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Summary Technical Report

to

NASA-Lewis Research Center

MECHANISMS OF DEVITRIFICATION OF GRAIN BOUNDARY GLASSY PHASES IN Si$_3$N$_4$ MATERIALS

NASA Grant #NSG 3254

Submitted by

L. L. Hench, Professor
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Department of Materials Science and Engineering
University of Florida
Gainesville, Florida 3261

December 17, 1982
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OVERVIEW

The objectives of this grant were: 1) to develop means for analyzing changes in the grain boundary (g.b.) phases of Si$_3$N$_4$, 2) to determine the effects of composition and thermal history on devitrification of the g.b. phases, and 3) relate devitrification of the g.b. phases of Si$_3$N$_4$ to mechanical behavior and oxidation sensitivity of the material. The series of six papers submitted for publication presented in this Summary Report describe our progress towards achieving our three objectives.

The first paper is a thorough review of the phase relationships that occur within the grain boundaries of Si$_3$N$_4$ containing various densification aids. Comparisons of the effects of MgO, Y$_2$O$_3$, CeO$_2$, and Y$_2$O$_3$ + Al$_2$O$_3$ are made in terms of the phase equilibria of the Si$_3$N$_4$ + SiO$_2$ + additive compositional system. Two new equilibrium phase diagrams for the Si$_3$N$_4$-SiO$_2$ and Y$_2$O$_3$ and Si$_3$N$_4$-SiO$_2$-Ce$_2$O$_3$ systems are presented in this work.

An experimental comparison of the effects of Y$_2$O$_3$ vs CeO$_2$ densification aids on the fracture surfaces of Si$_3$N$_4$ is made in the second paper. Auger electron spectroscopy showed that both oxides are concentrated within the fracture surface. Scanning electron microscopy showed evidence that Si$_3$N$_4$ with CeO$_2$ formed an intergranular structure of fine grained oxynitride reaction products, as predicted by phase equilibria, whereas the Y$_2$O$_3$ containing sample showed evidence of an intergranular glassy phase.

Several surface analysis methods were used in the third paper to show that Si$_3$N$_4$ fracture surfaces with Y$_2$O$_3$ possess a higher concentration of oxygen than the bulk and increasing concentrations of Y$_2$O$_3$ and Al$_2$O$_3$ increases the oxygen content of the intergranular phase. This paper provides direct evidence from fracture surfaces that certain compositions
of Si$_3$N$_4$ + Y$_2$O$_3$ + Al$_2$O$_3$ result in an amorphous grain boundary phase which is resistant to devitrification whereas other compositions devitrify more easily.

It was shown in the fourth paper of this series that devitrification of the amorphous g.b. phase of Si$_3$N$_4$ + 15% Y$_2$O$_3$ + 2% Al$_2$O$_3$ could be achieved by increasing the vacuum heat treating temperature to 1200°C. A new analytical method, Fourier transform infrared reflection spectroscopy (FTIRRS), was developed to analyze post-fracture surfaces in order to achieve this result.

A comparison of the effects of various levels of Al$_2$O$_3$ additions to the Si$_3$N$_4$ + Y$_2$O$_3$ system on sensitivity to grain boundary phase devitrification is presented in the fifth paper. It was shown that heat treatment at 1000°C produced some crystallization of Y$_2$Si$_2$O$_7$ in samples containing 2, 4, and 6% Al$_2$O$_3$ whereas material with 8% Al$_2$O$_3$ developed considerably more Y$_2$Si$_2$O$_7$ and some 10-9-1 phase. A 1200°C heat treatment produced more devitrification for all compositions. However, the 6% Al$_2$O$_3$ samples showed the most Y$_2$Si$_2$O$_7$ and 10-9-1 phases with the 2% Al$_2$O$_3$ sample second. The 4% Al$_2$O$_3$ additions result in the most resistance to devitrification.

FTIRRS analyses of the heat treated series showed that there is little shift of the spectra of the fracture surfaces of the 4% Al$_2$O$_3$ material whereas fracture surfaces of the 2% Al$_2$O$_3$ samples were affected considerably by devitrification. Little differences were noted for fracture surfaces of the 6 and 8% Al$_2$O$_3$ samples due to devitrification. However, the results that show splitting of primary Si-N molecular stretching vibration peaks of fracture surfaces confirm that crystallization is occurring in the grain boundary phase.

Both the compositional changes and the heat treatments affect room temperature and high temperature strengths of these materials. Heat
treatment of the 8% Al$_2$O$_3$ at 1200°C produced the highest room temperature (113,280 psi) and elevated temperature (20,010 psi) strengths of the series studied.

Finally, in the sixth paper of the series the oxidation sensitivity of Si$_3$N$_4$ + 15 w/o Y$_2$O$_3$ materials with 2, 4, 6, and 8% Al$_2$O$_3$ are compared at temperatures as low as 1000°C. It was found that concentrations of Al$_2$O$_3$ > 4% greatly retard the rate of oxidation and alter the mechanism of surface attack by promoting the surface formation of a glassy layer which contains mixed oxynitride bonds. The glassy layer retards heterogeneous oxidation attack and reduces the effect of an oxidation transition temperature between 1000°C and 1100°C for these materials.
Si$_3$N$_4$ GRAIN BOUNDARY PHASES

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Introduction

In order to fabricate fully dense hot-pressed Si$_3$N$_4$, currently a prime candidate for heat engine and gas turbine components, additives are required for sintering or hot-pressing. During the last several years, considerable experimental efforts have been made to improve the high-temperature properties of Si$_3$N$_4$ by: 1) synthesizing high-purity α-Si$_3$N$_4$ powders for use as starting materials; 2) using oxide additives which would allow the formation of refractory grain boundary phases, and 3) modifying processing techniques to improve product homogeneity. The use of high-purity Si$_3$N$_4$ powder improves high-temperature properties as well as increases oxidation resistance by eliminating undesirable impurity constituents such as SiO$_2$, CaO and alkaline oxides, and to a lesser extent Al$_2$O$_3$ and Fe$_2$O$_3$. This results in an increase in the refractoriness of the grain boundary phases formed during densification at elevated temperatures. However, the production of high-purity Si$_3$N$_4$ powder in bulk quantities may not be cost effective. Also, high-temperature properties, although improved, are still limited by the reaction products formed by the additives being used in densification. Thus, it is especially important to understand the behavior of additives which can simultaneously promote densification and enhance the refractoriness of the grain boundary phases.

Early attempts to hot-press Si$_3$N$_4$ to full density used a small amount of MgO as a densification aid which resulted in formation of a liquid phase at moderately high temperatures and promoted densification
of the sample. However, a major problem preventing the widespread use of MgO as a densification aid was the decrease in strength of the hot-pressed material above 1000°C. Evidence obtained with commercially hot-pressed Si₃N₄ containing MgO indicated that the decrease in strength was primarily due to the formation of the grain boundary glassy phase developed during hot-pressing. The viscosity of a silicate glassy phase decreases rapidly with increasing temperature and, as a result, the liquid starts to flow at elevated temperatures causing grain boundary sliding and ultimate mechanical failure of the Si₃N₄ material. Further understanding of the nature, distribution and characteristics of the grain boundary glassy phase has led to the conclusion that impurities in Si₃N₄ powders, particularly, CaO and alkaline oxides, decrease the viscosity and lower the softening temperature of the grain boundary phase.

Other oxides besides MgO which are known to be effective in hot-pressing or sintering Si₃N₄ include Y₂O₃. The improved properties of hot-pressed Si₃N₄ with Y₂O₃ have been attributed to the formation of highly refractory grain-boundary phases identified as yttrium silicon oxynitrides. However, addition of certain compositional ranges of Y₂O₃ degrade the oxidation resistance of Si₃N₄ materials. This undesirable effect on oxidation resistance can be eliminated by the use of CeO₂ or combined Y₂O₃+Al₂O₃ mixtures.

Because of the broad range of oxides studied as densification aids it is becoming important to establish the principles governing their behavior. The purpose of this article is to summarize the findings of the characteristics of the grain-boundary phase in dense Si₃N₄ based
materials and relate those characteristics to the phase equilibria of the systems. It will be shown that certain phase equilibria principles can be used to improve the refractoriness of the grain boundaries of these materials. For a review of other aspects of Si$_3$N$_4$ processing and properties the reader is referred to Messier and Croft (1).

**MgO Additives**

Hot-pressing of Si$_3$N$_4$ with MgO as an additive is one of the earliest approaches taken to produce dense Si$_3$N$_4$ materials (2). Early attempts of understanding the densification mechanisms of Si$_3$N$_4$ containing MgO have been made by Wild et al (3) and Terwillinger and Lange (4). These workers found that at 1 atm. of N$_2$ pressure, MgO reacts with SiO$_2$ present in the starting materials or formed by the decomposition of Si$_3$N$_4$ during hot-pressing at moderately high temperatures, to form forsterite, Mg$_2$SiO$_4$. The silicate phase becomes a liquid at hot-pressing temperatures and promotes densification of Si$_3$N$_4$ by a solution-reprecipitation process. This conclusion was confirmed by direct observation by Drew and Lewis (5) of the glassy phase in the grain-boundaries of Si$_3$N$_4$ containing MgO using transmission electron microscopy (TEM).

Hot-pressing of Si$_3$N$_4$ with 5% MgO at higher nitrogen pressure (>10 atm) was performed by Mitomo (6) with a view to suppress thermal decomposition of Si$_3$N$_4$, thereby permitting higher densification temperatures. He proposed that densification proceeds in two stages, namely, by initial particle rearrangement followed by a solution-reprecipitation process. Soon thereafter phase relations in the system Si$_3$N$_4$-SiO$_2$-MgO and their significance to strength and oxidation resistance of Si$_3$N$_4$
were investigated by Lange (7) who observed that the high temperature strength of Si₃N₄ containing MgO is strongly dependent on the MgO/SiO₂ molar ratio. It is this molar ratio which determines the amount of liquid phase present as predicted by the eutectics within the ternary compositional system.

The role of alkaline-doped MgSiO₃ on the densification process of Si₃N₄ has been studied by Kossowsky (8) who reported that the alkaline content should be kept below 50-100 ppm for optimum strength. Kossowsky's study used Auger electron spectroscopy (AES) to show that impurity constituents such as Ca, Mg, Na and K concentrated in the grain-boundaries of commercially hot-pressed Si₃N₄. Similar findings were reported by Hofmann et. al (9) using AES to study the fracture surfaces of hot-pressed Si₃N₄ containing MgO. Powell and Drew (10) also carried out AES studies on fracture surfaces of an Si₃N₄-7% MgO hot-pressed material and observed the presence of significant amounts of Ca, Mg and oxygen at the grain-boundaries. They estimated the probable chemical composition of the intergranular glassy phase to be 0.03 CaO: +0.1 MgO: +2 SiO₂ and attributed the reduction of strength of the hot-pressed material at temperatures above 1000°C to the decrease in viscosity of the grain boundary phase due to the presence of CaO.

A critical review of viscosity of glasses in the system CaO-MgO-Al₂O₃-SiO₂ was reported by Turkdogan and Bills (11) who observed that the viscosity of a CaO: Mg: 2 SiO₂ melt decreases rapidly within the temperature range between 1200°C and 1600°C. Measurements of the viscosity of the grain boundary glassy phase in hot-pressed Si₃N₄ containing MgO and CaO by an internal friction technique by Mosher et. al (12) confirm
that high temperature deformation occurs by this mechanism. The effect of impurity phases, particularly CaO, alkaline oxides, Al₂O₃ and Fe₂O₃ on the high-temperature mechanical properties of 5% MgO fluxed hot-pressed Si₃N₄ was also studied by Iskoe et. al(13). These workers observed that CaO, Na₂O and K₂O reduce the high-temperature strength of Si₃N₄ but Al₂O₃ and Fe₂O₃ have little effect.

Direct observations of the intergranular glassy phase in hot-pressed Si₃N₄ by transmission electron microscopy (TEM) have been carried out by several workers(5,14-17). Clarke and Thomas(14) used a lattice fringe imaging technique with TEM to identify the intergranular phases and reported that the second phase does not exist as a continuous wetting layer at the grain boundaries, but is generally localized at some of the multiple grain junctions and, occasionally, as a very thin layer (<10 Å) between the grains. This observation, however, contradicts the findings of Lou et. al(15) who reported that a continuous grain boundary glassy layer exists in the commercially hot-pressed Si₃N₄. The discrepancy was attributed by Lou et. al(16) to the difference in the TEM techniques used by the two groups of workers. Krivanek et. al(17) employed several analytical techniques to study the nature, distribution and composition of the grain-boundary phases in commercially hot-pressed Si₃N₄ and confirmed that a thin intergranular glassy layer exists at most grains and interphase boundaries. Further, the concentration of various impurity constituents, particularly CaO, in the glass exceeds several times their overall concentration so that even small amounts of impurity elements can have a disproportionately large influence on the properties of the glassy intergranular phase.
Lange (18) extended his phase equilibria work (7) to include the melting behavior of several compositions in the subsystem Si$_3$N$_4$-Si$_2$N$_2$O-Mg$_2$SiO$_4$ in which a ternary eutectic close to Mg$_2$SiO$_4$ was proposed as shown in Fig. 1. The temperature of the ternary eutectic was determined to be 1515°C. The presence of CaO as an impurity phase results in the further lowering of the melting temperature of the intergranular glassy phase to 1325°C. However, in a recent publication, Clarke et al. (19) examined the intergranular phases in MgO fluxed hot-pressed Si$_3$N$_4$ using several complementary TEM techniques and report that the composition of the intergranular non-crystalline phase can not be related to the ternary eutectic composition studied by Lange (18) but corresponds to a composition in the Si$_2$N$_2$O-SiO$_2$-MgSiO$_3$ phase field near the MgO-SiO$_2$ tie-line. On subsequent examination of the non-crystalline intergranular phase, Clarke et al. (20) further observed that it undergoes phase separation and possible crystallization to Si$_2$N$_2$O and MgSiO$_3$. The compositional range determined for the phase separation is close to the known miscibility gap existing in the pseudobinary system MgO-SiO$_2$.

