Comparison of the Tensile, Creep, and Rupture Strength Properties of Stoichiometric SiC Fibers

H.M. Yun
Cleveland State University, Cleveland, Ohio

J.A. DiCarlo
Glenn Research Center, Cleveland, Ohio

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H.M. Yun
Cleveland State University
Cleveland, Ohio 44115

J.A. DiCarlo
National Aeronautics and Space Administration
Glenn Research Center
Cleveland, Ohio 44135

Summary

Tensile strength, creep strength, and rupture strength properties were measured for the following types of polymer-derived stoichiometric SiC fibers: Hi-Nicalon Type S from Nippon Carbon, Tyranno SA from Ube, and Sylramic from Dow Corning. Also included in this study were an earlier version of the SA fiber plus two recent developmental versions of the Sylramic fiber. The tensile strength measurements were made at room temperature on as-received fibers and on fibers after high-temperature inert exposure. The creep-rupture property data were obtained at 1400 °C in air as well as argon. Some fiber types showed strong effects of environment on their strength properties. These results are compared and discussed in terms of underlying mechanisms and implications for ceramic composites.

Introduction

Continuous silicon carbide fibers are being developed as reinforcement of ceramic matrix composites (CMC) for high temperature structural applications. Small diameter SiC fibers (~10 to 14 μm) produced by polymer pyrolysis with little or no oxygen content display high stiffness, high room temperature strength, and improved thermal stability in comparison to other types of ceramic fibers. Recent examples of these low-oxygen types are the Hi-Nicalon fiber from Nippon Carbon (microcrystalline β-SiC with ~5 nm grain size plus excess carbon and trace oxygen) and the Sylramic fiber from Dow Corning (stoichiometric crystalline β-SiC with ~100 nm grain size plus trace boron as a sintering additive). They have displayed improved creep and rupture properties compared to the first-generation oxygen-containing Nicalon SiC fiber (refs. 1 and 2). However, at temperatures above ~1300 °C, the nonstoichiometric Hi-Nicalon fiber displays weight loss, grain growth, and contraction. A drop in the fast fracture strength accompanies these phenomena, even though the creep strength of the fiber improved (refs. 3 and 4). In contrast, the Sylramic fiber showed no weight loss, grain growth, or contraction below 1500 °C. Thus the coarser grained stoichiometric Sylramic fiber is more thermally stable than the finer grained nonstoichiometric Hi-Nicalon fiber.

More recently, Nippon Carbon and Ube Industries have introduced their own stoichiometric SiC fibers, namely Hi-Nicalon Type S and Tyranno SA, respectively, which have grain sizes and tensile strengths similar to those of the Sylramic fiber. These fiber types differ, however, in their production processes and in key details of their microstructures. Improved tensile and creep-rupture strengths have also been achieved for the Sylramic fibers by special developmental processes which yield microstructures with smaller boron content and process-related flaws (ref. 6). The reduction in boron in the Sylramic grain boundaries improves the fiber creep strength; whereas the reduction in flaw size improves the fiber fast fracture strength. Thus there exists various process routes and microstructural possibilities for thermally stable stoichiometric SiC fibers.

The objective of this study is to determine key strength properties for a variety of stoichiometric SiC fibers that are currently available in continuous lengths. These properties include (1) fiber tensile strength before and after thermal exposure and (2) fiber creep strength and rupture strength at 1400 °C in air and in argon. These
results along with other property data for the various fiber types are then compared and analyzed in order to 
obtain a better understanding of the current status of stoichiometric SiC fibers as reinforcement for high 
temperature CMC.

Experimental Procedure

Six types of stoichiometric SiC fibers were examined in this study. These include the most recent version 
of the Hi-Nicalon Type S fiber; an earlier and more recent version of the Tyranno SA fiber (SA(1) and SA(2)); 
the Sylramic fiber; and two recent types of developmental Sylramic fibers (Syl(1) and Syl(2)). The SA(1) 
fiber appears to be a finer grained version of the currently available SA fiber (called SA(2) in this study). The 
Syl(1) and Syl(2) fibers were specially processed to reduce boron content (ref. 6). Typical properties of the 
fibers are listed in table I. To determine tensile strength, single fibers were tested to fracture at room temper-
ature using a 25-mm gauge length and a constant displacement rate of 1.27 mm/min. The creep-rupture 
properties were determined for single fibers at 1400 °C using the same fiber creep-rupture facilities and 
procedures reported in detail elsewhere (refs. 7 and 8). Creep deformation versus time was recorded at a 
constant deadweight load using either 25 or 100 mm hot zone lengths for the air furnace and a 115 mm hot 
zone length for the argon furnace.

