AlN–SiC-particle-reinforced composites have been prepared at \( \leq 1400°C \) using submicrometer AlN, \(-325 \text{ mesh} \) \( \alpha \)-SiC particles, and polymethylsilane (PMS; \(-\text{(CH}_3\text{SiH)}_{n}-\)) via a polymer infiltration and pyrolysis (PIP) process. PMS is an organometallic SiC polymer precursor that can be modified with 16 wt% cross-linking aid to provide mPMS. mPMS converts to nanocrystalline \( \beta \)-SiC with >80% ceramic yield (1000°C in argon) with some excess (<5 wt%) graphitic carbon. mPMS has been used successfully as a nonfugitive binder for AlN–SiC compacts. Densities of 2.5 versus 2.1 g/cm\(^3\) have been obtained after nine PIP cycles for disk-shaped compacts formulated with and without mPMS binder, respectively. \( \alpha \)-SiC seeds crystallization of \( \beta \)-SiC derived from mPMS at temperatures as low as 1000°C. Some evidence suggests that AlN–SiC solid solutions form at particle/matrix interfaces.

I. Introduction

Particulate-reinforced ceramic-matrix composites (PRCs), when properly designed and processed, can offer superior high-temperature strength and stiffness, chemical inertness, and high specific strengths of potential use in advanced structural applications.\(^1\)–\(^6\) Unfortunately, conventional processing of PRCs often results in microstructures with broad grain-size distributions that severely limit mechanical properties.\(^6\) Efforts to realize PRCs with better mechanical properties have recently focused on producing nanocomposite materials. There are now several reports on the processing and properties of fully dense AlN–SiC and Al\(_2\)O\(_3–\)SiC nanocomposites.\(^7\)–\(^23\) As anticipated, these nanocomposite PRCs exhibit high-temperature fracture toughness, mechanical strength, and creep resistance superior to conventionally processed PRCs. These improvements are attributed to the refined microstructure and introduction of compressive stresses.\(^10\)\(^,\)\(^11\)\(^,\)\(^23\)\(^,\)\(^24\)

There appears to be significant impetus for developing a wide variety of nanocomposite materials. Unfortunately, the literature on nanocomposite materials processing and property evaluation is quite limited, most likely because of inherent processing difficulties. Conventional processing methods are rarely amenable to processing nanocomposites, because the target grain sizes in the finished PRC mandate either ultrafine (<100 nm), crystalline starting powders or amorphous powders that subsequently crystalize. Even with appropriate materials, processing conditions must be chosen to avoid uncontrollable grain growth at temperatures necessary for consolidation. Other issues include difficulties in obtaining low-cost, high-purity, unagglomerated, nanosized ceramic powders, especially for non-oxides.\(^25\)

Hot pressing (HPing) is the current method of choice for producing dense, nanocrystalline PRCs.\(^8\)–\(^16\)\(^,\)\(^20\)–\(^24\) For example, amorphous Si-C-N precursor powders can be HPed under nitrogen to form crystalline Si\(_3\)N\(_4\) matrices reinforced with nanosized SiC particles.\(^22\) Unfortunately, many processing constraints render HPing an impractical production method.\(^26\) A novel alternative to crystallization of amorphous powders relies on spinodal decomposition of solid solutions to provide nanocomposite PRCs, e.g., as found in the AlN–SiC system.

The structural similarity between AlN and 2H-\( \alpha \)-SiC favors interdiffusion and formation of solid solutions, rather than grain growth, when powders are HPed at >2000°C.\(^8\)–\(^10\)\(^,\)\(^12\)–\(^16\)\(^,\)\(^20\)–\(^23\) Annealing the resulting solid solution (\( \leq 1900°C \)) causes multidirectional spinodal decomposition to an AlN–SiC nanocomposite with thermomechanical properties superior to the individual components. Kou and Virkar\(^27\) report modulated AlN–SiC nanocomposite creep rates 3 times lower (values not provided) than AlN–SiC solid solutions with identical grain sizes. Huang and Jih\(^13\) find \(-20%\) increase in AlN–SiC nanocomposite hardnesses.

AlN–SiC PRCs can be obtained by pressureless sintering if additives (Y\(_2\)O\(_3\), Al\(_2\)O\(_3\), and CaO)\(^7\)\(^,\)\(^17\)–\(^19\)\(^,\)\(^28\)–\(^30\) are incorporated that allow liquid-phase sintering. These additives leave glassy phases at grain boundaries that provide improved toughness; however, their low viscosity can lead to unacceptable creep rates at high temperatures.\(^28\)–\(^30\)

Because of the above, considerable motivation remains to improve access to nanocomposite PRCs. Viable alternatives must offer short process times, simple processing methods, and access to structurally complex shapes. Low-cost, low-temperature (<1600°C), pressureless approaches that provide compositional and microstructural control are most preferable. Organometallic precursors offer one such alternative, because they transform at relatively low temperatures to nanocrystalline materials.\(^31\)–\(^35\) For example, the low-cost SiC precursor, polymethylsilane (PMS; \(-\text{(CH}_3\text{SiH)}_{n}-\)) transforms to phase-pure and nanocrystalline SiC at temperatures \( \approx 1000°C \).\(^36\)

Unfortunately, only one brief report describes a precursor route to AlN–SiC materials.\(^31\) and (to our knowledge) no reports discuss processing AlN–SiC composites via powder/precursor mixtures, despite the obvious attractions of this approach. Thus, the primary objectives of the work reported here are to (1) process AlN–SiC nanocomposites using polymer
infiltration and pyrolysis (PIP) of submicrometer AlN powder compacts.\textsuperscript{37–41} (2) determine if submicrometer AlN (crystallographically similar to α-SiC) seeds SiC (PMS) crystallization in heat-treated AlN–PMS mixtures, and (3) study the processability of AlN–SiC composites using PMS binder and α-SiC seeds. Secondary objectives include optimizing density gains per PIP cycle and evaluating the potential for forming AlN–SiC solid solutions at \(<1600°C.\) For practical purposes, in the work reported below, a 10 PIP cycle limit is the upper boundary chosen to achieve the above goals.

