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Equilibria in the Oxide Systems of Praseodymium and Americium

L. B. Asprey and B. B. Cunningham

April 14, 1949



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Equilibria in the Oxide Systems of Praseodymium and Americium

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#### ABSTRACT

The thermal decomposition of the higher oxides of praseodymium and americium have been investigated. The decomposition of  $PrO_{1.333}(Pr_6O_{11})$  to  $PrO_{1.5}(Pr_2O_3)$  was accomplished and a value of 47  $\pm$  5 kcals. for the heat of the reaction obtained. The reaction proved to be very complex. The decomposition of  $AmO_2$  was carried out only to  $AmO_{1.85}$  due to the greater stability of  $AmO_2$ . It was impossible  $\pm$  to extend the value = of 21 kcals. obtained for the partial heat of the reaction to obtain a value for the decomposition to  $AmO_{1.5}$ . The composition of the heat of the reaction to be  $AmO_{1.93} \pm 0.02$ .

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Equilibria in the Oxide Systems of Praseodynium and Americium

L. B. Asproy and B. B. Cunningham Radiation Laboratory and Department of Chemistry University of California, Berkeley, California

#### April 14, 1949

#### I. Introduction

An investigation of the thermal decomposition of the higher oxides of praseodymium and americium was undertaken with the following objectives in mind:

a) Equilibrium data would permit an evaluation of  $\Delta F$ ,  $\Delta H$  and  $\Delta S$  for the reactions

 $\frac{\Pr_{0_2} \longrightarrow 1/2 \operatorname{Pr}_{2^{0_3}} + 1/4 \operatorname{O}_{2}}{\operatorname{AmO}_{2} \longrightarrow 1/2 \operatorname{Am}_{2^{0_3}} + 1/4 \operatorname{O}_{2}}$ 

Such data would furnish a basis for estimating the free energies for the aqueous reactions

$$Pr^{+3}(aq) + H^{+} = Pr^{+4}(aq) + 1/2 H_2$$
  
Am^{+3}(aq) + H^{+} = Am^{+4}(aq) + 1/2 H\_2

b) In the case of americium this would constitute the first exidation-reduction equilibrium studies for this element, and would be of value in estimating the thermodynamic stability of other compounds (AmF<sub>1</sub>, for example).

c) The thermal stability and exact composition of the compound  $\times$  "AmO<sub>2</sub>" would be directly determined and the exygen pressure measured as a function of temperature. This would constitute a check on the validity of the assumption that the composition AmO<sub>2</sub> results from the direct ignition of the nitrate in air at  $800^{\circ}$  C -- an assumption made in calculating the half-life of the isotope Am<sup>241</sup> from specific activity measurements based on weighing the higher exide<sup>1</sup>.

It should be nontioned that although the americium oxide equilibria wore of primary interest, practical considerations required that most of the initial work on this problem be carried out with a "stand in" element. Because of the extremely limited availability and high toxicity of americium, the apparatus and techniques employed were developed using praseodymium as the most suitable "stand in" for americium.

Not all of the objectives mentioned previously were attained in this investigation. As will be made evident subsequently, the principle difficulty encountered arose from the (unanticipated) highly complex nature of the thermal decomposition of the higher praseodymium oxides. Because of the unique complexity of this reaction it was investigated in some detail. Another difficulty was the stability of  $AmO_2$  which proved to be much greater than previously believed. It was impossible to carry out the thermal decomposition of  $AmO_2$  beyond  $AmO_{1.35}$  at the temperatures available with the apparatus used.

Of the experimental methods which have been used to study equilibria between solids and gases, two were investigated as best fulfilling the requirements as to sample size and reduction of the hazard associated with handling the highly radioactive dry americium oxide.

In the first method, the course of the decomposition was followed with a simple guartz fiber cantilever balance inside a closed system by measuring the weight of the oxide under various pressures of oxygen. Suitable reaction rates could be attained only at elevated temperatures. At these temperatures thermal currents prevented any measurement of the weight of the oxide. Hence after a preliminary qualitative investigation on praseodymium, the method was abandoned.

