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ALKALI OXIDE-TANTALUM OXIDE AND ALKALI OXIDE-NIOBIUM OXIDE IONIC CONDUCTORS

by

R. S. Roth, H. S. Parker, W. S. Brower and D. Minor

prepared for

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### SUMMARY

This report summarizes work carried out between October 1972 and October 1973 under an agreement with the National Aeronautics and Space Administration, Lewis Research Center (Interagency Order C-29933-C) to study the phase equilibria of alkali oxide-tantalum and niobium oxide systems and synthesis of phases which might have interesting ionic conductivity.

The phase equilibrium relations of six systems were investigated in detail. These consisted of lithium, sodium and potassium tantalate, and niobate with the corresponding tantalum and niobium oxide. In addition, various other binary and ternary systems involving alkali oxides were examined in lesser detail. In all, thirteen phases were found to contain structures of sufficient interest to warrant further investigation. Nineteen phases were prepared as solid pellets and transmitted to LeRC for evaluation by dielectric loss measurement. Attempts were made to grow eight different phases as large single crystals, and two were submitted to LeRC for evaluation.

In terms of ionic conductivity, the most interesting phases from a structural point of view are those containing large tunnels only partially occupied by alkali ions. The potassium tantalate system contained four such phases, designated GTB, TTBs, HTB, and TTB (see table 6 for meaning of these initials). In all cases, the TB refers to a tungsten bronze type structure. Those phases which exhibit a maximum in the dielectric loss vs. temperature curves proved to be high temperature phases which are generally metastable at room temperature and do not have favorable melting characteristics for crystal growth directly from the melt.

In the coming year efforts will be expanded to include rubidium systems as well as antimonates and mixed tantalum-tungstates, in an effort to isolate an ionic conducting phase with favorable crystal growth possiblities.

### INTRODUCTION

In the search for new ionic conductors, compounds in oxide systems need to be examined for structure types where packing of the complex ions of the skeletal structure has the potential to produce loosely bound alkali ions.

The compounds of alkali oxides with tantalum oxide and with niobium oxide represent a class of such materials. The program summarized in this report involved the investigation of the phase diagrams to determine the phases which are actually contained in these systems and to define selected phases of potential interest as solid ionic conductors. Subsequent to this determination specimens were prepared of single phase polycrystalline ceramic pellets for screening measurements and, finally, attempts were made to grow single crystals of those materials which appeared most interesting and feasible. All the phases encountered were identified by single crystal x-ray diffraction patterns and/or identity with previously reported structures. For fast ion transport, it is probably advantageous to have an alkali ion in a "non-stoichiometric" crystallographic position. For commerical utilization of ionic conductivity, it is necessary to limit systems to those which will contain little or no electronic conductivity and, therefore, do not contain an element which is easily reduced during the synthesis. The Ta<sup>+5</sup> ion is apparently very difficult to reduce [1], and thus is ideally suitable as a host lattice for alkali "superionic-conductivity". Some preliminary work on alkali tantalate and niobate systems has been previously summarized by the present authors [2]. A complete detailed analysis of these systems is presently being prepared for publication and will be submitted to the Journal of Solid State Chemistry in the near future [3, 4, 5, 6].

### DISCUSSION OF RESULTS

### The System Nb<sub>2</sub>O<sub>5</sub>-LiNbO<sub>3</sub>

A phase equilibrium diagram for the system  $\text{Li}_2\text{O-Nb}_2\text{O}_5$  was published by Reisman and Holtzberg [7]. However, this diagram does not show the compound  $\text{LiNb}_3\text{O}_8$  which is known to occur in this system [8, 9, 10], nor does it show the solid solution of  $\text{Nb}_2\text{O}_5$  in  $\text{LiNbO}_3$  [11] which has been found to be important in the production of large optical quality crystals. A phase equilibrium diagram indicating these corrections was previously published [2]; and this diagram is reproduced here, essentially unchanged, in figure 1. The experiments performed in this laboratory to check this diagram are listed in table 1. The unit cell dimensions of the phases will be given in the summary table of x-ray data near the end of this report (Table 12).

The  $\mathrm{Nb}_2\mathrm{O}_5$ -LiNbO $_3$  system contains only two intermediate phases, the compound  $\mathrm{LiNb}_3\mathrm{O}_8$  and another phase which occurs at a very narrow compositional range near  $\mathrm{Li}_2\mathrm{O}:14\mathrm{Nb}_2\mathrm{O}_5$ . This phase was reported independently by the present authors [2] and by Norin and Nolander [12] to have the N-Nb $_2\mathrm{O}_5$  structure. Only this phase and LiNbO $_3$  itself may be considered to have any interesting non-stoichiometric properties.

A detailed analysis of the work performed in this system, including x-ray diffraction patterns, will be published in the near future [3].

### The System Ta<sub>2</sub>O<sub>5</sub>-LiTaO<sub>3</sub>

No phase equilibrium diagram for this system had been published prior to our preliminary work reported in reference [2]. The final phase diagram is shown in figure 2 and the experimental data from which it was constructed are listed in table 2. The present diagram differs from that given in [2] only in the first 5 mole percent Li<sub>2</sub>O content and near the melting point of LiTa<sub>3</sub>O<sub>8</sub>.

Lithium oxide has been found to stabilize the low temperature polymorph of  ${\rm Ta_2O_5}$  (L-Ta<sub>2</sub>O<sub>5</sub>); and, therefore, non-stoichiometric solid solutions appear on both sides of this system. There is only one intermediate compound, LiTa<sub>3</sub>O<sub>8</sub>, but it is trimorphic and apparently has no appre-

ciable non-stoichiometric region within the binary phase diagram. The low temperature form of  ${\rm LiTa_3O_8}$  is isostructural with  ${\rm LiNb_3O_8}$ , the intermediate form is isostructural with the mineral wodginite, and the high temperature form is similar in structure to  ${\rm LiTa_6O_15^F}$  and  ${\rm LiNb_6O_15^F}$  [13].

The unit cell dimensions will be listed in the summary table of x-ray data near the end of the report (Table 12) and a detailed analysis of the work performed in this system, including x-ray diffraction patterns, will be published in the near future [3].

# The System $Nb_2O_5$ -NaNbO3

This system has been studied by several different groups and phase equilibrium diagrams were published by Reisman, Holtzberg and Banks [14], and by Shafer and Roy [15]. The former group reported a 1:4 and a 1:14 compound whereas the second reference shows 1:4, 1:7, and 1:10 compounds. These type of ratios represent mole ratio of alkali oxide to niobium (or tantalum) oxide. Sten Andersson [16] has shown that the high Nb<sub>2</sub>O<sub>5</sub> content phase occurs at the 1:13 ratio (NaNb<sub>13</sub>O<sub>33</sub>) from crystal structure analyses [17] and found the other phase to occur at the 1:3 ratio (NaNb<sub>3</sub>O<sub>8</sub>). He found the compounds Na<sub>2</sub>Nb<sub>4</sub>O<sub>11</sub> and NaNb<sub>6</sub>O<sub>15</sub>(OH) only in hydrothermal experiments at 500-700°C and 2000 atm. In a crystal structure analysis of Andersson's NaNb<sub>3</sub>O<sub>8</sub> crystals, D. C. Craig and N. C. Stephenson decided that the composition of the crystal they examined was most likely Na<sub>13</sub>Nb<sub>35</sub>O<sub>94</sub> [18].

The phase equilibrium diagram which we have determined for this system is given in figure 3 and the experimental data from which it was determined are listed in table 3. We confirm the 1:13 compound but do not find any other between this and the "NaNb $_3$ O $_8$ " phase. The latter is apparently a non-stoichiometric solid solution with a tetragonal tungsten bronzetype substructure and an orthorhombic superstructure (TTBs). In addition Na $_2$ Nb $_4$ O $_{11}$  was found in equilibrium at low temperatures (below about 1000°C) and specimens near the 1:6 ratio was found to react with atmospheric moisture at temperatures between about 700-900°C to form NaNb $_6$ O $_{15}$ (OH) without the necessity for hydrothermal techniques.

The unit cell dimensions of these phases will be listed in the summary table of x-ray data near the end of the report and a detailed analysis of the work performed in this system, including x-ray diffraction patterns, will be published in the near future [4].

# The System Ta<sub>2</sub>C<sub>5</sub>-NaTaO<sub>3</sub>

The phase equilibrium diagram for this system had not been reported prior to our preliminary paper [2] although Reisman [19] had made a subsolidus study up to 1300°C. The diagram given in figure 4 is similar to that of [2] but has been reevaluated for the melting relations of the distorted tetragonal tungsten bronze-like phase (TTBs) between about 20 and 25 mole % Na<sub>2</sub>O. This is the only non-stoichiometric phase in the system although apparently a very small amount of Na<sub>2</sub>O may be incorporated in L-Ta<sub>2</sub>O<sub>5</sub>, increasing the phase transition temperature. The only other phase in the system is Na<sub>2</sub>Ta<sub>4</sub>O<sub>11</sub> which has a hexagonal structure similar to the monoclinic Na<sub>2</sub>Nb<sub>4</sub>O<sub>11</sub>.

The experimental data from which figure 4 has been constructed are listed in table 4. The unit cell dimensions of these phases will be listed in the summary table of x-ray data near the end of this report. A detailed analysis of the work performed in this system, including x-ray diffraction patterns, will be published in the near future [4].

# The System Nb<sub>2</sub>O<sub>5</sub>-KNbO<sub>3</sub>

A phase equilibrium diagram for this system was previously reported by Reisman and Holtzberg [20]. Additional information on compound formation, crystal growth, and unit cell dimensions has been published by Guerchais [21], Whiston and Smith [22], and Nassau et al [23]. The phase equilibrium diagram determined in the present study is shown in figure 5 and the experimental data from which it was constructed are listed in table 5.

The system was found to contain six intermediate phases. The unit cell dimensions of each phase are listed in the summary table near the end of this report. From these dimensions it is apparent that at least three of these phases have non-stoichiometric-type structures. These are designated in the phase diagram as GTB, TTBs, and TTB and occur at

about 11.5, 16-20, and 35 mole % K<sub>2</sub>O respectively. The former has a structure similar to that reported by Gatehouse for a rubidium niobate phase [24] and herein referred to as a Gatehouse tungsten bronze (GTB). The other two have a distorted tetragonal tungsten bronze structure (TTBs) and an undistorted 'bronze' structure (TTB) respectively.

The unit cell dimensions of the 2:3 phase indicate a layer of niobia octahedra with all the potash possibly located in planes. The hydrated form of this phase reveals different x-ray diffraction intensity distributions depending on the hydration direction and rate. This phenomena may possibly indicate different distributions of  $K^+$  ions which depend on the hydration. Ionic conductivity of this phase in a direction parallel to the cleavage, that is, parallel to the  $K^+$  and/or  $H_2^{0}$ 0 motion should be investigated more carefully.

A detailed analysis of the work performed in this system, including x-ray diffraction patterns, will be published in the near future [5].

# The System $Ta_2O_5$ -KTa $O_3$

The phase equilibrium diagram published for the system  $K_2O-Ta_2O_5$  by Reisman et al contained only two compounds between  $Ta_2O_5$  and  $KTaO_3$  [25]. The preliminary diagram published by the present authors [2] indicates nine equilibrium phases in this region and two other metastable phases. The present diagram, figure 6, differs from the previous only in the estimated width of the non-stoichiometric phases. The experimental data from which this diagram was constructed are listed in table 6.

The system was found to contain four stable non-stoichiometric regions in addition to five stable, apparently stoichiometric, phases. The phases with non-stoichiometric type structures include a Gatehouse tungsten bronze (GTB) at about 11.5 mole %  $\rm K_2O$ , an orthorhombic distorted tetragonal tungsten bronze with superstructure (TTBs) between about 15-20 mole %  $\rm K_2O$ , a hexagonal tungsten bronze (HTB) at about 21.75 mole %  $\rm K_2O$ , existing over a very narrow temperature region below the solidus, and an undistorted tetragonal tungsten bronze with no superstructure at  $\rm ^{\sim}34$  mole %  $\rm K_2O$ . Figures 7, 8, 9, and 10 show the results of measuring the dielectric loss (epsilon) versus temperature at a number of frequencies

for these four non-stoichiometric phases. The GTB and HTB phases show excellent ionic mobility. Unfortunately, single crystals of these phases have not yet been grown. The difficulty is due mainly to the very narrow primary phase field. An attempt must be made to find a system with wider primary phase regions for these structures, possibly in systems with Rb<sub>2</sub>O, which can then be ion exchanged.

### Other Systems

In the course of this study, a considerable number of specimens were investigated with compositions not included in the six systems described previously. The ternary systems involving these six systems with MoO<sub>3</sub> were all examined briefly for the purposes of growth of small crystals [2]. The compositions of the crystal growth preparations will be detailed at a later date [3, 4, 5, 6].

Several of these six binary systems were also examined in conjunction with WO<sub>3</sub>, especially the psuedo-binary joins NaTaO<sub>3</sub>-WO<sub>3</sub> and KTaO<sub>3</sub>-WO<sub>3</sub>. The NaTaO<sub>3</sub>-WO<sub>3</sub> system proved to be completely non-binary and contained mostly solid solutions of sodium tungstates in the sodium tantalate type phases. However, the KTaO<sub>3</sub>-WO<sub>3</sub> system is apparently a psuedo-binary system below the solidus and contains at least five phases. The compositions of the phases are about 25, 50, 50, 70, and 90 mole % WO<sub>3</sub>. The ternary system KTaO<sub>3</sub>-WO<sub>3</sub>-K<sub>2</sub>WO<sub>4</sub> has been found to be a promising flux system for crystal growth of both the pyrochlore and hexagonal tungsten bronze (HTB) type phases. Further work on this ternary will be conducted in the next year.

Some compositions in the  $\mathrm{Li_2O-Ta_2O_5-WO_3}$  system were prepared and examined for the purpose of introducing non-stoichiometry into the H-LiTa\_3O\_8 structure. Other compositions in the system  $\mathrm{Li_2O-Ta_2O_5-TiO_2}$  were also prepared for the same purpose. The WO\_3 gave single phase non-stoichiometric solid solutions but showed no ionic mobility (table 7). The  $\mathrm{TiO_2}$  specimens did not exhibit solid solution.

A study of the  $\mathrm{Li_2O\text{-}TiO_2}$  system or rather of the  $\mathrm{Li_2TiO_3\text{-}TiO_2}$  system revealed only one high temperature intermediate phase. This phase has the ramsdellite structure; but, unfortunately, melts incongruently with the peritectic very close in composition and temperature to the eutectic.

Thus, single phase crystals could not be pulled from the melt. Experiments in the  $\mathrm{Li}_2\text{O-TiO}_2\text{-MoO}_3$  ternary also did not reveal any ideal flux compositions for crystal growth of this phase. The ramsdellite lithium titanate did not exhibit appreciable ionic mobility (table 7). Other experiments with  $\mathrm{K}_2\mathrm{C-Li}_2\mathrm{C-TiO}_2$  and  $\mathrm{K}_2\mathrm{C-MgO-TiO}_2$  revealed a new structure type [26] which also failed the ionic mobility screening tests.

The only experiments performed under this contract which did not involve mixed oxides were conducted in the system NaF-AlF $_3$ -Na $_2$ GeF $_6$  in an attempt to produce Ge doped NaAlF $_4$ . These experiments were not successful and no NaAlF $_4$  was obtained.

A few compositions involving the systems  ${\rm NaSbO}_3{\rm -Sb}_2{\rm O}_4$  and  ${\rm KSbO}_3{\rm -Sb}_2{\rm O}_4$  have been prepared in order to survey possible compound formation in these systems, because Aurivillius [27] has reported K<sup>+</sup> ions to occur in large tunnels in some potassium antimonates. In addition, several specimens were examined in the ternary systems  ${\rm KSbO}_3{\rm -Sb}_2{\rm O}_4{\rm -SiO}_2$  and  ${\rm KSbO}_3{\rm -Sb}_2{\rm O}_4{\rm -Al}_2{\rm O}_3$  in an attempt to explain the published data of Spiegelberg [28] who reported a primitive cubic "polymorph" of  ${\rm KSbO}_3$  prepared in a porcelain crucible. These experiments were successful in demonstrating that a small amount of both  ${\rm Si}^{+4}$  and  ${\rm Al}^{+3}$  promote the formation of this cubic polymorph. These and similar antimonate systems will be examined in greater detail in the coming year.

