methylheptadecane				
		$\Delta_{\text{v}}H$	(432–581)	61.1	447	A [1987STE/MAL, 1959TER/BRI]
C₁₈H₃₈	[61868-02-8]	2,3-dimethylhexadecane				
		$\Delta_{\text{v}}H$	(466–583)	64.9	481	A [1987STE/MAL, 1959TER/BRI]
C₁₈H₃₈	[61868-08-4]	2,4-dimethylhexadecane				
		$\Delta_{\text{v}}H$	(434–562)	69.0	449	A [1987STE/MAL, 1959TER/BRI]
C₁₈H₃₈	[101882-67-1]	2,4,6-trimethylpentadecane				
		$\Delta_{\text{v}}H$	(420–550)	64.3	435	A [1987STE/MAL, 1999DYK/SVO, 1959TER/BRI]
C₁₈H₃₈	[na]	4,9-diisopropyldodecane				
		$\Delta_{\text{v}}H$	(368–424)	70.0	383	A [1987STE/MAL]

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		T _m (K)	Method	Reference
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)			
C ₁₈ H ₃₈	[62850-21-9]	1,1,2,2-tetra- <i>tert</i> -butylethane				
	$\Delta_{\text{sub}}H$	(303–366)	71.9	341		[1984FLA/BEC]
	$\Delta_{\text{sub}}H$		74.3	298		[1984FLA/BEC]
C ₁₈ H ₃₈ O	[112-92-5]	1-octadecanol				
	$\Delta_{\text{fus}}H$		40.1	330.1	DSC	[2004VEN/CAL]
	$\Delta_{\text{trs}}H$		26.5	329.5		
	$\Delta_{\text{fus}}H$		40.1	330.3		[2002VEN/RAM]
	$\Delta_{\text{fus}}H$		66.67	331.2		[2001VAN/OON2]
	$\Delta_{\text{fus}}H$		70.08	334.2		[1991CHI/BRA]
	$\Delta_{\text{sub}}H$	(318–329)	187.4 ± 1.3	324	ME	[1965DAV/KYB, 1987STE/MAL]
	$\Delta_{\text{sub}}H$		191.2 ± 1.3	298		[1965DAV/KYB]
	Δ_vH		116.8 ± 1.2	298	CGC	[2006NIC/KWE]
	Δ_vH	(435–504)	86.4	450	A	[1987STE/MAL]
	Δ_vH	(500–573)	76.3	515	A	[1987STE/MAL]
Δ_vH	(494–575)	76.9	509	A,EB	[1987STE/MAL, 1970AMB/SPR]	
Δ_vH	(334–356)	113.5	345	A, ME	[1987STE/MAL, 1965DAV/KYB]	
C ₁₈ H ₃₈ O ₂	[2136-71-2]	2-(hexadecyloxy)ethanol				
	$\Delta_{\text{trs}}H$		14.94	311.7		
	$\Delta_{\text{fus}}H$		37.32	318.5	DTA	[1979KUC/SKU]
C ₁₈ H ₃₈ O ₂	[3155-43-9]	1,18-octadecanediol				
	$\Delta_{\text{trs}}H$		38.7	366.1		
	$\Delta_{\text{fus}}H$		33.6	371.5	DSC	[1999OGA/NAK]
C ₁₈ H ₃₈ O ₄	[4161-35-5]	6,6'-[1,6-hexanediy]bis(oxy)]bis-1-hexanol				
	$\Delta_{\text{fus}}H$		57.96	329.1	DSC	[1991BED/BOO]
C ₁₈ H ₃₈ O ₄	[3055-94-5]	2-[2-(2-[dodecyloxy]ethoxy)ethoxy]ethanol				
	Δ_vH	(475–523)	102.7	490	A	[1987STE/MAL]
C ₁₈ H ₃₈ O ₄ S ₂	[na]	2-deoxy-(<i>D</i>)-glucose dihexyl dithioacetal				
	$\Delta_{\text{trs}}H$		16.4	376.2		
	$\Delta_{\text{fus}}H$		45.0	386.4	DSC	[1989VAN/VAN]
C ₁₈ H ₃₈ O ₄ S ₂	[na]	(<i>L</i>)-rhamnose dihexyl dithioacetal				
	$\Delta_{\text{fus}}H$		48.5	388.2	DSC	[1989VAN/VAN]
C ₁₈ H ₃₈ O ₅ S ₂	[115395-54-5]	(<i>D</i>)-glucose dihexyl dithioacetal				
	$\Delta_{\text{trs}}H$		5.9	373.6		
	$\Delta_{\text{fus}}H$		44.6	377	DSC	[1989VAN/VAN]
C ₁₈ H ₃₈ O ₅ S ₂	[na]	(<i>L</i>)-arabinose dihexyl dithioacetal				
	$\Delta_{\text{trs}}H$		6.7	345.1		
	$\Delta_{\text{trs}}H$		1.0	358.2		
	$\Delta_{\text{fus}}H$		39.2	367.2	DSC	[1989VAN/VAN]
C ₁₈ H ₃₈ O ₉	[25990-94-7]	1, ω -dimethoxyocta(oxyethylene)				
	$\Delta_{\text{fus}}H$		60.1	276.2		[1997SCH/VER]
C ₁₈ H ₃₈ S	[2885-00-9]	1-octadecanethiol				
	Δ_vH	(492–670)	77.1	507	EST	[1999DYK/SVO]
C ₁₈ H ₃₈ S ₂	[4485-77-2]	dioctyl disulfide				
	Δ_vH	(490–650)	78.3	514		[1999DYK/SVO]
C ₁₈ H ₃₉ N	[112-69-6]	N,N-dimethylhexadecylamine				
	Δ_vH	(483–671)	67.3	498	A	[1987STE/MAL]

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₈ H ₃₉ N	[2044-21-5] $\Delta_v H$	dinonylamine (486–676)	67.7	501	A	[1987STE/MAL]
C ₁₈ H ₃₉ N	[5877-76-9] $\Delta_v H$	N-ethylhexadecylamine (406–613)	66.4	421	A	[1987STE/MAL, 1947STU]
C ₁₈ H ₃₉ N	[124-30-1] $\Delta_v H$	octadecylamine (450–635)	76.2	465	A	[1987STE/MAL]
C ₁₈ H ₃₉ O ₄ P	[2528-39-4] $\Delta_v H$ $\Delta_v H$	trihexyl phosphate (483–513) (493–523)	104.8 104.9	298 298	CGC	[2007PAN/ANT2] [2007PAN/ANT2]
C ₁₉ H ₁₀ O	[3074-00-8] $\Delta_{\text{fus}} H$	6 <i>H</i> -benzo[<i>cd</i>]pyren-6-one 13.1	524.2	DSC	[2010KES/AUC]	
C ₁₉ H ₁₃ F ₃ O	[145698-49-3] $\Delta_{\text{fus}} H$	4-ethoxy-4'-trifluoromethyldiphenyl diacetylene 32.73	424.9	DSC	[1993JUA/CHE]	
C ₁₉ H ₁₃ NO	[846-63-9] $\Delta_v H$	2-(1-naphthyl)-5-phenyloxazole (510–595)	89.2	525	A	[1987STE/MAL]
C ₁₉ H ₁₃ NO	[5472-23-1] $\Delta_{\text{fus}} H$ $\Delta_{\text{sub}} H$	10-phenylacridin-9(10 <i>H</i>)-one 38.9 128	550	DSC DSC	[2003STO/KRZ] [2003STO/KRZ]	
C ₁₉ H ₁₄	[3351-31-3] $\Delta_{\text{fus}} H$	3-methylchrysene 16.5	445.0	DSC	[2010KES/AUC]	
C ₁₉ H ₁₄	[3351-30-2] $\Delta_{\text{fus}} H$	4-methylchrysene 18.3	424.0	DSC	[2010KES/AUC]	
C ₁₉ H ₁₄	[3697-24-3] $\Delta_{\text{fus}} H$	5-methylchrysene 19.0	390.7	DSC	[2010KES/AUC]	
C ₁₉ H ₁₄	[1705-85-7] $\Delta_{\text{fus}} H$	6-methylchrysene 22.7	432.5	DSC	[2010KES/AUC]	
C ₁₉ H ₁₄ F ₂	[145698-35-7] $\Delta_{\text{fus}} H$	4-propyl-3',4'-difluorodiphenyl diacetylene 22.03	343.7	DSC	[1993JUA/CHE]	
C ₁₉ H ₁₄ O ₄	[na] $\Delta_{\text{fus}} H$	4-hydroxyphenyl-4''-hydroxybiphenyl-4'-carboxylate 49.59	566.2		[2000PUN]	
C ₁₉ H ₁₅ Cl	[76-83-5] $\Delta_{\text{fus}} H$	triphenylchloromethane 27.9	376.8		[1996DOM/HEA]	
C ₁₉ H ₁₅ F ₉ OS	[246543-99-7] $\Delta_{\text{fus}} H$	4-[(3,3,4,4,5,5,6,6,6-nonafluorohexyl)thio]methoxy-1,1'-biphenyl 43.1	333.3	DTA	[1999DEG/GUI]	
C ₁₉ H ₁₅ F ₉ S	[246543-96-4] $\Delta_{\text{fus}} H$	4-[(3,3,4,4,5,5,6,6,6-nonafluorohexyl)thio]methyl-1,1'-biphenyl 41.0	307.1	DTA	[1999DEG/GUI]	
C ₁₉ H ₁₅ N	[574-45-8] $\Delta_{\text{fus}} H$ $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$	N-phenyl benzophenone imine 29.14 (348–387) 115.5 ± 1.8 119.7 ± 1.8	392.3 367 298	T	[1997VER/MOR] [1997VER/MOR] [1997VER/MOR]	
C ₁₉ H ₁₅ N ₃	[14309-25-2] $\Delta_{\text{sub}} H$	triphenylazidomethane (335–363)	120.6	349	A	[1987STE/MAL, 1974PEP/ERL]

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₉ H ₁₆	[519-73-3]	triphenylmethane				
	$\Delta_{\text{fus}}H$		20.7	367.2	DSC	[1999VER3]
	$\Delta_{\text{fus}}H$		21.97	365.3		[1996DOM/HEA]
	$\Delta_{\text{sub}}H$	(323–353)	109.1 ± 0.6	298	GS	[1999VER3]
	$\Delta_{\text{sub}}H$	(323–353)	106.7 ± 0.6	338	GS	[1999VER3]
	$\Delta_{\text{sub}}H$		112	298	CGC-DSC	[1998CHI/HES]
	$\Delta_{\text{sub}}H$	(343–363)	113.9	353	EM	[1989SAS/NGU]
	$\Delta_{\text{sub}}H$	(303–358)	106.8	330	T	[1986HAN/ECK]
	$\Delta_{\text{sub}}H$	(325–349)	100 ± 0.4	339	V	[1959AIH, 1970COX/PIL, 1987STE/MAL]
	$\Delta_{\text{sub}}H$		100.7	298		[1986MAR/LOE, 1936CUT/BEN]
	$\Delta_{\text{sub}}H$		105 ± 0.8			[1974PEP/ERL]
	Δ_vH		93.2 ± 2.2	298	CGC	[2008HAN/NUT]
	Δ_vH		94.6	298	CGC	[1998CHI/HES]
	Δ_vH	(453–503)	95.0	298	CGC	[1995CHI/HOS]
Δ_vH	(343–462)	82.0	403		[1989SAS/NGU]	
Δ_vH	(512–643)	58.6	527	A	[1987STE/MAL]	
C ₁₉ H ₁₆ ClNO ₄	[53-86-1]	1-(4-chlorobenzoyl)-5-methoxy-2-methyl-1 <i>H</i> -indole-acetic acid (indomethacin)				
	$\Delta_{\text{fus}}H$		36.5	435.2	DSC	[2010MUR/PIK]
	$\Delta_{\text{fus}}H$		43.5	434.5	DSC	[2009ACE/NIC]
	$\Delta_{\text{fus}}H$		39.99	434	DSC	[2008BAS/BOS]
	$\Delta_{\text{fus}}H$		37.9	433	DSC	[2006WAS/HOL]
	$\Delta_{\text{fus}}H$ (I)		36.85	432.3		
	$\Delta_{\text{fus}}H$ (II)		32.94	426.2		[2004HAM/FEU]
	$\Delta_{\text{fus}}H$		37.8	433		[2004LEG/FEU, 2007BER/WAS]
$\Delta_{\text{fus}}H$		36.49	435.2		[2000HAN/PAR]	
C ₁₉ H ₁₆ F ₈ N ₄ O ₂	[91488-85-6]	N-ethyl-4-[(4-nitrophenyl)azo]-N-(2,2,3,3,4,4,5,5-octafluoropentyl)benzenamine	112.6			[1984KAR/ROD]
C ₁₉ H ₁₆ F ₈ O ₂	[464213-30-7]	2,3-bis(trifluoromethyl)-4-methoxyphenyl- α , α -difluoro-4-n-propylbenzyl ether				
	$\Delta_{\text{fus}}H$		21.3	301.4	DSC	[2002MIY/KAT]
C ₁₉ H ₁₆ O	[76-84-6]	triphenylmethanol				
	$\Delta_{\text{fus}}H$		27.24	441.1	DSC	[1998VER3]
	$\Delta_{\text{sub}}H$		121.8 ± 1.7	298	GS	[1998VER3]
$\Delta_{\text{sub}}H$	(353–373)	122	363	A	[1987STE/MAL]	
C ₁₉ H ₁₆ O ₂	[160731-89-5]	2-fluorenyl-2-methyl-1,3-cyclopentandione				
	$\Delta_{\text{fus}}H$		24.6	395.2	DSC	[1995NOL/VER]
$\Delta_{\text{sub}}H$	(353–388)	122.3 ± 1.6	371	T	[1995NOL/VER]	
C ₁₉ H ₁₆ O ₅	[111171-32-5]	8-(hydroxymethyl)-6-phenyl-2-oxo-2 <i>H</i> -1-benzopyran-3-carboxylic acid, ethyl ester				
	$\Delta_{\text{fus}}H$		43.08	474.8	DSC	[1992HUA/ZHO2]
C ₁₉ H ₁₇ ClN ₂ O	[2955-38-6]	1-(cyclopropylmethyl)-5-phenyl-7-chloro-1,3-dihydro-2 <i>H</i> -1,4-benzodiazepin-2-one (prazepam)				
	$\Delta_{\text{fus}}H$		27.69	419	DSC	[2001VER/AUG]
C ₁₉ H ₁₇ F ₃ O	[126315-23-9]	4-pentoxy-2',3',4'-trifluorodiphenylacetylene				
	$\Delta_{\text{fus}}H$		33.1	315.8	DSC	[1995HSU/TSA]
C ₁₉ H ₁₇ NO ₂	[4946-83-2]	1-piperidinoanthraquinone				
	$\Delta_{\text{sub}}H$	(383–392)	U 18.3	387.5	A	[1987STE/MAL, 1977EIB/TRO]
Δ_vH	(395–404)	82.0	399	A	[1987STE/MAL]	
C ₁₉ H ₁₇ N ₃ O ₂	[na]	Disperse Yellow 50				

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
	$\Delta_{\text{sub}}H$	(464–484)	69.0	474	GS	[1989NIS/AND]
C ₁₉ H ₁₈ FNO ₃	[94611-29-7]	4-cyano-2-fluorophenyl 4-pentoxybenzoate				
	$\Delta_{\text{fus}}H$		36.2	359.7	DSC	[1984KEL]
C ₁₉ H ₁₈ FNO ₃	[94610-84-1]	4-cyano-3-fluorophenyl 4-pentoxybenzoate				
	$\Delta_{\text{fus}}H$		37.7	341.2	DSC	[1984KEL]
C ₁₉ H ₁₈ F ₂	[109970-66-3]	4-pentyl-3',4'-difluorodiphenylacetylene				
	$\Delta_{\text{fus}}H$		22.1	323.1	DSC	[1995HSU/TSA]
C ₁₉ H ₁₈ N ₂ O ₃	[na]	1',3'-dihydro-1',3',3'-trimethyl-6-nitrospiro[2 <i>H</i> -1-benzopyran-2,2'-(2 <i>H</i>)-indole]				
	$\Delta_{\text{fus}}H$		34.0	453.1		[2004KUL/MAR]
C ₁₉ H ₁₈ O ₂	[160731-87-3]	2-diphenylmethyl-2-methyl-1,3-cyclopentandione				
	$\Delta_{\text{fus}}H$		34.3	394.2		[1995NOL/VER]
	$\Delta_{\text{sub}}H$	(355–393)	120.2 ± 1.1	374	T	[1995NOL/VER]
C ₁₉ H ₁₈ O ₃	[568-72-9]	6,7,8,9-tetrahydro-1,6,6-trimethylphenanthro[1,2- <i>b</i>]furan-10,11-dione				
	$\Delta_{\text{fus}}H$		30.46	486.9		
	$\Delta_{\text{fus}}H$		29.17	486		[1992HUA/ZHO, 1988HUA/TAN]
C ₁₉ H ₁₈ O ₄	[17397-93-2]	6,7,8,9-tetrahydro-6-(hydroxymethyl)1,6-dimethylphenanthro-[1,2- <i>b</i>]-furan-10,11-dione				
	$\Delta_{\text{fus}}H$		23.74	479.7		[1992HUA/ZHO]
C ₁₉ H ₁₉ F	[109970-64-1]	4-pentyl-4'-fluorodiphenylacetylene				
	$\Delta_{\text{fus}}H$		25.6	337.4	DSC	[1995HSU/TSA]
C ₁₉ H ₁₉ FO	[139195-67-8]	4-pentoxy-4'-fluorodiphenylacetylene				
	$\Delta_{\text{fus}}H$		27.2	330.9	DSC	[1995HSU/TSA]
C ₁₉ H ₁₉ NO ₂	[483362-76-1]	1-[(4-nitrophenyl)ethynyl]-4-pentylbenzene				
	$\Delta_{\text{fus}}H$		21.46	342.7	DSC	[2002SPA/DZI]
C ₁₉ H ₁₉ NO ₃	[483362-80-7]	1-[(4-nitrophenyl)ethynyl]-4-pentylbenzene				
	$\Delta_{\text{fus}}H$		33.56	359.9	DSC	[2002SPA/DZI]
C ₁₉ H ₁₉ NO ₄ S	[na]	ethyl 5-phenylsulfonyl-4,7-dihydro-4,7-ethano-2 <i>H</i> -isoindole-1-carboxylate				
	$\Delta_{\text{fus}}H$		29.8	453.3		[2000UNO/ITO]
C ₁₉ H ₂₀ F ₃ N ₃ O ₃	[na]	2-[3-(trifluoromethyl)-phenyl]amino-3-pyridinecarboxylic acid β -morpholino-ethyl ester				
	$\Delta_{\text{fus}}H$		34.5	350		[1996DOM/HEA]
C ₁₉ H ₂₀ O ₂	[137932-36-6]	3-diphenylmethyl-3-methyl-2,4-pentanedione				
	$\Delta_{\text{fus}}H$		25.1	352.2		[1995NOL/VER]
	$\Delta_{\text{sub}}H$		114.4 ± 0.6	298	T,B	[1995NOL/VER]
	$\Delta_{\text{v}}H$	(353–386)	83.1 ± 0.5	370	GS	[1995NOL/VER]
C ₁₉ H ₂₀ O ₃	[35825-57-11]	1,2,6,7,8,9-hexahydro-1,6,6-trimethylphenanthro[1,2- <i>b</i>]furan-10,11-dione				
	$\Delta_{\text{fus}}H$		26.46	464.6	DSC	[1988HUA/TAN]
C ₁₉ H ₂₀ O ₃	[na]	1-(diphenylmethyl)-4-methyl-2,6,7-trioxabicyclo[2.2.2]octane				
	$\Delta_{\text{fus}}H$		32.6	443.2		[1995RAK/VER2]
C ₁₉ H ₂₀ O ₄	[85-68-7]	butylbenzyl phthalate				
	$\Delta_{\text{v}}H$	(416–516)	89.0	431	A	[1987STE/MAL]
C ₁₉ H ₂₀ O ₄	[74254-53-8]	dibenzyl ethylmalonate				
	$\Delta_{\text{v}}H$	(403–483)	94.1	418	A	[1987STE/MAL]

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₉ H ₂₁ F ₁₉	[139277-01-3]	1,1,1,2,3,3,4,4,5,5,6,6,7,7,8,8-hexadecafluoro-2-(trifluoromethyl)octadecane				
	$\Delta_{\text{trs}}H$		1.0	274		
	$\Delta_{\text{fus}}H$		25.0	298	DSC	[1992HOP/MOL]
C ₁₉ H ₂₁ IO ₃ S	[313057-07-7]	4-(7-octenyloxy)phenyl 5-iodo-2-thiophene carboxylate				
	$\Delta_{\text{fus}}H$		80.33	324.9	DSC	[2000WU/WAN]
C ₁₉ H ₂₁ NO	[127529-16-2]	(±)1,2-diphenyl-2-N-piperidinyl-1-ethanone				
	$\Delta_{\text{fus}}H$		33.93	349.2		[1994WEL/VER]
	$\Delta_{\text{sub}}H$		147.1 ± 1		B	[1994WEL/VER]
C ₁₉ H ₂₁ NO ₃	[483362-65-8]	2-(4-nitrophenyl)-1-(4-pentylphenyl)ethanone				
	$\Delta_{\text{fus}}H$		29.46	363.2	DSC	[2002SPA/DZI]
C ₁₉ H ₂₁ NO ₄	[483362-69-2]	2-(4-nitrophenyl)-1-(4-pentyloxyphenyl)ethanone				
	$\Delta_{\text{fus}}H$		27.07	353.4	DSC	[2002SPA/DZI]
C ₁₉ H ₂₂ FN ₃ O	[1649-18-9]	1-(4-fluorophenyl)-4-[4-(2-pyridinyl)-1-piperazinyl]-1-butanone (azaperone)				
	$\Delta_{\text{fus}}H$		30.5	366.2	DSC	[1981DRA/AZI]
C ₁₉ H ₂₂ F ₈ O ₂	[na]	2,3-bis(trifluoromethyl)-4-methoxyphenyl- <i>trans</i> -4-n-propyl-cyclohexyl- α,α -difluoromethyl ether				
	$\Delta_{\text{fus}}H$		21.4	314.7		[2002MIY/KAT]
C ₁₉ H ₂₃ NO	[na]	p-hexyloxybenzylideneaniline				
	$\Delta_{\text{trs}}H$		0.19	73.41		
	$\Delta_{\text{fus}}H$		30.91	321.6		[1996DOM/HEA]
C ₁₉ H ₂₃ NO	[29743-08-6]	4-butyl-N-[(4-ethoxyphenyl)methylene]benzenamine				
	$\Delta_{\text{sub}}H$			NA		[1981PIR/AZA]
C ₁₉ H ₂₃ N ₃	[33089-61-1]	N-methyl-N'-2,4-xylyl-N-(N-2,4-xylylformimodol)formamidine (amitraz)				
	$\Delta_{\text{fus}}H$ (I)		26.77	355.3		
	$\Delta_{\text{fus}}H$ (II)		19.47	344.4		
	$\Delta_{\text{fus}}H$ (III)		53.14	388.6		[2004DEV/VAN]
C ₁₉ H ₂₄	[25566-92-1]	dicumenylmethane				
	Δ_vH	(303–402)	71	318		[1999DYK/SVO]
	Δ_vH	(608–704)	57.9	623		[1999DYK/SVO]
	Δ_vH	(323–402)	73.7	338	A	[1987STE/MAL]
C ₁₉ H ₂₄ N ₂ O ₂	[55268-74-1]	(+) 2-(cyclohexylcarbonyl)-1,2,3,6,7,11b-hexahydro-4H-pyrazino-[2,1-a]isoquinolin-4-one ((+) praziquantel)				
	$\Delta_{\text{fus}}H$		30.8	412.2		[1998ELA/GIR]
C ₁₉ H ₂₄ N ₂ O ₂	[na]	(+) 2-(cyclohexylcarbonyl)-1,2,3,6,7,11b-hexahydro-4H-pyrazino-[2,1-a]isoquinolin-4-one ((+) praziquantel)				
	$\Delta_{\text{fus}}H$		23.9	386		[1998ELA/GIR]
C ₁₉ H ₂₄ N ₂ O ₂	[na]	(–) 2-(cyclohexylcarbonyl)-1,2,3,6,7,11b-hexahydro-4H-pyrazino[2,1-a]-isoquinolin-4-one ((–) praziquantel)				
	$\Delta_{\text{fus}}H$		24.15	385.5		[1998ELA/GIR]
C ₁₉ H ₂₄ O	[1706-65-6]	2- <i>tert</i> -butyl-4-methyl-6- α -methylbenzylphenol				
	$\Delta_{\text{fus}}H$		31.38	337.7	DTA	[1972INO/LIA]
C ₁₉ H ₂₄ O ₃	[104225-35-6]	3-[(2,3-dihydro-1 <i>H</i> -inden-5-yl)carbonyl]-1,2,2-trimethylcyclopentanecarboxylic acid				
	$\Delta_{\text{fus}}H$		22.5	404.3	DSC	[1992TER/PAU]
C ₁₉ H ₂₆ O ₄	[104225-25-4]	3-(4-methoxy-2,6-dimethylbenzoyl)-1,2,2-trimethylcyclopentanecarboxylic acid				
	$\Delta_{\text{fus}}H$		28.31	416.7	DSC	[1992TER/PAU]
C ₁₉ H ₂₆ O ₆	[104225-20-9]	1,2,2-trimethyl-3-(2,4,6-trimethoxybenzoyl)cyclopentanecarboxylic acid				
	$\Delta_{\text{fus}}H$		29.68	432.2	DSC	[1992TER/PAU]

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	Enthalpy							
C ₁₉ H ₂₇ NO ₃	[172589-18-3]	3-[(3,4-dimethylphenyl)(hydroxyimino)methyl]-1,2,2-trimethylcyclopentanecarboxylic acid methyl ester		39.14	426			[1995NUR/LEL]
	$\Delta_{\text{fus}}H$							
C ₁₉ H ₂₇ NO ₄	[172589-21-8]	3-[(hydroxyimino)(4-ethoxyphenyl)methyl]-1,2,2-trimethylcyclopentanecarboxylic acid methyl ester		36.75	401			[1995NUR/LEL]
	$\Delta_{\text{fus}}H$							
C ₁₉ H ₂₇ NO ₄	[252013-92-6]	(R)- β -cyano-3,4-dimethoxy- α , α -dimethyl- β -(1-methylethyl)benzenepropanoic acid, ethyl ester		36.9	386.4		DSC	[2003ROU/JIM2]
	$\Delta_{\text{fus}}H$			36.82	386.2			[1999ROS/MOL]
	$\Delta_{\text{fus}}H$							
C ₁₉ H ₂₇ NO ₅	[172589-23-0]	3-[(hydroxyimino)(3,4-dimethoxyphenyl)methyl]-1,2,2-trimethylcyclopentanecarboxylic acid methyl ester		36.2	393			[1995NUR/LEL]
	$\Delta_{\text{fus}}H$							
C ₁₉ H ₂₇ N ₃ O ₈	[53848-87-6]	dodecyl 2,4,6-trinitrobenzoate		7.64	325			
	$\Delta_{\text{trs}}H$			29.55	394		DSC	[1974WAR/WIL]
	$\Delta_{\text{fus}}H$							
C ₁₉ H ₂₈ N ₂	[na]	4-(4- <i>n</i> -heptyl-1-piperidinyl)benzotrile		29.01	326.2			[1991SHE/WEI]
	$\Delta_{\text{fus}}H$							
C ₁₉ H ₂₈ O ₂	[58-22-0]	testosterone		28.2	426.5		DSC	[2006WAS/HOL]
	$\Delta_{\text{fus}}H$			29.45	428			[1994REG/CHM]
	$\Delta_{\text{fus}}H$							
C ₁₉ H ₂₉ BrN ₂ O ₄	[138517-08-5]	(4-nitrophenyl)-12-bromododecyl carbamate		48.94	373.5		DSC	[1993TIE/FRA]
	$\Delta_{\text{fus}}H$							
C ₁₉ H ₃₀	[55030-46-1]	7-phenyl-6-tridecene (391–449)		77.2	406		A, MG	[1987STE/MAL, 1955SCH/WHI]
	Δ_vH							
C ₁₉ H ₃₀ O ₂	[na]	5 α -androstane-3-one-17 β -ol		27.15	455.5			[1996DOM/HEA]
	$\Delta_{\text{fus}}H$							
C ₁₉ H ₃₁ N ₃ O ₃	[138517-13-2]	1-dodecyl-3-(4-nitrophenyl) urea		40.88	390.8		DSC	[1993TIE/FRA]
	$\Delta_{\text{fus}}H$							
C ₁₉ H ₃₂	[123-02-4]	tridecylbenzene		72.0	488			[1999DYK/SVO]
	Δ_vH		(473–651)	90.0	358			[1990POM/PLA]
	Δ_vH		(343–463)	94.6	298			[1971WIL/ZWO]
	Δ_vH							
C ₁₉ H ₃₂	[2400-01-3]	7-phenyltridecane (413–470)		76.2	428		A, MG	[1987STE/MAL, 1955SCH/WHI]
	Δ_vH							
C ₁₉ H ₃₂ O ₂	[301-00-8]	methyl linolenate		110.5 ± 0.5	298		CGC	[2007LIP/KAP]
	Δ_vH		(423–503)	102.1	298		GC	[1997KRO/VEL]
	Δ_vH		(394–459)	87.7	409		A, MG, OM	[1987STE/MAL, 1952SCO/MAC]
	Δ_vH							
C ₁₉ H ₃₄	[1610-24-8]	tricyclohexylmethane		117.4	311			[1987STE/MAL, 1964MOR]
	$\Delta_{\text{sub}}H$		(301–321)	81.4	348		A	[1987STE/MAL, 1964MOR]
	Δ_vH		(333–365)	73.3	443		A	[1987STE/MAL]
	Δ_vH		(428–605)					
C ₁₉ H ₃₄ O ₂	[112-62-9]	linoleic acid, methyl ester (methyl linoleate)		107.8 ± 0.6	298		CGC	[2007LIP/KAP]
	Δ_vH		(423–503)	102.2	298		GC	[1997KRO/VEL]
	Δ_vH							

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		T _m (K)	Method	Reference	
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)				
		$\Delta_{\text{v}}H$	(453–543)	77.2	498	GC	[1993HUS/SAR]
		$\Delta_{\text{v}}H$	(391–459)	86.3	406	A, MG, OM	[1987STE/MAL, 1952SCO/MAC]
C ₁₉ H ₃₆	[2090-15-5]	1,1-dicyclohexylheptane					
		$\Delta_{\text{v}}H$	(293–368)	87.8	330	A	[1987STE/MAL, 1999DYK/SVO]
		$\Delta_{\text{v}}H$	(422–458)	73.8	437	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₁₉ H ₃₆	[26186-01-6]	1-nonadecyne					
		$\Delta_{\text{v}}H$	(462–637)	66.9	477		[1999DYK/SVO]
C ₁₉ H ₃₆	[61847-98-1]	2-nonadecyne					
		$\Delta_{\text{v}}H$	(469–648)	67.8	484		[1999DYK/SVO]
C ₁₉ H ₃₆	[61886-65-5]	3-nonadecyne					
		$\Delta_{\text{v}}H$	(460–635)	66.5	475		[1999DYK/SVO]
C ₁₉ H ₃₆ O	[6907-38-6]	cyclononadecanone					
		$\Delta_{\text{sub}}H$		82.4			[1938WOL/WEG, 1960JON]
C ₁₉ H ₃₆ O ₂	[112-62-9]	methyl cis-9-octadecenoate (methyl oleate)					
		$\Delta_{\text{v}}H$		106.2 ± 0.7	298	CGC	[2007LIP/KAP]
		$\Delta_{\text{v}}H$	(423–503)	103.3	298	GC	[1997KRO/VEL]
		$\Delta_{\text{v}}H$	(433–473)	99.6	298	CGC	[1995CHI/HOS]
		$\Delta_{\text{v}}H$	(453–543)	77.2	498	GC	[1993HUS/SAR]
		$\Delta_{\text{v}}H$		106.8 ± 1.0	298	GCC	[1980FUC/PEA]
		$\Delta_{\text{v}}H$	(428–486)	83.0	443	A	[1987STE/MAL, 1964ROS/SCH]
		$\Delta_{\text{v}}H$	(401–458)	86.7	416	MG, OM	[1952SCO/MAC]
C ₁₉ H ₃₆ O ₂	[1937-62-8]	methyl elaidate					
		$\Delta_{\text{v}}H$	(453–543)	77.2	498	GC	[1993HUS/SAR]
C ₁₉ H ₃₆ O ₃	[141-24-2]	methyl ricinoleate					
		$\Delta_{\text{v}}H$	(453–543)	89.3	498	GC	[1993HUS/SAR]
C ₁₉ H ₃₆ O ₅	[na]	undecyl[1-(butoxycarbonyl)ethyl]carbonate					
		$\Delta_{\text{v}}H$	(438–637)	77.0	453	A	[1987STE/MAL]
C ₁₉ H ₃₇ NO	[112-96-9]	octadecyl isocyanate					
		$\Delta_{\text{v}}H$	(388–494)	77.8	403	A	[1987STE/MAL]
C ₁₉ H ₃₇ NO ₃	[na]	2-[2-ethyl-(hexanoyloxy)]-N,N-dibutylpropionamide					
		$\Delta_{\text{v}}H$	(403–448)	83.0	418	A	[1987STE/MAL]
C ₁₉ H ₃₇ NO ₃	[56255-31-3]	N-hexadecanoyl-(<i>l</i>)-alanine					
		$\Delta_{\text{fus}}H$		65.3	374.1	DSC	[1986MIY/MAT]
C ₁₉ H ₃₇ NO ₃	[14379-30-7]	N-tetradecanoyl-(<i>l</i>)-valine					
		$\Delta_{\text{trs}}H$		14.9	44.53		
		$\Delta_{\text{fus}}H$		20.6	365.1	DSC	[1986MIY/MAT]
C ₁₉ H ₃₇ NO ₃	[83871-19-6]	N-tetradecanoyl-(<i>dl</i>)-valine					
		$\Delta_{\text{fus}}H$		68.1	370.1	DSC	[1986MIY/MAT]
C ₁₉ H ₃₈	[6006-33-3]	tridecylcyclohexane					
		$\Delta_{\text{v}}H$	(474–651)	72.2	489		[1999DYK/SVO]
		$\Delta_{\text{v}}H$		94.5	298		[1971WIL/ZWO]
C ₁₉ H ₃₈	[13151-92-3]	7-cyclohexyltridecane					
		$\Delta_{\text{v}}H$	(391–449)	75.6	406	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₁₉ H ₃₈	[1795-22-8]	tetradecylcyclopentane					
		$\Delta_{\text{v}}H$	(475–648)	73.6	490		[1999DYK/SVO]

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
	Enthalpy							
		$\Delta_{\text{v}}H$			95.4	298		[1971WIL/ZWO]
C ₁₉ H ₃₈	[55044-77-4]		7-(cyclopentylmethyl)tridecane	(389–446)	76.5	404	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₁₉ H ₃₈	[18435-45-5]		1-nonadecene	(560–604)	63.3	575	A	[1987STE/MAL]
		$\Delta_{\text{v}}H$			95	298		[1971WIL/ZWO]
C ₁₉ H ₃₈ O	[629-66-3]		2-nonadecanone		68.65	328	DSC	[1993VIL/HAM]
C ₁₉ H ₃₈ O	[504-57-4]		10-nonadecanone		66.67	330	DSC	[1993RUE/SAR, 1993VIL/HAM]
C ₁₉ H ₃₈ O ₂	[112-61-8]		methyl stearate		61.7	310.9	DSC	[2004CHI/ZHA]
		$\Delta_{\text{fus}}H$			71.1	311		[2003NIK/MAR]
		$\Delta_{\text{fus}}H$			64.4	310		[1936KIN/GAR]
		$\Delta_{\text{sub}}H$	(299–310)	158.2 ± 2.5	304			[1965DAV/KYB, 1987STE/MAL]
		$\Delta_{\text{v}}H$	(467–558)	109.5 ± 2.7	298		CGC	[2004CHI/ZHA]
		$\Delta_{\text{v}}H$		98	350			[2002VAN/VAN]
		$\Delta_{\text{v}}H$		90.0 ± 0.3	401			[2002VAN/VAN]
		$\Delta_{\text{v}}H$		105.9 ± 1.4	298			[2002VAN/VAN]
		$\Delta_{\text{v}}H$	(463–523)	106.2	298		GC	[1997KRO/VEL]
		$\Delta_{\text{v}}H$	(453–543)	75.4	498		GC	[1993HUS/SAR]
		$\Delta_{\text{v}}H$	(427–484)	83.2	442		A	[1987STE/MAL, 1964ROS/SCH]
C ₁₉ H ₃₈ O ₂	[14010-23-2]		ethyl margarate (ethyl heptadecanoate)		16.57	291.2		
		$\Delta_{\text{trs}}H$			36.2	298.4		[1936KIN/GAR]
C ₁₉ H ₃₈ O ₂	[2239-78-3]		propyl palmitate	(439–477)	74.5	454	A	[1987STE/MAL, 1948BON/ATH, 1984BOU/FRI]
C ₁₉ H ₃₈ O ₂	[142-91-6]		isopropyl palmitate	(433–471)	73.6	448	A	[1987STE/MAL, 1948BON/ATH, 1984BOU/FRI]
C ₁₉ H ₃₈ O ₂	[646-30-0]		nonadecanoic acid		7.4	339		
		$\Delta_{\text{fus}}H$			57.0	340.4	DSC	[2007GBA/NEG]
		$\Delta_{\text{trs}}H$			9.76	338		
		$\Delta_{\text{fus}}H$			57.62	341.2		[1996DOM/HEA]
		$\Delta_{\text{sub}}H$	(298–315)	143.6			TPTD	[2005CHA/ZIE]
		$\Delta_{\text{sub}}H$		198.7 ± 5				[1968BAC/NOV, 1970COX/PIL]
		$\Delta_{\text{v}}H$	(511–659)	94.4	526		A	[1987STE/MAL]
		$\Delta_{\text{v}}H$	(371–394)	121.8	386		ME, TE	[1982DEK/SCH]
C ₁₉ H ₃₈ O ₃	[35274-05-6]		hexadecyl lactate	(405–556)	90.5	420	A	[1987STE/MAL]
C ₁₉ H ₃₈ O ₃	[94434-74-9]		3-octyloxypropionic acid, octyl ester	(443–513)	73.6	458	A	[1987STE/MAL]
C ₁₉ H ₃₉ Br	[4434-66-6]		1-bromononadecane					

Note: Experimental values based on the TPTD method are often inconsistent with values determined using other experimental methods

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
	Enthalpy							
		$\Delta_{\text{v}}H$		(493–673)	77.9	508	A, EST	[1987STE/MAL, 1961LI/ROS, 1970DYK/VAN]
C ₁₉ H ₃₉ Cl	[62016-76-6]		1-chlorononadecane					
		$\Delta_{\text{v}}H$			114.7	298		[2006BOL/NER2]
		$\Delta_{\text{v}}H$		(483–673)	76.3	498	A	[1987STE/MAL, 1970DYK/VAN]
C ₁₉ H ₃₉ F	[1480-63-3]		1-fluorononadecane					
		$\Delta_{\text{v}}H$		(458–648)	72.5	473	A, EST	[1987STE/MAL, 1961LI/ROS, 1970DYK/VAN]
C ₁₉ H ₃₉ I	[62127-51-9]		1-iodononadecane					
		$\Delta_{\text{v}}H$		(506–673)	113.8	298	A,EST	[1987STE/MAL, 1961LI/ROS, 2006BOL/NER]
		$\Delta_{\text{v}}H$		(506–673)	79.1	521	A, EST	[1987STE/MAL, 1961LI/ROS, 1970DYK/VAN]
C ₁₉ H ₃₉ NO ₂	[6280-24-6]		N-hexadecyl lactamide					
		$\Delta_{\text{v}}H$		(423–508)	111	438	A	[1987STE/MAL]
C ₁₉ H ₃₉ NO ₂	[na]		N,N-dioctyl lactamide					
		$\Delta_{\text{v}}H$		(453–488)	99.3	468	A	[1987STE/MAL]
C ₁₉ H ₄₀	[629-92-5]		nonadecane					
		$\Delta_{\text{trs}}H$			12.7	294.8		
		$\Delta_{\text{fus}}H$			42.7	304.4	DSC	[2004MON/RAJ]
		$\Delta_{\text{trs}}H$			13.67	296		
		$\Delta_{\text{fus}}H$			47.4	305.3		[1996DOM/HEA]
		$\Delta_{\text{sub}}H$			143.6	298	C	[1972MOR3]
		$\Delta_{\text{sub}}H$		(288–303)	136.6	296		[1964MOR]
		$\Delta_{\text{v}}H$		(423–588)	76.2	438		[1994MOR/KOB]
		$\Delta_{\text{v}}H$			96.4	298		[1994RUZ/MAJ]
		$\Delta_{\text{v}}H$		(456–606)	73.0	471	A	[1987STE/MAL]
		$\Delta_{\text{v}}H$			95.8	298		[1971WIL/ZWO]
C ₁₉ H ₄₀	[1560-88-9]		2-methyloctadecane					
		$\Delta_{\text{v}}H$		(451–595)	67.5	466	A	[1987STE/MAL, 1999DYK/SVO, 1959TER/BRI]
C ₁₉ H ₄₀	[6561-44-0]		3-methyloctadecane					
		$\Delta_{\text{v}}H$		(455–597)	69.2	470	A	[1987STE/MAL, 1999DYK/SVO, 1959TER/BRI]
C ₁₉ H ₄₀	[10544-95-3]		4-methyloctadecane					
		$\Delta_{\text{v}}H$		(445–596)	63.3	460	A	[1987STE/MAL, 1999DYK/SVO, 1959TER/BRI]
C ₁₉ H ₄₀	[25117-35-5]		5-methyloctadecane					
		$\Delta_{\text{v}}H$		(445–595)	63.8	460	A	[1987STE/MAL, 1999DYK/SVO, 1959TER/BRI]
C ₁₉ H ₄₀	[61868-03-9]		5-methyloctadecane					
		$\Delta_{\text{v}}H$		(447–598)	64.1	462		[1999DYK/SVO, 1959TER/BRI]
		$\Delta_{\text{v}}H$		(493–598)	67.2	508	A	[1987STE/MAL]
C ₁₉ H ₄₀	[61868-09-5]		2,4-dimethylheptadecane					
		$\Delta_{\text{v}}H$		(444–574)	70.6	459	A	[1987STE/MAL, 1999DYK/SVO, 1959TER/BRI]
C ₁₉ H ₄₀	[102013-94-5]		2,4-dimethylheptadecane					
		$\Delta_{\text{v}}H$		(435–568)	67.3	450	A	[1987STE/MAL, 1999DYK/SVO, 1959TER/BRI]
C ₁₉ H ₄₀	[7225-66-3]		7-hexyltridecane					
		$\Delta_{\text{v}}H$		(411–444)	75.2	426	A	[1987STE/MAL]
C ₁₉ H ₄₀ O	[1454-84-8]		1-nonadecanol					
		$\Delta_{\text{fus}}H$			43.3	333.9	DSC	[2004VEN/CAL]
		$\Delta_{\text{trs}}H$			29.1	329.7		
		$\Delta_{\text{fus}}H$			43.3	333.9	DSC	[2002VEN/RAM]

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
			72.42	334.5		[2001VAN/OON2]
		Note: Value includes the enthalpy of a solid/solid transition that occurs at about 331 K				
			81.7	494	A	[1987STE/MAL]
			80.0	509	A	[1987STE/MAL]
C ₁₉ H ₄₀ O ₂	[7268-65-7]	1,19-nonadecanediol				
			37.1	358.9		
			35.7	373.9	DSC	[1999OGA/NAK]
C ₁₉ H ₄₀ S	[53193-23-0]	1-nonadecanethiol				
			79.2	517	EST	[1999DYK/SVO]
C ₁₉ H ₄₁ N	[14130-05-3]	nonadecylamine				
			72.7	547	A, EST	[1987STE/MAL, 1956MAN2]
C ₂₀ F ₄₂	[37589-57-4]	perfluoroicosane				
			0.67	149.5		
			11.25	202.9		
			80.33	437.9	DSC	[1986STA]
C ₂₀ H ₄ Cl ₄ F ₁₃ NO ₂	[433932-34-4]	4,5,6,7-dichloro-2-(4-n-tridecafluorohexylphenyl)isoindole-1,3-dione				
			45.1	512.8		[2002JOW/DIN]
C ₂₀ H ₆ Cl ₂ F ₁₃ NO ₂	[433932-32-2]	5,6-dichloro-2-(4-n-tridecafluorohexylphenyl)isoindole-1,3-dione				
			40.7	459.6		[2002JOW/DIN]
C ₂₀ H ₆ Cl ₂ F ₁₃ NO ₂	[433932-33-3]	4,7-dichloro-2-(4-n-tridecafluorohexylphenyl)isoindole-1,3-dione				
			18.4	413.2		
			19.9	417.2		[2002JOW/DIN]
C ₂₀ H ₁₀	[5821-51-2]	corannulene				
			17.3	542.3	DSC	[2002CHI/WEB]
			115.8	408	HSA	[2002CHI/WEB]
			119.5 ± 4.4	298	HSA	[2002CHI/WEB]
			116.3 ± 6.0	298	CGC+Fus	[2002CHI/WEB]
			115.5 ± 2.5	298	CGC	[2002CHI/WEB]
C ₂₀ H ₁₁ F ₁₄ N ₃	[502455-01-8]	2,2,3,3,4,4,4-heptafluoro-N-[2,2,3,3,4,4,4-heptafluoro-1-(phenylamino)-butylidene]-N'-phenylbutanimidamide				
			31.3	361		[2003SIE/WEB]
C ₂₀ H ₁₁ NO ₂	[63041-90-7]	6-nitrobenzo[a]pyrene				
			30.2	528.4	DSC	[2010KES/AUC]
C ₂₀ H ₁₂	[198-55-0]	perylene				
			31.88	551		[1996DOM/HEA, 1993ACR]
			132.6 ± 3.6	408	ME	[1998OJA/SUU]
			123.2	383	GS	[1995NAS/LEN]
			145.2 ± 2.5	298	C,ME	[1973GIG/MAL]
			125.5 ± 4.2	298	ME	[1967WAK/INO, 1970COX/PIL]
			139	418		[1958HOY/PEP, 1987STE/MAL]
			129.6 ± 2.1	415	ME	[1952INO/SHI]
			121.3	370	ME	[1951INO]
			119.5	298	CGC	[2008ZHA/UNH]
			123.1 ± 1.7	298	CGC	[2002CHI/WEB]
			89.9	398	GC	[2002LEI/CHA]

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
	Enthalpy							
C ₂₀ D ₁₂	[1520-96-3]	perylene - d ₁₂			119.5	298	CGC	[2008ZHA/UNH]
	$\Delta_{\text{v}}H$							
C ₂₀ H ₁₂	[50-32-8]	benzo[a]pyrene			14.7	451.8	DSC	[2010KES/AUC]
	$\Delta_{\text{fus}}H$				8.49	390.2		
	$\Delta_{\text{trs}}H$				17.32	454.2		[1991ACR]
	$\Delta_{\text{fus}}H$							
	$\Delta_{\text{sub}}H$	(313–453)		122.5	383	GS	[1995NAS/LEN]	
	$\Delta_{\text{sub}}H$	(358–431)		118.3	373	ME	[1987STE/MAL, 1974MUR/POL]	
	$\Delta_{\text{v}}H$			117.8 ± 1.0	298	CGC	[2008HAN/NUT]	
	$\Delta_{\text{v}}H$	(463–523)		105.0 ± 1.5	298	GC	[2006HAF/PAR]	
	$\Delta_{\text{v}}H$	(323–473)		91	398	GC	[2002LEI/CHA]	
$\Delta_{\text{v}}H$	(343–453)		95.5	398	GC	[1990HIN/BID2]		
C ₂₀ H ₁₂	[192-97-2]	benzo[e]pyrene			13.8	451.3	DSC	[2010KES/AUC]
	$\Delta_{\text{fus}}H$				2.51	426.2		
	$\Delta_{\text{trs}}H$				16.57	454.4		[1991ACR]
	$\Delta_{\text{fus}}H$							
	$\Delta_{\text{sub}}H$	(313–453)		117.9	383	GS	[1995NAS/LEN]	
	$\Delta_{\text{sub}}H$	(359–423)		119.1	373	ME	[1987STE/MAL, 1974MUR/POL]	
	$\Delta_{\text{v}}H$			118.2 ± 0.3	298	CGC	[2008HAN/NUT]	
	$\Delta_{\text{v}}H$	(463–523)		105.0 ± 1.5	298	GC	[2006HAF/PAR]	
	$\Delta_{\text{v}}H$	(343–453)		92	398	GC	[1990HIN/BID2]	
C ₂₀ H ₁₂	[207-08-9]	benzo[k]fluoranthene			32.4	489.7	DSC	[2010KES/AUC]
	$\Delta_{\text{fus}}H$				27.5	490.6		[2002DIO/MIN]
	$\Delta_{\text{fus}}H$							
	$\Delta_{\text{sub}}H$	(387–423)		124.2 ± 4.7	298	ME	[2002DIO/MIN]	
	$\Delta_{\text{sub}}H$	(363–430)		130	378		[1987STE/MAL]	
	$\Delta_{\text{sub}}H$			120 ± 10		TE	[1983FER/QUA]	
	$\Delta_{\text{v}}H$			117.4 ± 1.1	298	CGC	[2008HAN/NUT]	
	$\Delta_{\text{v}}H$	(463–513)		105.5 ± 1.5	298	GC	[2006HAF/PAR]	
	$\Delta_{\text{v}}H$	(323–473)		88.5	398	GC	[2002LEI/CHA]	
C ₂₀ H ₁₂	[205-99-2]	benzo[b]fluoranthene			19.6	441.5	DSC	[2010KES/AUC]
	$\Delta_{\text{fus}}H$							
	$\Delta_{\text{sub}}H$	(313–453)		119.2	383	GS	[1995NAS/LEN]	
	$\Delta_{\text{v}}H$			116.8 ± 1.6	298	CGC	[2008HAN/NUT]	
	$\Delta_{\text{v}}H$	(463–513)		104.0 ± 1.5	298	GC	[2006HAF/PAR]	
	$\Delta_{\text{v}}H$	(323–473)		89.7	398	GC	[2002LEI/CHA]	
C ₂₀ H ₁₂	[205-82-3]	benzo[j]fluoranthene			17.9	438.3	DSC	[2010KES/AUC]
	$\Delta_{\text{fus}}H$							
C ₂₀ H ₁₂ BrNO ₄	[59722-76-8]	1-amino-2-(4-bromophenoxy)-4-hydroxy-9,10-anthraquinone			163.6	413		[1978NIS/ISH]
$\Delta_{\text{sub}}H$	(473–543)							
C ₂₀ H ₁₂ O	[13345-21-6]	3-hydroxybenzo[a]pyrene			24.1	469.6	DSC	[2010KES/AUC]
	$\Delta_{\text{fus}}H$							
C ₂₀ H ₁₃ N	[194-59-2]	7H-dibenzo[c,g]carbazole			20.1	429.8	DSC	[2010KES/AUC]
	$\Delta_{\text{fus}}H$							
C ₂₀ H ₁₃ NO ₂	[109392-90-7]	phenyl acridine-9-carboxylate						

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	Enthalpy							
		$\Delta_{\text{fus}}H$			39.2	464	DSC	[2010KRZ/MAL]
C ₂₀ H ₁₃ NO ₄	[17418-58-5]		1-amino-4-hydroxy-2-phenoxy-9,10-anthraquinine (Disperse Red 60)					
		$\Delta_{\text{fus}}H$			30.79	458.2		[1991BAU/WEB]
		$\Delta_{\text{sub}}H$	(359–366)		152.5	362.5	A	[1987STE/MAL, 1977EIB/TRO]
		$\Delta_{\text{sub}}H$			141.8			[1984KAR/KRU]
C ₂₀ H ₁₃ N ₅ O ₃	[194785-03-0]		1-(2'-nitrobenzylidene)-2-phenazinoylhydrazine					
		$\Delta_{\text{fus}}H$	(373–453)		103.8	413		[1978NIS/ISH]
C ₂₀ H ₁₃ N ₅ O ₃	[194785-02-9]		1-(4'-nitrobenzylidene)-2-phenazinoylhydrazine					
		$\Delta_{\text{fus}}H$			44.8	548.2	DSC	[1997CIO/MEL]
C ₂₀ H ₁₄	[602-55-1]		9-phenylanthracene					
		$\Delta_{\text{sub}}H$	(313–453)		118.7	383	GS	[1995NAS/LEN]
		$\Delta_{\text{sub}}H$	(352–395)		119.7		TE	[1974SHI/GRE]
		$\Delta_{\text{sub}}H$	(353–426)		115.3	368		[1958KLO]
		Δ_vH	(323–473)		91.6	398	GC	[2002LEI/CHA]
		Δ_vH	(430–510)		84.4	445	A	[1987STE/MAL]
C ₂₀ H ₁₄	[477-75-8]		9,10-dihydro-9,10-(1',2') benzoanthracene (tryptycene)					
		$\Delta_{\text{fus}}H$			30.29	527.2		[1996DOM/HEA]
		$\Delta_{\text{sub}}H$			104.6 ± 12.6			[1973ROD/WES, 1977PED/RYL]
C ₂₀ H ₁₄	[11068-27-2]		binaphthalene					
		$\Delta_{\text{sub}}H$	(313–453)		138.3	383	GS	[1995NAS/LEN]
C ₂₀ H ₁₄	[604-53-5]		1,1'-binaphthyl					
		$\Delta_{\text{fus}}H$			19.2	431.2	DSC	[2005SAI/MAR]
		$\Delta_{\text{fus}}H$ (I)			30.5	418.2		
	$\Delta_{\text{fus}}H$ (II)			23.7	431.2	DSC	[1975WIL/PIN]	
C ₂₀ H ₁₄	[612-78-2]		β,β' -binaphthyl					
		$\Delta_{\text{fus}}H$			38.9	461.2		[1996DOM/HEA]
C ₂₀ H ₁₄ N ₂ O ₂	[4395-65-7]		1-anilino-4-aminoanthraquinone					
		$\Delta_{\text{sub}}H$			138.6			[1984KAR/KRU]
	$\Delta_{\text{sub}}H$	(473–553)		135.1		GS	[1977NIS/ISH, 1978NIS/ISH]	
C ₂₀ H ₁₄ N ₂ O ₄	[56405-27-7]		1-amino-2-(4-aminophenoxy)-4-hydroxy-9,10-anthraquinone					
		$\Delta_{\text{sub}}H$	(373–453)		U 50.2	413		[1978NIS/ISH]
C ₂₀ H ₁₄ N ₄	[101-60-0]		21 <i>H</i> ,23 <i>H</i> -porphine					
		$\Delta_{\text{sub}}H$	(424–507)		87 ± 3		Fluoresc	[2004STE/STI]
C ₂₀ H ₁₄ N ₄ O	[194784-97-9]		1-benzylidene-2-phenazinoylhydrazine					
		$\Delta_{\text{fus}}H$			45.76	512.1	DSC	[1997CIO/MEL]
C ₂₀ H ₁₄ O ₄	[94-01-9]		dibenzoyl resorcinol					
		$\Delta_{\text{sub}}H$	(323–399)		165.8	338	A	[1987STE/MAL]
		Δ_vH	(399–493)		76.0	414	A, UV	[1987STE/MAL, 1960SCH/HIR]
C ₂₀ H ₁₄ O ₄	[77-09-8]		phenolphthalein					
		$\Delta_{\text{fus}}H$			51.05	534		[1996DOM/HEA]
C ₂₀ H ₁₅ BrN ₄ O ₆	[na]		3,5-dinitro-4-(4-methoxyphenyl)aminobenzoyl (4-bromophenyl)amide					

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	Enthalpy							
		$\Delta_{\text{fus}}H$			50.67	522.5		[1999KOR/LEV]
C ₂₀ H ₁₅ BrN ₄ O ₆	[235114-51-9]		3,5-dinitro-2-(4-methoxyphenyl)aminobenzoyl (4-bromophenyl)amide		38.0	478.9		[1999KOR/LEV]
		$\Delta_{\text{fus}}H$						
C ₂₀ H ₁₅ F ₃	[68643-31-2]		1,1,1-trifluoro-2,2,2-triphenylethane		30.33	440.3		[1997SCH/VER]
		$\Delta_{\text{fus}}H$						
		$\Delta_{\text{sub}}H$			112.3 ± 1.0	298		[1997SCH/VER]
C ₂₀ H ₁₅ F ₃ O	[145698-50-6]		4-propoxy-4'-trifluoromethyldiphenyldiacetylene		18.81	315.9	DSC	[1993JUA/CHE]
		$\Delta_{\text{fus}}H$						
C ₂₀ H ₁₅ O ₅ P	[803-19-0]		bis(4-carboxyphenyl)phenylphosphine oxide		17.6	610.6	DSC	[2000WAN/WAN]
		$\Delta_{\text{fus}}H$						
C ₂₀ H ₁₆	[58-72-0]		triphenylethylene					
		$\Delta_{\text{fus}}H$			20.58	339.9	DSC	[1999VER/EBE]
		$\Delta_{\text{fus}}H$			20.35	341		[1998HIK/OKA]
		$\Delta_{\text{sub}}H$	(323–339)		110.1 ± 1.9	331	GS	[1999VER/EBE]
		$\Delta_{\text{sub}}H$	(323–339)		112.2 ± 1.9	298	GS	[1999VER/EBE]
		Δ_vH	(346–377)		88.0 ± 0.9	362	GS	[1999VER/EBE]
		Δ_vH	(346–377)		91.8 ± 0.9	298	GS	[1999VER/EBE]
		Δ_vH	(353–443)		89.7	398		[1989SAS/NGU]
C ₂₀ H ₁₆	[313-74-6]		7,12-dimethylbenz[a]anthracene					
		$\Delta_{\text{sub}}H$	(379–390)		135		A	[1987STE/MAL, 1964KEL/RIC]
		Δ_vH	(379–396)		107.8		A	[1987STE/MAL, 1964KEL/RIC]
		Δ_vH	(323–473)		88.9	398	GC	[2002LEI/CHA]
		Δ_vH	(396–408)		112.9	402	A, ME	[1987STE/MAL, 1964RAT/SHR, 1999DYK/SVO]
C ₂₀ H ₁₆	[3697-27-6]		5,6-dimethylchrysene					
		$\Delta_{\text{sub}}H$			130 ± 1.3			[1966GEI/QUI, 1970COX/PIL]
		$\Delta_{\text{sub}}H$			134 ± 1.3			[1966GEI/QUI, 1970COX/PIL]
		$\Delta_{\text{sub}}H$	(379–408)		135 ± 2.4	394	ME	[1964KEL/RIC]
		Δ_vH	(380–394)		121.7	387	A	[1987STE/MAL]
C ₂₀ H ₁₆	[313-74-6]		1',9-dimethyl-1,2-benzanthracene					
		$\Delta_{\text{sub}}H$			112.5 ± 3.3		ME	[1965KAR/KYB, 1970COX/PIL]
C ₂₀ H ₁₆	[316-51-8]		3',6-dimethyl-1,2-benzanthracene					
		$\Delta_{\text{sub}}H$			112.5 ± 3.3		ME	[1965KAR/KYB, 1970COX/PIL]
C ₂₀ H ₁₆ F ₂	[145698-36-8]		4- <i>n</i> -butyl-3',4'-difluorodiphenyldiacetylene		24.33	340.8	DSC	[1993JUA/CHE]
		$\Delta_{\text{fus}}H$						
C ₂₀ H ₁₆ N ₂ O ₅	[19685-09-7]		20-(S)-10-hydroxycamptothecin		53.38	467.2	DSC	[2010KUN/SAV]
		$\Delta_{\text{fus}}H$						
C ₂₀ H ₁₆ O ₄ S ₂	[3263-31-8]		6-ethoxy-2-(6-ethoxy-3-oxobenzothien-2(3 <i>H</i>)-ylidene)benzo[b]-thiophen-3(2 <i>H</i>)-one (C.I. Vat Orange 5)		65	577	GS	[1986NIS/AND]
		$\Delta_{\text{sub}}H$	(519–634)					
C ₂₀ H ₁₇ FO ₃ S	[38194-50-2]		(<i>Z</i>)-5-fluoro-2-methyl-1-[<i>p</i> -(methylsulfinyl)benzylidene]indene-3-acetic acid (sulindac)		33.4	460.2	DSC	[2006WAS/HOL]
		$\Delta_{\text{fus}}H$						
C ₂₀ H ₁₇ F ₂₅	[90499-31-3]		1,1,1,2,2,3,3,4,4,5,5,6,6,7,7,8,8,9,9,10,10,11,11,12,12-pentacosafuoroicosane		2.4	192		
		$\Delta_{\text{trs}}H$						

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
	$\Delta_{\text{trs}}H$		6.4	329		
	$\Delta_{\text{fus}}H$		23.7	361	DSC	[1991HOP/MOL]
	$\Delta_{\text{trs}}H$		5.6	324.2		
	$\Delta_{\text{fus}}H$		21.9	355.2	DSC	[1986RUS/RAB]
C ₂₀ H ₁₇ N ₃ O ₄	[na]	4,11-diamino-2-butyl-1 <i>H</i> -naphth[2,3- <i>f</i>]isoindole-1,3,5,10(2 <i>H</i>)-tetraone				
	$\Delta_{\text{fus}}H$		24.85	490.2		[1991BAU/WEB]
C ₂₀ H ₁₇ N ₅ O ₃	[na]	6-(2-naphthyl)-3,9-dihydro-3-[(2-hydroxyethoxy)methyl]-9-oxo-5 <i>H</i> -imidazol[1,2- <i>a</i>]pyrine				
	$\Delta_{\text{fus}}H$		37.64	493.4	DSC	[1999ZIE/GOL]
C ₂₀ H ₁₈	[15271-39-6]	1,1,1-triphenylethane				
	$\Delta_{\text{fus}}H$		19.95	375.9	DSC	[1999VER3]
	$\Delta_{\text{sub}}H$	(338–363)	108.6 ± 0.9	298	GS	[1999VER3]
	$\Delta_{\text{sub}}H$	(338–363)	105.4 ± 0.9	351	GS	[1999VER3]
C ₂₀ H ₁₈	[1520-42-9]	1,1,2-triphenylethane				
	$\Delta_{\text{fus}}H$		24.39	328.2	DSC	[1999VER3]
	$\Delta_{\text{sub}}H$	(335–368)	89.0 ± 0.5	351	GS	[1999VER3]
	$\Delta_{\text{sub}}H$	(335–368)	92.2 ± 0.5	298	GS	[1999VER3]
C ₂₀ H ₁₈ N ₂ O ₂	[7385-67-3]	9-(diethylamino)-5 <i>H</i> -benzo[<i>a</i>]phenoxazin-5-one (nile red)				
	$\Delta_{\text{sub}}H$	(427–515)		66 ± 2	Fluoresc	[2004STE/STI]
C ₂₀ H ₁₈ O ₂	[na]	2-fluorenyl-2-methyl-1,3-cyclohexanedione				
	$\Delta_{\text{fus}}H$		35.7	448.2		[1995NOL/VER]
C ₂₀ H ₁₈ O ₆	[170464-52-5]	9-fluorenyl- <i>tris</i> (methoxycarbonyl)methane				
	$\Delta_{\text{fus}}H$		32.3	407.2		[1995RAK/VER]
	$\Delta_{\text{sub}}H$		132.6	298	GS	[1995RAK/VER]
C ₂₀ H ₁₉ BrS	[na]	2- <i>n</i> -butyl-5-(4-bromobiphenyl-4-yl)thiophene				
	$\Delta_{\text{fus}}H$		21.4	501.4		[1993BRE/DUN]
C ₂₀ H ₁₉ F ₃ O	[172424-72-5]	4- <i>n</i> -hexyloxy-2',3',4'-trifluorodiphenylacetylene				
	$\Delta_{\text{fus}}H$		30.8	322	DSC	[1995HSU/TSA]
C ₂₀ H ₁₉ N ₇ O ₄	[41642-51-7]	Disperse Blue 165				
	$\Delta_{\text{sub}}H$	(464–484)	90.7	474	GS	[1989NIS/AND]
Note: The molecular structure of the compound given in the paper had a Cl group and only one CN functional group, which is different than the molecular structure given in Scifinder Scholar for Disperse Blue 165. The molecular formula for the structure in the paper is C ₂₀ H ₂₁ ClN ₆ O ₃ . The molecular weight given in the paper, 405, agrees with the molecular formula of C ₂₀ H ₁₉ N ₇ O ₄ . We have assumed that the chemical name is correct, but that the authors misdrew the molecular structure.						
C ₂₀ H ₂₀	[26902-55-6]	hexacyclopentane				
	$\Delta_{\text{v}}H$	(333–373)	85.8 ± 0.2	298	GS	[1995CHI/HES]
C ₂₀ H ₂₀	[89683-62-5]	pagodane (undecacyclo[9.9.0.0 ^{1,5} .0 ^{2,12} .0 ^{2,18} .0 ^{3,7} .0 ^{6,10} .0 ^{8,12} .0 ^{11,15} .0 ^{13,17} .0 ^{16,20}]eicosane)				
	$\Delta_{\text{sub}}H$	(418–473)	90.2 ± 2.3	446	T	[1994BEC/RUE]
C ₂₀ H ₂₀ FNO ₃	[94610-85-2]	4-cyano-3-fluorophenyl 4-hexyloxybenzoate				
	$\Delta_{\text{fus}}H$		35.56	332.7		[1984KEL]
C ₂₀ H ₂₀ F ₂	[145698-44-8]	4-hexyl-3',4'-difluorodiphenylacetylene				
	$\Delta_{\text{fus}}H$		24.3	314.9	DSC	[1995HSU/TSA]
C ₂₀ H ₂₀ F ₂ O	[145698-45-9]	4-hexyloxy-3',4'-difluorodiphenylacetylene				
	$\Delta_{\text{fus}}H$		33.1	323.6	DSC	[1995HSU/TSA]

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
C ₂₀ H ₂₀ F ₂ O	[172424-68-9]	4-hexyloxy-2',4'-difluorodiphenylacetylene	34.1	320.9	DSC	[1995HSU/TSA]
C ₂₀ H ₂₀ NP	[47182-04-7]	N-ethyl triphenylphosphine imine	75.3 ± 8.4	298		[1982PIL/SKI, 1960CLA/FOW]
C ₂₀ H ₂₀ O ₂	[160731-88-4]	2-diphenylmethyl-2-ethyl-1,3-cyclopentandione	28.2	382.2		[1995NOL/VER]
		(342–377)	122.8 ± 0.7	360	T	[1995NOL/VER]
C ₂₀ H ₂₀ O ₃	[87051-12-5]	4,4-dimethyl-1,8-diphenyl-2,3,5-trioxabicyclo[4.3.0]non-7-ene	22.1	369.2	DSC	[1991JEF/JAB]
C ₂₀ H ₂₀ O ₆	[170464-52-5]	1,1,1-tris(methoxycarbonyl)-2,2-diphenylethane	36.1	414.2		[1995RAK/VER]
			136	298	GS	[1995RAK/VER]
C ₂₀ H ₂₁ ClO ₄	[49562-28-9]	2-[4-(4-chlorobenzoyl)phenoxy]-2-methylpropanoic acid, isopropyl ester (fenofibrate)	32.4	353.7		[2002ZHO/ZHA]
C ₂₀ H ₂₁ F ₁₉ O	[144986-71-0]	1,1,1,2,2,3,3,4,4,5,5,6,6,7,7,8,8,9,9-nonadecafluoro-10-eicosanone	53.17	317.9		[1993VIL/HAM]
C ₂₀ H ₂₁ F ₂₁	[90499-29-9]	1,1,1,2,2,3,3,4,4,5,5,6,6,7,7,8,8,9,9,10,10-heneicosafuoroicosane	4	317		
			24.4	337	DSC	[1991HOP/MOL]
			2.2	306.5		
			26.7	336.7		[1989VIN/RUS]
C ₂₀ H ₂₁ NO ₃ S	[313057-11-3]	4-(7-octenyloxy)phenyl 5-cyano-2-thiophene carboxylate	68.2	332.7	DSC	[2000WU/WAN]
C ₂₀ H ₂₁ N ₃ O ₃	[198629-74-2]	pyrimethanil phenoxycetate (81–380)	34.28	349.4	AC	[2006SUN/LIU]
C ₂₀ H ₂₁ N ₃ O ₅ S	[87027-09-6]	2-methyl-1,1-dioxido-3-[(2-pyridinylamino)carbonyl]-2 <i>H</i> -1,2-benzothiazin-4-yl, 2,2-dimethylpropanoic acid ester (piroxicam pivalate)	32.74	427	DSC	[1998GIO/GAZ]
C ₂₀ H ₂₂ N ₂ O	[263896-41-9]	2,3-dihydro-2-[(3-exo)-8-methyl-8-azabicyclo[3.2.1]oct-3-yl]-1 <i>H</i> -benz[e]isoindol-1-one	29.1	448.2	DSC	[2006CAP/TRA]
C ₂₀ H ₂₂ N ₂ O ₄	[na]	1,4-bis(propylamino)anthraquinone (409–463)	118.3	424	A	[1987STE/MAL]
C ₂₀ H ₂₂ O ₂	[160731-83-9]	3-diphenylmethyl-3-ethyl-2,4-pentanedione	34.7	388.2		[1995NOL/VER]
		(349–387)	122.3 ± 1.5	368	T	[1995NOL/VER]
C ₂₀ H ₂₃ FN ₂ O	[2354-61-2]	1-(4-fluorophenyl)-4-[4-phenyl-1-piperazinyl]-1-butanone (butropipazone)	36.7	387.2		
			34.7	363.7	DSC	[1981DRA/AZI]
C ₂₀ H ₂₃ F ₁₉ O	[144986-72-1]	1,1,1,2,2,3,3,4,4,5,5,6,6,7,7,8,8,9,9-nonadecafluoro-10-eicosanol	3.6	346.2		
			33.5	356		[1992VIL/WEI]
C ₂₀ H ₂₃ NO ₄	[16590-41-3]	17-(cyclopropylmethyl)-4,5-epoxy-3,14-dihydroxy-morphinan-6-one (naltrexone)	15.64	448.2	DSC	[2004PIL/HAM]

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
	Enthalpy							
		$\Delta_{\text{fus}}H$			14.8	448.9	DSC	[2004HAM/HAM]
C ₂₀ H ₂₃ N ₃ O ₉	[142489-47-2]		N-[N-[[2-(1,3-dihydro-1,3-dioxo-2H-isoindol-2-yl)ethoxy]acetyl]-(l)-alanyl]-(d)-glutamic acid		54.53	NA		[1999ZAD/KER]
C ₂₀ H ₂₄	[115182-07-2]		8-(4-biphenyl)-1-octene		21	291.5	DSC	[1989MAL/KAN]
C ₂₀ H ₂₄ O ₂	[102607-41-0]		7-methyl-3-(1-methylethyl)-8-(4-methyl-3-pentenyl)-1,2-naphthalenedione (saprorthoquinone)		23.09	369.2		[1992HUA/ZHO]
C ₂₀ H ₂₄ O ₂	[57-63-6]		19-norpregna-1,3,5(10)-trien-20-yne-3,17-diol (ethinyl estradiol)		27.57	456.2		[2002VAN/KRU]
C ₂₀ H ₂₄ O ₃	[901-93-9]		3-(acetyloxy)-estra-1,3,5(10)-trien-17-one		15	399	DSC	[1990YAN/EIR]
C ₂₀ H ₂₄ O ₄ S	[313057-15-7]		4-(7-octenyloxy)phenyl 5-methoxy-2-thiophene carboxylate		76.57	332.8	DSC	[2000WU/WAN]
C ₂₀ H ₂₄ O ₆	[14187-32-7]		dibenzo-18-crown-6		57.45	435.8		[1998DOM]
		$\Delta_{\text{sub}}H$			178.8 ± 6.9	298	CGC-DSC	[2000NIC/ORF]
		$\Delta_{\text{v}}H$			137.0 ± 7.4	298	CGC	[2000NIC/ORF]
C ₂₀ H ₂₆	[1625-91-8]		4,4'-di- <i>tert</i> -butylbiphenyl		20.0	400.8	DSC	[2009MEL/PIM]
		$\Delta_{\text{trs}}H$			1.0	322		
		$\Delta_{\text{fus}}H$			18.8	402	DSC	[2002NAT/JES]
		$\Delta_{\text{sub}}H$			106.8 ± 3.2	298	C	[2009MEL/PIM]
		$\Delta_{\text{v}}H$			86.2 ± 3.2	298	S-V	[2009NTI/CHA]
C ₂₀ H ₂₆ O	[68-22-4]		19-nor-17 α -ethynyltestosterone		39.6	479		[1996DOM/HEA]
C ₂₀ H ₂₆ O ₂	[38107-76-5]		2- <i>tert</i> -butyl-4-methoxymethyl-6- α -methylbenzylphenol		29.4	371.7	DTA	[1972INO/LIA]
C ₂₀ H ₂₆ O ₃	[57078-10-1]		1,2,2-trimethyl-3-[(5,6,7,8-tetrahydro-2-naphthalenyl)carbonyl]cyclopentanecarboxylic acid		22.94	421.3	DSC	[1992TER/PAU]
C ₂₀ H ₂₆ O ₄	[84-61-7]		dicyclohexyl phthalate		97	406	A	[1987STE/MAL]
C ₂₀ H ₂₇ NO ₄	[135531-41-8]		(-)-1-[[2-(3,4-dimethoxyphenyl)ethyl]amino]-3-(3-methyl-phenoxy)-2-propanol (bevantolol)		43.22	348.3		[1999LI/ZEL]
C ₂₀ H ₂₇ NO ₄	[59170-23-9]		(+)-1-[[2-(3,4-dimethoxyphenyl)ethyl]amino]-3-(3-methyl-phenoxy)-2-propanol (bevantolol)		45.9	360.6		[1999LI/ZEL]
C ₂₀ H ₂₇ N ₅ O ₂	[73963-72-1]		6-[4-(1-cyclohexyl-1H-tetrazol-5-yl)butoxy]-3,4-dihydro-2(1H)-quinolinone (cilostazol)		4.72	432		
		$\Delta_{\text{fus}}H$ (I)			3.89	408.8		
		$\Delta_{\text{fus}}H$ (III)			4.28	419		[2002STO/BEH]
		Note: Reported fusion enthalpies are very small. Compound likely exhibits solid-solid transition(s) at lower temperature(s).						
C ₂₀ H ₂₈	[55000-56-1]		2-butyl-3-hexylnaphthalene		80.8	437		[1963DIX/YAR, 1984BOU/FRI, 1999DYK/SVO]

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		T _m (K)	Method	Reference
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)			
C ₂₀ H ₂₈	[55000-55-0]	7-butyl-1-hexylnaphthalene		433		[1963DIX/YAR, 1984BOU/FRI, 1999DYK/SVO]
	$\Delta_v H$	(418–481)	78.1			
C ₂₀ H ₂₈	[55000-53-8]	1,4-dimethyl-5-octylnaphthalene		447		[1963DIX/YAR, 1984BOU/FRI, 1999DYK/SVO]
	$\Delta_v H$	(432–496)	81.6			
C ₂₀ H ₂₈	[55000-54-9]	2,6-dimethyl-3-octylnaphthalene		445		[1963DIX/YAR, 1984BOU/FRI, 1999DYK/SVO]
	$\Delta_v H$	(430–494)	80.8			
C ₂₀ H ₂₈ N ₂	[na]	1-(4-cyanophenyl)-4-n-heptylpiperidine		326.2		[1991SHE/WEI]
	$\Delta_{\text{fus}} H$		30.2			
C ₂₀ H ₂₈ O ₂	[112018-00-5]	1-[3,5-di- <i>tert</i> -butyl-4-hydroxyphenyl]-5-hexyn-1-one (tebufelone)		342.2		[1993KEL/SAK]
	$\Delta_{\text{fus}} H$		25.14			
C ₂₀ H ₂₈ O ₃	[3129-42-8]	testosterone formate		398		[1994REG/CHM]
	$\Delta_{\text{fus}} H$		26.36			
C ₂₀ H ₂₈ O ₅	[104225-29-8]	3-(3,4-diethoxybenzoyl)-1,2,2-trimethylcyclopentanecarboxylic acid		389.3	DSC	[1992TER/PAU]
	$\Delta_{\text{fus}} H$		29.07			
C ₂₀ H ₃₀	[26902-55-6]	hexacyclopropylethane		109.0 ± 2.1		[1984BER/BEC]
	$\Delta_{\text{sub}} H$					
C ₂₀ H ₃₀	[3732-31-8]	1,1'-biadamantane		336.3		
	$\Delta_{\text{trs}} H$		1.15			
	$\Delta_{\text{trs}} H$		1.3	509.6		
	$\Delta_{\text{fus}} H$		70 ± 10	561	AC	[2007KAR/KAB]
	$\Delta_{\text{sub}} H$	(393–443)	109.1 ± 1.3	417.8	ME	[2007KAR/KAB]
	$\Delta_{\text{sub}} H$	(393–443)	113.8 ± 1.4	298	ME	[2007KAR/KAB]
C ₂₀ H ₃₀ N ₄ O ₄	[197300-58-6]	1,1'-(1,10-decanediyl)bisthymine		455		[2002ITA/KAM]
	$\Delta_{\text{fus}} H$		42.56			
C ₂₀ H ₃₀ O ₂	[58-18-4]	17-methyl testosterone		439		[1997CEN/MEL]
	$\Delta_{\text{fus}} H$		27.8			
C ₂₀ H ₃₀ O ₄	[84-75-3]	dihexyl phthalate		468	A	[1987STE/MAL]
	$\Delta_v H$	(453–533)	92	358	A, ME	[1987STE/MAL, 1948SMA/SMA]
	$\Delta_v H$	(343–387)	103			
C ₂₀ H ₃₂	[66538-96-3]	1,2,3,4-tetrahydro-6-butyl-7-hexylnaphthalene		428		[1963DIX/YAR, 1984BOU/FRI, 1999DYK/SVO]
	$\Delta_v H$	(413–475)	78.1			
C ₂₀ H ₃₂	[66205-02-5]	1,2,3,4-tetrahydro-7-butyl-1-hexylnaphthalene		424		[1963DIX/YAR, 1984BOU/FRI, 1999DYK/SVO]
	$\Delta_v H$	(409–471)	76.7			
C ₂₀ H ₃₂	[55255-59-9]	1,2,3,4-tetrahydro-2,6-dimethyl-7-octylnaphthalene		433		[1963DIX/YAR, 1984BOU/FRI, 1999DYK/SVO]
	$\Delta_v H$	(418–480)	79.4			
C ₂₀ H ₃₂	[55255-58-8]	1,2,3,4-tetrahydro-5,8-dimethyl-1-octylnaphthalene		434		[1963DIX/YAR, 1984BOU/FRI, 1999DYK/SVO]
	$\Delta_v H$	(419–481)	78.6			
C ₂₀ H ₃₂	[na]	10,10,13,13-tetramethylcyclohexadeca-1,5-diyne		323.2		[1975BJO/BOR]
	$\Delta_{\text{fus}} H$		18.83			
C ₂₀ H ₃₂ O ₄	[175848-66-5]	2,5-di- <i>n</i> -heptyloxy-1,4-benzoquinone		275.8		
	$\Delta_{\text{trs}} H$		3.6			
	$\Delta_{\text{trs}} H$		17.3	372.5		
	$\Delta_{\text{fus}} H$		38.4	406.2	DSC	[1996KEE/VAN]

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	Enthalpy							
C ₂₀ H ₃₄	[55255-70-4]		9-cyclohexyltetradecahydroanthracene	(419–488)	74.5	434	A	[1987STE/MAL]
C ₂₀ H ₃₄	[1459-10-5]		tetradecylbenzene	(485–665)	74.5	500		[1999DYK/SVO]
		$\Delta_v H$			99.6	298		[1971WIL/ZWO]
C ₂₀ H ₃₄ O ₂	[1191-41-9]		ethyl linolenate	(447–491)	72.7	462	A	[1987STE/MAL]
C ₂₀ H ₃₄ O ₁₁	[na]		diethylene glycol dicarboxylic acid, di[1-(butoxycarbonyl)ethyl] ester	(433–525)	103.6	448	A	[1987STE/MAL]
C ₂₀ H ₃₄ O ₁₁	[na]		diethylene glycol dicarboxylic acid, di[1-(sec-butoxycarbonyl)ethyl] ester	(418–513)	103.1	433	A	[1987STE/MAL]
C ₂₀ H ₃₄ O ₁₁	[na]		diethylene glycol dicarboxylic acid, di[1-(isobutoxycarbonyl)ethyl] ester	(415–513)	103.1	430	A	[1987STE/MAL]
C ₂₀ H ₃₆ N ₂	[85688-86-4]		tetraisobutylsuccinonitrile		34.31	360.2		[1983BAR/BEC]
C ₂₀ H ₃₆ O ₂	[544-35-4]		ethyl linoleate	(448–497)	72.6	463	A	[1987STE/MAL]
C ₂₀ H ₃₆ O ₂	[14113-56-5]		1,10-cycloeicosanedione		55.06	327.2		[1972ALV/BOR]
C ₂₀ H ₃₆ O ₂	[na]		1,9-cyclohexadecanedione <i>bis</i> ethylene ketal		42.13	404.2		[1972ALV/BOR]
C ₂₀ H ₃₆ O ₆	[na]		(<i>syn-cis/anti-cis</i>) dicyclohexano-18-crown-6		124.2 ± 4.0	298	CGC	[2000NIC/ORF]
C ₂₀ H ₃₈	[66455-55-8]		2-butyl-3-hexyldecahydronaphthalene	(407–472)	76.9	422		[1963DIX/YAR, 1984BOU/FRI, 1999DYK/SVO]
C ₂₀ H ₃₈	[66455-54-7]		7-butyl-1-hexyldecahydronaphthalene	(407–467)	80.0	422		[1963DIX/YAR, 1984BOU/FRI, 1999DYK/SVO]
C ₂₀ H ₃₈	[54964-83-9]		1,4-dimethyl-5-octyldecahydronaphthalene	(404–466)	73.9	419		[1963DIX/YAR, 1984BOU/FRI, 1999DYK/SVO]
C ₂₀ H ₃₈	[54964-85-1]		2,6-dimethyl-3-octyldecahydronaphthalene	(406–469)	76.4	421		[1963DIX/YAR, 1984BOU/FRI]
C ₂₀ H ₃₈	[26527-76-4]		3,4-dicyclohexyl-3,4-dimethylhexane	(343–365)	78.4	359		[1999DYK/SVO, 1980BEC/KRA]
C ₂₀ H ₃₈	[765-27-5]		1-eicosyne	(473–651)	68.9	488		[1999DYK/SVO]
C ₂₀ H ₃₈	[61847-99-2]		2-eicosyne	(480–661)	69.8	495		[1999DYK/SVO]
C ₂₀ H ₃₈	[61866-66-6]		3-eicosyne	(470–648)	68.4	485		[1999DYK/SVO]
C ₂₀ H ₃₈ O	[29171-23-1]		3,7,11,15-tetramethyl-1-hexadecyn-3-ol	(403–457)	43.8 ± 1.9	430		[1988BAG/GUR]
C ₂₀ H ₃₈ O ₂	[111-62-6]		ethyl oleate	(384–481)	92.4	399	A	[1987STE/MAL]

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		T _m (K)	Method	Reference
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)			
C ₂₀ H ₃₈ O ₂	[2495-27-4] $\Delta_v H$	hexadecyl methacrylate (431–541)	73.1	446	A	[1987STE/MAL]
C ₂₀ H ₃₈ O ₂	[na] $\Delta_v H$	(Z) 3-octadecenyl acetate (393–438)	108.7	298	GC	[1997KOU/HOS, 2000OVA/KOU]
C ₂₀ H ₃₈ O ₂	[na] $\Delta_v H$	(E) 3-octadecenyl acetate (393–438)	109.3	298	GC	[1997KOU/HOS, 2000OVA/KOU]
C ₂₀ H ₃₈ O ₂	[693-80-1] $\Delta_v H$	(Z) 9-octadecenyl acetate (393–438)	107.8	298	GC	[1997KOU/HOS, 2000OVA/KOU]
C ₂₀ H ₃₈ O ₂	[22147-38-2] $\Delta_v H$	(E) 9-octadecenyl acetate (393–438)	108.7	298	GC	[1997KOU/HOS, 2000OVA/KOU]
C ₂₀ H ₃₈ O ₂	[6186-98-7] $\Delta_v H$	(Z) 11-octadecenyl acetate (393–438)	108.4	298	GC	[1997KOU/HOS, 2000OVA/KOU]
C ₂₀ H ₃₈ O ₂	[69282-64-0] $\Delta_v H$	(E) 11-octadecenyl acetate (393–438)	109.1	298	GC	[1997KOU/HOS, 2000OVA/KOU]
C ₂₀ H ₃₈ O ₂	[60037-58-3] $\Delta_v H$	(Z) 13-octadecenyl acetate (393–438)	108.7	298	GC	[1997KOU/HOS, 2000OVA/KOU]
C ₂₀ H ₃₈ O ₂	[na] $\Delta_v H$	(E) 13-octadecenyl acetate (393–438)	109.8	298	GC	[1997KOU/HOS, 2000OVA/KOU]
C ₂₀ H ₃₈ O ₂	[na] $\Delta_v H$	(Z) 15-octadecenyl acetate (393–438)	110.2	298	GC	[1997KOU/HOS, 2000OVA/KOU]
C ₂₀ H ₃₈ O ₂	[na] $\Delta_v H$	(E) 15-octadecenyl acetate (393–438)	110.5	298	GC	[1997KOU/HOS, 2000OVA/KOU]
C ₂₀ H ₃₈ O ₂	[5561-99-9] $\Delta_{\text{trs}} H$ $\Delta_{\text{fus}} H$	<i>cis</i> -11-eicosenoic acid (gondoic acid)	9.0 49.7	270 296.5		[1997SAT/YAN]
C ₂₀ H ₃₈ O ₄	[14491-66-8] $\Delta_v H$	dioctyl succinate (503–523)	94.2	513	A	[1987STE/MAL]
C ₂₀ H ₃₈ O ₄	[6819-09-6] $\Delta_v H$	dipentyl sebacate (353–408)	99.2	368	A	[1987STE/MAL]
C ₂₀ H ₃₈ O ₄	[2424-92-2] $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$	eicosanedioic acid (380–395) (336–346)	165.7 ± 3.3 170.0 ± 3.3 199.5	388 298 341	ME A	[1960DAV/THO, 1987STE/MAL] [1960DAV/THO, 1999RIB/MON] [1987STE/MAL]
C ₂₀ H ₃₈ O ₅	[1086272-76-5] $\Delta_v H$	dodecyl[1-(butoxycarbonyl)ethyl] carbonate (408–498)	82.8	423	A	[1987STE/MAL]
C ₂₀ H ₃₉ NO ₃	[14379-41-0] $\Delta_{\text{fus}} H$	N-tetradecanoyl-(<i>l</i>)-leucine	32.4	377.5	DSC	[1986MIY/MAT]
C ₂₀ H ₃₉ NO ₃	[21394-55-8] $\Delta_{\text{trs}} H$ $\Delta_{\text{fus}} H$	N-tetradecanoyl-(<i>dl</i>)-leucine	1.8 54.8	320.1 349.6	DSC	[1986MIY/MAT]
C ₂₀ H ₄₀	[3452-07-1] $\Delta_v H$ $\Delta_v H$ $\Delta_v H$	1-eicosene (478–638) (573–615)	74.3 65 100	493 588 298	A	[1999DYK/SVO] [1987STE/MAL] [1971WIL/ZWO]

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		T _m (K)	Method	Reference	
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)				
C ₂₀ H ₄₀	[1795-18-2]	tetradecylcyclohexane					
	$\Delta_v H$	(486–665)	74.7	501		[1999DYK/SVO]	
	$\Delta_v H$		99.4	298		[1971WIL/ZWO]	
C ₂₀ H ₄₀	[4669-01-6]	pentadecylcyclopentane					
	$\Delta_v H$	(486–661)	76.5	501		[1999DYK/SVO]	
	$\Delta_v H$		100.3	298		[1971WIL/ZWO]	
C ₂₀ H ₄₀	[42506-54-7]	1,1,9,9-tetramethylcyclohexadecane					
$\Delta_{\text{fus}} H$			25.1	364.2		[1975BJO/BOR2]	
C ₂₀ H ₄₀	[42506-49-0]	1,1,4,4-tetramethylcyclohexadecane					
$\Delta_{\text{fus}} H$			25.1	303.2		[1975BJO/BOR2]	
C ₂₀ H ₄₀	[54157-03-8]	1,1-dimethylcyclooctadecane					
$\Delta_{\text{fus}} H$			23.85	283.2		[1974BJO/BOR]	
C ₂₀ H ₄₀ O	[60046-87-9]	3,7,11,15-tetramethyl-1-hexadecen-3-ol					
$\Delta_v H$		(439–468)	67.0 ± 2.0	453		[1988BAG/GUR]	
C ₂₀ H ₄₀ O ₂	[822-23-1]	octadecyl acetate					
	$\Delta_v H$	(393–438)	113.5	298	GC	[1997KOU/HOS, 2000OVA/KOU]	
	$\Delta_v H$	(341–500)	94.3	356	A	[1987STE/MAL]	
C ₂₀ H ₄₀ O ₂	[111-06-8]	butyl palmitate					
$\Delta_v H$		(353–383)	93.8	368	A	[1987STE/MAL]	
C ₂₀ H ₄₀ O ₂	[1654-86-0]	decyl decanoate					
$\Delta_v H$		(341–398)	97.8	356	A	[1987STE/MAL]	
C ₂₀ H ₄₀ O ₂	[506-30-9]	eicosanoic acid					
	$\Delta_{\text{trs}} H$		4.1	332.8			
	$\Delta_{\text{trs}} H$		6.1	333.3			
	$\Delta_{\text{fus}} H$		71.6	347.8	DSC	[2007MOR/COR]	
	$\Delta_{\text{fus}} H$		69.2	348.2		[1996DOM/HEA]	
	$\Delta_{\text{sub}} H$	(305–323)	148.4		TPTD	[2005CHA/ZIE]	
	Note: Experimental values based on the TPTD method are often inconsistent with values determined using other experimental methods						
	$\Delta_{\text{sub}} H$	(337–346)	199.6 ± 7.5	342	ME	[1961DAV/MAL, 1970COX/PIL]	
	$\Delta_v H$	(477–670)	114.5	492	A	[1987STE/MAL]	
	$\Delta_v H$	(380–404)	125.5	392	ME,TE	[1982DEK/SCH]	
C ₂₀ H ₄₀ O ₂	[111-61-5]	ethyl stearate					
	$\Delta_{\text{fus}} H$		59.83	307		[1967OMA]	
	$\Delta_{\text{sub}} H$	(297–306)	161.4	301.5	ME	[1987STE/MAL, 1967OMA]	
	$\Delta_v H$	(454–469)	111.9	461	A	[1987STE/MAL]	
	$\Delta_v H$	(310–328)	106.8	319	A, ME	[1987STE/MAL, 1967OMA]	
C ₂₀ H ₄₀ O ₂	[1731-94-8]	methyl nonadecanoate					
	$\Delta_{\text{fus}} H$		63.8	313.2	DSC	[2004CHI/ZHA]	
	$\Delta_{\text{trs}} H$		19.4	304.2			
	$\Delta_{\text{fus}} H$		42.8	313.2		[1936KIN/GAR]	
	$\Delta_v H$	(467–558)	109.5 ± 5.4	298	CGC	[2004CHI/ZHA]	
	$\Delta_v H$		101.2	350		[2002VAN/VAN]	
	$\Delta_v H$		105.0 ± 2.4	326		[2002VAN/VAN]	
$\Delta_v H$		109.5 ± 2.7	298		[2002VAN/VAN]		

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	Enthalpy							
		$\Delta_v H$		(441–529)	90.1	456	A, EST	[1987STE/MAL, 1963ROS/SCH]
C ₂₀ H ₄₀ O ₂	[20292-08-4]		2-ethylhexyl laurate					
		$\Delta_v H$		(371–452)	91.4	386		[2001BUR/JOS]
		$\Delta_v H$		(443–503)	104.5	298	GC	[1997KRO/VEL]
C ₂₀ H ₄₀ O ₄	[43091-29-8]		2,2,6,6,10,10,14,14-octamethyl-1,3,9,11-tetraoxacyclohexadecane					
		$\Delta_{\text{fus}} H$			24.69	406.9		[1973DAL/EKE]
C ₂₀ H ₄₀ O ₄	[56444-63-4]		2,2,12,12-tetramethyl-1,3,11,13-tetraoxacycloeicosane					
		$\Delta_{\text{fus}} H$			45.6	369.5		[1975BOR]
C ₂₀ H ₄₁ Br	[4276-49-7]		1-bromoeicosane					
		$\Delta_v H$		(502–673)	79.8	517	A, EST	[1987STE/MAL, 1961LI/ROS, 1970DYK/VAN]
C ₂₀ H ₄₁ Cl	[42217-02-7]		1-chloroeicosane					
		$\Delta_v H$			120.2	298		[2006BOL/NER2]
		$\Delta_v H$		(492–673)	78.3	507	A	[1987STE/MAL, 1970DYK/VAN]
C ₂₀ H ₄₁ F	[676-44-8]		1-fluoroicosane					
		$\Delta_v H$		(468–663)	74.3	483	A, EST	[1987STE/MAL, 1961LI/ROS, 1970DYK/VAN]
C ₂₀ H ₄₁ I	[34994-81-5]		1-iodoeicosane					
		$\Delta_v H$		(516–673)	118.5	298	A, EST	[1987STE/MAL, 1961LI/ROS, 1970DYK/VAN, 2006BOL/NER]
		$\Delta_v H$		(516–673)	80.9	531	A, EST	[1987STE/MAL, 1961LI/ROS, 1970DYK/VAN]
C ₂₀ H ₄₁ NO	[74534-12-6]		N-hexyl tetradecanamide					
		$\Delta_{\text{trs}} H$			8.0	310		
		$\Delta_{\text{trs}} H$			7.0	328		
		$\Delta_{\text{fus}} H$			35.0	334	DSC	[1980CAR/BUS]
C ₂₀ H ₄₁ NO	[146985-21-9]		N,N-di(2-ethylhexyl) isobutyramide					
		$\Delta_v H$		(463–513)	79.1 ± 0.9	298	CGC	[2009PAN/ANT]
C ₂₀ H ₄₁ NO	[75397-93-2]		N,N-dihexyl octanamide					
		$\Delta_v H$		(463–513)	82.9 ± 1.0	298	CGC	[2009PAN/ANT]
C ₂₀ H ₄₂	[6912-07-8]		5-butylhexadecane					
		$\Delta_v H$		(423–457)	77.3	438	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₂₀ H ₄₂	[61868-04-0]		2,3-dimethyloctadecane					
		$\Delta_v H$		(458–612)	65.7	473	A	[1987STE/MAL, 1959TER/BRI]
C ₂₀ H ₄₂	[61868-10-8]		2,4-dimethyloctadecane					
		$\Delta_v H$		(456–583)	75.8	471	A	[1987STE/MAL, 1959TER/BRI]
C ₂₀ H ₄₂	[112-95-8]		eicosane					
		$\Delta_{\text{fus}} H$			69.03	311.6		
		$\Delta_{\text{fus}} H$			69.8	310.2		[2006KHI/BOU, 2006GEN/AMA]
		$\Delta_{\text{fus}} H$			68.1	309.7	DSC	[2004MON/RAJ]
		$\Delta_{\text{fus}} H$			67.8	309.7		[1996DOM/HEA]
		$\Delta_{\text{sub}} H$		(302–308)	172.8 ± 3.0	305		[2009RAZ/NAC]
		$\Delta_{\text{sub}} H$			179.5 ± 2.0	367	B	[1994PIA/FON]
		$\Delta_{\text{sub}} H$		U	152.3 ± 5.0	298	B	[1991PIA/POM]
		$\Delta_{\text{sub}} H$			170.4	298	C	[1972MOR3]
		$\Delta_v H$		(313–373)	99.5 ± 1.1	343		[2009RAZ/NAC]
		$\Delta_v H$			102.6 ± 1.0	298	CGC	[2002CHI/WEB]
		$\Delta_v H$			102.8 ± 2.2	298	GS	[2001PUR/CHI]

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound		T _m (K)	Method	Reference	
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)				
		$\Delta_{\text{v}}H$		101.1 ± 2.0	298	CGC	[2000NIC/ORF]
		$\Delta_{\text{v}}H$	(453–503)	103.5	298	CGC	[1995CHI/HOS]
		$\Delta_{\text{v}}H$	(433–583)	78.0	448		[1994MOR/KOB]
		$\Delta_{\text{v}}H$		101.8	298		[1994RUZ/MAJ]
		$\Delta_{\text{v}}H$	(347–388)	110 ± 2	368	TE	[1994PIA/FON]
		$\Delta_{\text{v}}H$	(345–470)	79.0	360	TE,ME,GS	[1991PIA/POM]
		$\Delta_{\text{v}}H$	(388–625)	80.8	440	EB,IP	[1989CHI/NGU]
		$\Delta_{\text{v}}H$	(388–625)	68.3	540	EB,IP	[1989CHI/NGU]
		$\Delta_{\text{v}}H$	(363–460)	89.6	378		[1988SAS/JOS]
		$\Delta_{\text{v}}H$	(528–620)	71.1	543	A	[1987STE/MAL]
		$\Delta_{\text{v}}H$	(344–380)	93.3	359	A, GS	[1987STE/MAL, 1979MAC/PRA]
		$\Delta_{\text{v}}H$		100.8	298		[1971WIL/ZWO]
C₂₀H₄₂	[1560-86-7]	2-methylnonadecane					
		$\Delta_{\text{v}}H$	(465–607)	72.4	480	A	[1987STE/MAL, 1959PAR/MAC]
C₂₀H₄₂	[6418-45-7]	3-methylnonadecane					
		$\Delta_{\text{v}}H$	(463–609)	71.3	478	A	[1987STE/MAL, 1959TER/BRI]
C₂₀H₄₂	[25117-27-5]	4-methylnonadecane					
		$\Delta_{\text{v}}H$	(460–609)	68.4	475	A	[1987STE/MAL, 1959TER/BRI]
C₂₀H₄₂	[57160-72-2]	5-methylnonadecane					
		$\Delta_{\text{v}}H$	(462–609)	69.1	477	A	[1987STE/MAL, 1959TER/BRI]
C₂₀H₄₂	[55044-10-5]	4-propylheptadecane					
		$\Delta_{\text{v}}H$	(425–459)	79.2	440	A, MG	[1987STE/MAL, 1955SCH/WHI]
C₂₀H₄₂	[102155-32-8]	2,4,6-trimethylheptadecane					
		$\Delta_{\text{v}}H$	(449–579)	71.9		A	[1987STE/MAL, 1999DYK/SVO]
C₂₀H₄₂O	[629-96-9]	1-eicosanol					
		$\Delta_{\text{fus}}H$		43.6	336.6	DSC	[2004VEN/CAL]
		$\Delta_{\text{trs}}H$		28.4	335.5		
		$\Delta_{\text{fus}}H$		43.6	336.6		[2002VEN/RAM]
		$\Delta_{\text{fus}}H$		73.72	338.2		[2001VAN/OON2]
		$\Delta_{\text{sub}}H$	(327–341)	218 ± 3.8	332	ME	[1965DAV/KYB, 1987STE/MAL]
		$\Delta_{\text{sub}}H$		223 ± 3.8	298		[1965DAV/KYB]
		$\Delta_{\text{v}}H$		125.9 ± 0.8	298	CGC	[2006NIC/KWE]
		$\Delta_{\text{v}}H$	(488–653)	83.5	503	A	[1987STE/MAL]
		$\Delta_{\text{v}}H$	(493–648)	83.4	508	A	[1987STE/MAL]
		$\Delta_{\text{v}}H$	(339–358)	118.9	348	ME	[1987STE/MAL, 1965DAV/KYB]
C₂₀H₄₂O₂	[7735-43-5]	1,20-eicosanediol					
		$\Delta_{\text{trs}}H$		37	368.6		
		$\Delta_{\text{fus}}H$		39.7	376.1	DSC	[1999OGA/NAK]
C₂₀H₄₂O₅	[5274-68-0]	3,6,9,12-tetraoxa-1-tetracosanol					
		$\Delta_{\text{v}}H$	(501–543)	135.5	516	A	[1987STE/MAL]
C₂₀H₄₂O₁₀	[na]	1,ω-dimethoxynona(oxyethylene)					
		$\Delta_{\text{fus}}H$		73.9	289.2		[1996YAN/YU]
C₂₀H₄₂S	[13373-97-2]	1-eicosanethiol					
		$\Delta_{\text{v}}H$	(512–694)	81.3	527	EST	[1999DYK/SVO]
C₂₀H₄₂S₂	[10496-18-1]	dinonyl disulfide					
		$\Delta_{\text{v}}H$	(518–702)	83.4	533	EST	[1999DYK/SVO]

TABLE 10. Phase change enthalpies of C₁₅ to C₂₀ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
C ₂₀ H ₄₃ N	[1120-49-6] $\Delta_{\text{v}}H$	didecylamine (506–705)	70.9	521	A	[1987STE/MAL]
C ₂₀ H ₄₃ N	[30951-88-3] $\Delta_{\text{v}}H$	N,N-diethylhexadecylamine (412–628)	71.9	427	A	[1987STE/MAL, 1947STU]
C ₂₀ H ₄₃ N	[124-28-7] $\Delta_{\text{v}}H$	N,N-dimethyloctadecylamine (504–701)	74.7	519	A	[1987STE/MAL]
C ₂₀ H ₄₃ N	[10525-37-8] $\Delta_{\text{v}}H$	eicosylamine (543–659)	74.5	558	A	[1987STE/MAL, 1956MAN2]
C ₂₀ H ₄₈ O ₂	[302-79-4] $\Delta_{\text{trs}}H$ $\Delta_{\text{fus}}H$ (I) $\Delta_{\text{fus}}H$ (II)	3,7-dimethyl-9-(2,6,6-trimethyl-1-cyclohexen-1-yl)-2,4,6,8-nonatetraenoic acid (retinoic acid)	3.2 37.1 36.8	419.8 456.9 456.3		[2006CAV/PAN] [2006CAV/PAN]

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₂₁ H ₆ N ₁₂ O ₁₈	[49753-54-0] $\Delta_{\text{sub}}H$	2,4,6- <i>tris</i> (2,4,6-trinitrophenyl)-1,3,5-triazine (479–551)	167.9	494	A	[1987STE/MAL]
C ₂₁ H ₈ F ₂₈ O ₈	[na] Δ_vH	pentaerythritol, tetra-perfluorobutyrate (293–433)	35.5	308	I, A	[1987STE/MAL, 1957DOB/KEL]
C ₂₁ H ₁₃ F ₁₃ OS	[246543-94-2] $\Delta_{\text{fus}}H$ $\Delta_{\text{fus}}H$	2-(perfluoro- <i>n</i> -hexyl)ethylthiomethyl biphenyl-4-yl ether	40.6 40.4	344.8 344.8	DTA	[1999DEG/GUI] [1999TAF/GUI, 1999DEG/GUI]
C ₂₁ H ₁₃ F ₁₃ S	[na] $\Delta_{\text{fus}}H$	2-(perfluoro- <i>n</i> -hexyl)ethylthiomethyl biphenyl-4-yl	53.1	332.9	DTA	[1999TAF/GUI, 1999DEG/GUI]
C ₂₁ H ₁₃ N	[215-62-3] $\Delta_{\text{fus}}H$	dibenz[a,c]acridine	27.8	477.4	DSC	[2010KES/AUC]
C ₂₁ H ₁₃ N	[226-36-8] $\Delta_{\text{fus}}H$	dibenz[a,h]acridine	30.6	499.7	DSC	[2010KES/AUC]
C ₂₁ H ₁₃ N	[226-92-6] $\Delta_{\text{fus}}H$	dibenz[a,i]acridine	29.6	483.5	DSC	[2010KES/AUC]
C ₂₁ H ₁₃ N	[224-42-0] $\Delta_{\text{fus}}H$	dibenz[a,j]acridine	25.5	492.7	DSC	[2010KES/AUC]
C ₂₁ H ₁₄ N ₂ O ₃	[13494-38-7] $\Delta_{\text{sub}}H$	2-phenyl-3-benzoylquinoxaline-1,4-dioxide	167.4 ± 4.0	298	ME	[1997ACR/POW]
C ₂₁ H ₁₄ N ₂ O ₃	[na]	1,4-diamino-2-benzoyl-9,10-anthraquinone	168.5			[1984KAR/KRU]
C ₂₁ H ₁₅ BrN ₂ O ₂	[128-83-6] $\Delta_{\text{sub}}H$	1-amino-2-bromo-4-[(4-methylphenyl)amino]-9,10-anthraquinone (418–438)	167.0 ± 6.0	428		[1984KRI]
C ₂₁ H ₁₅ NO ₂	[158749-37-2] $\Delta_{\text{fus}}H$	2-methylphenyl acridine-9-carboxylate	30.5	415	DSC	[2010KRZ/MAL]
C ₂₁ H ₁₅ NO ₂	[158749-58-7] $\Delta_{\text{fus}}H$	3-methylphenyl acridine-9-carboxylate	32	429	DSC	[2010KRZ/MAL]
C ₂₁ H ₁₅ NO ₂	[158749-59-8] $\Delta_{\text{fus}}H$	4-methylphenyl acridine-9-carboxylate	30.7	446	DSC	[2010KRZ/MAL]
C ₂₁ H ₁₅ NO ₃	[na] $\Delta_{\text{sub}}H$	2-hydroxy-4-[(4-methylphenyl)amino]-9,10-anthraquinone (349–378)	121.0 ± 7.6	363		[1984KRI] Note: Compound is listed as the 2-hydroxy-derivative in the paper; however, it is listed as the 1-hydroxy-derivative in Chem. Abstracts
C ₂₁ H ₁₆	[56-49-5] $\Delta_{\text{sub}}H$ Δ_vH	3-methylcholanthrene (401–425) (323–473)	127.2 ± 2.4 93.8	413 398	A GC	[1987STE/MAL, 1964KEL/RIC] [2002LEI/CHA]
C ₂₁ H ₁₆	[611-48-3] $\Delta_{\text{fus}}H$	1,2'-dinaphthylmethane	30.54	369.6		[1996DOM/HEA]
C ₂₁ H ₁₆ N ₂	[484-47-9] $\Delta_{\text{fus}}H$ $\Delta_{\text{us}}H$ $\Delta_{\text{fus}}H$	2,4,5-triphenylimidazole	37.31 0.73 35.15	547.8 505.7 550.8	DSC	[2007SIF/AIT] [2002ROG/DOM]

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₂₁ H ₁₆ N ₂ O ₂	[na]	1-anilino-4-(N-methylamino)-9,10-anthraquinone 136.9 [1984KAR/KRU]				
C ₂₁ H ₁₆ N ₄ O ₂	[194784-98-0] $\Delta_{\text{fus}}H$	1-(4'-methoxybenzylidene)-2-phenazinoylhydrazine 63.26 534.3 DSC [1997CIO/MEL]				
C ₂₁ H ₁₇ F ₃ O	[145698-51-7] $\Delta_{\text{fus}}H$	4- <i>n</i> -butoxy-4'-trifluoromethyldiphenyldiacetylene 25.37 414.3 DSC [1993JUA/CHE]				
C ₂₁ H ₁₇ N ₃ O ₃	[82232-20-0] $\Delta_{\text{sub}}H$	(5-cyano-3,4-diphenyl-6-oxo-1,6-dihydropyridazin-1-yl)acetate (396–414) 131.9 ± 9.3 405 ME [1982DEP]				
C ₂₁ H ₁₈ F ₂	[193472-73-0] $\Delta_{\text{fus}}H$	1,1-difluoro-3,3,3-triphenylpropane 28.74 370.2 [1997SCH/VER]				
	$\Delta_{\text{sub}}H$	113.2 ± 1.7 298 [1997SCH/VER]				
C ₂₁ H ₁₈ F ₂	[145698-37-9] $\Delta_{\text{fus}}H$	4- <i>n</i> -pentyl-3',4'-difluorodiphenyldiacetylene 30.86 355.1 DSC [1993JUA/CHE]				
C ₂₁ H ₁₉ F	[193472-69-4] $\Delta_{\text{fus}}H$	1-fluoro-3,3,3-triphenylpropane 26.44 344.2 [1997SCH/VER]				
	$\Delta_{\text{sub}}H$	129.3 ± 0.6 298 [1997SCH/VER]				
	Δ_vH	(349–384) 95.9 ± 0.6 298 GS [1997SCH/VER]				
C ₂₁ H ₁₉ F	[193472-72-9] $\Delta_{\text{fus}}H$	2-fluoro-1,2,3-triphenylpropane 34.6 379.6 [1997SCH/VER]				
	$\Delta_{\text{sub}}H$	132.5 ± 3.0 298 [1997SCH/VER]				
C ₂₁ H ₂₀ BrN ₇ O ₆	[na] $\Delta_{\text{fus}}H$	N-[2-[(2-bromo-4,6-dinitrophenyl)azo]-[(2-cyanoethyl)-2-propenylamino]-4-methoxyphenyl] acetamide 59.08 465.2 [1991BAU/WEB]				
C ₂₁ H ₂₀ Br ₈ O ₂	[na] $\Delta_{\text{fus}}H$	2,2- <i>bis</i> [3,5-dibromo-4-(2,3-dibromopropoxy)phenyl]propane 39.63 Not given [1999TAH/TAK]				
		Note: Solid sample was precipitated from a methanol-dichloromethane mixture. Abstract implies that the compound may have other crystal forms.				
C ₂₁ H ₂₀ Cl ₂ O ₃	[61949-76-6] $\Delta_{\text{sub}}H$	(3-phenoxyphenyl)methyl- <i>cis</i> -3-(2,2-dichloroethenyl)-2,2-dimethylcyclopropanecarboxylate (<i>cis</i> -permethrin) (313–333) 108.8 323 GS,A [1986WEL/GRA]				
C ₂₁ H ₂₀ N ₄ O ₃	[32828-81-2] $\Delta_{\text{fus}}H$	4-methoxy-N,N- <i>bis</i> (3-pyridinylmethyl)-1,3-benzenedicarboxamide 28.43 403.9 DSC [1998MUR/BET]				
C ₂₁ H ₂₀ O ₁₂	[21637-25-2] $\Delta_{\text{fus}}H$	2-(3,4-dihydroxyphenyl)-3-(β -D-glucofuranosyloxy)-5,7-dihydroxy-4 <i>H</i> -1-benzopyran-4-one (isoquercitrin) 49.8 471.2 DSC [2007CHE/HUM]				
C ₂₁ H ₂₁ N	[1159-53-1] $\Delta_{\text{fus}}H$	4-methyl-N,N- <i>bis</i> (4-methylphenyl)benzenamine 22.96 388				
	$\Delta_{\text{fus}}H$	19.95 388.8 DSC [2004MAN/ROH, 2006MAN/ROH]				
	$\Delta_{\text{sub}}H$	92.53 DSC [2006MAN/ROH]				
	Δ_vH	72.57 DSC [2006MAN/ROH]				
C ₂₁ H ₂₁ N	[117597-62-3] $\Delta_{\text{fus}}H$	N-(3-methylphenyl)-N,N- <i>bis</i> (4-methylphenyl)amine 21.71 329.9 DSC [2006MAN/ROH]				

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₂₁ H ₂₁ N	$\Delta_{\text{sub}}H$		97.64		DSC	[2006MAN/ROH]
	Δ_vH		75.93		DSC	[2006MAN/ROH]
	[97413-60-0]	N,N-bis(3-methylphenyl)-N-(4-methylphenyl)amine				
	$\Delta_{\text{fus}}H$		26.39	362.7	DSC	[2006MAN/ROH]
	$\Delta_{\text{sub}}H$		95.69		DSC	[2006MAN/ROH]
C ₂₁ H ₂₁ N	Δ_vH		69.29		DSC	[2006MAN/ROH]
	[20676-79-3]	N,N,N-tris(3-methylphenyl)amine				
	$\Delta_{\text{fus}}H$		13.07	313	DSC	[2006MAN/ROH]
	$\Delta_{\text{sub}}H$		50.66		DSC	[2006MAN/ROH]
C ₂₁ H ₂₁ NO	Δ_vH		37.59		DSC	[2006MAN/ROH]
	[957-51-7]	N,N-dimethyl-2,2-diphenylbenzeneacetamide				
	$\Delta_{\text{fus}}H$		25.43	402	DSC	[1990DON/DRE]
C ₂₁ H ₂₁ O ₄ P	[78-30-8]	phosphoric acid, tri(2-tolyl) ester				
	Δ_vH	(293–700)	86.8	308	A, I	[1987STE/MAL, 1957DOB/KEL]
C ₂₁ H ₂₁ O ₄ P	[563-04-2]	phosphoric acid, tri(3-tolyl) ester				
	Δ_vH	(398–530)	123.2	413	A	[1987STE/MAL]
C ₂₁ H ₂₁ O ₄ P	[78-32-0]	phosphoric acid, tri(4-tolyl) ester				
	Δ_vH	(388–530)	104.9	408	A	[1987STE/MAL]
C ₂₁ H ₂₁ P	[1038-95-5]	tris(4-tolyl)phosphine				
	Δ_vH	(372–394)	126 ± 5	385	ME, TE	[1981DEK/HER]
C ₂₁ H ₂₃ BrFNO ₂	[10457-90-6]	1-(4-fluorophenyl)-4-[4-hydroxy-4-(4-bromophenyl)-1-piperidiny]-1-butanone (bromoperidol)				
	$\Delta_{\text{fus}}H$		50.8	432.7	DSC	[1981DRA/AZI]
C ₂₁ H ₂₃ ClFNO ₂	[52-86-8]	1-(4-fluorophenyl)-4-[4-hydroxy-4-(4-chlorophenyl)-1-piperidiny]-1-butanone (haloperidol)				
	$\Delta_{\text{fus}}H$		48	422.7	DSC	[1981DRA/AZI]
C ₂₁ H ₂₃ F ₂ NO ₂	[na]	1-(4-fluorophenyl)-4-[4-hydroxy-4-(4-fluorophenyl)-1-piperidiny]-1-butanone				
	$\Delta_{\text{fus}}H$		34	395.2	DSC	[1981DRA/AZI]
C ₂₁ H ₂₃ NO ₅	[561-27-3]	diacetylmorphine (heroin)				
	$\Delta_{\text{sub}}H$	(324–339)	144.5 ± 4.0	331	GS	[1984LWA/EL]
C ₂₁ H ₂₄ FNO ₂	[3109-12-4]	1-(4-fluorophenyl)-4-[4-hydroxy-4-phenyl-1-piperidiny]-1-butanone				
	$\Delta_{\text{fus}}H$ (I)		43.2	412.7		
	$\Delta_{\text{fus}}H$ (II)		35.2	385.2	DSC	[1981DRA/AZI]
C ₂₁ H ₂₄ O ₂	[160731-84-0]	3-(diphenylmethyl)-3-propyl-2,4-pentanedione				
	$\Delta_{\text{fus}}H$		27.1	349.2		[1995NOL/VER]
	$\Delta_{\text{sub}}H$		124.7	298	T,B	[1995NOL/VER]
C ₂₁ H ₂₅ FN ₂ O ₂	Δ_vH	(364–392)	96.7 ± 1.7	378	GS	[1995NOL/VER]
	[1480-19-9]	1-(4-fluorophenyl)-4-[4-(2-methoxyphenyl)-1-piperaziny]-1-butanone (fluanisone)				
	$\Delta_{\text{fus}}H$ (I)		27.3	348.7		
	$\Delta_{\text{fus}}H$ (II)		31.1	343.7		
C ₂₁ H ₂₅ F ₁₉	$\Delta_{\text{fus}}H$ (III)		15.7	323.7	DSC	[1981DRA/AZI]
	[139277-02-4]	1,1,1,2,3,3,4,4,5,5,6,6,7,7,8,8-hexadecafluoro-(trifluoromethyl)eicosane				
	$\Delta_{\text{fus}}H$		34	310.1	DSC	[1992HOP/MOL]

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₂₁ H ₂₅ NO	[122405-21-4]	4-(1-methylheptyloxy)-4'-cyanobiphenyl				
	$\Delta_{\text{fus}}H$ (I)		19.51	287.6		
	$\Delta_{\text{fus}}H$ (II)		17.07	294.3	DSC	[2004SAI/MAS]
	$\Delta_{\text{fus}}H$ (I)		20.8	287.8		
	$\Delta_{\text{fus}}H$ (II)		19.1	294.3		[2001SCI/SCI]
C ₂₁ H ₂₆	[6169-94-4]	[1,8]-para-cyclophane				
	$\Delta_{\text{sub}}H$	(354–376)	105 ± 1.3	365		[1969SHI/MCN, 1977PED/RYL]
	$\Delta_{\text{sub}}H$		110.9 ± 2.1	298		[1969SHI/MCN, 1977PED/RYL]
C ₂₁ H ₂₆ ClN ₃ OS	[58-39-9]	2-chloro-10-3-[1-(2-hydroxyethyl)-4-piperazinyl]propylphenothiazine (perphenazine)				
$\Delta_{\text{fus}}H$			41.8	370	DSC	[2006WAS/HOL]
C ₂₁ H ₂₆ FNO	[na]	4-octyloxy-N-(4-fluorobenzylidene) aniline				
	$\Delta_{\text{fus}}H$		44.7	360.5		[1991MIY/ENO]
C ₂₁ H ₂₆ FN ₃ O ₄	[143383-65-7]	[S-(R*,S*)]-1-cyclopropyl-6-fluoro-1,4-dihydro-8-methoxy-7-[3-[1-(methylamino)ethyl]-1-pyrrolidinyl]-4-oxo-3-quinolinecarboxylic acid (premafoxacin)				
	$\Delta_{\text{fus}}H$		60.52	471.9	DSC	[1997SCH/BER]
C ₂₁ H ₂₆ O ₂	[72-33-3]	3-methoxy-19-norpregna-1,3,5(10)-trien-20-yn-17-ol (mestranol)				
	$\Delta_{\text{fus}}H$		34.55	424.1		[1985DEM/CHA]
C ₂₁ H ₂₆ O ₂	[521-35-7]	cannabinol				
	$\Delta_{\text{fus}}H$		17	352.2		[2004STI/VAL]
C ₂₁ H ₂₆ O ₃	[2549-90-8]	2-hydroxy-4-(2-ethylhexyloxy)benzophenone				
	Δ_vH	(393–443)	98.7	418	ME	[1984SUR]
C ₂₁ H ₂₆ O ₃	[1843-05-6]	2-hydroxy-4-octyloxybenzophenone				
	Δ_vH	(413–453)	102.1	433	ME	[1984SUR]
C ₂₁ H ₂₆ O ₃	[68100-20-9]	2-hydroxy-4-butoxy-5- <i>tert</i> -butylbenzophenone				
	Δ_vH	(403–453)	90.2	428	ME	[1984SUR]
C ₂₁ H ₂₆ O ₃	[975-64-4]	3-(1-oxypropoxy)-estra-1,3,5(10)-trien-17-one				
	$\Delta_{\text{fus}}H$		23.0	409	DSC	[1990YAN/EIR]
C ₂₁ H ₂₆ O ₄	[6127-74-8]	2-hydroxy-4,4'-dibutoxybenzophenone				
	$\Delta_{\text{fus}}H$		54.0	372.1	DSC	[1999PRI/HAWN]
	$\Delta_{\text{sub}}H$		148		B	[1999PRI/HAWN]
C ₂₁ H ₂₆ O ₄	[96609-16-4]	4-{4-[4-(1,1-dimethylethyl)phenyl]-2-hydroxybutoxy}benzoic acid (lifibrol)				
	$\Delta_{\text{fus}}H$ (I)		38.1	415.2		
	$\Delta_{\text{fus}}H$ (II)		49.1	408.2		[2000BUR/LET]
C ₂₁ H ₂₇ FO ₆	[124-94-7]	triamcinolone				
	$\Delta_{\text{fus}}H$		42.56	543		[1994REG/CHM]
C ₂₁ H ₂₇ NO	[na]	4-octyloxy-N-benzylidene aniline				
	$\Delta_{\text{fus}}H$		41.47	342.7		[1991MIY/ENO]
C ₂₁ H ₂₈ O ₂	[5630-53-5]	(1 α ,17 α)-17-hydroxy-7-methyl-19-norpregn-5(10)-en-20-yn-3-one (tibolone)				
	$\Delta_{\text{fus}}H$ (monoclinic)		29.8	444.2	DSC	[2010BAR/ARA]
	$\Delta_{\text{fus}}H$ (triclinic)		21.0	421.2	DSC	[2010BAR/ARA]
	$\Delta_{\text{fus}}H$ (triclinic)		3.0	439.2	DSC	[2010BAR/ARA]
C ₂₁ H ₂₈ O ₃	[1097-51-4]	16 α ,17 α -epoxyprogesterone				
	$\Delta_{\text{fus}}H$		26.4	328.5		[2006NIE/GON]
C ₂₁ H ₂₈ O ₄	[19427-36-2]	11 α -hydroxy-16 α ,17 α -epoxyprogesterone				

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
	$\Delta_{\text{fus}}H$		44.8	522.2		[2006NIE/GON]
C ₂₁ H ₂₈ O ₅	[50-24-8]	prednisolone				
	$\Delta_{\text{fus}}H$		38.86	513		[1994REG/CHM]
C ₂₁ H ₂₈ O ₅	[53-06-5]	cortisone				
	$\Delta_{\text{fus}}H$		36.86	495		[1994REG/CHM]
C ₂₁ H ₂₉ NO ₃	[172589-24-1]	3-[(hydroxyimino)(5,6,7,8-tetrahydro-2-naphthalenyl)methyl-1,2,2-trimethylcyclopentanecarboxylic acid methyl ester				
	$\Delta_{\text{fus}}H$		38.37	425		[1995NUR/LEL]
C ₂₁ H ₂₉ N ₃ O	[3737-09-5]	α -[2-[bis(1-methylethyl)amino]ethyl]- α -phenyl-2-pyridineacetamide (disopyramide)				
	$\Delta_{\text{fus}}H$		26.7	363.7	DSC	[2008WAS/HOL]
C ₂₁ H ₃₀	[7225-71-0]	1-undecylnaphthalene				
	$\Delta_{\text{v}}H$	(436–502)	84.3	451	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₂₁ H ₃₀ O	[38256-01-8]	1,1'-diadamantyl ketone				
	$\Delta_{\text{rs}}H$		5.9	404.7		
	$\Delta_{\text{fus}}H$		15.7	470		[1997GAR/RED]
	$\Delta_{\text{sub}}H$	(362–378.8)	109.0 ± 1.8	298	ME	[1992ABB/JIM2]
C ₂₁ H ₃₀ O ₂	[57-83-0]	progesterone				
	$\Delta_{\text{fus}}H$ (I)		26.71	402.2	DSC	[2009BAR/ESP]
	$\Delta_{\text{fus}}H$ (II)		24.78	394.8	DSC	[2009BAR/ESP]
	$\Delta_{\text{fus}}H$ (I)		26.16	402.4	DSC	[2003LEG/FEU, 2004DEF/RAN]
	$\Delta_{\text{fus}}H$ (II)		21.42	395.4	DSC	[2003LEG/FEU, 2004DEF/RAN]
	$\Delta_{\text{fus}}H$ (III)		16.13	377	DSC	[2003LEG/FEU, 2004DEF/RAN]
	$\Delta_{\text{fus}}H$		26.99	404		[1994REG/CHM]
	$\Delta_{\text{fus}}H$ (I)		27.95	401		[1979MUR/IWA, 2009BAR/ESP]
	$\Delta_{\text{fus}}H$ (II)		23.43	395		[1979MUR/IWA, 2009BAR/ESP]
	$\Delta_{\text{fus}}H$ (I)		24.43	403.5		[1973CAM/GAM, 2009BAR/ESP]
	$\Delta_{\text{fus}}H$ (II)		21.32	396.2		[1973CAM/GAM, 2009BAR/ESP]
C ₂₁ H ₃₀ O ₂	[13956-29-1]	cannabidiol				
	$\Delta_{\text{fus}}H$		28.4	340.7		[2004STI/VAL]
C ₂₁ H ₃₀ O ₃	[64-85-7]	deoxycorticosterone				
	$\Delta_{\text{fus}}H$		27.98	414		[1994REG/CHM]
C ₂₁ H ₃₀ O ₃	[1045-69-8]	testosterone acetate				
	$\Delta_{\text{fus}}H$		27.88	413		[1994REG/CHM]
C ₂₁ H ₃₀ O ₄	[50-22-6]	corticosterone				
	$\Delta_{\text{fus}}H$		35.3	458.5	DSC	[2008WAS/HOL]
	$\Delta_{\text{fus}}H$		33.32	454		[1994REG/CHM]
C ₂₁ H ₃₀ O ₅	[50-23-7]	hydrocortisone				
	$\Delta_{\text{fus}}H$ (I)		44.65	497.7	DSC	
	$\Delta_{\text{fus}}H$ (II)		41.3	494.8	DSC	[2008SUI/JES]
	$\Delta_{\text{fus}}H$		35.84	486		[1994REG/CHM]
C ₂₁ H ₃₁ NO	[na]	6-dodecyloxyisoquinoline				
	$\Delta_{\text{fus}}H$		38.93	321.3		[1999LIN/KO]
C ₂₁ H ₃₂ O ₂	[2734-47-6]	methyl Z,Z,Z,Z,Z 5,8,11,14,17-eicosapentaenoate				
	$\Delta_{\text{v}}H$		121.0 ± 0.3	298	CGC	[2007LIP/KAP]
C ₂₁ H ₃₄ O ₂	[2566-89-4]	methyl Z,Z,Z,Z 5,8,11,14-eicosatetraenoate				

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$		118.3	298	CGC	[2007LIP/KAP]
C ₂₁ H ₃₅ N ₃ O ₂	[135742-56-2] $\Delta_{\text{fus}} H$	N-palmitoyl-pyrazinamide	51.82	362.7		[1991LIU/GUO]
C ₂₁ H ₃₆	[2131-18-2] $\Delta_v H$ $\Delta_v H$	pentadecylbenzene (495–677)	77.0 104.6	510 298		[1999DYK/SVO] [1971WIL/ZWO]
C ₂₁ H ₃₆ N ₂ OS	[442514-39-8] $\Delta_{\text{fus}} H$	N-[(3-methoxyphenyl)methyl]-N'-dodecylthiourea	52.87	361.7	DSC	[2002ABB/WHO]
C ₂₁ H ₃₆ O	[501-24-6] $\Delta_{\text{fus}} H$	3-pentadecylphenol	38.09	322.4	DSC	[2010MAO/LUO]
C ₂₁ H ₃₆ O ₂	[2566-89-4] $\Delta_v H$	methyl Z,Z,Z 11,14,17-eicosadienoate	122.6 ± 1.6	298	CGC	[2007LIP/KAP]
C ₂₁ H ₃₆ O ₆	[na] $\Delta_v H$	triisopentyl <i>trans</i> aconitate (396–499)	88.3	411	A	[1987STE/MAL]
C ₂₁ H ₃₆ O ₆	[64617-29-4] $\Delta_v H$	tripentyl <i>trans</i> aconitate (403–505)	91.4	418	A	[1987STE/MAL]
C ₂₁ H ₃₈ O ₂	[61012-46-2] $\Delta_v H$	methyl Z,Z 11,14-eicosadienoate	117.5 ± 0.5	298	CGC	[2007LIP/KAP]
C ₂₁ H ₃₈ O ₆	[621-70-5] $\Delta_v H$ $\Delta_v H$ $\Delta_v H$	glycerol tricaproate	99.9 108.3 ± 3.8 94.2	349 298 371	TGA TGA A, T	[1990KIS/SHO] [1990KIS/SHO] [1987STE/MAL, 1949PER/WEB2]
C ₂₁ H ₃₈ O ₆	[na] $\Delta_v H$	triisopentyl 1,2,3-propanetricarboxylate (396–508)	88.2	411	A	[1987STE/MAL]
C ₂₁ H ₃₈ O ₆	[5333-53-9] $\Delta_v H$	tripentyl 1,2,3-propanetricarboxylate (404–508)	90.2	419	A	[1987STE/MAL]
C ₂₁ H ₄₀	[66326-27-0] $\Delta_v H$	1-undecyldecahydronaphthalene (426–488)	83.3	411	A	[1987STE/MAL]
C ₂₁ H ₄₀	[95115-75-6] $\Delta_{\text{fus}} H$	<i>trans</i> -2-heptyl-6-butyldecalin	31.8	295.3		[1985VAR/BRI]
C ₂₁ H ₄₀	[95115-78-9] $\Delta_{\text{fus}} H$	<i>trans</i> -2-propyl-6-octyldecalin	41	308.8		[1985VAR/BRI]
C ₂₁ H ₄₀ O ₂	[2390-09-2] $\Delta_v H$	methyl Z 11-eicosenoate	115.8 ± 0.7	298	CGC	[2007LIP/KAP]
C ₂₁ H ₄₁ NO ₃	[45287-42-1] $\Delta_{\text{ms}} H$ $\Delta_{\text{fus}} H$	N-hexadecanoyl-(<i>l</i>)-valine	29.1 54.8	349.1 366.6		DSC [1986MIY/MAT]
C ₂₁ H ₄₁ NO ₃	[83871-20-9] $\Delta_{\text{fus}} H$	N-hexadecanoyl-(<i>dl</i>)-valine	80.5	375.1	DSC	[1986MIY/MAT]
C ₂₁ H ₄₂	[1599-68-4] $\Delta_v H$	1-heneicosene (392–628)	92.8	407		[1999DYK/SVO]
C ₂₁ H ₄₂	[6812-39-1] $\Delta_v H$	hexadecylcyclopentane (498–674)	79.2	513		[1999DYK/SVO]

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound					
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference	
		$\Delta_v H$	105.3	298		[1971WIL/ZWO]	
C ₂₁ H ₄₂	[6006-95-7]	pentadecylcyclohexane					
		$\Delta_{\text{fus}} H$	58.7	298.2	DSC	[2001YOU/SCH]	
		$\Delta_{\text{fus}} H$	58.3	NA	DSC	[2000YOU/DOL]	
		$\Delta_{\text{fus}} H$	58.3	301		[1991MIY/ENO]	
		$\Delta_v H$	(496–677)	77.2	511		[1999DYK/SVO]
		$\Delta_v H$	104.4	298		[1971WIL/ZWO]	
C ₂₁ H ₄₂ N ₃ PS ₆	[100575-31-3]	tris(dipropyldithiocarbamate)phosphorous					
		$\Delta_{\text{sub}} H$	127.4 ± 4.2		DSC,E	[1999NEV/GON]	
C ₂₁ H ₄₂ O	[22589-04-4]	2-heneicosanone					
		$\Delta_{\text{fus}} H$	77.65	333.9	DSC	[1993VIL/HAM]	
C ₂₁ H ₄₂ O	[19781-72-7]	11-heneicosanone					
		$\Delta_{\text{fus}} H$	76.2	336.7		[1993RUE/SAR]	
C ₂₁ H ₄₂ O ₂	[1120-28-1]	methyl eicosanoate					
		$\Delta_{\text{fus}} H$	74.3	319.2	DSC	[2004CHI/ZHA]	
		$\Delta_{\text{fus}} H$	73.7	319.2		[1936KIN/GAR]	
		$\Delta_{\text{sub}} H$	(311–318)	190.8 ± 10	314	ME	[1965DAV/KYB, 1987STE/MAL]
		$\Delta_v H$	(467–558)	120.9 ± 2.5	298	CGC	[2004CHI/ZHA]
		$\Delta_v H$		109.2	350		[2002VAN/VAN]
		$\Delta_v H$		97.8 ± 0.2	406		[2002VAN/VAN]
		$\Delta_v H$		116.4 ± 1.5	298		[2002VAN/VAN]
		$\Delta_v H$	(463–523)	116.2	298	GC	[1997KRO/VEL]
		$\Delta_v H$	(453–543)	76.9	498	GC	[1993HUS/SAR]
		$\Delta_v H$	(450–540)	92.4	465	A, EST	[1987STE/MAL, 1963ROS/SCH]
C ₂₁ H ₄₂ O ₂	[112-10-7]	isopropyl stearate					
		$\Delta_v H$	(453–483)	76.6	468	A	[1987STE/MAL]
C ₂₁ H ₄₂ O ₂	[18281-04-4]	ethyl nonadecanoate					
		$\Delta_{\text{fus}} H$	43.1	309		[1967OMA]	
		$\Delta_{\text{sub}} H$	(302–308)	149.7	305	ME	[1987STE/MAL, 1967OMA]
		$\Delta_v H$	(312–328)	111	320	A, ME	[1987STE/MAL, 1967OMA]
C ₂₁ H ₄₂ O ₂	[3634-92-2]	propyl stearate					
		$\Delta_v H$	(458–483)	87.9	470	A	[1987STE/MAL]
C ₂₁ H ₄₂ O ₂	[2363-71-5]	heneicosanoic acid					
		$\Delta_{\text{us}} H$	5.0	344.6			
		$\Delta_{\text{fus}} H$	63.0	346.7	DSC	[2007GBA/NEG]	
C ₂₁ H ₄₃ NO	[129392-93-4]	N-propylstearamide					
		$\Delta_{\text{us}} H$	16.02	348			
		$\Delta_{\text{fus}} H$	50.04	354	DSC	[1995CYP/JOH]	
C ₂₁ H ₄₃ NO	[173029-00-0]	N-heptylmyristamide					
		$\Delta_{\text{us}} H$	6.54	316			
		$\Delta_{\text{fus}} H$	49.02	343	DSC	[1995CYP/JOH]	
C ₂₁ H ₄₃ NO	[153929-66-9]	N-decylundecanamide					
		$\Delta_{\text{us}} H$	0.07	337			
		$\Delta_{\text{fus}} H$	42.45	344	DSC	[1995CYP/JOH]	

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₂₁ H ₄₃ NO	[173029-01-1]	N-laurylnonanamide				
	$\Delta_{\text{us}}H$		0.17	328		
	$\Delta_{\text{fus}}H$		66.91	341	DSC	[1995CYP/JOH]
C ₂₁ H ₄₃ NO	[173029-02-2]	N-myristylheptanamide				
	$\Delta_{\text{us}}H$		2.08	313		
	$\Delta_{\text{fus}}H$		52.68	334	DSC	[1995CYP/JOH]
C ₂₁ H ₄₃ NO	[79762-59-7]	N-stearylpropanamide				
	$\Delta_{\text{us}}H$		1.84	337		
	$\Delta_{\text{fus}}H$		56.03	350	DSC	[1995CYP/JOH]
C ₂₁ H ₄₃ NO ₂	[6280-27-9]	N-octadecyl lactamide (434–542)	112.8	449	A	[1987STE/MAL]
C ₂₁ H ₄₄	[629-94-7]	heneicosane				
	$\Delta_{\text{us}}H$		15.7	304.3		
	$\Delta_{\text{fus}}H$		46.6	313	DSC	[2004MON/RAJ]
	$\Delta_{\text{us}}H$		15.48	305.7		
	$\Delta_{\text{fus}}H$		47.7	313.7		[1996DOM/HEA]
	$\Delta_{\text{sub}}H$		141.8 ± 10	298	B	[1991PIA/POM]
	Δ_vH	(351–462)	93.7	368		[2006SAW/MOK]
	Δ_vH	(434–539)	106.8	298	CGC	[2004CHI/HAN]
	Δ_vH		109.4 ± 2.6	298	CGC	[1997CHI/WIL]
	Δ_vH	(365–400)	110 ± 2	382	TE	[1994PIA/FON]
	Δ_vH	(352–478)	84.7	367	TE,ME,GS	[1991PIA/POM]
Δ_vH	(422–630)	88.4	437	A, EST	[1987STE/MAL, 1966KUD/ZWO]	
C ₂₁ H ₄₄	[1560-84-5]	2-methyleicosane (473–621)	70.3	488	A	[1987STE/MAL]
C ₂₁ H ₄₄	[6418-46-8]	3-methyleicosane (477–620)	74.5	492	A	[1987STE/MAL, 1999DYK/SVO, 1959TER/BRI]
C ₂₁ H ₄₄	[25117-28-6]	4-methyleicosane (471–621)	70.2	486	A	[1987STE/MAL, 1999DYK/SVO, 1959TER/BRI]
C ₂₁ H ₄₄	[25117-36-6]	5-methyleicosane (519–621)	73.2	534	A	[1987STE/MAL, 1999DYK/SVO, 1959TER/BRI]
C ₂₁ H ₄₄	[75163-99-4]	2,3-dimethylnonadecane (493–635)	68.8	508	A	[1987STE/MAL, 1999DYK/SVO, 1959TER/BRI]
C ₂₁ H ₄₄	[115209-60-4]	2,4-dimethylnonadecane (465–594)	77.0	480	A	[1987STE/MAL, 1999DYK/SVO, 1959TER/BRI]
C ₂₁ H ₄₄	[11400-79-2]	2,4,6-trimethyloctadecane (460–576)	74.9	475	A	[1987STE/MAL, 1999DYK/SVO]
C ₂₁ H ₄₄	[13475-75-7]	8-hexylpentadecane (405–466)	78.5	420	A	[1987STE/MAL]
C ₂₁ H ₄₄ O ₂	[95008-70-1]	1,21-heneicosanediol	38.8	360		

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₂₁ H ₄₅ PO	$\Delta_{\text{fus}}H$		41.7	377.5	DSC	[1999OGA/NAK]
	[17262-51-0] Δ_vH	triheptylphosphine oxide (507–638)	70.8	573		[1971NAK/SMI]
		Note: The published abstract gives the formula for triheptylphosphine oxide; However, the title of the paper gives the name of trioctylphosphine oxide				
C ₂₂ H ₁₀ O ₂	[641-13-4] $\Delta_{\text{sub}}H$	anthanthrone (dibenzochrysenes-6,12-dione) (450–550)	152.2	465	A	[1987STE/MAL]
	C ₂₂ H ₁₂	[191-24-2] $\Delta_{\text{fus}}H$	benzo[ghi]perylene	17.37	554.2	
$\Delta_{\text{sub}}H$		(313–453)	129.9	383	GS	[1995NAS/LEN]
$\Delta_{\text{sub}}H$		(389–468)	127.8	404	ME	[1987STE/MAL, 1974MUR/POL]
$\Delta_{\text{sub}}H$		(450–510)	135.1	465	A	[1987STE/MAL]
$\Delta_{\text{sub}}H$		(454–502)	125.5	478	ME	[1967WAK/INO]
Δ_vH			128.9 ± 1.5	298	CGC	[2008HAN/NUT]
Δ_vH		(323–473)	96.1	398	GC	[2002LEI/CHA]
C ₂₂ H ₁₂	[191-26-4] $\Delta_{\text{sub}}H$	anthranthrene (dibenzo[def,mno]chrysenes)	135 ± 5	479	ME	[1952INO/SHI]
	C ₂₂ H ₁₂	[193-39-5] $\Delta_{\text{fus}}H$	<i>o</i> -phenylenepyrene	21.51	435.2	
C ₂₂ H ₁₂		[193-43-1] $\Delta_{\text{fus}}H$	indeno[1,2,3cd]fluoranthene	23.2	542.3	DSC
	C ₂₂ H ₁₂	[193-39-5] $\Delta_{\text{fus}}H$	indeno[1,2,3cd]pyrene	18.6	437	DSC
C ₂₂ H ₁₂ O ₂		[3029-32-1] $\Delta_{\text{sub}}H$	6,13-pentacenequinone	116.3 ± 5.9	298	
	C ₂₂ H ₁₄	[215-58-7] $\Delta_{\text{fus}}H$	dibenz[a,c]anthracene	25.82	553.5	
$\Delta_{\text{sub}}H$		(313–453)	135	383	GS	[1995NAS/LEN]
$\Delta_{\text{sub}}H$		(425–452)	159 ± 6	298	TE,ME	[1980DEK]
Δ_vH			132.3 ± 1.8	298	CGC	[2008HAN/NUT]
Δ_vH		(323–473)	97.5	398	GC	[2002LEI/CHA]
C ₂₂ H ₁₄	[53-70-3] $\Delta_{\text{fus}}H$	dibenz[a,h]anthracene	28.4	539.7	DSC	[2010KES/AUC]
	$\Delta_{\text{fus}}H$		31.16	544.2		[1991ACR]
	$\Delta_{\text{sub}}H$		134.1		GS	[1995NAS/LEN]
	$\Delta_{\text{sub}}H$	(436–462)	162 ± 6	298	TE,ME	[1980DEK]
	$\Delta_{\text{sub}}H$	(417–502)	141.8	457	ME	[1967WAK/INO]
	Δ_vH		131.1 ± 1.4	298	CGC	[2008HAN/NUT]
	Δ_vH	(323–473)	99.4	398	GC	[2002LEI/CHA]
C ₂₂ H ₁₄	[194-69-4] $\Delta_{\text{fus}}H$	benzo[c]chrysenes	22.7	398.5	DSC	[2010KES/AUC]
	C ₂₂ H ₁₄	[214-17-5] $\Delta_{\text{fus}}H$	1,2:6,7-dibenzophenanthrene (benzo[b]chrysenes)	25.3	574.2	DSC

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{sub}}H$		136.4	417	ME	[1967WAK/INO]
	$\Delta_{\text{sub}}H$	(398–513)	136.9	413	A	[1987STE/MAL]
C ₂₂ H ₁₄	[135-48-8]	pentacene				
	$\Delta_{\text{sub}}H$	(443–483)	156.9 ± 13.6	463	ME	[1998OJA/SUU]
	$\Delta_{\text{sub}}H$	(494–526)	154 ± 5	512	ME,TE	[1980DEK]
	$\Delta_{\text{sub}}H$	(495–530)	184 ± 10	298	ME,TE	[1980DEK]
	$\Delta_{\text{sub}}H$	(455–555)	157.7	505	ME	[1967WAK/INO]
C ₂₂ H ₁₄	[213-46-7]	picene				
	$\Delta_{\text{fus}}H$		35.19	637.2		[1980SMI]
	$\Delta_{\text{sub}}H$	(425–488)	140.1	456	ME	[1967WAK/INO]
	$\Delta_{\text{sub}}H$	(409–527)	140.7	424		[1987STE/MAL]
C ₂₂ H ₁₄ N ₄	[1154424-98-2]	1,4-bis((pyridine-3-yl)imino)methylbenzene				
	$\Delta_{\text{fus}}H$		40.7	438.2	DSC	[2008STI/CIN]
C ₂₂ H ₁₄ O ₄	[3363-97-1]	1,4-bis(phenylglyoxaloyl)benzene				
	$\Delta_{\text{fus}}H$		32.3	425.1		[1996DOM/HEA]
C ₂₂ H ₁₆	[68862-02-2]	2-(biphen-4-yl)naphthalene				
	$\Delta_{\text{fus}}H$		25.1	489.5	DSC	[2008ROU/LIM]
	$\Delta_{\text{sub}}H$	(405–437)	137.1 ± 0.4	421	ME	[2008ROU/LIM]
	$\Delta_{\text{sub}}H$	(405–437)	140.2 ± 1.3	298	ME	[2008ROU/LIM]
C ₂₂ H ₁₆	[87294-80-2]	2-(biphen-3-yl)naphthalene				
	$\Delta_{\text{fus}}H$		18.5	346.3	DSC	[2008ROU/LIM]
	Δ_vH		118.6 ± 1.5	298		[2008ROU/LIM]
C ₂₂ H ₁₆ O	[81-37-8]	3,8-dimethylnaphtho[3,2,1-kl]xanthene (3,8-dimethylceroxene)				
	$\Delta_{\text{sub}}H$	(373–433)	138.2	388	A	[1987STE/MAL]
C ₂₂ H ₁₇ NO ₂	[85084-64-7]	2-ethylphenyl acridine-9-carboxylate				
	$\Delta_{\text{fus}}H$		30.4	393	DSC	[2010KRZ/MAL]
C ₂₂ H ₁₇ NO ₂	[128649-37-6]	2,5-dimethylphenyl acridine-9-carboxylate				
	$\Delta_{\text{fus}}H$		37.9	457	DSC	[2010KRZ/MAL]
C ₂₂ H ₁₇ NO ₂	[na]	2,6-dimethylphenyl acridine-9-carboxylate				
	$\Delta_{\text{fus}}H$		29.1	435	DSC	[2010KRZ/MAL]
C ₂₂ H ₁₇ NO ₂	[na]	3,4-dimethylphenyl acridine-9-carboxylate				
	$\Delta_{\text{fus}}H$		29.2	442	DSC	[2010KRZ/MAL]
C ₂₂ H ₁₇ NO ₂	[na]	3,5-dimethylphenyl acridine-9-carboxylate				
	$\Delta_{\text{fus}}H$		38.4	469	DSC	[2010KRZ/MAL]
C ₂₂ H ₁₇ NO ₃ S	[36245-88-2]	2-(3-methoxypropyl)-1H-xantheno[2,2,9-def]-isoquinoline-1,3(2H)-dione				
	$\Delta_{\text{sub}}H$	(605–647)	111.8	620	A	[1987STE/MAL]
	$\Delta_{\text{sub}}H$	(647–685)	150.8	662	A	[1987STE/MAL]
C ₂₂ H ₁₈ F ₂ O	[145698-40-4]	4-(6-hexenyloxy)-3',4'-difluorodiphenyldiacetylene				
	$\Delta_{\text{fus}}H$		37.45	370	DSC	[1993JUA/CHE]
C ₂₂ H ₁₈ F ₂ O	[153038-12-1]	4-(cis-4-hexenyloxy)-3',4'-difluorodiphenyldiacetylene				
	$\Delta_{\text{fus}}H$		35.32	364.4	DSC	[1993JUA/CHE]
C ₂₂ H ₁₈ F ₂ O	[153038-13-2]	4-(cis-3-hexenyloxy)-3',4'-difluorodiphenyldiacetylene				
	$\Delta_{\text{fus}}H$		30.97	364.6	DSC	[1993JUA/CHE]

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₂₂ H ₁₈ N ₂ O ₂	[116-77-8] $\Delta_{\text{sub}}H$	1-amino-2-methyl-4-[(4-methylphenyl)amino]-9,10-anthraquinone (418–435)	142.2 ± 2.3	426		[1984KRI]
C ₂₂ H ₁₈ N ₂ O ₂	[6408-50-8] $\Delta_{\text{sub}}H$	1-(N-methylamino)-4-[(3-methylphenyl)amino]-9,10-anthraquinone (418–434)	129.0 ± 4.7	426		[1984KRI]
C ₂₂ H ₁₈ N ₂ O ₂	[128-85-8] $\Delta_{\text{sub}}H$	1-(N-methylamino)-4-[(4-methylphenyl)amino]-9,10-anthraquinone (403–426)	153.9 ± 3.9	414		[1984KRI]
C ₂₂ H ₁₈ O ₄	[523-31-9] Δ_vH	dibenzyl phthalate (445–513)	121.4	460	A	[1987STE/MAL]
C ₂₂ H ₁₉ Br ₂ NO ₃	[52918-00-5] $\Delta_{\text{fus}}H$	(S)- α -cyano-3-phenoxybenzyl (1R)- <i>cis</i> -3-(2,2-dibromovinyl)-2,2-dimethylcyclopropanecarboxylate 40.71	372.9		DSC	[1990DON/DRE]
		Note: Chemical Abstracts gives a CAN Registry Number of [52918-63-5] for the compound studied by the authors of Ref. [1990DON/DRE]. The authors of the paper; however, give a Registry Number of [52918-00-5]				
C ₂₂ H ₁₉ Br ₂ NO ₃	[52918-63-5] $\Delta_{\text{fus}}H$	(1R,3R)-3-(2,2-dibromoethyl)-2,2-dimethylcyclopropanecarboxylic acid, (S)-cyano(3-phenoxyphenyl)methyl ester (deltamethrin) (80–400)	26.73	372.2	AC	[2005XUE/WAN]
C ₂₂ H ₁₉ N ₅ O ₃	[244272-54-6] $\Delta_{\text{fus}}H$	6-(4-biphenyl)-3,9-dihydro-3-[(2-hydroxyethoxy)methyl]-9-oxo-5H-imidazol[1,2-a]pyrine 45.43	484.5		DSC	[1999ZIE/GOL]
C ₂₂ H ₂₀ N ₂ O ₄	[36360-34-6] $\Delta_{\text{sub}}H$	N,N'-bis(2-methoxyphenyl)terephthalamide (183–197)	197.5 ± 4.2		ME	[1973HAM/MIT, 1977PED/RYL]
C ₂₂ H ₂₀ N ₂ O ₄	[6957-81-9] $\Delta_{\text{sub}}H$	N,N'-bis(3-methoxyphenyl)terephthalamide (183–197)	209.2 ± 8.4		ME	[1973HAM/MIT2, 1977PED/RYL]
C ₂₂ H ₂₀ N ₂ O ₄	[7144-15-2] $\Delta_{\text{sub}}H$	N,N'-bis(4-methoxyphenyl)terephthalamide 227.6 ± 8.4			ME	[1973HAM/MIT2, 1977PED/RYL]
C ₂₂ H ₂₁ Cl ₃ N ₄ O	[168273-06-1] $\Delta_{\text{fus}}H$	5-(4-chlorophenyl)-1-(2,4-dichlorophenyl)-4-methyl-N-1-piperidinyl-1H-pyrazole-3-carboxamide (rimonabant) 36.1	427.9		DSC	[2007BER/WAS]
C ₂₂ H ₂₁ F	[193472-71-8] $\Delta_{\text{fus}}H$	2-benzyl-2-fluoro-1,3-diphenylpropane 24.35	363.6			[1997SCH/VER]
	$\Delta_{\text{sub}}H$	127.5 ± 0.8	298			[1997SCH/VER]
C ₂₂ H ₂₁ F ₂₅	[93454-72-9] $\Delta_{\text{trs}}H$	1,1,1,2,2,3,3,4,4,5,5,6,6,7,7,8,8,9,9,10,10,11,11,12,12-pentacosafuorodocosane 1	207			
	$\Delta_{\text{fus}}H$	9.5	342			
	$\Delta_{\text{fus}}H$	25.8	365		DSC	[1991HOP/MOL]
	$\Delta_{\text{trs}}H$	7.5	339.2			
	$\Delta_{\text{fus}}H$	22.2	357.2		DSC	[1986RUS/RAB]
C ₂₂ H ₂₂	[43044-69-5] $\Delta_{\text{fus}}H$	1,1,1-triphenylbutane 21.84	351.1		DSC	[1999VER3]
	$\Delta_{\text{sub}}H$	(323–347)	112.1 ± 1.1	335	GS	[1999VER3]
	$\Delta_{\text{sub}}H$	(323–347)	114.3 ± 1.1	298	GS	[1999VER3]
C ₂₂ H ₂₂	[4742-04-5] Δ_vH	tribenzylmethane (395–648)	79	521		[1999DYK/SVO]
C ₂₂ H ₂₃ NO ₃	[39515-41-8] $\Delta_{\text{fus}}H$	2,2,3,3-tetramethylcyclopropanecarboxylic acid, cyano(3-phenoxyphenyl)methyl ester (fenpropathrin) 18.57	322.5			[2001TAN/XUE, 1999XUE/TAN]

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₂₂ H ₂₄ N ₂ O ₂	[16087-30-2]	N,N'-ethylenebis(3-amino-1-phenylbut-2-en-1-one)				
	$\Delta_{\text{sub}}H$	(407–426)	192.9 ± 5.3	415	ME	[1995RIB/RIB]
	$\Delta_{\text{sub}}H$		198.8 ± 5.3	298		[1995RIB/RIB]
C ₂₂ H ₂₄ O ₃	[138306-50-0]	4-methyl-4-propyl-1,8-diphenyl-2,3,5-trioxabicyclo[4.3.0]non-7-ene				
	$\Delta_{\text{fus}}H$		16.6	351.2	DSC	[1991JEF/JAB]
C ₂₂ H ₂₄ O ₃	[104225-44-7]	3-([1,1-biphenyl]-4-ylcarbonyl)-1,2,2-trimethylcyclopentanecarboxylic acid				
	$\Delta_{\text{fus}}H$		27.69	444.2	DSC	[1992TER/PAU]
C ₂₂ H ₂₅ F ₂₁	[93454-71-8]	1,1,1,2,2,3,3,4,4,5,5,6,6,7,7,8,8,9,9,10,10-heneicosofluorodocosane				
	$\Delta_{\text{trs}}H$		6	334.1		
	$\Delta_{\text{fus}}H$		27	338.1	DSC	[1992HOP/MOL]
C ₂₂ H ₂₅ NO	[6018-34-4]	(+) 5,7,8,15-tetrahydro-3,4-dimethoxy-6,15-dimethyl-3]benzodioxolo-[5,6e] [2]benzazecin-14(6)-one (corycavidine)				
	$\Delta_{\text{fus}}H$		38.07	468.3		[2000KAM/YOS]
C ₂₂ H ₂₅ NO ₃	[126675-75-0]	2-(4-nitrophenyl)-1-[4-(trans-4-ethylcyclohexyl)phenyl]ethanone				
	$\Delta_{\text{fus}}H$		37.32	445.1	DSC	[2002SPA/DZI]
C ₂₂ H ₂₅ NO ₆	[836602-50-7]	methyl naltrexone-3-O-carbonate				
	$\Delta_{\text{fus}}H$		10.92	393.7	DSC	[2004PIL/HAM]
C ₂₂ H ₂₆	[59358-70-2]	1,1'-diphenyl-1,1'-bicyclopentyl				
	$\Delta_{\text{fus}}H$		31.38	414	DSC	[1983KRA/BEC]
	$\Delta_{\text{sub}}H$		141.4	141.4	E,B	[1983KRA/BEC]
C ₂₂ H ₂₆ FNO ₂	[1050-79-9]	1-(4-fluorophenyl)-4-[4-hydroxy-4-(4-methylphenyl)-1-piperidinyl]-1-butanone (moperone)				
	$\Delta_{\text{fus}}H$ (I)		37.4	398.2		
	$\Delta_{\text{fus}}H$ (II)		15	468.2	DSC	[1981DRA/AZI]
C ₂₂ H ₂₆ N ₂ O ₂	[17354-14-2]	1,4-bis(N-butylamino)-9,10-anthraquinone				
	$\Delta_{\text{sub}}H$	(389–398)	116.4 ± 2.3	394		[1984KRI]
C ₂₂ H ₂₆ N ₂ O ₂	[10720-45-7]	1,4-bis(N-isobutylamino)-9,10-anthraquinone				
	$\Delta_{\text{sub}}H$	(368–388)	96.4 ± 2.1	378		[1984KRI]
C ₂₂ H ₂₇ NO ₂	[1050-79-9]	pregna-2,4-dien-20-yno[2,3-d]isoxazol-17-ol (danazol)				
	$\Delta_{\text{fus}}H$		35.5	501.8	DSC	[2007BER/WAS]
C ₂₂ H ₂₈ N ₂ O	[437-38-7]	N-phenyl-N[1-(2-phenylethyl)-4-piperidinyl]propanamide (fentanyl)				
	$\Delta_{\text{fus}}H$		30.1	358.3	DSC	[2008GUP/GAN]
	$\Delta_{\text{fus}}H$		22.51	357.2	DTA	[1988ROY/FLY]
	$\Delta_{\text{sub}}H$	(423–493)	144.6 ± 7.2	298	Vap+Fus	[2008GUP/GAN]
	Δ_vH	(423–493)	107.2 ± 4.2	458	TGA	[2008GUP/GAN]
C ₂₂ H ₂₈ N ₂ O ₂	[145513-29-7]	(4R, 4'R, 5R, 5'R)-5,5-diphenyl-3,3',4,4'-tetramethyl-2,2'-bioxazolidine				
	$\Delta_{\text{fus}}H$		31.9	394		[1995TOR/GUD]
	$\Delta_{\text{sub}}H$	(349–358)	130.8 ± 0.8	356	ME	[1995TOR/GUD]
C ₂₂ H ₂₈ N ₂ O ₂	[145438-85-3]	(2R, 3R, 6R, 7R)-2,6-diphenyl-3,4,7,8-tetramethyl-cis-perhydro-[1,4]-oxazino-[3,2-b]-[1,4]-oxazine				
	$\Delta_{\text{fus}}H$		20.9	396.8		[1995TOR/GUD]
	$\Delta_{\text{sub}}H$	(353–364)	116.6 ± 1.0	358	ME	[1995TOR/GUD]
C ₂₂ H ₂₈ N ₂ O ₂	[145438-85-3]	(2R, 3S, 6R, 7S)-2,6-diphenyl-3,4,7,8-tetramethyl-cis-perhydro-[1,4]-oxazino-[3,2-b]-[1,4]-oxazine				
	$\Delta_{\text{fus}}H$		18.4	379.4		[1995TOR/GUD]

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{sub}}H$	(356–364)	123.1 ± 1.6	358	ME	[1995TOR/GUD]
C ₂₂ H ₂₈ O	[33574-11-7]	2,4,6-triisopropylbenzophenone				
	$\Delta_{\text{sub}}H$	(353–364)	116 ± 7	298	C	[1982INA/MUR2]
C ₂₂ H ₂₈ O	[33574-16-2]	3',5'-diisopropyl-4,4-dimethyl-3-phenyl-1,2-benzocyclobuten-3-ol				
	$\Delta_{\text{sub}}H$	(354–364)	117.9	298	C	[1982INA/MUR2]
C ₂₂ H ₂₈ O ₂	[54048-10-1]	13-ethyl-17-hydroxy-11-methylene-18,19-dinorpregn-4-en-20-yn-3-one (etonogestrel)				
	$\Delta_{\text{fus}}H$		31.15	472.2		[2002VAN/KRU]
C ₂₂ H ₂₈ O ₃	[2353-34-6]	3-[(1-oxobutyl)oxy]-estra-1,3,5(10)-trien-17-one				
	$\Delta_{\text{fus}}H$		22	381	DSC	[1990YAN/EIR]
C ₂₂ H ₂₈ O ₃	[na]	19-nor-17 α -ethynyl-17 β -acetoxy-4-androsten-3-one				
	$\Delta_{\text{fus}}H$		27.3	480		[1996DOM/HEA]
C ₂₂ H ₂₉ FO ₅	[50-02-2]	9-fluoro-11 β ,17,21-trihydroxy-16 α -methylpregna-1,2-diene-3,20-dione (dexamethasone)				
	$\Delta_{\text{fus}}H$		42.02	539		[1994REG/CHM]
C ₂₂ H ₂₉ NO ₂	[133544-40-8]	4- <i>n</i> -octyloxy-N-(4-methoxybenzylidene)aniline				
	$\Delta_{\text{fus}}H$		42.29	377.3		[1995MIY/NAK]
C ₂₂ H ₃₀ N ₂ O ₂ S	[56030-54-7]	N-[4-(methoxymethyl)-1-[2-(2-thienyl)ethyl]-4-piperidinyl]-N-phenylpropanamide (sufentanil)				
	$\Delta_{\text{fus}}H$		23.85	370.2	DTA	[1988ROY/FLY]
C ₂₂ H ₃₁ NO ₄	[23257-62-7]	N,N-bis(3-phenoxy-2-hydroxypropyl)butyl amine				
	$\Delta_{\text{sub}}H$	(363–411)	114.3	378	A	[1987STE/MAL]
	$\Delta_{\text{sub}}H$		146.0 ± 4.2			[1976KUZ/MIR]
C ₂₂ H ₃₂ O ₃	[57-85-2]	testosterone propionate				
	$\Delta_{\text{fus}}H$		25.64	393		[1994REG/CHM]
C ₂₂ H ₃₃ N ₃ O ₂	[765303-86-4]	pyrimethanil decylate				
	$\Delta_{\text{fus}}H$	(78–373)	45.88	311	AC	[2005SUN/LIU]
C ₂₂ H ₃₄ N ₄ O ₄	[501946-58-3]	1,1'-(1,12-dodecanediyl)bisthymine				
	$\Delta_{\text{fus}}H$		43.95	462	DSC	[2002ITA/KAM]
C ₂₂ H ₃₆ O ₄	[118476-23-6]	2,5-di- <i>n</i> -octyloxy-1,4-benzoquinone				
	$\Delta_{\text{fus}}H$		9.4	358.2		
	$\Delta_{\text{fus}}H$		43.0	405.8	DSC	[1996KEE/VAN]
C ₂₂ H ₃₈	[1459-09-2]	hexadecylbenzene				
	Δ_vH	(505–688)	79.5	520		[1999DYK/SVO]
C ₂₂ H ₃₈	[54934-70-2]	1,1-bis(decahydro-1-naphthyl)ethane				
	Δ_vH	(432–503)	77.3	447	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₂₂ H ₃₈	[54934-69-9]	1,2-bis(decahydro-1-naphthyl)ethane				
	Δ_vH	(440–507)	89.3	455	A	[1987STE/MAL]
C ₂₂ H ₃₈	[54934-71-3]	1,5-dicylopentyl-3-(2-cyclopentylethyl)-2-pentene				
	Δ_vH	(427–492)	81.4	442	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₂₂ H ₃₈	[62678-54-0]	<i>meso</i> 3,4-di(1-cyclohexen-1-yl)-2,2,5,5-tetramethylhexane				
	$\Delta_{\text{sub}}H$	(347–404)	117.2 ± 2.4	376	T	[1993HER/BEC]
C ₂₂ H ₃₈ N ₄ O ₂	[126235-07-2]	8-pentadecyltheophylline				
	$\Delta_{\text{fus}}H$		27.2	413.7	DSC	[1991ACR, 1989GON/KRA]
C ₂₂ H ₄₀	[55255-85-1]	1,5-dicylopentyl-3-(2-cyclopentylethyl)-2-pentane				
	Δ_vH	(430–494)	83.6	445	A, MG	[1987STE/MAL, 1955SCH/WHI]

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₂₂ H ₄₀ O ₂	[31067-26-2] $\Delta_{\text{fus}}H$	3,3,6,6,10,10,13,13-octamethylcyclotetradecane-1,8-dione	24.7	492.2		[1972BOR/DAL2]
C ₂₂ H ₄₀ O ₄	[38734-13-3] $\Delta_{\text{fus}}H$	1,10-cyclooctadecanedione <i>bis</i> ethylene ketal	33.56	378.2		[1972ALV/BOR]
C ₂₂ H ₄₂	[na] $\Delta_{\text{sub}}H$	<i>meso</i> -(E, E)-5,6-di- <i>tert</i> -butyl-2,2,9,9-tetramethyl-3,7-decadiene (297–353)	110.0 ± 1.7	325	T	[1995HER/VER]
C ₂₂ H ₄₂	[na] $\Delta_{\text{sub}}H$	(<i>dl</i>)-(E, E)-5,6-di- <i>tert</i> -butyl-2,2,9,9-tetramethyl-3,7-decadiene (297–346)	74.4 ± 1.7	307	T	[1995HER/VER]
C ₂₂ H ₄₂ O ₂	[142-77-8] Δ_vH	butyl oleate (353–393)	97.7	368	A	[1987STE/MAL]
C ₂₂ H ₄₂ O ₂	[112-86-7] $\Delta_{\text{us}}H$	<i>cis</i> 13-docosenoic acid (erucic acid)	8.9	282.2		
	$\Delta_{\text{fus}}H$		54	307.2		[1997SAT/YAN]
	Δ_vH	(479–655)	98.2	494	A	[1987STE/MAL]
C ₂₂ H ₄₂ O ₂	[506-33-2] Δ_vH	<i>trans</i> 13-docosenoic acid (482–656)	103.4	497	A	[1987STE/MAL]
C ₂₂ H ₄₂ O ₄	[123-79-5] Δ_vH	dioctyl adipate (373–493)	99	388	A	[1987STE/MAL]
C ₂₂ H ₄₂ O ₄	[2449-10-7] Δ_vH	dihexyl sebacate	99.9	344	TGA	[1990KIS/SHO]
	Δ_vH		106.4 ± 3.7	298	TGA	[1990KIS/SHO]
C ₂₂ H ₄₂ O ₆	[141-19-5] Δ_vH	<i>bis</i> (2-butoxyethyl) sebacate (368–423)	120.3	383	A, ME	[1987STE/MAL, 1948SMA/SMA]
C ₂₂ H ₄₃ NO ₃	[14379-42-1] $\Delta_{\text{fus}}H$	N-hexadecanoyl-(<i>l</i>)-leucine	46.1	367.1	DSC	[1986MIY/MAT]
C ₂₂ H ₄₃ NO ₃	[21394-54-7] $\Delta_{\text{us}}H$	N-hexadecanoyl-(<i>dl</i>)-leucine	4.3	333.1		
	$\Delta_{\text{fus}}H$		60.6	355.1	DSC	[1986MIY/MAT]
C ₂₂ H ₄₄	[1599-67-3] Δ_vH	1-docosene (401–640)	95.6	416		[1999DYK/SVO]
C ₂₂ H ₄₄	[6812-38-0] Δ_vH	hexadecylcyclohexane (507–689)	79.6	522		[1999DYK/SVO]
	Δ_vH		109.3	298		[1971WIL/ZWO]
C ₂₂ H ₄₄	[23014-56-4] $\Delta_{\text{fus}}H$	1,1,10,10-tetramethylcyclooctadecane	39.58	359.2		[1974BJO/BOR]
C ₂₂ H ₄₄ N ₂ O ₂	[31827-03-9] $\Delta_{\text{fus}}H$	N,N'-di-n-hexylsebacamide	53.56	415		[1996DOM/HEA]
C ₂₂ H ₄₄ O ₂	[123-95-5] $\Delta_{\text{us}}H$	butyl stearate	2.22	288.4		
	$\Delta_{\text{fus}}H$		37.48	299.7		[1986KAL/JAC]
	Δ_vH	(352–399)	99.9	367	A, T	[1987STE/MAL, 1949PER/WEB]
C ₂₂ H ₄₄ O ₂	[18281-05-5] $\Delta_{\text{fus}}H$	ethyl eicosanoate	68.62	315		[1967OMA]

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{sub}}H$	(307–313)	171.5	310	ME	[1987STE/MAL, 1967OMA]
	Δ_vH	(318–460)	113.7	333	A	[1987STE/MAL]
C₂₂H₄₄O₂	[6064-90-0]	methyl heneicosanoate				
	$\Delta_{\text{fus}}H$		75.1	321.2		[2004CHI/ZHA]
	Δ_vH	(459–529)	95.6	474	A, EST	[1987STE/MAL, 1963ROS/SCH]
C₂₂H₄₄O₂	[36528-28-6]	decyl dodecanoate				
	$\Delta_{\text{fus}}H$		63.67	293.2		[1999BAL/WEI]
C₂₂H₄₄O₂	[42232-25-7]	hexyl hexadecanoate				
	$\Delta_{\text{fus}}H$		56.19	287.6		[1999BAL/WEI]
C₂₂H₄₄O₂	[112-85-6]	docosanoic acid				
	Δ_vH	(373–600)	122.3	388	A	[1987STE/MAL]
C₂₂H₄₄O₄	[56444-64-5]	2,2,13,13-tetramethyl-1,3,12,14-tetraoxacyclodocosane				
	$\Delta_{\text{fus}}H$		61.9	374		[1975BOR]
C₂₂H₄₅Br	[6938-66-5]	1-bromodocosane				
	$\Delta_{\text{us}}H$		23.14	303.8		
	$\Delta_{\text{fus}}H$		44.98	317.1		[1953HOF/DEC]
C₂₂H₄₅NO	[3061-75-4]	docosamide				
	$\Delta_{\text{fus}}H$		63.3	383.3	DSC	[2008ABA/BAD]
C₂₂H₄₅NO	[74534-12-6]	N-hexyl hexadecanamide				
	$\Delta_{\text{fus}}H$		57	343.1	DSC	[1993ACR]
C₂₂H₄₅NO	[55334-54-8]	N,N-dioctyl hexanamide				
	Δ_vH	(463–513)	88.1 ± 1.0	298	CGC	[2009PAN/ANT]
C₂₂H₄₆	[629-97-0]	docosane				
	$\Delta_{\text{us}}H$		27.3	315.8		
	$\Delta_{\text{us}}H$		<0.3	316.1		
	$\Delta_{\text{fus}}H$		49.1	316.6	DSC	[2004MON/RAJ]
	$\Delta_{\text{us}}H$		29.51	315.2		
	$\Delta_{\text{fus}}H$		47.84	316.1		[1990DOM/HEA]
	$\Delta_{\text{sub}}H$		172.6 ± 2.0	391	B	[1994PIA/FON]
	$\Delta_{\text{sub}}H$		U 151.1 ± 10	298	B	[1991PIA/POM]
	Δ_vH	(434–539)	111.9	298	CGC	[2004CHI/HAN]
	Δ_vH		114.9 ± 0.3	298	CGC	[2002CHI/WEB]
	Δ_vH		115.6 ± 1.9	298	CGC	[1997CHI/WIL]
	Δ_vH	(453–503)	115.6	298	CGC	[1995CHI/HOS]
	Δ_vH	(453–573)	84.3	468		[1994MOR/KOB]
	Δ_vH	(372–410)	124 ± 2	391	TE	[1994PIA/FON]
	Δ_vH	(358–490)	89.9	373	TE,ME,GS	[1991PIA/POM]
	Δ_vH	(353–462)	100.9	368		[1988SAS/JOS]
	Δ_vH	(431–642)	91.3	446	A, EST	[1987STE/MAL, 1966KUD/ZWO]
C₂₂H₄₆	[1560-82-3]	2-methylheneicosane				
	Δ_vH	(485–640)	76.1	500	A	[1987STE/MAL]
C₂₂H₄₆	[6418-47-9]	3-methylheneicosane				
	Δ_vH	(484–631)	74.4	499	A	[1987STE/MAL, 1999DYK/SVO, 1959TER/BRI]
C₂₂H₄₆	[25117-29-7]	4-methylheneicosane				

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(497–632)	70.9	494	A	[1987STE/MAL, 1999DYK/SVO, 1959TER/BRI]
C ₂₂ H ₄₆	[25117-37-7]	5-methylheneicosane				
	$\Delta_v H$	(483–632)	73.9	498	A	[1987STE/MAL, 1999DYK/SVO]
C ₂₂ H ₄₆	[75163-98-3]	2,4-dimethyleicosane				
	$\Delta_v H$	(471–603)	77.5	486	A	[1987STE/MAL, 1999DYK/SVO, 1959TER/BRI]
C ₂₂ H ₄₆	[102886-19-1]	2,4,6-trimethylnonadecane				
	$\Delta_v H$	(470–587)	77.3	485	A	[1987STE/MAL, 1999DYK/SVO, 1959TER/BRI]
C ₂₂ H ₄₆	[71005-15-7]	8-heptylpentadecane				
	$\Delta_v H$	(298–313)	107.7	305	A	[1987STE/MAL]
C ₂₂ H ₄₆ O	[661-19-8]	1-docosanol				
	$\Delta_{\text{fus}} H$	(80–400)	85.07	340.8	AC	[2008TON/TAN3]
	$\Delta_{\text{fus}} H$		86.06	343.9		[2001VAN/OON2]
	$\Delta_{\text{us}} H$		17.24	333.9		
	$\Delta_{\text{fus}} H$		46.57	345.2		[1979KUC/SKU]
	$\Delta_{\text{sub}} H$	(335–341)	206.7 ± 10	330	ME	[1965DAV/KYB, 1987STE/MAL]
	$\Delta_{\text{sub}} H$		238.5 ± 10	298		[1965DAV/KYB]
	$\Delta_v H$		135.9 ± 0.8	298	CGC	[2006NIC/KWE]
	$\Delta_v H$	(344–459)	115.3	351	A, ME	[1987STE/MAL, 1965DAV/KYB]
C ₂₂ H ₄₆ O ₂	[22513-81-1]	1,22-docosanediol				
	$\Delta_{\text{us}} H$		39.8	369.7		
	$\Delta_{\text{fus}} H$		46.5	379.4	DSC	[1999OGA/NAK]
C ₂₂ H ₄₆ O ₄ S ₂	[na]	(l)-rhamnose dioctyl dithioacetal				
	$\Delta_{\text{fus}} H$		54.7	387.9	DSC	[1989VAN/VAN]
C ₂₂ H ₄₆ S	[7773-83-3]	1-docosanethiol				
	$\Delta_v H$	(437–680)	107.7	452	EST	[1999DYK/SVO]
C ₂₃ H ₁₅ ClO ₃	[3691-35-8]	2-[(4-chlorophenyl)phenylacetyl]-1 <i>H</i> -indene-1,3(2 <i>H</i>)-dione				
	$\Delta_{\text{fus}} H$		34.54	416.5	DSC	[1990DON/DRE]
C ₂₃ H ₁₅ F ₁₇ OS	[246544-01-4]	2-(perfluoro- <i>n</i> -octyl)ethylthiomethyl biphenyl-4-yl ether				
	$\Delta_{\text{fus}} H$		47.4	362.1	DTA	[1999DEG/GUI]
C ₂₃ H ₁₅ F ₁₇ S	[246543-98-6]	2-(perfluoro- <i>n</i> -octyl)ethylthiomethyl biphenyl-4-yl				
	$\Delta_{\text{fus}} H$		58.3	353.2	DTA	[1999DEG/GUI]
C ₂₃ H ₁₉ NO ₂	[1041479-15-5]	2-isopropylphenyl acridine-9-carboxylate				
	$\Delta_{\text{fus}} H$		29.0	396	DSC	[2010KRZ/MAL]
C ₂₃ H ₁₉ NO ₂	[1134294-42-0]	2,4,6-trimethylphenyl acridine-9-carboxylate				
	$\Delta_{\text{fus}} H$		24.4	405	DSC	[2010KRZ/MAL]
C ₂₃ H ₂₁ F ₃ O	[145698-52-8]	4- <i>n</i> -hexyloxy-4'-trifluoromethyldiphenyldiacetylene				
	$\Delta_{\text{fus}} H$		33.98	394.8	DSC	[1993JUA/CHE]
C ₂₃ H ₂₁ F ₇ N ₄ O ₃	[170729-80-3]	5-[[[(2 <i>R</i> ,3 <i>S</i>)-2-[(1 <i>R</i>)-1-[3,5-bis(trifluoromethyl)phenyl]ethoxy]-3-(4-fluorophenyl)-4-morpholinyl]methyl]-1,2-dihydro-3 <i>H</i> -1,2,4-triazol-3-one (Aprepitant)				
	$\Delta_{\text{fus}} H$ (I)		53.7	526.8	DSC	
	$\Delta_{\text{fus}} H$ (II)		52.4	526.2	DSC	[2008BRA/GEL]

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₂₃ H ₂₂ O ₆	[83-79-4]	[2R-(2a,6a α ,12a α)]-1,2,12,12a-tetrahydro-8,9-dimethoxy-2-(1-methylethenyl)-[1]benzopyrano[3,4-b]furo[2,3-h] [1]benzopyran-6(6aH)-one (Rotenone)				
	$\Delta_{\text{fus}}H$		35.64	437.9	DSC	[1990DON/DRE]
C ₂₃ H ₂₄ N ₆ O ₄	[na]	2-[[4-[[[(2-acetoxy)ethyl]butylamino]-2-methylphenyl]azo]-5-nitro-1,3-benzenedicarbonitrile				
	$\Delta_{\text{fus}}H$		37.88	424.2		[1991BAU/WEB]
C ₂₃ H ₂₄ O ₆	[170464-53-6]	<i>tris</i> (ethoxycarbonyl)-9-fluorenylmethane				
	$\Delta_{\text{sub}}H$		143.2	298	GS	[1995RAK/VER]
	Δ_vH	(359–393)	107.5 \pm 0.7		GS	[1995RAK/VER]
C ₂₃ H ₂₅ BrN ₆ O ₁₀	[na]	N-[5- <i>bis</i> [(2-acetyloxy)ethyl]amino]-2-[(2-bromo-4,6-dinitrophenyl)azo]-4-methoxyphenyl] acetamide				
	$\Delta_{\text{fus}}H$		57.28	421.2		[1991BAU/WEB]
C ₂₃ H ₂₅ F	[154393-25-6]	1-adamantylfluorodiphenylmethane				
	$\Delta_{\text{sub}}H$	(353–393)	125.9 \pm 1.3	373	T	[1994SCH/BEC]
C ₂₃ H ₂₆ O ₆	[183212-67-1]	1,1,1- <i>tris</i> (ethoxycarbonyl)-2,2-diphenylethane				
	$\Delta_{\text{fus}}H$		29.5	333.2		[1995RAK/VER]
	$\Delta_{\text{sub}}H$		140.1	298	GS	[1995RAK/VER]
	Δ_vH	(344–394)	109.3 \pm 1.0		GS	[1995RAK/VER]
C ₂₃ H ₂₇ Cl ₂ N ₃ O ₂	[129722-12-9]	7-[4-[4-(2,3-dichlorophenyl)-1-piperazinyl]butoxy]-3,4-dihydro-2(1H)-quinoline (aripiprazole)				
	$\Delta_{\text{fus}}H$ (I)		38.36	422.1	DSC	
	$\Delta_{\text{fus}}H$ (II)		41.51	416.3	DSC	
	$\Delta_{\text{fus}}H$ (III)		39.97	412.4	DSC	
	$\Delta_{\text{fus}}H$ (IV)		40.76	408.1	DSC	[2009BRA/GEL]
C ₂₃ H ₂₇ NO ₃	[126675-76-1]	2-(4-nitrophenyl)-1-[4-(<i>trans</i> -4-propylcyclohexyl)phenyl]ethanone				
	$\Delta_{\text{fus}}H$		38.87	436.5	DSC	[2002SPA/DZI]
C ₂₃ H ₂₇ NO ₃ S	[313057-12-4]	4-(7-undecenyloxy)phenyl 5-cyano-2-thiophenecarboxylate				
	$\Delta_{\text{fus}}H$		52.72	346.1	DSC	[2000WU/WAN]
C ₂₃ H ₂₇ NO ₆	[836602-51-8]	ethyl naltrexone-3-O-carbonate				
	$\Delta_{\text{fus}}H$		18.99	404.2	DSC	[2004PIL/HAM]
C ₂₃ H ₂₈ ClN ₃ O ₅ S	[10238-21-8]	5-chloro-N-[2-[4-[[[(cyclohexylamino)carbonyl]amino]sulfonyl]phenyl]ethyl]-2-methoxybenzamide (glyburide)				
	$\Delta_{\text{fus}}H$		46.3	446.8	DSC	[2007BER/WAS, 2006WAS/HOL]
	$\Delta_{\text{fus}}H$		53.35	450.2		[2000HAN/PAR]
C ₂₃ H ₂₈ N ₂ O ₅	[na]	N,N-dimethyl naltrexone-3-O-carbamate				
	$\Delta_{\text{fus}}H$		22.01	480.2	DSC	[2009VAD/BAN]
C ₂₃ H ₃₀ O ₃	[128788-26-1]	3-[(1-oxopentyl)oxy]-estra-1,3,5(10)-trien-17-one				
	$\Delta_{\text{fus}}H$		25	398	DSC	[1990YAN/EIR]
C ₂₃ H ₃₀ O ₄ S	[313057-16-8]	4-(7-undecenyloxy)phenyl 5-methoxy-2-thiophenecarboxylate				
	$\Delta_{\text{fus}}H$		61.92	334.1	DSC	[2000WU/WAN]
C ₂₃ H ₃₀ O ₆	[52-21-1]	prednisolone acetate				
	$\Delta_{\text{fus}}H$		42.3	515		[1997CEN/MEL]
	$\Delta_{\text{fus}}H$		38.67	511		[1994REG/CHM]
C ₂₃ H ₃₀ O ₆	[50-04-4]	cortisone acetate				
	$\Delta_{\text{fus}}H$		38.43	509		[1994REG/CHM]
C ₂₃ H ₃₁ NO	[164667-96-3]	4- <i>n</i> -octyloxy-N-(3,5-dimethylbenzylidene)aniline				
	$\Delta_{\text{fus}}H$		37.73	324.7		[1995MIY/NAK]

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound					
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference	
C ₂₃ H ₃₁ NO ₃	[164667-97-4] $\Delta_{\text{fus}}H$	4- <i>n</i> -octyloxy-N-(3,5-dimethoxybenzylidene)aniline	35.3	316.3		[1995MIY/NAK]	
C ₂₃ H ₃₂ O ₂	[119-47-1] $\Delta_{\text{fus}}H$	3,3'-di- <i>tert</i> -butyl-5,5'-dimethyl-2,2-dihydroxydiphenylmethane	29.33	403.7	DTA	[1972INO/LIA]	
C ₂₃ H ₃₂ O ₃	[27811-56-9] $\Delta_{\text{fus}}H$	estra-1,3,5(10)-triene-3-ol-17 β pentanoate	29.45	420.7	DSC	[1986DEM/MAS]	
C ₂₃ H ₃₂ O ₄	[56-47-3] $\Delta_{\text{fus}}H$	deoxycorticosterone acetate	29.66	430		[1994REG/CHM]	
C ₂₃ H ₃₂ O ₆	[50-03-3] $\Delta_{\text{fus}}H$ $\Delta_{\text{fus}}H$	hydrocortisone acetate	53.64	480		[1997CEN/MEL]	
			36.95	496		[1994REG/CHM]	
C ₂₃ H ₃₄ O ₂	[2566-90-7] Δ_vH	Z,Z,Z,Z,Z,Z 4,7,10,13,16,19-docosaheptaeneoate	131.8 ± 0.2	298	CGC	[2007LIP/KAP]	
C ₂₃ H ₃₄ O ₃	[3410-54-6] $\Delta_{\text{fus}}H$	testosterone butyrate	24.75	382		[1994REG/CHM]	
C ₂₃ H ₃₅ N ₃ O ₈	[53848-86-5] $\Delta_{\text{us}}H$ $\Delta_{\text{fus}}H$	hexadecyl 2,4,6-trinitrobenzoate	18.6	349.3			
			29.54	393.3	DSC	[1974WAR/WIL]	
C ₂₃ H ₃₆ N ₂ O ₂	[98319-26-7] $\Delta_{\text{us}}H$ (I) $\Delta_{\text{fus}}H$ (I) $\Delta_{\text{fus}}H$ (II) $\Delta_{\text{sub}}H$	(5 α ,17 β)-N-(1,1-dimethylethyl)-3-oxo-4-azaandrost-1-ene-17-carboxamide (finasteride)	4.1	503.2			
			33.2	530.2			
			32.8	530.2	DSC	[2000WEN/BAU]	
			(463–488)	143.7	TGA	[1997ELD]	
C ₂₃ H ₃₇ BrN ₂ O ₄	[138317-09-6] $\Delta_{\text{fus}}H$	(4-nitrophenyl)-16-bromohexadecyl carbamate	62.62	382.9	DSC	[1993TIE/FRA]	
C ₂₃ H ₃₉ N ₃ O ₃	[6313-97-9] $\Delta_{\text{fus}}H$	1-hexadecyl-3-(4-nitrophenyl) urea	53.94	392.6	DSC	[1993TIE/FRA]	
C ₂₃ H ₄₀	[14752-75-1] Δ_vH	heptadecylbenzene (414–664)	98.5	429		[1999DYK/SVO]	
C ₂₃ H ₄₂ O ₂	[61012-47-3] Δ_vH	methyl Z,Z 13,16-docosadienoate	127.9	298	CGC	[2007LIP/KAP]	
C ₂₃ H ₄₂ O ₃	[5420-17-7] Δ_vH	tetrahydrofurfuryl oleate (353–398)	98.7	368	A	[1987STE/MAL]	
C ₂₃ H ₄₄	[95115-76-7] $\Delta_{\text{fus}}H$	<i>trans</i> -2-heptyl-6-hexyldecalin	38.91	312.2		[1985VAR/BRI]	
C ₂₃ H ₄₄	[95115-79-0] $\Delta_{\text{fus}}H$	<i>trans</i> -2-pentyl-6-octyldecalin	43.51	314.2		[1985VAR/BRI]	
C ₂₃ H ₄₄ O ₂	[1120-34-9] Δ_vH Δ_vH Δ_vH	methyl erucate	125.6 ± 1.2	298	CGC	[2007LIP/KAP]	
			(463–523)	123.8	298	GC	[1997KRO/VEL]
			(453–543)	93.5	498	GC	[1993HUS/SAR]
C ₂₃ H ₄₄ O ₅	[820-17-7] $\Delta_{\text{fus}}H$	1-aceto-3-stearin	41.69	319.9		[1996DOM/HEA]	

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound					
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference	
C ₂₃ H ₄₅ NO ₃	[na] $\Delta_v H$	2-lauryloxy-N,N-dibutylpropionamide (443–458)	90.6	450	A	[1987STE/MAL]	
C ₂₃ H ₄₆	[55124-77-1] $\Delta_v H$	9-cyclohexylheptadecane (456–492)	83.9	471	A	[1987STE/MAL]	
C ₂₃ H ₄₆	[19781-73-8] $\Delta_v H$	hexadecylcyclohexane (414–664)	97.6	429		[1999DYK/SVO]	
C ₂₃ H ₄₆	[18835-32-0] $\Delta_v H$	1-tricosene (409–652)	98.5	424		[1999DYK/SVO]	
C ₂₃ H ₄₆ O	[540-09-0] $\Delta_{\text{fus}} H$	12-tricosanone	78.03	342.2		[1993RUE/SAR]	
C ₂₃ H ₄₆ O ₂	[929-77-1] $\Delta_{\text{fus}} H$	methyl docosanoate (methyl behenate)	83.5	327.2	DSC	[2004CHI/ZHA]	
	$\Delta_{\text{fus}} H$		82.3	327.2		[1936KIN/GAR]	
	$\Delta_v H$		(463–513)	126.0 ± 0.3	298	GC	[2006HAF/PAR]
	$\Delta_v H$		(467–558)	126.1 ± 2.5	298	CGC	[2004CHI/ZHA]
	$\Delta_v H$		(463–523)	126.1	298	GC	[1997KRO/VEL]
	$\Delta_v H$		(453–543)	81.0	498		[1993HUS/SAR]
C ₂₃ H ₄₆ O ₂	[2433-96-7] $\Delta_{\text{us}} H$	tricosanoic acid	2.5	349.9			
	$\Delta_{\text{fus}} H$		75.0	352	DSC	[2007GBA/NEG]	
C ₂₃ H ₄₆ O ₃	[102542-57-4] $\Delta_v H$	decyl 3-decyloxypropionate (453–523)	90.2	468	A	[1987STE/MAL]	
C ₂₃ H ₄₈	[55124-79-3] $\Delta_v H$	9-hexylheptadecane (450–486)	82.6	465	A	[1987STE/MAL]	
C ₂₃ H ₄₈	[1560-81-2] $\Delta_v H$	2-methyldocosane (495–652)	79.7	510	A	[1987STE/MAL]	
C ₂₃ H ₄₈	[25117-30-0] $\Delta_v H$	4-methyldocosane (493–643)	76.3	508	A	[1987STE/MAL, 1999DYK/SVO, 1959TER/BRI]	
C ₂₃ H ₄₈	[25163-52-4] $\Delta_v H$	5-methyldocosane (492–644)	75.6	507	A	[1987STE/MAL, 1999DYK/SVO, 1959TER/BRI]	
C ₂₃ H ₄₈	[638-67-5] $\Delta_{\text{us}} H$	tricosane	<0.3	310.5			
	$\Delta_{\text{us}} H$		19.6	312.4			
	$\Delta_{\text{us}} H$		<0.3	317.2			
	$\Delta_{\text{fus}} H$		52.6	320.2	DSC	[2004MON/RAJ]	
	$\Delta_{\text{us}} H$		21.76	313.7			
	$\Delta_{\text{fus}} H$		53.97	320.7		[1990DOM/HEA]	
	$\Delta_{\text{sub}} H$		U 146.8 ± 10	298	B	[1991PIA/POM]	
	$\Delta_v H$		(412–462)	93.5	427		[2006SAW/MOK]
	$\Delta_v H$		(434–539)	117	298	CGC	[2004CHI/HAN]
	$\Delta_v H$			118.7 ± 0.1	298	GS	[2001PUR/CHI]
$\Delta_v H$		119.7 ± 2.3	298	CGC	[2000NIC/ORF]		
$\Delta_v H$		120.5 ± 2.0	298	CGC	[1991DIK/KAB]		

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound					
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference	
		$\Delta_v H$	(370–416)	123 ± 1	393	TE	[1994PIA/FON]
		$\Delta_v H$	(370–490)	92	385	TE,ME,GS	[1991PIA/POM]
		$\Delta_v H$	(314–353)	110.4	329	A	[1987STE/MAL]
		$\Delta_v H$	(440–653)	94	455	A, EST	[1987STE/MAL, 1966KUD/ZWO]
C ₂₃ H ₄₈ O ₂	[95491-58-0]	1,23-tricosanediol					
		$\Delta_{\text{us}}H$		41.8	366.3		
		$\Delta_{\text{fus}}H$		46.5	380.7	DSC	[1999OGA/NAK]
C ₂₃ H ₄₈ S	[66375-01-7]	1-tricosanethiol					
		$\Delta_v H$	(444–690)	110.1	459	EST	[1999DYK/SVO]
C ₂₄ F ₅₀	[1766-41-2]	perfluorotetracosane					
		$\Delta_{\text{us}}H$		3.89	202.7		
		$\Delta_{\text{fus}}H$		100.8	465.2	DSC	[1986STA]
C ₂₄ H ₁₂	[191-07-1]	coronene					
		$\Delta_{\text{fus}}H$		21.2	709	DSC	[2009TOR/CAM]
		$\Delta_{\text{fus}}H$		19.2	710.5		[1991ACR]
		$\Delta_{\text{sub}}H$	(473–483)	126.6 ± 1.7	478	ME	[2009TOR/CAM]
		$\Delta_{\text{sub}}H$	(473–483)	131.0 ± 1.7	298	ME	[2009TOR/CAM]
		$\Delta_{\text{sub}}H$	(421–504)	133.1 ± 5.1	463	ME	[1998OJA/SUU]
		$\Delta_{\text{sub}}H$	(313–453)	143.2	383	GS	[1995NAS/LEN]
		$\Delta_{\text{sub}}H$	(427–510)	135.9	468	ME	[1987STE/MAL, 1974MUR/POL]
		$\Delta_{\text{sub}}H$		128.4		ME	[1967WAK/INO]
		$\Delta_{\text{sub}}H$	(433–513)	147	473		[1958HOY/PEP]
		$\Delta_{\text{sub}}H$	(476–555)	143.2	407	ME	[1952INO/SHI]
		$\Delta_{\text{sub}}H$		148.5	407	ME	[1951INO]
		$\Delta_v H$		148.0 ± 0.5	298	CGC	[2002CHI/WEB]
		$\Delta_v H$	(323–473)	104.2	398	GC	[2002LEI/CHA]
C ₂₄ H ₁₂	[102234-01-5]	<i>bis</i> -benzo[3,4]cyclobuta[1,2-a:1',2'-c]biphenylene ([4]phenylene)					
		$\Delta_{\text{sub}}H$		131.0 ± 4.2			[2000BEC/FAU]
C ₂₄ H ₁₂ O ₂	[3302-52-1]	3,4:9,10-dibenzpyrene-5,8-quinone					
		$\Delta_{\text{sub}}H$		112.5 ± 5.4			[1956MAG, 1970COX/PIL]
C ₂₄ H ₁₄	[192-65-4]	dibenzo[a,e]pyrene					
		$\Delta_{\text{fus}}H$		32.1	517.9	DSC	[2010KES/AUC]
		$\Delta_{\text{fus}}H$		30.5	520.2		[1991ACR]
		$\Delta_{\text{sub}}H$	(414–506)	146.4	429	A	[1987STE/MAL]
		$\Delta_{\text{sub}}H$	(434–526)	137.6	480	ME	[1967WAK/INO]
C ₂₄ H ₁₄	[192-51-8]	dibenzo[fg,op]naphthacene					
		$\Delta_{\text{sub}}H$	(430–555)	147.4	445	A	[1987STE/MAL]
		$\Delta_{\text{sub}}H$	(454–526)	146.9	490	ME	[1967WAK/INO]
		Note: called 1,2,6,7-dibenzpyrene in paper, which we have taken to be dibenzo[fo,op]naphthacene based upon the melting point temperature reported in the paper					
C ₂₄ H ₁₄	[191-30-0]	dibenzo[a,l]pyrene					
		$\Delta_{\text{fus}}H$		24.68	501.2		[1991ACR]
C ₂₄ H ₁₄	[189-55-9]	benzo[<i>rst</i>]pentaphene					
		$\Delta_{\text{fus}}H$		27.87	556.8		[1991ACR]
C ₂₄ H ₁₄	[5385-75-1]	dibenzo[a,e]fluoranthene					
		$\Delta_{\text{fus}}H$		24.0	505.5	DSC	[2010KES/AUC]

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₂₄ H ₁₄ S ₆	[88493-55-4]	2,2',5',2'',5'',2''',5''',2''',5''',2''''-sexithiophene				
	$\Delta_{\text{sub}}H$	(503–563)	207.1		ME	[1998KLO/LAU]
	$\Delta_{\text{sub}}H$	(543–573)	211.3		ME	[1998KLO/LAU]
C ₂₄ H ₁₆ N ₂ O ₂	[1806-34-4]	2,2'-(1,4-phenylene)bis(5-phenyl)oxazole				
	$\Delta_{\text{sub}}H$		140	480	Optical	[1989SCH/PEN]
	$\Delta_{\text{sub}}H$	(600–680)	94.4	615	A	[1987STE/MAL, 1975STE/SCH]
C ₂₄ H ₁₈	[1165-14-6]	1,2,3-triphenylbenzene				
	$\Delta_{\text{sub}}H$	(373–395)	131.7 ± 0.6	384	ME	[2010RIB/SAN2]
	$\Delta_{\text{sub}}H$	(373–395)	134.1 ± 1.1	298	ME	[2010RIB/SAN2]
C ₂₄ H ₁₈	[612-71-5]	1,3,5-triphenylbenzene				
	$\Delta_{\text{fus}}H$		32.6	445.2	DSC	[1997VER2]
	$\Delta_{\text{fus}}H$		33.4	446		[1982LEB/BYK]
	$\Delta_{\text{sub}}H$	(407–429)	141.2 ± 0.7	418	ME	[2006RIB/MON]
	$\Delta_{\text{sub}}H$	(407–429)	147.8 ± 0.7	298	ME	[2006RIB/MON]
	$\Delta_{\text{sub}}H$		150.9	298	CGC-DSC	[1998CHI/HES]
	$\Delta_{\text{sub}}H$	(364–388)	145.6 ± 0.9	376	T	[1997VER2]
	$\Delta_{\text{sub}}H$		150.3 ± 0.9	298		[1997VER2]
	$\Delta_{\text{sub}}H$		152 ± 0.3	298	C,ME	[1974MAL/BAR]
	$\Delta_{\text{sub}}H$	(410–444)	142	425	ME	[1974MAL/BAR, 1987STE/MAL]
	$\Delta_{\text{sub}}H$	(384–400)	142.2	422	ME	[1967WAK/INO]
	$\Delta_{\text{sub}}H$		149.7 ± 4.1	298	ME	[1958HOY/PEP, 1970COX/PIL]
	Δ_vH		133.4 ± 2.0	298	GCG	[2008HAN/NUT]
	Δ_vH		140	298	CGC	[1998CHI/HES]
Δ_vH	(500–735)	77.5	515	A	[1987STE/MAL, 1962VOH/KAN]	
Δ_vH	(454–500)	118	469	A	[1987STE/MAL, 1974MAL/BAR]	
C ₂₄ H ₁₈	[135-70-6]	<i>p</i> -quaterphenyl				
	$\Delta_{\text{trs}}H$		0.41	233		
	$\Delta_{\text{fus}}H$		37.8	587.2		[1985SAI/ATA, 1991ACR]
	Δ_vH		136.1 ± 1.6	298	CGC	[2008HAN/NUT]
C ₂₄ H ₁₈ N ₂ S ₂	[109538-09-7]	4,4'-bis-(2-thienylmethylidamino)- <i>trans</i> -stilbene				
	$\Delta_{\text{trs}}H$ (liq <i>cryst</i>)		44.9	567.2		
	$\Delta_{\text{trs}}H$ (liq <i>cr-liq</i>)		0.2	580.2		[1978KOS/BUD]
C ₂₄ H ₁₈ N ₂ S ₂	[109538-10-5]	1,2-bis-[5-(β -azastyryl)-2-thienyl]- <i>trans</i> -ethylene				
	$\Delta_{\text{fus}}H$		45.9	501.2		[1978KOS/BUD]
C ₂₄ H ₂₀ O ₆	[614-33-5]	glycerol tribenzoate				
	Δ_vH	(423–476)	123.5	438	A, T	[1987STE/MAL, 1949FOR/NOR]
C ₂₄ H ₂₁ NO ₂	[1041479-16-6]	2- <i>tert</i> -butylphenyl acridine-9-carboxylate				
	$\Delta_{\text{fus}}H$		39.8	462	DSC	[2010KRZ/MAL]
C ₂₄ H ₂₄ N ₂ O ₄	[na]	4-(methoxymethyl)-6-(phenylmethoxy)-9 <i>H</i> -pyrido[3,4- <i>b</i>]indole-3-carboxylic acid, 1-methylethyl ester				
	$\Delta_{\text{fus}}H$		38.83	424		[1999WIN]
C ₂₄ H ₂₄ O ₄	[89702-41-0]	syn 4,9-bis(methoxycarbonyl)pagodane (dimethyl undecacyclo[9.9.0.0 ^{1,5} .0 ^{2,12} .0 ^{2,18} .0 ^{3,7} .0 ^{6,10} .0 ^{8,12} .0 ^{11,15} .0 ^{13,17} .0 ^{16,20}]-eicosane-4- <i>syn</i> , 9- <i>syn</i> -dicarboxylate)				
	$\Delta_{\text{sub}}H$	(393–447)	146.1 ± 3.0	420	T	[1994BEC/RUE]
C ₂₄ H ₂₄ O ₄	[124316-65-0]	1,6-bis(methoxycarbonyl)dodecahedrane (dimethyl undecacyclo[9.9.0.0 ^{2,9} .0 ^{3,7} .0 ^{4,20} .0 ^{5,18} .0 ^{6,16} .0 ^{8,15} .0 ^{10,14} .0 ^{12,19} .0 ^{13,17}]-eicosane-1,6-dicarboxylate)				
	$\Delta_{\text{sub}}H$	(395–450)	139.7 ± 1.3	422	T	[1994BEC/RUE]

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₂₄ H ₂₅ F ₂₅	[89109-71-7]	1,1,1,2,2,3,3,4,4,5,5,6,6,7,7,8,8,9,9,10,10,11,11,12,12-pentacosuofluorotetracosane				
	$\Delta_{\text{trs}}H$		14.6	351		
	$\Delta_{\text{fus}}H$		23.2	361	DSC	[2008NUN/CLA]
	$\Delta_{\text{trs}}H$		10.0	352.1		
	$\Delta_{\text{fus}}H$		26.0	364.1	DSC	[1992HOP/MOL]
	$\Delta_{\text{fus}}H$		11.3	353.2		
C ₂₄ H ₂₅ F ₂₅	[116177-49-2]	1,1,1,2,2,3,3,4,4,5,5,6,6,7,7,8,8,9,9,10,10,11,11,12,12-pentacosuofluoro-14-methyltricosane				
	$\Delta_{\text{trs}}H$		9	220		
	$\Delta_{\text{fus}}H$		25	347.1	DSC	[1992HOP/MOL]
C ₂₄ H ₂₆ N ₂ O ₂	[14580-70-2]	1,5-dipiperidylantraquinone				
$\Delta_{\text{sub}}H$	(408–458)	173.3	428		[1958HOY/PEP, 1987STE/MAL]	
C ₂₄ H ₂₆ N ₂ O ₄	[72956-09-3]	1-(9 <i>H</i> -carbazol-4-yloxy)-3-[[2-(2-methoxyphenoxy)ethyl]amino-2-propanol (carvedilol)				
$\Delta_{\text{fus}}H$		57.6	387.3	DSC	[2007BER/WAS]	
C ₂₄ H ₂₇ NO ₄	[3088-05-9]	<i>bis</i> [N,N-(2-hydroxy-3-phenoxy)propyl]phenylamine				
Δ_vH	(388–423)	131	403	A	[1987STE/MAL]	
C ₂₄ H ₂₇ NO ₅ S	[97322-87-7]	5-[[4[(3,4-dihydro-6-hydroxy-2,5,7,8-tetramethyl-2 <i>H</i> -1-benzopyran-2-yl)methoxy]phenyl]methyl-2,4-thiazolidinedione (troglitazone)				
$\Delta_{\text{fus}}H$		48.8	412.4	DSC	[2007BER/WAS]	
C ₂₄ H ₂₈ FN ₃ O	[265667-22-9]	N-methyl-[1-[1-(2-fluorophenethyl)piperidin-4-yl]-1 <i>H</i> -indol-6-yl]acetamide				
	$\Delta_{\text{fus}}H$ (I)		38.2	421.3		
	$\Delta_{\text{fus}}H$ (II)		35.2	413		[2002KUS/ASH]
C ₂₄ H ₂₈ P ₂ O ₂	[7688-25-7]	1,4- <i>bis</i> (diphenylphosphino)butane				
$\Delta_{\text{sub}}H$		171.6	443	B	[1989HUI/VAN]	
C ₂₄ H ₂₉ NO ₃	[126675-77-2]	2-(4-nitrophenyl)-1-[4-(<i>trans</i> -4-butylcyclohexyl)phenyl]ethanone				
$\Delta_{\text{fus}}H$		36.4	426.9	DSC	[2002SPA/DZI]	
C ₂₄ H ₂₉ NO ₆	[836602-52-9]	propyl naltrexone-3- <i>O</i> -carbonate				
$\Delta_{\text{fus}}H$		20.97	379.2	DSC	[2004PIL/HAM]	
C ₂₄ H ₂₉ NO ₆	[836602-53-0]	isopropyl naltrexone-3- <i>O</i> -carbonate				
$\Delta_{\text{fus}}H$		26.62	427.2	DSC	[2004PIL/HAM]	
C ₂₄ H ₃₀	[59358-71-3]	1,1-diphenyl-1,1'-bicyclohexyl				
	$\Delta_{\text{fus}}H$		29.71	455	DSC	[1983KRA/BEC]
	$\Delta_{\text{sub}}H$		150.2		E,B	[1983KRA/BEC]
C ₂₄ H ₃₀ O ₄	[140-24-9]	dibenzyl sebacate				
	Δ_vH	(368–550)	114.3	383	A	[1987STE/MAL]
	Δ_vH	(405–463)	112.2	420	T	[1949PER/WEB]
	Δ_vH	(373–432)	121	388	T	[1939VER/MAR]
C ₂₄ H ₃₀ O ₄	[167321-36-0]	2,2'-diphenyl-bi-(5,5-dimethyl-1,3-dioxan-2-yl)				
$\Delta_{\text{sub}}H$	(372–420)	130.2 ± 1.8	396	T	[1995VER/DOG]	
C ₂₄ H ₃₁ FO ₆	[76-25-5]	triamcinolone acetonide				
$\Delta_{\text{fus}}H$		45.29	566		[1994REG/CHM]	
C ₂₄ H ₃₁ FO ₆	[1177-87-3]	dexamethasone acetate				
$\Delta_{\text{fus}}H$		37.72	503		[1994REG/CHM]	
C ₂₄ H ₃₂	[4384-23-0]	[6.6]-para-cyclophane				
	$\Delta_{\text{sub}}H$	(352–371)	108.8 ± 0.8			[1969SHI/MCN, 1977PED/RYL]

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{sub}}H$	(352–371)	115.1 ± 2.1	298		[1969SHI/MCN, 1977PED/RYL]
C ₂₄ H ₃₂	[115181-13-0]	8-[4-(4'- <i>n</i> -butylbiphenyl)]-1-octene				
	$\Delta_{\text{trs}}H$ (liq <i>cryst</i>)		2.2	248.6		
	$\Delta_{\text{trs}}H$ (liq <i>cr-liq</i>)		9.6	315.6	DSC	[1989MAL/KAN]
C ₂₄ H ₃₂ O ₃	[63058-78-6]	3-[(1-oxohexyl)oxy]-estra-1,3,5(10)-trien-17-one				
	$\Delta_{\text{fus}}H$		23	370	DSC	[1990YAN/EIR]
C ₂₄ H ₃₂ O ₄	[na]	2,2'-diphenyl-bi(5,5-dimethyl-1,3-dioxan-2-yl)				
	$\Delta_{\text{fus}}H$		49.8	507.1		[1995VER/DOG]
C ₂₄ H ₃₂ O ₄ S	[52-01-7]	17-hydroxy-7 α -mercapto-3-oxo-17 α -pregn-4-ene-21-carboxylic acid, γ -lactone, acetate				
	$\Delta_{\text{fus}}H$ (II)		22.9	480	DSC	[2007ESP/NIC]
	$\Delta_{\text{fus}}H$ (I)		20	478		
	$\Delta_{\text{fus}}H$ (II)		22.1	483		[1991AGA/LEG]
C ₂₄ H ₃₂ O ₈	[14174-09-5]	dibenzo[24-crown-8]				
	$\Delta_{\text{fus}}H$		62.5			[2002LEB/BYK, 2004BYK/LEB]
	$\Delta_{\text{trs}}H$		16.6	354.1		
	$\Delta_{\text{fus}}H$		52.25	375.4		[1998DOM, 1985RAE/SOL]
C ₂₄ H ₃₄	[1603-53-8]	1,1-diphenyldodecane				
	$\Delta_{\text{trs}}H$		1.92	191		
	$\Delta_{\text{fus}}H$		38.83	281.4		[1996DOM/HEA]
C ₂₄ H ₃₄ N ₄ O ₅ S	[93479-97-1]	3-ethyl-2,5-dihydro-4-methyl-N-[2-[4-[[[trans-4-methylcyclohexyl]amino]carbonyl]amino]sulfonyl]phenyl]ethyl-2-oxo-1 <i>H</i> -pyrrole-1-carboxamide (glimepiride)				
	$\Delta_{\text{fus}}H$		53.3	485.7	DSC	[2007BER/WAS]
C ₂₄ H ₃₆ O ₃	[3129-43-9]	testosterone valerate				
	$\Delta_{\text{fus}}H$		24.57	380		[1994REG/CHM]
C ₂₄ H ₃₆ O ₅	[75330-75-5]	2-methylbutanoic acid, (1 <i>S</i> ,3 <i>R</i> ,7 <i>S</i> ,8 <i>S</i> ,8 <i>aR</i>)-1,2,3,7,8,8 <i>a</i> -hexahydro-3,7-dimethyl-8-[2-[(2 <i>R</i> ,4 <i>R</i>)-tetrahydro-4-hydroxy-6-oxo-2 <i>H</i> -pyran-2-yl]ethyl]-1-naphthalenyl ester (lovastatin)				
	$\Delta_{\text{fus}}H$		36.53	444.3	DSC	[2008TUN/TAB]
	$\Delta_{\text{fus}}H$		43.14	445.5	DSC	[2008NTU/CHM]
C ₂₄ H ₃₇ N ₃ O	[218765-43-6]	pyrimethanil laurate				
	$\Delta_{\text{fus}}H$	(78–340)	67.24	321.5	AC	[2004SUN/LIU]
C ₂₄ H ₃₈ O ₄	[117-81-7]	<i>bis</i> (2-ethylhexyl) phthalate				
	Δ_vH	(373–660)	102.5	388	A	[1987STE/MAL]
	Δ_vH	(385–440)	110.7	390	T	[1949PER/WEB]
C ₂₄ H ₃₈ O ₄	[131-15-7]	<i>bis</i> (1-methylheptyl) phthalate				
	Δ_vH	(393–435)	93.1	408	A	[1987STE/MAL]
C ₂₄ H ₃₈ O ₄	[131-20-4]	<i>bis</i> (6-methylheptyl) phthalate				
	Δ_vH	(383–490)	92.4	398	A	[1987STE/MAL]
C ₂₄ H ₃₈ O ₄	[117-84-0]	dioctyl phthalate				
	Δ_vH	(423–523)	99.5	438	A	[1987STE/MAL]
	Δ_vH	(383–433)	107.6	398	T	[1949PER/WEB]
C ₂₄ H ₄₀	[62155-50-4]	1-cyclohexyl-1-phenyldodecane				
	$\Delta_{\text{fus}}H$		35.19	275.8		[1996DOM/HEA]
C ₂₄ H ₄₀ N ₈ O ₄	[58-32-2]	2,2',2'',2'''-[(4,8-di-1-piperidinylpyrimido[5,4- <i>d</i>]pyrimidine-2,6-diyl)-dinitrilo]- <i>tetrakis</i> -ethanol (dipyridamole)				
	$\Delta_{\text{fus}}H$ (I)		44.05	442.8		

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{fus}}H$ (II)		33.2	441.9		[2006ADH/BAS]
C ₂₄ H ₄₀ O ₃	[na] $\Delta_{\text{fus}}H$	5-(1,1-dimethylheptyl)-2-[(1R,2R,5R)-5-hydroxy-2-(3-hydroxypropyl)-cyclohexyl]phenol	18.4	357		[2004VAL/KIP]
C ₂₄ H ₄₀ O ₄	[175848-64-3] $\Delta_{\text{ms}}H$ $\Delta_{\text{ms}}H$ $\Delta_{\text{fus}}H$	2,5-di- <i>n</i> -nonyloxy-1,4-benzoquinone	8.0 24.2 47.1	352.6 383.8 402.7	DSC	[1996KEE/VAN]
C ₂₄ H ₄₂	[2456-68-0] Δ_vH	hexapropylbenzene (458–606)	68.4	473	A	[1987STE/MAL, 1937GRO/IPA]
C ₂₄ H ₄₂	[4445-07-2] Δ_vH	octadecylbenzene (423–675)	101	438		[1999DYK/SVO]
C ₂₄ H ₄₂ O ₆	[64617-30-7] Δ_vH	<i>trans</i> trihexyl aconitate (423–512)	98.2	438	A	[1987STE/MAL]
C ₂₄ H ₄₂ O ₁₁	[na] Δ_vH	di[1-(2-ethylbutyloxycarbonyl)ethyl] diethylene glycol dicarboxylate (448–538)	110.1	463	A	[1987STE/MAL]
C ₂₄ H ₄₂ O ₁₁	[na] Δ_vH	di[1-(2-hexyloxycarbonyl)ethyl] diethylene glycol dicarboxylate (443–548)	111	458	A	[1987STE/MAL]
C ₂₄ H ₄₄	[na] Δ_vH	9-decyltetradecahydroanthracene (501–536)	103.2	516	A	[1987STE/MAL]
C ₂₄ H ₄₄	[na] Δ_vH	9-decyltetradecahydrophenanthrene (502–542)	92.0	517	A	[1987STE/MAL]
C ₂₄ H ₄₄ O ₂	[na] $\Delta_{\text{fus}}H$	3,3,7,7,11,11,15,15-octamethylcyclohexadecane-1,9-dione	34.3	423.2		[1972BOR/DAL2]
C ₂₄ H ₄₄ O ₄	[na] $\Delta_{\text{fus}}H$	1,11-cycloicosanedione <i>bis</i> ethylene ketal	43.72	362.2		[1972ALV/BOR]
C ₂₄ H ₄₄ O ₆	[140-04-5] Δ_vH	O-acetylricinoleic acid, butyl ester (378–423)	105.2	393	A	[1987STE/MAL]
C ₂₄ H ₄₄ O ₆	[38094-13-2] Δ_vH	trihexyl 1,2,3-propanetricarboxylate (422–526)	98.1	437	A	[1987STE/MAL]
C ₂₄ H ₄₄ O ₆	[620-67-7] Δ_vH	glycerol triheptanoate (401–452)	84.4	416		[2001BUR/JOS]
C ₂₄ H ₄₆	[18254-57-4] $\Delta_{\text{fus}}H$	1,1-dicyclohexyldodecane	44.35	300.6		[1996DOM/HEA]
C ₂₄ H ₄₆	[95746-44-4] $\Delta_{\text{fus}}H$	2,11-dicyclohexyldodecane	43.93	300.6		[1996DOM/HEA]
C ₂₄ H ₄₆	[95115-77-8] $\Delta_{\text{fus}}H$	<i>trans</i> -2,6-diheptyldecalin	40.17	326.7		[1985VAR/BRI]
C ₂₄ H ₄₆ O ₄	[20270-50-2] Δ_vH	<i>bis</i> (3,5,5-trimethylhexyl)adipate (353–413)	107.6	368	A, ME	[1987STE/MAL, 1948SMA/SMA]
C ₂₄ H ₄₈	[4445-06-1] Δ_vH	octadecylcyclohexane (422–675)	100.3	437		[1999DYK/SVO]
C ₂₄ H ₄₈	[10192-32-2] Δ_vH	1-tetracosene (418–663)	101	433		[1999DYK/SVO]