In the PIP process, a precursor liquid or solution is infiltrated into a shaped, porous ceramic body by capillary action or vacuum impregnation.\textsuperscript{1,37–41} The filled ceramic body then is heat-treated to convert the incorporated polymer to a ceramic. Repeated cycles are required to produce sufficiently dense components. Acceptable densities depend on the actual application, with structural components typically requiring near theoretical densities.\textsuperscript{1,42}

PIP processing combines conventional shaping methods with low-temperature processing. Shaping and densification with PIP are always independent. Moreover, when the precursor is used as a nonfugitive precursor binder, (1) binder removal is obviated, (2) higher final densities are achievable, and (3) control of final microstructure is obtained.\textsuperscript{33} Thus, complex and near-net-shape parts can be fabricated in a cost-effective manner. In principle, most of the above criteria can be met with this approach. However, obtaining fully dense PRCs via PIP processing is nontrivial. The green densities of powder compacts (or fiber preforms for fiber-reinforced composites (FRCs)) are typically quite low, requiring multiple PIP cycles to achieve full density.\textsuperscript{39,41}

Fortunately, a wide variety of low-cost organometallic and metalloorganic precursors for PIP processing are now available. The literature describes precursors to SiC, Si\(_n\)N\(_n\), AlN, TiC, and aluminosilicates (celsian, mullite, and cordierite).\textsuperscript{35–38,41,44–48} To be useful for PIP processing, precursors must offer (1) ease of handling, (2) thermal stability, and environmental stability, (3) well-defined molecular weight (MW) distribution and controllable rheology, (4) matrix and reinforcement wetting, and (5) a well-defined composition upon pyrolysis.\textsuperscript{33,35,40} These attributes are typically controlled by the polymer chemistry, architecture, and processing conditions, as discussed in detail elsewhere.\textsuperscript{33}

Theoretically, the best precursor for SiC is PMS, with a number average molecular weight (\(M_n\)) of 2100, with some excess carbon to obtain phase-pure SiC, because unmodified PMS typically offers only moderate ceramic yields and generates SiC containing excess silicon.\textsuperscript{36}

II. Experimental Procedure

(1) General

(A) Materials: AlN powder (99% pure) was obtained from Tokuyama America, Inc. (San Mateo, CA), and α-SiC and β-SiC powders (both 99% pure) were obtained from CERAC, Inc. (Milwaukee, WI) and H. C. Starck, Inc. (Newton, MA), respectively. (See Fig. 1.) The reported average particle size of the Grade-F AlN powder (lot 48040) and α-SiC powder (lot X13385) were 0.3 and 10–20 μm, respectively. The as-received powders were heated (10°C/min under argon) to 1000°C in 1 h to remove adventitious water or organics.

(B) Pyrolysis Conditions: All pyrolyses \(<1200°C were conducted in a sealed and ported quartz tube (5.7 cm throat ID \(\times\) 71.1 cm length) under argon (100 mL/min) using a tube furnace (Furnace Model HTF 55322A and Controller Model CC58114PC, Lindberg/Blue M, Watertown, WI). Pyrolyses to 1400°C were run under argon (100 mL/min) in a high-temperature tube furnace (Model F54348CM, Barnstead/Thermolyne, Dubuque, IA) in a sealed and ported mullite tube equipped with a water-cooled aluminum end cap.

(C) Composite Processing: Two types of composites, with and without mPMS binder, were produced. No other additives were used to avoid contamination. The synthesis of mPMS has been described elsewhere.\textsuperscript{48} mPMS or mPMS-4, unless otherwise specified, refers to a precursor with C\(_{1,41}\)Si\(_{1,00}\)B\(_{0,020}\) stoichiometry. Three other derivatives were also studied (see Table I), but mPMS-4 was optimal for PIP processing.

(2) Sample Fabrication and Heat Treatment

Three types of composite disks were made: (1) pure AlN with mPMS-4 binder, (2) AlN–α-SiC without mPMS-4 binder, and (3) AlN–α-SiC with mPMS-4 binder. Disks fabricated with pure AlN had insufficient green strength and exhibited shear failure; pure AlN with mPMS-4 binder compacts pressed easily but were weak after heat treatment, and surface cracks were visible. AlN–α-SiC composite disks fabricated using a 9:1 (by weight) ratio powder mixture provided good green strength and ease of handling. This powder ratio was established empirically but not optimized. α-SiC was used as a reinforcement (1) because the AlN–α-SiC mixtures with mPMS-4 demonstrated the best handling characteristics and (2) to minimize confusion with polymer-derived β-SiC.

(A) Pure AlN and AlN–α-SiC Compacts with mPMS Binder

All operations were conducted in an argon glove box (Model MO-40-2 Dri-Lab glove box, Vacuum Atmospheres, Hawthorne, CA). Dilute mPMS-4 (~12 wt% polymer in toluene/tetrahydrofuran (THF)) was added to pure AlN to obtain a 3:1 and 1:3 composition of AlN and mPMS-4 after vacuum removal of solvent. A 9:1 (by weight) AlN–α-SiC powder mixture was mixed thoroughly. Then, dilute mPMS-4 was added to this mixture to obtain, after vacuum removal of solvent, (AlN–α-SiC)\(_{1-x}\) (mPMS-4) compositions where \(x = 10, 20, 25,\) or 30 wt%.

