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NATIONAL BUREAU OF STANDARDS REPORT

9803

PRELIMINARY REPORT ON THE THERMODYNAMIC PROPERTIES OF SELECTED LIGHT-ELEMENT AND SOME RELATED COMPOUNDS

(The previous semiannual reports in this series have the NBS Report Nos. 6297, 6484, 6645, 6928, 7093, 7192, 7437, 7587, 7796, 8033, 8186, 8504, 8628, 8919, 9028, 9389, 9500, and 9601.)



U.S. DEPARTMENT OF COMMERCE

NATIONAL BUREAU OF STANDARDS

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> Technical Summary Report on the Thermodynamic Properties of Light-Element Compounds

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U.S. DEPARTMENT OF COMMERCE NATIONAL BUREAU OF STANDARDS

ABSTRACT

Thermodynamic and related properties of substances important in current high-temperature research and development activities are being investigated under contract with the U. S. Air Force Office of Scientific Research (USAF Order No. ISSA 67-6) and the Advanced Research Projects Agency (ARPA Order No. 20). This research program is a direct contribution to the Interagency Chemical Rocket Propulsion Group (Working Group on Thermochemistry) and, simultaneously, to other organizations oriented toward acquiring the basic information needed to solve not only the technical problems in propulsion but also those associated with ballistics, reentry, and high-strength high-temperature materials. For given substances this needed basic information comprises an ensemble of closely related properties being determined by an extensive array of techniques. Some of these techniques, by relating thermodynamic properties to molecular or crystal structure, make it possible to tabulate these properties over far wider ranges of temperature and pressure than those actually employed in the basic investigations.

This report presents the results and interpretation of five recent NBS experimental studies of thermodynamic properties, two reviews which refine data treatment and evaluation, and one new critical assessment of published heats of formation. The heat capacity of beryllium nitride, Be₃N₂, was precisely measured from 0 to 315 K; after judicious correction for the irreducible impurities, the data led to the common thermodynamic properties from 0 to 315 K, complementing earlier NBS measurements from 273 to 1173 K. Precise high-temperature calorimetry gave the heat capacity of tungsten from 1100 to 2600 K (showing no detectable difference between two samples, one triple-zone-refined and 99.999% pure); finally corrected results are given also for solid and liquid beryllium aluminate, BeA l_2O_4 , from 1200 to 2400 K. A massspectrometric investigation of Al_2O_3 containing 5.8% BeO identified the high-temperature gas species and gave values for their standard heats of formation. As part of the current NBS microwave and infrared investigation of the alkali-hydroxide molecules, the high-temperature microwave spectra of four isotopic varieties of RbOH were observed and analyzed to give their molecular constants; like CsOH (studied earlier), RbOH is linear, with a low-frequency bending vibration of large amplitude. Two reviews deal with reaction calorimetry involving fluorine compounds: One documents the inadequacy of estimation rules for unmeasured inorganic fluorides, and recommends a systematic cataloging of results to achieve better consistency; while the other describes a new computer subroutine for the automatic reduction of bomb-calorimetry data on organic fluorine compounds. As the result of an up-to-date critical evaluation of the published data giving ΔHf_{298}^{0} , ΔGf_{298}^{0} , and S_{298}^{0} of magnesium compounds, values are documented and tabulated for 60 species of interest in "light-element" applications.

Thomas в.

Project Leader

Charles W. Beckett Assistant Division Chief for Thermodynamics Heat Division, Institute for Basic Standards

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Chapter 1

THE THERMOCHEMISTRY OF INORGANIC FLUORINE COMPOUNDS THE NEED FOR A CATALOG OF THERMOCHEMICAL PROCESSES¹

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1. ESTIMATES OF HEATS OF FORMATION

Some years ago we reviewed (see for example [1]) the effectiveness of estimates of the heats of formation of inorganic fluorine compounds by a procedure which we have not seen used elsewhere. We compared estimates of heats of formation for which no measured value existed at the time of the estimate, with later values measured experimentally. The estimates were, thus, <u>a priori</u> in a sense. By this tactic we hoped to avoid the possibility of (no doubt unpremeditated) influence of the measured value upon the quality of the estimate. We considered, for instance, that if a measured value were available the author of an estimate might be influenced favorably in his decision to publish by good agreement of his estimate; or, contrariwise, unfavorably and decide not to publish if agreement were poor; thus we would anticipate that estimates made after the availability of experimental data might tend to be better than estimates made prior to such availability.

Some new information has been added and the comparisons are shown in Table 1. The average error (without regard to sign) is 44 kcal mol⁻¹, and the average bias is 26 kcal mol⁻¹ positive, as compared with our earlier finding of 30 and 17 kcal mol⁻¹, respectively. Thus, the ability to estimate heats of formation was not particularly good, and recent data have continued to show large deviations. The tendency is for the compounds of fluorine to be more stable than estimated.

The difficulty may be due in part to the extreme position of fluorine in the periodic system, which causes all attempts to estimate on the basis of the properties of neighboring elements to be extrapolations rather than interpolations.

¹ Prepared for presentation at the Thermodynamic Symposium of the IUPAC Commission on Thermodynamics and Thermochemistry, Heidelberg, West Germany, September 1967.

Table 1

Accuracy of Estimates of Heats of Formation of Some Fluorine Compounds

$\begin{array}{c c c c c c c c c c c c c c c c c c c $	Compound	ΔH	kcal mol ⁻¹	Error	
$\begin{array}{cccccccccccccccccccccccccccccccccccc$		Estimate	Later Masurement		
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	$AcF_{3}(c)$	-420	-477	+57	
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$		-395		+82	
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	AlF ₃ (c)	-311	-361.0	+50	
$\begin{array}{cccccccccccccccccccccccccccccccccccc$		-323		+38	
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	AuF ₃ (c)	-100	- 83.3	-17	
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	BeF ₂ (c)	-227	-245.3	+18	
$\begin{array}{cccccccccccccccccccccccccccccccccccc$			•	+25	
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$				+26	
$\begin{array}{cccccccccccccccccccccccccccccccccccc$		-405		-16	
$\begin{array}{cccccccccccccccccccccccccccccccccccc$		17	8.9	+ 8	
-420 -39 RuF_5(c) -300 -213.4 -87 SF_4(g) -156 -171.7 $+16$ TaF_5(c) -300 -455 $+155$ -360 -455 $+95$ ThF_4(c) -477 -482 $+5$ TiF_4(c) -370 -394.2 $+24$ T&F(c) -65 -74.0 $+9$	NbF ₅ (c)	-342	-432	+90	
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	PF ₅ (g)	-315	-381.4	+66	
$\begin{array}{cccccccccccccccccccccccccccccccccccc$		-420		-39	
$TaF_5(c)$ -300-455+155-360+95 $ThF_4(c)$ -477-482+ 5 $TiF_4(c)$ -370-394.2+24 $T\ell F(c)$ - 65- 74.0+ 9	RuF ₅ (c)	-300	-213.4	-87	
-360 $+95$ $ThF_{4}(c)$ -477 -482 $+5$ $TiF_{4}(c)$ -370 -394.2 $+24$ $T\ell F(c)$ -65 -74.0 $+9$	SF ₄ (g)	-156	-171.7	+16	
$ThF_{l_1}(c)$ -477-482+ 5 $TiF_{l_1}(c)$ -370-394.2+24 $T\ell F(c)$ - 65- 74.0+ 9	TaF ₅ (c)	-300	-455	+155	
TiF-370-394.2+24TlF(c)-65-74.0+9		-360		+95	
TlF(c) - 65 - 74.0 + 9		-477	-482	+ 5	
	TiF ₁ (c)	-370	-394.2	+24	
	TLF(c)	- 65	- 74.0	+ 9	
$TLF_{3}(c)$ -175 -136.9 -38	TLF3(c)	-175	-136.9	-38	
$ZrF_{4}(c)$ -445 -456.8 +12		-445	-456.8	+12	
Average error 44 kcal mol-1			Average e	error 44	kcal mol-1
			Average b	ias 26	kcal mol-1

Estimates are selected from a review by Armstrong and Krieger [2].

We have not made a general study of the problem of predictions of binding energies or of the relative effectiveness of the different methods that have been used. This is a worthwhile exercise, and possibly if we had done so we would have somewhat less of a blunderbuss approach to the results. However, our concerns up to now at least have been primarily in other directions. The information which has been presented to you was enough to persuade us to adopt a skeptical attitude toward estimates. It is apparent that competent and well established reviewers have on the whole shown a lack of ability to predict <u>a priori</u> with reasonable accuracy the heats of formation of inorganic fluorine compounds.

2. SOME SOURCES OF EXPERIMENTAL DIFFICULTY IN MEASUREMENTS

The unusual bias of predictions, together with a curious regularity observed in successive experimental determinations of certain heats of formation, led to the formulation of a rather general rule, that members of cur laboratory have rather facetiously termed Armstrong's rule: as time goes by heats of formation tend to become more negative. In the denouement we hold no brief for this rule, but point it out as an interesting interlude in our examination of the findings of experimental thermochemistry.

An example that led to this attitude is the heat of formation of aluminum oxide (Table 2) which became significantly more negative over a period of 30 years. The heat of formation of $A&F_3$ has followed roughly this rule, we see semblences of it also in the heats of formation of BeO(c), $BeF_2(c)$, $CF_{l_1}(g)$, and HF(aq).

Table 2

Successive Determination	s AHr[Al ₂ 03(c,a)]
	kcal mol-1
Moose and Parr (1924) [3]	-376.9
Roth and Muller (1929) [1]	-380.1
Snyder and Seltz (1945) [5]	-399.09
Oketani and Maebashi (1950)	[6] -381.03
Holley and Huber (1951) [7]	-400.29
Schneider and Gattow (1954)	[8] -400.6 (-402)
Mah (1957) [9]	-400.4

The number in parenthesis included a questionable correction made by the authors for partial formation of a less stable phase.

When we examine these tendencies in greater detail, however, we find, perhaps not surprisingly, that the picture was oversimplified. Still, certain regularities of behavior do persist, and lead us to believe there is some rational explanation of these regularities. The explanations which finally appear to emerge are not the same in every case. That is, a variety of influences is apparently at work. An examination of a few of them may assist us in planning future experimental work.

We return to the heat of formation of Al_2O_3 briefly and note that each value quoted resulted from combustion of the metal in oxygen. With a little imagination we realize that probably the cause of progressively more negative heats of formation was better control of the amount of reaction as time passed. The less negative values probably represent smaller amounts of reaction than the experimenter presumed. There may be two reasons for this bias: (1) the sample metal may have contained larger amounts of inert impurities than were revealed by analysis; (2) the metal may have reacted less completely than presumed. The first source of error is rendered plausible by the finding of relatively large amounts of oxygen and nitrogen (probably as a surface coating) in metals even supposedly highly pure on the basis of spectrographic analysis. The second source of error is rendered plausible by the fact that the combustion product may completely cover small amounts of unburned metal and preclude further oxidation. On the basis of an argument like the above, we would anticipate in the combustion of refractory substances giving solid products, a bias in the direction of low heat of reaction which might be substantial in some cases. A notorious example of this is the combustion of boron in oxygen, for which the two processes may both be presumed to be severe. Let us turn to a more complicated and informative example, the heats of formation of CF1, and HF.

A few years ago, apparently valid but inconsistent experiments were reported by Jessup [10] on the one hand and by Good, Scott and Waddington [11] and by Neugebauer and Margrave [12] on the other hand, all of which gave values for the heat of formation of CF_{\downarrow} . See Table 3. In an attempt to reconcile them we observed the opposing sense in which the heat of formation of HF occurred in the relationships between the heats of reaction and the heat of formation of CF_{\downarrow} . These relationships are shown in Table 4, referring to reactions 2, 3b, and 4c. This led us to make the

suggestion that possibly the heat of formation used for HF in the calculation was in error, and it was no problem then to figure a reasonable change in the heat of formation of HF that would reconcile these inconsistent sets of experiments.

If we restrict ourselves to the reactions from which the heat of formation of CF_{\downarrow} is derived by its difference from HF (rather than from its sum with HF) as in reactions 3(a,b,d), 4(a,b,c), 5(a,b,c) and possibly 8, and 9, we find a rather nice consistency between the calculated values for CF_{\downarrow} no matter what value is selected for HF(aq). An illustration of this is given by Cox, Gundry, and Head [13], using an extreme value for the heat of formation of HF(aq).

The heat of formation of CF_{l_1} as inferred from these experiments is inextricably bound to the heat of formation of HF(aq). It is evident that we shall not learn anything further about it from them alone, so long as there is fluctuating information about the heat of formation of HF.

We may turn to the experiments that do not involve HF. Here we find ourselves in reaction sets 1, 6, and possibly 8 and 9, immediately faced with the same problems or at least similar problems to that mentioned earlier in the case of $A\ell_2O_3$. For instance, in reactions (8) and (9), involving the reduction of an organic compound and the formation of a solid residue carbon, the question arises, "to what extent has this reaction gone to completion?" Completion here includes two concepts: (1) has all of the F been extricated from the C residue? and (2) has the C residue been reduced to a recognizable, well defined standard state, or does it retain active centers (as a result of a large surface area, for example)? The experimental evidence, particularly with respect to the latter problem, as outlined in the notes to Table 1 is dishearteningly unsatisfactory. One is left with the feeling that the observed heat processes will tend to be biased toward the low side of the true heat of reaction.

Only by the most careful experimentation can one be sure that the observed heat actually corresponds to the expectation for a complete reaction described exactly by the equation.

Somewhat peripheral to our subject, the analysis of these equations led us to an understanding:

- (1) That the heat of formation of CF_{l_1} is best represented by the experiments on direct combustion of graphite in fluorine; and by a combination of the combustion of "Teflon" in oxygen and in fluorine, the only products being gaseous CO₂ and CF_{l_1} .
- (2) That other reactions not involving HF do not provide any conflicting evidence, but that the positive information they provide is limited by the lack of characterization of the reaction.
- (3) That reactions involving HF(aq) do not provide conflicting evidence, but that the positive information they supply is limited by uncertainty in the heat of formation of HF(aq). (Incidentally, the reaction by Jessup et al. [10] which was an initiator of the line of thought described here, turns out now to be inconsistent in the opposite sense from what it was at the outset, and no means of reconciliation is now apparent.).
 (4) That the reactions involving HF(aq). can be combined with independently arrived at values for heats of formation of CF₄ as well as BF₃ and NF₃) to arrive at new "consensus" values for the heat of formation of HF(aq).
 (5) That the explanation of the observed discrepancies in measurement

That the explanation of the observed discrepancies in measurements of the heat of formation of HF(g) or HF(aq) by various means may involve subtle arguments. The large deviation of the work of Cox and Harrop [19], for instance, may be due not to experimental error but to a residual entropy of $LiHF_2(c)$ at absolute zero, or to use of an inappropriate value for the heat of formation of gaseous HF. In the latter event, the heat of formation of gaseous HF is still a problem.

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3. CATALOGING THE CALORIMETRIC PROCESSES

A feature is present in Table 1 which, undoubtedly, is not new but suggests to us what we believe is a new concept of the appropriate forms for presenting calorimetric (and other thermodynamic) data. We have no question in our mind that the most useful form of thermodynamic information for the general user — the theoretician interested in energies of substances, and the engineer interested in design parameters is a set of tables of values of thermal functions of the substances in the style of NBS Circular 500 [20] (now Technical Note 270-1) [21] or of the JANAF Thermochemical Tables [22] from the U.S.A. or the Thermal Constants of Substances [23] and Thermodynamic Properties of Individual Substances [24] from the Academy of Sciences of the USSR. However, for the thermochemist, the experimental calorimetrist, or the selector of best values, another form seems at least as desirable, and in the latter case almost a necessary adjunct to his work. This form is a catalog of thermochemical processes.

We have seen in Section 2 that a set of equations together with the heat values associated with them is a flexible tool for attempting to determine true values of the heats of formation, and to appraise sources of errors. It is also apparent that critical substances for further study would be discernible in such a set of equations.

A catalog can be conceived of as a tabulation of observed processes chemical reactions, phase changes, and so on — listed as equations, with the value of the thermal effect observed. Limited amounts of selected information concerning the conditions of the experiment are also included. The tabulation of equations is indexed with respect to all substances and phases present, so that all processes involving a given substance can be located.

The limited list of equations involving $CF_{j_{4}}$ given in Table 3 represents a skeleton of a limited segment of such a catalog. To prepare a complete catalog including all the chemical substances for which measurements have been made is a very large task. In the following paragraphs we indicate certain special features of such a compilation and suggest the long term benefits which it might be expected to provide.

There is ample evidence that several groups now have essentially the framework of such a catalog. These are the groups which have been for some years compiling and critically evaluating thermodynamic data. The awkward feature of the present situation is that the information, except for the end result after it has been critically evaluated and released in tables of selected values, is essentially in the form of private files. In this form it is inaccessible to the general investigator; in the event of the formation of a new critical evaluation group, they must acquire their own file of such working information; in the event of the demise of one of these groups, its literature files would probably revert to inutility.

We suggest that much of the intermediate information gleaned from the thermochemical literature is of permanent value, subject to reuse in various ways, and susceptible to augmentation as time goes on without loss of any of the original body of information. In this respect, particularly it is more useful than a collection of selected values of thermodynamic functions, whose values change as new data are added.

A catalog in the form described can be looked upon as a set of mathematical equations or expressions of which a selection is given in Table 4. In general, most of the thermodynamic relationships involving heats of formation can be described as linear equations. The set of possible chemical reactions involving all the chemical substances may be looked upon as a set of simultaneous equations. Experimental enthalpy changes can be expressed in terms of the numerical coefficients and heats of formation of the substances present.

Because of the complexity and number of these equations, and the difficulty of finding simultaneous solutions which offer the best values of the heats of formation, the practice has been adopted of evaluating heats of formation sequentially, starting with certain "key" substances, to which values of heat of formation are assigned on the basis of carefully selected equations.

In principle, if the experimental enthalpy equations are known for any given set of substances, a set of best values can be selected which minimizes the ensemble of errors without predetermining any one key substance. This minimization is ordinarily not done because of its difficulty.

It is to be presumed that the use of high speed computers will ultimately allow selections of best values in this sense for the whole body of chemical substances for which data exist. A catalog of the type described above is a critical development if such a calculation is to be achieved. The catalog will essentially provide the input data for such a selection of best values.

With a workable program for making such selections, it should be possible to make selected best values annually, or more frequently; it should be possible to study the relative importances of various substances; and also to find those substances whose present values appear to be out of line and require redetermination.

Thus a catalog provides three services:

- (1) A resource for experimental calorimetrists for planning their work.
- (2) Source material for groups preparing best values of thermodynamic properties.
- (3) Input data for computerized selection of best values and other computerized analysis of data.

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	Reaction	AH298 Kcal mol-1	Reference
	$C(c,graphite) + 2F_2(g) = CF_1(g)$	-162±2ª	[1]
	2,0,7 4,07	(-183.5±2) ^b	[3]
		-222.87±0.38	[6]
•	$CH_{l_{1}}(g) + 4F_{2}(g) = CF_{l_{1}}(g) + 4HF(g)$	-459.3±9	[7]
(a)	$C_2F_4(g) = CF_4(g) + C(c, graphite)$	-61.43±1.4	· [8]
• •	2 4 - 4 -	-65.4±0.42	[9]
(b)	$C_2 F_{l_1}(g) + 2H_2(g) + 4[118.2H_20](l) = -$		
	4[HF+118.2H_0](l) + 2C(c,graphite)	-150.8±1.1	[9]
(c)	$C_2F_{l_1}(g)+2H_2(g) = 4HF(l)+2C(c,graphite)$	-132.72±0.7	[8]
	$4[HF+118.2H_0](l) + C(c,graphite) =$		
	$CF_{l_1}(g) + 2H_2(g) + 4[118.2H_20](l)$	+85.4±1.5	
(e)	$4HF(l)+C(c,graphite) = CF_{l}(g) + 2H_{2}(g)$	+67.32	
	$C_2F_{l_1}(solid polymer)+O_2(g) = CO_2(g)+CF_{l_1}(g)$		[10,11]
	C_2F_4 (solid polymer)+ $O_2(g) + 42H_2O(l) =$		
•	$2CO_{2}(g)+4[HF+1OH_{2}O](l)$	-160.3±0.9	[10,11]
(c)	$CO_2(g) + 4[HF + 10H_20](l) = CF_1(g) + 42H_20(l)$	+1,1.5±1.0	[10,11]
	$C_{12}F_{22}(l)+6.50_{2}(g)+401.8H_{2}0(l) = 0.60F_{1}(g)$		
	$11.4 CO_2(g) + 19.6[HF+20H_20](l)$	-4067.9/4.184	• [12]
(b)	$C_{12}F_{22}(l)+6.50_{g}(g)+131.2H_{2}O(l) = 3.9CF_{l}(g)$		
	$8.1CO_{2}(g) + 6.4[HF+20H_{2}O](l)$	-3496.5/4.184	
(c)	$CO_2(g) + 4[HF + 20H_0](l) = CF_1(g) + 82H_0(l)$	+41.38±0.32	[12]
	$C_2F_{l_1}(\text{solid polymer})+F_2(g) = 2CF_{l_1}(g)$	-247.85(-247.13)	0.30 [5,6]
	2 4 2 4	-246.84±0.07	[17]
(b)	$C_2F_{l_1}(solid polymer)+O_2(g) = CO_2(g)+CF_{l_1}(g)$) -118.8	[10,11]
	$CO_{2}(g) + F_{2}(g) = O_{2}(g) + CF_{1}(g)$	-129.05(128.63)	
	H ₂ , O ₂ , CO, CF ₁	-220.1±1.4	[13]
	$CF_{j}(g)+4K(c) = 4KF(c) + C(graphite)$	-307±!+	[2,4]
		-320±2	[15]
	$CF_{j}(g)+4Na(c) = 4NaF(c)+C(graphite)$	-325.5±2.2	[16]

* See separate set of references for Table 3.

Notes to Table 3

1. a. The original authors made an adjustment of 2.4 kcal mol⁻¹ for the estimated heat of formation of the Norite (activated wood charcoal) they burned. They obtained -165.0 ±1.5 kcal mol⁻¹ for the combustion of Norite in fluorine. When adjusted by +2.4 kcal mol⁻¹ this gives -162.6 kcal mol⁻¹ which they apparently rounded to -162 kcal mol⁻¹, increasing the uncertainty to 2 kcal mol⁻¹.

b. This is an adjustment by Ruff and Bretschneider [3] of the work of von Wartenberg and Schuette [1], taking into account the formation of several percent of higher fluorocarbons in the combustion of Norite. The recalculation changes the original data far too little. A recalculation using a more current value for the fluorination of a C-C bond (142.7 kcal mol⁻¹, instead of 107 kcal mol⁻¹ as was used by Ruff and Bretschneider) leads to $\Delta H_r = -190.5$ kcal mol⁻¹. Hence it appears that the formation of other products would have to be greater than was suggested by Ruff and Bretschneider. The uncertainty in amount of products seems to preclude further consideration of the work of von Wartenberg and Schuette.

2.

As a part of its uncertainty the work of Jessup, McCoskey, and Nelson [7] contains an uncertain correction of 1.5 to 2.0 kcal mol⁻¹ (of CF_{l_1}) for nonideality of HF. This now appears to have been an excessive correction. In the extreme case, if this correction is reduced to zero, the heat of reaction becomes -461.2 kcal mol⁻¹. The large uncertainty assigned by the authors is apparently a 95 percent confidence limit.

3. a. The carbon formed in the reaction by Duus [8] was weakly crystalline. Duus made no adjustment to the heat for the physical state of the graphite. An adjustment of the magnitude made by Neugebauer and Margrave [9] would bring Duus' value to -63.3 kcal mol⁻¹. Neugebauer and Margrave [9] measured the heat of reaction, with the formation amorphous carbon, to be -63.5 kcal mol⁻¹, and measured the heat of formation of the amorphous carbon to be 1.9 kcal mol⁻¹. We have combined their measurements to give the heat of decomposition to form graphite, -65.4 kcal mol⁻¹. b,c. Neugebauer and Margrave [9] measured the heat of hydrogenation of $C_2F_{l_4}(g)$, with the formation of HF(aq) and amorphous carbon, to be -lh7.8 kcal mol⁻¹, and measured the heat of formation of the amorphous carbon to be 3.0 kcal mol⁻¹. We have combined their measurements to give the heat of hydrogenation to form HF(aq) and graphite, $\Delta H = -150.8$ kcal mol⁻¹. Their reaction is definitely superior to reaction 3(c) as carried out by Duus, who was forced to make large corrections for the presence of HF(g), and did not attempt any correction for the physical state of the carbon formed. Neither reaction 3(b) nor 3(c) can be used by themselves to calculate the heat of formation of CF_{l_4} , but are included because they can be combined with other work done in the same laboratory to obtain a relationship leading to the desired value.

d. This reaction is obtained by subtracting 3(b) from 3(a). The heat value cited was obtained using the heat of reaction 3(a) determined by Neugebauer and Margrave [9].

e. The heat of this reaction was determined by combining the energy of reaction 3(c) with that for 3(a) determined by Duus [8] and reported in the same paper.