The second and more useful method consisted of following the course of the reaction by measurement of the pressure of oxygen in equilibrium with the solid in a system of known volume and temperature. Such investigations are ordinarily carried out with much larger samples than those used here, so that a relatively large change in oxygen pressure causes a very small change in composition of the

oxide. Particularly for the americium system but also for the praseodymium, since the amount of material available was limited to very small amounts, severe difficulties were placed in the way of accurate measurements. A system had to be designed so that the degradation of the oxide all the way to the sesquioxide would occur at readily measurable pressures, i.e. the volume of the system had to be small, of the order of one to two milliliters. Further, since the total weight of oxygen evolved during the transition to MO1.5 was of the order of 100-200 micrograms, it was necessary to prevent the reaction of this exygen with any other material in the system such as mercury vapor or stopcock grease or large errors in the calculation of the composition of the solid phase would result. As the vapor pressure of nercury at room temperature is  $10^{-3}$  mm and since a mercury manometer was used to measure pressures, it was obvious that this vapor must be kept from contact with the hot oxide. A cold trap maintained at the temperature of sublimation of solid CO2 was built into the system, separating both the stopcocks and the manometer from the part of the system undergoing heating. Due to volume considerations, small bore tubing was used for the manometer, introducing large corrections for the capillary depression of Hg. Since the weight of oxygen in the system was calculated from the known volume, pressure and density of the gas, it was necessary to correct for the offect of the cold trap and the heated Further errors resulted from absorption of oxygen on the section of the system. oxide itself and on the walls of the tubing of which the system was constructed. Since it was impossible to outgas the sample tube itself without subjecting the oxide to prolonged heating at high temperatures which would tend to decrease the reactivity of the sample, a further error entered due to the gases desorbed from the fused silica tube.



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#### II. Apparatus

The system consisted of a manometer, a cold trap with a 3-way stopcock connecting to gas and vacuum lines, and a quartz capillary tube containing the oxide whose vapor pressure was to be measured. A thermocouple and potentioneter more used for measuring the temperature of the oxide, and a platinum tube was placed around the sample end of the quartz tube and was surrounded in turn by a copper coil used for induction heating. Differences in Hg level were measured by means of an cathetometer equipped with a telescope.

The mercury manometer was constructed of 3.6 mm. I. D. precision bore tubing. It was of the closed type and could measure pressures up to 200 mm of mercury. The closed end **population** a section of capillary to remove desorbed gases<sup>2</sup>. The open end of the manometer was sealed onto a short piece of 1 mm capillary, bent at right angles, to which was attached a capillary socket joint designated as S 12/1.

To this nanonster, a cold trap made out of 1/2 nm capillary tubing was connected by an S 12/1 ball joint as shown in the diagram. The purpose of the trap was to keep mercury vapor from the hot end of the system and to remove any condensable gases. The middle heg of the trap was connected to a 3-way stopcock which in turn connected to gas and vacuum lines. By this means, any easily condensable gases were removed from exygen added to the system. The end of the trap opposite to the manometer was blown out and then partially melted to make a flared joint to which the guartz tube could be fastened by Apiezon wax. The trap was cooled to  $-78.5^{\circ}$  C by a mixture of crushed "dry ice" and ethyl alcohol.

A thick walled quartz tube, 23-25 cm. long and about 1 mm I. D. and 1.25 mm wall thickness was used to contain the sample. The end of the tube was sealed off and then an eight cm. piece of the same tubing welded on to reduce temperature end effects when the tubing was heated. The end of the tube to be fastened to the cold trap was rounded by grinding on emery paper.

The oxide was heated by means of a 3 KVA oscillator, feeding into a copper coil. The sample tube was surrounded by a 12 mm. O. D. cuartz tube over which was **ship**ped a platinum tube 9 cm. in length. The water cooled induction coil consisted of a 6.0 cm. diameter coil, 16 cm. long, containing 16 turns, of 1/4" O. D. copper tubing, connected to the oscillator. The coil was placed so that the oxide sample and the outer platinum tube were contered in the copper coil. Test showed that the temperature was constant to within <sup>±</sup> 1° C over the length of tube occupied by the sample.

