Multi-Scale Modeling of Metal-Composite Interfaces in Titanium-Graphite Fiber Metal Laminates Part I: Molecular Scale

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Received August 12, 2011; revised September 16, 2011; accepted September 25, 2011

Abstract

This study presents the first stage of a multi-scale numerical framework designed to predict the non-linear constitutive behavior of metal-composite interfaces in titanium-graphite fiber metal laminates. Scanning electron microscopy and x-ray diffraction techniques are used to characterize the baseline physical and chemical state of the interface. The physics of adhesion between the metal and polymer matrix composite components are then evaluated on the atomistic scale using molecular dynamics simulations. Interfacial mechanical properties are subsequently derived from these simulations using classical mechanics and thermodynamics. These molecular-level property predictions are used in a companion study to parameterize a continuum-level finite element model of the interface by means of a traction-separation constitutive law. Extension of the proposed approach to other dissimilar metal- or metal oxide-polymer interfaces is also discussed.

Keywords: Fiber Metal Laminates (FML), Titanium-Graphite (TiGr), Molecular Dynamics (MD), Adhesion

1. Introduction

Fiber metal laminates (FMLs) are a unique class of structural materials that combine traditional fiber-reinforced polymer matrix composite (PMC) laminae with thin, adhesively bonded metal layers. In these FML systems, the metal reinforcement layer(s) can be added to the PMC structure either internally or externally, provided that the fiber-reinforced polymer and metal layers are bonded together during the composite consolidation process. Consequently, the resultant structure is globally homogeneous but locally heterogeneous, with distinct material phases evident by visual inspection, Figure 1. First developed at Delft University of Technology in the late 1970s, commercially available examples of fiber metal laminates include GLARE (glass fiber-reinforced aluminum laminates) and ARALL (aramid fiber-reinforced aluminum laminates) [1]. Recently, a novel FML variant has been developed combining carbon fiber-reinforced polymer matrix composites with adhesively bonded titanium layers [2]. These FMLs are the focus of the present study and are commonly referred to as titanium-graphite (TiGr) or hybrid titanium composite laminates (HTCL). TiGr FMLs represent an intriguing material system as they possess increased stiffness and strength in comparison to ARALL and GLARE. They can also be utilized in elevated temperature applications, provided that the polymer matrix material has sufficient thermal degradation resistance.

The primary advantage of all fiber metal laminate variants is their ‘best of both worlds’ design, in which the beneficial properties of the individual metal and PMC constituents are combined into a single structure with customizable material properties [3]. In essence, fiber metal laminates utilize the synergistic enhancement of structural properties achieved by combining two ma-
terials (metals and PMCs) with counterbalanced strengths and weaknesses. Because the limitations of each constituent are effectively offset, FMLs can be used in applications which are typically considered ill-suited for either metals or polymer matrix composites, such as impact-critical or mechanically fastened structures [4-9]. For each of these scenarios, it is important to note that the amount of improvement offered by an FML over a traditional metal or composite structure is heavily dependent upon the state of the interface between the metal and PMC constituents [10]. For instance, in impact-critical structures the formation and propagation of a delamination zone between the metal and PMC layers reduces the overall damage tolerance and impact energy dissipation potential of the FML. In addition to this loss of damage tolerance, interfacial delamination resulting from impact has also shown to significantly reduce the residual static and fatigue strength of the FML [11]. For mechanically fastened fiber metal laminate structures, interfacial delamination prevents the metal reinforcement layers from alleviating the high stress concentration in the region surrounding the fastener. As such, the resulting delaminated FML is highly notch sensitive and will fail at lower than expected joint loads [6]. It is, therefore, highly desirable to form and maintain a strong interlaminar bond between the polymer matrix adhesive and the metal adhered in any FML system so that these components do not delaminate.

The importance of the metal-composite interface with respect to the structural performance of any FML variant underscores the evident need for accurate predictive models of interfacial mechanical behavior. However, it should be noted that prediction of adhesion between any two dissimilar components at a common interface is one of the most challenging and complex problems in the realm of materials modeling. The inherent difficulty is primarily due to the multiple length scales and damage modes associated with interfacial debonding and failure. For FMLs, this complexity means that any metal-composite interfacial model must be multi-scale. It must incorporate molecular-level considerations arising from separation of the interface, as well as continuum-level damage and failure phenomena due to microcracking and fracture of the polymer matrix. The present analysis is concerned with the development of one such multi-scale interface model, specifically applied and validated for use with TiGr FMLs. Establishment of the molecular-level simulation methodology and physical/chemical characterization of the metal-composite interface are described herein. The coupled continuum-level modeling approach and experimental validation of the multi-scale framework will be covered in detail in a companion study.

2. Tigr Processing and Interfacial Characterization

2.1. Titanium Surface Treatment

In the fabrication of any fiber metal laminate system, formation of a strong interlaminar bond between the metal and polymer matrix composite layers of the FML is of the utmost importance. In the specific case of titanium-graphite FMLs, the metal-composite interface must also be capable of withstanding the large residual thermal stress state that is introduced during the FML con-
solidation process. These residual stresses are caused by a sizable mismatch in the thermal expansion coefficients of adjacent titanium and PMC layers. In the as-received state, the interlaminar bond strength between the unmodified titanium and the polymer matrix is typically not sufficient to endure these residual thermal stresses, resulting in delamination after completion of the cure cycle [12,13]. Thus, in order to increase the strength and reliability of the TiGr metal-composite interface, a recently proposed chromium-free sequential acid-base titanium surface treatment procedure was employed [14]. It should be noted that the surface treatment process variables were slightly modified to adapt the process to the un-alloyed commercially pure (CP) titanium used in this study.

