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Report No AFRPL-TR-66-220

INVESTIGATION OF THE THERMODYNAMIC PROPERTIES OF ROCKET EXHAUST PRODUCTS

Milton Farber, M. A. Greenbaum, Margaret A. Frisch, et al.

FINAL REPORT Contract AF 04(611)-10929 September 1966

Department of the Air Force Air Force Rocket Propulsion Laboratory (AFSC) Edwards, California 93523

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Rocket Power, Inc 2275 E Foothill Blvd Pasadena, California 91107

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Rocket Power, Inc. 2275 E. Foothill Blvd. Pasadena, California 91107

FOREWORD

This report presents details of the work performed under Contract AF 04(611)-10929 during the period August 1, 1965 through July 31, 1966 at the Rocket Power, Inc. Research Laboratories, 2275 E. Foothill Blvd., Pasadena, California 91107. The technical monitor of this program was Mr. Curtis C. Selph, RPCL, Air Force Rocket Propulsion Laboratory, Air Force Systems Command, Edwards, California.

This technical report has been reviewed and is approved.

W. H. Ebelke, Colonel, USAF Chief, Propel'ant Division

ABSTRACT

A thermodynamic research program for the past twelve months has resulted in thermal data for the high temperature stable species BeOH(g), HBO(g), $B_2O_3(g)$, HOBO(g), $AlCl_2(g)$ and $Al_2Cl_4(g)$. In addition, infrared spectral data have been obtained for Be(OH)₂(g).

SUMMARY OF THERMODYNAMIC DATA

BeOH(g)

| ∆н ^о _{f2235} °к | <u>2nd Law</u> -55.7 <mark>+</mark> 3.8 kcal/mole | <u>3rd Law</u> -56.7 ⁺ .5 kcal/mole |
|--|--|---|
| ∆н ^о _{€298} ° _№ | -46.8 ⁺ 3.8 kcal/mole | -48.2 + .5 kcal/mole |
| ^{8°} 2235 [°] к | 75.3 + 1.7 cal/deg/mole | |
| ^{5°} 298 [°] к | 54.0 $\frac{+}{-}$ 1.7 cal/deg/mole | 53.3 cal/deg/mole (JANAF) |
| ∆н ^о _{f298} °к | AlCl ₂ (g) | -66.0 ⁺ _3 kcal/mole |
| | $Al_2Cl_4(q)$ | |
| ∆н ^о f900 ⁰ к | -194.8 + 1.0 kcal/mole | |
| ∆ н ^о _{f298} °к | -194.0 + 5 kcal/mole | |
| s ^о 900 ⁰ к | 116 + 1.0 cal/deg/mole | |

 $s^{\circ}_{298} s^{\circ}_{K}$ 90 $\frac{+}{5}$ cal/deg/mole

HBO(a)

3rd Law

| 2nd Law | | |
|------------------------------|--|--|
| -51 ⁺ 3 kcal/mole | | |
| | | |

 $\Delta H^{0}_{f298^{0}K}$ -50⁺ 3 kcal/mole

| | B203(0) |
|------------------------|--------------------|
| ∆H _{v1320} °K | 86.3 - 2 kcal/mole |
| ∆H _{v298} °K | 93.6 2 kcal/mole |

HOBO(a)

| △H ⁰ f1320 ⁰ K | -135.9 + 2 kcal/mole |
|--|----------------------|
| ∆H ⁰ ₁₂₉₈ ° _K | -133.5 + 2 kcal/mole |

$\frac{Be(OH)_2(g)}{2}$

| Experimental in | nfrared active v | ibrational fre | quency: | 649 cm^{-1} |
|-----------------------------------|------------------|----------------|------------|-----------------------|
| Estimated vibra | tional frequence | cies, bent mo | del: | |
| | 3600 | 1100 | 600 | |
| | 3500 | 900 | 600 | |
| | 1600 | 700 | 300 | |
| ^{8°} 1705°K | 93.3 cal | /deg/mole (b | ent model) | |
| 8 ⁰ 298 ⁰ K | 60.6 cal | /deg/mole (b | ent model) | |

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. INTRODUCTION

The extensive research and development efforts in the area of high energy propellant systems, solid, liquid and hybrid, have led to consideration of an increasingly large number of compounds containing an ever-widening spectrum of the elements. The accurate theoretical evaluation of these new propellent systems in terms of specific impulse, range and similar parameters, requires a knowledge of the thermodynamic properties of the reactants (heat of formation, ΔH_f , is the prime requisite here) and the thermodynamic and physical properties of all the possible products. Among the values needed are heat of formation, entropy, heat capacity, melting point, boiling point, and heats of vaporization, fusion and sublimation. Since the temperatures generally encountered in rockets and missiles fall in the range of 2000-5000[°]K the necessary thermodynamic properties for the exhaust products must be known in this temperature range.

The existence of thermodynamic data, whether theoretical or experimental, in the range of $20-600^{\circ}$ K for a given compound is not sufficient to permit the accurate theoretical evaluation of a propellant system which, on combustion, might form this compound at temperatures of 3000° K. Extrapolation of thermodynamic data over these temperature spans, i.e., 600 to 3000° K, are usually inaccurate. Of even greater importance is the fact that a large number of species produced in the combustion of propellants do not even exist at lower temperatures and thus their properties can only be determined at the high temperatures where they exist. It is for these reasons that research programs have been undertaken to determine the physical and thermodynamic properties of those species expected to be formed by the combustion of propellants.

The current interest in the use of the light metals and their compounds (Li, Be, Mg, Al, B and their hydrides) as fuels or additives in solid or hybrid propellants, has prompted the inauguration of numerous research programs designed to determine the requisite thermodynamic and physical properties of compounds arising from interaction of the metals with the oxidizers employed (generally oxygen or fluorine compounds). Thus, species of prime interest are those arising from reaction of a metal ion with fluorine, oxygen, nitrogen or combinations of these atoms. During the past 12 months thermal data was obtained for the high temperature stable species, BeOH(g), HBO(g), AlCl₂(g) and Al₂Cl₄(g). In addition, infrared spectral data were obtained for Be(OH)₂(g). All thermodynamic values reported at 298°K as well as those at the specified temperatures were calculated using the latest JANAF Tables¹, i.e. those available as of July 31, 1966 except where indicated in the text.

II. THERMODYNAMIC PROPERTIES OF BeOH(g)

A. Introduction

The literature does not report a value for the heat of formation for the subhydroxide, BeOH(g). However, the current JANAF Tables¹ report a value of $-25 \frac{+}{-}10$ kcal/mole for ΔH_{f298} of BeOH(g). This value is based on a minimum value obtained from a mass spectrometric study² of the reaction

$$Be_2O(g) + H_2O(g) = 2BeOH(g)$$
 (1)

Prior estimates based on bond energy relationships were reported in preceding JANAF gray Tables³ as -41 ± 15 kcal/mole for ΔH_{f298} of BeOH(g). Because of these rather large uncertainties and the fact that no values for the thermodynamic properties of BeOH can be found in the literature, a study of the reaction between solid beryllium oxide and gaseous hydrogen was undertaken in this laboratory. The reaction was studied employing the molecular flow effusion method⁴ in the temperature range 2107° K to 2368° K.

B. <u>Experimental</u>

1. Apparatus

A high temperature carbon rod furnace was assembled in a high vacuum system. The rods were supported vertically by tantalum sheets (five layers at the top and six layers at the bottom). Alumina caps were used to insulate the rods from the tantalum sheets. The carbon rods were shielded by five cylindrical layers of 5 mil tantalum separated by $1/4^{"}$ graphite spacers. The inner and outer shields were $2-1/2^{"}$ and $4-1/2^{"}$ in diameter, respectively. Tantalum bolts and graphite spacers supported every shield. The entire assembly was supported from the brass plate base by three $1/4^{"}$ diameter stainless steel rods attached to the outer shield. The leads for the carbon rod heating unit were extended through the five shields and attached to two copper rods, whic's were connected to the insulated feedthroughs attached to the brass plate. A detailed diagram of the apparatus is presented in Fig. 1.

The reaction cell was a 2" long BeO tube closed at one end, with an o.d. of approximately 0.4", and an i.d. of approximately 0.3". A hole approximately 1 mm in diameter was drilled through the closed end of the tube. The exact dimensions of the cell hole diameter and thickness were measured with a traveling microscope. Two BeO cells with different hole dimensions (as given in Table 1) were employed for the series of experiments.



.

The cell was loosely covered with a rhenium cell which had a 1.5 diameter mm hole drilled through the closed end. This rhenium cell in turn was covered with a tantalum cell with a 1.5 mm orifice at the center of the closed end. The entire unit was inserted tightly into a tantalum holder which was fitted to the end of the 1/4" o.d. tantalum flow tube (c.f. Fig. 1). A rhenium foil insert with a 0.2 mm hole was placed over the tantalum tube inlet to prevent flow of species to the tantalum tube. The flow tube penetrated the bottom tantalum sheets of the furnace and a thermocouple fabricated from tungsten, 5% rhenium-tungsten, 26% rhenium wire was used as the temperature sensing element. This thermocouple was inserted approximately 1" inside the inner shield at the bottom of the furnace. The temperature of the cell was measured with an optical pyrometer through a window at the top of the furnace. This measurement was checked against a calibrated thermocouple which was placed inside the cell just below the effusion hole. This thermocouple was checked against a calibrated Pt-Pt, 10% Rh. thermocouple. The results showed that at the experimental temperature range the deviation was within 5 degrees of the standard NBS values. Experiments were performed at 30° intervals for the total of more than 30 performed over the 300° temperature range. The 30° differential temperatures were reproducible to $\frac{1}{2}$ degrees.

The entire furnace assembly was fitted into a 17" long stainless steel cross pipe with 10-1/2" dia. brass plates at the vertical ends and 4" dia. flanges at the side arms. One of the side arms was connected to the pumping system, the other to the ionization and thermocouple gauges. The temperature of the cell could also be measured with an optical pyrometer by viewing through a quartz plate fitted at the top end plate of the pipe. The pressure of the system was usually maintained in the region of 10^{-6} to 10^{-7} mm Hg.

An 8 liter tank was filled with hydrogen (99.999% ultra pure grade) at the commencement of each H_2 experiment. A calibrated gauge was connected to the tank to permit calculation of the amount of hydrogen used during the run. A metering value was connected to the line immediately prior to the hydrogen entering the flow tube. Approximately the same value reading was used for each run to insure that the amount of hydrogen used would be fairly constant.

A schematic diagram of the entire apparatus is presented in Fig. 2 <u>Procedure</u>

In the temperature range of these experiments BeO has an appreciable weight loss due to vaporization, therefore, a blank run in which no

2.



hydrogen was passed through was performed at each temperature to determine the weight loss of BeO due to vaporization alone. The BeO cell was weighed and the reaction assembly was placed in the system. The system was pumped down to the desired pressure, heating was commenced and a constant temperature was maintained at a desired value. The blank or hydrogen experiment was then begun. Upon completion of the experiment, the system was allowed to cool down to near room temperature and the cell was removed and reweighed. Initial and final readings of the pressure in the hydrogen tank of calibrated volume were recorded. In this manner the number of moles of $H_2(g)$ used in each run was calculated.

C. Discussion

After a considerable number of preliminary experiments were performed to establish that a reaction was indeed occurring between H_2 and BeO, a series of over 30 experiments was made with two effusion cells over a 300° temperature range. Experiments below 2100°K yielded weight losses too small from which to obtain accurate data. Consequently, the experimental temperature range was chosen as 2107 to 2368°K. The experiments were performed at 30° temperature intervals and in random fashion. Blank runs were made before and after each H_2 -BeO reaction experiment. The data for these experiments are presented in Table II.