Auger electron spectroscopy (AES) depth-profile analysis and thermogravimetric analysis (TGA) were 
conducted on the different fiber types. For the TGA studies, weight change measurements were made in air 
and argon from room temperature to 1500 °C. Also TEM analysis was used to evaluate grain size and 
chemistry in the cross sections of the fibers. AES and TEM were conducted on single fibers; while TGA was 
conducted on tows wrapped into small volumes with initial weights from ~30 to 50 mg.

Results

Auger Electron Spectroscopy (AES)

AES analyses were conducted at every 10 nm from the fiber surface to ~300 nm deep. Figures 1(a) to (c), 
show the chemical composition of the surface zone in the (a) Sylramic, (b) Syl(1), and (c) Syl(2) fiber types. 
The Sylramic fiber surface was enriched by carbon ~10 nm deep. Beyond that, this fiber displayed a stoicho-
metric composition (nearly same Si and C atomic concentration). Boron existed in this Sylramic fiber up to at 
least 300 nm and noticeable amounts of boron and titanium were present on the fiber surface. The Syl(1) fiber 
(fig. 1(b)) showed a clear outer B-N rich layer which extended over 150 nm deep. This B-N rich area was not 
pure, but contained carbon and silicon with the first ~70 nm deep zone being silicon-depleted. The amount of 
boron on the fiber surface correlates well with amount of free boron originally present in the bulk of the 
Sylramic fiber (refs. 6 and 10). The Syl(2) fiber (fig. 3(c)) showed a nearly pure C-rich zone formed from 
about 20 nm up to ~100 nm deep. In comparison to the Sylramic fiber, the boron content below the surface of 
the Syl(2) fiber was reduced.

TEM Microstructures

TEM thin foils were prepared on fiber cross sections. Typical microstructures near the fiber surface zone 
are shown in figure 2 for (a) Sylramic, (b) Syl(1), and (c) SA(1). All fibers showed equiaxed and faulted 
β-SiC grains with various sizes. The range of grain sizes for the Sylramic, Syl(1), and Syl(2) fibers were 
comparable, from ~70 to ~170 nm; while the grain sizes for the SA(1) and SA(2) fibers were mostly above 
~150 and ~200 nm, respectively. Some grain junctions of the Sylramic and Syl(1) fibers were occupied by 
TiB2 and the frequency was higher for Sylramic than for Syl(1). This TiB2 phase was identified using both 
chemical elemental analysis and X-ray diffraction methods. The SA(1) fiber showed no second phase, even 
though aluminum compounds were reported to be utilized as sintering additives (ref. 9).
TGA Weight Change

The change of initial weight of the SiC fiber tows was measured from room temperature up to 1500 °C at a warm-up rate of ~5 °C/min. The results for various types of SiC fibers are shown in figure 3(a) for air and in figure 3(b) for argon. Weight changes above 100 percent indicate a weight increase, and below 100 percent, a weight decrease. For the SiC fibers in general, weight increases above 500 °C can be related to the formation of silica and other impurity oxides on the fiber surfaces; while the decreases below ~800 °C are typically related to removal of the initial sizing or excess carbon from the fiber surfaces.

Figure 3 shows that TGA behavior was dependent on the individual fiber types, with measurable differences beginning near ~250 °C. For example, the SA and Hi-Nicalon S fibers showed an abrupt weight decrease at ~250 °C, but the Sylramic fiber showed only a small decrease up to ~450 °C. This can be related to the amount of fiber sizing material; that is, less sizing for Sylramic and more sizing for SA and Hi-Nicalon S. From ~450 °C to ~1000 °C, the Sylramic fiber showed weight gain up to ~1.3 percent in air and ~0.4 percent in argon; however, the Syl(1), SA(2), and Hi-Nicalon S fibers showed only ~0.2 to 0.5 percent increase in air and <~0.1 percent increase in argon. (The weight gain in argon is believed to be due to trace oxygen in the argon.) The weight gain for the Sylramic fiber at intermediate temperatures in air and (impure) argon is probably due to the presence of boron and TiB2 on the fiber surface. That is, it is well known that liquid boria, which melts at 450 °C, can rapidly enhance silica growth on SiC materials (ref. 11). The developmental Syl(1,2) fibers with reduced boron appear to measurably reduce this intermediate temperature effect. The large weight loss near 700 °C in air for the Syl(2) fiber can be attributed to its carbon-rich surface layer (see fig. 1(c)). Weight changes for the various fibers at intermediate temperatures are summarized in table II.