### Polycrystalline Specimen Preparation

The sections of this report dealing with the phase equilibrium relationships in alkali niobate and tantalate systems have described several phases which, from structual considerations alone, would appear to be worthy of screening for their potential as fast ion conductors. Accordingly, polycrystalline specimens were prepared and submitted to Lewis Research Center for dielectric and conductivity measurements.

The choice of composition and thermal treatment for use in pellet fabrication of a desired phase was made on the basis of the compositional and temperature stability limits of the phase as determined in the phase equilibrium studies. These limits are not always compatible with the sintering characteristics of the powders and, as a consequence, it was

not possible to prepare high density pellets of some of the phases of interest. Several alkali titanate pellets of interest were also prepared as an outgrowth of work under a previous contract. All compositions for use in pellet fabrication were given blending and calcining treatments similar to those utilized in the preparation of compositions for phase equilibrium studies. In the case of phases wherein alkali loss was anticipated during sintering or where a minimum temperature of stability necessitated rapid cooling of the pellets, the exact composition of the pellet was chosen to give as much latitude as possible in the sintering treatment. The appropriate 1:1 alkali niobate or tantalate was used as one constituent of the batch in these systems to avoid the use of carbonates.

Conventional ceramic dry pressing and sintering techniques were utilized for pellet preparation. Dry pressing was performed in steel dies at nominal pressures of the order of 10,000 to 18,000 psi. No organic binders or other additives were utilized except as noted. In general, several test heatings were necessary to determine a suitable time-temperature relationship for sintering. In some cases, excessive recrystallization during sintering, regardless of the schedule followed, prevented the formation of physically sound pellets. As discussed below, hot pressing was found useful in some cases.

The starting compositions after calcining, as well as the as-fired surface and interior of the sintered pellets, were characterized by x-ray diffraction techniques to insure that the final pellets were homogeneous and single phase.

Tables 8, 9, and 10 summarize the compositions, heat treatments and x-ray characterizations of the pellets submitted for evaluation. The results of screening tests conducted by LeRC are given in Table 7.

### Crystal Growth

The growth of large single crystals of the various non-stoichiometric phases found in the alkali niobate and tantalate systems is complicated by many factors: (1) some of the desired phases do not exist in equilibrium with the liquid, (2) incongruency, and (3) volatility of the alkali. For these reasons only a few of the compounds found in the phase equilibrium

studies were grown successfully. These were the congruently melting compounds and those incongruent compounds which existed in equilibrium with a liquid over a wide temperature range. In all crystal growth attempts the molten material was contained in either platinum or iridium crucibles which were heated inductively and the desired phase pulled either by the conventional Czochralski technique, or by an approximation to top seeded solution growth.

A number of the more interesting phases do not lend themselves to growth by the more conventional techniques and it will be necessary to explore various flux growth techniques in an effort to obtain crystals of a size suitable for physical property measurements.

The crystal growth experiments (table 11) are summarized as follows:  ${\rm Nb_2O_5}{\rm -KNbO_3}$ 

### 7:13 Phase (TTB)

Growth of the 7:13 phase was attempted from a melt composition of  $36.5 \text{K}_2\text{O}:63.5 \text{Nb}_2\text{O}_5$ . The first material to crystallize was the 4:9 phase, on continued pulling a small amount of 7:13 phase crystallized out on the side of the 4:9 phase as a polycrystalline mass. As the pull continued the remainder consisted of single phase 2:3. Several different melt compositions were tried and the results were essentially the same, in all cases the 4:9 phase crystallizes out and as more material is removed the composition of the crystallizing phase shifts to the 2:3 phase. Apparently the temperature-composition region over which the 7:13 exists in equilibrium is too small to successfully obtain crystals by the pulling technique.

## $12K_2O:88Nb_2O_5$ (GTB)

Attempts to pull the (GTB) bronze-like phase  $12\text{K}_2\text{O}:88\text{Nb}_2\text{O}_5$  at a melt composition of  $15\text{K}_2\text{O}:85\text{Nb}_2\text{O}_5$  resulted in a polycrystalline multiphase mass. This result was not unexpected as an inspection of the phase diagram indicates a very narrow compositonal range where this phase is in equilibrium with the melt. This situation makes it extremely difficult to grow large crystals of the 12:88 phase any place in the  $\text{K}_2\text{O-Nb}_2\text{O}_5$ 

binary. Only by going to a flux system can one possibly expect to grow this phase.

# 17.5K<sub>2</sub>0:82.5Nb<sub>2</sub>0<sub>5</sub> (TTBs)

Attempts to grow the TTB phase from a melt composition of  $20K_2^{\circ}0:80{\rm Nb}_2^{\circ}0_5$  resulted in essentially single phase polycrystalline TTB with very fine grain size. This phase would be a likely candidate for top seeded solution growth (TSSG) or possibly the accelerated crucible rotation technique (ACR).

## Ta<sub>2</sub>O<sub>5</sub>:KTaO<sub>3</sub>

### TTB Phase

As determined from phase equilibria studies, the TTB phase melts incongruently. For the crystal growth attempts, a melt composition of  ${}^{45}{\rm K}_2{}^{0}:55{\rm Ta}_2{}^{0}{}_5$  was chosen. With this composition and using a pull rate of 0.2 in/hr crystals of the desired phase were grown. The maximum diameter attained was the order of 1/8 in. diameter at the stated pull rate. The growth of larger crystals would necessitate much slower growth rates, as with top seeded solution growth.

# ${\rm Ta_2O_5: LiTaO_3}$ System

### 1:3

Single crystals of the 1:3 phase were grown both "on composition" and from compositions slightly rich in Li<sub>2</sub>O. The "as-grown" crystals were light brown to colorless depending upon cooling rate. Subsequent annealing in air removed all traces of brown color. The reason for this change in color is not known.

#### 5:95 Phase

The 5:95 phase or L-Ta<sub>2</sub>O<sub>5</sub> has been grown in sizes up to as large as 2 cm long and 1.5 cm diameter. This is an incongruent melting compound and growth was by top seeded solution growth starting at a composition of  ${}^{15}\text{Li}_2\text{O}:85\text{Ta}_2\text{O}_5$ . These are the first large crystals of L-Ta<sub>2</sub>O<sub>5</sub> ever grown and are being used for a structure determination using neutron diffraction, in order to obtain a better knowledge of the mechanism of nonstoichiometry. The crystals are almost colorless when first grown but gradually turn yellow on exposure to light.

### Nb<sub>2</sub>O<sub>5</sub>-NaNbO<sub>3</sub> System

### 1:3 Phase

Single crystals of the 1:3 phase could be readily pulled from a melt of the stoichiometric composition, however, there was a problem with cracking which limited the size of usable crystals obtained. The cause of cracking was not investigated.

## Ta205-NaTa03 System

### 21:79 Phase (TTBs)

All attempts to grow crystals of the bronze-type phase by induction heating at ambient pressures were unsuccessful due to the rapid loss of  ${\rm Na}_2{\rm O}$  from the melt.

### Structural Mechanisms of Non-stoichiometry in Alkali-Niobates and Tantalates

An abnormally large number of phases have been found in the six systems examined and discussed in this report. In addition to the end members, the  $\text{Li}_2\text{O}$  systems contain two and three phases with  $\text{Nb}_2\text{O}_5$  and  $\text{Ta}_2\text{O}_5$  respectively whereas the  $\text{Na}_2\text{O}$  systems contain two and four phases each and the  $\text{K}_2\text{O}$  systems, six and nine phases each. The unit cell dimensions of each of these phases is listed in table 12 together with the pertinent crystallographic data where known.

The occurrence of the tungsten-bronze-type structures in these alkali niobate and tantalate binary systems is somewhat disconcerting, as the A\_xBO\_3 compositional range does not fall in these systems for any value other than x=1. It is quite evident, therefore, that it is impossible to have niobium and tantalum oxygen octahedra and pentagonal bipyramids form in a network structure in which only a non-stoichiometric amount of alkali ions compensate the framework for electrical neutrality. One possible hypothetical alternate of oxygen vacancies is both esthetically displeasing and structurally unsound. Furthermore, such structures have not been previously found to exist.

### HTB

The most likely explanation, and one based on preliminary structural evidence of a similar phase, is that of interstitial niobium (or tantalum) ions. B. M. Gatehouse [24] has shown that niobium can occur interstitially in a rubidium hexagonal tungsten bronze structure in the 9-fold tricapped-prism interstitial vacancy common to all of the tungsten bronze-type structures. Gatehouse has suggested that the composition of this hexagonal bronze phase would be RbNb<sub>3.4</sub>O<sub>9</sub> or 22.72 mole % Rb<sub>2</sub>O. Some hypotheses can be made, on the basis of Gatehouse's work, concerning the mechanism of non-stoichiometry in the phases observed in this study.

In the system  ${\rm Ta_2O_5}$ -KTaO\_3, a phase having the hexagonal tungsten bronze structure has been observed to occur at about 21.75 mole % K\_2O. By analogy to the RbNb\_3.4O\_9 phase, this can be considered to be  ${\rm KTa_3.4^O_9}$ . However, this formula would indicate 22.72 mole % K\_2O which is definitely not the case. The composition 21.75 mole % K\_2O would correspond to the formula K\_0.95 ${\rm Ta_3.41^O_9}$  and this non-stoichiometric phase must contain alkali vacancies as well as tantalum interstitials.

### TTB

This same analogy may be used to explain the non-stoichiometry in the tetragonal tungsten bronze phases which occur with no sign of superstructure in both the  ${\rm Ta_2O_5}$ -KTaO\_3 and  ${\rm Nb_2O_5}$ -KNbO\_3 systems. The formula for one unit cell of the tetragonal tungsten bronze structure is  ${\rm ^A6^B_{10}O_{30}}$ . If all the B ions have a valence of +5, then the formula for the ideal end member would be  ${\rm ^{K_1^{1}B_{10}^{+5}}}$  or 35.71 mole % K<sub>2</sub>O. In the  ${\rm ^{Nb_2O_5}}$ -KNbO\_3 system this phase has been observed at  ${\rm ^{Nb_2O_5}}$ -KTaO\_3 system, this phase has been observed at  ${\rm ^{Nb_2O_5}}$ -KTaO\_3 system, this phase has been observed at  ${\rm ^{Nb_2O_5}}$ -KTaO\_3 system, this phase has been observed at  ${\rm ^{Nb_2O_5}}$ -KTaO\_3 to the formula  ${\rm ^{K_5}}$ .45 ${\rm ^{Ta}_{10.91}}$ O<sub>30</sub>.

In his paper on rubidium niobates, Gatehouse discussed a new structure type containing 4-, 5-, 6-, and 7-sided tunnels which may be partially occupied by alkali ions. This phase has been called the Gatehouse tungsten bronze in the present paper and was observed to occur at about 11.5 mole %  $\rm K_2O$  in both the  $\rm Ta_2O_5$ -KTaO3 and  $\rm Nb_2O_5$ -KNbO3 systems. Gatehouse was unable to arrive at a correct composition for this phase, but in a personal communication confirmed the proposed structure: ". . . having recently obtained some magnificent lattice image photographs which completely confirm the basic structure -- 7-sided holes and all." The basic formula for this structure is  ${}^{A}_{8}{}^{B}_{54}{}^{O}_{146}$ . If all the B ions have a valence of +5 and the excess is supplied via interstitual B+5 ions then the ideal end member composition would be  $^{+1}_{8}^{+5}_{54}^{+5}_{4}$  + 2.8 $^{\circ}_{146}$ . This ideal formula would correspond to 12.35 mole % alkali oxide. The formula for the observed 11.5 mole % composition would be  $K_{7.4}B_{54} + 2.92O_{146}$  when B is either Nb<sup>+5</sup> or Ta<sup>+5</sup>. The solid solution region probably goes down to at least a composition of  $K_7B_{57}O_{146}$  (10.94) mole %  $K_2^0$ ) and possibly even slightly lower in  $K^+$  content. It should be pointed out that this hypothesis involves K ions in both the six and seven (or possible four) sided tunnels whereas Gatehouse explicitly stated that the seven- and four-sided tunnels appeared to be empty. It is obvious that the details of this structure need to be examined with more accurate data. Dr. Gatehouse has informed us that he is now refining three dimensional crystallographic data on his rubidium niobate, and NBS will attempt to do an accurate single crystal crystallographic study of the chemically well-characterized potassium niobate phase.

### TTBs

The case of the TTBs phase, that is, the orthorhombic tetragonal tungsten bronze with superstructure indicating that one of the <u>a</u> tetragonal axes is tripled, is more complex then the other three. This phase occurs in all four systems of sodium and potassium with niobium and tantalum. In the  $\mathrm{Nb}_2\mathrm{O}_5$ -NaNbO $_3$  system, it occurs between about 22.5 to 27.5 mole %  $\mathrm{Na}_2\mathrm{O}$  and between about 19.5 and 25.5 mole %  $\mathrm{Na}_2\mathrm{O}$  in the  $\mathrm{Ta}_2\mathrm{O}_5$ -NaTaO $_3$  system.

However, in the  $\mathrm{Nb}_2\mathrm{O}_5$ -KNbO $_3$  and  $\mathrm{Ta}_2\mathrm{O}_5$ -KTaO $_3$  systems, this same structure type occurs between about 15 and 20 mole %  $\mathrm{K}_2\mathrm{O}$ . Although the absolute limits of these have not been accurately located, these rough limits of about 15 to 27.5 mole % alkali oxide must be explained.

D. C. Craig and N. C. Stephenson [18] have examined the crystal structure of a specimen of NaNb<sub>3</sub>O<sub>8</sub> supplied by S. Andersson [16]. They concluded that the particular crystal examined had the composition Na<sub>13</sub>Nb<sub>35</sub>O<sub>94</sub> or 27.08 mole % Na<sub>2</sub>O rather than the 25 mole % Na<sub>2</sub>O suggested by Andersson. Stephenson reports that the tripled "tetragonal-tungsten-bronze" cell has a basic structure of  $^{\mathrm{Na}}_{6}$   $^{\mathrm{Nb}}_{34}$   $^{\mathrm{O}}_{94}$ and that twelve excess positive charges must be accommodated by Na and/or Nb distributed in the eight 5-sided tunnels. He considers only the cases of  $\mathrm{Nb}^{+5}$  +  $7\mathrm{Na}^{+}$  and  $2\mathrm{Nb}^{+5}$  +  $2\mathrm{Na}^{+}$  and says that the former more nearly coincides with the experimental intensities. In this paper [18], Craig and Stephenson dismiss the possibility of Nb in the three sided tunnels as being ". . . incompatible with the directional properties of the bonding orbitals of niobium (V)." This statement is subject to considerable doubt and leaves open the actual position of the "interstitial" niobium. Apparently Craig and Stephenson did not consider the possibility that this phase might have a non-integral number of  $\mathrm{Na}^{+1}$  and  $\mathrm{Nb}^{+5}$  ions, that is, be a composition in the middle of a non-stoichiometric solid solution.

The basic-structure of the tripled tetragonal tungsten bronze (TTBs) unit cell is, therefore,  $[A_6^{+1}B_{34}^{+5}O_{94}]^{-12}$  and contains a maximum of eight excess positions for alkali ions. The composition can, therefore, vary from a maximum alkali content of  $A_{14}^{+1}B_{34.8}^{+5}O_{94}$  with 28.7 mole % alkali oxide to a minimum alkali content of  $A_6^{+1}B_{36.4}^{+5}O_{94}$  with 14.15 mole % alkali oxide. It is noteworthy that the limits of the observed phases, 15 to 27.5 mole % alkali oxide, occur just within the theoretical minimum and maximum values of 14.15 to 28.7 mole % alkali oxide.

