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Thermochimica Acta xxx (2004) xxx-xxx

thermochimica acta

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4	The enthalpy of sublimation of cubane
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0	Received 1 March 2004; received in revised form 6 May 2004; accepted 8 May 2004
1	This paper is dedicated to Philip E. Eaton and Thomas W. Cole, Jr. on the occasion of the 40th anniversary
2	of the first successful synthesis of cubane and any of its derivatives

13 Abstract

The sublimation enthalpy of cubane, a key reference material for force field and quantum mechanical computations, was measured by combining the vaporization enthalpy at T = 298.15 K to the sum of the fusion enthalpy measured at T = 405 K and a solid-solid phase transition that occurs at T = 394 K. The fusion and solid-solid phase transitions were measured previously. A sublimation enthalpy value of (55.2 ± 2.0) kJ mol⁻¹ at T = 298.15 K was obtained. This value compares quite favorably the value obtained by comparing the sublimation enthalpy of similar substances as a function of their molar masses but is at odds with earlier measurements.

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20 Keywords: Cubane; Sublimation; Enthalpy

21 1. Introduction

This is the 40th anniversary of the first successful syn-22 thesis of the polycyclic hydrocarbon cubane [1]: two other 23 syntheses soon followed [2]. As befits the high symmetry 24 25 and accompanying esthetics, large strain energy and thus high energy, and eight tertiary carbons all capable of possi-26 ble functionalization, the chemistry of this seemingly sim-27 ple 8-carbon hydrocarbon and its derivatives has blossomed 28 as evidenced by numerous reviews in which it is featured 29 30 prominently [3]. The eponymic (i.e., cubical) symmetry of cubane results in there being a single type of carbon en-31 vironment and of bonded hydrogen, one type of C-C and 32 C-H bond and associated bond lengths, and one unique 33 C-C-C and C-C-H angle. Very few hydrocarbons have 34

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such a minimal description². Accordingly, paralleling the 35 "organic" chemistry (and related bio- and high energy chem-36 istry) interest in cubane and its derivatives, the physical 37 chemists have been active-soon after the first synthesis 38 of cubane itself there was a measurement of the enthalpies 39 of combustion and of sublimation of this hydrocarbon [4] 40 from which the gas phase enthalpy of formation of 622.2 41 \pm 4.2 kJ mol⁻¹ was derived. In turn, this quantity and the 42 molecular high symmetry have meant that molecular me-43 chanicians have been active: for example, cubane has been 44 important in the development of the recent molecular me-45 chanical model, MM4 [5a] and its predecessor, MM3 [5b] 46 and quantum chemists have likewise been active with high 47

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² The other known minimal hydrocarbons are methane, ethane, ethylene, acetylene, cyclopropane, neopentane, benzene, and dodecahedrane. Cyclohexane and cyclobutane, for example, do not qualify because there is a difference between equatorial and axial hydrogens; allene does not qualify because of two types of carbon, and cyclooctatetraene does not qualify because of two different types of carbon–carbon bonds. Ideally, diamond, graphite and polyethylene would qualify, however, there are end effects such as the finiteness of the sample.

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48 level methodologies [6a,b]. Calculations at the G2(MP2) 49 level give 606.7 or 625.9 kJ mol⁻¹ depending on whether the 50 atomization or bond separation method is used [6a], while 51 the G3(MP2) result is 610.9 kJ mol⁻¹, 11.3 kJ mol⁻¹ lower 52 than the reported experimental value [6b,c].

Recently using an analysis based on the sublimation enthalpies of other cyclic and polycyclic hydrocarbons, it has been suggested that the enthalpy of sublimation of cubane is seriously in error [7]. This raises considerable concern as to the enthalpy of formation of gaseous cubane and all related analyses. We recall that there is some controversy about the enthalpy of combustion, and thus formation of its

Table 1

Cubane mixture

1,4-dicarbomethoxy derivative [8]. The current study reports60a new experimental determination of the enthalpy of subli-
mation of cubane. Let us summarize our findings and anal-
yses that follow: the just enunciated literature suggestion is
verified and the derived concern is justified.61616262636364

2. Experimental

All standards were purchased from the Aldrich Chemical Company and were used without any further purification. Each was analyzed by gas chromatography and found 68

