Characterization of SiC (SCS-6) Fiber Reinforced Reaction-Formed Silicon Carbide Matrix Composites

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ABSTRACT

Silicon carbide (SCS-6) fiber reinforced-reaction formed silicon carbide matrix composites were fabricated using NASA’s reaction forming process. Silicon-2 at% niobium alloy was used as an infiltrant instead of pure silicon to reduce the amount of free silicon in the matrix after reaction forming. The matrix primarily consists of silicon carbide with a bi-modal grain size distribution. Minority phases dispersed within the matrix are niobium disilicide (NbSi$_2$), carbon and silicon. Fiber push-out tests on these composites determined a debond stress of $\sim$67 MPa and a frictional stress of $\sim$60 MPa. A typical four point flexural strength of the composite is 297 MPa (43.1 KSi). This composite shows tough behavior through fiber pull out.
INTRODUCTION

In recent years, there has been an increasing demand for high performance and light weight composite materials for aerospace and other high temperature structural applications. Silicon carbide-based advanced ceramics and composites which show high strength and toughness, oxidation resistance, and high thermal conductivity, have attracted attention for a variety of gas turbine engine applications [1-2]. Fabrication of gas turbine components requires a process with near-net or complex shape capabilities. Currently, silicon carbide matrix composites are fabricated by a number of techniques; hot pressing [3-4], polymer pyrolysis [5], chemical vapor infiltration [6], and melt infiltration [7,9]. Hot pressing, hot isostatic pressing, and sintering require high temperatures for processing (typically around 1500-1800°C). Also, it is very difficult to attain full density due to the constraint of the fibers. In addition, complex and near-net shape requirements are additional problems to overcome. In the polymer pyrolysis process, multiple infiltrations are needed. Further, this process yields microcrystalline and often microcracked matrices [8]. Another widely used technique is chemical vapor infiltration (CVI) which
is a very slow process requiring long processing time (often weeks). In addition, CVI cannot yield full density: composites typically have 10-15% residual porosity [8]. In the SILCOMP™ and other related processes, which are based on molten silicon infiltration, there is very little control over green body porosity and thick interface coatings are required to prevent molten silicon attack [7, 9].

From the above, it is clear that the majority of current composite processing techniques are either expensive or have limitations in complex shape and near-net-shape fabrication. Therefore, it is necessary to develop a processing approach which has near-net and complex shape capabilities and that is also cost effective.

The reaction forming process reported in this work has many advantages over the processing techniques described above. It can be used for the fabrication of silicon carbide matrix composites with large diameter monofilaments or small diameter fibers [10-12]. It also provides very good control of residual phases (refractory disilicides and silicon). Since the starting material is a liquid polymer, second phases can be incorporated
very easily. In this process, a microporous carbon preform containing fibers is infiltrated with molten silicon or silicon - refractory metal (molybdenum, niobium) alloys [10-12]. The infiltrating materials react with the carbon and form matrices of silicon carbide, refractory disilicides (MoSi$_2$, NbSi$_2$), and silicon. The relative amounts of these phases depends on the properties of the porous carbon preform and composition of the infiltration materials. These are controlled by the morphology and porosity of the carbon precursor matrix and the reaction conditions. This process yields composites which have full density.

The objective of this paper is to study the microstructure and mechanical properties of silicon carbide (SCS-6) fiber reinforced-reaction formed silicon carbide matrix composites fabricated by silicon-2 at% niobium alloy infiltration. Electron microscopy and microprobe analysis were used to determine the overall microstructure, the distribution of niobium disilicide, and to determine if there was any incorporation of the niobium apart from the disilicide. The interfacial shear strength and room temperature flexural properties of these composites are also discussed.
[2] EXPERIMENTAL PROCEDURES

To fabricate the composite matrix, a mixture of high char yield resin, a liquid pore-forming agent, and an acid catalyst is infused within a fiber preform. The resin mixture is polymerized and pyrolyzed to form a microporous carbon matrix. By varying the ratios of the constituents in the system, a wide variation of pore volumes, pore sizes, and porous carbon morphologies can be obtained. At the infiltration temperature, the molten silicon-2% niobium alloy reacts with the microporous carbon matrix to form silicon carbide. The rate of infiltration depends on the wetting of the preform by the melt. The wetting increases with increasing temperature. The selection of processing time and temperature are critical for the control of microstructure and interfacial reactions, and these control the mechanical properties of the composites.