In the above four cases, therefore, the most likely structural mechanism for non-stoichiometry is that of transition metal interstitials plus alkali ion vacancies. However, other mechanisms can also be found in the six systems discussed in this report.

### Other Phases

The N-Nb $_2$ O $_5$  phase stabilized by Li $_2$ O can be considered to be Li $_2$ O:14Nb $_2$ O $_5$  or Li $_1$ 8(Li $_0$ .45Nb $_3$ 1.55)O $_8$ 0 with Li $^+$  ions substituting for Nb $^+$ 5 and also present in interstitial four-sided channels. This non-stoichiometric phase is thus accounted for by a mechanism of alkali substitution plus alkali interstitials.

The high temperature form of LiTa $_3$ O $_8$  or Li $_2$ Ta $_6$ O $_{16}$  is apparently isostructural with LiTa $_6$ O $_{15}$ F and Ta $_4$ W $_2$ O $_{16}$ . Either fluorine or tungsten ions can be utilized to generate a non-stoichiometric phase and the structural mechanism is thus alkali ion vacancies plus cation (or anion) substitution.

The low temperature form of Ta<sub>2</sub>O<sub>5</sub> (L-Ta<sub>2</sub>O<sub>5</sub>) is stabilized by the addition of Li<sub>2</sub>O. This structure is apparently intrinsically non-stoichiometric with the formula Ta<sub>22</sub>O<sub>55</sub> and five oxygen "vacancies" [25]. The Li<sup>+</sup> ions either substitute for Ta<sup>+5</sup> creating more anion vacancies or more likely occur interstitially filling some of the anion "vacancies" already present or a mechanism of cation substitution plus anion interstitials.

It should be remembered that all of the mechanisms discussed in this section are hypothetical and must be checked by careful single crystal x-ray diffraction or even neutron diffraction studies. Until such studies can be made, our understanding of the structural mechanisms by which nature compensates a non-stoichiometric phase for electrical neutrality will remain poorly understood.

### SUMMARY OF RESULTS

1. The phase equilibria of various alkali tantalates have been studied up to and including liquidus temperatures and phase equilibrium diagrams have been constructed most consistent with the experimental data in the systems:

$${^{{\rm Ta}_2}{^{\rm O}_5}}^{-{\rm LiTaO}_3}$$
 ${^{{\rm Ta}_2}{^{\rm O}_5}}^{-{\rm NaTaO}_3}$ 
 ${^{{\rm Ta}_2}{^{\rm O}_5}}^{-{\rm KTaO}_3}$ 

2. The phase equilibria of various alkali niobates have been studied up to and including liquidus temperatures and phase equilibrium diagrams have been constructed most consistent with the experimental data in the systems:

$${\rm Nb_2^O_5}$$
-LiNbO $_3$ 
 ${\rm Nb_2^O_5}$ -NaNbO $_3$ 
 ${\rm Nb_2^O_5}$ -KNbO $_3$ 

