Silicide characterization at alumina–niobium interfaces

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Abstract Alumina–niobium interfaces formed by liquid-film-assisted joining with copper/niobium/copper interlayers exhibited microstructures that depend on the nature of the alumina components. Characterization of these interfaces in the transmission electron microscope provided insight on the relationship between interfacial microstructure and fracture performance. Interfaces between sapphire and niobium and those between high-purity (99.9%) polycrystalline alumina and niobium were free of secondary phases. However, niobium silicides were found at interfaces between lower-purity (99.5%) alumina and niobium, identified by electron diffraction analysis as the body-centered tetragonal \( \alpha \)-Nb\(_5\)Si\(_3\) phase. Spatially resolved compositional analysis was conducted on silicide particles at and away from the interface.

Introduction

The formation of secondary phases plays an important role in both the processing and properties of fabricated ceramic–metal interfaces. Alumina (\( \text{Al}_2\text{O}_3 \))–niobium (\( \text{Nb} \)) interfaces represent a model ceramic–metal system, where the effects of interface chemistry have been studied [1–7] and interfacial Nb silicides have been previously observed [1, 6, 7]. The combination of polycrystalline \( \text{Al}_2\text{O}_3 \) and polycrystalline Nb, and their contact along an interface, results in a wide range of potential nucleation sites for silicide formation. These include \( \text{Nb}–\text{Nb}, \text{Al}_2\text{O}_3–\text{Al}_2\text{O}_3 \), and \( \text{Al}_2\text{O}_3–\text{Nb} \) grain boundaries, triple junctions involving either two \( \text{Al}_2\text{O}_3 \) or two Nb grains, and four-grain corners involving \( \text{Al}_2\text{O}_3 \) triple lines or Nb triple lines. Each type of site has a distinct spatial density and a distinct nucleation barrier that will combine to define a spectrum of nucleation frequencies. The site providing the optimum combination of energetics and spatial density will have the highest overall nucleation rate [8]. Growth rates will be influenced by the site-specific diffusion paths and Si availability, and in conjunction with nucleation rates, will determine the nature of the site or sites that dominates the overall distribution of reaction products. The properties and microstructure of such reaction products, Nb silicides in the case of interest here, can, in turn, influence the fracture path and fracture energy.

Silicide formation

Studies of Nb silicide nucleation generally involve planar “bulk” reaction couples or planar “thin-film” couples produced by depositing thin Nb films onto single-crystal Si substrates, which are subsequently annealed at various temperatures. Theoretical consideration [9–11] based on
the equilibrium phase diagram and the interfacial energies that appear in the nucleation barrier to silicide formation predict that the first crystalline silicide phase to form in the Nb–Si binary system will be NbSi₂ (Fig. 1 gives the Nb–Si binary equilibrium phase diagram [12]). An effective heat of formation concept [13–16] making direct use of thermodynamic data has been successful in predicting silicide phase-formation sequences in multiple metal–Si systems, as well as explaining how the presence of impurities can alter this sequence. Experimentally, the nature of the phase(s) formed depends upon the nature of the experiment performed, e.g., a thin-film versus bulk reaction couple, the specific anneal time and temperature. In thin-film studies [17–21], various stable and metastable silicide phases and phase-formation sequences have been found, depending on deposition conditions and subsequent annealing conditions. The silicides found include NbSi₂ [17, 18], Nb₅Si₃ [17–19], Nb₃Si [17–20], amorphous Nb silicides of varying composition [17, 18], and a body-centered orthorhombic Nb₁₁Si₄ phase [21]. In thin-film (and other periodic) systems, interpretation can be difficult due to limited amounts of one or more components, phases forming sequentially rather than simultaneously, and the effects of interfaces on growth and diffusion [22]. To avoid these complications, bulk diffusion couples of Nb–Si and Nb–NbSi₂ were used to study diffusion-controlled reactive growth of NbSi₂ and Nb₅Si₃ [22]. In Nb–Si couples, NbSi₂ nucleated first and grew much faster than the Nb₅Si₃ layer that developed (the NbSi₂ layer was ≈1 order of magnitude thicker than the Nb₅Si₃ layer). In both diffusion couples, an Nb₅Si₃ layer developed between the Nb and NbSi₂. Preliminary experiments showed that the growth rate of Nb₅Si₃ was very low at temperatures below 1200 °C and obtaining a layer a few microns in thickness required very long anneal times.

It is to be expected that as systems under consideration become more chemically complex, further complexities can arise in the phase-formation preferences and sequences. In the interlayer used for joining, Cu is a minor constituent concentrated near the interface between the Nb-rich interlayer and a potentially silicate-containing Al₂O₃. This renders the Nb–Si–Cu system of interest. The ternary Nb–Si–Cu system has been investigated in both thin-film reaction couples [23] and “bulk” phase-equilibria studies [24, 25]. In the thin-film study, the phases formed and their formation sequence again depended on the experimental conditions. The observed phases were processing-dependent subsets of ternary Nb₅Cu₄Si₄ and binary Cu₃Si, NbSi₂, Nb₅Si₃, and Nb₃Si phases. The presence of Cu at the Nb–Si interface was found to affect the onset temperature for NbSi₂ and Nb₅Si₃ formation. Isothermal sections of the ternary Nb–Si–Cu phase diagram are available at 1500, 875, and 800 °C. Figure 2 shows the 1500 °C isothermal section. It should be noted that, according to the Nb–Cu binary phase diagram [12], there is very low solubility of Cu in Nb and Nb in Cu. The Nb₅Si₃ and Nb₃Si phases dissolve ≈4 at.% Cu in the temperature range 800–875 °C. The ternary tetragonal Nb₅Cu₄Si₄ phase was found to form only upon rapid cooling from the melt or by solid-state diffusion (as occurred in the thin-film diffusion couples) [24]. This phase had previously been observed at 800 °C [26] and