65

CH ₂ C1 ₂	$t_{\rm r}$ (min)						
	348.7	353.9	358.9	364	369.1	374.2	
(A) Mixture 1							
Methane ^a	1.328	1.349	1.353	1.373	1.341	1.359	
Nonane	3.87	3.105	2.826	2.621	2.449	2.312	
Cubane	5.485	4.25	3.799	3.453	3.168	2.935	
Decane	6.964	5.081	4.437	3.927	3.524	3.199	
exo-THDCPD ^b	10.105	7.271	6.286	5.495	4.865	4.35	
Undecane	13.53	9.102	7.645	6.488	5.594	4.88	
endo-THDCPD ^b	12.247	8.658	7.404	6.488	5.645	4.993	
(B) Mixture 2							
	343.6	348.6	353.8	358.8	363.9	368.9	374
Methane ^a	1.27	1.283	1.29	1.342	1.344	1.357	1.375
Norbornene	1.957	1.879	1.809	1.76	1.71	1.68	1.665
Octane	2.481	2.302	2.154	2.04	1.947	1.875	1.83
Cubane	5.472	4.779	4.215	3.77	3.4	3.113	2.895
Adamantane	11.43	9.585	8.124	6.967	6.049	5.315	4.929
Undecane	13.833	11.199	9.207	7.662	6.54	5.59	4.929
Naphthalene	21.913	17.759	14.562	12.06	10.17	8.624	7.485
Dodecane	27.976	21.921	17.44	14.015	11.562	9.535	8.117
(C) Mixture 3							
	343.7	348.8	353.85	359	364	369.05	
Methane ^a	1.319	1.317	1.329	1.341	1.351	1.35	
<i>n</i> -Heptane	1.805	1.735	1.688	1.653	1.623	1.59	
Methylcyclohexane	1.961	1.872	1.809	1.76	1.718	1.675	
1-Octene	2.398	2.234	2.101	1.998	1.915	1.844	
Nonane	3.863	3.421	3.078	2.805	2.587	2.406	
Cubane	5.481	4.78	4.219	3.772	3.413	3.12	
exo-THDCPD ^b	10.491	8.994	7.494	6.423	5.594	5.012	
endo-THDCPD ^b	12.463	10.508	8.751	7.447	6.43	5.678	
	343.6	348.8	353.9	359.0	364	369.1	374.2
(D) Mixture 4							
	343.6	348.8	353.85	359.0	364	369.1	374.2
Methane ^a	1.295	1.301	1.295	1.304	1.315	1.326	1.325
<i>n</i> -Heptane	1.854	1.78	1.720	1.674	1.636	1.608	1.575
Methylcyclohexane	2.018	1.924	1.845	1.784	1.734	1.693	1.652
1-Octene	2.464	2.286	2.142	2.028	1.934	1.855	1.79
Nonane	3.952	3.495	3.128	2.84	2.609	2.421	2.263
Cubane	5.59	4.862	4.340	3.815	3.507	3.159	2.872
exo-THDCPD ^b	10.689	8.95	7.582	6.504	5.632	4.882	4.378
endo-THDCPD ^b	12.715	10.654	8.868	7.54	6.478	5.593	4.946

^a On-retained reference.

^b Tetrahydrodicyclopentadiene.

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to be at least 99 mole percent pure. Cubane (+99 mol%) 69 was kindly supplied by Professor Phillip Eaton. Correla-70 tion gas chromatography experiments were performed on 71 an HP 5890A Series II Gas Chromatograph equipped with 72 a split/splitless capillary injection port and a flame ioniza-73 tion detector run at a split ratio of 100/1. Retention times 74 75 were recorded to three significant figures following the decimal point on a HP 3989A Integrator. The instrument was 76 run isothermally using either a 15 or 30 m SPB-5 capillary 77 column. Helium was used as the carrier gas. At the temper-78 atures of the experiments, the retention time of the solvent, 79 CH₂Cl₂, decreased with increasing temperature suggesting 80 that it was retained by the column. Methane was bubbled 81 prior to each run and its retention time was found to in-82 crease with temperature. A consequence of the increase in 83 viscosity of the carrier gas with temperature, this is the cri-84 85 terion used to confirm that a substance is not retained on the column and can be used to determine the dead volume 86 of the column. The gas chromatographic retention times of 87 cubane and the standards are summarized in Table 1. Ad-88 justed retention times, t_a , were calculated by subtracting the 89 90 measured retention time of methane from the retention time of each analyte as a function of temperature usually over a 91 30 K range. Column temperatures were controlled by the gas 92 chromatograph and were monitored independently by using 93 a Fluke 51 K/J thermometer. Temperature was maintained 94 constant by the gas chromatograph to ± 0.1 K. 95

