EFFECT OF PROCESSING PARAMETERS ON REACTION BONDING OF SILICON NITRIDE

SUBMITTED BY

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Reaction Bonding of Silicon Nitride

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Summary

An investigation into the development of reaction bonded silicon nitride has been conducted in which the relationship between the various processing parameters and the resulting microstructures has been used to design and synthesize reaction bonded materials with improved room temperature mechanical properties.

* This Final Report was written by all present members of the research group together. Alphabetically, they are Otto J. Gregory, Matthew B. Magida and Marc H. Richman.
1. **Introduction**

During recent years, silicon nitride (especially reaction bonded silicon nitride) as well as other ceramics, has emerged as a viable substitute material for gas turbine applications. Due to the characteristics of materials presently used, the overall efficiency of the system is limited. The high temperature strength and resistance to thermal shock, corrosion and creep properties of silicon nitride would allow higher operating temperatures and thus increases in efficiency. Furthermore, the superalloys currently used are expensive and require many materials in strategically scarce supply. Thus research and development of silicon nitride has been quite active.

The investigation into the development of reaction bonded silicon nitride at Brown University has been concerned with the relationship between the processing parameters and the resulting microstructures. These results are then correlated to the bulk mechanical properties of the material. Through an understanding of these relationships, a program can be implemented to produce a superior ceramic material.

Work completed in the early stages of this investigation utilized a starting silicon powder (K series) of relatively high purity with iron being the major (\(\sim1\) wt %) metallic impurity. It was confirmed that the purity and particle size distribution of the starting (green) silicon powder has a marked influence on the nitriding kinetics and resulting material. Since this investigation is concerned with the effects of various processing parameters on mechanical properties, a purer silicon powder (\(\sim0.03\) wt % Fe) was obtained in an attempt to remove the variances caused by impurity additions. Upon nitridation
of this new powder (E series), major differences were observed relative to the well-characterized K-series product.

The most striking difference between the two series was the color of the nitrided sample. The purer E-series powder nitrided to a dull goldish color in contrast to the normal gray color of the K series. The two series further differed in microstructure morphology and bulk mechanical properties. At this point, the investigation concentrated on the cause of these property variations and it was established that Fe as an impurity was directly responsible. With this information, the bulk of the later part of the investigation was concerned with producing, characterizing and testing silicon nitride produced from powder containing from .03 to 1 weight per cent iron. The per cent reacted, density, microstructure, strength and color of the nitrided product have been determined as a function of Fe concentration within the above range. The end results indicate that Fe must be present in the silicon powder in order to produce superior silicon nitride following conventional nitriding procedures.

2. Discussion of Results

The first silicon powder (K-series) used in this investigation was purchased from Kawecki Berylco Ind. Co. (Reading, Pa.) and specified as 99.4% pure (.35% Fe, .30% Al, .03% Ca). The as-received powder purity was determined prior to subsequent processing by neutron activation analysis performed at the University of Rhode Island Nuclear Facility. These results are included in Table I. The impurity levels after powder processing were similarly determined and are presented in Table II.
TABLE I

Neutron activation analysis of K series Si in the as received condition in ppm. (percent where noted).

<table>
<thead>
<tr>
<th>Element</th>
<th>Run 1</th>
<th>Run 2</th>
<th>Run 3</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aë</td>
<td>4.0% *</td>
<td>4.2% *</td>
<td>4.4% *</td>
</tr>
<tr>
<td>Mn</td>
<td>80</td>
<td>84</td>
<td></td>
</tr>
<tr>
<td>V</td>
<td>326</td>
<td>336</td>
<td>343</td>
</tr>
<tr>
<td>Ti</td>
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<td>292</td>
<td>418</td>
</tr>
<tr>
<td>Na</td>
<td>15</td>
<td>12</td>
<td>11</td>
</tr>
<tr>
<td>Sc</td>
<td>1.2</td>
<td>1.2</td>
<td>1.2</td>
</tr>
<tr>
<td>W</td>
<td>75</td>
<td>70</td>
<td>74</td>
</tr>
<tr>
<td>La</td>
<td>4.4</td>
<td>4.3</td>
<td>4.4</td>
</tr>
<tr>
<td>Fe</td>
<td>4822</td>
<td>5270</td>
<td>5117</td>
</tr>
<tr>
<td>Co</td>
<td>4.7</td>
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<tr>
<td>Cr</td>
<td>65</td>
<td>63</td>
<td>67</td>
</tr>
</tbody>
</table>

* Not Meaningful
TABLE II

Neutron activation analysis of the N, K, and E Series Si Powders as milled in ppm.

<table>
<thead>
<tr>
<th>Element</th>
<th>N</th>
<th>K</th>
<th>E</th>
</tr>
</thead>
<tbody>
<tr>
<td>Al</td>
<td>4414*</td>
<td>161.3*</td>
<td></td>
</tr>
<tr>
<td>Mn</td>
<td>126</td>
<td>2</td>
<td></td>
</tr>
<tr>
<td>V</td>
<td>1</td>
<td>377</td>
<td>5.4</td>
</tr>
<tr>
<td>Ti</td>
<td>439</td>
<td></td>
<td>13.3</td>
</tr>
<tr>
<td>Th</td>
<td>4.9</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fe</td>
<td>5226</td>
<td>11187</td>
<td></td>
</tr>
<tr>
<td>Co</td>
<td>205</td>
<td>306</td>
<td>6.7</td>
</tr>
<tr>
<td>Cr</td>
<td>956</td>
<td>1352</td>
<td>18.3</td>
</tr>
<tr>
<td>La</td>
<td>0.2</td>
<td>4.3</td>
<td></td>
</tr>
<tr>
<td>W</td>
<td>1376</td>
<td>780</td>
<td></td>
</tr>
<tr>
<td>Na</td>
<td>33</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sc</td>
<td>0.07</td>
<td>1.08</td>
<td>0.11</td>
</tr>
<tr>
<td>Sb</td>
<td>0.2</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ta</td>
<td>51</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

* Not Meaningful
The as-received powder contained particles with a mean size of 10-15 microns. It has been determined that smaller particle sizes are more easily nitrided to completion. Thus, processing to reduce the powder particle size was needed. It has also been established that nitrided strengths could be increased by increasing the green compact density. This relationship follows the classical fracture mechanics of brittle ceramic materials where the strength controlling flaws in this case are the pores. Thus, the powder was processed in such a way as to yield a higher green density.

White and Walton [1] have quantitatively shown that higher packing densities can be achieved using a mixture of different size particles. The smaller particles can pack, after thorough mixing, in the interstitial voids between the larger particles, leading to a more dense compact. They calculated that for optimum conditions, \( r_2 = 0.414 r_1 \) (\( r_2 \) and \( r_1 \) being the radii of the two-particle sizes). Using this particle packing theory, 100 grams of the as-received powder was ball milled in a Fritsch ball mill with 413 grams of WC balls and 100 ml. of acetone. After varying the ball milling time and determining the particle size distribution using a Fisher Sub Sieve Sizer, two particle size distributions were found in the ratio specified above. After driving off the acetone, the two powder lots were sized to -325 mesh and mixed together in a ratio of 1:1 by weight for 24 hours in a reaction bonded silicon nitride ball mill. The mill (Figure 1) was 3 in. (7.62 cm) in diameter and used five .75 in. (19 mm) reaction bonded silicon nitride cylinders as the grinding medium. It is interesting to note that particle size reduction using the silicon nitride ball mill resulted in a steeper and narrower particle size
Figure 1

$\text{Si}_3\text{N}_4$-ball mill with

$\text{Si}_3\text{N}_4$ balls.
distribution (as shown in Figure 2) than had been found after milling with the WC balls. In this system, silicon nitride is not considered an impurity. The resulting powder had an average particle size of 2.9 microns with the largest particles less than 20 microns. The particle size distribution is shown in Figure 2. Figure 3 is a photomicrograph of the as-milled K series powder.

The K series powder was isostatically pressed at Norton Co. in 6 in. (15.24 cm) diameter rubber bags approximately 8 in. (20.32 cm) long. Hard rubber plates were placed at the ends to insure a flat surface. These were then pressed at 35,000 psi (241 mm/m²), yielding a green compact density of 1.46 g/cc. These cylinders were then processed and nitrided as described below.

The mechanically strongest material obtained from this powder exhibited a room temperature modulus of rupture of 42.0 Ksi in four point bending. This value compares very favorably with those values reported in the literature [2]. The particular nitriding schedule and material properties of this sample is given in Figure 4 and Table III.

Another powder (N series) nominally -325 mesh as-received was also used in conjunction with the K series powder during the first part of the investigation. This powder was purchased from Electronic Space Products, Inc., as nominally 99.99% Si. After processing, the purity was determined using neutron activation analysis and is given in Table II. This powder was sized to -325 mesh and then ball milled in a Fritsch mill. One hundred grams of powder was milled for 4 hours

* Courtesy of Dr. M. Washburn.
Particle distribution for: the N series as received, N, K, and E series powders as milled and sieved; Si₃N₄ ball milled powder before and after milling.
Figure 3

Scanning electron micrograph of the K series Si powder as milled and sieved.
Figure 4
Time-Temperature Schedule for K-8 Run

TABLE III
Material Characteristics for Strongest RBSN to Date (Run K-8)

<table>
<thead>
<tr>
<th>$\alpha/\beta$</th>
<th>%a</th>
<th>$\alpha/\text{Si}$</th>
<th>MDR(ksi)</th>
<th>$K_{1c} \text{ MN/m}^{3/2}$</th>
<th>% Reacted</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.9</td>
<td>73</td>
<td>82</td>
<td>42.0</td>
<td>2.357</td>
<td>92</td>
</tr>
</tbody>
</table>
in a 4 inch (10 cm) diameter stainless steel jar using 320 grams of 0.25 inch (6 mm) WC balls as the grinding medium with 40 ml of acetone. After milling and upon removal of the acetone, the agglomerated Si powder was returned to the mill for 30 seconds. To further insure removal of powder agglomerates, the powder was again sized to -325 mesh. The resultant Si powder had a particle size distribution shown in Figure 2 with average and maximum particle sizes of 3.2 and 25 microns respectively.

The physical characteristics of the powder are shown in Figure 5. The powder was then isostatically pressed at the Norton Co. in 1 inch (25.4 mm) diameter cylindrical polyethylene bags approximately 10 in (254 mm) long at 40,000 psi (275 MN/m²) without a binder. This yielded a green compact with a density of 1.4 g/cc.

Using these two starting silicon powder compacts, experiments were designed and performed in an effort to establish the effects various processing parameters would have on the resulting reaction bonded silicon nitride. Since increased strength is the main objective of this investigation, it is important to establish the strength-controlling criterion. It is widely agreed, that the strength of brittle materials like silicon nitride is controlled, as described by the Griffith theory, by the stress necessary to propagate the largest surface or internal flaws. Thus, increased strengths could be achieved by controlling the size and shape of these flaws. One of the major failure-initiating flaws has been the porosity found in reaction bonded silicon nitride. It is therefore extremely important to understand the variations in the pore structure and understand its development.

Four types of porosity have been observed and quantified during this investigation. The largest, macropores, are associated with the formation
Figure 5

Scanning electron micrograph of the N series Si as milled and sieved.
of the green compact and are common to all ceramics (except those of theoretical density) formed by compacting powders. This post compaction porosity may vary in volume fraction from 0.27 to 0.44 porosity and in size from 10 to several hundred microns. Techniques for reducing the residual macro-porosity are well developed and include increased compacting pressures, decreased particle size and improved void filling processes (particle packing theory). Some densification aids such as MgO and CaO can be used, but they are detrimental at high temperatures due to their enhancement of creep.

The three other types of porosity have been labelled micro-pores, nano-pores and pico-pores in order of decreasing size (Figure 6). The micro-pores are associated with the melting of Si particles (either due to local impurity segregation or as a result of a large reaction exotherm) and may range in size (depending on original Si particle size) up to one hundred microns. The largest of the micro-pores are often found in materials formed from large starting Si powder particles and low α/β ratios. This is not to say that the β formation mechanisms require Si melting, but that this melting is associated with the formation of micro-pores.

