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ALKALI OXIDE-TANTALUM, NIOBIUM AND ANTIMONY OXIDE IONIC CONDUCTORS

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SUMMARY

This report summarizes work carried out between January 1, 1974 and December 31, 1974 under an agreement with the National Aeronautics and Space Administration, Lewis Research Center (Interagency Agreement PR545160) to study the phase equilibria of alkali oxide-tantalum, niobium and antimony oxide systems and synthesis of phases which might have interesting ionic conductivity. It also includes work performed since the last contract (Interagency Order C-29933-C [1]) which ended in October 1973, as well as that done since the formal ending of the present contract and the issuance of this summary report.

In addition to the six systems reported in the previous contract (Interagency Order C-29933-C [1]) the phase equilibrium relations of four additional systems were investigated in detail. These consisted of sodium and potassium antimonates with antimony oxide and tantalum and niobium oxide with rubidium oxide as far as the ratio $4Rb_20:11B_20_5$ (B=Nb, Ta). The ternary system NaSbO₃-Sb O_4 -NaF was also investigated extensively to determine the actual composition of the body centered cubic sodium antimonate. In addition, various other binary and ternary oxide systems involving alkali oxides were examined in lesser detail. The phases synthesized were screened by ion exchange methods to determine mobility of the alkali ion within the niobium, tantalum or antimony oxide (fluoride) structural framework.

Five structure types were found to be of sufficient interest to warrant further investigation. These structure types are (1) hexagonal tungsten bronze (HTB) (2) pyrochlore (3) the hybrid HTB-pyrochlore hexagonal ordered phases (4) body centered cubic antimonates and (5) $2K_20:3Nb_20_5$. Although size of these phases exhibit good ion exchange properties only the pyrochlore in the Sb_20_4 -NaSb0₃ system has so far been prepared with Na⁺ ions as an equilibrium phase and as a low porosity

- 3

ceramic. Unfortunately Sb^{+3} in the channel apparently interferes with ionic conductivity in this case, although relatively good ionic conductivity was found for the metastable Na⁺ ion exchanged analogs of RbTa₂O₅F and KTaWO₆ pyrochlore phases. Small crystals of the other phases can generally be prepared by flux techniques and ion exchanged with Na⁺. However, in the one case where congruency allows large crystals to be pulled from the melt (4Rb₂O:11Nb₂O₅) ion exchange techniques up to \sim 450°C are not sufficient to accomplish replacement with Na⁺ ions.

1.0 INTRODUCTION

The program described in the present report involves the preparation of single crystals and ceramic specimens of materials which have passed some preliminary screening tests and appear to be potential candidates as solid ionic conductors. Much of the preliminary screening was done under previous work order (No. C-29933-C) [1]. Furthermore the phase equilibrium diagrams have been studied for some systems in which promising candidates have been discovered but for which no phase data existed.

Crystals and ceramic specimens were prepared or synthesis attempted for various different structure types and compositions, including the hexagonal tungsten bronze and pyrochlore phases in the system $K_2O-Ta_2O_5-WO_3$, the $2K_2O:3Nb_2O_5$ type phase and the phases in the system $Rb_2O-Nb_2O_5$. In addition specimens of the pyrochlore phase and the body centered cubic phase of the $NaSbO_3-Sb_2O_4-NaF$ system were examined.

The phase equilibria diagrams of the $\text{Rb}_2\text{O}-\text{Nb}_2\text{O}_5$ and $\text{Rb}_2\text{O}-\text{Ta}_2\text{O}_5$ systems were examined as well as those of the $\text{Sb}_2\text{O}_4-\text{NaSbO}_3$ and $\text{Sb}_2\text{O}_4-\text{KSbO}_3$. In addition the equilibria in the system $\text{Sb}_2\text{O}_4-\text{NaSbO}_3-\text{NaF}$ were investigated. Other compositions examined included the alkali rare earth oxide systems and alkali oxide-bismuth oxide systems although no high temperature equilibrium phases were found in any of these systems.

The method of flux synthesis was used to prepare small single crystals for the first time of the pyrochlore phase $RbTa_2O_5F$ and the body centered cubic phases of F stabilized potassium and sodium antimonates.

In the following discussion all ratios (1:3, 3:5, etc.) refer to the alkali/metal ratio rather than to the particular starting material that may have been used.

2.0 DISCUSSION OF RESULTS

2.1 The System Nb₂O₅-4Rb₂O:11Nb₂O₅

The system $\text{Rb}_2\text{G-Nb}_2\text{O}_5$ was investigated from the region of about 27 mole % Rb_2O to the Nb_2O_5 end member. Ten compositions were prepared by dry mixing stoichiometric amounts of Rb_2CO_3 and Nb_2O_5 and calcining in open Pt crucibles at 500°C and 600°C for extended periods of time with intermediate grindings. The specimens were heated at higher temperatures in sealed Pt tubes in resistance wound quench-type furnaces (Table 1). The x-ray diffraction patterns of these quenched specimens were used to establish the phase equilibrium diagram shown in Figure 1.

Single crystals of the eleven layer (11-L) $(4Rb_20:11Nb_20_5)$, hexagonal tungsten bronze (HTB) $(21.75Rb_20:78.25Nb_20_5)$ and Gatehouse tungsten bronze (GTB) $(11.5Rb_20:88.5Nb_20_5)$ phases (see reference [1] for nomenclature of phases) were grown from the melt by modified Czochralski techniques. Single crystal fragments of the 11-L phase were used in ion exchange experiments but did not provide better results than isostatically pressed pellets prepared from calcined powder.

Unit cell dimensions for all phases are given in the summary table of x-ray data at the end of this report (Table 12). They were determined from both single crystal precession patterns and x-ray powder diffraction patterns (CuKa radiation, Ni filtered) with the aid of a computer least squares program. Only the phase occurring at low temperatures at about 15 mole % $Rb_{2}O$ has not been indexed, as no single crystals were obtained.

2.2 The System Ta₂O₅-4Rb₂O:11Ta₂O₅

The system $\text{Rb}_2\text{O}-\text{Ta}_2\text{O}_5$ was investigated from 26.67 mole % Rb_2O to 100 mole % Ta_2O_5 . Eight compositions were prepared and the system was investigated by conventional quenching methods. The samples were heated in open Pt trays at 500° and 600°C and then heated in sealed

Pt tubes at higher temperatures. The phases were identified by comparison with known structures in the $K_2O-Ta_2O_5$ and $Rb_2O-Nb_2O_5$ systems. Equilibria was assumed to have been established when successive heat treatments showed no change in the x-ray diffraction patterns. As in the $K_2O-Ta_2O_5$ [1,2] and $Rb_2O-Nb_2O_5$ systems a layer sequence 9-L, 16-L, 11-L (in order of increasing alkali content) was found to be present. The HTB and GTB-like phases [1] were also found to be present. The results of the experiments are given in Table 2 and the phase equilibrium diagram interpreted from this data is shown in Figure 2.

2.3 The System Sb₂O₄-NaSbO₃

The system between the compositional limits of $NaSbO_3$ and Sb_2O_4 has been examined in detail. Thirteen compositions were studied by air by conventional quenching techniques and the materials were examined by single crystal, powder, and high temperature x-ray diffraction techniques. The phase equilibrium diagram, Figure 3, has been constructed from the data given in Table 3. When Sb_2O_4 is reacted at low temperature (\sim 750-1000°C) with alkali carbonate it generally looses the CQ₂ and picks up oxygen from the atmosphere to satisfy equilibrium conditions of the phases formed, which may involve oxidation of the antimony ions. Therefore the antimonate systems reported here are not strictly binary.

The compound NaSbO₃ (ilmenite-type) was found in this work to melt at about 1555±5°C. An intermediate pyrochlore solid solution exists from about 37.5 mole % Na₂0:62.5 mole % Sb₂O₄ to 24 mole % Na₂0:76 mole % Sb₂O₄ at 1350°C. The 1:3 composition probably does not really correspond structurally to [NaSb⁺³]Sb₂⁺⁵O₇ although the 3:5 composition may be written as [Na_{1.5}Sb_{.5}]Sb₂⁺⁵O_{6.5} see Section 3.2. The 3Na₂O:5Sb₂O₄ composition apparently melts congruently at 1490±5°C. The solidus curve falls from this temperature to about 1340±5°C at 24 mole % Na₂O:76 mole % Sb₂O₄. A two phase region exists between the pyrochlore solid solution and Sb₂O₄. An unknown phase was found to occur in the system which could

be made approximately single phase by calcining the composition 15 mole % $Na_20:85$ mole % Sb_2O_4 at 750°C and reheating in a scaled Pt tube to 1000°C for 64 hours in the presence of PtO₂. This phase has an as yet unindexed x-ray diffraction pattern with the four strongest lines occurring at <u>d</u> values equal to 3.283, 2.798, 3.453, 8.23A.^{1/}

2.3.1 NaSb0,

The compound Na₂0:Sb₂0₅ was first reported to occur by Schrewelius [4] and to be hexagonal with a=5.316 and α =15.95A and an ilmenite structure. This compound was found in the present work to melt congruently at about 1555±5°C. No other stable polymorphs were encountered.

2.3.2 Pyrochlore Solid Solution

One intermediate phase, a cubic pyrochlore solid solution mentioned by Stewart and Knop [5], was characterized in the system. The compositional varies from approximately Na20:3Sb204 to 3Na20:5Sb204 with unit cell dimensions varying from 10.289 to 10.286A respectively. Since the pyrochlore is a tunnel structure and since this pyrochlore is the only one reported that contains sodium that can be formulated by direct synthesis it was worthy of further study as a possible ionic conductor. For ionic conductivity measurements dense materials were needed and several experiments were conducted with Na₂0:Sb₂0₄ (1:2) in an effort to determine its stability under high pressure and temperature. In the first experiment, 2.25 g of single phase pyrochlore material were prepressed at 10,000 psi, $\frac{2}{}$ packed into a Pt cylinder 3/8 0.D. x 1 1/2" long with PtO₂ and sealed. This sealed crucible was heated to 1000°C and 69,800 psi with catastrophic results. Apparently this phase is not completely stable in this temperature-pressure range and some of the material reacted with the Pt container resulting in a melting of the Pt. The remaining

 $\frac{1}{10\text{\AA}}$ = 1.0 nanometers

 $[\]frac{2}{1}$ The use of psi, bar, and kbar follows the current common practice of workers in the field. Note that 1 bar = 10^5 N/m^2 (or pascal) = 10^6 dyn/cm^2 = 0.9869 atm = 14.504 psi. The accepted international standard (SI) unit of pressure is the pascal or newton per meter squared.

specimen, slightly depleted in antimony oxide, was found by x-ray diffraction to contain a pyrochlore phase plus ilmenite (1:1).

The next experiment was scaled down in amount of material and pressure. A few milligrams of material were sealed in a 3 mm diameter Pt tube and heated to 700°C and 4,000 psi for 1 hour in the presence of PtO₂. The material remained 1:2 (pyrochlore) with slight sintering. A similar experiment was conducted at 1000°C and 4,000 psi for 2 hours. The PtO₂ was decomposed and there was grain growth. The x-ray pattern was of 1:2 pyrochlore. In the next experiment, two samples in sealed platinum tubes were heated at 1100°C and 5,000 psi for 3 hours. Platinum oxide was added to only one of the tubes before sealing. The specimen without PtO₂ was a single phase pyrochlore which appeared to be a very dense material. The measured density was 5.21 g/cm³. The specimen containing PtO₂ was not dense.

In the last experiment two 5 mm diameter x 1 1/4" long tubes were flattened (to a rectangular shaped cross section, 1.5 mm thick) packed with material, sealed and heated to 1100°C and 4,000 psi. After heating, this specimen was very thin and brittle and the specimen fragmented when the Pt was mechanically removed. Again the specimen was quite hard with a measured density of 5.26 to 5.29 g/cm³. The average density of four measured fragments was $5.26\pm.05$ g/cm³.

For ionic conductivity measurements, prilets of $Na_2O: 2Sb_2O_4$ (3/4" in diameter) were placed in sealed platinum fuil envelopes and hot pressed by a commercial company at 1100°C and 5,000 psi. The pellets were well formed single phase material with a density of 96% theoretical (See Section on Mechanisms of Non-stoichiometry 3.2). The ionic conductivity of these pellets was measured at NASA Lewis Research Center (LeRC) [6] and they were found to be essentially insulators.

The distribution of the various ions (i.e. Na⁺, Sb⁺³, Sb⁺⁵, 0^{-2}) in the Na₂0:2Sb₂0₄ has not as yet been determined from single crystal structure analysis. The structure of this material is currently

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being determined at NBS. Until the results of this analysis are forthcoming it may be assumed that the "lone pair" electrons associated with Sb^{+3} will not allow the passage of Na⁺ through the channels.

2.3.3 Polymorphism of $Sb_2O_4^{3/2}$

Two stable polymorphs of Sb_2O_4 have been reported in the literature. They are $\alpha-\text{Sb}_2O_4$ which is orthorhombic [7] a=5.436, b=11.76, c=4.810A and $\beta-\text{Sb}_2O_4$ which is monoclinic [8] a=11.905, b=4.834, c=5.383A and $\beta=101^\circ22^\circ$. From Table 4(a) it can readily be seen that specimens quenched from a temperature composition region represented on the phase diagram, Figure 3, as $\text{Sb}_2O_4 + \text{pyrochlore may contain either } \alpha-\text{Sb}_2O_4$ and/or $\beta-\text{Sb}_2O_4$ when quenched from high temperatures and ambient pressures and examined at room temperature. From this seemingly inconsistent data it would appear that the $\alpha-\text{Sb}_2O_4$ form and $\beta-\text{Sb}_2O_4$ form have a polytypic relationship. To help resolve this problem a high resolution electron microscope study should be done.

From the date in Table 4(b) it appears that the β form is the equilibrium high pressure form of Sb_2O_4 . Insufficient data has been collected to establish if an equilibrium boundary curve exists between $\alpha - \text{Sb}_2\text{O}_4$ and $\beta - \text{Sb}_2\text{O}_4$ at various temperatures and pressures. When specimens are sealed and heated under pressure in the presence of PtO_2 in either Pt or Au tubes single phase $\beta - \text{Sb}_2\text{O}_4$ is obtained. However when heated under pressure without the PtO_2 , a two phase specimen results, $\beta - \text{Sb}_2\text{O}_4$ and the dense high pressure form of Sb_2O_3 (valentinite). A similar polytypic relationship probably exists for the two polymorphs of Sb_2O_3 .

Some samples of Sb_2O_4 will be sent out of house for electron diffraction studies in an attempt to establish the polytypic relationship of $\alpha - \text{Sb}_2\text{O}_4$ and $\beta - \text{Sb}_2\text{O}_4$. Some additional experiments will be conducted to establish the high pressure boundary curve between α and $\beta - \text{Sb}_2\text{O}_4$.

 $[\]frac{3}{1000}$ Most of the polymorphism study of Sb₂O₄ was conducted prior to the issuance of Interagency Agreement PR545160, and is included in this report for the sake of completeness.

