High-Temperature Ternary Oxide Phases in Tantalum/Niobium–Alumina Composite Materials


Coarse-grained composites of refractory ceramics and refractory metals are a novel approach for materials at application temperatures up to 1500 °C. Al2O3 and the refractory metals Nb and Ta are suitable candidates for enhanced thermal shock capability, as they show similar thermal expansion. During fabrication, a key aspect to consider is the possible formation of additional phases upon interaction of the constituent phases as well as through reaction with the environment. X-Ray diffraction (XRD) and investigations of the microstructure with scanning electron microscopy methods unveil Al2O3–Nb composite to form NbO, whereas for Al2O3–Ta the ternary compound aluminum tantalate (AlTaO4) is found. Thermodynamic calculations show that the changing oxygen solubility in Nb accounts for the formation of NbO, and explain the absence of a corresponding niobate (AlNbO4) phase. AlTaO4 is identified as the disordered tetragonal high-temperature modification.

1. Introduction

Refractory metal (molybdenum, niobium, and tantalum) matrix composite materials in combination with refractory ceramics (alumina, zirconia, and mullite) are of particular interest for high-temperature applications, e.g., in refractory linings in the casting industry. However, besides the high-temperature applicability, other aspects need consideration as well for the choice of material combination, such as: 1) the reaction between the metal and ceramic particles, 2) the chemical interaction with the environment, and 3) the thermal mismatch between the ceramic and the metallic phases. The majority of the past publications dealt with so-called fine-grained refractory composites. However, their disadvantages are high shrinkage on sintering, thus resulting in limited thermal shock ability. The refractory metals Nb and Ta are promising candidates for application with Al2O3 due to their similar thermal expansion behavior. Wang et al. and White et al. showed for temperatures between 1000 and 2000 K a similarly linear increase of the linear thermal expansion coefficient for the three elements, with 8.8 to 11.2 × 10⁻⁶ K⁻¹ for α-Al2O3, 8.3 to 10.4 × 10⁻⁶ K⁻¹ for Nb, and 7.1 to 8.4 × 10⁻⁶ K⁻¹ for Ta. Recently, Zienert et al. demonstrated the fabrication of coarse-grained refractory composites based on alumina with niobium or tantalum, while Weidner et al. reported first mechanical properties at high temperatures (1300–1500 °C) under compressive load. In these coarse-grained refractory composites, alumina with different particle classifications from 0–20 μm up to 2–5 mm (with different volume fractions...
fractions) was combined with fine metal powders of Ta or Nb (d ≤ 75 μm). Moreover, Zienert et al.\cite{12} successfully fabricated niobium-alumina aggregate fractions with particle sizes up to 3 mm by crushing pre-synthesized fine-grained composites. A shrinkage of 1.6% was determined for such coarse-grained composites after sintering.

Weidner et al.\cite{11} showed that the coarse-grained Ta–Al2O3 and Nb–Al2O3 exhibited good mechanical properties depending on the volume fraction of refractory metals and temperature. Thus, compressive strengths of about 70 and 53 MPa were achieved at 1300 °C for 11 vol% Ta and Nb, respectively. 21 vol% metal yielded compressive strength values of 27 and 32 MPa, respectively. None of the tested specimens failed in a brittle manner, in all cases, certain plasticity was observed. However, one or two of the above-addressed issues seem to appear for this class of refractory metal (Nb, Ta) composites: 1) the reaction between the metal and ceramic particles and/or the chemical interaction with the environment, resulting in the apparent formation of new phases (see Figure 3 and 4 for Ta-composites and Nb-composites, respectively, in Ref. \cite{11}).

Earlier literature described the possible formation of such phases in the systems Ta–Al2O3\cite{13} and Nb–Al2O3.\cite{14} The consolidation of our samples takes place under a protective Ar atmosphere with residual oxygen (in the range of a few ppm) and, additionally, under the presence of different contaminants in the raw materials. One of the conclusions drawn in Ref. \cite{11} based on scanning electron microscope (SEM) investigations was the possible formation of tantalates and niobates during the sintering process in the fabrication route of these materials, but the exact identity of the phases remained unknown. At the time of writing, neither experimental evidence for the existence of these new phases nor thermodynamic calculations were available and will be presented in this article. In addition, it is of interest at which stage during preparation these phases form: During sintering or subsequent mechanical loading at high temperature?

The aims of the present study are thus twofold: 1) experimental analysis of the formed new phases in both systems Ta–Al2O3 and Nb–Al2O3 and 2) thermodynamic calculations of the formation of these new phases. We conducted microstructural investigations based on XRD and SEM using electron backscattered diffraction (EBSD) in combination with backscattered electron (BSE) contrast for the systems Nb–Al2O3 and Ta–Al2O3. Thermodynamic calculations of isothermal sections of the ternary phase diagrams Al–Nb–O and Al–Ta–O underpin our investigations.

