

Nano-engineered Composites: Interlayer Carbon Nanotubes Effect

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Received: November 3, 2012; Revised: December 19, 2012

The concept of carbon nanotube interlayer was successfully introduced to carbon fiber/epoxy composites. This new hybrid laminated composites was characterized by Raman spectroscopy, X-ray diffraction, scanning electron microscopy and tensile tests. An increase on peak stress close to 85% was witnessed when CNTs interlayer with 206.30 mg was placed to carbon fiber/epoxy laminates. The failure mechanisms are associated to CNTs distribution between and around carbon fibers. These CNTs are also responsible for crack bridging formation and the increase on peak stress. Initial stiffness is strongly affected by the CNT interlayer, however, changes on stiffness is associated to changes on nano/micro-structure due to damage. Three different behaviors can be described, i.e. for interlayers with ≈ 60 mg of CNT the failure mode is based on cracks between and around carbon fibers, while for interlayers with CNT contents between 136 mg and 185 mg cracks were spotted on fibers and inside the CNT/matrix mix. Finally, the third failure mechanism is based on carbon fiber breakage, as a strong interface between CNT/matrix mix and carbon fibers is observed.

Keywords: carbon nanotubes, carbon fiber/epoxy composites, interlayers, raman spectroscopy

1. Introduction

As described by Saito et al.¹, carbon nanotube is a honeycomb lattice rolled into a cylinder. Carbon nanotubes (CNTs) have been the center of many researches due to their dimensions and remarkable electro-mechanical properties. In general, a CNT diameter has a nanometer size and its length can be more than 1 μm . Its large aspect ratio (length/diameter) is appointed as one of the reasons for the CNTs notable properties. According to Kalamkarov et al.², single-walled nanotubes (SWNTs) have predicted specific strength around 600 times larger than steel. CNT capabilities have been observed experimentally and verified by numerical simulations. Frankland et al.³, Jin and Yuan⁴ and Agrawal et al.⁵ are among those researchers who employed molecular dynamics for analyzing CNTs. The atomistic simulation approach was employed by Belytschko et al.⁶, Lurie et al.⁷, Gates et al.⁸, while the nano-mechanics modeling was described by Liu et al.⁹, Ruoff and Pugno¹⁰, Li and Chou¹¹, Ávila et al.¹². Although CNTs have tremendous potential in a large variety of applications, e.g. aerospace and medical industries, there is no consensus about their exact mechanical properties. The experiments performed up to now have presented large variability due to the inherent complexity of manipulating these materials. However, their potential is unquestionable, in special for composites.

As mentioned by Ávila et al.¹³, carbon based nano-structures, i.e. carbon nanotubes and graphene nano sheets (GN), can be combined to traditional composites for a multi-scale reinforcement. Moreover, with the recent developments on CNT synthesis, its price per gram has dramatically reduced. As a consequence, the number of

researchers using carbon based nanostructures increased, and the results on nano-reinforcement of composites laminates are encouraging. Among those researchers are Kim and collaborators¹⁴ whom described no significant increase on tensile properties of the addition of CNTs to carbon fibers/epoxy laminates. Nonetheless, they noticed an enhancement on flexural modulus ($\approx 12\%$) and strength ($\approx 18\%$) with the addition of 0.3 wt. (%) of CNT to the epoxy system. This increase can be attributed to changes into flexural failure mechanisms. Following the same idea, Chou et al.¹⁵ discussed the influence of CNTs into the failure of laminated composites. They even proposed the concept of a hybrid inter-laminar architecture that can bridge inter-laminar cracks. Wicks and colleagues¹⁶ actually produced the hybrid nano reinforced laminated composites proposed by Chou et al.¹⁵ In their laminate, CNTs were grown *in situ* in all fibers leading to a “fuzzy” fibers configuration. As mentioned by Wicks, aligned CNTs bridges the plies interfaces, which can lead to an increase on toughness, for the steady state condition, 76% higher than the conventional laminated systems. Notice that for the interlayer nano reinforcement some issues must be considered, i.e. the interfacial bonds between carbon nanotubes and the fiber/matrix system and the length effect into this “grip condition”. To understand the failure mechanism, Shokrieh and Rafiee¹⁷ modeled the CNT length effect on reinforcement effectiveness. Moreover, they concluded that for carbon nanotubes with length less than 100 nm, the improvement on stiffness for CNT/polymeric systems is negligible. By taking into consideration Shokrieh and Rafiee¹⁷ conclusions and the work done by Ma et al.¹⁸ for dispersing carbon nanotubes

