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Differentiation Of Murine Erythroleukemic Cells During Exposure To Microwave Radiation

Roger F. Brown Missouri University of Science and Technology, rbrown@mst.edu

S. V. Marshall Missouri University of Science and Technology

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Table VII. Ultrasonic Velocity at 10 MHz in Mixtures of Sodium Dodecanoate in Dodecanoic Acid

<i>T</i> , K	<i>u</i> , m/s	10 ⁶ β _s , bar ⁻¹	<i>T</i> , K	<i>u</i> , m/s	10 ⁶ β _s , bar ⁻¹
		$x_{i} = 0$	9490		
318.4	1280.,	69.3	328.2	1248.	73.5
321.5	1272.	70.4	332.2	1237.	75.1
324.4	1262.2	71.7	335.7	1222.	77.2
		$x_{1} = 0$.8600		
327.2	1258.	71.4	337.7	1224.	76.2
330.2	1249.	72.6	341.4	1214.	77.7
333.7	1235.4	74.5		2	
		$x_{1} = 0$.7030		
333.2	1250	71.1	340.2	1230	73.8
336.7	1240.	72.5	344.2	1215.	76.1

the formula $\beta_s = (u^2 d)^{-1}$ is fitted by the equation $10^6 \beta_s = -87.7$ + 0.43517 + 13.368(1 + 0.001397) x_1 , where β_s is expressed in bar⁻¹, with a standard deviation of 0.2.

Acknowledgment

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Glossarv

d	density
η	coefficient of viscosity
β_s	coefficient of adiabatic compressibility
m	molality
М	molar weight
n	number of moles

- correlation coefficient
- cation radius r_+ Ν
 - Avogadro's number
- mole fraction of component 1 in the eutectic mixture *х* _{1Е} mole fraction of component 1 in the peritectic mix-X 1P
 - ture
- T^0 melting temperature of the pure compound
- $T_{\rm E}$ eutectic temperature
- T_{P} peritectic temperature
- и ultrasonic velocity
- ν V_m molar volume = $(\partial V / \partial n_j)_{T,P,n_j}$
- mean volume = $V/(n_1 + n_2)$
- Vintr intrinsic volume

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Relative Volatilities of the Ethane–Ethylene System from Total Pressure Measurements

David A. Barclay, Judith L. Flebbe, and David B. Manley*

Chemical Engineering Department, University of Missouri — Rolla, Rolla, Missouri 65401

A new apparatus has been developed for making vapor pressure measurements on high-pressure systems. Using this equipment, we have made a large number of new total pressure measurements on the ethane-ethylene binary from 198.15 to 278.15 K. By integration of the general coexistence equation, relative volatilities have been calculated for ethane-ethylene over the entire temperature range.

The viability of using the total pressure method for determining relative volatilities in systems at high pressure has been shown by Manley (1) and Manley and Swift (2). Walker (3) and Steele (4, 5) have developed an apparatus capable of producing rapid and accurate P-T-X measurements for moderate-pressure systems at near ambient temperatures. Barclay (6) incorporated the desirable features of that equipment into an apparatus useful at higher pressures and lower temperatures and provides new data on a system of economic significance.

No recent study of the ethane-ethylene system has been made which covers the entire region of industrial interest. Fredenslund et al. (7) produced high-quality VLE measurements for two isotherms, 263.15 and 293.15 K, the second being above the critical point of ethylene. Earlier measurements by Hanson et al. (8) at 273.15, 233.17, and 199.85 K are not of comparable accuracy. The results of this study should improve the consistency and the accuracy of the relative volatility information on the ethane-ethylene system and aid in the design of distillation columns for processing these two chemicals.

Experimental Equipment

The equipment developed to measure vapor pressures is very similar to that described by Walker and Steele. It was designed to operate at pressures up to 4.5 MPa with the capability of being easily modified to work almost to 7.0 MPa.

A major equipment modification was the redesign of the sample cell-transducer combination. Figure 1 is a proportional drawing of these.

The cell is constructed of 316 stainless steel with a 0.002-in. type 302 stainless diaphragm and is assembled by silver solder brazing. The cavity is machined from 2-in. disks approximately ¹/₈ in. thick. The diameter of the cavity is 1.50 in., and it is cut at a lathe setting of 1.5° for the entire radius to give a maximum span at the center of about 0.031 in. The sample chamber is made from 1-in. o.d. tubing and has an approximate



Figure 1. Cell and transducer for vapor pressure measurements.

volume of 6 cm³. The thermowell provides a way to get accurate temperature measurements of the fluid in the cell. Tare weight of the cell plus the valve attached to the fill line is less than 220 g. The maximum working pressure is about 7.0 MPa.

