

## ***In-Situ* Exfoliation of Graphene in Epoxy Resins:**

### **a Facile Strategy to Efficient and Large Scale Graphene Nanocomposites**

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#### **ABSTRACT**

Any industrial application aiming at exploiting the exceptional properties of graphene in composites or coatings is currently limited by finding viable production methods for large volumes of good quality and high aspect ratio graphene, few layer graphene (FLG) or graphite nanoplatelets (GNP). Final properties of the resulting composites are inherently related to those of the initial graphitic nanoparticles, which typically depend on time-consuming, resource-demanding and/or low yield liquid exfoliation processes. In addition, efficient dispersion of these nanofillers in polymer matrices, and their interaction, is of paramount importance. Here we show that it is possible to produce graphene/epoxy nanocomposites *in-situ* and with high conversion of graphite to FLG/GNP through the process of three-roll milling (TRM), without the need of any additives, solvents, compatibilisers or chemical treatments. This readily scalable

production method allows for more than 5 wt.% of natural graphite (NG) to be directly exfoliated into FLG/GNP and dispersed in an epoxy resin. The *in-situ* exfoliated graphitic nanoplatelets, with average aspect ratios of 300-1000 and thicknesses of 5-17 nm, were demonstrated to confer exceptional enhancements in mechanical and electrical properties to the epoxy resin. The above conclusions are discussed and interpreted in terms of simple analytical models.

## INTRODUCTION

As vastly reported, graphene's superb properties (theoretical specific area of *ca.* 2360 m<sup>2</sup>/g,<sup>1</sup> thermal conductivity of ~5000 W·m<sup>-1</sup>·k<sup>-1</sup>,<sup>2</sup> intrinsic charge mobility of 200,000 cm<sup>2</sup>·V<sup>-1</sup>·s<sup>-1</sup>,<sup>3</sup> Young's modulus of ~1.0 TPa and strength of ~130 GPa<sup>4</sup>), impermeability to gas or liquids<sup>5</sup> promise to have significant impact on a host of different industries and advanced application fields such as sensors, optoelectronics, batteries, transparent conductive electrodes/films or energy management.<sup>6-13</sup> However, its industrial adoption will only be possible when a method to produce 'good quality' (i.e. high aspect ratio, minimum thickness, few defects) graphene, few layer graphene (FLG) or graphite nanoplatelets (GNP) at low cost and in large-scale, will be developed.

If graphene is a single layer of *sp*<sup>2</sup> hybridised C atoms, FLG usually indicates a nanoparticle characterised by a number of graphene layers between 3 and 10 (< 5 nm thickness), which increases to 10 to 100 layers for GNP. Unfortunately the same physical properties (e.g. mechanical, electrical) that make graphene such an exciting material, deteriorate rather swiftly with number of stacked layers. It has been reported for instance that the Young's modulus decreases from 1 TPa to ~ 600 GPa and 400 GPa when going from an isolated graphene particle to a stack of 5 or 10 layers, respectively.<sup>14</sup> The problem is that single layer graphene is very difficult to produce - particularly in the case of large lateral dimensions - as well as to process. In fact it suffers from shape instability as it tends to roll, scroll, wrinkle or fold-up unless it is constrained onto a solid surface.<sup>15</sup> So a difficult compromise between properties, costs and processability has to be faced. FLG/GNP could well represent a more viable alternative to graphene if they could be produced cheaply and with large high aspect ratio as well as being processed easily.

Currently the most promising large scale production method for graphene/FLG/GNP is liquid exfoliation of graphite. Within liquid exfoliation, two possible routes can be distinguished: either graphite is first functionalised (mainly oxidised to produce graphene oxide) before being exfoliated in water<sup>16</sup> or it is directly exfoliated in organic solvents (notably NMP),<sup>17</sup> ionic liquids,<sup>18-19</sup> mixed solvents (acetone/water)<sup>20</sup> or in water/surfactants mixtures,<sup>21-22</sup> via contact or non-contact techniques.<sup>22-23</sup> The oxidising route is very efficient at producing predominantly monolayers and relatively large nanoparticles (50 nm to 3  $\mu\text{m}$ ),<sup>24-25</sup> but it introduces large quantities of defects which significantly and irreversibly compromise the physical properties of graphene. Graphene oxide, for instance, has a Young's modulus of about 250 GPa,<sup>26</sup> thermal conductivity of about 18  $\text{W}\cdot\text{m}^{-1}\cdot\text{k}^{-1}$  (46% carbon content),<sup>27</sup> and a low electrical conductivity which, for fully reduced monolayers, can only be partially recovered up to 2  $\text{S}\cdot\text{cm}^{-1}$ , with a field effect mobility of 2–200  $\text{cm}^2\cdot\text{V}^{-1}\cdot\text{s}^{-1}$  at room temperature.<sup>28</sup> The direct liquid exfoliation of graphite gives rise to nanoplatelets with few defects but with larger thickness (typically between 1 and 10 layers) and limited lateral size (typically between 100 nm to 1  $\mu\text{m}$ ). But high boiling point (organic) solvents are costly and difficult to handle and to extract while surfactants can essentially act as contaminants (for instance by limiting the electrical properties) if not carefully extracted. Moreover all liquid exfoliation methods suffer from a number of additional common problems including: low yield (typically 1-3%), use of energy intensive exfoliations techniques (usually ultrasonication but also high shear mixing) and/or long processing time (from several hours to several hundreds of hours) and low concentration of stable graphene/FLG/GNP liquid suspensions (typically up to few mg/ml).<sup>29</sup> Last but not least remains the problem of how to process a relatively low concentrated, only partially stable, graphene/FLG/GNP liquid suspension and convert it into a polymer nanocomposite or other assembly or device. All the above have effectively hindered any real industrial applications.

In order to overcome the above limitations, herein we present for the first time an effective and powerful route to produce *in-situ* exfoliated FLG/GNP directly into the polymer matrix of choice, by three-roll milling (TRM), avoiding any intermediate steps (e.g. filtering, removal of the dispersing liquid medium, purification, drying of powder, redispersion into the final matrix, etc.).

TRM has already been proven very effective in dispersing 1D nanofillers such as carbon nanotubes (CNTs)<sup>30</sup> or 2D nanoparticles such as nanoclays,<sup>31</sup> within epoxy resins. Unfortunately, only limited reports on the dispersion of graphitic materials are present in the scientific literature.<sup>23, 32</sup> Recently Throckmorton *et al.* claimed that TRM is capable to partially exfoliate and disperse graphite into FLG/GNP directly into an epoxy resin but only in presence of an ionic liquid as solvent/dispersant.<sup>33</sup> In absence of the ionic liquid, no electrical conductivity could be even detected for epoxy composites with filler content as high as 3 wt.%.