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into polymeric matrices, it is clear that although CNTs are a viable option for reinforcement of nanocomposites and its hybrid companions. However, the idea of dispersing nanotubes into polymeric matrices as described by Shokrieh and Rafiee¹⁷ and Ma et al.¹⁸ seems to be limited. The “fuzzy” fibers configuration developed by Wicks et al.¹⁶ is also limited as all plies have to be loaded with carbon nanotubes. This increase on “fiber density” due to the “CNTs loads” can lead to difficulties for using the vacuum assisted impregnation system, as resin flow channels will be reduced. It is clear that alternative techniques must be developed.

Different techniques have been tested for incorporating CNTs into composite materials. The CNT infusion into laminated composites and its alignment by applying an electric field after the infusion was studied by Domingues et al.¹⁹. The major criticism about Domingues’ work is the amount of CNT dispersed which is around 0.1 wt. (%). Another approach tried to link CNTs to laminated composites was implemented by Wu et al.²⁰. Wu’s work was based on electrochemical grafting of CNTs on carbon fibers surface. Although the technique described by Wu et al.²⁰ seems to be effective, it is limited to the CNT concentration into the solution. Moreover, as noticed by Wu, there were “preferential regions” for CNTs direct attachment to carbon fibers. These preferred sites were fibers’ grooves and edges. This phenomenon led to a non-uniform distribution of CNT on carbon fibers surface. Another technique used for attaching CNTs to carbon fibers was studied by De Riccardis et al.²¹ and Vilatela et al.²². In their case, the chemical vapor deposition (CVD) technique was employed for directly grown CNTs into carbon fibers. De Riccardis’ work was based on deposition of nickel clusters and later on the CNTs were grown by hot filament chemical vapor deposition (HFCVD) technique. By using ferrocene as precursor, and CVD as the growing process, Vilatela was also able to obtain good quality CNTs. Moreover, the CVD technique employed by Vilatela et al.²² seems to be much simpler and easier to control. Although the results presented by De Riccardis et al.²¹ and Vilatela et al.²² seem to be encouraging much work has to be done for applications to laminated composites, in special high performance carbon fiber/epoxy systems.

As commented by Khan and Kim²³, the incorporation of nanoscale CNTs with conventional micro scale fiber reinforcement in polymeric composites can be obtained by modifying either the matrix or the fibers. Qian et al.²⁴ mentioned that the most common manufacturing process for hybrid composites (composites with at least two levels of reinforcement, at micro and nano-scale) is based on shear mixing the CNTs into the resin system, followed by infusion/impregnation and consolidation processes. This entire manufacturing procedure is limited by the CNTs loadings due to the matrix viscosity increase. Karapappas et al.²⁵ employed a torous mill device for dispersing CNTs into an epoxy system, a later on the hand layup process was applied followed by an autoclave consolidation. For Mode I fracture toughness, the results were consistent, in other words, an increase on CNT levels will lead to a correspondent increase on Mode I fracture toughness. For Mode II, however, a different scenario was noticed. The intermediate concentration (0.5 wt. (%)) got the highest value. This could indicate some problems during the dispersion process and at interface nano-modified matrix/fibers. To try to overcome this problem Seyhan et al.²⁶ used the three-roll mill. Although the dispersion quality was

improved, the matters of CNTs loadings remained. They were able to disperse only 0.1 wt. (%) of CNTs.

Khan and Kim²³, however, discussed alternative techniques such as, direct CNTs in situ growth, manual spreading or spraying the CNTs among others. The work reported by Garcia et al.²⁷ can be described as the in situ CNTs growth process. In this case, they employed alumina fibers and hand layup process. They reported an increase on interlaminar shear strength (ILSS), close to 69% with small standard deviation.