The ability of this chromium-free chemical treatment process to physically alter the CP titanium surface is evident from comparison of Figure 2 and Figure 3, which show scanning electron microscope (SEM) images obtained using a JEOL 6700F field emission SEM for pre- and post-treated CP titanium foil. In Figure 2, the non-uniform preexisting surface oxide layer is preferentially aligned with the rolling direction of the foil, and only provides partial surface coverage. After completion of the surface treatment process, this preferential orientation disappears and full surface coverage is achieved, as shown in Figure 3. Coverage of the CP titanium surface with a protective titanium dioxide (TiO\textsubscript{2}) layer is the primary basis for enhanced interlaminar bond strength obtained using this sequential acid-base surface treatment method [14]. Additionally, comparison of the as-received and surface treated titanium foil surfaces shown respectively in Figure 2 and Figure 3 indicates that the surface treated titanium possesses a much higher degree of surface roughness in comparison the as-received titanium. This increased surface roughness also contributes to the interlaminar bond strength improvement, since it represents a larger active surface area between the polymer matrix adhesive and the titanium adhered.

2.2. Characterization of the Titanium Dioxide Surface Layer

In the preceding discussion, it was assumed a priori that changes in the CP titanium surface morphology, exhibited in Figure 3, were attributable to the formation of a titanium dioxide layer on the surface of the foil. However, it should be noted that such chemical changes cannot be determined through visual inspection alone (i.e. scanning electron or optical microscopy). Therefore, elemental characterization of the surface treated CP titanium foil was required to confirm an increase in oxygen content at the foil surface, consistent with the presence of a titanium dioxide layer. To quantify the chemical composition of the material, a total of four 6.35 mm by 6.35 mm samples were cut from random locations in two surface treated and as-received titanium foil sheets. These samples were then analyzed using an energy dispersive X-ray (EDX) spectroscopy system attached to the JEOL scanning electron microscope. Figure 4 provides the representative EDX spectra for both the as-received and surface treated titanium foil, with each spectrum normalized and offset for clarity. From this figure and the elemental reference peak energy values listed in Table 1, it is evident that the surface treated titanium foil exhibits a clear peak corresponding to the oxygen Kα x-ray radiation energy (0.525 keV). Comparatively, the as-received titanium foil contains only Ti peaks (Ti Kα, Ti Kβ, and Ti LI), which is expected for an unalloyed and theoretically uncontaminated material. From the EDX measurements, the oxygen Kα peak obtained for the surface treated samples indicates the presence of the previously assumed oxide layer on the titanium foil surface.

While the comparative EDX spectra of Figure 4 are ideal for determining the elemental composition of a given material, they provide no information as to its crystalline structure. This distinction is particularly important for titanium dioxide, since it naturally occurs in three distinct polymorphs; rutile, anatase, and brookite. Of these three, rutile, represents the most common phase [15]. Given that the positions of the titanium and oxygen atoms within the TiO\textsubscript{2} lattice and the atomic packing density are both functions of the crystalline phase, structural characterization is essential for accurate molecular-level modeling. In order to determine the crystalline structure of the titanium dioxide surface layer, a total of four 5.08 cm by 2.54 cm samples were cut from random locations in the same as-received and surface treated titanium foil sheets. These samples were subsequently analyzed using a Panalytical X’Pert Pro X-ray powder diffraction (XRD) system. X-ray intensity was measured for each sample at room temperature over a detector angle (2θ) range of 25° to 45° using a Cu-target X-ray tube with wavelength of 1.542 Å, operated at 1800 watts.

Normalized X-ray diffraction patterns obtained using these settings are shown in Figure 5 and Figure 6 for the as-received and surface treated CP titanium samples, respectively. As with the EDX spectra discussed previously, there is an evident distinction between the diffraction patterns of the as-received and surface treated titanium foil. As expected, detector angles corresponding to the X-ray intensity peaks of Figure 5 show good correlation with the powder diffraction file (PDF) reference peak locations for unalloyed α-phase titanium, with the highest intensity peaks occurring at detector angles of
Figure 2. Scanning electron micrograph (2000× magnification) of the as-received CP titanium foil surface showing preexisting oxide layer preferentially aligned with foil rolling direction (length of the page).

Figure 3. Scanning electron micrograph (2000× magnification) of the treated CP titanium foil surface.

38.3° and 40.2° [16]. The detector angles which correspond to observed peaks in the as-received samples and correspond to PDF reference peak locations for α-phase titanium are provided in Table 2. In comparison, investigation of the surface treated titanium XRD pattern of Figure 6 shows the same α-phase Ti peaks, although at reduced relative intensities and slightly shifted to lower detector angles. Additionally, a series of peaks attributable to the titanium dioxide surface layer are also evident in the treated CP titanium samples. In Figure 6, the highest intensity peak now resides at a detector angle of 27.4°. Comparison of this XRD pattern with the PDF
Figure 4. EDX spectra for as-received (bottom) and surface treated (top) CP titanium foil with elemental reference peak indicators.

Table 1. Measured EDX energy values for as-received and surface treated CP Ti.