From 1000 to 1500 °C in argon (plus trace oxygen), most SiC fibers tested showed a continuous weight increase of ~0.2 percent; while the Sylramic reached a maximum then showed a weight decrease above ~1200 °C. This drop might be attributed to the loss of boron from the borosilicate glass formed on the Sylramic fiber surface. In air near 1100 °C, the rate of weight change showed a minimum for most fibers, but not for the Sylramic and Syl(1,2) fibers. Also up to 1500 °C, the total amount of weight gains for the SA(2) and Hi-Nicalon S fibers were lower than for the three Sylramic fiber types. This enhanced oxidation might be attributed to the boron containing second phases that exist on the surfaces and in the bulk of the Sylramic fiber types.

Tensile Strength

The room-temperature tensile strengths are shown in figure 4 for the Hi-Nicalon S, SA(1,2), Sylramic, and Syl(1,2) fibers after fabrication and after exposure at 1400 °C for 1 hr in argon or at 1000 °C for 3 hr in vacuum. The strength values at each condition were the average of at least 12 single fibers removed from multifilament tows. Also shown are the standard deviations in strength. In general, the tensile strengths of the as-fabricated Sylramic, Syl(1,2), and SA(1) fibers were the highest, above 2800 MPa; while those of the as-fabricated SA(2) and Hi-Nicalon S fibers were below 1800 MPa, particularly the SA(2) fiber which showed a low tensile strength of ~1350 MPa. It should be noted that this as-fabricated strength for SA(2) is lower than that measured by the manufacturer (see table I), but is in qualitative agreement with the measured strength of SA(1) which is finer grained.

After exposure in vacuum or argon, the average tensile strength of the Sylramic fiber decreased by ~15 percent. A similar decrease was observed for the Hi-Nicalon S fiber after 1400 °C exposure. However, the Syl(1) fiber showed negligible tensile strength change after high-temperature exposure. These strength retention results are conflicting because most stoichiometric SiC fibers are reported to be stable after inert environmental exposures up to 1700 °C (ref. 9). It may be possible that at intermediate temperatures from 1000 to 1400 °C, the fiber surface morphology was altered, giving rise to new surface flaws that could lower the tensile strength (ref. 12). Thus the impurity type and content on the surface of a given fiber type plus the chemical composition of the exposure environment may play important roles in the fiber’s strength retention behavior.
Creep Strength

Typical creep curves are shown in figure 5 for the various stoichiometric SiC fibers at 1400 °C and 250 MPa in air. The Sylramic, Syl(1,2), and Hi-Nicalon S fibers displayed a short transient primary creep stage followed by pseudo-steady state creep; while the SA(1,2) fiber showed a large transient stage with no significant pseudo-steady state creep. Figure 5 shows that under the given test conditions, the Syl(1,2) fibers are the most creep-resistant stoichiometric SiC fibers. The next most creep resistant fiber was the Hi-Nicalon S. For prolonged times at 1400 °C, the Sylramic fiber ruptured without any tertiary creep; while the Syl(1,2) and Hi-Nicalon S fibers did not rupture up to 100 hr. As expected from the grain size difference, a measurable improvement in creep behavior was observed for the SA(2) fiber in comparison to the SA(1) fiber. However, both fibers crept to above 1 percent before rupturing. This amount of rupture strain appears to be abnormally large for a stoichiometric SiC fiber with a crystalline size of >100 nm.

The effect of stress on the 10 hr-creep strain at 1400 °C for the stoichiometric SiC fibers is shown in figure 6 for (a) air and (b) argon test conditions. Also included in this plot for comparison is the Hi-Nicalon nonstoichiometric fiber (ref. 7). With increasing stress, the creep strain increased by a power dependence of −2 for most SiC fibers both in air and argon (ref. 13). One exception is the Hi-Nicalon S fiber which indicated a lower stress dependency of −1. For all stresses, 10-hr creep strain of the Syl(1,2) fibers was reduced by a factor of −10 in comparison to the Sylramic fiber. Similarly the SA(2) fiber was more creep resistant than the SA(1) fiber by a factor of −3. Creep resistance of the Hi-Nicalon S fiber was between Sylramic and Syl(1,2). The creep resistance of the Syl(1,2) fibers was comparable to that of a low-boron containing α-SiC fiber with larger grain size (up to 2 μm) (ref. 7).