96 3. Results

⁹⁷ Cubane was analyzed using a series of different standards ⁹⁸ in four separate mixtures. The literature values of the stan-⁹⁹ dards are reported in Table 2. Standards were chosen on the ¹⁰⁰ basis of their relative retention times, our assessment of the ¹⁰¹ reliability of their vaporization enthalpies and their struc-¹⁰² tural similarities to cubane. A plot of $\ln(1/t_a)$ against 1/T(K)¹⁰³ resulted in straight lines characterized by the parameters

Table 2 Literature values used as reference for cubane; molar enthalpies in $kJ \text{ mol}^{-1}$ Avaptfm (298.15 K)

$\Delta_{\rm vap}H_{\rm m}(298.15{\rm K})$	Reference
35.1 ± 0.2	[9]
35.4	[10]
36.57 ± 0.18	[11]
40.3 ± 0.2	[12]
41.56 ± 0.2	[11]
46.55 ± 0.46	[11]
48.2	[13]
49.1 ± 2.3	[14]
50.2 ± 2.3	[14]
51.42 ± 0.26	[11]
55.65 ± 2.8	[15]
56.58 ± 0.56	[11]
61.52 ± 0.61	[11]
	$\begin{array}{l} \Delta_{\rm vap} H_{\rm m}(298.15{\rm K})\\ 35.1\pm0.2\\ 35.4\\ 36.57\pm0.18\\ 40.3\pm0.2\\ 41.56\pm0.2\\ 41.56\pm0.2\\ 46.55\pm0.46\\ 48.2\\ 49.1\pm2.3\\ 50.2\pm2.3\\ 51.42\pm0.26\\ 55.65\pm2.8\\ 56.58\pm0.56\\ 61.52\pm0.61\\ \end{array}$



Fig. 1. The correlation between experimental and calculated vaporization enthalpies of the standards used in four separate correlations. The equation of the line calculated by a linear regression analysis is given by: $\Delta_{vap}H_m$ (kJ mol⁻¹) (lit) = (1.004 ± 0.034 $\Delta_{vap}H_m$ (calcd) – (0.15 ± 0.98).

listed in the second and third columns of Table 3. Equations 104 for the correlation of enthalpies of transfer from solution 105 to the vapor, $\Delta_{sln}{}^{v}H_{m}(T_{m})$, against experimental vaporiza-106 tion enthalpies are given at the bottom of Table 3 for each 107 correlation. A graphical summary of how well experimental 108 vaporization enthalpies were reproduced is given in Fig. 1. 109 The equation describing the correlation between experimen-110 tal and calculated values of $\Delta_{vap}H_m$ (298.15 K) is provided 111 in the caption of Fig. 1. The mean vaporization enthalpy 112 of cubane at T = 298.15 K resulting from the four separate 113 correlations is $(44.7 \pm 1.6) \text{ kJ mol}^{-1}$ (Table 4). 114

Solid phase transitions of cubane have been previously 115 measured by adiabatic calorimetry and DSC [17]. Two 116 phase transitions have been observed in the solid state of 117 cubane, a solid-solid transition at $T_{\rm tr} = (394.02 \pm 0.04) \,\mathrm{K}$ 118 $(\Delta_{\rm tr} H_{\rm m}(5.94 \pm 0.02) \, \rm kJ \, mol^{-1})$ measured by adiabatic 119 calorimetry and $T_{\text{fus}}(\text{onset}) = (404.9 \pm 0.5) \text{ K} (\Delta_{\text{fus}} H_{\text{m}}(8.7 \pm 0.5) \text{ K})$ 120 \pm 0.3) kJ mol⁻¹) measured by DSC [17]. Since both of 121 these transitions occur above T = 298.15 K, both must be 122 taken into account in calculating the sublimation enthalpy of 123 cubane at T = 298.15 using the following thermodynamic 124 equality: 126