Although the two cells had different orifice dimensions, a further check for the establishment of equilibrium was made using BeO chips inside the BeO cell. The weight loss results for these experiments were essentially the same as those for the empty cells. Thus, within the experimental error, the surface area of several sq. cm. of the cells was large enough (i.e., reaction area to orifice area greater than 1000, to establish equilibrium with the hydrogen.

Having established that H_2 is reacting with the BeO, the treatment of the data requires a knowledge of the several equilibria that may be involved. The primary reaction for the formation of BeOH employing H_2 is

 $BeO(c) + 1/2H_2(g) = BeOH(g)$ (2)

Also to be considered are the reactions

 $BeO(c) + H_2(g) = Be(g) + H_2O(g)$ (3)

$$2BeO(c) + H_2(g) = Be_2O(g) + H_2O$$
(4)

and the dissociation of hydrogen

$$1/2H_{2}(g) = H(g)$$
 (5)

Of secondary importance is the reaction to form the hydroxide from the water formed as a result of reaction (3)

$$BeO(c) + H_2O(g) = Be(OH)_2(g)$$
 (6)

The measured quantities for an experiment included the weight loss of BeO and the number of moles of elemental hydrogen employed in the reaction. The total number of moles of H_2 , $n_{T(H_2)}$ entering the reaction zone is distributed over the reactions (2), (3) and (5) as

After consideration for the weight loss of BeO due to vaporization, the remainder of the weight loss of BeO could be attributed to reactions (2), (3), (4) and (6). The weight loss that would be attributed to reaction (6) as a result of the water formed in reaction (3) was negligible in the temperature range studied.⁵ Calculations also showed that the contribution due to reaction (4) was negligible. The value reported for the vaporization of liquid beryllium (ΔH_v of Be = 75.38 at 298°K kcal/mole) in the JANAF Tables is based on several recent measurements.⁶⁻¹⁰ Employing this value calculations were made for the weight loss of BeO resulting from reaction (3). The remaining weight loss of BeO was attributed to the formation of BeOH. In all experiments the total pressure inside the cell ranged from approximately 0.4 to 0.8 mm. Mean free path calculations in the experimental temperature range indicate that for the cell dimensions the flow for this pressure range is molecular. Therefore, all calculations were based on the modified Knudsen equation

where P is pressure in mm of Hg for the species inside the cell.

G is the weight, in grams, of the gas species escaping through the effusion hole

 W_0 is the Clausing factor. The dimensions of the cell orifice are given in Table I from which the Clausing factor was calculated as given in Dushman.¹¹

A is the area, in cm^2 , of the effusion hole

t is the total time of reaction, in seconds

T is temperature in ^OK

M is the molecular weight of the species

The contribution to the weight loss due to reaction (3) was approximately 10 %. Although the vaporization data for Be is quite definitive, it was decided to perform a further check to ascertain that the major reaction was not reaction (3). This check is based on the pronounced effect on reaction (3) which can be brought about by adding a small amount of water vapor to the H₂. A calculation showed that at 2235° K and a total pressure of 0.7 mm of Hg the addition of 3 mole percent water would cause a 75% reduction in weight loss, if this weight loss were due primarily to reaction (3). Such a reduction would occur despite the simultaneous enhancement of reaction (6). This water percentage represents an optimum since at higher percentages the ascendancy of reaction (6) begins to outweigh the reduction in reaction (3).

Thus for a 110 min. run at 2235°K and 0.7 mm total pressure, calculations based on JANAF data¹ for the formation of Be and water as a result of the reaction of hydrogen and BeO would predict a weight loss of 0.74 mg of BeO, while the 3 mole percent water and 97 mole percent H_2 mixture experiment would predict a weight loss of 0.2 mg of BeO. Since the total weight loss in the pure hydrogen experiments at this temperature ranged between 8 and 9 mg, a reduction of 6 to 7 mg would be expected if Be were the chief product formed. However, if the JANAF data for the Be correction were fairly accurate, a negligible change would be expected in the water experiments. Two water experiments (3 mole percent) were performed. The weight losses were 9.6 and 10.1 mg respectively, while the pure hydrogen experiments recorded weight losses of 8.1 and 8.8 mg, respectively.

(8)

Although a decrease of 0.5 mg was theoretically predicted, an actual increase of from 0.8 to 2 mg occurred. An examination of the 30 experiments performed in a 300° temperature range, shows that this discrepancy between the actual change and the predicted change of weight losses due to the addition of H₂O is within the scatter of the pure hydrogen experiments, i.e. 1 to 2 mg reproducibility at a given temperature.

The results of the water experiments indicated that use of the published value for the heat of vaporization of Be would be valid. Therefore, a weight loss of BeO was calculated for reaction (3) by means of the Knudsen equation (8) and the JANAF tables for the thermal data. The remaining weight loss of BeO was assumed to be due to the formation of BeOH. The partial pressure of H₂ shown in column 7 of Table III was calculated from the value of n_{H_2} obtained from equation (7). The partial pressure of BeOH(g) was then calculated and an equilibrium constant was obtained for reaction (2). These calculations are presented in Table III.

A plot of the experimental data in Table III in the form of $\log K_2$ vs 1/T is presented in Fig. 3. A least squares analysis of the data yields a value for the slope of the line which corresponds to 88.3 ± 3.8 kcal for ΔH_r at the average temperature of 2235° K. Similarly, a value for $\Delta S_{r2235} \circ_{\rm K}$ of 27.4 ± 1.7 cal/deg is obtained from the intercept of the curve. Employing the available thermodynamic values and thermal functions for BeO(c), $H_2(g)$ and thermal functions for BeOH(g)¹, the ΔH_{f298}° and S_{298}° for BeOH(g) are found to be -46.8 ± 3.8 kcal/mole and 54.0 ± 1.7 cal/deg/mole, respectively. The corresponding average 3rd Law value for the heat of reaction, $\Delta H_{r2235} \circ_{\rm K}$, is 87.2 ± 0.5 kcal. The free energy and heat of reaction for the 24 experiments are listed in Table IV. The Third Law ΔH_{f298} for BeOH(g) is calculated as -48.2 ± 0.5 kcal/mole. The entropy of BeOH(g) listed in the JANAF Tables is 53.29 cal/deg/mole. A summary of the thermodynamic functions for BeOH(g) is presented in Table V.

D. <u>Summary</u>

The excellent Second and Third Law agreement, supported by the experiments performed employing a mixture of hydrogen and water, establish the validity of the reaction of BeO with H_2 to produce BeOH. The recommended values for heat of formation and entropy of BeOH(g) at 298°K are -47.0 + 2 kcal/mole and 54.0 + 1 cal/deg/mole respectively.



THERMODYNAMIC PROPERTIES OF AIC1, AND ALC1

. Introduction

DT.

Several values have been published in the JANAF Tables for the heat of formation of AIC1 as a result of the reaction

$$AlCl_{3}(g) + 2Al(l) = 3AlCl(g)$$
 (1)

which has been studied by means of transpiration experiments. A recent study of this reaction in this laboratory¹² by means of molecular flow effusion has essentially confirmed these values. No data appears in the published literature for either $AlCl_2$ or its dimer Al_2Cl_4 . On the assumption made in the JANAF tables that the heat of reaction for

$$AlCl_{2}(g) + AlCl(g) = 2AlCl_{2}(g)$$
(2)

would be essentially zero, a value for ΔH_{f298} for AlCl₂ has been estimated as -75⁺ 20 kcal/mole. Based on this estimate it would appear that a high percentage of AlCl₂ would be produced in a reaction of aluminum chloride vapor and aluminum in the temperature range from 700 to 1,000^oK. Also, if the heat of dimerization

$$2AlCl_2(g) = Al_2Cl_4(g)$$
(3)

were approximately 50 kcal a considerable amount of the dimer would also be present in this same temperature range. There is no table for $Al_2Cl_4(g)$ in the JANAF Tables.¹

Therefore, in order to obtain thermal data for $AlCl_2$ and its dimer Al_2Cl_4 a number of transpiration experiments were performed at atmospheric pressure in the temperature range 800 to 1000° K. These experiments involved the use of argon as the carrier gas for aluminum chloride vapor and was passed over pure aluminum, both solid and liquid.

B. <u>Experimental</u>

1. Apparatus

The experimental apparatus employed in this study consisted of a dual heating system for: (1) generating aluminum chloride vapor, and (2) heating the cell containing aluminum to the desired transpiration temperature. A schematic diagram of the entire apparatus and a detailed diagram of the transpiration cell are shown in Figs. 4 and 5.





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0

1/2" Al₂O₃ Outlet Tube Fig. 5 Al - AlCl₃ Reaction Cell Assembly

The gaseous aluminum chloride was generated by heating the highly purified $AlCl_3(c)$ in a 250 ml glass flask by means of a heating mantle. A saturable core reactor-recorder combination was used to monitor and control the temperature of the flask. The flow rate of the gaseous $AlCl_3$ was controlled by controlling the temperature of the $AlCl_3$ generator. The flow rate of high purity argon was adjusted and measured employing very fine metering values and an accurately calibrated flow meter and was used to carry the $AlCl_3$ into the reaction zone.

The furnace was a heavily insulated 12" long by 1-1/4" diameter nichrome resistance type. The furnace tube was constructed of aluminum oxide and measured " i.d. Each end of the furnace tube was closed by a monel flange, which also provided the openings for the gas delivery and exit tubes, as well as for the chromelalumel thermocouple. The temperature of the cell was held constant by means of a silicon controlled rectifier-recorder combination.

The aluminum was contained in an aluminum oxide boat placed inside an aluminum oxide tube 10 mm i.d. The orifice and inlet hole of the reaction cell measured 1.5 mm in diameter, preventing any significant back diffusional losses in the reaction cell. The tube admitting the $AlCl_3$, argon mixture to the cell was also constructed of aluminum oxide.

The aluminum chloride in the argon-aluminum chloride vapor mixture was collected in a 250 ml flask after leaving the reaction zone. The aluminum chloride generator and collection flasks were connected to the transpiration apparatus with teflon and monel fittings. These fittings were heated with heating tapes and four immersion heaters were attached to the flanges in the ends of the furnace tube to prevent condensation of the aluminum chloride vapor entering or leaving the reaction zone. The temperature of these heating tapes and immersion heaters was maintained at approximately 230° C by the use of variable rheostats.

2. Procedure

The commerically obtained aluminum chloride (anhydrous) of 99.5% purity was further purified by sublimation prior to its use. The sublimation procedure also converted traces of moisture into HCl(g) which would be driven off as a vapor. As a further precaution against impurities, prior to each run approximately 1 gram of 99.9%, 200 mesh, aluminum powder was added to the sublimated aluminum chloride and thoroughly mixed by vigorous agitation for use in the generator flzsk.

The procedure employed for all experimental runs consisted of heating the sample of Al to the desired temperature under a stream of pure argon (99.998% purity). After the desired temperature was reached and automatic control established, the AlCl₃ boiler was heated rapidly to its predatermined temperature while argon was swept through it. At the desired temperature automatic control was established and the runs carried out for periods of 1-7 hours. Weight loss of the Al samples was established by weighing before and after each run. The amount of AlCl₃ passed over the sample was determined by weighing the generator before and after each run. After leaving the reaction the AlCl₃ in the gas mixture was collected in a second flask and weighed. (This weight gain was essentially equal to the weight loss in the first flask).

