

pair isomers are necessary to further elucidate the issue.

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Interface Structure and Atomic Bonding Characteristics in Silicon Nitride Ceramics

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Direct atomic resolution images have been obtained that illustrate how a range of rare-earth atoms bond to the interface between the intergranular phase and the matrix grains in an advanced silicon nitride ceramic. It has been found that each rare-earth atom bonds to the interface at a different location, depending on atom size, electronic configuration, and the presence of oxygen at the interface. This is the key factor to understanding the origin of the mechanical properties in these ceramics and will enable precise tailoring in the future to critically improve the materials' performance in wide-ranging applications.

Bulk silicon nitride (Si_3N_4) ceramics have been investigated extensively over the past two decades, largely because their mechanical and physical properties are relevant for many high-temperature applications, including high strength, high decomposition temperature (1900°C), good oxidation resistance, low coefficient of friction, negligible creep, good thermal shock properties, and good resistance to corrosive environments. In addition, thin silicon nitride films and coatings have been studied to understand electri-

cal and thermal conductivity properties. All of these properties, including the processing and sintering behavior, rely on the specific crystal structure, the local chemistry, and the local bonding at the interfaces.

The widespread use of these ceramics as structural components is currently limited by their brittleness, which limits reliability. Specifically, the toughness is too low and the sensitivity to flaws and cracks correspondingly too high, resulting in poor damage tolerance. Such properties can be improved by microstructural and compositional design. Indeed, the relation between microstructure, intergranular phases, and mechanical properties in ceramics is widely acknowledged (1, 2). Silicon nitride ceramic microstructures consist of elongated matrix grains that are randomly oriented, interlocked, and interspersed with a secondary phase (3–8). This intergranular phase in these materials is the key microstructural element that effectively controls most of the material's characteristics, most notably the mechanical properties, and is dependent on the local chemical composition, atomic structure, and bonding characteristics (9–13).

Previous investigations have shown empirically that the material's strength and toughness are markedly affected by the intergranular phase chemistry. An under-

standing of how this phase governs the macroscopic mechanical properties is limited, because the amorphous morphology of the phase and its small dimension make analysis difficult. Recent breakthroughs in scanning transmission electron microscopy (STEM) and associated chemical analysis (14–18) permit probing the local atomic structure and bonding characteristics with a resolution close to 1 Å.

We examine the immediate interface between matrix grains and the amorphous intergranular film, extracting structural and atomic bonding information. In particular, we relate the structure of the interface to the atomic size and electronic structure of the rare-earth elements in the intergranular film. The oxides of rare-earth elements are a very common type of sintering additive in Si_3N_4 , because it has been shown empirically that such additions improve the mechanical properties (12, 13). We investigate the interface between the intergranular phase and the Si_3N_4 matrix using high-angle annular dark-field STEM (HAADF-STEM) for imaging and electron-energy-loss spectroscopy (EELS) (14) for chemical analyses. Experimental EELS and incoherent high-angle ADF (Z-contrast) images (15) were obtained with a monochromated FEI Tecnai F20 STEM operating at 200 kV. The lens conditions in the microscope were defined for imaging and spectroscopy, with a probe size of 0.14 nm. The inner semi-angle of the ADF detector was chosen to be 74 mrad and 110 mrad. Images were acquired with an objective semi-angle of 13.5 mrad. These experimental conditions are enough to minimize the effects of strain fields on the Z-contrast image (16). In this experimental condition, the Z-contrast image allows the structure of the grains to be directly observed, and the image can also be used to position the electron probe for EELS (14).

