Ultra-thin Electrospun Nano-fibres for Development of Damage Tolerant Composite Laminates

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Abstract

The present article overcomes existing challenges ahead of inter-laminar toughening of novel multifunctional fibre-reinforced polymer composites via development and embedment of highly stretched, ultra-thin electrospun thermoplastic nanofibers made of polyamide 6.6. The nanofibers have exhibited significant enhancement of the composite laminate's structural integrity with almost zero weight penalty via ensuring a smooth stress transfer throughout the plies and serving tailoring mechanical properties in desired directions, with no interference with geometric features e.g. thickness. The findings for 1.5 grams per square meter (gsm) electrospun nanofibers have demonstrated, on test coupons specimens, improvements up to 85% and 43% in peak load and crack opening displacement, respectively, with significant improvement (> 25%) and no sacrifice of fracture toughness at both initiation and propagation phases. The initial stiffness for the modified specimens was improved by nearly 150%. The enhancement is mainly due to nano-fibres contributing to the stiffness of the resin rich area at the crack tip adjacent to the Polytetrafluoroethylene (PTFE) film. Glass fibre-reinforced woven phenolic preimpregnated composite plies have been modified with the nano-fibres (each layer having an average thickness of <1 micron) at 0.5, 1.0, 1.5, 2.0 and 4.0 gsm, electrospun at room temperature on each ply, and manufactured via autoclave vacuum bagging process. Inter-laminar fracture toughness specimens were manufactured for Mode I (double cantilever beam, DCB) fracture tests. It was found that there is threshold for electrospun nanofibers density, at which an optimum performance is reached in modified composite laminates. The threshold is influenced by the plastic deformation mechanism at the crack tip, the fibre bridging between the adjacent plies afforded by the nano-fibres and the density of the electrospun fibres. Such optimum performance was found linked to the nanofibers at a specific density. Excessively increasing above the threshold (herein > 2.0 gsm) degrades the adhesion properties (chemical bonding) between glass fibres and phenolic. The density of nano-fibres increases, so does the likelihood of forming a physical barrier between the plies resulting in the loss of resin flow and poor adhesion. Such effect was evident from microscopic investigations and reduction in fracture toughness data at initiation and propagation phases.

Keywords: electrospun nano-fibers, polymer-matrix composite, fracture toughness, chemical bonding, nanofiber deformation

1. Introduction

Fibre-reinforced polymer composite materials are extensively used in aerospace and general transport due to their high mechanical properties, low density and tailorability to align with a broad and diverse range of applications. Polymer matrix composites (PMCs) present excellent specific stiffness and strength which results in dramatic weight reduction in design. The use of PMCs on transport structures enables substantial fuel efficiency, and therefore have direct impact on CO₂ emission reduction However, one of the principal limitation of PMCs is the low damage tolerance of these materials and consequently a short lifetime. In particular, delamination growth between reinforcing plies in PMC laminates is considered one of the most predominant and life-limiting type of damage encountered in composite laminates during service that can be detrimental to flight safety [1-6]. Furthermore, while impact or dynamic events can significantly reduce mechanical properties, impact induced delamination damage is hardly detectable by the existing non-destructive inspection (NDI) techniques [7-9]. 87% of total composite damage is caused by impact with energies ranging from 5 to 100 joules, known as lowvelocity (energy) impacts [4-6, 10]. In such terms, several different methods have been developed to toughen composite materials to resist inter-laminar delamination [3, 11]. These delamination mechanisms are often driven by instantaneous development of interfacial cracks between two neighbouring plies. The crack propagation may be quantified by the strain energy release rate (G), which is the amount of energy needed to create a crack surface (J/m²), a.k.a. the delamination toughness between two plies, therefore, a function of crack face initiation and propagation. Its value can vary depending on the crack length and strengthening mechanisms such as adhesion between the plies,