For the first time we are able to demonstrate that graphite can be efficiently exfoliated and dispersed into an epoxy resin to *in-situ* produce FLG/GNP, without the need of any additives, solvents or compatibilisers and chemical or physical pre-treatments. This work presents a complete study of the relationship between TRM conditions, the structure/property of high-quality FLG/GNP and the properties achieved on the final nanocomposites, interpreted in terms of simple energy balances and geometrical arguments, a numerical model of the TRM process and the Hansen solubility parameters. Careful control over the TRM parameters results in relatively large aspect ratio FLG/GNP (up to 1000) combined with a relatively minimal thickness (minimum average thickness 5 nm) and, notably, 100 % conversion from the starting natural graphite powder.

## **EXPERIMENTAL**

### **Materials**

Natural graphite flakes (NG) were purchased from Alfa Aesar (Product No. 43319). The MVR444 two-part epoxy resin was supplied by Cytec (UK).

### **Exfoliation and dispersion protocols**

Dispersions with various NG contents were prepared using a three-roll mill (80E EXAKT GmbH, Germany). The process was done in three steps: **i)** the NG powder was pre-dispersed in the epoxy resin at 70 °C for 10 min using a magnetic stirrer to produce: **a)** composites with concentrations of 1, 2, 3, 4 and 5 wt.% and **b)** a 5 wt.% masterbatch; **ii)** this masterbatch was then diluted to the desired final loading (1 to 4 wt.%) for sake of comparison and **iii)** finally, all prepared samples were fed into the TRM. In a typical experiment, the epoxy/NG mixture was

passed 8-10 consecutive times through the TRM (referred to as 8-10 cycles), whereby the speed and roll-to-roll distance (gap size) as well as temperature were varied. A summary of the five different processing parameters is given in the Supporting Information (**Table S3 and S4**).

The final particle sizes were found to be affected predominantly by only two parameters: a) shear loading (controlled mainly by roll speed and gap size) and b) temperature. For sake of simplicity we will focus only on these two parameters which will be discussed separately in section 3.1, by analysing three series of samples following: Protocol I, II and III.

- Protocol I includes up to ten cycles, all done at a fixed gap distance of and fixed NG concentration (5 wt.%). The rotation speed of the apron roll was progressively increased from 30 to 60, 90, 150 and 200 rpm every two cycles.
- Protocol II includes up to eight cycles, with the same rotational speed (200 rpm) of the apron roll for all cycles and fixed NG concentration (5 wt.%). During Protocol II, the epoxy/NG mixtures were passed through the TRM twice in gap mode (fixed roll-to-roll distance) and then six cycles in force mode (fixed applied force of 5.0 N/mm) both done by using a ratio of 1:3:9 between  $N_1$  (feed roller) :  $N_2$  (central roller) :  $N_3$  (apron roller). During the first cycle in gap mode, gaps of  $N_1/N_2 = 120 \mu\text{m}$  and  $N_2/N_3 = 40 \mu\text{m}$  were used. For the second cycle,  $N_1/N_2$  and  $N_2/N_3$  were reduced to 60 and 20  $\mu\text{m}$ , respectively.
- Protocol III utilized the same process parameters of Protocol II except for the resin temperature, which was varied between 25 °C and 40 °C. 4 and 5 wt% graphite were added to epoxy for exfoliation via Protocol III.

### **Fabrication of epoxy nanocomposites**

After the exfoliation/dispersion conditions following Protocol I, II and III, the hardener was added to the epoxy in a 58:100 ratio after exfoliation/dispersion. The mixtures were degassed under gentle mechanical stirring at 70 °C for 60 min in a vacuum chamber (pressure of -1 bar). Mixtures were then casted into stainless steel moulds at room temperature and cured in an oven. The following curing conditions were applied: **i**) temperature ramp from RT to 120 °C (3 °C·min<sup>-1</sup>) followed by a 90 min isotherm, **ii**) temperature ramp from 120 °C to 180 °C at

(3 °C·min<sup>-1</sup>) followed by a 180 min isotherm and **iii**) cooling down from 180 °C to RT at 3 °C·min<sup>-1</sup>.

### **Characterisation techniques**

Morphological studies were carried out using optical microscopy (OM, Olympus BX 60) and/or scanning electron microscopy (FEI, Inspector-F) with an electron beam of 20 kV. SEM was conducted to assess the morphology and in particular the length (defined as the longest lateral dimension) of the particles. Specimens were prepared by “extracting” the particles from the different dispersions straight after TRM. Typically, a small amount of uncured epoxy/particle dispersion was immersed in acetone to dissolve the epoxy, followed by filtration using a 0.2 µm PA6 membrane. The obtained particles were then washed three times to remove any remaining epoxy, which was confirmed by optical microscopy. These particles were re-dispersed in acetone to a final concentration less than 5 mg·ml<sup>-1</sup>. The obtained suspension was casted onto an ITO coated glass substrates without gold coating at RT. After evaporation of acetone, the specimens were imaged as prepared. SEM was then used to estimate the length of the FLG/GNP by measuring 100-120 particles. Average values are reported.

Transmission electron microscopy (TEM) (JOEL JEM-2010) was used to see the morphology and different degrees of the exfoliated graphene. The FLG/GNP particles, extracted from uncured liquid epoxy according to the method described in the previous above, were dispersed in acetone (typical concentration 0.05 mg/ml) before being deposited on TEM grids (300 mesh, 3 mm, purchased from TAAB, C267/050) by drop casting. The copper grid with graphene dispersion was dried at RT for 10 min. 10-15 representative particles were collected for each specimen. The thickness of the FLG/GNP particles is estimated from the thickness of the edge of the particles sticking out of the plane of the copper grid.

Raman spectroscopy (Nicolet Almega XR, High-Performance Dispersive Raman Spectrometer) was utilized to characterise of natural graphite, exfoliated GNP. GNP sheet was prepared by vacuum filtration of the dispersion through a porous membrane (PVDF, pore size 0.45 µm). Raman measurements were performed with a wavelength of 532 nm.

The viscosity of the pure epoxy and the complex viscosity of the graphite/epoxy mixtures were measured by an AR2000 Rheometer equipped with 40 mm steel parallel plates. The

temperature was ramped both up and down at 3 °C·min<sup>-1</sup> from 25 to 100 °C and between 1-100 Hz, at a shear strain of 1%.

Drop shape analyzer (DSA100 KRÜSS, GmbH, Germany) was used to measure the contact angle between the liquid epoxy and a glass substrate. The glass substrates were thoroughly cleaned by acetone. Surface energies were calculated from contact angle data of sessile drops of 10 µL. In order to make experiments easier, we choose ethylene glycol as a non-volatile (boiling point: 197.3 °C) reference solvent, with a surface tension of 47.70 N/m at 20 °C. From the measured contact angles, the surface tension of the epoxy was extracted according to the Young-Laplace equation. For more details please refer to Supporting Information **Figure S5** and section: **Surface tension**.