- 4. a,b. The energies of these reactions were obtained by Good, et al. [10],
 [11] by extrapolating respectively to x = 1 and x = 0 from measurements covering the range x = 0.0285 to x = 0.8162 of the reactions
 C₂F₄(solid polymer) + 0₂(g) + 42(1-x)H₂O(1) = (2-x)CO₂(g) +xCF₄ + 4(1-x)[HF + 10H₂O](1). The experiments are carefully described.
 c. The heat of this reaction was obtained by combining the heats of reactions 4(a) and 4(b) determined in the same laboratory.
- 5. a. The heat of this reaction was obtained by Cox, Gundry, and Head [12] by appropriately combining reactions 5(a) and 5(b).
- 6. a. An earlier (in parentheses) and a more recent and more amply substantiated value are listed.

b. The heat of this reaction is obtained by combining the heats of reaction 6(a) and 4(a). This reaction does not in any way involve the heat of formation of HF, but in order for it to be valid, the C_2F_4 (solid polymer) used in the two different experiments must be similar.

Raibuz and Medvedev [14] recalculated the work of Baibuz [13] and reported $\Delta H_f[CF_{\downarrow_1}(g)] = -220.6$. The work of Baibuz is not available to us in sufficient detail to allow us to write the equation for the reaction or to know the dependence of the reported heat of formation of CF_{\downarrow_1} on other auxiliary data.

8.

9.

7.

. Von Wartenberg [2,4] reported $\Delta H = -307$ kcal mol⁻¹, and although he indicated the carbon was graphite, the evidence from other sources is that probably only a slight amount of graphite was present. On the basis of information supplied by Neugebauer and Margrave (1.5 to 1.9 kcal mol⁻¹) [9], von Wartenberg and Schuette (2.4 kcal mol⁻¹) [1], and Kirkbride and Davidson (2.5 kcal mol⁻¹) [15] if we assume the heat of formation of the carbon residue to be +2 ±1 kcal mol⁻¹ and apply a correction, the heat of reaction to that in which graphite is formed would be -307, which is still far less negative than the result of Kirkbride and Davidson.

Kirkbride and Davidson [15] reported $\Delta H_f[CF_{\downarrow}(g)] = -218 \pm 2$ kcal mol⁻¹, together with auxiliary data for XF(c) and amorphous carbon. We have back calculated to obtain $\Delta H_r = -320 \pm 2$ kcal mol⁻¹.

Vorob'ev and Skuratov [16] reported $\Delta H_r = -325.5 \pm 2.2$ kcal mol⁻¹ and that their reaction led to formation of β -graphite. If we presume, on the basis of the work of Kirkbride and Davidson [15], that only a small amount of crystalline material was present and apply a correction of 2 kcal mol⁻¹ for the heat of formation of amorphous carbon, the heat of reaction 9 would be -327.5 kcal mol⁻¹. See 8 above. References to Table 3

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Table 4

16

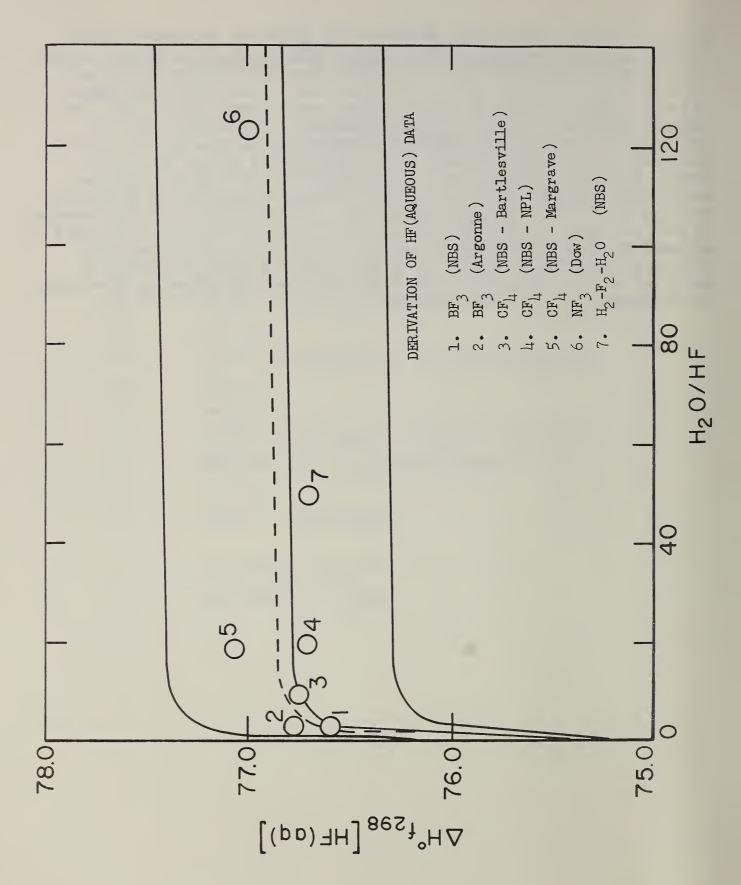
Process for $\Delta H_{f}[CF_{4}(g)]$ from Reactions of Table 3.

Reaction

*-

1	ΔΗ, (1)
2	$\Delta H_{r}(2) + \Delta H_{f}[CH_{l}(g)] - l \Delta H_{f}[HF(g)]$
3(a).	$\Delta H_{r}(3a) + \Delta H_{f}[C_{2}F_{h}(g)]$
3(6)	$\Delta H_{r}(3a) - \Delta H_{r}(3b) + 4\Delta H_{r}[HF in] 18.2 H_{2}0](\ell)$
3(c)	$\Delta H_{r}(3a) - \Delta H_{r}(3c) + 4\Delta H_{f}[HF](\ell)$
3(d)	$\Delta H_{r}(3d) + 4\Delta H_{f}[HF in 118.2 H_{2}0](\ell)$
3(e)	$\Delta H_{f}(3e) + 4\Delta H_{f}[HF(\ell)]$
4(a)	$\Delta H_r(\mu a) + \Delta H_r[C_2F_{\mu}(solid polymer)] - \Delta H_r[CO_2(g)]$
Ц(Ъ)	See 4(c)
4(c)	$\Delta H_{r}(\mu c) + \Delta H_{f}[CO_{2}(g)] + \mu \Delta H_{f}[HF in 10 H_{2}0](\ell)$
5(a)	See 5(c)
5(b)	See 5(c)
5(c)	$\Delta H_{r}(5c) + \Delta H_{f}[CO_{2}(g)] + \mu \Delta H_{f}[HF in 20 H_{2}0](l)$
6(a)	$\frac{1}{2}[\Delta H_r(6a) + \Delta H_r[C_2F_l(solid polymer)]$
6(b)	See 6(c)
6 (c)	$\Delta H_r(6c) + \Delta H_r[CO_2(g)]$
7	
8	$4\Delta H_{f}[KF(c)] - \Delta H_{r}(8)$
9	$4\Delta H_{n}[NaF(c)] - \Delta H_{n}(9)$

Figure 1. Recent information on the heat of formation of aqueous hydrofluoric acid. Upper curve, Cox and Harrop [19]; Lower curve, Wagman, et al. [21]; A cycle for BF₃-HF set up by Johnson, Feder and Hubbard [15] gives point (1) when applied to data of Domalski and Armstrong [14], and point (2) when applied to data of Johnson, Feder, and Hubbard [15] for BF₃. The heat of formation of CF₁ determined by Domalski and Armstrong [17] by combustion of graphite in fluorine is applied to data relating CF₁ to HF of Good, Scott and Waddington [11] to obtain point (3); to the data of Cox, Gundry, and Head [13] to obtain point (4); and to the data of Neugebauer and Margrave [12] to obtain point (5). A study relating NF₃ to HF reported by Sinke [16] is used to obtain point (6). A study of the H₂-F₂-H₂O reaction by King and Armstrong [18] gives point (7). The central solid curve shows an estimate of the probable best values for $\Delta H_f[HF(aq)]$. The dashed curve differs from the solid central curve by an amount showing the effect of using the heat of formation of CF₁ determined from the combustion of "Teflon" in fluorine [17] on the calculations.



Chapter 2

HEAT CAPACITY AND THERMODYNAMIC PROPERTIES OF BERYLLIUM NITRIDE, Be₃N₂, FROM 20 TO 315 K.

George T. Furukawa and Martin L. Reilly

Abstract

The heat capacity of beryllium nitride, Be_3N_2 , was determined from 20 to 315 K and the thermodynamic properties calculated from 0 to 315 K. The entropy at 298.15 K was found to be 34.4 ± 0.3 J deg⁻¹ mol⁻¹ (8.23 ±0.08 cal deg⁻¹ mol⁻¹).

I. Introduction

The results of heat-capacity measurements on beryllium nitride, Be₃N₂, presented between 20 and 315 K in this paper have been obtained in connection with a research program at the National Bureau of Standards to provide thermodynamic data on light-element compounds. In connection with this program, the high-temperature enthalpy measurements have previously been reported by Douglas and Payne [1]. Earlier, Satoh [9] reported relative enthalpy measurements at three temperatures in the range 273 to 773 K on a sample of 98.13 percent purity, the impurities being 0.78, 0.87, and 0.22 percent Be0, Fe_2O_3 , and SiO₂, respectively. Recently Justice [2] reported measurements in the range 25 to 310 K on a sample of 96.7 purity.

II. Apparatus and Method

The low-temperature heat-capacity measurements were made from about 20 to 315 K in an adiabatic calorimeter of a design similar in principle to that described previously [10]. (The measurements were made initially from about 80 K, but the measurements above 315 K were discontinued when small heat effects were observed above 320 K. The results presented in this paper are based on the data obtained subsequent to heating above 320 K.)

The calorimeter vessel, after filling with the Be_3N_2 sample, was sealed by means of a specially designed gold-gasket closure [7] and suspended within the adiabatic shield by means of a Nylon string instead of the filling tube shown in the above reference. The average temperature of the adiabatic shield was sensed relative to that of the outer surface of the double-walled sample vessel by means of fourjunction constantan-Chromel-P thermocouples and was controlled automatically by means of a combination of electronic and electromechanical equipment. Heater wires were wound bifilarly in grooves on the cylindrical surface and on the conical ends of the adiabatic shield to achieve good thermal contact. Details of the design of the calorimeter used, its operation, and the automatic adiabatic control system will be described in a later publication.

The platinum resistance thermometer (laboratory designation: 1057849) used in the measurements was calibrated in accordance with the International Practical Temperature Scale of 1948 [11]. The temperatures in degrees Kelvin (K) were obtained by adding 273.15 deg to the temperatures in degrees Celsius (C). Below 90 K, the thermometer was calibrated on the NBS-1955 provisional scale which is numerically 0.01 deg lower than the former NBS-1939 provisional scale [8]. The NBS-1939 provisional temperature scale is based on a set of platinumresistance thermometers that were compared against a helium gas thermometer. The thermometer resistance measurements were made to the nearest 0.00001 ohm. Initially a manual Mueller bridge was used and later an automatic Mueller bridge. The resistance-time observations obtained with the automatic bridge were analyzed by means of a computer to calculate the heat capacity. The observations with the manually operated bridge were processed by means of a desk calculator.

The electrical power introduced into the calorimeter heater (100 n) was measured by means of a Wenner potentiometer in conjunction with a volt box, standard resistor, and saturated standard cells, which were all calibrated at the National Bureau of Standards. A constant-current power supply stable to 10 ppm or better was used to facilitate the power measurements. The duration of each heating interval was measured by means of a high-precision interval timer operated on the 60-Hz frequency standard provided at the National Bureau of Standards. The 60-Hz frequency standard is based on a 100-kHz quartz oscillator which is stable to 0.5 ppm. An electronic counter containing a 1-MHz interval quartz oscillator was used simultaneously in the measurement of the heating interval. The estimated uncertainty in the determination of the heating interval was not greater than ± 0.01 sec for any heating period, none of which was less than 2 min in these experiments.

III. Sample

The Be₃N₂ sample was specially prepared for the measurements by the Brush Beryllium Company, Cleveland, Ohio, by reacting pure nitrogen and beryllium powder at elevated temperatures and crushing the product to particle sizes between 20 and 50 mesh (0.8 and 0.3 mm sieve openings, respectively). The material received was thoroughly mixed in a controlled-atmosphere box containing dry argon (dew-point: -40 C) and samples were apportioned for chemical analysis, for the hightemperature relative enthalpy measurements [1], and for the lowtemperature heat-capacity measurements described in this paper. The samples for the chemical analysis and the high-temperature enthalpy measurements were sealed in test tubes under dry argon. The sample that was poured into the low-temperature calorimeter vessel weighed 97.415g. The calorimeter vessel containing the sample was pumped at high vacuum and purged with dry helium gas several times and finally, helium gas at a pressure of 70 torr was sealed in the container with the sample. After completion of the measurements the calorimeter vessel was opened in the controlled-atmosphere box and the sample was divided and sealed in polyethylene bags for chemical analysis. The sealed polyethylene bags were placed individually in screw-cap jars which were in turn placed in desiccators, the whole operation being done in the controlled-atmosphere box.

The results of spectrochemical analysis of the sample are summarized in Table 1. Only the impurity elements that were detected are listed. The results of the chemical analyses on the material before and after the calorimetric measurements are given in Table 2. For the nitrogen and total beryllium analyses about 0.3 g samples were first dissolved in dilute hydrochloric acid and the solution diluted to 250 cc in a volumetric flask. A 10-cc aliquot was analyzed for nitrogen using the Kjeldahl method, the ammonia being titrated with 0.01 N hydrochloric acid that had been standardized against ammonium dihydrogen phosphate by means of the Kjeldahl procedure. The total beryllium was determined by precipitating the beryllium hydroxide in a 100-cc aliquot with ammonium hydroxide. The beryllium hydroxide was filtered, ignited, and weighed as Be0. The free beryllium was determined by dissolving 2-g samples in hydrochloric acid. The hydrogen formed was dried, converted to water, and weighed.

The analysis for water represents all forms of hydrogen present in the sample. Samples (20 mg) were heated at 1050 C in a carbonhydrogen-nitrogen analyzer and the amount of water driven off was determined. The oxygen was determined by a non-destructive ll₄-Mev neutron activation technique. Five determinations were made using 6-g samples.

The results of the chemical analysis were found to be inconsistent and unexpected. The decrease in the free beryllium content in the samples before and after the measurements can not be attributed to chemical reactions at temperatures of the measurements. (Small heat effects were observed above about 320 K. See later parts of this section for further details.) The difference is probably due to nonhomogeneity in the material, although relatively large samples (2-g) were used in this determination. The decrease in the total beryllium content because of a chemical reaction during the heat measurements does not seem possible. If the observed heat effects resulted from a chemical reaction within the sample, the expected reaction would be:

$$3Be(OH)_2 + Be_3N_2 \rightarrow 6BeO + 2NH_3$$
 (1)

This should lead to an increase in the apparent beryllium content, since the ammonia would be lost on opening the sample vessel.

For the material removed from the calorimeter vessel after the heat measurements, the mass balance for beryllium found is not consistent in terms of the analyses obtained. The analyses for N, O, and total Be sum to 100.01 percent and the addition of H (0.13 percent) found as water makes the total weight percent 100.14. This is within the expected limits of accuracy of the chemical analyses. However, inconsistencies arise when these analyses are used to obtain mass balance among the chemical species that are expected to be present $(Be_3N_2, Be0, Be(OH)_2,$ and Be metal). On the basis of the results of the oxygen and water analyses (2.38 and 1.14 percent, respectively), the percentage of beryllium as BeO and Be(OH)₂ sums to 0.77 percent. When the Be metal (0.04 percent) and beryllium (47.22 percent) as Be_3N_2 , based on the nitrogen analysis, are added to 0.77 percent, the total beryllium becomes 48.03 percent. This is considerably lower than the total beryllium (48.70 percent) from the chemical analysis. The masses of the substances present also sum to considerably less than 100 percent (96.15% - Be₃N₂, 2.72% - Be(OH)₂, 0.56% - BeO, and 0.04% - Be metal; 99.47% - total mass). These inconsistencies can be resolved if the Be metal analysis were higher (about 0.7 percent), but this is not possible since the estimated error in the chemical analysis for Be metal is much less than this value.

On the other hand, if the analysis shown in Table 2 for water is ignored, the remaining analyses would become consistent on the basis that the substances present are only Be_3N_2 , BeO, and Be metal. All of the oxygen (2.38 percent) would then be taken to be present as BeO, which corresponds to 3.73 percent BeO or 1.34 percent Be. When the percentages of Be as Be_3N_2 and as Be metal are added to 1.34 percent, the total "calculated" Be is 48.61 percent. This is within the limits of chemical analysis found for total Be (48.70 percent). When the 96.15 percent Be_3N_2 , based on the nitrogen analysis, is added to the percentages of BeO and Be metal, the total weight percent is 99.92 percent. The analyses for total beryllium, nitrogen, and oxygen, as mentioned earlier, sum to 100.01 percent. These values are all within the limits of the chemical analysis.

The source of heat effects observed during the measurements can not be explained in terms of the known possible chemical reaction within the sample. The possible chemical reaction depicted by equation (1), in which the water from $Be(OH)_2$ reacts with Be_3N_2 to form BeO and NH_3 , is not consistent with the chemical analyses. The reaction is obviously not possible if the water is securely held at the temperatures of the heat measurements. Duval and Duval [3] interpreted their thermal gravimetric analysis work to indicate that the decomposition of $Be(OH)_2$ starts around 150 C and that complete decomposition of $Be(OH)_2$ to BeO and H_2O occurs only on heating to 951 C. The observed loss in mass at temperatures below 150 C was attributed to adsorbed water in their sample. If any moisture were present in the sample that was used in the work reported here, it was adsorbed between the times the samples was removed from the furnace . and received from the supplier.

After careful consideration of the various possibilities that could be selected on the basis of the chemical analyses and the estimated uncertainties that would arise from them, the choice was made to ignore the analysis obtained for water in the reduction of the heat-capacity measurements. The sample was taken to be 96.23 percent Be₃N₂, 3.73 percent BeO, and 0.04 percent Be metal and corrections were applied to the observed values of heat capacity for BeO [5] and Be metal [6].

IV. Results

Two series of measurements were made, one on the calorimeter vessel with sample (gross) and the other on the empty vessel (tare). The measured energy increments (Q) and the corresponding thermometer resistances (R) before and after heating were analyzed on the highspeed digital computer to obtain dQ/dR of the sample as a function of the thermometer resistance [4]. The values of heat capacity (dQ/dT) were then calculated at even temperatures from the dR/dT and the R-T calibration for the thermometer using the relation:

$$dQ/dT = (dQ/dR)(dR/dT) .$$
(2)

Briefly, the procedure was to obtain first the best polynomial equation of the form

$$dQ/dR = \Sigma a_n R^n$$
(3)

to represent the experimental data on the empty calorimeter vessel (tare) [12]. The polynomial equation was then evaluated at the thermometer resistances observed for the gross measurements in order to obtain the energy increments for sample only from the observations in the calorimeter vessel plus sample experiments. The best polynomial equation giving dQ/dR for the sample only was then obtained.

Figure 1 shows the deviation of the experimental data on the empty container from corresponding values calculated from the polynomial equation fitted by the method of least squares to the energyresistance observations. The range of the polynomial is from n = -4to n = +10 (see eq. 3). Due to increasingly poorer precision at the lowest experimental temperatures the sample data were fitted with two overlapping equations. Figure 2 shows the deviation of the sample data from the polynomial (n ranging from 0 to 4) fitted to the data over the temperature range 20 to 110 K and figure 3 the deviation of the data from the polynomial (n ranging from -2 to +10) fitted over the temperature range 60 to 315 K. Heat capacity values calculated from the two equations agreed within 0.02% over the temperature range 80 to 100 K. The heat capacity values obtained from the two equations were joined at 90 K. Figures 2 and 3 show the precision of the measurements and the effect on the final smooth values of a change of 0.025 percent in the values of the heat-capacity measurements for the empty and filled sample vessel.

As mentioned earlier heat effects were observed during the measurements of the heat capacity above about 320 K, and the results that are presented are based on the data obtained between 20 and 315 K subsequent to heating above 320 K. In figure 4 are shown values of heat capacity initially obtained between 80 and 315 K prior to heating above 320 K. Above 150 K, the values of the initial measurements and those after heating above 320 K are shown to be essentially the same. Below 150 K, the values of the initial measurements are somewhat higher. Since no data were obtained initially below 80 K, no comparison can be made below 80 K between the values before and after heating above 320 K. In the range 80 to 315 K, however, the heating above 320 K did not make a large change in the heat capacity of the sample, at most about 0.5 percent at 110 K. The comparison shows that the heat capacity of the sample was not affected grossly when heated above 320 K.

The values of heat capacity of Be_3N_2 , that were generated at evenly spaced temperatures from dQ/dR in accordance with eq (2) were corrected for BeO [5] and Be metal [6] and converted to molal basis. The molecular weight was taken to be 55.05 g. These molal values were analyzed further for extrapolation to 0 K. The final selected values below 20 K are based on the extrapolation of the experimental heat capacity using the Debye heat-capacity function:

$$C = 1943.73 (705/T)^{3}$$
(4)
(J deg⁻¹mol⁻¹)

The values of heat capacity from 0 to 315 K were applied in the usual thermodynamic relations:

$$H_{\rm T} - H_{\rm OK}^{\rm o} = \int_{\rm O}^{\rm T} C \, dt , \qquad (5)$$

$$S_{T} = \int_{0}^{T} (C/T) dT , \qquad (6)$$

and

$$G_{\rm T} - H_{\rm OK}^{\rm o} = (H_{\rm T} - H_{\rm OK}^{\rm o}) - T S_{\rm T}$$
, (7)

to obtain the thermodynamic properties. The values are given in Table 3.

Justice [2] recently reported heat-capacity measurements on α -Be₃N₂ of 96.7 percent purity. The significant impurities were BeO - 2.7%, Be metal - 0.2%, C - 0.1%, and Fe - 0.3%. The entropy at 298.15 K that Justice [2] reported is 8.157 cal deg⁻¹ mol⁻¹, which is in good agreement with that (8.23 ±0.08 cal deg⁻¹ mol⁻¹) found in this investigation. The values of heat capacity obtained also join satisfactorily, in both the magnitude of heat capacity and its slope, with those derived from the high temperature relative enthalpy measurements reported by Douglas and Payne [1]. A combined table of thermodynamic properties from 0 to 1200 K will be submitted in the next report.

- V. References
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Table 1. Summary of Spectrochemical Analysis of the Sample of Beryllium Nitride, Be_3N_2 . Impurity Elements that were Detected.

<u>Element</u>	<u>Wt. %</u>	Element	<u>Wt. %</u>
AL	0.001 - 0.01	Mg	0.001 - 0.01
Ca	< 0.001	Min	< 0.001
Cr	0.001 - 0.01	Ni	0.001 - 0.01
Cu	< 0.001	Si	0.01 - 0.1
Fe	0.01 - 0.1	Ti	0.001 - 0.01

Analysis by: Elizabeth K. Hubbard Spectrochemical Analysis Section Table 2. Chemical Analysis of the Sample of Beryllium Nitride, Be_3N_2 .