Fig. 1 Nb–Si equilibrium binary phase diagram. Reproduced after Ref. [12]

![Phase Diagram Image](image-url)

Weight Percent Silicon

| Temperature (°C) | Composition |
|-----------------|-------------|
| 1200            | NbSi₂       |
| 1900            | Nb₅Si₃      |
| 2500            | Liquid      |

Notice that the phase diagram includes the following key points:
- NbSi₂:
- Nb₅Si₃:
- Liquid:

These points are crucial for understanding the phase behavior in the Nb–Si system.
this observation was confirmed in a later study [25]. A tetragonal Nb₄CuSi phase has been observed at 1500°C, in equilibrium with pure Nb and Nb₅Si₃ [25].

Intergranular glassy phase in alumina

The presence of a Si-containing intergranular glassy phase in Al₂O₃ has been shown to cause the formation of Nb silicide at Nb–Al₂O₃ interfaces [1, 6, 7]. Intergranular phases and films have been the subject of numerous investigations (as reviewed in Ref. [27]). Many have focused on Al₂O₃-based materials (e.g., [28–34]). Efforts have been made by Dillon et al. [32–34] to relate and categorize the effects of grain boundary “phases” on grain growth and transport kinetics along grain boundaries. In Al₂O₃ six distinct types of kinetic behavior were identified, each associated with a particular grain boundary “phase” that produced a characteristic grain boundary transport rate. These characteristic rates would, in turn, be expected to affect the mobility of impurities such as Nb along grain boundaries.

The wetting behavior, structure, and composition of a grain boundary in a system containing a glassy phase are generally determined by the boundary misorientation of adjacent grains [29, 30]. The majority of grain boundaries are wetted by a silicate glass phase, while low-angle and special (twin-related) boundaries are not wet. A continuous intergranular glassy phase is present at higher-energy boundaries. Isolated pockets of glassy phase and continuous films containing pockets at growth ledge steps and faceted grain boundaries have been observed [30, 31]. These could potentially serve as preferred nucleation sites for secondary phases.

It was proposed [35] that migration of an intergranular glass phase to a free surface (or interface) is driven by differences in wetting behavior of the surface (or interface) and the boundary, and this was observed for specific misorientations of sapphire bicrystals [29]. Intergranular films at metal–ceramic interfaces [31] that form as a result of an intergranular glass phase in alumina were found to have the same characteristic features as intergranular films at ceramic grain boundaries. The metal played an important role in determining the film chemistry and thickness, as well as the composition of glass pockets at triple junctions. Intergranular phases may affect the preferred nucleation sites for second phases and their growth kinetics.

Mechanical properties

In idealized systems in which two phases are in contact and form bimaterial interfaces, thermal-expansion mismatches can establish residual stresses that alone or in combination with applied stresses cause failure at or near the interface (see, e.g., [36]). Even when the coefficients of thermal expansion (CTEs) are closely matched, as is the case for Al₂O₃ and Nb, differences in other physical properties will affect the fracture path and resulting fracture energy. Fracture at a bimaterial interface is generally described by the complex stress-intensity factor [37–39], \( K \), in which the elastic mismatch across the interface is expressed using the nondimensional Dundurs parameters [40], \( \alpha \) and \( \beta \). It was noted by Kruzic et al. [6] that in sandwich-type specimens, for which complex stress-intensity solutions exist [41], analysis is complicated by deviations of cracks from the interface [42] and the validity of these solutions breaks down at distances ahead of the crack tip [43]. Using the strain energy release rate, \( G \), in mode I loading to evaluate crack driving forces [6] avoids these complications. In addition, \( G \) is easy to compute, invariant with respect to crack position, and unaffected by the presence of the interlayer in sandwich specimens when the interlayer thickness is small compared with other relevant specimen dimensions [41].

Cannon et al. [44, 45] determined that while linear-elastic calculations predict crack propagation along a path with maximum \( G \) and to be dependent on the Dundurs parameters (modulus mismatch), this analysis has limitations when applied to bimaterial systems and microstructural effects are important. Engineering interfaces with enhanced toughness involves generating plasticity in the metal interlayer, the competition between microstructurally weak crack paths and paths of maximum driving force, and encouraging crack jumping across the metal interlayer and
crack meandering. At Al₂O₃–Nb interfaces, it is expected that cracks that initiate close to the interface will be drawn to the Nb interlayer. The detailed crack path and fracture energy will depend on interlayer thickness and yield strength, as well as microstructural and chemical characteristics of the interface.