The transducer which attaches to the cell serves to sense the position of the cell diaphragm, to support the cell in the temperature bath, and to connect the upper cell cavity to the nitrogen system used for balancing the diaphragm. The push rod is located inside the connecting line and rests on the diaphragm. Movement of the diaphragm causes the rod to move a small powdered iron core connected to its upper end. a Trans-Tek Model 240-000 displacement transducer senses the core position. Output from the transducer is read on a digital millivoltmeter.

In actual operation, the cell-transducer combination is calibrated as a unit to find the diaphragm null as a function of temperature and pressure. The unit is sensitive to pressure differentials of less than 68.95 Pa at pressures up to 3.0 MPa. Sensitivity appears to decrease slightly at higher pressures. After a complete "overpressure" on the diaphragm of 7.0 MPa, the calibration accuracy is retained to better than \pm 138 Pa.

Constant temperature was provided in an air bath as shown in Figure 2. A small squirrel cage fan circulates air down the center or working space of the bath and then forces it back up the annulus where the cooling coil and the heaters are located. An FTS Systems two-stage refrigeration unit provides cooling. The heaters are bare wire nichrome heating elements. A Bayley proportional band controller is used for temperature control. The inner and outer shells are cast acrylic tubing. They measure 6×18 and 9×30 in., respectively. The internal parts are supported by six 0.25-in. stainless threaded rods while the outer shell is held only by the end plates and may be removed for easy access to the interior. The bath is insulated by stacking shaped foam planks around it. The range of op-



Figure 2. Air bath.



Figure 3. Schematic of system for vapor pressure measurements.

eration is -80 to 20 °C, and temperature control of the cells located in the bath was better than ± 0.01 °C.

Loading of cells proceeded in the following manner. The evacuated tare of the cell was established to better than 0.1 mg by replicate weighings over a period of 12 h. The cell was attached to a small fill manifold which connected the cell, the sample container, and a gauge to the vacuum-vent system. After evacuation, the system was flushed several times with the component to be charged to the cell. Sample was transferred to the cell by first filling the gauge and connecting lines with fluid from the sample cylinder, closing it off, and then opening the fill valve on the cell. The amount of sample was then determined by weighing the cell and contents. For binary mixtures ethane was added first with the sample cylinder being inverted to get liquid sample. Gentle heat was applied to increase cylinder pressure and improve the transfer of liquid ethane to the cell. This was not necessary for ethylene. When ethylene was added, the above process was repeated. The manifold would be filled with ethylene to a pressure above that in the cell and the cell valve then cracked open briefly to allow some transfer into the cell. The manifold was then filled to a higher pressure than before and the process repeated. When the maximum fill pressure was reached, the cell was weighed to find the amount of ethylene added.

A transducer was attached to the cell and the unit was placed in the air bath and connected to the pressure measuring system shown in Figure 3. After the cell was brought to temperature in the bath and the rough pressure found by using the gauge on the nitrogen system, the cell null was calculated. The displacement pump was used to make minor pressure adjustments to bring the diaphragm in the cell to a final null. The pressure was measured through the Ruska null cell by a Ruska oil dead-weight gauge.

Temperatures were measured by using Rosemount Model 146MA100F platinum resistance thermometers placed in the thermowells of the cells. Each thermometer was calibrated to IPTS-68 by comparison with a standard thermometer calibrated by NBS. Resistances were measured by using a Rubicon Instruments Mueller type bridge and a L & N dc null detector.

The ethane and the ethylene used in this study were research-grade purity, listed as 99.96% and 99.95% pure, respectively, purchased from Phillips Petroleum Co. Samples of each were prepared by subjecting them to a series of freezethaw cycles to remove traces of air. Chromatographic analyses of the gas phase of the ethane and of the ethylene revealed no detectable traces of air in either. This amounts to less than 6 ppm by weight. No other impurities were detected.

Theory

The rigorous form of the isothermal binary Gibbs-Duhem equation was integrated to find vapor compositions from the total pressure measurements

$$\frac{\mathrm{d}y}{\mathrm{d}x} = \frac{\left\{ y(1-y)\frac{\Delta' V}{(y-x)} - y(1-y) \left[\frac{\partial \ln (\phi_1/\phi_2)}{\partial P} \right]_y \right\} \frac{\mathrm{d}P}{\mathrm{d}x}}{1 + y(1-y) \left[\frac{\partial \ln (\phi_1/\phi_2)}{\partial y} \right]_P}$$
(1)

where the partial derivatives of ln (ϕ_1/ϕ_2) are constrained to constant vapor composition or constant pressure as indicated and

$$\Delta' V = (1/RT)(V^{\mathsf{v}} - V^{\mathsf{h}}) \tag{2}$$

The development of eq 1 has been given by Manley and Swift (2), who also derive the limiting forms as x approaches the composition extremes

$$\lim_{x \to 0} \frac{dy}{dx} = \lim_{x \to 0} \alpha = 1 + \Delta' V (\frac{dP}{dx})$$
(3)

$$\lim_{x \to 1} \frac{dy}{dx} = \lim_{x \to 1} \alpha = \frac{1}{[1 - \Delta' V (dP/dx)]}$$
(4)