The weight loss of aluminum due to vaporization by passing argon alone over the aluminum sample was found to be insignificant and in agreement with that predicted from the published vapor pressure of aluminum. To further check that no HCl or water vapor was present in the aluminum chloride generator flask argon gas was passed through the generator, which was heated to 50° C, and then passed over a sample of aluminum in the reaction cell, which was heated to 900° K. At this low generator temperature the quantity of AlCl₃ in the vapor state is very small and therefore the resulting quantity of aluminum lost due to the reaction with AlCl₃ would also be insignificant. Thus any large losses of aluminum would have to be attributed to the entrained HCl(g) which reacts quantitatively with aluminum at the experimental temperatures. Since no reaction was observed it can be concluded that the reaction taking place is due to AlCl₃ and not to impurities.

C. Discussion

The obtaining of the requisite thermal data required an analysis of a series of 33 transpiration experiments in the temperature range of 800 to 1000° K and at 25[°] temperature intervals. The flow rates were in the range 15 to 25 cc/min which were sufficient to establish equilibrium of the vapor species above the aluminum. The orifice in the cell prevented any appreciable back diffusion. The experimental data including AlCl₃ and Al weight losses and flow rates are presented in Table VI.

Four transpiration studies 13,14,15,16 and a spectrographic study 17 gave results of the heat of reaction (1) in good agreement with those of the effusion experiments previously performed in this laboratory 12 for the reaction in the temperature range 930 - 1034°K. The equilibrium constants obtained from the

effusion data were chosen as the basis of correcting the total weight loss of Al for this reaction, since at low pressure $\sim 10^{-5}$ atm., the dissociation of higher chlorides to AlCl would be more likely. Also, the dimerization equilibrium

$$Al_2Cl_6(g) = 2AlCl_3(g) \tag{4}$$

was employed in correcting the AlCl₃ flow data.

The remainder of the Al weight loss could then be distributed between $AlCl_2$ and its dimer Al_2Cl_4 . The appropriate reactions would be

 $2A1C1_{3}(g) + A1(1, s) = 3A1C1_{2}(g)$ (5)

and

$$4AlCl_{3}(g) + 2Al(1, s) = 3Al_{2}Cl_{4}(g)$$
 (6)

The partial pressures involved for the necessary equilibria were calculated from the weight loss data by means of the gas law equation,

$$P = \frac{nRT_o}{V_o}$$
(7)

where

n is the number of moles of the specie formed from liquid or solid Al reaction in unit time with a given volume V_0 of transpiring gas at a pressure of one atmosphere.

This gas includes all species, i.e., A, $AlCl_3(g)$, $Al_2Cl_6(g)$ and the products formed. T₀ : the ambient temperature. Thus the partial pressure of AlCl in atmospheres would be

$$P_{A1C1} = \frac{n_{A1C1}}{n_{T}}$$
(8)

where n_T is the total moles of gas in the transpiration mixture. A preliminary plot of the equilibrium constant for reaction (6) vs 1/T based on the assumption that reaction (5) was negligible showed extreme curvature indicating a possible mixture of the two species AlCl₂ and Al₂Cl₄. (See Fig. 6) Therefore, an iterative process was adopted for the solution of equations (5) and (6). After calculating a partial



pressure of the AlCl(g) from equation (1) to obtain a weight loss attributed to reaction (1), the remainder of the weight loss was distributed by means of an interative procedure between both AlCl₂ and Al₂Cl₄.

The iteration was based on the assumption that for the proper distribution of weight loss between the two species straight lines would occur for the slopes representing the ΔH_r for the reactions represented by equations (5) and (6).

The iteration was begun with the assumption that the entropy listed in the JANAF Tables¹ for the triatomic molecule $AlCl_2$ was sufficiently accurate (within 3 eu) to allow free energy calculations for reaction (5), based on assumed values for the heat of formation of $AlCl_2(g)$. After each iteration corrections were made for the Al weight loss due to $AlCl_2$ and the free energy for reaction (6) was calculated.

Table VII presents the partial pressures and weight losses calculated for the species involved in the various equilibria. The equilibrium constants K_1 for reaction (1) are from the effusion experiments¹². K_4 represents the equilibrium constant for the dimerization reaction (4) and are taken from the JANAF Tables¹. K_5 is the equilibrium constant for reaction (5) based on a value -66 kcal/mole for $\Delta H_f AlCi_2(g)$ at 298°K and on specific heats and entropies from the JANAF Tables¹. Finally, K_6 is the equilibrium constant for reaction (6) after corrections were completed for reactions (1) and (5).

A fairly straight slope was obtained for reaction (6) (see Fig. 7) based on an assumed value of $-66 \stackrel{+}{=} 1 \text{ kcal for } \Delta H_{f298} \text{ for AlCl}_2(g)$. The ΔH_r and ΔS_r of reaction (6) were found to be $-25.9 \stackrel{+}{=} 2.9 \text{ kcal and } -64.2 \stackrel{+}{=} 3 \text{ cal/deg/mole}$ respectively at 900°K. This yields values of $-194.8 \stackrel{+}{=} 1.0 \text{ kcal/mole}$ and 115.7 $\stackrel{+}{=} 1.1 \text{ cal/deg/mole}$ for ΔH_{f900} and S^{O}_{900} for Al₂Cl₄(g). A corresponding third law heat of reaction could not be calculated since no value for the entropy of Al₂Cl₄(g) is reported. However, the second law entropy of 116 cal/deg/mcle is a reasonable value for a molecule of this size and structure and may be compared to the estimated value of 115 cal/deg/mole (with an uncertainty of 5 e.u.) for $B_2Cl_4(g)$ listed in the JANAF Tables.¹

Since heat capacities have not been estimated for $Al_2Cl_4(g)$ calculations of the heat of formation and entropy at 298°K are speculative. However, a value for ΔH_{f298} of -194 $\frac{+}{2}$ 5 kcal/mole and S°₂₉₈ of 90 $\frac{+}{2}$ 5 cal/deg/mole are considered raliable.





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The heats and entropy of dimerization for reaction (3) are -62 kcal and -47 cal/deg/mole at 290° K, respectively. This compares to dimerization values for B_2Cl_4 of -77 kcal and -44 cal/deg/mole for the heat and entropy respectively. The thermodynamic functions for $AlCl_2(g)$ and $Al_2Cl_4(g)$ are presented in Table VIII.

Having obtained a fairly accurate value for ΔH_{f} of AlCl₂ it was possible to obtain a check on the assumption of equation (1) for the effusion data. In the temperature range of the effusion experiments 12 for reaction (1) a $\Delta H_{\rm r}$ of 95 kcal/mole was obtained. Employing a value of -66 kcal/mole for ΔH_{f298} of AlCl₂(g) a ΔH_r at an average temperature of 900°K for reaction (5) of 86 kcal is calculated. Since the slopes for equations (1) and (5) differ by only 9 kcal or 10%, curvature in reaction (1) would not be readily seen unless reaction (5) were contributing more than 10% to the Al weight loss. Therefore, in order to check the validity of the effusion experiments in the temperature range 850°K to 1000°K free energies (extrapolated from previous results at this laboratory¹²) were calculated for equation (2) at the highest and lowest temperatures, $850^{\circ}K$ and 1000⁰K. The calculations are presented in Table IX. Since the amounts of $AlCl_2(g)$ in equilibrium with AlCl and $AlCl_3$ range from only 5 to 8% over this temperature range the validity of the effusion experiments are further established. However, it would appear that at atmospheric pressure transpiration experiments would involve considerable quantities of both AlCl and AlCl₂.

D. <u>Summary</u>

An analysis of the effusion and the transpiration experiments to obtain thermal data for AlCl(g), $AlCl_2(g)$ and $Al_2Cl_4(g)$ shows that they are self-consistent. Thus, the recommended values for ΔH_{f298} for $AlCl_2(g)$ and $Al_2Cl_4(g)$ are $-66 \stackrel{+}{-} 3$ and $-194 \stackrel{+}{-} 5$ kcal/mole respectively. The recommended entropy S^{o}_{298} of $Al_2Cl_4(g)$ is $90 \stackrel{+}{-} 5$ cal/deg/ mole.

• THERMODYNAMIC PROPERTIES OF HBO(g)

A. Introduction

Ion intensities have been reported for the specie HBO(g) in mass spectrometric studies¹⁸ involving B_2O_3 and $H_2O(g)$. However, no thermal data have been published. The current JANAF Tables¹ report an estimated value of -20 \pm 20 kcal/mole while the previous gray Tables³ reported a value of -47 \pm 15 kcal/mole based on a National Bureau of Standards estimate.

A mass spectrometric study over the temperature range $1160 - 1405^{\circ}$ K has been made for the reaction of $B_2O_3(1)$ and $D_2(g)$ to obtain a heat of formation of the high temperature stable specie DBO. Initial experiments were performed on the reaction of B_2O_3 with H_2 which indicated the formation of HBO; however, the accuracy of the mass spectrometer intensities was impaired by the high background at mass 28, where the background intensity was 30 times larger than the background of mass 29. Since most compounds of deuterium have heats of formation nearly equal (within 1 kcal) of those of hydrogen it became apparent that a more precise value for HBO could be obtained by employing deuterium.

B. Experimental

The experimental procedure employed for the studies combines a high vacuum high temperature apparatus together with a quadrupole mass spectrometer. The vacuum furnace is similar to the one used for the BeOH experiments.

A graphite furnace mounted in a high vacuum chamber was constructed and tested for the purposes of this investigation. The furnace is capable of maintaining temper tures in excess of 2300° K, for long periods of time. A high vacuum system is necessary to keep the background or residual gas pressure low enough to obtain high sensitivities of the species produced. The vacuum system employed in the apparatus was capable of providing, with the aid of a cold trap, a vacuum of better than 1 x 10⁻⁷ mm Hg, at a cell temperature of 1800°K in the furnace, while at room temperature the pressure was 2 x 10⁻⁸ mm Hg.

Details of the various components of the experimental apparatus and their operating procedures are described in the following section.

1. <u>Apparatus</u> a. Vac

Vacuum System

The furnace chamber was fabricat \ldots in a cross configuration using 304L stainless steel 8" and 4" schedule 10 pipe and 3/8" thick plate. All seams were heli-arc welded on the inside to minimize the outgassing load. The quadrupole mass spectrometer probe was mounted on one of the 8" pipe plate flanges. The other 8" flange supported the furnace assembly, which is described in the following section. The cross was attached to the vacuum pump via one of the 4" branches while the other was sealed by a plate flange containing a 2" quartz window for visual temperature measurement. Only viton O-rings or teflon gaskets have been used for seals throughout the entire system. The cross section drawing of the chamber may be seen in Fig. 8 and photographs of the assembly are shown in Figs. 9 & 10.

During furnace operation, the 8" pipe was cooled with freon which was passed through a coil of 3/8" copper tubing soldered to the outside diameter of the cross while each plate flange of the 8" pipe was air cooled using high capacity blowers.

The furnace chamber was evacuated by a 4" high speed NRC diffusion pump (750 1/sec), charged with Dow Corning 705 oil. The diffusion pump is backed by a Welch Duo-Seal two stage fore pump (100 1/min). The back streaming of the oil vapor was essentially eliminated by the water cooled cap of the diffusion pump and the inclusion in the system of a NRC expanded water cooled chevron baffle. A 4" thimble cold trap was included in the system.

b. <u>Furnace</u>

The heating element of the furnace consisted of twelve 6" long x 1/8" o.d. high density graphite rods (resistivity 1500 x 10^{-6} ohm cm⁻¹) arranged in series and had a resistance of 2.9 ohms at 20° C. The rods were held 0.386" apart between centers in 1/4" o.d. graphite couplers. This distance is equivalent to a 30° chord of a 1.491" diameter circle. The heater rods were kept parallel to one another by additional 1/8" o.d. graphite rods placed in the center of each coupler. These support rods passed through 1/4" holes in the end shields and were capped at the ends with alumina tubes closed at one end (1/4" o.d. x 1/8" i.d. x 1" long). This arrangement allowed the positioning of the graphite support rods accurately in the end shields and the overall result was an extremely rigid furnace element. Figure 11 shows in detail the coupling arrangement between two graphite rods while in Fig. 12 the support rod relationship to the heater element and alumina spacer can be seen.