In the samples employed in the current study, Nb silicides along the ceramic-interlayer interface have the potential to provide microstructural weak links and can generate localized residual stresses that interact with an advancing crack. Nb silicides are brittle at room temperature. Vickers hardness values of 5.6–9.5 GPa [46] and KIC values of 2.3–5.5 MPa√/m [46, 47] were measured for pure Nb₅Si₃ samples. If one assumes an elastic modulus of 188 GPa for Nb₅Si₃ [48], the KIC values would imply mode I fracture energies, GIC, of 28–161 J/m². Investigations of the fracture energy of heterophase interfaces involving Nb silicide [49] found the interfacial fracture energy of Al₂O₃ with Nb₅Si₃ to be ≈16 J/m², while the interfacial fracture energy between Nb and Nb₅Si₃ was found to exceed 34 J/m².

Effects of silicides at processed interfaces

Even in a “simple” system such as Nb–Si, a range of phases and phase distributions develop in multilayer systems. The associated microstructural differences have the potential to induce differences in the response to loading, specifically, the fracture path and fracture energy.

In the Al₂O₃–Nb system, the potential Si source is an intergranular glassy phase in the Al₂O₃. In fracture studies of Al₂O₃–Nb interfaces in which the Al₂O₃ was heavily contaminated with silicate (97% pure Al₂O₃) [1], Nb embrittlement due to oxygen uptake was augmented relative to assemblies using higher-purity Al₂O₃ by flow of Si/SiO₂ to the Al₂O₃–Nb interface. Patches of Nb silicide (Nb₅Si₃ or Nb₅Si₄) were identified on fracture surfaces of specimens that failed within the Nb. These patches extended into the Nb in the vicinity of the Al₂O₃–Nb interface.

Collective observations from studies of Al₂O₃–Nb interfaces fabricated with Cu/Nb/Cu interlayers [6, 50–53] indicate a substantial sensitivity of the fracture behavior to the processing temperature, bonding atmosphere, interlayer design, post-bonding heat treatments, and testing method, even in situations where, from a macroscopic perspective, the system is approximated as an Al₂O₃–Nb bond (a bimaterial interface) and the same Al₂O₃ is used throughout. This suggests that the interfacial microstructure is sensitive to the details of the processing, and that changes in the processing parameters, broadly defined, influence the interplay between fracture paths of maximum driving force and those of microstructural weakness. Changes in the nature of the Al₂O₃ add further complexity. This work was undertaken in part to more fully characterize the nature of the ceramic–metal interfaces generated, particularly the interface between an Al₂O₃ containing an intergranular glassy phase and a Nb-based interlayer, and relate the observed microstructures to the fracture characteristics of the interfaces.

Experimental procedures

The materials and experimental procedures used to produce samples for the characterization studies described herein are identical to those used in prior studies to produce joined samples with Cu/Nb/Cu interlayers. Three types of Al₂O₃ were used: an undoped, high-purity, single-crystal, (0001)-oriented sapphire (≈99.996% pure, Insaco Inc., Quakertown, PA, USA); a high-purity, high-strength, fine-grain-size polycrystalline Al₂O₃ (≈99.9% pure, SSA-999W, Nikkato Corp., Osaka, Japan); and a lower-purity, lower-strength, coarse-grain-size polycrystalline Al₂O₃ (≈99.5% pure, AD995, Coors Technical Ceramic Co., Oak Ridge, TN, USA). The 99.9% pure material had a uniform microstructure with an average (mean linear intercept) grain size of ≈1 μm; the 99.5% pure Al₂O₃ exhibited a bimodal grain-size distribution, with the larger grains roughly 25 μm in size and the finer-grained regions typically consisting of grains ≈5 μm or less in size.

The as-received sapphire was polished to an optical finish. Surface roughness measurements using contact profilometry yielded average Ra values of 0.0036, 0.0037, and 0.0108 μm for line scans of 50, 500, and 5000 μm, respectively. The bonding surfaces of polycrystalline Al₂O₃ blocks were first ground flat with a 400 grit (37 μm) diamond wheel, then polished with successively finer-grit diamond (9, 6, 3, and 1 μm) paste and finished with 0.02 μm colloidal silica suspension. For the polycrystalline samples, average Ra values for line scans of 50, 500, and 5000 μm were 0.0089, 0.0105, and 0.0784 μm, respectively. The polished surfaces of the Al₂O₃ substrates were coated with ≈3.0-μm-thick Cu films. The coated Al₂O₃ substrates were joined using a 125-μm-thick Nb foil (≈99.99% pure, Goodfellow Corp., Malvern, PA, USA) in a graphite-element hot press. A bonding pressure of ≈2 MPa was maintained for 6 h at 1400 °C under a vacuum of ≈10⁻² Pa (10⁻⁷ atm). In-depth processing details are available in previous publications [50–55].

Cross-sectional TEM specimens of the interfaces were prepared by conventional grinding, polishing, and dimpling techniques. The thin foils were perforated using a low-energy Ar⁺ ion mill. Conventional TEM (imaging and diffraction analysis) was performed on a JEOL 200CX and a JEOL 2011 operating at 200 keV. Elemental analysis using energy-dispersive spectrometry (EDS) was performed on a Philips CM200 FEG (S)TEM operating at 200 keV with a probe size of 2 nm.
Results and discussion

The solubility–diffusivity product for Cu in Nb is sufficiently low at the bonding temperature (1400 °C) that, for a bonding time of 6 h, discrete particles of residual Cu decorate the sapphire–Nb and Al₂O₃–Nb interfaces. The morphology and projected (plan view) area fraction of these ductile interfacial Cu particles influence the extent of toughening obtained through metal plasticity and particle tearing, and thereby affect the fracture strength of the joined assemblies [6, 7, 49, 52, 56]. The ultimate spatial distribution of Cu along the ceramic-interlayer interface will depend upon the bonding temperature, pressure, and time, the bonding-surface roughness, and the microstructure and chemistry of the bulk ceramic [7, 51–53, 55, 57]. Well-polished samples with Rₐ much less than the Cu film thickness, fabricated using the same bonding conditions, were selected to isolate the effects of Al₂O₃ microstructure and purity.