$$\Delta_{\rm sub} H_{\rm m}(298.15\,{\rm K})$$
 127

$$= \Delta_{\rm vap} H_{\rm m}(298.15\,{\rm K}) + \Delta_{\rm cr}(2)^{-1} H_{\rm m}(298.15\,{\rm K}) \tag{1}$$
 128

where $\Delta_{\text{vap}}H_{\text{m}}$ (298.15 K) represents the vaporization enthalpy at T = 298.15 K and $\Delta_{\text{cr}(2)}{}^{1}H_{\text{m}}(298.15$ K) represents the sum of the measured phase transition and fusion enthalpy the sum of the measured phase transition and fusion enthalpy the sum of the measured phase transition and fusion enthalpy the sum of the measured phase transition and fusion enthalpy the sum of the measured phase transition and fusion enthalpy the sum of the measured phase transition and fusion enthalpy the sum of the measured phase transition and fusion enthalpy the sum of the measured phase transition and fusion enthalpy the sum of the measured phase transition and fusion enthalpy the sum of the measured phase transition and fusion enthalpy the sum of the measured phase transition and fusion enthalpy the sum of the measured phase transition and fusion enthalpy the sum of the measured phase transition and fusion enthalpy the sum of the sum of the measured phase transition and fusion enthalpy the sum of the sum of the measured phase transition and fusion enthalpy the sum of the sum of the measured phase transition and fusion enthalpy the sum of the sum of the sum of the measured phase transition and fusion enthalpy the sum of t

The vaporization enthalpy of cubane at T = 298.15 K is 133 obtained directly by correlation gas chromatography. Ad-134

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Table 3

	$\Delta_{\rm sln} {}^{v}H_{\rm m}(361~{\rm K})/R$	Intercept	$\Delta_{\rm vap}H_{\rm m}$ (298.15 K) (lit)	$\Delta_{\rm vap}H_{\rm m}$ (298.15 K) (calc)
(A) Mixture 1 ^a				
Cubane	4086.2	10.477		45.6
Nonane	4130.5	11.102	46.55	46.0
exo-THDCPD ^b	4533.4	11.028	49.1	49.7
endo-THDCPD ^b	4617.7	11.054	50.2	50.5
Decane	4713.1	11.996	51.42	51.4
Undecane	5232.7	12.733	56.58	56.2
(B) Mixture 2 ^c				
	$\Delta_{\rm sln}^{v} H_{\rm m}(359{\rm K})/R$	Intercept	$\Delta_{\rm vap}H_{\rm m}$ (298.15 K) (lit)	$\Delta_{\rm vap}H_{\rm m}$ (298.15 K) (calc)
Norbornene	3772.8	11.351	35.1	36.4
Octane	4214.6	12.077	41.56	420.
Cubane	4332.1	11.176		43.6
Adamantane	4539.4	10.909	48.2	46.2
Naphthalene	5163.4	12.008	55.65	54.3
Undecane	5358.1	13.074	56.58	56.8
Dodecane	5835.5	13.71	61.52	62.9
(C) Mixture 3 ^d				
	$\Delta_{ m sln}^{v} H_{ m m}(356{ m K})/R$	Intercept	$\Delta_{\rm vap}H_{\rm m}$ (298.15 K) (lit)	$\Delta_{\text{vap}}H_{\text{m}}$ (298.15 K) (calc)
Methylcyclohexane	3417.9	10.389	35.40	35.1
<i>n</i> -Heptane	3544.4	11.036	36.57	36.5
1-Octene	3951.2	11.421	40.3	41.0
Cubane	4292.3	11.065		44.7
Nonane	4411.4	11.904	46.55	46.1
exo-THDCPD ^b	4693.2	11.436	49.10	49.1
endo-THDCPD ^b	4793.4	11.535	50.20	50.2
(D) Mixture 4 ^e				
	$\Delta_{\rm sln}^{v} H_{\rm m}(356{\rm K})/R$	Intercept	$\Delta_{\rm vap}H_{\rm m}$ (298.15 K) (lit)	$\Delta_{\text{vap}}H_{\text{m}}$ (298.15 K) (calc)
Methylcyclohexane	3346.6	10.063	35.40	35.7
<i>n</i> -Heptane	3385.7	10.434	36.57	36.1
1-Octene	3898.0	11.19	40.3	41.0
Cubane	4256.3	10.927		44.4
Nonane	4386.0	11.789	46.55	45.7
exo-THDCPD ^b	4763.0	11.622	49.10	49.3
endo-THDCPD ^b	4870.3	11.736	50.20	50.3