Six five mil tantalum cylinders, 2-1/2" to 5" in diameter were used as the radiation shields. The lengths of the shields began at 8" and increased in 1/2" increments to 10-1/2" so that with the end shields in place





Fig. 9. Mass Spectrometer and the 4 Inch Vacuum System.



Fig.10. Furnace Port. The Reaction Cell and Flow System Sub-Assembly in Center.



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Fig. 11. Close Up View of Graphite Coupler used in Furnace Element



Fig. 12. View Showing Alignment Rod With Alumina Insulator

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the shields were a set of nested cylinders 1/4" apart. Tantalum rods 1/8" diameter, spaced 120° apart around both ends, and 1/4" o.d. x 0.245" long graphite spacers kept the shields concentric and rigid. These tantalum rods were bolted to three 1/4" o.d. stainless steel rods which supported the shield system 4" off the flange plate. Fig. 13 shows a top view of the shields with the furnace element in place. The complete furnace assembly is shown in Fig. 14.

In each of the top end radiation shields, there was a 1/2" hole which was coaxial with the mass spectrometer quadrupole analyzer. The cylindrical shields contained a set of 1/4" holes which were coaxial with the viewing port on the vacuum chamber. This hole provided a reasonable approximation to z black body cavity for visual temperature measurement.

c. <u>Temperature Measurement and Control</u>

The temperature was monitored by a tungsten-5% rhenium versus tungsten-26% rhenium thermocouple, mounted in the furnace just outside the heating zone. The thermocouple was calibrated against a 0.1% platinum versus platinum-10% Rhodium thermocouple at 30° intervals over the temperature range of the study. The tungsten-rhenium thermocouple was measured on a Leeds and Northrup K-3 potentiometer.

The K-3 bridge was set at the desired voltage and the imbalance plotted on a Leeds and Northrup Speedomax W recording potentiometer (center zero, 1 mV full scale). The control current to the 100 amp 230 volt Fincor SCR power supply was fed from a Leeds and Northrup Series 60 current adjusting type control system, acting on the information from the recording potentiometer. The sensitivity of the control loop at 1400° K. was about 0.1 to 0.2°C.

d. <u>Reaction Cell and Flow System</u>

The reaction cell and flow system were mounted on a separate plate which was sealed to the main flange plate via an O-ring. This arrangement (as seen in Fig. 15) facilitates the maintenance of the reaction system without disturbing the furnace assembly. A 1" hole in each of the bottom end shields permitted the assembly to pass into the furnace. Mounted on the 1/4" o.d. by 0.010" wall tantalum flow tube was a set of 5 mil tantalum 1" discs which sealed the hole in the radiation shields.

The reaction cell, fabricated from platinum, was 5 cm long by 1.27 cm o.d. by 1.12 cm i.d. At the closed end of the cell an exit tube 2.5 cm long by .25 cm o.d. by 10 cm i.d. was welded. The molten B_2O_3 was held in a



Fig. 13. Top View of Furnace Shields with Furnace Element in Place



Fig. 14. Complete Furnace Assembly Mounted on 8" Plate Flange




Fig. 15. View of Sample Port Assembly Showing Relationship of Flow Tube, Reaction Cell, Shields and Thermocouple

platinum boat 4.5 cm long and 0.5 cm deep. The reaction cell was joined to the 1/4 inch o.d. by 230 inch i.d. tantalum flow tube via a tantalum holder. The i.d. of the holder was machined .003" larger than 'he o.d. of the cell to allow 1 mil platinum foil to be placed around the cell and inserted into the holder for a distance of 1/8". The closeness of fit allowed moderately high pressures of D_2 gas to be used (e.g. 10mm) if desired.

The D_2 gas was fed to the cell from a two liter reservoir initially filled to a pressure of 0.1 atm via a Granville Phillips variable leak. Based on the Knudsen equation, the flow of D_2 gas through the orifice of the platinum cell at 1400°K and 10⁻⁴ atm. is approximately 2.0 cc hr⁻¹ at STP. Thus after one hour only a negligible change (1%) in the cell pressure occurs. The gas pressure was measured by a Veeco Dv-1 thermocouple gauge, calibrated for D_2 gas.

Quadrupole Mass Spectrometer

The quadrupole mass spectrometer used in this study, Model QUAD 200 (manufactured by Electronic Associates, Inc.) facilitates its installation in small vacuum systems, without sacrificing loss in sensitivity and resolution. This instrument has a resolution of 500 and a sensitivity of 10^{-14} to 10^{-15} torr for N₂, when an electron multiplier is used in conjunction with an electrometer amplifier. The quadrupole probe, factory mounted on a 4" Ultek flange, projects approximately 9" into the vacuum system (Figs. 16 & 17). The probe can be accurately aligned with the molecular beam from the reaction cell by sighting down the flow tube and adjusting the angle of the mass spectromete, with respect to the flange via a 3-1/2" i.d. bellows. The molecular beam can be intercepted to distinguish it from background by rotating a tantalum sectur directly in front of the furnace slits (Fig. 18). The vacuum seal used for the rotating shaft is a simple Swagelock device using a teflon front ferrule as the seal. There was no detectible air leak during rotation.

C. Discussion of Results

The intensities of the four principal species, D_2O , DBO, DOBO and B_2O_3 were measured with the quadrupole mass spectrometer at a series of temperatures covering a range of $250^{\circ}K$ (1160-1405°K). Twelve series of experiments were made encompassing a fifteen-fold change in the D_2 pressure (18-240 μ).

For the prime purpose of obtaining the heat of formation of DBO, only



Fig. 16. Quadrupole Probe



Fig. 17. Probe Mounted on Flange



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Fig. 18. Alignment Device with Shutters

the following three reactions were considered.

$$D_{2}(g) + B_{2}O_{3}(1) = DBO(g) + DOBO(g)$$
(1)

$$K_{1} = (^{I}DBO) (^{I}DOBO)$$

$$D_{2}O(g) + B_{2}O_{3}(1) = 2DOBO(g)$$
(2)

$$K_{2} = \frac{(^{I}DOBO)^{2}}{(^{I}D_{2}O)}$$

$$B_2O_3(1) = B_2O_3(g)$$
 (3)

 $K_3 = {}^{I}B_2O_3$

Since the $D_2(g)$ was maintained at a constant pressure its intensity was not included in K_1 . This assumption is not valid at the highest temperatures of the study where there is a significant decrease (> 5%) of the D_2 pressure in the cell due to reaction, therefore, these points have been ignored.

A typical set of data obtained during a run is given in Table X and the plots of the log K for reactions (1) through (3) as a function of 1/T are shown in Fig. 19. Ion intensities for species obtained at higher temperatures are shown in Table XI for a pressure of 240 μ and at a temperature of 1518⁰K.

The mass spectra measurements were all made at an electron energy of 70 volts and an emission current of 1 ma. A study was also made at 16 volts electron energy and 1 ma emission current at 75 μ D₂ pressure, but the data for reactions (1), (2) and (3) were essentially equal to those at 70 volts (deviation in the slope of less than 2 kcal) indicating no significant contribution of fragment species to the major peak intensities. Table XII summarizes the data obtained from nine determinations on reactions (1), (2) and (3). The heats of reaction were derived from least squares treatment of the data.



1.
$$\Delta H_v \text{ of } \tilde{b}_2 O_3(1)$$

The six runs obtained in this study agree with one another within $\frac{1}{2}$ kcal/mole. The average value for the second law ΔH^{0}_{V} at the mean temperature of 1320[°]K is 86.3 kcal/mole with an average deviation of $\frac{1}{2}$ 1.5 kcal/ mole. This value is in good agreement with most of the determinations. However, Buchler and Berkowitz-Mattuck¹⁹ report 93.6 $\frac{1}{2}$ 3 kcal/mole at 1300[°]K by mass spectrometry and Hildenbrand, Hall and Potter²⁰ 93.3 $\frac{1}{2}$ at 1500[°]K by Knudsen Vapor pressure measurements.

The reduction of ΔH_{v1320}^{o} to 298°K gives 93.6⁺ 2 kcal/ mole. This value compares favorably with 95.2⁺ 09 obtained by M.D. Sheer²¹; 90.5⁺ 1.5 from mass spectra measurements of White, Walsh, Goldstein and Dever²²; 93.1⁺ 0.7 given by Sommer²³; 92.8⁺ 1.4 obtained by Blackburn²⁴. Margrave in a recent article²⁵ quescions the validity of the current JANAF Table of B₂O₃(g) since he calculates 92⁺ 3 kcal/mole for ΔH_{v298} . His third law ΔH_{v298} is 104 kcal/ mole, which is in serious disagreement.

2. ΔH_{r}^{O} of $B_{2}O_{3}(1) + D_{2}O(g) = 2DOBO(g)$

The data obtained for this reaction was reproducible for the six series. The average value for the second law ΔH_{1320} is 81.5 kcal/mole with an average deviation $\stackrel{+}{-}$ 1.5 kcal/mole. This value is in excellent agreement with the mass spectra data of Meschi. Chupka and Berkowitz²⁶. They reported 84.6 $\stackrel{+}{-}$ 4 kcal/mole for the ΔH_{1250} of reaction (2). White, Mann, Walsh and Sommer²⁷ obtained 78 $\stackrel{+}{-}$ 5 kcal/mole from the variation of the 2030 cm⁻¹ band intensity of HOBO with temperature. Farber et al.²⁸ from measurements by the effusion method reported 80.8 $\stackrel{+}{-}$ 1.2 kcal/mole. The resultant ΔH_{f1320} of HOBO(g) is -135.9 $\stackrel{+}{-}$ 2 kcal/mole which reduces at 298°K to -133.5 $\stackrel{+}{-}$ 2 kcal/mole.

The data on reaction (1) gives an average second law $\triangle H_{f}^{o}$ at 1260°K for DBO(g) of -52 $\frac{+}{-}$ 3 kcal/mole and for HBO(g), -51 $\frac{+}{-}$ 3 kcal/mole. The high pressure runs at 240 μ D₂ pressure were discarded. The reduction of $\triangle H_{f}^{o}$ to 298°K for DBO(g) with the JANAF data gives -50.8 kcal/mole. Since the $\triangle H_{f}^{o}$ for most deuterium compounds is about 1 kcal/mole more negative, the $\triangle H_{f298}^{o}$ for HBO(g) is assigned a value of -50 $\frac{+}{-}$ 3 kcal/mole.

The JANAF¹ estimate of -20 kcal/mole for $\triangle H_{f298}$ is in serious disagreement with this experimental measurement; however, the previous estimate by W. H. Evans of -47 kcal/mole reported in the earlier Iterim JANAF Table³ is in remarkable agreement with the value obtained in this study.

