INTRODUCTION

Increasing aircraft engine efficiency and performance capabilities have been obtained for gas turbine components, such as, blades, vanes and disks, by clever materials and processing developments. Knowledge and understanding of processing-microstructure and mechanical properties relationships have been critical in these developments. The operating temperatures of superalloys have, however, now been pushed close to their incipient melting points. It is presently believed that further significant improvements in temperature capability can be obtained only by using the intermetallics, ceramics or refractory elements. These, however, are inherently brittle, at least at the room temperature. Effect of minor alloying additions (such as, boron, yttrium, in Ni$_3$Al) and innovative processing techniques (such as, rapid solidification, plasma processing) are being extensively investigated to overcome this major problem of alloy ductility.

The microstructures of these advanced composites are composed of micron size fibers distributed usually in a brittle matrix. Plasma sprayed cloth process combined with high temperature diffusion bonding is one of the leading composite fabrication technique. The nature of the fiber-matrix interface is very important in determining the physical and mechanical properties and environmental degradation of these composite materials. Study of the interface in materials with very diverse physical properties is therefore crucial in development of these composite materials. For example cracks may artificially be introduced into the material during specimen preparation because of the very brittle nature of the constituents and the interface. These cracks will make the selection of optimum material and its processing impossible, where occasionally the purpose is to quantify the
microcracks forming during the processing or thermal cycling of composite materials. Scanning electron and transmission electron microscopic examination of these materials requires skills which are very specialized. We have carried out such research under this cooperative agreement.

WORK PERFORMED DURING THE PROJECT:

(A) The first component, involving most of the effort, consisted of providing specialized scanning electron microscopy expertise and support for the other ongoing materials research programs in the Materials Division at the NASA-Lewis Research Center. Software and hardware updates to further extend the capability of the microscope were carried out. A range of materials such as, intermetallics, metal-matrix composites, ceramic-matrix composites, ceramics and intermetallic compounds, based on refractory elements were examined under this research. Crystal structure, size, shape and volume fraction distribution of various phases which constitute the microstructures were examined. Deformed materials were studied to understand the effect of interfacial microstructure on the deformation and fracture behavior of these materials. Specimens tested for a range of mechanical property requirements, such as, stress-rupture, creep, low cycle fatigue, high cycle fatigue, thermomechanical fatigue, etc, were examined. Microstructural and microchemical stability of these materials exposed to simulated operating environments were investigated. The EOIM Shuttle post-flight samples were also examined to understand the influence of low gravity processing on microstructure. In addition, fractographic analyses of Nb-Zr-W, titanium aluminate, molybdenum silicide and silicon carbide samples were carried out. Following are some of the specific highlights.

(1) Optimization of Microscope WDX Spectrometer on JEOL SEM: Optimization of the hardware and software in order to increase the capabilities of the scanning electron microscope resulted in use of synthetic layered diffracting crystal for the detection of C, O, N and B. The net result was a five fold increase in resolution, peak to background ratio, and almost a five fold decrease in the amount of current necessary to obtain satisfactory count rates.

(2) Seminar on Fractography of Composite Failure: A two day short course to introduce
fractography, the states of stress, fracture initiation and crack propagation in brittle, ductile, and composite materials, and composites was developed for presentation by NSF Center for Molecular and Microstructure of Composites which is a state/Industry/University consortium.

(3) SEM Examination of Post-Flight Shuttle Samples.

(4) Oxygen Embrittlement of Reinforced Orthorhombic Ti-Al-Nb Alloys.

(B) The second component consisted of extensive characterization of sapphire fibers in the fiber-reinforced composites made by powder cloth processing. Following conclusions can be drawn from this study on the deformation and fracture behavior during push-out testing of sapphire fibers in sapphire fiber reinforced Powder Cloth Processed (PM) and directionally solidified (DS) NiAl and NiAl(Yb) composites.

(1) Reaction between organic binder residues, used during powder cloth processing, and sapphire fibers results in the formation of gaseous porosities at the fiber-matrix interface during directional solidification of NiAl composites.

(2) Addition of ytterbium causes extensive reaction between the matrix and sapphire fibers. In the PM NiAl(Yb) composite the fiber is surrounded mostly by the oxygen rich NiAl layer, except for the Yb$_2$Al$_5$O$_{12}$ phase 'pegs' which grow from the fiber surface into the intergranular regions of the matrix. In the DS NiAl(Yb) composite the fiber is not in contact with NiAl; instead it is surrounded by a layer of Yb$_2$Al$_5$O$_{12}$ phase.

(3) The load versus displacement response during fiber pushout consists of three regimes: an initial region of constant slope, up to a stress $\tau_y$, followed by a region of decreasing slope up to a maximum stress $\tau_m$, the subsequent load drop and onset of constant stress, $\tau_f$. Occurrence of a strong acoustic emission signal near $\tau_y$ indicates that this stress corresponds to the onset of fiber-matrix debond. Fiber pushout occurs at $\tau_m$. The friction stress during sliding of the fiber is $\tau_f$.

(4) Grain boundaries in NiAl, in contact with the sapphire fibers, are the source of crack nucleation during fiber pushout. They are also the preferred path for the crack growth. The $\tau_y$ is therefore higher in the PM+DS NiAl (larger grain size and reduced number of grain boundaries in contact with the fiber) than in the PM NiAl.