Y$_2$O$_3$ Additive

The effectiveness of Y$_2$O$_3$ as an additive for hot-pressing Si$_3$N$_4$ has been discussed by several workers (21-25). Reaction between Si$_3$N$_4$ and Y$_2$O$_3$ was studied by Tsuge et al. (22) who identified a crystalline phase of composition Si$_3$N$_4$.Y$_2$O$_3$ in the hot-pressed samples and tentatively postulated the existence of another phase of composition Si$_3$N$_4$.2Y$_2$O$_3$. Preliminary property measurements at 1315°C reported by Gazza (23) showed that the high-temperature mechanical properties of Si$_3$N$_4$ can be significantly
improved by using $Y_2O_3$ as a densification aid. Rae et al.\(^{(24)}\) observed that densification of $Si_3N_4$ in presence of $Y_2O_3$ is enhanced by the formation of a transient silicon yttrium oxynitride melt at elevated temperatures and further reaction with more $Si_3N_4$ allows the formation a refractory silicon yttrium oxynitride phase, $Y_4(Si_2O_3N_4)$ which can accommodate impurity constituents into its structure that would otherwise degrade the high-temperature properties of $Si_3N_4$.

In an effort to establish the existence various silicon yttrium oxynitrides and their significance to hot-pressing of $Si_3N_4$ with $Y_2O_3$ additives, phase equilibria in the system $Si_3N_4$-$SiO_2$-$Y_2O_3$ have been investigated by several groups\(^{(26-29)}\). Although some disagreement still exists on the number and compositions of the various compounds occurring in the system, it is generally accepted that there are four pseudo-ternary compounds with compositions of $Y_2Si_3O_3N_4$, $YSiO_2N$, $Y_4Si_2O_7N_2$ and $Y_{10}(SiO_4)_6N_2$. The crystal structure of $Y_2Si_3O_3N_4$ was determined by Horiuchi and Mitomo\(^{(30)}\) who reported that the compound is isostructural with the melilite silicates: akermanite, $Ca_2MgSi_2O_7$ having a tetragonal symmetry. The crystal structure of $YSiO_2N$ was related to the $\alpha$-wollastonite, $CaSiO_3$, by Jack\(^{(27)}\) and the powder X-ray diffraction pattern has been indexed by Morgan et al.\(^{(31)}\) on the basis of the hexagonal pseudowollastonite structure. The compound $Y_4Si_2O_7N_2$ was reported by Wills et al.\(^{(26)}\) to be isostructural with the silicates of wohlerite-cuspidine, $Ca_4Si_2O_7F_2$, having a monoclinic structure. The compound $Y_{10}(SiO_4)_6N_2$ was reported to be isostructural with hexagonal fluorapatite, $Ca_5(PO_4)_3F$, by Thompson\(^{(32)}\) and the indexed X-ray powder diffraction pattern of the compound was presented by Wills et al.\(^{(26)}\).
With the available data on compound formation and compatibility relations existing between the various phases in the pseudo-ternary system $\text{Si}_3\text{N}_4-\text{SiO}_2-\text{Y}_2\text{O}_3$, a phase diagram has been deduced as shown in Fig. 2. Partial data on the liquid formation in the system are due to Gauckler et al.\(^{(29)}\) who observed two-liquid forming regions at 1550°C. The first is in the $\text{SiO}_2$-rich region within the compatibility-triangle $\text{Si}_2\text{N}_2\text{O}-\text{Y}_2\text{Si}_2\text{O}_7-\text{SiO}_2$ where compositions exhibited signs of bloating during annealing. A similar effect has been reported by Wills et al.\(^{(26)}\). The second liquid forming region was located in the compatibility-triangle $\text{Si}_3\text{N}_4-\text{Y}_2\text{Si}_2\text{O}_7-\text{Y}_{10}(\text{SiO}_4)\_6\text{N}_2$. Melting behavior of compositions within this area showed liquid formation at a temperature between 1480° and 1520°C.

The liquids in both the regions which form during hot-pressing become glass when cooled. From the phase equilibria data, it is generally understood that during hot-pressing $\text{Y}_2\text{O}_3$ reacts with $\text{Si}_3\text{N}_4$ and surface $\text{SiO}_2$ to form a liquid which allows densification of the materials. As the reaction proceeds, the liquid combines with excess $\text{Si}_3\text{N}_4$ to give one or more of the refractory silicon-yttrium-oxynitrides. Unreacted liquid cools to give a glass which remains in the grain-boundaries and impairs the high-temperature strength of $\text{Si}_3\text{N}_4$. TEM studies on identification and characterization of grain boundary phases in hot-pressed $\text{Si}_3\text{N}_4$ with 10% $\text{Y}_2\text{O}_3$ additive have been reported by Clarke and Thomas\(^{(33)}\). Detailed microstructural analysis has revealed that, in addition to the yttrium silicon oxynitride phase located at the multiple $\text{Si}_3\text{N}_4$ grain junctions, there exists a thin, probably noncrystalline
boundary phase separating the $Si_3N_4$ and the oxynitride grains. However, Clarke and Thomas could not determine the composition of this third phase and suggested that the presence of the amorphous film between the $Si_3N_4$ and crystalline yttrium silicon oxynitride grains is probably responsible for the decrease of the mechanical strength of the hot-pressed material.

The effect of the grain-boundary oxynitride phases on the oxidation resistance of $Si_3N_4$ has also been studied (28). Lange et al (28) observed that compositions within the $Si_3N_4-Y_2Si_2O_7-Y_2O_3$ phase field exhibited relatively poor oxidation resistance at 1000°C and above. The problem was attributed to the accelerated linear oxidation behavior of the grain boundary yttrium silicon oxynitride phases, particularly, $Si_3Y_2O_3N_4$.

At temperatures of 1000°C and above, the oxynitrides rapidly form non-protective and porous oxide scales causing catastrophic degradation and eventual disintegration of the material. In contrast, compositions within the compatibility triangle $Si_3N_4-Si_2N_2O-Y_2Si_2O_7$, in which none of the quaternary oxynitride compounds occur, exhibit excellent oxidation resistance. It was suggested that this was due to the compatibility of $SiO_2$, the oxidation product of $Si_3N_4$, with the secondary phases, $Y_2Si_2O_7$ and $Si_2N_2O$ and the relatively high melting temperature of the eutectic in this compositional area. However, Weaver and Lucek (34) disagreed with this finding and stress the need for property optimization through composition variation in hot-pressed $Si_3N_4$ with $Y_2O_3$ additives. These workers report that no deterioration of strength or oxidation resistance could be observed if the $Y_2O_3$ content was kept below a certain critical
limit. Based on their experimental data, it was suggested that 8% Y2O3 addition represents an optimized compositional level in which substantial improvements in both the strength and oxidation resistance have been observed.

CeO2 Additive

The use of CeO2 as a hot-pressing aid for densifying Si3N4 powder has been reported by several workers\(^{(35-39)}\). It is generally agreed that the addition of CeO2 significantly enhances densification of Si3N4 with improvements in high-temperature properties. In these studies it was assumed that CeO2 and SiO2 thus formed further combine with Si3N4 to give several quaternary cerium silicon oxynitrides. The existence of a compound having the chemical formula 3Ce2O3·2Si3N4 has been first reported by Wills and Cunningham\(^{(40)}\) who assigned an orthorhombic structure for the compound. However, Morgan and Carroll\(^{(41)}\) and Thompson\(^{(42)}\) re-examined the X-ray diffraction pattern for the compound and reported that the correct chemical composition of the compound is CeSiO2N having a pseudo-hexagonal structure similar to YSiO2N\(^{(31)}\).

Mah et al\(^{(43)}\) studied the densification behavior of Si3N4 with both CeO2 and Ce2O3 additives and suggest that a nitrogen containing cerium orthosilicate, Ce4.67(SiO4)3O, which forms as a grain-boundary glassy phase, was mainly responsible for the densification of Si3N4. Two cerium silicon oxynitrides, Ce2Si6O3N8 and Ce4Si2O7N2 were identified and the X-ray powder diffraction data for the compounds have been
presented. Guha et al. (44) hot-pressed Si$_3$N$_4$ with CeO$_2$ additive at temperatures between 1400° and 1750°C and identified a nitrogen containing apatite phase, Ce$_5$(SiO$_4$)$_3$N analogous to Y$_5$(SiO$_4$)$_3$N and the previously reported compound CeSiO$_2$N as reaction products. Evidence obtained by X-ray powder diffraction of the hot-pressed compositions indicated that both the oxynitrides constitute pseudo-binary tie-lines with Si$_3$N$_4$ in the system Si$_3$N$_4$-SiO$_2$-Ce$_2$O$_3$. The crystallographic data proposed by Mah et al. (43) for the compound Ce$_4$Si$_2$O$_7$N$_2$ have been re-examined by Morgan (45) and Guha (46) both of whom have concurrently assigned a monoclinic cuspidine type (Ca$_4$Si$_2$O$_7$F$_2$) unit-cell for the compound.

Subsolidus phase relations in the system Ce-Si-O-N have been reported by Jack (46) who proposed four quaternary cerium oxynitrides that are isostructural with the yttrium analogues plus an additional phase, Ce$_2$O$_3$.2Si$_3$N$_4$ which has no yttrium analogue. Lange (48,49) has also proposed a tentative phase diagram for the pseudo-ternary system Si$_3$N$_4$-SiO$_2$-Ce$_2$O$_3$ but failed to mention the compound Ce$_2$O$_3$.2Si$_3$N$_4$ and did not observe the compound Ce$_2$Si$_3$O$_3$N$_4$ and Ce$_4$Si$_2$O$_7$N$_2$ although these are expected in the diagram. The exact composition of the nitrogen apatite phase represented by Lange (49) in the diagram as Ce$_{10}$(SiO$_{3.67}$N$_{0.33}$)$_2$ (Ref. 47) and Ce$_5$(SiO$_4$)$_3$N still remains uncertain. However, the existence of a new group of silicon lanthanide oxynitrides of the general formula Ln$_5$(SiO$_4$)$_3$N which are isostructural with the hexagonal fluorapatite, Ca$_5$(PO$_4$)$_3$F has been confirmed by Hamon et al. (50) and the crystal structure of nitrogen apatites with particular reference to Y$_5$(SiO$_4$)$_3$N was
discussed by Jack (51). The nitrogen containing Ce-apatite phase, 
$\text{Ce}_5(\text{SiO}_4)_3\text{N}$ has been prepared by Guha (52) using solid state reaction techniques and the X-ray powder diffraction data for the compound which are similar to those of $\text{Y}_5(\text{SiO}_4)_3\text{N}$ were presented.

Based on the foregoing data on the chemical composition of the various phases in the pseudo-ternary system $\text{Si}_3\text{N}_4-\text{SiO}_2-\text{Ce}_2\text{O}_3$ and the compatibility relations between them, a phase diagram is deduced as shown in Fig. 3. As mentioned by the previous workers (47,48), the close resemblance between this and $\text{Si}_3\text{N}_4-\text{SiO}_2-\text{Y}_2\text{O}_3$ systems suggests that the densification behavior and ultimate properties of $\text{Si}_3\text{N}_4$ hot-pressed with ceria will be similar to those with yttria. The results obtained so far on the hot-pressing behavior of $\text{Si}_3\text{N}_4$ with $\text{Y}_2\text{O}_3$ and $\text{CeO}_2$ additives appear to confirm this supposition. Thus, it is now known that during hot-pressing, both of these oxides form a nitrogen containing apatite oxynitride phase which eventually allows the formation of a liquid phase in the grain-boundaries of $\text{Si}_3\text{N}_4$ and promotes densification of the material. Further reaction of the liquid with more $\text{Si}_3\text{N}_4$ yields one or more grain boundary refractory oxynitride phases which leads to improvement in high-temperature properties of the densified material. However, the advantage of $\text{CeO}_2$ over $\text{Y}_2\text{O}_3$ as an additive is that for each mole of $\text{CeO}_2$ added, corresponding molar proportions of $\text{Ce}_2\text{O}_3$ and $\text{SiO}_2$ will be formed according to the reaction:

$$\text{Si}_3\text{N}_4 + 12 \text{CeO}_2 \rightarrow 6 \text{Ce}_2\text{O}_3 + 3 \text{SiO}_2 + 2\text{N}_2$$  \hspace{1cm} (1)

Thus, the reaction of $\text{Si}_3\text{N}_4$ with $\text{CeO}_2$ allows the formation of the nitrogen containing Ce-apatite phase through reaction with the intermediate $\text{Ce}_2\text{O}_3$ phase. The Ce-apatite phase, in turn, will form the liquid phase
Fig. 3

SiO₂

Si₂ON₂

Ce₂Si₂O₇

Ce₄.₆₇(SiO₄)₃O

Ce₂SiO₅

Ce₅(SiO₄)₃N

CeSiO₂N

Ce₄Si₂O₇N₂

Si₃N₄ Ce₂SiO₃N₈ Ce₂Si₃O₇N₄ Ce₂O₃
required for densification of \( \text{Si}_3\text{N}_4 \). Further, due to the compositional limit for \( \text{SiO}_2 \) which is always present in all the compositions containing \( \text{CeO}_2 \), the quaternary oxynitrides located on the \( \text{Si}_3\text{N}_4-\text{Ce}_2\text{O}_3 \) join cannot be formed. This is in contrast to the case of \( \text{Y}_2\text{O}_3 \) in which the oxynitride \( \text{Si}_3\text{N}_4\cdot\text{Y}_2\text{O}_3 \) is known to be mainly responsible for the catastrophic oxidation of \( \text{Si}_3\text{N}_4 \) at 1000°C. Therefore, for the same concentrations of additive, \( \text{Si}_3\text{N}_4 \) densified with \( \text{CeO}_2 \) should be much more oxidation resistant than \( \text{Si}_3\text{N}_4 \) densified with \( \text{Y}_2\text{O}_3 \).