The dried powder/binder mixture (typically ~0.4 g) then was poured into a double-action stainless-steel die (12.72 mm ID ×
50.8 mm height). The assembled die was removed from the glove box and uniaxially compressed to 138 MPa (20 ksi) (Model C laboratory press, Carver, Wabash, IN). The compressive load was removed after the pressure had stabilized for ≥5 min and the disk ejected from the die under a protective inert atmosphere (typically nitrogen gas).

The disks (12.72 mm × 1.9 mm) were transferred rapidly onto graphite-foil-lined alumina holders inside a sealed and ported quartz furnace tube. The quartz tube was purged with argon (10 min) and then heated (10°C/min under argon) to 1000°C to convert the mPMS to β-SiC (SiC (mPMS)). After heat treatment, the quartz tube was returned to the glove box, following three degassing steps in the antechamber. The disk dimensions and weight were noted with digital calipers (Model CD-6° C, Mitutoyo Corp., Tokyo, Japan) and a precision balance (Model PJ360 DeltaRange®, Mettler, Hightstown, NJ).

(B) AlN–α-SiC Compacts without mPMS Binder: A 9:1 (by weight) ratio powder mixture was mixed thoroughly in air (~15 min) and then ultrasonicated for 10 min (Model 1200, Bransonic Ultrasonic Cleaner, Danbury, CT). The powder mixture (typically ~0.4 g) then was carefully poured into a double-action 316-stainless-steel die in air and compressed uniaxially, as above. The compressive load was removed after the pressure stabilized for ≥5 min and the disk ejected.

Disks (12.7 mm × 1.9 mm) were degassed under vacuum (~1 torr (~133 Pa)) in the antechamber of an argon glove box. They then were transferred into the glove box, placed on graphite-foil-lined alumina boats (6.0 cm × 3.0 cm × 1.4 cm), sealed in a ported quartz tube, and then transferred out for heat treatment. Disks without binder were heated in the quartz tube (10°C/min under argon to 1000°C) to eliminate adsorbed water, surface coatings, or organics introduced during sample preparation. The quartz tube was resealed and returned to the glove box after heat treatment. The disk dimensions were recorded as above.

(3) Polymer Infiltration and Pyrolysis Process

Disks (with and without α-SiC) were immersed for 12 h in dilute (12 wt% solids loading in toluene/THF) mPMS in tightly capped vials. The disks were removed from the vials, dried, placed into the quartz pyrolysis tube and transferred to a furnace. The heating schedules (Tables II and III) were designed according to the PMS thermogravimetric analysis (TGA) weight-loss behavior (Fig. 2). Disk weights and dimensions were recorded after each cycle and then polished (see below). The disks then were reimpregnated with mPMS solution as above and repyrolyzed.

Two different heating schedules were used. The initial 12 h program (Table II) was used to avoid damaging the relatively weak green compacts. The ramp rates for the second pyrolysis program were double those of the first.

Two procedures were used to ensure maximum polymer infiltration per cycle. The first involved gently polishing the disk (900 grit SiC paper, Mager Scientific, Dexter, MI) after each cycle to remove the thin SiC (mPMS) coatings that formed and inhibited subsequent infiltration. The second used intermediate 1200°C or 1400°C heat treatments to densify the SiC (mPMS) and reopen SiC (mPMS)-filled pores. The occurrence of the latter process was inferred from the improved infiltration behavior that results.

(4) Characterization

TGA studies were performed (Model Hi-Res TGA 2950 thermogravimetric analyzer, TA Instruments, Inc., New Castle, DE). Samples (15–30 mg) were placed in a platinum pan and heated under argon (60 mL/min) at 50°C/min (HiRes mode 4) from room temperature to 1000°C. Sample stabilities in air were assessed by placing 15–30 mg samples in the TGA furnace. A 30 s flow (60 mL/min) of prepurified, dry air (CO2 free) was introduced into the sample chamber after purging with argon (60 mL/min) for 5 min. The sample weight gain with time was recorded over a period of 5 h.

Differential thermal analyses (DTA) studies were performed (Model DSC 2910 module equipped with a 1600°C differential thermal analyzer cell, TA Instruments, Inc.). Samples (10–15 mg) were placed inside platinum sample cups lined with platinum foil and heated at 10°C/min under argon (50 mL/min) to 1400°C. α-Al2O3 was used as the reference material.

Diffuse reflectance infrared Fourier transform spectroscopy (DRIFTS) spectra of disk specimen surfaces were obtained (Model Galaxy Series FTIR 3020 spectrometer, Mattson Instruments, Inc., Madison, WI) equipped with a "praying mantis" DRIFTS accessory (Model DCA-02, Harrick Scientific Corp., Ossining, NY) and dry-nitrogen-gas purge assembly. Random cuttings of optical-grade KBr (International Crystal Laboratories, Garfield, NJ) were ground using a dry alumina mortar and pestle. The ground KBr powder (~0.150 g) was compacted uniaxially into a hardened 316-stainless-steel cylindrical die at 138 MPa (20 ksi) for 15 min. The resulting KBr pellet (~0.5 mm thick) then was placed above the specimen disk in a sample holder and transferred into the sample chamber. 250 scans were collected for each specimen at a gain of 20 and resolution of ±2 cm⁻¹.