(5) The PM NiAl(Yb) specimens show the highest, $\tau_y$, $\tau_m$ and $\tau_f$ values because of the
Yb₃Al₅O₁₂ 'pegs' between the fiber and the NiAl matrix, and also the surface roughness of the fibers due to the chemical reaction. These stress values are lower in the DS NiAl(Yb) composite because of the microcracks and shrinkage cavities at the Yb₃Al₅O₁₂-matrix interface.

(c) The third component was part of a general program at Cleveland State University to study pressure infiltration casting of metal matrix composites. A high pressure melt infiltration facility was designed and fabricated by support from NASA and other in-house funding. This facility was used to study the kinetics of pressure infiltration of ceramic fiber and particulate preforms by molten metallic and intermetallic alloys. This unique facility, is capable of casting low melting point alloys, such as copper or aluminum based, and also high temperature alloys, such as, superalloys and intermetallics. Refractory ceramic reinforced aluminum and copper alloys having potential for wear applications, and high temperature intermetallic matrix alloys for aircraft gas turbine applications, were examined using this facility. One particular aspect of pressure infiltrated composites was the potential damage on the fiber surface caused by the high melt temperature and pressures. This damage was studied by optical and scanning electron microscopy on the specimens prepared at Cleveland State University.
I. INTRODUCTION

The nickel base intermetallic, NiAl(β), is being extensively investigated for high-temperature structural applications because of its high melting point, low density, high modulus, good thermal conductivity, and outstanding oxidation resistance. However, NiAl suffers from low elevated-temperature strength and very poor low-to-intermediate-temperature toughness and plastic ductility. It is believed that these limitations can be overcome by refractory fiber reinforcement. Sapphire-fiber-reinforced NiAl composites, made by powder metallurgy techniques, are therefore being seriously investigated. The extent of strengthening and toughening imparted by the fibers is expected to depend on the microstructure of the fiber-matrix interface. Load-displacement measurements during fiber-pushout testing provide a measure of the interfacial shear strengths. However, a systematic study into the influence of the fiber-matrix interface on the deformation behavior during fiber pushout in sapphire-fiber-reinforced NiAl composites is lacking. Hence, the goal of the present study was to address these issues. In this study, two methods have been used to vary the microstructure at the fiber-matrix interface: (1) alloying of NiAl by ytterbium (Yb) and (2) use of solid-state heat treatment (HT) and directional solidification (DS) to modify the interfaces in the as-fabricated powder metallurgy (PM) sapphire-NiAl(Yb) composites. The powder-cloth (PC) process, which is essentially a PM technique, was selected for the fabrication of the composites in order to obtain fine, equiaxed-NiAl grains in contact with the sapphire fibers. Heat treatment and zone DS of the PC-processed composite feedstock were selected to enhance the extent of fiber-matrix reaction and to obtain large NiAl grains in contact with the fiber. Because NiAl does not react with alumina, the NiAl melt was not expected to wet the sapphire fiber. Ytterbium was selected as an alloying addition to NiAl, because ytterbium oxide is thermodynamically more stable than aluminum oxide and therefore a chemical reaction between the sapphire fibers and Yb in the matrix would be expected. It has been reported that the adhesion of the Al2O3 oxide scale to the Ni-Al alloys is enhanced by the addition of rare earth elements. Thus, the addition of Yb to the NiAl matrix could also lead to improved bonding between the sapphire fiber and the NiAl matrix. This study compares the fiber-matrix interfacial microstructure and the fiber-pushout response of PM, HT, and DS sapphire-NiAl(Yb) composites with interfacial microstructure and fiber-pushout behavior of PM and DS sapphire-NiAl(Yb) composites.

II. EXPERIMENTAL PROCEDURE

The composite feedstock specimens (fiber volume fraction: 28 vol pct) were fabricated by the PC process using powders of either near-stoichiometric, single-phase NiAl, or an aluminum-rich NiAl(Yb), and
combining these powders with single-crystal sapphire fibers (Saphikon, Milford, NH) of about 130-μm diameter. In the PC process, cloths of matrix alloy, formed by mixing powders combined with organic binders, are placed between plies made up of rows of alumina fibers and are diffusion bonded by application of high pressures and high temperatures. The PM composite plates (0.25 cm by 0.4-cm cross section) were subsequently used as feed material for HT and zone DS using a 30-kW RF generator. For the present set of experiments, solid-state HT and DS were carried out by creating a hot zone (molten, in the case of DS) of 0.6- to 0.8-cm length in the feedstock bar, using a copper concentrator (1.25-cm-diameter hole) and traversing this zone at a speed of 6.0 cm per hour along the length of the feed specimen. A positive 20 kPa pressure of ultra-high-purity argon was used to suppress the loss of aluminum because of its high vapor pressure. A radiation pyrometer was used to control the temperature of the zone to within ±6 K of the desired temperature.

The fiber-matrix interfacial shear strengths of PM, HT, and DS composites were measured using a desktop fiber-pushout-test apparatus. This device records the time dependence of load during pushing (compressing) of a single sapphire fiber embedded in the NiAl matrix by using a flat-bottomed punch of sintered tungsten carbide (102-μm diameter). The samples for pushout testing were in the form of flat discs (nominal thickness: 300 to 400 μm) with a metallographic surface finish. The disks were firmly glued to a steel sample holder having 400-μm-wide grooves, taking care to align the fibers over the grooves so as to enable their unobstructed sliding during the pushout test. A motorized traverse mechanism allowed the sample to be moved toward the stationary punch at a rate of 0.82 μm s⁻¹. The load vs time data were recorded at 50-ms intervals for a period of 180 seconds or less and stored in computer files. In addition, acoustic emission (AE) was also recorded during the fiber-pushout process. Generally, 10 to 30 tests were performed for each processing condition.

