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Controlled Structure Evolution of Graphene Networks in Polymer Composites

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Supporting Information

ABSTRACT: Exploiting graphene’s exceptional physical properties in polymer composites is a significant challenge because of the difficulty in controlling the graphene conformation and dispersion. Reliable processing of graphene polymer composites with uniform and consistent properties can therefore be difficult to achieve. We demonstrate distinctive regimes in morphology and nanocomposite properties, achievable through systematic control of shear rate and shear history. Remarkable changes in electrical impedance unique to composites of graphene nanoplatelets (GNPs) are observed. Low shear rates ≤0.1 s⁻¹ break up the typical GNP agglomerates found in graphene composites, partially exfoliate the GNPs to few-layer graphene, and reduce orientation, enhancing electrical conductivity in the composite materials, whereas at higher shear rates GNP orientation increases and the conductivity reduces by four orders of magnitude, as the graphene filler network is broken down. The structure of the composite continues to evolve, reflected in further changes in conductivity, after the shear force has been removed and the process temperature maintained. This work provides critical insights for understanding and controlling GNP orientation and dispersion within composites and will have important consequences in the industrial processing of graphene polymer composites via the informed design and choice of processing conditions.

INTRODUCTION

Polymer composites filled with graphene have received a great deal of attention due to the extraordinary physical properties of graphene, such as electrical and thermal conductivity, mechanical strength, and barrier properties, which are hoped to be transferred to the composite material. Unlocking these enhancements, while retaining the ease of processing of the host polymer matrix, remains a significant challenge within the polymer environment. A loss of interface due to aggregation, poor dispersion of the graphene within the composite, graphene orientation, and poor interaction between the polymer and the filler particle can all have a negative impact on the composite performance. Therefore, key to achieving the desired material properties is an understanding of how various processing techniques affect the end product. For example, aligning graphene platelets within a composite can almost double the Young’s modulus when compared to a randomly oriented sample. Composites with highly aligned graphene platelets have shown potential as electromagnetic interference shielding materials and in gas/liquid barrier applications. In contrast, for electrical conductivity to be established throughout the composite after processing induced alignment, annealing to restore a random orientation and percolated network is important.

The effect of processing on the conductivity of polymers filled with carbon nanotubes (CNTs) and carbon black (CB) has shown that shear can break down a conducting network within the polymer composite, followed by a subsequent recovery process upon cessation of the shearing. Similarly, graphite in polycarbonate composites has been seen to align following injection molding, followed by gradual disorientation of the graphite improving conductivity. For 2D layered materials, processing effects on the network structure can be even more complex, though few studies have reported these effects in graphene based composites.

The direct monitoring of flow effects on the electrical properties of graphene composites is highly valuable for understanding how the graphene structure and composite properties change under deformation. Despite this, very little work has been done on this so far. Our work focuses on the
steady-shear processing of composites of polystyrene (PS) and graphene nanoplatelets (GNPs). We study the impedance of our composite across a range of shear rates and, with it, find a number of different processes occurring during the shear and subsequent relaxation stages. We relate the change in impedance of the composite to changes in the GNP structure, orientation, and dispersion and show how processing can be used to control this. We support our conclusions with in situ small-angle X-ray scattering measurements. Our results present a significant step forward in understanding and designing appropriate processing conditions in order to achieve the desired enhancements in the properties of polymer–graphene composites.

**MATERIALS AND METHODS**

Polystyrene was supplied from Sigma-Aldrich (SKU 441147, measured $M_w = 273,000$ g mol$^{-1}$, $M_n = 113,000$ g mol$^{-1}$; see Supporting Information). GNP were supplied by XG Sciences (xGNP Graphene nanoplatelets grade M, 5 $\mu$m particle diameter, 5 nm thick). Carbon black was supplied by Sigma-Aldrich (SKU 699624), graphitized with a particle size $< 500$ nm. N-Methyl-2-Pyrrolidone (NMP) was supplied by Fisher Scientific (127630025 Acros Organics 99% extra pure). All materials were used as supplied.

**Composite Preparation.** Nanocomposite samples were prepared by solvent processing in NMP. Polystyrene was dissolved in NMP to a final concentration of 10 wt% polymer. The appropriate amount of GNPs or carbon black was added to the polymer solution to produce a final composite concentration of 5 vol% filler to polystyrene, and the sample transferred to a roller for 18 h. The sample was then sonicated with a solid probe sonicator (300 W, 20 min, 5 s pulses, Cole Parmer 750) to disperse the filler in the polymer solution. Sonication was done on no more than 50 mL of the dispersion at a time. The composite solution was then immediately precipitated dropwise into methanol (10 volume excess to NMP). The resulting precipitate was stirred in methanol for 30 min and then isolated by filtration. This was then stirred again in fresh methanol (18 h) before being isolated by filtration again. The resulting powder was then dried in vacuo (50 °C, 10$^{-2}$ mbar, 18 h).