The nano- and pico-pores result when silicon grains are reacted to form α-matte silicon nitride. The medium sized nano-pores are present in the unreacted silicon grain ahead of the Si - Si₃N₄ interface as shown in Figure 7. The finest pico-pores are found in the α-matte after the interface has passed through that portion of the silicon grain. Earlier mechanisms presented in the literature [5] for the growth of α-matte and those developed from this work jointly explain
Figure 6

Schematic of the various pore sizes associated with reaction bonded Si$_3$N$_4$. 
Figure 7

Scanning Electron Micrograph of Pore Structure Associated with $\alpha$-Si$_3$N$_4$ Advancing into a Si Grain
the formation of these pores. Briefly, the nano-pores are the result of the loss of silicon through either vaporization or diffusion (vacancy condensation) of the silicon to the nitridation sites. These pores are subsequently sealed off by the formation of the nitride. The pico-pores form as a result of further nitridation inside the nano-pores by either trapped or diffused nitrogen. The nano-pores are reduced to pico-pores through the 22% volume increase upon nitridation. With a knowledge of how these strength-reducing flaws are formed, it becomes important to understand the effect of nitriding conditions on these mechanisms.

At this point it should be emphasized that extensive material characterization was performed on the reaction bonded silicon nitride products. Optical, scanning and transmission electron microscopy were extensively used in the investigation and the experimental procedures will be discussed later. The observed characteristics from this study will be briefly reported below.

Silicon nitride exists in two crystallographic forms, \( \alpha \) and \( \beta \) which can form in distinct morphologies by several different reaction mechanisms. The morphologies that will now be discussed include \( \alpha \)-needles, \( \beta \)-spikes and \( \alpha \)-matte.

\( \alpha \)-needles have a high aspect ratio (>10) and tend to form at lower temperatures (<1300\(^\circ\)C) primarily in the internal porosity or on the external surface of the sample. Figures 8 and 9 show SEM and TEM images of these needles. In Figure 9 the drop at the end of the needle marked A indicates a vapor-liquid-solid (VLS) growth sequence. It has been shown that these beads contain high Fe concentrations which would facilitate
Figure 8

Scanning electron micrograph of α-needles growing into a pore

Figure 9

Transmission electron micrograph of α-needles, needle marked A has a bead at its terminus, B and C have crystalline inner cores.
the VLS mechanism by lowering the temperature of fusion of the reacting Si. Needles B and C have crystalline inner cores with amorphous outer sheaths (determined by lack of tilt contrast). Figure 10 shows a helical banded structure comprised of impurity bands that segregate out of the liquid as the VLS interface progresses. These needle morphologies observed may contribute to the mechanical integrity of the nitride in much the same way as interlocking plates in cement.

Beta silicon nitride has a hexagonal unit cell with large hexagonal tunnels in the <0001> direction. The formation of β silicon nitride is proposed as being controlled by a solid state diffusion mechanism. These β grains can grow very rapidly into the silicon grains at higher temperatures due to the ease of diffusion of N₂ down the β tunnels. β grains that have grown into the silicon grains can be seen in Figure 11. As the β spikes grow, they converge on one another combining to form a uniform, dense β-Si₃N₄ matrix (Figures 12 & 13). It appears from optical microscopy that the size of the β grains is limited mainly by the Si particle size. Figure 14 shows a finer β-grain cluster as a result of nitriding a fine starting Si powder, under conditions aimed at yielding a high α/β ratio.

The β grains not only grow larger than α silicon nitride but also contain much higher dislocation densities. These dislocations (Figures 15 & 16) are thought to be the result of the solid state diffusion growth mechanisms and compounded by the 22% specific volume increase upon nitridation.

In an additional morphology, α silicon nitride can grow as a very fine grained (<1 micron) α-matte (Figure 17). Contrasting the directional
Figure 10

Transmission electron micrograph of α-needles with helical impurity bands (10°C tilt).
Figure 11

Optical photomicrograph of large $\beta$-grains growing into a grain of unreacted Si (white is Si, gray is $\beta$-$\text{Si}_3\text{N}_4$, and black is porosity).
Optical photomicrograph of $\beta$-grains (gray) forming a dense $\beta$-matrix
(Si is white, and porosity is black).

Figure 12

Scanning electron micrograph of $\beta$-$\text{Si}_3\text{N}_4$
forming a dense $\beta$-matrix.

Figure 13
Figure 14

Scanning electron micrograph of finer β-grain cluster size associated with reduced starting Si particle size.
Figure 15

Transmission electron micrograph of large β-grains containing dislocations.

Figure 16

Transmission electron micrograph of large β-grains containing dislocations.
Figure 17

Scanning electron micrograph of fine grained α-matte $\text{Si}_3\text{N}_4$. 
growth of $\beta$-Si$_3$N$_4$ into the unreacted silicon, α-matte forms an even boundary on the silicon surface. This region then advances into the unreacted grain as shown in Figure 18.

The fine grained structure of the α-matte is clearly seen in Figures 19 and 20. The fact that the grain size is substantially less than the original silicon particles and the low dislocation density observed, supports a mechanism involving a vapor phase reaction. Such mechanisms have been proposed during this investigation and supported elsewhere in the literature [3].

Since all these phases and morphologies are simultaneously formed, knowledge of how to control the final microstructure would be advantageous. It is the goal of this investigation to be able to form a microstructure with the best resistance to crack propagation. In order to achieve this, understanding of how the nitriding parameters effect the microstructure is needed.

The nitriding variables of interest are the following: 1) Si particle size and distribution, 2) Si powder purity, 3) temperature of nitridation, 4) gas flow rate, 5) $O_2$ content of the nitriding atmosphere, 6) utilization of an argon pre-sinter, and 7) the $N_2$ content of the atmosphere. These parameters affect such features as the green and nitried density, the percentage reaction, the phase ratios, the grain size and the resultant pore size and distribution. All of these properties will affect the final mechanical properties of the reaction bonded silicon nitride.

Large Si particle sizes hinder successful compacting (particle packing) thus lowering the green density. Furthermore, complete
Figure 18

Optical photomicrograph of fine grained α-matte structure advancing into an unreacted Si grain (white).
Figure 19

Scanning electron micrograph of K series Si$_3$N$_4$.

Figure 20

Transmission electron micrograph of fine grained α-matte Si$_3$N$_4$. 
nitridation of the silicon is impeded by the larger particles. The vapor phase and solid state diffusion reaction mechanisms involved are accelerated by reduced particle size due to increased vapor pressure and reduced diffusion paths. The effect Si particle size has on the microstructure can be seen in Figures 21 through 24 where the particle sizes used were <75, <45, and average particle sizes of 3.2 and 2.9 microns respectively. These pictures clearly show that reduced Si grain size leads to a more refined and uniform microstructure. The results indicate that reducing the Si particle size promotes the reaction mechanisms, reduces the nitriding time, increases nitried density, refines grain size, reduces the general flaw size and hence promotes strength in reaction bonded silicon nitride.

The main concern in reducing the Si particle size is the maintenance of powder purity. Two main types of impurities were investigated. The first group, which are often added as densification aids, include MgO and CaO. These impurities are to be avoided as they tend to form a glassy grain boundary phase to the detriment of the material's ability to resist creep.

The second category includes impurities often present due to the processing techniques employed. These metallic impurities include Al, Cr, Fe, Mn, Ca, Ni, Co, etc. Iron is the most important impurity due to its high concentration (a few ppm to several percent) and also due to its effect on the kinetics of the nitridation reaction. The role of iron as an impurity was studied in the later part of this investigation and will be discussed later. Briefly, iron tends to increase the
Figure 21

Optical photomicrograph of reaction bonded Si₃N₄ made from >75 micron starting Si powder, white is Si and black is porosity.

Figure 22

Optical photomicrograph of reaction bonded Si₃N₄ made from >45 micron starting Si powder.
Figure 23
Optical photomicrograph of reaction bonded $\text{Si}_3\text{N}_4$ made from starting Si powder with an average size of 3.2 microns.

Figure 24
Optical photomicrograph of reaction bonded $\text{Si}_3\text{N}_4$ made from starting Si powder with an average size of 2.9 microns.
reaction rates as well as the resultant $\alpha/\beta$ ratio (see Table IV). Localized segregation of the iron can however be detrimental due to the formation of low melting point SiFx compounds. However, as will be shown later, Fe in the silicon powder is necessary in order to produce superior reaction bonded silicon nitride.

The ultimate temperature and the heating rates used during nitridation greatly affect the microstructure and resultant properties of the material. Observed changes in the $\alpha/\beta$ ratio as a function of the ultimate nitriding temperature is shown in Table V. The increase in $\beta$ content with temperature can be related to existing reaction mechanisms. The high temperatures associated with $\beta$ formation also promote melting of the Si and the formation of large strength reducing pores. If high temperatures are reached too quickly, large $\beta$ grains in an unreacted (melted) Si matrix result (Figure 25). Since $\alpha$ is not present in this region, it would seem that the $\beta$ formed as the silicon was melting. It is thus observed that high nitridation temperatures promote $\beta$ growth and large pore sizes. Unfortunately, it is often necessary to use such temperatures especially when nitriding compacts with high green densities (>1.6 g/cc). A way to minimize these high temperature effects is to hold the reaction at lower temperatures in an effort to completely react the silicon prior to reaching the high temperature.

The heating rates used during nitridation seem to have an effect similar to that of the ultimate temperature. High heating rates reduce the $\alpha$ content while promoting rapid $\beta$ growth. Large void formation is associated with high heating rates thus opposing the increase in reaction rates. Furthermore, a high heating rate in conjunction with the reaction
TABLE IV

Increase in the $\alpha/\beta$ Ratio Associated with Nitridation in the Presence of Fe. 
Pure Sample was Adjacent to Impure Samples Within Furnace Chamber.

<table>
<thead>
<tr>
<th>$\alpha/\beta$ Ratio of Sample Nitrided Alone</th>
<th>$\alpha/\beta$ Ratio of Sample Nitrided in Presence of Impure Si (Fe)</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.3</td>
<td>3.9</td>
</tr>
<tr>
<td></td>
<td>4.2</td>
</tr>
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<td></td>
<td>4.5</td>
</tr>
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</table>


<table>
<thead>
<tr>
<th>Temperature °C</th>
<th>α/β-Ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td>1300</td>
<td>4.4</td>
</tr>
<tr>
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<td>2.7</td>
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<td>1360</td>
<td>2.6</td>
</tr>
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<td>1410</td>
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</table>

<table>
<thead>
<tr>
<th>Temperature °C</th>
<th>α/β-Ratio</th>
</tr>
</thead>
<tbody>
<tr>
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<td>9</td>
</tr>
<tr>
<td>1300</td>
<td>9</td>
</tr>
<tr>
<td>1300</td>
<td>8.1</td>
</tr>
<tr>
<td>1360</td>
<td>2.9</td>
</tr>
<tr>
<td>1360</td>
<td>3.5</td>
</tr>
<tr>
<td>1400</td>
<td>3.5</td>
</tr>
<tr>
<td>1430</td>
<td>1.5</td>
</tr>
</tbody>
</table>
Figure 25

Optical photomicrograph of a sample heated too quickly to 1420°C; note unreacted Si.
exotherm can lead to severe silicon melting and possible damage to the nitriding furnace. Thus slow heating rates are desired.

The effect of gas flow rate on the nitridation of Si has been well established. Since α formation mechanisms are enhanced by SiO vapor, under flow conditions this vapor is likely removed. This inhibits α formation, reduces the reaction rates, and lowers the resultant weight gain and density of the nitried material. Reduction in the room temperature MOR is observed for material formed under flow conditions. This is explained by the effect gas flow has on the reaction kinetics which is to increase β formation with its accompanying increased flaw size. The more refined microstructure of reaction bonded silicon nitride formed with an effective gas flow rate of 0 is compared to that of a conventional cycle in Figures 26 and 27.