Table II. Slow Ramp Rate Heating Program with mPMS-4 Binder

<table>
<thead>
<tr>
<th>Temperature range (°C)</th>
<th>Ramp rate (°C/h)</th>
<th>Dwell time (h)</th>
</tr>
</thead>
<tbody>
<tr>
<td>20–100</td>
<td>60</td>
<td>0.5</td>
</tr>
<tr>
<td>100–200</td>
<td>120</td>
<td>0.5</td>
</tr>
<tr>
<td>200–250</td>
<td>60</td>
<td>0</td>
</tr>
<tr>
<td>250–350</td>
<td>120</td>
<td>0.5</td>
</tr>
<tr>
<td>350–470</td>
<td>60</td>
<td>0.5</td>
</tr>
<tr>
<td>470–550</td>
<td>120</td>
<td>0.5</td>
</tr>
<tr>
<td>550–1000</td>
<td>300</td>
<td>1.0</td>
</tr>
<tr>
<td>1000–20</td>
<td>300</td>
<td>End</td>
</tr>
</tbody>
</table>

Table III. Accelerated Heating Program for Composites during mPMS Processing

<table>
<thead>
<tr>
<th>Temperature range (°C)</th>
<th>Ramp rate (°C/h)</th>
<th>Dwell time (h)</th>
</tr>
</thead>
<tbody>
<tr>
<td>20–100</td>
<td>120</td>
<td>0.5</td>
</tr>
<tr>
<td>100–200</td>
<td>240</td>
<td>0.5</td>
</tr>
<tr>
<td>200–250</td>
<td>240</td>
<td>0</td>
</tr>
<tr>
<td>250–350</td>
<td>240</td>
<td>0.5</td>
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<td>350–470</td>
<td>300</td>
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<tr>
<td>470–550</td>
<td>300</td>
<td>0.5</td>
</tr>
<tr>
<td>550–1000</td>
<td>600</td>
<td>1.0</td>
</tr>
<tr>
<td>1000–20</td>
<td>600</td>
<td>End</td>
</tr>
</tbody>
</table>

Fig. 2. Decomposition profile of vacuum-dried mPMS-4 in argon.
Spectra of as-received powders and AlN–mPMS powder mixtures were obtained by grinding a 1.0 wt% sample with cuttings of optical-grade KBr in a dry alumina mortar and pestle in the glove box. The ground mixture then was packed into a sample cup and transferred into the specimen chamber. 500 scans were collected for each sample at a gain of 4 and resolution of ±4 cm⁻¹.

X-ray diffractometry (XRD) was used to follow development as samples were pyrolyzed to 1000°C–1400°C (under argon for 1 h). Spectra were collected using a rotating anode goniometer (Rigaku Denki Co., Ltd., Tokyo, Japan). Standard operating conditions are described elsewhere.36,45

Field-emission-gun scanning electron microscopy (SEM; Model S-800, Hitachi, Ltd., Tokyo, Japan) was used for microstructural characterization. Samples were cut from disks using a high-speed precision saw (Model Isomet 2000, Buehler, Ltd., Lake Bluff, IL) equipped with a diamond wafering blade. Specimen surfaces were polished with 900 grit SiC paper and ultrasonically cleaned. An operating voltage of 5.0 kV was used.

III. Results and Discussion

We describe here efforts to develop processing–microstructural relationships for PIP-processed AlN–SiC composites made with mPMS as binder and infiltrant. In particular, we are interested in the influence of variations in processing conditions and additives on composite densification and microstructural development.

Three different PIP-processed AlN–SiC systems were studied: (1) pure AlN with mPMS binder, (2) AlN–α-SiC (9:1 (by weight)) without mPMS binder, and (3) AlN–α-SiC (9:1 (by weight)) with mPMS-4 binder. The first system shows conclusively that AlN does not seed SiC (mPMS) crystallization, as reported by Bill and co-workers and by us.49,50

A DTA profile of vacuum-dried mPMS-4.

\[
\begin{align*}
\text{CH}_3 \quad | \quad \text{Si} \quad | \quad \text{H} \\
\downarrow \quad \quad \downarrow \quad \quad \downarrow \\
\beta\text{-SiC} + 2\text{H}_2 \quad (1)
\end{align*}
\]

\[\Delta \text{Ar}\]

(B) DTA: Figure 3 shows the DTA events that occur during the polymer-to-ceramic conversion of vacuum-dried mPMS. This DTA is substantially different from that reported previously by Zhang.38 The first broad exotherm, centered at 160°C, appears to result from cross-linking reactions that render the polymer infusible.38 The second, sharp exotherm at 250°C is associated with a second curing mechanism likely resulting from thermally promoted hydrolysis, as discussed below. A small exotherm at 400°–500°C can be assigned to the Kumada rearrangement.36 Hydrogen-gas evolution begins at ~700°C,36 as indicated by a broad exotherm centered at 790°C. A final exotherm for β-SiC crystallization occurs at ~1030°C (maximum), as confirmed by XRD (see below).