2. Experimental Section

The refractory ceramic–refractory metal composites investigated in this work were prepared by Zienert et al.\cite{10} from coarse-grained (i.e., coarse particle size) alumina and fine-grained niobium or tantalum, respectively. They were already part of investigations by Weidner et al.\cite{11} The raw materials used are listed in Table 1.

Alumina particles from four different classifications were used: 0–0.5 mm (21 vol%), 0.5–1 mm (10 vol%), 1–3 mm (13 vol%), and 2–5 mm (24 vol%). The refractory metals were added as two fine powder fractions, 0–20 μm (10 vol%) and 0–45 μm (11 vol%), yielding a total metal content in the castable of 21 vol%. 8 vol% CL370 and 3 vol% alphabond were used as additives. The powders with a volume of ≈350 cm³ were mixed in a concrete mixer (ToniMAX, Toni Baustoffprüfsysteme GmbH, Berlin, Germany) using 5 wt% water and 1 wt% of dispersing alumina as additives. The resulting blends were cast into prismatic (150 × 25 × 25 mm³) sample molds and left to rest for 1–3 days. The samples were removed, dried in air at 120 °C for 24 h, and sintered pressurelessly under an argon atmosphere at 1600 °C for 4 h. The following investigations focus on composite materials containing 21 vol% of Nb or Ta. The investigated samples are hereinafter called 21F0C-Nb and 21F0C-Ta, according to the nomenclature given in Refs. \cite{10,11}. For this purpose, the samples were cut parallel to the cylinder axis and embedded one half in EpoMet resin (Buehler, Esslingen am Neckar, Germany), ground to SiC grit P4000, and subsequently polished with 3 and 1 μm diamond suspension. The final polishing was performed with colloidal SiO2 suspension.

Crystal structure analyses were performed with XRD using a Bruker D2 phaser in Bragg–Brentano geometry with Cu Kα radiations at a step size of 0.01°. For electron microscopy imaging and EBSD the dual-beam scanning electron and focused ion beam microscope Zeiss Auriga 60 (Carl Zeiss AG, Oberkochen, Germany) was used equipped with a BSE imaging detector and an EDAX DigiView EBSD camera. The BSE images at three different contrast levels were used for phase segmentation based on gray-level binarization, employing a self-programmed MATLAB script, and compared to EBSD at 20 kV. Oxygen contents of the metals were determined with atom probe tomography (APT) measurements in a LEAP 4000X HR by Cameca.

Isothermal sections of the Al–Ta–O and Al–Nb–O ternary system at 1600 °C were obtained by CALPHAD modeling as described in Ref. \cite{15}.

3. Results

3.1. XRD

XRD phase analysis was performed for both, the 21F0C-Nb and 21F0C-Ta samples.

Figure 1 shows the XRD line profile of 21F0C-Nb and confirms the presence of α-Al2O3 (blue diamond symbols, corundum prototype, and space group 167), Nb (red squares,
W prototype, and space group 229), and NbO (yellow circles, NbO prototype, and space group 221), with the characteristic peaks of the latter at $2\theta = 30.0^\circ$, $48.3^\circ$, and $62.4^\circ$. In addition, the crystallographic data matches different modifications of Nb$_2$C (violet triangles) of the monoclinic (space group 6) and the orthorhombic (space group 33) variants, as well as of the hexagonal superstructure originally reported by Terao[16] (ε-Fe$_2$N prototype, space group 149). No unequivocal assignment to any of these prototypes is possible based on the assigned peaks at $2\theta = 33.5^\circ$, $36.8^\circ$, and $50.1^\circ$. While the peaks labeled by green squares result from K$_\beta$ diffraction, a few unidentified peaks remain, which may result from contaminations in the castable additives, notably the alumina binder, which contains traces of Na$_2$O, SiO$_2$, and CaO. No evidence of niobates (e.g., AlNbO$_4$) was found from XRD measurements.

For the 21F0C-Ta composite, XRD (Figure 2) confirms the presence of α-Al$_2$O$_3$, Ta, and, as Weidner et al.[10] suggested, the ternary oxide AlTaO$_4$ (aluminum tantalate, TiO$_2$ prototype, space group 136) with characteristic peaks at $2\theta = 27.4^\circ$, $35.8^\circ$, and $39.1^\circ$. K$_\beta$ signals are again marked in green, while a few peaks remain unassigned. A possible candidate for the unassigned peaks at $2\theta = 23.0^\circ$ and $32.6^\circ$ is sodium tantalate (NaTaO$_3$), with castable additives and Na contamination in the alumina as a sodium source.[17]

### 3.2. Microstructural Investigations

To study the presence of additional phases such as NbO or niobates and tantalates, as suggested by Weidner et al.,[11] SEM investigations were performed using BSE contrast and EBSD measurements.