between reinforcements and polymers and fibre bridging [3, 11] where reinforcements (e.g. carbon, glass or polymeric fibres) bridge across the faces of the neighbouring plies. Preventing delamination occurrence has to be taken into account during the design and conception of composite laminates, sources of out-of-plane stressing often being caused by load-path discontinuity. Another source is manufacturing induced defects as well as impact induced damage that cannot be entirely avoided [12-16] such as kissing bonds representative of pre-existing poor adhesion properties. Thus, increasing the intrinsic fracture toughness of composite laminates is crucial. Interleaving techniques have been developed via inserting inter-laminar layers, for example, poly-ether-imide veils [17], for improving toughness between the original composite plies. This method is effective as it enables impact damage absorption and hinders the damage at its initial stage, however these inserted layers are also relatively heavy and thick (almost the same thickness as the original composite ply, e.g. ~250 microns), and, therefore, reduce the specific stiffness and specific strength of the laminates [17].

When many successive plies have the same orientation, the remaining mismatched interfaces will suffer from extensive damage, and the laminate damage resistance is reduced, while increasing the number of dissimilar interfaces increase the energy absorbed during delamination [18]. That means that thicker plies would have less interfaces for the same structure's width that leads to relatively low strain energy absorption during delamination.

Interleaving techniques have shown encouraging results in terms of the laminate's fracture toughness enhancement [19-27]. To tackle its main drawback, the weight penalty in addition to thickness and geometry interference, researchers have studied the possibility to introduce a light layer of thermoplastic electrospun nanofibers which often enable multi-functionality such that in polyvinylidene fluoride (PVDF) materials e.g. enabling piezo-electricity and energy harvesting [19-27]. Electrospinning is an established and scalable technique to generate continuous fibres in nanoscale, with a broad range of constituents. The fibres are produced from a polymeric solution, stretched by the electrostatic repulsion between surface charges at relatively high voltages and the evaporation of solvent [28-30]. In particular, interfacial toughening based on thermoplastic nano-fibres electrospun from thermoplastics is of an increasing interest [31-41]. The advantage of this approach is that there is almost zero thickness or weight penalty (< 0.2%), since the nano-fibrous layer is in the order of 100nm thin, and is highly localised between the plies, leading to only marginal loss of in-plane properties. The nanofiber fabrication is cost efficient and rapid, and the existing composite processes do not have to be changed to meet the requirement of such toughening technique. Though the toughening process is efficient and straightforward, there is a research gap to

address the drawbacks of not having sufficient adhesion properties between PMC plies and the thermoplastic nanofibers. If the electrospinning density is not controlled properly, the modified laminate may re-possess interlayer defects due to the presence of nanofibers rather than chemical bonding, ductility and plasticity improvement between plies.

The aim of this study is to develop a modified PMC laminate with zero weight and thickness penalty, and quantify the effect of embedment of nanofibers on the fracture toughness of composite laminates made of glass fibre-reinforced PMC composites, at different nanofiber layer thickness and aerial density; gsm.

2. Materials and Manufacturing

The PMC pre-pregs used in this study were phenolic based glass fibre-reinforced composite laminates made of 16 plies of intermediate modulus, 0.25mm-thickness woven glass fibre composite pre-pregs having density of 2039 kg/m³, manufactured by BRP Composites Ltd. In a quasi-isotropic, symmetric stacking sequence ([45/0/90/45/0/90/45/0]s), the composite exhibited Young's modulus of E_{11} =22.9GPa, E_{22} =21.6GPa and E_{33} = 3.8GPa, respectively in longitudinal, lateral and through-thickness directions, and failing at approximately 2% glass fibre strain based on our primary investigations [21].

The inter-laminar toughening nanofibers were thermoplastic nylon 6,6 produced by high voltage, room temperature electrospinning technique at Munro Technology Limited. The process was capable of electrospun stretching of the polymer up to 500%. There is no additional mechanical stretching involved in the electrospinning, therefore the nano-fibres stretch as a consequence of the electrospinning process.

Nylon 6,6 powder sample exhibited a homogeneous sphere size with average diameter of approximately 100 μ m, as shown in Figure 1(a). The powders' electrospun nanofibers up to 300% stretching leads to the nano-fibre bundle shown in Figure 1(b). Comprising of nanofibers with diameters ranging from 400 nm to 3 microns. Further stretching (up to 500%) led to much smaller diameters (~100nm) and more consistent range of diameters, as shown in Figure 2.