Cross-sectional samples were metallographically mounted and polished for optical microscopy, scanning electron microscopy (SEM), and electron microprobe analysis (EMPA). The optical microscopy, performed on a Richert MeF3 A metallographic microscope, included interference
layering, in which chemical differences are enhanced in reflected light after the deposition of a thin oxide layer on the sample. SEM was performed on a JEOL JSM 840A and EMPA was done using an ARL SEMQ microprobe. Transmission electron microscopy (TEM) thin foils were prepared using standard techniques, finishing with Ar ion beam milling. TEM was accomplished on a Philips EM400T operating at 120 keV with a Kevex Quantum thin window x-ray energy dispersive spectrometer for chemical analysis. Fiber push-out tests were carried out on the apparatus described in [13]. Flexural strengths were measured using four point flexural tests on an INSTRON 7551 frame and load deflection behavior was recorded using a LVDT probe.

[3] RESULTS AND DISCUSSION

3.1 Microstructure

Fig. 1 shows a polished section of the composite formed by the infiltration of the preform with a silicon - 2 at% niobium alloy. The uniformity of the reaction formed material reflects the well-controlled
porosity of the carbon preform obtainable using the present method. The phase diagram of the Nb-Si system [14] indicates the existence of a large two phase (liquid silicon and solid niobium disilicide) region. Due to the reaction of silicon with carbon during processing, the local concentration of silicon in the alloy melt decreases and the melt composition shifts toward the two phase region (liquid silicon and niobium disilicide) where niobium disilicide starts to precipitate from the melt. Electron microprobe compositional dot maps of the composite (Fig. 2) showed that the niobium was incorporated in small clusters of particles in the matrix and had not diffused into or reacted with the SCS-6 fibers. The EMPA further revealed that any residual silicon and the NbSi$_2$ were dispersed within the matrix.

Higher magnification scanning electron images of the fiber/matrix interface (Fig. 3) revealed that some regions of the fiber outer carbon-rich coating had reacted to form scalloped regions of SiC. These reactions are very limited and also strongly depend on the chemistry and morphology of the porous carbon preforms and processing conditions (time and temperature). It was observed that the liquid resin provided a thin carbon coating on the fibers during composite fabrication. The presence of this
thin carbon coating coupled with the well controlled porosity of the carbon preform is thought to be the leading contributor to the significant reduction in the reaction of alloy infiltrants with the fibers during processing. It should be noted that in other melt infiltration processed composites, 3-5 μm thick interfacial coatings are required to sufficiently protect the fibers [7,9]. The interface between the outer carbon-rich coating layer and the inner carbon-rich coating layer must be maintained to obtain good fiber pullout and composite-like behavior. That interface was usually undisturbed in the present approach.

3.2 TEM Analysis

The details of the matrix microstructure were studied using transmission electron microscopy. Optical and SEM micrographs showed a slight variation in contrast within the silicon carbide (Fig. 3). In SEM, there were two shades of contrast when backscattered electrons (BSE) were used to form an image, which suggests two slightly different average atomic numbers. Through comparison of the SEM and TEM analyses, it was determined that the regions giving the appearance of a higher average
atomic number (being brighter in backscattered electron contrast) contained finer grained SiC while the SiC in the darker BSE regions was more coarse. The cause of this effect is not certain. The two regions were easily differentiated in the TEM (Fig. 4). The silicon carbide grain sizes in the finer regions varied over a range from approximately 30 nm in the center of the cluster to nearly 1 \( \mu \)m near the larger matrix material. The grain size ranges in the coarser material were narrower, generally staying between 1 and 5 \( \mu \)m. Electron diffraction patterns from the silicon carbide grains were almost always consistent with the cubic '\( \beta \)' polytype. A few grains, however, had the hexagonal 6H structure. Both finer and coarser SiC grains were highly faulted, as is shown in Figs. 5 (a) and (b). The expected minority phases, NbSi\(_2\) and residual carbon were confirmed crystallographically and compositionally in the TEM. The NbSi\(_2\) grains were relatively featureless internally and irregularly shaped. Residual carbon was fibrous in structure and usually neighbored by fine grained SiC.

TEM also revealed the fiber/matrix interface microstructure. The fiber core and surrounding silicon carbide were identical to as-received fibers,
as was the inner carbon-rich layer of the fiber/matrix interface (Fig. 6a). The outer carbon-rich layer, however, often reacted with the matrix to form very fine-grained silicon carbide (Fig. 6b), which was essentially identical to the finer grained matrix material. The boundary between the two carbon-rich layers, characteristic of the SiC (SCS-6) fiber coating system, was usually cracked in TEM thin foil samples. This indicates a relatively weak interface and possible residual stresses on the fiber by the matrix after processing.