2.4 The System Sb₂O₄-KSbO₃

This system has been examined between the compositional limits of KSb03 and Sb204. Sixteen compositions were studied by conventional quenching techniques and the materials were examined by powder and by single crystal x-ray diffraction techniques. Compositions which were initially calcined at 500 and 700°C for 60 hrs, were dried at 230°C for 1 hr, sealed in Pt tubes and reheated. The results of the experiments are given in the data presented in Table 5 from which the subsolidus relationships have been established as shown in Figure 4. The compound KSbO2 with an ilmenite structure has been previously reported [9]. A body centered cubic solid solution phase originally reported as KSb03 [9] has been found to occur metastably at about 47.5% K_2^{0} . The $3K_2^{0}$:55b $_2^{0}$ 5 compound reported previously [10,11] was found in this work to melt congruently at about 1400°C. The K20:2Sb205 compound reported previously [11] was found in this work to have a phase transition at about 1000°C and to dissociate to pyrochlore plus 3K20:5Sb205 at about 1150°C. The low temperature form of this compound labeled P21/c represents a monoclinic phase with a=7.178A, b=13.378A, c=11.985A and β =124°10'. Single crystals of this phase were grown by flux evaporation from the composition 50K20:5Sb204:45MoO3. The unit cell and space group were determined from these crystals and confirmed by least square indexing of the powder diffraction pattern of the low temperature form of the compound K20:2Sb205. The pyrochlore solid solution exists at 1150°C from about 15 mole % $K_20:85$ mole % Sb_20_4 to greater than 30 mole % $K_20:70$ mole % Sb_20_4 . The melting characteristics of these phases have been partially determined and are currently being reported.

2.4.1 Hyroxyl Ion Stabilization of Cubic Potassium Antimonate

The compound KSbO₃ was reported previously [12] as being cubic at ambient conditions when previously subjected to high temperatures and pressures. In this work two experiments were conducted, heating single phase $KSbO_3$ ilmenite in sealed platinum tubes at about 750°C and 00,000 psi, utilizing argon as a pressure medium. The first of these, a two-hour run, showed only one small cubic line in the x-ray pattern but the specimen quickly reacted with atmospheric moisture. This experiment was then repeated for a longer time, 7 hours. In this time the material apparently reacted with the sealed Pt tube. The specimen was ground, mixed with petroleum jelly, and subjected to x-ray diffraction analysis before too much hydration could take place. The x-ray pattern showed lines of a material not seen by us previously. It may be concluded that this is a somewhat reducing environment and some of the material has transformed to K_2O and Sb_2O_3 which in turn melted and reacted with the Pt container.

The body contered cubic phase was not formed at the above stated conditions. However, occasionally small amounts of a cubic phase were seen in the x-ray powder diffraction pattern of KSbO, ilmenite heated at ambient pressure. For these reasons specimens of 1:1 and 3:5 mole ratios K20:Sb204 were equilibrated in air at 750°C for 60 hours and picked up oxygen to form the phases KSb03 and K3Sb5018 and then were reheated for one hour at 1200°C to drive off all excess moisture. X-ray diffraction patterns of these specimens showed single phase ilmenite and the 3K20:55b205 compound. Portions of these 1200°C calcines were then saighed and mixed in acetone in the appropriate ratios to yield compositions of 46, 47, 47.5, 48 and 49 mole % K20. Each of these specimens was dried at 240°C for one hour and heated in open Pt tubes at 1200°C. for one hour. Only the x-ray pattern of the '46% specimen showed a little 3K20:5Sb205, the others contained only the cubic phase. A new specimen of 48 mole % K₂O was prepared in the same way except the Pt tube was sealed. After one hour at 1200°C, the x-ray pattern of the specimen showed only about 50% cubic. A new specimen of 48% was prepared by weighing the 1:1 and 3:5 phases immediately after removal from the

1200°C furnace and sealing the material in a flattened Pt tube within 1-2 minutes. This tube was then inflated at 1200° for a few minutes and the material mixed by shaking in a "wiggle-bug". The sealed specimen was then heated for 64 hours at 1200°C. The resultant specimen had exceedingly large grain growth indicating considerable solid state recrystallization but showed <u>no</u> cubic phase. The conclusion is inescapable that access to atmospheric moisture is probably <u>necessary</u> for the formation of the cubic phase at atmospheric pressure.

A paper entitled "Flux Synthesis of Cubic Antimonates" was published by the present authors during the course of this work [13]. In addition to the discovery that the F⁻ ion stabilized the formation of the body centered cubic phase of potassium antimonate it was disclosed that the cubic antimonate could also be obtained by reacting $KSbO_3$ with a small amount of other cations with small radii like B⁺³, Si⁺⁴, etc. It now appears obvious that in this reaction the boron or silicon (etc.) actually ties up some of the K⁺ ion in a second phase and allows the K⁺ deficient antimonate to react with atmospheric moisture to form the cubic antimonate previously thought to be "KSbO₃".

2.5 The System Sb_2O_4 -NaSbO₃-NaF

2.5.1 The System NaSbOg-NaF

To determine if NaF additions will stabilize the body-centered cubic phase similar to the $6KSbO_3:KF$ -phase, NaF was added to $NaSbO_3$ in the ratio of $3NaSbO_3:NaF$, $4NaSbO_3:NaF$, $5NaSbO_3:NaF$, and $6NaSbO_3:NaF$. The x-ray diffraction patterns of these specimens contained varying amounts of body contered cubic phase, ilmenite, and NaF. After heat dg at 750°C and 1000°C in sealed Pt tubes, the x-ray patterns showed only ilmenite and NaF, however at $\sim 1150°C$ all the compositions contained some body centered cubic-type phase. The compositions $3NaSbO_3:NaF$ and $4NaSbO_3:NaF$, when heated in sealed Pt tubes at $\sim 1250°C$, did not contain ilmenite and

appeared to be the closest to single phase cubic. The small crystals of 4NaSbO₂:NaF prepared by quenching in a small sealed tube appeared to be well formed truncated octahedrons. However, the room temperature x-ray diffraction pattern of the material had somewhat diffuse lines with the exception of the h00 lines which were reasonably sharp, suggesting rhombohedral symmetry. This material was placed on a hot stage microscope slide and analyzed by x-ray diffraction from room temperature up to 220°C. At 190°C the material appeared to start to go cubic and by 220°C a good quality cubic x-ray diffraction pattern was obtained (a=9.353A). When the material was cooled to room temperature the symmetry was again non-cubic. As the hOO lines deteriorate somewhat on cooling, the true symmetry of the room temperature form is probably no higher than monoclinic or triclinic rather than rhombohedral. It was therefore not unreasonable to expect that a body centered cubic phase could be obtained by direct synthesis with NaF without the necessity for Na^+ ion exchange.

2.5.2 The Ternary System

X-ray diffraction patterns (single crystal and powder) of selected NaFflux synthesized (see Section 2.8.2.4) washed crystals show only a truly cubic bodyccentered phase (a=9.334A). It must be postulated that the composition formed by this technique is slightly different from that made essentially single phase at $4NaSbO_3:NaF$ in a sealed tube. In an attempt to obtain a fluorine-substituted body centered cubic phase which exists at room temperature the compositions shown in Table 6 were prepared and show the reported phases when quenched from $1250^{\circ}C$. Equilibrium was not obtained in overnight heat treatments at $1200^{\circ}C$. At $1350^{\circ}C$ the body centered cubic phase started to decompose. The composition $68NaSbO_3:4Sb_2O_4:$ 28NaF (mole percent) was chosen as the best composition for further studies on ceramic procedures (Section 2.7).

The phases found in the specimens heated at $\sim 1250^{\circ}$ C are summarized in "equilibrium" diagrams for the quaternary system NaSb0₃-Sb₂0₃-Sb₂0₅-NaF (Figure 5) and the ternary plane of this system NaSb0₃-Sb₂0₄-NaF (Figure 6).

2.6 Other Systems

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During the course of this study other alkali-oxide systems were surveyed, to decide whether interesting phases existed in these systems. Among those investigated were sodium and potassium bismuthates and sodium and potassium with several rare earth oxides, Nd_2O_3 , Sm_2O_3 , and Gd_2O_3 as well as with Y_2O_3 .

2.6.1 Alkali Bismuthates

Samples of sodium bismuthates and potassium bismuthates were prepared from the respective alkali carbonates and Bi_2O_3 and heated in open Pt tubes at 500°C for various periods of time (Table 7). X-ray diffraction patterns showed essentially single phase Bi_2O_3 with several small unidentified peaks in the $\text{Na}_2\text{O}-\text{Bi}_2\text{O}_3$ samples. These peaks did not correspond with those of commercial NaBiO_3 (apparently prepared according to Scholder and Stobbe [14]), which decomposes to Bi_2O_3 .

2.6.2 Alkali-Rare Earth Systems

Alkali carbonates were dry mixed with the respective rare earth oxides and heated as shown in Table 8. X-ray diffraction patterns were recorded with Ni filtered Cu radiation. A "?" indicates an unknown phase which is present as one or two small peaks in the pattern.

The only significant result of these studies appears to be the presence of the cubic rare earth structure in Gd₂O₃ well above its equilibrium transition. Apparently the alkali ion enters the rare earth oxide lattice to some extent, altering the polymorphic conditions.

2.6.3 Further Studies in the System Nb₂O₅:KNbO₂

During the interim between this contract and the previous one, it became apparent that the compound $2K_20:3Nb_20_5$ might have interesting ionic conductivity. However, as the exact ratio $40K_20:60Nb_20_5$ did not yield a single phase, new compositions were prepared (Table 9) and the composition $41K_20:59Nb_20_5$ was further studied. The conclusion that this is probably a non-stoichiometric phase led to a revision of the Nb $_20_5$ -KNb0 $_3$ phase diagram (Figure 7).

2.7 Pellet Fabrication

2.7.1 41K₂0:59Nb₂O₅ ("2:3", K₄Nb₆O₁₇)

Evaluation by LeRC of the never-hydrated single crystal of $K_4Nb_6O_{17}$ supplied by NBS resulted in a request for single phase, polycrystalline, never-hydrated pellets of this same composition.

The starting material was prepared by dry mixing KNb0_3 and Nb_2O_5 in appropriate amounts, followed by calcining at 800°C for 10 hours and 1000°C for 30 hours with intermediate grinding. ?ellets were pressed without binder at 10,000 psi in a 5/8 inch diameter steel die. Sintering was done in air at 1125°C for 18 hours. The pellets were removed from the sintering furnace at temperature, immediately placed in a silica glass tube, evacuated to p $\approx 2 \times 10^{-5}$ torr and sealed for transmittal.

The composition $2K_20:3Nb_2O_5$ ($K_4Nb_6O_{17}$) was observed to contain a small amount of the tetragonal tungsten bronze (TTB) second phase. This second phase was completely eliminated by using a starting composition of $41K_2O:59Nb_2O_5$. A large batch of this 41:59 K/Nb composition was prepared and successfully exchanged as shown in Table 10. A pellet of the unexchanged 41:59 composition was prepared and sintered at $1100^{\circ}C$ for 1 hour and subsequently exchanged. However, the problem of hydration with the 41:59 composition is much more severe than with the 2:3. It was necessary to maintain the sample at temperatures $\approx 100^{\circ} - 150^{\circ}C$ to obtain a powder diffraction

pattern and to our dismay, the exchanged pellet hydrated and decrepitated while being x-rayed in laboratory air. More elaborate precautions were necessary to successfully characterize pellets of this composition. It is not known if the lack of the TTB second phase increased the susceptibility to hydration or if this is due to a slightly higher than stoichiometric amount of K^+ (Na⁺ in the exchanged material) in the lattice.

A series of heating experiments was performed to determine the optimum sintering conditions for pellets of nominal composition $41K_20:59Nb_20_5$. At a temperature of $1100^{\circ}C$ overfiring of the pellet was pronounced, as evidenced by the presence of a small amount of a liquid phase and marked reduction of niobium. Attempts to reoxidize the sample by annealing at lower temperatures were not completely successful. Temperatures of 1000° and $1040^{\circ}C$ (the melting point of KNb0₃) did not result in sintering sufficient to develop any mechanical strength. A temperature of 1050° and 2 hours time was found to result in sound pellets. X-ray characterization of the material after this heat treatment showed only single phase "2:3".

As stated above, the problem of hydration in this composition is severe at all stages of the fabrication and exchange. The following schedule was adopted in order to minimize the exposure to laboratory air, with each subsequent fabrication step performed with a minimum of delay.

| Single phase 41K ₂ 0:59Nb ₂ 0 ₅ powder | → prepress → press 700 psi 20,000 | ically dry at → 220°C psi 0.5 hr | + |
|---|---|--|---|
| sinter at 1050°C → 2 hrs | place in fused quartz tube, evacuate to $p \le 10^{-5}$ torr and seal off | | |

A similar procedure was used for the pellets to be exchanged, except that at the completion of sintering, the pellets were placed individually in

NaNO₃ and exchanged at 450°C for 64-68 hours. At the completion of the exchange period, the excess NaNO₃ was removed by leaching in methyl alcohol, the pellets were dried at 200°C for 0.5 hours and then placed in a fused quartz tube, evacuated and sealed.

During the exchange period, the pellets appeared to suffer no physical degradation and x-ray powder diffraction of a sample obtained from the center of a test pellet showed that complete exchange to the Na⁺ phase had occurred.

Six pellets of the "2:3" K^+ phase and three pellets of the exchanged "2:3" Na⁺ phase have been submitted for evaluation.

2.7.2 4Rb₂0:11Nb₂0₅ (11-layer phase)

Pellets of the 4:11 composition were prepared by using 1 weight percent stearic acid as a binder, pressed in steel dies at 10,000 psi and sintered at 1200°C in air for 56 hours. The pellets were sound and showed little tendency to spall or disintegrate during exchange in molten KNO_3 (see Table 10).

2.7.3 Large Pellets for Ionic Conductivity Measurements

As mentioned several times, the problems of hydration and compositional changes resulting from volatility or change in oxygen content in the alkali niobates, tantalates and antimonates necessitate the use of rather extreme measures to produce sound pellets. The complete encapsulation of the starting composition at the earliest stage possible in the fabrication process appears to be successful in some cases. In such cases, the specimen remains encapsulated until ready for evaluation.