To prepare samples for testing in the rheometer with in situ impedance or small-angle X-ray scattering measurements, the composites were heat pressed at a temperature of 160 °C with a pressure of 6 T for 30 min into 25 mm diameter discs of thickness approximately 0.5 mm.

**Combined Rheometry and Impedance Spectroscopy.** The pressed discs were tested in a TA Instruments AR 2000 rheometer at 200 °C using the environmental test chamber (ETC) with a nitrogen atmosphere. The bottom geometry comprised a ring electrode of outer diameter 25 mm and inner diameter 19 mm so as to have a better defined shear rate for the impedance measurements. The top plate acted as the sense electrode.

The impedance of the samples was recorded as a function of time using a Zurich Instruments HF2IS impedance spectrometer with an HF2TA transimpedance amplifier in the four wire mode. A voltage of 0.1 V was oscillated at a frequency of 10 Hz, and the gain of the amplifier matched to the requirements of the sample being measured. All samples were subject to an initial preshear for 5 min at a rate of 0.01 s$^{-1}$, and then annealed for 30 min to standardize their structure following pressing and loading in the rheometer. The samples were then sheared for 5 min before the relaxation of the stress was measured by the rheometer upon cessation of the shear, and the impedance measured by the HF2IS as a function of time over the course of the shear and relaxation steps. A median filter was applied to the GNP composite impedance data using Origin 8.6. Fits to the stress relaxation data were also performed using Origin 8.6 (see Supporting Information).

**Small-Angle X-ray Scattering.** SAXS experiments were conducted on beamline I22 at Diamond Light Source. The TA Instruments AR 2000 rheometer was aligned within the beamline and equipped with an electrically heated plate (EHP) in the parallel plate set up with a diameter of 25 mm. The X-ray beam was aligned 1 mm from the edge of the sample to ensure the beam was fully located within the sample while minimizing the path length. The high electron density of the GNPs gives good contrast for X-ray scattering in the polymer matrix. Experiments were conducted using a q range of 0.0018–0.128 Å$^{-1}$ with a beam energy of 12.4 keV and a sample detector distance of 7.629 m, and the data acquired for 0.1 s. The detector image was radially integrated into 45 bins using the Data Analysis WorkbeNch (DAWN) and the orientation determined using this software from the Cinader and Burghardt equation, where the orientation factor is given by the weighted average of the second moment tensor of u:

$$\langle u \rangle = \left( \frac{\cos^2 \beta}{\sin \beta \cos \beta} \right)$$

$$\langle \beta \rangle = \frac{1}{2\pi} \int_0^{2\pi} I(\beta) d(\beta)$$

where $\langle \beta \rangle$ is the azimuthal intensity distribution. $\beta$ is the azimuthal angle, and $I(\beta)$ is the azimuthal intensity distribution.

Samples were subjected to the same shear procedure as for the impedance measurements.

**RESULTS AND DISCUSSION**

Through the use of shear and relaxation processing, it is possible to control the structure of GNPs within the polymer composite. Such control is extremely important for achieving the best possible properties, and the complex nature of polymer–graphene interactions means that there may be little or no enhancement in the properties of the composite if the processing is inappropriate. We have developed a model describing the processes that occur within the composite during processing, as shown in Figure 1.