<table>
<thead>
<tr>
<th>EDX Spectrum Type</th>
<th>X-Ray Energy (keV)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Ti L</td>
</tr>
<tr>
<td>As-Received CP Titanium</td>
<td>0.39</td>
</tr>
<tr>
<td>Surface Treated CP Titanium</td>
<td>0.43</td>
</tr>
<tr>
<td>Reference Peak Energies</td>
<td>0.40</td>
</tr>
</tbody>
</table>

Table 2. Measured XRD peak detector angles for as-received surface treated CP Ti.

<table>
<thead>
<tr>
<th>Spectrum Type</th>
<th>Peak Locations - Detector Angle (2θ)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Peak 1</td>
</tr>
<tr>
<td>As-Received CP Titanium</td>
<td>-</td>
</tr>
<tr>
<td>Surface Treated CP Titanium</td>
<td>27.44</td>
</tr>
<tr>
<td>PDF 44-1294 (α-Ti)</td>
<td>-</td>
</tr>
<tr>
<td>PDF 21-1276 (Rutile TiO₂)</td>
<td>27.41</td>
</tr>
</tbody>
</table>
Figure 5. XRD pattern for as-received CP titanium with reference peak locations.

Figure 6. XRD pattern for surface treated CP titanium with reference peak locations.
references for titanium dioxide polymorphs indicates that the titanium dioxide surface layer is in the rutile crystalline phase, as indicated by the peak location detector angles provided in Table 2 [17].

2.3. Titanium-Graphite Fiber Metal Laminate Fabrication

Following characterization of the titanium dioxide surface layer, a set of two 10.16 cm by 15.24 cm titanium-graphite fiber metal laminate panels were fabricated, each with an identical [0/45/Ti/-45/90], stacking sequence. A [0/45/Ti/-45/90], layup was chosen since it had been shown previously to provide the optimum specific properties (property ratio with respect to density) for mechanically fastened titanium-graphite fiber metal laminate joints [18]. The polymer matrix composite layers of these TiGr FML panels consisted of unidirectional carbon fiber-reinforced toughened epoxy prepreg tape (CYCOM 977-3/IM7), obtained from Cytec Industries. Panel consolidation was performed in a 2 ft by 5 ft Thermal Equipment autoclave, according to the prepreg manufacturer’s suggested cure cycle, with a maximum temperature of 177°C and a sustained application pressure of 0.586 MPa [19]. After completion of the panel cure cycle, an optical microscopy investigation of each TiGr panel cross-section indicated that the panels were well consolidated, with the metal-composite interfaces free of discontinuities, as shown in Figure 7. In addition, enhanced magnification SEM images obtained at the metal-polymer interfaces revealed excellent continuity between the dissimilar materials down to the micron-scale, as evidenced by Figure 8. In this figure, the bright line at the boundary between the titanium reinforcement and the epoxy matrix corresponds to the titanium dioxide surface layer. Since titanium dioxide is highly resistive, particularly with respect to the titanium substrate and the IM7 carbon fibers, charging effects from the SEM electron beam make the TiO2 layer highly visible.

3. Simulation of the Interfacial Molecular Structure

3.1. Computational Methods for Molecular Simulation

In order to numerically reproduce the bonded metal-composite interface shown in Figure 8, the molecular basis for adhesion between the titanium dioxide and epoxy constituents was evaluated using molecular dynamics (MD) simulations. Molecular dynamics is essentially a numerical solution to a general n-body problem, in which the interaction physics between neighboring bodies (atoms) are approximated using a pre-defined force-field or potential energy expression. Given that these simulations are classical in nature, quantum mechanics
effects or other related phenomena are not considered. In this analysis, commercially available MD simulation software (Materials Studio) from Accelrys Inc. was used to generate molecular-level structure and property predictions for metal-composite interfaces in TiGr FMLs. The Materials Studio software suite contains a version of the condensed-phase optimized molecular potentials for atomistic simulation studies (COMPASS) forcefield, an ab initio forcefield which has been parameterized and validated using condensed-phase properties [20]. In the COMPASS forcefield, the general relationship between any two atoms in the simulation can be defined by the following functional form of the potential energy expression, which is a linear combination of all possible bonded (B), cross-term (CT), and non-bonded (NB) interactions [20]:

\[
U_{\text{total}} = U_B + U_{CT} + U_{NB}
\]

Specific forms of the bonded and cross-term interactions in the above potential energy expression can be obtained from literature. However, since this analysis is primarily concerned with the non-bonded interactions between atoms that describe the behavior of physisorbed interfaces, the functional form of \( U_{NB} \) is provided in Equation (2):

\[
U_{NB}(r_{ij}) = \frac{\alpha_i q_i q_j}{r_{ij}} + \left[ \frac{A_{ij}}{r_{ij}^{12}} - \frac{B_{ij}}{r_{ij}^{6}} \right]
\]  

In the above equation, the non-bonded interaction energy component consists of electrostatic and Leonard-Jones potential terms, where \( r_{ij} \) is the spatial position vector between the two atoms and \( q_i \) and \( q_j \) are the atomic charges. Additionally, \( A_{ij} \), \( B_{ij} \), and \( \varepsilon \) are material-dependant constants, which must be determined for any unique interaction between two atoms in the simulation. A major advantage of a commercially available software package, such as Materials Studio, is that the material constants necessary in the evaluation of Equation (2) have been previously determined for a variety of atomic configurations. Therefore, no additional parameterization of the COMPASS forcefield is required.