Original Sample			
	Wt. %		Wt. %
Be (total)	49.06 49.09	Ν	49.45 49.52 49.84
mean	49.08		49.04
		mean	49.60
Be (free)	0.09 0.11	CL	<0.02
	0.09	F	<0.01
mean	0.10		

Sample Removed from Calorimeter Vessel

Be	(total)	48.72 48.68	Ν	48.97 48.89
	mean	48.70	mean	48.93
Be	(free)	0.04 0.04	Н ₂ О	1.16 1.12
	mean	0.04	mean	1.14
0		2.38 (a	average of 5 determin	inations)

Chemical analysis by: R. A. Paulson Microchemical Analysis Section

Oxygen analysis by activation method: S. Nargolwalla Activation Analysis Section

TABLE 3

THERMODYNAMIC FUNCTIONS FOR BERYLLIUM NITRIDE (BE3N2)

SOLID PHASE

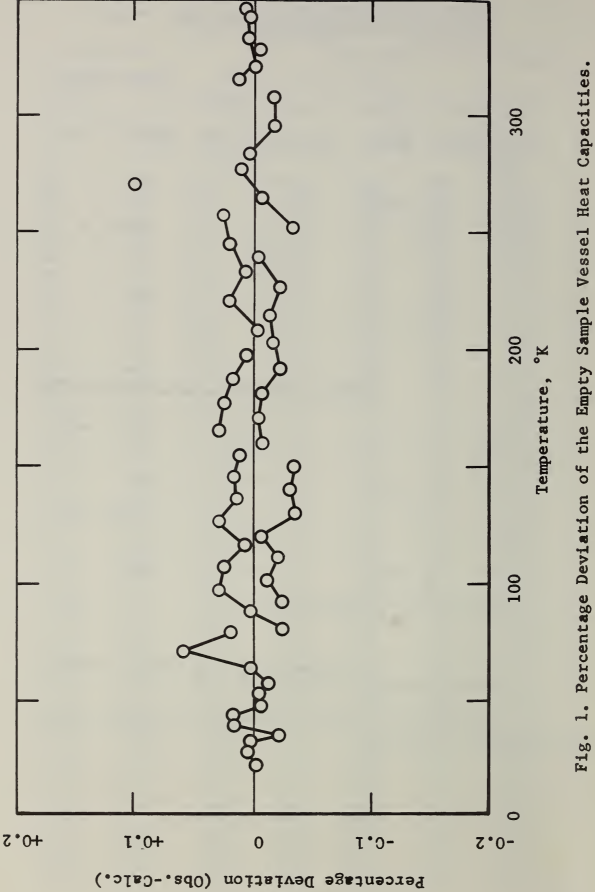
CAL=4.1840 ABS J

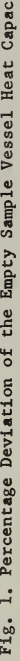
GRAM MOLECULAR WT.≈55.0500 GRAMS

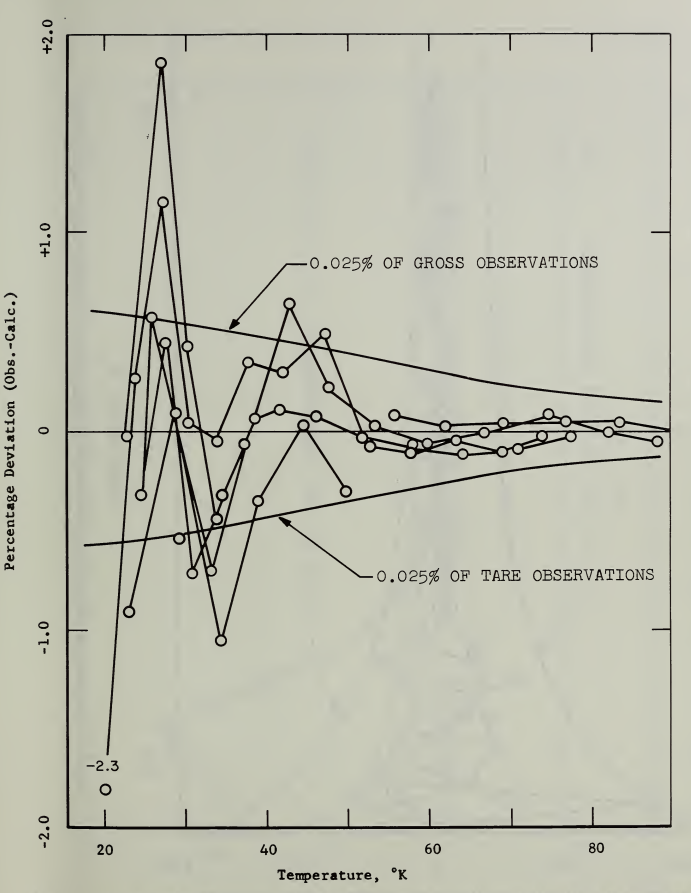
ТΟ	DEG	κ	=	273.15	+	Т	DEG	С
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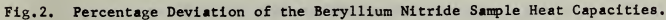
т	-(G ⁰ _T -H ^C ₀)/T	(H ⁰ _T −H ^C ₀)/T	$(s_{T}^{0}-s_{0}^{C})$	(н ⁰ -н ⁰)	cp0	-(G ⁰ _T -H ^C ₀)
DEG K	CAL DEG MOLE	CAL DEG MOLE	CAL DEG MOLE	CAL MOLE D	CAL DEG MOLE	MOLE
$\begin{array}{c} 0 \\ 0 \\ 0 \\ 0 \\ 0 \\ 0 \\ 0 \\ 0 \\ 0 \\ 0 $	000 000 000 000 000 000 002 003 005 007 010 014 018 029 037 045 055 067 080 095 111 130 151 173 198 2265 227 321 358 397 439 483 529 578 629 682 738 796 857 919 984 1051 11190 1263 338 1414 1492 1572 654 1572 655 2377 321 358 397 439 439 439 439 529 578 629 632 738 796 357 919 984 1051 11190 1263 338 1414 492 572 1654 7372 1654 7372 1654 2552 2682 2362 2556 25572 2682 2556 25572 2682 2556 25572 2682 2556 25572 2682 2556 25572 2682 2556 25572 2682 2556 25572 2682 2556 25572 2682 2556 25572 2682 2556 25572 2682 2556 25572 2682 2556 25572 2682 2556 25572 2682 2556 25572 2682 2556 25572 2682 2556 25572 2682 2556 25572 2682 2552 26848 2786 25572 26848 2786 25572 26848 2786 2786 25572 26848 2786 2786 2786 2786 2786 25572 26848 2786 2786 2786 2552 26848 27845 2786	000 000 000 001 003 005 009 014 021 029 040 053 069 088 111 139 171 208 251 299 352 412 478 549 626 710 799 893 994 1.0910 1.325 1.445 1.570 2.5544 2.510 3255 1.445 1.570 2.5544 2.5699 1.325 1.445 1.968 2.107 2.2506 2.544 2.5544 2.5549 3.517 3.962 2.5544 2.5544 2.5549 3.317 3.4767 3.9962 4.125 4.2599 3.317 3.4767 3.9962 4.1259 4.2599 3.317 3.4767 3.9962 4.2599 4.2599 3.317 3.4767 3.9962 4.2599 4.2599 3.317 3.4767 3.9962 4.2599 4.2599 3.317 3.4767 3.9962 4.2599 4.2599 3.317 3.4767 3.9962 4.259 4.2599 4.2599 3.317 3.4767 3.9962 4.259 4.2599 3.317 3.63799 3.962 4.259 5.2788 5.2788 5.442 5.6070 5.7710 5.9347 6.2500	000 000 001 003 007 012 019 028 039 053 071 092 118 148 148 148 226 275 331 393 464 542 628 722 825 1054 1181 1.315 1.457 1.667 1.764 1.928 2.099 2.276 2.460 2.650 2.460 2.650 2.460 2.650 3.463 3.678 3.679 3.678		000 001 004 011 021 035 055 081 115 158 212 280 364 464 584 724 886 1066 $1 \cdot 266$ $1 \cdot 485$ $1 \cdot 725$ $2 \cdot 865$ $3 \cdot 188$ $3 \cdot 525$ $3 \cdot 874$ $4 \cdot 234$ $4 \cdot 602$ $4 \cdot 978$ $5 \cdot 3625$ $3 \cdot 874$ $4 \cdot 234$ $4 \cdot 602$ $4 \cdot 978$ $5 \cdot 3625$ $5 \cdot 750$ $6 \cdot 143$ $6 \cdot 538$ $6 \cdot 935$ $7 \cdot 334$ $7 \cdot 732$ $8 \cdot 129$ $8 \cdot 5255$ $8 \cdot 919$ $9 \cdot 698$ $10 \cdot 465$ $10 \cdot 843$ $11 \cdot 216$ $12 \cdot 555$ $8 \cdot 919$ $9 \cdot 698$ $10 \cdot 843$ $11 \cdot 216$ $12 \cdot 555$ $12 \cdot 950$ $12 \cdot 309$ $12 \cdot 664$ $13 \cdot 3555$ $13 \cdot 693$ $13 \cdot 902$ $14 \cdot 024$ $14 \cdot 6705$ $15 \cdot 897$ $16 \cdot 468$	000 001 006 001 006 001 006 043 089 165 246 985 1.301 2.575 3.404 4.4277 7.186 8.994 1.13.645 16.568 19.941 23.805 28.223 3.172 38.755 44.991 51.919 59.576 68.000 77.228 87.2928 98.228 10.07 122.84 136.58 151.30 67.05 183.833 201.69 220.53 240.67 261.85 28.407 32.40 67.05 183.833 201.69 220.53 240.67 261.85 284.17 307.65 332.31 355.78 505.78 5560.65 573.58 609.39 646.47 684.85 724.52 755.48 807.74 851.31 896.18
285.00 290.00 295.00 298.15 300.00 305.00	2 • 268 2 • 362 2 • 456 2 • 516 2 • 552 2 • 648	5.278 5.442 5.607 5.710 5.771 5.934	7.546 7.804 8.063 8.226 8.322 8.583	1504 • 2 1578 • 3 1654 • 0 1702 • 5 1731 • 3 1810 • 0	14.670 14.985 15.294 15.486 15.598 15.897	646.47 684.85 724.52 750.17 765.48 807.74

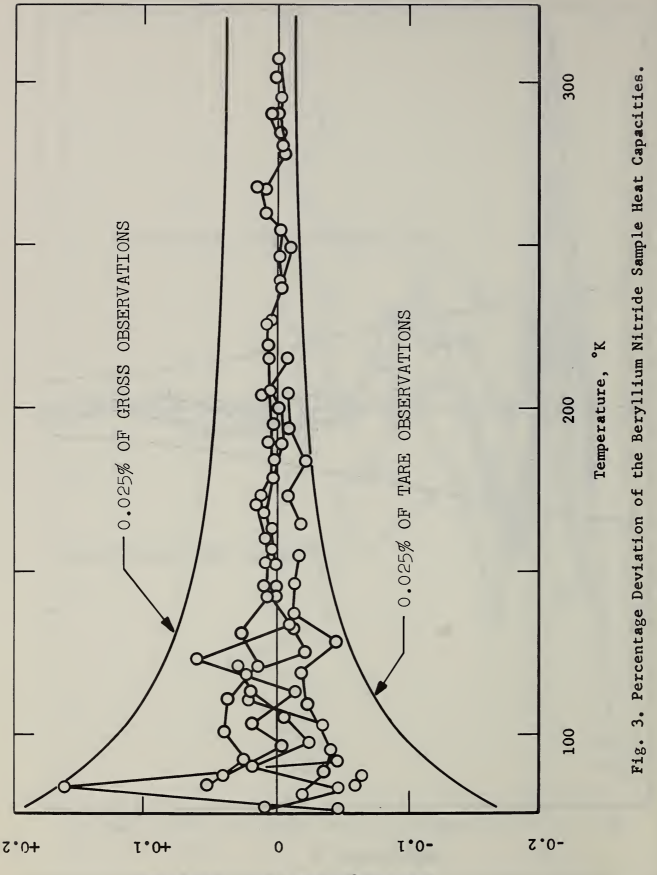
 ${\rm H}_0^{\rm C}$ and ${\rm S}_0^{\rm C}$ apply to the reference state of the solid at zero dek K.



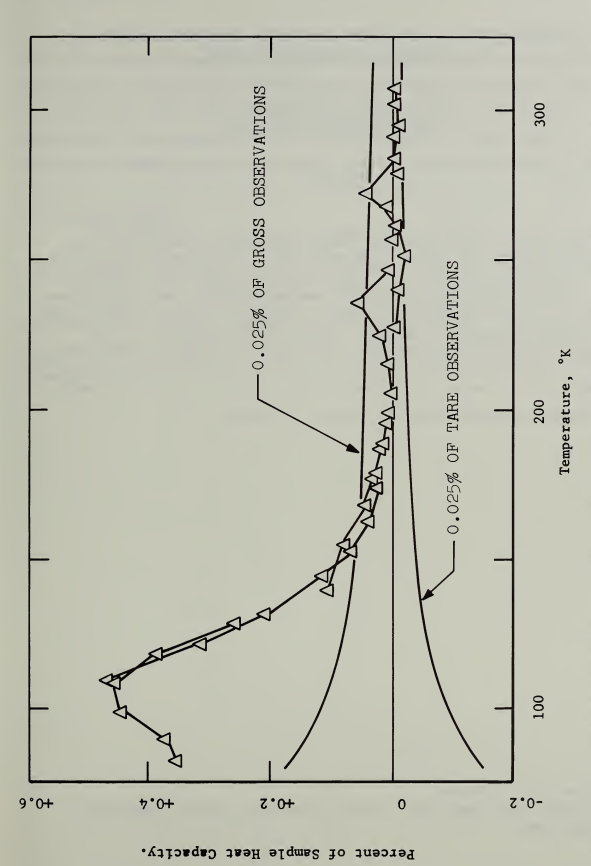


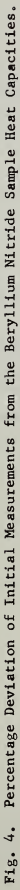






Percentage Deviation (Obs.-Calc.)





Chapter 3

AUTOMATIC DATA REDUCTION APPLIED TO BOMB CALORIMETRY OF ORGANIC FLUORINE COMPOUNDS*

George T. Armstrong Thermochemistry Section

Abstract

Subroutine FARMST, to calculate the combustion energy of organic fluorine compounds from oxygen-bomb calorimetric data, has been prepared following the computational procedure of Good and Scott. The subroutine is written in FORTRAN IV and conforms to the more general program of Shomate for C-H-O-N-S compounds. The Shomate program has been modified to allow introduction of subroutine FARMST, and to change the calculations in several procedural ways including presentation of the result in joules.

* This work was supported, in part, by funds from the Department of the Navy.

1. Introduction

Shomate [1] prepared a general program for reduction of the data from bomb combustion of organic compounds containing the elements C-H-O-N-S, following the computational procedure of Hubbard, Scott, and Waddington [2]. He recently [la] modified his program to FORTRAN IV, and segmented it into subroutines. We have made additional modifications in order to improve the versatility of the program, to render the results in joules, and to introduce the new subroutine FARMST.

We have written subroutine FARMST to allow the reduction of data involving fluorine as well as C-H-O-N. For this reduction a procedure for hand computation was previously reported by Good and Scott [3]. This procedure is followed in FARMST. The procedure does not make provision for handling compounds containing both S and F together.

In order to make the general modifications outlined above new variables were defined in the program. Many of these took the form of additions to the array W. It is useful to place variables here because W has the property that it can be modified by the input data cards, and the values in it are available for a memory print-out associated with any experiment. Array W has 999 terms, of which some 673 are used. A complete listing of the variables in this array is given in Table 1. Of these W(1) to W(22), W(109), W(206), W(226), W(246), W(266), W(286), and W(591) to W(673) have not previously been defined. In addition the definitions of W(572) and W(573) were changed. Tables of data needed in subroutine FARMST were introduced into DATA as block data. These are W51AF, a table of solubilities of CO2 in aqueous HF solutions, obtained from Cox and Head [4]; W87AF, a table of energies of solution of CO2 in aqueous HF solutions, also obtained from Cox and Head [4]; W71AF, a table of dilution energies for aqueous HF, taken from Wagman, et al. [5]; W72A, W73A, W74A, and W75A, tables of properties of aqueous HF solutions taken from Good and Scott [3]. Unfortunately the latter tables are given only at 25°C, so that the computation is limited in the adequacy with which it can be applied to other temperatures.

Values for the solubility of CO_2 are given by Cox and Head [4] for concentrations of HF from 0 to 5%. Values at 6% were estimated by extrapolation and inserted in the array of data in order to allow use of the quadratic interpolation formulas given in the program for interpolation in the region from 4 to 5 percent. The data for the energy of solution of CO_2 was similarly extrapolated.

Data from Wagman, et al. [5] were smoothed. The point at $HF-IOH_2O$ appears to be somewhat irregular and a fictitious value must be used at 0% HF, in order to allow reasonable interpolation from 1 to 2%.

Values at 1 percent intervals were taken from the smooth curve to form the data array to be used in the computations. In all the above cases quadratic interpolation is used. Where temperature is known as a variable, a two-fold quadratic interpolation is used. Table W71AF is not suitable for interpolation in the region 0 to 1% HF.

- 1

Table 1.

TABLE OF THERMOCHEMICAL VARIABLES EXPANDED FOR FARMST SUBROUTINE

Term of the W array	Value or meaning, an	nd units
1	l. The	e constant unity
· 2		e ice point temperature, °K
3		e standard reference temperature, 25°C.
4		e standard atmosphere, Nm ⁻²
5		e standard molar volume of a gas, cm ³
6		e thermochemical calorie, J
7		·
8	The gas constant, R	= 8.314325 J mol ⁻¹ , defined
	by $R = 0.000001$:	* W(4)/W(2).
9	The gas constant, R	= 1.9871715 cal mol ⁻¹ , defined
	by $R = W(8)/W(6)$	
10	The gas constant, R	$= 0.0820560 \text{ cm}^3 \text{ atm, defined}$
	by $R = W(5)/W(2)$	* 0.001
11	Molecular weight of	water, defined by
	W(11) = 2. * W(5)	59) + W(560)
12	Molecular weight of	carbon dioxide, defined by
	W(12) = W(558) +	2. * W(560)
13	Molecular weight of	nitrous acid, defined by
	W(13) = W(559) +	W(561) + 2. * W(560)
14	Molecular weight of	nitric acid, defined by
	W(14) = W(559) +	W(561) + 3. * W(560)
15	-	sulfuric acid, defined by
		59) + W(562) + 4 * W(560)
16	Molecular weight of	•
	W(16) = W(559) +	W(563)
17		
18		
19	38	
20	30	

21	$T_{\rm h}$ - 25. , defined by W(21) = W(148) - 25.
22	$T_h + 27315$, defined by $W(22) = W(148) + W(2)$
23	Mass of substance
24	Mass of substance
25	Moles of substance
26	$V_{1}[bomb]$, dm^{3}
27	$V_{i}[H_{2}O(tota1)]$, dm ³
28	P ₁ [gas], atm
29	M _i [H ₂ O(tota1)], g
30	N ₁ [H ₂ O(tota1)], mol
31	Vi[gas] , dm ³
32	N _i [H2O(vapor)], mol
33	N ₁ [H ₂ O(liquid)] , mol
34	N ₁ [gas], mol
35	$N_{i}[(O_{2} + N_{2}) \text{ (dissolved)}], \text{ mol}$
36	$N_1[O_2 + N_2)$ (total)], mol
37	Nf[HNO3], mol
38	$N_{f}[HNO_{2}]$, mol
39	N _f [H ₂ SO ₄] , mol
40	Nf[H20(liquid)] , mol
41	Mf[Solution)]
42	(Weight per cent) f [H ₂ SO4]
43	(Weight per cent) f [HNO3 + HNO2]
44	(Density) f [solution] , g dm ⁻³
45	Vf [Solution]
46	(Normality) f [H2SO4], mol dm ⁻³
47	(Normality) f [HNO ₃ + HNO ₂], mol dm ⁻³
48	N_{f} [H ₂ O (liquid)]/N _f [H ₂ SO ₄]
49	V_{f} [Gas], dm^{3}
50	N_{f} [CO ₂] (total)], mol
51	Kf [CO2]
52	D _f [CO ₂]
53	к <u>ё</u> [СО ₂]
54	N_{f} [CO ₂ (dissolved)], mol
55	Nf [CO ₂ (gas)], mol

	Table 1. (continued).
56	$N_{f} [(0_2 + N_2) (total)], mol$
57	к _f [0 ₂]
58	D _f [O ₂]
59	к <u>т</u> [0 ₂]
60	N_{f} [(O ₂ + N ₂) (dissolved)], mol
61	$N_{f} [(0_{2} + N_{2}) (gas)], mol$
62	N _f [gas] , mol
63	X_{f} [CO ₂] , (mole fraction)
64	U _f [gas]
65	P _f [gas] , atm
66	G
67	N _f [H ₂ O (vapor)] , mol
68	
69	$[(dE/dp)_T] f [solution], J mol-1 deg$
70	$\triangle E$ (dilution)[HNO ₃ + HNO ₂], J mol ⁻¹
71	ΔE (dilution)[H ₂ SO ₄], J mol ⁻¹
72	
73	
74	E [calorimeter] , J deg ⁻¹
75	E _i [contents] , J deg ⁻¹
76	E _f [contents] , J deg ⁻¹
77	$\triangle E$ (ignition), J
78	Ti , deg C
79	T _f , deg C
80	$ riangle extsf{T}$ (correction) , deg
81	$[\Delta E(vaporization)]_{i}$ [H ₂ 0] , J
82	$\triangle E_i$ [H ₂ O] from 1 to P ₁ , J
83	$ riangle {\tt E}_{i}$ [substence] from 1 to ${\tt P}_{i}$, J
84	[ΔE (solution)] $_{i}$ [$O_{2} + N_{2}$], J
85	$ riangle E_i$ (gas) from O to P $_i$, J
86	riangle E (isothermal bomb process) , J
87	[ΔE (solution)] _f [CO ₂] , J
88	[ΔE (solution)] f [O ₂ + N ₂], J
89	$\triangle E$ [solution] from 1 to P _f , J
90	[ΔE (dilution)] _f [HNO ₃ + HNO ₂], J
91	$[\triangle E (dilution)] f [H_2SO_4], J$

92	[ΔE (decomposition)] f [HNO ₃ + HNO ₂], J
93	ΔE_{f} [gas] from P_{f} to 0, J
94	[ΔE (vaporization)] f [H ₂ 0], J
95	Sum of W (81) to W (94) inclusive, J
96	Combustion energy from auxiliary material , J
97	Combustion energy from fuse , J
98	Combustion energy from compound , J
99	$\triangle Ec/M [compound] at Th , J$
100	$ riangle$ Ec [compound] at T $_{ m h}$, J
101	$ riangle Hc$ [compound] at $ extsf{T}_{ extsf{h}}$, J
102	\triangle Hc [compound] at 25 deg C , J
103	$ riangle H_{f}$ [compound] at 25 deg C , J
104	A subscript (\mathtt{C}_{A}) in empirical formula of bomb contents
105	в (н _в) и и и и и и
106	c '' (o _c) '' '' '' '' '' ''
107	D'' (N _D)''''''''''''''''''''''''''''''''''''
108	e '' (s _e)'' '' '' '' '' ''
109	F''(F _F)'''''''''''''''''''''''''''''''''''
110	() '' () '' '' '' '' ''
111	() '' () '' '' '' '' '' ''
112	() 11 () 17 17 11 11 11 11
113	N_{f} [N_{2}^{*}], extra nitrogen, mol
114	Weight of platinum in air , g
115	Weight of glass in air , g
116	Weight of other materials in air , g
117	Mass of platimum (weight in vacuum) , g
118	Mass of glass (weight in vacuum) , g
119	Mass of other materials (weight in vacuum) , g
120	Volume of platinum , dm ³
121	Volume of glass , dm ³
122	Volume of other materials , dm ³
123	Free space in rigid glass bulb , dm ³
124	Nitrogen in free space , mol
125	Initial period rate , deg min ⁻¹
126	Final period rate , deg min ⁻¹

127	$X [O_2 + N_2]$, mole fraction
128	X [N $\stackrel{\star}{2}$], mole fraction
129	$Mu [O_2]$, see item W(64)
130	A (C _p of solution) , see item W(76) , cal deg ⁻¹
131	E [calorimeter] * $(T_i - T_f - \Delta T (correction))$, J
132	E_i [contents] * ($T_i - T_h$), J
133	E_{f} [contents] * (T_{h} - T_{f} + $ riangle T$ (correction)) , J
134	$ riangle$ H (vaporization) [H $_2$ O] , for W(81) and W(94) ,
135	Cooling constant , min ⁻¹
136	Area under Temperature vs. Time curve, deg min
137	Time (start of main period), min
138	Time (end of main period) , min
139	Temperature (start of main period) , deg
140	Temperature (start of main period) , deg
141	Mid-time (time after start of main period) , min
142	Rotation time, min
143	Heat of bomb rotation , deg min ⁻¹
144	Sum of terms W(81) , W(82) , W(84) , W(85) , W(88) , W(89),
	W(93), W(94), W(83), J
145	Corrected temperature rise , deg
146	Pressure to which the bomb was filled , atm
147	Temperature at which the bomb was filled , deg C
148	\mathtt{T}_{h} , the selected temperature , deg C
149	\mathtt{T}_{h} , predetermined value of \mathtt{T}_{h} , deg C
150	Pressure of N2 in glass bulb , atm
151	Coefficients used to calculate values of items at
152	Coefficients used to calculate values of items at temperature T _h from values at 20, 25, and 30 deg C.
153	
154	Selected ratio N_{f} [H ₂ O]/N _f [H ₂ SO ₄]
155	Actual ratio N _f [H ₂ O]/N _f [H ₂ SO ₄]
156	ΔH_{f} [H ₂ SO ₄] at selected ratio, cal mol ⁻¹
157	$\triangle H_{f}$ [H ₂ SO ₄] at actual ratio , cal mol ⁻¹
158	Combustion energy from 1st impurity , J
159	Combustion energy from 2nd impurity , J
160	Density of air (if zero, item 567 is used), g cm ^{-3}

161	$C_{ m p}$ of H_2SO_4 solution (selected concentration) , cal g ⁻¹ deg ⁻¹
162	$\Delta c_p / \Delta T$ of H ₂ SO ₄ solution (selected concentration) , cal g ⁻¹ deg ⁻²
163	Density of H_2O , g dm ⁻³
164	Reference ratio N _f [H ₂ O] / N _f [HF]
165	
166	
167	
168	
169	
170	
171	
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185 186	
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	lable 1. (continued).
195	
196	
197	
198	
199	
200	
201	Number of C atoms in compound
202	rr rr _A rr rr rr
203	
204	ri ir N [°] ri ir ri
205	'' '' S '' '' ''
206	, , , , , , , , , , , , , , , , , , ,
207	··· ·· () ··· ··
208	
209	· · · · · · · · · · · · · · · · · · ·
210 .	Mass of compound (weight in vacuum) , g
211	Molecular weight of compound
212	Moles of compound
213	Density of compound , g cm ⁻³
214	Volume of compound , dm ³
215	Weight of compound (in air) , g
216	$c_{ m p}$ of compound , cal g ⁻¹ deg ⁻¹
217	$(\triangle V/\triangle T)_p$ of compound , dm ³ deg ⁻¹ g ⁻¹
218	$(\triangle E / \triangle P)_T$ of compound , J atm ⁻¹ g ⁻¹
219	ΔE_{c} [compound] , J g ⁻¹
220	$\triangle E_{c}$ [compound] , J mol ⁻¹
221	Number of C atoms in auxiliary material
222	11 11 H 11 11 11 11
223	··· ··· 0 ··· ··· ··· ··· ··
224	
225	11 11 S 11 11 11
226	11 11 F 11 11 11 11
227	
228	11 11 () 11 11 11
	11 11 () 11 11 11 11

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Table 1. (continued).