Our interpretation was developed following the combination of impedance spectroscopy with rheometry, enabling us to monitor changes in the composite electrical properties as a function of shear and subsequent annealing. In addition, we used in situ small-angle X-ray scattering (SAXS) under these conditions to support the model. Initially (top left), the composite starts as an agglomerated structure with a poor GNP dispersion. This secondary agglomeration of the GNPs forms an interconnected network, which provides a conducting path through the composite. At low shear rates (1) of ≤0.1 s$^{-1}$, GNP particles are released from the agglomerates, dispersing more GNPs through the composite, while graphene is also exfoliated from the GNPs. This increases the interfacial area of the GNPs within the composite, and their alignment is also reduced. This establishes a more effective GNP network, increasing the conductivity of the composite to its highest level. Following this shear, the GNP network will reagglomerate under annealing, with the conductivity reducing to levels seen prior to the shearing process. Above a critical shear rate (≥0.3 s$^{-1}$) the response to the shear changes (2). Instead of enhancing the network structure of the GNPs within the composite and increasing the conductivity, the conductivity decreases. This is a consequence of alignment of the GNPs. While the agglomerates themselves may be broken up and better dispersed within the composite at the higher shear rates, when aligned with the shear flow a conducting path is lost. Increasing shear rate leads to greater losses in the conductivity. Such aligned structures are,
however, preferable for enhancing the Young’s modulus of the composite, improving barrier properties, and for electromagnetic radiation shielding. Upon cessation of the shear, an initial rapid reduction in the impedance is observed upon relaxation of the polymer stress. This is ascribed to a rapid reorientation of the GNP particles, facilitated by the polymer stress relaxation. This effect peaks after 10−20 s of annealing at 200 °C. The impedance of the composite then increases again, a consequence of further agglomeration of the GNPs within the composite into isolated agglomerates. This leads to a minimum in the composite conductivity, but with more annealing the GNPs will reagglomerate into a connected network, enhancing the conductivity again. The best processing steps for particular properties are highlighted.

A schematic of the rheometer geometry developed to incorporate simultaneous measurements of impedance spectroscopy is shown in Figure 2. The bottom plate of the geometry acts as the source electrode, and the impedance is measured through the sample to the top plate, which acts as the sense electrode. A sliding contact was used on the top plate so that continuous shear could be applied. A ring electrode was used on the bottom plate so that the impedance was measured at a well-defined shear rate near the edge of the sample (Figure 2a). This set up allows evaluation of how the electrical properties of the composite changes as a function of shear conditions and history, with different structures giving different responses. For example, a well-developed network of the GNPs would have low impedance and high conductivity (Figure 2b). GNP particles aligned in the shear direction would have high impedance when the conducting path through the composite is broken (Figure 2c). The samples were sheared for 5 min at a constant shear rate. Upon cessation of the shear, the change in impedance continued to be monitored while annealed in the rheometer. The change in the impedance of the GNP composites during shear is shown in Figure 3a. A complex range of behaviors is observed, with a distinct shift between shear rates. For PS + 5 vol % GNPs, at shear rates ≤ 0.1 s⁻¹ the impedance of the composite is reduced during shear, enhancing the conductivity. This is consistent with the effects observed by Beckert et al. when shearing composites of PS/PS-grafted-functionalized graphite oxide and graphene nanocomposites. The required shear rate, γ, for graphene exfoliation can be estimated as

\[
\dot{\gamma} = \frac{[\sqrt{E_{5.6}} - \sqrt{E_{5.1}}]^2}{\eta L}
\]

(3)
where $E_{SG}$ and $E_{SL}$ are the surface energies of the graphene and the polystyrene, respectively, $\eta$ is the viscosity, and $L$ is the GNP length. The zero shear viscosity of the polymer is $\sim 18,000$ Pa s (see Supporting Information). $L = 5 \, \mu m$, $E_{SG} = 71 \, mJ \cdot m^{-2}$, and $E_{SL} = 42 \, mJ \cdot m^{-2}$. The process of exfoliation and the release of GNP particles from agglomerates would increase the interface of the filler within the composite, helping to build a more effective network, and increasing the conductivity of the composite.

At higher shear rates ($\geq 0.3 \, s^{-1}$) the impedance of the composite is increased, with the magnitude of the change increasing with the shear rate. This shows a loss of the conducting network within the composite. Following shear at 3 $s^{-1}$ for 5 min the impedance of the composite is four orders of magnitude greater than that of the composite following shear at a rate of 0.01 $s^{-1}$. This is also reflected by an increase in the phase angle of the network during shear. This can be seen in Figure 3b, where the phase angle of the composite increases from a value of $\sim 2.5$ degrees at rest to values over 40 degrees during shear. Such a change reveals a switch from predominantly resistive behavior to one with increasingly high capacitive contributions. This shows a change in the composite structure under shear. Alignment of the GNP, parallel to the rheometer plates, at higher shear rates would result in the loss of a resistive network and the formation of a layered structure of polymer and aligned GNP. This produces a capacitive structure, where the impedance is measured between the rheometer’s parallel plates, and would account for such an increase in the phase angle.

The impedance of the composite immediately following cessation of the shear is shown in Figure 4a (zones 1 and 2). Following an initial lag of $\sim 7$ s after the shear, the aligned samples show a reduction in the impedance, as highlighted by zone 3. The magnitude of this drop is dependent on the preceding shear rate: the faster the shear, the larger the drop in the impedance observed. This unprecedented change in impedance is consistent with the relaxation of shear induced
alignment within the composite, establishing a more effective conducting network. This is also supported by a decrease in the phase angle of the composite (Figure 4b), which suggests a change from an aligned capacitive structure to a more randomly oriented, resistive network structure.