(C) XRD: XRDs of mPMS-2 heated at 1000°–1400°C show the appearance of β-SiC (Table IV and Fig. 4). Similar to the work of Zhang, very broad peaks appear in samples heated to 1000°C, suggesting nanocrystalline material. Samples heated to 1200° or 1400°C undergo some grain growth, as evidenced by peak sharpening. However, the peaks remain broad, even at 1400°C, indicating that the material remains nanocrystalline. These observations are supported by TEM and nuclear magnetic resonance (NMR) studies to be published elsewhere.51 The average grain sizes (from TEM of SiC fibers from mPMS)36 are ~1 nm (1000°C), 10 nm (1200°C), 25 nm (1400°C), 60 nm (1600°C), and 200–500 nm (1800°C).51

The amount of cross-linking agent present in PMS strongly affects CY, phase purity (Fig. 5), and stoichiometry. Unmodified PMS exhibits CYs of ~55 wt% at 1000°C and forms β-SiC with excess silicon (up to 20 wt%45. In contrast, mPMS-2 (see Table I for composition) has a CY of ~80 wt% at 1000°C, with only β-SiC detected by XRD. mPMS-3 gives an 85 wt% CY with ~1 wt% carbon. Finally, mPMS (mPMS-4) provides a CY of ~85 wt% and forms β-SiC with ≤5 wt% graphitic carbon (see XRD section). The carbon contents are estimated from observed CYs and initial precursor compositions.

XRDs of mPMS-1 show only β-SiC at 1200°C (1 h), with excess silicon appearing only at 1400°C. There are several possible explanations for this behavior. We know from TGA results that an ~3% mass loss occurs at 1200°–1400°C. Below 1200°C, considerable hydrogenated, amorphous SiC is present, which is undetectable by XRD. At ≥1200°C, it appears that isolated hydrogen atoms begin to diffuse at reasonable rates, leading to formation and evolution of hydrogen gas and the generation of free silicon, which undergoes near-melt crystallization.

For the PIP studies, we select a version of mPMS (mPMS-4) that provides excess carbon on pyrolysis. It had been anticipated that this excess carbon would carbothermally reduce the oxide surface layers present on the particulate-reinforcing pow-

\[
\text{CH}_2\text{CH}_2 + \text{SiH}_4 \rightarrow \text{SiH}_4 \text{CH}_2\text{CH}_2
\]
ders, e.g., SiO$_2$ on SiC, providing access to well-controlled interfaces. Prochazka and Scanlan$^{52}$ find that submicrometer SiC powders sinter effectively when both boron and carbon additives are used. In contrast, precursor-derived oxide-free SiC sinters when boron is added but not carbon,$^{36}$ suggesting that excess carbon aids removal of surface oxide layers.

(D) **DRIFTS:**

DRIFTS spectra of SiC (mPMS) formed at $\approx 1000^\circ$C suggest the presence of some oxygen, as indicated by a small $v_{Si-O}$ band (1040 cm$^{-1}$). The oxygen content can be estimated from the $v_{Si-C}$ and $v_{Si-O}$ peak ratios. Tsuge $^{53}$ report that mixtures containing SiC and SiO$_2$ exhibit a linear relationship between the SiO$_2$ concentration and observed $v_{Si-O}$ intensity. On this basis, 1100°C SiC (mPMS) has <1 wt% oxygen.

Oxygen incorporation likely results from slow oxidation (by adventitious oxygen) of retained Si–H bonds, to form Si–O bonds. No $v_{Si-O}$ bands are present in samples heated to 1200°C, indicating oxidation is avoided or SiO loss occurs. This suggests that precursor processing must be conducted with care before full mineralization at $\approx 1200^\circ$C to limit impurity incorporation.

DRIFTS spectra of SiC (mPMS) heated to $\approx 1200^\circ$C (Fig. 6) show broad peaks attributable to SiC (900 and 780 cm$^{-1}$).$^{36,53,54}$ The $\delta_{Si-CH}$ (1400 cm$^{-1}$), $\delta_{Si-CH-Si}$ (1355 cm$^{-1}$), and $\nu_{Si-H}$ (2100 cm$^{-1}$) absorptions observed for 1000°C (2 h) samples are absent in spectra of 1200° and 1400°C samples, in accord with the mass loss attributed to hydrogen-gas evolution seen in the TGA. These changes indicate that the mPMS-to-SiC transformation is incomplete until $\approx 1200^\circ$C.

The breadth of the Si–C peak ($\approx 850$ cm$^{-1}$) observed in the 1000°C DRIFTS is a result of the highly disordered nanocrystalline material produced.$^{25,36}$ As the pyrolysis temperature is increased from 1200° to 1400°C, the 850 cm$^{-1}$ peak sharpens and becomes better resolved, indicating better ordering and crystallinity.

(3) **AlN–SiC Powder Characterization**

(A) **XRD Analyses:** The XRD patterns of as-received AlN and $\alpha$-SiC powders (average particle size of 0.3 and 10–20 $\mu$m, respectively, Fig. 1) are typical of highly crystalline hexagonal phases (see Table V for powder diffraction file cards). One difficulty with XRD analysis of the AlN–SiC system is that many peaks overlap. For example, (102), the most intense peak for $\alpha$-SiC (35.66° 2$\theta$) is obscured by the AlN (002) peak (36.04° 2$\theta$). The difficulty increases when both $\alpha$- and $\beta$-SiC are present with AlN, because all four peaks of the $\beta$ phase coincide with those of the $\alpha$-phase (Table IV). In principle, the microstructural evolution of SiC (mPMS) should be easy to follow given the broad XRD peaks typically generated from nanocrystalline $\beta$-SiC. However, this becomes impossible when seeding occurs.