Figure 3 depicts a phase segmentation micrograph of 21F0C-Nb. The three BSE images used can be found in the Figure S1, Supporting Information. A mixture of coarse-grained alumina (blue) particles and Nb (red) particles with metal grain sizes ranging from a few microns to up to 75 µm is visible. In addition, fine-grained alumina particles are present in between Nb particles. In yellow, an intermediate BSE contrast is assigned to an additional phase, which was suggested in Ref. [11] to be either NbO or Al niobate (AlNbO$_4$) based on SEM-EDS measurements. This will be further discussed in the context of Figure 4. Black areas are related to porosity remaining in the material after the sintering process. The lack of different contrast levels suggests that no additional phase is present within the resolution of the technique. Phase segmentations performed in these regions yield an average Nb volume fraction of about (27 ± 1) vol.-% and an alumina fraction of (44 ± 4) vol%. Porosity was determined to be (27 ± 4) vol%. The suspected NbO makes up roughly (0.2 ± 0.05) vol%, with a NbO/Nb ratio of 0.007 ± 0.001, and is located within or at Nb particles. Areal phase fractions are regarded to be equivalent to volume fractions due to the isotropy of the microstructure. However, the local heterogeneity of the material has to be taken into account, and, therefore, a single image resolving the finely distributed tertiary phase cannot be representative. Weidner et al.[11] mentioned an average porosity of about 17 vol% measured with the Archimedes principle for the complete sample, which gives a more representative value for the typical material than the 27 vol% obtained from local 2D SEM investigation but considers only open porosity. Due to higher shrinkage, porosity is more significant in regions with large amounts of fine-grained material. When only considering
coarse-grained alumina regions, the porosity measured with phase segmentation from SEM is 15 vol% and below.

Additional EBSD measurements confirmed the presence of the suspected NbO or AlNbO4 phases. The EBSD phase map and the electron backscatter diffraction patterns (EBSP) shown in Figure 4 confirm that the different phases indicated in Figure 3 correspond to α-Al2O3, Nb, and NbO. Thus, no evidence of AlNbO4 was found from EBSD measurements. When compared to Figure 3, some falsely assigned pixels in the EBSD map occur. This is due to unfavorable grain orientations and insufficient surface finish in certain regions, which stems from the largely varying strength and hardness of the phases involved.

Figure 5 depicts a representative region of the microstructure of 21F0C-Ta with Ta particles (red) mixed with a smaller fraction of alumina and neighbored by two major alumina particles (blue). The BSE images used for phase segmentation can be found in Figure S2, Supporting Information. Regions of a third, intermediate contrast in the vicinity of Al2O3 and Ta are colored yellow and are assigned to AlTaO4. As an average of three-phase segmentation, we find a Ta volume fraction of about (17 ± 2)% and an alumina fraction of (56 ± 5) vol%. The porosity amounts to (22 ± 3) vol%, while the AlTaO4 phase fraction is (0.2 ± 0.06) vol%. The ratio of ternary oxide to the refractory metal Ta amounts to 0.012 ± 0.002, which is considerably more than the NbO/Nb ratio in 21F0C-Nb. Similar to the Nb-containing composite, the heterogeneity of the material again complicates the determination of phase fractions. While Ta roughly corresponds to the initial amount of 21 vol%, the alumina fraction is lower than the amount in the castable. In coarse-grained alumina regions, the Al2O3 content exceeds values of 75 vol%. The observed porosity is roughly in the range of 17 vol% as reported by Weidner et al.[11]

The region depicted in Figure 5 was examined with EBSD to confirm the phase assignments. Figure 6 features a phase map of the region with EBSPs of all the phases present, i.e., Al2O3 (blue), Ta (red), and AlTaO4 (yellow), which confirms the findings from X-Ray diffraction. Predominantly within the oxides, some Al2O3 regions were falsely assigned to AlTaO4 due to poor Kikuchi pattern quality. These also do not appear as respective contrast for the phase segmentation map in Figure 5.

3. Thermodynamic Considerations

A discussion of the phases present in the microstructure of the refractory composites requires the phase diagrams of the systems Al–Nb–O and Al–Ta–O at the sintering temperature (1600 °C). These diagrams are not available in the literature and must be calculated using appropriate thermodynamic databases. Using the well-known CALPHAD method,[15] Gebauer et al.[18] assess thermodynamic datasets for both ternary systems. The databases consist of Gibbs energy functions that are temperature, pressure, and composition dependent. Thermodynamic equilibria are calculated by evaluating these functions for specified condition sets. Isothermal sections are then obtained by evaluating these equilibria at a given temperature for two independent composition variables. This dataset considers phase descriptions for intermetallic phases, binary and ternary oxides, and an emphasis on the quasi-binary section Al2O3–Nb2O5.