\[ \text{Y}_2\text{O}_3/\text{Al}_2\text{O}_3 \text{ Additive} \]

Silicon nitride densified with \( \text{Y}_2\text{O}_3/\text{Al}_2\text{O}_3 \) mixtures develops an extensive intergranular glassy phase which is very effective for obtaining highly dense products with excellent flexural strength. Early attempt to sinter \( \text{Si}_3\text{N}_4 \) with \( \text{Y}_2\text{O}_3 \) in which \( \text{Al}_2\text{O}_3 \) was incorporated as an impurity phase during ball milling was made by Tsuge et al\(^{(53)}\). They obtained a higher flexural strength for the material than previously reported data for \( \text{Si}_3\text{N}_4 \) containing \( \text{MgO} \). The extensive glassy phase formed at high temperatures crystallized to refractory grain-boundary phases containing \( \beta-\text{Si}_3\text{N}_4 \) and \( \text{Si}_3\text{N}_4\cdot\text{Y}_2\text{O}_3 \) during cooling. Wills\(^{(54)}\) has investigated the reaction of \( \text{Si}_3\text{N}_4 \) with \( \text{Y}_2\text{O}_3 \) and \( \text{Al}_2\text{O}_3 \) and suggested that a new phase exists with the probable composition \( 5\text{Y}_2\text{O}_3\cdot\text{Si}_3\text{N}_4\cdot\text{Al}_2\text{O}_3 \) content in the mixtures he studied. Morgan\(^{(55)}\), however, has disagreed with the occurrence of any new phase in the system and emphasizes that a solid solution series, \( \text{Y}_4\text{Al}_{2-x}\text{Si}_{x}0_{9-x}N_x \), exists instead.
Venables et al. (56) used 10% of the eutectic composition of $\text{Y}_2\text{O}_3$-$\text{Al}_2\text{O}_3$ as an additive and reported excellent sintering characteristics of $\text{Si}_3\text{N}_4$ due to the formation of a large volume of liquid glassy phase which completely wetted the solid and migrated rapidly through the $\text{Si}_3\text{N}_4$ grains. Jack (47) supported these findings and postulated a wide range of solid solution depicted as $\beta$-sialon in the pseudo-ternary system $\text{Si}_3\text{N}_4$-$\text{Y}_2\text{O}_3$-$\text{Al}_2\text{O}_3$. Compositions within the solid solution range showed none of the catastrophic oxidation at 1000°C experienced with some of the $\text{Si}_3\text{N}_4$-$\text{Y}_2\text{O}_3$ compositions without $\text{Al}_2\text{O}_3$. It was further suggested that the lowest liquidus in the $\text{Y}_2\text{O}_3$-$\text{SiO}_2$-$\text{Al}_2\text{O}_3$ system is at approximately 1350°C above which the oxidation of yttrium sialons involves liquid formation and hence becomes more pronounced. Milberg and Miller (57) prepared a composition containing 53% $\text{Si}_3\text{N}_4$, 27% $\text{Al}_2\text{O}_3$ and 20% $\text{Y}_2\text{O}_3$ by sintering for 3 hrs at 1600°C in an argon atmosphere. The XRD pattern showed the presence of $\beta$-sialon lines superimposed on a halo typical of a noncrystalline substance. When the same material was heated at 1200°C for 185 hr., the diffraction pattern showed the presence of $\text{Y}_2\text{Si}_2\text{O}_7$ which indicated that a glassy phase initially present in the material devitrified to yttrium disilicate during the heat-treatment. The preparation of Si-Y-Al-O-N glasses was reported by Shillito et al. (58) and hardness values obtained for these glasses were comparable to those of a Si-Y-Al-O glass.

An extensive study of the effect of $\text{Al}_2\text{O}_3$ on the mechanical properties of $\text{Si}_3\text{N}_4$ containing $\text{Y}_2\text{O}_3$ (or $\text{CeO}_2$) has been performed by Smith and Quackenbush (59). They found that $\text{Al}_2\text{O}_3$ preferentially goes into
the liquid phase formed at the sintering or hot-pressing temperatures and suppresses crystallization, thereby promoting glass formation. At elevated temperatures, this glassy phase causes structural degradation by slow crack growth and intergranular fracture. When phase purity is maintained, the decrease in Al$_2$O$_3$ content enhances crystallization of refractory grain-boundary phases containing Y$_2$Si$_2$O$_7$ and Y$_5$(SiO$_4$)$_3$N which were mainly responsible for the substantial improvement in the high-temperature strength of the Si$_3$N$_4$ material. In a concurrent publication, Quackenbush and Smith (60) reported the oxidation behavior of Si$_3$N$_4$ with Y$_2$O$_3$ and Al$_2$O$_3$ additives and observed that the Al$_2$O$_3$ bearing materials show good oxidation resistance at 1000°C. Above this temperature, oxidation of the materials follow parabolic kinetics and is controlled by oxygen diffusion through the surface silicate layer.

In a recent study, Hench et al (61) have employed several surface analytical techniques (XRD, IRRS and AES) to examine the fracture surfaces of Si$_3$N$_3$ hot-pressed with Y$_2$O$_3$ and Al$_2$O$_3$ densification aids. These workers have observed that the fracture occurs preferentially within the oxygen enriched grain boundaries. An increase in Y$_2$O$_3$ content increased the concentration of oxygen within the fracture surface. Additions of 13 to 15% Y$_2$O$_3$ and 6% Al$_2$O$_3$ to Si$_3$N$_4$ enhanced the formation of an amorphous grain-boundary phase which failed to devitrify when heat-treated for 10 hrs. at 1000°C in vacuum.

In a related study Hench, et al (62) investigated the oxidation resistance of Si$_3$N$_4$ containing 15 w/o Y$_2$O$_3$ and 2, 4, 6 and 8% Al$_2$O$_3$. Oxidation was followed at temperatures as low as 1000°C by use of
infrared reflection spectroscopy. It was found that Al$_2$O$_3$ additions of 4% or greater significantly retard oxidation of Si$_3$N$_4$+15% Y$_2$O$_3$ even though this composition is in the field where the destructive Si$_3$Y$_2$O$_3$N$_4$ phase can form. The oxidation protection is due to formation of a surface layer containing mixed silicon oxynitride bonds which retard a heterogeneous mode of attack of the Si$_3$N$_4$ structure. The mechanisms and the Al$_2$O$_3$ compositional effect are the same at both 1000°C and 1100°C confirming the previous conclusion(60) that development of surface glassy phases greatly reduce the importance of an oxidation transition temperature for these materials.

Acknowledgements

The authors gratefully acknowledge partial financial support of NASA Grant #NS63254.
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GRAIN-BOUNDARY PHASES IN HOT-PRESSED
SILICON NITRIDE CONTAINING Y$_2$O$_3$ AND CeO$_2$ ADDITIVES

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Introduction

The high-temperature strength and oxidation resistance of Si$_3$N$_4$ hot-pressed with Y$_2$O$_3$ and rare-earth oxides, particularly, CeO$_2$ have been the subject of numerous studies. It has been generally recognized that the usefulness of these oxides as an effective hot-pressing additive is largely dependent on the formation and stability of several intergranular phases identified as yttrium/cerium silicon oxynitrides which form during hot-pressing. Although the nature and characteristics of the individual oxynitride compounds have been widely reported, there is still much to learn of the composition of the grain-boundary phases that result from particular hot-pressing sequences and their overall effect on the high-temperature properties of Si$_3$N$_4$. Recently, direct microstructural analysis of the grain-boundary phases in Y$_2$O$_3$ fluxed hot-pressed Si$_3$N$_4$ by lattice fringe imaging using transmission electron microscopy has emerged as an useful technique for identifying various amorphous and crystalline phases and for demonstrating their overall distribution in the grain-boundaries of Si$_3$N$_4$. This technique in conjunction with selected area electron diffraction and energy dispersive X-ray microanalysis has been used to determine the composition of intergranular phases in Si$_3$N$_4$ sintered with Y$_2$O$_3$ additive. However, the heterogeneous distribution of the grain boundary phase in these materials makes quantitative analyses of their chemical composition difficult because of the small sampling volume in the TEM methods. Also, the large sampling depth usually makes TEM compositional analysis of fracture surfaces ambiguous because of the thin layer of grain
boundary phase on the fracture surface. Because of the surface sen-
sitivity of Auger electron spectroscopy it has been used by several
workers (14-16) to analyze the fracture surface of MgO doped Si₃N₄ hot-
pressed materials and to estimate the chemical composition of the grain-
boundary glass phase. The work described in this communication uses a
combination of Auger electron spectroscopy and scanning electron micro-
scopy to analyze the grain boundary phases of Y₂O₃ and CeO₂ doped Si₃N₄
hot-pressed materials. It demonstrates that the additives predominantly
concentrate in the grain-boundaries of Si₃N₄ in the form of various
oxynitrides phases.

Materials and Methods

Samples were prepared from high-purity Si₃N₄ powder to which 5 - 20
wt% Y₂O₃ and CeO₂ were added separately. The mixtures were blended for
24 hrs. in polyethylene containers using Si₃N₄ milling media to avoid
contamination. The samples were uniaxially hot-pressed in a graphite
mold coated with BN at 1700°C for 2 hrs. in a purified N₂ atmosphere at
a pressure of 28 MN/m², allowed to cool to room temperature inside the
furnace in flowing N₂ and then machined to remove BN from the surfaces.
A portion from each sample was crushed to fine powder and analyzed by
X-ray powder diffraction (XRD) using CuKα radiation to identify the
crystalline phases present. The grain boundary phases in the hot-
pressed samples were analyzed by Auger electron spectroscopy (AES).
Thin sections cut from the hot-pressed samples were fractured at room
temperature and directly mounted in the AES with the fracture surface
facing upward. For comparison, a portion of the bulk sample was metallographically
polished, cleaned and dried and mounted alongside the fractured specimen. The whole assembly was evacuated to $10^{-9}$ Torr. for 24 hrs. Prior to focusing the primary electron beam to the specimens, the surfaces, particularly, those of the polished specimens were sputtered with an Argon ion beam to remove the contamination due to handling and exposure to the atmosphere. AES was performed with the sputtering switched off but a partial pressure of Argon ($4 \times 10^{-5}$ Torr.) was maintained throughout the analysis. A beam current of 40 µA and primary beam energy of 3 KeV provided the optimum conditions for the resolution of the characteristic Auger peaks for both yttrium and cerium. The fracture surfaces of the hot-pressed samples were also examined by scanning electron microscopy (SEM) and secondary electron images were obtained.

Results

X-ray powder diffraction analysis of the hot-pressed samples of $\text{Si}_3\text{N}_4$ with $\text{Y}_2\text{O}_3$ additive revealed mainly $\beta$-$\text{Si}_3\text{N}_4$ as the major phase with two additional crystalline phases identified as $Y_2\text{Si}_3\text{O}_3\text{N}_4$ and $\text{YSiO}_2\text{N}$. The presence of the latter phase was detected only in those samples which contained a high-proportion of $\text{Y}_2\text{O}_3$. From the available phase equilibria data for the system $\text{Si}_3\text{N}_4$-$\text{SiO}_2$-$\text{Y}_2\text{O}_3$, it is evident that the surface $\text{SiO}_2$ present in the starting $\text{Si}_3\text{N}_4$ has shifted the overall composition within the compatibility triangle $\text{Si}_3\text{N}_4$-$Y_2\text{Si}_3\text{O}_3\text{N}_4$-$\text{YSiO}_2\text{N}$.

Samples hot-pressed with $\text{CeO}_2$ additive, on the other hand, were found to contain $\beta$-$\text{Si}_3\text{N}_4$ as the major phase with two crystalline phases identified as $\text{CeSiO}_2\text{N}$ and the nitrogen containing apatitie-type $\text{Ce}_5(\text{SiO}_4)_3\text{N}$.
As already known\(^{(9,11)}\), CeO\(_2\) decomposes to Ce\(_2\)O\(_3\) in presence of Si\(_3\)N\(_4\) with Ce\(_2\)O\(_3\) and SiO\(_2\) allows the formation of the quaternary oxynitride phases. Thus, it is apparent that the formation of SiO\(_2\) during hot-pressing has resulted in shifting the overall composition within the compatibility triangle Si\(_3\)N\(_4\)–CeSiO\(_2\)–Ce\(_5\)(SiO\(_4\))\(_3\)N in the system Si\(_3\)N\(_4\)–SiO\(_2\)–Ce\(_2\)O\(_3\).