^a $\Delta H_{\rm vap}$ (298.15 K) kJ mol⁻¹ = (1.117 ± 0.085) $\Delta_{\rm sln}^{v} H_{\rm m}$ (361 K) + (7.63 ± 0.56); r^2 = 0.9829.

^b Tetrahydrodicyclopentadiene.

^c $\Delta_{l}{}^{g}H_{m}$ (298.15 K) kJ mol⁻¹ = (1.55 ± 0.11) $\Delta_{sln}{}^{v}H_{m}$ (359 K) - (12.24 ± 1.55); r^{2} = 0.9809. ^d $\Delta_{l}{}^{g}H_{m}$ (298.15 K)/kJ mol⁻¹ = (1.321 ± 0.041) $\Delta_{sln}{}^{v}H_{m}$ (359 K) - (2.39 ± 0.45); r^{2} = 0.9961. ^e $\Delta_{l}{}^{g}H_{m}$ (298.15 K)/kJ mol⁻¹ = (1.157 ± 0.052) $\Delta_{sln}{}^{v}H_{m}$ (359 K) + (3.52 ± 0.64); r^{2} = 0.9921.

Table 4 Summary of vaporization enthalpies; by correlation and from the literature

	$\Delta_{\rm vap}H_{\rm m}$ (298.15 K) (lit)	Mix 1	Mix 2	Mix 3	Mix 4	$\Delta_{\rm vap}H_{\rm m}$ (298.15 K) mean
Norbornene	35.1		36.4			36.4
Methylcyclohexane	35.4			35.1	35.7	35.4
Heptane	36.57			36.5	36.1	36.3
1-Octene	40.3			41.0	41.0	41.0
Octane	41.56		42.0			42.0
Cubane		45.6	43.6	44.7	44.4	44.6 ± 0.8
Nonane	46.55	46.0		46.1	45.7	45.9
Adamantane	48.2		46.2			46.2
exo-THDCPD ^a	49.1	49.7		49.1	49.3	49.2
endo-THDCPD ^a	50.2	50.5		50.2	50.3	50.3
Decane	51.42	51.4				51.4
Naphthalene	55.65		54.3			54.3
Undecane	56.58	56.2	56.8			56.5
Dodecane	61.52		62.9			62.9

^a Tetrahydrodicyclopentadiene.

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Table 5 Phase change enthalpies of cubane; enthalpies in kJ mol⁻¹

$\overline{\Delta_{\rm cr(2)}}^{\rm cr(1)} H_{\rm m}(394{\rm K}) ({\rm lit})^{\rm a}$	5.94 ± 0.02
$\Delta_{\rm cr(1)}^{\rm cr(1)} H_{\rm m}(T_{\rm fus})$ (lit) ^a	8.7 ± 0.3
T _{fus} (K)	404.9
$\Delta_{\rm cr}^{1} C p_{\rm m} \Delta T {\rm kJ} {\rm mol}^{-1}{\rm b}$	-4.02 ± 1.2
$\Delta_{\rm cr(2)}{}^1 H_{\rm m}(298{\rm K})$	10.6 ± 1.2
$\Delta_{\rm vap}H_{\rm m}~(298{\rm K})^{\rm c}$	44.6 ± 1.6
$\Delta_{\rm sub} H_{\rm m}$ (298 K)	55.2 ± 2.0

^a [17].

