

Phase Equilibria as Related to Crystal Structure in the System Niobium Pentoxide-Tungsten Trioxide

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The phase equilibrium diagram for the binary system niobium pentoxide-tungsten trioxide has been constructed from results of x-ray diffraction studies on both single crystals and powders and from fusion characteristics. Twelve stable compounds have been found in the system. The exact composition of eight of these compounds has been established by single crystal analyses at the $\text{Nb}_2\text{O}_5 : \text{WO}_3$ ratios of 6:1, 13:4, 7:3, 8:5, 9:8, 1:1, 4:9, and 2:7. The approximate compositions of the remaining four other phases are 30:1, 6:11, 1:11, and 1:15. The 6:1, "6:11", 4:9, and 2:7 phases melt congruently at 1476, 1378, 1380, and 1357 °C, respectively. The "30:1", 7:3, 8:5, 9:8, "1:11", and "1:15" phases melt incongruently at 1470, 1440, 1385, 1375, 1356, and 1358 °C, respectively; and the 13:4 and 1:1 phases decompose before melting at 1435 and 1115 °C, respectively. The 8:5, 9:8, "6:11", 2:7, "1:11", and "1:15" compounds are shown on the phase diagram as having minimum temperatures of stability. One metastable phase having a narrow range of composition near the 3:8 ratio was also encountered. Although Nb_2O_5 apparently exhibits no solid solution, WO_3 was found to accept a maximum of three mole percent niobia in solid solution enabling all the reported polymorphs of WO_3 to be obtained at room temperature.

Key Words: Crystal structure, niobium pentoxide, phase equilibria, system, tungsten trioxide.

1. Introduction

A complete study of the phase relationships in the binary system $\text{Nb}_2\text{O}_5 - \text{WO}_3$ has been conducted as part of a continuing program of fundamental phase equilibria studies of ceramic materials. This particular system was selected for study in order to further the knowledge of the crystal chemistry of niobates in view of the results obtained from previous structural studies of single crystals [1-4].¹

Due to the complex nature of the x-ray diffraction powder patterns in this system, conflicting interpretations of the data have been reported [5-11]. In a previous study of the $\text{Nb}_2\text{O}_5 - \text{WO}_3$ system, Goldschmidt [5] concluded that Nb_2O_5 could accept more than 50 mole percent WO_3 in solid solution and reported the existence of a compound at approximately $\text{Nb}_2\text{O}_5 : 3\text{WO}_3$. He also found a limited solid solution of Nb_2O_5 in WO_3 . Kovba and Trunov [6] reported that the 1:3 compound was tetragonal with a structure related to that of a tetragonal tungsten-bronze. Fiegel et al. [7], studied the phase equilibria in the system at 1200 °C and below. They reported only about 33 mole percent solid solution of WO_3 in Nb_2O_5 with a 3:2 compound occurring at 1200 °C and a 1:1 compound at 1100 °C and below. They found very little, if any, solid solution of Nb_2O_5 in WO_3 and confirmed the existence of a compound at about 1:3 with a limited solid solution for both Nb_2O_5 and WO_3 .

Later [8] they reported that the 1:3 composition gave single crystal x-ray diffraction patterns which showed superstructure differing from one crystal to the next. Kovba et al. [9], also attempted to determine the phases formed in this system at about 1200 °C. They concluded that compounds existed at the $\text{Nb}_2\text{O}_5 : \text{WO}_3$ ratios of 4:1, 2:1, 4:7, and 1:3. They also reported a phase with variable WO_3 content occurring at high WO_3 concentrations and observed a small amount of solid solution in both Nb_2O_5 and WO_3 .

In a description of preliminary phase identification, Roth and Wadsley [1] on the basis of single crystal x-ray diffraction studies reported the existence of five compounds structurally related to Nb_2O_5 occurring at $\text{Nb}_2\text{O}_5 : \text{WO}_3$ ratios of 15:1, 6:1, 7:3, 8:5, and 9:8. They also confirmed the 1:1 phase and noted the existence of at least three phases related to the tetragonal tungsten bronze-type structure. The crystal structures of 6:1, 7:3, 8:5, and 9:8, were reported [2, 3] and the "building block" principle was elaborated as the basis of the crystal-chemistry of niobate compounds [4].

Schäfer and Gruehn [10] and Felten [11] have indicated in private communications that their interpretations of x-ray diffraction powder data for the $\text{Nb}_2\text{O}_5 - \text{WO}_3$ system differ in some respects from the previously mentioned published reports.

Because of the conflicting nature of the reported data, it was thought desirable to study completely the phase equilibria of the $\text{Nb}_2\text{O}_5 - \text{WO}_3$ system. With the use of the unit cell dimensions derived from

¹ Figures in brackets indicate the literature references at the end of this paper.

single crystal data the existence of the various phases in the powder patterns can now be established more readily. The exact compositions of phases can often be determined only by a solution of the crystal structure. The approximate composition and even the existence of a phase can sometimes be found only by a rather complete phase equilibria study. The two disciplines, crystal structure analysis and phase equilibria are therefore complimentary and no system can be considered to be well characterized unless both such studies have been made. Therefore, x-ray diffraction data together with the melting points of the compounds and solidus and liquidus temperatures at various compositions across the system have been obtained in order to construct an equilibrium diagram.

2. Sample Preparation and Test Methods

The following starting materials were employed for the preparation of the majority of the specimens:

Nb_2O_5 —high purity niobium pentoxide. Spectrographic analysis indicated less than about 0.01 percent Si; 0.001 percent Ca and Mg with As, Cu, and Ta only questionably present.

WO_3 —high purity tungsten anhydride. Spectrographic analysis indicated less than about 0.1 percent Si; 0.001 percent B, Ca, Cr, and Mg; 0.0001 percent Cu with Pb only questionably present.

A few specimens were prepared with less pure starting materials, in order to determine qualitatively the effect of impurities on the equilibrium products. The following is a typical example of the nature of the impurities present in two specimens of $Nb_2O_5:WO_3$ ratio of 16:5. For the less pure specimen the first series transition elements, in general, are present in amounts of an order of magnitude greater than for the more pure specimen.

$Nb_2O_5:WO_3$ (16:5)—higher purity end members. Spectrochemical analyses indicated less than about 0.01 percent Cr, Cu, and Si; 0.001 percent Al, B, Ca, Mg, and Ni; 0.0001 percent Mn with Pb only questionably present.

$Nb_2O_5:WO_3$ (16:5)—less pure end members. Spectrochemical analyses indicated less than about 0.01 percent Al, Ca, Cr, Cu, Fe, Mg, Ni, and Si; 0.001 percent Mn; 0.0001 percent B with Pb only questionably present.

For the preparation of the specimens, the weight percentages were calculated to within ± 0.01 percent, with no corrections made for percentage of purity of the starting materials except for loss on ignition.

For the higher purity specimens, the starting materials were weighed to the nearest ± 0.1 mg, in sufficient quantities to yield 3 g batches. Each batch was mixed in a mechanical shaker for about

15 min and pressed into a disk in a $\frac{3}{8}$ -in. diam mold at 10,000 lb/in². The disks were placed on Pt foil and calcined by heating in air at 700 °C for 19 hrs, with heating and cooling rates of approximately 4 °C/min.

The less pure specimens were weighed out in approximately 1 g batches and mixed with an alumina mortar and pestle. In order to minimize any possible loss of WO_3 these specimens were not calcined after mixing.

Subsolidus, as well as melting point data, were obtained by the quenching technique on samples sealed in platinum tubes. An electrically heated vertical tube furnace wound with 80 percent Pt-20 percent Rh wire was used for the quenching experiments. The furnace was controlled by an a-c Wheatstone bridge controller which was capable of holding the temperature to at least ± 2 °C for an extended period of time. Temperatures were measured with a Pt versus Pt-10 percent Rh thermocouple which had been calibrated against the melting points of Au (1063 °C) and Pd (1552 °C) [12]. The thermocouple was recalibrated several times during the course of the work. Specimens were suspended in the furnace by fine Pt wire. In order to quench the wire was fused allowing the sealed tubes to drop out of the heating chamber into a beaker of water. When the tubes were opened the specimens were examined for physical appearances of melting. The first sign of glazing of the surface of the specimen was interpreted as the first experimental evidence for the solidus temperature. Acceptance of this appearance as evidence of melting was found justified in many specimens by an abrupt difference in the x-ray diffraction powder patterns of the specimens. The formation of a concave meniscus, without the formation of relatively large crystals, indicated the liquidus temperature. The overall reproducibility of the temperature measurements for the experimental data points was within ± 2 °C or better and the overall accuracy of the reported temperatures was within ± 5 °C or better.

Equilibrium was considered to have been obtained when the x-ray diffraction patterns of specimens successively heated for longer times and/or at higher temperatures showed no change. X-ray diffraction powder patterns were made using a high-angle recording Geiger counter diffractometer and nickel-filtered copper radiation, with the Geiger counter traversing the specimen at $1/4^\circ 2\theta/\text{min}$ and the intensity of the radiation being recorded on the chart at $1^\circ 2\theta/\text{in}$. The unit cell dimensions reported can be considered accurate to about ± 2 in the last decimal place listed.

3. Compounds in the $Nb_2O_5-WO_3$ System

3.1. Nb_2O_5

The stability relations of the various reported polymorphs of Nb_2O_5 have been summarized by several workers [13-17]. It has been concluded that the high-temperature monoclinic form of Nb_2O_5 is the only

stable form at atmospheric pressure [14, 16, 17]. The crystal structure of the high-temperature modification of Nb_2O_5 has been described by Gatehouse and Wadsley [18]. They report this phase to be monoclinic, (space group P2) with 14 formula units in the unit cell and $a=21.16 \text{ \AA}$, $b=3.822 \text{ \AA}$, $c=19.35 \text{ \AA}$, $\beta=119^\circ 50'$. The x-ray diffraction powder pattern previously listed for Nb_2O_5 [19] can be more correctly indexed on the basis of the published single-crystal intensity data and is shown in table 1. This indexing leads to slightly revised values of the unit cell dimensions, as follows: $a=21.149 \text{ \AA}$, $b=3.823 \text{ \AA}$, $c=19.352 \text{ \AA}$, $\beta=119^\circ 48'$, in excellent agreement with the lattice constants given by Gatehouse and Wadsley [18]. The melting point of the pure Nb_2O_5 used for this study was previously determined to be 1485°C [20].

TABLE 1. X-ray diffraction powder data for Nb_2O_5 ($CuK\alpha$ radiation)

d^*	l^b	$\frac{1}{d^2}$ obs	$\frac{1}{d^2}$ calc	hkl^c	d^*	l^b	$\frac{1}{d^2}$ obs	$\frac{1}{d^2}$ calc	hkl^c
16.66	6	0.0036	0.0036	001				0.1528	802
10.517	5	0.0090	0.0090	201	2.543	36	0.1546	1.541	102
10.063	2	0.0099	0.0097	101				1.543	414
9.615	2	0.108	0.107	102	2.523	5	1.571	1.571	013
9.148	5	0.120	0.118	200	2.491	26	1.612	1.612	707
8.354	4	0.143	0.142	002	2.478	8	1.629	1.623	511
6.942	2	0.208	0.206	301				1.625	214
6.486	3	0.238	0.236	102	2.452	5	1.664	1.662	416
6.286	11	0.253	0.252	103				1.672	615
5.590	4	0.320	0.319	003	2.514	29	1.865	1.865	413
5.273	6	0.360	0.359	402				1.869	616
5.116	48	0.382	0.382	401	2.114	4	2.244	2.239	413
4.734	4	0.446	0.446	103	2.076	38	2.321	2.321	10, 0, 3
4.616	36	0.469	0.465	104	2.037	34	2.471	2.410	209
3.852	6	0.674	0.670	305	1.912	29	2.736	2.736	020
3.821	6	0.685	0.683	205				2.846	3, 0, 10
3.737	100	0.716	0.714	010	1.873	5	2.852	2.847	11, 0, 3
3.636	100	0.756	0.755	105				2.847	11, 0, 5
3.577	4	0.782	0.782	111	1.856	3	2.903	2.904	208
3.553	4	0.792	0.791	112	1.819	20	3.020	3.020	2, 0, 10
3.515	4	0.809	0.807	603				3.031	11, 0, 8
3.483	100	0.824	0.826	602	1.789	16	3.124	3.115	407
3.406	4	0.862	0.859	604				3.118	421
3.383	4	0.877	0.877	303				3.127	902
3.351	28	0.891	0.886	006	1.765	5	3.211	3.201	12, 0, 5
3.316	5	0.909	0.911	311				3.227	12, 0, 6
3.264	4	0.939	0.936	601				3.231	6, 0, 1
3.247	4	0.949	0.951	310	1.742	23	3.296	3.292	119
3.153	11	1.006	1.008	013	1.727	25	3.353	3.353	13, 0, 4
3.078	6	1.055	1.051	506	1.709	10	3.424	3.422	5, 0, 11
2.994	7	1.116	1.113	106	1.709	10	3.424	3.422	11, 1, 4
2.852	36	1.147	1.146	512	1.652	20	3.493	3.491	11, 1, 6
2.826	88	1.152	1.148	014	1.683	51	3.529	3.530	125
			1.162	513				3.531	3, 1, 10
			1.177	511	1.627	8	3.776	3.776	11, 1, 3
2.771	31	1.303	1.301	215	1.592	27	3.945	3.945	406
2.701	34	1.371	1.377	706	1.582	20	3.994	3.982	7, 1, 11
			1.401	415				3.995	11, 0, 1
2.668	3	1.405	1.405	207	1.579	18	4.012	4.012	318
2.644	4	1.431	1.433	510	1.556	22	4.130	4.013	10, 1, 1
			1.445	804				4.135	12, 1, 8
2.628	5	1.448	1.452	607					

* Interplanar spacing.

^b Intensity relative to the strongest peak(s).

^c Indexed on the basis of a monoclinic unit cell (space group P2) $a=21.149 \text{ \AA}$, $b=3.823 \text{ \AA}$, $c=19.352 \text{ \AA}$, $\beta=119^\circ 48'$ and the previously reported observed structure factors from the single crystal data [18].

3.2. Compounds Structurally Related to Nb_2O_5

a. Compounds Belonging to the Homologous Series $B_{2m+1}O_{3m-(n+m)+4}$

Roth and Wadsley [4] have shown that most of the phases formed by the addition to Nb_2O_5 of oxides with cations similar in size to Nb^{5+} , regardless of valence, can be described by the homologous series notation $B_{2m+1}O_{3m-(n+m)+4}$. Structurally, these phases are made up of blocks of octahedra in a given plane n long and m wide with a variable number (p) of blocks connected by edge sharing of octahedra at the corners of the blocks. Similar block units occur at two levels in the unit cell, zero and one-half, in a

TABLE 2. X-ray diffraction powder data for the compound $WNb_{13}O_{30}$ ($CuK\alpha$ radiation)

d^*	l^b	$\frac{1}{d^2}$ obs	$\frac{1}{d^2}$ calc	hkl^c	d^*	l^b	$\frac{1}{d^2}$ obs	$\frac{1}{d^2}$ calc	hkl^c
14.77	19	0.0046	0.0046	001	2.167	6	0.2138	0.2135	10, 0, 5
11.08	23	0.081	0.081	201				0.213	801
7.42	16	0.182	0.183	002	2.128	5	0.2309	0.2323	008
6.43	8	0.242	0.241	201				0.226	007
5.54	18	0.326	0.325	402	2.116	6	0.2233	0.2237	206
								0.224	517
5.35	6	0.349	0.348	401	2.112	6	0.2232	0.2244	517
5.04	18	0.394	0.394	403	2.072	67	0.2325	0.2328	10, 0, 7
4.937	12	0.410	0.411	003					
4.674	58	0.458	0.458	202	2.042	52	0.2398	0.2401	405
3.743	76	0.714	0.713	110	2.006	6	0.2484	0.2487	912
								0.269	802
3.729	96	0.719	0.718	111	1.9375	9	0.2664	0.2669	805
3.702	40	0.730	0.730	004					
3.697	40	0.732	0.732	603	1.9324	8	0.2678	0.2683	315
3.670	20	0.742	0.742	602	1.9121	60	0.2735	0.2735	020
3.616	284	0.765	0.766	203					
								0.2851	220
3.535	43	0.820	0.818	111	1.8664	5	0.2871	0.2873	220
3.510	300	0.812	0.811	604	1.8560	15	0.2899	0.2903	12, 0, 3
3.437	13	0.847	0.846	601				0.2910	227
3.415	15	0.857	0.857	205	1.8524	29	0.2914	0.2918	022
3.358	29	0.887	0.887	512				0.2920	028
								0.2972	12, 0, 4
3.205	27	0.974	0.975	112	1.8342	8	0.2969	0.2972	12, 0, 4
2.930	16	1.165	1.165	204	1.8144	8	0.3038	0.3040	12, 0, 7
2.901	5	1.189	1.189	512	1.8090	20	0.3064	0.3065	406
2.866	58	1.217	1.218	513	1.8007	8	0.3084	0.3093	209
2.836	37	1.243	1.243	113	1.7895	13	0.3126	0.3129	11, 1, 5
					1.7720	8	0.3185	0.3185	11, 1, 6
2.799	7	1.277	1.279	206					
2.735	41	1.337	1.337	514	1.7697	9	0.3193	0.3193	222
2.706	31	1.365	1.366	312	1.7640	22	0.3214	0.3217	8, 0, 10
					1.7567	19	0.3240	0.3245	12, 0, 8
					1.7403	21	0.3302	0.3305	316
2.682	17	1.390	1.392	805	1.7329	19	0.3330	0.3332	10, 0, 1
2.542	8	1.548	1.548	515	1.7311	18	0.3337	0.3339	11, 1, 7
								0.3500	223
								0.3541	515
2.520	116	1.575	1.574	801	1.6903	20	0.3500	0.3501	223
								0.3547	624
								0.3582	3585
								0.3664	3869
2.499	10	1.602	1.599	607	1.6709	26	0.3582	0.3585	11, 1, 8
					1.6688	21	0.3664	0.3669	10, 0, 2
2.462	16	1.650	1.651	511					
2.410	4	1.722	1.722	712	1.6070	23	0.3872	0.3880	2, 0, 10
2.339	6	1.828	1.831	404	1.5975	16	0.3919	0.3924	912
					1.5838	21	0.3987	0.3992	9, 1, 10
								0.4006	14, 0, 5
2.328	15	1.845	1.848	800	1.5801	20	0.4005	0.4017	10, 0, 11
								0.4049	119
2.316	29	1.864	1.868	516	1.5608	19	0.4105	0.4109	13, 1, 3
								0.4154	14, 0, 4
2.215	11	2.038	2.043	608	1.5327	9	0.4148	0.4154	14, 0, 4
2.183	7	2.096	2.099	710	1.5405	5	0.4214	0.4227	4, 0, 11

* Interplanar spacing.