Figure 6 also shows that for the Sylramic, Syl(1), and Hi-Nicalon S fibers, the test environment had a significant effect on creep and creep strength. For these stoichiometric fibers at the same stress, the creep in argon was generally greater than in air. The creep strengths for 0.1 percent creep in 10 hr at 1400 °C are summarized in table III. In air, creep strengths for the Sylramic, Hi-Nicalon S, and Syl(1) fibers were about 120, 310, and 520 MPa, respectively; while the strengths in argon were about <50, 210, 230 MPa, respectively. The significant drop in creep resistance for the Syl(1) fiber in argon is not fully understood. This behavior can be contrasted with the creep strengths for the SA fibers and the nonstoichiometric Hi-Nicalon fiber (ref. 7) which showed no significant air/argon effect.

Rupture Strength

Results for rupture time versus stress at 1400 °C are shown in figure 7 for the various fibers in (a) air and (b) argon. The Syl(1,2) fibers were the most rupture resistant in air. Below ~500 MPa, these fibers did not rupture; while the other fibers ruptured below ~250 MPa. In air, the next most rupture-resistant fiber was Hi-Nicalon S. Like creep strength, the Syl(1) fiber showed a lower rupture strength in argon than in air. In argon, the SA(1,2) and Hi-Nicalon S fibers exhibited higher rupture strength than the Sylramic and Syl(1) fibers. For the Sylramic, Syl(1), SA(2), and Hi-Nicalon S fibers, the 10 hr rupture strengths in air were about 200, 720, 230, 310 MPa; while those in argon were <50, 210, 310, and 280 MPa, respectively. The rupture strengths are summarized in table III. Generally for all fibers, the rupture strains in air were stress and time independent, but were about a factor of two larger in air than in argon. Apparently in the 1400 °C air environment, oxide formation blunted strength controlling flaws on the fiber surfaces; while in the argon environment, flaw blunting by silica formation was not available.

Discussion

Improvement in the fast fracture strength and long-term creep and rupture strengths of SiC fibers is desired for the successful application of structural CMC. Whereas these properties have been improved by the development of stoichiometric SiC fibers, further enhancement would be of benefit. This study has shown that one possible direction is by the use of improved processes for currently available fibers as, for example, those used to convert Sylramic to the developmental Syl(1,2), SA(1) to SA(2), and Hi-Nicalon to Hi-Nicalon S.
Clearly the high temperature creep and rupture properties of these new fiber types are improved; but the fast fracture tensile strengths of some fiber types, such as the SA and Hi-Nicalon S are reduced by ~20 to 40 percent. Currently it is believed that the enhanced creep-rupture properties for these new fibers are probably not all explained by the same mechanism. For example, based on a preliminary microstructural observations, the improvement of the creep and rupture strengths observed for Syl(1) was not directly related with an increase in grain size; while the SA(2) creep improvement can be related to an increase in grain size. On the other hand, the improvements observed for Hi-Nicalon S appear to be related to a variety of mechanisms: an increase in grain size, a reduction in oxygen impurity, and a restructuring of excess carbon (refs. 3 to 14). For the Sylramic fiber, which contains a small amount of boron sintering additives, it would appear that the creep-rupture improvements seen for Syl(1,2) can be related to a reduction in the boron content along the grain boundaries. AES depth profiles (fig. 1) do indeed indicate boron depletion inside the fibers. A similar effect might be expected for the SA fibers if the aluminum sintering additives could be removed (ref. 9). However, although this may be possible by simple heat treatments, there is the possibility that new pores or flaws will be created plus the SiC toughness could decrease (ref. 15). In addition, as with the transition from SA(1) to SA(2), the fiber grains may grow, resulting in fibers with lower tensile strengths.

Currently it appears that there does not exist a certain fiber microstructure which will allow one type of stoichiometric SiC fiber to behave the best under all CMC application conditions. The Syl(1,2) fibers possess the best tensile strengths and the best creep strengths in air and argon. These fiber types also possess the best rupture strengths in air, but in argon the rupture strengths of the SA fibers are best. Also, the most creep resistant fibers, Syl(1,2) and Hi-Nicalon S, display an air-argon environmental effect: that is, they creep more in argon. On the other hand, the more creep-prone SA(1,2) and Hi-Nicalon (ref. 7) fibers did not display a significant environment effect. For CMC applications, the results in argon might be considered representative of behavior of fibers embedded in an uncracked CMC; whereas the results in air are probably representative of the behavior of fibers in a cracked CMC exposed to an oxidizing environment. Clearly the fibers should be expected to experience a variety of CMC environments, ranging from inert to highly oxidizing. Thus, at the present time it is not clear which type of stoichiometric SiC fiber would be best for the majority of CMC applications. It is also unclear as to the mechanisms giving rise to the environmental effects observed for the some of the stoichiometric fibers of this study.