X-ray diffraction (XRD) (Philips PW 3830 automated powder diffraction) was used to characterize the thickness of exfoliated GNPs within the epoxy composites. The samples were cut into rectangular beams (dimensions 3.2 x 10 x 30 mm<sup>3</sup>). The X-ray texture scans were obtained between  $2\theta = 20-90^\circ$  at a scanning rate of 1°/min. The average out-of-plane crystallite thickness of the GNPs ( $t$ ) was estimated using the full width at half maximum (FWHM) of the (002) peak by Scherrer's equation:

$$t = \frac{K\lambda}{\beta \cos \theta} \quad \text{Equation 6}$$

where  $\beta$  is the line breadth (FWHM) in radians with the instrumental broadening subtracted,  $\lambda$  is the X-ray wavelength, and  $\theta$  is the diffraction angle of the (002) peak. The coefficient  $K$  was taken to be 0.89 according to Raza *et al.*<sup>34-35</sup> XRD result was the average value of three repetitions.

The conductivity of all samples was measured by a two probes method using a picoammeter (Keithley 6485) and a DC voltage source (Agilet 6614C). Samples were cut into beams of 3.2 x 10 x 30 mm<sup>3</sup> and silver paste was applied to the cross-section to ensure good contact between the electrodes and the sample. Voltages in the range 5 to 10 V were used. For specimens with resistances exceeding 10<sup>10</sup> Ohm, electrical resistivity was no longer measurable and the samples were considered as 'non-conductive'. Three specimens for each composite were tested in order to obtain average values of conductivity.

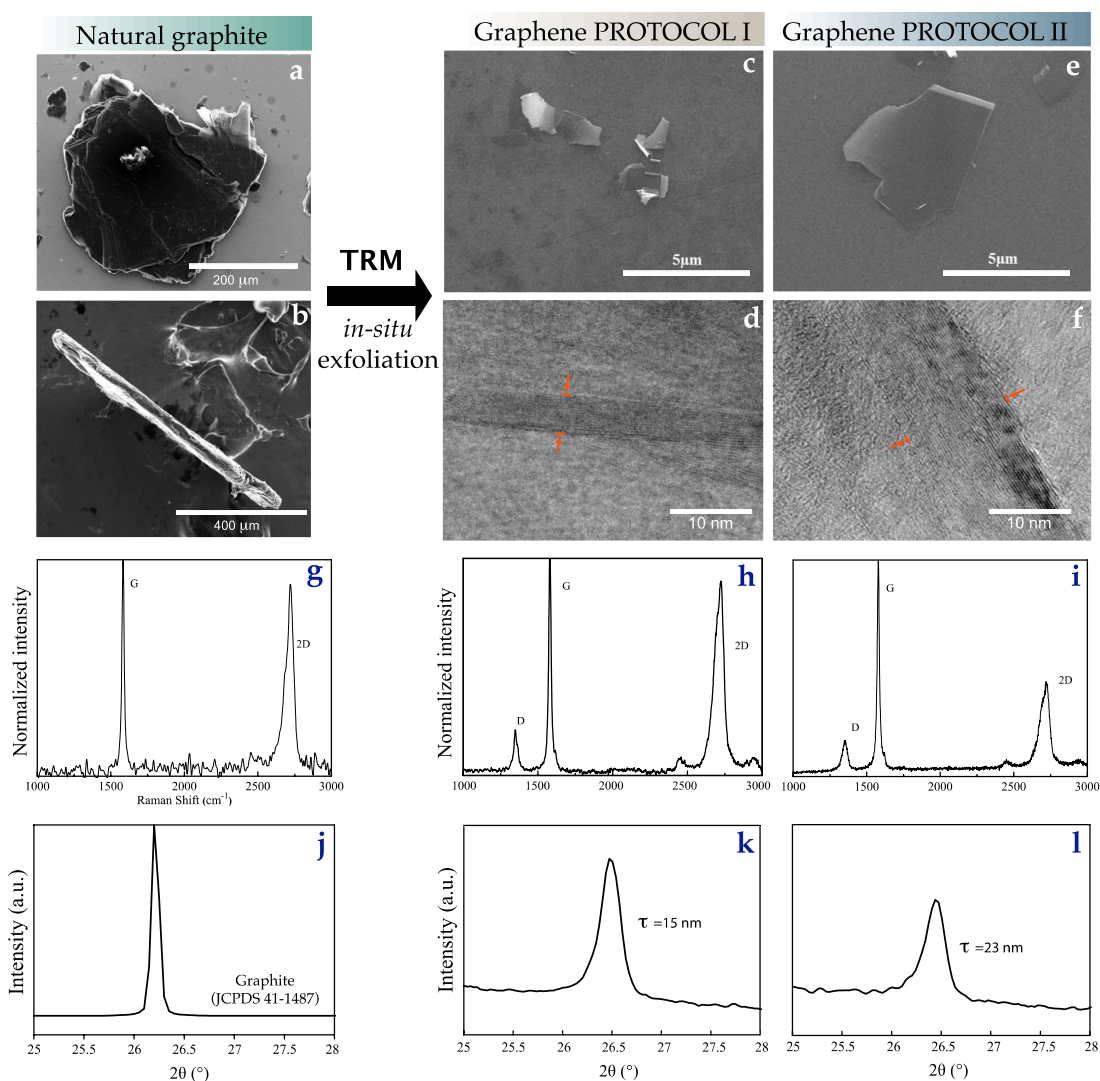
Flexural test specimens were prepared according to the ASTM-D 790 standard. The samples were cut into beams of dimensions 3.2 x 12.7 x 70 mm<sup>3</sup>. The cut surfaces were polished by hand using different grade abrasive paper (from 1000 to 4000 grit). The support span to depth ratio was 16:1 and a strain rate of 0.01 mm/min was used.

## RESULTS AND DISCUSSION

### **Effect of shear on *in-situ* exfoliation of natural graphite in epoxy resin**

SEM images in **Figure 1.a-b** present the typical morphology of the natural graphite flakes used in this study. With lateral dimensions varying between  $\sim 600 \pm 150 \mu\text{m}$  and  $\sim 800 \pm 200 \mu\text{m}$  and a thickness of circa  $40 \mu\text{m}$ , the initial natural graphite flakes present an aspect ratio of circa  $20 \pm 5$ . **Figure 1.g** shows a typical graphitic Raman spectra with the G and 2D peaks positioned at, respectively,  $1585 \text{ cm}^{-1}$  and  $2718 \text{ cm}^{-1}$ . No D peak (defect) was instead observed due to the high degree of crystallinity.





**Figure 1.** *In-situ* exfoliation of natural graphite (SEM images a and b) by TRM. SEM and TEM images of FLG/GNPs after exfoliation through Protocol I (c and d) and Protocol II (e and f). Comparison of Raman and XRD spectra ( $2\theta \approx 26.1^\circ$  indexed to the (0 0 2) planes of a hexagonal graphite lattice) before and after exfoliation: graphite (g and j), FLG/GNP Protocol I (h and k) and FLG/GNP protocol II (i and l), respectively.