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound						
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference		
C ₂₄ H ₄₈	[297-03-0]	cyclotetracosane						
	$\Delta_{\text{us}}H$		38.0	297				
	$\Delta_{\text{fus}}H$		10.8	322		[1987DRO/MOL]		
C ₂₄ H ₄₈ O ₂	[5908-87-2]	ethyl docosanoate						
	$\Delta_{\text{us}}H$		9.58	312.3				
	$\Delta_{\text{fus}}H$		19.16	321		[1996DOM/HEA]		
	$\Delta_{\text{fus}}H$		77.82	321		[1967OMA]		
	$\Delta_{\text{sub}}H$	(313–318)	196.5	315.5	ME	[1987STE/MAL, 1967OMA]		
C ₂₄ H ₄₈ O ₂	[2433-97-8]	methyl tricosanoate (473–528)	99.8	488	A, EST	[1987STE/MAL, 1963ROS/SCH]		
							Δ_vH	(327–344)
C ₂₄ H ₄₆ Cl	[6422-18-0]	1-chlorotetracosane (543–774)	72.4	558	A	[1987STE/MAL, 1970DYK/VAN]		
							Δ_vH	
C ₂₄ H ₅₀	[1928-30-9]	2-methyltricosane (450–664)	89.3	465	A	[1987STE/MAL]		
							Δ_vH	
C ₂₄ H ₅₀	[22331-09-5]	5-methyltricosane (503–653)	79.6	518	A	[1987STE/MAL, 1999DYK/SVO, 1959TER/BRI]		
							Δ_vH	
C ₂₄ H ₅₀	[646-31-1]	tetracosane						
			$\Delta_{\text{us}}H$		33.18	321		
			$\Delta_{\text{fus}}H$		59.31	324.1	DSC	[2007HAF/MAH]
			$\Delta_{\text{us}}H$		30.3	319		
			$\Delta_{\text{us}}H$		<0.3	319.6		
			$\Delta_{\text{fus}}H$		53.8	323.4	DSC	[2004MON/RAJ]
			$\Delta_{\text{us}}H$		31.3	321.3		
			$\Delta_{\text{fus}}H$		54.89	324.1		[1996DOM/HEA]
			$\Delta_{\text{sub}}H$	(308–323)	164.9 ± 1.8	315		[2009RAZ/NAC]
			$\Delta_{\text{sub}}H$		162 ± 12	298	B	[1991PIA/POM]
			Δ_vH	(333–373)	105.1 ± 0.5	353		[2009RAZ/NAC]
			Δ_vH	(334–452)	112	349	GC	[2007LEE/LAI]
			Δ_vH	(434–539)	121.9	298	CGC	[2004CHI/HAN]
			Δ_vH		126.8 ± 0.4	298	CGC	[2002CHI/WEB]
			Δ_vH		125.7 ± 1.6	298	CGC	[2000NIC/ORF]
			Δ_vH		126.2 ± 2.3	298	CGC	[1997CHI/WIL]
			Δ_vH	(453–588)	92.6	468		[1994MOR/KOB]
			Δ_vH	(386–425)	126 ± 2	405	TE	[1994PIA/FON]
			Δ_vH	(382–523)	95.2	397	TE,ME,GS	[1991PIA/POM]
Δ_vH	(451–497)	86.2 ± 4.6	474	GS	[1990PIA/SCA]			
Δ_vH	(373–463)	111.2	388		[1988SAS/JOS]			
Δ_vH	(498–573)	86.6	513	A, EST	[1987STE/MAL, 1966KUD/ZWO]			
C ₂₄ H ₅₀	[22331-52-8]	12-methyltricosane (435–454)	84.5	445	GC	[1982REC/GRE, 1999DYK/SVO]		
							Δ_vH	
C ₂₄ H ₅₀ O	[4543-57-8]	13-oxapentacosane						
			$\Delta_{\text{us}}H$		4.6	303.8		
			$\Delta_{\text{fus}}H$		92.88	304.62	DSC	[2004TYA/BIS]
C ₂₄ H ₅₀ O ₂	[2136-74-5]	2-(docosanoxy)ethanol						
			$\Delta_{\text{us}}H$		12.92	317.2		

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound					
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference	
		$\Delta_{\text{fus}}H$	43.93	335.9	DTA	[1979KUC/SKU]	
C ₂₄ H ₅₀ O ₂	[22513-82-2]	1,24-tetracosanediol					
		$\Delta_{\text{trs}}H$	42.7	372.7			
		$\Delta_{\text{fus}}H$	51.2	381.5	DSC	[1999OGA/NAK]	
C ₂₄ H ₅₀ O ₄ S ₂	[na]	(l)-rhamnose dinonyl dithioacetal					
		$\Delta_{\text{trs}}H$	24.7	342.1			
		$\Delta_{\text{fus}}H$	54.4	387.4	DSC	[1989VAN/VAN]	
C ₂₄ H ₅₀ S	[16331-24-1]	1-tetracosanethiol					
		(451–700)	112.2	466	EST	[1999DYK/SVO]	
C ₂₄ H ₅₁ N	[1116-76-3]	trioctylamine					
		(415–536)	110.4 ± 1.5	298	EB	[1996STE/CHI3]	
		(505–702)	70.6	520	A	[1987STE/MAL]	
C ₂₅ H ₂₀	[630-76-2]	tetraphenylmethane					
		$\Delta_{\text{fus}}H$	48.28	554.2	DSC	[1999VER3]	
		$\Delta_{\text{sub}}H$	(363–388)	140.0 ± 1.3	298	GS	[1999VER3]
		$\Delta_{\text{sub}}H$	(363–383)	135.4 ± 1.3	376	GS	[1999VER3]
		$\Delta_{\text{sub}}H$	(396–466)	150.6 ± 4	298	TE,ME	[1972KAN, 1977PED/RYL]
	$\Delta_{\text{sub}}H$	(404–466)	143.3	419		[1987STE/MAL]	
C ₂₅ H ₂₂ O ₁₀	[22888-70-6]	2-[(2R,3R)-2,3-dihydro-3-(4-hydroxy-3-methoxyphenyl)-2-(hydroxymethyl)-1,4-benzodioxin-6-yl]-2,3-dihydro-3,5,7-trihydroxy-4H-1-benzopyran-4-one (silybin)					
		$\Delta_{\text{fus}}H$	44.87	424	DSC	[2005ZHA/BAI, 2006BAI/YAN, 2005YAO/BAI]	
C ₂₅ H ₂₆	[55334-57-1]	3-phenylethyl-1,5-diphenyl-2-pentene					
		Δ_vH	(469–541)	86.8	484		[1999DYK/SVO]
C ₂₅ H ₂₈	[66374-88-7]	3-phenylethyl-1,5-diphenylpentane					
		Δ_vH	(498–542)	87.3	513	A	[1987STE/MAL, 1999DYK/SVO]
C ₂₅ H ₂₈ N ₆ O	[na]	irbesartan					
		$\Delta_{\text{sub}}H$ (α)	(403–421)	197.2 ± 20		ME	[2009TAU/SIT]
		$\Delta_{\text{sub}}H$ (β)	(403–421)	254.5 ± 20		ME	[2009TAU/SIT]
C ₂₅ H ₂₈ O ₃	[na]	estra-1,3,5(10)-triene-3,17-diol(17 β), 3-benzoate					
		$\Delta_{\text{fus}}H$	41.75	464.1		[1985DEM/CHA]	
C ₂₅ H ₃₁ FO ₈	[67-78-7]	triamcinolone diacetate					
		$\Delta_{\text{fus}}H$	38.31	508		[1994REG/CHM]	
C ₂₅ H ₃₁ NO ₃	[483362-62-5]	2-(4-nitrophenyl)-1-[4-[2-(trans-4-propylcyclohexyl)ethyl]phenyl]ethanone					
		$\Delta_{\text{fus}}H$	35.56	420	DSC	[2002SPA/DZI]	
C ₂₅ H ₃₁ NO ₃	[126675-78-3]	2-(4-nitrophenyl)-1-[4-(trans-4-pentylcyclohexyl)phenyl]ethanone					
		$\Delta_{\text{fus}}H$	34.48	415.8	DSC	[2002SPA/DZI]	
C ₂₅ H ₃₂ N ₂ O ₅	[na]	N,N-diethyl naltrexone-3-O-carbamate					
		$\Delta_{\text{fus}}H$	21.56	419.7	DSC	[2009VAD/BAN]	
C ₂₅ H ₃₂ O ₂	[na]	17-phenyl testosterone					
		$\Delta_{\text{trs}}H$	23.3	388.5			
		$\Delta_{\text{fus}}H$	44.2	450		[1997CEN/MEL]	
C ₂₅ H ₃₄ O ₃	[65445-09-2]	19-nor-17 α -ethynyl-17 β -(2,2-dimethylpropionyloxy)-4-androsten-3-one					
		$\Delta_{\text{fus}}H$	37.8	500		[1996DOM/HEA]	

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₂₅ H ₃₄ O ₃	[2985-59-3] $\Delta_v H$	2-hydroxy-4-dodecyloxybenzophenone (413–453)	115.5	433	ME	[1984SUR]
C ₂₅ H ₃₄ O ₃	[118924-66-6] $\Delta_{\text{fus}} H$	3-[(1-oxoheptyl)oxy]-estra-1,3,5(10)-trien-17-one	21	338	DSC	[1990YAN/EIR]
C ₂₅ H ₃₄ O ₆	[51333-22-3] $\Delta_{\text{fus}} H$	16 α ,17 α -(butylidenedioxy)-11 β ,21-dihydroxypregna-1,4-diene-3,20-dione (budesonide)	34.7	534	DSC	[2009MOT/CAR]
C ₂₅ H ₃₄ O ₈	[2203-97-6] $\Delta_{\text{fus}} H$	hydrocortisone hemisuccinate	41.34	444		[1997CEN/MEL]
C ₂₅ H ₃₆	[7225-70-9] $\Delta_v H$	1-phenyl-3-phenethylundecane (456–521)	91.9	471	A	[1987STE/MAL, 1999DYK/SVO]
C ₂₅ H ₃₆ O ₂	[119-47-1] $\Delta_{\text{sub}} H$	2,2'-methylenebis(6- <i>tert</i> -butyl-4-methylphenol) (383–403)	114	393	GS	[1971FEL/KUZ]
C ₂₅ H ₃₈	[55191-63-4] $\Delta_v H$ $\Delta_v H$	1-pentadecylnaphthalene (474–524) (474–540)	98.1 96.7	489 489	A A	[1987STE/MAL] [1987STE/MAL]
C ₂₅ H ₃₈ O ₅	[79902-63-9] $\Delta_{\text{fus}} H$ $\Delta_{\text{fus}} H$	2,2-dimethylbutanoic acid, (1S,3R,7S,8S,8aR)-1,2,3,7,8,8a-hexahydro-3,7-dimethyl-8-[2-[(2R,4R)-tetrahydro-4-hydroxy-6-oxo-2H-pyran-2-yl]ethyl]-1-naphthalenyl ester (simvastatin)	32.17 29.59	412.6 413.8	DSC DSC	[2009NTI/CHA] [2008TUN/TAB]
C ₂₅ H ₃₉ N ₃ O ₈	[53848-85-4] $\Delta_{\text{fus}} H$ $\Delta_{\text{fus}} H$	octadecyl 2,4,6-trinitrobenzoate 25.4 30.0	364.1 392.3		DSC	[1974WAR/WIL]
C ₂₅ H ₄₀	[55334-30-0] $\Delta_v H$	1-cyclohexyl-6-cyclopentyl-3-phenethylhexane (486–525)	87.7	501	A	[1987STE/MAL, 1999DYK/SVO]
C ₂₅ H ₄₀	[55334-31-1] $\Delta_v H$	1,7-dicyclopentyl-4-(2-phenethyl)heptane (487–525)	92.0	502	A, MG	[1987STE/MAL, 1955SCH/WHI, 1999DYK/SVO]
C ₂₅ H ₄₂	[334-29-2] $\Delta_v H$	1-hexadecylindane (495–536)	87.0	510	A, MG	[1987STE/MAL, 1955SCH/WHI, 1999DYK/SVO]
C ₂₅ H ₄₂	[66374-91-2] $\Delta_v H$	5-pentadecyl-1,2,3,4-tetrahydronaphthalene (471–534)	99.4	486	A	[1987STE/MAL, 1999DYK/SVO]
C ₂₅ H ₄₄	[29136-19-4] $\Delta_v H$	nonadecylbenzene (431–686)	103.6	446		[1999DYK/SVO]
C ₂₅ H ₄₄	[66374-92-3] $\Delta_v H$	1,5-dicyclohexyl-3-(2-cyclohexylethyl)-2-pentene (485–524)	88.0	500	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₂₅ H ₄₄	[66374-93-4] $\Delta_v H$	1,7-dicyclopentyl-4-(3-cyclopentylpropyl)-3-heptene (483–522)	89.0	498	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₂₅ H ₄₄	[5637-96-7] $\Delta_v H$	3-octyl-1-phenylundecane (476–513)	89.4	491	A	[1987STE/MAL]
C ₂₅ H ₄₄	[5637-96-7] $\Delta_v H$	9-(2-phenylethyl)heptadecane (448–513)	88.3	463	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₂₅ H ₄₄	[4445-08-3]	9-(4-tolyl)octadecane				

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(472–507)	92.0	487	A, MG	[1987STE/MAL, 1955SCH/WHI, 1999DYK/SVO]
C ₂₅ H ₄₄	[7225-65-2]	6-octyl(hexylhydrobenz[de]anthracene)				
	$\Delta_v H$	(467–534)	93.8	482		[1999DYK/SVO]
C ₂₅ H ₄₄ O ₈	[15834-04-5]	pentaerythritol tetrapentanoate				
	$\Delta_v H$	(334–413)	120.8	350		[2007RAZ/MOK]
C ₂₅ H ₄₆	[55401-70-2]	1-cyclohexyl-3-(cyclohexylethyl)-6-cyclopentylhexane				
	$\Delta_v H$	(487–524)	91.4	502	A	[1987STE/MAL]
C ₂₅ H ₄₆	[55401-72-4]	4-(2-cyclohexylethyl)-1,7-dicyclopentylheptane				
	$\Delta_v H$	(471–524)	88.8	486	A	[1987STE/MAL]
C ₂₅ H ₄₆	[2090-16-6]	1,5-dicyclohexyl-3-(2-cyclohexylethyl)pentane				
	$\Delta_v H$	(318–418)	107.6	333	A	[1987STE/MAL, 1964MOR]
	$\Delta_v H$	(488–528)	86.2	503	A	[1987STE/MAL]
C ₂₅ H ₄₆	[55429-35-1]	1,7-dicyclopentyl-4-(3-cyclopentylpropyl)heptane				
	$\Delta_v H$	(457–525)	87.6	472	A	[1987STE/MAL]
	$\Delta_v H$	(486–525)	88.9	501	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₂₅ H ₄₆ O ₆	[33599-07-4]	1,2-diaceto-3-stearin				
	$\Delta_{\text{fus}} H$		45.56	208.3		[1996DOM/HEA]
C ₂₅ H ₄₈	[7225-69-6]	1-cyclohexyl-3-(2-cyclohexylethyl)undecane				
	$\Delta_v H$	(480–516)	95.2	495	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₂₅ H ₄₈	[7225-68-5]	1-cyclopentyl-4-(3-cyclopentylpropyl)dodecane				
	$\Delta_v H$	(480–518)	88.5	495	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₂₅ H ₄₈	[55401-73-5]	1-hexyldecylhexahydroindane				
	$\Delta_v H$	(492–532)	87.6	507	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₂₅ H ₄₈	[66359-82-8]	1-pentadecyldecahydronaphthalene				
	$\Delta_v H$	(464–529)	93.4	479	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₂₅ H ₄₈ N ₆ O ₈	[70-51-9]	N'-[5-[[4-[[5-(acetylhydroxyamino)pentyl]amino]-1,4-dioxobutyl]-hydroxyamino]pentyl]-N-(5-aminopentyl)-N-hydroxybutanediamide (deferroxamine)				
	$\Delta_{\text{fus}} H$		105.3	411.1	DSC	[2000IHN/VEN]
C ₂₅ H ₄₈ O ₂	[2733-88-2]	methyl Z 15-tetracosenoate				
	$\Delta_v H$		135.3 ± 1.1	298	CGC	[2007LIP/KAP]
C ₂₅ H ₄₈ O ₄	[2064-80-4]	dioctyl nonanedioate				
	$\Delta_v H$	(393–523)	104.3	408	A	[1987STE/MAL]
C ₂₅ H ₅₀	[22349-03-7]	nonadecylcyclohexane				
	$\Delta_{\text{fus}} H$		79.9	315.7	DSC	[2001YOU/SCH]
	$\Delta_{\text{fus}} H$		78.8	313.2	DSC	[2000YOU/DOL]
	$\Delta_{\text{fus}} H$		77.79	316.2	DSC	[2000SIR/HER]
	$\Delta_{\text{fus}} H$		77.8	317		[1991MIY/ENO]
	$\Delta_v H$	(430–686)	102.8	445		[1999DYK/SVO]
C ₂₅ H ₅₀	[16980-85-1]	1-pentacosene				
	$\Delta_v H$	(426–674)	103.7	441		[1999DYK/SVO]
C ₂₅ H ₅₀	[25446-35-9]	9-(2-cyclohexylethyl)heptadecane				
	$\Delta_v H$	(490–513)	88.6	495		[1987STE/MAL, 1955SCH/WHI]
C ₂₅ H ₅₀	[5638-09-5]	9-(3-cyclopentylpropyl)heptadecane				

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound					
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference	
		$\Delta_v H$	(476–514)	86.9	491	A	[1987STE/MAL, 1999DYK/SVO]
C ₂₅ H ₅₀	[24306-18-1]	9-octyl-8-heptadecene					
		$\Delta_v H$	(441–500)	92.3	456	A	[1987STE/MAL, 1999DYK/SVO]
C ₂₅ H ₅₀ O	[2123-19-5]	13-pentacosanone					
		$\Delta_{\text{fus}} H$		96.17	347		[2000NAK/SHI]
C ₂₅ H ₅₀ O ₂	[2442-49-1]	methyl tetracosanoate					
		$\Delta_{\text{fus}} H$		90.0	331.2	DSC	[2004CHI/ZHA]
		$\Delta_v H$	(467–558)	136.6 ± 2.5		CGC	[2004CHI/ZHA]
		$\Delta_v H$	(422–452)	146.2	437		[2001BUR/JOS]
		$\Delta_v H$	(483–536)	100.8	498	A	[1987STE/MAL]
C ₂₅ H ₅₀ O ₂	[18281-07-7]	ethyl tricosanoate					
		$\Delta_{\text{fus}} H$		57.32	326		[1967OMA]
		$\Delta_{\text{sub}} H$	(316–322)	175.2	319	ME	[1987STE/MAL, 1967OMA]
		$\Delta_v H$	(336–359)	121.8	347	A, ME	[1987STE/MAL, 1967BER/WES]
C ₂₅ H ₅₂	[7225-64-1]	9-octylheptadecane					
		$\Delta_v H$	(470–505)	93.4	485	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₂₅ H ₅₂	[629-99-2]	pentacosane					
		$\Delta_{\text{urs}} H$		23.9	309		
		$\Delta_{\text{us}} H$		1.07	312.9		
		$\Delta_{\text{fus}} H$		55.53	325.9	DSC	[2006KHI/BOU]
		$\Delta_{\text{us}} H$		<0.4	310.5		
		$\Delta_{\text{us}} H$		0.4	319.4		
		$\Delta_{\text{us}} H$		23.6	320		
		$\Delta_{\text{us}} H$		<0.4	322.6		
		$\Delta_{\text{fus}} H$		57.8	326.4	DSC	[2004MON/RAJ]
		$\Delta_{\text{us}} H$		26.07	320.2		
		$\Delta_{\text{fus}} H$		57.74	326.7		[1996DOM/HEA]
		$\Delta_{\text{sub}} H$		173.6 ± 10	298	B	[1991PIA/POM]
		$\Delta_v H$	(382–462)	106.6	397		[2006SAW/MOK]
		$\Delta_v H$	(434–539)	126.8	298	CGC	[2004CHI/HAN]
		$\Delta_v H$		128.6 ± 2.2	298	GS	[2001PUR/CHI]
		$\Delta_v H$		127.6 ± 0.8	298	CGC	[2000NIC/ORF]
		$\Delta_v H$		129.8 ± 2.9	298	CGC	[1997CHI/WIL]
		$\Delta_v H$	(397–434)	126 ± 1	415	TE	[1994PIA/FON]
		$\Delta_v H$	(390–531)	97.6	405	TE, ME, GS	[1991PIA/POM]
		$\Delta_v H$	(461–498)	90.9 ± 5.7	479	GS	[1990PIA/SCA]
		$\Delta_v H$	(457–675)	99.2	472	A, EST	[1987STE/MAL, 1966KUD/ZWO]
C ₂₅ H ₅₂	[79370-85-7]	12-ethyltricosane					
		$\Delta_v H$	(435–454)	84.6	444	GC	[1982REC/GRE, 1999DYK/SVO]
C ₂₅ H ₅₂	[1560-78-7]	2-methyltetracosane					
		$\Delta_v H$	(425–670)	104.6	440		[1999DYK/SVO]
C ₂₅ H ₅₂	[126724-71-8]	5,5-bis(3,3'-dimethylbutyl)-2,2,8,8-tetramethylnonane					
		$\Delta_{\text{fus}} H$		48.53	472.7		[1990MEN/LIA]
		$\Delta_v H$		91.9 ± 1.8	298	CGC	[1995CHI/HES]
C ₂₅ H ₅₂	[163983-29-7]	7,7-dihexyltridecane					

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$		115.3 ± 1.8	298	CGC	[1995CHI/HES]
C ₂₅ H ₅₂ S	[66359-74-8]	1-pentacosanethiol				
	$\Delta_v H$	(458–709)	114.2	473	EST	[1999DYK/SVO]
C ₂₆ H ₁₄	[190-84-1]	1,12-phenyleneperylene (naphtha[1,2,3,4-ghi]perylene)				
	$\Delta_{\text{fus}} H$		17.28	541.5		[1991ACR]
C ₂₆ H ₁₅ Cl ₂ N ₅ O ₂	[68808-70-8]	1,3-bis(cyano-4-chlorophenylcarbamoyl-methylene)isoindolin				
	$\Delta_{\text{fus}} H$		123	679.2		[1993GRU]
C ₂₆ H ₁₆	[191-68-4]	dibenzo[g,p]chrysene				
	$\Delta_{\text{sub}} H$	(408–493)	142.2	423		[1987STE/MAL]
	$\Delta_{\text{sub}} H$	(417–500)	141.8	458	ME	[1967WAK/INO]
C ₂₆ H ₁₈	[1499-10-1]	9,10-diphenylanthracene				
	$\Delta_{\text{sub}} H$	(313–453)	137.5	383	GS	[1995NAS/LEN]
	$\Delta_{\text{sub}} H$		116.4			[1958KLO]
	$\Delta_{\text{sub}} H$	(393–433)	143.6	413		[1958HOY/PEP, 1987STE/MAL]
	$\Delta_{\text{sub}} H$	(481–502)	156.9 ± 4.2	492	HSA	[1953STE, 1970COX/PIL]
	$\Delta_v H$	(323–473)	102.7	398	GC	[2002LEI/CHA]
C ₂₆ H ₁₈	[1530-12-7]	9,9'-bifluorenyl				
	$\Delta_{\text{fus}} H$		36.9	519.2		[1994RAK/VER2]
	$\Delta_{\text{sub}} H$	(383–408)	131.8 ± 1.1	395	T	[1994RAK/VER2]
	$\Delta_{\text{sub}} H$		132.6 ± 1.1	298		[1994RAK/VER2]
	$\Delta_v H$	(383–408)	95.7		B	[1994RAK/VER2]
C ₂₆ H ₁₈ N ₂ O ₄	[6408-72-6]	disperse violet 31				
	$\Delta_v H$	(453–523)	59.9	468	A	[1987STE/MAL]
C ₂₆ H ₂₀	[632-51-9]	tetraphenylethene				
	$\Delta_{\text{fus}} H$		37.45	496.1	DSC	[1999VER/EBE]
	$\Delta_{\text{sub}} H$	(343–389)	129.3 ± 0.7	366	GS	[1999VER/EBE]
	$\Delta_{\text{sub}} H$		133.4 ± 0.7	298	GS	[1999VER/EBE]
C ₂₆ H ₂₀ N ₂ O ₂	[3073-87-8]	2,2'-(1,4-phenylene)bis(4-methyl-5-phenyl)oxazole				
	$\Delta_{\text{sub}} H$		150	480		[1989SCH/PEN]
C ₂₆ H ₂₂	[632-50-8]	1,1,2-tetraphenylethane				
	$\Delta_{\text{sub}} H$		136.8 ± 2.9	298	GS	[1990BEC/DOG2]
	$\Delta_{\text{sub}} H$	(370–423)	131.4 ± 2.1	396	GS	[1990BEC/DOG2]
C ₂₆ H ₂₂	[2294-94-2]	1,1,1,2-tetraphenylethane				
	$\Delta_{\text{sub}} H$		132.6 ± 2.1	298	GS	[1990BEC/DOG2]
	$\Delta_{\text{sub}} H$	(340–400)	128.7 ± 2.1	370	GS	[1990BEC/DOG2]
	$\Delta_{\text{sub}} H$		126.4 ± 1.7	434	HSA	[1956BEY/NIC]
C ₂₆ H ₂₂ N ₂ O ₂ S ₂	[109538-15-0]	1,2-bis-[5-(4-methoxy-β-azastyryl)-2-thienyl]-trans-ethylene				
	$\Delta_{\text{us}} H$ (liq <i>cryst</i>)		63.5	538.2		
	$\Delta_{\text{us}} H$ (liq <i>cr-liq</i>)		0.8	567.2		[1978KOS/BUD]
C ₂₆ H ₂₆	[35117-21-6]	pentacyclo[18.2.2.2(9,12).0(4,15).0(4,15).0(6,17)]hexacos-4,6(17),9,11,15,20,22,23,25-nonane (triple layered [2.2]paracyclophane)				
	$\Delta_{\text{sub}} H$	(299–412)	119.1 ± 1.5		TSGC	[1980NIS/SAK]
	$\Delta_{\text{sub}} H$		125.9 ± 2.5	298	TSGC	[1980NIS/SAK]
C ₂₆ H ₂₉ F ₂₅	[93454-73-0]	1,1,1,2,2,3,3,4,4,5,5,6,6,7,7,8,8,9,9,10,10,11,11,12,12-pentacosafuorohexacosane				

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound					
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference	
		$\Delta_{\text{us}}H$	16.3	363			
		$\Delta_{\text{fus}}H$	26.1	366	DSC	[1991HOP/MOL]	
		$\Delta_{\text{fus}}H$	26	359.2	DSC	[1986RUS/RAB]	
C ₂₆ H ₂₉ NO	[10540-29-1]	2-[4-[(1Z)-1,2-diphenyl-1-buten-1-yl]phenoxy]-N,N-dimethylethanamine (tamoxifen)					
		$\Delta_{\text{fus}}H$	34	371	DSC	[2007BER/WAS]	
C ₂₆ H ₃₂	[103042-85-9]	6-octyl-1,2,3,4-tetrahydronaphthacene					
		$\Delta_{\text{v}}H$	(503–574)	103.2	518	A	[1987STE/MAL]
C ₂₆ H ₃₂ O ₆	[na]	1,4,5,8-tetrakis(propoxy)-9,10-anthraquinone					
		$\Delta_{\text{fus}}H$	28.63	473.9		[2001NOR/TOU]	
C ₂₆ H ₃₃ NO ₃	[483362-63-6]	2-(4-nitrophenyl)-1-[4-[2-(trans-4-butylcyclohexyl)ethyl]phenyl]ethanone					
		$\Delta_{\text{fus}}H$	35.9	408.6	DSC	[2002SPA/DZI]	
C ₂₆ H ₃₄	[2883-70-7]	9-dodecylanthracene					
		$\Delta_{\text{v}}H$	(495–566)	99.4	510	A	[1987STE/MAL]
C ₂₆ H ₃₄	[3788-61-2]	9-dodecylphenanthrene					
		$\Delta_{\text{v}}H$	(495–568)	95.7	510	A	[1987STE/MAL]
C ₂₆ H ₃₄ O ₄	[na]	1,4,5,8-tetrapropoxyanthracene					
		$\Delta_{\text{fus}}H + \Delta_{\text{trs}}H$	43.93	410.2		[2001NOR/TOU]	
		Note: Authors report only the total enthalpy of melting. Numerical value contains enthalpies for two solid-solid transitions that occur at 370.7 K and 385.5 K. Larger tetraalkoxy-derivatives show liquid crystalline behavior.					
C ₂₆ H ₃₆ O ₃	[63042-30-8]	3-[(1-oxooctyl)oxy]-estra-1,3,5(10)-trien-17-one					
		$\Delta_{\text{fus}}H$	24.0	348	DSC	[1990YAN/EIR]	
C ₂₆ H ₃₈	[55268-63-8]	1,1-diphenyltetradecane					
		$\Delta_{\text{v}}H$	(467–530)	98.2	482	A	[1987STE/MAL]
C ₂₆ H ₃₈	[55268-62-7]	1,1-di(4-tolyl)dodecane					
		$\Delta_{\text{v}}H$	(466–529)	98.3	481	A	[1987STE/MAL]
C ₂₆ H ₃₈	[5171-91-5]	2,3-dimethyl-2,3-bis-(4-tert-butylphenyl)-butane					
		$\Delta_{\text{fus}}H$	43.93	493	DSC	[1983KRA/BEC]	
		$\Delta_{\text{sub}}H$	161.9		E,B	[1983KRA/BEC]	
C ₂₆ H ₃₈ O ₂	[3000-49-5]	3 β -octyloxy-estra-1,3,5(10)-trien-17-one					
		$\Delta_{\text{fus}}H$	19.0	331	DSC	[1990YAN/EIR]	
C ₂₆ H ₄₀	[95258-25-6]	5-octyl-1,2,3,4,4a,5,7,8,9,10,12,12a-dodecahydronaphthacene					
		$\Delta_{\text{v}}H$	(479–549)	91.9	494	A, MG	[1987STE/MAL, 1955SCH/WHI, 1999DYK/SVO]
C ₂₆ H ₄₀ O ₂	[128805-68-5]	3-(octyloxy)-estra-1,3,5(10)-trien-17-ol					
		$\Delta_{\text{fus}}H$	21.0	338	DSC	[1990YAN/EIR]	
C ₂₆ H ₄₂	[na]	1,1-bis(dodecahydroacenaphthylene-5-yl)ethane					
		$\Delta_{\text{v}}H$	(482–541)	110.9	497	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₂₆ H ₄₂ O	[141784-31-8]	<i>trans</i> -1-(4-heptanoylphenyl)-4-heptylcyclohexane					
		$\Delta_{\text{us}}H$ (liq <i>cryst</i>)	16.49	343.2			
		$\Delta_{\text{us}}H$ (liq <i>cr-liq</i>)	7.71	344.7	DSC	[1992FUR/BUT]	
C ₂₆ H ₄₂ O ₄	[14103-61-8]	<i>bis</i> (3,5,5-trimethylhexyl) phthalate					
		$\Delta_{\text{v}}H$	(333–393)	113.6	348	A	[1987STE/MAL]
C ₂₆ H ₄₂ O ₄	[84-76-4]	dinonyl phthalate					

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(333–393)	108.9	348	A	[1987STE/MAL]
C ₂₆ H ₄₆	[2655-95-0]	1,4-didecylbenzene				
	$\Delta_v H$	(468–536)	95.2	483	A	[1987STE/MAL]
C ₂₆ H ₄₆	[2398-68-7]	1-phenyleicosane				
	$\Delta_v H$	(499–538)	94.7	514	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₂₆ H ₄₆	[2398-66-5]	2-phenyleicosane				
	$\Delta_v H$	(492–531)	90.4	507	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₂₆ H ₄₆	[2400-02-4]	3-phenyleicosane				
	$\Delta_v H$	(489–526)	92.1	504	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₂₆ H ₄₆	[2400-03-5]	4-phenyleicosane				
	$\Delta_v H$	(487–527)	88.2	502	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₂₆ H ₄₆	[2400-04-6]	5-phenyleicosane				
	$\Delta_v H$	(485–521)	94.3	500	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₂₆ H ₄₆	[2398-64-3]	7-phenyleicosane				
	$\Delta_v H$	(483–520)	93.8	498	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₂₆ H ₄₆	[2398-65-4]	9-phenyleicosane				
	$\Delta_v H$	(483–520)	91.9	498	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₂₆ H ₄₆	[55191-36-1]	8-(4-tolyl)nonadecane				
	$\Delta_v H$	(482–517)	94.5	497	A	[1987STE/MAL]
C ₂₆ H ₄₈	[55401-75-7]	9-dodecyltetrahydroanthracene				
	$\Delta_v H$	(501–536)	102.7	519		[1999DYK/SVO]
C ₂₆ H ₄₈	[55334-01-5]	9-dodecyltetrahydrophenanthrene				
	$\Delta_v H$	(502–542)	90.8	522		[1999DYK/SVO]
C ₂₆ H ₄₈ N ₆ O ₉	[326813-31-4]	formamide deferoxamine				
	$\Delta_{\text{fus}} H$		92.93	430.8	DSC	[2000IHN/VEN]
C ₂₆ H ₄₈ O ₂	[25006-68-2]	4,4,7,7,13,13,16,16-octamethylcyclooctadecane-1,10-dione				
	$\Delta_{\text{fus}} H$		50.6	492.2		[1972BOR/DAL2]
C ₂₆ H ₅₀	[700004-11-1]	9-[α -(<i>cis</i> -bicyclo[3.3.0]octyl)methyl]heptadecane				
	$\Delta_v H$	(455–518)	92.3	470	A	[1987STE/MAL, 1999DYK/SVO]
C ₂₆ H ₅₀	[55334-09-3]	1,1- <i>bis</i> (4-methylcyclohexyl)dodecane				
	$\Delta_v H$	(484–520)	93.5	499	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₂₆ H ₅₀	[55334-08-2]	1,1-dicyclohexyltetradecane				
	$\Delta_v H$	(493–529)	97.7	508	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₂₆ H ₅₀	[55401-76-8]	1,1-dicyclopentylhexadecane				
	$\Delta_v H$	(471–525)	113.1	486	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₂₆ H ₅₀	[55334-11-7]	2-hexadecylbicyclopentyl				
	$\Delta_v H$	(495–532)	97.7	510	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₂₆ H ₅₀ N ₆ O ₁₀ S	[130281-67-3]	methylsulfonamide deferoxamine				
	$\Delta_{\text{fus}} H$		117.8	416	DSC	[2000IHN/VEN]
C ₂₆ H ₅₀ O ₄	[122-62-3]	(<i>dl</i>) <i>bis</i> (2-ethylhexyl) sebacate				
	$\Delta_v H$	(308–453)	114.9	323	A	[1987STE/MAL]
C ₂₆ H ₅₀ O ₄	[2432-87-3]	dioctyl sebacate				

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound					
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference	
		$\Delta_v H$	109.7	368	TGA	[1990KIS/SHO]	
		$\Delta_v H$	120.8 ± 4.2	298	TGA	[1990KIS/SHO]	
		$\Delta_v H$	(413–523)	107.1	428	A	[1987STE/MAL]
C ₂₆ H ₅₂	[4443-55-4]	1-cyclohexyleicosane					
		$\Delta_v H$	(499–538)	94.2	514	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₂₆ H ₅₂	[4443-56-5]	2-cyclohexyleicosane					
		$\Delta_v H$	(494–530)	98.3	509	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₂₆ H ₅₂	[4443-57-6]	3-cyclohexyleicosane					
		$\Delta_v H$	(492–530)	94	507	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₂₆ H ₅₂	[4443-58-7]	4-cyclohexyleicosane					
		$\Delta_v H$	(488–524)	98.3	503	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₂₆ H ₅₂	[4443-59-8]	5-cyclohexyleicosane					
		$\Delta_v H$	(488–524)	98.3	503	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₂₆ H ₅₂	[4443-60-1]	7-cyclohexyleicosane					
		$\Delta_v H$	(486–523)	93.6	501	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₂₆ H ₅₂	[4443-61-2]	9-cyclohexyleicosane					
		$\Delta_v H$	(486–523)	93.6	501	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₂₆ H ₅₂	[104338-48-9]	11-cyclohexyleicosane					
		$\Delta_{\text{fus}} H$	48.7	269.9		[1949PAR/MOO2]	
C ₂₆ H ₅₂	[6703-82-8]	1-cyclopentylheneicosane					
		$\Delta_v H$	(498–537)	93.8	513	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₂₆ H ₅₂	[6703-81-7]	11-cyclopentylheneicosane					
		$\Delta_v H$	(486–524)	92.4	501	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₂₆ H ₅₂	[18835-33-1]	1-hexacosene					
		$\Delta_v H$	(434–684)	106.1	449		[1999DYK/SVO]
C ₂₆ H ₅₂	[23014-57-5]	1,1,4,4,10,10,13,13-octamethylcyclooctadecane					
		$\Delta_{\text{us}} H$	6.74	427.2			
		$\Delta_{\text{fus}} H$	20.17	438.2		[1972BOR/DAL, 1969BOR/DAL]	
C ₂₆ H ₅₂ O ₂	[55373-89-2]	methyl pentacosanoate					
		$\Delta_{\text{fus}} H$	92.0	332.2	DSC	[2004CHI/ZHA]	
		$\Delta_v H$	(467–558)	142.0 ± 4.5	298	CGC	[2004CHI/ZHA]
C ₂₆ H ₅₂ O ₂	[24634-95-5]	ethyl tetracosanoate					
		$\Delta_{\text{us}} H$	11.2	317.7			
		$\Delta_{\text{fus}} H$	22.94	327.4		[1996DOM/HEA]	
C ₂₆ H ₅₃ NO	[74534-13-7]	N-decyl hexadecanamide					
		$\Delta_{\text{us}} H$	5.0	333			
		$\Delta_{\text{fus}} H$	63.0	347	DSC	[1980CAR/BUS]	
C ₂₆ H ₅₄	[55282-16-1]	5-butylidocosane					
		$\Delta_v H$	(482–518)	94.0	497	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₂₆ H ₅₄	[55282-15-0]	7-butylidocosane					
		$\Delta_v H$	(480–514)	97.2	495	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₂₆ H ₅₄	[55282-14-9]	9-butylidocosane					
		$\Delta_v H$	(479–516)	91.9	494	A, MG	[1987STE/MAL, 1955SCH/WHI]

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₂₆ H ₅₄	[13475-76-8] $\Delta_v H$	11-butylidocosane (480–516)	93.3	495	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₂₆ H ₅₄	[55282-13-8] $\Delta_v H$	5,14-dibutyloctadecane (458–508)	89.3	473	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₂₆ H ₅₄	[15874-03-0] $\Delta_v H$	6,11-dipentylhexadecane (468–504)	88.9	483	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₂₆ H ₅₄	[55282-12-7] $\Delta_v H$	3-ethyl-5-(2-ethylbutyl)octadecane (467–503)	88.4	482	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₂₆ H ₅₄	[55282-11-6] $\Delta_v H$	11-(1-ethylpropyl)heneicosane (474–509)	93.5	489	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₂₆ H ₅₄	[629-87-8] $\Delta_v H$	2-methylpentacosane (433–680)	107.2	448		[1999DYK/SVO]
C ₂₆ H ₅₄	[79370-84-6] $\Delta_v H$	12-propyltricosane (435–454)	91	445		[1982REC/GRE, 1999DYK/SVO]
C ₂₆ H ₅₄	[55282-17-2] $\Delta_v H$	3-ethyltetracosane (490–529)	90	505	A	[1987STE/MAL]
C ₂₆ H ₅₄	[630-01-3] $\Delta_{\text{us}}H$ $\Delta_{\text{fus}}H$ $\Delta_{\text{us}}H$ $\Delta_{\text{fus}}H$ $\Delta_{\text{us}}H$ $\Delta_{\text{fus}}H$ $\Delta_{\text{us}}H$ $\Delta_{\text{fus}}H$ $\Delta_{\text{sub}}H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$	hexacosane (434–539)	33.6 61.1 32.6 60.1 32.2 59.5 32.2 57.6 177.2 ± 10 131.7 139.3 ± 0.5 136.4 ± 0.2 140.0 ± 2.2 132 ± 1 97.6 99.0 ± 3.8 101.6 94.5	327.8 330.9 325 329.1 326.5 329.5 324.4 328.2 298 298 298 298 414 419 487 481 493	DSC DSC DSC DSC DSC DSC DSC DSC B CGC CGC GS CGC TE TE,ME,GS GS A, EST A	[2007GNA/PLA] [2004MON/RAJ] [1996DOM/HEA] [1992LOU/ROU] [1991PIA/POM] [2004CHI/HAN] [2002CHI/WEB] [2001PUR/CHI] [1997CHI/WIL] [1994PIA/FON] [1991PIA/POM] [1990PIA/SCA] [1987STE/MAL, 1966KUD/ZWO] [1987STE/MAL]
C ₂₆ H ₅₄	[55333-99-8] $\Delta_v H$	7-hexyleicosane (479–512)	101.1	494	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₂₆ H ₅₄	[55282-10-5] $\Delta_v H$	11-neopentylheneicosane (476–511)	93	491	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₂₆ H ₅₄	[14739-72-1] $\Delta_v H$	11-pentylheneicosane (478–512)	96.3	493	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₂₆ H ₅₄ O	[506-52-5] $\Delta_{\text{us}}H$ $\Delta_{\text{fus}}H$ $\Delta_v H$	1-hexacosanol 	16.74 67.78 148.0 ± 0.8	332.2 351.7 298		[1970TRA/LOM] [2006NIC/KWE]

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
C ₂₆ H ₅₄ O ₄ S ₂	[na]	<i>(l)</i> -rhamnose didecyl dithioacetal				
	$\Delta_{\text{fus}}H$		26.1	332.3		
	$\Delta_{\text{fus}}H$		53.2	385.2	DSC	[1989VAN/VAN]
C ₂₆ H ₅₄ S	[16331-25-2]	1-hexacosanethiol				
	$\Delta_{\text{v}}H$	(465–718)	116.2	480	EST	[1999DYK/SVO]
C ₂₇ H ₁₉ NO	[2083-09-2]	2,5- <i>bis</i> (1,1'-biphenyl)oxazole				
	$\Delta_{\text{v}}H$	(595–685)	109.7	610	A, I	[1987STE/MAL, 1975STE/SCH]
C ₂₇ H ₁₉ NO	[76733-99-8]	2-phenyl-5-(<i>p</i> -terphenyl-4-yl)oxazole				
	$\Delta_{\text{fus}}H$		42.0	504.2	DSC	[2001DIN/MUR]
C ₂₇ H ₁₉ NO	[362612-65-5]	2-(<i>p</i> -terphenyl-4-yl)-5-phenyloxazole				
	$\Delta_{\text{fus}}H$		37.0	485.2	DSC	[2001DIN/MUR]
C ₂₇ H ₃₀ N ₂	[221641-35-6]	2-(hept-1-ynyl)-5(4- <i>p</i> -hexylphenylbuta-1,3-diynyl) pyrimidine				
	$\Delta_{\text{fus}}H$		39.0	426	DSC	[1999HUD/SHE]
C ₂₇ H ₃₀ O ₃	[71203-39-9]	19-nor-17 α -ethynyl-17 β -(benzoyloxy-4-androsten-3-one				
	$\Delta_{\text{fus}}H$		41.5	531		[1996DOM/HEA]
C ₂₇ H ₃₀ O ₁₆	[153-18-4]	3-[[6-O-(6-deoxy- α -L-mannopyranosyl)- β -D-glucopyranosyl]-oxy]-2-(3,4-dihydroxyphenyl)-5,7-dihydroxy-4 <i>H</i> -1-benzopyran-4-one (rutin)				
	$\Delta_{\text{fus}}H$		82.3	450.2	DSC	[2007CHE/HUM]
C ₂₇ H ₃₂ O ₃	[138306-51-1]	spiro[8.5.0 ^(3,7)]-3,5-diphenyl-1,2,8-trioxa-10,12-tetramethyltridec-5-ene				
	$\Delta_{\text{fus}}H$		15.0	389.2	DSC	[1991JEF/JAB]
C ₂₇ H ₃₅ NO ₃	[483362-64-7]	2-(4-nitrophenyl)-1-[4-[2-(<i>trans</i> -4-pentylcyclohexyl)ethyl]phenyl]ethanone				
	$\Delta_{\text{fus}}H$		37.95	409	DSC	[2002SPA/DZI]
C ₂₇ H ₃₆ N ₂ O ₅	[na]	N,N-diisopropyl naltrexone-3-O-carbamate				
	$\Delta_{\text{fus}}H$		19.95	421.2	DSC	[2009VAD/BAN]
C ₂₇ H ₃₈ O	[3836-23-5]	19-nor-17 α -ethynyl-17 β -(heptanoyloxy-4-androsten-3-one				
	$\Delta_{\text{fus}}H$		21.6	340		[1996DOM/HEA]
C ₂₇ H ₃₈ O ₃	[105755-75-7]	3-[(1-oxononyl)oxy]-estra-1,3,5(10)-trien-17-one				
	$\Delta_{\text{fus}}H$		24.0	337	DSC	[1990YAN/EIR]
C ₂₇ H ₄₀	[55334-13-9]	5-pentadecylacenaphthene				
	$\Delta_{\text{v}}H$	(500–568)	105.7	105.7	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₂₇ H ₄₀ N ₄ O ₂	[182410-21-5]	2,2'-[(2,8,10-trimethylpyrido[3,2- <i>g</i>]quinoline-4,6-diyl) <i>bis</i> (oxy)]- <i>bis</i> [N,N-diethylethanamine]				
	$\Delta_{\text{fus}}H$		16.1	415.3	DSC	[2008ABB/KAL]
C ₂₇ H ₄₂ Cl ₂ N ₂ O ₆	[530-43-8]	hexadecanoic acid, [R-(R*,R*)]-2-[(dichloroacetyl)amino]-3-hydroxy-3-(4-nitrophenyl)propyl ester (chloramphenicol palmitate)				
	$\Delta_{\text{fus}}H$		64.02	368.2		[1998VAN/KEL, 1977BUR, 1970BOR]
	$\Delta_{\text{fus}}H$ (I)		51.04	367.3		
	$\Delta_{\text{fus}}H$ (II)		41.3	360.8		[1985OHM/LIP]
C ₂₇ H ₄₄ O	[313-04-2]	desmosterol				
	$\Delta_{\text{fus}}H$		15.9	388.2	DSC	[2009CHE/SU]
C ₂₇ H ₄₆ O	[57-88-5]	cholesterol				
	$\Delta_{\text{fus}}H$		28.5	421.7	DSC	[2009CHE/SU]
	$\Delta_{\text{fus}}H$	(300–440)	25.1	423.2	DSC	[2008PEN/JIA]
	$\Delta_{\text{us}}H$		2.71	311.7		
	$\Delta_{\text{fus}}H$		21.11	423.2	DSC	[2003KAL/PAU]
$\Delta_{\text{us}}H$	(5–425)	3.9	306.7			

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
		$\Delta_{\text{fus}}H$	28.4	422.3	AC	[1998VAN/VAN]
		$\Delta_{\text{trs}}H$	2.5	304.8		
		$\Delta_{\text{fus}}H$	27.41	420.2		[1996DOM/HEA]
		$\Delta_{\text{sub}}H$	(386–414)	142.5 ± 0.9	ME	[2009OJA/CHE]
		Δ_vH		153.7 ± 0.8	298	CGC [2006NIC/KWE]
		Δ_vH	(411–447)	114.9	426	A [1987STE/MAL]
C ₂₇ H ₄₆ O	[57-88-5]	5-cholesten-3 β -ol				
		$\Delta_{\text{fus}}H$	26.5	422.5		[2001IWA/MIN]
C ₂₇ H ₄₈	[481-21-0]	17-(1,5-dimethylhexyl)-10,13-dimethyl-hexahydro-1 <i>H</i> -cyclopenta[<i>a</i>]-phenanthrene (5- α -cholestane)				
		$\Delta_{\text{fus}}H$	25.4	351.8		[2000MOK/RUZ]
		$\Delta_{\text{sub}}H$		133.8	298	[2000MOK/RUZ]
		Δ_vH		108.4	352	[2000MOK/RUZ]
		Δ_vH	(481–538)	115.6	496	A [1987STE/MAL]
C ₂₇ H ₄₈	[40775-09-5]	heneicosylbenzene				
		Δ_vH	(446–705)	108.4	461	[1999DYK/SVO]
C ₂₇ H ₄₈	[6703-80-6]	11-phenylheneicosane				
		$\Delta_{\text{fus}}H$	64.77	294.3		[1949PAR/MOO2]
		Δ_vH	(491–529)	93.5	506	A, MG [1987STE/MAL, 1955SCH/WHI]
C ₂₇ H ₄₈ N ₂ OS	[467434-73-7]	N-[(3-methoxyphenyl)methyl]- <i>N'</i> -octadecylthiourea				
		$\Delta_{\text{fus}}H$	64.17	375.2	DSC	[2002ABB/WHO]
C ₂₇ H ₄₈ O	[80-97-7]	5 α -cholestan-3 β -ol				
		$\Delta_{\text{fus}}H$	22.6	413.5		[2001IWA/MIN]
C ₂₇ H ₄₈ O	[516-92-7]	5 β -cholestan-3 α -ol				
		$\Delta_{\text{fus}}H$	15.8	385.8		[2001IWA/MIN]
C ₂₇ H ₄₈ O	[360-68-9]	5 β -cholestan-3 β -ol				
		$\Delta_{\text{fus}}H$	16.1	373.8		[2002MIN/SAK]
C ₂₇ H ₅₀	[55282-69-4]	5-pentacyclododecahydroacenaphthalene				
		Δ_vH	(486–554)	98.1	501	A [1987STE/MAL]
C ₂₇ H ₅₀ N ₆ O ₉	[5722-48-5]	acetamide deferroxamine				
		$\Delta_{\text{fus}}H$	118.4	448.9	DSC	[2000IHN/VEN]
C ₂₇ H ₅₀ O ₆	[538-23-8]	glycerol trioctanoate				
		Δ_vH	118.7	386	TGA	[1990KIS/SHO]
		Δ_vH	135.4 ± 4.7	298	TGA	[1990KIS/SHO]
		Δ_vH	(396–453)	116	411	A, T [1987STE/MAL, 1949PER/WEB2]
C ₂₇ H ₅₄	[6703-99-7]	11-cyclohexylheneicosane				
		Δ_vH	(485–529)	107	500	A [1987STE/MAL]
C ₂₇ H ₅₄	[6703-79-3]	11-(cyclopentylmethyl)heneicosane				
		Δ_vH	(492–529)	94.4	507	A [1987STE/MAL]
C ₂₇ H ₅₄	[26718-82-1]	heneicosylcyclohexane				
		Δ_vH	(445–460)	107.8	460	[1999DYK/SVO]
C ₂₇ H ₅₄	[15306-27-1]	1-heptacosene				
		Δ_vH	(441–694)	108.7	456	[1999DYK/SVO]

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₂₇ H ₅₄	[163983-30-0] $\Delta_v H$	1-decyl-1-undecylcyclohexane	133.6 ± 1.8	298	CGC	[1995CHI/HES]
C ₂₇ H ₅₄ N ₃ PS ₆	[194281-16-8] $\Delta_{\text{sub}} H$	tris(diisobutylthiocarbamate)phosphorous	138 ± 3		DSC, E	[1997DES/DES]
C ₂₇ H ₅₄ N ₆	[38565-87-6] $\Delta_{\text{fus}} H$	tris N,N-diisobutylamino-1,3,5-triazine	35.81	372.6		[1986LAT/HOE]
C ₂₇ H ₅₄ O ₂	[5802-82-4] $\Delta_{\text{fus}} H$	methyl hexacosanoate	101.3	336.2	DSC	[2004CHI/ZHA]
	$\Delta_v H$	(467–558)	147.1 ± 4.5	298	CGC	[2004CHI/ZHA]
C ₂₇ H ₅₆	[593-49-7] $\Delta_{\text{us}} H$	heptacosane	0.3	312.9		
	$\Delta_{\text{us}} H$		2.5	322.3		
	$\Delta_{\text{us}} H$		27.1	325.9		
	$\Delta_{\text{fus}} H$		62.8	331.6	DSC	[2004MON/RAJ]
	$\Delta_{\text{us}} H$		2.26	318		
	$\Delta_{\text{us}} H$		26.28	325.4		
	$\Delta_{\text{fus}} H$		59.05	332.1	DSC	[1992LOU/ROU]
	$\Delta_{\text{sub}} H$		196.0 ± 30	298	B	[1991PIA/POM]
	$\Delta_v H$	(401–441)	132 ± 1	423	TE	[1994PIA/FON]
	$\Delta_v H$	(508–570)	94.2	523	ME,TE,GS	[1991PIA/POM]
C ₂₇ H ₅₆	[1561-02-0] $\Delta_v H$	2-methylhexacosane	109.6	456		[1999DYK/SVO]
	[55282-29-6] $\Delta_v H$	8-hexyl-8-pentylhexadecane	125.7 ± 1.8	298	CGC	[1995CHI/HES]
C ₂₇ H ₅₆	[55282-28-5] $\Delta_v H$	8,8-dipentylheptadecane	128.1 ± 1.8	298	CGC	[1995CHI/HES]
C ₂₇ H ₅₆	[55282-32-1] $\Delta_v H$	10-hexyl-10-methyleicosane	129.9 ± 1.8	298	CGC	[1995CHI/HES]
C ₂₇ H ₅₆	[55282-30-9] $\Delta_v H$	5-ethyl-5-methyltetracosane	133.8 ± 1.8	298	CGC	[1995CHI/HES]
C ₂₇ H ₅₆ S	[66291-85-8] $\Delta_v H$	1-heptacosanethiol	118.3	486	EST	[1999DYK/SVO]
C ₂₈ H ₁₂ Cl ₂ N ₂ O ₄	[130-20-1] $\Delta_{\text{sub}} H$	C.I. Vat Blue 6	199	577	GS	[1986NIS/AND]
C ₂₈ H ₁₄	[190-39-6] $\Delta_{\text{sub}} H$	phenanthro[1,10,9,8-opqra]perylene	180.5			[1987STE/MAL]
	$\Delta_{\text{sub}} H$	(580–630)	180.7 ± 5	605	ME	[1952INO/SHI]
	$\Delta_v H$	(580–630)	180.5	595	A	[1987STE/MAL]
C ₂₈ H ₁₄ N ₂ O ₄	[81-77-6] $\Delta_v H$	C.I. Vat Blue 4	167	577	GS	[1986NIS/AND]
C ₂₈ H ₁₆	[na]	1,2,4,5,7,8-tribenzopyrene				

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{fus}}H$		28.8	608		[1979SMI2]
C ₂₈ H ₁₈	[1055-23-8]	9,9'-bianthryl				
	$\Delta_{\text{sub}}H$		128.4 ± .2			[1970COX/PIL, 1958HOY/PEP]
	$\Delta_{\text{sub}}H$	(413–473)	127.9	443	A	[1958HOY/PEP, 1987STE/MAL]
	$\Delta_{\text{sub}}H$		148.1			[1951MAG/HAR, 1960JON]
C ₂₈ H ₁₈	[20532-03-0]	9,9'-biphenanthryl				
	$\Delta_{\text{sub}}H$		151.5			[1951MAG/HAR, 1960JON]
C ₂₈ H ₂₀ S	[362612-62-2]	3-(<i>p</i> -terphenyl-4-yl)-5-phenylthiophene				
	$\Delta_{\text{fus}}H$		43	561.2	DSC	[2001DIN/MUR]
C ₂₈ H ₂₀ S	[362612-62-2]	2-(<i>p</i> -terphenyl-4-yl)-4-phenylthiophene				
	$\Delta_{\text{fus}}H$		42	554.2	DSC	[2001DIN/MUR]
C ₂₈ H ₂₀ S	[56316-86-0]	2,5- <i>bis</i> (biphenyl-4-yl)thiophene				
	$\Delta_{\text{fus}}H$		39	595.2	DSC	[2001DIN/MUR]
C ₂₈ H ₂₂	[15300-82-0]	9,9'-dimethyl-9,9'-bifluorenyl				
	$\Delta_{\text{sub}}H$	(368–403)	118.7 ± 1.3	386	T	[1994RAK/VER2]
	$\Delta_{\text{sub}}H$	(368–403)	119.7 ± 1.3	298	T	[1994RAK/VER2]
	Δ_vH	(368–403)	94.6		B	[1994RAK/VER2]
C ₂₈ H ₂₂ N ₂ O ₂	[na]	1,4- <i>bis</i> [(4-methylphenyl)amino]-9,10-anthracenedione				
	$\Delta_{\text{fus}}H$		36.59	491.2		[1991BAU/WEB]
C ₂₈ H ₂₄ O ₄	[74568-07-3]	calix[4]arene-25,26,27,28-tetrol				
	$\Delta_{\text{sub}}H$		167 ± 2			[2008SUR]
C ₂₈ H ₂₄ O ₈	[125748-07-4]	2,8,14,20-tetramethyl-4,6,10,12,16,18,20,24-octahydroxyresorci[4]arene (calix[4]resorcinarene)				
	$\Delta_{\text{fus}}H$		38.2	578.6	DSC	[2010FRA/SAL]
C ₂₈ H ₂₄ O ₁₆ S ₄	[112269-92-8]	4-sulfonato-calix[4]arene				
	$\Delta_{\text{fus}}H$		192.4	549.8		[2005YAN/MAN]
C ₂₈ H ₂₆ N ₄ O ₈	[74734-27-3]	1,4- <i>bis</i> (3-phenylcarbzomoyl-2-oxo-5-oxazolidin-5-ylmethoxy)benzene				
	$\Delta_{\text{fus}}H$ (I)		10.1	475.2		[1990SHI/HAY]
	$\Delta_{\text{fus}}H$ (II)		5.1	502.2		
C ₂₈ H ₂₈ P	[7688-25-7]	1,4- <i>bis</i> (diphenylphosphino)butane				
	$\Delta_{\text{fus}}H$		45.3	405.9	DTA	[1989HUI/VAN]
	$\Delta_{\text{sub}}H$	(425–455)	171.6 ± 2.5	443	B	[1989HUI/VAN]
	Δ_vH	(425–455)	126.3 ± 2	443	ME	[1989HUI/VAN]
C ₂₈ H ₂₉ F ₂ N ₃ O	[2062-78-4]	1-[1-[4,4- <i>bis</i> (4-fluorophenyl)butyl]-4-piperidyl]-2-benzimidazolinone (pimozide)				
	$\Delta_{\text{fus}}H$		46.9	493.5	DSC	[2008THI/SUB]
C ₂₈ H ₃₀ N ₄	[1257-25-6]	2,3,7,8,12,13,17,18-octamethylporphyrin				
	$\Delta_{\text{sub}}H$	(593–653)	268 ± 11		GS	[2001NIK/SUL]
C ₂₈ H ₃₁ FN ₄ O	[na]	1-[(4-fluorophenyl)methyl]-N-[1-[2-(4-methoxyphenyl)ethyl]-4-piperidiny]-1 <i>H</i> -benzimidazol-2-amine (astemizole)				
	$\Delta_{\text{fus}}H$		51.1	447.6	DSC	[2007BER/WAS]
C ₂₈ H ₃₁ F ₂₅	[93454-74-1]	1,1,1,2,2,3,3,4,4,5,5,6,6,7,7,8,8,9,9,10,10,11,11,12,12-pentacosofluorooctacosane				
	$\Delta_{\text{fus}}H$		43.1	263.2	DSC	[1986RUS/RAB]
C ₂₈ H ₃₂	[55282-03-6]	1,7-diphenyl-4-(3-phenylpropyl)-3-heptene				
	Δ_vH	(488–556)	98.0	503	A, MG	[1987STE/MAL, 1955SCH/WHI]

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₂₈ H ₃₄	[5282-64-9] $\Delta_v H$	1,7-diphenyl-4-(3-phenylpropyl)heptane (490–557)	100.3	505	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₂₈ H ₃₈	[59358-73-5] $\Delta_{\text{fus}} H$	1,1-diphenyl-1,1'-bicyclooctyl	35.98	432	DSC	[1983KRA/BEC]
	$\Delta_{\text{sub}} H$		174.5		E,B	[1983KRA/BEC]
C ₂₈ H ₄₀ O ₃	[128788-27-2] $\Delta_{\text{fus}} H$	3-[(1-oxodecyl)oxy]-estra-1,3,5(10)-trien-17-one	29.0	344	DSC	[1990YAN/EIR]
C ₂₈ H ₄₄ O	[57-87-4] $\Delta_{\text{sub}} H$	ergosterol (318–412)	147.0 ± 0.9		ME	[2009OJA/CHE]
	$\Delta_v H$		(421–454)	118.7	436	A
C ₂₈ H ₄₆ O	[141784-32-9] $\Delta_{\text{us}} H$ (liq <i>cryst</i>)	<i>trans</i> -1-heptyl-4-(4-nonanoylphenyl)cyclohexane	20.8	343.4		
	$\Delta_{\text{us}} H$ (liq <i>cr-liq</i>)		11.32	353.3	DSC	[1992FUR/BUT]
C ₂₈ H ₄₆ O ₄	[26761-40-0] $\Delta_v H$	diisodecyl phthalate (371–496)	79.3	386	A	[1987STE/MAL]
C ₂₈ H ₄₈ O ₄	[175848-67-6] $\Delta_{\text{us}} H$	2,5-di- <i>n</i> -undecyloxy-1,4-benzoquinone	12.9	367.4		
	$\Delta_{\text{us}} H$		28.4	390		
	$\Delta_{\text{fus}} H$		52.1	397.2	DSC	[1996KEE/VAN]
C ₂₈ H ₅₀	[5634-22-0] $\Delta_v H$	docosylbenzene (453–715)	110.8	468		[1999DYK/SVO]
C ₂₈ H ₅₀	[55334-72-0] $\Delta_v H$	2-decyl-1-phenyldodecane (497–532)	102.5	512	A	[1987STE/MAL]
C ₂₈ H ₅₀ O ₁₁	[na] $\Delta_v H$	di[1-(2-ethylhexyl)oxycarbonyl]ethyl diethylene glycol dicarboxylate (463–553)	116.6	478	A	[1987STE/MAL]
C ₂₈ H ₅₀ O ₁₁	[na] $\Delta_v H$	di[1-(octyloxycarbonyl)ethyl] diethylene glycol dicarboxylate (463–564)	112.5	478	A	[1987STE/MAL]
C ₂₈ H ₅₂	[55334-73-1] $\Delta_v H$	1,7-dicyclohexyl-4-(3-cyclohexylpropyl)heptane (482–549)	98.7	497	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₂₈ H ₅₂ N ₆ O ₉	[326813-19-8] $\Delta_{\text{fus}} H$	propylamide deferoxamine	116.9	449.6	DSC	[2000IHN/VEN]
C ₂₈ H ₅₂ O ₂	[29844-60-8] $\Delta_{\text{fus}} H$	4,4,8,8,14,14,18,18-octamethylcycloeicosane-1,11-dione	36.8	418.2		[1972BOR/DAL2]
C ₂₈ H ₅₄	[55255-74-8] $\Delta_v H$	1-cyclohexyl-2-(cyclohexylmethyl)pentadecane (501–536)	105.4	516	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₂₈ H ₅₆	[61828-07-7] $\Delta_v H$	docosylcyclohexane (452–715)	110	467		[1999DYK/SVO]
C ₂₈ H ₅₆	[18835-34-2] $\Delta_v H$	1-octacosene (448–703)	111	463		[1999DYK/SVO]
C ₂₈ H ₅₆	[6704-00-3] $\Delta_v H$	11-(cyclohexylmethyl)heneicosane (499–538)	94.2	514	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₂₈ H ₅₆	[55255-73-7] $\Delta_v H$	2,2,4,10,12,12-hexamethyl-7-(3,5,5-trimethylhexyl)-6-tridecene (426–488)	83.8	441	A, MG	[1987STE/MAL, 1955SCH/WHI]

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound					
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference	
C ₂₈ H ₅₆	[29844-61-9] $\Delta_{\text{fus}}H$	1,1,5,5,11,11,15,15-octamethylcycloicosane	47.7	439.2		[1972BOR/DAL]	
C ₂₈ H ₅₆ O ₂	[55682-91-2] $\Delta_{\text{fus}}H$	methyl heptacosanoate	100.7	336.2	DSC	[2004CHI/ZHA]	
	Δ_vH		(467–558)	152.2 ± 4.5	298	CGC	[2004CHI/ZHA]
C ₂₈ H ₅₆ O ₂	[29030-81-7] $\Delta_{\text{us}}H$	ethyl hexacosanoate	13.22	322.7			
	$\Delta_{\text{fus}}H$			27.05	322.7		[1996DOM/HEA]
C ₂₈ H ₅₈	[1561-00-8] Δ_vH	2-methylheptacosane	111.9	463		[1999DYK/SVO]	
C ₂₈ H ₅₈	[3035-75-4] Δ_vH	2,2,4,10,12,12-hexamethyl-7-(3,5,5-trimethylhexyl)tridecane	(308–393)	98.5	323	A	[1987STE/MAL, 1964MOR]
	Δ_vH		(429–491)	84.9	444	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₂₈ H ₅₈	[55373-86-9] Δ_vH	7-hexyldocosane	(506–531)	100.7	518	A	[1987STE/MAL]
C ₂₈ H ₅₈	[630-02-4] $\Delta_{\text{us}}H$	octacosane	31.52	329.6			
	$\Delta_{\text{fus}}H$		67.38	334.2	DSC	[2007HAF/MAH]	
	$\Delta_{\text{us}}H$		35.2	329.4			
	$\Delta_{\text{fus}}H$		63.0	333.4	DSC	[2004MON/RAJ]	
	$\Delta_{\text{us}}H$		35.44	331.4			
	$\Delta_{\text{fus}}H$		64.64	334.5		[1996DOM/HEA]	
	$\Delta_{\text{sub}}H$		(323–329)	195.8 ± 2.2	326		[2009RAZ/NAC]
	$\Delta_{\text{sub}}H$			208.9 ± 10	298	B	[1991PIA/POM]
	Δ_vH		(339–412)	117.4 ± 1.2	376		[2009RAZ/NAC]
	Δ_vH		(354–517)	118.5	369	GC	[2007MOK/RAZ]
	Δ_vH		(434–539)	141.9	298	CGC	[2004CHI/HAN]
	Δ_vH			150.8 ± 0.5	298	CGC	[2002CHI/WEB]
	Δ_vH			150.7 ± 1.7	298	CGC	[2000NIC/ORF]
	Δ_vH			152.4 ± 2.9	298	CGC	[1997CHI/WIL]
	Δ_vH		(483–588)	100.5	498		[1994MOR/KOB]
	Δ_vH		(407–456)	135 ± 3	431	TE	[1994PIA/FON]
	Δ_vH		(426–493)	105.5	441	TE,ME,GS	[1991PIA/POM]
	Δ_vH		(473–515)	103.1 ± 3.0	494	GS	[1990PIA/SCA]
	Δ_vH		(450–575)	100.6	500	EB,IP	[1989CHI/NGU]
	Δ_vH		(450–575)	98.1	560	EB,IP	[1989CHI/NGU]
Δ_vH	(300–390)	131.7	315	A	[1987STE/MAL]		
Δ_vH	(481–705)	106.6	496	A, EST	[1987STE/MAL, 1966KUD/ZWO]		
C ₂₈ H ₅₈	[13475-77-9] Δ_vH	9-octyleicosane	(460–530)	106.8	475	A	[1987STE/MAL]
C ₂₈ H ₅₈ O	[5412-98-6] $\Delta_{\text{us}}H$	15-oxanonacosane	8.37	315.6			
	$\Delta_{\text{fus}}H$		113.39	316.8	DSC	[2004TYA/BIS]	
C ₂₈ H ₅₈ S	[16331-26-3] Δ_vH	1-octacosanethiol	(477–736)	120.2	492	EST	[1999DYK/SVO]
C ₂₉ H ₃₅ NO ₂	[84371-65-3] $\Delta_{\text{fus}}H$	17 β -hydroxy-11 β -[4-(dimethylamino)-phenyl]-17 α -(prop-1-ynyl)-estra-4,9-dien-3-one (Mifepristone)	31.7	467.1	DSC	[2006WAS/HOL]	

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₂₉ H ₄₁ NO ₄	[52485-79-7]	17-(cyclopropylmethyl)- α -(1,1-dimethylethyl)-4,5-epoxy-18,19-dihydro-6-methoxy- α -methyl-6,14-ethenomorphinan-7-methanol				
	$\Delta_{\text{fus}}H$		26.8	491.3		[1995STI/DUA]
C ₂₉ H ₄₂ O ₃	[105755-76-8]	3-[(1-oxoundecyl)oxy]-estra-1,3,5(10)-trien-17-one				
	$\Delta_{\text{fus}}H$		34.0	345	DSC	[1990YAN/EIR]
C ₂₉ H ₄₄ O ₂	[118-82-1]	3,3',5,5'-tetra- <i>tert</i> -butyldiphenylmethane-4,4'-diol				
	$\Delta_{\text{fus}}H$		42.97	447.7	DTA	[1972INO/LIA]
C ₂₉ H ₄₈ O	[83-48-7]	β -stigmaterol				
	$\Delta_{\text{sub}}H$	(390–417)	168.4 ± 1.4		ME	[2009OJA/CHE]
C ₂₉ H ₅₀	[55373-90-5]	11-(2,5-dimethylphenyl)-10-heneicosene				
	Δ_vH	(471–534)	99.2	486	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₂₉ H ₅₀ O	[83-46-5]	β -sitosterol				
	$\Delta_{\text{sub}}H$	(389–410)	143.8 ± 0.5		ME	[2009OJA/CHE]
C ₂₉ H ₅₂	[61828-04-4]	tricosylbenzene				
	Δ_vH	(459–724)	113.2	474		[1999DYK/SVO]
C ₂₉ H ₅₂	[18835-35-3]	1-nonacosene				
	Δ_vH	(455–713)	113.3	470		[1999DYK/SVO]
C ₂₉ H ₅₂	[55373-91-6]	11-(2,5-dimethylphenyl)heneicosane				
	Δ_vH	(472–535)	100.8	487	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₂₉ H ₅₂ N ₆ O ₁₁	[84211-47-2]	succinamide deferoxamine				
	$\Delta_{\text{fus}}H$		101	436.2	DSC	[2000IHN/VEN]
C ₂₉ H ₅₄ N ₆ O ₉	[326813-21-2]	butylamide deferoxamine				
	$\Delta_{\text{fus}}H$		111.4	451.1	DSC	[2000IHN/VEN]
C ₂₉ H ₅₈	[61828-08-8]	tricosylcyclohexane				
	Δ_vH	(459–724)	112.3	474		[1999DYK/SVO]
C ₂₉ H ₅₈ O ₂	[55682-92-3]	methyl octacosanoate				
	$\Delta_{\text{fus}}H$		109.7	340.2	DSC	[2004CHI/ZHA]
	Δ_vH	(467–558)	157.5 ± 4.5	298	CGC	[2004CHI/ZHA]
C ₂₉ H ₆₀	[1560-98-1]	2-methyloctacosane				
	Δ_vH	(455–709)	114.2	470		[1999DYK/SVO]
C ₂₉ H ₆₀	[630-03-5]	nonacosane				
	$\Delta_{\text{us}}H$		29.71	331.4		
	$\Delta_{\text{fus}}H$		66.11	336.6		[1996DOM/HEA]
	Δ_vH	(422–452)	112.5	437		[2006SAW/MOK]
	Δ_vH	(434–539)	147.1	298	CGC	[2004CHI/HAN]
	Δ_vH	(423–457)	137 ± 3	440	TE	[1994PIA/FON]
	Δ_vH	(423–456)	137.1 ± 3.0	439	TE	[1990POM/PIA]
Δ_vH	(488–714)	109	503	A, EST	[1987STE/MAL, 1966KUD/ZWO]	
C ₂₉ H ₆₀ S	[66213-92-1]	1-nonacosanethiol				
	Δ_vH	(483–744)	122	498	EST	[1999DYK/SVO]
C ₃₀ H ₁₄ O ₂	[128-70-1]	8,16-pyranthenedione (C. I. Vat Orange 9)				
	$\Delta_{\text{sub}}H$	(503–543)	197.7	518		[1987STE/MAL]
	$\Delta_{\text{sub}}H$		181.2	498	ME	[1951INO]

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₃₀ H ₁₆	[191-13-9] $\Delta_{\text{sub}}H$	pyranthrene	194.5 ± 6.7	595	ME	[1952INO/SHI]
C ₃₀ H ₂₂	[13476-68-1] $\Delta_{\text{fus}}H$	1,3-bis(biphenyl-4-yl)benzene	55.0	548.2	DSC	[2001DIN/MUR]
C ₃₀ H ₂₂	[13478-57-4] $\Delta_{\text{fus}}H$	1-(<i>p</i> -terphenyl-4-yl)-3-phenylbenzene	56.0	531.2	DSC	[2001DIN/MUR]
C ₃₀ H ₂₈ O ₄	[142433-64-5] $\Delta_{\text{sub}}H$	25,27-dimethoxycalix[4]arene-26,28-diol	75 ± 2			[2008SUR]
C ₃₀ H ₃₀	[2819-41-2] Δ_vH	1,1,6,6-tetraphenylhexane (511–579)	108.1	526	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₃₀ H ₃₂ P ₂	[19845-69-3] $\Delta_{\text{fus}}H$	1,6-bis(diphenylphosphino)hexane	66.8	399.4		[1998ZHA/TAN]
C ₃₀ H ₃₄	[40339-27-3] Δ_vH	1,10-di(1-naphthyl)decane (540–616)	108.6	555	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₃₀ H ₃₇ F ₂₅	[93454-75-2] $\Delta_{\text{fus}}H$	1,1,1,2,2,3,3,4,4,5,5,6,6,7,7,8,8,9,9,10,10,11,11,12,12-pentacosofluorotricontane	47.8	365.2	DSC	[1986RUS/RAB]
C ₃₀ H ₄₀ N ₄ O ₂	[na] $\Delta_{\text{fus}}H$	α,ω -bis(azobenzene-4-oxy)hexane	73.53	442.2		[2000BLA/LUC]
C ₃₀ H ₄₄ O ₃	[128788-28-3] $\Delta_{\text{fus}}H$	3-[(1-oxododecyl)oxy]-estra-1,3,5(10)-trien-17-one	31	342	DSC	[1990YAN/EIR]
C ₃₀ H ₄₆	[85668-74-2] $\Delta_{\text{fus}}H$ $\Delta_{\text{sub}}H$	3,4-diethyl-3,4-bis-(4- <i>tert</i> -butylphenyl)-hexane	29.71 167.8	400	DSC E,B	[1983KRA/BEC] [1983KRA/BEC]
C ₃₀ H ₄₆ F ₄ O ₂	[79312-06-4] $\Delta_{\text{fus}}H$	cholesteryl 2,2,3,3-tetrafluoropropionate	28.6	422.6		[1981YAN/NAB]
C ₃₀ H ₄₆ O ₂ S	[1620-93-5] $\Delta_{\text{fus}}H$	bis-[3,5-di- <i>tert</i> -butyl-4-hydroxybenzyl]sulfide	43.1	417.2	DTA	[1972INO/LIA]
C ₃₀ H ₄₉ BrO ₂	[73112-93-3] $\Delta_{\text{fus}}H$	cholesteryl α -bromopropionate	35.7	409.5		[1981YAN/NAB]
C ₃₀ H ₄₉ ClO ₂	[79312-05-3] $\Delta_{\text{fus}}H$	cholesteryl α -chloropropionate	48.5	409.1		[1981YAN/NAB]
C ₃₀ H ₅₄	[55268-64-9] Δ_vH	1,10-bis(decahydro-1-naphthyl)decane (520–583)	119.7	535	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₃₀ H ₅₄	[55281-91-9] Δ_vH	1,1,6,6-tetracyclohexylhexane (501–569)	103	516	A	[1987STE/MAL]
C ₃₀ H ₅₄	[61828-05-5] Δ_vH	tetracosylbenzene (466–732)	115.3	481		[1999DYK/SVO]
C ₃₀ H ₅₄ O ₆	[52193-50-7] Δ_vH	<i>trans</i> tris(2-ethylhexyl) aconitate (437–551)	97.1	452	A	[1987STE/MAL]
C ₃₀ H ₅₄ O ₆	[5400-99-7] Δ_vH	<i>tris</i> (2-ethylhexyl) 1,2,3-propanetricarboxylate (438–551)	97.9	453	A	[1987STE/MAL]
C ₃₀ H ₅₆ N ₆ O ₉	[103991-47-5]	valeramide deferoxamine				

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{fus}}H$		123.1	453.6	DSC	[2000IHN/VEN]
C ₃₀ H ₅₆ O ₂	[37465-23-9]	5,5,8,8,16,16,19,19-octamethylcyclodocosane-1,12-dione				
	$\Delta_{\text{fus}}H$		47.7	442.2		[1972BOR/DAL2]
C ₃₀ H ₅₈ O ₄	[2432-89-5]	didecyl sebacate				
	Δ_vH		120.5	405	TGA	[1990KIS/SHO]
	Δ_vH		138.7 ± 4.9	298	TGA	[1990KIS/SHO]
C ₃₀ H ₆₀	[61828-09-9]	tetracosylcyclohexane				
	Δ_vH	(465–733)	114.6	480		[1999DYK/SVO]
C ₃₀ H ₆₀	[18435-53-5]	1-tricontene				
	Δ_vH	(462–721)	115.4	477		[1999DYK/SVO]
C ₃₀ H ₆₀	[37590-56-0]	1,1,4,4,12,12,15,15-octamethylcycloodocosane				
	$\Delta_{\text{fus}}H$		58.58	411.2		[1972BOR/DAL]
C ₃₀ H ₆₀	[72443-19-7]	15-triacontene				
	$\Delta_{\text{us}}H$		30.96	324.2		
	$\Delta_{\text{fus}}H$		49.79	325.2	DSC	[2004TYA/BIS]
C ₃₀ H ₆₀ O ₄	[56444-66-7]	2,2,6,6,9,9,13,13,17,17,20,20-dodecamethyl-1,3,12,14-tetraoxacyclodocosane				
	$\Delta_{\text{fus}}H$		57.3	406.4		[1975BOR]
C ₃₀ H ₆₀ O ₁₅	[109635-67-8]	45-crown-15				
	$\Delta_{\text{fus}}H$		70.6	311.2	DSC	[1996YAN/YU]
C ₃₀ H ₆₁ Br	[62108-44-5]	1-bromotricosane				
	$\Delta_{\text{us}}H$		23.85	313.2		
	$\Delta_{\text{fus}}H$		79.5	339.6		[1953HOF/DEC]
C ₃₀ H ₆₂	[111-01-3]	2,6,10,15,19,23-hexamethyltetracosane (squalane)				
	Δ_vH	(363–513)	116.2	378	A	[1987STE/MAL]
C ₃₀ H ₆₂	[55319-83-0]	9-octyldocosane				
	Δ_vH	(518–588)	109.3	533	A	[1987STE/MAL]
C ₃₀ H ₆₂	[638-68-6]	triacontane				
	$\Delta_{\text{us}}H$		37.49	111.82		
	$\Delta_{\text{fus}}H$		68.83	338.7		[1996DOM/HEA]
	Δ_vH	(434–539)	152.3	298	CGC	[2004CHI/HAN]
	Δ_vH		164.5 ± 0.4	298	CGC	[2000NIC/ORF]
	Δ_vH	(422–487)	143 ± 2	454	TE	[1994PIA/FON]
	Δ_vH	(495–723)	111.3	510	A, E	[1987STE/MAL, 1966KUD/ZWO]
C ₃₀ H ₆₂	[1560-75-4]	2-methylnonacosane				
	Δ_vH	(461–718)	116.8	476		[1999DYK/SVO]
C ₃₀ H ₆₂ S	[66213-99-8]	1-triacontanethiol				
	Δ_vH	(488–751)	124	503	E	[1999DYK/SVO]
C ₃₀ H ₆₃ N	[2869-34-3]	tridecylamine				
	Δ_vH	(545–759)	76.8	560	A	[1987STE/MAL]
C ₃₁ H ₁₅ NO ₃	[3271-76-9]	C.I. Vat Green 3				
	$\Delta_{\text{sub}}H$	(519–634)	155	577	GS	[1986NIS/AND]
C ₃₁ H ₃₂ O ₂ P ₂	[32305-98-9]	(–) 2,3-O-isopropylidene-2,3-dihydroxy-1,4-bis(diphenylphosphino)butane				
	$\Delta_{\text{us}}H$	(78–380)	3.42	348.7		
	$\Delta_{\text{fus}}H$	(78–380)	38.61	364.2	AC	[2000WU/TAN]

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₃₁ H ₃₄	[56247-76-8] $\Delta_v H$	1,1-di(1-naphthyl)-1-undecene (518–588)	109.3	533	A	[1987STE/MAL]
C ₃₁ H ₄₃ NO ₅	[171018-28-3] $\Delta_{\text{fus}} H$	3-(acetyloxy)-17-(cyclopropylmethyl)- α -(1,1-dimethylethyl)-4,5-epoxy-18,19-dihydro-6-methoxy- α -methyl-6,14-ethenomorphinan-7-methanol	22.4	440.3		[1995STI/DUA]
C ₃₁ H ₄₄ O ₂	[na] $\Delta_{\text{fus}} H$	3,3'-bis-(1-cyclohexylethyl)-5,5'-dimethyldiphenylmethane-2,2'-diol	29.29	400.7		[1972INO/LIA]
C ₃₁ H ₄₈	[55319-81-8] $\Delta_v H$	1-(1-decylundec-1-enyl)naphthalene (499–567)	105.1	514	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₃₁ H ₅₂ O ₂	[1180-43-4] $\Delta_{\text{fus}} H$	cholesteryl α -methylpropionate 25.2		400.7		[1981YAN/NAB]
C ₃₁ H ₅₂ O ₃	[58-95-7] $\Delta_v H$	α -tocopherol acetate (466–524)	60.1 \pm 1.3	496		[1988BAG/GUR]
C ₃₁ H ₅₆	[55373-96-1] $\Delta_v H$	1,1-bis(decahydro-1-naphthyl)undecane (525–561)	110.5	540	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₃₁ H ₅₆	[61828-06-6] $\Delta_v H$	pentacosylbenzene (472–741)	117.5	487		[1999DYK/SVO]
C ₃₁ H ₅₆	[6006-90-2] $\Delta_v H$	13-phenylpentacosane (495–560)	106.7	510	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₃₁ H ₆₀	[55320-00-8] $\Delta_v H$	1-(1-decylundecyl)decahydronaphthalene (523–560)	107	538	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₃₁ H ₅₈ N ₆ O ₉	[326813-24-5] $\Delta_{\text{fus}} H$	caprolyamide deferoxamine 119.2		450.1	DSC	[2000IHN/VEN]
C ₃₁ H ₆₂	[18435-54-6] $\Delta_v H$	1-hentriacontene (468–730)	117.7	483		[1999DYK/SVO]
C ₃₁ H ₆₂	[61828-10-2] $\Delta_v H$	pentacosylcyclohexane (472–741)	116.6	487		[1999DYK/SVO]
C ₃₁ H ₆₂	[6697-15-0] $\Delta_v H$	13-cyclohexylpentaconsane (495–560)	106.7	510	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₃₁ H ₆₂ O	[502-73-8] $\Delta_{\text{fus}} H$	16-hentricontanone 117.1		356.4		[1994NAK/TAK]
C ₃₁ H ₆₄	[1560-72-1] $\Delta_v H$	2-methyltriacontane (468–726)	118.8	483		[1999DYK/SVO]
C ₃₁ H ₆₄	[55320-06-4] $\Delta_{\text{fus}} H$	11-decylheneicosane 71.13		282.3		[1996DOM/HEA]
	$\Delta_v H$	(298–313)	110.9	305	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₃₁ H ₆₄	[630-04-6] $\Delta_v H$	hentriacontane (534–565)	157.3	298	CGC	[2004CHI/HAN2]
	$\Delta_v H$	(433–474)	146 \pm 2	450	TE	[1994PIA/FON]
	$\Delta_v H$	(503–732)	113.8	518	A, E	[1987STE/MAL, 1966KUD/ZWO]
C ₃₁ H ₆₄ S	[534-24-9] $\Delta_v H$	1-hentriacontanethiol (494–759)	125.7	509	E	[1999DYK/SVO]
C ₃₂ H ₂ Br ₁₆ N ₈	[28746-04-5]	hexadecabromophthalocyanine				

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{sub}}H$	(438–493)	109.2 ± 16.3	453	ME	[1987STE/MAL, 1970BON/CAT]
C ₃₂ H ₂ Cl ₁₆ N ₈	[28888-81-5]	hexadecachlorophthalocyanine				
	$\Delta_{\text{sub}}H$	(398–443)	141.0 ± 17.6	413	ME	[1987STE/MAL, 1970BON/CAT]
C ₃₂ H ₁₄	[190-26-1]	ovalene				
	$\Delta_{\text{fus}}H$		8.08	729		
	$\Delta_{\text{fus}}H$		17.4	770.1		[1979SMI2]
	$\Delta_{\text{sub}}H$		211.7 ± 7.9	600	ME	[1952INO/SHI]
C ₃₂ H ₁₈ N ₈	[574-93-6]	β -29H,31H-phthalocyanine				
	$\Delta_{\text{sub}}H$	(598–698)	223.8 ± 1.3		ME	[2000SEM/BAS]
C ₃₂ H ₃₄	[116422-69-6]	1,8-bis-(4-biphenyl)octane				
	$\Delta_{\text{fus}}H$		56	415.2	DSC	[1989MAL/KAN]
C ₃₂ H ₃₄	[116422-70-9]	1,8-bis-[4-(4'-ethylbiphenyl)]butane				
	$\Delta_{\text{fus}}H$		46	454.2	DSC	[1989MAL/KAN]
C ₃₂ H ₃₈ N ₄	[1154424-99-3]	1,4-bis((1-benzylpiperidin-4-ylimino)methyl)benzene				
	$\Delta_{\text{fus}}H$		38.7	427.2	DSC	[2008STI/CIN]
C ₃₂ H ₃₈ O ₆	[99022-53-4]	hexa-2,4-diyne-1,6-diyl-bis(4-hexyloxybenzoate)				
	$\Delta_{\text{fus}}H$		49.27	334.2		[1990BEL/BAL]
C ₃₂ H ₃₉ ClO ₃	[71203-42-4]	norethindrone-6-(4-chlorophenyl)hexanoate				
	$\Delta_{\text{fus}}H$		28.8	413		[1996DOM/HEA]
C ₃₂ H ₄₁ F ₂₅	[89109-72-8]	1,1,1,2,2,3,3,4,4,5,5,6,6,7,7,8,8,9,9,10,10,11,11,12,12-pentacosofluorodotriacontane				
	$\Delta_{\text{fus}}H$		43.4	369.2	DSC	[1986RUS/RAB]
C ₃₂ H ₄₁ NO ₂	[50679-08-8]	α -[4-(1,1-dimethylethyl)phenyl]-4-(hydroxydiphenylmethyl)-1-piperidinebutanol (terfenadine)				
	$\Delta_{\text{fus}}H$		58.1	422.8	DSC	[2007BER/WAS]
C ₃₂ H ₄₅ NO ₅	[171018-29-4]	17-(cyclopropylmethyl)- α -(1,1-dimethylethyl)-4,5-epoxy-18,19-dihydro-6-methoxy- α -methyl (1-oxopropoxy)-6,14-ethenomorphinan-7-methanol				
	$\Delta_{\text{fus}}H$		27.1	410.2		[1995STI/DUA]
C ₃₂ H ₅₀	[85668-75-3]	2,4,5,7-tetramethyl-4,5-bis-(4-tert-butylphenyl)-octane				
	$\Delta_{\text{sub}}H$		182.8		E,B	[1983KRA/BEC]
C ₃₂ H ₅₀	[85668-73-1]	4,5-diethyl-4,5-bis-(4-tert-butylphenyl)-octane				
	$\Delta_{\text{sub}}H$		182.4		E,B	[1983KRA/BEC]
C ₃₂ H ₅₂ N ₆ O ₉	[105185-40-8]	benzoylamide deferroxamine				
	$\Delta_{\text{fus}}H$		107	453.8	DSC	[2000IHN/VEN]
C ₃₂ H ₅₈	[13024-80-1]	hexacosylbenzene				
	Δ_vH	(478–749)	119.6	493		[1999DYK/SVO]
C ₃₂ H ₆₀ O ₂	[na]	5,5,9,9,17,17,21,21-octamethylcyclotetrasane-1,13-dione				
	$\Delta_{\text{fus}}H$		32.6	380.2		[1972BOR/DAL2]
C ₃₂ H ₆₄	[18435-55-7]	1-dotriacontene				
	Δ_vH	(474–738)	119.8	489		[1999DYK/SVO]
C ₃₂ H ₆₄	[61828-11-3]	hexacosylcyclohexane				
	Δ_vH	(478–749)	118.6	493		[1999DYK/SVO]
C ₃₂ H ₆₄ O ₂	[7505-12-6]	ethyl triacontanoate				
	$\Delta_{\text{fus}}H$		16.2	334.7		
	$\Delta_{\text{fus}}H$		36.07	341.5		[1996DOM/HEA]

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound					
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference	
C ₃₂ H ₆₄ O ₄	[56444-67-8] $\Delta_{\text{fus}}H$	2,2,6,6,10,10,14,14,18,18,22,22-dodecamethyl-1,3,13,15-tetraoxacyclotetracosane	39.7	342.5		[1975BOR]	
C ₃₂ H ₆₄ O ₁₆	[71092-61-0] $\Delta_{\text{fus}}H$	48-crown-16	59.1	312.2	DSC	[1996YAN/YU]	
C ₃₂ H ₆₆	[55401-55-3] $\Delta_{\text{v}}H$	11-decyldocosane (523–559)	108.7	538	A, MG	[1987STE/MAL, 1955SCH/WHI]	
C ₃₂ H ₆₆	[544-85-4] $\Delta_{\text{us}}H$	dotriacontane	40.59	339.1			
	$\Delta_{\text{fus}}H$		79.74	341.9	DSC	[2003TOZ/INA]	
	$\Delta_{\text{us}}H$		41.38	338.7			
	$\Delta_{\text{fus}}H$		76.57	343.5		[1996DOM/HEA]	
	$\Delta_{\text{sub}}H$		271.1 ± 2.5			[1970COX/PIL]	
	$\Delta_{\text{v}}H$		(534–565)	162.5	298	CGC	[2004CHI/HAN2]
	$\Delta_{\text{v}}H$		(437–477)	147 ± 1	456	TE	[1994PIA/FON]
C ₃₂ H ₆₆	$\Delta_{\text{v}}H$	(361–395)	130.5	376	A	[1987STE/MAL]	
	$\Delta_{\text{v}}H$	(510–741)	116	535	A, E	[1987STE/MAL, 1966KUD/ZWO]	
C ₃₂ H ₆₆	[55401-54-2] $\Delta_{\text{v}}H$	9-octyltetracosane (501–563)	114.8	516	A, MG	[1987STE/MAL, 1955SCH/WHI]	
C ₃₂ H ₆₆	[1720-12-3] $\Delta_{\text{v}}H$	2-methylhentriacontane (474–735)	120.9	489		[1999DYK/SVO]	
C ₃₂ H ₆₆ O	[4113-12-6] $\Delta_{\text{us}}H$	17-oxatritriacontane	10.46	323.2			
	$\Delta_{\text{fus}}H$		116.73	324.7	DSC	[2004TYA/BIS]	
C ₃₂ H ₆₆ S	[66256-05-1] $\Delta_{\text{v}}H$	1-dotriacontanethiol (499–766)	127.5	514	E	[1999DYK/SVO]	
C ₃₃ H ₃₄ O ₃	[71203-40-2] $\Delta_{\text{fus}}H$	norethindrone-biphenyl-4-carboxylate	31.6	462		[1996DOM/HEA]	
C ₃₃ H ₄₀ O ₃	[71203-41-3] $\Delta_{\text{fus}}H$	norethindrone-4-cyclohexybenzoate	38.6	482		[1996DOM/HEA]	
C ₃₃ H ₄₆ O ₄	[128805-69-6] $\Delta_{\text{fus}}H$	17 β -4-heptoxybenzoyloxy testosterone	22.0	373	DSC	[1990YAN/EIR]	
C ₃₃ H ₄₇ NO ₅	[171018-30-7] $\Delta_{\text{fus}}H$	17-(cyclopropylmethyl)- α -(1,1-dimethylethyl)-4,5-epoxy-18,19-dihydro-6-methoxy- α -methyl-(1-oxobutoxy)-6,14-ethenomorphinan-7-methanol	32.4	422.1		[1995STI/DUA]	
C ₃₃ H ₄₈ O ₃	[71203-38-8] $\Delta_{\text{fus}}H$	norethindrone- <i>trans</i> -3-(4-butylcyclohexyl)propionate	22.5	374		[1996DOM/HEA]	
C ₃₃ H ₄₈ O ₃	[71203-37-7] $\Delta_{\text{fus}}H$	norethindrone- <i>trans</i> -hexylcyclohexylcarboxylate	22.6	398		[1996DOM/HEA]	
C ₃₃ H ₅₄ N ₆ O ₉	[326813-28-9] $\Delta_{\text{fus}}H$	phenylacetamide deferoxamine	119	447.7	DSC	[2000IHN/VEN]	
C ₃₃ H ₅₄ O ₆	[3319-31-1] $\Delta_{\text{v}}H$	tri(2-ethylhexyl)trimellitate (331–371)	81.1	346	ME	[2000LIA/MA]	
C ₃₃ H ₅₄ O ₆	[27251-75-8] $\Delta_{\text{v}}H$	triisooctyltrimellitate (331–372)	79.0	346	ME	[2000LIA/MA]	

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
C ₃₃ H ₆₀	[61828-25-9] $\Delta_{\text{v}}H$	heptacosylbenzene (484–756)	121.5	499		[1999DYK/SVO]
C ₃₃ H ₆₂ N ₆ O ₉	[99899-52-2] $\Delta_{\text{fus}}H$	octanoylamide deferoxamine	130.7	455.3	DSC	[2000IHN/VEN]
C ₃₃ H ₆₂ O ₆	[621-71-6] $\Delta_{\text{v}}H$	glycerol tricaprte (437–485)	130.5	411	TGA	[1990KIS/SHO]
	$\Delta_{\text{v}}H$		154.6 ± 5.4	298	TGA	[1990KIS/SHO]
	$\Delta_{\text{v}}H$		124.6	452	A	[1987STE/MAL]
C ₃₃ H ₆₆	[61828-12-4] $\Delta_{\text{v}}H$	heptacosylcyclohexane (484–757)	120.6	499		[1999DYK/SVO]
C ₃₃ H ₆₆	[61868-11-9] $\Delta_{\text{v}}H$	1-tritriacontene (480–746)	121.8	495		[1999DYK/SVO]
C ₃₃ H ₆₈	[630-05-7] $\Delta_{\text{fus}}H$	tritriacontane (534–565)	105.02	344		[1996DOM/HEA]
	$\Delta_{\text{v}}H$		167.6	298	CGC	[2004CHI/HAN2]
	$\Delta_{\text{v}}H$		148 ± 1	458	TE	[1994PIA/FON]
	$\Delta_{\text{v}}H$		118	532	A, E	[1987STE/MAL, 1966KUD/ZWO]
C ₃₃ H ₆₈	[1720-11-2] $\Delta_{\text{v}}H$	2-methyldotriacontane (480–743)	122.9	495		[1999DYK/SVO]
C ₃₃ H ₆₈ S	[66214-20-8] $\Delta_{\text{v}}H$	1-tritriacontanethiol (504–773)	129.1	519	E	[1999DYK/SVO]
C ₃₄ H ₁₆ O ₂	[116-71-2] $\Delta_{\text{sub}}H$	dibenzanthrone (violanthrone) (513–548)	208.8	528	A	[1987STE/MAL]
	$\Delta_{\text{sub}}H$		202.9	542	ME	[1951INO]
C ₃₄ H ₁₆ O ₂	[128-64-3] $\Delta_{\text{sub}}H$	isodibenzanthrone (isoviolanthrone) (523–553)	221.1	538		[1987STE/MAL]
	$\Delta_{\text{sub}}H$		215.5	537	ME	[1951INO]
C ₃₄ H ₁₈	[190-93-2] $\Delta_{\text{sub}}H$	benzo[<i>rst</i>]phenanthro[1,10,9- <i>cde</i>]pentaphene (478–603)	154.1	493	A	[1987STE/MAL]
C ₃₄ H ₁₈	[81-31-2] $\Delta_{\text{sub}}H$	violanthrene	223.8 ± 8.8	590		[1952INO/SHI, 1960JON]
C ₃₄ H ₁₈	[na] $\Delta_{\text{sub}}H$	violanthrene A (mp 478 °C) (562–724)	195.8	653	ME	[1967WAK/INO]
C ₃₄ H ₁₈	[na] $\Delta_{\text{sub}}H$	violanthrene B (mp 330 °C) (555–625)	153.5	590	ME	[1976WAK/INO]
Note: This entry is likely the original reference for benzo[<i>rst</i>]-phenanthro[1,10,9- <i>cde</i>]pentaphene listed in Ref. [1987STE/MAL]. Chemical Abstracts cites [1967WAK/INO] as reporting the heat of sublimation for benzo[<i>rst</i>]phenanthro[1,10,9- <i>cde</i>]pentaphene.						
C ₃₄ H ₁₈	[4430-29-9] $\Delta_{\text{sub}}H$	isoviolanthrene A (mp 510 °C) (588–724)	218	590	ME	[1952INO/SHI, 1960JON]
C ₃₄ H ₁₈	[191-79-7] $\Delta_{\text{sub}}H$	tetrabenzo[<i>de,hi,op,st</i>]pentacene (348–448)	118.5	363	A	[1987STE/MAL]
	$\Delta_{\text{sub}}H$		118	400	ME	[1967WAK/INO]
C ₃₄ H ₃₁ ClN ₂ O ₃	[119887-41-1]	spiro[isobenzofuran-1(3 <i>H</i>),9'(9 <i>H</i>)-7'-chloro-6'-(methylcyclohexylamino)-3'-methyl-2'-anilinoxanthene]-3-one				

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{fus}}H$		49.0	442.2	DSC	[1988NAK/KIT]
C ₃₄ H ₃₂ N ₂ O ₃	[55250-84-5]	spiro[isobenzofuran-1(3 <i>H</i>),9'(9 <i>H</i>)-6'-(methylcyclohexylamino)-3'-methyl-2'-anilinoxanthene]-3-one				
	$\Delta_{\text{fus}}H$		39.9	476.2	DSC	[1988NAK/KIT]
C ₃₄ H ₃₈	[116422-71-0]	1,6- <i>bis</i> -[4-(4'-ethylbiphenyl)]hexane				
	$\Delta_{\text{us}}H$		3.9	393.2		
	$\Delta_{\text{fus}}H$		35	422.2	DSC	[1989MAL/KAN]
C ₃₄ H ₄₉ NO ₅	[171018-31-8]	17-(cyclopropylmethyl)- α -(1,1-dimethylethyl)-4,5-epoxy-18,19-dihydro-6-methoxy- α -methyl-(1-oxopentoxo)-6,14-ethenomorphinan-7-methanol				
	$\Delta_{\text{fus}}H$		24.0	379.1		[1995STI/DUA]
C ₃₄ H ₅₂ N ₂ O ₄	[32687-78-8]	N,N'- <i>bis</i> [3-(3,5-di- <i>tert</i> -butyl-4-hydroxyphenyl)propionyl]hydrazine				
	$\Delta_{\text{us}}H$		19.47	474.1		
	$\Delta_{\text{fus}}H$		41.5	503	DSC	[2008COG/HIL]
C ₃₄ H ₅₄	[85668-72-0]	4,5-dipropyl-4,5- <i>bis</i> -(4- <i>tert</i> -butylphenyl)-octane				
	$\Delta_{\text{fus}}H$		40.58	419	DSC	[1983KRA/BEC]
	$\Delta_{\text{sub}}H$		198.3		E,B	[1983KRA/BEC]
C ₃₄ H ₆₂	[61828-26-0]	octacosylbenzene				
	Δ_vH	(490–764)	123.4	505		[1999DYK/SVO]
C ₃₄ H ₆₆ O ₄	[2432-88-4]	didodecyl sebacate				
	Δ_vH		131.9	420	TGA	[1990KIS/SHO]
	Δ_vH		154.5 ± 5.4	298	TGA	[1990KIS/SHO]
C ₃₄ H ₆₈	[61828-13-5]	octacosylcyclohexane				
	Δ_vH	(490–764)	122.4	505		[1999DYK/SVO]
C ₃₄ H ₆₈	[61868-12-0]	1-tetratriacontene				
	Δ_vH	(486–754)	123.7	501		[1999DYK/SVO]
C ₃₄ H ₆₈	[87292-56-6]	17-tetratriacontene				
	$\Delta_{\text{us}}H$		33.47	332.8		
	$\Delta_{\text{fus}}H$		51.46	334.3	DSC	[2004TYA/BIS]
C ₃₄ H ₆₈ O ₁₇	[109635-68-9]	51-crown-17				
	$\Delta_{\text{fus}}H$		66.6	301.2	DSC	[1996YAN/YU]
C ₃₄ H ₇₀	[55429-84-0]	11-decyltetracosane				
	Δ_vH	(537–574)	113.1	552	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₃₄ H ₇₀	[55429-83-9]	9-octylhexacosane				
	Δ_vH	(537–575)	110.3	552	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₃₄ H ₇₀	[14167-59-0]	tetratriacontane				
	$\Delta_{\text{us}}H$		0.41	342.8		
	$\Delta_{\text{us}}H$		46.65	343.5		
	$\Delta_{\text{fus}}H$		95.64	346.1	DSC	[2006WAN/TOZ]
	$\Delta_{\text{us}}H$		29.29	341.5		
	$\Delta_{\text{fus}}H$		79.96	345.9		[1996DOM/HEA]
	Δ_vH	(534–565)	172.7	298	CGC	[2004CHI/HAN2]
	Δ_vH	(446–497)	152 ± 2	471	TE	[1994PIA/FON]
	Δ_vH	(372–402)	149.7	387	A	[1987STE/MAL]
	Δ_vH	(523–756)	120.3	538	A, E	[1987STE/MAL, 1966KUD/ZWO]
C ₃₄ H ₇₀	[66214-27-5]	2-methyltritiacontane				
	Δ_vH	(486–750)	124.8	501		[1999DYK/SVO]

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₃₄ H ₇₀ S	[66214-28-6] $\Delta_v H$	1-tetratriacontanethiol (509–780)	130.7	524	E	[1999DYK/SVO]
C ₃₅ H ₂₈ Cl ₂ N ₈ O ₄	[84625-61-6] $\Delta_{\text{fus}} H$	4-[4-[4-[4-[[2-(2,4-dichlorophenyl)-2-(1 <i>H</i> -1,2,4-triazol-1-ylmethyl)-1,3-dioxolan-4-yl]methoxy]phenyl]-1-piperazinyl]phenyl]-2,4-dihydro-2-(1-methylpropyl)-3 <i>H</i> -1,2,4-triazol-3-one (itraconazole)	69.9	438.6	DSC	[2007BER/WAS]
C ₃₅ H ₃₇ N ₃	[500362-45-8] $\Delta_{\text{fus}} H$	4-butylphenyl-[6-(4-butylphenyl)methyl-9-methyl-9 <i>H</i> -carbazol-3-ylmethylene]amine	29.6	460.2	DSC	[2002BEL/MAN]
C ₃₅ H ₃₇ N ₃ O ₂	[500362-50-5] $\Delta_{\text{fus}} H$	4-butoxyphenyl-[6-(4-butoxyphenyl)methyl-9-methyl-9 <i>H</i> -carbazol-3-yl-methylene]amine	34.4	475.2	DSC	[2002BEL/MAN]
C ₃₅ H ₅₀ N ₂ O ₈	[874908-00-6] $\Delta_{\text{fus}} H$	2-methylacrylic acid 11-[4-(6,7,9,10,12,13,15,16-octahydro-5,8,11,14,17-pentaoxabenzocyclopentadecen-2-ylazo)phenoxy]undecyl ester	48.7	379.7	DSC	[2005NIS/WAT]
C ₃₅ H ₅₁ NO ₅	[171018-32-9] $\Delta_{\text{fus}} H$	17-(cyclopropylmethyl)- α -(1,1-dimethylethyl)-4,5-epoxy-18,19-dihydro-6-methoxy- α -methyl-(1-oxohexyloxy)-6,14-ethenomorphinan-7-methanol	22.6	352.6		[1995STI/DUA]
C ₃₅ H ₆₄	[56247-97-3] $\Delta_v H$	15-phenylnonacosane (523–550)	126.5	536	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₃₅ H ₆₄	[61828-27-1] $\Delta_v H$	nonacosylbenzene (495–771)	125.4	510		[1999DYK/SVO]
C ₃₅ H ₆₈ O ₄	[818-21-3] $\Delta_{\text{fus}} H$	1,3-propanediol, dipalmitate	133	329.8	DSC	[2007ABE/BOU]
C ₃₅ H ₇₀	[61828-14-6] $\Delta_v H$	nonacosylcyclohexane (495–771)	124.4	510		[1999DYK/SVO]
C ₃₅ H ₇₀	[61868-13-1] $\Delta_v H$	1-pentatriacontene (492–761)	125.5	507		[1999DYK/SVO]
C ₃₅ H ₇₀	[55521-27-2] $\Delta_v H$	15-cyclohexylnonacosane (548–581)	129	563	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₃₅ H ₇₂	[630-07-9] $\Delta_{\text{fus}} H$ $\Delta_{\text{fus}} H$ $\Delta_v H$ $\Delta_v H$	pentatriacontane (534–565) (529–764)	41.09 86.4 178 122.4	344.7 347.2 298 544	CGC A, E	[1996DOM/HEA] [2004CHI/HAN2] [1987STE/MAL, 1966KUD/ZWO]
C ₃₅ H ₇₂	[14167-65-8] $\Delta_v H$	2-methyltetratriacontane (491–758)	126.9	506		[1999DYK/SVO]
C ₃₅ H ₇₂ S	[66576-86-1] $\Delta_v H$	1-pentatriacontanethiol (514–787)	132.2	529	E	[1999DYK/SVO]
C ₃₆ H ₁₈	[191-48-0] $\Delta_{\text{fus}} H$	decacyclene	25.4	666		[1980SMI]
C ₃₆ H ₂₄	[7059-70-3] $\Delta_{\text{fus}} H$	1,3,5-tri- α -naphthylbenzene	42.26	472		[1967MAG]
C ₃₆ H ₃₂ N ₂ O ₄	[158547-46-7] $\Delta_{\text{fus}} H$ $\Delta_{\text{fus}} H$	3,3',4,4'-biphenyltetracarboxy- <i>N,N'</i> -bis-(4- <i>n</i> -butylphenyl)diimide	33.5 8.4	537.9 556.3		[1995EIS/DEN]
C ₃₆ H ₄₂	[116422-72-1]	1,8-bis-[4-(4'-ethylbiphenyl)]octane				

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₃₆ H ₄₂	$\Delta_{\text{us}}H$		8.4	402.2		
	$\Delta_{\text{fus}}H$		42	413.2	DSC	[1989MAL/KAN]
	[116445-91-1]	1,4- <i>bis</i> -[4-(4'- <i>n</i> -butylbiphenyl)]butane				
C ₃₆ H ₄₄ N ₂ S ₄	$\Delta_{\text{us}}H$ (liq <i>cryst</i>)		12	404.2		
	$\Delta_{\text{us}}H$ (liq <i>cr-liq</i>)		24	464.2	DSC	[1989MAL/KAN]
	[109537-97-5]	<i>bis</i> -[4-(5-heptyl-2-thienylmethylidenamino)phenyl]disulfide				
C ₃₆ H ₄₆ O ₄	$\Delta_{\text{fus}}H$		36.1	356.2		[1978KOS/BUD]
	[92341-28-1]	4,4'-didecanoyloxydiphenyldiacetylene				
	$\Delta_{\text{us}}H$		44.9	308		
C ₃₆ H ₅₃ NO ₅	$\Delta_{\text{fus}}H$		42.2	403		[1996DOM/HEA]
	[171018-33-0]	17-(cyclopropylmethyl)- α -(1,1-dimethylethyl)-4,5-epoxy-18,19-dihydro-6-methoxy- α -methyl-(1-oxoheptyloxy)-6,14-ethenomorphinan-7-methanol				
	$\Delta_{\text{fus}}H$		19.3	360		[1995STI/DUA]
C ₃₆ H ₅₄ O ₁₂	[65201-68-5]	benzene hexa- <i>n</i> -pentanoate				
	$\Delta_{\text{us}}H$	(13–390)	8.8	173.1		
	$\Delta_{\text{us}}H$	(13–390)	15.3	313.2		
	$\Delta_{\text{us}}H$	(13–390)	1.4	349.9		
C ₃₆ H ₆₀ O ₆	$\Delta_{\text{fus}}H$	(13–390)	30.3	379.5	AC	[2001ASA/SOR]
	[53894-23-8]	triisononyl trimellitate				
	Δ_vH	(334–372)	102.2	349	ME	[2000LIA/MA]
C ₃₆ H ₆₂ O ₄	[2915-60-8]	ditetradecyl phthalate				
	Δ_vH	(416–465)	126	431	T	[1987STE/MAL, 1949PER/WEB]
C ₃₆ H ₆₄ O ₄	[175848-69-8]	2,5-di- <i>n</i> -pentadecyloxy-1,4-benzoquinone				
	$\Delta_{\text{us}}H$		21.7	385.9		
	$\Delta_{\text{fus}}H$		101.7	393.5	DSC	[1996KEE/VAN]
C ₃₆ H ₆₆	[50715-02-1]	triacontylbenzene				
	Δ_vH	(501–778)	127	516		[1999DYK/SVO]
C ₃₆ H ₇₀ O ₄	[26719-63-1]	1,4-butanediol dipalmitate				
	$\Delta_{\text{fus}}H$		162.4	339	DSC	[2008ZHA/WUM]
C ₃₆ H ₇₂	[61868-14-2]	1-hexatriacontene				
	Δ_vH	(497–768)	127.4	512		[1999DYK/SVO]
C ₃₆ H ₇₂	[61828-15-7]	triacontylcyclohexane				
	Δ_vH	(500–778)	126.3	515		[1999DYK/SVO]
C ₃₆ H ₇₂ O ₁₈	[71092-62-1]	54-crown-18				
	$\Delta_{\text{fus}}H$		81.6	317.2	DSC	[1996YAN/YU]
C ₃₆ H ₇₄	[630-06-8]	hexatriacontane				
	$\Delta_{\text{us}}H$		11.73	347.3		
	$\Delta_{\text{us}}H$		24.72	348.3		
	$\Delta_{\text{fus}}H$		81.6	350.2	DSC	[2006KHI/BOU]
	$\Delta_{\text{us}}H$		0.47	343.6		
	$\Delta_{\text{us}}H$		15.02	348.8		
	$\Delta_{\text{us}}H$		28.52	347.1		
	$\Delta_{\text{fus}}H$		102.51	349.2	DSC	[2006WAN/TOZ]
	$\Delta_{\text{us}}H$		30.3	347		
	$\Delta_{\text{fus}}H$		87.6	349	DSC	[2004MAR/KAI]
	$\Delta_{\text{us}}H$	(80–370)	10.1	345.4		
	$\Delta_{\text{us}}H$	(80–370)	32.1	346.8		

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{fus}}H$	(80–370)	87.5	348.9	AC	[1999WAN/TAN]
	$\Delta_{\text{trs}}H$		9.92	345.4		
	$\Delta_{\text{trs}}H$		30.54	347.1		
	$\Delta_{\text{fus}}H$		88.83	349.2		[1996DOM/HEA]
	Δ_vH	(534–565)	182.8	298	CGC	[2004CHI/HAN2]
	Δ_vH	(452–516)	157 ± 2	484	TE	[1994PIA/FON]
	Δ_vH	(535–571)	124.4	550	A, E	[1987STE/MAL, 1966KUD/ZWO]
C ₃₆ H ₇₄	[55517-89-0]	13-undecylpentacosane				
	Δ_vH	(548–580)	132.9	563	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₃₆ H ₇₄	[66576-73-6]	2-methylpentatriacontane				
	Δ_vH	(497–765)	128.7	512		[1999DYK/SVO]
C ₃₆ H ₇₄	[67309-30-2]	18-methylpentatriacontane				
	$\Delta_{\text{fus}}H$		99	325.2	DSC	[2005IKE/YAM]
C ₃₆ H ₇₄ O	[6297-03-6]	19-oxaheptatriacontane				
	$\Delta_{\text{trs}}H$		10.88	333.2		
	$\Delta_{\text{fus}}H$		105.86	335.3	DSC	[2004TYA/BIS]
C ₃₆ H ₇₄ O ₁₈	[na]	1, ω -dimethoxyheptadeca(oxyethylene)				
	$\Delta_{\text{fus}}H$		136.6	301.2	DSC	[1996YAN/YU]
C ₃₆ H ₇₄ S	[66577-23-9]	1-hexatriacontanethiol				
	Δ_vH	(518–793)	134	533	E	[1999DYK/SVO]
C ₃₆ H ₇₅ N	[102-87-4]	tridodecylamine				
	Δ_vH	(579–807)	82.1	594	A	[1987STE/MAL]
C ₃₇ H ₄₈ N ₆ O ₅ S ₂	[155213-67-5]	2,4,7,12-tetraazatridecan-13-oic acid, 10-hydroxy-2-methyl-5-(1-methylethyl)-1-[2-(1-methylethyl)-4-thiazolyl]-3,6-dioxo-8,11-bis(phenylmethyl)-, 5-thiazolylmethyl ester (Ritonavir)				
	$\Delta_{\text{fus}}H$		57.9	395.5		[2002ZHO/ZHA]
C ₃₇ H ₅₂ O ₄	[129108-06-1]	3 β -n-octyloxy-17 β -butyloxybenzoyloxy estradiol				
	$\Delta_{\text{fus}}H$		27.0	332	DSC	[1990YAN/EIR]
C ₃₇ H ₆₈	[61828-28-2]	hexatriacontylbenzene				
	Δ_vH	(506–785)	128.8	521		[1999DYK/SVO]
C ₃₇ H ₆₈ O ₈	[7299-99-2]	pentaerythritol tetra-2-ethylhexanoate				
	Δ_vH	(355–443)	126.4	370		[2007RAZ/MOK]
C ₃₇ H ₇₀ O ₆	[30283-10-4]	1-caprylic-2-lauryl-3-myristic glycerol				
	Δ_vH	(464–526)	131.7	479	A, T	[1987STE/MAL, 1949PER/WEB2]
C ₃₇ H ₇₄	[61828-16-8]	hentriacontylcyclohexane				
	Δ_vH	(505–785)	128.1	520		[1999DYK/SVO]
C ₃₇ H ₇₄	[61868-15-3]	1-heptatriacontene				
	Δ_vH	(502–775)	129.2	517		[1999DYK/SVO]
C ₃₇ H ₇₆	[7194-84-5]	heptatriacontane				
	Δ_vH	(534–565)	187.5	298	CGC	[2004CHI/HAN2]
	Δ_vH	(471–511)	155 ± 2	491	TE	[1994PIA/FON]
	Δ_vH	(541–778)	126.4	556	A, E	[1987STE/MAL, 1966KUD/ZWO]
C ₃₇ H ₇₆	[66577-06-8]	2-methylhexatriacontane				
	Δ_vH	(502–772)	130.5	517		[1999DYK/SVO]
C ₃₇ H ₇₆ S	[66577-07-9]	1-heptatriacontanethiol				

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(523–799)	135.3	538	E	[1999DYK/SVO]
C ₃₈ H ₃₀	[18909-18-7]	1-diphenylmethylene-4-triphenylmethyl-2,5-cyclohexadiene				
	$\Delta_{\text{sub}} H$	(348–394)	114.6	363	A	[1987STE/MAL]
C ₃₈ H ₃₀ O ₂	[596-30-5]	<i>bis</i> (triphenylmethyl)peroxide				
	$\Delta_{\text{sub}} H$	(392–434)	158.1	407	A	[1987STE/MAL]
C ₃₈ H ₅₀ O ₄	[71332-86-0]	4,4'-diundecanoyloxydiphenyldiacetylene				
	$\Delta_{\text{us}} H$		18.1	339		
	$\Delta_{\text{us}} H$		7.59	359		
	$\Delta_{\text{fus}} H$		36.2	399		[1996DOM/HEA]
C ₃₈ H ₆₂	[85668-76-4]	5,6-dibutyl-5,6- <i>bis</i> -(4- <i>tert</i> -butylphenyl)-decane				
	$\Delta_{\text{fus}} H$		43.1	386	DSC	[1983KRA/BEC]
	$\Delta_{\text{sub}} H$		220.9		E,B	[1983KRA/BEC]
C ₃₈ H ₆₇ NO ₁₀	[150785-53-8]	8,9-didehydro-N-demethyl-9-deoxo-4'',6,12-trideoxy-6,9-epoxy-N-ethylerythromycin				
	$\Delta_{\text{fus}} H$		36.7	434.9		[2002ZHO/ZHA]
C ₃₈ H ₆₈ N ₂ O ₂	[312952-55-9]	N,N'-dihexadecanoylbenzene-1,2-diamine				
	$\Delta_{\text{us}} H$		78	367.2		
	$\Delta_{\text{fus}} H$		32	388.2	DSC	[2000AKU/IUC]
C ₃₈ H ₆₈ O ₄	[118476-26-9]	2,5-di- <i>n</i> -hexadecyloxy-1,4-benzoquinone				
	$\Delta_{\text{us}} H$		6.8	357.7		
	$\Delta_{\text{us}} H$		14.1	370.9		
	$\Delta_{\text{us}} H$		19	389		
	$\Delta_{\text{fus}} H$		83	394.2	DSC	[1996KEE/VAN]
C ₃₈ H ₆₈ S ₈	[105782-51-2]	2-[4,5- <i>bis</i> (octylthio)-1,3-dithiol-2-ylidene]-4,5- <i>bis</i> (octylthio)-1,3-dithiole				
	$\Delta_{\text{us}} H$	(10–330)	0.16	94.3		
	$\Delta_{\text{us}} H$	(10–330)	4.92	215.8		
	$\Delta_{\text{fus}} H$	(10–330)	89.3	322.5	AC	[1997TAN/ATA]
C ₃₈ H ₇₀	[61828-29-3]	dotriacontylbenzene				
	$\Delta_v H$	(511–791)	130.4	526		[1999DYK/SVO]
C ₃₈ H ₇₄ O ₄	[26719-47-1]	ditetradecyl sebacate				
	$\Delta_v H$		142.8	440	TGA	[1990KIS/SHO]
	$\Delta_v H$		170.6 ± 5.9	298	TGA	[1990KIS/SHO]
	$\Delta_v H$	(431–483)	135.5	446	A, T	[1987STE/MAL, 1949PER/WEB]
C ₃₈ H ₇₆	[61828-17-9]	dotriacontylcyclohexane				
	$\Delta_v H$	(510–792)	129.8	525		[1999DYK/SVO]
C ₃₈ H ₇₆	[61868-16-4]	1-octatriacontene				
	$\Delta_v H$	(507–782)	131	522		[1999DYK/SVO]
C ₃₈ H ₇₆	[62978-78-3]	19-octatriacontene				
	$\Delta_{\text{us}} H$		7.53	321		
	$\Delta_{\text{us}} H$		34.31	338.4		
	$\Delta_{\text{fus}} H$		66.53	340.8	DSC	[2004TYA/BIS]
C ₃₈ H ₇₈	[7194-85-6]	octatriacontane				
	$\Delta_v H$	(534–565)	192.6	298	CGC	[2004CHI/HAN2]
	$\Delta_v H$	(471–511)	160 ± 2	491	TE	[1994PIA/FON]
	$\Delta_v H$	(546–785)	128.5	561	A, E	[1987STE/MAL, 1966KUD/ZWO]
C ₃₈ H ₇₈	[66576-92-9]	2-methylheptatriacontane				

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
	$\Delta_{\text{v}}H$	(507–779)	132.2	522		[1999DYK/SVO]
C ₃₈ H ₇₈ S	[66576-93-0]	1-octatriacontanethiol				
	$\Delta_{\text{v}}H$	(527–805)	136.7	542	E	[1999DYK/SVO]
C ₃₉ H ₄₅ N ₃ O ₂	[500362-51-6]	4-hexyloxyphenyl-[6-(4-hexyloxyphenyl)methyl-9-methyl-9H-carbazol-3-yl-methylene]amine				
	$\Delta_{\text{fus}}H$		25.4	459.2	DSC	[2002BEL/MAN]
C ₃₉ H ₇₂	[55517-74-3]	17-phenyltritracontane				
	$\Delta_{\text{v}}H$	(544–571)	147.1	557	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₃₉ H ₇₂	[61828-30-6]	tritracontylbenzene				
	$\Delta_{\text{v}}H$	(516–798)	132	531		[1999DYK/SVO]
C ₃₉ H ₇₄ O ₆	[538-24-9]	glycerol trilaurate				
	$\Delta_{\text{fus}}H$		123.51	319.5		[1996DOM/HEA]
	$\Delta_{\text{v}}H$		221.1		TGA	[2002GOO/GEL]
	$\Delta_{\text{v}}H$		147.1	438	TGA	[1990KIS/SHO]
	$\Delta_{\text{v}}H$		180.0 ± 6.3	298	TGA	[1990KIS/SHO]
	$\Delta_{\text{v}}H$	(458–520)	137.6	473	A, T	[1987STE/MAL, 1949PER/WEB2]
C ₃₉ H ₇₆ O ₄	[17367-44-1]	1,3-propanediol, distearate				
	$\Delta_{\text{fus}}H$		110	329.8	DSC	[2007ABE/BOU]
C ₃₉ H ₇₈	[61868-17-5]	1-nontriacontene				
	$\Delta_{\text{v}}H$	(512–788)	132.5	527		[1999DYK/SVO]
C ₃₉ H ₇₈	[61828-18-0]	triatracontylcyclohexane				
	$\Delta_{\text{v}}H$	(515–798)	131.4	530		[1999DYK/SVO]
C ₃₉ H ₇₈	[55517-75-4]	17-cyclohexyltritracontane				
	$\Delta_{\text{v}}H$	(570–602)	131.9	585	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₃₉ H ₇₈ O	[22986-70-5]	20-nonatriacontanone				
	$\Delta_{\text{fus}}H$		153	365.8		[2000NAK/SHI]
	$\Delta_{\text{fus}}H$		153	365.6	DSC	[1994NAK/TAK]
C ₃₉ H ₈₀	[7194-86-7]	nonatriacontane				
	$\Delta_{\text{v}}H$	(552–791)	130.3	567	A, E	[1987STE/MAL, 1966KUD/ZWO]
C ₃₉ H ₈₀	[66576-59-8]	2-methyloctatriacontane				
	$\Delta_{\text{v}}H$	(512–785)	133.8	527		[1999DYK/SVO]
C ₃₉ H ₈₀	[857685-80-4]	18-butylpentatriacontane				
	$\Delta_{\text{fus}}H$		104	318.5	DSC	[2005IKE/YAM]
C ₃₉ H ₈₀ S	[66576-60-1]	1-nonatriacontanethiol				
	$\Delta_{\text{v}}H$	(531–811)	138.1	546	E	[1999DYK/SVO]
C ₄₀ H ₃₈ N ₄ O ₄	[130048-21-4]	bis-(4- <i>n</i> -butyl-1'-diazophenyl) isophthalate				
	$\Delta_{\text{fus}}H$		16.0	469.2	DSC	[1990JIN/KAN]
C ₄₀ H ₄₀ N ₂ O ₄	[158547-48-9]	3,3',4,4'-biphenyltetracarboxy- <i>N,N'</i> -bis-(4- <i>n</i> -hexylphenyl)diimide				
	$\Delta_{\text{us}}H$ (liq <i>cryst</i>)		19.9	432.4		
	$\Delta_{\text{us}}H$ (liq <i>cryst</i>)		26.2	513.8		
	$\Delta_{\text{us}}H$ (liq <i>cr-liq</i>)		9.5	563.3		[1995EIS/DEN]
C ₄₀ H ₄₆ O ₈	[161282-95-7]	6,7,9,10,12,13,15,16,18,19-decahydro-32,35-dimethoxy-31 <i>H</i> -4,21-(methano[1,3]benzenomethano)-26,30-metheno-25 <i>H</i> -dibenzo[<i>q,z</i>]-[1,4,7,10,13,16]hexaoxacycloheptacosin				
	$\Delta_{\text{sub}}H$		82 ± 2			[2008SUR]

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₄₀ H ₅₀	[116422-73-2]	1,8-bis-[4-(4'- <i>n</i> -butylbiphenyl)]octane				
	$\Delta_{\text{us}}H$ (liq <i>cryst</i>)		13.0	398.2		
	$\Delta_{\text{us}}H$ (liq <i>cr-liq</i>)		27.0	414.2	DSC	[1989MAL/KAN]
C ₄₀ H ₅₄ O ₄	[92341-29-2]	4,4'-didodecanoyloxydiphenyldiacetylene				
	$\Delta_{\text{us}}H$		50.2	374		
	$\Delta_{\text{fus}}H$		44.0	401	DSC	[1996DOM/HEA]
C ₄₀ H ₅₆	[7235-40-7]	β -carotene				
	$\Delta_{\text{fus}}H$		56.0	456		[2003TRE/KAS]
C ₄₀ H ₇₂ O ₄	[175848-70-1]	2,5-di-heptadecyloxy-1,4-benzoquinone				
	$\Delta_{\text{us}}H$		13.0	383.6		
	$\Delta_{\text{fus}}H$		120.9	395.3	DSC	[1996KEE/VAN]
C ₄₀ H ₇₄	[61828-31-7]	tetraatriacontylbenzene				
	Δ_vH	(520–804)	133.8	535		[1999DYK/SVO]
C ₄₀ H ₈₀	[61868-18-6]	1-tetracontene				
	Δ_vH	(517–794)	134	532		[1999DYK/SVO]
C ₄₀ H ₈₀	[61828-19-1]	tetraatriacontylcyclohexane				
	Δ_vH	(520–804)	132.9	535		[1999DYK/SVO]
C ₄₀ H ₈₂	[4181-95-7]	tetracontane				
	$\Delta_{\text{us}}H$		2.51	345		
	$\Delta_{\text{us}}H$		8.6	347.7		
	$\Delta_{\text{fus}}H$		143.94	354.6	DSC	[2006WAN/TOZ]
	$\Delta_{\text{us}}H$		14.02	345.4		
	$\Delta_{\text{fus}}H$		133.44	353.2	DSC	[1992LOU/ROU]
	Δ_vH	(323–523)	203.5 ± 0.2	298	CGC	[2008CHI/WAN]
	Δ_vH	(557–798)	132.2	572	A, E	[1987STE/MAL, 1966KUD/ZWO]
C ₄₀ H ₈₂	[66576-48-5]	2-methylnonatriacontane				
	Δ_vH	(517–791)	135.3	532		[1999DYK/SVO]
C ₄₀ H ₈₂	[304457-95-2]	20-methylnonatriacontane				
	$\Delta_{\text{fus}}H$		120	331.4		[2004YAM/NEM]
Note: DSC curve showed no solid-solid phase transition in the temperature range of T=(300 to 360) K						
C ₄₀ H ₈₂ S	[66576-49-6]	1-tetracontanethiol				
	Δ_vH	(535–817)	139.6	550	E	[1999DYK/SVO]
C ₄₁ H ₄₄ N ₂ O ₉	[832684-74-9]	3,3'-di(N-cyclopropylmethyl)-4,5-epoxy-14-hydroxymorphinan-6-one-3-yl)carbonate				
	$\Delta_{\text{fus}}H$		17.5	490.6	DSC	[2004HAM/HAM]
C ₄₁ H ₇₆	[61828-32-8]	pentatriacontylbenzene				
	Δ_vH	(525–810)	135.2	540		[1999DYK/SVO]
C ₄₁ H ₇₆ O ₈	[41058-87-1]	3,5,5-trimethylhexanoic acid, 2,2-bis[(3,5,5-trimethyl-1-oxohexyl)oxy]-methyl]-1,3-propanediyl ester				
	$\Delta_{\text{fus}}H$		51.3	304		[1996PYD/VAR]
C ₄₁ H ₈₂	[66576-37-2]	1-hentetracontene				
	Δ_vH	(521–800)	135.8	536		[1999DYK/SVO]
C ₄₁ H ₈₂	[61828-20-4]	pentatriacontylcyclohexane				
	Δ_vH	(524–810)	134.5	539		[1999DYK/SVO]
C ₄₁ H ₈₄	[7194-87-8]	hentetracontane				
	$\Delta_{\text{fus}}H$		147.2	357.5		[2000PAU/MEH]

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(562–804)	134.1	577	A, E	[1987STE/MAL, 1966KUD/ZWO]
C ₄₁ H ₈₄	[66575-38-3] $\Delta_v H$	2-methyltetracontane (521–797)	137.1	536		[1999DYK/SVO]
C ₄₁ H ₈₄	[857685-81-5] $\Delta_{\text{fus}} H$	18-hexylpentatricontane	107	314	DSC	[2005IKE/YAM]
C ₄₁ H ₈₄ S	[66576-39-4] $\Delta_v H$	1-hentetracontanethiol (539–822)	140.8	554	E	[1999DYK/SVO]
C ₄₂ H ₂₈	[517-51-1] $\Delta_{\text{sub}} H$	5,6,11,12-tetraphenyltetracene (453–523)	160.6 ± 4.2	488		[1958HOY/PEP, 1970COX/PIL]
C ₄₂ H ₃₆ O ₂₄ S ₆	[102088-39-1] $\Delta_{\text{fus}} H$	4-sulfonato-calix[6]arene	242.2	534.8		[2005YAN/MAN]
C ₄₂ H ₄₄ N ₂ O ₄	[158547-49-0] $\Delta_{\text{us}} H$ (liq <i>cryst</i>) $\Delta_{\text{us}} H$ (liq <i>cryst</i>) $\Delta_{\text{us}} H$ (liq <i>cr-liq</i>)	3,3',4,4'-biphenyltetracarboxy-N,N'-bis-(4- <i>n</i> -heptylphenyl)diimide	18.8 24.7 11.1	411 504.9 560.8		[1995EIS/DEN] [1995EIS/DEN]
C ₄₂ H ₅₂ N ₂ O ₄ S ₂	[109537-98-6] $\Delta_{\text{fus}} H$	1,2-bis-[4-(5-octyl-2-thienylmethylideneamino)-phenylcarboxyloxy]ethane	57.4	369.2		[1978KOS/BUD]
C ₄₂ H ₆₁ NO ₄	[425406-53-7] $\Delta_{\text{fus}} H$	2,7-dihexyloxy-9-(3,5-dihexyloxyphenyl)carbazole	41.02	350.2	DSC	[2002PER/LOP]
C ₄₂ H ₆₆ O ₁₂	[65201-69-6] $\Delta_{\text{us}} H$ $\Delta_{\text{us}} H$ $\Delta_{\text{us}} H$ $\Delta_{\text{fus}} H$	benzene-hexa- <i>n</i> -hexanoate (13–393) (13–393) (13–393) (13–393)	25.67 12.27 16.26 33.5	251.6 291.5 348.3 368.7	AC	[1996DOM/HEA]
C ₄₂ H ₇₈	[61828-33-9] $\Delta_v H$	hexatriacontylbenzene (529–815)	129	544		[1999DYK/SVO]
C ₄₂ H ₈₂ O ₄	[26719-48-2] $\Delta_v H$ $\Delta_v H$	dihexadecyl sebacate	149.8 183.8 ± 6.4	460 298	TGA TGA	[1990KIS/SHO] [1990KIS/SHO]
C ₄₂ H ₈₄	[21807-60-3] $\Delta_v H$	1-dotetracontene (526–806)	137.1	541		[1999DYK/SVO]
C ₄₂ H ₈₄	[61828-21-5] $\Delta_v H$	hexatriacontylcyclohexane (529–816)	135.8	544		[1999DYK/SVO]
C ₄₂ H ₈₆	[7098-20-6] $\Delta_{\text{us}} H$ $\Delta_{\text{us}} H$ $\Delta_{\text{fus}} H$ $\Delta_v H$ $\Delta_v H$	dotetracontane	2.11 8.46 165.97 213.5 136	344.6 348.6 357.3 298 582	DSC CGC A, E	[2006WAN/TOZ] [2008CHI/WAN] [1987STE/MAL, 1966KUD/ZWO]
C ₄₂ H ₈₆	[66576-40-7] $\Delta_v H$	2-methylhentetracontane (526–803)	138.5	541		[1999DYK/SVO]
C ₄₂ H ₈₆	[55470-97-8] $\Delta_v H$	2,2,4,15,17,17-hexamethyl-7,12-bis(3,5,5-trimethylhexyl)octadecane (512–575)	118.3	527	A, MG	[1987STE/MAL, 1955SCH/WHI]
C ₄₂ H ₈₆ S	[66576-41-8]	1-dotetracontanethiol				

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(543–828)	142.1	558	E	[1999DYK/SVO]
C ₄₂ H ₈₇ N	[27911-72-4]	tritradecylamine				
	$\Delta_v H$	(609–848)	86.6	624	A	[1987STE/MAL]
C ₄₃ H ₅₃ N ₃	[500362-46-9]	4-octylphenyl-[6-(4-octylphenyl)methyl-9-methyl-9H-carbazol-3-ylmethylene]amine				
	$\Delta_{\text{fus}} H$		33.1	425.2	DSC	[2002BEL/MAN]
C ₄₃ H ₈₀	[66576-74-7]	heptatriacontylbenzene				
	$\Delta_v H$	(533–821)	138.3	548		[1999DYK/SVO]
C ₄₃ H ₈₆	[66576-75-8]	heptatriacontylcyclohexane				
	$\Delta_v H$	(533–821)	137.4	548		[1999DYK/SVO]
C ₄₃ H ₈₆	[66576-76-9]	1-tritetracontene				
	$\Delta_v H$	(530–812)	138.7	545		[1999DYK/SVO]
C ₄₃ H ₈₈	[66576-76-9]	tritetracontane				
	$\Delta_v H$	(572–820)	137.7	587	A, E	[1987STE/MAL, 1966KUD/ZWO]
C ₄₃ H ₈₈	[66576-77-0]	2-methyldotetracontane				
	$\Delta_v H$	(530–809)	140	545		[1999DYK/SVO]
C ₄₃ H ₈₈ S	[66576-78-1]	1-tritetracontanethiol				
	$\Delta_v H$	(547–833)	143.2	562	E	[1999DYK/SVO]
C ₄₄ H ₂₆ Br ₄ N ₄	[68772-71-4]	5,10,15,20-tetrakis(3-bromophenyl)porphine				
	$\Delta_{\text{sub}} H$		204 ± 4		GS	[2000PER/GOL]
C ₄₄ H ₂₆ Br ₄ N ₄	[29162-73-0]	5,10,15,20-tetrakis(4-bromophenyl)porphine				
	$\Delta_{\text{sub}} H$		135 ± 4		GS	[2000PER/GOL]
C ₄₄ H ₂₆ Cl ₄ N ₄	[22112-77-2]	5,10,15,20-tetrakis(4-chlorophenyl)porphine				
	$\Delta_{\text{sub}} H$		311 ± 5		GS	[2000PER/GOL]
C ₄₄ H ₂₆ F ₄ N ₄	[27185-62-2]	5,10,15,20-tetrakis(2-fluorophenyl)porphine				
	$\Delta_{\text{sub}} H$		225 ± 8		GS	[2000PER/GOL]
C ₄₄ H ₂₆ F ₄ N ₄	[37095-43-5]	5,10,15,20-tetrakis(4-fluorophenyl)porphine				
	$\Delta_{\text{sub}} H$		178 ± 4		GS	[2000PER/GOL]
C ₄₄ H ₃₀ N ₄	[917-23-7]	5,10,15,20-tetraphenylporphine				
	$\Delta_{\text{sub}} H$	(626–707)	142 ± 3		Fluoresc	[2004STE/STI]
	$\Delta_{\text{sub}} H$	(543–555)	171 ± 2	550	ME	[2002TOR/CAM]
	$\Delta_{\text{sub}} H$		240 ± 7		GS	[2000PER/GOL]
	$\Delta_{\text{sub}} H$ (I)	(560–630)	267 ± 9			[1994GOL/PER]
	$\Delta_{\text{sub}} H$ (II)		185 ± 10			[1994GOL/PER]
C ₄₄ H ₄₈ N ₂ O ₄	[na]	3,3',4,4'-biphenyltetracarboxy-N,N'-bis-(4-n-octylphenyl)diimide				
	$\Delta_{\text{us}} H$ (liq cryst)		36.1	428.5		
	$\Delta_{\text{us}} H$ (liq cryst)		21.3	499.2		
	$\Delta_{\text{us}} H$ (liq cr-liq)		8.5	553.5		[1995EIS/DEN]
C ₄₄ H ₆₃ N ₃ O ₂	[na]	2-[3,5-bis[4-(dodecyloxy)phenyl]-1H-pyrazol-1-yl]pyridine				
	$\Delta_{\text{fus}} H$		47.3	344.2		[2003MAY/TOR]
C ₄₄ H ₈₀ O ₄	[175848-72-3]	2,5-di-n-nonadecyloxy-1,4-benzoquinone				
	$\Delta_{\text{us}} H$		16.2	385.5		
	$\Delta_{\text{fus}} H$		134	396.2	DSC	[1996KEE/VAN]
C ₄₄ H ₈₂	[66576-79-2]	octatriacontylbenzene				

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(537–826)	139.7	552		[1999DYK/SVO]
C ₄₄ H ₈₈	[66576-80-5]	octatriacontylcyclohexane				
	$\Delta_v H$	(537–827)	138.7	552		[1999DYK/SVO]
C ₄₄ H ₈₈	[66576-81-6]	1-tetratetracontene				
	$\Delta_v H$	(534–818)	140.1	549		[1999DYK/SVO]
C ₄₄ H ₉₀	[7098-22-8]	tetratetracontane				
	$\Delta_{\text{fus}} H$		145.5	360.9		[1995HAM/MEH]
	$\Delta_v H$	(323–523)	223.7 ± 0.9	298	CGC	[2008CHI/WAN]
	$\Delta_v H$		146	387		[1973IVA/GUJ]
	$\Delta_v H$	(577–821)	139.3	592	A, E	[1987STE/MAL, 1966KUD/ZWO]
C ₄₄ H ₉₀	[66576-82-7]	2-methyltritetracontane				
	$\Delta_v H$	(534–815)	141.5	549		[1999DYK/SVO]
C ₄₄ H ₉₀ S	[66576-83-8]	1-tetratetracontanethiol				
	$\Delta_v H$	(551–838)	144.1	566	E	[1999DYK/SVO]
C ₄₅ H ₈₄	[66576-61-2]	nontriacontylbenzene				
	$\Delta_v H$	(541–832)	141	556		[1999DYK/SVO]
C ₄₅ H ₈₆ O ₆	[60138-25-2]	(dl) 1-lauric-2-myristic-3-palmitic glycerol				
	$\Delta_v H$	(491–551)	147.8	506	A, T	[1987STE/MAL, 1949PER/WEB2]
C ₄₅ H ₈₆ O ₆	[na]	(dl) 1-myristic-2-capric-3-stearic glycerol				
	$\Delta_v H$	(490–551)	148.4	505	A, T	[1987STE/MAL, 1949PER/WEB2]
C ₄₅ H ₈₆ O ₆	[555-45-3]	glycerol trimyristate				
	$\Delta_{\text{fus}} H$		152.2	330.2		[1996DOM/HEA]
	$\Delta_v H$		199.05		TGA	[2002GOO/GEL]
	$\Delta_v H$		155.8	469	TGA	[1990KIS/SHO]
	$\Delta_v H$		199.2 ± 6.9	298	TGA	[1990KIS/SHO]
	$\Delta_v H$	(488–551)	147.8	503	A, T	[1987STE/MAL, 1949PER/WEB2]
C ₄₅ H ₉₀	[66576-62-3]	nonatriacontylcyclohexane				
	$\Delta_v H$	(541–832)	140.1	556		[1999DYK/SVO]
C ₄₅ H ₉₀	[66576-63-4]	1-pentatetracontene				
	$\Delta_v H$	(538–823)	141.5	553		[1999DYK/SVO]
C ₄₅ H ₉₂	[7098-23-9]	pentatetracontane				
	$\Delta_v H$	(582–827)	141	597	A, E	[1987STE/MAL, 1966KUD/ZWO]
C ₄₅ H ₉₂	[66576-64-5]	2-methyltetratetracontane				
	$\Delta_v H$	(538–820)	142.9	553		[1999DYK/SVO]
C ₄₅ H ₉₂ S	[66576-65-6]	1-pentatetracontanethiol				
	$\Delta_v H$	(554–843)	145.6	569	E	[1999DYK/SVO]
C ₄₆ H ₈₆	[66576-67-8]	tetracontylbenzene				
	$\Delta_v H$	(545–837)	142.3	560		[1999DYK/SVO]
C ₄₆ H ₉₀ O ₄	[3072-03-5]	dioctadecyl sebacate				
	$\Delta_v H$		157.5	480	TGA	[1990KIS/SHO]
	$\Delta_v H$		197.7 ± 6.9	298	TGA	[1990KIS/SHO]
C ₄₆ H ₉₂	[66576-68-9]	1-hexatetracontene				
	$\Delta_v H$	(542–828)	142.8	557		[1999DYK/SVO]

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₄₆ H ₉₂	[66576-69-0] $\Delta_v H$	tetracontylcyclohexane (545–837)	141.3	560		[1999DYK/SVO]
C ₄₆ H ₉₄	[7098-24-0] $\Delta_{\text{ms}} H$ $\Delta_{\text{fus}} H$ $\Delta_v H$ $\Delta_v H$	hexatetracontane (323–523) (586–832)	23.9 151.4 233.3 142.8	341.4 360.7 298 601		[2003BRI/BOU] CGC A, E [2008CHI/WAN] [1987STE/MAL, 1966KUD/ZWO]
C ₄₆ H ₉₄	[66564-10-1] $\Delta_v H$	2-methylpentatetracontane (542–826)	144.2	557		[1999DYK/SVO]
C ₄₆ H ₉₄ S	[66564-11-2] $\Delta_v H$	1-hexatetracontanethiol (557–847)	146.7	572	E	[1999DYK/SVO]
C ₄₇ H ₈₈	[66564-12-3] $\Delta_v H$	hentetracontylbenzene (549–842)	143.5	564		[1999DYK/SVO]
C ₄₇ H ₉₀ O ₆	[na] $\Delta_v H$	(dl) 1-myristic-2-lauric-3-stearic glycerol (493–558)	150.5	508	A, T	[1987STE/MAL, 1949PER/WEB2]
C ₄₇ H ₉₀ O ₆	[na] $\Delta_v H$	(dl) 1-palmitic-2-capric-3-stearic glycerol (507–559)	154.8	522	A, T	[1987STE/MAL, 1949PER/WEB2]
C ₄₇ H ₉₄	[66564-13-4] $\Delta_v H$	hentetracontylcyclohexane (548–842)	142.8	563		[1999DYK/SVO]
C ₄₇ H ₉₄	[66576-01-0] $\Delta_v H$	1-heptatetracontene (546–833)	143.9	561		[1999DYK/SVO]
C ₄₇ H ₉₆	[7098-25-1] $\Delta_v H$	heptatetracontane (591–837)	144.2	606	A, E	[1987STE/MAL, 1966KUD/ZWO]
C ₄₇ H ₉₆	[66576-02-1] $\Delta_v H$	2-methylhexatetracontane (546–831)	145.3	561		[1999DYK/SVO]
C ₄₇ H ₉₆ S	[66576-03-2] $\Delta_v H$	1-heptatetracontanethiol (561–852)	147.6	576	E	[1999DYK/SVO]
C ₄₈ H ₃₈ N ₄	[37083-40-2] $\Delta_{\text{sub}} H$	5,10,15,20-tetrakis(2-methylphenyl)porphine 159 ± 5			GS	[2000PER/GOL]
C ₄₈ H ₃₈ N ₄	[50849-45-1] $\Delta_{\text{sub}} H$	5,10,15,20-tetrakis(3-methylphenyl)porphine 177 ± 5			GS	[2000PER/GOL]
C ₄₈ H ₃₈ N ₄	[14527-51-6] $\Delta_{\text{sub}} H$	5,10,15,20-tetrakis(4-methylphenyl)porphine 178 ± 3			GS	[2000PER/GOL]
C ₄₈ H ₃₈ N ₄ O ₄	[22112-78-3] $\Delta_{\text{sub}} H$	5,10,15,20-tetrakis(4-methoxyphenyl)porphine (561–565)	213 ± 12	563	ME	[2002TOR/CAM]
C ₄₈ H ₄₀ P ₂	[na] $\Delta_{\text{fus}} H$	2,2'-bis(di-4-toluenephosphino)-1,1'-binaphthyl 41.98		528.3		[1997ZHA/GAO]
C ₄₈ H ₉₀	[66576-04-3] $\Delta_v H$	dotetracontylbenzene (552–846)	144.9	567		[1999DYK/SVO]
C ₄₈ H ₉₆	[66576-05-4] $\Delta_v H$	dotetracontylcyclohexane (552–847)	143.9	567		[1999DYK/SVO]
C ₄₈ H ₉₆	[66576-06-5] $\Delta_v H$	1-octatetracontene (549–838)	145.4	564		[1999DYK/SVO]

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₄₈ H ₉₈	[7098-26-2]	octatetracontane				
	$\Delta_v H$	(323–523)	243.0 ± 0.2	298	CGC	[2008CHI/WAN]
	$\Delta_v H$	(595–843)	145.9	610	A, E	[1987STE/MAL, 1966KUD/ZWO]
C ₄₈ H ₉₈	[66576-07-6]	2-methylheptatetracontane				
	$\Delta_v H$	(550–836)	146.5	565		[1999DYK/SVO]
C ₄₈ H ₉₈ S	[66576-08-7]	1-octatetracontanethiol				
	$\Delta_v H$	(564–856)	148.7	579	E	[1999DYK/SVO]
C ₄₉ H ₉₂	[66576-09-8]	tritetracontylbenzene				
	$\Delta_v H$	(556–851)	145.9	571		[1999DYK/SVO]
C ₄₉ H ₉₄ O ₆	[na]	(<i>dl</i>) 1-palmitic-2-lauryl-3-stearic glycerol				
	$\Delta_v H$	(506–567)	160	521	A, T	[1987STE/MAL, 1949PER/WEB2]
C ₄₉ H ₉₈	[66576-19-1]	1-nonatetracontene				
	$\Delta_v H$	(553–843)	146.4	568		[1999DYK/SVO]
C ₄₉ H ₉₈	[66576-11-2]	tritetracontylcyclohexane				
	$\Delta_v H$	(556–852)	144.9	571		[1999DYK/SVO]
C ₄₉ H ₁₀₀	[7098-27-3]	nonatetracontane				
	$\Delta_v H$	(599–847)	147.5	614	A, E	[1987STE/MAL, 1966KUD/ZWO]
C ₄₉ H ₁₀₀	[66576-12-3]	2-methyloctatetracontane				
	$\Delta_v H$	(553–840)	147.9	568		[1999DYK/SVO]
C ₄₉ H ₁₀₀ S	[66576-13-4]	1-nonatetracontanethiol				
	$\Delta_v H$	(567–861)	149.7	582	E	[1999DYK/SVO]
C ₅₀ H ₉₄	[66576-14-5]	tetratetracontylbenzene				
	$\Delta_v H$	(559–856)	147.1	574		[1999DYK/SVO]
C ₅₀ H ₁₀₀	[63911-02-4]	1-pentacontene				
	$\Delta_v H$	(556–848)	147.8	571		[1999DYK/SVO]
C ₅₀ H ₁₀₀	[66576-15-6]	tetratetracontylcyclohexane				
	$\Delta_v H$	(559–856)	146.2	574		[1999DYK/SVO]
C ₅₀ H ₁₀₂	[6596-40-3]	pentacontane				
	$\Delta_{\text{fus}} H$		185	366.9		[1995HAM/MEH]
	$\Delta_v H$	(323–523)	252.5 ± 0.2	298	CGC	[2008CHI/WAN]
	$\Delta_v H$	(603–852)	149	618	A, E	[1987STE/MAL, 1966KUD/ZWO]
C ₅₀ H ₁₀₂	[66576-16-7]	2-methylnonatetracontane				
	$\Delta_v H$	(557–845)	148.8	572		[1999DYK/SVO]
C ₅₀ H ₁₀₂ S	[66576-17-8]	1-pentacontanethiol				
	$\Delta_v H$	(570–865)	150.7	585	E	[1999DYK/SVO]
C ₅₁ H ₆₉ N ₃	[500362-48-1]	4-dodecylphenyl-[6-(4-dodecylphenyl)methyl-9-methyl-9H-carbazol-3-yl-methylene]amine				
	$\Delta_{\text{fus}} H$		34.5	412.2	DSC	[2002BEL/MAN]
C ₅₁ H ₉₆	[66576-18-9]	pentatetracontylbenzene				
	$\Delta_v H$	(562–860)	148.3	577		[1999DYK/SVO]
C ₅₁ H ₉₈ O ₆	[60138-20-7]	1-myristic-2-palmitic-3-stearic glycerol				
	$\Delta_v H$	(508–572)	157.9	523	A, T	[1987STE/MAL, 1949PER/WEB2]
C ₅₁ H ₉₈ O ₆	[555-44-2]	glycerol tripalmitate (tripalmitin)				
	$\Delta_{\text{fus}} H$		162.6	340.5	DSC	[2010HON/HUA]

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{fus}}H$		121	337.4	DSC	[2006LI/ROD]
	$\Delta_{\text{fus}}H$ (β)		177.2	339	DSC	[1999VAN/TEN]
	$\Delta_{\text{fus}}H$		179.37	338.9		[1996DOM/HEA]
	Δ_vH		U 474.3		TGA	[2002GOO/GEL]
	Δ_vH		166.3	483	TGA	[1990KIS/SHO]
	Δ_vH		217.1 \pm 7.6	298	TGA	[1990KIS/SHO]
	Δ_vH	(506–572)	160.8	521	A, T	[1987STE/MAL, 1949PER/WEB2]
C ₅₁ H ₁₀₀ CIN ₅	[106486-51-5]	2,4-bis N,N-didodecylamino-6-chloro-1,3,5-triazine				
	$\Delta_{\text{fus}}H$		34.25	307.5		[1986LAT/HOE]
C ₅₁ H ₁₀₂	[na]	1-henpentacontene				
	Δ_vH	(560–852)	148.6	575		[1999DYK/SVO]
C ₅₁ H ₁₀₂	[66576-20-3]	pentatetracontylcyclohexane				
	Δ_vH	(562–861)	147.4	577		[1999DYK/SVO]
C ₅₁ H ₁₀₂ N ₆	[38565-86-5]	tris N,N-dioctylamino-1,3,5-triazine				
	$\Delta_{\text{fus}}H$		74.25	312.7		[1986LAT/HOE]
C ₅₁ H ₁₀₄	[7667-76-7]	henpentacontane				
	$\Delta_{\text{us}}H$		1.75	337		
	$\Delta_{\text{us}}H$		5.3	343		
	$\Delta_{\text{fus}}H$		170.4	365.6		[1992SRC/KER]
	Δ_vH	(607–857)	150.6	622	A, E	[1987STE/MAL, 1966KUD/ZWO]
C ₅₁ H ₁₀₄	[66575-81-3]	2-methylpentacontane				
	Δ_vH	(560–850)	150	575		[1999DYK/SVO]
C ₅₁ H ₁₀₄ S	[66575-82-4]	1-henpentacontanethiol				
	Δ_vH	(573–869)	151.6	588	E	[1999DYK/SVO]
C ₅₂ H ₃₄ O ₂	[3432-73-3]	1,4-bis(2,4,5-triphenylcyclopentadienone-3-yl)benzene				
	$\Delta_{\text{fus}}H$	(300–650)	68.8	594.4		[2005SMI/KUL]
C ₅₂ H ₉₈	[66575-84-6]	hexatetracontylbenzene				
	Δ_vH	(566–864)	149.1	581		[1999DYK/SVO]
C ₅₂ H ₁₀₄	[66575-85-7]	1-dopentacontene				
	Δ_vH	(563–857)	149.8	578		[1999DYK/SVO]
C ₅₂ H ₁₀₄	[66575-86-8]	hexatetracontylcyclohexane				
	Δ_vH	(565–865)	148.5	580		[1999DYK/SVO]
C ₅₂ H ₁₀₆	[7719-79-1]	dopentacontane				
	$\Delta_{\text{us}}H$		17.1	352		
	$\Delta_{\text{fus}}H$		171.8	366.7		[1992SRC/KER]
	Δ_vH	(323–523)	261.8 \pm 1.5	298	CGC	[2008CHI/WAN]
	Δ_vH	(611–861)	152	626	A, E	[1987STE/MAL, 1966KUD/ZWO]
C ₅₂ H ₁₀₆	[66575-87-9]	2-methylhenpentacontane				
	Δ_vH	(563–854)	151.2	578		[1999DYK/SVO]
C ₅₂ H ₁₀₆ O ₂₆	[na]	1, ω -dimethoxypentacos(oxyethylene)				
	$\Delta_{\text{fus}}H$		209.7	316.2	DSC	[1996YAN/YU]
C ₅₂ H ₁₀₆ S	[66575-88-0]	1-dopentacontanethiol				
	Δ_vH	(575–873)	152.6	590	E	[1999DYK/SVO]

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
C ₅₃ H ₉₈ O ₆	[na] $\Delta_{\text{fus}}H$	1,3-bis-hexadecanoyloxy-2-(9-cis-octadecenoyloxy)propane				
			149.7	311.4		[1984PER, 1978TIM]
C ₅₃ H ₁₀₀	[66575-89-1] $\Delta_{\text{v}}H$	heptatetracontylbenzene				
		(569–868)	150.1	584		[1999DYK/SVO]
C ₅₃ H ₁₀₆	[66563-49-3] $\Delta_{\text{v}}H$	heptatetracontylcyclohexane				
		(568–869)	149.6	583		[1999DYK/SVO]
C ₅₃ H ₁₀₆	[66577-50-2] $\Delta_{\text{v}}H$	1-tripentacontene				
		(566–861)	150.9	581		[1999DYK/SVO]
C ₅₃ H ₁₀₈	[7719-80-4] $\Delta_{\text{v}}H$	tripentacontane				
		(615–866)	153.4	630	A, E	[1987STE/MAL, 1966KUD/ZWO]
C ₅₃ H ₁₀₈	[66575-90-4] $\Delta_{\text{v}}H$	2-methyldopentacontane				
		(566–858)	152.3	581		[1999DYK/SVO]
C ₅₃ H ₁₀₈ S	[66575-91-5] $\Delta_{\text{v}}H$	1-tripentacontanethiol				
		(578–877)	153.7	593	E	[1999DYK/SVO]
C ₅₄ H ₁₀₂	[66575-92-6] $\Delta_{\text{v}}H$	octatetracontylbenzene				
		(572–873)	151.1	587		[1999DYK/SVO]
C ₅₄ H ₁₀₈	[66575-93-7] $\Delta_{\text{v}}H$	octatetracontylcyclohexane				
		(571–873)	150.6	586		[1999DYK/SVO]
C ₅₄ H ₁₀₈	[66575-94-8] $\Delta_{\text{v}}H$	1-tetrapentacontene				
		(569–865)	151.9	584		[1999DYK/SVO]
C ₅₄ H ₁₀₈ O ₂₇	[182292-69-9] $\Delta_{\text{fus}}H$	81-crown-27				
			155.6	314.2	DSC	[1996YAN/YU]
C ₅₄ H ₁₁₀	[5856-66-6] $\Delta_{\text{ms}}H$ $\Delta_{\text{fus}}H$ $\Delta_{\text{v}}H$ $\Delta_{\text{v}}H$	tetrapentacontane				
			39	344.9		
			177.2	368		[2003BRI/BOU]
		(323–523)	271.0 ± 1.7	298	CGC	[2008CHI/WAN]
		(618–870)	155	633	A, E	[1987STE/MAL, 1966KUD/ZWO]
C ₅₄ H ₁₁₀	[66575-95-9] $\Delta_{\text{v}}H$	2-methyltripentacontane				
		(569–863)	153.4	584		[1999DYK/SVO]
C ₅₄ H ₁₁₀ S	[66575-96-0] $\Delta_{\text{v}}H$	1-tetrapentacontanethiol				
		(581–881)	154.4	596	E	[1999DYK/SVO]
C ₅₅ H ₁₀₂ O ₆	[na] $\Delta_{\text{fus}}H$	2-hexadecanoyloxy-1,3-bis-(9-cis-octadecenoyloxy)propane				
			125.5	291.9		[1984PER, 1978TIM]
C ₅₅ H ₁₀₄	[66575-98-2] $\Delta_{\text{v}}H$	nontetracontylbenzene				
		(575–877)	152.1	590		[1999DYK/SVO]
C ₅₅ H ₁₁₀	[66575-99-3] $\Delta_{\text{v}}H$	nonatetracontylcyclohexane				
		(574–877)	151.6	589		[1999DYK/SVO]
C ₅₅ H ₁₁₀	[66576-00-9] $\Delta_{\text{v}}H$	1-pentapentacontene				
		(572–869)	152.9	587		[1999DYK/SVO]
C ₅₅ H ₁₁₂	[5846-40-2] $\Delta_{\text{v}}H$	pentapentacontane				
		(622–874)	156.3	637	A, E	[1987STE/MAL, 1966KUD/ZWO]
C ₅₅ H ₁₁₂	[66575-60-8] $\Delta_{\text{v}}H$	2-methyltetrapentacontane				
		(572–867)	154.3	587		[1999DYK/SVO]

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₅₅ H ₁₁₂ S	[66575-61-9] $\Delta_v H$	1-pentapentacontanethiol (584–885)	155	599	E	[1999DYK/SVO]
C ₅₆ H ₄₈ O ₃₂ S ₈	[137407-62-6] $\Delta_{\text{fus}} H$	4-sulfonato-calix[8]arene	350.6	543.1		[2005YAN/MAN]
C ₅₆ H ₇₈ O ₈	[122356-76-7] $\Delta_{\text{sub}} H$	2,23,28,38-tetrakis(1,1-dimethylethyl)-6,7,9,10,12,13,15,16,18,19-decahydro-32,35-dimethoxy-31H-4,21-(methano[1,3]benzenomethano)-26,30-metheno-25H-dibenzo[q,z][1,4,7,10,13,16]hexaoxacycloheptacosin	78 ± 1			[2008SUR]
C ₅₆ H ₁₀₆	[66575-62-0] $\Delta_v H$	pentacontylbenzene (577–880)	153.2	592		[1999DYK/SVO]
C ₅₆ H ₁₀₈	[66575-63-1] $\Delta_v H$	1-hexapentacontene (575–873)	154.5	588		[1999DYK/SVO]
C ₅₆ H ₁₀₈	[66575-64-2] $\Delta_v H$	pentacontylcyclohexane (577–881)	152.4	592		[1999DYK/SVO]
C ₅₆ H ₁₁₄	[7719-82-6] $\Delta_v H$ $\Delta_v H$	hexapentacontane (323–523) (625–878)	279.7 ± 1.5 157.8	298 640	CGC A, E	[2008CHI/WAN] [1987STE/MAL, 1966KUD/ZWO]
C ₅₆ H ₁₁₄	[66575-65-3] $\Delta_v H$	2-methylpentapentacontane (575–871)	155.9	588		[1999DYK/SVO]
C ₅₆ H ₁₁₄ O ₂₈	[na] $\Delta_{\text{fus}} H$	1, ω -dimethoxyheptacos(oxyethylene)	224.6	315.2	DSC	[1996YAN/YU]
C ₅₆ H ₁₁₄ S	[66575-66-4] $\Delta_v H$	1-hexapentacontanethiol (586–888)	156	601	E	[1999DYK/SVO]
C ₅₇ H ₅₄ N ₆ O ₆	[130048-22-5] $\Delta_{\text{fus}} H$	1,3,5- <i>tris</i> -(4- <i>n</i> -butyl-4'-diazophenyl)benzenetricarboxylate	11.3	443.2	DSC	[1990JIN/KAN]
C ₅₇ H ₁₀₄ O ₆	[537-39-3] $\Delta_{\text{fus}} H$ (I) $\Delta_{\text{fus}} H$ (II)	1,2,3-tri(<i>trans</i> -9-octadecenoyl)glycerol (trilaidin)	157.07 84.2	314.8 288	DSC DSC	[2003VAN/VAN2]
C ₅₇ H ₁₀₈	[66575-67-5] $\Delta_v H$	heptapentacontylbenzene (580–884)	154.1	595		[1999DYK/SVO]
C ₅₇ H ₁₀₈ O ₆	[2846-04-0] $\Delta_v H$	1,3-distearic-2-oleic glycerol (523–593)	165.8	538	A, T	[1987STE/MAL, 1949PER/WEB2]
C ₅₇ H ₁₀₈ O ₆	[555-43-1] $\Delta_{\text{fus}} H$ $\Delta_v H$ $\Delta_v H$	glycerol tristearate (521–588)	203.26 220.8 167.5	345.7 536	TGA A, T	[1996DOM/HEA] [2002GOO/GEL] [1987STE/MAL, 1949PER/WEB2]
C ₅₇ H ₁₁₀ O ₆	[555-43-1] $\Delta_{\text{fus}} H$ (α) $\Delta_{\text{fus}} H$ (β') $\Delta_{\text{fus}} H$ (β) $\Delta_{\text{fus}} H$ (I) $\Delta_{\text{fus}} H$ (I) $\Delta_{\text{fus}} H$ (II) $\Delta_{\text{fus}} H$ (II) $\Delta_v H$	tristearin (10–370) (10–370)	115.4 144.8 293.5 197.6 195.8 129.1 114.1 174.9	328.1 338.5 345.6 345.9 346 327.3 327.3 506	AC DSC AC DSC DSC TGA	[2009DAS/BRE] [2005MAT/VAN] [1990KIS/SHO]

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$		236.2 ± 8.3	298	TGA	[1990KIS/SHO]
C ₅₇ H ₁₁₄	[66575-68-6] $\Delta_v H$	henpentacontylcyclohexane (580–885)	153.3	595		[1999DYK/SVO]
C ₅₇ H ₁₁₄	[66575-69-7] $\Delta_v H$	1-heptapentacontene (578–877)	154.6	593		[1999DYK/SVO]
C ₅₇ H ₁₁₆	[5856-67-7] $\Delta_v H$	heptapentacontane (629–882)	158.9	644	A, E	[1987STE/MAL, 1966KUD/ZWO]
C ₅₇ H ₁₁₆	[66575-70-0] $\Delta_v H$	2-methylhexapentacontane (578–875)	155.9	593		[1999DYK/SVO]
C ₅₇ H ₁₁₆ S	[66575-75-1] $\Delta_v H$	1-heptapentacontanethiol (589–892)	156.7	604	E	[1999DYK/SVO]
C ₅₈ H ₁₁₀	[66575-73-3] $\Delta_v H$	dopentacontylbenzene (583–888)	155.7	598		[1999DYK/SVO]
C ₅₈ H ₁₁₆	[66575-74-4] $\Delta_v H$	dopentacontylcyclohexane (582–888)	154.3	597		[1999DYK/SVO]
C ₅₈ H ₁₁₆	[66575-75-5] $\Delta_v H$	1-octapentacontene (580–881)	155.8	595		[1999DYK/SVO]
C ₅₈ H ₁₁₈	[7667-78-9] $\Delta_v H$ $\Delta_v H$	octapentacontane (323–523) (632–886)	288.5 ± 1.8 160.3	298 647	CGC A, E	[2008CHI/WAN] [1987STE/MAL, 1966KUD/ZWO]
C ₅₈ H ₁₁₈	[66575-76-6] $\Delta_v H$	2-methylheptapentacontane (581–879)	156.8	596		[1999DYK/SVO]
C ₅₈ H ₁₁₈ S	[66575-77-7] $\Delta_v H$	1-octapentacontanethiol (591–895)	157.4	606	E	[1999DYK/SVO]
C ₅₉ H ₈₅ N ₃	[500362-49-2] $\Delta_{\text{fus}} H$	4-hexadecylphenyl-[6-(4-hexadecylphenyl)methyl-9-methyl-9H-carbazol-3-yl-methylene]amine	52.1	405.2	DSC	[2002BEL/MAN]
C ₅₉ H ₉₀ O ₄	[303-98-0] $\Delta_{\text{fus}} H$	2-[(2E,6E,10E,14E,18E,22E,26E,30E,34E)-3,7,11,15,19,23,27,31,-35,39-decamethyl-2,6,10,14,18,22,26,30,34,38-tetracontadecaen-1-yl]-5,6-dimethoxy-3-methyl-2,5-cyclohexadiene-1,4-dione (Ubidecarenone)	82	322.5	DSC	[2006LI/ROD]
C ₅₉ H ₁₁₂	[66575-78-8] $\Delta_v H$	tripentacontylbenzene (585–891)	155.9	600		[1999DYK/SVO]
C ₅₉ H ₁₁₈	[66575-80-2] $\Delta_v H$	tripentacontylcyclohexane (585–892)	155	600		[1999DYK/SVO]
C ₅₉ H ₁₁₈	[66575-79-9] $\Delta_v H$	1-nonapentacontene (583–885)	156.4	598		[1999DYK/SVO]
C ₅₉ H ₁₂₀	[7667-70-0] $\Delta_v H$	nonapentacontane (635–890)	161.8	650	A, E	[1987STE/MAL, 1966KUD/ZWO]
C ₅₉ H ₁₂₀	[66575-49-3] $\Delta_v H$	2-methyloctapentacontane (583–882)	157.9	598		[1999DYK/SVO]
C ₅₉ H ₁₂₀ S	[66575-50-6] $\Delta_v H$	1-nonapentacontanethiol (593–899)	158.3	608	E	[1999DYK/SVO]
C ₆₀	[99685-96-8] $\Delta_{\text{us}} H$	buckminsterfullerene	9.0	261.4	DSC	[1993DEB/DWO]

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
	$\Delta_{\text{sub}}H$	(775–1033)	180 ± 2	298	ME	[2000SCH/MAT]
	$\Delta_{\text{sub}}H$	(789–907)	152.8 ± 0.1	897	GS	[1998PAN/MAL]
	$\Delta_{\text{sub}}H$		183.5 ± 1.0	298		[1998PAN/MAL]
	$\Delta_{\text{sub}}H$		179.2 ± 3.5	298		[1996GON/SUN, 1998PAN/MAL]
	$\Delta_{\text{sub}}H$	(730–990)	175.2 ± 2.9	860	ME,TE	[1995RIA/GIG]
	$\Delta_{\text{sub}}H$		181 ± 2.0	298	ME,TE	[1995RIA/GIG]
	$\Delta_{\text{sub}}H$		219.6		TGA	[1995YAS/TAK]
	$\Delta_{\text{sub}}H$	(546–722)	180 ± 10.0	634	UV	[94DAI/MAC]
	$\Delta_{\text{sub}}H$		158 ± 3.0	700	ME	[1994POP/DRA]
	$\Delta_{\text{sub}}H$		168.5 ± 1.2	298	ME	[1994POP/DRA, 1998PAN/MAL]
	$\Delta_{\text{sub}}H$		181.1 ± 2.6	298	ME	[1994KOR/SID, 1998PAN/MAL]
	$\Delta_{\text{sub}}H$		181.4 ± 2.3	700	MS	[1994SAI/LAK, 1992MAT/SAI]
	$\Delta_{\text{sub}}H$		158.6	773	ME	[1993MAT/SAI]
	$\Delta_{\text{sub}}H$		184.1 ± 3.1	298	GS	[1992PAN/CHA, 1998PAN/MAL]
	$\Delta_{\text{sub}}H$		183.2 ± 3.5	298	ME	[1992MAT/SAI, 1998PAN/MAL]
	$\Delta_{\text{sub}}H$		180.6 ± 1.5	298	ME	[1992ABR/OLA, 1998PAN/MAL]
	$\Delta_{\text{sub}}H$	(673–873)	159.0 ± 4.2		ME	[1992ABR/OLA2]
	$\Delta_{\text{sub}}H$		>163 (powder)		TGA	[1992CHE/KOR]
	$\Delta_{\text{sub}}H$	(640–800)	167.8 ± 5.4	707	ME,MS	[1991PAN/SAM]
	$\Delta_{\text{sub}}H$		U 138.5	600		[1991TOK/HAY]
	$\Delta_{\text{sub}}H$		U 90.0		ME,MS	[1990HAU/CON]
C₆₀F₁₆	[na]	hexadecafluorobuckminsterfullerene				
	$\Delta_{\text{sub}}H$		186 ± 9		ME,MS	[2000MAR/BOL]
C₆₀F₁₈	[172760-25-7]	octadecafluorobuckminsterfullerene				
	$\Delta_{\text{sub}}H$	(591–671)	197 ± 10	627	ME	[2002MAR/BOL]
C₆₀F₃₆	[na]	hexatriacontylfluorobuckminsterfullerene (4 isomer average)				
	$\Delta_{\text{sub}}H$	(422–525)	134 ± 6	473	MS	[1996BOL/MER]
C₆₀F₃₆	[150180-35-1]	hexatriacontylfluorobuckminsterfullerene				
	$\Delta_{\text{sub}}H$		139 ± 8			[2000PAP/KOL]
	$\Delta_{\text{sub}}H$	(408–539)	135 ± 8.0	466	ME,MS	[1999BOL/MAR]
C₆₀F₄₂	[na]	dotetracontylfluorobuckminsterfullerene				
	$\Delta_{\text{sub}}H$	(430–510)	110 ± 10		ME,MS	[2000EME/NIK]
C₆₀F₄₄	[na]	tetratetracontylfluorobuckminsterfullerene				
	$\Delta_{\text{sub}}H$	(430–510)	112 ± 6		ME,MS	[2000EME/NIK]
C₆₀F₄₄O	[na]	tetratetracontylfluorotetratetracontahydro-[5,6]fullereno-C60-1h-oxirene				
	$\Delta_{\text{sub}}H$	(430–510)	111 ± 3		ME,MS	[2000EME/NIK]
C₆₀F₄₆	[na]	hexatetracontylfluorobuckminsterfullerene				
	$\Delta_{\text{sub}}H$	(430–510)	114 ± 7		ME,MS	[2000EME/NIK]
C₆₀F₄₈	[143471-98-1]	octatetracontylfluorobuckminsterfullerene				
	$\Delta_{\text{sub}}H$	(395–528)	109 ± 7.0	476	ME,MS	[1999BOL/MAR, 2000BOL/GAL]
C₆₀F₄₈	[na]	fluorinated fullerene				
	$\Delta_{\text{fus}}H$		7	21.24		
	$\Delta_{\text{fus}}H$		NA			[1999DRU/GAL]
C₆₀H₁₆	[na]	hexadecahydrobuckminsterfullerene				
	$\Delta_{\text{sub}}H$		≥186		E	[2000KOR/DOR, 2001DOR/LOB]
C₆₀H₃₆	[na]	hexatriacontylhydrobuckminsterfullerene				
	$\Delta_{\text{sub}}H$	(560–680)	162 ± 5		MS	[2000KOR/DOR, 2001DOR/LOB]
	$\Delta_{\text{sub}}H$		152	630		[2001DOR/LOB]

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{sub}}H$		175	298		[2001DOR/LOB]
C ₆₀ H ₁₁₄	[66575-51-7] Δ_vH	tetrapentacontylbenzene (588–895)	156.6	603		[1999DYK/SVO]
C ₆₀ H ₁₂₀	[66575-52-8] Δ_vH	1-hexacontene (586–888)	157.1	601		[1999DYK/SVO]
C ₆₀ H ₁₂₀	[66575-53-9] Δ_vH	tetrapentacontylcyclohexane (587–895)	156	602		[1999DYK/SVO]
C ₆₀ H ₁₂₂	[7667-80-3] $\Delta_{\text{fus}}H$	hexacontane	186.8	373.2	DSC	[1992LOU/ROU]
	Δ_vH	(323–523)	299.9 ± 2.0	298	CGC	[2008CHI/WAN]
	Δ_vH	(638–893)	163	653	A, E	[1987STE/MAL, 1966KUD/ZWO]
C ₆₀ H ₁₂₂	[66575-54-0] Δ_vH	2-methylnonapentacontane (586–886)	158.5	601		[1999DYK/SVO]
C ₆₀ H ₁₂₂ S	[66575-55-1] Δ_vH	1-hexacontanethiol (595–902)	159.1	610	E	[1999DYK/SVO]
C ₆₁ H ₁₁₆	[66563-50-6] Δ_vH	pentapentacontylbenzene (590–898)	157.5	605		[1999DYK/SVO]
C ₆₁ H ₁₂₂	[66563-51-7] Δ_vH	1-henhexacontene (588–891)	158	603		[1999DYK/SVO]
C ₆₁ H ₁₂₂	[66563-52-8] Δ_vH	pentapentacontylcyclohexane (590–899)	156.6	605		[1999DYK/SVO]
C ₆₁ H ₁₂₄	[7667-81-4] Δ_vH	henhexacontane (642–897)	163.9	657	A, E	[1987STE/MAL, 1966KUD/ZWO]
C ₆₁ H ₁₂₄	[66563-53-9] Δ_vH	2-methylhexacontane (588–889)	159.4	603		[1999DYK/SVO]
C ₆₁ H ₁₂₄ S	[66563-54-0] Δ_vH	1-henhexacontanethiol (597–905)	159.6	612	E	[1999DYK/SVO]
C ₆₂ H ₁₁₈	[66563-55-1] Δ_vH	hexapentacontylbenzene (592–902)	158.4	607		[1999DYK/SVO]
C ₆₂ H ₁₂₄	[66563-56-2] Δ_vH	1-dohexacontene (590–895)	158.6	605		[1999DYK/SVO]
C ₆₂ H ₁₂₄	[66563-57-3] Δ_vH	hexapentacontylcyclohexane (592–902)	157.5	607		[1999DYK/SVO]
C ₆₂ H ₁₂₆	[7719-83-7] Δ_vH	dohexacontane (323–523)	306.8 ± 0.1	298	CGC	[2008CHI/WAN]
	Δ_vH	(645–901)	165.2	660	A, E	[1987STE/MAL, 1966KUD/ZWO]
C ₆₂ H ₁₂₆	[66563-58-4] Δ_vH	2-methylhenhexacontane (590–892)	160.1	605		[1999DYK/SVO]
C ₆₂ H ₁₂₂ S	[66563-59-5] Δ_vH	1-dohexacontanethiol (599–908)	160.1	614	E	[1999DYK/SVO]
C ₆₃ H ₁₂₀	[66563-60-8] Δ_vH	heptapentacontylbenzene (595–905)	158.9	610		[1999DYK/SVO]

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
C ₆₃ H ₁₂₆	[66563-61-9] $\Delta_{\text{v}}H$	heptapentacontylcyclohexane (594–905)	158.2	609		[1999DYK/SVO]
C ₆₃ H ₁₂₆	[66563-62-0] $\Delta_{\text{v}}H$	1-trihexacontene (593–899)	159.8	608		[1999DYK/SVO]
C ₆₃ H ₁₂₆ N ₆	[106486-49-1] $\Delta_{\text{fus}}H$	tris N,N-didecylamino-1,3,5-triazine	87.68	314.4		[1986LAT/HOE]
C ₆₃ H ₁₂₈	[7719-84-8] $\Delta_{\text{v}}H$	trihexacontane (647–904)	116.7	662	A, E	[1987STE/MAL, 1966KUD/ZWO]
C ₆₃ H ₁₂₈	[66563-63-1] $\Delta_{\text{v}}H$	2-methylidohexacontane (593–897)	161.3	608		[1999DYK/SVO]
C ₆₃ H ₁₂₈ S	[66563-64-2] $\Delta_{\text{v}}H$	1-trihexacontanethiol (602–911)	161.1	617	E	[1999DYK/SVO]
C ₆₄ H ₁₂₂	[66563-65-3] $\Delta_{\text{v}}H$	octapentacontylbenzene (597–908)	159.5	612		[1999DYK/SVO]
C ₆₄ H ₁₂₈	[66563-66-4] $\Delta_{\text{v}}H$	octapentacontylcyclohexane (596–908)	158.8	611		[1999DYK/SVO]
C ₆₄ H ₁₂₈	[66563-36-8] $\Delta_{\text{v}}H$	1-tetrahexacontene (595–902)	160.5	610		[1999DYK/SVO]
C ₆₄ H ₁₃₀	[7719-87-1] $\Delta_{\text{v}}H$ $\Delta_{\text{v}}H$	tetrahexacontane (323–523) (650–907)	315.4 ± 0.4 168.3	298 665	CGC A, E	[2008CHI/WAN] [1987STE/MAL, 1966KUD/ZWO]
C ₆₄ H ₁₃₀	[66563-37-9] $\Delta_{\text{v}}H$	2-methyltrihexacontane (595–900)	161.9	610		[1999DYK/SVO]
C ₆₄ H ₁₃₀ S	[66563-38-0] $\Delta_{\text{v}}H$	1-tetrahexacontanethiol (604–914)	161.6	619	E	[1999DYK/SVO]
C ₆₅ H ₁₂₄	[66563-39-1] $\Delta_{\text{v}}H$	nonapentacontylbenzene (599–911)	160.1	614		[1999DYK/SVO]
C ₆₅ H ₁₃₀	[66563-40-4] $\Delta_{\text{v}}H$	nonapentacontylcyclohexane (599–912)	159.9	614		[1999DYK/SVO]
C ₆₅ H ₁₃₀	[66563-41-5] $\Delta_{\text{v}}H$	1-pentaheptacontene (597–905)	161.1	612		[1999DYK/SVO]
C ₆₅ H ₁₃₂	[7719-88-2] $\Delta_{\text{v}}H$	pentaheptacontane (653–910)	169	668	A, E	[1987STE/MAL, 1966KUD/ZWO]
C ₆₅ H ₁₃₂	[66563-42-6] $\Delta_{\text{v}}H$	2-methyltetraheptacontane (597–903)	162.5	612		[1999DYK/SVO]
C ₆₅ H ₁₃₂ S	[66563-43-7] $\Delta_{\text{v}}H$	1-pentaheptacontanethiol (606–917)	162.1	621	E	[1999DYK/SVO]
C ₆₆ H ₈₀ O ₁₀	[na] $\Delta_{\text{fus}}H$	1,3-bis[4-(4-tetradecyloxybenxoyloxy)benxoyloxy]naphthalene	73.8	401.1	DSC	[2009GIM/CLE]
C ₆₆ H ₈₀ O ₁₀	[na] $\Delta_{\text{us}}H$ $\Delta_{\text{fus}}H$	1,7-bis[4-(4-tetradecyloxybenxoyloxy)benxoyloxy]naphthalene	15.4 54	418.2 428.4	DSC	[2009GIM/CLE]
C ₆₆ H ₁₀₉ NO ₄	[425406-55-9]	2,7-didodecyloxy-9-(3,5-didodecyloxyphenyl)carbazole				

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{fus}}H$		61.87	333.2	DSC	[2002PER/LOP]
C ₆₆ H ₁₂₆	[66563-44-8] Δ_vH	hexacontylbenzene (602–914)	161.2	617		[1999DYK/SVO]
C ₆₆ H ₁₃₂	[66563-45-9] Δ_vH	hexacontylcyclohexane (601–915)	160.5	616		[1999DYK/SVO]
C ₆₆ H ₁₃₂	[66563-46-0] Δ_vH	1-hexahexacontene (599–908)	161.7	614		[1999DYK/SVO]
C ₆₆ H ₁₃₄	[7719-89-3] Δ_vH Δ_vH	hexaheptacontane (323–523) (656–914)	324.0 ± 1.0 170	298 671	CGC A, E	[2008CHI/WAN] [1987STE/MAL, 1966KUD/ZWO]
C ₆₆ H ₁₃₄	[66563-47-1] Δ_vH	2-methylpentaheptacontane (599–906)	163.1	614		[1999DYK/SVO]
C ₆₆ H ₁₃₄ S	[66563-48-2] Δ_vH	1-hexaheptacontanethiol (607–920)	162.8	622	E	[1999DYK/SVO]
C ₆₇ H ₁₂₈	[66563-72-2] Δ_vH	heptaheptacontylbenzene (603–917)	162.1	618		[1999DYK/SVO]
C ₆₇ H ₁₃₄	[66563-73-3] Δ_vH	heptaheptacontylcyclohexane (603–917)	160.9	618		[1999DYK/SVO]
C ₆₇ H ₁₃₄	[66563-74-4] Δ_vH	1-heptaheptacontene (601–911)	162.3	616		[1999DYK/SVO]
C ₆₇ H ₁₃₆	[7719-90-6] Δ_vH	heptaheptacontane (659–937)	170.9	674	A, E	[1987STE/MAL, 1966KUD/ZWO]
C ₆₇ H ₁₃₆	[66563-75-5] Δ_vH	2-methylhexaheptacontane (601–909)	163.7	616		[1999DYK/SVO]
C ₆₇ H ₁₃₆ S	[66563-76-6] Δ_vH	1-heptaheptacontanethiol (609–922)	163.2	624	E	[1999DYK/SVO]
C ₆₈ H ₁₃₀	[66563-75-5] Δ_vH	doheptacontylbenzene (605–920)	162.6	620		[1999DYK/SVO]
C ₆₈ H ₁₃₆	[66563-78-8] Δ_vH	doheptacontylcyclohexane (605–920)	161.5	620		[1999DYK/SVO]
C ₆₈ H ₁₃₆	[66563-79-9] Δ_vH	1-octaheptacontene (603–913)	162.8	618		[1999DYK/SVO]
C ₆₈ H ₁₃₈	[7719-91-7] Δ_vH Δ_vH	octaheptacontane (323–523) (661–920)	331.9 ± 0.2 172.3	298 676	CGC A, E	[2008CHI/WAN] [1987STE/MAL, 1966KUD/ZWO]
C ₆₈ H ₁₃₈	[66563-80-2] Δ_vH	2-methylheptaheptacontane (603–912)	164.3	618		[1999DYK/SVO]
C ₆₈ H ₁₃₈ S	[66563-81-3] Δ_vH	1-octaheptacontanethiol (611–925)	163.6	626	E	[1999DYK/SVO]
C ₆₉ H ₁₃₂	[66563-82-4] Δ_vH	triheptacontylbenzene (607–923)	163.1	622		[1999DYK/SVO]
C ₆₉ H ₁₃₈	[66563-83-5] Δ_vH	1-nonheptacontene (605–916)	163.4	620		[1999DYK/SVO]

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₆₉ H ₁₃₈	[66563-93-7] $\Delta_v H$	trihexacontylcyclohexane (607–923)	162	622		[1999DYK/SVO]
C ₆₉ H ₁₄₀	[7719-92-8] $\Delta_v H$	nonahexacontane (664–923)	173.2	679	A, E	[1987STE/MAL, 1966KUD/ZWO]
C ₆₉ H ₁₄₀	[66563-94-8] $\Delta_v H$	2-methyloctahexacontane (605–914)	164.9	620		[1999DYK/SVO]
C ₆₉ H ₁₄₀ S	[66577-83-1] $\Delta_v H$	1-nonahexacontanethiol (612–928)	164.4	627	E	[1999DYK/SVO]
C ₇₀	[115383-22-7] $\Delta_{\text{sub}} H$	fullerene - C70 (864–1099)	199 ± 2	298	ME	[2000SCH/MAT]
	$\Delta_{\text{sub}} H$	(783–904)	189.8 ± 3.1	844	ME	[1996PIA/GIG]
	$\Delta_{\text{sub}} H$		200 ± 6.0	298		[1996PIA/GIG]
	$\Delta_{\text{sub}} H$		174 ± 3.0	740	ME	[1994POP/DRA]
	$\Delta_{\text{sub}} H$		193.4 ± 1.5	750	MS	[1994SAI/LAK]
	$\Delta_{\text{sub}} H$		186.6	788	ME	[1993MAT/SAI]
	$\Delta_{\text{sub}} H$	(673–873)	188.3 ± 4.2		ME	[1992ABR/OLA2]
	$\Delta_{\text{sub}} H$	(640–800)	180.0 ± 9.2	739	ME,MS	[1991PAN/SAM]
C ₇₀ H ₁₃₄	[66577-84-2] $\Delta_v H$	tetrahexacontylbenzene (609–925)	163.6	624		[1999DYK/SVO]
C ₇₀ H ₁₄₀	[66577-85-3] $\Delta_v H$	1-heptacontene (607–919)	163.9	622		[1999DYK/SVO]
C ₇₀ H ₁₄₀	[66577-86-4] $\Delta_v H$	tetrahexacontylcyclohexane (608–926)	162.8	623		[1999DYK/SVO]
C ₇₀ H ₁₄₂	[7719-93-9] $\Delta_v H$	heptacontane (323–523)	340.3 ± 0.3	298	CGC	[2008CHI/WAN]
	$\Delta_v H$	(666–926)	174.4	681	A, E	[1987STE/MAL, 1966KUD/ZWO]
C ₇₀ H ₁₄₂	[66577-87-5] $\Delta_v H$	2-methylnonahexacontane (607–917)	165.4	622		[1999DYK/SVO]
C ₇₀ H ₁₄₂ S	[66577-88-6] $\Delta_v H$	1-heptacontanethiol (614–930)	164.8	621	E	[1999DYK/SVO]
C ₇₁ H ₁₃₆	[66577-89-7] $\Delta_v H$	pentaheptacontylbenzene (611–928)	164.4	626		[1999DYK/SVO]
C ₇₁ H ₁₄₂	[66577-90-0] $\Delta_v H$	1-henheptacontene (609–922)	164.4	624		[1999DYK/SVO]
C ₇₁ H ₁₄₂	[66577-91-1] $\Delta_v H$	pentaheptacontylcyclohexane (610–928)	163.3	625		[1999DYK/SVO]
C ₇₁ H ₁₄₄	[7667-82-5] $\Delta_v H$	henheptacontane (669–928)	175.2	684	A, E	[1987STE/MAL, 1966KUD/ZWO]
C ₇₁ H ₁₄₄	[66577-92-2] $\Delta_v H$	2-methylheptacontane (609–920)	165.9	624		[1999DYK/SVO]
C ₇₁ H ₁₄₄ S	[66577-93-3] $\Delta_v H$	1-henheptacontanethiol (616–933)	165.1	631	E	[1999DYK/SVO]
C ₇₂ H ₉₆ O ₄₈	[na] $\Delta_{\text{fus}} H$	hexakis(2,3,4-tri-O-acetyl)- α -cyclodextrin	74.37	505.7		[2006BET/SOR]

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₇₂ H ₁₃₈	[66577-94-4] $\Delta_v H$	hexaheptacontylbenzene (613–931)	164.5	628		[1999DYK/SVO]
C ₇₂ H ₁₄₄	[66577-95-5] $\Delta_v H$	1-doheptacontene (610–924)	165.3	625		[1999DYK/SVO]
C ₇₂ H ₁₄₄	[66577-96-6] $\Delta_v H$	hexaheptacontylcyclohexane (612–931)	163.8	627		[1999DYK/SVO]
C ₇₂ H ₁₄₆	[7668-83-6] $\Delta_v H$ $\Delta_v H$	doheptacontane (323–523) (671–931)	348.4 ± 0.3 176.4	298 686	CGC A, E	[2008CHI/WAN] [1987STE/MAL, 1966KUD/ZWO]
C ₇₂ H ₁₄₆	[66577-97-7] $\Delta_v H$	2-methylheptacontane (611–923)	166.4	626		[1999DYK/SVO]
C ₇₂ H ₁₄₆ S	[66577-98-8] $\Delta_v H$	1-doheptacontanethiol (617–935)	165.8	632	E	[1999DYK/SVO]
C ₇₃ H ₁₀₈ O ₁₂	[6683-19-8] $\Delta_{\text{fus}} H$	tetrakis[methylene-3-(3,5-di- <i>tert</i> -butyl-4-hydroxyphenyl)propionate]methane 65.95	385.8	DSC		[2009WEI/CHE]
C ₇₃ H ₁₄₀	[66577-99-9] $\Delta_v H$	heptaheptacontylbenzene (614–933)	165.4	629		[1999DYK/SVO]
C ₇₃ H ₁₄₆	[66578-00-5] $\Delta_v H$	heptaheptacontylcyclohexane (614–933)	164.2	629		[1999DYK/SVO]
C ₇₃ H ₁₄₆	[66578-01-6] $\Delta_v H$	1-triheptacontene (612–927)	165.7	627		[1999DYK/SVO]
C ₇₃ H ₁₄₈	[7667-84-7] $\Delta_v H$	triheptacontane (674–934)	177.1	689	A, E	[1987STE/MAL, 1966KUD/ZWO]
C ₇₃ H ₁₄₈	[66578-02-7] $\Delta_v H$	2-methylheptacontane (613–926)	166.9	628		[1999DYK/SVO]
C ₇₃ H ₁₄₈ S	[66577-64-8] $\Delta_v H$	1-triheptacontanethiol (619–938)	166.2	634	E	[1999DYK/SVO]
C ₇₄ H ₁₄₂	[66577-65-9] $\Delta_v H$	octaheptacontylbenzene (616–936)	165.8	631		[1999DYK/SVO]
C ₇₄ H ₁₄₈	[66577-66-0] $\Delta_v H$	octaheptacontylcyclohexane (615–936)	165	630		[1999DYK/SVO]
C ₇₄ H ₁₄₈	[66577-67-1] $\Delta_v H$	1-tetraheptacontene (614–930)	166.2	629		[1999DYK/SVO]
C ₇₄ H ₁₅₀	[7667-85-8] $\Delta_v H$ $\Delta_v H$	tetraheptacontane (323–523) (676–936)	356.2 ± 0.1 178.2	298 691	CGC A, E	[2008CHI/WAN] [1987STE/MAL, 1966KUD/ZWO]
C ₇₄ H ₁₅₀	[66577-68-2] $\Delta_v H$	2-methyltriheptacontane (615–928)	167.4	630		[1999DYK/SVO]
C ₇₄ H ₁₅₀ S	[66577-69-3] $\Delta_v H$	1-tetraheptacontanethiol (620–940)	166.9	635	E	[1999DYK/SVO]
C ₇₅ H ₁₄₄	[66577-70-6] $\Delta_v H$	nonaheptacontylbenzene (618–938)	166.3	633		[1999DYK/SVO]
C ₇₅ H ₁₅₀	[66577-71-7]	nonaheptacontylcyclohexane				

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(617–632)	165.4	632		[1999DYK/SVO]
C ₇₅ H ₁₅₀	[66577-72-8] $\Delta_v H$	1-pentaheptacontene (616–932)	166.7	631		[1999DYK/SVO]
C ₇₅ H ₁₅₀ N ₆	[106486-50-4] $\Delta_{\text{fus}} H$	<i>tris</i> N,N-didodecylamino-1,3,5-triazine	119.19	320.3		[1986LAT/HOE]
C ₇₅ H ₁₅₂	[7667-86-9] $\Delta_v H$	pentaheptacontane (678–939)	179.4	693	A, E	[1987STE/MAL, 1966KUD/ZWO]
C ₇₅ H ₁₅₂	[66577-73-9] $\Delta_v H$	2-methyltetraheptacontane (616–931)	168.2	631		[1999DYK/SVO]
C ₇₅ H ₁₅₂ S	[66577-74-0] $\Delta_v H$	1-pentaheptacontanethiol (622–942)	167.2	637	E	[1999DYK/SVO]
C ₇₆	[135113-15-4] $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$	fullerene - C76 (637–911) (834–1069)	190 ± 7 206 ± 4.0	764 298	ME TE	[1998BOL/MAR] [1997BRU/GIG]
C ₇₆ H ₉₄ N ₄	[89372-90-7] $\Delta_{\text{sub}} H$	5,10,15,20- <i>tetrakis</i> (3,5- <i>di-tert</i> -butylphenyl)porphine	209 ± 5			[2000PER/GOL]
C ₇₆ H ₁₄₆	[66577-75-1] $\Delta_v H$	heptacontylbenzene (619–941)	167	634		[1999DYK/SVO]
C ₇₆ H ₁₅₂	[66577-76-2] $\Delta_v H$	heptacontylcyclohexane (619–941)	165.8	634		[1999DYK/SVO]
C ₇₆ H ₁₅₂	[66577-77-3] $\Delta_v H$	1-hexaheptacontene (617–935)	167.5	632		[1999DYK/SVO]
C ₇₆ H ₁₅₄	[7667-87-0] $\Delta_v H$ $\Delta_v H$	hexaheptacontane (323–523) (680–941)	364.3 ± 0.3 180.4	298 695	CGC A, E	[2008CHI/WAN] [1987STE/MAL, 1966KUD/ZWO]
C ₇₆ H ₁₅₄	[66577-78-4] $\Delta_v H$	2-methylpentaheptacontane (618–933)	168.7	633		[1999DYK/SVO]
C ₇₆ H ₁₅₄ S	[66577-79-5] $\Delta_v H$	1-hexaheptacontanethiol (623–945)	169.8	638		[1999DYK/SVO]
C ₇₇ H ₁₄₈	[66577-80-8] $\Delta_v H$	henheptacontylbenzene (621–943)	167.4	636		[1999DYK/SVO]
C ₇₇ H ₁₅₄	[66577-81-9] $\Delta_v H$	henheptacontylcyclohexane (620–943)	166.6	635		[1999DYK/SVO]
C ₇₇ H ₁₅₄	[66577-82-0] $\Delta_v H$	1-heptaheptacontene (619–937)	167.9	634		[1999DYK/SVO]
C ₇₇ H ₁₅₆	[7719-94-0] $\Delta_v H$	heptaheptacontane (682–944)	181.4	697	A, E	[1987STE/MAL, 1966KUD/ZWO]
C ₇₇ H ₁₅₆	[66575-56-2] $\Delta_v H$	2-methylhexaheptacontane (620–936)	169.1	635		[1999DYK/SVO]
C ₇₇ H ₁₅₆ S	[66575-57-3] $\Delta_v H$	1-heptaheptacontanethiol (625–947)	168.2	640	E	[1999DYK/SVO]
C ₇₈ H ₁₀₈	[125594-11-8] $\Delta_{\text{fus}} H$	2,3,6,7,10,11- <i>hexakis</i> (1-decynyl)triphenylene	63	314.2	DSC	[1996DOM/HEA, 1990PRA/KOH]

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
C ₇₈ H ₁₅₀	[66327-30-8] $\Delta_{\text{v}}H$	doheptacontylbenzene (622–945)	167.3	637		[1999DYK/SVO]
C ₇₈ H ₁₅₆	[66327-31-9] $\Delta_{\text{v}}H$	doheptacontylcyclohexane (622–945)	166.9	637		[1999DYK/SVO]
C ₇₈ H ₁₅₆	[66327-32-0] $\Delta_{\text{v}}H$	1-octaheptacontene (621–940)	168.3	636		[1999DYK/SVO]
C ₇₈ H ₁₅₈	[7719-85-9] $\Delta_{\text{v}}H$ $\Delta_{\text{v}}H$	octaheptacontane (638–691) (685–946)	372.1 ± 3.7 181.8	298 700	CGC A, E	[2008CHI/LIP] [1987STE/MAL, 1966KUD/ZWO]
C ₇₈ H ₁₅₈	[66327-33-1] $\Delta_{\text{v}}H$	2-methylheptaheptacontane (621–939)	169.9	636		[1999DYK/SVO]
C ₇₈ H ₁₅₈ S	[66375-13-1] $\Delta_{\text{v}}H$	1-octaheptacontanethiol (626–949)	168.8	641	E	[1999DYK/SVO]
C ₇₉ H ₁₅₂	[66327-34-2] $\Delta_{\text{v}}H$	triheptacontylbenzene (623–947)	168	638		[1999DYK/SVO]
C ₇₉ H ₁₅₈	[66327-35-3] $\Delta_{\text{v}}H$	1-nonaheptacontene (622–942)	169.1	637		[1999DYK/SVO]
C ₇₉ H ₁₅₈	[66327-36-4] $\Delta_{\text{v}}H$	triheptacontylcyclohexane (623–948)	167.7	638		[1999DYK/SVO]
C ₇₉ H ₁₆₀	[7719-86-0] $\Delta_{\text{v}}H$	nonaheptacontane (687–949)	182.7	702	A, E	[1987STE/MAL, 1966KUD/ZWO]
C ₇₉ H ₁₆₀	[66327-37-5] $\Delta_{\text{v}}H$	2-methyloctaheptacontane (622–940)	167.8	637		[1999DYK/SVO]
C ₇₉ H ₁₆₀ S	[66327-38-6] $\Delta_{\text{v}}H$	1-nonaheptacontanethiol (628–952)	169.1	643	E	[1999DYK/SVO]
C ₈₀ H ₁₅₄	[66327-39-7] $\Delta_{\text{v}}H$	tetraheptacontylbenzene (625–949)	168.4	640		[1999DYK/SVO]
C ₈₀ H ₁₆₀	[66327-40-0] $\Delta_{\text{v}}H$	1-octacontene (624–945)	169.4	639		[1999DYK/SVO]
C ₈₀ H ₁₆₀	[66327-41-1] $\Delta_{\text{v}}H$	tetraheptacontylcyclohexane (625–950)	168	640		[1999DYK/SVO]
C ₈₀ H ₁₆₂	[7667-88-1] $\Delta_{\text{v}}H$ $\Delta_{\text{v}}H$	octacontane (638–691) (689–951)	379.6 ± 3.8 183.6	298 704	CGC A, E	[2008CHI/LIP] [1987STE/MAL, 1966KUD/ZWO]
C ₈₀ H ₁₆₂	[66327-42-2] $\Delta_{\text{v}}H$	2-methylnonaheptacontane (624–943)	170.2	639		[1999DYK/SVO]
C ₈₀ H ₁₆₂ S	[66327-43-3] $\Delta_{\text{v}}H$	1-octacontanethiol (629–954)	169.6	644	E	[1999DYK/SVO]
C ₈₁ H ₁₅₆	[66327-44-4] $\Delta_{\text{v}}H$	pentaheptacontylbenzene (636–952)	169.1	641		[1999DYK/SVO]
C ₈₁ H ₁₆₂	[66327-45-5] $\Delta_{\text{v}}H$	1-henocentacontene (625–946)	169.3	640		[1999DYK/SVO]
C ₈₁ H ₁₆₂	[66327-46-6]	pentaheptacontylcyclohexane				

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(627–952)	168.4	642		[1999DYK/SVO]
C ₈₁ H ₁₆₄	[7667-89-2]	henoctacontane				
	$\Delta_v H$	(691–953)	184.5	706	A, E	[1987STE/MAL, 1966KUD/ZWO]
C ₈₁ H ₁₆₄	[66327-47-7]	2-methyloctacontane				
	$\Delta_v H$	(625–945)	170.9	640		[1999DYK/SVO]
C ₈₁ H ₁₆₄ S	[66327-48-8]	1-henoctacontanethiol				
	$\Delta_v H$	(630–955)	169.4	645	E	[1999DYK/SVO]
C ₈₂ H ₁₅₈	[66327-49-9]	hexaheptacontylbenzene				
	$\Delta_v H$	(628–954)	169.4	643		[1999DYK/SVO]
C ₈₂ H ₁₆₄	[66327-50-2]	1-dooctacontene				
	$\Delta_v H$	(626–949)	157.6	641		[1999DYK/SVO]
C ₈₂ H ₁₆₄	[66327-09-1]	hexaheptacontylcyclohexane				
	$\Delta_v H$	(627–954)	168.5	642		[1999DYK/SVO]
C ₈₂ H ₁₆₆	[7719-95-1]	dooctacontane				
	$\Delta_v H$	(638–691)	387.2 ± 3.8	298	CGC	[2008CHI/LIP]
	$\Delta_v H$	(693–955)	185.3	708	A, E	[1987STE/MAL, 1966KUD/ZWO]
C ₈₂ H ₁₆₆	[66327-10-4]	2-methylhenoctacontane				
	$\Delta_v H$	(627–947)	171.3	642		[1999DYK/SVO]
C ₈₂ H ₁₆₆ S	[66327-11-5]	1-dooctacontanethiol				
	$\Delta_v H$	(631–957)	170	646	E	[1999DYK/SVO]
C ₈₃ H ₁₆₀	[66327-12-6]	heptaheptacontylbenzene				
	$\Delta_v H$	(628–955)	169.6	643		[1999DYK/SVO]
C ₈₃ H ₁₆₆	[66327-13-7]	heptaheptacontylcyclohexane				
	$\Delta_v H$	(629–956)	168.9	644		[1999DYK/SVO]
C ₈₃ H ₁₆₆	[66327-14-8]	1-trioctacontene				
	$\Delta_v H$	(628–951)	170.4	643		[1999DYK/SVO]
C ₈₃ H ₁₆₈	[7667-90-5]	trioctacontane				
	$\Delta_v H$	(694–957)	186.5	709	A, E	[1987STE/MAL, 1966KUD/ZWO]
C ₈₃ H ₁₆₈	[66327-15-9]	2-methyldooctacontane				
	$\Delta_v H$	(628–949)	171.1	643		[1999DYK/SVO]
C ₈₃ H ₁₆₈ S	[66327-16-0]	1-trioctacontanethiol				
	$\Delta_v H$	(633–959)	170.2	648	E	[1999DYK/SVO]
C ₈₄	[135113-16-5]	fullerene - C84				
	$\Delta_{\text{sub}} H$	(658–980)	202 ± 4.0	853	ME	[1998BOL/MAR2]
	$\Delta_{\text{sub}} H$	(920–1190)	210 ± 6	950	TE	[1997PIA/PAL]
C ₈₄ H ₁₁₂ O ₅₆	[na]	heptakis(2,3,6-tri-O-acetyl)- β -cyclodextrin				
	$\Delta_{\text{fus}} H$		82.73	491.7		[2006BET/SOR]
C ₈₄ H ₁₆₂	[66327-17-1]	octaheptacontylbenzene				
	$\Delta_v H$	(630–957)	169.9	645		[1999DYK/SVO]
C ₈₄ H ₁₆₈	[66327-18-2]	octaheptacontylcyclohexane				
	$\Delta_v H$	(630–958)	169.5	645		[1999DYK/SVO]
C ₈₄ H ₁₆₈	[66327-19-3]	1-tetraoctacontene				
	$\Delta_v H$	(629–953)	170.1	644		[1999DYK/SVO]

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₈₄ H ₁₇₀	[7667-91-6]	tetraoctacontane				
	$\Delta_v H$	(638–691)	394.0 ± 3.9	298	CGC	[2008CHI/LIP]
	$\Delta_v H$	(696–960)	187.3	711	A, E	[1987STE/MAL, 1966KUD/ZWO]
C ₈₄ H ₁₇₀	[66327-20-6]	2-methyltrioctacontane				
	$\Delta_v H$	(629–951)	171.8	644		[1999DYK/SVO]
C ₈₄ H ₁₇₀ S	[66327-21-7]	1-tetraoctacontanethiol				
	$\Delta_v H$	(634–962)	170.8	649	E	[1999DYK/SVO]
C ₈₅ H ₁₆₄	[66327-22-8]	nonheptacontylbenzene				
	$\Delta_v H$	(631–960)	170.6	646		[1999DYK/SVO]
C ₈₅ H ₁₇₀	[66327-23-9]	nonheptacontylcyclohexane				
	$\Delta_v H$	(632–960)	169.8	647		[1999DYK/SVO]
C ₈₅ H ₁₇₀	[66327-24-0]	1-pentaoctacontene				
	$\Delta_v H$	(630–955)	170.9	645		[1999DYK/SVO]
C ₈₅ H ₁₇₂	[7719-96-2]	pentaoctacontane				
	$\Delta_v H$	(698–962)	187.9	713	A, E	[1987STE/MAL, 1966KUD/ZWO]
C ₈₅ H ₁₇₂	[66327-25-1]	2-methyltetraoctacontane				
	$\Delta_v H$	(631–953)	172.1	646		[1999DYK/SVO]
C ₈₅ H ₁₇₂ S	[66327-26-2]	1-pentaoctacontanethiol				
	$\Delta_v H$	(634–963)	170.8	649	E	[1999DYK/SVO]
C ₈₆ H ₁₆₆	[66327-27-3]	octacontylbenzene				
	$\Delta_v H$	(633–962)	170.9	648		[1999DYK/SVO]
C ₈₆ H ₁₇₂	[66327-28-4]	1-hexaoctacontene				
	$\Delta_v H$	(631–957)	171.5	646		[1999DYK/SVO]
C ₈₆ H ₁₇₂	[66327-29-5]	octacontylcyclohexane				
	$\Delta_v H$	(632–962)	169.9	647		[1999DYK/SVO]
C ₈₆ H ₁₇₄	[7667-92-7]	hexaoctacontane				
	$\Delta_v H$	(638–691)	402.1 ± 4.0	298	CGC	[2008CHI/LIP]
	$\Delta_v H$	(700–964)	188.6	715	A, E	[1987STE/MAL, 1966KUD/ZWO]
C ₈₆ H ₁₇₄	[66426-88-3]	2-methylpentaoctacontane				
	$\Delta_v H$	(632–956)	172.8	647		[1999DYK/SVO]
C ₈₆ H ₁₇₄ S	[66326-89-4]	1-hexaoctacontanethiol				
	$\Delta_v H$	(636–965)	171.1	651	E	[1999DYK/SVO]
C ₈₇ H ₁₆₈	[66326-90-7]	henoctacontylbenzene				
	$\Delta_v H$	(633–963)	171	648		[1999DYK/SVO]
C ₈₇ H ₁₇₄	[66326-91-8]	henoctacontylcyclohexane				
	$\Delta_v H$	(633–964)	170.6	648		[1999DYK/SVO]
C ₈₇ H ₁₇₄	[66326-92-9]	1-heptaoctacontene				
	$\Delta_v H$	(633–959)	171.8	648		[1999DYK/SVO]
C ₈₇ H ₁₇₆	[7667-93-8]	heptaoctacontane				
	$\Delta_v H$	(702–966)	189.3	717	A, E	[1987STE/MAL, 1966KUD/ZWO]
C ₈₇ H ₁₇₆	[66326-93-0]	2-methylhexaoctacontane				
	$\Delta_v H$	(633–957)	172.6	648		[1999DYK/SVO]
C ₈₇ H ₁₇₆ S	[66326-94-1]	1-heptaoctacontanethiol				

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(637–967)	171.7	652	E	[1999DYK/SVO]
C ₈₄ H ₁₄₄ O ₆	[na]	2,3,6,7,10,11-hexakis[(2R,4R,6R)-2,4,6-trimethyloctyl]oxy]-triphenylene				
	$\Delta_{\text{us}} H$		38	141.2		
	$\Delta_{\text{fus}} H$		5.2	237.2		[2002SCH/LAS]
C ₈₈ H ₁₇₀	[66326-95-2]	doioctacontylbenzene				
	$\Delta_v H$	(635–965)	171.3	650		[1999DYK/SVO]
C ₈₈ H ₁₇₆	[66326-96-3]	doioctacontylcyclohexane				
	$\Delta_v H$	(635–966)	170.8	650		[1999DYK/SVO]
C ₈₈ H ₁₇₆	[66326-97-4]	1-octaocantene				
	$\Delta_v H$	(634–961)	172.5	649		[1999DYK/SVO]
C ₈₈ H ₁₇₈	[7667-94-9]	octaocantane				
	$\Delta_v H$	(638–691)	409.2 ± 4.1	298	CGC	[2008CHI/LIP]
	$\Delta_v H$	(703–967)	190.4	718	A, E	[1987STE/MAL, 1966KUD/ZWO]
C ₈₈ H ₁₇₈	[66326-98-5]	2-methylheptaocantane				
	$\Delta_v H$	(634–959)	173.3	649		[1999DYK/SVO]
C ₈₈ H ₁₇₈ S	[66326-99-6]	1-octaocantethiol				
	$\Delta_v H$	(639–969)	171.9	654	E	[1999DYK/SVO]
C ₈₉ H ₁₇₂	[66327-00-2]	trioctacontylbenzene				
	$\Delta_v H$	(636–967)	172	651		[1999DYK/SVO]
C ₈₉ H ₁₇₈	[66327-01-3]	1-nonaocantene				
	$\Delta_v H$	(635–962)	172.3	650		[1999DYK/SVO]
C ₈₉ H ₁₇₈	[66327-02-4]	trioctacontylcyclohexane				
	$\Delta_v H$	(636–968)	171.5	651		[1999DYK/SVO]
C ₈₉ H ₁₈₀	[7719-76-8]	nonaocantane				
	$\Delta_v H$	(705–969)	190.9	720	A, E	[1987STE/MAL, 1966KUD/ZWO]
C ₈₉ H ₁₈₀	[66327-03-5]	2-methyloctaocantane				
	$\Delta_v H$	(636–962)	173.6	651		[1999DYK/SVO]
C ₈₉ H ₁₈₀ S	[66327-04-6]	1-nonaocantethiol				
	$\Delta_v H$	(639–970)	171.9	654	E	[1999DYK/SVO]
C ₉₀ H ₁₇₄	[66327-05-7]	tetraoctacontylbenzene				
	$\Delta_v H$	(637–968)	171.7	652		[1999DYK/SVO]
C ₉₀ H ₁₈₂	[7667-51-8]	nonaocantane				
	$\Delta_v H$	(638–691)	416.4 ± 4.3	298	CGC	[2008CHI/LIP]
	$\Delta_v H$	(707–971)	191.6	722	A, E	[1987STE/MAL, 1966KUD/ZWO]
C ₉₁ H ₁₈₄	[7719-97-3]	hennaocantane				
	$\Delta_v H$	(708–973)	192.5	723	A, E	[1987STE/MAL, 1966KUD/ZWO]
C ₉₂ H ₁₈₆	[7667-95-0]	dononaocantane				
	$\Delta_v H$	(638–691)	424.5 ± 4.0	298	CGC	[2008CHI/LIP]
	$\Delta_v H$	(710–975)	193	725	A, E	[1987STE/MAL, 1966KUD/ZWO]
C ₉₂ H ₁₈₆ O ₄₆	[na]	1,ω-dimethoxypentatetracos(oxyethylene)				
	$\Delta_{\text{fus}} H$		374.8	324.2	DSC	[1996YAN/YU]
C ₉₃ H ₁₈₈	[7667-96-1]	trinonaocantane				
	$\Delta_v H$	(711–977)	194.1	726	A, E	[1987STE/MAL, 1966KUD/ZWO]