The grain structure in the matrix was revealed by etching metallographically polished samples with a solution of 25 vol pct acetic acid, 25 vol pct nitric acid, and 50 vol pct distilled water. The debonded fiber and matrix surfaces from the pushout-test specimens were examined in a Cambridge 2000 scanning electron microscope (SEM). The sapphire fibers were extracted from the PM, HT, and DS composites for SEM examination by dissolving the matrix in a solution of 75 mL acetic acid (glacial), 25 mL concentrated HNO₃ acid, and 2 mL concentrated HCl. The fracture (debonding) behavior during fiber pushout was examined by interrupting the pushout test and examining the front faces (the sample surface in contact with the tungsten carbide punch) and back faces of the samples in the vicinity of the fibers by SEM. Energy dispersive X-ray spectroscopy (EDS) and X-ray diffraction were used to identify the phases present.

### III. RESULTS

#### A. Chemical Analyses and X-ray Diffraction of Composite Matrices

The chemical compositions of the PC processed PM, HT, and DS NiAl(Yb) matrices (after dissolving the matrix and removing the fibers), as determined by inductively coupled emission spectroscopy, are listed in Table I. The nickel and aluminum contents of the matrices did not change because of HT and DS; however, there was nearly a 50 to 70 pct decrease in the Yb content of the matrix in the DS and HT composites.

The results of X-ray analysis summarized in Table I show that both the PM and DS NiAl composites contained near-stoichiometric NiAl (with a lattice parameter of 0.2878 nm for PM and 0.2886 nm for DS material) and α-Al₂O₃. Also, the PM NiAl(Yb) composites contained NiAl, α-Al₂O₃, Yb₂O₃, and Yb₃Al₅O₁₂ phases. The lattice parameter of the NiAl(Yb) lattice, determined from the (100) and (110) reflections, was 0.2867 nm. In the case of HT NiAl(Yb) composites, the phases present were NiAl, α-Al₂O₃, NiAl martensite, and the spinel oxide Yb₂Al₅O₁₂, and the DS NiAl(Yb) composites contained NiAl, α-Al₂O₃, Yb₂Al₅O₁₂, and Ni₃Al phases. The NiAl(Yb) lattice parameters for the HT and DS NiAl(Yb) composites were 0.2861 and 0.2856 nm, respectively.

#### B. Microstructure

Figures 1(a) through (e) show typical transverse microstructures (normal to the fibers) of the sapphire fiber-reinforced composites. Each figure contains a low-magnification overall view that shows the distribution of fibers and also a high-magnification view that shows the matrix microstructure in the vicinity of the sapphire fibers. The average grain sizes of the PM NiAl and PM NiAl(Yb) specimens (measured on the transverse sections) were 50 and 80 μm, respectively. In comparison, the grain sizes of the DS NiAl and NiAl(Yb) specimens were 1000 and 230 μm, respectively. The HT-NiAl(Yb) specimens had an intermediate grain size of about 135 μm. Because of the smaller matrix grain size in the PM material compared to the HT and the DS material, the number of grain boundaries coincident on the fiber surface in PM material was much greater than in the HT and the DS specimens. There was occasional micro-porosity located at some of the grain boundary junctions in the matrices of PM specimens, marked by arrows in Figures 1(a) and (b). The DS (Figure 1(d)) and the HT (Figure 1(e)) sapphire-NiAl(Yb) specimens show micro-porosity and some matrix cracking.

The light phase along the grain boundaries in the PM, HT, and DS NiAl(Yb) specimens (marked by an arrow in Figures 1(b), (d), and (e)) was identified by EDS as Ni₃Al (the Ni₃Al phase was not detected in the PM-NiAl(Yb) by X-ray analysis described previously). Although the lenticular markings in the DS and HT NiAl(Yb) matrix (Figures 1(d) and (e)) strongly suggest the presence of martensite (which is expected to form in NiAl alloys with less than 37 at. pct Al¹²), X-ray analysis could not confirm its presence. Optical microscopy also revealed a severe chemical reaction between the fiber and matrix in the PM (Figure 1(b)), DS (Figure 1(d)), and HT (Figure 1(e)) sapphire-NiAl(Yb) specimens. In contrast, no fiber-matrix reaction was observed in the PM (Figure 1(a)) and DS (Figure 1(c)) sapphire-NiAl composites.
Table I. Summary of Chemical and X-Ray Phase Analysis of the PM and DS Sapphire-Reinforced NiAl(Yb) Composites*

<table>
<thead>
<tr>
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<tbody>
<tr>
<td>PM-NiAl</td>
<td>49.9</td>
<td>50.1</td>
<td>—</td>
<td>NiAl and Al₂O₃</td>
</tr>
<tr>
<td>DS-NiAl</td>
<td>50.3</td>
<td>49.7</td>
<td>—</td>
<td>NiAl and Al₂O₃</td>
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<tr>
<td>PM-NiAl(Yb)</td>
<td>62.1</td>
<td>36.8</td>
<td>0.36</td>
<td>NiAl, Al₂O₃, Yb₂O₃, and Yb₃Al₂O₆</td>
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<td>HT-NiAl(Yb)</td>
<td>62.6</td>
<td>37.3</td>
<td>0.11</td>
<td>NiAl, Al₂O₃, Yb₂Al₂O₇, and NiAl martensite</td>
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<tr>
<td>DS-NiAl(Yb)</td>
<td>62.5</td>
<td>36.4</td>
<td>0.19</td>
<td>NiAl, Al₂O₃, Yb₂Al₂O₇, and Ni₃Al</td>
</tr>
</tbody>
</table>

*The chemical compositions were determined to 5 pet relative accuracy.