Figure 7 shows the calculated isothermal section at 1600 °C and standard pressure for the Al–Nb–O system. At this
temperature, the maximum oxygen solubility in Nb, indicated by a red arrow, is about 6 at% but will decrease with decreasing temperature.\cite{19} APT measurements (see Figure S3, Supporting Information) in the Nb particles indeed yield a lower value of \((3.0 \pm 0.2)\) at% which corresponds to the solubility at about 1200 °C.\cite{19}

For the Al–Ta–O ternary system, the calculated isothermal section at 1600 °C and 10^5 Pa. L denotes liquid phase regions, whereas \(\varepsilon, \delta,\) and \(\sigma\) are Al–Nb intermetallic phases. The three-phase region Nb–Al2O3–NbO is marked yellow.

4. Discussion

The above-presented results of XRD and SEM have shown that reactions between the refractory metal and ceramic particles and/or possibly the chemical interaction with the environment lead to the formation of AlTaO4 in the system Ta–Al2O3 and NbO in the system Nb–Al2O3. Figure 1 and 2, moreover, indicate that these newly formed phases are not localized but are found throughout the material. Comparison of Figure 3 and 5 shows that NbO is present within or at Nb particles. In contrast, the phase identified as AlTaO4 is adjacent to both Al2O3 and Ta, suggesting a reaction involving alumina. Further phases, such as Nb2C, found in small amounts with XRD, could not be recognized with SEM methods.
The findings that these additional phases form during sintering are supported by thermodynamical considerations using the CALPHAD method. Whereas some NbO was already found in the original Nb powder,[12] its presence in the composites can be attributed to additional oxygen uptake of the samples during the sintering process, as is shown in Figure 7. The oxygen concentration of (3.0 ± 0.2) at% found in Nb with APT is considerably lower than the solubility limit of 6 at% at 1600 °C and corresponds to the limit at about 1200 °C. Thus, due to this decrease during cooling in the sintering furnace, an initially two-phase Nb–Al2O3 composite with 6 at% O in the metallic particles will form NbO, as the phase region boundary Nb–Al2O3 moves to the left upon cooling and the three-phase region marked in yellow extends to lower oxygen fractions. Several possible sources, like the sintering atmosphere, but also castable additives, e.g., water, may account for the increased oxygen level. Figure 7, moreover, shows why no ternary Al–Nb–O compound (aluminum niobate, AlNbO4, space group 12, indicated in blue) is formed, as it is not stable in the presence of metallic Nb. According to Figure 7, AlNbO4 is obtained through a reaction of Al2O3 and Nb2O5. Thus, the absence of the niobate precludes the occurrence of the latter during sintering. The experimentally determined occurrence of NbO, thus, matches the calculated three-phase equilibrium Nb–Al2O3–NbO from the database presented by Gebauer et al.[18].

Likewise, the experimental findings match calculations of the ternary system Al–Ta–O. Thus, as for Al–Nb–O, during cooling, the decrease of the oxygen solubility in the metallic regions, i.e., an extension of the yellow three-phase region to smaller oxygen fractions, will induce the formation of an additional oxide phase. However, unlike for Al–Nb–O, where NbO coexists with Al2O3 and Nb, here, it is the mixed oxide AlTaO4 that shares a three-phase field with Al2O3 and Ta. Furthermore, this compound can be obtained through a reaction of Al2O3 with Ta2O5.[13] as shown in Figure 8. This oxide may have formed through partial oxidation of the metal powder during the heat treatment but was not found in our investigations.

Notably, for the presented specimen, XRD and EBSD confirm that the structure of AlTaO4 is tetragonal. Ternary oxometallates of the formula AlMO4 (with M = Nb, Ta, among others) crystallize in a monoclinic crystal structure of the AlNbO4 prototype (space group 12).[21–23] However, Jasper-Tönnies and Müller-Buschbaum,[13] as well as Sarazin in an earlier publication,[24] reported the formation of an additional metastable tetragonal rutile-type AlTaO4 from Al2O3 (corundum) and Ta2O5 in the temperature range of 1300–1400 °C, and at 1800 °C. Roth et al.[25] and Wichmann et al.[26] correlated the preferred formation of this modification as grains at the alumina–metal interface. As expected from the calculations, a corresponding aluminum niobate (AlNbO4) was not found experimentally. This compound forms through a reaction of Al2O3 and Nb2O5, thus, its absence rules out the occurrence of the pentoxide.

Supporting Information
Supporting Information is available from the Wiley Online Library or from the author.

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Conflict of Interest
The authors declare no conflict of interest.

Data Availability Statement
The data that support the findings of this study are available from the corresponding author upon reasonable request.

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