^b Observed intensity.

^c Indexed on the basis of a monoclinic unit cell (space group C2) $a=22.282 \text{ \AA}$, $b=3.824 \text{ \AA}$, $c=17.734 \text{ \AA}$, $\beta=123^\circ 22'$ and the previously reported structure factors from the single crystal data [2].

plane perpendicular to one axis which always has a value of approximately 3.8 Å. This axis is the *b*-axis of a monoclinic cell or the *c*-axis of a tetragonal cell. The octahedra occurring at the two different levels are connected to each other by edge sharing except at the junction of four such blocks, two at each level, where they are bonded by a tetrahedrally coordinated cation. Those compounds in the Nb₂O₅-WO₃ system whose structures have been described all have *p* = 1 and the formula simplifies to B_{*n**m*+1}O_{3*n**m*-(*n*+*m*)+1}, where the tetrahedral position is always occupied by W⁶⁺ ions. In this arrangement when *n* = *m*, the unit cell is body-centered tetragonal as in PNb₁₀O₂₅ [21]. When *n* ≠ *m* the cell is C-centered monoclinic.

b. 6Nb₂O₅ · 3WO₃ (W₃Nb₁₂O₃₃)

The unit cell of this compound is C-centered monoclinic (probable space group C₂) having *n* = 3, *m* = 4 [2]. The indexed x-ray diffraction powder pattern obtained in the present study is listed in table 2. The unit cell dimensions derived from this pattern are *a* = 22.282 Å, *b* = 3.824 Å, *c* = 17.724 Å, β = 123°22'. WNb₁₂O₃₃ is apparently stable from room temperature to the congruent melting point at 1476 °C.

c. 7Nb₂O₅ · 3WO₃ (W₃Nb₁₄O₃₃)

The unit cell of this compound is body-centered tetragonal, space group I4/m or I4̄(*n* = 4, *m* = 4 [3]). The indexed x-ray diffraction powder pattern is listed in table 3 and the unit cell dimensions derived from this pattern are *a* = 21.002 Å, *c* = 3.820 Å.

d. 8Nb₂O₅ · 5WO₃ (W₅Nb₁₆O₃₅)

The unit cell of this phase is C-centered monoclinic space group C₂ (*n* = 4, *m* = 5, [2]). The indexed x-ray diffraction powder pattern is listed in table 4 and the unit cell dimensions derived from this pattern are *a* = 29.638 Å, *b* = 3.820 Å, *c* = 23.126 Å, β = 126°27'.

TABLE 4. X-ray diffraction powder data for the compound W₅Nb₁₆O₃₅ (CuKα radiation)

<i>d</i> ^a	<i>h</i> ^b	1/ <i>d</i> ² obs	1/ <i>d</i> ² calc	<i>hkl</i> ^c	<i>d</i> ^a	<i>h</i> ^b	1/ <i>d</i> ² obs	1/ <i>d</i> ² calc	<i>hkl</i> ^c
18.58	7	0.0029	0.0029	001	2.397	4	0.1741	0.1744	912
14.84	12	0.0465	0.0466	201	2.382	6	.1763	.1764	711
9.31	13	.0135	.0116	002	2.350	4	.1811	.1811	12,0,8
7.40	17	.0183	.0183	402				.1862	202
6.75	4	.0220	.0220	403	2.316	7	.1864	.1873	8,0,10
6.21	12	.0260	.0260	003	2.294	37	.1900	.1898	911
5.85	6	.0293	.0293	202				.1903	518
4.935	12	.0411	.0411	602				.1968	315
4.701	45	.0453	.0453	604	2.254	6	.1969	.1970	10,0,10
4.652	10	.0462	.0462	004	2.177	6	.2111	.2111	910
4.512	35	.0491	.0491	203	2.104	8	.2260	.2260	519
3.771	130	.0703	.0703	110	2.067	8	.2340	.2340	11,1,2
3.767	142	.0705	.0705	111	2.036	52	.2413	.2413	14,0,9
3.717	26	.0724	.0722	005	2.021	36	.2448	.2448	407
3.705	22	.0729	.0731	804	1.9453	4	.2642	.2638	606
3.660	250	.0747	.0743	803	1.9099	37	.2742	.2742	020
3.630	40	.0759	.0759	204	1.8827	6	.2821	.2823	10,0,3
3.587	222	.0777	.0777	111	1.8783	6	.2835	.2837	12,0,12
3.539	25	.0798	.0799	312	1.8602	17	.2890	.2890	0,0,10
3.506	8	.0813	.0813	022	1.8503	7	.2921	.2919	16,0,7
3.466	8	.0862	.0865	403	1.8329	18	.2977	.2972	16,0,6
3.386	10	.0872	.0872	112	1.8294	26	.2988	.2988	408
3.252	6	.0946	.0947	407	1.8007	12	.3084	.3083	16,0,5
3.188	16	.0984	.0984	513	1.7944	18	.3106	.3107	16,0,10
3.095	12	.1044	.1044	113				.3153	623
3.072	10	.1060	.1061	205	1.7691	7	.3195	.3195	4,0,12
2.980	7	.1126	.1126	800				.3233	223
2.897	8	.1192	.1192	10,0,6	1.7500	25	.3236	.3238	12,0,13
2.832	16	.1240	.1245	712	1.7452	8	.3285	.3285	119
2.817	47	.1260	.1260	714				.3292	12,0,2
2.804	40	.1272	.1274	408	1.7421	14	.3295	.3296	318
2.775	5	.1299	.1300	10,0,7	1.7311	24	.3337	.3336	15,1,9
2.740	54	.1332	.1332	713	1.7229	12	.3369	.3369	517
2.732	65	.1340	.1339	10,0,2	1.6943	29	.3484	.3489	224
2.727	71	.1346	.1345	313				.3515	15,1,10
2.615	8	.1462	.1463	716	1.6862	57	.3517	.3518	825
2.574	16	.1509	.1509	512	1.6303	12	.3763	.3762	12,0,3
2.562	84	.1524	.1520	10,0,1	1.6284	13	.3771	.3772	4,0,13
			.1528	606	1.6110	10	.3857	.3861	14,0,14
2.549	22	.1539	.1538	809	1.5888	12	.3962	.3958	11,1,3
			.1540	405				.4015	13,1,3
2.533	8	.1558	.1559	119	1.5785	14	.4014	.4015	428
			.1627	915				.4041	3,1,12
2.479	6	.1627	.1628	314	1.5732	26	.4041	.4078	17,1,6
			.1699	12,0,2	1.5654	30	.4081	.4081	10,2,2
2.425	11	.1701	.1704	916					

^a Interplaner spacing.

^b Observed intensity.

^c Indexed on the basis of a monoclinic unit cell (space group C₂) *a* = 29.638 Å, *b* = 3.820 Å, *c* = 23.126 Å, β = 126°27' and the previously reported observed structure factors from the single crystal data [2].

TABLE 3. X-ray diffraction powder data for the compound W₃Nb₁₄O₃₃ (CuKα radiation)

<i>d</i> ^a obs	<i>h</i> ^b obs	1/ <i>d</i> ² obs	1/ <i>d</i> ² calc	<i>hkl</i> ^c	<i>d</i> ^a obs	<i>h</i> ^b obs	1/ <i>d</i> ² obs	1/ <i>d</i> ² calc	<i>hkl</i> ^c	<i>d</i> ^a obs	<i>h</i> ^b obs	1/ <i>d</i> ² obs	1/ <i>d</i> ² calc	<i>hkl</i> ^c
14.87	25	0.0045	0.0045	110	2.560	22	0.1526	0.1524	611	1.8620	6	0.2884	0.2884	941
10.52	5	.0090	.0091	200	2.546	122	.1543	.1542	820	1.8564	15	.2902	.2902	880
7.42	33	.0181	.0181	220	2.488	10	.1616	.1615	541	1.8419	28	.2948	.2947	11,3,0,970
6.65	8	.0228	.0227	310	2.441	7	.1679	.1678	750	1.8367	15	.2964	.2967	312
4.951	23	.0408	.0408	330	2.421	16	.1706	.1705	631	1.8007	54	.3084	.3083	10,0,0
4.694	82	.0454	.0453	420	2.358	9	.1799	.1796	701	1.7694	16	.3194	.3194	422
4.118	5	.0590	.0589	510	2.347	8	.1814	.1815	840	1.7546	10	.3248	.3247	871
3.757	142	.0708	.0708	610	2.319	15	.1860	.1859	910	1.7499	40	.3266	.3265	12,0,0
3.712	37	.0726	.0725	440	2.302	41	.1887	.1887	721	1.7305	43	.3319	.3318	961
3.598	480	.0772	.0771	530	2.215	2	.2037	.2040	930	1.6874	65	.3512	.3511	532
3.539	52	.0798	.0799	211	2.152	9	.2159	.2159	811,741	1.6857	85	.3519	.3519	11,2,1,10,5,1
3.321	20	.0907	.0907	620	2.121	3	.2222	.2222	770	1.6612	4	.3624	.3627	12,4,0
3.195	42	.0980	.0980	321	2.100	11	.2258	.2267	10,0,0,860	1.6559	5	.3647	.3647	622
2.968	8	.1135	.1134	710,550	2.067	18	.2341	.2340	831	1.6396	5	.3720	.3718	10,8,0
2.912	2	.1179	.1179	640	2.039	130	.2405	.2408	950	1.6239	4	.3792	.3791	11,4,1
2.826	90	.1252	.1252	501,431	1.9497	3	.2631	.2630	10,4,0	1.6103	25	.3856	.3854	13,1,0,11,7,0
2.756	21	.1317	.1315	730	1.9102	40	.2741	.2743	002	1.5861	40	.3975	.3973	12,1,1,981
2.649	105	.1344	.1343	521	1.9004	10	.2769	.2766	11,1,0	1.5686	62	.4064	.4063	10,7,1
2.623	4	.1463	.1463	800	1.8797	5	.2830	.2831	202					

^a Interplaner spacings.

^b Observed intensities.

^c Indexed on the basis of a tetragonal unit cell (space group I4/m or I4̄) *a* = 21.002 Å, *c* = 3.820 Å.

TABLE 5. X-ray diffraction powder data for the compound $W_5Nb_{16}O_{35}$ ($CuK\alpha$ radiation)

d^*	I^*	$\frac{1}{d^2}$ obs	$\frac{1}{d^2}$ calc	hkl^*	d^*	I^*	$\frac{1}{d^2}$ obs	$\frac{1}{d^2}$ calc	hkl^*	d^*	I^*	$\frac{1}{d^2}$ obs	$\frac{1}{d^2}$ calc	hkl^*
18.43	4	0.0029	0.0029	110	2.731	119	0.1341	0.1340	631	1.9340	4	0.2674	0.2673	11,4,1
9.33	8	.0115	.0116	220	2.426	12	.1450	.1449	10,0,0/860	1.9068	40	.2750	.2750	002
6.20	10	.0260	.0261	330	2.422	18	.1455	.1455	721	1.8578	10	.2898	.2898	14,2,0/10,10,0
5.88	10	.0290	.0290	420	2.375	96	.1508	.1507	10,2,0	1.8482	12	.2928	.2927	11,9,0
4.650	8	.0464	.0464	440	2.522	11	.1373	.1372	651	1.8216	48	.3014	.3014	12,8,0
4.593	80	.0493	.0493	520	2.477	15	.1430	.1429	811,741	1.7791	12	.3160	.3159	13,7,0
4.375	5	.0522	.0522	600	2.458	7	.1744	.1743	831	1.7561	15	.3243	.3243	532
3.771	146	.0703	.0702	101	2.378	9	.1769	.1768	11,1,0	1.7474	26	.3275	.3275	15,1,0
3.712	25	.0726	.0725	710/550	2.304	6	.1885	.1884	11,3,0/970	1.7384	7	.3309	.3310	10,9,1
3.642	372	.0753	.0754	640	2.283	38	.1918	.1919	921/761	1.7235	28	.3366	.3367	14,5,0
3.450	15	.0840	.0840	730	2.253	11	.1970	.1971	10,6,0	1.6943	57	.3484	.3484	15,4,1/11,8,1
3.378	17	.0876	.0876	321	2.248	7	.1978	.1977	851	1.6697	65	.3503	.3504	12,7,1
3.185	12	.0986	.0985	820	2.242	6	.2066	.2067	12,0,0	1.6614	5	.3623	.3624	642
3.088	25	.1049	.1043	660	2.156	9	.2151	.2145	12,2,0	1.6532	6	.3689	.3688	15,5,0/13,9,0
3.054	15	.1072	.1060	501/431	2.100	12	.2267	.2267	10,1,1	1.6292	22	.3768	.3768	14,3,1/13,6,1
2.901	5	.1189	.1188	910	2.077	5	.2319	.2319	12,4,0	1.5801	28	.4005	.4006	16,2,0/14,8,0
2.793	82	.1282	.1282	541	2.015	85	.2463	.2463	13,1,0/11,7,0	1.5688	48	.4063	.4064	15,2,1
2.769	26	.1304	.1304	930	1.9584	5	.2607	.2608	12,6,0					13,2,1

* Interplanar spacing.

* Observed intensity.

* Indexed on the basis of a tetragonal unit cell (space group $I4_1$) $a = 26.270 \text{ \AA}$, $c = 3.814 \text{ \AA}$.

Due to the very large monoclinic cell this powder pattern can only be indexed unambiguously with the aid of single crystal data. By utilizing the published F_{obs} values for $W_5Nb_{16}O_{35}$ [2] it was possible to assign indices to the observed peaks in the powder pattern with reasonable certainty up to about $60^\circ 2\theta$ (Cu radiation). $W_5Nb_{16}O_{35}$ is probably not stable below about $1090^\circ C$ and melts incongruently at about $1385^\circ C$.

e. $9Nb_2O_5 \cdot 8WO_3$ ($W_5Nb_{16}O_{35}$)

The unit cell of the last compound observed in this structural series has $n=5$, $m=5$ [3] and is body-centered tetragonal with the most probable space group $I4_1$. The indexed x-ray diffraction powder pattern is given in table 5 and the unit cell dimensions derived from this pattern are $a = 26.270 \text{ \AA}$, $c = 3.814 \text{ \AA}$. This compound is not stable below about $1265^\circ C$ and melts incongruently at about $1375^\circ C$.