C. Interfacial Reaction

Figures 2(a) through (c) are back-scattered SEM images of the fiber-matrix interface in the PM, DS, and HT NiAl(Yb) composites, respectively. The sapphire fiber appears dark and the reaction-product layer appears white (or gray). It is apparent that the extent of chemical reaction is much greater in the HT and DS specimens when compared to the PM-NiAl(Yb) specimen. The various phases present in the matrix near the interface are labeled A, B, C, D, and E in the higher-magnification views shown in the inset for the PM, HT, and DS NiAl(Yb) composites. The corresponding EDS spectra of these five phases from the PM NiAl(Yb) specimen are presented in Figure 3. Phase A is present as a continuous layer surrounding the fiber (Figure 2(a)) and contains Ni, Al, and O (Figure 3(a)). Phase B, which is also present as a continuous layer beyond the A phase into the matrix (Figure 2(b)), comprises Yb, Al, and O (Figure 3(b)). Phase B also appears as "pegs" growing out from the sapphire surface into the intergranular regions of the NiAl matrix. Phase C is located within phase B (Figure 2(a)) and contains only Yb and O (Figure 3(c)). Phase D, which is the matrix B phase (Figures 2(a) and (b)) contains only Ni and Al (Figure 3(d)). A comparison of Figures 3(a) and (d) shows that phase A is richer in aluminum than the matrix phase D. In addition, EDS analysis showed that the nickel-to-aluminum peak-intensity ratio was >1 at the matrix grain boundaries, whereas the NiAl matrix (phase D) had a nickel-to-aluminum ratio of ≈1. Finally, the E phase, which was dispersed throughout the matrix and seen as bright white spots in Figure 2(a), contained Yb and F (Figure 3(e)).

Typical SEM views of fibers extracted from the PM and DS NiAl composites are shown in Figures 4(a) and 4(b) and fibers extracted from the PM, DS, and HT NiAl(Yb) composites are presented in Figures 4(c) through (e). Also shown is an SEM view of an as-received sapphire fiber (Figure 4(f)). These figures confirm the optical microscopy observations that fibers in the PM, HT, and DS NiAl(Yb) composites have undergone severe chemical attack by the matrix, resulting in extensive...
roughening of the fiber. In contrast, the as-received fibers (Figure 4(f)) generally had smooth surfaces, except for occasional asperities, as indicated in Figure 4(f). The surfaces of fibers extracted from the PM NiAl composites (Figure 4(a)) showed evidence of impressions from prior particle boundaries that formed during the powder cloth processing. On the other hand, the fibers extracted from the DS NiAl composites (Figure 4(b)) showed circular features that presumably formed from the gas porosity created during the zone DS process, as shown in
Fig. 2—Microstructures at the matrix-fiber interface in (a) PM NiAl(Yb), (b) DS NiAl(Yb), and (c) HT NiAl(Yb). Regions marked A through E denote same phases in PM, HT, and DS composites.
Figure 2(a). However, unlike the PM and DS NiAl(Yb) composites, no evidence of chemical attack of fibers was observed in the PM and DS NiAl composites.

D. Fiber-Pushout Behavior

Figure 5 shows typical shear stress-displacement plots and the accompanying AE from the fiber-pushout tests on PM and DS NiAl, and PM, DS, and HT NiAl(Yb) composite samples. Pushout stress values were obtained by dividing the pushout load values by the interfacial contact area, \( \pi D t \), where \( D \) is the fiber diameter and \( t \) is the disk thickness. Note that the disk thicknesses (300 to 400 \( \mu m \)) are much larger than the total fiber displacement during the pushout process. In general, a three-step stress-displacement response was observed in these composite materials. The three stages of failure process were as follows: an initial linear increase in stress (pseudelastic region) followed by a nonlinear increase in stress (inelastic region), up to a maximum, whereafter the stress dropped either abruptly, as in the case of NiAl composites (Figures 5(a) and (b)), or gradually, as in the case...
of NiAl(Yb) composites (Figures 5(e) through (e)), to some constant value. A gradual stress-drop-type behavior was also observed in a recent study on interfacial shear strength in the sapphire-Ti-48Al-2Cr-2Nb system, in which a severe interfacial reaction resulted in a very rough fiber surface.