Two compositions were prepared for large dense specimens to be sent to LeRC, $41K_20:59Nb_20_5$ and $NaSb0_{3-x}F_x$. The $NaSb0_{3-x}F_x$ was prepared by holding $7Sb_20_4:93KF$ at $\sim 1000^{\circ}C$ for less than 10 minutes and then

separating the fines from the coarser grains by washing, settling, and centrifuging the insoluble product. The fines were then ion exchanged two times with NaNO₃ at 450°C - 2 hrs. The exchanged phase (NaSbO_{3-x} F_x) gave an x-ray pattern which was single phase cubic. The powders were pressed into 3/4" diameter pellets which were repressed isostatically at 20,000 psi. These pellets were placed in platinum envelopes, sealed on three sides and were again isostatically pressed to remove most of the air from the container. The envelopes were sealed and sent to a commercial company for isostatic hot pressing at ~1100°C and 5,000 psi. The 41K₂0:59Nb₂O₅ specimen was submitted to LeRC, unopened, in the platinum envelope. Duplicate specimens were used to determine physical integrity and phase composition of the pellets. The $NaSbO_{3-x}F_x$ specimen was not single phase after hot pressing, apparently as a result of decomposition under pressure. Three phases were evident: a cubic phase, the pyrochlore solid solution and NaF. It was felt that the exchanged product had a composition too close to a phase boundary with resultant instability under pressure. For this reason, it was decided to directly react the constituents to form the desired sodium containing phase directly and then hot press the product. A composition of 68NaSbO3:4Sb204:28NaF was chosen.

Attempts to fabricate a large, dense specimen of the composition 62NaSbO₃:4Sb₂O₄:28NaF were largely unsuccessful. Direct reaction of the components at 750°C for 60 hours and 1000°C fo: 1 hour in open containers, with periodic grindings, resulted in a product containing NaSbO₃ (ilmenite) and a trace of NaF. Calcining temperatures as high as 1265°C were required to form a single phase material of the desired cubic structure. A trace of NaF was always present as a second phase. The use of such high calcining temperatures results in a coarse-grained product which could not easily be ground fine enough to produce a sinterable powder. Attempts to hot press the specimens in platinum envelopes,

using the same technique described above for the 41:59 and $\text{KSbO}_{3-x}F_x$ specimens, at a temperature of 1250°C for 10 minutes were unsuccessful. It is recommended that if these experiments are repeated, only the lower temperature calcines (at 750° and 1000°C) should be used, and the final desired phase in the pellet be produced by appropriate choice of time and temperature of hot pressing.

2.8 Crystal Growth and Synthesis

Many of the phases discovered during the phase equilibria studies of the alkali tantalate, niobate and antimonate systems have not been previously reported in the literature. One of the objectives of the crystal growth portion of the study has been to synthesize and grow small single crystals for x-ray diffraction studies to provide structural information on these materials and assist in the interpretation of x-ray diffraction powder patterns. The second, and equally important objective, has been to grow single crystals of the potentially useful K^+ and Rb^+ phases for use in studies of ion exchange behavior for the purpose of obtaining Na⁺ phases. These studies are described in Section 2.9 of this report.

In general the choice of technique was dictated by the characteristics of the desired phase, i.e. congruent or incongruent melting, volatility of one or more components of the phase, range of coexistence with liquid, and desired final size of crystal. Both fluxed melt and Czochralski techniques were utilized, and in some cases, the flux evaporation technique was used for the preparation of small crystals.

2.8.1 Crystal Growth by Pulling from the Melt

2.8.1.1 Hexagonal Tungsten Bronze in the K₂O-Ta₂O₅-WO₃ System

Selective phase equilibria experiments with mixtures of $KTaO_3$ (KT) and WO_3 in the neighborhood of $KT:2WO_3$ ($K_2O:Ta_2O_5:4WO_3$) during the previous contract confirmed the existence of a hexagonal tungsten bronze (HTB)

structure. Single phase compositions were easily prepared between the pyrochlore composition, $K_20:Ta_20_5:W0_3$, and a composition $KTa0_3:3W0_3$ where a second, unidentified phase appeared. At the same time, exploratory crystal growth runs at lower Ta_20_5 contents were encouraging and gave some promise of adaptation to the accelerated crucible rotation technique (ACRT) for single crystal growth of this bronze structure. However, considerably more work is required to determine the liquidus surface and primary field extent before such experiments would be definitive.

A 50 gram batch of $26.6K_20:3.3Ta_20_5:70.1W0_3$, prepared from KT, K_2W0_4 and $W0_3$ was premelted by induction heating in a platinum crucible. The crucible was then covered and transferred to a growth furnace. Following a 10 hour soak at 1200°C, the temperature was lowered at 2.5°C hr⁻¹ to 500°C and the crucible removed from the furnace. Results were essentially the same as obtained in previously reported work with a cooling rate of approximately 30° hr⁻¹. Crystals were in general small, the largest ranging from about 6x6x0.2 mm to 3x3x0.1 mm. The yield, based on visual estimate, was small.

Because of the small total yield and extensive nucleation and growth of small platelets, an attempt was made to pull crystals of the HTB from the fluxed melt. A 250 gram charge of composition $27K_20:5Ta_20_5:68W0_3$ was placed in a platinum crucible in a growth furnace which was baffled to provide an essentially isothermal chamber except directly above the center of the melt where the platinum pull rod was inserted. The charge was held at 1225° for three hours prior to attempting growth. Although no evidence of solid was present on the surface of the melt, extensive freezing of polycrystalline material occurred when the pull rod was inserted. The temperature was raised in increments and the polycrystalline mass redissolved. All attempts to achieve growth were unsuccessful at temperatures up to 1250°.

In the final attempt, at 1265° C, crystallization on the pull rod was slow enough to allow a 16 hour pull. X-ray diffraction examination of the results indicated that the crystals obtained were the desired phase. However, physically, the crystals were in the form of small plates, bound together by the frozen melt. It appears that a considerable effort would be required to establish suitable conditions for the growth of larger single crystals of this HTB. This would probably require a minimum of 1-2 man years.

2.8.1.2 The System KNb03-Nb205

Single crystals of $K_4Nb_6O_{17}$ have rather easily been grown both at other laboratories [15] and at NBS. However, the material hydrates readily in laboratory air with subsequent degradation of the crystalline perfection, which is not regained by drying. In order to provide LeRC with a large single crystal which had never been hydrated, a crystal was grown from a melt of the same composition, removed from the puller while still at a temperature of $300-400^{\circ}C$ and immediately placed in a fused silica ampoule, evacuated to $p \le 2x10^{-5} \frac{4}{7}$ torr and sealed for transmittal. Later conversations with LeRC indicated that the behavior of this crystal was indeed different, and as described in the section on pellet preparation, never hydrated pellets were also supplied for evaluation.

2.8.1.3 The System Rb₂O-Nb₂O₅

A 100 gram batch of $29Rb_20.71Nb_20_5$ was calcined at 400° for 60 hours and used for crystal pulling experiments. Five single and/or polycrystalline boules of the 4:11 (Rb:Nb) (11-layer) hexagonal phase were pulled from the melt by the top seeded solution method. The crystals were obtained using both seeded and unseeded pull rods. These crystals were used as "purified" material for the growth of single crystal 11L for use as a stable end member of the system and for chemical analysis.

 $[\]frac{4}{1}$ The use of torr follows the current common practice of workers in the field. Note that 1 torr = 1.32 x 10⁻³ atm = 1.34 x 10² N/m².

The crystals appear to be susceptible to thermal shock and therefore must be cooled relatively slowly. Self-seeded crystals show a marked preference for growth perpendicular to the c-axis. At least one excellent cleavage (basal-plane) is evident.

Crystals of the hexagonal tungsten bronze (HTB) phase in this system (21.75Rb₂0:78.25Nb₂0₅) were grown by pulling from a melt of the same composition (experiments performed by C. Jones while on an American University Research Participation Program). Large, clear water-white crystals were easily obtained but all fractured into rather large blocks upon cooling. All attempts to prevent cracking were unsuccessful and the cause remains unknown, but may be related to a symmetry change to orthorhombic on cooling.

Crystals of the Gatehouse tungsten bronze (GTB) $(11.5Rb_20:88.5Nb_20_5)$ phase were grown from a melt of composition $16Rb_20:84Nb_20_5$ (experiments performed at NBS by D. Klein while on an American University Research Participation Program). Small single crystal boules were successfully grown, but again fractures developed as soon as the boules were removed from the melt at the completion of the run in spite of all attempts at slow cooling of the boule.

2.8.2 Flux Growth

2.8.2.1 The System Rb₂O-Nb₂O₅-MoO₃

Nine different compositions in this system were prepared (2 gm batches) and heated by induction to temperatures in the 900°-1100°C range in small platinum crucibles. Recrystallization and/or volatility of the MoO₃ resulted in the formation of small crystals. Crystals (2mm+) of the 11 layer hexagonal (4:11) phase were obtained from compositions of $30Rb_2O:5Nb_2O_5:65MoO_3$ and $30Rb_2O:10Nb_2O_5:60MoO_3$. A 27.5Rb₂O:10Nb₂O₅:62.5MoO₃ composition yielded small crystals of the GTB

phase ($11.5 \text{ mole } \text{X} \text{ Rb}_20$) and H-Nb₂O₅ and $28\text{Rb}_20:5\text{Nb}_20:67\text{MoO}_3$ yielded small single crystals of the HTB ($21.75 \text{ mole } \text{X} \text{ Rb}_20$). No crystals of the unidentified phase occurring at about 15 mole $\text{X} \text{ Rb}_20$ in the binary system were obtained with any MoO₃ flux composition. This is further reason to believe this phase may not be an equilibrium compound in the binary system.

2.8.2.2 The System Rb₂O-Ta₂O₅-MoO₃

Six different compositions were prepared in the ternary system for flux evaporation crystal growth. Single crystals of the 9 layer phase (1:3) were grown using compositions of $30\text{Rb}_20:10\text{Ta}_20_5:60\text{MoO}_3$ and $35\text{Rb}_20:5\text{Ta}_20_5:60\text{MoO}_3$. Crystals of the other phases in the binary system $\text{Rb}_20-\text{Ta}_20_5$ could not be obtained using this method, probably because their primary fields do not extend to such high MoO₃ contents, or to the relatively low temperatures involved.

2.8.2.3 The System Rb₂O-Ta₂O₅-RbF

In view of the success obtained by the flux synthesis route in the $KSb0_3$ -KF system [13] similar attempts were made using RbF and Ta_2O_5 . A 95RbF:5Ta_2O_5 composition was completely liquid at about 925°C and on cooling yielded acicular crystals of an unknown phase. A 75RbF:25Ta_2O_5 composition was heated to about 1150°C for 30 seconds and quickly cooled. After leaching with water, a sizeable yield of clear, well-formed, octahedral crystals of the RbTa_2O_5F pyrochlore phase was obtained.

2.8.2.4 The Systems ASbO₃-Sb₂O₄-AF (A = Na,K,Rb)

The techniques for synthesis and growth of small crystals of $KSbO_{3-x}F_x$ by reaction between KF and Sb_2O_5 at temperatures in the 1000°C range, developed during this contract, have been described [13].

Attempts were made to melt the flux-synthesized $KSbO_{3-x}F_x$ in air with the thought it might be possible to pull crystals directly from

a melt of that particular composition. The material decomposed as the melting point was reached, indicating that pulling in air would not be feasible. A large batch of 50 grams was prepared with a composition of $70\text{KF}:30\text{Sb}_20_4$ and heated in an effort to obtain a melt (single phase liquid) suitable for crystal pulling. At all temperatures below which there was excessive volatilization a solid phase remained. It seems obvious now that we have to go much higher in KF concentration to obtain a composition which may be suitable for crystal growth.

Quenching experiments indicated that mostly ilmenite was formed at temperatures up to $\sim 1200^{\circ}$ C in the system NaF-NaSbO₃. We decided to attempt a direct synthesis of the sodium cubic phase in the same manner as for the KSbO_{3-x}F_x but at a higher temperature. A 3 gram batch of NaF-Sb₂O₄ (molar ratio 93:7) was heated in a platinum crucible to 1250°C and the NaF allowed to evaporate for 35 minutes. After leaching the residue in water we obtained essentially single phase NaSbO_{3-x}F_x.

The largest crystals were of the order of 1 mm. Some flat hexagonal plates of ilmenite are formed at the lip of the crucible, probably during volatilization of the sodium antimony fluor-oxide, however these can easily be separated mechanically from the cubic phase.

X-ray diffraction patterns (single crystal and powder) of selected washed crystals show only a truly cubic body centered phase (a=9.334A). It must be postulated that the composition formed by this techique is slightly different from that made essentially single phase at 4NaSbO₃:NaF in a sealed tube.

Exploratory flux evaporation heatings were made in the $KSb0_3-Sb_20_4-KF$ and $RbSb0_3-Sb_20_4-RbF$ systems as a preliminary step in the determination of suitable starting compositions for flux crystal growth of the desired $ASb0_{3-x}F_x$ phases. The results are summarized

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in Table 11. None of the reported compositions would appear to be suitable for top-seeded solution growth both from the standpoint of the excessive volatility of the alkali fluoride at the temperatures involved and the presence of a solid phase. The presence of a solid phase at such high concentrations of flux (alkali fluoride) indicates that either a gradient transfer technique in sealed crucibles must be adopted or the suitability of other flux systems investigated.

An attempt was made to synthesize "RbSb0₃" in the cubic polymorph by direct flux synthesis from the composition $95RbF:55b_20_4$ heated for one hour at about $900-1000^{\circ}C$ in an open Pt crucible. Well developed psuedo-octahedral crystals were formed and easily isolated although their exact composition is unknown. However single crystal x-ray diffraction patterns indicated that these crystals were probably triclinic, although the observed unit cell was C-centered.

2.9 Ion Exchange

One of the the best screening tests for ionic conductivity of a solid phase is to determine whether or not the alkali ion in the structure can be exchanged with an alkali ion of a different species. This may be tested by heating in a large excess of a molten salt (or solution) containing the second ion. A large number of experiments of this type were performed on many different compounds found in this and the previous year's [1] study. The results of these tests are found in Table 10(a).

Most of the exchange experiments have been performed on powdered materials. In those cases where single crystals could be grown either by flux techniques or melt techniques (see Section 2.8), attempts were also made to ion exchange the single crystals.

In general, the results of ion exchange experiments on single crystals were disappointing in that either disruption of the single crystal

occurred or exchange proceeded at extremely low rates. Thus it appears that a study of technique development for ion exchange of single crystals would be appropriate if these are to be useful in device applications. Consideration should be given to the use of an electric field or other technique in order to obtain a greater driving force for exchange.

Ion exchange experiments were conducted in the $Rb_2O-Nb_2O_5$ system with particular emphasis on the 11-layer compound occurring at or about $4Rb_2O:11Nb_2O_5$ (26.67 mole % Rb_2O). Yon exchange experiments were conducted on both single crystal fragments obtained from crystals grown from the melt and low temperature calcines of 4:11 powders. The single crystals underwent K⁺ exchange in molten KNO_3 but disintegrated during Na^+ exchange in NaNO₃ during various temperature and time combinations. Pellets of the 11-L phase were pressed both uniaxially and isostatically with and without polyvinyl alcohol binder. Pellets were fired and x-ray patterns made before any attempt was made to induce ion exchange. Generally the K⁺ exchange occurred without too much degradation of the specimen but the pellets disintegrated to a fine powder during Na^+ exchange. Results of these exchange experiments are also presented in Table 10.(b). The unit cell dimensions for complete ion exchange were developed from ion exchange in molten salts from very small single crystal fragments.