(B) **DRIFTS:**

The as-received AlN material exhibits a peak centered at $\approx 770$ cm$^{-1}$, with a smaller shoulder (possibly Al–O–N) as reported in literature.$^{55,56}$ This shoulder increases slightly after 1000°C for 1 h under argon treatment, perhaps from crystallization. The AlN DRIFT spectra also show low-intensity $\nu_{OH}$ and $\nu_{NH}$ peaks. The low intensities appear to be artifacts of the analytical method. DRIFTS (Fig. 7) of as-received and heat-treated AlN with $\approx 1$ wt% oxygen (manufacturer’s analysis) should exhibit more-intense $\nu_{OH}$ (3300–3500 cm$^{-1}$) and $\nu_{NH}$ peaks than actually observed.

Spectra of the coarse $\alpha$-SiC powders are difficult to obtain.
because the large particles cause significant specular reflectance and signal attenuation. Thus, submicrometer α-SiC particles (CERAC, Inc., Milwaukee, WI) were substituted (identical XRDs). Two main absorption peaks for α-SiC are present, with the primary peak centered at 900 cm⁻¹ and a smaller peak at 780 cm⁻¹. DRIFTS spectra for submicrometer α- and β-SiC (CERAC) are indistinguishable. Both materials appear oxygen-free, because no ν_{Si-O} peaks are visible, despite the likely presence of surface Si–O.

(4) Polymer Infiltration and Pyrolysis Processing

The green compact density is controlled by the (1) ceramic particle-size distribution, (2) concentration of mPMS-4 binder used, and (3) compaction pressure. A 138 mPa (20 ksi) uniaxial compaction pressure is most effective in producing strong and relatively dense compacts. Compacts pressed with AlN or AlN–α-SiC powders exhibit green densities of ~1.65 g/cm³ (51% of theory, rule of mixtures). The advantages of using mPMS-4 binder include (1) higher compact densities, (2) formation of more-uniform SiC (mPMS) coatings on the ceramic particles, (3) introduction of large charge of mPMS in the initial step, and (4) better potential for forming the continuous porous network required for PIP processing.

The green densities of disks with mPMS binder (20, 25, and 30 wt%) are slightly lower than the binder-free compacts at 1.60–1.65 g/cm³. However, the 1000°C densities (zero PIP cycles) are significantly higher, ~1.85 g/cm³ or 60% of theory. This increase results primarily from partial sintering, because disks typically lose ~20 vol%. Optimum densification is observed for 20–25 wt% binder loadings, which agrees well with theory.

To our knowledge, this apparent precursor-promoted sintering at 1000°C has not been reported before. There are several possible reasons sintering might occur, some based on reaction chemistry arguments and some based on physical rearrange-

(5) Infiltration Studies/Density Changes

Preliminary studies were made at shorter times; however, 12 h immersion times were then used for all PIP studies. Bubble streams appeared immediately upon sample immersion and occasionally obscured the sample completely. The rate of bubble evolution decreased slowly for the first four PIP cycles, but decreased rapidly after four cycles.

The rate of density gain from mPMS-4 uptake with the number of PIP cycles was similar for all samples with or without mPMS-4 binder and/or α-SiC particles. This result was significant, because the observed sintering in samples with mPMS-4 binder apparently did not hinder subsequent infiltration efforts. This implied that sintering was not accompanied by significant pore closure.

The gases evolved during mPMS conversion to SiC may have been important in maintaining a continuous pore structure. Because samples with mPMS-4 binder had higher green densities, fewer PIP cycles were required to achieve higher densities. Moreover, the maximum densities achieved were also higher with mPMS-4 binder. For example, the maximum densities obtained for mPMS binderless disks were ~2.2 g/cm³ after nine PIP cycles and 1400°C heat treatments (Fig. 8). In comparison, the highest densities obtained for samples with 20 wt% mPMS binder after nine PIP cycles were ~2.5 g/cm³ (theoretical density is ~3.1 g/cm³ by rule of mixtures).

The above densities were obtained using a staged heating/infiltration schedule. Disks were cycled at 1000°C for three or four cycles, followed by an intermediate cycle at 1200°C. The advantages of repeated PIP cycling to 1000°C included shorter processing times and minimum use of high temperatures. Processing at 1200° or 1400°C led to higher SiC (mPMS) densities (ρ = 2.4 g/cm³), and was used to reopen capillary pores. This resulted in improved precursor infiltration during subsequent cycles.

Slower heating rates were used during the first heat treatment of the more-fragile green compacts to minimize gas evolution. Faster heating rates (to 1000°C) were possible in subsequent cycles, because the compact’s mechanical integrity increased considerably beyond the zeroth PIP cycle. Heat treatments to 1400°C were performed at up to 25°/C/min. Samples were heat-treated at 1400°C for 1 h under argon when mPMS-4 uptake did not improve after a 1200°C heat treatment.

The 1400°C heat treatments gave no further sintering/
densification, in contrast to the work of Zhang,36 who showed that SiC (mPMS) continued to densify at higher temperatures, approaching full density (3.21 g/cm$^3$) at ~1800°C. The absence of densification could be explained if the added α-SiC seeded crystallization, because the SiC formed was fully dense, which appears to occur as discussed below.

The final densities of ~2.5–2.6 g/cm$^3$ obtained at nine PIP cycles indicates the presence of residual, closed macroporosity of ~20%, assuming all of the mPMS-4 transforms to fully dense SiC. Therefore, additional modifications are required to achieve higher density samples within the preset 10 PIP cycles limit, perhaps by using higher polymer concentrations during impregnation, higher heat-treatment temperatures, or HPing. However, this degree of porosity is typical of PIP, CVI, and CVD methods of producing composites.