Scanning electron micrographs of the fracture surfaces of hot-pressed Si\(_3\)N\(_4\) doped with Y\(_2\)O\(_3\) and CeO\(_2\) are shown in Fig. 1. Previous workers\(^{(14-16)}\) have reported that the fracture in hot-pressed Si\(_3\)N\(_4\) is predominantly intergranular and hence is expected to show a high proportion of grain-boundary area. In this study, hot-pressed specimens which were fractured at room temperatures and then examined by SEM revealed a combination of inter- and intragranular fracture for both the Y\(_2\)O\(_3\) and CeO\(_2\) doped Si\(_3\)N\(_4\) samples. As shown in Fig. 1(A), a SEM photomicrograph of a Y\(_2\)O\(_3\) doped Si\(_3\)N\(_4\) hot-pressed sample at high magnification exhibits large grains of Si\(_3\)N\(_4\) sintered together. In a few isolated areas, however, elongated pores preferentially located between the Si\(_3\)N\(_4\) grains show evidence of the presence of an intergranular liquid phase. In contrast, SEM examination of CeO\(_2\)-doped Si\(_3\)N\(_4\) hot-pressed samples, Fig. 1(B), revealed numerous areas in which the reaction between Si\(_3\)N\(_4\) and the additive oxide can be demonstrated. Fig. 1(B) shows the morphology of the reaction zone which primarily consists of large Si\(_3\)N\(_4\) grains surrounded by a porous and fine-grained layer containing the oxynitride phases formed as reaction products.
Fig. 1. Scanning electron micrographs of fracture surfaces of hot-pressed Si$_3$N$_4$ containing (A) 10 wt % Y$_2$O$_3$ and (b) 10 wt % CeO$_2$. 
Auger electron spectra obtained for both polished and fracture surfaces showed characteristic peaks for silicon, nitrogen and oxygen. The surfaces were found to be free from any detectable impurities or inclusions except for carbon which was present in significant quantities in all samples. The characteristic oxygen peak observed in the polished samples which persisted even after prolonged sputtering can be attributed to the presence of an oxide phase, presumably surface SiO$_2$ associated with the starting Si$_3$N$_4$ material. No peaks for either yttrium or cerium could be detected in the polished samples which confirms the earlier findings$^{(3)}$ that there is no solubility of these oxides in Si$_3$N$_4$.

In contrast, the characteristic spectra obtained from fracture surfaces showed the presence of yttrium and cerium in the respective samples. Typical Auger spectra of the fracture surfaces of hot-pressed Si$_3$N$_4$ doped with Y$_2$O$_3$ and CeO$_2$ are shown in Fig. 2 (A) and (B) respectively. The peak to peak amplitudes for oxygen (503 eV) and nitrogen (379 eV) varied significantly between the polished and the fracture surfaces. In general, a high oxygen content observed in the fracture surfaces is consistent with the existence of an oxygen enriched phase in the grain-boundaries. Furthermore, the presence of yttrium and cerium in the fracture surfaces and an overall increase in the O/N ratio imply that the additive oxides are predominantly concentrated in the intergranular phases. Thus the AES analysis used in conjunction with XRD and SEM techniques appear to be very effective in the identification and characterization of the grain-boundary phases in hot-pressed Si$_3$N$_4$ with
Fig. 2. Auger spectra of fracture surfaces of Si₃N₄ hot-pressed with (A) 10 wt % Y₂O₃ and (B) 10 wt % CeO₂.
Y$_2$O$_3$ and CeO$_2$ additives. Although no attempt was made in this study to perform quantitative analysis of the grain-boundary phases, an effort is now in progress to locate the area of interest in the sample and to carry out point by point Auger electron analysis using scanning AES to obtain a two dimensional mapping of the concentration of selected surface elements.

Acknowledgements

The authors wish to thank Susan Hofmiester for the Auger electron analysis and Professor Paul H. Holloway for helpful discussions. Partial financial support of NASA Grant #NS6 3254 is also gratefully acknowledged.
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Figure Captions

Figure 1. Scanning electron micrographs of fracture surfaces of hot-pressed Si$_3$N$_4$ containing (A) 10 wt % Y$_2$O$_3$ and (B) 10 wt % CeO$_2$.

Figure 2. Auger spectra of fracture surfaces of Si$_3$N$_4$ hot-pressed with (A) 10 wt % Y$_2$O$_3$ and (B) 10 wt % CeO$_2$. 
COMPOSITIONAL EFFECTS ON $\text{Si}_3\text{N}_4$ FRACTURE SURFACES

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November 23, 1982
Introduction

Studies of dense Si$_3$N$_4$ show that the extent of an intergranular amorphous silicate phase is a function of the concentration and type of densification aids such as MgO$_2$\(^{(1,2)}\), Y$_2$O$_3$\(^{(3,4)}\), CeO$_2$\(^{(5)}\) and Al$_2$O$_3$\(^{(5,6)}\). It has also been shown that the high temperature mechanical strength is also a function of the same compositional variables\(^{(4,5)}\) leading to the conclusion that the intergranular amorphous phase controls the thermal limits of performance. This conclusion is reinforced with evidence that crystallization of the grain boundary phase significantly improves high temperature mechanical strength\(^{(7)}\).

Although there have been efforts to analyze the intergranular phase on Si$_3$N$_4$ fracture surfaces\(^{(8,9)}\) the effects of densification aids and thermal history on extent of crystallinity of the fracture surface is still to be established. Smith and Quackenbush\(^{(5)}\), for example, showed that sintered Si$_3$N$_4$ containing Y$_2$O$_3$ and Al$_2$O$_3$ additions exhibited a 1400°C fracture surface containing many pullouts indicative of easy slippage between grains. Samples without Al$_2$O$_3$ showed much more transgranular fracture at both room temperature and 1400°C. However, identification of the intergranular phase responsible for these differences is extremely difficult since the thickness of the phase must be very small (100-200 Å) and may or may not be homogeneously distributed\(^{(2,6,10)}\).

The objectives of this investigation are twofold, (1) determine whether several surface analysis techniques (x-ray, IRRS, AES, etc.) can detect similar differences between fracture and non-fracture surfaces of dense Si$_3$N$_4$ and (2) establish whether variations in densification
additives or a crystallization heat treatment can alter the fracture surface analysis.

In understanding surfaces and interfaces it is important to recognize that analytical techniques sample to different depths within a surface. Secondary ion mass spectroscopy, ion scattering spectroscopy, and Auger electron spectroscopy (AES) obtain data from the near surface of a sample at a depth of 5-50 Å. The middle surface of the sample can be analyzed by using infrared reflection spectroscopy (RRS) to a depth of 0.5 μm. Standard scanning electron microscopy with energy dispersive x-ray analysis and electron microprobe techniques, penetrate to depths as large as 1.5 μm and such measurements are termed to be from the far surface(7).

Hence the techniques chosen were IRRS, AES and x-ray so that information could be obtained from the near to the far surface.

Materials

Three series of Si₃N₄ samples containing various concentrations of densification aids were compared in this study. Bars containing 8% (by weight) Y₂O₃ and very low (<1%) Al₂O₃ (1" x 1/4" x 1/8") were hot pressed at NASA Lewis Research Center. A GTE SN402 amorphous Si₃N₄ powder was used with Y₂O₃ blended by dry milling. Samples were first cold pressed into compacts followed by hot pressing at 1760°C for 3 hrs. and 5000 psi. The other two series of samples were from commercial lots of sintered Si₃N₄: 13-15% Y₂O₃, 6% Al₂O₃ (GTE SN 2502) and 20% Y₂O₃, 6% Al₂O₃ (GTE SN 3502). All samples were ground and polished to a final surface finish of 1 μm diamond paste.
Selected samples of each composition were also heat treated at 1000°C for 10 hrs. in vacuum, after densification, in order to alter the crystallization of the grain boundary phase.

Results

X-ray analysis

Standard powder diffraction analysis was performed on the samples before and after the 1000°C/10 hr. heat treatment. Figure 1 shows considerable difference in the phases present for the three materials. The sample with only 8% Y₂O₃ and little Al₂O₃ (Fig. 1A) shows -Si₃N₄ similar to the results of Smith and Quackenbush for Si₃N₄+6 Y₂O₃ with no Al₂O₃(5), and as predicted from the compatibility diagram of Wills et. al(12).

Increasing the Y₂O₃ content to 13-15% and addition of 6% Al₂O₃ resulted in only β-Si₃N₄ appearing (Fig. 1B), again similar to the results of Smith and Quackenbush(5) when Al₂O₃ was added to their Si₃N₄ +6% Y₂O₃ samples. The absence of minor phases in the diffraction pattern indicates that the intergranular phase developed during densification is retained as an amorphous phase. However, when sufficient Y₂O₃ is present (20%) the Al₂O₃ addition does not suppress crystallization of the intergranular phase and a significant quantity of the 10.9.1 phase appears (Fig. 1C).
Fig. 1A. X-ray diffraction pattern of Si₃N₄ + 8% Y₂O₃.

Fig. 1B. X-ray diffraction pattern of Si₃N₄ + 13-15% Y₂O₃ + Al₂O₃.

Fig. 1C. X-ray diffraction pattern of Si₃N₄ + 20% Y₂O₃ + 6% Al₂O₃.
A 10 hours heat treatment in vacuum at 1000°C shows a small effect on the extent of the minor phases for the 8% \( \text{Y}_2\text{O}_3 \) sample (Fig. 1A). There is an increase in both the \( \text{Y}_2\text{Si}_2\text{O}_7 \) and 10.9.1 phases. No differences were detected for the phases in the 15% and 20% \( \text{Y}_2\text{O}_3 \) samples after heat treatment (Figs. 1B, 1C).

**Infrared Reflection Spectroscopy (IRRS) Analysis**

Samples from the three compositional series before and after heat treatment were fractured in three point bending. The purpose of the three point bending fracture was to obtain a clean fracture surface that could be analyzed. The fracture surfaces were compared with polished surfaces using infrared reflection spectroscopy (IRRS). The technique was developed for the analysis of vitreous silicates \(^{(13)}\) and recently applied to studies of the oxidation and processing of \( \text{Si}_3\text{N}_4 \) \(^{(14)}\). In this analytical method a double beam IR spectrometer is used in either of two modes. The specimen for analysis can be placed on one aperture and a spectrum obtained in comparison with a highly polished reflective metal mirror. The spectrum that results is termed a single beam spectrum because the purpose of the mirror is simply to eliminate the errors associated with air scattering and instrumental variations. However, for very precise analytical work, or the elimination of certain complex features from spectra, it is possible to compare the surface of the unknown placed on the specimen aperture with a reference standard. The spectrum that results is called a difference spectra. The advantage of a difference spectrum is that it is possible to compare samples with and without various environmental changes in the surface of the sample \(^{(14)}\).
As the incident IR beam penetrates only 0.2-0.5 μm of the surface of the sample, the measurements are sensitive to surface films present on the sample. Consequently, if the Si3N4 fracture surfaces have an increased concentration of amorphous silicate phase over that of the bulk, the difference should appear in the comparison of spectra of a fracture surface versus a polished surface.

A single beam IRRS spectrum of vitreous SiO2 is compared to that of a polished surface of dense Si3N4 +8% Y2O3 in Figure 2. Two major peaks due to Si-O-Si bonds appear in the vitreous SiO2 spectrum. The (S) peak at 1110 cm⁻¹ is due to the molecular stretching vibration of the Si-O-Si bonds and the (R) peak at 470 cm⁻¹ is due to molecular rocking vibrations of the same bonds (8). Because of disorder in the structure of vitreous SiO2, there is considerable broadening of both of these reflection peaks.

The IRRS spectrum of Si3N4 shows two Si-N-Si molecular stretching vibrations (SN₁ and SN₂) located at 1020 cm⁻¹ and 900 cm⁻¹ respectively. Three Si-N-Si molecular rocking vibrations (SNR₁, SNR₂, SNR₃) are also present at 570 cm⁻¹, 430 cm⁻¹, and 370 cm⁻¹. The relationship between these vibrational modes with Si3N4 structures (15,16,17) are still being established.

Single beam IRRS spectra of Si3N4 fracture surfaces (Fig. 2) show a loss of most of the reflected intensity which is due to the roughened surface of the fracture. Nearly all of the intensity of the SN₁ peak
Fig. 2. Infra Red Spectra of $\text{Si}_3\text{N}_4 + 8\% \text{Y}_2\text{O}_3$
is gone whereas some reflection intensity at the location of the SN₂ peak is still present. This suggests that there may be some variation in the composition of the fracture surface compared with the non-fractured material. Only a small difference between samples with and without the 1000°C/10 hr. heat treatment is detected.

In order to emphasize the spectral differences between the polished and the fracture surfaces, a second set of IRRS spectra was obtained using the difference spectroscopy method(10). In this method only the molecular vibrations within the surface layer of the fractured vs non-fractured samples that are different are plotted as a function of the wavelength of the incident radiation.

Fig. 3A shows the IRRS difference spectra for the 8% Y₂O₃ sample with and without the 1000°C/10 hr. heat treatment. Before heat treatment some differences exist in the molecular vibrations in the 1400-1150 cm⁻¹ region for the fracture vs. non-fractured surfaces. This is the spectral region where Si-O-Si vibrations are intense (see Fig. 2) and indicates a larger concentration of Si-O bonds within the fracture surface than in the polished surface. Heat treatment eliminates these differences which indicates that there is less preferential fracture within the intergranular phase after 10 hrs. at 1000°C.

Increasing the concentration of oxide densification aids increases the magnitude of the difference spectra in the 1100 cm⁻¹ to 1300 cm⁻¹ range (Figs. 3B and 3C). This indicates that more Si-O bonds are present in the fracture surface of the samples containing the higher concentrations of Y₂O₃. This is as expected if the densification aids are concentrating
in the grain boundaries and fracture occurs preferentially through the g.b. phase. Heat treatment at 1000°C decreases the spectral difference between fracture and polished surfaces for both the 13-15% Y_2O_3 and the 20% Y_2O_3 samples. However, the phase on the fracture surface of the 13-15% Y_2O_3 samples is more resistant to heat treatment and therefore is indicative of a vitreous phase more resistant to devitrification.

Auger Electron Spectroscopy

Auger electron spectroscopy (AES) was used to determine the composition differences of the fractured versus polished surfaces of the Si_3N_4 samples. The experimental techniques employed in AES are well established\(^\text{18}\) and will not be discussed here. The Auger analysis was made with a single path cylindrical mirror analyzer (CMA5-110, Perkin-Elmer, Physical Electronics Div., Eden Prairie, MN) incorporating a scanning electron gun. The primary electron beam was incidental 35-45° to the fractured surface of the specimen. The primary energy of 5 K eV with the beam current density of 2x10^{-1} A/cm^2 was used throughout the experiments. The derivative of the energy distribution of the secondary electrons was plotted using standard techniques. The modulation signals were kept constant at 3 eV peak to peak up to 550 eV in the Auger spectrum and at 15 eV after 550 eV.