After the TRM processes the morphology of the natural graphite changes dramatically. The particle contour changes from relatively smooth and round to mostly sharp edged (**Figure 1.c**, **Figure 1.e**). Both lateral dimensions and thickness (**Figure 1.d**, **Figure 1.f**) are reduced but, interestingly, not in the same proportion. The platelet thickness is reduced by

approximately three orders of magnitude, hence demonstrating the success of the *in-situ* exfoliation, while the lateral dimension is reduced by about two orders of magnitude<sup>34</sup>.

**Table 1** summarises the average particle dimensions calculated from a statistical analysis of the SEM and TEM images and the X-ray data in **Figure 1**. An obvious difference in size of the graphitic particles obtained by the two processing protocols can be observed. Protocol I gives rise to slightly thinner nanoparticles (~ 6 nm instead of ~ 14 nm according to TEM observations). Conversely, Protocol II is able to achieve considerably larger nanoparticles (~ 3.51  $\mu\text{m}$  compared to ~ 0.76  $\mu\text{m}$ ). As a result the nanoparticles obtained by Protocol II possess a 3 times higher aspect ratio than those obtained by Protocol I, and 7.5 times higher than of original natural graphite particles. A discrepancy between the values of thickness as measured by TEM and XRD is noticed, consistent with previous reports which explained it in terms of differences in number and quality of sampled particles – which is inherent with the different preparation methods – and as well as the coefficient K selected in the Sherrer equation.<sup>34-35</sup> It is noted that the particles measured by TEM are extracted from uncured liquid resins and drop casted on copper grids from very diluted dispersions. The particles measured by XRD are embedded into the composites after curing the epoxy resins. A partial reagglomeration of particles is expected during curing, which will then result in an increase in the average particle thickness.

For sake of completeness, within this paper we will use both (average) values of thickness, as measured by XRD and TEM, and hence estimate two values of aspect ratio ( $L/T_{\text{XRD}}$  and  $L/T_{\text{TEM}}$ ) for each protocol.

**Figure 1.h-i** present also typical Raman spectra for the *in-situ* exfoliated particles obtained by the two protocols. Of interest here is the ratio of the intensities of the D and G bands,  $I_D/I_G$  (reported in **Table 1**), which gives indications of the quality of the particles; i.e. the lower the ratio the lower the defects in the graphitic structure. The particles obtained with Protocol II attain a lower  $I_D/I_G$  ratio (0.16 instead of 0.20). This difference can be explained by the reduced presence of edge defects for larger flakes (Protocol II), in analogy with previous reports. For instance Khan *et al.* showed a decrease of the  $I_D/I_G$  ratio from 0.22 to 0.08 when larger flakes were selectively separated from a solvent dispersion, by decreasing the centrifugation rotational speed from 4000 to 500 rpm.<sup>29</sup>

**Table 1.** Properties of initial NG and FLG/GNP particles exfoliated according to Protocol I & II.

Sample	Length by SEM L (μm)	Thickness by XRD T <sub>XRD</sub> (nm)	Thickness by TEM T <sub>TEM</sub> (nm)	Aspect Ratio (L/ T <sub>XRD</sub> - L/ T <sub>TEM</sub> )	I <sub>D</sub> /I <sub>G</sub> (-)
NG	~800	-	~ 40000*	N.A. - ~ 20	0
Protocol I	0.76 ± 0.34	~ 15 ± 3	6 ± 2	50 - 126	0.20 ± 0.05
Protocol II	3.51 ± 0.87	~ 23 ± 2	14 ± 5	150 - 250	0.16 ± 0.08

\*estimated from SEM micrographs.

In order to understand the exfoliation process shown above, let us start from a simple energy balance and geometrical argument.

Let us assume initial flakes of sizes  $L_x, L_y, L_z$  and define  $N_x, N_y, N_z$  as the number of cuts (or delamination, in the case of  $z$  axis) taking place along the related sides during the exfoliation and thus resulting in a total number of fragments equal to  $N = N_x \cdot N_y \cdot N_z$ . Indicating with  $\eta_f$  or  $\eta_d$  the energy fractions dissipated, respectively, during (in-plane) fracture of graphene (or other 2D materials) or during (out-of-plane) exfoliation (delamination) to separate the layers, the energy balance imposes  $\eta_f + \eta_d = \eta$ , where  $\eta$  is the efficiency of the process.

The energy dissipated by fracture is:

$$W_f = \gamma_f L_z (L_x (N_y - 1) + L_y (N_x - 1)) \quad \text{Equation 1}$$

whereas that dissipated by delamination is:

$$W_d = \gamma_d L_x L_y (N_z - 1) \quad \text{Equation 2}$$

where  $\gamma_f$  or  $\gamma_d$  are the surface energies of fracture or adhesion respectively and  $z$  is assumed to be perpendicular to the layers. Noting that  $N_{x,y,z} \gg 1$  and that  $L = L_x / N_x$ ,  $W = L_y / N_y$  and  $T = L_z / N_z$  are the final lateral sizes  $L$ ,  $W$  and thickness  $T$  of the flakes, we find the

following prediction of the flake aspect ratio ( $\lambda$ ) during exfoliation (independent from  $N$ ) imposing the energy balance:

$$\lambda \equiv \frac{\sqrt{LW}}{T} = \frac{\eta_d \gamma_f}{\eta_f \gamma_d} \frac{L/W + 1}{\sqrt{L/W}} \quad \text{Equation 3}$$

Posing  $L/W \approx 1$  (assuming cylindrical nanoplatelets),  $\eta_d/\eta_f \approx 1$  (assuming the energy dissipated by fracture is equal to the energy dissipated by exfoliation) and  $\gamma_d/\gamma_f \approx 10 - 100$ , results in  $\lambda \approx 20 - 200$  that is of the order of common experimental observations in the literature. On the other hand, values of  $\lambda$  above 1000 could be achieved if  $\eta_d/\eta_f$  is above 5-50, in other words if more energy is used for exfoliation (i.e. reduce thickness) rather than fracture (i.e. reduce lateral dimensions). We believe  $\eta_d/\eta_f$  is intimately linked with the specific processing methods and conditions used. In our case, the fact that Protocol I achieves lower aspect ratio than Protocol II suggests that  $(\eta_d/\eta_f)_{Protocol I} < (\eta_d/\eta_f)_{Protocol II}$ . By using the experimental values of aspect ratio found for Protocol I and II (**Table 1**), it is possible to estimate that the ratio  $\eta_d / \eta_f$  is as high as 2.5 - 6.3 for Protocol I and 7.5 - 12.5 for Protocol II.

But in order to explain the variation in nanoparticle aspect ratio obtained with our different protocols, it is necessary to better understand the actual process. In general it is expected that a higher shear loading should result in more efficient exfoliation, hence thinner graphitic particles. Recently Paton *et al.*<sup>22</sup> demonstrated that well exfoliated FLG/GNP particles could consistently be obtained when shear rates exceeding  $10^4 \text{ s}^{-1}$  were reached, independently from the mixing method and the dispersing liquid (NMP or water/NaC) used.