It should be noted that the binary equilibrium stability regions of this structural group of compounds tend to decrease with increasing W^{+6} content. This phenomenon is probably due to the increasing size of the basic "building block" unit, and therefore to the greater energy needed to maintain long range ordering.

f. Other Compounds Structurally Related to Nb_2O_5

In addition to the compounds with structures described by the homologous series formula $B_{2mp+1}O_{8mp+1}$ ($n+m$), $p+4$, the existence of several other phases has been reported, but their structures are, as yet, unknown. Notable among these is the $NbO_{2.482}$ (and $(Ti,Nb)O_{2.482}$) reported by Gruehn and Schäfer [22] which is apparently similar to the 'X' phase reported by Waring and Roth in the system vanadium-oxide-niobium oxide [23]. There are several stable phases in the $Nb_2O_5-WO_3$ system which fall in this category.

g. $13Nb_2O_5 \cdot 4WO_3$ ($W_5Nb_{16}O_{35}$)

Another phase, previously unreported, was found to occur between the known 6:1 and 7:3 compounds. The composition of this phase was deduced by Schäfer (private communication) to be 13:4 based on oxygen analyses. It was postulated that this compound might have an $Nb_2O_5:WO_3$ ratio of 16:5 and the structure would then be related to that of Nb_2O_5 by increasing the size of the basic "building blocks" from 3×4 and 3×5 (reported for Nb_2O_5 [18]) to 4×4 and 4×5 . However, single crystals obtained from a 16:5 composition were found by A. D. Wadsley (private communication) to have a unit cell which is C-centered monoclinic $a = 29.74 \text{ \AA}$, $b = 3.823 \text{ \AA}$, $c = 26.02 \text{ \AA}$, $\beta = 92^\circ 18'$. These values are not in agreement with the primitive space group and approximate unit cell dimensions which can be derived for the postulated 16:5 structure. The partially indexed powder pattern is given in table 6. It cannot be fully indexed without the aid of single crystal intensity data due to the very large size of the unit cell. The structure of this phase, is actually a mixture of blocks of 3×4 and 4×4 occurring in alternate sequence to make an "ordered intergrowth" structure [24]. The composition can be arrived at by adding the homologous series twice (as for Nb_2O_5 [4]) $n=3$, $m=4$, $p=1$ ($B_{13}O_{35}$) plus $n=4$, $m=4$, $p=1$ ($B_{17}O_{44}$) = $B_{30}O_{77}$ ($13Nb_2O_5 \cdot 4WO_3$). Although this compound does not form in short-time experiments (about 1 hr) it can be prepared readily by heating for several days at about $1400^\circ C$. This phase was observed to decompose at about $1435^\circ C$ into $6Nb_2O_5 \cdot WO_3$ and $7Nb_2O_5 \cdot 3WO_3$.

h. " $30Nb_2O_5:WO_3$ "?

Another phase was found to occur in the $Nb_2O_5-WO_3$ system at a ratio of approximately 3.5 mole

* Question marks are used around a composition whenever the phase referred to has not been completely characterized.

TABLE 6. X-ray diffraction powder data for the compound $W_4Nb_{25}O_{77}$ ($CuK\alpha$ radiation)

d^*	I^*	hkl^*	d^*	I^*	hkl^*
14.82	10	200	2.418	10	
12.93	8	002	2.546	7	
7.43	12	400	2.309	22	
6.55	5		2.277	5	
5.19	7		2.113	7	
4.951	14	600	2.067	13	
4.833	15	206	2.035	42	
4.686	60	602	2.042	66	
3.786	18	110	1.912	68	020
3.748	200	(111)	1.857	12	
		(111)			
3.711	40	(007)	1.854	15	
		(800)	1.848	25	
		(112)	1.835	5	
3.657	15	(207)	1.816	5	
		(207)	1.805	33	
3.610	250	802	1.781	16	
3.564	190	207	1.763	15	
		(802)	1.756	19	
3.542	50	(311)	1.744	13	
		(311)	1.736	16	
3.463	10	113	1.731	15	
3.367	10		1.706	5	
3.264	10	114	1.689	25	
3.199	16		1.684	68	
2.966	5		1.675	12	
2.919	15		1.610	25	
2.845	32		1.597	8	
2.832	48		1.592	20	
2.777	10		1.585	18	
2.731	52		1.571	25	
			1.565	38	
2.724	55				
2.529	90				

* Interplanar spacing.

* Observed intensity.

* Partially indexed on the basis of a C-centered monoclinic unit cell with $a=39.74$ Å, $b=3.825$ Å, $c=26.02$ Å, $\beta=92^\circ 18'$ by comparison with the single crystal intensities observed from a film taken with a Weissenberg camera (A. D. Wadley, private communication).

percent WO_3 . This phase is apparently the same as that previously reported to occur at about 15:1, the structure of which could not be verified [1]. The 30:1 ratio is that deduced by Schäfer (private communication). It is apparently composed of some complex packing of the "building blocks" previously described. One possible structure might be $n=3$, $m=4$, $p=2$ ($B_{25}O_{62}$) plus two blocks of $n=3$, $m=4$, $p=1$ ($B_{25}O_{60}$) = $B_{51}O_{128}$ ($25Nb_2O_5 \cdot WO_3$). Another possibility might be $n=3$, $m=4$, $p=2$ ($B_{25}O_{62}$) + $n=3$, $m=5$, $p=1$ ($B_{16}O_{41}$) = $B_{41}O_{105}$ ($20Nb_2O_5 \cdot WO_3$). The exact structure and composition remain to be proven because no good single crystal data is yet available. The "30:1" phase was found to melt incongruently at about 1470 °C. Its unindexed x-ray diffraction powder pattern is listed in table 7.

3.3. Compounds Containing Pentagonal Bipyramidal (Sevenfold) Coordinated Cations

a. $Nb_2O_5 \cdot WO_3$ (WNb_2O_8)

A phase isostructural with Nb_3O_7F [25] might be thought likely to occur in this system, but no such structure has ever been reported in pure oxide systems.

TABLE 7. X-ray diffraction powder data for the composition $30Nb_2O_5 \cdot WO_3$ ($CuK\alpha$ radiation)

d^*	I^*	d^*	I^*
16.20	10	2.435	9
13.93	4	2.376	40
10.78	16	2.212	6
8.96	7	2.100	8
8.10	5	2.076	74
6.37	14	2.041	90
5.36	8	1.999	5
5.10	50	1.956	6
4.797	7	1.913	48
4.645	60	1.871	10
3.742	132	1.866	10
3.638	312	1.857	12
3.579	10	1.825	18
3.556	10	1.819	42
3.493	228	1.796	8
3.354	37	1.791	8
3.173	16	1.782	23
3.093	7	1.769	7
2.984	14	1.746	14
2.845	55	1.743	25
2.836	47	1.731	30
2.763	28	1.716	10
2.707	46	1.693	22
2.693	18	1.684	30
2.647	8	1.680	60
2.614	7	1.623	15
2.597	6	1.603	8
2.538	65	1.595	30
2.503	45	1.581	44
2.476	10	1.558	28
2.458	10	1.548	8

* Interplanar spacing.

* Observed intensity.

Instead a new phase was found with unit cell dimensions related to the tetragonal tungsten bronze-type structures. Single crystals of the 1:1 compound were prepared by decomposing the 9:8 phase (made with the less pure end members) at about 1100 °C. Single crystal precession patterns of this phase made by A. Perloff of the National Bureau of Standards staff showed these crystals to be orthorhombic, probable space groups $Pmab$ or $P2_1ab$. From these data the x-ray diffraction powder pattern given in table 8 was indexed with $a=16.615$ Å, $b=17.616$ Å, $c=3.955$ Å. The compound $Nb_2O_5 \cdot WO_3$ could not be made as a single phase in the quenching experiments. X-ray diffraction powder patterns of these specimens always showed a trace of a metastable bronze-type phase plus some $7Nb_2O_5 \cdot 3WO_3$. However, high temperature x-ray powder patterns showed that the extra phases disappeared quickly above about 900 °C. The 1:1 compound was found to dissociate at about 1115 °C to $8Nb_2O_5 \cdot 5WO_3$ plus a bronze-like phase, probably $4Nb_2O_5 \cdot 9WO_3$.

WNb_2O_8 is related to WTa_2O_8 ($a=16.701$ Å, $b=8.864$ Å, $c=3.877$ Å) by a doubled b -axis. The crystal structure of $LiNb_6O_{15}F$ which is apparently isostructural with WTa_2O_8 was found by S. Andersson and M. Lundberg [26] to be made up of octahedrally coordinated cations. These octahedra are corner shared to form a ring of five octahedra. Within this ring is a cation in sevenfold coordination, in a penta-

TABLE 8. X-ray diffraction powder data for the compound WNb_2O_8 ($CuK\alpha$ radiation)

d^a	P^b	$\frac{1}{d^2}$ obs	$\frac{1}{d^2}$ calc	hkl^c	d^a	P^b	$\frac{1}{d^2}$ obs	$\frac{1}{d^2}$ calc	hkl^c	d^a	P^b	$\frac{1}{d^2}$ obs	$\frac{1}{d^2}$ calc	hkl^c
8.83	6	0.0128	0.0129	020	2.350	8	.1976	.1975	611	1.7546	16	.3248	.3246	851
8.30	25	.0145	.0145	200	2.223	8	.2023	.2024	451	1.7496	20	.3267	.3266	091
4.689	6	.0455	.0455	320	2.203	19	.2061	.2060	541	1.7444	23	.3279	.3281	481
4.403	5	.0516	.0516	040	2.196	19	.2070	.2072	621	1.7267	15	.3562	.3563	052
4.259	12	.0591	.0592	140	2.117	4	.2232	.2233	631				.3366	680
3.952	272	.0640	.0639	001	2.091	8	.2298	.2290	740	1.7154	16	.3398	.3394	291
3.994	46	.0660	.0660	240									.3399	342
3.759	112	.0708	.0708	420	2.078	14	.2316	.2317	800				.3472	841
3.606	4	.0768	.0768	021	2.050	6	.2381	.2379	462	1.6969	40	.3473	.3477	860
3.570	9	.0785	.0785	201	2.023	18	.2443	.2445	711	1.6885	9	.3507	.3508	252
3.449	202	.0841	.0841	340	1.9774	50	.2558	.2557	002	1.6740	7	.3569	.3573	761
3.270	20	.0930	.0929	031	1.9632	7	.2589	.2590	012	1.6684	22	.3593	.3591	522
3.215	8	.0967	.0966	131						1.6662	28	.3602	.3604	911
3.165	22	.0998	.0997	311	1.9473	18	.2637	.2642	480				.3607	301
3.110	98	.1034	.1034	520	1.9247	15	.2699	.2702	081	1.6551	27	.3607	.3607	501
3.051	16	.1075	.1074	231				.2702	202	1.6525	26	.3618	.3620	10,0,0
2.937	4	.1159	.1160	060				.2703	731	1.6498	5	.3632	.3632	442
2.890	80	.1195	.1196	160	1.8797	16	.2830	.2831	222	1.6434	4	.3703	.3701	921
2.825	11	.1253	.1251	411				.2833	840	1.6396	4	.3720	.3717	062
2.771	78	.1303	.1305	331	1.8743	15	.2847	.2846	281				.3749	10,2,0
2.725	50	.1347	.1348	600	1.8521	18	.2915	.2915	032	1.6326	12	.3752	.3752	532
2.654	16	.1420	.1421	240				.2915	312				.3754	162
2.644	18	.1430	.1432	620	1.8475	24	.2930	.2929	741	1.6234	14	.3795	.3802	4,10,0
2.633	12	.1443	.1445	051				.2934	760				.3861	502
					1.8294	15	.2991	.2956	801	1.6096	13	.3860	.3861	0,10,1
2.599	86	.1461	.1481	341	1.8178	6	.3026	.2988	811				.3862	262
2.573	8	.1511	.1509	431	1.8081	6	.3026	.2992	252	1.6029	9	.3892	.3893	612
2.518	8	.1577	.1577	511				.3027	381	1.5928	6	.3942	.3942	452
2.510	8	.1587	.1590	251	1.8017	8	.3061	.3073	042	1.5853	8	.3979	.3978	542
2.445	44	.1673	.1673	521				.3085	821				.3998	632
2.398	7	.1739	.1739	460	1.7908	7	.3118	.3109	142	1.5826	8	.3993	.3992	771
2.375	9	.1773	.1771	351	1.7852	6	.3138	.3123	571				.4005	491
2.334	40	.1835	.1834	531	1.7762	9	.3170	.3157	402	1.5808	10	.4002	.4006	2,10,1
2.293	5	.1901	.1836	161				.3169	412				.4087	941
2.268	23	.1943	.1903	720	1.7637	14	.3215	.3218	242	1.5640	14	.4088	.4093	960
			.1944	261				.3219	751	1.5592	26	.4114	.4116	861
								.3228	0,10,0	1.5326	12	.4257	.4260	10,0,1

^a Interplanar spacing.^b Observed intensity.^c Indexed on the basis of an orthorhombic unit cell (space group $Pnmb$ or $P2_1ab$) $a = 16.615 \text{ \AA}$, $b = 17.616 \text{ \AA}$, $c = 3.955 \text{ \AA}$.

gonal bipyramid configuration. These rings are connected by further corner sharing to form double chains. The doubled cell in WNb_2O_8 is probably due to an alternate puckering of the positions occupied by the sevenfold coordinated ions, but the proof of this must await a complete single crystal structure analysis.

b. $4Nb_2O_5 \cdot 9WO_3(Nb_4W_9O_{37})$

A compound having a general diffraction pattern similar to tetragonal potassium tungsten bronze (K_xWO_3) was reported by Roth and Wadsley [1] to occur at the composition $4Nb_2O_5 \cdot 9WO_3$. Sleight and Magneli [27] described the structure of this phase from a crystal obtained from a 1.3 composition which had been heated to a temperature of 1200–1400 °C. The structure of this crystal was found to be made up of three tetragonal-bronze-like unit cells with four out of the twelve possible five-fold rings being occupied by cations, with oxygens above and below forming pentagonal bipyramid coordination polyhedra. This structure represents the composition $Nb_{16}W_{18}O_{94}$ and has orthorhombic symmetry. The unit cell dimensions found in the present work are $a = 36.692 \text{ \AA}$, $b = 12.191 \text{ \AA}$, $c = 3.945 \text{ \AA}$, and were obtained from

the indexed x-ray diffraction powder pattern given in table 9. This compound was found to be stable from at least 1150 °C to the congruent melting point of about 1380 °C.

The presence of cations in the pentagonal holes of the tetragonal tungsten bronze structure suggests a possible homologous series which can be expressed as:



where n is the number of subcells of the tetragonal bronze type in the true unit cell and m is the number of pentagonal bipyramid polyhedra occupied by a cation. Each bronze-type subcell has only four pentagonal holes, therefore the maximum value of m must be equal to or less than $4n$.

c. $2Nb_2O_5 \cdot 7WO_3(Nb_4W_7O_{31})$

One method of reducing the general formula $B_{10n+m}O_{30n+3m}$ to include the 4:9 compound and to predict a *minimum* number of other phases is to assume $m = 4$. This results in the formula $B_{10n+4}O_{30n+4}$.

TABLE 9. X-ray diffraction powder data for the compound $W_3Nb_3O_{17}$ ($CuK\alpha$ radiation)

d^a	l^b	$\frac{1}{d^2}$ obs	$\frac{1}{d^2}$ calc	hkl^c	d^a	l^b	$\frac{1}{d^2}$ obs	$\frac{1}{d^2}$ calc	hkl^c	d^a	l^b	$\frac{1}{d^2}$ obs	$\frac{1}{d^2}$ calc	hkl^c	
11.59	4	0.0074	0.0075	110	2.835	5	.1244	0.1244	901	1.8671	15	.2869	.2869	222	
10.14	12	.0097	.0097	210			.1248	.1248	031			.2905	.2905	612	
9.15	9	.0119	.0119	400	2.827	9	.1251	.1256	131	1.8553	34	.2905	.2906	322	
8.64	8	.0134	.0134	310				.1275	721				.2919	15,3,1	
			.0186	500	2.799	7	.1277	.1278	231	1.8503	20	.2921	.2926	951	
			.0186	410					911	1.8219	34	.3013	.3012	18,3,0	
6.12	6	.0267	.0267	600	2.783	76	.1515	.1515	331	1.8178	53	.3026	.3024	960	
5.09	11	.0269	.0269	020					640	1.8040	6	.3073	.3072	161	
5.46	57	.0335	.0335	610	2.732	80	.1340	.1339	12,2,0				.3107	622	
5.08	5	.0388	.0388	420	2.727	78	.1344	.1344	640	1.7924	10	.3112	.3116	18,1,1	
4.314	22	.0537	.0537	620	2.684	12	.1388	.1387	821				.3132	361	
					2.643	7	.1432	.1434	531	1.7822	5	.3131	.3172	902	
								.1514	921				.3174	.3176	052
4.081	6	.0601	.0602	900					631	1.7749	6	.3174	.3239	912	
4.064	10	.0605	.0606	030	2.569	77	.1515	.1516	731				.3243	332	
4.044	10	.0612	.0613	150					940	1.7307	12	.3303	.3304	170	
3.945	212	.0643	.0643	001	2.490	6	.1613	.1612	731	1.7360	30	.3318	.3319	18,2,1	
3.867	176	.0669	.0669	910					940				.3332	661	
					2.442	5	.1677	.1678	940	1.7305	60	.3339	.3343	21,1,0	
3.858	195	.0672	.0672	390					141	1.7269	80	.3353	.3353	15,5,0	
			.0709	301	2.416	9	.1713	.1712	12,0,1				.3365	370	
3.754	6	.0710	.0710	911	2.411	10	.1720	.1719	041				.3390	15,4,1	
					2.405	6	.1729	.1727	141				.3394	12,5,1	
3.732	7	.0718	.0717	111	2.405	52	.1780	.1779	12,1,1	1.7238	55	.3365	.3364	370	
3.678	10	.0739	.0740	211	2.370	64	.1787	.1786	341	1.7238	55	.3365	.3364	370	
			.0743	10,0,0	2.366	22	.1850	.1850	931	1.7172	20	.3391	.3394	12,5,1	
3.660	28	.0746	.0746	820	2.325				941				.3441	922	
3.597	4	.0777	.0777	311				.1962	12,2,1				.3443	632	
3.559	14	.0790	.0791	530				.1987	641	1.7045	20	.3442	.3443	632	
3.474	6	.0829	.0829	501	2.245	33	.1984			1.6808	6	.3540	.3545	21,2,0	
				411	2.158	4	.2147	.2146	12,4,0				.3583	5584	
3.388	164	.0871	.0871	920	2.095	10	.2278	.2277	15,5,0	1.6703	7	.3583	.3584	16,5,0	
					2.091	14	.2286	.2284	950	1.6540	32	.3655	.3655	18,3,1	
3.384	166	.0873	.0873	630				.2314	15,0,1	1.6513	40	.3667	.3666	961	
3.315	5	.0909	.0910	601	2.078	10	.2316	.2318	12,2,1				.3707	12,1,2	
3.260	13	.0941	.0942	221				.2321	941	1.6417	24	.3710	.3714	542	
			.0977	511				.2325	051				.3732	932	
3.197	60	.0978	.0979	321	2.072	10	.2330	.2332	151	1.6273	7	.3776	.3777	932	
				10,2,0				.2570	002				.3900	12,2,2	
3.145	7	.1011	.1012	421	1.9725	54	.2570	.2570	002	1.5991	12	.3911	.3914	642	
				421	1.9681	35	.2582	.2583	15,2,1				.3947	171	
3.115	4	.1030	.1031	421	1.9681	35	.2582	.2582	051				.3985	21,1,1	
3.058	20	.1070	.1070	12,0,0	1.9644	12	.2591	.2592		1.5818	46	.3997	.3996	15,5,1	
3.048	25	.1076	.1077	040				.2637	012	1.5921	9	.3946	.3947	171	
3.019	6	.1097	.1098	521	1.9473	4	.2637	.2637	302	1.5841	40	.3985	.3985	21,1,1	
2.964	146	.1138	.1137	12,1,0				.2676	18,2,0	1.5798	42	.4007	.4006	371	
					1.9332	45	.2676	.2676	060				.4089	18,5,0	
2.956	158	.1143	.1143	340	1.9328	46	.2691	.2690	060				.4094	15,6,0	
2.912	17	.1180	.1179	621	1.9278			.2752	12,5,0						
2.878	50	.1208	.1207	950	1.9072	12	.2749	.2752							

^a Interplanar spacing.^b Observed intensity.^c Indexed on the basis of an orthorhombic unit cell with $a = 36.692 \text{ \AA}$, $b = 12.191 \text{ \AA}$, $c = 5.945 \text{ \AA}$.