Previous investigations of these interfaces have focused on two common additives, Y_2O_3 and La_2O_3 (7, 8), showing primarily that there is Y and La segregation in the intergranular phase. Furthermore, the most recent study (8) uses the HAADF-STEM imaging technique described above and first-principles calculations to show that there is

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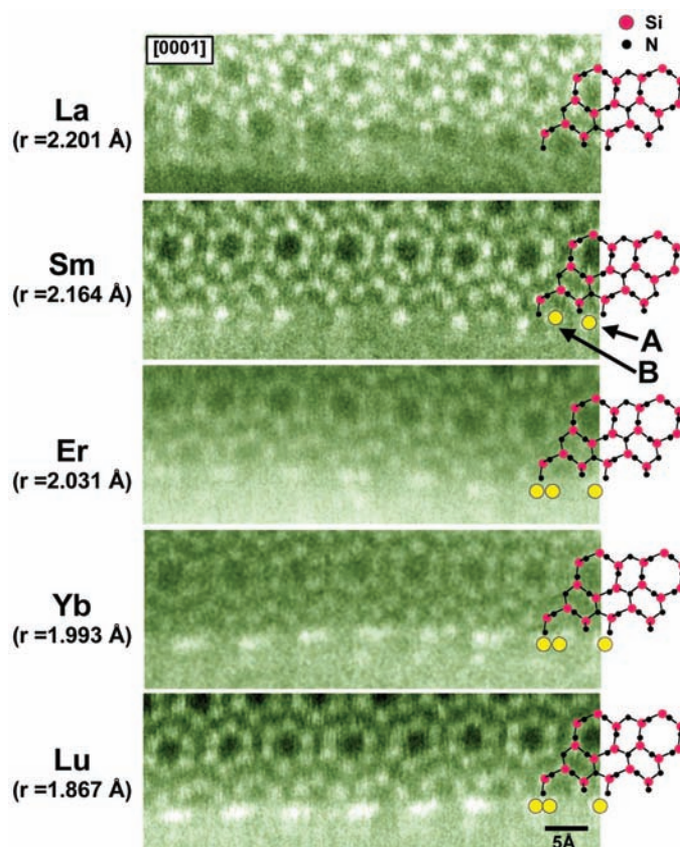
variability in this segregation at the interface between the matrix grains and the intergranular phase. However, the techniques employed did not give information about the electronic configuration or the specific atomic location for the additives at the interface. Our objective is the identification of bonding sites for a larger selection of rare-earth elements of the oxide sintering additives La_2O_3 , Sm_2O_3 , Er_2O_3 , Yb_2O_3 , and Lu_2O_3 and to extend previous investigations by analyzing their electronic configuration and determining the atomic locations of the additives at the interface with the intergranular phase. We take advantage of the “chemically sharp” interface between the intergranular phase and matrix grains caused by a miscibility gap between the Si_3N_4 crystal and any of the rare-earth atoms; that is, these atoms do not substitute for Si_3N_4 host atoms in their atomic positions nor do they sit at interstitial sites in the Si_3N_4 crystal structure.

The HAADF-STEM images in Fig. 1 depict an interface between the intergranular phase and a Si_3N_4 matrix grain for each of the five sintering additive types. The images confirm that the Si_3N_4 crystal structure ends at the interface with the intergranular phase, specifically with open hexagonal rings (7). The attachment of heavy atoms can be proven by the Z-contrast in the images that makes heavier atoms appear brighter. Concomitant large-area EELS measurements performed on these interfaces confirm that the heavy elements indeed segregate to the intergranular phase. Hence, the bright spots seen in the images are indeed related to the heavy atoms and their atomic positions. Accordingly, the atomic attachment at such an interface is different for each of the rare-earth elements (La, Sm, Er, Yb, and Lu) examined here. The STEM images of the sample containing La show no specific or periodic connection occurring between the Si_3N_4 matrix grain and the La atoms. The La atoms are present in the intergranular phase, as reported previously (8), yet they do not arrange into atomic columns that would increase contrast. Additionally, no lateral atomic ordering can be observed at this magnification. Previous investigations have shown that the La atom distribution along the intergranular phase fluctuates, with locations of high and low La atom concentration (8). The distance between the areas of high La concentration are rather far apart and as such cannot be associated with a particular atomic site along the interface. Concomitant theoretical calculations that attempt to determine specific atomic sites are of help; however, in agreement with the present results, a periodic atomic site for La atom attachment cannot be discerned unambiguously from the unaided STEM images.