The composition of the powder and nanofibers was obtained using energy dispersive Xray spectroscopy (EDS), and is tabulated in Table 1 taken from the surface of four randomly selected nanofibers.

The EDS data – performed multiple times per sample to generate a robust statistical distribution – for the powder and electrospun nanofiber bundles were compared. Results

from the powder indicated a carbon: fluorine ratio of between 4:1 and 2:1, indicating significant sample carbon-based contamination, as the expected result was a 1:1 ratio of carbon to fluorine (PVDF unit cell is -C2F2H2-). The contamination is attributed to carbon arising from manufacturing process, manipulation and detection of the carbon tape on which sample is stuck for being analysed. As no sample exceeded a fluorine ratio of 50%, it can be assumed that this is an accurate assumption. No peak that could be attributed to any other atomic element than carbon, fluorine or gold (coating) was identified. In particular, no Nitrogen was observed in any of the spectra of the prepared fibres, indicating that a negligible amount of solvent was present.

Fourier Transform Infra-Red Spectroscopy (FTIR) was carried out on the powder and the electrospun nanofiber bundles at different stretching up to 500%. Figure 3 compares the FTIR ATR ZnSe spectrums for the samples with most of the peaks are between 550cm⁻¹ and 1450cm⁻¹ (Below 550cm⁻¹ the quality of the measurement was too dependent on the setup device, and therefore makes it unreliable, and above 1450cm⁻¹, all spectrums were flat with limited peaks.).

It can be observed that the trend in the appearance of the peaks is almost identical for the spectrums. However, the powder spectrum exhibits slightly different spectrum to the others, with some peak shifts (at 870, 1070 or 1180 cm⁻¹) and higher additional peaks at approx. 612, 760, 795, 1207 or 1384 cm⁻¹. This is due to the fact that the powder was the raw material and has not been stretched unlike to the other samples.

Finally, the nano-fibres with the average diameter of 100nm (stretched up to 500%) were directly electrospun on the surface of each ply as shown in Figure 2 to create the modified composite pre-pregs. The pore size distribution of the electrospun mats of different density is a critical parameter affecting the fracture energy and toughening mechanisms. The nanofiber mat thickness was controlled via controlling the speed of electrospinning with respect to the pre-preg surface in a way to freely embed over the surface and avoid further stretching. Thereby, it was insured that the nanofibers mat thickness was nearly identical to the nanofibers average diameter.

Dynamic mechanical analysis (DMA) measurements were also carried out on the pristine and nanofiber modified pre-preg plies in order to identify any possible variation in curing parameters needed for the co-process. The results presented in Figure 4 shows a slight effect of nano-modification on glass temperature with maximum 30% increase in $tan(\delta)$ data.

To prepare the test specimens, the pre-preg lamina was cut into the plies. For the plies oriented at -45° or +45°, panels of 550mm × 424mm dimensions were cut. Plies were carefully stacked and the waxed paper protecting the nanofibers was released in a protective environment just before laying up over the middle ply, in order to avoid

moisture absorption. Post laying-up, the specimens where cured using vacuum bagging in the autoclave under pressure of 6 bar and at 125°C for two hours, with the initial heating rate of 1°C/min. Specimens were then cut to produce Mode I fracture toughness test specimens according to ASTM D5528 [42] with $a_0/W = 50/125 = 0.4$. A nonadhering release film, made of PTFE in thickness of 15 micron, was inserted over the mid-plane plies, on both panels to mimic pre-existing crack length of approx. 50 mm. Thus, the plies adjacent to this insert were not bonded post curing. This way, the insert simulated a stress concentration site to produce fracture test specimens in opening mode, DCB specimens. End blocks were manufactured to adhesively tab the specimens for mode I testing as shown in Figure 5.