In the ideal total pressure measurement experiment, the cell is filled with liquid so only a small vapor space remains. The overall cell composition, which is measured, then very closely approximates the liquid composition. For most cases a correction of the overall composition, z, to the liquid composition, x, is required. Using the material balance and phase relations for the contents of the cell, the correction of z to x is easily found

$$x = z\rho_{\rm o}/[X\rho_{\rm i} - X\rho_{\rm v}K + K\rho_{\rm v}]$$
⁽⁵⁾

where

$$X = (\rho_{o} - \rho_{v})/(\rho_{1} - \rho_{v})$$
 (5a)

Using eq 1 and 5 the integration at each isotherm is combined with the composition corrections as Gibbs and Van Ness (9) have indicated. The initial P-z data are curve fitted, and the integration is performed. With this initial estimate of K's

Table I. a Constant^a for Redlich-Kwong Equation

<i>T</i> , K	ethane	ethylene	<i>T</i> , K	ethane	ethylene
278.15	0.6095	0.4737	218.15	0.7226	0.5576
263.15	0.6349	0.4933	203.15	0.7548	0.5818
248.15	0.6646	0.5130	198.15	0.7660	0.5902
233.15	0.6921	0.5347			

^a Units: (L/mol)² MPa.

and y values, eq 5 is applied to give corrected x values. The process is repeated until the x values not longer change within established error bounds. Convergence was found to occur generally within 2-3 iterations.

The integration algorithm was an Adams–Moulton predictor– corrector procedure of order 5 (10). A fourth-order Runge– Kutta procedure was used to start the integration which proceeded in the direction of increasing pressure.

Curve fitting of the isothermal P-x data was done with an equation based on deviations from Raoult's law. A three-constant Redlich-Kister equation was used to fit the deviations

$$n (P/p_{\rm R}) = x(1-x)[B + C(2_x - 1) + D(2_x - 1)^2] \quad (6)$$

where $P_{\rm R}$ is the Raoult's law pressure. Equation 6 provided both the total pressure and its derivatives for the integration.

For saturated vapor volumes and the partial derivatives involving the fugacity coefficients, a Redlich–Kwong-type equation was used (11)

$$P = \frac{RT}{V-b} - \frac{a}{V(V+b)}$$
(7)

where

$$b_i = 0.08664RT_c/P_c$$
 (8)

and the *a* constant was adjusted at each isotherm to give the proper vapor compressibility factors for the pure components. The established mixing rules were relied upon for the mixture properties.

$$a = \sum_{i} y_{i} \left[\sum_{j} y_{i} (a_{j} a_{j})^{1/2} (1 - \delta_{ij}) \right]$$
(9)

$$b = \sum_{i} y p_i \tag{10}$$

Saturated liquid volumes were calculated from the correlation by Hankinson and Thomson (12)

$$V_{\rm s}/V^{*} = V_{\rm R}^{0}[1 - \omega V_{\rm R}^{\delta}]$$
(11)

where $V_{\rm R}^0$ and $V_{\rm R}^{\delta}$ are given as functions of reduced temperature. The adjustable parameter, V^* , was used to calibrate the equation to the pure-component volumes at each temperature. Their suggested mixing rules were used to calculate saturated volumes for mixtures of ethane and ethylene.

The pure-component P-V-T data on both ethane and ethylene given by Douslin and Harrison (13, 14) were used to calibrate the equations for saturated vapor and liquid volumes. These data were chosen not only for their accuracy and precision but for the fact that the saturated liquid and vapor volumes are internally consistent. Since eq 1 requires the difference between liquid and vapor volumes, this is very important.

Table I gives the *a* constants used in the vapor equation of state. The critical temperatures and pressures used were those given by Douslin and Harrison: for ethane, $T_c = 305.33$ K and $P_c = 4.8717$ MPa; for ethylene, $T_c = 282.35$ K and $P_c = 5.0419$ MPa. The interaction coefficient was estimated as 0.0125 by fitting the data of Fredenslund et al. (7) with the Soave-Redlich-Kwong VLE algorithm.

The liquid volume constants used are in Table II. Accentric factors were 0.0983 for ethane and 0.882 for ethylene as

Table II. Characteristic Volumes for Saturated Liquid Density Equation



Figure 4. Literature comparisons for ethane.

suggested in the original correlation.