^b The experimental heat capacity of the crystal at T = 298.15 K was obtained graphically [17]: Cp(cr) = 125 J mol⁻¹ K⁻¹; the heat capacity of the liquid phase of cubane was estimated [16]: CpQ(l) = 179.2 J mol⁻¹ K⁻¹; the total phase change enthalpy, $\Delta_{cr(2)}{}^{1}H_m(T_{fus})$ was adjusted to T= 298.15 K using Eq. (4) [18]; the uncertainty ($\pm 2\sigma$) assumed to be 0.3 of the magnitude of the temperature adjustment, see [18] for further details. ^c Uncertainty in vaporization enthalpy represents $\pm 2\sigma$ of the mean.

justment of the total phase change enthalpy from $T = T_{fus}$ to 135 T = 298.15 K is necessary because of the difference in heat 136 capacity of the crystalline and liquid phases. A protocol for 137 doing this is described below (Eqs. (4) and (5)); the method 138 has recently been tested [18]. Inclusion of the temperature 139 adjustment, which is small, results in a sublimation enthalpy 140 for cubane at T = 298.15 K of (55.2 ± 2.0) kJ mol⁻¹. These 141 results are summarized in Table 5. This value is considerably 142 smaller than the value of (80.3 \pm 1.7) kJ mol⁻¹ measured 143 by Knudsen effusion reported previously [4]. 144

145 **4. Discussion**

The sublimation enthalpy of cubane has previously been 146 147 calculated by the atom-atom potential method. The value calculated, 62.8 kJ mol^{-1} [19], is in reasonably good agree-148 ment with the value of (55.2 ± 2.0) kJ mol⁻¹ obtained in this 149 study. V.V. Diky et al. in their article questioning the subli-150 mation enthalpy of cubane demonstrate that the sublimation 151 enthalpies of a variety of saturated cyclic and polycyclic hy-152 drocarbons correlate on a qualitative basis, with their mo-153 lar mass. A graph similar to theirs is reproduced in Fig. 2 154 using the data in Table 6 obtained from recent compendia 155 [20–22] and includes the sublimation enthalpy of cubane 156 determined previously (\bullet) and by this work (\blacksquare) . Literature 157 vaporization and sublimation enthalpies, columns 2 and 3 of 158 Table 6, were adjusted to T = 298.15 K when necessary, us-159 ing Eqs. (2) and (3), respectively. The sublimation enthalpy 160 of the rigid solid, $[\Delta_{sub}H_m (298.15 \text{ K})]$ column 10, Table 6, 161 was calculated by combining the sublimation enthalpy with 162 all solid-solid phase transitions occurring between T = 0 K, 163 and the temperature(s) at which the sublimation enthalpy 164 was measured, columns 3 and 4. For compounds that are 165 liquids at T = 298.15 K, the vaporization enthalpy was ad-166 justed to T = 298.15 K using Eq. (2) when necessary and the 167 fusion enthalpy was adjusted to T = 298.15 K using Eq. (4) 168 [16,18]. Their sum (columns 2 and 9), combined with any 169 solid-solid phase transitions observed (column 4), are also 170



Fig. 2. Relation of enthalpies of sublimation of the rigid solid $([\Delta_{sub}H_m])$ at T = 298.15 K to their molar masses, M [7]. Cage hydrocarbons (\Box); bicyclic compounds (Δ); cyclic compounds (\bigcirc). Cubane: (\blacksquare) this work. Cubane: (\blacksquare) lit. [4]. The solid line was obtained by a linear regression analysis.

listed in column 10. Sublimation enthalpies of the rigid solid 171 were calculated in a slightly different manner by Diky et al. 172 [7]; however, the results compare favorably. All heat capacities, in column 6 were estimated unless noted otherwise. 175

$$\Delta_{\rm vap} H_{\rm m}(298.15 \,{\rm K}) \, ({\rm kJ \, mol}^{-1})$$
 176

$$= \Delta_{\rm vap} H_{\rm m}(T_{\rm m}) + \frac{(10.58 \pm 0.26 \times {\rm Cp}_{\rm l})(T_{\rm m} - 298.15)}{1000}$$
⁽⁷⁾