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230	Mass of auxiliary material (weight in vacuum) , g
231	Molecular weight of auxiliary material
232	Moles of auxiliary material
233	Density of auxiliary material , g cm ⁻³
234	Volume of auxiliary material , cm ³
235	Weight of auxiliary material (in air) , g
236	C_p of auxiliary material, cal g^{-1} deg ⁻¹
237	$(\Delta V/\Delta T)_p$ of auxiliary material , dm ³ deg ⁻¹ g ⁻¹
238	$(\Delta E / \Delta P)_T$ of auxiliary material , J atm-1 g ⁻¹
239	AE [auxiliary material], Jg ⁻¹
240	△E _c [auxiliary material] , J mol ⁻¹
241	Number of C atoms in fuse
242	11 11 H 11 11
243	
244	11 11 N 11 11 11
245	'' '' S '' '' ''
246	···············
247	
248	
249	
250	Mass of fuse (weight in vacuum) , g
251	Molecular weight of fuse
252	Moles of fuse
253	Density of fuse , g cm^{-3}
254	Volume of fuse , dm ⁻³
255	Weight of fuse (in air) , g
256	C_p of fuse , cal g ⁻¹ deg ⁻¹
257	$(\Delta V/\Delta T)_p$ of fuse, dm ³ deg ⁻¹ g ⁻¹
258	$(\Delta E / \Delta P)_T$ of fuse, dm ³ atm ⁻¹ g ⁻¹
259	ΔE_c [fuse], Jg ⁻¹
260	△E _c [fuse] , J mol ⁻¹
261	Number of C atoms in first impurity
262	ti ti H ti ti ti ti
263	
264	N
	•

265	Number of S atoms in first impurity
266	ri it Frintin in th
267	
268	· · · · · · · · · · · · · · · · · · ·
269	· · · · · · · · · · · · · · · · · · ·
270	Mass of first impurity (weight in vacuum) , g
271	Molecular weight of first impurity
272	Moles of first impurity
273	Density of first impurity , g cm ⁻³
274	Volume of first impurity , dm ³
275	Weight of first impurity (in air) , g
276	C_p of first impurity , cal g ⁻¹ deg ⁻¹
277	($\Delta V/\Delta T$) $_{ m p}$ of first impurity , dm 3 deg $^{-1}$ g $^{-1}$
278	$(\Delta E/\Delta P)_{\mathrm{T}}$ of first impurity , J atm ⁻¹ g ⁻¹
279	$\triangle E_c$ [first impurity] , J g ⁻¹
280	ΔE_c [first impurity] , J mol ⁻¹
281-	Number of C atoms in second impurity
282	II II H II II II
283	· · · · · · · · · · · · · · · · · · ·
284	11 11 N 11 11 11
285	11 [°] 11 S 11 11 11
286	ii ir Filli II II
287	···············
288	· · · · () · · · · · · · · · · · · · ·
289	
290	Mass of second impurity (weight in vacuum) , g
291	Molecular weight of second impurity
292	Moles of second impurity
293	Density of second impurity , g cm ^{-3}
294	Volume of second impurity , dm ³
295	Weight of second impurity (in air) , g
296	C_p of second impurity , cal g ⁻¹ deg ⁻¹
297	$(\Delta V/\Delta T)_p$ of second impurity , dm ³ deg ⁻¹ g ⁻¹
298	$(\Delta E / \Delta P)_T$ of second impurity , J atm ⁻¹ g ⁻¹
299	ΔE_{c} [second impurity], J g ⁻¹

. 46

300	ΔE_c [second impurity], J mol ⁻¹
301 - 400	Recorded times at which corresponding temperature
	readings in locations 401-500 were taken, minutes
401-500	Recorded temperatures of the initial, main, and final
	periods, introduced in ohms and converted by the
	program to degrees C.
501-550	Bridge dial corrections , ohms
551	Ten-ohm dial setting, nominal value, ohms
552	Ten-ohm dial setting correction , ohms
553	Average correction to convert N-only or R-only readings
	to $(N + R) / 2$.
554	Bridge-zero correction
555	
556	
557	
558	Atomic weight of C (12.01115)
559	Atomic weight of H (1.00797)
560	Atomic weight of O (15.9994)
561	Atomic weight of N (14.0067)
562	Atomic weight of S (32.064)
563	Atomic weight of F (18.9984)
564	Atomic weight of
565	Atomic weight of
566	Atomic weight of
567	Density of Air , 0.0012 g cm ⁻³
568	Density of weights , 8.4 g cm^{-3}
569	Density of platinum ,2145 g cm ⁻³
570	Density of glass , 2.24 g cm ⁻³
571	Density of other material , 1.0 g cm^{-3}
572	Density of water g/1000 cm ³
573	
574	0.0
575	$\Delta E_c/M$ for benzpic acid calculated by program , J g ⁻¹
576	$\Delta E_c/M$ for benzoic acid, certified value = 26.434 kJ g ⁻¹
577	C_p of platinum = 0.0325 cal g ⁻¹
578	C_p of glass = 0.17 cal g ⁻¹

Table 1.	(continued).
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579	C_p of other material , = 0.0 cal g ⁻¹
580	C_{p} of CO ₂ , gas, at 25°C , 0.20163 cal g ⁻¹ deg ⁻¹
581	$\triangle C_{\rm D}$ / $\triangle T$ of CO ₂ = 0.0 cal g ⁻¹ deg ⁻²
582	$C_{\rm p}^{\rm P}$ of O_2 , gas, at 25°C , 0.219375 cal g ⁻¹ deg ⁻¹
583	$\Delta C_p / \Delta T \text{ of } O_2 = 0.0 \text{ cal } g^{-1} \text{ deg}^{-2}$
584	$C_{\rm p}$ of N ₂ , gas, at 25°C, 0.248465 cal g ⁻¹ deg ⁻¹
585 .	$\Delta C_p / \Delta T$ of $N_2 = 0.0$ cal g ⁻¹ deg ⁻²
586	$C_{\rm p}$ of H ₂ O, liquid, at 25°C , 0.99895 cal g ⁻¹ deg ⁻¹
587	$\Delta C_p / \Delta T$ of $H_2 0 = 0.0$ cal g ⁻¹ deg ⁻²
588	$\Delta C_p / \Delta T$ of smaple = 0.0 cal g ⁻¹ deg ⁻²
589	$\Delta H_{f}[CO_{2} (gas)] = -2137.0552 \text{ cal g}^{-1}$
590	$\Delta H_{f}[H_{2}0(1)] = -3792.04 \text{ cal g}^{-1}$
591	$\triangle H \text{ vap } [H_2 0] = 10 \ 177 \ \text{cal mol}^{-1}$
592	$\triangle E \text{ decomp [HNO_3]} = 14 074 \text{ cal mol-1}$
593	$\triangle E$ decomp [HNO ₂] = (-6 600 cal mol ⁻¹
594	ΔE vap [HF from HF (aq)] = 11 000 cal mol ⁻¹
595	$\triangle E_{f}$ [HF in 10H ₂ 0] = -76 235 cal mol ⁻¹
596	\triangle H or \triangle E hydrolysis [CF ₄] = -41 500 cal m01 ⁻¹
597	C_{p} [CF ₄] = 14.7 cal mol ⁻¹ deg ⁻¹
598	$\Delta \hat{c}_p / \Delta T [CF_4] = 0$
599	
600	ar 14
601	C _i [HF (vap)]
602	(Normality) _i [HNO ₃], m mol cm ⁻³
603	K ₁ [0 ₂]
604	N _i [HF (vap)], mol
605	$D_i [O_2]$
606	N _f [HF (total)], mol
607	N _f [CF4 (total)], mol
608	N _i [HF (total)], mol
609	N _f [H ₂ O (total)], mol
610	(Weight per cent) _f .[INO3]
611	(Weight per cent) [HF]
612	N _f [HF (ag)] , mol
613	N _f [H ₂ 0] /N _f [HF], actual ratio

614	N _f [H ₂ O] / _f [HF] , selected ratio
615	C_{f} [IF (vap)]
616	(dE/dp) _i [solution], cal atm ⁻¹
617	(dE/dp) _f [solution], cal atm ⁻¹
618	A _i [solution]
619	A _f [solution]
620	N _f [IF (gas)], mol
621	(Normality) _f [HF], mmol cm ⁻³
622	$\Delta E_{f} [CO_{2}] / N_{f} [CO_{2}] cal$
623	Mole fraction (X) [CF ₄]
624	△E dilution [HF (final)],cal
625	N_{f} [H ₂ 0] /N _f [HF] , selected ratio
626	Weight per cent HF at selected ratio
627	Weight per cent HF at N_{f} [H ₂ 0] / N_{f} [HF] = 10
628	$\triangle H_{f}$ [HF (aq, selected ratio], cal mol ⁻¹
629	N _i [H ₂ 0], mol
630	$\triangle H_{f}$ [HF (aq, selected ratio)] - $\triangle H_{f}$ [HF (aq, 10H ₂ 0)], cal mol ⁻¹
631	$\triangle H_{f}$ [HF (aq, 10H ₂ 0)] - $\triangle H_{f}$ [HF (aq, actual ratio)],
	cal mol-1
632	M _i [solution], g
633	(△E vap) _i [HF] , cal mol ⁻¹
634	(AE vap) _f [HF] , cal mol ⁻¹
635	$\triangle E$ dilution [HF (total)], cal mol ⁻¹
636	M _i [HF], g
637	Molar coefficient of CF ₄ in equation
638	
639	Correction for CF4 hydrolysis , cal
640	C_p [HF (aq, selected ratio)], cal g ⁻¹
641	Molar coefficient of O ₂ in equation
642	Molar coefficient of H_2O adjusted for hydrolysis of CF_4 .
643	Molar coefficient of CO ₂ adjusted for hydrolysis of CF ₄
644	Molar coefficient of N ₂ in equation
645	Molar coefficient of HF adjusted for hydrolysis of CF ₄
646	4
647	

648	AN[gas], mol
649	ΔH_{f} [CO ₂ (g)], cal mol ⁻¹
650	ΔH_{f} [H ₂ O (1)], cal mol ⁻¹
651	Moles idealized actual HF (aq, selected ratio)
652	Moles idealized actual H20
653	Moles idealized actual CO2
654	Moles H ₂ O reacted (excludes water in HF (aq, selected ratio))
655	M _i [HNO ₃] , g
656	(Weight per cent) _i [HF]
657	(Weight per cent) _i [HNO ₃]
658	P_i [solution], g cm ⁻³
659	V _i [solution], 1000 cm ³
660	g _i
661	\triangle H at 25° C [adjusted reaction], J mol ⁻¹
662	$\triangle E$ at 25° C [adjusted reaction] , J mol ⁻¹
663	$\triangle E$ at 25° C [unadjusted reaction] , J mol ⁻¹
664	$\triangle H_{f}$ [compound] at 25° C, J mol ⁻¹
665	ΔE_{f} [compound] at 25° C, J mol ⁻¹
666	
667	
668	\triangle H [Reaction] at 25° C , adjusted for hydrolysis of CF4, cal mol ⁻¹
669	AE [Reaction] at 25°C, '' '' '' ''' ''' ''' cal mol ⁻¹
670	\triangle E [Reaction] at 25° C , unadjusted , cal mol ⁻¹
671	$\triangle H_{f}$ [Compound] at 25° C , cal mol ⁻¹
672	ΔE_{f} [Compound] at 25° C , cal mol ⁻¹
673	\triangle H [Reaction] at 25° C, unadjusted , cal mol ⁻¹

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3. Subroutine FARMST

The practice initiated by Shomate, of numbering variables in the W array, so far as possible, to conform to the number of the calculational step given by Hubbard, et al. [2] is continued here by numbering them to conform to the computational steps given by Good and Scott [3]. These authors have made their work mutually consistent to a large degree. It is not possible to follow this practice throughout, however, because in order to retain mutual consistency with Hubbard, et al., Scott and Good introduced multiple variables of the same step number. We have assigned some of these variables higher numbers (above W(590)) more or less as they are encountered in the calculation. The subroutine begins with the calculation of W(29), as all prior steps are identical in the two programs. The subroutine then bypasses the major part of the program called HOWARD, following a parallel but different set of calculations to arrive at final energy values. Where an identity could be established between a variable of FARMST and a variable of HOWARD, i.e. as to meaning, the same variable designation was used in FARMST.

In calculating the composition of the initial bomb contents and the final bomb contents, the program assigns temporary values to the amounts of water and HF present in the liquid phase, and of $CO_2(g)$, $[N_2 + O_2](g)$, $H_2O(g)$, and total gas, present in the gas phase. These are then adjusted after one pass through the calculation, and the calculation of the quantities is repeated.

In mathematical expressions used at various stages of the computations, Good and Scott [3] used expressions valid at 25° C, whereas in similar instances Hubbard, et al. [2] used expressions containing terms with temperature as a parameter. Where the correspondence between equations used showed that the equation used by Hubbard, et al. reduced at 25° C to that used by Good and Scott, we used the more general expression of Hubbard, et al., as did Shomate [1, 1a].

The modifications made in the main program HOWARD and subroutines INPUT, CONVRT, MIDTIM, OUTPUT, and DATA, may be classified as (1) those necessary to introduce an additional subroutine, which do not modify the mode of calculation, and (2) those which are matters of convenience or preference in causing the calculations to have certain desirable features.

In classification (1) we refer to changes such as program statements necessary to allow the program to recognize the new variables introduced in the subroutine. These are matters of programming technique and include definitions of certain variables at suitable times, in the program or in block DATA subroutine, transfer statements dependent on the presence of F in the compound, and suitable output statements to give the appropriate point out.

In classification (2) changes were introduced in order to change the details of the calculation in certain ways.

Calculations were put in terms of joules wherever practicable, Where input data, such as tables, were in calories in the original sources, these were converted to joules before finally being added as energy terms. The program was restated so that the energy equivalent of the calorimeter would be in joules per degree, the energies of combustion of fuse, auxiliary materials, and impurities were in joules per gram, the ignition energy was in joules, and so on. Output statements were modified to indicate the new units. A certain number of output variables were expressed in both joules and calories, and so new variables were introduced to allow this.

Universal constants and defined constants were introduced in the block data program where possible, rather than as terms in the numerical coefficients of equations throughout the program and subroutines. The algebraic variables were then used in the equations. The reason for this is to permit ready change of a constant throughout the program, if it should ever be desirable. Thus the value of the standard molar volume of a gas, $22413.6 \text{ cm}^3 \text{ mol}^{-1}$, is given the name W(5), and the

variable W(5) is used wherever the standard molar volume occurs in an equation. A change in the selected value of W(5) is thus readily accomplished. Other universal constants introduced in this way include the ice point, 273.15°K, the standard reference temperature, 298.15°K, the standard atmosphere,101325 Nm⁻², the ratio of the calorie to the joule, 4.184 J cal⁻¹, and the atomic weights of the elements. Derived quantities such as the gas constant, R, and the molecular weights of compounds involved in the calculations are calculated at an early stage of the program HOWARD.

The molar coefficients of the reactions involved and the change in moles of gas present were calculated early in the main program.

In INPUT subroutine a statement is introduced causing the machine to test whether Ti, Tf and ΔT (corr) have been specified. If all three are specified, as by means of a "9" card the program omits their calculation, thus bypassing subroutines CONVRT and MIDTIM.

5. Debugging

Subroutine FARMST and the whole program were "debugged" by running the sample calculation given by Good and Scott [3]. In order to do this, the data used by Good and Scott were carefully examined for conformity to the program requirements. Several adjustments in the program were made temporarily to assure that the program would duplicate the results obtained by Good and Scott. These included setting the density of air equal to zero for the buoyancy correction so that masses instead of weights in air could be used for the amounts of substances introduced. A small discrepancy in the bomb volume (apparently due to neglecting the platinum volume) was discovered. An adjusted bomb volume was then selected. Values of the CO₂ solubility and heat of solution were dubbed in to allow the calculation to simulate Good and Scott's.

6. Input Data Cards

Table 2 gives a printout of the input data cards for a run on perfluoropiperidine. Each card is coded in column 1 with a digit from 1 to 9 which defines the class of information found in it, following Shomate's practice. The "1" card contains data concerning the fuse and auxiliary material. The only deviation from Shomate's practice in the "1" card is the use of energies of combustion of these two substances in joules per gram instead of calories per gram. These values are found in columns 30-35 and 66-72 respectively. The "2" card contains the name of the compound, the numbers of C, H, O, N and S atoms in it, and some bomb conditions. The number of F atoms is introduced later in a "9" card. (Table 2 approximates the card format).

The "3" card contains data on the amounts of substances present. The format of this card has been slightly modified to permit weights to be expressed to micrograms. The revised format is as follows:

Column	Format	Quantity
1		3
2-3	A2	Run Number
4-11	F8.0	Weight of compound (in air), g
12-18	F7.0	Weight of fuse (in air), g
19 - 25	F7.0	Weight of auxiliary material (in air), g
26-32	F7.0	Weight of platinum (in air), g
33 - 39	F7.O	Weight of glass (in air), g
40-43	3 PF 4.0	Volume of water placed in bomb, cm ³
44-48	0 P F5.0	Ignition energy, J
49-53	6 p f7.0	Moles of HNO_3 formed (X 10^6)
54	Fl.O	Moles of HNO ₂ formed (X 10 ⁶)
55 - 59	10 PF 5.0	Pressure to which bomb is filled, atm

. 11.774.887	2 PERFLUOROPI	3 1 5.77703	4 28948	I SIHT /	' TEST OF	TEST OF	8 001673	81 001673	9 026 . 35381	9 078 2	9 649-94051.
t 。887	JOROPIPEF	20771	Ъ	ES A SAME		JF SHOMAT		N	35381.34	078 22.89480	+051.
1°5	[PE R I D I NE	,00114	Т	TE OF TI	SUBROU.	E PROGRA			074	670	650-(
• 4		.12864		HE PRIN	LINE US	AM USIN			1671	24.8	650-68317。
с °0				IO TUO-TV	ING DAT	IC TI, TH			16746。38	24 . 89078	
16945.10	2	19 . 592	3289	TAINED	A FROM W	, AND D			567 0	080	
ထိ		0°0	328948 20.	FROM TH	D GOOD	r (corr)			0°0	0° 00903	
4	1 1.730.	9.983.72		E COMBUST	AND DW SC	SUPPLIEI					
1.38	1 1.730.2514.955 25.			THIS IS A SAMPLE OF THE PRINT-OUT OBTAINED FROM THE COMBUSTION PROGRAM.	FARMST SUBROUTINE USING DATA FROM WD GOOD AND DW SCOTT EXAMPLE	SHOMATE PROGRAM USING TI, TF, AND DT (CORR) SUPPLIED ON '9' CARD					
° 315. 11	25.	1149. 40.0025.00		tam	JLE	ARD					

606 。045037 563 19。00

560 16.000 561 14.008

559 1.008

558 12.010

σ

9 614 10.

286 0.

266 0.

246 O.

226 O.

9 206 11.

Table 2

228lt2。2

60 - 64	F5.0	Temperature at which bomb is filled, C
65-71	F7.0	Density of air, for reducing weights to vacuum, g cm ⁻³
72	Fl.O	Free space, if any, in rigid glass sample bulb, cm ³

The moles of HNO_2 formed and the volume of free space in the bulb are usually zero, so those have been reduced to a single column each.

Card "4" contains temperature-time data points. The illustration shows a possible format if Ti, Tf and ΔT (corr) are given. A single temperature and time are given in the initial field (terminated by 1 in column 16) and in the final field (initiated by 3 in column 30). These may not be necessary. The actual values of Ti, Tf and ΔT (corr) are introduced later in a "9" card.

Cards "5" and "6" are not used in this program. Card "7" gives commentary which appears in the printout (see Table 3).

Cards "8" and "81" call for memory printout of the W array, as fixed point and floating point format, respectively. In the example, the W array from W(001) to W(673) inclusive is called for.

Cards "9" are used for direct introduction of values into the W array. The element of the W array is first indicated by its number, then the value to be introduced is listed. In the example, the bomb volume is given in W(026). (This vlaue differs from that used by Good and Scott, by the volume of platinum, which they apparently ignored). The energy equivalent of the calorimeter in joules per degree is given in W(074). A fictitious density of air is given in W(567) to permit masses (weight in vacuum) of the various substances in the bomb to be used as given by Good and Scott.

Ti, Tf, and ΔT (corr) are introduced in W(078), W(079), and W(080), respectively. Enthalpies of formation of carbon dioxide (g), and water (1) are introduced in cal mol⁻¹ in W(649) and W(650). W(614) is given the value for the selected ratio of H₂O/HF to be used in the calculations.

Atomic weights used by Good and Scott are introduced in W(558) to W(563). The number of F atoms in the compound of interest is placed in W(206). Other quantities can be introduced at will for checking or other purposes.

7. Output Data

Table 3 is the format of the summary printout of the results of the calculation. Some changes are made from the format suggested by Shomate. The number of decimal figures and the units of several of the quantities are different. Equations are given for the idealized bomb reaction and for the reaction adjusted for hydrolysis of any CF₄ formed. Certain corresponding energy changes are given. In the program these 'are calculated in joules first and then converted to calories. Values are given at Th and at 298.15°K, which in this case are the same. The results give for the internal energy change of the idealized reaction the value -354.54 kcal mol⁻¹, which may be compared with the value -354.51 given by Good and Scott.

The calculations were compared with those illustrated by Good and Scott at several stages in the development of the program. Only minor differences occur, mostly due to rounding. The difference of the value 354.54 from 354.51 kcal mol⁻¹ is apparently due to (1) an error in the value for item 93 listed by Good and Scott, and (2) the use of a different solution energy for CO₂ and a different dilution energy for HF (aq). (Table 3 approximates the actual format). TABLE 3

PERFLUOROPIPERIDINE

RUN NO. 1		22.89480 DEG. CENT.	24.89078 DEG. CENT.	.00903 DEG. CENT.	1.98695 DEG. CENT.	.000000 PER MIN.	16746.38 J/DEG		+ CF4	2.1983288	JOULES/GRAM	-5240.59	KCAL/MOLE		-352.61	-354° 54		+ N2 .	. 5000000	KCAL/MOLE	-4143° 84	-445.77
	(IN VACUO) 5.777030 GRAMS							CEPT N2	= CO2 + HF . 10H20	2.8016713 2.2066847	JOULES JOULES	K) -30275.05	KJ/MOLE	,	-1475.33	K) -1485.39	HYDROLYSIS OF CF4	= CO2 + HF, 10H20	0 5.0000000 10.999999	TION KJ/MOLE	-1857. 04	K) -1865.10
C5 NI FII	WEIGHT OF SAMPLE (IN AIR) 5.77030 (INITIAL TEMPERATURE	FINAL TEMPERATURE	CORRECTION ON TEMPERATURE RISE	CORRECTED TEMPERATURE RISE	COOLING CONSTANT X 10 EXP 3	ENERGY EQUIVALENT USED	COEFFICIENTS OF IDEALIZED REACTION EXCEPT N2	COMFOUND + 02 + H20	1.0000000 2.2500000 23.1701900 2.8016713	STANDARD ENERGY CHANGES OF ABOVE REACTION	INTERNAL ENERGY CHANGE (298.15 DEG K)	· · ·	ENTHALPY CHANGE (298.15 DEG K)	ENTHALPY CHANGE (298.15 DEG K)	INTERNAL ENERGY CHANGE (298.15 DEG K)	COEFFICIENTS OF REACTION ADJUSTED FOR HYDROLYSIS OF CF4	COMPOUND + 02 + H20	1.0000000 2.2500000 115.4999990	STANDARD ENERGY CHANGES OF ABOVE REACTION	ENTHALPY CHANGE (298.15 DEG K)	INTERNAL ENERGY CHANGE (298.15 DEG K)

Table 3 (continued)

STANDARD ENTHALPY OF FORMATION (298.15 DEG K)	-2047.03	-489.25
STANDARD INTERNAL ENERGY OF FORMATION (298.15)	-2076.78	-496° 36
CORRECTION FOR AUXILIARY MATERIAL(S)	-2938. 42	JOULES
CORRECTION FOR FUSE	-19.32	JOULES
CORRECTION FOR CARBON DIOXIDE	21.08	JOULES
CORRECTION FOR NITRIC ACID	67。66	JOULES
CORRECTION FOR DILUTION	1° 71	JOULES
CORRECTION FOR OTHER REDUCTIONS	74.02	JOULES
CHANGE OF INTERNAL ENERGY FOR ISOTHERMAL BOMB PROCESS IS -33274.22 -133.88 + 7.12 + 3.72, 0R -33397.26	CESS IS 3397.26	JOULES
MIDTIME .000		
אא מרושה שור און	UT COMPUTENCE	

TEST OF SHOMATE PROGRAM USING TI, TF, AND DT (CORR) SUPPLIED ON '9' CARD. D TEST OF FARMST SUBROUTINE USING DATA FROM WD GOOD AND DW SCOTT EXAMPLE THIS IS A SAMPLE OF THE PRINT-OUT OBTAINED FROM THE COMBUSTION PROGRAM

References

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STRUCTURE OF THE ALKALI HYDROXIDES. III. MICROWAVE SPECTRUM OF GASEOUS ROOH David R. Lide, Jr. and Chi Matsumura

National Bureau of Standards, Washington, D. C.