Figure 4 shows typical spectra after Ar sputter cleaning of the surfaces from the Si_3N_4-8% Y_2O_3 series. The oxygen content of the fracture surface (Fig. 4B) is considerably enhanced over that of the polished surface (Fig. 4A). There is almost a factor of 2x higher O/N ratio for the fracture surface. As the percentage of densification aid
Fig. 4A. Auger spectra of Si$_3$N$_4$ + 8% Y$_2$O$_3$ - polished surface.

Fig. 4B. Auger spectra of Si$_3$N$_4$ + 8% Y$_2$O$_3$ - fractured surface.
is increased (Table I) the O/N ratio increased in the polished surface of the Si$_3$N$_4$ samples but even more so in the fracture surface and grain boundary phase.

Vacuum heat treatment of the samples at 1000°C produced quite variable results in the AES analysis (Table 1). The O/N ratio of the 8% Y$_2$O$_3$ sample increased with heat treatment and the fracture and polished surfaces became more alike. This change is consistent with the spectral differences in the IRRS analysis. Heat treating the 20% Y$_2$O$_3$ samples produced similar changes, i.e. the O/N ratio increases with heating and there is less difference between polished and fracture surfaces. However, the 15% Y$_2$O$_3$ material showed a decrease in O/N ratio upon heating and a larger difference between fracture and polish surfaces, for unknown reasons.

The above results confirmed the presence of an oxygen enriched phase at the fracture surfaces of the Si$_3$N$_4$ samples similar to the findings of Powell and Drew(8) and Kosowsky(9). These results also show that the addition of larger quantities of Y$_2$O$_3$ enhanced the concentration of oxygen within the fracture surface. Heat treatment at 1600°C changed the oxygen content of both fracture and non-fracture surfaces. However these findings yield little additional understanding regarding the crystalline versus glassy nature of the fracture surface.

Therefore, another experiment was performed using the rate of Si dissociation from the fracture surface as an index of the amorphous versus crystalline nature of the fracture phase(s). Thomas(19) showed that under conditions of electron bombardment Si dissociated from the
<table>
<thead>
<tr>
<th>Y$_2$O$_3$ Concentration</th>
<th>(O/N) Polished Surface</th>
<th>O/N Fractured Surface</th>
</tr>
</thead>
<tbody>
<tr>
<td>8% Y$_2$O$_3$</td>
<td></td>
<td></td>
</tr>
<tr>
<td>No Heat Treatment</td>
<td>0.6</td>
<td>1.1</td>
</tr>
<tr>
<td>Heat Treated</td>
<td>1.7</td>
<td>2.0</td>
</tr>
<tr>
<td>1000°C/10 hrs</td>
<td></td>
<td></td>
</tr>
<tr>
<td>15% Y$_2$O$_3$</td>
<td></td>
<td></td>
</tr>
<tr>
<td>No Heat Treatment</td>
<td>1.6</td>
<td>2.2</td>
</tr>
<tr>
<td>Heat Treated</td>
<td>0.8</td>
<td>3.0</td>
</tr>
<tr>
<td>1000°C/10 hrs</td>
<td></td>
<td></td>
</tr>
<tr>
<td>20% Y$_2$O$_3$</td>
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<tr>
<td>No Heat Treatment</td>
<td>1.2</td>
<td>1.7</td>
</tr>
<tr>
<td>Heat Treated</td>
<td>3.8</td>
<td>3.1</td>
</tr>
<tr>
<td>1000°C/10 hrs</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
surface of quartz much more rapidly than from the surface of vitreous silica. Therefore, it was hypothesized that if an oxygen enriched Si$_3$N$_4$ fracture surface is amorphous, the rate of Si dissociation should be more rapid than from a devitrified fracture surface. Thus, a difference in the rate of dissociation may be associated with amorphous phase formation at the grain boundaries.

Samples of 8\% Y$_2$O$_3$-Si$_3$N$_4$, before and after heat treatment, were fractured in-situ at 10$^{-9}$ Torr in an Auger electron spectrometer. The decay of the 82 eV Si peak (characteristic of chemically bonded Si) and increase in the 92 eV Si peak (characteristic with elemental silicon) were monitored as a function of incident electron beam current (Fig. 5). Saturation was observed at approx. 200 c/cm$^2$. No effect of the 1000$^\circ$ C/10 hr. heat treatment was observed.

A similar Si dissociation experiment was performed on the fracture surface of the Si$_3$N$_4$ sample containing 13-15\% Y$_2$O$_3$+Al$_2$O$_3$. Saturation occurred at approx. 600 C/cm$^2$. Figure 6 compares results from the two compositions. The Si dissociation from the 8\% Y$_2$O$_3$ fracture surface (10$^{-9}$ Torr) was much more rapid (3x) than dissociation from the 15\% Y$_2$O$_3$ fracture surface. No effect of the 1000$^\circ$C/10 hr. heat treatment was observed in either sample.

Therefore, these results support the conclusion that the fracture surface of the 8\% Y$_2$O$_3$-Si$_3$N$_4$ material. However, they do not provide additional evidence regarding crystallization of the grain boundaries with the 1000$^\circ$C/10 hr. heat treatment.
FIG. 5

RATE OF DISSOCIATION
OF Si

Si$_3$N$_4$ + 8% Y$_2$O$_3$

82 ev Si W/O H.T.  ---
W/ H.T.  ---

92 ev Si W/H.T. ----
92 ev Si W/O H.T. ----

SATURATION ~ 200 C/cm$^2$

PEAK TO PEAK HEIGHT

5 min.  10  15  20  25  30 min.

42 C/cm$^2$  84  126  168  210 C/cm$^2$
SUMMARY OF AES STUDY

\[ \text{Normalized Peak to Peak Height} \]

\[ \text{Si}_3\text{N}_4 + 8\% \text{Y}_2\text{O}_3 \]

\[ \text{Si}_3\text{N}_4 + 13-15\% \text{Y}_2\text{O}_3 + \text{Al}_2\text{O}_3 \]

FIG. 6
Conclusions

Several surface analysis techniques (X-ray, IRRS, AES) applied to the same samples, show that fracture surfaces of Si₃N₄ with Y₂O₃ densification aids possess a higher concentration of oxygen than the bulk. Thus, the oxide densification aids concentrate in the grain boundaries and even low temperature fracture appears to occur preferentially within the oxygen enriched grain boundaries. Increasing concentrations of Y₂O₃ and Al₂O₃ increase the oxygen content of the fracture surface. A range of 13-15% Y₂O₃+6% Al₂O₃ produces an amorphous g.b. phase which is resistant to devitrification. Fracture occurs through the amorphous phase with a 1000°C heat treatment having little effect on the amorphous phase.

Acknowledgment

The authors acknowledge partial financial support of NASA Grant #NSG3254 and helpful discussions with Prof. Paul Holloway.
References


5. J. Thomas Smith & Carr Lane Quackenbush "Phase Effects in Si₃N₄ Containing Y₂O₃ or CeO₂", American Ceramic Society Bulletin 59(5) 529-532 (1980).


ANALYSIS OF GRAIN BOUNDARY PHASE DEVITRIFICATION

OF Y$_2$O$_3$ AND Al$_2$O$_3$ DOPED Si$_3$N$_4$

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Introduction

A number of studies have shown that $Y_2O_3 + Al_2O_3$ additions to $Si_3N_4$ enhance densification and produce an amorphous grain boundary phase.\(^{(1-5)}\)

It has also been shown that thermal treatments can be used to devitrify the grain boundary phase and thereby improve the high temperature mechanical properties.\(^{(4)}\)

Because of the very small scale of intergranular phases and their heterogeneous distribution it is difficult to follow devitrification processes. Previous studies have shown however that surface analysis method such as Auger electron spectroscopy (AES) can be used to analyze compositional differences between fracture surfaces and bulk, polished surfaces.\(^{(6-9)}\)

Differences due to extent of crystallinity are difficult to assess though since AES is primarily a method sensitive to differences in elemental compositions of the surface. It also has been shown that infrared reflection spectroscopy (IRRS) can be used in a difference method to compare the fracture surface vs polished surface of $Si_3N_4$ samples.\(^{(9,10)}\)

However, it is often difficult to interpret details of the difference spectra.

The objective of the present communication is to show that use of a Fourier transform IR spectrometer in a single beam reflection mode can be used for direct comparison of fracture vs non-fractured $Si_3N_4$ surfaces. This is possible because the FTIR method permits a digital summation of nearly 1000 scans of the fracture surface. Consequently even though the fracture surface is rough and scatters the incident IR beam there is sufficient intensity per scan to produce a reliable summation spectrum of the surface. Therefore, changes in the fracture surface resulting from devitrification of the grain boundary phase are observable.
Materials and Methods

The samples used in this study were obtained from NASA Lewis Research Center, Cleveland, Ohio. Commercial grade Si₃N₄, Y₂O₃ and Al₂O₃ were used. The sample contained 15% Y₂O₃ and 2% Al₂O₃ by weight. Mixtures for 100g batches were wet milled in polyethylene bottles for 17-20 hrs using high alumina grinding media and ethanol. The starting compositions were adjusted for Al₂O₃ pick up. After the powders were milled, the slurry was dried and sieved through a 60 mesh sieve to break up agglomerates. The powders were cold pressed at 414 MN/m² and sintered at 1750°C for 2 hrs under 1 atm nitrogen atmosphere. The specimens were then machined into test bars (2.54 by 0.64 by 0.32 inches) and the surfaces were subsequently ground to a surface finish of 10-15 μm.

The samples were heat treated in a vacuum induction heating furnace at either 1000°C for 10 hrs or 1200°C for 10 hrs each. The vacuum was maintained at 10⁻⁵ Torr. The samples were mounted on boron nitride dies during the heat treatment, to prevent contamination. The furnace was allowed to cool for 7 hrs before the samples were retrieved. The samples were analyzed by x-ray diffraction analysis (XRD) and by Fourier transform infrared reflection spectroscopic (FTIR) analysis. (11)

Results

The x-ray diffraction data of the non-heat treated and the heat treated samples are shown in Fig. 1. In the non-heat treated samples only the β-silicon nitride peaks are seen. However an amorphous hump is also observed indicating the presence of a glassy phase. When heat treated at 1000°C for 10 hrs, one additional peak is seen. However, after heat treating at 1200°C for 10 hrs, several additional peaks are observed.
X-RAY DIFFRACTION PATTERNS OF $\text{Si}_3\text{N}_4 + 15\% \text{Y}_2\text{O}_3 + 2\% \text{Al}_2\text{O}_3$

- 1200° C, 10hrs
- 1000° C, 10hrs
- No Heat Treatment
- Fused Si O$_2$

Fig. 1
indicating the presence of crystalline phases. The amorphous hump is also decreased, suggesting that the glassy phase has crystallized and thus given rise to the new peaks.

The samples were analyzed by the FTIR method which involved a computer summation of 960 spectral scans. In addition to analyzing the polished surfaces, the samples were fractured in three point bending and the fracture surfaces were also analyzed by mounting the broken end of the sample over the 3 mm specular reflection aperture. The results are presented in Fig. 2.

The dotted curve in Fig. 2 is the IRRS spectrum of vitreous, fused silica used for a standard of comparison of Si-O stretching bonds at 1120 cm\(^{-1}\) vs Si-N stretching bonds. As discussed previously\(^{(9,10)}\) there are two primary Si-N molecular stretching vibrations in Si\(_3\)N\(_4\), SN\(_1\) at 1050 cm\(^{-1}\) and SN\(_2\) at 900 cm\(^{-1}\). The dashed curves in Fig. 2 show that there is very little change in the SN\(_1\) peak during heat treatment at either 1000°C or 1200°C. However there is a sharpening of the SN\(_2\) peak, especially after the 1200°C treatment. This change is similar to that observed in the devitrification of Li\(_2\)O-2SiO\(_2\) glasses.\(^{(12)}\)

The FTIRRS spectra of the fracture surfaces of the unheat treated sample and the 1000°C/10 hr sample are noisy in spite of averaging 960 scans. In both cases there is a considerable difference in the breadth and location of the SN\(_1\) peak. The larger number of vibrational species in the 1100-1050 cm\(^{-1}\) region is characteristic of the presence of a considerable number of Si-O bonds in the fracture surface. After 10 hrs heat treatment at 1200°C the spectra of the fracture surface and the polished surface are nearly identical. The fracture surface is both smoother and the Si-O bonds present in the other samples no longer dominate the SN\(_1\) peak.
FOURIER TRANSFORM INFRARED SPECTRA OF \( \text{Si}_3\text{N}_4 + 15\% \text{Y}_2\text{O}_3 + 2\% \text{Al}_2\text{O}_3 \)

- --- Polished Surface
- --- Fracture Surface

1200 °C, 10 Hrs.
1000 °C, 10 Hrs.
No Heat Treatment
Fused SiO₂

\( \text{Fig. 2} \)
Discussion

The x-ray diffraction results indicate that crystallization occurred during heat treatment, especially at 1200°C. Even though the x-ray data was obtained from the bulk material, it can be assumed that the crystallization is at the grain boundary phase which is glassy before the heat treatment process.

The FTIRRS analysis shows that the devitrification did occur within the grain boundary phase. Prior to heat treatment the fracture surface shows spectral features characteristic of Si-O as well as Si-N bonds. This is direct evidence of preferential fracture through a phase of different composition than the bulk composition. After heat treatment there is no distinction between the fracture and non-fractured surface.