3.2. Molecular Modeling of the Metal-Composite Interface

To simulate the mechanics of adhesion at the titanium dioxide-epoxy interfaces in a TiGr FML, the COMPASS forcefield and the functional forms of the system’s total potential energy and the non-bond interaction energy provided in Equations (1)-(2) were used in a series of MD simulations to construct representative models of the molecular-level interfacial structure. Formation of a titanium dioxide simulation cell was the first step in the construction of this model, requiring knowledge of the TiO\(_2\) lattice parameters and atomic positions, as well as the exposed crystalline surface. From the X-ray diffraction pattern of Figure 6, it was previously determined that the CP titanium reinforcement layers in the TiGr
FMLs possessed a rutile-phase titanium dioxide surface structure. The unit cell of this particular TiO₂ polymorph is tetragonal, with a TiO₆ octahedron forming the basic structural unit of the crystal. The lattice parameters and atomic coordinates necessary for molecular modeling of rutile TiO₂ are well established in the literature, and are listed in Table 3 [22]. However, it should be noted that the exterior faces of the unit cell formed by these lattice parameters do not necessarily correspond to the exposed TiO₂ crystallographic plane at the surface of the CP titanium reinforcement. In rutile TiO₂, the predominately exposed crystalline cleave plane is the (110) face, and thus all titanium dioxide surface models presented in this study utilize TiO₂ crystals terminated with the (110) plane [23]. Because the dimensions of a single TiO₂ unit cell are much too small to produce physically relevant results, the simulation cell surface area (Aₛ) was increased by extending the crystal in each of the in-plane directions, as illustrated in Figure 9. This TiO₂ supercell was approximately tetragonal with dimensions of A=20.71 Å, B=19.47 Å, and C=14.94 Å. Prior to layering the polymer molecules on top of this TiO₂ supercell, a minimization of the supercell total potential energy was performed using the COMPASS forcefield to obtain the equilibrium configuration of the exposed surface. As expected, the positions of all atoms in the TiO₂ supercell did not significantly change following energy minimization, confirming the accuracy of the lattice parameters and atomic coordinates provided in Table 3.

After building the titanium dioxide surface model, the interface between the metal and composite constituents in a TiGr FML was replicated at the molecular-level by layering a cross-linked epoxy structure on top of the energy minimized TiO₂ surface. Exclusion of the reinforcing fibers from the molecular-level interface model is a reasonable assumption, since SEM investigation showed that the carbon fiber-titanium separation distance was several orders of magnitude larger than the MD unit cell. It should also be noted that the stipulation of a cross-linked epoxy layer complicated the analysis somewhat, since the number and location of these cross-links was not known a priori. Therefore, it was first necessary to model the transformation of the linear epoxy molecules into a highly networked structure before analyzing the resultant interface. Given that the behavior of the epoxy resin near the interface will, in general, deviate from the behavior of the bulk epoxy, the MD cross-linking simulations were performed in the presence of the TiO₂ surface. Although the actual formulation of the epoxy resin (CYCOM 977-3) is proprietary, it is widely assumed that this material is a tetraglycidyl methylene dianiline (TGMDA) and diamino-diphenyl sulfone (DDS) blend, where the chemical

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**Figure 9.** Titanium dioxide unit cell and supercell configurations used for molecular simulation.

**Table 3.** Lattice parameters and symmetry atomic coordinates for rutile TiO₂ [22].

<table>
<thead>
<tr>
<th>TiO₂ Lattice Quantity</th>
<th>a</th>
<th>b</th>
<th>c</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lattice Parameters (Å)</td>
<td>0.4595</td>
<td>0.4595</td>
<td>0.2959</td>
</tr>
<tr>
<td>Ti Atomic Position (Fractional Coordinates)</td>
<td>0.0000</td>
<td>0.0000</td>
<td>0.0000</td>
</tr>
<tr>
<td>O Atomic Position (Fractional Coordinates)</td>
<td>0.3048</td>
<td>0.3048</td>
<td>0.0000</td>
</tr>
</tbody>
</table>
structures and molecular models for each linear epoxy constituent are shown in Figure 10 [24]. The individual epoxy constituents presented in Figure 10 were then energy-minimized and a total of eight distinct amorphous polymer cells were created, containing 74.5% TGMDA to 25.5% DDS by weight. Due to the highly unstructured nature of the liquid resin, these eight independent epoxy configurations were required to effectively ‘average out’ configuration-space effects. The planar dimensions of the amorphous epoxy cells were set equal to the lattice parameters of the TiO2 surface (A=20.71 Å and B=19.47 Å), with the cell height constrained by the liquid-state density of the epoxy resin (1.16 g/cm³), determined from the manufacturer’s datasheet [19]. Each of the eight independent configurations were then layered on top of the titanium dioxide surface and a 20 Å vacuum spacer layer was added to the top of the simulation cells. Figure 11 provides two-dimensional views for two of the eight unique

![TGMDA Chemical Structure](image1)

![DDS Chemical Structure](image2)

![TGMDA Molecular Model](image3)

![DDS Molecular Model](image4)

**Figure 10.** Epoxy resin monomer (TGMDA) and cross-linking agent (DDS) molecular structure.

![Figure 11.](image5)

**Figure 11.** MD interface model for the TiO2-linear epoxy system.
TiO$_2$-linear epoxy interface configurations, each containing 1630 atoms. In the figure, it is important to note that periodic boundary conditions were applied for the in-plane directions (1- and 2-directions in Figure 11), but the presence of the vacuum spacer eliminated periodicity in the thickness (3) direction.