The pushout behavior will be described in terms of the proportional limit ($\tau_p$), corresponding to the stress at the transition point between the pseudoeastic and inelastic regions, maximum shear stress ($\tau_m$), corresponding to the maximum in the stress-displacement curves, and frictional shear stress ($\tau_f$), obtained at a displacement of 8 µm (10 seconds) beyond the stress drop. Table II compiles the average $\tau_p$, $\tau_m$, and $\tau_f$ values and their standard deviations for the composite specimens examined in this study. For some of the fibers in the PM NiAl(Yb) specimens, the maximum load available in the equipment (40 N) was insufficient to push the fiber out. For these
Fig. 5—Fiber-pushout response of the sapphire-fiber-reinforced NiAl and NiAl(Yb) composites: (a) PM NiAl, (b) DS NiAl, (c) PM NiAl(Yb), (d) DS NiAl(Yb), and (e) HT NiAl(Yb).
tests, the $\tau_m$ and $\gamma_l$ values could not be obtained. The number of such tests is indicated in Table II in parentheses, and the “>” sign has been used to indicate that the actual mean is probably higher than the values given in the table, which include only the pushed-out fibers. During pushout, the first distinct AE signal was generally detected near $\tau_p$. The ratios between the intensity of this AE signal and the background are also compiled in Table II. As indicated in Table II, and typically shown in Figure 5, the AE signal-to-noise ratio in the vicinity of $\tau_p$ was significantly larger in the PM and HT NiAl(Yb) specimens compared to the DS specimens. By comparison, both PM and DS NiAl specimens showed significantly weaker AE peaks.

Because the shear stresses at the onset of inelastic deformation (proportional limit $\tau_p$) represent the initial debond stress\(^{14}\) at which tensile cracks are nucleated at the sample backface, Table II shows that higher stresses were needed to nucleate cracks in the DS NiAl specimens ($\tau_p = 84 \pm 40$ MPa) as compared with the PM NiAl specimens ($\tau_p = 27 \pm 13$ MPa). In contrast, the NiAl(Yb) composites showed an opposite trend; the $\tau_p$ values were significantly higher in PM NiAl(Yb) ($177 \pm 64$ MPa) and HT NiAl(Yb) ($118 \pm 72$ MPa) as compared to the DS NiAl(Yb) ($47 \pm 28$ MPa). The maximum stress $\tau_m$ represents the final debond stress at which complete fiber-matrix debonding occurred.\(^{14}\) The $\tau_p$ values also show a pattern similar to the proportional limit stress $\tau_p$ for both materials. Considering only the pushed fibers, the average $\tau_p$ values for PM, HT, and DS NiAl(Yb) specimens were $205 \pm 69$, $164 \pm 64$, and $67 \pm 44$ MPa, respectively, compared to the average $\tau_m$ values for PM and DS NiAl specimens, which were $30 \pm 16$ and $138 \pm 62$ MPa, respectively.

The frictional stress $\gamma_l$ for PM and DS NiAl was nearly the same $20 \pm 13$ and $27 \pm 7$ MPa, respectively. In contrast, the $\gamma_l$ values for the PM NiAl(Yb) ($126 \pm 16$ MPa) and HT NiAl(Yb) ($135 \pm 47$ MPa) were considerably larger than the $\gamma_l$ for DS NiAl(Yb) ($53 \pm 41$ MPa). The frictional stresses for both the PM and DS NiAl(Yb) specimens were larger than those for the NiAl specimens.

The proportional shear stress $\tau_p$ corresponds to the initiation of the interfacial debond as a result of combined mechanical-loading and thermal-clamping stresses due to coefficient of thermal expansion (CTE) mismatch between the fiber and the matrix. The maximum shear stress $\tau_m$ corresponds to the completion of the debonding process. Thus, the stress range from $\tau_p$ to $\tau_m$ is related to the propagation of debond. The debond progression itself is a complex process and involves the growth of pre-existing interfacial microcracks, cracking of matrix grain boundaries that are coincident on the fiber surface, and interfacial and matrix microplasticity. The frictional shear stress $\gamma_l$ represents the “sliding stress” due to the combined influence of coulombic friction, asperity (roughness)-controlled friction, and the thermal-clamping stresses at asperity contact points during fiber sliding. The interpretation of test results to obtain bond strength is, therefore, not straightforward. Further complications may be introduced by the fragmentation of fiber into segments of unknown lengths during fiber pushout. Therefore, while the various shear stresses ($\tau_p$, $\tau_m$, and $\gamma_l$) obtained from the fiber-pushout test are measures of the strength of the interfacial bond between the fiber and the matrix, the precise relationship of these stresses to the “true” or “intrinsic” bond strength in a multiphase reactive system is difficult to establish.

### Table II. Fiber-Pushout Characteristics of Sapphire-Fiber-Reinforced NiAl and NiAl(Yb) Composites

<table>
<thead>
<tr>
<th>Material</th>
<th>Grain Size ($\mu$m)</th>
<th>AE Ratio</th>
<th>$\tau_p$ (MPa)</th>
<th>$\tau_m$ (MPa)</th>
<th>$\gamma_l$ (MPa)</th>
</tr>
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<tbody>
<tr>
<td>PM-NiAl</td>
<td>50</td>
<td>10 ± 4</td>
<td>27 ± 13</td>
<td>30 ± 16</td>
<td>20 ± 13</td>
</tr>
<tr>
<td>DS-NiAl</td>
<td>1000</td>
<td>8 ± 4</td>
<td>84 ± 40</td>
<td>138 ± 62</td>
<td>27 ± 7</td>
</tr>
<tr>
<td>PM-NiAl(Yb)</td>
<td>80</td>
<td>114 ± 81</td>
<td>177 ± 64</td>
<td>&gt;205 ± 69(24)</td>
<td>126 ± 16</td>
</tr>
<tr>
<td>HT-NiAl(Yb)</td>
<td>135</td>
<td></td>
<td>118 ± 72</td>
<td>164 ± 64</td>
<td>135 ± 47</td>
</tr>
<tr>
<td>DS-NiAl(Yb)</td>
<td>128 ± 77</td>
<td>13 ± 9</td>
<td>47 ± 28</td>
<td>67 ± 44</td>
<td>53 ± 41</td>
</tr>
</tbody>
</table>