TABLE 11. Phase change enthalpies of C₂₁ to C₁₉₂ organic compounds—Continued

Molecular Formula	CAS Reg No	Compound				
	Enthalpy	Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
C ₉₄ H ₁₉₀	[1574-32-9] $\Delta_{\text{v}}H$	tetranonacontane (713–978)	194.5	728	A, E	[1987STE/MAL, 1966KUD/ZWO]
C ₉₅ H ₁₉₂	[7667-97-2] $\Delta_{\text{v}}H$	pentanonacontane (714–980)	195.4	729	A, E	[1987STE/MAL, 1966KUD/ZWO]
C ₉₆ H ₁₉₄	[7763-13-5] $\Delta_{\text{v}}H$	hexanonacontane (716–982)	195.8	731	A, E	[1987STE/MAL, 1966KUD/ZWO]
C ₉₇ H ₁₉₆	[7670-25-9] $\Delta_{\text{v}}H$	heptanonacontane (717–983)	196.6	732	A, E	[1987STE/MAL, 1966KUD/ZWO]
C ₉₈ H ₁₉₈	[7670-26-0] $\Delta_{\text{v}}H$	octanonacontane (719–985)	196.9	734	A, E	[1987STE/MAL, 1966KUD/ZWO]
C ₉₉ H ₂₀₀	[7670-27-1] $\Delta_{\text{v}}H$	nonanonacontane (720–986)	197.8	735	A, E	[1987STE/MAL, 1966KUD/ZWO]
C ₁₀₀ H ₂₀₂	[6703-98-6] $\Delta_{\text{us}}H$	hectane (721–988)	54.8	365.5		
	$\Delta_{\text{fus}}H$		331.8	338.5		[1970HAY]
	$\Delta_{\text{v}}H$		198.5	736	A, E	[1987STE/MAL, 1966KUD/ZWO]
C ₁₀₂ H ₁₈₀ O ₆	[501447-89-8] $\Delta_{\text{us}}H$	2,3,6,7,10,11- <i>hexakis</i> [[<i>(2R,4R,6R,8R)</i> -2,4,6,8-tetramethyldecyl]oxy]triphenylene	23.4	139.2		
	$\Delta_{\text{fus}}H$		6.6	236.2	DSC	[2002SCH/LAS]
C ₁₉₂ H ₃₈₆	[96123-38-5] $\Delta_{\text{fus}}H$	<i>n</i> -dononacontahectane	661.1	399.1	DSC	[1989STA/MAN]

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds

Molecular Formula	CAS Reg No Enthalpy	Compound				Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)			
Ag (silver)							
(C ₅ H ₁₀ O ₂ Ag) ₂	[7324-58-5] $\Delta_{\text{sub}}H$	silver 2,2-dimethylpropanoate (dimer)	146.9 ± 8.9				[2001MAL/PAR]
Al (aluminum)							
C ₃ H ₉ Al	[75-24-1] $\Delta_{\text{fus}}H$	trimethylaluminum	17.1			Sub-Vap	[2003FUL/RUZ]
		Note: Authors noted the large discrepancy between their calculated value based on the difference in the enthalpy of sublimation minus the enthalpy of vaporization and the published literature data.					
	$\Delta_{\text{fus}}H$		8.79	288.4			[1996DOM/HEA]
	$\Delta_{\text{sub}}H$	(243–285)	60.1				[2003FUL/RUZ]
	Δ_vH	(288–293)	43.0				[2003FUL/RUZ]
	Δ_vH	(336–400)	39.8	351			[1963MCC/MES]
	Δ_vH		63.2 ± 1.7				[1963MOR/SEL, 1982PIL/SKI]
	Δ_vH		41.1			BG	[1946BAM/LEV]
C ₄ H ₁₀ AlCl	[96-10-6] Δ_vH	diethylaluminum chloride					
	Δ_vH	(278–318)	50.5	293			[1991BUC/POT]
	Δ_vH	(273–473)	53.9	373			[1991BUC/POT]
C ₄ H ₁₁ Al	[871-27-2] Δ_vH	diethylaluminum hydride					
	Δ_vH		57.7 ± 2.1				[1967PAW, 1982PIL/SKI, 1965SHA/SCH]
	Δ_vH		46.9				[1965SHA/SCH]
C ₅ H ₅ AlBr ₃ N	[15348-61-5] $\Delta_{\text{sub}}H$	aluminum tribromide- pyridine complex					
	$\Delta_{\text{sub}}H$	(501–633)	71.2 ± 0.6			T	[1989GRI/KON]
	$\Delta_{\text{sub}}H$		83.3			B,E	[1967WIL/WAR]
C ₆ H ₁₅ Al	[97-93-8] $\Delta_{\text{fus}}H$	triethylaluminum					
	$\Delta_{\text{fus}}H$		10.6	225			[1996DOM/HEA]
	Δ_vH		73.2 ± 2.1				[1967PAW, 1982PIL/SKI, 1965SHA/SCH]
	Δ_vH		60.2				[1965SHA/SCH]
	Δ_vH		54.1			BG	[1946BAM/LEV]
C ₆ H ₁₅ AlO	[1586-92-1] Δ_vH	diethylaluminum ethoxide					
	Δ_vH	(403–463)	48.7 ± 0.8	433			[1974SHM/GOL]
C ₇ H ₁₇ AlO	[6083-26-7] Δ_vH	diethylaluminum propoxide					
	Δ_vH	(398–463)	51.0 ± 0.8	430			[1974SHM/GOL]
C ₈ H ₁₉ Al	[1191-15-7] Δ_vH	diisobutylaluminum hydride					
	Δ_vH		42.3 ± 2.1				[1967PAW, 1982PIL/SKI, 1965SHA/SCH]
	Δ_vH		35.6				[1965SHA/SCH]
C ₉ H ₂₁ Al	[102-67-0] Δ_vH	tripropylaluminum					
	Δ_vH		43 ± 2.0				[2002BAE/SHI]
	Δ_vH		42.5 ± 1.2				[1967PAW, 1982PIL/SKI]
C ₉ H ₂₁ AlO ₃	[555-31-7] Δ_vH	aluminum isopropoxide					
	Δ_vH	(353–399)	48.1 ± 6.3	376			[1972BLE/FIE]
C ₁₂ H ₂₇ Al	[100-99-2] Δ_vH	triisobutylaluminum					
	Δ_vH	(273–322)	38.3	298			[1964SHA/TUB]
C ₁₂ H ₂₇ AlO ₃	[3085-30-1] Δ_vH	tributoxyaluminum					
	Δ_vH	(503–533)	104.1	518		A, I	[1987STE/MAL, 1957WIL]

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound		T_m (K)	Method	Reference	
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)				
$C_{12}H_{27}AlO_3$	[3453-79-0] $\Delta_v H$	triisobutoxyaluminum (500–550)	139.4	515	A, I	[1987STE/MAL, 1957WIL]	
$C_{12}H_{27}AlO_3$	[2269-22-9] $\Delta_v H$	tri-sec-butoxyaluminum (425–469)	81.5	440	A, I	[1987STE/MAL, 1957WIL]	
$C_{15}H_3AlF_{18}O_6$	[17786-67-3] $\Delta_{\text{sub}}H$	<i>tris</i> (1,1,1,5,5,5-hexafluoro-2,4-pentanedionato)aluminum(III) (333–363)	52.0		TGA	[2000FAH/BAR]	
	$\Delta_{\text{sub}}H$		(324–344)	77.6 ± 6.2	334	BG	[1987GRI/LAZ2, 1988LAZ/GRI]
	$\Delta_{\text{sub}}H$			79.0 ± 6.5	298		[1987GRI/LAZ2]
	$\Delta_{\text{sub}}H$			74.1 ± 2.5			[1985IGU/GER, 1987GRI/LAZ2]
	$\Delta_{\text{sub}}H$		(323–347)	109.6 ± 3.8	335		[1972FON/POM]
$C_{15}H_{12}AlF_9O_6$	[14354-59-7] $\Delta_{\text{sub}}H$	<i>tris</i> (1,1,1-trifluoro-2,4-pentanedionato)aluminum(III) (373–403)	74.0		TGA	[2000FAH/BAR]	
	$\Delta_{\text{sub}}H$		(363–423)	113.4 ± 1.3		GS	[1985MAT/KUW]
	$\Delta_{\text{sub}}H$		(369–392)	102.7 ± 3.2			[1978IGU/CHU2]
	$\Delta_{\text{sub}}H$			108 ± 2.0			[1977NAG, 1988GOL/SIT]
	$\Delta_{\text{sub}}H$			43.1	443		[1977VOL/MAZ]
	$\Delta_{\text{sub}}H$		(354–396)	93.7 ± 6.7	375		[1972FON/POM]
	$\Delta_{\text{sub}}H$			41.0			[1965FRA]
	$\Delta_{\text{sub}}H$			40.0			[1960BER/TRU, 1965BER/TRU]
	$\Delta_v H$		(349–411)	58.7 ± 0.7	380	BG	[1988LAZ/GRI]
	$\Delta_v H$		(392–484)	69.6 ± 0.5	438		[1978IGU/CHU2]
$C_{15}H_{21}AlO_6$	[13963-57-0] $\Delta_{\text{fus}}H$	<i>tris</i> (pentane-2,4-dionato)aluminum(III) 35.2	35.2	463	DSC	[2004SAB/MAR]	
	$\Delta_{\text{fus}}H$			32.7	458		[1971BEE/LIN2]
	$\Delta_{\text{fus}}H$			33.7	466.7		[1988LAZ/GRI]
	$\Delta_{\text{sub}}H$			107.1		DTA,TGA	[2009GAI/KUN]
	$\Delta_{\text{sub}}H$		(345–410)	101.8	378	ME	[2007SID/SID]
	$\Delta_{\text{sub}}H$		(376–467)	121.8 ± 1.5	298		[2006SEM/IGU]
	$\Delta_{\text{sub}}H$		(413–443)	93		TGA	[2000FAH/BAR]
	$\Delta_{\text{sub}}H$			120 ± 3.0	298	ME	[1977NAG, 1988RIB/FER4]
	$\Delta_{\text{sub}}H$		(432–464)	102.0 ± 3.2	448	BG	[1988LAZ/GRI]
	$\Delta_{\text{sub}}H$			47.1 ± 1.0			[1981TEG/FER]
	$\Delta_{\text{sub}}H$			118.9 ± 7.9			[1980SAC/HIL]
	$\Delta_{\text{sub}}H$			24.3	458		[1977VOL/MAZ]
	$\Delta_{\text{sub}}H$			121.7 ± 4.2	298		[1975IRV/RIB2]
	$\Delta_{\text{sub}}H$		(383–413)	66.1 ± 3.3	398		[1972FON/POM]
	$\Delta_{\text{sub}}H$			23.4			[1965FRA]
	$\Delta_{\text{sub}}H$		(417–476)	20.5			[1960BER/TRU, 1965BER/TRU]
	$\Delta_v H$			80.2		DTA,TGA	[2009GAI/KUN]
	$\Delta_v H$		(430–530)	78.7 ± 0.9	298	T	[1986GRI/LAZ]
$C_{16}H_{40}Al_2N_2$	[115381-27-6] $\Delta_{\text{sub}}H$		tetramethyl <i>bis</i> [μ -[N-(1-methylethyl)-2-propanaminto]]dialuminum(III) 99.2	99.2		ME	[1988BRA/FAK]
$C_{18}H_{15}Al$	[841-76-9] $\Delta_{\text{sub}}H$		triphenylaluminum (432–477)	172 ± 5	455	ME,TE	[1984GOV/KAN]
$C_{24}H_{12}AlF_9O_6S_3$	[14054-83-2] $\Delta_{\text{sub}}H$	<i>tris</i> (1-(2-thenoyl)-4,4,4-trifluoro-1-,3-butanedione)aluminum(III) U 46.4	U 46.4			[1960BER/TRU, 1965BER/TRU]	
$C_{27}H_{18}AlN_3O_3$	[2085-33-8] $\Delta_{\text{sub}}H$	<i>tris</i> (8-hydroxyquinolinato)aluminum(III) 137.7	137.7		TGA	[1995YAS/TAK]	

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
$C_{30}H_{18}AlF_9O_6$	[14323-12-7] $\Delta_{\text{sub}}H$	<i>tris</i> (1-phenyl-4,4,4-trifluoro-1,3-butanedione)aluminum(III) U 55.2 [1960BER/TRU, 1965BER/TRU]				
$C_{30}H_{27}AlO_6$	[14376-06-8] $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$	<i>tris</i> (1-phenyl-1,3-butanedionato)aluminum(III) (462–478) 186.8 ± 2.1 470 ME,TE [1995RIB/MON2] 195.2 ± 2.1 298 [1995RIB/MON2] 193.7 ± 0.3 298 [1983RIB/REI]				
$C_{30}H_{30}F_{21}AlO_6$	[18716-26-2] $\Delta_{\text{sub}}H$	<i>tris</i> (1,1,1,2,2,3,3-heptafluoro-7,7-dimethyl-4,6-octanedionato)aluminum(III) (363–398) 71.1 ± 2.5 381 [1972FON/POM]				
$C_{32}H_{16}AlClN_8$	[14154-42-8] $\Delta_{\text{sub}}H$	aluminum(III)-(phthalocyaninato)chloro complex (588–703) 236.4 ± 1.7 ME [2000SEM/BAS]				
$C_{32}H_{16}AlFN_8$	[51961-93-4] $\Delta_{\text{sub}}H$	aluminum(III)-(phthalocyaninato)fluoro complex (658–768) 266.9 ± 2.5 ME [2000SEM/BAS]				
$C_{33}H_{57}AlO_6$	[14319-08-5] $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$	<i>tris</i> (2,2,6,6-tetramethyl-3,5-heptanedionato)aluminum(III) (413–443) 88 TGA [2000FAH/BAR] 119 ± 3.0 [1977NAG, 1983RIB/REI]				
AlB_3H_{12}	[16962-07-5] Δ_vH	aluminum borohydride (231–290) 30.0 260 [1940SCH/SAN]				
Am (americium)						
$(C_{15}H_3AmF_{18}O_6)-2(C_{12}H_{27}O_4P)$	[58760-64-8] $\Delta_{\text{sub}}H$	<i>tris</i> (1,1,1,5,5,5-hexafluoro-2,4-pentanedionato)americium(III)- 2(tributylphosphate) complex (425–511) 133.9 ± 1.7 468 TRM [1978DAV/TRA]				
$(C_{15}H_{12}AmF_9O_6)-2(C_{12}H_{27}O_4P)$	[75101-27-8] $\Delta_{\text{sub}}H$	<i>tris</i> (1,1,1-trifluoro-2,4-pentanedionato)americium(III)- 2(tributylphosphate) complex (509–545) 222.6 ± 29.2 527 TRM [1978DAV/TRA]				
$C_{24}H_{30}AmF_9O_6)-2(C_{12}H_{27}O_4P)$	[75101-26-7] $\Delta_{\text{sub}}H$	<i>tris</i> (1,1,1-trimethyl-5,5,5-trifluoro-2,4-pentanedionato)americium(III)- 2(tributylphosphate) complex (438–493) 129.7 ± 23.4 465 TRM [1978DAV/TRA]				
$C_{33}H_{57}AmO_6$	[71817-66-8] $\Delta_{\text{sub}}H$	<i>tris</i> (2,2,6,6-tetramethyl-3,5-heptanedionato)americium(III) (373–423) 200.8 ME [1979AMA/SAT]				
As (arsenic)						
$CaS_2Cl_2F_3S$	[762-86-7] Δ_vH	dichloro(trifluoromethylthio)arsine (293–373) 37.1 333 [1960EME/PUG]				
CH_3AsBr_2	[676-70-0] Δ_vH	methyl dibromoarsine (293–333) 49.9 313 [1948RED/CHA2]				
CH_3AsCl_2	[593-89-5] Δ_vH	methyl dichloroarsine (273–313) 41.0 293 [1948RED/CHA2]				
CH_3AsF_2	[420-24-6] Δ_vH	methyl difluoroarsine (244–350) 35.5 297 MM [1946LON/EME]				
$C_2AsClF_6S_2$	[819-39-6] Δ_vH	chloro <i>bis</i> (trifluoromethylthio)arsine (293–373) 39.6 333 [1960EME/PUG]				
$C_2H_2AsCl_3$	[na] Δ_vH	β -chlorovinyl dichloroarsine (339–383) U88.3 354 [1947GOU/HOL]				
$C_2H_2AsCl_3$	[34461-56-8] Δ_vH	<i>cis</i> -2-chlorovinyl dichloroarsine (341–382) 48.5 356 [1948WHI]				
$C_2H_2AsCl_3$	[50361-05-2] Δ_vH	<i>trans</i> -2-chlorovinyl dichloroarsine (323–423) 54.5 338 [1950MAT/SUM]				

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound			Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)		
C ₂ H ₂ AsCl ₃	[na] $\Delta_{\text{fus}}H$	<i>trans</i> - β -(chlorovinyl)dichloroarsine				[1996LEB/KUL]
C ₂ H ₂ AsCl ₃	[541-25-3] Δ_vH	2-chlorovinyl-dichloroarsine (293–333)	53.4	313		[1948RED/CHA2]
C ₂ H ₂ AsF ₆ N	[1648-73-3] Δ_vH	(amino) <i>bis</i> (trifluoromethyl)arsine (313–358)	31.8	335		[1959CUL/EME]
C ₂ H ₅ AsCl ₂	[598-14-1] Δ_vH	ethyl dichloroarsine (293–333)	44.6	313		[1948RED/CHA2]
C ₂ H ₅ AsF ₂	[430-40-0] Δ_vH	ethyl difluoroarsine (248–367)	33.9	307	MM	[1946LON/EME]
C ₂ H ₇ AsO ₂	[75-60-5] $\Delta_{\text{fus}}H$	hydroxydimethyl arsine oxide	24.46	470.8	DSC	[1990DON/DRE]
C ₃ AsF ₉ S	[75-60-5] Δ_vH	<i>bis</i> (trifluoromethyl) trifluoromethylthioarsine (263–312)	34	287		[1962EME/PAC]
C ₃ AsF ₉ Se	[816-45-5] Δ_vH	<i>bis</i> (trifluoromethyl) trifluoromethylselenoarsine (227–295)	34.8	261		[1962EME/PAC]
C ₃ H ₄ AsF ₆ N	[684-21-9] Δ_vH	(methylamino) <i>bis</i> (trifluoromethyl)arsine (293–355)	34.9	324		[1959CUL/EME]
C ₃ H ₇ AsCl ₂	[926-53-4] Δ_vH	propyl dichloroarsine (293–333)	49.2	313		[1948RED/CHA2]
C ₃ H ₉ As	[593-88-4] $\Delta_{\text{fus}}H$	trimethyl arsine	8.96	186.6		[1988NIS/SHE]
	Δ_vH	(240–280)	27.7 ± 0.2	260		[2001BAE]
	Δ_vH		28.9 ± 1.3			[1956LON/SAC, 1982PIL/SKI]
C ₃ H ₉ AsO ₃	[6596-95-8] Δ_vH	trimethyl arsenite (300–335)	42.3 ± 1.3	298		[1953CHA/MOR, 1970COX/PIL]
C ₄ As ₄ F ₁₂	[7547-15-1] $\Delta_{\text{sub}}H$	<i>tetrakis</i> (trifluoromethyl)tetraarsene (317–354)	76.6	335		[1966COW/BUR]
C ₄ HAs ₂ F ₁₂ N	[3892-55-5] Δ_vH	iminobis[<i>bis</i> (trifluoromethyl)arsine] (357–398)	38.9	377		[1959CUL/EME]
C ₄ H ₆ AsF ₆ N	[1537-49-1] Δ_vH	(ethylamino) <i>bis</i> (trifluoromethyl)arsine (292–368)	32.8	330		[1959CUL/EME]
C ₄ H ₆ AsF ₆ N	[1537-48-0] Δ_vH	(dimethylamino) <i>bis</i> (trifluoromethyl)arsine (296–358)	35.6	327		[1959CUL/EME]
C ₄ H ₁₁ As	[692-42-2] Δ_vH	diethyl arsine (281–366)	35.2	273	MM	[2001BAE2]
	Δ_vH	(281–366)	34.2	298	MM	[2001BAE2]
C ₄ H ₁₁ AsO ₂	[4964-27-6] $\Delta_{\text{fus}}H$	diethylarsinic acid	19.9	411	DTA	[1970SMI/IRG]
C ₄ H ₁₂ AsN	[30880-19-4] Δ_vH	(dimethylamino) dimethylarsine (274–342)	36.7			[59M6D]
C ₅ AsF ₁₃ Se	[679-01-6] Δ_vH	heptafluoropropylseleno <i>bis</i> (trifluoromethyl)arsine (277–348)	40.3	312		[1962EME/PAC]

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound		T_m (K)	Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)			
C ₅ H ₇ AsCl ₂	[170135-56-5] $\Delta_v H$	bis(2-chlorovinyl)methylarsine (293–333)	55.6	313		[1948RED/CHA2]
C ₅ H ₁₁ AsBr ₂	[64047-02-5] $\Delta_v H$	pentyl dibromoarsine (293–333)	60	313		[1948RED/CHA2]
C ₅ H ₁₅ AsN ₂	[41813-33-6] $\Delta_v H$	bis(dimethylamino) methylarsine (273–333)	39.2			[1959M6D]
C ₆ H ₅ AsCl ₂	[696-28-6] $\Delta_v H$ $\Delta_v H$	phenyl dichloroarsine (313–333) (335–529)	58.4 48.7	323 350		[1948RED/CHA2] [1947STU]
C ₆ H ₉ As	[13652-20-5] $\Delta_v H$	trivinylarsine (295–339)	35.6	310		[1957MAI/SEY, 1984BOU/FRI]
C ₆ H ₁₂ AsN	[64049-16-7] $\Delta_v H$	cyano(ethyl)propylarsine (293–313)	54.6	303		[1948RED/CHA2]
C ₆ H ₁₅ As	[617-75-4] $\Delta_{\text{fus}} H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$	triethylarsine (273–339) (290–379)	11.06 38.1 ± 1.5 38.5 ± 0.7 43.1 ± 4.2	181.8 306 334		[1996DOM/HEA] [2001BAE] [2001BAE] [1963LAU/TRO, 1982PIL/SKI]
C ₆ H ₁₅ AsO ₂	[2870-87-3] $\Delta_{\text{fus}} H$	dipropylarsinic acid	22.1	408	DTA	[1970SMI/IRG]
C ₆ H ₁₅ AsO ₃	[3141-12-6] $\Delta_v H$ $\Delta_v H$	arsenic (III) triethoxide (305–340)	47.9 ± 1.1 50.6 ± 4.2	298	DSC	[1996DES/BRA] [1953CHA/MOR, 1970COX/PIL]
C ₆ H ₁₈ AsN ₃	[6596-96-9] $\Delta_{\text{fus}} H$ $\Delta_v H$	tris(dimethylamino)arsine (288–359)	13.31 45.8	222.6		[2002SHE/KAR] [1959M6D]
C ₈ H ₁₂ AsNO ₃	[na] $\Delta_v H$	dimethyl arsanilate (288–433)	48.8	303		[1947STU]
C ₈ H ₁₈ AsO ₂	[2850-61-5] $\Delta_{\text{fus}} H$	dibutylarsinic acid	29.5	412	DTA	[1970SMI/IRG]
C ₉ H ₂₁ As	[57538-64-4] $\Delta_v H$	triisopropylarsine (346–405)	45.2 ± 0.5	376		[2001BAE]
C ₉ H ₂₁ As	[5852-57-3] $\Delta_{\text{fus}} H$ $\Delta_v H$	tripropylarsine (314–420)	14.6 44.0 ± 0.7	180 367		[2002SHE/KAR2] [1995BAE/MIK, 2001BAE]
C ₉ H ₂₁ AsO ₃	[15606-91-4] $\Delta_v H$	arsenic (III) tripropoxide	51.2 ± 1.8		DSC	[1996DES/BRA]
C ₉ H ₂₁ AsO ₃	[39936-83-9] $\Delta_v H$	arsenic (III) triisopropoxide	80.1 ± 0.9		DSC	[1996DES/BRA]
C ₁₀ H ₁₆ AsNO ₃	[na] $\Delta_v H$	diethyl arsanilate (311–454)	54.2	326	A	[1987STE/MAL, 1947STU]
C ₁₀ H ₂₃ AsO ₂	[4964-30-1] $\Delta_{\text{fus}} H$	dipentyl arsenic acid	36	405	DTA	[1970SMI/IRG]

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound		T_m (K)	Method	Reference	
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)				
C ₁₂ H ₂₇ AsO ₂	[6727-92-0]	dihexyl arsenic acid					
			16.4	393			
			24.35	405	DTA	[1970SMI/IRG]	
C ₁₂ H ₂₇ AsO ₃	[3141-10-4]	arsenic (III) tributoxide					
			64.0 ± 1.8		DSC	[1996DES/BRA]	
C ₁₂ H ₂₇ AsO ₃	[51587-28-1]	arsenic (III) triisobutoxide					
			75.7 ± 1.2		DSC	[1996DES/BRA]	
C ₁₃ H ₁₀ AsN	[23525-22-6]	diphenylarsine carbonitrile (296–326)		84.6	311	A	[1987STE/MAL]
C ₁₄ H ₃₁ AsO ₂	[6757-53-5]	diheptyl arsenic acid					
			30.1	299			
			20.3	389	DTA	[1970SMI/IRG]	
C ₁₅ H ₃₀ AsN ₃ S ₆	[17767-20-3]	<i>tris</i> (N,N-diethylthiocarbamate)arsenic(III)					
			124 ± 3	298		[1987AIR/DES]	
C ₁₅ H ₃₃ As	[5852-59-5]	tripentylarsine (408–466)		62.3	432		[1932JON/DYK]
C ₁₈ H ₁₅ As	[603-32-7]	triphenylarsine					
			98.3 ± 8.4			[1982PIL/SKI, 1964MOR/SEL]	
			75.7	508	A	[1987STE/MAL, 1949FOR/BOW]	
C ₁₈ H ₁₅ AsO	[1153-05-5]	triphenylarsine oxide					
			149.0 ± 5.4			[1994LIE/MAR]	
C ₁₈ H ₃₉ AsO ₂	[6727-94-2]	dinonyl arsenic acid					
			24.3	383			
			38.1	399	DTA	[1970SMI/IRG]	
C ₁₉ H ₃₇ AsO ₇	[155325-38-5]	(R)-1,2-dicapryloxypropyl-3-arsonic acid					
			41.8	347.7	DSC	[1993SER/SOT]	
C ₁₉ H ₃₇ AsO ₇	[155325-39-6]	(S)-1,2-dicapryloxypropyl-3-arsonic acid					
			37.66	346.7	DSC	[1993SER/SOT]	
C ₂₀ H ₄₃ AsO ₂	[6727-95-3]	di- <i>n</i> -decylarsinic acid					
			24.5	380			
			42.3	400	DTA	[1970SMI/IRG]	
C ₂₁ H ₄₂ AsN ₃ S ₆	[86431-46-1]	<i>tris</i> (dipropylthiocarbamate)arsenic(III)					
			145.1 ± 5.3	298	DSC,E	[1999NEV/GOU]	
C ₂₂ H ₄₇ AsO ₂	[6727-96-4]	di- <i>n</i> -undecyl arsenic acid					
			30.1	384			
			45.1	396	DTA	[1970SMI/IRG]	
C ₂₃ H ₄₅ AsO ₇	[155325-40-9]	(R)-1,2-dicaproyloxypropyl-3-arsonic acid					
			68.2	358.3	DSC	[1993SER/SOT]	
C ₂₃ H ₄₅ AsO ₇	[155325-41-0]	(S)-1,2-dicaproyloxypropyl-3-arsonic acid					
			54.39	358.5	DSC	[1993SER/SOT]	
C ₂₄ H ₅₁ AsO ₂	[6727-97-5]	di- <i>n</i> -dodecyl arsenic acid					
			31.4	385			
			49.4	398	DTA	[1970SMI/IRG]	

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound		T_m (K)	Method	Reference	
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)				
C ₂₆ H ₅₃ AsO ₂	[6727-98-6]	di- <i>n</i> -tridecyl arsenic acid					
			$\Delta_{\text{us}}H$	36.5	388		
			$\Delta_{\text{fus}}H$	52.7	396	DTA	[1970SMI/IRG]
C ₂₇ H ₅₃ AsO ₇	[155325-42-1]	(R)-1,2-dilauryloxypropyl-3-arsonic acid					
			$\Delta_{\text{fus}}H$	76.99	364.9	DSC	[1993SER/SOT]
C ₂₇ H ₅₃ AsO ₇	[155325-43-2]	(S)-1,2-dilauryloxypropyl-3-arsonic acid					
			$\Delta_{\text{fus}}H$	65.69	363.3	DSC	[1993SER/SOT]
C ₂₇ H ₅₄ AsN ₃ S ₆	[48233-55-2]	<i>tris</i> (N,N-dibutylthiocarbamate)arsenic(III)					
			$\Delta_{\text{sub}}H$	128 ± 3	298		[1994LIE/MAR]
C ₂₇ H ₅₄ AsN ₃ S ₆	[41582-74-5]	<i>tris</i> (N,N-diisobutylthiocarbamate)arsenic(III)					
			$\Delta_{\text{sub}}H$	128 ± 2	298	DSC,E	[1997DES/DES]
C ₂₈ H ₅₉ AsO ₂	[6727-99-7]	di- <i>n</i> -tetradecyl arsenic acid					
			$\Delta_{\text{us}}H$	39.3	390		
			$\Delta_{\text{fus}}H$	58.2	397	DTA	[1970SMI/IRG]
C ₃₀ H ₆₃ AsO ₂	[6757-54-6]	di- <i>n</i> -pentadecyl arsenic acid					
			$\Delta_{\text{us}}H$	46.4	390		
			$\Delta_{\text{fus}}H$	63.6	396	DTA	[1970SMI/IRG]
C ₃₁ H ₆₁ AsO ₇	[146863-97-0]	(R)-1,2-dimyristoyloxypropyl-3-arsonic acid					
			$\Delta_{\text{fus}}H$	289.5	373.7	DSC	[1992SER/TSI]
C ₃₁ H ₆₁ AsO ₇	[146863-98-1]	(S)-1,2-dimyristoyloxypropyl-3-arsonic acid					
			$\Delta_{\text{fus}}H$	267.8	374.7	DSC	[1992SER/TSI]
C ₃₂ H ₆₇ AsO ₂	[6728-00-3]	di- <i>n</i> -hexadecyl arsenic acid					
			$\Delta_{\text{us}}H$	47.4	389		
			$\Delta_{\text{fus}}H$	66.8	395	DTA	[1970SMI/IRG]
C ₃₄ H ₇₁ AsO ₂	[6728-01-4]	di- <i>n</i> -heptadecyl arsenic acid					
			$\Delta_{\text{us}}H$	50.9	390		
			$\Delta_{\text{fus}}H$	68.6	393	DTA	[1970SMI/IRG]
C ₃₅ H ₆₉ AsO ₇	[146863-99-2]	(R)-1,2-dipalmitoyloxypropyl-3-arsonic acid					
			$\Delta_{\text{fus}}H$	250.6	377.3	DSC	[1992SER/TSI]
C ₃₅ H ₆₉ AsO ₇	[146864-00-8]	(S)-1,2-dipalmitoyloxypropyl-3-arsonic acid					
			$\Delta_{\text{fus}}H$	206.3	377.2	DSC	[1992SER/TSI]
C ₃₆ H ₇₅ AsO ₂	[6728-02-5]	di- <i>n</i> -octadecyl arsenic acid					
			$\Delta_{\text{fus}}H$	128.9	394	DTA	[1970SMI/IRG]
C ₃₈ H ₇₉ AsO ₂	[6728-03-6]	di- <i>n</i> -nonadecyl arsenic acid					
			$\Delta_{\text{fus}}H$	144	393	DTA	[1970SMI/IRG]
C ₃₉ H ₇₇ AsO ₇	[146864-01-9]	(R)-1,2-distearoyloxypropyl-3-arsonic acid					
			$\Delta_{\text{fus}}H$	274.9	382.8	DSC	[1992SER/TSI]
C ₃₉ H ₇₇ AsO ₇	[146864-02-0]	(S)-1,2-distearoyloxypropyl-3-arsonic acid					
			$\Delta_{\text{fus}}H$	180.3	382.4	DSC	[1992SER/TSI]
C ₄₀ H ₈₃ AsO ₂	[6728-04-7]	di- <i>n</i> -eicosanyl arsenic acid					
			$\Delta_{\text{us}}H$	40.0	383		
			$\Delta_{\text{fus}}H$	76.9	393	DTA	[1970SMI/IRG]
AsF ₃	[7784-35-2]	arsenic trifluoride					
			Δ_vH	35.8	293		[1941RUS/RUN]

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound		T_m (K)	Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)			
AsH ₃	[7784-42-1] $\Delta_v H$	arsine	16.7	210		[1955SHE/GIA]
AsH ₃ Si ₃	[15110-34-6] $\Delta_v H$	trisilylarsine (258–287)	41	272		[1962AMB/BOE]
As ₂ S ₂	[1303-32-8] $\Delta_v H$	arsenic (II) sulfide (663–838)	69.6	750		[1968KUA/UST]
Au (gold)						
C ₇ H ₁₀ AuF ₃ O ₂	[63470-53-1] $\Delta_{\text{sub}} H$	dimethyl(1,1,1-trifluoro-2,4-pentanedionato)gold(III) (265–300)	83.5			[2000OHT/CIC]
C ₈ H ₁₈ Au ₂ F ₆ O ₄	[1095578-82-7] $\Delta_{\text{sub}} H$	tetramethyl bis[μ -2,2,2-trifluoroacetato]digold (296–325)	103.6 ± 2.9		ME	[2008BES/MOR]
C ₈ H ₁₈ Au ₂ O ₄	[1067677-78-4] $\Delta_{\text{sub}} H$	bis[(μ -acetato)dimethylgold] (291–332)	100.9 ± 0.8	312	ME	[2007BES/BAI, 2008BES/MOR, 2007BES/MOR]
C ₁₁ H ₁₂ AuNO	[21158-63-4] $\Delta_{\text{sub}} H$	dimethyl(8-quinolinolato)gold (353–388)	121.2 ± 1.9		ME	[2008BES/MOR2]
C ₁₁ H ₁₂ AuNS	[1135482-91-5] $\Delta_{\text{sub}} H$	dimethyl(8-mercaptoquino)gold (359–418)	120.5 ± 1.7		ME	[2008MOR/ZHE]
C ₁₄ H ₃₀ Au ₂ O ₄	[1067677-79-5] $\Delta_{\text{sub}} H$	bis[μ -(2,2-dimethylpropanoato)]tetramethyldigold (295–323)	109.1 ± 2.1		ME	[2008BES/MOR, 2007BES/MOR]
C ₁₈ H ₂₂ Au ₂ O ₄	[1095578-84-9] $\Delta_{\text{sub}} H$	bis[μ -(benzoate)]tetramethyldigold (363–403)	154.5 ± 1.5		ME	[2008BES/MOR]
C ₂₀ H ₃₄ AuO ₉ PS	[34031-32-8] $\Delta_{\text{fus}} H$	5-triethylphosphine gold-2,3,4,6-tetra-O-acetyl-1-thio- β -(<i>d</i>)-glucopyranoside (auranofin)	37.82	385		[1985LIN/RAT]
B (boron)						
CH ₃ BO	[13205-44-2] $\Delta_v H$	borine carbonyl (134–209)	19.7	194		[1947STU]
(CH ₃ N)–(BH ₃)	[1722-33-4] $\Delta_{\text{sub}} H$	methylamine- borane complex (273–318)	78.7 ± 4.2		ME	[1959ALT/BRO]
CH ₅ BO ₂	[13061-96-6] $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$	dihydroxymethylborane (293–362) (298–338)	64.1 65.2	308 318	A	[1987STE/MAL] [1940BUR]
(CH ₃ N)–(C ₃ H ₉ BO ₃)	[na] $\Delta_{\text{sub}} H$	methylamine- trimethylborate complex	58.2			[1951GOU/LIN]
CH ₁₁ B ₅	[19495-55-7] $\Delta_v H$	1-methylpentaborane (9) (241–349)	32.7	295		[1963RYS/HAR]
C ₂ BF ₅	[32038-87-2] $\Delta_v H$	perfluorovinyl difluoroborane (177–238)	26.6	207	T	[1960STA/STO]
C ₂ BCl ₂ F ₃	[758-99-6] $\Delta_v H$	perfluorovinyl dichloroborane (238–301)	31.5	269	T	[1960STA/STO]
C ₂ H ₃ BF ₂	[358-95-2] $\Delta_v H$	vinyl difluoroborane (178–228)	22.6	203	T	[1960BRI/STO]
C ₂ H ₃ BCl ₂	[3677-80-3]	vinyl dichloroborane				

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)			
	$\Delta_v H$	(237–282)	27.7	260		T	[1960BRI/STO]
(C ₂ H ₃ OF)–(BF ₃)	[353-44-6]	methylfluorocarbonyl- trifluoroboron complex					
	$\Delta_{\text{sub}} H$	(223–273)	26.3	248			[1957SUS/WUH]
C ₂ H ₅ BCl ₂ O	[16339-28-9]	ethoxydichloroborane					
	$\Delta_v H$		35.1 ± 0.8	298			[1931WIB/SUT, 1970COX/PIL]
(C ₂ H ₅ B ₃)–(C ₃ H ₉ N)	[na]	1,5-dicarbopentaborane(5)- trimethylamine complex					
	$\Delta_{\text{sub}} H$	(220–253)	49.7	236			[1972BUR/REI]
C ₂ H ₆ BCl ₂ N	[1113-31-1]	dimethylaminodichloroborane					
	$\Delta_v H$		37.2 ± 1.3	298			[1951BUR/RAN, 1970COX/PIL]
C ₂ H ₆ BF ₂ N	[na]	dimethylaminodifluoroborane					
	$\Delta_{\text{sub}} H$	(308–353)	76.5	333			[1954BUR/BAN]
C ₂ H ₆ B ₄	[na]	carborane-4					
	$\Delta_v H$	(241–287)	26.2	272			[1963SHA/KEI]
C ₂ H ₆ B ₄	[20693-67-8]	1,6-dicarbahexaborane					
	$\Delta_{\text{sub}} H$	(190–209)	31.2	198	A		[1987STE/MAL]
(C ₂ H ₆ O)–(BF ₃)	[na]	dimethyl ether—boron trifluoride complex					
	$\Delta_v H$	(311–346)	53.1	328			[1960MCL/TAM]
C ₂ H ₆ ClBO ₂	[868-81-5]	dimethoxychloroborane					
	$\Delta_v H$		34.3 ± 1.2	298			[1931WIB/SUT, 1970COX/PIL]
(C ₂ H ₆ S)–(BH ₃)	[13292-87-0]	dimethyl sulfide—borane complex					
	$\Delta_v H$	(273–314)	44.9	293			[1999DYK/SVO]
C ₂ H ₇ B ₅	[20693-69-0]	2,4-dicarbocloso-heptaborane					
	$\Delta_v H$	(273–323)	31.6	288	I		[1976SHM/SHL]
(C ₂ H ₇ N)–(BH ₃)	[74-94-2]	dimethylamine- borane complex					
	$\Delta_{\text{sub}} H$	(273–308)	77.4 ± 2.9			ME	[1969KEI/KAN]
(C ₂ H ₇ N)–(C ₃ H ₉ BO ₃)	[na]	dimethylamine—methylborate complex					
	$\Delta_{\text{sub}} H$		70.3				[1951GOU/LIN]
C ₂ H ₈ BSb	[60646-39-1]	dimethylstibinoborane					
	$\Delta_v H$	(234–273)	32.1	254			[1959BUR/GRA]
C ₂ H ₁₀ BP	[4268-35-3]	dimethylphosphine borane					
	$\Delta_v H$	(303–383)	45.1	318			[1953HER/MAR]
C ₂ H ₁₁ B ₂ N	[23273-02-1]	N-dimethylaminodiborane					
	$\Delta_{\text{fus}} H$		1.41	218			[1955FUR/MCC]
	$\Delta_v H$	(220–267)	29.3	252			[1955FUR/MCC]
C ₂ H ₁₂ B ₁₀	[16872-09-6]	1,2-dicarbododecaborane (<i>o</i> -carborane)					
	$\Delta_{\text{trs}} H$	(5–310)	0.6	160			
	$\Delta_{\text{trs}} H$	(5–310)	3.77	275	AC		[2003YAM/HAY]
	$\Delta_{\text{sub}} H$	(283–333)	50.3	318	A		[1987STE/MAL]
	$\Delta_{\text{sub}} H$	(333–423)	49.4	348	A		[1987STE/MAL]
	$\Delta_{\text{sub}} H$		65.4 ± 1.0	298			[1982PIL/SKI, 1976MIR/PAV]
C ₂ H ₁₂ B ₁₀	[16986-24-6]	1,7-dicarbododecaborane (<i>m</i> -carborane)					
	$\Delta_{\text{trs}} H$	(5–310)	2.61	170			
	$\Delta_{\text{trs}} H$	(5–310)	4.41	284	AC		[2003YAM/HAY]

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound			Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)		
	$\Delta_{\text{sub}}H$	(283–333)	67.5	298	A	[1987STE/MAL]
	$\Delta_{\text{sub}}H$	(333–423)	63.3	348	A	[1987STE/MAL]
	$\Delta_{\text{sub}}H$		58.5 ± 1.0	298		[1982PIL/SKI, 1976MIR/PAV]
C₂H₁₂B₁₀	[20644-12-6]	1,12-dicarbododecaborane (<i>p</i> -carborane)				
	$\Delta_{\text{sub}}H$		61.3 ± 1.0	298		[1982PIL/SKI, 1976MIR/PAV]
C₂H₁₃B₅	[na]	1-ethylpentaborane (9)				
	Δ_vH	(273–383)	35.0	328		[1963RYS/HAR]
C₃BF₉S₃	[36884-78-3]	<i>tris</i> (trifluoromethylthio)borane				
	Δ_vH	(242–298)	33.9	270		[1999DYK/SVO]
C₃H₅BF₂	[819-69-2]	(allyl)difluoroborane				
	Δ_vH	(194–249)	28	221	T	[1960BRI/STO]
C₃H₇BF₂	[691-36-1]	(propyl)difluoroborane				
	Δ_vH	(195–248)	29.4	221	T	[1960BRI/STO]
(C₃H₇N)–(BH₃)	[na]	azetidine- borane complex				
	$\Delta_{\text{sub}}H$	(297–321)	67.9	309		[1956BUR/GOO]
C₃H₉BO₃	[121-43-7]	trimethylborate				
	Δ_vH	(304–340)	34.2	319		[1967CHR/SHI]
C₃H₉B	[593-90-8]	trimethylborane				
	$\Delta_{\text{fus}}H$		3.25	113.2		[1996DOM/HEA]
	Δ_vH		20.2 ± 0.1			[1961JOH/KIL, 1961SCO/MES]
	Δ_vH		23.9		BG	[1946BAM/LEV]
(C₃H₉B)–(C₂H₉NSi)	[na]	trimethylboron—silyldimethylamine complex				
	$\Delta_{\text{sub}}H$	(243–268)	51.4	255		[1954SUJ/WIT]
(C₃H₉B)–(C₇H₁₃N)	[na]	trimethylboron—azabicyclo[2.2.2]octane complex				
	$\Delta_{\text{sub}}H$	(273–388)	79.6			[1948BRO/SUJ]
C₃H₉BS	[19163-05-4]	dimethyl(methylthio)borane				
	Δ_vH	(227–304)	31.6	265		[1999DYK/SVO]
C₃H₉BS₂	[19163-08-7]	methylbis(methylthio)borane				
	Δ_vH	(300–373)	44.7	315		[1999DYK/SVO]
C₃H₉BS₃	[997-49-9]	<i>tris</i> (methylthio)borane				
	Δ_vH	(325–462)	44.9	394		[1999DYK/SVO]
	Δ_vH	(303–493)	51.6	398		[1967FIN/GAR2]
	Δ_vH	(303–493)	54.0 ± 0.8	298		[1967FIN/GAR2]
C₃H₉B₃Cl₃N₃	[703-86-6]	2,4,6-trichloro-1,3,5-trimethylborazine				
	$\Delta_{\text{sub}}H$	(363–404)	57.9	383.5	A	[1987STE/MAL, 1950BUR/KUL]
C₃H₉B₃O₃	[na]	methylboric acid anhydride				
	Δ_vH	(273–327)	37.4	288		[1940BUR]
(C₃H₉N)–(BF₃)	[420-20-2]	trimethylamine- boron trifluoride complex				
	$\Delta_{\text{sub}}H$	(373–413)	68.9	393	A	[1987STE/MAL, 1943BUR3]
(C₃H₉N)–(B₂F₄)	[3801-72-7]	trimethylamine- diboron tetrafluoride (tetramer)				
	$\Delta_{\text{sub}}H$	(366–399)	65.1	382		[1958FIN/SCH]
(C₃H₉N)–(BH₃)	[75-22-9]	trimethylamine- borane complex				
	$\Delta_{\text{sub}}H$	(273–363)	56.9 ± 0.8		ME	[1959ALT/BRO]

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{sub}}H$	(296–367)	57.0	311	A	[1987STE/MAL, 1937BUR/SCH]
(C ₃ H ₉ N)–(C ₃ H ₆ BCl ₂ N)	[na]	trimethylamine- dimethylaminoboron dichloride complex				
	$\Delta_{\text{sub}}H$	(293–342)	66.1 ± 1.7	317		[1952BRO/OST]
C ₃ H ₁₀ BN	[4023-40-9]	N-methylaminodimethylborane				
	$\Delta_{\text{sub}}H$		56.9 ± 0.8	298		[1988GOL/SIT, 1966GOO/MAN]
C ₃ H ₁₂ BN	[na]	borine trimethylamine				
	Δ_vH	(136–195)	19.9	180		[1937BUR/SCH]
C ₃ H ₁₂ B ₁₀ O ₂	[18178-04-6]	<i>o</i> -carboranecarboxylic acid				
	$\Delta_{\text{sub}}H$		97.0 ± 1.7	298		[1982PIL/SKI, 1970GAL/MAR]
C ₃ H ₁₂ B ₁₀ O ₂	[18581-81-2]	<i>m</i> -carboranecarboxylic acid				
	$\Delta_{\text{sub}}H$		97.7 ± 0.7	298		[1982PIL/SKI, 1970GAL/MAR]
C ₃ H ₁₂ B ₁₀ O ₂	[23087-98-1]	<i>p</i> -carboranecarboxylic acid				
	$\Delta_{\text{sub}}H$		96.3 ± 0.7	298		[1982PIL/SKI, 1970GAL/MAR]
C ₃ H ₁₄ B ₁₀	[16872-10-9]	methyl- <i>o</i> -carborane				
	$\Delta_{\text{sub}}H$		63.8 ± 0.6	298		[1982PIL/SKI, 1976MIR/PAV]
C ₃ H ₁₄ B ₁₀ O	[19610-34-5]	hydroxymethyl- <i>o</i> -carborane				
	$\Delta_{\text{sub}}H$		77.0 ± 1.3	298		[1982PIL/SKI, 1976MIR/PAV]
C ₃ H ₁₄ B ₁₀ O	[53257-04-8]	hydroxymethyl- <i>m</i> -carborane				
	$\Delta_{\text{sub}}H$		78.3 ± 1.3	298		[1982PIL/SKI, 1976MIR/PAV]
C ₃ H ₁₄ B ₁₀ O	[35795-98-3]	hydroxymethyl- <i>p</i> -carborane				
	$\Delta_{\text{sub}}H$		83.9 ± 1.3	298		[1982PIL/SKI, 1976MIR/PAV]
C ₃ H ₁₅ B ₅	[na]	1-isopropylpentaborane (9)				
	Δ_vH	(273–398)	37.2	335		[1963RYS/HAR]
C ₄ BClF ₆	[669-89-6]	<i>bis</i> (perfluorovinyl)chloroborine				
	Δ_vH	(280–322)	35.6	301	T	[1960STA/STO]
C ₄ H ₆ BCl	[10147-89-4]	divinylchloroborane				
	Δ_vH	(275–298)	33.0	286	T	[1960BRI/STO]
C ₄ H ₆ BF	[1537-50-4]	divinylfluoroborane				
	Δ_vH	(193–273)	25.8	233	T	[1960BRI/STO]
(C ₄ H ₁₀ O)–(BF ₃)	[na]	diethyl ether—boron trifluoride complex				
	Δ_vH	(283–353)	55.1	318		[1960MCL/TAM]
C ₄ H ₁₀ BClO ₂	[20905-32-2]	diethoxychloroborane				
	Δ_vH		38.9 ± 0.8	298		[1931WIB/SUT, 1970COX/PIL]
C ₄ H ₁₁ BO ₂	[4426-47-5]	dihydroxy- <i>n</i> -butylborane				
	$\Delta_{\text{sub}}H$	(303–340)	69.9 ± 0.8	321	BG	[1956MAT/ERI]
(C ₄ H ₁₁ N)–(C ₃ H ₉ B)	[na]	N,N-dimethylethylamine—trimethylborane complex				
	Δ_vH	(303–339)	58.2	321		[1960KAE/STO]
C ₄ H ₁₂ BClN ₂	[6562-41-0]	<i>bis</i> (dimethylamino)chloroborane				
	Δ_vH		41.8 ± 2.1	298		[1951BUR/RAN, 1970COX/PIL]
C ₄ H ₁₂ B ₂ Br ₄ N ₂	[25928-66-9]	dibromo(dimethylamino)borane dimer				
	$\Delta_{\text{sub}}H$		87.4 ± 22.2		BG	[1983SPI/KOL]
C ₄ H ₁₂ B ₂ O ₄	[7318-94-7]	tetramethoxydiboron				
	Δ_vH		44.7			[1972FIN/GAR]

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound			Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)		
	$\Delta_v H$	(273–348)	44.0	310		[1960BRO/MCC]
(C ₄ H ₁₂ GeO)–(BF ₃)	[na]	trimethylmethoxygermane—boron trifluoride complex				
	$\Delta_{\text{sub}} H$	(289–306)	59.5	297	SG	[1961GRI/ONY]
C ₄ H ₁₆ B ₁₀	[17032-21-2]	dimethyl- <i>o</i> -carborane				
	$\Delta_{\text{sub}} H$		65.3 ± 07	298		[1982PIL/SKI, 1976MIR/PAV]
C ₄ H ₁₇ B ₅	[na]	1-sec-butylpentaborane (9)				
	$\Delta_v H$	(299–428)	41.4	364		[1963RYS/HAR]
C ₄ H ₁₈ B ₄ N ₂	[na]	1,4-piperazinediyl bis(diborane(6))				
	$\Delta_{\text{sub}} H$	(318–346)	63.9	332		[1968BUR/IAC]
(C ₅ H ₅ N)–(BBr ₃)	[3022-54-6]	boron tribromide- pyridine complex				
	$\Delta_{\text{sub}} H$	(523–602)	65.8 ± 0.2		T	[1989GRI/KON]
			105.5 ± 1.1	393	C	[1989GRI/KON]
(C ₅ H ₁₀ O)–(BF ₃)	[na]	tetrahydropyran—boron trifluoride complex				
	$\Delta_v H$	(323–368)	60.9	345		[1960MCL/TAM2]
(C ₅ H ₁₁ N)–(BCl ₃)	[na]	piperidine- boron trichloride complex				
	$\Delta_{\text{sub}} H$		76.1		GS	[1960GRE/WAD]
(C ₅ H ₁₁ N)–(BH ₃)	[na]	piperidine- borane complex				
	$\Delta_{\text{sub}} H$	(342–380)	87.8	361		[1956BUR/GOO]
C ₅ H ₁₆ B ₁₀	[na]	isopropenyl- <i>o</i> -carborane				
	$\Delta_v H$	(323–473)	36.7	398		[1963FEI/BOB]
C ₅ H ₁₆ B ₁₀ O ₂	[19528-60-0]	1-acetoxymethyl- <i>o</i> -carborane				
	$\Delta_v H$		56.5	569		[1974DIT/SKO4]
C ₅ H ₁₉ B ₅	[na]	1-methyl-2-sec-butylpentaborane				
	$\Delta_v H$	(301–423)	41.0	362		[1963RYS/HAR]
C ₅ H ₂₁ B ₃ N ₂ S	[37956-18-6]	1,2,3,3,4,4,5,5,6,6-decahydro-1,3,3,5,5-pentamethyl-2 <i>H</i> -1,3,5,2,4,6-thiadiazatriborine				
	$\Delta_{\text{sub}} H$		57.7			[1972BUR]
C ₆ BF ₉	[815-70-3]	tris(perfluorovinyl)borine				
	$\Delta_v H$	(297–335)	41.1	316	T	[1960STA/STO]
C ₆ H ₅ BBr ₂	[4151-77-3]	phenylboron dibromide				
	$\Delta_v H$	(391–433)	43.9 ± 2.1	412	T	[1967FIN/GAR]
C ₆ H ₅ BCl ₂	[873-51-8]	phenylboron dichloride				
	$\Delta_v H$	(273–318)	33.7 ± 0.8	296	T	[1967FIN/GAR]
C ₆ H ₁₀ B ₂ N ₄	[16998-91-7]	pyrazabole				
	$\Delta_{\text{fus}} H$		11.83	354.3	DSC	[1993DOM/SER]
C ₆ H ₁₂ BCl ₃ O ₃	[na]	tris(2-chloroethyl) orthoborate				
	$\Delta_v H$	(390–448)	57.7	419		[1946JON/THO]
C ₆ H ₁₂ BNO ₃	[283-56-7]	2,8,9-trioxa-5-aza-1-boratricyclo[3.3.3.0 ^{1,5}]undecane				
	$\Delta_{\text{ms}} H$		4.54	466.7		
	$\Delta_{\text{fus}} H$		8.78	499.4	DSC	[1984WEI/LEF]
	$\Delta_{\text{sub}} H$		111.9 ± 0.9	418	C	[1984VOR/MIR]
C ₆ H ₁₃ BO ₂	[10173-39-4]	1-butaneboronic acid, cyclic ethylene ester				
	$\Delta_v H$		40.2	329		[1970FIN/GAR]

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₆ H ₁₅ B	[97-94-9]	triethylborane				
	$\Delta_{\text{fus}}H$		11.85	180.3		[1996DOM/HEA]
	Δ_vH		33.6	293		[1983HOU2]
			36.8 ± 0.4			[1963POP/SKI, 1982PIL/SKI]
C ₆ H ₁₅ BO ₃	[150-46-9]	triethylborate				
	Δ_vH	(302–382)	41.0	317		[1967CHR/SHI]
	Δ_vH	(302–382)	38.2	391		[1967CHR/SHI]
C ₆ H ₁₅ BS ₃	[998-26-5]	triethylthioborane				
	Δ_vH		61.5 ± 2.1			[1966FIN/GAR, 1970COX/PIL]
C ₆ H ₁₅ B ₃ O ₃	[3043-60-5]	triethylboroxin				
	Δ_vH	(347–424)	46.0	362	EB	[1990SPR/GRE]
C ₆ H ₁₆ BN	[na]	(N-ethylamino)diethylborane				
	Δ_vH		60.7 ± 0.8			[1967SMI/GOO, 1982PIL/SKI]
C ₆ H ₁₇ B ₅ Br ₂ Si ₂	[66798-29-6]	2,4-bis(bromodimethylsilyl)-2,4-dicarbo-closo-heptaborane (7)				
	Δ_vH	(388–463)	53.1	403	I	[1979GOL/SHM]
C ₆ H ₁₇ B ₅ Cl ₂ Si ₂	[28699-83-4]	2,4-bis(chlorodimethylsilyl)-2,4-dicarbo-closo-heptaborane (7)				
	Δ_vH	(359–439)	46.2	374	I	[1979GOL/SHM]
C ₆ H ₁₈ BN	[1722-26-5]	triethylaminoborane				
	Δ_vH		69.7 ± 0.8			[1967SMI/GOO, 1970COX/PIL]
C ₆ H ₁₈ BN ₃	[4254-29-6]	tris(triethylamino)borane				
	Δ_vH		46.9 ± 0.8			[1951BUR/RAN, 1970COX/PIL]
C ₆ H ₁₉ B ₅ Si ₂	[59351-11-0]	2,4-bis(dimethylsilyl)-2,4-dicarba-closo-heptaborane				
	Δ_vH	(373–453)	41.3	388	I	[1976SHM/SHL]
C ₆ H ₂₀ B ₂ N ₂	[na]	dimethylaminomethyl borane cyclic dimer				
	Δ_vH	(311–357)	57.8	314		[1966MIL/MUR]
C ₆ H ₂₀ B ₁₀	[23835-38-3]	1-butyl- <i>o</i> -carbaborane (12)				
	Δ_vH	(433–534)	77.3 ± 3.8	298	EB	[1980SHU/VAR]
	Δ_vH	(433–534)	50.6 ± 1.3	571	EB	[1980SHU/VAR]
C ₆ H ₂₀ B ₁₀	[51952-46-6]	1-isobutyl- <i>o</i> -carbaborane (12)				
	Δ_vH	(427–536)	72.8 ± 2.1	298	EB	[1980SHU/VAR]
	Δ_vH	(427–536)	49.1 ± 0.9	564	EB	[1980SHU/VAR]
C ₆ H ₂₀ B ₁₀	[70312-25-3]	1-butyl- <i>m</i> -carbaborane (12)				
	Δ_vH	(406–527)	67.7 ± 0.8	298	EB	[1980SHU/VAR]
	Δ_vH	(406–527)	46.7 ± 0.6	537	EB	[1980SHU/VAR]
C ₆ H ₂₀ B ₁₀	[na]	1-isobutyl- <i>m</i> -carbaborane (12)				
	Δ_vH	(400–488)	64.1 ± 2.8	298	EB	[1980SHU/VAR]
	Δ_vH	(400–488)	44.6 ± 1.3	532	EB	[1980SHU/VAR]
C ₇ H ₇ BCl ₂	[na]	<i>p</i> -tolylidichloroborane				
	$\Delta_{\text{fus}}H$		4.39	301		[1973FIN/GAR]
(C ₇ H ₉ N)–(BH ₃)	[na]	2,6-dimethylpyridine- borane complex				
$\Delta_{\text{sub}}H$	(358–378)	83.8	368	T	[1956BRO/DOM]	
C ₇ H ₁₄ BNO ₃	[283-62-5]	2,9,10-trioxa-5-aza-1-boratricyclo[4.3.3.0 ^{1,6}]dodecane				
	$\Delta_{\text{sub}}H$		105.2 ± 0.6	390	C	[1984VOR/MIR]

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound		T_m (K)	Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)			
$C_7H_{15}BO_2$	[30169-71-2] $\Delta_v H$	1-butaneboronic acid, cyclic trimethylene ester	43.1	348		[1970FIN/GAR]
$C_7H_{15}B_3F_3N_3$	[20453-68-3] $\Delta_v H$	1,2,3,4,5-pentamethyl-6-(trifluorovinyl)borazaine	18.4	302		[1999DYK/SVO]
$C_7H_{22}B_{10}$	[75482-33-6] $\Delta_v H$ $\Delta_v H$	1-pentyl- <i>o</i> -carbaborane (12)	84.3 ± 6.0	298	EB	[1980SHU/VAR]
			52.0 ± 1.5	571	EB	[1980SHU/VAR]
$C_7H_{22}B_{10}$	[75482-35-8] $\Delta_v H$ $\Delta_v H$	1-pentyl- <i>m</i> -carbaborane (12)	74.6 ± 2.4	298	EB	[1980SHU/VAR]
			48.6 ± 0.8	555	EB	[1980SHU/VAR]
$C_8H_{12}B_2Cl_6O_5$	[na] $\Delta_{\text{fus}} H$	1,3-diethyl-1,3- <i>bis</i> (trichloroacetoxy)-1,3-diboroxane	24.22	327.2		[1995DAB/DOM]
$C_8H_{16}BNO_3$	[283-64-7] $\Delta_{\text{sub}} H$	2,10,11-trioxa-5-aza-1-boratricyclo[4.4.3.01,6]tridecane	102.2 ± 1.0	390	C	[1984VOR/MIR]
$C_8H_{16}B_2O_5$	[na] $\Delta_{\text{fus}} H$	1,3-diacetoxy-1,3-diethyl-1,3-diboroxane	21.6	377.2		[1995DAB/DOM]
$C_8H_{17}BO_2$	[31044-62-9] $\Delta_v H$	1-butaneboronic acid, cyclic tetramethylene ester	76.6	364		[1970FIN/GAR]
$C_8H_{18}BBr$	[5674-70-4] $\Delta_v H$	dibutylboron bromide	50.6	328		[1953SKI/TEE]
$C_8H_{18}BCl$	[1730-69-4] $\Delta_v H$	dibutylboron chloride	48.2	328		[1953SKI/TEE]
$C_8H_{18}BNO_3$	[283-65-8] $\Delta_{\text{sub}} H$	2,10,11-trioxa-5-aza-1-boratricyclo[4.4.4.0 ^{1,6}]tetradecane	97.9 ± 1.0	418	C	[1984VOR/MIR]
$C_8H_{18}B_{10}O_3$	[146959-04-8] $\Delta_{\text{fus}} H$ $\Delta_{\text{sub}} H$	1,2-dicarbadodecaborane(12)-1-carboperoxoic acid 1,1-dimethyl-2-propynyl ester	16	374		[1999DIB/PIS2]
			120.7 ± 7.4		ME	[1999DIB/PIS]
$C_8H_{18}B_{10}O_3$	[146959-05-9] $\Delta_{\text{fus}} H$ $\Delta_{\text{sub}} H$	1,7-dicarbadodecaborane(12)-1-carboperoxoic acid 1,1-dimethyl-2-propynyl ester	29.4	360		[1999DIB/PIS2]
			80.1 ± 6.1		ME	[1999DIB/PIS]
$C_8H_{19}BO_2$	[10394-51-1] $\Delta_v H$	1-butaneboronic acid, diethyl ester	43.3	346		[1970FIN/GAR]
$C_8H_{20}B_2O$	[7318-84-5] $\Delta_v H$	tetraethyldiboroxane	42.9	358	EB	[1990SPR/GRE]
$C_8H_{20}B_2O_4$	[1630-81-5] $\Delta_v H$	tetraethoxydiboron	52.9	315		[1960BRO/MCC]
$C_8H_{24}B_2N_4$	[1630-79-1] $\Delta_v H$	tetra(dimethylamino)diboron	52.7	352		[1960BRO/MCC2]
$C_8H_{24}B_{10}$	[20740-05-0] $\Delta_v H$ $\Delta_v H$ $\Delta_v H$	1-hexyl- <i>o</i> -carbaborane (12)	86.2 ± 1.4	298		[1982PIL/SKI, 1978GAL/PAV]
			93.5 ± 6.0	298	EB	[1980SHU/VAR]
			54.1 ± 2.1	601	EB	[1980SHU/VAR]
$C_8H_{24}B_{10}$	[75482-36-9]	1-hexyl- <i>m</i> -carbaborane (12)				

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(434–544)	79.8 ± 2.4	298	EB	[1980SHU/VAR]
	$\Delta_v H$	(434–544)	50.7 ± 1.0	572	EB	[1980SHU/VAR]
C₈H₂₃B₅Si₂	[59351-10-9]	2,4-bis(trimethylsilyl)-2,4-dicarba-closo-heptaborane				
	$\Delta_v H$	(373–473)	45.0	388	I	[1976SHM/SHL]
C₉H₁₁BO₂	[4406-77-3]	benzeneboronic acid, cyclic trimethylene ester				
	$\Delta_v H$		47.3	426		[1970FIN/GAR]
C₉H₁₅BCl₆O₃	[na]	<i>tris</i> (2,2'-dichloroisopropyl) orthoborate				
	$\Delta_v H$	(488–513)	77.0	465		[1946JON/THO]
C₉H₂₁B	[1116-61-6]	tripropylborane				
	$\Delta_v H$		41.8 ± 1.3			[1963GAL/VAR, 1982PIL/SKI]
	$\Delta_v H$	(273–393)	40.0		BG	[1946BAM/LEV]
C₉H₂₁B	[1776-66-5]	triisopropylborane				
	$\Delta_v H$		41.8 ± 1.3			[1963GAL/VAR, 1982PIL/SKI]
	$\Delta_v H$	(273–393)	40.0		BG	[1946BAM/LEV]
C₉H₂₁BO₃	[688-71-1]	tripropylborate				
	$\Delta_v H$	(340–453)	52.3	355		[1980THO/SMI]
	$\Delta_v H$	(358–452)	47.6	452		[1967CHR/SHI]
C₉H₂₁BO₃	[5419-55-6]	triisopropylborate				
	$\Delta_v H$	(338–412)	42.4	412		[1967CHR/SHI]
C₉H₂₁BS₃	[998-38-9]	tri(propylthio)borane				
	$\Delta_v H$	(423–483)	76.2	453		[1967FIN/GAR2]
	$\Delta_v H$	(423–483)	87.0 ± 2.1	298		[1967FIN/GAR2]
C₉H₂₂BNO	[na]	butyl(dimethylamino)methoxyborane				
	$\Delta_v H$	(369–427)	48.1	384	EB	[1973GAL/BRY]
	$\Delta_v H$		58.2 ± 2.5	298		[1973GAL/BRY]
C₁₀H₁₃BO₂	[4406-76-2]	benzeneboronic acid, cyclic tetramethylene ester				
	$\Delta_v H$		57.3	441		[1970FIN/GAR]
C₁₀H₁₅BO₂	[31044-59-4]	benzeneboronic acid, diethyl ester				
	$\Delta_v H$		67.4	332		[1970FIN/GAR]
C₁₁H₂₄B₁₀O₃	[na]	3-methyl-3-(7-isopropyl- <i>m</i> -carboranoylperoxy)-1-butyne				
	$\Delta_v H$	(353–368)	140.6 ± 4.4	360	ME	[1999DIB/PIS2]
C₁₁H₂₄B₁₀O₃	[146959-06-0]	1,2-dicarbadodecaborane(12)-1-carboperoxoic acid 2-(1-methylethyl)-1,1-dimethyl-2-propynyl ester				
	$\Delta_{\text{sub}} H$	(345–362)	125.1 ± 7.0		ME	[1999DIB/PIS]
C₁₁H₂₄B₁₀O₃	[na]	1,7-dicarbadodecaborane(12)-1-carboperoxoic acid, 7-(1-methylethyl)-1,1-dimethyl-2-propynyl ester				
	$\Delta_{\text{fus}} H$		32.4	368		[1999DIB/PIS2]
C₁₂H₁₀BBr	[5123-17-1]	diphenylboron bromide				
	$\Delta_v H$	(436–516)	60.2 ± 2.5	476	T	[1967FIN/GAR]
C₁₂H₁₀BCl	[3677-81-4]	diphenylboron chloride				
	$\Delta_v H$	(363–485)	41.4 ± 2.1	424	T	[1967FIN/GAR]
C₁₂H₂₁B	[16664-33-8]	dodecahydro-9-boraphenalene				
	$\Delta_v H$	(304–404)	53.1	319	A	[1987STE/MAL]
C₁₂H₂₇B	[122-56-5]	tributylboron				
	$\Delta_v H$	(293–363)	54.7	328		[1953SKI/TEE]

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound			Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)		
C ₁₂ H ₂₇ BO ₃	[688-74-4]	tributylborate				
	$\Delta_v H$	(380–504)	58.1	395		[1980THO/SMI]
	$\Delta_v H$	(390–491)	55.9	405		[1967CHR/SHI]
C ₁₂ H ₂₇ BO ₃	[13195-76-1]	triisobutylborate				
	$\Delta_v H$	(372–472)	51.7	483		[1967CHR/SHI]
C ₁₂ H ₂₇ BS ₃	[998-46-9]	tri(butylthio)borane				
	$\Delta_v H$	(440–503)	83.9	471		[1967FIN/GAR2]
	$\Delta_v H$	(440–503)	95.8 ± 2.1	298		[1967FIN/GAR2]
C ₁₂ H ₃₀ B ₈	[223268-31-3]	1,10-dipentyl-1,10-dicarbadeborane				
	$\Delta_{\text{fus}} H$		14.7	269.7		[1999DOU/BOT]
C ₁₄ H ₂₆ B ₂ N ₄	[14695-69-3]	4,4,8,8-tetraethylpyrazabole				
	$\Delta_{\text{fus}} H$		28.61	342.4		
	$\Delta_{\text{fus}} H$		3.22	379.2	DSC	[1993DOM/SER]
C ₁₅ H ₃₂ B ₁₀ O ₅	[141695-58-1]	2,5-dimethyl-(2- <i>tert</i> -butylperoxy-5- <i>m</i> -carboranoylperoxy)-3-hexyne				
	$\Delta_v H$	(353–366)	86.8 ± 5.4	360	ME	[1999DIB/PIS2]
C ₁₅ H ₃₃ BO ₃	[621-78-3]	tripentylborate				
	$\Delta_v H$	(410–505)	67.7	425		[1980THO/SMI]
C ₁₅ H ₃₃ BS ₃	[1116-74-1]	tri(pentylthio)borane				
	$\Delta_v H$	(446–503)	92.3	474		[1967FIN/GAR2]
	$\Delta_v H$	(446–503)	104.6 ± 2.1	298		[1967FIN/GAR2]
C ₁₈ H ₁₂ BCl ₃ O ₃	[7539-58-2]	<i>tris</i> (4-chlorophenoxy)borane				
	$\Delta_v H$	(428–476)	30.6 ± 0.9	452	MM	[1973WIL/FEN]
C ₁₈ H ₁₂ BCl ₃ O ₃	[42080-72-8]	<i>tris</i> (3-chlorophenoxy)borane				
	$\Delta_v H$	(476–524)	49.6 ± 1.6	500	MM	[1973WIL/FEN]
C ₁₈ H ₁₅ B	[960-71-4]	triphenylborane				
	$\Delta_{\text{sub}} H$		103.8 ± 2.5	360	TE,ME	[1984GOV/KAN2]
	$\Delta_{\text{sub}} H$		92.1 ± 2.5	298		[1978STE3]
	$\Delta_{\text{sub}} H$		81.6 ± 2.1			[1982PIL/SKI, 1967FIN/GAR]
	$\Delta_v H$	(423–568)	64.3	438	A	[1987STE/MAL]
C ₁₈ H ₃₃ B	[1088-01-3]	tricyclohexylboron				
	$\Delta_{\text{sub}} H$		81.6 ± 4.2	298		[1982PIL/SKI, 1967FIN/GAR]
C ₁₈ H ₃₄ B ₂ N ₄	[77189-78-7]	4,4,8,8-tetrapropylpyrazabole				
	$\Delta_{\text{fus}} H$		33.0	382.2	DSC	[1993DOM/SER]
C ₂₁ H ₁₂ BN ₃ O ₃	[42080-77-3]	<i>tris</i> (4-cyanophenoxy)borane				
	$\Delta_v H$	(448–506)	46.2 ± 2.0	477	MM	[1973WIL/FEN]
C ₂₁ H ₂₁ BO ₃	[14643-62-0]	<i>tris</i> (4-methylphenoxy)borane				
	$\Delta_v H$	(475–525)	76.1 ± 1.7	500	MM	[1973WIL/FEN]
C ₂₁ H ₂₁ BO ₃	[14750-98-2]	<i>tris</i> (3-methylphenoxy)borane				
	$\Delta_v H$	(477–523)	77.1 ± 2.2	500	MM	[1973WIL/FEN]
C ₂₁ H ₂₁ BO ₆	[42080-76-2]	<i>tris</i> (3-methoxyphenoxy)borane				
	$\Delta_v H$	(440–496)	57.8 ± 2.4	468	MM	[1973WIL/FEN]
C ₂₁ H ₂₁ BO ₆	[42080-75-1]	<i>tris</i> (4-methoxyphenoxy)borane				
	$\Delta_v H$	(448–500)	42.4 ± 2.7	474	MM	[1973WIL/FEN]

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)			
C₂₃H₂₄BNO₂	[345342-83-8] $\Delta_{\text{fus}}H$	4-benzyl-5,6-dimethyl-2,5-diphenyl-1,3-dioxo-4-aza-2-boracyclohexane	13	408.2		[2001KLI/LUB]	
C₂₉H₂₆BCl₂NO₂	[345342-96-3] $\Delta_{\text{fus}}H$	4-benzhydryl-2,5-di(4'-chlorophenyl)-4,5-dimethyl-1,3-dioxo-4-aza-2-boracyclohexane	23.96	453.2		[2001KLI/LUB]	
		Note: Sample may have experienced partial decomposition as authors report a mass decrease at melting.					
C₂₉H₂₈BNO₂	[345342-93-0] $\Delta_{\text{fus}}H$	4-benzhydryl-5,6-dimethyl-2,5-diphenyl-1,3-dioxo-4-aza-2-boracyclohexane	28.43	431.2		[2001KLI/LUB]	
C₃₁H₃₂BNO₂	[345342-82-7] $\Delta_{\text{fus}}H$	4-benzhydryl-5,6-dimethyl-2,5-di(4'-methylphenyl)-1,3-dioxo-4-aza-2-boracyclohexane	32.48	453.2		[2001KLI/LUB]	
C₃₃H₃₀BNO₂	[345342-99-6] $\Delta_{\text{fus}}H$	4-benzhydryl-5,6-dimethyl-2-(1'-naphthyl)-5-phenyl-1,3-dioxo-4-aza-2-boracyclohexane	25.48	454.2		[2001KLI/LUB]	
BH₃O₃	[10043-35-3] $\Delta_{\text{sub}}H$	boric acid (326–363)	174.1 ± 4.7	345	GS	[2007PAN/ANT]	
B₂F₄	[13965-73-6] $\Delta_{\text{sub}}H$	diboron tetrafluoride (178–209.5)	35.5	193		[1958FIN/SCH]	
B₂H₆	[19287-45-7] Δ_vH Δ_vH Δ_vH Δ_vH Δ_vH	diborane (118–179)	15.3 14.2 12.6 10.5 7.3	164 180 210 240 270	C C C C C	[1961DIT/PER] [1959PAR/MAC] [1959PAR/MAC] [1959PAR/MAC] [1959PAR/MAC]	
B₂D₆	[20396-66-1] Δ_vH	perdeuterodiborane (118–179)	15.3	164		[1961DIT/PER]	
B₃Br₃H₃N₃	[13703-88-3] $\Delta_{\text{sub}}H$ Δ_vH	2,4,6-tribromoborazine (342–395) (404–415)	86.2 ± 0.4 47.0 ± 5.1	409	I I	[1966LAU/SCA] [1966LAU/SCA]	
B₃Cl₃H₃N₃	[933-18-6] $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$ Δ_vH Δ_vH	2,4,6-trichloroborazine (303–353) (313–357) (363–409) (360–386)	70.5 ± 0.4 71.1 49.6 ± 0.2 47.8	386 373	I I I I	[1966LAU/SCA] [1955BRO/LAU] [1966LAU/SCA] [1955BRO/LAU]	
B₃F₃H₃N₃	[13779-24-3] $\Delta_{\text{sub}}H$	2,4,6-trifluoroborazine (273–454)	63.1 ± 0.1		I	[1966LAU/SCA]	
B₃H₆N₃	[6569-51-3] $\Delta_{\text{fus}}H$	borazaole (13–310)	10.61	215.8	AC	[1992KUL/LEB, 1991LEB/KUL]	
B₃H₁₂N₃	[13871-09-5] $\Delta_{\text{sub}}H$	hexahydroborazine (321–349)	104.6 ± 12.6		ME	[1969LEA/LON, 1971LEA]	
(NH₃)–(B₃H₇)	[na] $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$	ammonia-triborane complex (306–328) (304–327)	71.5 ± 0.4 71.5		ME	[1959ALT/BRO] [1959WES/LEV]	
Ba (barium)							
(C₁₀H₂BaF₁₂O₄)–(C₁₂H₂₄O₆)	[143737-48-8] $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$	bis(1,1,1,5,5-hexafluoro-2,4-pentanedionato)barium(II)-18-crown-6 complex (412–468) (428–473)	104.9 ± 1.3 115 ± 2	440 450	T	[1995TOB/WAT] [1993SYO/GOL]	

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound		T_m (K)	Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)			
C₂₂H₃₈BaO₄	[155138-07-1]	<i>bis</i> (2,2,6,6-tetramethylheptan-3,5-dionato)barium(II)				
	$\Delta_{\text{sub}}H$		NA			[1994ALI/MAL]
	$\Delta_{\text{sub}}H$		90.2			[1993TOB/LAN]
C₃₄H₄₂BaCu₂F₂₄O₈	[16034-35-2]	<i>tetrakis</i> (hexafluoroisopropoxy) <i>bis</i> (2,2,6,6-tetramethylheptan-3,5-dionato)barium(II)dicopper(II)				
	$\Delta_{\text{sub}}H$	(383–448)	102.7	416		[1996LAB/HUB]
C₅₆H₈₀BaF₂₄O₁₂Y₂	[160669-81-8]	<i>tetrakis</i> (hexafluoroisopropoxy) <i>tetrakis</i> (2,2,6,6-tetramethylheptan-3,5-dionato)barium(II)diyttrium(III)				
	$\Delta_{\text{sub}}H$	(360–403)	84.8	382		[1996LAB/HUB]
Be (beryllium)						
C₂H₆Be	[506-63-8]	dimethyl beryllium				
	Δ_vH	(373–453)	88.7	388		[1952COA/GLO]
C₁₀H₂BeF₁₂O₄	[19648-82-9]	<i>bis</i> (1,1,1,5,5,5-hexafluoro-2,4-pentanedionato)beryllium(II)				
	$\Delta_{\text{sub}}H$	(289–349)	66.1	319	BG	[1987GRI/LAZ2]
C₁₀H₈BeF₆O₄	[13939-10-1]	<i>bis</i> (1,1,1-trifluoro-2,4-pentanedionato)beryllium(II)				
	$\Delta_{\text{sub}}H$	(354–383)	85.3 ± 6.3	368	BG	[1987GRI/LAZ2, 1988LAZ/GRI]
	$\Delta_{\text{sub}}H$		88.0 ± 6.5	298		[1987GRI/LAZ2]
	$\Delta_{\text{sub}}H$		U 30.5			[1960BER/TRU, 1965BER/TRU]
	Δ_vH	(387–474)	59.8 ± 0.4	431	BG	[1988LAZ/GRI]
C₁₀H₁₄BeO₄	[10210-64-7]	<i>bis</i> (2,4-pentanedionato)beryllium(II)				
	$\Delta_{\text{fus}}H$		15.7	381.2	DSC	[1983MUR/HIL]
	$\Delta_{\text{sub}}H$		95.3 ± 2.0			[1988RIB/PIL]
	$\Delta_{\text{sub}}H$		94 ± 1.0	298	ME	[1977NAG, 1988RIB/FER4]
	$\Delta_{\text{sub}}H$		82.3		BG	[1988LAZ/GRI]
	$\Delta_{\text{sub}}H$		91 ± 1.4	298	C	[1985MUR/SAK]
	$\Delta_{\text{sub}}H$		85.3 ± 3.5		DSC	[1983MUR/HIL]
	$\Delta_{\text{sub}}H$		U 35.6			[1960BER/TRU, 1965BER/TRU]
Δ_vH	(382–511)	65.7 ± 1.1	447	BG	[1988LAZ/GRI]	
C₁₂H₁₈Be₄O₁₃	[19049-40-2]	<i>hexakis</i> (aceto)-oxotetraberyllium				
	$\Delta_{\text{sub}}H$	(390–451)	115.3	420.5	A	[1987STE/MAL]
	$\Delta_{\text{sub}}H$ (<i>monoclinic</i>)		115.3			[1959SEM/GOR]
	$\Delta_{\text{sub}}H$ (<i>I</i>)	(394–422)	132.6	408		[1955MOM/SEK]
	$\Delta_{\text{sub}}H$ (<i>II</i>)	(426–446)	113.4	436		[1955MOM/SEK]
C₂₀H₁₂BeF₆O₄	[14052-07-4]	<i>bis</i> (1-phenyl-4,4,4-trifluoro-1,3-butanedionato)beryllium(II)				
	$\Delta_{\text{sub}}H$		U 35.8		I	[1960BER/TRU, 1965BER/TRU]
C₂₀H₁₈BeO₄	[14128-75-7]	<i>bis</i> (benzoylacetato)beryllium(II)				
	$\Delta_{\text{sub}}H$	(416–438)	151.6 ± 1.8	427	TE,ME	[1995RIB/MON2]
	$\Delta_{\text{sub}}H$		158.0 ± 1.8	298		[1995RIB/MON2]
	$\Delta_{\text{sub}}H$		142.3 ± 1.4	298	C	[1983RIB/REI]
C₂₂H₃₈BeO₄	[36915-22-7]	<i>bis</i> (2,2,6,6-tetramethylheptane-3,5-dionato)beryllium				
	$\Delta_{\text{sub}}H$		84.2		BG	[1988LAZ/GRI]
	Δ_vH	(383–525)	65.1	454	BG	[1988LAZ/GRI]
BeF₂	[7787-49-7]	beryllium fluoride				
	$\Delta_{\text{sub}}H$	(713–795)	236.4 ± 2.9	750	TE	[1965BLA/GRE]
	$\Delta_{\text{sub}}H$		231.8 ± 1.7	755	MS	[1965BLA/GRE]
	Δ_vH	(823–1223)	222.8	923	TE,ME,GS	[1963GRE/FOS]
	Δ_vH	(802–1021)	209.6	911		[1958SEN/STO]

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)			
	$\Delta_v H$	(821–1002)	196.6	911		[1958NOV/SEM]	
	$\Delta_v H$	(745–968)	212.9	856	GS	[1954SEN/SNY]	
Bi (bismuth)							
CH₅Bi	[60458-17-5] $\Delta_v H$	methylbismuth (190–258)	29.9	224		[1961AMB]	
C₂H₇Bi	[1438-45-4] $\Delta_v H$	trimethylbismuth (206–250)	32.7	228		[1961AMB]	
C₃H₉Bi	[593-91-9] $\Delta_v H$	trimethylbismuth (215–380)	35.8	298		[1961AMB]	
	$\Delta_v H$		36.0 ± 1.3			[1954LON/SAC, 1982PIL/SKI]	
	$\Delta_v H$		34.8		BG	[1946BAM/LEV]	
C₆H₉Bi	[65313-35-1] $\Delta_v H$	trivinylbismuth (293–346)	48.5	308		[1957MAI/SEY, 1984BOU/FRI]	
C₆H₁₅Bi	[617-77-6] $\Delta_{\text{fus}} H$	triethylbismuth	8.7	145.8		[1989NIS/RAB]	
	$\Delta_v H$		46.0 ± 4.2			[1963LAU/TRO, 1982PIL/SKI]	
	$\Delta_v H$	(301–343)	43.9	322		[1957MAI/SEY]	
C₁₅H₃₀BiN₃S₆	[20673-31-8] $\Delta_{\text{sub}} H$	<i>tris</i> (N,N-diethylthiocarbamate)bismuth(III)	213 ± 3	298		[1994LIE/MAR]	
C₁₈H₁₅Bi	[603-33-8] $\Delta_{\text{sub}} H$	triphenylbismuth	110.9 ± 8.4	298		[1982PIL/SKI, 1979STE]	
C₂₁H₄₂BiN₃S₆	[57407-97-3] $\Delta_{\text{sub}} H$	<i>tris</i> (dipropylthiocarbamate)bismuth(III)	285.2 ± 5.0		DSC,E	[1999NEV/GOU]	
C₂₇H₅₄BiN₃S₆	[34410-99-6] $\Delta_{\text{sub}} H$	<i>tris</i> (N,N-dibutylthiocarbamate)bismuth(III)	202 ± 3	298		[1994LIE/MAR]	
C₂₇H₅₄BiN₃S₆	[90285-80-6] $\Delta_{\text{sub}} H$	<i>tris</i> (N,N-diisobutylthiocarbamate)bismuth(III)	147 ± 3	298	DSC,E	[1997DES/DES]	
BiCl₃	[7787-60-2] $\Delta_{\text{sub}} H$	bismuth (III) chloride (371–468)	124.7		ME	[1966CUB, 1959DAR/YOS]	
	$\Delta_{\text{sub}} H$	(371–468)	118.8 ± 0.4	420	ME	[1959DAR/YOS]	
Br (bromine)							
BrFO₃	[25251-03-0] $\Delta_v H$	perbromyl fluoride (188–291)	25.3	250		[1972JOH/OHA]	
BrF₃	[7787-71-5] $\Delta_v H$	bromine trifluoride (311–428)	45.9	326		[1952OLI/GRI]	
BrF₅	[7789-30-2] $\Delta_v H$	bromine pentafluoride (297–314)	30.6	304		[1956ROG/SPE]	
	$\Delta_v H$	(213–297)	31.2	255		[1931RUF/MEN]	
BrN₃O₆	[66794-51-2] $\Delta_v H$	bromine(III) nitrate	81.1	271		[1961SCH/TAG]	
Br₂	[7726-95-6] $\Delta_v H$	bromine (343–383)	29.8	358		[1973BLA/IHL]	
	$\Delta_v H$	(297–389)	31.3	312		[1955FIS/BIN]	

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound		T_m (K)	Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)			
HBr	[7726-95-6] $\Delta_v H$	hydrogen bromide	17.6	206	C	[1928GIA/WIE]
Ca (calcium)						
C₂₂H₃₈CaO₄	[3618-89-0] $\Delta_{\text{sub}} H$	<i>bis</i> (2,2,6,6-tetramethylheptan-3,5-dionato)calcium(II)	72		GS	[1990YUH/KIK]
Cd (cadmium)						
C₂H₆Cd	[506-82-1] $\Delta_{\text{us}} H$ $\Delta_{\text{fus}} H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$	dimethyl cadmium	1.52 7.84 37.1 ± 0.1 38.9 37.9 ± 0.1 35.4	254.4 270.5 324 282		[1956LI] [1985SOK/BAE, 2001BAE] [1956LI] [1949CAR/HAR2, 1982PIL/SKI] [1946BAM/LEV]
C₄H₁₀Cd	[592-02-9] $\Delta_v H$ $\Delta_v H$	diethyl cadmium		324 46.0 ± 2.1		[1985SOK/BAE, 2001BAE] [1949CAR/HAR, 1982PIL/SKI]
C₄H₁₆CdCl₂N₈S₄	[28813-21-0] $\Delta_{\text{sub}} H$	<i>trans</i> -dichloro- <i>tetrakis</i> (thiourea)cadmium(II)		75 ± 20		[1970ASH]
C₆H₁₄Cd	[5905-48-6] $\Delta_v H$	dipropyl cadmium		54.2 ± 0.4 342		[1985SOK/BAE, 2001BAE]
C₈H₁₈Cd	[3431-67-2] $\Delta_v H$	dibutyl cadmium		67.7 ± 1.2 356		[1985SOK/BAE, 2001BAE]
C₁₀H₁₄CdCl₂N₆O₂	[na] $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$	[cadmium(1-methylcytosine) ₂ Cl ₂]		135.3 ± 20 493 145 ± 20 298	ME	[1984BUR/MOR] [1984BUR/MOR]
C₁₀H₁₄CdO₄	[14689-45-3] $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$	<i>bis</i> (2,4-pentanedionato)cadmium(II)		144.9 ± 22 443 154 ± 22 298	ME	[1984BUR/MOR] [1984BUR/MOR]
C₁₀H₂₀CdN₂S₄	[14239-68-0] $\Delta_{\text{sub}} H$	<i>bis</i> (diethyldithiocarbamate)cadmium(II)		133.2 451		[1987STE/MAL]
C₁₄H₂₈CdN₂S₄	[55519-99-8] $\Delta_{\text{sub}} H$	<i>bis</i> (dipropyldithiocarbamate)cadmium(II)		199 ± 1 298	DSC,E	[1992DEC/AIR]
C₁₈H₁₂CdN₂O₂	[14245-29-5] $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$	<i>bis</i> (8-hydroxyquinolinato)cadmium(II)		201.7 ± 7.5 298 144.9 ± 22 443 154 ± 22 298	ME ME	[1994RIB/MAT] [1984BUR/MOR] [1984BUR/MOR]
C₁₈H₃₆CdN₂S₄	[14566-86-0] $\Delta_{\text{sub}} H$	<i>bis</i> (dibutyldithiocarbamate)cadmium(II)		123 ± 3 298	DSC,E	[1991DES/DES]
C₁₈H₃₆CdN₂S₄	[69090-75-1] $\Delta_{\text{sub}} H$	<i>bis</i> (diisobutyldithiocarbamate)cadmium(II)		281 ± 2 298	DSC,E	[1994SOU/PIN]
C₂₀H₁₆CdN₂O₂	[15685-78-6] $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$	<i>bis</i> (8-hydroxy-2-methylquinolinato)cadmium(II)		190.9 ± 7.3 546 203.3 ± 7.3 298	ME	[1998RIB/MAT3] [1998RIB/MAT3]
C₄₄H₂₈CdN₄	[14977-07-2]	5,10,15,20-tetraphenylporphine cadmium(II)				

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound		T_m (K)	Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)			
	$\Delta_{\text{sub}}H$		222 ± 6		GS	[2000PER/GOL]
CdCl₂	[10108-64-2]	cadmium chloride (875–1026)	132.6	950		[1958BLO/WEL]
Ce (cerium)						
C₁₅H₁₅Ce	[1298-53-9]	<i>tris</i> (cyclopentadienyl)cerium (528–653)	104.6 ± 2.1			[1973BOR/KRA]
CeBr₃	[14457-87-5]	cerium(III) bromide (887–1003)	300 ± 10	298	TE	[2000VIL/BRU]
CeCl₃	[7790-86-5]	cerium(III) chloride (955–1070)	331 ± 5	298	TE	[2000VIL/BRU]
CeI₃	[7790-87-6]	cerium(III) iodide (910–1031)	295 ± 10	298	TE	[2000VIL/BRU]
Cf (californium)						
(C ₁₅ H ₃ CfF ₁₈ O ₆)-2(C ₆ H ₁₄ OS)	[123611-97-2]	<i>tris</i> (1,1,1,5,5,5-hexafluoro-2,4-pentanedionato)californium-dipropyl sulfoxide (1:2) complex (402–434)	93.6 ± 6.0		GS,TRM	[1989AIZ/FED]
(C ₁₅ H ₃ CfF ₁₈ O ₆)-2(C ₁₂ H ₂₇ OP)	[123628-36-4]	<i>tris</i> (1,1,1,5,5,5-hexafluoro-2,4-pentanedionato)californium-tributylphosphine oxide (1:2) complex (431–485)	130.6 ± 1.9		GS,TRM	[1989AIZ/FED]
(C ₁₅ H ₃ CfF ₁₈ O ₆)-2(C ₁₂ H ₂₇ O ₄ P)	[123712-43-6]	<i>tris</i> (1,1,1,5,5,5-hexafluoro-2,4-pentanedionato)californium-tributylphosphate (1:2) complex (413–451)	133.0 ± 6.1		GS,TRM	[1989AIZ/FED]
Cl (chlorine)						
ClFO₃	[7616-96-6]	perchloryl fluoride (164–228)	19.2	226	MM	[1958KOE/GIA]
	Δ_vH		19.3	226	C	[1958KOE/GIA]
ClF₃	[7790-91-2]	chlorine trifluoride (299–317)	27.5	313		[1997SAK/HOR]
	Δ_vH	(226–303)	28.4	288		[1951GRI/BER]
ClNO	[2696-92-6]	nitrosyl chloride (203–258)	25.3	230		[1949PAR/WYN]
ClNO₃	[14545-72-3]	chlorine nitrate	30.5			[1961SCH/BRA]
Cl₂O₆	[12442-63-6]	dichlorine hexaoxide (273–318)	52.3	295		[1990LOP/SIC]
HCl	[7647-01-0]	hydrogen chloride (121–133)	19.7	127		[1990SER/LAR]
	$\Delta_{\text{sub}}H$	(134–150)	19.6	142		[1990SER/LAR]
	Δ_vH		16.2	188	C	[1928GIA/WIE2]
Co (cobalt)						
C₃CoNO₄	[14096-82-3]	cobalt nitrosyl tricarbonyl (272–353)	36.3	287		[1947STU]
C₄HCoO₄	[16842-03-8]	hydridocobalt tetracarbonyl (273–295)	28.0		GS	[1980ROT/ORC]

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound		T_m (K)	Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)			
C₄H₃CoO₄Si	[14652-62-1] $\Delta_v H$	silyl tetracarbonyl cobalt (263–357)	37.8	310	T	[1969AYL/CAM]
C₄H₁₆Cl₂CoN₈S₄	[22738-43-8] $\Delta_{\text{sub}} H$	<i>trans</i> -dichloro- <i>tetrakis</i> (thiourea)cobalt(II) (356–382)	129 ± 20			[1970ASH]
C₇H₅CoO₂	[na] $\Delta_v H$	(cyclopentadienyl) cobalt dicarbonyl (313–369)	52.1 ± 0.7			[2000SZT/BAE]
C₈Co₂O₈	[10210-68-1] $\Delta_{\text{sub}} H$	octacarbonyldicobalt (264–278)	84.3 ± 0.5	271	TE	[1995GAR/CHA]
	$\Delta_{\text{sub}} H$	(288–315)	103.8	301.5	A	[1987STE/MAL, 1968BAE]
	$\Delta_{\text{sub}} H$		65.2 ± 3.3	298		[1982PIL/SKI, 1975GAR/CAR]
	$\Delta_{\text{sub}} H$	(207–287)	75.3 ± 6.3		EM	[1973CAR/ROB]
C₈H₂Co₂O₈Si	[23591-62-0] $\Delta_v H$	silylene <i>bis</i> (tetracarbonylcobalt) (297–335)	38.7	316	T	[1969AYL/CAM]
C₈H₁₀Cl₂CoN₆O₂	[74543-51-4] $\Delta_{\text{sub}} H$	[cobalt(cytosine) ₂ Cl ₂] (483–523)	151.8 ± 14	503	ME	[1984BUR/MOR]
	$\Delta_{\text{sub}} H$	(483–523)	162 ± 14	298	ME	[1984BUR/MOR]
C₉CoMnO₉	[35646-82-3] $\Delta_{\text{sub}} H$	nonacarbonylcobaltmanganese 85 ± 2		308	C	[1998ADD/CON]
	$\Delta_{\text{sub}} H$	72 ± 2		298	C	[1998ADD/CON]
C₉CoO₉Re	[15039-80-2] $\Delta_{\text{sub}} H$	nonacarbonylcobaltrhenium 94 ± 4		313	C	[1998ADD/CON]
	$\Delta_{\text{sub}} H$	83 ± 4		298	C	[1998ADD/CON]
C₁₀BrCo₃O₉	[19439-14-6] $\Delta_{\text{sub}} H$	(bromomethylidene)tricobalteneacarbonyl 99.6 ± 1.7		298		[1982PIL/SKI, 1975GAR/CAR]
C₁₀ClCo₃O₉	[13682-02-5] $\Delta_{\text{sub}} H$	(chloromethylidene)tricobalteneacarbonyl 117.6 ± 2.5		298		[1982PIL/SKI, 1975GAR/CAR]
C₁₀H₈Cl₄CoN₂	[14361-73-0] $\Delta_{\text{sub}} H$	[cobalt(2-chloropyridine) ₂ Cl ₂] (345–365)	101.2 ± 6.7	355	DSC	[1982MOR]
C₁₀H₈Cl₄CoN₂	[14361-78-5] $\Delta_{\text{sub}} H$	[cobalt(3-chloropyridine) ₂ Cl ₂] (345–365)	77.0 ± 4.2	355	DSC	[1982MOR]
C₁₀H₁₀Co	[1277-43-6] $\Delta_{\text{sub}} H$	dicyclopentadienyl cobalt 72.1 ± 0.1				[1988TOR/BAR2]
	$\Delta_{\text{sub}} H$	70.3 ± 4.2		298		[1982PIL/SKI, 1975TEL/KIR]
C₁₀H₁₄CoO₄	[14024-48-7] $\Delta_{\text{sub}} H$	<i>bis</i> (2,4-pentanedionato)cobalt(II) (433–463)	149		TGA	[2000FAH/BAR]
	$\Delta_{\text{sub}} H$	(322–371)	130.1 ± 6.3	298	ME	[1990MAL/ALI]
	$\Delta_{\text{sub}} H$		118.7 ± 2.2	298		[1985MUR/SAK]
	$\Delta_{\text{sub}} H$		81.2	370		[1970GOE/BLO]
	$\Delta_{\text{sub}} H$		U 62.8			[1960BER/TRU, 1965BER/TRU]
C₁₂Co₄O₁₂	[17786-31-1] $\Delta_{\text{sub}} H$	tetracobaltdodecacarbonyl 96.2 ± 4.2		298		[1982PIL/SKI, 1974CON/SKI]
C₁₂H₁₄Cl₂CoN₂	[13869-67-5] $\Delta_{\text{sub}} H$	cobalt(2-methylpyridine) ₂ Cl ₂ (345–365)	86.6 ± 3.8	355	DSC	[1982MOR]
C₁₄H₁₀Br₂CoN₂S₂	[21422-14-0] $\Delta_{\text{sub}} H$	[cobalt(benzothiazole) ₂ Br ₂] (381–399)	124.7 ± 4.1	390	DSC	[1973MOR/MCN]

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
$C_{15}H_3CoF_{18}O_6$	[16702-37-7] $\Delta_{\text{sub}}H$	<i>tris</i> (1,1,1,5,5,5-hexafluoro-2,4-pentanedionato)cobalt(III) (333–363)	73.0		TGA	[2000FAH/BAR]
$C_{15}H_{12}CoF_9O_6$	[16827-64-8] $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$	<i>tris</i> (1,1,1-trifluoro-2,4-pentanedionato)cobalt(III) (373–403) (383–433)	168 ± 2.0 114 ± 4.0 108.8 ± 0.4	119 407 298	TGA C GS	[2000FAH/BAR] [1988RIB/FER] [1988RIB/FER] [1985MAT/KUW]
$C_{15}H_{21}CoO_6$	[21679-46-9] $\Delta_{\text{fus}}H$	<i>tris</i> (2,4-pentanedionato)cobalt(III) Note: Value is abnormally large compared with Cr(acac) ₃ - may undergo decomposition (433–463)	93.9 138	478	DSC TGA	[2004SAB/MAR] [2000FAH/BAR] [1994GER/GER]
	$\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$	(318–382)	134.6 ± 4.0 142.6 ± 6.9 86.3 107.1 74.9 ± 4.6 U 13.0	298 471 390	ME DSC	[1990MAL/ALI] [1987MUR/HIL] [1971ASH] [1970GOE/BLO] [1964WOO/JON] [1961BER/DOW]
$C_{15}H_{30}CoN_3S_6$	[13963-60-5] $\Delta_{\text{sub}}H$	<i>tris</i> (diethyldithiocarbamato)cobalt(III) (448–587)	95 ± 6	518		[1979CAV/HIL2]
$C_{16}H_{14}Br_2CoN_2O_2$	[22974-96-5] $\Delta_{\text{sub}}H$	[cobalt(2-methylbenzoxazole) ₂ Br ₂] (345–390)	111.1 ± 4.2	368	DSC	[1982MOR, 1974MOR/MCN]
$C_{16}H_{14}Cl_2CoN_2O_2$	[52657-96-2] $\Delta_{\text{sub}}H$	[cobalt(2-methylbenzoxazole) ₂ Cl ₂] (345–390)	92.4 ± 2.5	368	DSC	[1982MOR, 1974MOR/MCN]
$C_{16}H_{14}Br_2CoN_2S_2$	[26225-02-5] $\Delta_{\text{sub}}H$	[cobalt(2-methylbenzothiazole) ₂ Br ₂] (335–354)	115.1 ± 4.1	345	DSC	[1973MOR/MCN]
$C_{16}H_{14}Cl_2CoN_2S_2$	[26225-01-4] $\Delta_{\text{sub}}H$	[cobalt(2-methylbenzothiazole) ₂ Cl ₂] (332–356)	122.6 ± 1.2	345	SC	[1973MOR/MCN]
$C_{18}H_{12}CoN_2O_2$	[13978-88-6] $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$	<i>bis</i> (8-hydroxyquinolinato)cobalt(II) (533–569)	205.3 ± 4.0 185.7 ± 9 200 ± 10	298 551 298	ME ME	[1994RIB/MAT] [1984BUR/MOR] [1984BUR/MOR]
$C_{18}H_{14}CoN_4$	[41283-94-7] $\Delta_{\text{sub}}H$	dibenzotetra-aza-annulene cobalt(II) complex (433–463)	178.2 ± 16.7	360		[1982ZVE/VIN]
$C_{18}H_{18}Br_2CoN_2O_2$	[52230-48-5] $\Delta_{\text{sub}}H$	[cobalt(2,5-dimethylbenzoxazole) ₂ Br ₂] (345–390)	95.4 ± 4.6	368	DSC	[1982MOR, 1974MOR/MCN]
$C_{18}H_{18}Cl_2CoN_2O_2$	[52230-47-4] $\Delta_{\text{sub}}H$	[cobalt(2,5-dimethylbenzoxazole) ₂ Cl ₂] (345–390)	104.6 ± 5.8	368	DSC	[1982MOR, 1974MOR/MCN]
$C_{20}H_{16}CoN_2O_2$	[17992-18-6] $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$	<i>bis</i> (8-hydroxy-2-methylquinolinate)cobalt(II) (457–473)	196.1 ± 5.9 204.4 ± 5.9	465 298	ME	[1998RIB/MAT3] [1998RIB/MAT3]
$C_{22}H_{38}CoO_4$	[13986-53-3] $\Delta_{\text{sub}}H$	<i>bis</i> (2,2,6,6-tetramethyl-3,5-heptanedionato)cobalt(II) (433–463)	143		TGA	[2000FAH/BAR]
$C_{24}H_{12}CoF_9O_6S_3$	[41875-84-7] $\Delta_{\text{sub}}H$	<i>tris</i> (1-(2-thenoyl)-4,4,4-trifluoro-1,3-butanedione)cobalt(III) (433–463)	45.6			[1961BER/DOW]

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound			Method	Reference	
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)			
$C_{24}H_{12}CoF_9O_9$	[64137-83-3] $\Delta_{\text{sub}}H$	<i>tris</i> (2-furoyltrifluoroacetato)cobalt(III)	35.6			[1961BER/DOW]	
$C_{30}H_{18}CoF_9O_6$	[31125-84-5] $\Delta_{\text{sub}}H$	<i>tris</i> (1-phenyl-4,4,4-trifluoro-1,3-butanedionato)cobalt(III)	51.0			[1961BER/DOW]	
$C_{30}H_{27}CoO_6$	[14524-55-1] $\Delta_{\text{sub}}H$	<i>tris</i> (1-phenyl-1,3-butanedionato)cobalt(III)	39.0			[1961BER/DOW]	
$C_{32}H_{16}CoN_8$	[3317-67-7] $\Delta_{\text{sub}}H$	cobalt (II) phthalocyanine	183.7 ± 13.8		ME	[1970BON/CAT]	
$C_{32}H_{46}CoN_2O_4$	[18347-53-8] $\Delta_{\text{sub}}H$	<i>bis</i> (2,2,6,6-tetramethyl-3,5-heptanedionato)(2,2'-bipyridyl)cobalt(II)	126 ± 4.0		B	[1996CHA/EMM]	
	$\Delta_{\text{sub}}H$		130.3		UV/Vis	[1996CHA/EMM]	
	$\Delta_{\text{sub}}H$		124.4		MEM	[1996CHA/EMM]	
$C_{33}H_{57}CoO_6$	[14877-41-3] $\Delta_{\text{sub}}H$	<i>tris</i> (2,2,6,6-tetramethyl-3,5-heptanedionato)cobalt(III)	(433–463)	132	TGA	[2000FAH/BAR]	
	$\Delta_{\text{sub}}H$			126 ± 3.0	298	C	[1988RIB/FER]
$CoBr_2$	[7789-43-7] $\Delta_{\text{sub}}H$	cobalt(II) bromide	(764–911)	207 ± 4.0	802	TE	[1997BAR/BRU]
	$\Delta_{\text{sub}}H$			216 ± 1.0	298		[1997BAR/BRU]
Cr (chromium)							
C_6CrO_6	[13007-92-6] $\Delta_{\text{sub}}H$	chromium hexacarbonyl	(309–347)	63.3	328	GS	[2002PAN/MAL]
	$\Delta_{\text{sub}}H$		(266–272)	65.7	269	TE	[1995GAR/CHA]
	$\Delta_{\text{sub}}H$		(323–391)	68.5 ± 1.1			[1993BAE]
	$\Delta_{\text{sub}}H$		(288–423)	68.5	355.5	A	[1987STE/MAL]
	$\Delta_{\text{sub}}H$			68.9 ± 2	298		[1984ALT/CON]
	$\Delta_{\text{sub}}H$			70.0 ± 2	298	C	[1983RIB/REI]
	$\Delta_{\text{sub}}H$		(240–280)	71.6 ± 1.7	260	ME	[1980BOX/ERN, 1979DAA/ERN]
	$\Delta_{\text{sub}}H$			69.5	298	C	[1975ADE/BRO]
	$\Delta_{\text{sub}}H$			72.0 ± 4.2	298		[1982PIL/SKI, 1975PIT/PIL]
	$\Delta_{\text{sub}}H$		(274–301)	71.5 ± 0.8	288	BG	[1966BON]
	$\Delta_{\text{sub}}H$		(319–411)	69.3			[1952REZ/SHV]
	$\Delta_{\text{sub}}H$			71.9			[1935HIE/ROM]
	$\Delta_{\text{sub}}H$		(308–408)	63.6	358	MM	[1934WIN/BLA]
Δ_vH	(309–424)	62.5	324		[1947STU]		
$C_8H_3CrNO_5S$	[55293-31-7] $\Delta_{\text{sub}}H$	thiazole(pentacarbonyl)chromium	(270–301)	102.0 ± 2.7	286	ME	[1979DAA/ERN]
$C_8H_4CrN_2O_5$	[71127-65-6] $\Delta_{\text{sub}}H$	pyrazole(pentacarbonyl)chromium	(270–303)	88.4 ± 1.8	287	ME	[1979DAA/ERN]
$C_8H_9CrNO_5$	[15228-26-9] $\Delta_{\text{sub}}H$	trimethylamine(pentacarbonyl)chromium	(248–293)	80.2 ± 0.7	271	ME	[1980BOX/ERN]
$C_8H_9CrO_5P$	[26555-09-9] $\Delta_{\text{sub}}H$	trimethylphosphine(pentacarbonyl)chromium		91.2 ± 1.6		ME	[1980BOX/ERN]
$C_8H_{12}CrMoO_8$	[71561-64-3] $\Delta_{\text{sub}}H$	chromium molybdenum tetraacetate		165.0 ± 8.4			[1982PIL/SKI]
$C_8H_{12}Cr_2O_8$	[15020-15-2] $\Delta_{\text{sub}}H$	tetra- μ -acetatodichromium(II)	(330–340)	299.6 ± 10	335	ME,TE	[1984CAR]

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound			Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)		
	$\Delta_{\text{sub}}H$		313.8 ± 27.0	298		[1982PIL/SKI, 1979DAA/ERN]
	$\Delta_{\text{sub}}H$		145		E	[1979CAV/GAR]
C ₉ H ₄ CrN ₂ O ₅	[66179-02-0]	pyrazine(pentacarbonyl)chromium				
	$\Delta_{\text{sub}}H$		99.7		ME	[1979DAA/ERN]
C ₉ H ₅ ClCrO ₃	[12082-03-0]	chlorobenzenechromium tricarbonyl				
	$\Delta_{\text{sub}}H$		102.5 ± 4.2	298		[1982PIL/SKI, 1975ADE/BRO]
C ₉ H ₆ CrO ₃	[12082-08-5]	benzene chromium tricarbonyl				
	$\Delta_{\text{sub}}H$		91.2	298	C	[1975ADE/BRO]
	$\Delta_{\text{sub}}H$		U 58.6			[1961FIS/FRI, 1973CON/SKI]
	$\Delta_{\text{sub}}H$	(364–370)	97.9		TE	[1959COR/SCH, 1973CON/SKI]
C ₁₀ H ₅ CrNO ₅	[14740-77-3]	pyridine(pentacarbonyl)chromium				
	$\Delta_{\text{sub}}H$	(294–317)	103.2 ± 1.8	306	ME	[1979DAA/ERN]
C ₁₀ H ₈ CrO ₃	[12125-72-3]	cycloheptatriene chromium tricarbonyl				
	$\Delta_{\text{sub}}H$		94.1	298	C	[1975ADE/BRO]
C ₁₀ H ₈ CrO ₃	[12125-87-0]	η^6 -toluene(tricarbonyl)chromium				
	$\Delta_{\text{sub}}H$		93.0 ± 2.0	298	C	[1984ALT/CON]
	$\Delta_{\text{sub}}H$		94.6 ± 4.2	298		[1982PIL/SKI, 1975ADE/BRO]
C ₁₀ H ₈ CrO ₃	[12116-44-8]	η^6 -anisole(tricarbonyl)chromium				
	$\Delta_{\text{sub}}H$		104.2 ± 2.0	298	C	[1984ALT/CON]
C ₁₀ H ₁₀ Cr	[1271-24-5]	chromocene				
	$\Delta_{\text{sub}}H$		71.0	298		[1984BAE/BAR2]
	$\Delta_{\text{sub}}H$		62.8 ± 4.2	298		[1982PIL/SKI, 1975TEL/KIR]
	$\Delta_{\text{sub}}H$		69.9 ± 1.7			[1977TEL/RAB]
	Δ_vH	(452–519)	49.5 ± 1.5	485		[1984BAE/BAR2]
C ₁₀ H ₁₄ CrO ₄	[14024-50-1]	bis(2,4-pentanedionato)chromium(II)				
	$\Delta_{\text{sub}}H$	(330–370)	129.8 ± 8.7	298	ME	[1990MAL/ALI]
	$\Delta_{\text{sub}}H$		111	439	T	[1981MAS/BAR]
C ₁₀ H ₁₁ CrNO ₅	[15710-39-1]	piperidine(pentacarbonyl)chromium				
	$\Delta_{\text{sub}}H$	(265–298)	93.5 ± 1.9	282	ME	[1979DAA/ERN]
C ₁₁ H ₈ CrO ₃	[na]	styrenetricarbonyl chromium				
	$\Delta_{\text{fus}}H$		25.28	354.4		[2003SMI/LEB]
C ₁₁ H ₈ CrO ₄	[12146-36-0]	norbornadienechromium tetracarbonyl				
	$\Delta_{\text{sub}}H$		89.0 ± 4.0	298		[1982PIL/SKI, 1977BRO/CON]
C ₁₁ H ₈ CrO ₄	[12153-11-6]	η^6 -acetophenone(tricarbonyl)chromium				
	$\Delta_{\text{sub}}H$		107.0 ± 0.6	298	C	[1984ALT/CON]
C ₁₁ H ₈ CrO ₅	[12125-87-0]	η^6 -methyl benzoate(tricarbonyl)chromium				
	$\Delta_{\text{sub}}H$		114.0 ± 5.0	298	C	[1984ALT/CON]
C ₁₁ H ₁₁ CrNO ₃	[12109-10-3]	η^6 -N,N-dimethylaniline(tricarbonyl)chromium				
	$\Delta_{\text{sub}}H$		118.4 ± 10	298	C	[1984ALT/CON]
C ₁₂ H ₁₀ CrO ₃	[na]	α -methylstyrenetricarbonyl chromium				
	$\Delta_{\text{fus}}H$		27.4	360		[2003SMI/LEB]
C ₁₂ H ₁₂ Cr	[1271-54-1]	dibenzenechromium				
	$\Delta_{\text{sub}}H$	(323–363)	89.4	343	A	[1987STE/MAL]
	$\Delta_{\text{sub}}H$		78.2 ± 6.3	298		[1982PIL/SKI, 1973CON/SKI]

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound			Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)		
	$\Delta_{\text{sub}}H$		82.0 ± 2.1		ME	[1973UMI/FED]
	$\Delta_{\text{sub}}H$		90.6 ± 0.3			[1969AND/WES2]
	$\Delta_{\text{sub}}H$		78.2 ± 6.2	298		[1958FIS/SCH]
C₁₂H₁₂CrO₃	[12129-67-8]	mesitylene chromium tricarbonyl				
	$\Delta_{\text{sub}}H$		108.4	298	C	[1975ADE/BRO]
	$\Delta_{\text{sub}}H$		U 64.4			[1961FIS/SCH, 1977TEL/RAB]
C₁₂H₁₂CrO₃	[32913-41-0]	(1,2,4-trimethylbenzene) chromium tricarbonyl				
	$\Delta_{\text{sub}}H$		U 33.5			[1961FIS/SCH, 1977TEL/RAB]
C₁₃H₈CrO₃	[12110-37-1]	(1,2,3,4,4a,8a-h-naphthalene)tricarbonyl chromium				
	$\Delta_{\text{sub}}H$		107 ± 3	298	C	[1979CON/MAR]
C₁₅H₃F₁₈CrO₆	[14592-80-4]	<i>tris</i> (1,1,1,5,5,5-hexafluoro-2,4-pentanedionato)chromium(III)				
	$\Delta_{\text{sub}}H$	(333–363)		46	TGA	[2000FAH/BAR]
	$\Delta_{\text{sub}}H$		164 ± 4.0	426	C	[1987RIB/FER]
	$\Delta_{\text{sub}}H$		112 ± 4.0	298		[1987RIB/FER]
	$\Delta_{\text{sub}}H$	(333–360)	123.0 ± 1.3	335		[1972FON/POM]
C₁₅H₁₂CrF₉O₆	[14592-89-2]	<i>tris</i> (1,1,1-trifluoro-2,4-pentanedionato)chromium(III)				
	$\Delta_{\text{sub}}H$	(373–403)	71		TGA	[2000FAH/BAR]
	$\Delta_{\text{sub}}H$		182 ± 4.0	426	C	[1987RIB/FER]
	$\Delta_{\text{sub}}H$		117 ± 4.0	298		[1987RIB/FER]
	$\Delta_{\text{sub}}H$	(373–438)	115.1 ± 0.8		GS	[1985MAT/KUW]
	$\Delta_{\text{sub}}H$	(403–423)	112.5 ± 4.8			[1978CHU/IGU]
	$\Delta_{\text{sub}}H$		53.6	447		[1977VOL/MAZ]
	$\Delta_{\text{sub}}H$	(377–413)	108.8 ± 1.3	395		[1972FON/POM]
	Δ_vH	(424–486)	76.7 ± 0.6	455		[1978CHU/IGU]
C₁₅H₁₈CrO₃	[12088-11-8]	hexamethylbenzene chromium tricarbonyl				
	$\Delta_{\text{sub}}H$		123.0 ± 4.0	298	C	[1975ADE/BRO, 1977BRO/CON]
C₁₅H₂₁CrO₆	[21679-31-2]	<i>tris</i> (2,4-pentanedionato)chromium(III)				
	$\Delta_{\text{fus}}H$		35.9	486	DSC	[2004SAB/MAR, 1988LAZ/GRI]
	$\Delta_{\text{fus}}H$		34	489		[1987LAZ/GRE]
	$\Delta_{\text{fus}}H$		28.7	487		[1984MUR/HIL]
	$\Delta_{\text{fus}}H$		35.2	490		[1971BEE/LIN2]
	$\Delta_{\text{fus}}H$		28.4	489		[1970MEL/MER2]
	$\Delta_{\text{sub}}H$		120.8		TGA,DTA	[2009GAI/KUN]
	$\Delta_{\text{sub}}H$	(345–410)	128.2	378	ME	[2007SID/SID]
	$\Delta_{\text{sub}}H$	(320–388)	127.6	354	ME	[2005SEM/IGU]
	$\Delta_{\text{sub}}H$	(374–418)	111.6	396	GS	[2002PAN/MAL]
	$\Delta_{\text{sub}}H$		133.8 ± 4.2			[2001FED/GEL]
	$\Delta_{\text{sub}}H$	(413–443)	91.0		TGA	[2000FAH/BAR]
	$\Delta_{\text{sub}}H$	(350–375)	126.8 ± 4.2	298	ME	[1990MAL/ALI]
	$\Delta_{\text{sub}}H$	(457–486)	113.0 ± 4.8		BG	[1988LAZ/GRI, 1987GRI/LAZ]
	$\Delta_{\text{sub}}H$		132.1 ± 1.9	298	C	[1985MUR/SAK]
	$\Delta_{\text{sub}}H$		28.9	463		[1977VOL/MAZ]
	$\Delta_{\text{sub}}H$		112.1	390		[1970GOE/BLO]
	$\Delta_{\text{sub}}H$	(363–393)	40.2 ± 1.7	378		[1972FON/POM]
	$\Delta_{\text{sub}}H$		110.9 ± 0.8	298	HSA	[1970MEL/MER, 1970MEL/MER2]
	$\Delta_{\text{sub}}H$		123 ± 3.0	298	ME	[1977NAG, 1988RIB/FER4, 1967HIL/IRV]
	Δ_vH		89.9		DTA,TGA	[2009GAI/KUN]

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound		T_m (K)	Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)			
	$\Delta_v H$		79.4 ± 4.2			[2001FED/GEL]
	$\Delta_v H$	(490–536)	82.2 ± 2.0	513	BG	[1988LAZ/GRI]
C₁₆H₂₀Cr	[na]	<i>bis</i> (ethylbenzene)chromium				
	$\Delta_v H$		75.3 ± 8.4			[1973TEL/RAB, 1982PIL/SKI]
C₁₈H₂₄Cr	[1274-07-3]	<i>bis</i> (η^6 -1,3,5-trimethylbenzene)chromium				
	$\Delta_{\text{sub}} H$		104 ± 1	298	C	[1979CON/MAR]
C₂₀H₁₆Cr	[33085-81-3]	<i>bis</i> (naphthalene)chromium				
	$\Delta_{\text{sub}} H$		105.0 ± 10			[1979CON/MAR]
C₂₀H₂₈Cr	[na]	<i>bis</i> (1,2-diethylbenzene)chromium				
	$\Delta_v H$		75.3 ± 8.4			[1973TEL/RAB, 1982PIL/SKI]
C₂₁H₃₀Cr	[na]	(1,2-diisopropylbenzene)isopropylbenzenechromium				
	$\Delta_v H$		100.4 ± 8.4			[1973TEL/RAB, 1982PIL/SKI]
C₂₃H₁₅CrO₅P	[14917-12-5]	triphenylphosphine(pentacarbonyl)chromium				
	$\Delta_{\text{sub}} H$	(324–347)	170.2 ± 6.8	336	ME	[1980BOX/ERN]
C₂₄H₂₄Cr₂N₄O₄	[67634-82-6]	<i>tetrakis</i> (6-methyl-2-hydroxypyridyl)dichromium(II)				
	$\Delta_{\text{sub}} H$		150.0 ± 4.0	298		[1982PIL/SKI, 1981CAV/GAR]
C₂₄H₃₆Cr	[na]	<i>bis</i> (1,2-diisopropylbenzene)chromium				
	$\Delta_v H$		100.4 ± 8.4			[1973TEL/RAB, 1982PIL/SKI]
C₂₄H₃₆Cr	[12156-66-0]	<i>bis</i> (η^6 -hexamethylbenzene)chromium				
	$\Delta_{\text{sub}} H$		119 ± 4	298	C	[1979CON/MAR]
C₃₀H₂₇CrO₆	[16432-36-3]	<i>tris</i> (1-phenyl-1,3-butanedionato)chromium(III)				
	$\Delta_{\text{sub}} H$		186 ± 2	298	C	[1987RIB/FER]
C₃₀H₃₀F₂₁CrO₆	[17966-86-8]	<i>tris</i> (1,1,1,2,2,3,3-heptafluoro-7,7-dimethyl-4,6-octanedionato)chromium(III)				
	$\Delta_{\text{sub}} H$	(323–353)	37.7 ± 0.8	338		[1972FON/POM]
C₃₃H₅₇CrO₆	[14434-47-0]	<i>tris</i> (2,2,6,6-tetramethyl-3,5-heptanedionato)chromium(III)				
	$\Delta_{\text{sub}} H$	(413–443)	85		TGA	[2000FAH/BAR]
	$\Delta_{\text{sub}} H$		133 ± 2	298	C	[1987RIB/FER]
CrI₂	[13478-28-9]	chromium(II) iodide				
	$\Delta_{\text{sub}} H$	(943–1054)	298.7	298		[1956ALL]
Cs (cesium)						
C₅H₉CsO₂	[20442-70-0]	cesium pivalate				
	$\Delta_{\text{sub}} H$		163.5 ± 7.2			[1998KHO/RYK]
CsI	[7789-17-5]	cesium iodide				
	$\Delta_{\text{sub}} H$		195.6	298	GS	[1998PAN/MAL]
	$\Delta_{\text{sub}} H$		193.1	298	T	[1985VEN/PRA, 1998PAN/MAL]
	$\Delta_{\text{sub}} H$		193.1	298	T	[1984COR, 1998PAN/MAL]
	$\Delta_{\text{sub}} H$		191.1	298	MS	[1984VIS/HIL, 1998PAN/MAL]
Cu (copper)						
C₆H₁₂CuN₂S₄	[137-29-1]	<i>bis</i> (dimethyldithiocarbamate)copper				
	$\Delta_{\text{sub}} H$		156.0 ± 0.3	298	C	[1995RIB/REI]
	$\Delta_{\text{sub}} H$	(443–473)	147.4 ± 0.8	458	A	[1987STE/MAL, 1978TAV/NEE]
	$\Delta_{\text{sub}} H$		149.0 ± 2.5		GC	[1976TAV/NEE]
	$\Delta_v H$	(443–473)	147.4	458		[1999DYK/SVO]

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₂ H ₁₈ CuO ₄	[14781-49-8]	<i>bis</i> (3-methyl-2,4-pentanedionato)copper(II)				
	$\Delta_{\text{sub}}H$		130.7 ± 1	396.7	ME	[1992RIB/FER3]
	$\Delta_{\text{sub}}H$		135.6 ± 1	298	ME	[1992RIB/FER3]
	$\Delta_{\text{sub}}H$		132.7 ± 2.5	298	C	[1992RIB/FER3]
C ₁₄ H ₁₆ CuF ₆ O ₄	[33896-35-4]	<i>bis</i> (1,1,1-trifluoro-5-methylhexane-2,4-dione)copper(II)				
	$\Delta_{\text{sub}}H$		122.4 ± 0.9	298	ME	[1998RIB/GON]
C ₁₄ H ₂₈ CuN ₂ S ₄	[14354-07-5]	<i>bis</i> (dipropyldithiocarbamate)copper				
	$\Delta_{\text{sub}}H$	(440–465)	129.5 ± 2.9	452.5	A	[1987STE/MAL, 1978TAV/NEE]
	$\Delta_{\text{sub}}H$		118.4 ± 3.3			[1978TAV/NEE]
	Δ_vH	(422–453)	118.4	437		[1999DYK/SVO]
C ₁₆ H ₈ CuF ₆ O ₄ S ₂	[13928-09-1]	<i>bis</i> (thenoyltrifluoroacetate)copper(II)				
	$\Delta_{\text{sub}}H$		167.9 ± 7.4	298	C	[2006RIB/SAN2]
C ₁₆ H ₈ CuF ₆ O ₆	[13928-10-4]	<i>bis</i> (4,4,4-trifluoro-1-(2-furanyl)butane-1,3-dione)copper(II)				
	$\Delta_{\text{sub}}H$		161.1 ± 2.1	298	ME	[1998RIB/GON]
C ₁₆ H ₂₀ CuF ₆ O ₄	[na]	<i>bis</i> (pivaloyltrifluoroacetate)copper				
	Δ_vH	(381–443)	76.5 ± 2.0		GS	[1993SYO/GOL]
C ₁₆ H ₂₀ CuF ₆ O ₄	[150026-91-8]	<i>bis</i> (1,1,1-trimethyl-5,5,5-trifluoro-2,4-pentanedionato)copper (II)				
	$\Delta_{\text{sub}}H$		120.2 ± 1.0	298	ME	[1998RIB/GON]
	$\Delta_{\text{sub}}H$	(353–379)	102 ± 3	366	T	[1993SYO/GOL]
	Δ_vH	(381–443)	76.5 ± 2	412	T	[1993SYO/GOL]
C ₁₆ H ₂₀ CuF ₆ O ₄	[220869-88-5]	<i>bis</i> (1,1,1-trifluoro-5-methylheptane-2,4-dione)copper(II)				
	$\Delta_{\text{sub}}H$		122.5 ± 0.9	298	ME	[1998RIB/GON]
(C ₁₆ H ₂₀ CuF ₆ O ₄)-(C ₁₀ H ₂₀ O ₅)	[na]	<i>bis</i> (1,1,1-trimethyl-5,5,5-trifluoro-2,4-pentanedionato)copper (II)-15-crown-5 complex				
	Δ_vH	(368–443)	80.2 ± 2	405	T	[1993SYO/GOL]
C ₁₆ H ₂₀ CuF ₆ O ₄ -(C ₁₀ H ₂₀ O ₅)	[na]	<i>bis</i> (pivaloyltrifluoroacetate)copper- 15-crown-5 complex				
	Δ_vH	(368–443)	80.2 ± 2.0		GS	[1993SYO/GOL]
C ₁₆ H ₂₆ CuO ₄	[15716-72-0]	<i>bis</i> (5,5-dimethyl-2,4-hexanedionato)copper(II)				
	$\Delta_{\text{sub}}H$		NA			[1978IGU/CHU]
C ₁₈ H ₁₂ CuN ₂ O ₂	[10380-26-6]	<i>bis</i> (8-hydroxyquinolino)copper(II)				
	$\Delta_{\text{sub}}H$		168.7 ± 7.3	298	ME	[1994RIB/MAT]
	$\Delta_{\text{sub}}H$	(478–503)	160.3 ± 3	491	ME	[1984BUR/MOR]
	$\Delta_{\text{sub}}H$		170 ± 3	298		[1984BUR/MOR]
C ₁₈ H ₁₄ CuN ₄	[41283-96-9]	dibenzotetra-aza-annulene copper(II) complex				
	$\Delta_{\text{sub}}H$	(493–553)	99.7 ± 8.7	523	T	[1983FER/QUA]
C ₁₈ H ₃₀ CuO ₄	[15321-96-7]	<i>bis</i> (2,2-dimethylheptan-3,5-dionato)copper(II)				
	$\Delta_{\text{sub}}H$	(344–364)	125.0 ± 1.3	354	TE	[1995RIB/MON]
	$\Delta_{\text{sub}}H$	(344–364)	127.8 ± 1.3	298	TE	[1995RIB/MON]
	$\Delta_{\text{sub}}H$	(344–364)	125.1 ± 0.5	354	ME	[1995RIB/MON]
	$\Delta_{\text{sub}}H$	(344–364)	127.9 ± 0.5	298	ME	[1995RIB/MON]
C ₁₈ H ₃₀ CuO ₄	[17653-77-9]	<i>bis</i> (2,6-dimethylheptan-3,5-dionato)copper(II)				
	$\Delta_{\text{sub}}H$		118.0 ± 347	298		[1984RIB/RIB]
C ₁₈ H ₃₆ CuN ₂ S ₄	[13927-71-4]	<i>bis</i> (dibutyldithiocarbamate)copper				
	Δ_vH	(423–468)	121.8	445		[1999DYK/SVO]

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound					
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference	
C₁₈H₃₆CuN₂S₄	[51205-55-1] $\Delta_v H$	<i>bis</i> (diisobutyldithiocarbamate)copper (425–445)	101.8	435		[1999DYK/SVO]	
C₂₀H₁₂CuF₆O₄	[14126-89-7] $\Delta_{\text{sub}} H$	<i>bis</i> (4,4,4-trifluoro-1-phenylbutane-1,3-dione)copper(II)	172.1 ± 3.1	298	ME	[1998RIB/GON]	
C₂₀H₁₆CuN₂O₂	[14522-43-1] $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$	<i>bis</i> (8-hydroxy-2-methylquinoline)copper(II) (402–419)	166.5 ± 3.4	410	ME	[1998RIB/MAT3]	
			172.1 ± 3.4	298	ME	[1998RIB/MAT3]	
C₂₀H₁₈CuO₄	[14128-84-8] $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$	<i>bis</i> (1-phenylbutane-1,3-dionato)copper(II) (429–450)	152.2 ± 1.7	439	TE	[1995RIB/MON]	
			159.3 ± 1.7	298	TE	[1995RIB/MON]	
			152.2 ± 1.9	439	ME	[1995RIB/MON]	
			159.3 ± 1.9	298	ME	[1995RIB/MON]	
			160 ± 4	298	C	[1979RIB/REI]	
C₂₀H₂₀CuF₁₄O₄	[38926-19-1] $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$	<i>bis</i> (1,1,1,2,2,3,3-hetafluoro-7,7-dimethyloctane-4,6-dionato)copper(II)	122.8 ± 0.7	298	ME	[1998RIB/GON]	
			NA			[1978IGU/CHU]	
C₂₀H₃₄CuO₄	[141752-16-3] $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$	<i>bis</i> (2,2,6-trimethylheptan-3,5-dionato)copper(II) (346–362)	127.4 ± 0.7	354	ME	[1995RIB/MON]	
			130.2 ± 0.7	298		[1995RIB/MON]	
			127.8 ± 1.5	354	TE	[1995RIB/MON]	
			130.6 ± 1.5	298		[1995RIB/MON]	
			129.0 ± 1.3	351	ME	[1995RIB/MON]	
			131.7 ± 1.3	298		[1995RIB/MON]	
			126.4 ± 1.1	298		[1984RIB/RIB]	
C₂₂H₂₄CuN₂O₂	[15214-38-7] $\Delta_{\text{sub}} H$	<i>bis</i> [(4-phenylimino)-2-pentanoato]copper(II)	128.1 ± 0.8	298	ME,TE	[1990RIB/RIB]	
C₂₂H₃₆CuF₂O₄	[1148044-73-8] $\Delta_{\text{sub}} H$	<i>bis</i> (2,2,6,6-tetramethyl-4-fluoroheptane-2,4-dionato)copper(II) (392–453)	115.6 ± 1.1	422	ME	[2008ZHE/MOR2]	
C₂₂H₃₈CuO₄	[14040-05-2] $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$	<i>bis</i> (2,2,6,6-tetramethyl-heptane-3,5-dionato)copper(II) (375–435)		96 ± 2	GS	[2009JOH/SEL]	
				127.6 ± 0.4	361	TE	[2001COL/LAU]
				127.2 ± 1.7	351	TE	[2001COL/LAU]
				120		TGA	[2000FAH/BAR]
				74.8		TGA,DTA	[1996YUA/MEN]
				100	404	T	[1996RAP/DES]
				124.5 ± 0.8	372	ME	[1995RIB/MON]
				129.1 ± 0.8	298		[1995RIB/MON]
				124.6	407		[1993TOB/LAN]
				123.6	445		[1992WAF/MUS]
				105.9		GS	[1990YUH/KIK]
				111.6			[1988FED/VOI, 1993TOB/LAN]
				122.8 ± 6.5	298	C	[1984RIB/RIB]
				112		C	[1979IGU/CHU]
C₂₄H₃₀CuN₄O₄	$\Delta_{\text{sub}} H$	<i>bis</i> (N-benzoyl-N',N'-diethylureato)copper(II)	180.9 ± 3.7	298	C	[2001RIB/RIB2]	
C₂₈H₁₆CuF₆O₄	[30983-56-3] $\Delta_{\text{sub}} H$	<i>bis</i> (4,4,4-trifluoro-1-(2-naphthalenyl)butane-1,3-dione)copper(II)	208.4 ± 4.9	298	ME	[1998RIB/GON]	

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound						
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference		
C₃₀H₂₂CuO₄	[58179-06-9] $\Delta_{\text{sub}}H$	<i>bis</i> (dibenzoylmethanato)copper(II)		230.7 ± 8.2	298	C	[2006RIB/SAN2]	
C₃₀H₅₄CuO₆	[952723-42-1] Δ_vH	<i>bis</i> (2,6-dimethyl-6-methoxydodecane-3,5-dionato) copper (II)		(340–395)	126.6 ± 1.9	367	ME	[2006LIS/SEM]
C₃₂H₁₆CuN₈	[na] $\Delta_{\text{sub}}H$	copper(II) α -phthalocyanine		114			TGA	[1995YAS/TAK]
C₃₂H₁₆CuN₈	[147-14-8] $\Delta_{\text{sub}}H$	copper(II) β -phthalocyanine		(618–713)	231.8 ± 2.1		ME	[2000SEM/BAS]
	$\Delta_{\text{sub}}H$				211.1		TGA	[1995YAS/TAK]
	$\Delta_{\text{sub}}H$			(657–863)	266.1			[1969HAM]
	$\Delta_{\text{sub}}H$			(657–722)	266.1 ± 5.1		ME	[1965CUR/SHA, 1970BON/CAT]
C₃₉H₅₉F₁₂O₈CuY	[160364-36-3] $\Delta_{\text{sub}}H$	<i>bis</i> (hexafluoroisopropoxy) <i>tris</i> (2,2,6,6-tetramethylheptan-3,5-dionato)copper(II)yttrium(III)		(370–410)	81.2	390		[1996LAB/HUB]
C₄₄H₂₈CuN₄	[14172-91-9] $\Delta_{\text{sub}}H$	5,10,15,20-tetraphenylporphine copper(II)		160 ± 5			GS	[2000PER/GOL]
Dy (dysprosium)								
C₁₅H₁₅Dy	[12088-04-9] $\Delta_{\text{sub}}H$	<i>tris</i> (cyclopentadienyl)dysprosium(III)		105.0 ± 2.1				[1973DEV/BOR]
C₃₀H₃₀DyF₂₁O₆	[18232-98-3] $\Delta_{\text{sub}}H$	<i>tris</i> (1,1,1,2,2,3,3-heptafluoro-7,7-dimethyloctane-4,6-dione)dysprosium(III)		(370–385)	156.5 ± 2.9		ME	[1971SWA/KAR]
C₃₃H₅₇DyO₆	[15522-69-7] $\Delta_{\text{sub}}H$	<i>tris</i> (2,2,6,6-tetramethylpentane-2,4-dionato)dysprosium(III)		(373–388)	171.5	380	ME	[1981AMA/SAT]
	$\Delta_{\text{sub}}H$			(388–413)	152.7	400	ME	[1981AMA/SAT]
	$\Delta_{\text{sub}}H$			(410–456)	133.5	433	BG	[1969SIC/DUB]
	Δ_vH			(456–500)	86.2		BG	[1969SIC/DUB]
DyBr₃	[14456-48-5] $\Delta_{\text{sub}}H$	dysprosium tribromide		(878–1151)	289 ± 6.0	298	TE	[1999BRU/VAS]
DyCl₃	[10025-74-8] $\Delta_{\text{sub}}H$	dysprosium trichloride		(924–1214)	283 ± 5.0	298	TE	[1999BRU/VAS]
DyI₃	[15474-63-2] $\Delta_{\text{sub}}H$	dysprosium triiodide		(889–1157)	282 ± 4.0	298	TE	[1999BRU/VAS]
Er (erbium)								
C₁₅H₁₂ErF₉O₆	[70332-27-3] $\Delta_{\text{sub}}H$	<i>tris</i> (1,1,1-trifluoro-2,4-pentanedionato)erbium(III)		(473–494)	79.5 ± 11.5	484		[1996ZVE/CHE]
C₁₅H₁₅Er	[39330-74-0] $\Delta_{\text{sub}}H$	<i>tris</i> (cyclopentadienyl)erbium(III)		(503–558)	97.2 ± 3.2	530		[1996ZVE/CHE]
	$\Delta_{\text{sub}}H$				97.1 ± 3.3			[1973DEV/BOR]
C₂₄H₃₃Er	[130521-76-5] $\Delta_{\text{sub}}H$	<i>tris</i> [(1,2,3,4,5- η)-1-(1-methylethyl)-2,4-cyclopentadien-1-yl]erbium(III)		(464–502)	78.6 ± 3.0	483		[1996ZVE/CHE]
C₃₀H₃₀ErF₂₁O₆	[17978-75-5] $\Delta_{\text{sub}}H$	<i>tris</i> (1,1,1,2,2,3,3-heptafluoro-7,7-dimethyloctane-4,6-dione)erbium(III)		(349–362)	154.8 ± 4.2		ME	[1971SWA/KAR]
C₃₃H₅₇ErO₆	[14319-09-6] $\Delta_{\text{sub}}H$	<i>tris</i> (2,2,6,6-tetramethylheptane-3,5-dionato)erbium(III)			130.8 ± 2.2	298	DSC	[1999SAN/PET]

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound		T_m (K)	Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)			
	$\Delta_{\text{sub}}H$	(471–505)	93.9 ± 4.6	488		[1996ZVE/CHE]
	$\Delta_{\text{sub}}H$	(363–418)	154	390	ME	[1981AMA/SAT]
	$\Delta_{\text{sub}}H$	(358–381)	149.3 ± 1.7		ME	[1971SWA/KAR]
	$\Delta_{\text{sub}}H$	(410–454)	133.2	432	BG	[1969SIC/DUB]
	Δ_vH	(454–490)	85.6		BG	[1969SIC/DUB]
Eu (europium)						
C₁₅H₃EuF₁₈O₆	[14592-81-5]	<i>tris</i> (1,1,1,5,5,5-hexafluoro-2,4-pentanedionato)europium(III)				
	$\Delta_{\text{sub}}H$	(340–380)	129.4 ± 9.5			[2007MAL/ALI]
		Note: Authors report that the vapor phase consists largely of dimers				
C₃₃H₅₇EuO₆	[15522-71-1]	<i>tris</i> (2,2,6,6-tetramethylheptane-3,5-dionato)europium(III)				
	$\Delta_{\text{sub}}H$	(363–433)	179.9	398	ME	[1981AMA/SAT]
	$\Delta_{\text{sub}}H$	(373–423)	180		ME	[1979AMA/SAT]
	$\Delta_{\text{sub}}H$	(430–466)	165.4	448	BG	[1969SIC/DUB]
	Δ_vH	(466–490)	87.4		BG	[1969SIC/DUB]
F (fluorine)						
F₃NO	[13847-65-9]	trifluoroamine oxide				
	Δ_vH	(116–191)	16.1			[1968FOX/MAC]
F₅I	[7783-66-6]	iodine pentafluoride				
	Δ_vH	(283–378)	39.3	330		[1971OSB/SCH]
HF	[7664-39-3]	hydrogen fluoride				
	Δ_vH	(240–290)	25.2	265		[1934CAM/CAM]
	Δ_vH	(190–320)	25.2	255		[1924SIM]
Fe (iron)						
C₂FeN₂O₄	[13682-74-1]	dicarbonyldinitrosyl iron				
	$\Delta_{\text{sub}}H$	(272–291)	47.2	281.5	A	[1987STE/MAL]
C₄FeI₂O₄	[14878-30-9]	iron tetracarbonyl diiodide				
	$\Delta_{\text{sub}}H$		86.0 ± 4.0	298		[1982PIL/SKI, 1979CON/DEM]
C₄H₆FeO₄Si₂	[26469-80-7]	tetracarbonyl disilyl iron				
	Δ_vH	(329–377)	43.8	353	T	[1969AYL/CAM3]
C₄H₁₆Cl₂FeN₈S₄	[28813-18-5]	<i>trans</i> -dichloro- <i>tetrakis</i> (thiourea)iron(II)				
	$\Delta_{\text{sub}}H$	(372–405)	110 ± 20			[1970ASH]
C₅FeO₅	[13463-40-6]	iron pentacarbonyl				
	Δ_vH	(254–304)	40.1 ± 0.5	279		[1974GIL/SUL]
	Δ_vH	(266–353)	39.0	309		[1970VAL/KIL]
	Δ_vH		38.1 ± 0.4	298		[1959LEA/SPI]
	Δ_vH		40.2 ± 0.8			[1959COT/FIS, 1982PIL/SKI]
	Δ_vH	(266–378)	37.6	281		[1947STU]
C₆H₅FeO₃	[12189-10-5]	allyliron tricarbonyl iodide				
	$\Delta_{\text{sub}}H$		84.5 ± 4.0	298		[1982PIL/SKI, 1979CON/DEM]
C₇H₆FeO₃	[na]	1,3-butadiene iron tricarbonyl				
	Δ_vH		49.0 ± 4.2			[1976BRO/CON, 1982PIL/SKI]
C₈H₆Fe₂O₆S₂	[14878-96-7]	hexacarbonyl <i>bis</i> (methanethiolato)diiron				
	$\Delta_{\text{sub}}H$		102.8	333	C	[1995CON/GOB]
	$\Delta_{\text{sub}}H$		109.8	298		[1995CON/GOB]

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No	Compound					
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference	
C ₉ Fe ₂ O ₉	[15321-51-4]	diiron nonacarbonyl					
			(296–314)	135.3	305	A	[1987STE/MAL]
				75.3 ± 21.0	298		[1982PIL/SKI, 1972CON/SKI]
C ₉ H ₁₂ FeO	[na]	<i>bis</i> (1,3-butadiene)ironcarbonyl					
				76.1 ± 4.2	298		[1982PIL/SKI, 1976BRO/CON]
C ₁₀ H ₁₀ Fe	[102-54-5]	ferrocene					
				17.8	448.5	DSC	[2008LOU/PIN]
				17.49	447.6	DSC	[2001DAB/MIS]
				0.9	163.9		
				4.14	242		
				17.78	448.2		[1981OGA/SOR, 1969JOE/GJA]
				73.3 ± 1.9	298	ME	[2008LOU/PIN]
			(298–304)	73.2 ± 1.9	301	ME	[2008LOU/PIN]
				72.7 ± 0.2	298	C	[2008LOU/PIN]
			(290–363)	72.7 ± 0.2	298	GS	[2007EME/VER]
				72.6 ± 0.1	313	C	[2006MON/SAN]
				73.3 ± 0.1	298	C	[2006MON/SAN]
				74.9 ± 1.7	298	C	[2004SAN/SCH]
				73.3 ± 0.1	298	C	[2001KIY/MIN]
				73.1 ± 1.4	333	DSC	[2001ROJ/ORO, 2007EME/VER]
				74.1 ± 1.4	298	DSC	[2001ROJ/ORO, 2007EME/VER]
				74.3 ± 0.4	298	ME	[1995TOR/GUD2]
				73.2 ± 0.7	298	C	[1995TOR/GUD2]
			(292–300)	72.5 ± 1.0	296	ME	[1990RIB/MON]
				72.4 ± 1.0	298		[1990RIB/MON]
			(294–302)	70.3 ± 1.0	298	ME	[1989MIN, 1990RIB/MON]
			(278–309)	72.1 ± 0.4	294	ME	[1988TOR/BAR2]
				71.9 ± 0.4	298		[1988TOR/BAR2]
			(348–446)	64.6	397	A	[1987STE/MAL]
				75.6 ± 0.4	298	TE,ME,DM	[1983JAC/VAN]
				74.0 ± 2	298	TE	[1981PEL/TOM]
			(328–398)	70.0 ± 2		DSC	[1980MUR/CAV]
			(328–398)	71.9 ± 2.0	298	DSC	[1980MUR/CAV, 2007EME/VER]
				72.6 ± 1.4	298	ME	[1980CAL/DIA]
			(348–451)	67.9	298		[1977BAR/GAI, 2007EME/VER]
				73.6 ± 0.4	298		[1982PIL/SKI, 1975TEL/KIR]
				74.1 ± 1.7	298	TCM	[1973DEK/OON]
			(385–455)	84.0 ± 2.0		DSC	[1971BEE/LIN]
			(385–455)	87.6 ± 2.0	298	DSC	[1971BEE/LIN, 2007EME/VER]
				72.7 ± 2	298	ME	[1969AND/WES]
				76.6 ± 1	298	ME	[1962EDW/KIN]
			(295–303)	76.8 ± 0.9	298	ME	[1960EDW/KIN, 2007EME/VER]
			(323–367)	83.3		ME	[1959COR/SCH]
			(323–367)	84.7	298	ME	[1959COR/SCH, 2007EME/VER]
			(357–455)	70.5	406		[1952KAP/KES]
	(357–455)	73.7	298	BG	[1952KAP/KES, 2007EME/VER]		
	(456–523)	47.3	471	A	[1987STE/MAL, 1999DYK/SVO]		
	(451–523)	49.8	466	A	[1987STE/MAL, 1977BAR/GAI]		
	(519–604)	44.7	561	EB	[1972NIS/SOK]		
	(519–604)	64.7 ± 0.4	298	EB	[1972NIS/SOK, 2007EME/VER]		
		47.3	456		[1952KAP/KES]		
C ₁₀ H ₁₀ Fe ₂ O ₆ S ₂	[28829-01-8]	hexacarbonyl <i>bis</i> (ethanethiolato)diiron					

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound		T_m (K)	Method	Reference	
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)				
C₁₀H₁₄FeO₄	$\Delta_{\text{sub}}H$		105.4	340	C	[1995CON/GOB]	
	$\Delta_{\text{sub}}H$		112	298	C	[1995CON/GOB]	
C₁₀H₁₄FeO₄	[14024-17-0]	<i>bis</i> (2,4-pentanedionato)iron(II)					
	$\Delta_{\text{sub}}H$	(330–368)	131.2 ± 8.7	298	ME	[1990MAL/ALI]	
	$\Delta_{\text{sub}}H$		117.6	385		[1970GOE/BLO]	
C₁₁H₈FeO₃	[12093-05-9]	cyclooctatetraeneirontricarbonyl					
	$\Delta_{\text{sub}}H$		87.0 ± 4.0	298		[1982PIL/SKI, 1979CON/DEM]	
C₁₁H₁₀FeO	[12093-10-6]	ferrocenecarboxaldehyde					
	$\Delta_{\text{trs}}H$		11.6	317.4			
	$\Delta_{\text{fus}}H$		2.5	397.2	DSC	[2008LOU/PIN]	
	$\Delta_{\text{trs}}H$		0.89	293.5			
	$\Delta_{\text{trs}}H$		13.3	317			
	$\Delta_{\text{fus}}H$	(13–405)	2.76	397.6	AC	[2007KAN/SOR]	
	$\Delta_{\text{trs}}H$		11.7	316.4			
	$\Delta_{\text{fus}}H$		2.05	396.7		[1978DAN/LEA]	
	$\Delta_{\text{sub}}H$		89.7 ± 5.1	298	ME	[2008LOU/PIN]	
	$\Delta_{\text{sub}}H$	(302–312)	89.9 ± 5.1	305	ME	[2008LOU/PIN]	
	$\Delta_{\text{sub}}H$		87.9 ± 3.5	298	ME	[2008LOU/PIN]	
	$\Delta_{\text{sub}}H$	(302–312)	87.3 ± 3.5	310	ME	[2008LOU/PIN]	
	C₁₂H₁₂FeO	[1271-55-2]	acetylferrocene				
		$\Delta_{\text{sub}}H$	(329–358)	115.6 ± 2.5	298		[1981PEL/TOM]
C₁₂H₁₂FeO	[1273-86-5]	ferrocenemethanol					
	$\Delta_{\text{fus}}H$		23.82	351.4	DSC	[2001DAB/MIS]	
	$\Delta_{\text{sub}}H$	(313–320)	102.8 ± 0.5	298	GS	[2007EME/VER]	
	Δ_vH	(353–393)	87.0 ± 0.8	298	GS	[2007EME/VER]	
C₁₂H₁₄Fe	[1273-89-9]	ethylferrocene					
	$\Delta_{\text{fus}}H$		12.29	273.9		[2003KOZ/KAR, 2003KAR/KOZ]	
	Δ_vH	(297–320)	65.1 ± 2.7	308	ME	[2003KAR/KOZ]	
C₁₂H₁₄Fe	[1291-47-0]	1,1'-dimethylferrocene					
	$\Delta_{\text{fus}}H$		17.66	312.6	DSC	[2008LOU/PIN]	
	$\Delta_{\text{sub}}H$		84.5 ± 1.9	298	ME	[2008LOU/PIN]	
C₁₂H₁₄FeO	[1277-49-2]	1-ferrocenyl ethanol					
	$\Delta_{\text{fus}}H$		14.75	343.7	DSC	[2008LOU/PIN]	
	$\Delta_{\text{fus}}H$		26.65	366.5	DSC	[2001DAB/MIS]	
	$\Delta_{\text{sub}}H$		102.4 ± 0.9	298	C	[2008LOU/PIN]	
C₁₂Fe₃O₁₂	[17685-52-8]	triiron dodecacarbonyl					
	$\Delta_{\text{sub}}H$		96.0 ± 21.0	298		[1982PIL/SKI, 1972CON/SKI]	
C₁₃H₁₄FeO	[1271-79-0]	propanoyl ferrocene					
	$\Delta_{\text{fus}}H$		19.2	311.6	AC	[2009KRO/DRU]	
	$\Delta_{\text{fus}}H$		19.7	311.6	DSC	[2009KRO/DRU]	
	$\Delta_{\text{sub}}H$		99.0 ± 1.5	298	GS	[2009KRO/DRU]	
	Δ_vH		80.7 ± 0.8	298	GS	[2009KRO/DRU]	
C₁₃H₁₆Fe	[1273-92-3]	propylferrocene					

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound		Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)		
	$\Delta_{\text{trs}}H$		1.23	186.7	
	$\Delta_{\text{fus}}H$		12.7	276.8	AC [2009KRO/DRU]
	$\Delta_{\text{fus}}H$		14.6	278.2	DSC [2009KRO/DRU]
	Δ_vH		69.4 ± 0.8	298	GS [2009KRO/DRU]
C₁₃H₁₆FeO	[34978-83-1]	<i>bis</i> (1,3-cyclohexadiene)ironcarbonyl			
	$\Delta_{\text{sub}}H$		95.0 ± 4.2	298	[1982PIL/SKI, 1976BRO/CON]
C₁₃H₁₇NFe	[1271-86-9]	N,N-dimethyl(aminomethyl) ferrocene			
	$\Delta_{\text{fus}}H$		14.6	281.5	DSC [2008LOU/PIN]
	$\Delta_{\text{trs}}H$		0.32	134	
	$\Delta_{\text{fus}}H$		15.01	280.9	[2003KAR/KOZ]
	$\Delta_{\text{fus}}H$		15.01	279.9	[2002KAR/SHE]
	Δ_vH		73.8 ± 0.4	298	C [2008LOU/PIN]
	Δ_vH	(295–319)	66.3 ± 3.9	307	ME [2003KAR/KOZ]
C₁₄H₁₀Fe₂O₄	[12154-95-9]	<i>bis</i> (η^5 -cyclopentadienyl)iron(II)dicarbonyl			
	$\Delta_{\text{fus}}H$		30.8	472.9	[2008KOZ/MAR]
C₁₄H₁₄FeO₂	[1273-94-5]	1,1'-diacetylferrocene			
	$\Delta_{\text{sub}}H$	(360–400)	91.9 ± 2.5	298	[1981PEL/TOM]
C₁₄H₁₈Fe	[31904-29-7]	n-butylferrocene			
	$\Delta_{\text{fus}}H$		21.43	281.5	[2002KOZ/KAR, 2003KAR/KOZ]
	Δ_vH	(315–333)	75.0 ± 3.0	324	ME [2003KAR/KOZ]
C₁₄H₁₈Fe	[1273-97-8]	1,1'-diethylferrocene			
	$\Delta_{\text{fus}}H$	(5–300)	21.03	236.9	AC [1999DOM/KAR]
C₁₅H₃F₁₈FeO₆	[17786-67-3]	<i>tris</i> (1,1,1,5,5,5-hexafluoro-2,4-pentanedionato)iron(III)			
	$\Delta_{\text{sub}}H$	(333–363)	60.0		TGA [2000FAH/BAR]
C₁₅H₁₂F₉FeO₆	[14526-22-8]	<i>tris</i> (1,1,1-trifluoro-2,4-pentanedionato)iron(III)			
	$\Delta_{\text{sub}}H$	(373–403)	96.0		TGA [2000FAH/BAR]
	$\Delta_{\text{sub}}H$	(378–438)	104.6 ± 0.8		GS [1985MAT/KUW]
	$\Delta_{\text{sub}}H$		80.3	433	[1977VOL/MAZ]
	$\Delta_{\text{sub}}H$		128.9	345	[1970GOE/BLO]
	$\Delta_{\text{sub}}H$		87.0		[1960BER/TRU, 1965BER/TRU]
	Δ_vH	(392–428)	87.0 ± 1.2	410	[1978CHU/IGU]
C₁₅H₁₈FeOS₂	[122380-51-2]	1,4,6-oxadithiacyclooctan-5-ylferrocene			
	$\Delta_{\text{fus}}H$		24.4	383.3	DSC [1992HUA/WAN]
C₁₅H₂₁FeO₆	[14024-18-1]	<i>tris</i> (2,4-pentanedionato)iron(III)			
	$\Delta_{\text{fus}}H$		30.1	459	DSC [2004SAB/MAR]
	$\Delta_{\text{fus}}H$		25.3	460	DSC [2004SAB/MAR]
	$\Delta_{\text{fus}}H$		22.6	462	[1984MUR/HIL]
	$\Delta_{\text{fus}}H$		25.9	454	[1970MEL/MER2]
	$\Delta_{\text{sub}}H$	(413–443)	112		TGA [2000FAH/BAR]
	$\Delta_{\text{sub}}H$		118		TGA [1997GIL/BOT]
	$\Delta_{\text{sub}}H$	(369–388)	124.6 ± 0.9	378	TE, ME [1996RIB/MON]
	$\Delta_{\text{sub}}H$		128.6 ± 0.9	298	[1996RIB/MON]
	$\Delta_{\text{sub}}H$	(338–355)	114.2 ± 1.5		[1992GER/GER]
	$\Delta_{\text{sub}}H$	(309–360)	126.4 ± 3.1	298	ME [1990MAL/ALI]
	$\Delta_{\text{sub}}H$		138.4 ± 5.2	298	C [1985MUR/SAK]
	$\Delta_{\text{sub}}H$		100	395	T [1981MAS/BAR]

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound		T_m (K)	Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)			
	$\Delta_{\text{sub}}H$		113.6 ± 3.8			[1980SAC/HIL]
	$\Delta_{\text{sub}}H$		99 ± 0.8			[1979RIB/REI, 1981MAS/BAR, 1970MEL/MER2]
	$\Delta_{\text{sub}}H$		114.2	385		[1970GOE/BLO]
	$\Delta_{\text{sub}}H$		65.3 ± 3.3	298		[1982PIL/SKI, 1968HIL/IRV2]
	$\Delta_{\text{sub}}H$		97.9		I	[1964FAR/JON]
	$\Delta_{\text{sub}}H$		81.6			[1960BER/TRU, 1965BER/TRU]
C₁₅H₃₀FeN₃S₆	[34768-31-5]	<i>tris</i> (diethyldithiocarbamato)iron(III)				
	$\Delta_{\text{sub}}H$		65.7 ± 1.7	246		[1970DAS/WEN]
C₁₇H₁₄FeO	[1272-44-2]	benzoylferrocene				
	$\Delta_{\text{fus}}H$	(6–372)	29.9	380.7	AC	[2008KRO/DRU]
	$\Delta_{\text{sub}}H$	(342–379)	119.9 ± 0.7	298	GS	[2007EME/VER]
	$\Delta_{\text{sub}}H$	(358–382)	116.3 ± 6	298	TE,ME	[1983PEL/GIG]
	Δ_vH	(384–429)	98.2 ± 0.3	298	GS	[2007EME/VER]
C₁₇H₁₆Fe	[32994-54-0]	(phenylmethyl)ferrocene				
	$\Delta_{\text{fus}}H$	(6–372)	26.8	349.9	AC	[2008KRO/DRU]
	$\Delta_{\text{sub}}H$	(312–341)	109.3 ± 0.7	298	GS	[2007EME/VER]
	Δ_vH	(351–377)	90.6 ± 0.6	298	GS	[2007EME/VER]
C₁₇H₂₂FeO₂S₂	[122395-65-7]	1,9-dioxa-4,6-dithiacyclundecan-5-ylferrocene				
	$\Delta_{\text{fus}}H$		29.4	371.7	DSC	[1992HUA/WAN]
C₁₈H₂₇FeO₆	[13978-46-6]	<i>tris</i> (3-methylpentane-2,4-dionato)iron(III)				
	$\Delta_{\text{sub}}H$		164.5	422		[1992RIB/FER]
C₁₉H₂₆FeO₂S₃	[122395-70-4]	1,9-dioxa-4,6,12-trithiacyclotetradecan-5-ylferrocene				
	$\Delta_{\text{fus}}H$		40.0	367.1	DSC	[1992HUA/WAN]
C₁₉H₂₆FeO₃S₂	[122395-66-8]	1,9,12-trioxa-4,6-dithiacyclotetradecan-5-ylferrocene				
	$\Delta_{\text{fus}}H$		32.1	349.7	DSC	[1992HUA/WAN]
C₂₀H₃₀Fe	[12126-50-0]	<i>bis</i> (η^5 -pentamethylcyclopentadienyl)iron				
	$\Delta_{\text{us}}H$		4.3	402.6		
	$\Delta_{\text{us}}H$		4.87	503.7	DSC	[2008LOU/PIN]
		Note: Decomposed upon melting				
	$\Delta_{\text{sub}}H$		99.0 ± 2.4	298	ME	[2008LOU/PIN]
	$\Delta_{\text{sub}}H$	(355–376)	95.7 ± 2.4	365	ME	[2008LOU/PIN]
	$\Delta_{\text{sub}}H$		96.8 ± 0.6	298	C	[2001KIY/MIN]
C₂₄H₁₂F₉FeO₆S₃	[14319-78-9]	<i>tris</i> (1-(2-thenoyl)-4,4,4-trifluoro-1,3-butanedione)iron(III)				
	$\Delta_{\text{sub}}H$		U 46.4			[1960BER/TRU, 1965BER/TRU]
C₂₄H₁₈FeO₂	[12180-80-2]	1,1'-dibenzoylferrocene				
	$\Delta_{\text{sub}}H$	(358–381)	109.3 ± 6	298	TE,ME	[1983PEL/GIG]
C₃₀H₂₇FeO₆	[14323-17-2]	<i>tris</i> (benzoylacetato)iron(III)				
	$\Delta_{\text{sub}}H$		U 45.6		I	[1964FAR/JON]
C₃₃H₅₇FeO₆	[14876-47-2]	<i>tris</i> (2,2,6,6-tetramethylheptan-3,5-dionato)iron(III)				
	$\Delta_{\text{sub}}H$	(413–443)	111		TGA	[2000FAH/BAR]
	$\Delta_{\text{sub}}H$	(360–378)	128.5 ± 0.9	369	TE,ME	[1996RIB/MON]
	$\Delta_{\text{sub}}H$		129.3 ± 1.2	298		[1996RIB/MON]
	$\Delta_{\text{sub}}H$		106.7		ME	[1973BRU/CUR]

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound			Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)		
C₄₅H₃₃FeO₆	[14405-49-3] $\Delta_{\text{sub}}H$	<i>tris</i> (dibenzoylmethano)iron(III)	U 31.8		I	[1964FAR/JON]
Br₂Fe	[7789-46-0] $\Delta_{\text{sub}}H$	iron(II) dibromide	(655–833)	197.6 ± 2.0	744	TE,ME [1996BAR/BRU]
	$\Delta_{\text{sub}}H$			208 ± 2.0	298	[1996BAR/BRU]
	$\Delta_{\text{sub}}H$		(680–720)	196 ± 8	700	TE [1996BAR/BRU, 1960SIM/GRE]
	$\Delta_{\text{sub}}H$		(673–962)	197 ± 2	817	GS [1996BAR/BRU, 1955MAC/GRE]
	$\Delta_{\text{sub}}H$			210 ± 6	298	[1996BAR/BRU, 1955MAC/GRE]
	$\Delta_{\text{sub}}H$		(623–718)	197 ± 4	670	ME [1996BAR/BRU, 1955MAC/GRE]
FeCl₂	[7758-94-3] $\Delta_{\text{sub}}H$	iron(II) dichloride	(693–866)	198.9 ± 2.0	780	TE,ME [1996BAR/BRU]
	$\Delta_{\text{sub}}H$			204 ± 4.0	298	[1996BAR/BRU]
	$\Delta_{\text{sub}}H$		(694–745)	189 ± 8	719	TE [1960SIM/GRE, 1996BAR/BRU]
	$\Delta_{\text{sub}}H$		(621–658)	186 ± 12	640	MS [1958SCH/POR, 1996BAR/BRU]
	$\Delta_{\text{sub}}H$			193 ± 12	298	[1958SCH/POR, 1996BAR/BRU]
FeF₂	[7789-28-8] $\Delta_{\text{sub}}H$	iron(II) difluoride	(958–1178)	263 ± 2.0	1068	TE,ME [1996BAR/BRU]
	$\Delta_{\text{sub}}H$			271 ± 2.0	298	[1996BAR/BRU]
	$\Delta_{\text{sub}}H$		(848–1142)	263 ± 3	995	ME [1976ZHU/ALI, 1996BAR/BRU]
	$\Delta_{\text{sub}}H$			270 ± 3	298	[1976ZHU/ALI, 1996BAR/BRU]
Ga (gallium)						
C₃H₉Ga	[1445-79-0] $\Delta_{\text{fus}}H$	trimethyl gallium		11.8		Sub-Vap [2003FUL/RUZ]
	$\Delta_{\text{fus}}H$			11.05	257.9	[1996DOM/HEA]
	$\Delta_{\text{sub}}H$		(225–257)	47.4		[2003FUL/RUZ]
	$\Delta_{\text{sub}}H$		(247–257)	45.2	252	A [1987STE/MAL]
	Δ_vH		(259–263)	35.6		[2003FUL/RUZ]
	Δ_vH			33.1 ± 0.8		[1958LON/SAC, 1982PIL/SKI]
	Δ_vH			32.6		[1933KRA/TOO, 1958FOW/MOR]
C₄H₁₀ClGa	[30914-08-0] Δ_vH	diethylgallium chloride	(273–333)	59.9	303	[1991BUC/POT]
C₆H₉Ga	[1188-13-2] Δ_vH	trivinyl gallium	(298–373)	U 72.6	335	[1962OLI/STE]
		Note: Decomposition noted above 333 K				
C₆H₁₅Ga	[1115-99-7] $\Delta_{\text{fus}}H$	triethyl gallium		11.64	193.5	[1996DOM/HEA]
	Δ_vH		(299–387)	43.1 ± 1.6	343	[2001BAE, 2001BAE/CHE]
	Δ_vH			38.5 ± 0.4		[1973KOL/RAB, 1982PIL/SKI]
C₉H₂₁Ga	[54514-59-9] Δ_vH	triisopropyl gallium	(298–373)	49.0	335	[1962OLI/STE]
C₉H₂₁Ga	[29868-77-7] Δ_vH	tripropyl gallium	(316–385)	46.6 ± 0.5	350	[2001BAE]
	Δ_vH		(298–373)	49.2	335	[1962OLI/STE]
C₁₁H₂₄GaNS₂	[253595-30-1] Δ_vH	di- <i>tert</i> -butyl gallium dimethyldithiocarbamate	(374–427)	43 ± 1		TGA [1999KEY/BOT]
C₁₁H₂₄GaNS₂	[253595-34-5]	dibutyl gallium dimethyldithiocarbamate				

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound		T_m (K)	Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)			
	$\Delta_v H$	(385–424)	53 ± 1		TGA	[1999KEY/BOT]
C₁₁H₂₄GaNS₂	[253595-35-8]	di-sec-butyl gallium dimethyldithiocarbamate				
	$\Delta_v H$	(366–425)	44 ± 1		TGA	[1999KEY/BOT]
C₁₂H₂₇Ga	[15677-44-8]	tributyl gallium				
	$\Delta_v H$	(330–378)	51.6 ± 1.3	354		[2001BAE]
	$\Delta_v H$	(426–507)	56.2	441	A	[1987STE/MAL]
C₁₃H₂₈GaNS₂	[253595-32-3]	di-tert-butyl gallium diethyldithiocarbamate				
	$\Delta_v H$	(372–419)	48 ± 6		TGA	[1999KEY/BOT]
C₁₅H₃F₁₈GaO₆	[19648-92-1]	tris(1,1,1,5,5,5-hexafluoro-2,4-pentanedionato)gallium(III)				
	$\Delta_{\text{sub}} H$	(333–363)	53.0		TGA	[2000FAH/BAR]
C₁₅H₁₂F₉GaO₆	[15453-83-5]	tris(1,1,1-trifluoro-2,4-pentanedionato)gallium(III)				
	$\Delta_{\text{sub}} H$	(373–403)	75.0		TGA	[2000FAH/BAR]
	$\Delta_{\text{sub}} H$	(378–433)	118.8 ± 1.7		GS	[1985MAT/KUW]
	$\Delta_{\text{sub}} H$	(386–401)	89.4 ± 6.7			[1978IGU/CHU2]
	$\Delta_v H$	(401–459)	75.6 ± 0.5	430		[1978IGU/CHU2]
C₁₅H₂₁GaO₆	[14405-43-7]	tris(pentane-2,4-dionato)gallium(III)				
	$\Delta_{\text{sub}} H$	(413–433)	90.0		TGA	[2000FAH/BAR]
C₁₅H₃₂GaNS₂	[253595-33-4]	di-tert-butyl gallium dipropyldithiocarbamate				
	$\Delta_v H$	(365–424)	46 ± 1		TGA	[1999KEY/BOT]
C₁₆H₃₆Ga₄S₄	[135283-83-9]	[((CH ₃) ₃ C)Ga(^μ 3-S)] ₄				
	$\Delta_{\text{sub}} H$	(367–380)	110	373	TGA	[1997GIL/BOT]
C₁₆H₃₆Ga₄Se₄	[13528-84-0]	[((CH ₃) ₃ C)Ga(^μ 3-Se)] ₄				
	$\Delta_{\text{sub}} H$	(375–388)	119	381	TGA	[1997GIL/BOT]
C₁₆H₃₆Ga₄Te₄	[135258-40-1]	[((CH ₃) ₃ C)Ga(^μ 3-Te)] ₄				
	$\Delta_{\text{sub}} H$	(391–422)	131	406	TGA	[1997GIL/BOT]
C₂₀H₄₄Ga₄S₄	[166331-96-0]	[(C ₂ H ₅ (CH ₃) ₂ C)Ga(^μ 3-S)] ₄				
	$\Delta_{\text{sub}} H$	(369–382)	124	375	TGA	[1997GIL/BOT]
C₂₀H₄₄Ga₄Se₄	[176100-40-6]	[(C ₂ H ₅ (CH ₃) ₂ C)Ga(^μ 3-Se)] ₄				
	$\Delta_{\text{sub}} H$	(395–407)	137	375	TGA	[1997GIL/BOT]
C₂₀H₄₄Ga₄Te₄	[176100-41-7]	[(C ₂ H ₅ (CH ₃) ₂ C)Ga(^μ 3-Te)] ₄				
	$\Delta_{\text{sub}} H$	(416–432)	140	324	TGA	[1997GIL/BOT]
C₂₄H₅₂Ga₄S₄	[166331-97-1]	[((C ₂ H ₅) ₂ (CH ₃)C)Ga(^μ 3-S)] ₄				
	$\Delta_{\text{sub}} H$	(407–420)	137	413	TGA	[1997GIL/BOT]
C₂₄H₅₂Ga₄Se₄	[187612-49-3]	[((C ₂ H ₅) ₂ (CH ₃)C)Ga(^μ 3-Se)] ₄				
	$\Delta_{\text{sub}} H$	(388–420)	147	404	TGA	[1997GIL/BOT]
C₂₄H₅₂Ga₄Te₄	[176100-42-8]	[((C ₂ H ₅) ₂ (CH ₃)C)Ga(^μ 3-Te)] ₄				
	$\Delta_{\text{sub}} H$	(432–447)	151	439	TGA	[1997GIL/BOT]
C₂₈H₆₀Ga₄S₄	[187612-47-1]	[((C ₂ H ₅) ₃ C)Ga(^μ 3-S)] ₄				
	$\Delta_{\text{sub}} H$	(432–444)	149	438	TGA	[1997GIL/BOT]
C₂₈H₆₀Ga₄Se₄	[187612-51-7]	[((C ₂ H ₅) ₃ C)Ga(^μ 3-Se)] ₄				
	$\Delta_{\text{sub}} H$	(452–464)	156	458	TGA	[1997GIL/BOT]
C₂₈H₆₀Ga₄Te₄	[187612-52-8]	[((C ₂ H ₅) ₃ C)Ga(^μ 3-Te)] ₄				
	$\Delta_{\text{sub}} H$	(444–456)	156	450	TGA	[1997GIL/BOT]

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)			
C₃₃H₅₇O₆Ga	[34228-15-4]	<i>tris</i> (2,2,6,6-tetramethylheptan-3,5-dionato)gallium(III)					
	$\Delta_{\text{sub}}H$	(413–443)	87.0		TGA	[2000FAH/BAR]	
	$\Delta_{\text{sub}}H$		102.1		ME	[1973BRU/CUR]	
GaBr₃	[13450-88-9]	gallium tribromide					
	$\Delta_{\text{sub}}H$	(300–357)	92.5 ± 2.0	298	TE	[2009BRU/PIA]	
(GaBr₃)–(NH₃)	[54955-92-9]	gallium tribromide- ammonia complex					
	$\Delta_{\text{sub}}H$		67.4 ± 1.3			[1975TRU/SUV]	
GaCl₃	[13450-90-3]	gallium trichloride					
	$\Delta_{\text{fus}}H$		11.12	349.6	DSC	[2007CHU/ZEL]	
	$\Delta_{\text{sub}}H$	(289–308)	89 ± 2	298	TE	[2010BRU/PIA]	
	$\Delta_{\text{sub}}H$	(313–349)	87.1 ± 1.2	298	T	[2007CHU/ZEL]	
	Δ_vH	(351–421)	72.7 ± 0.2	349	T	[2007CHU/ZEL]	
(GaCl₃)–(NH₃)	[50599-24-1]	gallium trichloride- ammonia complex					
	$\Delta_{\text{sub}}H$		75.6 ± 1.3			[1975TRU/SUV]	
GaF₃	[7783-51-9]	gallium trifluoride					
	$\Delta_{\text{sub}}H$	(808–958)	252 ± 4	298	TE	[2010BRU/PIA]	
GaI₃	[13450-91-4]	gallium triiodide					
	$\Delta_{\text{sub}}H$	(345–401)	100.5 ± 2.0	298	TE	[2010BRU/PIA]	
Gd (gadolinium)							
C₁₀H₁₀ClGd	[11087-14-2]	<i>bis</i> (cyclopentadienyl)gadolinium chloride					
	$\Delta_{\text{sub}}H$	(338–438)	138.5 ± 2.1		ME	[1971HAU]	
C₁₅H₁₅Gd	[1272-21-5]	<i>tris</i> (cyclopentadienyl)gadolinium					
	$\Delta_{\text{sub}}H$	(513–623)	106.7 ± 2.9			[1973BOR/KRA]	
C₃₀H₃₀F₂₁GdO₆	[17631-67-3]	<i>tris</i> (1,1,1,2,2,3,3-heptafluoro-7,7-dimethyloctane-4,6-dione)gadolinium(III)					
	$\Delta_{\text{sub}}H$	(362–385)	154.8 ± 0.8		ME	[1971SWA/KAR]	
C₃₃H₅₇GdO₆	[14768-15-1]	<i>tris</i> (2,2,6,6-tetramethylpentane-2,4-dionato)gadolinium(III)					
	$\Delta_{\text{sub}}H$		166.1 ± 3.5	298	DSC	[1999SAN/PET]	
	$\Delta_{\text{sub}}H$		78.8 ± 1.5			[1996TSY/DYA2, 2000GIE]	
	$\Delta_{\text{sub}}H$	(383–418)	181.2	400	ME	[1981AMA/SAT]	
	$\Delta_{\text{sub}}H$			163.6	ME	[1973BRU/CUR]	
	$\Delta_{\text{sub}}H$	(420–456)	161.3	438	BG	[1969SIC/DUB]	
	Δ_vH	(456–500)	90.2		BG	[1969SIC/DUB]	
Ge (germanium)							
CHCl₃Ge	[21572-22-5]	trichloro(dichloromethyl)germane					
	Δ_vH	(303–423)	47.9	318		[1975SOK/KAR]	
CH₂Cl₄Ge	[21572-18-9]	trichloro(chloromethyl)germane					
	Δ_vH	(303–423)	45.9	318		[1975SOK/KAR]	
CH₃Cl₃Ge	[993-10-2]	methyltrichlorogermane					
	Δ_vH	(293–385)	37.4	308		[1971GON/KAR]	
CH₄Cl₂Ge	[1111-82-6]	methyldichlorogermane					
	Δ_vH	(281–346)	34.5	313	SG	[1961GRI/ONY]	
	Δ_vH	(273–290)	33.1	281		[1961AMB/BOE]	
CH₅BrGe	[30123-09-2]	methylbromogermane					

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound		T_m (K)	Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)			
	$\Delta_v H$	(273–333)	33.3	303	SG	[1961GRI/ONY]
CH ₅ ClGe	[29914-10-1] $\Delta_v H$	methyl chlorogermane (241–263)	25.8	252		[1961AMB/BOE]
CH ₆ Ge	[1449-65-6] $\Delta_v H$ $\Delta_v H$	methylgermane (159–230) (164–197)	16.6 21.4	194 181	SG	[1961GRI/ONY] [1961AMB/BOE]
CH ₆ GeS	[16643-16-8] $\Delta_v H$	(methylthio)germane (223–291)	29.8	257		[1999DYK/SVO]
CH ₁₂ Ge ₃ Si	[20576-06-1] $\Delta_v H$	trigermymethylsilane (301–378)	39.1	339		[1968DUT/ONY]
C ₂ H ₅ Cl ₃ Ge	[993-42-0] $\Delta_v H$	trichloro(ethyl)germane (293–415)	41.9	308		[1971GON/KAR]
C ₂ H ₇ ClGe	[21961-73-9] $\Delta_v H$	dimethylchlorogermane (273–288)	29.4	280		[1961AMB/BOE]
C ₂ H ₈ Ge	[1449-64-5] $\Delta_v H$	dimethylgermane (196–228)	26.5	212		[1961AMB/BOE]
C ₂ H ₁₀ Ge ₂	[23830-51-5] $\Delta_v H$	1,1-dimethyldigermane (259–295)	31.8	277		[1969GEO/MAC]
C ₂ H ₁₀ Ge ₂	[23830-52-6] $\Delta_v H$	1,2-dimethyldigermane (258–295)	29.3	277		[1969GEO/MAC]
C ₂ H ₁₂ Ge ₂ Si	[23830-52-6] $\Delta_v H$	digermyldimethylsilane (297–381)	34.4	339		[1968DUT/ONY]
C ₃ H ₉ ClGe	[1529-47-1] $\Delta_v H$ $\Delta_v H$	trimethylchlorogermane (293–363) (273–341)	36.3 34.4	308 307	SG	[1972DIT/SKO2] [1961GRI/ONY]
C ₃ H ₉ FGe	[661-37-0] $\Delta_{\text{sub}} H$ $\Delta_v H$	trimethylfluorogermane (250–284) (285–345)	40.0 32.4	267 315	SG SG	[1987STE/MAL, 1961GRI/ONY] [1961GRI/ONY]
C ₃ H ₁₂ GeSi	[18365-18-0] $\Delta_v H$	(trimethylsilyl)germane (288–314)	30.3	301		[1968DUT/ONY]
C ₃ H ₁₂ Ge ₂	[20478-15-3] $\Delta_v H$	1,1,1-trimethyldigermane (273–327)	36.1	300		[1968DUT/ONY]
C ₃ H ₁₂ Ge ₂	[23830-53-7] $\Delta_v H$	1,1,2-trimethyldigermane (268–294)	33.5	281		[1969GEO/MAC]
C ₄ H ₉ Cl ₃ Ge	[4872-26-8] $\Delta_v H$ $\Delta_v H$	butyltrichlorogermanium (313–453) (337–377)	49.2 45.8	328 352		[1975SOK/KAR2] [1972GON/KAR]
C ₄ H ₁₂ Ge	[865-52-1] $\Delta_{\text{fus}} H$ $\Delta_v H$ $\Delta_v H$	tetramethylgermane	7.45 28.1 ± 0.1 27.6 ± 2.1	184.4 285		[1970VAL/KIL] C [1970VAL/KIL] [1969SHA/FED, 1982PIL/SKI]
C ₄ H ₁₂ GeO	[6163-67-3] $\Delta_v H$	trimethylmethoxygermane (273–335)	32.4	304	SG	[1961GRI/ONY]

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound		T_m (K)	Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)			
C ₄ H ₁₂ GeO ₄	[992-91-6]	tetramethoxygermane				
	$\Delta_v H$	(264–303)	48.7	298		[2008PAN/FUL]
	$\Delta_v H$		40.2 ± 0.4			[1970SHA/FED, 1977PED/RYL]
C ₄ H ₁₂ GeS ₄	[21736-70-9]	tetra(methylthia)germane				
	$\Delta_{\text{fus}} H$		14.2	284.1	DSC	[1998FUE/STR]
C ₅ H ₁₁ Cl ₃ Ge	[25425-26-7]	pentyltrichlorogermanium				
	$\Delta_v H$	(323–473)	51.9	338		[1975SOK/KAR2]
C ₆ H ₅ Cl ₃ Ge	[1074-29-9]	phenyltrichlorogermane				
	$\Delta_v H$	(343–473)	55.4	358		[1972SOK/KAR]
C ₆ H ₁₃ Cl ₃ Ge	[35460-93-6]	hexyltrichlorogermanium				
	$\Delta_v H$	(315–491)	51.1	329		[1972GON/KAR]
C ₆ H ₁₅ BrGe	[1067-10-3]	bromotriethylgermane				
	$\Delta_v H$	(303–463)	48.3	318		[1971GON/KAR]
C ₆ H ₁₈ Ge ₂ O	[2237-93-6]	hexamethyldigermoxane				
	$\Delta_v H$	(291–345)	44.1	318	SG	[1961GRI/ONY]
C ₇ H ₇ Cl ₃ Ge	[6181-21-1]	benzyltrichlorogermane				
	$\Delta_v H$	(373–473)	58.8	388		[1972SOK/KAR]
C ₇ H ₁₅ Cl ₃ Ge	[1190-86-9]	heptyltrichlorogermanium				
	$\Delta_v H$	(323–506)	52.3	338		[1972GON/KAR]
C ₈ H ₂₀ Ge	[57596-76-6]	pentyl(trimethyl)germane				
	$\Delta_v H$	(303–423)	44.3	318		[1975SOK/KAR2]
C ₈ H ₂₀ Ge	[597-63-7]	tetraethylgermane				
	$\Delta_{\text{fus}} H$		12.31	180.3		[1996DOM/HEA]
	$\Delta_v H$	(253–293)	43.4	273	GS	[1992GAZ/SCH]
	$\Delta_v H$		45.7 ± 0.4	298	C	[1977PEA/FUC]
	$\Delta_v H$	(337–436)	46.1	352		[1974MOG/HOC]
C ₈ H ₂₀ GeO ₄			44.8 ± 1.3			[1964BIL/COT, 1982PIL/SKI]
	[14165-55-0]	tetraethoxygermane				
	$\Delta_v H$	(259–303)	56.7	298		[2008PAN/FUL]
	$\Delta_v H$		53.9	389		[1988GRI/CHE, 2008PAN/FUL]
	$\Delta_v H$		53.6	406		[1977BAL/RUD, 2008PAN/FUL]
C ₁₀ H ₂₄ GeO ₂			43.1 ± 0.4			[1970SHA/FED, 1977PED/RYL]
			47.9	371		[1958BRA/KAY]
		(328–414)	46.6	403		[1953JOH/FRI, 2008PAN/FUL]
	[26452-74-4]	<i>tert</i> -butylperoxytriethylgermane				
	$\Delta_v H$		43.5 ± 4.2			[1971RAB/KIP, 1982PIL/SKI]
C ₈ H ₂₄ Ge ₄ O ₄	[7749-82-8]	octamethyltetragermoxane				
	$\Delta_{\text{sub}} H$		68.2 ± 4.2	298		[1982PIL/SKI, 1972VOL/SMO]
C ₁₀ H ₂₅ GeN			51.4	403		[1972VOL/SMO]
	[756-66-1]	triethyl(diethylamino)germane				
C ₁₂ H ₁₂ Ge			50.9	318		[1970GON/KAR]
	$\Delta_v H$	(303–463)	46.0 ± 4.8			[1971KOL/RAB, 1982PIL/SKI]
	[1675-58-7]	diphenylgermane				

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound		T_m (K)	Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)			
	$\Delta_{\text{fus}}H$		11.91	240.2		[1980LEB/KIP]
C₁₂H₂₈Ge	[994-65-0]	tetrapropylgermane				
	Δ_vH	(353–493)	54.7	368	A	[1987STE/MAL]
	Δ_vH		61.5 ± 4.2			[1964POP/SKI, 1982PIL/SKI]
C₁₂H₂₈GeO₄	[128426-02-8]	tetrapropoxygermane				
	Δ_vH	(343–453)	63.3	358	A	[1987STE/MAL]
	Δ_vH	(369–465)	55.0	417		[1958BRA/KAY]
	Δ_vH		55.6			[1953JOH/FRI]
C₁₂H₂₈GeO₄	[na]	tetraisopropoxygermane				
	Δ_vH	(313–453)	60.4	328	A	[1987STE/MAL]
	Δ_vH	(355–444)	54.9	400		[1958BRA/KAY]
C₁₂H₃₀ClGeN₃	[na]	<i>tris</i> (diethylamino)chlorogermane				
	Δ_vH	(363–493)	64.4	378		[1970GON/KAR]
C₁₂H₃₀Ge₂Hg	[4149-28-4]	<i>bis</i> (triethylgermyl)mercury				
	Δ_vH	(383–403)	64.8	393		[1972BRA/KAR]
	Δ_vH		62.8 ± 4.2			[1972KOL/RAB, 1982PIL/SKI]
C₁₂H₃₀Ge₂	[993-62-4]	hexaethyldigermane				
	Δ_vH		62.8			[1963RAB/TEL, 1982PIL/SKI]
C₁₂H₃₀Ge₂O	[2538-70-7]	hexaethyldigermoxane				
	Δ_vH		58.6 ± 4.2			[1971RAB/KIP, 1982PIL/SKI]
C₁₆H₁₂Ge	[1675-59-8]	(diethynyl)diphenylgermane				
	$\Delta_{\text{fus}}H$	(8–326)	20.1	320		[1975LEB/MIL]
	$\Delta_{\text{sub}}H$		133.9		BE	[1975LEB/MIL]
	Δ_vH	(305–337)	110.8	320	A	[1987STE/MAL]
C₁₆H₁₈Ge	[na]	1,1-diphenylgermanocyclopentane				
	$\Delta_{\text{fus}}H$		14.45		DSC	[1988CAR/DYS]
	$\Delta_{\text{sub}}H$		104.6 ± 2.8	298		[1988CAR/DYS]
	Δ_vH	(294–322)	87.6 ± 2.8		ME	[1988CAR/DYS]
C₁₆H₃₆Ge	[1067-42-1]	tetrabutylgermane				
	$\Delta_{\text{fus}}H$		19.1	198.6		[1971SHA/YAK]
C₁₆H₃₆GeO₄	[25063-27-8]	tetrabutoxygermane				
	Δ_vH	(394–519)	62.4	456		[1958BRA/KAY]
	Δ_vH		59.6			[1953JOH/FRI]
C₁₆H₃₆GeO₄	[1085941-13-4]	tetraisobutoxygermane				
	Δ_vH	(369–482)	59.9	426		[1958BRA/KAY]
C₁₆H₃₆GeO₄	[na]	tetra- <i>sec</i> -butoxygermane				
	Δ_vH	(365–475)	59.9	420		[1958BRA/KAY]
C₁₆H₃₆GeO₄	[1085941-54-3]	tetra- <i>tert</i> -butoxygermane				
	Δ_vH	(364–460)	53.8	412		[1958BRA/KAY]
C₁₈HF₁₅Ge	[42371-50-6]	<i>tris</i> (pentafluorophenyl)germane				
	$\Delta_{\text{fus}}H$	(7–500)	34.9	405	AC	[1997SMI/LEB2]
C₁₈H₄₂Ge₂Hg	[24004-54-4]	<i>bis</i> (triisopropylgermyl)mercury				
	Δ_vH	(373–483)	68.7	388		[1972BRA/KAR]

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)			
	$\Delta_v H$		54.4 ± 4.2				[1972KOL/RAB, 1982PIL/SKI]
C₂₀H₁₈Ge	[4049-97-2]	triphenyl vinylgermanium					
	$\Delta_{\text{sub}} H$		98.7 ± 1.6	298	ME,TE		[1988CAR/JAM2]
C₂₄H₂₀Ge	[1048-05-1]	tetraphenylgermane					
	$\Delta_{\text{sub}} H$	(402–480)	148.6	441	A		[1987STE/MAL]
	$\Delta_{\text{sub}} H$		156.9 ± 4.2	298			[1982PIL/SKI, 1969ADA/CAR]
C₂₄H₂₀GeO₄	[1085941-60-1]	tetraphenoxygermane					
	$\Delta_v H$		37.4 ± 0.4				[1970SHA/FED, 1977PED/RYL]
C₂₆H₂₀Ge	[4131-49-1]	triphenyl phenylethynylgermane					
	$\Delta_{\text{sub}} H$		107.5 ± 1.5	298	ME,TE		[1988CAR/JAM2]
C₂₈H₂₈Ge	[1048-05-1]	tetrabenzylgermane					
	$\Delta_{\text{sub}} H$		168.6 ± 8.4	298			[1982PIL/SKI, 1970CAR/CAR]
C₃₂H₁₆Cl₂GeN₈	[19566-97-3]	dichlorophthalocyaninatogermane					
	$\Delta_{\text{sub}} H$			147.4			[1972MAR/LOP]
C₃₆H₃₀Ge₂O	[2181-40-0]	bis(triphenyl germanium) oxide					
	$\Delta_{\text{sub}} H$		98.0 ± 1.5	298	ME,TE		[1988CAR/JAM2]
C₃₆H₃₀Ge₂	[2816-39-9]	hexaphenyldigermane					
	$\Delta_{\text{sub}} H$		151.3				[1972MAR/LOP]
	$\Delta_{\text{sub}} H$		209.2 ± 4.2	298			[1982PIL/SKI, 1970CAR/CAR]
GeBr₂	[24415-00-7]	germanium dibromide					
	$\Delta_{\text{fus}} H$		11.6	409.2	DSC		[2006ZEL/CHU2]
GeBr₄	[13450-92-5]	germanium tetrabromide					
	$\Delta_{\text{fus}} H$	(5–315)	12.85	299.3	AC		[1999BER/ZEL]
	$\Delta_{\text{sub}} H$	(273–299)	58.6 ± 1.2	298			[2004ZEL/CHU]
	$\Delta_v H$	(299–373)	46.6 ± 0.3	298			[2004ZEL/CHU]
GeCl₄	[10038-98-9]	germanium tetrachloride					
	$\Delta_{\text{fus}} H$		8.52	221.7			[1986DEV/GUS]
	$\Delta_{\text{sub}} H$	(187–221)	44.6 ± 0.2		MG		[1964BAL/DON]
GeF₂	[13940-63-1]	germanium difluoride					
	$\Delta_{\text{fus}} H$		9.25	(corrected to 298 K)			[1971ADA/MAR]
	$\Delta_{\text{sub}} H$		82.8 ± 4.2	298	MS		[1971ADA/MAR]
	$\Delta_{\text{sub}} H$		93.3 ± 10.5	298			[1971ADA/MAR]
GeI₂	[13573-08-5]	germanium diiodide					
	$\Delta_{\text{fus}} H$		33.3	701.2			[2003ZEL/TIT]
GeI₄	[13573-08-5]	germanium tetraiodide					
	$\Delta_{\text{fus}} H$		19.1	419			[1998ZEL/MIN]
	$\Delta_{\text{sub}} H$		87.1 ± 3	298			[1999TIT/ZEL]
	$\Delta_{\text{sub}} H$		86.7 ± 3	298			[1999TIT/ZEL]
	$\Delta_{\text{sub}} H$	(323–420)	76.5 ± 5.7	298	TE		[1987FER/STR]
	$\Delta_v H$	(419–613)	64.2 ± 0.2	419			[1999TIT/ZEL2]
Ge₄H₁₀	[14691-47-5]	tetragermane					

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No	Compound		T_m (K)	Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)			
	$\Delta_v H$		32.8			[1959AMB]
Ge₅H₁₂	[15587-39-0]	pentagermane				
	$\Delta_v H$		34.6			[1959AMB]
GeH₆Si	[13768-63-3]	germylsilane				
	$\Delta_v H$	(190–250)	25.0	220		[1963SPA/MAC]
Ha (hahnium)						
HaCl₅	[146837-09-4]	hahnium(V) pentachloride				
	$\Delta_{\text{sub}} H$		<120	298		[1996TUR/EIC]
HaOCl₃	[143928-41-0]	hahnium(V) oxychloride				
	$\Delta_{\text{sub}} H$	(298–607)	152 ± 18	298		[1996TUR/EIC]
Hf (hafnium)						
C₁₀H₁₀Cl₂Hf	[12116-66-4]	<i>bis</i> (cyclopentadienyl)hafnium dichloride				
	$\Delta_{\text{sub}} H$		110.2 ± 2.9	298	ME	[2001DIO/PIE]
	$\Delta_{\text{sub}} H$	(394–447)	100.3	420.5	A	[1987STE/MAL]
	$\Delta_{\text{sub}} H$		106.7 ± 2.1	298		[1982PIL/SKI, 1976KIR/TEL]
	$\Delta_{\text{sub}} H$		100.4 ± 1.3			[1977BAL/BAR]
	$\Delta_{\text{sub}} H$		107.3 ± 2.4	298		[1968KIS/DIL, 2001DIO/PIE]
C₁₂H₁₆Hf	[37260-88-1]	<i>bis</i> (cyclopentadienyl)dimethyl hafnium				
	$\Delta_{\text{sub}} H$	(295–316)	81.1 ± 1.9	303	ME	[2008MOR/ZHE]
C₁₈H₃₀HfN₂	[159338-62-2]	<i>bis</i> (cyclopentadienyl)hafnium <i>bis</i> (diethylamide)				
	$\Delta_{\text{sub}} H$	(328–365)	130.5 ± 1.0	346	ME	[2008MOR/ZHE]
C₂₀H₁₆F₁₂HfO₈	[17475-68-2]	<i>tetrakis</i> (1,1,1-trifluoro-2,4-pentanedionato)hafnium(IV)				
	$\Delta_{\text{fus}} H$		40.1	398	DSC	[2008ZHE/MOR]
	$\Delta_{\text{sub}} H$	(358–393)	133.0 ± 1.8	376	GS	[2008MOR/ZHE]
	$\Delta_{\text{sub}} H$	(358–398)	126.5 ± 1.8			[2008ZHE/MOR]
	$\Delta_{\text{sub}} H$	(383–438)	129.7 ± 3.8		GS	[1985MAT/KUW]
	$\Delta_{\text{sub}} H$	(383–438)	124.7 ± 3.8		GS	[1985MAT/KUW]
	$\Delta_v H$	(403–423)	84.7 ± 3.1	413	GS	[2008MOR/ZHE]
	$\Delta_v H$	(403–423)	83.2 ± 2.0	413		[2008ZHE/MOR]
C₂₀H₂₈HfO₈	[17475-67-1]	<i>tetrakis</i> (pentane-2,4-dionato)hafnium(IV)				
	$\Delta_{\text{sub}} H$	(408–433)	138.7 ± 7.4	420	GS	[2008MOR/ZHE]
	$\Delta_{\text{sub}} H$	(408–443)	130.4 ± 6.1	425		[2008ZHE/MOR]
	$\Delta_{\text{sub}} H$		150.6 ± 4.2			[1991TEL/LAR]
C₃₂H₄₀F₁₂HfO₈	[916441-69-5]	<i>tetrakis</i> (1,1,1-trifluoro-5,5-dimethyl-2,4-hexanedionato)hafnium(IV)				
	$\Delta_{\text{sub}} H$	(383–423)	135.3 ± 1.7	403	GS	[2008MOR/ZHE]
		Note: the enthalpy of sublimation that was reported in the paper, 97.5 ± 1.7 kJ/mole, was not consistent with the reported Antoine constants. We have recalculated the the enthalpy of sublimation assuming that the reported Antoine constants are correct.				
	$\Delta_{\text{sub}} H$	(386–423)	121.5 ± 0.8	404		[2008ZHE/MOR]
	$\Delta_v H$	(426–493)	92.1 ± 1.3	460	GS	[2008MOR/ZHE]
	$\Delta_v H$	(424–472)	91.2 ± 0.3	448		[2008ZHE/MOR]
C₄₄H₇₆HfO₈	[63370-90-1]	<i>tetrakis</i> (2,2,6,6-tetramethyl-3,5-heptanedionaot)hafnium(IV)				
	$\Delta_{\text{tr}} H$		15.8	446		
		Note: The enthalpy includes a solid/solid transition that occurs at 433 K				
	$\Delta_{\text{fus}} H$		5.4	630	DSC	[2008ZHE/MOR]

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{sub}}H$	(453–553)	103.5 ± 0.6	503	GS	[2008MOR/ZHE]
	$\Delta_{\text{sub}}H$	(368–428)	136.6 ± 4.2	398	ME	[2008MOR/ZHE]
HfCl₄	[13499-05-3]	hafnium tetrachloride				
	$\Delta_{\text{sub}}H$	(398–500)	97.9 ± 1.2	499	T	[1994TAN/BOS]
	$\Delta_{\text{sub}}H$	(353–433)	107.9 ± 0.8			[1973IZM/KHO]
Hg (mercury)						
CH₃BrHg	[506-83-2]	methylmercuric bromide				
	$\Delta_{\text{sub}}H$	(258–297)	67.6 ± 1.6	277.5	V	[1987STE/MAL, 1951CHA/SKI]
CH₃ClHg	[115-09-3]	methylmercuric chloride				
	$\Delta_{\text{sub}}H$	(278–307)	64.9 ± 1.6	298	V	[1987STE/MAL, 1982PIL/SKI, 1950HAR/PRI, 1951CHA/SKI]
CH₃HgI	[143-36-2]	methylmercuric iodide				
	$\Delta_{\text{sub}}H$	(263–290)	65.3 ± 1.6	276	V	[1951CHA/SKI]
C₂H₅BrHg	[107-26-6]	ethylmercuric bromide				
	$\Delta_{\text{sub}}H$	(285–303)	76.5 ± 2.9	294	V	[1987STE/MAL, 1982PIL/SKI, 1951HAR/PRI, 1951CHA/SKI]
C₂H₅ClHg	[107-27-7]	ethylmercuric chloride				
	$\Delta_{\text{sub}}H$	(283–303)	76.2 ± 2.9	293	V	[1987STE/MAL, 1982PIL/SKI, 1951HAR/PRI, 1951CHA/SKI]
C₂H₅HgI	[2440-42-8]	ethylmercuric iodide				
	$\Delta_{\text{sub}}H$	(286–303)	79.7 ± 2.9	294.5	V	[1987STE/MAL, 1982PIL/SKI, 1951HAR/PRI, 1951CHA/SKI]
C₂H₆Hg	[593-74-8]	dimethyl mercury				
	Δ_vH	(275–367)	36.7 ± 0.1	321		[2001BAE]
	Δ_vH		34.6 ± 0.8			[1950HAR/PRI, 1982PIL/SKI]
C₂F₆HgS₂	[1085746-33-3]	<i>bis</i> (trifluoromethylthio)mercury				
	Δ_vH	(353–423)	49.9	368		[1999DYK/SVO]
C₄H₁₀Hg	[627-44-1]	diethyl mercury				
	Δ_vH		44.8 ± 1.7			[1951HAR/PRI, 1982PIL/SKI]
	Δ_vH		44.9			[1935JON/EVA]
C₄H₁₆Cl₂HgN₈S₄	[28813-22-1]	<i>trans</i> -dichloro- <i>tetrakis</i> (thiourea)mercury(II)				
	$\Delta_{\text{sub}}H$		101 ± 20			[1970ASH]
C₆H₁₄Hg	[628-85-3]	dipropyl mercury				
	Δ_vH		55.2 ± 1.3			[1952MOR/PRI, 1982PIL/SKI]
C₆H₁₄Hg	[1071-39-2]	diisopropyl mercury				
	Δ_vH		53.6 ± 1.7			[1952MOR/PRI, 1982PIL/SKI]
C₁₀H₁₄Cl₂HgN₆O₂	[na]	[mercury(1-methylcytosine) ₂ Cl ₂]				
	$\Delta_{\text{sub}}H$	(428–443)	150.8 ± 19	435	ME	[1984BUR/MOR]
	$\Delta_{\text{sub}}H$		159 ± 19	298		[1984BUR/MOR]
C₁₀H₂₀HgN₂S₄	[14239-51-1]	<i>bis</i> (diethyldithiocarbamate) mercury complex				
	$\Delta_{\text{sub}}H$	(378–403)	47.6	390.5		[1987STE/MAL]
C₁₂H₁₂Hg	[587-85-9]	diphenylmercury				
	$\Delta_{\text{sub}}H$	(314–303)	112.8 ± 0.8	298	ME	[1958CAR/STR]
C₁₂H₃₀Ge₂Hg	[4149-28-4]	<i>bis</i> (triethylgermyl)mercury				
	Δ_vH	(383–403)	64.8	393		[1972BRA/KAR]
C₁₂H₃₀HgSi₂	[4149-29-5]	<i>bis</i> (triethylsilyl)mercury				

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound			Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)		
	$\Delta_v H$	(383–433)	64.0	398		[1972BRA/KAR]
C₁₄H₁₄Hg	[780-24-5] $\Delta_{\text{sub}} H$	<i>bis</i> (benzyl)mercury (350–390)	88.7 ± 2.1		ME, TE	[1984CAR/SPE]
C₁₄H₂₈HgN₂S₄	[21439-56-5] $\Delta_{\text{sub}} H$	<i>bis</i> (dipropyldithiocarbamate)mercury(II)	200 ± 2	298	DSC, E	[1992DEC/AIR]
C₁₆H₁₀Hg	[6077-10-7] $\Delta_{\text{sub}} H$	<i>bis</i> (phenylethynyl)mercury (350–390)	99.2 ± 1.4		ME, TE	[1984CAR/SPE]
C₁₈H₃₆HgN₂S₄	[21439-58-7] $\Delta_{\text{sub}} H$	<i>bis</i> (dibutyldithiocarbamate)mercury(II)	193 ± 3	298	DSC,E	[1991DES/DES]
C₁₈H₃₆HgN₂S₄	[79001-48-2] $\Delta_{\text{sub}} H$	<i>bis</i> (diisobutyldithiocarbamate)mercury(II)	247 ± 1	298	DSC,E	[1994SOU/PIN]
C₁₈H₄₂Ge₂Hg	[24004-54-4] $\Delta_v H$	<i>bis</i> (triisopropylgermyl)mercury (373–483)	68.7	388		[1972BRA/KAR]
HgF₂	[7783-39-3] $\Delta_{\text{sub}} H$	mercuric fluoride (496–629)	136 ± 4	298		[2008BRU/PIA]
HgI₂	[7774-29-0] $\Delta_{\text{fus}} H$	mercuric iodide	20.3	537	Sub-Vap	[2002SU/ZHU]
	$\Delta_{\text{sub}} H$		84.4		UV	[2002SU/ZHU]
	$\Delta_v H$	(537–610)	64.0	574	UV	[2002SU/ZHU]
Ho (holmium)						
C₁₅H₁₅Ho	[1272-22-6] $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$	<i>tris</i> (cyclopentadienyl)holmium(III) (338–348)	102.1 ± 2.1 119.7 ± 2.1			[1973DEV/BOR] ME [1971HAU, 1971HAU2]
C₃₃H₅₇HoO₆	[15522-73-3] $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$	<i>tris</i> (2,2,6,6-tetramethylheptane-3,5-dionato)holmium(III) (363–418) (420–458)	131.0 ± 2.9 152.7 131.4	390 439	DSC ME BG	[1993AIR/SAN] [1981AMA/SAT] [1969SIC/DUB]
	$\Delta_v H$	(458–500)	84.6		BG	[1969SIC/DUB]
I (iodine)						
HI	[10034-85-2] $\Delta_v H$	hydrogen iodide	19.8	238	C	[1929GIA/WIE]
In (indium)						
C₃H₉In	[3385-78-2] $\Delta_{\text{fus}} H$ $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$	trimethyl indium (274–313) (328–362)	14.3 62.7 48.5 ± 2.5 57.7	358.7 294 298 344		[1991URY/RON] [2004FUL/RUZ] [1982PIL/SKI, 1968CLA/PRI] [1987STE/MAL, 1941LAU/GIL]
C₆H₁₅In	[923-34-2] $\Delta_{\text{fus}} H$ $\Delta_{\text{fus}} H$ $\Delta_v H$	triethyl indium (326–376)	13.09 13.01 45.0 ± 0.7	237.6 237.6 351		[1996DOM/HEA] [1996DOM/HEA] [2001BAE, 2001BAE/CHE]

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)			
C₉H₂₁In	[3015-98-3] $\Delta_v H$	tripropyl indium (400–483)	52.0	441		[1999DYK/SVO]	
C₉H₂₁In	[17144-80-8] $\Delta_v H$ $\Delta_v H$	triisopropyl indium (318–366) (394–478)	52.3 ± 0.7 51.0	342 336		[2001BAE] [1999DYK/SVO]	
C₁₂H₂₇In	[15676-66-1] $\Delta_v H$	tributyl indium (444–539)	58.5	459	A	[1987STE/MAL]	
C₁₅H₁₂F₉InO₆	[15453-87-9] $\Delta_{\text{sub}} H$ $\Delta_v H$	<i>tris</i> (1,1,1-trifluoro-2,4-pentanedionato)indium(III) (378–428) (398–478)	112.1 ± 1.3 77.4 ± 0.6	438	GS	[1985MAT/KUW] [1978IGU/CHU2]	
C₁₅H₃₀InN₃S₆	[15741-07-8] $\Delta_{\text{sub}} H$	<i>tris</i> (diethyldithiocarbamate)indium(III) 176.7 ± 3.3	298		DSC,E	[2000SOU/OLI]	
C₂₀H₄₈In₂P₄	[115381-28-7] $\Delta_{\text{sub}} H$	<i>bis</i> [μ -[<i>bis</i> (1,1-dimethylethyl)phosphino]]tetramethyldiindium(III) 130.0			ME	[1988BRA/FAK]	
C₂₁H₄₂InN₃S₆	[87052-01-5] $\Delta_{\text{sub}} H$	<i>tris</i> (dipropyldithiocarbamate)indium(III) 372.8 ± 3.4	298		DSC,E	[2000SOU/OLI]	
C₂₁H₄₂InN₃S₆	[85883-33-6] $\Delta_{\text{sub}} H$	<i>tris</i> (diisopropyldithiocarbamate)indium(III) 279.5 ± 3.5	298		DSC,E	[2000SOU/OLI]	
C₂₇H₅₄InN₃S₆	[85883-33-6] $\Delta_{\text{sub}} H$	<i>tris</i> (dibutyldithiocarbamate)indium(III) 358.3 ± 3.2	298		DSC,E	[2000SOU/OLI]	
C₂₇H₅₄InN₃S₆	[85129-27-7] $\Delta_{\text{sub}} H$	<i>tris</i> (diisobutyldithiocarbamate)indium(III) 182.0 ± 3.3	298		DSC,E	[2000SOU/OLI]	
C₃₃H₅₇InO₆	[34269-03-9] $\Delta_{\text{sub}} H$	<i>tris</i> (2,2,6,6-tetramethylheptan-3,5-dionato)indium(III) 129.3			ME	[1973BRU/CUR]	
InBr₃	[13465-09-3] $\Delta_{\text{sub}} H$	indium(III) bromide 147 ± 4	298		TE	[1997BRU/PAL]	
InCl₃	[10025-82-8] $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$	indium(III) chloride (495–648) 152 ± 4 158 ± 4 150.4 161.1 (453–572) 151.1 ± 1.2 155.6 ± 1.2 (478–563) 161.1 ± 1.6 168.5 ± 1.6 (623–773) 156.3 166.6	570 298 710 298 489 298 524 298 698 298		TE MS	[1998BRU/PIA] [1998BRU/PIA] [1994OPP/KRA] [1994OPP/KRA, 1998BRU/PIA] [1988DEF/CHA] [1988DEF/CHA, 1998BRU/PIA] [1988DEF/CHA] [1988DEF/CHA, 1998BRU/PIA] [1974KUN/HOS] [1974KUN/HOS, 1998BRU/PIA]	
InI₃	[13510-35-5] $\Delta_{\text{sub}} H$	indium(III) iodide (399–479)	136 ± 5.0	298	TE,ME	[1997BRU/GIU]	
Ir (iridium)							
C₇H₇IrO₄	[14023-80-4] $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$	dicarbonyl-2,4-pentanedionato iridium complex (306–333) (286–325)	94.1 ± 2.7 92.0 ± 1.3	306	ME ME	[2009MOR/SEM] [1978JES/ERN, 1987STE/MAL]	
C₇H₁₃Cl₂IrO₂	$\Delta_{\text{sub}} H$	<i>bis</i> (chloroethylene)-2,4-pentanedionato iridium complex (281–298)	89.5 ± 4.2	290	ME	[1978JES/ERN, 1987STE/MAL]	

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
C₉H₁₅IrO₂	[52654-27-0] $\Delta_{\text{sub}}H$	<i>bis</i> (ethylene)-2,4-pentanedionato iridium complex (283–311)	82.8 ± 4.2	297	ME	[1978JES/ERN, 1987STE/MAL]
C₁₁H₁₉IrO₂	[66467-05-8] $\Delta_{\text{sub}}H$	<i>bis</i> (propylene)-2,4-pentanedionato iridium complex (269–304)	90 ± 1.3	287	ME	[1978JES/ERN]
C₁₂O₁₂Ir₄	[11065-24-0] $\Delta_{\text{sub}}H$	tetrairidiumdodecacarbonyl 104.6 ± 20		298		[1982PIL/SKI, 1974CON/SKI]
C₁₂H₁₅IrO₂	[32660-96-1] $\Delta_{\text{sub}}H$	(pentamethylcyclopentadienyl)(dicarbonyl)iridium (I) (297–332)	105.0 ± 3.4		ME	[2009MOR/SEM]
C₁₃H₁₉IrO₂	[12154-84-6] $\Delta_{\text{sub}}H$	(acetylacetonato)(1,5-cyclooctadiene)iridium (I) (335–370)	111.7 ± 1.7		ME	[2009MOR/SEM]
C₁₃H₁₉IrO₆	[66467-07-0] $\Delta_{\text{sub}}H$	<i>bis</i> (vinyl acetate)-2,4-pentanedionato iridium complex (325–344)	120.5 ± 2.9	333	ME	[1978JES/ERN]
C₁₃H₁₉IrO₆	[66467-08-1] $\Delta_{\text{sub}}H$	<i>bis</i> (methyl acrylate)-2,4-pentanedionato iridium complex (311–335)	117.2 ± 5	323	ME	[1978JES/ERN]
C₁₄H₁₉Ir	[122644-88-3] $\Delta_{\text{sub}}H$ $\Delta_{\text{v}}H$	(methylcyclopentadienyl)(1,5-cyclooctadiene)iridium (I) (304–310) (310–330)	124.6 ± 5.0 88.1 ± 1.3		ME ME	[2009MOR/SEM] [2009MOR/SEM]
C₁₅H₂₁IrO₆	[15635-87-7] $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$	<i>tris</i> (2,4-pentanedionato)iridium(III) 128 (423–473) (383–433) (387–513) (468–518)	129.3 ± 0.8 130.5 ± 3.4 101.6 ± 1.8 86.6 ± 1.7 NA		GS ME MCV SMZG	[2001MOR/ZHA] [2000MOR/SEM] [2000MOR/SEM] [2000MOR/SEM] [2000MOR/SEM] [1994GER/GER]
La (lanthanum)						
C₁₅H₁₅La	[1272-23-7] $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$	<i>tris</i> (cyclopentadienyl)lanthanum (548–663)	114.6 ± 4.0 102.1 ± 2.9	298		[1982PIL/SKI, 1974DEV/RAB] [1973BOR/KRA]
C₃₀H₃₀F₂₁LaO₆	[19106-89-9] $\Delta_{\text{sub}}H$	<i>tris</i> (1,1,1,2,2,3,3-heptafluoro-7,7-dimethyloctane-4,6-dione)lanthanum(III) (387–403)	145.2 ± 2.9		ME	[1971SWA/KAR]
C₃₃H₅₇LaO₆	[14319-13-2] $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$	<i>tris</i> (2,2,6,6-tetramethylheptan-3,5-dionato)lanthanum(III) (388–423) (450–520)	156.0 ± 4.6 116.1 ± 8.4 107.9 ± 4.6 179.5 164.4 143.6	405 485	DSC ME ME BG	[1997SAN/ROC, 2000GIE] [1996TSY/DYA] [1996TSY/DYA2, 2000GIE] [1981AMA/SAT] [1973BRU/CUR] [1969SIC/DUB]
Li (lithium)						
C₂H₅Li	[811-49-4] $\Delta_{\text{sub}}H$	ethyl lithium (298–333)	116.6	315.5	A	[1987STE/MAL, 1962CHA]
C₄H₉Li	[109-72-8] $\Delta_{\text{sub}}H$ $\Delta_{\text{v}}H$	butyl lithium (333–368)	109.7 107.1 ± 2.9	350.5	A	[1987STE/MAL, 1962LEB/MIR] [1961FOW/MOR, 1982PIL/SKI]

Note: Authors of Ref. [1961FOW/MOR] noted that the experimental data was not very reproducible, and subject to considerable error.

