CROATICA CHEMICA ACTA CCACAA 81 (2) 273–275 (2008) ISSN-0011-1643 CCA-3240 Original Scientific Paper

The Melting Points of the Inert Gas Solids

Lawrence J. Dunne,^{a,b} John N. Murrell,^{b,*} and George Manos^c

^aDepartment of Engineering Systems, London South Bank University, London SE1 0AA, UK

^bDepartment of Chemistry, University of Sussex, Falmer, Brighton, BN1 9QJ, UK

^cDepartment of Chemical Engineering, University College London, Torrington Place, London WC1E 7JE, UK

RECEIVED NOVEMBER 26, 2007; REVISED FEBRUARY 13, 2008; ACCEPTED FEBRUARY 15, 2008

Keywords heavy inert gases melting points diatomic potential wells Lindemann's theory of melting

The melting points of the heavy inert gases and of some other simple molecules show an excellent linear correlation with the depths of their diatomic potential wells, and the slope of the correlation line is in accord with Lindemann's theory of melting

INTRODUCTION

Here is a simple question.^{\neq} Why do the melting points of the inert gases (Ne–Xe) increase as we ascend the periodic table? A common response is 'they get heavier'. This is not the correct answer. Isotopes and isotopomers interact through the same potentials, and at the classical level, the Gibbs free energy of solids and liquids have a very weak (logarithmic) dependence on mass through their translational motion, yet they have very similar melting points. Moreover, at the classical level *p*-*V* isotherms are mass independent. Notably, mass is not a parameter in the Monte Carlo computer simulation of solid-liquid melting.

Here are some examples of melting points (in brackets) taken from Ref. 1 and the NIST physical properties data base:² CH₄ (90.6 K), CD₄ (89.8 K), C₆H₆ (278.6 K), C₆D₆ (280 K), H₂O (273.15 K), H₂O¹⁸ (273.4 K), D₂O (277 K). Significant increases in mass lead to changes in the melting point that are usually only about one percent

or less, sometimes up, sometimes down. Any differences are primarily due to vibrational zero point energies, which are a quantum effect. The largest mass difference in the above list is for H_2O/D_2O because hydrogen bonds have large zero point energies. Helium is also an exception because it is a quantum system.

An early, phenomenological theory of melting was given by Lindemann.³ This was based on the idea that if the average displacement of the atoms from their lattice sites exceeded a certain fraction of the lattice spacing (later studies showed this to be about 1/10), then above the sublimation pressure the solid would melt. This idea has been supported by theoretical calculations, and by neutron diffraction experiments on supercooled liquids and frozen glasses.^{4,5}

Lindemann used his idea to produce a mathematical expression for the melting point, which in the classical limit is indeed mass independent. Although very simple, Lindemann's assumption is still the basis of many melting models, as illustrated by a recent publication.⁶ In this

^{*} Author to whom correspondence should be addressed. (E-mail: j.n.murrell@sussex.ac.uk)

[≠] This question was posed by John N. Murrell at the Brijuni Conference on Interdisciplinary Topics in Chemistry and Physics 2006.

paper we go further and show that his model gives a simple and surprisingly accurate explanation of the melting points of inert gas solids, and indeed of some other simple solids, a result that to our knowledge has not been published before.

Although this paper emphasises melting points a similar mass dependence applies to boiling points as illustrated by a few examples:^{1,2,7} ²⁰Ne (27.084 K), ²²Ne (27.211 K), CH₄ (111.6 K), CD₄ (111.7 K), C₆H₆ (353 K), C₆D₆ (352 K).

A MODIFIED LINDEMANN MODEL

For small displacements from an equilibrium position we can restrict the potential to harmonic terms as

$$V(r) = -\varepsilon + (1/2)(r - r_{\rm e})^2 k \tag{1}$$

where ε is the depth of the well, r_e the equilibrium distance and k the force constant. Now k will usually depend on the form of the potential (e.g., the exponent in the Morse function), but for the well-established Lennard-Jones (m,n) family of potentials k depends only on ε and r_e . For example, for the (6,12) potential, which we can write in the form

$$V(r) = \varepsilon((r_e/r)^{12} - 2(r_e/r)^6)$$
(2)

$$k = 72\varepsilon/r_{\rm e}^2 \tag{3}$$

The equipartition theorem shows that at equilibrium both the potential and kinetic energies are equal to $k_{\rm B}T/2$, where $k_{\rm B}$ is Boltzmann's constant. The Lindemann hypothesis is that melting occurs when the root mean square displacement exceeds a fraction *f* of the equilibrium distance, hence we can make the equality

$$k_{\rm B} T_{\rm m} = k r_{\rm e}^2 f^2 \tag{4}$$

and for the (6,12) potential

$$T_{\rm m} = 72\varepsilon f^2 / k_{\rm B} \tag{5}$$

We see again that the melting point $T_{\rm m}$ is independent of mass and for the (6,12) potential (or others of that family), is proportional to ε .

Figure 1 shows that this prediction holds very well for the inert gases, and from the slope of the line we deduce f = 0.09, which is very close to the generally accepted Lindemann value of 0.1. The well depths were taken from the compilation given in Ref. 9.

The figure also includes data for O_2 , N_2 , CH_4 , and CF_4 , and although the simple (6,12) law is not expected



Figure 1.

to be as accurate for these systems, their melting points lie surprisingly close to the inert gas line.

A simple thermodynamic argument supports the idea that ε is the most important factor in determining melting points. At the melting point the Gibbs energy of solid and liquid are the same, hence we can write

$$T_{\rm m} = \Delta H_{\rm m} \,/\, \Delta S_{\rm m} \tag{6}$$

And as the volume change on melting is small, $\Delta H_{\rm m}$ is approximately equal to $\Delta U_{\rm m}$, which, in the case of the inert gas solids, will be proportional to ε . Thus the linear relation between $T_{\rm m}$ and ε in our figure is to be expected. However, the fact that the slope of the line is in accord with Lindemann's model has implications for $\Delta S_{\rm m}$, and suggests a common factor for the entropy of melting, just as there is for the entropy of vaporisation, (Trouton's rule). For the inert gases the entropies of fusion per atom lie between 1.64 $k_{\rm B}$ (Ne) and 1.71 $k_{\rm B}$ (Xe).⁷

REFERENCES

- R. C. Reid, J. M Prausnitz, and B. E Poling, *The Properties of Gases and Liquids*, 4th Ed., Mc GrawHill, New York, 1987.
- 2. NIST database (http://webbook.nist.gov/chemistry/).
- 3. F. A. Lindemann, Phys. Z. 11 (1910) 609.
- S. Rabinovich, D. Berrebi, and A. Voronel, J. Phys. Condensed Matter 1 (1989) 6881.
- 5. V. Lubchenko, J. Phys. Chem. B 110 (2006) 18779-18786.
- C. Chakravarty, P. G. Debendetti, and F. H. Stillinger, J. Chem. Phys. 126 (2007) 204508–204518.
- P. Haussinger, R. Glatthaar, W. Rhode, H. Kick, C. Benkmann, J. Weber, H. J. Wunschel, and V. Stenke, *Nobel Gases*, Ullmann's Encyclopedia of Industrial Chemistry, Wiley, March, 2001.

SAŽETAK

Tališta krutina plemenitih plinova

Lawrence J. Dunne, John N. Murrell i George Manos

Tališta krutina teških plemenitih plinova i nekih jednostavnih molekula linearno se koreliraju s dubinom njihovih potencijalnih jama, a nagib korelacijskoga pravca slaže s Lindemannovom teorijom taljenja.