### E. Fracture Behavior

Unlike the sapphire-NiAl composites in which the fracture behavior during the pushout test was different for the PM and the DS specimens,\(^{16}\) no differences were observed in the fracture behavior of PM, HT, and DS NiAl(Yb) specimens. Figure 6 shows typical SEM views of the debonded surfaces for PM and DS NiAl(Yb) specimens after fiber pushout beyond $\tau_m$. They both showed backface cracking, similar to the PM and DS NiAl specimens reported earlier.\(^{17}\) The front faces were generally crack-free, although occasional cracks were observed at the interfacial regions on the front face. Both the fiber and matrix in the fractured regions showed considerable surface heterogeneity, which resulted from the chemical interactions. The debonded matrix surface (Figure 6(c)) showed fine wear tracks parallel to the fiber-sliding direction. Because the asperities generated on the fiber surface resulting from the chemical reaction between the fiber and the matrix in NiAl(Yb) composites were much coarser than the growth-related surface defects on the as-received fibers, the plastic grooving of the matrix and the formation of wear tracks were significantly greater in the PM, HT, and DS NiAl(Yb) (Figure 6(c)) compared to the DS NiAl, where fiber-surface roughening resulting from chemical reactions was absent.\(^{17}\)

The fact that no differences in the failure modes could be detected from SEM examination of tested specimens of PM, HT, and DS NiAl(Yb) materials indicates that the failure processes in these materials are far more complex than those in the NiAl material. It is conceivable that matrix grain boundaries coincident on the fiber surface, the different extents of interfacial chemical reactions, somewhat different reaction products, and the
Fig. 6—Fracture behavior of the sapphire-fiber-reinforced NiAl(Yb) composites: (a) front face and (b) back face. (c) Wear tracks resulting from microplastic grooving from fiber asperities during fiber sliding.

IV. DISCUSSION

An examination of the Ni-Al phase diagram (Figure 7(a)) shows that the NiAl composites (Ni-50 at. pct Al) had a single β-phase matrix. The NiAl(Yb) composites (Ni-37 at. pct Al) are, however, expected to consist of NiAl (β) and Ni3Al (γ′) phases (Figure 7(a)). As presented in Table I, γ′ was detected only in the HT and DS NiAl(Yb) by X-ray diffraction and not in PM NiAl(Yb), although optical microscopy indicated γ′ in the PM NiAl(Yb), which was confirmed by the EDS analysis of the light phase at the grain boundaries (Figure 2(a)). Because this phase showed only nickel and aluminum, and its nickel-to-aluminum ratio was higher than that in the NiAl matrix, it can be assumed that this phase is Ni3Al. Also, X-ray analysis did not show the presence of martensite in the DS NiAl(Yb), although its presence was revealed by metallography (Figure 1(d)).

A. Interfacial Microstructure

The following sequence of reactions can be hypothesized for the formation of the interfacial microstructures (Figures 2(a) through (c)) in the PM, DS, and HT NiAl(Yb) composites. Ytterbium is a chemically active rare-earth element, as well as a strong oxide former. The standard free energies of formation of Y2O3 and Al2O3 are \(-1727.50 \text{ kJ} \cdot \text{mol}^{-1}\) and \(-1583.10 \text{ kJ} \cdot \text{mol}^{-1}\), respectively. In the PM NiAl(Yb) composite, the reaction between ytterbium in the matrix and Al2O3 fibers leads to the dissolution of the fiber and to aluminum and oxygen enrichment of the surrounding NiAl ("A" in Figure 2(a)). The diffusion of oxygen into the matrix (away from the sapphire fibers) and of ytterbium toward the fibers leads to the formation of a Yb2O3 layer around the sapphire fiber ("C" in Figure 2(a)). Because of the oxygen enrichment of the original powder surface, the Yb2O3 phase is also formed on the grain boundaries (prior powder boundaries) throughout the matrix. The Yb2O3, located on the matrix grain boundaries which were in
contact with the sapphire fibers, initiates the formation of the pegs described previously. The Yb₂O₃ is subsequently consumed because of a reaction with Al₂O₃, leading to the formation of the Yb₃Al₅O₁₂, which is essentially a spinel oxide (Yb₂O₃ to Al₂O₃ ratio of 3:5) (Figure 7(b)). Elevated-temperature exposure of PM NiAl(Yb) composite, in the HT NiAl(Yb) specimens, for example, has been observed to result in an almost complete absence of Yb₂O₃ in the vicinity of the fiber and its conversion into Yb₃Al₅O₁₂ (Figure 2(c)). The presence of fluorine in the TEFLON* binder, used during the PC processing of composites, accounts for the ytterbium- and fluorine-rich dispersoids observed in the NiAl matrix.