3. Pellets of nineteen phases were transmitted to the sponsoring agency for evaluation and the following three compositions were found to show peaks in the dielectric loss vs. temperature, indicating some ionic mobility.

```
11K_2O:89Ta_2O_5 - Gatehouse tungsten bronze structure (GTB) 21.75K_2O:78.25Ta_2O_5 - Hexagonal tungsten bronze structure (HTB) 34K_2O:66Ta_2O_5 - Tetragonal tungsten bronze structure (TTB)
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4. Attempts were made to grow large crystals of eight different phases and two crystals were transmitted to the sponsor for evaluation.

#### FUTURE WORK

- 1. The most interesting phases in the K<sub>2</sub>O-Ta<sub>2</sub>O<sub>5</sub> system (GTB and HTB) did not have phase equilibria relations favorable for crystal growth from the melt. Preliminary examination of the Rb<sub>2</sub>O-Nb<sub>2</sub>O<sub>5</sub> and Rb<sub>2</sub>O-Ta<sub>2</sub>O<sub>5</sub> systems suggests that these systems may be more favorable for melt growth of the GTB and HTB type phases. The phase equilibria in these systems will be examined in more detail and attempts made to grow crystals which might then be ion exchanged.
- 2. Small crystals of the hexagonal tungsten bronze phase in the system KTaO<sub>3</sub>-WO<sub>3</sub> have been successfully grown from a potassium tungsten oxide flux. Attempts will be made to grow larger single crystals of this phase for dielectric loss and/or ionic conductivity measurements.
- 3. A cubic phase of KSbO<sub>3</sub>, similar to the high pressure polymorph previously demonstrated to exhibit ionic mobility, has been successfully synthesized by the addition of a small amount of either SiO<sub>2</sub> or Al<sub>2</sub>O<sub>3</sub>. The phase equilibria of the Sb<sub>2</sub>O<sub>4</sub>-NaSbO<sub>3</sub> and Sb<sub>2</sub>O<sub>4</sub>-KSbO<sub>3</sub> systems will be studied and the effects will be studied of various impurities on the stabilization of the cubic modifications of KSbO<sub>3</sub> and NaSbO<sub>3</sub>. Attempts will be made to prepare single crystals of these cubic phases and/or other interesting alkali antimonates.

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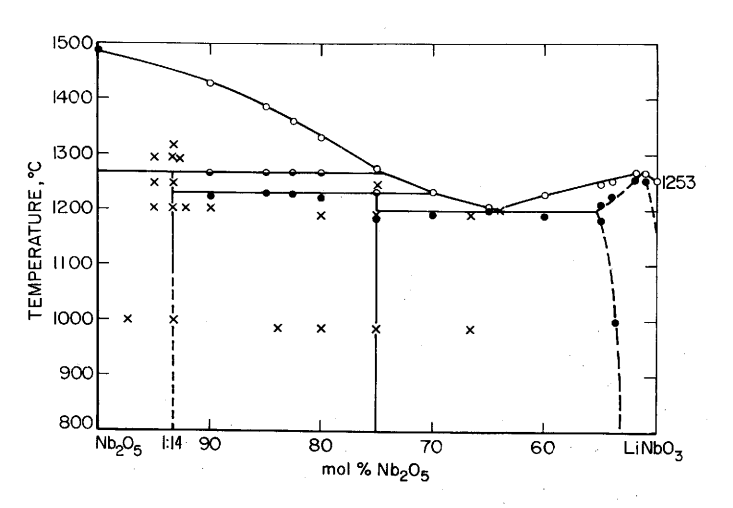
### FIGURE CAPTIONS

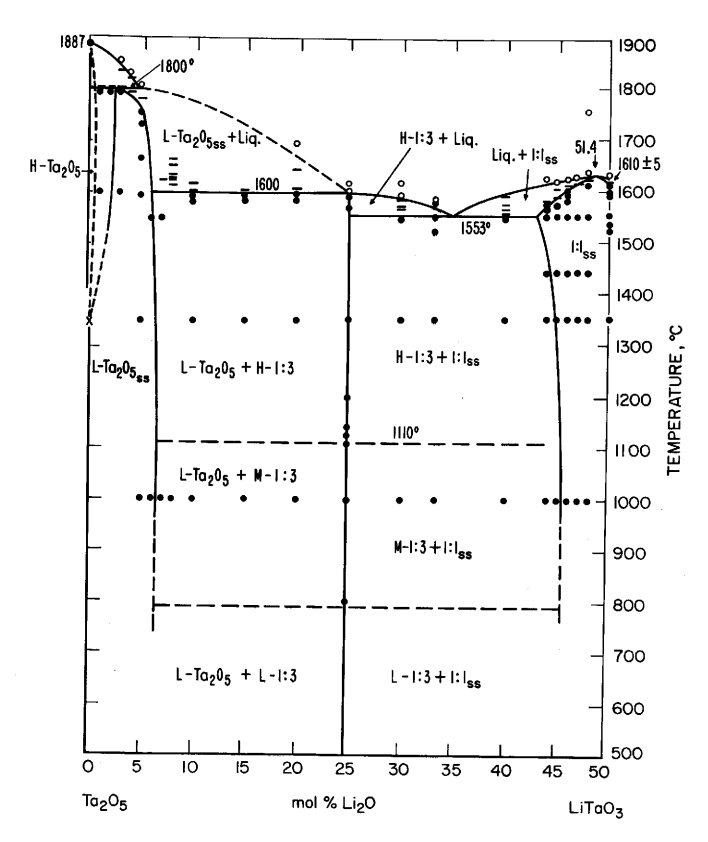
- Figure 1 Phase equilibrium diagram for the system  ${\rm Nb_2O_5}{\rm -LiNbO_3}$  redrawn to conform with accepted published data.
  - o liquidus [7]
  - e transition [7]
  - - solidus and solid solution boundary [7,11]
  - X present work
- Figure 2 Phase equilibrium diagram of the system Ta<sub>2</sub>O<sub>5</sub>-LiTaO<sub>3</sub>, mostly from reference [2].
  - o completely melted
  - - partially melted
  - - no melting
- Figure 3 Phase equilibrium diagram of the system Nb<sub>2</sub>O<sub>5</sub>-NaNbO<sub>3</sub>
  - X liquidus values from reference [4]
  - o completely melted
  - e partially melted
  - no melting

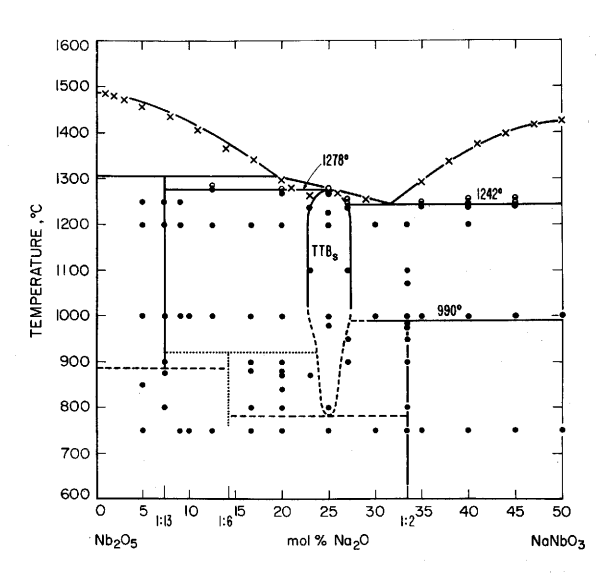
dotted line - a phase resulting from reaction with atmospheric moisture

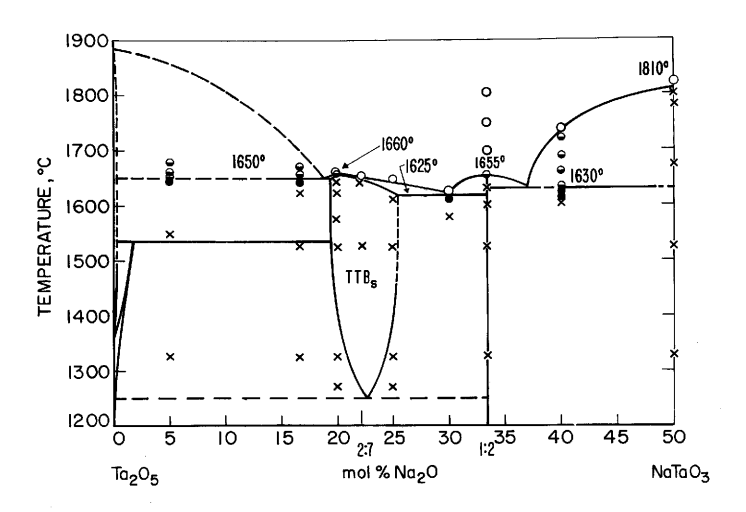
- Figure 4 Phase equilibrium diagram for the system  $Ta_2O_5$ -NaTaO3
  - o completely melted
  - partially melted
  - no melting
  - X quenched specimen
- Figure 5 Phase equilibrium diagram for the system Nb<sub>2</sub>O<sub>5</sub>-KNbO<sub>3</sub>
  - X liquidus values from reference [20]
  - o completely melted
  - e partially melted
  - no melting

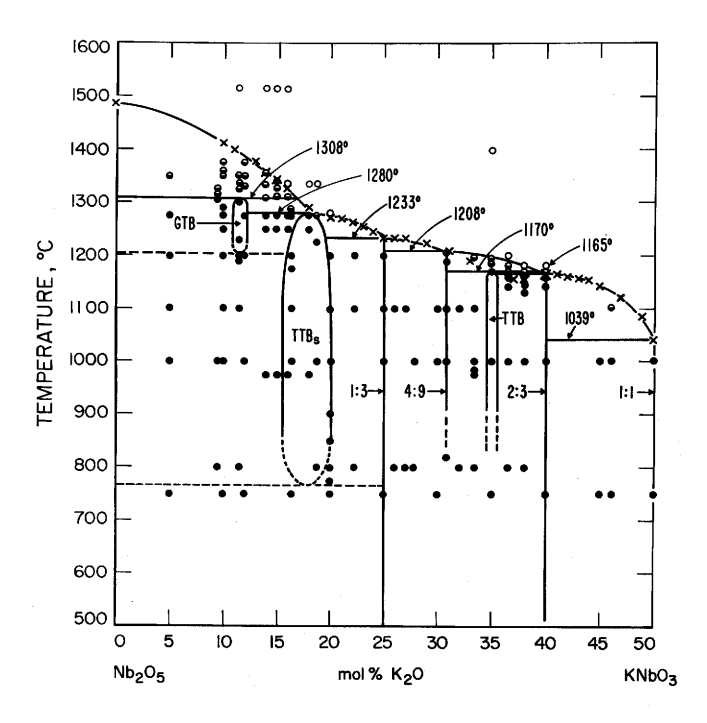
- Figure 6 Phase equilibrium diagram for the system  ${\rm Ta_2O_5}$ -KTaO3
  - solidus and liquidus values from reference [25]
  - o completely melted
  - partially melted
  - X no melting
- Figure 7 Dielectric loss (epsilon-2) versus reciprocal temperature (theta =  $10^3$  x degrees Kelvin<sup>-1</sup>) for a specimen of  $11K_2^0:89Ta_2^0_5$  (sample No. 27) having the structure of the Gatehouse Tungsten Bronze (GTB).
- Figure 8 Dielectric loss (epsilon-2) versus reciprocal temperature (theta =  $10^3$  x degrees Kelvin<sup>-1</sup>) for a specimen of  $16.67 \text{K}_2\text{O}:83.33 \text{Ta}_2\text{O}_5$  (sample No. 23) having the structure of a Tetragonal Tungsten Bronze modified by superstructure to a tripled cell with orthorhombic (TTBs).
- Figure 9 Dielectric loss (epsilon-2) versus reciprocal temperature (theta =  $10^3$  x degrees Kelvin<sup>-1</sup>) for a specimen of  $21.75K_20:78.25Ta_2O_5$  (sample No. 39) having the structure of a Hexagonal Tungsten Bronze (HTB).
- Figure 10 Dielectric loss (epsilon-2) versus reciprocal temperature (theta =  $10^3$  x degrees Kelvin<sup>-1</sup>) for a specimen of  $34K_2O:66Ta_2O_5$  (sample No. 28) having the structure of a Tetragonal Tungsten Bronze with no superstructure (TTB).

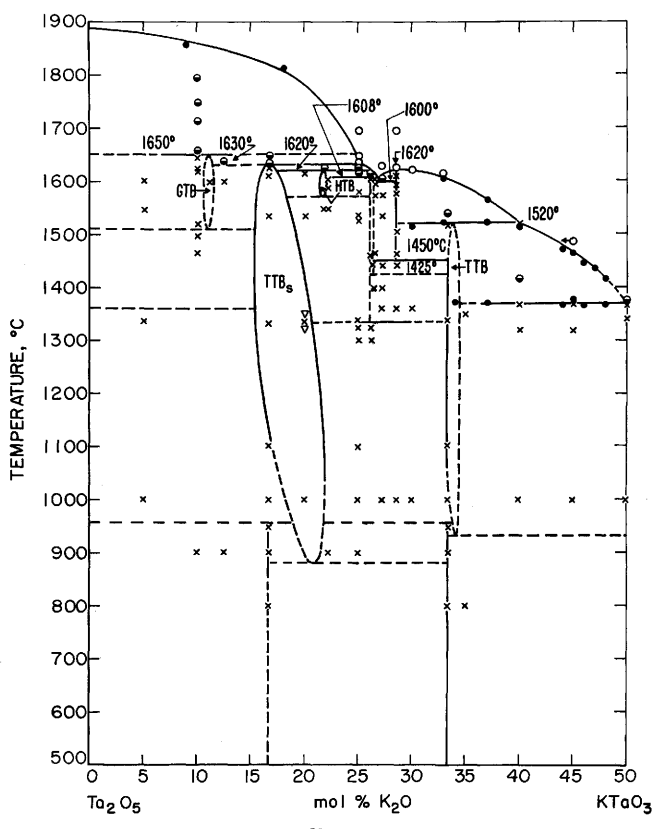


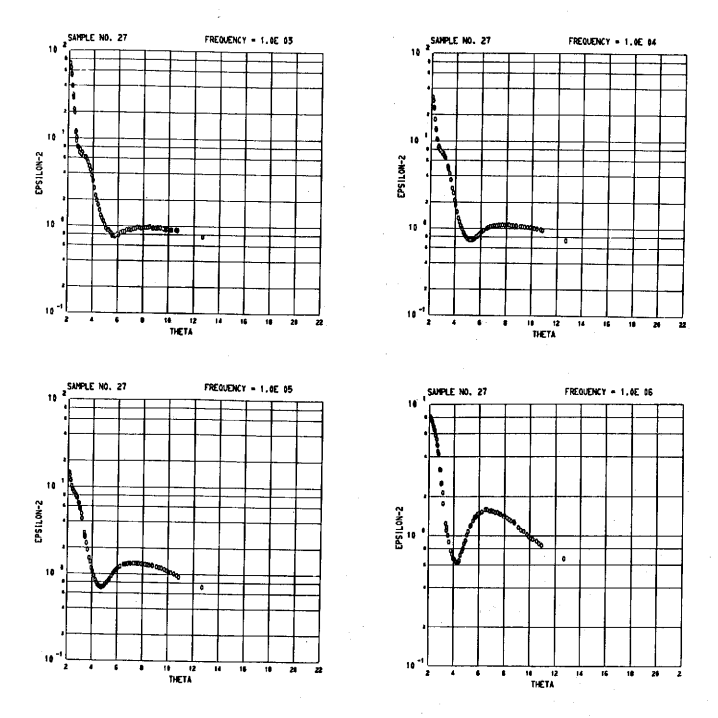




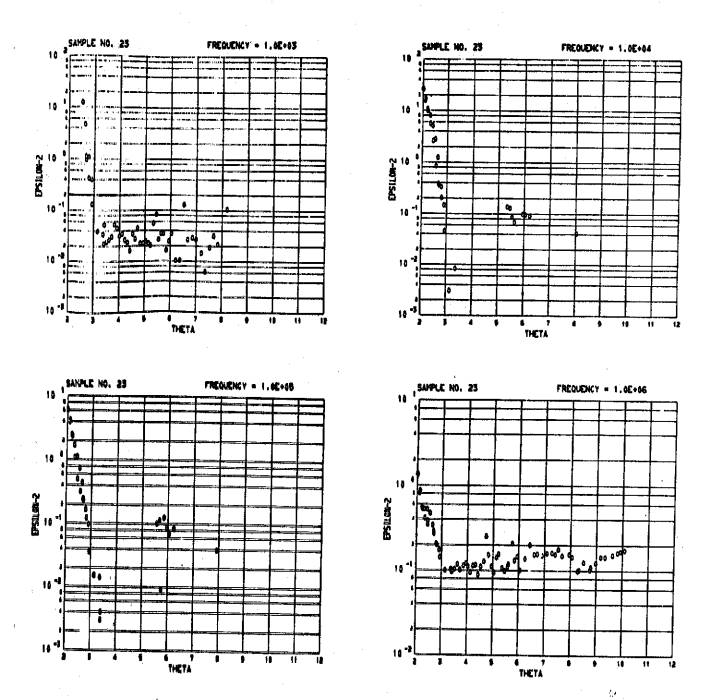




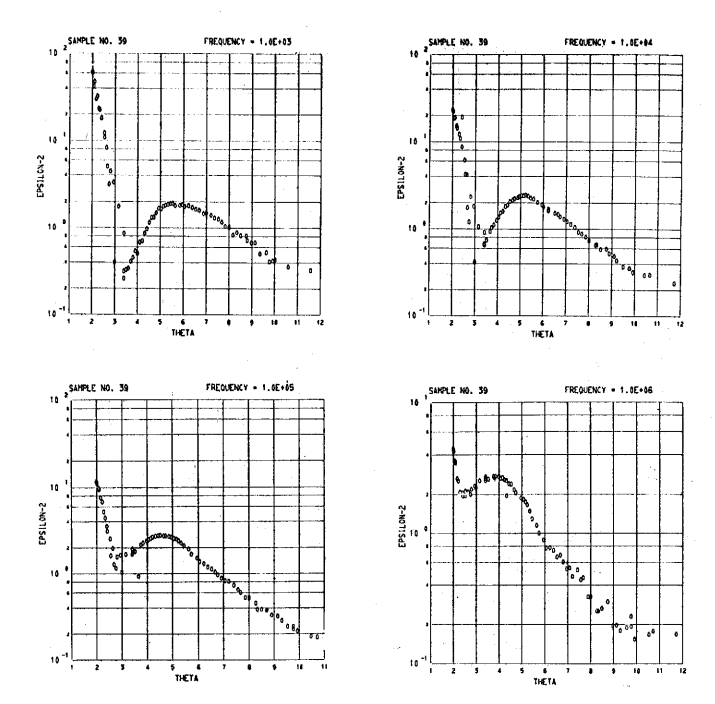




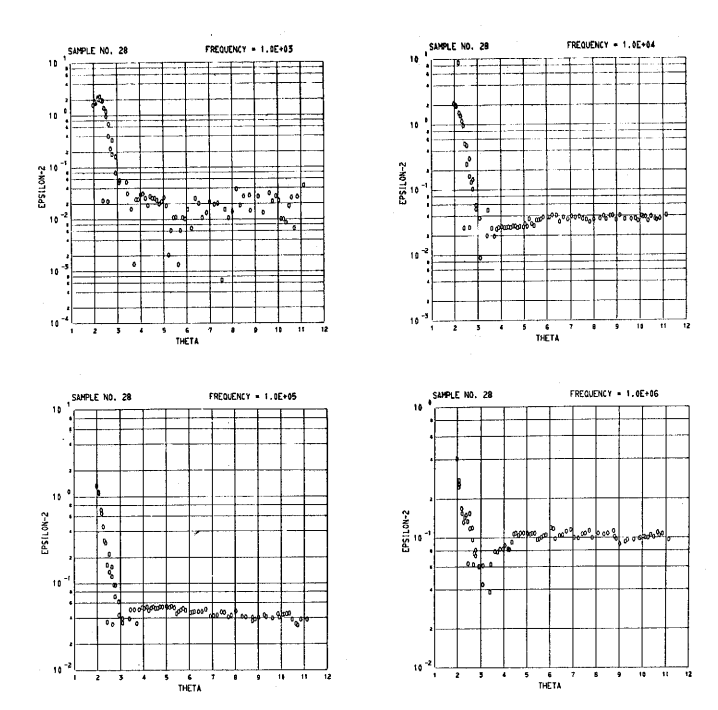
DIELECTRIC LOSS (EPSILON-2) VERSUS RECIPROCAL TEMPERATURE (THETA =  $10^3$  x degrees Kelvin<sup>-1</sup>) for a specimen of  $11K_20:89Ta_20_5$  (sample No. 27) maving the structure of the Gatehouse Tungsten Bronze (GTB).



DIELECTRIC LOSS (EPSILON-2) VERSUS RECIPROCAL TEMPERATURE (THETA =  $10^3$  x degrees Kelvin  $^1$ ) for a specimen of  $16.67 K_2 0.63.33 Ta_2 0_5$  (Sample No. 23) having the structure of a Tetragonal Tungsten Bronze Modified by superstructure to a tripled cell with Orthorhombic symmetry (TTBs)



Bielectric loss (epsilon-2) versus reciprocal temperature (theta =  $10^3$  x degrees Kelvin<sup>-1</sup>) for a specimen of  $21.75K_28:78.25Ta_20_5$  (sample No. 39) maying the structure of a Hexagonal Tungsten Bronze (HTB).



Dielectric loss (epsilon-2) versus reciprocal temperature (theta =  $10^3$  x degrees Kelvin<sup>-1</sup>) for a specimen of  $34K_20:66Ta_20_5$  (sample No. 28) having the structure of a Tetragonal Tungsten Bronze with no superstructure (TTB)

TABLE 1: EXPERIMENTAL DATA FOR THE SYSTEM Nb205-Linbo3

Composition		Initia	l Heat		Heat ment ⊈/	Results of Physical Observation	Results of X-Ray Diffraction Analyses 2					
Mole 4 ≛/		Temp.	Time Hrs.	Temp,	Tima Hre.	COMESVACION	vueri\heb					
Nb205	Li 20	•		-								
97,50	2.50	1000	96				$H-Mb_2O_5 + 1;3 + N-Nb_2O_5$ (tr)					
95,00	5,00	1000	63									
				1202	20	<b></b>	N-Mb2O5+ H-Mb2O5					
				1249	19	Not melted						
				1274	24	Not visibly melted						
				1294	22	Not visibly melted	N-Nb2O5 + H-Nb2O5					
				1318	24	Not visibly melted						
				1336	23	Some liquid present	<del></del>					
93,33	6,67	1000	63				H-Mb <sub>2</sub> O <sub>5</sub> + 1:3					
				1202	20		N-Nb2O5					
		·		1249	19	Not melted	N-Nb2O5					
				1274	24	Not melted						
				1294	22	Not visibly melted	N-Nb205 4/					
				1318	24	Partially melted	$H-Nb_2O_5 + 1:3 + N-Nb_2O_5$					
				1.336	23	Partially melted						
92.86	7.14	1000	60			#==						
				1294	22	Not melted	N-Nb <sub>2</sub> O <sub>5</sub> + H-Nb <sub>2</sub> O <sub>5</sub> + 1:3					
92.31	7.69	1000	63									
				1202	20	Not melted	$N-Nb_2O_5 + 1:3$					
				1248	18.75	Partially melted						
90.00	10.00	1000	63				****					
				1202	20	Not melted	N-Mb <sub>2</sub> O <sub>5</sub> + 1:3					
83.33	16.67	985	64			Not melted						
				1244	20	Partially melted						
80.00	20,00	985	65			Not melted	1:3 + H-ND <sub>2</sub> O <sub>5</sub>					
				1190	96	Not melted	1:3 + N-Nb <sub>2</sub> 0 <sub>5</sub>					
				1224	25	Not melted	7-7-5					
75.00	25.00	985	65			Not melted	1:3					
				1190	96	Not melted	1:3					
				1224	25	Not melted						
				1244	20	Partially melted	$N-Nb_2O_5 + 1:3 + 1:1$					
70.00	30.00	1000	70.5			Not melted						
				1183	23	Not melted						
				1195	24	No visible melting						
				1203	25	Some melting						
				1207	23	Some melting						
				1215	24	Some melting	WI 60 50 60 mm					
66,67	33.33	985	65		<del></del>		1:1 + 1:3					
				1190	96	Not melted	1:1 + 1:3					
				1224	25	Melted						
63.99	36.01	1000	70.5				:					
				1183	23	Not melted .						
				1195	24	Partially melted	=++					
				1203	25	Melted	$1_{1}1 + 1_{1}3$					
				1207	23	Melted	*****					

 $<sup>\</sup>underline{\mathbf{a}}'$  For ease and accuracy of weighing Li<sub>2</sub>O was added to Nb<sub>2</sub>O<sub>5</sub> as LiNbO<sub>3</sub> not as the oxide end member.

 $b^\prime$  All specimens were initially calcined in Pt crucibles at the indicated temperature and

 $<sup>\</sup>underline{c}/$  All subsequent heat treatments were quenched in sealed Pt tubes from the indicated

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

H-Nb<sub>2</sub>O<sub>5</sub> - the high temperature form of Nb<sub>2</sub>O<sub>5</sub>

N-Nb<sub>2</sub>O<sub>5</sub> - a metastable form of Nb<sub>2</sub>O<sub>5</sub> apparently stabilized by Li<sub>2</sub>O<sub>5</sub>

1:3 - LiNb<sub>3</sub>O<sub>8</sub>

1:1 - LiNb<sub>3</sub>O<sub>8</sub> solid solution

 $<sup>^{\</sup>underline{e}\prime}$  These experiments suggest that the N-Nb<sub>2</sub>O<sub>5</sub> phase melts incongruently between 1294° and 1318° rather than at the 1268° value given by Reisman.

TABLE 2: EXPERIMENTAL DATA FOR THE SYSTEM Ta205-LITAO3

				4			
Composition		Initia Treat	l Heat ment b/		L Heat	Results of Physical Observation	Results of X-Ray Diffraction Analyses
Mo	le 🕽 🌁	Temp.	Time	Temp.	Time		
Ta <sub>2</sub> O <sub>5</sub>	Li <sub>2</sub> 0	*¢	Hrs.	*C	Hrs.		•
14205	4420			4,5	6		
<b>5</b> 9	1	1000	60				
				1600	19.00	No melting	H-Ta <sub>2</sub> O <sub>5</sub> + L-Ta <sub>2</sub> O <sub>5</sub> ss
				1795	0.17	No melting	м 11
				1805	0.17	Partially melted	
98	2	1000	60				
				1600	24.00	No melting	L-Ta205 ss + H-Ta205
				1795	0.17	No melting	er H
				1806	0,25	Partially melted	и н
97	3	1000	60			*****	
		<del>-</del>		1600	24.00	No melting	L-Ta2O5 ss
				1795	0.17	No melting	
				1803	0.17	Partially melted	
				1842	0.03	Partially melted	
				1855	0.03	Completely melted	
96	4	1000	60				
				1794	0.17	Partially melted	
				1820	0.33	Partially melted	
				1834	0.33	Completely melted	
95	5	1000	10				
	_	1000	10	1350	24.00	No melting	ITa_O_ SS
				1594	16.00	No melting	L-Ta <sub>2</sub> O <sub>5</sub> ss
				1597	0.50	No melting	***
				1667	0.17	No melting	
				1732 1757	0.17 0.17	No melting No melting	
				1782	0.17	Partially melted	I-Ta O ee
				1809	0.17	Completely melted	L-Ta <sub>2</sub> O <sub>5</sub> ss
	_					- <del>-</del>	
94	6	1000	10				L-Ta <sub>2</sub> O <sub>5</sub> ss
				1550	16.00	No melting	
93	7	1000	10			<u>-</u>	L-Ta <sub>2</sub> O <sub>5</sub> ss + M-1:3
				1550	24.00	No melting	1 14702 22 . W.I.I.
				1625	16.00	Partially melted	L-Ta <sub>2</sub> O <sub>5</sub> ss
92	8	1000	10				
	•	1000	10	1615	1.00	Partially melted	<b>*</b>
				1625	0.75	Partially melted	
				1627	0.75	Partially melted	
				1635	0.50	Partially melted	
				1653	0.50	Partially melted	
				1663	1.00	Partially melted	L-Ta <sub>2</sub> O <sub>5</sub> ss + H-1:3
90	10	1000	10				
				1350	24.00	No melting	L-Ta <sub>2</sub> O <sub>5</sub> ss + H-1:3
		•		1583	0.50	No melting	
				1590	16.00	No melting	$L-Ta_2O_5$ ss + H-1:3
				1593 1607	0.50 0.75	No melting Partially melted	F-Ma-O- no ( V 3-2 ) 1 2
	•			1618	0.50	Partially melted	L-Ta <sub>2</sub> O <sub>5</sub> ss + H-1:3 + 1:1
85	15	1000	10				
	•			1350	24.00	No melting	
				1583 1593	0.50 0.75	No melting No melting	
				1605	0.50	Partially melted	
80	20	1000	10				M-1:3 + L-Ta <sub>2</sub> O <sub>5</sub> ss
				1350	24.00	No melting	H-1:3 + L-Ta <sub>2</sub> O <sub>5</sub> ss
				1580 1595	1.00 1.00	No melting	W III
				1609	0.50	No melting Partially melted	
				1642	0.50	Partially melted	
				1646	0.50	Partially melted	<del></del> ·
				1695	0.75	Probably completely	
						melted	
					35		