In **Figure 2a**, the estimated shear rates achieved in the two TRM Protocols is presented. Shear rates and shear stresses are calculated by modelling the TRM process in analogy to a recent work of Magnier *et al.*,<sup>36</sup> who developed a model to describe the calendaring process (in particular the rolls separating force) of power-law non-Newtonian fluids between counter rotating rolls at different velocities. The isothermal model was based on the lubrication approximation, as in classical calendaring models.<sup>37</sup> But, due to the asymmetry caused by the different velocities of the rolls, the generalised Reynolds equation had to be solved taking into account various velocities profiles in different zones (4-5 zones). In zone 3, at a certain

horizontal distance  $x=x^*$  (please note that  $x$  is negative until  $x = 0$  at the nip region) before the nip region, in correspondence of a vertical distance (height) between rolls of  $2h=2h^*$  (see Supporting Information **Figure. S1**), the pressure is maximum ( $dp/dx=0$ , see Supporting Information **Figure. S1**) and the velocity profile is linear. Only in this region the shear rate assumes the simple expression:

$$\dot{\gamma} = \frac{U_2 - U_1}{2h^*} \quad \text{Equation 4}$$

Where  $U_1$  and  $U_2$  are the linear velocities of the two rolls. For a power law non-Newtonian fluid the shear stress can be expressed as:

$$\tau = k(\dot{\gamma})^m \quad \text{Equation 5}$$

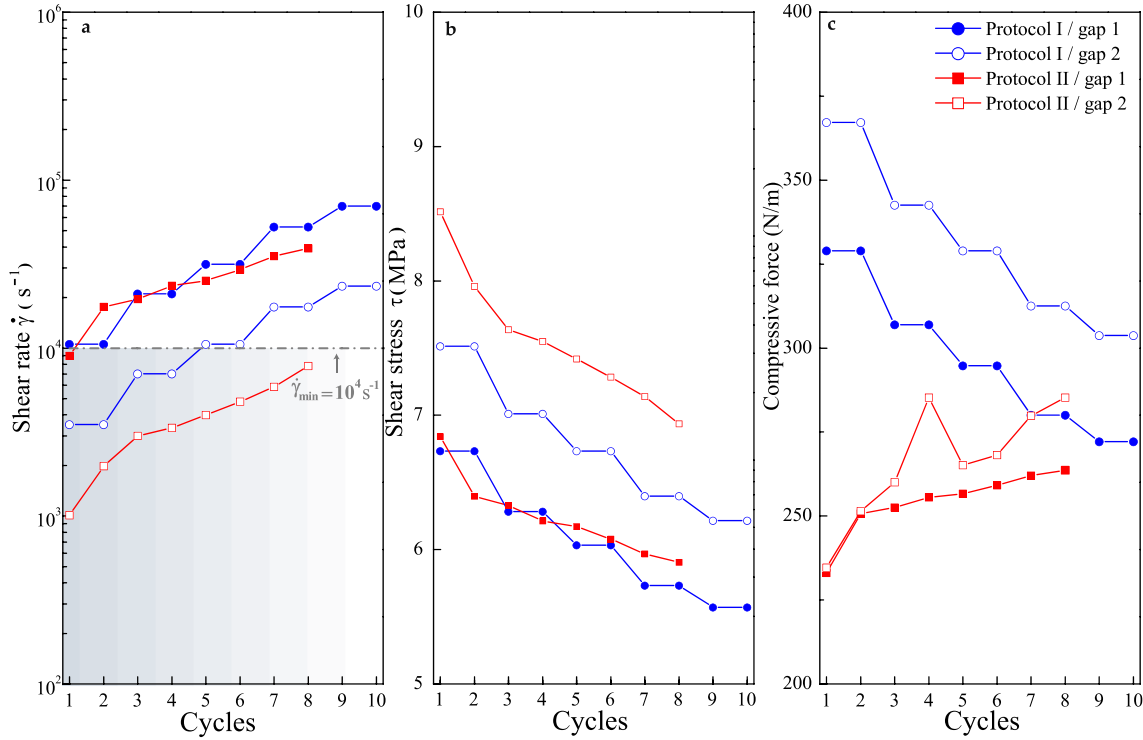
Where  $m$  is the power law index and  $K$  is the power law coefficient. In our case,  $K$  and  $m$  have been measured experimentally (see Supporting Information **Figure S3-4**).

Referring back to our process, it is noted that for each cycle the material goes through two gaps (gap 1, between apron roll and central roll, and gap 2, between central roll and feed roll), hence experiencing two different shear rates and shear stresses. Using Equation 4, it can be seen that Protocol I reaches higher shear rates; it effectively always exceeds, at both gaps and for each cycle, the threshold of  $10^4 \text{ s}^{-1}$  indicated by Paton *et al.*<sup>22</sup>, and approaches values of  $10^5 \text{ s}^{-1}$  in the last 4 cycles. Protocol II, instead, is relatively mild both in terms of shear rates and number of cycles (8 instead of 10). For each cycle, the shear rate at gap 2 never exceeds  $10^4 \text{ s}^{-1}$  (from  $10^3 \text{ s}^{-1}$  to  $10^4 \text{ s}^{-1}$ ) while the shear rate at gap 1 progressively increases from  $10^4 \text{ s}^{-1}$  to  $5 \cdot 10^4 \text{ s}^{-1}$ .

Using Equation 5, it can be shown that the shear stress ranges between 5 and 10 MPa, which is well above the minimum resistance of the graphene stack under pure shear (about 0.25 MPa<sup>33</sup>) (**Figure 2b**).

This shows that both protocols should be able to achieve exfoliation of graphite. However, since Protocol I is characterised by higher shear rates, shear stresses and number of cycles,

compared to Protocol II, it is reasonable to expect thinner graphitic nanoparticles, in agreement with experimental observations (**Figure 2, Table 1**).