### Summary and Conclusions

The stoichiometric SiC fibers evaluated in this study were found to display many **similar** properties that are beneficial for CMC fabrication and structural performance:

- Small diameter: 10-14 μm. A good size for weaving and braiding multifilament tows into CMC fiber architectures.
- Grain size: ~100 nm. Probably maximum size for good as-produced fiber strength and good creep resistance.
- Tensile strength (as-produced and after thermal exposure): >2 GPa for all fibers except the SA(2) fiber of this study. Important for providing CMC graceful failure after CMC fabrication.

The stoichiometric SiC fibers, however, also displayed many **dissimilar** properties which must be examined further under actual CMC fabrication and application conditions in order to fully understand their impact on CMC performance:

- Thermal conductivity: SA > Syl(1,2) > Syl > Hi-Nic-S. Important for reducing thermal stresses in high-temperature CMC applications.
- Surface roughness: not measured here but differences are expected based on surface grain size. High roughness can increase CMC interfacial shear behavior.
• Surface chemistry: Sylramic fibers with boron sintering additives display enhanced oxidation between 450 and 1000 °C due to boria formation. May result in enhanced fiber strength degradation and fiber-fiber and fiber-matrix bonding in cracked CMC.

• Creep strength: depends on fiber environment and grain boundary phases introduced during fiber processing. Syl(I,2) fibers with reduced boron sintering additives are best in air and argon; SA fibers with aluminum sintering additives are worst under same conditions. For creep-prone matrices, fiber creep controls CMC creep.

• Rupture strength: depends on fiber environment, creep resistance, and rupture strain. Creep-prone SA fibers rank highest because of relatively high high rupture strains in air and argon. Fiber rupture directly controls CMC rupture life.

References


### TABLE I. NOMINAL PROPERTIES OF NEAR STOICHIOMETRIC β-SiC FIBERS

(Currently available in continuous lengths)

<table>
<thead>
<tr>
<th>Trade name</th>
<th>Manufacturer</th>
<th>Process, (Sintering T. &gt;1600 °C)</th>
<th>Average grain size, nm</th>
<th>Second phases or impurity conc. (wt.%)</th>
<th>Density, g/cm³</th>
<th>Average diameter, mm</th>
<th>Number of filaments</th>
<th>PVA sizing, wt %</th>
<th>Modulus at RT (GPa)</th>
<th>Tensile strength, MPa</th>
<th>Tensile elongation, percent</th>
<th>Thermal cond. (W/mK)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hi-Nicalon S</td>
<td>Nippon carbon Polymer D./Sintering</td>
<td>22 (XRD)</td>
<td>Al (&lt;2), O(0.2)</td>
<td>3.0</td>
<td>13</td>
<td>500</td>
<td>PVA</td>
<td>400-420</td>
<td>120</td>
<td>&lt;0.6</td>
<td>18.4</td>
<td></td>
</tr>
<tr>
<td>SA(1)</td>
<td>Ube Industries Polymer D./Sintering</td>
<td>~150</td>
<td>TiB₂ &lt;0.1</td>
<td>3.02</td>
<td>10</td>
<td>800</td>
<td>PVA</td>
<td>375</td>
<td>520</td>
<td>-0.9</td>
<td>64.6</td>
<td></td>
</tr>
<tr>
<td>SA(2)</td>
<td>Ube Industries Polymer D./Sintering</td>
<td>&gt;~150</td>
<td>O(0.3), TiB₂</td>
<td>3.1</td>
<td>10</td>
<td>800</td>
<td>PEO</td>
<td>280</td>
<td>510</td>
<td>-0.7</td>
<td>46.0</td>
<td></td>
</tr>
<tr>
<td>Sylramic</td>
<td>Dow Corning Polymer D./Sintering</td>
<td>~100</td>
<td>Reduced B, TiB₂</td>
<td>3.1</td>
<td>10</td>
<td>800</td>
<td>PEO</td>
<td>&gt;400</td>
<td>&gt;310</td>
<td>-0.8</td>
<td>&gt;46</td>
<td></td>
</tr>
<tr>
<td>Syl.(1) Devel.</td>
<td>Dow Corning Polymer D./Sintering</td>
<td>~100</td>
<td>Reduced B, TiB₂, B-N</td>
<td>&gt;3.1</td>
<td>10</td>
<td>800</td>
<td>PEO</td>
<td>&gt;3200</td>
<td>&gt;3200</td>
<td>-0.8</td>
<td>&gt;46</td>
<td></td>
</tr>
<tr>
<td>Syl.(2) Devel.</td>
<td>Dow Corning Polymer D./Sintering</td>
<td>~100</td>
<td>Reduced B, TiB₂, C</td>
<td>&gt;3.1</td>
<td>10</td>
<td>800</td>
<td>PEO</td>
<td>&gt;3200</td>
<td>&gt;3200</td>
<td>-0.8</td>
<td>&gt;46</td>
<td></td>
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* Measured on TEM microstructures
* PEO sizing.
* At room temperature.