Prior work has shown that in all three types of samples, the area fraction of residual Cu particles at the ceramic-interlayer interface is typically ≤5% for the processing conditions described. As a result, a plan view of the ceramic-interlayer interface is well suited to detecting and characterizing the Cu particle size and distribution. Comparisons of sapphire-interlayer interfaces and polycrystalline Al₂O₃-interlayer interfaces reveal that the sizes and distributions of particles are affected by the ceramic microstructures. Specifically, the average size of Cu particles, and to a lesser degree the projected area fraction, decreases with the grain size. This, coupled with observations of the fracture-surface topography, suggests that grain-boundary grooving in both the metal foil [7, 55] and on the ceramic surface plays a role in defining the average size and spacing of the Cu particles. Decreasing the average Cu particle size will in turn increase the amount of Cu-interlayer contact area per unit area of interface, which may increase the total flux of Cu into the Nb core layer and contribute to decreasing area fractions of Cu with decreasing Al₂O₃ grain size.

As a result of the low area fraction of Cu particles, assumed to be ≤0.05, the average nearest-neighbor particle separation distance is estimated to be of the order of 3–5× the average particle diameter (≈10 μm in samples using sapphire substrates to ≤1 μm in samples using high-purity Al₂O₃ substrates). A random cut perpendicular to the plane of the interface, designed to allow a cross-sectional view, will generally intersect relatively few and widely separated particles. In an SEM, it is possible to view in cross-section an interface that is multiple millimeters in length, and those segments in which particles are intersected can be examined. In the TEM work described here, a typical length of ceramic-interlayer interface is in the range of 2–20 μm. It follows that the likelihood of a particle intersection is very low. This is consistent with our observations (or lack thereof). Moreover, due to the relatively higher milling rate of Cu, even when Cu particles are intersected, there is a high likelihood that preferential etching of a through-thickness particle will generate a particle-shaped perforation along the ceramic-interlayer interface. Isolated observations of such features have been made.

In view of the understandable absence of Cu particles, the primary differences in the interfacial microstructures and chemistries among the three types of samples fabricated reflect the effects of the Al₂O₃ microstructure on the interface “roughness” and of the Al₂O₃ chemistry on Al₂O₃-interlayer compatibility and the potential for reaction-product formation along the interface. Figure 3 presents bright-field TEM images of the three types of fabricated interfaces for comparison of the microstructures.

Fig. 3 Bright-field TEM images of a a (0001) sapphire–Nb interface, b a high-purity Al₂O₃–Nb interface (note the preferential thinning at the interface), and c a lower-purity Al₂O₃–Nb interface revealing the presence of Nb silicides.
Figure 3a shows a (0001) sapphire–Nb interface. The interface appears distinct and smooth with no reaction phases present [3, 55, 58, 59], as sapphire eliminates the possibility of a glassy phase flowing to and filling interfacial voids [55]. EDS confirmed that silicides and other secondary phases were not present at the sapphire–Nb interface.

Figure 3b shows a bright-field TEM image of a high-purity Al$_2$O$_3$–Nb interface. Again, there was no evidence of silicide or other secondary-phase formation at the interface, as expected for a high-purity Al$_2$O$_3$. EDS confirmed this. The interface appears “rough” relative to the sapphire–Nb interface due to grooving of the fine-grained Al$_2$O$_3$ during the bonding cycle, with the Nb conforming to the Al$_2$O$_3$ grain-boundary grooves. This roughness was even more evident in fracture surfaces when joined assemblies prepared from this Al$_2$O$_3$ failed along the ceramic-interlayer interface [55]. As observed in previous studies [60–62], preferential thinning of the interface occurs during TEM specimen preparation.

Figure 3c reveals the microstructure along a lower-purity Al$_2$O$_3$–Nb interface. Again, the Nb has conformed to the Al$_2$O$_3$ grain-boundary grooves, and the interface appears “rough” relative to the single-crystal sapphire–Nb interface. Nb silicides are evident at and slightly away from the Al$_2$O$_3$–Nb interface, consistent with previous studies [6, 7] in which Nb-silicide formation was observed in small-grained regions of the Al$_2$O$_3$. This higher grain-boundary density in these regions leads to a higher flux and local concentration of Si per unit area of Al$_2$O$_3$ at the interface as the glassy silicate phase present at Al$_2$O$_3$ grain boundaries flows to the interface. Chemical analysis using EDS [6, 7] indicated a Nb:Si ratio of ~4.5/3 (~1.5) and the silicide particles were determined to be the Nb$_5$Si$_3$ phase (see Fig. 1). At these interfaces, nucleation likely occurred where Nb metal was in contact with Al$_2$O$_3$ grain boundaries, and thus silicate that flowed to the interface. Formation of pockets at triple lines involving two Nb–Al$_2$O$_3$ and one Al$_2$O$_3$–Al$_2$O$_3$ grain boundary and four-grain junctions where three Al$_2$O$_3$–Al$_2$O$_3$ and one Nb–Al$_2$O$_3$ interface come together will produce larger localized reservoirs of Si (silicate). The proximity of these pockets to the Nb source suggests that nucleation should be preferred at the interface rather than at subsurface triple lines or four-grain junctions.

