Research Article

Graphene Nanoplatelets in Epoxy System: Dispersion, Reaggregation, and Mechanical Properties of Nanocomposites

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The use of graphene nanocomposites in advanced applications has attracted much attention in recent years. However, in order to substitute traditional epoxy reinforcements with graphene, there are still some issues like dispersion, homogenization, and reaggregation. In this paper, graphene bundles dispersed in two-component epoxy system by bath sonication, dispersion state, and reaggregation behavior of graphene in this system have been studied. Light transmittance in ultraviolet-visible spectroscopy has been used to quantify the reaggregation by a series of controlled experiments. After 18 mins sonication of 0.005 wt% graphene dispersion at 20°C, the light transmittance decreased from 68.92% to 54.88% in liquid epoxy and decreased from 72.80% to 46.42% in hardener; while increasing the temperature from 20°C to 60°C, the light transmittance in liquid epoxy decreased from 65.96% to 53.21% after 6 mins sonication. With the incorporation of 0.3 wt% graphene, the tensile strength of nanocomposites increased from 57.2 MPa to 64.4 MPa and the storage modulus increased from 1.66 GPa to 2.16 GPa. The results showed that the dispersion state depends on the function of sonication time and temperature, and graphene has a significant reinforcement effect on epoxy.

1. Introduction

Since its discovery in 2004 [1], graphene has rapidly attracted both academic interest and industrial interest because of its outstanding properties such as high surface to volume ratio, high aspect ratio, low electrical resistivity, high thermal conductivity, high strength, and modulus. Figure 1 [2] shows graphene as the building block of all graphitic carbon allotropes with different dimensionalities. For composites, graphene can significantly improve the physical and chemical properties of matrix at extremely small loadings when incorporated appropriately [3–5].

Epoxy is one of the most adaptable and universal high performance thermosets available and has been widely used in aerospace, automotive, marine, construction, electrical, and electronic systems, biomedical devices, adhesives, paints and coatings, and other general consumer products [6, 7]. The combination of graphene and epoxy has led to a new class of nanocomposites for advanced engineering applications. This has now become an avid area of scientific research as evident in Figure 2. Figure 2 shows the dramatic increase in epoxy/graphene nanocomposites research during the last decade.

However, in practical terms, graphene is not suitable to disperse in epoxy just by simple mixing. This is due to graphene’s pronounced tendency to reaggregate in the matrix due to the strong van der Waals force and pi-stacking between separately dispersed graphene sheets [8, 9]. The maximum improvements in final properties could only be achieved when graphene is homogeneously dispersed in the matrix and external stresses are efficiently transferred through strong graphene-epoxy interface [10, 11]. This can also be seen for other polymer nanocomposites where it is critical to achieve homogenization and thorough dispersion. Therefore, the dispersion state of graphene in the matrix plays a crucial role for achieving superior properties in epoxy/graphene nanocomposites [12, 13].

According to the published research, processing methods significantly affect dispersion of graphene, and it is generally agreed that mechanical stirring is one of the suitable techniques to disperse graphene. However, Yang et al. [14] reported that mechanical stirring of suspended graphene
Appropriate covalent functionalization of graphene with chemicals retards reaggregation [19]. Appropriately surface modified graphene shows rich solution chemistry, with properties such as dispersibility in solvents and polymers [20]. However, the main drawback of covalent functionalization is the functional groups on the graphene surface which disrupt the band structure and degrade the electronic properties of graphene. Wajid et al. [21] even reported that some surface functionalized graphene acts as insulator due to stronger contribution of the attached chemical groups.

Stable dispersion can also be achieved by wrapping graphene surface by stabilizers such as polymers or surfactants [22, 23]. As compared to the covalent functionalization of graphene sheets, this kind of noncovalent surface modification can disperse graphene without oxidation or defect formation, thus preserving the integrity of the structure and electronic properties [24]. However, it is generally recognized that the presence of foreign stabilizers is undesirable for most applications because they could decrease the purity of graphene, subsequently deteriorating the unique and exceptional properties of graphene [25, 26].

could result in reaggregation between flakes, especially for toluene or benzene suspensions. As a widely adopted technique, sonication proved to be effective to disperse graphene. Several researchers [15–18] reported sonication to be an effective way to suppress the reaggregation of graphene.
Using solvents has been widely accepted and regarded as the simplest method to disperse graphene. Choi et al. [27] dispersed graphene in 1-propanol and showed long-term stability of the solution. Other groups dispersed graphene in acetone [28–30], THF [31–33], and DMF [34–36]. However, it should be noted that such dispersions require another step to remove the solvents prior to curing. This can lead to operational complexities and lengthy processing durations which are not desirable. The residual solvent also has detrimental effect on the final properties of the composite material [37, 38]. Beside that, some solvents are expensive and toxic, which may lead to environmental and health and safety implications [39–41].