Earlier experiments had employed a nickel block 5" long and 1-1/4" in diameter containing 2 holes, one for the sample tube and another for the thermocouple. The Ni block was surrounded by an asbestos insulated tube furnace of the ordinary type. The temperature gradient was too large and the time consumed in coming to an equilibrium temperature was too great so the change was made to induction heating.

Attempts to measure the temperature of the oxide sample by means of an exterior thermocouple gave variable results so an interior thermocouple was used. Five mil Pt, Pt-10% Rh wire was inserted down the length of the quartz sample tube. At the lower end of the thermocouple one wire was insulated from contact with the other by means of fine quartz capillary, and near the waxed joint by a coat of glyptal varnish. The hot junction was buried in the oxide or was within 2 mm of it for all runs. A cold junction was maintained at 0° C by means of an ice and water bath. The effect of the inductive field on the thermocouple was small and it was felt that the thermal conductivity between the junction and sample was sufficiently great so no appreciable error in the measurement of the temperature would result.

The potential developed was measured by a Rubicon Portable Precision Potentiometer of more than adequate sensitivity, since measurements were made only to the nearest degree Centigrade.

The difference in the level of the 2 Hg arms of the manometer was measured by means of a Gaertner cathetometer, equipped with a telescope and a vernier so that readings could be made accurately to the nearest 0.05 mm and estimated still more closely.

Since it was necessary to know the total volume at any pressure in order to calculate the composition, all parts of the system were calibrated by filling with morcury which was then removed and weighed. A reference mark was scratched on the procision bore section of the manometer near the open end. The volume of the manometer up to this point was measured before the manometer was filled with mercury. Changes in the volume of the system due to pressure changes could then easily be calculated. An assumption was made that the menisci were spherical segments and their volume calculated on this basis. The volume of the cuartz insulation on the thermocouple wire was calculated from a measurement of its The overall uncertainty in the volume of the system at diameter and length. any pressure was estimated to be - 2 µl. which varies from - 0.4% to = 0.2% depending on the pressure in the system. The volume of the cuartz sample tube was approximately 1/4 ml and the cold trap and manometer to the reference point another 1/4 ml. At a pressure of approximately 100 mm. the total volume of the system was equal to 1 ml.

The capillary depression of mercury in tubes of 3.6 mm diameter is a large offect which must be corrected for. The system was pumped to  $< 10^{-3}$  mm and the mercury levels of the manometer road, including the height of the menisci. A series of readings at zero pressure with different menisci heights were obtained and the correction to be applied determined from these. It is estimated that

the pressure read from the manometer should be accurate to  $\frac{+}{-}$  0.2 nm for all cases and nearer  $\frac{+}{-}$  0.1 mm for most.

The weight of oxygen in the system was calculated from the pressure at some known temperature, the density of oxygen at this temperature, and the volume of the Since a cold trap was added to the system, cooling only system at this pressure. a fraction of it, and the part containing the sample was heated to different temperatures, it was necessary to determine the extent of these two effects. This was done empirically by first filling the system with a known pressure of oxygen, then adding the cold trap and measuring the decrease in pressure. This was done over the range of pressures that would be worked with and the results plotted as % correction versus pressure in mm of Hg, giving a straight line. Calculations of the corrections were in close agreement with the measured values. The corrections for the increase in pressure due to heating the sample were determined in the same way. A certain pressure of  $O_2$  was admitted to the system with the cold trap in place, and then the temperature raised over the desired This was done for a series of pressures. A series of straight lines rango. were obtained by plotting T vs. % correction, one line for each pressure measured. The correction for the measured pressure could be obtained from this plot by interpolation. The over -all accuracy of these corrections probably falls within + 0.5%.