Conclusion

Use of Fourier transform IR reflection spectroscopy (FTIRRS) and x-ray diffraction shows that 10 hrs at 1200°C is sufficient to densify the amorphous grain boundary phase of Si₃N₄ containing 15% Y₂O₃ + 2% Al₂O₃ densification aids.

Acknowledgments

The authors acknowledge the assistance of Sunil Dutta of NASA Lewis Research Center in obtaining the Si₃N₄ samples and partial financial support of NASA Grant NSG 3254.
References


EFFECTS OF DEVITRIFICATION OF THE
AMORPHOUS PHASE OF $\text{Si}_3\text{N}_4 + \text{Y}_2\text{O}_3 + \text{Al}_2\text{O}_3$ CERAMICS

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Previous studies have shown the effectiveness of $\text{Y}_2\text{O}_3$ as a densification aid for $\text{Si}_3\text{N}_4$ ceramics.\(^{(1-6)}\) It has also been demonstrated that addition of $\text{Al}_2\text{O}_3$ with $\text{Y}_2\text{O}_3$ greatly improves oxidation resistance of these materials.\(^{(3)}\) However, $\text{Al}_2\text{O}_3$ also significantly affects the relative proportion of amorphous phase produced during high temperature densification.\(^{(3)}\) Because the grain boundary amorphous phase is widely implicated in the degradation of mechanical properties of $\text{Si}_3\text{N}_4$ at high temperatures,\(^{(4)}\) it is important to understand the effect of varying the $\text{Al}_2\text{O}_3$ concentration on the extent of glassy phase. Also, since it has been shown that devitrification of the grain boundary phase can improve high temperature mechanical properties,\(^{(4)}\) it is also important to understand the effect of $\text{Al}_2\text{O}_3$ on control of grain boundary devitrification. Both of these topics are investigated in this paper.

Fully densified $\text{Si}_3\text{N}_4$ bodies were prepared by sintering at $1750^\circ\text{C}$ in $\text{N}_2$ with $15\% \text{Y}_2\text{O}_3$ and 2, 4, 6 or $8\% \text{Al}_2\text{O}_3$ as the densification aid. Machined bars were then fractured at room temperature and $1370^\circ\text{C}$ and phase analyses made of polished surfaces and fracture surfaces using X-ray diffraction (XRD) and Fourier transform infrared reflection spectroscopy (FTIRRS). Bars from the same compositions were also heat treated in vacuum at $1000^\circ\text{C}$ or $1200^\circ\text{C}$ for 10 hrs to promote devitrification of the grain boundary (g.b.) glassy phase. Phase analyses of polished and fracture surfaces were also made of these samples. Thus, the primary objective of this study is to relate the changes in room temperature and elevated temperature mechanical strength with the composition and extent of devitrification of the g.b. glassy phase.
Materials and Methods

The compositions were formulated to have a constant amount of 15% Y_2O_3 and varying amounts of Al_2O_3. Table I summarizes the nominal compositions produced.

Commercial grade Si_3N_4, Y_2O_3, and Al_2O_3 powder (see Table II for sources) were used in the fabrication. An impurity analysis of the "as-received" powders is shown in Table II. Si_3N_4 and Al_2O_3 powders were a higher purity with respect to metal contaminants. Y_2O_3 powder contained Al, Si, and Fe as major impurities. The Si_3N_4 powder had an oxygen content of 2.7 wt % and a specific surface area of 11.84 m/g (3-point BET method).

The Si_3N_4 powder was totally amorphous; the powder particles are spherical and often agglomerated (Fig. 1), with individual particle sizes ranging from 0.05 to 1.0 μm. Mixtures for 10 g batches were wet milled in polyethylene bottles for 17-20 h using high alumina grinding media and ethanol. The starting compositions were adjusted to allow for pick up of Al_2O_3 from the mills. After the powders were milled, the slurry was dried on a heated aluminum plate and sieved through a 60 mesh sieve to break up agglomerates. Seventeen grams of mixed powder was cold die pressed into rectangular blocks 7.6 x 2.5 x 0.64 cm followed by cold isostatic pressing at a total pressure of 414 MN/m^2. The compacts were pressureless sintered in a "cold-wall" furnace at 1750°C for 2 h under nitrogen pressure of 1 atm.

Sintered specimens were machined into test bars (2.54 x 0.64 x 0.32 cm), and the surfaces were subsequently ground with a 220-grit wheel to a final surface roughness of 10-15 μ in rms.
<table>
<thead>
<tr>
<th>COMPOSITION</th>
<th>WEIGHT PERCENT</th>
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<tbody>
<tr>
<td>SNYAL-2%</td>
<td>$\text{Si}_3\text{N}_4 - 15\text{ Y}_2\text{O}_3 - 2\text{ Al}_2\text{O}_3$</td>
</tr>
<tr>
<td>SNYAL-4%</td>
<td>$\text{Si}_3\text{N}_4 - 15\text{ Y}_2\text{O}_3 - 4\text{ Al}_2\text{O}_3$</td>
</tr>
<tr>
<td>SNYAL-6%</td>
<td>$\text{Si}_3\text{N}_4 - 15\text{ Y}_2\text{O}_3 - 6\text{ Al}_2\text{O}_3$</td>
</tr>
<tr>
<td>SNYAL-8%</td>
<td>$\text{Si}_3\text{N}_4 - 15\text{ Y}_2\text{O}_3 - 8\text{ Al}_2\text{O}_3$</td>
</tr>
</tbody>
</table>
TABLE II - TRACE IMPURITY ANALYSIS OF
RAW POWDERS (ppm)

<table>
<thead>
<tr>
<th>Element</th>
<th>$\text{Si}_3\text{N}_4$</th>
<th>$\text{Al}_2\text{O}_3^+$</th>
<th>$\text{Y}_2\text{O}_3^{++}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Al</td>
<td>640</td>
<td>Major</td>
<td></td>
</tr>
<tr>
<td>Co</td>
<td>50</td>
<td>--</td>
<td>--</td>
</tr>
<tr>
<td>Cu</td>
<td>--</td>
<td>--</td>
<td>--</td>
</tr>
<tr>
<td>Cr</td>
<td>--</td>
<td>--</td>
<td>90</td>
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<td>Fe</td>
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<td>160</td>
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<tr>
<td>Mg</td>
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<td>110</td>
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<tr>
<td>Mn</td>
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</tr>
<tr>
<td>Mo</td>
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<td>--</td>
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</tr>
<tr>
<td>Ni</td>
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<td>--</td>
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<tr>
<td>Si</td>
<td>Major</td>
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<td>Ti</td>
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<td>--</td>
<td>--</td>
</tr>
<tr>
<td>V</td>
<td>--</td>
<td>--</td>
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</tr>
<tr>
<td>W</td>
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<td>--</td>
<td>--</td>
</tr>
<tr>
<td>Zr</td>
<td>--</td>
<td>--</td>
<td>--</td>
</tr>
</tbody>
</table>

---- = not determined

*GTE, Sylvania, Towanda, PA
+Linde A, Union Carbide Corporation, New York, NY
++United Mineral & Chemical Corporation, New York, NY
Figure 1. Transmission micrograph of commercial silicon nitride powder.
In order to ensure that all samples were homogeneous, infrared reflection analyses were made on 3 or 4 areas on each side of each sample using a 3 mm aperture. Nearly all samples showed no more than ±1% variation in intensity of the SN_1 and SN_2 peaks. Figure 2 illustrates the results of the homogeneity test for 4 samples, SNYAL-2% Al_2O_3, SNYAL-3% Al_2O_3, SNYAL-4% Al_2O_3, and SNYAL-8% Al_2O_3. The data were obtained with a grating IR spectrometer run on medium scan rate. A vitreous silica standard was also run along with each sample as well in order to normalize the data. Previous studies using this method showed very large variations in the microstructure and IRRS spectra for reaction bonded Si_3N_4. The conclusions from the present homogenization study are that the samples prepared by sintering with the Y_2O_3 + Al_2O_3 are uniformly dense regardless of the percentage of Al_2O_3 in the composition.

Two-thirds of the samples from the SNYAL composition series were selected at random for a 10 hour thermal treatment in vacuum induction heating furnace at either 1000°C or 1200°C. The vacuum was maintained at 10^{-5} Torr. During heat treatment the samples were mounted on BN dies to prevent contamination. The furnace was cooled to room temperature during a seven hour period.

Fracture of the samples with and without heat treatment was done on an Instron testing machine in four point bending using SiC loading platens for the 1370°C tests and tool steel for the room temperature fracture. A 3/8" gauge length was used in both cases and a cross heat speed of 0.02"/min. Both low and high temperature fracture was done in air resulting in approximately 5 minutes exposure at 1370°C and 1.5 hours for the high temperature thermal cycle.
ORIGINAl PAGE IS OF POOR QUALITY.
Results

X-ray analysis. Effects of the 1000°C and 1200°C, 10 hour heat treatments on the phases in the SNYAL series were determined by X-ray diffraction. Figures 2-3 and Table III summarizes the findings. The numbers shown in Table III for each 2θ value correspond to the relative intensity of the various X-ray peaks.

Prior to heat treatment only \( \beta \text{Si}_3\text{N}_4 \) was present in the samples containing 2, 4, and 6\% \( \text{Al}_2\text{O}_3 \) along with an extensive glassy XRD hump from 24°-36° 2θ. However even without heat treatment, the 8\% \( \text{Al}_2\text{O}_3 \) material had significant quantities of \( \text{Y}_2\text{Si}_2\text{O}_7 \).

Heating for 10 hours at 1000°C induced formation of a very small amount of \( \text{Y}_2\text{Si}_2\text{O}_7 \) in the 2, 4, and 6\% \( \text{Al}_2\text{O}_3 \) samples. The 8\% \( \text{Al}_2\text{O}_3 \) material developed considerably more \( \text{Y}_2\text{Si}_2\text{O}_7 \) and also some 10-9-1 phase as a result of the 1000°C heat treatment. This large increase in devitrification was probably due to nucleation of the phases already being present in the unheattreated material.

The 1200°C heat treatment results in devitrification of all compositions, however the relative extent of crystallization depends considerably on the \% \( \text{Al}_2\text{O}_3 \) present. Samples with 6\% \( \text{Al}_2\text{O}_3 \) show extensive development of both \( \text{Y}_2\text{Si}_2\text{O}_7 \) and 10-9-1 phases (Fig. 5, Table III). The 2\% \( \text{Al}_2\text{O}_3 \) composition also developed considerable amounts of both \( \text{Y}_2\text{Si}_2\text{O}_7 \) and 10-9-1 phases. The 4\% \( \text{Al}_2\text{O}_3 \) material had more second phases after 1200°C/10 hr than after 1000°C/10 hr, but it was quite resistant to devitrification.

Heating at 1200°C seemed to reduce the extent of second phase crystallization for the 8\% \( \text{Al}_2\text{O}_3 \) material. Figure 6 and Table III show that the relative intensities of both \( \text{Y}_2\text{Si}_2\text{O}_7 \) phases are decreased from
SNYAL - 2% Al₂O₃

ORIGINAl PAGe IS OF POOR QUALITY

1200°C/10hr

1000°C/10hr

No Heat Treatment

Fig. 3

2θ
### Table III. X-ray Diffraction Analysis of Snyal Series

<table>
<thead>
<tr>
<th>°20</th>
<th>d(A)</th>
<th>Snyal-2%</th>
<th></th>
<th>Snyal-4%</th>
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<th>Snyal-6%</th>
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<th>Snyal-8%</th>
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<tbody>
<tr>
<td></td>
<td></td>
<td>No</td>
<td>1000°C</td>
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<td>1000°C</td>
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</tr>
<tr>
<td></td>
<td></td>
<td>H.T. 10 Hr.</td>
<td>10 Hr.</td>
<td>H.T. 10 Hr.</td>
<td>10 Hr.</td>
<td>H.T. 10 Hr.</td>
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<td>-</td>
<td>-</td>
<td>15</td>
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<td>26.5</td>
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<td>Y₂Si₂O₇</td>
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<td>-</td>
<td>8</td>
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<td>Y₂Si₂O₇</td>
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<td>29.9</td>
<td>2.99</td>
<td>Y₂Si₂O₇</td>
<td>-</td>
<td>-</td>
<td>15</td>
<td>-</td>
<td>10</td>
<td>-</td>
</tr>
<tr>
<td>32.8</td>
<td>2.73</td>
<td>10-9-1</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>33.3</td>
<td>2.69</td>
<td>10-9-1</td>
<td>-</td>
<td>-</td>
<td>52</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>33.7</td>
<td>2.65</td>
<td>βSi₃N₄</td>
<td>97</td>
<td>100</td>
<td>98</td>
<td>87</td>
<td>100</td>
<td>100</td>
</tr>
<tr>
<td>36.2</td>
<td>2.48</td>
<td>βSi₃N₄</td>
<td>100</td>
<td>89</td>
<td>100</td>
<td>100</td>
<td>85</td>
<td>98</td>
</tr>
<tr>
<td>37.7</td>
<td>2.38</td>
<td>Y₂Si₂O₇</td>
<td>-</td>
<td>9</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>7</td>
</tr>
</tbody>
</table>
SNYAL - 6% $\text{Al}_2\text{O}_3$

Fig. 5
SNYAL - 8% Al₂O₃

1200°C / 10 hr

1000°C / 10 hr

No Heat Treatment

Fig. 6
those obtained after the 1000°C treatment. Some peaks are even less than observed without heat treatment. This suggests dissolution of the devitrified g.b. phase may be occurring.