3. Experiments

3.1. Fracture toughness specimens and tests

Fracture toughness specimens for Mode I testing (opening mode) followed the ASTM D5528 standard's specifications [42] using the DCB specimens in 20mm × 125mm dimensions, presented in Figure 5 at unloaded and loaded conditions. Edge side release films were embedded onto the DCB specimens over the mid-ply, to create pre-existing crack faces before the materials were co-processed. Four specimens per category were manufactured to examine the repeatability of the test data. Pristine reference specimens were also manufactured for testing to create a baseline for comparative studies. The same manufacturing protocol as that for the nanofiber modified samples was used (outlined in section 2) to manufacture the pristine reference samples using the autoclave under 6 bar pressure, and at 125°C for two hours with the heating rate of 1°C/min.

Fracture toughness tests were performed using a Zwick 10kN force machine equipped with a load cell of 2.0 kN. The crosshead opening displacement speed for all the tests was set to be at or below 1 mm/min to ensure quasi-static loading condition. Great care was taken to avoid initial loading on the pre-existing crack and the nanofiber layers when fitting the load blocks on the jigs; and to verify the pins inserted in the blocks were parallel, in order to avoid torsional moment. Load and displacement data were recorded using the TestXpert v5.01 software on a Desktop computer. In both testing, the delaminated crack length measurement was carried out using continuous visual observations from one side of the specimens, Figure 5 via an optical magnifier. The load levels were automatically recorded against the crack growth data.

Mode I fracture toughness (G_I) was then evaluated based on the modified beam theory (Eq. (1)), to account for end blocks and large deformation conditions as the specimens'

crack were allowed to propagate longer than the minimum required. The distance between the loading pin hole and the front of the tab was 4 mm, and between the hole and the surface of the DCB specimen was approximately 3.5 mm. Such values were taken into account for theoretical corrections of the G_I values given by:

$$G_I = \frac{3P\delta}{2b(a+\Delta)} \times \frac{F}{N} \tag{1}$$

where *P* is the applied load, δ is the load point displacement measured using the machine head's LVDT, *b* is the specimen's width, and *a* is the delamination crack length measured optically. Δ is the correction factor that accounts for the fact that opening DCB is not a perfect cantilever with its end totally fixed. It is determined experimentally by generating a least squared plot of the cube root of compliance, $C^{1/3}$ where $C = \delta/P$, as a function of delamination length. The large deformation factor, *F* in Eq. (1), corrects the fact that as the angle of the end blocks changes during loading, their orientation also changes that influences the distance between the crack front and the loading pin. The use of end blocks, which can cause a stiffening effect of the specimen arms or different displacement relative to the ideal loading point, is corrected by the *N* factor. Values for *F* and *N* are given by:

$$F = 1 - \frac{3}{10} \left(\frac{\delta}{a}\right)^2 - \frac{2}{3} \left(\frac{\delta l_1}{a^2}\right)$$
(2)

$$N = 1 - \left(\frac{L}{a}\right)^3 - \frac{9}{8} \left[1 - \left(\frac{L}{a}\right)^2\right] \left(\frac{\delta l_1}{a^2}\right) - \frac{9}{35} \left(\frac{\delta}{a}\right)^2 \tag{3}$$

where *L* is the specimen effective length (horizontal distance between the load pin's centre and the end side of specimens), and l_1 and l_2 are the distances from centre of loading pin to mid-plane of the specimen and to the edge of the loading blocks, respectively.

The initial fracture toughness in opening mode, $G_{I,ini}$, is calculated based on the Standard's recommendation, and is given by:

$$G_{I,ini} = \frac{3P\delta}{2ba} \tag{4}$$

3.2. Microscopy

Optical microscopy and scanning electron microscopy (SEM) were used to identify damage mechanisms occurred in the specimens post fracture toughness testing. The

specimens for SEM were plasma coated with a thin gold layer, ~15 nm thickness at 20 mA, to improve the imaging of the polymers via discharging surface electrons.

4. Results and Discussion

Raw data of applied force are plotted versus crack mouth opening displacement in Figure 6 for all the modified phenolic composite DCB specimens. The data shown in all sub-figures for the modified specimens (dashed lines) are compared with that of the pristine specimens (solid lines).