The microwave spectrum of gaseous RbOH and RbOD have been studied in a high temperature microwave spectrometer. The spectrum indicates a linear or very near linear molecule with a large amplitude, low-frequency bending vibration. The RbO bond length is 2.30 Å and the OH bond distance is 0.97 Å. The observed frequencies and their assignments for ⁸⁵RbOH and ⁸⁵RbOD are given in Table 1. Similar data for ⁸⁷RbOH and ⁸⁷RbOD are given in Table 2. The effective rotational constants and nuclear quadrupole coupling constants for various vibrational states of ⁸⁵RbOH and ⁸⁷RbOD are found in Table 3. Table 4 gives the values of q for excited vibrational states involving v_2 . A fit of $B_0 v_2^{\frac{4}{2}0}$ to a power series for ⁸⁵RbOH and ⁸⁷RbOD is given in Table 5. Variation of rotational constants with v_1 is shown in Table 6, while the structural parameters found for the species is given in Table 7.

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Observed frequencies in ⁸⁵ RbOH and ⁸⁵ RbOD					
	J = 2 → 3	. •			
State	$F \rightarrow F$	⁸⁵ _{RЬОН}	85 _{Rbod}		
	5/2 5/2	37744.8			
2	7/2 (7/2, 9/2) 9/2 11/2	37741.2	34324.6		
(00 ⁰ 0)	5/2 7/2	37738.3	34321.6		
	3/2 5/2 1/2 3/2	37734.9	34317.9		
	9/2 (7/2, 9/2)	37728.6	34311.9		
	5/2 3/2 3/2 1/2	37600.7			
	5/2 5/2 9/2 11/2	37594.3	34301.9		
(01 ¹ 0) <i>l</i> +	7/2 (7/2, 9/2)	37591.3	34299.0		
	5/2 7/2 3/2 5/2	37589.4	34297.3		
	9/2 (7/2, 9/2)	37584.7	34292.3		
	5/2 3/2 3/2 1/2	37541.0			
	5/2 5/2 9/2 11/2	37534.9	34242.9		
(01 ¹ 0) <i>l</i> -	7/2 (7/2, 9/2)	37532.0	34240.1		
	5/2 7/2 3/2 5/2 9/2 (7/2, 9/2)	37530.6	34238.5		
	9/2 (7/2, 9/2)	37525.5	34233.5		
	7/2 (7/2, 9/2)	37430.5	34242.8		
(02 ⁰ 0)	9/2 11/2 5 /2 7/2	37417.1	34240.0		
	$3/2 5/2 \\ 1/2 3/2 $		(34236.0)		
	1/2 (1/2, 3/2)	37394.0	34228.8		
(02 ² 0)	3/2 (1/2,3/2,5/2) 9/2 (7/2,9/2,11/2)	37385.2	34220.1		
	5/2 (3/2,5/2,7/2)	37375.6	34210.4		
	7/2 (5/2,7/2,9/2)	37372.2	34 207 . 1		
	5/2 5/2 } 9/2 11/2 }	37362.2	34278.3		
(03 ¹ 0) <i>l</i> +	7/2 (7/2,9/2)	37358.9	34275.4		
	5/2 7/2 }	37357.2	34273.5		
	9/2 (7/2,9/2)	37352.5			

Table I

Table I - cont.

(03 ¹ 0) <i>l</i> -	5/2 5/2 9/2 11/2 7/2 (7/2,9/2) 5/2 7/2 3/2 5/2	37238.1 37235.0 37233.0	34152.2 34149.3 34147.8
(04 ² 0)	1/2 (1/2,3/2) 3/2 (1/2,3/2,5/2) 9/2 (7/2,9/2,11/2) 5/2 (3/2,5/2,7/2) 7/2 (5/2,7/2,9/2)	37179.4 37170.7 37160.9 37157.8	34189.2 34179.2 34174.8
(10 ⁰ 0)	7/2 (7/2,9/2) 9/2 11/2 3/2 5/2 1/2 3/2 9/2 7/2 9/2 9/2	37479.1 37472.7 37466.7	34095.4 34089.0 34082.9
(11 ¹ 0) <i>l</i> +	5/2 5/2 9/2 11/2 7/2 (7/2,9/2) 5/2 7/2 3/2 5/2 9/2 (7/2,9/2)	37333.6 37330.8 37329.0 37324.1	34071.0 34068.2 34066.4
(11 ¹ 0) <i>l</i> -	5/2 5/2 9/2 11/2 7/2 (7/2,9/2) 5/2 7/2 3/2 5/2 9/2 (7/2,9/2)	37269.2 37266.4 37264.9 37260.0	34010.2 34007.5 34006.1
(12 ⁰ 0)	7/2 (7/2,9/2) 9/2 11/2 5/2 7/2	37164.5	34010 .5 34007.6
(12 ² 0)	1/2 (1/2,3/2) 3/2 (1/2,3/2,5/2) 9/2 (7/2,9/2,11/2) 5/2 (3/2,5/2,7/2) 7/2 (5/2,7/2,9/2)	37140.7 37132.0 37122.3 37119.2	33995.2 33986.8 33977.3 33974.1
(200)	$7/2 (7/2,9/2) \\ 9/2 11/2 \\ 5/2 7/2 \\ 3/2 5/2 \\ 1/2 3/2 \\ 9/2 (7/2,9/2) \\ 65$	37216.9 37213.9 37210.8 37204.8	33866.8 33863.9 33860.7

Table II

Observed frequencies in $^{87}\mathrm{RbOH}$ and $^{87}\mathrm{RbOD}$

$J = 2 \rightarrow 3$					
State	$F \rightarrow F$	8 ⁷ кьон	⁸⁷ Rbod		
(vo ⁰ o)	7/2 9/2 5/2 7/2 3/2 5/2	37597.8	34188.4		
	1/2 3/2	37595.6	34186.6		
	3/2 3/2	37455.1			
1	$ \begin{array}{ccc} 7/2 & 9/2 \\ 1/2 & 3/2 \end{array} $	37450.8	34165.1		
(01 ¹ 0) <i>l</i> +	5/2 7/2 3/2 5/2	37448.7	34163.4		
	7/2 7/2	37444.0			
	7/2 9/2 1/2 3/2	37391.7	34106.9		
(01 ¹ 0) <i>L</i> -	5/2 7/2 3 3/2 5/2	37389.7	34105.1		
	7/2 9/2 5/2 7/2	37287.5	34106.9		
(02 ⁰ 0)	$\begin{array}{c} 3/2 & 7/2 \\ 3/2 & 5/2 \\ 1/2 & 3/2 \end{array}$	37285.6	34105.2		
	1/2 3/2 7/2 (5/2,7/2,9/2)	37246.8 37241.1	34082.5		
(02 ² 0)	3/2 (3/2,5/2) 5/2 (3/2,5/2,7/2)	37238.3 37232.8	34074.2		
	5/2 (5/2,5/2,1/2)	57252.0	J+U/+ , Z		
0	7/2 9/2 5/2 7/2	37337.1	33960.8		
(10 ⁰ 0)	5/2 7/2 3/2 5/2		33959.0		

Table III

Effective rotational constants and nuclear quadrupole coupling constants for various vibrational states of RbOH and RbOD.

State		Вv (⁸⁵ RbOH)	Bv (⁸⁵ RbOD)	eqQ
0000	Σ	6290.15	5720.77	-67.9
01 ¹ 0	Π	6260.55	5711.69	-67.6
02 ⁰ 0	Σ	6238.37	5707.08	(-67.9)
02 ² 0	\bigtriangleup	6230.13	5702.57	-67.4
03 ¹ 0	Π	6216.43	5702.34	(-67.6)
04 ² 0	\bigtriangleup	6194.38	5697.42	- 67
10 ⁰ 0	Σ	6246.45	5682.50	-65.9
11 ¹ 0	Π	6216.70	5673.24	-67
12 ⁰ 0	Σ	6194.03	5668.37	(-66)
1220	Δ_{-}	6187.95	5663.75	-66.3
2000	Σ	6202.75	5644.40	-64.2

State		Вv (⁸⁷ кьон)	Bv (⁸⁷ RbOD)	eqQ 🔇
0000	Σ	6266.35	5698.13	-35
01 ¹ 0	π	6236.84	5689.32	-33
02 ⁰ 0	Σ	6214.63	5684.53	-31
02 ² 0	\bigtriangleup	6206.48	5680.02	-32
10 ⁰ 0	Σ	6222.90	5660.18	(-33)

State	q (⁸⁵ RЬОН)	q (⁸⁵ RbOD)	q(⁸⁷ RbOH)	q (⁸⁷ RbOD)
(01 ¹ 0)	9.87	9.82	9.85	9.70
(03 ¹ 0)	10.33	10.51		
(11 ¹ 0)	10.73	10.12		

l-Doubling constants

Table V

	⁸⁵ _{RЬОН}		⁸⁵ _{RbOD}	
State	calc BOv ₂ ^l O	Obs - calc	calc BOv ₂ ^l O	Obs - calc
00 ⁰ 0	6290.10	0.05	5720.77	0.00
01 ¹ 0	6260.38	0.17	5711.70	-0.01
02 ⁰ 0	6238.57	-0,20	5707.09	-0.01
02 ² 0	6230.38	-0.25	5702.49	-0.08
03 ¹ 0	6216.48	-0.05	5702.34	0.00
04 ² 0	6194.10	0.28	5697.45	-0.03

Fit of $BOv_2^2 O$ to power series

Ta	Ь1	е	VI
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v ₂ ^ℓ	BOv ₂ ^l O	Blv ₂ ^l 0	
	⁸⁵ _{RЬОН}		
0 ⁰	47.70	43.70	
1 ¹ 2 ⁰	43.85 44.34		
2 ²	42.18		
	⁸⁵ _{RbOD}		
o ^o	38.27	38.10	
1 ¹ 2 ⁰	38.45 38.71		
2 ²	38.82		
	⁸⁷ кьон		
o ^O	43.45		
	⁸⁷ кьор		
0 ⁰	37.95		

Variation of rotational constants with V_1

Table VII

2.316	2.305
0.913	0.965

Structural parameters of $R_{\rm ubidium}\ {\rm Hydroxide}$

a) The first terms in equations (1) were used as ${}^{\prime\prime}B_{}{}^{\prime\prime}$.

Chapter 5

MASS SPECTROMETRIC STUDY AT HIGH TEMPERATURES OF THE Al₂ O₃ -BeO SYSTEM

by

J. Efimenko

INTRODUCTION

Data have been obtained for the vapor species existing at high temperatures for a number of selected compositions in the $Al_2 O_3$ -BeO system (Report No. 8628, 9028, 9389). This report presents data and calculated thermodynamic quantities for the eutectic mixture, BeO: $Al_2 O_3$ (94.2% $Al_2 O_3$), melting point 1890 ± 10°C, S. M. Lang, C. L. Fillmore, and L. H. Maxwell, J. Res. NBS <u>48</u>, 301 (1952). The initial purpose of these investigations was to observe and study any complex vapor species in this system. The following is a preliminary report of these data.

EXPERIMENTAL

The procedure followed is similar to that used previously, Report No. 9389. The mass spectrometer had been disassembled completely and reassembled in the new laboratory previous to this experiment. Before the eutectic mixture was placed in the instrument a blank run was carried out. It was necessary to check whether contamination from previously used materials was serious. In addition, a silver calibration was performed in the same effusion cell before the experiment with this eutectic mixture was concluded.

RESULTS

Experimental data are shown in Table 1 and auxiliary data used for computation in Table 2. Equilibria considered for this system are the following:

(1) $A1(g) + O(g) + Be(g) \neq A10Be(g)$

- (2) $2A1(g) + O(g) \neq Al_2 O(g)$
- (3) $Al_2 O(g) + Be(g) \neq A10Be(g) + A1(g)$
- (5) $Al_2 O_3 \cdot BeO(\ell) \neq 2A1(g) + Be(g) + 4O(g)$

Equilibria constants corresponding to these reactions were computed and are presented in Table 3. A second law treatment, (LnK vs 1/T) yielded enthalpies as shown in Table 4. For comparison, enthalpies for the same equilibria, observed over the compound, $Al_2 O_3 .BeO$ (80.3% $Al_2 O_3$), are presented. On the basis of the reaction enthalpy for (1) this experiment indicates a value of $\Delta H_{Of}^0 = -41$ kcal/mol for the standard heat of formation of AlOBe. The earlier experiments gave a value of -6.2 kcal/mol.

DISCUSSION

In the temperature range investigated, $2177-2562^{\circ}K$, the only complex specie which appeared in addition to the species from the individual reactants was AlOBe. At the higher end of the temperature range, tungsten oxides appeared also but their intensities were minor in comparison to the species Be⁺, O⁺, Al⁺.

During the period that the data in Table 1 were obtained, two shut-downs occurred. One was to replace the electron bombardment heating filaments that failed and the second was to introduce a weighed quantity of silver for a calibration. The data collected in any one run are indicated by the sequence in the index number and the separate runs by the tens series.

From Table 1 it is observed that there are 34 rows of data to compute enthalpies for reactions 1 and 5, while only eight rows for reactions 2 and 3. Since the specie of interest was AlOBe, more time was spent accumulating data for reactions 1 and 5. One set of data, index numbers 50-57, does not have a temperature cycle.

The computed equilibrium constants for reaction 1 show a rather large scatter for check temperature points. Two factors can be responsible for such a scatter. The effusion cell may have had a serious temperature gradient, before or after changing the bombardment filaments and secondly, temperature equilibrium may not have been established inside the effusion cell. Whenever a temperature cycle was performed during a run, the down temperature data do not always follow the same line as the up temperature data. Attainment of constancy of ion intensity may not be a true indication of thermal equilibrium after intentionally changing temperature. The equilibrium constants for reactions 2 and 3 show much less scatter because no temperature cycle was performed for the data, index numbers 50-57.

Since no composition analysis was made of the condensed phase with time and temperature, reaction 5 will not be discussed here. Using the eutectic composition $Be0:Al_2O_3$ (94.2% Al_2O_3) gave a starting composition with specific properties at equilibrium. Reaction 3, which is not independent but is obtainable from combination of reactions 1 and 2, is included as a check. These enthalpies need to be re-examined because of the indicated discrepancies. Reaction 2 has been studied independently by other workers, reporting values ΔH_0^0 , -254 ± 7 and -243 ± 7 kcal/mol. A third law computation will be made for the enthalpies of these reactions for analysis and comparison. As pointed out previously, the data points for reactions 2 and 3 were taken without a temperature cycle and include much fewer readings. Data were obtained on two other eutectic compositions, BeO:Al₂O₃ (85.9% Al₂O₃) and BeO:Al₂O₃ (75% Al₂O₃), as well as another compound, BeO.Al₂O₃ (80.3% Al₂O₃). This additional data will be analyzed for the same gaseous equilibria.

Ion Intensities Over Eutectic BeO:Al₂ O₃ (94.2% Al₂ O₃)

(intensities in volts)

$I_{Al_2}^+$ 0	28 × 10 ⁻² 14 × 10 ⁻² 34 × 10 ⁻³ 14 × 10 ⁻³ 32 × 10 ⁻³ 12 × 10 ⁻⁴ 10 × 10 ⁻⁴
I ⁺ A10	28×10^{-2} 13.5×10^{-3} 67.5×10^{-3} 33×10^{-3} 14×10^{-4} 9×10^{-4} 9×10^{-4}
I ⁺ AlOBe	$\begin{array}{cccccccccccccccccccccccccccccccccccc$
r ⁺ Al	$\begin{array}{c} 4.2 \times 10^{-1} \\ 2.2 \times 10^{-1} \\ 1.2 \times 10^{-1} \\ 60.5 \times 10^{-3} \\ 77 \times 10^{-3} \\ 77 \times 10^{-3} \\ 36.5 \times 10^{-3} \\ 39 \times 10^{-3} \\ 61 \times 10^{-3} \\ 27 \times 10^{-3} \\ 10 \times 10^{-3} \\ 27 \times 10^{-3} \\ 28 \times 10^{-3} \\ 28 \times 10^{-3} \\ 36 \times 10^{-3} \\ 36 \times 10^{-3} \\ 36 \times 10^{-3} \\ 24 \times 10$
I ⁺ 0	$\begin{array}{cccccccccccccccccccccccccccccccccccc$
I ^t Be	$\begin{array}{cccccccccccccccccccccccccccccccccccc$
T, °C	2190 2135 2085 2085 2030 1985 1855 1850 1890 1890 1980 1920 1880 1928 1920 1880 1880 1880 1880 1880 1880 1880 18
Index No.	57 57 57 57 57 57 57 57 57 57 57 57 57 5

TABLE 1 (continued)

×	×	28×10^{-3}	×	×	×	×	×	×
4	×	55×10^{-1}	×	×	×	×	×	×
×	×	67×10^{-2}	×	×	×	×	×	×
×	×	10×10^{-2}	×	×	×	×	×	×
1935	2035	2200	2150	2080	2040	1975	2025	1920
48	47	46	45	44	43	42	41	40

0 0

76

Auxiliary Data

Sensitivity, S _{Ag}	1.7796 × 10 ⁻⁸ amp/mmHg
Temp Silver Calib.	1305K

Window + Prism Transmissivity $1.40 \times 10^{-5} \text{K}^{-1}$

Specie	Ionization Cross Section σ_i (1)	Multipli er Sensitivity γ ₁ (2)
Ag	34.8	296
Ве	6.3	276
0	3.3	288
A1	15.4	292
Al ₂ 0	34.1	226
AlÕBe	25.0	93

- (1) J. W. Otvos and D. P. Stevenson, J. Am. Chem. Soc. <u>78</u>, 546-551 (1956).
- (2) Experimental values, author.

Equilibria Constants

 K_i subscripts refer to the reactions listed below

Index <u>No.</u>	1/T, °K	K1	K2	Kg	K ₅ × 10 ²⁰
57	3.9201	1.2730	3.6402	3.4971	5.7040 × 10 ⁻³⁰
56	4.0128	4.0242	1.2165	3.3082	9.5881 × 10 ⁻³²
55	4.1009	1.1166	3.9963	2.7941	1.5154 × 10 ⁻³³
54	4.2022	3.7407	1.4808	2.5261	1.5368 × 10 ⁻³⁵
53	4.2985	1.5657	6.6793	2.3441	7.3905 × 10 ⁻³⁸
52	4.4725	1.3175	6.0730	2.1694	2.4400×10^{-41}
51	4.5592	7.9974	2.4020	3.3294	1.1349×10^{-43}
50	4.5703	1.1798	1.6978	6.9490	1.0738×10^{-43}
10	4.5482	1.3564			4.1156×10^{-43}
11	4.5926	3.0777			1.5066×10^{-44}
12	4.4789	4.2548			1.8209×10^{-42}
13	4.4832	3.2166			2.9582×10^{-42}
14	4.2595	1.1251			4.9912×10^{-38}
15	4.1648	2.2541			8.5291×10^{-36}
16	4.2887	1.2952			1.8948×10^{-38}
17	4.3993	6.5663			9.4411 \times 10 ⁻⁴¹
18	4.5047	4.4072			3.5974 × 10 ⁻⁴³ 1.4185 × 10 ⁻⁴⁴
20	4.5482	1.1973			1.4185×10^{-41} 1.1580×10^{-41}
21	4.4200	7.9299			1.6163×10^{-43}
22	4.5047	4.4311			9.7115×10^{-45}
23	4.5703	9.0211			1.7242×10^{-34}
30 31	4.1555	3.8737 1.9521			6.4258×10^{-39}
32	4.2789 4.4034	1.8653			4.0287×10^{-40}
33	4.4034	9.1351			3.3718 × 10 ⁻⁴²
48	4.3890	1.3427			6.5356×10^{-40}
47	4.1928	8.5192			1.0185×10^{-35}
46	3.9037	1.0857			3.5567 × 10 ⁻²⁹
45	3.9871	3.7652			3.3823×10^{-33}
44	4.1099	1.6873			1.6897 × 10 ⁻³⁵
43	4.1834	6.7253			1.4244 × 10 ⁻³⁵
42	4.3084	4.2802			2.6880×10^{-38}
41	4.2116	1.1728			2.6290 × 10 ⁻³⁶
40	4.4200	1.8497			1.1650 × 10 ⁻³⁸
	1. A1 + 0 + 2. 2A1 + 0 =	Be = A10Be, Al ₂ 0, K ₂	K ₁		

- 2. $2A1 + 0 = Al_2 0$, K_2 3. $Al_2 0 + Be = A10Be + A1$, K_3 5. $Al_2 0_3 \cdot Be0(\ell) = Be + 2A1 + 40$, K_5

Reaction Enthalpies

Results from eutectic BeO:Al₂ O₃ (94.2 wgt % Al₂ O₃, melting point 1890 \pm 10°C)

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Results* from compound BeO.Al₂ O₃ (80.3 wgt % Al₂ O₃, melting point 1870 ± 10°C)

Reaction	ΔH_{2350}^{o} , kcal/mol	ΔH_0^o , kcal/mol
(1) (2)	-222 ± 4.6 -257 ± 6	-216 -252
(3) (5)	+35.5 ± 4 +952 ± 71	+36.2

* From Report No. 9028.

Chapter 6

THE ENTHALPY OF TUNGSTEN FROM 1100 TO 2600°K

E. D. West and S. Ishihara

The high-temperature enthalpy of tungsten is important not only because of its use in high-temperature engineering applications but also because of its use as a container material in the high-temperature enthalpy measurements of other materials and its possible use as a reference material for these measurements to permit a check on the capabilities of various laboratories.

In earlier work in this program [1,2], tungsten was used as a container for liquid $A\ell_2O_3$ and liquid BeO·A ℓ_2O_3 . Revised results for the latter are reported in a subsequent chapter in this report.

The work reported in this chapter was carried out as part of a National Bureau of Standards program related to work done for AFOSR.

Method and Apparatus

The method and apparatus have been described briefly [1,2,3]. Essentially, the apparatus consists of a furnace, in which a specimen in a capsule is held at various high temperatures, and an adiabatic calorimeter, operating near room temperature, which measures the heat given up when the hot specimen is lifted into it. Heat lost by the container in the transfer process is accounted for by separate experiments with the empty capsule. If all the heat lost comes from the capsule wall in both experiments, it is accounted for when the difference in heats is ascribed to the specimen.

The temperature in the furnace is measured by a Leeds & Northrup automatic photoelectric pyrometer sighted on the bottom of the capsule. The output signal from the pyrometer is used in a control loop to maintain a selected constant temperature while the capsule is in the furnace. Some improvements have been made in the pyrometry since the earlier report [1]. Frequent balancing of the integrator circuit has apparently improved the precision of the temperature measurement. Reported temperatures are based on a calibration made by the NBS Radiation Thermometry Section in 1967. The conditions of this calibration are closer to the conditions of use than those of the 1965 calibration although the maximum difference between the two is 0.2% at 1108°K. Four spot checks between these two calibrations do not show any consistent trend. They differ from the 1967 calibration by -0.16, -0.95, +0.31 and +0.66°K. At high temperatures, there is considerable risk of contamination or mechanical damage to both specimen and container. With tungsten we found it helpful to use a smaller specimen for higher temperatures to reduce breakage of the suspension wire. To minimize the risk and permit this flexibility, we use a method which yields in one day's work all the information required for the calculation of the enthalpy at the operating temperature for that day. Usually four experiments are made at the same temperature in a day--two with the capsule alone and two with the capsule plus specimen. For some higher temperatures, only two experiments are made because of the time required for additional weighing. Weights of the capsule and the specimen can be made after each experiment to check for contamination or vaporization. Allowance for variations in the heat for the capsule can be made by correlating weight gains and observed heats for the empty capsule.