The decomposition,  $2 \text{ CuO} = \text{Cu}_2 + 1/2 \text{ O}_2$ , was carried out in an apparatus of this type. The AH value obtained by plotting 1/T versus log P agreed with the value found by other workers<sup>3</sup> to within 0.2%, indicating that the system is capable of giving good results with samples of the order of a few milligrams.

When used for the decomposition of americium oxide, the system was surrounded by a box made from plywood and lucite, fitted with sliding doors on the front and connected to the exhaust system of the room so that air was constantly being drawn through the box.

#### III. Preparation of Sample

The praseodymium used in this investigation was obtained from Johnson-Matthey and Co., Ltd. of London, and was of their best quality. Spectroscopic analysis showed several percent of sodium and potassium and less than 1% of other rare earths. This material was further purified by R. C. Lilly and D. C. Stewart of this laboratory by means of ion exchange methods, yielding a praseodymium oxide that was pure within the sensitivity of the spectroscopic method.

#### Table I

#### Spectrographic Analysis of Praseodymium reported as unicrograms per fifty micrograms of sample (ND = not detected)

A1	-	ND	Gd	ND< 0.1	Ta	-	ND	< 0.5
Ba		ND	Ho -	ND <0.1	Tb	-	ND	< 0.1
Be	-	ND < 0.005	К -	ND < 0. Ł	Yb	-	ND	<0.1
Ca	-	< 0.01	LA -	ND < 0.1	Y	•	ND	<0.1
Ce	-	ND < 0.1	Lu -	ND <0.1				
Dy	-	ND < 0.1	Na -	ND < 0.01				
Er	••	ND <0.1	Nd -	ND < 0. 0]				
Eu	-	ND < 0.1	Sm -	ND <0.1				
Fe		ND < 0.1	Sr -	ND <b>≼</b> 0.01				

The americium used was obtained from waste solutions left from the purification of plutonium at Los Alamos. It was separated by long and arduous procedures to give, for the final product, a spectroscopically pure solution of americium. This americium was precipitated as the oxalate, and ignited to give AmO<sub>2</sub>.

#### Table II

#### Spectrographic Analysis of Americium

reported as micrograms per fifty micrograms of sample (ND = not detected)

Al		< 0.01	К -	ND < 0.1
Be	-	ND <0.005	Mn -	ND < 0.01
Ca	-	0.02	Na -	ND < 0.01
Co		ND < 0.1	Ni -	ND < 0.01
Cr	-	ND <0.01	Pb -	ND <0.1
Fe	-	ND <0.1		

In the case of praseodymium, the uncertainty in the composition of the compound usually designated as " $Pr_6O_{11}$ ", made it designable to use  $Pr_0$  as the primary standard for weighing. The sesquioxide was obtained by hydrogen reduction of the higher oxide at 500° C. This was weighed in an air-tight platinum capsule. In one case, the loose salt was removed in a dry N2 atmosphere and its This weight was known to ± 20 µg for an S mg weight obtained by difference. sample of Pr203. This salt was added by means of a platinum funnel to the guartz sample tube of the apparatus. This tube was then vigorously tapped until all the oxide appeared to drop to the bottom of the tube. Due to the uncertainty as to whether all the salt had fallen to the bottom of the tube and due to the interaction of metal salts and SiO at high temperatures, another procedure was adopted for a second sample. A small platinum tube about 1.5 cm long and about 1 mm O.D. was fabricated from 1/2 mil platinum foil. This tube was placed inside the larger platinum weighing vessel and subjected to the treatment to be used for the H<sub>2</sub> reduction, then capped and weighed. It was opened and the small inner tube removed, then recapped and weighed again, giving the weight of the tube by difference. After constant weight was obtained for the tube, it was filled with

previously prepared  $Pr_2O_3$ , treated with H<sub>2</sub> at 500° C, and the weight of the salt found by difference. This procedure was repeated until constant weight was obtained. The weight of about 5 mg of  $Pr_2O_3$  was known to  $\frac{1}{2}$  6 µg. The small tube was placed in the guartz sample tube and allowed to fall down to the bottom. In this case all of the oxide was in the zone that was to be heated.