Results

Vapor pressures were measured for both pure ethane and ethylene and 13 mixtures at 7 temperatures. Each set of measurements was at constant overall composition. Duplicate points were taken for each temperature, the first while proceeding in the direction of decreasing temperature and the second while proceeding with increasing temperature. Two separate sets of measurements were made on the pure components, one at the beginning of the study and one at the end, using different cells and different thermometers. This helped to establish the precision of the equipment and verified the purity of the ethane and ethylene. All of the raw data were corrected to even temperatures before further use.

In order to make the correction from overall composition to liquid composition, it was necessary to find the volumes of the cells. Calibration of the cell volumes as a function of temperature and pressure was done with compressed liquid propane. The densities of propane were taken from Ely and Kobayashi (*15*).

Figure 4 shows the comparison for ethane vapor pressures of more recent literature data with this study. The data of Douslin and Harrison (13), Straty and Tsumura (16), Pal et al. (17), and this work all show general agreement within $\pm 0.25\%$. The API-44 (18) correlation represents earlier data. Reference pressures were calculated by using the Goodwin equation (20) fitted to the data from this investigation

$$\ln (P/P_r) = A \chi + B \chi^2 + C \chi^3 + D \chi (1-\chi)^{\epsilon}$$
(12)

where

$$\chi = (1 - T_{\rm b}/T)/(1 - T_{\rm b}/T_{\rm c})$$
(13)

 P_r , T_b , T_c , and ϵ were set at 0.101325 MPa, 184.563 K, 305.33 K, and 1.43, respectively. The calculated constants are A = 3.4245, B = 0.5643, C = -0.11647, and D = 0.53079. The equation was used only for the literature comparisons.

Pure-component ethylene literature comparisons are in Figure 5. The Goodwin constants found were A = 3.4172, B = 0.6557, C = -0.16647, and D = 0.58704. $T_{\rm b}$, $T_{\rm c}$, $P_{\rm r}$, and ϵ were set at 169.389 K, 282.35 K, 0.101325 MPa, and 1.46 as



Figure 5. Literature comparisons for ethylene.



Figure 6. Raoult's law deviations for ethane--ethylene binary.

Douslin and Harrison suggest. All deviations between data sets are within $\pm 0.2\%$.

Table III shows the measured vapor pressures for the ethane-ethylene binary. The estimated probable error in pressure is given by (0.0132%)P + 0.00024 MPa. Errors in temperature are believed to be within ± 0.01 K. Pressures are reported to more places than accuracy justifies to retain internal consistency. Total moles in the cell and overall composition remain constant for all temperatures. The estimated error in overall mole fractions is ± 0.0001 mole fraction units. The cell volumes are believed to be within 0.2%. The corrected x values are estimated to be accurate to ± 0.0005 mole fraction units as an average value. Since the estimate is a function of the amount of liquid in the cell, this value can vary. In the region of high ethylene composition at 278.15 and 263.15 K, which is near the critical point, the errors in x may be larger. These should average less than 0.5% of x.

The Redlich–Kister constants used to fit the P-x data for each isotherm are in Table IV. These were used in eq 6 for integration of the coexistence equation.

Figures 6–9 show the deviations from Raoult's law for all of the isotherms. These illustrate the precision of the data. For comparison, the Redlich-Kister curve fit is also plotted.

Results of the numerical integration of eq 1 are in Table V. Figure 10 is a plot of these same results.

To investigate the effect of systematic errors and errors in the calculated physical properties from the correlations on the



Figure 7. Raoult's law deviations for ethane-ethylene binary.



Figure 8. Raoult's law deviations for ethane-ethylene binary.

integration, we deliberately introduced several types of errors into the integration. (1) The pure-component vapor pressure of ethylene was changed by 0.1%. A similar error in ethane should have the same effect. (2) The saturated vapor volume equation, eq 7, was adjusted to give vapor compressibility factors for the pure components which were in error by 0.6%. (3) The pressure equation was adjusted to give pressures in error by 0.1% at x = 0.5. Other types of errors were assumed to be negligible compared to these. The results are shown in Table VI.

Effects of the errors vary with temperature and composition. The error introduced in vapor compressibilities dominates at 278.15 K while the error in total pressure is largest at 198.15 K. The average error is 0.4% of α .

Figure 11 is a comparison of results of this study with literature data. Those isotherms which correspond to relative volatilities derived directly from the pressure measurements are shown as solid lines. The dashed lines were interpolated by



Figure 9. Raoult's law deviations for ethane-ethylene binary.