For the 4:9 compound $n=3$. The composition of $n=2$ is $Nb_2O_5 \cdot WO_3$ and the composition of $n=4$ would be $2Nb_2O_5 \cdot 7WO_3$. The 1:1 compound does not belong to the general homologous series but has a unique structure only vaguely related to the tetragonal tungsten bronze type, as previously described. The structure predicted for $2Nb_2O_5 \cdot 7WO_3$ would be either orthorhombic with one a -axis multiplied by four or tetragonal with both a -axes doubled. The latter structure, which exhibits a higher symmetry than the former, was found to occur for the 2:7 composition between a minimum temperature of about 1245°C and the probable congruent melting point of 1357°C . A crystal structure analysis of this compound is currently being conducted by N. Stephensen [28]. The unit cell dimensions of $Nb_4W_7O_{31}$ are $a = 24.264 \text{ \AA}$, $c = 3.924 \text{ \AA}$, and were obtained from the indexed x-ray diffraction powder pattern listed in table 10.

d. " $6Nb_2O_5 \cdot 11WO_3$ " ($Nb_{12}W_{11}O_{63}$)

Roth and Wadsley [1] reported another compound in this system occurring at a ratio of about $13Nb_2O_5 : 24WO_3$. Kovba and Trunov [9] described a phase to which they assigned the composition $4Nb_2O_5 \cdot 7WO_3$. The only logical ratio, near this composition, which could belong to the general formula $B_{10n+m}O_{30n+m}$ is $6Nb_2O_5 \cdot 11WO_3$. In order to include this composition with the 4:9 and 2:7 compounds in one simplified formula, it is sufficient to postulate a series limited to the case $m=n+1$. The new homologous series formula then becomes $B_{11n+1}O_{31n+1}$. In this case for the 2:7 compound, $n=\infty$ and for $4Nb_2O_5 \cdot 9WO_3$, $n=3$. When $n=2$ the composition is $B_{25}O_{65}$ and the predicted structure has two bronze-like subcells in the unit cell, with three filled pentagonal holes. Such a structure would have a very low symmetry and is apparently not formed

Table 10. X-ray diffraction powder data for the compound $W_7Nb_4O_{31}$ ($CuK\alpha$ radiation)

d^a	h^b	$\frac{1}{d^2}$ obs	$\frac{1}{d^2}$ calc	hkl^c	d^a	h^b	$\frac{1}{d^2}$ obs	$\frac{1}{d^2}$ calc	hkl^c	d^a	h^b	$\frac{1}{d^2}$ obs	$\frac{1}{d^2}$ calc	hkl^c
10.84	10	0.0085	0.0085	210	2.860	31	0.1223	0.1223	660	1.9664	9	0.2671	0.2669	402
8.56	8	.0136	.0136	230	2.838	8	.1242	.1240	830	1.8606	11	.2897	.2896	412
6.06	8	.0272	.0272	400	2.822	8	.1256	.1257	750	1.8447	24	.2939	.2937	422
5.48	35	.0340	.0340	420	2.816	10	.1261	.1261	601	1.8364	11	.2959	.2960	10,6,1
4.508	8	.0493	.0493	520	2.743	98	.1329	.1329	621	1.8189	12	.3023	.3022	502/432
4.289	19	.0544	.0544	440	2.713	102	.1359	.1359	840	1.8087	36	.3057	.3058	12,6,0
4.046	5	.0611	.0612	500	2.660	10	.1414	.1414	631	1.7878	7	.3129	.3129	11,5,1
3.996	5	.0626	.0628	610	2.651	18	.1500	.1499	711/551					942
3.924	194	.0649	.0649	901	2.555	50	.1532	.1533	641	1.7939	8	.3142	.3142	13,4,0/11,8,0
3.834	256	.0679	.0679	620	2.473	8	.1635	.1635	731					622
3.792	25	.0696	.0696	540	2.399	5	.1737	.1737	801	1.7468	20	.3277	.3277	12,4,1
3.735	10	.0717	.0717	291	2.369	50	.1804	.1804	821	1.7252	50	.3368	.3367	
3.617	16	.0764	.0764	630	2.311	10	.1872	.1872	661	1.7160	104	.3396	.3397	14,2,0/10,10,0
3.439	25	.0850	.0849	710/550	2.290	5	.1907	.1906	751	1.6949	15	.3481	.3481	642
3.366	136	.0883	.0883	640	2.232	50	.2008	.2008	841	1.6825	6	.3533	.3533	12,8,0
3.333	8	.0900	.0900	720	2.177	5	.2209	.2208	11,3,0/970	1.6426	30	.3706	.3707	802
3.265	10	.0938	.0938	411	2.138	12	.2308	.2310	10,6,0					12,6,1
3.181	52	.0989	.0989	421	2.064	14	.2347	.2348	10,0,1/861	1.6326	15	.3752	.3753	822
3.052	7	.1074	.1074	501/431	1.9620	42	.2568	.2568	602	1.6177	4	.3822	.3821	662
3.034	18	.1086	.1087	800	1.9182	60	.2718	.2718	12,4,0	1.5896	12	.3958	.3957	842
2.942	146	.1155	.1155	820	1.8667	7	.2786	.2786	11,2,1/10,5,1	1.5720	63	.4047	.4047	14,2,1/10,10,1
2.890	5	.1191	.1193	441					10,8,0	1.5535	7	.4144	.4145	12,10,0

^a Interplanar spacing^b Observed intensity^c Indexed on the basis of a tetragonal unit cell with $a = 24.264 \text{ \AA}$, $c = 3.924 \text{ \AA}$.

in this system. Experimentally the compound " $6Nb_2O_5 \cdot 11WO_3$ " was found to be orthorhombic, probably with the a -axis of the tetragonal bronze subcell tripled. It is stable from about 1210 °C to the apparently congruent melting point of 1378 °C. The x-ray diffraction powder pattern listed in table 11 is indexed with $a = 36.740 \text{ \AA}$, $b = 12.195 \text{ \AA}$, $c = 3.951 \text{ \AA}$. It may be concluded from the size and symmetry of the unit cell, that the homologous series formula $B_{11n+1}O_{31n+1}$ is not correct for this compound. Therefore, " $6Nb_2O_5 \cdot 11WO_3$ " either has a defect structure or belongs to some other structural homologous series, as yet unknown. A single crystal structure analysis is currently under study by N. Stephenson [28].

e. Metastable Tetragonal Bronze-Type Solid Solution (" $3:8$ ")

In addition to the ordered bronze-like phases which have been found in the Nb_2O_5 - WO_3 system, another, apparently disordered, phase has also been reported at about the $Nb_2O_5:3WO_3$ composition [5, 6, 7]. In the present study a tetragonal bronze-type phase without any indication of superstructure in the powder pattern was found to occur from about 72 to 74 mole percent WO_3 in a temperature range from about 1100 to 1250 °C. The unit cell dimensions of this phase were found to vary from about $a = 12.190 \text{ \AA}$, $c = 3.968 \text{ \AA}$ for the composition containing 72 mole percent WO_3 to $a = 12.178 \text{ \AA}$, $c = 3.930 \text{ \AA}$ for the composition containing 75 mole percent WO_3 . It must be concluded that the lower temperature phase is only metastable, because it decomposes at high temperatures into the two ordered compounds, $4Nb_2O_5 \cdot 9WO_3$ and $2Nb_2O_5 \cdot 7WO_3$.

3.4. Compounds Related to the ReO_3 Structure-Type. The Magneli "Shear Phases"

In the temperature interval from about 1270 to 1358 °C, at least two phases have been found to occur with x-ray diffraction powder patterns suggestive of the Magneli "shear phases" [29], which are structurally related to WO_3 . These phases have been found to occur between about 91 and 94 mole percent WO_3 . Due to the difficulty of obtaining equilibrium and the complex nature of the diffraction patterns, it is very difficult to decipher the exact composition of the phases by powder data alone. Although no single crystals have been examined, the best interpretation of the data suggests that there are two equilibrium phases having the $Nb_2O_5:WO_3$ ratios of 1:11 and 1:15. These compositions would correspond to the members $n = 13$ and $n = 17$ of the homologous series B_nO_{3n-1} . It is possible that other structurally related phases are formed in this compositional region which have very little or no thermal stability. Unindexed x-ray diffraction powder patterns for the "1:11" and "1:15" phases are given in tables 12 and 13, respectively.

3.5. Polymorphs of WO_3 and WO_2 Solid Solutions

WO_3 has been reported to occur in many different polymorphs [30-32]. None of the various high- or low-temperature polymorphs have been previously reported to be quenchable to room temperature. However, many of these reported phases have been found at room temperature in the Nb_2O_5 - WO_3 system.

TABLE 11. X-ray diffraction powder data for the phase $6\text{Nb}_2\text{O}_5 \cdot 11\text{WO}_3$ ($\text{CuK}\alpha$ radiation)

d^a	h^b	$\frac{1}{d^2}$ obs	$\frac{1}{d^2}$ calc	hkl^c	d^a	h^b	$\frac{1}{d^2}$ obs	$\frac{1}{d^2}$ calc	hkl^c
11.62	5	0.0074	0.0075	110	2.328	15	1.846	1.846	931
10.18	14	.0097	.0097	210	2.248	25	1.978	1.977	14,2,1
9.16	8	.0119	.0119	400	2.246	22	1.983	1.983	641
8.62	8	.0135	.0134	310	2.097	7	2.273	2.272	15,3,0
5.47	36	.0334	.0334	610	2.095	8	2.282	2.281	950
4.320	16	.0536	.0536	620	2.072	6	2.350	2.329	151
4.070	8	.0604	.0600	900	1.9754	85	2.563	2.563	002
3.952	256	.0640	.0605	080	1.9355	22	2.669	2.669	10,2,0
3.865	140	.0669	.0641	001	1.9290	26	2.688	2.688	660
3.742	7	.0714	.0672	910	1.9087	8	2.745	2.745	12,5,0
3.682	8	.0738	.0707	330	1.8696	7	2.861	2.861	222
3.663	19	.0745	.0707	301	1.8586	13	2.895	2.897	612
3.596	6	.0774	.0715	011	1.8528	15	2.913	2.913	15,3,1
3.562	10	.0798	.0738	211	1.8496	11	2.923	2.922	951
3.388	156	.0871	.0743	10,0,0	1.8299	25	3.006	3.006	10,3,0
3.223	12	.0962	.0820	820	1.8195	32	3.021	3.021	960
3.201	24	.0976	.0775	311	1.7593	29	3.231	3.230	912
3.061	14	.1067	.0920	530	1.7394	22	3.309	3.309	170
3.049	22	.1076	.0969	920	1.7332	33	3.329	3.328	661
2.968	100	.1135	.1076	611	1.7317	35	3.335	3.334	21,1,0
2.959	106	0.1142	.1076	321	1.7284	52	3.348	3.348	15,4,0
2.917	10	.1175	.1076	12,0,0	1.7244	51	3.363	3.362	370
2.880	33	.1205	.1134	040	1.7184	6	3.386	3.384	15,4,1
2.803	5	.1273	.1205	12,1,0	1.7072	22	3.431	3.432	922
2.763	76	.1310	.1273	721	1.6559	15	3.647	3.647	631
2.735	45	.1337	.1308	911	1.6523	25	3.663	3.662	961
2.730	40	.1342	.1318	331	1.6442	22	3.699	3.697	12,1,2
2.687	12	.1385	.1336	12,2,0	1.6294	8	3.766	3.768	932
2.644	5	.1431	.1343	640	1.6012	12	3.901	3.899	12,2,2
2.573	62	.1511	.1510	921	1.5926	5	3.943	3.943	171
2.492	5	.1610	.1513	631	1.5853	25	3.977	3.975	21,1,1
2.421	6	.1706	.1609	131	1.5801	24	4.005	4.002	15,5,1
2.414	7	.1716	.1609	131	1.5654	5	4.081	4.081	18,5,0
2.373	49	.1776	.1717	041	1.5640	7	4.089	4.088	15,6,0
2.368	59	.1783	.1783	341					

^a Interplaner spacing.
^b Observed intensity.
^c Indexed on the basis of an orthorhombic unit cell with $a = 36.740 \text{ \AA}$, $b = 12.195 \text{ \AA}$, $c = 3.951 \text{ \AA}$.

a. Room Temperature Monoclinic Polymorph

A monoclinic polymorph of WO_3 was reported by Tanisaki [32] to have a monoclinic unit cell with $a = 7.30 \text{ \AA}$, $b = 7.53 \text{ \AA}$, $c = 7.68 \text{ \AA}$, $\beta = 90^\circ 54'$ at room temperature. However, the x-ray diffraction powder pattern can be completely indexed with the c -axis equal to one-half that of the true unit cell. The presence of superstructure requiring the doubled cell can apparently only be found with single crystal data. For this reason, the x-ray diffraction powder pattern listed in table 14 has been indexed on the basis of one-half the real c -axis value. The unit cell dimensions obtained from this pattern were found to be $a = 7.299 \text{ \AA}$, $b = 7.535 \text{ \AA}$, $c = 7.688 (3.844) \text{ \AA}$, $\beta = 90^\circ 54'$. With 1 mole percent solid solution of Nb_2O_5 in WO_3 , the parameters are changed to a

$= 7.317 \text{ \AA}$, $b = 7.532 \text{ \AA}$, $c = 7.684 (3.842) \text{ \AA}$, $\beta = 90^\circ 55'$. For pure WO_3 this monoclinic polymorph is apparently stable from about 17°C [32] to about 310°C .

TABLE 12. X-ray diffraction powder data for the composition $\text{Nb}_2\text{O}_5:11\text{WO}_3$ ($\text{CuK}\alpha$ radiation)

d^a	h^b	d^a	h^b
4.638	5	1.927	90
4.308	7	1.910	50
3.968	25	1.855	58
3.850	350	1.845	90
3.726	250	1.812B	18
3.690	350	1.712	32
3.116B	15	1.700	14
2.764B	10	1.677	100
2.677	178	1.664	50
2.670	190	1.569	8
2.579	28	1.541	18
2.202	25	1.530	25
2.190	25	1.493B	10
2.139	12		
1.998B	10		

B = broad.
^a Interplaner spacing.
^b Observed intensity.