(6) Microstructural Changes

(A) SEM: SEM of AlN–α-SiC–mPMS disk cross sections (Fig. 9; $p = 2.5$ g/cm$^3$ after nine PIP cycles at 1400°C maximum temperature) reveals few macroscopic voids. However, substantial micrometer-sized porosity is evident. Some clogged pores are visible (Fig. 9(a)) near the specimen surface (within ~100 μm).

Three separate phases can be identified by their distinct morphologies. The submicrometer AlN particles are the most abundant and appear unchanged by the PIP process (Fig. 9(c) and (d)). The edges of several α-SiC particles are visible. These large α-SiC particles are almost completely surrounded by the much smaller AlN particles. Nanocrystalline SiC appears in the third phase, primarily in the filled pore channels (Fig. 9(b)).

The observed macroscopic voids (arrows in Fig. 9(a)) are likely cracks from compressive stresses caused by the drying polymer. Similar cracks are observed by Tu and Lange39,40 in polymer-infiltrated powder compacts. These authors find that removing the surface polymer layer before pyrolysis reduces the compressive stresses and prevents surface cracks from occurring. The cracks in AlN–SiC compacts are filled during subsequent PIP cycles. The presence of the cracks was discovered only on sectioning. Crack healing near the surface appears complete, because the original crack edges are barely observable. However, healing appears limited to cracks ≤400 μm deep (Fig. 9(a)). The limit on infiltration depth may result because of surface pore collapse and residual gases trapped within the crack, which prevent complete pore filling. If true, improved densities may be accessible simply by pressure- or vacuum-assisted infiltration.

(B) XRD: Work on SiC–SiC composites conducted in our laboratory and a related HRTEM study by Bill and co-workers10 indicates that SiC powders seed SiC (mPMS) crystallization. The similarity in the basal plane lattice parameters12,15–18 between 2H-AlN and α-SiC suggests that AlN may seed SiC (mPMS) crystallization. This possibility has been tested by mixing submicrometer AlN powder and mPMS-4 in a 3:1 ratio (by weight) and heating the mixture to different temperatures. The resulting materials have been characterized by XRD, TGA, and DRIFTS. Three AlN–mPMS-4 experi-

![Fig. 9. SEMs of AlN-α-SiC disk after nine PIP cycles: (a) primary area of observation, (b) view of a pore channel filled with SiC (mPMS), (c) magnified view of a crack region, and (d) phases present near the crack.](image-url)
ments have been conducted to determine the effect of AlN concentrations on the crystallization behavior of SiC (mPMS). XRD analyses of 3:1 (by weight) mixtures of AlN–SiC (mPMS) pyrolyzed at 1200°C and further heated to 1400°C suggest that SiC (mPMS) does not nucleate on AlN particles. Broad peaks, typical of SiC (mPMS) are visible at 1200°C and remain broad even at 1400°C. The graphitic carbon normally seen with SiC (mPMS) derived from mPMS-4 is missing, presumably because of re-oxidation at the surface as the Al–N and Si–C bands are much diminished.

The second possibility is that nanocrystalline SiC seeds mixed with AlN–mPMS-4; XRDs of pyrolyzed AlN–α-SiC disks infiltrated with mPMS-4 (Fig. 10) suggest that seeded crystallization occurs. Figure 10(a) is from a green, as-pressed AlN–α-SiC disk with no mPMS-4 binder. The other XRD patterns are of a single disk with mPMS-4 binder at different processing stages.

No SiC peaks result from the as-pressed disk, despite the 10 wt% α-SiC. After three 1000°C PIP cycles, peaks at 35.6° (assigned to β-SiC) and 59.98° ± 2° appear. The (101) peak for 6H-α-SiC (34.09° ± 2°, 35% of most intense peak) is used to distinguish between the α and β phases. The enhanced intensity and narrow β-SiC peaks shown in Fig. 10 indicate much greater crystallinity than shown in Fig. 5. No changes are observed in the XRD pattern when the same disk is heated to 1200°C, without additional PIP cycles (Fig. 10(c)).

However, significant changes are detected after a fourth 1000°C PIP cycle followed immediately by heating to 1400°C. The (101) peak for 6H-α-SiC (34.09° ± 2°) is now present (Fig. 10(d)), while the SiC peak at 35.6° ± 2° increases slightly; the AlN peak at 36.04° ± 2° is used as a reference.

Several explanations can account for these observations. The first is that the β-SiC phase is present at 1000° and 1200°C, and partially transforms to α phase after heating to 1400°C. The second possibility is that nanocrystalline α-SiC is formed with β-SiC, even at 1000°C, and grain growth occurs when heated to 1400°C, leading to the appearance of the peak at 34° ± 2°. SiC (mPMS) typically contains minor amounts (~15 wt%) of α-SiC. 

The first explanation is considered more plausible. Because AlN does not seed crystallization of SiC (mPMS), the 10 wt% α-SiC CERAC powder used in the compact must be responsible for the enhanced crystallization observed. However, the absence of the 34° ± 2° peak at 1000° and 1200°C suggests that α-SiC promotes crystallization of β-SiC (in accord with Bill and co-workers50), and the β- to α-phase transformation occurs only at 1400°C. The observed peaks are much sharper than normally expected for SiC (mPMS).

The (110) peak splitting observed at 59.40° and 59.95° ± 2° (in Figs. 10(b–d)) is unusual and, based on the work of Miura et al.,17 suggests that the formation of an AlN–SiC solid solution has occurred. Normally, the AlN and SiC (110) peaks overlap (as in Fig. 10(a)); however, XRD results by Miura et al.17 indicate similar (110) peak separation at 59.46° and 59.93° ± 2°. These authors claim that the peak shifts result because AlN–SiC composites with modulated nanoscale AlN-rich–SiC-rich texture are produced by spinodal decomposition of AlN–SiC solid solutions. Thus, one interpretation of the peak splitting we observe is that an AlN–SiC solid solution forms that subsequently produces an AlN–SiC nanocomposite. Further work is required to validate this possibility.