The thermal profile imposed on the composite specimens during zone DS leads to an initial solid-state heating of the specimen before the formation of the melt zone. Apparently, the NiAl layer surrounding the sapphire fibers and also the Yb₂O₃ layer in the PM composite are completely consumed during this high-temperature exposure, leading to the formation of a thick Yb₃Al₅O₁₂ layer that surrounds the fibers and the Yb₃Al₅O₁₂ pegs protruding from the fiber surface. Subsequently, during the DS, the NiAl alloy melts around these Yb₃Al₅O₁₂ layers and goes through the process of solidification and cooling to the room temperature. Solidification shrinkages result, especially in the crevices in the vicinity of these pegs (marked by arrows in Figure 2(b)) because of an insufficient metallocstatic head. The pegs also contribute to the shrinkage of the matrix during cooling to room temperature in both HT and DS specimens, and the stresses thus generated cause severe cracking of the NiAl matrix, as observed in the DS and HT NiAl(Yb) composites (Figures 1(d) and (e)). The reaction sequence previously proposed for the formation of various reaction products is, however, conjectural but not implausible. Detailed research of the reaction mechanisms and reaction paths was not attempted in the present study.

In summary, the following microstructural features are of interest in the NiAl composites. (1) There was minimal degradation of the sapphire fiber surface, and (2) the number of grain boundaries in contact with the fiber in PM composites was much greater than in the HT and DS composites. In the PM NiAl(Yb) composite, the fiber was surrounded mostly by the oxygen-rich NiAl layer, except for the Yb₃Al₅O₁₂ phase pegs, which grew from the fiber surface into the intergranular regions of the matrix. In the HT and DS NiAl(Yb) composite, the fiber was not in contact with NiAl; instead, it was surrounded by a layer of Yb₃Al₅O₁₂ phase, about 5-μm thick in the DS NiAl(Yb) and <0.5-μm thick in the HT NiAl(Yb) composite.

B. Fiber Pushout Behavior

Recent finite element modeling studies[17] of fiber-pushout behavior in a "thin-slice" test configuration suggest that thermal stresses (due to CTE mismatch) and mechanical stresses (due to loading) can generate radial stresses large enough to nucleate fine cracks at the sample back face that propagate along the interface and lead to interface failure. Interrupted pushout test and fractography of sapphire-NiAl composites[17] failed to detect any front- or back-face cracks on the test specimens when the test was interrupted at τp. However, a high AE signal-to-noise ratio was generally observed in the vicinity of τp (Table II), suggesting that some micromechanical events, possibly too weak to leave gross microstructural manifestation, do occur at the fiber-matrix interface, at τp. On the other hand, when the test was interrupted at τ > τp, both the front (Figure 8(a)) and back faces (Figure 8(b)) of PM-NiAl specimens showed[17] extensive interfacial cracking along grain boundaries that were coincident on the fiber. The fibers in the DS NiAl showed both grain boundary cracking and circumferential decohesion along part of the fiber-matrix interface on the front (Figure 8(c)) and back (Figure 8(d)) faces of the samples. For the fibers completely embedded within
the NiAl grains in the DS specimens, the front faces were free of interfacial damage (Figure 8(e)), but the back faces (Figure 8(f)) showed partial fiber-matrix debonding together with fine radial cracks; these fibers displayed the highest interfacial shear strength. Thus, radial grain boundaries in contact with the fiber surface are not only the preferred site for nucleation of these cracks but also provide an easy path for crack growth. The sapphire fibers in the PM NiAl, because of a small matrix grain size, had the maximum number of such grain boundaries, resulting in easy nucleation of such cracks and the lowest $\tau_p$ values. The fibers in the DS materials, with a fewer number of such grain boundaries, showed larger $\tau_p$ values.\(^1\) Also, in both PM and DS sapphire-NiAl composites, localized microplastic grooving of the matrix by fiber asperities was observed (Figure 8(g)), in a manner similar to the present sapphire-NiAl(Yb) system (Figure 6(c)).

These results show that the deformation processes that occur during fiber pushout are influenced by the nature of the fiber-matrix interface and the matrix microstructure, both of which are influenced by processing. In the absence of an interfacial chemical reaction (as in PM and DS NiAl), the progressive elimination of matrix grain boundaries by DS enhanced the fiber-matrix interfacial shear strength.\(^1\) However, when an extensive chemical reaction between the fiber and the matrix was induced by alloying NiAl with Yb, the influence of interfacial reactions dominated the matrix microstructure in the vicinity of the fiber in determining the interfacial shear strength.

As described in Section III, only the DS and the HT
NiAl(Yb) test specimens showed matrix cracking; the PM material did not show cracking, although it had numerous grain boundaries coincident on the fiber-matrix interface in a manner similar to the PM NiAl specimens. However, unlike the grain boundaries in the PM NiAl specimens, the grain boundaries in the NiAl(Yb) matrix showed presence of the hard and brittle intermetallic Ni$_3$Al (Figures 1(b), (d), and (e)). The role of grain boundaries in the failure processes associated with fiber debonding during fiber pushout is, therefore, expected to be different in the NiAl(Yb) and NiAl materials. In particular, the grain boundaries coincident on the fiber surface in the NiAl(Yb) are not expected to serve as sites favorable for crack nucleation and growth, as they did in the NiAl material, because of the presence of the Ni$_3$Al phase at grain boundaries. Thus, despite the presence of numerous grain boundaries in both PM NiAl and PM NiAl(Yb) materials, the latter material, by comparison, showed significantly greater $\tau_p$ and $\tau_m$ values, as well as AE ratios (Table II), because the strengthening effects of chemical bonding were not offset by the weakening effects of grain boundaries as a result of the presence of Ni$_3$Al phase at grain boundaries. On the other hand, the cracks formed during processing of HT and DS NiAl(Yb) materials did not have the Ni$_3$Al phase present. Therefore, such cracks are expected to overcome the advantage of enhanced chemical bonding in the NiAl(Yb) material by assisting the debonding processes during fiber pushout. This is revealed most dramatically when the $\tau_p$ values and the AE ratios of PM NiAl(Yb) (no cracking) materials and DS NiAl(Yb) (extensive cracking) materials are compared (Table II).