In this paper, pristine graphene was dispersed in two-component epoxy system without using any solvent. The effect of sonication time, storage time, graphene concentration, and sonication temperature on the dispersion and reaggregation of graphene in epoxy, hardener, and their mixtures have been extensively analyzed here.

2. Experimental Details

2.1. Materials

2.1.1. Epoxy Matrix System. The epoxy matrix used in this study consists of EPOPHEN EL5 bisphenol A based liquid epoxy (EP) and EPOPHENEHA 57 diamine hardener (HD), purchased from Polyfibre UK Ltd. This epoxy system is a multipurpose resin offering good all-round properties with the epoxy group content of 4.76–5.25 mol/kg. The viscosities of liquid epoxy and hardener are 12000–15000 cps and 45 cps, respectively, at room temperature. To prepare epoxy material (EP + HD), the mix proportions are 50 parts by weight of liquid epoxy to 100 parts by weight of liquid epoxy.

2.1.2. Graphene. Graphene was purchased from Graphene Laboratories Inc., USA (product name: AO-3). The graphene nanoplatelets, according to the manufacturer’s technical data sheet, have a specific surface area of 80 m²/g; the average lateral size and thickness are 4.5 μm and 12 nm, respectively. Figure 3 shows the SEM image of a raw graphene nanoplatelet.

2.2. Sample Preparation. Graphene samples were weighed in Sartorius MC210S analytical balance (with the readability of 0.01 mg) and dispersed in EP by hand mixing for 5 s gently and then sonicated through a bath sonicator (Grant MXB6) for uniform dispersion. The bath sonicator was rated for an average working power output of 89 W in aqueous media.

For studying the influence of sonication time on the dispersibility, 0.005 wt% dispersions were sonicated for different durations ranging from 6 mins to 60 mins at 20°C. Another part of dispersion was sonicated for 30 mins and then stored for 10 days for studying the reaggregation against storage time. For studying the influence of concentration on the dispersibility, different concentrations ranging from 0.005 wt% to 0.1 wt% of samples had been prepared and then sonicated for 30 mins at 20°C. For studying the influence of sonication temperature on the dispersibility, 0.005 wt% dispersions were sonicated for 6 mins from 20°C to 60°C. Graphene-HD, graphene-EP + HD dispersions were prepared by the same method accordingly. All samples had been degassed at −0.1 MPa to remove the entrapped air.

To prepare epoxy/graphene nanocomposites, graphene was first dispersed in hardener by bath sonication for 30 mins at room temperature. Then the suspensions were mixed with liquid epoxy by the ratio of EP : HD of 2 : 1. Following thorough hand mixing for 10 mins, vacuum degassing was carried out to remove the entrapped air. The mixtures were then mould cast and cured at room temperature for 6 h followed by postcuring at 80°C for 6 h.

2.3. Characterization. Light transmittance in the UV-visible spectroscopy (HITACHI U-3000) has been used to quantify the reaggregation of graphene in epoxy system through a series of controlled experiments. Tests were always carried out immediately after the sonication of each dispersion. Standard polystyrene cuvettes with an optical path length of 10 mm were used for transmittance measurements. Light transmittance of graphene dispersions was recorded versus blank EP, HD, and EP + HD at fixed wavelength of 450 nm. Five specimens were tested for each set of conditions and mean values were then reported. Graphene dispersions were also systematically characterized by optical microscopy.

Tensile, three-point bend tests were conducted by Universal Testing Machine (Instron 3382); the crosshead speed was kept at 2 mm/min for all the tests. Tensile properties were measured according to ASTM D638 (Type V geometry) with specimen thickness 4 mm. Three-point bend test was conducted according to ASTM D790 with specimen dimensions of 3 × 12.7 × 48 mm. Five specimens were tested for each set of conditions and mean values were then reported.

Dynamic Mechanical Analyser (DMA) (Model 8000, Perkin Elmer) was used to determine the storage modulus (E’), and loss factor tan δ. Rectangular specimens with dimensions of 2.5 × 8 × 30 mm were tested in single cantilever mode. All tests were carried out by temperature sweep method (temperature ramp from 30 to 180°C at 10°C/min) at a constant frequency of 1 Hz.
3. Results and Discussion

3.1. Reaggregation as a Function of Sonication Time. Sonication is the most widely adopted method to disperse graphene in liquid matrix and has proved to be of high efficiency. Figure 4 shows graphene dispersion in EP and HD before and after sonication.