In order to imitate formation of the networked epoxy structure, molecular dynamics cross-linking simulations were performed using the COMPASS forcefield with an isochoric-isobaric (NVT) canonical ensemble for each of the eight TiO$_2$-linear epoxy simulation cells. In each case, the equilibrium configuration of the interface was determined from 100 ps of dynamics time at 450K, which corresponded to the maximum applied temperature in the TiGr consolidation process [19]. The time step between successive dynamics increments was set at 1 fs, and a cutoff radius of 9.5 Å was used to determine the maximum separation distance ($r_{ij}$) at which the non-bond energy of Equation (2) was evaluated for each atom pair. Given that the potential energy of the TiO$_2$ surface had previously been minimized, the positions of all atoms in the titanium dioxide supercell were fixed in place during the molecular dynamics run. A fixed surface assumption is valid since the TiO$_2$ lattice parameters and atomic coordinates do not significantly vary (0.16%) over the temperature range of interest (298K to 450K) [22].

During the MD analysis, trajectories for all epoxy atoms in the model were stored at every 5 ps of simulation time, and were used to predict the resulting cross-linked structure according to the method outlined by Yarovsky and Evans [25]. In this method, cross-link formation in a thermosetting polymer is assumed to be directly related to the distance between the active sites of the linear polymer and cross-linking agent molecules. When these sites are within a specified interaction radius (6.0 Å in this study), cross-linking will occur. For a TGMDA/DDS system, the active sites for each molecule correspond to the epoxy (-C-O-C-) and amine (NH$_2$) functional groups, as illustrated in Figure 12. Replication of this cross-linking process was performed by calculating the position of each active epoxy site from the MD trajectories and applying a nearest neighbor search algorithm to identify the corresponding amine site in closest proximity. If the distance between these two groups was less than or equal to the interaction radius, then the corresponding TGMDA and DDS molecules were connected in the MD simulation cell according to Figure 12. Application of this method to all eight TiO$_2$-linear epoxy interface models resulted in an average cross-link efficiency of 83.9% after 100 ps of dynamics time.

It is important to point out that the geometric approximation of an essentially chemical process was necessitated by the classical nature of molecular dynamics simulations. Because MD analyses are based upon numerical solution of the Newtonian equations of motion, charge transfer phenomena associated with the formation and breakage of chemical bonds cannot be explicitly modeled using MD techniques without the inclusion of quantum mechanics effects. Given that quantum calculations for a system containing thousands of atoms become prohibitively expensive from a computational standpoint, the above geometric method represents the best predictive strategy for obtaining the final cross-linked configuration of the epoxy resin. Once the networked epoxy configurations had been formed, the final structure of each TiO$_2$-epoxy interface was determined from a 200 ps molecular dynamics run at room temperature (298K). MD analyses of the cooled interface were, once again, performed using the COMPASS forcefield and an NVT ensemble. In this manner, a total of 300 ps of MD simulation time were used to determine the equilibrium state of each interface configuration; 100 ps for the linear epoxy at 450K and 200 ps for the cross-linked system at 298K.

![Figure 12. TGMDA/DDS epoxy cross-linking reaction.](image-url)
4. Molecular-Level Interfacial Property Predictions

4.1. Interfacial Predictions across Multiple Length Scales

The room temperature molecular configurations obtained for each of the eight simulation cells were then used to generate mechanical property predictions for the metal-composite interfaces in TiGr FMLs. In this analysis, the properties of interest are the transverse elastic constants ($E_{33}$, $G_{13}$, $G_{23}$), ultimate normal and shear interfacial strength values ($\sigma_{33}^{\text{int}}$, $\sigma_{13}^{\text{int}}$, $\sigma_{23}^{\text{int}}$), corresponding strains at failure ($\epsilon_{33}^{\text{int}}$, $\epsilon_{13}^{\text{int}}$, $\epsilon_{23}^{\text{int}}$), and fracture energies ($U_{33}^{\text{int}}$, $U_{13}^{\text{int}}$, $U_{23}^{\text{int}}$). However, it should be noted that only three of these component groups are independent, due to the integral relationship between fracture energy and the stress-strain curves in each of the normal and shear modes. In this analysis, strain at failure was chosen as the interfacial dependent variable set, since its resolution from MD simulations contains the greatest ambiguity. To determine the remaining independent interfacial variables, a series of calculations were performed using the predicted TiO2-epoxy molecular structures obtained after 300 ps of MD simulation time. As will be discussed in the companion study, these MD-derived mechanical properties were integrated into a continuum-level finite element analysis via a traction-separation constitutive law. This propagation of molecular-level considerations up to the continuum-level is what gives the current model its multi-scale nature.