In contrast to the NiAl, where DS resulted in improved $\tau_p$ because of the elimination of transverse grain boundaries, the DS NiAl(Yb) specimens showed a reduction in $\tau_p$ and $\tau_m$ as compared with the PM NiAl(Yb). Once a chemical bond is established between the fiber and the matrix in the PM NiAl(Yb) composite, the increase in the extent of chemical attack of the fiber by the matrix during the HT and zone DS is not expected to provide any additional interfacial strengthening. Consequently, no change in the $\tau_p$ and $\tau_m$ values would be expected between the PM, HT, and DS NiAl(Yb) composites. The fact that the DS composites had, in fact,
lower interfacial shear strengths compared to the PM specimens can be attributed to the extensive interfacial cracking between Yb₃Al₂O₉ and NiAl matrix and also to the presence of shrinkage cavities located there in the DS samples (marked by arrows in Figure 2(b)).

As the onset of inelastic instability at τ₂ indicates initial debonding, a higher τ₂ value is expected to be associated with a stronger AE signal. This is indeed the case, as noted from Table II. Of all the composite specimens examined in this study, the PM and HT NiAl(Yb) specimens (with the highest τ₂ values) showed largest values of the signal-to-noise ratios. The DS NiAl(Yb) composite showed a lower AE signal-to-noise ratio because of the relatively smaller energy released at the onset of debonding. On the other hand, the relatively smooth fiber-matrix interfaces of the PM and DS NiAl composites, as compared with the NiAl(Yb) composites, yielded the lowest AE signal-to-noise ratios, as seen from Table II.

It is well known that a certain amount of chemical interaction between the fiber and the matrix is essential for improving the strength of the interfacial bond; however, the extent of this interaction must be carefully controlled to achieve an optimum bond. An excessive chemical attack of the fiber by the matrix degrades the fiber strength because of surface "notches," which cause premature fiber failure without yielding any load-bearing advantage of the fiber. In the PM NiAl(Yb) composites, the existence of a chemical bond as opposed to a purely "frictional" bond suggests that the interfacial shear strength should be high. This is indeed the case and is reflected in the high values of τ₂ and τ₃ for this composite. However, as a result of the extreme reactivity of the sapphire-NiAl(Yb) system, the advantage of improved chemical bonding is realized at the expense of the fiber strength and surface quality. As the focus in the present work was the strength of the interface rather than that of the bulk composite, the highly reactive sapphire-NiAl(Yb) model system was chosen. As a cautionary note, we reiterate that no strengthening of the bulk composite is to be expected by reinforcing the NiAl(Yb) matrix by sapphire fibers because of the extreme degradation of the latter during composite fabrication.

Once the fibers are completely debonded at τ₃, the frictional sliding of the fiber in the matrix is affected mainly by the fiber-surface roughness. Therefore, the friction stress in the PM and DS NiAl composites would be expected to be about the same, as is observed (Table II), because of the relatively smooth fiber-matrix interface in the NiAl composites. On the other hand, the rougher fiber surface and the presence of pegs in the NiAl(Yb) composites are responsible for much larger friction stresses observed in the NiAl(Yb) composites.

V. CONCLUSIONS

The following conclusions can be drawn from this study on the deformation and fracture behavior during pushout testing of sapphire fibers in sapphire-fiber-reinforced Powder Cloth processed, heat treated, and directionally solidified NiAl(Yb) composites.

1. The addition of ytterbium causes extensive reaction between the matrix and sapphire fibers. In the PM NiAl(Yb) composite, the fiber is surrounded mostly by the oxygen-rich NiAl layer, except for the Yb₃Al₂O₉ phase pegs, which grow from the fiber surface into the intergranular regions of the matrix. In the HT and the DS NiAl(Yb) composites, the fiber is not in contact with NiAl but, instead, is surrounded by a layer of Yb₃Al₂O₉ phase.
2. The stress vs displacement response during fiber pushout consists of three regimes: (a) an initial region of constant slope up to a stress τ₄, followed by a region of decreasing slope up to a maximum stress τ₅, and (c) a subsequent gradual stress drop and onset of frictional sliding at a stress τ₆. The occurrence of a strong AE signal near τ₄ indicates that this stress corresponds to the onset of fiber-matrix debond. Complete debonding occurred at τ₅ and large-scale, postdebond fiber sliding occurred at a progressively decreasing friction stress τ₆.
3. The PM and HT NiAl(Yb) specimens showed higher τ₅ and τ₆ values because of the formation of a chemical bond between the fiber and the matrix, as opposed to the sapphire-NiAl composite in which the bond is purely frictional in nature. These stress values are, however, lower in the DS NiAl(Yb) composite because of the microcracks and shrinkage cavities at the Yb₃Al₂O₉-matrix interface. The frictional stress for postdebond fiber sliding in the sapphire-NiAl(Yb) composites was significantly higher than that in the sapphire-NiAl composites.

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