Figure 10. Relative volatility of ethylene in ethane.

using the pure-component equations (eq 12) and by curve fitting the constants in Table IV with temperature. Integration of the coexistence equation then used these interpolated constants and pressures to calculate the intermediate isotherms. The data of Fredenslund et al. (7) and Hanson et al. (8) are results of sampling liquid and vapor phases at equilibrium.

Agreement is generally within the combined experimental error. At 255.38 K, the data of Hanson appear to be high at the ethylene end of the isotherm.

Conclusions

Experimental equipment has been developed which may be used to measure vapor pressures almost to 7.0 MPa and temperatures between -80 and 20 °C. This should be useful in generating more accurate and precise data on high-pressure

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Table III.	Vapor Pressure	s for the	Ethane-Ethylene	Binary

x	press., M	IPa	cell vol, cm ³	total mol in cell	overall mole fractio of ethylene	n 9	x	press., M	IPa	cell vol, cm ³	total mol in cell	overall mole fraction of ethylene
0.0 0.0458 0.1233 0.2274 0.3703 0.3982 0.4067 0.5194	2.691 22, 2.6 2.691 91, 2.6 2.780 35, 2.7 2.928 90, 2.9 3.131 61, 3.1 3.399 14, 3.3 3.449 70, 3.4 3.465 02, 3.4 3.678 17, 3.6	592 18 592 17 780 92 930 42 130 50 399 41 447 97 467 39 579 28	6.230 5.906 6.237 6.241 5.903 6.242 6.246	0.042 169 0.042 274 0.068 138 0.063 423 0.057 226 0.053 484 0.056 838	0.0483 0.1289 0.22934 0.3740 0.4029 0.4142 0.5255	278.15	K 0.5478 0.6298 0.7440 0.8154 0.9113 0.9677 1.00	3.728 10, 3. 3.884 98, 3. 4.097 49, 4.1 4.235 57, 4. 4.417 72, 4. 4.525 81, 4. 4.590 00, 4. 4.588 89, 4.	725 47 886 09 097 65 232 14 417 11 525 78 591 03 589 96	5.901 6.250 5.899 5.898 6.259 6.261	0.049 741 0.054 091 0.046 900 0.043 044 0.044 138 0.039 679	0.5557 0.63646 0.7508 0.8220 0.9150 0.9695
	X	p	ress., MPa	v	cell ol, cm ³			x	p	ress., MP	8	cell vol, cm ³
	0.0	20 1.861 1.861 1.929	63.15 K 14, 1.861 40, 1.861 64, 1.930	. 79 . 67 . 42	6.218			0.0	0.46 0.46 0.48	218.15 K 0 36, 0.46 0 48, 0.46 5 54, 0.48	50 43 50 50 35 69	6.199
	0.1240 0.2269 0.3699 0.3983 0.4070	2.043 2.195 2.396 2.431 2.443	45, 2.043 00, 2.194 54, 2.394 75, 2.431	22 01 71	6.222 6.226 5.909 6.227			0.1264 0.2278 0.3716 0.4002 0.4106	0.52 0.57 0.64 0.65	5 70, 0.52 6 50, 0.52 2 83, 0.64 5 74, 0.65	25 79 76 57 42 88 55 32	5.921 6.201 6.202 5.920 6.202
	0.5194 0.5486 0.6300 0.7447	2.598 2.637 2.748 2.901	375, 2.600 42, 2.637 72, 2.749 29, 2.901	003 702 965 116	6.229 5.908 6.232 5.906			0.5224 0.5519 0.6333 0.7478	0.70 0.72 0.75 0.80	8 31, 0.70 0 73, 0.72 5 09, 0.75 0 34, 0.80	08 37 21 25 54 95 00 50	6.202 6.203 5.920 6.204 5.919