In order to obtain a second and third law correlation, the free energy of reaction

 $D_2(g) + B_2O_3(g) = DBO(g) + DOBO(g)$ (4)

at the temperature of 1400°K was combined with thermal functions of the reactants and products. The equilibrium constant for this reaction is approximately 0.2 within a factor of 5 at 1400°K. Since Margrave²⁵ had pointed out that a considerable discrepancy appears in the thermal functions for both liquid and gaseous B_2O_3 , calculations were made with entropies listed in the latest JANAF Tables¹ and those in the earlier JANAF grey tables²⁹. The entropy in the grey tables at 1400°K for $B_2O_3(g)$ is 92.9 e.u. while the value in the most recent JANAF Tables¹ is 99.5 e.u. Employing S_{1400} cf $B_2O_3(g)$ of 92.9 cal/deg/mole a third law value of 17.3 kcal/mole for the heat of reaction (4) was obtained. This compares with 18.5 kcal/mole, for the average ΔH_r derived from the six experimental second law heat of reaction.

The $\triangle H_r$ which is obtained with the entropy value of 99.5 e.u. for $B_2O_3(g)$ yields a $\triangle H_r$ of only 8.1 kcal which is considerably lower than the second law value. Since Margrave also found a discrepancy of 12 kcal in the vaporization data of B_2O_3 between his second and third law values, it is likely that the JANAF entropy data for B_2O_3 is in error. The estimated entropies for HBO and HOBO kury also be in error by as much as 2 to 3 entropy units (total) contributing from 3 to 4 kcal error in the $\triangle H_r$.

THERMODYNAMIC PROPERTIES OF $Be(OH)_2(g)$

A. Introduction

The heat of formation of Be(OH)₂(g) ($\Delta H^{O}_{f298} = -158.5 + 1.5$ kcal/ mole) has been calculated from equilibrium data obtained from flow experiments performed by Grossweiner and Seifert³⁰ involving water vapor and crystalline beryllium oxide,

$$H_2O(g) + BeO(c) = Be(OH)_2(g)$$
(1)

Young³¹ in 1960 performed similar flow experiments and obtained a ΔF_{1673} of 27.7 kcal/mole for the free energy for reaction (1). This compared to Grossweiner and Seifert's value of 29 kcal/mole at the same temperature. Young's data resulted in ΔH_{f298}^{o} of -159.2 ⁺ .7 kcal/mole for Be(OH)₂(g). Regarding the entropy of Be(OH)₂(g), the JANAF Thermochemical Tables¹ have estimated frequencies which yield an entropy (57.97 cal/deg/mole) in good agreement with these second law heats.

Recently, an extensive experimental program was conducted in these laboratories⁵ which resulted in a second law value for ΔH_{f298} of 161⁺ 7 kcal/mole for Be(OH)₂(g) and a second law entropy at the average experimental temperature of 1705^oK of 92.6⁺ 4 cal/deg/mole. This compares with the entropy value from the JANAF Tables at 1705^oK of 89.93 cal/deg/mole. Based on the JANAF entropy values, data of Blauer, et al, ⁵ yield an average third law $\Delta H_{f298}^{o}{}_{K}$ of -165.9⁺ 0.7 kcal/mole, a difference of nearly 5 kcal between the second and third law heats of formation.

In an effort to obtain more reliable entropy data for $Be(OH)_2(g)$ an experimental spectroscopic investigation was undertaken in these laboratories. Once spectra are obtained they can be resolved by means of statistical mechanics which provide the fundamental equations necessary for the evaluation of the entropy of a gas. These equations relate the energies of the various states - translational, rotational, vibrational and electronic - to the probability of their propulation under specified conditions.

Vibrational-rotational molecular spectra of polyatomic molecules normally appear in the near infrared. These spectra, when obtained with highdispersion, high-resolution instruments, can be analyzed to provide the requisite data for entropy calculations. The experimental program involved two important experimental considerations for the spectroscopic study which include: (1) an abundance of the species under study and (2) spectral resolution of these from spectra occurring from other species which may complicate the spectrum of the compound in question. The thermodynamics of the BeO-water vapor reaction are such that water will be present in the system, regardless of the method of generation of the Be(OH)₂. Therefore, the operating conditions under which Be(OH)₂ is generated must be chosen in such a manner that the Be(OH)₂ concentration is maximized and the H₂O is minimized.

An extrapolation of the data of Blauer et al⁵ shows that 5% of Be(OH)₂ will be present in an equilibrium consisting of H₂O and Be(OH)₂ at 2500^oK.

The highest yield of $Be(OH)_2(g)$ is produced in the reaction of $H_2O(g)$ and BeO at high temperatures. Spectroscopic measurement: may thus be made directly on this high temperature gas mixture or the specie may be trapped in a suitable matrix and the spectral data of this material may be obtained at cryogenic temperatures. Flame temperatures in excess of 2750^OK have been reported for the H_2-O_2 flame; therefore, this flame, producing water vapor at high temperature, was chosen for the production of the Be(OH)₂ specie.

B. <u>Experimental</u>

Since a high temperature flame is capable of providing the necessary spectral data, the experimental apparatus was constructed so that either absorption or emission measurements could be made. The burner assembly and housing were designed for mounting in the sample space of a Perkin-Elmer 421 dual grating infrared spectrometer.

Figure 20 is a cross-sectional view of the H_2-O_2 burner assembly and housing Space limitations necessitated the inclusion of several mirrors to adapt the burner assembly light path to that of the spectrometer. For the absorption studies, light from the black-body radiator is deflected from the usual, straight-through path to a path which intercepts the flame and its products. In order to improve the amount of light reaching the infrared spectrometer (Perkin Elmer Infrared Grating Spectrograph Model 421) a collimated lens made from NaCl was placed at the exit of the light path from the apparatus. Photographs of the completed apparatus are shown in Figs. 21 and 22.



Fig. 20 Apparatus for Be(OH)₂ Generation



Fig.22. Complete Set Up for Be(OH) 2 Emission Studies The primary site of the $BeO-H_2O$ reaction is the inside wall of the oxygen-flow tube, above the end of the hydrogen tube, and a BeO target rod was inserted into the H_2-O_2 flame at this point. The burner tip was adjusted vertically, relative to the light path, to permit various portions of the flame to be examined.

The burner was housed in a chamber with a single exhaust port through which the lame products were removed by means of a vacuum-pressure pump. The burner exhaust gases were drawn through an aerosol filter to a fume exhaust blower.

Emission s pectra were observed for BeO in an oxyhydrogen flame in the region from 4000 cm⁻¹ to 550 cm⁻¹. Since atmospheric and reaction water absorption masked the 4000-3400 cm⁻¹ and the 1900-1300 cm⁻¹ ranges, the only new emission bands observed were in the 600-700 cm⁻¹ range. The band system is quite complex and is centered at 650 cm⁻¹. The band, as it appears (Fig. 23), could be a single frequency with complex rotational components, or several frequencies, and their rotational components. The spectrum (Fig. 23) was referenced against the spectrophotometer's internal glower. Both of these beams were reflected off the same mirrors and therefore had approximately the same path lengths. The sample beam, however, passes through an NaCl lens and an NaCl window of total thickness of ~ 10 mm.

Several experiments were performed employing this laboratory's three meter visible and U.V. grating spectrograph in an attempt to obtain the vibrational components of the electronic transitions of $Be(OH)_2$. A band had previously been reported resulting from flame emission spectra of beryllium salts in an oxyhydrogen flame located at 4710Å.³² However, exposures of one hour (103af plates) employing the same flame apparatus that was used in the infrared measurements showed only a faint band in the 4650-4800Å region. The band appeared as a continuum with no apparent line structure.

C. Discussion

Employing the data obtained from the experimental spectroscopic investigation of $Be(OH)_2(g)$ in these laboratories, together with data from the literature, a calculation has been made for the entropy.

The calculation of the spectroscopic entropy requires a knowledge of the structure of gas molecules. When a complete rotational-vibrational spectra is available, the structure may be deduced from an analysis of the spectra.



Since these spectra obtained were far from complete, a definite determination of the structure was not possible. Therefore, two molecular structural models have been assumed; a linear model, and a bent model. The bent model is the transconfiguration, but this form would give the same rotational contribution to the entropy as the cis, and the vibrational contributions would differ only slightly. The calculated entropies for the bent and linear forms may be sufficiently different, so that by comparison with the experimental values⁵ the correct model may be distinguished. The calculation of the spectroscopic entropy for both models can be divided into three parts: translational, rotational, and vibrational. The electronic and nuclear spin contributions have been disregarded. The calculations are based on the harmonic-oscillator rigid rotator approximation. At 298°K this approximation is very satisfactory. At 1705°K the influence of anharmonicity will make itself felt and the calculated vibrational entropy will no longer give the accurate representation as it did at 298⁰K. However, this disregard of anharmonicity at 1705[°]K is negligible compared with the other approximations that have been made.

The translational entropy for both models is the same, since it depends only on the temperature and molecular weight of the gaseous $Be(OH)_2$. The value at 298.15 ^OK is 37.190 cal/deg/mole, and at 1705 ^OK it is 45.849 cal/deg/mole.

The rotational entropy calculation involves estimates of the bond angles and bond distances. For both models the Be-O distance has been taken to be 1.5A. The JANAF Tables use 1.4A in their calculations. Since BeF₂ has a bond distance of 1.43A and the double bond distance in BeO is 1.4A, ¹ it seems unlikely that Be(OH)₂ would be the same as in these two molecules. The O-H distance has been taken to be 0.96A on the basis of compiled data.³³ Most oxygen bond angles are close to 105° : the smallest oxygen bond angle reported, 100° , is for OF₂.³³ Therefore, the Be-O-H angle has been taken to be 105° . This angle applies to the bent model. The linear model, of course, has an angle of 180° . The calculated rotational entropies for the bent model at 298° K and 1705° K are 20.597 and 25.792 cal/deg/mole, respectively. For the linear model these entropies are 14.397 and 17.860 cal/deg/mole. Fortunately, the moments of inertia, which depend on the bond angles and distances, represent roughly 15% of the rotational entropy in the bent case and 20% in the linear case. Errors of 0.1A and 5^o in estimating the distances and angles in calculating the moments

of inertia result in about 1.4 cal/deg/mole or 5-6% error in the rotational entropies for the bent model. For the linear model the error in estimating leads to 0.5 cal/ deg/mole or 2-3% error in the rotational entropies.

The vibrational entropy calculation requires a knowledge of the vibrational frequencies associated with the intramolecular motions of the molecule. On the basis of the two models (Fig. 24 and 25), the linear model should have seven frequencies independent of the cis or trans forms. A careful analysis of the frequencies of similar molecules was the basis for the assignment of values to these vibrational modes. The contribution to the vibrational entropy is greatest for the frequencies below 1000 $\rm cm^{-1}$. As discussed in the experimental section, this investigation centered on this region to the lower limit of 550 cm^{-1} . Newkirk³⁴ has reported an O-H stretch of 3500 for Be(OH)₂. This data allows the assignment for frequencies of 3500 and 3600 which are the symmetric O-H stretching vibrations, y_1 , for both models and the asymmetric O-H stretch, y_5 , for the bent model and ν_3 for the linear model. This allows an assignment of 1600 as a reasonable frequency for the asymmetric BeO stretch of y_{2} (bent) and y_{4} (linear). Values of 1100 for an OH torsion y_8 and 900 for Be-O symmetric stretch y_2 have been selected for the bent model. A value of 900 for symmetric Be-O stretch y_2 and 650 as an OA torsion \mathcal{Y}_5 (doubly degenerate) for the linear model. The experimental value of 650 is either a unique frequency or a group of frequencies as discussed previously in the experimental section. Therefore, for the bent model, values of 700,600 and 600 have been selected for Be-O bending, y_2 Be-O bending y_{9} and OH torsion y_{4} . For the linear model the value at 650 y_{6} (doubly degenerate) has been assigned as a Be-O bending mode. (These frequencies are listed in Table XIID.