TABLE 13. X-ray diffraction powder data for the composition $\text{Nb}_2\text{O}_5:15\text{WO}_3$ ($\text{CuK}\alpha$ radiation)

d^a	h^b	d^a	h^b
3.931	25	1.898	27
3.839	204	1.869	65
3.760	114	1.855	20
3.682	536	1.841	40
3.134	10	1.812B	15
3.089	23	1.799	25
2.763	7	1.711	21
2.676	140	1.695	20
2.647	96	1.671B	65
2.584	18	1.652	15
2.214	14	1.573	8
2.181	30	1.555	10
2.152	12	1.546	15
2.006B	10	1.529	34
1.922	58	1.525	30

B = broad.
^a Interplaner spacing.
^b Observed intensity.

b. Low-Temperature Triclinic Polymorph

According to Tanisaki [32] the room temperature polymorph of WO_3 transforms to a triclinic form at about 17°C , on cooling. However, on reheating there is still some of the triclinic form remaining at room temperature, indicating some hysteresis in the phase transition. It was found in the present study, that the room temperature monoclinic polymorph could be partially transformed to the triclinic modification by grinding the specimen in a mortar and pestle, without lowering the temperature. The unit cell dimensions of the triclinic form were reported [32] as $a = 7.30 \text{ \AA}$, $b = 7.52 \text{ \AA}$, $c = 7.69 \text{ \AA}$, $\alpha = 88^\circ 50'$, $\beta = 90^\circ 55'$, $\gamma = 90^\circ 56'$. The indexed x-ray diffraction

TABLE 14. X-ray diffraction powder data for the room temperature (monoclinic) form of WO_3 ($CuK\alpha$ radiation)

d^a	l^b	$\frac{1}{d^2}$ obs	$\frac{1}{d^2}$ calc	hkl^c	d^a	l^b	$\frac{1}{d^2}$ obs	$\frac{1}{d^2}$ calc	hkl^c
3.840	154	0.0678	0.0677	001	1.8111	45	0.3049	0.3050	112
3.761	160	.0707	.0705	020	1.7977	42	.3094	.3094	112
3.646	268	.0752	.0751	300					
3.419	5	.0856	.0853	011	1.7116	48	.3414	.3413	022
3.344	65	.0894	.0892	120				.3414	202
3.114	70	.1031	.1030	111	1.6914	40	.3495	.3495	041
3.083	50	.1052	.1052	111	1.6894	32	.3504	.3504	202
2.989	132	.1383	.1382	021					
2.667	72	.1407	.1406	210	1.6740	35	.3569	.3569	240
2.679	54	.1447	.1450	201	1.6581	34	.3637	.3636	401
					1.6504	24	.3671	.3672	151
2.620	152	.1457	.1455	220	1.6455	42	.3693	.3694	141
2.533	22	.1559	.1558	121	1.6423	60	.3708	.3708	430
2.514	27	.1582	.1580	121	1.6385	50	.3725	.3725	401
2.176	38	.2112	.2110	221	1.6231	5	.3919	.3918	331
2.154	36	.2156	.2155	221	1.5943	5	.3984	.3985	331
					1.5984	30	.4117	.4119	222
2.103	6	.2262	.2262	031	1.5419	32	.4206	.4206	222
2.045	17	.2395	.2394	320					
2.025	12	.2438	.2439	131	1.5384	27	.4225	.4224	241
2.016	14	.2461	.2461	131	1.5312	10	.4265	.4268	241
1.9955	20	.2511	.2509	311	1.5179	15	.4341	.4341	421
					1.5029	20	.4427	.4430	421
1.9709	12	.2574	.2576	311	1.4981	22	.4456	.4459	132
1.9216	44	.2708	.2708	002					
1.8838	60	.2818	.2818	040					
					1.4899	42	.4505	.4503	132
								.4507	312
1.8246	133	.3004	.3004	400				.4508	340
					1.4679	14	.4641	.4641	312

^a Interplaner spacing.
^b Observed intensity.
^c Indexed on the basis of a monoclinic unit cell with $a = 7.299 \text{ \AA}$, $b = 7.535 \text{ \AA}$, $c = 3.844 \text{ \AA}$ ($\beta = 91^\circ 54'$).

powder pattern of the triclinic polymorph obtained at room temperature from a specimen which had been removed from liquid nitrogen, is listed in table 15. The value of the c -axis was again halved, as the larger value is not necessary to index the powder pattern. According to Tanisaki, this phase is stable from about -40°C to about 17°C . The triclinic polymorph was never observed in any of the Nb_2O_5 - WO_3 solid solutions.

TABLE 15. X-ray diffraction powder data for the triclinic form of WO_3 ($CuK\alpha$ radiation)

d^a	l^b	$\frac{1}{d^2}$ obs	$\frac{1}{d^2}$ calc	hkl^c	d^a	l^b	$\frac{1}{d^2}$ obs	$\frac{1}{d^2}$ calc	hkl^c	d^a	l^b	$\frac{1}{d^2}$ obs	$\frac{1}{d^2}$ calc	hkl^c
3.840	284	0.0678	0.0677	001	2.159	20	0.2145	0.2145	221	1.8040	26	0.3073	0.3068	112
3.760	234	.0707	.0707	020	2.154	53	.2155	.2158	221	1.8007	43	.3084	.3086	112
3.652	350	.0750	.0749	300	2.150	51	.2163	.2163	221	1.7951	26	.3110	.3113	112
3.262	20	.0885	.0885	120	2.120	3	.2224	.2227	031	1.7244	17	.3363	.3360	022
3.322	25	.0906	.0906	130	2.082	5	.2306	.2310	031	1.7113	24	.3415	.3420	202
3.143	42	.1012	.1013	111	2.059	4	.2358	.2360	320	1.7013	23	.3455	.3451	041
3.099	20	.1041	.1042	111	2.046	8	.2380	.2389	131	1.6975	22	.3470	.3470	022
3.085	65	.1051	.1051	111	2.032	5	.2422	.2423	340	1.6897	17	.3503	.3506	041
3.070	42	.1061	.1059	111	2.025	10	.2439	.2439	131	1.6825	13	.3533	.3536	240
2.714	55	.1359	.1358	021	2.011	9	.2474	.2481	311	1.6653	16	.3606	.3608	141
2.667	108	.1407	.1407	201	2.005	12	.2487	.2490	151					
2.660	102	.1413	.1413	021	1.9985	10	.2504	.2503	131					
2.640	80	.1435	.1435	220	1.9844	9	.2540	.2540	311	1.6612	52	.3624	.3620	240
2.632	93	.1443	.1443	201	1.9736	12	.2565	.2564	311	1.6570	31	.3642	.3635	401
2.600	58	.1480	.1477	220	1.9713	14	.2573	.2568	311	1.6537	28	.3657	.3660	320
										1.6502	33	.3672	.3669	141
2.562	13	.1524	.1524	121	1.9216	48	.2708	.2708	002	1.6358	30	.3738	.3736	141
2.527	11	.1567	.1564	121	1.8901	52	.2829	.2829	040					
2.501	15	.1598	.1598	121				.2995	400					
2.500	17	.1600	.1600	121	1.8273	148	.2995	.2995	140	1.6337	35	.3747	.3744	420
2.202	20	.2063	.2066	221	1.8198	44	.3020	.3021	112	1.6303	18	.3763	.3760	141

^a Interplaner spacing.
^b Observed intensity.
^c Indexed on the basis of a triclinic unit cell with $a = 7.30 \text{ \AA}$, $b = 7.52 \text{ \AA}$, $c = 3.845 \text{ \AA}$ ($\beta = 7.69^\circ$, $\alpha = 88^\circ 50'$, $\beta = 90^\circ 55'$, $\gamma = 90^\circ 56'$).

c. Lowest Temperature Polymorph (Monoclinic)

From about -40°C to below liquid nitrogen temperature, another monoclinic polymorph occurs in pure WO_3 which was reported by Tapisaki [32] to have $a = 5.27 \text{ \AA}$, $b = 5.16 \text{ \AA}$, $c = 7.67 \text{ \AA}$, $\beta = 91^\circ 43'$. This polymorph can be obtained essentially single phase at room temperature by quenching a solid solution of $2Nb_2O_5 : 98WO_3$ from the temperature interval 1230°C to about 1385°C , the solidus temperature. This polymorph also occurs as a mixture with other polymorphs in specimens containing 1 and 3 mole percent Nb_2O_5 . The unit cell dimensions obtained from the powder pattern for the 2 mole percent Nb_2O_5 specimen are $a = 5.305 \text{ \AA}$, $b = 5.192 \text{ \AA}$, $c = 7.671 \text{ \AA}$, $\beta = 91^\circ 33'$. The indexed x-ray diffraction powder pattern is given in table 16, utilizing the 7.671 \AA value for c as several peaks necessitating this doubled value are observed in the powder pattern. High-temperature x-ray patterns show this phase to be apparently stable in the 2 mole percent Nb_2O_5 composition up to a temperature of about 735°C , at which temperature it transforms to a tetragonal polymorph. If the temperature is not raised above about 750°C , this transition is reversible. However, if the temperature is raised to 900°C the low-temperature monoclinic polymorph is not recovered and the tetragonal phase transforms to the orthorhombic polymorph.

Tungsten trioxide has been reported to be ferroelectric below about -40°C [33]. Specimens of sintered WO_3 containing 2 to 4 mole percent Ta_2O_5 were reported to exhibit ferroelectric behavior at room temperature [34]. The low-temperature monoclinic polymorph formed by quenching a specimen containing 2 mole percent Nb_2O_5 was examined for evidence of ferroelectricity [35]. However, no conclusive ferroelectric properties could be found.

TABLE 16. X-ray diffraction powder data for the low-temperature monoclinic form of WO_3 taken from $2Nb_2O_5 \cdot 98WO_3$ solid solution ($CuK\alpha$ radiation)

d^a	I^b	$\frac{1}{d^2}$ obs	$\frac{1}{d^2}$ calc	hkl^c	d^a	I^b	$\frac{1}{d^2}$ obs	$\frac{1}{d^2}$ calc	hkl^c
3.824	408	0.0680	0.0680	002	1.7990	54	0.3090	0.3090	014
3.703	388	.0729	.0729	110	1.7875	29	.3130	.3130	104
3.145	25	.1011	.1011	102	1.7163	37	.3395	.3394	114
3.083	50	.1052	.1051	012	1.6900	25	.3501	.3501	114
3.067	32	.1063	.1064	102	1.6805	37	.3541	.3542	222
2.691	138	.1381	.1382	112	1.6692	41	.3589	.3592	310
2.641	144	.1433	.1432	200	1.6557	28	.3648	.3648	222
2.594	40	.1486	.1484	020	1.6448	26	.3697	.3696	130
2.458	6	.1655	.1654	021	1.6179	4	.3820	.3821	302
2.294	10	.1900	.1900	013	1.5851	9	.3980	.3981	302
					1.5776	5	.4018	.4018	032
2.236	5	.2000	.1998	121	1.5729	10	.4042	.4044	206
2.206	22	.2056	.2058	211	1.5644	18	.4186	.4182	312
2.148	70	.2167	.2163	022	1.5432	21	.4285	.4282	024
2.120	12	.2429	.2429	212	1.5326	6	.4257	.4257	204
2.002	14	.2496	.2495	122	1.5101	28	.4351	.4349	132
								.4352	312
1.9860	17	.2535	.2536	212	1.5056	26	.4412	.4412	132
1.9815	18	.2547	.2548	122	1.4995	13	.4508	.4507	214
1.9178	68	.2719	.2719	004	1.4700	24	.4621	.4614	124
1.8521	100	.2915	.2915	220				.4628	214
1.8192	27	.3022	.3023	104					

^a Interplaner spacing

^b Observed intensity

^c Indexed on the basis of a monoclinic unit cell with $a = 5.305 \text{ \AA}$, $b = 5.192 \text{ \AA}$, $c = 7.671 \text{ \AA}$, $\beta = 91^\circ 33'$.

d. High-Temperature Orthorhombic Polymorph

Although no thermal effect was indicated with DTA, the room-temperature monoclinic polymorph of pure WO_3 is observed in the high temperature x-ray furnace to transform reversibly at 310°C to an orthorhombic phase in agreement with Wyart and Foex [31]. This orthorhombic form is observed at room temperature in the solid solutions whenever the low-temperature monoclinic polymorph is heated to about 900°C and cooled by removing from the furnace. The indexed x-ray diffraction powder pattern of a specimen of this polymorph containing 2 mole percent Nb_2O_5 is listed in table 17. It shows no indication of a doubled c -axis and is indexed on the basis of the smaller orthorhombic cell. The unit cell dimensions obtained for this composition, at room temperature, are $a = 7.384 \text{ \AA}$, $b = 7.512 \text{ \AA}$, $c = 7.692 (3.846) \text{ \AA}$. The orthorhombic phase is observed in this composition to transform to tetragonal in the high temperature x-ray furnace at 510°C and the reverse transition was observed to take place at 440°C .

e. Tetragonal High-Temperature Polymorph(s)

In pure WO_3 the orthorhombic polymorph was found to undergo a transition to the tetragonal form at about 750°C in the high-temperature x-ray furnace. The DTA data indicated that the phase transition occurred at a temperature of 740°C on heating and 730°C on cooling. These results are in reasonable agreement with those of previous workers [7, 31]. The tetragonal polymorph is found as a single phase as low as 440°C in the specimen containing 2 mole

percent Nb_2O_5 . However, in the 3 mole percent Nb_2O_5 specimen, the low-temperature monoclinic polymorph transforms after heating at about 900°C into a mixture consisting predominantly of a tetragonal modification with only a trace of the orthorhombic form. The unit cell dimensions obtained at room temperature from this composition are $a = 5.265 \text{ \AA}$, $c = 3.846 \text{ \AA}$, and the x-ray diffraction powder pattern from which the data were obtained, indexed on the basis of the smaller cell, is listed in table 18.

In addition to the transformation at $730\text{--}740^\circ\text{C}$, another transition is observed at about 900°C in pure WO_3 , as indicated by a very small peak in the DTA pattern. High-temperature x-ray diffraction powder patterns indicate that WO_3 is tetragonal both above and below this minor thermal discontinuity. It is possible that the unit cell has a doubled c -axis below 900°C and only above this temperature does the powder pattern yield the correct unit cell. However, this hypothesis cannot be verified without single crystal data.

Although WO_3 might be expected to have a cubic polymorph of the ReO_3 -type, there is no evidence in either the present work or previously reported studies that such a polymorph exists.

TABLE 17. X-ray diffraction powder data for the orthorhombic form of WO_3 taken from $2Nb_2O_5 \cdot 98WO_3$ solid solution ($CuK\alpha$ radiation)

d^a	I^b	$\frac{1}{d^2}$ obs	$\frac{1}{d^2}$ calc	hkl^c
3.845	248	0.0676	0.0676	001
3.754	156	.0740	.0709	020
3.691	232	.0734	.0734	200
3.427	5	.0852	.0853	011
3.346	18	.0893	.0892	120
3.104	60	.1038	.1037	111
2.986	35	.1086	.1085	021
2.862	80	.1141	.1109	201
2.633	126	.1343	.1342	220
2.525	10	.1568	.1568	121
2.173	42	.2119	.2118	221
2.099	5	.2270	.2271	031
2.056	5	.2365	.2359	320
2.018	20	.2455	.2454	131
1.9976	14	.2506	.2504	311
1.9232	48	.2704	.2704	002
1.8779	33	.2836	.2836	040
1.8461	46	.2934	.2934	400
1.8198	21	.3020	.3019	140
1.8064	37	.3065	.3064	112
1.7119	24	.3412	.3412	022
1.7057	25	.3437	.3437	202
1.6879	28	.3510	.3512	041
1.6740	26	.3569	.3569	240
1.6645	32	.3609	.3610	401
1.6568	34	.3643	.3643	420
1.6453	14	.3694	.3695	141
1.5968	5	.3922	.3922	331
1.5530	10	.4147	.4146	222
1.5349	17	.4244	.4245	241
1.5217	20	.4319	.4319	421
1.4933	16	.4484	.4482	132
1.4858	13	.4530	.4531	340
				312

^a Interplaner spacing

^b Observed intensity

^c Indexed on the basis of an orthorhombic unit cell with $a = 7.384 \text{ \AA}$, $b = 7.512 \text{ \AA}$, $c = 3.846 \text{ \AA}$ ($\neq 7.692 \text{ \AA}$).

TABLE 18. X-ray diffraction powder data for the tetragonal form of WO_3 taken from $3\text{Nb}_2\text{O}_5 \cdot 97\text{WO}_3$ solid solution ($\text{CuK}\alpha$ radiation)

d^a	f^b	$\frac{1}{d^2}$ obs	$\frac{1}{d^2}$ calc	hkl^c
3.847	280	0.0676	0.0676	001
3.725	252	.0721	.0722	110
3.106	58	.1037	.1037	101
2.674	116	.1396	.1396	111
2.658	96	.1438	.1443	200
2.173	40	.2118	.2119	201
2.008	25	.2481	.2480	211
1.923	46	.2705	.2705	002
1.861	48	.2896	.2896	220
1.806	42	.3067	.3066	102
1.707	26	.3430	.3426	112
1.675	40	.3566	.3563	201
1.665	48	.3606	.3608	310
1.597	5	.3922	.3923	301
1.552	8	.4150	.4148	202
1.528	28	.4284	.4284	311
1.489	15	.4510	.4509	212

^a Interplanar spacing.

^b Observed intensity.

^c Indexed on the basis of a tetragonal unit cell with $a = 5.265 \text{ \AA}$, $c = 3.846 \text{ \AA}$.