	0.8163 0.9116 0.9679 1.00	2.996 3.126 3.200 3.244 3.244	35, 2.996 02, 3.126 60, 3.201 57, 3.245 25, 3.245	552 526 59 503 503	5.906 6.239 6.240			0.8193 0.9136 0.9688 1.00	0.82 0.86 0.88 0.89 0.90	9 45, 0.82 6 11, 0.86 7 36, 0.88 9 67, 0.89 0 83 0 89	29 53 56 15 37 53 99 84 99 87	5.919 6.206 6.206
	0.0	24 1.235 1.235 1.235	48.15 K 74, 1.235 93, 1.236	5 79 5 00 7 4 2	6 209			0.0	0.25 0.25 0.25	203.15 K 0 65, 0.25 0 75, 0.25	50 33 50 78	6 107
	0.1247 0.2270 0.3702 0.3984	1.287 1.370 1.480 1.624 1.652	0 27, 1.28 0 27, 1.37 0 24, 1.48 0 54, 1.62 2 09, 1.65) 88) 15 5 38 2 5 9	5.916 6.212 6.215 5.914			0.1272 0.2282 0.3723 0.4010	0.20 0.29 0.32 0.36 0.37	1 83, 0.29 3 49, 0.32 4 33, 0.36 2 44, 0.37	91 90 23 55 54 38 72 48	5.922 6.198 6.199 5.922
	0.4081 0.5202 0.5492 0.6309	1.661 1.771 1.799 1.877	29, 1.661 44, 1.772 65, 1.799 73, 1.878	1 84 2 16 9 44 3 03	6.215 6.217 5.913 6.219			0.4117 0.5234 0.5531 0.6343	0.37 0.40 0.41 0.43	4 61, 0.37 4 05, 0.40 1 67, 0.41 1 94, 0.43	74 58 04 01 11 63 32 06	6.199 6.200 5.922 6.200
	0.7454 0.8170 0.9123 0.9682 1.0	1.984 2.050 2.139 2.190 2.219 2.222	02, 1.984 76, 2.05(20, 2.139 67, 2.19(94, 2.22(204, 2.22(419 068 066 093 039 051	5.912 5.911 6.224 6.224			0.7488 0.8202 0.9141 0.9691 1.00	0.45 0.47 0.49 0.51 0.51	8 98, 0.45 6 71, 0.47 8 02, 0.49 0 61, 0.51 7 97, 0.51 8 52, 0.51	59 05 76 66 98 06 10 66 17 99 18 89	5.921 5.921 6.201 6.202
	0.00	2: 0.779 0.779	33.15 K 915, 0.779 906, 0.779	9 1 3 9 3 1				0.0	0.19 0.20	198.15 9 21, 0.20 0 31, 0.20	K)0 00)0 33	
	0.0468 0.1255 0.2274 0.3709 0.3993 0.4094 0.5213	0.814 0.875 0.951 1.052 1.071 1.077	54, 0.816 37, 0.875 91, 0.951 34, 1.052 63, 1.072 97, 1.078	5 33 5 56 1 92 2 51 2 25 3 25 3 31	6.203 5.919 6.205 6.207 5.917 6.207 6.207			0.0476 0.1274 0.2284 0.3725 0.4012 0.4120 0.5237	0.21 0.23 0.26 0.29 0.30 0.30	3 40, 0.21 4 91, 0.23 1 49, 0.26 5 56, 0.29 1 96, 0.30 4 02, 0.30 8 47 0 37	13 39 34 88 51 47 95 61 92 27 94 07 28 49	6.197 5.923 6.198 6.199 5.922 6.199 6.199
	0.5506 0.6321 0.7467 0.8182 0.9130	1.172 1.225 1.296 1.341 1.400	2 29, 1.172 5 34, 1.225 5 75, 1.297 5 87, 1.341 5 42, 1.400	2 45 5 45 7 17 1 98 0 64	5.917 6.210 5.916 5.916 6.213			0.5535 0.6346 0.7490 0.8205 0.9142	0.32 0.33 0.35 0.37 0.38 0.40	5 24, 0.32 5 24, 0.33 1 65, 0.35 3 89, 0.37 8 64, 0.38 6 09, 0.40	35 25 51 67 73 90 38 19 96 10	5.922 6.200 5.922 5.922 6.201
	0.9686 1.00	1.434 1.453 1.455	51, 1.434 66, 1.454 43, 1.454	+ 42 + 20 + 23	6.213			0.9691 1.00	0.410 0.422 0.422	5 57, 0.41 2 55, 0.42 2 85, 0.42	16 54 22 52 22 88	6.201