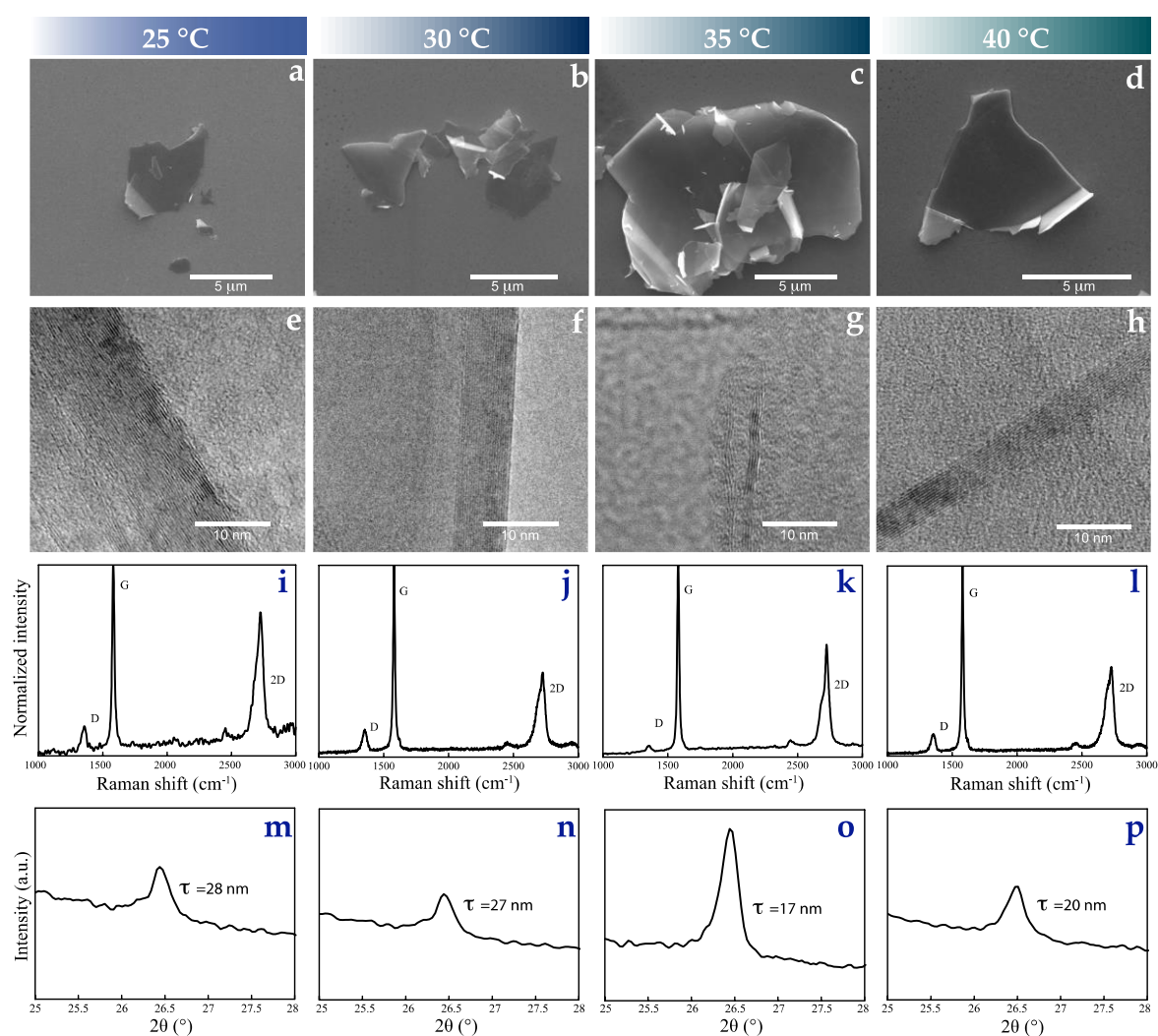


**Figure 2.** (a) Shear rate (b) Shear stress and (c) compressive force (per unit roll width) experienced by the epoxy composites mixtures as a function of different cycles of Protocol I and II.

Unfortunately more severe processing conditions might also induce breaking down of the graphitic particles (reduction in lateral dimension) along with the reduction of their thickness. In fact it is noted that Protocol I is characterised by much higher compressive forces (**Figure 2c**), particularly at the first cycles, which might induce severe fracturing of the natural graphite particles apart from simple exfoliation. This is in agreement with the previous conclusion  $(\eta_d/\eta_f)_{Protocol I} < (\eta_d/\eta_f)_{Protocol II}$  which states that Protocol I uses a higher fraction of energy for platelet fracture rather than exfoliation, compared with Protocol II.

## Effect of temperature on *in-situ* exfoliation of natural graphite in epoxy resin

Another important processing parameter found to significantly affect the *in-situ* exfoliation process was temperature. SEM and TEM micrographs in **Figure 3a-d** show the typical morphology of FLG/GNP particles obtained at different resin temperatures, between 25 °C and 40 °C (Protocol III). The average particle dimensions calculated from a statistical analysis of the SEM and TEM images and the X-ray data are reported in **Table 2**, in analogy with the previous section.



**Figure 3.** Exfoliation through Protocol III. Effect of resin temperature on morphology (SEM and TEM), Raman spectroscopy and XRD for FLG/GNP produced at: 25 °C (a, e, i and m), 30 °C (b, f, j and n), 35 °C (c, g, k and o) and 40 °C (d, h, l and p), respectively.

Interestingly, the FLG/GNP size strongly depends on processing temperature. Moreover this dependency is non-monotonic. The development of thickness and the lateral dimension experience, respectively, a minimum and a maximum in correspondence of a resin temperature of 35 °C. To the best of the authors' knowledge it is the first time that such an effect is reported. Notably, the FLG/GNP particles obtained at 35 °C reach an aspect ratio of up to 1000, 20 folds larger than Protocol I and 50 folds larger than the original natural graphite particles. **Figure 3.i-I** also presents typical Raman spectra for the *in-situ* exfoliated particles obtained at different temperatures. Also in this case the ratio of the intensities of the D and G bands,  $I_D/I_G$  (reported in **Table 2**), is an inverse function of the flakes' lateral dimensions. The  $I_D/I_G$  ratio attains a minimum of 0.07 in correspondence of the largest flakes (*in-situ* exfoliated at 35 °C), suggesting a reduction of edge defects.<sup>29</sup>

**Table 2.** Properties of FLG/GNP particles obtained by Protocol III.

Sample	T (°C)	Length, L (µm)	Thickness by XRD, T <sub>XRD</sub> (nm)	Thickness by TEM, T <sub>TEM</sub> (nm)	Aspect Ratio (L/ T <sub>XRD</sub> - L/ T <sub>TEM</sub> )	$I_D/I_G$
Protocol III - 1	25	3.5 ± 1.5	28 ± 8	14 ± 8	125 - 250	0.16 ± 0.08
Protocol III - 2	30	4.0 ± 1.3	27 ± 6	12 ± 4	150 - 333	0.15 ± 0.04
Protocol III - 3	35	5.2 ± 2.0	17 ± 5	5 ± 4	306 - 1040	0.07 ± 0.05
Protocol III - 4	40	4.6 ± 1.8	20 ± 10	9 ± 5	230 - 511	0.12 ± 0.04

In order to explain these exceptional results, it is noted that an increase in temperature monotonically decreases the epoxy viscosity (see Supporting Information **Figure S4**) which, as a consequence, decreases the shear stress for a given shear rate imposed by the TRM process (**Figure 4**). One would therefore expect the exfoliation efficiency to decrease monotonically with increasing temperature, which is in disagreement with the experimental results of **Figure 3** and **Table 2**. Hence, the reduction in viscosity of the epoxy resin cannot explain the difference in *in-situ* exfoliated FLG/GNP particle sizes.