The role of O₂ in the nitriding system is not fully understood at this time. However, it is known to greatly affect the reaction rates and the resultant α/β ratio. Sources of O₂ in the system include: oxygen in the nitriding gas, the furnace liner (Al₂O₃ will liberate some O₂ at elevated temperatures), and the silicon powder in the green compact (in the form of SiO₂ or adsorbed onto the silicon particle surface). The major role that oxygen plays during the nitridation reaction seems to lie in its effect on the stability of the layer of SiO₂ on the silicon surface. If the O₂ concentration is maintained below the equilibrium vapor pressure of SiO₂, then the silicon will volatilize forming SiO vapor which aids in the formation of α. This will increase the α/β ratio while achieving complete nitridation to yield reduced flaw
Figure 26

Optical Photomicrographs of Two Ford Motor Co. Reaction Bonded Si$_3$N$_4$ Samples Nitrided Using a Conventional Cycle.
Figure 27

Optical Photomicrograph of Ford Motor Co. supplied Green Si Compact After Nitridation in the K Series System; Note the Relatively Small Pore Size Distribution in Comparison to Figure 26.
sizes and refined microstructures. It should be noted that this is an upper limit on $O_2$ concentration and that much less $O_2$ in the system reduces the $\alpha/\beta$ ratio. The results from the investigation show the maximum $\alpha/\beta$ ratio for the N series (1.5 wt % $O_2$) was 4, while the K series (2.4 wt % $O_2$) was 9. Although the higher $\alpha/\beta$ ratio reduces the pore size distribution, large amounts of unreacted Si are also found and this is detrimental to the strength of the finer material. Thus, $O_2$ can be both beneficial and detrimental to the formation of quality reaction bonded silicon nitride.

Often, in an effort to impart greater strength to the green silicon compact for machining, a presinter in argon is performed. Table VI summarizes the results of an argon presinter of the N series material and shows a decrease in the $\alpha$ content as a result. Argon sintering will reduce the surface area of the silicon particles which in effect lowers the vapor pressure (larger radii of curvature) thus reducing the formation of $\alpha$. The formation of $\alpha$ can be further reduced if argon acts to effectively "clean" the silicon surface resulting in the loss of $SiO_2$ and leaving virgin silicon, a condition that will encourage $\beta$ formation. The sintering of the silicon particles is shown in Figures 28 and 29 as indicated by the inter-particle neck formations. These are the result of a 48-hour 1200°C argon presinter of the K series Si compact.

The addition of $H_2$ to the nitriding atmosphere has been known to be beneficial to both the room and high temperature properties of reaction bonded silicon nitride. This has been attributed to a finer more uniform grain structure and reduced grain boundary impurity
TABLE VI

The Effect of an Argon Pre-Sintering Treatment
On the $\alpha/\beta$-Ratio for N Series Si Compacts.

<table>
<thead>
<tr>
<th>Pre-Sinter</th>
<th>$N_2$ Flow Rate cc/min</th>
<th>Temperature $^\circ$C</th>
<th>$\alpha/\beta$-Ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td>None</td>
<td>0</td>
<td>1390</td>
<td>2.3</td>
</tr>
<tr>
<td>None</td>
<td>130</td>
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<td>3.5</td>
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<td>1100$^\circ$C</td>
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<td>1</td>
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<tr>
<td>1100$^\circ$C</td>
<td>0</td>
<td>1460</td>
<td>1</td>
</tr>
<tr>
<td>1200$^\circ$C</td>
<td>44</td>
<td>1415</td>
<td>2</td>
</tr>
</tbody>
</table>
Figure 28
Optical Photomicrograph of Polished K Series Si Compact After an Argon Pre-Sinter; Note the Neck Formation

Figure 29
Scanning Electron Micrograph of K Series Si Compact After an Argon Pre-Sinter; Note the Neck Formation and the Effectively Large Grain Size.
segregation. It is believed that the H₂ can readily promote increased SiO vapor concentration (through reaction with impurities and oxides which liberate H₂O and other lower more volatile oxides) accounting for the more refined microstructure (Figures 30 and 31). Table VII shows that under those nitriding conditions favoring β formation, the addition of H₂ lowered the α/β ratio. Furthermore, under those conditions favoring the formation of α, the H₂ addition accelerated the rate of α silicon nitride formation.

The results of the K series experiments reveal that there was no significant variation in α/β or α/Si ratio with or without H₂ in the nitriding atmosphere. It appears that these ratios are largely determined by the Si particle size, temperature and oxygen content of the silicon powder. Indications are that the role of H₂ is the refinement of the microstructure, flaw size and grain size through the increased nucleation of silicon nitride by vapor phase transport.

The results presented below are from experiments designed to correlate the reaction mechanisms with the resultant microstructures. The representative curves in Figure 32 show the possible variations in pore size distribution as a function of α/β ratio at constant density. In Figure 33, it may be seen that C (defined as the pore size where there was an average of 1 pore per 0.018 mm²) is independent of density. This further shows the importance of pore size and distribution on the strength of reaction bonded silicon nitride. In Figure 34, C is plotted against α/β ratio for two silicon particle sizes and constant density. It is observed that as the α/β ratio increases, C decreases and that for constant α/β, the pore size
Figure 30
Scanning Electron Micrograph of K Series Si$_3$N$_4$ Nitrided Without H$_2$; Note the Coarse Structure in Comparison to Figure 31.

Figure 31
Scanning Electron Micrograph of K Series Si$_3$N$_4$ Nitrided With 1% H$_2$ Added to the Atmosphere; Note the Fine Structure in Comparison to Figure 30.
TABLE VII

The Effects of H\textsubscript{2} Additions to the Atmosphere on the α/β-Ratio for the N Series Si\textsubscript{2}N\textsubscript{4} Material.

<table>
<thead>
<tr>
<th>Gas</th>
<th>Flow Rate (cc/min)</th>
<th>Temperature (°C)</th>
<th>α/β-Ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td>N\textsubscript{2}</td>
<td>0</td>
<td>1360</td>
<td>4</td>
</tr>
<tr>
<td>N\textsubscript{2}</td>
<td>0</td>
<td>1400</td>
<td>4</td>
</tr>
<tr>
<td>N\textsubscript{2}</td>
<td>10</td>
<td>1400</td>
<td>10</td>
</tr>
<tr>
<td>N\textsubscript{2} - 1% H\textsubscript{2}</td>
<td>0</td>
<td>1450</td>
<td>3.2</td>
</tr>
<tr>
<td>N\textsubscript{2} - 1% H\textsubscript{2}</td>
<td>2</td>
<td>1450</td>
<td>2.0</td>
</tr>
</tbody>
</table>
Figure 32

Pore Size Distribution Curves vs. 
α/β-Ratio for the N Series Si₃N₄.
Figure 33

Pore Size Parameter C vs. Density for the N Series Si$_3$N$_4$. 
Figure 34
Pore Size Parameter C vs. $\alpha/\beta$-Ratio for the N Series $\text{Si}_3\text{N}_4$ at Constant Density.

Figure 35
Optical Photomicrograph of the Pore Size Distribution of N Series $\text{Si}_3\text{N}_4$ with an $\alpha/\beta$-Ratio of 1.
distribution decreases as the silicon particle size decreases. Typical microstructures and pore distributions for the N series material is shown in Figures 35 and 36 with α/β ratios of 1 and 4 respectively. Similar results were observed for the K series material and these in turn were related to the reaction mechanisms involved.

A thorough understanding of the relationships between the processing parameters, reaction mechanisms and microstructures indicated that microstructural control could be employed here. The results have been the production of an N series material with a density of 2.4 g/cc and a room temperature MOR of 42 ksi (275 MN/m²) in three point bending. With a density of 2.46 g/cc, K series material exhibited an MOR of 50 ksi (344 MN/m²) in three point bending. Following is a more detailed presentation of the effects previously mentioned with respect to mechanical properties.

The N series modulus of rupture was determined using a three point bend rig (Figure 37) while the K series was tested using a four point bend rig designed through the courtesy of the NASA Lewis Research Center (Figure 38). Figure 37 also reveals the specimen geometry and the test parameters used. It is important to note that while any particular MOR test may be used to compare silicon nitride materials, only carefully standardized tests can be used to quantitatively compare results from different investigations.

The K series bend bars were diamond ground to final dimensions of 1.250 x 0.250 x .1250 ± 0.001 in. (31.75 x 6.35 x 3.17 ± 0.0254 mm) with a 0.005 in (0.127 mm) 45° edge chamfer to reduce edge failure. An Instron testing machine with a cross head speed of 0.02 in/min
Figure 36

Optical Photomicrograph of the Pore Size Distribution of N Series Si₃N₄ with an α/β-Ratio of 4.
Figure 37

Schematic of N Series Three Point Bend Fixture.
Figure 38

Schematic of Four Point Bend Fixture.
(0.5 mm/min) was used for both the three and four point testing.

When testing a brittle material such as silicon nitride, the surface finish of the test bar is of major concern. Damage caused by grinding will often be the failure initiating flaw. The bars tested during this investigation were prepared following the grinding specifications shown in Table VIII. Figures 39 and 40 show failure initiating from a surface defect while Figure 41 shows an internal failure initiation site.

As previously noted, the pore size C for the N series decreased with increasing α/β ratio. Thus, it would be expected that the N series MOR would increase with decreasing pore size and increasing α content. Figures 42 and 43 confirm this observation.

The results for the K series material, however, do not yield such a direct relationship. The K series MOR varied from 30 to 42 ksi (in four point bending) but did not show a continuing increase with increasing α/β ratios. Samples with high α/β ratios (8 to 9) had MOR values of 30.3-33.6 ksi (209-231 MN/m²). Those samples with maximum MOR had α/β ratios from 3 to 5 while those with ratios below 3 had reduced MOR values. This data is shown in Figure 44.

One possible explanation of the K series results is the amount and role of the unreacted silicon. The K series material with high α-content are also found to contain larger amounts of unreacted Si. Figure 45 is a photomicrograph of such a material and Figure 46 shows an unreacted Si particle as the failure initiating flaw. Samples of silicon nitride with low α/β ratios (<3) are well reacted but exhibit reduced strength.
TABLE VIII
Grinding Specifications for K Series
Four Point Bend Bars.

<table>
<thead>
<tr>
<th>Rough Grind</th>
<th>Intermediate Grind</th>
<th>Finish Grind</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wheel Spec.</td>
<td>100 grit</td>
<td>Wheel Spec.</td>
</tr>
<tr>
<td>Wheel Speed</td>
<td>5000-6000 sfpm</td>
<td>Wheel Speed</td>
</tr>
<tr>
<td>Downfeed</td>
<td>0.002-0.003 in inches/pass</td>
<td>Downfeed</td>
</tr>
<tr>
<td>Crossfeed</td>
<td>1/8-1/4 in inches/pass</td>
<td>Crossfeed</td>
</tr>
<tr>
<td>Table Speed</td>
<td>300-400 inches/min</td>
<td>Table Speed</td>
</tr>
</tbody>
</table>
Figure 39
Scanning Electron Micrograph of Failure Initiating Flaw at Corner for K Series Sample.

Figure 40
Scanning Electron Micrograph of Failure Initiating Flaw at Corner for K Series Sample.
Figure 41

Scanning Electron Micrograph of Internal Failure Initiating Flaw for K Series Sample.
Figure 42

MOR vs. Pore Size Parameter C-0.5
for the N Series Reaction Bonded Si₃N₄

Figure 43

MOR vs. α/β-Ratio for the N Series Si₃N₄
Figure 44

MOR vs. α/β-Ratio for K Series Si₃N₄ Samples;
Note Maximum MOR Does Not Coincide
With Maximum α/β-Ratio.