Due to health hazards involved in the manipulations necessary to carry out the reduction, it was decided to weigh the americium oxide as  $AmO_2$ . The composition of this compound seems well established. No change in lattice constants for the oxide were detected after treatment of the compound under 50 atmospheres of  $O_2$  at 200° C<sup>4</sup>.

A small platinum tube was weighed repeatedly after being treated in the same manner as was to be used for the weighing of the americium oxide. After constant weight was obtained, the tube was filled with the americium oxide, heated to  $700^{\circ}$  C and cooled slowly to  $250^{\circ}$  C where it was held for 15 min. to allow maximum absorption of  $0_2$ . It was then put in the weighing capsule and capped. After checking by the health group for outside contamination, it was weighed. The vessel was opened, the inner tube removed and transferred to the cuartz sample tube. The empty outer vessel was again weighed and the weight of  $AmO_2$  determined by difference. A 5 mg sample of  $AmO_2$  was weighed is  $\frac{t}{2}$  30 µg.

All weighings were made on the Ainsworth F D J micro-balance, with the exception of the second praseodymium oxide sample called Sample 2 of Series 1 which was weighed on the American Balance Corp. semi-micro-balance sensitive to 10 µg.

#### IV. Experimental Results

The oxide decomposition data obtained from these studies is presented as a sories of plots of the log of the pressure in mm of mercury versus the reciprocal of the absolute temperature. (Figs. I through IX).

The various samples and apparati used were as follows:

#### Pr Oxide Decomposition

Series 1 -- Preliminary apparatus - approximate calibration.

Sample 1 --  $8.037 \stackrel{+}{-} 0.020$  mg weighed as  $Pr_2^{0}_{3}$  directly into the quartz tube in the form of pellets. (Fig. I).

Sample 2 -- 8.11 - 0.30 mg weighed as Pr<sub>6</sub>O<sub>11</sub> directly into the cuartz tube as a powder. (Fig. II).

Series 2 -- Improved apparatus as shown in drawing - Accurately calibrated within the limits given in Section II.

Sample 1 -- 8.76 <sup>±</sup> .020 mg weighed as Pr<sub>2</sub>O<sub>3</sub>, added directly to quartz sample tube (Figs. III, IV, V, VI).

Sample 2 -- 5.145  $\stackrel{+}{-}$  0.006 mg weighed as  $Pr_2O_3$  in small Pt tube which was then placed in the sample tube (Figs. VII, VIII, IX).

#### Am Oxide Decomposition

Same apparatus as in Series 2 above.

Sample 1 -- 5.250 ± 0.030 mg weighed as AmO<sub>2</sub> in small Pt tube which was then placed in the sample tube. (Figs. XIII and XIV).

Since the volume of the system was known at any pressure of O2, the composition of the oxide could be calculated. Due to the fact that in some cases oxygen appeared to react with substances within the system and also due to outgassing of the absorbed gases in the fused silica tube, the composition of the oxide is not known with high accuracy, particularly at temperatures above  $1,000^{\circ}$  C. The compositions given on Fig. XI represent the best values obtained from the average of

several different runs. Compositions from a decomposition run which did not agree with any other runs were not considered in this compilation. Compositions given to only two decimal places are from only one or two separate runs and the degree of error is quite uncertain.

The x-ray data were obtained only for the praseodymium oxide decomposition. One set of samples was taken at  $PrO_{1.66}$  and another at  $PrO_{1.73}$ . These data along with lattice constants of  $Pr_6O_{11}$ ,  $PrO_2$ , and  $Pr_2O_3(0 \text{ form})$  from measurements<sup>5</sup> made in these laboratories are presented in Fig. XII. The lattice constants given for the body-centered cubic structures are actually one-half of the true lattice constant. Another sample was taken at what corresponded to  $PrO_{1.53}$ but was inhomogeneous yielding hexagonal  $Pr_2O_3$  and a fluorite structure with the same lattice constant as that of  $PrO_{1.73}$ . This sample was cooled very rapidly which probably explains the inhomogeneity.