All DCB specimens, whether with (modified) and without (pristine) nanofibers, have the same physical dimensions and same initial crack lengths of 50 mm, however the initial stiffness values of the four pristine specimens vary substantially (>50%). Same disparity is seen for the modified specimens with densities of 1.0 gsm and 5.0 gsm, Figure 6(b) and 6(e). Such disparity rises for both pristine and modifies specimens:

The PTFE tape was inserted to simulate a pre-existing crack, and therefore had no influence in driving the initial stiffness, i.e. before crack initiation. The panels from which the specimens were extracted, however, showed different quality post process, in different locations, apparently a sign of non-uniform cure at the centre and the edge sides due to non-uniform heat transfer in the conventional oven. As the panels were sufficiently large, the specimens cut near the edge of the panels are believed to attain different quality, thus a different quality which may have led to the disparity in the initial stiffness data. However, it is acknowledged that non-uniform cure in relatively large panels in actual applications is a common challenge especially in non-flat panels with back structures.

As seen, in comparison with the pristine data, the peak load increases with the increasing nanofiber density from 0.5 to 2.0 gsm (Figure 6(a) to (d)), and decreases for densities > 2.0 gsm (Figure 6(e)), as also observed for the increasing displacement data (x-axis).

Figure 7 suggests that the 1.5 gsm modified composite exhibits a better performance with 85% improvement in the peak load (= $100 \times (48N - 26N)/26N$) whilst having the lowest disparity (scatter in data), a sign of robust reliability based on repeatable test data. Moreover, before any crack initiation the initial stiffness of the modified composite has significantly been improved at 1.5 and 2.0 gsm with nearly 145% and 110% improvements, respectively. Such increase in the opening stiffness before crack is initiated suggests a high modulus compared to that of the pristine specimen within the elastic regime, attributed to high adhesion properties (e.g. interfacial strength) in the opening mode.

The maximum force and opening displacement data extracted from Figure 6, and averaged data of the four specimens are presented in Figure 7. As seen, the peak load for the 1.0 gsm reached a high value of 52N for one specimen whilst the other specimens data suggest slightly lower level than the pristine samples. (Figure 6(b) and 5(a) showing high scattered data as result of such high disparity).

As seen, in Figure 6, the significant stiffening occurs only at 1.5gsm and 2.0gsm (Fig. 6 (c, d)). Taking the 1.5gsm as an example, this is in a consistent trend with the improvements in the peak load by 85% and the maximum displacement by 43%. As the Stiffness is the ratio of force over displacement, the stiffness for 1.5gsm is almost doubled, by ~198% (=85/43). It may be noted that this occurs in the linear elastic regime where mainly deformation mechanisms are in play, rather than damage. The authors have observed that in such elasticity improvement scenarios, two microstructural mechanisms are prevailing for the 1.5gsm and 2.0gsm which are rarely observed in other density levels: adhesion of nanofiber layers to the polymer (and in some cases to fibres) and intact fibre bridging, also consistent with the initiation of crack surfaces which requires 25% higher energy to initiate fracture (such as that in Figure 8(a)), meaning that the energy dissipated initially by the existing deformation mechanisms in the cases of 1.5gsm and 2.0 gsm prior to damage.

The fracture toughness data based on the measurement of crack length in opening mode has been given in Eq. (1) for propagation and Eq. (4) for initiation phases, and are plotted in Figure 8 as a function of the increasing nanofiber density (0.5 gsm to 4.0 gsm). As seen, the fracture response at both initiation and propagation phases have improved for the 1.5 gsm specimens by more than 25% compared to the pristine samples. Such improvement may seem trivial compared to the existing interleaving veils however it should be noted that the 25% improvement is obtained at the zero weight and thickness penalty, and in laboratory-scale specimens. Scale-up analysis for large structural scale components in actual applications requires further investigation. The initiation fracture toughness is slightly higher than the propagation one almost for all cases, due to the nanofibers induced crack tip being blunt shape rather than sharp and straight. The fibre bridging was evident during fracture toughness testing of 1.5 and 2.0 gsm specimens, significantly occurring due to the nylon nano-fibres sticking to the glass fibre and the phenolic matrix. This has contributed to hindering instantaneous separation via introducing plasticity mechanisms surrounding the crack tip edges, as postulated in [15]. Consequently, the G_I value increased during the propagation phase.