A principal advantage of this method is that loss of a capsule or container on one day does not invalidate data taken previously as would be the case if all the data were obtained for the empty capsule before any measurements were made with the specimen. In this case, loss of the capsule might mean loss of the data taken with it. In experiments at 2700°K, a tungsten capsule developed a crack which allowed contamination of specimen D with carbon from the furnace atmosphere, but experiments at lower temperatures were not affected.

Another advantage of the method is that it facilitates intercomparison of different specimens under nearly identical conditions. At 1613°K, for example, two different specimens were used the same day.

Materials

Four specimens were used in this work. These are described below with the same designations used in table 1. All machining was done by grinding with aluminum oxide wheels.

Specimen A: Commercial "pure", nominally 99.95% tungsten. Spectrochemical analysis of another piece of the same rod for 51 elements found a maximum of 0.01% impurity. Surface was cleaned by rinsing in carbon tetrachloride. Weight (corrected for buoyancy) 23.0134 g, constant to 0.2 mg, which is about the limit of the balance used.

<u>Specimen B</u>: Triple zone-refined tungsten nominally 99.999% tungsten, in the form of one-eighth inch diameter rods. The rods were cut to length, but not otherwise machined. The surface was treated with one part 48% HF and two parts concentrated HNO₃ for one minute. The weight was 13.3911 g constant ± within 0.2 mg. Specimen C: Same as A but surface treated with one part HF and three parts HNO_3 . Weight 13.8482 g constant within 0.2 mg until it gained 0.6 mg in the experiment at 2403°K.

Specimen D: Same as C, except weight 10.6807 g. The specimen gained 0.4 mg at 2450°K, 0.4 and 2.9 mg at 2600, 15.5 mg at 2700°K and 20.6 mg at 2800°K. Analysis in a carbon-hydrogen combustion train found 0.74% carbon in a sample 1 mm long cut from the ends of the specimen. A second sample 2 mm long adjacent to the first showed 0.12% carbon. The analysis indicates that carbon vapor reached the specimen through the crack in the capsule wall near one end.

The atomic weight used for tungsten was 183.85.

Results

The corrected experimental data for the enthalpy of tungsten are shown in table 1. Three capsules and four specimens were used, as indicated in the last column. The specimens are described above.

The temperature in the first column has been corrected for the loss of radiation in a prism in the optical path. Corrections made for variation in room temperature in earlier work [1] are no longer necessary.

The quantity of heat in the second column includes corrections for the enthalpy of the capsule and specimen between the actual final calorimeter temperature and 298.15 °K. Data required for this correction are taken from Stull and Sinke [4]. Corrections made in the earlier work [1] for small variations in the furnace temperature are no longer required. The enthalpy calculated from one experiment with the capsule and one with the capsule and specimen are shown in the third column.

The data for the enthalpy of tungsten in kilojoules per gramatomic weight are represented by the equation

$$H_{T} - H_{298.15} = -7.115 + (23.041)10^{-3}T + (2.9836)10^{-0}T^{2} - (.82203)10^{-9}T^{3} + (.21967)10^{-12}T^{4} kJ/gaw$$
(1)

The standard deviation is 0.17%. Deviations are shown in the fourth column of table 1.

The table contains data for very pure tungsten (sample B) at four temperatures. The percentage deviations are -0.03, -0.20, +0.09, -0.23, and -0.12. We regard these differences as insignificant and the data for sample B have been used in obtaining equation (1). One point at 2353°K was omitted in determining equation (1) because it differs by more than three times the standard deviation from the corresponding value from equation (1), although the experiment does not have any apparent defect.

Equation (1) was obtained by the method of least squares, minimizing the sum of the squares of the percentage deviations. This procedure is preferable to working with the deviations themselves because the uncertainty in temperature measurements increases with the temperature. Such a procedure would tend to give too much weight to the experiments at high temperatures.

Discussion

Kirillin, et al. [5] summarize the measurements made before 1963, including their own. They represent the data of several observers with two smoothing equations, one for the range from 250 to 385° K and one from 385 to 3100° K. The maximum difference between their equation and equation (1) is 0.59% at 2050° K. For their data for the range 1275 to 2600° K, the standard deviation is 0.46%. From the deviation plot given [5], it appears that the data in this range are more precise than the standard deviation indicates. For this reason, we have obtained equation (2) for the enthalpy in kilojoules/gram-atomic weight which fits their data for the range 630 to 3093° K with a standard deviation of 0.33%:

$$H_{T}-H_{298.15} = -3.854 + (25.848)10^{-3}T + (0.72405)10^{-6}T^{2} + (.30599)10^{-9}T^{3} - .68886 \ln T kJ/gaw$$
(2)

The difference between equations (1) and (2) is +0.15% at 1100° K, decreasing to -.22% at 1900°K, then increasing to +0.29% at 2500 and 0.51% at 2600°K.

The agreement between the two equations is quite good except at the highest temperature. That agreement is better between equations (1) and (2) than between equation (1) and the equation of Kirillin, et al. [5] indicated that the latter may have given too much weight to older data in the temperature range below 1100°K.

Since tungsten is used as a container for liquid oxides, the data for the enthalpy of tungsten affect the liquid data obtained for $A\ell_2O_3$ and BeO·A ℓ_2O_3 previously reported [1,2]. Revisions of those data are presented in subsequent chapters.

References

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- [2] S. Ishihara and E. D. West, NBS Report 9601 to AFOSR, 1 July 1967, p. 55.
- [3] E. D. West and S. Ishihara in "Advances in Thermophysical Properties at High Temperatures and Pressures," Am. Soc. Mech. Engr. 1965, p. 146.
- [4] D. R. Stull and G. C. Sinke, Thermodynamic Properties of the Elements, Am. Chem. Soc., Washington, D.C. (1956).
- [5] V. A. Kirillin, A. E. Sheindlin, V. Ya. Chekhovskoi, and
 V. A. Petrov, J. Phys. Chem. USSR <u>37</u>, 1212, 1963,
 translated from Zh. Fiz Khim 37, 2249 (1963).
- [6] V. Ya. Chekhovskoi, B. Ya. Shumyatskii, and K. A. Yakimovich, Inzh. Fiz. Zh. Akad. Nauk Belorussk SSR 5, 13 (1962).

Table 1. Experimental Data for the Enthalpy of Tungsten

Furn. temp.°K Date of run	Heat to Calorim- eter at 298.15°K Joules	HT-H298.15 kJoules/at.wt.	% Deviations Obs-Calc. from eq (1)	Specimen and Capsule Material
1108.8 3 -1 6-66	4599.4 1933.3 1933.3	21.299	•02	A, Mo
	4599.6	21.301	.03	A, Mo
1183.8 2-23-66	2129.6 5056.6 5055.9 2125.2	23.383 23.413	11 .02	A, Mo A, Mo
1183.8 2 -1 6-66	2121.4 5050.9 5052.0 2119.7	23.403 23.426	03 .07	A, Mo A, Mo
1333.6 2-14-66	2508.3 5967.8 5970.0 2510.6	27.638 27.637	09 09	A, Mo A, Mo
1405.9 3-7-66	6427.7 2705.1 2706.7 4872.9	29.739 29.741	03 03	А, Мо В, Мо
1514.3 2 -11- 66	3001.6 7136.7 7142.9 3008.8	33.035 33.027	•35 •33	А, Мо А, Мо
1613.2 3 -18- 66	7758.5 3283.9 3285.9 5892.3	35•747 35•784	30 20	A, Mo B, Mo
1726.8 2-9-66	3605.4 8529.1 8531.2 3607.2	39•335 39•337	.12 .12	А, Мо А, Мо
1815.7 3 -11- 66	9127.9 3860.3 3866.2 9109.9	42.082 41.891	•13 -•32	А, Мо А, Мо
				,

Table 1 (Cont'd)

Furn. temp.°K Date of run	Heat to Calorim- eter at 298.15°K Joules	HT-H298.15 kJoules/at.wt.	% Deviations Obs-Calc. from eq(1)	Specimen and Capsule Material
1915.4 3-14-66	4166.9 9812.0 9827.3 4163.8	45.097 45.245	10 .23	А, Мо А, Мо
1998.1 3-21-66	4400.6 7883.8 7893.2 442 1. 4	47.821 47.665	.09 23	B, W(1) B, W(1)
2107.4 3-23-66	86 15.3 4753.7 4764.4	51.268	12	B, W(l)
	8502.1	51.317	03	C, W(l)
2107.5 4 -18- 66	8874.4 5021.0 5023.9	51.155	35	C, W(l)
	8891.8	51.347	02	C, W(l)
2210.0 3-31-66	5059.4 9186.0 9215.1 5095.0	54.785 54.700	.08 08	C, W(l) C, W(l)
2209.1 4-4-66 (Slow lift time)	5078.5 9215.6 9232.3 5102.6	54.924 54.825	•38 •20	C, W(1) C, W (1)
2267.7 4-14-66	3473.0 6770.5	56.761	.11	D, W(2)
2300.8 4 - 6-66	9803.9 5451.5	57.783	09	C, W(l)
2403.3 4-8-66	5944.5 10569.2	61.394	02	C. W(l)

Table 1 (Cont'd)

	eat to Calorim- ter at 298.15°K	HT-H298.15 kJoules/at.wt.	%Deviations Obs-Calc. from eq(1)	Specimen and Capsule Material
	Joules		11 01 04 (1)	10001 101
2460.8 4 -21- 66	4101.4 7799.1	63.649	•30	D, W(2)
2503.1 4-12-66	7741.8 3984.8	64.671	48	D, W(2)
2604.4 4-25-66	5167.1 9171.6 8896.6 4912.7	68.931 68.577	.31 20	D, W(2) D, W(2)
2353 .11[*] 4 -15- 66	7126.2 3685.1	59.232	70	D, W(2)

^a Atomic wt. of tungsten 183.85.

* This point was given zero weight for determining the smoothed curve.

Chapter 7

THE ENTHALPY OF SOLID AND LIQUID BeO·A&203 FROM 1200° TO 2400°K - REVISION

S. Ishihara and E. D. West

The data presented for the enthalpy of solid and liquid $Be0 \cdot Al_2O_3$

¹ Ishihara and West, NBS Report 9601 to the AFOSR, 1 July 1967.

have been revised slightly as a result of a calibration of the automatic pyrometer carried out following completion of the measurements and also as a result of new data for the enthalpy of tungsten, which affects only the data for the liquid. The revisions affect the enthalpy relative to 298.15°K by less than 0.1%. They are therefore probably not important in engineering calculations, but may be useful in checks of consistency of thermodynamic data.

The enthalpy of solid $Be0 \cdot A\ell_2O_3$ from 1180 to 2143°K relative to 298.15°K is given by the equation:

$$H_{T} - H_{298.15} = -(0.255187)T + (3.46055)10^{-4}T^{2} - (1.44123)10^{-7}T^{3} + (2.43189)10^{-11}T^{4} + 72.2028 \ln T - 362.433 kJ/M$$
(1)

The enthalpy calculated from equation (1) will be approximately 0.05% lower than that calculated from the earlier equation.^{*} With the increased precision from the new calibration, an additional constant is warranted (F test at 95% confidence level).² The constants in

² <u>M. G. Natrella, Experimental Statistics NBS Handbook 91</u> (U.S. Dept. of Commerce, 1963).

equation (1) are therefore quite different from those in the earlier equation, but the difference in enthalpy calculated is small. Equation (1) fits the data with a standard deviation for an individual measurement of 0.16%, which is very slightly smaller than that given earlier.

^{*} In the earlier report a minus sign was omitted for the logarithmic term.

The enthalpy of liquid $Be0 \cdot A\ell_2 O_3$ has been corrected for new data for tungsten, which was used as a container material, and also for the new pyrometer calibration. The enthalpy for the range 2143 to 2400°K is given by the equation:

$$H_{\rm T} - H_{298.15} = 1299 - (0.9376)T + (0.2637)10^{-3}T^{2} kJ/M$$
 (2)

The enthalpy calculated from (2) will differ by less than 0.1% from the enthalpy calculated by the earlier equation. The root-mean-square deviation from equation (2) for a single determination consisting of three experiments is 0.03%.

The heat of fusion at 2143°K calculated from the difference between equations (1) and (2) is 172.3 kJ/mole, lower by 1.5 kJ/mole than the value reported earlier. This value depends slightly on the temperature taken for the melting point and increases 0.04 kJ/mole for each degree decrease in the melting point.

Chapter 8

THE STATUS OF THE THERMOCHEMICAL DATA ON SOME Mg COMPOUNDS by Vivian B. Parker

I Introduction

As part of the revision of NBS Circular 500 "Selected Values of Chemical Thermodynamic Properties", we have completed a review of the available data on all magnesium compounds. We present here a summary of values selected from this review for substances pertinent to the Light Element Thermodynamics Program. We are including also a brief discussion of the sources from which these values have been calculated. It is hoped that this report will indicate where additional research is required. We will appreciate it if data that have been overlooked or errors in the values are brought to our attention before the final tables are published.

All auxiliary data and constants used in the calculations are given in NBS Technical Note 270-3 [1]. All values of \triangle Hf° and \triangle Gf° are in kcal/mol for the molecular formula written and at 298°K, unless otherwise specified. Caution should be exercised if these values must be combined with data taken from other compilations or sources, in order to avoid errors caused by a lack of consistency between the tables.

II Discussion of Data

<u>Mg(c)</u> This is taken as the standard state. The thermal functions were taken from the evaluation of Hultgren et al. [2,3].

<u>Mg(g)</u> The $\triangle H^{\circ}(\text{subl}) = 35.3$ kcal has been taken from the tabulations of Hultgren et al. [2,3], as have the thermal functions of the ideal monatomic gas. Hultgren et al. [3] revised their original selection of 35.3 to 35.0 kcal. We prefer 35.3.

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 $Mg_{-}^{n+}(g)$ The successive ionization energies were taken from Moore [4].

 $\frac{Mg^{2+}}{(aq, std. state, m = 1)}$ See MgCl₂(aq), Mg(ClO₄)₂(aq, std. state, m = 1) and Mg(NO₃)₂(aq).

Let us first consider \triangle Hf° from the aqueous MgCl₂ system. The values given in Table II were corrected to \triangle Hf°. Representative values from the MgCl₂·6H₂O system are: Samoilov -191.17 kcal; Voskresenskaya and Ponomareva -191.10; Thomsen -191.42; Mishchenko and Yakovlev -191.17. From the MgCl₂·4H₂O(c) and MgCl₂·2H₂O(c) systems we obtain from Thomsen's measurements -191.77 and -191.20. From MgCl₂(c) we obtain: Mishchenko and Yakovlev -190.92; Mishchenko et al. -191.17; Thomsen -190.28. From Richards et al. we obtain -189.56. A tentative "best" value for \triangle Hf° MgCl₂ is -191.22 kcal with \triangle Hf° Mg²⁺(aq, std. state, m = 1) = -111.32 kcal.

For Mg(ClO₄)₂ \triangle Hf° is -173.63 kcal from which we obtain \triangle Hf° Mg²⁺ = -111.81 kcal.

From the Mg(NO₃) system, we obtain for \triangle Hf°: Pickering -210.67; Thomsen -211.00; Ewing et al. -210.57 and -210.72. These result in a \triangle Hf° = -210.74 and \triangle Hf° Mg²⁺ = -111.62 kcal.

We have selected as "best" value for $\triangle Hf^{\circ} Mg^{2+}$ -111.58 kcal.

From the solubilities given by Linke [5] and the activity coefficients and osmotic coefficients given by Robinson and Stokes [6] we obtain ΔG° soln MgCl₂·6H₂O(c) = -6.09 kcal and ΔG° soln MgSO₄·6H₂O(c) (metastable at 25°C) = +2.18 kcal. Combined with the appropriate heat of solution data we obtain ΔS° soln = +6.70 e.u. and -10.64 e.u., respectively. These result in S° Mg²⁺ = -33.1 and -32.5 e.u. We have selected S° = -33.0, with ΔGf° Mg²⁺ = -108.7 kcal.

MgO(macrocrystal) and MgO(microcrystal)

The thermal functions for the macrocrystal were taken from the tabulation of the Dow Chemical Co. [7]; those for the microcrystal from the evaluation of Furukawa et al. [8].

There are many determinations for $\triangle Hf^{\circ} MgO(c)$; however, very few of them distinguish between the macro and micro crystal forms. The following values were reported without any indication which form was obtained or the percent of each obtained in the direct combustion of Mg(c); Muthman and Weiss [9], -143.0 kcal; Moose and Paar [10] -146.1; von Wartenberg [11] -143.9; Rogers [12] -146.4; Vorob'ev and Skuratov [13] -143.9; Shchukarev et al. [14] -143.23; Holley and Huber [15] -143.65 kcal.

Shomate and Huffman [16] measured the heats of solution of Mg(c) and MgO in aqueous HCl, from which we obtain Mg(c) + $H_2O(\text{liq}) \rightarrow \text{MgO(c)}$ + $H_2(g)$, $\Delta H^\circ = -75.494$ kcal and $\Delta Hf^\circ = -143.81$. No mention is made of the crystal size of MgO; however, the MgO used was made from igniting Mg(OH)₂(c) at high temperature; the macro crystal form is usually obtained by this method. We are therefore adopting this value as the ΔHf° of the macro form. It is the only definite one for all macro MgO(c).

Giauque [17] showed that finely divided MgO prepared from the reduction of Mg(OH)₂ at low temperature gives a different S° and Δ H soln in HCl. He reported Δ H° = 0.89 kcal for MgO(macro) \rightarrow MgO(micro). This results in Δ Hf° = -142.92 kcal.

<u>MgO(g)</u> Brewer and Porter [18] from their vapor pressure measurements have shown that the solid vaporizes mainly into molecular species. They gave spectroscopic evidence that the ${}^{1}\Sigma$ electronic state is not the principal vaporizing species. From their measurements and a ${}^{3}\Sigma$ ground state we obtain a 3rd law $\triangle H^{\circ}$ subl = 129 kcal, and a $\triangle Hf^{\circ}$ = -15 kcal. From Alexander et al.'s [19] H₂O(g) transpiration study we obtain a 2nd law $\triangle Hf^{\circ}$ = -4 and a 3rd law $\triangle Hf^{\circ}$ = +5 kcal. Altman's [20] Knudsen effusion and O₂ transpiration studies indicate that MgO(g) is of little importance at 2000°K (in contrast to Brewer and Porter) but from the experiments in which O₂(g) is passed over MgO(c) the 3rd law $\triangle H^{\circ}$ subl = 162 kcal and $\triangle Hf^{\circ}$ = +18 kcal. From Porter et al. [21] we obtain $\triangle H^{\circ}$ subl > 148.6 and $\triangle Hf^{\circ}$ > 4.8 kcal.

Bulewicz and Sugden [22] and Veits et al. [23] reported $D_0^{\circ} = 98$ and 100 kcal for a ${}^{3}\Sigma$ and ${}^{1}\Sigma$ ground state, respectively. These result in $\triangle Hf^{\circ} = -4$ kcal and -2 kcal (corrected to a ${}^{3}\Sigma$ ground state).

Drowart et al. [24] used the Knudsen effusion-mass spectrometer technique to determine relative ion intensities and enthalpies for the reactions $\log O(g) + O(g) \rightarrow Mg(g) + O_2(g)$, and $MgO(g) + WO_2(g) \rightarrow Mg(g) + WO_3(g)$. We obtain, corrected for a $3 \ge$ ground state, $\Delta Hf^\circ = +12$ and +20 kcal from the two reactions.

As is evident, the values are very discordant. We are taking $\triangle Hf^{\circ} = +4$ kcal. The uncertainty is large, ± 20 kcal.

Brewer and Trajmor [25] feel that for the purposes of high temperature calculations, the most reliable calculations are obtained by the assumption of a ${}^{3}\pi$ ground state for MgO(g). This would increase \triangle Hf° approximately 3 kcal.

The thermal functions used in the calculations were obtained from the Dow Chemical Co. [7].

MgH₂(c) The thermal functions were obtained from the Dow Chemical Co. [26] which also cites the references and values obtained for \triangle Hf°. We have taken \triangle Hf° = -18.0 kcal.

In addition Pepekin et al. [27] reported $\triangle H^\circ = -190.4$ kcal for the oxidation of MgH₂ to MgO(c) + H₂O(liq). This results in $\triangle Hf^\circ = -21.7$ kcal.

<u>MgOH</u>⁺(aq, std. state, m = 1) From Sillen and Martell [28] we obtain logK = 2.58 for Mg²⁺(aq) + OH⁻ → MgOH⁺. Hostetler's [29] value is in agreement.

 $Mg(OH)_2(c)$ The thermal functions were obtained from the Dow Chemical Co. [30].

Wells and Taylor [31] measured the heats of solution of $Mg(OH)_2(c)$ and MgO(macro) in 2N HC1. We obtain for $MgO(c) + H_2O(1iq) \rightarrow Mg(OH)_2(c)$, $\Delta H^\circ = -9.02$ kcal and $\Delta Hf^\circ = -221.15$ kcal. Similarly, from Taylor and Wells' [32] measurements, we obtain $\Delta H^\circ = -9.07$, using macro MgO and brucite ($\Delta Hf^\circ = -221.20$) and -8.83 kcal using macro MgO and synthetic $Mg(OH)_2$ ($\Delta Hf^\circ = -220.96$ kcal). From Giauque and Archibald [33] we obtain $\Delta H^\circ = -9.735$ kcal, using micro MgO, and $\Delta Hf^\circ = -220.97$. Torgeson and Sahama [34] measured $\Delta H = -26.95$ kcal for the solution of synthetic Mg(OH)_2 in 1N HC1. Using the results of Shomate and Huffman [16] on the solution of macro MgO, we obtain $\Delta H^{\circ} = -8.84$ kcal for the formation of Mg(OH)₂ from macro MgO and H₂O(liq) and $\Delta Hf^{\circ} =$ -220.97 kcal. From Roth and Chall's [35] measurements we obtain $\Delta H^{\circ} =$ -7.96 kcal and $\Delta Hf^{\circ} = -220.1$ kcal (this is assuming macro MgO(c) is used). We have selected a "best" value of -220.97 kcal.

Giauque and Archibald's decomposition pressure measurements are in agreement with this value. The Dow Chemical Co. [30] cites other decomposition vapor pressure measurements.

 $Mg(OH)_2$ (aq,std. state, m = 1) The values were obtained from the ions.

 $M_g(OH)_2(g)$ Alexander et al. [19] determined the Kp's (from 1780°K to 2010°K by the transpiration method) of the reaction MgO(c) + $H_2O(g) \rightarrow M_g(OH)_2(g)$. $\triangle H_{1900°K} = 66.8$ kcal from which we obtain $\triangle H^\circ = 67.6$ kcal and $\triangle Hf^\circ = -134$ kcal.

The values of H - H_{298} used in correcting $\triangle H$ to 298°K were taken from the estimated functions of the Dow Chemical Co. [30].

<u>MgF(g)</u> Ehlert et al. [38] reported the K's of the reactions, $AlF_3(g) + 2Mg(g) \rightarrow AlF(g) + 2MgF(g)$ (1238°K to 1301°K), and $2MgF(g) \rightarrow$ $Mg(g) + MgF_2(c)$ (1205°K to 1242°K). Third law values for ΔH° are 47.5 kcal and -126.65 kcal, which lead to -54.1 and -53.3 kcal for ΔHf° . An additional reaction, studied in Ehlert's Dissertation,

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is $2MgF(g) \rightarrow Mg(g) + MgF_2(g)$ for which the Dow Chemical Co. [30] reported $M^\circ = -34.76$ kcal. This results in $\Delta Hf^\circ = -51.5$ kcal.

Gaydon [39] and Herzberg [40] report $D_0^{\circ} = 3.2 \text{ ev}$ and 4.2 ev, respectively or $\triangle \text{Hf}^{\circ} = -21$ and -41 kcal.

We have taken $\triangle Hf^{\circ} = -53.0$ kcal.

The thermal functions were taken from the Dow Chemical Co. [30].

 $MgF_2(c)$ The thermal functions were taken from the Dow Chemical Co. [41].

Rudzitis et al. [42], from fluorine bomb calorimetry, reported \triangle Hf° = -268.7 kcal. Gross et al. [43] reported \triangle H° = -109.5 kcal for Mg(c) + PbF₂(c) \rightarrow MgF₂(c) + Pb(c) which results in \triangle Hf° = -268.2 kcal. From Kapustinskii and Samplavskaya [44] we obtain \triangle H = -94.8 kcal for Mg₂SiO₄(c) + 725.2(HF + 4.442 H₂O) \rightarrow 2MgF₂(c) + H₂SiF₆ + 4H₂O + 715.2(HF + 4.510 H₂O) and \triangle Hf° = -266.3 kcal.