The light transmittance of graphene dispersion against sonication time is shown in Figure 5. The graphs show a significant drop in the transmittance for the graphene dispersion in HD within the first 12 mins. This high magnitude slope suggests a much higher tendency of graphene to disperse in HD. Before sonication, graphene aggregates lowered light absorbance because of the shielding effects of the bundles [42]. After sonication, the aggregates are dispersed into small aggregates/flakes causing higher light absorption or lower light transmittance. A similar trend was also observed for the graphene dispersion in EP, where light transmittance also decreased with the sonication time. However, the magnitude of the slope is much lower than that of HD due to the high viscosity of EP, making it more difficult for graphene to disperse. It is noteworthy that, as compared to 15 percent drop in EP and 26 percent drop in HD, there is just 11 percent drop.
Figure 6: Optical microscopic analyses: (a) graphene-EP before sonication; (b) graphene-HD before sonication; (c) graphene-EP after sonication; and (d) graphene-HD after sonication.

Figure 7 shows the light transmittance of graphene dispersion against storage time. Within the first 5 days, the behavior of graphene in EP and that in HD are similar. Both dispersions showed slight increment in light transmittance, 5% and 4% in EP and HD, respectively. This increment indicated that some level of reaggregation took place during this time, but very slightly. During 5 to 10 days, the light transmittance did not change, indicating that the dispersions are stable over this time period. The light transmittance of graphene-EP + HD dispersion kept constant, because the system became stable after the epoxy resin was fully cured within 24 hours.

Optical microscopy further confirmed the stability of the dispersion. Figures 8(a) and 8(b) show the graphene dispersion in EP and HD tested within 1 min after sonication and Figures 8(c) and 8(d) show the dispersion after 10-day storage. It can be seen that there is no obvious change in the dispersion state, which indicates that the dispersions are in general stable during this time period.

3.3. Reaggregation as a Function of Graphene Concentration. Five series of graphene dispersion with concentrations between 0.005% and 0.1% were prepared. Figure 9 shows the
changes in transparency of graphene dispersions at different concentrations. As can be noticed from the image, suspensions with higher concentration have less light transmission as compared to low concentration suspensions. Samples with concentration higher than 0.025 wt% are visually all black with none light transmission. It should be noted here that though this is a qualitative analysis, it yields some very meaningful findings for further research work.

As mentioned in Section 3.1, at low concentration, the light transmittance decreased with the decrease of agglomerate/flake size. Figure 10 shows the measured light transmittance against concentration. The light transmittance decreased with the increase of concentration.

Figure II depicts the optical photograph of graphene dispersion in EP and HD with increased concentration after sonicating for 30 mins. We can see that graphene was dispersed in the dispersion because of the sonication. Reaggregation behavior cannot be seen directly here. Because the interparticle distance between dispersed graphene sheets is small at higher concentration, making graphene sheets easier to attract each other, it can be deduced that reaggregation behavior can be more pronounced at higher concentrations. Beside reaggregation, higher graphene concentration also means increased difficulty for uniform dispersion [43], which further hinders the stability of the dispersion.

3.4. Reaggregation as a Function of Sonication Temperature. High temperature accelerates chemical reactions and mobility of molecules in liquid system, which would make graphene nanoplatelets easier to disperse. Five series of samples were prepared with sonications at 20 to 60°C and sonicated for 6 mins. The results are shown in Figure 12. For graphene-HD dispersion, the light transmittance is 60.32% at 20°C after 6 mins sonications; however, it reaches 46.42% at 50°C within 6 mins. This value could only be achieved after 18 mins sonications at 20°C as can be seen in Figure 5 (Section 3.1). Similarly, the light transmittance of graphene-EP dispersion was 53.21% at 60°C after 6 mins sonications, which was only achieved after 24 mins at 20°C (Figure 5). For graphene-EP + HD dispersion, longer sonication duration will lead to the curing of the resin, which would hinder any dispersion. Under higher temperature, sonication at 50°C for 6 mins is enough to reach uniform dispersion state as confirmed in Figure 12.

These results confirm that dispersion is strongly dependent on the sonication temperature. Theoretically, appropriate dispersion is achieved by providing the right energy to the system, which is normally accomplished by sonication. However, for fine powders or strongly bonded aggregates, higher temperatures are preferred for increased mobility of chemical species and effective debundling of agglomerates.

3.5. Mechanical Properties as a Function of Concentration in Nanocomposites. The degree of dispersion correlates directly with the mechanical properties of nanocomposites; however, the reinforcement effect at low graphene concentration might be tiny. For analysing the effect of dispersion on the mechanical properties, nanocomposites with higher loading were prepared, at graphene concentrations of 0.1 wt%, 0.3 wt%, 0.5 wt%, and 1 wt%, respectively.