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C₅HF₆LiO₂	[22466-51-9]	(1,1,1,5,5,5-hexafluoro-2,4-pentanedionato)lithium				
	$\Delta_{\text{sub}}H$	(453–523)	149.5 ± 3.5			[2006FIL/SYS]
Note: Above value calculated assuming the vapor phase is a dimer						
C₅H₄F₃LiO₂	[127892-64-2]	(1,1,1-trifluoro-2,4-pentanedionato)lithium				
	$\Delta_{\text{sub}}H$	(453–523)	145.6 ± 3.0			[2006FIL/SYS]
Note: Above value calculated assuming the vapor phase is a dimer						
C₈H₁₀F₃LiO₂	[22441-09-4]	(2,2-dimethyl-6,6,6-trifluoro-3,5-hexanedionato)lithium				
	$\Delta_{\text{sub}}H$	(453–523)	182.0 ± 2.3			[2006FIL/SYS]
Note: Above value calculated assuming the vapor phase is a dimer						
C₁₁H₁₉LiO₂	[22441-13-0]	(2,2,6,6-tetramethylheptane-3,5-dionato)lithium				
	$\Delta_{\text{sub}}H$	(400–450)	191.2 ± 7.1		ME,MS	[2006FIL/STA]
	$\Delta_{\text{sub}}H$	(444–543)	181.1 ± 2.8			[2006FIL/SYS]
	$\Delta_{\text{sub}}H$	(537–545)	198.0 ± 15.1			[2006FIL/SYS]
	$\Delta_{\text{sub}}H$	(444–549)	178.3 ± 2.0			[2006FIL/SYS]
	$\Delta_{\text{sub}}H$		174.5		ME	[1973BRU/CUR]
Note: Above four values calculated assuming the vapor phase is a tetramer						
	$\Delta_v H$	(550–581)	95.6 ± 1.3			[2006FIL/SYS]
Note: Above value calculated assuming the vapor phase is a tetramer						
LiF	[7789-24-4]	lithium fluoride				
	$\Delta_{\text{sub}}H$	(1073–1121)	268.2 ± 4.2			[1959SCH/MAR, 1958EIS/ROT]
	$\Delta_{\text{sub}}H$	(957–1113)	267.8 ± 4.2			[1958POR/SCH]
Lu (lutetium)						
C₁₅H₁₅Lu	[1272-24-8]	<i>tris</i> (cyclopentadienyl)lutetium(III)				
	$\Delta_{\text{sub}}H$		123.0 ± 2.9			[1973DEV/BOR]
C₁₅H₂₁LuO₆	[17966-84-6]	<i>tris</i> (2,4-pentanedionato)lutetium(III)				
	$\Delta_{\text{sub}}H$	(403–433)	79 ± 13	418		[1983TRE/BER]
C₃₃H₅₇LuO₆	[15497-45-2]	<i>tris</i> (2,2,6,6-tetramethylpentane-2,4-dionato)lutetium(III)				
	$\Delta_{\text{sub}}H$		135.8 ± 2.9	298	DSC	[1999SAN/PET]
	$\Delta_{\text{sub}}H$	(363–413)	154.8	390	ME	[1981AMA/SAT]
	$\Delta_{\text{sub}}H$	(420–448)	134.2	434	BG	[1969SIC/DUB]
	$\Delta_v H$	(448–490)	83.6		BG	[1969SIC/DUB]
Mg (magnesium)						
C₁₀H₁₀Mg	[1284-72-6]	<i>bis</i> (cyclopentadienyl) magnesium				
	$\Delta_{\text{sub}}H$		68.2 ± 1.3	298		[1982PIL/SKI, 1967HUL/REI, 1967TUR]
C₁₀H₂₂Mg	[19978-31-5]	<i>bis</i> (2,2-dimethylpropyl)magnesium				
	$\Delta_{\text{sub}}H$	(318–348)	160.0 ± 2.0	333	ME	[1983AKK/SCH]
C₁₈H₁₂MgN₂O₂	[14639-28-2]	<i>bis</i> (8-hydroxyquinolino)magnesium(II)				
	$\Delta_{\text{sub}}H$		230.2 ± 4.0	298	ME	[1994RIB/MAT]
C₂₀H₁₆MgN₂O₂	[14406-92-9]	<i>bis</i> (8-hydroxy-2-methylquinolino)magnesium(II)				
	$\Delta_{\text{sub}}H$	(533–549)	212.2 ± 6.5	541	ME	[1998RIB/MAT3]
	$\Delta_{\text{sub}}H$	(533–549)	224.3 ± 6.5	298	ME	[1998RIB/MAT3]
C₂₈H₅₄MgN₂O₄	[302351-10-6]	(N,N,N',N'-tetramethylethylenediamine) <i>bis</i> ((2,2,6,6-tetramethyl-3,5-heptanedionato)magnesium				
	$\Delta_{\text{fus}}H$		58.3 ± 5.2		DTA	[2008MAR/SEL]
	$\Delta_{\text{sub}}H$		83.2 ± 2.3		TGA	[2008MAR/SEL]

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)			
C₃₁H₆₁MgN₃	$\Delta_{\text{fus}}H$ $\Delta_{\text{sub}}H$ Δ_vH	(N,N,N',N',N''-pentamethyldiethylenetriamine) bis(2,2,6,6-tetramethyl-3,5-heptanedionato)magnesium				DTA	[2009MAR/SEL]
			19.0	352.7		Fus+ Vap	[2009MAR/SEL]
		(373–441)	59 ± 1		TGA	[2009MAR/SEL]	
C₄₄H₇₆Mg₂O₈	[236095-55-9] Δ_vH	tetrakis(2,2,6,6-tetramethyl-3,5-heptanedionato)dimagnesium				TGA	[2009MAR/SEL]
		(395–476)	67 ± 2				
MgF₂	[7783-40-6] $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$	magnesium fluoride					
		(1220–1450)	359.8	1330	MS	[1962BER/MAR]	
		(1273–1513)	327.3 ± 4.3	1400	TE	[1964GRE/KO]	
			348.2 ± 4.3	298		[1964GRE/KO]	
Mn (manganese)							
C₄H₁₆Cl₂MnN₈S₄	[28813-17-4] $\Delta_{\text{sub}}H$	trans-dichloro-tetrakis(thiourea)manganese(II)					
		(382–409)		133 ± 20		[1970ASH]	
C₅BrMnO₅	[14516-54-2] $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$	bromo(pentacarbonyl)manganese					
			58.6 ± 8.4	298		[1982PIL/SKI, 1972CON/SKI]	
			88.0 ± 2.0	298	C	[1982CON/ZAF]	
C₅ClMnO₅	[14100-30-2] $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$	chloro(pentacarbonyl)manganese					
			58.6 ± 8.4	298		[1982PIL/SKI, 1972CON/SKI]	
			91 ± 9			[1982CON/ZAF]	
C₅IMnO₅	[14879-42-6] $\Delta_{\text{sub}}H$	iodo(pentacarbonyl)manganese					
			77.4 ± 1.4	298	C	[1982CON/ZAF]	
C₅H₃MnO₅Si	[15770-61-3] Δ_vH	silyl pentacarbonyl manganese					
		(294–391)	39.6	343	T	[1969AYL/CAM2, 1967AYL/CAM]	
C₆F₃MnO₅	[13601-14-4] $\Delta_{\text{sub}}H$	pentacarbonyl(trifluoromethyl)manganese					
			77.8 ± 1.0	298	C	[1982CON/ZAF]	
C₆H₃MnO₅	[13601-24-6] $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$	methyl(pentacarbonyl)manganese					
			60.3 ± 1.0			[1982PIL/SKI, 1974BRO/CON]	
		(293–403)	60.2			[1958HIE/WAG]	
C₇F₃MnO₆	[14099-62-8] $\Delta_{\text{sub}}H$	pentacarbonyl(trifluoroacetyl)manganese					
			79 ± 5	298	C	[1982CON/ZAF]	
C₇H₃MnO₆	[13963-91-2] $\Delta_{\text{sub}}H$	acetyl(pentacarbonyl)manganese					
			80 ± 7	298	C	[1982CON/ZAF]	
C₈H₅MnO₃	[12079-65-1] $\Delta_{\text{fus}}H$ $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$ Δ_vH	cyclopentadienyl(tricarbonyl)manganese				DSC	[2008PIC/CAN]
			18.9	350.1			
			72.2 ± 3.9	294	ME	[2008PIO/CAN]	
			72.0 ± 3.9	298	ME	[2008PIO/CAN]	
			75.8 ± 0.4	305	C	[2008PIO/CAN]	
			76.0 ± 0.4	298	C	[2008PIO/CAN]	
		(323–353)	52.7 ± 3.1	338		[1982PIL/SKI, 1965EVS/SHM, 1970BAE/DEM]	
		(335–343)	64.2 ± 11.6	339	ME	[1959COR/SCH, 2008PIO/CAN]	
C₈H₁₀Cl₂MnN₆O₂	[74543-44-5] $\Delta_{\text{sub}}H$	[manganese-(cytosine) ₂ Cl ₂]					
		(433–453)	U 146 ± 21	443	ME	[1984BUR/MOR]	

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
C₁₀MnO₁₀Re	[14693-30-2]	decacarbonylmanganeserhenium				
	$\Delta_{\text{sub}}H$		109 ± 4	363	C	[1998ADD/CON]
	$\Delta_{\text{sub}}H$		86 ± 4	298		[1998ADD/CON]
	$\Delta_{\text{sub}}H$	(363–440)	68.6	401	MM	[1971BAE/DEM]
	$\Delta_{\text{v}}H$	(440–463)	56.5	451		[1971BAE/DEM]
C₁₀Mn₂O₁₀	[10170-69-1]	decacarbonyldimanganese				
	$\Delta_{\text{sub}}H$		80.3 ± 4.2	298		[1982PIL/SKI, 1958GOO/FAI]
	$\Delta_{\text{sub}}H$		92.3 ± 2.1	298	C	[1982CON/ZAF]
	$\Delta_{\text{sub}}H$	(351–428)	80.3 ± 2.1	390	MM	[1971BAE/DEM]
	$\Delta_{\text{sub}}H$		62.8 ± 4.2			[1960COT/MON]
	$\Delta_{\text{v}}H$	(428–463)	60.7 ± 1.3	446		[1971BAE/DEM]
C₁₀H₆Mn₂O₈S₂	[21321-38-0]	<i>bis</i> (μ -methanethiolato)octacarbonyldimanganese				
	$\Delta_{\text{sub}}H$		114.2 ± 0.8	340	C	[1995CON/GOB2]
C₁₀H₁₀Mn	[1271-27-8]	<i>bis</i> (cyclopentadienyl)manganese				
	$\Delta_{\text{sub}}H$	(298–445)	72.4	371.5	A	[1987STE/MAL]
	$\Delta_{\text{sub}}H$		75.7 ± 1.7	298		[1982PIL/SKI, 1971TEL/RAB]
	$\Delta_{\text{sub}}H$		72.4			[1956WIL/COT]
	$\Delta_{\text{v}}H$	(378–435)	58.0	393	A	[1987STE/MAL]
C₁₀H₁₄MnO₄	[14024-58-9]	<i>bis</i> (2,4-pentanedionato) manganese(II)				
	$\Delta_{\text{sub}}H$	(390–440)	139.3 ± 2.5	298	ME	[1990MAL/ALI]
	$\Delta_{\text{sub}}H$		87.0	343		[1981MAS/BAR]
	$\Delta_{\text{sub}}H$		88.7			[1972BOL, 2000DUN]
	$\Delta_{\text{sub}}H$		88.7	400		[1970GOE/BLO]
C₁₁H₅MnO₅	[13985-77-8]	phenyl(pentacarbonyl)manganese				
	$\Delta_{\text{sub}}H$		84.9 ± 4.4	298	C	[1982CON/ZAF]
C₁₂H₅MnO₆	[15612-92-7]	benzoyl(pentacarbonyl)manganese				
	$\Delta_{\text{sub}}H$		123 ± 3	298	C	[1982CON/ZAF]
C₁₂H₇MnO₅	[14049-86-6]	benzyl(pentacarbonyl)manganese				
	$\Delta_{\text{sub}}H$		84.5 ± 0.7	298	C	[1982CON/ZAF]
C₁₅H₁₂F₉MnO₆	[14526-24-0]	<i>tris</i> (1,1,1-trifluoro-2,4-pentanedionato) manganese(III)				
	$\Delta_{\text{sub}}H$	(378–413)	120.5 ± 9.2		GS	[1985MAT/KUW]
	$\Delta_{\text{sub}}H$		117.3			[1971ASH]
	$\Delta_{\text{sub}}H$		77.8			[1964WOO/JON]
C₁₅H₂₁MnO₆	[14284-89-0]	<i>tris</i> (2,4-pentanedionato) manganese(III)				
	$\Delta_{\text{fus}}H$		27.7	421.9	DSC	[2004SAB/MAR]
	$\Delta_{\text{sub}}H$	(320–380)	124.7 ± 3.8	298	ME	[1990MAL/ALI]
	$\Delta_{\text{sub}}H$		120 ± 10	298	E	[1988RIB/FER2]
	$\Delta_{\text{sub}}H$		99.0	392		[1981MAS/BAR]
	$\Delta_{\text{sub}}H$		113	370		[1970GOE/BLO]
	$\Delta_{\text{sub}}H$		77.8 ± 0.8	298		[1982PIL/SKI, 1968HIL/IRV]
C₁₈H₁₂MnN₂O₂	[14495-13-7]	<i>bis</i> (8-hydroxyquinolinato) manganese(II)				
	$\Delta_{\text{sub}}H$		194.6 ± 10.4	298	ME	[1994RIB/MAT]
	$\Delta_{\text{sub}}H$	(615–650)	208.4 ± 14	633	ME	[1984BUR/MOR]
	$\Delta_{\text{sub}}H$		226 ± 14	298		[1984BUR/MOR]
C₂₀H₁₆MnN₂O₂	[14515-78-7]	<i>bis</i> (8-hydroxy-2-methylquinolinato)manganese(II)				

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound		T_m (K)	Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)			
	$\Delta_{\text{sub}}H$	(521–541)	199.6 ± 7.2	531	ME	[1998RIB/MAT3]
	$\Delta_{\text{sub}}H$		211.2 ± 7.2	298		[1998RIB/MAT3]
C₃₀H₂₇MnO₆	[14376-07-9]	<i>tris</i> (1-phenylbutane-1,3-dionato)manganese(III)				
	$\Delta_{\text{sub}}H$		195 ± 10	298	E	[1988RIB/FER2]
C₃₃H₅₇MnO₆	[14324-99-3]	<i>tris</i> (2,2,6,6-tetramethylheptane-3,5-dionato)manganese(III)				
	$\Delta_{\text{sub}}H$		140 ± 10	298	E	[1988RIB/FER2]
C₄₄H₂₈MnN₄	[31004-82-7]	5,10,15,20-tetraphenylporphine manganese (II)				
	$\Delta_{\text{sub}}H$		175 ± 1		UV	[1993SHE/KAR]
MnF₂	[7782-64-1]	manganese (II) fluoride				
	$\Delta_{\text{sub}}H$		318.4 ± 8.4	298		[1964KEN/EHL]
Mo (molybdenum)						
C₆MoO₆	[13939-06-5]	molybdenum hexacarbonyl				
	$\Delta_{\text{sub}}H$	(265–300)	77.7			[2000OHT/CIC]
	$\Delta_{\text{sub}}H$	(316–423)	69.1	331	A	[1987STE/MAL]
	$\Delta_{\text{sub}}H$	(240–285)	76.9 ± 0.9	263	ME	[1979DAA/ERN, 1980BOX/ERN]
	$\Delta_{\text{sub}}H$		73.8 ± 1.0			[1975PIL/WAR, 1974BAR/PIL]
	$\Delta_{\text{sub}}H$	(343–383)	69.7	363		[1960MON/COT]
	$\Delta_{\text{sub}}H$	(323–403)	72.5			[1952REZ/SHV]
	$\Delta_{\text{sub}}H$	(292–308)	72.8			[1947LAN/GER]
	$\Delta_{\text{sub}}H$		68.2			[1935HIE/ROM]
C₇H₃MoNO₅	[17594-16-0]	acetonitrile molybdenum pentacarbonyl				
	$\Delta_{\text{sub}}H$	(260–279)	105.8 ± 5.6	298		[1980CAV/ERN]
C₈F₁₂Mo₂O₈	[36608-07-8]	dimolybdenum tetratetrafluoroacetate				
	$\Delta_{\text{sub}}H$	(330–370)	113.6 ± 1.7	350	ME,TE	[1984CAR]
C₈H₁₂CrMoO₈	[71561-64-3]	chromium molybdenum tetraacetate				
	$\Delta_{\text{sub}}H$		165.0 ± 8.4			[1982PIL/SKI]
C₈H₁₂Mo₂O₈	[na]	dimolybdenum tetraacetate				
	$\Delta_{\text{sub}}H$	(400–420)	170.5 ± 7	410	ME,TE	[1984CAR]
C₈H₁₂Mo₂O₈	[14221-06-8]	tetra- μ -acetatodimolybdenum(II)				
	$\Delta_{\text{sub}}H$		129 ± 1	491	C	[2008SLY/KON]
	$\Delta_{\text{sub}}H$		165.0 ± 8.4	298		[1982PIL/SKI, 1979CAV/GAR]
C₈H₂₄MoN₄	[100207-68-9]	<i>tetrakis</i> (dimethylamino)molybdenum				
	$\Delta_{\text{sub}}H$		88.4 ± 3	376	C	[1979ADE/CAV]
	$\Delta_{\text{sub}}H$		72.4 ± 6	298	C	[1979ADE/CAV, 1982PIL/SKI]
C₉H₉MoN₃O₃	[15038-48-9]	<i>tris</i> (acetonitrile) molybdenum tricarbonyl				
	$\Delta_{\text{sub}}H$	(283–308)	111.3 ± 3.0	298		[1980CAV/ERN]
	$\Delta_{\text{sub}}H$		96.0 ± 10.0	298		[1982PIL/SKI, 1978ADE/CON]
C₁₀H₅MoNO₅	[14324-76-6]	pyridine(pentacarbonyl)molybdenum				
	$\Delta_{\text{sub}}H$	(283–299)	102.0 ± 2.0	291	ME	[1979DAA/ERN]
C₁₀H₈MoO₃	[12125-77-8]	cycloheptatriene(tricarbonyl)molybdenum				
	$\Delta_{\text{sub}}H$		88.0 ± 4.0	298		[1982PIL/SKI, 1977BRO/CON]
C₁₀H₁₀Cl₂Mo	[12184-22-4]	dichloro <i>bis</i> (η^5 -2,4-cyclopentadien-1-yl)molybdenium				
	$\Delta_{\text{sub}}H$		100.4 ± 4.2	298	E	[1976TEL/RAB]
C₁₀H₁₀I₂Mo	[12184-29-1]	<i>bis</i> (η^5 -2,4-cyclopentadien-1-yl)diiodomolybdenium				
	$\Delta_{\text{sub}}H$		100.4 ± 4.2	298	E	[1976TEL/RAB]

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound			Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)		
C₁₀H₁₁MoNO₅	[19456-57-6] $\Delta_{\text{sub}}H$	piperidine(pentacarbonyl)molybdenum (275–289)	121.9 ± 5.1	282	ME	[1979DAA/ERN]
C₁₀H₁₂Mo	[1291-40-3] $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$	<i>bis</i> (η^5 -2,4-cyclopentadien-1-yl)dihydromolybdenum	81.4 ± 1.0 92.5 ± 2.1		ME	[1990DIA/DIO] [1976TEL/RAB]
C₁₁H₈MoO₄	[12146-37-1] $\Delta_{\text{sub}}H$	norbornadienemolybdenumtetracarbonyl	92.0 ± 4.0	298		[1982PIL/SKI, 1977BRO/CON]
C₁₂H₁₂Mo	[12129-68-9] $\Delta_{\text{sub}}H$	dibenzene molybdenum	94.6 ± 8			[1970COX/PIL, 1961FIS/FRI]
C₁₂H₁₆Mo	[39333-52-3] $\Delta_{\text{sub}}H$	dimethyldicyclopentadienylmolybdenum	70.4 ± 4.2	298		[1982PIL/SKI, 1980DEP]
C₁₂H₂₀Mo₂O₈	[41880-55-1] $\Delta_{\text{sub}}H$	<i>tetakis</i> [μ -(propanoato-O:O')]dimolybdenum	129.0 ± 1.1	491	C	[2008SLY/KON]
C₁₂H₃₆Mo₂N₆	[51956-20-8] $\Delta_{\text{sub}}H$	<i>hexakis</i> (dimethylamine)dimolybdenum(II)	111 ± 8	298	C	[1979ADE/CAV, 1981CAV/CON]
C₁₄H₂₀Mo₂O₈	[na] $\Delta_{\text{sub}}H$	di- μ -acetatobis(pentane-2,4-dionato)dimolybdenum(II)	163.0 ± 5.0	298		[1982PIL/SKI, 1979CAV/GAR]
C₁₆H₁₄Mo₂N₂O₄	[na] $\Delta_{\text{sub}}H$	di(6-methyl-2-hydroxypyridyl)diacetatodimolybdenum(II)	161.0 ± 4.0	298		[1982PIL/SKI, 1981CAV/GAR]
C₁₈H₁₅MoN₃O₃	[15279-79-5] $\Delta_{\text{sub}}H$	<i>tris</i> (pyridine)tricarbonylmolybdenum	142.0 ± 10.0	298		[1982PIL/SKI, 1978ADE/CON]
C₁₈H₄₂Mo₂O₆	[62521-20-4] $\Delta_{\text{sub}}H$	<i>hexakis</i> (isopropoxy)dimolybdenum	113 ± 10	298	C	[1981CAV/CON]
C₂₄H₂₄Mo₂N₄O₄	[67634-80-4] $\Delta_{\text{sub}}H$	tetra(6-methyl-2-hydroxypyridyl)dimolybdenum(II)	157.0 ± 3.0	298		[1982PIL/SKI, 1981CAV/GAR]
C₂₄H₅₆Mo₂O₈	[79376-50-4] $\Delta_{\text{sub}}H$	<i>octakis</i> (isopropoxy)dimolybdenum(II)	137.0 ± 15	298	C	[1981CAV/CON]
MoF₆	[7783-77-9] Δ_vH	molybdenum hexafluoride (318–363)	27.4	340		[1968NIS/NIK]
N (nitrogen)						
BrClFN	[145543-68-6] Δ_vH	bromochlorofluoroammonia (240–310)	30.2	275		[1996SLA/NOV]
BrF₂N	[15605-95-5] Δ_vH	bromodifluoroammonia (180–250)	23.2	215		[1996SLA/NOV]
Br₂FN	[145543-67-5] Δ_vH	dibromofluoroammonia (280–350)	33.6	315		[1996SLA/NOV]
Br₃N	[15162-90-0] Δ_vH	nitrogen tribromide (380–450)	44.1	415		[1996SLA/NOV]
Cl₂FN	[17417-38-8] Δ_vH	dichlorofluoroammonia (200–280)	25.7	240		[1996SLA/NOV]
Cl₃N	[10025-85-1] Δ_vH	nitrogen trichloride (280–440)	32.9	360		[1996SLA/NOV]

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound		T_m (K)	Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)			
HNO ₃	[7697-37-2] $\Delta_v H$	nitric acid (273–356)	38.6	312		[1966HOL]
NH ₃	[7664-41-7] $\Delta_{\text{sub}} H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$	ammonia (177–195) (293–392) (199–241)	31.2 22.7 23.5 23.4	308 239 239		[1937OVE/GIA] [1979ZAN/THO] [1937OVE/GIA] [1937OVE/GIA]
NH ₃ O	[7803-49-8] $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$	hydroxylamine (261–280) (273–298)	64.2 U 46.5	285		[1965BAC/BET] [1941BOD]
NH ₄ Br	[12124-97-9] $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$	ammonium bromide	183.7 187.9	550 298	I	[1971CAL/SMI] [1955LUF]
NH ₄ Cl	[12125-02-9] $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$	ammonium chloride (308–363)	168.6 176.6 ± 0.4 177	550 298	I TE	[1971CAL/SMI] [1961WAG/NEU] [1955LUF]
NH ₄ I	[12027-06-4] $\Delta_{\text{sub}} H$	ammonium iodide	182	298		[1955LUF]
NH ₄ CN	[12211-52-8] $\Delta_{\text{sub}} H$	ammonium cyanide	84.5	298		[1955LUF]
NH ₄ SCN	[1762-95-4] $\Delta_{\text{sub}} H$	ammonium thiocyanate	133.9	298		[1955LUF]
NO	[10102-43-9] $\Delta_v H$	nitric oxide	13.8	212	C	[1929JOH/GIA]
N ₂	[7727-37-9] $\Delta_v H$ $\Delta_v H$	nitrogen (63–126)	6.1 5.6	78 77		[1967EDE/THO] [1933GIA/CLA]
N ₂ F ₄	[10036-47-2] $\Delta_v H$	tetrafluorohydrazene	26.4	200		[1958COL/KEN]
N ₂ H ₄	[302-01-2] $\Delta_{\text{sub}} H$ $\Delta_v H$	hydrazene (288–353)	U 46.0 44.5	303		[1941GIG/RUN, 2001GIO] [1949SCO/OLI]
N ₂ H ₄ O ₂ S	[7803-58-9] $\Delta_{\text{sub}} H$	sulfamide (347–358)	101.5 ± 1.0			[1997DEZ/POI, 1959TAK/SHI]
N ₂ H ₄ O ₃	[6484-52-2] $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$	ammonium nitrate (349–438)	178.7 174.9	298		[1962BRA/JUN] [1955LUF]
N ₂ O	[10024-94-2] $\Delta_{\text{fus}} H$ $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$ $\Delta_v H$	nitrous oxide (68–80) (148–182) (103–123)	6.5 25.1 ± 0.4 24.6 23.6 16.5	182.4 74 161 113 184.7		[1974ATA/CHI] LE [1974BRY/CAZ] [1935BLU/GIA] MG [1930BLA/VAN] [1974ATA/CHI]

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)			
	$\Delta_v H$	(182–236)	16.1	221		[1945HOG]	
Na (sodium)							
C₄H₉ONa	[865-48-5] $\Delta_{\text{sub}} H$	sodium <i>tert</i> -butoxide	NA			[1990VOR/ZVE]	
C₆H₁₃ONa	[67638-48-6] $\Delta_{\text{sub}} H$	sodium methyl-diethylmethoxide	NA			[1990VOR/ZVE]	
C₇H₁₅ONa	[53535-82-3] $\Delta_{\text{sub}} H$	sodium triethylmethoxide	NA			[1990VOR/ZVE]	
C₁₆H₁₃O₃Na	[57495-14-4] $\Delta_{\text{fus}} H$	sodium ketoprofen	20.8	457		[1997HIL/MUL]	
C₃₂H₄₀F₁₂NaO₈Pr	[93557-93-8] $\Delta_{\text{sub}} H$	sodium <i>tetrakis</i> (1,1,1-trimethyl-5,5,5-trifluoro-2,4-pentanedionato)praseodymate	(423–483) 155 ± 2	453	T	[1993SYO/GOL]	
C₃₂H₄₀F₁₂NaO₈Tb	[12576-88-4] $\Delta_{\text{sub}} H$	sodium <i>tetrakis</i> (1,1,1-trimethyl-5,5,5-trifluoro-2,4-pentanedionato)terbate	(418–473) 163 ± 3	445	T	[1993SYO/GOL]	
C₃₂H₄₀F₁₂NaO₈Y	[12576-89-5] $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$	sodium <i>tetrakis</i> (1,1,1-trimethyl-5,5,5-trifluoro-2,4-pentanedionato)yttrate	(418–503) 130 ± 3 (463–503) 142 ± 12	460 483	T	[1993SYO/GOL] [1993SYO/GOL]	
Nb (niobium)							
C₅H₁₅NbO₅	[1066-25-7] $\Delta_{\text{sub}} H$	niobium pentamethoxide	80.3 ± 10.5		ME,E	[1972TEL/RAB, 1977TEL/RAB]	
C₁₀H₁₀Cl₂Nb	[12793-14-5] $\Delta_{\text{sub}} H$	<i>bis</i> (cyclopentadienyl)niobium dichloride	127.4 ± 4.4	298	ME	[2001DIO/PIE]	
C₁₀H₂₅NbO₅	$\Delta_v H$	pentaethylniobate	(376–414) 107.6	391	A	[1987STE/MAL]	
NbBr₅	[13478-45-8] $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$	niobium(V) pentabromide	(298–479) 115 ± 18 112.5	298 298		[1996TUR/EIC] [1996TUR/EIC, 1991KNA/KUB]	
NbCl₅	[10026-12-7] $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$	niobium(V) pentachloride	(298–479) 94 95 ± 16	298 298		[1996TUR/EIC, 1991KNA/KUB] [1996TUR/EIC]	
NbCl₃O	[113597-20-1] $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$	niobium(V) oxychloride	(298–607) 128.5 124 ± 16	298 298		[1996TUR/EIC, 1991KNA/KUB] [1996TUR/EIC]	
Nd (neodymium)							
C₁₅H₁₅Nd	[1273-98-9] $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$	<i>tris</i> (cyclopentadienyl)neodymium(III)	(533–633) 108.8 ± 3.8 (338–438) 134.7 ± 2.1			[1973BOR/KRA] ME [1971HAU, 1971HAU2]	
C₃₀H₃₀F₂₁NdO₆	[17978-76-6] $\Delta_{\text{sub}} H$	<i>tris</i> (1,1,1,2,2,3,3-heptafluoro-7,7-dimethyloctane-4,6-dione)neodymium(III)	(387–402) 155.2 ± 2.9		ME	[1971SWA/KAR]	
C₃₃H₅₇NdO₆	[15492-47-4] $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$	<i>tris</i> (2,2,6,6-tetramethylheptane-3,5-dionato)neodymium(III)	159.1 ± 3.4 92.9 ± 2.5 (378–423) 177	298 400	DSC ME	[1999SAN/PET] [1996TSY/DYA2, 2000GIE] [1981AMA/SAT]	

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound		T_m (K)	Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)			
	$\Delta_{\text{sub}}H$	(430–491)	158.4	460	BG	[1969SIC/DUB]
	Δ_vH	(491–510)	99.1		BG	[1969SIC/DUB]
Ni (nickel)						
(C ₃ NiO ₃)–(C ₃ F ₉ P)	[na]	<i>tris</i> (trifluoromethyl)phosphine—nickel tricarbonyl complex				
	Δ_vH	(273–323)	31.2	298		[1958EME/SMI]
C ₄ NiO ₄	[13463-39-3]	nickel tetracarbonyl				
	$\Delta_{\text{sub}}H$		41.6 ± 0.5			[1953WAL]
	Δ_vH	(277–412)	29.8	344		[1970BAE]
	Δ_vH		27.6 ± 1.3			[1957FIS/COT, 1982PIL/SKI]
	Δ_vH		28.0			[1955SPI/STA]
	Δ_vH		27.2	298		[1955SPI/STA]
	Δ_vH		30.2 ± 0.1			[1953WAL]
	Δ_vH	(250–315)	29.5	265		[1947STU]
	Δ_vH		30.1			[1942SUG/SAT, 1955SPI/STA]
	Δ_vH		27.2			[1903DEW/JON, 1955SPI/STA]
C ₄ H ₁₆ Cl ₂ N ₈ NiS ₈	[28813-19-6]	<i>trans</i> -dichloro- <i>tetrakis</i> (thiourea)nickel(II)				
	$\Delta_{\text{sub}}H$	(409–447)	74 ± 20			[1970ASH]
C ₆ H ₁₂ N ₂ NiS ₄	[15521-65-0]	<i>bis</i> (dimethyldithiocarbamate)nickel				
	Δ_vH	(448–478)	139.9 ± 2.1	463	A	[1987STE/MAL, 1978TAV/NEE, 1999DYK/SVO]
C ₈ F ₁₈ NiO ₂ P ₂	[15188-79-1]	dicarbonyl <i>bis</i> [<i>tris</i> (trifluoromethyl)phosphine]nickel				
	$\Delta_{\text{sub}}H$	(293–302)	47.2	298		[1966BUR/STR]
C ₈ F ₂₈ NiP ₄	[14917-18-1]	<i>tetrakis</i> [<i>bis</i> (trifluoromethyl)phosphinous fluoride]nickel				
	$\Delta_{\text{sub}}H$	(305–331)	66.6	318		[1966BUR/STR]
C ₁₀ H ₈ F ₆ NiO ₄	[14324-83-5]	<i>bis</i> (1,1,1-trifluoro-2,4-pentanedionato)nickel(II)				
	$\Delta_{\text{sub}}H$	(416–473)	157.7 ± 3.3		GS	[1985MAT/KUW]
C ₁₀ H ₁₀ Ni	[1271-28-9]	<i>bis</i> (cyclopentadienyl) nickel				
	$\Delta_{\text{sub}}H$		71.5 ± 0.6			[1988TOR/BAR2]
	$\Delta_{\text{sub}}H$		70.2 ± 1.5	298		[1984BAE/BAR2]
	$\Delta_{\text{sub}}H$	(353–419)	72.4 ± 1.3	298	MM	[1982PIL/SKI, 1975TEL/KIR, 1967TUR]
C ₁₀ H ₁₄ NiO ₄	[3264-82-2]	<i>bis</i> (2,4-pentanedionato)nickel(II)				
	$\Delta_{\text{sub}}H$	(357–420)	126.4 ± 4.4	298	ME	[1990MAL/ALI]
	$\Delta_{\text{sub}}H$		108.2 ± 5	207	DSC	[1987MUR/HIL]
	$\Delta_{\text{sub}}H$		108.2 ± 4.9	480	DSC	[1987RIB/FER]
	$\Delta_{\text{sub}}H$		155 ± 80	298	C	[1985MUR/SAK]
	$\Delta_{\text{sub}}H$	(378–403)	127.7 ± 10	381	ME	[1984BUR/MOR]
	$\Delta_{\text{sub}}H$		132 ± 10	298	ME	[1984BUR/MOR]
	$\Delta_{\text{sub}}H$		69.0		I	[1971ASH]
	$\Delta_{\text{sub}}H$		95.4	400		[1970GOE/BLO]
	$\Delta_{\text{sub}}H$		69.0			[1960BER/TRU, 1965BER/TRU]
C ₁₀ H ₂₀ N ₂ NiS ₄	[14267-17-5]	<i>bis</i> (diethyldithiocarbamate)nickel(II)				
	$\Delta_{\text{sub}}H$	(448–478)	157.3 ± 6.0		C	[1989RIB/REI]
	$\Delta_{\text{sub}}H$	(440–478)	152 ± 0.8	459	A	[1987STE/MAL, 1978TAV/NEE]
	$\Delta_{\text{sub}}H$	(507–650)	98.8 ± 6	579	DSC	[1979CAV/HIL2]
	$\Delta_{\text{sub}}H$	(443–543)	91.9 ± 6	493	DSC	[1979CAV/HIL2]
	$\Delta_{\text{sub}}H$		151.9 ± 2.1			[1976TAV/NEE]
	$\Delta_{\text{sub}}H$		61.1 ± 1.7		I	[1969DAS/WEN]

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₁₂ H ₈ N ₂ NiO ₄	[17653-01-9] $\Delta_{\text{sub}}H$	<i>bis</i> (picolinato)nickel(II)	76.6		I	[1963WOO/JON]
C ₁₃ H ₆ F ₂₄ N ₂ Ni ₂ O ₃ P ₄	[14402-98-3] $\Delta_{\text{sub}}H$	μ -carbonyldicarbonyl <i>bis</i> [μ -[(methylimino) <i>bis</i> [<i>bis</i> (trifluoromethyl)phosphine]]]dinickel	92.3	380		[1968SIN/BUR]
C ₁₄ H ₁₀ NiO ₄	[14263-01-5] $\Delta_{\text{sub}}H$	<i>bis</i> (salicyladehydato)nickel(II)	85.4		I	[1963WOO/JON]
C ₁₄ H ₁₂ N ₂ NiO ₂	[14283-99-9] $\Delta_{\text{sub}}H$	<i>bis</i> (salicyliminato)nickel(II)	158.2		I	[1963WOO/JON]
C ₁₄ H ₁₂ N ₂ NiO ₄	[14363-30-5] $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$	<i>bis</i> (salicylaldoximato)nickel(II)	106.6 ± 29 112 ± 29	413 298		[1984BUR/MOR] [1984BUR/MOR]
C ₁₄ H ₂₈ N ₂ NiS ₄	[14516-30-4] $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$ Δ_vH	<i>bis</i> (dipropyldithiocarbamate)nickel	147.2 ± 5.0 126.1 ± 0.8		C	[1989RIB/REI] [1978TAV/NEE]
C ₁₄ H ₂₈ N ₂ NiS ₄	[15694-55-0] $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$	<i>bis</i> (diisopropyldithiocarbamate) nickel complex	148.0 ± 5.0 143.4 ± 2.1	459.5	C A	[1989RIB/REI] [1987STE/MAL, 1978TAV/NEE]
C ₁₆ H ₈ F ₆ NiO ₂ S ₄	[14239-90-8] $\Delta_{\text{sub}}H$	<i>bis</i> (monothiothenoyltrifluoroacetate)nickel (II)	161.0 ± 5.1	298	C	[2007RIB/SAN2]
C ₁₆ H ₁₄ N ₂ NiO ₂	[14167-20-5] $\Delta_{\text{sub}}H$	<i>N,N-bis</i> (salicylidene)ethylenediaminonickel(II)	149.8 ± 7.0		ME	[1999ALI/MAL]
C ₁₆ H ₁₆ N ₂ NiO ₂	[57377-56-7] $\Delta_{\text{sub}}H$	<i>bis</i> (2-hydroxyacetophenamine)nickel(II)	130.2 ± 7.2		GS	[2009ARO/MAL]
C ₁₈ H ₁₂ N ₂ NiO ₂	[14100-15-3] $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$	<i>bis</i> (8-hydroxyquinolinato)nickel(II)	175.4 ± 6.7 129.9 ± 6 139 ± 6	298 486 298	ME ME	[1994RIB/MAT] [1984BUR/MOR] [1984BUR/MOR]
C ₁₈ H ₁₄ N ₄ Ni	[39251-81-5] $\Delta_{\text{sub}}H$	dibenzotetra-aza-annulene nickel(II) complex	116.6 ± 5.5	508	T	[1983FER/QUA]
C ₁₈ H ₂₀ N ₂ NiO ₂	[1161880-17-6] $\Delta_{\text{sub}}H$	<i>bis</i> (2-hydroxypropiofenamine)nickel (II)	113.0 ± 7.5		GS	[2009ARO/MAL]
C ₁₈ H ₃₆ N ₂ NiS ₄	[13927-77-0] $\Delta_{\text{sub}}H$ Δ_vH	<i>bis</i> (dibutyldithiocarbamate)nickel	132.6 ± 5.0 136.6	500	C	[1989RIB/REI] [1999DYK/SVO]
C ₁₈ H ₃₆ N ₂ NiS ₄	[28371-07-5] $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$ Δ_vH	<i>bis</i> (diisobutyldithiocarbamate)nickel	133.6 ± 5.0 152.1 ± 1.3 124	433	C	[1989RIB/REI] [1987STE/MAL, 1978TAV/NEE] [1999DYK/SVO]
C ₂₀ H ₁₆ N ₂ NiO ₂	[15200-70-1] $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$	<i>bis</i> (8-hydroxy-2-methylquinolinato)nickel(II)	170.9 ± 3.7 180.9 ± 3.7	496 298	ME	[1998RIB/MAT3] [1998RIB/MAT3]
C ₂₂ H ₃₈ NiO ₄	[14481-08-4]	<i>bis</i> (2,2,6,6-tetramethylheptane-3,5-dionato)nickel(II)				

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound		T_m (K)	Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)			
	$\Delta_{\text{sub}}H$	(453–493)	111		MEM	[1999EMM/PIC]
	$\Delta_{\text{sub}}H$		145.2 ± 10		ME	[1978IRV/SCH]
C₂₂H₄₂N₂NiO₂S₄	[1005000-26-9]	N,N-dibutyl-N'-thenoylthiourea				
	$\Delta_{\text{sub}}H$		204.7 ± 3.4	298	C	[2008RIB/SCH]
C₂₂H₄₂N₂NiO₂S₄	[1005000-14-5]	N,N-diisobutyl-N'-thenoylthiourea				
	$\Delta_{\text{sub}}H$		203.2 ± 2.4	298	C	[2008RIB/SCH]
C₂₂H₄₄N₂NiS₄	[55935-69-8]	<i>bis</i> [<i>bis</i> (3-methylbutyl)dithiocarbamate]nickel				
	$\Delta_{\text{sub}}H$	(429–468)	164.5	448		[1999DYK/SVO]
C₃₂H₁₆N₈Ni	[14055-02-8]	nickel(II) phthalocyanine				
	$\Delta_{\text{sub}}H$		144.6		TGA	[1995YAS/TAK]
C₄₄H₂₈N₄Ni	[14172-92-0]	5,10,15,20-tetraphenylporphine nickel(II)				
	$\Delta_{\text{sub}}H$		152 ± 4		GS	[2000PER/GOL]
NiBr₂	[13462-88-9]	nickel(II) bromide				
	$\Delta_{\text{sub}}H$	(714–969)	207 ± 4.0	841	TE	[1997BAR/BRU]
	$\Delta_{\text{sub}}H$		226 ± 1.0	298		[1997BAR/BRU]
NiFr₂	[10028-18-9]	nickel(II) fluoride				
	$\Delta_{\text{sub}}H$	(1054–1106)	332.2 ± 4.1		ME	[1964EHL/KEN]
Np (neptunium)						
(C₁₀H₂F₁₂NpO₆)–(C₃H₉OP)	[106617-32-7]	<i>bis</i> (1,1,1,5,5,5-hexafluoro-2,4-pentanedionato)neptunium(IV) dioxide- trimethylphosphine oxide adduct				
	$\Delta_{\text{sub}}H$	(370–418)	90 ± 3			[1988GRE/SID]
C₂₀H₄F₂₄NpO₈	[110900-26-0]	<i>tetrakis</i> (1,1,1,5,5,5-hexafluoro-2,4-pentanedionato)neptunium(IV)				
	$\Delta_{\text{sub}}H$	(314–375)	81 ± 3			[1988GRE/SID, 1987GRE/SID]
(C₂₀H₄F₂₄NpO₈)–(C₃H₉OP)	[110934-11-7]	<i>tetrakis</i> (1,1,1,5,5,5-hexafluoro-2,4-pentanedionato)neptunium(IV)- trimethylphosphine oxide adduct				
	$\Delta_{\text{sub}}H$	(353–404)	100 ± 4			[1988GRE/SID]
C₃₂H₄₀F₁₂NpO₈	[99791-99-8]	<i>tetrakis</i> (1,1,1-trimethyl-5,5,5-hexafluoro-2,4-pentanedionato)neptunium(IV)				
	$\Delta_{\text{sub}}H$	(374–424)	106 ± 3			[1988GRE/SID, 1987GRE/SID]
C₄₀H₄₀F₂₈NpO₈	[27988-02-9]	<i>tetrakis</i> (1,1,1,2,2,3,3-heptafluoro-7,7-dimethyloctane-4,6-dione)neptunium(IV)				
	$\Delta_{\text{sub}}H$	(350–368)	147.7 ± 2.9	359	ME	[1970SWA/KAR]
O (oxygen)						
H₂O₂	[7722-84-1]	hydrogen peroxide				
	Δ_vH	(277–363)	48.5	320		[1924MAA/HIE]
Os (osmium)						
C₁₀H₁₀Os	[1273-81-0]	<i>bis</i> (cyclopentadienyl)osmium (osmocene)				
	$\Delta_{\text{sub}}H$	(393–506)	72.9 ± 1.4			[1984BAE/BAR]
	$\Delta_{\text{sub}}H$		80.5 ± 1.7	298		[1984BAE/BAR]
	$\Delta_{\text{sub}}H$		75.3			[1959FIS/GRU]
	Δ_vH	(506–563)	56.3 ± 1.3	535		[1984BAE/BAR]
C₁₂O₁₂Os₃	[15696-40-9]	triosmium dodecacarbonyl				
	$\Delta_{\text{sub}}H$	(349–396)	134.4 ± 0.4			[1999CHA/GAR]
	$\Delta_{\text{sub}}H$		104.6 ± 20	298		[1982PIL/SKI, 1974CON/SKI]
	$\Delta_{\text{sub}}H$	(423–543)	108.4	483		[1974GAI/BAE]
	Δ_vH	(497–543)	101.7	520		[1974GAI/BAE2]

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound			Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)		
P (phosphorous) (see main table for organophosphorous compounds)						
Br₃P	[7789-60-8] $\Delta_v H$	tribromophosphine	48.5			[1996OVC/MAK, 1963HAR/HOL]
PBr₃S	[3931-89-3] $\Delta_{\text{sub}} H$	thiophosphoryl bromide	NA		GSM	[1941NIT/SEK]
Cl₃P	[7719-12-2] $\Delta_v H$	trichlorophosphine	32.6			[1996OVC/MAK, 1963HAR/HOL]
Cl₅P	[10026-13-8] $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$	pentachlorophosphorous	67.4 ± 2.3	390		[1973POL/POL]
			71.1 ± 5.0	298		[1973POL/POL]
F₂HOP	[14939-34-5] $\Delta_v H$	hydrophosphoryl difluoride (220–271)	36.1	245	T	[1967CHA/CAV]
F₂HPS	[13780-63-7] $\Delta_v H$	hydrothiophosphoryl difluoride (188–258)	29.1	223	T	[1967CHA/CAV]
F₂N₃OP	[38005-28-6] $\Delta_v H$	difluorophosphoryl azide	36.4	296		[1972ONE/SHR]
F₅P	[7647-19-0] $\Delta_v H$	phosphorous pentafluoride (179–189)	17.2	184	QM	[1937LIN/ROH]
F₆NP₃	[56564-56-8] $\Delta_v H$	tris(difluorophosphino)amine	31.2			[1975ARN/RAN]
F₆N₃P₃	[15599-91-4] $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$ $\Delta_v H$	trimeric phosphonitrilic fluoride (273–300)	53.6		T	[1958HAB/UEN]
			NA			[1958STE/LAN]
			32.1			[1958HAB/UEN]
F₈N₄P₄	[14700-00-6] $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$ $\Delta_v H$	tetrameric phosphonitrilic fluoride (273–303)	58.2		T	[1958HAB/UEN]
			NA			[1958STE/LAN]
			37.3			[1958HAB/UEN]
P₃Cl₆N₃	[940-71-6] $\Delta_{\text{sub}} H$	phosphonitrilic chloride (trimer)	76.1			[1943AUD/STE]
P₃Cl₆N₃	[940-71-6] $\Delta_{\text{fus}} H$	hexachlorocyclotriphosphazene	23.5	388.6	AC,DC	[1999LEB/KUL2]
P₄Cl₈N₄	[2950-45-0] $\Delta_{\text{fus}} H$	octachlorocyclotetraphosphazene	32.2	400.6	AC,DC	[1999LEB/KUL2]
PH₃	[7803-51-2] $\Delta_{\text{sub}} H$ $\Delta_v H$ $\Delta_v H$	phosphine (129–140)	17.2		MM	[1937STE/GIA]
			14.6 ± 0.1	186		[1939FRA/CLU]
			14.6	185		[1937STE/GIA]
Pb (lead)						
C₂H₈Pb	[30691-92-0] $\Delta_v H$	dimethylplumbane (173–223)	25.5	198		[1960AMB]
C₃H₁₀Pb	[7442-13-9]	trimethylplumbane				

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound		T_m (K)	Method	Reference	
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)				
	$\Delta_v H$	(193–243)	31.1	218		[1960AMB]	
C₄H₁₂Pb	[75-74-1]	tetramethyllead					
	$\Delta_{\text{fus}} H$		10.8	242.9		[1996DOM/HEA]	
	$\Delta_v H$		38.1 ± 0.4			[1959GOO/SCO, 1982PIL/SKI]	
	$\Delta_v H$	(298–308)	35.7	303		[1929TAN/NAG]	
C₅H₉F₅Pb	[812-34-0]	(pentafluoroethyl)trimethyllead					
	$\Delta_v H$	(295–329)	39.1	312	T	[1960KAE/PHI]	
C₈H₂₀Pb	[78-00-2]	tetraethyllead					
	$\Delta_{\text{fus}} H$		9.11	141.4		[1996DOM/HEA]	
	$\Delta_v H$		56.6 ± 1.0	298	C	[1980ABR/IRV]	
	$\Delta_v H$	(311–456)	57.3	326		[1947STU]	
	$\Delta_v H$		56.9 ± 2.5			[1956GOO/SCO, 1982PIL/SKI]	
	$\Delta_v H$	(273–343)	56.3	308		[1936BUC/NOR]	
C₁₀H₂F₁₂O₄Pb	[19648-88-5]	<i>bis</i> (1,1,1,5,5,5-hexafluoro-2,4-pentanedionato)lead(II)					
	$\Delta_{\text{sub}} H$	(368–413)	111.7 ± 1.3		GS	[1997KRI/SYS]	
	C₁₀H₁₄O₄Pb	[15282-88-9]	<i>bis</i> (2,4-pentanedionato)lead(II)				
		$\Delta_{\text{sub}} H$	(393–444)	102.4 ± 5.0		GS	[1997KRI/SYS]
		$\Delta_{\text{sub}} H$		87.0		LE	[1994GER/GER2, 1997KRI/SYS]
$\Delta_{\text{sub}} H$							
C₁₀H₂₀N₂PbS₄	[17549-30-3]	<i>bis</i> (diethyldithiocarbamate)lead complex					
	$\Delta_{\text{sub}} H$	(444–482)	129.9 ± 2.5	463	A	[1987STE/MAL, 1978TAV/NEE]	
C₁₀H₂₀O₄Pb	[56767-12-5]	lead(II) pentanoate					
	$\Delta_{\text{fus}} H$		12.6	355.6	DSC	[2008MAR/RAM]	
C₁₂H₁₀Br₂Pb	[3134-29-6]	diphenyl lead dibromide					
	$\Delta_{\text{sub}} H$	(298–398)	141.8 ± 0.8	298	ME	[1988GOL/SIT, 1976BUT/CAR]	
C₁₆H₂₀F₆O₄Pb	[21751-12-2]	<i>bis</i> (1,1,1-trimethyl-5,5,5-trifluoro-2,4-pentanedionato)lead(II)					
	$\Delta_{\text{sub}} H$	(393–463)	117.5 ± 2.8		GS	[1997KRI/SYS]	
C₁₈H₁₂N₂PbO₂	[14976-96-6]	<i>bis</i> (8-hydroxyquinolino)lead(II)					
	$\Delta_{\text{sub}} H$		187.1 ± 6.2	298	ME	[1994RIB/MAT]	
C₁₈H₁₅BrPb	[894-06-4]	triphenyl lead bromide					
	$\Delta_{\text{sub}} H$	(298–398)	134.7 ± 3.3	298	ME	[1988GOL/SIT, 1976BUT/CAR]	
C₁₈H₁₅IPb	[894-07-5]	triphenyl lead iodide					
	$\Delta_{\text{sub}} H$	(298–398)	130.1 ± 0.4	298	ME	[1988GOL/SIT, 1976BUT/CAR]	
C₂₀H₂₀F₁₄O₄Pb	[21600-78-2]	<i>bis</i> (6,6,7,7,8,8,8-heptafluoro-2,2-dimethyl-3,5-octanedionato)lead(II)					
	$\Delta_{\text{sub}} H$		75.0			[1992NYM/DES]	
C₂₂H₃₈O₄Pb	[21319-43-7]	<i>bis</i> (2,2,6,6-tetramethyl-3,5-heptanedionato)lead(II)					
	$\Delta_{\text{sub}} H$	(373–398)	117.5 ± 2.8		GS	[1997KRI/SYS]	
	$\Delta_{\text{sub}} H$		66.9		LE	[1994GER/GER2, 1997KRI/SYS]	
	$\Delta_{\text{sub}} H$		86.0			[1992NYM/DES]	
C₂₄H₂₀Pb	[595-89-1]	tetraphenyl lead					
	$\Delta_{\text{sub}} H$	(412–480)	151	446		[1987STE/MAL]	
	$\Delta_{\text{sub}} H$	(412–474)	159 ± 1	298	ME,TE	[1977KAN/MOR]	
	$\Delta_{\text{sub}} H$		194.6 ± 6.3	298	E	[1982PIL/SKI, 1972CAR/LAY]	
	$\Delta_{\text{sub}} H$	(298–316)	U 80.2	298	ME	[1962CAR/COO]	

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)			
	$\Delta_{\text{sub}}H$		82.8	298		[1972NEW]	
C₃₂H₁₆N₈Pb	[15187-16-3]	lead(II) phthalocyanine					
	$\Delta_{\text{sub}}H$		156.3		TGA	[1995YAS/TAK]	
	$\Delta_{\text{sub}}H$	(542–663)	195.7			[1984MRW/STA]	
PbF₂	[7783-46-2]	lead(II) fluoride					
	$\Delta_{\text{sub}}H$		267.8			[1969ZMB/HAS, 1971ADA/MAR]	
PbI₂	[10101-63-0]	lead(II) iodide					
	$\Delta_{\text{sub}}H$	(598–640)	173.1 ± 1.6	298	ME	[1996KON/COR]	
	$\Delta_{\text{sub}}H$	(474–582)	167.7 ± 1.3	298	MS	[1996KON/COR, 1985HIL/BEN]	
	$\Delta_{\text{sub}}H$	(900–1150)	182.5 ± 1.0	298		[1996KON/COR, 1979ABA/MAL]	
	$\Delta_{\text{sub}}H$	(563–613)	165.2 ± 1.8	298	ME	[1996KON/COR, 1964DUN/THO]	
	$\Delta_{\text{sub}}H$	(579–650)	166.4 ± 1.0	298	ME	[1996KON/COR, 1939NIW/SAT]	
PbSe	[12069-00-0]	lead selenide					
	$\Delta_{\text{sub}}H$	(835–1047)	226 ± 1		TE	[1993BRU/PIA]	
Pd (palladium)							
C₈H₁₀Pd	[1271-03-0]	(cyclopentadienyl)allyl palladium					
	$\Delta_{\text{sub}}H$	(291–333)	49.9	312	A	[1987STE/MAL, 1976ZOR/RAC]	
C₁₀H₂F₁₂O₄Pd	[64916-48-9]	bis(1,1,1,5,5,5-hexafluoro-2,4-pentanedionato)palladium(II)					
	$\Delta_{\text{sub}}H$	(318–368)	93.5 ± 0.6			[2005ZHA/STA]	
	$\Delta_{\text{sub}}H$	(293–313)	84.6 ± 1.6		ME	[2000ZHA/STA]	
	$\Delta_{\text{v}}H$	(371–398)	67.8 ± 1.4			[2005ZHA/STA]	
C₁₀H₈F₆O₄Pd	[63742-52-9]	bis(1,1,1-trifluoro-2,4-pentanedionato)palladium(II)					
	$\Delta_{\text{sub}}H$	(332–378)	115.2 ± 1.4		ME	[2000ZHA/STA]	
	$\Delta_{\text{sub}}H$	(423–448)	105.0 ± 0.8		GS	[1985MAT/KUW]	
C₁₀H₁₀F₆N₂O₂Pd	[203874-01-5]	bis(1,1,1-trifluoro-4-imino-2-pentanonato)palladium(II)					
	$\Delta_{\text{sub}}H$	(332–386)	110.9 ± 0.7		ME	[2000ZHA/STA]	
C₁₀H₁₄O₄Pd	[14024-61-4]	bis(2,4-pentanedionato)palladium(II)					
	$\Delta_{\text{sub}}H$	(402–452)	121.5 ± 1.5			[2005ZHA/STA]	
	$\Delta_{\text{sub}}H$		111.6			[2001MOR/ZHA]	
	$\Delta_{\text{sub}}H$	(347–416)	130.1 ± 2.8		ME	[2000ZHA/STA]	
	$\Delta_{\text{sub}}H$	(330–394)	122.7 ± 8.6	298		[1991MAL/ALI]	
	$\Delta_{\text{sub}}H$	(363–393)	127.6 ± 17	378	ME	[1984BUR/MOR]	
C₁₀H₂₀N₂PdS₄	[15170-78-2]	bis(diethyldithiocarbamate)palladium(II)					
	$\Delta_{\text{sub}}H$	(493–517)	153.1 ± 1.9			[2005ZHA/STA]	
	$\Delta_{\text{sub}}H$		153.1 ± 1.9			[1999ZEM/STA]	
	$\Delta_{\text{v}}H$	(520–558)	107.6 ± 1.2			[2005ZHA/STA]	
C₁₂H₂₈O₄P₂PdS₄	[52442-37-2]	palladium(II) diisopropylthiophosphate					
	$\Delta_{\text{sub}}H$	(384–413)	137.2 ± 5.6			[2005ZHA/STA]	
C₁₃H₁₈O₂Pd	[12130-90-4]	acetylacetonato(2,4-cyclooctadienyl)palladium(II)					
	$\Delta_{\text{sub}}H$	(344–362)	130.1 ± 6.3			[2005ZHA/STA]	
C₁₆H₂₀F₆O₄Pd	[77964-87-5]	bis(1,1,1-trimethyl-5,5,5-trifluoro-2,4-pentanedionato)palladium(II)					
	$\Delta_{\text{sub}}H$	(315–357)	131.4 ± 1.9		ME	[2000ZHA/STA]	
C₁₆H₂₀F₆O₆Pd	[301198-67-4]	bis(1,1-dimethyl-1-methoxy-5,5,5-trifluoro-2,4-pentanedionato)palladium(II)					

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)			
	$\Delta_{\text{sub}}H$	(315–369)	113.8 ± 1.2			ME	[2000ZHA/STA]
C₁₈H₁₂N₂O₂Pd	[14638-30-3]	<i>bis</i> (8-hydroxyquinolato)palladium(II)					
	$\Delta_{\text{sub}}H$	(483–503)	158.5 ± 4	493		ME	[1984BUR/MOR]
	$\Delta_{\text{sub}}H$		168 ± 4	298			[1984BUR/MOR]
C₂₀H₁₂F₆O₄Pd	[85159-01-9]	<i>bis</i> (4,4,4-trifluoro-1-phenyl-1,3-butanedionato)palladium(II)					
	$\Delta_{\text{sub}}H$	(386–452)	148.6 ± 1.4			ME	[2000ZHA/STA]
C₂₀H₁₈O₄Pd	[15186-07-9]	<i>bis</i> (1-phenyl-1,3-butanedionato)palladium(II)					
	$\Delta_{\text{sub}}H$	(410–471)	152.9 ± 1.4			ME	[2000ZHA/STA]
C₂₂H₃₈O₄Pd	[15214-66-1]	<i>bis</i> (2,2,6,6-tetramethyl-2,4-heptanedionato)palladium(II)					
	$\Delta_{\text{sub}}H$	(343–401)	125.4 ± 1.4			ME	[2000ZHA/STA]
C₄₄H₂₈N₄Pd	[76775-77-4]	5,10,15,20-tetraphenylporphine palladium(II)					
	$\Delta_{\text{sub}}H$		207 ± 5			GS	[2000PER/GOL]
Pm (promethium)							
C₃₃H₅₇O₆Pm	[67840-53-3]	<i>tris</i> (2,2,6,6-tetramethylheptan-3,5-dionato)promethium(III)					
	$\Delta_{\text{sub}}H$	(433–463)	131.8				[1979LEB/BER]
Pr (praseodymium)							
C₁₅H₁₅Pr	[11077-59-1]	<i>tris</i> (cyclopentadienyl)praseodymium					
	$\Delta_{\text{sub}}H$		125.5 ± 3.0	298			[1982PIL/SKI, 1974DEV/RAB]
	$\Delta_{\text{sub}}H$	(533–653)	113.0 ± 1.7				[1973BOR/KRA]
	$\Delta_{\text{sub}}H$	(338–438)	131.0 ± 2.1			ME	[1971HAU, 1971HAU2]
C₃₂H₄₀F₁₂O₈NaPr	[93557-93-8]	sodium <i>tetrakis</i> (1,1,1-trimethyl-5,5,5-trifluoro-2,4-pentanedionato)praseodymate					
	$\Delta_{\text{sub}}H$	(423–483)	155 ± 2	453		T	[1993SYO/GOL]
C₃₃H₅₇O₆Pr	[15492-48-5]	<i>tris</i> (2,2,6,6-tetramethylpentane-2,4-dionato)praseodymium(III)					
	$\Delta_{\text{sub}}H$		104.3 ± 2.6				[1996TSY/DYA2, 2000GIE]
	$\Delta_{\text{sub}}H$		163.0 ± 3.6			DSC	[1993AIR/SAN, 2000GIE]
	$\Delta_{\text{sub}}H$	(383–423)	178.7	403		ME	[1981AMA/SAT]
	$\Delta_{\text{sub}}H$	(450–495)	165.4	473		BG	[1969SIC/DUB]
	Δ_vH	(495–530)	109.2			BG	[1969SIC/DUB]
PrBr₃	[13536-53-3]	praseodymium(III) bromide					
	$\Delta_{\text{sub}}H$		288 ± 4	900		TE	[2000VIL/BRU2]
	$\Delta_{\text{sub}}H$		306 ± 4	298			[2000VIL/BRU2]
	$\Delta_{\text{sub}}H$		292	298			[2000VIL/BRU2]
PrCl₃	[10361-79-2]	praseodymium(III) chloride					
	$\Delta_{\text{sub}}H$		317 ± 4	1000		TE	[2000VIL/BRU2]
	$\Delta_{\text{sub}}H$		340 ± 4	298			[2000VIL/BRU2]
	$\Delta_{\text{sub}}H$		324	298			[2000VIL/BRU2]
PrI₃	[13813-23-5]	praseodymium(III) iodide					
	$\Delta_{\text{sub}}H$		263 ± 4	900		TE	[2000VIL/BRU2]
	$\Delta_{\text{sub}}H$		280 ± 4	298			[2000VIL/BRU2]
	$\Delta_{\text{sub}}H$		275	298			[2000VIL/BRU2]
Pt (platinum)							
C₈H₁₄Pt	[1271-07-4]	cyclopentadienyltrimethylplatinum					
	$\Delta_{\text{sub}}H$		77.8 ± 2.0	298			[1982PIL/SKI, 1977TEL/RAB]
C₁₀H₈F₆O₄Pt	[67596-99-0]	<i>cis-bis</i> (trifluoroacetylacetonato) platinum					

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)			
	$\Delta_{\text{sub}}H$	(412–461)	106.2 ± 2.1				[2006ZHA/BAI]
C₁₀H₈F₆O₄Pt	[76740-70-0]	<i>trans</i> -bis(trifluoroacetylacetonato) platinum					
	$\Delta_{\text{sub}}H$	(437–496)	109.9 ± 2.9				[2006ZHA/BAI]
C₁₀H₁₄O₄Pt	[15170-57-7]	<i>bis</i> (2,4-pentanedionato)platinum(II)					
	$\Delta_{\text{sub}}H$		105.9				[2001MOR/ZHA]
	$\Delta_{\text{sub}}H$	(363–383)	129.4 ± 9	373	ME		[1984BUR/MOR]
	$\Delta_{\text{sub}}H$		133 ± 9	298			[1984BUR/MOR]
C₁₀H₂₀N₂PtS₄	[15730-38-8]	<i>bis</i> (diethyldithiocarbamate)platinum(II)					
	$\Delta_{\text{sub}}H$		157.1 ± 2.0				[1999ZEM/STA]
C₁₂H₁₆Pt	[42613-14-9]	dicyclopentadienyldimethylplatinum					
	$\Delta_{\text{sub}}H$		83.7 ± 3.5	298			[1982PIL/SKI, 1977TEL/RAB]
Pu (plutonium)							
C₄₀H₄₀F₂₈O₈Pu	[28041-99-8]	<i>tetrakis</i> (1,1,1,2,2,3,3-heptafluoro-7,7-dimethyloctane-4,6-dione)plutonium(IV)					
	$\Delta_{\text{sub}}H$	(349–363)	153.5 ± 7.9	356	ME		[1970SWA/KAR]
Rb (rubidium)							
C₅H₉O₂Rb	[70205-79-7]	rubidium pivalate					
	$\Delta_{\text{sub}}H$		167.1 ± 5.6				[1998KHO/RYSK]
Re (rhenium)							
C₄H₆Br₄O₄Re₂	[75027-96-2]	<i>bis</i> (μ -acetato)tetrabromodirhenium stereoisomer (or 75081-56-0)					
	$\Delta_{\text{sub}}H$ (<i>cis</i>)	(410–510)	66.6				[1984STE/ALI]
	$\Delta_{\text{sub}}H$ (<i>trans</i>)	(410–510)	59.9				[1984STE/ALI]
C₄H₆Cl₄O₄Re₂	[62320-69-8; 100495-10-1]	<i>bis</i> (μ -acetato)tetrachlorodirhenium stereoisomer					
	$\Delta_{\text{sub}}H$ (<i>cis</i>)	(450–560)	72.8				[1984STE/ALI]
	$\Delta_{\text{sub}}H$ (<i>trans</i>)	(450–560)	64.7				[1984STE/ALI]
C₅BrO₅Re	[14220-21-4]	bromopentacarbonylrhenium					
	$\Delta_{\text{sub}}H$		92.1 ± 2		C		[1983ALT/CON]
C₅ClO₅Re	[14099-01-5]	chloropentacarbonylrhenium					
	$\Delta_{\text{sub}}H$		110.9 ± 2		C		[1983ALT/CON]
C₅HO₅Re	[16457-30-0]	rhenium hydride pentacarbonyl complex					
	$\Delta_{\text{sub}}H$	(279–369)	45.1	324	A		[1987STE/MAL]
C₆H₃O₅Re	[14524-92-6]	rhenium methylpentacarbonyl complex					
	$\Delta_{\text{sub}}H$	(315–380)	65.2	347.5	A		[1987STE/MAL, 1960HIE/BRA]
	$\Delta_{\text{sub}}H$		70.0 ± 2	298	C		[1983ALT/CON]
	$\Delta_{\text{sub}}H$		65.3 ± 1.0	298			[1982PIL/SKI, 1974BRO/CON]
	$\Delta_{\text{sub}}H$	(313–383)	64.9				[1958HIE/WAG]
C₁₀O₁₀MnRe	[14693-30-2]	manganese rhenium decacarbonyl					
	$\Delta_{\text{v}}H$	(440–463)	56.5	451			[1971BAE/DEM]
C₁₀O₁₀Re₂	[14285-68-8]	dirhenium decacarbonyl					
	$\Delta_{\text{sub}}H$		100.9 ± 2	298			[1983ALT/CON]
	$\Delta_{\text{sub}}H$		93.3 ± 4.2	298			[1982PIL/SKI, 1974BRO/CON]
	$\Delta_{\text{sub}}H$	(363–450)	77.6	406	MM		[1971BAE/DEM]
	$\Delta_{\text{sub}}H$		79.5				[1961GIN, 1971BAE/DEM]
	$\Delta_{\text{v}}H$	(454–483)	68.7	468			[1971BAE/DEM]

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
Rh (rhodium)						
C₇H₇O₄Rh	[14874-82-9] $\Delta_{\text{sub}}H$	dicarbonyl-2,4-pentanedionatorhodium complex (276–301)	87 ± 2.9	289	ME	[1978JES/ERN, 1987STE/MAL]
C₉H₁₃Cl₂O₂Rh	[12282-04-1] $\Delta_{\text{sub}}H$	<i>bis</i> (chloroethylene)-2,4-pentanedionatorhodium complex (275–288)	117.2 ± 7.1	281	ME	[1978JES/ERN, 1987STE/MAL]
C₉H₁₅O₂Rh	[12082-47-2] $\Delta_{\text{sub}}H$	<i>bis</i> (ethylene)-2,4-pentanedionatorhodium complex (282–301)	97.9 ± 7.1	292	ME	[1978JES/ERN, 1987STE/MAL]
C₁₀H₁₄O₄Rh	[69047-66-1] $\Delta_{\text{sub}}H$	<i>bis</i> (2,4-pentanedionato)rhodium(II) (383–447)	173.2 ± 7.0	298		[1991MAL/ALI]
C₁₁H₁₉O₂Rh	[12282-38-1] $\Delta_{\text{sub}}H$	<i>bis</i> (propylene)-2,4-pentanedionatorhodium complex (270–296)	86.2 ± 1.7	283	ME	[1978JES/ERN, 1987STE/MAL]
C₁₃H₁₉O₆Rh	[31724-87-5] $\Delta_{\text{sub}}H$	<i>bis</i> (vinylacetate)-2,4-pentanedionatorhodium complex (309–328)	121.3 ± 3	319	ME	[1978JES/ERN]
C₁₃H₁₉O₆Rh	[31724-88-6] $\Delta_{\text{sub}}H$	<i>bis</i> (methyl acrylate)-2,4-pentanedionatorhodium complex (311–327)	111.7 ± 4.6	319	ME	[1978JES/ERN]
C₁₅H₂₁O₆Rh	[14284-92-5] $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$	<i>tris</i> (2,4-pentanedionato)rhodium(II)	118.8 NA			[2001MOR/ZHA] [1994GER/GER]
C₁₆O₁₆Rh₆	[28407-51-4] $\Delta_{\text{sub}}H$	hexarhodiumhexadecacarbonyl (117.2 ± 20.0)		298		[1982PIL/SKI, 1975BRO/CON]
Ru (ruthenium)						
C₅O₅Ru	[16406-48-7] Δ_vH	ruthenium pentacarbonyl (243–323)	42.2	283		[1991KOE/BOR]
C₁₀H₁₀Ru	[1287-13-4] $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$ Δ_vH	<i>bis</i> (cyclopentadienyl)ruthenium (ruthenocene) (383–479)	76.2 ± 1.4 82.7 ± 1.7 77.6 ± 1.6	298		[1984BAE/BAR] [1984BAE/BAR] [1967TUR]
C₁₅H₃F₁₈O₆Ru	[16827-63-7] $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$	<i>tris</i> (1,1,1,5,5,5-hexafluoro-2,4-pentanedionato)ruthenium(III) (299–313)	114.1 ± 1.0 114.5 ± 1.0	306 298	ME ME	[2001RIB/MON] [2001RIB/MON]
C₁₅H₁₂F₉O₆Ru	[16702-38-8] $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$	<i>tris</i> (1,1,1-trifluoro-2,4-pentanedionato)ruthenium(III) (346–467)	131.4 ± 4.6		ME	[2009MOR/ZHE]
		(350–369)	126.8 ± 1.0	360	ME	[2001RIB/MON]
			129.9 ± 1.0	298	ME	[2001RIB/MON]
		(383–423)	90.0 ± 3.0			[1996BYK/MOR]
C₁₅H₂₁O₆Ru	[14284-93-6] $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$	<i>tris</i> (2,4-pentanedionato)ruthenium(III) (374–434)	129.1 ± 2.0		ME	[2009MOR/ZHE]
		(394–441)	148.8 ± 1.7	418	ME	[2009SID/SID]
		(377–435)	128.9 ± 1.9		ME	[2007IGU/SEM]
			126.6			[2001MOR/ZHA]
		(423–493)	127.0 ± 0.9			[1996BYK/MOR]
		(398–413)	139.7 ± 2.5	406	ME	[1993RIB/GIE]
			145.1 ± 2.5	298	ME	[1993RIB/GIE]
C₂₄H₃₆F₃O₆Ru	[na]	<i>tris</i> (1,1,1-trifluoro-5,5-dimethyl-2,4-hexandianato)ruthenium(III)				

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(322–347)	75.7 ± 3.3		ME	[2009MOR/ZHE]
C₃₃H₅₄F₃O₆Ru	[na] $\Delta_{\text{sub}} H$	<i>tris</i> (2,2,6,6-tetramethyl-4-fluoro-3,5-heptanedionato)ruthenium(III) (353–393)	130.0 ± 2.7		ME	[2009MOR/ZHE]
C₃₃H₅₇O₆Ru	[38625-54-6] $\Delta_{\text{sub}} H$	<i>tris</i> (2,2,6,6-tetramethyl-3,5-heptanedionato)ruthenium(III) (353–393)	149.2 ± 2.2		ME	[2009MOR/ZHE]
S (sulfur)						
Br₂OS	[507-16-4] $\Delta_v H$	thionyl bromide (313–439)	43.6	330		[1999DYK/SVO]
Br₂S₂	[13172-31-1] $\Delta_v H$	disulfur dibromide (365–503)	53.9	380		[1999DYK/SVO]
Br₂FO₂S	[13536-61-3] $\Delta_v H$	sulfuryl bromide fluoride (236–333)	32.0	251		[1999DYK/SVO]
ClF₂NO₂S	[30913-20-3] $\Delta_v H$	difluoroamidodisulfuryl chloride (232–290)	31.2	261		[1971ZAB/SHR]
ClFOS	[14177-25-4] $\Delta_v H$	thionyl chloride fluoride (212–304)	27.7	227		[1999DYK/SVO]
ClFO₂S	[13637-84-8] $\Delta_v H$	sulfuryl chloride fluoride (211–300)	29.0	226		[1999DYK/SVO]
ClFO₅S₂	[13637-85-9] $\Delta_v H$	pyrosulfuryl chloride fluoride (284–396)	40.8	299		[1999DYK/SVO]
ClHO₃S	[7790-94-5] $\Delta_v H$	chlorosulfonic acid (324–454)	45.8	339		[1999DYK/SVO]
Cl₂OS	[7719-09-7] $\Delta_v H$	thionyl chloride (257–372)	32.4	272		[1999DYK/SVO]
Cl₂O₂S	[7791-25-5] $\Delta_v H$	sulfuryl chloride (357–365)	34.5	272		[1999DYK/SVO]
Cl₂O₅S₂	[7791-27-7] $\Delta_v H$	pyrosulfuryl dichloride (325–450)	44.7	340		[1999DYK/SVO]
Cl₂S	[10545-99-0] $\Delta_v H$	sulfur chloride (265–348)	43.8	280		[1999DYK/SVO]
Cl₂S₂	[10025-67-9] $\Delta_v H$	disulfur dichloride (306–439)	41.1	321		[1999DYK/SVO]
FHO₃S	[7789-21-1] $\Delta_v H$	fluorosulfonic acid (343–459)	55.7	358		[1999DYK/SVO]
FNS	[18820-63-8] $\Delta_v H$	thiazyl fluoride (270–299)	21.7	285		[1999DYK/SVO]
F₂HPS	[13780-63-7] $\Delta_v H$	hydrothiophosphoryl difluoride (188–258)	29.1	223	T	[1967CHA/CAV]
F₂N₂S	[500010-01-5] $\Delta_v H$	dinitrogen sulfur difluoride (192–281)	23.9	207		[1999DYK/SVO]
F₂OS	[7783-42-8] $\Delta_v H$	thionyl fluoride (173–244)	23.7	188		[1999DYK/SVO]

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound		T_m (K)	Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)			
F_2O_2S	[2699-79-8] Δ_vH	sulfuryl fluoride (160–233)	20.0	175		[1999DYK/SVO]
F_2O_4S	[13997-94-9] Δ_vH	peroxysulfuryl difluoride (198–248)	25.7	223		[1975GAM/SIC]
$F_2O_5S_2$	[13036-75-4] Δ_vH	pyrosulfuryl difluoride (240–346)	31.4	255		[1999DYK/SVO]
$F_2O_8S_3$	[13709-33-6] Δ_vH	trisulfur octoxide difluoride (296–419)	40.7	311		[1999DYK/SVO]
F_2S_2	[16860-99-4] Δ_vH	disulfur difluoride (153–196)	14.9	168		[1999DYK/SVO]
F_3NO_3S	[6816-12-2] Δ_vH	N,N-difluorohydroxylamine-O-fluorosulfonate (206–272)	24.6	239		[1963LUS/CAD]
F_3NS	[501679-94-3] Δ_vH	N-fluorosulfur difluoride amide (213–246)	24.1	230		[1969GLE/MEW]
F_3NS	[15930-75-3] Δ_vH	nitrogen fluoride sulfide (184–268)	23.1	199		[1999DYK/SVO]
F_4OS	[13809-54-1] Δ_vH	sulfur oxide tetrafluoride (166–240)	21.4	181		[1999DYK/SVO]
$F_4O_5S_2$	[44982-62-9] Δ_vH	disulfur pentoxide tetrafluoride (246–353)	18.0	261		[1999DYK/SVO]
F_4S	[7783-60-0] Δ_vH Δ_vH	sulfur tetrafluoride (170–250) (160–224)	21.1 24.6	185 192		[1999DYK/SVO] [1955BRO/ROB]
$F_6O_3S_2$	[81439-35-2] Δ_vH	pentafluorosulfur fluorosulfane (228–273)	32.2	250		[1962COH/MAC]
F_6S	[2551-62-4] $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$	sulfur hexafluoride (175–207)	23.2 ± 0.01 23.3	186 191		[1994OHT/YAM] [1932KLE/HEN]
$F_{10}O_2S_2$	[12395-41-4] Δ_vH	thiosulfuryl decafluoride (239–344)	31.8	242		[1999DYK/SVO]
$F_{10}S_2$	[5714-22-7] Δ_vH Δ_vH	disulfur decafluoride (226–322) (222–273)	30.1 29.6	241 237		[1999DYK/SVO] [1962COH/MAC]
$F_{14}O_2S_3$	[108021-40-5] Δ_vH	SF ₅ OSF ₄ OSF ₅	33.4			[1963PAS/ROB]
$F_{18}O_4S_4$	[na] Δ_vH	SF ₅ OSF ₄ OOSF ₅ OSF ₅	47.5			[1963PAS/ROB]
H_2S	[7783-06-4] $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$ Δ_vH Δ_vH Δ_vH	hydrogen sulfide (128–142) (164–187) (185–228) (228–363) (187–213)	22.5 25.4 19.5 18.6 21.9	135 175 200 243 200	MG	[1951CLA/COC] [1936GIA/BLU] [1999DYK/SVO] [1999DYK/SVO] [1936GIA/BLU]

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)			
H₂S₂	[13465-07-1]	dihydrogen disulfide					
	$\Delta_v H$	(256–367)	34.0	271		[1999DYK/SVO]	
	$\Delta_v H$		33.8 ± 0.1	293	C	[1958FEH/HIT]	
H₂S₂O₇	[7783-05-3]	pyrosulfuric acid					
	$\Delta_{\text{fus}} H$		23.81	308.4		[1961DAC/WYA]	
H₂S₃	[13845-23-3]	dihydrogen trisulfide					
	$\Delta_v H$	(328–474)	43.1	343		[1999DYK/SVO]	
	$\Delta_v H$		45.5 ± 0.2	293	C	[1958FEH/HIT]	
H₂S₄	[13465-25-5]	dihydrogen tetrasulfide					
	$\Delta_v H$	(384–547)	52.2	399		[1999DYK/SVO]	
	$\Delta_v H$		56.8 ± 0.3	293	C	[1958FEH/HIT]	
H₂S₅	[13465-24-4]	dihydrogen pentasulfide					
	$\Delta_v H$	(426–592)	61.5	441		[1999DYK/SVO]	
	$\Delta_v H$		68.4 ± 0.6	293	C	[1958FEH/HIT]	
NHS₇	[293-42-5]	heptasulfur imide					
	$\Delta_{\text{fus}} H$		18.83	386.7		[1975HAM/KUD]	
SO₂	[7446-09-5]	sulfur dioxide					
	$\Delta_v H$	(200–263)	24.9	263		[1938GIA/STE]	
	$\Delta_v H$		24.9	263	C	[1938GIA/STE]	
SO₃	[7446-11-9]	sulfur trioxide					
	$\Delta_v H$	(290–318)	46.7	290		[1985KON/STR]	
	$\Delta_v H$	(290–318)	45.5 ± 0.8	298		[1985KON/STR]	
	$\Delta_v H$	(353–473)	32.4	368		[1963ABE/TIL]	
SO₃	[7446-11-9]	γ -sulfur trioxide					
	$\Delta_{\text{fus}} H$	(16–332)	9.35	290.2	AC	[1989KON/STR]	
Sb (antimony)							
CH₅Sb	[23362-09-6]	methylstibine					
	$\Delta_v H$	(223–273)	27.4	248		[1959BUR/GRA]	
C₂H₇Sb	[23362-10-9]	dimethylstibine					
	$\Delta_v H$	(241–273)	30.8	257		[1959BUR/GRA]	
C₂H₈BSb	[60646-39-1]	dimethylstibinoborine					
	$\Delta_v H$	(234–273)	32.1	254		[1959BUR/GRA]	
C₃Cl₂F₉Sb	[420-74-6]	<i>tris</i> (trifluoromethyl)antimony dichloride					
	$\Delta_v H$	(243–323)	38.8	283		[1957DAL/EME]	
C₃F₉Sb	[432-05-3]	<i>tris</i> (trifluoromethyl)stibine					
	$\Delta_v H$	(215–343)	34.7	279		[1957DAL/EME]	
C₃H₉Sb	[594-10-5]	trimethylstibine					
	$\Delta_v H$	(249–296)	32.5 ± 0.01	298		[2010FUL/MOR]	
	$\Delta_v H$		28.9 ± 1.3			[1955LON/SAC, 1982PIL/SKI]	
	$\Delta_v H$		31.2		BG	[1946BAM/LEV]	
C₄H₁₂Sb₂	[41422-43-9]	tetramethylbistibine					
	$\Delta_v H$	(325–358)	46.9	341		[1959BUR/GRA]	
C₆H₉Sb	[5613-68-3]	trivinylstibine					
	$\Delta_v H$	(293–363)	38.7	308		[1957MAI/SEY, 1984BOU/FRI]	

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound		T_m (K)	Method	Reference	
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)				
C₆H₁₅Sb	[617-85-6]	triethylstibine					
	$\Delta_{\text{fus}}H$		9.45	153.9		[1996DOM/HEA]	
	Δ_vH	(238–309)	47.6	260		[2006FUL/RUZ]	
	Δ_vH	(238–309)	45.6	298		[2006FUL/RUZ]	
	Δ_vH	(238–309)	44.2	320		[2006FUL/RUZ]	
	Δ_vH	(238–309)	41.4	360		[2006FUL/RUZ, 1946BAM/LEV]	
	Δ_vH	(238–309)	38.3	400		[2006FUL/RUZ, 1946BAM/LEV]	
	Δ_vH	(193–333)	39.9 ± 1.3	306		[2001BAE]	
C₆H₁₅Sb	[138260-00-1]	<i>tert</i> -butyldimethylantimony					
	$\Delta_{\text{sub}}H$	(248–283)	43.5 ± 0.01	266		[2010FUL/MOR]	
	Δ_vH	(288–308)	41.1 ± 0.01	298		[2010FUL/MOR]	
	C₁₅H₃₀N₃S₆Sb	[22914-48-3]	<i>tris</i> (N,N-diethyldithiocarbamate)antimony(III)				
		$\Delta_{\text{sub}}H$		160 ± 2	298		[1994LIE/MAR]
	C₁₈H₁₅Sb	[603-36-1]	triphenylantimony				
		$\Delta_{\text{sub}}H$		106.3 ± 8.4	298		[1982PIL/SKI, 1960BIR]
	C₂₁H₄₂N₃S₆Sb	[226980-30-9]	<i>tris</i> (dipropyldithiocarbamate)antimony(III)				
$\Delta_{\text{sub}}H$			169.5 ± 6.1		DSC,E	[1999NEV/GOU]	
C₂₇H₅₄N₃S₆Sb	[14907-93-8]	<i>tris</i> (N,N-dibutyldithiocarbamate)antimony(III)					
	$\Delta_{\text{sub}}H$		179 ± 3	298		[1994LIE/MAR]	
C₂₇H₅₄N₃S₆Sb	[41594-79-0]	<i>tris</i> (N,N-diisobutyldithiocarbamate)antimony(III)					
	$\Delta_{\text{sub}}H$		157 ± 3	298	DSC, E	[1997DES/DES]	
Sc (scandium)							
C₁₅H₃F₁₈O₆Sc	[18990-42-6]	<i>tris</i> (1,1,1,5,5,5-hexafluoro-2,4-pentadionato)scandium(III)					
	$\Delta_{\text{sub}}H$	(333–363)	55.0		TGA	[2000FAH/BAR]	
	$\Delta_{\text{sub}}H$	(313–348)	60.2 ± 1.2		I	[1978KOM/GUR]	
C₁₅H₁₂F₉O₆Sc	[14634-68-5]	<i>tris</i> (1,1,1-trifluoro-2,4-pentanedionato)scandium(III)					
	$\Delta_{\text{sub}}H$	(373–403)	78.0		TGA	[2000FAH/BAR]	
	$\Delta_{\text{sub}}H$	(363–433)	117.6 ± 1.7			[1985MAT/KUW]	
	$\Delta_{\text{sub}}H$	(366–413)	53.2 ± 1.0		I	[1978KOM/GUR]	
	Δ_vH	(397–457)	82.2 ± 0.8	427		[1978CHU/IGU]	
C₁₅H₁₅Sc	[1298-54-0]	<i>tris</i> (cyclopentadienyl)scandium					
	$\Delta_{\text{sub}}H$		97.1 ± 3.5	298		[1982PIL/SKI, 1974DEV/RAB]	
C₁₅H₂₁O₆Sc	[14284-94-7]	<i>tris</i> (2,4-pentanedionato)scandium(III)					
	$\Delta_{\text{fus}}H$		28.8	460		[1970MEL/MER2]	
	$\Delta_{\text{sub}}H$	(413–443)	95		TGA	[2000FAH/BAR]	
	$\Delta_{\text{sub}}H$	(393–453)	58.2 ± 0.8		I	[1978KOM/GUR]	
	$\Delta_{\text{sub}}H$		99.6 ± 0.8	298	HSA	[1970MEL/MER, 1970MEL/MER2]	
C₃₃H₅₇O₆Sc	[15492-49-6]	<i>tris</i> (2,2,6,6-tetramethylheptan-3,5-dionato)scandium(III)					
	$\Delta_{\text{sub}}H$	(375–424)	97 ± 1		TG-TS	[2009SEL/RAG]	
	$\Delta_{\text{sub}}H$	(413–443)	90		TGA	[2000FAH/BAR]	
			79.6 ± 2.4			[1997SAN/ROC]	

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound			Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)		
	$\Delta_v H$	(434–465)	77 ± 2		TG-TS	[2009SEL/RAG]
Se (selenium)						
CBrF₃Se	[753-95-7] $\Delta_v H$	trifluoromethylselenyl bromide	30.9			[1980GOM/WEI]
CClF₃Se	[1495-26-7] $\Delta_v H$	trifluoromethylselenyl chloride	27.6			[1980GOM/WEI]
COSe	[1603-84-5] $\Delta_v H$ $\Delta_v H$ $\Delta_v H$	carbon oxyselenide	22.1 21.7 22.1	236 236 211		[1999DYK/SVO] [1947STU] [1937PUR/ZAH]
CSSe	[5951-19-9] $\Delta_v H$ $\Delta_v H$	carbon selenide sulfide	35.5 33.6	241 288		[1999DYK/SVO] [1914STO/WIL, 1984BOU/FRI]
CSe₂	[506-80-9] $\Delta_{\text{sub}} H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$	carbon diselenide	46.3 39.1 35.9 37.2 ± 0.8 39.0	224 245 305 288	A	[1987STE/MAL, 1966GAT/DRA] [1999DYK/SVO] [1999DYK/SVO] [1966GAT/DRA, 1982PIL/SKI] [1947IVE/PIT, 1984BOU/FRI]
CHF₃Se	[55446-31-6] $\Delta_v H$	trifluoromethaneselenol	22.5			[1980GOM/WEI]
CH₃FO₃Se	[17697-13-1] $\Delta_v H$	fluoroselenic acid, methyl ester	46.9			[1967PAE/KUR]
CH₃F₃SeSi	[753-96-8] $\Delta_v H$	silyl trifluoromethyl selenide	28.0	245		[1962EBS/EME]
C₂BrF₅Se	[6123-59-7] $\Delta_v H$	(pentafluoroethane) selenyl bromide	34.5	267		[1999DYK/SVO]
C₂ClF₅Se	[6123-50-8] $\Delta_v H$	(pentafluoroethane) selenyl chloride	30.3	252		[1999DYK/SVO]
C₂F₃NOSe	[20334-48-9] $\Delta_v H$	trifluoromethyl selenium isocyanate	29.5	259		[1968WEL/WUL]
C₂F₃NSSe	[21438-06-2] $\Delta_v H$	trifluoromethyl selenium thiocyanate	25.9	258		[1968WEL/WUL]
C₂F₃NSSe	[691-07-6] $\Delta_v H$	trifluoromethane sulphenyl selenocyanate	33.3	288		[1963EME/HAA]
C₂F₃NSe	[1717-49-3] $\Delta_v H$ $\Delta_v H$	trifluoromethyl selenocyanate	37.9 37.6	253		[1980GOM/WEI] [1968WEL/WUL]
C₂F₃NSe₂	[20563-91-1] $\Delta_v H$	trifluoromethyl selenium selenocyanate	26.6	245		[1968WEL/WUL]
C₂F₆Se	[371-79-9] $\Delta_v H$	bis(trifluoromethyl)selenide	24.4			[1980GOM/WEI]
CF₆Se₂	[372-65-6]	bis(difluoromethyl) diselenide				

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound			Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)		
	$\Delta_v H$		33.1			[1980GOM/WEL]
$C_2H_3F_3Se$	[1544-45-2] $\Delta_v H$	methyl(trifluoromethyl)selenide (209–294)	27.7	251		[1999DYK/SVO, 1963EME/WEL]
C_2H_6Se	[593-79-3] $\Delta_{\text{fus}} H$	dimethyl selenide	8.5	185.1		[1991RAB/SHE]
	$\Delta_v H$	(280–318)	30.3 ± 0.1	295		[1999DYK/SVO, 1997BAE]
	$\Delta_v H$	(278–313)	31.9	295	I	[1994KAR/FRA]
$C_2H_6Se_2$	[7101-31-7] $\Delta_{\text{fus}} H$	dimethyl diselenide	8.55	190.8		[1991RAB/SHE]
	$\Delta_v H$	(288–313)	74.9	300	I	[1994KAR/FRA]
	$\Delta_v H$		42.0 ± 1.0	298	C	[1989VOR/KLY]
C_3AsF_9Se	[816-45-5] $\Delta_v H$	bis(trifluoromethyl) trifluoromethylselenoarsine (227–295)	34.8	261		[1962EME/PAC]
C_3BrF_7Se	[662-44-2] $\Delta_v H$	(heptafluoro-1-propane) selenyl bromide (251–298)	35	274		[1999DYK/SVO, 1963EME/WEL]
C_3ClF_7Se	[662-46-4] $\Delta_v H$	(heptafluoro-1-propane) selenyl chloride (223–289)	33.4	256		[1999DYK/SVO, 1963EME/WEL]
C_3F_5NSe	[20334-51-4] $\Delta_v H$	pentafluoroethyl selenocyanate (254–293)	32.0	273		[1968WEL/WUL]
$C_3H_2F_6Se_2$	[691-25-8] $\Delta_v H$	bis[(trifluoromethyl)seleno]methane (273–359)	35.4	315		[1999DYK/SVO, 1963EME/WEL]
$C_3H_3F_5Se$	[6123-56-4] $\Delta_v H$	methyl pentafluoroethyl selenide (234–286)	31.9	260		[1999DYK/SVO]
$C_3H_3F_7SeSi$	[1647-59-2] $\Delta_v H$	(heptafluoropropyl)selenyl silane (233–393)	33.1	263		[1999DYK/SVO, 1962EBS/EME]
$C_3H_4F_5NSe$	[6123-53-1] $\Delta_v H$	(pentafluoroethyl)seleno methylamine (243–318)	33.8	280		[1999DYK/SVO]
C_3H_5FOSe	[367-52-2] $\Delta_v H$	fluoroselenoacetic acid, Se-methyl ester (273–333)	46.3	303		[1999DYK/SVO]
$C_3H_5F_3Se$	[690-25-5] $\Delta_v H$	ethyl(trifluoromethyl)selenide (223–254)	31.6	238		[1999DYK/SVO, 1963EME/WEL]
$C_3H_6F_3NSe$	[690-32-4] $\Delta_v H$	N,N-dimethyl(trifluoromethyl)selenenamide (231–321)	28.1	276		[1963EME/WEL]
$C_4F_{10}Se$	[6123-61-1] $\Delta_v H$	bis(pentafluoroethyl)selenide (232–295)	31.6	263		[1999DYK/SVO]
$C_4F_{10}Se_2$	[6123-49-5] $\Delta_v H$	bis(pentafluoroethyl) diselenide (272–318)	40.0	295		[1999DYK/SVO]
$C_4HF_{10}NSe_2$	[6123-55-3] $\Delta_v H$	bis[(pentafluoroethyl)seleno]amine (270–322)	38.3	296		[1999DYK/SVO]
$C_4H_3F_7Se$	[662-45-3] $\Delta_v H$	methyl(heptafluoropropyl) selenide (232–324)	30.8	278		[1999DYK/SVO, 1963EME/WEL]
$C_4H_4N_2O_2Se$	[92754-59-1]	selenobarbituric acid				

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{sub}}H$	(449–486)	141 ± 4.0	466	TE	[1999BRU/PIA]
C ₄ H ₄ Se	[288-05-1]	selenophene				
	$\Delta_{\text{sub}}H$	(208–243)	47.1	225		[1951MIL/PAO]
	Δ_vH	(234–300)	40.8	272		[1999DYK/SVO]
	Δ_vH		38.1 ± 0.7	298	C	[1989VOR/KLY]
	Δ_vH	(234–387)	34.0	249		[1947STU]
C ₄ H ₅ F ₅ Se	[6123-57-5]	ethyl(pentafluoroethyl) selenide				
	Δ_vH	(241–311)	34.8	276		[1999DYK/SVO]
C ₄ H ₆ F ₅ NSe	[6123-52-0]	1,1,2,2,2-pentafluoro-N,N-dimethylethane selenamide				
	Δ_vH	(256–320)	34.8	288		[1999DYK/SVO]
C ₄ H ₆ Se	[57796-75-5]	divinyl selenide				
	Δ_vH		42.0 ± 1.0	298	C	[1989VOR/KLY]
C ₄ H ₈ OSe	[5368-46-7]	1,4-oxaselenane				
	Δ_vH	(352–429)	46.6	367		[1999DYK/SVO]
C ₄ H ₁₀ Se	[627-53-2]	diethyl selenide				
	Δ_vH	(243–381)	39.7	258		[1999DYK/SVO]
	Δ_vH		38.9 ± 1.0	298	C	[1989VOR/KLY]
	Δ_vH		38.9 ± 4.2			[1936MER/SCH, 1982PIL/SKI]
	Δ_vH	(298–308)	36.8	303		[1929TAN/NAG]
C ₄ H ₁₀ Se ₂	[628-39-7]	diethyl diselenide				
	Δ_vH		47.1 ± 0.9	298	C	[1989VOR/KLY]
C ₅ AsF ₁₃ Se	[679-01-6]	heptafluoropropylseleno bis(trifluoromethyl)arsine				
	Δ_vH	(277–348)	40.3	312		[1962EME/PAC]
C ₅ H ₃ F ₁₀ NSe ₂	[6123-54-2]	N,N-bis[(pentafluoroethyl)seleno]methylamine				
	Δ_vH	(282–324)	38.3	303		[1999DYK/SVO]
C ₅ H ₅ F ₇ Se	[755-44-2]	ethyl(heptafluoropropyl) selenide				
	Δ_vH	(243–333)	36.0	288		[1999DYK/SVO, 1963EME/WEL]
C ₅ H ₆ F ₇ NSe	[755-79-3]	N,N-dimethyl(heptafluoropropyl)selenenamide				
	Δ_vH	(228–321)	30.8	274		[1963EME/WEL]
C ₆ F ₁₄ Se	[755-81-7]	bis(heptafluoropropyl) selenide				
	Δ_vH	(228–343)	34.5	286		[1999DYK/SVO, 1963EME/WEL]
C ₆ F ₁₄ Se ₂	[755-51-1]	bis(heptafluoropropyl) diselenide				
	Δ_vH	(260–348)	37.7	304		[1999DYK/SVO, 1963EME/WEL]
C ₆ H ₆ Se	[645-96-5]	benzene selenol				
	Δ_vH	(331–458)	45.4	395		[1999DYK/SVO]
C ₆ H ₁₄ Se	[37773-02-7]	diisopropyl selenide				
	Δ_vH		43.1 ± 1.0	298	C	[1989VOR/KLY]
C ₇ H ₈ Se	[4346-64-9]	methyl phenyl selenide				
	Δ_vH	(273–291)	52.5	282		[1999DYK/SVO]
C ₈ H ₆ N ₂ Se	[25660-64-4]	4-phenyl-1,2,3-selenadiazole				
	$\Delta_{\text{sub}}H$	(275–343)	91.2 ± 1.7	309	ME	[1974ARS]
	$\Delta_{\text{sub}}H$		94.1 ± 0.8	298	GS	[1973ARS/SHA]
	$\Delta_{\text{sub}}H$	(327–345)	90.7	336	A	[1987STE/MAL]

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound		T_m (K)	Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)			
C₈H₁₈Se	[14835-66-6] $\Delta_v H$	dibutyl selenide	47.3 ± 1.0	298	C	[1989VOR/KLY]
C₁₀H₂₂Se	[14835-67-7] $\Delta_v H$	dipentyl selenide	51.9 ± 1.0	298	C	[1989VOR/KLY]
C₁₂H₁₀Se	[1132-39-4] $\Delta_{\text{sub}} H$	diphenyl selenide	(302–324) 116.7 ± 2.5	313	ME	[1980MOR/WAT]
	$\Delta_v H$		(379–575) 63.4	394		[1999DYK/SVO]
	$\Delta_v H$		(378–575) 61.9	393	A	[1987STE/MAL]
	$\Delta_v H$		63.6 ± 2.5			[1973BAR/MOR, 1982PIL/SKI]
C₁₄H₁₄Se₂	[1482-82-2] $\Delta_{\text{sub}} H$	dibenzyl diselenide	(291–330) 130.5		ME	[1974ARS, 1973ARS/SHA]
Cl₂OSe	[7791-23-3] $\Delta_v H$	selenium oxychloride	(352–476) 59.1	367		[1999DYK/SVO]
	$\Delta_v H$		(353–453) 46.9	403		[1971NIS/TRE]
D₂Se	[13536-95-3] $\Delta_v H$	hydrogen selenide-d ₂	(202–256) 22.2	217		[1999DYK/SVO]
F₂OSe	[7783-43-9] $\Delta_{\text{fus}} H$	seleninyl difluoride	8.08	288		[1979CAR/CLA, 1977BOU/CAR]
	$\Delta_v H$		(316–420) 52.1	331		[1999DYK/SVO]
	$\Delta_v H$		46.9 ± 0.8	298	C	[1979CAR/CLA]
	$\Delta_v H$		46.7			[1977BOU/CAR]
F₄Se	[13465-66-2] $\Delta_v H$	selenium tetrafluoride	(297–398) 46.4	312		[1999DYK/SVO]
F₆Se	[7783-79-1] $\Delta_{\text{sub}} H$	selenium hexafluoride	24.96 ± 0.04	205	C	[1996OHT/OSA]
	$\Delta_{\text{sub}} H$		(194–226) 23.5	210		[1932KLE/HEN]
F₆O₂Se	[27069-91-6] $\Delta_v H$	<i>trans</i> bis(fluoroxy) tetrafluoroselenium	(241–286) 26.5	263		[1970SMI/CAD]
Si (silicon)						
CH₂Cl₄OSi	[18157-08-9] $\Delta_v H$	chloromethoxytrichlorosilane	(273–323) 9.3	288		[1958FRO/ROC]
CH₃Cl₃Si	[75-79-6] $\Delta_{\text{fus}} H$	methyltrichlorosilane	8.95	197.4		[1996DOM/HEA]
	$\Delta_v H$		(328–358) 30.7	343		[1967GOL/LAP]
	$\Delta_v H$		(287–337) 31.2	302	I	[1954JEN/CHA]
	$\Delta_v H$		31.0 ± 2.1			[1969AGA/HAJ, 1982PIL/SKI]
(CH₃Cl₃Si)₂–(C₆H₁₅N₃)	[na] $\Delta_{\text{sub}} H$	<i>bis</i> -1,3,5-trimethyl-1,3,5-triazacyclohexane- methyltrichlorosilane	(298–354) 74.0 ± 2.8			[1984GOL/LEV]
CH₃NSi	[18081-38-4] $\Delta_{\text{sub}} H$	isocyanosilane	(253–304) 48.8	279	A	[1987STE/MAL, 1956MAC]
CH₄Cl₂Si	[42430-97-7] $\Delta_v H$	(dichloromethyl)silane	(283–319) 32.5	301		[1957KAE/STO]
CH₄Cl₂Si	[75-54-7]	methyldichlorosilane				

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(275–314)	28.3	290	I	[1954JEN/CHA]
CH ₅ BrSi	[1631-88-5] $\Delta_v H$	methylbromosilane (283–295)	28.5	289		[1958EBS/EME]
CH ₅ ClSi	[10112-09-1] $\Delta_v H$	(chloromethyl)silane (246–297)	27.5	271		[1957KAE/STO]
CH ₆ OSi	[2171-96-2] $\Delta_v H$	methoxysilane (184–216)	25.8	201		[1961STE/MAC]
CH ₈ Si ₂	[13498-43-6] $\Delta_v H$	methyldisilane (190–273)	26.8	231	T	[1966ABE/MAC]
C ₂ H ₃ Cl ₃ Si	[75-94-5] $\Delta_v H$	trichlorovinylsilane (291–356)	34.2	306	I	[1954JEN/CHA]
C ₂ H ₃ Cl ₅ Si	[684-00-4] $\Delta_v H$	1,2-dichloroethyltrichlorosilane (375–453)	45.7	390	I	[1954JEN/CHA]
C ₂ H ₄ Cl ₆ Si ₂	[2504-64-5] $\Delta_v H$	bis(trichlorosilyl)ethane (364–432)	48.6	379	I	[1954JEN/CHA]
C ₂ H ₄ Si	[1066-27-9] $\Delta_v H$	silylacetylene (215–251)	22.0	233		[1963EBS/FRA]
C ₂ H ₅ Cl ₃ Si	[115-21-9] $\Delta_{\text{fus}} H$	ethyltrichlorosilane	6.96	165.3		[1996DOM/HEA]
	$\Delta_v H$	(303–363)	35.1	318		[1970SOK/KAR]
	$\Delta_v H$	(301–368)	35.9	316	I	[1954JEN/CHA]
C ₂ H ₅ F ₃ OSi	[460-55-9] $\Delta_v H$	ethoxytrifluorosilane (206–248)	26.8	227		[1949EME/HEA]
C ₂ H ₃ F ₃ O ₂ Si	[6876-44-4] $\Delta_v H$	silyl trifluoroacetate (273–293)	30.7	283	A	[1987STE/MAL, 1967EBS/THO]
C ₂ H ₆ Cl ₂ Si	[75-78-5] $\Delta_{\text{fus}} H$	dichlorodimethylsilane	8.83	199		[1996DOM/HEA]
	$\Delta_v H$	(301–345)	31.5	316	I	[1954JEN/CHA]
C ₂ H ₆ Cl ₂ Si	[1789-58-8] $\Delta_v H$	dichloroethylsilane (279–346)	31.5	294	I	[1954JEN/CHA]
	$\Delta_v H$	(301–345)	31.6	316		[1954DAV/JEN]
C ₂ H ₆ Cl ₄ Si ₂	[4518-98-3] $\Delta_v H$	1,1,2,2-tetrachloro-1,2-dimethyldisilane (300–375)	42.4	337		[1967REE/URR]
C ₂ H ₆ F ₃ NSi	[812-14-6] $\Delta_v H$	1,1,1-trifluoro-N,N-dimethylaminosilane (225–288)	28.5	273		[1961GRO/KLE]
C ₂ H ₉ NSi	[2875-98-1] $\Delta_{\text{sub}} H$	dimethylaminosilane (228–264)	58.8	246	A	[1987STE/MAL, 1954SUJ/WIT]
C ₂ H ₁₀ Si ₂	[870-26-8] $\Delta_v H$	1,2-dimethyldisilane (227–273)	25.4	258		[1962CRA/MAC]
C ₂ H ₁₁ NSi ₂	[14396-26-0] $\Delta_v H$	N,N-dimethyldisilanylamine (207–273)	35.4	240	T	[1963ABE/MAC]
C ₃ H ₄ Cl ₃ NSi	[2621-01-4]	trichloro- β -cyanoethylsilane				

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound		T_m (K)	Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)			
	$\Delta_v H$	(343–443)	53.5	358		[1978SHM/SHL]
$C_3H_4Cl_3NSi$	[1071-22-3] $\Delta_{\text{fus}} H$	β -trichlorosilylpropionitrile	21.24	307.9		[1975KOS/SAM]
$C_3H_5Cl_3Si$	[107-37-9] $\Delta_v H$	allyltrichlorosilane	40.1	333	I	[1954JEN/CHA]
$C_3H_6Cl_4Si$	[2550-06-3] $\Delta_v H$ $\Delta_v H$	γ -chloropropyltrichlorosilane	49.7 46.4	328 375		[1972SOK/BRA] I [1954JEN/CHA]
$C_3H_6Cl_4Si$	[na] $\Delta_v H$	β -chloropropyltrichlorosilane	46.9	328		[1972SOK/BRA]
$C_3H_8Cl_2OSi$	[1825-75-8] $\Delta_v H$ $\Delta_v H$	dichloroethoxymethylsilane	45.4 38.0	328 254	EB	[2010DON/WU] [1947STU]
C_3H_9BrSi	[2857-97-8] $\Delta_v H$	bromotrimethylsilane	32.6 ± 2.1			[1967BAL/LAP, 1982PIL/SKI]
C_3H_9ClSi	[75-77-4] $\Delta_{\text{us}} H$ $\Delta_{\text{fus}} H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$	chlorotrimethylsilane	0.7 9.68 30.8 30.2 30.1 ± 1.7	185.1 218 289 291		[1996DOM/HEA] [1964CAP/FRI] [1954JEN/CHA] [1967BAL/LAP, 1982PIL/SKI]
$C_3H_{10}OSi$	[1066-40-6] $\Delta_v H$ $\Delta_v H$ $\Delta_v H$	trimethylsilanol	46.8 45.6 ± 1.7 44.2	306 324	A I	[1987STE/MAL] [1969AGA/HAJ, 1982PIL/SKI] [1953GRU/OST]
$C_3H_{11}NSi$	[74897-01-1] $\Delta_v H$	N,N-dimethyl(methylsilyl)amine	28.2	296		[1958EBS/EME]
$C_3H_{13}NSi_2$	[18145-61-4] $\Delta_v H$	N-methyldi(methylsilyl)amine	32.2	327		[1958EBS/EME]
$C_3H_{15}NSi_3$	[18145-64-7] $\Delta_v H$	tri(methylsilyl)amine	33.7	350		[1958EBS/EME]
$C_4H_2Cl_6SSi_2$	[18145-50-1] $\Delta_v H$	2,5-bis(trichlorosilyl)thiophene	55.6	388		[1981DIT/SKO]
$C_4H_8Cl_2Si$	[10138-21-3] $\Delta_v H$	dichloroethylvinylsilane	38.1	333	I	[1954JEN/CHA]
$C_4H_9Cl_3Si$	[5936-98-1] $\Delta_{\text{us}} H$ $\Delta_{\text{fus}} H$	(trichloromethyl)trimethylsilane	11.16 7.36	285.3 405.3	DTA,DSC	[1994BRA/DOU]
$C_4H_9Cl_3Si$	[18171-74-9] $\Delta_{\text{us}} H$ $\Delta_{\text{fus}} H$	<i>tert</i> -butyltrichlorosilane	6.01 6.69	206.4 373.4	DTA,DSC	[1994BRA/DOU]
$C_4H_9F_6NSi_2$	[28245-41-2] $\Delta_v H$	1,1,1-trifluoro-N-(1-methylpropyl)-N-(trifluorosilyl)silanamine	25.8	213		[1973BEC/RUC]
$C_4H_{10}Cl_2Si$	[1719-53-5]	dichlorodiethylsilane				

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound			Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)		
	$\Delta_v H$	(321–401)	39.2	336	I	[1954JEN/CHA]
$C_4H_{10}F_3NSi$	[28245-37-6] $\Delta_v H$	(N,N-diethylamino)trifluorosilane (208–274)	27.4	241		[1974DIT/SKO3]
$C_4H_{10}F_3NSi$	[28245-40-1] $\Delta_v H$	(N- <i>tert</i> -butylamino)trifluorosilane (208–250)	33.6	229		[1973AYL/ELL]
$C_4H_{10}Si$	[765-33-3] $\Delta_v H$	1-methylsilacyclobutane	25.1	298	C	[1991VOR/KLY3]
$C_4H_{12}Si$	[75-76-3] $\Delta_{\text{fus}} H$	tetramethylsilane	6.74	174		[1996DOM/HEA]
$C_4H_{12}Si$	[542-91-6] $\Delta_v H$	diethylsilane	30.0 ± 0.4			[1972PED/ISE, 1982PIL/SKI]
$C_4H_{12}Cl_2OSi_2$	[2401-73-2] $\Delta_v H$	1,3-dichlorotetramethyldisiloxane (303–403)	40.3	318		[1971SOK/KAR]
$C_4H_{12}O_3Si$	[1185-55-3] $\Delta_v H$ $\Delta_v H$	methyltrimethoxysilane	34.3 ± 0.6 34.3 ± 0.3	298 298	C EB	[1988VOR/BAR] [1985KLY/DAN]
$C_4H_{12}O_4Si$	[681-84-5] $\Delta_v H$ $\Delta_v H$ $\Delta_v H$ $\Delta_v H$	tetramethoxysilane (364–393)	38.0 41.4 ± 0.7 41.4 ± 0.2 41.0	379 298 298 324	EB C EB	[1989KAT/TAN] [1988VOR/BAR] [1985KLY/DAN] [1980THO/SMI]
$C_4H_{12}Si$	[75-76-3] $\Delta_v H$ $\Delta_v H$ $\Delta_v H$	tetramethylsilane	26.0 ± 0.6 26.2 ± 0.4 24.2 ± 0.1	298 298 299	C C C	[1988VOR/BAR] [1972PED/ISE, 1982PIL/SKI] [1941AST/KEN]
$C_4H_{12}S_4Si$	[3931-76-8] $\Delta_{\text{us}} H$ $\Delta_{\text{fus}} H$	tetra(methylthia)silane	11.63 2.18	288.6 304.5		DSC [1998FUE/STR]
$C_4H_{13}NSi$	[16513-17-0] $\Delta_v H$ $\Delta_v H$	N,1,1,1-tetramethylsilanamine	37.4 ± 0.8 36.0 ± 2.1	298	C	[1991VOR/KLY] [1967BAL/LAP, 1982PIL/SKI]
$C_4H_{14}N_2Si$	[4693-04-3] $\Delta_v H$	<i>bis</i> (dimethylamino)silane (288–344)	32.4	316	T	[1964AYL/PET]
$C_4H_{16}N_2Si_2$	[18148-05-5] $\Delta_v H$	N,N,N',N'-tetramethyldisilanyldiamine (311–354)	39.3	332	T	[1963ABE/MAC]
$C_5H_6Cl_2SSi$	[18243-72-6] $\Delta_v H$	2-(methylchlorosilyl)thiophene (341–467)	46.4	356		[1981DIT/SKO]
$C_5H_9F_6NOSSi$	[34556-30-4] $\Delta_v H$	S,S- <i>bis</i> (trifluoromethyl)-N-(trimethylsilyl)sulfoximine	33.5	378	I	[1972SAU/SHR]
$C_5H_9F_6PSSi$	[38680-96-5] $\Delta_v H$	<i>bis</i> (trifluoromethyl)(trimethylsilylthio)phosphine (273–328)	46.6	301		[1973GOS/MIL]
$C_5H_{10}F_3NSi$	[33552-49-7] $\Delta_v H$	1-(trifluorosilyl)piperidine (250–282)	33.9	266		[1973AYL/ELL]
$C_5H_{12}Si$	[na]	1,2-dimethylsilacyclobutane				

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)			
	$\Delta_v H$		33.1	298		C	[1991VOR/KLY3]
			Note: The authors list the compound as 1,2-dimethylsilacyclobutane in Table 1 and as 1,1-dimethylsilacyclobutane in Table 2 of their paper.				
C₅H₁₂Si	[2295-12-7] $\Delta_{\text{fus}} H$	1,1-dimethylsilacyclobutane	6.76	155.5			[1975GUS/KAR]
	$\Delta_v H$		32.1	356			[1975GUS/KAR]
	$\Delta_v H$		33.0 ± 0.8	298		I	[1974BES/MAR]
	$\Delta_v H$		34.7 ± 2.1				[1972PED/ISE, 1982PIL/SKI]
C₅H₁₂Si	[754-05-2] $\Delta_{\text{fus}} H$	vinyltrimethylsilane	7.66	141.7			[1996DOM/HEA]
	$\Delta_v H$		33.1 ± 0.6	298		C	[1988VOR/BAR]
C₅H₁₃NSi	[216-90-7] $\Delta_{\text{fus}} H$	trimethylsilylethyleneimine	10.5	192.5			[1999KUL/LEB]
C₅H₁₄OSi	[1825-62-3] $\Delta_v H$	ethoxytrimethylsilane	38.4 ± 0.6	298		C	[1988VOR/BAR]
	$\Delta_v H$		38.4 ± 0.3	298		EB	[1985KLY/DAN]
	$\Delta_v H$	(223–349)	35.1	238			[1947STU]
C₅H₁₄O₃SSi	[57557-66-1] $\Delta_v H$	trimethoxy[(methylthio)methyl]silane	40.2 ± 0.6	298		C	[1989VOR/SOR]
C₅H₁₄Si	[760-32-7] $\Delta_v H$	methyldiethylsilane	34.6 ± 0.7	298		C	[1988VOR/BAR]
C₅H₁₅NSi	[2083-91-2] $\Delta_v H$	pentamethylsilanamine	33.6 ± 0.8	298		C	[1991VOR/KLY]
	$\Delta_v H$		31.8 ± 1.7				[1967BAL/LAP, 1982PIL/SKI]
	$\Delta_v H$	(313–357)	31.7	335			[1958EBS/EME]
C₅H₂₀O₅Si₅	[6166-86-5] $\Delta_v H$	1,3,5,7,9-pentamethylcyclopentasiloxane	47.0 ± 0.9	298		C	[1991VOR/KLY2]
C₆H₄Cl₄Si	[2003-90-9] $\Delta_v H$	(2-chlorophenyl)trichlorosilane	52.1	439		EB	[1974BES/MAR]
C₆H₄Cl₄Si	[2003-89-6] $\Delta_v H$	(3-chlorophenyl)trichlorosilane	50.7	430		EB	[1974BES/MAR]
C₆H₅Cl₃Si	[98-13-5] $\Delta_{\text{fus}} H$	phenyl trichlorosilane	11.66	233.4			[1996DOM/HEA]
	$\Delta_v H$	(333–453)	51.1	348			[1970SOK/KAR]
	$\Delta_v H$	(375–470)	47.9	390		I	[1954JEN/CHA]
C₆H₅F₃Si	[368-47-8] $\Delta_v H$	trifluorophenylsilane	40.1	257			[1947STU]
C₆H₈Cl₄SSi₂	[4480-01-7] $\Delta_v H$	2,5-bis(methyldichlorosilyl)thiophene	55.7	420			[1981DIT/SKO]
C₆H₈Si	[694-53-1] $\Delta_{\text{fus}} H$	phenylsilane	8.4	201			[2006ZEL/CHU]
	$\Delta_v H$	(238–390)	36.6 ± 0.3	314			[2006ZEL/CHU]
C₆H₉F₆NSi	[17599-55-2]	1,1,1-trimethyl-N-[2,2,2-trifluoro-1-(trifluoromethyl)ethylidene]silanamine					

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound		T_m (K)	Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)			
	$\Delta_v H$		30.5	358	I	[1972SWI/BAB]
$C_6H_{10}Cl_2Si$	[3651-23-8] $\Delta_v H$	diallyldichlorosilane (254–390)	47.9	269		[1947STU]
$C_6H_{11}NSi_2$	[4459-07-8] $\Delta_v H$	N-phenyldisilazane (298–356)	34.9	327	T	[1969AYL/HAK]
$C_6H_{12}Si$	[6224-91-5] $\Delta_{\text{fus}} H$	1-trimethylsilyl-1-propyne	10.96	204.5		[1993KUL/LEB, 1997LEB/KUL]
$C_6H_{12}Si$	[10654-12-9] $\Delta_{\text{fus}} H$	1,1-dimethyl-1-silacyclopent-3-ene	7.77	166.8		[2000BYK/LEB]
$C_6H_{12}Si$	[3514-67-8] $\Delta_v H$	1-methyl-1-vinylsilacyclobutane	33.1	298	C	[1991VOR/KLY3]
$C_6H_{12}Si_2$	[1627-98-1] $\Delta_v H$	1,1,3,3-tetramethyl-1,3-disilacyclobutane	39.5	391		[1975GUS/KAR]
$C_6H_{14}Si$	[30681-90-4] $\Delta_v H$	1,1,2-trimethylsilacyclobutane	36.0	298	C	[1991VOR/KLY3]
$C_6H_{14}Si$	[1072-54-4] $\Delta_v H$	1,1-dimethylsilacyclopentane	37.7 ± 2.1			[1972PED/ISE, 1982PIL/SKI]
$C_6H_{14}Si$	[2295-13-8] $\Delta_v H$	1,1,3-trimethylsilacyclobutane	35.5	298	C	[1991VOR/KLY3]
$C_6H_{15}ClSi$	[994-30-9] $\Delta_v H$	chlorotriethylsilane (268–419)	42.9	419		[1947STU]
$C_6H_{15}ClSi$	[18162-48-6] $\Delta_{\text{us}} H$ $\Delta_{\text{fus}} H$	<i>tert</i> -butyldimethylchlorosilane	7.75 5.6	203.6 358.1	DTA,DSC	[1994BRA/DOU]
$C_6H_{15}FO_3Si$	[358-60-1] $\Delta_v H$	triethoxyfluorosilane (291–373)	40.3	332	I	[1949EME/HEA]
$C_6H_{15}NOSi_2$	[na] $\Delta_v H$	pentamethylsilyl isocyanate (320–377)	44.2	348		[1963URE/MAC]
$C_6H_{15}NSi_2$	[1560-29-8] $\Delta_v H$	pentamethylsilyl cyanide (335–402)	46.9	350		[1962CRA/URE]
$C_6H_{16}OSi$	[1825-63-4] $\Delta_v H$ $\Delta_v H$	propoxytrimethylsilane	34.3 ± 0.6 34.3 ± 0.3	298 298	C C	[1988VOR/BAR] [1985KLY/DAN]
$C_6H_{16}OSi$	[1825-64-5] $\Delta_v H$ $\Delta_v H$	isopropoxytrimethylsilane	31.8 ± 0.6 31.8 ± 0.4	298 298	C EB	[1988VOR/BAR] [1985KLY/DAN]
$C_6H_{16}OSi$	[597-52-4] $\Delta_v H$	triethylsilanol (298–413)	50.6	355	I	[1953GRU/OST]
$C_6H_{16}O_2Si$	[78-62-6] $\Delta_v H$ $\Delta_v H$ $\Delta_v H$	diethoxydimethylsilane	43.1 ± 0.7 43.1 ± 0.3 43.3	298 298 269	C EB	[1988VOR/BAR] [1985KLY/DAN] [1947STU]
$C_6H_{16}O_3SSi$	[66785-19-1]	trimethoxy[2-(methylthio)ethyl]silane				

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound		T_m (K)	Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)			
	$\Delta_v H$		45.2 ± 0.7	298	C	[1989VOR/SOR]
C₆H₁₆O₃SSi	[53696-79-0]	[(ethylthio)methyl]trimethoxysilane				
	$\Delta_v H$		41.4 ± 0.6	298	C	[1989VOR/SOR]
C₆H₁₆Si	[756-81-0]	dimethyldiethylsilane				
	$\Delta_v H$		38.9 ± 0.6	298	C	[1988VOR/BAR]
C₆H₁₆Si	[617-86-7]	triethylsilane				
	$\Delta_v H$		37.4 ± 0.6	298	C	[1988VOR/BAR]
	$\Delta_v H$	(303–373)	33.5		EB, I	[1975BRA/KAR]
	$\Delta_v H$		36.4 ± 1.3			[1972PED/ISE, 1982PIL/SKI]
C₆H₁₆Si₂	[1627-98-1]	1,1,3,3-tetramethyl-1,3-disilacyclobutane				
	$\Delta_{\text{fus}} H$		10.26	266		[1996DOM/HEA, 1975GUS/KAR]
	$\Delta_v H$		36.7 ± 1.1	298	I	[1974SHM/SHL]
	$\Delta_v H$		41.0 ± 2.1			[1972PED/ISE, 1982PIL/SKI]
C₆H₁₇B₅Br₂Si₂	[66798-29-6]	2,4-bis(bromodimethylsilyl)-2,4-dicarbo-closo-heptaborane (7)				
	$\Delta_v H$	(388–463)	53.1	403	I	[1979GOL/SHM]
C₆H₁₇B₅Cl₂Si₂	[28699-83-4]	2,4-bis(chlorodimethylsilyl)-2,4-dicarbo-closo-heptaborane (7)				
	$\Delta_v H$	(359–439)	46.2	374	I	[1979GOL/SHM]
C₆H₁₈Cl₂O₂Si₃	[na]	1,5-dichlorohexamethyltrisiloxane				
	$\Delta_v H$	(299–457)	49.8	314		[1947STU]
C₆H₁₈OSi₂	[107-46-0]	hexamethyldisiloxane				
	$\Delta_{\text{fus}} H$		11.92	204.9		[1996DOM/HEA]
	$\Delta_v H$	(300–383)	36.9	315	EB	[1986FLA]
	$\Delta_v H$	(293–361)	33.1	327		[1971DIT/SKO]
	$\Delta_v H$		37.2 ± 1.7			[1964GOO/LAC, 1982PIL/SKI]
	$\Delta_v H$	(309–411)	36	324	EB	[1961SCO/MES]
	$\Delta_v H$		34.6 ± 0.1	332	C	[1961SCO/MES]
	$\Delta_v H$		33.1 ± 0.1	351	C	[1961SCO/MES]
	$\Delta_v H$		31.3 ± 0.1	373	C	[1961SCO/MES]
	$\Delta_v H$		37.2 ± 1.7			[1947STU]
C₆H₁₈O₃Si₃	[541-05-9]	hexamethylcyclotrisiloxane				
	$\Delta_{\text{fus}} H$		11.8	335.2		
	$\Delta_{\text{fus}} H$		16.61	335.2		[1996DOM/HEA, 1971ALV/DAL]
	$\Delta_{\text{sub}} H$	(297–335)	55.2 ± 0.4	316		[1953OST/GRU]
	$\Delta_v H$	(342–419)	40.8	357	EB	[1986FLA]
	$\Delta_v H$	(353–403)	39.7	368		[1974BRA/KAR]
	$\Delta_v H$	(339–407)	39	373		[1971DIT/SKO]
	$\Delta_v H$	(343–388)	39.7	365		[1953OST/GRU]
C₆H₁₈Si₂	[1450-14-2]	hexamethyldisilane				
	$\Delta_{\text{fus}} H$		9.75	221.8		
	$\Delta_{\text{fus}} H$		3.02	287.7		[1996DOM/HEA]
	$\Delta_v H$	(305–387)	36.3	320	EB	[1986TAK/ISH]
	$\Delta_v H$		37.4 ± 0.4			[1972PED/ISE, 1982PIL/SKI]
	$\Delta_v H$	(288–310)	37.2			[1959SUG/SEK, 1986TAK/ISH]
	$\Delta_v H$	(294–334)	36.8			[1941BRO/DAV, 1986TAK/ISH]
C₆H₁₉B₅Si₂	[59351-11-0]	2,4-bis(dimethylsilyl)-2,4-dicarbo-closo-heptaborane				

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound		T_m (K)	Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)			
	$\Delta_v H$	(373–453)	41.3	388	I	[1976SHM/SHL]
C₆H₁₉NSi₂	[999-97-3]	hexamethyldisilazane				
	$\Delta_v H$		42.2 ± 0.9	298	C	[1991VOR/KLY]
	$\Delta_v H$	(294–395)	36.0	344		[1972DIT/SKO2]
	$\Delta_v H$		41.4 ± 2.1			[1966BEE/MOR, 1982PIL/SKI]
C₆H₁₉N₃Si	[na]	<i>tris</i> (dimethylamino)silane				
	$\Delta_v H$	(309–387)	41.1	348	T	[1964AYL/PET]
C₆H₂₁N₃Si₃	[1009-93-4]	hexamethylcyclotrisilazane				
	$\Delta_{\text{fus}} H$		15.17	254.4		[1996DOM/HEA]
	$\Delta_v H$	(342–456)	45.6	399		[1972DIT/SKO2]
C₇H₈Cl₂Si	[18173-99-4]	benzyl dichlorosilane				
	$\Delta_v H$	(318–467)	58.5	333		[1947STU, 1999DYK/SVO]
C₇H₈Cl₂Si	[149-74-6]	phenyldichloromethylsilane				
	$\Delta_v H$	(309–479)	51.2	323		[1947STU, 1999DYK/SVO]
C₇H₈Cl₂Si	[13272-80-5]	dichloro-4-tolylsilane				
	$\Delta_v H$	(319–469)	58	334		[1947STU, 1999DYK/SVO]
C₇H₈F₂Si	[13272-80-5]	difluoromethylphenylsilane				
	$\Delta_v H$	(303–413)	44.6	318		[1999DYK/SVO]
C₇H₉F₈NOSSi	[77589-40-3]	2,2,3,3,4,4,5,5-octafluoro-1,1,2,3,4,5-hexahydro-1-[(trifluoromethyl)silyl]imino]thiophene-1-oxide				
	$\Delta_v H$		31.4	383		[1981ABE/SHR2]
C₇H₉F₉N₂OSSi	[62609-67-0]	1,1,1-trifluoro-N-[2,2,2-trifluoro-1-(trifluoromethyl)ethylidene]-N'-(trimethylsilyl)methanesulfonimidamide				
	$\Delta_v H$		39.3	429	I	[1977KIT/SHR, 1999DYK/SVO]
C₇H₁₅NO₃Si	[2288-13-3]	1-methyl-2,8,9-trioxa-5-aza-1-silabicyclo[3.3.3]undecane				
	$\Delta_{\text{sub}} H$		82 ± 0.8			[1989VOR/BAR]
C₇H₁₆O₃SSi	[57877-58-4]	trimethoxy[(2-propenylthio)methyl]silane				
	$\Delta_v H$		38.6 ± 0.5	298	C	[1989VOR/SOR]
C₇H₁₇ClSi	[18817-17-9]	(1-chloroethyl)diethylmethylsilane				
	$\Delta_v H$	(353–445)	41.8	400		[1999DYK/SVO]
C₇H₁₇NSi	[na]	N-(β -trimethylsilyl)ethylenimine				
	$\Delta_{\text{fus}} H$		10.62	176.5		[1996DOM/HEA]
C₇H₁₈OSi	[1825-65-6]	butyl trimethylsilyl ether				
	$\Delta_v H$	(344–397)	38.5	359	EB	[1969SHE/LAN, 1984BOU/FRI]
C₇H₁₈O₃SSi	[94358-36-8]	trimethoxy[3-(methylthio)propyl]silane				
	$\Delta_v H$		43.5 ± 0.6	298	C	[1989VOR/SOR]
C₇H₁₈O₃SSi	[40532-52-3]	[2-(ethylthio)ethyl]trimethoxysilane				
	$\Delta_v H$		41.4 ± 0.7	298	C	[1989VOR/SOR]
C₇H₁₈O₃Si	[2031-67-6]	triethoxymethylsilane				
	$\Delta_v H$		45.1 ± 0.7	298	C	[1988VOR/BAR]
	$\Delta_v H$		45.1 ± 0.4	298	EB	[1985KLY/DAN]
	$\Delta_v H$	(272–416)	45.2	287		[1947STU]
C₇H₁₈SSi	[3553-78-4]	<i>n</i> -butylthio)trimethylsilane				
	$\Delta_v H$		40.6 ± 2.1			[1967BAL/LAP, 1982PIL/SKI]
C₇H₁₈Si	[757-21-1]	methyltriethylsilane				

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound		T_m (K)	Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)			
	$\Delta_v H$		40.5 ± 0.6	298	C	[1988VOR/BAR]
C₇H₁₈Si	[999-03-1]	methyldipropylsilane				
	$\Delta_v H$		35.9 ± 0.7	298	C	[1988VOR/BAR]
C₇H₁₈Si	[18442-00-7]	methyldiisopropylsilane				
	$\Delta_v H$		32.4 ± 0.8	298	C	[1988VOR/BAR]
C₇H₁₉NSi	[996-50-9]	N,N-diethyl-1,1,1-trimethylsilanamine				
	$\Delta_v H$		37.9 ± 0.8	298	C	[1991VOR/KLY]
C₇H₂₀Si₂	[2117-28-4]	methylene-bis(trimethylsilane)				
	$\Delta_v H$	(323–407)	40.3 ± 0.3	365	QM	[1975GUS/KUL, 1975GUS/KAR]
C₇H₂₀Si₂	[2117-28-4]	hexamethyldisilylmethane				
	$\Delta_{\text{fus}} H$		11.11	140.7		[1996DOM/HEA, 1975GUS/KAR]
C₇H₂₁NSi₂	[920-68-3]	N,1,1,1-tetramethyl-N-(trimethylsilyl)silanamine				
	$\Delta_v H$		38.1 ± 0.8	298	C	[1991VOR/KLY]
	$\Delta_v H$		38.9 ± 2.1			[1967BAL/LAP, 1982PIL/SKI]
C₈H₁₀Cl₂OSi	[18236-80-1]	dichloroethoxyphenylsilane				
	$\Delta_v H$	(325–496)	56.3	340		[1999DYK/SVO]
C₈H₁₀Cl₂Si	[1125-27-5]	dichloroethylphenylsilane				
	$\Delta_v H$	(316–503)	51.3	331		[1999DYK/SVO]
C₈H₁₁ClSi	[768-33-2]	chlorodimethylphenylsilane				
	$\Delta_v H$	(302–467)	52.2	317		[1999DYK/SVO]
	$\Delta_v H$	(303–466)	49.7	318		[1947STU]
C₈H₁₁FSi	[454-57-9]	fluorodimethylphenylsilane				
	$\Delta_v H$	(303–423)	49.6	318		[1999DYK/SVO]
C₈H₁₂Si	[1112-55-6]	tetravinylsilane				
	$\Delta_v H$		42.7 ± 0.7	298	C	[1988VOR/BAR]
C₈H₁₂Si	[766-77-8]	dimethylphenylsilane				
	$\Delta_v H$	(298–432)	45.3	293		[1947STU]
C₈H₁₅NO₃Si	[2097-18-9]	1-ethenyl-2,8,9-trioxa-5-aza-1-silabicyclo[3.3.3]undecane				
	$\Delta_{\text{sub}} H$		85 ± 0.8			[1989VOR/BAR]
C₈H₁₆Cl₄O₄Si	[18290-84-1]	tetrakis(2-chloroethoxy)silane				
	$\Delta_v H$	(447–500)	81.1	473		[1999DYK/SVO, 1946JON/THO]
C₈H₁₇NO₃Si	[2097-16-7]	1-ethyl-2,8,9-trioxa-5-aza-1-silabicyclo[3.3.3]undecane				
	$\Delta_{\text{sub}} H$		81 ± 0.9			[1989VOR/BAR]
C₈H₁₇NO₃Si	[18225-19-9]	1,7-dimethyl-2,8,9-trioxa-5-aza-1-silabicyclo[3.3.3]undecane				
	$\Delta_{\text{sub}} H$		92 ± 0.8			[1989VOR/BAR]
C₈H₁₇NO₄Si	[3463-21-6]	1-ethoxy-2,8,9-trioxa-5-aza-1-silabicyclo[3.3.3]undecane				
	$\Delta_{\text{sub}} H$		81 ± 0.8			[1989VOR/BAR]
C₈H₁₈F₃NOSi₂	[na]	CF ₃ C[OSi(CH ₃) ₃]=NSi(CH ₃) ₃				
	$\Delta_v H$	(316–350)	41.8	333		[1970VON/GLE]
C₈H₁₈O₃Si	[78-08-0]	vinyltriethoxysilane				
	$\Delta_v H$		50.2 ± 0.8	298	C	[1988VOR/BAR]
	$\Delta_v H$		50.2 ± 0.4	298	EB	[1985KLY/DAN]
	$\Delta_v H$	(334–421)	46.2	349	I	[1954JEN/CHA]

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)			
C ₈ H ₁₉ NSi	[42525-64-4] $\Delta_{\text{fus}}H$	N-(β -trimethylsilylethyl)trimethylenimine (7–305)	12.9	199.4		AC	[1996DOM/HEA]
C ₈ H ₁₉ NSi	[na] $\Delta_{\text{fus}}H$	triethylsilylethyleneimine	14.25	183.6			[2001SMI/LEB]
C ₈ H ₂₀ Cl ₂ OSi ₂	[18825-03-1] Δ_vH	1,3-dichloro-1,1,3,3-tetraethyldisiloxane (343–463)	53.6	358			[1971SOK/KAR, 1999DYK/SVO]
C ₈ H ₂₀ O ₃ SSi	[57557-74-1] Δ_vH	[3-(ethylthio)propyl]trimethoxysilane	41.8 ± 0.6	298		C	[1989VOR/SOR]
C ₈ H ₂₀ O ₃ SSi	[57557-68-3] Δ_vH	[(butylthio)methyl]trimethoxysilane	41.6 ± 0.6	298		C	[1989VOR/SOR]
C ₈ H ₂₀ O ₃ SSi	[57557-69-4] Δ_vH	trimethoxy[[2-methylpropyl]thio]methylsilane	38.7 ± 0.6	298		C	[1989VOR/SOR]
C ₈ H ₂₀ O ₃ SSi	[57557-70-7] Δ_vH	[[1,1-dimethylethyl]thio]methyltrimethoxysilane	50.6 ± 0.7	298		C	[1989VOR/SOR]
C ₈ H ₂₀ O ₃ Si	[78-07-9] Δ_vH	ethyltriethoxysilane (338–426)	47.0	353		I	[1954JEN/CHA]
C ₈ H ₂₀ O ₄ Si	[78-10-4] $\Delta_{\text{us}}H$ $\Delta_{\text{fus}}H$ Δ_vH Δ_vH Δ_vH Δ_vH Δ_vH Δ_vH	tetraethoxysilane	13.2 11.14 53.9 52.3 40.9 48.5 ± 0.3 50.0 U 33.9 49.5	187.7 191 298 298 419 298 328 304			[1992VAN/COR] [1995VAN/COR] [1995VAN/COR] [1989KAT/TAN] [1985KLY/DAN] [1980THO/SMI] [1973DYA/VIG] [1947STU]
C ₈ H ₂₀ Si	[995-89-1] Δ_vH	dimethyldipropylsilane	40.2 ± 0.6	298		C	[1988VOR/BAR]
C ₈ H ₂₀ Si	[631-36-7] $\Delta_{\text{fus}}H$ Δ_vH Δ_vH Δ_vH	tetraethylsilane	13.01 39.0 ± 0.7 39.7 ± 2.1 43.3	189.4 298 287			[1990DOM/HEA] [1988VOR/BAR] [1972PED/ISE, 1982PIL/SKI] [1947STU]
C ₈ H ₂₀ Si	[998-14-1] Δ_vH	ethyldipropylsilane	37.9 ± 0.6	298		C	[1988VOR/BAR]
C ₈ H ₂₀ Si	[17591-40-1] Δ_vH	ethyldiisopropylsilane	38.1 ± 0.7	298		C	[1988VOR/BAR]
C ₈ H ₂₀ Si	[30736-07-3] Δ_vH	di- <i>tert</i> -butylsilane (242–288)	41.4	257			[2005FUL/RUZ]
C ₈ H ₂₁ NO ₃ Si	[na] Δ_vH	γ -aminopropyltriethoxysilane (363–492)	55.8	388			[1976DIT/SKO]
C ₈ H ₂₃ B ₅ Si ₂	[59351-10-9] Δ_vH	2,4- <i>bis</i> (trimethylsilyl)-2,4-dicarba-closo-heptaborane (373–473)	45.0	388		I	[1976SHM/SHL]
C ₈ H ₂₄ Cl ₂ O ₃ Si ₄	[2474-02-4]	1,7-dichloro-1,1,3,3,5,5,7,7-octamethyltetrasiloxane					

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound		T_m (K)	Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)			
	$\Delta_v H$	(326–495)	53.8	341		[1999DYK/SVO]
$C_8H_{24}N_4Si$	[1624-01-7]	tetrakis(dimethylamino)silane				
	$\Delta_v H$	(361–415)	40.0	388	T	[1964AYL/PET]
$C_8H_{24}O_2Si_3$	[107-51-7]	octamethyltrisiloxane				
	$\Delta_v H$	(346–446)	43.2	361	EB	[1986FLA]
	$\Delta_v H$		39.7 ± 2.1			[1972PED/ISE, 1982PIL/SKI]
	$\Delta_v H$	(345–417)	40.2	381		[1971SKO/DIT]
$C_8H_{24}O_4Si_4$	[556-67-2]	octamethylcyclotetrasiloxane				
	$\Delta_{\text{trs}} H$		4.87	258		
	$\Delta_{\text{fus}} H$		23.77	290.5		[1996DOM/HEA, 1975MEK/KAR, 1971ALV/DAL]
	$\Delta_{\text{sub}} H$		64 ± 2		B	[1953OST/GRU, 1960JON]
	$\Delta_v H$		57.0 ± 0.8	298	C	[1991VOR/KLY2]
	$\Delta_v H$	(361–469)	47.6	376	EB	[1986FLA]
	$\Delta_v H$	(334–423)	44.1	378		[1971DIT/SKO]
	$\Delta_v H$	(303–428)	56.1	298	I	[1954OST/GRU]
	$\Delta_v H$	(303–428)	48.5	373	I	[1954OST/GRU]
	$\Delta_v H$	(303–428)	45.6	398	I	[1954OST/GRU]
$C_8H_{24}O_{12}Si_8$	[57348-79-5]	octamethyldecaoxooctasilicon				
	$\Delta_{\text{sub}} H$	(463–563)	110.5	513	A	[1987STE/MAL, 1975TIT/CHU]
$C_8H_{24}Si_3$	[3704-44-7]	octamethyltrisilane				
	$\Delta_v H$		46.0 ± 0.8			[1972PED/ISE, 1982PIL/SKI]
$C_8H_{28}N_4Si_4$	[1020-84-4]	octamethylcyclotetrasilazane				
	$\Delta_{\text{fus}} H$		25.05	367.7		[1996DOM/HEA]
	$\Delta_v H$	(388–513)	52.3	450		[1972DIT/SKO2]
$C_9H_9F_5Si$	[1206-46-8]	pentafluorophenyl(trimethyl)silane				
	$\Delta_{\text{fus}} H$		8.4	233		[2006ZEL/CHU]
	$\Delta_v H$	(273–440)	40.6 ± 0.3	357		[2006ZEL/CHU]
$C_9H_{14}OSi$	[1529-17-5]	phenoxytrimethylsilane				
	$\Delta_v H$		56.9 ± 0.8	298	C	[1988VOR/BAR]
$C_9H_{19}NO_3Si$	[26053-77-0]	1-propyl-2,8,9-trioxa-5-aza-1-silabicyclo[3.3.3]undecane				
	$\Delta_{\text{sub}} H$		84 ± 0.8			[1989VOR/BAR]
$C_9H_{19}NO_3Si$	[2097-17-8]	1-(1-methylethyl)-2,8,9-trioxa-5-aza-1-silabicyclo[3.3.3]undecane				
	$\Delta_{\text{sub}} H$		92 ± 0.9			[1989VOR/BAR]
$C_9H_{19}NO_3Si$	[56492-01-4]	1,3,7-trimethyl-2,8,9-trioxa-5-aza-1-silabicyclo[3.3.3]undecane				
	$\Delta_{\text{sub}} H$		101 ± 0.8			[1989VOR/BAR]
$C_9H_{20}OSi$	[13871-89-1]	cyclohexyl trimethylsilyl ether				
	$\Delta_v H$	(364–441)	45.1	379	EB	[1969SHE/LAN]
$C_9H_{22}O_3SSi$	[53696-83-6]	triethoxy[(ethylthio)methyl]silane				
	$\Delta_v H$		42.3 ± 0.6	298	C	[1989VOR/SOR]
$C_9H_{22}Si$	[994-44-5]	propyltriethylsilane				
	$\Delta_v H$		40.0 ± 0.7	298	C	[1988VOR/BAR]
$C_9H_{22}Si$	[998-29-8]	tripropylsilane				

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)			
	$\Delta_v H$		39.1 ± 0.7	298		C	[1988VOR/BAR]
C ₉ H ₂₂ Si	[999-35-9]	methyl dibutylsilane					
	$\Delta_v H$		36.2 ± 0.7	298		C	[1988VOR/BAR]
C ₉ H ₂₃ NSi	[5277-20-3]	1,1,1-triethyl-N-(1-methylethyl)silanamine					
	$\Delta_v H$		38.6 ± 0.8	298		C	[1991VOR/KLY]
C ₉ H ₂₃ NSi	[17887-11-5]	1,1,1-triethyl-N-propylsilanamine					
	$\Delta_v H$		41.5 ± 0.8	298		C	[1991VOR/KLY]
C ₉ H ₂₄ Si ₂	[2295-05-8]	1,3-propanediyl-bis(trimethylsilane)					
	$\Delta_v H$	(338–443)	43.1 ± 0.5	390		QM	[1975GUS/KUL, 1975GUS/KAR]
C ₉ H ₂₄ Si ₂	[2295-05-8]	1,3-hexamethyl disilylpropane					
	$\Delta_{\text{fus}} H$		16.05	223.7			[1996DOM/HEA, 1975GUS/KAR]
C ₉ H ₂₄ Si ₃	[1627-99-2]	1,1,3,3,5,5-hexamethyl-1,3,5-trisilacyclocyclohexane					
	$\Delta_{\text{fus}} H$		16.5	269.3			[1996DOM/HEA, 1975GUS/KAR]
C ₉ H ₂₇ NSi ₃	[1586-73-8]	tris(trimethylsilyl)amine					
	$\Delta_{\text{us}} H$		7.9	244.2			
	$\Delta_{\text{fus}} H$		1.77	337.2			[1971MUR/BRE]
	$\Delta_v H$		54.4 ± 8.4				[1967BAL/LAP, 1982PIL/SKI]
C ₁₀ H ₉ F ₇ Si	[122571-42-0]	trimethyl(4-trifluoromethylphenyl)silane					
	$\Delta_v H$	(296–460)	47.4 ± 0.2	378			[2006ZEL/CHU]
C ₁₀ H ₁₄ Si	[3944-08-9]	1-phenyl-1-methyl-1-silacyclobutane					
	$\Delta_{\text{fus}} H$		12.28	210			[1996DOM/HEA]
C ₁₀ H ₁₄ Si	[1125-26-4]	vinyl dimethyl phenyl silane					
	$\Delta_{\text{fus}} H$		12.26	190.7			[1996DOM/HEA]
C ₁₀ H ₁₆ OSi	[704-43-8]	(2-methoxyphenyl)trimethylsilane					
	$\Delta_v H$		59.4 ± 0.8	298		C	[1988VOR/BAR]
C ₁₀ H ₁₆ OSi	[17876-90-3]	(3-methoxyphenyl)trimethylsilane					
	$\Delta_v H$		56.1 ± 0.8	298		C	[1988VOR/BAR]
C ₁₀ H ₁₆ OSi	[877-68-9]	(4-methoxyphenyl)trimethylsilane					
	$\Delta_v H$		56.9 ± 0.8	298		C	[1988VOR/BAR]
C ₁₀ H ₁₆ OSi	[17902-31-7]	<i>m</i> -tolyl trimethylsilyl ether					
	$\Delta_v H$	(371–398)	49.7	384		EB	[1969SHE/LAN]
C ₁₀ H ₁₆ OSi	[17902-32-8]	<i>p</i> -tolyl trimethylsilyl ether					
	$\Delta_v H$	(374–402)	49.8	388		EB	[1969SHE/LAN]
C ₁₀ H ₁₆ O ₃ SSi	[57557-71-8]	trimethoxy[(phenylthio)methyl]silane					
	$\Delta_v H$		56.4 ± 0.7	298		C	[1989VOR/SOR]
C ₁₀ H ₁₈ Si	[na]	5-trimethylsilyl-2-norbornene					
	$\Delta_{\text{fus}} H$		6.84	201.6			[1994LEB/SMI2]
C ₁₀ H ₂₀ O ₂ Si	[13081-67-9]	diallyl(diethoxy)silane					
	$\Delta_v H$	(342–459)	48.3	357		A	[1987STE/MAL]
C ₁₀ H ₂₁ NO ₃ Si	[71229-51-1]	1,3,7,10-tetramethyl-2,8,9-trioxa-5-aza-1-silabicyclo[3.3.3]undecane					
	$\Delta_{\text{sub}} H$		115 ± 0.9				[1989VOR/BAR]
C ₁₀ H ₂₄ O ₂ Si	[2031-63-2]	dipropyldiethoxysilane					

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No	Compound		T _m (K)	Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)			
	$\Delta_{\text{v}}H$		46.5 ± 0.7	298	C	[1988VOR/BAR]
	$\Delta_{\text{v}}H$		46.4 ± 0.3	298	EB	[1985KLY/DAN]
C ₁₀ H ₂₄ O ₃ SSi	[57557-72-9]	triethoxy[2-(ethylthio)ethyl]silane				
	$\Delta_{\text{v}}H$		46.9 ± 0.7	298	C	[1989VOR/SOR]
C ₁₀ H ₂₄ Si	[994-59-2]	diethyldipropylsilane				
	$\Delta_{\text{v}}H$		41.5 ± 0.7	298	C	[1988VOR/BAR]
C ₁₀ H ₂₄ Si	[995-24-4]	methyltripropylsilane				
	$\Delta_{\text{v}}H$		42.6 ± 0.6	298	C	[1988VOR/BAR]
C ₁₀ H ₂₄ Si	[998-61-8]	ethylidibutylsilane				
	$\Delta_{\text{v}}H$		39.9 ± 0.7	298	C	[1988VOR/BAR]
C ₁₀ H ₂₄ Si	[17591-42-3]	ethyl-diisobutylsilane				
	$\Delta_{\text{v}}H$		39.8 ± 0.7	298	C	[1988VOR/BAR]
C ₁₀ H ₂₅ NO ₂ Si ₃	[27602-22-8]	1,1,1,3,5,5,5-heptamethyl-3-(2-cyanoethyl)trisiloxane				
	$\Delta_{\text{v}}H$	(367–511)	59.5	382	A	[1987STE/MAL]
C ₁₀ H ₂₅ NSi	[6022-10-2]	pentaethylsilanamine				
	$\Delta_{\text{v}}H$		42.2 ± 1.0	298	C	[1991VOR/KLY]
C ₁₀ H ₂₅ NSi	[17940-20-4]	N-(1,1-dimethylethyl)-1,1,1-triethylsilanamine				
	$\Delta_{\text{v}}H$		40.3 ± 0.9	298	C	[1991VOR/KLY]
C ₁₀ H ₂₆ O ₃ Si ₃	[110505-51-6]	1,1,3,3-tetraethyl-5,5-dimethylcyclotrisiloxane				
	$\Delta_{\text{us}}H$		0.13	195		
	$\Delta_{\text{fus}}H$		9.52	260		[1996DOM/HEA]
C ₁₀ H ₂₈ O ₄ Si ₃	[17928-13-1]	1,5-diethoxy-1,1,3,3,5,5-hexamethyltrisiloxane				
	$\Delta_{\text{v}}H$	(314–470)	56.2	329	A	[1987STE/MAL]
C ₁₀ H ₃₀ OSi ₄	[1360-31-2]	bis[(pentamethyl)disilanyl] ether				
	$\Delta_{\text{v}}H$	(376–456)	49.3	376		[1962CRA/URE]
C ₁₀ H ₃₀ O ₃ Si ₄	[17928-28-8]	methyl tris(trimethylsiloxy)silane				
	$\Delta_{\text{v}}H$	(362–476)	49.5	377	EB	[1986FLA]
C ₁₀ H ₃₀ O ₃ Si ₄	[141-62-8]	decamethyl tetrasiloxane				
	$\Delta_{\text{v}}H$	(366–479)	50.3	381	EB	[1986FLA]
	$\Delta_{\text{v}}H$	(343–454)	48.2	358	A	[1987STE/MAL, 1971SKO/DIT]
C ₁₀ H ₃₀ O ₅ Si ₅	[541-02-6]	decamethyl cyclopentasiloxane				
	$\Delta_{\text{fus}}H$		20.37	226.2		[1971ALV/DAL]
	$\Delta_{\text{v}}H$		59.0 ± 1.0	298	C	[1991VOR/KLY2]
	$\Delta_{\text{v}}H$	(383–496)	52.1	398	EB	[1986FLA]
	$\Delta_{\text{v}}H$	(364–472)	49.0	379	A	[1987STE/MAL, 1971DIT/SKO]
	$\Delta_{\text{v}}H$		48.1 ± 2.1			[1972PED/ISE, 1982PIL/SKI]
C ₁₀ H ₃₀ Si ₄	[865-76-9]	decamethyltetrasilane				
	$\Delta_{\text{v}}H$		52.3 ± 1.7			[1972PED/ISE, 1982PIL/SKI]
C ₁₀ H ₃₁ NSi ₄	[1787-38-8]	1,1,3,3-tetramethyl-1,3-bis(trimethylsilyl)disilazane				
	$\Delta_{\text{v}}H$	(378–435)	58.0	393	A	[1987STE/MAL, 1963URE/MAC]
		Note: Molecular formula given in paper is not consistent with chemical name				
C ₁₁ H ₁₆ Si	[na]	vinyl dimethylbenzylsilane				
	$\Delta_{\text{fus}}H$		11.6	204.1		[1996DOM/HEA]

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
C ₁₁ H ₁₈ O ₃ SSi	[53696-80-3] $\Delta_{\text{v}}H$	trimethoxy[[phenylmethyl]thio]methylsilane	56.1 ± 0.7	298	C	[1989VOR/SOR]
C ₁₁ H ₂₀ OSi	[17962-20-8] $\Delta_{\text{v}}H$	triallylethoxy silane	48.4	364	A	[1987STE/MAL]
C ₁₁ H ₂₀ OSi ₂	[14920-92-4] $\Delta_{\text{v}}H$ $\Delta_{\text{v}}H$	pentamethylphenyl disiloxane	53.3 44.4	362 410	A	[1987STE/MAL] [1974ENG/WOO]
C ₁₁ H ₂₀ O ₃ Si ₃	[na] $\Delta_{\text{v}}H$	1,1,3,3,5-pentamethyl-5-phenylcyclotrisiloxane	48.0	450		[1974DIT/SKO]
C ₁₁ H ₂₄ O ₃ Si	[13080-95-0] $\Delta_{\text{v}}H$ $\Delta_{\text{v}}H$	vinyltripropoxysilane	52.3 ± 0.9 52.3 ± 0.4	298 298	C EB	[1988VOR/BAR] [1985KLY/DAN]
C ₁₁ H ₂₆ Si	[994-63-8] $\Delta_{\text{v}}H$	ethyltripropylsilane	41.0 ± 0.7	298	C	[1988VOR/BAR]
C ₁₁ H ₂₆ Si	[1001-48-5] $\Delta_{\text{v}}H$	methyldipentylsilane	40.3 ± 0.7	298	C	[1988VOR/BAR]
C ₁₁ H ₂₆ Si	[na] $\Delta_{\text{v}}H$	methyl-di(2,2-dimethylpropyl)silane	38.1 ± 0.1	298	C	[1988VOR/BAR]
C ₁₁ H ₂₇ NSi	[133943-80-3] $\Delta_{\text{v}}H$	1,1,1-triethyl-N-(1-methylbutyl)silanamine	46.9 ± 1.0	298	C	[1991VOR/KLY]
C ₁₁ H ₂₈ O ₄ Si ₄	[35331-58-9] $\Delta_{\text{v}}H$ $\Delta_{\text{v}}H$	8,8,10,10,12,12-hexamethyl-7,9,11,13-tetrasiloxa-6,8,10,12-tetrasilaspiro[5,7]tridecane	47.6 48.8	408 449	A	[1987STE/MAL] [1974DIT/SKO]
C ₁₁ H ₂₈ O ₄ Si ₄	[na] $\Delta_{\text{v}}H$	hexamethyl(silacyclohexyl)cyclotetrasiloxane	48.89	453		[1974DIT/SKO2]
C ₁₂ H ₉ Cl ₃ Si	[18030-62-1] $\Delta_{\text{fus}}H$ $\Delta_{\text{fus}}H$ $\Delta_{\text{v}}H$	2-(trichlorosilyl)biphenyl	0.06 20.72	289.5 339.2		[1974GEI/DZH, 1996DOM/HEA]
C ₁₂ H ₉ Cl ₃ Si	[18030-61-0] $\Delta_{\text{fus}}H$ $\Delta_{\text{v}}H$	4-(trichlorosilyl)biphenyl	18.57 75.7	372.9 494	A	[1996DOM/HEA, 1974GEI/DZH] [1987STE/MAL]
C ₁₂ H ₁₀ Cl ₂ Si	[80-10-4] $\Delta_{\text{v}}H$ $\Delta_{\text{v}}H$	dichlorodiphenylsilane	62.5 69.5 ± 4.2	480	A, I	[1987STE/MAL, 1954JEN/CHA] [1966RIN/ONE, 1982PIL/SKI]
C ₁₂ H ₁₀ F ₂ Si	[312-40-3] $\Delta_{\text{v}}H$	difluorodiphenylsilane	50.7	407	A	[1987STE/MAL]
C ₁₂ H ₁₃ NSi	[na] $\Delta_{\text{v}}H$	(N,N-diphenylamino)silane	50.4	460	T	[1969AYL/HAK2]
C ₁₂ H ₂₀ Cl ₈ O ₄ Si	[na] $\Delta_{\text{v}}H$	tris(2,2'-dichloroisopropyl) orthosilicate	U 172.7	524		[1946JON/THO]
C ₁₂ H ₂₀ O ₃ Si	[780-69-8] $\Delta_{\text{v}}H$	triethoxyphenylsilane	58.3 ± 0.9	298	C	[1988VOR/BAR]

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound		T_m (K)	Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)			
	$\Delta_v H$	(344–506)	61.8	359		[1947STU]
C₁₂H₂₀Si	[1112-66-9]	tetraallylsilane				
	$\Delta_{\text{fus}} H$		25.5	244		[2004SMI/LEB]
C₁₂H₂₈O₄Si	[682-01-9]	tetrapropoxysilane				
	$\Delta_v H$		49.8 ± 0.8	298	C	[1988VOR/BAR]
	$\Delta_v H$	(307–563)	66.9	322	A	[1987STE/MAL]
	$\Delta_v H$		49.8 ± 0.4	298	EB	[1985KLY/DAN]
C₁₂H₂₈O₄Si	[1992-48-9]	tetraisopropoxysilane				
	$\Delta_v H$	(327–438)	52.7	342		[1980THO/SMI]
C₁₂H₂₈Si	[994-66-1]	tetrapropylsilane				
	$\Delta_v H$		42.2 ± 0.7	298	C	[1988VOR/BAR]
C₁₂H₂₈Si	[998-41-4]	tributylsilane				
	$\Delta_v H$		42.9 ± 0.7	298	C	[1988VOR/BAR]
C₁₂H₂₈Si	[6485-81-0]	triisobutylsilane				
	$\Delta_v H$		40.0 ± 0.7	298	C	[1988VOR/BAR]
C₁₂H₂₈Si	[17591-43-4]	ethyldipentylsilane				
	$\Delta_v H$		41.2 ± 0.7	298	C	[1988VOR/BAR]
C₁₂H₂₈Si	[18159-61-0]	ethyldiisopentylsilane				
	$\Delta_v H$		42.6 ± 0.7	298	C	[1988VOR/BAR]
C₁₂H₃₀HgSi₂	[4149-29-5]	bis(triethylsilyl)mercury				
	$\Delta_v H$	(383–433)	64.0	398		[1972BRA/KAR]
2(C₆H₁₅N₃)–Cl₄Si	[na]	bis-1,3,5-trimethyl-1,3,5-triazacyclohexane- tetrachlorosilane complex				
	$\Delta_{\text{sub}} H$	(298–354)	76.1 ± 4.6			[1984GOL/LEV]
C₁₂H₃₀O₃Si₃	[2031-79-0]	hexaethyl cyclotrisiloxane				
	$\Delta_{\text{us}} H$		0.46	160		
	$\Delta_{\text{us}} H$		11.82	242.3		
	$\Delta_{\text{fus}} H$		11.42	280.2		[1990DOM/HEA]
	$\Delta_v H$	(385–524)	57.9	400	A	[1987STE/MAL]
	$\Delta_v H$	(434–516)	58.7	449	A	[1987STE/MAL, 1954JEN/CHA]
C₁₂H₃₁N₃Si	[15730-66-2]	N,N,N',N',N'',N'-hexamethylsilanetriamine				
	$\Delta_v H$		58.4 ± 1.0	298	C	[1991VOR/KLY]
C₁₂H₃₆O₄Si₅	[141-63-9]	dodecamethyl pentasiloxane				
	$\Delta_v H$	(395–515)	55.4	410	EB	[1986FLA]
	$\Delta_v H$	(389–498)	50.3	404	A	[1987STE/MAL, 1971SKO/DIT]
	$\Delta_v H$		53.1 ± 2.1			[1972PED/ISE, 1982PIL/SKI]
C₁₂H₃₆O₄Si₅	[3555-47-3]	tetrakis(trimethylsiloxy)silane				
	$\Delta_v H$	(398–494)	52.3	413	EB	[1986FLA]
C₁₂H₃₆O₆Si₆	[540-97-6]	dodecamethyl cyclohexasiloxane				
	$\Delta_{\text{fus}} H$		28.58	269		[1971ALV/DAL]
	$\Delta_v H$	(411–531)	56.1	426	EB	[1986FLA]
	$\Delta_v H$	(340–509)	62.6	355	A	[1987STE/MAL, 1971DIT/SKO]
C₁₂H₃₆Si₅	[4098-98-0]	tetrakis(trimethylsilyl)silane				
	$\Delta_{\text{us}} H$	(203–298)	42	243.7	DSC	[2001SON/HUA]
	$\Delta_{\text{us}} H$		11.57	241.2	DTA	[1971MUR/BRE]

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				Method	Reference	
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)				
		Note: Compound sublimed, fusion values not reported.						
	$\Delta_{\text{sub}}H$		83.7 ± 20.9	298			[1982PIL/SKI, 1972PED/ISE]	
C ₁₂ H ₃₆ Si ₆	[4098-30-0]	dodecamethylcyclohexasilane						
	$\Delta_{\text{trs}}H$		16.7	352.4				
	$\Delta_{\text{fus}}H$		4.2	528.8			[1986CAO/WES]	
C ₁₃ H ₁₄ Si	[776-76-1]	methyldiphenylsilane						
	Δ_vH		64.6 ± 0.8	298	C		[1988VOR/BAR]	
C ₁₃ H ₁₉ NO ₄ Si	[63071-93-2]	1-(2-phenoxy)methyl-2,8,9-trioxa-5-aza-1-silabicyclo[3.3.3]undecane						
	$\Delta_{\text{sub}}H$		108 ± 0.8				[1989VOR/BAR]	
C ₁₃ H ₂₆ O ₂ Si ₃	[546-44-1]	1,1,1,3,5,5,5-heptamethyl-3-phenyl trisiloxane						
	$\Delta_{\text{fus}}H$		18.29	226.8			[1996DOM/HEA]	
	Δ_vH	(357–492)	61.5	372	A		[1987STE/MAL]	
C ₁₃ H ₂₆ O ₄ Si ₄	[10448-09-6]	2,4,4,6,6,8,8-heptamethyl-2-phenylcyclotetrasiloxane						
	Δ_vH	(397–514)	65.6	412	A		[1987STE/MAL]	
C ₁₃ H ₃₀ O ₃ SSi	[57557-75-2]	[3-(butylthio)propyl]triethoxysilane						
	Δ_vH		47.1 ± 0.6	298	C		[1989VOR/SOR]	
C ₁₃ H ₃₀ Si	[18414-75-0]	decyltrimethylsilane						
	Δ_vH	(340–513)	57.8	355			[1947STU]	
C ₁₃ H ₃₀ Si	[1001-46-3]	methyldihexylsilane						
	Δ_vH		42.6 ± 0.7	298	C		[1988VOR/BAR]	
C ₁₄ H ₆ F ₁₀ Si	[10536-62-6]	di(pentafluorophenyl)dimethylsilane						
	Δ_vH	(366–463)	55.3 ± 0.6	414			[2006ZEL/CHU]	
C ₁₄ H ₁₆ O ₂ Si	[6843-66-9]	diphenoxydimethylsilane						
	Δ_vH		64.4 ± 0.9	298	C		[1988VOR/BAR]	
C ₁₄ H ₁₆ Si	[7535-07-1]	ethylidiphenylsilane						
	Δ_vH		66.1 ± 0.8	298	C		[1988VOR/BAR]	
C ₁₄ H ₁₉ NO ₅ Si	[79791-55-2]	2,8,9-trioxa-5-aza-1-silabicyclo[3.3.3]undecane-1-methanol benzoate ester						
	$\Delta_{\text{sub}}H$		109 ± 0.9				[1989VOR/BAR]	
C ₁₄ H ₂₁ NO ₃ Si	[63330-92-7]	1-(2-phenylethyl)-2,8,9-trioxa-5-aza-1-silabicyclo[3.3.3]undecane						
	$\Delta_{\text{sub}}H$		108 ± 0.9				[1989VOR/BAR]	
C ₁₄ H ₃₂ Si	[10175-53-8]	triethyloctylsilane						
	Δ_vH	(347–535)	56.1	361			[1947STU]	
C ₁₄ H ₃₂ Si	[17591-45-6]	ethylidihexylsilane						
	Δ_vH		44.8 ± 0.7	298	C		[1988VOR/BAR]	
C ₁₄ H ₃₂ Si	[994-76-3]	dipropyldibutylsilane						
	Δ_vH		44.0 ± 0.8	298	C		[1988VOR/BAR]	
C ₁₄ H ₃₃ NSi	[17995-32-3]	N,N-dibutyl-1,1,1-triethylsilanamine						
	Δ_vH		56.3 ± 1.0	298	C		[1991VOR/KLY]	
C ₁₄ H ₃₃ NSi	[133943-79-0]	1,1,1-triethyl-N,N-bis(1-methylpropyl)silanamine						
	Δ_vH		51.4 ± 0.9	298	C		[1991VOR/KLY]	
C ₁₄ H ₃₃ NSi	[133943-81-4]	1,1,1-triethyl-N-octylsilanamine						
	Δ_vH		59.1 ± 1.0	298	C		[1991VOR/KLY]	

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)			
C ₁₄ H ₄₂ O ₂ Si ₆	[1787-37-7] $\Delta_v H$	<i>bis</i> [(pentamethyl)disilanoxy]disilane (442–474)	62.2	457		[1962CRA/URE]	
C ₁₄ H ₄₂ O ₅ Si ₆	[107-52-8] $\Delta_v H$	tetradecamethyl cycloheptasiloxane (449–545)	56.9	464	EB	[1986FLA]	
	$\Delta_v H$	(397–522)	56.6	412	A	[1987STE/MAL, 1971SKO/DIT]	
C ₁₄ H ₄₂ O ₇ Si ₇	[107-50-6] $\Delta_{\text{fus}} H$	tetradecamethyl cycloheptasiloxane	20.88	237.7		[1971ALV/DAL]	
	$\Delta_v H$	(359–537)	58.6	374	A	[1987STE/MAL]	
	$\Delta_v H$	(431–548)	60.6	446	EB	[1986FLA]	
C ₁₅ H ₁₈ OSi	[1825-59-8] $\Delta_v H$	methyldiphenylethoxysilane (373–512)	72.9			[2008WU/JIA]	
C ₁₅ H ₂₁ NO ₅ Si	[100446-65-9] $\Delta_{\text{sub}} H$	4-methylbenzoic acid 2,8,9-trioxa-5-aza-1-silabicyclo[3.3.3]undec-1-ylmethyl ester	123 ± 0.9			[1989VOR/BAR]	
C ₁₅ H ₂₁ NO ₆ Si	[94697-86-6] $\Delta_{\text{sub}} H$	4-methoxybenzoic acid 2,8,9-trioxa-5-aza-1-silabicyclo[3.3.3]undec-1-ylmethyl ester	143 ± 0.9			[1989VOR/BAR]	
C ₁₅ H ₃₄ Si	[994-78-5] $\Delta_v H$	propyltributylsilane	45.0 ± 0.8	298	C	[1988VOR/BAR]	
C ₁₅ H ₃₄ Si	[6485-78-5] $\Delta_v H$	tripentylsilane	48.1 ± 0.8	298	C	[1988VOR/BAR]	
C ₁₅ H ₃₄ Si	[17922-08-6] $\Delta_v H$	triisopentylsilane	43.8 ± 0.7	298	C	[1988VOR/BAR]	
C ₁₅ H ₃₄ Si	[17908-09-7] $\Delta_v H$	dodecyltrimethylsilane (364–546)	62.0	379		[1947STU]	
C ₁₆ H ₁₂ Si	[1675-57-6] $\Delta_{\text{fus}} H$	diphenyldiethynylsilane	19.67	316.2		[1996DOM/HEA]	
C ₁₆ H ₂₀ O ₂ Si	[17964-48-6] $\Delta_v H$	<i>bis</i> (2-methylphenoxy)dimethylsilane	63.6 ± 0.8	298	C	[1988VOR/BAR]	
C ₁₆ H ₂₀ O ₂ Si	[17964-47-5] $\Delta_v H$	<i>bis</i> (3-methylphenoxy)dimethylsilane	61.5 ± 0.8	298	C	[1988VOR/BAR]	
C ₁₆ H ₂₀ O ₂ Si	[17964-49-7] $\Delta_v H$	<i>bis</i> (4-methylphenoxy)dimethylsilane	65.3 ± 0.9	298	C	[1988VOR/BAR]	
C ₁₆ H ₂₀ O ₂ Si	[2553-19-7] $\Delta_v H$	diethoxydiphenylsilane (385–569)	71.5	399		[1947STU]	
C ₁₆ H ₂₂ O ₃ Si ₃	[1693-51-2] $\Delta_{\text{fus}} H$	1,1,3,3-tetramethyl-5,5-diphenylcyclotrisiloxane	22.19	338		[1996DOM/HEA]	
	$\Delta_v H$	(439–523)	66.9	481		[1974DIT/SKO]	
C ₁₆ H ₂₂ O ₃ Si ₃	[31751-60-7] $\Delta_v H$	<i>cis</i> 1,1,3,5-tetramethyl-3,5-diphenylcyclotrisiloxane (423–541)	66	532		[1972DIT/SKO2]	
C ₁₆ H ₂₂ O ₃ Si ₃	[31751-59-4] $\Delta_v H$	<i>trans</i> 1,1,3,5-tetramethyl-3,5-diphenylcyclotrisiloxane (397–535)	66.4	466		[1972DIT/SKO2]	
C ₁₆ H ₃₂ O ₄ Si ₄	[177-49-1] $\Delta_v H$	6,12,18,24-tetraoxa-5,7,13,19-tetrasilatetraspiro[4,1,4,1,4,1,4,1]tetracosane (452–583)	67.6	467	A	[1987STE/MAL]	
	$\Delta_v H$	(452–583)	69.5	518		[1974DIT/SKO2]	

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_{\text{m}}$ (kJ mol ⁻¹)	T_{m} (K)	Method	Reference
$\text{C}_{16}\text{H}_{32}\text{O}_4\text{Si}_4$	[na] $\Delta_{\text{v}}H$	tetra(silacyclopentyl)cyclotetrasiloxane (452–583)	69.5	517		[1974DIT/SKO2]
$\text{C}_{16}\text{H}_{36}\text{O}_3\text{Si}_2$	[349140-64-3] $\Delta_{\text{fus}}H$	11-(1,1,3,3-tetramethyldisiloxanyl)undecanoic acid, methyl ester	23.5	233.3		[2004RYA/LEB]
$\text{C}_{16}\text{H}_{36}\text{O}_4\text{Si}$	[4766-57-8] $\Delta_{\text{v}}H$ $\Delta_{\text{v}}H$	tetrabutoxysilane (333–479)	52.0 ± 1.0 79.6	298 348	C A	[1988VOR/BAR] [1987STE/MAL]
$\text{C}_{16}\text{H}_{40}\text{O}_4\text{Si}_4$	[1451-99-6] $\Delta_{\text{trs}}H$ $\Delta_{\text{fus}}H$ $\Delta_{\text{v}}H$	octaethyl cyclotetrasiloxane (420–574)	12.22 13.71 69.2	208.2 213.4 435		[1990DOM/HEA] [1987STE/MAL]
$\text{C}_{16}\text{H}_{46}\text{O}_7\text{Si}_6$	[18143-15-2] $\Delta_{\text{v}}H$	1,11-diethoxy-1,1,3,3,5,7,7,9,9,11,11-dodecamethylhexasiloxane (376–547)	66.9	391	A	[1987STE/MAL]
$\text{C}_{16}\text{H}_{48}\text{O}_6\text{Si}_7$	[541-01-5] $\Delta_{\text{v}}H$ $\Delta_{\text{v}}H$	hexadecamethylheptasiloxane (443–468) (443–551)	63.8 60.8	459 458	EB A	[1986FLA] [1987STE/MAL, 1971SKO/DIT]
$\text{C}_{16}\text{H}_{48}\text{O}_8\text{Si}_8$	[556-68-3] $\Delta_{\text{v}}H$ $\Delta_{\text{v}}H$	hexadecamethyl cyclooctasiloxane (378–563) (454–576)	66.6 64.5	391 469	A EB	[1987STE/MAL] [1986FLA]
$\text{C}_{17}\text{H}_{26}\text{O}_4\text{Si}_4$	[13093-12-4] $\Delta_{\text{v}}H$	hexamethyl(silaacenaphthenyl)cyclotetrasiloxane (466–548)	68.6	507		[1974DIT/SKO2]
$\text{C}_{17}\text{H}_{32}\text{O}_2\text{Si}$	[na] $\Delta_{\text{v}}H$	3-methyl-3-[2-cyclohexylpropylperoxy]-1-trimethylsilyl-1-butyne (307–318)	74.2 ± 2.0		ME	[1999DIB/LUS]
$\text{C}_{17}\text{H}_{38}\text{Si}$	[18558-18-4] $\Delta_{\text{v}}H$	tetradecyltrimethylsilane (393–573)	70.9	408		[1947STU]
$\text{C}_{18}\text{H}_{15}\text{ClSi}$	[76-86-8] $\Delta_{\text{fus}}H$	triphenylchlorosilane	26.88	370.6		[1996DOM/HEA]
$\text{C}_{18}\text{H}_{28}\text{O}_2\text{Si}_3$	[797-77-3] $\Delta_{\text{fus}}H$	1,1,1,5,5,5-hexamethyl-3,3-diphenyltrisiloxane	22.75	270.5		[1996DOM/HEA]
$\text{C}_{18}\text{H}_{28}\text{O}_4\text{Si}_4$	[18604-02-9] $\Delta_{\text{v}}H$	2,2,4,4,6,8-hexamethyl-6,8-diphenylcyclotetrasiloxane (459–576)	70.5	474	A	[1987STE/MAL]
$\text{C}_{18}\text{H}_{28}\text{O}_4\text{Si}_4$	[1693-44-3] $\Delta_{\text{fus}}H$	1,1,3,3,5,5-hexamethyl-7,7-diphenylcyclotetrasiloxane	42.73	305		[1996DOM/HEA]
$\text{C}_{18}\text{H}_{40}\text{Si}$	[2929-52-4] $\Delta_{\text{v}}H$	trihexylsilane	51.0 ± 0.7	298	C	[1988VOR/BAR]
$\text{C}_{18}\text{H}_{40}\text{Si}$	[51502-64-8] $\Delta_{\text{v}}H$	ethyldioctylsilane	47.3 ± 0.7	298	C	[1988VOR/BAR]
$\text{C}_{18}\text{H}_{48}\text{Si}_6$	[76750-22-6] $\Delta_{\text{trs}}H$ $\Delta_{\text{fus}}H$	1,2,3,4,5,6-hexamethyl-1,2,3,4,5,6-hexaethylcyclohexasilane	3.8 1.8	226.3 439.2		[1986CAO/WES]
$\text{C}_{18}\text{H}_{54}\text{O}_7\text{Si}_8$	[556-69-4] $\Delta_{\text{v}}H$ $\Delta_{\text{v}}H$	octadecamethyl octasiloxane (378–563) (464–586)	67.7 68.4	393 479	A EB	[1987STE/MAL] [1986FLA]

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound			Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)		
C ₁₈ H ₅₄ O ₉ Si ₉	[556-71-8]	octadecamethyl cyclononasiloxane				[1971ALV/DAL]
	$\Delta_{\text{fus}}H$		25.64	246.2		
	Δ_vH	(463–584)	67.9	478	A	[1987STE/MAL]
		(473–578)	68	488	EB	[1986FLA]
C ₁₉ H ₁₈ O ₃ Si	[3439-97-2]	methyltriphenoxysilane				
	Δ_vH		71.5 ± 0.9	298	C	[1988VOR/BAR]
C ₂₀ H ₂₀ OSi	[1516-80-9]	ethoxytriphenylsilane				
	$\Delta_{\text{sub}}H$		142.7 ± 1.0			[1988VOR/BAR]
		(440–617)	89.7	455		[1947STU]
C ₂₀ H ₂₁ ClOSi ₂	[53634-34-7]	1,3-dimethyl-1,1,3-triphenyl-3-chlorodisiloxane				
	Δ_vH	(468–626)	69.0	547		[1974DIT/SKO]
C ₂₀ H ₃₀ O ₃ Si ₃	[108543-32-4]	1,1,3,3-tetraethyl-5,5-diphenylcyclotrisiloxane				
	$\Delta_{\text{fus}}H$		18.37	279.1		[1996DOM/HEA]
C ₂₀ H ₄₄ O ₄ Si	[na]	<i>tetrakis</i> (1-ethylprooxy)silane				
	Δ_vH	(371–427)	89.2	386	A	[1987STE/MAL]
C ₂₀ H ₅₀ Si ₅	[75217-22-0]	decaethylcyclopentasilane				
	$\Delta_{\text{fus}}H$		16.3	254.8		
	$\Delta_{\text{fus}}H$		1.4	440.1		[1986CAO/WES]
C ₂₀ H ₅₈ O ₉ Si ₈	[18724-16-6]	1,15-diethoxy-1,1,3,3,5,5,7,7,9,9,1,1,13,13,15,15-hexadecamethyloctasiloxane				
	Δ_vH	(406–585)	79.7	421	A	[1987STE/MAL]
C ₂₀ H ₆₀ O ₈ Si ₉	[2652-13-3]	eicosamethylnonasiloxane				
	Δ_vH	(417–581)	85.9	432	A	[1987STE/MAL]
C ₂₀ H ₆₀ O ₁₀ Si ₁₀	[18772-36-6]	eicosamethylcyclodecasiloxane				
	$\Delta_{\text{fus}}H$		39.76	265.8		[1971ALV/DAL]
	Δ_vH	(480–603)	71.3	495	A	[1987STE/MAL]
C ₂₁ H ₂₂ Si	[1747-92-8]	tribenzylsilane				
	Δ_vH	(460–637)	81.9	475	A	[1987STE/MAL]
C ₂₁ H ₂₄ OSi ₂	[14920-93-5]	1,1,3-trimethyl-1,3,3-triphenyl disiloxane				
	Δ_vH	(494–624)	80	509	A	[1987STE/MAL]
	Δ_vH	(495–624)	64.4	560		[1974DIT/SKO]
C ₂₁ H ₂₄ O ₃ Si ₃	[6138-53-0]	<i>trans</i> 1,3,5-trimethyl-1,3,5-triphenylcyclotrisiloxane				
	$\Delta_{\text{fus}}H$		43.66	320.9		[1996DOM/HEA, 1975MEK/KAR2]
	Δ_vH	(483–586)	76.1	534		[1972DIT/SKO]
C ₂₁ H ₂₄ O ₃ Si ₃	[3424-57-5]	<i>cis</i> 1,3,5-trimethyl-1,3,5-triphenylcyclotrisiloxane				
	$\Delta_{\text{fus}}H$		43.07	374.3		[1996DOM/HEA, 1975MEK/KAR2]
	Δ_vH	(473–551)	80.6	512		[1972DIT/SKO]
C ₂₁ H ₄₆ Si	[18753-02-1]	triheptylsilane				
	Δ_vH		57.4 ± 0.8	298	C	[1988VOR/BAR]
C ₂₁ H ₄₆ Si	[51502-65-9]	methyldidecylsilane				
	Δ_vH		57.4 ± 0.8	298	C	[1988VOR/BAR]
C ₂₂ H ₂₄ O ₃ Si	[55893-94-2]	methyltris(2-methylphenoxy)silane				
	Δ_vH		68.2 ± 0.9	298	C	[1988VOR/BAR]

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C ₂₂ H ₂₄ O ₃ Si	[55893-95-3] $\Delta_v H$	methyltris(3-methylphenoxy)silane	66.9 ± 0.8	298	C	[1988VOR/BAR]
C ₂₂ H ₂₄ O ₃ Si	[55893-96-4] $\Delta_v H$	methyltris(4-methylphenoxy)silane	70.3 ± 0.9	298	C	[1988VOR/BAR]
C ₂₂ H ₄₀ O ₄ Si	[na] $\Delta_v H$	dimethyldi-[3-methyl-3-tert-amylperoxy-1-butynyl]silane	(318–338) 92.0 ± 1.6		ME	[1999DIB/LUS]
C ₂₂ H ₄₈ Si	[51502-66-0] $\Delta_v H$	ethylididecylsilane	58.7 ± 0.8	298	C	[1988VOR/BAR]
C ₂₂ H ₆₆ O ₁₁ Si ₁₁	[18766-38-6] $\Delta_{\text{fus}} H$ $\Delta_v H$	docosamethyl cycloundecasiloxane	17.73 74.5	216.2 511		[1971ALV/DAL] A [1987STE/MAL]
C ₂₃ H ₃₀ O ₃ Si ₃	[67102-99-2] $\Delta_v H$	1,1,1,3,5-pentamethyl-3,5,5-triphenyltrisiloxane	(521–678) 69.8	536	A	[1987STE/MAL]
C ₂₄ F ₂₀ Si	[1524-78-3] $\Delta_{\text{fus}} H$ $\Delta_{\text{sub}} H$ $\Delta_v H$	tetra(pentafluorophenyl)silane	46.9 (433–517) 128 ± 1.2 80.6 ± 0.4	518 475 514		[2006ZEL/CHU] [2006ZEL/CHU] [2006ZEL/CHU]
C ₂₄ H ₂₀ O ₄ Si	[1174-72-7] $\Delta_{\text{sub}} H$	tetraphenoxysilane	124.7 ± 1.2			[1988VOR/BAR]
C ₂₄ H ₂₀ Si	[1048-08-4] $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$	tetraphenylsilane	(428–484) 51.2 156.9 ± 1.7 149.4 ± 1.7 (428–489) 51.0 51.0	456 298 298	A ME,TE MG	[1987STE/MAL] [1978STE4] [1974CAL/KAN] [1973MCC/SMI] [1972NEW, 1986MAR/LOE]
C ₂₄ H ₂₂ N ₂ Si	[22519-45-5] $\Delta_v H$ $\Delta_v H$	N,N,N'N'-tetraphenyl silane diamine	(410–473) 59.1 (410–473) 57.3	425 441	A T	[1987STE/MAL] [1969AYL/HAK2]
C ₂₄ H ₅₂ O ₄ Si	[7425-86-7] $\Delta_v H$	tetrahexyloxysilane	(454–573) 87.0	469	A	[1987STE/MAL]
C ₂₄ H ₅₂ Si	[18765-09-8] $\Delta_v H$	trioctylsilane	59.8 ± 0.8	298	C	[1988VOR/BAR]
C ₂₄ H ₇₂ O ₁₂ Si ₁₂	[18919-94-3] $\Delta_{\text{fus}} H$ $\Delta_v H$	tetracosamethyl cyclododecasiloxane	15.45 76.6	234.2 523		[1971ALV/DAL] A [1987STE/MAL]
C ₂₅ H ₄₀ O ₂ Si ₂	[na] $\Delta_{\text{fus}} H$	norethindrone pentamethyldisiloxy ether	22.9	355		[1996DOM/HEA]
C ₂₆ H ₂₆ OSi ₂	[807-28-3] $\Delta_{\text{fus}} H$ $\Delta_v H$ $\Delta_v H$	1,3-dimethyl-1,1,3,3-tetraphenyldisiloxane	26.58 (518–616) 93.3 (518–685) 64.4	322 533 602		[1996DOM/HEA] A [1987STE/MAL] [1974DIT/SKO]
C ₂₆ H ₂₆ O ₃ Si ₃	[1438-86-4] $\Delta_{\text{fus}} H$	1,1-dimethyl-3,3,5,5-tetraphenylcyclotrisiloxane	28.2	361.1		[1996DOM/HEA]

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound		T _m (K)	Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)			
C ₂₇ H ₃₀ O ₂ Si	[na] $\Delta_v H$	3-methyl-3- <i>tert</i> -butylperoxy-1-triphenylsilyl-1-butyne (378–398)	115.9 ± 3.2		ME	[1999DIB/LUS]
C ₂₇ H ₅₈ Si	[51502-67-1] $\Delta_v H$	trinonylsilane	61.8 ± 0.8	298	C	[1988VOR/BAR]
C ₂₈ H ₂₈ O ₄ Si	[16714-40-2] $\Delta_v H$	tetrakis(2-methylphenoxy)silane	76.2 ± 1.0	298	C	[1988VOR/BAR]
C ₂₈ H ₂₈ O ₄ Si	[16714-54-8] $\Delta_v H$	tetrakis(3-methylphenoxy)silane	73.6 ± 0.9	298	C	[1988VOR/BAR]
C ₂₈ H ₂₈ O ₄ Si	[16714-41-3] $\Delta_v H$	tetrakis(4-methylphenoxy)silane	97.1 ± 1.0	298	C	[1988VOR/BAR]
C ₂₈ H ₃₂ O ₂ Si	[na] $\Delta_v H$	3-methyl-3- <i>tert</i> -amylperoxy-1-triphenylsilyl-1-butyne (378–393)	120.3 ± 5.8		ME	[1999DIB/LUS]
C ₂₈ H ₃₂ O ₂ Si ₃	[67103-00-8] $\Delta_v H$	1,1,1,3-tetramethyl-3,5,5,5-tetraphenyltrisiloxane (549–678)	82.6	564	A	[1987STE/MAL]
C ₂₈ H ₃₂ O ₂ Si ₃	[67142-05-6] $\Delta_v H$	1,1,3,5-tetramethyl-1,3,5,5-tetraphenyltrisiloxane (566–666)	90.9	581	A	[1987STE/MAL]
C ₂₈ H ₃₂ O ₂ Si ₃	[3982-82-9] $\Delta_v H$	1,3,3,5-tetramethyl-1,1,5,5-tetraphenyltrisiloxane (544–686)	88.3	559	A	[1987STE/MAL]
C ₂₈ H ₃₂ O ₄ Si ₄	[1693-47-6] $\Delta_{\text{us}}H$ $\Delta_{\text{us}}H$ $\Delta_{\text{fus}}H$	1,1,3,3-tetramethyl-5,5,7,7-tetraphenylcyclotetrasiloxane	0.24 1.05 27.05	186.5 271.5 346.2	AC	[1996DOM/HEA, 1976KUL/DZH]
C ₂₈ H ₃₂ O ₄ Si ₄	[77-63-4] $\Delta_{\text{fus}}H$	1,1,5',7'-tetramethyl-1',3',5,7-tetraphenylcyclotetrasiloxane	24.62	373.4	AC	[1996DOM/HEA]
C ₂₉ H ₃₄ O ₂ Si	[261766-67-0] $\Delta_v H$	3-methyl-3- <i>tert</i> -hexylperoxy-1-triphenylsilyl-1-butyne (383–398)	126.3 ± 3.0		ME	[1999DIB/LUS]
C ₂₉ H ₄₄ O ₆ Si ₂	[179108-75-9] $\Delta_{\text{us}}H$ $\Delta_{\text{fus}}H$	4-[[1-oxo-11-(1,1,3,3-tetramethyldisiloxanyl)undecyl]oxy]benzoic acid, 4-methoxyphenyl ester	20.1 5.4	322 331		[2004RYA/LEB]
C ₂₉ H ₄₄ O ₆ Si ₂	[349149-95-7] $\Delta_{\text{us}}H$ $\Delta_{\text{fus}}H$	4-methoxybenzoic acid, 4-[[1-oxo-11-(1,1,3,3-tetramethyldisiloxanyl)undecyl]oxy]phenyl ester	24.3 4.7	324 334		[2004RYA/LEB]
C ₃₀ H ₄₀ F ₃₀ O ₃ Si ₄	[na] $\Delta_v H$	1,1,1,3,5,5,5-hepta(γ -trifluoropropyl)-3- <i>tris</i> (γ -trifluoropropyl)siloxyltrisiloxane (502–652)	64.3	671		[1974DIT/SKO3]
C ₃₀ H ₆₄ Si	[18765-73-6] $\Delta_v H$	<i>tris</i> (decyl)silane	65.3 ± 0.8	298	C	[1988VOR/BAR]
C ₃₂ H ₁₆ Cl ₂ N ₈ Si	[19333-10-9] $\Delta_{\text{sub}}H$	silicon phthalocyanine dichloride	151.3			[1972MAR/LOP]
C ₃₂ H ₇₀ Si ₁₀	[206868-23-7] $\Delta_{\text{fus}}H$	1,1,2,2,3,3,4,4,5,5,6,6,7,7,8,8,9,9,10,10-eicosamethyl-1,10-diphenyldecasilane	56.1	391.2		[2001YAT/MIN]
C ₃₃ H ₃₄ O ₂ Si ₃	[67103-01-9] $\Delta_v H$	1,1,3-trimethyl-1,3,5,5,5-pentaphenyltrisiloxane (603–711)	91.3	618	A	[1987STE/MAL]
C ₃₃ H ₃₄ O ₂ Si ₃	[3390-61-2]	1,3,5-trimethyl-1,1,3,5,5-pentaphenyltrisiloxane				

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(575–625)	87.3	590	A	[1987STE/MAL]
C₃₃H₃₄O₄Si₄	[32395-60-1]	1,3,5-trimethyl-1,3,5,7,7-pentaphenylcyclotetrasiloxane				
	$\Delta_v H$	(523–676)	86.9	600		[1974DIT/SKO]
C₃₄H₇₆Si₁₁	[386719-88-6]	1,1,2,2,3,3,4,4,5,5,6,6,7,7,8,8,9,9,10,10,11-docosamethyl-1,11-diphenylundecasilane				
	$\Delta_{\text{fus}} H$		57.8	398.2		[2001YAT/MIN]
C₃₆H₃₀Si₂	[1450-23-3]	hexaphenyldisilane				
	$\Delta_{\text{sub}} H$		209.2 ± 2.1	298	ME,TE	[1974CAL/KAN]
C₃₆H₃₀Si₃O₃	[512-63-0]	hexaphenylcyclotrisiloxane				
	$\Delta_{\text{us}} H$		1.13	455		
	$\Delta_{\text{fus}} H$		39.3	466		[2000LEB/SMI]
C₄₈H₄₀Si₄O₄	[546-56-5]	octaphenylcyclotetrasiloxane				
	$\Delta_{\text{us}} H$		44.8	459.2		
	$\Delta_{\text{fus}} H$		1.26	473.2	DSC	[2001MAT/SHC]
	$\Delta_{\text{us}} H$		2.22	348		
	$\Delta_{\text{us}} H$		46.4	463		
	$\Delta_{\text{fus}} H$		1.13	475		[2000LEB/SMI]
	$\Delta_{\text{us}} H$		2.9	349.8		
	$\Delta_{\text{us}} H$		43.8	462.8		
	$\Delta_{\text{fus}} H$		1.95	478.1		[1979SMI]
Br₃HSi	[7789-57-3]	tribromosilane				
	$\Delta_v H$	(273–393)	34.8	333		[1934SCH/BIC]
Cl₂H₂Si	[4109-96-0]	dichlorosilane				
	$\Delta_v H$	(290–350)	22.2 ± 0.7			[1986VOR/BAL]
Cl₃HSi	[10025-78-2]	trichlorosilane				
	$\Delta_v H$	(303–325)	24.9	314		[1967LAP/NIS]
	$\Delta_v H$	(275–305)	27.2	290	I	[1954JEN/CHA]
Cl₄Si	[10026-04-7]	silicon tetrachloride				
	$\Delta_{\text{fus}} H$		7.72	204.7		[1985DEV/GUE]
	$\Delta_{\text{sub}} H$	(175–204)	43.3 ± 0.1		MG	[1964BAL/DON]
	$\Delta_v H$	(298–313)	29.9	305		[1973JAI/YAD]
	$\Delta_v H$	(273–326)	30.4	288		[1964CAP/FRI]
	$\Delta_v H$	(275–330)	30.2	290	I	[1954JEN/CHA]
	$\Delta_v H$	(273–333)	30.1	303		[1936KEA]
FH₃Si	[13537-33-2]	fluorosilane				
	$\Delta_v H$	(145–167)	18.5	156		[1944EME/MAD]
FH₅Si₂	[14537-73-6]	disilanyl fluoride				
	$\Delta_v H$	(178–227)	26.3	202	T	[1963ABE/MAC]
F₂H₂Si	[13824-36-7]	difluorosilane				
	$\Delta_v H$	(151–167)	19.9	159		[1944EME/MAD]
F₂H₄NPSi	[36875-96-4]	silylaminodifluorophosphine				
	$\Delta_v H$	(200–273)	34.3	236		[1972ARN/EBS]
F₃ISi	[16865-60-4]	trifluoroiodosilane				
	$\Delta_v H$	(139–227)	21.3	183		[1973AYL/ELL2]
F₃HSi	[13465-71-9]	trifluorosilane				
	$\Delta_v H$	(156–168)	20.1	162		[1944EME/MAD]

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound		T_m (K)	Method	Reference	
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)				
F₃H₃Si₂	[15195-26-3] $\Delta_{\text{sub}}H$	1,1,1-trifluorodisilane (195–209)	39.2	202		[1972SOL/BUR]	
F₄Si	[7783-61-1] $\Delta_{\text{sub}}H$	silicon tetrafluoride (148–183)	25.8			[1930PAT/PAP]	
H₅ClSi₂	[14565-98-1] Δ_vH	disilanyl chloride (227–273)	29.3			[1962CRA/URE]	
H₅ISi₂	[14380-76-8] Δ_vH	disilanyl iodide (274–363)	33.9	318		[1960WAR/MAC]	
H₇NSi₂	[5702-11-4] Δ_vH	disilazane (177–250)	23.4	213	SG	[1969AYL/HAK3]	
H₉PSi₃	[15110-33-5] Δ_vH	trisilylphosphine (243–284)	36.4	263		[1962AMB/BOE2]	
H₉SbSi₃	[14798-31-3] Δ_vH	trisilylstibine	32.0			[1963AMB/BOE]	
H₁₀Si₄	[7783-29-1] Δ_vH	tetrasilane (273–369)	35.6		T	[1946EME/MAD]	
H₁₀OSi₄	[14809-36-0] Δ_vH	bis(disilanyl) ether (273–363)	36.4	318		[1960WAR/MAC]	
Sm (samarium)							
C₁₅H₁₅Sm	[1298-55-1] $\Delta_{\text{sub}}H$	tris(cyclopentadienyl)samarium(III) (513–633)	109.6 ± 1.7			[1973BOR/KRA]	
C₁₅H₂₁O₆Sm	[14589-42-5] $\Delta_{\text{sub}}H$	tris(2,4-pentanedionato)samarium(III) (293–413)	U 20 ± 2			[1985SER/ZAG]	
C₃₀H₃₀F₂₁O₆Sm	[17631-69-5] $\Delta_{\text{sub}}H$	tris(1,1,1,2,2,3,3-heptafluoro-7,7-dimethyloctane-4,6-dione)samarium(III) (379–394)	158.6 ± 1.7		ME	[1971SWA/KAR]	
C₃₃H₅₇O₆Sm	[15492-50-9] $\Delta_{\text{sub}}H$	tris(2,2,6,6-tetramethylheptan-3,5-dionato)samarium(III)	149.7 ± 3.3	298	DSC	[1999SAN/PET]	
	$\Delta_{\text{sub}}H$		(378–418)	180.7	398	ME	[1981AMA/SAT]
	$\Delta_{\text{sub}}H$		(430–468)	150.6	447	BG	[1969SIC/DUB]
	Δ_vH		(468–500)	93.0		BG	[1969SIC/DUB]
SmI₂	[32248-43-4] Δ_vH	samarium diiodide (1008–1155)	255.9 ± 5.3	1082		[1974HIR/CAS]	
Sn (tin)							
C₃H₉BrSn	[1066-44-0] Δ_vH	trimethyltin bromide	47.3 ± 4.2			[1957PED/SKI, 1982PIL/SKI]	
C₃H₉ISn	[811-73-4] Δ_vH	trimethyltin iodide	48.1 ± 4.2			[1957PED/SKI, 1982PIL/SKI]	
C₄H₉F₃Sn	[754-25-6] Δ_vH	(trifluoromethyl)trimethyltin (276–323)	37.5	300	T	[1960KAE/PHI]	
C₄H₁₂S₄Sn	[210298-57-0] $\Delta_{\text{fus}}H$	tetra(methylthia)tin	24.1	307.5	DSC	[1998FUE/STR]	
C₄H₁₂Sn	[594-27-4]	tetramethyltin					

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound		T_m (K)	Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)			
	$\Delta_{\text{fus}}H$		9.23	218.2	DSC	[1989SHE/RAB, 1974UTS/BAC]
	Δ_vH	(273–350)	32.6 ± 0.2	311		[2001BAE]
	Δ_vH	(313–393)	31.3		GC	[1992HAW]
	Δ_vH		31.1 ± 0.1	298	C	[1980ABR/IRV]
	Δ_vH		32.8 ± 0.1	298		[1970VAL]
	Δ_vH		33.1 ± 1.3			[1963DAV/POP, 1982PIL/SKI]
	Δ_vH		30.5	298		[1936THO/LIN]
	Δ_vH	(273–308)	33.4	290	I	[1930BUL/HAU]
	Δ_vH	(308–355)	31.6	331	I	[1930BUL/HAU]
	Δ_vH		33.1	298	I	[1930BUL/HAU]
	Δ_vH	(298–308)	31.6	303		[1929TAN/NAG]
C₅H₉F₅Sn	[812-35-1]	(pentafluoroethyl)trimethyltin				
	Δ_vH	(295–329)	35.6	312	T	[1960KAE/PHI]
C₅H₁₂Sn	[754-06-3]	trimethylvinyltin				
	Δ_vH		37.2 ± 2.1			[1959PED/SKI, 1982PIL/SKI]
C₅H₁₄Sn	[3531-44-0]	ethyl trimethyltin				
	Δ_vH		37.7 ± 1.7			[1963DAV/POP, 1982PIL/SKI]
	Δ_vH	(243–381)	38.4	258		[1947STU]
	Δ_vH	(273–336)	37.0	304	I	[1930BUL/HAU]
	Δ_vH	(336–384)	34.9	360	I	[1930BUL/HAU]
C₆H₁₆Sn	[3531-45-1]	trimethylpropyltin				
	Δ_vH	(261–405)	43.8	276		[1947STU]
	Δ_vH	(286–328)	41.4	307	I	[1930BUL/HAU]
	Δ_vH	(328–405)	38.0	366	I	[1930BUL/HAU]
C₆H₁₆Sn	[3531-46-2]	trimethylisopropyltin				
	Δ_vH		40.6 ± 2.1			[1966COL/SKI, 1982PIL/SKI]
C₆H₁₈Sn₂	[661-69-8]	hexamethyldistannane				
	Δ_vH		50.2 ± 4.2			[1957PED/SKI, 1982PIL/SKI]
C₇H₁₈OSn	[1067-21-6]	triethylmethoxystannane				
	Δ_vH	(312–435)	49.9	273	MM	[2001BAE2]
	Δ_vH	(312–435)	48.7	298	MM	[2001BAE2]
C₇H₁₈Sn	[3531-47-3]	<i>tert</i> -butyltrimethyltin				
	Δ_vH		54.0 ± 4.2			[1966COL/SKI, 1982PIL/SKI]
C₈H₁₂Sn	[1112-55-6]	tetravinyltin				
	Δ_vH	(313–393)	40.5		GC	[1992HAW]
C₈H₁₅F₅Sn	[2925-46-4]	(pentafluoroethyl)triethyltin				
	Δ_vH	(303–343)	39.2	323	T	[1960KAE/PHI]
C₈H₁₈Cl₂Sn	[683-18-1]	di- <i>n</i> -butylindichloride				
	$\Delta_{\text{fus}}H$		22.75	316.2		[1974UTS/BAC2]
C₈H₂₀Sn	[597-64-8]	tetraethyltin				
	$\Delta_{\text{fus}}H$		9.15	142.1		[1996DOM/HEA]
	Δ_vH	(293–455)	46.6 ± 0.6	374		[2001BAE]
	Δ_vH	(313–393)	42.4		GC	[1992HAW]
	Δ_vH		50.6 ± 0.2	298	C	[1980ABR/IRV]
	Δ_vH		51.0 ± 2.1			[1963DAV/POP, 1982PIL/SKI]
C₉H₁₄Sn	[934-56-5]	phenyltrimethyltin				

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound		T_m (K)	Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)			
	$\Delta_v H$		52.3 ± 4.2			[1959PED/SKI, 1982PIL/SKI]
C₁₀H₁₆Sn	[4314-94-7]	benzyltrimethyltin				
	$\Delta_v H$		56.5 ± 4.2			[1959PED/SKI, 1982PIL/SKI]
C₁₀H₂₄O₂Sn	[14570-10-6]	triethyltin <i>tert</i> -butylperoxide				
	$\Delta_v H$		48.8 ± 2.1			[1971RAB/KIP, 1982PIL/SKI]
C₁₀H₂₅N₂Sn	[na]	(N,N-diethylamino)triethyltin				
	$\Delta_v H$		50.2 ± 4.2			[1971KOL/RAB, 1982PIL/SKI]
C₁₂H₂₇BrSn	[na]	tributyltin bromide				
	$\Delta_v H$		83.7 ± 12.6			[1959PED/SKI, 1982PIL/SKI]
C₁₂H₂₀Sn	[7393-43-3]	tetraallyltin				
	$\Delta_v H$	(333–393)	52.0		GC	[1992HAW]
C₁₂H₂₇ClSn	[1461-22-9]	tri- <i>n</i> -butyltin chloride				
	$\Delta_{\text{fus}} H$		11.43	260.2		[1974UTS/BAC2]
C₁₂H₂₈Sn	[2176-98-9]	tetrapropyl tin				
	$\Delta_v H$	(343–457)	55.0 ± 0.7	400		[2001BAE]
	$\Delta_v H$	(333–393)	60.8		GC	[1992HAW]
	$\Delta_v H$	(361–470)	52.5	376	A	[1987STE/MAL]
	$\Delta_v H$		65.4 ± 2.5	298	C	[1980ABR/IRV]
	$\Delta_v H$		66.9 ± 2.1			[1963DAV/POP, 1982PIL/SKI]
	$\Delta_v H$		60.7			[1935JON/EVA]
C₁₂H₂₈Sn	[2949-42-0]	tetraisopropyl tin				
	$\Delta_v H$	(342–441)	48.0 ± 0.7	392		[2001BAE]
	$\Delta_v H$	(333–393)	56.4		GC	[1992HAW]
	$\Delta_v H$		64.9 ± 4.2			[1966COL/SKI, 1982PIL/SKI]
C₁₂H₃₀OSn₂	[1112-63-6]	<i>bis</i> (triethyltin)oxide				
	$\Delta_v H$		52.3 ± 2.1			[1971RAB/KIP, 1982PIL/SKI]
C₁₂H₃₀Sn₂	[993-63-5]	hexaethyldistannane				
	$\Delta_v H$		62.8 ± 4.2			[1966TEL/RAB, 1982PIL/SKI]
C₁₅H₂₆O₂Sn	[na]	triethyltin dimethylphenylperoxide				
	$\Delta_v H$		56.5 ± 2.1			[1971RAB/KIP, 1982PIL/SKI]
C₁₆H₁₈Sn	[53561-93-6]	1,1-diphenylstannolane				
	$\Delta_{\text{sub}} H$		106.8 ± 5.5	298	B	[1988CAR/JAM]
C₁₆H₃₆Sn	[1461-25-2]	tetrabutyl tin				
	$\Delta_v H$	(389–462)	67.8 ± 0.5	425		[2001BAE]
	$\Delta_v H$		82.8 ± 2.1			[1963DAV/POP, 1982PIL/SKI]
C₁₆H₃₆Sn	[3531-43-9]	tetraisobutyl tin				
	$\Delta_v H$	(391–451)	53.6 ± 1.1	421		[2001BAE]
C₁₇H₂₀Sn	[19814-46-1]	hexahydro-1,1-diphenylstannin				
	$\Delta_v H$		75.0 ± 1.5			[1988CAR/JAM]
C₂₀H₁₈Sn	[2117-48-8]	triphenyl vinyl tin				
	$\Delta_{\text{sub}} H$		114.1			[1985CAR/LAY]
C₂₀H₁₈O₂Sn	[900-95-8]	(acetyloxy)triphenylstannane				
	$\Delta_{\text{fus}} H$		41.92	397.6		[1990DON/DRE]
C₂₄H₂₀Sn	[595-90-4]	tetraphenyl tin				

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)			
	$\Delta_{\text{fus}}H$		37.2	502.2		DSC	[1969VIC/WAL]
	$\Delta_{\text{sub}}H$	(393–461)	151.7	427		A	[1987STE/MAL]
	$\Delta_{\text{sub}}H$		161.1 ± 4.2	298			[1982PIL/SKI, 1969ADA/CAR2]
	$\Delta_{\text{sub}}H$	(428–454)	152.5 ± 0.6			TE	[1969KEI/KAN]
	$\Delta_{\text{sub}}H$	(393–461)	151.8 ± 1.1			ME	[1969KEI/KAN]
	$\Delta_{\text{sub}}H$		59.5	298			[1972NEW, 1986MAR/LOE]
	$\Delta_{\text{sub}}H$	(298–316)	U	298		ME	[1962CAR/COO, 1970CAR/LAY]
			66.0 ± 21.2				
C₂₇H₂₀Sn	[1247-08-1]	triphenyl phenylethynyl tin					
	$\Delta_{\text{sub}}H$		137.6				[1985CAR/LAY]
C₃₂H₁₆Cl₂N₈Sn	[18253-54-8]	tin(IV) phthalocyanine dichloride					
	$\Delta_{\text{sub}}H$		218.4 ± 17.6			ME	[1970BON/CAT]
C₃₂H₁₆N₈Sn	[15304-57-1]	tin(II) phthalocyanine					
	$\Delta_{\text{sub}}H$		123.4 ± 10.0			ME	[1970BON/CAT]
C₃₆H₃₀Sn₂	[1064-10-4]	hexaphenyl ditin					
	$\Delta_{\text{sub}}H$		188.3 ± 4.2	298		ME,TE	[1969KEI/KAN]
C₄₄H₂₆N₈Sn	[219130-47-0]	diphenyl tin(IV) phthalocyanine					
	$\Delta_{\text{sub}}H$		174.9 ± 18.8			ME	[1970BON/CAT]
C₆₀H₇₈OSn₂	[na]	hexakis(2-methyl-2-phenylpropyl)distanoxane					
	$\Delta_{\text{fus}}H$		71.81	417.7		DSC	[1990DON/DRE]
SnBr₄	[7789-67-5]	stannic bromide					
	$\Delta_{\text{sub}}H$	(257–299)	62.4	278			[1941SEK]
SnI₄	[7790-47-8]	stannic iodide					
	Δ_vH	(418–523)	57.2	423			[1936NEG]
	$\Delta_{\text{sub}}H$	(366–414)	75.6	390			[1941SEK]
Sr (strontium)							
SrCl₂	[10476-85-4]	strontium chloride					
	$\Delta_{\text{sub}}H$		328.9 ± 4.8	298		LE	[1965LOE/KEN]
Ta (tantalum)							
C₅H₁₅O₅Ta	[865-35-0]	tantalum pentamethoxide					
	$\Delta_{\text{sub}}H$		88.3 ± 13.4			ME,E	[1972TEL/RAB]
C₁₀H₂₅O₅Ta	[na]	pentaethyltantalate					
	Δ_vH	(388–424)	72.6	403		A	[1987STE/MAL]
TaBr₅	[13451-11-1]	tantalum(V) pentabromide					
	$\Delta_{\text{sub}}H$		127 ± 18	298			[1996TUR/EIC]
	$\Delta_{\text{sub}}H$		121.9	298			[1996TUR/EIC, 1991KNA/KUB]
TaI₅	[14693-81-3]	tantalum(V) pentaiodide					
	$\Delta_{\text{sub}}H$	(573–655)	120.9				[1978ABA/MAL]
Tb (terbium)							
C₁₅H₁₅Tb	[1272-25-9]	tris(cyclopentadienyl)terbium(III)					
	$\Delta_{\text{sub}}H$		103.8 ± 1.7				[1973DEV/BOR]
C₃₂H₄₀F₁₂NaO₈Tb	[12576-88-4]	sodium tetrakis(1,1,1-trimethyl-5,5,5-trifluoro-2,4-pentanedionato)terbate					
	$\Delta_{\text{sub}}H$	(418–473)	163 ± 3	445		T	[1993SYO/GOL]

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)			
C₃₃H₅₇O₆Tb	[15492-51-0]	<i>tris</i> (2,2,6,6-tetramethylheptane-3,5-dionato)terbium(III)					
	$\Delta_{\text{sub}}H$		138.4 ± 2.6	298	DSC	[1999SAN/PET]	
	$\Delta_{\text{sub}}H$	(373–420)	173.6	396	ME	[1981AMA/SAT]	
	$\Delta_{\text{sub}}H$	(420–433)	151	426	ME	[1981AMA/SAT]	
	$\Delta_{\text{sub}}H$	(420–454)	141.5	437	BG	[1969SIC/DUB]	
	Δ_vH	(454–500)	87		BG	[1969SIC/DUB]	
Te (tellurium)							
C₂H₆Te	[593-80-6]	dimethyl telluride					
	$\Delta_{\text{fus}}H$		0.7	164.1			
	$\Delta_{\text{fus}}H$		7.77	201.9		[1998SHE/NIS]	
	Δ_vH	(298–367)	34.4	313		[1999DYK/SVO]	
	Δ_vH	(273–372)	35.6 ± 0.1	323		[1997BAE, 1996BAE/POD]	
	Δ_vH	(267–369)	36.9	282	BG	[1996VAN/COR]	
	Δ_vH	(267–369)	36.1 ± 1.0	298	BG	[1996VAN/COR]	
	Δ_vH		37.4 ± 0.7	298	C	[1989VOR/KLY]	
	Δ_vH		36.0 ± 2.1			[1988TEL/LAR]	
C₄H₆Te	[63000-06-6]	divinyl telluride					
	$\Delta_{\text{fus}}H$		9.59	182.6		[1999NIS/SHE]	
	Δ_vH		44.8 ± 0.8	298	C	[1989VOR/KLY]	
	Δ_vH		38.1 ± 2.1			[1988TEL/LAR]	
C₄H₁₀Te	[627-54-3]	diethyl telluride					
	$\Delta_{\text{fus}}H$		7.62	161.5		[1996SHE/KAM]	
	Δ_vH	(295–411)	41.8	310		[1999DYK/SVO]	
	Δ_vH	(273–415)	41.6 ± 0.2	344		[1996BAE/POD]	
	Δ_vH		41.6 ± 0.8	298	C	[1989VOR/KLY]	
C₆H₁₄Te	[64501-17-3]	dipropyl telluride					
	Δ_vH	(298–434)	45.5 ± 0.3	366		[1996BAE/POD]	
	Δ_vH		46.5 ± 0.7	298	C	[1989VOR/KLY]	
C₆H₁₄Te	[51112-72-2]	diisopropyl telluride					
Δ_vH	(298–399)	40.4 ± 0.1	349		[1996BAE/POD]		
C₆H₁₄Te₂	[79971-42-9]	dipropyl ditelluride					
Δ_vH		52.7 ± 1.0	298	C	[1989VOR/KLY]		
C₈H₁₈Te	[38788-38-4]	dibutyl telluride					
	Δ_vH	(303–423)	53.4 ± 0.1	358		[1996BAE/POD]	
	Δ_vH		51.0 ± 1.0	298	C	[1989VOR/KLY]	
C₈H₁₈Te	[82817-01-0]	diisobutyl telluride					
Δ_vH	(303–410)	47.6 ± 0.1	356		[1996BAE/POD]		
C₈H₁₈Te	[83817-20-3]	di-sec-butyl telluride					
Δ_vH	(303–372)	49.6 ± 0.9	338		[1996BAE/POD]		
C₈H₁₈Te₂	[77129-69-2]	dibutyl ditelluride					
Δ_vH		57.3 ± 1.0	298	C	[1989VOR/KLY]		
C₁₀H₂₂Te	[71475-88-2]	dipentyl telluride					
	$\Delta_{\text{fus}}H$		23.1	215.4		[1994TEL/SHE]	
	Δ_vH	(343–403)	59.5 ± 0.8	373		[1996BAE/POD]	

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
C₁₀H₂₂Te	[110346-75-3] $\Delta_v H$	diisopentyl telluride (343–403)	51.9 ± 0.7	373		[1996BAE/POD]
C₁₂H₁₀Te	[1202-36-4] $\Delta_{\text{fus}} H$	diphenyl telluride	15.35	268.4		[1996TEL/SHE]
TeCl₄	[10026-07-0] $\Delta_{\text{sub}} H$ $\Delta_v H$	tellurium tetrachloride (506–660)	105 ± 2 77	298 583		[1994DAL/FER] GS [1930SIM]
TeF₆	[7783-80-4] $\Delta_{\text{sub}} H$	tellurium hexafluoride (194–233)	25.6	214		[1932KLE/HEN]
TeI₄	[7790-48-9] $\Delta_{\text{sub}} H$	tellurium tetraiodide (420–480)	95.1			[2007KUT/POL]
Th (thorium)						
C₂₀H₁₆F₁₂O₈Th	[17500-72-0] $\Delta_{\text{sub}} H$	<i>tetrakis</i> (1,1,1-trifluoropentan-2,4-dionato)thorium(IV)	154.6	298	GS,HA	[1986GAR/JAN]
C₄₀H₄₀F₂₈O₈Th	[23841-30-7] $\Delta_{\text{sub}} H(\alpha)$ $\Delta_{\text{sub}} H(\beta)$ $\Delta_{\text{sub}} H$	<i>tetrakis</i> (1,1,1,2,2,3,3-heptafluoro-7,7-dimethyloctane-4,6-dionato)thorium (IV) (344–367)	151.2 130.6 138.5 ± 3.3	298 298 355	GS,HA GS,HA ME	[1986GAR/JAN] [1986GAR/JAN] [1970SWA/KAR]
C₄₄H₇₆O₈Th	[18865-73-1] $\Delta_{\text{sub}} H$	<i>tetrakis</i> (2,2,6,6-tetramethylheptan-3,5-dionato)thorium(IV) (391–409)	152.3 ± 3.3	400	ME	[1970SWA/KAR]
Ti (titanium)						
(C₂H₃N)–Cl₄Ti	[13682-81-0] $\Delta_{\text{sub}} H$	titanium trichloride- acetonitrile (1:1 complex)	123		T	[1970HES/PER]
2(C₂H₃N)–Cl₄Ti	[15227-64-2] $\Delta_{\text{sub}} H$	titanium trichloride- acetonitrile (1:2 complex)	171.5		T	[1970HES/PER]
(C₄H₈O)–(Cl₄Ti)	[15005-09-1] $\Delta_{\text{sub}} H$	titanium trichloride- tetrahydrofuran (1:1 complex)	140.2		T	[1970HES/PER]
2(C₄H₈O)–(Cl₄Ti)	[31011-57-1] $\Delta_{\text{sub}} H$	titanium trichloride- tetrahydrofuran (1:2 complex)	205.4		T	[1970HES/PER]
(C₄H₈S)–(Cl₄Ti)	[14281-72-2] $\Delta_{\text{sub}} H$	titanium trichloride- tetrahydrothiophene (1:1 complex)	124.3		T	[1970HES/PER]
2(C₄H₈S)–(Cl₄Ti)	[16893-00-8] $\Delta_{\text{sub}} H$	titanium trichloride- tetrahydrothiophene (1:2 complex)	181.2		T	[1970HES/PER]
C₅H₅Cl₃Ti	[1270-98-0] $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$ $\Delta_{\text{sub}} H$	cyclopentadienyltitanium trichloride (354–404)	89.8 104.6 ± 8.4 89.1 ± 0.8	379 298	A	[1987STE/MAL] [1982PIL/SKI, 1977TEL/RAB] [1977BAL/BAR]
(C₅H₁₀O)–(Cl₄Ti)	[22538-12-1] $\Delta_{\text{sub}} H$	titanium trichloride- tetrahydropyran (1:1 complex)	139.3		T	[1970HES/PER]
2(C₅H₁₀O)–(Cl₄Ti)	[31011-56-0] $\Delta_{\text{sub}} H$	titanium trichloride- tetrahydropyran (1:2 complex)	305.4		T	[1970HES/PER]
(C₈H₈O)–(Cl₄Ti)	[31011-60-6] $\Delta_{\text{sub}} H$	titanium trichloride- acetophenone (1:1 complex)	163.6		T	[1970HES/PER]