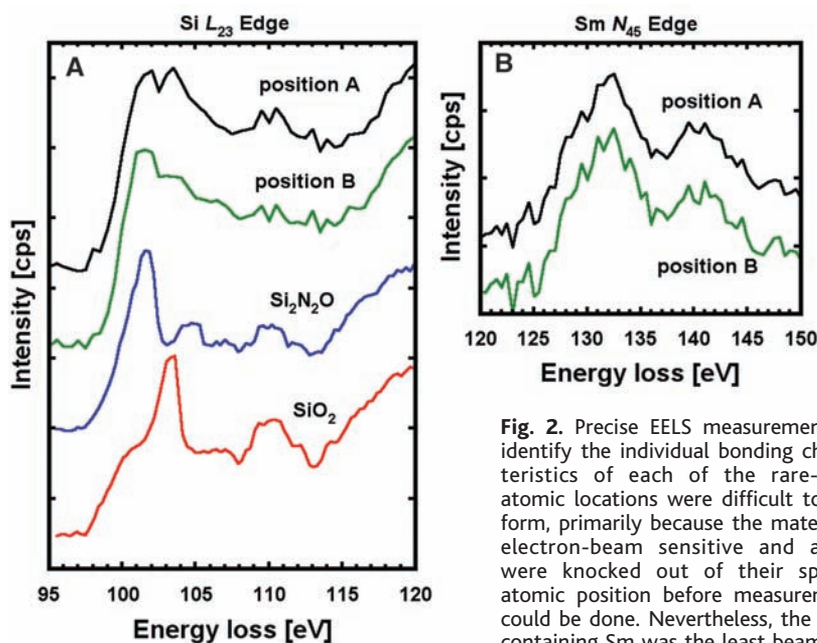
In contrast, the images of the other four compositions containing Sm, Er, Yb, and Lu exhibit very different features. Here, the bright spots that appear at the interface in the amorphous intergranular phase are attributed to columns of Sm, Er, Yb, and Lu atoms, respectively, because of their high contrast. These images clearly show that the atomic bonding of those atoms along the Si_3N_4 prismatic plane is periodic and occurs at two very specific atomic sites, A and B (Fig. 1). Sm atoms bond in single-atom configuration to both positions. This changes with the slightly heavier, but smaller, atoms Er, Yb, and Lu, as they bond in pairs at position A with the pair axis oriented parallel to the prismatic plane. It cannot be determined to what degree the axis is also parallel to the image plane, but observing how close the bright spots are and considering the atomic size of these elements, it follows that the atom-pair axes are inclined and are seen in projection. The atom pairs of Lu, Yb, and Er appear to be separated differently: Lu, $1.43 \pm 0.07 \text{ \AA}$; Yb, $1.46 \pm 0.05 \text{ \AA}$; and Er, $1.48 \pm 0.04 \text{ \AA}$. This growing pair separation can be related to the increase of the valence shell radius from Lu to Er (Fig. 1), which may also be the cause for the occasional pair-splitting that can be observed in the images, i.e., some atom pairs are separated

more than others. For Er and Yb, this results in an occasional larger atom-pair separation of $2.06 \pm 0.03 \text{ \AA}$ and $2.15 \pm 0.1 \text{ \AA}$, respectively. Upon close examination of these split atom pairs, there is indication that beyond the first row of attached atoms there is a second set of rare-earth atoms, reaching even farther into the amorphous intergranular phase—possibly a partial ordering imposed by the Si_3N_4 crystal structure onto the amorphous intergranular phase. However, precise atomic positions beyond the first layer of atomic columns cannot be discerned at this point. These observations demonstrate that the bonding of the rare-earth elements to the interface is controlled by atomic size. The small atoms attach in pairs at specific and periodic sites along the interface. With increasing atom size, the pair separation increases to the point that larger atoms, as in the case of Sm, are not able to bond in pairs at position A and have to adapt a new single-atom periodic bonding configuration. Consequently, one of them has to move to position B. With a further increase in atom size, that is, La atoms, the attachment to the interface changes again, this time forfeiting all periodicity and site-specific atomic bonding. The question remains, however, why the small atoms preferably bond to position A and not B.