BARS ARE ± STANDARD DEVIATION

α/β RATIO
K-SERIES  ρ = 2.46 g/cm³
Figure 45

Optical Photomicrograph of a High $\alpha$-Si$_3$N$_4$
K Series Material; Note the Unreacted Si (white)

Figure 46

Scanning Electron Micrograph of Large Si Grain
as the Failure Initiating Flaw in the K Series.
as a result of large pore size distributions. Thus, for maximum MOR, one must insure both minimum flaw size and complete reaction.

Efforts at determining the elastic modulus from the four point bend load-elongation curves were unsuccessful. This was mainly due to the large compliance of the test fixture with respect to the material being tested. The elastic constant $E$ was successfully measured using elastic wave propagation. $E$ was measured for the K-series material as $24.2 \times 10^6$ psi (166 GN/m$^2$).

When considering the use of ceramics in structural applications, component design and fabrication as well as failure prediction and analysis become quite important. To aid these analyses, the evaluation of the plane strain fracture toughness ($K_{lc}$) is required. $K_{lc}$ is a measure of the material's inherent resistance to the propagation of a crack. Once $K_{lc}$ is determined, the critical flaw size, $2a$, can be calculated in conjunction with the material's strength. Furthermore, if the elastic modulus, $E$, is known, the effective surface energy for failure initiation can be determined.

In order to measure $K_{lc}$, a 0.075 in (1.958 mm) notch was cut in the center of the K series bend specimens using a .006 in (0.15 mm) thick diamond wafering blade (Figure 47). The resulting notch root is shown in Figure 48. As can be seen, the region just ahead of the notch root contains small natural cracks. The $K_{lc}$ tests were performed using the four point fixture with a cross head speed of 0.02 in (0.5 mm) per minute. The test bar had a width of 0.125 in (3.175 mm) and a thickness of 0.250 in (6.35 mm).
Figure 47

Specimen Configuration for $K_{IC}$ Test
Figure 48

Optical Photomicrograph of Damage at Notch Root
It is seen in Figure 49 that the fracture toughness is constant with respect to the α/β ratio of the material tested. The average $K_{lc}$ value is 2107.4 ± 160 lbs/in$^{1.5}$ (2.316 ± 0.175 MN/m$^{1.5}$). The variations found for the MOR results with pore size and α/β ratio are not present for the fracture toughness. The main variation in $K_{lc}$ observed was for those samples which were argon sintered prior to nitridation. These samples had an average $K_{lc}$ value of 2,318 ± 232 lbs/in$^{1.5}$ (2.55 ± 0.3 MN/m$^{1.5}$). The exact role argon has in this increase is presently unknown. It is thought that the argon may yield increases in grain boundary energy and through its grain coarsening effect, increase the frictional energy term.

The fracture surface energy $\gamma_1$ for the materials tested was determined yielding an average value of $9.08 \times 10^{-2} \pm 1.8 \times 10^{-2}$ in-lbs/in$^2$ (15.9 ± 3.25 J/m$^2$). This value is in good agreement with results found elsewhere for other Si$_3$N$_4$ materials. The argon sintered material exhibited an increase in $\gamma_1$ to $1.12 \times 10^{-1} \pm 2.2 \times 10^{-2}$ in-lbs/in$^2$ (19.6 ± 3.9 J/m$^2$) as would be expected from the $K_{lc}$ data.

Using these values, the critical flaw sizes were evaluated and found to range from 106 microns for an MOR of 30.3 ksi (209 MN/m$^2$ with α/β = 9) to 52 microns for an MOR of 42 ksi (290 MN/m$^2$ with α/β = 3.5). The size of these critical flaws are comparable to the largest microstructural features observed.

This data indicates that through control of the various processing parameters and an understanding of their effect on the reaction mechanisms, one can design and produce an improved microstructure that will encourage superior mechanical properties.
Figure 49

Fracture Toughness $K_{lc}$ vs. $\alpha/\beta$ Ratio for K Series $Si_3N_4$
The next stage of this research effort concerns the effect of impurities in the starting silicon powder on the reaction mechanisms (of nitride formation) and the properties of the resultant material. In an effort to remove the variances caused by impurity additions, 99.999% pure silicon powder was purchased from Electronic Space Products, Inc., Los Angeles, California. The as received powder purity was determined by neutron activation analysis and is given in Table IX. The powder (E series) was nominally -325 mesh (<44μ) but further reduction in particle size was required for optimum nitridation. The powder was, therefore, further processed at NASA Lewis Research Center by a method developed there [4]. Using an attritor mill (similar to a ball mill, but one in which the action of the balls and additional force is accomplished by an internal rotating oar or paddle) modified to maintain a nitrogen atmosphere, it was found that the steel balls used became intimately coated with a continuous layer of silicon. This protective coating was also established on the walls of the mill. Thus, a situation developed such that silicon was grinding silicon, maintaining the purity of the powder. Table II gives the purity levels after milling.

During the milling process, air cannot be allowed to enter the system or the silicon will be rapidly oxidized and the protective layer will spall off. This oxidation phenomenon represents a problem of major concern with this method of comminution. After milling, the silicon powder is extremely pyroforic. The oxidation process is

* This work was performed by Drs. T. I. bell and T. Glasgow.
# TABLE IX


<table>
<thead>
<tr>
<th>Element</th>
<th>Before</th>
<th>After</th>
</tr>
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<tbody>
<tr>
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<td>95 *</td>
<td>161.3 *</td>
</tr>
<tr>
<td>Mn</td>
<td>2</td>
<td></td>
</tr>
<tr>
<td>V</td>
<td>2.7</td>
<td></td>
</tr>
<tr>
<td>Ti</td>
<td>13.3</td>
<td></td>
</tr>
<tr>
<td>Th</td>
<td>5.4</td>
<td>4.9</td>
</tr>
<tr>
<td>Fe</td>
<td></td>
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<td>6.7</td>
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<td>Cr</td>
<td></td>
<td>18.3</td>
</tr>
<tr>
<td>Sc</td>
<td>0.39</td>
<td>0.11</td>
</tr>
</tbody>
</table>

* Not meaningful
strongly exothermic and the powder can glow white hot if introduced to air in an uncontrolled manner. Hence, the powder was removed in a sealed nitrogen atmosphere, evacuated and then introduced to air slowly. The temperature and pressure of the sealed container was carefully monitored during this step so as to allow the reaction to go to completion to avoid runaway oxidation. This process resulted in a higher O$_2$ content (in the form of SiO$_2$) of the powder than the 1.6 or 2.2% O$_2$ contained in those other powders used previously in the investigation.

Another problem associated with this process is that the mean particle size is not reduced with more grinding time below the low micron range. This has been attributed to the fact that since the powder is milled dry, agglomerates or weak welds between particles develop and are not broken up as shown in Figure 50. Nevertheless, this process was successful in maintaining the basic purity of the powder. Since purity was of major concern, and the powder exhibited excellent compaction properties, no further refining was performed prior to compaction.

The processed powder had an average particle size of 4 microns and a maximum particle size of 30 microns with a size distribution shown in Figure 2. This powder was then compacted using isostatic pressing at the NASA Lewis Center. Two cylinders 3 1/4 inches in diameter and approximately 8 inches long were pressed at 40,000 psi (275 MN/m$^2$) and 60,000 psi (355 MN/m$^2$). This led to a green density of 1.33 and 1.38 g/cm$^3$ respectively. These cylinders were then cut into wafers using a clean hacksaw blade. These wafers were cut using the same blade and shaped into blocks 1.3 x 1.3 x 0.6 in (3.3 x 3.3 x
Figure 50

Scanning Electron Micrograph of the E Series Si Powder as Milled at NASA Lewis
1.5 mm) by hand grinding on 60 grit SiC papers. These blocks were then nitrided to form reaction bonded silicon nitride.

Upon removal from the furnace, the nitrided E series silicon compact was covered with the normal layer of light blue gray silicon oxide fur. After light polishing removed this coating, the reaction product exhibited a dull gold color. All of the reaction bonded silicon nitride made during this investigation and those reported throughout the literature were gray in color. After x-ray diffraction determined that the compact was indeed silicon nitride, an investigation into what was different about the E series began.

The material characteristics and properties of the E series product will be reported first, and then the results of the investigation into the gold silicon nitride will be presented. On a macroscopic level, there often appeared small circular areas of dark gray color (characteristic of the K series nitride) within the gold material. These gray freckles may be associated with localized impurity segregation. Magnetic susceptibility and emission spectroscopy (performed at Texas Instruments, Inc., Attleboro, Massachusetts) were performed on three darker areas and the results are listed in Table X. Upon microscopic examination, no change in microstructure across the freckle-matrix interface could be discerned.

The microstructure of the E series reaction bonded silicon nitride exhibited three main morphologies. Most striking was the observation of large interconnected pore channels. This type of pore structure is shown in Figure 51 and was not apparent in the K-series material. In another morphology, large pores are present but are not as interconnected
TABLE X

Analyses of Impurities in Gray Freckles
Associated with E Series Silicon Nitride.

**Magnetic Susceptibility**

<p>| | | |</p>
<table>
<thead>
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<th></th>
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<tbody>
<tr>
<td>Gold Si₃N₄ (E series)</td>
<td>3.5 x 10⁻⁴ % Fe*</td>
<td></td>
</tr>
<tr>
<td>Gray in Gold Si₃N₄ (E series)</td>
<td>0.02% Fe*</td>
<td></td>
</tr>
<tr>
<td>Gray Si₃N₄ (K series)</td>
<td>0.24% Fe*</td>
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</table>

* It was assumed that the mass of ferromagnetic impurity as reflected by susceptibility was due to Fe.

**Emission Spectroscopy**

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<th>Minor</th>
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<td>Fe, Al, Ni, Ti, Co</td>
<td>Mg, Cr</td>
</tr>
<tr>
<td>Gray in Gold Si₃N₄ (E series)</td>
<td>Si</td>
<td>Mg</td>
<td>Ni, Al, Fe, Ti, Co</td>
</tr>
<tr>
<td>Gray Si₃N₄ (K series)</td>
<td>Si</td>
<td>Mg</td>
<td>Ni, Al, Fe, Ti, Co</td>
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</tbody>
</table>

NA

<p>| | | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
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</thead>
<tbody>
<tr>
<td>Gold Si₃N₄ (E series)</td>
<td>Low</td>
<td></td>
</tr>
<tr>
<td>Gray in Gold Si₃N₄ (E series)</td>
<td>High</td>
<td></td>
</tr>
</tbody>
</table>
Figure 51

Interconnected Pore Channel Structure in
E Series $\text{Si}_3\text{N}_4$ Compacted at 40 Ksi.
as the channels mentioned above. This structure (Figure 52) and the pore channels could have been caused by poor compaction. However, other evidence including the green density achieved indicates that these morphologies are due to reaction mechanisms peculiar to the purer E series compacts. Figure 53 shows the pore structure associated with the E-series compact that was pressed at 60 ksi (355 MN/m²) was similar to that of Figure 52. The third morphological feature observed was more comparable to the K-series material. This more refined and smaller pore structure is shown in Figure 54 along with an unreacted silicon grain into which the nitride was advancing.