Also, it was observed that the nanofiber modified specimens exhibited a more stable crack propagation at 1.5 and 2 gsm, and a more unstable instantaneously occurring crack propagation at densities above 2.0 gsm. New mechanisms induced by the nano-

fibres within the interlayer have also allowed increasing the potential for elastic and plastic yield of the laminate in opening mode during the propagation phase. It may be envisaged that the nanofibers ahead of the tip of a propagating crack effectively dissipate the strain energy via extensive deformation rather than damage, therefore creating plastic zone ahead of the crack edge tip. Such deformation mechanism, in the subsequent loading, creates bridges between the two neighbouring plies, holding the fracture surfaces together and mitigating the creation of new crack faces. The presence of this bridging phenomenon would be a mark of improvement of the resistance to Mode I loads, resulting in greater steps of load increase. In order to observe such phenomenon at microscale, fractography using SEM was carried out to support the link between the nano-fibres and the variation of fracture toughness.

4.1. Microscopy data

To observe a correlation between variation of G_I and nanofiber density, specimens were carefully cut on their crack faces post failure to fit the SEM cavity for microstructural fractography. The dominant mechanism was observed as interfacial fracture between the composite plies visible in Figure 9 for a typical modified specimen. Generally in most of the specimens, it was observed that any force drop throughout the test (see-saw drops in Figure 6) was associated with an instantaneously occurring delamination as opposed to the increase in force levels, due to progressive (gradual) creation of crack faces.

Figure 10 compares the fracture surfaces in pristine and nanofiber modified specimens. A closer look on the glass fibre reinforcement reveals a significant distinction on the surfaces morphology: unlike the surface of the reinforcements in the pristine specimen, the glass fibres in the nanofiber modified specimens exhibit a surface containing numerous pores made of nylon 6, 6. Figure 11 shows the SEM image for glass fibre rich area in a pristine specimen where instantaneously occurred delamination has bared the glass reinforcements off the phenolic matrix. However for the modified specimens having nanofiber density of greater than 1.5 gsm, the nano-fibres form a barrier for the phenolic resin to flow through and adequately wet the glass fibres, so they have resulted in reduction of peak load and fracture toughness (Figures 7 and 8). A low wettability of the nano-fibres by the resin at high densities results in a weak interface and, therefore, in a preferential path for the crack propagation. In addition to the wettability issue, in an analogous way to high glass fibre volume fraction, the proportion of nanofibers areal distribution (e.g. 4 gsm) should be considered more carefully. Nano-fibres have huge surface area, potentially conferring a strong interface. In case of excessive proportion of nano-fibres, the resin becomes soaked up on the fibres, leading to weak bonds between

the plies during the co-process. Consequently, delamination becomes very likely to occur in high nanofiber density as well as in low density. We have then taken this extreme case of 4.0 gsm to investigate the performance reduction:

Figure 12 presents fracture surface images of 5.0 gsm nano-modified specimens that have failed at the lowest load assessed herein with the lowest G_I . The nanofibers are not present in pristine specimens (Figure 11), and thus no residue of matrix remains stuck to the surface of the fibres, making a smooth surface whereas the electrospun, co-processed nano-fibres gather around the fibres and create a spider web shaped structure. Also, the nanofibers at 4.0 gsm are seen having created a porous and deformable structure, however, due to the excess of the nanofibers diffused within the phenolic resin during the co-process and covering the glass fibres, chemical disbond is induced between the fibres and the phenolic matrix, a well-known phenomenon (see e.g. [43]). The porous structure represents nanoscale fibres that have failed in a brittle manner since no reduction of the cross-section was observed. Such phenomenon is more evident in Figure 13 in the magnified nanofibers area. A deformation of the ductile matrix would have given a less regular failure.

Generally, by comparing the matrix/fibre interface in the two laminates (pristine and modified), any new bond in the interfacial area would be directly related to the addition of nanofibers, which at densities above 2.0 gsm would create disbond due to excessive diffusion of nanofibers through the matrix and over the reinforcement surface, thus results in degradation in load carrying capacity and fracture response.