We calculate the rotational diffusivity, \( \Theta \), of our GNPs to be \( \sim 2.5 \times 10^{-9} \) s\(^{-1}\). Remarkably we see the relaxation process occur over the course of \( \sim 10 \) s, which exceeds the calculated reorientation rate for GNPs under quiescent conditions by approximately eight orders of magnitude. We estimate \( \Theta \) from the equation for rotational diffusion of ellipsoid particles:\(^{26} \)

\[
\Theta = \frac{3kT}{16 \pi \eta_{\text{matrix}} a^4} \left[ 2 \ln \left( \frac{2a}{b} \right) - 1 \right]
\]

(4)

where \( k \) is the Boltzmann constant, \( \eta \) the matrix viscosity (\( \sim 18000 \) Pa s), \( a \) the diameter (5 \( \mu \)m), and \( b \) the thickness (5 nm) of the GNPs. Rapid losses in alignment of clay particles in polymer composites have previously been linked to the polymer stress relaxation,\(^ {26} \) or a coupling between the polymer chains and the clay accelerating the particle relaxation.\(^ {53} \) Here, we have observed the relaxation of the polymer stress following cessation of the shear over the course of the time scale observed for the change in impedance measured (see Supporting Information). The relaxation of the polymer stress may therefore lead to an initial rapid orientation relaxation of the GNPs within the composite. Gradually, however, following this initial decrease, the impedance increases to a maximum. This is also seen at the lower shear rates (zone 4) before then reducing again to levels below that seen immediately following the shear step. Clearly there are multiple structural evolutions within the composite during the relaxation stage. Following any randomization of the GNPs, we propose that a two-step agglomeration process occurs. Initially there is a loss of network structure and a collection of discrete agglomerates, which reduces the composite conductivity. A secondary agglomeration step results in the formation of an interconnected network of the discrete agglomerates, and the formation of a conducting network (\( S \)).

Changes in the dispersion of graphene in polypropylene composites upon annealing at 200 °C have shown the formation of an interconnected, macroscopic network from smaller aggregates, increasing composite conductivity,\(^ {30} \) while recovery of conductivity following deformation in graphene based composites has been observed following various tensile, bending, compression, and oscillatory shear tests.\(^ {37,31,32} \) Oscillatory shear can have markedly different effects on the conductivity of polymer composites than steady shear, as network deformation can be reversible,\(^ {9} \) described as an agglomerated network memory effect.\(^ {33} \) Using steady-shear, our results are more akin to processing conditions than oscillation. The complexity of relaxation behavior seen in our work has not been observed before and is unique to these GNP composites. A control composite made with carbon black did not show this relaxation behavior, instead showing monotonic reductions in impedance following all the preceding shear rates studied (see Supporting Information). As with the rapid disorientation (step 3) the time scale for the GNP agglomeration processes is much quicker than expected by Brownian motion and is likely to be driven by attractive interparticle interactions in the polymer matrix. Similar conclusions were formed from the structural evolution of polypropylene–clay nanocomposites, but this is the first time that such behavior has been verified through dramatic changes to impedance.\(^ {17,18} \)

The processing is seen to be cyclic. For example, Figure 5 shows the change in the impedance of the composite under repeated shear and annealing steps. During step (1) the composite is sheared at a rate of 0.3 s\(^{-1}\), resulting in an increase in the impedance of the composite (and a reduction in conductivity). Upon cessation of the shear, the sample is annealed (2). During this stage the impedance increases to a maximum, before decreasing again, as previously described. In stage (3) the sample is sheared at a rate of 0.01 s\(^{-1}\). There is a brief spike in the impedance upon start-up of the shear, before the impedance of the composite is reduced over the course of the shear to its lowest level during the experiment, increasing the composite conductivity. This matches with the behavior described in Figure 3a. Once this shear stage is stopped, (4) we find the impedance increases to a maximum under annealing, before then reducing again.