The nitridation reaction was generally not as complete for the E-series material as it was for the K series. Following the nitriding schedule that yielded the best K-series material (Figure 4) resulted in an E series compact that was 86% reacted. The mechanisms mentioned earlier could explain this as being caused by the increased O₂ content of the E series powder. The oxide layer on the silicon particles could remain stable longer due to the increased amount of O₂ and hence inhibit the extent of the reaction. It was observed that just prior to the 1200°C nitriding hold, a large amount of fur formed in the hot zone of the furnace surrounding the sample (as observed through an optical pyrometer). This phenomenon coincided with a surge in the nitrogen demanded by the reaction. It is proposed that these occurrences are related to the higher O₂ content and the absence of volatilizing (and cleaning of SiO₂) aids such as Fe. It is interesting to note that one sample was nitrided at high temperature to 78% completion and then renitrided to 89% reacted following a more conventional nitriding schedule.
Figure 52

Large Pores Associated With E Series Nitrided From 40 Ksi Compact

Figure 53

Large Pores Associated With E Series Nitrided From 60 Ksi Compact
Figure 54

Photomicrograph of E Series Si$_3$N$_4$ With
A Morphology Similar to K Series
The ratio of the two phases (α/β) present in reaction bonded silicon nitride ranged from 1.96 to 5.53 for the E series material. As was the case for the K series, the role H₂ had was to promote formation of the phase which was favored by the nitriding schedule. For example, two nitriding runs made using the 4%H₂ gas resulted in α/β ratios of 3.79 and 1.96: the former following the standard schedule (Figure 4) and the latter modified towards longer times at the higher temperatures.

When the E series compact was subjected to an argon pre-sinter, the α/β ratio increased (>5). This is contrary to earlier results and is not fully understood at this time. Furthermore, the argon pre-sinter seemed to refine the E series microstructure (Figure 55) although Figure 56 shows a microstructure more consistent with the mechanisms proposed earlier.

Since the E series reaction bonded silicon nitride exhibited a complicated and overwhelming pore structure, it was not possible to determine the pore size distributions as was done for the K series. However, as would be expected, the enlarged pore structure had an adverse effect on the room temperature mechanical properties. The average modulus of rupture for the E-series material was 15.85 ksi (109 MN/m²) and the fracture toughness was measured as 1640.6 lbs/in¹.5 (1.803 MN/m¹.5). The strongest E series product exhibited a MOR of 17.6 ksi (121 MN/m²) and corresponded to an argon pre-sinter.

As was the case with the N series, the MOR of the E-series material increased with increasing α content (strongest MOR at α/β = 5.12). The
Figure 55
Nitrided E Series Microstructure as Refined by Argon Presinter (9 hrs. at 1000° C)

Figure 56
Nitrided E Series Microstructure Showing β and Large Pores Associated with Argon Presinter
results may also be comparable to the K series but no tests were conducted on E-series material with larger $a/\beta$ ratios. The density of the nitrided E-series material was fairly constant, despite various nitriding schedules, with the average being $2.14 \text{ g/cm}^3$. The two densest nitrides, 2.166 and 2.165 g/cm$^3$, were from E-series material that was argon presintered and made from the compact pressed at 60 ksi, respectively.

These results indicate that the purer E-series silicon compacts nitrided to form reaction bonded silicon nitride containing a large pore structure with reduced room temperature mechanical properties.

When the furnace was first opened after the initial E series run, a strong smell of rotten eggs was observed. This, with the goldish color, pointed to the possibility that sulfur had been introduced into the system. As a test, the nitriding gas used was bubbled through lead acetate in order to precipitate sulfides. No positive results were obtained. Due to the background nitrogen would produce, mass spectrometry of the nitriding gas used was not performed. Eventually, a second E series compact was nitrided using a new supply of gas and produced gold silicon nitride.

This gas had previously been used to nitride a K series compact resulting in the typical gray color. Thus, the nitriding gas used was no longer considered to be the cause of the color change.

A comparison of x-ray diffraction patterns of the K and E series starting silicon powders was made. The E series pattern was unusual in that the silicon peaks were very broad especially at their base. This typ. of peak broadening could be caused by reduced particle size,
residual stresses and strains or an amorphous component in the powder. Another set of x-ray patterns was obtained after a 16 hour 800°C argon sintering of the K and E series powders. The amorphous nature of the E series powder was no longer present. The sintering process would have relieved residual stresses and could have crystallized any amorphous material. It is possible that the two powder lots contained different oxides of silicon, which could effect the reaction kinetics, but no evidence of this was found.

A piece of each nitrided series was metallographically polished for reflectivity measurements. This test revealed that both reaction bonded silicon nitride materials absorbed the same part of the spectrum. However, it was observed that the two patterns differed in amplitude with the K series nitride being approximately one percent more reflective.

The x-ray anomaly mentioned above indicated a need to measure the crystallization temperature of the two powders. A Vacuum Technology Assoc. RF diode sputtering unit was used and operated at a frequency of 13.56 MHz to sputter an amorphous silicon film. The targets made from the E and K series silicon compacts were 2 x 2 . 0.25 in (5.08 x 5.08 x 0.635 cm) and were cleaned by sputtering onto dummy substrates. Two (per series) sapphire single crystal (Al₂O₃) substrates were attached to the vertical plate anode using Dow Corning 340, a silicone heat sink paste. With a plate current of 110 ma, RF voltage of 800v and pressure maintained at 4.0 x 10⁻² torr (argon flow), a sputtering rate of ~3μm/hr. was achieved. The final film thickness measured optically and by using a Tallysurf surface profilometer was between 10 and 15 μm.
X-ray diffraction analysis confirmed that both the K and E series films were initially amorphous. Two films (one from each series) were then heated in a flowing argon atmosphere to 700°C and held there for three hours. After slow cooling, the samples were analyzed by x-ray diffraction. The films appeared to be deteriorating and spalling off. This could have been due to some crystallization but was not supported by the x-ray data.

Several other attempts were made in a similar fashion but no difference was detected between the two films. Due to time limitations, however, no further work was conducted along this line. The ability to sputter both silicon powders successfully was established and it is recognized that this could be important for future silicon nitride development.

Throughout these experiments investigating the E series reaction bonded silicon nitride, work was being done to establish the actual purity of the silicon powder (Table XI). The purity of the powder was measured through the various processing steps and is shown for both the K and E series starting silicon powder.

As already noted, both magnetic susceptibility and emission spectroscopy analyses were made on the E and K series silicon nitride. These tests were in close agreement with the results obtained from the neutron activation analyses. Two other tests* to determine purity were attempted without success. By using an Auger analysis* in conjunction with an ion mill, it would have been possible to obtain an elemental profile of the nitrided sample. Unfortunately, the reaction bonded silicon nitride was not conductive enough. The attempt to use plasma emission spectroscopy*

* Performed at Texas Instruments, Inc., Attleboro, Mass. - Dr. S. S. White
Table XI

Neutron Activation Analysis of the K and E Series
Through the Processing and Nitriding Steps

<table>
<thead>
<tr>
<th>Element</th>
<th>K series green powder</th>
<th>K series argon sintered only</th>
<th>K series Nitrided only</th>
<th>K series argon sintered &amp; nitrided</th>
<th>E series as received</th>
<th>E series after ball milling</th>
<th>E series argon sintered</th>
<th>E series nitrided only</th>
</tr>
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<tr>
<td>Fe</td>
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<td>10,037</td>
<td>7,775</td>
<td>7,083</td>
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<td>603</td>
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<tr>
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<td>-</td>
<td>-</td>
<td>83</td>
<td>53</td>
<td>80</td>
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<td>Mn</td>
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<td>-</td>
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<td>9</td>
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<td>Mg</td>
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<td>2,820</td>
<td>1,000</td>
<td>1,592</td>
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<td>50</td>
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<td>V</td>
<td>547</td>
<td>561</td>
<td>388</td>
<td>387</td>
<td>-</td>
<td>101</td>
<td>101</td>
<td>67</td>
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<td>Ti</td>
<td>422</td>
<td>430</td>
<td>340</td>
<td>342</td>
<td>-</td>
<td>-</td>
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<td>1,149</td>
<td>1,036</td>
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<td>23</td>
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<tr>
<td>W</td>
<td>1,055</td>
<td>1,044</td>
<td>953</td>
<td>880</td>
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<td>-</td>
<td>280</td>
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<tr>
<td>Co</td>
<td>300</td>
<td>293</td>
<td>260</td>
<td>242</td>
<td>-</td>
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<td>6</td>
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<tr>
<td>Ni</td>
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<td>1,053</td>
<td>706</td>
<td>650</td>
<td>-</td>
<td>101</td>
<td>44</td>
<td>-</td>
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<tr>
<td>Th*</td>
<td></td>
<td></td>
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<td></td>
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<tr>
<td>Sc*</td>
<td>34</td>
<td>31</td>
<td>23</td>
<td>21</td>
<td>-</td>
<td>1</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>

*Concentration in PPM
failed as the silicon nitride could not be made to go into solution.

Based on the results that were obtained, it became apparent that Fe as an impurity was most important. The results from the magnetic susceptibility test (Table X) strongly substantiates this fact. The gray in the gold E series nitride exhibited a susceptibility, attributed to Fe, between the values attained for the K and E series nitrides. Thus, a series of experiments were designed to try to verify the role of iron as an impurity.

The E-60-Fe experiment was designed to try to introduce electrolytic grade iron into the E series silicon compact pressed at 60,000 psi. Three silicon samples were prepared according to Table XII. These samples were dried in an oven at 90°C prior to nitriding using the 1% H₂ gas according to the following schedule: 1000°C, 2 hrs.; 1100°C, 24 hrs.; 1190°C, 18 hrs.; 1360°C, 3 hrs.

Macroscopically, samples A and B appeared light gray around the edge and the bulk was brownish gold. Sample C was basically light gray with many small dark dots and some isolated golden areas. Samples A, B and C did not react to completion and were 59%, 46% and 38% reacted respectively. Their respective α/β ratios were 12.5, 7.7 and 13.1.

Microscopic observation of the microstructures revealed a morphology of disjointed material (Figure 57) that was later seen to be connected by an underlying matrix. Figures 58 and 59 represent the microstructures of samples B and C respectively. The open, interconnected pore channels associated with the E series silicon nitride were not as prevalent in these samples.
Table XII

E-60-Fe Sample Preparation

Sample A. 1 small silicon wafer soaked in 125 ml HCl + 375 ml distilled water + 5 g. electrolytic grade iron for 64 hours

Sample B. 1 large silicon wafer soaked in above solution plus methanol (1:1) to break surface tension for 64 hours

Sample C. E-60 forming powder soaked in solution for 40 hours. Repressed using die at 6,000 psi.
Figure 57
Photomicrograph of E-60-Fe Sample C Showing Disjointed Material Appearance

Figure 58
Photomicrograph of E-60-Fe Sample B Showing More Refined Microstructure as Compared to E Series Material. Note Unreacted Silicon (white)
Figure 59

Photomicrograph of E-60-Fe Sample C

Note Large Pores But Absence of Pore Channels
The results from the E-60-Fe experiment offer positive proof that Fe as an impurity in the starting silicon powder is responsible for the observed color changes in reaction bonded silicon nitride. The next group of experiments was designed to repeat the E-60-Fe program in a more controlled manner. The sample preparation for this GTG series is listed in Table XIII. These samples were nitrided using the 1% H₂ gas following a schedule of: 1100°C, 21 hrs.; 1200°C, 10 hrs.; 1300°C, 10 hrs.; and 1390°C, 48 hrs.

GTG-1 reacted to the typical gold color associated with the E series silicon nitride. On the top edge of the sample, a layer of gray silicon nitride had formed. It is believed that since all of the GTG samples were nitrided together, the vapors associated with each sample may have mixed. Thus, it becomes apparent that some Fe transported to the sample in the vapor state causing the shallow layer of gray silicon nitride. The microstructure of this sample was characteristic of the E series but the pore structure was more refined (Figure 60). This could also be a result of increased vapor activity. The sample was 82.4% reacted with an α/β ratio of 7.2.

The macroscopic appearance of GTG-2 was typical of the K series silicon nitride. The microstructure, however, exhibited the disjointed morphology shown in Figure 61. Observations made using the SEM revealed that a typical matrix of silicon nitride lay below and in between the isolated islands of material (Figure 62). This sample was 89.4% reacted with an α/β ratio of 3.7.