3.0 <u>RELATION OF STRUCTURAL MECHANISMS OF NON-STOICHIOMETRY TO</u> IONIC CONDUCTIVITY

It is probably generally accepted that a phase which exhibits unusual ionic conductivity must necessarily be structurally nonstoichiometric. Unfortunately the opposite is not necessarily true. Nevertheless a crystallographic understanding of non-stoichiometric phases is an obvious necessity to the tailoring of new fast-ion conductors. For this reason it is worthwhile to discuss the nature of the non-stoichiometry which has been observed in this study for those phases which seem to be of interest.

3.1 Hexagonal Tungsten Bronze-type Phases (HTB)

It has already been mentioned in the previous summary report [1] that the hexagonal tungsten bronze-type phase (HTB) found in binary alkali niobate and tantalate systems has alkali ions in non-stoichiometric positions and excess Nb⁺⁵ or Ta⁺⁵ ions. These excess pentavalent ions may well block the alkali ion conductivity as these hexagonal bronzes cannot be ion exchanged. There are two mechanisms that have proved effective in altering the ion exchange characteristics of the hexagonal tungsten bronze-tupe structures. One is to change the total alkali:other cation valence ratio by substituting W^{+6} for Ta⁺⁵, for instance in the system KTa03-W03. In this case the HTB phase occurs at about the 1:2 ratio and the K^+ ion can be replaced with Na⁺ by heating in a large excess of NaNO, . Unfortunately the Na-tantalumtungstate is not stable above about 450°C and therefore cannot be formed into low porosity ceramics by conventional techniques. The single crystals of HTB grown with the potassium tungstate flux (Section 2.8.2.1) need some greater driving force then temperature (<450°) to obtain complete exchange with Na⁺ ions. The alternate to replacing Ta^{+5} ions with W⁺⁶ ions is to change the structure enough to allow ion exchange (See Section 3.3).

3.2 Pyrochlore Phases

In the $\text{KTaO}_3-\text{WO}_3$ system a pyrochlore phase also occurs at about the 1:1 ratio or $\text{K}_{1.0}[\text{TaW}]\text{O}_6$. Unfortunately the pyrochlore in this system transforms to a tetragonal tungsten bronze (TTB) at high temperatures. Although it can be ion exchanged with Na⁺, this pyrochlore phase also is not stable above about 450°C. The only stable Na⁺ containing pyrochlore is the one in the Sb₂O₄-NaSbO₄ system and apparently this one is not a good ionic conductor.

The distribution of Na⁺, Sb⁺³, Sb⁺⁵ and O⁻² ions in a pyrochlore single crystal is currently under evaluation by the Crystallography Section at NBS. However, certain assumptions can be made which may enable us to postulate the approximate distribution. The formula for the compositions observed to result in a pyrochlore structure might be postulated to be $[NaSb^{+3}]Sb_2O_7$ for the Na/Sb ratio of 1:3, $[Na_{1\ 1/3}Sb_{2/3}^{+3}]Sb_2O_{6.67}$ for 1:2, and $[Na_{1\ 1/2}Sb_{1/2}]Sb_2O_{6.5}$ for 3:5. However, these compositions do not illustrate the structural nature of pyrochlore nor account for the observation that the "lone pair" electrons associated with Sb⁺³ will not allow O⁻² ions to completely coordinate the antimony and result in apparent vacancies.

The structural formula of pyrochlore should be written as $[A_2X]$ $[B_2X_6]$ to emphasize the fact that the octahedral network of B_2X_6 is required to be complete if the structure is to be stable. The A_2X ions fill the intersecting channels in this B_2X_6 framework. In our material the B_2X_6 framework must be represented as $[Sb_2^{+5}O_6]^{-2}$ and <u>must</u> be stoichiometric. All remaining Na⁺ and O⁻² ions, as well as Sb⁺³, must be in the $[A_2X]^{+2}$ portion of the formula. All Sb⁺⁵ must be in B_2X_6 and only Sb⁺³ in A_2X . Furthermore the <u>maximum</u> number of the sum of Na⁺¹, Sb⁺³ excess O⁻² (beyond O₆⁻²) and "lone pair" electrons cannot exceed three. One can then write the general formula as $[A_2O]^{+2}[Sb_2O_6]^{-2}$ with $[A_2O]^{+2}$ equal to

 $Na_{2/k}^{\pm 1} + Na_{x}^{\pm 1} + Sb_{kx}^{\pm 3} + O_{y}^{\pm 2} + L.P_{kx} \le 3$

where k equals the ratio Sb/Na. Using the ionic valences and the sum of the ions equal to three, maximum densities can be calculated and compared with the observed to test the structural hypothesis. The maximum density for the Na/Sb ratio of 1:3 represented by the formula $[Na_{0.917}^{+1}Sb_{0.75}^{+3}o_{0.583}^{-2}]_{0.75}^{+2}[Sb_2^{+5}o_6]^{-2}$ is calculated to be 5.469 units.

For the Na/Sb ratio of 3:5 with the formula

 $[Na_{1.5}^{+1}Sb_{0.5}0_{0.5}^{-2}\Box_{0.5}]^{+2} [Sb_{2}0_{6}]^{-2}$ the density is calculated as 5.406. For the intermediate composition with the Na/Sb ratio of 1:2 and a formula of $[Na_{1.294}^{+1}Sb_{0.588}^{+3}0_{0.529}^{-2}\Box_{0.588}]^{+2} [Sb_{2}0_{6}]^{-2}$ the maximum density is found to be 5.481. The density found for our isostatically hot pressed specimens is 96.0% of the maximum theoretical density. It should be remembered however that the true theoretical density of any given Sb/Na ratio will decrease with decrease in temperature. Thus the densities obtained on our hot pressed specimens are, in all probability, greater than 96% of theoretical in view of the expected increased oxidation of the Sb at the relatively low temperatures involved.

As stated above, the pyrochlore structural formula should be written as $[A_2X]$ $[B_2X_6]$. The RbTa₂O₅ is apparently equivalent to the formula

[[2^{2^{Rb}]⁺¹ [Ta₂⁰5^F]⁻¹}

as shown by Hong's structural study [16] of single crystals prepared by the flux synthesis method described by NBS [17]. The Rb^+ ion is apparently too large for the A sites and it is this preference for the larger anion site that makes this compound stable. During ion exchange in KNO₃ the K^+ ion apparently enters the A site and, upon exposure to atmospheric moisture, an H₂O molecule occupies the site formerly containing Rb⁺. The formula is then

 $[\Box K H_{2}0]^{+1} [Ta_{2}0_{5}F]^{-1}$

During sodium exchange it is apparently possible to obtain a non-stoichiometric amount of Na in this lattice. The formula for this phase can probably be written as

 $[Na_{1+x}OH_x]^{+1} [Ta_2O_5F]^{-1}$.

The product obtained by acid leaching of this pyrochlore is apparently

 $[\Box_{2^{H_{3}0}}]^{+1} [T_{a_{2}0_{5}F}]^{-1}.$

Although infra-red analysis of the Na⁺ exchange product does not indicate (OH)⁻ [18] it would probably be worthwhile to examine this product with NMR for hydrogen resonance, a much more sensitive method than infrared adsorption.

3.3 Hexagonal Tungsten Bronze - Pyrochlore Series

The $[B_2X_6]$ framework of the pyrochlore structure can be described as being made up of alternating layers of the hexagonal tungsten bronze structure separated by layers of isolated octahedra sharing only two corners with the adjacent HTB layers. If this structure is modified by increasing the sequence number of either of these types of layers from ABAB to AABAAB or AAABAAAB etc., a sequence of hexagonal phases would be formed having the same a axis as the HTB structure and with varying but integral multiplicities of the c axis dimension. Such phases are actually encountered in the K20-Ta205 [1,2] Rb20-Nb205 and Rb₂O-Ta₂O₅ systems and have been estimated by us to represent 9-layer, 16-layer and 11-layer sequences. All of the phases can be ion exchanged for Na⁺ unlike the HTB in the same systems. The reason for this appears to be that a rotation or translation of a portion of the layer sequence allows the isolated vertical channels found in the HTB structure to be changed to intersecting channels as in the cubic pyrochlore structure. The exact crystallographic nature of these compounds is currently under investigation by Dr. B. T. Gatehouse at Monash University, Melbourne, Australia. Unfortunately, however, the Na⁺ ion exchanged products are again not stable above about 450°C.

As is the case with the pyrochlore compounds the K_2^0 containing phases have unit cell dimensions very similar to or even larger than the corresponding Rb_0^0 containing phases. For instance the c-axis for

the 11-layer phase in the $K_2O-Ta_2O_5$ system is 43.512A while that in the $Rb_2O-Nb_2O_5$ system is 43.18A and in the $Rb_2O-Ta_2O_5$ 43.19A. This may be due to a real difference in total alkali content. It may also be due to hydration of the K_2O phases. No high temperature x-ray data is available to check this hypothesis but small single crystals of the 2:5 $K_2O:Ta_2O_5$ phase have been noted to crack, spall and jump on exposure to air.

3.4 Body Centered Cubic Antimonates

A successful method of synthesizing cubic potassium antimonate by heating in molten KF was published by the present authors during this contractural period [13]. The major reason for the success in obtaining completely single phase fluorine stabilized cubic potassium antimonate is that the potassium ilmenite is H_2O soluble and may be easily separated from the cubic material.

An examination of the structural model of the octahedral framework of the body centered cubic antimonate phase suggests that this structure <u>must</u> always have some anion (X) occupancy in the 000, $1/2 \ 1/2 \ 1/2$ position. The structural formula thus appears to be $[A_{16}X_2]^{+12}[Sb_{12}o_{36}]^{-12}$ with the alkali ion in position (A) located at (or just off) the juncture of the open cages. However, it seems very likely from both structural reasons (bond lengths, etc.) and valency considerations that either or both of the non-framework positions will be nonstoichiometric. Valency considerations require that at least two out of 16 alkali ions must be missing and the structural formula then be $[\Box_2A_{14}X_2]^{+12}[Sb_{12}O_{36}]^{-12}$. This formula corresponds to the composition reported by the Lincoln Laboratory report [19] for the single crystal x-ray diffraction analyses of the phase synthesized with KF according to the NBS method [13]:

 $K_{12}Sb_{12}O_{36}.2KF$ or $[\Box_{2}K_{14}F_{2}]^{+12}[Sb_{12}O_{36}]^{-12}$.

It seems quite likely, however, that this general formula does not completely account for all of the preparations which have been observed to form this structure, whether body centered or primitive. The observation that a primitive phase can be formed, in air, by reaction with atmospheric moisture at a 48:52 ratio suggests that this phase may well have considerably less than 14 alkali ions per unit cell. The formula <u>must</u> be compensated, in this case, by a substitution of a monovalent anion (OH, F) in the <u>octahedral framework</u>. The general formula then becomes $[\Box_{2+x}A_{14-x}X_2]^{+(12-x)}[Sb_{12}O_{36-x}X_x]^{-(12-x)}$. The composition found at $\vee 48:52$ in the potassium antimonate system can be written (assuming a ratio of 11:12 K/Sb or $47.826\% \text{ K}_2\text{O}$): $\text{K}_{22}\text{Sb}_{24}O_{71}^{+5H}2^{O} \neq \text{K}_{22}\text{Sb}_{24}O_{66}(\text{OH})_{10}$ or $[\Box_5\text{K}_{11}(\text{OH})_2]^{+9}[\text{Sb}_{12}O_{33}(\text{OH})_3]^{-9}$ which also can be described as $6\text{KSbO}_3:3\text{Sb}_2O_5:5\text{KOH}$ (see Figure 8). The general formula describing the K⁺ containing compositions is then

 $[\Box_{2+x}K_{14-x}X_{2}']^{+(12-x)}[sb_{12}^{+5}o_{36-x}X_{x}'].$

The above formula contains only pentavalent antimony and apparently does not completely explain the compositions which form a 'stable' body centered cubic phase in the system $NaSbO_3:Sb_2O_{4+x}:NaF$. The only formula which does not involve the loss or gain of O^{-2} (or F⁻) when the Sb_2O_4 is added in a sealed tube corresponds to:

$$\left[\Box_{2^{Na}14}F_{2}\right]\left[sb_{y}^{+3}sb_{12-y}^{+5}O_{36-2y}F_{2y}\right]$$

which is represented by the join 6:1 - 3:7 on Figures 5 and 6. There is really no place in the framework structure for Sb^{+3} and it is difficult to believe that octahedrally coordinated antimony can be Sb^{+3} . The lone pair electrons can be attached to the sodium ions instead of the antimony or just in the vacancies. However, for convenience, the

formulas can be written involving Sb⁺³. The new formula would then have two variables:

$$[\bigsqcup_{2+x^{A_{14}-x^{X_{2}'}}} [sb_{y}^{+3}sb_{12-y}^{+5}o_{36-(x+2y)} x'_{(x+2y)}]$$

represented by the plane in the quaternary system $NaSbO_3:Sb_2O_3:NaF$ bounded by the 6:1 -- 3:4 and 6:1 -- 3:7 joins of Figures 5 and 6. However the single phase region in this system actually appears to contain more NaF than described by this general formula. Apparently some O^{-2} is evolved in the sealed Pt tubes, the amount depending on uncontrolled variables such as the amount of free volume in the tube and on changes from the original composition during treatment. The absolute maximum amount of NaF which can be accommodated structurally by the body centered cubic phase can be described by the formula

$$[Na_{16}F_2]^{+14}[Sb^{+3}_{2-x}Sb_{10-x}O_{34-2x}F_{2+2x}]^{-14}$$

which represents a line in the system shown by the join 3:1 -- 3:8 in Figure 6 and involves the evolution of one molecule of gas (O_2) per formula unit. The results of our investigations so far suggest that the body centered phase approaches this formula as a limit. The composition of the cubic phase in equilibrium with excess Sb_2O_4 and molten NaF actually appears to touch this line at approximately $10NaSbO_3:Sb_2O_4:6NaF$ or

 $[Na_{16}F_2]$ $[Sb_3^{+3}Sb_9^{+5}O_{32}F_4] + O_2.$

The single phase distorted cubic material on the binary join NaSbO₃:NaF appears to have a composition between 6:1 and 5:1 or approximately 11NaSbO₃:2NaF or

 $[Na_{14.18}F_2]$ $[Sb_{12}O_{35.818}F_{0.1818}] + .0909 O_2.$

The compositions in the quaternary system thus probably lie on a join between these two end members.

A large number of experiments were performed in an attempt to synthesize ceramic products of single phase body centered cubic sodium antimonate "stabilized" with F^+ . Although we have not yet succeeded in preparing such a ceramic all indications point to a short time heat treatment with specific specimen preparation techniques. These almost unique conditions indicate that the body centered cubic phase is not really stable under the conditions at which it is formed. Thus either its true stability field may lie at some higher or lower partial pressure of O_n (or OH) or it may always be metastable under any conditions.