Following room temperature fracture, the ends of the fracture samples were oriented incident to the X-ray beam. The objective was to determine whether the fracture surface had a different distribution of crystalline phases than the bulk material. Because of the very small area exposed to the X-rays there was very little diffraction intensity. Table IV summarizes the findings, even though the results must be considered inconclusive. Only \( \beta \) Si\(_3\)N\(_4\) peaks were observed on any of the fracture surfaces. Changing the \( \% \) Al\(_2\)O\(_3\) or increasing crystallization did not show any additional peaks on the fracture surfaces. It is tempting to conclude that the heat treatment decreased the detection of Si\(_3\)N\(_4\) on the fracture surface, see Table IV. However, the intensities were so low that such a conclusion is very tentative.

**FTIRRS analysis.** A Fourier transform IR spectrometer with a specular reflection stage was used to analyze the SNYAL series before and after heat treatment on both polished and fracture surfaces. Because of the roughness of a fracture surface there is considerable scattering of the incident IR beam. However, as discussed in previous papers from this investigation, (#3 and #4) an advantage in the Fourier transform IR reflection spectroscopy (FTIRRS) method is the ability to scan the same area on the sample for numerous times, store the digital spectral data in the computer, and plot a relatively noise-free spectrum of the sample. In order to ascertain an optimal number of scans the fracture surfaces of the SNYAL-2% samples were analyzed for a total of 480, 960, 1440, and 1920 scans. It was found the 960 scans were sufficient to obtain reasonably noise-free fracture surface spectrum.
<table>
<thead>
<tr>
<th>Material</th>
<th>Heat Treatment</th>
<th>27°(βSi₃N₄)</th>
<th>33.7°(βSi₃N₄)</th>
<th>36°(βSi₃N₄)</th>
</tr>
</thead>
<tbody>
<tr>
<td>SNYAL-2%</td>
<td>None</td>
<td>ND</td>
<td>ND</td>
<td>ND</td>
</tr>
<tr>
<td>SNYAL-2%</td>
<td>1000°C/10 Hr.</td>
<td>ND</td>
<td>33.7°(βSi₃N₄)</td>
<td>36°(βSi₃N₄)</td>
</tr>
<tr>
<td>SNYAL-2%</td>
<td>1200°C/10 Hr.</td>
<td>ND</td>
<td>ND</td>
<td>ND</td>
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<tr>
<td>SNYAL-4%</td>
<td>None</td>
<td>ND</td>
<td>ND</td>
<td>36°(βSi₃N₄)</td>
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<tr>
<td>SNYAL-4%</td>
<td>1000°C/10 Hr.</td>
<td>ND</td>
<td>ND</td>
<td>ND</td>
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<tr>
<td>SNYAL-4%</td>
<td>1200°C/10 Hr.</td>
<td>ND</td>
<td>ND</td>
<td>ND</td>
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<tr>
<td>SNYAL-6%</td>
<td>None</td>
<td>ND</td>
<td>ND</td>
<td>36°(βSi₃N₄)</td>
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<tr>
<td>SNYAL-6%</td>
<td>1000°C/10 Hr.</td>
<td>27°(βSi₃N₄)</td>
<td>34°(βSi₃N₄)</td>
<td>36°(βSi₃N₄)</td>
</tr>
<tr>
<td>SNYAL-6%</td>
<td>1200°C/10 Hr.</td>
<td>ND</td>
<td>ND</td>
<td>ND</td>
</tr>
<tr>
<td>SNYAL-8%</td>
<td>None</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td>SNYAL-8%</td>
<td>1000°C/10 Hr.</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td>SNYAL-8%</td>
<td>1200°C/10 Hr.</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
</tr>
</tbody>
</table>

**Note:** ND = No Peaks Detected  
NA = Not Analyzed
This procedure was followed in previous paper #4 and is the basis for
the data presented herein for the other samples in the SNYAL series.

Results from the SNYAL-2% Al₂O₃ materials are shown in Fig. 7 over
the spectral range of 1400-400 cm⁻¹. The fracture surface spectra are
shown only over the range 1300-600 cm⁻¹ for sake of clarity. The FTIRSS
spectrum of a fused, vitreous silica standard is also shown in Fig. 7
for comparison of the molecular vibrational modes on the Si₃N₄ surfaces.
The two primary Si-N molecular stretching vibrations, SN₁ and SN₂, are
located at 1050 cm⁻¹ and 900 cm⁻¹ respectively whereas the Si-O molecular
stretching vibration is at 1110 cm⁻¹.

Figure 7 shows that heat treatment of the SNYAL-2% Al₂O₃ material
makes the fracture surface much more like the polished bulk surface,
especially after 10 hours at 1200°C. The XRD analysis, Table III,
provides the evidence that considerable crystallization has occurred with
both Y₂Si₂O₇ and 10-9-1 phases appearing. The FTIRSS results show that
the Si-O enriched spectral region from 1100-1000 cm⁻¹ present in the
fracture surfaces is eliminated by the crystallization treatment at
1200°C. Thus, the fracture mode which produces an oxide-enriched surface
layer has been replaced by a fracture mode that has equivalence to the
bulk Si₃N₄ structure.

There is much less devitrification of the SNYAL-4% Al₂O₃ material,
Table III, at ether heat treatment. The FTIRSS results, Fig. 8, also
show no shift of the fracture surface spectra of the 4% material in
contrast to that observed in the 2% Al₂O₃ samples with extensive
devitrification. There is little change in the fracture surface spectra
for the 4% Al₂O₃ material; if anything, the sample treat heated at 1200°C
indicates more of a glassy phase on the fracture surface.
FOURIER TRANSFORM INFRARED SPECTRA OF $\text{Si}_3\text{N}_4 + 15\% \text{Y}_2\text{O}_3 + 2\% \text{Al}_2\text{O}_3$

- Polished Surface
- Fracture Surface

1200°C, 10 Hrs.

1000°C, 10 Hrs.

No Heat Treatment

Fused $\text{SiO}_2$

Wavenumbers

% Reflectance

- 1400
- 1200
- 1000
- 800
- 600
- 400

Fig. 7
SNYAL-4% Al₂O₃, 1200°C

Fractured at Room Temperature

Polished

Fig. 8C
Neither the 6% $\text{Al}_2\text{O}_3$ series or the 8% $\text{Al}_2\text{O}_3$ series shows an obvious correlation of the FTIRRS results with the change in crystal phases shown in Table III. There is no appreciable difference between the polished and fracture surfaces in the unheat treated, 6% sample or the 1000°C/10 hr sample (Fig. 9). The extensive devitrification of the 6% $\text{Al}_2\text{O}_3$ sample produces a major change in the $\text{SN}_2$ vibrational mode. It has split into two peaks with one at 975 cm$^{-1}$ and the other at 900 cm$^{-1}$. The structural origin of this spectral splitting is not known at this time. There is much less evidence of the peak splitting on the fracture surface which is more like the unheat treated $\text{Si}_3\text{N}_4$ spectrum.

The 8% $\text{Al}_2\text{O}_3$ series, Fig. 10, also shows little evidence of the fracture surface having a major change in g.b. crystallinity. Splitting of the $\text{SN}_2$ Si-N molecular stretching vibrations is much more extensive for the heat treated material. There is a substantial change of the relative proportion of the split $\text{SN}_2$ peaks for the fracture surface. In contrast, the non-fracture surface of the 8% series changes only a small amount with heat treatment. This finding confirms that the crystallization occurring from the 1000°C and 1200°C heating is primarily located in the g.b. phase and alters the fracture mode for the material.

Fracture at 1370°C in air results in both oxidation of the bulk surface of the $\text{Si}_3\text{N}_4$ samples and the fracture surface. Figure 11 is an example of the FTIRRS spectra obtained from the SNYAL 1370°C fracture series. The sample is SNYAL-4% $\text{Al}_2\text{O}_3$ heat treated for 1200°C for 10 hrs. Comparison with Fig. 8 shows that almost none of the original Si-N vibrational modes are present. Instead a significant peak has developed at 1090 cm$^{-1}$ due to formation of Si-O molecular vibrations. The remainder of the spectrum is complicated and will require additional systematic studies of $\text{Si}_3\text{N}_4$ oxidation to establish the mechanisms of the surface alteration.
SNYAL-6% Al$_2$O$_3$, 1000°C

Polished

Fractured at Room Temperature

Fig. 9B.
Fig. 9c

SNYAL-68 Al₂O₃, 1200°C

Polished at Room Temperature

Fractured at Room Temperature
SNYAL-8% Al₂O₃  
Non-Heat Treated

Polished  
Fractured at Room Temperature

Fig. 10A
SNYAL-8% Al₂O₃, 1200°C

Fractured at Room Temperature

Polished

Fig. 10C
Fig. 12

SNYAL-2, 80%, 1200°C

1108 (0.32)
970 (0.22)
850 (0.26)
SNYAL-1, Non-Heat Treated

Fig. 13
### TABLE V. 4-PT BEND STRENGTH OF SNYAL SERIES

<table>
<thead>
<tr>
<th>Material</th>
<th>No Heat Treatment</th>
<th>Room Temperature Strength (psi)</th>
<th>1370°C Strength</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>No Heat Treatment</td>
<td>1000°C 10 Hrs.</td>
<td>1200°C 10 Hrs.</td>
</tr>
<tr>
<td>SNYAL-2%</td>
<td>No Heat Treatment</td>
<td>70,080 (483 MPa)</td>
<td>79,680 (550 MPa)</td>
</tr>
<tr>
<td>SNYAL-4%</td>
<td>No Heat Treatment</td>
<td>98,880 (682 MPa)</td>
<td>77,760 (536 MPa)</td>
</tr>
<tr>
<td>SNYAL-6%</td>
<td>No Heat Treatment</td>
<td>95,040 (655 MPa)</td>
<td>88,320 (609 MPa)</td>
</tr>
<tr>
<td>SNYAL-8%</td>
<td>No Heat Treatment</td>
<td>69,120 (477 MPa)</td>
<td>21,120* (146 MPa)</td>
</tr>
</tbody>
</table>

*Specimen contained crack prior to loading
TABLE VI. RATIO OF HIGH TEMPERATURE (1370°C) TO ROOM TEMPERATURE BEND STRENGTH

<table>
<thead>
<tr>
<th>Material</th>
<th>No Heat Treatment</th>
<th>1000°C 10 Hrs.</th>
<th>1200°C 10 Hrs.</th>
</tr>
</thead>
<tbody>
<tr>
<td>SNYAL-2%</td>
<td>.04</td>
<td>.03</td>
<td>.07</td>
</tr>
<tr>
<td>SNYAL-4%</td>
<td>.07</td>
<td>.10</td>
<td>.09</td>
</tr>
<tr>
<td>SNYAL-6%</td>
<td>.14</td>
<td>.09</td>
<td>.13</td>
</tr>
<tr>
<td>SNYAL-8%</td>
<td>.23</td>
<td>–</td>
<td>.17</td>
</tr>
</tbody>
</table>

TABLE VII. RATIO OF HIGH TEMPERATURE (1370°C) STRENGTH AFTER HEAT TREATMENT TO UNHEAT TREATED ROOM TEMPERATURE STRENGTH.

<table>
<thead>
<tr>
<th>Material</th>
<th>1000°C 10 Hrs.</th>
<th>1200°C 10 Hrs.</th>
</tr>
</thead>
<tbody>
<tr>
<td>SNYAL-2%</td>
<td>.03</td>
<td>.05</td>
</tr>
<tr>
<td>SNYAL-4%</td>
<td>.08</td>
<td>.08</td>
</tr>
<tr>
<td>SNYAL-6%</td>
<td>.09</td>
<td>.13</td>
</tr>
<tr>
<td>SNYAL-8%</td>
<td>.27</td>
<td>.29</td>
</tr>
</tbody>
</table>
8% Al$_2$O$_3$ material. Heat treatment at 1200°C did not enhance the strength of the other compositions. The most important finding is the improved high temperature strength of the heat treated samples with 8% Al$_2$O$_3$, Table V. There is almost a 7X improvement in high temperature strength for the 1200°C/10 hr heat treated 8% Al$_2$O$_3$ samples over the values for the 2% Al$_2$O$_3$ material. This finding indicates that an optimization of the devitrification schedule of the 8% Al$_2$O$_3$-15% Y$_2$O$_3$ material may yield even more significant advances in the high temperature performance of this material. Other work in this program (paper 416) shows that oxidation resistance is also greatly improved with this material. Consequently, extension of this work should be pursued in order to optimize thermal induced devitrification and the oxidation resistance of 8% Al$_2$O$_3$ material.

References


EFFECT OF $Y_2O_3$ and $Al_2O_3$ ON THE OXIDATION RESISTANCE OF $Si_3N_4$

by

L. L. Hench*, P. N. Vaidyanathan*, and Sunil Dutta**

ABSTRACT

Oxidation of cold pressed and sintered $Si_3N_4$ containing 15 w/o $Y_2O_3$ and 2, 4, 6 and 8% $Al_2O_3$ is observed at temperatures as low as $1000^\circ C$ with infrared reflection spectroscopy. Concentrations of $Al_2O_3 > 4\%$ greatly retard the rate of oxidation and alter the mechanism of surface attack by promoting formation of a glassy layer on the surface containing mixed oxynitride bonds. The glassy layer retards heterogeneous attack and reduces the effect of an oxidation transition temperature between $1000^\circ C$ and $1100^\circ C$ for these materials.

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**NASA Lewis Research Center, Cleveland, Ohio
Introduction

A number of investigations have shown that use of $Y_2O_3$ as a densification aid in $Si_3N_4$ materials can improve high temperature strengths relative to $Si_3N_4$ containing $MgO$. However, instability of certain grain boundary compositions make some $Si_3N_4 + Y_2O_3$ materials susceptible to structural degradation under oxidizing conditions. The accelerated oxygen attack occurs at or below a transition temperature ($T_c$). Above $T_c$, the oxidation kinetics are parabolic with time and the oxide layer formed is dense, coherent and protective. Below $T_c$, oxidation kinetics are linear, the oxide layer has open, connected porosity and is nonprotective.