The work done by Arai et al.²⁸ fits on second category, as they dispersed a CNT powder using a sifter into carbon/epoxy prepregs. They discussed the influence of the testing method into the interlaminar shear strength. A similar approach, the sifter usage into nano/micro structures (in this case vapor grown carbon fibers - VGCF) was employed by Li et al.²⁹. Based on mode I fracture toughness results presented by Li et al., it seems that a saturation limit on VGCF is around $\text{g}\cdot\text{m}^{-2}$. Moreover, the zigzag fracture pattern could be the reason for the increase on fracture toughness, as the energy release rate is higher. The work done by Joshi and Dikshit³⁰ followed the idea of dispersing/transfer CNTs into prepreg surfaces. They sprayed an ethanol/CNT solution into a peel cloth and then the CNTs were mechanically transferred to the prepregs by applying a pressure. This procedure has two major problems: first ethanol is harmful for prepregs and the second problem is that by spraying the solution into the cloth and later transferring to the prepregs, there is always the possibility of agglomeration and non-uniform CNT distribution. This seems the case for high CNTs concentrations ($2.25 \text{ g}\cdot\text{m}^{-2}$ and $3.62 \text{ g}\cdot\text{m}^{-2}$) employed by Joshi and Dikshit³⁰. To avoid the problem of prepregs degradation by ethanol, Davis and Whean³¹, used not impregnated carbon fibers. The amount of CNTs employed was 0.3 wt. (%) and 0.5 wt. (%), a very small percentage. After spraying the ethanol/CNT solution into the fibers, they allowed the ethanol evaporation, and later employed the heated vacuum resin transfer molding process (HVRTM). Although they reported an increase on failure initiation close to 23%, their standard deviation indicates problems into the CNTs homogeneity. The resin flow during the HVRTM could simply wash out the CNTs in different regions of the composite panel. A more efficient way to transfer CNTs to prepregs and/or not impregnated fibers was described by Garcia et al.³². In their case, they did an in situ CNT growth in a Si substrate, and the transfer were performed by rolling the prepregs into a cylinder and then the cylinder was rolled over the Si substrate while applying a small pressure. They reported an increase on energy release rate in Mode II close to 300% with small standard deviation.

This paper discusses the advantages and disadvantages of CNTs in situ growth into not impregnated carbon fibers where the CNT forest is partially filled by the net resin during the impregnation process. Differently from the work done by Garcia et al.³² where the CNT forest was completely filled with resin, in this paper the CNT forest was partially filled by the net resin. By employing this strategy, the bridging condition is explicitly imposed.

2. Material and Methods

As discussed by Mathur et al.³³, thermal CVD has high growth rate, i.e. $10 \mu\text{m}/\text{min}$, and the CNTs diameters ranges

from ≈ 40 -60 nm. Moreover, Raman spectroscopy indicates a good crystallinity of multiwall carbon nanotubes (CNT's). In this research, a thermal CVD device from *FirstNano* was used for growing CNTs directly into glass fibers. CNTs can be grown over a temperature range of 600-900 °C using thermal CVD. After a series of tests, the optimum temperature, considering the CNTs alignment, was selected as 750 °C at an inert atmosphere (argon). The precursor employed was a mix of ferrocene and toluene (2%). The precursor was evenly distributed into the fibers' cloth, and later on the fibers' cloth with the precursor was placed in the furnace main heating zone. The temperature gradient from the furnace center to the edge was ≈ 10 °C.cm⁻¹, at a nominal temperature of 750 °C. The CNTs grown for this research were mainly Multiwall Carbon Nanotubes (MWNT).

The plain weave fiber glass used in this research has an areal density of 180 g.m⁻². To be able to understand the carbon nanotubes effect as interlayer into carbon/epoxy laminated composites, five different CNTs forests were grown *in situ* into the plain weave fiber glass. These fiber glass fabrics loaded with CNTs were placed between two layers of carbon fibers. The carbon fibers have a plain weave configuration and areal density of 200 g/m². The epoxy system employed here is based on *diglycidil ether bisphenol A* (DGBA) resin and an amine hardener, i.e. AR300 and a mix of AH300/AH150 supplied by Barracuda Composites Inc. The resin/hardener ratio employed was 100:27. The fiber/epoxy system ratio is equal to 50:50. The resin impregnation is based on hand lay-up while the composite final consolidation is performed by cure under vacuum. The total time for cure was 24 hours, 6 under vacuum and the remaining under air.