3.5 The Phase 2K₂0:3Nb₂0₅

In the previous contract summary report [1] it was reported that the compound occurring at about 2K20:3Nb205 had the unit cell dimensions a=7,822, b=33,019, c=6.481A when not hydrated. Only the b axis expanded upon exposure to moisture as previously mentioned by Nassau [15]. From these unit cell dimensions several postulations can be made. The a direction corresponds to 2x3.911 or two O-Nb-O distances and probably represents two Nb-octahedra sharing oxygen at their corners. The c direction corresponds to 2x3.2405 or two times the 0-0 distance corresponding to edge sharing of the Nb-octahedra. The b direction is \sim 10 times the value for edge sharing. It may be assumed that the structure is made up of layers or slabs perpendicular to b, composed of octahedra edge-shared in the c direction and corner shared in the a direction. These slabs must be separated by layers of K⁺ ions and the K^+ ions should then be rather mobile. That this is actually the case is demonstrated by our reported ion exchange with Na⁺ where the b axis changes to 30.78A and the conductivity measurements made by LeRC [20]. The actual crystal structure of this phase is currently under investigation by Dr. N.C. Stephenson of the New South Wales Institute of Technology, Sydney, Australia.

Although no ionic-conductivity or even ion exchange studies have been performed on the compound $K_20:3Nb_20_5$ it is sufficiently similar to the 2:3 phase to warrant some additional study.

4.0 SUMMARY OF RESULTS

- Single crystals of hexagonal tungsten bronze (HTB) were synthesized in the system KTaO₃-WO₃-K₂WO₄ by the flux technique. No crystals large enough to measure conductivity by the presently utilized techniques were successfully obtained.
- 2. The phase equilibria up to and including the liquidus of portions of the systems Rb₂O-Nb₂O₅ and Rb₂O-Ta₂O₅ have been studied and phase diagrams have been constructed most consistent with the experimental data. Single crystals were grown of most of the compounds in the niobate system and the phases were investigated for ion exchange properties. Single crystal and polycrystalline specimens were submitted to the sponsor for evaluation.
- 3. The phase equilibria up to and including the liquidus of the systems Sb_2O_4 -NaSbO₃ and Sb_2O_4 -KSbO₃ have been studied and a phase diagram has been constructed most consistent with the experimental data. Polycrystalline ceramic specimens of the pyrochlore phase were prepared and submitted to the sponsor for evaluation.
- 4. The stabilization of cubic antimonates by addition of F^+ was studied in an attempt to provide a low porosity Na⁺ ion ceramic specimen for evaluation. Although small single crystals were easily synthesized by the alkali fluoride flux technique, no crystals larger than \sim 1-2 mm were obtained.
- 5. Many other alkali-oxide rare earth oxide and Bi₂O₃ systems were examined but no new interesting phases identified.
- 6. The phase occurring at about $2K_20:3Nb_20_5$ was found to have interesting ion exchange and ionic conductivity properties.

5.0 FUTURE WORK

- Develop techniques to grow large crystals from a flux. Such techniques have only been reported for a very few phases and generally involve an inordinate amount of man hours to produce results.
- 2. Develop techniques to ion exchange large crystals, perhaps by utilizing an electronic current with a hot alkali salt or solution.
- 3. Investigate and develop improved fabrication techniques to produce dense, sound pellets of the 68NaSbO₃:4Sb₂O₄:28NaF composition in a size suitable for definitive ionic conductivity measurements and examine the effect of varying composition on conductivity.
- Obtain crystallographic structure analyses on 2K₂O:3Nb₂O₅ and on 4Rb₂O:11Nb₂O₅ (now being conducted by N.C. Stephenson and B. Gatehouse).

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Figure Captions

Figure 1. Phase equilibrium diagram for the system
$$Nb_20_5-4Rb_20:11Nb_20_5$$
.
• - liquidus values from reference [21]
• - completely melted
• - partially melted
• - no melting
Figure 2. Phase equilibrium diagram for the system $Ta_20_5-4Rg_20:11Ta_20_5$.
• - partially melted
• - no melting
Figure 3. Phase equilibrium diagram for the system $Sb_20_4-NaSb0_3$.
Not necessarily a true binary system. (L = liquid, 3
S = solid, V = vapor)
• - melting
X = no melting
Figure 4. Phase equilibrium diagram for the system $Sb_20_4-KSb0_4$.
Not necessarily a true binary system.
• - melting
x = no melting
ss = solid solution
1:2 - K,0:28b_02
3:5 - 3K_0:58b_20
Figure 5. Phase relations in the quaternary system NaSb0_3-Sb_20_3-Sb_20_5-NaF.
The join 6:1--3:4 represents the formula
 $[\Box_{2+x}Na_{14-x}F_2]$ $[Sb_{12}^{+5}o_{36-x}F_x]$
The join 6:1--3:7 represents the formula
 $[\Box_{2+x}Na_{14-x}F_2]$ $[Sb_{12}^{+5}o_{36-x}F_x]$
Figure 6. Phase relations in the ternary system NaSb0_3-Sb_20_4-NaF
The join 6:1--3:4 represents the formula
 $[\Box_{2+x}Na_{14-x}F_2]$ $[Sb_{12}^{+5}o_{36-x}F_x]$

The join 6:1--3:', represents the formula

$$\begin{bmatrix} \Box_{2}Na_{14}F_{2} \end{bmatrix} \begin{bmatrix} sb_{y}^{+3}sb_{12}^{+5}o_{36-(x+2y)}F_{(x+2y)} \end{bmatrix}$$

The join 3:1 --3:8 represents the formula

$$[Na_{16}F_2]$$
 $[Sb_{2+x}^{+3}Sb_{10-x}^{+3}O_{34-2x}F_{2+2x}] + O_2$

Figure 7. Phase equilibrium diagram for the system Nb₂O₅-KNbO₃

- x liquidus values from reference [22]
- o completely melted
- e partially melted
- - no melting
- Figure 8. Representation of the ternary system KSb0₃-Sb₂0₅-KOH illustrating the hydration of the 11:12 mixture to form the single phase cubic compound when heated in air.

















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| 7/ | ible 1, Xeperi | imental Da | ta for th | a Aystem) | Nag0g-28820121NB205 | |
|---------------------------------------|-------------------------------------|----------------------------------|--|---|--|---|
| Composition | Provid Trad Tamp, *C | nus Heat Liment Time hr | Final Tree Temp. "C | Nest iment Tine br | Results of Physical Observation | Results of X-ray Diffraction Analysis |
| 38820183H82(| 500 500 600 | 120 170 90 | 300 600 | 120b/ 90b/ | | H-H9205 + 213 H-H9205 + 3117 |
| | | | 800 900 1000 1000 | 66 40 80 111£/ | | H+Hb205 + 3117 H-Hb205 + 3117 H-Hb205 + 3117 |
| | _ | | 1200 | 88 · | | H-H0203 + CTB |
| TORP ³ 01404P ³ | 03 500 500 600 | 120 120 90 | \$00 \$00 | 902/ * | | H-HD ₂ O ₅ + 213 H-HD ₂ O ₅ + 213 |
| | | | \$00 \$00 1000 1075 1200 | 60 40 80 1115/ 60 88 | *. | 3:17 + M-Hb ₂ O ₅ 3:17 + H-Hb ₂ O ₅ 3:17 + CTB + H-Hb ₂ O ₅ CTB + 3:17 + N-Hb ₂ O ₅ CTB + 3:17 + N-Hb ₂ O ₅ CTB + H-Hb ₂ O ₅ |
| 11.576 ₂ 0;66, | 500 500 500 | 120 120 90 | 500 600 | 1200 | • | H-ND205 + 213 H-ND205 + 213 |
| | | | 800 900 | 60 40 | | 3:17 + H-WB.05 3:17 + H-WB.05 |
| | 500 600 | 120 90 | 1000 1000 | 80 111£/ | | GTB + 3:17 ² GTB + 3:17 |
| | 1000 | *** | 1145 1200 1280 | 16 85 40 | | GTB GTB GTB |
| | 4 | | 1340 1349 | 23 | Not melted | GTB . |
| | e. E | | 1352 | ,25 ,33 | Not melted Not melted | |
| ه ۲۰ ۰ د در | • | | 1369 1375 1380 | ,5 1,5 2 | Not melted Fartial melted Completed melted | GTS |
| 158620185862 | 0 ₅ 500 500 | 120 120 | 500 600 | 120k/ 90k/ | , | H-Hb205 + 213 H-Hb205 + 213 |
| | | • | 750 730 800 900 1000 | 165/ 165/ 44 40 80 1119/ | • | 11-L + K-Kb ₂ 0 ₃ 3117 3117 3117 3117 3117 3117 3117 |
| | 500 600 | 120 90 | 1100 | 44 44 | | 3117 GTB + NTB |
| , , | - | | 1156 1177 1180 1183 1226 1262 | 36,5 24 23 64 18 18 | | 5:17 3:17 + GTB 3:17 + GTB HTB + GTB HTB + GTB |
| | 500 600 1000 1200 | 120 90 111 16,5 | | | · · · | RIB - CIB |
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| 208620180062 | 5 500 500 | 120 120 | 500 600 | 120 <u>b/</u> 90 ^{b/} | , · | H-Hb203 + 215 H-Hb203 + 213 |
| • | | yu | 800 900 1000 1000 | 66 40 80 1115/ | | 3:17 + 11-L 3:17 + 12-L 3:17 + 11-L |
| · · · · · · · · · · · · · · · · · · · | | | 1200 | 40 57 | | 9L + GTB + HTB |
| 21,7586 ₂ 0;78, | ,2586 ₂ 03 500 600 | 120 120 90 | 500 600 | 120 ²⁷ 90 ⁸ / | | |
| | | | 800 900 | 65 40 | | 3:17 + 11-L |
| | | | 1000 | 111 <u>e</u> / | | 3127 + 9-L + 11-L |
| | | | 1200 | 88 16 | | 371/ 4 9-6 9-6 + 678 878 + 9-1 |
| DAC | E IS | | 1250 | 16 | | HTB + 9-L HTB + 9-L |
| DRIGINAL FAR | TTTY | | 1300 | 16,5 | | HTB + 11-L + 16-L |
| OF POOR QUA | | | | 50 | :) | |

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| | | 500 600 1000 | 120 90 111 | | | | |
|---------------|---|--------------------|--------------------|--------------------------------------|-------------------------------------|---|--|
| | | | | 1149 1169 1181 1184 1313 | 23 18 89 3,5 64 | N-s -stead | 9-L + 3:17 9-L + 3:17 9-L + GTB HTB + 9-L HTB + 11-L |
| | | | | 1320 1320 1328 1347 | 0.5 22 1.5 16.5 | Fartially malted Completely malted Completely malted Completely malted | HTS + 11-L HTB + 11-L |
| 23. | 191620176,81N52 | 1000 | 1.263 | | •1 | | 9-1, + HTB |
| | | | | 1200 1255 1290 1306 | 215) 18 18,5 22 | | 9-1 + HTB 9-1 + HTB 9-1 + HTB 9-1 + HTB HTB + 11-1 |
| 258 | ^b 2 ^{0:75Nb} 2 ⁰ 5 | 500 500 600 | 120 - 120 90 | 500 600 | 120 <u>b</u> / 90 ^b / | | $\frac{Nb_2O_3}{Nb_2O_3} + 213$ $\frac{Nb_2O_3}{2} + 213$ |
| | | | | 800 900 1000 1000 | 66 40 80 1115/ | | 11-L + 3;17 11-L + 9-L |
| | | | | 1200 1260 1300 | 88 1 40 | | 9-L 9-L + HTB 9-1. |
| | | 500 600 | 120 90 | | | | |
| | | 1000 | 111 | 1290 1300 | 18.5 | | 11-L + HTB |
| 25. | 58520174.585205 | 500 500 | 63 63 | 500 600 | 63 ^{b)} | | |
| | | 800 | 24 | 1000 1249 | 69 ^C / 22 | | 9-L + 11-L 9-L |
| | | | | 1258 | 67 16 17 | | 9-1 + 16-1 9-1 + 16-1 9-1 + 16-1 |
| | | | | 1275 | 117 | | 16-L + 1tr 16-L + 1tr |
| | | | | 1285 1287 | 18 67 | | 16-L + 11-L 16-L + 11-L |
| • | | | | 1291 1294 1297 | 21 16 22 | | 16-L + 11-L 16-L + 11-L 16-L + 11-L |
| 26. | 67Rb20173.33Mb20 | 5500 500 | 120 120 | 500 600 | 120 <u>b/</u> 90 <u>b</u> / | | H-Nb205 + 213 H-Nb205 + 213 |
| | | | | 750 800 | 60 66 | | 11-L 11-L |
| | | | | 900 1000 | 40 80 _c / | • | 11-L 11-L |
| | | | | 1000 | 60 88 | | 11-L 11-L |
| | | | | 1241 1260 | 18 | | 11-L 11-L |
| TAL PAU | LTTY. | 500 | 120 | 1300 | 40 | | H-L |
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| OK FO | | | | 1306 1316 1316 1320 | 22 0.5 16 0.5 | Not melted Partially maited Partially melted Completely melted | 11-L |

"Except as indicated by a footnote the samples were quenched into water from the temperature indicated.