3.3 *Interfacial Shear Strengths*

As was mentioned earlier, the silicon carbide (SCS-6) fiber used in this study has an outer carbon-rich coating applied by the manufacturer. In a brittle fiber-brittle matrix composite, it is necessary that a low shear strength material separate the fiber and matrix to give good mechanical properties to the composite. Fig. 7 (a) shows a typical load-displacement curve for fiber push-out tests. The average debond and frictional stresses for a 0.65 mm sample are $67.2 \pm 15.1$ MPa and $60.8 \pm 14$ MPa, respectively. Fig. 7 (b) and (c) show typical micrographs of pushed-out fibers. The
majority of fibers debond between the outer carbon layers of the SiC (SCS-6) fibers with some debonding occurring between the fiber and its carbon layers.

3.4 Flexural Properties

The mechanical behavior of the silicon carbide (SCS-6) fiber reinforced -reaction formed silicon carbide composites was determined using four-point flexural tests at room temperature. Fig. 8 (a) shows a room temperature flexural stress vs. displacement curve for the silicon-2 at% niobium alloy infiltrated composite. The average room temperature flexural strength of composites with ~18 vol% fibers is 43.1 KSI. The stress-displacement curve (in Fig. 8a) shows 'non-elastic deflection' or strain behavior in these composites, demonstrating an improvement in toughness of these composites due to addition of fibers. The fractograph in Fig. 8 (b) also shows fiber pull-out and tough composite behavior.

[4] SUMMARY OF RESULTS
The microscopic and EPMA results indicate that the niobium from the silicon-niobium alloy precipitates as niobium disilicide (NbSi$_2$) in the matrix. There was no evidence of niobium carbide (NbC, Nb$_2$C) formation at the interface. The silicon carbide matrix has a bi-modal grain size distribution with grain sizes in the range of 1-5 $\mu$m (large grains) and approximately 30 nm (small grains).

Fiber push-out tests indicate that the majority of fibers debond between the inner and outer carbon layers of the SiC (SCS-6) fibers. The average debond stress was $67.2 \pm 15.1$ MPa and the frictional stress was $60.8 \pm 14$ MPa. The room temperature flexural strength of this composite with $\sim 18$ vol% fibers is 43.1 KSi. Fractographs show fiber pull-out and tough composite behavior.

[5] CONCLUSIONS

The findings of this microstructural and mechanical property study suggest that dense SiC (SCS-6) fiber reinforced reaction formed silicon carbide matrix composites can be fabricated by the NASA reaction forming
process without the need for any additional interface coating on the SCS-6 fibers. Minimal damage to the fiber coating has been attributed to the microstructural uniformity of the precursor carbon matrix in the preform. In addition, the silicon-niobium alloy infiltrant results in the formation of a uniformly distributed niobium disilicide phase which could be advantageous in improving the fracture toughness of the matrix. Residual free silicon is also uniformly distributed throughout the silicon carbide matrix. This composite shows tough composite behavior through fiber pull out.

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REFERENCES


Figure Captions

Fig. 1: Interference layering optical micrograph of composite infiltrated with molten silicon-2 at% Nb alloy.

Fig. 2: Electron microprobe images of the composite formed using x-ray signals from niobium, carbon, and silicon as well as backscattered electrons (BSE's).

Fig. 3: Scanning electron micrograph of the fiber/matrix interface using BSE contrast showing scalloped reaction region of the outer carbon-rich layer.

Fig. 4: Transmission electron micrograph (TEM) showing the bimodal and separate SiC grain size distributions.

Fig. 5: Intragranular faulted microstructures shown using TEM of the finer (6a) and coarser (6b) SiC grains.

Fig. 6: Bright field TEM of the fiber/matrix interface. The SiC fiber/inner carbon-rich layer (7a) and outer carbon-rich layer/matrix SiC regions are shown.

Fig. 7: (a) Fiber push-out curve for composite infiltrated with Si-2 at% Nb alloy (sample thickness = 0.65mm) and (b-c) micrographs showing pushed fibers.

Fig. 8: (a) Load-deflection curve for silicon-2 at% niobium alloy infiltrated composites, and (b) fractopgraph showing tough composite behavior.
Fig. 2  Electron Microprobe Analysis of SiC Matrix Composites

Composite

Nobium

Carbon

Silicon
SEM Micrograph of the Fiber/Matrix Interface
(Backscattered Electron Contrast)

Note: Scalloped Reaction of Outer Carbon Layer with Matrix
TEM Analysis of Reaction-Formed Silicon Carbide Matrix

Fine and Coarser SiC Grains
TEM of Reaction-Formed Silicon Carbide Matrix Grains

Fine SiC Cluster

Coarser SiC Grain

Figs. 5
TEM Analysis of Fiber/Matrix Interface in Composites

Fiber / Inner Carbon Layer

(a)

Reacted Outer Carbon Layer / Matrix SiC

(b)
Fig. 7(a)
Scanning Electron Micrographs Showing Pushed-Out Fibers

Fig. 7