4. Discussion of Phase Equilibria

The postulated phase equilibrium diagram of the binary system $\text{Nb}_2\text{O}_5\text{-WO}_3$ is shown in figure 1. The experimental data from which this diagram was constructed are given in table 19. The system contains four compounds which are shown as melting congruently, $6\text{Nb}_2\text{O}_5 \cdot \text{WO}_3$, " $6\text{Nb}_2\text{O}_5 \cdot 11\text{WO}_3$," $4\text{Nb}_2\text{O}_5 \cdot 9\text{WO}_3$, and $2\text{Nb}_2\text{O}_5 \cdot 7\text{WO}_3$. Six compounds melt incongruently " $30\text{Nb}_2\text{O}_5 \cdot \text{WO}_3$," $7\text{Nb}_2\text{O}_5 \cdot 3\text{WO}_3$, $8\text{Nb}_2\text{O}_5 \cdot 5\text{WO}_3$, $9\text{Nb}_2\text{O}_5 \cdot 8\text{WO}_3$, and the two Magneli phases estimated to be " $\text{Nb}_2\text{O}_5 \cdot 11\text{WO}_3$ " and " $\text{Nb}_2\text{O}_5 \cdot 15\text{WO}_3$." In addition, two stable phases, $13\text{Nb}_2\text{O}_5 \cdot 4\text{WO}_3$ and $\text{Nb}_2\text{O}_5 \cdot \text{WO}_3$ dissociate before melting as does the apparently metastable disordered bronze type solid solution labeled " $3:8$ " in the phase diagram. Six of the compounds ($8:5$, $9:8$, " $6:11$ ", $2:7$, " $1:11$," and " $1:15$ ") are shown on the phase diagram as having minimum stability temperatures. In many of these cases, it is impossible to determine whether or not the minimum temperature represents a stable transition. The existence of solid solution could neither be definitely established in any of the compounds nor in Nb_2O_5 . However, WO_3 accepts a maximum of about 3 mole percent Nb_2O_5 in solid solution.

The solidus temperature between the " $30:1$ " phase and $6\text{Nb}_2\text{O}_5 \cdot \text{WO}_3$ has been found to be about 1464°C . The exact eutectic composition has not been determined but probably exists between about 7 and 10 mole percent WO_3 . The peritectic corresponding to the incongruent melting point of $7\text{Nb}_2\text{O}_5 \cdot 3\text{WO}_3$ occurs at 1440°C and about 40 mole percent WO_3 . However, the composition of the two peritectics at 1385 and 1375°C and of the eutectic at 1364°C , were not determined exactly. They probably occur between about 60 and 63 mole percent WO_3 .

Both the $8:5$ and $9:8$ compounds apparently have minimum temperatures of stability. However, all efforts to locate accurately the temperature of these

minimums proved inconclusive (table 19). For both compounds, the experimental temperature at which the phase begins to form on heating and the temperature at which it begins to dissociate on cooling had a wide range of overlap. Furthermore, the actual temperature of the minimum stability seemed to be strongly dependent on the purity content (table 19, parts I and II). For these reasons the minimum stability temperatures of the $8:5$ and $9:8$ are indicated on the phase diagram by dashed lines and the temperatures assigned to each, ~ 1090 and $\sim 1265^\circ\text{C}$, respectively, represent the best compromise between the inconsistencies in the experimental data.

None of the ordered bronze-type phases can be prepared in reasonable laboratory time at 1100°C . The " $6:11$ " phase was not found to be well crystallized below about 1210°C . The minimum line at this temperature is dashed to indicate uncertainty in both the temperature value and the stability relation. The same discussion also is relative for the minimum value of the $2:7$. In this case, however, the superstructure lines characteristic of this phase do not begin to appear below about 1245°C . The three ordered bronze-type phases, " $6:11$ ", $4:9$, and $2:7$, all appear to melt congruently at 1378 , 1380 and 1357°C , respectively. The solidus temperatures between these phases at 1365 and 1335°C appear to be measurably lower than the observed melting temperature of the compounds. However, no eutectic compositions could be experimentally determined, possibly due to reduction and, therefore, the liquidus curves are necessarily dashed (see discussion on reduction in sec. 5).

The ability to distinguish a two-phase region between the two bronze-type phases $4\text{Nb}_2\text{O}_5 \cdot 9\text{WO}_3$ and $2\text{Nb}_2\text{O}_5 \cdot 7\text{WO}_3$ was greatly increased by a separation of those phases in the experiments of long-time duration. The top of the specimens within the two-phase region showed only $4:9$ in the x-ray diffraction pattern while the bottom of the specimen showed mostly $2:7$. This apparent gravity separation is probably enhanced by an appreciable vapor phase in the sealed Pt tube and is considered to be due mainly to vapor transport. A similar gravity separation takes place throughout most of the system and adds more evidence to the probability of a two-phase region between " $6\text{Nb}_2\text{O}_5 \cdot 11\text{WO}_3$ " and $4\text{Nb}_2\text{O}_5 \cdot 9\text{WO}_3$. This two-phase region is based mostly on the different morphology and color of these phases, as well as the slight differences in the superstructure exhibited by single crystal patterns. Almost all of the high-temperature phases in the $\text{Nb}_2\text{O}_5\text{-WO}_3$ system are needle formers. Most of those which contain appreciable WO_3 are yellow-green or blue-green in color. However, the " $6:11$ " phase is white and forms tabular crystals rather than needles.

A eutectic exists between the $2:7$ compound and the Magneli "shear" phases at about 1340°C and 83 mole percent WO_3 . However, the exact compositions of the peritectics at 1356 and 1358°C , corresponding to the decomposition temperatures have not been determined.

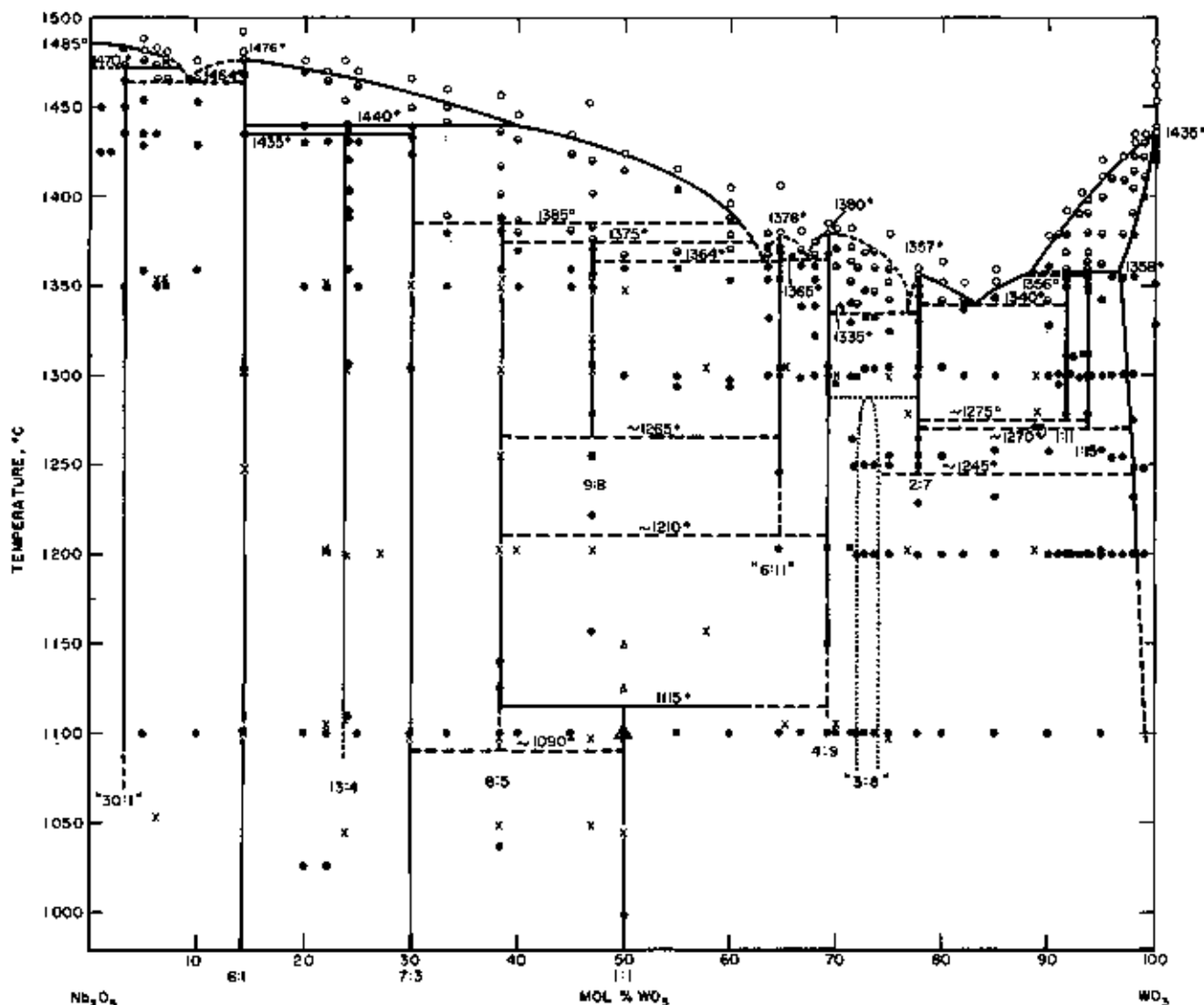


FIGURE 1. Postulated phase equilibrium diagram of the binary system $Nb_2O_5-WO_3$.

Heavy lines represent well-established portions of the system and dashed lines represent not as well-established portions. Dotted lines represent persistent metastable equilibria.
 ●—not melted.
 ◐—partially melted.
 ○—completely melted.
 ×—experimental data obtained from "less-pure" materials.
 △—high-temperature x-ray data.

Tungsten trioxide has been found to accept a maximum of about 3 mole percent Nb_2O_5 in solid solution at the solidus temperature of 1358 °C. The amount of solid solution decreases with decreasing temperature to about 2 mole percent at 1230 °C and less than 1 mole percent at 1100 °C. The melting point of WO_3 was found, in the present study, to be 1435 °C. This value was independently confirmed by E. M. Levin [36] and is in disagreement with the previously published value of 1472 °C (Jaeger and Germs) [37]. In a discussion of the 1472 °C value for the melting point of WO_3 , Phillips and Chang [38] concluded that this temperature was probably too low and might actually represent the eutectic between WO_3 and $W_{20}O_{58}$. However, they did not attempt to experi-

mentally determine the melting points of the compounds WO_3 and $W_{20}O_{58}$ or the solidus temperature between them. The 1435 °C value for the melting point of WO_3 is internally consistent with the experimental phase diagram of both the $Nb_2O_5-WO_3$ and $WO_3-B_2O_3$ systems [36]. This value apparently represents the practical melting point for the experimental conditions utilized: sealed Pt tube, heated for very short intervals of time (≤ 5 min). This method is preferable to that of Jaeger and Germs [37] who utilized an open Pt crucible and much longer heating time. However, the true melting point of WO_3 can probably only be established by utilizing an oxygen pressure furnace and extrapolating to one atmosphere pressure.

TABLE 19. Experimental data for compositions in the binary system Nb₂O₅-WO₃. Part I

Composition		Heat treatments				Results	
		Initial ^a		Final ^b			
Nb ₂ O ₅	WO ₃	Temp.	Time	Temp.	Time		
Mole %	Mole %	°C	Hour	°C	Hour		
99	1			1425	65	No melting	Nb ₂ O ₅
				1450	66	do.	Nb ₂ O ₅
98	2			1425	65	No melting	Nb ₂ O ₅ + "30:1"
96.77	3.23	1350	2			No melting	
	(30:1)			1435	46	do.	"30:1"
				1450	66	do.	"30:1"
				1465	1	No melting (?)	"30:1"
				1474	0.5	Partially melted	"30:1"
				1483	.5	do.	"30:1"
95	5	1359	2	1100 ^d	60	No melting	"30:1" + Nb ₂ O ₅ (?) + 6:1 (?)
				1429	88	do.	"30:1" + 6:1 (?)
				1435	46	do.	"30:1" + 6:1 (?)
				1454	1	do.	"30:1" + 6:1 (?)
				1476	1	Partially melted	"30:1" + 6:1 (?)
				1481	1	Completely melted	"30:1" + 6:1
				1488	0.833	do.	Low Nb ₂ O ₅
93.75	6.25	1550	2				
	(15:1)			1435	46	No melting	"30:1" + 6:1
				1468	0.5	Partially melted	"30:1" + 6:1
				1474	.5	do.	"30:1" + 6:1
				1483	.5	Completely melted	
92.86	7.14	1350	2				
	(13:1)			1466	0.5	Partially melted	6:1 + "30:1" (?)
				1473	.117	do.	
				1476	.5	do.	6:1 (+ unknown)
				1478	15.5	do.	6:1
				1481	0.5	Completely melted	
90	10	1359	2	1100 ^e	60	No melting	6:1 + "30:1"
				1429	88	do.	6:1 + "30:1" (?)
				1453	1	do.	6:1 + "30:1" (?)
				1466	1	Partially melted	
				1476	1	Completely melted	6:1 + low Nb ₂ O ₅
85.71	14.29	1304	2	1100 ^e	60	No melting	6:1
	(6:1)			1435	0.25	do.	6:1
				1467	.25	do.	6:1
				1476	.5	Partially melted	
				1481	.5	Completely melted	6:1 + low Nb ₂ O ₅
				1492	.25	do.	Low Nb ₂ O ₅
80	20			1026	136	No melting	6:1 + 7:3 (+ unknown trf.)
		1550	2	1100 ^d	60	do.	6:1 + 7:3
				1102	65	do.	6:1 + 7:3 + 13:4 (+ unknown trf.)
				1430	1	do.	6:1 + 7:3 + 13:4 (trf.)
				1431	17	do.	6:1 + 13:4 + 7:3 (trf.)
				1440	1	do.	6:1 + 7:3
				1470	0.5	Partially melted	
				1476	.5	Completely melted	
77.78	22.22			1026	136	No melting	6:1 + 7:3 (+ unknown trf.)
	(7:2)	1350	2	1100 ^d	60	do.	6:1 + 7:3 + 13:4 (+ unknown trf.)
				1102	65	do.	6:1 + 7:3 + unknown + 13:4 (trf.)
				1202	30	do.	6:1 + 7:3 (+ unknown trf.)
				1431	0.5	do.	13:4 + 6:1 + 7:3
	7:2 (lean)			1465	.5	Partially melted	
				1470	.5	Completely melted	
76.19	23.81			1109	2	No melting	6:1 + 7:3 + unknown ^e
	(16:5)	1360	2	1110	18	do.	7:3 + unknown + 6:1 (?)
				1307	1	do.	6:1 + 7:3
				1589	119	do.	
				1593	88	do.	13:4
				1404	0.25	do.	
				1421	.25	do.	
				1431	17	do.	13:4 + 6:1 + 7:3 (?)
				1452	0.5	Partially melted	
				1476	.25	Completely melted	
		1360	2				
		1389	119			No melting	13:4
		900	0.33			do.	13:4
				1435	1.25	do.	13:4 + 7:3 + 6:1 (?)
				1439	17	do.	13:4 + 7:3 + 6:1 (?)
				1440	1	do.	13:4 + 6:1 + 7:3 (?)
				1454	1	Partially melted	13:4 + 6:1 + 7:3 (?)
75	25			1100 ^d	60	No melting	7:3 + 6:1 + 13:4
		1550	2	1102	65	do.	7:3 + 6:1 + 13:4 + unknown ^e
				1431	17	do.	13:4 + 7:3 (?)
				1462	0.5	Partially melted	
				1470	.5	Completely melted	
70	30			1100 ^d	60	No melting	7:3 + "leucite" (trf.)
	(17:3)	1304	2			do.	7:3
				1423	1	do.	

TABLE 19. Experimental data for compositions in the binary system Nb₂O₅-WO₃. Part I—Continued.