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound			Method	Reference	
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)			
2(C ₈ H ₈ O)–(Cl ₄ Ti)	[31011-61-7] $\Delta_{\text{sub}}H$	titanium trichloride- acetophenone (1:2 complex)	277.8		T	[1970HES/PER]	
C ₈ H ₂₄ N ₄ Ti	[3275-24-9] Δ_vH	titanium tetradimethylamide (353–418)	53.8 ± 3.0	383		[1984BAE/MIK, 2001BAE/MIK]	
C ₁₀ H ₁₀ Ti	[1271-29-0] $\Delta_{\text{sub}}H$	<i>bis</i> (cyclopentadienyl)titanium	58.5 ± 8.0	298		[1982PIL/SKI, 1971TEL/RAB]	
C ₁₀ H ₁₀ Cl ₂ Ti	[1271-19-8] $\Delta_{\text{sub}}H$	<i>bis</i> (cyclopentadienyl)titanium dichloride (418–533)	124.4 ± 2.9	298	ME	[2001DIO/PIE]	
	$\Delta_{\text{sub}}H$		124.4	475.5	A	[1987STE/MAL]	
	$\Delta_{\text{sub}}H$		118.8 ± 2.1	298		[1982PIL/SKI, 1977TEL/RAB]	
	$\Delta_{\text{sub}}H$		111.7 ± 1.7			[1977BAL/BAR]	
	$\Delta_{\text{sub}}H$		96.2			[1969DIL/KIS]	
	$\Delta_{\text{sub}}H$		102 ± 13	298		[1968KIS/DIL, 2001DIO/PIE]	
C ₁₂ H ₁₀ O ₂ Ti	[12129-51-0] $\Delta_{\text{sub}}H$	<i>bis</i> (cyclopentadienyl)dicarbonyl titanium	84.2 ± 3.5	298	ME	[1987DIA/DIA]	
C ₁₂ H ₁₆ Ti	[1271-66-5] $\Delta_{\text{sub}}H$	<i>bis</i> (cyclopentadienyl)dimethyltitanium	79.5 ± 8.4	298		[1982PIL/SKI, 1977TEL/RAB]	
C ₁₂ H ₂₈ O ₄ Ti	[546-68-9] Δ_vH	tetraisopropyl titanate (336–459)	62.3	351	A	[1987STE/MAL]	
C ₁₂ H ₂₈ O ₄ Ti	[3087-37-4] Δ_vH	tetrapropyl titanate (411–479)	111.9	426	A	[1987STE/MAL]	
(C ₁₃ H ₁₀ O)–(Cl ₄ Ti)	[23368-15-2] $\Delta_{\text{sub}}H$	titanium trichloride- benzophenone (1:1 complex)	249.4		T	[1970HES/PER]	
2(C ₁₃ H ₁₀ O)–(Cl ₄ Ti)	[31011-63-9] $\Delta_{\text{sub}}H$	titanium trichloride- benzophenone (1:2 complex)	287.9		T	[1970HES/PER]	
C ₁₄ H ₁₀ F ₆ O ₄ Ti	[1282-45-7] $\Delta_{\text{sub}}H$	<i>bis</i> (cyclopentadienyl)titanium <i>bis</i> (trifluoroacetate)	108.0 ± 8.0	298		[1982PIL/SKI, 1981CAL/DIA]	
C ₁₆ H ₃₆ O ₄ Ti	[5593-70-4] Δ_vH	tetrabutoxy titanium (462–564)	89.7	477	A	[1987STE/MAL]	
	Δ_vH		(443–493)	85.0 ± 3.1	458	A	[1987STE/MAL, 1978GRA/KON]
C ₁₆ H ₃₆ O ₄ Ti	[7425-80-1] Δ_vH	tetraisobutoxy titanium (436–529)	77.4	451	A	[1987STE/MAL]	
C ₁₆ H ₃₆ O ₄ Ti	[873376-17-1] Δ_vH	tetra- <i>sec</i> -butoxy titanium (378–414)	76.8	393	A	[1987STE/MAL]	
	Δ_vH		(370–476)	67.1	385	A	[1987STE/MAL]
C ₁₆ H ₃₆ O ₄ Ti	[119279-48-0] Δ_vH	tetra- <i>tert</i> -butoxy titanium (386–486)	55.9	401	A	[1987STE/MAL]	
	Δ_vH		(322–388)	62.6	337	SG	[1958BRA/SWA, 1984BOU/FRI]
	Δ_vH			66.1 ± 3.3	298	SG	[1958BRA/SWA, 1966BRA/HIL]
C ₁₆ H ₃₆ O ₄ Ti	[5593-70-4] Δ_vH	titanium (IV) tetrabutylate (323–418)	47.6 ± 0.7	370		[2002BAE/SHI2]	
C ₁₆ H ₄₀ N ₄ Ti	[na] Δ_vH	titanium (IV) <i>tetrakis</i> (diethylamide) (423–463)	94.6 ± 4.0	443		[2001BAE/MIK]	
C ₂₀ H ₄₄ O ₄ Ti	[na] Δ_vH	<i>tetrakis</i> (1,1-dimethylpropoxy)titanium (397–430)	67.8	412	A	[1987STE/MAL]	

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_v H$	(361–423)	71.0	376	SG	[1958BRA/SWA, 1984BOU/FRI]
	$\Delta_v H$		77.4 ± 3.8	298	SG	[1958BRA/SWA, 1966BRA/HIL]
C₂₀H₄₄O₄Ti	[na]	<i>tetrakis</i> (1-ethylpropoxy)titanium				
	$\Delta_v H$	(385–445)	103.6	400	A	[1987STE/MAL]
C₂₀H₄₄O₄Ti	[na]	<i>tetrakis</i> (3-methylbutoxy)titanium				
	$\Delta_v H$	(407–493)	119.7	422	A	[1987STE/MAL]
C₂₀H₄₄O₄Ti	[10585-24-7]	tetrapentoxytitanium				
	$\Delta_v H$	(484–558)	103.4	499	A	[1987STE/MAL]
C₂₀H₄₄O₄Ti	[10585-26-9]	tetra- <i>tert</i> -pentoxytitanium				
	$\Delta_v H$	(361–423)	71.1	376	A	[1987STE/MAL]
C₂₂H₂₀Ti	[1273-09-2]	<i>bis</i> (cyclopentadienyl)diphenyltitanium				
	$\Delta_{\text{sub}} H$		88 ± 8			[1982DIA/SAL]
C₂₂H₄₀O₈Ti	[852536-12-0]	<i>bis</i> [1,1-diemthylethyl-3-oxobutanoato] <i>bis</i> (2-propanolato)titanium				
	$\Delta_{\text{sub}} H$		85.4			[2008KUN/SHI]
C₂₄H₂₀O₄Ti	[12156-48-8]	<i>bis</i> (benzoato) <i>bis</i> (η^5 -2,4-cyclopentadien-1-yl)titanium				
	$\Delta_{\text{sub}} H$		112 ± 8			[1981CAL/DIA]
C₂₄H₂₄Ti	[See Note]	<i>bis</i> (cyclopentadienyl)dibenzyltitanium				
	$\Delta_{\text{sub}} H$		83.7 ± 8.4	298		[1982PIL/SKI, 1977TEL/RAB]
		Note: There is no reference to [1977TEL/RAB] in Chemical Abstracts under the given chemical name. Rather, Chemical Abstracts lists the paper under <i>bis</i> (cyclopentadienyl)diphenyltitanium.				
C₂₄H₅₂O₄Ti	[na]	<i>tetrakis</i> (1,1-dimethylbutoxy)titanium				
	$\Delta_v H$	(414–454)	94.6	429	A	[1987STE/MAL]
C₂₄H₅₂O₄Ti	[na]	<i>tetrakis</i> (1-methyl-ethylpropoxy)titanium				
	$\Delta_v H$	(412–460)	86.2	427	A	[1987STE/MAL]
C₂₄H₅₂O₄Ti	[na]	tetrahexyloxy titanium				
	$\Delta_v H$	(520–581)	94.8	535	A	[1987STE/MAL]
C₂₈H₅₂O₆Ti	[80570-88-3]	diisopropoxy <i>bis</i> (2,2,6,6-tetramethyl-3,5-heptanedionato) titanium				
	$\Delta_{\text{sub}} H$	(353–413)	98.6 ± 2.7	383		[2001TUR/KRI]
	$\Delta_{\text{sub}} H$	(273–403)	104.1	338		[1997KOJ/KAD, 2001TUR/KRI]
C₃₀H₂₈Fe₂Ti	[65274-19-3]	<i>bis</i> (cyclopentadienyl)diferrocenyl titanium				
	$\Delta_{\text{sub}} H$		150 ± 15			[1982DIA/SAL]
Br₄Ti	[7789-68-6]	titanium (IV) tetrabromide				
	$\Delta_{\text{sub}} H$	(283–306)	62.4	294		[1941SEK]
Cl₄Ti	[7550-45-0]	titanium (IV) tetrachloride				
	$\Delta_v H$	(250–423)	37.5	265		[1966LUC]
	$\Delta_v H$	(363–415)	37.9	378		[1959PIK/FOS]
	$\Delta_v H$	(313–357)	39.8	335	I	[1953SCH/ZEP]
F₃Ti	[13470-08-1]	titanium (III) trifluoride				
	$\Delta_{\text{sub}} H$	(759–865)	237.2 ± 1.7	810		[1967ZMB/MAR]
I₄Ti	[7720-83-4]	titanium (IV) tetraiodide				
	$\Delta_v H$	(433–643)	58.5	538		[1947BLO/CAM]
Tl (thallium)						
C₃H₉Tl	[3003-15-4]	trimethylthallium				

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound		T_m (K)	Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)			
	$\Delta_{\text{fus}}H$		16.74	311.2		[1965PRI/JAC]
	$\Delta_{\text{sub}}H$	(258–304)	57.3	285	CATH	[1965PRI/JAC, 1987STE/MAL]
	Δ_vH	(311–360)	40.6	335	I,MM	[1965PRI/JAC]
	Δ_vH	(328–349)	37.9	338	I	[1946GIL/JON]
C₆H₁₅Tl	[687-82-1]	triethylthallium				
	Δ_vH	(282–465)	41.9	297		[1947STU]
TlF	[7789-27-7]	thallium(I) fluoride				
	$\Delta_{\text{sub}}H$		142.7	298		[1967KEN/CUB]
Tm (thulium)						
C₁₅H₁₅Tm	[1272-26-0]	<i>tris</i> (cyclopentadienyl)thulium				
	$\Delta_{\text{sub}}H$		111.3 ± 3.5	298		[1982PIL/SKI, 1974DEV/RAB]
	$\Delta_{\text{sub}}H$		98.7 ± 1.7			[1973DEV/BOR]
	$\Delta_{\text{sub}}H$	(338–438)	109.2 ± 2.1		ME	[1971HAU, 1971HAU2]
C₃₃H₅₇O₆Tm	[15631-58-0]	<i>tris</i> (2,2,6,6-tetramethylheptane-3,5-dionato)thulium(III)				
	$\Delta_{\text{sub}}H$		131.3 ± 2.9	298	DSC	[1999SAN/PET]
	$\Delta_{\text{sub}}H$	(363–418)	156.1	390	ME	[1981AMA/SAT]
	$\Delta_{\text{sub}}H$	(410–446)	131.4	428	BG	[1969SIC/DUB]
	Δ_vH	(446–490)	84.1		BG	[1969SIC/DUB]
U (uranium)						
C₆H₁₈O₆U	[69644-82-2]	uranium hexamethoxide				
	$\Delta_{\text{sub}}H$		102.9 ± 8.4			[1991TEL/LAR]
C₁₀H₂F₁₂O₆U	[67316-66-9]	<i>bis</i> (1,1,1,5,5,5-hexafluoro-2,4-pentanedionato)uranium dioxide complex				
	$\Delta_{\text{sub}}H$	(370–425)	147	397.5	A	[1987STE/MAL]
	$\Delta_{\text{sub}}H$	(423–470)	147 ± 4			[1978EKS/RAN]
C₁₅H₁₅ClU	[11087-14-2]	<i>tris</i> (cyclopentadienyl)uranium chloride				
	$\Delta_{\text{sub}}H$	(338–348)	115.9 ± 2.1		ME	[1971HAU, 1971HAU2]
C₁₆H₁₆U	[na]	<i>bis</i> (cyclooctatetraene)uranium				
	$\Delta_{\text{sub}}H$	(400–500)	107.9 ± 3.3			[1979TEL/RAB, 1977BED]
	$\Delta_{\text{sub}}H$		114.2 ± 4.8	298		[1979TEL/RAB, 1977BED]
C₂₀H₂₀F₃₀O₁₀U₂	[137220-74-7]	<i>bis</i> [<i>pentakis</i> (trifluoroethoxy)]diuranium				
	$\Delta_{\text{sub}}H$		NA			[1991SEV/ALI]
C₂₀H₂₂Cl₂F₁₂O₆U	[136211-24-0]	<i>bis</i> (1,1,1,5,5,5-hexafluoro-2,4-pentanedionato)dichlorouranium- <i>bis</i> (tetrahydropyran)				
	$\Delta_{\text{sub}}H$	(316–387)	79.1	352	T	[1991GIL/SAG]
C₂₀H₂₈O₈U	[65137-03-3]	<i>tetrakis</i> (pentane-2,4-dionato)uranium(IV)				
	$\Delta_{\text{sub}}H$		148.1 ± 4.6			[1991TEL/LAR]
C₂₂H₃₈O₆U	[50707-86-9]	<i>bis</i> (2,2,6,6-tetramethylheptane-3,5-dionato)dioxouranium				
	$\Delta_{\text{sub}}H$	(370–412)	151.6 ± 1.9	404	ME	[1993RIB/MON]
	$\Delta_{\text{sub}}H$		156.9 ± 1.9	298		[1993RIB/MON]
	$\Delta_{\text{sub}}H$		126 ± 9			[1978EKS/RAN]
C₄₀H₄₀F₂₈O₈U	[23797-50-4]	<i>tetrakis</i> (1,1,1,2,2,3,3-heptafluoro-7,7-dimethyloctane-4,6-dione)uranium(IV)				
	$\Delta_{\text{sub}}H$		68.2		BG	[1977DES]
	$\Delta_{\text{sub}}H$		64.3 ± 3.2		C	[1977DES]
	$\Delta_{\text{sub}}H$	(343–367)	143.5 ± 1.3	355	ME	[1970SWA/KAR]

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)			
C ₄₀ H ₆₈ O ₁₂ U	[133952-93-9] $\Delta_{\text{sub}}H$	<i>tetrakis</i> (2,6-dimethyl-2-methoxy-3,5-heptanedionato)uranium(IV) (344–377)	121.7 ± 18	350		[1991SEV/KRA]	
C ₄₄ H ₇₆ O ₈ U	[na] $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$	<i>tetrakis</i> (2,2,6,6-tetramethyl-3,5-heptanedionato)uranium(IV) (372–478) (392–409)	152.2 ± 3.3 149.0 ± 1.3	425 400	ME ME	[1977BED/HUS] [1970SWA/KAR]	
C ₄₄ H ₇₆ O ₁₂ U	[133952-92-8] $\Delta_{\text{sub}}H$	<i>tetrakis</i> (2,6,6-trimethyl-2-methoxy-3,5-heptanedionato)uranium(IV) (387–428)	160.7 ± 6.3	408		[1991SEV/KRA]	
UF ₆	[7783-81-5] Δ_vH	uranium hexafluoride (337–389)	29.5	352		[1953OLI/MIL]	
V (vanadium)							
C ₁₀ H ₈ F ₆ O ₅ V	[52081-94-4] $\Delta_{\text{sub}}H$	<i>bis</i> (1,1,1-trifluoro-2,4-pentanedionato)oxovanadium(IV) (423–473)	119.7 ± 0.8			[1985MAT/KUW]	
C ₁₀ H ₁₀ Cl ₂ V	[12083-48-6] $\Delta_{\text{sub}}H$	<i>bis</i> (cyclopentadienyl)vanadium dichloride 140.1 ± 7.4		298	ME	[2001DIO/PIE]	
C ₁₀ H ₁₀ V	[1277-47-0] $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$	<i>bis</i> (cyclopentadienyl)vanadium (323–338)	57.4 58.6 ± 4.2	330.5 298	A	[1987STE/MAL] [1982PIL/SKI, 1971TEL/RAB]	
C ₁₀ H ₁₄ O ₅ V	[3153-26-2] $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$	<i>bis</i> (2,4-pentanedionato)oxovanadium(IV) (418–443)	140.7 ± 4.0 91.5 192.4 140.4 ± 1.1	493 430.5 461 298	DSC A C C	[1987MUR/HIL2] [1987STE/MAL] [1986JAM/PIL] [1986JAM/PIL]	
C ₁₀ H ₁₇ NO ₅ V	[122343-53-7] $\Delta_{\text{sub}}H$	amine <i>bis</i> (pentane-2,4-dionato)oxovanadium 29.0		370	DSC	[1989SHI/SHI]	
C ₁₂ H ₁₂ V	[12129-72-5] $\Delta_{\text{sub}}H$	<i>bis</i> (benzene)vanadium 70 ± 10				[1982PIL/SKI]	
C ₁₂ H ₂₇ O ₄ V	[1801-76-9] Δ_vH	vanadic acid, tributyl ester (395–435)	90.2	410	A	[1987STE/MAL]	
C ₁₂ H ₂₇ O ₄ V	[19120-62-8] Δ_vH	vanadic acid, triisobutyl ester (383–418)	82.2	398	A	[1987STE/MAL]	
C ₁₂ H ₂₇ O ₄ V	[17838-66-3] Δ_vH	vanadic acid, tri-sec-butyl ester (378–413)	82.4	393	A	[1987STE/MAL]	
C ₁₂ H ₂₇ O ₄ V	[1686-24-4] Δ_vH	vanadic acid, tri-tert-butyl ester (348–385)	71.4	363	A	[1987STE/MAL]	
C ₁₄ H ₁₆ V	[36955-47-2] $\Delta_{\text{sub}}H$	benzene(ethylbenzene)vanadium (453–483)	69.5	468		[1972UMI/VAN]	
C ₁₅ H ₁₂ F ₉ O ₆ V	[15695-88-2] $\Delta_{\text{sub}}H$	<i>tris</i> (1,1,1-trifluoro-2,4-pentanedionato)vanadium(III) (383–433)	118.4 ± 2.1			[1985MAT/KUW]	
C ₁₅ H ₁₈ V	[36955-49-4] $\Delta_{\text{sub}}H$	benzene(isopropylbenzene)vanadium (453–483)	83.7	468		[1972UMI/VAN]	
C ₁₅ H ₁₈ BrNO ₅ V	[24263-16-9] $\Delta_{\text{sub}}H$	3-bromopyridine <i>bis</i> (acetylacetonato)oxovanadium 59.4		402	DSC	[1989SHI/SHI]	
C ₁₅ H ₁₉ NO ₅ V	[24263-31-8]	pyridine <i>bis</i> (acetylacetonato)oxovanadium					

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
	$\Delta_{\text{sub}}H$				47.8	404	DSC	[1989SHI/SHI]
$C_{15}H_{21}O_6V$	[13476-99-8]	<i>tris</i> (2,4-pentanedionato)vanadium(III)						
	$\Delta_{\text{fus}}H$				30.0	460		[1971BEE/LIN2]
	$\Delta_{\text{fus}}H$				23.8	460		[1970MEL/MER2]
$C_{16}H_{18}N_2O_5V$	$\Delta_{\text{sub}}H$				102.9 ± 0.8	298	HSA	[1970MEL/MER, 1970MEL/MER2]
	[24263-13-6]	3-cyanopyridine <i>bis</i> (acetylacetonato)oxovanadium						
$C_{16}H_{18}N_2O_5V$	$\Delta_{\text{sub}}H$				79	391	DSC	[1989SHI/SHI]
	[24263-14-7]	4-cyanopyridine <i>bis</i> (acetylacetonato)oxovanadium						
$C_{16}H_{20}V$	[36955-48-3]	<i>bis</i> (ethylbenzene)vanadium						
	$\Delta_{\text{sub}}H$		(453–483)		72.0	468		[1972UMI/VAN]
$C_{16}H_{21}NO_5V$	[24263-33-0]	4-methylpyridine <i>bis</i> (acetylacetonato)oxovanadium						
	$\Delta_{\text{sub}}H$				56.9	421	DSC	[1989SHI/SHI]
$C_{18}H_{24}V$	[36472-53-4]	<i>bis</i> (isopropylbenzene)vanadium						
	$\Delta_{\text{sub}}H$		(453–483)		86.2	468		[1972UMI/VAN]
$C_{32}F_{16}N_8OV$	[128675-60-5]	(hexadecafluorophthalocyaninato)oxovanadium						
	Δ_vH		(590–670)		52.7 ± 1.0	630	ME	[2008SEM/BAS]
$C_{32}H_{16}N_8OV$	[13930-88-6]	oxovanadium phthalocyanine						
	Δ_vH		(578–672)		46.5 ± 0.7	625	ME	[2008SEM/BAS]
W (tungsten)								
C_6O_6W	[14040-11-0]	tungsten hexacarbonyl						
	$\Delta_{\text{sub}}H$		(265–288)		77.7	276	TE	[1995GAR/CHA]
	$\Delta_{\text{sub}}H$		(338–423)		74.9 ± 1.3			[1993BAE]
	$\Delta_{\text{sub}}H$		(333–433)		74.4	348	A	[1987STE/MAL]
	$\Delta_{\text{sub}}H$		(250–292)		78.9 ± 1.1	271	ME	[1980BOX/ERN, 1979DAA/ERN]
	$\Delta_{\text{sub}}H$				73.2	298	C	[1975ADE/BRO]
	$\Delta_{\text{sub}}H$				76.5 ± 1.3			[1975PIL/WAR]
	$\Delta_{\text{sub}}H$		(339–410)		69.7			[1952REZ/SHV]
$\Delta_{\text{sub}}H$				74.1			[1935HIE/ROM]	
$C_7H_3NO_5W$	[15096-68-1]	acetonitrile tungsten pentacarbonyl						
	$\Delta_{\text{sub}}H$		(271–303)		98.1 ± 2.0	298		[1980CAV/ERN]
$C_8H_4N_2O_5W$	[39017-11-3]	pyrazole(pentacarbonyl)tungsten						
	$\Delta_{\text{sub}}H$		(287–327)		112.5 ± 2.4	307	ME	[1979DAA/ERN]
$C_8H_6N_2O_4W$	[16800-45-6]	<i>bis</i> (acetonitrile)tetracarbonyltungsten						
	$\Delta_{\text{sub}}H$		(294–313)		131.0 ± 6.0	298		[1980CAV/ERN]
$C_8H_9NO_5W$	[15228-32-7]	trimethylamine(pentacarbonyl)tungsten						
	$\Delta_{\text{sub}}H$				89.1 ± 2.1			[1979DAA/ERN, 1980BOX, 1980BOX/ERN]
$C_8H_9O_5PW$	[26555-11-3]	trimethylphosphine(pentacarbonyl)tungsten						
	$\Delta_{\text{sub}}H$		(283–327)		93.8 ± 1.5	305	ME	[1980BOX/ERN]
$C_9H_4N_2O_5W$	[65761-19-5]	pyrazine(pentacarbonyl)tungsten						
	$\Delta_{\text{sub}}H$		(287–321)		108.4 ± 1.3	304	ME	[1979DAA/ERN]
$C_9H_4N_2O_5W$	[65761-20-8]	pyridazine(pentacarbonyl)tungsten						
	$\Delta_{\text{sub}}H$				106.4 ± 2.5			[1979DAA/ERN, 1980BOX, 1980BOX/ERN]
$C_9H_9N_3O_3W$	[16800-47-8]	<i>tris</i> (acetonitrile) tungsten tricarbonyl						
	$\Delta_{\text{sub}}H$		(308–333)		103.4 ± 6.0	298		[1980CAV/ERN]

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)	Method	Reference
$C_{10}H_5NO_5W$	[14586-49-3] $\Delta_{\text{sub}}H$	pyridine(pentacarbonyl)tungsten (285–313)	109.7 ± 2.7	299	ME	[1979DAA/ERN]
$C_{10}H_8O_3W$	[12128-81-3] $\Delta_{\text{sub}}H$	cycloheptatrienetungstentricarbonyl	92	298	C	[1977BRO/CON, 1982PIL/SKI]
$C_{10}H_{10}Cl_2W$	[12184-31-5] $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$	dichlorobis(η^5 -2,4-cyclopentadien-1-yl)tungsten	120.7 ± 8.6 104.6 ± 4.2	298	ME	[2001DIO/PIE] [1976TEL/RAB]
$C_{10}H_{10}I_2W$	[12184-31-5] $\Delta_{\text{sub}}H$	bis(η^5 -2,4-cyclopentadien-1-yl)diiodotungsten	104.6 ± 4.2			[1976TEL/RAB]
$C_{10}H_{11}NO_5W$	[31082-68-5] $\Delta_{\text{sub}}H$	piperidine(pentacarbonyl)tungsten (289–327)	106.4 ± 1.0	308	ME	[1979DAA/ERN]
$C_{10}H_{12}W$	[1271-33-6] $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$	dicyclopentadienyttungsten dihydride (313–323)	84.6 ± 1.6 96.2 ± 2.1	298	ME	[1990DIA/DIO] [1982PIL/SKI, 1979CAL/DIA, 1976TEL/RAB]
$C_{12}H_{12}W$	[12089-23-5] $\Delta_{\text{sub}}H$	dibenzene tungsten	106	298	ME	[1974ZOR/UMI]
$C_{12}H_{16}W$	[39333-53-4] $\Delta_{\text{sub}}H$	bis(η^5 -2,4-cyclopentadien-1-yl)dimethyltungsten	74.6 ± 4.2			[1980DEP]
$C_{12}H_{36}N_6W$	[68941-84-4] $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$	hexakis(dimethylamino)tungsten	164.0 ± 5 89.1 ± 7	461 298	C C	[1979ADE/CAV] [1979ADE/CAV]
$C_{12}H_{36}N_6W_2$	[54935-70-5] $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$	hexakis(dimethylamino)ditungsten	186.5 ± 5 113.3 ± 6	451 298	C C	[1979ADE/CAV] [1979ADE/CAV]
$C_{23}H_{15}O_5PW$	[26555-11-3] $\Delta_{\text{sub}}H$	triphenylphosphine(pentacarbonyl)tungsten (340–364)	162.2 ± 8.3	352	ME	[1980BOX/ERN]
$C_{23}H_{15}O_8PW$	[23306-41-4] $\Delta_{\text{sub}}H$	triphenylphosphite(pentacarbonyl)tungsten (308–348)	120.2 ± 6.6	328	ME	[1980BOX/ERN]
WCl_4O	[13520-78-0] $\Delta_{\text{sub}}H$	tungsten(IV) oxychloride (396–447)	63.7 ± 1.7	421	DSM	[1983CAS/PON]
WF_6	[7783-82-6] Δ_vH	tungsten hexafluoride (290–343)	25.8	316		[1968NIS/NIK]
Xe (xenon)						
XeF_2	[13709-36-9] Δ_vH	xenon difluoride (553–663)	53.5	568		[1983HOU]
XeF_4	[13709-61-0] Δ_vH	xenon tetrafluoride (553–663)	60.0	568		[1983HOU]
Y (yttrium)						
$C_{15}H_3F_{18}O_6Y$	[18911-76-7] $\Delta_{\text{sub}}H$	tris(1,1,1,5,5,5-hexafluoro-2,4-pentanedionato)yttrium(III) (310–365)	91.6 ± 8.5		ME	[1999ALI/MAL]
$C_{15}H_{15}Y$	[1294-07-1] $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$	tris(cyclopentadienyl)yttrium	111.7 ± 3.5 99.2 ± 3.3	298		[1982PIL/SKI, 1974DEV/RAB] [1973DEV/BOR]

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)			
C₁₅H₂₁O₆Y	[15554-47-9]	<i>tris</i> (2,4-pentanedionato)yttrium(III)					
	$\Delta_{\text{sub}}H$		98 ± 16				[1984TRE/BER]
C₃₃H₅₇O₆Y	[15632-39-0]	<i>tris</i> (2,2,6,6-tetramethylheptan-3,5-dionato)yttrium(III)					
	$\Delta_{\text{fus}}H$		51.8	440			[2004FUL/RUZ2]
	$\Delta_{\text{sub}}H$	(395–434)	141.6				[2004FUL/RUZ2]
	$\Delta_{\text{sub}}H$	(358–387)	151.0 ± 0.8	372	TE		[2001COL/LAU]
	$\Delta_{\text{sub}}H$	(357–377)	153.1 ± 0.4	366	TE		[2001COL/LAU]
	$\Delta_{\text{sub}}H$	(403–433)	135.9		TG,DTA		[1997YUA/YAN]
	$\Delta_{\text{sub}}H$		117.0				[1997SAN/ROC]
	$\Delta_{\text{sub}}H$	(382–412)	126.0	397	T		[1996RAP/DES]
	$\Delta_{\text{sub}}H$		117.0				[1993TOB/LAN]
	$\Delta_{\text{sub}}H$		115.7				[1993TOB/LAN]
	$\Delta_{\text{sub}}H$		138.5		GS		[1990YUH/KIK]
	$\Delta_{\text{sub}}H$	(363–418)	156.9	388	ME		[1981AMA/SAT]
	$\Delta_{\text{sub}}H$		130.8		ME		[1973BRU/CUR]
	Δ_vH	(450–455)	89.5				[2004FUL/RUZ2]
Δ_vH		66.5		GS		[1990YUH/KIK]	
C₃₂H₄₀F₁₂O₈NaY	[12576-89-5]	sodium <i>tetrakis</i> (1,1,1-trimethyl-5,5,5-trifluoro-2,4-pentanedionato)yttrate					
	$\Delta_{\text{sub}}H$	(418–503)	130 ± 3	460	T		[1993SYO/GOL]
	$\Delta_{\text{sub}}H$	(463–503)	142 ± 12	483			[1993SYO/GOL]
Yb (ytterbium)							
C₁₅H₁₅Yb	[1295-20-1]	<i>tris</i> (cyclopentadienyl)ytterbium					
	$\Delta_{\text{sub}}H$		108.8 ± 3.5	298			[1982PIL/SKI, 1974DEV/RAB]
	$\Delta_{\text{sub}}H$		96.2 ± 2.9				[1973DEV/BOR]
C₁₅H₂₁O₆Yb	[14284-98-1]	<i>tris</i> (2,4-pentanedionato)ytterbium(III)					
	$\Delta_{\text{sub}}H$	(364–404)	93.3				[1981SMI/MAR]
C₃₀H₃₀F₂₁O₆Yb	[18323-96-1]	<i>tris</i> (1,1,1,2,2,3,3-heptafluoro-7,7-dimethyloctane-4,6-dionato)ytterbium(III)					
	$\Delta_{\text{sub}}H$	(339–356)	154.8 ± 3.3			ME	[1971SWA/KAR]
C₃₃H₅₇O₆Yb	[15492-52-1]	<i>tris</i> (2,2,6,6-tetramethylpentane-2,4-dionato)ytterbium(III)					
	$\Delta_{\text{sub}}H$		131.1 ± 2.7	298	DSC		[1999SAN/PET]
	$\Delta_{\text{sub}}H$	(363–413)	156.9	388	ME		[1981AMA/SAT]
	$\Delta_{\text{sub}}H$	(410–444)	133.3	427	BG		[1969SIC/DUB]
	Δ_vH	(444–494)	82.8		BG		[1969SIC/DUB]
Zn (zinc)							
C₂H₆Zn	[544-97-8]	dimethyl zinc					
	$\Delta_{\text{trs}}H$		1.06	210.3			
	$\Delta_{\text{fus}}H$		6.83	230.1			[1996DOM/HEA]
	Δ_vH	(273–313)	30.4 ± 0.1				[1997BAE]
	Δ_vH		29.5 ± 0.4				[1949CAR/HAR, 1982PIL/SKI]
	Δ_vH	(248–318)	29.9	283	BG		[1946BAM/LEV]
C₄H₁₀Zn	[557-20-0]	diethyl zinc					
	$\Delta_{\text{fus}}H$	(18–273)	18.05	239.8			[1987GIB/GRI]
	Δ_vH		37.9	298			[1983HOU2]
	Δ_vH		40.2 ± 2.1				[1949CAR/HAR, 1982PIL/SKI]
	Δ_vH	(250–391)	39.9	265			[1947STU]
	Δ_vH		40.2		BG		[1946BAM/LEV]

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)			
C₄H₁₆Cl₂N₈S₄Zn	[28813-20-9] $\Delta_{\text{sub}}H$	<i>trans</i> -dichloro- <i>tetrakis</i> (thiourea) zinc(II) (351–382)	90 ± 20				[1970ASH]
C₆H₁₄Zn	[628-91-1] Δ_vH Δ_vH Δ_vH Δ_vH	dipropyl zinc (313–370)	42.1 ± 0.4 45.6 ± 2.5 39.5 40.3	341			[1984SOK/BAE2] [1949CAR/HAR, 1982PIL/SK1] [1949HAT/SUT] [1946BAM/LEV]
C₆H₁₄Zn	[625-81-0] Δ_vH Δ_vH	diisopropyl zinc (303–345) (310–338)	41.8 ± 0.5 47.4	324 324			[1984SOK/BAE2] [1946THO]
C₈H₁₈Zn	[1119-90-0] Δ_vH Δ_vH Δ_vH Δ_vH	dibutyl zinc (305–379)	50.7 ± 0.3 54.4 ± 3.3 45.3 42.9	342			[1984SOK/BAE] [1949CAR/HAR, 1982PIL/SK1] [1949HAT/SUT] [1946BAM/LEV]
C₈H₁₈Zn	[7446-94-8] Δ_vH	di- <i>sec</i> -butyl zinc (287–372)	40.9 ± 0.2	330			[1984SOK/BAE]
C₈H₁₈Zn	[na] Δ_vH	diisobutyl zinc (288–372)	44.6 ± 0.2	330			[1984SOK/BAE]
C₈H₁₈Zn	[16636-96-7] $\Delta_{\text{fus}}H$ Δ_vH	di- <i>tert</i> -butyl zinc (300–322)	45.3 49.3 ± 0.8	300 311			[1984SOK/BAE] [1984SOK/BAE]
C₁₀H₁₄O₄Zn	[14024-63-6] $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$	<i>bis</i> (2,4-pentanedionato)zinc(II)	132.6 ± 8 117 ± 3	298		C	[1985MUR/SAK, 1988RIB/PIL] [1980SAC/HIL]
C₁₀H₂₀N₂S₄Zn	[14324-55-1] $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$	<i>bis</i> (diethyldithiocarbamate)zinc(II) (401–444)	115 ± 15 143.1 142.7 ± 2.5	298 422.5		DSC,E A GC	[2000DEA/SOU] [1987STE/MAL] [1976TAV/NEE]
C₁₀H₂₂Zn	[14402-93-8] Δ_vH	dipentyl zinc	48.6				[1949HAT/SUT]
C₁₂H₂₆Zn	[13822-55-4] Δ_vH	dihexyl zinc	56.2				[1949HAT/SUT]
C₁₄H₂₈N₂S₄Zn	[15694-56-1] $\Delta_{\text{sub}}H$	<i>bis</i> (dipropyldithiocarbamate)zinc(II)	147 ± 2	298		DSC, E	[1992DEC/AIR]
C₁₄H₃₀Zn	[14402-95-0] Δ_vH	diheptyl zinc	62.3				[1949HAT/SUT]
C₁₈H₁₂N₂O₂Zn	[13978-85-3] $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$ $\Delta_{\text{sub}}H$	<i>bis</i> (8-hydroxyquinolino)zinc(II) (473–513)	183.2 ± 6.3 167.9 ± 6 178 ± 6	298 493 298		ME ME	[1994RIB/MAT] [1984BUR/MOR] [1984BUR/MOR]
C₁₈H₃₆N₂S₄Zn	[136-23-2] $\Delta_{\text{sub}}H$	<i>bis</i> (dibutyldithiocarbamate)zinc(II)	107 ± 3	298		DSC,E	[1991DES/DES]
C₁₈H₃₆N₂S₄Zn	[36190-62-2] $\Delta_{\text{sub}}H$	<i>bis</i> (diisobutyldithiocarbamate)zinc(II)	283 ± 2	298		DSC,E	[1994SOU/PIN]

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound		T_m (K)	Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)			
C₂₀H₁₆N₂O₂Zn	[14128-73-5]	<i>bis</i> (8-hydroxy-2-methylquinolate)zinc(II)				
	$\Delta_{\text{sub}}H$	(437–556)	172.0 ± 5.0	541	ME	[1998RIB/MAT3]
	$\Delta_{\text{sub}}H$		179.4 ± 5.0	298		[1998RIB/MAT3]
C₂₂H₃₈O₄Zn	[14363-14-5]	<i>bis</i> (2,2,6,6-tetramethylheptan-3,5-dianato)zinc(II)				
	$\Delta_{\text{sub}}H$		136		ME	[1973BRU/CUR]
C₂₂H₄₄N₂S₄Zn	[15337-18-5]	<i>bis</i> (dipentylidithiocarbamate)zinc(II)				
	$\Delta_{\text{sub}}H$		127 ± 3	298	DSC,E	[2000DEA/SOU]
C₃₂F₁₆N₈Zn	[14320-04-8]	1,2,3,4,8,9,10,11,15,16,17,18,22,23,24,25-hexadecafluorophthalocyanine zinc(II)				
	Δ_vH	(658–711)	56.5 ± 0.4	685	ME	[2008SEM/BAS]
C₃₂H₁₆N₈Zn	[14320-04-8]	zinc(II) phthalocyanine				
	Δ_vH	(664–709)	48.1 ± 1.8	686	ME	[2008SEM/BAS]
C₄₄H₂₈N₄Zn	[14074-80-7]	5,10,15,20-tetraphenylphosphine zinc (II)				
	$\Delta_{\text{sub}}H$	(555–567)	183 ± 3	559	ME	[2002PAT/CAM]
	$\Delta_{\text{sub}}H$	(555–567)	196 ± 3	298	ME	[2002PAT/CAM]
	$\Delta_{\text{sub}}H$	(563–663)	213 ± 3			[1994PER/NAN, 2002PAT/CAM]
	$\Delta_{\text{sub}}H$		109	666	UV/Vis	[1971EDW/DOL, 2002PAT/CAM]
	$\Delta_{\text{sub}}H$		208 ± 4		GS	[2000PER/GOL]
Cl₂Zn	[7646-85-7]	zinc chloride				
	Δ_vH	(695–826)	134.5	760		[1958BLO/WEL]
F₂Zn	[7783-49-5]	zinc fluoride				
	$\Delta_{\text{sub}}H$	(901–1125)	252.4	1015	ME	[1973BIE/EIC]
Zr (zirconium)						
C₁₀H₁₀Cl₂Zr	[1291-32-3]	<i>bis</i> (cyclopentadienyl)zirconium dichloride				
	$\Delta_{\text{sub}}H$		108.5 ± 4.6	298	ME	[2001DIO/PIE]
	$\Delta_{\text{sub}}H$	(393–457)	100.3	425	A	[1987STE/MAL]
	$\Delta_{\text{sub}}H$		105.0 ± 2.1	298		[1982PIL/SKI, 1976KIR/TEL]
	$\Delta_{\text{sub}}H$		100.4 ± 1.7			[1977BAL/BAR]
	$\Delta_{\text{sub}}H$		96.7			[1969DIL/KIS]
	$\Delta_{\text{sub}}H$		103 ± 13	298		[1968KIS/DIL, 2001DIO/PIE]
	Δ_vH	(394–425)	71 ± 5			[2008ARU/MAT]
C₁₂H₁₆Zr	[1291-32-3]	<i>bis</i> (cyclopentadienyl)dimethylzirconium				
	$\Delta_{\text{sub}}H$		81.2 ± 2.1	298		[1982PIL/SKI, 1976KIR/TEL]
C₁₂H₃₆N₄Zr	[175923-04-3]	<i>tetrakis</i> (methylethylamino)zirconium (IV)				
	$\Delta_{\text{sub}}H$	(278–333)	79.4 ± 2.4		ME	[2009MON/NUT]
Note: Authors state in the paper that the compound is a liquid at room temperature. Figure 3 in the paper shows that the plot of ln P versus 1/T is linear over the entire temperature range. The authors refer to the enthalpy as the enthalpy of sublimation. We have taken the value to be the enthalpy of vaporization given the authors' statement that the compound is a liquid. The compound's melting point temperature is unknown.						
C₁₆H₃₆O₄Zr	[na]	tetra- <i>tert</i> -butoxy zirconium				
	Δ_vH	(374–587)	56.6	389	A	[1987STE/MAL]
C₂₀H₄F₂₄O₈Zr	[19530-02-0]	<i>tetrakis</i> (1,1,1,5,5,5-hexafluoro-2,4-pentanedionato)zirconium(IV)				
	$\Delta_{\text{sub}}H$	(333–363)	59.0		TGA	[2000FAH/BAR]
	Δ_vH	(366–456)	48.6 ± 0.6	411	T	[1996MOR/SYS]
C₂₀H₁₆F₁₂O₈Zr	[17499-68-2]	<i>tetrakis</i> (1,1,1-trifluoro-2,4-pentanedionato)zirconium(IV)				
	$\Delta_{\text{sub}}H$	(373–403)	94		TGA	[2000FAH/BAR]

TABLE 12. Phase change enthalpies of organometallic and inorganic compounds—Continued

Molecular Formula	CAS Reg No Enthalpy	Compound				Method	Reference
		Temp (K) Range	$\Delta_{\text{trans}}H_m$ (kJ mol ⁻¹)	T_m (K)			
	$\Delta_{\text{sub}}H$	(368–398)	133.6 ± 2.0	383		SMZG	[1996MOR/SYS]
	$\Delta_{\text{sub}}H$		118.7 ± 3.1	298		C	[1992RIB/FER2]
	$\Delta_{\text{sub}}H$	(383–438)	126.4 ± 1.7			GS	[1985MAT/KUW]
	$\Delta_{\text{sub}}H$	(383–438)	119.2 ± 1.7			GS	[1985MAT/KUW]
C₂₀H₂₈O₈Zr	[17501-44-9]	<i>tetrakis</i> (2,4-pentanedionato)zirconium(IV)					
	$\Delta_{\text{sub}}H$	(413–443)	126			TGA	[2000FAH/BAR]
	$\Delta_{\text{sub}}H$	(403–433)	138.8 ± 2	418		SMZG	[1996MOR/SYS]
	$\Delta_{\text{sub}}H$		125.8 ± 2.9	298		C	[1992RIB/FER2]
	$\Delta_{\text{sub}}H$		132.0 ± 6.8	463			[1987MUR/HIL2]
	$\Delta_{\text{sub}}H$		116 ± 34				[1984TRE/BER]
C₂₀H₄₄O₄Zr	[na]	<i>tetrakis</i> (1,1-dimethylpropoxy)zirconium					
	Δ_vH	(392–426)	68	407		A	[1987STE/MAL]
C₂₀H₄₄O₄Zr	[na]	tetra- <i>tert</i> -pentoxyzirconium					
	Δ_vH	(361–435)	74.1	361		A	[1987STE/MAL]
C₂₂H₂₀Zr	[51177-89-0]	<i>bis</i> (cyclopentadienyl)diphenylzirconium					
	$\Delta_{\text{sub}}H$		92.0 ± 4.2				[1976KIR/TEL]
C₂₄H₅₂O₄Zr	[na]	<i>tetrakis</i> (1,1-dimethylbutoxy)zirconium					
	Δ_vH	(406–449)	93.3	421		A	[1987STE/MAL]
C₂₄H₅₂O₄Zr	[na]	<i>tetrakis</i> (1-methyl-1-ethylpropoxy)zirconium					
	Δ_vH	(423–460)	91.4	438		A	[1987STE/MAL]
C₃₂H₄₀F₁₂O₈Zr	[56044-44-1]	<i>tetrakis</i> (1,1,1-trimethyl-5,5,5-trifluoro-2,4-pentanedionato) zirconium(IV)					
	$\Delta_{\text{sub}}H$	(408–449)	87 ± 1				[2008ARU/MAT]
	$\Delta_{\text{sub}}H$	(388–423)	134.9 ± 1.6	406		SMZG	[1996MOR/SYS]
C₄₄H₇₆O₈Zr	[18865-74-2]	<i>tetrakis</i> (2,2,6,6-tetramethylheptan-3,5-dionato)zirconium(IV)					
	$\Delta_{\text{us}}H$		11.6	446			
		Note: value also includes the enthalpy of the solid/solid transition at 438 K					
	$\Delta_{\text{fus}}H$		5.7	616		DSC	[2008ZHE/MOR]
	$\Delta_{\text{us}}H$		4.86	387.1			
	$\Delta_{\text{us}}H$		16.38	430.3			[2004FUL/RUZ2]
	$\Delta_{\text{sub}}H$	(411–463)	85.4	437		GS	[2008JEE/ARO]
	$\Delta_{\text{sub}}H$	(413–443)	120			TGA	[2000FAH/BAR]
ZrCl₄	[10026-11-6]	zirconium tetrachloride					
	$\Delta_{\text{sub}}H$	(405–518)	98.9 ± 0.5	512		T	[1994TAN/BOS]
ZrF₄	[7783-64-4]	zirconium tetrafluoride					
	$\Delta_{\text{sub}}H$	(696–856)	240.0 ± 0.1	298		TE	[1994KON/HIL]
	$\Delta_{\text{sub}}H$	796	243	298		MS	[1965SID/AKI, 1994KON/HIL]
	$\Delta_{\text{sub}}H$	(983–1177)	241.1 ± 0.1	298			[1964FIS/PET, 1994KON/HIL]
	$\Delta_{\text{sub}}H$	(681–913)	242.6 ± 1.7	298		MS	[1963AKA/BEL, 1994KON/HIL]
	$\Delta_{\text{sub}}H$	(713–873)	232.3 ± 1.2	298			[1963GAL/TUM, 1994KON/HIL]
	$\Delta_{\text{sub}}H$	(983–1081)	239.9 ± 0.2	298			[1958CAN/NEW, 1994KON/HIL]
	$\Delta_{\text{sub}}H$	(890–1150)	241.8 ± 0.6	298		GS	[1954SEN/SNY2, 1994KON/HIL]

5. References

- 1882KRA F. Krafft, *Berichte der Deutschen Chemischen Gesellschaft* **15**, 1687 (1882).
- 1883KAH G. W. A. Kahlbaum, *Berichte der Deutschen Chemischen Gesellschaft* **16**, 2476 (1883).
- 1886RIC A. Richardson, *J. Chem. Soc.* 761 (1986).
- 1889EYK J. F. Eykman, *Z. Phys. Chem., Stoechiom. Ver-*
wandtschaftsl. **4**, 497 (1889).
- 1894KAH G. W. A. Kahlbaum, *Z. Phys. Chem., Stoechiom. Ver-*
wandtschaftsl. **13**, 14 (1894).
- 1898KAH G. W. A. Kahlbaum, *Z. Phys. Chem., Stoechiom. Ver-*
wandtschaftsl. **26**, 577 (1898).
- 1898LOU W. Louguinine, *Ann. Chim. (Paris)* **7**, 334 (1898).
- 1901DEF M. De Forcrand, *Compt. Rend.* **132**, 688 (1901).
- 1903DEW/JON J. Dewar and H. O. Jones, *Proc. R. Soc. London* **71A**, 427

- (1903).
- 1904JQAQ/WAS A. Jaquero and E. Wassmer, *Berichte der Deutschen Chemischen Gesellschaft* **3**, 2531 (1904).
- 1906REX A. Rex, *Z. Phys. Chem., Stoechiom. Verwandtschaftsl.* **55**, 355 (1906).
- 1910BIN E. C. Bingham, *J. Am. Chem. Soc.* **43**, 287 (1910).
- 1910MEY J. Meyer, *Z. Phys. Chem., Stoechiom. Verwandtschaftsl.* **72**, 225 (1910).
- 1911LOU/DUP W. Louguine and G. Dupont, *Bull. Soc. Chim. Fr.* **9**, 219 (1911); *Chem. Abstr.* **5**, 10166 (1911).
- 1912HER/RAT W. Herz and W. Rathmann, *Chem. Ztg.* **36**, 1417 (1912).
- 1913MUN C. F. Mündel, *Z. Phys. Chem., Stoechiom. Verwandtschaftsl.* **85**, 435 (1913).
- 1914STO/WIL A. Stock and E. Willfroth, *Berichte der Deutschen Chemischen Gesellschaft* **47**, 144 (1914).
- 1915CRA J. M. Crafts, *J. Chim. Phys. Phys.-Chim. Biol.* **13**, 105 (1915).
- 1916TER J. W. Terwen, *Z. Phys. Chem., Stoechiom. Verwandtschaftsl.* **91**, 469 (1916).
- 1919BEL/SAW J. M. Bell and J. P. Sawyer, *J. Ind. Eng. Chem.* **11**, 1025 (19).
- 1919MOL/BAT E. Moles and T. J. Batuecas, *J. Chim. Phys. Phys.-Chim. Biol.* **17**, 537 (1919).
- 1920BAX/BEZ G. P. Baxter, F. K. Bezenberger, and C. H. Wilson, *J. Am. Chem. Soc.* **42**, 1386 (20).
- 1920MON K. P. Monroe, *J. Ind. Eng. Chem.* **12**, 969 (1920).
- 1921MAA/WRI O. Maass and C. H. Wright, *J. Am. Chem. Soc.* **43**, 1098 (1921).
- 1921STO/HEN A. Stock, F. Henning, and E. Kuss, *Berichte der Deutschen Chemischen Gesellschaft* **54**, 1119 (1921).
- 1922NEL/SEN O. A. Nelson and C. E. Senseman, *Ind. Eng. Chem.* **14**, 58 (1922).
- 1922TAY/SMI R. S. Taylor and L. B. Smith, *J. Am. Chem. Soc.* **44**, 2450 (1922).
- 1923MOR/MUR F. S. Mortimer and R. V. Murphy, *Ind. Eng. Chem.* **15**, 1140 (1923).
- 1923SEN/NEL C. E. Senseman and O. A. Nelson, *Ind. Eng. Chem.* **15**, 382 (1923).
- 1924MAA/HIE O. Maas and P. G. Hiebert, *J. Am. Chem. Soc.* **46**, 2693 (1924).
- 1924SIM J. Simons, *J. Am. Chem. Soc.* **46**, 2179 (1924).
- 1925PER/BAR J. H. Perry and D. C. Bardwell, *J. Am. Chem. Soc.* **47**, 2629 (1925).
- 1925SWA/MAC T. H. Swan and H. Mack, Jr., *J. Am. Chem. Soc.* **47**, 2112 (1925).
- 1925VOL/KIR M. Volmer and P. Kirchhoff, *Z. Phys. Chem., Stoechiom. Verwandtschaftsl.* **115**, 223 (1925).
- 1926AND M. R. Andrews, *J. Phys. Chem.* **30**, 1497 (1926).
- 1926AND/LYN D. H. Andrews, G. Lynn, and J. Johnston, *J. Am. Chem. Soc.* **48**, 1274 (1926).
- 1926EVA/AYL W. V. Evans and M. B. Aylesworth, *Ind. Eng. Chem.* **18**, 24 (1926).
- 1926FEL/DUR W. A. Felsing and S. A. Durban, *J. Am. Chem. Soc.* **48**, 2885 (1926).
- 1926LOO/WAL A. G. Loomis and J. E. Walters, *J. Am. Chem. Soc.* **48**, 2051 (1926).
- 1926MAT J. H. Mathews, *J. Am. Chem. Soc.* **48**, 562 (1926).
- 1926NOY/WEB W. A. Noyles, Jr. and D. E. Webbe, *J. Am. Chem. Soc.* **48**, 1882 (1926).
- 1926PER/POR J. H. Perry and F. Porter, *J. Am. Chem. Soc.* **48**, 299 (1926).
- 1926SIN/HAR H. Sinozaki, R. S. Hara, and S. Mitsukuri, *Bull. Chem. Soc. Jpn.* **1**, 59 (1926).
- 1926STR F. Straus, *Berichte der Deutschen Chemischen Gesellschaft* **59**, 1664 (1926).
- 1926TAY/RIN C. A. Taylor and W. H. Rinkenbach, *J. Chem. Soc.* 1305 (1926).
- 1926YAT G. W. C. Yates, *Philos. Mag.* **2**, 817 (1926).
- 1927BER/MAY J. F. T. Berliner and O. E. May, *J. Am. Chem. Soc.* **49**, 1007 (1927).
- 1927COO/COO A. S. Coolidge and M. S. Coolidge, *J. Am. Chem. Soc.* **49**, 100 (1927).
- 1927KLO/WOO S. Klosky, L. P. L. Woo, and R. J. Flanigan, *J. Am. Chem. Soc.* **49**, 1280 (1927).
- 1927MAY/BER O. E. May, J. F. T. Berliner, and D. F. J. Lynch, *J. Am. Chem. Soc.* **49**, 1012 (1927).
- 1927NAG N. N. Nagornov, *Ann. Inst. Anal. Phys. Chem. (Leningrad)* **3**, 562 (1927); *Chem. Zentr.* **II**, 2668 (1927); *Chem. Abstr.* **22**, 4298 (1928).
- 1927PER J. H. Perry, *J. Phys. Chem.* **31**, 1737 (1927).
- 1927RIN W. H. Rinkenbach, *Ind. Eng. Chem.* **19**, 474 (1927).
- 1928GIA/WIE W. F. Giauque and R. Wiebe, *J. Am. Chem. Soc.* **50**, 2193 (1928).
- 1928GIA/WIE2 W. F. Giauque and R. Wiebe, *J. Am. Chem. Soc.* **50**, 101 (1928).
- 1928NEL O. A. Nelson, *Ind. Eng. Chem.* **20**, 1382 (1928).
- 1928NEL2 O. A. Nelson, *Ind. Eng. Chem.* **20**, 1380 (1928).
- 1928PAR/BAR G. S. Parks and B. J. Barton, *J. Am. Chem. Soc.* **50**, 24 (1928).
- 1928ROL M. Roland, *Bull. Soc. Chim. Belg.* **37**, 117 (1928).
- 1928STE/JOH L. E. Steiner and J. J. Johnston, *J. Phys. Chem.* **32**, 912 (1928).
- 1929GIA/WIE W. F. Giauque and R. Wiebe, *J. Am. Chem. Soc.* **51**, 1441 (1929).
- 1929HAG/WEI C. F. Haggerty and J. F. Weiler, *J. Am. Chem. Soc.* **51**, 1623 (1929).
- 1929JEL/RUD K. Jellinek and A. Rudat, *Z. Phys. Chem., Stoechiom. Verwandtschaftsl.* **143**, 55 (1929).
- 1929JOH/GIA H. L. Johnston and W. F. Giauque, *J. Am. Chem. Soc.* **51**, 3194 (1929).
- 1929PEA/PET J. N. Pearce and P. E. Peters, *J. Phys. Chem.* **33**, 873 (1929).
- 1929SMY/ENG C. P. Smyth and E. W. Engel, *J. Am. Chem. Soc.* **51**, 2646 (1929).
- 1929TAN/NAG Y. Tanaka and Y. Nagai, *Proc. Imp. Acad. (Tokyo)* **5**, 78 (1929); *Chem. Abstr.* **23**, 2622 (1929).
- 1930BLA/VAN H. K. Black, G. van Praagh, and B. Topley, *Trans. Faraday Soc.* **26**, 196 (1930).
- 1930BUL/HAU R. H. Bullard and A. C. Haussman, *J. Phys. Chem.* **34**, 743 (1930).
- 1930COO A. S. Coolidge, *J. Am. Chem. Soc.* **52**, 1874 (1930).
- 1930CUN G. V. Cunningham, *Power* **72**, 374 (1930), as cited in Ref. [1985KAI/HAD].
- 1930DRU/FLA C. Drucker and Th. Flade, *Z. wiss. Phot., Schaum Festschr.* **29**, 29 (1930).
- 1930LIV/HEI R. Livingston and J. B. Heisig, *J. Am. Chem. Soc.* **52**, 2409 (1930).
- 1930NEL O. A. Nelson, *Ind. Eng. Chem.* **22**, 971 (1930).
- 1930PAT/PAP W. I. Patnode and J. Papish, *J. Phys. Chem.* **34**, 1494 (1930).
- 1930SCH/THO H. A. Schuette and R. W. Thomas, *J. Am. Chem. Soc.* **52**, 2028 (1930).
- 1930SIM J. H. Simons, *J. Am. Chem. Soc.* **52**, 3488 (1930).
- 1930SOU/AND J. C. Southard and D. H. Andrews, *J. Franklin Inst.* **209**, 349 (1930).
- 1930ZMA M. A. Zmaczynski, *J. Chim. Phys. Phys.-Chim. Biol.* **27**, 503 (1930).
- 1931BUF/FLE R. M. Buffington and J. Fleischer, *J. Ind. Eng. Chem.* **23**, 1290 (1931).
- 1931COS V. E. Cosslett, *Z. Anorg. Allg. Chem.* **201**, 75 (1931); *Chem. Abstr.* **26**, 1535 (1932).
- 1931CRO/BIJ C. A. Crommelin, W. J. Bijleveld, and E. G. Brown, *Proc. R. Acad. Sci. Amsterdam* **34**, 1314 (1931); *Chem. Abstr.* **26**, 2904 (1931).
- 1931HIL A. E. Hill, *J. Am. Chem. Soc.* **53**, 2598 (1931).
- 1931LIN E. G. Lindner, *J. Phys. Chem.* **35**, 531 (1931).
- 1931RUF/MEN O. Ruff and W. Menzel, *Z. Anorg. Allg. Chem.* **202**, 49 (1931).
- 1931SCH/COW H. A. Schuette and M. A. Cowley, *J. Am. Chem. Soc.* **53**, 3485 (1931).
- 1931VAN/MAN P. A. van der Meulen and R. F. Mann, *J. Am. Chem. Soc.* **53**, 451 (1931).
- 1931WIB/SUT E. Wiberg and W. Sütterlin, *Z. Anorg. Allg. Chem.* **202**, 1 (1931).

- 1932CLA/GIA J. O. Clayton and W. F. Giauque, *J. Am. Chem. Soc.* **54**, 2610 (1932).
- 1932ELL/REI L. M. Ellis, Jr. and E. E. Reid, *J. Am. Chem. Soc.* **54**, 1674 (1932).
- 1932JON/DYK W. J. Jones, W. J. C. Dyke, G. Davies, D. C. Griffiths, and J. H. Webb, *J. Chem. Soc.* 2285 (1932).
- 1932KLE/HEN W. Klemm and P. Henkel, *Z. Anorg. Allg. Chem.* **207**, 73 (1932).
- 1932KUB T. Kubota, *J. Chem. Soc. Jpn.* **53**, 404 (1932).
- 1932NEU/VOL K. Neumann and E. Volker, *Z. Phys. Chem. Abt. A* **161**, 33 (1932).
- 1932SPA/THO M. E. Spaght, S. B. Thomas, and G. S. Parks, *J. Phys. Chem.* **36**, 882 (1932).
- 1932VAU W. E. Vaughan, *J. Am. Chem. Soc.* **54**, 3863 (1932).
- 1933BAR/BUR E. G. V. Barrett and J. L. Burrage, *J. Phys. Chem.* **37**, 1029 (1933).
- 1933BOO/BUR H. S. Booth, P. E. Burchfield, E. M. Bixly, and J. B. MacKelvey, *J. Am. Chem. Soc.* **53**, 2231 (1933).
- 1933BRO/FRA L. F. Broadway and R. G. J. Fraser, *J. Chem. Soc.* 429 (1933).
- 1933COW/SCH M. A. Cowley and H. A. Schuette, *J. Am. Chem. Soc.* **55**, 387 (1933).
- 1933DEI V. R. Deitz, *J. Am. Chem. Soc.* **55**, 472 (1933).
- 1933GIA/CLA W. F. Giauque and J. O. Clayton, *J. Am. Chem. Soc.* **55**, 4875 (1933).
- 1933HEI G. Heim, *Bull. Soc. Chim. Belg.* **42**, 467 (1933).
- 1933HEI/MUR G. B. Heisig and O. D. Murd, *J. Am. Chem. Soc.* **55**, 3485 (1933).
- 1933HEI2 G. B. Heisig, *J. Am. Chem. Soc.* **55**, 2304 (1933).
- 1933HEN/MUR H. R. Henze and T. J. Murchison, *J. Am. Chem. Soc.* **55**, 4255 (1933).
- 1933HOR/GEI F. F. Horvorka E. Geiger, *J. Am. Chem. Soc.* **55**, 4759 (1933).
- 1933HOR/LAN F. Hovorka, H. P. Lankelma, and C. K. Naujoks, *J. Am. Chem. Soc.* **53**, 4820 (1933).
- 1933KRA/TOO C. A. Kraus and F. E. Toonder, *Proc. Natl. Acad. Sci. U.S.A.* **19**, 292 (1933); *Chem. Abstr.* **27**, 2646 (1933).
- 1933MEN/MOH W. Menzel and F. Mohry, *Z. Anorg. Allg. Chem.* **210**, 257 (1933).
- 1933MIL/MEN F. T. Miles and A. W. C. Menzies, *J. Phys. Chem.* **37**, 435 (1933).
- 1933NEL/YOU O. A. Nelson and H. D. Young, *J. Am. Chem. Soc.* **55**, 2429 (1933).
- 1933PAR/HUF G. S. Parks, H. M. Huffman, and M. Barmore, *J. Am. Chem. Soc.* **55**, 2733 (1933).
- 1933TAN H. Tanneberger, *Chem. Ber.* **66B**, 484 (1933); *Chem. Abstr.* **27**, 3374 (1933).
- 1933TAY/LAY A. Taylor and E. T. Layng, *J. Chem. Phys.* **1**, 798 (1933).
- 1933TOR/MOL M. T. Toral and E. Moles, *An. R. Soc. Esp. Fis. Quim.* **31**, 735 (1933); *Chem. Abstr.* **28**, 954 (1934).
- 1933WHI/FLE F. C. Whitmore and G. H. Fleming, *J. Am. Chem. Soc.* **55**, 3803 (1933).
- 1933YOS/STO D. M. Yost and W. E. Stone, *J. Am. Chem. Soc.* **55**, 1889 (1933).
- 1934CAM/CAM A. N. Campbell and A. J. Campbell, *Trans. Faraday Soc.* **30**, 1109 (1934).
- 1934GOO J. W. Goodeve, *Trans. Faraday Soc.* **30**, 501 (1934). (Vapor pressure data are presented graphically in paper as log P versus 1/T.)
- 1934GOR S. V. Gorbachev, *Educ. Psychol. Meas.* **7**, 388 (1934); *Chem. Abstr.* **29**, 13996 (1934).
- 1934HIE/WOE W. Hieber and A. Woerner, *Z. Elektrochem. Angew. Phys. Chem.* **40**, 252 (1934).
- 1934HIR H. Hirsbrunner, *Helv. Chim. Acta* **17**, 477 (1934).
- 1934LEW/SCH G. N. Lewis and P. W. Schultz, *J. Am. Chem. Soc.* **56**, 1002 (1934).
- 1934MEH W. Mehl, *Z. Phys. Chem. Abt. A* **164**, 312 (1934).
- 1934OGU/ANJ S. Oguri, S. Anjo, and Y. Kuwabara, *Bull. Waseda Appl. Chem. Soc.* **22**, 1 (1934); *Chem. Abstr.* **28**, 4298 (1934).
- 1934RIG/FEL N. E. Rigler, W. A. Felsing, and H. R. Henze, *J. Am. Chem. Soc.* **56**, 1499 (1934).
- 1934ROT/NAG L. Rotinjanz and N. Nagornow, *Z. Phys. Chem. Abt. A* **169**, 20 (1934).
- 1934SCH/BIC W. C. Schumb and F. A. Bickford, *J. Am. Chem. Soc.* **56**, 852 (1934).
- 1934WIN/BLA M. M. Windsor and A. A. Blanchard, *J. Am. Chem. Soc.* **56**, 823 (1934).
- 1934WOL/TRI K. L. Wolf and H. G. Trieschmann, *Z. Phys. Chem. Abt. B* **27**, 376 (1934).
- 1935BAU/BUR H. Bauer and K. Burschkies, *Berichte der Deutschen Chemischen Gesellschaft* **68**, 1238 (1935).
- 1935BLU/GIA R. W. Blue and W. F. Giauque, *J. Am. Chem. Soc.* **57**, 991 (1935).
- 1935BUT/RAM J. A. V. Butler, C. N. Ramchandani, and D. W. Thompson, *J. Chem. Soc.* 280 (1935).
- 1935CAR/DIC J. L. Carrico and R. G. Dickinson, *J. Am. Chem. Soc.* **57**, 1343 (1935).
- 1935HIE/ROM W. Hieber and E. Romberg, *Z. Anorg. Allg. Chem.* **221**, 332 (1935); *Chem. Abstr.* **29**, 2467 (1935).
- 1935JON/EVA W. J. Jones, D. P. Evans, T. Gulwell, and D. C. Griffiths, *J. Chem. Soc.* 39 (1935).
- 1935KIR/POP V. A. Kireev and A. A. Popov, *Russ. J. Gen. Chem.* **5**, 1399 (1935); *Chem. Abstr.* **30**, 2441 (1936).
- 1935LEE P. Z. Lee, *Z. Anorg. Allg. Chem.* **223**, 213 (1935); *Chem. Abstr.* **29**, 6118 (1935).
- 1935SCH/STA H. Schierholtz and M. L. Staples, *J. Am. Chem. Soc.* **57**, 2709 (1935).
- 1935SPE/WIL R. Spence and W. Wild, *J. Chem. Soc.* 506 (1935).
- 1935THO/LIN H. W. Thompson and J. W. Linnett, *Trans. Faraday Soc.* **31**, 1743 (1935).
- 1935TRI H. G. Trieschmann, Ph.D. dissertation, Inst. Fur Phys. Chem. and Electrochem. der Universitat Kiel, Germany, 1935.
- 1935WIL A. L. Wilson, *Ind. Eng. Chem.* **27**, 867 (1935).
- 1936AST/MES J. G. Aston and G. H. Messerly, *J. Am. Chem. Soc.* **58**, 2354 (1936).
- 1936BEN/CUT H. E. Bent, G. R. Cuthbertson, M. Dorfman, and R. E. Leary, *J. Am. Chem. Soc.* **58**, 165 (1936).
- 1936BUC/NOR E. J. Buckler and R. G. W. Norrish, *J. Chem. Soc.* 1567 (1936).
- 1936CUT/BEN G. R. Cuthbertson and H. E. Bent, *J. Am. Chem. Soc.* **58**, 2000 (1936).
- 1936DEB J. H. De Boer, *Trans. Faraday Soc.* **32**, 10 (1936).
- 1936EWE M. Ewert, *Bull. Soc. Chim. Belg.* **45**, 493 (1936).
- 1936GIA/BLU W. F. Giauque and R. W. Blue, *J. Am. Chem. Soc.* **58**, 831 (1936).
- 1936GRE M. A. Grekhnev, *Lesokhim. Prom.* **5**, 11 (1936); *Chem. Abstr.* **32**, 5271 (1938).
- 1936HOV/SCH F. Hovorka, R. A. Schaefer, and D. Dreisback, *J. Am. Chem. Soc.* **58**, 2264 (1936).
- 1936KEA K. Kearby, *J. Am. Chem. Soc.* **58**, 374 (1936).
- 1936KIN/GAR A. M. King and W. E. Garner, *J. Chem. Soc.* 1372 (1936).
- 1936MER/SCH H. Merten and H. Schlüter, *Chem. Ber.* **69B**, 1364 (1936).
- 1936NEG G. R. Negishi, *J. Am. Chem. Soc.* **58**, 2293 (1936).
- 1936THO/LIN H. W. Thompson and J. W. Linnett, *Trans. Faraday Soc.* **32**, 681 (1936).
- 1937AST/SIL J. G. Aston, C. W. Siller, and G. H. Messerly, *J. Am. Chem. Soc.* **59**, 1743 (1937).
- 1937BUR/SCH A. B. Burg and H. I. Schlesinger, *J. Am. Chem. Soc.* **59**, 780 (1937).
- 1937DEW J. H. de Wilde, *Z. Anorg. Allg. Chem.* **233**, 411 (1937).
- 1937DUN/WOL H. Dunken and K. L. Wolf, *Z. Phys. Chem. Abt. B* **38**, 441 (1937).
- 1937EGA/KEM C. J. Egan and J. D. Kemp, *J. Am. Chem. Soc.* **59**, 1264 (1937).
- 1937GAL/HIB A. F. Gallagher and H. Hibbert, *J. Am. Chem. Soc.* **59**, 2521 (1937).
- 1937GAR/BRE G. S. Gardner and J. E. Brewer, *Ind. Eng. Chem.* **29**, 179 (1937).
- 1937GIA/EGA W. F. Giauque and C. J. Egan, *J. Chem. Phys.* **5**, 45 (1937).
- 1937GRO/IPA A. V. Grosse and V. N. Ipatieff, *J. Org. Chem.* **2**, 447 (1937).
- 1937GUY/SCH A. Guyer, W. Schutze, and M. Weidenmann, *Helv. Chim.*

- Acta **20**, 1936 (1937).
- 1937KEM/GIA J. D. Kemp and W. F. Giauque, *J. Am. Chem. Soc.* **59**, 79 (1937).
- 1937KLE/WAG A. Klemenc and G. Wagner, *Berichte der Deutschen Chemischen Gesellschaft* **70B**, 1880 (1937); *Chem. Abstr.* **31**, 8412 (1937).
- 1937LIN/ROH R. Linke and W. Rohrmann, *Z. Phys. Chem. Abt. B* **35**, 256 (1937).
- 1937MOO/KAN V. G. Moor, E. K. Kanep, and I. E. Dobkin, *Trans. Exptl. Research Lab. Khemgas, Materials on Cracking and Chemical Treatment of Cracking Products U.S.S.R.* **3**, 320 (1937); *Chem. Abstr.* **31**, 6072 (1937).
- 1937OVE/GIA R. Overstreet and W. F. Giauque, *J. Am. Chem. Soc.* **59**, 254 (1937).
- 1937PUR/ZAH R. H. Purcell and F. D. Zahoorbux, *J. Chem. Soc.* 1029 (1937).
- 1937RIN/SAY J. C. Rintelen Jr., J. H. Saylor, and P. M. Gross, *J. Am. Chem. Soc.* **59**, 1129 (1937).
- 1937STE/GIA C. C. Stephenson and W. F. Giauque, *J. Chem. Phys.* **5**, 149 (1937).
- 1937THO/DAI H. W. Thompson and F. S. Dainton, *Trans. Faraday Soc.* **33**, 1546 (1937).
- 1937WIT/KEM R. K. Witt and J. D. Kemp, *J. Am. Chem. Soc.* **59**, 273 (1937).
- 1938BRA J. D. Brandner, *Ind. Eng. Chem.* **30**, 681 (1938).
- 1938EFT E. Efring, Ph.D. dissertation, University of Lund, 1938, as quoted in Ref. [1956KIR].
- 1938EGA/KEM C. G. Egan and J. D. Kemp, *J. Am. Chem. Soc.* **60**, 2097 (1938).
- 1938GIA/STE W. F. Giauque and C. C. Stephenson, *J. Am. Chem. Soc.* **60**, 1389 (1938).
- 1938HOV/LAN F. Hovorka, H. P. Lankelma, and S. C. Stanford, *J. Am. Chem. Soc.* **60**, 820 (1938).
- 1938KUC K. Kuchinskaya, *Sbornik Trudov. Opytnogo Zavoda im. Akad. S. V. Lebedeva*, 27 (1938); *Chem. Abstr.* **34**, 3147 (1940).
- 1938LEV/SHT E. Levin and I. Shtern, *Educ. Psychol. Meas.* **11**, 426 (1938).
- 1938LIN M. Linhard, *Z. Anorg. Allg. Chem.* **236**, 200 (1938).
- 1938RAD/ALE D. Radulescu and M. Alexa, *Bull. Chem. Soc. Romania* **20A**, 89 (1938).
- 1938REI L. Reidel, *Z. Ges. Kalte-Ind.* **45**, 221 (1938).
- 1938SCT/RAY G. Scatchard and C. L. Raymond, *J. Am. Chem. Soc.* **60**, 1278 (1938); **60**, 3099(E) (1938).
- 1938UBB A. R. Ubbelohde, *Trans. Faraday Soc.* **34**, 282 (1938).
- 1938WEG H. Weghofer, Ph.D. dissertation, Phys. Chem. Institut der Universitat Halle, 1938.
- 1938WOL/WEG K. L. Wolf and H. Z. Weghofer, *Z. Phys. Chem. Abt. B* **39**, 194 (1938).
- 1939AST/EID J. G. Aston, M. L. Eidinoff, and W. S. Forster, *J. Am. Chem. Soc.* **61**, 1539 (1939).
- 1939FRA/CLU A. Frank and K. Clusius, *Z. Phys. Chem. Abt. B* **42**, 395 (1939).
- 1939GOL/MAR K. B. Goldblum, R. W. Martin, and R. B. Young, *Ind. Eng. Chem.* **39**, 1474 (1939).
- 1939NIW/SAT K. Niwa, M. Sato, and M. Yosiya, *J. Chem. Soc. Jpn.* **60**, 918 (1939).
- 1939PAT/SCH W. Patnode and W. J. Scheiber, *J. Am. Chem. Soc.* **61**, 3449 (1939).
- 1939POW/GIA T. M. Powell and W. F. Giauque, *J. Am. Chem. Soc.* **61**, 2366 (1939).
- 1939REI L. Reidel, *Bull. Intern. Inst. Refrig.* **20**, 1 (1939).
- 1939RUE/GIA R. A. Ruehrwein and W. F. Giauque, *J. Am. Chem. Soc.* **61**, 2940 (1939).
- 1939VAN A. Van de Vloed, *Bull. Soc. Chim. Belg.* **48**, 229 (1939).
- 1939VER/MAR F. H. Verhoek and A. L. Marshall, *J. Am. Chem. Soc.* **61**, 2736 (1939).
- 1940AST/KEN J. G. Aston, R. M. Kennedy, and S. C. Schumann, *J. Am. Chem. Soc.* **62**, 2059 (1940).
- 1940AST/MES J. G. Aston and G. H. Messerly, *J. Am. Chem. Soc.* **62**, 1917 (1940).
- 1940BEN/MCH A. F. Benning and R. C. McHarness, *Ind. Eng. Chem.* **32**, 497 (1940).
- 1940BUR A. B. Burg, *J. Am. Chem. Soc.* **62**, 2228 (1940).
- 1940DAR/VER F. R. Darkis, H. E. Vermillion, and P. M. Gross, *Ind. Eng. Chem.* **32**, 946 (1940).
- 1940HOV/LAN F. Hovorka, H. P. Lankelma, and A. E. Axelrod, *J. Am. Chem. Soc.* **62**, 187 (1940).
- 1940HOV/LAN2F. Hovorka, H. P. Lankelma, and I. Schneider, *J. Am. Chem. Soc.* **62**, 1096 (1940).
- 1940HOV/LAN3F. Hovorka, H. P. Lankelma, and W. R. Smith, *J. Am. Chem. Soc.* **62**, 2372 (1940).
- 1940LAM/ROP A. B. Lamb and E. E. Roper, *J. Am. Chem. Soc.* **62**, 806 (1940).
- 1940MES/AST G. H. Messerly and J. G. Aston, *J. Am. Chem. Soc.* **62**, 886 (1940).
- 1940MES/KEN G. H. Messerly and R. M. Kennedy, *J. Am. Chem. Soc.* **62**, 2988 (1940).
- 1940PIT K. S. Pitzer, *J. Am. Chem. Soc.* **62**, 1224 (1940).
- 1940SCH/SAN H. I. Schlesinger, R. T. Sanderson, and A. B. Burg, *J. Am. Chem. Soc.* **62**, 3421 (1940).
- 1940SMI E. R. Smith, *J. Res. Natl. Bur. Stand.* **24**, 229 (1940).
- 1940STU/SAY J. M. Stuckey and J. H. Saylor, *J. Am. Chem. Soc.* **62**, 2922 (1940).
- 1940ZIB A. A. Zibberman-Granovskaya, *Russ. J. Phys. Chem.* **14**, 759 (1940); **14**, 768 (1940); **14**, (1940); *Chem. Abstr.* **35**, 3867 (1941).
- 1941AST/KEN J. G. Aston, R. M. Kennedy, and G. H. Messerly, *J. Am. Chem. Soc.* **63**, 2243 (1941).
- 1941BEL A. F. Belyaev, *Acta Physicochim. URSS* **14**, 523 (1941).
- 1941BOD M. Z. Bodenstein, *Z. Elektrochem. Angew. Phys. Chem.* **47**, 501 (1941); *Chem. Abstr.* **36**, 4014 (1942).
- 1941BRO/DAV L. O. Brockway and N. R. Davidson, *J. Am. Chem. Soc.* **63**, 3287 (1941).
- 1941GIG/RUN P. A. Giguere and R. E. Rundle, *J. Am. Chem. Soc.* **63**, 1135 (1941).
- 1941HEI C. B. Heisig, *J. Am. Chem. Soc.* **63**, 1698 (1941).
- 1941KEN/SAG R. M. Kennedy, M. Sagenkahn, and J. G. Aston, *J. Am. Chem. Soc.* **63**, 2267 (1941).
- 1941KIR/SIT V. A. Kireev and I. P. Sitnikov, *Russ. J. Appl. Chem.* **14**, 483 (1941); *Chem. Abstr.* **36**, 2189 (1942).
- 1941LAU/GIL A. W. Laubengayer and W. F. Gilliam, *J. Am. Chem. Soc.* **63**, 477 (1941).
- 1941LIS M. W. Lister, *J. Am. Chem. Soc.* **63**, 143 (1941).
- 1941MIL G. Milazzo, *Boll. Sci. Facolta Chim. Ind. Bologna*, 94 (1941).
- 1941NIT/SEK I. Nitta and S. Seki, *J. Chem. Soc. Jpn.* **62**, 581 (1941); *Chem. Abstr.* **36**, 15 (1942).
- 1941OSB/GAR D. W. Osborne, C. S. Garner, R. N. Doecher, and D. M. Yost, *J. Am. Chem. Soc.* **63**, 3496 (1941).
- 1941RAL/SEL A. W. Ralston, W. M. Selby, and W. O. Pool, *J. Ind. Eng. Chem.* **33**, 682 (1941).
- 1941RIE L. Riedel, *Z. Ges. Kalte-Ind.* **48**, 9 (1941).
- 1941RUS/RUN H. Russell, Jr., R. E. Rundle, and D. M. Yost, *J. Am. Chem. Soc.* **63**, 2825 (1941).
- 1941SEK S. Seki, *J. Chem. Soc. Jpn.* **62**, 789 (1941).
- 1941SEK/NIT S. Seki and I. Nitta, *J. Chem. Soc. Jpn.* **62**, 907 (1941).
- 1941SMI E. R. Smith, *J. Res. Natl. Bur. Stand.* **26**, 129 (1941).
- 1941STO/FIS J. W. Stout and L. H. Fisher, *J. Chem. Phys.* **9**, 163 (1941).
- 1941YOU/GAR D. M. Youst, C. S. Garner, D. W. Osborne, T. R. Rubin, and H. Russel, Jr., *J. Am. Chem. Soc.* **63**, 2267 (1941).
- 1942BEA/ING J. A. Beattie, H. G. Ingersoll, and W. H. Stockmayer, *J. Am. Chem. Soc.* **64**, 546 (1942).
- 1942BEN S. W. Benson, *Ind. Eng. Chem.* **13**, 189 (1942).
- 1942BEN/KIS S. W. Benson and G. B. Kistiakowsky, *J. Am. Chem. Soc.* **64**, 80 (1942).
- 1942BUR P. E. Burchfield, *J. Am. Chem. Soc.* **64**, 2501 (1942).
- 1942NEL/SMI O. A. Nelson and L. E. Smith, *J. Am. Chem. Soc.* **64**, 3035 (1942).
- 1942OSB/DOE D. W. Osborne, R. W. Doescher, and D. M. Yost, *J. Am. Chem. Soc.* **64**, 169 (1942).
- 1942RUS/OSB H. D. Russell, Jr., D. W. Osborne, and D. M. Yost, *J. Am. Chem. Soc.* **64**, 165 (1942).
- 1942SCH/AST S. C. Schumann, J. G. Aston, and M. Sagenkahn, *J. Am.*

- 1942SUG/SAT B. Suginuma and K. Satozaki, *Rikagaku Kenkyusho Iho* **41**, 432 (1942).
- 1942WIL/GIL G. E. Williams and E. C. Gilbert, *J. Am. Chem. Soc.* **64**, 2776 (1942).
- 1942ZIE/AND W. T. Ziegler and D. H. Andrews, *J. Am. Chem. Soc.* **64**, 2482 (1942).
- 1943AST/FIN J. G. Aston, H. L. Finke, and S. C. Schumann, *J. Am. Chem. Soc.* **65**, 341 (1943).
- 1943AST/SZA J. G. Aston, G. J. Szasz, and H. L. Finke, *J. Am. Chem. Soc.* **65**, 1135 (1943).
- 1943AUD/STE L. F. Audrieth, R. Steinman, and A. D. F. Toy, *Chem. Rev. (Washington, D.C.)* **32**, 109 (1943).
- 1943BUR P. Z. Burbo, *Russ. J. Phys. Chem.* **18**, 286 (1943); *Russ. J. Phys. Chem.* **18**, 253 (1944); *Chem. Abstr.* **39**, 1581 (1945).
- 1943BUR2 A. B. Burg, *J. Am. Chem. Soc.* **65**, 1629 (1943).
- 1943BUR3 A. B. Burg, *J. Am. Chem. Soc.* **63**, 1838 (1943).
- 1943CRA K. S. N. Cramer, *Recl. Trav. Chim. Pays-Bas* **62**, 606 (1943); *Chem. Zentr.* **II**, 2234 (1943); *Chem. Abstr.* **38**, 6148 (1944).
- 1943HAL/REI W. P. Hall and E. E. Reid, *J. Am. Chem. Soc.* **65**, 1466 (1943).
- 1943HOL/MEL G. Holst and L. Melander, *Svensk. Kem. Tid.* **55**, 131 (1943); *Chem. Abstr.* **40**, 2706 (1946).
- 1943KET/KRU J. A. A. Ketelaar and S. Kruyer, *Recl. Trav. Chim. Pays-Bas* **62**, 550 (1943); *Chem. Abstr.* **38**, 1926 (1944).
- 1943LEM/FEL J. F. Lemons and W. A. Felsing, *J. Am. Chem. Soc.* **65**, 46 (1943).
- 1943NIT/SEK I. Nitta and S. Seki, *J. Chem. Soc. Jpn.* **64**, 475 (1943); *Chem. Abstr.* **41**, 3359a (1947).
- 1943PIT/SCO K. S. Pitzer and D. W. Scott, *J. Am. Chem. Soc.* **65**, 803 (1943).
- 1943STE D. R. Stevens, *Ind. Eng. Chem.* **35**, 655 (1943).
- 1944ALT/TRI P. M. Althouse and H. O. Triebold, *Ind. Eng. Chem.* **16**, 605 (1944).
- 1944AST/SAG J. G. Aston, M. L. Sagenkahn, G. J. Szasz, G. W. Moessen, and H. F. Zuhr, *J. Am. Chem. Soc.* **66**, 1171 (1944).
- 1944EME/MAD H. J. Emeleus and A. G. Maddock, *J. Chem. Soc.* 293 (1944).
- 1944IOF/YAM I. I. Ioffe and E. S. Yamp'olskaya, *Zh. Prikl. Khim. (S.-Peterburg)* **17**, 527 (1944); *Chem. Abstr.* **39**, 3986 (1945).
- 1944MCD H. J. McDonald, *J. Phys. Chem.* **48**, 47 (1944).
- 1944MIL G. Milazzo, *Gazz. Chim. Ital.* **74**, 58 (1944).
- 1944MIL2 G. Milazzo, *Gazz. Chim. Ital.* **74**, 49 (1944).
- 1944RUB/LEV T. R. Rubin, B. D. Levedahl, and D. M. Yost, *J. Am. Chem. Soc.* **66**, 279 (1944).
- 1944RUS/GOL H. Russell, Jr., D. R. V. Golding, and D. M. Yost, *J. Am. Chem. Soc.* **66**, 16 (1944).
- 1944SCO/FER R. B. Scott, W. J. Ferguson, and F. G. Brickwedde, *J. Res. Natl. Bur. Stand.* **33**, 1 (1944).
- 1945DAV/WIE H. S. Davis and O. F. Wiedeman, *Ind. Eng. Chem.* **37**, 482 (1945).
- 1945FAW/RAS F. S. Fawcett and H. E. Rasmussen, *J. Am. Chem. Soc.* **67**, 1705 (1945).
- 1945GUT/PIT L. Guttman and K. S. Pitzer, *J. Am. Chem. Soc.* **67**, 324 (1945).
- 1945HOG H. J. Hoge, *J. Res. Natl. Bur. Stand.* **34**, 281 (1945).
- 1945PRO/ROS E. J. Prosen and F. D. Rossini, *J. Res. Natl. Bur. Stand.* **34**, 59 (1945).
- 1945SWI/HOC E. S. Swift, Jr. and H. P. Hochanadel, *J. Am. Chem. Soc.* **67**, 880 (1945).
- 1945WAC/LIN R. C. Waskher, C. B. Linn, and A. V. Grosse, *Ind. Eng. Chem.* **37**, 464 (1945).
- 1945WIL/TAY C. B. Willingham, W. J. Taylor, J. M. Pignocco, and F. D. Rossini, *J. Res. Natl. Bur. Stand.* **35**, 219 (1945).
- 1946ALB/WIL A. Albert and J. B. Willis, *Nature (London)* **157**, 341 (1946); see also J. B. Willis, *Trans. Electrochem. Soc.* **43**, 97 (1946).
- 1946AST/FIN J. G. Aston, H. L. Finke, A. B. Bestul, E. L. Pace, and G. J. Szasz, *J. Am. Chem. Soc.* **68**, 52 (1946).
- 1946BAM/LEV C. H. Bamford, D. L. Levi, and D. M. Newitt, *J. Chem. Soc.* 468 (1946).
- 1946CRO/FEE D. A. Crooks and F. M. Feetham, *J. Chem. Soc.* 899 (1946).
- 1946DZU L. S. Dzung, *Brown Boveri Rev.* **33**, 158 (1946).
- 1946EME/MAD H. J. Emeleus and A. G. Maddock, *J. Chem. Soc.* 1131 (1946).
- 1946FIE/SAY F. H. Field and J. H. Saylor, *J. Am. Chem. Soc.* **68**, 2649 (1946).
- 1946GIL/JON H. Gilman and R. G. Jones, *J. Am. Chem. Soc.* **68**, 517 (1946).
- 1946JON/THO W. J. Jones, L. H. Thomas, E. H. Pritchard, and S. T. Bowden, *J. Chem. Soc.* 824 (1946).
- 1946KIL/PIT J. E. Kilpatrick and K. S. Pitzer, *J. Am. Chem. Soc.* **68**, 1066 (1946).
- 1946KNO/SCH J. W. Knowlton, N. C. Schieltz, and D. MacMillan, *J. Am. Chem. Soc.* **68**, 208 (1946).
- 1946LON/EME L. H. Long, H. J. Emeleus, and H. V. A. Briscoe, *J. Chem. Soc.* 1123 (1946).
- 1946PIT/GUT K. S. Pitzer, L. Guttman, and E. F. Westrum, Jr., *J. Am. Chem. Soc.* **68**, 2209 (1946).
- 1946RUE/POW R. A. Ruehrkeim and T. M. Powell, *J. Am. Chem. Soc.* **68**, 1063 (1946).
- 1946SCO/BRI R. B. Scott and R. G. Brickwedde, *J. Res. Natl. Bur. Stand.* **35**, 501 (1946).
- 1946SPI/PIT R. Spitzer and K. S. Pitzer, *J. Am. Chem. Soc.* **68**, 2537 (1946).
- 1946THO G. W. Thompson, *Chem. Rev. (Washington, D.C.)* **38**, 1 (1946).
- 1947AST/SZA J. G. Aston and G. J. Szasz, *J. Am. Chem. Soc.* **69**, 3108 (1947).
- 1947BAL E. W. Balson, *Trans. Faraday Soc.* **43**, 54 (1947).
- 1947BAL/DEN E. W. Balson, K. G. Denbigh, and N. K. Adam, *Trans. Faraday Soc.* **43**, 42 (1947).
- 1947BEE/JUN J. Beersmans and J. C. Jungers, *Bull. Soc. Chim. Belg.* **56**, 238 (1947).
- 1947BLO/CAM J. M. Blocher, Jr. and I. E. Campbell, *J. Am. Chem. Soc.* **69**, 2100 (1947).
- 1947FOW/HAM R. D. Fowler, J. M. Hamilton, Jr., J. S. Kasper, C. E. Weber, W. B. Burford III, and H. C. Anderson, *Ind. Eng. Chem.* **39**, 375 (1947).
- 1947GOU/HOL C. Gould, G. Holzman, and C. Niemann, *Anal. Chem.* **19**, 204 (1947).
- 1947GRAN A. Granovskaya, *Russ. J. Phys. Chem.* **21**, 967 (1947); *Chem. Abstr.* **42**, 2486d (1948).
- 1947HEI/WIE E. Heilbronner and K. Wieland, *Helv. Chim. Acta* **30**, 947 (1947).
- 1947IVE/PIT D. J. G. Ives, R. W. Pittman, and W. Wardlaw, *J. Chem. Soc.* 1080 (1947).
- 1947IVI/DAI K. J. Ivin and F. S. Dainton, *Trans. Faraday Soc.* **43**, 32 (1947).
- 1947JON/GIA W. M. Jones and W. F. Giauque, *J. Am. Chem. Soc.* **69**, 983 (1947).
- 1947KET/VAN J. A. A. Ketelaar, P. F. Van Velden, and J. S. Zalm, *Recl. Trav. Chim. Pays-Bas* **66**, 721 (1947).
- 1947LAN/GER J. J. Lander and L. H. Germer, American Institute of Mining Metallurgical Engineers: Institute of Metals Division, Metal Technology 14 (No. 6), Technical Publication No. 2259, 1947, as quoted in Ref. [1960MON/COT].
- 1947LEI/SHO A. G. Leibush and E. D. Shorina, *Zh. Prikl. Khim. (S.-Peterburg)* **20**, 69 (1947).
- 1947OSB/GIN N. S. Osborne and D. C. Ginnings, *J. Res. Natl. Bur. Stand.* **39**, 453 (1947); *Chem. Abstr.* **42**, 1795i (1948).
- 1947STR/GAB F. H. Stross, C. M. Gable, and G. C. Rounds, *J. Am. Chem. Soc.* **69**, 1629 (1947).
- 1947STR/MON F. H. Stross, J. M. Monger, and H. De V. Finch, *J. Am. Chem. Soc.* **69**, 1627 (1947).
- 1947STU D. R. Stull, *Ind. Eng. Chem.* **39**, 517 (1947).
- 1947WAD/DOU G. Waddington and D. R. Douslin, *J. Am. Chem. Soc.* **69**, 2275 (1947).
- 1947WAD/TOD G. Waddington, S. S. Todd, and H. M. Huffman, *J. Am. Chem. Soc.* **69**, 22 (1947).
- 1948BAN/EME A. A. Banks, H. J. Emeleus, R. N. Haszeldine, and V. Kerrigan, *J. Chem. Soc.* 2188 (1948).

- 1948BEN/FRA H. E. Bent and R. J. Francel, *J. Am. Chem. Soc.* **70**, 634 (1948).
- 1948BON/ATH C. E. Bonhorst and P. M. Athouse, *Ind. Eng. Chem.* **40**, 2379 (1948).
- 1948BRO/SUJ H. C. Brown and S. Sujishi, *J. Am. Chem. Soc.* **70**, 2878 (1948).
- 1948COA/SUT G. E. Coates and L. E. Sutton, *J. Chem. Soc.* 1187 (1948).
- 1948DAY/NIC H. O. Day, D. E. Nicholson, and W. A. Felsing, *J. Am. Chem. Soc.* **70**, 1784 (1948).
- 1948DOU T. B. Douglas, *J. Am. Chem. Soc.* **70**, 2001 (1948).
- 1948EME/WOO H. J. Emeleus and J. F. Wood, *J. Chem. Soc.* 2183 (1948).
- 1948GAN/JUN J. F. Ganefff and J. C. Jungers, *Bull. Soc. Chim. Belg.* **57**, 82 (1948).
- 1948GIA/JON W. F. Giauque and W. M. Jones, *J. Am. Chem. Soc.* **70**, 120 (1948).
- 1948GLE/HAU O. Glemser and V. Häuser, *Z. Naturforsch. B* **3B**, 159 (1948).
- 1948GOR/GIA J. Gordon and W. F. Giauque, *J. Am. Chem. Soc.* **70**, 1506 (1948).
- 1948HOP/SEA E. R. Hopke and G. W. Sears, *J. Am. Chem. Soc.* **70**, 3801 (1948).
- 1948KEL/CAD K. B. Kellogg and G. H. Cady, *J. Am. Chem. Soc.* **70**, 3986 (1948).
- 1948KLA/MOH F. Klages and K. Mohler, *Chem. Ber.* **81**, 411 (1948).
- 1948MIC/WAS A. Michels and T. Wassenaar, *Physica (Amsterdam)* **14**, 104 (1948).
- 1948NIT/SEK I. Nitta and S. Seki, *J. Chem. Soc. Jpn.* **69**, 85 (1948); *Chem. Abstr.* **44**, 9204e (1950).
- 1948NIT/SEK2 I. Nitta and S. Seki, *J. Chem. Soc. Jpn. Pure Chem. Sect.* **69**, 141 (1948); see also **69**, 143 (1948); *Chem. Abstr.* **46**, 1320i (1952).
- 1948PAC/AST E. L. Pace and J. G. Aston, *J. Am. Chem. Soc.* **70**, 566 (1948).
- 1948RAL/RUS J. H. Raley, F. F. Rust, and W. E. Vaughan, *J. Am. Chem. Soc.* **70**, 88 (1948).
- 1948RED/CHA C. E. Redemann, S. W. Chaikin, and R. B. Fearing, *J. Am. Chem. Soc.* **70**, 631 (1948).
- 1948RED/CHA2 C. E. Redemann, S. W. Chaikin, R. B. Fearing, and D. M. Benedict, *J. Am. Chem. Soc.* **70**, 637 (1948).
- 1948RED/CHA3 C. E. Redemann, S. W. Chaikin, and R. B. Fearing, *J. Am. Chem. Soc.* **70**, 1648 (1948).
- 1948RED/CHA4 C. E. Redemann, S. W. Chaikin, R. B. Fearing, J. W. Rotariu, J. Savit, and D. Van Hoesen, *J. Am. Chem. Soc.* **70**, 3604 (1948).
- 1948SEA/HOP G. W. Sears and E. R. Hopke, *J. Phys. Chem.* **52**, 1137 (1948).
- 1948SMA/SMA P. A. Small, K. W. Small, and P. Cowley, *Trans. Faraday Soc.* **44**, 810 (1948).
- 1948WHI G. H. Whiting, *J. Chem. Soc.* 1209 (1948).
- 1949BIR/BRA J. Birks and R. S. Bradley, *Proc. R. Soc. London* **A198**, 226 (1949).
- 1949BRA/SHE R. S. Bradley and A. D. Shellard, *Proc. R. Soc. London* **A198**, 239 (1949).
- 1949BRASHE2 R. S. Bradley and A. D. Shellard, *Trans. Faraday Soc.* **45**, 501 (1949).
- 1949BUC/COL F. R. Buck, K. F. Coles, G. T. Kennedy, and F. J. Morton, *J. Chem. Soc.* 2377 (1949).
- 1949CAR/HAR A. S. Carson, K. Hartley, and H. A. Skinner, *Trans. Faraday Soc.* **45**, 1159 (1949).
- 1949CAR/HAR2 A. S. Carson, K. Hartely, and H. A. Skinner, *Proc. R. Soc. London* **A195**, 500 (1949).
- 1949DRE/MAR R. R. Dreisbach and R. A. Martin, *Ind. Eng. Chem.* **41**, 2875 (1949).
- 1949DRE/SHR R. R. Dreisbach and S. A. Shrader, *Ind. Eng. Chem.* **41**, 2879 (1949).
- 1949EME/HEA H. J. Emeleus and H. G. Heal, *J. Chem. Soc.* 1696 (1949).
- 1949FOR/BOW M. V. Forward, S. T. Bowden, and J. Jones, *J. Chem. Soc.* S121 (1949).
- 1949FOR/NOR A. F. Forziati, W. R. Norris, and F. D. Rossini, *J. Res. Natl. Bur. Stand.* **43**, 555 (1949).
- 1949GIA/GOR W. F. Giauque and J. Gordon, *J. Am. Chem. Soc.* **71**, 2176 (1949).
- 1949HAT/SUT L. F. Hatch, G. Sutherland, and W. J. Ross, *J. Org. Chem.* **14**, 1130 (1949); (Note: Vapor pressure data presented graphically as log P versus 1/T in paper).
- 1949HIG/END T. Higuchi, N. Endow, and J. E. Willard, *J. Am. Chem. Soc.* **71**, 365 (1949).
- 1949HOL/DOR D. E. Holcomb and C. L. Dorsey, Jr., *Ind. Eng. Chem.* **41**, 2788 (1949); **42**, 570(E) (1950).
- 1949KLA F. Klages, *Chem. Ber.* **82**, 358 (1949).
- 1949KRE/WIE C. B. Kretschmer and R. Wiebe, *J. Am. Chem. Soc.* **71**, 1793 (1949).
- 1949KUH/MAS W. Kuhn and P. Massini, *Helv. Chim. Acta* **32**, 1530 (1949).
- 1949NIC J. C. Nicholson, *J. Chem. Soc.* 1553 (1949).
- 1949NIC/LAF E. Nicolini and P. Laffitte, *Compt. Rend.* **229**, 757 (1949); see also **229**, 935 (1949).
- 1949NIT/SEK I. Nitta, S. Seki, and H. Chihara, *J. Chem. Soc. Jpn. Pure Chem. Sect.* **70**, 387 (1949); *Chem. Abstr.* **45**, 2737d (1951).
- 1949PAR/MOO G. S. Parks and G. E. Moore, *J. Chem. Phys.* **17**, 1151 (1949).
- 1949PAR/MOO2 G. S. Parks, G. E. Moore, M. L. Renquist, B. F. Naylor, L. A. McClaine, P. S. Fujii, and J. A. Hatton, *J. Am. Chem. Soc.* **71**, 3386 (1949).
- 1949PAR/WYN J. R. Partington and A. L. Wynes, *J. Phys. Chem.* **53**, 500 (1949).
- 1949PER/WEB E. S. Perry and W. H. Weber, *J. Am. Chem. Soc.* **71**, 3726 (1949).
- 1949PER/WEB2 E. S. Perry, W. H. Weber, and B. F. Baubert, *J. Am. Chem. Soc.* **71**, 3720 (1949).
- 1949SCO/GRO D. W. Scott, M. E. Gross, G. D. Oliver, and H. M. Huffman, *J. Am. Chem. Soc.* **71**, 1634 (1949).
- 1949SCO/OLI D. W. Scott, G. D. Oliver, M. E. Gross, W. N. Hubbard, and H. M. Huffman, *J. Am. Chem. Soc.* **71**, 2293 (1949).
- 1949SCO/WAD D. W. Scott, G. Waddington, J. C. Smith, and H. M. Huffman, *J. Am. Chem. Soc.* **71**, 2767 (1949).
- 1949SEA/HOP G. W. Sears and E. R. Hopke, *J. Am. Chem. Soc.* **71**, 2575 (1949).
- 1949SEA/HOP2 G. W. Sears and E. R. Hopke, *J. Am. Chem. Soc.* **71**, 1632 (1949).
- 1949THO/EME J. Thompson and H. J. Emeleus, *J. Chem. Soc.* 3080 (1949).
- 1949WAD/KNO G. Waddington, J. W. Knowlton, D. W. Scott, S. D. Oliver, S. S. Todd, W. N. Hubbard, J. C. Smith, and H. M. Huffman, *J. Am. Chem. Soc.* **71**, 797 (1949).
- 1949WAD/SMI G. Waddington, J. C. Smith, D. W. Scott, and H. M. Huffman, *J. Am. Chem. Soc.* **71**, 3902 (1949).
- 1949WIN/KUL L. O. Winstrom and L. Kulp, *Ind. Eng. Chem.* **41**, 2584 (1949).
- 1949YAR/FED N. L. Yarym-Agaev, N. N. Fedos'ev, and K. G. Skorikov, *Zh. Fiz. Khim.* **11**, 1257 (1949).
- 1950AST/MAS J. G. Aston, S. V. R. Mastrangelo, and G. W. Moessen, *J. Am. Chem. Soc.* **72**, 5287 (1950).
- 1950BRI/CAR N. F. H. Bright, T. Carson, and T. A. Dyson, *Research (London)* **3**, 185 (1950); *Chem. Abstr.* **44**, 9367c (1950).
- 1950BUR/KUL A. B. Burg and E. S. Kuljian, *J. Am. Chem. Soc.* **72**, 3103 (1950).
- 1950COL/POP K. F. Coles and F. Popper, *Ind. Eng. Chem.* **42**, 1434 (1950).
- 1950CON/ELV A. Z. Conner, P. J. Elving, J. Benischeck, P. E. Tobias, and S. Steingiser, *Ind. Eng. Chem.* **42**, 106 (1950).
- 1950CRO/SMY R. W. Crowe and C. P. Smyth, *J. Am. Chem. Soc.* **72**, 1098 (1950).
- 1950DAY/FEL H. O. Day and W. A. Felsing, *J. Am. Chem. Soc.* **72**, 1698 (1950).
- 1950EDW G. T. Edwards, *Trans. Faraday Soc.* **46**, 423 (1950).
- 1950FOR/CAM A. F. Forziati, D. L. Camin, and F. D. Rossini, *J. Res. Natl. Bur. Stand.* **45**, 406 (1950).
- 1950HAR/PRI K. Hartley, H. O. Pritchard, and H. A. Skinner, *Trans. Faraday Soc.* **46**, 1019 (1950).
- 1950MAT/SUM J. B. Matthews, J. F. Sumner, and E. A. Moellyn-Hughes, *Trans. Faraday Soc.* **46**, 797 (1950).
- 1950MIC/WAS A. Michels and T. Wassenaar, *Physica (Amsterdam)* **16**,

- 221 (1950).
- 1950NIT/SEK I. Nitta, S. Seki, M. Momotani, and K. Sato, *J. Chem. Soc. Jpn.* **71**, 378 (1950); *Chem. Abstr.* **45**, 6448g (1951).
- 1950NIT/SEK2 I. Nitta, S. Seki, M. Momotani, K. Suzuki, and S. Nakagawa, *Proc. Jpn. Acad.* **26**, 11 (1950); *Chem. Abstr.* **45**, 4545b (1950).
- 1950NIT/SEK3 I. Nitta, S. Seki, and M. Momotani, *J. Chem. Soc. Jpn. Pure Chem. Sect.* **71**, 430 (1950); *Chem. Abstr.* **45**, 6448i (1951).
- 1950NOY/NOY R. M. Noyes, W. A. Noyes, and H. Steinmetz, *J. Am. Chem. Soc.* **72**, 33 (1950).
- 1950SCO/FIN D. W. Scott, H. L. Finke, M. E. Gross, and H. M. Huffman, *J. Am. Chem. Soc.* **72**, 2424 (1950).
- 1950SCO/FIN2 D. W. Scott, H. L. Finke, W. N. Hubbard, J. P. McCullough, M. E. Gross, K. D. Williamson, G. Waddington, and H. M. Huffman, *J. Am. Chem. Soc.* **72**, 4664 (1950).
- 1950SCO/WAD D. W. Scott and G. Waddington, *J. Am. Chem. Soc.* **72**, 4310 (1950).
- 1950WIS/PUC H. Wise, T. T. Puck, and C. F. Farley, *J. Phys. Chem.* **54**, 734 (1950).
- 1951AST/FIN J. G. Aston, H. L. Finke, G. J. Janz, and K. E. Russell, *J. Am. Chem. Soc.* **73**, 1939 (1951).
- 1951AST/JAN J. G. Aston, G. J. Janz, and K. E. Russell, *J. Am. Chem. Soc.* **73**, 1943 (1951).
- 1951BAR/MCC G. M. Barrow and A. L. McClellan, *J. Am. Chem. Soc.* **73**, 573 (1951).
- 1951BRA/WAG R. S. Bradley and G. S. C. Waghorn, *Proc. R. Soc. London* **A206**, 65 (1951).
- 1951BRI N. F. H. Bright, *J. Chem. Soc.* 624 (1951).
- 1951BRY/HOW D. Bryce-Smith and K. E. Howlett, *J. Chem. Soc.* 1141 (1951).
- 1951BUR/CAD L. L. Burger and G. H. Cady, *J. Am. Chem. Soc.* **73**, 4243 (1951).
- 1951BUR/RAN A. B. Burg and C. L. Randolph, *J. Am. Chem. Soc.* **73**, 953 (1951).
- 1951CHA/SKI T. Charnley and H. A. Skinner, *J. Chem. Soc.* 1921 (1951).
- 1951CLA/COC A. M. Clark, A. H. Cockett, and H. S. Eisner, *Proc. R. Soc. London* **A209**, 408 (1951).
- 1951COA/HAR G. E. Coates, J. Harris, and T. Sutcliffe, *J. Chem. Soc.* 2762 (1951).
- 1951EGE/EMT A. C. Egerton, W. Emte, and G. J. Minkoff, *Discuss. Faraday Soc.* **10**, 278 (1951).
- 1951FLO/ALP D. G. Flom, N. Alpert, and P. J. Elving, *Ind. Eng. Chem.* **43**, 1178 (1951).
- 1951GOU/LIN J. Goubeau and R. Link, *Z. Anorg. Allg. Chem.* **267**, 27 (1951).
- 1951GRI/BER J. W. Grisard, H. A. Bernhardt, and G. D. Oliver, *J. Am. Chem. Soc.* **73**, 5725 (1951).
- 1951HAR/PRI K. Hartley, H. O. Pritchard, and H. A. Skinner, *Trans. Faraday Soc.* **47**, 254 (1951).
- 1951HOP/SEA E. R. Hopke and G. W. Sears, *J. Chem. Phys.* **19**, 1345 (1951).
- 1951INO H. Inokuchi, *J. Chem. Soc. Jpn. Pure Chem. Sect.* **72**, 552 (1951).
- 1951MAG/HAR A. Magnus, H. Hartman, and F. Becker, *Erdol u Kohle* **4**, 115 (1951); see also *Z. Phys. Chem. (Leipzig)* **197**, 75 (1951).
- 1951MCC/PER J. P. McCullough, W. B. Person, and R. Spitzer, *J. Am. Chem. Soc.* **73**, 4069 (1951).
- 1951MCE/RIG W. S. McErvan and M. W. Rigg, *J. Am. Chem. Soc.* **73**, 4725 (1951).
- 1951MIL/PAO G. Milazzo and L. Paoloni, *Ann. Chim. (Rome)* **41**, 673 (1951); see also *Rend. Ist. Super. Sanita Rome* **14**, 673 (1951); *Chem. Abstr.* **46**, 3355f (1952).
- 1951NIC G. R. Nicholson, *J. Chem. Soc.* 503 (1951).
- 1951NIT/SEK I. Nitta, S. Seki, H. Chihara, and K. Suzuki, *Science Papers Osaka University No. 29*, (1951), p. 1; *Chem. Abstr.* **46**, 3820 (1952).
- 1951NIT/SEK2 I. Nitta, S. Seki, and K. Suzuki, *Bull. Chem. Soc. Jpn.* **24**, 63 (1951).
- 1951OLI/GRI G. D. Oliver and J. W. Grisard, *J. Am. Chem. Soc.* **73**, 1688 (1951).
- 1951POT/SAY J. C. Potter and J. H. Saylor, *J. Am. Chem. Soc.* **73**, 90 (1951).
- 1951SCO/FIN D. W. Scott, H. L. Finke, J. P. McCullough, M. E. Gross, K. D. Williamson, G. Waddington, and H. M. Huffman, *J. Am. Chem. Soc.* **73**, 261 (1951).
- 1951SMI/BON T. F. Smith and R. F. Bonner, *Ind. Eng. Chem.* **43**, 1169 (1951).
- 1951TIC/LOS A. W. Tickner and F. P. Lossing, *J. Phys. Chem.* **55**, 733 (1951).
- 1951VAU W. E. Vaughan, *Discuss. Faraday Soc.* **10**, 330 (1951).
- 1952AIH A. Aihara, *J. Chem. Soc. Jpn., Pure Chem. Sect.* **73**, 855 (1952); *Chem. Abstr.* **47**, 3634d (1953).
- 1952BRA/COT R. S. Bradley, S. Cotson, and E. G. Cox, *J. Chem. Soc.* 740 (1952).
- 1952BRA/EME G. A. R. Brandt, H. J. Emeleus, and R. N. Haszeldine, *J. Chem. Soc.* 2198 (1952).
- 1952BRA/PLE D. S. Brackman and P. H. Plesch, *J. Chem. Soc.* 2188 (1952).
- 1952BRO I. Brown, *Aust. J. Sci. Res., Ser. A* **5**, 530 (1952).
- 1952BRO/OST C. A. Brown and R. C. Osthoff, *J. Am. Chem. Soc.* **74**, 2340 (1952).
- 1952BYW S. Bywater, *J. Polym. Sci.* **9**, 417 (1952).
- 1952COA/GLO G. E. Coates, F. Glocling, and N. D. Huck, *J. Chem. Soc.* 4496 (1952).
- 1952EDW G. Edwards, *Trans. Faraday Soc.* **48**, 513 (1952).
- 1952FIN/SCO H. L. Finke, D. W. Scott, M. E. Gross, G. Waddington, and H. M. Huffman, *J. Am. Chem. Soc.* **74**, 2804 (1952).
- 1952FUG/BOW E. T. G. Fuge, S. T. Bowden, and W. J. Jones, *J. Phys. Chem.* **56**, 1013 (1952).
- 1952GOT R. Goto, *Bull. Inst. Chem. Res., Kyoto Univ.* **28**, 68 (1952); *Chem. Abstr.* **47**, 34027 (1953).
- 1952GUT/SCO D. W. Scott, W. N. Hubbard, C. Katz, J. P. McCullough, M. E. Gross, K. D. Williamson, and G. Waddington, *J. Am. Chem. Soc.* **74**, 4662 (1952).
- 1952GUT/SCO2 G. B. Guthrie, D. W. Scott, and G. Waddington, *J. Am. Chem. Soc.* **74**, 2795 (1952).
- 1952HUB/FIN W. N. Hubbard, H. L. Finke, D. W. Scott, J. P. McCullough, C. Katz, M. E. Gross, J. F. Messerly, R. E. Pennington, and G. Waddington, *J. Am. Chem. Soc.* **74**, 6025 (1952).
- 1952INO/SHI H. Inokuchi, S. Shiba, T. Handa, and H. Akamatu, *Bull. Chem. Soc. Jpn.* **25**, 299 (1952).
- 1952JON/TAM W. S. Jones and W. S. Tamplin, (glycols) American Chemical Society's Series of Chemistry monographs, 1952.
- 1952KAP/KES L. Kaplan, W. L. Kester, and J. J. Katz, *J. Am. Chem. Soc.* **74**, 5531 (1952).
- 1952LEI/MOR L. C. Leitch and A. T. Morse, *Can. J. Chem.* **30**, 924 (1952).
- 1952MCC/SCO J. P. McCullough, D. W. Scott, H. L. Finke, M. E. Gross, K. D. Williamson, R. E. Pennington, G. Waddington, and H. M. Huffman, *J. Am. Chem. Soc.* **74**, 2801 (1952).
- 1952MOR/PRI C. T. Mortimer, H. O. Pritchard, and H. A. Skinner, *Trans. Faraday Soc.* **48**, 220 (1952).
- 1952OLI/GRI G. D. Oliver and J. W. Grisard, *J. Am. Chem. Soc.* **74**, 2705 (1952).
- 1952REZ/SHV T. N. Rezukhina and V. V. Shyrev, *Vestn. Moskov. Univ.* **7**, 41 (1952); *Chem. Abstr.* **47**, 4676h (1953).
- 1952SCO/DOU D. W. Scott, D. R. Douslin, M. E. Gross, G. D. Oliver, and H. M. Huffman, *J. Am. Chem. Soc.* **74**, 883 (1952).
- 1952SCO/FIN D. W. Scott, H. L. Finke, W. N. Hubbard, J. P. McCullough, G. D. Oliver, M. E. Gross, C. Katz, K. D. Williamson, G. Waddington, and H. M. Huffman, *J. Am. Chem. Soc.* **74**, 4656 (1952).
- 1952SCO/FIN2 D. W. Scott, H. L. Finke, J. P. McCullough, M. E. Gross, R. E. Pennington, and G. Waddington, *J. Am. Chem. Soc.* **74**, 2478 (1952).
- 1952SCO/MAC T. A. Scott, Jr., D. MacMillan, and E. H. Melvin, *Ind. Eng. Chem.* **44**, 172 (1952).
- 1952SIM/HAU J. H. Simons and J. W. Hausteller, *J. Chem. Phys.* **20**, 1516 (1952).
- 1952STI/CAD V. E. Stiles and G. H. Cady, *J. Am. Chem. Soc.* **74**, 3771 (1952).

- 1952WHI G. H. Whipple, *Ind. Eng. Chem.* **44**, 1664 (1952).
- 1952WHI/BER P. T. White, D. G. Bernard-Smith, and F. A. Fidler, *Ind. Eng. Chem.* **44**, 1430 (1952).
- 1953AIH A. Aihara, *J. Chem. Soc. Jpn. Pure Chem. Sect.* **74**, 437 (1953); see also **74**, 631 (1953); **74**, 634 (1953); *Chem. Abstr.* **48**, 1088h (1954).
- 1953AST/WOO J. G. Aston, J. L. Wood, and T. P. Zolki, *J. Am. Chem. Soc.* **75**, 6202 (1953).
- 1953BAR/BRO J. A. Barker, I. Brown, and F. Smith, *Discuss. Faraday Soc.* **15**, 142 (1953).
- 1953BRA/CAR R. S. Bradley and A. D. Care, *J. Chem. Soc.* 1688 (1953).
- 1953BRA/CLE R. S. Bradley and T. G. Cleasby, *J. Chem. Soc.* 1690 (1953).
- 1953BRA/CLE2 R. S. Bradley and T. G. Cleasby, *J. Chem. Soc.* 1681 (1953).
- 1953BRA/COT R. S. Bradley and S. Cotson, *J. Chem. Soc.* 1684 (1953).
- 1953CHA/MOR T. Charnley, C. T. Mortimer, and H. A. Skinner, *J. Chem. Soc.* 1181 (1953).
- 1953CLE/WIS H. E. Clements, K. V. Wise, and S. E. Johnson, *J. Am. Chem. Soc.* **75**, 1593 (1953).
- 1953COP/EVE J. L. Copp and D. H. Everett, *Discuss. Faraday Soc.* **15**, 174 (1953).
- 1953DAV/SCH R. T. Davis, Jr. and R. W. Schiessler, *J. Phys. Chem.* **57**, 966 (1953).
- 1953EDW G. Edwards, *Trans. Faraday Soc.* **49**, 152 (1953).
- 1953EWA A. H. Ewald, *Trans. Faraday Soc.* **49**, 1401 (1953).
- 1953FUR/MCC G. T. Furukawa, R. E. McCoskey, and M. L. Reilly, *J. Res. Natl. Bur. Stand.* **51**, 69 (1953).
- 1953GRU/OST W. T. Grubb and R. C. Osthoff, *J. Am. Chem. Soc.* **75**, 2230 (1953).
- 1953HER/MAR E. F. G. Herington and J. F. Martin, *Trans. Faraday Soc.* **49**, 154 (1953).
- 1953HOF/DEC J. D. Hoffman and B. F. Decker, *J. Chem. Phys.* **57**, 520 (1953).
- 1953JEN/STY A. D. Jenkins and D. W. G. Style, *J. Chem. Soc.* 2337 (1953).
- 1953JOH/FRI O. H. Johnson and H. E. Fritz, *J. Am. Chem. Soc.* **75**, 718 (1953).
- 1953KAR/SAY S. Kardon and J. H. Saylor, *J. Am. Chem. Soc.* **75**, 1997 (1953).
- 1953MCC/SCO J. P. McCullough, D. W. Scott, H. L. Finke, W. N. Hubbard, M. E. Gross, C. Katz, R. E. Pennington, J. F. Messerly, and G. Waddington, *J. Am. Chem. Soc.* **75**, 1818 (1953).
- 1953MIC/WAS A. Michels, T. Wassenaar, P. Louwerse, R. J. Lunbeck, and G. W. Wolkers, *Physica (Amsterdam)* **19**, 287 (1953).
- 1953MIL/PAO G. Milazzo and L. Paoloni, *Gazz. Chim. Ital.* **83**, 867 (1953).
- 1953OLI/MIL G. D. Oliver, H. T. Milton, and J. W. Grisard, *J. Am. Chem. Soc.* **75**, 2827 (1953).
- 1953OST/GRU R. C. Osthoff, W. T. Grubb, and C. A. Burkhard, *J. Am. Chem. Soc.* **75**, 2227 (1953).
- 1953RAT/GWI G. W. Rathjens and W. D. Gwinn, *J. Am. Chem. Soc.* **75**, 5629 (1953).
- 1953SCH/ZEP H. Schaefer and F. Z. Zeppernick, *Z. Anorg. Allg. Chem.* **272**, 274 (1953).
- 1953SCO/FIN D. W. Scott, H. L. Finke, W. N. Hubbard, J. P. McCullough, C. Katz, M. E. Gross, J. P. Messerly, R. E. Pennington, and G. Waddington, *J. Am. Chem. Soc.* **75**, 2795 (1953).
- 1953SEK/SUZ S. Seki and K. Suzuki, *Bull. Chem. Soc. Jpn.* **26**, 209 (1953); **26**, 372 (1953).
- 1953SER/VOI V. V. Serpinski, S. A. Voitkevich, and N. Y. Lyuboshits, *Zh. Fiz. Khim.* **27**, 1032 (1953); *Chem. Abstr.* **49**, 3594e (1955).
- 1953SKI/TEE H. A. Skinner and T. F. S. Tees, *J. Chem. Soc.* 3378 (1953).
- 1953STA/MUL H. Stage, E. Müller, and P. Faldix, *Erdol u Kohle* **6**, 375 (1953); (Enthalpies of vaporization were calculated by fitting published vapor pressure data, taken from various sources to the Antoine equation. Some of the references are to unpublished data measured by chemical companies.)
- 1953STE B. Stevens, *J. Chem. Soc.* 2973 (1953).
- 1953WAG/BUR R. I. Wagner and A. B. Burg, *J. Am. Chem. Soc.* **75**, 3869 (1953).
- 1953WAL K. A. Walsh, U.S. Atomic Energy Commission Report No. LA-1649, 1953, p. 1; *Chem. Abstr.* **55**, 19319c (1961).
- 1954ABR/DAV A. Abrams and T. W. Davis, *J. Am. Chem. Soc.* **76**, 5993 (1954); see also R. B. Mooney and E. B. Ludlam, *Proc. R. Soc. Edinburgh* **49**, 160 (1929).
- 1954BUK/MAJ M. Bukala, J. Majewski, and W. Rodzinski, *Przem. Chem.* **10**, 6 (1954).
- 1954BUR/BAN A. B. Burg and J. Banus, *J. Am. Chem. Soc.* **76**, 3903 (1954).
- 1954CAM/FOR D. L. Camin, A. F. Forziati, and F. D. Rossini, *J. Phys. Chem.* **58**, 440 (1954).
- 1954CRO/JON T. I. Crowell and G. L. Jones, *J. Phys. Chem.* **58**, 666 (1954).
- 1954DAV/JEN M. Davies and D. G. Jenkin, *J. Chem. Soc.* 2374 (1954).
- 1954DAV/JON M. Davies and J. I. Jones, *Trans. Faraday Soc.* **50**, 1042 (1954).
- 1954DUN S. A. Dunn, *J. Am. Chem. Soc.* **76**, 6191 (1954).
- 1954FRI/PIC V. Fried, J. Pick, E. Hala, and O. Vilim, *Chem. Listy* **48**, 774 (1954).
- 1954FUR/MCC G. T. Furukawa, R. E. McCoskey, and M. L. Reilly, *J. Res. Natl. Bur. Stand.* **52**, 11 (1954).
- 1954HUB/WAD W. N. Hubbard and G. Waddington, *Recl. Trav. Chim. Pays-Bas* **73**, 910 (1954).
- 1954JAN/HAS J. Jander and R. N. Haszeldine, *J. Chem. Soc.* 696 (1954).
- 1954JEN/CHA A. C. Jenkins and G. F. Chambers, *Ind. Eng. Chem.* **46**, 2367 (1954).
- 1954JOR T. E. Jordan, *Vapour Pressure of Organic Compounds* (Wiley, New York, 1954).
- 1954LON/SAC L. H. Long and J. F. Sackman, *Trans. Faraday Soc.* **50**, 1177 (1954).
- 1954LOR/WOO G. Lord and A. A. Woolf, *J. Chem. Soc.* 2546 (1954).
- 1954MCC/FIN J. P. McCullough, H. L. Finke, W. N. Hubbard, W. D. Good, R. E. Pennington, J. F. Messerly, and G. Waddington, *J. Am. Chem. Soc.* **76**, 2661 (1954).
- 1954MCC/SCO J. P. McCullough, D. W. Scott, R. E. Pennington, I. A. Hossenlopp, and G. Waddington, *J. Am. Chem. Soc.* **76**, 4791 (1954).
- 1954NIC/SZA G. R. Nicholson, M. Szarc, and J. W. Taylor, *J. Chem. Soc.* 2767 (1954).
- 1954OST/GRU R. C. Osthoff and W. T. Grubb, *J. Am. Chem. Soc.* **76**, 399 (1954).
- 1954POR/RIT A. E. Potter, Jr. and H. L. Ritter, *J. Phys. Chem.* **58**, 1040 (1954).
- 1954SEA/HOP G. W. Sears and E. R. Hopke, *J. Am. Chem. Soc.* **76**, 2026 (1954).
- 1954SEN/SNY K. A. Sense, M. J. Snyder, and J. W. Clegg, *J. Phys. Chem.* **58**, 223 (1954).
- 1954SEN/SNY2 K. A. Sense, M. J. Snyder, and R. B. Filbert, Jr., *J. Phys. Chem.* **58**, 995 (1954).
- 1954SER/VOI V. V. Serpinski, S. A. Voitkevich, and N. Y. Lyuboshits, *Zh. Fiz. Khim.* **28**, 810 (1954); see also **28**, 1969 (1954); *Chem. Abstr.* **49**, 6677C (1955); *Chem. Abstr.* **50**, 4573h (1956).
- 1954SPR/WHI H. D. Springall and T. R. White, *J. Chem. Soc.* 2764 (1954).
- 1954SUJ/WIT S. Sujishi and S. Witz, *J. Am. Chem. Soc.* **76**, 4631 (1954).
- 1954TSU/KAT Y. Tsuzuki, S. Kata, and H. Okazaki, *Kagaku (Kyoto, Jpn.)* **24**, 523 (1954); *Chem. Abstr.* **48**, 13740h (1954).
- 1954WIS K. V. Wise, *J. Am. Chem. Soc.* **76**, 3094 (1954).
- 1955AIH A. Aihara, *J. Chem. Soc. Jpn. Pure Chem. Sect.* **76**, 495 (1955).
- 1955AIH2 A. Aihara, *J. Chem. Soc. Jpn. Pure Chem. Sect.* **76**, 497 (1955).
- 1955AIH3 A. Aihara, *J. Chem. Soc. Jpn., Pure Chem. Sect.* **76**, 492 (1955); *Chem. Abstr.* **49**, 15323c (1955).
- 1955ARM/BRI G. T. Armstrong, F. G. Brickwedde, and R. B. Scott, *J. Res. Natl. Bur. Stand.* **55**, 39 (1955).
- 1955AST/WIL J. G. Aston, P. E. Wills, and T. P. Zolki, *J. Am. Chem. Soc.*

- 77, 3939 (1955).
- 1955AST/ZOL J. G. Aston, T. P. Zolki, and J. L. Wood, *J. Am. Chem. Soc.* **77**, 281 (1955).
- 1955BRO/LAU C. A. Brown and A. W. Laubengayer, *J. Am. Chem. Soc.* **77**, 3699 (1955).
- 1955BRO/ROB F. Brown and P. L. Robinson, *J. Chem. Soc.* 3147 (1955).
- 1955CAM/ROS D. L. Camin and F. D. Rossini, *J. Phys. Chem.* **59**, 1173 (1955).
- 1955CAS/SPR R. C. Cass, H. D. Springall, and P. G. Quincey, *J. Chem. Soc.* 1188 (1955).
- 1955CUM/MCL G. A. N. Cummings and E. McLaughlin, *J. Chem. Soc.* 1391 (1955).
- 1955DRE/MAR R. Dreyer, W. Martin, and U. von Weber, *Chung Hua Nei Ko Tsa Chih* **273**, 324 (1955).
- 1955ENG J. J. Engelsman, Ph.D. thesis, Vrije Universiteit Te Amsterdam, 1955.
- 1955FIS/BIN J. Fischer and J. Bingle, *J. Am. Chem. Soc.* **77**, 6511 (1955).
- 1955FOZ/MOR O. R. Foz Gazulla, J. Morcilio, A. Perez-Masia, and A. Mendes, *Anales Real Soc. Espan. Fis. Quim. (Madrid)* **50B**, 23 (1955).
- 1955FUR/MCC G. T. Furukawa, R. E. McCoskey, M. L. Reilly, and A. W. Herman, *J. Res. Natl. Bur. Stand.* **55**, 201 (1955).
- 1955HAR/SWI N. H. Hartshorne and P. McL. Swift, *J. Chem. Soc.* 3705 (1955).
- 1955ISH/MAT T. Ishiguro and K. Matsumoto, *Yakugaku Zasshi* **75**, 1414 (1955); *Chem. Abstr.* **50**, 53875 (1956).
- 1955JAS/MIL J. J. Jasper and G. B. Miller, *J. Phys. Chem.* **59**, 441 (1955).
- 1955KOS G. H. Kosolapoff, *J. Chem. Soc.* 2964 (1955).
- 1955LON/SAC L. H. Long and J. F. Sackman, *Trans. Faraday Soc.* **51**, 1062 (1955).
- 1955LUF N. W. Luft, *Ind. Chemist* **31**, 502 (1955); *Chem. Abstr.* **50**, 5388c (1956).
- 1955MAC/GRE R. O. MacLaren and N. W. Gregory, *J. Phys. Chem.* **59**, 184 (1955).
- 1955MCC/FIN J. P. McCullough, H. L. Finke, J. F. Messerly, R. E. Pennington, I. A. Hossenlopp, and G. Waddington, *J. Am. Chem. Soc.* **77**, 6119 (1955).
- 1955MEA/STA W. H. Mears, R. F. Stahl, S. R. Orfeo, R. C. Shair, L. F. Kells, W. Thompson, and H. McCann, *Ind. Eng. Chem.* **47**, 1449 (1955).
- 1955MOM/SEK M. Momotani, S. Seki, H. Chihara, and H. Suga, *Bull. Chem. Soc. Jpn.* **28**, 325 (1955).
- 1955MOR/AYS A. T. Morse, P. B. Ayscough, and L. C. Leitch, *Can. J. Chem.* **33**, 453 (1955).
- 1955NEA/WIL E. Neale and L. T. D. Williams, *J. Chem. Soc.* 2485 (1955).
- 1955SCH/WHI R. W. Schiessler and F. C. Whitmore, *Ind. Eng. Chem.* **47**, 1660 (1955); (Note: Data are deposited as supplementary material, Document No. 4597).
- 1955SCO/FIN D. W. Scott, H. L. Finke, J. P. McCullough, M. E. Gross, J. F. Messerly, R. E. Pennington, and G. Waddington, *J. Am. Chem. Soc.* **77**, 4993 (1955).
- 1955SER/VOI V. V. Serpinski, S. A. Voitkevich, and N. Yu Lyuboshits, *Zh. Fiz. Khim.* **29**, 653 (1955); *Chem. Abstr.* **50**, 16231e (1956).
- 1955SHE/GIA R. H. Sherman and W. F. Giaque, *J. Am. Chem. Soc.* **77**, 2154 (1955).
- 1955SPI/STA J. E. Spice, L. A. Staveley, and G. A. Harrow, *J. Chem. Soc.* 100 (1955).
- 1955TSY O. Tsyplikina and J. Ya, *J. Appl. Chem. USSR* **28**, 167 (1955); *Chem. Abstr.* **49**, 9967h (1955).
- 1955VON/GEB E. Von Terres, F. Gebert, H. Hulsemann, H. Petereit, H. Toepsch, and W. Ruppert, *Brennst.-Chem.* **36**, 272 (1955).
- 1956ALL T. L. Allen, *J. Am. Chem. Soc.* **78**, 5476 (1956).
- 1956AMB D. Ambrose, *Trans. Faraday Soc.* **52**, 772 (1956).
- 1956BAR/CAD E. J. Barber and C. H. Cady, *J. Phys. Chem.* **60**, 504 (1956).
- 1956BAR/HAS D. A. Barr and R. N. Haszeldine, *J. Chem. Soc.* 3416 (1956).
- 1956BEY/NIC J. H. Beynon and G. R. Nicholson, *J. Sci. Instrum.* **33**, 376 (1956).
- 1956BRE/UBB D. Brennan and A. R. Ubbelohde, *J. Chem. Soc.* 3011 (1956).
- 1956BRO/DOM H. C. Brown and L. Domash, *J. Am. Chem. Soc.* **78**, 5384 (1956).
- 1956BUR/GOO A. B. Burg and C. D. Good, *J. Inorg. Nucl. Chem.* **2**, 237 (1956).
- 1956CAM/ROS D. L. Camin and F. D. Rossini, *J. Phys. Chem.* **60**, 1446 (1956).
- 1956CAP/JAC R. H. Capps and W. M. Jackson, *J. Phys. Chem.* **60**, 811 (1956).
- 1956DIC W. Dickinson, *Trans. Faraday Soc.* **52**, 31 (1956).
- 1956FIN/SCO H. L. Finke, D. W. Scott, M. E. Gross, J. F. Messerly, and G. Waddington, *J. Am. Chem. Soc.* **78**, 5469 (1956).
- 1956FRO/LOE F. Fromm and M. C. Loeffler, *J. Phys. Chem.* **60**, 252 (1956).
- 1956GAR/SCH D. Garvin and C. Schubert, *J. Phys. Chem.* **60**, 807 (1956).
- 1956GLE/REE D. N. Glew and L. W. Reeves, *J. Phys. Chem.* **60**, 615 (1956).
- 1956GOO/SCO W. D. Good, D. W. Scott, and G. Waddington, *J. Phys. Chem.* **60**, 1090 (1956).
- 1956GRA/PRA P. Gray, M. W. Pratt, and M. J. Larkin, *J. Chem. Soc.* 210 (1956).
- 1956KIR F. W. Kirkbride, *J. Appl. Chem.* **6**, 11 (1956).
- 1956KLO V. P. Klochkov, *Zh. Fiz. Khim.* **30**, 2823 (1956); *Chem. Abstr.* **51**, 13495g (1957).
- 1956LI J. C. M. Li, *J. Am. Chem. Soc.* **78**, 1081 (1956).
- 1956LI/PIT J. C. M. Li and K. S. Pitzer, *J. Am. Chem. Soc.* **78**, 1077 (1956).
- 1956LON/SAC L. H. Long and J. F. Sackman, *Trans. Faraday Soc.* **52**, 1201 (1956).
- 1956MAC A. G. MacDiarmid, *J. Inorg. Nucl. Chem.* **2**, 88 (1956).
- 1956MAG A. Magnus, *Z. Phys. Chem. (Munich)* **9**, 141 (1956).
- 1956MAJ T. G. Majury, *Chem. Ind.* 349 (1956).
- 1956MAJ2 T. G. Majury, *J. Soc. Dyers Colour.* **72**, 41 (1956).
- 1956MAN Manufacturing Chemists Association. Manufacturing Chemists Association Research Project, (1956); see Tables 23-9-2(2.0111)-k, 23-9-2(2.0112)-k, 23-10-2(2.0111)-k, 23-10-2(2.0112)-k, 23-11-2(2.0111)-k, 23-11-2(2.0112)-k; (Note: The authors suspect that the tabulated Antoine Constants for the various compounds were generated in the manner discussed in Ref. [1961LI/ROS]).
- 1956MAN2 Manufacturing Chemists Association, Manufacturing Chemists Association Research Project 1956, Tables 23-18-2(1.01111)-k and 23-18-2(1.01112)-k. The authors suspect that the tabulated Antoine Constants for the various compounds were generated in the manner discussed in Ref. [1961LI/ROS].
- 1956MAT/ERI H. C. Mattraw, C. E. Erickson, and A. W. Laubengayer, *J. Am. Chem. Soc.* **78**, 4901 (1956).
- 1956MIL G. Milazzo, *Ann. Chim. (Rome)* **46**, 1105 (1956); *Chem. Abstr.* **51**, 7791b (1957).
- 1956NEA/WIL E. Neale, L. T. D. Williams, and V. T. Moores, *J. Chem. Soc.* 422 (1956).
- 1956PEN/FIN R. E. Pennington, H. L. Finke, W. N. Hubbard, J. F. Messerly, F. R. Frow, I. A. Hossenlopp, and G. Waddington, *J. Am. Chem. Soc.* **78**, 2055 (1956).
- 1956PEN/SCO R. E. Pennington, D. W. Scott, H. L. Finke, J. P. McCullough, J. F. Messerly, I. A. Hossenlopp, and G. Waddington, *J. Am. Chem. Soc.* **78**, 3266 (1956).
- 1956PIC/FRI J. Pick, V. Fried, E. Hala, and O. Vilim, *Collect. Czech. Chem. Commun.* **21**, 260 (1956).
- 1956ROG/SPE M. T. Rogers and J. L. Speirs, *J. Phys. Chem.* **60**, 1462 (1956).
- 1956SCO/MCC D. W. Scott, J. P. McCullough, W. N. Hubbard, J. F. Messerly, I. A. Hossenlopp, F. R. Frow, and G. Waddington, *J. Am. Chem. Soc.* **78**, 5463 (1956).
- 1956SCO/MCC2D W. Scott, J. P. McCullough, W. D. Good, J. F. Messerly, R. E. Pennington, T. C. Kincheloe, I. A. Hossenlopp, D. R. Douslin, and G. Waddington, *J. Am. Chem. Soc.* **78**, 5457 (1956).
- 1956SEK/SUZ S. Seki, K. Suzuki, and T. Koide, *J. Chem. Soc. Jpn. Pure*

- Chem. Sect. **77**, 346 (1956); Chem. Abstr. **50**, 5343i (1956).
- 1956SER/VOI V. V. Serpinskiĭ, S. A. Voitkevich, and N. Y. Lyuboshits, Zh. Fiz. Khim. **30**, 177 (1956).
- 1956SMI L. Smith, *Acta Chem. Scand.* (1947-1973) **10**, 884 (1956).
- 1956SUZ/ONI K. Suzuki, S. Onishi, T. Koide, and S. Seki, *Bull. Chem. Soc. Jpn.* **29**, 127 (1956).
- 1956TOO E. M. Toops, Jr., *J. Phys. Chem.* **60**, 304 (1956).
- 1956VON/JEN A. Von Brockhaus and E. Jenckel, *Macromol. Chem. Phys.* **18/19**, 262 (1956).
- 1956WIL/COT G. Wilkinson, F. A. Cotton, and J. Birmingham, *J. Inorg. Nucl. Chem.* **2**, 95 (1956).
- 1957AYL J. Aylett, *J. Chem. Soc.* 4152 (1957).
- 1957BAR/SEF W. S. Barnhart, R. J. Seffl, R. F. Wade, F. W. West, and J. L. Zollinger, *J. Chem. Eng. Data* **2**, 80 (1957).
- 1957BES/KOC S. D. Beskov, L. I. Kochetkova, and R. M. Golubeva, *Uchenye Zapiski Moskov. Gosudarst. Pedagog. Inst. Im. V. I. Lenina* **99**, 147 (1957); Chem. Abstr. **53**, 20993d (1959).
- 1957BIS/PAR T. C. Bissot, R. W. Parry, and D. H. Campbell, *J. Am. Chem. Soc.* **79**, 796 (1957).
- 1957BRO/SMI I. Brown and F. Smith, *Aust. J. Chem.* **10**, 423 (1957).
- 1957CAL F. Call, *J. Sci. Food. Agric.* **8**, 81 (1957).
- 1957CAR/DAV E. L. Carpenter and H. S. Davis, *J. Appl. Chem.* **7**, 671 (1957).
- 1957DAL/EME J. W. Dale, H. J. Emeleus, R. N. Haszeldine, and J. H. Moss, *J. Chem. Soc.* 3708 (1957).
- 1957DAY/OES J. H. Day and C. Oestrich, *J. Org. Chem.* **22**, 214 (1957).
- 1957DOB/KEL A. Dobry and R. Keller, *J. Phys. Chem.* **61**, 1448 (1957).
- 1957DYK/SEP J. Dykij, M. Seprakova, and J. Paulech, *Chem. Zvesti* **11**, 461 (1957).
- 1957EAS/HAR B. C. Easton, M. K. Hargreaves, and R. K. S. Mitchell, *J. Appl. Chem.* **7**, 198 (1957).
- 1957FIS/COT A. K. Fischer, F. A. Cotton, and J. Wilkinson, *J. Am. Chem. Soc.* **79**, 2044 (1957).
- 1957FRA/SAN J. W. Frazer and R. H. Sanborn, U.S. Atomic Energy Commission Report No. UCRL-4978, 1957, p. 1; Chem. Abstr. **52**, 8936a (1958).
- 1957GLA/RUL F. Glaser and H. Ruland, *Chem.-Ing.-Tech.* **29**, 772 (1957).
- 1957GRA/PRA P. Gray and M. W. T. Pratt, *J. Chem. Soc.* 2163 (1957).
- 1957KAE/STO H. D. Kaesz and F. G. A. Stone, *J. Chem. Soc.* 1433 (1957).
- 1957KEM/GOL M. D. Kemp, S. Goldhagen, and F. A. Zihlman, *J. Phys. Chem.* **61**, 240 (1957).
- 1957LIT R. Littlewood, *J. Chem. Soc.* 2419 (1957).
- 1957LON/SAC L. H. Long and J. F. Sackman, *Trans. Faraday Soc.* **53**, 1606 (1957).
- 1957MAI/SEY L. Maier, O. Seyforth, F. C. A. Stone, and E. Rochov, *J. Am. Chem. Soc.* **79**, 5884 (1957).
- 1957MCC/DOU J. P. McCullough, D. R. Douslin, J. F. Messerly, I. A. Hossenlopp, T. C. Kincheloe, and G. Waddington, *J. Am. Chem. Soc.* **79**, 4289 (1957).
- 1957MCC/HUB J. P. McCullough, W. N. Hubbard, F. R. Frow, I. A. Hossenlopp, and G. Waddington, *J. Am. Chem. Soc.* **79**, 561 (1957).
- 1957NEI/WHI E. F. Neilson and D. White, *J. Am. Chem. Soc.* **79**, 5618 (1957).
- 1957PED/SKI J. B. Pedley, H. A. Skinner, and C. L. Chernick, *Trans. Faraday Soc.* **53**, 1612 (1957).
- 1957POR/CAD R. S. Porter and G. H. Cady, *J. Am. Chem. Soc.* **79**, 5625 (1957).
- 1957ROS/ACC A. Rose, J. A. Acciari, R. C. Johnson, and W. W. Sanders, *Ind. Eng. Chem.* **49**, 104 (1957).
- 1957ROW/THA J. S. Rowlinson and R. Thacker, *Trans. Faraday Soc.* **53**, 1 (1957).
- 1957SAG A. J. Saggiomo, *J. Org. Chem.* **22**, 1171 (1957).
- 1957SCO/FIN D. W. Scott, H. L. Finke, J. P. McCullough, J. F. Messerly, R. E. Pennington, I. A. Hossenlopp, and G. Waddington, *J. Am. Chem. Soc.* **79**, 1062 (1957).
- 1957SER/VOI V. V. Serpinskiĭ, S. A. Voitkevich, and N. Yu. Lyuboshits, Zh. Fiz. Khim. **31**, 1278 (1957); Chem. Abstr. **52**, 3270f (1958).
- 1957SHE/BRY T. K. Sherwood and J. H. Bryant, Jr., *Can. J. Chem. Eng.* **35**, 51 (1957); Chem. Abstr. **52**, 92a (1958).
- 1957SPI L. H. Spinar, Ph.D. dissertation, University of Wisconsin, 1957; as cited in W. S. Johnson, J. L. Margrave, V. J. Bauer, M. A. Frisch, L. H. Dreger, and W. N. Hubbard, *J. Am. Chem. Soc.* **82**, 1255 (1960).
- 1957SUS/WUH B. P. Susz and J.-J. Wuhrmann, *Helv. Chim. Acta* **40**, 722 (1957).
- 1957VAN A. VanKamp, Ph.D. dissertation, Free University of Amsterdam, 1957.
- 1957WIL R. C. Wilhoit, *J. Phys. Chem.* **61**, 114 (1957).
- 1957WIL/HAR K. D. Williamson and R. H. Harrison, *J. Chem. Phys.* **26**, 1409 (1957).
- 1958/PEP W. J. Peppel, *Ind. Eng. Chem.* **50**, 767 (1958).
- 1958BID/MAR D. P. Biddiscombe and J. F. Martin, *Trans. Faraday Soc.* **54**, 1316 (1958).
- 1958BLO/WEL H. Bloom and B. J. Welch, *J. Phys. Chem.* **62**, 1594 (1958).
- 1958BRA/KAY D. C. Bradley, L. J. Kay, J. D. Swanick, and W. Wardlaw, *J. Chem. Soc.* 3656 (1958).
- 1958BRA/SWA D. C. Bradley and J. D. Swanwick, *J. Chem. Soc.* 3207 (1958).
- 1958BRO/MEA J. A. Brown and W. H. Mears, *J. Phys. Chem.* **62**, 960 (1958).
- 1958BUR/SLO A. B. Burg and P. J. Slota, Jr., *J. Am. Chem. Soc.* **80**, 1107 (1958).
- 1958CAN/NEW S. Cantor, R. F. Newton, W. R. Grimes, and F. F. Blankenship, *J. Phys. Chem.* **62**, 96 (1958).
- 1958CAR/STR A. S. Carson, D. R. Stranks, and B. R. Wihmshurst, *Proc. R. Soc. London* **244**, 72 (1958).
- 1958CAS/FLE R. C. Cass, S. E. Fletcher, C. T. Mortimer, P. G. Quincey, and H. D. Springall, *J. Chem. Soc.* 958 (1958).
- 1958CAS/FLE2 R. C. Cass, S. E. Fletcher, C. T. Mortimer, P. G. Quincey, and H. D. Springall, *J. Chem. Soc.* 2595 (1958).
- 1958CAS/FLE3 R. C. Cass, S. E. Fletcher, C. T. Mortimer, H. D. Springall, and T. R. White, *J. Chem. Soc.* 1406 (1958).
- 1958COL/KEN C. B. Colburn and A. Kennedy, *J. Am. Chem. Soc.* **80**, 5004 (1958).
- 1958DUN/HAN A. S. Dunn and A. Hanrahan, *J. Chem. Soc.* 534 (1958).
- 1958DUN/MUR R. D. Dunlap, C. J. Murphy, Jr., and R. C. Bedford, *J. Am. Chem. Soc.* **80**, 83 (1958).
- 1958EBS/EME E. A. V. Ebsworth and H. J. Emeleus, *J. Chem. Soc.* 2150 (1958).
- 1958EIS/ROT M. Eisenstadt, G. M. Rothberg, and P. Kusch, *J. Chem. Phys.* **29**, 797 (1958).
- 1958EME/SMI H. J. Emeleus and J. D. Smith, *J. Chem. Soc.* 527 (1958).
- 1958FEH/HIT F. Feher and G. Hitzemann, *Z. Anorg. Allg. Chem.* **294**, 50 (1958).
- 1958FIN/SCH A. Finch and H. I. Schlesinger, *J. Am. Chem. Soc.* **80**, 3573 (1958).
- 1958FIS/SCH E. O. Fischer and S. Schreiner, *Chem. Ber.* **91**, 2213 (1958).
- 1958FOW/MOR P. A. Fowell and C. T. Mortimer, *J. Chem. Soc.* 3734 (1958).
- 1958FRO/ROC R. E. Frost and E. G. Rochow, *J. Inorg. Nucl. Chem.* **52**, 201 (1958); Chem. Abstr. **52**, 6999c (1958).
- 1958GOO/FAI W. D. Good, D. M. Fairbrother, and G. Waddington, *J. Phys. Chem.* **62**, 853 (1958).
- 1958GRA/PRA P. Gray and M. W. T. Pratt, *J. Chem. Soc.* 3403 (1958).
- 1958HAB/UEN C. P. Haber and R. K. Uenishi, *Chem. Eng. Data Ser.* **3**, 323 (1958); Chem. Abstr. **53**, 16781b (1959).
- 1958HIE/WAG W. Hieber and G. Wagner, *Chem. Ber.* **618**, 24 (1958).
- 1958HOY/PEP H. Hoyer and W. Peperle, *Z. Elektrochem.* **62**, 61 (1958).
- 1958HUB/DOU W. N. Hubbard, D. R. Douslin, J. P. McCullough, D. W. Scott, S. S. Todd, J. F. Messerly, I. A. Hossenlopp, A. George, and G. Waddington, *J. Am. Chem. Soc.* **80**, 3547 (1958).
- 1958KLI/STR V. G. Klipping and I. N. Stranski, *Z. Anorg. Allg. Chem.* **297**, 23 (1958).
- 1958KLO V. P. Klochkov, Zh. Fiz. Khim. **32**, 1177 (1958); Chem. Abstr. **52**, 19394f (1958).

- 1958KOE/GIA J. K. Koehler and W. F. Giauque, *J. Am. Chem. Soc.* **80**, 2659 (1958).
- 1958LON/SAC L. H. Long and J. F. Sackman, *Trans. Faraday Soc.* **54**, 1797 (1958).
- 1958LOO/DOW C. E. Looney and J. R. Downey, *J. Am. Chem. Soc.* **80**, 2840 (1958).
- 1958MAH/BUR W. Mahler and A. B. Burg, *J. Am. Chem. Soc.* **80**, 6161 (1958).
- 1958MCC/FIN J. P. McCullough, H. L. Finke, D. W. Scott, R. E. Pennington, M. E. Gross, J. F. Messerly, and G. Waddington, *J. Am. Chem. Soc.* **80**, 4786 (1958).
- 1958MOL L. Molard, *Mem. Poudres* **40**, 13 (1958), as cited in Ref. [1977PEL].
- 1958NOV/SEM A. V. Novoselova, K. N. Semenenko, and O. Ya. Turova, *Vestn. Mosk. Univ., Ser. Mat., Mekh., Astron., Fiz., Khim.* **13**, 139 (1958).
- 1958POR/SCH R. F. Porter and R. C. Schoonmaker, *J. Chem. Phys.* **29**, 1070 (1958).
- 1958ROS/PAP A. Rose, B. T. Paphronis, and E. T. Williams, *Chem. Eng. Data Ser.* **3**, 216 (1958).
- 1958SCH/POR R. C. Schoonmaker and R. F. Porter, *J. Chem. Phys.* **29**, 116 (1958).
- 1958SCO/MCC D. W. Scott, J. P. McCullough, J. F. Messerly, R. E. Pennington, I. A. Hossenlopp, H. L. Finke, and G. Waddington, *J. Am. Chem. Soc.* **80**, 55 (1958).
- 1958SEN/STO K. A. Sense and R. W. Stone, *J. Phys. Chem.* **62**, 453 (1958).
- 1958SER/VOI V. V. Serpinski, S. A. Voitkevich, and N. Yu. Lyuboshits, *Trudy Vsesoyuz. Nauch.-Issledovatel. Inst. Sintet. I. Natural. Dushistykh Veshchestv* **4**, 125 (1958).
- 1958SKL/MAR S. I. Sklyarenko, B. I. Markin, and L. B. Belyaeva, *Zh. Fiz. Khim.* **32**, 1916 (1958); *Chem. Abstr.* **53**, 4848f (1959).
- 1958STE/LAN F. Steel and J. Langer, *Z. Anorg. Allg. Chem.* **295**, 315 (1958).
- 1958UNG/MCB H. Ungnade and E. T. McBee, *Chem. Rev. (Washington, D.C.)* **58**, 249 (1958).
- 1959AIH A. Aihara, *Bull. Chem. Soc. Jpn.* **32**, 1242 (1959).
- 1959ALT/BRO E. R. Alton, R. D. Brown, J. C. Carter, and R. C. Taylor, *J. Am. Chem. Soc.* **81**, 3550 (1959).
- 1959AMB E. Amberger, *Angew. Chem.* **71**, 372 (1959); *Chem. Abstr.* **53**, 16780c (1959).
- 1959ARO/KAS Kh. A. Aronovich, L. P. Kastorskii, and K. F. Fedorova, *Zh. Fiz. Khim.* **41**, 20 (1959).
- 1959BIS/FIN E. R. Bissell and M. Finger, *J. Org. Chem.* **24**, 1259 (1959).
- 1959BRA/DRU R. S. Bradley and T. Drury, *Trans. Faraday Soc.* **55**, 1844 (1959).
- 1959BRO/SMI I. Brown and F. Smith, *Aust. J. Chem.* **12**, 407 (1959).
- 1959BUR/GRA A. B. Burg and L. R. Grant, *J. Am. Chem. Soc.* **81**, 1 (1959).
- 1959CHA/VAN P. Chaiyavech and M. Van Winkle, *J. Chem. Eng. Data* **4**, 53 (1959).
- 1959COR/SCH J. F. Cordes and S. Schreiner, *Z. Anorg. Allg. Chem.* **299**, 87 (1959).
- 1959COT/FIS F. A. Cotton, A. K. Fischer, and J. Wilkinson, *J. Am. Chem. Soc.* **81**, 800 (1959).
- 1959CUL/EME W. R. Cullen and H. J. Emeleus, *J. Chem. Soc.* 372 (1959).
- 1959DAR/YOS A. J. Darnell and S. J. Yosim, *J. Phys. Chem.* **63**, 1813 (1959).
- 1959DAV/JON M. Davies and A. H. Jones, *Trans. Faraday Soc.* **55**, 1329 (1959).
- 1959DAV/JON2 M. Davies, A. H. Jones, and G. H. Thomas, *Trans. Faraday Soc.* **55**, 1100 (1959).
- 1959EME/SMI H. J. Emeleus and J. D. Smith, *J. Chem. Soc.* 375 (1959).
- 1959EVA/SKI F. W. Evans and H. A. Skinner, *Trans. Faraday Soc.* **55**, 255 (1959).
- 1959FIS/GRU E. O. Fischer and H. Grubert, *Chem. Ber.* **92**, 2302 (1959).
- 1959FLE/MOR S. E. Fletcher, C. T. Mortimer, and H. D. Springall, *J. Chem. Soc.* 580 (1959).
- 1959FOL/MOR P. A. Folwell and C. T. Mortimer, *J. Chem. Soc.* 2913 (1959).
- 1959GOO/DOU W. D. Good, D. R. Douslin, D. W. Scott, A. George, J. L. Lacina, J. P. Dawson, and G. Waddington, *J. Phys. Chem.* **63**, 1133 (1959).
- 1959GOO/SCO W. D. Good, D. W. Scott, J. L. Lacina, and J. P. McCullough, *J. Phys. Chem.* **63**, 1139 (1959).
- 1959HAR N. Harutada, *Takamine Kenkyusho Nempo* **11**, 66 (1959); *Chem. Abstr.* **55**, 10026e (1961).
- 1959HIL/MCD D. L. Hildenbrand and R. A. McDonald, *J. Phys. Chem.* **68**, 1521 (1959).
- 1959HIL/SIN D. L. Hildenbrand, G. C. Sinke, R. A. McDonald, and W. R. Kramer, *J. Chem. Phys.* **31**, 650 (1959).
- 1959LEA/SPI A. J. Leadbetter and J. E. Spice, *Can. J. Chem.* **37**, 1923 (1959).
- 1959MCC/DOU J. P. McCullough, D. R. Douslin, W. N. Hubbard, S. S. Todd, J. F. Messerly, I. A. Hossenlopp, F. R. Frow, J. P. Dawson, and G. Waddington, *J. Am. Chem. Soc.* **81**, 5884 (1959).
- 1959MCC/PEN J. P. McCullough, R. E. Pennington, J. C. Smith, I. A. Hossenlopp, and G. Waddington, *J. Am. Chem. Soc.* **81**, 5880 (1959).
- 1959MCD/SHR R. A. McDonald, S. A. Shrader, and D. R. Stull, *J. Chem. Eng. Data* **4**, 311 (1959).
- 1959MD K. Mödritzer, *Chem. Ber.* **92**, 2637 (1959); *Chem. Abstr.* **54**, 3180e (1959).
- 1959PAR/MAC L. J. Paridon, G. E. MacWood, and J.-H. Hu, *J. Phys. Chem.* **63**, 1998 (1959).
- 1959PED/SKI J. B. Pedley and H. A. Skinner, *Trans. Faraday Soc.* **55**, 544 (1959).
- 1959PIK/FOS F. P. Pike and C. F. Foster, Jr., *J. Chem. Eng. Data* **4**, 305 (1959).
- 1959SCH/MAR R. S. Scheffee and J. L. Margrave, *J. Chem. Phys.* **31**, 1682 (1959).
- 1959SCO/DOU D. W. Scott, D. R. Douslin, J. F. Messerly, S. S. Todd, I. A. Hossenlopp, T. C. Kincheloe, and J. P. McCullough, *J. Am. Chem. Soc.* **81**, 1015 (1959).
- 1959SEM/GOR K. N. Semenenko and I. V. Gordeev, *Zh. Neorg. Khim.* **4**, 952 (1959); *Chem. Abstr.* **54**, 8206d (1960).
- 1959SEP/PAU M. Seprakova, J. Paulech, and J. Dykyj, *Chem. Zvesti* 313 (1959).
- 1959STE/GRE J. H. Stern and N. W. Gregory, *J. Phys. Chem.* **63**, 556 (1959).
- 1959SUG/SEK H. Suga and S. Seki, *Bull. Chem. Soc. Jpn.* **32**, 1088 (1959).
- 1959TAK/CHI S. Takagi, H. Chihara, and S. Seki, *Bull. Chem. Soc. Jpn.* **32**, 84 (1959).
- 1959TAK/SHI S. Takagi, R. Shintani, H. Chihara, and S. Seki, *Bull. Chem. Soc. Jpn.* **32**, 137 (1959); *Chem. Abstr.* **54**, 1958g (1960).
- 1959TER/BRI E. Terres, L. Brinkmann, D. Fischer, D. Hüllstrung, W. Loz, and G. Weisbrod, *Brennst.-Chem.* **40**, 279 (1959).
- 1959URB L. Urbancova, *Chem. Zvesti* **13**, 224 (1959).
- 1959USA/DEM M. Usanovich and A. I. Dembitskii, *Zh. Obshch. Khim.* **29**, 1781 (1959).
- 1959USA/DEM2 M. Usanovich and A. I. Dembitskii, *Zh. Obshch. Khim.* **29**, 1771 (1959).
- 1959WES/LEV E. F. Westrum, Jr. and N. W. Levitin, *J. Am. Chem. Soc.* **81**, 3544 (1959).
- 1959YEN/REE L. C. Yen and T. M. Reed III, *J. Chem. Eng. Data* **4**, 102 (1959).
- 1960AIH A. Aihara, *Bull. Chem. Soc. Jpn.* **33**, 194 (1960).
- 1960AIH2 A. Aihara, *Bull. Chem. Soc. Jpn.* **33**, 1188 (1960).
- 1960AMB E. Amberger, *Angew. Chem.* **72**, 494 (1960); *Chem. Abstr.* **55**, 22103f (1961); **60**, 4171d (1964).
- 1960AND/BID R. J. L. Andon, D. P. Biddiscombe, J. D. Cox, R. Handley, D. Harrop, E. F. G. Herington, and J. F. Martin, *J. Chem. Soc.* 5246 (1960).
- 1960BED/MOR A. F. Bedford and C. T. Mortimer, *J. Chem. Soc.* 1622 (1960).
- 1960BER/TRU E. W. Berg and J. T. Truemper, *J. Phys. Chem.* **64**, 487 (1960).
- 1960BIR K.-H. Birr, *Z. Anorg. Allg. Chem.* **306**, 21 (1960).
- 1960BRA/BIR R. S. Bradley, C. L. Bird, and F. Jones, *Trans. Faraday Soc.* **56**, 23 (1960).

- 1960BRI/STO F. E. Brinckman and F. G. A. Stone, *J. Am. Chem. Soc.* **82**, 6218 (1960).
- 1960BRO/MCC R. J. Brotherton, A. L. McCloskey, J. L. Boone, and H. M. Manasevit, *J. Am. Chem. Soc.* **82**, 6245 (1960).
- 1960BRO/MCC2R. J. Brotherton, A. L. McCloskey, L. L. Petterson, and H. Steinberg, *J. Am. Chem. Soc.* **82**, 6242 (1960).
- 1960BUD S. Budurov, *Izv. Khim. Inst. Bulg. Akad. Nauk* **7**, 281 (1960).
- 1960CAM/ROS D. L. Camin and F. D. Rossini, *J. Chem. Eng. Data* **5**, 368 (1960).
- 1960CHA/WES S. S. Chang and E. F. Westrum, *J. Phys. Chem.* **94**, 1547 (1960).
- 1960CLA/FOW A. P. Claydon, P. A. Fowell, and C. T. Mortimer, *J. Chem. Soc.* 3284 (1960).
- 1960COT/MON F. A. Cotton and R. R. Monchamp, *J. Chem. Soc.* 533 (1960).
- 1960DAV/THO M. Davies and G. H. Thomas, *Trans. Faraday Soc.* **56**, 185 (1960).
- 1960DJK J. Djkyj, J. Paulech, and M. Seprakova, *Chem. Zvesti* **14**, 327 (1960).
- 1960EDW/KIN J. W. Edwards, G. L. Kington, and R. Mason, *Trans. Faraday Soc.* **56**, 660 (1960).
- 1960EME/NAB H. J. Emelcus and S. N. Nabi, *J. Chem. Soc.* 1103 (1960).
- 1960EME/PUG H. J. Emelcus and H. Pugh, *J. Chem. Soc.* 1108 (1960).
- 1960FRA J. W. Frazer, *J. Inorg. Nucl. Chem.* **16**, 63 (1960).
- 1960GIA/OTT W. F. Giauque and J. B. Ott, *J. Am. Chem. Soc.* **82**, 2689 (1960).
- 1960GLE D. N. Glew, *Can. J. Chem.* **38**, 208 (1960).
- 1960GRE J. H. S. Green, *Chem. Ind. (London)*, 1215 (1960).
- 1960GRE/WAD N. N. Greenwood and K. Wade, *J. Chem. Soc.* 1130 (1960).
- 1960GRI/HAZ C. E. Griffin and R. N. Hazeldine, *J. Chem. Soc.* 1398 (1960).
- 1960HIE/BRA W. Hieber, G. Braun, and W. Beck, *Chem. Ber.* **93**, 901 (1960).
- 1960HIR/STE R. C. Hirt, J. E. Steger, and G. L. Simard, *J. Polym. Sci.* **43**, 319 (1960); *Chem. Abstr.* **54**, 16968b (1960).
- 1960JON A. H. Jones, *J. Chem. Eng. Data* **5**, 196 (1960).
- 1960KAE/PHI H. D. Kaesz, J. P. Phillips, and F. G. A. Stone, *J. Am. Chem. Soc.* **82**, 6228 (1960).
- 1960KAE/STO H. D. Kaesz and F. G. A. Stone, *J. Am. Chem. Soc.* **82**, 6213 (1960).
- 1960KLY/MIS M. L. Klyueva, K. P. Mischenko, and M. K. Fedorov, *Zh. Prikl. Khim. (S.-Peterburg)* **3**, 473 (1960); *Chem. Abstr.* **54**, 11616g (1960).
- 1960LYN/WIL E. J. Lynch and C. R. Wilke, *J. Chem. Eng. Data* **5**, 300 (1960).
- 1960MAC/MAY H. Mackle, R. G. Mayrick, and J. Rooney, *Trans. Faraday Soc.* **56**, 115 (1960).
- 1960MCK/SAG R. A. McKay and B. H. Sage, *J. Chem. Eng. Data* **5**, 21 (1960).
- 1960MCL/TAM D. E. McLaughlin and M. Tamres, *J. Am. Chem. Soc.* **82**, 5618 (1960).
- 1960MCL/TAM2D. E. McLaughlin, M. J. Tamres, and S. Searless, Jr., *J. Am. Chem. Soc.* **82**, 5621 (1960).
- 1960MON/COT R. R. Monchamp and F. A. Cotton, *J. Chem. Soc.* 1438 (1960).
- 1960MUE/IGN C. R. Mueller and J. Ignatowski, *J. Chem. Phys.* **32**, 1430 (1960).
- 1960NEG/MIK H. Negoro, T. Miki, and S. Ueda, *Yakugaku Zasshi* **80**, 665 (1960); *Chem. Abstr.* **54**, 20424f (1960).
- 1960NEG/MIK2H. Negoro, T. Miki, S. Ueda, T. Sanada, and R. Okada, *Yakugaku Zasshi* **80**, 670 (1960).
- 1960NIC G. R. Nicholson, *J. Chem. Soc.* 2377 (1960).
- 1960NOV/MAT J. Novak, J. Matous, and J. Pick, *Collect. Czech. Chem. Commun.* **25**, 583 (1960).
- 1960NOV/MAT2J. Novak, J. Matous, and J. Pick, *Collect. Czech. Chem. Commun.* **25**, 2405 (1960).
- 1960SCH/HIR R. G. Schmitt and R. C. Hirt, *J. Polym. Sci.* **45**, 35 (1960).
- 1960SCH/LEG K. Schwabe and C. Legler, *Z. Elektrochem.* **64**, 902 (1960).
- 1960SIM/GRE R. G. Sime and N. W. Gregory, *J. Phys. Chem.* **64**, 86 (1960).
- 1960STA/STO S. L. Stafford and F. G. A. Stone, *J. Am. Chem. Soc.* **82**, 6238 (1960).
- 1960THO L. H. Thomas, *J. Chem. Soc.* 4906 (1960), as quoted in [2000BRU/DEL].
- 1960TRE/MIL F. M. Trent, F. D. Miller, and G. H. Brown, *J. Chem. Eng. Data* **5**, 110 (1960).
- 1960VAN/CAD W. P. Van Meter and G. H. Cady, *J. Am. Chem. Soc.* **82**, 6005 (1960).
- 1960WAD I. Wadso, *Acta Chem. Scand.* **14**, 566 (1960).
- 1960WAD/KIS T. Wada, E. Kishida, Y. Tomie, H. Suga, S. Seki, and I. Nitta, *Bull. Chem. Soc. Jpn.* **33**, 1317 (1960).
- 1960WAR/MAC L. G. L. Ward and A. G. MacDiarmid, *J. Am. Chem. Soc.* **82**, 2151 (1960).
- 1960WEY/KLI F. Weygand, G. Klipping, and D. Palm, *Chem. Ber.* **93**, 2619 (1960).
- 1960WOO/MUR A. L. Woodman, W. L. Murbach, and M. H. Kaufman, *J. Phys. Chem.* **64**, 658 (1960).
- 1960YAR/KAY R. M. Yarrington and W. B. Kay, *J. Chem. Eng. Data* **5**, 24 (1960).
- 1961AMB E. Amberger, *Chem. Ber.* **94**, 1447 (1961); *Chem. Abstr.* **56**, 8279e (1962).
- 1961AMB/BOE E. Amberger and H. Boeters, *Angew. Chem.* **73**, 114 (1961); *Chem. Abstr.* **55**, 16403h (1961).
- 1961BAN/GIN R. E. Banks, A. E. Ginsberg, and R. N. Haszeldine, *J. Chem. Soc.* 1740 (1961).
- 1961BAR/HAS D. A. Barr, R. N. Haszeldine, and C. J. Willis, *J. Chem. Soc.* 1351 (1961).
- 1961BER/DOW E. W. Berg and H. W. Dowling, *J. Chem. Eng. Data* **6**, 556 (1961).
- 1961BER/SCO W. T. Berg, D. W. Scott, W. N. Hubbard, S. S. Todd, J. F. Messerly, I. A. Hossenlopp, A. Osborn, D. R. Douslin, and J. P. McCullough, *J. Phys. Chem.* **65**, 1425 (1961).
- 1961BUR/GRI A. B. Burg and J. E. Griffiths, *J. Am. Chem. Soc.* **83**, 4333 (1961).
- 1961BUS/IVI W. K. Busfield, K. J. Ivin, H. Mackel, and P. A. G. O'Hare, *Trans. Faraday Soc.* 1058 (1961).
- 1961CAS/RAY J. R. Case, N. H. Ray, and H. L. Roberts, *J. Chem. Soc.* 2070 (1961).
- 1961CAS/RAY2 J. R. Case, N. H. Ray, and H. L. Roberts, *J. Chem. Soc.* 2066 (1961).
- 1961DAC/WYA B. Dacre and P. A. Wyatt, *Trans. Faraday Soc.* **57**, 1281 (1961).
- 1961DAV/MAL M. Davies and V. E. Malpass, *J. Chem. Soc.* 1048 (1961).
- 1961DIT/PER J. F. Ditter, J. C. Perrine, and I. Shapiro, *J. Chem. Eng. Data* **6**, 271 (1961).
- 1961DYK/SEP J. Dykvy, M. Seprakova, and J. Paulech, *Chem. Zvesti* **15**, 465 (1961); *Chem. Abstr.* **56**, 10930b (1962).
- 1961FIS/FRI E. O. Fischer and H. P. Fritz, *Angew. Chem.* **73**, 353 (1961).
- 1961FIS/SCH E. O. Fischer, S. Schreiner, and A. Reckziegel, *Chem. Ber.* **94**, 258 (1961).
- 1961FOW/MOR P. A. Fowell and C. T. Mortimer, *J. Chem. Soc.* 3793 (1961).
- 1961FRO/LOG A. F. Frolov, M. A. Loginova, and M. M. Kiseleva, *Zh. Phys. Khim.* **35**, 1784 (1961).
- 1961GIN A. A. Ginzburg, *Zh. Prikl. Khim. (S.-Peterburg)* **34**, 2569 (1961).
- 1961GOO/LAC W. D. Good, J. L. Lacina, and J. P. McCullough, *J. Phys. Chem.* **65**, 2229 (1961).
- 1961GRI/ONY J. E. Griffiths and M. Onyszczuk, *Can. J. Chem.* **39**, 339 (1961).
- 1961GRO/KLE H. Grosse-Ruyken and R. Kleesaat, *Z. Anorg. Allg. Chem.* **308**, 122 (1961).
- 1961HEI/ILA J. Heinrich, J. Ilavsky, and J. Surovy, *Chem. Zvesti* **15**, 414 (1961).
- 1961HES/WHI P. Hestermans and D. White, *J. Phys. Chem.* **65**, 362 (1961).
- 1961HUF/GRO H. M. Huffman, M. E. Gross, D. W. Scott, and J. P. McCullough, *J. Phys. Chem.* **65**, 495 (1961).
- 1961JOH/KIL W. H. Johnson, M. V. Kilday, and E. J. Prosen, *J. Res. Natl. Bur. Stand., Sect. A* **65A**, 215 (1961).

- 1961LAB/GRE A. Labbauf, J. B. Greenshields, and F. D. Rossini, *J. Chem. Eng. Data* **6**, 261 (1961).
- 1961LI/ROS J. C. M. Li and F. D. Rossini, *J. Chem. Eng. Data* **6**, 268 (1961).
- 1961MAL S. Malanowski, *Bull. Acad. Pol. Sci. Ser. Sci. Chim.* **9**, 71 (1961).
- 1961MAL2 S. Malanowski, *Bull. Acad. Pol. Sci. Ser. Sci. Chim.* **9**, 83 (1961).
- 1961MAT/MCK J. F. Mathews and J. J. McKetta, *J. Phys. Chem.* **65**, 758 (1961).
- 1961MCC/FIN J. P. McCullough, H. L. Finke, W. N. Hubbard, S. S. Todd, J. F. Messerly, D. R. Douslin, and G. Waddington, *J. Phys. Chem.* **65**, 784 (1961).
- 1961NIC/KOB J. K. Nickerson, K. A. Kobe, and J. J. McKetta, *J. Phys. Chem.* **65**, 1037 (1961).
- 1961PAC/BOB E. L. Pace and R. J. Bobka, *J. Chem. Phys.* **35**, 454 (1961).
- 1961ROS/SUP A. Rose and W. R. Supina, *J. Chem. Eng. Data* **6**, 173 (1961).
- 1961ROZ/POL I. L. Rozenfel'd, M. N. Polteva, and V. P. Persiantseva, *Zh. Fiz. Khim.* **35**, 1474 (1961).
- 1961SCH/BOT I. M. Schiopu, O. Bot, and V. Onu, *Bull. Inst. Politeh. Iasi.* **7**, 115 (1961); *Chem. Abstr.* **58**, 10793c (1963).
- 1961SCH/BRA M. Schmeisser and K. Braendle, *Angew. Chem.* **73**, 388 (1961).
- 1961SCH/TAG M. Schmeisser and L. Taglinger, *Chem. Ber.* **94**, 1533 (1961).
- 1961SCO/MES D. W. Scott, J. F. Messerly, S. S. Todd, G. B. Guthrie, I. A. Hossenlopp, R. T. Moore, A. Osborn, W. T. Berg, and J. P. McCullough, *J. Phys. Chem.* **65**, 1320 (1961).
- 1961SMU/BON E. J. Smutny and A. Bondi, *J. Phys. Chem.* **65**, 546 (1961).
- 1961STE/MAC B. Sternback and A. G. MacDiarmid, *J. Am. Chem. Soc.* **83**, 3384 (1961).
- 1961VAN G. Van Dyke Tiers, *J. Org. Chem.* **26**, 2538 (1961).
- 1961WAG/NEW H. Wagner and K. Neumann, *Z. Phys. Chem. [Frankfurt am Main]* **28**, 51 (1961).
- 1961WAL/SMI P. N. Walsh and N. O. Smith, *J. Chem. Eng. Data* **6**, 33 (1961).
- 1961ZIM/GEI H. Zimmermann and H. Geisenfelder, *Z. Elektrochem.* **65**, 368 (1961).
- 1962AMB/BOE E. Amberger and H. Boeters, *Angew. Chem.* **74**, 293 (1962); *Chem. Abstr.* **57**, 4276f (1962).
- 1962AMB/BOE2E E. Amberger and H. Boeters, *Angew. Chem.* **74**, 32 (1962); *Chem. Abstr.* **57**, 12522e (1962).
- 1962BAU/GUN A. Bauder and H. H. Gunthard, *Helv. Chim. Acta* **45**, 1698 (1962).
- 1962BER/MAR J. Berkowitz and J. R. Marquart, *J. Chem. Phys.* **37**, 1853 (1962).
- 1962BER/MCK N. S. Berman and J. J. McKetta, *J. Phys. Chem.* **66**, 1444 (1962).
- 1962BIT/KAU H.-J. Bittrich and E. Kauer, *Z. Phys. Chem. [Frankfurt am Main]* **219**, 224 (1962).
- 1962BIT/KAU2 H.-J. Bittrich, E. Kauer, M. Kraft, G. Schoeppe, W. Soell, and A. Ullrich, *J. Prakt. Chem.* **17**, 250 (1962).
- 1962BRA/JUN J. D. Brandner, N. N. Junk, J. W. Lawrence, and J. Robins, *J. Chem. Eng. Data* **7**, 227 (1962).
- 1962CAR/COO A. Carson, R. Cooper, and D. R. Stranks, *Trans. Faraday Soc.* **58**, 2125 (1962).
- 1962CHA A. M. Chaikin, *Zh. Fiz. Khim.* **36**, 130 (1962); *Chem. Abstr.* **58**, 7387e (1962).
- 1962COH/MAC B. Cohen and A. G. MacDiarmid, *Inorg. Chem.* **1**, 754 (1962).
- 1962CRA/MAC A. D. Craig and A. G. MacDiarmid, *J. Inorg. Nucl. Chem.* **24**, 161 (1962).
- 1962CRA/URE A. D. Craig, J. V. Urenovitch, and A. G. MacDiarmid, *J. Chem. Soc.* 548 (1962).
- 1962DOW A. J. Downs, *J. Chem. Soc.* 4361 (1962).
- 1962EBS/EME E. A. V. Ebsworth, H. J. Emeleus, and N. J. Welman, *J. Chem. Soc.* 2290 (1962).
- 1962EDW/KIN J. W. Edwards and G. L. Kingston, *Trans. Faraday Soc.* **58**, 1323 (1962).
- 1962EME/PAC H. J. Emeleus, K. H. Packer, and N. Welman, *J. Chem. Soc.* 2529 (1962).
- 1962GEI/QUI G. Geiseler, K. Quitzsch, J. Hoffmann, and W. Moeschler, *Z. Phys. Chem. (Leipzig)* **200**, 391 (1962).
- 1962GEI/QUI2 G. Geiseler, K. Quitzsch, J. Hesselbach, and R. Huettig, *Z. Phys. Chem. (Leipzig)* **200**, 79 (1962).
- 1962GOO/DOU W. D. Good, D. R. Douslin, and J. P. McCullough, *J. Phys. Chem.* **66**, 958 (1962).
- 1962GRI/BUR J. E. Griffiths and A. B. Burg, *J. Am. Chem. Soc.* **84**, 3442 (1962).
- 1962HAT/DOU W. E. Hatton, D. R. Douslin, G. C. Sinke, and D. R. Stull, *J. Chem. Eng. Data* **7**, 229 (1962).
- 1962JOH/MCE I. B. Johns, E. A. McElhill, and J. O. Smith, *J. Chem. Eng. Data* **7**, 277 (1962).
- 1962JON/YOW M. M. Jones, B. W. Yow, and W. R. May, *Inorg. Chem.* **1**, 166 (1962).
- 1962KRA/BER M. Kraus, L. Beranek, K. Kochloff, and V. Bazant, *Chem. Prumsyl* **12**, 649 (1962); *Chem. Abstr.* **58**, 10746c (1963).
- 1962KRE A. Kreglewski, *Bull. Acad. Pol. Sci., Ser. Sci. Chem.* **10**, 629 (1962).
- 1962LEB/MIR Y. A. Lebedev, E. A. Miroshnichenko, and A. M. Chaikin, *Dokl. Akad. Nauk SSSR* **145**, 1288 (1962); *Chem. Abstr.* **58**, 2906g (1963).
- 1962MAC/MAY H. Mackle and R. G. Mayrick, *Trans. Faraday Soc.* **58**, 33 (1962).
- 1962MAC/MAY2 H. Mackle and R. G. Mayrick, *Trans. Faraday Soc.* **58**, 230 (1962).
- 1962MAC/MAY3 H. Mackle and R. G. Mayrick, *Trans. Faraday Soc.* **58**, 238 (1962).
- 1962MAN/SUN M. Mansson and S. Sunner, *Acta Chem. Scand.* **16**, 1863 (1962).
- 1962MAR J. J. Martin, *J. Chem. Eng. Data* **7**, 68 (1962).
- 1962OLI/STE J. P. Oliver and L. G. Stevens, *J. Inorg. Nucl. Chem.* **24**, 953 (1962).
- 1962PAS/THO G. J. Pasek and G. L. Thodes, *J. Chem. Eng. Data* **7**, 21 (1962).
- 1962RAB/TEL I. B. Rabinovich, V. I. Tel'noi, L. M. Terman, A. S. Kirillova, and G. A. Razuvaev, *Dokl. Akad. Nauk SSSR* **143**, 133 (1962); see also *Dokl. Chem.* **143**, 171 (1962); *Chem. Abstr.* **57**, 4557e (1962).
- 1962ROS/HEI G. R. Ross and W. G. Heideger, *J. Chem. Eng. Data* **7**, 505 (1962).
- 1962SCO/DOU D. W. Scott, D. R. Douslin, H. L. Finke, W. N. Hubbard, J. F. Messerly, I. A. Hossenlopp, and J. P. McCullough, *J. Phys. Chem.* **66**, 1334 (1962).
- 1962SCO/GOO D. W. Scott, W. D. Good, S. S. Todd, J. F. Messerly, W. T. Berg, I. A. Hossenlopp, J. L. Lucina, A. G. Osborn, and J. P. McCullough, *J. Chem. Phys.* **36**, 406 (1962).
- 1962SEL/SUN P. Sellers and S. Sunner, *Acta Chem. Scand.* **16**, 46 (1962).
- 1962SIN/HIL G. C. Sinke and D. L. Hildenbrand, *J. Chem. Eng. Data* **7**, 74 (1962).
- 1962STE/DOR J. H. Stern and F. H. Dorer, *J. Phys. Chem.* **66**, 97 (1962).
- 1962TJE J. Tebbes, *Acta Chem. Scand.* **16**, 916 (1962).
- 1962TJE2 J. Tebbes, *Acta Chem. Scand.* **16**, 953 (1962).
- 1962VAL/BRO R. H. Valentini, G. E. Brodale, and W. R. Giauque, *J. Phys. Chem.* **66**, 392 (1962).
- 1962VOH/KAN S. P. Vohra, T. L. Kang, K. A. Kobe, and J. J. McKetta, *J. Chem. Eng. Data* **7**, 150 (1962).
- 1962VON/AYM A. J. Von Arvia, P. J. Aymoning, and H. J. Schumacher, *Z. Anorg. Allg. Chem.* **316**, 325 (1962).
- 1962WAD/SMI G. Waddington, J. C. Smith, K. D. Williamson, and D. W. Scott, *J. Phys. Chem.* **66**, 1074 (1962).
- 1962WIL/CAD S. M. Williamson and G. H. Cady, *Inorg. Chem.* **1**, 673 (1962).
- 1963ABE/MAC M. Abedini and A. G. MacDiarmid, *Inorg. Chem.* **2**, 608 (1963).
- 1963ABE/TIL D. C. Abercromby and P. F. Tiley, *J. Chem. Soc.* 4902 (1963).
- 1963AKA/BEL P. A. Akashin, V. I. Belousov, and L. N. Sidorov, *Russ. J. Inorg. Chem.* **8**, 789 (1963).
- 1963AMB/BOE E. Amberger and H. Boeters, *Z. Naturforsch. B* **18B**, 157

- (1963); Chem. Abstr. **59**, 5194g (1963).
- 1963AMB/TOW D. Ambrose and R. J. Townsend, *J. Chem. Soc.* 3614 (1963).
- 1963BEN/MCK E. T. Benyon, Jr. and J. J. McKetta, *J. Phys. Chem.* **67**, 2761 (1963).
- 1963BES R. J. Best, *J. Chem. Eng. Data* **8**, 267 (1963).
- 1963BID/COL D. P. Biddiscombe, R. R. Collerson, R. Handley, E. F. G. Herington, J. F. Martin, and C. H. S. Sprake, *J. Chem. Soc.* 1954 (1963).
- 1963BID/HAN D. P. Biddiscombe, R. Handley, D. Harrop, A. J. Head, G. B. Lewis, J. F. Martin, and C. H. S. Sprake, *J. Chem. Soc.* 5764 (1963).
- 1963BON A. Bondi, *J. Chem. Eng. Data* **8**, 371 (1963).
- 1963BOT/SEI G. A. Bottomley and G. H. F. Seiflow, *J. Appl. Chem.* **13**, 399 (1963).
- 1963BOY R. H. Boyd, *J. Chem. Phys.* **38**, 2529 (1963).
- 1963BRO J. A. Brown, *J. Chem. Eng. Data* **8**, 106 (1963).
- 1963BUL/SER S. S. Bull, I. I. Seregnnaja, and P. R. Tsherbakora, *Khim. Prom. (Moscow)* **7**, 507 (1963).
- 1963CAP/FRI A. Capkova and V. Fried, *Collect. Czech. Chem. Commun.* **28**, 2235 (1963).
- 1963DAN/FLU W. Dannhauser and A. F. Fluechlinger, *J. Chem. Phys.* **38**, 69 (1963).
- 1963DAV/POP J. V. Davies, A. E. Pope, and H. A. Skinner, *Trans. Faraday Soc.* **59**, 2233 (1963).
- 1963DIX/YAR J. A. Dixon, G. R. Yarnell, and J. A. Mountain, *J. Chem. Eng. Data* **8**, 572 (1963).
- 1963EBS/FRA E. A. V. Ebsworth and S. G. Frankiss, *J. Chem. Soc.* 661 (1963).
- 1963EME/HAA H. J. Emeleus and A. Haas, *J. Chem. Soc.* 1272 (1963).
- 1963EME/WEL H. J. Emeleus and N. Welman, *J. Chem. Soc.* 1268 (1963).
- 1963FEI/BOB M. M. Fein, J. Bobinski, N. Mayes, N. Schwartz, and M. S. Cohen, *Inorg. Chem.* **2**, 1111 (1963).
- 1963GAL/TUM N. P. Galkin, Y. N. Tumanov, V. I. Tarasov, and Y. D. Shishov, *Russ. J. Inorg. Chem.* **8**, 1054 (1963).
- 1963GAL/VAR G. L. Gal'chenko and R. M. Varushchenko, *Zh. Fiz. Khim.* **37**, 2513 (1963).
- 1963GIL/CAD W. P. Gilbreath and G. H. Cady, *Inorg. Chem.* **2**, 496 (1963).
- 1963GOO/DOU W. D. Good, D. R. Douslin, and J. P. McCullough, *J. Phys. Chem.* **67**, 1312 (1963).
- 1963GOO/TOD W. D. Good, S. S. Todd, J. F. Messerly, J. L. Lacina, J. P. Dawson, D. W. Scott, and J. P. McCullough, *J. Phys. Chem.* **67**, 1306 (1963).
- 1963GRE/FOS M. A. Greenbaum, J. N. Foster, M. L. Arin, and M. Farber, *J. Phys. Chem.* **67**, 36 (1963).
- 1963HAL/COX J. L. Hales, J. D. Cox, and E. B. Lees, *Trans. Faraday Soc.* **59**, 1544 (1963).
- 1963HAR/HOL S. B. Hartley, W. S. Holmes, J. K. Jaques, M. F. Mole, and J. C. McCoubrey, *Q. Rev., Chem. Soc.* **17**, 204 (1963).
- 1963HIR/HIL H. Hiraoka and J. H. Hildebrand, *J. Phys. Chem.* **67**, 916 (1963).
- 1963HOL/WAG R. R. Holmes and R. P. Wagner, *Inorg. Chem.* **2**, 384 (1963).
- 1963HOR/WEN G. R. Horton and W. W. Wendlandt, *J. Inorg. Nucl. Chem.* **25**, 241 (1963).
- 1963LAU/TRO W. F. Lautsch, A. Tröber, W. Zimmer, L. Mehner, W. Linck, H. M. Lehman, H. Brandenberger, H. Korner, H.-J. Metschker, K. Wagner, and R. Kaden, *Z. Chem.* **3**, 415 (1963).
- 1963LUS/CAD M. Lustig and G. H. Cady, *Inorg. Chem.* **2**, 388 (1963).
- 1963MAR/FRI J. L. Margrave, M. A. Frisch, R. G. Bautista, R. L. Clarke, and W. S. Johnson, *J. Am. Chem. Soc.* **85**, 546 (1963).
- 1963MCC/LAI K. G. McCurdy and K. J. Laidler, *Can. J. Chem.* **41**, 1867 (1963).
- 1963MCC/MES J. P. McCullough, J. F. Messerly, R. T. Moore, and S. S. Todd, *J. Phys. Chem.* **67**, 677 (1963).
- 1963MIL G. A. Miller, *J. Chem. Eng. Data* **8**, 69 (1963).
- 1963MOR/SEL C. T. Mortimer and P. Sellers, *J. Chem. Soc.* 1978 (1963).
- 1963NOB/REE P. Noble, Jr., W. I. Reed, C. J. Hoffman, J. A. Gallagher, and F. G. Borgardt, *AIAA J.* **1**, 395 (1963); Chem. Abstr. **58**, 13701b (1963).
- 1963PAS/ROB G. Pass and H. L. Roberts, *Inorg. Chem.* **2**, 1016 (1963).
- 1963POP/SKI A. E. Pope and H. A. Skinner, *J. Chem. Soc.* 3704 (1963).
- 1963QUI/NOW K. Quitzsch, C. Nowak, P. Winkler, and G. Geiseler, *J. Prakt. Chem.* **20**, 92 (1963).
- 1963RAB/TEL I. B. Rabinovich, V. I. Tel'noi, N. V. Karyakin, and G. A. Razuvaev, *Dokl. Akad. Nauk SSSR* **149**, 324 (1963).
- 1963RAS/NIG R. P. Rastogi, R. P. Nigam, R. N. Sharma, and H. L. Girdhar, *J. Chem. Phys.* **39**, 3042 (1963).
- 1963ROS/SCH A. Rose and V. N. Schrodt, *J. Chem. Eng. Data* **8**, 9 (1963).
- 1963RYS/HAR G. E. Ryschkewitsch, S. W. Harris, E. J. Mezley, H. H. Sisler, E. A. Weilmuenster, and A. B. Garrett, *Inorg. Chem.* **2**, 890 (1963).
- 1963SCO/GOO D. W. Scott, W. D. Good, G. B. Gutherie, S. S. Todd, I. A. Hossenlopp, A. G. Osborn, and J. P. McCullough, *J. Am. Chem. Soc.* **67**, 685 (1963).
- 1963SCO/HUB D. W. Scott, W. N. Hubbard, J. F. Messerly, S. S. Todd, I. A. Hossenlopp, W. D. Good, D. R. Douslin, and J. P. McCullough, *J. Am. Chem. Soc.* **67**, 680 (1963).
- 1963SCO/MES D. W. Scott, J. F. Messerly, S. S. Todd, I. A. Hossenlopp, A. Osborn, and J. P. McCullough, *J. Chem. Phys.* **38**, 532 (1963).
- 1963SHA/KEI I. Shapiro, B. Keilin, R. E. Williams, and C. D. Good, *J. Am. Chem. Soc.* **85**, 3167 (1963).
- 1963SOK/ZHI V. V. Sokolov, L. P. Zhilina, and K. P. Mischenko, *Zh. Prikl. Khim. (S.-Peterburg)* **36**, 750 (1963).
- 1963SPA/MAC E. J. Spanier and A. G. MacDiarmid, *Inorg. Chem.* **2**, 214 (1963).
- 1963URE/MAC J. V. Urenovitch and A. G. MacDiarmid, *J. Chem. Soc.* 1091 (1963).
- 1963VIN/MAR C. G. Vinson, Jr. and J. J. Martin, *J. Chem. Eng. Data* **8**, 74 (1963).
- 1963VLA/GRA O. N. Vlasov, V. A. Granzhan, and L. M. Savenko, *Zh. Prikl. Khim. (S.-Peterburg)* **36**, 2311 (1963).
- 1963VOI S. A. Voitkevich, *Tr. Vses. Nauchn.-Issled. Inst. Sintetich. i Natural'n Dushistykh Veshchestv.*, 91 (1963); Chem. Abstr. **61**, 13121e (1964).
- 1963WOJ J. G. Wojtasinski, *J. Chem. Eng. Data* **8**, 381 (1963).
- 1963WOO/ADI A. L. Woodman and A. Adicoff, *J. Chem. Eng. Data* **8**, 241 (1963).
- 1963WOO/JON J. L. Wood and M. M. Jones, *J. Phys. Chem.* **67**, 1049 (1963).
- 1963WUL/WES C. A. Wulff and E. F. Westrum, Jr., *J. Phys. Chem.* **67**, 2376 (1963).
- 1964AMB/TOW D. Ambrose and R. Townsend, *Trans. Faraday Soc.* **60**, 1025 (1964).
- 1964AYL/PET B. J. Aylett and L. K. Peterson, *J. Chem. Soc.* 3429 (1964).
- 1964BAL/DON P. Balk and D. Dong, *J. Phys. Chem.* **68**, 960 (1964).
- 1964BAN/CHE R. E. Banks, W. M. Cheng, and R. N. Haszeldine, *J. Chem. Soc.* 2485 (1964).
- 1964BER/LAR N. S. Berman, C. W. Larkam, and J. J. McKetta, *J. Chem. Eng. Data* **9**, 218 (1964).
- 1964BIL/COT J. A. Bills and F. A. Cotton, *J. Phys. Chem.* **68**, 806 (1964).
- 1964CAP/FRI A. Capkova and V. Fried, *Collect. Czech. Chem. Commun.* **29**, 336 (1964).
- 1964CAV R. G. Cavell, *J. Chem. Soc.* 1992 (1964).
- 1964CAV/EME R. G. Cavell and H. J. Emeleus, *J. Chem. Soc.* 5825 (1964).
- 1964CAV/EME2 R. G. Cavell and H. J. Emeleus, *J. Chem. Soc.* 5896 (1964).
- 1964CLY/SVE D. D. Clyde and H. Svec, *U. S. Atomic Energy Comm. IS-790*, 1 (1964); Chem. Abstr. **61**, 1293b (1964).
- 1964DUN/CAD L. C. Duncan and G. H. Cady, *Inorg. Chem.* **3**, 850 (1964).
- 1964DUN/CAD2 L. C. Duncan and G. H. Cady, *Inorg. Chem.* **3**, 1045 (1964).
- 1964DUN/THO J. F. Duncan and F. G. Thomas, *J. Chem. Soc.* 360 (1964).
- 1964EHL/KEN R. C. Ehlert, R. A. Kent, and J. L. Margrave, *J. Am. Chem. Soc.* **86**, 5093 (1964).
- 1964ENS/DUR J. V. Enstein, L. I. Duronina, and A. S. Pashinkin, *Zh.*

- Prikl. Khim. (S.-Peterburg) **37**, 2543 (1964).
- 1964FAR/JON D. T. Farrar and M. M. Jones, *J. Phys. Chem.* **68**, 1717 (1964).
- 1964FAW/LIP F. S. Fawcett and R. D. Lipscomb, *J. Am. Chem. Soc.* **86**, 2576 (1964).
- 1964FIE/MAC P. E. Fielding and A. G. Mackay, *Aust. J. Chem.* **17**, 1288 (1964).
- 1964FIS/BUC G. Fisher and A. S. Buchanan, *Aust. J. Chem.* **17**, 481 (1964).
- 1964FIS/PET W. Fischer, T. Petzel, and S. Lauter, *Z. Anorg. Allg. Chem.* **333**, 326 (1964).
- 1964FRI/BUA M. A. Frisch, R. G. Bautista, J. L. Margrave, C. G. Parsons, and J. H. Wotez, *J. Am. Chem. Soc.* **86**, 335 (1964).
- 1964FRI/SAM K. Friedrich and K. Sambach, *J. Chromatogr.* **16**, 22 (1964).
- 1964FUR/REI G. T. Furukawa, M. L. Reilly, H. H. Piccirelli, and M. J. Tenenbaum, *J. Res. Natl. Bur. Stand., Sect. A* **68**, 367 (1964).
- 1964GEI/KON G. Geiseler and W. Konig, *Z. Phys. Chem. (Leipzig)* **227**, 81 (1964).
- 1964GOL/GOR N. A. Goldberg, V. A. Gorbushchenkov, and Z. G. Teplova, *Zh. Prikl. Khim. (S.-Peterburg)* **37**, 745 (1964).
- 1964GOO/LAC W. D. Good, J. L. Lacina, B. L. De Prater, and J. P. McCullough, *J. Phys. Chem.* **68**, 579 (1964).
- 1964GRE/KO M. A. Greenbaum, H. C. Ko, M. Wong, and M. Farber, *J. Phys. Chem.* **68**, 965 (1964).
- 1964GUB/FER A. N. Gubkov, N. A. Fermor, and N. I. Smirnov, *Zh. Prikl. Khim. (S.-Peterburg)* **37**, 2204 (1964).
- 1964HAN/HAR R. Handley, D. Harrop, J. F. Martin, and C. H. S. Sprake, *J. Chem. Soc.* 4404 (1964).
- 1964JON/SED F. Jones and R. Seddon, *Text. Res. J.* **34**, 373 (1964).
- 1964KEL/RIC J. D. Kelley and F. O. Rice, *J. Phys. Chem.* **68**, 3794 (1964).
- 1964KEN/EHL R. A. Kent, T. C. Ehlert, and J. L. Margrave, *J. Am. Chem. Soc.* **86**, 5090 (1964).
- 1964KLE A. V. Kletskii, *Inzh. Fiz. Zh. Akad. Nauk Bel'drussk. SSR* **7**, 40 (1964).
- 1964KLI/FRI V. Kliment, V. Fried, and J. Pick, *Collect. Czech. Chem. Commun.* **29**, 2008 (1964).
- 1964MAC/MCC H. Mackle and R. T. B. Mc Clean, *Trans. Faraday Soc.* **60**, 817 (1964).
- 1964MAC/OHA H. Mackle and P. A. G. O'Hare, *Trans. Faraday Soc.* **60**, 506 (1964).
- 1964MAN E. G. Mangalin, *Khim. Prom.* **4**, 304 (1964).
- 1964MOR D. W. Morecroft, *J. Chem. Eng. Data* **9**, 488 (1964).
- 1964MOR/SEL C. T. Mortimer and P. Sellers, *J. Chem. Soc.* 1965 (1964).
- 1964MUR K. P. Murphy, *J. Chem. Eng. Data* **9**, 259 (1964).
- 1964PAT/PRO C. R. Patrick and G. S. Prosser, *Trans. Faraday Soc.* **60**, 700 (1964).
- 1964PET/BUR L. K. Peterson and A. B. Burg, *J. Am. Chem. Soc.* **86**, 2587 (1964).
- 1964POP/SKI A. E. Pope and H. A. Skinner, *Trans. Faraday Soc.* **60**, 1404 (1964).
- 1964RAT/SHR C. T. Ratcliffe and J. M. Shreeve, *Inorg. Chem.* **3**, 631 (1964).
- 1964ROS/SCH A. Rose and V. Schrod, *J. Chem. Eng. Data* **9**, 12 (1964).
- 1964SER/TIM L. A. Serafimov, V. S. Timofeev, M. P. Strukova, and S. V. L'vov, *Russ. J. Phys. Chem.* **38**, 1018 (1964).
- 1964SEV/SOK N. N. Sevruгова, V. A. Sokorskii, and N. M. Zhavoronkov, *Zh. Prikl. Khim. (Leningrad)* **37**, 1989 (1964).
- 1964SHA/TUB Y. K. Shaulov, V. S. Tubyanskaya, E. V. Evstegneeva, and G. O. Shmyreva, *Russ. J. Phys. Chem.* **38**, 967 (1964).
- 1964SMI/GOR N. K. Smith, G. Gorin, W. D. Good, and J. P. McCullough, *J. Phys. Chem.* **68**, 940 (1964).
- 1964WOO/JON J. L. Wood and M. M. Jones, *Inorg. Chem.* **3**, 1553 (1964).
- 1965AYL/PET B. J. Aylett and L. K. Peterson, *J. Chem. Soc.* 4043 (1965).
- 1965BAC/BET R. A. Back and J. Betts, *Can. J. Chem.* **43**, 2157 (1965).
- 1965BAN/BAR R. E. Banks, M. G. Barlow, and R. N. Haszeldine, *J. Chem. Soc.* 6149 (1965).
- 1965BER/TRU E. W. Berg and J. T. Truemper, *Anal. Chim. Acta* **32**, 245 (1965).
- 1965BLA/GRE J. A. Blauer, M. A. Greenbaum, and M. Farber, *J. Phys. Chem.* **69**, 1069 (1965).
- 1965BOY/CHR R. H. Boyd, R. L. Christensen, and R. Pua, *J. Am. Chem. Soc.* **87**, 3554 (1965).
- 1965BUR/HEN A. B. Burg and J. Heners, *J. Am. Chem. Soc.* **87**, 3092 (1965).
- 1965CAR/WES H. G. Carlson and E. F. Westrum, Jr., *J. Chem. Eng. Data* **10**, 134 (1965).
- 1965CLA/PES L. B. Clark, G. G. Peschel, and I. Tinoco, *J. Phys. Chem.* **69**, 3615 (1965).
- 1965COL/COU R. R. Collerson, R. Handley, J. F. Martin, and C. H. S. Sprake, *J. Chem. Soc.* 3697 (1965).
- 1965COU/GRE J. F. Councell, J. H. S. Green, J. L. Hales, and J. F. Martin, *Trans. Faraday Soc.* **61**, 212 (1965).
- 1965COU/HAL J. F. Councell, J. L. Hales, and J. F. Martin, *Trans. Faraday Soc.* **61**, 1869 (1965).
- 1965CUR/SHA J. Curry and R. W. Shaw, *J. Phys. Chem.* **69**, 344 (1965).
- 1965DAV/KYB M. Davies and B. Kybett, *Trans. Faraday Soc.* **61**, 1608 (1965); see also *Trans. Faraday Soc.* **61**, 1893 (1965).
- 1965DOU/OSB D. R. Douslin and A. Osborn, *Rev. Sci. Instrum.* **42**, 369 (1965).
- 1965EVS/SHM E. V. Evstigneeva and G. O. Shmyreva, *Russ. J. Phys. Chem.* **39**, 529 (1965).
- 1965FIN/HOS H. L. Finke, I. A. Hossenlopp, and W. T. Berg, *J. Phys. Chem.* **69**, 3030 (1965).
- 1965FIN/MES H. L. Finke, J. F. Messerly, and S. S. Todd, *J. Phys. Chem.* **60**, 2094 (1965).
- 1965FRA W. A. Frankhauser, MS Dissertation, Air Force Inst of Technology, Wright Patterson Air Force Base, Ohio; as cited in [1972FON/POM].
- 1965FRA/AST M. Frankosky and J. G. Aston, *J. Phys. Chem.* **69**, 3126 (1965).
- 1965FRA/SHR G. W. Fraser and J. M. Shreeve, *Inorg. Chem.* **4**, 1497 (1965).
- 1965GAK/BAB I. J. Gakh and E. P. Babin, *Russ. J. Phys. Chem.* **39**, 924 (1965).
- 1965HAS/TIP R. N. Haszeldine and A. E. Tipping, *J. Chem. Soc.* 6141 (1965).
- 1965HEI/SUR J. Heinrich, J. Surovy, and J. Dojcansky, *Chem. Zvesti* **19**, 462 (1965).
- 1965HUL/REI H. S. Hull, A. F. Reid, and A. G. Turnbull, *Aust. J. Chem.* **18**, 249 (1965).
- 1965KAL/ROZ L. A. Kalashnikova, E. G. Rozantsev, and A. M. Chaikin, *Izv. Akad. Nauk SSSR*, 800 (1965); *Chem. Abstr.* **63**, 4972c (1965).
- 1965KAR/KYB H. A. Karnes, B. Kybett, M. H. Wilson, J. L. Margrave, and M. S. Newman, *J. Am. Chem. Soc.* **87**, 5554 (1965).
- 1965KIL/BIT H. Kilian and H. J. Bittrich, *Z. Phys. Chem.* **230**, 383 (1965).
- 1965KUD/SAV I. V. Kudryashov and R. I. Savechenko, *Izv. Vyssh. Uchebn. Zaved. Khim. Khim. Tekhnol.* **8**, 602 (1965).
- 1965LIN/FRI J. Linek, V. Fried, and J. Pick, *Collect. Czech. Chem. Commun.* **30**, 1358 (1965).
- 1965LOE/KEN R. E. Loehman, R. A. Kent, and J. L. Margrave, *J. Chem. Eng. Data* **10**, 296 (1965).
- 1965LUT/KOL N. V. Lutugina, V. N. Kolbina, and L. I. Reshetova, *Zh. Prikl. Khim. (S.-Peterburg)* **38**, 1541 (1965).
- 1965MAR P. Martin, *J. Chem. Eng. Data* **10**, 292 (1965).
- 1965MAR/SUS A. N. Marinichev and M. P. Susarev, *Zh. Prikl. Khim. (S.-Peterburg)* **38**, 378 (1965).
- 1965MCD/KIL L. A. McDougall and J. E. Kilpatrick, *J. Chem. Phys.* **42**, 2311 (1965).
- 1965MER/POL I. Mertl and J. Polak, *Collect. Czech. Chem. Commun.* **30**, 3526 (1965).
- 1965MOR G. C. Morris, *J. Mol. Spectrosc.* **18**, 42 (1965).
- 1965NAR P. A. Naro, *J. Chem. Eng. Data* **10**, 86 (1965).
- 1965NIS/LAP L. A. Nisel'son and I. I. Lapivus, *Russ. J. Phys. Chem.* **39**, 80 (1965).
- 1965NIS/LAP2 L. A. Nisel'son and I. I. Lapivus, *Russ. J. Phys. Chem.* **39**, 931 (1965).
- 1965NIX J. F. Nixon, *J. Inorg. Nucl. Chem.* **27**, 1281 (1965).
- 1965NOF/CAD R. Nofle and G. H. Cady, *Inorg. Chem.* **4**, 1010 (1965).
- 1965PRI/JAC S. J. Price and M. G. Jacko, *Can. J. Chem.* **43**, 3481

- (1965).
- 1965PUT/MCE W. E. Putnam, D. M. McEachern, and J. E. Kiplatrick, *J. Chem. Phys.* **42**, 749 (1965).
- 1965RON/HAR R. E. Rondeau and L. A. Harrah, *J. Chem. Eng. Data* **10**, 84 (1965).
- 1965SER/BYK I. I. Serebryannaya and S. Sh. Byk, *Khim. Prom.* **23**, 828 (1965).
- 1965SHA/SCH Y. K. Shaulov, G. O. Schmyreva, and V. S. Tubyanskaya, *Zh. Fiz. Khim.* **39**, 105 (1965).
- 1965SHR/DUN J. M. Shreeve, L. C. Duncan, and G. H. Cady, *Inorg. Chem.* **4**, 1516 (1965).
- 1965SID/AKI L. N. Sidorov, P. A. Akishin, V. B. Shol'ts, and Y. M. Kotenev, *Russ. J. Phys. Chem.* **39**, 1146 (1965).
- 1965SVE/CLY H. Svec and D. D. Clyde, *J. Chem. Eng. Data* **10**, 151 (1965).
- 1965SWA/VAN P. A. Swamy and M. J. Van Winkle, *J. Chem. Eng. Data* **10**, 214 (1965).
- 1965WAD I. Wadso, *Acta Chem. Scand.* **9**, 1079 (1965).
- 1966ABE/MAC M. Abedini and A. G. MacDiarmid, *Inorg. Chem.* **5**, 2040 (1966).
- 1966BAN/BAR R. W. Banks, M. G. Barlow, W. R. Deem, R. N. Haszeldine, and D. R. Taylor, *J. Chem. Soc. C* **1966**, 981.
- 1966BEE/MOR A. E. Beezer and C. T. Mortimer, *J. Chem. Soc. A* 514 (1966).
- 1966BON A. A. Boni, *J. Electrochem. Soc.* **113**, 1089 (1966); *Chem. Abstr.* **65**, 17721h (1966).
- 1966BOR/NAK B. Börjesson, Y. Nakase, and S. Sunner, *Acta Chem. Scand.* **20**, 803 (1966).
- 1966BOT/ADL T. R. Bott and H. N. Adler, *J. Chem. Eng. Data* **11**, 25 (1966).
- 1966BOY R. H. Boyd, *Tetrahedron* **22**, 119 (1966).
- 1966BRA/HIL D. C. Bradley and M. J. Hillyer, *Trans. Faraday Soc.* **62**, 2374 (1966).
- 1966BRA/SEM V. A. Branzhan, S. V. Semenenko, and O. G. Kirillova, *Educ. Psychol. Meas.* **39**, 1399 (1966); *Chem. Abstr.* **65**, 9820f (1966).
- 1966BRO/FRA D. H. Brown, G. W. Fraser, and G. W. A. Sharp, *J. Chem. Soc. A* 171 (1966).
- 1966BUR A. B. Burg, *J. Am. Chem. Soc.* **88**, 4298 (1966).
- 1966BUR/STR A. B. Burg and G. B. Street, *Inorg. Chem.* **5**, 1532 (1966).
- 1966CAR/STE J. L. Carson, R. C. Stewart, and A. G. Williamson, *J. Chem. Eng. Data* **11**, 231 (1966).
- 1966COL/SKI D. J. Coleman and H. A. Skinner, *Trans. Faraday Soc.* **62**, 1721 (1966).
- 1966COW/BUR A. H. Cowley, A. B. Burg, and W. R. Cullen, *J. Am. Chem. Soc.* **88**, 3178 (1966).
- 1966CUB D. Cubicciotti, *J. Phys. Chem.* **70**, 2410 (1966).
- 1966DEL/SHR J. J. Delfino and J. M. Shreeve, *Inorg. Chem.* **5**, 308 (1966).
- 1966DES/CAD D. D. DesMarteau and G. H. Cady, *Inorg. Chem.* **5**, 169 (1966).
- 1966DOB R. C. Dobbie, *J. Chem. Soc. A* 1555 (1966).
- 1966DOB/EME R. C. Dobbie and H. J. Emeleus, *J. Chem. Soc. A* 933 (1966).
- 1966EAR/HIL B. L. Earl, B. K. Hill, and J. M. Shreeve, *Inorg. Chem.* **5**, 2184 (1966).
- 1966EME/ONA H. J. Emeleus and T. Onak, *J. Chem. Soc. A* 1291 (1966).
- 1966EME/TAT H. J. Emeleus and B. W. Tattershall, *J. Inorg. Nucl. Chem.* **28**, 1823 (1966).
- 1966EVA/TIL F. D. Evans and P. F. Tiley, *J. Chem. Soc. B* **1966**, 134.
- 1966FIN/GAR A. Finch, P. J. Gardner, and E. J. Pearn, *Trans. Faraday Soc.* **62**, 1072 (1966).
- 1966GAT/DRA G. Gattrow and M. Draeger, *Z. Zorn. Allgem. Chem.* **343**, 232 (1966); *Chem. Abstr.* **64**, 18513f (1966).
- 1966GEI/FRU G. Geiseler, J. Fruwert, and R. Huetting, *Chem. Ber.* **99**, 1594 (1966).
- 1966GEI/QUI G. Geiseler, K. Quitzsch, H.-J. Rauh, H. Schaffernicht, and H. Walther, *Ber. Bunsenges. Phys. Chem.* **70**, 551 (1966).
- 1966GOO/DEP W. D. Good and B. L. DePrater, *J. Phys. Chem.* **70**, 3606 (1966).
- 1966GOO/MAN W. D. Good and M. Mansson, *J. Phys. Chem.* **70**, 97 (1966).
- 1966GRA/BUR D. N. Gray and G. Burton, *J. Chem. Eng. Data* **11**, 59 (1966).
- 1966HAS/TIP R. N. Haszeldine and A. E. Tipping, *J. Chem. Soc. C* 1236 (1966).
- 1966HOL I. Holec, *Chem. Prumysl* **16**, 267 (1966).
- 1966JEN/SCH D. J. Jensen and E. D. Schall, *J. Agric. Food Chem.* **14**, 123 (1966).
- 1966JON/KRA F. Jones and J. Kraska, *J. Soc. Dyers Colour.* **82**, 333 (1966).
- 1966KUD/ZWO A. P. Kudchadker and B. J. Zwolinski, *J. Chem. Eng. Data* **11**, 253 (1966).
- 1966KYB/CAR B. D. Kybett, S. Carroll, P. Natalis, D. W. Bonnell, J. L. Margrave, and J. L. Franklin, *J. Am. Chem. Soc.* **88**, 626 (1966).
- 1966LAU/SCA A. W. Laubengayer and C. W. Scaife, *J. Chem. Eng. Data* **11**, 172 (1966).
- 1966LEB/ROS Y. A. Lebedev, E. G. Rosantsev, L. A. Kalashnikova, V. P. Lebedev, M. B. Nieman, and Y. A. Apin, *Dokl. Akad. Nauk SSSR* **168**, 104 (1966); see also *Int. J. Comput. Math.* **168**, 460 (1966).
- 1966LUC G. P. Luchinskii, *Zh. Fiz. Khim.* **40**, 593 (1966).
- 1966MAN/SUN M. Mansson and S. Sunner, *Acta Chem. Scand.* **20**, 845 (1966).
- 1966MEA/ROS W. H. Mears, E. Rosenthal, and J. V. Sinka, *J. Chem. Eng. Data* **11**, 338 (1966).
- 1966MEY/FRA M. D. Meyers and S. Frank, *Inorg. Chem.* **5**, 1455 (1966).
- 1966MEY/MET R. Meyer and J. C. R. Metzger, *Acad. Sci. Paris Ser. C* **263**, 1333 (1966).
- 1966MEY/WAG E. F. Meyer and R. E. Wagner, *J. Phys. Chem.* **70**, 3162 (1966).
- 1966MIL/MUR N. E. Miller, M. D. Murphy, and D. L. Reznicek, *Inorg. Chem.* **5**, 1832 (1966).
- 1966MOR/TAM H. L. Morris, M. Tamres, and S. Searles, *Inorg. Chem.* **5**, 2156 (1966).
- 1966OSB/DOU A. N. Osborn and D. R. Douslin, *J. Chem. Eng. Data* **11**, 502 (1966).
- 1966PIR J. Pirsch, *Monatsch. Chem.* **97**, 260 (1966).
- 1966RIN/ONE M. A. Ring, H. E. O'Neal, A. H. Kadhim, and F. J. Jappe, *J. Organomet. Chem.* **5**, 124 (1966).
- 1966ROS J. M. Rosen, private communication, as quoted in Ref. [1968MAR/ARM].
- 1966TEL/RAB V. I. Tel'noi and I. B. Rabinovich, *Zh. Fiz. Khim.* **40**, 1556 (1966).
- 1966THO/MEA L. M. Thomas and R. Meatyard, *J. Chem. Soc. Inorg. Phys. Theoret.* **1966**, 92.
- 1966WAD I. Wadso, *Acta Chem. Scand.* **20**, 544 (1966).
- 1966ZAL/STR A. A. Zalykin and Y. A. Strepikheev, *Russ. J. Appl. Chem.* **39**, 2448 (1966).
- 1966ZIM/ROB M. F. Zimmer, R. A. Robb, and G. A. Carpenter, *J. Chem. Eng. Data* **11**, 577 (1966).
- 1967AND/HAM M. L. Anderson and R. N. Hammer, *J. Chem. Eng. Data* **12**, 442 (1967).
- 1967AYL/CAM B. J. Aylett and J. M. Campbell, *Inorg. Nucl. Chem. Lett.* **3**, 137 (1967).
- 1967BAL/LAP J. C. Baldwin, M. F. Lappert, J. B. Pedley, and J. A. Treverton, *J. Chem. Soc. A* 1980 (1967).
- 1967BAN/HAS R. E. Banks, R. N. Haszeldine, and R. Hatton, *J. Chem. Soc. C* 427 (1967).
- 1967BER/WES H. A. Berman and E. D. West, *J. Chem. Eng. Data* **12**, 197 (1967).
- 1967BOY/GUH R. H. Boyd, K. R. Guha, and R. Wuthrich, *J. Phys. Chem.* **71**, 2187 (1967).
- 1967BRA/SZI W. K. Bratton, I. Szilard, and C. A. Cupas, *J. Org. Chem.* **32**, 2019 (1967).
- 1967BUC/COX E. Buckley and J. D. Cox, *Trans. Faraday Soc.* **63**, 895 (1967).
- 1967CHA/CAV T. L. Charlton and R. G. Cavell, *Inorg. Chem.* **6**, 2204 (1967).
- 1967CHI/SIM Y. T. Chia and H. E. Simmons, *J. Am. Chem. Soc.* **89**, 2638 (1967).
- 1967CHR/SHI P. M. Christopher and A. Shilman, *J. Chem. Eng. Data* **12**, 333 (1967).