Direct confirmation of the internal orientation of GNPs during shear processing was determined by SANS. Our rheometer was aligned within the beamline I22 at Diamond Light Source to measure the SANS of the composite samples 1 mm from their edge while being sheared or annealed. Graphene materials have been shown to have different structures depending on the scattering length scale observed.\(^ {34} \) Here, we monitor the orientation at our largest length scale, in the \( q \) range 0.002–0.0035 Å\(^{-1}\) (length scale of \( \sim 2000 \) Å) so as to avoid the contribution of wrinkles or edges to the scattering.\(^ {34} \) GNP orientation factors were calculated using the Cinar and Burghardt equation.\(^ {23} \) A value of +1 indicates perfect orientation in the flow direction, −1 perfect orientation in the transverse direction, and a value of zero indicates no alignment. The results show a number of interesting effects. Following shear at 0.01 s\(^{-1}\), where the sample has its highest conductivity, it is apparent from the detector image there is some alignment present in the sample (Figure 6). A randomly oriented sample would give a perfectly isotropic scattering pattern. At a shear rate of 0.3 s\(^{-1}\), it can be seen that the anisotropy of the scattering has increased (Figure 6). Two peaks are observed at angles of −90° and +90° as the scattering intensity is plotted as a function of the azimuthal angle. This highlights the orientation of the GNPs within the composite and shows the increase in alignment of the sample following

![Figure 5](image-url)
shear at 0.3 s⁻¹. The calculated orientation factor increases from 0.27 to 0.47 at 0.01 and 0.3 s⁻¹ shear, respectively.

Interestingly, we find that shearing the sample at the lowest rates reduces the pre-existing alignment of the GNPs within the composite. It can be seen in Figure 7a that before the preshear stage the composite retains a high degree of alignment from the initial pressing. Upon start-up of the shear at a rate of 0.01 s⁻¹, the orientation initially increases slightly before reducing for the rest of the shear stage. Upon cessation of the shear the orientation monotonically decays, but at a much lower rate than during shear. Further shear of the sample at 0.01 s⁻¹ reduces the composite alignment further. The reduction in alignment corresponds with the decrease in impedance of the composite (Figure 3a) at this shear rate and highlights GNP structural changes leading to increasing conductivity.

At higher shear rates, the orientation increases during shear. The speed at which the orientation increases is also seen to increase with the shear rate, as shown in Figure 7b. Upon start-up of the shear, the orientation factor increases instantly and is much quicker at a shear rate of 1.8 s⁻¹ than at 0.3 s⁻¹. This is also matched by the impedance response of the composite, where the 0.3 s⁻¹ shear impedance increases at a lower rate than the 1.8 s⁻¹ sample. The impedance eventually plateaus, while the orientation factor at both rates seems to average around a plateau of ~0.5, suggesting a limit to the orientation of the GNPs in the system. The higher impedance at a shear rate of 1.8 s⁻¹ compared to 0.3 s⁻¹ shear rate is also likely to be a consequence of increased breakup of GNP networks, in addition to the increase in alignment. The orientation of the GNPs under the action of shear, however, suggests that these actions are closely linked to the effect on the impedance of the composites during processing.

The changes in alignment of the GNPs at different shear rates show the importance of the shear processing to the structure of the composite, which directly relates to the ability to enhance various properties of the polymer.

■ CONCLUSIONS

We have shown a series of property changes through continuous shear and anneal processing in composites filled with GNP. We have combined impedance spectroscopy with rheometry and used small-angle X-ray scattering to investigate the GNP orientation within the composites during these processing steps. They have a huge impact on the composite impedance, with reversible changes of several orders of magnitude, and the conductivity of the composite can be enhanced or diminished. Low shear rates help to build a network, improving conductivity, while higher rates break the network and align the GNPs, reducing conductivity. Upon annealing post-shear, several more changes are observed comprising disorientation and agglomeration steps. Our insights are important for understanding how the GNP structure evolves and can be controlled within the polymer environment. If the composite structure is fixed at specific points through the processing cycle, it will be possible not only to gain control over the electrical properties of the composite, but to optimize production for a whole range of different properties. These results therefore present a significant advancement in our understanding of the relationship between processing, structure, and properties in polymer–graphene composites. As our insights hinge predominantly on the physical shape and aspect ratio of the GNP (rather than its precise chemical interactions with the matrix), it presents a model system with wide applicability and impact upon the
family of composites formed from two-dimensional filler materials, platelets, and nanoparticles.

ASSOCIATED CONTENT

Supporting Information
The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.chemmater.7b04343.

Rheo-impedance measurements for composites of PS + carbon black, used as a control sample. Effect of temperature on the processing of PS + GNP composites. Zero shear viscosity measurement for the PS. Composite stress relaxation times as a function of shear rate. Size exclusion chromatography of the PS. Rheological and impedance data for percolation of as pressed samples. Resistance and reactance of impedance under shear (PDF).

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Notes
The authors declare no competing financial interest.

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