Silicide formation requires a sufficiently low O$_2$ activity; the equilibrium O$_2$ partial pressure to form Nb$_5$Si$_3$ from a reaction between pure SiO$_2$ and Nb is of the order of $10^{-14}$ atm (10$^{-9}$ Pa) at 1400 °C. The O$_2$ partial pressure for Nb–NbO equilibrium at 1400 °C is $\approx 6 \times 10^{-18}$ atm (6 $\times$ 10$^{-13}$ Pa) at 1400 °C [63]. The Nb foil is thus expected to establish a low O$_2$ activity along the Al$_2$O$_3$–Nb interface, which in turn promotes the reaction of an intergranular silicate with Nb to form a silicide in the near-interfacial region. Moreover, oxygen readily dissolves in and is a fast diffuser in Nb [64, 65]. The dissolution makes the reaction more favorable, and the rapid diffusion prevents a build-up of dissolved oxygen at the interface. The graphite-element environment coupled with a 10$^{-7}$ atm (10$^{-2}$ Pa) vacuum pressure during bonding is expected to establish a sufficiently low ambient O$_2$ pressure to allow oxygen removal from the foil edges. Under these conditions, the observed formation of interfacial silicides is expected.

The local microstructure seen in Fig. 3c suggests that following nucleation the particles grew into the Al$_2$O$_3$ along grain boundaries, where the Si concentration will be higher as the reaction progresses. The silicide particles also grew laterally along the Al$_2$O$_3$–Nb interface and slightly into the Nb. Lateral expansion of the silicide along the Al$_2$O$_3$–Nb interface is likely due to enhanced diffusion of Si along the Al$_2$O$_3$–silicide boundary or flanking Al$_2$O$_3$–Nb boundary, as the rate of diffusion along this boundary can be expected to be much greater than diffusion of Si through the silicide lattice. Faceting at the interface between Nb and the Nb silicide was observed (see, for example, Fig. 7a), which may limit the growth rate of the silicide into the Nb foil. Grain boundaries in the silicide particles suggest multiple nucleation sites along Al$_2$O$_3$ grain boundaries.

Chemical analysis was performed on silicide particles at and away from the Al$_2$O$_3$–Nb interface using EDS in the TEM. Figure 4a shows EDS spectra obtained from multiple regions of a silicide particle at the interface. The spectra show the presence of Nb, Si, and Cu, and no dissolved Al in the particle. When the spectra are normalized by their maximum Nb$_L$ counts to reduce specimen-thickness effects, the normalized spectra overlap, indicating constant Nb:Si and Nb:Cu ratios in the silicide particle. Figure 4b presents an EDS line scan across the silicide particle. The dashed line in the figure indicates the position of the interface between the Nb metal and silicide particle. The interface does not appear distinct due to the combined effects of specimen thickness and tilting of the interface (the interface is not edge-on) relative to the incident electron beam. Minimizing these thickness effects was again achieved by normalizing the Cu and Si counts by the Nb counts at each position in the scan. The Nb:Si and Nb:Cu ratios appear constant across the silicide particle. The nominal composition of this particle was 54 (±2) at.% Nb, 40 (±2) at.% Si, and 6 (±5) at.% Cu, with a Nb:Si ratio of 1.35. The presence of dissolved Cu is a result of the cladding layers of the interlayer used to join the Al$_2$O$_3$.

Small, isolated particles on Al$_2$O$_3$ grain boundaries away from the Al$_2$O$_3$–Nb interface, believed to have formed during cooling from the processing temperature [7].
were also analyzed. Figure 5 shows an EDS line scan across a particle that lies \( \approx 1 \mu m \) from the Al\(_2\)O\(_3\)-Nb interface, with the associated HAADF image. The EDS scan again reveals the presence of Nb, Si, and trace amounts of Cu. The dashed line in b indicates the interface between the Nb and the silicide particle.