4.2. Calculation of Interfacial Fracture Energy

Of these interfacial mechanical properties, fracture energy is by far the most straightforward to determine from a molecular dynamics simulation. This is due to the analogous relationship between fracture energy and work of adhesion ($W_{ad}$) for a bonded interface, in which ‘failure’ is defined by separation of the heterogeneous interface into two distinct homogeneous components. For a chemical or mechanical bond between two materials, work of adhesion is the fundamental thermodynamic quantity which defines their dissociation at the molecular-level. Experimental measurements of the work of adhesion are difficult to obtain and often overestimate the stability of the interface. This is due to secondary energy dissipation mechanisms in the sample which do not contribute to debonding (e.g. specimen relaxation and plasticity) [26]. Calculation of the work of adhesion in a molecular dynamics simulation is comparatively simple, since $W_{ad}$ is linearly dependent on the interaction energy ($U^{\text{int}}$) between the interface components:

$$W_{ad}^I = \frac{-U^{\text{int}}}{(A^I)^n}$$

Where $(A^I)$ represents the surface area of the interface with normal direction $(I)$ and the interaction energy is defined by the total system energy and the surface free energies of each component of the interface (TiO2 and epoxy) as per the Dupré relationship [27]:

$$U^{\text{int}} = U_{\text{total}} - (U_{\text{epoxy}} + U_{\text{TiO2}})$$

Computation of each term in Equation (4) in an MD analysis is straightforward because potential energy must be determined at each time step in order to solve the governing equations of motion. Therefore, for each of the eight TiO2-epoxy interface configurations, the work of adhesion at 298K was determined by calculating the potential energy of the total system as well as that of the epoxy and titanium dioxide surfaces using the COMPASS forcefield at the final MD time step ($t = 200$ ps). Using this method, the predicted work of adhesion for a TiO2-epoxy interface was found to be $1.12 \pm 0.03$ J/m², where the uncertainty represents one standard deviation over all eight configurations. Because the surface area component of $W_{ad}$ in Equation (3) is always normal to the interface, it was assumed that the work of adhesion and fracture energy were mode-independent (i.e. $W_{ad} = U_{33}^{\text{int}}$, $U_{13}^{\text{int}}$, $U_{23}^{\text{int}}$). It should also be noted that while use of the final molecular trajectory ($t = 200$ ps) in the work of adhesion calculations was somewhat arbitrary, it did not greatly impact the predictions. This is due to the fact that the potential energy of each configuration converged to an equilibrium state after approximately 7 ps of dynamics time, as shown in Figure 13.

4.3. Calculation of Interfacial Normal and Shear Strength

In addition to fracture energy, the interfacial strength in the shear and normal deformation modes was also calculated from the equilibrium configurations of the TiO2-epoxy interface using energy considerations and the COMPASS forcefield [28]. From the potential energy form in Equation (1), the inter- and intra-molecular forces ($F_i$) acting on any atom in the MD simulation are related to the atomic total potential energy through the gradient operator:

$$F_i = -\nabla_i \left[ U_{\text{total}} \left( r_i \right) \right]$$

where summation is not applied for all lower case subscripts. While this equation is essential for the solution of the classical equations of motion in any MD analysis, it can also be used to calculate the tensile or shear stresses.
Figure 13. TiO$_2$-epoxy total potential energy as a function of MD simulation time at 298K for a representative interface configuration.

acting on an interface. Focusing solely on the interaction between the TiO$_2$ surface and the cross-linked epoxy layer, the intra-atomic forces required for bonding ($F_{\text{int}}^i$) can be obtained from Equation (5) if the total potential energy is replaced by the interaction energy, previously defined in Equation (4). Utilizing the classical mechanics definition of stress ($\sigma_{ij}$) as the action of a force ($F_j$) on a differential area with normal vector ($n_i$), the state of stress at the interface can be completely described by the interaction energy between the constituents:

$$\sigma_{ij} = -\nabla_j \left[U_{\text{total}}^i \left(r_g^i\right)\right]$$

(6)

Since the interfacial surface area remains approximately constant in an MD simulation performed using the NVT canonical ensemble, its value can be incorporated into the gradient operator. The interfacial stress tensor can be further simplified using the definition of work of adhesion provided in Equation (3):

$$\sigma_{ij} = -\nabla_j \left[W_{\text{ad}}^i \left(r_g^i\right)\right]$$

(7)

This represents the general form of the stress tensor associated with any arbitrary deformation of the interface. However, this analysis is only concerned with three distinct stress states ($\sigma_{33}^\alpha, \sigma_{13}^\alpha, \sigma_{23}^\alpha$), and thus the gradient in Equation (7) can be replaced with an ordinary differential operator, provided that deformation of the interface occurs in a purely normal or shear mode. For example, considering a state of pure tensile stress applied to the interface:

$$\sigma_{33} = -\frac{d}{d\delta_3} \left[W_{\text{ad}}^\alpha \left(\delta_3\right)\right]$$

(8)

where $\delta_3$ in the above equation represents the interfacial spacing between the titanium dioxide surface and the cross-linked epoxy layer. In order to evaluate the right-hand side of Equation (7), the work of adhesion was calculated at a finite number of interfacial separation distances, shown schematically for two extreme values of $\delta_3$ in Figure 14. Interfacial stresses were then computed at these discrete sampling points using a second-order centered finite difference approximation of Equation (7). Considering a purely tensile deformation mode, Equation (7) reduces to Equation (8) and this approximation becomes:

$$\sigma_{33} \cong \frac{-W_{\text{ad}}^\alpha \left(\delta_3 + 2\Delta\right) + 8W_{\text{ad}}^\alpha \left(\delta_3 + \Delta\right) - 8W_{\text{ad}}^\alpha \left(\delta_3 - \Delta\right) + W_{\text{ad}}^\alpha \left(\delta_3 - 2\Delta\right)}{12\Delta}$$