Be(OH)₂ should have a slightly weaker force constant than BeF₂, however, the OH group has a mass of 17 compared to 19 for F. Therefore, the frequency of the low frequency mode in Be(OH)₂ should be quite close to that of BeF₂. Snelson³⁵ reports a frequency of 320 cm⁻¹ for the BeF₂. Therefore, the value chosen for the low frequency mode, V_7 , for the linear model is 300 cm⁻¹. The remaining frequency, V_5 , for the bent model has been assigned a value of 300 cm⁻¹ based on the bond character and distances Letween OH groups.

Based on these frequencies, the calculated vibrational entropy for the bent model yields values of 2.830 and 21.679 cal/deg/mole at 298° K and 1705° K, respectively. For the linear model the values are 4.463 at 298° K and











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27.231 cal/deg/mole at 1705° K. For a deviation of 100 cm^{-1} for each frequency in the bent model, the vibrational contribution to the entropy would change 0.8 e.u. at 298°K and 2.1 e.u. at 1705° K or approximately 30% at 298°K and 10% at 1705°K. For the linear model the changes would be 1.4 cal/deg/mole at 298°K and 2.1 cal/deg/mole at 1705° K or 30% and 8%. The choice of the 300 cm⁻¹ rather than the 600 cm⁻¹ choice of JANAF for one vibration contributes an additional 1.4 cal/deg/mole at 1705° K and 1.0 cal/deg/mole at 298°K for the bent model, and for the linear model they are 2.8 e.u. at 1705° K and 2.0 e.u. at 298° K.

The total discrepancies in entropy from the estimates of angles, distance and frequencies are 2.2 cal/deg/mole at 298° K and 3.5 cal/deg/mole at 1705° K for the bent model, which are roughly 4% of the total entropy. For the linear model the discrepancies are 1.9 cal/deg/mole at 298° K and 2.6 cal/deg/mole at 1705° K.

The resultant entropy at 298.15°K is 60.6 cal/deg/mole for the bent model and 56.1 cal/deg/mole for the linear model. At 1705° K the values are 93.3 cal/deg/mole for the bent model and 90.9 cal/deg/mole for the linear model.

The experimental value of Blauer, et al,⁵ of 92.6 ± 4 cal/deg/mole at 1705° K is nearly equal to that of the bent model 93.3 e.u. and somewhat higher than that of the linear mode 90.9 e.u.

This new calculated entropy allows a re-examination of the third "law heat of formation obtained by Blauer, et al.⁵ If the calculated entropy for the bent model of 93.3 e.u. is used, then a third law value at 1705° K for ΔH_{f1705° K of Be(OH)₂(g) of -166.4 kcal/mole is obtained. This is equal to the second law value at the average temperature of 1705° K. The third law value reduces to -160.9⁺ .7 at 258^oK.

REFERENCES

1. JANAF Thermochemical Tables.

| 2. | D. L. | Hilde | nbrand, | et ai, | Final | Technical | Report, | AFRPL- | TR-65-95 , |
|----|--------|--------|---------|--------|-------|-----------|---------|--------|-------------------|
| | Contra | act AF | 04(611) | -8523, | April | 1965. | | | |

3. JANAF Thermochemical Tables, December 31, 1960.

4. M. Farber, J. Chem. Phys. <u>36</u>, 1101 (1962).

5. J. Blauer, M. A. Greenbaum and M. Farber, J. Phys. Chem. <u>70</u>, 973 (1966).

6. E. Baur and R. Brunner, Helv. Chim. Acta. <u>17</u>, 958 (1934).

7. C. L. McCabe and C. E. Birchenall, J. Metals, <u>5</u>, 707 (1953).

8. R. B. Holden, R. Speiser and H. L. Johnston, J. Am. Chem. Soc. <u>70</u>, 3897 (1948).

9. E. A. Gulbransen and K. F. Andrew, J. Electrochem. Soc. 97, 383 (1950).

- 10. A. N. Nesmeyanov, <u>Vapor Pressure of the Chemical Elements</u>, Ed. by R. Gary, Elsevier Publishing Co., 1963, p. 172.
- 11. S. Dushman, <u>Vacuum Technique</u>, John Wiley and Sons, Inc., New York, 1949.
 - 12. Margaret A. Frisch, M. A. Greenbaum and M. Farber, J. Phys. Chem. <u>69</u>, 3001 (1965).
 - 13. P. Gross, C. S. Campbell, P. J. C. Kent and D. L. Levi, Disc. Fara. Soc., <u>4</u>, 206 (1948).
 - 14. A. S. Russell, K. E. Martin, and C. N. Cochran, J. Am. Chem. Soc. <u>73</u>, 1466 (1951).
 - S. A. Semenkovich, Zh. Prikl, Khim., <u>33</u>, 1281 (1960); Chem. Abstr. <u>54</u>, 1925d (1960).
 - 16. D. Bhogeswara Rao and V. V. Dadape, J. Phys. Chem. <u>70</u>, 1349 (1966).
- 17. A. G. Gaydon, <u>Dissociation Energies and Spectra of Diatomic Molecules</u>, Chapman and Hall, Ltd., London, 1953.
- 18. W. P. Sholette and R. F. Porter, J. Phys. Chem. <u>67</u>, 177 (1963).
- 19. A. Buchler and J. B. Berkowitz-Mattuck, J. Chem. Phys. <u>39</u>, 286 (1963).
- 20. D. L. Hildenbrand, W. F. Hall and N. D. Potter, J. Chem. Phys. <u>39</u>, 296 (1963).
- 21. M. D. Sheer, J. Phys. Chem. <u>61</u>, 1184 (1957).

| 22. | D. White, P. N. Walsh, H. W. Goldstein and D. F. Dever, J. Phys. Cham. <u>65</u> , 1404 (1961). |
|-----|--|
| 23. | A. Sommer, Ph.D. Thesis, The Ohio State University, 1962. |
| 24. | P. E. Blackburn, Arthur D. Little, Inc. Quarterly Report No. 4, Mar 1 - May 31, 1963, "Research on Thermodynamics of the Al-B-O and Be-B-O Systems." |
| 25. | John L. Margrave and Associates, "Molecular Parameters and Thermodynamic Properties of Light Element Molecules," Proceedings of the Third Meeting, ICRPG Working Group on Thermochemistry, March 1965. |
| 26. | O. J. Meschi, W. A. Chupka, and J. Berkowitz, J. Chem. Phys. <u>33</u> , 530 (1960). |
| 27. | D. White, D. C. Mann, P. N. Walsh and A. Sommer, J. Chem. Phys. 33, 488 (1960). |
| 28. | M. Farber, et al, Final Technical Report, Contract DA-04-495-ORD-1987, May 1960 through June 1962, Rocket Power, Inc. |
| 29. | JANAF Thermochemical Tables, March 31, 1961. |
| 30. | L. I. Grossweiner and R. L. Seifert, J. Am. Chem. Soc. 74, 2701 (1952). |
| 31. | W. A. Young, J. Phys. Chem. <u>64</u> , 1003 (1960). |
| 32. | Beckman Instruments, Inc. Bulletin No. 736c. |
| 33. | L. Pauling, <u>The Nature of the Chemical Bond</u> , Cornell University Press, Ithaca, N. Y. (1951), p. 168. |
| 34. | H. W. Newkirk, Inorg. Chem. <u>3</u> , 1041 (1964). |
| 35. | A. Snelson, IITRI Report C6013-3, Contract DA-31-124-ARO(D). |
| | |

TABLE I

EFFUSION CELL CHARACTERISTICS

| | Diameter of Effusion Hole | Depth of Hole | Clausing Factor, W _o |
|------------|---------------------------|---------------|---------------------------------|
| Cell No. 1 | 1.065 mm | 1.27 mm | 0.471 |
| Cell No. 2 | 0.990 mm | 2.0 mm | 0.359 |

TABLE II

| Run No. | (^{CII}) | Time (min) | Total Wt. Loss of BeO (mg) | Blank (mg) | Total Moles of H ₂ x 10 ² | Net Wt. Loss of BeO (110 min) | No. of Moles of H ₂ (110 min) |
|------------|--------------------|---------------|-------------------------------------|---------------|---|--|---|
| 29** | 2107 | 290 | 6.1 | 0.2 | 3.567 | 2.24 | 1.353 |
| 5** | 2139 | 230 | 7.0 | 0.5 | 2.459 | 3.11 | 1.176 |
| 30** | 2139 | 230 | 7.2 | 0.5 | 2.840 | 3.20 | 1.358 |
| 6** | 2171 | 230 | 8.9 | 1.2 | 2.344 | 3.68 | 1.121 |
| · 7** | 2171 | 230 | 9.0 | 1.2 | 2.524 | 3.73 | 1.207 |
| 8** | 2171 | 230 | 8.7 | 1.2 | 2.944 | 3.59 | 1.408 |
| 9** | 2203 | 110 | 5.8 | 1.0 | 1.372 | 4.80 | 1.372 |
| 10** | 2203 | 230 | 12.1 | 2.6 | 2.745 | 4.54 | 1.313 |
| 11** | 2235 | 110 | 8.5 | 2.5 | 1.487 | 6.0 | 1.487 |
| 12** | 2235 | 110 | 8.0 | 2.0 | 1.372 | 6.0 | 1.372 |
| 13** | 2235 | 230 | 18.4 | 5.0 | 2.613 | 6.41 | 1.250 |
| 14** | 2235 | 230 | 17.0 | 5.0 | 2.548 | 5.74 | 1.218 |
| 15** | 2268 | 110 | 11.9 | 4.3 | 1.425 | 7.60 | 1.425 |
| 17* | 2268 | 110 | 13.3 | 2.2 | 1.487 | 11.1 | 1.487 |
| 18* | 2268 | 110 | 11.4 | 2.2 | 1.434 | 9.2 | 1.434 |
| 19* | 2268 | 110 | 12.0 | 2.2 | 1.449 | 10.0 | 1.449 |
| 16** | 2301 | 110 | 14.6 | 5.2 | 1.349 | 9.4 | 1.349 |
| 20* | 2301 | 110 | 21.1 | 4.3 | 1.420 | 16.8 | 1.420 |
| 21* | 2301 | 110 | 20.9 | 4.3 | 1.420 | 16.6 | 1,420 |
| 22* | 2301 | 110 | 19.8 | 4.3 | 1.482 | 15.5 | 1.482 |
| 23* | 2335 | 110 | 25.8 | 6.8 | 1.430 | 19.0 | 1.430 |
| 24* | 2335 | 110 | 27.6 | 6.8 | 1.415 | 20.8 | 1.415 |
| 25* | 2368 | 110 | 31.8 | 8.5 | 1.372 | 23.3 | 1.372 |
| 26* | 2368 | 110 | 30.0 | 8.5 | 1.377 | 21.5 | .377 |