- 1967CRO/TAY G. A. Crowder, Z. L. Taylor, T. M. Reed III and J. A. Young, *J. Chem. Eng. Data* **12**, 481 (1967).
- 1967CRU/CUT A. J. B. Cruickshank and A. J. B. Cutler, *J. Chem. Eng. Data* **12**, 326 (1967).
- 1967DAN/GOL S. M. Danov and Y. D. Golubev, Tr. Khim. Khim. Tekhnol, 52 (1967); Chem. Abstr. **68**, 95253a (1968).
- 1967DAT/KAN K. V. Datye, P. J. Kangle, and B. Milicevic, Textilveredlung **2**, 263 (1967).
- 1967EBS/THO E. A. V. Ebsworth and J. C. Thompson, J. Chem. Soc. A 69 (1967).
- 1967EDE/THO M. P. Edejer and G. Thodos, *J. Chem. Eng. Data* **12**, 206 (1967).
- 1967ERM/SKR N. V. Ermakov and V. P. Skripov, Zh. Fiz. Khim. **41**, 77 (1967).
- 1967FIN/GAR A. Finch, P. J. Gardner, and G. B. Watts, *Trans. Faraday Soc.* **63**, 1603 (1967).
- 1967FIN/GAR A. Finch, P. J. Gardner, E. J. Pearn, and G. B. Watts, *Trans. Faraday Soc.* **63**, 1880 (1967).
- 1967FIN/GRA D. H. Fine and P. Gray, *Combust. Flame* **11**, 71 (1967); Chem. Abstr. **66**, 117537j (1967).
- 1967FRI/GAL V. Fried, P. Gallant, and G. B. Schneier, *J. Chem. Eng. Data* **12**, 504 (1967).
- 1967GEI/SCH G. Geiseler, H. Schaffernicht, and H. Walther, Ber. Bunsenges. Phys. Chem. **71**, 196 (1967).
- 1967GLE/BIE O. Glemser, U. Biermann, and M. Fild, *Chem. Ber.* **100**, 1082 (1967).
- 1967GOE/SCH K. H. Goebel, J. Schaffenger, and G. Opel, Chem. Tech. (Leipzig) **19**, 307 (1967).
- 1967GOL/LAP Y. V. Golubkov, I. I. Lapidus, and L. A. Nisel'son, Russ. J. Phys. Chem. **41**, 1122 (1967).
- 1967GRE/JON H. S. Green and F. Jones, *Trans. Faraday Soc.* **63**, 1612 (1967).
- 1967HAC/MAT M. Haccuria and M. P. Mathieu, Ind. Chim. Belg. **32**, 165 (1967).
- 1967HAL/LEE J. L. Hales, E. B. Lees, and D. J. Ruxton, *Trans. Faraday Soc.* **63**, 1876 (1967).
- 1967HER A. J. Herbert, *Trans. Faraday Soc.* **63**, 555 (1967).
- 1967HIL/IRV J. O. Hill and R. J. Irving, J. Chem. Soc. A 1413 (1967).
- 1967HIR/SUD M. Hirata and S. Suda, Kagaku Kogaku Ronbunshu **31**, 339 (1967).
- 1967HUL/REI H. S. Hull, A. F. Reid, and A. G. Turnbull, *Inorg. Chem.* **6**, 805 (1967).
- 1967KEN/CUB F. J. Keneshea and D. Cubicciotti, *J. Phys. Chem.* **71**, 1958 (1967).
- 1967KLE/PET A. B. Kletshii and L. E. Petric, Zh. Fiz. Khim. **41**, 1183 (1967).
- 1967KON/ZHU I. I. Konstantinov, E. Z. Zhuravlev, and Z. G. Teplova, Educ. Psychol. Meas. **40**, 1084 (1967).
- 1967LAP/NIS L. L. Lapidus, L. A. Nisel'son, and A. A. Karateeva, Russ. J. Phys. Chem. **41**, 241 (1967).
- 1967LES/OGO T. M. Lesteva, S. K. Ogorodnikov, and A. I. Morozova, Zh. Prikl. Khim. (Leningrad) **40**, 891 (1967).
- 1967LUS M. Lustig, *Inorg. Chem.* **6**, 1064 (1967).
- 1967MAG J. H. Magill, *J. Chem. Phys.* **47**, 2802 (1967).
- 1967MEE/GOL A. C. Meeks and I. J. Goldfarb, *J. Chem. Eng. Data* **12**, 196 (1967).
- 1967MIR/LEB E. A. Miroshnichenko, Y. A. Lebende, S. A. Shevelev, V. I. Gulevskaya, A. A. Fainzil'berg, and A. Y. Apin, Zh. Fiz. Khim. **41**, 1477 (1967). Russ. J. Phys. Chem. 1967, **41**, 783 (1967).
- 1967MUL/MUK F. I. Mullayanov, R. K. Mukhutdinov, M. F. Mazitov, M. F. Buz'ko, V. P. Borschenko, and V. N. Krashennikov, Khim. Prom. (Moscow) **43**, 505 (1967).
- 1967MUR/KIV J. Murto and A. Kivinen, Suomen Kemistil. B **40**, 258 (1967).
- 1967NIK/RAB P. N. Nikolaev and I. B. Rabinovich, Zh. Fiz. Khim. **49**, 2191 (1967).
- 1967OMA M. M. Omar, J. Chem. Soc. C 2038 (1967).
- 1967PAC/PLA E. L. Pace and A. C. Plaush, *J. Chem. Phys.* **47**, 38 (1967).
- 1967PAE/KUR V. R. Paetzold, R. Kurze, and G. Engelhardt, *Z. Anorg. Allg. Chem.* **353**, 62 (1967).
- 1967PAW S. Pawlenko, *Chem. Ber.* **100**, 3591 (1967).
- 1967PIH/HEI K. Pihlaja and J. Heikkila, *Acta Chem. Scand.* **21**, 2430 (1967).
- 1967PIN/WIL P. E. Pincock, K. R. Wilson, and T. E. Kiovisky, *J. Am. Chem. Soc.* **89**, 6890 (1967).
- 1967PLA/PAC A. C. Plaush and E. L. Pace, *J. Chem. Phys.* **47**, 44 (1967).
- 1967REE/URR D. Reedy and G. Urry, *Inorg. Chem.* **6**, 2117 (1967).
- 1967RIB/WES A. Ribner and E. F. Westrum, Jr., *J. Phys. Chem.* **71**, 1208 (1967).
- 1967SCO/BER D. W. Scott, W. T. Berg, I. A. Hossenlopp, W. N. Hubbard, J. F. Messerly, S. S. Todd, D. R. Douslin, J. P. McCullough, and G. Waddington, *J. Phys. Chem.* **71**, 2263 (1967).
- 1967SHA R. L. Shak, *J. Chem. Eng. Data* **12**, 474 (1967).
- 1967SIM/KNO M. Simon, C. M. Knobler, and A. G. Duncan, *Cryogenics* **7**, 138 (1967).
- 1967SIV/MAT M. Sivokova, A. Matejicek, and F. Juracka, Chem. Prum. **17**, 213 (1967).
- 1967SMI/GOO N. K. Smith and W. D. Good, *J. Chem. Eng. Data* **12**, 570 (1967).
- 1967TUR A. G. Turnbull, Aust. J. Chem. **20**, 2757 (1967).
- 1967TUR2 C. Turrión, Presiones de vapor y calores de sublimación de los ácidos benzenecarboxílicos y sus ésteres metílicos. Beca "Juan March" de estudios en España, grupo B. 1967, Madrid.
- 1967VAN W. A. Van Hook, *J. Chem. Phys.* **46**, 1907 (1967).
- 1967VAN/SOC H. C. Van Ness, C. A. Soczek, G. L. Peloquin, and R. L. Machado, *J. Chem. Eng. Data* **12**, 217 (1967).
- 1967WAK/INO N. Wakayama and H. Inokuchi, *Bull. Chem. Soc. Jpn.* **40**, 2267 (1967); Chem. Abstr. **68**, 38935a (1968).
- 1967WES/RIB E. F. Westrum and A. Ribner, *J. Phys. Chem.* **71**, 1216 (1967).
- 1967WIL/WAR J. M. Wilson and I. J. Worrall, J. Chem. Soc. A 392 (1967).
- 1967ZMB/MAR K. F. Zmbov and J. L. Margrave, *J. Phys. Chem.* **71**, 2893 (1967).
- 1968ADA/SUG K. Adachi, H. Suga, and S. Seki, *Bull. Chem. Soc. Jpn.* **41**, 1073 (1968).
- 1968AMB D. Ambrose, J. Chem. Soc. 1381 (1968).
- 1968AND/COU R. J. L. Andon, J. F. Councell, J. L. Hales, E. B. Lees, and J. F. Martin, J. Chem. Soc. A 2357 (1968).
- 1968BAC/NOV D. P. Baccanari, J. A. Novinski, Y.-C. Pan, M. M. Yevitz, and H. A. Swain, *Trans. Faraday Soc.* **64**, 1201 (1968).
- 1968BAE A. K. Baev, Vestsi Akad. Navak Belarus. SSR, Ser. Khim. Navuk, 76 (1968); Chem. Abstr. **70**, 61310t (1969).
- 1968BAR/CAR E. E. Baroody, G. A. Carpenter, R. A. Robb, and M. F. Zimmer, *J. Chem. Eng. Data* **13**, 215 (1968).
- 1968BUR/IAC A. B. Burg and B. Iachia, *Inorg. Chem.* **7**, 1670 (1968).
- 1968CHE/TUR E. L. Cherkasskaya, A. M. Tur, Z. F. Petrenkova, and V. I. Lyubomilov, Educ. Psychol. Meas. **41**, 2553 (1968).
- 1968CIH/VOJ M. Cihova, J. Vojtko, and M. Hrusovsky, Zb. Pr. Chemickotechnol. Fak. SVST (Slov. Vys. SK. Tech.), 135 (1968); Chem. Abstr. **72**, 136676u (1970).
- 1968CLA/PRI W. D. Clarke and S. J. W. Price, *Can. J. Chem.* **46**, 1633 (1968).
- 1968COU/HAL J. F. Councell, J. L. Hales, E. B. Lees, and J. F. Martin, J. Chem. Soc. A 2994 (1968).
- 1968COU/LEE J. F. Councell, E. B. Lees, and J. F. Martin, J. Chem. Soc. A 1819 (1968).
- 1968DES/WIL P. D. Desai, R. C. Wilhoit, and B. J. Zwolinski, *J. Chem. Eng. Data* **13**, 334 (1968).
- 1968DUT/ONY W. A. Dutton and M. Onyszczuk, *Inorg. Chem.* **7**, 1735 (1968).
- 1968FLO W. Z. Florian, Phys. Chem. [Frankfurt am Main] **61**, 319 (1968); Chem. Abstr. **70**, 81136u (1969).
- 1968FOW/TRU L. Fowler, W. N. Trump, and C. E. Vogler, *J. Chem. Eng. Data* **13**, 209 (1968).
- 1968FOX/MAC W. B. Fox, J. S. MacKenzie, E. R. McCarthy, J. R. Holmes, R. F. Stahl, and R. Juurik, *Inorg. Chem.* **7**, 2064 (1968).
- 1968FRE/TIP J. Freear and A. E. Tipping, J. Chem. Soc. C 1096 (1968).
- 1968GAW/SWI W. J. Gaw and F. L. Swinton, *Trans. Faraday Soc.* **64**, 637 (1968).
- 1968GAW/SWI2 W. J. Gaw and F. L. Swinton, *Trans. Faraday Soc.* **64**,

- 2023 (1968).
- 1968GEI/HOF G. Geiseler and J. Hoffmann, *Z. Phys. Chem. [Frankfurt am Main]* **57**, 318 (1968).
- 1968GLE/VON O. Glemser and S. P. von Halasz, *Z. Naturforsch. B* **23B**, 743 (1968).
- 1968GLE/VON2 O. Glemser, S. P. von Halasz, and U. Biermann, *Z. Naturforsch. B* **23B**, 1381 (1968).
- 1968GOP/RIZ F. Gopal and S. A. Rizvi, *J. Indian Chem. Soc.* **45**, 13 (1968).
- 1968GOU/WES P. Goursot and E. F. Westrum, Jr., *C. R. Acad. Sci. Paris, Ser. C* **266**, 667 (1968).
- 1968GOU/WES2 P. Goursot and E. F. Westrum, Jr., *C. R. Acad. Sci. Paris Ser. C* **266**, 590 (1968).
- 1968HAM/FAG J. V. Hamilton and T. F. Fagley, *J. Chem. Eng. Data* **13**, 523 (1968).
- 1968HAS/TIP R. N. Haszeldine and A. E. Tipping, *J. Chem. Soc. C* **398** (1968).
- 1968HIL/IRV J. O. Hill, and R. J. Irving, *J. Chem. Soc. A* 3116 (1968).
- 1968HIL/IRV2 J. O. Hill, and R. J. Irving, *J. Chem. Soc. A* 1052 (1968).
- 1968KAC/NEM R. V. Kachalova and M. S. Nemtsov, *Zh. Prikl. Khim. (Leningrad)* **41**, 2315 (1968).
- 1968KAR/RAB N. V. Karyakin, I. B. Rabinovich, and L. G. Pakhomov, *Russ. J. Phys. Chem.* **42**, 954 (1968).
- 1968KHO/BYK A. S. Khorevskaya and S. S. Byk, *Zh. Prikl. Khim. (Leningrad)* **41**, 2566 (1968); *Chem. Abstr.* **70**, 61317a (1969).
- 1968KIS/DIL R. W. Kiser, J. G. Dillard, and D. L. Dugger, *Adv. Chem. Ser.* **72**, 153 (1968).
- 1968KUA/UST B. M. Kuadzhe, G. P. Ustyugov, and A. A. Kudryavtsev, *Tr. Mosk. Khim. Tekhnol. Inst.* **58**, 27 (1968); *Chem. Abstr.* **71**, 6728 (1969).
- 1968KUS/MIS V. V. Kushchenko and K. P. Mishchenko, *Educ. Psychol. Meas.* **41**, 646 (1968); *Chem. Abstr.* **68**, 117314a (1968).
- 1968LAP/NIS I. I. Lapidus and L. A. Nisel'son, *Russ. J. Phys. Chem.* **42**, 733 (1968).
- 1968LUC/LEW C. A. Lucchesi and W. T. Lewis, *J. Chem. Eng. Data* **13**, 389 (1968).
- 1968MAK Y. Y. Maksimov, *J. Phys. Chem.* **42**, 1550 (1968).
- 1968MAL/MEU P. F. Malbrunot, P. A. Meunier, G. M. Scatena, W. H. Mears, K. P. Murphy, and J. V. Sinka, *J. Chem. Eng. Data* **13**, 16 (1968).
- 1968MAN/NAK M. Mansson, Y. Nakase, and S. Sunner, *Acta Chem. Scand.* **22**, 171 (1968).
- 1968MAR/ARM S. Marantz and G. T. Armstrong, *J. Chem. Eng. Data* **13**, 118 (1968).
- 1968MAS/RAE R. Mason and A. I. M. Rae, *Proc. R. Soc. London, Ser. A* **304**, 501 (1968).
- 1968MER I. Mertl, *Chem. Listy* **62**, 584 (1968).
- 1968MOR E. Morawetz, *Acta Chem. Scand.* **22**, 1509 (1968).
- 1968NAS/BAB L. L. Nash, D. P. Babb, J. J. Couville, and J. M. Shreeve, *J. Inorg. Nucl. Chem.* **30**, 3373 (1968).
- 1968NIS/NIK L. A. Nisel'son, R. K. Nikolaev, T. D. Sokolova, V. I. Stolyarov, and Y. M. Korolev, *Izv. Sib. Otd. Akad. Nauk SSSR Ser. Khim. Nauk*, 109 (1968); L. A. Nisel'son, R. K. Nikolaev, T. D. Sokolova, V. I. Stolyarov, and Y. M. Korolev, *Chem. Abstr.* **69**, 69834s (1969).
- 1968OSB/DOU A. G. Osborn and D. R. Douslin, *J. Chem. Eng. Data* **13**, 534 (1968).
- 1968PAC/REN E. L. Pace and M. A. Reno, *J. Chem. Phys.* **48**, 1231 (1968).
- 1968PEP/MIR V. I. Pepekin, E. A. Miroshnichenko, Y. A. Levedev, and A. Y. Apin, *J. Chem. Phys.* **42**, 1583 (1968).
- 1968PLA/WIL R. D. Place and S. M. Williamson, *J. Am. Chem. Soc.* **90**, 2550 (1968).
- 1968RAT/SHR C. Ratcliffe and J. M. Shreeve, *J. Am. Chem. Soc.* **90**, 5403 (1968).
- 1968RAV/DAN A. A. Ravdel and V. V. Danilov, *Izv. Vyssh. Uchebn. Zaved. Khim. Khim. Tekhnol.* **11**, 642 (1968); *Chem. Abstr.* **69**, 90232.
- 1968SIN/BUR R. A. Sinclair and A. B. Burg, *Inorg. Chem.* **7**, 2160 (1968).
- 1968TSU/KOJ S. Tsuruoka and H. Kojima, *Sen'i Gakkaishi* **24**, 27 (1968).
- 1968VAN/HOE H. Van Bekkum, M. A. Hoefnagel, L. de Lavieter, A. Van Veen, P. E. Verkade, A. Wemmers, B. M. Wepster, J. H. Palm, L. Schafer, H. Dekker, C. Mosselman, and G. Sommen, *Recl. Trav. Chim. Pays-Bas* **87**, 1363 (1968).
- 1968WAD I. Wadso, *Acta Chem. Scand.* **22**, 2438 (1968).
- 1968WEL/WUL N. Welcman and M. Wulf, *Isr. J. Chem.* **6**, 37 (1968).
- 1968ZHU/KON E. Z. Zhuravlev and I. I. Konstantinov, *Educ. Psychol. Meas.* **41**, 1170 (1968).
- 1969ADA/CAR G. P. Adams, A. S. Carson, and P. G. Laye, *Trans. Faraday Soc.* **65**, 113 (1969).
- 1969ADA/CAR2 G. P. Adams, A. S. Carson, and P. G. Laye, *J. Chem. Thermodyn.* **1**, 393 (1969).
- 1969AGA/HAJ M. Y. Agarunov and S. N. Hajiev, *Dokl. Akad. Nauk SSSR* **185**, 221 (1969).
- 1969AND/BRA D. N. Andreevskii, M. M. Brazhnikov, and V. N. Il'nchik, *Vestn. Beloruss. Univ.*, 3 (1969); *Chem. Abstr.* **73**, 113607 (1969).
- 1969AND/WES J. T. S. Andrews and E. F. Westrum, Jr., *J. Organomet. Chem.* **17**, 349 (1969).
- 1969AND/WES2 J. T. S. Andrews, E. F. Westrum, Jr., and N. Bjerrum, *J. Organomet. Chem.* **17**, 293 (1969).
- 1969AYL/CAM B. J. Aylett and J. M. Campbell, *J. Chem. Soc. A* 1910 (1969).
- 1969AYL/CAM2 B. J. Aylett and J. M. Campbell, *J. Chem. Soc. A* 1916 (1969).
- 1969AYL/CAM3 B. J. Aylett, J. M. Campbell, and A. Walton, *J. Chem. Soc. A* 2110 (1969).
- 1969AYL/HAK B. J. Aylett and M. J. Hakim, *J. Chem. Soc. A* 800 (1969).
- 1969AYL/HAK2 B. J. Aylett and M. J. Hakim, *J. Chem. Soc. A* 636 (1969).
- 1969AYL/HAK3 B. J. Aylett and M. J. Hakim, *J. Chem. Soc. A* 639 (1969).
- 1969BOR/DAL G. Borgen and J. Dale, *Chem. Commun.* 447 (1969).
- 1969BRE/HAG W. Bremser, R. Hagen, E. Heilbronner, and E. Vogel, *Helv. Chim. Acta* **52**, 418 (1969).
- 1969BRO/FOC I. Brown, W. Fock, and F. Smith, *J. Chem. Thermodyn.* **1**, 273 (1969).
- 1969CAL/VAL G. C. Calero, M. M. Valle, and C. G. Losa, *Rev. Acad. Cienc. Fis. Quim. Nat. Zaragoza* **24**, 137 (1969).
- 1969CHA/STE R. Chastel, F. Steckel, and H. Tachoire, *Proceedings of the First International Conference on Calorimetry, Thermodynamics Polish Scientific, Warszawa, 1969*; as cited in [1982COL/JIM].
- 1969CID/POL J. Cidlinsky and J. Polak, *Collect. Czech. Chem. Commun.* **34**, 1317 (1969).
- 1969COR/FRA P. Corradini, A. Frasci, and E. Martuscelli, *J. Chem. Soc. D* 778 (1969).
- 1969COX/GUN J. D. Cox, H. A. Gundry, D. Harrop, and A. J. Head, *J. Chem. Thermodyn.* **1**, 77 (1969).
- 1969CRI F. T. Crimmons, *U.S. Atomic Comm. UCRL-50704* (1969); *Chem. Abstr.* **72**, 70876a (1970).
- 1969DAN/MAT S. M. Danov, V. B. Matin, P. V. Efremov, and N. K. Slashtshina, *Zh. Fiz. Khim.* **43**, 733 (1969); *Chem. Abstr.* **71**, 6729g (1969).
- 1969DAS/WEN G. D'Ascenzo and W. W. Wendlandt, *J. Therm. Anal.* **1**, 423 (1969).
- 1969DAV/MAK Z. V. Davydova, I. A. Makolkin, and P. S. Bataev, *Zh. Obshch. Khim.* **39**, 1668 (1969).
- 1969DAV/MAK2 Z. V. Davydova, I. A. Makolkin, and P. S. Bataev, *Russ. J. Gen. Chem.* **39**, 1636 (1969).
- 1969DEV/ONE J. A. Devore and H. E. O'Neal, *J. Phys. Chem.* **73**, 2644 (1969).
- 1969DIL/KIS J. G. Dillard and R. W. Kiser, *J. Organomet. Chem.* **16**, 265 (1969).
- 1969FIN T. J. V. Findlay, *J. Chem. Eng. Data* **14**, 229 (1969).
- 1969FRA/WAT F. Franks and B. Watson, *Trans. Faraday Soc.* **65**, 2339 (1969).
- 1969FRE/TIP J. Freear and A. E. Tipping, *J. Chem. Soc. C* 411 (1969).
- 1969GEO/MAC R. D. George and K. M. Mackay, *J. Chem. Soc. A* 2122 (1969).
- 1969GLE/MEW O. Glemser, R. Mews, and H. W. Roesky, *J. Chem. Soc., Chem. Commun.* 914 (1969).
- 1969GLE/VON O. Glemser and S. P. Von Halasz, *Inorg. Nucl. Chem. Lett.* **5**, 393 (1969).

- 1969HAM C. Z. Hamann, *Chem.* **9**, 195 (1969); *Chem. Abstr.* **71**, 24938d (1969).
- 1969JOE/GJA O. Joens and J. Gjaldback, *Chr. Dan. Tidsskr. Farm.* **43**, 151 (1969); *Chem. Abstr.* **72**, 6776r (1969).
- 1969KEI/KAN D. Keiser and A. S. Kana'an, *J. Phys. Chem.* **73**, 4264 (1969).
- 1969KEM/KRE R. H. Kemme and S. I. Kreps, *J. Chem. Eng. Data* **14**, 98 (1969).
- 1969KON/PRO J. Konicek, M. Prochazka, V. Krestanova, and M. Smisek, *Collect. Czech. Chem. Commun.* **34**, 2249 (1969).
- 1969KRE/WOO W. Kreis and R. H. Wood, *J. Chem. Thermodyn.* **1**, 523 (1969).
- 1969KRO N. G. Krokhin, *Russ. J. Phys. Chem.* **43**, 1342 (1969).
- 1969LAM/PER R. Lamartine and R. Perrin, *Bull. Soc. Chim. Fr.*, 443 (1969).
- 1969LEA/LON D. R. Leavers, J. R. Long, S. G. Shore, and W. J. Taylor, *J. Chem. Soc.* 1580 (1969).
- 1969LEB/DNE N. D. Lebedeva, A. S. Dneprovskii, and Y. A. Katin, *Russ. J. Phys. Chem.* **43**, 770 (1969).
- 1969LES/MOR T. M. Lesteva, A. I. Morozova, V. I. Morozova, S. K. Ogorodnikov, and T. M. Tyvina, *Zh. Prikl. Khim. (S.-Peterburg)* **42**, 533 (1969).
- 1969MAC/MCN H. Mackle, D. V. McNally, and W. V. Steele, *Trans. Faraday Soc.* **65**, 2060 (1969).
- 1969MAC/MCN H. Mackle and D. V. McNally, *Trans. Faraday Soc.* **65**, 1738 (1969).
- 1969MAC/STE H. Mackle and W. V. Steele, *Trans. Faraday Soc.* **65**, 2069 (1969); see also *Trans. Faraday Soc.* **65**, 2073 (1969).
- 1969MAN M. Mansson, *J. Chem. Thermodyn.* **1**, 141 (1969).
- 1969MAN/MOR M. Mansson, E. Morawetz, Y. Nakase, and S. Sunner, *Acta Chem. Scand.* **23**, 56 (1969).
- 1969MAS J. Mason, *J. Chem. Soc. A* 1587 (1969).
- 1969MEL/MER T. Melia and R. Merrifield, *J. Appl. Chem.* **19**, 79 (1969).
- 1969MES S. Meszaros, *Period. Polytech., Chem. Eng.* **13**, 79 (1969); *Chem. Abstr.* **72**, 66308z (1970).
- 1969MIR/LEB E. A. Miroshnichenko and Y. A. Lebedev, *Khim. Geterotsikl. Soedin.* **5**, 963 (1969).
- 1969OSB/DOU A. G. Osborn and D. R. Doustin, *J. Chem. Eng. Data* **14**, 208 (1969).
- 1969OVE/STE J. E. Overberger, W. A. Steele, and J. G. Aston, *J. Chem. Thermodyn.* **1**, 535 (1969).
- 1969PLA/GLA C. Plato and A. R. Glasgow, Jr., *Anal. Chem.* **41**, 330 (1969).
- 1969ROB/SCO P. M. Robinson and H. G. Scott, *Mol. Cryst. Liq. Cryst.* **5**, 387 (1969).
- 1969ROS J. M. Rosen, *Microscope* **117**, 141 (1969).
- 1969ROS/DIC J. M. Rosen and C. Dickinson, *J. Chem. Eng. Data* **14**, 120 (1969).
- 1969ROS/HOL J. M. Rosen, J. R. Holden, and H. T. Simmons, *Mol. Cryst. Liq. Cryst.* **6**, 205 (1969).
- 1969RUE B. G. Rueben, *J. Chem. Eng. Data* **14**, 235 (1969).
- 1969SHA/FED Y. K. Shaulov, A. K. Federov, and V. G. Genchel, *Zh. Fiz. Khim.* **185**, 1336 (1969).
- 1969SHE/LAN R. J. Sheehan and S. H. Langer, *J. Chem. Eng. Data* **14**, 248 (1969).
- 1969SHI/MCN C.-F. Shieh, D. McNally, and R. H. Boyd, *Tetrahedron* **25**, 3653 (1969).
- 1969SIC/DUB J. E. Sicre, J. T. Dubois, K. J. Eisentraut, and R. E. Sievers, *J. Am. Chem. Soc.* **91**, 3476 (1969).
- 1969SMI/PAC J. H. Smith and E. L. Pace, *J. Phys. Chem.* **73**, 4232 (1969).
- 1969SOU/GOU M. A. Soulie, P. Goursot, A. Peneloux, and J. Metzger, *J. Chim. Phys. Phys.-Chim. Biol.* **66**, 607 (1969).
- 1969SOU/GOU2 M. A. Soulie, P. Goursot, A. Peneloux, and J. Metzger, *J. Chim. Phys. Phys.-Chim. Biol.* **66**, 603 (1969).
- 1969SPE/CLI W. F. Spencer and M. M. Cliath, *Environ. Sci. Technol.* **3**, 670 (1969).
- 1969STU/WES D. R. Stull, E. F. Westrum, Jr., and G. C. Sinke, *The Chemical Thermodynamics of Organic Compounds* (Wiley, New York, 1969).
- 1969VIC/WAL M. Victoria and J. Walkley, *Trans. Faraday Soc.* **65**, 57 (1969).
- 1969VOJ/CIH J. Vojtko, M. Cihova, and M. Hrusovsky, *Zb. Pr. Chem. Tech. Fak. Svst.* 179 (1969); *Chem. Abstr.* **76**, 76492b (1972).
- 1969WAD I. Wadso, *Acta Chem. Scand.* **23**, 2061 (1969).
- 1969WOO/ADI A. L. Woodman and A. Adicoff, *J. Chem. Eng. Data* **14**, 479 (1969).
- 1969ZMB/HAS K. F. Zmbov, J. W. Hastie, R. Hauge, and J. L. Margrave, in *High Temperature Technology*, Proceedings of the Third IUPAC Conference on High Temperature Technology (Butterworths, London, 1969), p. 345; as cited in [1971ADA/MAR].
- 1970AMB/SPR D. Ambrose and C. H. S. Sprake, *J. Chem. Thermodyn.* **2**, 631 (1970).
- 1970AMB/COU D. Ambrose, J. F. Councell, and A. J. Davenport, *J. Chem. Thermodyn.* **2**, 283 (1970).
- 1970AND/BRA D. N. Andreevskii and M. M. Brazhnikov, *Vestn. Beloruss. Univ.* **2**, 14 (1970); *Chem. Abstr.* **76**, 158651s (1972).
- 1970AND/WES J. T. S. Andrews and E. F. Westrum, *J. Phys. Chem.* **74**, 2170 (1970).
- 1970ASH S. J. Ashcroft, *J. Chem. Soc. A* 1020 (1970).
- 1970BAE A. K. Baev, *Obshch. Prikl. Khim.*, 146 (1970); *Chem. Abstr.* **74**, 91934z (1971).
- 1970BAE/DEM A. K. Baev and V. V. Demyanchuk, *Obshch. Prikl. Khim.* **6**, 65 (1970); *Chem. Abstr.* **74**, 103799 (1971).
- 1970BAR/HAS M. G. Barlow, R. N. Haszeldin, and R. Hubbard, *J. Chem. Soc. C* 1232 (1970).
- 1970BON/CAT D. Bonderman, E. D. Cater, and W. Bennett, *J. Chem. Eng. Data* **15**, 396 (1970).
- 1970BOR L. Borcka, *Acta Pharm. Suec* **7**, 1 (1970).
- 1970BOR/DAL G. Borgen and J. Dale, *J. Chem. Soc., Chem. Commun.* 1340 (1970).
- 1970BUR/KAN A. B. Burg and D.-K. Kang, *J. Am. Chem. Soc.* **92**, 1901 (1970).
- 1970BUR/PAR A. B. Burg and D. M. Parker, *J. Am. Chem. Soc.* **92**, 1898 (1970).
- 1970CAR/CAR A. S. Carson, E. M. Carson, P. G. Laye, J. A. Spencer, and W. Steele, *Trans. Faraday Soc.* **66**, 2459 (1970).
- 1970CAR/LAY A. S. Carson, P. G. Laye, J. A. Spencer, and W. Steele, *J. Chem. Thermodyn.* **2**, 659 (1970).
- 1970CAR/ZIM G. A. Carpenter, M. F. Zimmer, E. E. Baroody, and R. A. Robb, *J. Chem. Eng. Data* **15**, 553 (1970).
- 1970CHA/MCN S.-J. Chang, D. McNally, S. Sharry-Tehrany, M. J. Hickey, and R. H. Boyd, *J. Am. Chem. Soc.* **92**, 3109 (1970).
- 1970COU/FEN J. F. Councell, J. O. Fenwick, and E. B. Lees, *J. Chem. Thermodyn.* **2**, 367 (1970).
- 1970COX/PIL J. D. Cox and G. Pilcher, *Thermochemistry of Organic and Organometallic Compounds* (Academic, London, 1970).
- 1970DAS/WEN G. D'Ascenzo and W. W. Wendlandt, *J. Inorg. Nucl. Chem.* **32**, 2431 (1970).
- 1970DYK J. Dykyj, *J. Petrochemica* **10**, 51 (1970).
- 1970DYK/VAN J. Dykyj and A. Vanko, *Petrochemica* **10**, 3 (1970).
- 1970EIS/ORA O. Eisen and A. Orav, *Eesti Tead. Akad. Toim. Keem. Geol.* **19**, 202 (1970).
- 1970FIN/GAR A. Finch, P. J. Gardner, P. McNamara, and G. R. Wellum, *J. Chem. Soc. A* 3339 (1970).
- 1970GAL/MAR G. L. Gal'chenko, L. N. Martynovskaya, V. I. Stanko, and A. I. Klimova, *Dokl. Akad. Nauk SSSR* **193**, 483 (1970).
- 1970GEN/DUV B. Genot and X. Duval, *J. Chim. Phys. Phys.-Chim. Biol.* **67**, 1332 (1970).
- 1970GOE/BLO H. J. Goetze, K. Bloss, and H. Molketin, *Z. Phys. Chem. [Frankfurt am Main]* **73**, 314 (1970).
- 1970GON/KAR A. K. Goncharov, M. K. Karapet'yants, and G. M. Kol'yakova, *Russ. J. Phys. Chem.* **44**, 466 (1970).
- 1970GOO/MOR W. D. Good and R. T. Moore, *J. Chem. Eng. Data* **15**, 150 (1970).
- 1970GUA/SAR G. Guarini and P. Sarti-Fantoni, *Mol. Cryst. Liq. Cryst.* **6**, 423 (1970).
- 1970HAR/HEA D. Harrop, A. J. Head, and G. B. Lewis, *J. Chem. Thermodyn.* **2**, 203 (1970).
- 1970HAY J. N. Hay, *J. Polym. Sci., Part B: Polym. Lett.* **8**, 395 (1970).
- 1970HES/PER B. Hessett and P. G. Perkins, *J. Chem. Soc. A* 3229 (1970).

- 1970HOW/WADP. B. Howard and I. Wadso, *Acta Chem. Scand.* **24**, 145 (1970).
- 1970IRV/WAD R. J. Irving and I. Wadso, *Acta Chem. Scand.* **24**, 589 (1970).
- 1970KOI/OUC E. Koizumi and S. Ouchi, *Nippon Kagaku Kaishi* **91**, 501 (1970); *Chem. Abstr.* **74**, 46115x (1971).
- 1970KOJ H. Kojima, *Sen'i Gakkaishi* **26**, 530 (1970).
- 1970KON/WAD J. Konicek and I. Wadso, *Acta Chem. Scand.* **24**, 2612 (1970).
- 1970KUS/WAD K. Kusano and I. Wadso, *Acta Chem. Scand.* **24**, 2037 (1970).
- 1970LEN/VEL C. Lenchitz and R. Velicky, *J. Chem. Eng. Data* **15**, 401 (1970).
- 1970LIN/SIL D. C.-K. Lin, I. H. Silberberg, and J. J. McKetta, *J. Chem. Eng. Data* **15**, 483 (1970).
- 1970MAN/RAP M. Mansson, N. Rapport, and E. F. Westrum, Jr., *J. Am. Chem. Soc.* **92**, 7296 (1970).
- 1970MAY/VEN J. N. Maycock and V. R. P. Venecker, *Thermochim. Acta* **1**, 191 (1970).
- 1970MEL/MER T. P. Melia and R. Merrifield, *J. Inorg. Nucl. Chem.* **32**, 1489 (1970).
- 1970MEL/MER2T. P. Melia and R. Merrifield, *J. Inorg. Nucl. Chem.* **32**, 2573 (1970).
- 1970MER/BRE E. Murrill and L. Breed, *Thermochim. Acta* **1**, 232 (1970).
- 1970MOI/ANT V. D. Moisev and N. D. Antonova, *Zh. Fiz. Khim.* **44**, 2912 (1970); *Chem. Abstr.* **69**, 80487e (1971).
- 1970MUL/GAL F. I. Mullayanov, G. V. Galagan, M. F. Buz'ko, Y. V. Churkin, and A. A. Shiryaela, *Tr. Nauch. Issled. Inst. Neftekhim. Proizvod.*, 95 (1970); *Chem. Abstr.* **74**, 91930v (1971).
- 1970POL/MUR J. Polak, S. Murakami, V. T. Lam, and G. C. Benson, *J. Chem. Eng. Data* **15**, 323 (1970).
- 1970POL/PER M. N. Polteva, V. P. Persiantseva, and I. L. Rozenfel'd, *Corrosion and Protection of Metals* (Nauka, Moscow, 1970), p. 140; as cited in [1972ROZ/POL].
- 1970PRO/KRE M. Prochazka, V. Krestanova, J. Konicek, and M. Smisek, *Collect. Czech. Chem. Commun.* **35**, 727 (1970).
- 1970QUI/HOF K. Quitzsch, H. P. Hofmann, D. Hering, R. Salzer, and G. Geiseler, *Z. Phys. Chem. (Leipzig)* **243**, 321 (1970).
- 1970SCO D. W. Scott, *J. Chem. Thermodyn.* **2**, 833 (1970).
- 1970SEL P. Sellers, *Acta Chem. Scand.* **24**, 2453 (1970).
- 1970SEL2 P. Sellers, *J. Chem. Thermodyn.* **2**, 211 (1970).
- 1970SHA/FED Y. K. Shaulov, A. K. Federov, G. Y. Zueva, G. V. Bori-syuk, and V. G. Genchen, *Russ. J. Phys. Chem.* **44**, 1181 (1970); see also *Zh. Fiz. Khim.* **44**, 2081 (1970); *Chem. Abstr.* **74**, 7230m (1971).
- 1970SHA/MCG F. Shafizadeh, G. D. McGinnis, R. A. Susott, and C. W. Philpot, *Carbohydr. Res.* **15**, 165 (1970).
- 1970SMI/CAD J. E. Smith and G. H. Cady, *Inorg. Chem.* **9**, 1293 (1970).
- 1970SMI/IRG M. R. Smith, K. J. Irgolic, E. A. Meyers, and R. A. Zingaro, *Thermochim. Acta* **1**, 51 (1970).
- 1970SMI/THO E. D. Smith and W. L. Thornsberry, *J. Chem. Eng. Data* **15**, 296 (1970).
- 1970SOK/KAR V. B. Sokolov, M. K. Karapet'yants, and A. K. Goncharov, *Tr. Mosk. Khim. Tekhnol. Inst.* **67**, 37 (1970); *Chem. Abstr.* **75**, 122149z (1971).
- 1970SPE/CLI W. F. Spencer and M. M. Cliath, *J. Agric. Food Chem.* **18**, 529 (1970).
- 1970SWA/KAR H. A. Swain, Jr. and D. G. Karraker, *Inorg. Chem.* **9**, 1766 (1970).
- 1970TRA/LOM A. A. Trapeznikov and T. A. Lomonosova, *Zh. Fiz. Khim.* **44**, 2121 (1970).
- 1970VAL A. J. Valerga, *Dissertation Abstr.* **31**, 3316-B (1970).
- 1970VAL/KIL A. J. Valerja and J. E. Kilpatrick, *J. Chem. Phys.* **52**, 4545 (1970).
- 1970VAR/BEL R. M. Varushchenko, N. A. Belikova, S. M. Skuratov, and A. F. Plate, *Zh. Fiz. Khim.* **44**, 3022 (1970).
- 1970VOJ/CIH J. Vojtko, M. Cihova, and M. Hrusovsky, *Chem. Zvesti* **24**, 173 (1970).
- 1970VON/GLE S. P. Von Halasz and O. Glemser, *Chem. Ber.* **103**, 553 (1970).
- 1970VON/WIL P. R. von Schleyer, J. E. Williams, and K. R. Blanchard, *J. Am. Chem. Soc.* **92**, 2377 (1970).
- 1970WES/WON E. F. Westrum, Jr., W. E. Wong, and E. Morawetz, *J. Phys. Chem.* **74**, 2542 (1970).
- 1970WIE/WAU A. G. Wiedemann and H. P. Waughna, *Proceedings of the Third Toronto Symposium on Thermal Analysis, 1970* (unpublished), p. 233 (as quoted in [1999ZIE/PER]).
- 1970ZAB/SHR L. M. Zaborowski and J. M. Shreeve, *J. Am. Chem. Soc.* **92**, 3665 (1970).
- 1971ADA/MAR G. P. Adams, J. L. Margrave, R. P. Steiger, and P. W. Wilson, *J. Chem. Thermodyn.* **3**, 297 (1971).
- 1971ALV/DAL T. Alvik and J. Dale, *Acta Chem. Scand.* **25**, 2131 (1971).
- 1971AND/CON R. J. L. Andon, J. E. Connett, J. F. Councell, E. B. Lees, and J. F. Martin, *J. Chem. Soc. A* 661 (1971).
- 1971ASH S. J. Ashcroft, *Thermochim. Acta* **2**, 512 (1971).
- 1971BAE/DEM A. K. Baev, V. V. Dem'yanchuk, G. Mirzoev, G. I. Noikov, and N. E. Kolobova, *Russ. J. Phys. Chem.* **45**, 777 (1971) [see *Chem. Abstr.* **75**, 80991k (1971)] as cited by D. S. L. Brown, J. A. Connor, and H. A. Skinner, *J. Organomet. Chem.* **81**, 403 (1974).
- 1971BAY/LET J. W. Bayles and T. M. Letcher, *J. Chem. Eng. Data* **16**, 266 (1971).
- 1971BEE/LIN G. Beech and R. M. Lintonbon, *Thermochim. Acta* **2**, 86 (1971).
- 1971BEE/LIN2 G. Beech and R. M. Lintonbon, *Thermochim. Acta* **3**, 97 (1971).
- 1971BER/GIR G. Bertholon, M. Giray, R. Perrin, and M. F. Vincent-Falquet-Berny, *Bull. Soc. Chim. Fr.*, 3180 (1971).
- 1971BOY/SAN R. H. Boyd, S. N. Sanwal, S. Shary-Tehrany, and D. McNally, *J. Phys. Chem.* **75**, 1264 (1971).
- 1971BUT/CAR R. S. Butler, A. S. Carson, P. G. Laye, and W. V. Steele, *J. Chem. Thermodyn.* **3**, 277 (1971).
- 1971CAL/SMI J. E. Callanan and N. O. Smith, *J. Chem. Thermodyn.* **3**, 531 (1971).
- 1971CAR/FIN A. S. Carson, D. H. Fine, P. Gray, and P. G. Laye, *J. Chem. Soc. B*, 1611 (1971).
- 1971CAR/LAY A. S. Carson, P. G. Laye, W. V. Steele, D. E. Johnston, and M. A. McKervey, *J. Chem. Thermodyn.* **3**, 915 (1971).
- 1971CHO/JON J. K. Choi and M. J. Joncich, *J. Chem. Eng. Data* **16**, 87 (1971).
- 1971COU/LEE J. F. Councell, D. A. Lee, and J. F. Martin, *J. Chem. Soc. A* 313 (1971).
- 1971DEM/SHR R. A. De Marco and J. M. Shreeve, *Inorg. Chem.* **10**, 911 (1971).
- 1971DIN/STA R. H. Dinegar and M. Stammier, *Explosivstoffe* **1**, 14 (1971).
- 1971DIT/SKO V. E. Ditsent, I. I. Skorokhodov, N. A. Terent'eva, and M. N. Zolotareva, *Russ. J. Phys. Chem.* **45**, 901 (1971).
- 1971DYK J. Dykyj, *Petrochemia* **11**, 27 (1971).
- 1971EDW/DOL L. Edwards, D. H. Dolphin, M. Gouterman, and A. D. Adler, *J. Mol. Spectrosc.* **38**, 16 (1971).
- 1971EON/POM C. Eon, C. Pommier, and G. Guiochon, *J. Chem. Eng. Data* **16**, 408 (1971).
- 1971FEL/KUZ L. S. Fel'dshtein and A. S. Kuz'minskii, *Kauch. Rezina* **30**, 15 (1971).
- 1971GEL/SIM Z. I. Geller, V. D. Simonov, M. V. Lyubarskii, E. A. Bra-khfogel, L. M. Meizler, and I. A. Parmonov, *Izv. Vyssh. Uchebn. Zaved. Nept. Gaz* **7**, 71 (1971).
- 1971GON/KAR A. K. Goncharov, M. K. Karapet'yants, and T. K. Gar, *Russ. J. Phys. Chem.* **45**, 104 (1971).
- 1971GOO/MOO W. D. Good and R. T. Moore, *J. Chem. Thermodyn.* **3**, 701 (1971).
- 1971HAL/BAL H. K. Hall, Jr. and J. H. Baldt, *J. Am. Chem. Soc.* **93**, 140 (1971).
- 1971HAM/WIT W. S. Hamilton and L. C. Witt, *J. Chem. Eng. Data* **16**, 234 (1971).
- 1971HAU H. O. Haug, *Ges. Kernforsch. m.b.H. [Rep.] KFK-1283*, 23 (1971); *Chem. Abstr.* **75**, 122126q (1971).
- 1971HAU2 H. O. Haug, *J. Organomet. Chem.* **30**, 53 (1971); *Chem. Abstr.* **75**, 122161x (1971).
- 1971JAC/HUN W. Jackson, T. S. Hung, and H. P. Hopkins, Jr., *J. Chem. Thermodyn.* **3**, 347 (1971).
- 1971KAM/MIT H. Kambe, I. Mita, and R. Yokata, *Proceedings of the*

- Third ICTAC, Davos Swizz, 1971, pp. 387–395 (unpublished); Chem. Abstr. **79**, 5824n (1973).
- 1971KIP/RAB E. G. Kiparisova and I. B. Rabinovitch, Dokl. Akad. Nauk SSSR **199**, 1075 (1971); see also Dokl. Chem. **199**, 675 (1971).
- 1971KOL/RAB G. M. Kol'yakova, I. B. Rabinovich, and N. S. Vyazankin, Dokl. Akad. Nauk SSSR **200**, 111 (1971).
- 1971KUS/WAD K. Kusano and I. Wadso, *Acta Chem. Scand.* **25**, 219 (1971).
- 1971KUS/WAD2K. Kusano and I. Wadso, *Bull. Chem. Soc. Jpn.* **44**, 1705 (1971).
- 1971LEA D. R. Leavers, Ph.D. dissertation, The Ohio State University, 1971.
- 1971LEB/KAT N. D. Lebedeva, Y. A. Katin, and G. Y. Akhmedova, Russ. J. Phys. Chem. **45**, 1192 (1971).
- 1971LEB/KAT2 N. D. Lebedeva, Y. A. Katin, and G. Y. Akhmedova, Russ. J. Phys. Chem. **45**, 771 (1971).
- 1971LEB/RYA N. D. Lebedeva, V. L. Ryadnenko, and I. N. Kuznetsova, Russ. J. Phys. Chem. **45**, 549 (1971).
- 1971LEN/VEL C. Lenchitz, R. W. Velicky, G. Silvestro, and L. P. Schlosberg, *J. Chem. Thermodyn.* **3**, 689 (1971).
- 1971MAN/RIN M. Mansson, B. Ringner, and S. Sunner, *J. Chem. Thermodyn.* **3**, 547 (1971).
- 1971MAT/PEP Y. N. Matyushin, V. I. Pepekin, S. P. Golova, T. I. Godovikova, and L. I. Khmel'nitskii, *Bull. Acad. Sci. USSR Div. Chem. Sci.* **20**, 162 (1971).
- 1971MEY/REN E. F. Meyer, T. A. Renner, and K. S. Stec, *J. Phys. Chem.* **76**, 642 (1971).
- 1971MIT/IMA I. Mita, I. Imai, and I. Kambe, *Thermochim. Acta* **2**, 337 (1971).
- 1971MOR E. Morawetz, *Chem. Scr.* **1**, 103 (1971).
- 1971MOR2 E. Morawetz, Ph.D. dissertation, University of Lund, 1971; see also Ref. [1983DEW/VAN].
- 1971MUR/BRE E. Murrill and L. W. Breed, *Inorg. Chem.* **10**, 641 (1971).
- 1971NAK/SMI I. E. Nakhutin, N. M. Smirnova, V. I. Krivenko, and G. A. Loshakov, Zh. Obshch. Khim. **41**, 940 (1971); Chem. Abstr. **75**, 122154x (1971).
- 1971NIS/TRE L. A. Nisel'son, K. V. Tret'yakova, E. P. Paremuzov, and E. N. Torbina, Izv. Akad. Nauk SSSR, Neorg. Mater. **7**, 792 (1971); Chem. Abstr. **75**, 67746q (1971).
- 1971OSB/SCH D. W. Osborne, F. Schreiner, and H. Selig, *J. Chem. Phys.* **54**, 3790 (1971).
- 1971PAR/ROC G. H. Parsons, C. H. Rochester, and C. E. C. Wood, *J. Chem. Soc. B* **1974**, 533.
- 1971PIH/TUO K. Pihlaja and M.-L. Tuomi, *Acta Chem. Scand.* **25**, 465 (1971).
- 1971PIN/TON R. E. Pincock, M.-M. Tong, and K. R. Wilson, *J. Am. Chem. Soc.* **93**, 1669 (1971).
- 1971POL/BEN J. Polak and G. C. Benson, *J. Chem. Thermodyn.* **3**, 235 (1971).
- 1971PRI V. P. Privalko, Russ. J. Phys. Chem. **45**, 900 (1971).
- 1971RAB/KIP I. B. Rabinovich, E. G. Kiparisov, and Y. A. Aleksandrov, Dokl. Akad. Nauk SSSR **200**, 1116 (1971).
- 1971RAD L. G. Radchenko, Zh. Fiz. Khim. **45**, 1310 (1971); Chem. Abstr. **75**, 122104f (1971).
- 1971RAP/WES N. J. Rapport, E. F. Westrum, Jr., and J. T. S. Andrews, *J. Am. Chem. Soc.* **93**, 4363 (1971).
- 1971RIN/SUN B. Ringner, S. Sunner, and H. Watanabe, *Acta Chem. Scand.* **25**, 141 (1971).
- 1971ROG F. E. Rogers, *J. Phys. Chem.* **75**, 1734 (1971).
- 1971ROS/HOL J. M. Rosen, J. R. Holden, and H. T. Simmons, Sr., *Microscope* **19**, 151 (1971).
- 1971SAU/SHR D. T. Sauer and J. M. Shreeve, *J. Fluorine Chem.* **1**, 1 (1971).
- 1971SAU/SHR2 D. T. Sauer and J. M. Shreeve, *Inorg. Chem.* **10**, 358 (1971).
- 1971SEL P. Sellers, *Acta Chem. Scand.* **25**, 2099 (1971).
- 1971SEL2 P. Sellers, *Acta Chem. Scand.* **25**, 2194 (1971); **25**, 2291 (1971).
- 1971SEL3 P. Sellers, *Acta Chem. Scand.* **25**, 2189 (1971).
- 1971SHA/YAK Y. K. Shaulov and E. S. Yakubov, Zh. Fiz. Khim. **45**, 188 (1971).
- 1971SKO/DIT I. I. Skorokhodov, V. E. Ditsent, N. A. Teret'eva, and M. N. Zolotarev, Russ. J. Phys. Chem. **45**, 902 (1971).
- 1971SOK/KAR V. B. Sokolov, M. K. Karapet'yants, N. D. Rummyantseva, and V. A. Drozdov, Russ. J. Phys. Chem. **45**, 1332 (1971).
- 1971SUN/EIS C. Sunwoo and H. Eisen, *J. Pharm. Sci.* **60**, 238 (1971).
- 1971SWA/KAR H. A. Swain, Jr., and D. G. Karkker, *J. Inorg. Nucl. Chem.* **33**, 2851 (1971).
- 1971SWI/ZAB F. Swindell, L. M. Zaborowski, and J. M. Shreeve, *Inorg. Chem.* **10**, 1635 (1971).
- 1971TEL/RAB V. I. Tel'noi, I. B. Rabinovitch, V. N. Latyaeva, and A. N. Lineva, Dokl. Akad. Nauk SSSR **197**, 353 (1971).
- 1971VOI/SHC S. A. Voitkevich, M. M. Shchedrina, N. P. Solov'eva, and T. A. Rudol'fi, Maslo-Zhir. Prom. **37**, 27 (1971).
- 1971WIL/ZWO R. C. Wilhoit and B. J. Zwolinski, *Handbook of Vapor Pressures and Heats of Vaporization of Hydrocarbons and Related Compounds* (API 44-TRC Publications in Science and Engineering, College Station, TX, 1971).
- 1971WON/WES W.-K. Wong and E. F. Westrum, Jr., *J. Chem. Thermodyn.* **3**, 105 (1971).
- 1971WU/HSU P.-J. Wu, L. Hsu, and D. A. Dows, *J. Chem. Phys.* **54**, 2714 (1971).
- 1971ZAB/SHR L. M. Zaborowski and J. M. Shreeve, *Inorg. Chem.* **10**, 407 (1971).
- 1972AHM/EAD A. M. I. Ahmed and R. G. Eades, *J. Chem. Soc., Faraday Trans. 2* **68**, 1623 (1972).
- 1972AHM/EAD2A. M. I. Ahmed and R. G. Eades, *J. Chem. Soc., Faraday Trans. 2* **68**, 2017 (1972).
- 1972ALV/BOR T. Alvik, G. Borgen, and J. Dale, *Acta Chem. Scand.* **126**, 1805 (1972).
- 1972AMB/SPR D. Ambrose, C. H. S. Sprake, and R. Townsend, *J. Chem. Thermodyn.* **4**, 247 (1972).
- 1972AMI/VAK A. V. Amitin, E. A. Vakurova, V. K. Katunin, and N. S. Afanas'eva, Zh. Fiz. Khim. **46**, 1054 (1972); Chem. Abstr. **77**, 66425p (1972).
- 1972ARN/EBS D. E. J. Arnold, E. A. V. Ebsworth, H. F. Jessep, and D. W. H. Rankin, *J. Chem. Soc. Dalton Trans.* 1681 (1972).
- 1972ARN/JON P. R. Arnold and F. Jones, *Mol. Cryst. Liq. Cryst.* **19**, 133 (1972).
- 1972BLE/FIE R. H. T. Bleyerveld and W. Fiegeen, Recl. Trav. Chim. Pays-Bas **91**, 477 (1972); Chem. Abstr. **76**, 145167p (1972).
- 1972BOL M. W. G. Bolster, Ph.D. thesis, University of Leiden, 1972; as quoted in Ref. [2000DUN].
- 1972BOR/DAL G. Borgen and J. Dale, *Acta Chem. Scand.* **26**, 1149 (1972).
- 1972BOR/DAL2G. Borgen and J. Dale, *Acta Chem. Scand.* **26**, 1799 (1972).
- 1972BOU/AIM T. Boublik and K. Aim, Collect. Czech. Chem. Commun. **37**, 3513 (1972).
- 1972BRA/KAR G. P. Brajin, M. K. Karapet'yants, A. K. Goncharov, and G. M. Kol'yakova, Tr. Mosk. Khim. Tekhnol. Inst. **71**, 10 (1972); Chem. Abstr. **80**, 100458a (1974).
- 1972BUR A. B. Burg, *Inorg. Chem.* **11**, 2283 (1972).
- 1972BUR/REI A. B. Burg and T. J. Reilly, *Inorg. Chem.* **11**, 1962 (1972).
- 1972BYR/CUR S. R. Byrn, D. Y. Curtin, and I. C. Paul, *J. Am. Chem. Soc.* **94**, 890 (1972).
- 1972CAR/LAY A. S. Carson, P. J. Laye, J. A. Spencer, and W. J. Steele, *J. Chem. Thermodyn.* **4**, 783 (1972).
- 1972CHA/BES S. S. Chang, and A. B. Bestul, *J. Chem. Phys.* **56**, 503 (1972).
- 1972COL/LAY M. Colomina, J. L. Laynez, R. Perez-Ossorio, and C. Turrión, *J. Chem. Thermodyn.* **4**, 499 (1972).
- 1972COL/MON M. Colomina, C. Monzon, C. Turrión, and J. Laynez, Paper presented at the Fifth Experimental Thermodynamics Conference, Lancaster (unpublished); as cited in Ref. [1982COL/JIM].
- 1972CON/SKI J. A. Connor, H. A. Skinner, and Y. Virmani, *J. Chem. Soc., Faraday Trans. 1* **68**, 1754 (1972).
- 1972COU/LEE J. F. Cuncell and D. A. Lee, *J. Chem. Thermodyn.* **4**, 915 (1972).
- 1972DAL/KRI J. Dale and P. O. Kristiansen, *Acta Chem. Scand.* **26**, 1471 (1972).

- 1972DEM/SHR R. A. De Marco and J. M. Shreve, *J. Fluorine Chem.* **1**, 269 (1972).
- 1972DIT/SKO V. E. Ditsent, I. I. Skorokhodov, N. A. Terent'eva, and M. N. Zolotareva, *Russ. J. Phys. Chem.* **46**, 317 (1972); see also *Zh. Fiz. Khim.* **46**, 544 (1972); *Chem. Abstr.* **77**, 9812p (1972).
- 1972DIT/SKO2 V. E. Ditsent, I. I. Skorokhodov, N. A. Terent'eva, and M. N. Zolotareva, *Russ. J. Phys. Chem.* **46**, 1088 (1972); see also *Zh. Fiz. Khim.* **46**, 1887 (1972); *Chem. Abstr.* **77**, 106168r (1972); **77**, 106166p (1972).
- 1972DYK J. Dykyj, *Petrochemia* **12**, 13 (1972); **12**, 34 (1972).
- 1972FIN/GAR A. Finch, P. J. Gardner, and A. F. Webb, *J. Chem. Thermodyn.* **4**, 495 (1972).
- 1972FIN/MCC H. L. Finke, J. P. McCullough, J. F. Messerly, A. Osborn, and D. R. Douslin, *J. Chem. Thermodyn.* **4**, 477 (1972).
- 1972FON/POM R. Fontaine, C. Pommier, and G. Guiochon, *Bull. Soc. Chim. Fr.*, 3011 (1972).
- 1972GAR/HUS P. J. Gardner and K. S. Hussain, *J. Chem. Thermodyn.* **4**, 819 (1972).
- 1972GEI/RAU G. Geiseler, and H. J. Rauh, *Z. Phys. Chem. (Leipzig)* **249**, 376 (1972).
- 1972GEI/SAW G. Geiseler and J. Sawistowsky, *Z. Phys. Chem. (Leipzig)* **250**, 43 (1972).
- 1972GON/KAR A. K. Goncharov and M. H. Karapet'yants, *Zh. Fiz. Khim.* **46**, 228 (1972).
- 1972GOO W. D. Good, *J. Chem. Eng. Data* **17**, 158 (1972).
- 1972GRA/SAL V. A. Granzhan and E. A. Salganikov, *Tr. Nauch. Issled. Proekt. Inst. Azotn. Prom. Prod. Org. Sin.*, 85 (1972); *Chem. Abstr.* **79**, 118908m (1973).
- 1972HAM/WIT W. S. Hamilton and L. C. Witt, *J. Chem. Eng. Data* **17**, 138 (1972).
- 1972INO/LIA P. P. Inozemtsev, A. G. Liakumovich, and Z. D. Gracheva, *Russ. J. Phys. Chem.* **46**, 914 (1972).
- 1972IRV R. J. Irving, *J. Chem. Thermodyn.* **4**, 793 (1972).
- 1972JAK/VAN G. Jakli and W. A. Van Hook, *J. Chem. Thermodyn.* **4**, 857 (1972).
- 1972JOH/OHA G. K. Johnson, P. A. G. O'Hare, and E. H. Appelman, *Inorg. Chem.* **11**, 800 (1972).
- 1972KAN A. S. Kana'an, *J. Chem. Thermodyn.* **4**, 893 (1972).
- 1972KOL/RAB G. M. Kol'yakova, I. B. Rabinovich, E. N. Gladyshev, and N. S. Vyazankin, *Dokl. Akad. Nauk SSSR* **204**, 356 (1972).
- 1972KRE/PRI M. Krech, S. J. W. Price, and W. F. Yared, *Can. J. Chem.* **50**, 2935 (1972).
- 1972LAY/WAD J. Laynez and I. Wadso, *Acta Chem. Scand.* **26**, 3148 (1972).
- 1972LEB/KAT N. D. Lebedeva and Y. A. Katin, *Zh. Fiz. Khim.* **46**, 1060 (1972); see also *Russ. J. Phys. Chem.* **46**, 623 (1972).
- 1972LEB/KAT2 N. D. Lebedeva and Y. Katin, *Zh. Fiz. Khim.* **46**, 1088 (1972).
- 1972MAN M. Mansson, *J. Chem. Thermodyn.* **4**, 865 (1972).
- 1972MAN2 M. Mansson, *Acta Chem. Scand.* **26**, 1707 (1972).
- 1972MAR/LOP I. Y. Markova, I. L. Lopatkina, Y. K. Shaulov, and Y. A. Priselkov, *Tr. Khim. Khim. Tekhnol.*, 61 (1972); *Chem. Abstr.* **79**, 832724u (1973).
- 1972MCF/SOM F. R. McFeely and G. A. Somorjee, *J. Phys. Chem.* **76**, 914 (1972).
- 1972MET/SHR S. G. Metcalf and J. M. Shreeve, *Inorg. Chem.* **11**, 1631 (1972).
- 1972MIL G. D. Mills, *Diss. Abstr. Int.*, B **33**, 1485 (1972); *Chem. Abstr.* **78**, 42653f (1973).
- 1972MOR E. Morawetz, *J. Chem. Thermodyn.* **4**, 455 (1972).
- 1972MOR2 E. Morawetz, *J. Chem. Thermodyn.* **4**, 139 (1972).
- 1972MOR3 E. Morawetz, *J. Chem. Thermodyn.* **4**, 461 (1972).
- 1972NAK/TOY K. Nakanishi and O. Toyama, *Bull. Chem. Soc. Jpn.* **45**, 3210 (1972).
- 1972NEE/HAL W. C. Neely and T. D. Hall, *J. Chem. Eng. Data* **17**, 294 (1972).
- 1972NEW H. W. Newkirk, *J. Organomet. Chem.* **44**, 263 (1972).
- 1972NIS/HAK M. Nishimura, M. Hakayama, and T. Yano, *J. Chem. Eng. Jpn.* **5**, 223 (1972); *Chem. Abstr.* **78**, 48659j (1973).
- 1972NIS/SOK L. A. Nisel'son, T. D. Sokolova, and R. K. Nikolaev, *Vestn. Mosk. Univ., Ser. 2: Khim.* **13**, 432 (1972); *Chem. Abstr.* **78**, 8069k (1973).
- 1972ONE/SHR S. R. O'Neill and J. M. Shreeve, *Inorg. Chem.* **11**, 1629 (1972).
- 1972PAC/HOD E. L. Pace and P. E. Hodge, *J. Chem. Thermodyn.* **4**, 441 (1972).
- 1972PED/ISE J. B. Pedley and B. S. Iseard, *CATCH Tables, Silicon Compounds* (University of Sussex, Brighton, 1972).
- 1972PEP/MAT V. I. Pepekin, Y. N. Matyushin, G. G. Rozantsev, S. A. Shevelev, and A. Y. Apin, *Izv. Akad. Nauk SSSR Ser. Khim.* **21**, 2703 (1972); see also *Izv. Akad. Nauk SSSR Ser. Khim. Eng. Edit.* **21**, 2634 (1972).
- 1972PLA C. Plato, *Anal. Chem.* **44**, 1531 (1972).
- 1972PRY/GOO R. Prydz and R. D. Goodwin, *J. Chem. Thermodyn.* **4**, 123 (1972).
- 1972REG J. Regnier, *J. Chim. Phys. Phys.-Chim. Biol.* **69**, 942 (1972).
- 1972ROG2 F. E. Rogers, *J. Phys. Chem.* **76**, 106 (1972).
- 1972ROM/SUK B. C. Romanov, B. I. Sukhorukov, E. A. Miroshnichenko, and Y. A. Lebedev, Fourth International Biophysics Congress Abstracts, 1972 (unpublished), Paper No. E7 (a4/7); as cited in Ref. [2000BRU/PIA].
- 1972ROZ/POL I. L. Rozenfel'd, M. N. Polteva, and V. P. Persiantseva, *Izv. Akad. Nauk SSSR Ser. Khim.* **720** (1972); *Chem. Abstr.* **77**, 66496n (1972). (Note: compound is incorrectly named in the abstract.)
- 1972SAB/CHA R. Sabbah, R. Chastel, and M. Laffitte, *Thermochim. Acta* **5**, 117 (1972).
- 1972SAU/SHR D. T. Sauer and J. M. Shreeve, *Inorg. Chem.* **11**, 238 (1972).
- 1972SOK/BRA V. B. Sokolov, G. P. Bragin, and M. K. Karapet'yants, *Tr. Mosk. Khim. Tekhnol. Inst.* **71**, 8 (1972); *Chem. Abstr.* **80**, 100457z (1974).
- 1972SOK/KAR V. B. Sokolov, M. K. Karapet'yants, S. P. Kolesnikov, B. L. Perl'mutter, and O. M. Nefedov, *Tr. Mosk. Khim. Tekhnol. Inst.* **71**, 6 (1972); *Chem. Abstr.* **80**, 100456y (1974).
- 1972SOL/BUR D. Solan and A. B. Burg, *Inorg. Chem.* **11**, 1253 (1972).
- 1972SPE/CLI W. F. Spencer and M. M. Cliath, *J. Physiol. (London)* **20**, 645 (1972).
- 1972STRE/NOV Y. A. Strepikheev, O. P. Novikova, and A. L. Chimishkyan, *Khim. Prom. (Moscow)* **48**, 230 (1972); *Chem. Abstr.* **69**, 145717t (1972).
- 1972SWI/BAB R. F. Swindell, D. B. Babb, T. J. Ouellette, and J. M. Shreeve, *Inorg. Chem.* **11**, 242 (1972).
- 1972SWI/SHR R. F. Swindell and J. M. Shreeve, *J. Am. Chem. Soc.* **94**, 5713 (1972).
- 1972TEL/RAB V. I. Tel'noi, I. B. Rabinovich, B. I. Kozyrkin, B. A. Salamant, and K. V. Kir'yanov, *Dokl. Akad. Nauk SSSR* **205**, 364 (1972); *Chem. Abstr.* **77**, 131500a (1972).
- 1972UMI/VAN V. A. Umilin and V. K. Vanchagova, *Izv. Akad. Nauk SSSR Ser. Khim.*, 651 (1972).
- 1972URB/IAN S. Urban, J. A. Janik, J. Lenik, J. Mayer, T. Baluga, and S. Wrobel, *Phys. Status Solidi A* **10**, 271 (1972).
- 1972VAR/DRU R. M. Varuschenko and A. I. Druzhinina, *Zh. Fiz. Khim.* **46**, 1313 (1972).
- 1972VOL/SMO E. A. Volchkova, D. D. Smol'yanimova, V. G. Genchel, K. Lapatkina, and Y. K. Shaulov, *Russ. J. Phys. Chem.* **46**, 1053 (1972).
- 1972WIE H. G. Wiedemann, *Thermochim. Acta* **3**, 355 (1972).
- 1972WOL G. Wolf, *Helv. Chim. Acta* **55**, 1446 (1972).
- 1973ABE/SHR T. Abe and J. M. Shreeve, *J. Fluorine Chem.* **3**, 17 (1973/1974).
- 1973AMB/SPR D. Ambrose, C. H. S. Sprake, and R. J. Townsend, *J. Chem. Soc., Faraday Trans. 1* **69**, 839 (1973).
- 1973AND/COU R. J. L. Andon, J. F. Councell, D. A. Lee, and J. F. Martin, *J. Chem. Soc., Faraday Trans. 1* **69**, 1721 (1973).
- 1973AND/MAR R. J. L. Andon and J. F. Martin, *J. Chem. Soc., Faraday Trans. 1* **69**, 761 (1973).
- 1973ARS/SHA M. R. Arshadi and M. Shabrang, *J. Chem. Soc., Perkin Trans. 2* **2**, 1732 (1973).
- 1973AYL/ELL B. J. Aylett, I. A. Ellis, and C. J. Porritt, *J. Chem. Soc. Dalton Trans.* 83 (1973).

- 1973AYL/ELL2 B. J. Aylett, I. A. Ellis, and J. R. Richmond, *J. Chem. Soc. Dalton Trans.* 981 (1973).
- 1973BAR/MAL G. Bardi, L. Malaspina, and V. Piacenti, *J. Chem. Eng. Data* **18**, 126 (1973).
- 1973BAR/MOR D. S. Barnes and C. T. Mortimer, *J. Chem. Thermodyn.* **5**, 371 (1973).
- 1973BAR/MOR2 D. S. Barnes, C. T. Mortimer, and E. Mayer, *J. Chem. Thermodyn.* **5**, 481 (1973).
- 1973BEC/RUC H.-D. Beckhaus and C. Rüchardt, *Tetrahedron Lett.* **14**, 1971 (1973).
- 1973BEH/GAT W. Behrendt and G. Gattow, *Z. Anorg. Allg. Chem.* **398**, 198 (1973).
- 1973BER/DES P. A. Bernstein and D. D. Desmarreau, *J. Fluorine Chem.* **2**, 315 (1973).
- 1973BIE/EIC R. M. Biefeld and H. A. Eick, *J. Chem. Thermodyn.* **5**, 353 (1973).
- 1973BLA/IHL A. Blair and H. Ihle, *J. Inorg. Nucl. Chem.* **35**, 3795 (1973).
- 1973BOR/KRA G. K. Borisov, S. G. Krasnova, and G. G. Devyatykh, *Russ. J. Phys. Chem.* **18**, 346 (1973).
- 1973BRU/CUR H. R. Brunner and B. J. Curtis, *J. Therm. Anal.* **5**, 111 (1973).
- 1973CAM/GAM R. Cameroni, G. Gamberini, M. T. Bernabei, and M. Facchini, *Farmaco* **28**, 621 (1973).
- 1973CAR/KOB G. F. Carruth and R. Kobayashi, *J. Chem. Eng. Data* **18**, 115 (1973).
- 1973CAR/ROB A. Cartner, B. Robinson, and P. J. Gardner, *J. Chem. Soc., Chem. Commun.* 317 (1973).
- 1973CAS/VEC F. Casellato, C. Vecchi, and A. Girell, *Thermochim. Acta* **6**, 361 (1973).
- 1973CON/SKI J. A. Connor, H. A. Skinner, and Y. Virmani, *J. Chem. Soc., Faraday Trans. 1* **69**, 1218 (1973).
- 1973COU/LEE J. F. Councell and D. A. Lee, *J. Chem. Thermodyn.* **5**, 583 (1973).
- 1973DAL/EKE J. Dale and T. Ekeland, *Acta Chem. Scand.* **27**, 1519 (1973).
- 1973DAW/SIL P. D. Dawson, Jr., I. M. Silberberg, and J. J. McKetta, *J. Chem. Eng. Data* **18**, 7 (1973).
- 1973DEK/OON C. G. DeKruif and H. A. Oonk, *J. Chem. Ing. Techn.* **45**, 455 (1973); *Chem. Abstr.* **78**, 152198u (1973).
- 1973DEM/LEH G. R. De Mare, T. Lehman, and M. Termonia, *J. Chem. Thermodyn.* **5**, 829 (1973).
- 1973DEM/SHR R. A. De Marco and J. M. Shreeve, *Inorg. Chem.* **12**, 1896 (1973).
- 1973DEV/BOR G. G. Devyatykh, G. K. Borisov, L. F. Zyuzina, and S. G. Krasnov, *Dokl. Phys. Chem.* **212**, 703 (1973).
- 1973DIE/MAR E. A. Dietz and D. R. Martin, *J. Inorg. Nucl. Chem.* **35**, 3681 (1973).
- 1973DYA/VIG N. N. D'yachkova, E. N. Vigdorovich, and L. A. Ivanyutin, *Russ. J. Phys. Chem.* **47**, 258 (1973).
- 1973FEL/SAV I. N. Fel'dman, V. V. Savko, U. I. Mamai, and M. F. Finkel'shtein, *Russ. J. Phys. Chem.* **47**, 1531 (1973).
- 1973FIN/GAR A. Finch, P. J. Gardner, N. Hill, and K. S. Hussian, *J. Chem. Soc. Dalton Trans.* 2543 (1973).
- 1973FRA/KRZ H. P. Frank, K. Krzemicki, and H. Voellenkle, *Chem. Ztg.* **97**, 206 (1973).
- 1973GAL/BRY G. L. Gal'chenko, E. P. Brykina, R. M. Varushchenko, L. S. Vasil'ev, and B. N. Mikhailov, *Russ. J. Phys. Chem.* **47**, 914 (1973).
- 1973GEI/DZH K. I. Geidarov, O. I. Dzhabarov, K. A. Karasharli, and V. N. Kostryukov, *Zh. Fiz. Khim.* **47**, 275 (1973); *Chem. Abstr.* **78**, 129067s (1973).
- 1973GEI/QUI G. Geiseler, K. Quitzsch, H.-P. Hofmann, and R. Z. Pfestorf, *Z. Phys. Chem. (Leipzig)* **252**, 170 (1973).
- 1973GEI/SAW G. Geiseler and J. Sawistowsky, *Z. Phys. Chem. (Leipzig)* **253**, 333 (1973).
- 1973GIG/MAL R. Gigli, L. Malaspina, and G. Bardi, *Ann. Chim. (Rome)* **63**, 627 (1973); *Chem. Abstr.* **82**, 77775g (1975).
- 1973GOS/MIL K. Gosling and J. L. Miller, *Inorg. Nucl. Chem. Lett.* **9**, 355 (1973).
- 1973GOT/MEN F. A. Gothard, D. C. Mentianu, D. G. Breban, and C. I. Cristea, *J. Chem. Eng. Data* **18**, 381 (1973).
- 1973HAL/SMI H. K. Hall, C. D. Smith, and J. H. Baldt, *J. Am. Chem. Soc.* **95**, 3197 (1973).
- 1973HAM/AYE J. V. Hamilton and D. A. Ayers, *J. Chem. Eng. Data* **18**, 366 (1973).
- 1973HAM/MIT J. V. Hamilton and G. M. Mitchell, *J. Chem. Eng. Data* **18**, 38 (1973).
- 1973HAM/MIT2 J. V. Hamilton, G. M. Mitchell, and D. A. Ayers, *J. Chem. Eng. Data* **18**, 364 (1973).
- 1973IVA/GUJ G. A. Ivanov, V. G. Gujlya, and A. A. Zhukhovitskii, *Zavodskaya Lab.* **39**, 15 (1973); *Chem. Abstr.* **78**, 129058 (1973).
- 1973IZM/KHO A. S. Izmailovich, Y. S. Khodeev, and V. I. Tsirel'nikov, *Zh. Fiz. Khim.* **47**, 1565 (1973); *Chem. Abstr.* **79**, 108298y (1973).
- 1973JAI/YAD D. V. S. Jain and O. P. Yadav, *Indian J. Chem.* **11**, 28 (1973).
- 1973KIS/SUG K. Kishimoto, H. Suga, and S. Seki, *Bull. Chem. Soc. Jpn.* **46**, 3020 (1973).
- 1973KKY/REP J. Kkykj and M. Repas, *Petrochemia* **13**, 179 (1973).
- 1973KOL/RAB G. M. Kol'yakova, I. B. Rabinovich, and E. N. Zorina, *Dokl. Akad. Nauk SSSR* **209**, 616 (1973).
- 1973KON J. Konicek, *Acta Chem. Scand.* **27**, 1496 (1973).
- 1973KRE/PRI M. J. Krech, S. J. W. Price, and W. F. Yared, *Can. J. Chem.* **51**, 3662 (1973).
- 1973KRI/LIC G. Krien, H. H. Licht, and J. Zierath, *Thermochim. Acta* **6**, 465 (1973).
- 1973LEB/KAT N. D. Lebedeva and Y. A. Katin, *Zh. Fiz. Khim.* **47**, 1620 (1973); see also *Russ. J. Phys. Chem.* **47**, 922 (1973).
- 1973LEB/KAT2 N. D. Lebedeva and Y. A. Katin, *Russ. J. Appl. Chem.* **46**, 2131 (1973).
- 1973LIN/WIC J. Linek and I. Wichterle, *Collect. Czech. Chem. Commun.* **38**, 1846 (1973).
- 1973MAJ/SHR A. Majid and J. M. Shreeve, *J. Org. Chem.* **38**, 4028 (1973).
- 1973MAL/BAR L. Malaspina, R. Gigli, and G. Bardi, *J. Chem. Thermodyn.* **5**, 845 (1973).
- 1973MAL/GIG L. Malaspina, R. Gigli, and G. Bardi, *J. Chem. Phys.* **59**, 387 (1973).
- 1973MAL/GIG2 L. Malaspina, R. Gigli, G. Bardi, and G. DeMaria, *J. Chem. Thermodyn.* **5**, 699 (1973).
- 1973MCC/SMI J. A. McCauley and N. O. Smith, *J. Chem. Thermodyn.* **5**, 31 (1973).
- 1973MCD W. McDowell, *J. Soc. Dyers Colour.* **895**, 177 (1973).
- 1973MCE/SAN D. M. McEachern and O. Sandoval, *J. Phys. E* **6**, 155 (1973).
- 1973MEY/HOT E. F. Meyer and R. D. Hotz, *J. Chem. Eng. Data* **18**, 359 (1973).
- 1973MOR/MCN C. T. Mortimer and J. L. McNaughton, *Thermochim. Acta* **6**, 269 (1973).
- 1973PEP/GAF V. I. Pepekin, R. G. Gafurov, Y. A. Lebedev, L. T. Eremenko, E. M. Sogomonyan, and A. Y. Apin, *Izv. Akad. Nauk SSSR Ser. Khim.* **22**, 318 (1973); see also *Izv. Akad. Nauk SSSR Ser. Khim. Eng. Edit.* **22**, 304 (1973).
- 1973PEP/LEB V. I. Pepekin, Y. A. Lebedev, and A. Y. Apin, *Dokl. Phys. Chem.* **208**, 35 (1973).
- 1973POL/POL L. D. Polyachenok and O. G. Polyachenok, *Zh. Fiz. Khim.* **47**, 498 (1973).
- 1973RAU/GEI H.-J. Rauh and G. Geiseler, *Z. Phys. Chem. (Leipzig)* **252**, 395 (1973).
- 1973RAU/GEY H.-J. Rauh, W. Geyer, H. Schmidt, and G. Geiseler, *Z. Phys. Chem. (Leipzig)* **253**, 43 (1973).
- 1973ROC/SYM D. H. Rochester and J. R. Symonds, *J. Chem. Soc., Faraday Trans. 1* **1**, 1267 (1973).
- 1973ROD/WES D. L. Rodgers, E. F. Westrum, Jr., and J. T. S. Andrews, *J. Chem. Thermodyn.* **5**, 733 (1973).
- 1973ROG/QUA F. E. Rogers and S. W. Quan, *J. Phys. Chem.* **77**, 828 (1973).
- 1973SCH/PIL C. J. Schack, D. Pilipovich, and J. F. Hon, *Inorg. Chem.* **12**, 897 (1973).
- 1973SPR/WRI G. H. Sprenger, K. J. Wright, and J. M. Shreeve, *Inorg. Chem.* **12**, 2890 (1973).
- 1973SVO/VES V. Svoboda, F. Vesely, R. Holub, and J. Pick, *Collect.*

- Czech. Chem. Commun. **38**, 3539 (1973).
- 1973TEL/RAB V. I. Tel'noi, I. B. Rabinovich, and V. A. Umilin, Dokl. Akad. Nauk SSSR **209**, 127 (1973).
- 1973UMI/FED V. A. Umilin, I. E. Fedorov, Y. B. Zverev, and A. Gorbunova, Tr. Khim. Khim. Tekhnol., 63 (1973); Chem. Abstr. **81**, 140914y (1974).
- 1973VAS/KOR I. A. Vasil'ev and A. D. Korkhov, Zh. Fiz. Khim. **47**, 2710 (1973).
- 1973WAN/SHR C. S.-C. Wang and J. M. Shreeve, *Inorg. Chem.* **12**, 81 (1973).
- 1973WAR/SKU W. Waradzin and P. Skubla, Chem. Prumsyl **23**, 556 (1973); Chem. Abstr. **80**, 112780n (1974).
- 1973WIL/FEN J. W. Wilson and J. T. F. Fenurck, *J. Chem. Thermodyn.* **5**, 341 (1973).
- 1973WIL/ZWO R. C. Wilhoit and B. J. Zwolinski, *J. Phys. Chem. Ref. Data Suppl.* **1**, 1 (1973). (Note: the authors report Antoine constants for numerous alcohols that were calculated from vapor pressure data taken from the published literature. For several of the alcohols, the authors had only a very limited number of experimental data points that were pooled from several sources.)
- 1973WRI/SHR K. J. Wright and J. M. Shreeve, *Inorg. Chem.* **12**, 77 (1973).
- 1974AMB/ELL D. Ambrose and J. H. Ellender, *J. Chem. Thermodyn.* **6**, 909 (1974).
- 1974AMB/SPR D. Ambrose and C. H. S. Sprake, *J. Chem. Thermodyn.* **6**, 453 (1974).
- 1974AMB/SPR2 D. Ambrose, C. H. S. Sprake, and R. J. Townsend, *J. Chem. Thermodyn.* **6**, 693 (1974).
- 1974AND/COU R. J. L. Andon, J. F. Councell, D. A. Lee, and J. F. Martin, *J. Chem. Soc., Faraday Trans. 1* **70**, 1914 (1974).
- 1974ARS M. R. Arshadi, *J. Chem. Soc., Faraday Trans. 1* **70**, 1569 (1974).
- 1974ATA/CHI T. Atake and H. Chihara, *Bull. Chem. Soc. Jpn.* **49**, 2126 (1974).
- 1974AUG/BOR E. Augdahl, G. Borgen, J. Dale, and J. Krane, *Acta Chem. Scand., Ser. B* **28B**, 125 (1974).
- 1974BAR/PIL D. S. Barnes, G. Pilcher, D. A. Pittam, H. A. Skinner, D. Todd, and Y. Virmani, *J. Less-Common Met.* **36**, 177 (1974); *J. Less-Common Met.* **38**, 53 (1974).
- 1974BEA/MUE P. Beak, D. S. Mueller, and J. Lee, *J. Am. Chem. Soc.* **96**, 3867 (1974).
- 1974BES/CHE N. A. Bessarab, F. S. Chernoglazova, and Y. M. Martynov, Zh. Fiz. Khim. **48**, 235 (1974); Chem. Abstr. **80**, 112833g (1974).
- 1974BES/MAR N. A. Bessareb, and Y. M. Martynov, Russ. J. Phys. Chem. **48**, 1547 (1974).
- 1974BJO/BOR S. L. Björnstad, G. Borgen, and G. Gaupset, *Acta Chem. Scand., Ser. B* **28B**, 821 (1974).
- 1974BLA/LEV R. L. Blaine and P. Levy, *Anal. Calorim.* **3**, 185 (1974).
- 1974BOR G. Borgen, *Acta Chem. Scand., Ser. B* **28B**, 13 (1974).
- 1974BRA/KAR G. P. Bragin and M. K. Karapet'yants, Russ. J. Phys. Chem. **48**, 612 (1974).
- 1974BRO/CON L. S. Brown, J. A. Connor, and H. A. Skinner, *J. Organomet. Chem.* **81**, 403 (1974).
- 1974BRY/CAZ C. E. Bryson III, V. Cazcarra, and L. L. Levenson, *J. Chem. Eng. Data* **19**, 107 (1974).
- 1974CAL/KAN L. M. Calle and A. S. Kana'an, *J. Chem. Thermodyn.* **6**, 935 (1974).
- 1974CAN/JAC J. Canceill, J. Jacques, and M. C. Perucaud-Brianso, *Bull. Soc. Chim. Fr.*, 2833 (1974).
- 1974CLA/MCK T. Clark, M. A. McKervey, H. Mackle, and J. Rooney, *J. Chem. Soc., Faraday Trans. 1* **70**, 1279 (1974).
- 1974COL/ROU M. Colomina, M. V. Roux, and C. Turrion, *J. Chem. Thermodyn.* **6**, 149 (1974).
- 1974COL/ROU2 M. Colomina, M. V. Roux, and C. Turrion, *J. Chem. Thermodyn.* **6**, 571 (1974).
- 1974CON/SKI J. A. Connor, H. A. Skinner, and Y. Virmani, *Faraday Symposium 8, High Temperature Chemistry, 1974* (unpublished), Vol. 18.
- 1974COX C. D. Cox, *Pure Appl. Chem.* **40**, 399 (1974).
- 1974DEV/RAB C. G. Devyatikh, I. B. Rabinovitch, V. I. Tel'noi, G. K. Borisov, and L. F. Zyazina, Dokl. Akad. Nauk SSSR **217**, 673 (1974).
- 1974DIT/SKO V. E. Ditsent, I. I. Skorokhodov, N. A. Terent'eva, and M. N. Zolotareva, Russ. J. Phys. Chem. **48**, 447 (1974); see also Zh. Fiz. Khim. **48**, 2152 (1974); Chem. Abstr. **81**, 158939v (1974).
- 1974DIT/SKO2 V. E. Ditsent, I. I. Skorokhodov, N. A. Terent'eva, M. N. Zolotareva, and M. B. Lotarev, Russ. J. Phys. Chem. **48**, 1279 (1974); Chem. Abstr. **81**, 158938u (1974).
- 1974DIT/SKO3 V. E. Ditsent, I. I. Skorokhodov, N. A. Terent'eva, M. N. Zolotareva, N. E. Rodzeczich, and O. I. Minosyan, Zh. Fiz. Khim. **48**, 2153 (1974); Chem. Abstr. **81**, 158940p (1974).
- 1974DIT/SKO4 V. E. Ditsent, I. I. Skorokhodov, N. A. Terent'eva, M. N. Zolotareva, B. N. Parfenov, and O. I. Minosyan, Zh. Fiz. Khim. **48**, 2151 (1974); Chem. Abstr. **81**, 158937t (1974).
- 1974DOJ/HEI J. Dojcansky and J. Heinrich, *Chem. Zvesti* **28**, 157 (1974); Chem. Abstr. **82**, 64723p (1975).
- 1974ENG/WOO P. S. Engel, J. L. Wood, J. A. Sweet, and J. L. Margrave, *J. Am. Chem. Soc.* **96**, 2381 (1974).
- 1974FRO H. Frostling, *Acta Chem. Scand.* **28**, 83 (1974).
- 1974GAI/BAE I. L. Gaidym, A. K. Baev, V. G. Syrkin, A. A. Uel'skii, and A. V. Medvedeva, Zh. Fiz. Khim. **48**, 1871 (1974); Chem. Abstr. **81**, 160019p (1974).
- 1974GAI/BAE2 I. L. Gaidym, A. K. Baev, V. G. Syrkin, A. A. Uel'skii, and A. E. Medvedeva, Russ. J. Phys. Chem. **48**, 1111 (1974); Chem. Abstr. **81**, 160019p (1974).
- 1974GEI/DZH K. I. Geidarov, O. I. Dzhaforov, K. A. Karasharli, and V. N. Kostyukov, Zh. Fiz. Khim. **48**, 1058 (1974).
- 1974GIB/CRE H. F. Gibbard and J. L. Creek, *J. Chem. Eng. Data* **19**, 308 (1974).
- 1974GIL/SUL A. G. Gilbert and K. G. P. Sulzmann, *J. Electrochem. Soc.* **121**, 832 (1974).
- 1974GOO/MOO W. D. Good, R. T. Moore, A. G. Osborn, and D. R. Douglas, *J. Chem. Thermodyn.* **6**, 303 (1974).
- 1974GUS/REN J. L. Gustin and H. Renon, *Bull. Soc. Chim. Fr.*, 2719 (1974).
- 1974HIR/CAS C. Hirayama, P. M. Castle, R. W. Liebermann, R. J. Zollweg, and F. E. Camp, *Inorg. Chem.* **13**, 2804 (1974).
- 1974JAC A. W. Jackowski, *J. Chem. Thermodyn.* **6**, 49 (1974).
- 1974KIV/NAD N. N. Kiva, S. I. Nadutkina, L. V. Ivanova, and V. S. Bogdanov, Russ. J. Phys. Chem. **148**, 297 (1974).
- 1974KOZ/BYC M. P. Kozina, L. V. Bychikhina, G. L. Gal'chenko, A. N. Kalinichenko, A. A. Bobyleva, N. A. Belikova, and A. F. Plate, Zh. Fiz. Khim. **48**, 2075 (1974); Chem. Abstr. **81**, 159978u (1974).
- 1974KUN/HOS Y. Kuniya, S. Hosoda, and M. Hosaka, *Denki Kagaku* **42**, 20 (1974); see also [1998BRU/PIA].
- 1974KUS/SAI K. Kusano and Y. Saito, Preprints of the Tenth Conference on Chemical Thermodynamics and Thermal Analysis, Japan, 1974 (unpublished), p. 155; as cited in Ref. [1985MAJ/SVO].
- 1974LET/MAR T. M. Letcher and F. Marsicano, *J. Chem. Thermodyn.* **6**, 509 (1974).
- 1974MAJ/SHR A. Majid and J. M. Shreeve, *Inorg. Chem.* **13**, 2710 (1974).
- 1974MAL/BAR L. Malaspina, G. Bardi, and R. Gigli, *J. Chem. Thermodyn.* **6**, 1053 (1974).
- 1974MAN M. Mansson, *Acta Chem. Scand., Ser. B* **28**, 905 (1974).
- 1974MAN2 M. Mansson, *Acta Chem. Scand., Ser. B* **28**, 895 (1974).
- 1974MAN3 M. Mansson, *Acta Chem. Scand., Ser. B* **28**, 677 (1974).
- 1974MAN4 M. Mansson, *J. Chem. Thermodyn.* **6**, 1153 (1974).
- 1974MES/FIN J. F. Messerly, H. L. Finke, and S. S. Todd, *J. Chem. Thermodyn.* **6**, 635 (1974).
- 1974MOG/HOC P. H. Mogul, M. C. Hochberg, R. Michiel, G. K. Nestel, B. L. Wamsley, and S. D. Coren, *J. Chem. Eng. Data* **19**, 4 (1974).
- 1974MOR/MCN C. T. Mortimer and J. L. McNaughton, *Thermochim. Acta* **8**, 265 (1974).
- 1974MOS/MOU C. Mosselman and J. Mouric, *J. Chem. Thermodyn.* **6**, 477 (1974).
- 1974MOZ/KOL L. V. Mozhginskay and L. E. Kolysko, Zh. Fiz. Khim. **48**,

- 1506 (1974).
- 1974MUR/POL J. J. Murray, R. F. Pollie, and C. Pupp, *Can. J. Chem.* **52**, 557 (1974).
- 1974MUR/TUD R. A. Murogova, G. L. Tudorovskaya, V. V. Laufer, V. D. Koxlova, and L. A. Seraphimov, *Zh. Prikl. Khim. (S.-Peterburg)* **47**, 2024 (1974).
- 1974MYA/SCH L. F. Myasinkova, V. A. Schmelev, I. L. Vaisman, V. I. Bushinskii, and D. A. Novokhatka, *Zh. Prikl. Khim. (Leningrad)* **47**, 2604 (1974).
- 1974OSB/DOU A. G. Osborn and D. R. Douslin, *J. Chem. Eng. Data* **19**, 114 (1974).
- 1974PEP/ERL V. I. Pepekin, R. D. Erlikh, Y. N. Matyushin, and Y. A. Lebedev, *Dokl. Phys. Chem.* **214**, 123 (1974).
- 1974PEP/LEB V. I. Pepekin, V. P. Lebedev, A. A. Balepin, and Y. A. Lebedev, *Dokl. Akad. Nauk SSSR* **221**, 1118 (1974).
- 1974PEP/MAT V. I. Pepekin, Y. N. Matyushin, and Y. A. Lebedev, *Izv. Akad. Nauk SSSR Ser. Khim.* **23**, 1786 (1974); see also *Izv. Akad. Nauk SSSR Ser. Khim. Eng. Edit.* **23**, 1707 (1974).
- 1974PET/SHR K. E. Peterman and J. M. Shreeve, *Inorg. Chem.* **13**, 2705 (1974).
- 1974PRI/POU J. Pribilova and J. Pouchly, *Collect. Czech. Chem. Commun.* **39**, 1118 (1974).
- 1974RAD/KAT L. G. Radchenko and A. I. Kataigorodskii, *Zh. Fiz. Khim.* **48**, 2702 (1974); see also *Russ. J. Phys. Chem.* **48**, 1595 (1974).
- 1974ROU/TUR M. V. Roux, C. Turrion, M. Colomina, and R. Perez-Ossario, *An. Quim.* **70**, 201 (1974); *Chem. Abstr.* **81**, 119718c (1974).
- 1974SAB R. Sabbah, *Energetique de Liaison Inter-et Intramoleculaires* (CNRS, Marseille, 1974).
- 1974SAB/CHA R. Sabbah, R. Chastel, and M. Laffite, *Can. J. Chem.* **52**, 2201 (1974).
- 1974SAB/CHA2 R. Sabbah, R. Chastel, and M. Laffite, *Thermochim. Acta* **10**, 353 (1974).
- 1974SAB/GIL R. Sabbah and E. M. Gilbert, *Thermochim. Acta* **10**, 345 (1974).
- 1974SAS/KON Y. Sassa, R. Konishi, and T. Katayama, *J. Chem. Eng. Data* **19**, 44 (1974).
- 1974SHI/GRE C. F. Shieh and N. W. Gregory, *J. Chem. Eng. Data* **19**, 11 (1974).
- 1974SHM/GOL G. O. Shmyreva, R. M. Golosova, G. B. Sakharovskaya, A. F. Popov, N. N. Korneev, and A. A. Smolyaninova, *Russ. J. Phys. Chem.* **48**, 447 (1974).
- 1974SHM/SHL G. O. Shmyreva, G. B. Sakharovskaya, R. M. Golosova, G. G. Filippov, E. A. Volnina, E. N. Burdasov, and L. E. Gusel'nikov, *Russ. J. Phys. Chem.* **48**, 1699 (1974).
- 1974SIN G. C. Sinke, *J. Chem. Thermodyn.* **6**, 311 (1974).
- 1974SPE/SHR G. H. Spenger and J. M. Shreeve, *J. Fluorine Chem.* **4**, 201 (1974).
- 1974SUN/WUL S. Sunner and C. A. Wulff, *J. Chem. Thermodyn.* **6**, 287 (1974).
- 1974TEP/SUK A. B. Teplitskii, L. F. Sukhodub, and I. K. Yanson, *Fiz. Kondens. Sostoyaniya* **32**, 68 (1974); *Chem. Abstr.* **84**, 170535e (1976).
- 1974UTS/BAC H. Utschick, G. Bachmann, and H. Kapitza, *Chem. Tech. (Leipzig)* **26**, 423 (1974).
- 1974UTS/BAC2 H. Utschick, G. Bachmann, and H. Kapitza, *Chem. Tech. (Leipzig)* **26**, 422 (1974).
- 1974VAR/BUL R. M. Varushchenko and L. L. Bulgakova, *Tr. Khim. Khim. Tekhnol.* **1**, 69 (1974).
- 1974VAR/DRU R. M. Varushchenko, A. I. Druzhimina, O. Y. Kovner, E. M. Mil'viitskaya, A. A. Bobileva, N. A. Belikova, and G. L. Gal'chenko, *Russ. J. Phys. Chem.* **48**, 1121 (1974).
- 1974VAS/KOR I. A. Vasil'ev and A. D. Korkhov, *Tr. Khim. Khim. Tekhnol.* **103** (1974); *Chem. Abstr.* **82**, 65233r (1975).
- 1974WAR/WIL M. D. Warren and W. S. Wilson, *Thermochim. Acta* **10**, 33 (1974).
- 1974YAN/VER I. K. Yanson, B. I. Verkin, O. I. Shlyarevskii, and A. B. Teplitskii, *Stud. Biophys.* **46**, 29 (1974); *Chem. Abstr.* **83**, 8884a (1975).
- 1974ZHU/MON E. Z. Zhuravlev, N. M. Moncharzh, and V. G. Vodop'yanov, *Trudy Khim. Khim. Tekhnol.*, 102 (1974).
- 1974ZOR/UMI A. D. Zorin, V. A. Umilin, and V. K. Vanchagova, *Zh. Obshch. Khim.* **44**, 592 (1974); as cited in J. A. Connor, M. I. El Saied, J. A. Martinho-Simoes, and H. A. Skinner, *J. Organomet. Chem.* **212**, 405 (1981).
- 1975ADE/BRO F. A. Adedeji, D. L. S. Brown, J. A. Connor, M. L. Leung, I. M. Paz-Andrade, and H. A. Skinner, *J. Organomet. Chem.* **97**, 221 (1975).
- 1975AMB/CON D. Ambrose, J. E. Connett, J. H. S. Green, J. L. Hales, A. J. Head, and J. F. Martin, *J. Chem. Thermodyn.* **7**, 1143 (1975).
- 1975AMB/ELL D. Ambrose, J. H. Ellender, E. B. Lees, C. H. S. Sprake, and R. J. Townsend, *J. Chem. Thermodyn.* **7**, 453 (1975).
- 1975AMB/ELL2 D. Ambrose, J. W. Ellender, C. H. S. Sprake, and R. I. Townsend, *J. Chem. Soc., Faraday Trans. 1* **71**, 35 (1975).
- 1975AMB/LAW D. Ambrose, I. J. Lawrenson, and C. H. S. Sprake, *J. Chem. Thermodyn.* **7**, 1173 (1975).
- 1975ANA/GRO S. C. Anand, J.-P. E. Grolier, O. Kiyohara, C. J. Halpin, and G. C. Benson, *J. Chem. Eng. Data* **20**, 184 (1975).
- 1975ANT/CAR M. E. Anthony, A. S. Carson, and P. G. Laye, *Conf. Int. Thermodyn. Chim. C. R. 4th* **1**, 99 (1975); *Chem. Abstr.* **84**, 163961n (1976).
- 1975ARN/RAN D. E. Arnold and D. W. H. Rankin, *J. Chem. Soc. Dalton Trans.* 889 (1975).
- 1975ARR/MEL J. Arro, L. Melder, and H. Tamvelius, *Trudy Tallinskogo Instit. Toimetict.*, 37 (1975).
- 1975AUB/MAY M. Aubry, M. N. Mayoral, and P. Villardry, *Bull. Soc. Chim. Fr.* **112**, 500 (1975).
- 1975BAG/AND G. V. Bagrov, S. N. Andreev, M. F. Smirnova, and L. A. Vol'f, *Deposited Doc. VINITI*, 1196 (1975); *Chem. Abstr.* **87**, 59208y (1977).
- 1975BAR/PIL D. S. Barnes and G. Pilcher, *J. Chem. Thermodyn.* **7**, 377 (1975).
- 1975BER/OLO S. Bergstrom and G. Olofsson, *J. Solution Chem.* **4**, 535 (1975).
- 1975BER/PER G. Bertholon and R. Perrin, *Bull. Soc. Chim. Fr.* **7/8**, 1537 (1975).
- 1975BJO/BOR S. L. Björnstad and G. Borgen, *Acta Chem. Scand., Ser. B* **29B**, 13 (1975).
- 1975BJO/BOR2 S. L. Björnstad, G. Borgen, J. Dale, and G. Gaupset, *Acta Chem. Scand., Ser. B* **29B**, 320 (1975).
- 1975BOG/BER K. A. Bogdanova, A. A. Berlin, V. Z. Kompanets, G. V. Rakova, E. A. Miroshnichenko, Y. A. Lebedev, and N. S. Enikolopyan, *Vysokomol. Soedin., Ser. A* **17**, 658 (1975); *Chem. Abstr.* **83**, 43841f (1975).
- 1975BOR G. Borgen, *Acta Chem. Scand., Ser. B* **29B**, 265 (1975).
- 1975BRA/AND M. M. Brazhnikov, D. N. Andreevskii, A. I. Sachek, and A. D. Peshchenko, *Zh. Prikl. Khim. (Leningrad)* **48**, 2181 (1975).
- 1975BRA/KAR G. P. Bragin and M. K. Karapet'yants, *Tr. Khim. Khim. Tekhnol.*, 78 (1975); *Chem. Abstr.* **85**, 83381d (1976).
- 1975BRO/CON D. L. S. Brown, J. A. Connor, and H. A. Skinner, *J. Chem. Soc., Faraday Trans. 1* **71**, 699 (1975).
- 1975CAB/CON T. Cabani, G. Conti, D. Giannessi, and L. Lepori, *J. Chem. Soc., Faraday Trans. 1* **71**, 1154 (1975).
- 1975CAB/CON2 S. Cabani, G. Conti, V. Mollica, and L. Lepori, *J. Chem. Soc., Faraday Trans. 1* **71**, 1943 (1975).
- 1975CAR/LAY A. S. Carson, P. G. Laye, and H. Morris, *J. Chem. Thermodyn.* **7**, 993 (1975).
- 1975CHI J. S. Chickos, *J. Chem. Educ.* **52**, 134 (1975).
- 1975CHU/DRU K. W. Chun, J. C. Drummond, W. H. Smith, and R. R. Davison, *J. Chem. Eng. Data* **20**, 58 (1975).
- 1975CLA/KNO T. Clark, T. Knox, H. Mackle, M. A. McKervey, and J. J. Rooney, *J. Chem. Soc., Faraday Trans. 1* **71**, 2107 (1975).
- 1975CUC A. Cuccuru, *Thermochim. Acta* **13**, 96 (1975).
- 1975CUC2 A. Cuccuru, *Thermochim. Acta* **11**, 247 (1975).
- 1975DEK/VAN C. G. DeKruif and C. H. D. van Ginkel, and J. Voogd, *Fourth International Conference on Chemical Thermodynamics, Montpellier, France, 1975* (unpublished); as cited in Ref. [1982COL/JIM].
- 1975DEM/KOV R. A. De Marco, T. A. Kovacina, and W. B. Fox, *J. Fluorine Chem.* **5**, 221 (1975).

- 1975DEM/KOV2R. A. De Marco, T. A. Kovacina, and W. B. Fox, *J. Fluorine Chem.* **6**, 93 (1975).
- 1975FEN/HAR J. O. Fenwick, D. Harrop, and A. J. Head, *J. Chem. Thermodyn.* **7**, 943 (1975).
- 1975FRE/ADA H. K. Frensdorff and R. K. Adams, *J. Chem. Eng. Data* **20**, 13 (1975).
- 1975GAM/SIC M. Gambaruto, J. E. Sicre, and H. J. Schumacher, *J. Fluorine Chem.* **5**, 175 (1975).
- 1975GAR/CAR P. J. Gardner, A. Cartner, R. G. Cunninghame, and B. H. Robinson, *J. Chem. Soc. Dalton Trans.* 2582 (1975).
- 1975GEI/DZH K. I. Geidarov, O. I. Dzhafarov, and K. A. Karasharli, *Russ. J. Phys. Chem.* **49**, 197 (1975).
- 1975GOO/MES W. D. Good, J. F. Messerly, A. G. Osborn, and D. R. Douslin, *J. Chem. Thermodyn.* **7**, 285 (1975).
- 1975GRO H. A. J. Grovers, *Acta Crystallogr., Sect. A: Cryst. Phys., Diffraction, Theor. Gen. Crystallogr.* **31**, 380 (1975).
- 1975GUS/KAR A. Z. Guseinov, K. A. Karasharli, O. I. Dzhafarov, G. G. Nurullaev, N. S. Nametkin, L. E. Gusel'nikov, E. A. Volnina, E. N. Burdasov, and V. N. Vdovin, *Dokl. Akad. Nauk SSSR* **222**, 1369 (1975).
- 1975GUS/KUL A. Z. Guseinov, A. A. Kuliev, K. A. Karasharli, and O. I. Dzhafarov, *Azerb. Khim. Zh.*, 72 (1975); *Chem. Abstr.* **84**, 65359c (1976).
- 1975HAM/KUD S. Hamada, Y. Kudo, and M. Kawano, *Bull. Chem. Soc. Jpn.* **48**, 2963 (1975).
- 1975HOE/HOE C. Hoerner, A. Hoepfner, and B. Schmeiser, *Ber. Bunsenges. Phys. Chem.* **79**, 222 (1975).
- 1975HOE/PAR A. Hoepfner, N. Parekh, C. Hoerner, and A. Abdel-Hamid, *Ber. Bunsenges. Phys. Chem.* **79**, 216 (1975).
- 1975IRV/RIB R. J. Irving and M. A. V. Ribeiro da Silva, *J. Chem. Soc. Dalton Trans.* 798 (1975).
- 1975IRV/RIB2 R. J. Irving and M. A. V. Ribeiro da Silva, *J. Chem. Soc. Dalton Trans.* 1257 (1975).
- 1975IWA/DAT H. Iwasaki and K. Date, *Koatsu Gasu* **12**, 374 (1975).
- 1975KIR/LAS R. L. Kirchmeier, U. I. Lasouris, and J. M. Shreeve, *Inorg. Chem.* **14**, 592 (1975).
- 1975KON/SEL I. I. Konstantinov, V. D. Selivanov, and T. I. Melent'eva, *Zh. Fiz. Khim.* **49**, 1058 (1975).
- 1975KOS/SAM V. N. Kostyukov, O. P. Samorukov, N. K. Samorukova, and E. K. Chaplygina, *Russ. J. Phys. Chem.* **49**, 944 (1975).
- 1975KUN/LIL H. Kundel, U. Lille, and N. Kaidas, *Tr. Tallin. Politekh. Inst.* **390**, 107 (1975).
- 1975KUS/SAI K. Kusano and Y. Saito, *Preprints 33rd Ann. Meeting Chem. Soc. Japan* **1**, 123 (1975); as cited in Ref. [1985MAJ/SVO].
- 1975LEB/MIL B. V. Lebedev, V. I. Milov, L. Y. Tsvetkova, N. K. Lebedev, Y. G. Kirparisova, and A. A. Evstropov, *Zh. Fiz. Khim.* **49**, 1591 (1975).
- 1975LEB/TSV B. V. Lebedev, L. Y. Tsvetkova, E. G. Kiparisova, and N. K. Lebedev, *Russ. J. Phys. Chem.* **49**, 1265 (1975).
- 1975LEE/SLU W. Y. Lee and L. J. Slutsky, *J. Phys. Chem.* **79**, 2602 (1975).
- 1975MAL/GIG L. Malaspina, R. Gigli, G. Bardi, and G. DeMaria, *Conf. Int. Thermodyn. Chim. C.R. 4th* **1**, 54 (1975); *Chem. Abstr.* **84**, 179387r (1976).
- 1975MAS/SCO J. F. Masi and R. B. Scott, *J. Res. Natl. Bur. Stand., Sect. A* **79A**, 619 (1975).
- 1975MCE/INI D. M. McEachern, J. C. Iniguez, and H. C. Ornelas, *J. Chem. Eng. Data* **20**, 226 (1975).
- 1975MCE/SAN D. M. McEachern, O. Sandoval, and J. C. Iniguez, *J. Chem. Thermodyn.* **7**, 299 (1975).
- 1975MEK/KAR S. A. Mekhtiev, K. A. Karasharli, and D. I. Dzhafarov, *Russ. J. Phys. Chem.* **49**, 259 (1975).
- 1975MEK/KAR2S. A. Mekhtiev, K. A. Karasharli, D. I. Dzhafarov, and A. G. Kuznetsova, *Russ. J. Phys. Chem.* **49**, 1914 (1975).
- 1975MES/BAE U. Messow, J. Baer, K. Quitzsch, and G. Geiseler, *J. Prakt. Chem.* **317**, 114 (1975).
- 1975MES/FIN J. F. Messerly, H. L. Finke, A. G. Osborn, and D. R. Douslin, *J. Chem. Thermodyn.* **7**, 1029 (1975).
- 1975OBS/DOU A. G. Osborn and D. R. Douslin, *J. Chem. Eng. Data* **20**, 229 (1975).
- 1975PAR/STE W. Parker, W. V. Steele, W. Stirling, and I. Watt, *J. Chem. Thermodyn.* **7**, 795 (1975).
- 1975PEP/MAT V. I. Pepekin, Y. N. Matyushin, A. D. Nikolaeva, A. P. Kirsanov, L. V. Platonova, and Y. A. Lebedev, *Izv. Akad. Nauk SSSR Ser. Khim.*, 1870 (1975).
- 1975PET/SAN V. M. Petrov and L. Sandler, *Zh. Fiz. Khim.* **49**, 2797 (1975).
- 1975PET/SHR K. E. Peterman and J. M. Shreeve, *Inorg. Chem.* **14**, 1106 (1975).
- 1975PET/SHR2 K. E. Peterman and J. M. Shreeve, *Inorg. Chem.* **14**, 1223 (1975).
- 1975PET/SHR3 K. E. Peterman and J. M. Shreeve, *J. Fluorine Chem.* **6**, 83 (1975).
- 1975PIL/WAR G. Pilcher, M. J. Ware, and D. A. Pittam, *J. Less-Common Met.* **42**, 223 (1975).
- 1975PIS/ROZ V. V. Pisarev, A. M. Rozhnov, R. M. Varushchenko, and A. G. Sarkisov, *Russ. J. Phys. Chem.* **49**, 1450 (1975).
- 1975PIS/ROZ2 V. V. Pisarev, A. M. Rozhnov, R. M. Varushchenko, and A. G. Sarkisov, *Russ. J. Phys. Chem.* **49**, 1605 (1975).
- 1975PIT/PIL D. A. Pittam, G. Pilcher, D. S. Barnes, H. A. Skinner, and D. Todd, *J. Less-Common Met.* **42**, 217 (1975).
- 1975RIV F. Rivenq, *Bull. Soc. Chim. Fr.* **1**, 2433 (1975).
- 1975SOK/KAR V. B. Sokolov, M. K. Karapet'yants, T. N. Sergeeva, S. P. Kolesnikov, and B. L. Perl'mutter, *Izv. Vyssh. Uchebn. Zaved. Khim. Khim. Tekhnol.* **18**, 1319 (1975); *Chem. Abstr.* **84**, 22433e (1976).
- 1975SOK/KAR2 V. B. Sokolov, M. K. Karapet'yants, T. N. Sergeeva, S. P. Kolesnikov, and B. L. Perl'mutter, *Izv. Vyssh. Uchebn. Zaved. Khim. Khim. Tekhnol.* **18**, 1234 (1975); *Chem. Abstr.* **84**, 9167b (1976).
- 1975STE/SCH B. Steyer and F. P. Schafer, *Appl. Phys.* **7**, 113 (1975).
- 1975STRI/SUN G. Stridh and S. Sunner, *J. Chem. Thermodyn.* **7**, 161 (1975).
- 1975SUB/ZWO D. J. Subach and B. J. Zwolinski, *J. Chem. Eng. Data* **20**, 232 (1975).
- 1975TEL/KIR V. I. Tel'noi, K. V. Kirynov, V. I. Ermolaev, and I. B. Rabinovich, *Dokl. Akad. Nauk SSSR* **220**, 1088 (1975).
- 1975TEP/YAN A. B. Teplitskii and I. K. Yanson, *Biofizika* **20**, 189 (1975); *Chem. Abstr.* **83**, 16712h (1975).
- 1975TIT/CHU V. A. Titov, T. P. Chusova, and G. A. Kokovin, *Izv. Sib. Otd. Akad. Nauk SSSR Ser. Khim. Nauk*, 3 (1975); *Chem. Abstr.* **83**, 183512z (1975).
- 1975TRU/SUV V. I. Trusov, A. V. Suvorov, and R. N. Abakumova, *Zh. Neorg. Khim.* **20**, 501 (1975); *Chem. Abstr.* **82**, 161007r (1975).
- 1975VAN/DEK C. H. D. van Ginkel, G. C. DeKruif, and F. E. B. DeWaal, *J. Phys. E* **8**, 490 (1975).
- 1975VIL/PER R. Vilcu, S. Perisanu, and I. Ciocazanu, *Conf. Int. Thermodyn. Chim. C. R. 4th* **1**, 105 (1975).
- 1975WAL/DES N. S. Walker and D. D. DesMarteau, *J. Fluorine Chem.* **5**, 127 (1975).
- 1975WAL/DES2 N. S. Walker and D. D. DesMarteau, *J. Fluorine Chem.* **5**, 135 (1975).
- 1975WIL/PIN K. R. Wilson and R. E. Pincock, *J. Am. Chem. Soc.* **97**, 1474 (1975).
- 1975YAN/TEP I. K. Yanson and A. B. Teplitskii, *Zh. Fiz. Khim.* **49**, 736 (1975); see also *Russ. J. Phys. Chem.* **49**, 428 (1975).
- 1976AMB/ELL D. Ambrose, J. H. Ellender, C. H. S. Sprake, and R. J. Townsend, *J. Chem. Thermodyn.* **8**, 165 (1976).
- 1976AMB/LAW D. Ambrose, I. J. Lawrenson, and C. H. S. Sprake, *J. Chem. Thermodyn.* **8**, 503 (1976).
- 1976AMB/SPR D. Ambrose and C. H. S. Sprake, *J. Chem. Thermodyn.* **8**, 601 (1976).
- 1976AMM/BUL M. M. Ammar, L. L. Bulgakova, and R. M. Varushchenko, *Russ. J. Phys. Chem.* **50**, 1466 (1976).
- 1976ANT/CAR M. E. Anthony, A. S. Carson, and P. G. Laye, *J. Chem. Soc., Perkin Trans. 1* 1032 (1976).
- 1976ANT/CAR2 M. E. Anthony, A. S. Carson, P. G. Laye, and M. Yurekli, *J. Chem. Thermodyn.* **8**, 1009 (1976).
- 1976ASH S. J. Ashcroft, *J. Chem. Eng. Data* **21**, 397 (1976).
- 1976ATA/CHI T. Atake and H. Chihara, *Chem. Lett.* **5**, 683 (1976).
- 1976BAR/BOU M. A. Barnard, Y. Boukari, and F. Busnot, *Thermochim.*

- Acta* **16**, 267 (1976).
- 1976BER/BER G. Berchiesi, M. A. Bercuiesi, G. G. Lobbia, and D. Leonesi, *Gazz. Chim. Ital.* **106**, 549 (1976).
- 1976BOR/CHU G. K. Borisov and S. G. Chugunova, *Russ. J. Phys. Chem.* **50**, 1791 (1976).
- 1976BOR/DAL G. Borgen and J. Dale, *Acta Chem. Scand., Ser. B* **30B**, 711 (1976).
- 1976BRA/PES M. M. Brazhnikov, A. D. Peshchenko, and O. V. Ral'ko, *Zh. Prikl. Khim. (Leningrad)* **49**, 1041 (1976).
- 1976BRO/CON D. L. S. Brown, J. A. Connor, M. L. Leung, M. I. Paz-Andrade, and H. A. Skinner, *J. Organomet. Chem.* **110**, 79 (1976).
- 1976BUR/SHR C. A. Burton and J. M. Shreeve, *J. Am. Chem. Soc.* **98**, 6545 (1976).
- 1976BUT/CAR R. S. Butler, A. S. Carson, P. G. Laye, and W. V. Steele, *J. Chem. Thermodyn.* **8**, 1153 (1976).
- 1976CIH/HYN J. Cihlar, V. Hynek, V. Svoboda, and R. Holub, *Collect. Czech. Chem. Commun.* **41**, 1 (1976).
- 1976COL/JIM M. Colomina, P. Jimenez, R. Perez-Ossorio, and C. Turion, *J. Chem. Thermodyn.* **8**, 439 (1976).
- 1976CON/COU J. E. Connett, J. F. Councell, and D. A. Lee, *J. Chem. Thermodyn.* **8**, 1199 (1976).
- 1976DAV/FIN R. H. Davies, A. Finch, P. J. Gardner, A. Hameed, and M. Stephens, *J. Chem. Soc. Dalton Trans.* 556 (1976).
- 1976DEP R. S. Depablo, *J. Chem. Eng. Data* **21**, 141 (1976).
- 1976DIT/SKO V. E. Ditsent, I. I. Skorokhodov, N. A. Terent'eva, M. N. Zolotareva, Z. V. Belyakova, and Z. V. Belikova, *Zh. Fiz. Khim.* **50**, 1905 (1976); *Chem. Abstr.* **85**, 166806x (1976).
- 1976ENG/MEL P. S. Engel, R. A. Melaugh, M. Mansson, J. W. Timberlake, A. Garner, and F. D. Rossini, *J. Chem. Thermodyn.* **8**, 607 (1976).
- 1976FAL/DES E. R. Falardeau and D. D. Desmarteau, *J. Fluorine Chem.* **7**, 185 (1976).
- 1976FAL/DES2 E. R. Falardeau and D. D. Desmarteau, *J. Fluorine Chem.* **7**, 409 (1976).
- 1976FER/PIA D. Ferro, V. Piacente, R. Gigli, and G. D'Ascenzo, *J. Chem. Thermodyn.* **8**, 1137 (1976).
- 1976HA/MOR H. Ha, J. A. Morrison, and E. L. Richards, *J. Chem. Soc., Faraday Trans. 1* **72**, 1051 (1976).
- 1976HAM/THO W. S. Hamilton, P. Thompsom, and S. Pustejovsky, *J. Chem. Eng. Data* **21**, 428 (1976).
- 1976HON/SIN H. C. Hon, R. P. Singh, and A. P. Kudchadker, *J. Chem. Eng. Data* **21**, 430 (1976).
- 1976HOP/BOS H. P. Hopkins, Jr., D. Bostwick, and C. J. Alexander, *J. Am. Chem. Soc.* **98**, 1355 (1976).
- 1976HOP/DES M. J. Hopkinson and D. D. Desmarteau, *J. Fluorine Chem.* **7**, 501 (1976).
- 1976KIP/TSV E. G. Kiparisova, L. Y. Tsvetkova, and B. V. Lebedev, *Russ. J. Phys. Chem.* **50**, 638 (1976).
- 1976KIR/TEL K. V. Kir'yanov, V. I. Tel'noi, G. A. Vasil'eva, and I. B. Rabinovich, *Dokl. Akad. Nauk SSSR* **231**, 1021 (1976).
- 1976KOL/SLA V. P. Kolesov, G. M. Slavutskaya, and L. N. Dityat'eva, *J. Chem. Thermodyn.* **8**, 907 (1976).
- 1976KOZ/BYC M. P. Kozina, L. V. Bychikhina, G. V. Gal'chenko, E. M. Milwitskaya, M. Ordubade, and A. F. Plate, *Dokl. Akad. Nauk SSSR* **226**, 1105 (1976); see also *Dokl. Chem.* **226**, 143 (1976).
- 1976KUL/DZH A. M. Kuliev, O. I. Dzhaferov, K. A. Karasharli, and A. G. Kuznetsova, *Zh. Fiz. Khim.* **50**, 1903 (1976); *Chem. Abstr.* **85**, 183105 (1976).
- 1976KUZ/MIR V. P. Kuznetsova, E. A. Miroshinichenko, A. Z. Zenlenetskii, G. V. Rokova, Y. A. Lebedev, and N. S. Enikolopyan, *Dokl. Phys. Chem.* **226**, 147 (1976).
- 1976LEC/COL M. Leclercq, A. Collet, and J. Jacques, *Tetrahedron* **32**, 821 (1976).
- 1976MEL/MAN R. A. Melaugh, M. Mansson, and F. D. Rossini, *J. Chem. Thermodyn.* **8**, 623 (1976).
- 1976MEY/HOT E. F. Meyer and C. A. Hotz, *J. Chem. Eng. Data* **21**, 274 (1976).
- 1976MIR/LEB E. A. Miroshnichenko, and Y. A. Lebedev, *Mezhnol. Vzaimodeistvie Konform. Mol. Tezisy Dokl. Vses. Simp. 3rd*, 1976 (unpublished), pp. 40–41; *Chem. Abstr.* **91**, 56104h (1979).
- 1976MIR/PAV E. A. Miroshnichenko, V. K. Pavloich, A. Yu, Y. A. Lebedev, V. I. Stanko, and G. L. Gal'chenko, *Termodin. Org. Soedin* **5**, 3 (1976).
- 1976MIS/RIE S. Miskiewicz, K. Rieser, and T. Dorfmueller, *Ber. Bunsenges. Phys. Chem.* **80**, 395 (1976).
- 1976PEL P. A. Pella, *Anal. Chem.* **48**, 1632 (1976).
- 1976RAO/CHI K. V. Rao and C. Chiranjivi, *J. Chem. Eng. Data* **21**, 393 (1976).
- 1976SHM/SHL G. O. Shmyreva, V. G. Shlyakova, R. M. Golosova, A. B. Petrunin, D. B. Bekker, and A. F. Zhigach, *Russ. J. Phys. Chem.* **50**, 478 (1976).
- 1976STA/MEW I. Stahl, R. Mews, and O. Glemser, *J. Fluorine Chem.* **7**, 55 (1976).
- 1976STE/POL K. Steele, B. E. Poling, and D. B. Manley, *J. Chem. Eng. Data* **21**, 399 (1976).
- 1976STR G. Stridh, *J. Chem. Thermodyn.* **8**, 193 (1976).
- 1976STR2 G. Stridh, *J. Chem. Thermodyn.* **8**, 895 (1976).
- 1976STR3 G. Stridh, *J. Chem. Thermodyn.* **8**, 901 (1976).
- 1976TAV/NEE A. Tavlaridis and R. Neeb, *Naturwiss.* **63**, 146 (1976), as quoted in Ref. [79LAR].
- 1976TAY/CRO J. W. Taylor and R. J. Crookes, *J. Chem. Soc., Faraday Trans. 1* **72**, 723 (1976).
- 1976TEL/RAB V. I. Tel'noi, I. B. Rabinovich, K. V. Kir'yanov, and A. S. Smirnov, *Dokl. Akad. Nauk SSSR* **231**, 733 (1976).
- 1976VAR/BUL R. M. Varushchenko and L. L. Bulgakova, *Tr. Khim. Khim. Tekhnol.* **3**, 29 (1976).
- 1976ZHU/ALI L. V. Zhuravleva, A. S. Alikhanyan, and L. N. Sidorov, *Russ. J. Phys. Chem.* **50**, 818 (1976).
- 1976ZOR/RAC A. D. Zorin, O. F. Rachkova, and N. S. Belousova, *Termodin. Org. Soedin*, 40 (1976).
- 1977ALE/KOZ V. A. Aleshina, M. P. Kozina, and G. L. Gal'chenko, *Vses. Konf. Kalorim. [Rasshir. Tezisy Dokl.] 7th 1*, 85 (1977); *Chem. Abstr.* **92**, 94582r (1980).
- 1977BAL/BAR G. Balducci, G. Bardi, G. de Maria, and R. Gigli, VII All-Union Calorimetry Conference, 1977 (unpublished).
- 1977BAL/RUD T. S. Balanevskaya, V. V. Rudnev, N. G. Klishina, and A. A. Efremov, *Tr. VNI Khim. Reakt. Osobo Chist. Khim. Veshehestv*, 98 (1977).
- 1977BAR/GAI A. A. Barkatin, I. L. Gaidym, and A. K. Baev, *Khim. Khim. Technol. (USSR)* **12**, 44 (1977).
- 1977BED R. G. Bedford, *J. Phys. Chem.* **81**, 1284 (1977).
- 1977BED/HUS R. G. Bedford and E. B. Huss, *J. Chem. Eng. Data* **22**, 239 (1977).
- 1977BOL/MAK N. I. Boldenkov and Y. Y. Maksimov, *Zh. Fiz. Khim.* **51**, 1502 (1977).
- 1977BOU/CAR J. Bousquet, J. Carre, P. Claudy, M. Kollmanns-Berger, J. Thourey, and P. Barberi, *J. Therm. Anal. Calorim.* **8**, 83 (1977).
- 1977BRO/CON D. L. S. Brown, J. A. Connor, C. P. Demain, M. L. Lueng, J. A. Martinho-Simoes, H. A. Skinner, and M. T. Zarfarani-Moattar, *J. Organomet. Chem.* **142**, 321 (1977).
- 1977BUR A. Burger, *Sci. Pharm.* **45**, 269 (1977).
- 1977BUR/SHR C. A. Burton and J. M. Shreeve, *Inorg. Chem.* **16**, 1039 (1977).
- 1977BUR/SHR2 C. A. Burton and J. M. Shreeve, *Inorg. Chem.* **16**, 1408 (1977).
- 1977CAM/SCH H. K. Cammenga, F. W. Schulze, and W. Theuerl, *J. Chem. Eng. Data* **22**, 131 (1977).
- 1977CAR/LAY A. S. Carson, P. G. Laye, and M. Yurkeli, *J. Chem. Thermodyn.* **9**, 827 (1977).
- 1977CAS/VEC F. Casellato, C. Vecchi, and A. Girelli, *Thermochim. Acta* **21**, 195 (1977).
- 1977CLA/KNO T. Clark, T. M. O. Knox, H. Mackle, and M. A. McKervey, *J. Chem. Soc., Faraday Trans. 1* **73**, 1224 (1977).
- 1977DAS/REE T. R. Das, C. O. Reed, Jr., and P. T. Eubank, *J. Chem. Eng. Data* **22**, 3 (1977).
- 1977DAS/REE2 T. R. Das, C. O. Reed, Jr., and P. T. Eubank, *J. Chem. Eng. Data* **22**, 9 (1977).
- 1977DAS/REE3 T. R. Das, C. O. Reed, Jr., and P. T. Eubank, *J. Chem. Eng. Data* **22**, 16 (1977).

- 1977DEK/VAN C. G. DeKruif and C. H. D. van Ginkel, *J. Chem. Thermodyn.* **9**, 725 (1977).
- 1977DES M. L. Desauge, *J. Therm Anal. Calorim.* **8**, 89 (1977); *Chem. Abstr.* **87**, 206628r (1977).
- 1977DIT/KOL L. N. Dityat'eva and V. P. Kolesov, *Vses. Konf. Kalorim. [Rasshir. Tezisy Dokl.]* 7th **1**, 95 (1977); *Chem. Abstr.* **92**, 58009a (1980).
- 1977DWO/FUC A. Dworkin and A. H. Fuchs, *J. Chem. Phys.* **67**, 1789 (1977).
- 1977DYG/STE R. S. Dygdala, K. Stefanski, and J. Wonikowski, *Bull. Acad. Pol. Sci., Ser. Sci., Math., Astron. Phys.* **25**, 439 (1977).
- 1977EIB/TRO J. Eibl, W. Tröger, and K.-H. Sender, *Melliand Textilber.* **58**, 844 (1977).
- 1977FIN/MES H. L. Finke, J. F. Messerly, S. H. Lee, A. G. Osborn, and D. R. Douslin, *J. Chem. Thermodyn.* **9**, 937 (1977).
- 1977FRA J. Franklin, Ph.D. thesis, University of Leeds, 1977.
- 1977GAF/PIE J. S. Gaffney, R. C. Pierce, and L. Friedman, *J. Am. Chem. Soc.* **99**, 4293 (1977).
- 1977KAN/MOR A. S. Kana'an and T. I. Morrison, *J. Chem. Thermodyn.* **9**, 423 (1977).
- 1977KIM/KIM B. C. Kim and D. H. Kim, *Hwahak Kwa Hwahak Kongop* **20**, 232 (1977); *Chem. Abstr.* **88**, 95565s (1978).
- 1977KIT/SHR T. Kitazume and J. M. Shreeve, *Inorg. Chem.* **16**, 1818 (1977).
- 1977KIT/SHR2 T. Kitazume and J. M. Shreeve, *J. Am. Chem. Soc.* **99**, 3690 (1977).
- 1977KIT/SHR3 T. Kitazume and J. M. Shreeve, *J. Am. Chem. Soc.* **99**, 4194 (1977).
- 1977KOZ/BYC M. P. Kozina, V. L. Bychikhina, and G. L. Gal'chenko, *Zh. Fiz. Khim.* **51**, 2142 (1977).
- 1977KRE/PRI M. J. Krech, S. W. Price, and H. Sapiano, *Can. J. Chem.* **55**, 4222 (1977).
- 1977LEB/EVS B. V. Lebedev, A. A. Evstropov, G. B. Sadikov, and L. F. Larina, *Russ. J. Phys. Chem.* **51**, 762 (1977).
- 1977LEB/NAZ N. D. Lebedeva, L. F. Nazarova, and Y. Katin, *Org. Soedin.* **6**, 72 (1977).
- 1977LEB/RYA N. D. Lebedeva, V. L. Ryadnenko, N. N. Kiseleva, and L. F. Nazarova, *Vses. Konf. Kalorim. [Rasshir. Tezisy Dokl.]* 7th **1**, 91 (1977); *Chem. Abstr.* **92**, 75617q (1980).
- 1977LYU/SMO M. V. Lyubarskii and R. I. Smolyanets, *Vses. Konf. Kalorim. [Rasshir. Tezisy Dokl.]* 7th **1**, 119 (1977); *Chem. Abstr.* **92**, 75619s (1980).
- 1977MAN/SEL M. Mansson, P. Sellers, G. Stridh, and S. Sunner, *J. Chem. Thermodyn.* **9**, 91 (1977).
- 1977MEY/GEN E. F. Meyer and T. H. Gens, *J. Chem. Eng. Data* **22**, 30 (1977).
- 1977NAB/SAB M. Nabavian, R. Sabbah, R. Chastel, and M. Laffitte, *J. Chim. Phys. Phys.-Chim. Biol.* **74**, 115 (1977).
- 1977NAG H. Naghibi-Bidokhti, Ph.D. thesis, the University of Surrey, 1977.
- 1977NAG/SAB S. N. Ngauv, R. Sabbah, and M. Laffitte, *Thermochim. Acta* **20**, 371 (1977).
- 1977NIS/ISH K. Nishida, E. Ishihara, T. Osaka, and M. Koukitu, *J. Chem. Dyers Colour* **93**, 52 (1977).
- 1977NOV/NOV N. Novakova and J. Novak, *J. Chromatogr.* **135**, 13 (1977).
- 1977PAR/STE W. Parker, W. V. Steele, and I. Watt, *J. Chem. Thermodyn.* **9**, 307 (1977).
- 1977PEA/FUC L. A. Peacock and R. Fuchs, *J. Am. Chem. Soc.* **99**, 5524 (1977).
- 1977PED/RYL J. B. Pedley and J. Rylance, *N. P. L. Computer Analysed Thermochemical Data: Organic and Organometallic Compounds* (School of Molecular Sciences, University of Sussex, Brighton, 1977).
- 1977PEL P. A. Pella, *J. Chem. Thermodyn.* **9**, 301 (1977).
- 1977PIS/ROZ V. V. Pisarev, A. M. Rozhnov, and A. G. Sarkisov, *Russ. J. Phys. Chem.* **51**, 323 (1977).
- 1977RAO/VIU J. J. Rao and D. S. Viuswanath, *J. Chem. Eng. Data* **22**, 36 (1977).
- 1977SAB/NAB R. Sabbah, M. Nabavian, and M. Laffitte, *C. R. Hebd. Seances Acad. Ser. C.* **284**, 953 (1977); *Chem. Abstr.* **87**, 151519y (1977).
- 1977SAS/FAL T. M. Sas, V. A. Falin, N. E. Mazepova, I. A. Krasavin, and I. I. Bogomolov, *Zh. Fiz. Khim.* **51**, 1273 (1977).
- 1977SCH/PET F.-W. Schulze, H.-J. Petrick, H. K. Canmenga, and H. Klinge, *Z. Phys. Chem. (Wiesbaden)* **107**, 1 (1977). P. Sellers, *J. Chem. Thermodyn.* **9**, 139 (1977).
- 1977SEL W. V. Steele, *J. Chem. Thermodyn.* **9**, 311 (1977).
- 1977STE W. V. Steele and I. Watt, *J. Chem. Thermodyn.* **9**, 843 (1977).
- 1977STE2 H. Steinwandter, *Chemosphere* **6**, 59 (1977); *Chem. Abstr.* **87**, 17285r (1977).
- 1977STRI/SUN G. Stridh, S. Sunner, and C. Svenson, *J. Chem. Thermodyn.* **9**, 1005 (1977).
- 1977SVO/MAJ V. Svoboda, V. Majer, F. Vesely, and J. Pick, *Collect. Czech. Chem. Commun.* **42**, 1755 (1977).
- 1977SVO/VES V. Svoboda, F. Vesely, R. Holub, and J. Pick, *Collect. Czech. Chem. Commun.* **42**, 943 (1977).
- 1977TEL/RAB V. I. Tel'noi and I. B. Rabinovich, *Usp. Khim.* **46**, 1337 (1977); see also *Prog. Chem.* **46**, 689 (1977).
- 1977VAR/AMM R. M. Varushchenko, M. M. Ammar, and L. L. Bulgakova, *Zh. Fiz. Khim.* **51**, 278 (1977).
- 1977VAR/AMM R. M. Varushchenko, M. M. Ammar, and L. L. Bulgakova, *Russ. J. Phys. Chem.* **51**, 167 (1977).
- 1977VAS/KOT T. F. Vasil'eva and V. I. Kotov, *Vses. Konf. Kalorim. [Rasshir. Tezisy Dokl.]* 7th **1**, 102 (1977); *Chem. Abstr.* **92**, 58010u (1980).
- 1977VOL/MAZ S. V. Volkov, E. A. Mazurenko, and Z. N. Bublik, *Str. Svoistva Primen. B[Beta]-Diketonatov Met., [Mater. Vses. Semin.]* 3rd, 119 (1977); *Chem. Abstr.* **91**, 79727c (1977).
- 1977WON/WES W.-K. Wong and E. F. Westrum, Jr., *J. Phys. Chem.* **74**, 1303 (1970).
- 1977ZHU/MEL E. Z. Zhuravlev, T. L. Melent'eva, P. V. Mulyanov, A. I. Kormushechkina, T. M. Bogdanova, and I. I. Konstantinov, *Zh. Prikl. Khim. (Leningrad)* **50**, 2032 (1977).
- 1978ABA/MAL A. S. Abakumov and M. L. Malyshev, *Russ. J. Inorg. Chem.* **23**, 1442 (1978).
- 1978ADE/CON F. A. Adedeji, J. A. Connor, C. P. Demain, J. A. Martinho-Simoes, H. A. Skinner, and M. T. Zafarani-Moattar, *J. Organomet. Chem.* **149**, 333 (1978).
- 1978AND/CAR J. T. S. Andrews, R. E. Carpenter, T. M. Martinko, R. C. Fort, Jr., T. A. Flood, and M. G. Adlington, *Mol. Cryst. Liq. Cryst.* **41**, 257 (1978).
- 1978ARO/STE M. Arora and W. V. Steele, *J. Chem. Thermodyn.* **10**, 403 (1978).
- 1978BER/SPI C. V. Berney and D. Spickerman, *J. Chem. Thermodyn.* **10**, 637 (1978).
- 1978BOU/LEC D. Bougeard, N. A. Le Calve, A. Novak, and B. C. Nguyen, *Mol. Cryst. Liq. Cryst.* **44**, 113 (1978).
- 1978CAL/CAL C. D. H. Calis-Van Ginkel, G. H. M. Calis, C. W. M. Timmermans, C. G. DeKruif, and H. A. J. Oonk, *J. Chem. Thermodyn.* **10**, 1083 (1978).
- 1978CAN S. Cantor, *Thermochim. Acta* **26**, 39 (1978).
- 1978CAV/CHA S. D. Cave, A. Chaine, and A. Prantera, *J. Chem. Eng. Data* **23**, 279 (1978).
- 1978CHI/SHE J. S. Chickos and D. E. Sherwood, *J. Org. Chem.* **43**, 1146 (1978).
- 1978CHU/IGU Y. V. Chumachenko, I. K. Igumenov, and S. V. Zemskov, *Russ. J. Phys. Chem.* **52**, 1393 (1978).
- 1978COL/JIM M. Colomina, P. Jimenez, M. V. Roux, and C. Turrion, *J. Chem. Thermodyn.* **10**, 661 (1978).
- 1978COR/PER R. P. Corbally, M. J. Perkins, A. S. Carson, P. G. Laye, and W. V. Steele, *J. Chem. Soc., Chem. Commun.* 778 (1978).
- 1978CUN/PAL R. B. Cundall, T. F. Palmer, and C. E. C. Wood, *J. Chem. Soc., Faraday Trans. 1* **74**, 1339 (1978).
- 1978DAN/LEA D. M. Daniel, A. L. Leadbetter, R. E. Meads, and W. G. Parker, *J. Chem. Soc., Faraday Trans. 2* **74**, 456 (1978).
- 1978DAV/TRA A. V. Davydov, S. S. Travnikov, E. V. Fedoseev, L. A. Ivanova, and B. F. Myasoedov, *Probl. Khim. Primen. B [Beta]-Diketonatov Met., [Mater. Vses. Semin.]* 02-100 (1978).
- 1978DEM/FOX R. A. De Marco and W. B. Fox, *J. Fluorine Chem.* **12**, 137 (1978).

- 1978DEP R. S. DePablo, *J. Phys. D: Appl. Phys.* **11**, 193 (1978).
- 1978DOZ/FUJ Y. Dozen, H. Fujishima, and H. Shingu, *Thermochim. Acta* **25**, 209 (1978).
- 1978EKS/RAN A. Ekstrom and C. H. Randall, *J. Phys. Chem.* **82**, 2180 (1978).
- 1978ENG/MON P. S. Engel, R. L. Montgomery, M. Mansson, R. A. Leckonby, H. L. Foyt, and F. D. Rossini, *J. Chem. Thermodyn.* **10**, 205 (1978).
- 1978FUC/PEA R. Fuchs and L. A. Peacock, *Can. J. Chem.* **56**, 2493 (1978).
- 1978GAL/PAV G. L. Gal'chenko, V. K. Pavlovich, V. N. Siryatskaya, and A. F. Zhigach, *Termodin. Org. Soedin.* **7**, 38 (1978).
- 1978GRA/KON T. D. Grabik, S. G. Konstantinov, G. P. Dudchik, and O. G. Polyachenok, *Russ. J. Phys. Chem.* **52**, 894 (1978).
- 1978GRO/SPE R. Grover, W. F. Spencer, W. J. Farmer, and T. D. Shoup, *Weed Sci.* **26**, 505 (1978).
- 1978GUZ/LAR J. Guzman and J. J. Largo-Cabrerizo, *J. Heterocycl. Chem.* **15**, 1531 (1978).
- 1978HAM/BEN W. S. Hamilton, S. Benton, J. French, D. McCormick, S. Pustejovsky, and P. J. Thomspson, *J. Chem. Eng. Data* **23**, 201 (1978).
- 1978HEA/HEF G. A. Heath, G. T. Hefter, and W. V. Steele, *J. Chem. Thermodyn.* **10**, 395 (1978).
- 1978IGU/CHU I. K. Igumenov, Y. V. Chumachenko, and S. V. Zemskov, *Koord. Khim.* **4**, 164 (1978); **5**, 34 (1979).
- 1978IGU/CHU2 I. K. Igumenov, Y. U. Chumanchenko, and S. V. Zemskov, *Zh. Fiz. Khim.* **52**, 2664 (1978); see also *Russ. J. Phys. Chem.* **52**, 1531 (1978).
- 1978IND/STO D. Indritz, J. Stone, and F. Williams, *J. Chem. Eng. Data* **23**, 6 (1978).
- 1978IRV/SCH R. J. Irving, R. A. Schulz, and H. Naghihi (unpublished); cited in R. Irving and M. A. V. Riberio Da Silva, *J. Chem. Soc. Dalton Trans.* 399 (1978).
- 1978JES/ERN A. C. Jesse, J. M. Ernstring, D. J. Stufkens, and K. Vrieze, *Thermochim. Acta* **25**, 69 (1978).
- 1978JOR/AIR R. A. Jorge, C. Airoldi, and A. P. Chagas, *J. Chem. Soc. Dalton Trans.* 1102 (1978).
- 1978KAR/KAM N. V. Karyakin, G. P. Kamelova, and V. N. Sapozhnikov, *Termodin. Org. Soedin.* **7**, 52 (1978); *Chem. Abstr.* **92**, 75658d (1980).
- 1978KIS/SUG K. Kishimoto, H. Suga, and S. Seki, *Bull. Chem. Soc. Jpn.* **51**, 1691 (1978).
- 1978KIT/SHR T. Kitazume and J. M. Shreeve, *Inorg. Chem.* **17**, 2173 (1978).
- 1978KOL/DIT V. P. Kolesov and L. N. Dityat'eva, *Termodin. Org. Soedin.* **7**, 44 (1978); *Chem. Abstr.* **92**, 75657c (1980).
- 1978KOM/GUR L. N. Komissarova, M. Z. Gurevich, T. S. Sas, and B. D. Stepin, *Zh. Neorg. Khim.* **23**, 3145 (1978); *Chem. Abstr.* **90**, 61378j (1979).
- 1978KOS/BUD G. Koßmehl and D. Z. Budwell, *Z. Naturforsch. B* **42B**, 478 (1978). (Note: names of the compounds were in German and were translated.)
- 1978LEB/TSV B. V. Lebedev, L. Y. Tsvetkova, and I. B. Rabinovich, *J. Chem. Thermodyn.* **10**, 809 (1978).
- 1978LEB/TSV2 B. V. Lebedev, L. Y. Tsvetkova, I. B. Rabinovich, E. S. Finkel'shtein, and B. S. Strel'chick, *Termodin. Org. Soedin.* **7**, 3 (1978); *Chem. Abstr.* **92**, 83507u (1980).
- 1978MAN M. Mansson, unpublished work as referenced in Ref. [1978MON/ROS].
- 1978MCC/HAM D. G. McCormick and W. S. Hamilton, *J. Chem. Thermodyn.* **10**, 275 (1978).
- 1978MON/ENG R. L. Montgomery, P. S. Engel, R. A. Leckonby, F. D. Rossini, M. Mansson, S. Szilagyi, and J. W. Timberlake, *J. Chem. Eng. Data* **23**, 129 (1978).
- 1978MON/ROS R. L. Montgomery, F. D. Rossini, and M. Mansson, *J. Chem. Eng. Data* **23**, 125 (1978).
- 1978MOU A. H. N. Mousa, *J. Chem. Eng. Data* **23**, 133 (1978).
- 1978NIS/ISH K. Nishida, E. Ishihara, M. Osaka, and M. Koukitu, *Galaxia* **72**, 13 (1978).
- 1978NOW/SZC M. J. Nowak, K. Szczepaniak, A. Barski, and D. Z. Shugar, *Z. Naturforsch. C* **33C**, 876 (1978).
- 1978OSB/SCO A. G. Osborn and D. W. Scott, *J. Chem. Thermodyn.* **10**, 619 (1978).
- 1978RIB/IRV M. A. V. Ribeiro da Silva and R. G. Irving, *Rev. Port. Quim.* **20**, 36 (1978); *Chem. Abstr.* **90**, 211054s (1979).
- 1978SAB/LAF R. Sabbah and M. Laffite, *Bull. Soc. Chim. Fr.* **1**, 50 (1978).
- 1978SAB/LAF2 R. Sabbah and M. Laffite, *Thermochim. Acta* **23**, 196 (1978).
- 1978SHM/SHL G. O. Shmyreva, V. G. Shlyakova, and R. M. Golosova, *Russ. J. Phys. Chem.* **52**, 135 (1978).
- 1978STA/MEW I. Stahl, R. Mews, and O. J. Glemser, *J. Fluorine Chem.* **11**, 455 (1978).
- 1978STE W. V. Steele, *J. Chem. Thermodyn.* **10**, 585 (1978).
- 1978STE2 W. V. Steele, *J. Chem. Thermodyn.* **10**, 919 (1978).
- 1978STE3 W. V. Steele, *J. Chem. Thermodyn.* **10**, 441 (1978).
- 1978STE4 W. V. Steele, *J. Chem. Thermodyn.* **10**, 445 (1978).
- 1978SUN/VIS S. Sundaram and D. S. Viswanath, *J. Chem. Eng. Data* **23**, 63 (1978).
- 1978TAV/NEE A. Tavlaridis and R. Z. Neeb, *Anal. Chem.* **293**, 211 (1978) as quoted in Ref. [1999ZEM/STA].
- 1978TIM R. E. Timms, *Chem. Phys. Lipids* **21**, 113 (1978).
- 1978TRI/VOO L. E. Trimble and R. J. H. Voorhoeve, *Analyst (Cambridge, U.K.)* **103**, 759 (1978).
- 1978VIL/PER R. Vilcu, S. Perisanu, and I. Ciocazanu, *Bull. Inst. Politek "Gheorghe Gheorghiu-Dej" Bucuresti Ser. Chim.-Metal* **40**, 9 (1978); *Chem. Abstr.* **91**, 28258t (1979).
- 1979ABA/MAL A. S. Abakumov and M. L. Malyhev, *Zh. Neorg. Khim.* **24**, 642 (1979).
- 1979ADE/CAV F. A. Adedeji, J. K. Cavell, S. Cavell, J. A. Connor, G. Pilcher, H. A. Skinner, and M. T. Zafarani-Moattar, *J. Chem. Soc., Faraday Trans. 1* **75**, 603 (1979).
- 1979AMA/SAT R. Amano, A. Sato, and S. Suzuki, *Radiochem. Radioanal. Lett.* **39**, 441 (1979); *Chem. Abstr.* **91**, 181625e (1979).
- 1979ARM/JAM N. A. Armstrong, K. C. James, and C. K. Wong, *J. Pharm. Pharmacol.* **31**, 627 (1979).
- 1979BAL/FRI M. Balish and V. J. Fried, *J. Chem. Eng. Data* **24**, 91 (1979).
- 1979BER/ANG G. Berthon, V. Angot, V. Beden, and O. Enea, *J. Chem. Thermodyn.* **11**, 539 (1979).
- 1979BLU/BAE V. I. Bludilina, A. K. Baev, V. K. Matveev, I. L. Gaidym, and E. I. Shcherbina, *Zh. Fiz. Khim.* **53**, 1052 (1979); *Chem. Abstr.* **91**, 9585f (1979).
- 1979BOT/CAM H. Bothe and H. K. Cammenga, *J. Therm. Anal.* **16**, 267 (1979).
- 1979CAL/DIA J. C. G. Calado, A. R. Dias, A. R. Martinho-Simoes, and M. A. V. Riberio da Silva, *J. Organomet. Chem.* **174**, 77 (1979).
- 1979CAR/CLA J. Carre, P. Claudy, J. M. Letoffe, M. Kollmannsberger, and J. Bousquet, *J. Fluorine Chem.* **14**, 139 (1979).
- 1979CAV/GAR K. J. Cavell, C. D. Garner, G. Pilcher, and S. Parks, *J. Chem. Soc. Dalton Trans.* 1714 (1979).
- 1979CAV/HIL K. J. Cavell, J. O. Hill, and R. J. Magee, *Thermochim. Acta* **33**, 155 (1979).
- 1979CAV/HIL2 K. J. Cavell, J. O. Hill, and R. J. Magee, *Thermochim. Acta* **33**, 383 (1979).
- 1979CLA/KNO T. Clark, T. M. Knox, M. A. McKervey, H. Mackle, and J. Rooney, *J. Am. Chem. Soc.* **101**, 2404 (1979).
- 1979COL/JIM M. Colomina, P. Jimenez, R. Perez-Ossorio, and C. Turrión, *J. Chem. Thermodyn.* **11**, 1179 (1979).
- 1979COL/JIM2 M. Colomina, P. Jimenez, M. V. Roux, and C. Turrión, *An. Quim.* **75**, 620 (1979); *Chem. Abstr.* **92**, 75685k (1980).
- 1979CON/DEM J. A. Connor, C. P. Demain, H. A. Skinner, and M. T. Zafarani-Moattar, *J. Organomet. Chem.* **170**, 117 (1979).
- 1979CON/MAR J. A. Connor, J. A. Martinho-Simoes, H. A. Skinner, and M. T. Zafarani-Moattar, *J. Organomet. Chem.* **179**, 331 (1979).
- 1979DAA/ERN H. Daamen, J. M. Ernstring, and A. Oskam, *Thermochim. Acta* **33**, 217 (1979).
- 1979DAA/VAN H. Daamen, H. Van der Poel, D. J. Stufkens, and A. Oskam, *Thermochim. Acta* **34**, 69 (1979).
- 1979DAS/DHA D. Das, S. R. Dharwadkar, and M. S. Chandrasekharaiah, *Thermochim. Acta* **30**, 371 (1979).
- 1979DEK/OON C. G. De Kruif and H. A. Oonk, *J. Chem. Thermodyn.* **11**,

- 287 (1979).
- 1979DEK/VOO C. G. DeKruif, J. Voogd, and J. C. A. Offringa, *J. Chem. Thermodyn.* **11**, 651 (1979).
- 1979FUC/PEA R. Fuchs and L. A. Peacock, *Can. J. Chem.* **57**, 2302 (1979).
- 1979GAR/LAW R. G. Garza, Lawrence Livermore National Laboratory UCRL, 1 (1979); as cited in Ref. [2008RAI/BHA].
- 1979GOL/SHM R. M. Golosova, G. O. Shmyreva, V. G. Shlyakova, A. B. Petrunin, D. B. Bekker, and A. F. Zhigach, *Zh. Fiz. Khim.* **53**, 766 (1979).
- 1979GON/CHA A. S. Goncalves, A. P. Chagas, and C. Airoidi, *J. Chem. Soc. Dalton Trans.* 159 (1979).
- 1979IGU/CHU I. K. Igumenov, Y. V. Chumachenko, and S. V. Zemskov, *Khim. Termodin. Termokhim.*, 65 (1979); *Chem. Abstr.* **91**, 217803c (1979).
- 1979JAK/TZI G. Jakli, P. Tzias, and W. A. Van Hook, *Acta Chim. Acad. Sci. Hung.* **99**, 121 (1979).
- 1979KIM/TAK T. Kimura and S. J. Takagi, *J. Chem. Thermodyn.* **11**, 47 (1979).
- 1979KUC/SKU Y. K. Kuchhal, R. N. Shukla, and A. B. Biswas, *Thermochim. Acta* **31**, 61 (1979).
- 1979KUD/KUD A. P. Kudchadker, S. A. Kudchadker, R. P. Shukla, and P. R. J. Patnaik, *J. Phys. Chem. Ref. Data* **8**, 499 (1979).
- 1979KUD/KUD2S A. Kudchadker, A. P. Kudchadker, and B. J. Zwolinski, *J. Chem. Thermodyn.* **11**, 1051 (1979).
- 1979LAR S. V. Larionov, *Russ. J. Inorg. Chem.* **24**, 802 (1979).
- 1979LEB/BER S. Y. Lebedev, S. S. Berdonosov, I. V. Melikhov, N. B. Mekheev, and I. A. Rumer, *Radiokhimiya* **21**, 470 (1979); *Chem. Abstr.* **91**, 146146c (1979).
- 1979LEE/HOS S. H. Lee-Bechtold, I. A. Hossenlop, D. W. Scott, A. G. Osborn, and W. D. Good, *J. Chem. Thermodyn.* **11**, 469 (1979).
- 1979LET/ORC T. M. Letcher, S. W. Orchard, and M. J. Albers, *J. Chem. Thermodyn.* **11**, 173 (1979).
- 1979MAC/PRA A. B. Macknick and J. M. Prausnitz, *J. Chem. Eng. Data* **24**, 175 (1979).
- 1979MAH/SMI P. J. Maher and B. D. Smith, *J. Chem. Eng. Data* **24**, 16 (1979).
- 1979MAJ/SVO V. Majer, V. Svoboda, S. Hala, and J. Pick, *Collect. Czech. Chem. Commun.* **44**, 637 (1979).
- 1979MAJ/SVO2 V. Majer, V. Svoboda, J. Koubek, and J. Pick, *Collect. Czech. Chem. Commun.* **44**, 3521 (1979).
- 1979MAR/SAC V. S. Markovnik, A. I. Sachek, A. D. Peshchenko, O. V. Shvaro, D. N. Andreevskii, and N. M. Olizarevich, *Termodin. Org. Soedin.*, 107 (1979).
- 1979MAS/MAL R. Masse, R. Malaviole, and A. Chauvet, *J. Therm. Anal.* **16**, 341 (1979).
- 1979MJO C. C. Mjojo, *J. Chem. Soc., Faraday Trans. 2* **75**, 692 (1979).
- 1979MUR/IWA M. Muramatsu, M. Iwahashi, and U. Takeuchi, *J. Pharm. Sci.* **68**, 175 (1979).
- 1979PET/MAJ L. Petros, V. Majer, J. Koubek, V. Svoboda, and J. Pick, *Collect. Czech. Chem. Commun.* **44**, 3533 (1979).
- 1979PRI/SAP S. J. W. Price and H. J. Sapiiano, *Can. J. Chem.* **57**, 685 (1979).
- 1979RIB/REI M. A. V. Riberio da Silva and A. M. M. V. Reis, *Bull. Chem. Soc. Jpn.* **52**, 3080 (1979).
- 1979SAB R. Sabbah, *Bull. Chem. Soc. Jpn.* **9-10**, 434 (1979).
- 1979SAB2 R. Sabbah, C. R. Seances Acad. Sci., Ser. C **289**, 153 (1979); *Chem. Abstr.* **92**, 58049p (1980).
- 1979SAL/PEA P. P. S. Saluja, L. A. Peacock, and R. Fuchs, *J. Am. Chem. Soc.* **101**, 1958 (1979).
- 1979SAN/EPS D. J. Sandman, A. J. Epstein, J. S. Chickos, J. Ketchum, J. S. Fu, and H. A. Scheraga, *J. Chem. Phys.* **70**, 305 (1979).
- 1979SCO/OSB D. W. Scott and A. G. Osborn, *J. Phys. Chem.* **83**, 2714 (1979).
- 1979SEKDES A. Sekiya and D. D. Desmarteau, *J. Fluorine Chem.* **14**, 289 (1979).
- 1979SEV C. Svensson, *J. Chem. Thermodyn.* **11**, 593 (1979).
- 1979SMI G. W. Smith, *Phase Transitions* **1**, 107 (1979).
- 1979SPE/SHO W. F. Spencer, T. D. Shoup, M. M. Cliath, W. J. Farmer, and R. J. Hague, *J. Agric. Food Chem.* **27**, 273 (1979).
- 1979STE W. V. Steele, *J. Chem. Thermodyn.* **11**, 187 (1979).
- 1979SUK/VLA S. I. Sukhova, O. N. Vlasov, and L. V. Li, *Nov. Khim. Sredstva Azsch Rast.* 51 (1979); *Chem. Abstr.* **92**, 83359x (1980).
- 1979SUN/SVE S. Sunner and C. Svensson, *J. Chem. Soc., Faraday Trans. 1* **75**, 2359 (1979).
- 1979SUN/SVE2 S. Sunner, C. Svensson, and A. S. Zelepuga, *J. Chem. Thermodyn.* **11**, 491 (1979).
- 1979TEL/RAB V. I. Tel'noi, I. B. Rabinovich, M. R. Leonov, G. V. Solov'eva, and N. I. Gramoteeva, *Dokl. Phys. Chem.* **245**, 363 (1979).
- 1979THO/MEA L. H. Thomas, R. Meatyrd, H. Smith, and G. H. Davies, *J. Chem. Eng. Data* **24**, 159 (1979).
- 1979YAN/TEP I. K. Yanson, A. B. Teplitsky, and L. F. Sukhodub, *Biopolymers* **18**, 1149 (1979).
- 1979ZAN/THO M. Zander and W. Thomas, *J. Chem. Eng. Data* **24**, 1 (1979).
- 197BAG/MAL A. Baghdoyan, J. Malik, and V. Fried, *J. Chem. Eng. Data* **16**, 96 (1971).
- 1980ABE/SHR T. Abe and J. M. Shreeve, *Inorg. Chem.* **19**, 3063 (1980).
- 1980ABR/IRV M. H. Abraham and R. J. Irving, *J. Chem. Thermodyn.* **12**, 539 (1980).
- 1980AIH A. Aihara, *Denki Tsushin Daigaku Gakuho* **31**, 65 (1980); *Chem. Abstr.* **94**, 174045z (1981).
- 1980AND/PIL F. Andruzzi, G. Pilcher, J. M. Hacking, and S. Cavell, *Makromol. Chem.* **181**, 923 (1980); *Chem. Abstr.* **93**, 8632y (1980).
- 1980ARN/SCH H. Arntz and G. M. Schneider, *Faraday Discuss. Chem. Soc.* **69**, 139 (1980).
- 1980ARS M. R. Arshadi, *J. Chem. Thermodyn.* **12**, 903 (1980).
- 1980BAL/LEB A. A. Balepin, V. P. Lebedev, A. A. Kuznetsova, K. K. Venter, M. A. Trusule, D. O. Lolya, and Y. A. Lebedev, *Izv. Akad. Nauk. SSR, Ser. Khi.* **4**, 848 (1980).
- 1980BAR/BEN G. Bardi, L. Bencivenni, D. Ferro, B. Martini, C. S. Nunziante, and R. Teghil, *Thermochim. Acta* **40**, 275 (1980).
- 1980BECKRA H.-D. Beckhaus, G. Kratt, K. Lay, J. Greiselmann, C. Rüchardt, B. Kitschke, and H. Lindner, *J. Chem. Ber.* **113**, 3441 (1980).
- 1980BOX G. Boxhoorn, Dissertation, University of Amsterdam, 1980, Appendix, p. 149, as cited in Ref. [1980CAV/ERN].
- 1980BOX/ERN G. Boxhoorn, J. M. Ernsting, D. J. Stufkens, and A. Os-kam, *Thermochim. Acta* **42**, 315 (1980).
- 1980BYS K. Bystrom, *J. Chem. Soc., Faraday Trans. 1* **76**, 1986 (1980).
- 1980CAL/DIA J. C. R. Calado, A. R. Dias, M. E. Minas da Piedade, and J. A. Martinho Simoes, *Rev. Port. Quim.* **22**, 53 (1980); see also Ref. [1983JAC/VAN]; *Chem. Abstr.* **96**, 52441w (1981).
- 1980CAR/BUS C. Carfagna, V. Busico, V. Salerno, and M. Vacatello, *Thermochim. Acta* **37**, 31 (1980).
- 1980CAV/ERN K. J. Cavell, J. M. Ernsting, and D. J. Stufkens, *Thermochim. Acta* **42**, 343 (1980).
- 1980CLA/KNO T. Clark, T. M. Knox, M. A. McKervey, and H. Mackle, *J. Chem. Soc., Perkin Trans. 2* 1686 (1980).
- 1980COL/JIM M. Colomina, P. Jimenez, M. V. Roux, and C. Turrion, *J. Therm. Anal. Calorim.* **11**, 3-19-1/3-19-16 (1980); *Chem. Abstr.* **95**, 121878n (1981); see Ref. [1981COL/JIM].
- 1980COL/JIM2 M. Colomina, P. Jimenez, C. Turrion, J. A. Fernandez, and C. Monzon, *An. Quim. Ser. A* **76**, 245 (1980).
- 1980DAV/FIN R. H. Davies, A. Finch, and P. G. Gardner, *J. Chem. Thermodyn.* **12**, 291 (1980).
- 1980DEK C. G. DeKruif, *J. Chem. Thermodyn.* **12**, 243 (1980).
- 1980DEK/GOV C. G. DeKruif and H. A. J. Govers, *J. Chem. Phys.* **73**, 553 (1980).
- 1980DEP R. S. DePablo, *J. Phys. D: Appl. Phys.* **13**, 313 (1980).
- 1980DYA/VAS G. N. D'yakova and I. A. Vasil'ev, *Neftepererab. Neftekhim. (Moscow)*, 51 (1980); *Chem. Abstr.* **92**, 170269m (1980).
- 1980DYG/STE R. S. Dygdala and K. Stefanski, *Chem. Phys.* **53**, 51 (1980).
- 1980FAR/YAN W. J. Farmer, M. S. Yang, J. Letey, and W. F. Spencer, *Soil Sci. Soc. Am. J.* **44**, 676 (1980).

- 1980FER/BEN D. Ferro, L. Bencivenni, R. Teghil, and R. Mastromarino, *Thermochim. Acta* **42**, 75 (1980).
- 1980FER/BEN2 D. Ferro, L. Bencivenni, R. Teghil, and M. Pelino, *J. Indian Chem. Soc.* **57**, 629 (1980).
- 1980FRA/CAS R. Francesconi, C. Castellari, A. Arcelli, and F. Comelli, *Can. J. Chem. Eng.* **58**, 113 (1980).
- 1980FUC/PEA R. Fuchs and L. A. Peacock, *Can. J. Chem.* **58**, 2796 (1980).
- 1980GOM/WEI W. Gombler and H.-U. Weiler, *J. Fluorine Chem.* **15**, 279 (1980).
- 1980JAR/AFA N. L. Jarim-Agaev, L. D. Afanasienko, V. P. Kalinichenko, and G. B. Tolmacheva, *Ukr. Khim. Zhur.* **46**, 1331 (1980).
- 1980JON/MAT J. A. Jonsson, L. Mathiasson, and A.-M. Olsson, *Acta Chem. Scand., Ser. A* **34A**, 147 (1980).
- 1980KRA/PIG A. Krajewska and K. Pigon, *Thermochim. Acta* **41**, 187 (1980).
- 1980LEB/KAL N. D. Lebedeva, Y. A. Kalin, V. L. Raydnenko, V. B. Stapanov, and N. M. Kozlova, *Zh. Prikl. Khim. (Leningrad)* **53**, 2588 (1980); *Chem. Abstr.* **94**, 156270q (1981).
- 1980LEB/KIP N. K. Lebedev, E. G. Kiparisova, B. V. Lebedev, A. M. Sladkov, and N. A. Vasmeva, *Bull. Acad. Sci. USSR Div. Chem. Sci.* **29**, 374 (1980).
- 1980LEB/NAZ N. D. Lebedeva, L. F. Nazarova, and Y. A. Katin, *Zh. Prikl. Khim. (Leningrad)* **53**, 1394 (1980).
- 1980MAJ/SVA V. Majer, L. Svab, and V. Svoboda, *J. Chem. Thermodyn.* **12**, 843 (1980).
- 1980MAJ/SVO V. Majer, V. Svoboda, A. Posta, and J. Pick, *Collect. Czech. Chem. Commun.* **45**, 3063 (1980).
- 1980MAJ/WAN V. Majer, Z. Wanger, V. Svoboda, and V. Cadek, *J. Chem. Thermodyn.* **12**, 387 (1980).
- 1980MEY/AWE E. F. Meyer, M. J. Awe, and R. E. Wagner, *J. Chem. Eng. Data* **25**, 371 (1980).
- 1980MOR/WAT C. T. Mortimer and J. Waterhouse, *J. Chem. Thermodyn.* **12**, 961 (1980).
- 1980MUR/CAV J. P. Murray, K. J. Cavell, and J. O. Hill, *Thermochim. Acta* **36**, 97 (1980).
- 1980NAK/SUG N. Nakamura, H. Suga, and S. Seki, *Bull. Chem. Soc. Jpn.* **53**, 2755 (1980).
- 1980NAS/HWA P. Nasir, S. C. Hwang, and R. Kobayashi, *J. Chem. Eng. Data* **25**, 298 (1980).
- 1980NIG/DEP G. D. Nigam and B. Deppisch, *Z. Kristallogr.* **151**, 185 (1980).
- 1980NIS/SAK K. Nishiyama, N. Sakiyama, S. Seki, H. Horita, T. Otsubo, and S. Misumi, *Bull. Chem. Soc. Jpn.* **53**, 869 (1980).
- 1980OSB/SCO A. G. Osborn and D. W. Scott, *J. Chem. Thermodyn.* **12**, 429 (1980).
- 1980PAT/TOM C. R. Patrick and F. J. Tomes, *J. Fluorine Chem.* **15**, 267 (1980).
- 1980PIT/KIS V. S. Pityugin, T. L. Kislytsyna, R. F. Shakirov, L. N. Sharif'yanova, and E. N. Maslennikov, *Khim. Prom-st. Ser. Khlor'naya Prom-st.*, 11 (1980).
- 1980ROD/KRU G. N. Rodionova, I. V. Krutovskaya, A. N. Rodionov, Y. G. Tuchin, and V. V. Karpov, *Zh. Prikl. Spektrosk.* **32**, 623 (1980).
- 1980ROT A. M. Rothman, *J. Agric. Food Chem.* **28**, 1225 (1980).
- 1980ROT/ORC J. A. Roth and M. Orchin, *J. Organomet. Chem.* **187**, 103 (1980).
- 1980SAB R. Sabbah, *Thermochim. Acta* **35**, 73 (1980).
- 1980SAB/KOM R. Sabbah and S. Komorowski, *Thermochim. Acta* **41**, 379 (1980).
- 1980SAB/SKO R. Sabbah and S. Skoulika, *Thermochim. Acta* **36**, 179 (1980).
- 1980SAB2 R. Sabbah, *Thermochim. Acta* **41**, 33 (1980).
- 1980SAC/HIL J. Sachinides and J. O. Hill, *Thermochim. Acta* **35**, 59 (1980).
- 1980SAT/SAK T. Sato-Toshima, M. Sakiyama, and S. Saki, *Bull. Chem. Soc. Jpn.* **53**, 2762 (1980).
- 1980SHA/SAD L. N. Sharif'yanova, F. M. Sadykova, and S. G. Akhmerova, *Khim. Prom-st. Ser. Khlor'naya Prom-st.*, 16 (1980).
- 1980SHU/VAR P. Shul'tse, R. M. Varushchenko, G. L. Gal'chenko, T. V. Klimova, and V. I. Stanko, *Russ. J. Gen. Chem.* **50**, 1482 (1980); see also *Zh. Obshch. Khim.* **50**, 1818 (1980).
- 1980SMI G. W. Smith, *Mol. Cryst. Liq. Cryst.* **64**, 15 (1980).
- 1980SMI/STE N. K. Smith, R. C. Stewart, Jr., A. G. Osborn, and D. W. Scott, *J. Chem. Thermodyn.* **112**, 919 (1980).
- 1980STE W. V. Steele, *J. Chem. Thermodyn.* **12**, 187 (1980).
- 1980SVO/UCH V. Svoboda, V. Uchtylova, V. Majer, and J. Pick, *Collect. Czech. Chem. Commun.* **145**, 3233 (1980).
- 1980SWA/KWA H. A. Swain, Jr., Y. C.-Y. Kwan, and H.-N. Sung, *J. Phys. Chem.* **84**, 1347 (1980).
- 1980TEP/YAN A. B. Teplitskii, I. K. Yanson, O. T. Glukhova, A. Zielenkiewicz, W. Zielenkiewicz, and K. L. Wierzchowski, *Biophys. Chem.* **11**, 17 (1980).
- 1980THO/SMI L. H. Thomas, H. Smith, and G. H. Davis, *J. Chem. Technol. Biotechnol.* **30**, 476 (1980).
- 1980URB/GIG M. Urbani, R. Gigli, and V. Picente, *J. Chem. Eng. Data* **25**, 97 (1980).
- 1980VAN/PRA C. Van de Rostyne and J. M. Prausnitz, *J. Chem. Eng. Data* **25**, 1 (1980).
- 1980VIL/PER R. Vilcu and S. Perisanu, *Rev. Roum. Chim.* **25**, 619 (1980).
- 1980YEV/LEB A. A. Yevstropov, B. V. Lebedev, Y. G. Kiparisova, V. A. Alekseyev, and G. A. Stashina, *Polym. Sci. U.S.S.R.* **22**, 2685 (1980).
- 1981ABE/SHR T. Abe and J. M. Shreeve, *Inorg. Chem.* **20**, 2894 (1981).
- 1981ABE/SHR2 T. Abe and J. M. Shreeve, *Inorg. Chem.* **20**, 2432 (1981).
- 1981AMA/SAT R. Amano, A. Sato, and S. Suzuki, *Bull. Chem. Soc. Jpn.* **54**, 1368 (1981).
- 1981AMB/ELL D. Ambrose, J. H. Ellender, H. A. Gundry, D. A. Lee, and R. J. Townsend, *J. Chem. Thermodyn.* **13**, 795 (1981).
- 1981AMB/HAL D. Ambrose and D. J. Hall, *J. Chem. Thermodyn.* **13**, 61 (1981).
- 1981BRO/MCE W. Brostow, D. M. McEachern, and J. A. Valdez, *Mater. Chem.* **6**, 187 (1981).
- 1981BYS K. Bystrom, *J. Chem. Thermodyn.* **13**, 139 (1981).
- 1981CAL/DIA J. C. G. Calado, A. R. Dias, M. S. Salema, and J. A. Martinho-Simoes, *J. Chem. Soc. Dalton Trans.* 1174 (1981).
- 1981CAV/CON K. J. Cavell, J. A. Connor, G. Pilcher, M. A. V. Riberio Da Silva, M. D. M. C. Riberio Da Silva, H. A. Skinner, Y. Virmani, and M. T. Zafarami-Moattar, *J. Chem. Soc., Faraday Trans. 1* **77**, 1585 (1981).
- 1981CAV/GAR K. J. Cavell, C. D. Garner, J. A. Martinho-Simoes, G. Pilcher, H. Al-Samman, H. A. Skinner, G. Al-Takhin, I. B. Walton, and M. T. J. Zafarami-Moattar, *J. Chem. Soc., Faraday Trans. 1* **77**, 2927 (1981).
- 1981CHI/GAR J. S. Chickos, D. L. Garin, M. Hitt, and G. Schilling, *Tetrahedron* **37**, 2255 (1981).
- 1981CHI/HYM J. S. Chickos, A. S. Hyman, L. H. Ladon, and J. F. Liebman, *J. Org. Chem.* **46**, 4294 (1981).
- 1981COL/JIM M. Colomina, P. Jimenez, M. V. Roux, and C. Turrión, *An. Quim. Ser. A* **77**, 114 (1981).
- 1981DEK/HER K. G. De Kruif, J. M. Herman, and P. J. Van den Berg, *J. Chem. Eng. Data* **26**, 359 (1981).
- 1981DEK/KUI C. G. De Kruif, T. Kuipers, J. C. van Miltenburg, R. C. F. Schaake, and G. Stevens, *J. Chem. Thermodyn.* **13**, 1081 (1981).
- 1981DEK/SMI C. G. DeKruif, E. J. Smit, and H. A. Gover, *J. Chem. Phys.* **74**, 5838 (1981).
- 1981DEK/VAN C. G. De Kruif, A. C. G. Van Genderen, J. C. W. G. Bink, and H. A. J. Oonk, *J. Chem. Thermodyn.* **13**, 457 (1981).
- 1981DEP R. S. DePablo, *J. Chem. Eng. Data* **26**, 237 (1981).
- 1981DIT/SKO V. E. Ditsent, I. I. Skorokhodov, M. N. Zolotareva, V. I. Savuskina, and B. N. Tabenko, *Zh. Prikl. Khim. (Leningrad)* **54**, 1617 (1981).
- 1981DRA/AZI M. Dragnet-Brughmans, M. Azibi, and R. Bouche, *Proc. Eur. Symp. Thermal Anal.* **2**, 615 (1981).
- 1981DYA/KOR G. N. D'yakova, G. L. Korichev, A. D. Korkhova, T. F. Vasil'eva, and I. A. Vasil'ev, *Zh. Prikl. Khim. (Leningrad)* **54**, 1644 (1981).
- 1981EDW/PRA D. R. Edwards and J. M. Prausnitz, *J. Chem. Eng. Data* **26**, 121 (1981).
- 1981FER/PIA D. Ferro, V. Piacente, and M. Pelino, *Rev. Roum. Chim.*

- 26, 9 (1981).
- 1981GAT/STR D. Gatta, L. Stradella, and P. Venturello, *J. Solution Chem.* **10**, 209 (1981).
- 1981GUS/KAS S. N. Guseinova, V. V. Kas'yanov, and Y. V. Politanskii, *Khim. Prom-st. Ser. Khlor'naya Prom-st.*, 28 (1981); *Chem. Abstr.* **95**, 225804b (1981).
- 1981HAL/COG J. L. Hales, R. C. Cogman, and W. J. Frith, *J. Chem. Thermodyn.* **13**, 591 (1981).
- 1981HOS/SCO I. A. Hossenlopp and D. W. Scott, *J. Chem. Thermodyn.* **13**, 405 (1981).
- 1981HOS/SCO2 I. A. Hossenlopp and D. W. Scott, *J. Chem. Thermodyn.* **13**, 415 (1981).
- 1981HOS/SCO3 I. A. Hossenlopp and D. W. Scott, *J. Chem. Thermodyn.* **13**, 423 (1981).
- 1981INO/ARA M. Inoue, Y. Arai, S. Saito, and J. Suzuki, *J. Chem. Eng. Data* **26**, 287 (1981).
- 1981JOO/ARL H.-J. Joo and W. Arlt, *J. Chem. Eng. Data* **26**, 138 (1981).
- 1981LEB/YEV B. V. Lebedev, A. A. Yevstropov, and Y. G. Kiparisova, *J. Chem. Thermodyn.* **13**, 1185 (1981).
- 1981MAJ/SVO V. Majer, V. Svoboda, A. Posta, and J. Pick, *Collect. Czech. Chem. Commun.* **46**, 817 (1981).
- 1981MAN/SUN M. Mansson and S. Sunner, *J. Chem. Thermodyn.* **13**, 671 (1981).
- 1981MAR/SAC V. S. Markovnik, A. I. Sachek, A. D. Peshchenko, O. V. Shvaro, and D. N. Andreevskii, *Termodin. Org. Soedin. Gor'kii* **10**, 54 (1981).
- 1981MAS/BAR J. Maslowska and J. B. Baranowski, *Chemia Analityczna* **26**, 1017 (1981).
- 1981MAS/OLE T. N. Masalitinova, T. P. Oleinikova, V. L. Ryadnenko, N. N. Kiseleva, and N. D. Lebedeva, *Zh. Prikl. Khim. (Leningrad)* **54**, 1799 (1981).
- 1981NAV/HAU P. Navard and J. M. Haudin, *J. Therm. Anal.* **22**, 107 (1981).
- 1981OGA/SOR K. Ogasahara, M. Sorai, and H. Suga, *Mol. Cryst. Liq. Cryst.* **71**, 189 (1981).
- 1981PAP/ERA T. S. Papina, P. A. Erastov, and V. P. Kolevsov, *J. Chem. Thermodyn.* **13**, 683 (1981).
- 1981PEL/TOM M. Pelino, M. Tomassetti, V. Piacente, and G. D'Ascenzo, *Thermochim. Acta* **44**, 89 (1981).
- 1981PIR/AZA A. I. Pirogov, G. I. Azarova, and V. I. Klopov, *Izv. Vyssh. Uchebn. Zaved. Khim. Khim. Tekhnol.* **24**, 827 (1981); *Chem. Abstr.* **95**, 138901n (1981).
- 1981SAB/MIN R. Sabbah and C. Minadakis, *Thermochim. Acta* **43**, 269 (1981).
- 1981SHC/RUD M. M. Shchedrina, T. A. Rudol'fi, L. O. Mindlin, and K. K. Prilepskaya, *Maslozhirovays Promyshlennost* **10**, 33 (1981).
- 1981SHI/SAI M. Shimizu, Y. Saito, and K. Kusano, *Preprints 17th Conf. Chem. Thermodyn. Thermal Anal., Japan, 1981* (unpublished), p. 50; as cited in Ref. [1985MAJ/SVO].
- 1981SMI/MAR E. V. Smirnov, I. A. Marav'eva, L. I. Martynenko, and V. I. Spitsyn, *Zh. Neorg. Khim.* **26**, 1709 (1981); *Chem. Abstr.* **95**, 86430m (1981).
- 1981SVO/CHA V. Svoboda, V. Charvatova, V. Majer, and J. Pick, *Collect. Czech. Chem. Commun.* **46**, 2983 (1981).
- 1981TEG/FER R. Teghil, D. Ferro, L. Bencivenni, and M. Pelino, *Thermochim. Acta* **44**, 213 (1981).
- 1981TEK/MAJ V. Tekac, V. Majer, V. Svoboda, and V. Hynek, *J. Chem. Thermodyn.* **13**, 659 (1981).
- 1981VAR/BUL R. M. Varushchenko, L. L. Bulgakova, P. S. Minzabekyants, and K. N. Makarov, *Russ. J. Phys. Chem.* **55**, 1480 (1981).
- 1981WES/SIM J. W. Westcott, C. G. Simon, and J. F. Bidleman, *Environ. Sci. Technol.* **15**, 1375 (1981).
- 1981WIE/KOB S. A. Wiczorek and P. Kobayashi, *J. Chem. Eng. Data* **26**, 8 (1981); **26**, 11 (1981).
- 1981WIL/JOH G. M. Wilson, R. H. Johnston, S.-C. Hwang, and C. Tsouopoulos, *Ind. Eng. Chem. Process Des. Dev.* **20**, 94 (1981).
- 1981YAN/NAB S. Yano, Y. Nabata, and K. Aoki, *Mol. Cryst. Liq. Cryst.* **70**, 163 (1981).
- 1982AIR/CHA C. Airoidi, A. P. Chagas, and F. P. Assuncao, *Can. J. Chem.* **60**, 2132 (1982).
- 1982BAL/MRA M. Balty, S. Mraw, L. A. K. Staveley, A. H. Overs, M. C. Owen, R. K. Thomas, and J. W. White, *Mol. Phys.* **45**, 1015 (1982).
- 1982BYS K. Bystroem, *J. Chem. Thermodyn.* **14**, 865 (1982).
- 1982BYS/MAN K. Bystrom and J. Mansson, *J. Chem. Soc., Perkin Trans. 2* **565** (1982).
- 1982CAS/FRA C. Castellari, R. Francesconi, and F. Comelli, *J. Chem. Eng. Data* **27**, 156 (1982).
- 1982COL/JIM M. Colomina, P. Jimenez, and C. Turrión, *J. Chem. Thermodyn.* **14**, 779 (1982).
- 1982CON/ZAF J. A. Connor, M. T. Zafarani-Moattar, J. Bickerton, N. I. El-Saied, S. Suradi, R. Carson, G. Al Takkhin, and H. A. Skinner, *Organometallics* **1**, 1166 (1982).
- 1982CUE/SOL M. A. Cuevas, X. Solans, and L. Artus, *Afinidad* **39**, 406 (1982).
- 1982DEK/BLO C. G. DeKruif and J. G. Blok, *J. Chem. Thermodyn.* **14**, 201 (1982).
- 1982DEK/SCH G. C. DeKruif, R. C. F. Schaake, J. C. Van Meltenburg, K. Van der Klauw, and J. G. Blok, *J. Chem. Thermodyn.* **14**, 791 (1982).
- 1982DEM/FOX R. A. De Marco and W. B. Fox, *J. Org. Chem.* **47**, 3772 (1982).
- 1982DEP R. S. DePablo, *J. Chem. Eng. Data* **27**, 374 (1982).
- 1982DIA/SAL A. R. Dias, M. S. Salema, and J. A. Martinho Simoes, *Organometallics* **1**, 971 (1982).
- 1982DWO/FUC A. Dworkin, A. H. Fuchs, M. Ghelfenstein, and H. Szwarc, *J. Physique Lettres* **43**, L21 (1982).
- 1982FUC/HAL R. Fuchs, J. H. Hallman, and M. O. Perlman, *Can. J. Chem.* **60**, 832 (1982).
- 1982FUC/PEA R. Fuchs, L. A. Peacock, and W. K. Stephenson, *Can. J. Chem.* **60**, 1953 (1982).
- 1982FUR/SAK J. Furukawa, M. Sakiyama, S. Seki, Y. Saito, and K. Kusano, *Bull. Chem. Soc. Jpn.* **55**, 3329 (1982).
- 1982GAM/CAL B. E. Gammon, J. E. Callanan, I. A. Hossenlopp, A. G. Osborn, and W. D. Good, *Proc. Symp. Thermophys. Prop.* **8**, 402 (1982).
- 1982GRA/FOS B. T. Grayson and L. A. Fosbraey, *Pestic. Sci.* **13**, 269 (1982).
- 1982GUT/KNA B. Gutsche and H. Knapp, *Fluid Phase Equilib.* **8**, 285 (1982).
- 1982HON/WAK G. S. Hong, R. Waksak, H. Finston, and V. Fried, *J. Chem. Eng. Data* **27**, 146 (1982).
- 1982INA/MUR S. Inagaki, S. Murata, and M. Sakiyama, *Bull. Chem. Soc. Jpn.* **55**, 2808 (1982).
- 1982INA/MUR2 S. Inagaki, S. Murata, M. Sakiyama, Y. Ito, Y. Umehara, T. Hijiya, and T. Matsuura, *Bull. Chem. Soc. Jpn.* **55**, 2803 (1982).
- 1982INI/LOP J. C. Iniguez, M. E. Lopez, and D. M. McEachern, *Rev. Soc. Quim. Mex.* **26**, 122 (1982); *Chem. Abstr.* **97**, 144252 j (1982).
- 1982INV C. Invernizzi, *Termotecnica* **4**, 78 (1982).
- 1982JAL/ZOG I. M. Jalal, G. Zografi, A. K. Rakshit, and F. D. Gunston, *Chem. Phys. Lipids* **31**, 395 (1982).
- 1982JOC/DEK R. Jochems, H. Dekker, C. Mosselman, and G. Somsen, *J. Chem. Thermodyn.* **14**, 395 (1982).
- 1982JOC/DEK2 R. Jochems, H. Dekker, C. Mosselman, and G. Somsen, *J. Chem. Thermodyn.* **14**, 799 (1982).
- 1982KAR/SHV N. V. Karyakin, K. G. Shvetsova, and E. S. Dzharimova, *Termodin. Org. Soedin.* **33** (1982); *Chem. Abstr.* **99**, 129172f (1983).
- 1982LEB/BYK B. V. Lebedev, T. A. Bykova, N. N. Smirnova, and T. G. Kulagina, *Zh. Obshch. Khim.* **52**, 2630 (1982).
- 1982MAR/AND J. F. Martin and R. J. L. Andon, *J. Chem. Thermodyn.* **14**, 679 (1982).
- 1982MAR/MIR A. Martin and M. J. Miralles, *J. Pharm. Sci.* **71**, 439 (1982).
- 1982MEN J. Menaucourt, *J. Chim. Phys. Phys.-Chim. Biol.* **79**, 531 (1982).
- 1982MIN/SAB C. Minadakis and R. Sabbah, *Thermochim. Acta* **55**, 147 (1982).
- 1982MOR C. T. Mortimer, in *Thermochemistry and Its Applications*

- to *Chemical and Biochemical Systems*, Series C: Mathematical and Physical Sciences Vol. 119, edited by Riberio Da Silva (Reidel, Boston, MA, 1982), pp 47–60.
- 1982MUR/SAK S. Murata, M. Sakiyama, and Seki, *J. Chem. Thermodyn.* **14**, 707, 723–731 (1982).
- 1982PIL/SKI G. Pilcher and H. A. Skinner, in *The Chemistry of the Metal-Carbon Bond*, edited by F. R. Hartley and S. Patai (Wiley, New York, 1982), Chap. 2.
- 1982POE/FAN G. Poeti, E. Faneli, and M. Braghetti, *J. Therm. Anal.* **24**, 273 (1982).
- 1982REC/GRE P. Reche and M.-F. Grenier-Loustalot, *J. Chromatogr.* **238**, 317 (1982).
- 1982ROO F. L. Rook, *J. Chem. Eng. Data* **27**, 72 (1982).
- 1982SAB/TOR R. Sabbah and L. A. Torres Gomez, *Thermochim. Acta* **52**, 285 (1982).
- 1982SAC/PES A. I. Sachek, A. D. Peshenko, V. S. Markovnik, O. V. Ral'ko, D. N. Andreevskii, and A. A. Leont'eva, *Termodin. Org. Soedin.*, 94 (1982).
- 1982SUR/SAI S. Suradi, N. El Saïad, G. Pilcher, and H. A. Skinner, *J. Chem. Thermodyn.* **14**, 45 (1982).
- 1982SVO/CHA V. Svoboda, V. V. Charvatova, V. Majer, and V. Hynek, *Collect. Czech. Chem. Commun.* **47**, 543 (1982).
- 1982TOR/SAB L. A. Torres Gomez and R. Sabbah, *Thermochim. Acta* **57**, 67 (1982).
- 1982TOR/SAB2 L. A. Torres Gomez and R. Sabbah, *Thermochim. Acta* **58**, 311 (1982).
- 1982TOU/OKA H. Touhara, S. Okazaki, F. Okino, H. Tanaka, K. Ikari, and K. Nakanishi, *J. Chem. Thermodyn.* **14**, 145 (1982).
- 1982VAR/PUC R. M. Varushchenko, S. S. Puchkov, and A. I. Druzina, *Zh. Fiz. Khim.* **56**, 2934 (1982).
- 1982WAY M. Wayaku, *Aroamatikku* **34**, 219 (1982).
- 1982WRE/VIK D. J. Wren and A. C. Vikis, *J. Chem. Thermodyn.* **14**, 435 (1982).
- 1982ZVE/VIN V. V. Zvezdina, E. A. Vinogradova, V. P. Bochinn, B. D. Berezin, and O. A. Golubchikov, *Zh. Neorg. Khim.* **27**, 2818 (1982).
- 1983AKK/SCH O. S. Akkerman, G. Schat, E. A. I. M. Evers, and F. Bickelhaupt, *Recueil. J. Royal Netherlands Chem. Soc.* **102**, 109 (1983).
- 1983ALT/CON G. Al-Takhin, J. A. Connor, and H. A. Skinner, *J. Organomet. Chem.* **259**, 313 (1983).
- 1983ALT/PIL G. Al-Takhin, G. Pilcher, J. Bickerton, and A. A. Zaki, *J. Chem. Soc. Dalton Trans.* 2657 (1983).
- 1983AN/MAN X.-W. An and M. Mansson, *J. Chem. Thermodyn.* **15**, 287 (1983).
- 1983BAR/BEC W. Barbe, H.-D. Beckhaus, H. J. Lindner, and C. Rüchardt, *Chem. Ber.* **116**, 1017 (1983).
- 1983BEN/BIE R. Bender, V. Bieling, and G. Mauer, *J. Chem. Thermodyn.* **15**, 585 (1983).
- 1983CAS/PON A. E. Castro Luna, M. I. Ponzi, and J. B. Rivarola, *J. Chem. Eng. Data* **28**, 349 (1983).
- 1983CHA S. S. Chang, *J. Chem. Phys.* **79**, 6229 (1983).
- 1983CLA/COR R. M. Clay, S. Corr, G. Keenan, and W. V. Steele, *J. Am. Chem. Soc.* **105**, 2070 (1983).
- 1983COL/JIM M. Colomina, P. Jimenez, C. Turrion, M. Kaminski, and W. Zielenkiewicz, *Thermochim. Acta* **68**, 371 (1983).
- 1983DEK/VAN C. G. DeKruif, J. C. Van Miltenburg, and J. G. Blok, *J. Chem. Thermodyn.* **15**, 129 (1983).
- 1983DEW/BOW H. G. M. DeWit, J. G. Bowstra, and C. G. DeKruif, *J. Chem. Phys.* **78**, 1470 (1983).
- 1983DEW/DEK H. G. M. De Wit, C. G. De Kruif, and J. C. Van Miltenburg, *J. Chem. Thermodyn.* **15**, 891 (1983).
- 1983DEW/VAN H. G. M. DeWit, J. C. Van Miltenburg, and C. G. DeKruif, *J. Chem. Thermodyn.* **15**, 651 (1983).
- 1983FER/IMP D. Ferro, P. Imperatori, and C. Quagliata, *J. Chem. Eng. Data* **28**, 242 (1983).
- 1983FER/PIA D. Ferro, V. Piacente, and P. Scardala, *Thermochim. Acta* **68**, 329 (1983).
- 1983FER/QUA D. Ferro, C. Quagliata, and M. R. Conte, *Thermochim. Acta* **60**, 211 (1983).
- 1983FOK/VAN J. G. Fokkens, J. G. M. Van Amelsfoort, C. J. De Blaey, C. G. De Kruif, and J. Wilting, *Int. J. Pharm.* **14**, 79 (1983).
- 1983FUC/HAL R. Fuchs and J. H. Hallman, *Can. J. Chem.* **61**, 503 (1983).
- 1983GEI/NUR Kh. I. Geidarov, G. G. Nurullaev, M. S. Salakhov, Z. Guseinov, and K. A. Karasharli, *Azerb. Khim. Zh.*, 120 (1983).
- 1983GRA/ABO D. J. Grant and I. K. A. Abougela, *Int. J. Pharm.* **17**, 77 (1983).
- 1983HAL/STE J. H. Hallman, W. K. Stepenson, and R. Fuchs, *Can. J. Chem.* **61**, 2044 (1983).
- 1983HOL M. R. Holdiness, *Thermochim. Acta* **68**, 375 (1983).
- 1983HOU J. E. House, Jr., *J. Fluorine Chem.* **22**, 299 (1983).
- 1983HOU2 J. E. House, Jr., *Thermochim. Acta* **71**, 215 (1983).
- 1983JAC/VAN M. H. G. Jacobs, P. J. van Ekeren, and C. G. DeKruif, *J. Chem. Thermodyn.* **15**, 619 (1983).
- 1983KHA.KHE M. S. Khanna, S. C. Khetarpal, K. Lal, and H. L. Bhatnagar, *Indian J. Pure Appl. Phys.* **20**, 503 (1983).
- 1983KRA/BEC G. Kratt, H.-D. Bechhaus, W. Bernloehr, and C. Rüchardt, *Thermochim. Acta* **62**, 279 (1983).
- 1983MAC V. Machat, Thesis Utzcht, Prague, 1983.
- 1983MAJ/AZZ J. R. Majer and A. S. P. Azzouz, *J. Chem. Soc., Faraday Trans. 1* **79**, 675 (1983).
- 1983MAS/CHA J. Masse and A. Chauvet, *Ann. Pharm. Fr.* **41**, 579 (1983).
- 1983MAS/STE H. Maskill and W. V. Steele, *J. Chem. Thermodyn.* **15**, 481 (1983).
- 1983MUR/HIL J. P. Murray and J. O. Hill, *Thermochim. Acta* **63**, 211 (1983).
- 1983NAT/VIS G. Natarajan and D. S. Viswanath, *Rev. Sci. Instrum.* **54**, 1175 (1983).
- 1983OI/SHU T. Oi, J. Shulman, A. Popowicz, and T. Ishida, *J. Phys. Chem.* **87**, 3153 (1983).
- 1983OLS/JON A.-M. Olsson, J. A. Jonsson, B. Thelin, and T. Liljefors, *Chem. Ecol.* **9**, 375 (1983).
- 1983ORO/MRA D. R. O'Rourke and S. C. Mraw, *J. Chem. Thermodyn.* **15**, 489 (1983).
- 1983PAL/CHO M. Palczewska-Tulinska, J. Cholinski, A. Szafranski, and D. Wyrzykowska-Stankiewicz, *Fluid Phase Equilib.* **11**, 233 (1983).
- 1983PEL/GIG M. Pelino, R. Gigli, and M. Tomassette, *Thermochim. Acta* **61**, 301 (1983).
- 1983PRI/WOO J. G. Priest, E. M. Wooley, J. B. Ott, and J. R. Goates, *J. Chem. Thermodyn.* **15**, 357 (1983).
- 1983RAE/SCH A. Raemy and T. F. Schweizer, *J. Therm. Anal.* **28**, 95 (1983).
- 1983RED/MUR G. Om. Reddy, B. K. M. Murali, and A. K. Chatterjee, *Propellants, Explos., Pyrotech.* **8**, 29 (1983).
- 1983RIB/REI M. A. V. Ribeiro Da Silva, and A. M. M. V. Reis, *J. Chem. Thermodyn.* **15**, 957 (1983).
- 1983SCH/STR T. Schmeling and R. Strey, *Ber. Bunsenges. Phys. Chem.* **87**, 871 (1983).
- 1983SIV/KOB A. Sivaraman and R. Kobayashi, *J. Chem. Thermodyn.* **15**, 1127 (1983).
- 1983SIV/MAR A. Sivaraman, R. J. Martin, and R. Kobayashi, *Fluid Phase Equilib.* **12**, 175 (1983).
- 1983SKO/SAB S. Skoulika and R. Sabbah, *Thermochim. Acta* **61**, 203 (1983).
- 1983SON/ZOL W. J. Sonnefeld, W. H. Zoller, and W. E. May, *Anal. Chem.* **55**, 275 (1983).
- 1983SPE/CLI W. F. Spencer and M. M. Cliath, *Residue Rev.* **85**, 57 (1983).
- 1983SPI/KOL V. I. Spitsyn, I. D. Kolli, E. A. Balabanova, E. V. Smirnov, and I. A. Murav'eva, *Zh. Neorg. Khim.* **28**, 2697 (1983).
- 1983TAM/DRA A. Tamir, C. Dragoescu, A. Apelblat, and J. Wisniak, *Fluid Phase Equilib.* **10**, 9 (1983).
- 1983TRE/BER G. V. Trembovetskii, S. S. Berdonosov, I. Murav'eva, and L. I. Martynenko, *Zh. Neorg. Khim.* **28**, 3032 (1983); *Chem. Abstr.* **100**, 39999r (1984).
- 1983TSO/WIL C. Tsionopoulos and G. M. W. Wilson, *AIChE J.* **29**, 990 (1983).
- 1983UCH/MAJ V. Uchytlova, V. Majer, V. Svoboda, and V. Hynek, *J. Chem. Thermodyn.* **15**, 853 (1983).
- 1983VAN/JAC P. J. Van Ekeren, M. H. G. Jacobs, J. C. A. Offringa, and C. G. DeKruif, *J. Chem. Thermodyn.* **15**, 409 (1983).

- 1983VAN/KAC Yu. Ya. Van-Chin-Syan, N. S. Kachurina, G. A. Petrovskaya, and S. K. Chuchmarev, *Russ. J. Phys. Chem.* **57**, 1751 (1983).
- 1983VAS T. F. Vasel'eva, in *Teplofiz Svoistva Uglevodorodov Nefteprod*, edited by G. I. Cherednichenko (Ts NIIT Eneftekhim, Moscow, 1983) pp. 85–94.
- 1983ZAM/KAI M. Zamkanej, J. H. Kaiser, H. Birkhofer, H.-D. Beckhaus, and C. Rüchardt, *Chem. Ber.* **116**, 3216 (1983).
- 1983ZIE/ZIE A. Zielenkiewicz, W. Zielenkiewicz, M. Colomina, P. Jimenez, E. Cesari, A. Sanahiya, and V. Torra, *An. Quim. Ser. A* **79** [3 Suppl. 2], 677 (1983); *Chem. Abstr.* **101**, 22920f (1984).
- 1984ALT/CON G. Al-Takhin, J. A. Connor, H. A. Skinner, and M. T. Zafarani-Moattan, *J. Organomet. Chem.* **260**, 189 (1984).
- 1984BAE/BAR A. K. Baev and A. A. Barkatin, *Russ. J. Phys. Chem.* **58**, 195 (1984).
- 1984BAE/BAR2 A. K. Baev, A. A. Barkatin, and L. M. Dyagileva, *Vesti Akad. Navuk BSSR Ser. Khim.*, 60 (1984).
- 1984BAE/MIK A. K. Baev, V. E. Mikhailov, and A. A. Baev, *Vesti Akad. Navuk BSSR Ser. Khim. Navuk*, 75 (1984).
- 1984BEC/RUC H.-D. Beckhaus, C. Rüchardt, and M. Smisek, *Thermochim. Acta* **79**, 149 (1984).
- 1984BER/BEC W. Bernloehr, H.-D. Beckhaus, K. Peters, H. G. Von Schnering, and C. Ruechardt, *Chem. Ber.* **117**, 1013 (1984).
- 1984BER/BEC2 W. Bernloehr, H.-D. Beckhaus, and C. Rüchardt, *Chem. Ber.* **117**, 1026 (1984).
- 1984BIC/MIN J. Bickerton, M. E. Minas da Piedade, and G. Pilcher, *J. Chem. Thermodyn.* **16**, 661 (1984).
- 1984BIC/PIL J. Bickerton, G. Pilcher, and G. Al-Takhin, *J. Chem. Thermodyn.* **16**, 373 (1984).
- 1984BOS/TUR S. R. Bosco and A. G. Turner, *J. Chem. Eng. Data* **29**, 242 (1984).
- 1984BOU/FRI T. Boublik, V. Fried, and E. Hala, *The Vapour Pressures of Pure Substances: Selected Values of the Temperature Dependence of the Vapour Pressures of Some Pure Substances in the Normal and Low Pressure Region*, 2nd ed. (Elsevier, Amsterdam, 1984). The vaporization enthalpies were calculated from the vapor pressures obtained from the Antoine constants reported in this compendium. In cases where the Antoine constant $C=0$, the Antoine Equation ($\log_{10} P=A-B/(C+T)$) reduces to the integrated form of the Clausius Clapeyron equation directly. This was the case for many vaporization enthalpies. In those cases where $C=0$, the vaporization enthalpy was calculated as $\Delta_{\text{vap}}H_m(T)=2.303 R B[T/(T+C)]^2$.
- 1984BOU/OON J. A. Bouwstra, H. A. J. Oonk, J. G. Blok, and C. G. De Kruij, *J. Chem. Thermodyn.* **16**, 403 (1984).
- 1984BUR/ARM L. P. Burkhard, D. E. Armstrong, and A. Andren, *J. Chem. Eng. Data* **29**, 248 (1984).
- 1984BUR/MOR P. M. Burkinshaw and C. T. Mortimer, *J. Chem. Soc. Dalton Trans.* 75 (1984).
- 1984CAR A. S. Carson, *J. Chem. Thermodyn.* **16**, 427 (1984).
- 1984CAR/SPE A. S. Carson and J. A. Spencer, *J. Chem. Thermodyn.* **16**, 423 (1984).
- 1984CAS/FRA C. Castellari, R. Francesconi, and F. Comelli, *J. Chem. Eng. Data* **29**, 126 (1984).
- 1984CAS/FRA2 C. Castellari, R. Francesconi, and F. Comelli, *J. Chem. Eng. Data* **29**, 90 (1984).
- 1984CAS/FRA3 C. Castellari, R. Francesconi, F. Comelli, and S. Ottani, *J. Chem. Eng. Data* **29**, 283 (1984).
- 1984CER/BOU I. Cervenková and T. Boublik, *J. Chem. Eng. Data* **29**, 425 (1984).
- 1984COL/JIM M. Colomina, P. Jimenez, M. V. Roux, and C. Turrion, *J. Chem. Thermodyn.* **16**, 379 (1984).
- 1984COL/JIM2 M. Colomina, P. Jimenez, M. V. Roux, and C. Turrion, *J. Chem. Thermodyn.* **16**, 1121 (1984).
- 1984COR E. H. P. Cordfunke, *Thermochim. Acta* **108**, 45 (1984).
- 1984DOM/EVA E. S. Domalski, W. H. Evans, and E. D. Hearing, *J. Phys. Chem. Ref. Data Suppl.* **13**, 1 (1984).
- 1984EBE/FRA H. Ebeling and E. U. Franck, *Ber. Bunsenges. Phys. Chem.* **88**, 862 (1984).
- 1984ENG/SAN R. Eng and S. I. Sandler, *J. Chem. Eng. Data* **29**, 156 (1984).
- 1984EUB/CED P. T. Eubank, L. E. Cediel, J. C. Holste, and K. R. Hall, *J. Chem. Eng. Data* **29**, 389 (1984).
- 1984FLA/BEC M. A. Flamm-Ter Meer, H.-D. Beckhaus, and C. Rüchardt, *Thermochim. Acta* **80**, 81 (1984).
- 1984FOU/AMO M. Foulon, J. P. Amoureux, J. L. Sauvajol, J. P. Cavrot, and M. Muller, *J. Phys. C* **17**, 4213 (1984).
- 1984GOL/LEV G. S. Gol'din, E. I. Levites, A. I. L'vov, and N. L. Sin'ko, *Koord. Khim.* **20**, 563 (1984).
- 1984GOV/KAN S. W. Govorchin and A. S. Kana'an, *J. Chem. Thermodyn.* **16**, 437 (1984).
- 1984GOV/KAN2S W. Govorchin, A. S. Kana'an, and J. M. Kanameuller, *J. Chem. Thermodyn.* **16**, 703 (1984).
- 1984GRI/KON A. A. Grigor'ev, Yu. V. Konrat'ev, and A. V. Suvorov, *Zh. Obshch. Khim.* **54**, 1935 (1984); as cited in D. R. Kirklin and E. S. Domalski, *J. Chem. Thermodyn.* **20**, 743 (1988).
- 1984HES/WIS W. Hessler, *Wissenschaftliche Zeitschrift der Wilhelm-Piech Universität, Rostoch* **33**, 9 (1984).
- 1984HOL M. R. Holdiness, *Thermochim. Acta* **78**, 435 (1984).
- 1984HUL/LU D. M. Hull and B. C.-Y. Lu, *J. Chem. Eng. Data* **29**, 417 (1984).
- 1984KAR/KRU V. V. Karpov, I. V. Krutovskaya, and G. N. Rodionova, *Zh. Prikl. Khim. (Leningrad)* **57**, 1348 (1984).
- 1984KAR/ROD V. V. Karpov, G. N. Rodionova, I. V. Krutovskaya, L. Z. Gandel'sman, L. A. Khomenko, and L. M. Yagupol'skii, *Dyes Pigm.* **5**, 285 (1984).
- 1984KAT/HAR H. Katayama and Y. Harada, *J. Chem. Eng. Data* **29**, 373 (1984).
- 1984KEL S. M. Kelly, *Helv. Chim. Acta* **67**, 1572 (1984).
- 1984KER/OPP von F. Kersten and H. Oppermann, *Z. Phys. Chem. (Leipzig)* **265**, 1207 (1984).
- 1984KIM/WOO Y. H. Kim, J. E. Woodrow, and J. N. Seiber, *J. Chromatogr.* **314**, 37 (1984).
- 1984KOZ/TIM M. P. Kozina, L. P. Timofeeva, G. L. Gal'chenko, E. S. Balenkova, and M. D. Ordubadi, *Vestn. Mosk. Univ., Ser. 2: Khim.* **25**, 364 (1984); *Chem. Abstr.* **101**, 190955q (1984).
- 1984KRI G. Krien, *Thermochim. Acta* **81**, 29 (1984).
- 1984LEB/GUT N. D. Lebedeva, N. M. Gutner, N. M. Katin, Yu. A. Yu, N. M. Kozlova, N. N. Kiseleva, E. F. Machinerya, and S. L. Dobychin, *Zh. Prikl. Khim. (Leningrad)* **57**, 2297 (1984).
- 1984LEB/GUT2 N. V. Lebedeva, N. M. Gutner, Yu. A. Katin, N. M. Kozlova, N. N. Kiseleva, E. F. Makhina, and S. L. Dobychin, *Russ. J. Phys. Chem.* **57**, 2118 (1984).
- 1984LAW/ELI A. H. Lawrence, L. Elias, and M. Authier-Martin, *Can. J. Chem.* **62**, 1886 (1984).
- 1984MAJ/SVO V. Majer, V. Svoboda, and V. Hynek, *J. Chem. Thermodyn.* **16**, 1059 (1984).
- 1984MAJ/SVO2 V. Majer, V. Svoboda, and M. Lencka, *J. Chem. Thermodyn.* **16**, 1019 (1984).
- 1984MAJ/SVO3 V. Majer, V. Svoboda, J. Pechacek, and S. Hala, *J. Chem. Thermodyn.* **16**, 567 (1984).
- 1984MAJ/UCH V. Majer, V. Uchytilova, V. Svoboda, and A. Posta, *J. Chem. Thermodyn.* **16**, 761 (1984).
- 1984MAR/MEL M. Marcos, E. Melendez, J. L. Serrano, P. Camps, M. Figueredo, and C. Jaime, *J. Chem. Soc., Perkin Trans. 2* **7** (1984).
- 1984MIC/JOS M.-A. Michou-Saucet, J. Jose, C. Michou-Saucet, and J. C. Merlin, *Thermochim. Acta* **75**, 85 (1984).
- 1984MOU A. H. N. Mousa, *J. Fluorine Chem.* **25**, 165 (1984).
- 1984MRA/KEW S. C. Mraw and C. F. Keweshan, *J. Chem. Thermodyn.* **16**, 873 (1984).
- 1984MRW/STA A. Mrwa, M. Starke, and C. Harmann, *Z. Phys. Chem. (Leipzig)* **265**, 577 (1984).
- 1984MUR/HIL J. P. Murray and J. O. Hill, *Thermochim. Acta* **72**, 341 (1984).
- 1984NUR/MEK G. G. Nurullaev, S. A. Mekhtiev, K. A. Karasharli, N. F. Musaeva, and M. S. Salakhov, *Azerb. Khim. Zh.*, 132 (1984).
- 1984PAL/CHO M. Palczewska-Tulinska, J. Cholinski, A. M. Szafranski, and D. Wyrzkowska-Stankiewicz, *Fluid Phase Equilib.* **15**,

- 295 (1984).
- 1984PER R. R. Perron, *Revue Francaise des CORPS GRAS* **31**, 171 (1984).
- 1984PIN/POS N. Pingel, U. Poser, and A. Wurflinger, *J. Chem. Soc., Faraday Trans. 1* **80**, 3221 (1984).
- 1984PUT/IVA S. Putcha, R. V. Ivanturi, and R. Machiraju, *J. Chem. Eng. Data* **29**, 135 (1984).
- 1984RIB/RIB M. D. M. C. Ribeiro Da Silva, M. A. V. Ribeiro Da Silva, A. P. S. M. C. Carvalho, M. J. Akello, and G. Pilcher, *J. Chem. Thermodyn.* **16**, 137 (1984).
- 1984RIB/RIB2 M. D. M. C. Ribeiro Da Silva, M. A. V. Ribeiro Da Silva, and G. Pilcher, *J. Chem. Thermodyn.* **16**, 1149 (1984).
- 1984SAC/MAR A. I. Sachek, V. S. Markovnik, A. D. Peshchenko, A. V. Shvaro, and D. N. Andreevskii, *Khim. Prom-st. (Moscow)*, 337 (1984).
- 1984SHC/KAP A. E. Shcherbina, L. M. Kaporovskii, and E. A. Shcherbina, *Zh. Prikl. Khim. (Leningrad)* **57**, 1910 (1984); *Chem. Abstr.* **101**, 177895a (1984).
- 1984SIP/WIE J. T. Sipowska and S. A. Wiczorek, *J. Chem. Thermodyn.* **16**, 693 (1984).
- 1984SIV/KOB A. Sivaraman and R. Kobayashi, *J. Chem. Eng. Data* **29**, 375 (1984).
- 1984SOK/BAE A. E. Sokolovskii and A. K. Baev, *Russ. J. Phys. Chem.* **58**, 1635 (1984).
- 1984SOK/BAE2 A. E. Sokolovskii and A. K. Baev, *Vesti Akad. Navuk BSSR Ser. Khim. Navuk*, 115 (1984).
- 1984STA/WAD P. Starzewski, I. Wadso, and W. Zielenkiewicz, *J. Chem. Thermodyn.* **16**, 331 (1984).
- 1984STE/ALI A. V. Steblevskii, A. S. Alikhanyan, L. G. Vedenkina, A. V. Shtemenko, A. S. Kotel'nikova, and V. I. Gorgoraki, *Koord. Khim.* **10**, 72 (1984).
- 1984SUR E. Surova, *Chem. Zvesti* **38**, 93 (1984).
- 1984TRE/BER G. V. Trembovetskii, S. S. Berdonosov, I. A. Murav'eva, and L. I. Martynenko, *Zh. Neorg. Khim.* **29**, 2159 (1984); *Chem. Abstr.* **101**, 137451w (1984).
- 1984UCH/MAJ V. Uchytlova, V. Majer, V. Svoboda, and I. Hemer, *J. Chem. Thermodyn.* **16**, 475 (1984).
- 1984VIS/HIL R. Viswanathan and K. Hilpert, *Ber. Bunsenges. Phys. Chem.* **88**, 125 (1984).
- 1984VOR/MIR V. P. Vorob'eva, E. A. Miroshnichenko, I. I. Solomennikova, F. I. Zelchan, E. Lukevits, and Yu. A. Lebedev, *Bull. Acad. Sci. USSR Div. Chem. Sci.* **33**, 1372 (1984).
- 1984WEI/LEF D. I. Weinstein, A. J. Leffler, and J. A. Currie, *Mol. Cryst. Liq. Cryst.* **104**, 95 (1984).
- 1984WIS/TAM J. Wisniak and A. Tamir, *J. Chem. Eng. Data* **29**, 19 (1984).
- 1984ZIE/ZIE A. Zielenkiewicz, W. Zielenkiewicz, L. F. Sukhodub, O. T. Glukhova, A. B. Teplitsky, and K. L. Wierzchowski, *J. Solution Chem.* **13**, 757 (1984).
- 1985APP/MEN E. H. Appelman, M. H. Mendelsohn, and H. Kim, *J. Am. Chem. Soc.* **107**, 6515 (1985).
- 1985BAR/CAS G. Barone, G. Castronuovo, G. Della Gatta, V. Elia, and A. Iannone, *Fluid Phase Equilib.* **21**, 157 (1985).
- 1985BOU/DEL J. A. Bouwstra, V. V. De Leeuw, and J. C. van Miltenburg, *J. Chem. Thermodyn.* **17**, 685 (1985).
- 1985BRO/INI W. Brostow, J. C. Iniguez, M. A. Shmorhun, and J. A. Valdez, *Mater. Chem. Phys.* **12**, 557 (1985).
- 1985CAM/FER A. R. Campanelli, D. Ferro, and N. V. Pavel, *Thermochim. Acta* **87**, 231 (1985); *Chem. Abstr.* **103**, 93861q (1985).
- 1985CAO/BAC J.-R. Cao and R. A. Back, *Can. J. Chem.* **63**, 2945 (1985).
- 1985CAR/LAY A. S. Carson, P. G. Laye, and J. A. Spencer, *J. Chem. Thermodyn.* **17**, 277 (1985).
- 1985COL/JIM M. Colomina, P. Jimenez, M. V. Roux, and C. Turrión, *J. Chem. Thermodyn.* **17**, 1091 (1985).
- 1985DEA J. A. Dean, *Lange's Handbook of Chemistry*, 13th ed. (McGraw-Hill, New York, 1985).
- 1985DEM/CHA G. De Maury, A. Chauvet, and J. Masse, *Thermochim. Acta* **87**, 189 (1985).
- 1985DEV/GUE G. G. Devyatikh, A. V. Gusev, A. M. Gibin, N. V. Zhermenkov, L. M. Zakharov, M. Yu. Antipin, and Yu. M. T. Struchov, *Russ. J. Inorg. Chem.* **30**, 780 (1985).
- 1985FER/PIA D. Ferro and V. Piacente, *Thermochim. Acta* **90**, 387 (1985).
- 1985FIG/SZW P. Figuiere, H. Szwarc, M. Ogumi, and H. Suga, *J. Chem. Thermodyn.* **17**, 949 (1985).
- 1985GLU/ARK O. T. Glukhova, N. M. Arkhangelova, A. B. Teplitsky, L. F. Sukhodub, I. K. Yanson, and M. Kaminski, *Thermochim. Acta* **95**, 133 (1985).
- 1985HAN Y. P. Handa, *Can. J. Chem.* **63**, 68 (1985).
- 1985HIL/BEN K. Hilpert, L. Bencivenni, and B. Saha, *Ber. Bunsenges. Phys. Chem.* **89**, 1292 (1985).
- 1985IGU/GER I. K. Igumenov, T. Yu. Gerasimenko, and V. G. Isakova, *Izv. Sib. Otd. Akad. Nauk SSSR Ser. Khim. Nauk*, 42 (1985).
- 1985KAI/HAD E. Kaiserberger, W. Hadrich, and W.-D. Emmerich, *Thermochim. Acta* **95**, 331 (1985).
- 1985KAM/ZIE M. Kaminski and W. Zielenkiewicz, *Calorim. Anal. Therm.* **16**, 281 (1985); *Chem. Abstr.* **104**, 96844t (1986).
- 1985KLY/DAN V. A. Klyuchnikov, T. F. Danilova, M. G. Voronkov, and V. M. D'yakov, *Dokl. Phys. Chem.* **281**, 373 (1985).
- 1985KON/STR S. N. Kondrat'ev and B. V. Strizhov, *Zh. Fiz. Khim.* **59**, 2933 (1985).
- 1985KUS K. Kusano, *Thermochim. Acta* **88**, 109 (1985).
- 1985LEB/BYK B. V. Lebedev, T. A. Bykova, E. G. Kiparisova, Y. A. Chernomordik, A. S. Kurapov, and V. A. Sergeev, *Bull. Acad. Sci. USSR Div. Chem. Sci.* **34**, 274 (1985).
- 1985LIN/RAT S. Lindenbaum, E. S. Rattie, G. E. Zuber, M. E. Miller, and L. J. Ravin, *Int. J. Pharm.* **26**, 123 (1985).
- 1985MAJ/SVO V. Majer and V. Svoboda, *Enthalpies of Vaporization of Organic Compounds: A Critical Review and Data Compilation* (Blackwell Scientific, Oxford, 1985).
- 1985MAJ/SVO2 V. Majer, V. Svoboda, and M. Lencka, *J. Chem. Thermodyn.* **17**, 365 (1985).
- 1985MAJ/SVO3 V. Majer, V. Svoboda, V. Uchytlova, and M. Finke, *Fluid Phase Equilib.* **20**, 111 (1985).
- 1985MAR/MANK N. Marsh and M. Mansson, *J. Chem. Thermodyn.* **17**, 995 (1985).
- 1985MAR/WU A. Martin, P. L. Wu, and T. Velasquez, *J. Pharm. Sci.* **74**, 277 (1985).
- 1985MAT/KUW N. Matsubara and T. Kuwamoto, *Inorg. Chem.* **24**, 2697 (1985).
- 1985MAT/KUW2 N. Matsubara and T. Kuwamoto, *Thermochim. Acta* **83**, 193 (1985).
- 1985MOU A. H. N. Mousa, *J. Fluorine Chem.* **30**, 29 (1985).
- 1985MUR/SAK S. Murata, M. Sakiyama, and S. Seki, *Thermochim. Acta* **88**, 121 (1985).
- 1985NAT/VIS G. Natarajan and D. S. Viswanath, *J. Chem. Eng. Data* **30**, 137 (1985).
- 1985OHM/LIP A. Ohm and B. C. Lippold, *Int. J. Pharm. Tech. Prod. Mfg.* **6**, 1 (1985).
- 1985PAS/VAR L. L. Pashchenko and R. M. Varushchenko, *Zh. Obshch. Khim.* **55**, 721 (1985).
- 1985PIA/SCA V. Piacente, P. Scardala, D. Ferro, and R. Gigli, *J. Chem. Eng. Data* **30**, 372 (1985).
- 1985PRI/PUC V. P. Privalko, G. A. Puchkovskaya, E. N. Shermatov, and A. A. Yakubov, *Mol. Cryst. Liq. Cryst.* **126**, 289 (1985).
- 1985RAE/SOL O. A. Raewskii, B. P. Solov'ev, and L. W. Govorkova, *Zh. Obshch. Khim.* **55**, 1381 (1985).
- 1985RAV/RAO A. Raviprasad and K. V. Rao, *J. Chem. Thermodyn.* **17**, 117 (1985).
- 1985RED/RAO K. D. Reddy, M. V. P. Rao, and M. Ramakrishna, *J. Chem. Eng. Data* **30**, 394 (1985).
- 1985ROR B. F. Rordorf, *Thermochim. Acta* **85**, 435 (1985).
- 1985SAI/ATA K. Saito, T. Atake, and H. Chihara, *J. Chem. Thermodyn.* **17**, 539 (1985).
- 1985SCH/BRU A. G. R. Scholz and E. Brunner, *J. Chem. Eng. Data* **30**, 72 (1985).
- 1985SCH/HIL J. M. Schroy, F. D. Hileman, and S. C. Cheng, *Chemosphere* **14**, 877 (1985).
- 1985SER/ZAG G. B. Sergeev, V. V. Zagorskii, A. M. Kosolapov, and S. E. Kondakov, *Zh. Neorg. Khim.* **30**, 3212 (1985); *Chem. Abstr.* **104**, 121894u (1986).
- 1985SKI/PIL H. A. Skinner, G. Pilcher, and G. Al-Takhin, personal comment, as cited in M. Faour and T. S. Akasheh, *J. Chem.*

- Soc., Perkin Trans. 2 811 (1985).
- 1985SOK/BAE A. E. Sokolovski and A. K. Baev, *Vesti Akad. G. Emii Navuk BSSR Ser. Khim. Navuk*, 1112 (1985).
- 1985STE/GAN W. V. Steele, B. E. Gannon, N. K. Smith, A. Greenburg, J. S. Chickos, and J. F. Liebman, *J. Chem. Thermodyn.* **17**, 505 (1985).
- 1985TRU/KRA V. I. Trusov, I. M. Krasikova, A. A. Grigor'ev, A. I. Altsybeeva, A. V. Suvorov, A. A. Khonikevich, and P. A. Vinogradov, *Zashch. Met.* **21**, 299 (1985); *Chem. Abstr.* **102**, 173623h (1985).
- 1985VAR/BRI D. Varech, M. J. Brienne, and J. Jacques, *Tetrahedron Lett.* **26**, 61 (1985).
- 1985VEN/PRA V. Venugopal, R. Prasad, and D. D. Sood, *J. Nucl. Mater.* **130**, 115 (1985).
- 1985WIB/WAS K. B. Wiberg, D. J. Wasserman, E. J. Martin, and M. A. Murcko, *J. Am. Chem. Soc.* **107**, 6019 (1985).
- 1985WIE/SIP S. A. Wiczorek and J. T. Sipowska, *J. Chem. Thermodyn.* **17**, 255 (1985).
- 1985WOR/YER C. J. Wormald and T. K. Yerlett, *J. Chem. Thermodyn.* **17**, 1171 (1985).
- 1985YAN/GU H. K. Yan, J. K. Gu, and R. H. Fu, *J. Environ. Eng.* **1**, 543 (1985).
- 1985ZAR A. P. Zaraiskii, *Zh. Fiz. Khim.* **59**, 2087 (1985).
- 1985ZVE/MOT V. V. Zvezdina, O. R. Motorina, G. Yu. Guara, V. P. Boshin, O. A. Golubchikov, and B. D. Berezin, *Russ. J. Phys. Chem.* **57**, 1422 (1983).
- 1986ACH/HAS P. J. Achorn, W. G. Haseltine, and J. K. Miller, *J. Chem. Eng. Data* **31**, 385 (1986).
- 1986ALL/JOS N. Allemand, J. Jose, and J. C. Merlin, *Thermochim. Acta* **105**, 79 (1986).
- 1986BAL/GNA V. Baliah and K. Gnanasekaran, *Indian J. Chem., Sect A: Inorg., Phys., Theor. Anal.* **25A**, 673 (1986).
- 1986BAR/FER G. Barone, D. Ferro, P. Fiorani, C. A. Mattia, and R. Puliti, *J. Calorim. Anal. Therm. Thermodyn. Chim.* **17**, 471 (1986).
- 1986BER/COL M. Bertaut, M. Collet, and R. Nowak, *Solid State Commun.* **58**, 613 (1986).
- 1986CAO/WES M. Cao, J. Wesson, K. Loufakis, B. Wunderlich, and M. Moller, *Mol. Cryst. Liq. Cryst.* **140**, 231 (1986).
- 1986CHI/ANN J. S. Chickos, R. S. Annunziata, L. H. Ladon, A. S. Hyman, and J. F. Liebman, *J. Org. Chem.* **51**, 4311 (1986).
- 1986CHU/DEM A. Chuvet, G. De Maury, A. Terol, and J. Masse, *Thermochim. Acta* **97**, 143 (1986).
- 1986CIH/VOJ M. Cihova, J. Vojtko, and J. Ilavsky, *Petrochimica* **26**, 69 (1986).
- 1986COL/JIM M. Colomina, P. Jimenez, M. V. Roux, and C. Turrión, *An. Quim.* **82**, 126 (1986).
- 1986DEM/MAS G. De Maury and J. Masse, *J. Therm. Anal.* **31**, 1263 (1986).
- 1986DEV/GUS G. G. Devyatykh, A. V. Gusev, A. M. Gibin, N. V. Zhernenkov, and A. V. Kabanov, *Russ. J. Inorg. Chem.* **31**, 1281 (1986).
- 1986DMI/KAC Yu. G. Dmitriev, N. S. Kachurina, C. H. Wang, and V. V. Kouchukei, *Vestn. L'vov. Politekh. Inst.* **201**, 29 (1986).
- 1986FLA O. L. Flanagan, *J. Chem. Eng. Data* **31**, 266 (1986).
- 1986FLA/BEC M. A. Flammtter Meer, H.-D. Beckhaus, and C. Rüchardt, *Thermochim. Acta* **107**, 331 (1986).
- 1986GAR/JAN G. Gärtner, P. Janiel, H. Rau, H. A. M. van Hal, and H. J. P. Nabben, *Ber. Bunsenges. Phys. Chem.* **90**, 459 (1986).
- 1986GON/SZW A. Gonthier-Vassal and H. Szwarc, *Chem. Phys. Lett.* **129**, 5 (1986).
- 1986GRI/LAZ Ya. Kh. Grinberg, V. B. Lazarev, A. Yu. Zavernyaev, V. A. Shreider, and S. D. Chepik, *Zh. Fiz. Khim.* **60**, 1386 (1986); *Chem. Abstr.* **105**, 86041k (1986).
- 1986HAN/ECK P. C. Hansen and C. A. Eckert, *J. Chem. Eng. Data* **31**, 1 (1986).
- 1986HAR/GIL P. D. Harvey, D. F. R. Gilson, and I. S. Butler, *J. Phys. Chem.* **90**, 136 (1986).
- 1986HES/LIC W. Hessler and W. Lichtenstein, *Wissenschaftliche Zeitschrift der Wilhelm-Piech Universität Rostich* **35**, 27 (1986).
- 1986JAM/PIL E. H. Jamea and G. Pilcher, *Thermochim. Acta* **97**, 77 (1986).
- 1986JIM/ROU P. Jimenez, M. V. Roux, C. Turrión, and F. Gomis, *J. Calorim. Anal. Therm. Thermodyn. Chim.* **17**, 469 (1986).
- 1986KAL/JAC B. Kalinowska and A. W. Jackowski, *J. Chem. Thermodyn.* **18**, 715 (1986).
- 1986KIR/ACR J. J. Kirchner, W. E. Acree, Jr., G. Pilcher, and S. Li, *J. Chem. Thermodyn.* **18**, 793 (1986).
- 1986KOZ/DAL A. A. Kozyro, S. V. Dalidovich, and A. P. Krausulin, *Zh. Prikl. Khim. (S.-Peterburg)* **59**, 1456 (1986).
- 1986KRA/KOZ A. P. Krasulin, A. A. Kozyro, and S. V. Dalidovich, *Zh. Fiz. Khim.* **60**, 2580 (1986).
- 1986KRA/KOZ2 A. P. Krasulin and A. A. Kozyro, Proceedings of the 11th All-Union Conference on Calorimetry and Chemical Thermodynamics, Novosibirsk, USSR, 17 June 1986 (unpublished), p. 129; see also [1990PIA/FER].
- 1986KRE/PRA D. H. Krevor and J. M. Prausnitz, *J. Chem. Eng. Data* **31**, 349 (1986).
- 1986KUN/YUK H. Kunisada and Y. Yuki, *Kobunshi Rombunshu* **43**, 897 (1986); *Chem. Abstr.* **106**, 85109.
- 1986LAT/HOE G. Latterman and H. Hoecker, *Mol. Cryst. Liq. Cryst.* **133**, 245 (1986).
- 1986MAR/LOE Y. Marcus and A. Loewenschuss, *J. Chem. Soc., Faraday Trans. 1* **82**, 993 (1986).
- 1986MEY/MEY E. F. Meyer and J. J. Meyer, *J. Chem. Eng. Data* **31**, 273 (1986).
- 1986MIC/JOS M.-A. Michou-Saucet, J. Jose, and C. Michou-Saucet, *Thermochim. Acta* **102**, 271 (1986).
- 1986MIY/MAT S. Miyagishi, S. Matumura, T. Asakawa, and M. Nishida, *Bull. Chem. Soc. Jpn.* **59**, 557 (1986).
- 1986NIL/WAD S.-O. Nilsson and I. Wadso, *J. Chem. Thermodyn.* **18**, 673 (1986).
- 1986NIS/AND K. Nishida, Y. Ando, S. Sunagawa, A. Ogihara, I. Tanaka, and A. Koukitsu, *J. Soc. Dyers Colour.* **102**, 18 (1986).
- 1986NUN/BAR L. Nunez, L. Barral, L. S. Gavelanes, and G. Pilcher, *J. Chem. Thermodyn.* **18**, 575 (1986).
- 1986PAU/KRU H.-I. Paul, J. Krug, and H. Knapp, *Thermochim. Acta* **108**, 9 (1986).
- 1986PED/NAY J. P. Pedley, R. D. Naylor, and S. P. Kirby, *Thermochemical Data of Organic Compounds*, 2nd ed. (Chapman and Hall, New York, 1986).
- 1986PIM/DOM S. M. Pimenova, I. N. Domnin, A. M. Lakshin, and V. V. Takhistov, Abstract of Papers, XI All-Union Conference on Calorimetry and Thermodynamics Novosibirsk, 1986 (unpublished), Vol. 2, p. 41.
- 1986RIB/RIB M. D. M. C. Ribeiro Da Silva, M. A. V. Ribeiro Da Silva, and G. Pilcher, *J. Chem. Thermodyn.* **18**, 295 (1986).
- 1986ROR B. F. Rordorf, *Chemosphere* **15**, 1325 (1986).
- 1986ROR/SAR B. F. Rordorf, L. P. Sarna, and G. R. B. Webster, *Chemosphere* **15**, 2073 (1986).
- 1986ROR2 B. F. Rordorf, Proceedings of the fifth International Symposium on Chlorinated Dioxins and Related Compounds, Bayreuth, FRG, 16-19 September 1986 (unpublished).
- 1986RUS/RAB T. P. Russell, J. F. Rabolt, R. J. Twieg, R. L. Siemens, and B. L. Farmer, *Macromolecules* **19**, 1135 (1986).
- 1986SAT/INO N. Sato, H. Inomata, K. Arai, and S. Saito, *J. Chem. Eng. Jpn.* **19**, 145 (1986).
- 1986SHA/BRI A. C. Shah and N. J. Britten, *J. Pharm. Pharmacol.* **39**, 736 (1986).
- 1986STA H. W. Starkweather, Jr., *Macromolecules* **19**, 1131 (1986).
- 1986TAK/ISH S. Takagi, M. Ishikawa, M. Kumada, T. Kimura, and R. Fujishiro, *Thermochim. Acta* **109**, 55 (1986).
- 1986TSV/NAZ V. S. Tsvetkov, A. G. Nazmutdinov, K. S. Sharonov, and A. M. Rozhnov, *Termodin. Org. Soedin.*, 71 (1986).
- 1986VAR R. M. Varushchenko, *Zh. Fiz. Khim.* **60**, 291 (1986).
- 1986VOR/BAL V. M. Vorotyntsev, V. V. Balabanov, and D. A. Shamrakov, *Zh. Fiz. Khim.* **60**, 234 (1986); *Chem. Abstr.* **104**, 96047k (1986).
- 1986WEL/GRA D. Wells, B. T. Grayson, and E. Langner, *Pestic. Sci.* **17**, 473 (1986).
- 1986WEN/SCH U. Wenzel and G. M. Schneider, *Thermochim. Acta* **109**, 111 (1986).
- 1986WHI C. M. White, *J. Chem. Eng. Data* **31**, 198 (1986).