In order to accurately determine the specific Nb-silicide phase, crystallographic analysis was conducted using electron diffraction. Based on the Nb:Si ratios, which will be further discussed subsequently, attention turned to the \( \gamma \)-Nb\(_5\)Si\(_3\) phase. There are two stable allotropes of Nb\(_5\)Si\(_3\), both with a body-centered tetragonal crystal structure (space group 14/mcm): \( \gamma \)-Nb\(_5\)Si\(_3\) is stable below 1940 °C with lattice parameters \( a = 6.571 \) Å and \( c = 11.889 \) Å while \( \beta \)-Nb\(_5\)Si\(_3\) is stable above 1650 °C with lattice parameters \( a = 10.018 \) Å and \( c = 5.072 \) Å \([12, 66, 67]\). Metastable \( \gamma \)-Nb\(_5\)Si\(_3\) is hexagonal (space group P6\(_3\)/mcm) with lattice parameters \( a = 7.536 \) Å, \( c = 5.249 \) Å, and \( \gamma = 120^\circ \) and has been found to be stabilized by interstitial impurities, particularly C \([66, 67]\). As a specific example of
stabilization by interstitial impurities, the addition of Hf was found to stabilize \( c_\text{Nb}_5\text{Si}_3 \) up to 1500 \( ^\circ \text{C} \) \[68\]. At the processing temperature employed in this study, \( \text{Nb}_2\text{Si}_2 \), with a hexagonal crystal structure (space group \( P6_22 \)) and lattice parameters \( a = 4.819 \text{\AA} \), \( c = 6.592 \text{\AA} \), and \( \gamma = 120^\circ \), is also stable \[66\]. In diffusion couples involving bulk \( \text{Nb}/\text{Si} \) and bulk \( \text{Nb}_2\text{Si}_2 \) phases in the temperature range from 1200–1350 \( ^\circ \text{C} \), only \( \text{Nb}_2\text{Si}_2 \) and \( \text{Nb}_5\text{Si}_3 \) formed \[22\]. Similarly, in samples prepared by field-activated pressure-assisted combustion synthesis of elemental \( \text{Nb} \) and \( \text{Si} \) powders at 1400 \( ^\circ \text{C} \), either \( \text{Nb}_2\text{Si}_3 \) or a combination of \( \text{Nb}_5\text{Si}_3 \) and \( \text{Nb}_2\text{Si}_2 \) were formed \[46\]. We note that reports of multiple other phases exist in the literature, but the phases appear in samples processed either at higher or lower temperatures. \( \text{Nb}_2\text{Si} \) appears in the binary phase diagram between 1770 and 1980 \( ^\circ \text{C} \), and both stable tetragonal primary \( \text{Nb}_3\text{Si} \) \[20, 69\] and finer metastable orthorhombic \( \text{Nb}_2\text{Si}_2 \) particles \[20\] have been observed in solidified materials, but these materials were processed at higher temperatures than the joints produced for this study. A body-centered orthorhombic \( \text{Nb}_{11}\text{Si}_4 \) \[21\] phase has recently been reported to form as a precipitate in \( \text{Nb}–\text{Si} \) solid solutions. The tetragonal \( \text{Nb}_3\text{Cu}_4\text{Si}_4 \) phase has been reported in both phase-equilibria studies \[25\] and thin-film reaction couples \[23\] processed at temperatures lower than the bonding temperature used in the present work. Additionally, tetragonal \( \text{Nb}_3\text{Cu}_4\text{Si}_4 \) \[25\] has been observed. Consequently, it was desirable to identify the crystallographic form(s) of the silicides in our material.

The silicide particle shown in Fig. 4, from which EDS data was obtained, was chosen for crystallographic analysis. The lateral expansion of the silicide particle at the interface decreases the likelihood of surrounding \( \text{Al}_2\text{O}_3 \) or \( \text{Nb} \) either above or below the silicide through the thickness of the TEM foil. The bright-field image in Fig. 7a shows the silicide particle, recorded in the [001] zone axis. There was no low-index plane-on-plane orientation relationship between this specific silicide and \( \text{Nb} \) pair, nor was any preferred orientation relationship evident for any silicide particles along the interface or alumina grain boundaries. This may have implications as to whether the silicide phase nucleated on one of the “substrate” phases (\( \text{Nb} \) or \( \text{Al}_2\text{O}_3 \)) or within the glassy phase. Examinations of \( \text{Nb}–\text{Al}_2\text{O}_3 \) fracture surfaces using Auger electron spectroscopy \[6\] suggest that the glassy phase does not wet the \( \text{Nb}–\text{Al}_2\text{O}_3 \) interface and is largely confined to pockets. Experiments of the type performed in Refs. \[29–31\] may be better suited for determining whether the glassy phase wets the silicide/contacting-phase interface. If the glassy phase wets this interface, nucleation may occur within the glass pocket. If the silicide nucleated within the glassy phase, no preferred orientation relationship would be expected.

Four experimental zone-axis diffraction patterns were obtained, presented in Fig. 7b; these zone axes are, as indicated in the figure, [001], [111], [211], and [201]. The diffraction patterns are superimposed on a quadrant of the (001) stereographic projection of a tetragonal crystal, which also illustrates the relation between the four orientations. The four respective simulated zone-axis diffraction patterns for the \( x\text{-}\text{Nb}_5\text{Si}_3 \) phase are shown in Fig. 7c, and they are consistent with the experimental diffraction data. The diffraction analysis revealed the silicide particle to be the low-temperature body-centered tetragonal \( x\text{-}\text{Nb}_5\text{Si}_3 \) phase, confirming prior EDS results \[6, 7\] that found a Nb:Si ratio consistent with \( \text{Nb}_5\text{Si}_3 \).

Although EDS analysis indicated differing Nb:Si ratios in the silicides at and slightly away from the ceramic-interlayer interface, no difference in their crystal structures was evident. The small size of intergranular silicides, combined with the lack of a preferred orientation relationship between the silicide and either of the adjoining, misoriented \( \text{Al}_2\text{O}_3 \) grains, introduced diffraction from the surrounding \( \text{Al}_2\text{O}_3 \) matrix that complicated diffraction analysis. Figure 8 shows bright-field and dark-field images of a small silicide particle \( \approx 500 \text{ nm} \) from the \( \text{Al}_2\text{O}_3\)-interlayer interface. Diffraction patterns revealed this particle to also be the \( x\text{-}\text{Nb}_5\text{Si}_3 \) phase. The images were
obtained by tilting into a (200) systematic row condition from the [001] zone axis, as shown in the diffraction patterns inset in the figures.