Composition		Heat treatments				Results	
Nb ₂ O ₅	WO ₃	Initial ^a		Final ^b		Physical observation	X-ray diffraction analysis ^c
		Temp.	Time	Temp.	Time		
Mole %	Mole %	°C	Hour	°C	Hour		
66.67	33.33	1350	2	1434	1	do	7:3
				1439	17	do	7:3
				1449	0.503	Partially melted	
				1466	.33	Completely melted	
				1100 ^d	60	No melting	7:3 + 1:1 + "bronze" (tr) ^e
				1381	1	do	7:3 + 8:5
				1389	1	Partially melted (?)	
				1442	0.5	Partially melted	
				1450	.5	do	
				1461	.5	Completely melted	
61.54	38.46	1350	2	1037	286	No melting	7:3 + 1:1
				1100 ^d	60	do	7:3 + 1:1 + "bronze" (tr) ^e
				1100	167	do	7:3 + 1:1 + 8:5 ^e
				1125	160	do	8:5 + 1:1 + 7:3 ^e
				1140	65	do	8:5 + 1:1 + 7:3 ^e
				1037	236	do	
				1100	167	do	8:5 + 1:1 (tr) ^e
				1125	160	do	8:5 + 1:1 (tr) ^e
				1140	65	do	
				1359	2	do	8:5
60	40	1350	2	1381	2	do	8:5 + 7:3 ^f
				1388	2	Partially melted (?)	7:3 + 8:5 (tr)
				1402	2	do	7:3
				1417	1	do	
				1437	1	do	
				1437	0.5	Completely melted	
				1100 ^d	60	No melting	7:3 + 1:1 + "bronze" (tr) ^e
				1371	1	do	8:5 + 9:8 (tr)
				1381	1	Partially melted (?)	8:5
				1387	1	Partially melted	
55	45	1350	2	1433	1	do	
				1446	0.5	Completely melted	
				1100 ^d	60	No melting	1:1 + 7:3 + "bronze" (tr) ^e
				1360	0.67	do	
				1381	.5	Partially melted (?)	9:8 + 8:5
				1424	.5	Partially melted	
				1435	.5	Completely melted	
				1157	89	No melting	8:5 + 1:1 + "bronze" ^e
				1222	168	do	9:8 + 8:5 + "6:11" ^e
				1255	89	do	9:8 + 8:5 + "6:11" ^e
52.94	47.06	1350	2	1279	64	do	9:8 + 8:5 + "6:11" ^e
				1307	1	do	9:8 + 8:5 + "6:11" ^e
				1350	1	do	9:8 + 8:5 (tr)
				1351	2	do	9:8 + 8:5 (tr)
				1351	18	do	9:8 + 8:5 (tr)
				1222	168	No melting	
				1255	89	do	9:8 + 8:5 + "6:11" ^e
				1279	64	do	9:8 + 8:5 + "6:11" ^e
				1371	1	do	9:8 + 8:5 (tr)
				1377	1	Partially melted (?)	9:8 + 8:5
50	50	1300	2	1383	1	Partially melted	9:8 + 8:5
				1402	0.25	do	8:5 + Q - liq ^f
				1422	.25	do	
				1452	.25	Completely melted	Poorly crystalline 8:5 ^f
				998	144	No melting	1:1 + 7:3 + "bronze" ^e
				1100 ^d	60	do	1:1 + 7:3 + "bronze" ^e
				1361	0.5	do	9:8 + "6:11" ^e
				1368	.67	Partially melted (?)	
				1414	.5	Partially melted	
				1424	.5	Completely melted	
45	55	1300	2	1100 ^d	60	No melting	1:1 + "bronze" ^e
				1294	65	do	9:8 + "6:11" ^e
				1404	0.5	Partially melted	
				1415	.503	Completely melted	9:8 + Q - liq ^f
				1260	0.5	do	9:8 + "6:11" ^e
				1370	.5	Partially melted (?)	
				1100 ^d	60	No melting	1:1 + "bronze" ^e
				1294	65	do	"6:11" + 9:8
				1405	0.5	Completely melted	Q - liq + "bronze"
				1353	1	No melting	"6:11" + 9:8 (7) + 8:5 (tr) ^e
40	60	1298	2	1379	1	do	"6:11" + 9:8
				1388	1	Partially melted	"6:11" + 9:8
				1379	1	do	9:8 + "6:11" ^e + 8:5 tr + Q - liq ^f
				1388	1	do	Q - liq + 9:8 (7) ^e
				1396	0.833	Completely melted	"bronze" + Q - liq
				1360	0.5	No melting	
				1370	.5	Partially melted (?)	
				1100 ^d	60	No melting	1:1 + "bronze" ^e
				1294	65	do	"6:11" + 9:8
				1405	0.5	Completely melted	Q - liq + "bronze"
36.36	63.64	1300	2	1353	1	No melting	"6:11" + 9:8 (7)
				1371	1	do	"6:11" + 9:8 (7)
				1388	1	do	"6:11" + 9:8 (7)
				1396	0.833	Completely melted	"6:11" + 9:8 (7)
				1368	.5	Partially melted	"6:11" + 9:8 (7)
				1332	17	do	
				1354	64	do	
				1361	0.5	do	
				1368	.5	Partially melted	

TABLE 19. *Experimental data for compositions in the binary system Nb₂O₅-WO₃. Part I—Continued.*

Composition		Heat treatments				Results					
Nb ₂ O ₅	WO ₃	Initial*		Final*		Physical observation	X-ray diffraction analyses†				
		Temp.	Time	Temp.	Time						
Mole %	Mole %	°C	Hour	°C	Hour						
35.20 (6:11)	64.71	1298	2	1372	.5	do.	"6:11" + 9:8 (?)				
				1360	.5	do.	"6:11" + 9:8 (?)				
				1100 ^d	60	No melting.	"bronze" + 1:1.				
				1100	64	do.	"bronze" + 1:1.				
				1203	19	do.	4:9 + 8:5 tr.				
				1246	165	do.	"6:11".				
				1305	48	do.	"6:11".				
				1246	165	No melting.					
				1358	1	do.	"6:11".				
				1369	93	do.	"6:11".				
33.33	66.67	1296	2	1371	1	do.	"6:11".				
				1360	1	Completely melted.	"bronze" + Q - liq.				
				1405	0.5	do.	"bronze" + Q - liq.				
				1100 ^d	60	No melting.	"bronze" + 1:1.				
				1339	166	do.	"6:11" + 4:9 (?)				
				1362	1	do.	"6:11" + 4:9 (?)				
				1371	1.5	Partially melted.	"6:11" + Q - liq.				
				1381	1	Completely melted.	"bronze" + Q - liq (tr).				
				32	68	1300	2	1522	17	do.	4:9 + "6:11" (?)
								1579	166	do.	4:9 + "6:11" (?)
1354	64	do.									
1361	0.5	do.									
1368	.5	Partially melted.									
1375	.5	do.									
1100 ^d	60	No melting.	"bronze" + 1:1 (tr).								
1100	64	do.	"bronze" + 1:1 (tr).								
1150	160	do.	Poorly crystalline 4:9.								
1203	19	do.	4:9 + "bronze" + 8:5 (?) †.								
30.77 (4:9)	69.23	1299	2	1300 ^d	48	do.	4:9.				
				1305	48	do.	4:9.				
				1150	160	do.	4:9.				
				1368	1	do.					
				1379	0.75	Partially melted.	4:9.				
				1405	1	Completely melted.	"bronze" + Q - liq (tr).				
				1100 ^d	60	No melting.	"bronze" + 1:1 (tr).				
				28.57 (2:5)	71.43	1296	2	1361	1	Partially melted.	
								1371	1	do.	4:9 + Q - liq.
								1382	1	Completely melted.	"bronze" + Q - liq.
1100 ^d	60	No melting.	"bronze" + 1:1 (tr).								
1203	19	do.	"bronze" + 4:9 (?)								
1300 ^d	48	do.	4:9 + 2:7 (?)								
1305	48	do.	4:9 + 2:7 (?)								
1249	70	do.									
1265	1	do.	4:9 + 2:7 (?)								
1330	0.5	do.	"bronze" + 4:9 (?)								
28	72	1296	2	do.		do.	4:9 + 2:7 (?)				
				1329	16.5	do.	4:9 + 2:7 (?)				
				1341	1	Partially melted.	4:9 + Q - liq.				
				1353	1	do.	4:9 + Q - liq.				
				1364	1	do.					
				1372	1.5	do.					
				1382	1	Completely melted.	"bronze" + Q - liq.				
				1100 ^d	60	No melting.	"bronze" + 1:1 (tr) + WO ₃ (tr) †.				
				1200	70	do.					
				1300	2	do.					
27.27 (5:8)	72.73	1296	2	1340	0.5	Partially melted.	4:9 + Q - liq.				
				1361	.5	do.					
				1100 ^d	60	No melting.	"bronze" + WO ₃ + 1:1 (tr) †.				
				1200	70	do.	"bronze".				
				1347	64	Partially melted.	4:9 + "bronze".				
				1335	67	do.	4:9 + 2:7.				
				1200	2	No melting.					
				1250	65	do.					
				1200	2	No melting.					
				1304	16	do.					
26.42 (14:39)	73.58	1296	2	1369	0.5	Partially melted.	"bronze" + WO ₃ + 1:1 (tr) †.				
				1100 ^d	60	No melting.	"bronze".				
				1200	70	do.					
				1200	2	do.					
				1250	65	do.					
				1335	67	do.	4:9 + 2:7.				
				1347	64	Partially melted.					
				1361	0.5	do.					
				1200	2	No melting.					
				1304	16	do.					
25 (1:3)	75	1296	2	1370	0.5	Partially melted.					
				1100 ^d	60	No melting.	"bronze" + WO ₃ + 1:1 (tr) †.				
				1250	2	do.	"bronze + L.MonWO ₃ + "1:11" + WO ₃ ".				
				1250	260	Partially melted.					

TABLE 19. Experimental data for compositions in the binary system Nb₂O₅-WO₃. Part I—Continued.

Composition		Heat treatments				Results	
Nb ₂ O ₅	WO ₃	Initial ^a		Final ^b		Physical observation	X-ray diffraction analyzer ^c
		Temp.	Time	Temp.	Time		
Mole %	Mole %	°C	Hour	°C	Hour		
22.22 (2:7)	77.78	1200	2	1255	65	No melting	"bronzes" + 2:1
				1305	16	do	4:9 + 2:7 (?)
				1343	0.75	Partially melted	4:9 + Q - liq.
				1353	.5	do	4:9 + Q - liq.
				1360	.5	do	
				1379	.5	Completely melted	
		1200	60	1225	97	No melting	"bronzes" + L-MonWO ₃ (tr)
				1100 ^d	60	No melting	4:9 + 2:7 (?)
				1160	64	do	"bronzes" + WO ₃ + 1:1 (tr) ^e
				1202	20	do	"bronzes" + WO ₃
				1250	260	Partially melted ^f	"bronzes" + L-MonWO ₃ + WO ₃ (?)
				1265	1	do	"bronzes" + L-MonWO ₃ ^g
		1200	2	1300 ^h	40	do	
				1343	0.5	do	
				1360	1	Completely melted	
				1229	67	No melting	"bronzes" + Tet WO ₃
				1256	66	do	2:7
				1305	16	do	"bronzes" + "1:11" ⁱ
		1200	2	1342	0.5	Partially melted ^j	
				1361	.583	Completely melted ^k	
1229	67			No melting	2:7		
1331	64			do	2:7		
1344	0.75			do	2:7		
1350	.5			do	2:7		
20	80	1200	2	1357	.5	Partially melted	
				1100 ^l	60	No melting	"bronzes" + WO ₃ + 1:1 (tr) ^e
				1200	2	do	
				1343	0.5	Partially melted	
				1255	65	No melting	"bronzes" + 2:7 + L-MonWO ₃ ^m
				1305	16	do	"bronzes" + 2:7 + "1:11" + "1:15" (?) ⁿ
18	82	1200	2	1342	0.5	Partially melted	
				1352	.5	Completely melted (?)	
				1365	.5	Completely melted	
				1200	2	No melting	2:7 + "1:11"
				1300	2	do	2:7 + "1:11"
				1337	0.5	Partially melted	2:7 + Q - liq.
15	85	1200	2	1352	.5	Completely melted	
				1100 ^o	60	No melting	WO ₃ + "bronzes"
				1232	17	do	"bronzes" + L-MonWO ₃ + WO ₃ (tr)
				1258	16	do	"bronzes" + L-MonWO ₃ + 2:7 (?) ^p
				1300	16	do	2:7 + "1:11"
				1343	0.5	Partially melted	
10	90	1200	2	1352	.5	Completely melted	
				1359	.5	do	
				1100 ^q	60	No melting	WO ₃ + "bronzes"
				1100	20	do	WO ₃ + "bronzes"
				1200	90	do	L-MonWO ₃ + "bronzes" + WO ₃ (?)
				1250	17	Partially melted ^r	L-MonWO ₃ + "bronzes"
9	91	1200	2	1275	144	No melting	
				1258	16	do	L-MonWO ₃ + "bronzes" + 2:7 (?) ^s
				1300	16	do	"1:11" + "1:15" + 2:7 ^t
				1301	69	do	"1:11" + 2:7
				1328	0.5	do	
				1342	.5	Partially melted (?)	"1:11" + Q - liq.
8.33 (1:11)	91.67	1200	2	1343	17	Partially melted	"1:11" + Q - liq.
				1361	0.67	do	
				1378	.5	Completely melted	
				1295	67	No melting	"1:11" + 2:7 (tr)
				1301	69	do	"1:11" + 2:7 (tr)
				1379	0.083	No melting	
1200	2	1279	85	Partially melted			
		1311	70	do	"1:11" + 2:7 + "1:15" (?) ^u		
		1350	17	do	"1:11" + 2:7 (tr) ^v		
		1350	17	do	"1:11"		
		1354	0.33	No melting	"1:11"		
		1355	.5	do	"1:11"		
1200	2	1357	.5	do	"1:11"		
		1359	.5	Partially melted	"1:15" + L-MonWO ₃ + Q - liq.		
		1359	.5	do	L-MonWO ₃ + Q - liq.		
		1369	.5	do	L-MonWO ₃ + Q - liq.		
		1379	.083	do			
		1392	.083	Completely melted	Q - liq.		

TABLE 19. Experimental data for compositions in the binary system Nb₂O₅-WO₃. Part I—Continued.

Composition		Heat treatments				Remarks					
		Initial ^a		Final ^b		Physical observation	X-ray diffraction analyses ^c				
Nb ₂ O ₅	WO ₃	Temp.	Time	Temp.	Time						
Mole %	Mole %	°C	Hour	°C	Hour						
8	92	1200	2	1301	69	No melting.					
7.69	92.31	1200	2	1311	70	No melting.	"1:11" + "1:15" (?).				
7.14	92.86	1200	2	1311	70	No melting.					
-7	93	1200	2	1299	67	No melting.					
		1300	2			do	"1:15" + "1:11".				
6.66	93.33	1200	2	1391	0.063	No melting.					
		1300	2	1402	0.063	Partially melted.					
6.25	92.75	1200	16	1312	68	Completely melted.					
				1279	65	No melting.	"1:15" + "1:11".				
6	94	1200	2	1312	68	do	Tet WO ₃ + "bronze" + 17 WO ₃ .				
				1312	68	do	"1:15" + L-MonWO ₃ + "1:11" + 2:7 (tr).				
				1350	17	do	"1:15" + L-MonWO ₃ (?).				
				1200	2	do	"1:15" + L-MonWO ₃ (?).				
				1300	2	do	"1:15" + L-MonWO ₃ (?).				
				1355	0.6	do	"1:15" + L-MonWO ₃ (?).				
				1369	5	do	"1:15" + L-MonWO ₃ (?).				
				1380	0.063	do	"1:15" + L-MonWO ₃ (?).				
				1391	0.063	do	L-MonWO ₃ + Q - 14.				
				1398	0.063	do					
				1200	2	do	No melting.				
				1300	2	do	No melting.				
5	95	1200	2	1347	4.5	do	"1:15" + L-MonWO ₃ (tr).				
				1357	0.5	do	"1:15".				
				1359	5	do	L-MonWO ₃ + Q - liq.				
				1364	5	do	L-MonWO ₃ + Q - 14.				
				1299	67	do	No melting.				
				1108 ^a	60	do	"1:15" + L-MonWO ₃ .				
				1300	30	do	WO ₃ + "bronze".				
				1260	30	do	WO ₃ + "bronze".				
				1202	30	do	L-MonWO ₃ + "bronze" + WO ₃ (tr).				
				1250	30	do	L-MonWO ₃ + "bronze" + WO ₃ (tr).				
				1250	16	do	L-MonWO ₃ + Q - 14.				
				1275	144	do	L-MonWO ₃ + "1:15" + WO ₃ (tr).				
4	96	1200	2	1200	20	Partially melted (?).	L-MonWO ₃ + WO ₃ + "bronze".				
				1250	16	do	L-MonWO ₃ + WO ₃ + "bronze".				
				1299	67	do	L-MonWO ₃ + WO ₃ + "bronze".				
				1380	16	do	L-MonWO ₃ + "bronze".				
				1343	0.5	do	"1:15" + L-MonWO ₃ .				
				1343	17	do	L-MonWO ₃ + "1:15" + WO ₃ + "1:11" (?).				
				1362	0.75	do	"1:15" + L-MonWO ₃ + "1:11" (?).				
				1379	0.75	do	L-MonWO ₃ .				
				1400	5	do	L-MonWO ₃ + Q - 14 (?).				
				1411	0.063	do	Completely melted.				
				1420	25	do	do				
				3	97	1200	2	1254	66	No melting.	L-MonWO ₃ (?).
1355	19	do	L-MonWO ₃ + "1:15".								
1410	0.063	do	Partially melted.								
1200	2	do	No melting.								
2	98	1200	2	1301	68	do	L-MonWO ₃ + "1:15".				
				900 ^b	0.25	do	Tet WO ₃ + "1:15".				
				1254	66	do	No melting.				
				1355	19	do	L-MonWO ₃ + 2:7.				
				1379	0.063	do	L-MonWO ₃ + Orth WO ₃ (?).				
				1409	0.063	do	do				
				1422	0.063	do	Completely melted.				
				1300	2	do	No melting.				
				1301	68	do	No melting.				
				1	99	1200	2	900 ^b	0.25	do	L-MonWO ₃ + "1:15".
								1200	58	do	Tet WO ₃ + "1:15".
								1232	17	do	L-MonWO ₃ + WO ₃ + "bronze".
1275	19	do	L-MonWO ₃ .								
1355	19	do	L-MonWO ₃ + WO ₃ + "bronze".								
1379	0.063	do	do								
1391	0.063	do	do								
1405	0.063	do	Partially melted.								
1414	0.063	do	do								
1422	0.063	do	do								
1430	0.063	do	Completely melted.								
1435	0.063	do	do								
1200	2	do	No melting.								
1248	67	do	No melting.								
1	100	1200	2	900 ^b	0.25	do	L-MonWO ₃ .				
				900	25	do	Orth WO ₃ + Tet WO ₃ (tr).				
				900	25	do	Orth WO ₃ .				
1	100	1200	2	900 ^b	0.25	do	No melting.				
1	100	1301	68	900	25	do	No melting.				