- 1986WIS/LEN B. Wisniewska, M. Lencka, and M. Rogalski, *J. Chem. Thermodyn.* **18**, 703 (1986).
- 1986YER/WOR Y. K. Yerlett and C. J. Wormald, *J. Chem. Thermodyn.* **18**, 719 (1986).
- 1986YER/WOR2T. K. Yerlett and C. J. Wormald, *J. Chem. Thermodyn.* **18**, 371 (1986).
- 1987AIR/DES C. Airoldi and A. G. DeSouza, *J. Chem. Soc. Dalton Trans.* 2955 (1987).
- 1987ALL/FIN P. H. Allot, A. Finch, G. Pilcher, L. Numez, and L. Barral, *J. Chem. Thermodyn.* **19**, 771 (1987).
- 1987ALN/ALS H. Al-Najjar and D. Al-Sammerrai, *J. Chem. Technol. Bio-technol.* **37**, 145 (1987).
- 1987AMB/GHI D. Ambrose and N. B. Ghiassee, *J. Chem. Thermodyn.* **19**, 911 (1987).
- 1987AMB/GHI2 D. Ambrose and N. B. Ghiassee, *J. Chem. Thermodyn.* **19**, 903 (1987).
- 1987AMB/GHI3 D. Ambrose and N. B. Ghiassee, *J. Chem. Thermodyn.* **19**, 505 (1987).
- 1987AN/HU O. An, R. Hu, H. Wang, M. Wu, and Y. Zou, *Acta Phys. Chim. Sinica* **3**, 668 (1987).
- 1987AN/ZHU X. An, I. Zhu, and R. Hu, *Thermochim. Acta* **121**, 473 (1987).
- 1987ATI/SAI Z. Atik, Y. Saito, and K. Kusano, *J. Chem. Thermodyn.* **19**, 99 (1987).
- 1987AZA E. C. Azandegbe, *Analisis* **15**, 370 (1987).
- 1987CHI J. S. Chickos, in *Molecular Structure and Energetics*, edited by J. F. Liebman and A. Greenberg (VCH, New York, 1987), Vol. 2, Chap. 3.
- 1987COL/JIM M. Colomina, P. Jimenez, R. Perez-Ossorlo, M. V. Roux, and C. Turrion, *J. Chem. Thermodyn.* **19**, 155 (1987).
- 1987COL/JIM2 M. Colomina, P. Jimenez, M. V. Roux, and C. Turrion, *J. Chem. Thermodyn.* **19**, 1139 (1987).
- 1987CUR/ASR S. A. Curran and J. Asrar, *Mol. Cryst. Liq. Cryst.* **148**, 255 (1987).
- 1987DAU/JAL T. E. Daubert, J. W. Jalowka, and V. Goren, *AIChE Symp. Ser.* **83**, 128 (1987).
- 1987DIA/DIA A. R. Dias, P. B. Dias, H. P. Diogo, A. M. Galvao, M. E. Minas da Piedade, and J. A. Martinho Simoes, *Organometallics* **6**, 1427 (1987).
- 1987DRO/MOL H. Drotloff and M. Moller, *Thermochim. Acta* **112**, 57 (1987).
- 1987ECO/BER C. Ecolivet, A. Bertaut, A. Miezeyewski, and A. Collet, in *Dynamics of Molecular Crystals*, edited by J. Lascombe (Elsevier, Amsterdam, 1987), p. 187.
- 1987FER/DEL D. Ferro and G. Della Gatta, *Thermochim. Acta* **122**, 189 (1987).
- 1987FER/DEL2 D. Ferro, G. Della Gatta, and V. Piacente, *J. Chem. Thermodyn.* **19**, 915 (1987).
- 1987FER/PIL M. L. C. C. H. Ferrao and G. Pilcher, *J. Chem. Thermodyn.* **19**, 543 (1987).
- 1987FER/STR D. Ferro and S. Stranges, *Thermochim. Acta* **119**, 373 (1987).
- 1987FIO/FER P. Fiorani and D. Ferro, *Thermochim. Acta* **112**, 387 (1987).
- 1987FUC/CHA R. Fuchs, E. J. Chambers, and W. K. Stepenson, *Can. J. Chem.* **65**, 2624 (1987).
- 1987GAR/TRE F. Garcia-Sanchez and A. Trejo, *J. Chem. Thermodyn.* **19**, 359 (1987).
- 1987GIB/GRI A. M. Gibin, N. D. Grishnova, A. V. Gusev, A. N. Moiseev, I. I. Remeshkova, and Y. M. Salganskii, *Vysokochist. Veshchestva*, 28 (1987); *Chem. Abstr.* **109**, 136102 (1988).
- 1987GRE/SID N. R. Grebenshchikov, G. V. Sidorenko, and D. N. Suglobov, *Radiokhimiya* **29**, 296 (1987).
- 1987GRI/LAZ Ya. Kh. Grinberg and V. B. Lazarev, *Zh. Neorg. Khim.* **32**, 3110 (1987).
- 1987GRI/LAZ2 Ya. Kh. Grinberg, V. B. Lazarev, V. V. Petukhov, G. Ya. Novikova, G. A. Sharpataya, and Z. P. Ozerova, *Dokl. Akad. Nauk SSSR* **297**, 633 (1987).
- 1987HAU/WU T. Hauschild, H. S. Wu, and S. I. Sandler, *J. Chem. Eng. Data* **32**, 226 (1987).
- 1987JIM/ROU P. Jimenez, M. V. Roux, C. Turrion, and F. Gomis, *J. Chem. Thermodyn.* **19**, 985 (1987).
- 1987KAF/DOR V. V. Kafarov, I. N. Dorokhov, V. N. Vetokhin, and L. P. Volkov, *Dokl. Phys. Chem.* **298**, 77 (1987).
- 1987KAM/ZIE M. Kaminski and W. Zielenkiewicz, *Bull. Pol. Acad. Sci., Chem.* **35**, 583 (1987).
- 1987KAN/OI A. Kanungo, T. Oi, A. Popowicz, and T. Ishida, *J. Phys. Chem.* **91**, 4198 (1987).
- 1987KLA/MOH S. M. Klara, R. S. Mohamed, D. M. Dempsey, and G. D. Holder, *J. Chem. Eng. Data* **32**, 143 (1987).
- 1987KNE/ZON P. Kneisi and J. W. Zondio, *J. Chem. Eng. Data* **21**, 11 (1987).
- 1987LAZ/GRE V. B. Lazarev, J. H. Greenberg, Z. P. Ozerova, and G. A. Sharpataya, *J. Therm. Anal.* **33**, 797 (1987).
- 1987MEI/DOG M. Meier, B. Dogan, H. D. Beckhaus, and C. Rüchardt, *New J. Chem.* **11**, 1 (1987).
- 1987MIH/BAS P. Mihailovic, P. Bassoul, and J. Simon, *Chem. Phys. Lett.* **141**, 462 (1987).
- 1987MIL/FEN P. L. Mills and R. L. Fenton, *J. Chem. Eng. Data* **32**, 266 (1987).
- 1987MIL/FEN2 P. L. Mills, R. L. Fenton, and F. G. Schaefer, *J. Chem. Eng. Data* **32**, 251 (1987).
- 1987MUR/HIL J. P. Murray and J. O. Hill, *Thermochim. Acta* **109**, 383 (1987).
- 1987MUR/HIL2 J. P. Murray and J. O. Hill, *Thermochim. Acta* **109**, 391 (1987).
- 1987OYU/BRI Y. Oyumi and T. B. Brill, *Thermochim. Acta* **116**, 125 (1987).
- 1987RIB/FER M. A. V. Ribeiro da Silva and M. L. C. C. H. Ferrao, *J. Chem. Thermodyn.* **19**, 645 (1987).
- 1987SAB/ANT R. Sabbah and I. Antipine, *Bull. Soc. Chim. Fr.*, 392 (1987).
- 1987SHI/OHK T. Shimizu, S. Ohkubo, M. Kimura, I. Tabata, and T. Hori, *J. Soc. Dyers Colour.* **103**, 132 (1987).
- 1987SMI/TER G. G. Smirnova, G. F. Tereshchenko, and M. A. Blagushina, *Zh. Prikl. Khim. (S.-Peterburg)* **62**, 182 (1987).
- 1987STE/MAL R. M. Stephenson and S. Malanowski, *Handbook of the Thermodynamics of Organic Compounds* (Elsevier: New York, 1987). The data in this listing were obtained directly from this compendium which does not contain literature references. The sublimation enthalpies were calculated from the vapor pressures obtained from the Antoine constants reported in this compendium. In cases where the Antoine constant $C=0$, the Antoine Equation ($\log_{10}P=A-B/(C+T)$) reduces to the integrated form of the Clausius Clapeyron equation directly. This was the case for most sublimation enthalpies. In those cases where this condition was not met, vapor pressures were calculated over a 30 K temperature range closest to $T=298$ K and the temperature dependence was reformulated in terms of the integrated form of the Clausius Clapeyron equation.
- 1987TRE/LU T. Treszczanowicz and B. C.-Y. Lu, *J. Chem. Thermodyn.* **19**, 391 (1987).
- 1987VAN/KAC Yu. Ya. Van-Chin-Syan and N. S. Kachurina, *Russ. J. Phys. Chem.* **61**, 622 (1987).
- 1987VAR/LOS R. M. Varushchenko, O. L. Loseva, and A. I. Druzhinina, *Zh. Fiz. Khim.* **61**, 31 (1987).
- 1987VAR/LOS2 R. M. Varushchenko, O. L. Loseva, and A. I. Druzhinina, *Zh. Fiz. Khim.* **61**, 638 (1987).
- 1987YAN/GU H. Yan, J. Gu, X. An, and R.-H. Hu, *Huaxue Xuebao* **45**, 1184 (1987).
- 1988AFZ/BUT M. Afzal, P. K. Butt, and M. J. Iqbal, *J. Sci. Ind. Res.* **31**, 404 (1988).
- 1988AMB/GHI D. Ambrose and N. B. Ghiassee, *J. Chem. Thermodyn.* **20**, 1231 (1988).
- 1988AMB/GHI2 D. Ambrose and N. B. Ghiassee, *J. Chem. Thermodyn.* **20**, 765 (1988).
- 1988AMB/GHI3 D. Ambrose, N. B. Ghiassee, and R. Tuckerman, *J. Chem. Thermodyn.* **20**, 767 (1988).
- 1988ASK/DAU C. F. Askonas and T. E. Daubert, *J. Chem. Eng. Data* **33**, 225 (1988).
- 1988BAG/GUR A. K. Baglay, L. L. Gurarly, and G. G. Kuleshov, *J. Chem. Eng. Data* **33**, 512 (1988).
- 1988BAS/NIL M. Bastos, S. O. Nilsson, M. D. M. C. Ribeiro da Silva, M.

- A. V. Ribeiro da Silva, and J. Wadso, *J. Chem. Thermodyn.* **20**, 1353 (1988).
- 1988BAU/PER G. L. Baughman and T. A. Perenich, *Envir. Toxicol. Chem.* **7**, 183 (1988).
- 1988BRA/FAK D. C. Bradley, M. M. Faktor, D. M. Frigo, and K. V. Young, *Chemtronics* **3**, 50 (1988); *Chem. Abstr.* **109**, 43735e (1988).
- 1988CAR/DYS A. S. Carson, J. Dyson, P. G. Laye, and J. A. Spencer, *J. Chem. Thermodyn.* **20**, 1423 (1988).
- 1988CAR/JAM A. S. Carson, E. H. Jamea, P. G. Laye, and J. A. Spencer, *J. Chem. Thermodyn.* **20**, 923 (1988).
- 1988CAR/JAM2 A. S. Carson, E. H. Jamea, P. G. Laye, and J. A. Spencer, *J. Chem. Thermodyn.* **20**, 1223 (1988).
- 1988COL/JIM M. Colomina, P. Jimenez, R. Perez-Ossorio, M. V. Roux, and C. Turrion, *J. Chem. Thermodyn.* **20**, 575 (1988).
- 1988DAV/EWI D. R. M. Davies, B. Ewing, J. A. Hugill, and M. L. McGlashan, *Can. J. Chem.* **66**, 760 (1988).
- 1988DEF/CHA F. Defoort, C. Chatillon, and C. Bernard, *J. Chem. Thermodyn.* **20**, 1443 (1988).
- 1988DON/LIN J.-Q. Dong, R.-S. Lin, and W.-H. Yen, *Can. J. Chem.* **66**, 783 (1988).
- 1988FED/VOI N. E. Fedotova, A. A. Voityuk, A. A. Blitzyuk, and I. K. Igumenov, *Koord. Khim.* **14**, 1493 (1988).
- 1988FER/DEL D. Ferro, G. Della Gatta, and G. Barone, *J. Therm. Anal.* **34**, 835 (1988).
- 1988GAL/GON F. Galan-Estella, J. Gonzalez-Julian, and P. Aquado-Rodriguez, *Photochemistry* **27**, 3069 (1988); *Chem. Abstr.* **110**, 102153h (1989).
- 1988GOL/SIT O. F. Golovanova, G. V. Sitonina, V. I. Pepekin, and F. I. Korsunskii, *Izv. Akad. Nauk SSSR Ser. Khim.*, 1012 (1988); *Chem. Abstr.* **110**, 38466j (1989).
- 1988GRE/SID N. R. Grebenshchikov, G. V. Sidorenko, D. N. Suglobov, L. L. Shcherbakova, and V. M. Adamov, *Inorg. Chim. Acta* **145**, 315 (1988).
- 1988GRI/CHE E. E. Grinberg, N. G. Chernaya, and A. A. Efremov, *Vysokochist. Veshchestva*, 180 (1988).
- 1988HOP/PUG J. Höpken, C. Pugh, W. Richtering, and M. Möller, *Makromol. Chem.* **189**, 911 (1988).
- 1988HOS/ARC I. A. Hossenlopp and D. G. Archer, *J. Chem. Thermodyn.* **20**, 1061 (1988).
- 1988HUA/TAN W. Huang, F. Tan, and H. Luo, *Zhongguo Yaoke Daxue Xuebao* **19**, 90 (1988).
- 1988IMA/MUR A. Imamura, S. Murata, and M. Sakiyama, *J. Chem. Thermodyn.* **20**, 389 (1988).
- 1988ISM/GAB T. S. Ismailov, N. R. Gabzalilova, and Kh. M. Makhkamov, *Uzb. Khim. Zh.* **4**, 48 (1988).
- 1988KAT H. Katayama, *Bull. Chem. Soc. Jpn.* **61**, 3326 (1988).
- 1988KAT2 H. Katayama, *J. Chem. Eng. Data* **33**, 75 (1988).
- 1988KHU V. L. Khudyakov, *Russ. J. Phys. Chem.* **62**, 1743 (1988).
- 1988KIR/DOM D. R. Kirklin and E. S. Domalski, *J. Chem. Thermodyn.* **20**, 743 (1988).
- 1988KNA/SAB P. Knauth and R. Sabbah, *Bull. Soc. Chim. Fr.*, 834 (1988).
- 1988LAZ/GRI V. B. Lazarev, Y. K. Grinberg, Z. P. Ozerova, and G. A. Sharpataya, *J. Therm. Anal.* **33**, 797 (1988).
- 1988LEB/BYK B. V. Lebedev, T. A. Bykova, E. G. Kiparisova, M. Frenkel, A. M. Fainleib, and V. A. Pankratov, *Bull. Acad. Sci. USSR Div. Chem. Sci.* **37**, 1082 (1988).
- 1988LEB/KAL B. V. Lebedev, T. G. Kalagina, and N. N. Smirnova, *J. Chem. Thermodyn.* **20**, 1383 (1988).
- 1988LET/SEW T. M. Letcher, J. Sewry, and S. W. Orchard, *J. Chem. Thermodyn.* **20**, 1115 (1988).
- 1988LIC/RIT H. H. Licht, and H. Ritter, *Propellants, Explos., Pyrotech.* **13**, 25 (1988).
- 1988LUS/RUB V. N. Lushnikov, Y. I. Rubstov, L. T. Dremenko, and A. M. Korolev, *Zh. Fiz. Khim.* **62**, 1209 (1988).
- 1988MES/FIN J. F. Messerly, H. L. Finke, W. D. Good, and B. Gammon, *J. Chem. Thermodyn.* **20**, 485 (1988).
- 1988NAK/KIT H. Nakazuma, T. Kitao, and K. Ohnishi, *J. Heterocycl. Chem.* **25**, 1422 (1988).
- 1988NIS/SHE V. P. Nistratov, M. S. Sheiman, I. B. Rabinovich, V. G. Vasil'ev, E. N. Karataev, and I. A. Feshchenko, *Zh. Fiz. Khim.* **62**, 2219 (1988).
- 1988NUN/BAR L. Nunez, L. Barral, and G. Pilcher, *J. Chem. Thermodyn.* **20**, 1211 (1988).
- 1988PAR/KAW R. M. Paroli, N. T. Kawai, I. S. Butler, and D. F. R. Gilson, *Can. J. Chem.* **66**, 1973 (1988).
- 1988PAU/KRU H. I. Paul, J. Krug, and H. Knapp, *J. Chem. Eng. Data* **33**, 435 (1988).
- 1988PES/SHV A. D. Peschenko, O. V. Shvaro, and V. I. Zubkov, *Termodin. Org. Soedin. (Gorky)*, 39 (1988).
- 1988RIB/FER M. A. V. Ribeiro da Silva and M. L. C. C. H. Ferrao, *Bull. Chem. Soc. Jpn.* **61**, 1755 (1988).
- 1988RIB/FER2 M. A. V. Ribeiro da Silva and M. L. C. C. H. Ferrao, *Can. J. Chem.* **66**, 651 (1988).
- 1988RIB/FER3 M. A. V. Ribeiro da Silva, M. L. C. C. H. Ferrao, and C. C. H. Luisa, *J. Chem. Thermodyn.* **20**, 359 (1988).
- 1988RIB/FER4 M. A. V. Ribeiro da Silva and M. L. C. C. H. Ferrao, *Pure Appl. Chem.* **60**, 1225 (1988).
- 1988RIB/PIL M. A. V. Ribeiro da Silva, G. Pilcher, and R. J. Irving, *J. Chem. Thermodyn.* **20**, 95 (1988).
- 1988RIB/REI M. A. V. Ribeiro da Silva, A. M. Reis, and G. Pilcher, *Thermochim. Acta* **124**, 319 (1988).
- 1988RIB/RIB M. A. V. Ribeiro da Silva, M. D. M. C. Ribeiro da Silva, and G. Pilcher, *J. Chem. Thermodyn.* **20**, 969 (1988).
- 1988ROY/FLY S. D. Roy and G. L. Flynn, *Pharm. Res.* **5**, 580 (1988).
- 1988SAB/ELW R. Sabbah and L. El Watik, *Bull. Soc. Chim. Fr.*, 626 (1988).
- 1988SAB/ELW2 R. Sabbah, L. El Watik, and C. C. R. Minadakis, *Acad. Sci., Paris, C. R.* **307**, 239 (1988).
- 1988SAI/ATA K. Saito, T. Atake, and H. Chihara, *Bull. Chem. Soc. Jpn.* **61**, 2327 (1988).
- 1988SAS/JOS K. Sasse, J. Jose, and J.-C. Merlin, *Fluid Phase Equilib.* **42**, 287 (1988).
- 1988SHA/DIE R. H. Shay, B. N. Diel, D. M. Schubert, and A. D. Norman, *Inorg. Chem.* **27**, 2378 (1988).
- 1988SHA/PIL L. Shaofeng and G. Pilcher, *J. Chem. Thermodyn.* **20**, 463 (1988).
- 1988SOP/KEA P. A. Sopade, M. W. Kearsley, and G. A. Le Grys, *Int. Sugar J.* **90**, 38 (1988).
- 1988STE/ARC W. V. Steele, D. G. Archer, R. D. Chirico, W. B. Collier, I. A. Hossenlopp, A. Nguyen, K. N. Smith, and B. E. Gammon, *J. Chem. Thermodyn.* **20**, 1233 (1988).
- 1988TEL/LAR V. I. Tel'noi, V. N. Larina, E. N. Karataev, and E. N. Deryagina, *Zh. Fiz. Khim.* **62**, 3108 (1988).
- 1988TOR/BAR L. A. Torres-Gomez, G. Barreiro-Rodriguez, and A. Galarza-Mondragon, *Thermochim. Acta* **124**, 229 (1988).
- 1988TOR/BAR2 L. A. Torres-Gomez, G. Barreiro-Rodriguez, and F. Mendez-Ruiz, *Thermochim. Acta* **124**, 179 (1988).
- 1988VAR/LOS R. M. Varushchenko, O. L. Loseva, A. I. Druzhinina, and E. F. Zorina, *Zh. Fiz. Khim.* **62**, 1776 (1988).
- 1988VOR/BAR M. G. Voronkov, V. P. Baryshok, V. A. Klyuchnikov, T. F. Danilova, V. I. Pepekin, A. NI. Korchagina, and Y. I. Khudobin, *J. Organomet. Chem.* **345**, 27 (1988).
- 1988WHI/PER M. A. White and A. Perrott, *Can. J. Chem.* **66**, 729 (1988).
- 1989ABB/JIM J.-L. Abboud, P. Jimenez, M. V. Roux, C. C. Turrion, and C. Lopez-Mardomingo, *J. Chem. Thermodyn.* **21**, 859 (1989).
- 1989ACR/KIR W. E. Acree, Jr., J. J. Kirchner, S. A. Tucker, G. Pilcher, and D. M. C. Riberio da Silva, *J. Chem. Thermodyn.* **21**, 443 (1989).
- 1989AIZ/FED M. I. Aizenberg, E. V. Fedoseev, S. S. Travnikov, A. V. Davydov, and B. F. Myasoedov, *J. Radioanal. Nucl. Chem.* **136**, 405 (1989).
- 1989AN/HU X. An and H. Hu, *Wuli Huaxue Xuebao* **5**, 565 (1989); *Chem. Abstr.* **112**, 118206f (1990).
- 1989AN/ZHE X.-W. An and X.-L. Zheng, *Acta Phys. Chim. Sinica* **5**, 487 (1989).
- 1989AZA E. C. Azandegbe, *Analisis* **17**, 285 (1989).
- 1989BRA/RYT B. K. Braxton and J. H. Rytting, *Thermochim. Acta* **154**, 27 (1989).
- 1989BRE/LIC P. Bret-Dibat and A. Lichanot, *Thermochim. Acta* **147**, 261 (1989).
- 1989BRO/CON J. M. Brown, A. D. Conn, G. Pilcher, M. L. P. Leitao, and M. Y. J. Yang, *J. Chem. Soc., Chem. Commun.* 1817

- (1989).
- 1989CEP/GON E. Cepeda, C. Gonzalez, and J. M. Resa, *J. Chem. Eng. Data* **34**, 270 (1989).
- 1989CHA/DEM A. Chauvet, A. De Maury, and J. Masse, *Thermochim. Acta* **147**, 17 (1989).
- 1989CHI/GRO M. P. Chiarelli and M. L. Gross, *Anal. Chem.* **61**, 1895 (1989).
- 1989CHI/HOS R. D. Chirico, I. A. Hossenlopp, A. Nguyen, W. V. Steele, and B. E. Gammon, *J. Chem. Thermodyn.* **21**, 179 (1989).
- 1989CHI/KRI R. D. Chirico, S. E. Kripmeyer, A. Nguyen, and W. V. Steele, *J. Chem. Thermodyn.* **21**, 1307 (1989).
- 1989CHI/NGU R. D. Chirico, A. Nguyen, W. V. Steele, M. M. Strube, and C. Tsonopoulos, *J. Chem. Eng. Data* **34**, 149 (1989).
- 1989COL/JIM M. Colomina, P. Jimenez, M. V. Roux, and C. Turrion, *J. Chem. Thermodyn.* **21**, 275 (1989).
- 1989GON/KRA T. Gondova, P. Kralik, and J. Gonda, *Thermochim. Acta* **156**, 147 (1989).
- 1989GRI/KON A. A. Grigor'ev, Y. V. Kondrat'ev, A. V. Suvorov, and V. I. Nikolaeva, *Zh. Obshch. Khim.* **59**, 1834 (1989).
- 1989HUI/VAN J. Huinink, K. Van Miltenburg, H. A. Oonk, and A. Schuijff, *J. Chem. Eng. Data* **34**, 99 (1989).
- 1989IMA/TAK A. Imamura, K. Takahashi, S. Murata, and M. Sakiyama, *J. Chem. Thermodyn.* **21**, 237 (1989).
- 1989JIM/ROU P. Jimenez, M. V. Roux, and C. Turrion, *J. Chem. Thermodyn.* **21**, 759 (1989).
- 1989KAT/TAN M. Kato and M. Tanaka, *J. Chem. Eng. Data* **34**, 206 (1989).
- 1989KIR/DOM D. R. Kirklin and E. S. Domalski, *J. Chem. Eng. Data* **21**, 449 (1989).
- 1989KON/STR S. N. Kondrat'ev and B. V. Strizhov, *Russ. J. Phys. Chem.* **63**, 656 (1989).
- 1989KRE/AZA A. G. Krestov and G. I. Azarova, *Izv. Vyssh. Uchebn. Zaved. Khim. Khim. Tekhnol.* **32**, 61 (1989); *Chem. Abstr.* **111**, 219794f (1989).
- 1989LUB/JAN J. Lubkowski, T. Janiak, J. Czerminski, and J. Blazejowski, *Thermochim. Acta* **155**, 7 (1989).
- 1989MAL/KAN J. J. Mallon and S. W. Kanto, *Macromolecules* **22**, 2070 (1989).
- 1989MIN M. E. Minas da Piedade, Ph.D. thesis, Technical University, Lisbon, 1989.
- 1989NIS/AND K. Nishida, Y. Ando, K. Ohwada, T. Mori, M. Koide, and A. Koukitsu, *J. Soc. Dyers Colour.* **105**, 112 (1989).
- 1989NIS/RAB V. P. Nistratov, I. B. Rabinovich, M. S. Sheiman, N. N. Smirnova, I. A. Zelyaev, and I. A. Feshchenko, *Zh. Fiz. Khim.* **63**, 1779 (1989).
- 1989PAR/GME S. J. Park and J. Gmehling, *J. Chem. Eng. Data* **34**, 399 (1989).
- 1989PIN/GON J. J. Pinvidic, A. Gonthier-Vassal, H. Szwarc, R. Ceolin, P. Toffoli, J. M. Teulon, and C. Guechot, *Thermochim. Acta* **153**, 37 (1989).
- 1989RIB/MON M. A. V. Ribeiro da Silva, M. J. S. Monte, and M. A. R. Matos, *J. Chem. Thermodyn.* **21**, 159 (1989).
- 1989RIB/REI M. A. V. Ribeiro da Silva and A. M. M. V. Reis, *J. Chem. Thermodyn.* **21**, 423 (1989).
- 1989RIB/RIB M. A. V. Ribeiro da Silva, M. D. M. C. Ribeiro da Silva, J. A. S. Teixeira, J. M. Bruce, P. M. Guyan, and G. Pilcher, *J. Chem. Thermodyn.* **21**, 265 (1989).
- 1989RIB/SOU M. D. M. C. Ribeiro Da Silva, P. Souza, and G. Pilcher, *J. Chem. Thermodyn.* **21**, 173 (1989).
- 1989ROR B. F. Rordorf, *Chemosphere* **18**, 783 (1989).
- 1989ROR/RUT B. F. Rordorf, S. Rutschmann, and P. Schiess, *Int. J. Mass Spectrom. Ion Process.* **95**, 211 (1989).
- 1989SAB/ELW R. Sabbah and E. El Watik, *Thermochim. Acta* **138**, 241 (1989).
- 1989SAK/IWA A. Sakoguichi, Y. Iwai, and J. Takenaka, *Kagaku Kogaku Ronbunshu* **15**, 166 (1989).
- 1989SAL/ABA S. R. Salman and K. F. Abas, *Thermochim. Acta* **152**, 381 (1989).
- 1989SAL/ABA2 S. R. Salman and K. F. Abas, *Thermochim. Acta* **142**, 245 (1989).
- 1989SAS/NGU K. Sasse, J. N'Guimbi, J. Jose, and J. C. Merlin, *Thermochim. Acta* **146**, 53 (1989).
- 1989SCH/PEN J. Schmidt and A. Penzkofer, *J. Chem. Phys.* **91**, 1403 (1989).
- 1989SHE/RAB M. S. Sheiman, I. B. Rabinovich, V. P. Nistratov, G. P. Kamelova, E. N. Karataev, and A. I. Feshchenko, *Zh. Fiz. Khim.* **63**, 836 (1989).
- 1989SHI/SHI Y. Shibutani and K. Shinra, *Bull. Chem. Soc. Jpn.* **62**, 1477 (1989).
- 1989STA/MAN G. M. Stack, L. Mandelkern, C. Krohnke, and G. Wegner, *Macromolecules* **22**, 4351 (1989).
- 1989STE/CHI W. V. Steele, R. D. Chirico, I. A. Hossenlopp, A. Nguyen, N. K. Smith, and B. E. Gammon, *J. Chem. Thermodyn.* **21**, 81 (1989).
- 1989STE/CHI2 W. V. Steele, R. D. Chirico, A. Nguyen, I. A. Hossenlopp, and N. K. Smith, *AIChE Symp. Ser.* **85**, 140 (1989).
- 1989STE/CHI3 W. V. Steele, R. D. Chirico, I. A. Hossenlopp, A. Nguyen, N. K. Smith, and B. E. Gammon, *J. Chem. Thermodyn.* **21**, 1121 (1989).
- 1989SUZ/SHI E. Suzuki, K. Shimomura, and K. Sekiguchi, *Chem. Pharm. Bull. (Tokyo)* **37**, 493 (1989).
- 1989VAN/VAN H. A. Van Doren, R. Van der Geest, C. A. Keuning, R. M. Kellogg, and H. Wynberg, *Liq. Cryst.* **5**, 265 (1989).
- 1989VAR/PAS R. M. Varushchenko and L. L. Pashchenko, *Russ. J. Phys. Chem.* **63**, 964 (1989).
- 1989VAR/SOM B. Varughese and J. T. Sommerfeld, *J. Chem. Eng. Data* **34**, 25 (1989).
- 1989VAS/LEB V. G. Vasil'ev and B. V. Lebedev, *Zh. Obshch. Khim.* **59**, 2415 (1989).
- 1989VIN/RUS C. Viney, T. P. Russel, L. E. Depero, and R. J. Twieg, *Mol. Cryst. Liq. Cryst.* **168**, 63 (1989).
- 1989VIN/WOR M. D. Vine and C. J. Wormald, *J. Chem. Thermodyn.* **21**, 1151 (1989).
- 1989VOR/BAR M. G. Voronkov, V. P. Baryshok, V. A. Klyuchnikov, A. N. Korchagina, and V. I. Pepekin, *J. Organomet. Chem.* **359**, 169 (1989).
- 1989VOR/KLY M. G. Voronkov, V. A. Klyuchnikov, S. N. Kolabin, G. N. Shvets, P. I. Varushin, E. N. Deryagina, N. A. Korchevin, and S. I. Tsvetnitskaya, *Dokl. Phys. Chem.* **307**, 650 (1989).
- 1989VOR/SOR M. G. Voronkov, M. S. Sorokin, V. A. Klyuchnikov, G. N. Shvets, and V. I. Pepekin, *J. Organomet. Chem.* **359**, 301 (1989).
- 1989WAN/YIN K. Wang, X.-G. Ying, M.-S. Xia, J.-Z. Xia, and Y. Hu, *J. Chem. Eng. Data* **34**, 126 (1989).
- 1989WU/SAN H. S. Wu and S. I. Sandler, *J. Chem. Eng. Data* **34**, 209 (1989).
- 1989ZHA/YAN Z. Y. Zhang and M. L. Yang, *Thermochim. Acta* **156**, 157 (1989).
- 1989ZN/ZHE X. Zn and X. Zheng, *Wuli Huaxue Xuebao* **5**, 487 (1989).
- 1990AMB/EWI D. Ambrose, M. B. Ewing, N. B. Ghiasee, and J. C. S. Ochoa, *J. Chem. Thermodyn.* **22**, 589 (1990).
- 1990AMB/GHI D. Ambrose and N. B. Ghiasee, *J. Chem. Thermodyn.* **22**, 307 (1990).
- 1990AN/HE X. An, J. He, and R.-H. Hu, *Thermochim. Acta* **169**, 331 (1990).
- 1990BAR/DEL G. Barone, G. Della Gatta, D. Ferro, and V. Piacente, *J. Chem. Soc., Faraday Trans.* **86**, 75 (1990).
- 1990BEC/DOG H. D. Beckhaus, B. Dogan, J. Pakusch, S. P. Verevkin, and C. Rüchardt, *Chem. Ber.* **123**, 2153 (1990).
- 1990BEC/DOG2 H. D. Beckhaus, B. Dogan, J. Schaetzer, S. Hellman, and C. Rüchardt, *Chem. Ber.* **123**, 137 (1990).
- 1990BEL/BAL K. Belina, G. Balazs, G. Varga, Gy. Hardy, J. Horvath, K. Nyitrai, A. R. Werninck, A. J. Shand, and G. H. W. Milburn, *Liq. Cryst.* **7**, 399 (1990).
- 1990BHA/LAL D. V. Bhalla and J. K. Lalla, *Drug Dev. Ind. Pharm.* **16**, 115 (1990).
- 1990BRI/WAD L. E. Briggner and I. Wadso, *J. Chem. Thermodyn.* **22**, 143 (1990).
- 1990CAB/BEL J. L. Cabezas and S. Beltran, *J. Chem. Eng. Data* **35**, 389 (1990).
- 1990CHA/GAD J. Chao, N. A. M. Gadalla, B. E. Gammon, K. N. Marsh, A. S. Rodgers, G. R. Somayajulu, and R. C. Wilhoit, *J. Phys. Chem. Ref. Data* **19**, 1547 (1990).

- 1990CHI/ARC R. D. Chirico, D. G. Archer, I. A. Hossenlopp, A. Nguyen, W. V. Steele, and B. E. Gammon, *J. Chem. Thermodyn.* **22**, 665 (1990).
- 1990CHI/GAM R. D. Chirico, B. E. Gammon, S. E. Kripmeyer, A. Nguyen, M. M. Strube, C. Tsonopoulos, and Steele, *J. Chem. Thermodyn.* **22**, 1075 (1990).
- 1990DAU/HUT T. E. Daubert and G. Hutchison, *AIChE Symp. Ser.* **279**, 93 (1990).
- 1990DAV/FIN R. H. Davies, A. Finch, and K. Salem, *Thermochim. Acta* **170**, 213 (1990).
- 1990DIA/DIO A. R. Dias, H. P. Diogo, M. E. Minas da Piedade, J. A. Martinho-Simoes, A. S. Carson, and E. H. Jamea, *J. Organomet. Chem.* **391**, 361 (1990).
- 1990DOG/BEC B. Dogan, H. D. Beckhaus, H. Birkhofer, and C. Rüchardt, *Chem. Ber.* **123**, 1365 (1990).
- 1990DOM/HEA E. S. Domalski and E. D. Hearing, *J. Phys. Chem. Ref. Data* **19**, 881 (1990).
- 1990DON/DRE J. R. Donnelly, L. A. Drewes, R. L. Johnson, W. D. Munslow, K. K. Knapp, and G. W. Sovocool, *Thermochim. Acta* **167**, 155 (1990).
- 1990DUT/KAH N. V. Dutt, A. P. Kahol, P. J. Reddy, and K. S. Raghunandan, *J. Chem. Eng. Data* **35**, 114 (1990).
- 1990FRI/DOG K. Fritzsche, H.-D. Dogan, Beckhaus, and C. Rüchardt, *Thermochim. Acta* **160**, 147 (1990).
- 1990HAI/GIL J. Haines and D. F. R. Gilson, *J. Chem. Soc., Faraday Trans.* **86**, 2617 (1990).
- 1990HAU/CON R. E. Hauffer, J. Conceicao, L. P. F. Chibante, Y. Chai, N. E. Byrne, S. Flanagan, M. M. Haley, S. C. O'Brien, C. Pan, Z. Xiao, W. E. Billups, M. A. Cinfolini, R. H. Hauge, J. L. Margrave, L. J. Wilson, R. F. Curl, and R. E. Smalley, *J. Phys. Chem.* **94**, 8634 (1990).
- 1990HIN/BID2 D. A. Hinckley, T. F. Bidleman, W. T. Foreman, and J. R. Tuschall, *J. Chem. Eng. Data* **35**, 232 (1990).
- 1990HWA/YOS D. R. Hwang, F. Yoshizawa, and M. Tamura, *Anzen Kogaku* **29**, 168 (1990); *Chem. Abstr.* **115**, 186382x (1991).
- 1990JIM/ROU P. Jimenez, M. V. Roux, and C. Turrión, *J. Chem. Thermodyn.* **22**, 721 (1990).
- 1990JIN/KAN J.-I. Jin, C.-S. Kang, and B. Y. Chung, *Bull. Korean Chem. Soc.* **11**, 245 (1990).
- 1990KAB/MIR G. Y. Kabo, E. A. Miroshnichenko, M. L. Frenkel, A. A. Kozyro, V. V. Simirskii, A. P. Krasulin, V. P. Vorob'eva, and Y. A. Lebedev, *Izv. Akad. Nauk SSSR Ser. Khim.* **39**, 750 (1990).
- 1990KAB/MIR2 G. Y. Kabo, E. A. Miroshnichenko, M. L. Frenkel, A. A. Kozyro, V. V. Simirskii, A. P. Krasulin, V. P. Vorob'eva, and Y. A. Lebedev, *Bull. Acad. Sci. USSR Div. Chem. Sci.* **39**, 662 (1990).
- 1990KAL/DRE J. Kalinowski, Z. Dreger, J. Godlewski, and P. Mondalski, *Chem. Phys. Lett.* **172**, 522 (1990).
- 1990KIS/SHO K. Kishore, H. K. Shobha, and G. J. Mattamal, *J. Phys. Chem.* **94**, 1642 (1990).
- 1990KNA/SAB P. Knauth and R. Sabbah, *Can. J. Chem.* **68**, 731 (1990).
- 1990KNA/SAB2 P. Knauth and R. Sabbah, *J. Struct. Chem.* **1**, 43 (1990).
- 1990KNA/SAB3 P. Knauth and R. Sabbah, *Thermochim. Acta* **164**, 145 (1990).
- 1990KOZ/SIM A. A. Kozyro, V. V. Simirskii, A. P. Krasulin, V. M. Sevruck, G. Y. Kabo, M. L. Frenkel, P. N. Gaponik, and Y. V. Grigor'ev, *Russ. J. Phys. Chem.* **64**, 348 (1990).
- 1990KOZ/SIM2 A. A. Kozyro, V. V. Simirskii, G. Y. Kabo, M. L. Frenkel, A. P. Krasulin, V. M. Sevruck, and N. A. Sokolov, *Zh. Fiz. Khim.* **64**, 2360 (1990).
- 1990KOZ/SIM3 A. A. Kozyro, V. V. Simirskii, A. P. Krasulin, V. M. Sevruck, G. Y. Kabo, M. L. Frenkel, P. N. Gaponik, and Y. V. Grigor'ev, *Zh. Fiz. Khim.* **64**, 656 (1990).
- 1990LEI/PIL M. L. P. Leitao, G. Pilcher, W. E. Acree, Jr., A. I. Zvaigzne, S. A. Tucker, and M. D. M. C. Ribeiro da Silva, *J. Chem. Thermodyn.* **22**, 923 (1990).
- 1990LEI/PIL2 M. L. P. Leitao, G. Pilcher, Y. Meng-Yan, J. M. Brown, and A. D. Conn, *J. Chem. Thermodyn.* **22**, 885 (1990).
- 1990LEN M. Lenka, *J. Chem. Thermodyn.* **22**, 473 (1990).
- 1990LOP/SIC M. I. Lopez and J. E. Sicre, *J. Phys. Chem.* **94**, 3860 (1990).
- 1990LYM/REE *Handbook of Chemical Property Estimation Methods*, edited by W. J. Lyman, W. F. Reehl, and D. H. Rosenblatt (American Chemical Society, Washington, D.C., 1990).
- 1990MAL/ALI I. P. Malkerova, A. S. Alikhanyan, V. G. Sevvast'yanov, Y. K. Grinberg, and V. I. Gorgoraki, *Zh. Neorg. Khim.* **35**, 413 (1990); *Chem. Abstr.* **112**, 186852p (1990).
- 1990MAN/AHU R. H. Manzo and A. A. Ahumanda, *J. Pharm. Sci.* **79**, 1109 (1990).
- 1990MEN/LIA G. F. Mendenhall, S. X. Liang, and E. H. T. Chen, *J. Org. Chem.* **55**, 3697 (1990).
- 1990MEN/PIL Y. Meng-Yan and G. Pilcher, *J. Chem. Thermodyn.* **22**, 893 (1990).
- 1990MES/TOD J. F. Messerly, S. S. Todd, H. L. Finke, S. H. Lee-Bechtold, G. B. Guthrie, W. V. Steele and R. D. R. D. Chirico, *J. Chem. Thermodyn.* **22**, 1107 (1990).
- 1990MEV/LIC I. M. Meva'a and A. Lichanot, *Thermochim. Acta* **158**, 335 (1990).
- 1990MID/KAT Y. Mido, H. Katano, T. Isono, M. Hashimoto, and H. Matsuura, *J. Phys. Chem.* **94**, 8070 (1990).
- 1990MIH/BAS P. Mihailovic, P. Bassoul, and J. Simon, *J. Phys. Chem.* **94**, 2815 (1990).
- 1990NES/NAZ T. N. Nesterova, A. G. Nazmutdinov, V. S. Tsvetkov, A. M. Rozhnov, and I. Yu. Roshchupkina, *J. Chem. Thermodyn.* **22**, 365 (1990).
- 1990ORT/SUS J. Ortego, P. Susial, and C. de Alfonso, *J. Chem. Eng. Data* **35**, 216 (1990).
- 1990PIA/FER V. Piacente, D. Ferro, and G. Della Gatta, *Thermochim. Acta* **158**, 79 (1990).
- 1990PIA/SCA V. Piacente and P. Scardala, *Thermochim. Acta* **159**, 193 (1990).
- 1990POM/PIA T. Pompili and V. Piacente, *Thermochim. Acta* **170**, 289 (1990).
- 1990PRA/KOH K. Praefcke, B. Kohne, and D. Singer, *Angew. Chem.* **102**, 200 (1990).
- 1990PUL/MAT R. Puliti, C. A. Mattia, G. Barone, G. Della Gatta, and D. Ferro, *Thermochim. Acta* **162**, 229 (1990).
- 1990RAM/KUD D. H. S. Ramkumar and A. P. Kudchadker, *Fluid Phase Equilib.* **55**, 207 (1990).
- 1990RIB/MAT M. A. V. Ribeiro Da Silva, M. A. R. Matos, and M. J. S. Monte, *J. Chem. Thermodyn.* **22**, 609 (1990).
- 1990RIB/MON M. A. V. Ribeiro Da Silva and M. J. S. Monte, *Thermochim. Acta* **171**, 169 (1990).
- 1990RIB/RIB M. A. V. Ribeiro Da Silva, M. D. M. C. Ribeiro Da Silva, M. C. S. S. Rangel, G. Pilcher, M. J. Akello, A. S. Carson, and E. H. Jamea, *Thermochim. Acta* **160**, 267 (1990).
- 1990RIC/YAN M. F. Richardson, O.-C. Yang, E. Novotny-Bregger, and J. D. Dunitz, *Acta Crystallogr., Sect. B: Struct. Sci.* **46**, 653 (1990).
- 1990ROZ/BAR A. M. Rozhnov, V. I. Barkov, K. G. Sharonov, and V. S. Tsvetkov, *J. Chem. Thermodyn.* **22**, 327 (1990).
- 1990SER/LAR F. Ser and Y. Larher, *J. Chem. Thermodyn.* **22**, 407 (1990).
- 1990SHI/HAY C. Shimasaki, S. Hayase, A. Murai, J. Takai, E. Tsukurimichi, and T. Yoshimura, *Bull. Chem. Soc. Jpn.* **63**, 1074 (1990).
- 1990SIN/GLI N. B. Singh and M. E. Glicksman, *Thermochim. Acta* **159**, 93 (1990).
- 1990SMI N. O. Smith, *J. Chem. Eng. Data* **35**, 387 (1990).
- 1990SOH/OKA M. Sohda, M. Okazaki, Y. Iwai, Y. Arai, A. Sakoguchi, R. Ueoka, and Y. Kato, *J. Chem. Thermodyn.* **22**, 607 (1990).
- 1990SOL/KAB T. V. Soldatova, G. Y. Kabo, A. A. Kozyro, and M. L. Frenkel, *Russ. J. Phys. Chem.* **64**, 177 (1990).
- 1990SPR/GRE A. Sprozynski and J. Gregorowicz, *Main Group Met. Chem.* **13**, 1 (1990).
- 1990SUE/MAT K. Suenaga, T. Matsuo, and H. Suga, *Thermochim. Acta* **163**, 263 (1990).
- 1990VAN/KEL R. van der Geest, R. M. Kellogg, and H. Wynberg, *Recl. Trav. Chim. Pays-Bas* **109**, 197 (1990).
- 1990VAN/PAV Y. Y. Van-Chin-Syan, Y. P. Pavlovskii, N. S. Kachurina, M. A. Dikii, Y. V. Panchenko, and G. A. Petrovskaya, *Russ. J. Phys. Chem.* **64**, 295 (1990).

- 1990VOR/ZVE V. M. Vorotyntsev, Y. B. Zverev, and D. A. Shamrakov, *Vysokochist. Veshchestva*, 123 (1990).
- 1990YAN/EIR S. Yano, M. Eiraku, M. Tomiyama, and T. Sugihara, *Liq. Cryst.* **7**, 537 (1990).
- 1990YAW/YAN C. L. Yaws, H. C. Yang, and W. A. Cowley, *Hydrocarbon Process., Int. Ed.*, 87 (1990).
- 1990YIN/LIN Z. Z. Ying and Y. M. Lin, *J. Chem. Thermodyn.* **22**, 617 (1990).
- 1990YUH/KIK S. Yuha, K. Kikuchi, M. Yoshida, and Y. Sugawara, *Mol. Cryst. Liq. Cryst.* **184**, 231 (1990).
- 1990ZHA/YAN Z. Zhang and M. L. Yang, *Thermochim. Acta* **169**, 263 (1990).
- 1991ACR W. E. Acree, Jr., *Thermochim. Acta* **189**, 37 (1991), and references therein.
- 1991ACR/TUC W. E. Acree, Jr., S. A. Tucker, A. I. Zvaigzne, Y. Mengyan, G. Pilcher, and M. D. M. C. Ribeiro da Silva, *J. Chem. Thermodyn.* **23**, 31 (1991).
- 1991AGA/LEG V. Agafonov, B. Legendre, D. Rodier, D. Wouessidgewe, and J. M. Dense, *J. Pharm. Sci.* **80**, 181 (1991).
- 1991BAS/SVO P. Basarova and V. Svoboda, *Fluid Phase Equilib.* **68**, 13 (1991).
- 1991BAU/WEB G. L. Baughman and E. J. Weber, *Dyes Pigm.* **16**, 261 (1991).
- 1991BED/BOO A. D. Bedells, C. Booth, and K. Viras, *Makromol. Chem.* **192**, 2099 (1991).
- 1991BUC/POT N. I. Buchan, R. M. Potemski, and T. F. Kuech, *J. Chem. Eng. Data* **36**, 372 (1991).
- 1991BYK/KIP T. A. Bykova, Y. G. Kiparisova, B. V. Lebedev, K. A. Mager, and Y. G. Gololobov, *Polym. Sci. U.S.S.R.* **33**, 537 (1991).
- 1991CEO/DUG R. Ceolin, J. Dugue, J. C. Rouland, C. Ralambosoa, and F. Lepage, *Int. J. Pharm.* **74**, 77 (1991).
- 1991CHI/BRA J. S. Chickos, C. M. Braton, D. G. Hesse, and J. F. Liebman, *J. Org. Chem.* **56**, 927 (1991) and references and supplemental material therein.
- 1991CHI/KNI R. D. Chirico, S. E. Knipmeyer, A. Nguyen, and W. V. Steele, *J. Chem. Thermodyn.* **23**, 431 (1991).
- 1991CHI/KNI2 R. D. Chirico, S. E. Knipmeyer, A. Nguyen, and W. V. Steele, *J. Chem. Thermodyn.* **23**, 759 (1991).
- 1991DES/DES A. G. De Souza, J. H. De Souza, and C. Airoldi, *J. Chem. Soc. Dalton Trans.* 1751 (1991).
- 1991DIK/KAB V. V. Dikii and G. Ya. Kabo, *Dokl. Akad. Nauk SSSR* **35**, 158 (1991); *Chem. Abstr.* **114**, 206233j.
- 1991ELG/YRA J. Elguero, G. I. Yranzo, J. Laynez, P. Jimenez, M. Menendez, J. Catalan, J. L. G. de Paz, F. Anvia, and R. W. Taft, *J. Org. Chem.* **56**, 3942 (1991).
- 1991ELI L. Elias, AH-DE/5-WP/17 International Civil Aviation Organization, Montreal, Canada, 23 September 1991 (unpublished), [as quoted in Ref. [2002JON/LIG]].
- 1991ELW/SAB L. El Watik and R. Sabbah, *Bull. Soc. Chim. Fr.* **128**, 344 (1991).
- 1991EWI/GOO M. B. Ewing and A. R. H. Goodwin, *J. Chem. Thermodyn.* **23**, 1163 (1991).
- 1991GIL/SAG M. S. Gill and V. S. Sagoria, *Indian J. Chem.* **30A**, 727 (1991); *Chem. Abstr.* **115**, 149065a (1991).
- 1991HEN/TSC F. Hentrich, C. Tschierske, and H. Zaszke, *Angew. Chem., Int. Ed. Engl.* **30**, 440 (1991).
- 1991HOP/MOL J. Hopken, M. Moller, and S. Boileau, *New Polymeric Mater.* **2**, 339 (1991).
- 1991HOR T. Hori, *Shikizai Kyokaishi* **64**, 83 (1991).
- 1991JEF/JAB C. W. Jefford, A. Jaber, J. Boukouvalas, and P. Tissot, *Thermochim. Acta* **188**, 337 (1991).
- 1991KAW/GIL N. T. Kawai, D. F. R. Gilson, and I. S. Butler, *Can. J. Chem.* **69**, 1758 (1991).
- 1991KNA/KUB *Thermochemical Properties of Inorganic Substances II*, edited by O. Knacke, O. Kubaaschewski, and K. Hesselmann, (Springer-Verlag, Berlin, 1991).
- 1991KOE/BOR R. Koelliker and G. Bor, *Inorg. Chem.* **30**, 2236 (1991).
- 1991LAB/WES A. K. Labban, E. F. Westrum, Jr., and J. A. R. Cheda, *Can. J. Chem.* **69**, 1796 (1991).
- 1991LEB/KUL B. V. Lebedev and T. G. Kulagina, *J. Chem. Thermodyn.* **23**, 1097 (1991).
- 1991LIU/GUO Z. Z. Liu, X. D. Guo, L. E. Straub, G. Erdos, R. J. Prankerd, R. J. Gozalez-Rothi, and H. Schreier, **8**, 57 (1991).
- 1991LUK/TIM V. A. Luk'yanova, L. P. Timofeeva, M. P. Kozina, V. N. Kirin, and A. V. Tarakanova, *Russ. J. Phys. Chem.* **65**, 439 (1991).
- 1991MAL/ALI I. P. Malkerova, A. S. Alikhanyan, I. Y. Filatov, I. L. Kazanskaya, and V. G. Sevast'yanov, *Zh. Neorg. Khim.* **36**, 3112 (1991); *Chem. Abstr.* **116**, 114007b (1991).
- 1991MIY/ENO S. Miyajima, T. Enomoto, T. Kusanagi, and T. Chiba, *Bull. Chem. Soc. Jpn.* **64**, 679 (1991).
- 1991PAN/SAM C. Pan, M. P. Sampson, Y. Chai, R. H. Hauge, and J. L. Margrave, *J. Phys. Chem.* **95**, 2944 (1991).
- 1991PIA/POM V. Piacente, T. Pompili, P. Scardala, and D. Ferro, *J. Chem. Thermodyn.* **23**, 379 (1991).
- 1991PIC/RYL M. Pickering, J. Rylance, R. W. H. Small, and D. Stubbley, *Acta Crystallogr., Sect. B: Struct. Sci.* **47**, 782 (1991).
- 1991RAB/SHE I. B. Rabinovich, M. S. Sheiman, G. P. Kamelova, V. P. Nistratov, and E. N. Karataev, *Zh. Fiz. Khim.* **65**, 2071 (1991).
- 1991RAD/RAD M. Radomska and R. Radomski, *J. Therm. Anal.* **37**, 693 (1991).
- 1991RAI/GEO U. S. Rai and S. George, *Cryst. Res. Technol.* **26**, 511 (1991).
- 1991ROZ/SAF A. M. Rozhnov, V. V. Safronov, S. P. Verevkin, K. G. Sharonov, and V. I. Alenin, *J. Chem. Thermodyn.* **23**, 629 (1991).
- 1991SAB/AN R. Sabbah and X. W. An, *Thermochim. Acta* **178**, 339 (1991).
- 1991SAB/AN2 R. Sabbah and X. W. An, *Thermochim. Acta* **179**, 81 (1991).
- 1991SAB/BUL R. Sabbah and E. N. L. E. Buluku, *Can. J. Chem.* **69**, 481 (1991).
- 1991SAB/HIR R. Sabbah, and H. Hirtz, *Bull. Chem. Soc. Jpn.*, 26 (1991).
- 1991SAB/WAT R. Sabbah and L. E. Watik, *Thermochim. Acta* **176**, 163 (1991).
- 1991SEV/ALI V. G. Sevast'yanov, A. S. Alikhanyan, T. I. Krasovskaya, I. Y. Filatov, S. V. Krasnodub-skaya, and N. T. Kuznetsov, *Vysokochist. Veshchestva*, 137 (1991); *Chem. Abstr.* **115**, 246646y (1991).
- 1991SEV/KRA V. G. Sevast'yanov, S. V. Krasnodubskaya, O. V. Kurtasov, A. S. Alikhanyan, N. T. Kuznetsov, and A. S. Zanina, *Vysokochist. Veshchestva*, 190 (1991); *Chem. Abstr.* **114**, 239291e (1991).
- 1991SHA/MIS K. G. Sharonov, Y. B. Mishentseva, A. M. Rozhnov, E. A. Miroshnichenko, and L. I. Korchatova, *J. Chem. Thermodyn.* **23**, 636 (1991).
- 1991SHE/WEI B. M. Sheikh-Ali and R. G. Weiss, *Liq. Cryst.* **10**, 575 (1991).
- 1991STE/CHI W. V. Steele, R. D. Chirico, S. E. Knipmeyer, and A. Nguyen, *J. Chem. Thermodyn.* **23**, 957 (1991).
- 1991TEL/LAR V. I. Tel'noi, V. N. Larina, N. T. Kuznetsov, V. G. Sevast'yanov, S. V. Krasnodubskaya, and I. Yu. Felatov, *Vysokochist. Veshchestva*, 194 (1991).
- 1991TOK/HAY A. Tokmakoff, D. R. Haynes, and S. M. George, *Chem. Phys. Lett.* **186**, 450 (1991).
- 1991URY/RON V. F. Ur'yash, O. V. Ronina, E. N. Karataev, I. V. Zimina, T. V. Kuznetsova, T. V. Petrun, I. A. Feshchenko, and A. D. Zorin, *Metalloorganicheskaya Khimiya* **4**, 1050 (1991); *Chem. Abstr.* **116**, 6605 (1992).
- 1991VOR/KLY M. G. Voronkov, V. A. Klyuchnikov, L. I. Marenkova, T. F. Danilova, G. N. Shvets, S. I. Tsvetnitskaya, and Yu. I. Khudobin, *J. Organomet. Chem.* **406**, 99 (1991).
- 1991VOR/KLY2 M. G. Voronkov, V. A. Klyuchnikov, E. V. Mironenko, G. N. Shvets, T. F. Danilova, and Yu. I. Khudobin, *J. Organomet. Chem.* **406**, 91 (1991).
- 1991VOR/KLY3 M. G. Voronkov, V. A. Klyuchnikov, E. V. Sokolova, T. F. Danilova, G. N. Shvets, A. N. Korchagina, L. E. Gussel'nikov, and V. V. Volkova, *J. Organomet. Chem.* **401**, 245 (1991).
- 1991WHI/PER M. A. White and A. Perrott, *J. Solid State Chem.* **90**, 87 (1991).

- 1991WIB/HAO K. B. Wiberg and S. Hao, *J. Org. Chem.* **56**, 5108 (1991).
- 1991WIB/WAL K. B. Wiberg and R. F. Waldron, *J. Am. Chem. Soc.* **113**, 7697 (1991).
- 1991WIL/WIL W. V. Wilding, L. C. Wilson, and G. M. Wilson, *AIChE Data Ser.* **1**, 6 (1991).
- 1991WU/LOC H. S. Wu, W. E. Locke III, and S. I. Sandler, *J. Chem. Eng. Data* **36**, 127 (1991).
- 1991WU/PIV H. S. Wu, K. A. Pividal, and S. I. Sandler, *J. Chem. Eng. Data* **36**, 418 (1991).
- 1991YAS/BYK V. G. Yasil'ev, T. A. Bykova, and B. V. Lebedev, *Zh. Fiz. Khim.* **65**, 51 (1991).
- 1991ZHA/HUA X. Zhang, W. Huang, F. Tan, G. Xu, and Z. Wu, *Acta Phys. Chim. Sinica* **1**, 106 (1991).
- 1992ABB/JIM J. L. M. Abboud, P. Jimenez, M. V. Roux, C. Turrión, and C. Lopez-Mardomingo, *J. Chem. Thermodyn.* **24**, 1299 (1992).
- 1992ABB/JIM2 J.-L. M. Abboud, P. Jimenez, M. V. Roux, C. Turrión, C. Lopez-Mardomingo, and G. Sanz, *J. Chem. Thermodyn.* **24**, 217 (1992).
- 1992ABR/OLA J. Abrefah, D. R. Olander, M. Balooch, and W. J. Siekhaus, *Appl. Phys. Lett.* **96**, 3566 (1992).
- 1992ABR/OLA2 J. Abrefah, D. R. Olander, M. Balooch and W. J. Siekhaus, *Appl. Phys. Lett.* **60**, 1313 (1992).
- 1992ACR/SIM W. E. Acree, Jr., V. V. Simirsky, A. A. Kozyro, A. P. Krasulin, G. J. Kabo, and M. L. Frenkel, *J. Chem. Eng. Data* **37**, 131 (1992).
- 1992ACR/TUC W. E. Acree, Jr., S. A. Tucker, and G. Pilcher, *J. Chem. Thermodyn.* **24**, 213 (1992).
- 1992BAB/WHA M. W. Babich, S. W. Hwang, and R. D. Mounts, *Thermochim. Acta* **210**, 83 (1992).
- 1992BAR/GIA G. Barone, C. Giancola, T. H. Lilley, C. A. Mattia, and R. Puliti, *J. Therm. Anal.* **38**, 2771 (1992).
- 1992CHA/MOU A. Chauvet, A. El Moussaoui, and J. Masse, *Thermochim. Acta* **210**, 133 (1992).
- 1992CHE/KOR H. S. Chen, A. R. Kortan, R. C. Haddon, and D. A. Fleming, *J. Phys. Chem.* **96**, 1016 (1992).
- 1992CHI/HES J. S. Chickos, D. G. Hesse, S. Y. Panshin, D. W. Rogers, M. Saunders, P. M. Uffer, and J. F. Liebman, *J. Org. Chem.* **57**, 1897 (1992).
- 1992DEC/AIR M. A. R. P. de Carvalho, C. Airoidi, and A. G. de Souza, *J. Chem. Soc. Dalton Trans.* 1235 (1992).
- 1992DIA/MIN A. R. Dias, M. E. Minas da Piedade, J. A. Martinho Simoes, J. A. Simoni, C. Teixeira, H. P. Diogo, Y. Meng-Yan, and G. Pilcher, *J. Chem. Thermodyn.* **24**, 439 (1992).
- 1992ELS/PRA M. Elsabee and R. Prankerd, *Int. J. Pharm.* **86**, 221 (1992). [Note: experimental fusion enthalpy (in J g⁻¹) was taken from the graph.]
- 1992FUR/BUT I. Furman, R. J. Butcher, R. M. Catchings, and R. G. Weiss, *J. Am. Chem. Soc.* **114**, 6023 (1992).
- 1992GAZ/SCH M. Gazicki, H. Schalko, P. Svasek, F. Olcaytug, and F. Kohl, *J. Vac. Sci. Technol. A* **10**, 51 (1992).
- 1992GER/GER P. A. Gerasimova, A. I. Gerasimova, N. E. Fedotova, N. B. Morozova, and I. K. Igumenov, *Izv. Vyssh. Uchebn. Zaved. Khim. Khim. Tekhnol.* **35**, 38 (1992).
- 1992GRA/SAN M. Gracia, F. Sanchez, P. Perez, J. Valero, and C. Gutierrez Losa, *J. Chem. Thermodyn.* **24**, 463 (1992).
- 1992HAW D. C. Hawker, *Chemosphere* **25**, 427 (1992).
- 1992HE/AN J. He, X.-W. An, and R.-H. Hu, *Acta Chim. Sin.* **50**, 943 (1992).
- 1992HEA/SIN E. A. Heath, P. Singh, and Y. Ebisuzaki, *Acta Crystallogr., Sect. C: Cryst. Struct. Commun.* **48**, 1960 (1992).
- 1992HOP/FAU J. Hopken, S. Faulstich, and M. Moller, *Mol. Cryst. Liq. Cryst. Sci. Technol., Sect. A* **210**, 59 (1992).
- 1992HOP/MOL J. Hopken and M. Moller, *Macromolecules* **25**, 2482 (1992).
- 1992HUA/WAN Z. Huang, Y. Wang, W. Huang, Y. Zhou, and F. Tan, *Huaxue Wuli Xuebao* **5**, 445 (1992).
- 1992HUA/ZHO W. Huang, Y. Zhou, F. Tan, X. Huang, and B. Yang, *Acta Phys. Chim. Sinica* **8**, 795 (1992).
- 1992HUA/ZHO2 W. Huang, Y. Zhou, F. Tan, G. Xu, Y. Hu, and H. Hu, *Huaxue Wuli Xuebao* **5**, 178 (1992).
- 1992JAD/FRA R. Jadot and M. Fraiha, *J. Chem. Eng. Data* **37**, 509 (1992).
- 1992JIM/ROU P. Jimenez, M. V. Roux, and C. Turrión, *J. Chem. Thermodyn.* **24**, 1145 (1992).
- 1992KAB/KOZ G. J. Kabo, A. A. Kozyro, V. S. Krouk, V. M. Sevruck, I. A. Yursha, V. V. Simirsky, and V. I. Gogolinsky, *J. Chem. Thermodyn.* **24**, 1 (1992).
- 1992KAM M. Kaminski, *Bull. Pol. Acad. Sci., Chem.* **40**, 149 (1992).
- 1992KAT H. Katayama, *J. Chem. Eng. Jpn.* **25**, 366 (1992).
- 1992KIM/SZY W. Kimizuka and J. Szydłowski, *Fluid Phase Equilib.* **77**, 261 (1992).
- 1992KOZ/KAB A. A. Kozyro, G. J. Kabo, V. S. Krouk, M. S. Sheiman, I. A. Yursha, V. V. Simirsky, A. P. Krasulin, V. M. Sevruck, and V. I. Gogolinsky, *J. Chem. Thermodyn.* **24**, 883 (1992).
- 1992KUL/LEB T. G. Kulagina, B. V. Lebedev, E. N. Karataev, and Yu. I. Amosov, *Zh. Fiz. Khim.* **66**, 1694 (1992).
- 1992LEB/SMI B. Lebedev, N. Smirnova, Y. Kiparisova, and K. Makovetsky, *Makromol. Chem.* **193**, 1399 (1992).
- 1992LEE/CHE C.-H. Lee, Q. Chen, R. S. Mohamed, and G. D. Holder, *J. Chem. Eng. Data* **37**, 179 (1992).
- 1992LEE/DEM C.-H. Lee, D. M. Dempsey, R. S. Mohamed, and G. D. Holder, *J. Chem. Eng. Data* **37**, 183 (1992).
- 1992LOU/ROU D. Lourdin, A. H. Roux, J.-P. E. Grolier, and J.-M. Buisine, *Thermochim. Acta* **204**, 99 (1992).
- 1992MAR/KOZ L. I. Marachuk, A. A. Kozyro, and V. V. Simirsky, *Zh. Prikl. Khim. (S.-Peterburg)* **64**, 875 (1992).
- 1992MAT/SAI C. K. Mathews, M. Sai Baba, T. S. L. Narasimhan, R. Balasubramanian, N. Sivaraman, T. G. Srinivasan, and P. R. V. Rao, *J. Phys. Chem.* **96**, 3566 (1992).
- 1992NGU/KAS J. N'Guimbi, H. Kasehgari, I. Mokbel, and J. Jose, *Thermochim. Acta* **196**, 367 (1992).
- 1992NYM/DES M. Nyman and S. B. Desu, *Mater. Res. Soc. Symp. Proc.* **243**, 381 (1992); *Chem. Abstr.* **119**, 172856a.
- 1992OGU/YAM K. Oguchi, M. Yamagishi, and A. Murano, *Fluid Phase Equilib.* **80**, 131 (1992).
- 1992OKA/OGU N. Okamoto, M. Oguni, and H. Suga, *Thermochim. Acta* **202**, 215 (1992).
- 1992PAN/CHA C. Pan, M. S. Chandasekharaiyah, D. Agan, R. H. Hauge, and J. L. Margrave, *J. Phys. Chem.* **96**, 6752 (1992).
- 1992PAP/PIM T. S. Papina, S. M. Pimenova, V. Yu. Zakharov, and V. P. Kolesov, *Zh. Khim. Termodin. Termokhim.* **1**, 207 (1992).
- 1992RAI/GEO U. S. Rai and S. George, *Can. J. Chem.* **70**, 2869 (1992).
- 1992REI/HAN H. Reinke, M. Hans, and H. Dehne, *Mikrochim. Acta* **108**, 143 (1992).
- 1992RIB/FER M. A. V. Ribeiro da Silva, M. L. C. C. H. Ferrao, and R. M. G. Esteves da Silva, *J. Chem. Thermodyn.* **24**, 1293 (1992).
- 1992RIB/FER2 M. A. V. Ribeiro Da Silva, M. L. C. C. H. Ferrao, R. M. C. Marques, and J. M. T. Lima, *J. Chem. Thermodyn.* **24**, 595 (1992).
- 1992RIB/FER3 M. A. V. Ribeiro Da Silva, M. L. C. C. H. Ferrao, M. J. S. Monte, R. M. G. Esteves Da Silva, and J. C. Ribeiro, *J. Chem. Thermodyn.* **24**, 585 (1992).
- 1992RIB/MAT M. A. V. Ribeiro Da Silva, A. R. Matos, Y. Meng-Yan, and G. Pilcher, *J. Chem. Thermodyn.* **24**, 107 (1992).
- 1992RIB/MON M. A. V. Ribeiro da Silva and M. J. S. Monte, *J. Chem. Thermodyn.* **24**, 1219 (1992).
- 1992RIB/MON2 M. A. V. Ribeiro Da Silva and M. J. S. Monte, *J. Chem. Thermodyn.* **24**, 715 (1992).
- 1992RIB/REI M. A. V. Ribeiro Da Silva, A. M. V. Reis, M. J. S. Monte, M. S. S. F. Bartolo, and J. A. Rodrigues, *J. Chem. Thermodyn.* **24**, 653 (1992).
- 1992RIC/MCC P. J. Richardson, D. F. McCafferty, and A. D. Woolfson, *Int. J. Pharm.* **78**, 189 (1992).
- 1992SAB/ELW R. Sabbah and L. El Watik, *Can. J. Chem.* **70**, 24 (1992).
- 1992SAB/ELW2 R. Sabbah and L. El Watik, *Thermochim. Acta* **197**, 381 (1992).
- 1992SAB/WAT R. Sabbah and L. E. Watik, *J. Therm. Anal.* **38**, 803 (1992).
- 1992SAL/WAN M. Salvi-Narkhede, B.-H. Wang, J. L. Adcock, and W. H. Van Hook, *J. Chem. Thermodyn.* **24**, 1065 (1992).
- 1992SER/TSI S. V. Serves, G. M. Tsivoulis, D. N. Sotiropoulos, P. V.

- Ioannou, and M. K. Jain, *Phosphorus, Sulfur Silicon Relat. Elem.* **77**, 99 (1992).
- 1992SMI/LEB N. N. Smirnova, B. V. Lebedev, E. G. Kiparisova, K. L. Makovetskii, and L. I. Gorbacheva, *Polym. Sci. U.S.S.R.* **34**, 72 (1992).
- 1992SRC/KER S. Srcic, J. Kerc, U. Urleb, I. Zupancic, G. Lahajnar, B. Kofler, and J. Smid-Korbar, *Int. J. Pharm.* **87**, 1 (1992).
- 1992STE/CHI W. V. Steele, R. D. Chirico, S. E. Knipmeyer, and A. Nguyen, *J. Chem. Thermodyn.* **24**, 245 (1992).
- 1992STE/CHI2 W. V. Steele, R. D. Chirico, S. E. Knipmeyer, and A. Nguyen, *J. Chem. Thermodyn.* **24**, 499 (1992).
- 1992SVO/KUB V. V. Svoboda, Kubes, and P. Basarova, *J. Chem. Thermodyn.* **24**, 333 (1992).
- 1992SVO/KUB2 V. V. Svoboda, Kubes, and P. Basarova, *J. Chem. Thermodyn.* **24**, 555 (1992).
- 1992TER/PAU A. Terol, B. Pauvert, A. Bouassab, P. Chevallet, and G. Cassanas, *Thermochim. Acta* **211**, 279 (1992).
- 1992VAN/COR M. G. M. Van der Vis, E. H. P. Cordfunke, R. J. M. Konings, G. J. K. Van der Berg, and J. C. Van Miltenburg, *J. Chem. Thermodyn.* **24**, 1103 (1992).
- 1992VER/BEC S. P. Verevkin, H.-D. Beckhaus, and C. Rüchardt, *Thermochim. Acta* **197**, 27 (1992).
- 1992VIL/WEI P. M. Vilalta and R. G. Weiss, *Liq. Cryst.* **12**, 531 (1992).
- 1992WAF/MUS E. Waffenschmidt, J. Musolf, M. Heuken, and K. Heime, *J. Supercond.* **5**, 119 (1992).
- 1992WEB L. A. Weber, *Fluid Phase Equilib.* **80**, 141 (1992).
- 1992WHI/WAS M. A. White, R. E. Wasylshen, P. E. Eaton, Y. Xiong, K. Pramod, and N. Nodari, *J. Phys. Chem.* **96**, 421 (1992).
- 1992ZHU/WU M.-S. Zhu, J. Wu, and Y.-D. Fu, *Fluid Phase Equilib.* **80**, 99 (1992).
- 1993ABB/JIM J.-L. M. Abboud, P. Jimenez, M. V. Roux, C. C. Turrión, and C. Lopez-Mardomingo, *Calor. Anal. Therm.* **24**, 181 (1993).
- 1993ACR W. E. Acree, Jr., *Thermochim. Acta* **219**, 97 (1993); and references therein.
- 1993ACR/SEV W. E. Acree, Jr., V. M. Sevruck, A. A. Kozyro, A. P. Krasulin, G. J. Kabo, and M. L. Frenkel, *J. Chem. Eng. Data* **38**, 101 (1993).
- 1993ACR/TUC W. E. Acree, Jr., S. A. Tucker, G. Pilcher, M. I. P. Andrade, and M. D. C. Ribeiro da Silva, *J. Chem. Thermodyn.* **25**, 653 (1993).
- 1993ACR/TUC2 W. E. Acree, Jr., S. A. Tucker, G. Pilcher, A. Chowdhary, M. D. C. Ribeiro Da Silva, and M. J. S. Monte, *J. Chem. Thermodyn.* **25**, 1253 (1993).
- 1993AIR/SAN C. Airoldi and L. S. Santos, *Struct. Chem.* **4**, 323 (1993).
- 1993AN/XIE X.-W. An and Y.-D. Xie, *Thermochim. Acta* **220**, 17 (1993).
- 1993AUC/MON A. Aucejo, J. B. Monton, R. Munoz, and M. Sanchoello, *J. Chem. Eng. Data* **38**, 160 (1993).
- 1993BAE A. K. Baev, *Russ. J. Phys. Chem.* **67**, 2161 (1993).
- 1993BRE/DUN R. Brettle, D. A. Dunmur, C. M. Marson, M. Pinol, and K. Toriyama, *Liq. Cryst.* **13**, 515 (1993).
- 1993BRU/MON M. C. Burguet, J. B. Monton, M. Sanchoello, and M. I. Vazquez, *J. Chem. Eng. Data* **38**, 328 (1993).
- 1993BRU/PIA B. Brunetti and V. Piacente, *J. Mater. Sci. Lett.* **12**, 1738 (1993).
- 1993BYK/KIP T. A. Bykova, E. G. Kiparisova, B. V. Lebedev, N. G. Senchenya, K. A. Mager, and Y. G. Gololobov, *Polym. Sci. U.S.S.R.* **35**, 8 (1993).
- 1993CHI/HES J. S. Chickos, D. G. Hesse, and J. F. Liebman, *Struct. Chem.* **4**, 261 (1993).
- 1993CHI/HOS J. S. Chickos, S. Hosseini, D. G. Hesse, and J. F. Liebman, *Struct. Chem.* **4**, 271 (1993). Heat capacity corrections to 298 K were estimated by the procedure outlined in the article.
- 1993CHI/KN1 R. D. Chirico, S. E. Knipmeyer, A. Nguyen, and W. V. Steele, *J. Chem. Thermodyn.* **25**, 1461 (1993).
- 1993CHI/KN2 R. D. Chirico, S. E. Knipmeyer, A. Nguyen, N. K. Smith, and W. V. Steele, *J. Chem. Thermodyn.* **25**, 729 (1993). (Note: calculated from the vapor pressure—temperature data.)
- 1993CON/VIA M. D. Contreras Claramonte, A. P. Vialard, and F. G. Vilchez, *Int. J. Pharm.* **94**, 23 (1993).
- 1993DEB/DWO J. De Brulin, A. Dworkin, H. Szwarc, J. Godard, R. Ceolin, C. Fabre, and A. Rassat, *Europhys. Lett.* **24**, 551 (1993).
- 1993DIK/KAB V. V. Diky, G. J. Kabo, A. A. Kozyro, A. P. Krasulin, and V. M. Sevruck, *J. Chem. Thermodyn.* **25**, 1169 (1993).
- 1993DIO/MIN H. P. Diogo, M. E. Minas da Piedade, A. C. Fernandes, J. A. Martinho Simoes, M. A. V. Riberio da Silva, and M. J. S. Monte, *Thermochim. Acta* **228**, 15 (1993).
- 1993DOM/SER U. Domanska, J. Serwatowski, A. Sporzynski, and M. Dabrowski, *Thermochim. Acta* **222**, 279 (1993).
- 1993ELM/CHA A. El Moussaoui, A. Chauvet, and J. Masse, *J. Therm. Anal.* **39**, 619 (1993).
- 1993ENG/WAN P. S. Engel, C. Wang, Y. Chen, C. Rüchardt, and H.-D. Beckhaus, *J. Am. Chem. Soc.* **115**, 65 (1993).
- 1993FAR/WIC J. Farkova and I. Wichterle, *Fluid Phase Equilib.* **90**, 143 (1993).
- 1993FER/MOR N. C. Ferreira, C. A. Moreira dos Santos, and I. C. C. Calegao, *Eclética Quim.* **18**, 49 (1993).
- 1993FUJ/OGU H. Fujimori and M. Oguni, *J. Phys. Chem. Solids* **54**, 607 (1993).
- 1993GRU A. Grubenmann, *Dyes Pigm.* **21**, 273 (1993).
- 1993HAK G. Hakvoort, *Calorim. Anal. Therm.* **24**, 177 (1993).
- 1993HE/AN J. He, X. An, and R. Hu, *Huaxue Xuebao* **51**, 1059 (1993); *Chem. Abstr.* **120**, 87930w (1994).
- 1993HER/BEC C. Herberg, H.-D. Beckhaus, T. Koertvelysi, and C. Rüchardt, *Chem. Ber.* **126**, 117 (1993).
- 1993HUS/SAR S. Husain, P. N. Sarma, G. Y. S. K. Swamy, and K. S. Devi, *J. Am. Oil Chem. Soc.* **70**, 149 (1993).
- 1993IGA/LOP J. M. Igartua, A. Lopez-Echarri, T. Breczewski, and I. Ruiz-Larrea, *Phase Transitions* **46**, 47 (1993).
- 1993JUA/CHE T. M. Juang, Y. N. Chen, S. H. Lung, Y. H. Lu, C. S. Hsu, and S. T. Wu, *Liq. Cryst.* **15**, 529 (1993).
- 1993KAB/KOZ G. J. Kabo, A. A. Kozyro, A. P. Krasulin, V. M. Sevruck, and L. S. Ivashkevich, *J. Chem. Thermodyn.* **25**, 485 (1993).
- 1993KAS/MOK H. Kasehgari, I. Mokbel, C. Viton, and J. Jose, *Fluid Phase Equilib.* **87**, 133 (1993).
- 1993KEL/SAK G. R. Kelm and A. A. Sakr, *Drug Dev. Ind. Pharm.* **19**, 809 (1993).
- 1993KOZ/KAB A. A. Kozyro, G. J. Kabo, A. P. Krasulin, V. M. Sevruck, V. V. Simirsky, M. S. Sheiman, and M. L. Frenkel, *J. Chem. Thermodyn.* **25**, 1409 (1993).
- 1993KUL/LEB T. T. Kulagina and B. V. Lebedev, *Journées Méditerranéennes de Calorimétrie et d'Analyse Thermique* **24**, 253 (1993); *Chem. Abstr.* **122**, 57054n (1995).
- 1993LEB/VAS B. Lebedev, V. Vasil'yev, and N. Novosyolova, *Makromol. Chem.* **194**, 739 (1993).
- 1993LEE/HOL C.-H. Lee and G. D. Holder, *J. Chem. Eng. Data* **38**, 320 (1993).
- 1993MAT/SAI C. K. Mathews, M. Sai Baba, T. S. Lakshmi Narasimhan, R. Balasubramanian, N. Sivaraman, T. G. Srinivasan, and P. R. Vasudeva Rao, *Fullerene Sci. Technol.* **1**, 101 (1993); see also *J. Phys. Chem.* **96**, 3566 (1992).
- 1993NAD/BER M. H. Nadais and M. G. Bernardo-Gil, *Fluid Phase Equilib.* **91**, 321 (1993).
- 1993OIS/HOR T. Oishi, H. Horie, and H. Shuyama, *Polym. J. (Tokyo, Jpn.)* **25**, 781 (1993).
- 1993OVC/SOB V. V. Ovchinnikov, A. A. Sobanov, and A. N. Pudovik, *Dokl. Phys. Chem.* **333**, 483 (1993).
- 1993PIA/FER V. Piacente, D. Ferro, and G. Della Gatta, *Thermochim. Acta* **223**, 65 (1993).
- 1993PIL/PAR G. Pilcher, O. G. Parchment, I. H. Hillier, F. Heatley, D. Fletcher, M. A. V. Ribeiro da Silva, M. L. C. C. H. Ferrao, M. J. S. Monte, and F. Jiye, *J. Phys. Chem.* **97**, 243 (1993).
- 1993RAM/BOU J. Rambaud, A. Bouassab, B. Pauvert, P. Chevallet, J.-P. Declercq, and A. Terol, *J. Pharm. Sci.* **82**, 1262 (1993).
- 1993REI/SAN R. Reich and V. Sanhueza, *J. Chem. Eng. Data* **38**, 341 (1993).
- 1993RIB/FER M. A. V. Ribeiro da Silva, M. L. C. C. H. Ferrao, and A. J. M. Lopes, *J. Chem. Thermodyn.* **25**, 229 (1993).

- 1993RIB/GIE M. A. V. Ribeiro da Silva, E. Giera, and M. J. S. Monte, *J. Alloys Compd.* **197**, 105 (1993).
- 1993RIB/MAT M. A. V. Ribeiro da Silva, M. A. R. Matos, M. J. S. Monte, M. C. B. Alves, and J. M. A. P. Vieira, *J. Chem. Thermodyn.* **25**, 579 (1993).
- 1993RIB/MON M. A. V. Ribeiro Da Silva, M. J. S. Monte, E. Giera, and W. Kakolowicz, *J. Chem. Thermodyn.* **25**, 1263 (1993).
- 1993RIB/RIB M. A. V. Ribeiro da Silva, M. D. M. C. Ribeiro da Silva, J. P. A. Paiva, I. M. C. S. Nogueira, A. M. Damas, J. V. Barley, M. M. Harding, M. J. Akello, and G. Pilcher, *J. Chem. Soc., Perkin Trans. 2* 1765 (1993).
- 1993RUE/SAR P. Ruelle, E. Sarraf, L. van den Berge, K. Seghers, M. Buchmann, and U. W. Kesselring, *Pharm. Acta Helv.* **68**, 49 (1993).
- 1993SAB/LE R. Sabbah and T. H. D. Le, *Can. J. Chem.* **71**, 1378 (1993).
- 1993SAB/PEM R. Sabbah and O. Pemenzi, *C. R. Acad. Sci. Paris Ser. II* **317**, 575 (1993).
- 1993SCH/BEC R. Schulze, H.-D. Beckhaus, and C. Rüdhardt, *Chem. Ber.* **126**, 1031 (1993).
- 1993SCH/KVA B. Schneider, J. Kvarda, P. Schmidt, M. Suchoparek, and I. Prokopova, *Collect. Czech. Chem. Commun.* **58**, 2403 (1993).
- 1993SER/SOT S. V. Serves, D. N. Sotiropoulos, P. V. Ioannou, and M. K. Jain, *Phosphorus, Sulfur Silicon Relat. Elem.* **81**, 181 (1993).
- 1993SHE/KAR V. B. Sheinin, V. V. Karasyov, S. A. Klyuyev, and B. D. Berezin, *Zh. Khim. Termodin. Termokhim.* **2**, 186 (1993); *Chem. Abstr.* **122**, 116291n (1995).
- 1993SIL/WEB A. M. Silva and L. A. Weber, *J. Chem. Eng. Data* **38**, 644 (1993).
- 1993SIN/CAR M. Siniti, J. Carre, J. M. Letoffe, J. P. Bastide, and P. Claudy, *Thermochim. Acta* **224**, 97 (1993).
- 1993STE/CHI W. E. Steele, R. D. Chirico, S. E. Knipmeyer, and A. Nguyen, *J. Chem. Thermodyn.* **25**, 965 (1993).
- 1993SUS/ORT P. Susial and J. Ortego, *J. Chem. Eng. Data* **38**, 434 (1993).
- 1993SUS/ORT2 P. Susial and J. Ortego, *J. Chem. Eng. Data* **38**, 647 (1993).
- 1993SYO/GOL S. V. Syoev, A. N. Golubenko, L. D. Nikulina, and T. N. Martynova, *Thermochim. Acta* **225**, 137 (1993).
- 1993TER/BOU A. Tero, A. Bouassab, B. Pauvert, P. Chevallet, and G. Cassanas, *Thermochim. Acta* **225**, 97 (1993).
- 1993TIE/FRA G. V. D. Tiers and C. V. Francis, *Thermochim. Acta* **226**, 311 (1993).
- 1993TOB/LAN P. Tobaly and G. Lanchec, *J. Chem. Thermodyn.* **25**, 503 (1993).
- 1993VAR/PUC R. M. Varushchenko, G. A. Puchkova, and A. I. Druzhinina, *Zh. Fiz. Khim.* **67**, 897 (1993).
- 1993VIL/HAM P. M. Vilalta, G. S. Hammond, and R. G. Weiss, *Langmuir* **9**, 1910 (1993).
- 1993WOL/KIM H. Wolff, W. Kimizuka, and J. Szydowski, *Fluid Phase Equilib.* **90**, 163 (1993).
- 1994ACR/TUC W. E. Acree, Jr., S. A. Tucker, G. Pilcher, and G. Toole, *J. Chem. Thermodyn.* **26**, 85 (1994).
- 1994AIM K. Aim, *J. Chem. Eng. Data* **39**, 591 (1994).
- 1994AIM2 K. Aim, *J. Chem. Thermodyn.* **26**, 977 (1994).
- 1994ALI/MAL A. S. Alikhanyan, I. P. Malkerova, N. P. Kuz'mina, V. K. Ivanov, and A. R. Kaul, *Zh. Neorg. Khim.* **39**, 1534 (1994).
- 1994ANT/SAN M. Antosik and S. I. Sandler, *J. Chem. Eng. Data* **39**, 584 (1994).
- 1994AUC/BUR A. Aucejo, M. C. Burguet, J. B. Monton, R. Munoz, M. Sanchothello, and M. I. Vazquez, *J. Chem. Eng. Data* **39**, 578 (1994).
- 1994BEC/RUE H.-D. Beckhaus, C. Rüdhardt, D. R. Lagerwall, L. A. Pauette, F. Wahl, and H. Prinzbach, *J. Am. Chem. Soc.* **116**, 11775 (1994).
- 1994BLA/BEL B. Blanco, S. Beltran, J. L. Cabezas, and J. Coca, *J. Chem. Eng. Data* **39**, 23 (1994).
- 1994BLU/PRA D. Blunk, K. Praefcke, and G. Legler, *Liq. Cryst.* **17**, 841 (1993).
- 1994BRA/DOU S. Braeuninger, S.-Q. Dou, H. Fuess, W. Schmahl, R. Strauss, and A. Weiss, *Ber. Bunsenges. Phys. Chem.* **98**, 1096 (1994).
- 1994CAR/LAY A. S. Carson, P. G. Laye, J. B. Pedley, A. M. Welsby, J. S. Chickos, and S. Hosseini, *J. Chem. Thermodyn.* **26**, 1103 (1994).
- 1994CHE/WES J. A. R. Cheda and E. F. Westrum, Jr., *J. Phys. Chem.* **98**, 2482 (1994).
- 1994CHI/GAM R. D. Chirico, B. E. Gammon, I. A. Hossenlopp, and W. V. Steele, *J. Chem. Thermodyn.* **26**, 469 (1994).
- 1994CHI/HOS R. D. Chirico, I. A. Hossenlopp, B. E. Gammon, S. E. Knipmeyer, and W. V. Steele, *J. Chem. Thermodyn.* **26**, 1187 (1994).
- 1994CHI/HOS2 R. D. Chirico, I. A. Hossenlopp, B. E. Gammon, S. E. Knipmeyer, and W. V. Steele, *J. Chem. Thermodyn.* **26**, 1219 (1994).
- 1994DAI/MAC S. Dai, L. Mac Toth, G. D. Del Cul, and D. H. Metcalf, *J. Chem. Phys.* **101**, 4470 (1994).
- 1994DAL/FER L. D'Alessio, D. Ferro, and V. Piacente, *J. Alloys Compd.* **209**, 207 (1994).
- 1994DIK/KAB V. V. Diky, G. J. Kabo, A. A. Kozyro, A. P. Krasulin, and V. M. Sevruk, *J. Chem. Thermodyn.* **26**, 1001 (1994).
- 1994DOU/FUE S. Q. Dou, H. Fuess, W. Schmahl, R. Strauss, and A. Weiss, *Z. Naturforsch., A: Phys. Sci.* **49A**, 594 (1994).
- 1994FAL/BID R. L. Falconer and T. F. Bidleman, *Atmos. Environ.* **28**, 547 (1994).
- 1994FER/MAR D. Ferro, R. Martino, and G. Della Gatta, *J. Chem. Thermodyn.* **26**, 183 (1994).
- 1994FON/MUN J. Font and J. Muntasell, *Mater. Res. Bull.* **29**, 1091 (1994).
- 1994FON/MUN2 J. Font and J. Muntasell, *Thermochim. Acta* **246**, 57 (1994).
- 1994FUJ/OGU H. Fujimori and M. Oguni, *J. Chem. Thermodyn.* **26**, 367 (1994).
- 1994GER/GER P. A. Gerasimov, A. I. Gerasimova, V. G. Isakova, and I. K. Igumenov, *Izv. Vyssh. Uchebn. Zaved. Khim. Khim. Tekhnol.* **37**, 48 (1994); *Chem. Abstr.* **120**, 228675f (1994).
- 1994GER/GER2 P. A. Gerasimov, A. I. Gerasimova, V. V. Krisyuk, and I. K. Igumenov, *Russ. J. Phys. Chem.* **68**, 199 (1994).
- 1994GOL/PER O. A. Golubchikov, G. L. Perlovich, and B. K. Naneishvili, *Zh. Fiz. Khim.* **68**, 2120 (1994).
- 1994GRA/PER M. Gracia, P. Perez, and J. Valero, *Fluid Phase Equilib.* **97**, 147 (1994).
- 1994HEN/DIE F. Hentrich, S. Diele, and C. Tschierske, *Liq. Cryst.* **17**, 827 (1994).
- 1994HUA/CHE W. Huang, S. Chen, F. Tan, and Z. Yang, *Wuli Huaxue Xuebao* **10**, 151 (1994).
- 1994KAB/KOZ G. J. Kabo, A. A. Kozyro, A. P. Marchand, V. V. Diky, V. V. Simirsky, L. S. Ivashkevich, A. P. Krasulin, V. M. Sevruk, and M. L. Frenkel, *J. Chem. Thermodyn.* **26**, 129 (1994).
- 1994KAR/FRA U. Karlson, W. T. Frankenberger, Jr., and W. F. Spencer, *J. Chem. Eng. Data* **39**, 608 (1994).
- 1994KIM/LEE K.-J. Kim, C.-H. Lee, and S.-K. Ryu, *J. Chem. Eng. Data* **39**, 228 (1994).
- 1994KON/HIL R. J. M. Konings and D. L. Hildenbrand, *J. Chem. Thermodyn.* **26**, 155 (1994).
- 1994KOR/SID M. V. Korobov and L. N. Sidorov, *J. Chem. Thermodyn.* **26**, 61 (1994).
- 1994KOU/HOS B. Koutek, M. Hoskovec, P. Vrckova, K. Konecny, and L. Feltl, *J. Chromatogr. A* **679**, 307 (1994).
- 1994KRA/GME M. A. Krähenbühl and J. Gmehling, *J. Chem. Eng. Data* **39**, 759 (1994).
- 1994LAR/MAR S. Larsen and K. Marthi, *Acta Crystallogr., Sect. B: Struct. Sci.* **50**, 373 (1994).
- 1994LEB/SMI B. Lebedev and N. Smirnova, *Macromol. Chem. Phys.* **195**, 35 (1994).
- 1994LEB/SMI2 B. Lebedev, N. Smirnova, N. Novosyolova, K. Makovetskii, and I. Ostrovskaya, *Macromol. Chem. Phys.* **195**, 1807 (1994).
- 1994LEB/SMI3 B. V. Lebedev, N. N. Smirnova, V. G. Yasil'ev, E. G. Ki-

- parisova, and V. I. Kleiner, *Polym. Sci., Ser. A Ser. B* **36**, 1171 (1994).
- 1994LEE/LIE M.-J. Lee, P.-J. Lien, and W.-K. Huang, *Ind. Eng. Chem. Res.* **33**, 2853 (1994).
- 1994LIE/MAR J. F. Liebman, J. A. Martinho-Simoes, and S. W. Slayden, in *The Chemistry of Organic Arsenic, Antimony and Bismuth Compounds*, edited by S. Patai (Wiley, New York, 1994), Chap. 4.
- 1994LIU/DIC K. Liu and R. M. Dickhut, *Chemosphere* **29**, 581 (1994).
- 1994LOP/VAN D. O. Lopez, J. Van Braak, J. L. L. Tamarit, and H. A. J. Oonk, *CALPHAD: Comput. Coupling Phase Diagrams Thermochem.* **18**, 387 (1994).
- 1994MEL/LIT A. P. Melissaris and M. H. Litt, *Macromolecules* **27**, 888 (1994).
- 1994MOR/KOB D. L. Morgan and R. Kobayashi, *Fluid Phase Equilib.* **97**, 211 (1994).
- 1994NAK/TAK K. Nakasone, K. Takamizawa, K. Shiokawa, and Y. Urabe, *Thermochim. Acta* **233**, 175 (1994).
- 1994OHT/YAM T. Ohta, O. Yamamuro, and H. Suga, *J. Chem. Thermodyn.* **26**, 319 (1994).
- 1994OPP/KRA H. Opperman, R. Krausz, U. Bruhn, and M. Balarin, *Z. Anorg. Allg. Chem.* **620**, 1110 (1994).
- 1994ORT/GAL J. Ortega and S. Galvan, *J. Chem. Eng. Data* **39**, 907 (1994).
- 1994PER/NAN G. L. Perlovich, B. K. Naneishvili, and O. A. Golubchikov, *Zh. Fiz. Khim.* **68**, 1932 (1994).
- 1994PIA/FER V. Piacente, D. Ferro, and G. Della Gatta, *Thermochim. Acta* **232**, 317 (1994); **235**, 292(E) (1994).
- 1994PIA/FON V. Piacente, D. Fontana, and P. Scardala, *J. Chem. Eng. Data* **39**, 231 (1994).
- 1994POP/DRA A. Popovic, G. Drazic, and J. Marsel, *Rapid Commun. Mass Spectrom.* **8**, 985 (1994).
- 1994RAK/VER K. Rakus, S. P. Verevkin, H.-D. Beckhaus, and C. Rüchardt, *Chem. Ber.* **127**, 2225 (1994).
- 1994RAK/VER2 K. Rakus, S. P. Verevkin, J. Schaezter, H.-D. Beckhaus, and C. Rüchardt, *Chem. Ber.* **127**, 1095 (1994).
- 1994REG/CHM A. Regosz, A. Chmielewska, T. Pelplinska, and P. Kowalski, *Pharmazie* **49**, 371 (1994).
- 1994RIB/FER M. A. V. Ribeiro da Silva and M. L. C. C. H. Ferrao, *J. Chem. Thermodyn.* **26**, 315 (1994).
- 1994RIB/FER2 M. A. V. Riberio da Silva, M. L. C. C. H. Ferrao, and J. H. Fang, *J. Chem. Thermodyn.* **26**, 839 (1994).
- 1994RIB/MAT M. A. V. Ribeiro da Silva, M. A. R. Matos, J. M. Goncalves, P. M. Johnson, G. Pilcher, P. M. Burkinshaw, and C. T. Mortimer, *Thermochim. Acta* **247**, 245 (1994).
- 1994RUZ/MAJ K. Ruzicka and V. Majer, *Space Sci. Rev.* **23**, 1 (1994).
- 1994RUZ/ZAB V. Ruzicka, Jr., M. Zabransky, K. Ruzicka, and V. Majer, *Thermochim. Acta* **245**, 121 (1994).
- 1994SAB/GOU R. Sabbah and M. Gouali, *Aust. J. Chem.* **47**, 1651 (1994).
- 1994SAB/TAB R. Sabbah, D. Tabet, and S. T. Belaadi, *Thermochim. Acta* **247**, 193 (1994).
- 1994SAB/TAB2 R. Sabbah, D. Tabet, and S. T. Belaadi, *Thermochim. Acta* **247**, 201 (1994).
- 1994SAI/LAK M. Sai Baba, T. S. Lakshmi Narasimhan, R. Balasubramanian, N. Sivaraman, and C. K. Mathews, *J. Phys. Chem.* **98**, 1333 (1994).
- 1994SAN/DEF N. Santilli, P. de Filippo, and A. Chianese, *J. Chem. Eng. Data* **39**, 179 (1994).
- 1994SCH/BEC F. Schaffer, H.-D. Beckhaus, H.-J. Rieger, and C. Rüchardt, *Chem. Ber.* **127**, 557 (1994).
- 1994SMI/MAT S. P. Smirnov, Y. N. Matiushin, and I. Z. Akmetov, International Civil Aviation Organization, Montreal, Canada, 14–18 February 1994 (unpublished), paper No. AH-DE/8-WP/12, as quoted in Ref. [2002JON/LIG].
- 1994SOU/PIN A. G. Souza, C. D. Pinheiro, L. C. Santos, and M. L. M. Melo, *Thermochim. Acta* **231**, 31 (1994).
- 1994SPI/LUI W. Spieksma, R. Luijk, and H. A. J. Govers, *J. Chromatogr. A* **672**, 141 (1994).
- 1994STE/CHI W. V. Steele, R. D. Chirico, A. Nguyen, and S. E. Knipmeyer, *J. Chem. Thermodyn.* **26**, 515 (1994).
- 1994STE/CHI2 W. V. Steele, R. D. Chirico, I. A. Hossenlopp, S. E. Knipmeyer, A. Nguyen, and N. K. Smith, in *Experimental Results for DIPPR 1990–1991 Projects on Phase Equilibria and Pure Component Properties*, edited by J. R. Cunningham and D. K. Jones (AIChE, New York, NY, 1994), pp. 188–215.
- 1994TAN/BOS R. P. Tangri and D. K. Bose, *Thermochim. Acta* **244**, 249 (1994).
- 1994TAN/SAB Z. C. Tan and R. Sabbah, *Sci. China, Ser. B: Chem., Life Sci., Earth Sci.* **37**, 641 (1994).
- 1994TAN/SAB2 Z. C. Tan and R. Sabbah, *Thermochim. Acta* **231**, 109 (1994).
- 1994TEL/SHE V. I. Tel'noi, M. S. Sheiman, G. P. Kamelova, and V. N. Larina, *Russ. J. Gen. Chem.* **64**, 977 (1994).
- 1994TER/CAS A. Terol, G. Cassanas, J. Nurit, B. Pauvert, A. Bouassab, J. Rambaud, and P. Chevallet, *J. Pharm. Sci.* **83**, 1437 (1994).
- 1994TER/PIA L. Terenzi and V. Piacenti, *Thermochim. Acta* **235**, 61 (1994).
- 1994TIP/JIM A. E. Tipping, P. Jimenez, E. Ballesteros, J.-L. M. Abboud, M. Yanez, M. Esseffar, and J. Elguero, *J. Org. Chem.* **59**, 1039 (1994).
- 1994TOR/HER L. A. Torres, I. Hernandez-Contreras, J. A. Guardado, and M. G. Gonzalez, *Meas. Sci. Technol.* **5**, 51 (1994).
- 1994WAN/KUO C.-C. Wang, J.-F. Kuo, and C.-Y. Chen, *Macromol. Chem. Phys.* **195**, 1493 (1994).
- 1994WAN/SHU F. Wania, W.-Y. Shui, and D. Mackay, *J. Chem. Eng. Data* **39**, 572 (1994).
- 1994WEL/VER F. Welle, S. P. Verevkin, M. Keller, H.-D. Beckhaus, and C. Rüchardt, *Chem. Ber.* **127**, 697 (1994).
- 1994WIB/MOR K. B. Wiberg, K. M. Morgan, and H. Maltz, *J. Am. Chem. Soc.* **116**, 11067 (1994).
- 1994YU/ZIP X. Yu, G. L. Zipp, and G. W. R. Davidson III, *Pharm. Res.* **11**, 522 (1994).
- 1994ZHI/SAB T. Zhi-Cheng and R. Sabbah, *Chin. Sci. Bull.* **39**, 1003 (1994).
- 1994ZIE/ZIE W. Zielenkiewicz, A. Zielenkiewicz, and K. L. Wierzchowski, *J. Solution Chem.* **23**, 1125 (1994).
- 1995ABB/JIM J.-L. M. Abboud, P. Jimenez, M. V. Roux, C. Turrión, C. Lopez-Mardomingo, A. Podosenin, D. W. Rogers, and J. F. Liebman, *J. Phys. Org. Chem.* **8**, 15 (1995).
- 1995ACR/BOT W. E. Acree, Jr., S. G. Bott, S. A. Tucker, M. D. M. C. Ribeiro da Silva, M. A. R. Matos, and G. Pilcher, *J. Chem. Thermodyn.* **27**, 1433 (1995).
- 1995ACR/TUC W. E. Acree, Jr., S. A. Tucker, M. D. M. C. Ribeiro da Silva, M. A. R. Matos, J. M. Goncalves, M. A. V. Ribeiro da Silva, and G. Pilcher, *J. Chem. Thermodyn.* **27**, 391 (1995).
- 1995ALM/FIN H. M. A. Al-Maydama, A. Finch, P. J. Gardner, and A. J. Head, *J. Chem. Thermodyn.* **27**, 575 (1995).
- 1995ALM/FIN2 H. M. A. Al-Maydama, A. Finch, P. J. Gardner, and A. J. Head, *J. Chem. Thermodyn.* **27**, 173 (1995).
- 1995ARC/BLA A. Arce, A. Blanco, J. Martinez-Ageitos, and A. Soto, *J. Chem. Eng. Data* **40**, 515 (1995).
- 1995AUC/GON A. Aucejo, V. Gonzalez-Alfaro, J. B. Monton, and M. I. Vazquez, *J. Chem. Eng. Data* **40**, 332 (1995).
- 1995BAE/MIK A. K. Baev, V. E. Mikhailov, and I. N. Chernyak, *Russ. J. Gen. Chem.* **65**, 977 (1995).
- 1995BAH/DUP A. Bah and N. Dupont-Pavlovsky, *J. Chem. Eng. Data* **40**, 869 (1995).
- 1995BEC/RUC H.-D. Beckhaus, C. Rüchardt, S. I. Kozhushkov, V. N. Belov, S. P. Verevkin, and A. de Meijere, *J. Am. Chem. Soc.* **117**, 11854 (1995).
- 1995BEL/AIT F. B. Belaribi, A. Ait-Kaci, and J. Jose, *J. Therm. Anal.* **44**, 1177 (1995).
- 1995BOW/HER C. J. Bowden, T. M. Herrington, A. M. Moseley, and R. Richardson, *Liq. Cryst.* **18**, 825 (1995).
- 1995BUS/ESC P. Bustamante and B. Escalera, *J. Pharm. Pharmacol.* **47**, 550 (1995).
- 1995CHE/WAN G.-H. Chen, Q. Wang, Z.-M. Ma, X.-H. Yan, and S.-J. Han, *J. Chem. Eng. Data* **40**, 361 (1995).
- 1995CHI/HES J. S. Chickos, D. G. Hesse, S. Hosseini, J. F. Liebman, G. D. Mendenhall, S. P. Verevkin, K. Rakus, H.-D. Beckhaus, and C. Rüchardt, *J. Chem. Thermodyn.* **27**, 693 (1995).

- 1995CHI/HOS J. S. Chickos, S. Hosseini, and D. G. Hesse, *Thermochim. Acta* **249**, 41 (1995).
- 1995CON/GOB J. A. Connor and A. Göbel, *J. Chem. Thermodyn.* **27**, 605 (1995).
- 1995CON/GOB2 J. A. Connor and A. Göbel, *Polyhedron* **14**, 3107 (1995).
- 1995CYP/JOH C. C. Cypcar, G. Johnson, and L. Mathias, *Polym. Prepr. (Am. Chem. Soc. Div. Polym. Chem.)* **36**, 402 (1995).
- 1995DAB/DOM M. Dabrowski, U. Domanska, J. Serwatowski, and A. Sporzynski, *Thermochim. Acta* **250**, 19 (1995).
- 1995DEJ/BUR A. Dejoj, M. C. Burguet, R. Munoz, and M. Sanchoello, *J. Chem. Eng. Data* **40**, 290 (1995).
- 1995DEL/SAB P. Del Vecchio, R. Sabbah, L. Abate, G. Garone, and G. Della Gatta, *Calorim. Anal. Therm.* **26**, 158 (1995).
- 1995DIO/MIN H. P. Diogo, M. E. Minas da Piedade, J. A. Martinho Simoes, and Y. Nagana, *J. Chem. Thermodyn.* **27**, 597 (1995).
- 1995DIO/SAN H. P. Diogo, R. C. Santos, P. M. Nunes, and M. E. Minas da Piedade, *Thermochim. Acta* **249**, 113 (1995).
- 1995EIS/DEN P. Eiselt, S. Denzinger, and H.-W. Schmidt, *Liq. Cryst.* **18**, 257 (1995).
- 1995ESP/BIS F. Espitalier, B. Biscans, P. S. Peyrigain, and C. Laguerie, *Fluid Phase Equilib.* **113**, 151 (1995).
- 1995FER/DEL P. Ferloni and G. Della Gatta, *Thermochim. Acta* **266**, 203 (1995).
- 1995FON/MUN J. Font and J. Muntassel, *J. Mater. Chem.* **5**, 1137 (1995).
- 1995GAB/MAR C. Gabaldon, P. Marzal, and J. B. Monton, *J. Chem. Eng. Data* **40**, 190 (1995).
- 1995GAR/CHA M. L. Garner, D. Chandra, and K. H. Lau, *J. Phase Equilib.* **16**, 24 (1995).
- 1995GUT/LIU J. P. Guthrie and Z. Liu, *Can. J. Chem.* **73**, 1395 (1995).
- 1995HAM/MEH A. Hammami and A. K. Mehrotra, *Fluid Phase Equilib.* **111**, 253 (1995).
- 1995HER/VER C. Herberg, S. P. Verevkin, M. Nölke, H.-D. Beckhaus, and C. Rüchardt, *Liebigs Ann.* **1995**, 515 (2006).
- 1995HSU/TSA C. S. Hsu, K. T. Tsay, A. C. Chang, S. R. Wang, and S. T. Wu, *Liq. Cryst.* **19**, 409 (1995).
- 1995JAB/LET S. Jabrane, J. M. Letoffe, and P. Claudy, *Thermochim. Acta* **258**, 33 (1995).
- 1995JIM/MEN P. Jiménez, V. Menéndez, M. V. Roux, and C. Turrión, *J. Chem. Thermodyn.* **27**, 679 (1995).
- 1995KAB/KOZ G. J. Kabo, A. A. Kozyro, V. V. Diky, V. V. Simirsky, L. S. Ivashkevich, A. P. Krasulin, V. M. Sevruck, A. P. Marchand, and M. L. Frenkel, *J. Chem. Thermodyn.* **27**, 707 (1995).
- 1995KAB/KOZ2 G. J. Kabo, A. A. Kozyro, V. V. Diky, and V. V. Simirsky, *J. Chem. Eng. Data* **40**, 371 (1995).
- 1995KEL/SCH S. M. Kelly, M. Schadt, and H. Seiberle, *Liq. Cryst.* **18**, 581 (1995).
- 1995KOB/OGU K. Kobashi and M. Oguni, *J. Chem. Thermodyn.* **27**, 979 (1995).
- 1995KRI/VES A. Kristl and G. Vesnaver, *J. Chem. Soc., Faraday Trans.* **91**, 995 (1995).
- 1995KSI/NAG A. Ksiazczak and I. Nagata, *Thermochim. Acta* **254**, 31 (1995).
- 1995LEB/BYK B. V. Lebedev, T. A. Bykova, E. A. Kiparisova, B. G. Velen'kaya, and V. N. Filatova, *Polym. Sci., Ser. A Ser. B* **37**, 126 (1995).
- 1995LEB/BYK2 B. V. Lebedev, T. A. Bykova, N. V. Novoselova, N. G. Senchenya, and K. A. Mager, *Polym. Sci., Ser. A Ser. B* **37**, 886 (1995).
- 1995LEB/CHI L. P. Lebedev, V. V. Chironov, A. N. Kizin, I. F. Falyakhov, I. Sh. Saifullin, O. R. Klyuchnikov, Y. D. Orlov, and A. Y. Lebedev, *Izv. Akad. Nauk Ser. Khim.* **4**, 660 (1995).
- 1995LEB/KUL B. V. Lebedev, T. G. Kulagina, N. N. Smirnova, E. G. Kiparisova, and N. V. Novoselova, *Russ. J. Phys. Chem.* **69**, 525 (1995).
- 1995LEB/KUL2 B. V. Lebedev, T. G. Kulagina, V. I. Telnoy, and V. G. Vasil'ev, *Macromol. Chem. Phys.* **196**, 3487 (1995).
- 1995LUK/KOZ V. A. Luk'yanova and M. P. Kozina, *Russ. J. Phys. Chem.* **69**, 1906 (1995).
- 1995LUK/KOZ2 V. A. Luk'yanova and M. P. Kozina, *Zh. Fiz. Khim.* **69**, 2094 (1995); *Chem. Abstr.* **124**, 68127f (1996).
- 1995MAC/JOY J. I. Macnab and J. A. Joy, *Thermochim. Acta* **259**, 31 (1995).
- 1995MAR/COS T. M. R. Maria, F. S. Costa, M. L. P. Leitao, and J. S. Redinha, *Thermochim. Acta* **269/270**, 405 (1995).
- 1995MAR/GAB P. Marzal, C. Gabaldon, A. Seco, and J. B. Monton, *J. Chem. Eng. Data* **40**, 589 (1995).
- 1995MIY/NAK S. Miyajima, A. Nakazato, N. Sakoda, and T. Chiba, *Liq. Cryst.* **18**, 651 (1995).
- 1995MOK/GUE I. Mokbel, T. Guetachew, and J. Jose, *ELDATA: Int. Electron. J. Phys.-Chem. Data* **2**, 167 (1995).
- 1995MOK/PAU I. Mokbel, V. Pauchon, and J. Jose, *ELDATA: Int. Electron. J. Phys. Chem. Data* **1**, 53 (1995).
- 1995MUR/PAI S. S. N. Murthy, A. Paikaray, and N. Arya, *J. Chem. Phys.* **102**, 8213 (1995).
- 1995NAS/LEN K. Nass, D. Lenoir, and A. Kettrup, *Angew. Chem., Int. Ed. Engl.* **34**, 1735 (1995).
- 1995NOL/VER M. Nolke, S. P. Verevkin, H.-D. Beckhaus, and C. Rüchardt, *Liebigs Ann.* **1995**, 41 (2006).
- 1995NUR/LEL J. Nurit, J. M. Leloup, B. Pauvert, O. Doumbia, P. Chevillet, and A. Terol, *J. Therm. Anal.* **45**, 535 (1995).
- 1995OHT/YAM T. Ohta, O. Yamamuro, and T. Matsuo, *J. Phys. Chem.* **99**, 2403 (1995).
- 1995ORT/GAL J. Ortego and S. Galvan, *J. Chem. Eng. Data* **40**, 699 (1995).
- 1995PAP/PIM T. S. Papina, S. M. Pimenova, V. A. Luk'yanova, and V. P. Kolesov, *Zh. Fiz. Khim.* **69**, 2148 (1995); *Chem. Abstr.* **124**, 68148p (1996).
- 1995PIA/GIG V. Piacente, G. Gigli, P. Scradala, A. Giustini, and D. Ferro, *J. Phys. Chem.* **99**, 14052 (1995).
- 1995RAK/VER K. Rakus, S. P. Verevkin, M. Keller, H.-D. Beckhaus, and C. Rüchardt, *Liebigs Ann.* **1995**, 1483 (2006).
- 1995RAK/VER2 K. Rakus, S. P. Verevkin, W.-H. Peng, H.-D. Beckhaus, and C. Rüchardt, *Liebigs Ann.* **1995**, 2059 (2006).
- 1995RIB/FER M. A. V. Ribeiro da Silva, M. L. C. C. H. Ferrao, and F. Jiye, *J. Chem. Eng. Data* **40**, 426 (1995).
- 1995RIB/MAT M. A. V. Ribeiro da Silva, M. A. R. Matos, and L. M. P. F. Amaral, *J. Chem. Thermodyn.* **27**, 565 (1995).
- 1995RIB/MAT2 M. A. V. Ribeiro da Silva, M. A. R. Matos, and L. M. P. F. Amaral, *J. Chem. Thermodyn.* **27**, 1187 (1995).
- 1995RIB/MAT3 M. A. V. Ribeiro da Silva, M. A. R. Matos, and L. M. P. F. Amaral, *J. Chem. Thermodyn.* **27**, 1141 (1995).
- 1995RIB/MAT4 M. A. V. Ribeiro da Silva, M. A. R. Matos, and V. M. F. Morais, *J. Chem. Soc., Faraday Trans.* **91**, 1907 (1995).
- 1995RIB/MON M. A. V. Ribeiro da Silva, M. J. S. Monte, and J. Huinink, *J. Chem. Thermodyn.* **27**, 175 (1995).
- 1995RIB/MON2 M. A. V. Ribeiro da Silva, M. J. S. Monte, and J. Huinink, *J. Alloys Compd.* **224**, 181 (1995).
- 1995RIB/MOR M. A. V. Ribeiro da Silva, V. M. F. Morais, M. A. R. Matos, and C. M. A. Rio, *J. Org. Chem.* **60**, 5291 (1995).
- 1995RIB/REI M. A. V. Ribeiro da Silva, A. M. M. V. Reis, and R. I. M. C. P. Faria, *J. Chem. Thermodyn.* **27**, 1365 (1995).
- 1995RIB/RIB M. A. V. Ribeiro da Silva, M. D. M. C. Ribeiro da Silva, M. J. S. Monte, J. M. Goncalves, and E. M. R. Fernandes, *J. Chem. Thermodyn.* **27**, 613 (1995).
- 1995RUL/RAK J. Rulewski, J. Rak, P. Dorurno, P. Skurski, and J. Blazejowski, *J. Therm. Anal.* **45**, 839 (1995).
- 1995SAB/AGU R. Sabbah and A. R. Aguilar, *Can. J. Chem.* **73**, 1538 (1995).
- 1995SAK/UEO A. Sakoguchi, R. Ueoka, Y. Kato, and Y. Arai, *Kagaku Kogaku Ronbunshu* **21**, 219 (1995).
- 1995SCH/PUS J. Schmelzer and J. Pusch, *Fluid Phase Equilib.* **110**, 183 (1995).
- 1995SCI/MAY J. Sciesinski, J. Mayer, T. Wasiutynski, E. Sciesinska, and J. Wojtowicz, *J. Phase Trans.* **54**, 15 (1995).
- 1995SKA/GOL S. A. Skackelford and J. F. Goldman, *Propellants, Explos., Pyrotech.* **20**, 1 (1995).
- 1995STE/CHI W. V. Steele, R. D. Chirico, A. B. Cowell, A. Nguyen, and S. E. Knipmeyer, *J. Chem. Thermodyn.* **27**, 1407 (1995).
- 1995STE/CHI2 W. V. Steele, R. D. Chirico, A. Nguyen, and S. E. Knipmeyer, *J. Chem. Thermodyn.* **27**, 311 (1995).
- 1995STI/DUA A. L. Stinchcomb, R. Dua, A. Paliwal, R. W. Woodard, and G. L. Flynn, *Pharm. Res.* **12**, 1526 (1995).
- 1995STR/ARG L. Stradella and M. Argentero, *Thermochim. Acta* **268**, 1

- (1995).
- 1995TAK/YAM S. Takahara, O. Yamamuro, and T. Matsuo, *J. Phys. Chem.* **99**, 9589 (1995).
- 1995TOB/WAT P. Tobaly and I. M. Watson, *J. Chem. Thermodyn.* **27**, 1211 (1995).
- 1995TOR/GUD L. A. Torres, G. Gudiño, I. Hernandez-Contreras, M. E. Ochoa, N. Farfan, and R. L. Santillan, *J. Chem. Thermodyn.* **27**, 779 (1995).
- 1995TOR/GUD2 L. A. Torres, R. Gudino, R. Sabbah, and J. A. Guardado, *J. Chem. Thermodyn.* **27**, 1261 (1995).
- 1995VAN/COR M. G. M. Van der Vis and E. H. P. Cordfunke, *Thermochim. Acta* **265**, 129 (1995).
- 1995VAR/DRO R. M. Varushchenko and A. I. Droujinina, *J. Chem. Thermodyn.* **27**, 355 (1995).
- 1995VER/BEC S. P. Verevkin, H.-D. Beckhaus, and C. Rüchardt, *Thermochim. Acta* **249**, 1 (1995).
- 1995VER/DOG S. P. Verevkin, B. Dogan, J. Hadrich, H.-D. Beckhaus, and C. Rüchardt, *J. Prakt. Chem.* **337**, 93 (1995).
- 1995VIE/CDE E. F. S. Vieira, J. C. de Queiroz, and F. S. Dias, *Thermochim. Acta* **256**, 249 (1995).
- 1995WOJ/TOU G. Wojcik, L. Toupet, C. Gors, and M. Foulon, *Phys. Status Solidi A* **147**, 99 (1995).
- 1995WOL/LAN H. Wolff, H. Landeck, H.-P. Frerichs, and E. Wolff, **109**, 245 (1995).
- 1995XUW/DAJ A. Xu-Wu and G. Da-Jun, *Thermochim. Acta* **253**, 235 (1995).
- 1995YAM/KIT K. Yamamoto, H. Kitamura, M. Momota, and K. Narita, *Thermochim. Acta* **267**, 313 (1995).
- 1995YAN/YIN L. Yang, H. Yin, W. Zhu, and S. Niu, *J. Therm. Anal.* **45**, 207 (1995).
- 1995YAS/TAK K. Yase, Y. Takahashi, N. Ara-kato, and A. Kawazu, *Jpn. J. Appl. Phys., Part 1* **34**, 636 (1995).
- 1996ACR/BOT W. E. Acree, Jr., S. G. Bott, S. A. Tucker, M. D. M. C. Ribeiro da Silva, M. A. R. Matos, and G. Pilcher, *J. Chem. Thermodyn.* **28**, 673 (1996).
- 1996BAC/GRZ D. D. Back, L. R. Grzyll, and M. Corrigan, *Thermochim. Acta* **272**, 53 (1996).
- 1996BAE/POD A. K. Baev, A. I. Podkovirov, and V. I. Kosirkin, *Izv. Vyssh. Uchebn. Zaved., Khim. Khim. Tekhnol.* **39**, 10 (1996).
- 1996BAR/BRU G. Bardi, B. Brunetti, and V. Piacente, *J. Chem. Eng. Data* **41**, 14 (1996).
- 1996BAU A. Bauer-Brandl, *Int. J. Pharm.* **140**, 195 (1996).
- 1996BEL/UFN B. Belhachemi, W. Ufnalsky, and Z. Derriche, *J. Chim. Phys. Phys.-Chim. Biol.* **93**, 1117 (1996).
- 1996BOE/MAR A. Boehncke, K. Martin, M. G. Muller, and H. G. Cammenga, *J. Chem. Eng. Data* **41**, 543 (1996).
- 1996BOL/MER O. V. Boltalina, V. Y. Merkov, A. Y. Borschevskiy, and A. Popovitch, *Mendeleev Commun.* **6**, 253 (1996).
- 1996BUR/DAG H. D. Burrows, M. da Graca, M. Miguel, A. P. Varela, and R. S. Becker, *Thermochim. Acta* **279**, 77 (1996).
- 1996BUR/KOL A. Burger, K. T. Koller, and W. M. Schiermeir, *Eur. J. Pharm. Biopharm.* **42**, 142 (1996).
- 1996BUR/MON M. C. Burguet, J. B. Monton, R. Munoz, J. Wisniak, and H. Segura, *J. Chem. Eng. Data* **41**, 1191 (1996).
- 1996BYK/MOR A. F. Bykov, N. B. Morozova, I. K. Igumenov, and S. V. Sysoev, *J. Therm. Anal.* **46**, 1551 (1996).
- 1996CAM/FIG H. K. Cammenga, L. O. Figura, and B. Zielasko, *J. Therm. Anal.* **47**, 427 (1996).
- 1996CEO/AGA R. Ceolin, V. Agafonov, D. Louer, V. A. Dzyabchenko, S. Toscani, and J. M. Cense, *J. Solid State Chem.* **122**, 186 (1996).
- 1996CHA/EMM P. Chassot and F. Emmenegger, *Inorg. Chem.* **35**, 5931 (1996).
- 1996CHI/SAB J. S. Chickos, R. Sabbah, S. Hosseini, and J. F. Liebman, *Struct. Chem.* **7**, 391 (1996).
- 1996CHI/STE R. D. Chirico, W. V. Steele, A. Nguyen, T. D. Klots, and S. E. Knipmeyer, *J. Chem. Thermodyn.* **28**, 797 (1996).
- 1996CIO/MEL I. Ciocazanu and V. Meltzer, *J. Therm. Anal.* **47**, 1755 (1996).
- 1996DAH/WIC O. Dahmani, I. Wichterle, and A. Ait-Kaci, *Fluid Phase Equilib.* **124**, 135 (1996).
- 1996DES/BRA A. G. de Souza, M. G. A. Brasilino, and C. Airoidi, *J. Chem. Thermodyn.* **28**, 1359 (1996).
- 1996DOM/HEA E. S. Domalski and E. D. Hearing, *J. Phys. Chem. Ref. Data* **25**, 1 (1996), and references therein.
- 1996DOM/MOOU U. Domanska, W. C. Moollan, and T. M. Letcher, *J. Chem. Eng. Data* **41**, 261 (1996).
- 1996DOU/FUE S.-Q. Dou, H. Fuess, R. Strau, and A. Weiss, *Z. Naturforsch., Z. Naturforsch., A: Phys. Sci.* **51A**, 777 (1996).
- 1996FIE/JOH C. Fiege, R. Joh, M. Petri, and J. Gmehling, *J. Chem. Eng. Data* **41**, 1431 (1996).
- 1996FON/ROS L. Fontana, L. Rosati, A. Sala, and L. Dobbetti, *Pharm. Acta Helv.* **71**, 341 (1996).
- 1996GAL/BOU H. E. Gallis, F. Bougrioua, H. A. J. Oonk, P. J. van Ekeren, and J. C. van Miltenburg, *Thermochim. Acta* **274**, 231 (1996).
- 1996GAL/VAN H. E. Gallis, G. J. K. van den Berg, and H. A. J. Oonk, *J. Chem. Eng. Data* **41**, 1303 (1996).
- 1996GIL/WIL N. F. Giles, H. L. Wilson, and W. V. Wilding, *J. Chem. Eng. Data* **41**, 1223 (1996).
- 1996GON/SUN Q. Gong, Y. Sun, Z. Huang, X. Zhou, Z. Gu, and D. J. Qiang, *Phys. Rev. B* **29**, 4981 (1996).
- 1996GOV/RUT H. A. J. Govers, E. Ruts, F. W. M. van der Wielen, and A. G. van Haelst, *Polycyclic. Aromat. Compd.* **9**, 75 (1996).
- 1996GRA B. Granzow, *J. Mol. Struct.* **381**, 127 (1996).
- 1996JIM/ROU P. Jimenez, M. V. Roux, J. Kulhanek, and O. Exner, *Struct. Chem.* **7**, 375 (1996).
- 1996KAM/ZIE M. Kaminski and W. Zielenkiewicz, *J. Chem. Thermodyn.* **28**, 153 (1996).
- 1996KEE/VAN E. M. D. Keegstra, V. van der Mieden, J. W. Zwikker, L. W. Jenneskens, A. Schouten, H. Kooijman, N. Veldman, and A. L. Spek, *Chem. Mater.* **8**, 1092 (1996).
- 1996KIR/CHI D. R. Kirklin, J. S. Chickos, and J. F. Liebman, *Struct. Chem.* **7**, 355 (1996).
- 1996KON/COR R. J. M. Konings, E. H. P. Cordfunke, and R. R. van der Laan, *Thermochim. Acta* **273**, 231 (1996).
- 1996KOU/HOS B. Koutek, M. Hoskovec, P. Vrkocova, K. Konecny, L. Feltl, and J. Vrkoc, *J. Chromatogr. A* **719**, 391 (1996).
- 1996LAB/HUB F. Labrize, L. G. Hubert-Pfalzgraf, L.-C. Daran, S. Halut, and P. Tobaly, *Polyhedron* **15**, 2707 (1996).
- 1996LEB/BYK B. V. Lebedev, T. A. Bykova, E. G. Kiparisova, and V. G. Vasil'yev, *Macromol. Chem. Phys.* **197**, 1553 (1996).
- 1996LEB/KUL B. V. Lebedev, T. G. Kulagina, A. A. Cheremukhina, and E. N. Karataev, *Russ. J. Gen. Chem.* **66**, 880 (1996).
- 1996LEB/SMI B. Lebedev, N. Smirnova, and V. Telnoy, *Macromol. Chem. Phys.* **197**, 3807 (1996).
- 1996LIN/WIC J. Linek, I. Wichterle, and K. Marsh, *J. Chem. Eng. Data* **41**, 1212 (1996).
- 1996LUE/MAG T. O. Lueddecke and J. W. Magee, *Int. J. Thermophys.* **17**, 823 (1996).
- 1996MIR/ORL E. A. Miroshinchenko, Yu. D. Orlov, L. I. Korchatova, V. P. Vorob'eva, and Yu. A. Lebedev, *Zh. Fiz. Khim.* **70**, 1583 (1996).
- 1996MOR/SYS N. B. Morozova, S. V. Sysoev, I. K. Igumenov, and A. N. Golubenko, *J. Therm. Anal.* **46**, 1367 (1996).
- 1996OHT/OSA T. Ohta, Y. Osamu, T. Matsuo, and H. Suga, *J. Phys. Chem.* **100**, 2353 (1996).
- 1996OLS J. D. Olson, *Fluid Phase Equilib.* **116**, 414 (1996).
- 1996OVC/MAK V. V. Ovchinnikov, T. B. Makeeva, L. I. Lapteva, and A. I. Konovalov, *Thermochim. Acta* **277**, 145 (1996).
- 1996PAP/KOL T. S. Papina, V. P. Kolesov, V. P. Vorobieva, and V. F. Golovkov, *J. Chem. Thermodyn.* **28**, 307 (1996).
- 1996PIA/GIG V. Piacente, G. Gigli, P. Scardala, A. Giustini, and G. Bardi, *J. Phys. Chem.* **100**, 9815 (1996).
- 1996POL/GUE M. Polednicek, T. Guetachew, J. Jose, V. Ruzicka, V. Rohac, and M. Zabransky, *ELDATA: Int. Electron. J. Phys. Chem. Data* **2**, 41 (1996).
- 1996PUL/BAR R. Puliti, G. Barone, C. Giancola, and C. A. Mattia, *J. Mol. Struct.* **382**, 197 (1996).
- 1996PYD/VAR M. Pyda, M. Varma-Nair, W. Chen, H. S. Aldrich, R. H. Schlosberg, and B. Wunderlich, *J. Therm. Anal.* **46**, 1093 (1996).
- 1996RAP/DES B. J. Rappoli and W. DeSisto, *Appl. Phys. Lett.* **68**, 2726

- (1996).
- 1996RIB/MON M. A. V. Ribeiro da Silva, M. J. S. Monte, and J. Huinink, *J. Chem. Thermodyn.* **28**, 413 (1996).
- 1996RIB/MOR M. A. V. Ribeiro da Silva, V. M. F. Morais, M. A. R. Matos, C. M. A. Rio, and C. M. G. S. Piedade, *Struct. Chem.* **7**, 329 (1996).
- 1996RIB/RIB M. A. V. Ribeiro da Silva, M. D. M. C. Ribeiro da Silva, M. F. B. M. Monteiro, M. L. A. C. N. Gomes, J. S. Chiccos, A. P. Smith, and J. F. Liebman, *Struct. Chem.* **7**, 367 (1996).
- 1996ROD/BER M. F. Rodrigues and M. G. Bernardo-Gil, *J. Chem. Eng. Data* **41**, 581 (1996).
- 1996ROU/JIM M. V. Roux, P. Jimenez J. Z. DaValos, J. L. M. Abboud, and M. T. Molina, *J. Chem. Thermodyn.* **28**, 1029 (1996).
- 1996ROU/JIM2 M. V. Roux, P. Jimenez, J. Z. Davalos, O. Castano, M. T. Molina R. Notario, M. Herrerros, and J. L. M. Abboud, *J. Am. Chem. Soc.* **118**, 12735 (1996).
- 1996SAB/GOU R. Sabbah and M. Gouali, *Can. J. Chem.* **74**, 500 (1996).
- 1996SCH F. P. Schwarz, *J. Solution Chem.* **25**, 471 (1996).
- 1996SHE/KAM M. S. Sheiman, G. P. Kamelova, K. G. Shvetsova, and V. P. Nistratov, *Russ. J. Phys. Chem.* **70**, 1105 (1996).
- 1996SLA/NOV I. B. Sladkov and N. Yu. Novikova, *Zh. Prikl. Khim. (S.-Peterburg)* **96**, 1840 (1996).
- 1996STE/CHI W. V. Steele, R. D. Chirico, S. E. Knipmeyer, and A. Nguyen, *J. Chem. Eng. Data* **41**, 1255 (1996).
- 1996STE/CHI2 W. V. Steele, R. D. Chirico, S. E. Knipmeyer, A. Nguyen, and N. K. Smith, *J. Chem. Eng. Data* **41**, 1285 (1996).
- 1996STE/CHI3 W. V. Steele, R. D. Chirico, S. E. Knipmeyer, A. Nguyen, N. K. Smith, and I. R. Tasker, *J. Chem. Eng. Data* **41**, 1269 (1996).
- 1996STR/BRA R. Strauss, S. Braun, S.-Q. Dou, H. Fuess, and A. Weiss, *Z. Naturforsch., A: Phys. Sci.* **51A**, 871 (1996).
- 1996SUS/SMI S. R. Susay, M. A. Smith, and G. G. Lockwood, *Zh. Fiz. Khim.* **83**, 864 (1996).
- 1996TEL/SHE V. I. Tel'noi, M. S. Sheiman, V. N. Larina, and G. P. Kamelova, *Russ. J. Phys. Chem.* **70**, 1261 (1996).
- 1996TOG/TOG R. K. Toghiani, H. Toghiani, and G. Verkateswarlu, *Fluid Phase Equilib.* **122**, 157 (1996).
- 1996TSY/DYA E. I. Tsyganova and L. M. Dyagileva, *Russ. Chem. Rev.* **65**, 315 (1996).
- 1996TSY/DYA2 E. I. Tsyganova and L. M. Dyagileva, *Usp. Khim.* **65**, 334 (1996), as quoted in Ref. [2000GIE].
- 1996TUR/EIC A. Türler, B. Eichler, D. T. Jost, D. Piquet, H. W. Gäggeler, K. E. Gregorich, B. Kadkhodayan, S. A. Kreek, D. M. Lee, M. Mohar, E. Sylwester, D. C. Hoffman, and S. Hübener, *Radiochim. Acta* **73**, 55 (1996).
- 1996UL/KLU P. Ulbig, M. Klüppel, and S. Schulz, *Thermochim. Acta* **271**, 9 (1996).
- 1996USH/SED V. S. Ushakov, S. M. Sedov, B. A. Knyazev, and B. I. Kuchkaev, *Zh. Fiz. Khim.* **70**, 1573 (1996).
- 1996VAN/ALV J. C. van Miltenburg, A. Alvarez-Larena, M. Labrador, L. Palacios, J. Rodriguez-Romero, E. Tauler, and E. Estop, *Thermochim. Acta* **273**, 31 (1996).
- 1996VAN/COR M. G. M. Van der Vis, E. H. P. Cordfunke, R. J. M. Konings, and A. Oskam, *J. Chem. Soc., Faraday Trans.* **92**, 973 (1996).
- 1996VAN/VAN A. G. van Haelst, F. W. M. van der Wielen, and H. A. J. Govers, *J. Chromatogr. A* **727**, 265 (1996).
- 1996VAN/YU Yu. Ya. Van-Chin-Syan, V. V. Kochubei, V. V. Sergeev, Yu. A. Raevskii, S. I. Gerasimchuk, and Kh. Z. Kotovich, *Russ. J. Phys. Chem.* **70**, 1789 (1996).
- 1996VAR/DRU R. M. Varushchenko, A. I. Druzhinina, and I. L. Pashchenko, *Fluid Phase Equilib.* **126**, 93 (1996).
- 1996VAR/PAS R. M. Varushchenko, L. L. Pashchenko, and A. I. Druzhinina, *Russ. J. Phys. Chem.* **70**, 208 (1996).
- 1996VER/BEC S. P. Verevkin, H.-D. Beckhaus, R. S. Belen'kaya, K. Rakus, and C. Rüchardt, *Thermochim. Acta* **279**, 47 (1996).
- 1996VER/PEN S. P. Verevkin, W.-H. Peng, H.-D. Beckhaus, and C. Rüchardt, *Struct. Chem.* **7**, 397 (1996).
- 1996VER/ZUF S. P. Verevkin, S. Züffle, H.-D. Beckhaus, and C. Rüchardt, *Thermochim. Acta* **285**, 1 (1996).
- 1996VIT/CHA C. Viton, M. Chavret, and J. Jose, *ELDATA: Int. Electron. J. Phys.-Chem. Data* **2**, 103 (1996).
- 1996WEB/DEF L. A. Weber and D. R. Defibaugh, *J. Chem. Eng. Data* **41**, 1477 (1996).
- 1996WEB/DEF2 L. A. Weber and D. R. Defibaugh, *J. Chem. Eng. Data* **41**, 382 (1996).
- 1996YAN/YU Z. Yang, C.-E. Yu, J. Cooke, Z. Ali-Adib, K. Viras, H. Matsuura, A. J. Ryan, and C. Booth, *J. Chem. Soc., Faraday Trans.* **92**, 3173 (1996).
- 1996YUA/MEN Z. Yuan and G. Meng, *Huaxue Shiji* **18**, 76 (1996); *Chem. Abstr.* **125**, 47592j (1996).
- 1996ZHA/HU J. Zhang, R. Hu, C. Zhu, F. Geng, and Q. Long, *Theory Pract. Energ. Mater.*, 133 (1996); *Chem. Abstr.* **126**, 133206p (1997).
- 1996ZIE/EIK T. Ziegler, O. Eikenberg, U. Bilitewski, and M. Grol, *Analyst (Cambridge, U.K.)* **121**, 119 (1996).
- 1996ZVE/CHE Yu. B. Zverev and S. G. Chesnokova, *Vysokochist. Veshchestva*, 75 (1996).
- 1997ACR/POW W. E. Acree, Jr., J. R. Powell, S. A. Tucker, M. D. M. C. Ribeiro da Silva, M. A. R. Matos, J. M. Goncalves, L. M. N. B. F. Santos, V. M. F. Morais, and G. Pilcher, *J. Org. Chem.* **62**, 3722 (1997).
- 1997ART/LAF H. Artigas, C. Lafuente, P. Cea, F. M. Royo, and J. S. Urieta, *J. Chem. Eng. Data* **42**, 132 (1997).
- 1997BAE A. E. Baev, *Russ. J. Inorg. Chem.* **42**, 587 (1997).
- 1997BAR/BRU G. Bardi, B. Brunetti, E. Ciccariello, and V. Piacente, *J. Alloys Compd.* **247**, 202 (1997).
- 1997BLA/BEL B. Blanco, S. Beltran, J. L. Cabezas, and J. Coca, *J. Chem. Eng. Data* **42**, 938 (1997).
- 1997BLO/KAB A. V. Blokhin, G. K. Kabo, A. A. Kozyro, L. S. Ivashkevich, A. P. Krasulin, V. V. Diky, V. Yu, and V. Vaksimuk, *Thermochim. Acta* **292**, 19 (1997).
- 1997BRU/GIG B. Brunetti, G. Gigli, E. Giglio, V. Piacente, and P. Scardala, *J. Phys. Chem. B* **101**, 10715 (1997).
- 1997BRU/GIU B. Brunetti, A. Giustini, and V. Piacente, *J. Chem. Thermodyn.* **29**, 239 (1997).
- 1997BRU/PAL B. Brunetti, C. Palchetti, and V. Piacente, *J. Mater. Sci. Lett.* **16**, 1395 (1997).
- 1997BUR/ROL A. Burger, J. M. Rollinger, and R. Brueggellar, *J. Pharm. Sci.* **86**, 674 (1997).
- 1997BUS/HAM D. Busing, F. Hamann, and A. Wurfinger, *Thermochim. Acta* **299**, 33 (1997).
- 1997CAL/FIL J. C. G. Calado, E. J. M. Filipe, and J. N. C. Lopes, *J. Chem. Thermodyn.* **29**, 1435 (1997).
- 1997CEN/MEL Z. Cenuse and V. Meltzer, *Analele Universit. F. Atii Bucure W. Sti. Chimie* **6**, 101 (1997).
- 1997CHA/YUE K.-T. Chan, K.-H. Yuen, H. Takayangi, S. Janadasa, and K.-K. Peh, *Phytochemistry* **46**, 1209 (1997).
- 1997CHE/LIA J. Chen, Y. Liang, and P. Ma, *Huagong Xuebao* **48**, 622 (1997).
- 1997CHI/WIL J. S. Chickos and J. A. Wilson, *J. Chem. Eng. Data* **42**, 190 (1997).
- 1997CIO/MEL I. Ciocazan, V. Meltzer, A. Nicolae, and R. Vilcu, *Calorim. Anal. Therm.* **28**, 267 (1997).
- 1997DAS/DAS J. P. Dasilva and A. M. Dasilva, *J. Chem. Eng. Data* **42**, 538 (1997).
- 1997DEF/CAR D. R. Defibaugh, E. Carrillo-Nava, J. J. Hurly, M. R. Moldover, J. W. Schmidt, and L. A. Weber, *J. Chem. Eng. Data* **42**, 488 (1997).
- 1997DEL A. Delle Site, *J. Phys. Chem. Ref. Data* **26**, 157 (1997).
- 1997DES/DES A. G. de Souza, F. de Souza Neto, J. H. de Souza, R. O. Macedo, J. B. L. de Oliveira, and C. D. Pinheiro, *J. Therm. Anal.* **49**, 679 (1997).
- 1997DEZ/POI V. de Zea Bermudez, C. Poinson, and M. B. Armand, *J. Mater. Chem.* **7**, 1677 (1997).
- 1997DOM/GON U. Domanska and J. A. Gonzalez, *Fluid Phase Equilib.* **129**, 139 (1997).
- 1997DUA/HWA H. A. Duarte-Garza, C.-A. Hwang, S. A. Kellerman, R. C. Miller, K. R. Hall, J. C. Holste, K. N. Marsh, and G. E. Gammon, *J. Chem. Eng. Data* **42**, 497 (1997).
- 1997ELD J. P. Elder, *J. Therm. Anal.* **49**, 897 (1997).
- 1997FIN/GAR A. Finch, P. J. Gardner, A. J. Head, and W. Xiaoping, *Thermochim. Acta* **298**, 191 (1997).

- 1997GAR/RED V. M. Garcia, M. I. Redondo, M. V. Redondo, and M. V. Roux, *J. Chem. Soc., Faraday Trans.* **93**, 2509 (1997).
- 1997GIL/BOT E. G. Gillan, S. G. Bott, and A. R. Barron, *Chem. Mater.* **9**, 796 (1997).
- 1997GOO M. A. Goodman, *J. Chem. Eng. Data* **42**, 1227 (1997).
- 1997GUD/TOR R. Gudino, L. A. Torres, M. Campos, R. L. Santillan, and N. Farfan, *J. Chem. Thermodyn.* **29**, 565 (1997).
- 1997HAL/WAN M. Hallquist, I. Wangberg, and E. Ljungstrom, *Environ. Sci. Technol.* **31**, 3166 (1997).
- 1997HER/ORT P. Hernández and J. Ortega, *J. Chem. Eng. Data* **42**, 1090 (1997).
- 1997HIL/MUL G. E. Hildebrand and C. C. Muller-Goymann, *J. Pharm. Sci.* **86**, 854 (1997).
- 1997IBR/FRA W. G. A. Ibrom and A. W. Frahm, *Arzeim. Forsch. Drug Res.* **47**, 662 (1997).
- 1997JEN/SAN M. Jenau, M. Sandmann, A. Wurflinger, and J. L. Tamarit, *Z. Naturforsch., A: Phys. Sci.* **52A**, 493 (1997).
- 1997JIM/ROU P. Jimenez, M. V. Roux, J. Z. Davalos, M. A. Marrtin-Luengo, and J. L. M. Abboud, *J. Chem. Thermodyn.* **29**, 1281 (1997).
- 1997KER/LOC P. Kerkoc, S. Lochran, R. T. Bailey, F. R. Cruickshank, D. Pugh, and J. Sherwood, *J. Appl. Phys.* **81**, 624 (1997); *Chem. Abstr.* **126**, 136332n (1997).
- 1997KLE A. Klein, Thermodynamics of liquid mixtures with lactones, *Fortschr.-Berich VDI Reihe 3: Verfahrenstechnik* **479**, 1 (1997).
- 1997KOJ/KAD Y. Kojima, H. Kadokura, Y. Okuhara, M. Matsumoto, and T. Mogi, *Integr. Ferroelectr.* **18**, 183 (1997).
- 1997KOR/API B. L. Korsounskii and T. A. Apina, International Annual Conference of ICT, 1997 (unpublished), p. 46; *Chem. Abstr.* **127**, 163997k (1997).
- 1997KOU/HOS B. Koutek, M. Hoskovec, P. Vrkocova, and L. Feltl, *J. Chromatogr. A* **759**, 93 (1997).
- 1997KRI/SYS V. V. Krisyuk, S. V. Sysoev, N. E. Fedotova, I. K. Iguenov, and N. V. Grigorieva, *Thermochim. Acta* **307**, 107 (1997).
- 1997KRO/VEL H. B. Krop, M. J. M. Velzen, J. R. Parsons, and H. A. J. Govers, *J. Am. Oil Chem. Soc.* **74**, 309 (1997).
- 1997KUL/LEB T. T. Kulagina and B. V. Lebedev, *Russ. J. Phys. Chem.* **71**, 520 (1997).
- 1997KUL/LEB2 T. G. Kulagina and B. V. Lebedev, *Russ. J. Phys. Chem.* **71**, 709 (1997).
- 1997LEB/KUL B. V. Lebedev and T. G. Kulagina, *J. Chem. Thermodyn.* **29**, 595 (1997).
- 1997LEE/CHA M.-J. Lee, Y.-K. Chang, H.-M. Lin, and C.-H. Chen, *J. Chem. Eng. Data* **42**, 349 (1997).
- 1997LIM/TUN L.-T. Lim and M. A. Tung, *J. Food. Sci.* **62**, 1061 (1997).
- 1997LOH/JOH J. Lohmann, R. Joh, and J. Gmehling, *J. Chem. Eng. Data* **42**, 1170 (1997).
- 1997LUK/KOL V. A. Luk'yanova, V. P. Kolesov, N. V. Avramenko, and V. F. Vorob'eva, *Russ. J. Phys. Chem.* **71**, 338 (1997).
- 1997MAK/KAB Y. V. Maksimuk, G. J. Kabo, A. A. Kozyro, V. V. Simirsky, A. P. Krasulin, and V. M. Sevruk, *J. Chem. Thermodyn.* **29**, 687 (1997).
- 1997MIN/BEH L. M. G. Minier and R. Behrens, Jr., *Propellants, Explos., Pyrotech.* **22**, 23 (1997).
- 1997NEA/BHA S. H. Neau, S. V. Bhandarkar, and E. W. Hellmuth, *Pharm. Res.* **14**, 601 (1997).
- 1997NEM/ACS K. Nematik, M. Acs, D. Kozma, and E. Fogassy, *J. Therm. Anal.* **48**, 691 (1997).
- 1997PAN/BOS F. Pan, C. Bosshard, M. S. Wong, C. Serbutoviez, K. Schenk, V. Gramlich, and P. Günter, *Chem. Mater.* **9**, 1328 (1997).
- 1997PEY/LET L. Peyrot, J. M. Letoffe, M. Elkhatib, J. P. Scharff, and H. Delalu, *Calorim. Anal. Therm.* **28**, 298 (1997).
- 1997PFE/SAB G. Pfefer, R. Sabbah, and R. Boistelle, *J. Appl. Crystallogr.* **30**, 527 (1997).
- 1997PIA/PAL V. Piacente, C. Palchetti, G. Gigli, and P. Scardala, *J. Phys. Chem. A* **101**, 4303 (1997).
- 1997PUL/DES R. Puliti, C. De Sena, and C. Giancola, *J. Therm. Anal.* **48**, 1249 (1997).
- 1997REU/BUS J. Reuter, D. Büsing, J. Ll. Tamarit, and A. Würflinger, *J. Mater. Chem.* **7**, 41 (1997).
- 1997RIB/GON M. A. V. Ribeiro da Silva, J. M. Goncalves, and G. Pilcher, *J. Chem. Thermodyn.* **29**, 253 (1997).
- 1997RIB/MAT M. A. V. Ribeiro da Silva, M. A. R. Matos, and L. M. P. F. Amaral, *J. Chem. Thermodyn.* **29**, 1545 (1997).
- 1997RIB/MAT2 M. A. V. Ribeiro da Silva, M. A. R. Matos, and L. M. P. F. Amaral, *J. Chem. Thermodyn.* **29**, 1535 (1997).
- 1997RIB/MAT3 M. A. V. Ribeiro da Silva, M. A. R. Matos, and L. M. P. F. Amaral, *J. Chem. Thermodyn.* **29**, 1129 (1997).
- 1997RIB/MAT4 M. A. V. Ribeiro da Silva, M. A. R. Matos, C. M. A. do Rio, and V. M. F. Morais, *J. Chem. Soc., Faraday Trans. 1* **93**, 3061 (1997).
- 1997RIB/MAT5 M. A. V. Ribeiro da Silva, M. A. R. Matos, and L. M. P. F. Amaral, *J. Chem. Thermodyn.* **29**, 295 (1997).
- 1997RIB/SAN M. A. V. Ribeiro da Silva, L. M. N. B. F. Santos, and G. Pilcher, *J. Chem. Thermodyn.* **29**, 757 (1997).
- 1997ROU/JIM M. V. Roux, P. Jimenez, M. A. Martin-Luengo, J. Z. Davalos, Z. Sun, R. S. Hosmane, and J. F. Liebman, *J. Org. Chem.* **62**, 2732 (1997).
- 1997SAB/PER R. Sabbah and L. Perez, *Can. J. Chem.* **75**, 357 (1997).
- 1997SAK/HOR T. Sako, S. Horiguchi, H. Ichimaru, and S. Nakagawa, *J. Chem. Eng. Data* **42**, 169 (1997).
- 1997SAN/ROC L. S. Santos, Jr., S. Roca, and C. Airoidi, *J. Chem. Thermodyn.* **29**, 661 (1997).
- 1997SAT/YAN K. Sato, J. Yano, I. Kawada, M. Kawano, F. Kaneko, and M. Suzuki, *J. Am. Oil Chem. Soc.* **74**, 1153 (1997).
- 1997SCH/BER W. C. Schinzer, M. S. Bergren, D. S. Aldrich, F. S. Chao, M. J. Dunn, A. Jeganathan, and L. M. Madden, *J. Pharm. Sci.* **86**, 1426 (1997).
- 1997SCH/VER F. Schaffer, S. P. Verevkin, H.-J. Rieger, H.-D. Beckhaus, and C. Rüchardt, *Liebigs Ann. Recueil* **1997**, 1333 (2006).
- 1997SMI/LEB N. N. Smirnova, B. V. Lebedev, E. G. Kiparisova, K. L. Makovetskii, and I. Y. Ostrovskay, **39**, 893 (1997).
- 1997SMI/LEB2 N. N. Smirnova, B. V. Lebedev, and V. G. Vasil'ev, *Zh. Obshch. Khim.* **66**, 199 (1997).
- 1997STE/CHI W. V. Steele, R. D. Chirico, A. B. Cowell, S. E. Knipmeyer, and A. Nguyen, *J. Chem. Eng. Data* **42**, 1053 (1997).
- 1997STE/CHI2 W. V. Steele, R. D. Chirico, S. E. Knipmeyer, and A. Nguyen, *J. Chem. Eng. Data* **42**, 1008 (1997).
- 1997STE/CHI3 W. V. Steele, R. D. Chirico, S. E. Knipmeyer, and A. Nguyen, *J. Chem. Eng. Data* **42**, 1021 (1997).
- 1997STE/CHI4 W. V. Steele, R. D. Chirico, S. E. Knipmeyer, A. Nguyen, and N. K. Smith, *J. Chem. Eng. Data* **42**, 1037 (1997).
- 1997TAN/ATA T. Tanaka, T. Atake, Z. Shi, C. Nakano, T. Enoki, G. Saito, and H. Inokuchi, *J. Phys. Chem. Solids* **58**, 295 (1997).
- 1997TES/PIK M. Tesconi, M. J. Pikal, and S. H. Yalkowsky, *J. Pharm. Sci.* **86**, 1299 (1997).
- 1997UKR/SOL E. A. Ukrainseva, D. V. Soldatov, and Yu. A. Dyadin, *Zh. Neorg. Khim.* **42**, 283 (1997).
- 1997VAR/DRU R. M. Varushchenko, A. I. Druzhinina, and E. L. Sorkin, *J. Chem. Thermodyn.* **29**, 623 (1997).
- 1997VAR/PAS R. M. Varushchenko, L. L. Pashchenko, V. M. Yuldasheva, and A. P. Orlov, *Russ. J. Phys. Chem.* **71**, 539 (1997).
- 1997VER S. P. Verevkin, *J. Chem. Thermodyn.* **29**, 891 (1997).
- 1997VER/MOR S. P. Verevkin, J. Morgenthaler, and C. Rüchardt, *J. Chem. Thermodyn.* **29**, 1175 (1997).
- 1997VER/NOL S. P. Verevkin, M. Nölke, H.-D. Beckhaus, and C. Rüchardt, *J. Org. Chem.* **62**, 4683 (1997).
- 1997VER2 S. P. Verevkin, *J. Chem. Thermodyn.* **29**, 1495 (1997).
- 1997VER3 S. P. Verevkin, *Thermochim. Acta* **307**, 17 (1997).
- 1997WEL/VER F. M. Welle, S. P. Verevkin, H.-D. Beckhaus, and C. Rüchardt, *Liebigs Ann. Recueil* **1997**, 155 (2006).
- 1997YUA/YAN Z. Yuan, P. Yang, G. Meng, and D. Peng, *Wuji Huaxue Xuebao* **13**, 168 (1997); *Chem. Abstr.* **127**, 144299t (1997).
- 1997YUA/ZHA J. Yuan, G.-L. Zhang, D.-Y. Huang, and H. Z. Zhang, *Liq. Cryst.* **22**, 693 (1997).
- 1997ZEM S. Zeman, *Thermochim. Acta* **302**, 11 (1997).
- 1997ZHA/GAO J.-B. Zhang, X.-G. Gao, Z.-C. Tan, S.-H. Meng, L.-M. Zhang, and S.-H. Li, *Thermochim. Acta* **297**, 1 (1997).
- 1997ZHA/HU J. Zhang, R. Hu, C. Zhu, G. Feng, and Q. Long, *Thermochim. Acta* **298**, 31 (1997).

- 1997ZHA/TAN J.-B. Zhang, Z.-C. Tan, S.-H. Meng, S.-H. Li, and L.-M. Zhang, *Thermochim. Acta* **307**, 11 (1997).
- 1998ABD/MEI M. A. Abdi and A. Meisen, *J. Chem. Eng. Data* **43**, 133 (1998).
- 1998ADD/CON S. J. Addison, J. A. Connor, and J. A. Kinkaid, *J. Organomet. Chem.* **554**, 123 (1998).
- 1998ASA/SOR S. Asahina, M. Sorai, and R. Eidenshink, *Liq. Cryst.* **24**, 201 (1998).
- 1998AUC/LOR A. Aucejo, S. Loras, R. Munoz, P. Reich, and H. Segura, *J. Chem. Eng. Data* **43**, 973 (1998).
- 1998BLA/KLI W. Blanke, G. Klingenberg, and F. Weber, *Int. J. Thermophys.* **19**, 653 (1998).
- 1998BOL/MAR O. V. Boltalina, V. Y. Markov, A. Y. Borschevskii, L. N. Sidorov, V. N. Bezmelnitsin, A. V. Elets'kii, and R. Taylor, *Rapid Commun. Mass Spectrom.* **12**, 1028 (1998).
- 1998BOL/MAR20 O. V. Boltalina, V. Y. Markov, A. Y. Borschevskii, V. Y. Davydov, L. N. Sidorov, V. N. Bezmelnitsin, A. N. Elets'kii, and R. Taylor, *Mendeleev Commun.* **8**, 141 (1998).
- 1998BOL/WIE A. Boller and H. G. Wiedemann, *J. Therm. Anal.* **53**, 431 (1998).
- 1998BOT/ELL M. Botoshansky, A. Ellern, N. Gasper, J.-O. Henck, and F. H. Herbstein, *Acta Crystallogr., Sect. B: Struct. Sci.* **54**, 277 (1998).
- 1998BRU/PIA B. Brunetti, V. Piacenti, and P. Scardala, *J. Chem. Eng. Data* **43**, 101 (1998).
- 1998BUS/PEN P. Bustamante, M. A. Pena, and J. Barra, *Int. J. Pharm.* **174**, 141 (1998).
- 1998BUS/ROM P. Bustamante, S. Romero, A. Pena, B. Escalera, and A. Reillo, *J. Pharm. Sci.* **87**, 1590 (1998).
- 1998CHI J. S. Chickos, *Thermochim. Acta* **313**, 19 (1998).
- 1998CHI/COW R. D. Chirico, A. B. Cowell, W. D. Good, T. D. Klots, S. E. Knipmeyer, A. A. Nguyen, R. Rau, J. W. Reynolds, N. K. Smith, and W. V. Steele, *J. Chem. Thermodyn.* **30**, 1423 (1998).
- 1998CHI/HES J. S. Chickos, D. Hesse, S. Hosseini, G. Nichols, and P. Webb, *Thermochim. Acta* **313**, 101 (1998).
- 1998DOM U. Domanska, *Pol. J. Chem.* **72**, 925 (1998).
- 1998DRO/TOM K. G. Drouillard, G. T. Tomy, D. C. G. Muir, and K. Friesen, *Envir. Toxicol. Chem.* **17**, 1252 (1998).
- 1998ELA/GIR S. K. El-Arini, D. Giron, and H. Leuenberger, *Eur. J. Pharm. Sci.* **3**, 557 (1998).
- 1998EWI/SAN M. B. Ewing and J. C. Sanchez Ochoa, *J. Chem. Thermodyn.* **30**, 189 (1998).
- 1998FUE/STR H. Fuess and R. Strauss, *J. Phys. Chem. B* **102**, 5329 (1998).
- 1998GEO/YOU M. A. George, K. M. Young, E. A. Robertson III, S. E. Beck, and G. Voloshin, *J. Chem. Eng. Data* **43**, 60 (1998).
- 1998GIO/GAZ F. Giordano, A. Gazzaniga, J. R. Moyano, P. Ventura, M. Zanol, T. Peveri, and L. Carima, *J. Pharm. Sci.* **87**, 333 (1998).
- 1998GIU/BRU A. Giustini, B. Brunetti, and V. Piacenti, *J. Chem. Eng. Data* **43**, 447 (1998).
- 1998GON/SZW A. Gonthier-Vassal and H. Szwarc, *Thermochim. Acta* **320**, 141 (1998).
- 1998GUD/TOR R. Gudino, L. A. Torres, R. L. Santillan, and N. Farfan, *J. Chem. Thermodyn.* **30**, 671 (1998).
- 1998HAT/SUZ T. Hatanaka, R. Suzuki, K. Katayama, and T. Koizumi, *Int. J. Pharm.* **168**, 199 (1998). (Experimental enthalpies of fusion and melting point temperatures were obtained through correspondence with the authors.)
- 1998HEL/OWE H. C. Helgeson, C. E. Owens, A. M. Knox, and L. Richard, *Geochim. Cosmochim. Acta* **1998**, 62, 985 (1998).
- 1998HIK/OKA T. Hikima, N. Okamoto, M. Hanaya, and M. Oguni, *J. Chem. Thermodyn.* **30**, 509 (1998).
- 1998ISS/ELA Y. M. Issa, A. L. El-Ansary, and W. Selim, *Anal. Lett.* **31**, 131 (1998).
- 1998JAB/LET S. Jabrane, J. M. Letoffe, and P. Claudy, *Thermochim. Acta* **311**, 121 (1998).
- 1998JAM/PAL M. E. Jamroz, M. Palczewska-Tulinska, D. Wyrzykowska-Stankiewicz, A. M. Szafranski, J. Polaczek, J. C. Dobrowolski, M. H. Jamroz, and A. P. Mazurek, **152**, 307 (1998).
- 1998KAB/BLO G. J. Kabo, A. V. Blokhin, A. A. Kozyro, V. V. Diky, L. S. Ivashkevich, A. P. Krasulin, V. M. Sevruck, and M. Frenkel, *Thermochim. Acta* **313**, 111 (1998).
- 1998KHO/RYS N. M. Khoretonenko, A. N. Rykov, and Y. M. Korenev, *Zh. Neorg. Khim.* **43**, 584 (1998); *Chem. Abstr.* **129**, 45952k (1998).
- 1998KIS/KAS V. D. Kiselev, E. A. Kashava, and A. I. Kononov, *Russ. J. Gen. Chem.* **68**, 1246 (1998).
- 1998KLO/LAU C. Kloc and R. A. Laudise, *J. Cryst. Growth* **193**, 563 (1998).
- 1998KOB/KYO K. Kobashi, T. Kyomen, and M. Oguni, *J. Phys. Chem. Solids* **59**, 667 (1998).
- 1998KOL/PIM V. P. Kolesov, S. M. Pimenova, V. A. Lukyanova, T. S. Kuznetsova, and M. P. Kozina, *J. Chem. Thermodyn.* **30**, 1455 (1998).
- 1998LEB/CHI V. P. Lebedev, V. V. Chirnonov, V. P. Vorob'eva, and Yu. N. Matyushin, *Khim. Fiz.* **17**, 54 (1998).
- 1998LIN/BEC A. Linden, H.-D. Beckhaus, S. P. Verevkin, C. Rüchardt, B. Ganguly, and B. Fuchs, *J. Org. Chem.* **63**, 8205 (1998).
- 1998LOB/BOH S. Löbbecke, M. A. Bohn, A. Pfeil, and H. Krause, *Proceedings of the 29th International Annual Conference of ICT, Karlsruhe, Federal Republic of German, 1998* (unpublished).
- 1998LUS/OLI Z. Lus, L. Olivier, R. Poupko, K. Muller, C. Krieger, and H. Zimmerman, *J. Am. Chem. Soc.* **120**, 5526 (1998).
- 1998MAG J. W. Magee, *Int. J. Thermophys.* **19**, 1397 (1998).
- 1998MAK/KAB Y. V. Maksimuk, G. J. Kabo, V. V. Simirsky, A. A. Kozyro, and V. M. Sevruck, *J. Chem. Eng. Data* **43**, 293 (1998).
- 1998MCK/FLO R. L. McKenney, Jr., T. G. Floyd, W. E. Stevens, T. G. Archibald, A. P. Marchand, G. V. M. Sharma, and S. G. Bott, *J. Energ. Mater.* **16**, 1 (1998).
- 1998MOK/RAU I. Mokbel, E. Rauzy, J. P. Meille, and J. Jose, *Fluid Phase Equilib.* **147**, 271 (1998).
- 1998MOR/KOP K. M. Morgan and D. A. Kopp, *J. Chem. Soc., Perkin Trans. 2* **2**, 2759 (1998).
- 1998MUR/BET P. Mura, G. P. Bettinetti, M. T. Faucci, and P. L. Parrini, *Thermochim. Acta* **321**, 59 (1998).
- 1998MUR/BET2 P. Mura, G. P. Bettinetti, A. Manderioli, M. T. Faucci, G. Bramanti, and M. Sorrenti, *Int. J. Pharm.* **166**, 189 (1998).
- 1998NIL/SOE F. Nilsson, O. Söderman, and I. Johansson, *J. Colloid Interface Sci.* **203**, 131 (1998).
- 1998NOL/VAL O. Noll, A. Valtz, D. Richon, T. Getachew-Sawaya, I. Mokbel, and J. Jose, *ELDATA: Int. Electron. J. Phys. Chem. Data* **4**, 105 (1998).
- 1998OJA/SUU V. Oja and E. M. Suuberg, *J. Chem. Eng. Data* **43**, 486 (1998).
- 1998OLS J. D. Olson, *Fluid Phase Equilib.* **150/151**, 713 (1998).
- 1998OON/VAN H. A. J. Oonk, P. R. van der Linde, J. Huinink, and J. G. Blok, *J. Chem. Thermodyn.* **30**, 897 (1998).
- 1998PAL/WEH G. R. Palmieri, P. Wehrle, and S. Martelli, *Drug Dev. Ind. Pharm.* **24**, 653 (1998).
- 1998PAN/MAL R. Pankajavalli, C. Mallika, O. M. Sreedharan, M. Premila, and P. Gopalan, *Thermochim. Acta* **316**, 101 (1998).
- 1998PAP/KOL T. S. Papina, V. P. Kolesov, V. A. Lukyanova, V. F. Golovkov, C. A. Chernov, and V. P. Vorobieva, *J. Chem. Thermodyn.* **30**, 431 (1998).
- 1998PAP/LUK T. S. Papina, V. A. Luk'yanova, V. P. Golovkov, S. A. Chernov, and V. P. Vorob'eva, *Zh. Fiz. Khim.* **72**, 7 (1998).
- 1998PAR/GIL R. M. Paroli, D. F. R. Gilson, and I. S. Butler, *Can. J. Chem.* **76**, 1365 (1998). (Note: the authors report only an entropy of transition and a temperature range for the first two transitions. The temperature that is listed is the midpoint of the reported temperature range, and the transition enthalpy is computed as the product of the reported transition entropy times transition temperature as defined above.)
- 1998PAR/GIL2 R. M. Paroli, D. F. R. Gilson, and I. S. Butler, *J. Solid State Chem.* **136**, 16 (1998).
- 1998PRI/HAW D. M. Price and M. Hawkins, *Thermochim. Acta* **315**, 19 (1998).

- 1998RAI/RAI U. S. Rai and R. N. Rai, *J. Therm. Anal.* **53**, 883 (1998).
- 1998RIB/CAR M. A. V. Ribeiro da Silva, A. P. S. M. C. Carvalho, M. J. S. Monte, and E. Giera, *J. Chem. Thermodyn.* **30**, 815 (1998).
- 1998RIB/GON M. A. V. Ribeiro da Silva and J. M. Goncalves, *J. Chem. Thermodyn.* **30**, 1465 (1998).
- 1998RIB/MAT M. D. M. C. Ribeiro da Silva, M. A. R. Matos, M. C. Vas, L. M. N. B. F. Santos, G. Pilcher, W. E. Acree, Jr., and J. R. Powell, *J. Chem. Thermodyn.* **30**, 869 (1998).
- 1998RIB/MAT2 M. D. M. C. Ribeiro da Silva, M. A. R. Matos, G. Pilcher, and W. E. Acree, Jr., *J. Chem. Thermodyn.* **30**, 271 (1998).
- 1998RIB/MAT3 M. A. V. Ribeiro da Silva, M. A. R. Matos and J. M. Goncalves, *J. Chem. Thermodyn.* **30**, 299 (1998).
- 1998ROH/RUZ V. Rohac, V. Ruzicka, K. Ruzicka, and K. Aim, *J. Chem. Eng. Data* **43**, 770 (1998).
- 1998ROU/JIM M. V. Roux, P. Jimenez, J. Z. Davalos, C. Turrión, H. Y. Afeefy, and J. F. Liebman, *J. Chem. Soc., Faraday Trans. 1* **94**, 887 (1998).
- 1998RUU/MOK K. Ruuzicka, I. Mokbel, V. V. Majer, Ruuzicka, J. Jose, and M. Zabransky, *Fluid Phase Equilib.* **148**, 107 (1998).
- 1998SAB/DAS R. Sabbah and M. E. da Silva Eusebio, *Can. J. Chem.* **76**, 18 (1998).
- 1998SAB/HEV R. Sabbah and R. Hevia, *Thermochim. Acta* **313**, 131 (1998).
- 1998SAB/HEV2 R. R. Sabbah, Hevia, and D. Tabet, *Thermochim. Acta* **316**, 1 (1998).
- 1998SAB/KUA R. Sabbah, D. N. Kuakuvu, and L. Perez, *Thermochim. Acta* **316**, 137 (1998).
- 1998SAB/TAB R. Sabbah, D. Tabet, and M. E. S. Eusebio, *Thermochim. Acta* **315**, 93 (1998).
- 1998SEM/WIL B. Semeniuk and H. Wilczura-Wachnik, *Fluid Phase Equilib.* **152**, 337 (1998).
- 1998SHE/NIS M. S. Sheiman, V. P. Nistratov, and G. P. Kamelova, *Russ. J. Phys. Chem.* **72**, 1620 (1998).
- 1998SOR/KIM M. Sorai, K. Kimura, A. Weiss, and R. Strauss, *J. Chem. Thermodyn.* **30**, 1441 (1998).
- 1998STE/ZAW Z. Stec and J. Zawadiak, *Pol. J. Appl. Chem.* **42**, 237 (1998).
- 1998STO/KRZ P. Storoniak, K. Krzyminski, and J. Blazejowski, *J. Therm. Anal.* **54**, 183 (1998).
- 1998STO/NG L. D. Stockton, T. L. Ng, N. Maung, I. B. Poole, J. O. Williams, A. C. Wright, D. R. Foster, and D. J. Cole-Hamilton, *J. Cryst. Growth* **183**, 95 (1998).
- 1998SVO/HYN V. V. Svoboda, Hyněk, and B. Koutek, *J. Chem. Thermodyn.* **30**, 1411 (1998).
- 1998VAN/AUG G. Van den Mooter, P. Augustijns, and R. Kinget, *Int. J. Pharm.* **164**, 81 (1998).
- 1998VAN/KEL P. J. C. M. van Hoof and E. Kellenbach, *J. Therm. Anal. Calorim.* **53**, 957 (1998).
- 1998VAN/VAN J. C. van Miltenburg, A. C. G. van Genderen, and G. J. K. van den Berg, *Thermochim. Acta* **319**, 151 (1998).
- 1998VAR/DRU R. M. Varushchenko and A. I. Druzhinina, *J. Chem. Thermodyn.* **30**, 697 (1998).
- 1998VAS/LEB V. G. Vasil'ev and B. V. Lebedev, *Polym. Sci., Ser. A Ser. B* **40**, 464 (1998).
- 1998VER S. P. Verevkin, *J. Chem. Thermodyn.* **30**, 1029 (1998).
- 1998VER/BEC S. P. Verevkin, H.-D. Beckhaus, U. Schüle, and C. Rüchardt, *Struct. Chem.* **9**, 1 (1998).
- 1998VER/KUM S. P. Verevkin, M. Kümmerlin, H.-D. Beckhaus, C. Galli, and C. Rüchardt, *Eur. J. Org. Chem.*, 579 (1998).
- 1998VER/PEN S. P. Verevkin, W.-H. Peng, H. D. Beckhaus, and C. Rüchardt, *Eur. J. Org. Chem.* 2323 (1998).
- 1998VER/WEL S. P. Verevkin and F. M. Welle, *Struct. Chem.* **9**, 215 (1998).
- 1998VER2 S. P. Verevkin, *J. Chem. Thermodyn.* **30**, 1069 (1998).
- 1998VER3 S. P. Verevkin, *Struct. Chem.* **9**, 375 (1998).
- 1998VER4 S. P. Verevkin, *Thermochim. Acta* **310**, 229 (1998).
- 1998VER5 S. P. Verevkin, *J. Chem. Thermodyn.* **30**, 389 (1998).
- 1998VER6 S. P. Verevkin, *Struct. Chem.* **9**, 113 (1998).
- 1998VER7 S. P. Verevkin, *Thermochim. Acta* **316**, 131 (1998).
- 1998ZEL/MIN L. N. Zelenina, Y. F. Minenkov, Y. G. Stenin, V. A. Titov, and T. P. Chusova, *Zh. Fiz. Khim.* **72**, 803 (1998).
- 1998ZHA/MO X. Zhang, H. Mo, K. Yang, and F. An, *Huanjing Huaxue* **17**, 50 (1998); *Chem. Abstr.* **128**, 318311d (1998). (Note: the value is referred to as a vaporization enthalpy).
- 1998ZHA/TAN J.-B. Zhang, Z.-C. Tan, X.-G. Gao, S.-H. Meng, and L. Li, *Thermochim. Acta* **322**, 89 (1998).
- 1998ZIE/WSZ A. Zielenkiewicz, M. Wszelaka-Rylik, J. Poznanski, and W. Zielenkiewicz, *J. Solution Chem.* **27**, 235 (1998).
- 1999AHL/LOH J. Ahlers, J. Lohmann, and J. Gmehling, *J. Chem. Eng. Data* **44**, 727 (1999).
- 1999ALI/MAL A. S. Alikhanyan, I. P. Malkerova, N. P. Kuz'mina, A. Gleizes, M. Julve-Olsina, J. L. Sanz, and I. L. Eremenko, *Zh. Neorg. Khim.* **44**, 969 (1999); *Chem. Abstr.* **131**, 234185q (1999).
- 1999ANT/FRA M. Antosik, Z. Frasz, and S. K. Malanowski, *J. Chem. Eng. Data* **44**, 368 (1999).
- 1999AUC/LOR A. Aucejo, S. Loras, R. Munoz, and L. M. Ordonez, *Fluid Phase Equilib.* **156**, 173 (1999).
- 1999BAL/WEI J. E. Baldvins and R. G. Weiss, *Liq. Cryst.* **26**, 897 (1999).
- 1999BAU/MAR A. Bauer-Brandl, E. Marti, A. Geoffroy, A. Poso, J. Suurkusk, E. Wappler, and K. H. Bauer, *J. Therm. Anal. Calorim.* **57**, 7 (1999).
- 1999BER/ZEL G. A. Berezovskii, L. M. Zelenina, T. P. Chusova, and I. E. Paukov, *Russ. J. Phys. Chem.* **73**, 1347 (1999).
- 1999BOL/MAR O. V. Boltalina, V. Y. Markov, A. Y. Borschevskii, N. A. Galeva, L. N. Sidorov, G. Gigli, and G. Balducci, *J. Phys. Chem. B* **103**, 3828 (1999).
- 1999BRU/PIA B. Brunetti and V. Piacenti, *J. Chem. Eng. Data* **44**, 809 (1999).
- 1999BRU/VAS B. Brunetti, P. Vassallo, V. Piacenti, and P. Scardala, *J. Chem. Eng. Data* **44**, 509 (1999).
- 1999CHA/GAR D. Chandra, M. L. Garner, and K. H. Lau, *J. Phase Equilib.* **20**, 565 (1999).
- 1999CHI/ACR J. S. Chickos, W. E. Acree, Jr., and J. F. Liebman, *J. Phys. Chem. Ref. Data* **28**, 1535 (1999).
- 1999CHI/KNI R. D. Chirico, S. E. Knipmeyer, A. Nguyen, and W. V. Steele, *J. Chem. Thermodyn.* **31**, 339 (1999).
- 1999CHI/NIC J. Chickos, G. Nichols, J. Wilson, M. Orf, P. Webb, and J. Wang, in *Energetics of Stable Molecules and Reactive Intermediates*, NATO Science Series C, Vol. 535, edited by M. da Piedade (Kluwer Academic, Boston, MA, 1999), pp. 177–202.
- 1999COS/EUS F. S. Costa, M. E. Eusebio, J. S. Redinha, and M. L. P. Leitao, *J. Chem. Thermodyn.* **31**, 895 (1999).
- 1999DAV/FLO J. Z. Davalos, H. Flores, P. Jimenez, R. Notano, M. V. Roux, E. Juaristi, R. S. Hosmane, and J. F. Liebman, *J. Org. Chem.* **64**, 9328 (1999).
- 1999DEF/DEO R. F. de Farias, O. A. de Oliveira, J. V. Medeiros, and C. Airoldi, *Thermochim. Acta* **328**, 241 (1999).
- 1999DEG/GUI E. T. De Givenchy, F. Guittard, F. Bracon, and A. Cambon, *Liq. Cryst.* **26**, 1371 (1999).
- 1999DEL/BAR P. Del Vecchio, B. Barone, R. Sabbah, G. Della Gatta, and L. Abate, *J. Chem. Thermodyn.* **31**, 1001 (1999).
- 1999DIA/GUE M. A. E. Diaz, T. Guetachew, P. Landy, J. Jose, and A. Voilley, *Fluid Phase Equilib.* **109**, 469 (1999).
- 1999DIB/LUS V. N. Dibrivnyi, G. V. Lustiv, Yu. Ya. Van-Chin-Syan, A. P. Yuvchenko, and E. A. Dikusar, *Russ. J. Phys. Chem.* **73**, 2040 (1999).
- 1999DIB/PIS V. N. Dibrivnyi, Z. E. Pistun, Yu. Ya. Van-Chin-Syan, A. P. Yuvchenko, and T. D. Zvereva, *Zh. Fiz. Khim.* **73**, 2257 (1999).
- 1999DIB/PIS2 V. N. Dibrivnyi, Z. E. Pistun, Yu. Ya. Van-Chin-Syan, A. Yuvchenko, and T. D. Zvereva, *Russ. J. Phys. Chem.* **73**, 2043 (1999).
- 1999DOL/LEC G. Dollo, P. Le Corre, M. Chollet, F. Chevanne, M. Bertault, J.-L. Burgot, and R. Le Verge, *J. Pharm. Sci.* **88**, 889 (1999).
- 1999DOM/KAR L. G. Domracheva, N. V. Karyakin, M. S. Sheiman, G. V. Kamelova, V. N. Larina, O. N. Suvorova, and G. A. Domrachev, *Russ. Chem. Bull.* **48**, 1647 (1999).
- 1999DOU/BOT A. G. Douglass, B. Both, and P. Kaszynski, *J. Mater. Chem.* **9**, 683 (1999).
- 1999DRU/GAL A. I. Druzhinina, N. A. Galeva, R. M. Varushchenko, and

- O. V. Boltalina, *J. Chem. Thermodyn.* **31**, 1469 (1999).
- 1999DRU/VAR A. I. Druzhinina and R. M. Varushchenko, *Russ. J. Phys. Chem.* **73**, 1367 (1999).
- 1999DYK/SVO J. Dykyj, J. Svoboda, R. C. Wilhoit, M. L. Frenkel, and K. R. Hall, *Vapor Pressure of Chemicals: Part A. Vapor Pressure and Antoine Constants for Hydrocarbons and Sulfur, Selenium, Tellurium and Hydrogen Containing Organic Compounds* (Springer, Berlin, 1999). The vaporization enthalpies were calculated from the vapor pressures obtained from the Antoine constants reported in this compendium. In cases where the Antoine constant $C=0$, the Antoine Equation ($\log_{10} P=A-B/(C+T)$) reduces to the integrated form of the Clausius Clapeyron equation directly. This was the case for many vaporization enthalpies. In those cases where this condition was not met the vaporization enthalpy was calculated as $\Delta_{\text{vap}}H_m(T)=2.303RB[T/(T+C)]^2$.
- 1999EMM/PIC F. Emmenegger and M. Piccand, *J. Therm. Anal. Calorim.* **57**, 235 (1999).
- 1999ESC/SAN G. N. Escobedo-Alvarado and S. I. Sandler, *J. Chem. Eng. Data* **44**, 319 (1999).
- 1999FAT D. Fatu, *J. Therm. Anal. Calorim.* **56**, 739 (1999).
- 1999GAL/COL Z. Galewski and H. J. Coles, *J. Mol. Liq.* **79**, 77 (1999).
- 1999GAR/AND R. Garriga, A. C. Andres, P. Perez, and M. Gracia, *J. Chem. Eng. Data* **44**, 296 (1999).
- 1999GIO/BET F. Giordano, R. Bettini, C. Donini, A. Gazzaniga, M. R. Caira, G. G. Z. Zhang, and D. J. W. Grant, *J. Pharm. Sci.* **88**, 1210 (1999).
- 1999GIR/PIE D. Giron, P. Piechon, C. Goldbronn, and S. Pfeffer, *J. Therm. Anal. Calorim.* **57**, 61 (1999).
- 1999GOT/BUH A. Gotze, C. Buhmester, I. Svoboda, and H. Fuess, *Croatica Chim. Acta* **72**, 443 (1999).
- 1999GRI/AUE U. J. Griesser, M. E. Auer, and A. Burger, *Sci. Pharm.* **67**, 319 (1999).
- 1999GRI/SZE U. J. Griesser, M. Szelagiewicz, U. Ch. Hofmeier, C. Pitt, and S. Ciaferani, *J. Therm. Anal.* **57**, 45 (1999).
- 1999HAM/WURF. Hamann and A. Wurflinger, *Z. Phys. Chem.* **211**, 85 (1999).
- 1999HEI/FIS A. Heine, K. Fischer, and J. Gmehling, *J. Chem. Eng. Data* **44**, 373 (1999).
- 1999HEN/KUH J.-O. Henck and M. Kuhnert-Brandstaetter, *J. Pharm. Sci.* **88**, 103 (1999).
- 1999HUD/SHE C. M. Hudson, R. A. Shenoy, M. E. Neubert, and R. G. Petschek, *Liq. Cryst.* **26**, 241 (1999).
- 1999HUZ/SAI S. Huzisawa, K. Saito, and I. Ikemoto, *J. Phys. Chem. Solids* **60**, 723 (1999).
- 1999JIM/ROU P. Jimenez, M. V. Roux, J. Z. Davalos, J. L. M. Abboud, and M. T. Molina, *J. Chem. Thermodyn.* **31**, 263 (1999).
- 1999JON/FEN D. E. G. Jones, H. T. Feng, R. A. Austen, and R. C. Fouchard, *J. Therm. Anal. Calorim.* **55**, 9 (1999); *Chem. Abstr.* **132**, 182729u (2000).
- 1999KAB/KOZ G. J. Kabo, A. A. Kozyro, M. Frenkel, and A. V. Blokhin, *Mol. Cryst. Liq. Cryst.* **326**, 333 (1999).
- 1999KEY/BOT A. Keys, S. G. Bott, and A. R. Barron, *Chem. Mater.* **11**, 3578 (1999).
- 1999KIM/HIR K. Kimura, F. Hirayama, and K. Uekama, *J. Pharm. Sci.* **88**, 385 (1999).
- 1999KOB/OGU K. Kobashi and M. Oguni, *J. Phys. Chem. B* **103**, 7687 (1999).
- 1999KOL/DOR V. P. Kolesov, O. V. Dorofeeva, V. S. Iorish, T. S. Papina, V. A. Lukyanova, and S. V. Melkhanova, *Mendeleev Commun.* **9**, 143 (1999).
- 1999KOR/LEV A. P. Korobko, I. V. Levakova, S. V. Krashennnikov, A. I. Stash, N. A. Kon'kova, V. V. Kuz'min, N. V. Kozlova, and T. A. Korobko, *Kristallografiya* **44**, 262 (1999).
- 1999KUL/LEB T. G. Kulagina and B. V. Lebedev, *Russ. J. Phys. Chem.* **73**, 1909 (1999).
- 1999LEB/KUL B. V. Lebedev, T. G. Kulagina, and E. G. Kiparisova, *Russ. J. Phys. Chem.* **73**, 521 (1999).
- 1999LEB/KUL2 B. V. Lebedev, T. G. Kulagina, and D. R. Tur, *J. Chem. Thermodyn.* **31**, 697 (1999).
- 1999LEI/WAN Y. D. Lei, F. Wania, and W. Y. Shiu, *J. Chem. Eng. Data* **44**, 577 (1999).
- 1999LEI/WAN2 Y. D. Lei, F. Wania, W. Y. Shiu, and D. G. B. Boocock, *J. Chem. Eng. Data* **44**, 200 (1999).
- 1999LI/ZEL Z. J. Li, M. T. Zell, E. J. Munson, and D. J. W. Grant, *J. Pharm. Sci.* **88**, 337 (1999).
- 1999LIM/PAR J. S. Lim, Y.-Y. Park, B.-G. Lee, Y.-W. Lee, and J.-D. Kim, *J. Chem. Eng. Data* **44**, 1226 (1999).
- 1999LIN/KO H. C. Lin, C.-W. Ko, K. Guo, and T. W. Cheng, *Liq. Cryst.* **26**, 613 (1999).
- 1999LOR/AUC S. Loras, A. Aucejo, and R. Munoz, *Fluid Phase Equilib.* **156**, 185 (1999).
- 1999MAR/BAS G. Marchionni, M. Bassi, G. Fontana, P. Maccone, and G. Ajroldi, *J. Fluorine Chem.* **41**, 98 (1999).
- 1999MAT/PEP Y. N. Matyushin, V. I. Pepekin, V. P. Lebedev, V. V. Chironov, L. M. Kostikova, Y. O. Inozemtcev, T. S. Pivina, and A. B. Sheremetev, International Annual Conference of ICT, 1999 (unpublished), Vol. 30, p. 77/1.
- 1999MAY/WIT J. Mayer, W. Witko, M. Massalska-Arodz, G. Williams, and R. Dabrowski, *Phase Transitions* **69**, 199 (1999).
- 1999MEN/LIA S. H. Meng, P. Liang, Z. C. Tan, Y. J. Song, L. Li, and L. Wang, *Thermochim. Acta* **342**, 47 (1999).
- 1999MIC/NEG F. Michaud, Ph. Negrier, D. Mikailitchenko, A. Marbeuf, Y. Haget, M. Cuevas-Diarte, and H. A. J. Oonk, *Mol. Cryst. Liq. Cryst.* **326**, 409 (1999).
- 1999MIR/VOR E. A. Miroshnichenko and V. P. Vorob'eva, *Russ. J. Phys. Chem.* **73**, 349 (1999).
- 1999MIR/VOR2 E. A. Miroshnichenko and V. P. Vorob'eva, *Zh. Fiz. Khim.* **73**, 419 (1999).
- 1999MO/YAN O. Mo, M. Yanez, M. V. Roux, P. Jimenez, J. Z. Davalos, M. A. V. Ribeiro da Silva, M. D. M. C. Ribeiro da Silva, M. A. R. Matos, L. M. P. F. Amaral, A. Sanchez-Migallon, P. Cabildo, R. Claramunt, J. Elguero, and J. F. Liebman, *J. Phys. Chem. A* **103**, 9336 (1999).
- 1999MOK/PRC L. Mokbel, S. Pcedda, T. Guetachew, B. Marongiu, and J. Jose, *ELDATA: Int. Electron. J. Phys. Chem. Data* **5**, 79 (1999).
- 1999MON/DEL J. B. Monton, J. de la Torre, M. C. Burguet, R. Munoz, and S. J. Loras, *J. Chem. Eng. Data* **44**, 1158 (1999).
- 1999MON/HIL M. J. S. Monte and D. M. Hillesheim, *J. Chem. Thermodyn.* **31**, 1433 (1999).
- 1999MUR/FAU P. Mura, M. T. Faucci, P. L. Parrini, S. Furlanetto, and S. Penzauti, *Int. J. Pharm.* **179**, 117 (1999).
- 1999NGU/BER J. N'Guimbi, C. Berro, I. Mokbel, E. Rauzy, and J. Jose, *Fluid Phase Equilib.* **162**, 143 (1999).
- 1999NIS/SHE V. P. Nistratov, M. S. Sheiman, and G. P. Kamelova, *Russ. J. Phys. Chem.* **73**, 842 (1999).
- 1999NVEV/GOU L. P. M. Neves, A. Gouveia de Souza, J. B. L. de Oliveira, and C. Airolidi, *Thermochim. Acta* **328**, 217 (1999).
- 1999OGA/NAK Y. Ogawa and N. Nakamura, *Bull. Chem. Soc. Jpn.* **72**, 943 (1999).
- 1999OJA/SUU V. Oja and E. M. Suuberg, *J. Chem. Eng. Data* **44**, 26 (1999).
- 1999ORT/HER J. Ortega and P. Hernandez, *J. Chem. Eng. Data* **44**, 757 (1999).
- 1999PRI/BAS D. M. Price, S. Bashir, and P. R. Derrick, *Thermochim. Acta* **327**, 167 (1999).
- 1999PRI/HAW D. M. Price and M. Hawkins, *Thermochim. Acta* **329**, 73 (1999).
- 1999RAI/RAI U. S. Rai and R. N. Rai, *J. Mater. Res.* **14**, 1299 (1999).
- 1999RIB/FER M. D. M. C. Ribeiro da Silva, M. L. c. C. H. Ferrao, M. J. S. Monte, J. M. Goncalves, and F. Jiye, *J. Chem. Thermodyn.* **31**, 1067 (1999).
- 1999RIB/MAT M. A. V. Ribeiro da Silva, M. A. R. Matos, M. J. S. Monte, D. M. Hillesheim, M. C. P. O. Marques, and N. F. T. G. Vieira, *J. Chem. Thermodyn.* **31**, 1429 (1999).
- 1999RIB/MAT2 M. A. V. Ribeiro da Silva, M. A. R. Matos, V. M. F. Moraes, and M. S. Miranda, *J. Org. Chem.* **64**, 8816 (1999).
- 1999RIB/MON M. A. V. Ribeiro da Silva, M. J. S. Monte, and J. R. Ribeiro, *J. Chem. Thermodyn.* **31**, 1093 (1999).
- 1999RIB/RIB M. A. V. Ribeiro da Silva, M. D. M. C. Ribeiro da Silva, M. A. R. Matos, P. Jimenez, M. V. Roux, J. Elguero, R. Claramunt, P. Cabildo, and A. Sanchez-Migallon, *J. Chem.*

- Thermodyn.* **31**, 129 (1999).
- 1999ROH/MUS V. Rohac, J. E. Musgrove, K. Ruzicka, V. Ruzicka, M. Zabransky, and K. Aim, *J. Chem. Thermodyn.* **31**, 971 (1999).
- 1999ROH/RUZ V. Rohac, V. Ruzicka, K. Ruzicka, M. Polednicek, K. Aim, J. Jose, and M. Zabransky, *Fluid Phase Equilib.* **157**, 121 (1999).
- 1999ROM/ESC S. Romero, B. Escalera, and P. Bustamante, *Int. J. Pharm.* **178**, 193 (1999).
- 1999ROS/MOL F. Ros and M. T. Molina, *Eur. J. Org. Chem.* 3179 (1999).
- 1999ROU/DAV M. V. Roux, J. Z. Davalos, P. Jimenez, H. Flores, J. L. Saiz, J. L. M. Abboud, and E. Juaristi, *J. Chem. Thermodyn.* **31**, 635 (1999).
- 1999ROU/JIM M. V. Roux, P. Jimenez, J. Z. Davalos, R. Notaro, and J.-L. M. Abboud, *J. Chem. Thermodyn.* **31**, 1457 (1999).
- 1999SAB/IDE R. Sabbah and S. Ider, *Can. J. Chem.* **77**, 249 (1999).
- 1999SAB/PER R. Sabbah and L. Perez, *Aust. J. Chem.* **52**, 235 (1999).
- 1999SAB/PER2 R. Sabbah and L. Perez, *Can. J. Chem.* **77**, 1508 (1999).
- 1999SAB/XU R. Sabbah, A. Xu-Wu, J. S. Chickos, M. L. Plana Leitao, M. V. Roux, and L. A. Torres, *Thermochim. Acta* **331**, 93 (1999).
- 1999SAL/LOP J. Salud, D. O. Lopez, M. Barrio, and J. I. Tamarit, *J. Mater. Chem.* **9**, 909 (1999).
- 1999SAN/PET L. S. Santos, Jr., G. P. Petrucelli, and C. Airoidi, *Polyhedron* **18**, 969 (1999); *Chem. Abstr.* **131**, 50237 (1999).
- 1999SHE/KAM M. S. Sheiman, G. P. Kamelova, I. A. Zelyaev, and V. M. Sheiman, *Russ. J. Phys. Chem.* **73**, 512 (1999).
- 1999SUG S. Suga, *Thermochim. Acta* **328**, 9 (1999).
- 1999TAF/GUI E. Taffin de Givenchy, F. Guittard, F. Bracon, and A. Cambon, *Liq. Cryst.* **26**, 1163 (1999).
- 1999TAH/TAK Y. Tahira and Y. Taketani, Patent, *Chem. Abstracts* **130**, 124888 (1999).
- 1999TIT/ZEL V. A. Titov, L. N. Zelenina, and T. P. Chusova, *Zh. Fiz. Khim.* **73**, 951 (1999).
- 1999TIT/ZEL2 V. A. Titov, L. N. Zelenina, and T. P. Chusova, *Russ. J. Phys. Chem.* **73**, 835 (1999).
- 1999TOC/AKI K. Tochigi, K. Akimoto, K. Ochi, F. Liu, and Y. Kawase, *J. Chem. Eng. Data* **44**, 588 (1999).
- 1999VAN/TEN J. C. van Miltenburg and E. ten Grotenhuis, *J. Chem. Eng. Data* **44**, 721 (1999).
- 1999VER S. P. Verevkin, *J. Chem. Thermodyn.* **31**, 1397 (1999).
- 1999VER/EBE S. P. Verevkin and J. Ebenhoch, *Struct. Chem.* **10**, 401 (1999).
- 1999VER/HEI S. P. Verevkin and A. Heintz, *J. Chem. Eng. Data* **44**, 1240 (1999).
- 1999VER2 S. P. Verevkin, *J. Chem. Thermodyn.* **31**, 559 (1999).
- 1999VER3 S. P. Verevkin, *J. Chem. Eng. Data* **44**, 557 (1999).
- 1999VER4 S. P. Verevkin, *J. Chem. Eng. Data* **44**, 1245 (1999).
- 1999VER5 S. P. Verevkin, *J. Chem. Eng. Data* **44**, 175 (1999).
- 1999VER6 S. P. Verevkin, *Thermochim. Acta* **326**, 17 (1999).
- 1999VER7 S. P. Verevkin, *Phys. Chem. Chem. Phys.* **1**, 127 (1999).
- 1999VER8 S. P. Verevkin, *Thermochim. Acta* **332**, 27 (1999).
- 1999WAN/TAN L. Wang, Z.-C. Tan, S.-H. Meng, and D.-B. Liang, *Thermochim. Acta* **342**, 59 (1999).
- 1999WEB F. Weber, *PTB-Mitteilungen* **109**, 469 (1999).
- 1999WEL/DRU D. Wells and C. J. Drummond, *Langmuir* **15**, 4713 (1999).
- 1999WIN G. Winter, in *Reactivity of Molecular Solids*, edited by E. Boldyreva and V. Boldyrev (Wiley and Sons, NY, 1999) Chap. 7.
- 1999XUE/TAN B. Xue, Z.-C. Tan, S.-W. Lu, S.-H. Meng, and X.-H. Yuan, *Huaxue Xuebao* **57**, 881 (1999); *Chem. Abstr.* **131**, 224854 (1999).
- 1999ZAD/KER D. Zadravec, J. Kerc, and J. Kristl, *Farm. Vest. [Ljubljana]* **50**, 350 (1999); *Chem. Abstr.* **132**, 313492.
- 1999ZAR/CHA M. I. Zaretskii, E. M. Chartov, L. A. Pushkina, and V. V. Elkin, *Russ. J. Phys. Chem.* **72**, 1702 (1999).
- 1999ZEM/STA S. M. Zemskova, P. A. Stabnikov, S. V. Susoev, and I. K. Igumenov, *Proc.-Electrochem. Soc.* **98**, 286 (1999).
- 1999ZHE/KAT V. P. Zhelezny, Y. A. Katchurka, and M. V. Pybnikov, *High Temp. - High Press.* **31**, 169 (1999).
- 1999ZIE/GOL W. Zielenkiewicz, B. Golankiewicz, G. L. Pelovich, and M. Kozbial, *J. Solution Chem.* **28**, 731 (1999).
- 1999ZIE/PER W. Zielenkiewicz, G. L. Perlovich, and M. Wszelaka-Rylik, *J. Therm Anal. Calorim.* **57**, 225 (1999).
- 2000AGU/GUA A. R. Aguilar and E. O. Guareno, *J. Chem. Thermodyn.* **32**, 767 (2000).
- 2000AKU/IUC T. Akutagawa, K. Iuchi, and Y. Matsunaga, *Liq. Cryst.* **27**, 1399 (2000).
- 2000ATA/KAW T. Atake, H. Kawaji, T. Tojo, K. Kawasaki, Y. Ootsuka, M. Katou, and Y. Koga, *Bull. Chem. Soc. Jpn.* **73**, 1987 (2000).
- 2000BEC/FAU H.-D. Beckhaus, R. Faust, D. L. Matzger, D. L. Mohler, D. W. Rogers, C. Rhardt, A. K. Sawhney, S. p. Verevkin, K. P. C. Volhardt, and S. Wolff, *J. Am. Chem. Soc.* **122**, 7819 (2000).
- 2000BEL/BEL B. F. Belaribi, G. Belaribi-Boukais, A. A. Kaci, and J. Jose, *J. Therm Anal. Calorim.* **61**, 787 (2000).
- 2000BLA/LUC A. E. Blatch and G. R. Luckhurst, *Liq. Cryst.* **27**, 775 (2000).
- 2000BOB/CAM S. Bobbo, R. Camporese, and R. Stryjek, *J. Chem. Thermodyn.* **32**, 1647 (2000).
- 2000BOL/GAL O. V. Boltalina and N. A. Galeva, *Russ. Chem. Rev.* **69**, 609 (2000).
- 2000BOU/YE E. Bourret-Courchesne, Q. Ye, D. W. Peters, J. Arnold, M. Ahmed, S. J. C. Irvine, R. Kanjolia, L. M. Smith, and S. A. Rushworth, *J. Cryst. Growth* **217**, 47 (2000).
- 2000BRO/DU S. Brownridge, H. Du, S. A. Fairhurst, R. C. Haddon, H. Oberhammer, S. Parson, J. Passmore, M. J. Schriver, L. H. Sutcliffe, and N. P. C. Westwood, *J. Chem. Soc. Dalton Trans.* 3365 (2000).
- 2000BRU/DEL B. Brunetti, G. Della Gatta, and V. Piacente, *J. Chem. Eng. Data* **45**, 237 (2000).
- 2000BRU/PIA B. Brunetti, V. Piacente, and G. Portalone, *J. Chem. Eng. Data* **45**, 242 (2000).
- 2000BUR/LET A. Burger and A. Lettenbichler, *Eur. J. Pharm. Biopharm.* **49**, 65 (2000).
- 2000BUR/VAN A. Burger and C. Van den Boom, *Mikrochim. Acta* **135**, 63 (2000).
- 2000BYK/LEB T. A. Bykova, B. V. Lebedev, N. V. Ushakov, and E. Sh. Finkel'shtein, *Polymer Sci.* **42**, 863 (2000). (Note: the compound used was only 83% crystalline. The authors reported a corrected value of 7.77 kJ mol⁻¹ for the enthalpy of fusion for a sample of 100 % degree of crystallinity.)
- 2000CHA/SOS C. Chaimbault, J. J. Sosc, J. M. Leger, P. Negrier, F. Capelle, and C. Jarry, *J. Pharm. Sci.* **89**, 1496 (2000).
- 2000DAN/PRO G. H. Danila, L. Profire, G. G. Bumbu, and C. Vasile, *Thermochim. Acta* **343**, 69 (2000).
- 2000DEA/SOU J. A. de A. Imeida Sales, A. G. Souza, G. F. Goncalves de Freitas, S. Prasad, M. F. S. Trindade, L. H. Carvalho, and P. O. Dunstan, *Thermochim. Acta* **356**, 9 (2000).
- 2000DEF/VAN K. M. De Fina, T. T. Van, K. A. Fletcher, and W. E. Acree, Jr., *Can. J. Chem.* **78**, 449 (2000).
- 2000DEL/JOZ G. Della Gatta, M. Jozwiak, B. Brunetti, and L. Abate, *J. Chem. Thermodyn.* **32**, 979 (2000).
- 2000DI/LI Y.-Y. Di, S. Li, S.-H. Meng, Z.-C. Tan, and S.-S. Qu, *Huaxue Xuebao* **58**, 1380 (2000); *Chem. Abstr.* **134**, 10187r (2001).
- 2000DI/TAN Y. Y. Di, Z.-C. Tan, X.-M. Xu, S.-H. Meng, and S.-S. Qu, *Thermochim. Acta* **356**, 143 (2000).
- 2000DI/TAN2 Y.-Y. Di, Z.-C. Tan, Z.-N. Li, Y.-J. Song, Y. Zheng, S.-H. Meng, and S.-S. Qu, *Thermochim. Acta* **362**, 7 (2000).
- 2000DRU/VAR A. I. Druzhinina, R. M. Varushchenko, V. S. Sarisova, and A. A. Pimerzin, *Zh. Fiz. Khim.* **74**, 404 (2000).
- 2000DRU/VAR2 A. I. Druzhinina, R. M. Varushchenko, V. S. Sarkisova, and A. A. Pimerzin, *Russ. J. Phys. Chem.* **74**, 333 (2000).
- 2000DUN P. O. Dunstan, *Thermochim. Acta* **356**, 19 (2000).
- 2000EME/NIK A. L. Emelina, M. I. Nikitin, A. S. Alikhanyan, and V. F. Sukhoverkhov, *Russ. J. Inorg. Chem.* **45**, 1565 (2000).
- 2000FAH/BAR B. D. Fahlman and A. R. Barron, *Adv. Mater. Opt. Electron.* **10**, 223 (2000).
- 2000GAN/BOG R. B. Gandhi, J. B. Bogardus, D. E. Bugay, R. K. Perrone, and M. A. Kaplan, *Int. J. Pharm.* **201**, 221 (2000).
- 2000GIE E. Giera, *J. Chem. Thermodyn.* **32**, 821 (2000).
- 2000GRI/AUE U. J. Griesser, M. E. Auer, and A. Burger, *Microchem. J.*

- 65, 283 (2000).
- 2000HAN/BOT E. Handelsman-Benory, M. Botoshansky, M. Greenberg, V. Shteiman, and M. Kafory, *Tetrahedron* **56**, 6887 (2000).
- 2000HAN/PAR B. C. Hancock and M. Parks, *Pharm. Res.* **17**, 397 (2000).
- 2000IHN/VEN P. M. Ihnat, J. L. Vennerstrom, and D. H. Robinson, *J. Pharm. Sci.* **89**, 1525 (2000).
- 2000JAR/MAR S. Jarmelo, T. M. R. Maria, M. L. P. Leitao, and R. Fausto, *Phys. Chem. Chem. Phys.* **2**, 1155 (2000).
- 2000KAB/BLO G. J. Kabo, A. V. Blokhin, M. B. Charapennikau, A. G. Kabo, and V. M. Sevruck, *Thermochim. Acta* **345**, 125 (2000).
- 2000KAM/YOS M. Kamigauchi, M. Yoshida, K. Saiki, M. Sugiura, J. Nishijo, Y. In, and T. Ishida, *Bull. Chem. Soc. Jpn.* **73**, 1233 (2000).
- 2000KAN/SAM S. K. Kang and E. T. Samulski, *Liq. Cryst.* **27**, 371 (2000).
- 2000KIR D. Kirklın, *J. Chem. Thermodyn.* **32**, 701 (2000).
- 2000KOR/DOR M. V. Korobov, P. A. Dorozhko, A. S. Lobach, A. A. Popov, and V. M. Senyavin, *Schr. Forschungszent. Juelich Energietechn./Energy Technol.* **15**, 371 (2000).
- 2000KOZ/MAK A. A. Kozyro, Yu. V. Maksimuk, and G. Ya. Kabo, *Zh. Prikl. Khim. (S.-Peterburg)* **73**, 199 (2000).
- 2000LAR/LER L. Larachi, M. Leroux, S. Hamoudi, A. Bernis, and A. Sayari, *J. Chem. Eng. Data* **45**, 404 (2000).
- 2000LEB/SMI B. V. Lebedev, N. N. Smirnova, V. S. Papkov, M. I. Buzin, and I. I. Dubovik, *Polym. Sci., Ser. A Ser. B* **42**, 1111 (2000).
- 2000LIA/MA Y.-H. Liang and P.-S. Ma, *Shiyou Huagong* **29**, 939 (2000).
- 2000LIS/JAM Z. Lisicki and M. E. Jamroz, *J. Chem. Thermodyn.* **32**, 1335 (2000).
- 2000MAC/COU J. J. B. Machado, J. A. Coutinho, and E. A. Macedo, *Fluid Phase Equilib.* **173**, 121 (2000).
- 2000MAH/SOL R. Mahmoud, R. Solimando, M. Bouroukba, and M. Rogalski, *J. Chem. Eng. Data* **45**, 433 (2000).
- 2000MAR/BOL V. Yu. Markov, O. V. Boltalina, A. A. Gorjunktov, A. Y. Lukonin, L. N. Sidorov, G. Gigli, G. Balducci, and R. Taylor, *Proc.-Electrochem. Soc.* **2000-12**, 109 (2000); *Chem. Abstr.* **134**, 121607x (2001).
- 2000MAT/MIR M. A. R. Matos, M. S. Miranda, and V. M. F. Morais, *J. Phys. Chem. A* **104**, 9260 (2000).
- 2000MEL/PIM S. V. Melkhanova, S. M. Pimenova, V. P. Kolesov, A. A. Pimerzin, and V. S. Sarkisova, *J. Chem. Thermodyn.* **32**, 1311 (2000).
- 2000MOK/RUZ I. Mokbel, K. Ruzicka, V. V. Majer, Ruzicka, M. Ribeiro, J. Jose, and M. Zabransky, *Fluid Phase Equilib.* **169**, 191 (2000).
- 2000MON/HIL M. J. S. Monte and D. M. Hillesheim, *J. Chem. Thermodyn.* **32**, 1727 (2000).
- 2000MON/HIL2 M. J. S. Monte and D. M. Hillesheim, *J. Chem. Eng. Data* **45**, 1088 (2000).
- 2000MOR/HAR K. Moriya, F. Harada, S. Yano, and S. Kagabu, *Liq. Cryst.* **27**, 1647 (2000).
- 2000MOR/SEM N. B. Morozova, P. P. Semyannikov, S. V. Sysoev, V. M. Grankin, and I. K. Igumenov, *J. Therm Anal. Calorim.* **60**, 489 (2000).
- 2000NAK/SHI N. Nakasone, K. Shiokawa, Y. Urabe, and N. Nemoto, *J. Phys. Chem. B* **104**, 7843 (2000).
- 2000NIC/ORF G. Nichols, J. Orf, S. M. Reiter, J. Chickos, and G. W. Gokel, *Thermochim. Acta* **346**, 15 (2000).
- 2000OHT/CIC T. Ohta, F. Cicoira, P. Doppelt, L. Beitone, and P. Hoffman, *Chem. Vap. Deposition* **7**, 33 (2000); *Chem. Abstr.* **134**, 153247w (2001).
- 2000OON/VAN H. A. J. Oonk, A. C. G. van Genderen, J. G. Blok, and P. R. van der Linde, *Phys. Chem. Chem. Phys.* **2**, 5614 (2000).
- 2000OVA/KOU P. V. Ova, B. Koultek, and M. Hoskovec, in *Practice Oriented Results on Use and Production of Neem Ingredients and Pheromones VI*, edited by H. Kleeberg and C. P. W. Zebitz (Druck and Graphic, Giessen, 2000), pp. 211–218.
- 2000PAL/SZA M. Palczewska-Tulinska and A. M. Szafranski, *J. Chem. Eng. Data* **45**, 988 (2000).
- 2000PAP/KOL T. S. Papaina, V. P. Kolesova, V. A. Lukyanova, O. V. Boltalina, A. Y. Lukonin, and L. N. Sidorov, *J. Phys. Chem. B* **104**, 5403 (2000).
- 2000PAU/MEH I. Paunovic and A. K. Mehrotra, *Thermochim. Acta* **356**, 27 (2000).
- 2000PER/GOL G. L. Perlovich, O. A. Golubchikov, and M. E. Klueva, *J. Porphyr. Phthalocyanines* **4**, 699 (2000).
- 2000PUN P. Panchaipetch, Ph.D. Dissertation, University of North Texas, 2000.
- 2000REI/CAR R. Reich, M. Cartes, H. Segura, and J. Wisniak, *Phys. Chem. Liq.* **38**, 217 (2000).
- 2000RES/GON J. M. Resa, C. Gonzalez, S. Ortiz de Landaluce, and J. Lanz, *J. Chem. Eng. Data* **45**, 867 (2000).
- 2000RIB/GON M. D. M. C. Ribeiro da Silva, J. M. Goncalves, and W. E. Acree, Jr., *J. Chem. Thermodyn.* **32**, 1071 (2000).
- 2000RIB/MAT M. A. V. Ribeiro da Silva, M. A. R. Matos, C. M. A. Rio, M. S. Miranda, and V. M. F. Morais, *J. Phys. Chem. A* **104**, 6644 (2000).
- 2000RIB/MAT2 M. A. V. Ribeiro da Silva, M. A. R. Matos, C. A. Rio, V. M. F. Morais, J. Wang, G. Nichols, and J. S. Chickos, *J. Phys. Chem. A* **104**, 1774 (2000).
- 2000RIB/MON M. A. V. Ribeiro da Silva, M. J. S. Monte, and J. R. Ribeiro, *J. Chem. Eng. Data* **45**, 756 (2000).
- 2000RIB/RIB M. A. V. Ribeiro da Silva, M. D. M. C. Ribeiro da Silva, L. C. M. Silva, F. Dietz, and E. Hoyer, *J. Chem. Thermodyn.* **32**, 1113 (2000).
- 2000RIB/RIB2 M. A. V. Ribeiro da Silva, M. D. M. C. Ribeiro da Silva, M. A. R. Matos, P. Jimenez, M. V. Roux, M. A. Martin-Luengo, J. Elguero, R. Claramut, and P. Cabildo, *J. Chem. Thermodyn.* **32**, 237 (2000).
- 2000ROD/ART S. Rodriguez, H. Artigas, C. Lafuente, A. M. Mainar, and F. M. Royo, *Thermochim. Acta* **362**, 153 (2000).
- 2000ROD/VEC F. Rodante, S. Vecchio, G. Catalani, and M. Guidotti, *J. Therm Anal. Calorim.* **60**, 605 (2000).
- 2000ROH/CEN V. Rohac, M. Censky, D. Zala, V. Ruzicka, K. Ruzicka, K. Sporcka, and K. Aim, *J. Chem. Eng. Data* **45**, 1205 (2000).
- 2000ROU/JIM M. A. Roux, P. Jimenez, J. Z. Davalos, M. A. Martin-Luengo, V. M. Rotello, A. O. Cuello, and J. F. Liebman, *Struct. Chem.* **11**, 1 (2000).
- 2000SCH/MAT E. Schonherr, K. Matsumoto, and K. Murakami, *Proc.-Electrochem. Soc.* **2000-12**, 89 (2000); *Chem. Abstr.* **134**, 121606w (2001).
- 2000SEM/BAS P. P. Semyannikov, T. V. Basova, V. M. Grankin, and I. K. Igumenov, *J. Porphyr. Phthalocyanines* **4**, 271 (2000).
- 2000SIR/HER E. B. Sirota, A. B. Herhold, and M. Varma-Nair, *J. Chem. Phys.* **113**, 8225 (2000).
- 2000SOU/OLI A. G. Souza, M. M. Oliveira, A. D. Gondim, P. O. Dunstan, and D. M. de A. Melo, *Thermochim. Acta* **344**, 29 (2000).
- 2000SZT/BAE B. Sztaray and T. Baer, *J. Am. Chem. Soc.* **122**, 9219 (2000).
- 2000TAM/LOP J. Ll. Tamarit, D. O. Lopez, X. Alcobe, M. Barrio, J. Salud, and L. C. Pardo, *Chem. Mater.* **12**, 555 (2000).
- 2000TOZ/AKU Y. Tozuka, H. Akutsu, Y. Yamamura, K. Saito, and M. Sorai, *Bull. Chem. Soc. Jpn.* **73**, 2279 (2000).
- 2000UNO/IITO H. Uno, S. Ito, M. Wada, H. Watanabe, M. Nagai, A. Hayashi, T. Murashima, and N. Uno, *J. Chem. Soc., Perkin Trans. 1* **1**, 4347 (2000).
- 2000URY/KUP V. F. Ur'yash, V. F. Kupriyanov, N. Y. Kokurina, A. G. Smirnov, and T. M. Kuleshova, *Russ. J. Gen. Chem.* **70**, 719 (2000).
- 2000VAN/OON J. C. van Miltenburg, H. A. J. Oonk, and G. J. K. van den Berg, *J. Chem. Eng. Data* **45**, 704 (2000).
- 2000VER S. P. Verevkin, *J. Chem. Thermodyn.* **32**, 207 (2000).
- 2000VER/HEI S. P. Verevkin and A. Heintz, *J. Chem. Thermodyn.* **32**, 1169 (2000).
- 2000VER/SCH S. P. Verevkin and C. Schick, *J. Chem. Eng. Data* **45**, 946 (2000).
- 2000VER/WAN S. P. Verevkin, D. Wandschneider, and A. Heintz, *J. Chem. Eng. Data* **45**, 618 (2000).
- 2000VER2 S. P. Verevkin, *J. Chem. Eng. Data* **45**, 953 (2000).
- 2000VER3 S. P. Verevkin, *J. Chem. Thermodyn.* **32**, 247 (2000).