In view of the presence of Nb, Si, and, to a more limited degree, Cu in the silicide particles, the Nb:Si ratios and the overall compositions were assessed in the context of the available Nb–Si and Nb–Si–Cu phase diagrams. The measured Nb:Si ratios, 1.35 and 1.21, place the composition within the two-phase field between Nb5Si3 and NbSi2 on the binary equilibrium phase diagram. Two experimental studies [24, 25] of phase equilibria in the Nb–Si–Cu system are available. Experimental evidence indicates that Nb5Si3 can incorporate up to 12 at.% Cu at 1500 °C. At 875 °C, the compositions would lie in a three-phase stability field of Nb5Si3, NbSi2, and a ternary liquid; the phase diagram suggests a solubility limit of Cu in Nb5Si3 of \( \approx 2 \) at.% [24]. At 800 °C, the nominal compositions would lie in a three-phase stability field of Nb5Si3, NbSi2, and Cu3Si, with the Nb5Si3 phase incorporating up to \( \approx 2–3 \) at.% Cu [24].

The EDS results, when assessed in the context of available phase diagrams, predict a multiphase equilibrium microstructure. However, evidence of silicide phases other than the Nb5Si3 phase was not found using electron diffraction. Contrast in bright-field images also did not reveal a second silicide phase, either within the Nb5Si3 particles or at the interfaces between Nb5Si3 particles with the Al2O3 and Nb. While it is possible that a much more extensive examination of silicide particles (a dozen were examined using electron diffraction) may reveal other phases, a key
A calculation using the range of measured compositions (including error) gives values of 0.02–0.33 for the expected phase fraction of the NbSi2 phase. Examination of a dozen particles suggests that this phase did not form. Several other factors are likely to contribute to this apparent disparity between the observations and expectations based on equilibrium. The phase diagram assessments (1500, 875, and 800 °C sections) did not include compositions lying in the "Nb5Si3–NbSi2" two-phase region. Thus, there is uncertainty in the limits of the Nb5Si3 phase stability field in the ternary diagram. However, a more significant source of error will stem from difficulties in obtaining quantitative EDS data from small, dispersed silicide phases within a matrix phase (surrounding Nb and Al2O3), leading to large uncertainty in the measured compositions. This will cause shifts in the apparent compositions of the silicide particles. Previously [20, 69], compositions of binary silicides were determined by assuming an exact stoichiometry to calculate k-factors, thus providing a range of compositions that fit this stoichiometry. Furthermore, the analysis of a multiphase specimen in which the electron beam passes through one or more phases will lead to an incorrect measurement of composition for the phases. In prior studies [69] efforts were made to examine only silicides that projected out from the edge of the TEM foil, avoiding the complications of a multiphase analysis. In this study, it was not possible to obtain spectra from isolated silicide particles unless the silicides were at the ceramic–metal interface. The quantitative correction software makes the implicit assumption that the specimen has a composition given by the combination of all the elements present in the spectrum, which will lead to erroneous application of the ZAF corrections [70].

In parallel, one can also compare the findings to those of prior studies of niobium-silicide nucleation and growth involving Nb/Si planar reaction couples [11], bulk diffusion couples of Nb/Si and Nb/NbSi2 [22], or the deposition of thin Nb films onto single-crystal substrates [9], followed by annealing at various temperatures. As discussed by Milanese et al. [22], thin-film studies can provide limited amounts of one or more components, phases may form sequentially rather than simultaneously, and interfaces may affect growth and diffusion. The discussion in the literature of the considerations that enter into determining the first phase to nucleate in thin-film structures [9, 11, 17, 18] provides an example of such effects. Nucleation energetics may favor metastable phases [20] or extend the stability field. Moreover, in the present work the Si source is a silicate phase, the Nb is alloyed with Cu and O, and growth is along either Al2O3–Nb or Al2O3–Al2O3 interfaces. Thus, chemical differences and microstructural factors may also play a role.

As mentioned previously, changes in the bonding temperature (from 1150 to 1400 °C), changes in the atmosphere during bonding or post-bonding anneals (vacuum versus gettered Ar), increases in the Cu film thickness (1.5 to 5.5 μm), and changes in fracture test method (bend versus fatigue tests) all appear to influence the dominant path of crack propagation in these materials, with an overall increase in the average fracture strength and a transition from all interfacial to predominantly ceramic failure for samples using the optimum processing conditions (McKeown et al., unpublished). Depending upon the bonding and test conditions, significant tearing of the Nb foil, propagation of the crack along the ceramic-interlayer...
interface, or significant excursions into the Al₂O₃ was observed.

Several studies have revealed that the interfacial silicides tend to form most easily in the finer-grained regions of the Al₂O₃. In view of the bimodal character of the Al₂O₃ grain size, this suggests that a substantial fraction of the total interface could potentially include silicide particles. This is consistent with the relative ease of finding the particles in TEM cross-sections. A reduced milling rate relative to Cu is also expected, which facilitates observation. The microscopy indicates that the Nb₂Si₃ grows both laterally along the ceramic–metal interface and into the Al₂O₃ along grain boundaries. As shown in this study and prior work [6, 7, 52], the silicides do not provide a continuous, low-energy fracture path. The silicides are, however, effective in altering the crack path when fracture occurs at or near the ceramic-interlayer interface.