Sample GTC-3 was light gray in color with a slightly pitted surface. The color of this silicon nitride indicates again that Fe as an impurity
### TABLE XIII

**Sample Preparation for GTG Experiment**

<table>
<thead>
<tr>
<th>Sample</th>
<th>Purpose</th>
<th>Processing</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>E series control brick</td>
<td>10 g. brick</td>
</tr>
<tr>
<td>2</td>
<td>K series control brick</td>
<td>11 g. brick</td>
</tr>
<tr>
<td>3</td>
<td>To introduce Fe impurity to E series powder</td>
<td>10 g. of E powder for 23 hours in solution* plus 3.00 grams of Fe** powder, repressed at 6000 psi</td>
</tr>
<tr>
<td>4</td>
<td>To remove metallic impurities from K series powder</td>
<td>10 g. K series powder soaked in 1:1 HF:H$_2$O (distilled) for 22 hrs. Powder repressed at 6000 psi</td>
</tr>
<tr>
<td>5</td>
<td>To chemically etch metallic impurities</td>
<td>10 g. K series powder soaked for 41.5 hrs. in solution*</td>
</tr>
<tr>
<td>6</td>
<td>To introduce Fe into E series compact</td>
<td>9 g. brick soaked in 100 mL of solution + 50 mL ethyl alcohol + 3 g. Fe** for 20 hrs.</td>
</tr>
<tr>
<td>7</td>
<td>To remove metallic impurities from K series brick</td>
<td>10.5 g. brick soaked in 100 mL of solution* for 20 hrs.</td>
</tr>
</tbody>
</table>

* Solution: 250 mL reagent grade HC$l$ + 750 mL distilled H$_2$O

** Fe: Electrolytic grade iron
Figure 60
Photomicrograph of GTG-1 Microstructure

Figure 61
Photomicrograph of GTG-2 Microstructure
Figure 62
Scanning Electron Photomicrograph of GTG-2
Revealing Underlying Si₃N₄ Matrix

Figure 63
Photomicrograph of GTG-4 Microstructure
was present. The microstructure observed was similar to that of GTG-2 except that the isolated islands of material were larger. GTG-3 was 76.6% reacted with $\alpha/\beta = 4.21$.

GTG-4 exhibited a gold color with a shallow gray area on the top face (similar to GTG-1). A small internal gray area was also present and indicates a local concentration of Fe. The microstructure of the sample is shown in Figure 63 as being porous but very tight. The porosity may be due to the low compacting pressure used. This material was 87.6% reacted and had a high $\alpha/\beta$ ratio of 11.1.

Although GTG-5 was prepared to remove metallic impurities, it was macroscopically similar to K-series silicon nitride. The powder used here was washed in a HC$l$ solution. The results, however, indicate that the HC$l$ was never able to "clean" the silicon particles due to its inability to penetrate the native SiO$_2$ layer on the particle surface. The microstructure of GTG-5 was similar to that of GTG-2 as previously reported. This sample reacted to 79% completion with an $\alpha/\beta$ ratio of 3.0.

The center of GTG-6 was a light gold color that darkens progressively to gray at the edges. This indicates that the Fe transporting solution did not thoroughly penetrate the compact. The microstructure exhibited porosity and isolated material morphologies. The sample was 88.1% reacted and had an $\alpha/\beta$ equal to 2.38.

The GTG-7 sample was gray throughout indicating the lack of HC$l$ penetration mentioned for GTG-5. The bulk of the material was very porous and exhibited the island morphology. This material was 88.1% reacted with a resultant $\alpha/\beta$ ratio of 3.9.
In order to use plasma emission analysis to determine the impurity content of the GTG reaction bonded silicon nitrides, the material first had to be put into solution. Attempts were made to fuse the silicon nitride powders by mixing with sodium carbonate (Na₂CO₃) and heating in a platinum crucible. Solution was not achieved by this or several other methods, and a more indirect method had to be used.

To determine if the GTG sample preparations did indeed accomplish the objectives, plasma emission analysis was conducted on the original solutions used in the sample preparation (see Table XIV) and these results were compared to the plasma emission results on the filtrates. This comparison indicated clearly that the goals of the sample preparations were achieved (except as noted above for GTG-S and 7). Further confirmation of this was attained by microprobe elemental analysis of the GTG nitrided samples using an ARL microprobe unit.

The results of the GTG experiment shows that when Fe is removed from the K series silicon powder (which normally nitrides to a gray color), gold reaction bonded silicon nitride results. Furthermore, the gold silicon nitride produced from the E series silicon powder becomes gray when Fe is added. Thus, Fe as an impurity in the starting silicon powder has a major effect on the production of reaction bonded silicon nitride. These effects were further investigated and will be reported later.

Since various impurities greatly affect the nitridation of silicon, an investigation concerning the nitridation of single crystal silicon was initiated. To better understand the nitridation mechanisms responsible for the densification of silicon compacts during the reaction
### TABLE XIV

**Plasma Emission Analysis**

Values in μg/ml

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<thead>
<tr>
<th>Sample</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Solution (250 ml HCl + 750 ml H₂O) + 3 g. iron</td>
</tr>
<tr>
<td>2</td>
<td>GTG-3 Filtrate</td>
</tr>
<tr>
<td>3</td>
<td>GTG-4 Filtrate</td>
</tr>
<tr>
<td>4</td>
<td>GTG-6 Filtrate</td>
</tr>
<tr>
<td>5</td>
<td>GTG-7 Filtrate</td>
</tr>
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</table>

<table>
<thead>
<tr>
<th>Sample</th>
<th>Cr</th>
<th>Fe</th>
<th>Mg</th>
<th>Ni</th>
<th>A%</th>
<th>Si</th>
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<tbody>
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<td>905</td>
<td>2.4</td>
<td>143</td>
<td>241</td>
<td>30.3</td>
</tr>
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</table>

*Performed at Texas Instruments, Attleboro, Massachusetts: Dr. A. Savolainen.*
bonding process. A series of nitriding experiments were carried out on Czochralski grown single crystals of Si (semiconductor grade). Wafers were metallographically polished and then nitrided using a heating rate of 50°C per hour in each of the following schedules: 1) 1100°C, 20 hrs.; 1360°C, 90 hrs. or 2) 1100°C, 24 hrs.; 1200°C, 40 hrs.; 1330°C, 24 hrs. The nitriding gas used throughout these experiments was N₂ - 1% H₂ which had been dried over P₂O₅ and gettered for O₂ over pure Si metal.

In addition to the usual characterization of the nitride by x-ray diffraction and SEM, Laue (back-reflection) x-ray diffraction patterns of the polished silicon wafer faces were taken to determine the orientation of each face and thus the nitriding direction. The wafers had been cut such that the (111) planes of the silicon crystal were parallel to the faces of the wafer and this established the nitride growth direction as being parallel to the [111] silicon direction.

The silicon nitride formed on single crystal silicon was quite different from the nitride developed from Si powder compacts, especially with respect to phase content and microstructure. Due to the different crystal structures of silicon nitride and silicon, there was no evidence of epitaxial growth of the silicon nitride from the silicon surface [5]. The thickness of the silicon nitride layers varied from about 10 to 16 μm depending on the schedule used, with thicker layers being developed at higher temperatures. The extent of nitridation depends primarily on the solid state diffusion of nitrogen through the silicon nitride already formed as also indicated by the kinetics of nitridation.
X-ray diffraction analysis of the Si$_3$N$_4$ layers revealed a predominantly $\beta$-Si$_3$N$_4$ product. The presence of a large amount of $\beta$-Si$_3$N$_4$ in the reacted layer was verified by examining fracture and nitrided surfaces using SEM. A typical SEM micrograph of a fracture surface of the wafer is shown in Figure 64. The morphology of this Si$_3$N$_4$ layer is characteristic of these products on single crystals, in that they are composed of a dense, continuous glassy like phase ($\beta$-Si$_3$N$_4$) which appears smooth and with very small pores. Associated with these pores at the free surface is a small amount of $\alpha$-Si$_3$N$_4$ (needles), as seen from the SEM micrograph of the free nitride surface shown in Figure 65. X-ray diffraction revealed only small amounts of $\alpha$-Si$_3$N$_4$ and thus small $\alpha/\beta$ ratios. This result is consistent with the $\alpha$-Si$_3$N$_4$ formation mechanisms proposed by Jennings [6] and Danforth [7]. Only when elemental Si is volatilized from the bulk crystal through the vapor phase to beads surrounded by nitrogen gas can the $\alpha$ phase develop. The amount of $\alpha$ formed is limited by the nitride product already formed on the crystal. Silicon must diffuse through this layer, most likely through any porosity or breaks in the film to a free surface where there is a supply of N$_2$ gas, and volatilize at the exterior of the product. Since there is only a single flat grain, there is little opportunity for further surface reaction and thus minimal $\alpha$ formation. The largest of these pores is on the order of 5 $\mu$m. A high magnification SEM micrograph of one of these pores associated with $\alpha$ formation is shown in Figure 66. Evidence of crystal facets below the surface of the pore suggests that it is in a $\beta$-Si$_3$N$_4$ matrix and that it may have developed due to a break in the dense $\beta$-Si$_3$N$_4$ grain structure or other pore mechanisms already discussed.
SEM Micrograph of Fracture
Formed from a Single Plate

α-Needles Formed in Region
Figure 66
SEM Micrograph of a Pore Associated with the 
a-Needles Formed on a Free Si₃N₄ Surface

Figure 67
SEM Micrograph of the Si-Si₃N₄ Interface Showing 
Small Pores Formed from Vacancy Condensation
The SEM micrographs shown in Figures 67 and 68 reveal the morphology in the vicinity of the Si-Si$_3$N$_4$ interface. This structure can account for some of the porosity in the nitride layer and, at the same time, verify the existence of a vacancy-condensation process ahead of the nitride growth front.

Just ahead of the α-matte growth front which is growing in the [111] direction with respect to the Si, there appear several small pores isolated in the bulk silicon crystal. According to Atkinson [3] and Danforth [7], the Si migrates out through this region to form subsequent Si$_3$N$_4$ on the exterior. This migration leaves vacancies in the bulk Si single crystal which condense to form nano-porosity ahead of the interface. The reaction of N$_2$ with Si vapor within the internal porosity causes infilling of the nano-pores to produce pico-pores in a fine grained α Si$_3$N$_4$. These particular pores observed in the single crystal were only associated with the α-matte formation and were small enough to be characterized as pico porosity.

Close examination of the Si-Si$_3$N$_4$ interface in the SEM micrograph shown in Figure 69 reveals the presence of randomly orientated, hexagonal prisms, characteristic of β-Si$_3$N$_4$. In addition, very small nuclei of β-Si$_3$N$_4$ appeared to be forming just ahead of the predominantly β-Si$_3$N$_4$ growth front. The hexagonal crystals at the growth front are not oriented in any given direction and they lose their prismatic character at the silicon surface. Since there is no epitaxial growth of Si$_3$N$_4$ from the silicon, there may be distortions of the crystal facets due to the nucleation and growth process. Away from the interface, these crystals became more regular.
Figure 68
SEM Micrograph of Pores Formed by Vacancy Condensation Just Ahead of the α-Matte Growth Front

Figure 69
SEM Micrograph of Fracture Surface of Si₃N₄ Layer Showing β-Si₃N₄ Growth Front and Small Prismatic Nuclei Forming Just Ahead of it. Note Facets of the Hexagonal Crystals in the Nitride Layer.
The SEM micrographs shown in Figures 67 and 68 reveal the morphology in the vicinity of the Si-Si$_3$N$_4$ interface. This structure can account for some of the porosity in the nitride layer and, at the same time, verify the existence of a vacancy-condensation process ahead of the nitride growth front.