From Guntz' [45,46] reactions we obtain $\triangle H = 2.8$ kcal for MgSO₄ (440 H₂O) + 2KF(220 H₂O) \rightarrow MgF₂(c) + K₂SO₄(880 H₂O) and $\triangle H = -30.2$ kcal for MgO + 2HF(110 H₂O) \rightarrow MgF₂(c) + H₂O. These result in $\triangle Hf^{\circ} = -266.7$ and -257.5 kcal, respectively. von Wartenberg [47] reported $\triangle H = -109.6$ kcal for the solution of Mg in aqueous HF, from which we calculate $\triangle Hf^{\circ} = -262.0$ kcal. Petersen's [48] $\triangle H$ for MgCl₂(440 H₂O) + 2AgF(220 H₂O) \rightarrow 2AgCl(c) + MgF₂(c) results in $\triangle H = -28.1$ kcal and $\triangle Hf^{\circ} = -265.2$ kcal. Torgeson and Sahama's [34] reaction of Mg(OH)₂ (c, 25°C) with HF(4.44 H₂O) at 73.7°C results in $\triangle H = -30.9$ kcal and $\triangle Hf^{\circ} = -267.5$ kcal. We have taken $\triangle Hf^{\circ} = -268.5$ kcal.

 $MgF_2(g)$ The thermal functions used were obtained from the Dow Chemical Co. [30].

There have been many measurements of the vapor pressure over the crystal and liquid. Table I summarizes the 2nd and 3rd law ΔH° subl obtained. The "best" value is 95.5 kcal and $\Delta Hf^{\circ} = -173.0$ kcal.

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			∆H° s	∆H° sub l	
Investigator	Temperature °K	Method	2nd Law	3rd Law	
Berkowitz and Marquart [49]	1450	Torsion effusion	91.3	94.68	
Green et al. [50]	1241 - 1492	Knudsen effusion	93.8	96.1	
Hammer and Pask [51]	1451 - 1533(c)	11 11	98.8	96.1	
	1568-1613(liq)	п п	90.4	96.2	
	1413 - 1518	Torsion effusion	92.9	96.2	
	1539 - 1614	11 11	96.4	96.2	
Greenbaum et al. [52]	1273-1513	н н	84.0	91.5	
Margrave [53]	1241-1492	Knudsen effusion	93.8	96.1	
Le Boucher [54]	1934-2129	Manometric	99.9	96.0	
Hildenbrand et al. [55]	1425-1533(c)	Torsion effusion	104.8	96.0	
	1540-1606(liq)	пп	102.8	95.9	
Hildenbrand and Theard [56]	1326-1527(c)	п п	91.5	93.3	
	1544-1604(liq)	11 11	82.4	95.1	
Gunther [57]	1284 - 1530	Knudsen effusion	95.4	95.6	

Table I

<u>MgCl(g)</u> Herzberg [40] and Gaydon [39] select D°_{0} to be 3.2 ev and 2.7 ev, respectively. These result in $\triangle Hf^{\circ} = -10$ and +1 kcal.

The thermal functions were obtained from the Dow Chemical Co. [41].

 $MgCl_2(c)$ The thermal functions were obtained from the Dow Chemical Co. [58].

Bommer and Hohman [59] measured the $\triangle H$ at 293°K for Mg(c) and MgCl₂(c) in 0.1N HCl to be -112.5 and -37.85 kcal, respectively. We obtain $\triangle H =$ -74.3 kcal for Mg(c) + 2HCl(0.1N) \rightarrow MgCl₂(c) + H₂(g) and \triangle Hf° = -153.9 kcal. Similarly Shomate and Huffman [16] obtained, in solutions involving 77.2HCl·4190 H₂O, $\triangle H = -111.285$ and -36.560 kcal, respectively, or $\triangle H = -74.725$ kcal for Mg(c) + 77.2(HCl + 54.275) $-MgCl_2(c) + 75.2$ (HCl + 55.718 H₂O) + H₂(g) and \triangle H^o_f = -153.28 kcal.

Sano [60] measured the equilibrium $MgCl_2(c) + H_2O(g) \rightarrow MgO(c) + 2HCl(g)$ in the range 796°K to 894°K. We obtain 2nd and 3rd law $\Delta H^\circ = 19.7$ and 23.14 kcal/mol MgCl₂ and $\Delta Hf^\circ = -149.9$ and -153.28 kcal.

Treadwell [61,62] and Sano [63] measured the equilibrium for $MgCl_2(c)$ + $\frac{1}{2}O_2 \rightarrow MgO(c) + Cl_2(g)$ in the temperature ranges 320°K to 976°K and 365°K to 943°K, respectively. The 2nd and 3rd law ΔH° 's are 9.0 and 9.42 kcal, from Treadwell's measurements, and 8.3 and 9.72 kcal from Sano's, which lead to ΔHf° = -152.8 and -153.23, and -152.1 and -153.6 kcal, respectively. Similarly from the results of Tsuchiya [64] in the temperature range 823°K to 973°K we obtain ΔHf° = -151.6 and -153.6 kcal from the 2nd and 3rd law approaches. In the temperature range 1013°K to 1153°K (MgCl_2(liq)) we obtain ΔHf° = -161.6 and -157.6 kcal, respectively. Reznikov's [65] measurements at 1023°K to 1223°K lead to 2nd law and 3rd law ΔHf° 's = -160.3 and -152.1 kcal, respectively.

Egan et al. [66] from EMF measurements on the reaction Mg(c) + $2AgCl(NaCl) \rightarrow MgCl_2(c) + 2Ag(c)$ combined with other data on the Ag/(AgCl - NaCl) half cell obtain the $\triangle Gf^{\circ}$ of MgCl_2(c) at 673°, 723° and 773°K. Their values lead to a 3rd law $\triangle Hf^{\circ} = -154.14$ kcal. From the cell measurements of Treadwell et al. [67], Markov et al. [68], Lorenz and Velde [69], and Takahashi [70] on the reaction Mg(c,liq) + $Cl_2(g) \rightarrow MgCl_2(liq)$ we obtain 3rd law $\triangle Hf^{\circ}$'s = -159.9, -154.9, -152.96, and -153.18 kcal, respectively.

 $\frac{\text{MgCl}_2(g)}{\text{were obtained from the Dow Chemical Co. [7]}}$.

Schrier and Clark [71] have shown that the gaseous species over $MgCl_2(liq)$, in the range 1200°K to 1400°K, is about 30% dimer. They have also measured the total vapor pressure. Correcting for the dimerization we obtain 2nd and 3rd law ΔH° subl = 57.6 kcal and 58.0 kcal, respectively. Using their equilibrium constants for the dimerization process we have corrected the measurements of Kushkin et al.[72] (1173°K to 1323°K) to 43.6 and 57.1 kcal, respectively, the measurements of Fischer et al. [73] (1136°K to 1435°K) to 58.0 and 56.9 kcal, Maier [74] (1056°K to 1400°K) to 50.6 and 57.2 kcal.

Berkowitz and Marquart [49], from Knudsen effusion-mass spectroscopy measurements, reported a monomer pressure at 921°K of 3.84 x 10^{-5} atm.

and $\Delta Hsubl = 54.6$ keal at 800°K. These measurements result in a 3rd law $\Delta H^\circ = 56.4$ keal and a 2nd law $\Delta H^\circ = 55.4$ keal.

Hildenbrand et al. [55] reported vapor pressure results from torsion effusion measurements over the temperature range 800°K to 985°K. Monomeric vapor was assumed. Second and third law ΔH° 's of 61.8 kcal and 57.8 kcal are obtained. We have selected ΔH° subl = 57.6 kcal.

 $\frac{\text{MgCl}_2(\text{aq, std. state, m = 1})}{\text{the ions. See Mg}^{2+}(\text{aq, std. state}), \text{MgCl}_2(\text{aq}), \text{Mg(ClO}_4)_2(\text{aq, std. state}), \text{MgCl}_2(\text{aq}), \text{Mg(ClO}_4)_2(\text{aq, std. state, m = 1}), \text{ and Mg(NO}_3)_2(\text{aq}).}$

 $\frac{\text{MgCl}_2(\text{aq})}{\text{Thomsen [36], and Fricke [77] measured \triangle Hdiln of aqueous $MgCl_2$.}$ $\frac{\text{Mishchenko and Yakovlev [78] measured the \triangle Hsoln at varying concentrations.}$

Measurements of the \triangle Hsoln MgCl₂·rH₂O(c) lead to the following values for \triangle Hf (aq, nH₂O), where n is the final concentration. Corrections to 25°C have been made. (See Table II.)

Auzhbikovich [86] reported $\triangle H^\circ = 16.4$ kcal/mol H₂O for the decomposition of the dihydrate to the anhydrous salt, or $\triangle Hf^\circ = -306.3$ kcal.

System	Inv	estigator	∆llf <u>kcal/mol</u>	/	°C of Meas.
MgC12•6H20	1.	Samoilov [79]	-190.79	1110	25
2 2	2.	Voskresenskaya and			
		Ponomareva [80]	-190.52	333	25
	3.	Thomsen [36]	-190.88	400	18
	4.	Richardson and			
		Wells [81]	-188.30	22	22.8
	5.	Berthelot and			
		Ilosvay [82]	-190.49	200	12.7
	6.	Mishchenko and			
		Yakovlev [78]	-190.79	1176	2
			-189.95	52	2
			-187.15	11.45	2
MgC1 ₂ •4H ₂ 0(c)	1.	Thomsen [36]	-191.23	400	18
$MgC1_2 \cdot 2H_2O(c)$	1.	Thomsen [36]	-190.66	400	18
MgCl ₂ (c)	1.	Mishchenko and			
-		Yakovlev [78]	-190.92	2270	2
	2.	Mishchenko et al. [83]	-190.8	1110	25
	3.	Thomsen [36]	-189.85	800	18
Mg(c) + equiv.					
am't. HC1.200 H ₂ 0	1.	Richards et al. [84]	-189.56	400	20

 $\frac{\text{MgCl}_{2} \cdot 4\text{H}_{2} \text{O}(c)}{\text{As for MgCl}_{2} \cdot 2\text{H}_{2} \text{O}(c), \text{ we obtain, from Shomate and Huffman's [16]}}$ $\frac{\text{MgCl}_{2} \cdot 2\text{H}_{2} \text{O}(c), \text{ we obtain, from Shomate and Huffman's [16]}}{\text{measurements } \Delta \text{H} = -102.056 \text{ kcal for Mg(c)} + 77.2(\text{HCl} + 54.275) + 4\text{H}_{2} \text{O}(1\text{iq}) \rightarrow \text{MgCl}_{2} \cdot 4\text{H}_{2} \text{O}(c) + 75.2(\text{HCl} + 55.718 \text{ H}_{2} \text{O}) \text{ and } \Delta \text{Hf}^{\circ} = -453.87 \text{ kcal.}}$

Kondirev and Berezovskii [87], Derby and Yngve [88], Sano [89], and Auzhbikovich [86] measured the decomposition pressure of the hexahydrate to the tetrahydrate. A log p vs 1/T plot of the data results in $\Delta Hf^{\circ} =$ -453.99 kcal. Kondirev and Berezovskii [87] and Sano [90] measured the decomposition pressures of the tetrahydrate to the dihydrate. A log p vs 1/T plot results in $\Delta H f^\circ = -454.05$. Auzhbikovich's measurements result in $\Delta H f^\circ = -457.1$.

 $\frac{\text{MgCl}_{2} \cdot 6\text{H}_{2}\text{O}(c)}{\text{As for MgCl}_{2} \cdot 4\text{H}_{2}\text{O}(c)}$ The Cp° and S° were obtained from Kelley and King [85]. As for MgCl}_{2} \cdot 4\text{H}_{2}\text{O}(c) we obtain from Shomate and Huffman's [16] measurements $\Delta \text{H} = -108.836$ kcal for Mg(c) + 77.2(HCl + 54.275) + 6H_{2}\text{O}(liq) \rightarrow \text{MgCl}_{2} \cdot 6\text{H}_{2}\text{O}(c) + 75.2(HCl + 55.718 \text{ H}_{2}\text{O}) and $\Delta \text{Hf}^{\circ} = -597.28$ kcal.

Mg(ClO₄)₂(c) Vorob'ev et al. [91] reported \triangle Hsoln in 800 H₂O to be -37.08 kcal. This results in \triangle Hf° = -135.97 kcal. Drakin and Chzhan [92] reported four measurements in the range from m = .02 to .09 (where m is the molality), which lead to \triangle Hf° = -135.68 kcal. Smeets' [93] measurements for m = 0.12 result in \triangle Hf° = -135.53 kcal. Measurements have also been reported by Smith et al. [94]. We have selected the first mentioned value.

 $Mg(ClO_4)_2$ (aq, std. state, m = 1) The final values were obtained from the ions. See Mg²⁺ (aq, std. state, m = 1), MgCl₂(aq) and Mg(NO₃)₂(aq). Jongenburger and Wood [95] measured the \triangle H diln from 3.2 to .0018 molal.

Vorob'ev et al. [91] reported $\triangle H = -36.11$ kcal for the reaction, MgO(c) + 2HClO₄ · 800 H₂O \rightarrow Mg(ClO₄)₂ · 800 H₂O + H₂O. This results in $\triangle Hf = -173.28$ kcal for Mg(ClO₄)₂ · 800 H₂O and $\triangle Hf^{\circ} = -173.63$ kcal.

Mg(Cl0₄)₂·6H₂O(c) Smeets [93] reported measurements of △Hsoln at 18°C for the hexahydrate and the anhydrous salt. We obtain △H° = -38.71 kcal for Mg(Cl0₄)₂(c) + 6H₂O(liq) → Mg(Cl0₄)₂·6H₂O(c) and △Hf° = -584.57 kcal. Similarly from Smith et al. [94] we obtain △H° = -32.7 kcal and △Hf° = -578.6 kcal. Their value for △H soln = 0.8 kcal for the hexahydrate leads to △Hf° = -583.8 kcal. Nosova [96] reported △H soln to a final concentration of 0.07 m = 1.45 kcal, from which we obtain △Hf° = -584.39 kcal. Linke [5] lists the saturation concentration to be 4.48 molal. Using this and the activity coefficient and osmotic coefficient from Robinson and Stokes [6] we obtain ΔG° soln = -7.60 kcal, which results in ΔGf° = -445.34 kcal and S° = 124.5 e.u.

<u>MgOHCl</u>(c) Kelley [97] evaluated the data on the MgOHCl(c) system and obtained for MgCl₂·H₂O(c) \rightarrow MgOHCl(c) + HCl(g) \triangle H° = 17.82 and \triangle G° = 8.34 kcal. These lead to \triangle Hf° = -191.1 kcal, \triangle Gf° = -174.9, and S° = 20.0 e.u.

Savinkova and Dagtyareva [98] reported the equilibrium from 743°K to 1023°K for MgCl₂(solid soln) + H₂O(g) \rightarrow MgOHCl(solid soln) + HCl(g). The K's include the activities. Their results give $\triangle H^\circ = -1.5$ kcal and $\triangle Hf^\circ = -191.1$ kcal. Vel'nyanskii and Savinkova [99] do not give the activities; hence, their values have not been used.

MgBr₂(c) Beketov [100] reported △Hsoln in 800 H₂O = -43.30 kcal, from which we obtain △Hf° = -125.26 kcal. Finch et al. [101] report △H = -35.23 kcal for the reaction of MgO(c) in excess HBr(m = 0.5) to form MgBr₂·2000 H₂O, and △H = -43.14 kcal for the solution of MgBr₂(c) in excess HBr(m = 0.5) to form MgBr₂·9000 H₂O. Correcting for the concentration change, we obtain △H = 7.78 kcal for MgO(c) + 2HBr(aq) → MgBr₂(c) + H₂O(liq), and △Hf° = -125.36 kcal.

Toguri et al. [102] reported $\triangle G^{\circ}_{1073^{\circ}K} = 0.267$ kcal for $\frac{1}{2}MgBr_{2}(liq)$ + HCl(g) $\rightarrow \frac{1}{2}MgCl_{2}(liq)$ + HBr(g). Using estimated thermal functions from the Dow Chemical Co. [58] we obtain $\triangle H^{\circ} = -2.3$ kcal and $\triangle Hf^{\circ} = -121.9$ kcal.

S° has been estimated as 28.0 e.u. by Kelley and King [85].

MgBr₂(g) Berkowitz and Marquart [49] obtained the vapor pressure by the mass spectroscopy-Knudsen effusion method. A 3rd law ΔH° from measurements at 798°K and 842°K is 49.7 kcal. The 2nd law value, at 727°K, results in $\Delta H^{\circ} = 52.2$ kcal. We have taken ΔH° subl = 51.3 kcal and $\Delta Hf^{\circ} = -74.0$ kcal. MgBr₂(aq, std. state, m = 1) The values were obtained from the ions.

MgBr₂(aq) Lange and Streek [75] measured the heats of dilution.

MgBr₂·6H₂O(c) The saturation solubility was obtained from Linke [5]; the activity and osmotic coefficients at saturation from Robinson and Stokes [6]. The resultant △G°soln = -7.13 kcal. We have estimated S° = 95. e.u. This results in △Hf° = -575.9 kcal. A value of 40.7 kcal is obtained for the △H° for the reaction MgBr₂·6H₂O(c) → MgBr₂(c) + $6H_2O(1iq)$. This appears high in relation to △H° = 34.1 for the comparable chloride system; however, from the differential △H soln (obtained from the solubility measurements), which was corrected to infinite dilution we obtain △H°soln = -3.4 kcal and △Hf° = -576.2 kcal.

MgI₂(c) From Beketov's [100] measured △H soln in 880 H₂O, we have $\overline{\Delta Hf^{\circ}} = -87.1$ kcal. Finch et al. [101] report △H = -35.10 kcal for the reaction of MgO(c) in excess HI (m = 0.5) to form MgI₂·2000 H₂O; and $\Delta H = -49.75$ kcal for the solution of MgBr₂(c) in HI to form MgBr₂·9000 H₂O. Correcting for the concentration change, we obtain $\Delta H = 14.50$ kcal for MgO(c) + 2HI(aq) \rightarrow MgBr₂(c) + H₂O(liq) and $\Delta Hf^{\circ} = -87.0$ kcal.

Kelley and King [85] have estimated S° = 31.0 e.u.

 $MgI_2(g)$ Berkowitz and Marquart [49] measured the vapor pressure by the mass spectroscopy-Knudsen effusion method. A 3rd law ΔH° from measurements at 757°K and 734°K is 45. kcal(the $\Delta(G^\circ-H^\circ_{298})/T$ was estimated). The 2nd law value at 640°K results in 46.4 kcal. We have taken ΔH° subl = 46 kcal from which we obtain ΔHf° = -41 kcal.

 $MgI_2(aq, std. state, m = 1)$ The values were obtained from the ions.

 $MgI_2(aq)$ The values were estimated from a consideration of the $\triangle Hdiln$ of $MgCl_2$ and $MgBr_2$, and the Cl-Er-I system of the monovalent electrolytes.

MgSO₄(c) The Cp° and S° were obtained from Kelley and King [85]. The following values for Δ Hsoln have been obtained: Thomsen [36] -20.7 kcal (nH₂O = 420); Berthelot and Ilosvay [82] -20.8 (nH₂O = 100); Pickering [103] -20.9 and [104] -20.8 kcal (nH₂O = 420). These result in Δ Hf° = -307.1, -306.8, -307.0, and -307.1 kcal, respectively.

Kelley [105] evaluated the data on the equilibrium $2MgSO_4 \rightarrow 2MgO(c)$ + $2SO_2(g) + O_2(g)$ and obtained $\triangle H^\circ = 196.0$ kcal, which leads to $\triangle Hf^\circ = -312.8$ kcal. The results of Dewing and Richardson [106] yield $\triangle H^\circ = 189.6$ kcal and $\triangle Hf^\circ = -309.6$ kcal.

From Knopf and Staude's [107]equilibrium measurements for MgSO₄(c) \rightarrow MgO(c) + SO₃(g) we obtain \triangle H° = 58.3 kcal and \triangle Hf° = -296.7 kcal.

 $\frac{MgSO_4}{4}$ (aq, std. state, m = 1) The final values were obtained from the ions. See Mg^{2+} (aq, std. state, m = 1).

<u>MgSO₄(aq)</u> The heats of dilution were measured by Lange [109], Lange and Streeck [110], Lange and Messner [111,112], Plake [113], and Thomsen [36]. Kaganovich and Mishchenko [114] measured △Hsoln of MgSO₄·7H₂O(c) as a function of the concentration. From Thomsen's [36] reported values for the reactions: (1) MgCl₂(200 H₂O) + H₂SO₄(200 H₂O) \rightarrow MgSO₄(200 H₂O) + 2HCl(100 H₂O); (2) MgCl₂(800 H₂O) + H₂SO₄(400 H₂O) \rightarrow MgSO₄(400 H₂O) + 2HCl(400 H₂O); and (3) MgSO₄(400 H₂O) + 2HNO₃(400 H₂O) \rightarrow Mg(NO₃)₂(800 H₂O) + H₂SO₄(400 H₂O), we obtain (1) △H = -3.82 kcal, (2) △H = -3.76 kcal, and (3) △H = +3.99 kcal, which result in (1) △Hf (200 H₂O) = -327.70, (2) △Hf (400 H₂O) = -327.92 and (3) △Hf (400 H₂O) =

-328.12 kcal. These are in good agreement with the values obtained from the use of the heats of diln and the ions.

 $\frac{\text{MgSO}_4 \cdot \text{H}_2 \text{O(c)}}{\text{The Cp}^\circ \text{ and S}^\circ \text{ were obtained from Kelley and King [85].}}$ Thomsen [36] and Pickering [103] measured ΔH soln in 400 H₂O. Their results lead to $\Delta \text{Hf}^\circ = -382.5$ and -384.0 kcal. Jamieson and Frost [115] reported ΔH (soln, ∞) = -14.26 kcal which results in $\Delta \text{Hf}^\circ = -382.9$ kcal.

Kohler and Zaske [116] reported $\Delta H = 15.0$ kcal for the decomposition to the anhydrous sulfate and $H_0O(g)$. This results in $\Delta Hf^\circ = -379.9$ kcal.

 $\frac{MgSO_4 \cdot H_2O(\text{amorph})}{-20.8 \text{ kcal}; \Delta Hf^\circ, \text{ then, is -376.4 kcal}} = \frac{MgSO_4 \cdot H_2O(\text{amorph})}{-20.8 \text{ kcal}; \Delta Hf^\circ, \text{ then, is -376.4 kcal}}$

Frost et al. [117] report $S^{\circ} = 33.0 \text{ e.u.}$

 $\frac{MgSO_4 \cdot 2H_2O(c)}{\Delta Hf^\circ = -453.2 \text{ kcal.}}$ Thomsen's [36] ΔH soln in 400 H₂O is -11.4 kcal, or

Kohler and Zaske [116] reported $\triangle H = 25.3$ kcal for the decomposition to the anhydrous salt and $H_2O(g)$; this results in $\triangle Hf^\circ = -455.8$ kcal.

 $\frac{\text{MgSO}_4 \cdot 4\text{H}_2\text{O(c)}}{\Delta \text{Hf}^\circ = -596.7 \text{ kcal.}}$ Thomsen's [36] ΔH soln in 400 H₂O is -4.45 kcal or

Kohler and Zaske [116] reported $\triangle H = 25.8$ kcal for the decomposition of the hexahydrate to the tetrahydrate and $H_2O(g)$. $\triangle Hf^\circ$, then, = -597.4 kcal.

 $\frac{MgSO_4 \cdot 6H_2O(c)}{The Cp^{\circ} \text{ and } S^{\circ} \text{ were taken from Kelley and King [85].}}{Thomsen [36] reported <math>\triangle H$ soln in 400 H_2O to be +0.02 kcal from which we obtain $\triangle Hf^{\circ} = -737.8$ kcal.

Carpenter and Jette [118], Rolla and Accame [119], Bell [120], and Kohler and Zaske [116], reported values for ΔH for decomposition of the heptahydrate to the hexahydrate and $H_2O(g)$. Their values are 14.0, 14.6, 14.1, and 14.7 kcal, or $\Delta Hf^\circ = -738.1$, -737.5, -738.0, and -737.5kcal, respectively. MgSO₄ $^{\circ}$ 7H₂O(c) Kaganovich and Mishchenko [114] measured \triangle Hsoln from m = .25 to m = 2.2. Their values at these concentrations are 3.12 and 4.54 kcal. The 21 measurements result in \triangle Hf° = -809.92 kcal. Kapustinskii and Samoilov [121] reported \triangle Hsoln = 1.82 kcal at m = .03. This results in \triangle Hf° = -808.20 kcal. From Perreu's [122] \triangle Hsoln measurements at 284°K, from nH₂O = 75 to nH₂O = 21, we obtain \triangle H = 4.0 to 4.5 kcal or \triangle Hf° = -809.74 to -809.88 kcal. From Thomsen's [36] and Pickering's [103] \triangle Hsoln, 400 H₂O we have \triangle H = 3.74 and 3.90 kcal or \triangle Hf° = -809.8 and -810.00 kcal.

From the solubility given by Linke[5] and the activities and osmotic coefficients given by Robinson and Stokes [6] we obtain $\triangle G^\circ$ soln = 2.29 kcal and $\triangle Gf^\circ$ = -685.8 kcal. Using this value for $\triangle Gf^\circ$ and a $\triangle Hf^\circ$ = -809.92 we obtain S° = 87 e.u. which appears low. From the decomposition pressure measurements of the heptahydrate to the hexahydrate of Bell [120] we obtain $\triangle H^\circ$ = 14.1 and $\triangle G^\circ$ = 2.42 kcal from which we obtain S° = 89.0 e.u. Kohler and Zaske's [116] decomposition pressure is in agreement.