5. Conclusions

The manufacturing process of handling pre-pregs electrospun with thermoplastic nanofibers via room temperature electrospinning was carried out on phenolic preimpregnated glass fibre-reinforced composite plies, followed by a co-process in autoclave and using vacuum bagging. It was observed that though the nanofibers were ultra-thin and low-density, apparent improvement mechanical performance, damage tolerant and fracture toughness can be achieved provided that the density of nanofibers is taken into control. Nano-fibre density levels at 0.5, 1.0, 1.5, 2.0 and 4.0 gsm were examined.

The findings for 1.5 gsm demonstrated the most promising and reliable nanomodification for the proposed laminate in coupon test specimens, via improvements up to 85% and 43% in peak load and crack opening displacement, respectively, with improvement (>25%) of fracture toughness at both initiation and propagation phases, and initial stiffness improvement by nearly 150% in the linear regime prior damage initiation. The stiffening mechanism was observed due to the enhancement of adhesion properties and contribution of deformation mechanisms (plastic zone creation and fibre bridging) at the right density of 1.5 and 2.0 gsm. Excessive electrospun nanofibers (above 2.0 gsm) were found degrading rather than improving the performance, dominantly due to the diffusion of nanofiber onto the glass reinforcements and creation of interfacial defects (disbond). Such conclusion must emphasise that the 1.5 gsm is an optimum nano-fibre density for the phenolic resin. Other matrix such as epoxy and polyesters will require similar approach to determine the optimum nano-fibre loading.

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Figure 1: Comparison of morphology of (a) nylon 6, 6 powder and (b) electrospun nanofibers, stretched up to 300%



5

Figure 2: Nylon 6, 6 nanofibres electrospun, stretched up to 500%, deposited directly over the phenolic composite plies at density of 2.0 gsm and 5.0 gsm



Figure 3: Comparison of FTIR spectrums for the powder and nanofibres, ranged from 550cm⁻¹ to 1450 cm⁻¹



Figure 4: Effect of electrospun nanofibers on the glass transition temperature (DMA data) showing approximately identical temperature of 170°C



Figure 5: Mode-I DCB specimen test setup demonstrating the composite specimen with the white-painted edge marked up at every millimetre for crack propagation studies





Figure 6: Comparison of evolution of driven force with crack mouth opening displacement in pristine (non-modified) and modified glass fibre-reinforced phenolic composites with electrospun thermoplastic nanofibers at densities of (a) 0.5 gsm, (b) 1.0 gsm, (c) 1.5 gsm, (d) 2.0 gsm, and (e) 5.0 gsm



Figure 7: Comparison of pristine and electrospun nanofiber modified glass fibre-reinforced phenolic composites at nanofiber densities of 0.5, 1.0, 1.5, 2.0 and 5.0 gsm: (a) averaged maximum load data, (b) averaged maximum crack mouth opening displacement data



Figure 8: Evolution of fracture toughness with the increasing electrospun nanofiber density; (a) $G_{I,ini}$ at initiation, (b) G_I at propagation stage



20 mm

Figure 9: Crack open DCB specimen laps post failure



Figure 10: Comparison of fracture surfaces comprising of glass fibre reinforcements and phenolic matrix in (a) pristine specimen; and (b) nanofiber modified specimen



25 µm

Figure 11: SEM image of typical DCB pristine specimen at ultimate failure in glass fibres area (Fig. 10(a))



Figure 12: SEM images of typical DCB nanofiber modified specimen (5.0 gsm herein) at ultimate failure in glass fibres area (Fig. 10(b))



5 µm

2.5 µm





Figure 13: SEM images showing the presence of deformed and broken nanoscale structures at ultimate failure from different locations, appearing to be electrospun nanofibers at diameters of 100-200nm

Spectrum Label	C (%)	N (%)	O (%)	F (%)	AI (%)	Si (%)	Ca (%)
Spectrum 1	28.77	3.85	36.74	0.04	4.4	12.59	13.61
Spectrum 2	41.23	9.21	25.03	0	1.03	19.88	3.62
Spectrum 3	35.65	3.34	26.66	0.35	1.21	29.7	3.09
Spectrum 4	50.2	12.48	21.17	0.5	0.32	14.59	0.74

Table 1: EDS data from surfaces of electrospun nanofibers