(9)
where (Δ) represents the incremental spacing between successive measurements of $W_{ad}$. Application of Equation (9) to the TiO$_2$-epoxy MD equilibrium structures generated in the preceding MD analysis produced a series of tensile stress separation curves, shown for two representative interface configurations in Figure 15. As expected, the interfacial stress is highly compressive at small separation distances ($\delta_3 \leq 1.5\text{Å}$), with its limit becoming unbounded as $\delta_3$ decays to zero. This behavior is a consequence of the repulsive component of the Lennard-Jones potential energy, which dominates non-bonded interactions at small atomic separation distances. Focusing on the bounded region of Figure 15, the interfacial tensile strength for each configuration was determined from the maximum point on the respective stress-separation curve. The average predicted tensile strength ($\sigma_{33}$) over all eight configurations was found to be $23.77 \pm 2.34$ MPa, where the uncertainty once again represents the measured standard deviation. This same analysis was also applied to determine the shear strength of the interface ($\sigma_{13}$, $\sigma_{23}$), with periodicity in one of the in-plane directions eliminated to simulate the presence of a free surface. The numerical methods outlined in Equations (8)-(9) were analogous for the determination of shear strength, with the only difference being that the interfacial separation distance ($\delta_3$) was replaced in these equations by the interfacial shear displacement between the TiO$_2$ surface and the cross-linked epoxy layer. The calculated strength values in each shear mode were approximately equal, and thus it was assumed that the transverse shear strength showed no directional dependence ($\sigma_{13}^{\sigma} = \sigma_{23}^{\sigma}$). Averaged over both shear deformation modes for all eight configurations, the predicted shear strength of the interface was equal to $18.61 \pm 1.54$ MPa.

4.4. Calculation of the Interfacial Elastic Constants

After determination of the fracture energy and strength in both the normal and shear modes, the interfacial elastic constants ($E_{33}$, $G_{13}$, $G_{23}$) represented the only remaining parameters required for characterization of the traction-separation constitutive model. In this analysis, the elastic constants were determined from the MD-predicted structure of the TiO$_2$-epoxy interface using the small-strain procedure of Theodorou and Suter [29]. In this method, the stiffness tensor of a material can be atomistically determined through slight perturbations of the system away from equilibrium, with subsequent measurements of the resulting change in total potential energy. The ‘small-strain’ provision of this method implies equivalence of the material and spatial configurations, such that all second-order deformation gradient terms are much smaller than their first-order counterparts. Assuming that this assumption is valid, the isothermal stiffness tensor ($C_{ijkl}$) of the interface can be defined in terms of the second derivative of the Helmholtz free energy ($A$) with respect to the strain tensor ($\varepsilon_{ij}$), normalized by the volume of the simulation cell ($V$):

$$
C_{ijkl} = \frac{1}{V} \left[ \frac{\partial^2 A}{\partial \varepsilon_{ij} \partial \varepsilon_{kl}} \right]_T
$$

In the above equation, the isothermal requirement is relatively inconsequential in this analysis since all MD simulations were performed using the NVT ensemble. Thus, the temperature of the system rapidly converged to its specified value after only a few picoseconds of dynamics time, illustrated for a representative interface configuration in Figure 16. Equation (10) can be further generalized using the definition of Helmholtz free energy as a function of the system total potential energy, temperature ($T$), and entropy ($S$):

$$
A = U_{total} - TS
$$

Substitution of Equation (11) into Equation (10) and expansion of the second-order strain tensor derivative for an isothermal system results in:
Figure 15. TiO$_2$-epoxy interfacial tensile stress vs. separation distance for selected configurations.

Figure 16. TiO$_2$-epoxy simulation cell temperature convergence to the NVT control point (298K) for a representative configuration.
In their detailed analysis of amorphous glassy polymers, Theodorou and Suter showed that entropic contributions to the above equation are relatively minor, and thus their effects can be neglected [29]. Using the same assumption, the material stiffness tensor can be approximated by the second derivative of the system’s total potential energy with respect to the strain tensor, provided that the vibratory contributions to the internal energy of the system are negligible:

\[
C_{ijkl} \equiv \frac{1}{V} \left[ \frac{\partial^2 U^{\text{total}}}{\partial \epsilon_{ij} \partial \epsilon_{kl}} \right]_T
\]  

(13)

Expression of any mechanical quantity in terms of the system’s total potential energy is highly advantageous since determination of the potential energy is a necessary condition for solution of the classical equations of motion in an MD analysis. Using the approximation of Equation (13), the individual stiffness components were determined by subjecting the equilibrium interface configurations to a series of small strain deformations and measuring the resultant change in potential energy. Switching from tensorial to reduced index notation and considering a state of pure tensile strain normal to the interface, the strain second-derivative term greatly simplifies, resulting in the following diagonal component of the stiffness matrix:

\[
C_{33} \equiv \frac{1}{V} \left[ \frac{\partial^2 U^{\text{total}}}{\partial \epsilon_3^2} \right]_T
\]  

(14)