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Figure 68
SEM Micrograph of Pores Formed by Vacancy Condensation Just Ahead of the α-Matte Growth Front

Figure 69
SEM Micrograph of Fracture Surface of Si₃N₄ Layer Showing β-Si₃N₄ Growth Front and Small Prismatic Nuclei Forming Just Ahead of it. Note Facets of the Hexagonal Crystals in the Nitride Layer.
The observation of such prismatic nuclei forming ahead of the $\beta$-$\text{Si}_3\text{N}_4$ growth front has not appeared in the literature to date. The mechanisms for $\beta$ formation already discussed are mainly concerned with the growth of $\beta$ spikes ($\beta$ crystals) once they are formed. It appears from the single crystal nitridation experiments that a nucleation process involving the adsorption of nitrogen gas, the formation of $\beta$ embryo and the stabilization of these $\beta$ embryo of a critical size must be an integral part of the formation of $\beta$-$\text{Si}_3\text{N}_4$. Only those $\beta$ nuclei of critical size will continue to grow by the mechanism already mentioned while those unstable nuclei will disappear. It has not been determined whether or not a favorable orientation of silicon will result in the stabilization of these $\beta$ nuclei but it can be stated conclusively that $\beta$-silicon nitride forms by a nucleation and growth process and needs no Fe impurities in the silicon for its subsequent formation as mentioned in the literature [8].

A final experiment was designed to measure the effect of iron as an impurity on the microstructure and mechanical properties of reaction bonded silicon nitride. As previously noted, the E series silicon powder had an iron concentration of $\approx 0.03$ weight percent iron and the K series powder $\approx 1.0$. These two powders were mixed together so that a range of composition from 0.03 to 1.0 weight percent iron in 0.097 increments was achieved. These eleven samples were nitrided and then characterized.

These EOI powder mixtures were combined in the appropriate ratios so that the final weight equaled 10 grams. The samples were labeled from EOI-1 to EOI-11 corresponding to the end points of 0.03 to 1.00 percent Fe respectively. The two powders were mixed together by hand using a mortar and pestle. Acetone was added to insure thorough mixing.
and the mixing was continued until most of the acetone had evaporated (approximately 10 minutes) and the mixture was quite muddy, to minimize powder segregation. After drying the powder in an oven at 55°C, the mortar and pestle were used to break up the caked surface.

The powder was then pressed, with a little acetone, into 1 in. (2.54 cm) diameter cylinders in a split die under 6,000 psi using a standard laboratory press. Care was taken to insure the release of air from the compact by raising the pressure slowly and holding it for ten minutes at each 1,000 psi increment. This yielded a green density from 1.19 to 1.33. The variation of green density with iron content is shown in Figure 70.

After pressing, the powder compact was placed in vacuum to remove any residual acetone. All of the samples were nitrided using the 1% \( \text{H}_2 \) gas and in accordance with the nitridation schedule shown in Figure 4. (The 1360°C hold was maintained for 24 hours.) As a direct result of the vapor interaction experienced during the GTG experiments, each EOI sample was nitrided separately. Weight gain was used to monitor the extent of the reaction and Figure 71 shows the variation in percent reacted as a function of iron concentration.

After nitridation, the color of the reaction bonded silicon nitride was examined. As anticipated from the earlier experimental results, a complete color spectrum from gold to gray (EOI-1 to EOI-11, respectively) silicon nitride resulted. The iron responsible for the change in color was not found to cause any observable change in the x-ray diffraction pattern.

The presence of Fe in silicon nitride can add more energy levels within the band gap of silicon nitride. In an effort to verify this
hypothesis, a gold and gray piece of silicon nitride was mechanically thinned for transmission measurements in the optical wavelength range. Since the sample needed to be translucent, further thinning was attempted using an ion mill. This experiment is still in progress at this date. Once the samples are thinned, it is believed that the two materials will exhibit different transmission cutoffs representative of their respective band gaps.

X-ray diffraction analysis was also used to determine the phase ratios ($\alpha/\beta$, $\alpha/\mathrm{Si}$) of the resultant reaction bonded silicon nitride. The $\alpha/\beta$ measurements ranged from 4.5 to 8.7. These values are high compared to the averages normally found and may be a direct result of the compacting pressure. No trends could be established relating the $\alpha/\beta$ ratio to the concentration of iron. However, both the $\alpha/\mathrm{Si}$ (7.9-64.6) and $\beta/\mathrm{Si}$ (1.7-15) ratios tended to increase until EOI-8 (0.709 wt % Fe) and then decreased. The reason for this is not understood at this time, but it does indicate that within the range being considered, the amount of unreacted silicon decreases to a minimum.

As previously discussed, Figure 71 shows the variation of the extent of the nitridation reaction as a function of the iron content. It is noted that the percent reacted greatly increases with increasing iron content from powder nitrided with from 0.03 to 0.321 wt. % Fe. The trend continues throughout the EOI series but to a much lesser degree. These observations are not unexpected and are well explained by the kinetics of the nitridation kinetics.
The nitrided density was found to increase with increased iron content. Figure 70 illustrates this and also shows the green density and the ratio of nitrided density to green density as a function of iron concentration. The increase in nitrided density is partially due to the observed increase in green density. However, since the ratio of the densities also increases with iron indicates that the green density is not the sole cause. It is consistent with present theories to note that the increasing density ratio curve closely resembles the percent reacted pattern.

Prior to presenting the mechanical properties of the reaction bonded silicon nitride products, the morphological observations made using the optical and scanning electron microscope will be presented.

All of the microstructures discussed were observed throughout the EOI samples. As indicated by the results cited above and suggested by the formation mechanisms presented, the occurrence of open microstructures (isolated islands of material, pore channels, large pores) decreased with increasing iron content. Figures 72 through 75 show the densification and refinement of the reaction bonded silicon nitride with the addition of iron to the silicon powder.

A series of high magnification optical photomicrographs (Figures 76-78) clearly elucidate the nature of the disjointed microstructure previously discussed. As the series progresses (focusing further into the void), the underlying silicon nitride matrix is revealed. These void areas (more prevalent in the purer materials) may be the result of compaction macropores compounded by the diffusion of silicon to reaction sites and not filled in due to less formation by vapor-condensation mechanisms associated with the lack of iron.
Figure 72
Photomicrograph of E01-2 Microstructure wt. % Fe = 0.127 α/β = 8.7

Figure 73
Photomicrograph of E01-5 Microstructure wt. % Fe = 0.418 α/β = 6.0
Figure 74
Photomicrograph of EOI-8 Microstructure wt. % Fe = 0.709  \( \alpha/\beta = 5.7 \)

Figure 75
Photomicrograph of EOI-11 Microstructure wt. % Fe = 1.00  \( \alpha/\beta = 6.46 \)
Figure 76
Focus on Top Surface of E01-2 (0.127 st. % Fe) Microstructure. Note Comparison with Figures 77 and 78

Figure 77
Same Area as Figure 76 but Focusing Deeper into Microstructure
Figure 78
Same Area as in Figures 76 and 77
Focus is on Bottom of Microstructure

Figure 79
EOI-4 (0.321 wt. % Fe) Microstructure Exhibiting 'Ghost' Grains (Lighter Regions)
Figures 79 and 80 show a microstructural component that is similar to that representing prior austenite grains seen in various steel alloys. These 'ghost' grains (or colonies) appear as the lighter phase in Figure 79. It is possible that these are partially reacted silicon grains, but close observation reveals smaller prismatic white regions within these ghosts that are more typical of unreacted silicon. Furthermore, the ghost in Figure 80 does not resemble unreacted silicon. It is possible that these areas are the result of powder agglomerates in the silicon compacts as well as subtle differences in the impurity content.

A more obvious effect of local impurity gradients is shown in Figure 81. This type of microstructure is present throughout the EOI series reaction bonded silicon nitride. From a microstructural viewpoint, the darker region is representative of a higher iron concentration. This is arrived at for two reasons. First, as has been shown, the darker color is associated with iron impurities. Secondly, the kinetics of the reaction mechanisms suggest that with iron present, a finer more uniform material is produced. This is clearly seen in the photomicrograph. It is interesting to note that the effect iron has is extremely localized as inferred from the sharp boundary observed. The association of this darker area with iron was strongly supported by the microprobe elemental analysis performed during the GTG experiments. As the elemental profiles were determined across the sample surface, readings were taken for neighboring light and dark areas (as seen through the microscope attached to the unit). The darker areas were
Figure 80
EOI-11 (1.00 wt. % Fe) Microstructure
Exhibiting a 'Ghost' Grain (Lower Center)

Figure 81
Photomicrograph of EOI-6 (0.515 wt. % Fe) Revealing
Microstructures Representative of High and Low Iron Levels
determined to contain more than twice the amount of iron contained in the lighter areas.

The scanning electron micrographs shown in Figures 82 and 83 were taken from neighboring areas. An EDAX (Energy Dispersive Analysis of X-Rays) profile of the two regions revealed iron to be present in one (Figure 83) but not the other (Figure 82). Again, the refining effect in the presence of iron can be seen. Furthermore, the high magnification of these pictures emphasizes the very localized effect of iron.

The mechanical properties of the EOI reaction bonded silicon nitride samples investigated include modulus of rupture, hardness and the fracture toughness parameter. These properties were related to the amount of iron present in the silicon compact and the material characteristics observed.

The hardness of the various samples was measured using a Wilson Hardness Tester with a Vickers Diamond Pyramid Indentor ($136^\circ$) and a 20 Kg load. The Vickers Hardness Number, $H$, was calculated by measuring the diagonal lengths of several indentations. These lengths were often difficult to discern due to the inherent porosity of the material. Furthermore, the presence of this porosity leads to the crushing of the material and can alter the hardness measured. This effect was noted when the same material was tested under different loads and did not exhibit a constant hardness number. Nevertheless, the measurements made can be used for qualitative comparisons.

The results obtained for the EOI samples show that the hardness increases with increasing iron content. This increase becomes greater at the higher iron concentrations. This trend is understandable in light of the results and mechanisms previously discussed.
Figure 82
Scanning Electron Micrograph of EOI-2 (0.127 wt. % Fe)
EDAX Analysis Showed No Iron

Figure 83
Region Directly Adjacent to That Shown in Figure 82.
EDAX Analysis Revealed the Presence of Iron
Due to the size limitations of the EOI reaction bonded silicon nitride cylinders, there were not enough bend bars to determine the fracture toughness according to the procedure previously outlined. Thus a method developed by Evans and Charles [9] to measure the effective fracture toughness of ceramic materials using indentation fracture was attempted. Once again, the pore structure of the silicon nitride precluded the measurement of the cracks emanating from the indented diamond corners needed, and the fracture toughness was not successfully measured.

The modulus of rupture of the EOI samples was determined in three point bending. The bend rig used was similar to that shown in Figure 37 but here the span between the lower pins was 0.625 in (15.9 mm). Any bend bars which were observed to contain pressing cracks were not included in the final calculations reported.

Figure 84 is a compilation of the modulus of rupture results for the EOI reaction bonded silicon nitride. The strongest material corresponded to EOI-11 (1.0 wt. % Fe) which had an average MOR of 24.2 Ksi (165 MN/m²). This value is lower than expected for the K series silicon nitride and is a direct result of the compacting technique.

The EOI-5 (0.418 wt. % Fe) run was repeated using powder that was sieved to -325 mesh prior to compacting. The resultant increase in strength due to the processing is indicated in Figure 84. The effect of processing was to remove powder agglomerates to aid compaction and resulted in a 15.6% increase in nitried density over the EOI-5 sample originally processed.
Figure 84

Modulus of Rupture as a Function of Fe

Content for the EOI Reaction Bonded Silicon Nitride Series
As expected, the strength of the reaction bonded material continually increased with increasing iron content. The low strength shown for 0.515 wt. % Fe (EOI-6) can be explained by the presence of an unreacted core in the center of the bars. Although scanning electron microscopy was used to survey the fracture surfaces, it was difficult to identify the failure initiating flaw. This was partially due to the mottled appearance of the surfaces caused by porosity and further compounded by the disruptive nature of the three point bend test (avoided by four point bending).

Figure 85 shows the variation of MOR with respect to the weight percent of the resultant phases. The patterns associated with the weight percents of α and β are not fully understood at this time. The exponentially decaying silicon weight percent curve indicates, as previously established, that the modulus of rupture drastically decreases with increasing amounts of unreacted silicon.