The storage modulus ($E'$) and tan δ values with respect to temperature were studied to assess the role of the nanocomposites under stress (Figure 13). As summarized in Table 1, the storage modulus of the nanocomposites at room temperature increased with increasing graphene concentration up to 0.3 wt% (Figure 13(a)). The nanocomposites with a graphene concentration of 0.3 wt% show the highest storage modulus of $E' = 2.16$ GPa, while the pure epoxy shows $E' = 1.66$ GPa; that is, 30% increment. The value reached maximum at the loading of 0.3 wt%. This was due to the homogeneous dispersion being limited up to the graphene concentration of 0.3 wt%.
Figure 8: Optical microscopic analyses: (a) graphene-EP storage for 1 min; (b) graphene-HD storage for 10 days; (c) graphene-EP storage for 1 min; and (d) graphene-HD storage for 10 days.

Figure 9: Qualitative analyses of (a) graphene-EP and (b) graphene-HD (concentration from left to right: 0.005 wt%; 0.0125 wt%; 0.025 wt%; 0.05 wt%; 0.1 wt%).

Figure 13(b) shows the temperature dependence of epoxy/graphene nanocomposites’ tanδ values. The tanδ value is defined as loss factor, which is the ratio of loss modulus to storage modulus (\(\frac{E''}{E'}\)), and is very sensitive to structural transformation of solid materials. The maximum peak of tanδ represents the glass transition temperature \(T_g\) of material and is shown in Table 1. The \(T_g\) value of nanocomposites shifts to higher temperature, from 93.37°C (pure epoxy) to 99.08°C (graphene concentration 0.3 wt%). The rise in \(T_g\) is associated with restriction in molecular motion and reduction in free volume due to the uniform dispersion of fillers. This is responsible for decreasing the mobility of the polymer chains in response to an applied thermal energy. For concentrations above 0.5 wt%, a drop
The incorporation of graphene also significantly influenced the tensile and flexural properties of nanocomposites, as shown in Figure 14. The figure shows the average tensile strength for pure epoxy is 57.2 MPa, and the tensile strength for nanocomposites with 0.3 wt% graphene loading is 64.4 MPa, which is an increment of 12.6%. The flexural strength also shows a 10% increment at graphene loading of 0.3 wt%. These remarkable increments in the tensile and flexural strengths were due to the uniform dispersion of graphene in the epoxy matrix, which formed a perfect cocontinuous structure. Further increasing the graphene loading above 0.5 wt% led to a decrease in both the tensile and flexural strengths. This is due to the nonuniform dispersion of graphene and the aggregates may have acted as the defect center; thus the increased stress concentration led to the decreasing of the properties. As a result, it was suggested that graphene can be effectively dispersed in epoxy matrix.

Table 1: Thermal mechanical properties of epoxy/graphene nanocomposites.

<table>
<thead>
<tr>
<th>Graphene concentration (wt%)</th>
<th>Storage modulus (GPa)</th>
<th>$T_g$ (°C)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>1.66</td>
<td>93.37</td>
</tr>
<tr>
<td>0.1</td>
<td>2.06</td>
<td>97.15</td>
</tr>
<tr>
<td>0.3</td>
<td>2.16</td>
<td>99.08</td>
</tr>
<tr>
<td>0.5</td>
<td>1.96</td>
<td>98.35</td>
</tr>
<tr>
<td>1</td>
<td>1.86</td>
<td>97.12</td>
</tr>
</tbody>
</table>

in $T_g$ was recorded. This is due to the presence of larger aggregation sites within the nanocomposites as reported in the literature [44, 45]. The aggregates are physical defects in the material, which hinder the curing and lead to the decrease in crosslinking density [46, 47].
when the loading is lower than 0.3 wt%, which is consistent with the DMA test. While increasing the concentration of graphene above this loading, the restacking of graphene leads to the aggregates, saturating the reinforcing efficiency of the nanocomposites. However, in the case of tensile and flexural moduli, the value reached maximum at 1 wt% of graphene loading. This continuous increment shows that, with the incorporation of rigid fillers, the epoxy/graphene nanocomposites are resilient towards elastic deformation. The results (i.e., changes of modulus and strength) are in accordance with other reports [31, 48–50]. Overall, as compared to pure epoxy, both the tensile and flexural properties of the nanocomposites were improved. These increments suggest strong reinforcement effect of graphene in epoxy matrix.

4. Conclusion

Dispersion of graphene in matrix plays crucial role for the performance of nanocomposites. Reaggregation, that is, agglomeration with the passage of time, has often been underestimated and even ignored. This work has measured the reaggregation of graphene in two-component epoxy
system using optical transmittance spectroscopy. The results show that temperature and viscosity significantly contribute to the dispersion of graphene. Graphene tends to disperse easily under high temperature in low viscosity system; beside this, lower concentration yielded lower reaggregation profile (size and trend) and vice versa. The mechanical properties of the nanocomposites were significantly improved as compared to pure epoxy with the incorporation of graphene, which suggested that uniform dispersion plays a critical role.

Conflict of Interests
The authors declare that there is no conflict of interests regarding the publication of this paper.

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References