EXPERIMENTAL DATA OF MOLECULAR FLOW EFFUSION EXPERIMENTS

* Cell 1

** Cell 2

TABLE III

CALCULATED DATA FOR THE REACTION

 $BeO(c) + 1/2H_2(g) = BeOH(g)$

| Run No. | т (⁰ К) | Loss of BeO (mg/110 min) | PBe, atm x 10 ⁶ | Be Wt. Loss due to Be Formation (mg/110 min) | BeO Wt. Loss due to BeOH Formation (mg/110 min) | P_{H_2} atm x 10 ⁴ | PBEOH, atm x 10 ⁵ | K ₂ x 10 ³ |
|-------------|------------------------|-----------------------------------|----------------------------------|---|--|---------------------------------------|------------------------------------|----------------------------------|
| 29** | 2107 | 2.24 | 1.94 | 0.28 | 1.96 | 10.29 | 22.69 | 0.707 |
| 5** | 2139 | 3.11 | 2.36 | 0.34 | 2.77 | 8.849 | 3.232 | 1.086 |
| 30** | 2139 | 3.20 | 2,55 | 0.37 | 2.83 | 10.28 | 3.302 | 1.029 |
| 6** | 2171 | 3.68 | 2.99 | 0.43 | 3.25 | 8.362 | 3.821 | 1.322 |
| 7** | 2171 | 3.73 | 3.12 | 0.45 | 3.28 | 9.031 | 3.856 | 1.283 |
| 8** | 2171 | 3.59 | 3.38 | 0.49 | 3.10 | 10.61 | 3.644 | 1.120 |
| S ** | 2203 | 4.80 | 4.30 | 0.62 | 4.18 | 10.22 | 4.949 | 1.546 |
| 10** | 2203 | 4.54 | 4.19 | 0.60 | 3.94 | 9.773 | 4.665 | 1.495 |
| 11** | 2235 | 6.0 | 5.71 | 0.81 | 5.19 | 10.99 | 6.188 | 1.867 |
| 12** | 2235 | 6.0 | 5.46 | 0.78 | 5.27 | 10.08 | 6.224 | 1.957 |
| 13** | 2235 | 6.41 | 5.18 | 0.74 | 5.67 | 9.129 | 6.701 | 2.239 |
| 14** | 2235 | 5.74 | 5.11 | 0.73 | 5.01 | 8.890 | 5.974 | 2.005 |
| 15** | 2268 | 7.60 | 7.09 | 1.00 | 6.60 | 10.34 | 7.929 | 2.466 |
| 17* | 2268 | 11.1 | 5.72 | 1.23 | 9.87 | 6-905 | 7.825 | 2.981 |
| 18* | 2268 | 9.2 | 5.61 | 1.20 | 8.0 | 6.690 | 6.342 | 2.453 |
| 19* | 2268 | 10.0 | 5.64 | 1.21 | 8.79 | 6.712 | 6.969 | 2.691 |
| 16** | 2301 | 9.4 | 8.63 | 1.21 | 8.19 | 9.534 | 9.913 | 3.213 |
| 20* | 2301 | 16.8 | 6.94 | 1.48 | 15.32 | 6.336 | 12.24 | 4.857 |
| 21* | 2301 | 16.6 | 6,94 | 1.48 | 15.12 | 6.342 | 12.08 | 4.794 |
| 22* | 2301 | 15.5 | 7.12 | 1.52 | 13.98 | 6.663 | 11.17 | 4.329 |
| 23* | 2335 | 19.0 | 8.67 | 1.83 | 17.17 | 6.176 | 13.82 | 5.573 |
| 24* | 2335 | 20.8 | 8.59 | 1.32 | 18.98 | 6.082 | 15.27 | 6.195 |
| 25* | 2368 | 23.3 | 10.27 | 2.16 | 21.14 | 5.621 | 17.14 | 7.232 |
| 26* | 2368 | 21.5 | 10.32 | 2.17 | 19.33 | 5.666 | 15.67 | 6.584 |

* Cell 1

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** Cell 2

TABLE V

FREE ENERGY AND HEAT OF REACTION FOR THE EQUILIBRIUM

 $BeO(c) + 1/2H_2(g) = BeOH(g)$

| т°к | ΔF_r (kcal) | ∆ H _r (kcal) | ΔH_r^{298} (kcal) |
|------|---------------------|-------------------------|---------------------------|
| 2107 | 30.37 | 87.207 | 94.603 |
| 2139 | 29.01 | 86.553 | 94.109 |
| 2139 | 29.25 | 86.788 | 94.344 |
| 2171 | 28.59 | 86.834 | 94.549 |
| 2171 | 28.73 | 86.973 | 94.688 |
| 2171 | 29.32 | 87.560 | 95.275 |
| 2203 | 28.34 | 87.276 | 95.151 |
| 2203 | 28.48 | 87.418 | 95.293 |
| 2235 | 27.91 | 87.542 | 95.580 |
| 2235 | 27.70 | 87.328 | 95.366 |
| 2235 | 27.10 | 86.734 | 94.772 |
| 2235 | 27.59 | 87.225 | 95.262 |
| 2268 | 27.07 | 87.947 | 96.153 |
| 2268 | 26.22 | 86.556 | 94.762 |
| 2268 | 27.10 | 87.438 | 95.644 |
| 2268 | 26.67 | 87.012 | 95.218 |
| 2301 | 26.26 | 87.308 | 95.684 |
| 2301 | 24.37 | 85.413 | 93.789 |
| 2301 | 24.31 | 85.360 | 93.736 |
| 2301 | 24.88 | 85.929 | 94.305 |
| 2335 | 24.08 | 85.821 | 94.409 |
| 2335 | 23.59 | 85.329 | 93.917 |
| 2368 | 23.20 | 85.602 | 94.395 |
| 2368 | 23.64 | 86.046 | 94.839 |

TABLE V

SUMMARY OF THERMODYNAMIC DATA FOR THE REACTION

 $BeO(c) = 1/2H_2(g) = BeOH(g)$

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| ∆ н [°] _{r2235} ° _К | 88.3 ⁺ 3.8 kcal | 87.2 ⁺ 0.5 kcal |
|---|--------------------------------------|----------------------------------|
| [∆] ,H [°] _{r298} °K | 96.3 ± 3.8 kcal | 94.9 ⁺ 0.5 kcal |
| $\Delta \operatorname{H}^{o}_{f2235} \operatorname{o}_{K} \operatorname{BeOH(g)}$ | -55.7 + 3.8 kcal/mole | -56.7 ± 0.5 kcal/mole |
| △ H ^o _{f298} o _K BeOH(g) | -46.8 [±] 3.8 kcal/mole | -48.2 ⁺ 0.5 kcal/mole |
| ∆s ^o r2235 ^o K | 27.4 ⁺ 1.7 cal/deg/mole | |
| S ⁰ 2235 ⁰ K BeOH(g) | 75.3 ⁺ _1.7 cal/deg/mole | |
| S ⁰ 298 ⁰ K ВеОН(g) | 54.0 ⁺ _ 1.7 cal/deg/mole | 53.29 cal/deg/mole(JANAF) |

TABLE VI

EXPERIMENTAL DATA FOR THE REACTION OF AICI₃ VAPOR AND Al(s, l)

| C | -0 | Time | Wt. Loss of AlCl ₃ | Wt. Loss of Al | Flow of A | Moles of A/hour 2 | Moles of Al/hour (wt. loss) |
|---|-------------|-------|----------------------------------|-------------------|-----------|---------------------|--------------------------------|
| | Т°К | (hrs) | (gm) | (mg) | (cc/hr) | $(N_A) \times 10^2$ | $(N_{Al}) \times 10^{\circ}$ |
| | 800 | 6,0 | 28.85 | 5.9 | 1338 | 5.48 | े .643 |
| | 800 | 6.0 | 47.95 | 8.7 | 1338 | 5.48 | 5.370 |
| | 800 | 5.0 | 54.41 | 6.3 | 1440 | 5.91 | 4.670 |
| | 800 | 5.0 | 34.10 | 4.0 | 1482 | 6.07 | 2.960 |
| | 800 | 3.0 | 29.08 | 3.9 | 1030 | 4.22 | 4.820 |
| | 800 | 0.67 | 11.00 | 1.7 | 1026 | 4.20 | 9.450 |
| | 800 | 5.0 | 16.99 | 2.3 | 938 | 3.85 | 1.705 |
| | 825 | 5.0 | 38.20 | 55.7 | 1338 | 5.48 | 4.220 |
| | 825 | 5.0 | 28.05 | 4.6 | 1338 | 5.48 | 3.410 |
| | 851 | 5.0 | 47,05 | 8.9 | 1060 | 4.34 | 6.59(|
| | 851 | 4.5 | 36.33 | 6.0 | 1020 | 4.18 | 4.930 |
| | 851 | 3.5 | 23.70 | 4.8 | 1092 | 4.47 | 5.070 |
| C | 875 | 5.0 | 29.05 | 6.2 | 912 | 3.74 | 4.0 00 |
| | 875 | 3.0 | 25.80 | 5.6 | 1194 | 4.90 | 6.910 |
| | 875 | 4.0 | 36,90 | 7.8 | 1194 | 4.90 | 7.225 |
| | 901 | 4.5 | 26.10 | 7.7 | 913 | 3.74 | 6.330 |
| | 901 | 4.0 | 26.55 | 7.5 | 956 | 3.92 | 6.960 |
| | 901 | 4.0 | 24.22 | 7.3 | 1020 | 4.18 | 6.780 |
| | 925 | 4.0 | 30.90 | 17.2 | 1338 | 5.48 | 15.94 |
| | 925 | 4.0 | 37.40 | 11.0 | 1194 | 4.90 | 10.20 |
| | 925 | 4.0 | 23.02 | 10.6 | 1194 | 4.90 | 9.82 |
| | 949 | 3.0 | 18.35 | 15.5 | 1338 | 5.48 | 9.15 |
| | 949 | 5.0 | 11.07 | 7.7 | 1290 | 5.29 | 5 71 |
| | 949 | 1.25 | 13.41 | 5.1 | 942 | 3.86 | 15.10 |
| | 949 | 2.5 | 21.47 | 10.2 | 942 | 3.86 | 15.10 |
| | 975 | 3.0 | 26.80 | 19.7 | 1338 | 5.48 | 24.35 |
| | 975 | 3.5 | 39.00 | 26.3 | 1338 | 5.48 | 27.85 |
| 4 | 975 | 4.0 | 22.33 | 5.5 | 1250 | 5.12 | 14.38 |
| | 9 75 | 3.0 | 29.40 | 21.4 | 1338 | 5.48 | 26.42 |
| | 1000 | 4.0 | 39.45 | 38.9 | 1338 | 5.48 | 36.00 |
| | 1000 | 4.0 | 30.65 | 29.9 | 1338 | 5.48 | 27.65 |