Another physical property which was found to be significantly affected by the temperature was the surface tension of the epoxy resin used. The surface tension, estimated from contact angle measurements (see Supporting Information. **Figure S5** and Section: **Surface**



**tension**), varied between 65 mJ/m<sup>2</sup> at 25 °C to 30 mJ/m<sup>2</sup> at 40 °C (**Figure 4**). Interestingly, for temperatures between 35-40 °C, the surface tension assumed values of 40 - 50 mJ/m<sup>2</sup>, which coincides with the optimal surface tension range for liquids to disperse graphene, as proposed by Hernandez *et al.*<sup>17</sup> The same authors explained this behaviour by invoking the Hildebrand–Scratchard equation.<sup>38</sup> The latter establishes a relationship between the enthalpy of mixing and the balance of graphene and solvent surface energies, with an energetic minimum expected for solvents whose surface energy matches that of graphene. The same authors could demonstrate and predict the energetic minimum to be in correspondence of a liquid surface tension of 40 - 50 mJ m<sup>-2</sup>.

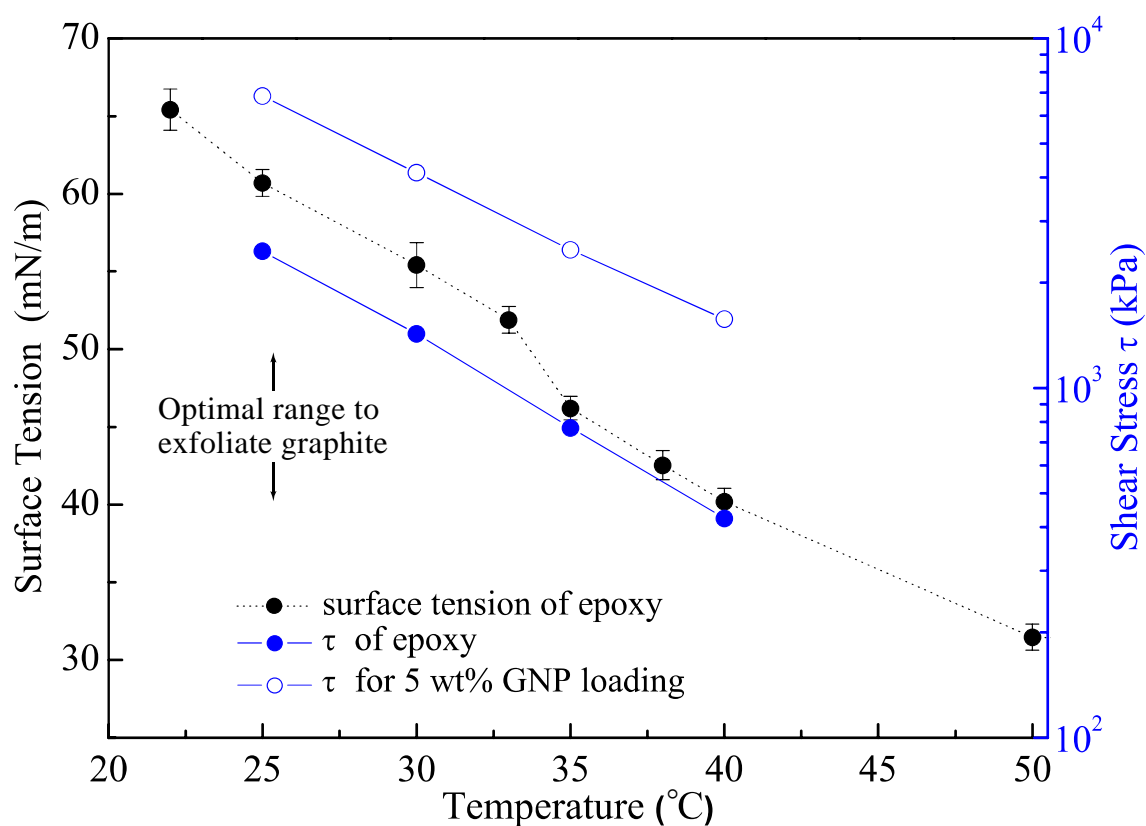
In analogy to the work of Hernandez *et al.*<sup>17</sup> we therefore believe that the maximum FLG/GNP aspect ratio found in our case can be explained by a minimisation of the surface energy difference between graphene and the epoxy resin, which assists the *in-situ* exfoliation process. The matching of the surface energy helps dispersing graphene nanoparticles while they are produced by mechanically exfoliation.

Interestingly, the existence of this optimal temperature in Protocol III is in agreement with our previous energetic model introduced in section 3.1, since  $\gamma_d \propto \gamma_{\text{graphite}} - \gamma_{\text{resin}} \cos \mathcal{G}$ , where  $\mathcal{G}$  is the contact angle (small under large spreading), is minimized. By using the experimental values of aspect ratio found (**Table 2**), it is possible to estimate that the ratio  $\eta_d / \eta_f$  is as high as 15 - 50 for Protocol III-3.

Another approach reported in the scientific literature to interpret improved graphene dispersion, apart from the Hildebrand–Scratchard equation, is based on the Hansen solubility parameters (HSP).<sup>20, 39</sup> HSP are widely used to predict the compatibility between two materials.

For each material, three Hansen parameters can be defined:  $\delta_D$ ,  $\delta_P$  and  $\delta_H$ , which can be located in the 3D Hansen space just as co-ordinates.<sup>39-40</sup> In the Hansen space a HSP distance  $R_a$ , in general between solvent 1 (or any dispersing media) and solute 2 (in our case graphene), can be defined as  $R_a = (4(\delta_{D1} - \delta_{D2})^2 + (\delta_{P1} - \delta_{P2})^2 + (\delta_{H1} - \delta_{H2})^2)^{1/2}$ . The smaller  $R_a$ , the higher the solubility. In other words a good dispersing media should have HSP matching that of graphene.<sup>41</sup> The HSP of graphene<sup>41</sup> has been estimated as  $\delta_D \sim 18 \text{ MPa}^{1/2}$ ,  $\delta_P \sim 9.3 \text{ MPa}^{1/2}$  and  $\delta_H \sim 7.7 \text{ MPa}^{1/2}$ . In a previous publication, using an epoxy resin (bisphenol A-diglycidylether)

similar to the one used in this study, HSP parameters were measured as follows:  $\delta_D \sim 20 \text{ MPa}^{1/2}$ ,  $\delta_P \sim 10 \text{ MPa}^{1/2}$  and  $\delta_H \sim 8 \text{ MPa}^{1/2}$ .<sup>42</sup> It is interesting to notice the relative close similarity of the HSP of graphene and epoxy, demonstrating that in general epoxy resins are potentially good dispersing media for graphene. In this paper we build on this relative compatibility and demonstrate that by fine tuning the temperature the dispersability of graphene in epoxy can be further improved, which in turn assists the *in-situ* exfoliation process.



**Figure 4.** Surface tension and estimated shear stress achieved in Protocol III at different temperatures. The vertical range 40-50 mJ/m<sup>2</sup> represents the optimal range for graphene dispersion/exfoliation in liquids.