and low-temperature systems for design and correlation work. A significant number of new vapor pressure measurements have been made on the ethane-ethylene binary. By integration of the general coexistence equation, relative volatilities have been determined over the range 198.15-278.15 K with a probable error of $\pm 0.4\%$.

Glossary

κ	y/x	
Ρ	pressure,	MPa

Table IV. Constants for Total Pressure Equation

	pure-compone	ent press., MPa	1000(Redlich-Kister eq constants)			
<i>Т</i> , К	ethane	ethylene	B	С	D	
278.15	2.691 87	4.589 97	-0.199 512 8	-17.406 32	4.372 535	<u> </u>
263.15	1.861 50	3.244 71	29.998 38	-26.62831	4.513 233	
248.15	1.235 86	2.220 72	54.79805	-36.285 65	5.796 761	
233.15	0.77916	1.454 38	80.019 92	-52.05210	9.858 286	
218.15	0.460 45	0.90006	108.491 7	-73.57278	21.324 98	
203.15	0.25063	0.51834	140.0844	-103.8521	30.00963	
198.15	0.19997	0.422 70	155.0662	-116.0000	35.684 77	

Table V. Results of Integration of Binary Gibbs-Duhem Equation

x	у	Р	V^1	Ζ	α	x	У	Р	$V^{\mathbf{i}}$	Ζ	α
		278.1	15 K			0.5000	0.6206	1.139 34	0.061 72	0.8200	1.636
0.0000	0.0000	2.691 87	0.077 34	0.6525	1.409	0.6000	0.7051	1.204 36	0.061 50	0.8130	1.594
0.1000	0.1329	2.885 96	0.07771	0.6364	1.379	0.7000	0.7839	1.267 89	0.061 30	0.8062	1.555
0.2000	0.2523	3.077 30	0.07818	0.6196	1.350	0.8000	0.8587	1.33043	0.06112	0.7996	1.519
0.3000	0.3615	3.266 41	0.07876	0.6022	1.321	0.9000	0.9305	1.39245	0.060 95	0.7931	1.487
0.4000	0.4629	3.453 97	0.079 47	0.5842	1.293	1.0000	1.0000	1.454 38	0.060 81	0.7867	1.462
0.5000	0.5586	3.64073	0.080 36	0.5653	1.265						
0.6000	0.6500	3.827 50	0.081 47	0.5455	1.238			218.1	5 K		
0.7000	0.7387	4.01508	0.082 87	0.5243	1.212	0.0000	0.0000	0.460 45	0.06013	0.9022	2.027
0.8000	0.8258	4.204 24	0.084 72	0.5013	1.185	0.1000	0.1805	0.51269	0.059 81	0.8960	1.982
0.9000	0.9124	4.395 67	0.087 23	0.4757	1.158	0.2000	0.3258	0.56262	0.05949	0.8899	1.933
1.0000	1.0000	4.5 89 9 7	0.090 95	0.4462	1.128	0.3000	0.4463	0.61017	0.05917	0.8840	1.880
		262.1	6.77			0.4000	0.5491	0.655 52	0.058 87	0.8784	1.827
0 0000	0.0000	203.1	LD K.	0.7264	1 6 4 4	0.5000	0.6395	0.69895	0.05857	0.8729	1.774
0.0000	0.0000	1.861.50	0.070.98	0.7364	1.544	0.6000	0.7209	0.740 85	0.058 29	0.8677	1.722
0.1000	0.1439	2.009.60	0.07098	0.7240	1.513	0.7000	0.7961	0.781 59	0.058 01	0.8627	1.674
0.2000	0.2703	2.154 49	0.071.01	0.7115	1.481	0.8000	0.8669	0.821 55	0.057 74	0.8578	1.629
0.3000	0.3833	2.296 33	0.071.09	0.6988	1.450	0.9000	0.9346	0.86098	0.05748	0.8531	1.588
0.4000	0.4862	2.435 45	0.071.21	0.6862	1.419	1.0000	1.0000	0. 9 00 06	0.057 23	0.8485	1.549
0.5000	0.5815	2.57233	0.071 38	0.6735	1.390			203.1	5 K		
0.6000	0.6712	2.707.53	0.07161	0.6608	1.361	0.000	0.0000	0.250.63	0.05772	0 0360	2 246
0.7000	0.7568	2.84170	0.071 91	0.6480	1.334	0.0000	0.0000	0.23003	0.05736	0.9309	2.240
0.8000	0.8396	2.975 51	0.072 29	0.6353	1.309	0.1000	0.1901	0.205 52	0.057.01	0.9322	2.195
0.9000	0.9205	3.109.63	0.07277	0.6223	1.286	0.2000	0.3400	0.344.16	0.057.01	0.9277	2.133
1.0000	1.0000	3.244 71	0.073 36	0.6091	1.265	0.5000	0.5716	0.371.90	0.05631	0.9233	2.070
		248.1	15 K			0.1000	0.6590	0.398.19	0.055.98	0.9152	1 933
0.0000	0.0000	1.235.86	0.066.47	0.8022	1 681	0.5000	0.0350	0.423 33	0.055.65	0.9132	1.955
0.1000	0.1548	1.344 90	0.066.30	0.7926	1 649	0.0000	0.8079	0.423 55	0.055.32	0.9113	1.800
0.2000	0.2875	1.450.98	0.066.14	0 7829	1 614	0.000	0.8748	0 471 43	0.055.00	0.9045	1 747
0.3000	0.4037	1.554.08	0.066.01	0.7732	1 579	0.0000	0.9386	0 4 9 4 9 5	0.053.69	0.9043	1.698
0.4000	0.5073	1 654 35	0.065.89	0.7637	1 544	1 0000	1 0000	0.51834	0.054 38	0.9011	1.658
0.5000	0.6015	1.752.13	0.065.80	0.7543	1 509	1.0000	1.0000	0.010 04	0.004 50	0.0770	1.050
0.6000	0.6888	1.847.84	0.065 73	0.7451	1 476			198.1	5 K		
0.7000	0.7711	1.942.00	0.065.69	0.7361	1 444	0.0000	0.0000	0.19997	0.056 99	0.9465	2.335
0.8000	0.8498	2.03515	0.065.69	0 7273	1 4 1 4	0.1000	0.2022	0.22772	0.05662	0.9423	2.281
0.9000	0.9259	2 1 2 7 8 7	0.065.72	0.7186	1 388	0.2000	0.3566	0.253 99	0.056 26	0.9382	2.217
1.0000	1.0000	2 220 72	0.065.78	0.7099	1 366	0.3000	0.4790	0.27865	0.055 90	0.9342	2.145
1.0000	110000	2.220 / 2	0.000 / 0	0.7077	1.500	0.4000	0.57 9 8	0.301 80	0.055555	0.9305	2.070
		233.1	5 K			0.5000	0.6660	0.32364	0.055 20	0.9270	1.994
0.0000	0.0000	0.77916	0.06298	0.8578	1.840	0.6000	0.7422	0.344 45	0.054 86	0.9237	1.920
0.1000	0.1669	0.856 49	0.06271	0.8500	1.802	0.7000	0.8119	0.364 53	0.054 52	0.9206	1.850
0.2000	0.3059	0.93115	0.06244	0.8422	1.763	0.8000	0.8773	0.38415	0.054 19	0.9176	1.788
0.3000	0.4245	1.00307	0.06219	0.8346	1.721	0.9000	0. 9398	0.403 51	0.053 87	0.9147	1.733
0.4000	0.5281	1.07237	0.061 9 5	0.8272	1.678	1.0000	1.0000	0.42270	0.053 55	0.9118	1.684