Fig. 1. In these STEM images of La-, Sm-, Er-, Yb-, and Lu-doped Si_3N_4 , the matrix grain is oriented along the [0001] zone axis such that the open Si_3N_4 crystal structure is clearly visible at the atomic level and its prismatic plane faces the amorphous intergranular phase. The Si_3N_4 crystal structure ends with open hexagonal rings at the interface with the intergranular phase (7), which indicates that the prismatic plane may contain dangling bonds that can attract atoms from the intergranular phase. The attachment of heavy atoms in the form of atomic columns oriented normal to the image plane—here La, Sm, Er, Yb, and Lu listed with their respective valence-shell radii—is shown by the strong Z-contrast in the images that makes heavier atoms appear brighter. Two distinct atom positions, A and B, can be identified along the interface.



The initial EELS measurements, performed by area-scanning a rather large region of the intergranular films, indicated atom segregation to the interface. Attempting to identify specific atomic bonding configurations, we performed more precise EELS measurements on the Sm-containing sample with a 1.4 Å probe size, because this sample was the least electron-beam sensitive and it showed rare-earth atom attachment to positions A and B. Such a small probe size allows positioning of the electron beam over a single atom column analyzing the very local bonding characteristics (14). The results show the electron loss spectra for the Si L_{23} and the Sm N_{45} edges, respectively, measured on both identified atomic positions (A and B) along the interface (Fig. 2). There is no difference between the spectra at the Sm N_{45} edge, which suggests that the atomic environment and bonding characteristics are the same for the Sm atoms at both locations, that is, Sm exists in the same formal valence state in each location. The difference in the electron-loss spectra from positions A and B is at the Si L_{23} edge. A double peak (102 eV and 103.5 eV) appears in the spectrum taken from atom position A. The first peak can be identified with a Si-N bond, whereas the second peak is associated with a Si-O bond because its position in the spectrum closely coincides with measurements taken on SiO_2 (19). This



measurement. The results for (A) the Si L_{23} and (B) the Sm N_{45} edges identify the individual bonding characteristics at positions A and B. The Si L_{23} edge shows a double peak at position A but not at position B. The first peak can be identified with a Si-N bond because it coincides with measurements taken on the Si_3N_4 crystal structure. Identification of the second peak is achieved by comparing the results to reference spectra obtained from SiO_2 and $\text{Si}_2\text{N}_2\text{O}$. Accordingly, the peak in question can be associated with a Si-O bond. Regarding the electronic environment for the Sm atoms at positions A and B, there is no difference between the spectra at the Sm N_{45} edge.

indicates that the atomic environment around the terminating Si atoms on the Si_3N_4 prism plane that surround position A is most likely an oxygen atom. In contrast, the atomic environment of the Si atoms on the prism plane that surround position B appears to have no oxygen atoms involved, only nitrogen.

The observation that the Er, Yb, and Lu atom pairs preferably bond to position A suggests that the interaction of the rare-earth atom with the oxygen at site A plays a critical role in controlling atom attachment at the interface. Thus, a combination of both factors, that is, the difference in atomic size and the presence of oxygen, govern atom attachment and site selection along the interface. This results in different atom concentrations at the individual positions A and B. The rare-earth atom concentration is high for position A and comparably low at position B. The interaction of electronic configuration, atom size, and position of the rare-earth atoms on the prismatic plane is currently being investigated by atomic-structure calculations of this interface.

This information about the specific atomic structure and bonding characteristics in advanced ceramics has been missing for many years and should now aid the development of improved ceramics in many ways. Most important, it will assist in understanding how ceramic microstructures evolve during fabrication, especially how grain growth and

microstructural evolution is affected by different sintering additives at the atomic level. Control over evolution and the ability to precisely tailor ceramic microstructures is critical in improving mechanical properties of ceramics, which invariably limits their use. From previous empirical investigations (1, 2, 9, 10, 12, 13), it is known that ceramics' fracture is highly sensitive to the intergranular phase chemistry and therefore to its atomic structure and bonding characteristics. The identification of precise atom locations from the present study should also prove invaluable for atomic-structure calculations. Such computational methods (20–26) seek to calculate individual atomic bonding strengths so as to explain the particular fracture mechanisms that occur along and across the intergranular phase. However, lacking essential experimental information about the atomic structure and bonding characteristics, such calculations have been incomplete.

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