The other 35 diagonal and off-diagonal components of \( C_{ij} \) were determined analogously using small perturbations of the interface away from its equilibrium configuration in various deformation modes. It is important to note that calculation of all 36 components of the stiffness matrix was necessary, since the method outlined in Equations (10)-(13) will generally produce a fully populated, asymmetric stiffness matrix due to spatial non-uniformity on the molecular level. For each of the eight TiO\(_2\)-epoxy interface configurations, the system potential energy was measured using the COMPASS forcefield, with the strain second derivative evaluated using a finite difference approximation, akin to that of Equation (9). This produced the following average and standard deviation values (in GPa) of the TiO\(_2\)-epoxy interfacial stiffness matrix components: (Equation (15))

\[
[C] = \begin{bmatrix}
11.06 & 6.12 & 6.79 & 0 & 0 & 0 \\
6.12 & 12.60 & 6.57 & 0 & 0 & 0 \\
6.79 & 6.57 & 13.04 & 0 & 0 & 0 \\
0 & 0 & 0 & 2.38 & 0 & 0 \\
0 & 0 & 0 & 0 & 2.37 & 0 \\
0 & 0 & 0 & 0 & 0 & 2.44
\end{bmatrix}
\]  

(15)

Comparison of Equations (15) and (16) shows that averaging the non-zero off-diagonal components \((C_{12}|C_{21}, C_{13}|C_{31}, C_{23}|C_{32})\) does not significantly change any of their values; therefore averaging was performed solely to conform to the tenets of classical mechanics. Inversion of the stiffness matrix in Equation (16) yields the compliance matrix, from which the requisite elastic constants of the interface were found to be: \(E_{33} = 8.01\) GPa, \(G_{13} = 2.38\) GPa, and \(G_{23} = 2.37\) GPa. As was the case for the interfacial shear strength and fracture energy, the shear moduli in each mode are effectively equivalent, and thus their average was used as the ‘true’ value of the mode-independent interfacial shear modulus. As a validation of the methodology of Equations (10)-(16), the same molecular-level analysis was repeated for the bulk epoxy material (i.e. no TiO\(_2\) surface structure included).
The bulk system, shown in Figure 17, underwent an identical simulation procedure; amorphous cell construction, elevated temperature cross-linking, room temperature cooldown, and small-strain perturbations from equilibrium, for a total of eight different spatial configurations. The resultant stiffness matrix for the bulk epoxy was passed through the same anisotropy and asymmetry filters. Neglecting small molecular-level spatial non-uniformity, the isotropic elastic constants of the material \(E = 3.66\) GPa and \(G = 1.41\) GPa were very close to the values provided by the manufacturer \(E = 3.52\) GPa and \(G = 1.30\) GPa), thus validating the procedures outlined in Equations (10)-(16) [19]. With the calculation and validation of these interfacial elastic constants, all requisite molecular mechanical properties of the TiO\(_2\)–epoxy interface were fully determined, as outlined in the methodology of Equations (3)-(16). These MD obtained material properties were subsequently used in a continuum-level finite element model to define the mechanical behavior of the metal-composite interfaces in a TiGr FML, described in the second portion of this study.

5. Summary and Conclusions

Titanium-graphite fiber metal laminates are promising candidate materials for structural applications in which either polymer matrix composite or monolithic metal components are considered unfit. However, TiGr implementation has been slowed by the relatively poor metal-composite interlaminar bond strength obtained during the FML consolidation process. This interface often represents the ‘weakest link’ in the titanium-graphite layup, and its comparatively poor mechanical performance diminishes the strength and weight advantages associated with TiGr FMLs. Therefore, a comprehensive understanding of the mechanics of adhesion at these metal-composite interfaces represents the first step towards enhancement of TiGr FML interlaminar properties.

In this analysis, construction of a molecular-level interface model was discussed, with interfacial mechanical properties obtained from thermodynamics and structural mechanics considerations. The properties derived herein will be propagated up to the continuum-level in the second portion of this study, allowing for experimental validation of the multi-scale method. It should be noted that, as a standalone system, the molecular-level model showed good correlation when used to predict the behavior of the bulk TiGr polymer matrix material. As such, it can be concluded that the presented molecular-level model can be utilized independent of the multi-scale framework. Furthermore, there is no need for this model to be restricted only to bonded metal-composite interfaces in titanium-graphite fiber metal laminates, or any other FML variant. In fact, the methodology presented herein could easily be applied to any physisorbed metal-polymer interface; provided that the length scales associated with individual constituent materials are such that their structure can be completely defined within the confines of a molecular dynamics simulation cell. The physisorption requirement associated with this model symbolizes one potential area of improvement, as only secondary bonding effects (van der Waals and electrostatic) are presently considered between the two components of the interface. Inclusion of covalent bonding between the constituents would allow for extension of this method to functionalized metal-polymer interfaces, which possess a mixture of chemical, mechanical, and electrostatic effects. Recent advances in computational methods and chemical simulation techniques have made this prospect feasible, although a significant amount of research is required for its actual implementation.

6. Acknowledgements

This study was principally supported by Raytheon Missile Systems with Andrew Facciano as the program monitor. Additional funding was provided by the United States Department of Education through the Graduate Assistance in Areas of National Need (GAANN) fellowship program. SEM and EDX results presented in this study utilized additional support from NSF IGERT: Materials Creation Training Program (MCTP) — DGE-0654431 and the California NanoSystems Institute (CNSI). The authors would like to thank Julio de Unamuno IV of Accelrys Inc. for his helpful discussion of the molecular
dynamics simulations employed in this study. Additionally, special thanks go to Rachel Hundley for her invaluable editorial comments.

7. References