Additions of $Al_2O_3$ to the $Si_3N_4 + Y_2O_3$ materials suppresses the destructive low temperature oxidation. It is proposed that this is due to formation of a glassy surface phase which spreads at a low temperature to form a dense, pore-free surface layer.

The objective of this study is to examine the effects of $Al_2O_3$ content on the low temperature oxidation of $Si_3N_4 + 15\% Y_2O_3$. This composition is in the $Si_3N_4-Y_2O_3$ compositional field which shows minimal oxidation resistance due to formation of the deleterious $YSiO_2N$ phase. However, addition of $Al_2O_3$ both aids in the formation of a grain boundary liquid and inhibits crystallization of the g.b. phase. Thus, a study of the effects of progressive concentrations of $Al_2O_3$ to $Si_3N_4 + 15\% Y_2O_3$ may yield an understanding of the protective mechanism for low temperature oxidation proposed by Quackenbush and Smith if a glassy surface phase is involved.
The experimental approach used is analysis of the oxidized surface with infrared reflection analysis (IRRS). Previous investigation of oxidized silicon nitride with this method showed that two major modes of attack were present. Si$_3$N$_4$ with MgO additives generally showed a uniform degradation of the IRRS spectrum whereas Si$_3$N$_4$ with ZrO$_2$ showed that a glassy surface layer developed which inhibited oxidation and preserved the strength of the material$^{13,14}$.

**Experimental Procedure**

Four compositions of Si$_3$N$_4$ + Y$_2$O$_3$ and Al$_2$O$_3$ (Table I) were prepared as follows:

Commercial grade Si$_3$N$_4$, Y$_2$O$_3$, and Al$_2$O$_3$ powders were used in the fabrication studies. An impurity analysis and the sources of the "as-received" powders are shown in Table II. Si$_3$N$_4$ and Al$_2$O$_3$ powders were a higher purity with respect to metal contaminants. Y$_2$O$_3$ powder contained Al, Si and Fe as major impurities. The Si$_3$N$_4$ powder had an oxygen content of 2.7 wt% and a specific surface area of 11.84 m$^2$/g (3-point BET method). The Si$_3$N$_4$ powder was totally amorphous; the powder particles were spherical and often agglomerated, with individual particle sizes ranging from 0.05 to 1.0 µm. Mixtures for 100 - g batches were wet milled in polyethylene bottles for 17 - 20 h. using high alumina grinding media and ethanol. The starting compositions were adjusted to allow for pick up of Al$_2$O$_3$ from the mills. After the powders were milled, the slurry was dried on a heated aluminum plate and sieved through a 60 mesh sieve to
### TABLE I

<table>
<thead>
<tr>
<th>MATERIAL COMPOSITIONS (WEIGHT %)</th>
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<tr>
<td>SNA1-1</td>
</tr>
<tr>
<td>SNA1-2</td>
</tr>
<tr>
<td>SNA1-3</td>
</tr>
<tr>
<td>SNA1-4</td>
</tr>
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TABLE II - TRACE IMPURITY ANALYSIS OF
RAW POWDERS (ppm)

<table>
<thead>
<tr>
<th>Element</th>
<th>Si$_3$N$_4$ *</th>
<th>Al$_2$O$_3$ +</th>
<th>Y$_2$O$_3$ ++</th>
</tr>
</thead>
<tbody>
<tr>
<td>Al</td>
<td>--</td>
<td>Major</td>
<td>640</td>
</tr>
<tr>
<td>Co</td>
<td>50</td>
<td>-----</td>
<td>---</td>
</tr>
<tr>
<td>Cu</td>
<td>--</td>
<td>-----</td>
<td>---</td>
</tr>
<tr>
<td>Cr</td>
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<td>-----</td>
<td>90</td>
</tr>
<tr>
<td>Fe</td>
<td>70</td>
<td>70</td>
<td>160</td>
</tr>
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<td>Mg</td>
<td>--</td>
<td>110</td>
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<td>Mn</td>
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</tr>
<tr>
<td>Ni</td>
<td>--</td>
<td>-----</td>
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</tr>
<tr>
<td>Si</td>
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<td>154</td>
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<tr>
<td>Ti</td>
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<td>-----</td>
<td>---</td>
</tr>
<tr>
<td>V</td>
<td>--</td>
<td>-----</td>
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</tr>
<tr>
<td>W</td>
<td>--</td>
<td>-----</td>
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</tr>
<tr>
<td>Zr</td>
<td>--</td>
<td>-----</td>
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</tr>
</tbody>
</table>

---- = not determined

*GTE, Sylvania, Towanda, PA
+Linde A, Union Carbide Corporation, New York, NY
++United Mineral & Chemical Corporation, New York, NY
break up agglomerates. Seventeen grams of mixed powder was cold pressed into rectangular blocks 7.6 by 2.5 by 0.64 cm followed by cold isostatic pressing at a total pressure of 414 MN/m². The compacts were pressureless sintered in a "cold-wall" furnace at 1750°C for 2 h under nitrogen pressure of 1 atm.

Sintered specimens were machined into test bars (2.54 by 0.64 by 0.32 cm), and the surfaces were subsequently ground with a 220-grit wheel to a surface roughness of 10-15 μm in rms.

The samples were oxidized in ambient laboratory air at 1000°C and 1100°C for the following times: 5, 10, 15, 30, 45, 60, 90, 120 minutes.

Before and after oxidation all samples were examined with infrared reflection spectroscopy (IRRS) over the spectral range from 1400 cm⁻¹ to 200 cm⁻¹. The spectra were calibrated to a vitreous silica standard by adjusting a shutter in the IR beam such that the peak for the Si-O-Si molecular stretching vibration at 1120 cm⁻¹ was 80% reflectance.

Results

All four compositions showed nearly identical IRRS spectra, Fig. 1. Multiple scans along the four sides of each sample showed no more than ± 2% variation in IRRS intensity, evidence of excellent homogeneity. Previous investigations of samples with varying degrees of homogeneity showed variations in IRRS intensity of as much as 60% reflection. Thus, the present series of specimens gave highly reproducible oxidation results because of their excellent homogeneity.
Five peaks characterize the IRRS spectra of the silicon nitride samples (Fig. 1). There are two Si–N–Si molecular stretching vibrations, labeled SN₁ and SN₂, and three Si–N–Si molecular rocking vibrations, SNR₁, SNR₂ and SNR₃. The location and relative intensity of these peaks is independent of the concentration of Al₂O₃. Figure 1 also shows the IRRS spectrum of vitreous SiO₂ which consists of two primary peaks. The peak at 1120 cm⁻¹, designated S, is assigned to the Si–O–Si molecular stretching vibration. A single peak (R) at 470 cm⁻¹ is assigned to the Si–O–Si molecular rocking vibration. Thus uniform coherent oxidation of a Si₃N₄ surface should result in a gradual replacement of the five SN and SR peaks with a broadened spectrum containing only S and R peaks.

Figure 2 shows however that oxidation at 1000°C for the sample with 2% Al₂O₃ results in a gradual deterioration of the Si₃N₄ spectrum with no S or R peaks appearing. The progressive decrease in intensity of both molecular stretching and molecular rocking peaks with no shift in peak location can generally be ascribed to increased scattering of the IR beam incident on the surface.

Increasing the Al₂O₃ content to 4 weight % or greater provides a remarkable increase in protection of the surface, Fig. 3. Very little reduction in the relative intensity of the spectrum of samples with 4, 6 and 8% Al₂O₃ is observed after 120 minutes oxidation at 1000°C. In fact the major SN₂ stretching peak is increased, due to a reduction in scattering from surface features that are covered with the glassy oxidation layer.
Fig. 3. IRRS spectra of $\text{Si}_3\text{N}_4 + \text{Y}_2\text{O}_3 + \text{Al}_2\text{O}_3$ samples oxidized at $1000^\circ\text{C}$ for 120 mins.
An important difference is present in the SN₁ peak for the 6% and 8% Al₂O₃ samples 1000°C after oxidation. A second peak located at 1100 cm⁻¹ to 1140 cm⁻¹ has developed, Fig. 3. This change occurs between 90 and 120 minutes at 1000°C. Note however that a new peak at 470 cm⁻¹ has not appeared during the 1000°C oxidation for any of the compositions. Consequently it is unlikely that the new peak developing near 1120 cm⁻¹ is due to formation of Si-O-Si bonds. It is more likely to be due to mixed silicon, oxynitride vibrational species, similar to that observed for low temperature oxidized Si₃N₄ with either MgO or ZrO₂ additives¹⁹. Thus this peak is labeled SNO, in Fig. 3.

The time dependent changes of the intensity of the SN₂ (normalized to pre-oxidation intensity) and the SNO peaks for the four compositions are shown for the first 90 minutes of oxidation in Fig. 4. There is a rapid reduction of the SN₂ peak for the 2, 4 and 6% Al₂O₃ samples followed by a slower rate of attack. The 8% Al₂O₃ sample exhibits only the slower rate of attack of the SN₂ peak. For all four compositions there is no evidence of either the SNO peak, S, or R peaks during the 90 minute time period of 1000°C oxidation. After 120 minutes only the 8% Al₂O₃ sample shows a SNO peak.

Increasing the oxidation temperature to 1100°C greatly accelerates the attack of the composition containing 2% Al₂O₃ (Fig. 5). After just 15 minutes severe degradation of SN₁, SN₂ and all three SNR peaks has occurred. In contrast, the major SN₂ peak of the 4, 6 and 8% Al₂O₃ samples has increased in reflection intensity and the SNR₁,2,3 peaks show very little alteration. The SN₁ peak of the higher Al₂O₃ samples
Fig. 4. Time dependent change of IRRS $SN_2$ peak and SNO peak due to 1000°C oxidation.
Fig. 5. IRRS spectra of $\text{Si}_3\text{N}_4 + \text{Y}_2\text{O}_3 + \text{Al}_2\text{O}_3$ samples oxidized at 1100°C for 15 mins.
has developed into a doublet due to formation of the SNO silicon-oxynitride bonds in the surface. These bonds appear within 5 minutes of 1100°C oxidation of the 8% Al₂O₃ sample (Fig. 6) and are associated with the longer term protection of the surface. After 60 minutes some deterioration of the SN₂ peak has occurred for the 8% Al₂O₃ material (Fig. 7) but the rate of attack is very low (Fig. 6). The intensity of the SNO peak remains approximately constant as the SN₁ peak is completely replaced by the SNO peak as discussed in a subsequent paper on high temperature oxidation of Si₃N₄.

The materials with 4% and 6% Al₂O₃ also show appearance of the SNO peak by 15 minutes oxidation (Fig. 5). The peak remains stable for a short period of time for these compositions (Fig. 6) but is eventually decreased in intensity (Figs. 6, 7) along with the SN₂ peak. This is due to an increase in roughness and scattering of the IR beam by the surface and is an index of overall surface attack for these compositions.

Discussion

These results show that Al₂O₃ significantly increases the oxidation resistance of Si₃N₄ materials containing a high percentage of Y₂O₃. The effectiveness of the Al₂O₃ additions appears to be related to enhanced formation of a surface silicon oxynitride or mixed silicon dioxide-silicon nitride layer, associated with the SNO peak in the IRRS spectra. The fact that an SiO₂ related peak does not develop at 470cm⁻¹ during formation of the SNO peak favors the assignment of the new 1100cm⁻¹ and 1140cm⁻¹ is a mixed silicon oxynitride species.
Fig. 6. Time dependent change of IRRS SN\textsubscript{2} peak and SN\textsubscript{0} peak due to 1100°C oxidation.
Fig. 7. IRRS spectra of $\text{Si}_3\text{N}_4 + \text{Y}_2\text{O}_3 + \text{Al}_2\text{O}_3$ samples oxidized at 1100°C for 60 mins.
A critical concentration of $\text{Al}_2\text{O}_3$ is necessary to form the SNO peak and provide protection against attack of the $\text{Si}_3\text{N}_4$ structure. Without the critical (>4%) amount of $\text{Al}_2\text{O}_3$, surface attack is observed, Fig. 2, as quickly as 5 minutes at 1000°C. The general deterioration of the IRRS spectra is concluded to be due to roughening of the surface indicative of heterogeneous attack of grain boundaries and formation of the porous surface layer described by Quackenbush and Smith\(^1\). When this mode of attack is initiated, it is rapid. Subsequent surface damage is a much slower process (Figs. 4 and 7).

Thus, it is proposed that the region of low temperature linear oxidation kinetics results from the onset of the heterogeneous surface attack. Addition of a critical amount of $\text{Al}_2\text{O}_3$ greatly retards this process even in compositions containing sufficient $\text{Y}_2\text{O}_3$ to be in the $\text{Si}_3\text{Y}_2\text{O}_5\text{N}_4$ field.

These processes and compositional effect appear to be the same at both 1000°C and 1100°C confirming previous the conclusion\(^1\) that development of surface glassy phases greatly reduces the importance of an oxidation transition temperature for these materials.

**Conclusions**

$\text{Al}_2\text{O}_3$ additions of 4% or greater greatly retard oxidation of $\text{Si}_3\text{N}_4$

+ 15% $\text{Y}_2\text{O}_3$ even though this composition is in the field where the destructive $\text{Si}_3\text{Y}_2\text{O}_5\text{N}_4$ phase can form. The oxidation protection is due to formation of a surface layer containing mixed silicon oxynitride bonds which retards a heterogeneous mode of attack of the $\text{Si}_3\text{N}_4$
structure. The mechanisms and compositional effects are the same for both 1000°C and 1100°C indicating that any oxidation transition temperature is masked by the formation of the protective surface layer.

Acknowledgement

The authors gratefully acknowledge partial financial support of NASA Grant No. NSG 3254.
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