(C) TGA: The predicted and actual weight losses for these mixtures are consistent (within 1%) with that expected from the behavior of the individual components on heating to 1000°C. For example, the actual AlN–SiC (mPMS) weight (1000°C, TGA) is 0.350 g, whereas the weight predicted assuming an 85 wt% CY for mPMS-4 (1000°C) is 0.351 g. This suggests that the entire processing system is well behaved. No further weight loss is detected above 1000°C, in keeping with the seeded crystallization. This also suggests that hydrogen loss is accelerated by seeding.

(D) DRIFTS: DRIFTS has been used to complement the XRD data. Because many different peaks (Al–N, Al–O, Si–C, Al–O) overlap, care must be taken in making peak assignments. The first step is to establish baseline spectra for the individual materials. Peaks obtained are also compared with literature values.33,55

DRIFTS can be highly sensitive to sample crystallinity, particle size, and bond type but is not as structure sensitive as XRD, because it probes at the atomic scale. For instance, minute amounts of SiO2, not detected by XRD, are detectable by infrared methods. In contrast, α- and β-SiC, easily distinguishable by XRD, are indistinguishable by infrared methods.

Application of DRIFTS to the disks requires a modified, nondestructive, nonstandard approach. The sample thickness and opacity rule out use of transmission infrared methods. Instead, a method based on Koenig and co-workers57–61 is used, wherein thin layers of ground KBr powder are used to differentiate between glass substrates and polymeric surface coatings. The amount of powder (or KBr layer thickness) used controls the probing depth.59,60

To maintain sample consistency, a fixed amount of ground KBr powder is pressed into a die to give KBr pellets with a consistent thickness and good surface uniformity. These pellets can be used repeatedly, further reducing the error within a group of samples tested. The sample surfaces are not polished, because flat surfaces give specular rather than diffuse reflectance and lead to a lower signal-to-noise ratio. Moreover, the sampling depth for DRIFTS can be quite shallow (several micrometers) and is affected by sample reflectivity.57–61

Figure 11(a) is a DRIFTS spectrum for a green AlN–α-SiC disk with no SiC (mPMS). Two main absorption bands at 800–900 cm−1 (νSi–C) and centered at 700 cm−1 (νAl–N) are observed. Figure 11(b) is the spectrum of a sample heat-treated to 1000°C. The broad band (mixture of νSi–O and νAl–O) at 1040 cm−1 results as Si–H groups react with the 1 wt% oxygen initially present in the AlN. The AlN bands are likely masked by the larger Al–N and Si–C) and centered at 700 cm−1 (νAl–N) are observed. Figure 11(b) is the spectrum of a sample heat-treated to 1000°C. The broad band (mixture of νSi–O and νAl–O) at 1040 cm−1 results as Si–H groups react with the 1 wt% oxygen initially present in the AlN. The AlN bands are likely masked by the larger Al–N and Si–C bands. The predicted and actual weight losses for these mixtures are consistent (within 1%) with that expected from the behavior of the individual components on heating to 1000°C. For example, the actual AlN–SiC (mPMS) weight (1000°C, TGA) is 0.350 g, whereas the weight predicted assuming an 85 wt% CY for mPMS-4 (1000°C) is 0.351 g. This suggests that the entire processing system is well behaved. No further weight loss is detected above 1000°C, in keeping with the seeded crystallization. This also suggests that hydrogen loss is accelerated by seeding.

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Figure 11(c) is similar to Fig. 11(a), except that a new peak
The feasibility of forming nanocrystalline AlN–SiC composites and solid solutions via a low-temperature PIP process with mPMS was studied. The use of organometallic polymer precursors allowed processing of nanocrystalline CMCs at relatively low temperatures (<1600°C). A carbon-rich mPMS (mPMS-4) was used as a reinforcing phase and nonfugitive binder in AlN–α-SiC composites. The excess carbon was added to provide phase-pure SiC (mPMS) by eliminating silicon formation and to aid removal of oxide layers on the particulate phases. Higher densities (~2.5 g/cm³ or 78% of theory) were obtained in samples with mPMS-4 binder, compared to samples without binder (~2.2 g/cm³). The mPMS binder caused the powder compacts to partially sinter at 1000°C without appreciable pore collapse, because density gains from subsequent PIP processing were unaffected.

The effects of adding 10 wt% (~325 mesh) of α-SiC particles to AlN–mPMS-4 powder compacts were considerable. XRD studies indicated that the α-SiC seeded crystallization of SiC(mPMS). In contrast, AlN particles were found to have no effect on SiC(mPMS). Higher green compact strengths and crack healing also were attained with α-SiC addition. However, surface cracks in samples with α-SiC healed during PIP processing. In contrast, the surface cracks grew, rather than healed, in samples of AlN with mPMS-4.

XRD and density gain results showed that composites of AlN–SiC with 20–30 wt% SiC (mPMS) were formed, rather than complete solid solutions at the temperatures used, i.e., ~1400°C. However, there was one piece of evidence suggesting that partial solid-solution formation may have occurred, namely, the peak separation observed in at 60° 2θ. XRD data further suggested that β-phase SiC (mPMS) formed at 1000°C and 1200°C but partially transformed to α-phase at 1400°C.

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