```
75
          25
                            307
                    800
                                                                               L=1:3 + M=1:3
                   1000
                             10
                                                                               M-1:3
                                    1077
                                           312.00
                                                      No melting
                                                                               H
                                    1110
                                            42.00
                                                      No melting
                                    1111
                                           307,00
                                                      No melting
                                    1126
                                            96,00
                                                      No melting
                                    1130
                                            16.00
                                                      No melting
                                    1143
                                            16.00
                                                      No melting
                                    1144
                                           115,00
                                                      No malting
                                    1205
                                            18.00
                                                      No melting
                                    1265
                                            18,00
                                                      No melting
                                    1350
                                             24.00
                                                      No melting
                                    1573
                                            16.00
                                                      No melting
                                                                                ----
                                    1592
                                             1.00
                                                      No melting
                                    1601
                                             1.00
                                                      Completely melted
                                    1616
                                             1.50
                                                      Completely melted
                                                                              R-1:3 + L-Ta_2O_5 ss + 1:1
                   1000
                             10
                                     --
                                             --
                                                          ----
                                                                                ----
                   1205
                             18
                                    1130
                                            16,00
                                                      No melting
                                                                                 ----
                                                      No melting
                                                                                 ----
                                    1143
                                            16.00
                                                                                 ----
                                    1144
                                           115.00
                                                      No melting
                   1000
                                                                                ----
                             10
                                     --
                                             --
                                                          ----
                                                           ----
                   1250
                             30
                                                                              H-1:3
                                    1077
                                           312.00
                                                           ____
                                                                              H = 1 : 3
                                    1111
                                           307.00
                                                                                ____
                                            96.00
70
                   1000
                             10
                                                           ____
                                                                                 ----
                                    1350
                                            24.00
                                                      No melting
                                                                                 ----
                                    1545
                                             0.50
                                                      No melting
                                    1555
                                             0.50
                                                      Partially melted
                                                                                 ----
                                    1566
                                             1.00
                                                      Partially melted
                                                                                 ----
                                    1572
                                             0.75
                                                      Partially melted
                                                                                 ----
                                    1587
                                                                                ----
                                                      Partially melted
                                    1594
                                             0.25
                                                     Probably completely melted
                                    1616
                                             0.75
                                    1616
                                             0.50
                                                                              H-1:3 + 1:1 ss
66.67
          33.33
                  1000
                             10
                                                                              M-1:3 + 1:1 ss
                                    1350
                                            24.00
                                                     No melting
                                                                              H-1:3 + 1:1 as
                                    1525
                                             2,50
                                                     No melting
                                                                                ____
                                    1551
                                             2.50
                                                      No melting
                                    1555
                                             0.50
                                                     Probably some melting
                                    1567
                                             1.00
                                                      Considerably melted
                                                                                ----
                                    1578
                                                                                -----
                                             2.00
                                                     Completely melted
                                    1581
                                             0.25
                                                     Completely melted
                                    1583
                                                     Completely melted
                                             3,00
                                                                              H-1:3 + 1:1 ss
60
          40
                  1000
                          . 10
                                    1350
                                            24.00
                                                     No melting
                                                                                ----
                                                     No melting
                                                                                ----
                                    1547
                                             0.50
                                    1555
                                             0.50
                                                     Partly melted
                                    1566
                                             0.50
                                                     Partly melted
                                                                                -----
                                    1572
                                             0.50
                                                                                ----
                                                     Partly melted
                                    1590
                                             1.00
                                                     Considerably melted
56
                  1000
                            10
          44
                                    1348
                                            66.00
                                                     No melting
                                                                              1:1 ss + H-1:3
                                    1440
                                            70.00
                                                     No melting
                                    1550
                                             0.75
                                                     No melting
                                                                                ----
                                    1565
                                             0.50
                                                     No melting
                                                                                ----
                                                     No melting
                                    1576
                                             0.50
                                    1581
                                             0.50
                                                     Partially melted
                                    1622
                                             0.50
                                                     Completely melted
55
          45
                  1000
                                                                                ----
                                    1348
                                            66.0
                                                     No melting
                                                                              1:1 ss
                                    1350
                                            24.00
                                                     No melting
                                                                              1:1 \text{ ss} + H-1:3 (tr)
                                                     No melting
                                    1440
                                            70.00
                                                                              l:l ss
                                    1550
                                             1.00
                                                     No melting
                                    1572
                                             0.75
                                                     No melting
                                                                              1:1 ss
                                    1587
                                             0.50
                                                     Partially melted
                                                                                ---
                                    1606
                                             0.50
                                                     Partially melted
                                    1617
                                             0.75
                                                     Completely melted
54
          46
                  1000
                            10
                                   1348
                                            66.00
                                                     No melting
                                                                              1:1 88
                                   1440
                                            70,00
                                                     No melting
                                   1550
                                            1,00
                                                     No melting