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TABLE VII

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EQUILIBRIUN DATA FOR THE REACTION OF AICI3 AND AI

| Average Log K ₆ | - 6.84 | -7.35 | -7.66 |
|--|---|--|--|
| К ₆ х 10 ⁸ | 34.20 28.90 07.85 06.16 12.98 24.55 09.12 | 05.39 06.57 05.40 02.92 05.77 | 02.22 02.32 01.98 |
| N _{Al} × 10 ⁵ per hr due to Reaction (6) | 3.433 5.077 4.306 2.679 4.436 9.016 1.550 | 3.630 2.917 5.380 3.845 4.081 | 4.923 4.705 3.042 |
| N _{Al} × 10 ⁵ per hr due to Reaction (5) | 0.119 0.162 0.205 0.153 0.168 0.168 0.251 0.251 | 0.301 0.246 0.597 0.532 0.478 | 1.055 1.008 0.706 |
| ĸs | 3.2×10^{-12} | 1.446×10^{-11} 1.446×10^{-11} 6.31×10^{-11} 6.31×10^{-11} 6.31×10^{-11} 6.31×10^{-11} | 2.28×10^{-10} 2.28 × 10^{-10} 2.28 × 10^{-10} |
| N _{Al} x 10 ⁵ per hr due to Reaction (1) | 0.098 0.131 0.160 0.128 0.128 0.184 0.072 | 0.289 0.247 0.613 0.553 0.511 | 1.247 1.197 0.852 |
| ۴ı | $\begin{array}{c} 4.34 \times 10^{-14} \\ 4.34 \times 10^{-14} \\ 4.34 \times 10^{-14} \\ 4.37 \times 10^{-14} \end{array}$ | 3.90×10^{-13} 3.90×10^{-13} 2.82×10^{-12} 2.82×10^{-12} 2.82×10^{-12} 2.82×10^{-12} | 1.70×10^{-11} 1.70 × 10^{-11} 1.70 × 10^{-11} |
| $ \begin{array}{c} {}^{\rm N}{}_{\rm Al_2Cl_6} \\ \\ {}^{\rm x} {}_{\rm 10}{}^{\rm 3} \\ {}^{\rm (hr)} \end{array} $ | 11.40 20.30 28.45 16.75 25.30 45.40 08.09 | 17.20 12.03 19.80 16.60 13.21 | 16.25 14.85 09.76 |
| NAIC1 ₃ x 10 ² (hr) | 1.335 1.950 2.490 1.780 2.120 3.320 0.937 | 2.310 1.814 3.110 2.740 2.388 | 3.690 3.500 2.418 |
| K 4 | 0.199 991.0 991.0 991.0 991.0 991.0 | 0.323 0.323 0.523 0.523 0.523 | 0.845 0.845 0.845 |
| T ^o K | 500 800 800 800 800 800 | 825 825 851 851 851 | 875 875 875 |

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TABLE VII - continued

No. of Concession, Name

Average Log K₆ -7.76 -7.59 -8.40 -8.23 -8.21 x 10⁸ 02.02 00.14 01.18 01.95 01.57 01.74 02.88 02.90 00.25 00.90 00.83 00.10 00.95 00.33 00.77 01.61 yo Y per hr due to Reaction (6) $N_{A1} \times 10^{5}$ 7.970 3.870 3.508 4.630 3.934 0.785 1.745 3.767 5.530 5.980 5.170 3.560 7.120 6.040 7.750 3.861 per hr due to Reaction (5) NAIX 10⁵ 1.293 2.020 1.766 3.920 1.352 1.580 4.150 4.505 8.450 10.310 1.214 2.580 4.690 7.090 7.850 6.580 8.42×10^{-10} 8.42×10^{-10} 8.42×10^{-10} 2.57×10^{-9} 2.57×10^{-9} 5.92 × 10⁻⁸ 5.92×10^{-8} 2.57×10^{-9} 7.4×10^{-9} 2.2×10^{-8} 7.4×10^{-9} 2.2×10^{-8} 7.4×10^{-9} 2.2×10^{-8} 2.2×10^{-8} 7.4×10^{-9} KS per hr due to Reaction (1) $N_{A1} \times 10^{5}$ 1.774 6.175 1.608 3.345 6.210 8.120 3.170 2.820 7.830 11.830 12.880 11.730 15.330 .720 6.850 17.940 4.57×10^{-10} 4.57×10^{-10} 9.34×10^{-11} 4.57×10^{-10} 9.34 × 10⁻¹¹ 1.78 × 10⁻⁹ 9.34 × 10⁻¹¹ 1.78×10^{-9} 1.78×10^{-9} 1.78 × 10⁻⁹ 6.31×10^{-9} 6.31×10^{-9} 6.31×10^{-9} 6.31×10^{-9} 2.0 × 10⁻⁸ 2.0×10^{-8} Y. NA12C16 × 10³ (hr) 05.76 01.08 03.66 09.23 96.98 06.90 03.33 08.16 06.13 05.23 08.13 07.80 04.87 08.33 08.32 10.94 NAIC13 x 10² 2.925 3.168 4.806 5.028 2.726 1 450 5.990 3.534 6.743 6.365 3.154 4.144 5.254 5.892 5.494 2.800 2.000 2.920 2.000 2.000 2.920 2.920 4.270 4.270 6.250 1.370 4.270 4.270 6.250 i.370 1.370 2.920 K4 925 925 925 949 949 949 975 975 975 975 1000 1000 106 106 949 $\mathbf{T}^{\mathbf{O}}\mathbf{K}$ 106

TABLE VIII

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THERMODYNAMIC DATA FOR AICI2(g) AND AI2CI4(g)

| ΔH _{f900} °K ^{A1} 2 ^{C1} 4 ^(g) | -194.8 <mark>+</mark> 1.0 kcal/mole |
|--|-------------------------------------|
| S ^o 900 ^o K ^{A1} 2 ^{C1} 4 ^(g) | -115.7 ⁺ 1.1 cal/deg/mol |
| $\Delta H_{f298} \circ_{K} Al_2 Cl_4(g)$ | -194 ± 5 kcal/mole |
| S ⁰ 298 ⁰ K Al ₂ Cl ₄ (g) | 90 ⁺ 5 cal/deg/mole |
| $\Delta H_{f298} \circ_{K} AlCl_{2}(g)$ | -66 ⁺ _3 kcal/mole |

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TABLE IX

EQUILIBRIUM DATA FOR THE REACTION

$AlCl_3(g) + AlCl(g) = 2AlCl_2(g)$

| т ^о к | PAICI ₃ (atm) | PAIC1 (atm) | P _{AlCl2} (atm) | % AlC1 ₂ Compared to AlC1 |
|------------------|-----------------------------|----------------------|-----------------------------|--|
| 850 | 3.0×10^{-5} | 4.2×10^{-6} | 3.9×10^{-7} | 8.4% |
| 1000 | 3.7×10^{-5} | 8.2×10^{-5} | 4.3×10^{-6} | 5.0% |

TABLE X

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 $(D_2 \text{ Pressure} = 75 \mu)$

A TYPICAL MASS SPECTROMETRIC DETERMINATION OF

THE SPECIES RESULTING FROM THE REACTION $B_2O_3(1)$ and $D_2(g)$

| | log K ₃ | | | | | | 0.954* | 1.290 | 1.580 | 1.857 | 2.130 | |
|------------|---|------------------------|----------------------|------------------------|------------------------|------------------------|------------------------|------------------------|------------------------|------------------------|------------------------|--|
| | log K ₂ | 0.477* | 0.772* | 1.188* | 1.604* | 1.894* | 2.272 | 2.556 | 2.803 | 3.090 | 3.352 | |
| | K ₂ | 3.00 | 5.20 | 1.54×10^{1} | 4.02×10^{1} | 7.84×10^{1} | 1.87×10^{2} | 3.60×10^{2} | 6.35×10^{2} | 1.23×10^{3} | 2.25 x 10 ³ | |
| | log K ₁ | 3.393 | 3.840 | 4.288 | 4.702 | 5.072 | 5.491 | 5 . 85 ົ | 6.190 | 6.550 | 6.824* | |
| | К ₁ | 2.48 x 10 ³ | 7.01×10^{3} | 1.94 x 10 ⁴ | 5.04 x 10 ⁴ | 1.18 × 10 ⁵ | 3.10 × 10 ⁵ | 7.11 × 10 ⁵ | 1.55 × 10 ⁶ | 3.55 × 10 ⁶ | 6.67 × 10 ⁶ | |
| Mass 70 | $^{\mathrm{I}}\mathrm{B}_{2}\mathrm{O}_{3}$ | | | | | | 0.6 | 19.5 | 38. | 72. | 135. | |
| Mass 45 | 08001 | 15. | 25.5 | 48. | 88.5 | 145.5 | 265. | 435. | .069 | 1110. | 1710. | |
| Mass 29 | 080 | 165. | 275. | 405. | 570. | 610. | 1170. | 1635. | 2250. | 3200. | 3900. | |
| Mass 20 | $^{I}D_{2}O$ | 75. | 110. | 150. | 195. | 270. | 375. | 525. | 750. | 1000. | 1300. | |
| 1/T | × 10 ³ | .8380 | .8200 | .8026 | .7859 | .7698 | .7542 | .7392 | .7246 | .7106 | .6970 | |
| Ŀ | У _о К | 1161.1 | 1187.3 | 1213.8 | 1240.4 | 1267.2 | 1294.3 | 1321.5 | 1349.4 | 1377.3 | 1405.5 | |

*Discarded

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TABLE XI

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INTENSITIES OF MAJOR SPECIES OF REACTION $B_2O_3(1) + D_2(g)$ (TEMP. 1518°K, PRESSURE 240 μ)

| D ₂ O | 2700 | .23 |
|--------------------|-------|-------|
| BO | 1260 | .11 |
| DBO | 11850 | 1.00 |
| BO ₂ | 585 | .049 |
| DOBO | 7600 | . 64 |
| B(OD) ₂ | 162 | .014 |
| (BO) ₂ | 75 | .0063 |
| B202 | 555 | .047 |

The trimers $(BOD)_3$ and $(DOBO)_3$ were not observed.

TABLE XII

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THERMODYNAMIC DATA FOR THE REACTIONS

OF $P_{j}O_{3}(1)$ AND $D_{2}(g)$

| Run | D ₂ (g) | Tavg. ^o K | ▲ H ₁ kcal/mole | Tavg. ^o K | ∆H ₂ kcal/mole | Tavg. ok | ∆ H ₃ kcal∕mole |
|--------|--------------------|-------------------------|-------------------------------|-------------------------|------------------------------|-------------|-------------------------------|
| 1 | 75 | 1269 | 106.6±0.5 | 1350 | 80,6±1.3 | 1364 | 85.1 ± 0.2 |
| 2 | 75 | 1256 | 103.7 ± 0.9 | 1329 | 82.9 ± 1.2 | 1364 | 86.2 1 3.5 |
| * ° | 240 | 1310 | 96.4 - 0.9 | 1270 | 80.9 ± 0.8 | 1364 | 84.8+.3 |
| 4* | 240 | 1271 | 100.2 ± 0.7 | 1271 | 80.1 ± 1.2 | 1351 | 85.2 ± 1.0 |
| S | 18 | 1215 | 104.0 + 1.1 | 1296 | 79.5 ± 1.2 | 1324 | 84.2 [±] 0.6 |
| 9 | 18 | 1245 | 102.1±0.6 | 1311 | 83.0±0.8 | 1284 | 91.4 - 6.8** |
| 2 | 18 | 1296 | 103.5 ± 1.2 | | | 1298 | 85.1±0.8 |
| 80 | 240 | 1311 | 98.8±0.6 | | | 1338 | 88.9 1 2.4 |
| 6 | . 5 | 1257 | 106.4 ± 0.6 | | | 1298 | 88.3 ± 0.9 |
| verage | | 1260 | 104.4 ± 1.5 | 1320 | 81.5 ⁺ 1.5 | 1320 | 86.3 ± 1.5 |
| *Run d | Iscarded | | | | | | |

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**Point discarded

TABLE XIII

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ESTIMATED FREQUENCIES FOR $Be(OH)_2(g)$

| Bent Mo | odel | Linear M | lodel | - |
|---------|-----------------------|----------|-----------------------|-----|
| 3600 | ν ₆ | 3600 | V ₃ | |
| 3500 | ν_1 | 3500 | $\boldsymbol{\nu}_1$ | |
| 1600 | <i>L</i> ₇ | 1600 | \mathcal{V}_4 | |
| 1100 | ۲ ₈ | 900 | L 2 | |
| 900 | L ₂ | 650 | $\boldsymbol{\nu}_6$ | (2) |
| 700 | ل ₃ | 650 | 25 | (2) |
| 600 | \mathcal{V}_4 | 300 | <i>ل</i> 7 | (2) |
| 600 | L ₉ | | | |
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