The composition of the higher oxide of americium was determined as follows: The higher oxide was reduced with  $H_2$  at 550° C to  $Am_2O_3$ . Oxygen was then admitted to the system but in the first instance reaction occurred so rapidly that the pressure of oxygen could not be accurately measured and only an approximation of the pressure could be obtained. This observation gave a minimum O:Am ratio of 1.95 to 1.0. A second experiment where the reduction was carried out at 750° C followed by heating one hour under high vacuum at 990° C, gave a reduction product which did not react rapidly with oxygen at room temperature. This allowed a more accurate measurement to be made. The result of this second measurement gave an oxygen to americium ratio of 1.95  $\stackrel{+}{-}$  0.02 to 1.

#### The AH of the decomposition

Slopes of the various curves were measured by visually drawing the best straight line through the points of the log p vs. 1/T plots and those curves were

used which had the best defined slopes. By plotting the points at which points of inflection at the beginning of curves F and H of Fig. XI first appeared for all runs at many different pressures, two straight lines were obtained on which most of the points fell. (Fig. X) The slopes of these two curves were calculated from this collected data. The lines of lesser slope (Curves A, B, C, D and E. Fig. XI) contributed only small amounts to the  $\Delta H$  value and thus small inaccuracies in measurement of the slope were unimportant in the  $\Delta H$  for the total reaction. The slopes of the curves beyond H were obtained from runs where only relatively small amounts of oxygen were present in the system and the decomposition could be effected at lower temperatures. Slight inaccuracies in these curves would cause large errors in the  $\Delta H$  value as would errors in the composition at these points of inflection. The value of AH calculated using these slopes and compositions gave 47  $\div$  5 kcals. as the AH for the decomposition of  $Pr_{60}_{11}$  to  $Pr_2O_2$  over a temperature range of 400° to 1200° C. The AH obtained from heats of solution given in Bichovsky and Rossini<sup>6</sup> is 48 kcals. A further check on these values was the measurement of the heat of solution of  $Pr_2O_3$  by L. Eyring<sup>4</sup>, which agreed with the previous value. The conclusion is drawn that for much of the decomposition, at least, equilibrium pressures were measured.

### Table III

Phaso	Fraction of Total Reaction	log P1/P2	1/T <sub>2</sub> - 1/T <sub>1</sub>	-AH(kcal) for frac- tion decomposed
А	0.021	0.439	0.30	0.157
В	0.039	0.363	0.20	0.775
C + D*	0.093	0.439	0.30	0.624
F	0.201	0.512	0.15	3.142
Н	0.171	0.924	0.06	12.053
I	0.144	0.494	0.20	1.628
J	0.210	0.823	0.03	26.376
K	0.120	1.2505	0.30	2.292
Total	0.999			47.05

ΔH Values for the Docomposition of Pr6<sup>0</sup>11 Letters refer to Fig. XI

\*Inclusion of the slope of curve E causes an increase of 0.3 kcal in the total  $\Delta H.$ 

#### V. <u>Discussion</u>

One original objective of this study was to obtain a  $\Delta H$  value for the thermal decomposition of americium dioxide to americium sesquioxide. By assuming that there is the same difference between the heats of formation of  $AmO_2$  and  $Am^{+4}(aq)$  and  $1/2 Am_2O_3$  and  $Am^{+3}(aq)$  as there is for the corresponding plutonium compounds and ions and that the  $\Delta S$  values are the same, a value for the potential of the III-IV couple could be estimated.

Although the praseodymium-oxide system proved very complex and there was a lack of rapid, reversible equilibrium pressures over certain portions of the system, a reasonably accurate approximation of the  $\Delta H$  for the decomposition of Pr60<sub>11</sub> to Pr<sub>2</sub>0<sub>3</sub> was obtainable. It was significant that a  $\Delta H$  based on an idealized picture of the system agreed with the  $\Delta H$  obtained from heats of solution to within the experimental error.