In cases where the fracture has involved excursions of the crack front into the metal foil, one can speculate that the failure proceeded through the patches of silicide along the ceramic–metal interface that extend into the metal. Nb₂Si₃ is known to be brittle, and the fracture toughness has been reported to be between 2.3–5.5 MPa/√m [46, 47]. Equilibration of the silicide with the adjoining niobium foil will increase the Si levels and locally harden the niobium. EDS results are unlikely to be sufficiently accurate to distinguish between a failure that propagates through the silicide, along the interface, or deviates to a limited extent into the Si-saturated niobium.

In assessing the fatigue behavior, Kruzic et al. [6] observed that cracks repeatedly extended into the Al₂O₃ but were drawn back to the interface. It was concluded that although the modulus mismatch between Al₂O₃ and Nb would draw the crack to the interface, the lower intrinsic fatigue-crack propagation resistance of some Al₂O₃ grain boundaries allowed the crack to deviate into the ceramic. Preexistent flaws were implicated as possibly expediting these excursions. It appears possible, based on observations in this study, that in addition to pores or near-interfacial pockets of glassy phase, silicide particles located along grain boundaries could also serve as “preexistent flaws.” In addition to having low fracture toughness, the coefficient of thermal expansion of Nb₂Si₃ is reported to be 6.2 x 10⁻⁶ K⁻¹ [47], and thus lower than that of Al₂O₃ (≈ 8.2 x 10⁻⁶ K⁻¹). Both the portions of the Nb₂Si₃ that extend parallel to the Al₂O₃-interlayer interface and the wedge-like particles that extend down the grain boundaries would generate tensile stresses in the Al₂O₃. Since these particles do not extend to any significant depth below the Al₂O₃-interlayer interface, this may be a factor in keeping the cracks in the Al₂O₃ to a substantial degree and preventing more substantial deviations away from the interlayer. The lower interfacial fracture energy of Nb₂Si₃ with Al₂O₃ (≈ 16 J/m²) relative to that of Nb₂Si₃ with Nb (≥ 34 J/m²) [49] may also help to account for crack deviations into the alumina.

A much more limited set of bend tests was conducted in which the finer-grain-size 99.9% pure Al₂O₃ was bonded with Cu/Nb/Cu interlayers for varying times (5 min, 30 min, 6 h) at 1400 °C. For the 6 h samples [55], a limited number of room-temperature tests resulted in exclusively ceramic failure and an average fracture strength of ≈ 360 MPa (versus 240 MPa for the 99.5% pure Al₂O₃). The transition to ceramic failure suggests a substantial strengthening of the ceramic-interlayer interface. Whether this is a consequence of fewer or less severe flaws along or near the interface as a result of grain-size refinement and altered Cu-particle characteristics or whether this may in part be due to the lack of periodic arrays of silicide particles is uncertain. Further studies of this behavior that would improve the statistics would be useful, as would more detailed fracture studies. In a system where fracture can involve a wide range of homophase and heterophase interfaces (Al₂O₃/Nb, Al₂O₃/Nb₂Si₃, Al₂O₃/Cu, Nb/Nb₅Si₃, Nb/Cu, and Nb₂Si₃/Cu), understanding microstructural effects and the character of phases that are present becomes a necessary precursor toward fabricating strong, reliable ceramic–metal interfaces.

Summary

Prior work, based largely on optical and scanning electron microscopy, suggested that Al₂O₃–Nb interfaces processed with single-crystal sapphire or high-purity, polycrystalline Al₂O₃ were free of secondary phases. In contrast, interfaces fabricated with a lower-purity, polycrystalline Al₂O₃ formed Nb silicide particles along the Al₂O₃–Nb interface and Al₂O₃ grain boundaries. In the present work, TEM-based methods were used to examine these interfaces at higher spatial resolution, and to provide a more detailed and complete characterization of the crystallographic and chemical characteristics of secondary phases that were present.

The TEM results confirm the absence of silicides (and other secondary phases) in both the sapphire and high-purity Al₂O₃ samples, and the presence of silicides when lower-purity Al₂O₃ is used. Diffraction analysis of a silicide particle at the Al₂O₃–Nb interface revealed that the silicide was the body-centered tetragonal z-Nb₂Si₃ phase. Energy-dispersive spectrometry of silicides both at and slightly away from the interface showed that the range of Nb:Si ratios would place the composition in the two-phase region between Nb₂Si₃ and NbSi₂ on the binary Nb–Si phase diagram at all temperatures from the 1400 °C processing temperature down to room temperature. However,
evidence for a two-phase structure was not found. The silicide incorporates a small amount of Cu, which modifies the phase-stability field, and may extend the stability range of the "\(\alpha\)-Nb\(_5\)Si\(_3\)" structure. Unfortunately, the only isothermal sections available are at temperatures above (1500 °C) or well below (875 °C, 800 °C) the joining temperature. The ternaries also suggest a multiphase microstructure should form. The difficulties of quantitative EDS for small dispersed phases of the type examined and resulting shifts in the apparent composition of the silicide, as well as the role of kinetic barriers to achieving equilibrium in severely constrained systems, emerge as possible explanations for the disparity. Studies of samples subjected to longer anneals and phase-equilibria studies of relevant Nb–Si–Cu compositions at intermediate temperatures would provide useful insights on the origins of the observed microstructures.

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