Table VI. Errors in Relative Volatility Generated by Errors in Correlations

	x = 0.10		x = 0.50		x = 0.90		x = 0.95			
	α	Δ	α	Δ	α	Δ	α	Δ	error	
					278.15 K					
	1.3792		1.2653		1.1575		1.1431			
	1.3800	0.0008	1.2654	0.0001	1.1566	0.0009	1.1419	0.0012	1	
	1.3819	0.0027	1.2680	0.0027	1.1595	0.0020	1.1449	0.0018	2	
	1.3810	0.0018	1.2648	0.0005	1.1557	0.0018	1.1414	0.0017	3	
sum		0.0053		0.0033		0.0047		0.0047		
	•				198.15 K					
	2.2808		1.9939		1.7335		1.7089			
	2.2828	0.0020	1.9957	0.0018	1.7348	0.0013	1.7101	0.0012	1	
	2.2792	0.0016	1.9921	0.0018	1.7320	0.0015	1.7075	0.0014	2	
	2.2846	0.0038	1.9957	0.0018	1.7275	0.0060	1.7012	0.0077	3	
sum		0.0074		0.0054		0.0088		0.0103		



Figure 11. Literature comparison of relative volatilities for ethaneethylene system.

reference pressure in Goodwin equation Raoult's law pressure
critical pressure
gas constant
temperature
critical temperature
volume, L/mol
liquid mole fraction, unsubscripted means most volatile component
vapor mole fraction, unsubscripted means most volatile component
overall mole fraction of ethylene in sample cell
vapor compressibility factor
relative volatility
binary interaction parameter
density
fugacity of component / in a mixture

Subscripts

v

- 1,2 components in binary system, 1 is most volatile
- liquid 1
- overali n
- P constraint on partial derivative to constant pressure vapor
- constraint on partial derivative to constant compov sition

Superscripts

- liquid
- vapor

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Osmotic and Activity Coefficients of Some Sulfamates and Sulfanilates at 298.15 K

Oscar D. Bonner

Department of Chemistry, University of South Carolina, Columbia, South Carolina 29208

Osmotic and activity coefficients are reported for solutions of the sodium and potassium saits of sulfamic and sulfanilic acids and for the parent acid. The data are in agreement with the evidence in the literature that sulfamic acid is a moderately strong acid with an ionization constant which is not too different from that of iodic acid.

are quite weak or quite strong, there are few acids with ionization constants in the range of 10⁻¹-10. A recent survey article (1) reminded one that sulfamic and sulfanilic acids exist as zwitterions in solution and suggested that a study of the properties of solutions of these acids and their salts might be of interest. The ionization constant for the equilibrium

Moderately strong acids such as iodic or the trihaloacetic acids are a small and unique group in that, while many acids

$$NH_3^+SO_3^- = NH_2SO_3^- + H^+$$

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has been reported to be 0.1006 from conductance measure-