According to Koo et al.³⁴, during the nanoparticles dispersion into polymeric matrices nano-structures are formed. The two most common detection techniques to nano-structures identification are X-ray diffraction and electron microscopy. In this research, X-ray diffraction (XRD) experiments were carried out on a Shimadzu XRD-6000 X-ray diffract meter with Cu ($\lambda = 0.154$ nm) irradiation at 40 kV and 30 mA using a Ni filter. Data were recorded in the range from 2 to 80 deg in a continuous scanning at 2 degrees per minute and sampling pitch of 0.02 deg.. The high resolution scanning electron microscope (HRSEM) used was a Quanta 200 - FEG - FEI, while the transmission electron microscope (TEM) employed was a

Tecnai – G2-20-FEI. As mentioned by Dresselhaus et al.³⁵, Raman spectroscopy is also a powerful tool for studying carbon based nanostructures, i.e. CNTs and graphene nanosheets. In this study, the Raman spectroscopy analysis was performed using a DilorXY 800 Raman spectrograph.

DilorXY 800 Raman spectrograph has an 800 mm focal length triple spectrograph, f/6 subtractive or additive fore-monochromator – Gratings 1200 g.mm⁻¹, spectral range 27,000 cm⁻¹ to 5 cm⁻¹, and it is equipped with a 2 watts argon laser with wavelength of 514.5 nm or 488 nm, a 10 nW HeNe laser with wavelength of 632.8 nm. The scanning is performed by a motorized xy stage for confocal microscope with 0.1 μ m resolution and software for Raman map.

The hybrid laminated composite mechanical characterization is based on tensile tests following ASTM D 3039 standard³⁶. An EMIC DL-10000 universal testing machine with 10 KN and a 500N load cells was used to perform the tensile tests at constant displacement of 0.5 mm/min. Load and displacement were continually measured by the load cell and extensometer, respectively. Furthermore, as this research focuses on understanding the CNT inter-lamina effect into laminated composites, the concept of representative volume element (RVE) described by Ávila et al.¹² was employed. The hybrid composite laminate was composed of two layers of carbon fiber/epoxy with an extra inter-layer of fiber glass fabric where CNTs were grown *in situ* by thermal CVD. The total average thickness was around 0.145 mm. The other dimensions followed the ASTM D 3039 standard. By applying the RVE concept, it was possible to obtain a direct relation between the CNTs effects into the composite's macroscopic behavior.

3. Data Analysis

As this research deals with multi-scale composite reinforcement, i.e. from nano to macro, two different approaches are employed. The first one is related to nano and micro structure analysis based on HSEM, Raman spectroscopy and X-ray diffraction analysis. The second one is based to tensile tests and failure mode analysis. By analyzing the two sets of data, it is possible to correlate the nano/micro structure formed and the hybrid composite macroscopic behavior.

Figures 1a, b show the Raman spectroscopy and the X-ray diffraction signatures for the hybrid laminated

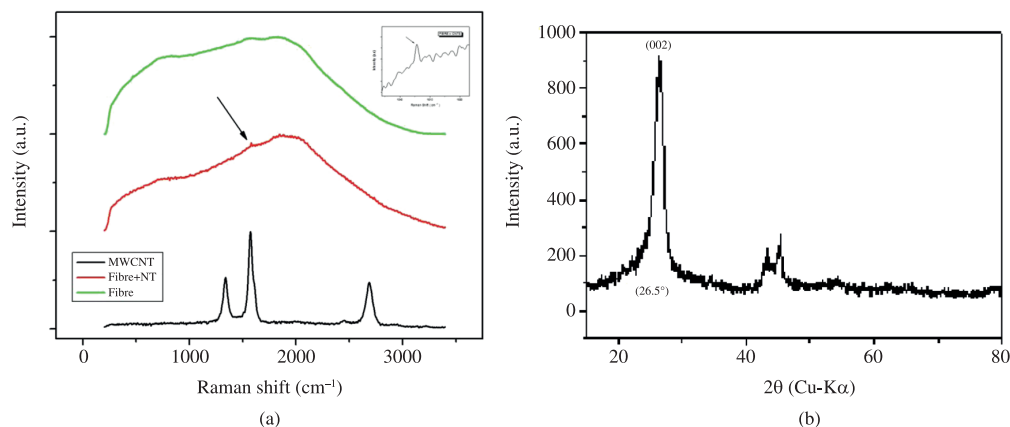


Figure 1. Fiber and CNTs characterization. (a) Raman spectroscopy; (b) CNTs' XRD signature.