- 2000VIL/BRU A. R. Villani, B. Brunetti, and V. Piacente, *J. Chem. Eng. Data* **45**, 823 (2000).
- 2000VIL/BRU2 A. R. Villani, B. Brunetti, and V. Piacente, *J. Chem. Eng. Data* **45**, 1167 (2000).
- 2000WAN/WAN L.-S. Wang and X.-L. Wang, *J. Chem. Eng. Data* **45**, 743 (2000).
- 2000WEN/BAU R. M. Wenslow, M. W. Baum, R. G. Ball, J. M. McCauley, and R. J. Varsolona, *J. Pharm. Sci.* **89**, 1271 (2000).
- 2000WOR/FEN C. J. Wormald and D. P. Fennell, *Int. J. Thermodyn.* **21**, 767 (2000).
- 2000WOR/JAM C. J. Wormald and G. F. James, *J. Chem. Eng. Data* **45**, 348 (2000).
- 2000WOR/VIN C. J. Wormald and M. D. Vine, *J. Chem. Thermodyn.* **32**, 329 (2000).
- 2000WOR/VIN2 C. J. Wormald and M. D. Vine, *J. Chem. Thermodyn.* **32**, 659 (2000).
- 2000WU/TAN X.-M. Wu, Z.-C. Tan, Y. Ji-Song, S.-H. Meng, and S.-S. Qu, *Thermochim. Acta* **346**, 57 (2000).
- 2000WU/WAN L.-H. Wu, Y.-C. Wang, and C.-S. Hsu, *Liq. Cryst.* **27**, 1503 (2000).
- 2000YAM/TAN M. Yamazaki, M. Tanaka, T. Inoue, Y. Suzuki, Y. Nibu, H. Shimada, and R. Shimadu, *Bull. Chem. Soc. Jpn.* **73**, 837 (2000).
- 2000YOU/DOL P. H. Young, D. Dollimore, and C. A. Schall, *J. Therm Anal. Calorim.* **62**, 163 (2000).
- 2000YU/MEN S. Yu, S. Meng, Z. Tan, L. Li, and J. Zhang, *Taiyangneng Xuebao* **21**, 171 (2000); *Chem. Abstr.* **133**, 166195.
- 2000YU/STE L. Yu, G. A. Stephenson, C. A. Mitchell, C. A. Bunnell, S. V. Snorek, J. J. Bowyer, T. B. Borchardt, J. G. Stowell, and S. R. Byrn, *J. Am. Chem. Soc.* **122**, 585 (2000).
- 2000ZHA/STA G. I. Zharkova, P. A. Stabnikov, V. M. Grankin, P. P. Semynnikov, and I. K. Igumenov, *Russ. J. Coord. Chem.* **26**, 576 (2000).
- 2000ZIE W. Zielenkiewicz, *J. Chem. Eng. Data* **45**, 626 (2000).
- 2001AHN/HAR S. Ahn, K. D. M. Harris, B. M. Kariuki, and D. M. S. Zin, *J. Solid State Chem.* **156**, 10 (2001).
- 2001ALB K. C. Albyn, *J. Chem. Eng. Data* **46**, 1415 (2001).
- 2001ALB/HAH M. Albert, I. Hahnenstein, H. Hasse, and G. Maurer, *J. Chem. Eng. Data* **46**, 897 (2001).
- 2001ASA/SOR S. Asahina and M. Sorai, *Liq. Cryst.* **28**, 1085 (2001).
- 2001BAE A. K. Baev, *Izv. Vyssh. Uchebn. Zaved., Khim. Khim. Tekhnol.* **44**, 3 (2001).
- 2001BAE/CHE A. K. Baev and I. N. Chernyak, *Russ. J. Inorg. Chem.* **46**, 310 (2001); *Zh. Neorg. Khim.* **46**, 361 (2001). [Note: enthalpies of vaporization are presented graphically in paper for binary mixtures.]
- 2001BAE/MIK A. K. Baev and V. E. Mikhailov, *Russ. J. Appl. Chem.* **74**, 1435 (2001).
- 2001BAE2 A. K. Baev, *Russ. J. Appl. Chem.* **74**, 162 (2001).
- 2001BER/WIC S. Bernatova and I. Wichterle, *Fluid Phase Equilib.* **189**, 111 (2001).
- 2001BLO/VAN J. G. Blok, A. C. G. van Genderen, P. R. van der Linde, and H. A. J. Oonk, *J. Chem. Thermodyn.* **33**, 1097 (2001).
- 2001BUR/JOS N. Bureau, J. Jose, I. Mokbel, and J.-C. de Hemptinne, *J. Chem. Thermodyn.* **33**, 1485 (2001).
- 2001CAL/MEL M. Calligaris, A. Melchior, and S. Geremia, *Inorg. Chim. Acta* **323**, 89 (2001).
- 2001CEN/LIP M. Censky, M. Lipovska, H.-G. Schmidt, V. Ruzicka, and G. Wolf, *J. Therm Anal. Calorim.* **63**, 879 (2001).
- 2001CHA/DOL K. Chatterjee, D. Dollimore, and K. Alexander, *Instrum. Sci. Technol.* **29**, 133 (2001); *Chem. Abstr.* **135**, 16629t (2001). (Note: the reported values are referred to as enthalpies of vaporization).
- 2001CHA/TOB S. Chattopadhyay, H. J. Tobias, and P. J. Ziemann, *Anal. Chem.* **73**, 3797 (2001).
- 2001CHE/HUA Y. Cheng, Y. Huang, K. Alexander, and D. Dollimore, *Thermochim. Acta* **367/368**, 23 (2001).
- 2001CHY/FRA K. Chylinski, Z. Frasz, and S. K. Malanowski, *J. Chem. Eng. Data* **46**, 29 (2001).
- 2001COL/LAU C. Colominas, K. H. Lau, D. L. Hildenbrand, S. Crouch-Baker, and A. Sanjurjo, *J. Chem. Eng. Data* **46**, 446 (2001).
- 2001DAB/MIS M. Dabrowski, B. Misterkiewicz and A. Sporzynski, *J. Chem. Eng. Data* **46**, 1627 (2001).
- 2001DAM/BLA F. Damian, N. Blaton, P. Augustijns, L. Naesens, J. Balzarini, R. Kinget, and G. Van den Mooter, *Thermochim. Acta* **366**, 61 (2001).
- 2001DI/SUN Y.-Y. Di, X.-H. Sun, Y.-F. Liu, Z.-C. Tan, S.-H. Meng, S.-L. Gao, and S.-S. Qu, *Huaxue Xuebao* **59**, 496 (2001); *Chem. Abstr.* **134**, 372309w.
- 2001DI/SUN2 Y.-Y. Di, X.-H. Sun, Z.-C. Tan, Y.-F. Liu, S.-H. Meng, S.-L. Gao, and S.-S. Qu, *Thermochim. Acta* **369**, 25 (2001).
- 2001DIN/MUR T. J. Dingemans, N. S. Murthy, and E. T. Samulski, *J. Phys. Chem. B* **105**, 8845 (2001).
- 2001DIO/PIE H. P. Diogo, M. E. Minas da Piedade, J. M. Goncalves, M. J. S. Monte, and M. A. V. Ribeiro da Silva, *Eur. J. Inorg. Chem.* 257 (2001).
- 2001DOR/LOB P. A. Dorozhko, A. S. Lobach, A. A. Popov, V. M. Senyavin, and M. V. Korobov, *Chem. Phys. Lett.* **336**, 39 (2001).
- 2001FED/GEL N. E. Fedotova, N. V. Gelfond, I. K. Igumenov, A. N. Mikheev, N. B. Morozova, and R. H. Tuffias, *Int. J. Therm. Sci.* **40**, 469 (2001).
- 2001GIO D. Giordano, *J. Chem. Eng. Data* **46**, 486 (2001).
- 2001HE/GRI X. He, U. J. Griesser, J. G. Stowell, T. B. Borchardt, and S. B. Byrn, *J. Pharm. Sci.* **90**, 371 (2001).
- 2001HE/STO X. He, J. G. Stowell, K. R. Morris, R. R. Pfeiffer, H. Li, P. Stahly, and S. R. Byrn, *Cryst. Growth Des.* **1**, 305 (2001).
- 2001HEI/VER A. Heintz and S. P. Verevkin, *Fluid Phase Equilib.* **179**, 85 (2001).
- 2001HOR/GAR S. Horstmann, H. Gardeler, K. Fischer, F. Köster, and J. Gmehling, *J. Chem. Eng. Data* **46**, 337 (2001).
- 2001IWA/MIN M. Iwahashi, H. Minami, T. Suzuki, M. Koyanagi, H. Hao, K. Ema, and K. Ashizawa, *J. Oleo Sci.* **50**, 693 (2001).
- 2001JAM/DOB M. E. Jamroz, J. C. Dobrowolski, J. Placzek, A. M. Szafranski, J. K. Kazimirski, and Z. Lisicki, *J. Chem. Thermodyn.* **33**, 565 (2001).
- 2001KAM/SUE R. Kamae, K. Suenaga, T. Matsuo, and H. Suga, *J. Chem. Thermodyn.* **33**, 471 (2001).
- 2001KIY/MIN T. Kiyobayashi and M. E. Minas da Piedade, *J. Chem. Thermodyn.* **33**, 11 (2001).
- 2001KLI/LUB W. Kliegel, G. Lubkowitz, J. O. Pokriefke, S. J. Retting, and J. Trotter, *Can. J. Chem.* **79**, 226 (2001).
- 2001KUL/DES I. Kul, D. D. DesMarteau, and A. L. Beyerlein, *Fluid Phase Equilib.* **185**, 241 (2001).
- 2001KUL/VER D. Kulikov, S. P. Verevkin, and A. Heintz, *J. Chem. Eng. Data* **46**, 1593 (2001).
- 2001KUL/VER2 D. Kulikov, S. P. Verevkin, and A. Heintz, *Fluid Phase Equilib.* **192**, 187 (2001).
- 2001LAG/DIO A. L. C. Lagoa, H. P. Diogo, M. P. Dias, M. E. Minas da Piedade, L. M. P. E. Amaral, M. A. V. Ribeiro da Silva, J. A. M. Simoes, R. C. Guedes, B. J. Costa Cabral, K. Schwarz, and M. Eppli, *Chem.-Eur. J.* **7**, 483 (2001).
- 2001LAI/LEE L.-L. Lai, L.-J. Lee, E. Wang, and F.-Y. Su, *Liq. Cryst.* **28**, 381 (2001).
- 2001LAL/SCH J. O. Lalah, K.-W. Schramm, D. Lenoir, B. Henkelmann, N. Hertkorn, G. Matuschek, A. Kettrup, and K. Günther, *Ceram. Eng. Sci. Proc.* **7**, 4790 (2001).
- 2001LEB/VAN N. Lebrun and J. C. van Miltenburg, *J. Alloys Compd.* **320**, 320 (2001).
- 2001LEG/BAZ B. Legendre, G. Baziard-Mouysson, M. Anastassiadou, J. M. Leger, and M. Payard, *J. Therm Anal. Calorim.* **66**, 659 (2001).
- 2001LI/HE J. Li, Y. He, K. Ishida, T. Yamane, and Y. Inoue, *Polym. J. (Tokyo, Jpn.)* **33**, 773 (2001).
- 2001LOR/AUC S. Loras, A. Aucejo, J. B. Monton, J. Wisniak, and H. Segura, *J. Chem. Eng. Data* **46**, 1351 (2001).
- 2001MAL/PAR I. P. Malkerova, S. E. Paramonov, A. S. Alikhanyan, and N. P. Kuz'mina, *Zh. Neorg. Khim.* **46**, 1700 (2001).
- 2001MAT/SHC E. V. Matukhina, O. I. Shchegolikhina, N. N. Makarova, Y. A. Pozdniakova, and D. E. Katsoulis, *Liq. Cryst.* **28**, 869 (2001).
- 2001MON/HIL M. J. S. Monte and D. M. Hillesheim, *J. Chem. Eng. Data*

- 46, 1601 (2001).
- 2001MON/HIL2 M. J. S. Monte and D. M. Hillesheim, *J. Chem. Thermodyn.* **33**, 745 (2001).
- 2001MON/HIL3 M. J. S. Monte and D. M. Hillesheim, *J. Chem. Thermodyn.* **33**, 103 (2001).
- 2001MON/HIL3 M. J. S. Monte and D. M. Hillesheim, *J. Chem. Thermodyn.* **33**, 837 (2001).
- 2001MOR/ZHA N. B. Morozova, S. I. Zharkova, P. P. Semyannikov, I. K. Igumenov, N. E. Fedotova, and N. V. Gelford, *J. Phys. IV* **11**, Pr3/609 (2001); *Chem. Abstr.* **136**, 43187 (2001).
- 2001MUN/KRA L. A. L. Munoz and M. A. Krahenbuhl, *J. Chem. Eng. Data* **46**, 120 (2001).
- 2001NIK/SUL G. E. Nikitina, B. Suleiman, A. S. Semeikin, and O. A. Golubchikov, *Russ. J. Phys. Chem.* **75**, 675 (2001).
- 2001NOR/TOU S. Norvez, F.-G. Tournilhac, P. Bassoul, and P. Herson, *Chem. Mater.* **13**, 2552 (2001).
- 2001ORT/GON J. Ortega, C. Gonzalez, and S. Galvan, *J. Chem. Eng. Data* **46**, 904 (2001).
- 2001OXL/SMI J. C. Oxley, J. L. Smith, J. Zhang, and C. Bedford, *J. Phys. Chem. A* **105**, 579 (2001). (The authors report that compounds exhibit melting endotherm well below their decomposition exotherm.)
- 2001PER/BAU G. L. Perlovich and A. Bauer-Brandl, *J. Therm Anal. Calorim.* **63**, 653 (2001).
- 2001PUR/CHI S. Puri, J. S. Chickos, and W. Welsh, *Anal. Chem.* **73**, 1480 (2001).
- 2001RAI/VAR R. N. Rai and K. B. R. Varma, *Mater. Lett.* **48**, 356 (2001).
- 2001RIB/FER M. D. M. C. Ribeiro da Silva, S. C. C. Ferreira, I. A. P. Rodrigues, L. C. M. da Silva, W. E. Acree Jr., S. Pandey, and L. E. Roy, *J. Chem. Thermodyn.* **33**, 1227 (2001).
- 2001RIB/GON M. D. M. C. Ribeiro da Silva, J. M. Goncalves, S. C. C. Ferreira, L. C. M. da Silva, M. J. Sottomayor, G. Pilcher, W. E. Acree, Jr., and L. E. Roy, *J. Chem. Thermodyn.* **33**, 1263 (2001).
- 2001RIB/MAT M. A. V. Ribeiro da Silva, M. A. R. Matos, M. S. Miranda, M. H. F. A. Sousa, R. M. Borges dos Santos, M. M. Bizarro, and J. A. Martinho *Struct. Chem.* **12**, 171 (2001).
- 2001RIB/MON M. A. V. Ribeiro da Silva, M. J. S. Monte, and E. Giera, *J. Chem. Thermodyn.* **33**, 369 (2001).
- 2001RIB/RIB M. A. V. Ribeiro da Silva, M. D. M. C. Ribeiro da Silva, L. C. M. da Silva, F. Dietz, E. Hoyer, L. Beyer, B. Schröder, A. M. Damas, and J. F. Liebman, *J. Chem. Soc., Perkin Trans. 2* **2**, 2174 (2001).
- 2001RIB/RIB2 M. A. V. Ribeiro da Silva, M. D. M. C. Ribeiro da Silva, L. C. M. da Silva, F. Dietz, and E. Hoyer, *Thermochim. Acta* **378**, 45 (2001).
- 2001RIT L. Rittfeldt, *Anal. Chem.* **73**, 2405 (2001).
- 2001ROJ/ORO A. Rojas-Aguilar, E. Orozco-Guareno, and M. Martinez-herra, *J. Chem. Thermodyn.* **33**, 1405 (2001).
- 2001ROL/BUR J. M. Rollinger and A. Burger, *J. Pharm. Sci.* **90**, 949 (2001).
- 2001ROT/GLA M. K. Rotich, B. D. Glass, and M. E. Brown, *J. Therm Anal. Calorim.* **64**, 681 (2001).
- 2001ROU/JIM M. V. Roux, P. Jimenez, J. Z. Davalos, R. Notario, and E. Juaristi, *J. Org. Chem.* **66**, 5343 (2001).
- 2001ROU/JIM2 M. V. Roux, P. P. Jimenez, A. Mayorga, J. Z. Davalos, S. Böhm, and O. Exner, *J. Phys. Chem. A* **105**, 7926 (2001).
- 2001SCI/SCI J. Sciesinski, E. Sciesinski, M. Massalska-Arodz, T. Wasutynski, P. M. Zielinski, and W. Witko, *IEEE Trans. Dielectr. Electr. Insul.* **8**, 522 (2001).
- 2001SEG/LAM H. Segura, E. Lam, R. Reich, and J. Wisniak, *Phys. Chem. Liq.* **39**, 43 (2001).
- 2001SMI/LEB N. N. Smirnova and B. V. Lebedev, *Russ. J. Phys. Chem.* **75**, S118 (2001).
- 2001SON/HUA Y. Song, Y. Huang, E. A. Havenga, and I. S. Butler, *Vib. Spectrosc.* **27**, 127 (2001).
- 2001TAN/ZUE Z.-C. Tan, B. Xue, S.-W. Lu, S. H. Meng, X.-H. Yuan, and Y. J. Song, *J. Therm Anal. Calorim.* **63**, 297 (2001).
- 2001TIT/TOM S. A. Tittlemier and G. T. Tomy, *Envir. Toxicol. Chem.* **20**, 146 (2001).
- 2001TUR/KRI A. E. Turgambaeva, V. V. Krisyuk, S. V. Sysoev, and I. K. Igumenov, *Chem. Vap. Deposition* **7**, 121 (2001).
- 2001UUS/POK P. Uusi-Kyyny, J.-P. Pokki, J. Aittaman, and S. Liukkonen, *J. Chem. Eng. Data* **46**, 1244 (2001).
- 2001VAL/SMI E. J. Valente, T. N. Smith, and M. E. Harris, *Chirality* **13**, 244 (2001).
- 2001VAN/OON J. C. van Miltenburg, H. A. J. Oonk, and B. J. K. Van den Berg, *J. Chem. Eng. Data* **46**, 84 (2001).
- 2001VAN/OON2 J. C. van Miltenburg, H. A. J. Oonk, and L. Ventola, *J. Chem. Eng. Data* **46**, 90 (2001).
- 2001VAN/PAS R. M. Varushchenko, L. L. Paschenko, A. I. Druzhinina, A. V. Abramnikov, and A. A. Pimersin, *J. Chem. Thermodyn.* **33**, 733 (2001).
- 2001VAR/DRU R. M. Varushchenko, A. I. Druzhinina, A. Yu. Churkina, and Z.-C. Tan, *Russ. J. Phys. Chem.* **75**, 1223 (2001).
- 2001VER/AUG S. Verheyen, P. Augustijns, R. Kinget, and G. Van den Mooter, *Int. J. Pharm.* **228**, 199 (2001).
- 2001VER/HEI S. P. Verevkin and A. Heintz, *J. Chem. Eng. Data* **46**, 984 (2001).
- 2001VER/HEI2 S. P. Verevkin and A. Heintz, *J. Chem. Eng. Data* **46**, 41 (2001).
- 2001VER/VAZ E. Vercher, M. I. Vazquez, and A. Martinez-Andreu, *J. Chem. Eng. Data* **46**, 1584 (2001).
- 2001WON/LEI A. Wong, Y. D. Lei, M. Alaei, and F. Wania, *J. Chem. Eng. Data* **46**, 239 (2001).
- 2001YAT/MIN T. Yatabe, N. Minami, H. Okumoto, A. Kaito, K. Ueno, and Y. Tanabe, *Mol. Cryst. Liq. Cryst.* **365**, 7 (2001).
- 2001YOU/SCH P. H. Young and C. A. Schall, *Thermochim. Acta* **367/368**, 387 (2001).
- 2001ZHU/LI L. Zhu, H. Li, C. Wang, and S. Han, *J. Chem. Eng. Data* **46**, 1231 (2001).
- 2001ZOR/COS H. E. Zorel, Jr., A. G. C. Costalonga, M. S. Crespi, and C. A. Ribiero, *Quim. Nova* **24**, 599 (2001).
- 2002ABB/CAS J.-L. M. Abboud, O. Castao, J. Z. Davalos, P. Jimenez, R. Gomperts, P. Muller, and M. V. Roux, *J. Org. Chem.* **67**, 1057 (2002).
- 2002ABB/WOH T. P. Abbott, A. Wohlman, T. Isbell, F. A. Momany, C. Cantrell, D. Garlotta, and D. Weisleder, *Ind. Crop Products* **16**, 43 (2002).
- 2002ANT/FRA M. Antosik, Z. Frasz, and S. K. Malanowski, *J. Chem. Eng. Data* **47**, 757 (2002).
- 2002BAE/SHI A. K. Baev and M. A. Shishko, *Russ. J. Appl. Chem.* **75**, 25 (2002).
- 2002BAE/SHI2 A. K. Baev and M. A. Shishko, *Russ. J. Appl. Chem.* **75**, 156 (2002).
- 2002BAT I. Batiu, *Fluid Phase Equilib.* **198**, 111 (2002).
- 2002BEL/MAN M. Belloni, M. Manickam, and J. A. Preece, *Ferroelectrics* **276**, 103 (2002).
- 2002BLO/PAU A. V. Blokhin, Y. U. Paulechka, G. J. Kabo, and A. A. Kozyro, *J. Chem. Thermodyn.* **34**, 29 (2002).
- 2002BOB/ART S. Bobbo, G. Artico, L. Fedele, M. Scattolini, and R. Camporese, *J. Chem. Eng. Data* **47**, 839 (2002).
- 2002BOB/FED S. Bobbo, L. Fedele, M. Scattolini, and R. Camporese, *J. Chem. Eng. Data* **47**, 179 (2002).
- 2002BOR/CES B. Borde and A. Cesaro, *J. Therm Anal. Calorim.* **69**, 267 (2002).
- 2002BOU/SAI M. Bouvet and H. Said, *Phys. Chem. News* **8**, 79 (2002); *Chem. Abstr.* **138**, 344563 (2002).
- 2002BRO/DU S. Brownridge, H. Du, S. A. Fairhurst, R. C. Haddon, H. Oberhammer, S. Parson, J. Passmore, M. J. Schriver, L. H. Sutcliffe, and N. P. C. Westwood, *J. Chem. Soc. Dalton Trans.* 3365 (2002).
- 2002BRU/POR B. Brunetti, G. Portalone, and V. Piacente, *J. Chem. Eng. Data* **47**, 17 (2002).
- 2002CAI/BET M. R. Caira, G. Bettinetti, and M. Sorrenti, *J. Pharm. Sci.* **91**, 467 (2002).
- 2002CHA/DOL K. Chatterjee, D. Dollimore, and K. S. Alexander, *Thermochim. Acta* **392-393**, 107 (2002).
- 2002CHA/MAN D. Chandra, H. Mandalia, W.-M. Chien, D. W. Lindle, and R. Rudman, *Z. Phys. Chem.* **216**, 1389 (2002).
- 2002CHI/ACR J. S. Chickos and W. E. Acree, Jr., *J. Phys. Chem. Ref. Data* **31**, 537 (2002).
- 2002CHI/HIL J. S. Chickos, D. M. Hillesheim, G. Nichols, and M. J. Zehe, *J. Chem. Thermodyn.* **34**, 1647 (2002).

- 2002CHI/KN1 R. D. Chirico, S. E. Knipmeyer, and W. V. Steele, *J. Chem. Thermodyn.* **34**, 1873 (2002).
- 2002CHI/KN12 R. D. Chirico, S. E. Knipmeyer, and W. V. Steele, *J. Chem. Thermodyn.* **34**, 1885 (2002).
- 2002CHI/WEB J. S. Chickos, P. Webb, G. Nichols, T. Kiyobayashi, P.-C. Cheng, and L. Scott, *J. Chem. Thermodyn.* **34**, 1195 (2002).
- 2002CON/WIC D. Constantinescu and I. Wichterle, *Fluid Phase Equilib.* **203**, 71 (2002).
- 2002DAH/MOK A. Dahmani, I. Mokbel, and J. Jose, *Fluid Phase Equilib.* **202**, 193 (2002).
- 2002DAL/DEL L. Dall'acqua, G. Della Gatta, B. Nowicka, and P. Ferloni, *J. Chem. Thermodyn.* **34**, 1 (2002).
- 2002DAS/ZAL Z. Daszkiewicz, J. Zaleski, E. M. Nowakowska, and J. B. Kyziol, *Pol. J. Chem.* **76**, 1113 (2002).
- 2002DEN/LI D. Deng, H. Li, and S. Han, *J. Chem. Thermodyn.* **34**, 1431 (2002).
- 2002DIA/LOP J. Diaz-Hernandez, A. Lopez-Echarri, I. Ruiz-Lauren, A. Fraile-Rodriguez, and T. Breczewski, *Ferroelectrics* **268**, 753 (2002).
- 2002DIN/PAS G. Di Nicola and G. Passerini, *J. Chem. Eng. Data* **47**, 882 (2002).
- 2002DIO/KIY H. P. Diogo, T. Kiyobayashi, M. E. Minas da Piedade, N. Burlak, D. W. Rogers, D. McMasters, G. Persy, J. Wirz, and J. F. Liebman, *J. Am. Chem. Soc.* **124**, 2065 (2002).
- 2002DIO/MIN H. P. Diogo and M. E. Minas da Piedade, *J. Chem. Thermodyn.* **34**, 173 (2002).
- 2002DOM/KOZ U. Domanska, M. K. Kozłowska, and M. Rogalski, *J. Chem. Eng. Data* **47**, 456 (2002).
- 2002GOO/GEL J. W. Goodrum and D. P. Geller, *Bioresour. Technol.* **84**, 75 (2002). (Note: the vaporization enthalpies that were calculated with the Clausius–Clapeyron equation coefficients given in the paper were less than 1 kJ/mole. Through personal correspondence with the authors we learned that the coefficients are to be multiplied by 1000).
- 2002GOO/LAI C. Goosen, T. J. Laing, J. du Plessis, T. C. Goosen, and J. L. Flynn, *Pharm. Res.* **19**, 13 (2002).
- 2002GRA/RAS S. Gracin and A. C. Rasmuson, *J. Chem. Eng. Data* **47**, 1379 (2002).
- 2002HAN/HIK M. Hanaya, T. Hikira, M. Hatase, and M. Oguni, *J. Chem. Thermodyn.* **34**, 1173 (2002).
- 2002HAZ/DOL A. Hazra, D. Dollimore, and K. Alexander, *Thermochim. Acta* **392/393**, 221 (2002).
- 2002HER/ACK A. K. Hermetet, L. J. Ackerman, J. K. Swearingen, C. A. Presto, D. R. Kelman, J. M. Giesen, K. I. Goldberg, W. Kaminsky, and D. M. West, *J. Chem. Cryst.* **32**, 17 (2002).
- 2002ITA/KAM T. Itahara, J. Kamada, S. Ikeda, and T. Ueda, *Chem. Lett.* **2002**, 1122.
- 2002JIM/ROU P. Jimenez, M. V. Roux, J. Z. Davalos, and M. T. Molina, *J. Chem. Thermodyn.* **34**, 1117 (2002).
- 2002JIM/ROU2 P. Jimenez, M. V. Roux, J. Z. Davalos, and M. Temprado, *Thermochim. Acta* **394**, 25 (2002).
- 2002JON/COO S. O. Jonsdottir, S. A. Cooke, and E. A. Macedo, *Carbohydr. Res.* **337**, 1563 (2002).
- 2002JON/LIG D. E. G. Jones, P. D. Lightfoot, R. C. Fouchard, and Q. S. M. Kwok, *Thermochim. Acta* **388**, 159 (2002).
- 2002JOW/DIN K. G. Jow and T. J. Dingemans, *Liq. Cryst.* **29**, 573 (2002).
- 2002KAR/SHE N. V. Karyakin, M. S. Sheiman, M. S. Kozlova, G. P. Kamelova, V. N. Larina, and A. S. Smirnov, *Russ. J. Phys. Chem.* **76**, 1061 (2002).
- 2002KEL/SZC D. R. Kelman, L. F. Szczepura, K. I. Goldberg, W. Kaminsky, A. K. Hermetet, L. G. Ackerman, J. K. Swearingen, and D. X. West, *J. Mol. Struct.* **610**, 143 (2002).
- 2002KOZ/KAR M. S. Kozlova, N. V. Karyakin, M. S. Sheiman, G. P. Kamelova, and V. N. Larina, *Russ. J. Phys. Chem.* **76**, 1068 (2002).
- 2002KRA/VAS E. L. Krasnykh, T. V. Vasil'tsova, S. P. Verevkin, and A. Heintz, *J. Chem. Eng. Data* **47**, 1372 (2002).
- 2002KUS/ASH I. Kushida and K. Ashizawa, *J. Pharm. Sci.* **91**, 2193 (2002).
- 2002LAG/DIO A. L. C. Lagoa, H. P. Diogo, M. E. Minas da Piedade, L. M. P. F. Amaral, R. C. Guedes, B. J. C. Cabral, D. V. Kulikov, S. P. Verevkin, M. Siedler, and M. Epple, *J. Phys. Chem. A* **106**, 9855 (2002).
- 2002LEB/BYK B. V. Lebedev, T. A. Bykova, A. V. Markin, A. R. Korogodskii, V. V. Kireev, and N. N. Sanina, *Polym. Sci., Ser. A Ser. B* **44**, 1327 (2002).
- 2002LEI/CHA Y. D. Lei, R. Chankalal, A. Chan, and F. Wania, *J. Chem. Eng. Data* **47**, 801 (2002).
- 2002LOR/AUC S. Loras, A. Aucejo, J. B. Monton, J. Wisniak, and H. Segura, *J. Chem. Eng. Data* **47**, 1256 (2002).
- 2002LU/SON Q.-Y. Lu, Y.-J. Song, S.-H. Meng, J. Xing, and Z.-C. Tan, *Wuli Huaxue Xuebao* **18**, 166 (2002); *Chem. Abstr.* **136**, 134441e (2002).
- 2002LUB/BAN M. Lubomska, A. Banas, and S. K. Malanowski, *J. Chem. Eng. Data* **47**, 1466 (2002).
- 2002MAR/BOL V. Yu. Markov, O. V. Boltalina, A. A. Goryunkov, N. Sidorov, and R. Taylor, *J. Chem. Thermodyn.* **34**, 57 (2002).
- 2002MAR/GOM F. Martinez and A. Gomez, *Pattern Anal. Appl.* **40**, 411 (2002).
- 2002MAR/OLI I. M. Marrucho, N. S. Oliveira, and R. Dohrn, *J. Chem. Eng. Data* **47**, 554 (2002).
- 2002MEN/DOL D. Menon, D. Dollimore, and K. S. Alexander, *Thermochim. Acta* **392/393**, 237 (2002).
- 2002MIN/SAK H. Minami, Y. Sakamoto, and M. Iwahashi, *J. Oleo Sci.* **51**, 11 (2002).
- 2002MIR/LEB E. A. Miroshnichenko, V. P. Lebedev, and Y. N. Matyushin, *Dokl. Akad. Nauk* **382**, 497 (2002).
- 2002MIY/KAT K. Miyazawa, T. Kato, M. Itoh, and Ushioda, *Liq. Cryst.* **29**, 1483 (2002).
- 2002MUR/YAM J. Murata, S. Yamashita, M. Akiyama, S. Katayama, T. Hiaki, and A. Sekiya, *J. Chem. Eng. Data* **47**, 911 (2002).
- 2002MUS/RAZ A. M. Musuc, D. Razus, and D. Oancea, *Analele Universitatiei Bucuresti Chimie* **11**, 147 (2002).
- 2002NAT/JES C. Nather, I. Jess, Z. Havlas, M. Bolte, N. Nagel, and S. Nick, *Solid State Sci.* **4**, 859 (2002).
- 2002PAN/MAL R. Pankajavalli, C. Mallika, O. M. Sreedharan, V. S. Raghunathan, P. A. Premkumar, and K. S. Negaraja, *Chem. Eng. Sci.* **57**, 3603 (2002).
- 2002PAT/CAM R. Patino, M. Campos, and L. A. Torres, *J. Chem. Thermodyn.* **34**, 193 (2002).
- 2002PER/HAN G. L. Perlovich, L. Kr. Hansen, and A. Bauer-Brandl, *J. Pharm. Sci.* **91**, 1036 (2002).
- 2002PER/LOP E. Perea, F. Lopez-Calahorra, and D. Velasco, *Liq. Cryst.* **29**, 421 (2002).
- 2002PFO/RIE O. Pfohl, C. Riebesell, and R. Dohrn, *Fluid Phase Equilib.* **202**, 289 (2002).
- 2002PIM/MEL S. M. Pimehova, S. V. Melkhanova, V. P. Kolesov, P. I. Demyanov, A. N. Fedotov, and V. P. Vorbieva, *J. Chem. Thermodyn.* **34**, 385 (2002).
- 2002POK/UUS J.-P. Pokki, P. Uusi-Kyyny, J. Aittamaa, and S. Liukkonen, *J. Chem. Eng. Data* **47**, 371 (2002).
- 2002RAI/PAN U. S. Rai, P. Pandey, and R. N. Rai, *Mater. Lett.* **53**, 83 (2002).
- 2002RIB/AMA M. A. V. Ribeiro da Silva, L. M. P. F. Amaral, and I. M. C. L. Ferreira, *J. Chem. Thermodyn.* **34**, 119 (2002).
- 2002RIB/RIB M. A. V. Ribeiro da Silva, M. D. M. C. Ribeiro da Silva, L. C. M. da Silva, and F. Dietze, *J. Chem. Thermodyn.* **34**, 155 (2002).
- 2002ROC/GRI G. D. Rockwell, T. B. Grindley, K. C. Smith, and M. A. White, *J. Phys. Chem. B* **106**, 12311 (2002).
- 2002ROD/CAN A. Rodriguez, J. Canosa, A. Dominguez, and J. Tojo, *Fluid Phase Equilib.* **198**, 95 (2002).
- 2002ROD/CAN2 A. Rodriguez, J. Canosa, A. Dominguez, and A. Tojo, *J. Chem. Eng. Data* **47**, 1098 (2002).
- 2002ROG/DOM M. Rogalski, U. Domanska, D. Czyrny, and D. Dyczko, *Chem. Phys.* **285**, 355 (2002).
- 2002ROL/GST J. M. Rollinger, E. M. Gstrein, and A. Burger, *Eur. J. Pharm. Biopharm.* **53**, 75 (2002).
- 2002ROU/DAV M. V. Roux, J. Z. Davalos, and P. Jimenez, *Thermochim. Acta* **394**, 19 (2002).
- 2002ROY/RIG S. Roy, A. T. Riga, and K. S. Alexander, *Thermochim. Acta* **392/393**, 399 (2002).

- 2002SCH/LAS A. Schulz, S. Laschat, M. Morr, S. Diele, M. Dreyer, and G. Bringmann, *Helv. Chim. Acta* **85**, 3909 (2002).
- 2002SEG/GAL H. Segura, G. Galindo, R. Reich, J. Wisniak, and S. Loras, *Phys. Chem. Liq.* **40**, 277 (2002).
- 2002SEG/WIS H. Segura, J. Wisniak, G. Galindo, and R. Reich, *Phys. Chem. Liq.* **40**, 67 (2002).
- 2002SHE/KAR V. M. Sheiman, N. V. Karyakin, B. I. Kozyrkin, and I. A. Zelyaev, *Russ. J. Phys. Chem.* **76**, 885 (2002).
- 2002SHE/KAR2 V. M. Sheiman, N. V. Karyakin, M. S. Sheiman, and I. A. Zelyaev, *Russ. J. Phys. Chem.* **76**, 881 (2002).
- 2002SII/KIR E. Siimer, H. Kirss, M. Kuus, and L. Kudryavtseva, *Proc. Est. Acad. Sci. Chem.* **51**, 19 (2002).
- 2002SOR/DOL T. V. Sorokina, D. Dollimore, and K. S. Alexander, *Thermochim. Acta* **392/393**, 315 (2002).
- 2002SPA/DZI A. Spadlo, J. Dziaduszek, R. Dabrowski, K. Cyprynski, T. Stolarz, and S. T. Wu, *Proc. SPIE* **4759**, 79 (2002).
- 2002STE/CHI W. V. Steele, R. D. Chirico, S. E. Knipmeyer, and A. Nguyen, *J. Chem. Eng. Data* **47**, 648 (2002).
- 2002STE/CHI2 W. V. Steele, R. D. Chirico, A. B. Cowell, S. E. Knipmeyer, and A. Nguyen, *J. Chem. Eng. Data* **47**, 667 (2002).
- 2002STE/CHI3 W. V. Steele, R. D. Chirico, S. E. Knipmeyer, and A. Nguyen, *J. Chem. Eng. Data* **47**, 689 (2002).
- 2002STE/CHI4 W. V. Steele, R. D. Chirico, A. B. Cowell, S. E. Knipmeyer, and A. Nguyen, *J. Chem. Eng. Data* **47**, 700 (2002).
- 2002STE/CHI5 W. V. Steele, R. D. Chirico, S. E. Knipmeyer, and A. Nguyen, *J. Chem. Eng. Data* **47**, 715 (2002).
- 2002STE/CHI6 W. V. Steele, R. D. Chirico, A. B. Cowell, S. E. Knipmeyer, and A. Nguyen, *J. Chem. Eng. Data* **47**, 725 (2002).
- 2002STO/BEH G. W. Stowell, R. J. Behme, S. M. Denton, I. Pfeiffer, F. D. Sancilio, L. B. Whittall, and R. R. Whittle, *J. Pharm. Sci.* **91**, 2481 (2002).
- 2002SU C. C. Su, MS thesis, National Taiwan University of Science and Technology, 2002.
- 2002SU/ZHU C.-H. Su, S. Zhu, N. Ramachandran, and A. Burger, *J. Cryst. Growth* **235**, 313 (2002).
- 2002SWI/MAL B. E. Swiatek and S. K. Malanowski, *J. Chem. Eng. Data* **47**, 478 (2002).
- 2002SZC/KEL L. F. Szczepura, D. R. Kelman, A. K. Hermetet, L. J. Ackerman, K. I. Goldberg, K. A. Claborn, W. Kaminsky, and D. X. West, *J. Mol. Struct.* **608**, 245 (2002).
- 2002SZT/KAM P. Szterner, M. Kaminski, and A. Zielenkiewicz, *J. Chem. Thermodyn.* **34**, 1005 (2002).
- 2002TAN/SUN Z.-C. Tan, L.-X. Sun, S.-H. Meng, L. Li, F. Xu, P. Yu, B.-P. Liu, and J.-B. Zhang, *J. Chem. Thermodyn.* **34**, 1417 (2002).
- 2002TAT/DOL A. S. Tatavarti, D. Dollimore, and K. S. Alexander, *J. Pharm. Sci.* **4**, 1 (2002).
- 2002TOR/CAM L. A. Torres, M. Campos, E. Enriquez, and R. Patino, *J. Chem. Thermodyn.* **34**, 293 (2002).
- 2002VAL/COQ A. Valtz, C. Coquelet, A. Baba-Ahmed, and D. Richon, *Fluid Phase Equilib.* **202**, 29 (2002).
- 2002VAL/HER J. Vales-Martinez, S. Hernandez-Ortega, G. Espinosa-Perez, C. A. Presto, A. K. Hermetet, K. D. Haslow, L. J. Ackerman, L. F. Szczepura, K. I. Goldberg, W. Kaminsky, and D. X. West, *J. Mol. Struct.* **608**, 77 (2002).
- 2002VAN/KRU J. A. H. van Laarhoven, M. A. B. Krufft, and H. Vromans, *Int. J. Pharm.* **232**, 163 (2002).
- 2002VAN/PAR A. van Roon, J. R. Parsons, and H. A. J. Govers, *J. Chromatogr. A* **955**, 105 (2002).
- 2002VAN/VAN A. C. G. van Genderen, J. C. van Miltenburg, J. G. Blok, M. J. van Bommel, P. J. van Ekeren, G. J. K. van den Berg, and H. A. J. Oonk, *Fluid Phase Equilib.* **202**, 109 (2002).
- 2002VAN/VAN2 J. C. van Miltenburg and P. J. van Ekeren, *Thermochim. Acta* **385**, 11 (2002).
- 2002VAR/AIT R. M. Varuschenko, C. A. Aitkeeva, A. I. Druzhinina, Y. A. Myshenseva, and L. L. Pashchenko, *Russ. J. Phys. Chem.* **76**, 724.
- 2002VAR/DRU R. M. Varuschenko and A. I. Druzhinina, *Fluid Phase Equilib.* **199**, 109 (2002).
- 2002VAR/PAS R. M. Varushchenko, L. L. Pashchenko, A. Y. Churkina, and A. V. Shabanova, *Russ. J. Phys. Chem.* **76**, 915 (2002).
- 2002VAR/PAS2 R. M. Varuschenko, L. L. Pashchenko, A. I. Druzhinina, A. Yu. Churkina, I. I. Vorob'ev, and S. N. Kravhun, *Russ. J. Phys. Chem.* **76**, 546 (2002).
- 2002VEN/RAM L. Ventola, M. Ramirez, T. Calvet, X. Solans, M. A. Cuevas-Diarte, N. Negrier, D. Mondieig, J. C. van Miltenburg, and H. A. J. Oonk, *Chem. Mater.* **14**, 508 (2002).
- 2002VER S. P. Verevkin, *J. Chem. Eng. Data* **47**, 1071 (2002).
- 2002VER/HEI S. P. Verevkin and A. Heintz, *J. Chem. Soc., Perkin Trans. 2* **2**, 728 (2002).
- 2002VER/KUM S. P. Verevkin, M. Kummerlin, E. Hickl, H.-D. Beckhaus, C. Ruchardt, S. I. Kozhushkov, R. Haag, R. Boese, J. Benet-Buchholz, K. Nordhoff, and A. de Meijere, *Eur. J. Org. Chem.* **2280** (2002).
- 2002VER2 S. P. Verevkin, *J. Chem. Thermodyn.* **34**, 263 (2002).
- 2002WIL/VON G. M. Wilson, D. M. Von Niederhausern, and N. F. Giles, *J. Chem. Eng. Data* **47**, 761 (2002).
- 2002ZHO/PEN C.-R. Zhou, G.-S. Peng, Y.-D. Zhang, D. G. Jiang, F. Wang, J.-J. Li, and X.-Q. Shen, *Gaoxiao Huaxue Gongcheng Xuebao* **16**, 237 (2002); *Chem. Abstr.* **137**, 316567d.
- 2002ZHO/ZHA D. Zhou, G. G. Z. Zhang, D. Law, D. J. W. Grant, and E. A. Schmitt, *J. Pharm. Sci.* **91**, 1863 (2002).
- 2003ARA/STO A. A. S. Araujo, S. Storpirtis, L. P. Mercuri, F. M. S. Carvalho, M. dos Santos Filho, and J. R. Matos, *Int. J. Pharm.* **260**, 303 (2003).
- 2003BRI/BOU A.-J. Briard, M. Bouroukba, D. Petitjean, N. Hubert, and M. Dirand, *J. Chem. Eng. Data* **48**, 497 (2003).
- 2003CAC/BAU C. Cacula, A. Baudot, M. L. Duarte, A. M. Matos-Beja, M. R. Silva, J. A. Paixo, and R. Fausto, *J. Mol. Struct.* **649**, 143 (2003).
- 2003CAR/DES L. Carpentier, S. Desprex and M. Descamps, *J. Therm Anal. Calorim.* **73**, 577 (2003).
- 2003CEN/RUZ M. Censky, K. Ruzicka, and M. Zabransky, *Thermochim. Acta* **408**, 45 (2003).
- 2003CHA/BLO M. B. Charapennikau, A. V. Blokhin, A. G. Kabo, and G. J. Kabo, *J. Chem. Thermodyn.* **35**, 145 (2003).
- 2003CHA/BLO2 M. B. Charapennikau, A. V. Blokhin, G. J. Kabo, V. M. Sevruk, and A. P. Krasulin, *Thermochim. Acta* **405**, 85 (2003).
- 2003CHA/GUP C. Chawla, P. Gupta, R. Thilagavathi, A. K. Chakraborti, and A. K. Bansal, *Eur. J. Pharm. Sci.* **20**, 305 (2003).
- 2003CHI/ACR J. S. Chickos and W. E. Acree, Jr., *Thermochim. Acta* **395**, 59 (2003).
- 2003CHI/ACR2 J. S. Chickos and W. E. Acree, Jr., *J. Phys. Chem. Ref. Data* **32**, 519 (2003).
- 2003CHI/KN I R. D. Chirico, S. E. Knipmeyer, and W. V. Steele, *J. Chem. Thermodyn.* **35**, 1059 (2003).
- 2003CLO/JAN K. Clou, J. F. Janssens, N. Blaton, A. T. H. Lenstra, and H. O. Desseyn, *Thermochim. Acta* **298**, 47 (2003).
- 2003DEM/KOZ A. de Meijere, S. I. Kozhushkov, K. Rauch, H. Schill, S. P. Verevkin, M. Kuemmerlin, H.-D. Beckhaus, C. Rchardt, and D. S. Yufit, *J. Am. Chem. Soc.* **125**, 15110 (2003).
- 2003DER/MIC S. O. Derawi, M. L. Michelsen, G. M. Kontogeorgis, and E. H. Stenby, *Fluid Phase Equilib.* **209**, 163 (2003).
- 2003DIK/FRE V. V. Diky, M. Frenkel, and L. S. Karpushenkava, *Thermochim. Acta* **408**, 115 (2003).
- 2003EUS/LOP E. M. Eusebio, A. J. Lopes Jesus, M. S. C. Cruz, M. L. P. Leitao, and R. J. Simoes, *J. Chem. Thermodyn.* **35**, 123 (2003).
- 2003EWI/OCH M. B. Ewing and J. C. S. Ochoa, *Fluid Phase Equilib.* **210**, 277 (2003).
- 2003FUL/RUZ M. Fulem, K. Ruzicka, E. Hulicius, T. Simecek, K. Melichar, J. Pangrac, S. A. Rushworth, and L. M. Smith, *J. Cryst. Growth* **248**, 99 (2003).
- 2003GRZ/LAN A. L. Grzesiak, M. Lang, K. Kim, and A. J. Matzger, *J. Pharm. Sci.* **92**, 2260 (2003).
- 2003HIN/RAF D. Hinks, M. I. Rafiq, D. M. Price, G. A. Montero, and B. Smith, *Color. Technol.* **119**, 89 (2003).
- 2003HUA C.-Y. Huang, M.S. thesis, Chinese Culture University, 2003.
- 2003KAL/PAU J. Kaloustian, A.-M. Pauli, P. Lechene de la Porte, H. Lafont, and H. Portugal, *J. Therm Anal. Calorim.* **71**, 341 (2003).

- 2003KAR/KOZ N. V. Karyakin, M. S. Kozlova, M. S. Sheiman, G. P. Kamelova, and V. N. Larina, *Zh. Fiz. Khim.* **77**, 1375 (2003).
- 2003KON/TAN L.-G. Kong, Z.-C. Tan, J. Xu, S.-H. Meng, and X.-H. Bao, *J. Chem. Thermodyn.* **35**, 1897 (2003).
- 2003KOZ/KAR M. S. Kozlova, N. V. Karyakin, M. S. Sheiman, G. P. Kamelova, V. N. Larina, and A. S. Smirnov, *Russ. J. Phys. Chem.* **77**, 26 (2003).
- 2003LEB/VAN N. Lebrun, J. C. Van Miltenburg, D. Bustin, and M. Descamps, *Phase Transitions* **76**, 841 (2003).
- 2003LEG/FEU B. Legendre, Y. Feutelais, and G. Defossemont, *Thermochim. Acta* **400**, 213 (2003).
- 2003MAR/AVI F. Martinez, C. M. Avila, and A. Gomez, *J. Braz. Chem. Soc.* **4**, 803 (2003).
- 2003MAT/MIR M. A. R. Matos, M. S. Miranda, and V. M. F. Morais, *J. Chem. Eng. Data* **48**, 669 (2003).
- 2003MAT/MIR2 M. A. R. Matos, M. S. Miranda, V. M. F. Morais, and J. F. Liebman, *Org. Biomol. Chem.* **1**, 2566 (2003).
- 2003MAY/TOR M. J. Mayoral, M. C. Torralba, M. Cano, J. A. Campo, and J. V. Heras, *Inorg. Chem.* **6**, 626 (2003).
- 2003MOR/MIR V. M. F. Morais, M. S. Miranda, and M. A. R. Matos, *Org. Biomol. Chem.* **1**, 4329 (2003).
- 2003NIK/MAR R. Nikolic, M. Marinovic-Cincovic, S. Gadzuric, and I. Zsigrai, *Sol. Energy Mater. Sol. Cells* **79**, 285 (2003).
- 2003NOK/BOL A. Nokhodchi, N. Bolourtchian, and R. Dinarvand, *Int. J. Pharm.* **250**, 85 (2003).
- 2003ORT/ESP J. Ortega, F. Espiau, and M. Postigo, *J. Chem. Eng. Data* **48**, 916 (2003).
- 2003ORT/ESP2 J. Ortega, F. Espiau, J. Togo, J. Canosa, and A. Rodriguez, *J. Chem. Eng. Data* **48**, 1183 (2003).
- 2003PAG/BEN S. Pagola, A. Benavente, A. Raschi, E. Romano, M. A. A. Molina, and P. W. Stephens, *AAPS PharmSciTech* **2003**, 5 (2003).
- 2003PAJ/ROS J. Pajak, M. Rospenk, Z. Galewski, and L. Sobczyk, *J. Mol. Liq.* **105**, 53 (2003).
- 2003PEN/RIB R. Pena, J. P. Ribet, J. L. Mural, L. Valat, F. Lacouloncha, and A. Chauvet, *Thermochim. Acta* **408**, 85 (2003).
- 2003PER/BAU G. L. Perlovich and A. Bauer-Brandl, *Pharm. Res.* **20**, 471 (2003). [Note: the authors calculated the ideal mole fraction solubility of $x=0.0311$ at 298.2 K using their previously published enthalpy of fusion data, which was presumably reported previously (Ref. [2001PER/BAU]). The referenced paper does not contain the experimental enthalpy of fusion data. The enthalpy of fusion value for acetylsalicylic acid was calculated using the thermodynamic relationship: $\ln x = -\frac{\Delta_{\text{fus}}H_m(T)(T_{\text{fus}} - 298.2)}{(8.314 \times 298.2 \times T_{\text{fus}})}$.]
- 2003PER/CON S. Perisanu, I. Contineanu, M. D. Banciu, J. F. Liebman, B. S. Farivar, M. A. Mullan, J. S. Chickos, N. Rath, and D. M. Hillesheim, *Thermochim. Acta* **400**, 109 (2003).
- 2003PER/KUR G. L. Perlovich, S. V. Kurkov, and A. Bauer-Brandl, *Eur. J. Pharm. Sci.* **19**, 423 (2003).
- 2003PERKUR2 G. L. Perlovich, S. V. Kurkov, A. N. Kinchin, and A. Bauer-Brandl, *J. Pharm. Sci.* **92**, 2502 (2003).
- 2003RIB/LIM M. A. V. Ribeiro da Silva, L. M. S. S. Lima, L. M. P. F. Amaral, A. I. M. C. L. Ferreira, and J. R. B. Gomes, *J. Chem. Thermodyn.* **35**, 1343 (2003).
- 2003RIB/SAN M. D. M. C. Ribeiro da Silva, L. M. N. B. F. Santos, A. L. R. Silva, O. Fernandez, and W. E. Acree, Jr., *J. Chem. Thermodyn.* **35**, 1093 (2003).
- 2003ROJ/ORO A. Rojas and E. Orozco, *Thermochim. Acta* **405**, 93 (2003).
- 2003ROU/JIM M. V. Roux, P. Jimenez, J. Z. Davalos, M. Temprado, and J. F. Liebman, *J. Chem. Thermodyn.* **35**, 803 (2003).
- 2003ROU/JIM2 M. V. Roux, P. Jimenez, A. Vacas, F. H. Cano, M. del Carmen Aprea-Rojas, and F. Ros, *Eur. J. Org. Chem.* **2003**, 2084 (2003).
- 2003ROU/TEM M. V. Roux, M. Temprado, P. Jimenez, J. Z. Davalos, C. Foces-Foces, M. V. Garcia, and M. I. Redondo, *Thermochim. Acta* **404**, 235 (2003).
- 2003ROU/TEM2 M. V. Roux, M. Temprado, P. Jimenez, R. Guzman, E. Juaristi, and J. S. Chickos, *Thermochim. Acta* **406**, 9 (2003).
- 2003ROU/TEM3 M. V. Roux, M. Temprado, P. Jimenez, J. Perez-Parajon, and R. Notario, *J. Phys. Chem. A* **107**, 11460 (2003).
- 2003RUT/SAL M. A. Rute, J. Salud, P. Negrier, D. O. Lopez, J. Li. Tamarit, R. Puertas, M. Barrio, and D. Mondieig, *J. Phys. Chem. B* **107**, 5914 (2003).
- 2003SHA/KAN B. L. Sharma, R. Kant, R. Sharma, and S. Tandon, *Mater. Chem. Phys.* **82**, 216 (2003).
- 2003SIE/WEB A. R. Siedle, R. J. Webb, F. E. Behr, R. A. Newmark, D. A. Weil, K. Erikson, R. Naujok, M. Bronstrom, M. Mueller, S.-H. Chou, and V. G. Young, Jr., *Inorg. Chem.* **42**, 932 (2003).
- 2003SMI/LEB N. N. Smirnova and B. V. Lebedev, *Russ. J. Gen. Chem.* **73**, 247 (2003).
- 2003STA/BAI P. A. Stabnikov, I. A. Baidina, S. V. Sysoev, N. S. Vanina, N. B. Morozova, and I. K. Igumenov, *J. Struct. Chem.* **44**, 1054 (2003).
- 2003STE/CHI W. V. Steele, R. D. Chirico, A. B. Cowell, A. Nguyen, and S. E. Knipmeyer, *J. Chem. Thermodyn.* **35**, 1253 (2003).
- 2003STO/KRZ P. Storoniak, K. Krzyminski, A. Bouzyk, E. P. Koval'chuk, and J. Blaziejowski, *J. Therm. Anal. Calorim.* **74**, 443 (2003).
- 2003SUC/RAJ M. Suceška, M. Rajic, S. Matecic-Musanic, S. Zeman, and Z. Jalovy, *J. Therm. Anal. Calorim.* **74**, 853 (2003).
- 2003SUN/SON X.-H. Sun, J.-R. Song, Z.-C. Tan, Y.-Y. Di, H.-X. Ma, M.-H. Wang, and L.-X. Sun, *Huaxue Xuebao* **61**, 1897 (2003).
- 2003TEO/WIL M. Teodorescu, M. Wilken, and R. Wittig, *Fluid Phase Equilib.* **204**, 267 (2003).
- 2003TOZ/INA K.-I. Tozaki, H. Inaba, H. Hayashi, C. Quan, N. Nemoto, and T. Kimura, *Thermochim. Acta* **397**, 155 (2003).
- 2003TRE/KAS T. Treszczanowicz, T. Kasprzycka-Guttman, and A. Treszcyanowicz, *J. Chem. Eng. Data* **48**, 1517 (2003) (secondary reference).
- 2003VAN/GAB J. C. van Miltenburg, H. Gabrielova, and K. Ruzicka, *J. Chem. Eng. Data* **48**, 1323 (2003).
- 2003VAN/VAN J. C. Van Miltenburg, G. J. K. van den Berg, and M. Ramirez, *J. Chem. Eng. Data* **48**, 36 (2003).
- 2003VAN/VAN2 J. C. van Miltenburg, P. J. van Ekeren, F. G. Gandolfo, and E. Flter, *J. Chem. Eng. Data* **48**, 1245 (2003).
- 2003VER S. P. Verevkin, *J. Chem. Thermodyn.* **35**, 1237 (2003).
- 2003VER/KRA S. P. Verevkin, E. L. Krasnykh, T. V. Vasil'tsova, and A. Heintz, *J. Chem. Eng. Data* **48**, 591 (2003).
- 2003VER/KRA2 S. P. Verevkin, E. L. Krasnykh, T. V. Vasil'tsova, B. Koutek, J. Doub'sky, and A. Heintz, *Fluid Phase Equilib.* **206**, 331 (2003).
- 2003VER/ROU S. P. Verevkin, M. V. Roux, R. Notario, D. E. Demasters, and J. F. Liebman, *Mol. Phys.* **21**, 3231 (2003).
- 2003VER/SCH S. P. Verevkin and C. Schick, *Fluid Phase Equilib.* **211**, 161 (2003).
- 2003VER2 S. P. Verevkin, *Phys. Chem. Chem. Phys.* **5**, 710 (2003).
- 2003WAN/LI C. Wang, H. Li, L. Ma, and S. Han, *J. Chem. Thermodyn.* **35**, 131 (2003).
- 2003WIL/VAN H. T. A. Wilderbeek, M. G. M. Van der Meer, M. A. G. Jansen, L. Nelissen, H. R. Fisher, J. J. G. S. Van Es, C. W. M. Bastiaansen, J. Lub, and D. J. Broer, *Liq. Cryst.* **30**, 93 (2003).
- 2003YAM/HAY O. Yamamura, M. Hayashi, T. Matsuo, and P. Lunkenheimer, *J. Chem. Phys.* **119**, 4775 (2003).
- 2003YU/TAN H.-G. Yu, Z.-C. Tan, Y. Liu, X.-Z. Lan, F. Xu, X.-M. Huang, and L.-X. Sun, *Thermochim. Acta* **404**, 89 (2003).
- 2003YU/TAN2 P. Yu, Z. C. Tan, S. H. Meng, S. W. Lu, X. Z. Lan, L. X. Sun, F. Xu, T. Zhang, and S. X. Hu, *J. Therm. Anal. Calorim.* **74**, 867 (2003).
- 2003ZAI/KAB Dz. Zaitsau, G. J. Kabo, A. A. Kozyro, and V. M. Sevruk, *Thermochim. Acta* **406**, 17 (2003).
- 2003ZAI/VER D. H. Zaitsau, S. P. Verevkin, Y. U. Paulechka, G. J. Kabo, and V. M. Sevuk, *J. Chem. Eng. Data* **48**, 1393 (2003).
- 2003ZEL/TIT L. N. Zelenina, V. A. Titov, T. P. Chusov, Yu. G. Stenin, and A. A. Titov, *J. Chem. Thermodyn.* **35**, 1601 (2003).
- 2003ZHO/ZHA C. Zhou, Y. Zhang, and D. Jiang, *Chin. J. Chem. Eng.* **11**, 598 (2003).
- 2004AHM/BAR A. Ahmed, B. W. Barry, A. C. Williams, and A. F. Davis,

- J. Pharm. Biomed. Anal.* **34**, 945 (2004).
- 2004AHM/GIE S. Ahmad, R. Giesen, and K. Lucas, *J. Chem. Eng. Data* **49**, 826 (2004).
- 2004AMB/MAH A. A. Ambike, K. R. Mahadik, and A. Paradkar, *Int. J. Pharm.* **282**, 151 (2004).
- 2004ANT/GAL M. Antosik, M. Galka, and S. K. Malanowski, *J. Chem. Eng. Data* **49**, 11 (2004).
- 2004BAS/CHI A. Bashir-Hashemi, J. S. Chickos, W. Hanshaw, H. Zhao, B. S. Farivar, and J. F. Liebman, *Thermochim. Acta* **424**, 91 (2004).
- 2004BEN/KHI M. Benkhennouf, K. Khimeche, and A. Dahmani, *J. Phys. Chem. A* **108**, 7 (2004).
- 2004BYK/LEB T. A. Bykova and B. V. Lebedev, *Russ. J. Gen. Chem.* **74**, 250 (2004).
- 2004BYK/SMI T. A. Bykova, N. N. Smirnova, L. V. Nikishchenkova, L. V. Belova, and G. P. Novikova, *Russ. J. Phys. Chem.* **78**, 1712 (2004).
- 2004CAI/BOU M. R. Caira, S. A. Bourne, and C. L. Oliver, *J. Therm. Anal. Calorim.* **77**, 597 (2004).
- 2004CAI/FOP M. R. Caira, A. Foppoli, M. E. Sangalli, L. Zema, and F. Giordano, *J. Therm. Anal. Calorim.* **77**, 653 (2004).
- 2004CER/PER L. Cerioni, S. Perez, and A. Wolfenson, *J. Phys. Chem. Solids* **65**, 1133 (2004).
- 2004CHE/CLE Y. Chernyak and J. H. Clements, *J. Chem. Eng. Data* **49**, 1180 (2004).
- 2004CHE/MA M. Chen and P. Ma, *J. Chem. Eng. Data* **49**, 756 (2004) (secondary reference).
- 2004CHI/HAN J. S. Chickos and W. J. Hanshaw, *J. Chem. Eng. Data* **49**, 77 (2004).
- 2004CHI/HAN2 J. S. Chickos and W. Hanshaw, *J. Chem. Eng. Data* **49**, 620 (2004).
- 2004CHI/STE R. D. Chirico and W. V. Steele, *J. Chem. Thermodyn.* **36**, 633 (2004).
- 2004CHI/ZHA J. S. Chickos, H. Zhao, and G. Nichols, *Thermochim. Acta* **424**, 111 (2004).
- 2004CHY/FRA K. Chylinski, Z. Frasz, and S. K. Malanowski, *J. Chem. Eng. Data* **49**, 2 (2004).
- 2004CHY/FRA2 C. Chylinski, Z. Frasz, and S. K. Malanowski, *J. Chem. Eng. Data* **49**, 18 (2004).
- 2004CLI/RAM C. L. Clifford, D. Ramjugernath, and J. D. Raal, *J. Chem. Eng. Data* **49**, 1189 (2004).
- 2004COS/SCH S. Coste, J.-M. Schneider, M.-N. Petit, and G. Coquerel, *Cryst. Growth Des.* **4**, 1237 (2004).
- 2004COV/MOK M. Covarrubias-Cervantes, I. Mokbel, A. Champion, J. Jose, and A. Voilley, *Food Chem.* **85**, 221 (2004).
- 2004DEF/RAN G. Defossemont, S. L. Randzio, and B. Legendre, *Cryst. Growth Des.* **4**, 1169 (2004).
- 2004DEV/VAN M. M. de Villiers, C. M. Van Eeden, W. Liebenberg, M. Song, W. M. Kolling, and M. R. Caira, *J. Agric. Food Chem.* **52**, 7362 (2004).
- 2004DI/TAN Y.-Y. Di, Z.-C. Tan, X.-H. Sun, M.-H. Wang, F. Xu, Y.-F. Liu, L.-X. Sun, and H.-T. Zhang, *J. Chem. Thermodyn.* **36**, 79 (2004).
- 2004DIA/CAC A. M. A. Dias, A. I. Caco, J. A. P. Coutinho, L. M. N. B. F. Santos, M. M. Pineiro, L. F. Vega, M. F. Costa Gomes, and I. M. Manucho, *Cryst. Growth Des.* **4**, 39 (2004).
- 2004DIN M. S. Ding, *J. Chem. Eng. Data* **49**, 276 (2004).
- 2004DOR/YAN D. V. Dorofeeva, V. S. Yangman, R. M. Varushehenko, and A. I. Druzhinina, *Int. J. Thermophys.* **25**, 1097 (2004).
- 2004FLO/AMA H. Flores, and P. Amador, *J. Chem. Thermodyn.* **36**, 1019 (2004).
- 2004FOP/SAN A. Foppoli, M. E. Sangalli, A. Maroni, A. Gazzaniga, M. R. Caira, and F. Giordano, *J. Pharm. Sci.* **93**, 521 (2004).
- 2004FRI/KAP N. Fridman, M. Kapon, Y. Sheynin, and M. Kaftory, *Acta Crystallogr., Sect. B: Struct. Sci.* **60**, 97 (2004).
- 2004FUJ/TOD H. Fujimori, A. Todoroki, T. Asaji, and M. Oguni, *AIP Conf. Proc.* **708**, 685 (2004).
- 2004FUL/RUZ M. Fulem, K. Ruzicka, V. Ruzicka, E. Hulicius, T. Simecek, J. Pangrac, S. A. Rushworth, and L. M. Smith, *J. Cryst. Growth* **272**, 42 (2004).
- 2004FUL/RUZ2 M. Fulem, K. Ruzicka, V. Ruzicka, T. Simecek, E. Hulicius, and J. Pangrac, *J. Cryst. Growth* **264**, 192 (2004).
- 2004GEN/PEI Z. Geng-qu, C. Pei, H. Rong-zu, L. Yang, Z. Zhi-zhong, Z. Yan-shui, Y. Xu-wu, G. Yin, G. Sheng-li, and S. Qu-zhen, *J. Hazard. Mater.* **113**, 67 (2004).
- 2004GON/KOS T. Gondova and D. Koscakova, *J. Therm. Anal. Calorim.* **76**, 133 (2004).
- 2004GRA/RAS S. Gracin and A. C. Rasmuson, *Cryst. Growth Des.* **4**, 1013 (2004).
- 2004GRI/WEI U. J. Griesser, D. Weigand, J. M. Rollinger, M. Haddow, and E. Gstrein, *J. Therm. Anal. Calorim.* **77**, 511 (2004).
- 2004HAM/FEU N. Hamdi, Y. Feutelais, N. Yagoubi, D. de Girolamo, and B. Legendre, *J. Therm. Anal. Calorim.* **76**, 985 (2004).
- 2004HAM/HAMD C. Hammell, M. Hamad, H. K. Vaddi, P. A. Crooks, and A. L. Stinchcomb, *Telemet. J.* **97**, 283 (2004).
- 2004HAN/BEY A. R. Hansen and K. D. Beyer, *J. Phys. Chem. A* **108**, 3457 (2004).
- 2004HOR/FIS S. Horstman, K. Fisher, and J. Gmehling, *J. Chem. Eng. Data* **49**, 1494 (2004).
- 2004KAO/SIE C.-P. C. Kao, A. C. Sievert, M. Schuller, and J. F. Sturgis, *J. Chem. Eng. Data* **49**, 532 (2004).
- 2004KIM/KES Y. Kim, K. I. Keskinen, and J. Aittamaa, *J. Chem. Eng. Data* **49**, 1273 (2004).
- 2004KON/TAN L. G. Kong, Z. C. Tan, W. Zhang, F. Xu, M. H. Wang, X. H. Bao, T. Zhang, and L. X. Sun, *Chem. Pap.* **58**, 295 (2004).
- 2004KON/TAN2 L.-G. Kong, Z.-C. Tan, J.-T. Mei, L.-X. Sun, and X.-H. Bao, *Thermochim. Acta* **414**, 131 (2004).
- 2004KUL/MAR T. G. Kulagina, A. V. Markin, T. A. Bykova, N. N. Smirnova, L. A. Smirnova, and V. A. Barachevskii, *Russ. J. Phys. Chem.* **78**, 139 (2004).
- 2004KUR/MAE H. Kuramochi, K. Maeda, and K. Kawamoto, *Envir. Toxicol. Chem.* **23**, 1386 (2004).
- 2004KUR/MAE2 H. Kuramochi, K. Maeda, and K. Kawamoto, *J. Chem. Eng. Data* **49**, 720 (2004).
- 2004LAU/HIL K. H. Lau, D. L. Hildenbrand, S. Crouch-Baker, and A. Sanjurjo, *J. Chem. Eng. Data* **49**, 544 (2004).
- 2004LEE/SU M.-J. Lee, C.-C. Su, and H.-M. Lin, *J. Chem. Eng. Data* **49**, 588 (2004).
- 2004LEG/FEU B. Legendre and Y. Feutelais, *J. Therm. Anal. Calorim.* **76**, 255 (2004).
- 2004LI/SHI X.-W. Li, E. Shibata, E. Kasai, and T. Nakamura, *Envir. Toxicol. Chem.* **23**, 348 (2004).
- 2004LU/TAN X.-C. Lu, Z.-C. Tan, Y.-Y. Di, Q. Shi, L.-X. Shi, L.-X. Sun, and T. Zhang, *J. Chem. Thermodyn.* **36**, 787 (2004).
- 2004LUB/MAL M. Lubomska and S. K. Malanowski, *J. Chem. Eng. Data* **49**, 1488 (2004).
- 2004MA/LIU X.-B. Ma, X.-G. Liu, Z.-H. Li, and G.-H. Xu, *Fluid Phase Equilib.* **221**, 51 (2004).
- 2004MAN/ROH T. Manifar, S. Rohani, and M. Saban, *Ind. Eng. Chem. Res.* **44**, 970 (2004).
- 2004MAR/KAI E. Marti, E. Kaisersberger, and W.-D. Emmerich, *J. Therm. Anal. Calorim.* **77**, 905 (2004).
- 2004MAS/NAK M. Massalska-Arodz, T. Nakamoto, T. Wasiutynski, J. Mayer, J. Krawczyk, and M. Sorai, *J. Chem. Thermodyn.* **36**, 877 (2004).
- 2004MAT/MIR M. A. R. Matos, M. S. Miranda, V. M. F. Morais, and J. F. Liebman, *Org. Biomol. Chem.* **2**, 1647 (2004).
- 2004MAT/MIR2 M. A. R. Matos, M. S. Miranda, D. V. S. S. Martins, N. A. B. Pinto, V. M. F. Morais, and J. F. Liebman, *Mar. Fish. Rev.* **2**, 1353 (2004).
- 2004MAT/MIR3 M. A. R. Matos, M. S. Miranda, V. M. F. Morais, and J. F. Liebman, *Eur. J. Org. Chem.* **3340** (2004).
- 2004MAT/MON M. A. R. Matos, M. J. S. Monte, C. C. S. Sousa, A. R. R. P. Almeida, and V. M. F. Morais, *Mar. Fish. Rev.* **2**, 908 (2004).
- 2004MIR/MOR M. S. Miranda, V. M. F. Morais, and M. A. R. Matos, *J. Chem. Thermodyn.* **36**, 431 (2004).
- 2004MON/ALM M. J. S. Monte, A. R. R. P. Almeida, and M. A. V. Ribeiro da Silva, *J. Chem. Thermodyn.* **36**, 385 (2004).
- 2004MON/RAJ D. Mondieig, F. Rajabalee, V. Metivaud, H. A. J. Oonk, and M. A. Cuevas-Diarte, *Chem. Mater.* **16**, 786 (2004).
- 2004MOR/MAT V. M. F. Morais, M. A. R. Matos, M. S. Miranda, and J. F. Liebman, *Mol. Phys.* **102**, 525 (2004).

- 2004MOR/MIR V. M. F. Morais, M. S. Miranda, and M. A. R. Matos, *J. Chem. Thermodyn.* **36**, 377 (2004).
- 2004NAS/ZIM K. Nasirzadeh, D. Zimin, R. Neueder, and W. Kunz, *J. Chem. Eng. Data* **49**, 607 (2004).
- 2004NUN/EUS S. C. C. Nunes, M. E. Eusebio, M. L. P. Leitão, and J. S. Redinha, *Int. J. Pharm.* **285**, 13 (2004).
- 2004PAU/ZAI Y. U. Paulechka, Dz. H. Zaitsau, and G. J. Kabo, *J. Mol. Liq.* **115**, 105 (2004).
- 2004PER/KUR G. L. Perlovich, S. V. Kurkov, A. N. Kinchin, and A. Bauer-Brandl, *Eur. J. Pharm. Biopharm.* **57**, 411 (2004).
- 2004PIA/SUG X. Piao, Y. Sugihara, and J. Nakayama, *Heteroat. Chem.* **15**, 424 (2004).
- 2004PIL/HAM O. Pillai, M. O. Hamad, P. A. Crooks, and A. L. Stinchcomb, *Pharm. Res.* **21**, 1146 (2004). (Enthalpies of fusion were calculated from the tabulated melting point temperature and thermodynamic activity data given in the paper.)
- 2004PRO/RAS V. M. Profir and A. C. Rasmuson, *Cryst. Growth Des.* **4**, 315 (2004).
- 2004QU/BAI H. Qu, P. Bai, Z. Yang, and G. Yu, *Chin. J. Chem. Eng.* **12**, 294 (2004).
- 2004RAM/COR J. J. M. Ramos, N. T. Correia, and H. P. Diogo, *Phys. Chem. Chem. Phys.* **6**, 793 (2004).
- 2004RAM/DIO K. J. M. Ramos, H. P. Diogo, M. H. Godinho, C. Cryz, and K. Merkel, *J. Phys. Chem. B* **108**, 7955 (2004).
- 2004RIB/GOM M. D. M. C. Ribeiro da Silva, J. R. B. Gomes, J. M. Goncalves, E. A. Sousa, S. Pandey, and W. E. Acree, Jr., *Org. Biomol. Chem.* **2**, 2507 (2004).
- 2004RIB/GOM2 M. D. M. C. Ribeiro da Silva, J. R. B. Gomes, J. M. Goncalves, E. A. Sousa, S. Pandey, and W. E. Acree, Jr., *J. Org. Chem.* **69**, 2785 (2004).
- 2004RIB/GON M. D. M. C. Ribeiro da Silva, J. M. Goncalves, A. R. L. Silva, A. M. R. O. A. Silva, P. C. F. C. Oliveira, and M. A. V. M. A. V. Ribeiro da Silva, *Thermochim. Acta* **420**, 67 (2004).
- 2004RIB/MAT M. D. M. C. Ribeiro da Silva, M. A. R. Matos, M. S. Miranda, V. M. F. Morais, and W. E. Acree, Jr., *J. Chem. Thermodyn.* **36**, 107 (2004).
- 2004RIB/RIB M. A. V. Ribeiro da Silva, M. D. M. C. Ribeiro da Silva, L. M. P. F. Amaral, P. Jimenez, M. V. Roux, J. Z. Davalos, M. Temprado, P. Cabildo, R. M. Claramunt, J. Elguero, O. Mó, and M. Yáñez, *J. Chem. Thermodyn.* **36**, 533 (2004).
- 2004RIB/SAN M. A. V. Ribeiro da Silva, L. M. N. B. F. Santos, B. Schroder, and L. Beyer, *J. Chem. Thermodyn.* **36**, 555 (2004).
- 2004RIB/SAN2 M. A. V. Ribeiro da Silva, L. M. N. B. F. Santos, B. Schroder, F. Dietze, and L. Beyer, *J. Chem. Thermodyn.* **36**, 491 (2004).
- 2004RIB/SAN3 M. A. V. Ribiero da Silva and L. M. N. B. F. Santos, *J. Chem. Thermodyn.* **36**, 447 (2004).
- 2004ROH/RUZ V. Rohac, K. Ruzicka, V. Ruzicka, D. H. Zaitsau, G. J. Kabo, V. Dicky, and K. Aim, *J. Chem. Thermodyn.* **36**, 929 (2004).
- 2004ROJ/FOR A. Rojas-Aguilar, H. Fores-Lara, M. Martinez-Herrera, and F. Ginez-Carbajal, *J. Chem. Thermodyn.* **36**, 453 (2004).
- 2004ROM/BUS S. Romero, P. Bustamante, B. Escalera, M. Cirri, and P. Mura, *J. Therm Anal. Calorim.* **77**, 541 (2004).
- 2004ROM/BUS2S. Romero, P. Bustamante, B. Escalera, P. Mura, and M. Cirri, *J. Pharm. Biomed. Anal.* **35**, 715 (2004).
- 2004ROM/ROC J. J. M. Romos, S. Rocha, and H. P. Diogo, *J. Non-Cryst. Solids* **344**, 119 (2004).
- 2004ROU/TEM M. V. Roux, M. Temprado, R. Notario, S. P. Verevkin, V. N. Emel'yanenko, D. Demasters, and J. F. Liebman, *Mol. Phys.* **102**, 1909 (2004).
- 2004ROU/TEM2M. V. Roux, M. Temprado, P. Jimenez, C. Foces-Foces, M. V. Garcia, and M. I. Redondo, *Thermochim. Acta* **420**, 59 (2004).
- 2004ROU/TEM3M. V. Roux, M. Temprado, P. Jimenez, R. Notario, R. Guzman-Mejia, and E. Juaristi, *J. Org. Chem.* **69**, 1670 (2004).
- 2004RYA/LEB M. V. Ryabkov, B. V. Lebedev, N. N. Smirnova, E. V. Agina, S. Ponomarenko, E. A. Makeev, N. I. Boiko, and V. P. Shibaev, *Russ. J. Phys. Chem.* **78**, 1718 (2004).
- 2004SAB/MAR J. Sabolovi, E. Marak, S. Kotrun, and A. Janekovi, *Inorg. Chem.* **43**, 8479 (2004).
- 2004SAI/MAS K. Saito, M. Massalska-Arodz, S. Ikeuchi, M. Maekawa, J. Sciesinski, E. Sciesinska, J. Mayer, T. Wasiutynski, and M. Sorai, *J. Phys. Chem. B* **108**, 5785 (2004).
- 2004SAN/SCH L. M. N. B. F. Santos, B. Schroder, O. O. P. Fernandez, and M. A. V. Ribeiro da Silva, *Thermochim. Acta* **415**, 15 (2004).
- 2004SAW/MOK T. Sawaya, I. Mokbel, E. Rauzy, J. Saab, C. Berro, and J. Jose, *Fluid Phase Equilib.* **226**, 283 (2004).
- 2004SHA/JAM B. L. Sharma, R. Jamwal, and R. Kant, *Cryst. Res. Technol.* **39**, 454 (2004).
- 2004SMI/LEB N. N. Smirnova, B. V. Lebedev, O. V. Stepanova, A. M. Muzafarov, and E. A. Tatarinova, *Russ. J. Phys. Chem.* **78**, 12 (2004).
- 2004SMI/MAR N. N. Smirnova, A. V. Markin, K. V. Kandeev, H. Hocker, and H. Keul, *Thermochim. Acta* **409**, 55 (2004).
- 2004SOH/SEO Y. T. Sohn and H. O. Seo, *High Temp. Mater. Sci.* **27**, 357 (2004).
- 2004SON/TAN Y.-J. Song, Z. C. Tan, S. W. Lu, and Y. Xue, *J. Therm Anal. Calorim.* **77**, 873 (2004).
- 2004STE/CHI W. V. Steele, R. D. Chirico, S. E. Knipmeyer, and A. Nguyen, *J. Chem. Thermodyn.* **36**, 845 (2004).
- 2004STE/CHI2 W. V. Steele, R. D. Chirico, A. B. Cowell, A. Nguyen, and S. E. Knipmeyer, *J. Chem. Thermodyn.* **36**, 497 (2004).
- 2004STE/STI A. Stefanov, A. Stibor, A. Dominguez-Clarimon, and M. Arndt, *J. Chem. Phys.* **121**, 6935 (2004).
- 2004STE/SUN F. Steyer and K. Sundmacher, *J. Chem. Eng. Data* **49**, 1675 (2004).
- 2004STI/VAL A. L. Stinchcomb, S. Valiveti, D. C. Hammell, and D. R. Ramsey, *J. Pharm. Pharmacol.* **56**, 291 (2004).
- 2004SUN/LIU X.-H. Sung, Y.-F. Liu, Z.-C. Tan, Y.-Y. Di, H.-F. Wang, and M.-H. Wang, *J. Chem. Thermodyn.* **36**, 895 (2004).
- 2004SUN/SON X.-H. Sun, J.-R. Song, Z.-C. Tan, Y.-Y. Di, M.-H. Wang, M.-X. Ma, H.-F. Wang, and L.-X. Sun, *Youji Huazue* **24**, 409 (2004).
- 2004SUN/SON2 X.-H. Sun, J.-R. Song, Z.-C. Tan, Y.-Y. Di, H.-X. Ma, M.-H. Wang, and L.-X. Sun, *Thermochim. Acta* **413**, 261 (2004).
- 2004TAN/LI T. Tan, H. Li, C. Wang, H. Jiang, and S. Han, *Fluid Phase Equilib.* **224**, 279 (2004).
- 2004TYA/BIS O. S. Tyagi, H. S. Bisht, and A. K. Chatterjee, *J. Phys. Chem. B* **108**, 3010 (2004).
- 2004UUS/POK P. Uusi-Kyyny, J.-P. Pokki, Y. Khim, and J. Aittamaa, *J. Chem. Eng. Data* **49**, 251 (2004).
- 2004VAL/KIP S. Valiveti, P. K. Kiptoo, D. C. Hammell, and A. L. Stinchcomb, *Int. J. Pharm.* **278**, 173 (2004).
- 2004VAN/MAL E. V. Van Tonder, T. S. P. Maleka, W. Liebenberg, M. Song, D. E. Wurster, and M. M. de Villiers, *Int. J. Pharm.* **269**, 417 (2004).
- 2004VAN/VAN J. C. Van Miltenburg and G. J. K. Van den Berg, *J. Chem. Eng. Data* **49**, 735 (2004).
- 2004VEC/CAT S. Vecchio, A. Catalani, V. Rossi, and M. Tomassetti, *Thermochim. Acta* **420**, 99 (2004).
- 2004VEN/CAL L. Ventola, T. Calvet, M. A. Cuevas-Diarte, M. Ramirez, H. A. J. Oonk, D. Mondieig, and Ph. Negrier, *Phys. Chem. Chem. Phys.* **6**, 1786 (2004).
- 2004VER S. P. Verevkin, *Fluid Phase Equilib.* **225**, 145 (2004).
- 2004VER/EME S. P. Verevkin and V. E. Emel'yanenko, *J. Phys. Chem. A* **108**, 6575 (2004).
- 2004VER/SCH S. P. Verevkin and C. Schick, *Thermochim. Acta* **415**, 35 (2004).
- 2004VER2 S. P. Verevkin, *Fluid Phase Equilib.* **224**, 23 (2004).
- 2004WAN/TAN M.-H. Wang, Z.-C. Tan, X.-H. Sun, F. Xu, L.-B. Kong, L.-X. Sun, and T. Zhang, *Thermochim. Acta* **411**, 203 (2004).
- 2004WAN/TAN2M.-H. Wang, Z.-C. Tan, X.-H. Sun, F. Xu, Y.-F. Liu, L.-X. Sun, and T. Zhang, *Thermochim. Acta* **414**, 25 (2004).
- 2004WAN/TAN3S. X. Wang, Z. C. Tan, Y. Y. Di, F. Xu, M. H. Wang, L. X. Sun, and T. Zhang, *J. Therm Anal. Calorim.* **76**, 335 (2004).
- 2004WAN/TAN4S.-X. Wang, Z.-C. Tan, Y.-Y. Di, F. Xu, H.-T. Zhang, L.-X.

- Sun, and T. Zhang, *J. Chem. Thermodyn.* **36**, 393 (2004).
- 2004WAN/WAN L.-C. Wang and F. A. Wang, *Fluid Phase Equilib.* **226**, 289 (2004).
- 2004WAN/WIE X. Wang, H. Wiehler, and C. B. Ching, *Chirality* **16**, 220 (2004).
- 2004XIN/TAN J. Xing, Z.-C. Tan, Y.-Y. Di, X.-H. Sun, L.-X. Sun, and Z. Tao, *Huaxue Xuebao* **62**, 2415 (2004).
- 2004XU/SUN F. Xu, L.-X. Sun, Z.-C. Tan, J.-G. Liang, and R.-L. Li, *Thermochim. Acta* **412**, 33 (2004).
- 2004XU/SUN2 F. Xu, L.-X. Sun, Z.-C. Tan, J.-G. Liang, D.-H. Zhou, Y.-Y. Di, X. Z. Lan, and T. Zhang, *Wuji Huaxue Xuebao* **20**, 50 (2004); *Chem. Abstr.* **141**, 42690.
- 2004XUE/WAN B. Xue, Y.-Y. Wang, Z.-C. Tan, S.-W. Lu, and S.-H. Meng, *J. Therm. Anal. Calorim.* **76**, 965 (2004).
- 2004YAM/NEM H. Yamamoto, N. Nemoto, and K. Tashiro, *J. Phys. Chem. B* **108**, 5827 (2004).
- 2004ZEL/CHU L. N. Zelenina, T. P. Chusova, and Yu. G. Stenin, *Zh. Fiz. Khim.* **78**, 598 (2004).
- 2004ZIE/SZT W. Zielenkiewicz and P. Szterner, *J. Chem. Eng. Data* **49**, 1197 (2004).
- 2005ALT/COP M. Altamura, G. Coppini, P. Cuda, P. Dapporto, A. Guerri, A. Guidi, C. Nativi, P. Paoli, and P. Rossi, *J. Mol. Struct.* **749**, 20 (2005).
- 2005BAZ/BLO A. B. Bazyleva, A. V. Blokhin, G. J. Kabo, A. G. Kabo, and Y. U. Paulechka, *J. Chem. Thermodyn.* **37**, 643 (2005).
- 2005BAZ/KAB A. B. Bazyleva, G. J. Kabo, Y. U. Paulechka, D. H. Zait-sau, A. V. Blokhin, and V. M. Sevruk, *Thermochim. Acta* **436**, 56 (2005).
- 2005BEN/AIT H. Ben-makhlouf-Hakem, A. Ait-Kaci, and J. Jose, *Fluid Phase Equilib.* **232**, 189 (2005).
- 2005CHA/ZIE S. Chattopadhyay and P. Ziemann, *Aerosol Sci. Technol.* **39**, 1085 (2005).
- 2005CHE/TAN Y.-P. Chen, M. Tang, and J.-C. Kuo, *Fluid Phase Equilib.* **232**, 182 (2005).
- 2005CHI/STE R. D. Chirico and W. V. Steele, *J. Chem. Eng. Data* **50**, 697 (2005).
- 2005CHI/STE2 R. D. Chirico and W. V. Steele, *J. Chem. Eng. Data* **50**, 1052 (2005).
- 2005CON/CHI J. Contineanu, L. Chivu, and S. Perisanu, *Revista di Chim.* **56**, 719 (2005).
- 2005DES/COR M. Descamps, N. T. Correia, P. Derollez, F. Danede, and F. Capet, *J. Phys. Chem. B* **109**, 16092 (2005).
- 2005DIA/GON A. M. A. Dias, C. M. B. Goncalves, A. I. Caco, L. M. N. B. F. Santos, M. M. Pineiro, L. F. Vega, J. A. P. Coutinho, and I. M. Marrucho, *J. Chem. Eng. Data* **50**, 1328 (2005).
- 2005EME/VER V. N. Emel'yanenko, S. P. Verevkin, B. Koutek, and J. Doubsky, *J. Chem. Thermodyn.* **37**, 73 (2005).
- 2005EME/VER2 V. N. Emel'yanenko and S. P. Verevkin, *J. Phys. Chem. A* **109**, 3960 (2005).
- 2005FAT/KAS A. Fattahi, S. R. Kass, J. F. Liebman, M. A. R. Matos, M. S. Miranda, and V. M. F. Morais, *J. Am. Chem. Soc.* **127**, 6116 (2005).
- 2005FUL/RUZ M. Fulem, K. Ruzicka, V. Ruzicka, T. Sinecsk, E. Hulicius, J. Pangrac, J. Becker, J. Koch, and A. Salzmann, *J. Chem. Eng. Data* **50**, 1613 (2005).
- 2005GOM/AMA J. R. B. Gomes, L. M. P. F. Amaral, and M. A. V. Ribeiro da Silva, *Chem. Phys. Lett.* **406**, 154 (2005).
- 2005HAS/TAJ M. Hashimoto, T. Tajima, K. Eda, K. Yamamura, and T. Okazaki, *J. Mol. Struct.* **734**, 23 (2005).
- 2005HOS/GRY M. Hoskovec, D. Grygarova, J. Cvacka, L. Strinz, J. Zima, S. P. Verevkin, and B. Koutek, *J. Chromatogr. A* **1083**, 161 (2005).
- 2005HOS/NAG S. M. A. Hosseini, H. Naghibi, M. H. Peikar, and E. Jamalizadeh, *Asian J. Chem.* **18**, 351 (2005).
- 2005HUA/SIM D. Huang, S. L. Simon, and G. B. McKenna, *J. Chem. Phys.* **122**, 084907 (2005).
- 2005HUA/TAN C.-Y. Huang, M. Tang, and Y.-P. Chen, *J. Chem. Eng. Data* **50**, 40 (2005). (Note: numerical values of the enthalpy of fusion obtained through correspondence with Y.-P. Chen.)
- 2005IKE/YAM K. Ikedou, H. Yamamoto, H. Nagashima, N. Nemoto, and K. Tashiro, *J. Phys. Chem. B* **109**, 10668 (2005).
- 2005KAP/SLO S. Kapteina, K. Slowik, S. P. Verevkin, and A. Heintz, *J. Chem. Eng. Data* **50**, 398 (2005).
- 2005KLO/BRO M. G. Klous, G. M. Bronner, B. Nuijen, J. M. van Ree, and J. H. Beijnen, *J. Pharm. Biomed. Anal.* **39**, 944 (2005).
- 2005LIZ/ZAB E. Lizarraga, C. Zabaleta, and J. A. Palop, *Thermochim. Acta* **427**, 171 (2005).
- 2005LOP/TOM A. J. Lopes Jesus, L. I. N. Tomé, M. E. Eusébio, and J. S. Redinha, *J. Phys. Chem. B* **109**, 18055 (2005).
- 2005LU/TAN X.-C. Lu, Z.-C. Tan, Q. Shi, H.-T. Zhang, L.-X. Sun, and T. Zhang, *J. Chem. Eng. Data* **50**, 932 (2005).
- 2005MAR/AVA G. Marchionni, M. Avataneo, U. De Patta, P. Maccone, and G. Pezzin, *J. Fluorine Chem.* **126**, 465 (2005).
- 2005MAR/ZAT S. A. Markarian, A. L. Zatikyan, V. V. Grigoryan, and G. S. Grigoryan, *J. Chem. Eng. Data* **50**, 23 (2005).
- 2005MAT/MIR M. A. R. Matos, M. S. Miranda, V. M. F. Morais, and J. F. Liebman, *Mol. Phys.* **103**, 221 (2005).
- 2005MAT/MIR2 M. A. R. Matos, M. S. Miranda, N. A. B. Pinto, V. M. F. Morais, N. Dhananjaya, and J. F. Liebman, *Mol. Phys.* **103**, 1885 (2005).
- 2005MAT/MOR M. A. R. Matos, V. M. F. Morais, M. D. M. C. Ribeiro da Silva, M. C. F. Margues, E. A. Sousa, J. P. Castieiras, C. P. Santos, and W. E. Acree, Jr., *J. Chem. Eng. Data* **50**, 1184 (2005).
- 2005MAT/VAN M. Matovic, J. C. van Miltenburg, J. Los, F. G. Gandolfo, and E. Flöter, *J. Chem. Eng. Data* **50**, 1624 (2005).
- 2005MON/MUN J. B. Monton, R. Munoz, M. C. Burguet, and J. de la Torre, *Fluid Phase Equilib.* **227**, 19 (2005).
- 2005MON/SOU M. J. S. Monte and C. A. D. Sousa, *J. Chem. Eng. Data* **50**, 2101 (2005).
- 2005MUN/MON R. Munoz, J. B. Monton, M. C. Burguet, and J. de la Torre, *Fluid Phase Equilib.* **232**, 62 (2005).
- 2005NAK/SHI K. Nakajoh, E. Shibata, T. Todoroki, A. Ohara, K. Nishizawa, and T. Nakamura, *Envir. Toxicol. Chem.* **24**, 1602 (2005).
- 2005NAS/NEU K. Nasirzadeh, R. Neueder, and W. Kunz, *J. Chem. Eng. Data* **50**, 26 (2005).
- 2005NIS/WAT Y. Nishikawa, T. Watanabe, H. Yoshida, and M. Ikeda, *Thermochim. Acta* **431**, 81 (2005).
- 2005PEN/MUR S. Pena-Tejedor, R. Murga, M. T. Sanz, and S. Beltran, *J. Chem. Eng. Data* **50**, 197 (2005).
- 2005PER/BAN S. Perisanu, M. D. Banciu, I. Contineanu, A. Neascu, and L. Chivu, *Rev. Chim. (Bucuresti)* **56**, 813 (2005).
- 2005PER/ROD G. L. Perlovich, S. V. Rodionov, and A. Bauer-Brandl, *Eur. J. Pharm. Sci.* **24**, 25 (2005).
- 2005PIN/DIO S. S. Pinto, H. P. Diogo, R. C. Guedes, B. J. C. Cabral, M. E. Minas de Piedade, and J. A. M. Simoes, *J. Phys. Chem. A* **109**, 9700 (2005).
- 2005RES/GON J. M. Resa, C. Gonzalez, S. Ortiz de Landaluce, and J. M. Goenaga, *J. Chem. Eng. Data* **50**, 319 (2005).
- 2005RIB/FON M. A. V. Ribeiro da Silva, J. M. S. Fonseca, R. P. B. M. Carvalho, and M. J. S. Monte, *J. Chem. Thermodyn.* **37**, 271 (2005).
- 2005RIB/GOM M. A. V. Ribeiro da Silva, J. R. B. Gomes, and A. I. M. C. L. Ferreira, *J. Phys. Chem. B* **109**, 13356 (2005).
- 2005RIB/MAT M. A. V. Ribeiro da Silva, M. A. R. Matos, and L. M. P. F. Amaral, *J. Chem. Thermodyn.* **37**, 1312 (2005).
- 2005RIB/MIR M. D. M. C. Ribeiro da Silva, M. S. Miranda, C. M. V. Vaz, M. A. R. Matos, and W. E. Acree, Jr., *J. Chem. Thermodyn.* **37**, 49 (2005).
- 2005RIB/RIB M. A. V. Ribeiro da Silva, M. D. M. C. Ribeiro da Silva, L. M. P. F. Amaral, J. Elguero, P. Jimenez, M. V. Roux, J. Z. Davalos, M. Temprado, P. Cabildo, R. M. Claramunt, O. Mo, and M. Yanez, *J. Chem. Thermodyn.* **37**, 1168 (2005).
- 2005ROG/PIS G. N. Roganov, P. N. Pisarev, V. N. Emel'yanenko, and S. P. Verevkin, *J. Chem. Eng. Data* **50**, 1114 (2005).
- 2005ROJ/GIN A. Rojas-Aguilar, F. Ginez-Carbajal, E. Orozco-Guareno, and H. Flores-Sejura, *J. Therm. Anal. Calorim.* **79**, 95 (2005).
- 2005RON/GU H.-R. Rong and H. Gu, *Thermochim. Acta* **428**, 19 (2005).
- 2005ROU/DAV M. V. Roux, J. Z. Davalos, P. Jimenez, R. Notorio, O. Castano, J. S. Chickos, W. Hanshaw, H. Zhao, N. Rath, J. F. Liebman, B. S. Farivar, and A. Bashir-Hashemi, *J. Org.*

- Chem.* **70**, 5461 (2005).
- 2005ROU/TEM M. V. Roux, M. Temprado, and J. S. Chickos, *J. Chem. Thermodyn.* **37**, 941 (2005).
- 2005SAI/KE H. Saitoh, S. Ikeuchi, and K. Saito, *J. Therm Anal. Calorim.* **81**, 511 (2005).
- 2005SAI/MAR C. I. Sainz-Diaz, A. P. Martin-Islan, and J. H. E. Cartwright, *J. Phys. Chem. B* **109**, 18758 (2005).
- 2005SBI/VEC N. Sbirrazzuoli, S. Vecchio, and A. Catalani, *Int. J. Chem. Kinet.* **37**, 74 (2005).
- 2005SCH A. C. Schmidt, *Int. J. Pharm.* **298**, 186 (2005).
- 2005SEM/IGU P. P. Semyannikov, I. K. Igumenov, S. V. Trubin, T. P. Chusova, and Z. I. Semenova, *Thermochim. Acta* **432**, 91 (2005).
- 2005SIK/MOD M. Sikorska-Iwan and B. Modzelewska-Banachewicz, *J. Therm Anal. Calorim.* **81**, 119 (2005).
- 2005SMI/KAN N. N. Smirnova, K. V. Kandehev, and T. A. Bykova, *Russ. J. Phys. Chem.* **79**, 857 (2005).
- 2005SMI/KAN2 N. N. Smirnova, K. V. Kandehev, T. A. Bykova, and T. G. Kulagina, *J. Chem. Thermodyn.* **38**, 376 (2005).
- 2005SMI/KUL N. N. Smirnova, T. G. Kulagina, A. V. Markin, Z. B. Shifrina, and A. L. Rusanov, *Thermochim. Acta* **425**, 39 (2005).
- 2005STE/SUN F. Steyer and K. Sundmacher, *J. Chem. Eng. Data* **50**, 1277 (2005).
- 2005STR/SPO A. Strutynska, A. Sporzynski, J. Serwatowski, and S. P. Verevkin, *Fluid Phase Equilib.* **227**, 283 (2005).
- 2005SUN/LIU X.-H. Sun, Y.-F. Liu, Z.-C. Tan, M.-H. Wang, and J.-Q. Jia, *Chem. Res. Chin. Univ.* **21**, 697 (2005).
- 2005SUN/LIU2 X.-H. Sun, Y.-F. Liu, Z.-C. Tan, Y.-Q. Jia, M.-H. Wang, and Y.-Y. Di, *Chin. J. Chem.* **23**, 23 (2005).
- 2005SUN/LIU3 X.-H. Sun, F.-Y. Liu, Z.-C. Tan, P.-J. Ji, and M.-H. Wang, *Chin. J. Chem.* **23**, 1490 (2005).
- 2005SUN/LIU4 X.-H. Sun, Y.-F. Liu, Z.-C. Tan, Y.-Q. Jia, J.-W. Yang, and M.-H. Wang, *Chin. J. Chem.* **23**, 501 (2005).
- 2005TEM/CHI M. Temprado and J. S. Chickos, *Thermochim. Acta* **435**, 49 (2005).
- 2005TEM/ROU M. Temprado, M. V. Roux, P. Umnahanant, H. Zhao, and J. S. Chickos, *J. Phys. Chem. B* **109**, 12590 (2005).
- 2005TIA/TAN Q.-F. Tian, Z.-C. Tan, Q. Shi, F. Xu, L.-X. Sun, and T. Zhang, *Thermochim. Acta* **430**, 53 (2005).
- 2005TOM/MIZ S. Tomitaka, M. Mizukami, F. Paladi, and M. Oguni, *J. Therm Anal. Calorim.* **81**, 637 (2005).
- 2005TUR/VAC R. Turcotte, M. Vachon, Q. S. M. Kwok, R. Wang, and D. E. G. Jones, *Thermochim. Acta* **433**, 105 (2005).
- 2005VAL/QUI E. Valez, J. Quijano, J. Vaviria, M. V. Roux, P. Jimenez, M. Temprado, G. Martin-Valcarcel, J. Perez-Parajon, and R. Notario, *J. Phys. Chem. A* **109**, 7832 (2005).
- 2005VAN/OON J. C. van Miltenburg and H. A. J. Oonk, *J. Chem. Eng. Data* **50**, 1348 (2005).
- 2005VAN/VAN P. R. van der Linde, J. C. van Miltenburg, G. J. K. van den Berg, and H. A. J. Oonk, *J. Chem. Eng. Data* **50**, 164 (2005).
- 2005VAR/DRU R. M. Varushchenko, A. I. Druzhinina, V. M. Senyavin, and V. S. Sarkisova, *J. Chem. Thermodyn.* **37**, 141 (2005).
- 2005VAS/VER T. V. Vasil'tsova, S. P. Verevkin, E. Bich, A. Heintz, R. Bogel-Lukasik, and U. Domanska, *J. Chem. Eng. Data* **50**, 142 (2005).
- 2005VEC/BRU S. Vecchio and B. Brunetti, *J. Chem. Eng. Data* **50**, 666 (2005). (Note: complete chemical numbers for several of the compounds were obtained through personal correspondence with S. Vecchio.)
- 2005VOG/COH F. G. Vogt, D. E. Cohen, J. D. Bowman, G. P. Spoor, G. E. Zuber, G. A. Trescher, P. C. Dell'orco, L. M. Katrinic, G. W. Debrosse, and R. C. Haltiwanger, *J. Pharm. Sci.* **94**, 651 (2005).
- 2005WAN/TAN M.-H. Wang, Z.-C. Tan, X.-H. Sun, L.-X. Sun, Y.-F. Liu, and T. Zhang, *J. Environ. Eng.* **21**, 573 (2005).
- 2005WAN/TAN2M.-H. Wang, Z.-C. Tan, X.-H. Sun, H.-T. Zhang, B.-P. Liu, L.-X. Sun, and T. Zhang, *J. Chem. Eng. Data* **50**, 270 (2005).
- 2005WAN/TAN3S.-X. Wang, Z.-C. Tan, Q. Shi, Y.-Y. Di, H.-T. Zhang, F. Xu, L.-X. Sun, and T. Zhang, *J. Chem. Thermodyn.* **37**, 349 (2005).
- 2005XU/SUN F. Xu, L.-X. Sun, Z.-C. Tan, R.-L. Li, Q.-F. Tian, and T. Zhang, *J. Environ. Eng.* **21**, 1 (2005).
- 2005XUE/WAN B. Xue, J. Wang, Z.-C. Tan, T. Wu, Z. Zhang, and H. Chen, *Huagong Xuebao* **56**, 570 (2005).
- 2005YAN/DEV W. Yang and M. M. de Villiers, *AAPS J.* **7**, E241 (2005).
- 2005YAN/MAN W. Yang, R. Manek, W. M. Kolling, M. Brits, W. Liebenberg, and M. M. De Villiers, *Supramol. Chem.* **17**, 485 (2005).
- 2005YAO/BAI W.-W. Yao, T.-C. Bai, J.-P. Sun, C.-W. Zhu, J. Hu, and H.-L. Zhang, *Thermochim. Acta* **437**, 17 (2005).
- 2005ZHA/BAI H.-L. Zhang, T.-C. Bai, G.-B. Yan, and J. Hu, *Fluid Phase Equilib.* **238**, 186 (2005).
- 2005ZHA/STA G. I. Zharkova, P. A. Stabnikov, S. A. Sysoev, and I. K. Igumenov, *J. Struct. Chem.* **46**, 320 (2005).
- 2005ZIE/SZT W. Zielenkiewicz and P. Szterner, *J. Chem. Eng. Data* **50**, 1139 (2005).
- 2006ADH/BAS R. Adhiyaman and S. K. Basu, *Int. J. Pharm.* **321**, 27 (2006).
- 2006BAD/DEL E. Badea, G. Della Gata, D. D. Angelo, B. Brunetti, and Z. Reckova, *J. Chem. Thermodyn.* **38**, 1546 (2006).
- 2006BAI/YAN T.-C. Bai, G.-B. Yan, J. Hu, H.-L. Zhang, and C.-G. Huang, *Int. J. Pharm.* **308**, 100 (2006).
- 2006BAR/BLO A. B. Bazyleva, A. V. Blokhin, G. J. Kabo, A. G. Kabo, and V. M. Sevruk, *Thermochim. Acta* **451**, 65 (2006).
- 2006BAR/DAV J. Baran, N. A. Davydova, M. Drozd, and A. Pietraszko, *J. Phys.: Condens. Matter* **18**, 5695 (2006).
- 2006BET/SOR G. Bettinetti, M. Sorrenti, L. Catenacci, F. Ferrari, and S. Rossi, *J. Pharm. Biomed. Anal.* **41**, 1205 (2006).
- 2006BOL/NER M. F. Bolotnikov and Yu. A. Neruchev, *Fluid Phase Equilib.* **243**, 121 (2006).
- 2006BOL/NER2 M. F. Bolotnikov and Yu. A. Neruchev, *Russ. J. Phys. Chem.* **80**, 1191 (2006).
- 2006BOU/BEL G. Boukais-Belaribi, B. F. Belaribi, A. Ait-Kaci, and J. Jose, **248**, 181 (2006).
- 2006BUC/BUE J. H. Buchaman, L. C. Buettner, and D. E. Trevault, *J. Chem. Eng. Data* **51**, 1331 (2006).
- 2006CAM A. G. Camacho, Ph.D. thesis, Universidad Nacional de Tucuman, 2006.
- 2006CAM/MAR A. G. Camacho, A. Mariano, L. Mussari, and M. A. Postigo, *J. Chem. Eng. Data* **51**, 1536 (2006).
- 2006CAP/TRA A. Cappelli, V. Travagli, I. Zanardi, M. Anzini, G. Giorgi, A. Donati, M. Aggravi, M. Casolaro, M. Fresta, E. Paccagnini, F. Makovec, and S. Vomero, *J. Pharm. Sci.* **95**, 2706 (2006).
- 2006CAV/PAN G. Caviglioli, M. Pani, P. Gatti, B. Parodi, S. Cafaggi, and G. Bignardi, *J. Pharm. Sci.* **95**, 2207 (2006).
- 2006CHE/OJA X. Chen, V. Oja, W. G. Chan, and M. R. Hajaligol, *J. Chem. Eng. Data* **51**, 386 (2006).
- 2006COR/LOP P. Correia, C. Lopez, M. E. M. Piedade, J. A. A. Lourenco, and M. L. Serrano, *J. Chem. Eng. Data* **51**, 1306 (2006).
- 2006DAB/SPO A. Dabrowska, A. Sporzynski, and S. P. Verevkin, *Fluid Phase Equilib.* **249**, 115 (2006).
- 2006DEB/MED A. L. F. de Barros, A. Medina, F. Zappa, J. M. Pereira, E. Bessa, M. H. P. Martins, L. F. S. Coelho, W. Wolff, and N. V. de Castro Faria, *Nucl. Instrum. Methods Phys. Res. A* **560**, 219 (2006).
- 2006DIO/PIN H. P. Diogo, S. S. Pinto, and J. J. M. Ramos, *J. Therm Anal. Calorim.* **83**, 361 (2006).
- 2006DIV/CHE S. Divi, R. Chellappa, and D. Chandra, *J. Chem. Thermodyn.* **38**, 1312 (2006).
- 2006DOM/MOR U. Domanska, P. Morawski, and R. Wierzbicki, *Fluid Phase Equilib.* **242**, 154 (2006).
- 2006DRE/SHA V. A. Drebushchak, T. P. Shakhshneider, S. A. Apenina, T. N. Dresushchak, A. S. Medvedeva, L. P. Safronova, and V. V. Boldyrev, *J. Therm Anal. Calorim.* **84**, 643 (2006).
- 2006DRU/DOR A. I. Druzhinina, O. V. Dorofeeva, R. M. Varushchenko, and E. L. Krasnykh, *J. Chem. Thermodyn.* **38**, 10 (2006).
- 2006DRU/KRO A. I. Druzhinina, O. V. Krol, A. A. Efimova, R. M. Varushchenko, and L. L. Gervits, *Russ. J. Phys. Chem.* **80**, 1742 (2006).
- 2006EME/KAB V. N. Emel'yanenko, G. J. Kabo, and S. P. Verevkin, *J.*

- Chem. Eng. Data **51**, 79 (2006).
- 2006FIL/STA E. S. Filatov, P. A. Stabnikov, P. P. Semyannikov, S. V. Trubin, and I. K. Igumenov, *Russ. J. Coord. Chem.* **32**, 126 (2006).
- 2006FIL/SYS E. S. Filatov, S. V. Sysoev, L. N. Zelenina, T. P. Chusova, V. A. Logvinenko, P. P. Semyannikov, and I. K. Igumenov, *J. Therm Anal. Calorim.* **86**, 537 (2006).
- 2006FUJ/MAT H. Fujimori, K. Matsuda, A. Todoroki, T. Asaji, and M. Oguni, *J. Non-Cryst. Solids* **352**, 4790 (2006).
- 2006FUL/RUZ M. Fulem, K. Ruzicka, V. Ruzicka, T. Simecek, E. Huličius, and J. Pangrac, *J. Chem. Thermodyn.* **38**, 312 (2006).
- 2006GEN/AMA A. Genovese, G. Amarasinghe, M. Glewis, D. Mainwaring, and R. A. Shanks, *Thermochim. Acta* **443**, 235 (2006).
- 2006HAF/PAR J. J. H. Haftka, J. R. Parsons, and H. A. J. Govers, *J. Chromatogr. A* **1135**, 91 (2006).
- 2006HUO/ZEN W.-L. Huo, Z.-X. Zeng, and W.-L. Xue, *J. Chem. Eng. Data* **51**, 2110 (2006).
- 2006KHI/BOU K. Khimeche, Y. Boumrah, M. Benziane, and A. Dahmani, *Thermochim. Acta* **444**, 166 (2006).
- 2006KHI/DAH K. Khimeche and A. Dahmani, *J. Therm Anal. Calorim.* **84**, 47 (2006).
- 2006KHI/DAH2 K. Khimeche and A. Dahmani, *J. Chem. Eng. Data* **51**, 382 (2006).
- 2006KRA/VER E. L. Krasnykh, S. P. Verevkin, B. Koutek, and J. Doub-sky, *J. Chem. Thermodyn.* **38**, 717 (2006).
- 2006KUR/PER S. V. Kurkov, G. L. Perlovich, and W. Zielenkiewicz, *J. Therm Anal. Calorim.* **83**, 549 (2006).
- 2006LAH/PAA A. Lahtinen, J. Paasivirta, and V. A. Nikiforov, *Thermochim. Acta* **447**, 5 (2006).
- 2006LI/ROD J. Li, M. Rodrigues, A. Paiva, H. A. Matos, and E. G. de Azevedo, *Fluid Phase Equilib.* **241**, 196 (2006).
- 2006LI/STO H. Li, J. G. Stowell, T. B. Borchardt, and S. R. Bryn, *Cryst. Growth Des.* **11**, 2469 (2006).
- 2006LIS/SEM T. I. Liskovskaya, P. P. Semyannikov, I. A. Baidina, P. A. Stabnikov, N. V. Pervukhina, V. A. Logvinenko, I. K. Igumenov, and I. E. Sokolov, *J. Struct. Chem.* **47**, 726 (2006).
- 2006MAN/CUT A. Mandanici, M. Cutroni, A. Triolo, V. Rodriguez-Mora, and M. A. Ramos, *J. Chem. Phys.* **125**, 054514 (2006).
- 2006MAN/ROH T. Manifar, S. Rohani, M. Jennings, D. Hairsine, and I. Dance, *Cryst. Eng. Comm.* **8**, 59 (2006).
- 2006MAT/MIR M. A. R. Matos, M. S. Miranda, V. M. F. Morais, and J. F. Liebman, *Mol. Phys.* **104**, 1833 (2006).
- 2006MAT/MIR2 M. A. R. Matos, M. S. Miranda, V. M. F. Morais, and J. F. Liebman, *Mol. Phys.* **104**, 2855 (2006).
- 2006MIR/MOR M. S. Miranda, V. M. F. Morais, and M. A. R. Matos, *J. Chem. Thermodyn.* **38**, 559 (2006).
- 2006MON/SAN M. J. S. Monte, L. M. N. B. F. Santos, M. Fulem, J. M. S. Fonseca, and C. A. D. Sousa, *J. Chem. Eng. Data* **51**, 757 (2006).
- 2006MOR/MIR V. M. F. Morais, M. S. Miranda, and M. A. R. Matos, *J. Chem. Thermodyn.* **38**, 450 (2006).
- 2006MOR/MIR2 V. M. F. Morais, M. S. Miranda, M. A. R. Matos, and J. F. Liebman, *Mol. Phys.* **104**, 325 (2006).
- 2006MUL/MOZ P. Mulder, O. Mozenon, S. Lin, C. E. S. Bernardes, M. E. Minas da Piedade, A. F. L. O. Santos, M. A. V. Ribeiro da Silva, G. A. DiLabio, H.-G. Korth, and K. U. Ingold, *J. Phys. Chem. A* **110**, 9949 (2006).
- 2006NAS/NEU K. Nasurzadeh, R. Neueder, and W. Kunz, *J. Chem. Eng. Data* **51**, 7 (2006).
- 2006NIC/KWE G. Nichols, S. Kweskin, M. Frericks, S. Reiter, G. Wang, J. Orf, B. Carvallo, D. Hillesheim, and J. S. Chickos, *J. Chem. Eng. Data* **51**, 475 (2006).
- 2006NIE/GON Q. Nie, J. B. Gong, J. K. Wang, and S. Wang, *Ind. Eng. Chem. Res.* **45**, 432 (2006).
- 2006NOR/RAS F. L. Nordstrom and A. C. Rasmuson, *J. Chem. Eng. Data* **51**, 1775 (2006).
- 2006NOR/RAS2 F. L. Nordstrom and A. C. Rasmuson, *J. Pharm. Sci.* **95**, 748 (2006).
- 2006PAL/ORA M. Palczewska-Tuliuska and P. Oracz, *J. Chem. Eng. Data* **51**, 639 (2006).
- 2006PER/CON S. Perisanu, I. Continanu, M. D. Banciu, H. Zhao, N. Ruth, and J. S. Chickos, *Struct. Chem.* **17**, 639 (2006).
- 2006PIN/DIO S. P. Pinto and H. P. Diogo, *J. Chem. Thermodyn.* **38**, 1515 (2006).
- 2006PRO/RAS V. M. Profrir and A. C. Rasmuson, *Cryst. Growth Des.* **6**, 1143 (2006).
- 2006RIB/AMA M. A. V. Ribeiro da Silva, L. M. P. F. Amaral, and J. R. B. Gomes, *J. Phys. Chem. A* **110**, 9301 (2006).
- 2006RIB/AMA2 M. A. V. Ribeiro da Silva, L. M. P. F. Amaral, A. F. L. O. M. Santos, and J. R. B. Gomes, *J. Chem. Thermodyn.* **38**, 367 (2006).
- 2006RIB/AMA3 M. A. V. Ribeiro da Silva, L. M. P. F. Amaral, A. F. L. O. M. Santos, and J. R. B. Gomes, *J. Chem. Thermodyn.* **38**, 748 (2006).
- 2006RIB/CAB M. A. V. Ribeiro da Silva and J. I. T. A. Cabral, *J. Chem. Eng. Data* **51**, 1556 (2006).
- 2006RIB/CAB2 M. A. V. Ribeiro da Silva and J. I. T. A. Cabral, *J. Chem. Eng. Data* **51**, 767 (2006).
- 2006RIB/CAB3 M. A. V. Ribeiro da Silva and J. I. T. A. Cabral, *J. Chem. Thermodyn.* **38**, 1008 (2006).
- 2006RIB/CAB4 M. A. V. Ribeiro da Silva and J. L. T. A. Cabral, *J. Chem. Thermodyn.* **38**, 1461 (2006).
- 2006RIB/CAB5 M. A. V. Ribeiro da Silva, J. I. T. A. Cabral, P. Gomes, and J. R. B. Gomes, *J. Org. Chem.* **71**, 3677 (2006).
- 2006RIB/CAB6 M. A. V. Ribeiro da Silva, J. I. T. A. Cabral, and J. R. B. Gomes, *J. Chem. Thermodyn.* **38**, 1072 (2006).
- 2006RIB/FER M. A. V. Ribeiro da Silva, A. I. M. C. L. Ferreira, and J. R. B. Gomes, *Bull. Chem. Soc. Jpn.* **79**, 1852 (2006).
- 2006RIB/FER2 M. A. V. Ribeiro da Silva, A. I. M. C. L. Ferreira, and J. R. B. Gomes, *Chem. Phys. Lett.* **422**, 565 (2006).
- 2006RIB/MAT M. A. V. Ribeiro da Silva, M. A. R. Matos, and L. M. P. F. Amaral, *J. Chem. Thermodyn.* **38**, 49 (2006).
- 2006RIB/MON M. A. V. Ribeiro da Silva, M. J. S. Monte, and L. M. N. B. F. Santos, *J. Chem. Thermodyn.* **38**, 778 (2006).
- 2006RIB/SAN M. A. V. Ribeiro da Silva, C. P. F. Santos, M. J. S. Monte, and C. A. D. Sousa, *J. Therm Anal. Calorim.* **83**, 533 (2006).
- 2006RIB/SAN2 M. A. V. Ribeiro da Silva and L. M. N. F. F. Santos, *J. Chem. Thermodyn.* **38**, 817 (2006).
- 2006RIB/SAN3 M. A. V. Ribeiro da Silva, L. M. N. B. F. Santos, and B. Schoder, *J. Chem. Thermodyn.* **38**, 1455 (2006).
- 2006ROD/GIN S. Rodriguez, B. Giner, M. Haro, S. Martin, and H. Artigas, *Phys. Chem. Liq.* **44**, 275 (2006).
- 2006ROU/TEM M. V. Roux, M. Temprado, P. Jimenez, R. Notario, R. Guzman-Mejia, and E. Juaristi, *J. Org. Chem.* **71**, 2581 (2006).
- 2006SAW/MOK T. Sawaya, I. Mokbel, N. Ainous, E. Rauzy, C. Berro, and J. Jose, *J. Chem. Eng. Data* **51**, 854 (2006).
- 2006SCH/SCH A. C. Schmidt, I. Schwarz, and K. Mereiter, *J. Pharm. Sci.* **95**, 1097 (2006).
- 2006SED/MAI Z. Sedlakova, I. Majjevska, K. Rehak, and P. Vrbka, *Collect. Czech. Chem. Commun.* **71**, 1350 (2006).
- 2006SEM/IGU P. P. Semyannikov, I. K. Igumenov, S. V. Trubin, and Z. I. Semenova, *Thermochim. Acta* **451**, 80 (2006).
- 2006SHE/KAB M. P. Shevelyova, G. J. Kabo, A. V. Blokhin, A. G. Kabo, J. A. Jursha, and A. A. Rajko, *J. Chem. Eng. Data* **51**, 40 (2006).
- 2006SUN/LIU X.-H. Sun, Y.-F. Liu, Z.-C. Tan, Y.-Q. Jia, and M.-H. Wang, *Gaodeng Xuexiao Huaxue Xuebao* **27**, 1109 (2006); *Chem. Abstr.* **145**, 426927 (2006).
- 2006TEI/GON A. C. T. Teixeira, A. M. P. S. Goncalves da Silva, and A. C. Fernandes, *Chem. Phys. Lipids* **144**, 160 (2006).
- 2006TEM/ROU M. Temprado, M. V. Roux, P. Jimenez, R. Guzman-Mejia, and E. Juaristi, *Thermochim. Acta* **441**, 20 (2006).
- 2006TEO/BAR M. Teodorescu, A. Barhala, and D. Dragoescu, *J. Chem. Thermodyn.* **38**, 1432 (2006).
- 2006UMN/KWE P. Umnahanant, S. Kweskin, G. Nichols, M. J. Dunn, H. Smart-Ebinne, and J. S. Chickos, *J. Chem. Eng. Data* **51**, 2246 (2006).
- 2006VAN/VAN P. J. van Ekeren, A. C. G. van Genderen, and G. J. K. van den Berg, *Thermochim. Acta* **446**, 33 (2006).
- 2006VAS/VER T. V. Vasiltsova, S. P. Verevkin, E. Bich, A. Heintz, R. Bogel-Lukasik, and U. Domanska, *J. Chem. Eng. Data* **51**,

- 213 (2006).
- 2006VEG/PET D. Vega, A. Petragalli, D. Fernandez, and J. A. Ellena, *J. Pharm. Sci.* **95**, 1075 (2006).
- 2006VEG/POL D. R. Vega, G. Polla, A. Martinez, E. Mendioroz, and M. Reinoso, *Int. J. Pharm.* **328**, 112 (2006).
- 2006VEN/MET L. Ventola, V. Metivaud, L. Bayes, R. Benzges, M. A. Cuevas-Diarte, T. Calvet, and D. Mondieig, *Helv. Chim. Acta* **89**, 2027 (2006).
- 2006VER S. P. Verevkin, *J. Chem. Thermodyn.* **38**, 1111 (2006).
- 2006VER/KOZ S. P. Verevkin, S. A. Kozlova, V. N. Emel'yanenko, E. D. Nikitin, A. P. Popov, and E. L. Krasnykh, *J. Chem. Eng. Data* **51**, 1896 (2006).
- 2006WAN/FAN Z. Wang, W. Fang, R. Lin, Y. Guo, and X. Zhou, *Energy Fuels* **85**, 1794 (2006).
- 2006WAN/TAN M.-H. Wang, Z.-C. Tan, Q. Shi, L.-X. Sun, and T. Zhang, *J. Therm Anal. Calorim.* **84**, 413 (2006).
- 2006WAN/TOZ S. Wang, K. Tozaki, H. Hayashi, H. Inaba, and H. Yamamoto, *Thermochim. Acta* **448**, 73 (2006).
- 2006WAS/HOL C. M. Wassvik, A. G. Holmen, C. A. S. Bergstrom, I. Zamora, and P. Artursson, *Eur. J. Pharm. Sci.* **29**, 294 (2006).
- 2006XU/SUN F. Xu, L. X. Sun, Z. C. Tan, J. G. Liang, and T. Zhang, *J. Therm Anal. Calorim.* **83**, 187 (2006).
- 2006ZAI/PAU D. H. Zaitsau, Y. U. Paulechka, G. J. Kabo, A. N. Kolpikau, V. N. Emel'yanenko, A. Heintz, and S. P. Verevkin, *J. Chem. Eng. Data* **51**, 130 (2006).
- 2006ZAK/LAZ D. V. Zakharychev, S. N. Lazarev, Z. A. Bredikhina, and A. A. Bredikhin, *Russ. Chem. Bull.* **55**, 230 (2006).
- 2006ZEL/CHU L. N. Zelenina, T. P. Chusova, Yu. G. Stenin, and V. V. Bakovets, *Russ. J. Phys. Chem.* **80**, 139 (2006).
- 2006ZEL/CHU2 L. N. Zelenina, T. P. Chusova, Yu. G. Stenin, and G. A. Berezovskii, *Russ. J. Phys. Chem.* **80**, 1911 (2006).
- 2006ZHA/BAI G. I. Zharkova, I. A. Baidina, P. A. Stabnikov, and I. K. Igumenov, *J. Struct. Chem.* **47**, 716 (2006).
- 2007ABE/BOU M. Abes, L. Bouzidi, and S. S. Narine, *Chem. Phys. Lipids* **150**, 89 (2007).
- 2007BAB/SUB P. S. Babu, C. V. S. Subrahmanyam, J. Thimmasetty, R. Manavalan, and K. Valliappan, *Pak. J. Pharm. Sci.* **20**, 311 (2007).
- 2007BAD/BLA E. Badea, I. Blanco, and G. Della Gatta, *J. Chem. Thermodyn.* **39**, 1392 (2007).
- 2007BAS/AND A. Bashkirava, P. C. Andrews, P. C. Junk, E. G. Robertson, L. Spiccia, and N. Vanderhoek, *Chem. Asian J.* **2**, 530 (2007).
- 2007BER/MIN C. E. S. Bernardes, M. E. Minas da Piedade, L. M. P. F. Amaral, A. I. M. C. L. Ferreira, M. A. V. Ribeiro da Silva, H. P. Diogo, and B. J. C. Cabral, *J. Phys. Chem. A* **111**, 1713 (2007).
- 2007BER/WAS C. A. S. Bergstrom, C. A. M. Wassvik, K. Johansson, and I. Hubatsch, *J. Med. Chem.* **50**, 5858 (2007).
- 2007BES/BAI A. A. Bessonov, I. A. Baidina, N. B. Morozova, P. P. Semyannikov, S. V. Trubin, N. V. Gelfond, and I. K. Igumenov, *J. Struct. Chem.* **48**, 282 (2007).
- 2007BES/MOR A. A. Bessonov, N. B. Morozova, N. V. Gelfond, P. P. Semyannikov, S. V. Trubin, Yu. V. Shevtsov, Yu. V. Shubin, and I. K. Igumenov, *Surf. Coat. Technol.* **201**, 9099 (2007).
- 2007BON/CAT M. G. Bonicelli, A. Catalani, G. Mariano, and S. Vecchio, *Thermochim. Acta* **466**, 69 (2007).
- 2007BOU/BEL G. Boukais-Belaribi, B. F. Belaribi, J. Lohmann, and J. Jose, *Fluid Phase Equilib.* **262**, 180 (2007).
- 2007CAI/BET M. R. Caira, G. Bettinetti, M. Sorrenti, and L. Catenacci, *J. Pharm. Sci.* **96**, 996 (2007).
- 2007CAM/MOL A. G. Camacho, J. M. Moll, S. Canzonieri, and M. A. Postigo, *J. Chem. Eng. Data* **52**, 871 (2007).
- 2007CAN/EYL A. I. Canizo, G. N. Eyler, and G. P. Barreto, *Chromatographia* **65**, 21 (2007).
- 2007CAP/LOV C. D. Cappa, E. R. Lovejoy, and A. R. Ravishankara, *J. Phys. Chem. A* **111**, 3099 (2007).
- 2007CHE/HUM L. Chebil, C. Humeau, J. Anthoni, F. Dehez, J.-M. Engasser, and M. Ghoul, *J. Chem. Eng. Data* **52**, 1552 (2007).
- 2007CHE/KIM Y.-H. Cheon and K.-J. Kim, *J. Chem. Eng. Data* **52**, 1390 (2007).
- 2007CHI/JOH R. D. Chirico, R. D. Johnson III, and W. V. Steele, *J. Chem. Thermodyn.* **39**, 698 (2007).
- 2007CHU/ZEL T. P. Chusova, L. N. Zelenina, Yu. G. Stenin, Z. I. Semenova, and V. A. Titov, *Russ. Chem. Bull.* **56**, 1313 (2007).
- 2007DON/LI J.-X. Dong, Q. Li, Z.-C. Tan, Z.-H. Zhang, and Y. Liu, *J. Chem. Thermodyn.* **39**, 108 (2007).
- 2007DRU/EFI A. I. Druzhinina, A. A. Efimova, R. M. Varushchenko, and N. V. Chelovskaya, *Russ. J. Phys. Chem.* **81**, 2090 (2007).
- 2007ECK/HUA B. Eckl, Y.-L. Huang, J. Vrabec, and H. Hasse, **260**, 177 (2007).
- 2007EFI/PAS A. A. Efimova, L. L. Pashchenko, R. M. Varushchenko, E. Krasnyh, and S. V. Levanova, *J. Chem. Thermodyn.* **39**, 142 (2007).
- 2007EME/DAB V. N. Emel'yanenko, A. Dabrowska, S. P. Verevkin, M. O. Hertel, H. Scheuren, and K. Sommer, *J. Chem. Eng. Data* **52**, 468 (2007).
- 2007EME/KOZ V. N. Emel'yanenko, S. A. Kozlova, S. P. Verevkin, and G. N. Roganov, *J. Chem. Thermodyn.* **39**, 10 (2007).
- 2007EME/VER V. N. Emel'yanenko, S. P. Verevkin, O. V. Krol, R. M. Varushchenko, and N. V. Chelovskaya, *J. Chem. Thermodyn.* **39**, 594 (2007).
- 2007ESP/NIC P. Espeau, B. Nicolai, R. Ceolin, M.-A. Perrin, L. Zaska, J. Giovannini, and F. Leveiller, *J. Therm Anal. Calorim.* **90**, 341 (2007).
- 2007FER/BAD D. Ferro, E. Badea, M. Jozwiak, and J. V. Rau, *Thermochim. Acta* **460**, 50 (2007).
- 2007FRE/OLI V. L. S. Freitas, L. I. P. Oliveira, and M. D. M. C. Ribeiro da Silva, *J. Chem. Thermodyn.* **39**, 39 (2007).
- 2007GBA/NEG G. Gbabode, P. Negrier, D. Mondieig, E. M. Calvo, T. Calvet, and M. A. Cuevas-Diarte, *Chem.-Eur. J.* **13**, 3150 (2007).
- 2007GNA/PLA I. I. Gnatyuk, N. V. Platovova, G. A. Puchkovskaya, E. N. Kotelnikova, S. K. Filatov, J. Baran, and M. Drozd, *J. Struct. Chem.* **48**, 654 (2007).
- 2007GOE/MCC A. Goel, L. L. McConnell, and A. Torrents, *J. Environ. Sci. Health, Part B* **42**, 342 (2007).
- 2007GUP/SIN R. K. Gupta, S. K. Singh, and R. A. Singh, *J. Cryst. Growth* **300**, 415 (2007).
- 2007HAF/MAH S. L. Hafsaoui and R. Mahmoud, *J. Therm Anal. Calorim.* **88**, 565 (2007).
- 2007HEI/KAP A. Heintz, S. Kapteina, and S. P. Verevkin, *J. Phys. Chem. A* **111**, 6552 (2007).
- 2007HUL/JOH A. T. Hulme, A. Johnston, A. J. Florence, P. Fernandes, K. Shankland, C. T. Bedford, G. W. A. Welch, G. Sadiq, D. A. Haynes, W. D. S. Motherwell, D. A. Toucher, and S. L. Price, *J. Am. Chem. Soc.* **129**, 3649 (2007).
- 2007IGU/SEM I. K. Igumenov, P. P. Semyannikov, S. V. Trubin, N. B. Morozova, N. V. Gelfond, A. V. Mischenko, and J. A. Norman, *Surf. Coat. Technol.* **201**, 9003 (2007).
- 2007KAN/SOR Y. Kaneko and M. Sorai, *Phase Transitions* **80**, 517 (2007).
- 2007KAR/KAB L. S. Karpushenkava, G. J. Kabo, A. B. Bazyleva, A. V. Blokhin, A. G. Kabo, D. H. Zaitsau, A. A. Pinerzin, and V. S. Sarkisova, *Thermochim. Acta* **459**, 104 (2007).
- 2007KUT/POL A. M. Kut'in, V. S. Polyakov, M. F. Churbanov, and G. E. Snopatin, *Inorg. Mater.* **43**, 1018 (2007).
- 2007LAR/PER F. Lara-Ochoa, G. E. Perez, and F. Miyangos-Santiago, *J. Mol. Struct.* **840**, 97 (2007).
- 2007LEE/LAI M.-J. Lee, C.-H. Lai, T.-B. Wang, and H.-M. Lin, *J. Chem. Eng. Data* **52**, 1291 (2007).
- 2007LIP/KAP D. Lipkind, Y. Kapustin, P. Umnahanant, and J. S. Chickos, *Thermochim. Acta* **456**, 94 (2007).
- 2007LIU/LIU L.-S. Liu, W.-S. Liu, N. Zhou, X.-N. Fan, H. Song, C. Fu, and Y.-Y. Yang, *Gaoxiao Huaxue Gongcheng Xuebao* **21**, 875 (2007); *Chem. Abstr.* **148**, 222587 (2007).
- 2007LU/WAN J. Lu, X.-J. Wang, X. Yang, and C.-B. Ching, *Cryst. Growth Des.* **7**, 1590 (2007).
- 2007MAL S. K. Malanowski, *J. Chem. Eng. Data* **52**, 239 (2007).
- 2007MAL/ALI I. P. Malkerova, A. S. Alikhanyan, S. V. Eliseeva, V. A. Ketsko, and N. P. Kuz'mina, *Russ. J. Inorg. Chem.* **52**,

- 918 (2007).
- 2007MAT/MIR M. A. R. Matos, M. S. Miranda, S. M. M. Pereira, V. M. F. Morais, and J. F. Liebman, *J. Phys. Chem. A* **111**, 7181 (2007).
- 2007MAT/MOR M. A. R. Matos, V. M. F. Morais, C. C. S. Sousa, M. V. Roux, R. Notario, and J. F. Liebman, *Mol. Phys.* **105**, 1789 (2007).
- 2007MAT/SOU M. A. R. Matos, C. C. S. Sousa, and V. M. F. Morais, *J. Chem. Eng. Data* **52**, 1089 (2007).
- 2007MAT/MIR2 M. A. R. Matos, M. S. Miranda, M. J. S. Monte, L. M. N. B. F. Santos, V. M. F. Morais, J. S. Chickos, P. Umnahant, and J. F. Liebman, *J. Phys. Chem. A* **111**, 11153 (2007).
- 2007MIS/MIS A. K. Misra, M. Misra, G. M. Panpalia, and A. K. Dorle, *J. Macromol. Sci.* **44A**, 685 (2007).
- 2007MOK/RAZ M. Mokbel, A. Razzouk, A. Hajjaji, N. Misakni, and J. Jose, *J. Chem. Eng. Data* **52**, 1720 (2007).
- 2007MON/PER M. Moneghini, B. Perissutti, F. Vecchione, I. Kikic, P. Alessi, and A. Cortesi, *Curr. Drug Del.* **4**, 241 (2007).
- 2007MOO/KOE D. J. Moore, D. Koelmel, D. Laura, and E. Bedford, *Chem. Phys. Lipids* **150**, 109 (2007).
- 2007MOR/COR E. Moreno, R. Cordobilla, T. Calvet, M. A. Cuevas-Diarte, G. Gbabode, P. Negrier, D. Mondieig, and H. A. J. Oonk, *New J. Chem.* **31**, 947 (2007).
- 2007MOR/MIR V. M. F. Morais, M. S. Miranda, and M. A. R. Matos, *J. Chem. Eng. Data* **52**, 627 (2007).
- 2007PAL/ORA M. Palczewska-Tulinska and P. Oracz, *J. Chem. Eng. Data* **52**, 645 (2007).
- 2007PAL/ORA2 M. Palczewska-Tuliuska and P. Oracz, *J. Chem. Eng. Data* **52**, 2468 (2007).
- 2007PAN/ANT R. Pankajavalli, S. Anthonysamy, K. Ananthasivan, and P. P. V. Rao, *J. Nucl. Mater.* **362**, 128 (2007).
- 2007PAN/ANT2 K. Panneerselvam, M. P. Antony, T. G. Srinivasan, and P. R. Vasudeva Rao, *Thermochim. Acta* **466**, 49 (2007).
- 2007PAS/BET I. Pasquali, R. Bettini, and F. Giordano, *J. Therm. Anal. Calorim.* **90**, 903 (2007).
- 2007PAS/KUZ L. L. Pashchenko and T. S. Kuznetsova, *Russ. J. Phys. Chem.* **81**, 1238 (2007).
- 2007PER/HAN G. L. Perlovich, L. Kr. Hansen, T. V. Volkova, S. Mirza, A. N. Manin, and A. Bauer-Brandl, *Cryst. Growth Des.* **7**, 2643 (2007).
- 2007PER/STR G. L. Perlovich, N. N. Strakhova, V. P. Kazachenko, T. V. Volkova, V. V. Tkachev, K.-J. Schaper, and O. A. Raevsky, *Int. J. Pharm.* **334**, 115 (2007).
- 2007PER/SUR G. L. Perlovich, A. O. Surov, L. Kr. Hansen, and A. Bauer-Brandl, *J. Pharm. Sci.* **96**, 1031 (2007).
- 2007PER/SUR2 G. L. Perlovich, A. O. Surov, and A. Bauer-Brandl, *J. Pharm. Biomed. Anal.* **45**, 679 (2007).
- 2007PER/VOL G. L. Perlovich, T. V. Volkova, and A. Bauer-Brandl, *Mol. Pharmacol.* **4**, 929 (2007).
- 2007PIN/BER S. S. Pinto, C. E. S. Bernardes, H. P. Diogo, and M. E. Minas da Piedade, *J. Chem. Thermodyn.* **39**, 1384 (2007).
- 2007RAM/ROJ L. G. Ramirez-Verduzco, A. Rojas-Aguilar, J. A. de los Reyes, J. A. Munoz-Arroyo, and F. Murrieta-Guevara, *J. Chem. Eng. Data* **52**, 2212 (2007).
- 2007RAU/BEH R. B. Rauch and R. Behrens, *Propellants, Explos., Pyrotech.* **32**, 97 (2007).
- 2007RAZ/MOK A. Razzouk, I. Mokbel, J. Garcia, J. Fernandez, N. Msakni, and J. Jose, *Fluid Phase Equilib.* **260**, 248 (2007).
- 2007RIB/AMA M. A. V. Ribeiro da Silva, L. M. P. F. Amaral, and J. R. B. Gomes, *J. Phys. Chem. B* **111**, 792 (2007).
- 2007RIB/AMA2 M. A. V. Ribeiro da Silva, L. M. P. F. Amaral, and J. R. B. Gomes, *J. Phys. Chem. B* **111**, 13033 (2007).
- 2007RIB/AMA3 M. A. V. Ribeiro da Silva, L. M. P. F. Amaral, and J. R. B. Gomes, *Pol. J. Chem.* **81**, 865 (2007).
- 2007RIB/ARA M. D. M. C. Ribeiro da Silva and N. R. M. Araujo, *J. Chem. Thermodyn.* **39**, 1372 (2007).
- 2007RIB/CAB M. A. V. Ribeiro da Silva and J. I. T. A. Cabral, *J. Therm. Anal. Calorim.* **90**, 865 (2007).
- 2007RIB/CAB2 M. A. V. Ribeiro da Silva and J. I. T. A. Cabral, *Thermochim. Acta* **453**, 147 (2007).
- 2007RIB/FER M. A. V. Ribeiro da Silva, A. I. M. C. L. Ferreira, and J. R. B. Gomes, *J. Phys. Chem. B* **111**, 2052 (2007).
- 2007RIB/FER2 M. A. V. Ribeiro da Silva, A. I. M. C. L. Ferreira, and J. R. B. Gomes, *J. Phys. Chem. B* **111**, 6444 (2007).
- 2007RIB/FRE M. D. M. C. Ribeiro da Silva, V. L. S. Freitas, L. M. N. B. F. Santos, M. Fulem, M. J. Sottomayor, M. J. S. Monte, and W. E. Acree, Jr., *J. Chem. Eng. Data* **52**, 580 (2007).
- 2007RIB/MON M. A. V. Ribeiro da Silva, I. M. M. Monteiro, L. M. N. B. F. Santos, and B. Schroder, *J. Chem. Thermodyn.* **39**, 767 (2007).
- 2007RIB/SAN M. A. V. Ribeiro da Silva and A. F. L. O. M. Santos, *J. Therm. Anal. Calorim.* **88**, 7 (2007).
- 2007RIB/SAN2 M. A. V. Ribeiro da Silva, L. M. N. B. F. Santos, and E. Giera, *J. Chem. Thermodyn.* **39**, 361 (2007).
- 2007RIB/SAN3 M. A. V. Ribeiro da Silva, C. P. F. Santos, M. J. S. Monte, and C. A. D. Sousa, *J. Chem. Thermodyn.* **39**, 1363 (2007).
- 2007RIC/BER L. S. Richard, C. E. S. Bernardes, H. P. Diogo, J. P. Leal, and M. E. Minas da Piedade, *J. Phys. Chem. A* **111**, 8741 (2007).
- 2007ROU/NOT M. V. Roux, R. Notario, E. Velez, M. Temprado, A. Guerrero, S. P. Verevkin, J. Quijano, and J. Gaviria, *J. Chem. Thermodyn.* **39**, 1377 (2007).
- 2007ROU/TEM M. V. Roux, M. Temprado, P. Jimenez, R. Notario, J. S. Chickos, A. F. L. O. M. Santos, and M. A. V. Ribeiro da Silva, *J. Phys. Chem. A* **111**, 11084 (2007).
- 2007ROU/TEM2 M. V. Roux, M. Temprado, P. Jimenez, R. Notario, R. Guzman-Mejia, and E. Juaristi, *J. Org. Chem.* **72**, 1143 (2007).
- 2007ROU/TEM3 M. V. Roux, M. Temprado, R. Notario, J. S. Chickos, A. F. L. O. M. Santos, and M. A. V. Ribeiro da Silva, *J. Phys. Chem. A* **111**, 5280 (2007).
- 2007SAP/ZAY E. Sapei, A. Zaytseva, P. Uusi-Kyyny, and K. I. Keshinen, *Fluid Phase Equilib.* **252**, 130 (2007).
- 2007SCH/DOE B. Schmid, M. Doeker, and J. Gmehling, *Fluid Phase Equilib.* **258**, 115 (2007).
- 2007SHE/ZAI M. P. Shevelyova, D. H. Zaitsau, Y. U. Paulechka, A. V. Blokhin, G. J. Kabo, S. P. Verevkin, and A. Heintz, *J. Chem. Eng. Data* **52**, 1360 (2007).
- 2007SHI/TAN Q. Shi, Z.-C. Tan, Y.-Y. Di, B. Tong, Y.-S. Li, and S.-X. Wang, *J. Chem. Eng. Data* **52**, 941 (2007).
- 2007SHI/TAN2 Q. Shi, Z.-C. Tan, Y.-Y. Di, B. Tong, S.-X. Wang, and Y.-Z. Li, *Thermochim. Acta* **463**, 6 (2007).
- 2007SHI/TAN3 Q. Shi, Z.-C. Tan, B. Tong, Y.-Y. Di, Z.-H. Zhang, J.-L. Zeng, L.-X. Sun, and Y.-S. Li, *J. Chem. Thermodyn.* **39**, 817 (2007).
- 2007SID/SID M. A. Siddiqui, R. A. Siddiqui, and B. Atakan, *Surf. Coat. Technol.* **201**, 9055 (2007).
- 2007SIF/AIT H. Sifaoui, A. Ait-Kaci, A. Mondarressi, and M. Rogalski, *Thermochim. Acta* **456**, 114 (2007).
- 2007SMI/TSV N. N. Smirnova, L. Ya. Tsvtkova, T. A. Bykova, and Y. Marcus, *J. Chem. Thermodyn.* **39**, 1508 (2007).
- 2007STR/RUZ M. Straka, K. Ruzicka, and V. Ruzicka, *J. Chem. Eng. Data* **52**, 1375 (2007).
- 2007STR/RUZ2 M. Straka, A. van Genderen, K. Ruzicka, and V. Ruzicka, *J. Chem. Eng. Data* **52**, 794 (2007).
- 2007TOM/ROS L. I. N. Tome, M. T. S. Rosado, S. C. C. Nunes, T. M. R. Maria, J. Canotilho, and M. E. S. Eusebio, *J. Chem. Thermodyn.* **39**, 1354 (2007).
- 2007TON/TAN B. Tong, Z.-C. Tan, X. C. Lv, L. X. Sun, F. Xu, Q. Shi, and Y. S. Li, *J. Therm. Anal. Calorim.* **90**, 217 (2007).
- 2007TON/TAN2 B. Tong, Z.-C. Tan, Q. Shi, Y.-S. Li, D.-T. Yue, and S.-X. Wang, *Thermochim. Acta* **457**, 20 (2007).
- 2007VAR/DRU R. M. Varushchenko, A. I. Druzhinina, G. M. Kuramshina, and O. V. Dorofeeva, *Fluid Phase Equilib.* **256**, 112 (2007).
- 2007VEC S. Vecchio, *J. Therm. Anal. Calorim.* **87**, 79 (2007).
- 2007VEC/BRU S. Vecchio and B. Brunetti, *J. Chem. Eng. Data* **52**, 1585 (2007).
- 2007VER S. P. Verevkin, *J. Chem. Eng. Data* **52**, 301 (2007).
- 2007VER/EME S. P. Verevkin, V. N. Emel'yanenko, and A. Klamt, *J. Chem. Eng. Data* **52**, 499 (2007).
- 2007VER/GEO S. P. Verevkin, M. Georgieva, and S. V. Melkhanova, *J.*

- Chem. Eng. Data **52**, 286 (2007).
- 2007VER/SCH S. P. Verevkin and C. Schick, *J. Chem. Thermodyn.* **39**, 758 (2007).
- 2007WAN/TAN S.-X. Wang, Z.-C. Tan, Y.-S. Li, Y. Li, Q. Shi, and B. Tong, *Thermochim. Acta* **463**, 21 (2007).
- 2007XU/ZEN X.-G. Xu, Z.-X. Zeng, W.-I. Xue, and H. Y. Zhang, *J. Chem. Eng. Data* **52**, 1189 (2007).
- 2007XUE/WAN B. Xue, J. Y. Wang, Z.-C. Tan, and T.-H. Wu, *Chem. Res. Chin. Univ.* **23**, 460 (2007).
- 2007ZHA/TAN X.-S. Zhao, Z.-C. Tan, Y.-S. Lu, B. Tong, Q. Shi, and Y. Li, *Wuli Huaxue Xuebao* **23**, 1459 (2007).
- 2007ZIE/SZT W. Zielenkiewicz and P. Szterner, *J. Chem. Eng. Data* **52**, 624 (2007).
- 2008ABA/BAD L. Abate, E. Badea, I. Blando, and G. Della Gatta, *J. Chem. Eng. Data* **53**, 959 (2008).
- 2008ABB/KAL D. Abbas, J. Kaloustian, C. Orneto, P. Piccerelle, H. Portugal, and A. Nicolay, *J. Therm. Anal. Calorim.* **93**, 353 (2008).
- 2008ARU/MAT T. S. Arul Jeevan, T. Mathews, V. S. Raghunathan, and K. S. Nagaraja, *Thermochim. Acta* **478**, 41 (2008).
- 2008BAE A. K. Baev, *Russ. J. Phys. Chem.* **82**, 1266 (2008).
- 2008BAR/BER I. Barsky and J. Bernstein, *Cryst. Eng. Comm.* **10**, 669 (2008).
- 2008BAR/BOT C. A. Barton, M. A. Botelho, and M. A. Kaiser, *J. Chem. Eng. Data* **53**, 939 (2008).
- 2008BAS/BOS S. Basavoju, D. Bostrom, and S. Velaga, *Pharm. Res.* **25**, 530 (2008).
- 2008BAZ/BLO A. B. Bazleva, A. V. Blokhin, A. G. Kabo, G. J. Kabo, V. N. Emel'yanenko, and S. P. Verevkin, *J. Chem. Thermodyn.* **40**, 509 (2008).
- 2008BER/MIN C. E. S. Bernardes and M. E. Minas da Piedade, *J. Phys. Chem. A* **112**, 10029 (2008).
- 2008BER/PIE C. E. S. Bernardes, M. F. M. Piedade, and M. E. Minas da Piedade, *Cryst. Growth Des.* **8**, 2419 (2008).
- 2008BES/MOR A. A. Bessonov, N. B. Morozova, P. P. Semyannikov, S. V. Trubin, N. V. Gel'fond, and I. K. Igumenov, *J. Therm. Anal. Calorim.* **92**, 751 (2008).
- 2008BES/MOR2 A. A. Bessonov, N. B. Morozova, P. P. Semyannikov, S. V. Trubin, N. V. Gel'fond, and I. K. Igumenov, *Russ. J. Coord. Chem.* **34**, 186 (2008).
- 2008BIA/CEZ T. F. Biazus, A. M. Cezaro, G. R. Borges, J. P. Bender, E. Franceschi, M. L. Corazza, and V. Oliveira, *J. Chem. Thermodyn.* **40**, 437 (2008).
- 2008BRA/GEL D. E. Braun, T. Gelbrich, V. Kahlenberg, G. Laus, J. Wieser, and U. J. Griesser, *New J. Chem.* **32**, 1677 (2008).
- 2008BRE/BRE A. A. Bredikhin, Z. A. Bredikhina, V. G. Novikova, A. V. Pashagin, D. V. Zakharychev, and A. T. Gubaidullin, *Chirality* **20**, 1092 (2008).
- 2008BRU/PIA B. Brunetti, V. Piacente, A. Latini, and P. Scardala, *J. Chem. Eng. Data* **53**, 2943 (2008).
- 2008CAP/LOV C. D. Cappa, E. R. Lovejoy, and A. R. Ravishankara, *J. Phys. Chem. A* **112**, 3959 (2008).
- 2008CHI/LIP J. S. Chickos and D. Lipkind, *J. Chem. Eng. Data* **53**, 2432 (2008).
- 2008CHI/STE R. D. Chirico and W. V. Steele, *J. Chem. Thermodyn.* **40**, 806 (2008).
- 2008CHI/WAN J. S. Chickos, T. Wang, and E. Sharma, *J. Chem. Eng. Data* **53**, 481 (2008).
- 2008COG/HIL J. M. Cogen and A. Hilmer, *Polym. Degrad. Stab.* **93**, 2193 (2008).
- 2008DI/WAN Y.-Y. Di, D.-Q. Wang, S. Quan, and Z.-C. Tan, *Chin. Phys. B* **17**, 2859 (2008).
- 2008DIO/PIN H. P. Diogo, S. S. Pinto, and J. J. M. Ramos, *Int. J. Pharm.* **358**, 192 (2008).
- 2008DOM/MOR U. Domanska, P. Morawski, and M. Piekarska, *J. Chem. Thermodyn.* **40**, 710 (2008).
- 2008EME/KOZ V. N. Emel'yanenko, S. A. Kozlova, S. P. Verevkin, and G. N. Roganov, *J. Chem. Thermodyn.* **40**, 911 (2008).
- 2008EME/TOK V. N. Emel'yanenko, A. V. Toktonov, S. A. Kozlova, S. P. Verevkin, V. Andrushko, N. Andrushko, and A. Borner, *J. Phys. Chem. A* **112**, 4036 (2008).
- 2008EME/VER V. N. Emel'yanenko, S. P. Verevkin, E. N. Burakova, G. N. Roganov, and M. K. Georgieva, *Russ. J. Phys. Chem.* **82**, 1521 (2008).
- 2008FAN/WAN R. L. Fan, L.-S. Wang, and M.-Y. Li, *J. Chem. Eng. Data* **53**, 228 (2008).
- 2008FAV/FRE R. Favareto, P. H. Fregadolli, V. F. Cabral, O. A. C. Antunes, and L. Cardozo-Filho, *J. Chem. Eng. Data* **53**, 1080 (2008).
- 2008FRE/KEB A. Freedman, P. L. Kebarian, Z. Li, W. A. Robinson, and J. C. Wormhoudt, *Meas. Sci. Technol.* **19**, 125102/1 (2008).
- 2008FRI/ACR V. L. S. Frietas, W. E. Acree, Jr., and M. D. M. C. Ribeiro da Silva, *J. Chem. Eng. Data* **53**, 1820 (2008).
- 2008GBA/NEG G. Gbabode, P. Negrier, D. Mondieig, E. Moreno, T. Calvet, and M. A. Cuevas-Diarte, *Chem. Phys. Lipids* **154**, 68 (2008).
- 2008GOL/SUU J. L. Goldfarb and E. M. Suuberg, *Envir. Toxicol. Chem.* **27**, 1244 (2008).
- 2008GOL/SUU2 J. L. Goldfarb and E. M. Suuberg, *J. Chem. Thermodyn.* **40**, 460 (2008).
- 2008GOM/AMAJ R. B. Gomes, L. M. P. F. Amaral, and M. A. V. Ribeiro da Silva, *J. Phys. Org. Chem.* **21**, 365 (2008).
- 2008GUA/YAN Y. Gua, F. Yang, Y. Xing, D. Li, W. Fang, and R. Lin, *Energy Fuels* **22**, 510 (2008).
- 2008GUP/GAN P. K. Gupta, K. Ganesan, P. K. Gutch, L. Manral, and D. K. Dubey, *J. Chem. Eng. Data* **53**, 841 (2008).
- 2008HAN/NUT W. Hanshaw, M. Nutt, and J. S. Chickos, *J. Chem. Eng. Data* **53**, 1903 (2008); **53**, 2721(E) (2008).
- 2008JEE/ARO T. S. A. Jeevan, S. Arockiasamy, T. Mathews, V. S. Raghunathan, and K. S. Nagaraja, *Mater. Lett.* **62**, 4170 (2008).
- 2008KIM/SVE I. Kim, H. F. Svendsen, and E. Borresen, *J. Chem. Eng. Data* **53**, 2521 (2008).
- 2008KOL/KUK Z. Kolsa, J. Kukal, M. Zabransky, and V. Ruzicka, *Ind. Eng. Chem. Res.* **47**, 2075 (2008).
- 2008KOZ/EME S. A. Kozlova, V. N. Emel'yanenko, M. Georgieva, S. P. Verevkin, Y. Chernyak, B. Schaffner, and A. Borner, *J. Chem. Thermodyn.* **40**, 1136 (2008).
- 2008KOZ/MAR M. S. Kozlova, A. V. Markin, V. N. Larina, and N. V. Karyzkin, *Russ. J. Phys. Chem.* **82**, 1993 (2008).
- 2008KRO/DRU O. V. Krol, A. I. Druzhinina, R. M. Varushchenko, O. V. Dorofeeva, M. D. Reshetova, and N. E. Borisova, *J. Chem. Thermodyn.* **40**, 549 (2008).
- 2008KUN/SHI G. V. Kunte, S. A. Shivashankar, and A. M. Umarji, *Meas. Sci. Technol.* **19**, 025704/1 (2008).
- 2008KUR/KAW H. Kuramochi, K. Kawamoto, K. Miyazaki, K. Nagahama, K. Maeda, X.-W. Li, E. Shibata, T. Nakamura, and S.-I. Sakai, *Envir. Toxicol. Chem.* **27**, 2413 (2008).
- 2008KUR/PER S. V. Kurkov and G. L. Perlovich, *Int. J. Pharm.* **357**, 100 (2008).
- 2008LAG/JIM A. F. Lago, P. Jimenez, R. Herrero, J. Z. Davalos, and J.-L. M. Abboud, *J. Phys. Chem. A* **112**, 3201 (2008).
- 2008LIP/KRA S. V. Lipp, E. L. Krasnykh, and S. V. Levanova, *Russ. J. Phys. Chem.* **82**, 2025 (2008).
- 2008LLA/MON E. Lladosa, J. B. Monton, and M. C. Burguet, *J. Chem. Eng. Data* **53**, 1897 (2008).
- 2008LOU/PIN C. M. Lousada, S. S. Pinto, J. N. Canongia Lopes, M. F. Minas da Piedade, H. P. Diogo, and M. E. Minas da Piedade, *J. Phys. Chem. A* **112**, 2977 (2008).
- 2008LU/GAO X.-C. Lu, X.-H. Gao, Z.-C. Tan, Y.-S. Li, and L.-X. Sun, *J. Therm. Anal. Calorim.* **92**, 523 (2008).
- 2008MAR/EUS T. M. R. Maria and M. E. S. Eusebio, *J. Chem. Eng. Data* **53**, 1316 (2008).
- 2008MAR/RAM F. J. Martinez Casado, M. Ramos Riesco, A. Sanchez Arenas, M. V. Garci Perez, M. I. Redondo, S. Lopez-Andras, L. Garrido, and J. A. R. Cheda, *J. Phys. Chem. B* **112**, 16601 (2008).
- 2008MAR/SEL M. Maria, J. Selvakumar, V. S. Raghunathan, T. Mathews, and K. S. Nagaraja, *Thermochim. Acta* **474**, 87 (2008).
- 2008MAT/IVA V. E. Matulis, O. A. Ivashkevich, P. N. Gaponik, P. D. Elkind, G. T. Sukhanov, A. B. Bazyleva, and D. H. Zait-sau, *THEOCHEM* **854**, 18 (2008).
- 2008MAT/MIR M. A. R. Matos, M. S. Miranda, D. A. P. Fonseca, V. M. F. Morais, and J. F. Liebman, *J. Phys. Chem. A* **112**, 10053

- (2008).
- 2008MAT/MIR2 M. A. R. Matos, M. S. Miranda, and V. M. F. Morais, *J. Chem. Thermodyn.* **40**, 625 (2008).
- 2008MAT/SOU M. A. R. Matos, C. C. S. Sousa, and V. M. F. Morais, *J. Chem. Thermodyn.* **40**, 1485 (2008).
- 2008MAT/SOU2 M. A. R. Matos, C. C. S. Sousa, and V. M. F. Morais, *J. Phys. Chem. A* **112**, 7961 (2008).
- 2008MEN/FLO J. Mentado, H. Flores, and P. Amador, *J. Chem. Thermodyn.* **40**, 1106 (2008).
- 2008MON/SAN M. J. S. Monte, L. M. N. B. F. Santos, C. A. D. Sousa, and M. Fulem, *J. Chem. Eng. Data* **53**, 1922 (2008).
- 2008MOR/ZHE N. B. Morozova, K. V. Zherikova, I. A. Baidina, S. V. Sysoev, P. P. Semyannikov, L. V. Yakovkina, T. P. Smirnova, N. V. Gelfond, I. K. Igumenov, G. Carta, and G. Rossetto, *J. Phys. Chem. Solids* **69**, 673 (2008).
- 2008MOT/QUE F. L. Mota, A. J. Queimada, S. P. Pinho, and E. A. Macedo, *Ind. Eng. Chem. Res.* **47**, 5182 (2008).
- 2008NIC/BEL S. Nicoli, S. Belzi, P. Santi, M. R. Caira, J. Li, and R. Bettini, *J. Pharm. Sci.* **97**, 4830 (2008).
- 2008NTI/CHM J. Nti-Gyabaah, R. Chmielowski, V. Chan, and Y. C. Chiew, *Int. J. Pharm.* **359**, 111 (2008).
- 2008NUN/CLA E. Nunez, C. G. Clark, Jr., W. Cheng, A. Best, G. Floudas, A. N. Semenov, G. Fytas, and K. Mullen, *J. Phys. Chem. B* **112**, 6542 (2008).
- 2008OSM/CAT A. Osmont, L. Catoire, and I. Gokalp, *Energy Fuels* **22**, 2241 (2008).
- 2008PAN/FUL J. Pangrac, M. Fulem, E. Hulcius, K. Melichar, T. Simceek, K. Ruzicka, P. Moravek, V. Ruzicka, and S. A. Rushworth, *J. Cryst. Growth* **310**, 4720 (2008).
- 2008PEN/JIA L. Peng, X. Jianjun, M. Fangquan, L. Xi, and Z. Chaocan, *J. Therm Anal. Calorim.* **93**, 485 (2008).
- 2008PER/STR G. L. Perlovich, N. N. Strakhova, V. P. Kazachenko, T. V. Volkova, V. V. Tkachev, K.-J. Shaper, and O. A. Raevsky, *Int. J. Pharm.* **349**, 300 (2008).
- 2008PER/VOL G. L. Perlovich, T. V. Volkova, A. N. Manin, and A. N. Bauer-Brandl, *J. Pharm. Sci.* **97**, 3883 (2008).
- 2008PIN/DIO S. S. Pinto and H. P. Diogo, *J. Pharm. Sci.* **97**, 5354 (2008).
- 2008PIO/CAN R. Piciocchi, J. N. Cananaja Lopes, H. P. Diogo, and M. E. Minas da Piedade, *J. Phys. Chem. A* **112**, 10429 (2008).
- 2008RAI/BHA N. Rai, D. Bhatt, J. I. Siepmann, and L. E. Fried, *J. Chem. Phys.* **129**, 194510 (2008).
- 2008RIB/AMA M. A. V. Ribeiro da Silva and L. M. P. F. Amaral, *J. Therm Anal. Calorim.* **92**, 53 (2008).
- 2008RIB/AMA2 M. A. V. Ribeiro da Silva, L. M. P. F. Amaral, C. R. P. Boaventura, and J. R. B. Gomes, *J. Chem. Thermodyn.* **40**, 1226 (2008).
- 2008RIB/AMA3 M. A. V. Ribeiro da Silva, L. M. P. F. Amaral, and A. F. L. O. M. Santos, *J. Chem. Thermodyn.* **40**, 1588 (2008).
- 2008RIB/CAB M. A. V. Ribeiro da Silva and J. I. T. A. Cabral, *J. Therm Anal. Calorim.* **92**, 59 (2008).
- 2008RIB/CAB2 M. A. V. Ribeiro da Silva and J. I. T. A. Cabral, *J. Chem. Thermodyn.* **40**, 829 (2008).
- 2008RIB/CAB3 M. A. V. Ribeiro da Silva, J. I. T. A. Cabral, and J. R. B. Gomes, *J. Phys. Chem. A* **112**, 12263 (2008).
- 2008RIB/FER M. A. V. Ribeiro da Silva and A. I. M. C. L. Ferreira, *J. Chem. Thermodyn.* **40**, 924 (2008).
- 2008RIB/FER2 M. A. V. Ribeiro da Silva and A. I. M. C. L. Ferreira, *J. Chem. Thermodyn.* **40**, 362 (2008).
- 2008RIB/FER3 M. A. V. Ribeiro da Silva, A. I. M. C. L. Ferreira, L. M. S. S. Lima, and S. M. M. Sousa, *J. Chem. Thermodyn.* **40**, 137 (2008).
- 2008RIB/FIG M. A. V. Ribeiro da Silva, D. F. Figueriedo, and J. I. T. A. Cabral, *J. Chem. Thermodyn.* **40**, 369 (2008).
- 2008RIB/RIB M. D. M. C. Ribeiro da Silva, M. A. V. Ribeiro da Silva, V. L. S. Freitas, M. V. Roux, P. Jimenez, M. Temprado, J. Z. Davalos, P. Cabildo, R. M. Claramunt, and J. Elguero, *J. Chem. Thermodyn.* **40**, 386 (2008).
- 2008RIB/RIB2 M. D. M. C. Ribeiro da Silva, M. A. V. Ribeiro da Silva, V. L. S. Freitas, M. V. Roux, P. Jimenez, J. Z. Davalos, P. Cabildo, R. M. Claramunt, and J. Elguero, *J. Chem. Thermodyn.* **40**, 1378 (2008).
- 2008RIB/SAN M. A. V. Ribeiro da Silva and A. F. L. O. M. Santos, *J. Chem. Thermodyn.* **40**, 225 (2008).
- 2008RIB/SAN2 M. A. V. Ribeiro da Silva and A. F. L. O. M. Santos, *J. Chem. Thermodyn.* **40**, 917 (2008).
- 2008RIB/SAN3 M. A. V. Ribeiro da Silva and A. F. L. O. M. Santos, *J. Chem. Thermodyn.* **40**, 1217 (2008).
- 2008RIB/SAN4 M. A. V. Ribeiro da Silva and A. F. L. O. M. Santos, *J. Chem. Thermodyn.* **40**, 1309 (2008).
- 2008RIB/SAN5 M. A. V. Ribeiro da Silva and A. F. L. O. M. Santos, *J. Chem. Thermodyn.* **40**, 1451 (2008).
- 2008RIB/SAN6 M. A. V. Ribeiro da Silva, L. M. N. B. F. Santos, and L. M. S. S. Lima, *J. Chem. Thermodyn.* **40**, 375 (2008).
- 2008RIB/SCH M. A. V. Ribeiro da Silva, B. Schroder, V. B. M. Castro, and L. N. B. F. Santos, *J. Chem. Thermodyn.* **40**, 599 (2008).
- 2008ROU/LIM A. A. Roucha, C. F. R. A. C. Lima, and L. M. N. B. F. Santos, *J. Chem. Thermodyn.* **40**, 1458 (2008).
- 2008ROU/TEM M. V. Roux, M. Temprado, J. S. Chickos, and Y. Nagano, *J. Phys. Chem. Ref. Data* **37**, 1855 (2008).
- 2008ROU/TEM2 M. V. Roux, M. Temprado, R. Notario, C. Foces-Foces, V. N. Emel'yanenko, and S. P. Verevkin, *J. Phys. Chem. A* **112**, 7455 (2008).
- 2008SAG/SAF E. V. Sagadeev and Yu. G. Safina, *Russ. J. Gen. Chem.* **78**, 65 (2008).
- 2008SEM/BAS P. P. Semyannikov, T. V. Basova, S. V. Trubin, E. K. Kol'tsov, V. A. Plyashkevich, and I. K. Igumenov, *Russ. J. Phys. Chem.* **82**, 159 (2008).
- 2008SHA/GUP B. L. Sharma, S. Gupta, S. Tandon, and R. Kant, *Mater. Chem. Phys.* **111**, 423 (2008).
- 2008SHA/WU X.-Z. Shao, J.-S. Wu, G.-Q. Zhang, and L.-S. Wang, *J. Chem. Eng. Data* **53**, 1012 (2008).
- 2008SIN/DAS N. B. Singh, S. S. Das, P. Gupta, and M. K. Dwivedi, *J. Cryst. Growth* **311**, 118 (2008).
- 2008SIN/DAS2 N. B. Singh, S. S. Das, N. P. Singh, and T. J. Agrawal, *J. Cryst. Growth* **310**, 2878 (2008).
- 2008SIN/MUR L. P. Singh and S. S. N. Murthy, *J. Phys. Chem. B* **112**, 2606 (2008).
- 2008SIN/MUR2 L. P. Singh and S. S. N. Murthy, *J. Chem. Phys.* **129**, 094501 (2008).
- 2008SLY/KON I. V. Slyusarev, Yu. V. Kondrat'ev, A. O. Kozin, and L. P. Belorukova, *Vestn. St. Petersburg. Universit. Ser. 4 Fiz. Khim.*, 64 (2008).
- 2008SON/RAM M. Soni, D. Ramjugernath, and J. D. Raal, *J. Chem. Eng. Data* **53**, 745 (2008).
- 2008STI/CIN V. Stilinovic, D. Cincic, and B. Kaitner, **55**, 874 (2008).
- 2008STU/ROR H. K. Stulzer, P. O. Roriguez, T. M. Cardoso, J. S. R. Matos, and M. A. S. Silva, *J. Therm Anal. Calorim.* **91**, 323 (2008).
- 2008SUI/JES V. Suitchmezian, I. Jess, and C. Naether, *J. Pharm. Sci.* **97**, 4516 (2008).
- 2008SUR O. V. Surov, *Russ. J. Gen. Chem.* **78**, 602 (2008).
- 2008TEM/ROU M. Temprado, M. V. Roux, and J. S. Chickos, *J. Therm Anal. Calorim.* **94**, 257 (2008).
- 2008TEM/ROU2 M. Temprado, M. V. Roux, P. Jimenez, C. Foces-Foces, and R. Notario, *J. Phys. Chem. A* **112**, 10378 (2008).
- 2008TEM/ROU3 M. Temprado, M. V. Roux, A. R. Parameswar, A. V. Demchenko, J. S. Chickos, and J. F. Liebman, *J. Therm Anal. Calorim.* **91**, 471 (2008).
- 2008THI/SUB J. Thimmasetty, C. V. S. Subrahmanyam, P. R. S. Babu, M. A. Maulik, and B. A. Viswanath, *J. Solution Chem.* **37**, 1365 (2008).
- 2008TON/TAN B. Tong, Z.-C. Tan, and S.-X. Wang, *Chin. J. Chem.* **26**, 1561 (2008).
- 2008TON/TAN2 B. Tong, Z. C. Tan, Q. Shi, Y. S. Li, and S. X. Wang, *J. Therm Anal. Calorim.* **91**, 463 (2008).
- 2008TON/TAN3 B. Tong, Z.-C. Tan, and S.-X. Wang, *J. Environ. Eng.* **24**, 1699 (2008); *Chem. Abstr.* **150**, 11821 (2008).
- 2008TUN/TAB H.-H. Tung, J. Tabora, N. Variankaval, D. Bakken, and C.-C. Chen, *J. Pharm. Sci.* **97**, 1813 (2008).
- 2008VEN/BAY L. Ventola, L. Bayes, R. Benzges, F. J. Novegil-Anleo, M. A. Cuevas-Diarte, T. Calvet, and D. Mondieig, *Helv. Chim. Acta* **91**, 1286 (2008).

- 2008VER/EME S. P. Verevkin and V. N. Emel'yanenko, *Fluid Phase Equilib.* **266**, 64 (2008).
- 2008VER/EME2 S. P. Verevkin, V. N. Emel'yanenko, and S. A. Kozlova, *J. Phys. Chem. A* **112**, 10667 (2008).
- 2008VER/KOZ S. P. Verevkin and S. A. Kozlova, *Thermochim. Acta* **471**, 33 (2008).
- 2008VER/KOZ2 S. P. Verevkin, S. A. Kozlova, V. N. Emel'yanenko, P. Goodrich, and C. Hardacre, *J. Phys. Chem. A* **112**, 11273 (2008).
- 2008VER/TOK S. P. Verevkin, A. V. Toktonov, Y. Chernyak, B. Schaffner, and A. Borner, *Fluid Phase Equilib.* **268**, 1 (2008).
- 2008WAN/TAN S.-X. Wang, Z.-C. Tan, Y.-S. Li, B. Tong, Y. Li, Q. Shi, and J.-N. Zhang, *Chin. J. Chem.* **26**, 2016 (2008).
- 2008WAS/HOL C. M. Wassvik, A. G. Holmen, R. Draheim, P. Artursson, and C. A. S. Bergstrom, *J. Med. Chem.* **51**, 3035 (2008).
- 2008WEI D. Wei, *Thermochim. Acta* **479**, 32 (2008).
- 2008WIS/BER S. Wishkerman and J. Bernstein, *Chem.-Eur. J.* **14**, 197 (2008).
- 2008WU/JIA C. Wu, J. Jiang, G. Lia, and H. Qiu, *Huaxue Gongcheng* **36**, 47 (2008); *Chem. Abstr.* **150**, 475170 (2008).
- 2008XIA/ZHA Q. Xia, F.-B. Zhang, G.-L. Zhang, J.-C. Ma, and L. Zhao, *J. Chem. Eng. Data* **53**, 838 (2008).
- 2008XIN/TAN J. Xing, Z.-C. Tan, Q. Shi, B. Tong, S.-X. Wang, and Y.-S. Li, *J. Therm. Anal. Calorim.* **92**, 375 (2008).
- 2008ZAI/PAU D. H. Zaitsau, Y. U. Paulechka, G. J. Kabo, A. V. Blokhin, V. N. Emel'yanenko, S. P. Verevkin, and A. Heintz, *J. Chem. Eng. Data* **53**, 694 (2008).
- 2008ZHA/UNH H. Zhao, P. Unhannanant, W. Hanshaw, and J. S. Chickos, *J. Chem. Eng. Data* **53**, 1545 (2008).
- 2008ZHA/WAN G. Q. Zhang, L. S. Wang, R.-L. Fan, X.-Z. Shao, and X.-F. Wang, *J. Chem. Eng. Data* **53**, 1192 (2008).
- 2008ZHA/WUML. Zhang, E. Wumanjian, Q. Meng, and G. Cai, *Yingyong Huagong* **37**, 802 (2008).
- 2008ZHE/MOR K. V. Zherikova, N. B. Morozova, L. N. Zelenina, S. V. Sysoev, T. P. Chusova, and I. K. Igumenov, *J. Therm. Anal. Calorim.* **92**, 729 (2008).
- 2008ZHE/MOR2 K. V. Zherikova, N. B. Morozova, I. A. Baidina, S. V. Sysoev, and I. K. Igumenov, *J. Struct. Chem.* **49**, 58 (2008).
- 2008ZHU/XU R. Zhu, X. Xu, Z. Dong, and Y. Tian, *Fluid Phase Equilib.* **272**, 60 (2008).
- 2009ACE/NIC J. M. Aceves-Hernandez, I. Nicolas-Vazquez, F. J. Aceves, J. Hinojosa-Torres, M. Paz, and V. M. Castano, *J. Pharm. Sci.* **98**, 2448 (2009).
- 2009ARO/MAL S. Arockiasamy, C. Mallika, O. M. Sreetharan, V. S. Raghunathan, and K. S. Nagaraja, *Inorg. Chim. Acta* **362**, 1977 (2009).
- 2009BAR/ESP M. Barrio, P. Espeau, J. L. Tamarit, M.-A. Perrin, N. Veglio, and R. Ceolin, *J. Pharm. Sci.* **98**, 1657 (2009).
- 2009BAR/GAM C. Baraldi, M. C. Gamberini, A. Tinti, F. Palazzoli and V. Ferioli, *J. Mol. Struct.* **918**, 88 (2009).
- 2009BEL/RAZ A. Belabbaci, A. Razzouk, I. Mokbel, J. Jose, and L. Negadi, *J. Chem. Eng. Data* **54**, 2312 (2009).
- 2009BRA/GEL D. E. Braun, T. Gelbrich, V. Kahlenberg, R. Tessadri, J. Wieser, and U. J. Griesser, *J. Pharm. Sci.* **98**, 20 (2009).
- 2009BRI H. G. Brittain, *Cryst. Growth Des.* **9**, 2492 (2009).
- 2009BRI2 H. G. Brittain, *Cryst. Growth Des.* **9**, 3497 (2009).
- 2009BRU/PIA B. Brunetti, V. Piacente, and P. Scardala, *J. Chem. Eng. Data* **54**, 2273 (2009).
- 2009BUT/BUC A. B. Butrow, J. H. Buchanan, and D. E. Trevault, *J. Chem. Eng. Data* **54**, 1876 (2009).
- 2009CAM/EME H. Cammenga, V. N. Emel'yanenko, and S. P. Verevkin, *Ind. Eng. Chem. Res.* **48**, 10120 (2009).
- 2009CHE/SU W. Chen, B. Su, H. Xing, Y. Yang, and Q. Ren, *Fluid Phase Equilib.* **287**, 1 (2009).
- 2009CHE/XIA S. Chen, Q. Xia, D. Li, W.-G. Yan, F.-B. Zhang, and G.-L. Zhang, *J. Chem. Eng. Data* **54**, 1395 (2009).
- 2009CHI/ACR J. S. Chickos and W. E. Acree, Jr., *Thermochim. Acta* **495**, 5 (2009) and supplemental information.
- 2009CHI/STE R. D. Chirico and W. V. W. V. Steele, *J. Chem. Thermodyn.* **41**, 392 (2009).
- 2009CLA/GOM R. A. Clara, A. C. Gomez Mariagliano, and H. N. Solimo, *J. Chem. Eng. Data* **54**, 1087 (2009).
- 2009DAS/BRE E. da Silva, S. Bresson, and D. Rousseau, *Chem. Phys. Lipids* **157**, 113 (2009).
- 2009DOM/POB U. Domanska, A. Pobudkowska, A. Pelczarska, and P. Gierycz, *J. Phys. Chem. B* **113**, 8941 (2009).
- 2009EFI/DRU A. A. Efimova, A. I. Druzhinina, R. M. Varushchenko, O. V. Dorofeeva, and E. L. Krasnyh, *J. Chem. Eng. Data* **54**, 2457 (2009).
- 2009EME/VER V. N. Emel'yanenko, S. P. Verevkin, E. N. Burakova, G. N. Roganov, and M. K. Georgieva, *Russ. J. Phys. Chem.* **83**, 598 (2009).
- 2009FLO/CAM H. Flores, E. A. Camarillo, and J. Mentado, *Thermochim. Acta* **493**, 76 (2009).
- 2009FRE/GOM V. L. S. Freitas, J. R. B. Gomes, and M. D. M. C. Ribeiro da Silva, *J. Chem. Thermodyn.* **41**, 1199 (2009).
- 2009FRE/GOM2 V. L. S. Freitas, J. R. B. Gomes, and M. D. M. C. Ribeiro da Silva, *J. Therm. Anal. Calorim.* **97**, 827 (2009).
- 2009FRE/MON V. L. S. Freitas, M. J. S. Monte, L. M. V. B. F. Santos, J. R. B. Gomes, and M. D. M. C. Ribeiro da Silva, *J. Phys. Chem. A* **113**, 12988 (2009).
- 2009GAI/KUN A. Gairola, G. V. Kunte, A. M. Umarji, and S. A. Shivasankar, *Thermochim. Acta* **488**, 17 (2009).
- 2009GBA/NEG G. Gbabode, P. Negrier, D. Mondieig, E. Moreno, T. Calvet, and M. A. Cuevas-Diarte, *J. Alloys Compd.* **469**, 539 (2009).
- 2009GIE/KOS P. Gierycz, A. Kosowski, and R. Swietlik, *J. Chem. Eng. Data* **54**, 2996 (2009).
- 2009GIM/CLE N. Gimeno, M. J. Clemente, P. Forcen, J. L. Serrano, and M. B. Ros, *New J. Chem.* **33**, 2007 (2009).
- 2009GOM/MONJ. R. B. Gomes, A. R. Monteiro, B. B. Campos, P. Gomes, and M. D. M. C. Ribeiro da Silva, *J. Phys. Org. Chem.* **22**, 17 (2009).
- 2009GOO/ROD D. J. Good and N. Rodriguez-Hornedo, *Cryst. Growth Des.* **9**, 2252 (2009).
- 2009HA/HAN J.-M. Ha, B. D. Hamilton, M. A. Hillmyer, and M. D. Ward, *Cryst. Growth Des.* **9**, 4766 (2009).
- 2009HE/ZHU Q. He, J. Zhu, H. Goma, M. Jennings, and S. Rohani, *J. Pharm. Sci.* **98**, 1835 (2009).
- 2009JOH/SEL M. G. Johnson, J. Selvakumar, and K. S. Nagaraja, *Thermochim. Acta* **495**, 38 (2009).
- 2009KRO/DRU O. V. Krol, A. I. Druzhinina, R. M. Varushchenko, M. D. Reshetova, N. E. Borisova, O. V. Dorofeeva, and S. I. Troyanov, *J. Chem. Eng. Data* **54**, 263 (2009).
- 2009LAH/RAU A. Lahde, J. Raula, J. Malm, E. I. Kauppinen, and M. Karppinen, *Thermochim. Acta* **482**, 17 (2009).
- 2009LEG/BAC B. Legendre, P. Bac, M. German, and Y. Feutelais, *J. Therm. Anal. Calorim.* **98**, 91 (2009).
- 2009LI/FAN D. Li, W. Fang, W. Xie, Y. Wing, Y. Guo, and R. Lin, *Energy Fuels* **23**, 794 (2009).
- 2009LIP/CHI D. Lipkind and J. S. Chickos, Abstract from 38th Great Lakes Regional Meeting of the American Chemical Society, Chicago, IL, 13-16 May 2009 (unpublished). Abstract available online through Scifinder Scholar.
- 2009LIP/CHI2 D. Lipkind and J. S. Chickos, *Struct. Chem.* **20**, 49 (2009).
- 2009LIP/HAN D. Lipkind, W. Hanshaw, and J. S. Chickos, *J. Chem. Eng. Data* **54**, 2930 (2009) and references cited.
- 2009MAR/AUC B. Marrufo, A. Aucejo, M. Sanchoello, and S. Loras, *Fluid Phase Equilib.* **279**, 11 (2009).
- 2009MAR/LLA N. F. Martinez, E. Lladosa, M. C. Burguet, J. B. Monton, and M. Yazimon, *Fluid Phase Equilib.* **277**, 49 (2009).
- 2009MAR/SEL M. Maria, J. Selvakumar, V. S. Raghunathan, and K. S. Nagaraja, *Surf. Coat. Technol.* **204**, 222 (2009).
- 2009MAT/SOU M. A. R. Matos, C. C. S. Sousa, and V. M. F. Morais, *J. Chem. Thermodyn.* **41**, 308 (2009).
- 2009MAT/SOU2 M. A. R. Matos, C. C. S. Sousa, and V. M. F. Morais, *J. Chem. Thermodyn.* **41**, 69 (2009).
- 2009MAT/SOU3 M. A. R. Matos, C. C. S. Sousa, and V. M. F. Morais, *J. Chem. Eng. Data* **54**, 2189 (2009).
- 2009MAT/SOU4 M. A. R. Matos, C. C. S. Sousa, M. S. Miranda, V. M. F. Morais, and J. F. Liebman, *J. Phys. Chem. B* **113**, 11216 (2009).
- 2009MEL/PIM S. V. Melkhanova, S. M. Pimenova, N. V. Chelovskaya, E.

- A. Miroshichenko, L. L. Pashehenko, I. A. Nesterov, and P. V. Naumkin, *J. Chem. Thermodyn.* **41**, 651 (2009).
- 2009MEN/TAN Q. Meng, Z. Tan, X. Wang, Y. Dong, W. Li, and Q. Shi, *Chin. J. Chem.* **27**, 1225 (2009).
- 2009MOK/RAZ I. Mokhel, A. Razzouk, T. Sawaya, and J. Jose, *J. Chem. Eng. Data* **54**, 819 (2009).
- 2009MON/NUT D. Monnier, I. Nuta, C. Chatillon, M. Gross-Jean, F. Volpi, and E. Blanquet, *J. Electrochem. Soc.* **156**, H71 (2009).
- 2009MOR/SEM N. B. Morozova, P. P. Semyannikov, S. V. Trubin, P. P. Stabnikov, A. A. Bessonov, K. V. Zherikova, and I. K. Igumenov, *J. Therm Anal. Calorim.* **96**, 261 (2009).
- 2009MOR/ZHE N. B. Morozova, K. V. Zherikova, P. P. Semyannikov, S. V. Trubin, and I. K. Igumenov, *J. Therm Anal. Calorim.* **98**, 395 (2009).
- 2009MOT/CAR F. L. Mota, A. P. Carneiro, A. J. Queimada, S. P. Pinho, and E. A. Macedo, *Eur. J. Pharm. Sci.* **37**, 499 (2009).
- 2009NEZ/AER A. Nezzal, L. Aert, M. Verspaille, G. Hendrickx, and A. Redl, *J. Cryst. Growth* **311**, 3863 (2009).
- 2009NTI/CHA J. Nti-Gyabaah, V. Chan, and Y. C. Chiew, *Fluid Phase Equilib.* **280**, 35 (2009).
- 2009OJA/CHE V. Oja, X. Chen, M. R. Hajaligol, and W. G. Chan, *J. Chem. Eng. Data* **54**, 730 (2009).
- 2009OLI/BER P. R. Oliveira, L. S. Bernadi, F. S. Murakami, C. Mendes, and M. A. S. Silva, *J. Therm Anal. Calorim.* **97**, 741 (2009).
- 2009PAN/ANT K. Panneerselvam, M. P. Antony, T. G. Srinivasan, and P. R. V. Rao, *Thermochim. Acta* **495**, 1 (2009).
- 2009PEN/ESC M. A. Pena, A. Escalera, A. Reillo, A. B. Sanchez, and P. Bustamante, *J. Pharm. Sci.* **98**, 1129 (2009).
- 2009PER/MAG R. A. Perkins and J. W. Magee, *J. Chem. Eng. Data* **54**, 2646 (2009).
- 2009PER/OCH R. A. Perkins, J. C. S. Ochoa, and J. W. Magee, *J. Chem. Eng. Data* **54**, 3192 (2009).
- 2009PER/TKA G. L. Perlovich, V. V. Tkachev, N. N. Strakhova, V. P. Kazachenko, T. V. Volkova, O. V. Surov, K.-J. Schaper, and O. A. Raevsky, *J. Pharm. Sci.* **98**, 4738 (2009).
- 2009POS/MAR M. A. Postigo, A. B. Mariano, A. F. Jara, and N. Zurakowski, *J. Chem. Eng. Data* **54**, 1575 (2009).
- 2009QUE/MOT A. J. Queimada, F. L. Mota, S. P. Pinho, and E. A. Macedo, *J. Phys. Chem. B* **113**, 3469 (2009).
- 2009RAI/RED R. N. Rai and R. S. B. Reddi, *Thermochim. Acta* **496**, 13 (2009).
- 2009RAZ/NAC A. Razzouk, R. A. Naccoul, I. Mokbel, J. Saab, and J. Jose, *J. Chem. Eng. Data* **54**, 1214 (2009).
- 2009RAZHAJ A. Razzouk, A. Hajjaji, I. Mokbel, P. Mougain, and J. Jose, *Fluid Phase Equilib.* **282**, 11 (2009).
- 2009RIB/AMA M. A. V. Ribeiro da Silva and L. M. P. F. Amaral, *J. Chem. Thermodyn.* **41**, 26 (2009).
- 2009RIB/AMA2 M. A. V. Ribeiro da Silva and L. M. P. F. Amaral, *J. Chem. Thermodyn.* **41**, 349 (2009).
- 2009RIB/CAB M. A. V. da Silva Ribeiro and J. I. T. A. Cabral, *J. Chem. Thermodyn.* **41**, 355 (2009).
- 2009RIB/CAB2 M. A. V. Ribeiro da Silva, J. I. T. A. Cabral, and J. R. B. Gomes, *J. Chem. Thermodyn.* **41**, 1193 (2009).
- 2009RIB/FER M. A. V. Ribeiro da Silva and A. I. M. C. L. Ferreira, *J. Chem. Eng. Data* **54**, 2517 (2009).
- 2009RIB/FER2 M. A. V. Ribeiro da Silva and A. I. M. C. L. Ferreira, *J. Chem. Thermodyn.* **41**, 361 (2009).
- 2009RIB/FER3 M. A. V. Ribeiro da Silva and A. I. M. C. L. Ferreira, *J. Chem. Thermodyn.* **41**, 499 (2009).
- 2009RIB/FER4 M. A. V. Ribeiro da Silva and A. I. M. C. L. Ferreira, *J. Chem. Thermodyn.* **41**, 1096 (2009).
- 2009RIB/FER5 M. A. V. Ribeiro da Silva and A. I. M. C. L. Ferreira, *J. Chem. Thermodyn.* **41**, 1104 (2009).
- 2009RIB/FER6 M. A. V. Ribeiro da Silva, A. I. M. C. L. Ferreira, J. I. T. A. Cabral, A. F. L. M. O. Santos, A. R. G. Moreno, T. L. P. Galvao, I. M. Rocha, P. M. V. Fernandes, S. Q. Salgueiro, V. A. F. de Moura, I. M. S. C. Oliveira, P. C. Cotelto, and M. R. A. Ribeiro, *J. Chem. Thermodyn.* **41**, 984 (2009).
- 2009RIB/FER7 M. A. V. Ribeiro da Silva, A. I. M. C. L. Ferreira, and A. R. G. Moreno, *J. Chem. Thermodyn.* **41**, 109 (2009).
- 2009RIB/FER8 M. A. V. Ribeiro da Silva, A. I. M. F. Ferreira, A. F. L. O. M. Santos, and I. M. Rocha, *J. Chem. Thermodyn.* **41**, 1239 (2009).
- 2009RIB/FER9 M. A. V. Ribeiro da Silva, A. I. M. C. L. Ferreira, and A. R. G. Moreno, *J. Chem. Thermodyn.* **41**, 904 (2009).
- 2009RIB/RIB M. A. V. Ribeiro da Silva, M. D. M. C. Ribeiro da Silva, A. I. M. C. L. Ferreira, A. F. L. O. M. Santos, and T. L. P. Galvao, *J. Chem. Thermodyn.* **41**, 1074 (2009).
- 2009RIB/RIB2 M. A. V. Ribeiro da Silva, M. D. M. C. Ribeiro da Silva, A. I. M. C. L. Ferreira, A. F. L. O. M. Santos, and T. L. P. Galvao, *J. Chem. Thermodyn.* **41**, 1247 (2009).
- 2009RIB/RIB3 M. D. M. C. Ribeiro da Silva, M. A. V. Ribeiro da Silva, V. L. S. Freitas, M. V. Roux, P. Jimenez, M. Temprado, J. Z. Davalas, P. Cabildo, R. M. Calramunt, and J. Elguero, *J. Chem. Thermodyn.* **41**, 1400 (2009).
- 2009RIB/SAN M. A. V. Ribeiro da Silva, A. F. L. O. M. Santos, J. R. B. Gomes, M. V. Roux, M. Temprado, P. Jimenez, and R. Notario, *J. Phys. Chem. A* **113**, 11042 (2009).
- 2009RIB/SAN2 M. A. V. Ribeiro da Silva and A. F. L. O. M. Santos, *J. Chem. Thermodyn.* **41**, 926 (2009).
- 2009ROU/TEM M. V. Roux, M. Temprado, P. Jimenez, C. Foces-Foces, R. Notario, A. R. Parameswar, A. V. Demchenko, J. S. Chicos, C. A. Deakyne, A. K. Ludden, and J. F. Liebman, *J. Phys. Chem. A* **113**, 10772 (2009).
- 2009SAN/FIG R. C. Santos, R. M. B. B. M. Figueira, M. F. M. Piedade, H. P. Diogo, and M. E. Minas da Piedade, *J. Phys. Chem. B* **113**, 14291 (2009).
- 2009SAN/GOM A. F. L. O. M. Santos, J. R. B. Gomes, and M. A. V. Ribeiro da Silva, *J. Phys. Chem. A* **113**, 3630 (2009).
- 2009SAN/RIB A. F. L. O. M. Santos and M. A. V. Ribeiro da Silva, *J. Phys. Chem. A* **113**, 9741 (2009).
- 2009SAP/UUS2 E. Sapei, P. Uusi-Kyyny, K. I. Keskinen, and J. Aittamaa, *Fluid Phase Equilib.* **279**, 81 (2009).
- 2009SEL/RAG J. Selvakumar, V. S. Raghunathan, and K. S. Nagaraja, *J. Phys. Chem. C* **113**, 19011 (2009).
- 2009SID/SID M. A. Siddiqi, R. A. Siddiqui, and B. Atakan, *J. Chem. Eng. Data* **54**, 2795 (2009).
- 2009SIN/MUR L. P. Singh and S. S. N. Murthy, *Phys. Chem. Chem. Phys.* **11**, 5110 (2009).
- 2009SOU/MAT C. C. S. Sousa, M. A. R. Matos, and V. M. F. Morais, *J. Chem. Thermodyn.* **41**, 1408 (2009).
- 2009SUR/SZT A. O. Surov, P. Szterner, W. Zielenkiewicz, and G. L. Perlovich, *J. Pharm. Biomed. Anal.* **50**, 831 (2009).
- 2009SUR/TER A. O. Surov, I. V. Terekhova, A. Bauer-Brandl, and G. L. Perlovich, *Cryst. Growth Des.* **9**, 3265 (2009).
- 2009SZT P. Szterner, *J. Therm Anal. Calorim.* **98**, 337 (2009).
- 2009TAM/MIR R. E. Tamagawa, E. A. Miranda, C. C. Santana, and M. Giulietti, *J. Chem. Eng. Data* **54**, 16 (2009).
- 2009TAU/SIT P. Taulelle, G. Sitja, G. Pepe, E. Garcia, C. Hoff, and S. Veessler, *Cryst. Growth Des.* **9**, 4706 (2009).
- 2009TOR/CAM L. A. Torres, M. Campos, M. Martinez, and A. Rojas, *J. Chem. Thermodyn.* **41**, 957 (2009).
- 2009VAD/BAN H. K. Vaddi, S. L. Banks, J. Chen, D. C. Hammell, P. A. Crooks, and A. L. Stinchcomb, *J. Pharm. Sci.* **98**, 2611 (2009).
- 2009VAN/WES J. Th. H. van Eupen, R. Westheim, M. A. Deij, A. Meekes, P. Bennema, and E. Vlieg, *Int. J. Pharm.* **368**, 146 (2009).
- 2009VEC/BRU S. Vecchio and B. Brunetti, *J. Chem. Thermodyn.* **41**, 880 (2009).
- 2009VEC/TOM S. Vecchio and M. Tomassetti, *Fluid Phase Equilib.* **279**, 64 (2009).
- 2009VER/EME S. P. Verevkin, V. N. Emel'yanenko, E. N. Stepurko, R. V. Rulys, D. H. Zaitsau, and A. Stark, *Ind. Eng. Chem. Res.* **48**, 10087 (2009).
- 2009VER/EME2 S. P. Verevkin, V. N. Emel'yanenko, A. V. Toktonov, A. S. Leolko, J. Duwensee, U. Kragl, and S. M. Sarge, *Ind. Eng. Chem. Res.* **48**, 7388 (2009).
- 2009VER/EME3 S. P. Verevkin, V. N. Emel'yanenko, A. V. Toktonov, P. Goodrich, and C. Hardacre, *J. Phys. Chem. B* **113**, 12704 (2009).
- 2009VER/EME4 S. P. Verevkin, V. N. Emel'yanenko, A. V. Toktonov, P. Goodrich, and C. Hardacre, *Ind. Eng. Chem. Res.* **48**, 9809 (2009).

- 2009VER/KOZ S. P. Verevkin, S. A. Kozlova, V. N. Emel'yanenko, and G. Nell, *J. Chem. Thermodyn.* **41**, 1125 (2009).
- 2009WEI/CHE D. Wei and L. Chen, *Fluid Phase Equilib.* **277**, 9 (2009).
- 2009WEI/JIN D. Wei and K. Jin, *J. Chem. Thermodyn.* **41**, 145 (2009).
- 2009WU/MEH W. Wu, S. J. Mehrman, Y. Zhou, S. X. Pu, L. Huang, A. Fermier, and S. Karki, *J. Cryst. Growth* **311**, 3435–3444 (2009).
- 2009XIN/FAN Y. Xing, W. Fang, D. Li, Y. Guo, and R. Lin, *J. Chem. Eng. Data* **54**, 1865 (2009).
- 2009ZAI/PAU D. H. Zaitsau, Y. U. Paulenchka, A. V. Blokhin, A. V. Yermalayeu, A. G. Kabo, and M. R. Ivanets, *J. Chem. Eng. Data* **54**, 3026 (2009).
- 2009ZEN/CAO J. L. Zeng, Z. Cao, D. W. Yang, F. Xu, L. X. Sun, L. Zhang, and X. F. Zhang, *J. Therm Anal. Calorim.* **95**, 501 (2009).
- 2010AND/ABU G. P. Andrews, O. A. Abudiak, and D. S. Jones, *J. Pharm. Sci.* **99**, 1322 (2010).
- 2010BAR/ARA G. L. Barros de Araju, D. L. Araujo de Faria, M. H. Zaim, F. M. de Souza Carvalho, F. R. Dias de Andrade, and J. do Rosario Matos, *J. Therm Anal. Calorim.* **102**, 233 (2010).
- 2010BRU/PIA2 B. Brunetti, V. Piacente, and P. Scardala, *J. Chem. Eng. Data* **55**, 98 (2010).
- 2010CAB/MON J. I. T. A. Cabral, R. A. R. Monteiro, M. A. A. Rocha, L. M. N. B. F. Santos, W. E. Acree, Jr., and M. D. M. C. Ribeiro da Silva, *J. Adv. Oxid. Technol.* **100**, 431 (2010).
- 2010CHA/LAY E. L. Charsley, P. G. Laye, and T. Le Goff, *Thermochim. Acta* **497**, 72 (2010).
- 2010CHI/STE R. D. Chirico, A. F. Kazakov, and W. V. Steele, *J. Chem. Thermodyn.* **42**, 581 (2010).
- 2010CHI/STE2 R. D. Chirico, A. F. Kazakov, and W. V. Steele, *J. Chem. Thermodyn.* **42**, 571 (2010).
- 2010CIL/ALB F. Cilirzo, E. Alberti, P. Minghetti, C. G. M. Gennari, A. Casiraghi, and L. Montanari, *Int. J. Pharm.* **386**, 71 (2010).
- 2010DON/WU H. Dong, C. Wu, X. Yang, and G. Lai, *J. Chem. Eng. Data* **55**, 889 (2010).
- 2010EFI/EME A. A. Efimova, V. N. Emel'yanenko, S. P. Verevkin, and Y. Chernyak, *J. Chem. Thermodyn.* **42**, 330 (2010).
- 2010FAN/WAN C. L. Fan and L. S. Wang, *J. Chem. Eng. Data* **55**, 479 (2010).
- 2010FRA/SAL L. S. Franco, Y. P. Salamanca, M. Maldonado, and E. F. Vargas, *J. Chem. Eng. Data* **55**, 1042 (2010).
- 2010FRE/GOM V. L. S. Freitas, J. R. B. Gomes, and M. D. M. C. Ribeiro da Silva, *J. Chem. Thermodyn.* **42**, 251 (2010).
- 2010FUL/MOR M. Fulem, P. Moravec, J. Pangrac, E. Hulicius, T. Simceek, K. Ruzicka, V. Ruzicka, B. Kozyrkin, and V. Shatunov, *J. Chem. Eng. Data* **55**, 362 (2010).
- 2010GAO/LIN G.-Y. Gao and S.-Y. Lin, *J. Pharm. Sci.* **99**, 256 (2010).
- 2010GRI/TSA J. R. Griffiths, J. Tsanaktsidis, G. P. Savage, and R. Priefer, *Thermochim. Acta* **499**, 15 (2010).
- 2010GUO/SAD K. Guo, G. Sadiq, C. Seaton, R. Davey, and Q. Yin, *Cryst. Growth Des.* **10**, 268 (2010).
- 2010GUO/WAN X.-Z. Guo, L.-S. Wang, and N.-N. Tian, *J. Chem. Eng. Data* **55**, 1745 (2010).
- 2010HON/HUA J. Hong, D. Hua, X. Wang, H. Wang, and J. Li, *J. Chem. Eng. Data* **55**, 297 (2010).
- 2010HU/WAN F.-H. Hu, L.-S. Wang, and S.-F. Cai, *J. Chem. Eng. Data* **55**, 492 (2010).
- 2010KES/AUC V. Kestens, G. Auclair, K. Drozdowska, A. Held, G. Roebben, and T. Linsinger, *J. Therm Anal. Calorim.* **99**, 245 (2010).
- 2010KRZ/MAL K. Krzyminski, P. Malecha, P. Storoniak, B. Zadykiewicz, and J. Blazejowski, *J. Therm Anal. Calorim.* **100**, 217 (2010).
- 2010KUN/SAV S. Kunadharaju and M. Savva, *J. Chem. Eng. Data* **55**, 103 (2010).
- 2010LAZ/RIE M. Lazerges, I. V. Rietveld, Y. Corvis, R. Ceolin, and P. Espeau, *Thermochim. Acta* **497**, 124 (2010).
- 2010LUT/MAT K. M. Lutker and A. J. Matzger, *J. Pharm. Sci.* **99**, 794 (2010).
- 2010MAO/LUO Z.-B. Mao, T.-L. Luo, T.-B. Cui, Y. Wang, and G.-J. Liu, *J. Chem. Eng. Data* **55**, 543 (2010).
- 2010MAR/LOR B. Marrufo, S. Loras, and M. Sanchoello, *J. Chem. Eng. Data* **55**, 62 (2010).
- 2010MEJ/SEG A. Mejia, H. Segura, and M. Cartes, *J. Chem. Eng. Data* **55**, 428 (2010).
- 2010MON/ALM M. J. S. Monte, A. R. R. P. Almeida, and M. A. R. Matos, *J. Chem. Eng. Data* **55**, 2246 (2010).
- 2010MON/GON M. J. S. Monte, M. V. Goncalves, and M. D. M. C. Ribeiro da Silva, *J. Chem. Eng. Data* **55**, 2246 (2010).
- 2010MUE/ESC S. Muela, B. Escalera, M. A. Pena, and P. Bustamante, *Int. J. Pharm.* **394**, 93 (2010).
- 2010MUR/PIK S. B. Murdande, M. J. Pikal, R. M. Shanker, and R. H. Bogner, *J. Pharm. Sci.* **99**, 1254 (2010).
- 2010PAN/SAR E. Paneteli, P. Saratsioti, H. Stamatis, and E. Voutsas, *J. Chem. Eng. Data* **55**, 745 (2010).
- 2010PET/REY M. Petitjean, E. Reyes-Perez, D. Perez, Ph. Mariabel, and C. Le Calve, *J. Chem. Eng. Data* **55**, 852 (2010).
- 2010RIB/FER M. A. V. Ribeiro da Silva, A. I. M. C. L. Ferreira, A. F. L. O. M. Santos, C. M. A. Ferreira, D. C. B. Barros, J. A. C. Reis, J. C. S. Costa, M. M. G. Calvino, S. I. A. Rocha, S. P. Pinto, S. S. L. Freire, S. M. Almeida, V. S. Guimaraes, and V. N. M. Almeida, *J. Chem. Thermodyn.* **42**, 371 (2010).
- 2010RIB/FER2 M. A. V. Ribeiro da Silva and A. I. M. C. Ferreira, *J. Chem. Thermodyn.* **42**, 182 (2010).
- 2010RIB/FER3 M. A. V. Ribeiro da Silva, A. I. M. C. L. Ferreira, and F. M. Maciel, *J. Chem. Thermodyn.* **42**, 220 (2010).
- 2010RIB/FER4 M. A. V. Ribeiro da Silva, A. I. M. L. Ferreira, A. F. L. O. M. Santos, and I. M. Rocha, *J. Chem. Thermodyn.* **42**, 169 (2010).
- 2010RIB/GON M. D. M. C. Ribeiro da Silva, M. V. Goncalves, and M. J. S. Monte, *J. Chem. Thermodyn.* **42**, 472 (2010).
- 2010RIB/RIB M. A. V. Ribeiro da Silva, M. D. M. C. Ribeiro da Silva, A. F. L. O. M. Santos, A. I. M. C. Ferreira, and T. L. P. Galvao, *J. Chem. Thermodyn.* **42**, 496 (2010).
- 2010RIB/RIB2 M. D. M. C. Ribeiro da Silva, M. A. V. Ribeiro da Silva, V. L. S. Freitas, M. V. Rous, P. Jimenez, J. Z. Davalos, P. Cabildo, R. M. Claramunt, E. Pinilla, M. R. Torres, and J. Elguero, *J. Chem. Thermodyn.* **42**, 536 (2010).
- 2010RIB/SAN M. A. V. Ribeiro da Silva and A. F. L. O. M. Santos, *J. Chem. Thermodyn.* **42**, 128 (2010).
- 2010RIB/SAN2 M. A. V. Ribeiro da Silva, L. M. N. B. F. Santos, and L. M. S. S. Lima, *J. Chem. Thermodyn.* **42**, 134 (2010).
- 2010RIB/SAN3 M. A. V. Ribeiro da Silva, A. F. L. O. M. Santos, and L. M. P. F. Amarsal, *J. Chem. Thermodyn.* **42**, 564 (2010).
- 2010SAP/UUS E. Sapei, P. Uusi-kyyny, K. I. Keskinen, and V. Alopaeus, *Fluid Phase Equilib.* **288**, 155 (2010).
- 2010SVA/NOR M. Svard, F. L. Nordstrom, T. Jasnobulka, and A. C. Ras-muson, *Cryst. Growth Des.* **10**, 195 (2010).
- 2010SZT/LEG P. Szterner, B. Legendre, and M. Sghaier, *J. Therm Anal. Calorim.* **99**, 325 (2010).
- 2010TON/LIU B. Tong, R.-B. Liu, C.-G. Meng, F.-Y. Yu, S.-H. Ji, and Z.-C. Tan, *J. Chem. Eng. Data* **55**, 119 (2010).
- 2010TON/YU B. Tong, Y. Yu, Z.-C. Tan, C.-G. Meng, L.-J. Cui, G. Xia and R.-B. Liu, *Thermochim. Acta* **499**, 117 (2010).
- 2010WID/BRU J. A. Widegren and T. J. Bruno, *J. Chem. Eng. Data* **55**, 159 (2010).
- 2010ZEN/GEL N. Zencirci, T. Gelbrich, D. C. Apperley, R. K. Harris, V. Kalenberg, and U. J. Briesser, *Cryst. Growth Des.* **10**, 302 (2010).
- UR/CHI J. S. Chickos (unpublished).
- UR/DEK C. G. de Kruif (unpublished).
- UR/FUC R. H. Fuchs (unpublished).
- UR/MAC H. Mackle, W. V. Steele, and D. V. McNally (unpublished).
- UR/MCC J. W. McCargar and W. E. Acree, Jr. (unpublished).
- UR/STU D. Stull (unpublished).
- UR/VER S. P. Verevkin, E. Krasnykh, T. Vasiltsova, and A. Heintz, unpublished data as quoted in [2002VER].