182

186

$$\Delta_{\rm sub} H_{\rm m}(298.15 \,\text{K}) \,(\text{kJ mol}^{-1})$$

$$= \Delta_{\rm sub} H_{\rm m}(T_{\rm m}) + \frac{(0.75 + 0.15 \text{Cp}_{\rm cr})(T_{\rm m} - 298.15)}{1000} \quad (3)$$
180
181

$$\Delta_{\rm fur} H_{\rm m}(298.15\,{\rm K})/{\rm kJ\,mol^{-1}}$$
183

$$= \Delta_{\rm fus} H_{\rm m}(T_{\rm fus}) + \Delta_{\rm cr}^{-1} C p_{\rm m} \Delta T, \qquad (4) \quad 184$$

where

$$\Delta_{\rm cr}{}^{1}{\rm Cp}_{\rm m}\Delta T \ (\rm kJ \ mol^{-1})$$

$$= [0.15{\rm Cp}({\rm cr}) - 0.26{\rm Cp}(1) - 9.83] \left(\frac{T_{\rm fus} - 298.15}{1000}\right).$$
187
188

A treatment of the data in the graph by a linear regression analysis, excluding cubane from the analysis, results 191 in the following relationship between sublimation enthalpy 192 and molar mass (M): 193

$$\Delta_{\rm sub} H_{\rm m}(298.15\,{\rm K})\,({\rm kJ\,mol}^{-1})$$
 195

$$= (0.425 \pm 0.023)M + (6.64 \pm 4.6);$$
 $r^2 = 0.9535$ 196

(6) 197

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Table 6						
Enthalpies of sublimation	of saturated	cvclic h	hydrocarbons	and	auxiliarv	data ^a

	$\Delta_{\rm vap}H_{\rm m}$ (298 K)	$\Delta_{\rm sub}H_{\rm m}$ (298 K)	$\Delta_{\rm tr} H_{\rm m}$ ($T_{\rm tr}$)	T _{tr} (K)	Cp(l)/ Cp(cr)	$\Delta_{\rm fus} H_{\rm m}$ ($T_{\rm fus}$)	T _{fus} (K)	$\Delta_{\rm fus}H_{\rm m}$ (298 K)	$[\Delta_{\rm sub}H_{\rm m}]^{\rm b}$ (298 K)	M (g/mol)	$\Delta_{\rm sub}H_{\rm m}$ (298 K) Calc.
Cyclic compounds			,		/	(120)	. ,	. ,	. ,		
Cyclobutane	24.1		5.71	145.7	104/98.4	1.09	182.4	5.4	35.3	56.11	30.5
Cyclopentane	28.5		4.9	122	130/123						
			0.34	138		0.6	179.7	5.84	39.6	70.1	36.4
Cyclohexane	33.04		6.74	186.1	155/148	2.68	279.8	3.62	43.4	84.16	42.4
Cycloheptane	38.53		4.97	134.8							
			0.29	198.2							
			0.45	212.4	181/172	1.89	265.1	3.8	48.0	98.19	48.4
Cyclooctane	43.35		6.31	166.5	207/197						
			0.48	183.8		2.41	287.9	3.1	53.25	112.21	54.3
Cyclododecane		76.1	0.6	199					76.7	168.32	78.2
Cyclotetradecane		96.8							96.8	192.3	88.4
Bicyclic and tricyclic	compounds										
Bicyclooctane	1	47.7	4.6	164.3			52.3	110.2	53.5		
cis-PHI ^c	46.12		8.26	182.3							
			0.39	184.5	223/196	1.4	236.5	3.3	58.1	124.2	59.4
trans-PHI ^c	44.76				223/196	10.9	213.9	14.1	58.9	124.2	59.4
m-Decalin	50.1		2.14	216.1	248/220	9.49	230.2	14.6	66.8	138.3	65.4
trans-Decalin	48.5				248/220	14.4	242.8	18.6	67.1	138.3	65.4
endo- THDCPD ^d		51	10.7	213.8					61.7	136.2	64.5
exo-THDCPD ^d	49.1		3.18	162.1	238/194	1.2	183.2	6.35	58.6	136.2	64.5
Cage compounds											
Cubane									55.2	104.2	50.9
Adamantane		59.3 ^e	3.38	208.7					62.6	136.2	64.5
PCU ^f		55.85	4.86						60.7	146.2	68.8
HCTD ^g		79.29							79.3	184.3	84.9
Diamantane		96.77							96.77	188.3	86.7
Pagodane ^h		116.8			/263				116.8	260.4	117.3

^a Enthalpies in kJ mol⁻¹; estimated heat capacities of the liquid/crystal (Cp(l)/Cp(cr) in J mol⁻¹ K⁻¹ [16]; phase change enthalpies were obtained from references [20–22] unless referenced otherwise.