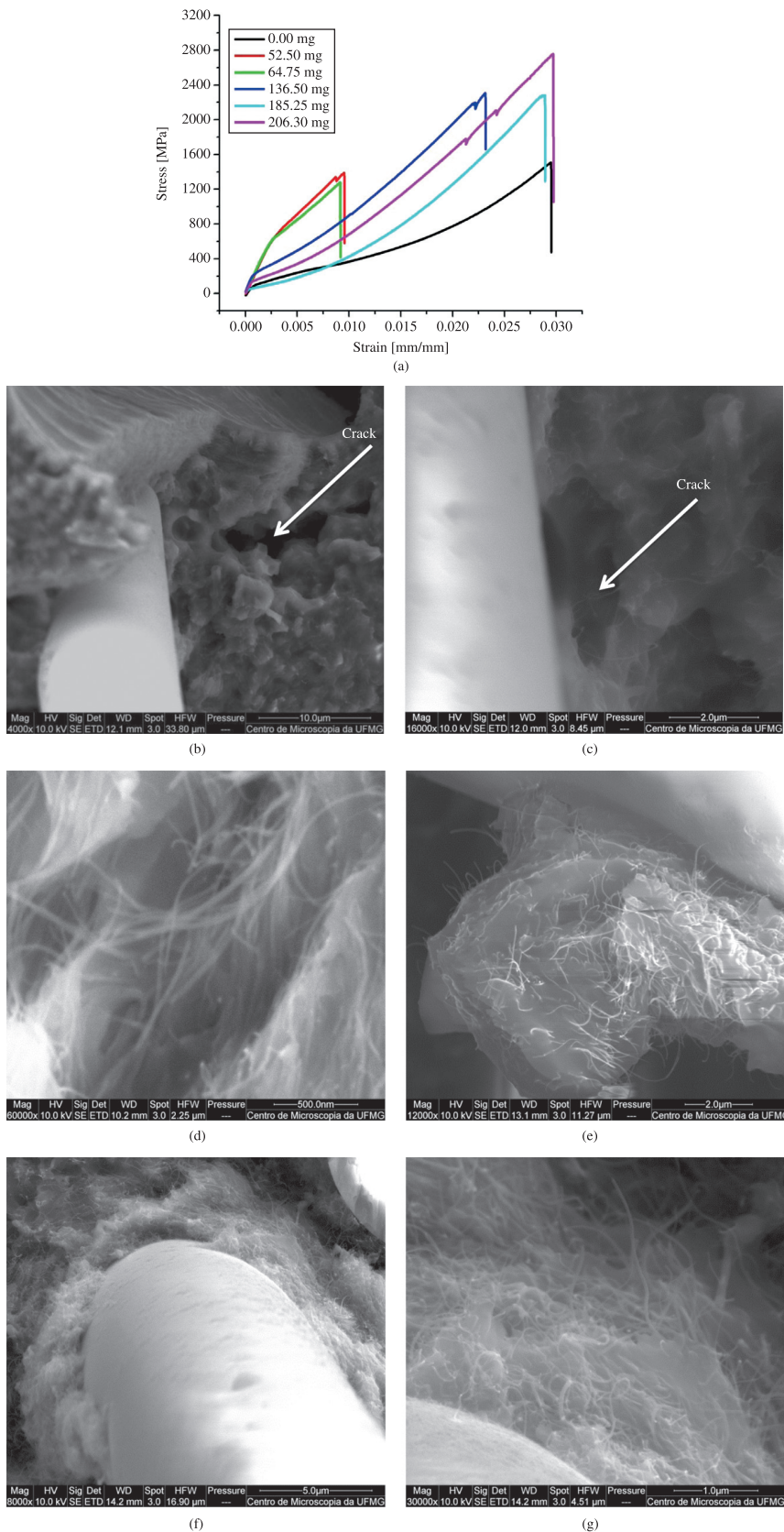


Figure 2. Hybrid laminated composite with CNT inter-layer. (a) Stress-strain curves; (b) Fiber/CNTs for 52.50 mg; (c) Fiber/CNTs for 64.75.50 mg; (d-e) Fiber/CNTs for 136.50 mg; (f) Fiber/CNTs for 185.25 mg; (g) Fiber/CNTs for 206.30 mg.