Unfortunately the thermal decomposition of  $AmO_2$  could be carried out only to  $AmO_{1.85}$  at the temperatures possible with this apparatus. Within the range of decomposition investigated ( $AmO_2 \longrightarrow AmO_{1.35}$ ),  $\Delta H$  for the total reaction assuming the  $\Delta H$  of the partial reaction amounts to 21 kcals. This may be compared with 49 kcals from estimations based on the heat of solution of  $AmO_2$ <sup>4</sup>.

The data presented in preceding sections show conclusively that the course of the thermal decomposition

 $Pr_6^{0}_{11} = 3Pr_2^{0}_3 + 0_2$ 

is highly complex. The most interesting feature of this reaction is that it suggests that it is possible to have immiscibility of oxides of verw similar crystal structure and composition. In support of this hypothesis the data of McCullough showing immiscibility of  $Pr60_{11}$  and  $Pr0_2$  may be presented.5

Although no final interpretation of the data can be made until the system is examined by means of a high temperature x-ray camera, a tentative explanation based on meagre x-ray data taken at room temperature may be advanced. From the fact that both  $PrO_{1.333}(Pr_6O_{11})$  and  $PrO_{1.73}$  have fluorite structures, it would appear that all intermediate compositions have the same fluorite type structure.

The regions between Prol. 833 and Prol. 732 on the one hand and Prol. 715 and Prol. 51 on the other would appear to be regions of continuous solid solution formation, although if this is the case the P-T relationships are of an unusual nature. There is a certain inertia in the system in the area C, the upper curve being obtained by continually raising the temperature until the curve D is reached. Upon cooling from the point of intersection of C and D, the absorption of oxygen follows the lower dotted line which has the same slope as D and is reversible in that region. Again, as decomposition proceeds further, the curves C and D are followed up to the point of inflection between D and F, which apparently represents a limit of solid solution formation decomposition. The curve E was obtained in only two of the many decompositions effected and may represent another possible path for the decomposition. The loss of oxygen atoms as the decomposition proceeds eventually results in the formation of a new phase with a decomposition slope H. The actual experimental curve F shows a considerable curvature as it approaches curve G which may be an indication of solid solution formation. However, x-ray evidence shows only one prominent phase at Prol.73 which is near the composition Proj 715 at which the horizontal curve G intersects curve F.

The structure obtained at  $PrO_{1.73}$  is the fluorite type with a coordination number of eight for the metal ion, while the structure at  $PrO_{1.66}$  is of the C type with a coordination number of 6 for the metal ion. The pressure of oxygen over the new phase represented by the decomposition curve H seems to be less than 1 mm of Hg at about 650° C. This was shown by the data presented in Fig. I, Runs A and B, where the oxygen pressure was lowered to zero by evacuation of the system at a temperature just high enough to have decomposed all of the previous phase, i.e. to the right of curve G. The pressure of oxygen then increased along the curve obtained in run B, Fig. I, which could be extrapolated to intersect the point of inflection between G and H and which had the same slope as curve H of Fig. XI.

Since the C type structure occurs at  $PrO_{1.66}$ , and a C type structure is known for  $PrO_{1.5}(Pr_2O_3)$  it is probable that intermediate compositions also have the C type lattice. At high temperatures, the final product,  $PrO_{1.5}$ , has the A type structure (hexagonal) and presumably a transition to this lattice type must occur. From the data available, it is impossible to tell where this transition does take place. There are three points of inflection between phase H and tho final phase  $Pr_2O_3$ , all having different slopes. As the temperature is increased, the same sort of decomposition takes place for the fluorite structures.

The fact that the compositions of oxide existing at the intersections of the various straight lines conform to no simple stoichiometric ratios of  $Pr_2O_3$  to  $PrO_2$  seems analogous to the situation which has been demonstrated for many different oxides such as those of Ti, V, Mo, W, Mn, Cr, Ni, and Pb oxides<sup>7</sup>.

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VII. <u>Bibliography</u>

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FIG. XI

FIG. XII







FIG. XIV