^b $[\Delta_{sub}H_m](298) = \Delta_{vap}H_m$ (298) $+ \Delta_{fus}H_m$ (298) $+ \Sigma \Delta_{trr}H_m$ (T_{tr}) or $\Delta_{sub}H_m$ (298) $+ \Sigma \Delta_{tr}H_m$ (T_{tr}); sublimation of the rigid crystal.

^c Perhydroindane [7].

^d Tetrahydrodicyclopentadiene [14].

^e Average of six values reported at T = 298.16 K [20].

^f Pentacyclo[5.4.0.0^{2,6}.0^{3,10}.0^{5,9}]undecane [26].

^g Heptacyclo[6.6.0², ⁶.0³, ¹³.0^{4,11}.0^{5,9}.0^{10,14}]tetradecane [28].

^h Adjusted to T = 298.15 K using Eq. (3) using an estimated Cp(cr) of $263 \text{ J} \text{ mol}^{-1} \text{ K}^{-1}$ [29].

The sublimation enthalpy of cubane calculates to (50.9 ± 4.6) kJ mol⁻¹ according to this equation. The results calculated using this equation are included as the last column in Table 6. The experimental sublimation enthalpy of cubane is within the uncertainty of the correlation.

Using the group contribution method reported by Diky 204 et al. [7] for estimating the sublimation enthalpy of the rigid 205 crystalline state, a group value of (5.52 ± 0.41) kJ mol⁻¹ 206 is derived for a cyclic tertiary sp³ carbon from bicyclic and 207 polycyclic model compounds containing five and six mem-208 bered rings. This computes to a sublimation enthalpy of 209 (44.2 ± 3.3) kJ mol⁻¹ estimated for cubane. The experimen-210 tal value determined is about 10 kJ mol^{-1} larger. Using the 211 values reported in Table 6 to generate group values for cyclic 212 tertiary and secondary sp³ hybridized carbon atoms result 213 in near identical group values, 5.59 and 6.87 kJ mol⁻¹, re-214 spectively. 215

The sublimation enthalpy of cubane can also be estimated by combining an estimated vaporization enthalpy with the experimental fusion enthalpy. Using the following equation for predicting the vaporization enthalpy of a hydrocarbon [23]: 220

$$\Delta_{\rm vap} H_{\rm m}(298.15 \,{\rm K}) \,({\rm kJ \, mol}^{-1})$$
 222

$$= 4.69(n_{\rm C} - n_{\rm Q}) + 1.3n_{\rm Q} + 3.0 \tag{7}$$

where $n_{\rm C}$ equals the number of carbon atoms and $n_{\rm Q}$ refers 224 to the number of quaternary carbons, a vaporization enthalpy of 40.5 kJ mol⁻¹ results. Addition of the temperature 226 adjusted fusion enthalpy of 10.6 kJ mol⁻¹ result in an estimated sublimation enthalpy of 51.1 kJ mol⁻¹, a value in 228 good agreement with the experimental determination of 229 (55.2 \pm 2.0) kJ mol⁻¹. 230

As noted by Diky et al. [7], the sublimation enthalpy 231 of cyclotetradecane, also measured by Knudsen effusion by 232

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the same laboratory reporting the cubane results [24], is 233 similarly much larger than results reported by others for the 234 same compound [25–27]. 235

The enthalpy of formation of crystalline cubane [4], is 236 also subject to question. The value of $541.8 \text{ kJ mol}^{-1}$ de-237 rived from combustion measurements ($\Delta_c E^\circ = -4828.3 \text{ kJ}$ 238 mol^{-1}), was corrected for an unspecified amount of ill-239 defined carbon adhering to the walls of the bomb. Unravel-240 ing of this portion of the problem will also require additional 241 experimental measurements. 242

Acknowledgements 243

Financial support from the Research Board of the Uni-244 versity of Missouri is gratefully acknowledged. We would 245 246 also like to thank Professors Phillip E. Eaton for a sample of cubane Donald Rogers (Emeritus) for communicating the re-247 sults of his computations and Vladimir Diky and his cowork-

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ers for communicating their results prior to publication. 249

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