                                    1581
                                             0.50
                                                     No melting
                                                                                ----
                                    1592
                                             0.50
                                                     No melting
                                                                                ----
                                   1605
                                             0.50
                                                     No melting
                                   1612
                                             0,50
                                                     Partially melted
                                   1624
                                             0.50
                                                     Completely melted
```

53	47	1000	10					
		-,		1348	66.00	No melting		l:l ss
				1440	70.00	No melting		10
				1550	1.00	No melting		
				1612	0.50	Just begun	to melt	
				1623	0.50	Completely	melted	
52	48	1000	10		·			1:1 ss
- •	_			1348	66.00	No melting		*1
				1440	70.00	No melting		ri
				1550	1.00	No melting		
		•		1612	0.75	No melting		
				1623	0.50	Just begun	to melt	
				1637	0.33	Completely	melted	<b></b>
				1754	0.17	Completely	melted	
50	50 S	Starting ma	terial	***		Powder		1:1
		_		1348	19.00	No melting		
				1522	16.00	No melting		"
				1538	2.50	No melting		"
				1553	20.00	No melting		
				1592	1.00	No melting		
				1598	1.75	No melting		
				1607	1.00	No melting		
				1612	1.00	No melting		
				1616	1.00	Partially r	nelted	
				1630	0.50	Completely		
				1633	1.00	Completely	melted	

 $<sup>\</sup>frac{a}{2}$  For ease and accuracy of weighing,  $\text{Li}_2\text{O}$  was added to  $\text{Ta}_2\text{O}_5$  as  $\text{LiTaO}_3$  not as the oxide end member.

b/ All specimens were initially calcined in Pt crucibles at the indicated temperatures and time.

Specimens were heated in both open and sealed Pt tubes and seemed to show no difference or discrepancy in results, as very little, if any, volatilization takes place even from the melt.

 $<sup>\</sup>frac{d}{d}$  All phases identified are given in order of amount present at room temperature (greatest amount first). The phases are not necessarily those present at the temperature to which the specimen was heated. H-Ta $_2$ O $_5$  - The high temperature polymorph of Ta $_2$ O $_5$ .

L-Ta<sub>2</sub>O<sub>5</sub> = The high temperature polymorph of Ta<sub>2</sub>O<sub>5</sub>.

L-Ta<sub>2</sub>O<sub>5</sub> ss - A solid solution of the low temperature polymorph of Ta<sub>2</sub>O<sub>5</sub> stabilized by Li<sub>2</sub>O.

L-1:3 - The low temperature polymorph of LiTa<sub>3</sub>O<sub>8</sub> isostructural with LiNb<sub>3</sub>O<sub>8</sub>.

M-1:3 - The medium temperature polymorph of LiTa<sub>3</sub>O<sub>8</sub> isostructural with the mineral woodgenite.

H-1:3 - The high temperature polymorph of LiTa<sub>3</sub>O<sub>8</sub> isostructural with LiTa<sub>6</sub>O<sub>15</sub>F and Ta<sub>4</sub>W<sub>2</sub>O<sub>16</sub>.

<sup>1:1</sup> ss - A solid solution of LiTaO3.

TABLE 3: EXPERIMENTAL DATA FOR THE SYSTEM Nb2O5-NANDO3

Compos	ition	Initia: Tréats	l Heat ment b/	Final Treat	Heat ment ©/	Results of Physical Observation	Results of X-Ray Diffraction Analyses d
Mole	• •	Temp.	Time Hrs.	Temp. °C	Time Hrs.		
Nb <sub>2</sub> O <sub>5</sub>	Na <sub>2</sub> O						
95	5	750	69				H-Nb <sub>2</sub> O <sub>5</sub> + 1:2
		1000	69			Not melted	1:13 + H-Nb <sub>2</sub> O <sub>5</sub>
				1200	21	Not melted	11
				1250	21	Not melted	
92.86	7.14	800	69				H-Nb <sub>2</sub> O <sub>5</sub> + 1:6
				875	19.5	Not melted	ă
				900	140	Not melted	$1:13 + H-Nb_2O_5 + 1:6$
				1200	21	Not melted	1:13
		1000	164	1250	67	Not melted Not melted	u
		1225	64	1250		Not melted	
			•	1000	234	Not malted	1:13
						•	
90.01	9.09	750	69				H-Nb <sub>2</sub> O <sub>5</sub> + 1:2
		1000	69	1200	22	Not melted Not melted	1:13 + TTBs + H-Nb <sub>2</sub> O <sub>5</sub> 1:13 + TTBs
				1250	21	Not melted	11
90	10	750	64				H-Nb <sub>2</sub> O <sub>5</sub> + 1:2
		1000	69			Not melted	1:13 + TTBs
87.5	12.5	750	69				U-mb () + 1.2
0,.5	14.5	1000	69			Not melted	H-Nb <sub>2</sub> O <sub>5</sub> + 1:2 1:13 + TTBs
				1200	22	Not melted	
				1278	72	Partially melted	•
				1285	1	Partially melted	it.
		800	114			·	
		1225	8			Not melted	1:13 + TTBs
83.33	16.67	750	64				H-Nb <sub>2</sub> O <sub>5</sub> + 1:2
				800	113	Not melted	1:6 + H-Nb,Og + TTBs
				880	20	Not melted	TTBs + H-Nb <sub>2</sub> O <sub>5</sub> + 1:6
		1000	69	900	18,5	Not melted	1:13 + TTBs
•		1000	02	1200	22	Not melted	1:13 * 1108
		,					
80	20	750	64				H-Nb <sub>2</sub> O <sub>5</sub> + 1:2
				800	113	Not melted	1:6 + H-Nb <sub>2</sub> O <sub>5</sub> + TTBs
				840 869	70 19	Not melted Not melted	TTBs + 1:6 + H-Nb <sub>2</sub> O <sub>5</sub>
				880	20	Not melted	10
				900	18.5	Not melted	
			•	1000	67	Not melted	TTBs + 1:13
		1000	69				н
				1200	22	Not melted	•
				1270 1275	1 1.5	Not melted Not melted	
	•			1278	2.5	Partially melted	
77	23	870	91			safe Chi mai dan ma	TTBs + $H-Nb_2O_5 + 1:2$
		1100	21			·	TTBs + 1:13
		1100	44	 1238	19		TTBs "
				1236	1.7		
75	25	750	64				H-Nb2O5 + 1:2
		,		800	113	Not melted	1:6 + TTBs + 1:2
				980	18.5	Not melted	TTBs "
		1000	69	1000	20	Not melted Not melted	н
			J.,	1200	21	Not melted	u
				1270	1	Not melted	
				1275	1.5	Not melted	
				1278	2,5	Completely melted	
		1000		1280	1	Completely melted	
		1200	21	800	115	Not melted	T <b>TB</b> s
		900 1000	62 96			Not melted	TTBs
			,,,	1225	1	Not melted	111111
				1225	5	Not melted	u

72.92	27.08	900 900	19 135				TTBs + 1:2 + H-Nb <sub>2</sub> O <sub>5</sub> TTBs + 1:2
		300	, 133	950	91		4
		1100	44			Not melted	TTBs (+ 1:1 ?) 💇
				1238	19	Not melted	
				1249	3.5	Partially melted	~~~ <del>~</del>
				1255	16.5	Partially melted	
70	30	750	64	-			1:2 + 1:1 + H-Nb <sub>2</sub> O <sub>5</sub> ,
		1000	69			Not melted	1:2 + TTBs (+1:1) #/
				1000	20	Not melted	/م
				1200	22	Not melted	TTBs (+1:1 ?) $e/$
66.67	33,33	800	62			an ex ma ex ex	
		900	93			Not melted	
				950	93	Not melted	1:2
		800	62				ade was per hije ger
		1100	62			Not melted	TTBs + 1:1
				975	21	Not melted	1:2
				985	22	Not melted	1:2 (+TTBs + 1:1) trace
		800	89				1:2
				985	45	Not melted	1:2 (+1:1 + TTBs) trace
				1000	169	Not melted	TTBs + 1:1
				1073	67	Not melted	"
				1200	21	Not melted	"
65	35	750	64				$1:2 + 1:1 + H-Nb_2O_5$
	•			1241	19	Not melted	<del></del>
				1254	19	Partially melted	
		1000	69			Not melted	1:2 + 1:1
				1250	21	Partially melted	TTBs + 1:1 (?)
60	40	750	64			<del></del>	1:2 + 1:1
				1241	19	Not melted	1:1 + TTBs
				1245	65	Partially melted	
				1254	19	Partially melted	
		1000	69			Not melted	1:2 + 1:1 (+TTBs) e/
	,			1200	21	Not melted	1:1 + TTBs
				1250	21	Partially melted	n
55	45	750	64				1:1 + 1:2
				1241	19	Not melted	1:1 + TTBs
				1245	<b>6</b> 5	Partially melted	
				1254	19	Partially melted	<del></del>
		1000	69			Not melted	1:1 + 1:2
50	. 50	750	64				1:1
•		1000	69			Not melted	1:1

 $<sup>\</sup>frac{a}{}$  For ease and accuracy of weighing Na<sub>2</sub>O was added to Nb<sub>2</sub>O<sub>5</sub> as NaNbO<sub>3</sub> not as the oxide end member.

b/ All specimens were initially calcined in Pt crucibles at the indicated temperatures and time.

c/ All subsequent heat treatments were quenched in sealed Pt tubes from the indicated temperature.

 $<sup>\</sup>underline{d}'$  All phases identified are given in order of amount present at room temperature (greatest amount first). The phases are not necessarily those present at the temperature to which the specimen was heated. H-Nb<sub>2</sub>O<sub>5</sub> - The high temperature form of Nb<sub>2</sub>O<sub>5</sub>

<sup>1:13 -</sup> NaNb<sub>13</sub>O<sub>33</sub>

<sup>1:6 -</sup> NaNb<sub>6</sub>O<sub>15</sub>(OH). The presence of this phase indicates that the specimen has reacted with atmospheric moisture.

TTBs - A nonstoichiometric solid solution having an orthorhombic distortion of a tetragonal tungsten bronze-type lattice with superstructure indicating a tripled unit cell.

 $<sup>1:2 -</sup> Na_2Nb_4O_{11}$ 

<sup>1:1 -</sup> NaNbO3

e/ The presence of a small amount of either 1:1 or TTBs in equilibrium with a large amount of the other cannot be determined because of a complete overlap of all of the strongest peaks.

TABLE 4: EXPERIMENTAL DATA FOR THE SYSTEM Ta205-Nata03

Composition		Initia		Final	L Heat	Results of Physical	Results of X-Ray Diffraction
Mole % a/		Temp.	ment 🖭		ment <sup>C</sup> / Time	Observation	Analyses 🚭
MOTE	·	°C	Time Hrs.	Temp.	Hrs.		
<b>Ta</b> <sub>2</sub> O <sub>5</sub>	Na <sub>2</sub> O	J		Ū	112.0		
95	5	1000	109				L-Ta <sub>2</sub> O <sub>5</sub> + 1:2
				1328	16.00	Not melted	<del></del> -
				1549	16.00	Not melted	H-Ta <sub>2</sub> O <sub>5</sub> (tri.) + TTBs
				1648	0,08	Not melted	
				1653	0.08	Partially melted	
				1664	0.08	Partially melted	# 17 m m =
				1679	0.08	Partially melted	
83.33	16.67	1000	109			 Watlt-d	
				1328	16.00	Not melted	TTBs + L-Ta <sub>2</sub> O <sub>5</sub>
				1527	16.00	Not melted	TTBs + L-Ta <sub>2</sub> O <sub>5</sub> ss
				1625	41.00	Not melted	TTBs + H-Ta2O5
	•			1643 1658	0.08 0.08	Not melted Partially melted	<del></del>
				1670	0.08	Partially melted	
00	20	1000	100			_	10.700
80	20	1000	109	 1195	552.00	No melting	1:2 + L-Ta <sub>2</sub> O <sub>5</sub>
				1270	360.00	No melting	TTBs + L-Ta <sub>2</sub> O <sub>5</sub>
				1329	64.00	No melting	11
				1527	16.00	No melting	TTBs
		•		1577	19.00	No melting	0
				1623	17.00	No melting	et .
				1643	0.16	No melting	
				1654	0.16	Partially melted	
				1660	0.08	Completely melted	
				1670	0.08	Completely melted	
	•	1527	16	1195 1270	552.00 360.00	No melting No melting	TTBs + L-Ta <sub>2</sub> O <sub>5</sub>
77.78	22.22	800	112				
				1350	48.00	Not melted	TTBs
				1527	16.00	Not melted	
				1642	0.16	Not melted	
				1654	0.16	Completely melted	
75	25	1000	109				1:2 + L-Ta <sub>2</sub> O <sub>5</sub>
				1195	552.00	Not melted	$1:2 + L-Ta_2^2O_5 + TTBs (tr)$
				1270	360.00	Not melted	TTBs + 1:2
				1329	64.00	Not melted	10
				1527	6.00	Not melted	TTBs
				1613	17.00	Not melted	11
				1643	0.25	Completely melted	<del></del>
		1527	6	1722 1195	80.0	Completely melted	
		1327	ъ	1270	552.00 360.00	Not melted Not melted	TTBs + 1:2 (tr)
	•	1722	0.08	1576	16,00	Not melted	TTBs
70	30	1000	109			, 	
				1580	64.00	Not melted	1:2 + TTBs
				1612	0.08	Not melted	
				1627	0.08	Completely melted	
66.67	33.33	1000	109			Not melted	1:2
,		•	•	1329	64.00	Not melted	11
				1524	7.00	Not melted	tI .
			`	1601	1.00	Not melted	
				1620	0.75	Not melted	
				1633	0.08	Not melted	
				1655	0.08	Completely melted	
		٠		1698	0.08	Completely melted	
				1750	0.08	Completely melted	
•				1805	0.08	Completely melted	

60	40	1000	109				
				1602	5.00	Not melted	1:2 + 1:1
				1617	0.08	Not melted	
				1628	0.08	Not melted	
				1632	0.08	Partly melted	
				1664	0.08	Partly melted	
				1685	0.08	Partly melted	
				1690	0.08	Partly melted	<b>*</b>
				1722	0.08	Partially melted	
				1737	0.08	Completely melted	
50	50	600	4				1:1
				1328	16.00	Not melted	
				1527	0.50	Not melted	
				1622	0.50	Not melted	
				1676	0.08	Not melted	
				1782	0.08	Not melted	
	•			1800	0.08	Not melted	
				1821	0.08	Completely melted	

 $<sup>\</sup>frac{a}{2}$  For ease and accuracy of weighing Na<sub>2</sub>O was added to Ta<sub>2</sub>O<sub>5</sub> as NaTaO<sub>3</sub> not as the oxide end member.

L-Ta<sub>2</sub>O<sub>5</sub> - The low temperature polymorph of Ta<sub>2</sub>O<sub>5</sub> H-Ta<sub>2</sub>O<sub>5</sub> - The high temperature polymorph of Ta<sub>2</sub>O<sub>5</sub>

ss - Solid solution

tri - Triclinic

tr - Trace

TTBs - A nonstoichiometric solid solution having an orthorhombic distortion of a tetragonal tungsten bronze-type lattice with superstructure indicating a tripled cell.

1:2 - Na2Ta4011 1:1 - NaTaO3

 $<sup>\</sup>underline{b}^{\prime}$  All specimens were initially calcined in Pt crucibles at the indicated temperatures and time.

 $<sup>\</sup>frac{\text{c}'}{\text{All}}$  subsequent heat treatments below about 1650° were quenched in sealed Pt tubes from the indicated temperatures. Experiment above about 1650° were performed in an inductively heated Ir crucible using sealed 80/20 Pt/Rh tubes.

 $<sup>\</sup>frac{d}{d}$  All phases identified are given in order of amount present at room temperature (greatest amount first). The phases are not necessarily those present at the temperature to which the specimen was heated.

TABLE 5: EXPERIMENTAL DATA FOR THE SYSTEM Mb2O5-KNbO3

Composition		Initial Heat Treatment		Final Heat Treatment S		Results of Physical Observation	Results of X-Ray Diffraction Analyses	
Mole %	, ≛⁄	Temp.	Time Hrs.	Temp.	Time Hrs.			
Nb205	K <sub>2</sub> O	7		•			+ +	
	-	750					H-Nb <sub>2</sub> O <sub>5</sub> + 1:3	
95	5	750	69			No malaina	H-Nb <sub>2</sub> O <sub>5</sub> + TTBs	
		1000	69	1100	114	No melting No melting	#-MD202 + 11BB	
				1200	69	No malting	· u	
				1275	20	No melting	H-Nb <sub>2</sub> O <sub>5</sub> + GTB	
				1350	20	Partially melted (?)	N	
90.54	9.46	800	62					
		1000	96		7.40	No melting	H-Nb2O5 + TTBs	
				1305 1315	140 46.5	No melting Partially melted	GTB (+H-Nb <sub>2</sub> O <sub>5</sub> ?) H-Nb <sub>2</sub> O <sub>5</sub> + HTB + ?	
				1325	17	Partially melted	H-Nb <sub>2</sub> O <sub>5</sub> + GTB + HTB + ?	
90	10	750	69				H-Nb <sub>2</sub> O <sub>5</sub> + 1:3	
		1000	69			No melting	TTBs + H-Nb <sub>2</sub> O <sub>5</sub>	
				1100	23	No melting	<b>*</b>	
				1200	69	No melting	Comp. 1 to Miss Oc. (1 months 2)	
				1250 1275	40 20	No melting No melting	GTB + H-Nb <sub>2</sub> O <sub>5</sub> (+TTBs ?) GTB + H-Nb <sub>2</sub> O <sub>5</sub> (trace)	
				1290	16	No melting	# n-ND205 (CLACE)	