²⁷ Except as indicated by a footnote the samples were quenches into water from the traperous indicated by a footnote the samples were quenches into water from the furnaces at the temperatures indicated. Rb₂O may have been volatilized during these calcines thereby changing the composition to one greater in mole percent Nb₂O₅.
²⁷ The sample was a large sealed Pt tube and pulled from the furnace at the temperature and cooled rapidly on a chill block.
⁴⁷ The sample was removed from the furnace at the temperature indicated and immediately dunked into a beaker of water.

of vater.

| 18048 4 | · ckyet | Tealizer | hare tot | tua skaram | 1205-4802011112205 | |
|--------------------|----------------------------|----------------------|--|--|--|--|
| Composition | Initial Treatm Temp, | lieat ant Time | Final Tree Temp. | Hest ^A / tment Time | Results of Physical Observation | Results of X-ray Diffraction Analysis |
| 5852019574205 | | 116 | 500 | 120 <u>b</u> / | | L-Te205 |
| | 500 500 600 | 120 120 110 | 600 | 1102 | | |
| | | | 1340 1500 | 120 144 | | L-Ta ₂ O ₅ + GTB + H-Ta ₂ O ₅ + GTB |
| 108520:9074205 | 500 500 600 | 120 120 90 | 500 600 | 120b/ 110b/ | | L-Ta205 |
| | | | 800 900 1365 1500 1700 | 80 60 120 120 1 , | | L-Ta ₂ O ₅ + 9L L-Ta ₂ O ₅ + 9L GTB # L-Ta ₂ O ₅ GTB + H-Ta ₂ O ₅ (tr) GTB |
| 11.5Rb20:88.5T4205 | 500 500 600 | 120 120 110 | 500 600 | 120 <u>2/</u> 110 ^{2/} | | |
| | | | 1098 1255 1318 1340 1500 1603 | 136 27 16 120 144 24 | | L-Ta ₂ O ₅ + 9L. 9L + ² L ² Ta ₂ O ₅ 9L + L-Ta ₂ O ₅ GTB GTB GTB |
| | 1200 | 150 | 1296 1297 1307 1317 | 91 117 64 67 | | L-Ta,0, + 9L + GTB (tr) GTB + L-Ta,0, GTB + L-Ya,0, GTB + L-Ya,0, |
| • • • • | • | | 1327 1330 1332 1340 | 18 115 16 72 | • | $\begin{array}{l} GTB + L - Ta_{2}O_{5}^{2} \\ GTB + L - Ta_{2}O_{5}^{2} \\ GTB + L - Ta_{2}O_{5}^{2} \\ GTB \end{array}$ |
| | | | 1704 1709 1714 | 0.5 | not melted not melted not melted | · · |
| 11 - A | 1110 | 118 | 1721 1722 | 2 | completely melted | CTB |
| | 1340 | 72 | 1317 | 67 | • | GTB · |
| | 1277 | 136 | 1277 1297 | 136 117 | • | GTB |
| | | | 1331 | 115 | | GTB |
| 158520185T#205 | 500 500 600 | 120 120 110 | 600 | 1105 | | · |
| • | • | | 800 900 1008 1360 1473 1500 1500 1500 1500 | 80 60 136 120 65 17.5 18 144 2 | | L-Ta ₂ O ₅ + 9L L-Ta ₂ O ₅ + 9L L-Ta ₂ O ₅ + 9L L-Ta ₂ O ₅ + 9L GTB + 9L GTB + 9L GTB + HTB GTB + HTB GTB + HTB GTB + HTB |
| | 1200 | 150 | 1707 1711 1714 1715 | 0.5 0.5 0.5 0.3 | not melted not melted completely melted completely melted | |
| 208520:8074205 | 500 500 | 120 120 | 500 600 | 120 <u>b</u> / 110 <u>b</u> / | ini di s | |
| • | 600 - | 110 | 1340 | 120 | | 9L + GTB |

erimental Data for the System Ta.O Table -486 0+117

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| 21.758620178.2576205 500 | 120 | 300 600 | 1202/ | | |
|---|-------------------|--|--|--|--|
| 400 | 110 | 800 900 1100 1360 1473 1500 1643 1700 1700 | 80 40 138 120 63 18 24 2 1 1 | | 9L + L-Ta ₂ O ₃ 9L + L-Ta ₂ O ₅ 9L + L-Ta ₂ O ₅ 9L + CTB ₂ O ₅ 9L + CTB 9L + CTB 9L + CTB HTB + 9L (tr) HTB HTB |
| 1200 | 130 | 1474 1477 1477 1479 1480 1482 1484 1484 1486 1488 1488 1498 1498 1706 1716 1717 1721 | 67 23 18 17 19 17 18 45 20 41 16 16 21 1 1 0.5 2 | not melted not melted partially melted partially melted | 9L + GTB 9L + HTB 9L + HTB 9L + HTB HTB |
| 2586 ₂ 0:7578 ₂ 05 500 500 600 | 120 120 110 | 500 600 | 120 b/ 110 b/ | | L-Ta ₂ 05 |
| | | 1000 1100 1344 1498 1675 1700 1700 1715 | 40 138 120 144 18 1 1 1 | not ne lted | 9L + L-Ta ₂ O ₃ (tr) 9L 9L + 7 (tr) 16L + HTB 16L + HTB 16L + HTB |
| 1200 | 150 | 1600 1621 1624 1659 1663 1677 1682 1722 1725 | 20 3 17 21 19 67 29 18 16 16 | partially melted pertially melted | 9L 9L 9L 9L 9L 9L + HTB (cr) 9L + HTB + 16L 16L + HTB 16L + HTB |
| 26.78520:73.318205 500 500 600 | 120 120 110 | 500 600 | 120 <u>b</u> / 110 <u>b</u> / | | L-Ta ₂ 0 ₅ + 213 |
| | 170 | 800 900 1098 1360 1500 1596 1625 1640 1647 1700 1710 | 80 60 40 136 120 120 24 16 2 1 1 1 1 | not melted | 111 + 213 $111 + 213$ $111 + 213$ $111 + 213$ $111 + 213$ $161 + 111$ $161 + 111$ $161 + 111$ $161 + 111$ $161 + 111$ $161 + 111$ $161 + 111$ 111 |
| 1200 | 120 | 1283 1672 1719 1725 1726 | 72 1 0.5 16 | not belted | 16L + 11L 11L 11L 11L |

A Except as indicated by b/ the samples were quenched into water from the temperature indicated.

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 \underline{b}' The samples were heated at the temperatures indicated in open Pt crucibles. Due to the volatility of Rb₂O the bulk composition may have shifted to one higher in mole percent Ta₂O₅.

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| Conpos | itton | Heat Tree | tsent. | H | -1 |
|-----------------|-------------|------------|------------|-------------------------------|---------------------------------------|
| ma ju ma IoX | molei | 1630 *C | hr | Thysical Observation | X-ray Diffraction Analysis |
| 50 | 50 | 1213 | | not welted | · · · · · · · · · · · · · · · · · · · |
| | | 1264 | 3 | 14 14 | N#\$603 |
| | | 1433 | 1 | 98 BÁ | |
| | | 1502 | .05 | 29 BÁ | |
| | | 1542 | .08 | 4 H | NalbO ₃ + unknown |
| | | 1569 | .08 .08 | Helted | • |
| 45. | 13 | 1000 | 44 | ngt melted | |
| | | 1100 | 44 | n n | NaibO, + pyrachlare es |
| | | 1473 | 123 | 11 II | , |
| | | 1495 | .08 | completely melted | |
| 40 | 60 | 3102 | 20 | ngt melted | pyrochiore se + Naub0, |
| | | 1305 | 19 | 01 11 61 14 | n n - , |
| | | 1470 | -08 | ** 15 | |
| | | 1488 | .08 | 14 H | |
| | | 1495 | .03 | completely melted | |
| 37,5 | 62.5 (315) | 1100 | 48 | not welted | Fyrochlore se |
| | | 1306 | 19 | ai 11 | synochlore es |
| | | 1326 | 20 | at 14 | F) |
| | | 1351 | 1 | 99 la | |
| | | 1373 | 2 | not melted (reneat of 1100-4) | pyrochlore as |
| | | 1392 | .16 | | pyrochlore as |
| | | 1412 | .16 | H 11 | |
| | | 1447 | -16 | 11 F1 | |
| | | 1434 | .08 | N 11 | • |
| | | 1464 | .08 | 67 P 3 | |
| | | 1476 | .08 | <u>н</u> н | • |
| | | 1487 | -08 | 11 H | |
| | | 1490 | -08 | , melted | |
| 33.33 | 66.67 (112) | 10000 | 8 | not melted | nyrochiere se |
| | | 1100 | 3 | H 4 | Pyternicite of |
| | | 1103 | 91 | 41 1 4 | 41 |
| | • | 1287 | 2 | H H | 11 |
| | | 1292 | 1.5 | 41 H4 x | •• |
| | | 1307 | 19 | FC 14 | pyrochlore se |
| | | 1316 | .5 | u h , . | ** ** |
| | | 1317 | 3.5 | 11 . 11 11 . 14 | |
| | | 1334 | 24 | .н. 4 | N N |
| | | 1376 | .3 | , н н | 91 |
| | • | 1378 | .5 | 69 ¹⁰ | |
| | • | 1411 | 19 | 4 4 | |
| | | 1437 | 24 | pertially melted | |
| | | 1475 | .02 | completely melted | |
| 25 | 75 (1:3) | 750 | 60 | not melted | pyrochlore ss + unknown // |
| | | 800 | 60 | 47 H | pyrochlore as + unknown |
| | | 800 | 60 | n 11 | pyrochiors ss + unknown |
| | | 1098 | 16 | 0 W | pyrochlore s* |
| | | 1192 | ĩ | 10 H | * <i>#</i> *1 |
| | | 1200 | 24 | 4 H | 14 |
| | - | 1220 | 2 | P 11 | 11 |
| | | 1306 | 24 | te ai | |
| | | 1307 | .08 | 60 H | |
| | | 1317 | 16 | 11 14 | pyrochlore as |
| | · | 1339 | .08 | n () | |
| | | 1345 | 25 | FT 61 | pyrochlore se |
| | | 1346 | .08 | 17 +1 19 -21 | |
| | | 1358 | .08 | sertially matrix | pyrochlore as + M-ShO |
| | | 1427 | .02 | Hairtary Bairag | Naranitore an a second |
| 71 | 77 | 1200 | 24 | not melted | - vchlore ss + B-Bb.O. |
| 43 | ** | 1266 | Â, | | chlore + q,+ 81/2 4 |
| | | 1267 | 19 | 24 14 14 14 | Jochlore sel |
| | | 1299 | .08 | 17 18 - 71 | |
| | | 1304 | .08 | 0 4 | |
| | | 1322 | ,00 | U Q | 1/ |
| | | 1332 | .08 | 97 76 . An Ar | pyrochlors as' |
| | • | 1338 | .08 | W 74 | |

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Table 3. Experimental Data for Compositions in the System Sodium Antimonate-Antimony Tetroxide."

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| 20 | 60 | 1099 | 672 | not malted | pyrochlore ank/ |
|----|----|------|----------------|---------------------------------------|--|
| | | 1107 | 244 | | |
| | | 1200 | 24 ' | not melted | pyrochlore as + a |
| | | 1220 | 2,5 | 17 19 | pyrochiore ss + 0-5b_04 |
| | | 1234 | 2.5 | H H | 1 U U U U U U U U |
| | | 1277 | 16 | 11 H | pyrochlore as $+\beta-5b_0$, |
| | | 1301 | .5 | not malted | |
| | | 1305 | 19 | | NaBbo E/ |
| | | 1106 | 26 | es es | pyrochlore as + B-5b.C. |
| | | 1116 | 1 08 | | pyrochlore as $d = \frac{1}{2}$ |
| | | 1010 | | *1 15 | pyrocitore an 4 d-ob204 |
| | | 1310 | 100 | 0 D | . • |
| | | 1333 | .00 | 6 U | |
| | | 2339 | •4 | | |
| | | 1340 | •08 | , , , , , , , , , , , , , , , , , , , | |
| | | 1345 | 12 | partially malted | |
| | | 1362 | .5 | ** ** | |
| 15 | 65 | 600 | 74 | not melted | unknown + pyrochlors as + og5b.0, \$11/ |
| | | 800 | 60 | , и и | a + pyrochlore se + unknown***** |
| | | 1000 | 64 | | unknown + tr α -Sb ₂ O ₁ (dried 240) $\frac{1}{240}$ |
| | | 1000 | 64 | | unknown + tr α -Sb- 0 , α |
| | | 1007 | 48 | not melted | nyrochlore + 0-5b-0, + anknown |
| | | 1107 | 144 | H H | pyrochlore as $+ = 55.0. + 8-55.0$. |
| | | 1200 | 24 | 0 0 | py |
| | | 1200 | 60 | 11. ji | syrablars as + 0-th 0 |
| | | 1337 | ů, | +1 + | pyrocatore and 0-00204 |
| | | 1337 | • | H 11 | |
| | | 1340 | 15 | | |
| | | 1398 | •4 | percially metted | • • |
| 10 | 90 | 800 | 74 | not melted | $a-5b_0$, $+$ unknown ¹ / |
| | | 1007 | 48 | ····· | α -5b.0, + 6-5b.0, + pyrochlors = $\frac{1}{2}$ |
| | | 1107 | 166 | n (t | |
| | | 1236 | 5 | te și | d-Sh 0 + pyrochlore es |
| | | 1101 | • •• | 11 D | |
| | | 1200 | . 13 | 11 11 | |
| | | 1200 | | H D | a fb a t unus tiens as |
| | | 1111 | .7 | 17 ET | u-so ₂₀₄ + pyrochiore == |
| | | 1310 | 33 | 10 H | |
| | | 1336 | | н и | |
| | | 1121 | | 1) (I | a_Sh 0 + overachlore es |
| | | 133/ | ; * | het is a second second | a + Superhises as + superhad liquid |
| | | 1321 | 1 | baterarry marced | a + pyrochiore ss + queboned iiquid- |
| 5 | 95 | 1007 | 48 | not melted | $\beta - 5b_00_1 + \alpha - 5b_00_1 + pyrochlore ee^{1/2}$ |
| | | 1107 | 144 | 47 44 | α -sb ₂ 0 ⁷ ₄ + pyrothiore as + trace β -sb ₂ 0 ⁴ ₄ |
| | | 1234 | 3,5 | H H. | н 1124 н н |

All specimens were preheated to 750°C for 60 hours and 1200°C for 19 hours unless otherwise footnoted. Rate of heating and cooling was approximately 3°/min. For higher heat treatments, speciments were heated in scaled Pt tubes and quenched from temperatures indicated.

- b/ The phases identified are given in the order of the amount present (greatest amount first) at room temperature. These phases are not necessarily those present at the temperature to which the specimen was heated.
- C/ Specimen heated with PtO, at 68,900 psi in scaled Pt tube.
- $\frac{df}{df}$ Specimen heated in scaled Pt tube at 5,000 psi.
- e Specimen previously heated at 1292*C for 1.5 hours.
- ${\cal U}$ Specimen heated in scaled Pt tube in presence of water. The unknown phase formed is probably a hydrate.
- E' Specimen heated in scaled Pt tube in PtO2.
- $\frac{h}{2}$ Specimen heated in presence of 5:95 Na₂O:Sb₂O₄ which served as a buffer.
- 1/ In spite of extensive x-ray study it has not been determined which of the polymorphic forms of Sb204 is the stable form.
- \mathcal{U} Sb₂Q₄ probably mosked into Pt container and the composition changed to pyrochlore set
- k/ Platinum tube leaked.
- 1/ Unknown phase, d-spacing of major lines given in text. This phase is probably a hydrated phase which exists in the presence of moisture and/or PtO, and can be aliminated by an solitional calcining of 1200°C for several hours. Once eliminated this phase does not appear to reform at lower temperatures in laboratory time.
- ^{m/} Specimen contained non-equilibrium material derived from a liquid when quenched from above the liquidus and examined at room temperature.