composite with CNTs as interlayer. As it can be observed in Figure 1a, for CNTs (the lower curve), three bands can be easily identified. The D band is around 1340 cm^{-1} , while the G band is located around 1580 cm^{-1} and the G' band is nearby 2685 cm^{-1} . As described by Malard et al.³⁷ and Dresselhaus et al.³⁵ these peak values are typical from carbon based materials. Furthermore, the narrow and intense D band could be associated to the small nanotube diameter ($\approx 20\text{ nm}$) and the high intensity can be related to CNTs length. Moreover, the narrow G band is related to the CNT's crystallinity, which is confirmed by the XRD signature shown in Figure 1b. By analyzing the second curve (fibers + CNTs), it is possible to observe the carbon fibers' amorphous behavior represented by the smooth curve. However, the carbon nanotubes G band is also observed superposed to the smooth curve. The G band is indicated by the arrow and a region's zoom is shown on Figure 1a upper right corner. Finally, the last curve (upper curve) indicates the fiber glass amorphous behavior.

To understand the how the interlayer nanotubes impact the laminated composites' macroscopic behavior under tension five different CNTs forests were grown into the fiber glass cloths. The amount of CNTs grown in each fiber glass interlayer can be described as 52.50 mg, 64.75 mg, 136.50 mg, 185.25 mg, and 206.30 mg, respectively.

Figure 2a shows the stress-strain curves for the hybrid laminated composite. As it can be noticed as we are dealing with a RVE (a thin laminate), the stresses are "naturally" high. Moreover, changes on peak stress and stiffness are observed. The peak stress changes seem to be attributed to the failure modes. As it can be noticed in Figure 2b cracks between fibers and also around fibers were observed. CNTs, however, are everywhere and this distribution lead to bridges between cracks (see CNTs at Figure 2c) and consequently and increase on peak stress. As the failure mechanism is the same for both cases, the peak stress for CNTs concentrations of 52.50 and 64.75 mg are statistically similar. An increase on CNTs concentration to 136.50 mg and 185.25 mg lead to another level of peak stress, i.e. from $\approx 1350\text{ MPa}$ to $\approx 2300\text{ MPa}$. Again the failure mechanism seems to be the same for both cases (136.50 mg and 185.25 mg). No cracks are noted between fibers, but cracks were spotted on fibers and inside the CNT/matrix mix (Figures 2d, e). Finally, as expected, the increase on CNTs concentration to 206.30 mg leads to the highest peak stress ($\approx 2800\text{ MPa}$). This

performance can be attributed to the strong bond between fibers and CNTs as it can be observed in Figures 2g, h.

In summary, when peak stresses are considered, it is possible to conclude that the addition of the CNT interlayer lead to an increase close to 85%. However, changes on stiffness during the tensile test indicate changes on nano/micro-structures due to damage. The increase on stiffness from 33.99 GPa (from samples with no CNTs) to 235.13 GPa (from samples with CNT concentrations of 52.50 mg and 64.75 mg) was not only due to the CNT concentration but it is also related to the CNTs distribution. As cited by Thostenson et al.³⁸, an increase on stiffness is associated to a decrease on displacement and strain, and vice-versa (as shown in Figure 2a).

4. Conclusions

CNT interlayers were successfully introduced to carbon fibers/epoxy laminated composites. An increase on peak stress close to 85% was witnessed when CNTs interlayer with 206.30 mg was placed in carbon fiber/epoxy laminates. The failure mechanisms are associated to CNTs distribution between and around carbon fibers. These CNTs are also accountable for crack bridging formation and the increase on peak stress. The initial stiffness is strongly affected by the CNT interlayer, however, changes on stiffness is related to changes on nano/micro-structure due to damage. Cracks between fibers were observed in hybrid composites with interlayers made of 52.50 mg and 64.75 mg. When the amount of nanotubes is increased to 136.50 mg and 185.50 mg, no cracks are noted between fibers, but cracks were spotted on fibers and inside the CNT/matrix mix. Finally, when the CNT amount reached 206.30 mg, the main failure mode was fiber breaking, as a strong bond between fiber/CNTs was observed by SEM.

Acknowledgements

This research was supported in part by the AFOSR under contract FA9550-10-1-0050, the Brazilian Research Council (CNPq) under grants number 303447/2011-7 and 472583/2011-5, and the Minas Gerais State Research Foundation (FAPEMIG) grant TEC-PPM00192-12. The authors would like to recognize the technical support provided by the Nanomaterial's Laboratory from the Physics department of Universidade Federal de Minas Gerais where the CNT were in situ grown on the surface of glass fibers.

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