 $\frac{Mg_{3}N_{2}(c, \alpha)}{\text{ing reactions to be -287.89 kcal and -32.08 kcal, respectively.}}$

(1) $Mg_{3}N_{2}(c) + 231.6HC1 \cdot 12570 H_{2}O \rightarrow 224.6HC1 \cdot 3MgC1_{2} \cdot 2NH_{4}C1 \cdot 12570 H_{2}O$

(2) $2NH_3(1iq) + 226.6HC1 \cdot 12570 H_2O \rightarrow 224.6HC1 \cdot 3MgC1_2 \cdot 2NH_4C1 \cdot 12570 H_2O$ Using the ΔH of Shomate and Huffman [16] for the reaction of Mg(c) in HC1(aq) and the ΔH vap of $NH_3(1iq)$, he obtains $\Delta H^\circ = -88.05$ kcal for $3Mg + 2NH_3(g) \rightarrow Mg_3N_2(c) + 3H_2(g)$ from which we obtain $\Delta Hf^\circ = -110.09$ kcal.

Neumann et al. [124] reported the values of $\triangle H$ for the following to be -284.8 and -336.0 kcal, respectively.

(1) $Mg_{3}N_{2} + 8(HC1 + 20 H_{2}O) \rightarrow (3MgC1_{2} + 2NH_{4}C1)(aq)$

(2) $3Mg(c) + 6(HC1 + 20 H_2O) \rightarrow 3MgC1_2(aq) + 3H_2(g)$

Using the $\Delta H = 4.14$ kcal from Juza et al. [125] for the solution of $NH_4Cl(c)$ in HCl(aq) they obtain $\Delta H = -42.9$ kcal for $3Mg(c) + 2NH_4Cl(c) \rightarrow Mg_3N_2(c) + 2(HCl + 20 H_2O) + 3H_2(g)$ from which $\Delta Hf^\circ = -114.9$ kcal.

Neumann et al. [126] directly determined the value of △Hf° to be -116.6 kcal.

The results of Brunner are cited by Neumann et al. [124]. $\Delta H = -177.0 \text{ kcal for } \text{Mg}_3\text{N}_2(c)+5\text{H}_2O(c) \rightarrow 3\text{Mg}(OH)_2(\text{ppt.}) + 2\text{NH}_3(aq) \text{ from which}$ we obtain $\Delta Hf^\circ = -111.4 \text{ kcal.}$ Matignon [127] and Moser and Herzner [128] measured the $\Delta H = -279.7$ and -269.9 kcal, respectively for the reaction of Mg_3N_2 with $\text{H}_2\text{SO}_4 \cdot 200 \text{ H}_2\text{O}$. The values for ΔHf° are -133.0 and -143.0 kcal.

Dreger [129] measured the decomposition pressure to Mg(g) and $N_{2}(g)$ from 1118° to 1254°K, and obtained a $\triangle H^\circ$ apparent = 295.0 kcal, or a △Hf° = -189.1 kcal. Blank [130] obtained nonequilibrium values from Torsion effusion measurements in the range 1000-1250 °K of 286.3 and 218.7 kcal/mol $Mg_{3}N_{2}$ for the 3rd law and 2nd law values, respectively. Hildenbrand and Theard [56], in similar measurements, obtained a 2nd law non equilibrium $\triangle H^\circ = 309$ kcal. These values lead to $\triangle Hf^\circ = -180.4$, -112.8, and -203.1 kcal. Both Blank and Hildenbrand determined the condensation coefficients and corrected their vapor pressures to obtain equilibrium values. Blank's results now yield $\Delta H^{\circ} = 242.3$ kcal (3rd law) and 239.1 kcal (2nd law) and △Hf°= -136.4 and -133.2 kcal, respectively. Hildenbrand's measurements result in an equilibrium $\triangle H^{\circ} = 243.5$ kcal and a $\triangle Hf^{\circ} = -137.6$ kcal. The latter three values are in good agreement; however, as Hildenbrand points out, there is a possibility that there is a secondary reaction going on that would interfere with the decomposition reaction. Until there is further evidence supporting this, we have selected the first mentioned AHf° = -110.1 kcal.

The Dow Chemical Co. [131] tabulates the thermal functions.

(3) $Mg(NO_3)_2(c) + 77.2HC1.4190 H_2O \rightarrow Mg(NO_3)_2.77.2HC1.4190 H_2O; \Delta H = -20.634$ kcal Using the reaction of Shomate and Huffman [16], Mg(c)+77.2HC1.4190 H_2O $\rightarrow MgC1_2.75.2$ HC1.4190 H_2O+H_2(g); $\Delta H = -111.285$ kcal, we obtain $\Delta H = -90.340$ kcal for Mg(c) + 2HNO_3.26.5 H_2O $\rightarrow Mg(NO_3)_2(c) + H_2(g)$ and $\Delta Hf^\circ = -188.97$ kcal.

 $\frac{Mg(NO_3)_2}{aq}$, std. state, m = 1) The final values were obtained from the ions. See Mg²⁺(aq, std. state, m = 1)

 $Mg(NO_3)_2$ (aq) Lange and Streeck [110], Hammerschmid and Lange [133], Ewing et al. [134], and Thomsen [36] measured $\triangle H$ diln.

Pickering [135], Thomsen [36], and Ewing et al. [134] measured $\triangle H$ soln. of the hexahydrate. Their results lead to $\triangle Hf = -210.25$ kcal (400 H₂O), -210.58 kcal (400 H₂O), and -210.04 kcal (130 H₂O), respectively. Ewing also measured the $\triangle H$ soln. of the anhydrous salt, from which we obtain $\triangle Hf(130 H_2O) = -210.20$ kcal.

Lloyd et al. [136] reported decomposition pressure measurements to the anhydrous salt in the temperature range from 293°K to 333 °K. We obtain Δ Hf° = -618.3 kcal.

From the solubility given by Linke [5] and the activity coefficients and osmotic coefficients given by Robinson and Stokes [6] we obtain $\Delta G^{\circ}soln = -4.11$ kcal or $\Delta Gf^{\circ} = -497.93$ kcal. This results in $\Delta Sf^{\circ} =$ -425.3 e.u. and $S^{\circ} = 109.5$ e.u. This appears a little high. We have estimated $S^{\circ} = 103$ e.u. and have adjusted ΔCf° accordingly.

 $MgCO_3(c)$ The Cp° and S° were obtained from Kelley and King [85]; the high temperature functions from Kelley [138].

Kapustinskii and Stakhanova [139] measured the following AH's:

- (1) $MgCO_3(c) + 2HCl(aq) \rightarrow MgCl_2(aq) + H_2O(liq) + CO_2(g); \Delta H = -12.80$ kcal
- (2) $MgO(c) + 2HC1(aq) \rightarrow MgCl_2(aq) + H_2O(liq); \Delta H = -36.32$ kcal

This results in $\triangle H = +23.52$ kcal for $MgCO_3(c) \rightarrow CO_2(g) + MgO(c)$, or $\triangle Hf^\circ = -261.4$ kcal.

Roth [140] reported the $\triangle H^{\circ} decomp. = +23.6 \text{ kcal or } \triangle Hf^{\circ} = -261.5 \text{ kcal.}$ Berthelot [141] reported $\triangle H = 2.7 \text{ kcal for the precipitation}$ of MgCO₃ from K₂CO₃(220 H O) and MgSO₄(220 H₂O). We obtain $\triangle Hf^{\circ} = -268.1 \text{ kcal.}$ Ellis and Mortland [142] and Foldvari-Vogl and Kliburszky [143] obtained $\triangle H^{\circ} decomp.$ from differential thermal analysis methods, to be 27.6 kcal and 27.0 kcal, respectively, or $\triangle Hf^{\circ} = -265.5 \text{ and } -264.9 \text{ kcal.}$

We obtain from Marc and Simek's [144] vapor pressure measurements $\triangle H^{\circ}$ decomp. = 28.2 kcal (3rd law) and 26.4 kcal (2nd law) or $\triangle Hf^{\circ}$ = -266.1 kcal and -264.3 kcal.

Langmuir [145] reviews the above data as well as the solubility measurements and the derived $\triangle Gf^{\circ}$ values and points out the inconsistencies in the stability relationships in the MgO-CO₂-H₂O system. We have accepted his interpretation and calculations to obtain $\triangle G^{\circ} =$ 0.665 kcal for MgCO₃·3H₂O(c) \rightarrow MgCO₃(c) + 3H₂O(liq). Using our value for $\triangle Gf^{\circ}$ for the trihydrate = -412.6 kcal, we obtain $\triangle Gf^{\circ} = -241.9$ kcal. Mg₂Si(c) Kubaschewski and Villa [146] reported \triangle Hf° = -18.6 kcal. Eremenko and Lukashenko [147], from ENF measurements obtained E(in ev) = 0.2075 - 0.77x10⁻⁴T for the range 673°K to 873°K, from which we obtain \triangle Hf°_{773°K} = -19.1 kcal. This corrects to \triangle Hf° = -18.7 kcal. Grjotheim et al. [148] measured the decomposition pressure in the range 1130 to 1273°K by the transpiration technique. We obtain \triangle H° = 41.1 kcal/mol Mg(g) or \triangle Hf° = -11.6 kcal.

From Eremenko and Lukashenko we also obtain a $\Delta Sf^{\circ}(773^{\circ}K) = -7.1$ e.u. which, assuming a $\Delta Cp = -1$ cal/mol deg, corrects to $\Delta Sf^{\circ} = -6.1$ and $S^{\circ} = 14.0$ e.u. This value appears low. Mannchen and Jacobi [149] reported Cp measurements from 12°K to 300°K on Mg_{2.12}Si(c) and S° = 19.8 cal/mol. Assuming their reported S° is for Mg_{2.12}Si(c) we obtain S° Mg₂Si(c) = 18.6 e.u. and a $\Delta Sf^{\circ} = -1.5$ e.u.

For the corresponding Ge compound $(S^{\circ}_{Ge}(c) = 7.43 \text{ e.u.})$ we have $S^{\circ} = 20.7 \text{ e.u.}$ and $\triangle Sf^{\circ} = -2.35 \text{ e.u.}$ In comparison with these figures we feel 18.6 e.u. is high (and -1.5 e.u. too positive). We have therefore adjusted S° to = 18.0 e.u.

 $MgSiO_3(c)$ Furukawa and Reilly [150] evaluated the low temperature Cp and high temperature heat content and have tabulated the thermal functions up to 2000°K.

Torgeson and Sahama [34] by HF soln calorimetry of $Mg(OH)_2(c)$, SiO₂(c, quartz), and MgSiO₃(c) obtained $\triangle H = 0.16$ kcal for Mg(OH)₂(c) + SiO₂(c) \rightarrow MgSiO₃(c) + H₂O(liq). This results in $\triangle Hf^\circ = -370.22$ kcal.

Henderson and Taylor [151] reported $\triangle G^{\circ}_{1823^{\circ}K} = -5.1$ kcal for the formation from the oxides. This results in $\triangle Hf^{\circ} = -368.5$ kcal.

 Mg_2SiO_4 (c) Furukawa and Reilly [150] tabulated the thermal functions up to 2000°K.

Torgeson and Sahama [34] by HF soln. calorimetry of Mg(OH)₂(c), SiO₂(c, quartz), and Mg₂SiO₄(c) obtained $\triangle H = 2.58$ kcal for 2Mg(OH)₂(c) + SiO₂(c) \rightarrow Mg₂SiO₄(c) + 2H₂O(liq), from which we obtain $\triangle Hf^{\circ} = -520.45$ kcal. King et al. [152] from HF soln. calorimetry obtained $\Delta H^{\circ} = -13.86$ kcal for $2MgO(c) + SiO_2(c) \rightarrow Mg_2SiO_4(c)$ which leads to $\Delta Hf^{\circ} = -519.20$ kcal. Henderson and Taylor [151] reported $\Delta G^{\circ}_{1823^{\circ}K} = -9.4$ kcal for the formation of $Mg_2SiO_4(c)$ from the oxides. This results in $\Delta Hf^{\circ} = -515.4$ kcal.

The Cp° and S° were also obtained from King et al.

 $\frac{Mg_{3}Si_{4}O_{10}(OH)_{2}(c)}{-44.9 \text{ kcal for } 3MgO(c) + 4SiO_{2}(c) + H_{2}O(1iq) \rightarrow Mg_{3}Si_{4}O_{10}(OH)_{2}(c) \text{ from } Which we obtain <math>\Delta Hf^{\circ} = -1415.5 \text{ kcal.}$

The Cp° and S° were obtained from Robie and Stout [154].

 $\frac{MgB_{12}}{[155]}$ (c) The thermal functions were obtained from Germaidze et al. [155] who also measured the Mg(g) pressure over the MgB₁₂ to be 10⁻⁵ atm. at 1700°K. This results in Δ Hf° = -53.4 kcal.

 $\frac{\text{MgAl}_{2}\text{O}_{4}(c)}{\text{Navrotsky and Kleppa [157] measured the } \Delta \text{H soln. of MgO(c), Al}_{2}\text{O}_{3}}$ (c, α) and MgAl}_{2}\text{O}_{4}(c) in 9PbO·3CdO·4B $_{2}\text{O}_{3}$ (melt) and obtained $\Delta \text{H}_{970^{\circ}\text{K}}^{=}$ -8.72 kcal for the formation from the component oxides. This results in $\Delta \text{H}^{\circ} = -8.6$ kcal and $\Delta \text{Hf}^{\circ} = -552.9$ kcal.

Altman [20] from Knudsen effusion decomposition pressure measurements $[MgAl_2O_4(c) \rightarrow Mg(g) + \frac{1}{2}O_2(g) + Al_2O_3(c)]$, supports a $\Delta H^\circ = -5$ kcal for the formation from the oxides. Altman [158] from transpiration experiments on the Al-MgO and Al-MgAl_2O_4 systems obtained $\Delta H^\circ = -7$ kcal. Grjotheim et al. [159], by the transpiration method, measured the K's for $4MgO(c) + 2Al(1iq) \rightarrow MgAl_2O_4(c) + 3Mg(g)$. They considered the system ideal. On this basis we obtain $\Delta H^\circ = 7.3$ kcal (2nd law) and -7.6 kcal (3rd law). Altman [20] corrects their results for the activity of Al. Using the thermal functions mentioned above we obtain a 3rd law $\Delta H^{\circ} = -8.2$ kcal and $\Delta ME^{\circ} = -552.5$ kcal. Grjotheim et al. [160] repeat their measurements and obtain a $\Delta M^{\circ} = +1.1$ kcal. In a later series of measurements on the Al-Mg-SiO system they [161] obtain the activity of Al and pressure of Mg(g) for the equilibrium process, $4MgO(c) + 2AlSi(liq) \neg MgAl_2O_4(c) + 3Mg(g)$. This results in $\Delta M^{\circ} = -7.4$ kcal for formation from the oxides and $\Delta ME^{\circ} = -551.7$ kcal. In a reaction involving MgO(c) and $Al_4C_3(c)$ they [162] obtain $\Delta M^{\circ} = -11.1$ kcal and $\Delta HE^{\circ} = -555.4$ kcal. Rao and Dadape [163] measured the K's of the reaction, $4MgO(c) + 3AlCl(g) \neq MgAl_2O_4(c) + AlCl_3(g) + 3Mg(g)$ in the range 1298°K to 1465°K. We obtain $\Delta ME^{\circ} = -554.0$ kcal (3rd law) and a $\Delta H^{\circ} = -9.7$ kcal for formation from the oxides and $\Delta HE^{\circ} = -554.7$ kcal (2nd law) and a $\Delta H^{\circ} = -10.4$ kcal.

Zaionchovskii and Rubal'skaya [164] reported \triangle Hf° = -567.1 kcal. The first mentioned \triangle Hf° was selected.

 $\frac{Mg_2Al_4Si_5O_{18}(c)}{[165]}$; the high temperature heat content from Pankratz and Kelley [166].

Henderson and Taylor [151] reported $\triangle G^{\circ}_{1733^{\circ}K} = -12.5$ kcal for $2MgO(c) + 2Al_2O_3(c) + 5SiO_2(c) \rightarrow Mg_2Al_4Si_5O_{18}(c)$ from which we obtain $\triangle Hf^{\circ} = -2177$.kcal.

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Selected Values for Some Mg Compounds at 298°K

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Compound	State	∆Hf° kcal/mole	∆Gf° kcal/mole	S° cal/deg mol
Mg	с	0	0	7.81
Mg	g	35.30	27.04	35.502
Mg+	g	213.100		
Mg ²⁺	g	561.299		
Mg ²⁺	aq std. state, m =	1 -111.58	-108.7	-33.0
MgO	macro crystal (periclase)	-143.81	-136.10	6.44
MgO	micro crystal	-142.92	-135.27	6.67
MgO	g	4.		
MgH2	с	-18.0	-8.6	7.43
MgOH+	aq std. state, m =	1	-149.8	
Mg(OH) ₂	с	-220.97	-199.23	15.10
Mg(OH) ₂	ppt	-220.0		
Mg(OH) ₂	aq std. state, m = 1	1 - 221.52	-183.9	-35.6
Mg(OH) ₂	g	-134		
MgF	g	-53.0	-59.2	52.79
MgF ₂	с	-268.5	-255.8	13.68
MgF2	g	-173.0	-174.6	61.73
MgC1	g	-10.	-16.	55.8
MgC12	c	-153.28	-141.45	21.42
MgCl ₂	g	~ 95 . 7		
MgC12	aq			ť
	std. state, m = 1	1 -191.48	-171.4	-6.0
	50 H ₂ 0	-190.23		•
	100 H ₂ О	-190.59		
	400 H ₂ O	- 190,•94		
	. 1,000 H ₂ 0	-191.080		
	5,000 H ₂ O	-191.269		
	10,000 H ₂ O	-191.323		
	∞ H ₂ O	-191.48		
		120		

Compound	State	∆Hf° kcal/mole	∆Gf° kcal/mole	S° cal/deg mol
MgC12•H20	с	-231.03	-205.98	32.8
MgC1 ₂ •2H ₂ O	с	-305.86	-267.24	43.0
MgC12•4H20	с	-453.87	-388.03	63.1
MgC12•6H20	с	-597.28	-505.49	87.5
Mg(C10 ₄) ₂	с	-135.97		
Mg(C10 ₄) ₂	aq std. state, m =	1 -173.40	-112.8	54.0
Mg(C104) ₂ •6H ₂ 0	с	-584.5	-445.3	124.5
MgOHC1	с	-191.1	-174.9	20.0
MgBr ₂	с	-125.3	-120.4	28.0
MgBr ₂	g	-74.0		
MgBr ₂	aq			
	std. state, m =		-158.4	6.4
	100 H ₂ 0	-168.98		
	400 H ₂ 0	-169.18		
	1,000 H ₂ 0	-169.304		
	5,000 H ₂ O	-139.476		
	10,000 H ₂ 0	-169.528		
	∞ H2O	-169.68		
MgBr2·6H2O	С	-576.0	-491.4	95
MgI ₂	С	-87.0	-85.6	31.0
MgI2	g	-41		
MgI ₂	aq std. state, m =	1 -137.96	-133.4	20.2
	400 H ₂ 0	-137.49		
	1,000 H ₂ 0	-137.59		
	∞ H ₂ O	-137.96		
MgSO4	с	-307.1	-279.8	21.9
MgSO ₄	aq., std. stat m = 1, undisso		-289.74	-1.7

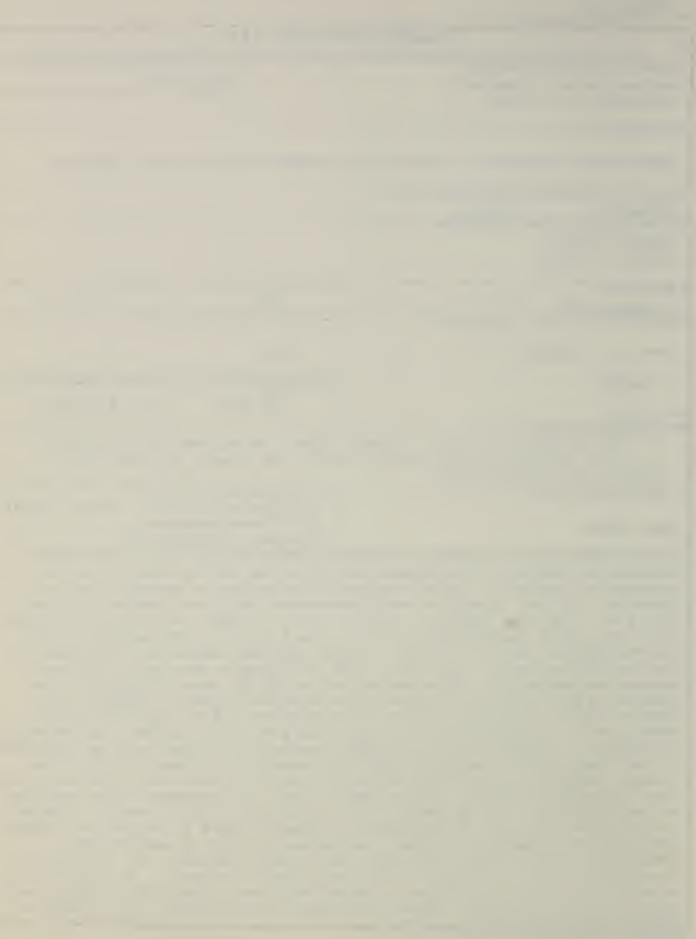
Compound	State	∆Hf° kcal/mole	∆Gf° kcal/mole	S° cal/deg mol
MgS04	aq	1 200 00	206 7	00.0
	std. state, m =		-286.7	-28.2
	50 H ₂ O			
	100 H ₂ 0			
	400 H ₂ 0			
	1,000 H ₂ 0			
	5,000 H ₂ 0			
	10,000 H ₂ O			
	∞ H ₂ O .			
$MgSO_4 \cdot H_2O$	С	-382.9	-341.5	30.2
MgSO ₄ •H ₂ O	amorph	-376.4	-335.8	33.0
$MgSO_4 - 2H_2O$	с	-453.2		
MgSO4•4H20	С	-596.7		
$MgSO_4 \cdot 6H_2O$	с	-737.8	-629.1	83.2
MgS04•7H20	с	~809.9 2	-686.4	89.
Mg3N2	с	-110.1		
Mg(NO3)2	с	-188.97	-140.9	39.2
$Mg(NO_3)_2$	aq			
	std. state, m =		-161.9	37.0
	50 H ₂ O	-210.14		
	100 H ₂ О	-210.16		
	400 H ₂ 0	-210.28	•	
	1,000 H ₂ 0	- 210.364		
	5,000 H ₂ 0	-210,502	·e	
	10,000 H ₂ 0	-210.551		
	∞ H ₂ O	-210.70		
Mg(NO ₃) ₂ •6H ₂ O	с	- 624 . 59	-497.3	108
MgC2	с	20		
Mg ₂ C ₃	с	17		
MgCO3	c, magnesite	-261.9	-241.9	15 . 7

Compound	State	Alf° kcal/mole	∆Gf° kcal/mole	S° cal/dcg mol
Mg ₂ Si	с	-18.6	-18.0	18.
MgSiO3	c, clinoensta	tite -370.22	~ 349 . 46	16.19
Mg2SiO4	c, forsterite	-519.6	-491.2	22.74
Mg3Si205(OH)4	c, chrysotile	-1043.4	-965.1	52,9
Mg3Si4010(OH)2	c, talc	-1415.5	-1324.8	62.3
MgB ₁₂	с	- 53.	-52.	21.4
Mg0•A1203	с	∽ 552.9	-523.1	19.27
Mg2A14Si5018	c, cordierite	-2177	- 2055	97.3

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13. ABSTRACT This report presents the results	and interpre	etation of	five recent NBS
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treatment and evaluation, and one new crit	ical assess	ment of pul	blished heats of
formation. The heat capacity of beryllium	nitride, Be	e3N2, was p	precisely measured
from 0 to 315 K; after judicious correctio			
led to the common thermodynamic properties	from 0 to 3	315 K, com	plementing earlier
NBS measurements from 273 to 1173 K. Prec	ise high-ter	nperature d	calorimetry gave the
heat capacity of tungsten from 1100 to 260	0 K (showing	g no detect	table difference
between two samples, one triple-zone-refin	ed and 99.99	9% pure);	finally corrected
results are given also for solid and liqui			
1200 to 2400 K. A mass-spectrometric inve			
identified the high-temperature gas specie			
of formation. As part of the current NBS	•		
of the alkali-hydroxide molecules, the hig			
isotopic varieties of RbOH were observed a			
like CsOH (studied earlier), RbOH is linea			
of large amplitude. Two reviews deal with		-	
compounds: One documents the inadequacy of			
fluorides and recommends a systematic cata			
consistency; while the other describes a n			
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