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| Composition | Heat 7 | freatment | | Results | к.] |
|--|-------------------|------------------|------------------------------------|---|---|
| Starting Material | Temp °C | Time hr | Environment | Physical Observation | X-ray Diffraction Analysis |
| α-\$b _{i12} 04 | 1223 " | .5 | sealed Pt tube unsealed Pt tube | not melted | α + tr β α |
| ^{β-5} ^b ,2 ⁰ 4 | 1223 | ,5 " | scaled Pt tube unscaled Pt tube | not melted volatilized | β + tr α |
| β-sb ₂ 0 ₄ α-sb ₂ 0 ₄ | 1223 " | 2. | sealed Pt tube sealed Pt tube | not melted | β + tr α α + Sb ₂ 0 ₃ |
| α-Sb204 β-Sb204 | 1303 | 19 [.] | scaled Pt tube | not melted | β + α β |
| $\alpha-sb_2O_4$ $\beta-sb_2O_4$ | 1327 JI | .08 | sealed Pt tube | not melted | α+β , β+α |
| a-sb ₂ 0 ₄ | 1330 | .25 | sealed Pt tube | not melted | β + α |
| α-sb ₂ 0 ₄ β-sb ₂ 0 ₄ | 1339 " | •08 11 | sealed Pt tube | not melted | α + β β + α |
| β-sb204 | 1345 | .08 | sealed Pt tube | , not melted | β + α |
| α-5b2 ⁰ 4 | 1350 | , <u>1</u> , 9 | sealed Pt tube | melted (vapor soaked into Pt) | |
| β-sb204 | 1350 | .08 | sealed Pt tube | melted ? large tabular vapor grown crystals | |
| α-sb ₂ 04 ^{<u>b</u>/} | 1200 | • - | high temperature x-ray | · | α (starting material remained α up to 1200°C) |
| a-sb204 c/ | 750 | 24 | open tray | • | ä |
| 976 9 11 11 | 800 900 950 | 97 * 11 87 | 11 TT 11 11 11 11 | • | α + β " " |

Table 4a. Experimental Data for Polymorphism in Antimony Tetroxide

 $\frac{\alpha}{2}$ The phases identified are given in the order of the amount present (greatest amount first) at room temperature. These phases are not necessarily those present at the temperature to which the specimen was heated. α refers to α -Sb₂O₄ polymorph and β to the β -Sb₂O₄ polymorph.

 $\frac{b}{b}$ Material placed on platinum slide and heated and examined by x-ray diffraction at various temperatures.

<u>C</u>/ Poorly crystalline as received Sb₂O₄ was heated 750°C - 24 hours and the same specimen which was never ground was reheated at 800°C - 24 hours, then 900°C - 64 hours and finally 950°C - 24 hours.

| Composition Starting Material | Heat Tr Temp °C | eatment Time hrs | Environme | nt | Pressure psi | Results ^{b/} X-ray Diffraction Analysis |
|-----------------------------------|-----------------------|------------------------|-------------------|---------------------|-----------------|--|
| α-Sb ₂ 0, ^A | 700 | 24 | Sealed Au t | ube | BB,000 | $\beta^{\underline{c}'} + sb_2 0_3^{\underline{d}'}$ |
| 2 4 | 750 | 48 | 98 B1 | #1 | 59,680 | n n "n" |
| 11 | 750 | 96 | 1) E | " | 73,200 | 10 01 00 |
| 13 | 750 | 16 | 11 11 | 11 | 89,400 | β + trace Sb ₂ O ₃ |
| N. | 751 | 116 | 11 11 | | 109,000 | β + sb ₂ 0 ₃ |
| U U | 760 | 96 | Sealed Au tube wi | th PtO, | 80,000 | β |
| ti , | 766 | 96 | Sealed Au t | ube | 88,000 | $\beta + Sb_2O_3$ |
| U | 775 | 115 | " Pt | ** | 47,500 | $\alpha + 5b_2O_3$ |
| u | 775 | 48 | " Au | H | 54,760 | $\beta + sb_2 o_3$ |
| 11 | 775 | 48 | " Pt | 11 | 66,500 | # N _ U |
| н | 800 | -24 | " Au | 44 | 93,000 | 16 41 II |
| \$ 7 | 800 | 24 | Sealed Au tube wi | th PtO ₂ | 105,000 | β |
| н | 850 | 16 | Sealed Au t | ube | 82,500 | $\beta + sb_2 0_3$ |
| 4 H | 900 | 72 | Sealed Pt tube wi | th PtO2 | 104,000 | β |
| β-Sb ₂ 04 | 900 | 72 | n 14 n | n u | 104,000 | ß |

Table 4b. Experimental High Pressure Data for Polymorphism in Antimony-Tetroxide.

 $\frac{a}{\alpha} - 5b_20_4$ prepared by the oxidation of 5b at 530°C on Pt tray. This material was reheated at 800°C - 60 hrs.

b/ The phases identified are given in the order of the amount present (greatest amount first) at room temperature. These phases are not necessarily those present at the temperatures to which the specimen was heated.

⊆ β form of Sb₂0₄.

 $\frac{d}{d}$ High pressure form of sb_2o_3 (valentinite).

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| Composition | | Heat Tre | atment ^a / | Results | | | | |
|---------------------------|-----------------------------|--|---|--|---|--|--|--|
| K ₂ 0 HõleX | Sb ₂ 0, Molež | Temp °C | Time hr | Physical Observation | X-ray Diffraction Analysis ²⁷ | | | |
| 5 | 95 | 950 1168 | 60 48 | not melted | pyrochlore as $+ \alpha$ -Sb ₂ O ₄ + β -Sb ₂ O ₄ = α -Sb ₂ O ₄ + β -Sb ₂ O ₄ = β | | | |
| 10 | 90 • | 950 1168 | 60 48 | not melted | pyrochlore as $+ \alpha - 5b_20_4 + \beta - 5b_20_4 \frac{c}{2}$ | | | |
| 15 | 85 | 853 950 966 | 24 60 4 | not melted """ """ | pyrochlore ss | | | |
| | | 1200 | 48 19 | U H | pyrochlore ss + α -Sb ₂ 0 ₄ + β -Sb ₂ 0 ₄ | | | |
| 20 | 80 | 950 116B | 60 48 | not melted | pyrochlore ss | | | |
| 25 | 75 | 950 1179 1361 1375 | 60 48 •08 •08 | not melted """" """ partially melted | P2 ₁ /c ^{d/} + pyrochlore ss pyfochlore ss pyrochlore ss | | | |
| | | 1385 1403 | .08 .08 | completely melted | | | | |
| 30 | 70 | 950 1178 1366 | 60 48 •08 | not melted H H H H | P2 ₁ /c ^{<u>d</u>/ + pyrochlore ss 1:2 + pyrochlore ss} | | | |
| · | | 1380 1382 1399 | .08 .08 .08 | partially melted """ completely melted | pyrochlore ss + 3:5 | | | |
| 33,33 | 66.67 | 950 950 998 1050 1050 1102 | 60 64 70 168 168 1 | not melted U 91 U 11 U 11 U 11 U 11 U 11 U 11 | $3:5 + P2_1/c^{d/}$ $P2_1/c^{d/}$ $1:2\frac{c}{c}/+3:5 + pyrochlore = + P2_1/c^{d/}$ | | | |
| · , | • | 1106 1106 <u></u> 1160 <u></u> 1179 1214 <u></u> 1214 | 64 64 1 48 1 2 | 11 17 11 11 11 15 11 15 11 11 11 11 | 1:227 + 3:5 + pyrochlore 1:2 + 3:5 3:5 + pyrochlore ss 1:2 + 3:5 + pyrochlore ss 3:5 + pyrochlore ss "" | | | |
| 35 | 65 ' | 950 1178 1380 1397 1409 | 60 48 .08 .08 .08 | not melted """ partially melted "" completely melted | pyrochlore + 3:5 + pyrochlore ss 1:2 + 3:5 3:5 + pyrochlora | | | |
| 37.5 | 62.5 | 950 1174 1195 1208 950 1310 1352 1379 1399 1416 | 60 88 19 1 64 45 .08 .08 .08 .08 | not melted """""""""""""""""""""""""""""""""""" | 3:5 3:5 + trace cubic 3:5 + trace 1:1 ^{E/} | | | |
| 40 | 60 | 950 1174 1208 1295 <u>e</u> / 1362 <u>e</u> / 1375 <u>e</u> / | 60 88 1 20 .5 | not melted """" """ """ """ partially melted | 3:5 + cubic """" 3:5 + 1:1 """ | | | |

| 45 | 55 | 950 1174 1208 1311 <u></u> | 60 88 1 1 | not melted U U U U U U | 1:1 + cubic + P2 ₁ /c cubic + 3:5 3:5 + cubic 3:5 + 1:1 |
|------|------|---|---|---|---|
| 46 | 54 | 1200 <u>h</u> / | l | not melted | cubic + 3:5 |
| 47 | 53 | 1194 <u>h</u> / 1200 | 3 1 | not melted # | cubic + trace 3:5 cubic |
| 47.5 | 52.5 | 1212 <u>h</u> / 1218 <u>h</u> / 1310 <u>8.h</u> / | 88 17 45 | not melted | cubic + 3:5 + 1:1 cubic + 1:1 + 3:5 1:1 |
| 48 | 52 | 1198 1200 <u>1</u> / 1200 <u>1</u> / 1308 <u>c</u> / 1103 <u>1</u> / | 3 1 1:5 .5 1 3 | not melted """""""""""""""""""""""""""""""""""" | cubic " cubic + 3:5 ilmenite 1:1 cubic + ilmenite + pyrochlore ilmenite + pyrochlore |
| 49 | 51 | 1200 | 1 | not melted | cubic |
| 50 | 50 | 750 800 921 946 950 1103 1104 1150 1174 1194 1202 1214 1298 1363 1403 1421 1426 | 70 24 1 21 60 1 22 1 88 1 1 .5 .5 .08 .08 | not melted """" """" """" """" """" """" """" """" """" """" """" """" """" """" """" """" """"" """""" | |

- <u>a</u>/ All specimens were preheated to 500 and 700°C for 60 hours unless otherwise footnoted. Rate of heating and cooling were approximately 3°/min. Specimens were heated in sealed Pt tubes and quenched from temperatures indicated.
- b/ The phases identified are given in order of the amount present (greatest amount first) at room temperature. These phases are not necessarily those present at the temperature to which the specimen was heated. 1:2 - K₂0.2Sb₂0₅; 3:5 - 3K₂0.5Sb₂0₅ and 1:1 - KSb0₃ - ilmenite structure.

c/ Non-equilibrium mixture - see Discussion in text.

<u>d</u>/ The phase was indexed from single crystal x-ray precession data which has shown the compound is monoclinic space group P2₁/c a=7.178, b=13.37B, c=11.985, B=124°10'.

 $\frac{a}{T}$ This specimen was previously heated to 500°, 700° and 1200°C - 19 hours in a sealed Pt tube.

 \underline{f}' Specimen heated in open Pt tube.

g/ Specimen leaked and changed composition.

 $\frac{h}{composition}$ prepared from a mixture 1:1 and 3:5 - see text for explanation.

 $\frac{1}{2}$ Specimen calcined and examined by x-ray diffraction while in form of pellet.

| | | Heat Tr | eatment ^a / | |
|---|-------------------------|--------------------------------------|----------------------------|---|
| | | Temp | Time | |
| Composition | Mole X | •C | hr | X-ray Analysis |
| NaSb03 Sb204 NaF | 75.08 3.15 21.77 | 1250 | 19 | single phase distorted cubic |
| NaSb03 Sb204 NaF | 67.79 6.25 25.96 | 1250 | 19 | body centered cubic + pyrochlore + ilmenite |
| NaSb03 Sb204 NaF | 53,50 12,34 34,16 | 1250 | 19 | body centered cubic + pyrochlore + sodium fluoride |
| NaSbO3 Sb204 Naf | 39.59 18.27 42.14 | 1250 | 19 | body centered cubic + pyrochlore + sodium fluoride |
| NaSb03 Sb204 NaF | 69.05 2.90 28.05 | 1250 | 19 | body centered cubic + trace godium fluoride |
| NaSb03 Sb204 NaF | 49.28 11.37 39.35 | 1250 | 19 | pyrochlore + body centered cubic + eodium fluoride |
| NaSbO3 Sb2O4 NaF | 31.20 28.87 39.93 | 1250 | 19 | pyrochlore + sodium fluoride |
| NaSb03 Sb204 NaF | 84.62 | 1268 | 19 | ilmenite + cubic |
| NaSbO3 Sb204 NaF | 74.42 2.32 23.26 | 1261 1268 | 1 19 | distorted cubic + ilmenits distorted cubic + NaF |
| NaSbO ₃ Sb ₂ O ₄ NaF | 70.00 3.33 26.67 | 1264 | 1 | cubic + ilmenite |
| NaSb03 Sb204 NaF | 65.96 4.26 29.78 | 1266 1267 | 1 19 | cubic + ilmenite cubic + NeF |
| Nasbo ₃ Sb.,04 Naf | 62.96 4.94 32.10 | 1266 1267 | 2 19 | cubic + NaF cubic + NaF |
| Na5b03 Sb204 NaF | 58.82 5.89 35.29 | 1267 | 19 | cubic + NaF |
| NaSbO3 Sb204 NaF | 68.00 4.00 28.00 | 1000 1252 1265 1265 1265 | 1 16 .1 1.5 72 | ilmenite + trace NaF cubic + trace ilmenite cubic + NaF cubic + NaF cubic + NaF |

Table 6. Experimental Data for the Ternary System NaSb03-5b204-NaF.

 $\stackrel{\text{al}}{=}$ Preheated at 750°C for 60 hours open.

| Composition | Temp. °C | Time hr | Results of X-ray Diffraction Analysis |
|--|-------------|------------|--|
| Na20:Bi203 | 500 500 | 96 120 | $Bi_{2}O_{3} + tr unknown Bi_{2}O_{3}$ |
| ^K 2 ^{0:B1} 2 ⁰ 3 | 500 500 | 66 120 | ^{Bi} ₁₂ ^O 3 |
| ^K 2 ^{0:3B1} 2 ⁰ 3 | 500 | 66 | Bi203 |

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Table 7. Experimental Data for Alkali Bismuth Oxide Systems.

Table 8. Experimental Data for Alkali-Rare Earth Systems.

| Composition | Temp °C | Time hr | Results of X-ray Diffraction Analysis |
|--|-----------------------------|---|--|
| Na20:Nd203 | 800 | 23 | a-Nd203 |
| K201Nd203 | 950 | 66 | $\alpha - Nd_2O_3 + ?$ |
| ^{K20:3Nd203} | 950 | 66 | $\alpha - Nd_2O_3 + ?$ |
| 3K ₂ 0:Y ₂ 0 ₃ | 600 700 800 950 | 20 20 20 66 | $Y_{203} + K_{203}$ """" |
| ^к 2 ^{0:Ч2⁰3} | 500 | 24 | ¥2 ⁰ 3 |
| ^{Na} 2 ^{O:Sm} 2 ^O 3 | 800 950 | 23 1.5 | Sm203 |
| Na20:35m203 | 800 950 | 23 1.5 | Sm 2 ⁰ ₁₁ 3 |
| 89.6Na20:10.4Sm203 | ∿1350 | 1 <u>*</u> / | Sm203 |
| ^K 2 ^{O:3Sm} 2 ^O 3 | 800 950 | 23 1.5 | Sm2013 |
| κ ₂ 0:Sm ₂ 0 ₃ | 800 950 1250 1350 | 23 1.5 16.5 <u>b</u> / 17.5 <u>b</u> / | $\frac{\text{Sm}_{2_{11}}^{0}}{\text{Sm}_{2_{12}}^{0}}$ + ? $\frac{\text{Sm}_{2_{12}}^{0}}{\text{Sm}_{2_{13}}^{0}}$ + ? |
| к ₂ 0:Gd ₂ 0 ₃ | 750 1000 1365 1370 | 90 92 67 <u>b</u> / 16 <u>b</u> / | $B-Gd_2O_3 + C-Gd_2O_3$ |

 $\frac{a}{a}$ The sample was heated in an open Pt crucible in an induction heater and the temperature recorded via an optical pyrometer,

 $\frac{b}{d}$ Quenched from temperature indicated.

| Composition | Temp °C | Time hr | Results of X-ray Diffraction Analysis |
|-------------------|------------|---------------------|--|
| 40.5K_0:59.5Nb_0_ | 600 | 40 | |
| 2 2 2 5 | 750 | 84 | |
| | 1000 | 48 . | 2:3 hydrate + TTB |
| | 1000 | 48 <mark>a</mark> / | 2:3 anhydrous + TTB |
| 41K_0:59Nb_0_ | 600 | 48 | |
| 2 2 5 | 700 | 85 | |
| | 750 | 60 | |
| | 800 | 90, | |
| | 1000 | 48 <mark>a</mark> / | 2:3 anhydrous |
| | 1000 | 48, | 2:3 hydrate |
| | 1000 | . 48 <u>.</u> | 2:3 anhydrous |
| | 1000 | 60 <u>b</u> / | 11 11 |

Table 9. Further Experimental Data for the System Nb205-KNb03.