				1350	20	Partially melted	$H-Nb_2O_5 + GTB + TTBs (?) + HTB (?)$	
				1360	20	Partially melted	H-Nb <sub>2</sub> O <sub>5</sub> + TTBs + ?	
				1375	40	Partially melted	H-Nb <sub>2</sub> O <sub>5</sub> + HTB	
88.5	11.5	800	62					
		1000	96	1200		No melting	TTBs + H-Nb <sub>2</sub> O <sub>5</sub> TTBs + H-Nb <sub>2</sub> O <sub>5</sub>	
•				1200 1300	66 18	No melting No melting	GTB	
	•			1515	1	Completely melted		
		1300	18				GTB	
				1190	18	No melting		
				1200 1200	18 66	No melting	GTB	
				1210	71	No melting No melting	GTB	
				1230	71	No melting	GTB,	
				1325	17	Partially melted	TTBs + H-Nb <sub>2</sub> O <sub>5</sub> + ?	
				1335	70.5	Partially melted	<del></del>	
			-	1350	43	Partially melted	<del></del>	
		1515	1	1300	72	Melted Not melted	GTB	
an.	1.7	350						
88	12	750 1000	69 69			No melting	H-Nb <sub>2</sub> O <sub>5</sub> + 1:3 + 1:1 TTBs + H-Nb <sub>2</sub> O <sub>5</sub>	
		1000		1100	45	No melting	" II-MD205	
				1200	69	No melting	"	
				1275	20	No melting	GTB + TTBs	
				1330	18	Partially melted	GTB	
				1335	71	Partially melted	H-Nb <sub>2</sub> O <sub>5</sub> + HTB + ?	
				1350 1360	20 20	Partially melted	GTB $+$ $H-Nb2O5 (?)$	
				1375	40	Partially melted Partially melted	H-Nb <sub>2</sub> O <sub>5</sub> + HTB	
86	14	975	23					
50	4-4	1250	91			Not melted	H-Nb <sub>2</sub> O <sub>5</sub> + TTBs + 1:3 TTBs + GTB	
,		· .	•	1275	19	Not melted		
				1310	19	Partially melted (?)	GTB + TTBs	
	•	•		1332	18	Partially melted	TTBs + H-Nb <sub>2</sub> O <sub>5</sub> + ?	
				1335	120	Partially melted	<del></del>	
				1355 1515	125 1	Partially melted Completely melted		
85	15	975	23				H=Nh_O_ + TTRO ± 1.3 ± 5.3 5.3	
		1250	91			Not melted	$H-Nb_2O_5 + TTBs + 1:3 + 2:3 hyd.$ $TTBs + GTB$	
				1275	19	Not melted		
				1310	19	Partially melted	GTB + TTBs	
				1325	16	Partially melted	71 Mb-0- 1 mm- 1 3	
				1335 1335	19 125	Partially melted	H-Nb <sub>2</sub> O <sub>5</sub> + TTBs + ?	
				1515	125	Partially melted Completely melted		
84	16	975	23				$\text{H-Nb}_2\text{O}_5$ + TTBs + 1:3 + 2:3 hyd.	
		1250	91			Not melted	TTBs + GTB (7)	
				1275	19	Not melted		
		4		1310	19	Partially melted	GTB + TTBs	
				1335 1515	19 1	Completely melted		
				2010	-	Completely melted	· .	

83.66	16.34	750	69				
		1000	69			Not melted	TTBs
				1100	114	Not melted	•
				1176	24	Not melted	<b>a</b>
				1200	69	Not malted	
				1275	21	Not melted	4
				1,279	ī	Not melted	
				1295	1	Partially melted	
				1287	1	Partially melted	
82	18	975	16	w.e-			H-Nb <sub>2</sub> O <sub>5</sub> + TTBs + 1:3 + 2:3 hyd.
		1250	91			Not melted	7-7-3
		4		1275	19	Not melted	
				1335	18	Completely melted	
21 20							
81.25	18.75	800	62				
		1000	96			Not melted	TTBs
				1225	5	Not melted	
				1275 1335	1 <del>9</del> 18	Partially melted (?) Completely melted	TTBS + HTB
				1,33	10	combideet merced	1183 7 1118
80	20	750	69				$1:3 + H-Nb_2O_5$
				1000	69	Not melted	TTBs + 1:3 + 2:3
				1100	114	Not melted	TTBs
				1200	69	Not melted	TTBs + 1:3
				1275	21	Partially melted	TTBs
				1279	1	Completely melted	
		900	24				TTBs + 1:3 + H-Nb <sub>2</sub> O <sub>5</sub>
		1000	59				TTBs
				775	42		n 
				800	65		
		,		850	40		
		(uncal	cined)	775	42		TTBs + 1:3 + H-Nb <sub>2</sub> O <sub>5</sub>
				800	65		"
				850	40		. "
77,78	22,22	800	69			****	
			**	1100	45		TTBs + 1:3
				1200	69	Not melted	
75	25	750	69				1:3 + 2:3 + H-Nb <sub>2</sub> O <sub>5</sub>
		1000	69			Not melted	1:3 + TTB
				1100	114	Not melted	ti
				1200	69	Not melted	1:3
74	26	800	45				
	20	800	43	1000			7 3.2 ± 4.0
				1000	45	Not melted	1:3 + 4:9
73	27	800	45				
				1100	45	Not melted	1:3 + 4:9
72.2	27 <b>.</b> B	800	62				777 <del>7</del>
		1000	96	~-		Not melted	1:3 + 4:9
70	30	750	69				4:9 + 1:3
		1000	69			Not melted	*1
				1100	114	Not melted	<del></del>
69.23	30.77	820	73				
		1000	79			<del></del>	
				1100	21	Not melted	4:9
				1100	45	Not melted	4:2
				1188	43	Not melted	4:9
				1205	2	Not melted	•
					•		
68	32	800	45		***		
				1100	45	Not melted	4:9 + TTB
66 67	33.33	200	00		_		3.3.4.5.3
66.67	33.33	800 1000	89 164			Not welfed	1:3 + 2:3
		T000	194	 975	21	Not melted	TTB + 4:9
				975 985	45	Not melted	TTB + 4:9 TTB + 4:9
				1100	114	Not melted Not melted	11B T 4:9
				1195	20	Partially melted	4:5
				1470		rarerary mercea	4.5
65	35	750	64			=====	70555
		1000	69			Not melted	TTB
				1100	21	Not melted	TTB
				1195	18	Partially melted	TTB
				1215	0.17	Melted	
		1398	ì			Melted	(?)
				1170	2	Not melted	TTB
				1170	19	Not melted	TTB
				1183	19	Partially melted	TTB + 4:9 + 2:3 hyd.
				1189	2	Partially melted	
		900	47	1191	67	Partially melted	4:9 + 2:3 hyd.
		800 1000	47 70				
		1000	70	1125	8	Not malted	TTB
				****	v	Not melted	TTB

63.5	36.5	800	65		10° 100		
		1000	45			Not melted	TTB + 2:3 hyd.
		4000	1	1143	2	Not melted	
				1158	ĭ	Not melted	Ħ
				1165	ī	Not melted	
				1170	2	Partially melted	TTB
				1180	2	Partially melted	(?)
				1200	0.5	Melted	TTB
				1200	0.5	MAT CAT	
62	38	800	65				
		1000	45			Not melteđ	2:3 hyd. + TTB
				1130	16	Not melted	**
				1143	2	Not melted	
				1158	1	Not melted	
				1160	1	Not melted	
				1165	1	Not melted	
		-		1170	2	Completely melted	
				1180	2	Completely melted	<b></b>
				1200	0.5	Completely melted	2:3 + TTB
60	40	750	69	·			1:3 + 2:3 + 1:1
		1000	69			Not melted	2:3 + 2:3 hyd.
				1000	20	Not melted	2:3
				1143	2	Not melted	
				1158	1	Not melted	
				1165	1	Completely melted	
				1170	2	Completely melted	
				1172	2	Completely melted	
				1180	2	Completely melted	
55	45	750	69				1:1 + 1:3 + 2:3
		1000	69			Not melted	1:1 + 2:3 + 2:3 hyd.
ED 05	46.15	750	60				1:1 + 2:3 + 1:3
53.85	46.15	750	69			Not malted	1:1 + 2:3 + 1:3 1:1 + 2:3 + 2:3 hyd.
		1000	69			Not melted	1:1 + 2:3 + 2:3 hyd.
				1000	20	Not melted	
				1100	23	Partially melted	1:1 + 2:3
50	50	750	69				1:1
		1000	69			Not melted	1:1

 $<sup>\</sup>frac{a}{}$  For ease and accuracy of weighing  $K_2O$  was added to  $Nb_2O_5$  as  $KNbO_3$  not as the oxide end member.

 $<sup>\</sup>frac{b}{A}$  All specimens were initially calcined in Pt crucibles at the indicated temperatures and time.

 $<sup>^{</sup>m c/}$  All subsequent heat treatments were quenched in sealed Pt tubes from the indicated temperatures.

All phases identified were given in order of amount present at room temperature (greatest amount first). The phases were not necessarily those present at the temperature to which the specimen was heated. H-Nb<sub>2</sub>O<sub>5</sub>. - The high temperature form of Nb<sub>2</sub>O<sub>5</sub>.

<sup>1:3 -</sup> NaNb<sub>3</sub>O<sub>8</sub>.

GTB - Gatehouse Tungsten Bronze - A nonstoichiometric solid solution having a large tetragonal unit cell with 7-sided tunnels first described by B. M. Gatehouse for a rubidium niobate of unknown composition.

TTBs - A nonstoichiometric solid solution having an orthorhombic distortion of a tetragonal tungsten bronze-type lattice with superstructure indicating a tripled cell.

<sup>4:9 -</sup> A compound having the apparent composition 4K2O:9Nb2O5 (K8Nb18O49).

TTB - Tetragonal Tungsten Bronze - A nonstoichiometric solid solution having an undistorted tetragonal lattice with no superstructure.

<sup>2:3 -</sup> K4Nb6O17.

<sup>2:3</sup> hyd. - The hydrated form of  $K_4 Nb_6 O_{17}$  in equilibrium with atmospheric moisture at room temperature.

<sup>1:1 -</sup> KNbO3.

HTB - A phase resulting from a quenched liquid with an x-ray pattern resembling a hexagonal tungsten bronze.

<sup>? -</sup> An unknown phase which apparently results from quenching a liquid.

TABLE 6: EXPERIMENTAL DATA FOR THE SYSTEM Ta205-KTa03

Compos	ition	Initial Treats	l Heat		Heat ment⊆⁄	Results of Physical Observation	Results of X-Ray Diffraction Analyses 💇	
Mole * a/		Temp.	Time	Temp.	Time	Opedivacion	Analyses 😅	
		°C	Hrs.	*C	Hrs.			
Ta205	K20							
95	5	1000	109				L-Ta2O5 + 1:5 + TTBs	
			<del></del> -	1337	16	No melting	L-Ta <sub>2</sub> O <sub>5</sub> + TTBs	
				1549	16	No melting	H-Ta205 + GTB	
				1602	16	No melting	11	
90	10	900	168		- <b>-</b>	<del></del>		
,,,		300	*00	1466	168	No melting	TTBs + L-Ta <sub>2</sub> O <sub>5</sub> + H-Ta <sub>2</sub> O <sub>5</sub>	
				1500	17	No melting	TTBs + H-Ta <sub>2</sub> O <sub>5</sub> + L-Ta <sub>2</sub> O <sub>5</sub>	
				1553	1.0	No melting		
				1609	17	No melting	GTB + H-Ta <sub>2</sub> O <sub>5</sub> (?)	
				1715	1.5	Partially melted	**************************************	
				1747	1.5	Partially melted	HTB + H-Ta <sub>2</sub> O <sub>5</sub>	
		1000	109	1795 1521	1.5 44	Partially melted No melting	HTB + H-Ta <sub>2</sub> O <sub>5</sub> + L-Ta <sub>2</sub> O <sub>5</sub>	
		1000	109	1601	0.75	No melting		
				1619	64	No melting	GTB + H-Ta <sub>2</sub> O <sub>5</sub>	
				1620	16	No melting	GTB + H-Ta <sub>2</sub> O <sub>5</sub>	
		1609	17	1466	168	No melting	GTB + TTBs	
				1500	1.7	No melting		
				1521	44	No melting		
				1624 1647	0.17 0.17	No melting		
				1657	0.17	No melting Partially melted		
89	11	800	115		<b></b>	~~~	1:5 + L-Ta <sub>2</sub> O <sub>5</sub>	
				1600	12	No melting	GTB	
87.5	12.5	900	216				1.5 A I-We O	
		200	210	1601	17	No melting	1:5 + L-Ta <sub>2</sub> O <sub>5</sub> GTB + TTBs	
				1640	19	Partially melted	GTB	
83.33	16.67	800	192				L-Ta <sub>2</sub> O <sub>5</sub> + 1:2	
		900	168				1:5	
				950 1100	360 64	No melting	1:5 + TTBs (tr)	
		1000	109			No melting	1:5 + TTBs + L-Ta <sub>2</sub> O <sub>5</sub>	
				1337	16		TTBs + 1:5 + L-Ta <sub>2</sub> O <sub>5</sub> TTBs	
				1532	16		TTBs	
				1610	16		TTBs	
				1624	0.17	Not melted		
				1634	6.5	Partially melted	GTB + HTB	
				1635 1647	0.17 0.17	Partially melted Partially melted		
				2021	0.11	rarcially matcad		
80	20	1000	109				TTBs + 1:5	
	•		•	1337	16	Not melted	TTBs	
				1538	16	Not melted	TTBs + 9L	
				1614	16	Not melted	HTB + TTBs	
				1617 1618	16 16	Not melted	te ne	
		1617	16	1325	168	Not melted Not melted	TTBs	
	•	1618	16	1350	144	Not melted	"	
78.25	21.75	800	91	1600	6		HTB	
77.78	22.22	900	168					
	<b></b>			1549	1.0	Not melted	TTBs + 9L + HTB	
				1571	1.0	Not melted		
				1575	10	Not melted		
				1581	19	Not melted	нтв .	
				1602	6.5	Not melted	нтв	
				1607 1620	0.5 16	Not melted Partly melted (?)	UMB + Or	
				1632	19	Partly melted (?)	HTB + 9L	
		1581	19	1549	19	Not melted	HTB + TTBs	
		1602	6.5	1549	1.0	Not melted		
				1571	1.0	Not melted	HTB + TTBs	
				1603	0.17	Not melted		
				1613	0,17	Not melted	**************************************	
				1624	0.17	Partly melted		

	0.5	000	160		***		1:2 + 1:5 + TTBs
75	25	900	168	1304	96	Not melted	TTBs + TTB
				1340	19	Not melted	
		•		1579	16	Not melted	9L + HTB
				1600	68	Not melted	
				1692	2	Completely melted	llL + HTB
		1000	109				TTBs + TTB + 3L
		1000	103	1100	64	Not melted	TTBs + TTB
				1327	120	Not melted	TTBs + TTB
				1340	19	Not melted	9L + TTBs
				1340	72	Not melted	n
				1538	64	Not melted	D .
				1575	10	Not melted	16
				1598	0.5	Not melted	
				1611	5.5	Not melted	9L + HTB
				1634	16	Melted	HTB + H-1:3
				1646	1.0	Melted	W
		1340	72	1040			***
		1340	14	1600	68	Not melted	9L + HTB
					16	Not melted	9L + HTB
		1240	73	1606	10	NOC METCEC	32 · 112
		1340	72	1610		Not melted	9L + HTB
		900	1.5	1610		HOC WEIGH	28
		1000	109	3.500	70	Mak malkad	9L + TTBs
		1538	64	1528	72	Not melted	70 1 11BB
		1600	68	1603	0.25	Not melted	
				1613	0.25	Melted	
				1624	0.25	Melted	<del></del>
73.85 <u>e</u> /		1450				Net1403	9L
73.85 <del>-</del>	26.15	1462	6 <del>9</del>			Not melted	
				1304	96	Not melted	9L + TTB
			,	1327	120	Not melted	9L + TTB
				1603	0.17	Not melted	
				1613	0.17	Partially melted	
e/							A 16*
73.67 º/	26.33	1443	89			Not melted	9L + 16L
<b>-/</b>							
<sub>73.5</sub> €/	26.5	1466	64				16L + 9L (tr)
				1400	48	Not melted	16L + 9L + 11L
				1574	0.5	Not melted	<del></del>
				1574	16	Not melted	16L
				1586	16	Not melted	
				1594	4	Not melted	
				1603	0.17	Not melted	<del></del>
			•				
72.73	27, 27	1000	68				
				1361	24	Not melted	9L + TTB
				1400	48 .	Not melted	**
				1573	0.17	Not melted	
				1573	16	Not melted	11L + 16L
				1589	0.17	Not melted	
				1629	0.17	Completely melted	
				1632	0.08	Completely melted	
						·	* 10 °
71.43	28.57	1000	68				TTBs + TTB + 1:2 + 1:5 + 3L
				1361	24	Not melted	9L + TTB
				1438	75	Not melted	16L + TTB
				1465	336	Not melted	11L
				1507	139	Not melted	11L
				1578	1.0	Not melted	*
				1583	3.5	Not melted	11L + TTB
				1591	88	Not melted	11L
				1611	0.08	Not melted	
				1613	0.17	Not melted	
				1618	0.08	Not melted	<u>-</u>
	•	•		1618	16	Not melted	
				1624	0.08	Melted	
				1626	1.5	Melted	
				1694	1.0	Melted	11L
		1575	10	:			11L + TTB
		1583	3.5	1438	75		11L + 16L + TTB
				1465	336		11L
				1507	137		11L
	,					•	
70	30	1000	68				
	-			1361	24	Not melted	TTB + 9L
				1622	1.0	Melted	11L + TTB (tr)
					4.5	•	
					46		

66.67	33.33	800	192			****	1:2 + 1:1 + L-Ta <sub>2</sub> O <sub>5</sub>
		900	168				1:2
			è	950	360	Not melted	ti
				1100	64	Not melted	TTB
		1000	109				1:2 + TTB + 3L
				798	163	Not melted	н
				1340	19	Not melted	TTB
				1340	72 .	Not melted	, m
				1515	64	Not melted	TTB + 11L
				1538	20	Partially melted	llL + TTB
				1616	1.0	Completely melted	llL + TTB + 9L
66	34	800	90	1400	10	Not melted	TTB
65	35	800	112		~~	10 TO 10 TO 10	1:1 + 1:2 + L-Ta <sub>2</sub> O <sub>5</sub>
				1350	66	Not melted	TTB (+1:1 ?)
60	40	1000	92				TTB + 1:1
				1318	64	Not melted	TTB + 1:1
				1368	0.5	Not melted	· · · · · · · · · · · · · · · · · · ·
				1412	16	Partially melted	
55	45	1000	92			4744	1:1 + TTB
				1318	64	Not melted	**
				1368	0.5	Not melted	<b>-</b>
				1480	18	Completely melted	
50	50	600	4				1:1
				1340	19	Not melted	11
				1368	0.5	Not melted	
				1375	0.5	Completely melted	

 $<sup>\</sup>underline{a}$  For ease and accuracy of weighing  $K_2O$  was added to  $Ta_2O_5$  as  $KtaO_3$  not as the oxide end member.

 $<sup>\</sup>underline{\mathbf{b}}'$  All specimens were initially calcined in Pt crucibles at the indicated temperature and time.