As already established for the N series, the EOI reaction bonded silicon nitride exhibited a linear increase in MOR with increasing α/β ratios at constant density. This is depicted in Figure 86 where the modulus of rupture has been normalized to constant density.

The results of the EOI experiments indicate that Fe as an impurity in the starting silicon powder refines the microstructure of reaction bonded silicon nitride and encourages improved room temperature mechanical properties.
Figure 85

Modulus of Rupture vs. wt. % α; wt. % β;
and wt. % Si for the EOI Reaction Bonded Silicon Nitride
Figure 86
MOR Data (Normalized to Nitrided Density) as a Function of $\alpha/\beta$-Ratio for the EOI Reaction
Bonded $Si_3N_4$ Series
The above investigation has revealed that through thorough control of the various processing parameters, improved understanding of the relationships between these variables and the reaction mechanisms involved in the growth of $\alpha$ and $\beta$-reaction bonded silicon nitride, one can successfully design and fabricate a reaction bonded $\text{Si}_3\text{N}_4$ material with improved microstructural features and increased MOR, $K_{1c}$, $\gamma_i$, and 2a (at room temperature).

It has been determined that the formation of $\beta$ silicon nitride involves a nucleation and growth process and can occur in the absence of iron and without the melting of silicon.

Through a systematic evaluation of the effect of iron as an impurity in the starting silicon powder, it has been determined that improved microstructural and mechanical properties of reaction bonded silicon nitride are achieved with iron additions. The gold color of the reaction bonded silicon nitride was caused by the absence of iron.
References


Personnel

The following personnel participated in this research project:

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Appendix A

Experimental Procedures

The furnace system used in the nitridation process (see Figs. A-C) was capable of allowing various atmospheres, flow rates and heating schedules. The furnace chamber was an open ended recrystallized Al₂O₃ tube, 3.125 in ID x 3.5 in OD x 48 in (7.9 x 8.9 x 122 cm). The ends of the tube were sealed using rubber gaskets and stainless steel end caps modified for gas and viewing ports. Alumina silicate (Al₂O₃SiO₄) heat shields were placed into the tube in order to achieve a large, constant hot zone by reducing radiant heat losses. The heat shields also protected the Al₂O₃ tube by leveling the thermal gradient along the length of the tube.

The central 20 inches of the Al₂O₃ tube was placed inside the stainless steel shell of the furnace and surrounded by eight kanthal 33 molybdenum disilicide resistance elements connected in series. These elements were surrounded by fibrous aluminum silicate and zirconia insulation. The temperature of the furnace was controlled using a R.I. Controls, silicon controlled rectifier MPRY thermac controller. This unit compared the electrical signals from an external programmer (representing the desired temperature) to those of a type S (platinum-platinum/10% rhodium) thermocouple (representing actual furnace temperature). The power for the system was supplied via a 6KVA stepdown transformer.

The external programmer consisted of a motor box and potentiometer variac geared to a heating rate of 50-75°C per hour. This heating rate was used in order to protect the Al₂O₃ muffle tube from thermal shock and, as previously pointed out, for better nitridation results. The temperature of the furnace was monitored to within ± 4°C using the type S thermocouple and a Honeywell 2705 potentiometer. The temperature of the sample being nitrided was measured
Figure A

Schematic of the furnace system used during the K series experiments

Figure B

A portion of the K series furnace system
Figure C

A portion of the K series furnace system used.
using the view port and a micro optical pyrometer measuring to \( \pm 5^\circ C \).

Three nitriding gases were used in the course of the investigation: \( N_2, 99\% N_2 - 1\% H_2 \) and \( 96\% N_2 - 4\% H_2 \). The system was also capable of supplying Ar gas for inert atmospheres. Prior to entering the reaction tube, the gases were first restricted by means of a Mathewson flow meter and valve and then passed over Si granules in a quartz tube maintained at 800\(^\circ C\) by tube furnace. This acted as an oxygen getter with the partial pressure of \( O_2 \) over Si at 800\(^\circ C\) being \( 1 \times 10^{-33} \) atmospheres. Next, the gases were passed over \( P_2O_5 \) which acted as an \( H_2O \) getter and then through a controlling Mathewson flow meter. The gas flow could then be passed by the furnace entrance (a zero flow rate in the furnace) or directly through the reaction tube. Upon exiting the system, the gas bubbled through 21 inches (53 cm) of water to an exit through the roof. The column of water was used to maintain a positive pressure throughout the system and to aid in leak detection.

Teflon tubing, .25 in (6.35 mm) in diameter was used throughout the system and connections made using .375 in (9.5 mm) brass swagelock fittings and valves.

Prior to nitriding, the green silicon bricks were weighed on a Christian Becker Chainomatic Scale to an accuracy of \( \pm 1 \) mg. The brick is then put on top of a silicon nitride sample holder which is placed on an \( Al_2O_3 \) sleeve. This sleeve acts to protect the \( Al_2O_3 \) muffle tube in case the exothermic nitridation reaction causes the Si to melt. The brick, sample holder and sleeve were then inserted into the reaction tube centering in the hot zone. The \( Al_2O_3SiO_4 \) heat shields were then inserted and the end caps made secure. The system was then made leak tight and the run started.

Generally, argon was used to purge the system and was allowed to flow through the tube at a rate between 40 and 70 cc/min. After purging for approximately 5-7 hours, the programmer was activated and the temperature has increased. When the system approached 800-900\(^\circ C\), (\( \sim 15 \) hours) the gas
was switched to one of the nitriding atmospheres which was allowed to flow through the tube at a rate of 100 cc/min. When the temperature exceeded 1000°C, the flow was generally reduced to 60-85 cc/min and rerouted to pass by the front of the reaction tube. This "easy exit" flow was used as an approach to a zero flow rate in the tube. As the reaction calls for nitrogen, it sucks it in from the infinite source. Otherwise, the gas does not see the sample at all. Whenever the 96% N₂ - 4% H₂ gas was used, a small flow 30-70 cc/min was maintained to avoid an explosive situation. During the reaction, N₂ is consumed but the H₂ is passive and will build up. Thus, a slight flow needs to be maintained. From this point, various time/temperature/atmosphere schedules were used lasting from hours to days and temperatures up to 1370°C. After the schedule was completed, the furnace was cooled via the programmer (again at 50-75°C per hour) until the temperature dropped below 200°C. The sample was removed, weighed and prepared for further investigations.

After nitridation, various metallurgical techniques were used to enable both qualitative and quantitative characterization of the material. This included a weight determination of percent reacted, x-ray diffraction for the relative amounts of phases present, optical and scanning electron microscopy for morphology identification as well as energy dispersive analysis of x-rays and electron microprobe analysis for elemental information. The mechanical properties of the nitrided samples were determined by four point bend tests for modulus, of rupture (MOR) and fracture toughness (K₁c) characteristics. The hardness of the reaction bonded material was also recorded. These methods were conducted similarly for both the K and E series nitrided samples.

Immediately after nitridation, the weight of the sample was recorded and compared to the previously recorded green weight. Assuming stoichiometric silicon nitride formed, the percent reacted can be calculated according to the equation:
\[
\frac{\text{nitrided weight} - \text{green weight}}{(.667) \text{ green weight}} \times 100
\]

It should be noted, that this method does not account for any material loss during the reaction (vapor mechanisms). Hence, 100% reacted is never attained.

A piece of the nitrided sample was ground into a fine powder by hand using an agate mortar and pestle. The mortar and pestle were only used for the grinding of silicon nitride and were carefully cleaned with acetone before and after each use. The resulting powder was then analyzed using a Phillips x-ray diffractometer. A Cu target tube with a Ni filter produced Cu-K\(\alpha\) radiation with a wavelength of 1.54 Å.

From the plot of angle (2\(\Theta\)) versus intensity, the interplanar spacings (d) of the prominent peaks were determined according to Bragg's laws \(\lambda = 2d\sin\Theta\), consulting the tabulated values of d spacings for silicon nitride in the powder diffraction files, specific intensity peaks were identified with the correlating diffraction planes. This data was then normalized following the work by Gazzara and Messier [10] to obtain the weight fraction of the three phases (\(\alpha, \beta, \text{unreacted Si}\)). From these values, \(\alpha/\beta\) and \(\alpha/\text{Si}\) ratios were calculated.

The x-ray data compiled above was also used to determine the lattice spacings of the phases present. A computer program developed by Dr. Kirby Dwight [11] yielded a and c values for the hexagonal \(\alpha\) and \(\beta\) phases from input data consisting of the diffracting plane and the angle at which its peak was found. This method was capable of offsetting the variances caused during the set up and running of the diffraction apparatus.

Another piece of material was cut from each nitrided brick and mounted in bakelite. This sample was then polished following standard metallographic techniques on unlubricated 320, 600 and 600 soft grit silicon carbide papers.
with intermediate ultrasonically cleaning in acetone. Final polishing was accomplished using a Buehler AB Vibromet polisher. The platen was covered with a Jarrett System Pellon pan-w polishing pad on which the samples were loaded under 666 grams of pressure. A slurry of 1-4 micron cerium oxide (CeO) powder and distilled water was used as the polishing medium. The use of CeO required increased polishing times of from 3 to 5 days. However, this method avoided pitting that resulted from the use of the more conventional alumina mixtures. Afterwards, the samples were again ultrasonically cleaned in acetone in preparation for microscopic examination at magnifications of up to 750 diameters. While this method does not allow the identification of $\alpha$ and $\beta - \text{Si}_3\text{N}_4$ (the grain sizes being generally: $\alpha < 0.5 \text{ microns}$ and $\beta < 5 \text{ microns}$), it does reveal important morphological characteristics including pore size and structure and unreacted silicon.

An AMR 100A scanning electron microscope was used to achieve greater magnification and resolution of the various morphologies of the phases present in silicon nitride. Since silicon nitride is on the insulator side of semiconductors, a conductive coating of approximately 300 angstroms of gold was sputtered onto the silicon nitride prior to examination using a p5-2 sputtering unit.

The polished samples (now coated with gold) were examined and general morphologies photographed. Further, more detailed SEM observations were made on the fracture surfaces from the MOR and fracture toughness samples. Pores that were too small to see optically became visible using the SEM and $\alpha$ and $\beta$ grains could be distinguished. Furthermore, by careful examination of the fracture surface, the failure initiation flaw could often be determined.

Frequently during optical or scanning electron microscopy, investigation of localized peculiar morphological areas was carried out using analytical elemental analysis. An EDAX (Energy Dispersive Analysis of X-rays) unit
attached to the scanning electron microscope allowed direct elemental identification of specific areas. Occasionally, another elemental determination based on characteristic wavelengths was employed using an ARL EMX-SEM microprobe unit. The unusual morphologies were often associated with localized impurities by these methods.

The density of the reaction bonded silicon nitride was obtained by simple weight-to-volume ratios. The more accurate methods involving liquid displacements and dry-to-wet weights were precluded due to the complicated pore structure characteristic of this material. Green densities were often based on nitrided volumes since in the green state, the bricks are extremely fragile. This is considered valid in light of the small (< 1%) volume change upon nitridation.

Characterization of the mechanical properties of the reaction bonded silicon nitride was accomplished using three and four point bend tests. The sample configurations and the machine parameters have been described in the body of this report (see page 58). From the load at failure; the modulus of rupture (MOR) and the fracture toughness (K$_{1c}$) were calculated.

As noted elsewhere, the surface finish of the bend bar has a marked effect on the failure strength. Throughout most of the later part of this investigation, the bend bar preparation was performed in-house. Two grinding rigs were designed and machined, one for the width (0.250 in. face) and the other for the height (0.125 in. face). Final surface finish and sample dimensions were achieved with a diamond embedded grinding wheel adapted onto a micromatic wafering machine. The bars were then tested for MOR in the as-ground condition. K$_{1c}$ measurements were made on bend bars after the introduction of a machined notch.