 \underline{a}^{\prime} Slide heated @ 220° for 96 hours.

 \underline{b} / Slide heated > 125°C.

c' Slide heated to 110°C.

Table 10s. Summary of Ion Exchange Experiments

| EXCHANCE MEDIUM | TPC | TIKE | RESULTS |
|------------------------------|-------------------------|-----------------------|---|
| | 7K2011 | | |
| XaNO 3 | 340* | 2 hr | Decomposed to KMb0 ₃ + KMb ₃ 0 ₅ + H-Mb ₃ 0 ₅ |
| 3HaHO212NaMO3 | 244* 244* | 24 hr 72 hr | Unchanged 7:13(TTB)+Na-TTB(?) Unchanged 7:13 + NaMbO ₃ (tr) |
| нано ₃ (аq)≜/ | 104* | 2 hr | No change |
| | 18,75K ₂ 0;8 | 1,25%b205 | |
| NaNOJ | ~\$00" 330" | 2 hr 2 hr | NaMbOg H-Mb ₂ O ₃ + residual TTB _p |
| | 11,3K20:80 | 1.5Nb205 | |
| HaNO3(ad)=/ | 100* | 2 hr | Decomposed to $H-Nb_2O_5 + TTB_g$ |
| | 2K201 | 3ND 205 | |
| LOHTH CHAR | ~500* 340* | 2 hr 1 hr | NaNbO, perovakite Decomposed to KNb ₃ 0 ₈ + KNbO ₃ |
| NaNO3 (aq) | 100" 25" 25" | 2 hr 72 hr 4 hr | Decomposed to NaHbO, higher hydrats (b=47.2) + TTBd/ higher hydrate (b=42.2) + TTBd/ |
| NaOH (aq) b/ | 25* | 18 hr | higher hydrate + TTEd/ |
| H ₂ O (deionized) | 25* | 18 hr | No change |
| NaI (ac) ^{_/} | 25* | 18 hr | Unchanged 2:3 + TTB + higher hydrate (acetonstel)4 |

| | 41K20159M | 2 ⁰ 5 | |
|-------------------|----------------------------|-----------------------|--|
| | As prepared As prepared | hydrated anhydrous | Hydrated "2:3" Single phase "2:3" |
| Nano ₃ | 450 500 450 | 3,5 1 4 | Complete exchange Complete exchange Pallete hydrated and decrepitated while being x-rayed |

| | KTAWO6 | | |
|------------------------------|--------------------------------------|--------------|--|
| C ^{OKaH} | ~500" 315" | 2 hr 2 hr | NeTeO ₂ two places: KTaVO ₆ + NaTaVO ₆ |
| HaNO ₃ (aq)#/ | 105* | 1 hr | NaTaWO aw10,375 |
| RbNO3(sq)#/ | 105* | 1 hŕ | RbTaWO, (a=10.352) + unknown phase |
| H ₂ O (deionised) | 100* | 1 hr | KTAWO6 |
| | K,3 ^{TA} .3 ^W .7 | °, ' | • |
| HaNO3 | **500* 340* | 2 hr 2 hr | NeTEO, + HTB No chinge |
| | K20:2T8203 | | |
| Hano3 | ~500 * | 2 hr | No-change using sither 900° (rhomb) or 1300° (TTB) calcine starting material |

48520:11N5205#/

| <u>Bingle Zu</u> | change | |
|--------------------------|---------------------------------|--|
| 400* | 3 hr | K exch. 11 L phase a=7.354 |
| 400* | 16 hr | K exch. 11 L phase |
| 400* 450* | 16 hr 36 | K exch. 11 L phase K exch, 11 L phase |
| 500 450 414 400 | 1 hr 64 hr 17 hr 16 hr | Partially exch. e=7.4 c=43.6 Partially decomposed Partially exch. e=7.458 c=43.19 Partially exch. e=7.472 c=43.23 |

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NaNO3

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Double Exchange

| $KNO_3/400^*/3$ hr + NaNO_3/400*/1.5 hr + NaNO_3/400*/16 hr | 4-7.458 a=7.368 | c=43.398 c=43.869 | |
|--|--------------------|----------------------|--|
| KNO ₃ /400*/16 hr + NaNO ₃ /450*/66 hr | a=7.366 | c=43.898 | |
| KNO ₃ /400°/16 hr + NaNO ₃ /450°/98 hr | a=7.332 | c=43.997 | |
| Rb2013Hb205 | | | |

450*

450 450

450*

KNO₃

21.75Rb_0:78.25Nb_05

Single Exchange

KNO 3

NaNO₃

Partial exchange, increase in 'c' and partial decomposition

Double Exchange

 $KNO_{3}/450^{\circ}/18 \text{ hr} \rightarrow NaNO_{3}/450^{\circ}/23 \text{ hr}$

Partial exchange

No exchange (?)

RbTa₂O₅F (a=7,519 c=3.888)

18 hr 99 hr

99 hr

Single Exchange

| KNO3 | 340* | 2.5 | Partial e: | Kch. (7) |
|---|--------------|----------|------------------------|----------|
| | 340° 450° | 15 16 | | 0 |
| NaNO ₃ (aq. 20 w %) NaNO ₃ | 340* | 3.5 | No exchan No exchan | Be Be |
| | | | | |

Double Exchange

 $KNO_3/450^{\circ}/16$ hr + $NaNO_3/340^{\circ}/2$ hr

KN03/450*/16 hr + NaN03/450*/44.5 hr

No exchange

Some decomposition + partial exchange a=7.51 c=3.814

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Rb1-5xNb3.4+x09 Unchanged HTB + $Na_2Nb_4O_{11}$ (tr) + $NaNbO_3$ (tr) ~500° 2 hr NaNO₃ 330* 2 hr No change 3NaNO2:2NaNO3 240° 22 hr No change ∿500° 2 hr KNbO₃ + unchanged HTB KNO₃

 $\frac{m}{20}$ percent by weight of nitrate in deionized water

 $\frac{b}{20}$ percent by weight of hydroxide in deionized water

e/ 10 percent by weight salt in acètone

d/ when dried at 220°C this 'higher hydrate" or "acetonate" changes to a 2:3 type structure with <u>a</u> and <u>c</u> the same as original but with a <u>very</u> small <u>b</u> axis of 30.78A by comparison with the original <u>b</u>=33.019A.

e/ single crystal fragments

| Starting Moterial | Pressing History | Firing History | K ⁺ Exchange History | Na Exchange History | Remarks |
|--|--|---|---|--|--|
| 4Rb ₂ 0:11Nb ₂ 0 ₅ 500 °C-90 hr <u>c</u> / | 10000 psi <u>a</u>) 20000 psi <u>b</u> / | 1200 *-7 hr | | | Pellets cracked during firing a = 7.522 c = 43.180 |
| | | | RNO3-20hr @ 400° partial exchange ≞/ | | Pellet fragment disintegrated a = 7.128, c = 43.469 |
| | | + 15°/hr 1200 *-7 hr | KNO ₃ -16 hr @ 400 ⁴ partial ex- change | NeNO -18 hr @400 ^{3.} C partial ex- change | Fellet fragment did not dimintagrate K ⁺ a = 7.575, c = 43.469 Na ⁺ a = 7.390, c = 43.663 |
| | | †5°/minute 1200°∼1hr †5°/minute | KN0,-1 hr @400* complete exchange | | pellet did not disintegrate K ⁺ a = 7.545, c = 43.332 |
| | | †3°/minute 1200-1 hr †3° minute | KNO ₂ -19 hr 0400° complete exchange | | pellet displayed some spalling K ⁺ a = 7.555 c = 43.428 |
| 500-120 hr 600-90 hr <u>d</u> / | 10000 psi <u>s</u> / | 1100°C- 1 hr | KNO ₃ -16 hr G400° complete exchange | NaNO ₃ -21 hr G400 ³ | K^+ = 7.551, c = 43.401 Na ⁺ cells not celculated due to broadness of lines in diffraction |
| | | 1100°C-1.5 hr | KNO, 16 hr @400 partial exchange | NaNO41 hr @400 ³ partial exchange | Na exchanged pellet decomposed $K^+ = 7.547$, c = 43.428 Na ⁺ a = 7.404, c = 43.332 |
| | | 1100°C-1 hr | KNO ₁ -72 hr Ø400* completa exchange | NaNO32 hr 6300 ³ partial exchange | K ⁺ .z = 7.557, c = 43.442 Na ⁺ a = 7.484, c = 43.482 |
| · | 10000 pei <u>a/</u> 20000 pei <u>b</u> / | 1100-1hr 1200-1 hr 1100 - 16 hr 1200 -2 hr | KNO ₃ - 16 hr @400 ^s complete ex- change | | Pellet cracked during final 1200 ° firing K ⁺ a = 7,557 c = 43,442 |
| 500-90 | | †5°/minute 1200-5 hr †5° minute | KNO ₃ -2 hr 400°C ³ complete exchange | | powder fired in open tube K ⁴ a = 7.549, c = 43.387 |
| | | | KNO ₃ -18hr <u>d</u> / G400° complete exchange | NaNO ₃ =16 hr 0400 ³ complete exchange | $k^+ a = 7.553$ c = 43.387 $Na^+ a = 7.375$ c = 43.830 |
| 1 | a/= hydraulic | | | | c - 43,030 |
| | • pressure <u>b</u> /= Isostatic | | | | |
| | pressure c/= Without PV | ٨ | | | |
| | d/ with PVA | | | | |
| · · · · | e/ Complete exchange | | | | |
| | RbNb - KNb - NaNb - | 4 b 7.522Å 43.1 7.554Å 43.3 7.368Å 43.8 | 80Å 98Å 69Å | · · | • · · |

Table 10b. Summary of Ion Exchange Experiments with 4Rb_0:11Nb_0, Pellets,

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Table 11. Alkali Fluoride - Sb_2O_4 : Flux Evaporation

| Composition | Temp/Time | Results | | | |
|-----------------------------------|---|---|--|--|--|
| 98KF:2Sb204 | 1100°-1200°C 6 hrs | Solid + liquid x-ray shows KSbO small crystals form | | | |
| 88.0KF:7.2KSb0 ₃ :4.88 | ³⁶ 2 ⁰ 4 1100° - 1 hr | Solid + liquid small crystals formed | | | |
| 90KF:2Sb204:8B203 | 1100° - 2 hr, slow cool to 1000° | Solid + liquid small crystals formed | | | |
| 95RbF:58b204 | 900° - 1000°C 1 hr | Solid + liquid small crystals formed | | | |

Table 12. Summary of X-ray Data.

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| | Composition | | | | | | | Cozditiona | | |
|------------|--------------------------|-----------------------------|----------------|---------------|----------------------|--------|----------|-------------|---|-------------------|
| \$y+:+= | Designation | Alkeli Hetel Oxide Oxide | Hetel Orida | fundada e 710 | Unit Cell Dimensions | | | Liniting | Probable | |
| | | | •••••• | | b | 4 | F | Reflections | | |
| | | Holel | Molei | | 1 | 1 | 1 | | | |
| xb20+Xb303 | 11-L | 26.67 | 73,33 | Hemagona 1 | 7.522 | • | 43.18 | • | half-ktatl=3n | 22,83,832,838,836 |
| | 10-0 | 2212 | 12.2 | Rentform | 7.310 | - | 02.12 | - | hht:t=2n | 26,86,7426,763mc |
| | RTA | 21.75 | 11.3 | Orthorhombic | 12.991 | 7.530 | 7,794 | - | hkQih+k=2n | 76362,7676,763mmc |
| | | | | | | | | | hOL:1=7n | 200 00 |
| | GT B | 11.5 | 88,5 | Tetragonal | \$7,484 | - | 7,9737 | • | h0014+2n | 742,2,722,8 |
| Rb20-Ta305 | 11-L | 26.67 | 73.33 | Hazagenel | 7.506 | - | 43,19 | - | hk[1-h+k+1=3n | A1.FJ.(11.A30.A30 |
| | 9-1. | | • | Hexagonal | 7.306 | * | 36,41 | • | hhljf=2n | Hime Hiller Forme |
| | HTD | 21.75 | 78,23 | Hexegonal | 7.331 | • | 3,907 | + | Dene | P 6/ Spiele |
| | GTB | 11-2 | 48.5 | Tetragonal | 27.573 | • | 3.9018 | • | h00th=2a | 242122523m |
| ¥20~8b204 | pyrachiore as | (veriable) | | | | | | | | - • |
| | | 15 | - 4 5 | Cubic | 10.331 | - | - | - | hklib+k,k+l=2n | 7d3a |
| | | 30 | 70 | Cubin | 10,381 | • | • | | Oktik+l-én | |
| | 117 | 33.33 | 46,47 | Monoclinic | 19,473 | 7.452 | 7,198 | 94"54.4" | hklih+k=2n | 62/# (ij) |
| | ¥11/c | 33*33 | 66.67 | Nondellaic | 7,178 | 13,378 | 11.945 | 124*10' | hklino conditions hOlil=2n OLOIk=2n | 221/e |
| | 315 | 37.5 | 62.3 | Orthorhombic | 26.274 | 7.157 | 7.334 | - | hkino conditions Okitk=2n hOlih=2n hkOlno conditione | 6 1 /101 |
| | | | | | | | | | OOLIND conditions | (21) |
| ¥420-83204 | pyrachlore en (verieble) | | | | | | | | | |
| | | 25 | 75 | Cubic | 10,289 | • | - | | hkt;h+k,k+t=2n | ¥43- |
| | | 37,5 | 62,5 | CAPTE | 10,206 | * | - | - | Obliktiein | * = ** |

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