 $rac{ extsf{c}'}{ extsf{All}}$  subsequent heat treatments were quenched in sealed Pt tubes from the indicated temperatures.

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

L-Ta<sub>2</sub>O<sub>5</sub> - The low temperature polymorph of Ta<sub>2</sub>O<sub>5</sub>. H-Ta<sub>2</sub>O<sub>5</sub> - The high temperature polymorph of Ta<sub>2</sub>O<sub>5</sub>.

<sup>1:5 -</sup> KTa<sub>5</sub>O<sub>13</sub> - An orthorhombic compound of undetermined structure.

GTB - Gatehouse Tungsten Bronze - A nonstoichiometric solid solution having a large tetragonal unit cell with 7-sided tunnels first described by B. M. Gatehouse for a rubidium niobate of unknown composition.

TTBs - A nonstoichiometric solid solution having an orthorhombic distortion of a tetragonal tungsten bronze-type lattice with superstructure indicating a tripled cell.

HTB - Hexagonal Tungsten Bronze - A nonstoichiometric solid solution with an x-ray pattern resembling a hexagonal tungsten bronze.

<sup>9-</sup>L - A hexagonal phase with a c-axis ~ 9 x 4 Å.

<sup>16-</sup>L - A hexagonal phase with a c-axis ~ 16 x 4 Å.

<sup>11-</sup>L - A hexagonal (rhombohedral) phase with a c-axis ~ 11 x 4 Å.

<sup>3-</sup>L - An apparently metastable hexagonal phase with a c-axis  $^{\circ}$  3 x 4 Å.

 $<sup>1:2 -</sup> K_2 Ta_4 O_{11}$ .

H-1:3 - A monoclinic phase which apparently results from quenching a liquid near the composition K20:3Ta205.

TTB - Tetragonal Tungsten Bronze - A nonstoichiometric solid solution having an undistorted tetragonal lattice with no superstructure.  $1:1 - KTaO_3.$ 

e/ Made from the 1000° calcines of the 75:25 and 72.73:27.27 mixtures.

TABLE 7

Summary of LeRC Measurements a/ of Dielectric Loss on Polycrystalline NBS Samples

Composition		Frequency, Hz b/				Resonance c/		
		10 <sup>3</sup>	104	105	10 <sup>6</sup>			
6Li <sub>2</sub> O:17TiO <sub>2</sub>	ε' ε"	54 622	30 67	22 9.8	19 1.9	No Peaks		
K <sub>1.55</sub> Li <sub>1.26</sub> (LiTi)O <sub>16</sub>	ε"	796 401	418 194	225 9 <b>4</b>	132 43	No Peaks		
K <sub>1.6</sub> Mg <sub>0.8</sub> Ti <sub>3.8</sub> O <sub>8</sub>	٤ ' د	80 4.0	74 3.1	71 1.4	70 0.9	$\Delta E = 15.1 \text{ kcal/mole}$ $\rho_{25} = 2.7 \times 10^9 \text{ N-cm}$		
5Li <sub>2</sub> O:95Ta <sub>2</sub> O <sub>5</sub> d/	ε¹ ε"	64 7	59 2	57 0.7	58 0.3	No Peaks		
5Li <sub>2</sub> O:95Ta <sub>2</sub> O <sub>5</sub>						No Peaks		
Li <sub>1.8</sub> Ta <sub>5.8</sub> W <sub>0.2</sub> O <sub>16</sub>	ε' ε"	19 0.3	18 0.02	18 < 0.002	18 0.05	No Peaks		
Na <sub>2</sub> 0:13Nb <sub>2</sub> 0 <sub>5</sub>	ε' ε"	20 0.9	19 0. <b>4</b>	19 0.3	19 0.2	No Peaks		
NaNb <sub>3</sub> O <sub>8</sub>			<del>-</del>			No Peaks		
Na <sub>10.8</sub> Nb <sub>34</sub> W <sub>1.8</sub> O <sub>95.6</sub>						$\Delta E > 10$ $\rho_{25} \approx 3 \times 10^7 \Omega$ -cm		
21Na <sub>2</sub> O:79Ta <sub>2</sub> O <sub>5</sub>	ε" ε"	22 0.3	22 0.2	21 0.1	21 0.2	No Peaks		
7K <sub>2</sub> O:13Nb <sub>2</sub> O <sub>5</sub>						Slight Peaks $\Delta E = 5.6$ to $6.8$ $\rho_{25} \sim 1$ to $5 \times 10^4$ $\Omega$ cm		
11.5K <sub>2</sub> 0:88.5Nb <sub>2</sub> 0 <sub>5</sub>	ε" ε"	25 5.1				No Peaks		
3K <sub>2</sub> O:13Nb <sub>2</sub> O <sub>5</sub>						No Peaks		
KNb308 €/	ε" ε"	< 100 < 2	< 100 < 0.1		< 100 < 0.1	No Peaks, two orientations.		
к <sub>4</sub> Nb <sub>6</sub> O <sub>17</sub> •хн <sub>2</sub> О <u>е</u> /	ε' ε"	536 854	264 64	102 < 1	64 < 1	No Peaks, two orientations.		
11K20:89Ta2O5	٤'	52	43	38	36	High Temp. $\Delta E = 10.3 \text{ kcal/mole}$ $\rho_{2.5} = 2 \times 10^8 \Omega - \text{cm}$		
	ε"	6.1	4.7	2.6	1.1	Low Temp. $\Delta E = 8 \text{ kcal/mole}$ $\rho_{25} = 1 \times 10^8 \Omega \text{-cm}$		
к <sub>2</sub> 0:5та <sub>2</sub> 0 <sub>5</sub>	€"	18 0.03	18 0.01	18 0.01	18 0.1	No Peaks		
к <sub>2</sub> 0:4та <sub>2</sub> 0 <sub>5</sub>	€"	22 0.04		22 0.1		No Peaks		
21.75K <sub>2</sub> O:78.25Ta <sub>2</sub> O <sub>5</sub>				<del></del>		$\Delta E = 7.9 \text{ kcal/mole}$ $\rho_{25} = 1 \times 10^4 \Omega \text{-cm}$		
34K <sub>2</sub> O:66Ta <sub>2</sub> O <sub>5</sub>	•					$\Delta E \approx 20$ to 25 kcal/mole $\rho_{25} = 6 \times 10^{13} \Omega$ -cm		
K <sub>0.51</sub> Ta <sub>0.51</sub> W <sub>0.49</sub> O <sub>3</sub>						No Peaks.		

a/ Measurements made by H. E. Kautz, LeRC.

b/ at 25°C.

 $<sup>\</sup>mathfrak{L}^{\prime}$  Values of  $\Delta E$  obtained from  $\epsilon$  vs temperature measurements.

d√ hot pressed specimen.

 $<sup>\</sup>frac{e}{}$  Single crystal specimens, supplied by K. Nassau, BTL.

TABLE 8
ALKALI TANTALATE PELLET FABRICATION

Composition	Calcine	Forming	Sintering	X-Ray	
5 Li <sub>2</sub> O:95 Ta <sub>2</sub> O <sub>5</sub>	1000°C - 144 hours	Hot pressed 1200°C	Refire 1500°C	Low Ta <sub>2</sub> O <sub>5</sub> , crystallinity improved by refiring	
		Hot pressed 1300°C	Rafire 700°C 18 hours	Low Ta <sub>2</sub> O <sub>5</sub> , crystallinity improved by refiring	
Li <sub>2</sub> O:3 Ta <sub>2</sub> O <sub>5</sub> (LiTa <sub>3</sub> O <sub>6</sub> )	1000°C - 23 hours 1050°C - 44 hours	10,000 psi	1300°C - 22 hours air quenched	Single phase, LiTa <sub>6</sub> O <sub>15</sub> F- type	
"Substituted LiTa <sub>3</sub> O <sub>8</sub> " Li <sub>1.8</sub> Ta <sub>5.8</sub> W <sub>0.2</sub> O <sub>16</sub>	1000°C - 23 hours 1050°C - 44 hours	10,000 psi	1400°C - 23 hours air quenched	Single phase, LiTa <sub>6</sub> O <sub>15</sub> F- type	
"Tetragonal Bronze" 21 Na <sub>2</sub> O:79 Ta <sub>2</sub> O <sub>5</sub>	1000°C - 23 hours 1050°C - 44 hours	10,000 psi	1400°C - 17 hours air quenched	Single phase bronze	
11 K <sub>2</sub> 0:89 Ta <sub>2</sub> O <sub>5</sub>	800°C - 90 hours	10,000 psi	1600°C - 12 hours Removed at tem- perature and placed on chill block for rapid cooling	Single phase	
K <sub>2</sub> O:5 Ta <sub>2</sub> O <sub>5</sub>	800°C - 90 hours	10,000 psi	1500°C - 20 hours air quenched	Tetragonal bronze	
K <sub>2</sub> O:4 Ta <sub>2</sub> O <sub>5</sub>	800°C - 90 hours	10,000 psi	1400°C - 20 hours air quenched	Tetragonal bronze	
21.75 K <sub>2</sub> Q:78.25 Ta <sub>2</sub> Q <sub>5</sub>	800°C - 91 hours	10,000 psi	1600°C - 6 hours Pellet sealed in platinum, welded closed. Removed at temperature and water-quenched	After removal of surface layer by grinding, single phase hexagonal bronze	
34 K <sub>2</sub> O:66 Ta <sub>2</sub> O <sub>5</sub>	800°C - 90 hours	10,000 psi	1400°C - 10 hours Cooled at 120°C/hour	Single phase	
K <sub>.51</sub> Ta <sub>.51</sub> W <sub>.49</sub> O <sub>3</sub> (Pyrochlore)	800°C - 34 hours	10,000 psi	950°C - 27 hours Removed at tem- perature and placed in quartz tube, evacuated to p < 10 <sup>-5</sup> Torr and sealed to prevent hydration	X-ray of pellet im- pregnated with silicone resin to prevent hydration showed single phase pyrochlore	

 $<sup>\</sup>underline{a\prime}$  Hot pressing performed by the Haselden Co., San Jose, California.

TABLE 9

ALKALI NIOBATE PELLET FABRICATION

Composition	Calcine	Forming	Sintering	X-Ray	
Li <sub>2</sub> O:14 Nb <sub>2</sub> O <sub>5</sub>	1000°C - 18 hours 1250°C - 68 hours	10,000 psi	1250°C - 12 hours Cooled at 180°C/Hr.	Single phase N-Nb <sub>2</sub> O <sub>5</sub> after surface grinding	
Na <sub>2</sub> O:13 Nb <sub>2</sub> O <sub>5</sub>	800°C - 114 hours	10,000 psi	1225°C - 8 hours Cooled at 180°C/Hr.	Single phase except for one line of phase next highest in Na <sub>2</sub> O content.	
NaNb <sub>3</sub> O <sub>8</sub>	800°C - 62 hours 1000°C - 96 hours	18,000 psi	1225°C - 1 hour air quenched	Single phase 1:3	
Na <sub>10.8</sub> (NbO) 4 (WO) <sub>1.8</sub> Nb <sub>30</sub> O <sub>90</sub>	800°C - 89 hours 1000°C - 70 hours	18,000 psi	1225°C - 3 hours air quenched	Single phase, pattern similar to NaNb3O8	
11.5 K <sub>2</sub> 0:88.5 Nb <sub>2</sub> O <sub>5</sub>	800°C - 62 hours 1000°C - 96 hours	18,000 psi	1300°C - 6.5 hours air quenched. Specimens poorly sintered. Excessive grain growth under all conditions. Hot pressing in progress.	Gatehouse tungsten bronze	
3 K <sub>2</sub> O:13 Nb <sub>2</sub> O <sub>5</sub>	800°C - 62 hours 1000°C - 96 hours	18,000 psi	1225°C - 1 hour air quenched	Tetragonal tungsten bronze with super- structure.	
7 K <sub>2</sub> 0:13 Nb <sub>2</sub> 0 <sub>5</sub>	800°C - 47 hours 1000°C - 70 hours	18,000 psi	1225°C - 1 hour air quenched	Tetragonal tungsten bronze without super- structure.	

TABLE 10

## ALKALI TITANATE PELLET FABRICATION

Composition	Calcine	Forming	Sintering	X-Ray
Ramsdellite 6 Li <sub>2</sub> O:17 TiO <sub>2</sub> (Li <sub>2</sub> 9(Li <sub>3</sub> Ti <sub>1,7</sub> )O <sub>4</sub> )	9 pbw 6Li <sub>2</sub> 0:17TiO <sub>2</sub> calcined: 800°C - 2.5 Hr. 1050°C - 68.5 Hr. 1 pbw raw batch, 6 Li <sub>2</sub> CO <sub>3</sub> :17 TiO <sub>2</sub>	10,000 psi	1200°C - 17 hours air quenched.	Single phase Ramedellite
K <sub>2</sub> O:MgO:4 TiO <sub>2</sub> [K <sub>1.6</sub> (Mg <sub>.8</sub> Ti <sub>3.2</sub> )O <sub>8</sub> ]	800°C - 18 hours 1000°C - 24 hours	10,000 psi 0.5 percent by weight stearic acid as binder.	1200°C - 13 hours air quenched	Single phase, "Cmcm-phase"

TABLE 11 Summary of Crystal Growth Experiments

System	Phase	Melt Composition	Method	Comments
к <sub>2</sub> 0-мь <sub>2</sub> 0 <sub>5</sub>	7:13 (TTB) 17.5:82.5 (TTBs) 17.5:82.5 (TTBs) 12.5:81.5 (GTB)	36.5K <sub>2</sub> O:63.5Nb <sub>2</sub> O <sub>5</sub> 17.5K <sub>2</sub> O:82.5Nb <sub>2</sub> O <sub>5</sub> 20.0K <sub>2</sub> O:80.0Nb <sub>2</sub> O <sub>5</sub> 15.0K <sub>2</sub> O:85.0Nb <sub>2</sub> O <sub>5</sub>	TSSG* TSSG TSSG TSSG	Yield: 4:9 + small amount 7:13 + 2:3 Yield: polycrystalline multiphase Yield: polycrystalline TTB Yield: Nb <sub>2</sub> O <sub>5</sub>
к <sub>2</sub> о-та <sub>2</sub> о <sub>5</sub>	1:2 (TTB)	45K <sub>2</sub> O:55Ta <sub>2</sub> O <sub>5</sub>	TSSG	Yield: single phase
Li20-TA205	1:3 5:95 (L-Ta <sub>2</sub> O <sub>5</sub> )	25Li <sub>2</sub> O: 75Ta <sub>2</sub> O <sub>5</sub> 15Li <sub>2</sub> O: 85Ta <sub>2</sub> O <sub>5</sub>	Czochralski TSSG	Yield: single crystals Yield: single crystals
Na <sub>2</sub> O-Nb <sub>2</sub> O <sub>5</sub>	1:3 (TTBs)	25Na <sub>2</sub> O: 75Nb <sub>2</sub> O <sub>5</sub>	Czochralski	Yield: single crystals
Na <sub>2</sub> O-Ta <sub>2</sub> O <sub>5</sub>	21:79 (TTBs)	25Na <sub>2</sub> O: 75Ta <sub>2</sub> O <sub>5</sub>	Czochralski	Excessive vaporization of Na <sub>2</sub> O

<sup>\*</sup> Top seeded solution growth.

TABLE 12
Crystallographic Date for Phases in the NbgOg-Alkali Niobate and TegOg-Alkali Tantalate Systems

System	Designation	Composition Mol %	Syrmetry	*	Unit C	ell Dimensi C Ā	one u	ß	Y	Conditions Limiting Possible Reflections	Probable Space Groupe
Nb <sub>2</sub> O <sub>5</sub> -LiNbO <sub>3</sub>	N-ND <sub>2</sub> O <sub>5</sub> 1:3	Nb <sub>2</sub> O <sub>5</sub> Li <sub>2</sub> O 93.33 6.67 75 25	Monoclinic Monoclinic	25.518 7.457	3.827 5.035	17,554 15.264		124°59.1° 107°18.7′		hkl:h+k=2n h01:t=2n 0k0:k=2n	C2,Cm,C2/m P2 <sub>1</sub> /c
Ta <sub>2</sub> O <sub>5</sub> -LiTaO <sub>3</sub>	I # 0	Ta <sub>2</sub> O <sub>5</sub> Li <sub>2</sub> O 95 5	Orthorhombic	6.198	40.29	3.888				None	pm, pq 1/
	L-Ta <sub>2</sub> O <sub>5</sub> ss L-1:3	75 25	Monoclinic	7.41	5.10	15.12		107°12'		h0l:l=2n OkO:k=2n	P2 <sub>1</sub> /c
	M-1:3	75 25	Monoclinic	9,420	11.536	5.055		91°32'		hkt:h+k=2n h0t:t=2n	Cc.C2/c
	H-1:3	75 25	Orthorhombic	16.716	8. 941	3.840				Ok 2 : 2 = 2n	Pmma,P2 <sub>1</sub> ma, Pm2a
Nb <sub>2</sub> O <sub>5</sub> -NaNbO <sub>3</sub>	•	Nb205 Na20									
* *	1:13	92.86 7.14 80 20	Monaclinic Orthorhombic	22.40 ∿14.7	3.834 ∿10.2	15.37 v3.9		91°28.2°		hkl:h+k=2n hkl:h+k=2n	C2,Cm,C2/m C222,Cm2m, Cmm2,Cmam
•	TTBs	75 25	Orthorhombic	12.364	36.992	3.955				0k l : k=2n h0 l : h=2n	Pham, Pha21
	1:2	66.67 33.33	Monoclinic	10.840	6.162	12,745		106"13.2"		hkl:h+k≠2n h0l:l=2n	Cc,C2/c
Ta <sub>2</sub> O <sub>5</sub> -NaTaO <sub>3</sub>		Ta <sub>2</sub> O <sub>5</sub> Na <sub>2</sub> O					100				
impos maraos	TTBs	80 20	Orthorhombic	12.397	37.34	3,903	-;			h0t:h=2n	Pmam, P2 <sub>1</sub> am, Pma2
	TTBs 1:2	75 25 66.67 33,33	Orthorhombic Hexagonal	12.398 6.120	37.28	3, 899 36, 629				h01:h=2n hk1:-h+k+1=3n h01:1=2n	R3c,R3c
Nb <sub>2</sub> O <sub>5</sub> +KNbO <sub>3</sub>	GTB TTBs	Nb <sub>2</sub> O <sub>5</sub> K <sub>2</sub> O 88.5 11.5 83.33 16.67	Tetragonal Orthorhombic	27.518 12.519	 37 <b>.</b> 558	3.9687 3.952				h00:h=2n h01:h=2n	P42 <sub>1</sub> 2,P42 <sub>1</sub> m Pmam,P2 <sub>1</sub> am, Pma2
	TTBs	80 20	Orthorhombic	12.545	37.636	3.957				a	4 11
	НТВ 1:3	Unknown* 75 25	Kexagonal Orthorhombic	7.511 8.925	21.232	3.889 3.808				None hkl:k+l=2n h0l:h=2n	P6/mmm Amam, A2 <sub>1</sub> am,
	4:9 TTB	69.23 30.77 65 35	Triclinic Tetragonal	13.353 12.589	13,915 	15.022 3.981	82°11.8°	69°42'	89°4.3' 	None h0%:h=2n	Ama2 P <sub>1</sub> ,P1 Pmam,P2 <sub>1</sub> am, Pma2
	2:3	60 40	Orthorhombie	7.822	33.019	6.481				h0£:h+£=2n hk0:k=2n	Pmnb, P21nb
	2:3 hyd.	60 40	Orthorhombic	7.824	38.073	6.485				h00: h=2n 0k0: k=2n 00l: k=2n	P2 <sub>1</sub> 2 <sub>1</sub> 2 <sub>1</sub>
Ta,05~KTa03		Τα <sub>2</sub> 0 <sub>5</sub> κ <sub>2</sub> 0									
2-53	GTB	88.5 11.5	Tetragonal	27.55		3.899				h00:h=2n	P42 <sub>1</sub> 2, P42 <sub>1</sub> m
	1:5	83.33 16.67	Orthorhombic	5.654	10.713	16.80				h0&:	Pbcm, Pbc2
	TTBs	80 20	Orthorhombie	12.547	37.641	3.922				h01:h=2n	Pmam, P2 <sub>1</sub> am, Pma2
	HTB 9L	78.25 21.75 73.85 26.15		7.527 7.55	3	3,901 36,583				None hhl:l=2n	P6/mmm P63mc, P62c,
	16L	73.5 26.5	Hexagonal	7.542	*	65.57				hhl: l=2n	P63mmc
	11L 1:2	71.43 28.57 66.67 33,33		7.54 6.283		43.512 36.878				hkl:-h+k+l=3n hkl:-h+k+l=3n	R3,R3,R32, R3m,R3m
	TTB	66 34	Tetragonal	12.569	*	3.957				h0l:h=2n	Pmam,P2 <sub>1</sub> am, Pma2
	3L*	75 25	Hexagonal	9.051		12.284				hOl: l=2n	P63cm.P6c2, P63mcm
•	н-1:3*	75 25	Monoclinic	14.615	3.774	6.557		98°30'		None	P2, Pm, P2/m

 $<sup>\</sup>frac{1}{2}$  Two dimensional plane groups.

<sup>\*</sup> Metastable phase obtained from quenched liquid.

<sup>\*\*</sup> Probably due to reaction with atmospheric moisture-NaNb  $_{6}o_{16}\left( \text{OH}\right) .$