### TABLE II. WEIGHT GAIN OF VARIOUS SiC TOWS AT INTERMEDIATE TEMPERATURES

<table>
<thead>
<tr>
<th>Fiber type</th>
<th>Sizing material</th>
<th>Test conditions, temperatures, °C</th>
<th>Weight gain, percent</th>
</tr>
</thead>
<tbody>
<tr>
<td>SYL(RAMIC)</td>
<td>PVA</td>
<td>320 to 1000</td>
<td>Air 1.28</td>
</tr>
<tr>
<td>SYL.(2)</td>
<td>460 to 1000</td>
<td>-0.73</td>
<td></td>
</tr>
<tr>
<td>SYL.(1)</td>
<td>500 to 1000</td>
<td>Air 0.24</td>
<td>Argon -0.04</td>
</tr>
<tr>
<td>Hi-NIC. S</td>
<td>PVA</td>
<td>430 to 980</td>
<td>Air 0.18</td>
</tr>
<tr>
<td>SA(2)</td>
<td>PEO</td>
<td>470 to 1000</td>
<td>Air 0.26</td>
</tr>
</tbody>
</table>

* Determined by TGA.
* Argon contains trace oxygen.

### TABLE III. STRENGTH PROPERTIES OF AS-RECEIVED AND TREATED SYLRAMIC FIBERS

<table>
<thead>
<tr>
<th>Fiber type</th>
<th>Creep-rupture Type</th>
<th>Creep-rupture strength at 1400 °C, MPa</th>
<th>Hi-NIC. Type S</th>
</tr>
</thead>
<tbody>
<tr>
<td>10 hr/0.1 percent creep</td>
<td>Air</td>
<td>120</td>
<td>520</td>
</tr>
<tr>
<td>10 hr-rupture</td>
<td>Air</td>
<td>200</td>
<td>720</td>
</tr>
<tr>
<td>10 hr/0.1 percent creep</td>
<td>Argon</td>
<td>&lt;50</td>
<td>230</td>
</tr>
<tr>
<td>10 hr-rupture</td>
<td>Argon</td>
<td>&lt;50</td>
<td>210</td>
</tr>
</tbody>
</table>
Figure 1.—AES depth profile of surface of Sylramic SiC fiber types, (a) Sylramic, (b) Syl. (1), and (c) Syl. (2).

Figure 2.—TEM microstructures of SiC fibers, fiber cross-sections.
Figure 3.—TGA weight change of sized SiC fibers (a) in dry air and (b) in argon. Warm-up rate—5 °C/min.
Figure 4.—Room temperature tensile strength of SiC fibers: as-received and after exposure in inert environments.

Figure 5.—Typical creep-rupture curves for various stoichiometric SiC fibers.
Figure 6.—Stress-effects on creep strain of SiC fibers at 10 hour and 1400 °C: (a) in air and (b) in argon.

Figure 7.—Rupture strengths of SiC fibers at 1400 °C: (a) in air and (b) in argon.
Tensile strength, creep strength, and rupture strength properties were measured for the following types of polymer-derived stoichiometric SiC fibers: Hi-Nicalon Type S from Nippon Carbon, Tyranno SA from Ube, and Sylramic from Dow Corning. Also included in this study were an earlier version of the SA fiber plus two recent developmental versions of the Sylramic fiber. The tensile strength measurements were made at room temperature on as-received fibers and on fibers after high-temperature inert exposure. The creep-rupture property data were obtained at 1400 °C in air as well as argon. Some fiber types showed strong effects of environment on their strength properties. These results are compared and discussed in terms of underlying mechanisms and implications for ceramic composites.