TABLE 19. Experimental data for compositions in the binary system Nb₂O₅-WO₃. Part I—Continued.

Composition		Heat treatments				Results			
Nb ₂ O ₅	WO ₃	Initial ^a		Final ^b		Physical observation	X-ray diffraction analyses ^c		
		Temp. °C	Time Hour	Temp. °C	Time Hour				
1	99	1200	2	1200	90	No melting	WO ₃ + L-MonWO _{3m}		
				1400	0.083	do	WO ₃ + L-MonWO _{3m}		
				1411	0.083	Partially melted			
				1422	0.083	do			
				1430	0.083	do			
				1435	0.083	Completely melted			
				1200	2				
				1248	67			No melting	WO ₃ + L-MonWO _{3m}
						900 ^b		do	WO ₃
						1000	10	No melting	WO ₃
0	100			1329	2	do	WO ₃ + TrWO ₃		
				1351	48	do	WO ₃ + TrWO ₃		
				1420	0.083	do			
				1424	0.083	do			
				1430	0.083	do			
				1433	0.083	do			
				1435	05	Completely melted			
				1435	0.083	do			
				1436	0.166	do			
				1437	0.083	do			
				1439	0.083	do			
				1453	0.083	do			
				1462	0.083	do			
				1470	0.083	do			
				1485	0.083	do	WO ₃ + TrWO ₃		

Experimental data for compositions in the binary system Nb₂O₅-WO₃. Part II (less pure end members)

Composition		Heat treatments				Results	
		Initial ^a		Final		Physical observation	X-ray diffraction analyses ^c
		Temp. °C	Time Hour	Temp. °C	Time Hour		
93.75 (15:1)	6.25	1354	48	1053	336	No melting	6:1 + Nb ₂ O ₅
				1053	336	do	"20:1" + 6:1 (?)
				1354	40	do	"20:1" + 6:1 (?)
92.86 (13:1)	7.14					No melting	"20:1" + 6:1
85.71 (6:1)	14.29			1100	66	No melting	6:1
				1248	64	do	6:1
				1352	66	do	6:1
77.78 (7:2)	22.22			1104	60	No melting	6:1 + 7:3 + 13:4
				1203	89	do	6:1 + 13:4 + 7:3 (?)
				1352	21	do	6:1 + 7:3 + 13:4 (tr)
76.19 (16:5)	23.81			1044	336	No melting	13:4 + 6:1 (?) + 7:3 (?)
				1109	65	do	13:4 + 6:1 (?) + 7:3 (?)
				1200	130	do	13:4 + 6:1 + 7:3
72.73 (8:2)	27.27			1203	71	do	6:1 + 7:3 + 13:4 (tr)
				1201	89	No melting	13:4 + 7:3
70 (7:3)	30			1097	94	No melting	7:3 + 13:4 (tr)
				1346	23.5	do	7:3 + 6:1 (tr)
				1097	94	No melting	7:3 + 1:1
61.54 (8:5)	38.46			1201	113	do	8:5 + 7:3 (tr)
				1202	42	do	8:5 + 7:3 + "brassite"
				1255	65	do	8:5 + 7:3 (tr)
				1255	92	do	8:5 + 7:3 (tr)
				1303	16	do	8:5 + 7:3 (tr)
				1350	19	do	8:5 + 7:3 (tr)
				1354	16	No melting	8:5 + 7:3 (tr)
						do	8:5 + 7:3 + "brassite" + 1:1
						do	8:5 + "brassite" (tr)
60 (3:2)	40			1048	336	No melting	7:3 + 1:1 + "brassite" (tr)
				1203	89	do	8:5 + "brassite"
				1097	70	do	8:5 + "brassite" + 9:8 (?)
				1261	113	do	9:8 + 8:5 (tr)
				1255	65	do	9:8 + 8:5 (tr)
				1303	16	do	9:8 + 8:5 (tr)
				1310	67	do	9:8 + 8:5 (tr)
				1321	19	do	9:8 + 8:5 (tr)
				1348	20	do	9:8 + 8:5 (tr)
				1350	19	do	9:8 + 8:5 (tr)
52.94 (9:8)	47.06			1352	21	do	9:8 + 8:5 (tr)
				1352	68	do	9:8 + 8:5 (tr)
				1354	16	do	9:8 + 8:5 (tr)
						do	1:1 + 7:3 + "brassite" + 9:8 (tr) + 8:5 (tr)
						do	1:1 + "brassite" + 7:3
						do	1:1 + "brassite" + 7:3
50	50			1048	336	No melting	
				1090	336	do	
				1101	66	do	

Experimental data for compositions in the binary system $Nb_2O_5-WO_3$. Part II (less pure end members) - Continued.

Composition		Heat treatments				Results	
		Initial ¹		Final		Physical observation	X-ray diffraction analyses ²
		1368	20	1044	336		
42.31 (11:15)	57.69			1044	336	do	9:8 + "6:11"
35.14 (13:26)	64.86			1157	89	No melting	8:5 + "6:11" + 1:1
30	70			1305	22	do	"bronze" + 1:1 + 8:5
25	75			1305	67	No melting	"6:11" + 9:8
23.17 (19:63)	76.83			1305	22	do	"6:11"
11.11 (1:8)	88.89			1106	67	No melting	4:9 + "bronze" (?)
				1501	70	do	4:9 + "bronze" (?)
				1697	70	No melting	"bronze" + WO_3 (tr)
				1300	44	Partially melted (?)	"bronze"
				1500	70	do	"bronze"
				1302	24	No melting	"bronze" + WO_3 (tr)
				1279	46	Partially melted (?)	"bronze" + WO_3 (tr)
				1202	24	No melting	"bronze" + L-Mon WO_3 + WO_3
				1279	46	Partially melted (?)	"1:1" + "bronze"
				1300	44	do	"1:1" + 2:7

¹ All specimens were initially calcined at 700 °C for 19 hr with heating and cooling rates of approximately 4 °C/min. Small portions of this calcine were then reheated for the designated time at one or more designated higher temperatures and generally quenched in sealed Pt tubes.

² After the initial heat treatment(s) all specimens were reheated at the indicated temperature and quenched in sealed Pt tubes, unless otherwise specified.

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

(?) = interpretation is not certain.
 sr = solid solution.

(tr) = trace just barely discernible in x-ray pattern.

"30:1" = a phase of unknown structure occurring at a mole ratio of approximately 30:1 $Nb_2O_5:WO_3$.

(unknown) = extra peaks in the x-ray pattern which cannot be assigned to any recognized phase.

6:1 = $6Nb_2O_5 \cdot WO_3$ ($WNb_{12}O_{34}$).

low Nb_2O_5 = nonequilibrium phase (similar to the low temperature form of Nb_2O_5), when quenched from above the liquidus and examined at room temperature.

7:3 = $7Nb_2O_5 \cdot 3WO_3$ ($W_3Nb_{14}O_{41}$).

13:4 = $13Nb_2O_5 \cdot 4WO_3$ ($W_4Nb_{26}O_{77}$).

"bronze" = a metastable disordered phase having a variable composition and an undistorted tetragonal-tungsten-bronze type x-ray pattern (designated as "3:8" on the phase diagram).

8:5 = $8Nb_2O_5 \cdot 5WO_3$ ($W_5Nb_{16}O_{49}$).

9:8 = $9Nb_2O_5 \cdot 8WO_3$ ($W_8Nb_{18}O_{59}$).

1:1 = $Nb_2O_5 \cdot WO_3$ (WNb_2O_6).

"6:11" = a tungsten bronze type phase of orthorhombic symmetry and with superstructure lines indicative of the unit cell being tripled in the *a* and *b* directions.

Q-liq = metastable phase or phases which form only from the quenched liquid and give very poorly defined x-ray powder patterns, having the general appearance of the ReO_3 subcell characteristic of most of the compounds in this system.

4:9 = $4Nb_2O_5 \cdot 9WO_3$ ($Nb_8W_9O_{37}$).

L-Mon WO_3 = the monoclinic polymorph of WO_3 reported [30] to occur below -40 °C "stabilized" by a small amount of solid solution of Nb_2O_5 in WO_3 .

2:7 = $2Nb_2O_5 \cdot 7WO_3$ ($W_7Nb_4O_{29}$).

"1:11" = Magneli "shear" phase having the approximate composition $1Nb_2O_5 \cdot 11WO_3$.

"1:15" = Magneli "shear" phase having the approximate composition $1Nb_2O_5 \cdot 15WO_3$.

Tot WO_3 = the high temperature polymorph of WO_3 stabilized by solid solution of Nb_2O_5 in WO_3 .

Orth WO_3 = the orthorhombic polymorph of WO_3 ordinarily stable between about 320 °C and ~745 °C, "stabilized" by addition of Nb_2O_5 in solid solution.

Tr WO_3 = the polymorph of WO_3 which is stable just below room temperature and occurs at room temperature when pure WO_3 is ground in a mortar and pestle for a short time.

³ Specimen heated and cooled at about 4 °C/min rather than quenched.

⁴ Nonequilibrium, probably due to incomplete reaction.

⁵ Nonbinary equilibrium, postulated as being due to reduction.

⁶ Metastable melting.

⁷ Pt tube not sealed.

⁸ Did not receive preliminary 700° heat treatment.

5. Reduction

As has been previously mentioned there is some tendency for WO_3 to reduce on heating to high temperatures, even in the sealed Pt tubes. This tendency is greatly enhanced by subjecting the specimens to high temperatures for extended periods of time. The problem of reduction of WO_3 is apparently carried into the $Nb_2O_5-WO_3$ system to such an extent that the system may not be completely binary at high temperatures.

The experimental data indicate that the system probably remains essentially binary as long as no melting occurs. However, above the solidus, reduction very likely takes place and the measured liquidus values may not represent binary equilibrium. The melting relations around the ordered bronze-type phases, from about 60 to 80 mole percent WO_3 , were particularly difficult to obtain experimentally. The three compounds at "6:11", 4:9 and 2:7 all appeared to melt congruently with binary solidus relations between them. However, the liquidus values appear to form smooth curves between these congruent melting points without any indication of a change in slope toward a eutectic composition. The phenomena is characteristic of a nonbinary system. For the sake of simplicity, the phase diagram has been drawn to indicate the most likely binary relationships and liquidus curves have been dashed to illustrate the uncertainty in values.

Another indication of reduction occurs in the melting relations of the $B_{n+1}O_{2nm-(n+w)+4}$ homologous series compounds. The phases $W_2Nb_{14}O_{44}$, $W_5Nb_{16}O_{88}$, and $W_8Nb_{18}O_{99}$ were all observed to melt incongruently. Experimentally these compound compositions were observed to begin to form a second phase at about 1440°, 1385°, and 1375°, respectively. However, compositions in two-phase regions between these compounds did not begin to form the new phase until considerably higher temperatures, even though the specimen may have shown an appearance of partial melting. This apparent increase in the stability temperature of the phases is dependent on the amount of time the specimen is held at temperature, and binary melting relationships can be approximated by only holding the experimental specimens for relatively short periods of time. It may therefore be concluded that reduction takes place in these specimens, when held above the solidus temperatures. However, it has previously been concluded [1-4] that the crystal structure of these compounds precludes any appreciable nonstoichiometry. Therefore, it appears likely that, as the W^{+6} ion is reduced, the Nb/W ratio of these phases shifts towards higher tungsten content, maintaining an essentially constant cation: anion ratio.

In addition to the relatively large amount of reduction which apparently takes place above melting, there is possibly some tendency towards reduction at lower temperatures. The amount of reduction which can take place experimentally below melting is probably several orders of magnitude less than that which occurs above melting because it seems in no way to affect

the equilibrium relationships. The only indication of this minor reduction is in a tendency for discoloration of the specimens. Almost any composition in the system, held for long periods of time in a sealed Pt tube will show a much darker green or blue color than does the same composition either held for shorter periods or not sealed. This darker color can always be lightened by annealing the specimen in an open Pt tube, at about 1000°C.

It is intriguing to postulate that the $Nb_2O_5-WO_3$ system might more closely approach binary conditions if the experiments were performed under conditions of high oxygen pressure, such as that utilized by Van Hook [39]. Such studies will probably become more common in the near future.

6. Metastable Phases and Nonequilibrium

6.1. Metastable Melting

It can be seen from table 19, that all compositions containing more than about 72 mole percent WO_3 were calcined for about 2 hr at 1200°C prior to attempting to obtain equilibrium data. This was because the few preliminary heat treatments, performed at temperatures above 1250°C, without the 1200°C calcine, exhibited some indication of melting, whereas those with a prior 1200°C calcine did not show melting until much higher temperatures.

The metastable melting apparently is due to a eutectic type reaction between the metastable bronze-type solid solution (labeled "3:8" on fig. 1) and pure WO_3 . It must be emphasized that in order to obtain equilibrium products in the region 72 to 100 mole percent WO_3 , this preliminary calcine at 1200°C is imperative. Without such preliminary heat treatment, neither the 2:7 compound nor the "Magneli-shear" structures can be prepared as single phases.

6.2. Metastable Phases

The bronze-type solid solution previously mentioned ("3:8" of fig. 1) has been interpreted as a metastable phase for several reasons. It appears to be responsible for very low temperature metastable melting which takes place between about 72 and 100 mole percent WO_3 . The x-ray powder pattern of the bronze-type solid solution shows no superstructure and the phase must contain disordered pentagonal bipyramid polyhedra. It must be concluded that the disordered phase is metastable because it dissociates on heating to two ordered phases. All compositions between about the 6:11 and 2:7 ratios, when quenched from above the liquidus exhibit the tetragonal bronze-type "disordered" phase with no indication of any superstructure peaks in the x-ray diffraction powder pattern. In addition, many specimens also show a small amount of a quenched liquid phase which exhibits a poorly defined x-ray pattern characteristic of a disordered ReO_3 structure, the basic building unit of all the compounds in the system.

From about 5 mole percent WO_3 to at least 15 mole percent WO_3 another metastable phase is formed from the quenched liquid. This phase has an x-ray pattern, characteristic of the orthorhombic low-temperature form of Nb_2O_5 . The diffraction pattern shows diffuse peaks and little or no superstructure lines. This phase never appears below the melting point and only forms as a single phase if the quenching temperature is considerably above the liquidus. The ability of a specimen to be quenched as a phase with a powder pattern similar to the low-temperature form of Nb_2O_5 seems to be dependent on cooling rate and composition. This phenomena has been previously reported for other niobate systems [23, 40].

As shown in table 19, another unknown phase has been found occasionally in nonequilibrium mixtures between about 7 and 25 mole percent Nb_2O_5 . All efforts to establish an equilibrium region for this phase have proved negative. It is possible that the unknown compound may represent a complex structure involving a mixture of 3×3 blocks along with the 3×4 and/or 4×4 blocks normally found in other compounds occurring in this composition range. Such a structure would contain a considerable number of Nb^{+5} ions in tetrahedral coordination and would not be expected to form a stable assemblage.

It has been previously mentioned [1] that $\text{WNb}_{12}\text{O}_{33}$ appears to have a small region of homogeneity, in that the 8:1 composition showed only a single phase. However, no change in lattice parameters could be detected. This apparent region of solid solution is most likely due to the very similar powder patterns of the 6:1 compound and the "30:1" phase. A solid solution structure containing either a considerable number of oxygen vacancies or cation interstitials (in either the tetrahedral or octahedral position) would be expected to cause some measurable change in the lattice parameters. One other possible mechanism which has been suggested for solid solution [41] is again raised by the appearance of the "ordered intergrowth" structure of the 13:4 compound, and the suggested possible structure of a 25:1 or 20:1 compound. It is possible that a "disordered intergrowth" of double 3×4 blocks might occur in association with the isolated 3×4 blocks of the $\text{WNb}_{12}\text{O}_{33}$ compound [24]. Such a disordered structure or "compositional stacking fault" would cause only a slight broadening of some of the lines of an x-ray diffraction powder pattern, as would a mixture of a small amount of the ordered second phase. It is problematical as to whether such a picture of solid solution would represent a stable or only a metastable condition. Nevertheless, proof of such a postulated disorder would necessarily depend upon other data, outside the scope

of this project. A careful electron diffraction and/or electron microscope study might prove of value in deciphering the nature of any possible disorder of this type.

7. References

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