### Effect of *in-situ* exfoliation on epoxy nanocomposites properties

As explained in the introduction section, in order to fully exploit the potential of graphene or graphitic nanoparticles in a number of application fields including composites and coatings,

it is necessary to produce thin ( $< 10$  layers, preferably  $< 3-5$  layers) and large nanoparticles (aspect ratios  $> 1000$ , preferably  $> 10000$ )<sup>43</sup> at an industrial-scale. In fact, it is well known from micromechanical composite theories<sup>44</sup> and percolation theories<sup>45-46</sup> that the larger the filler aspect ratio, the higher the mechanical reinforcement is (asymptotically approaching the rule-of-mixture limit), as well as the electrical and thermal conductivity, and the lower the percolation threshold. The filler aspect ratio also plays a fundamental role in a number of other properties, particularly in transport-dominated properties like gas/liquid barrier properties,<sup>47</sup> fire retardancy,<sup>48</sup> corrosion resistance,<sup>49</sup> etc.

In the previous sections, an industrially viable strategy to minimise the thickness and maximise the lateral dimensions of FLG/GNP particle has been presented. **Figure 5.a** compares the morphology (thickness and aspect ratio) of the best *in-situ* exfoliated FLG/GNP particles of this work (Protocol III-3) with the most representative results reported in the literature<sup>17, 21, 23, 29, 34, 50-57</sup> and commercially<sup>58,59</sup>. It can be seen that our *in-situ* exfoliated FLG/GNP have comparable thicknesses than average reported values (5 nm compared to minimum values of 1-3 nm), and generally higher aspect ratios (up to 1000 compared to  $\sim 200$ ). It should be noted that our results are averages obtained without any process (centrifugation,<sup>17</sup> sedimentation,<sup>29</sup> etc.) of particle selection on the basis of their size, as often used and reported in literature. In other words we attain full conversion of the initial natural graphite (100 % yield, **Figure 5.a**) without any losses. To the best of our knowledge no other methodology achieves a comparable combination of production yield and size (aspect ratio and thickness).

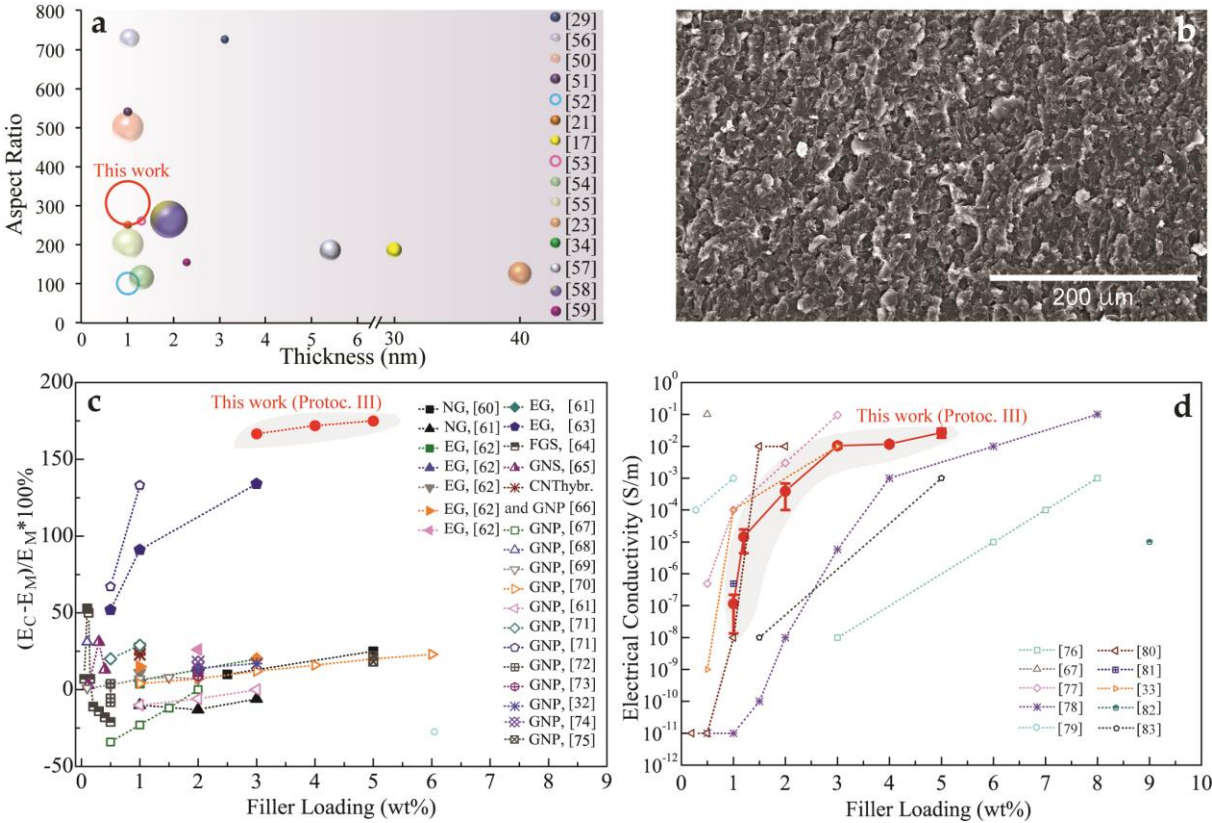
The question remains how these interesting topological features and ‘quality’ of our *in-situ* exfoliated FLG/GNP particles translate into macroscopic properties of epoxy based nanocomposites.

**Figure 5.c** compares the bending elastic moduli of epoxy based nanocomposites filled with FLG/GNP particles obtained with Protocol III-3, as a function of filler content.<sup>33, 60-75</sup> To put these results into context, the mechanical reinforcement achieved (defined as  $(E_C - E_M)/E_M$ , where  $E_C$  and  $E_M$  are the flexural moduli of the composites and neat matrix, respectively) is compared with the best results found in the scientific literature for epoxy/graphitic particles nanocomposites. Interestingly, our FLG/GNP particles achieve the highest mechanical reinforcement ever reported, demonstrating the potential of our *in-situ* exfoliation process in

future applications. This is also the consequence of a very homogenous dispersion of the nanoparticles in epoxy, as shown in **Figure 5.b**.

**Figure 5.d** shows the electrical conductivity values of epoxy based nanocomposites filled with FLG/GNP particles obtained with Protocol III-3, compared with literature.<sup>33, 67, 76-83</sup> With a max electrical conductivity (relative to 5 wt.% of FLG/GNP) exceeding  $10^{-2}$  S/m and a percolation threshold of  $\sim 1$  wt.% our epoxy nanocomposites closely approach the best results ever reported in the literature ( $\sim 10^{-1}$  S/m and  $\sim 0.5$  wt.%, respectively).

It is noted that for sake of brevity only a limited number of results from the scientific and commercial literature could be used for comparison. A more complete collection of experimental data is included in our Supporting Information (**Table S1-2**).



**Figure 5.** a) Comparison of morphological features (thickness, aspect ratio) and yield of *in-situ* exfoliated FLG/GNP produced by Protocol III-3 with graphitic nanoparticles from literature, b) SEM micrograph of the cross-sectional area of an epoxy nanocomposite containing with 5 wt.% FLG/GNP, produced by Protocol III-3, c) relative percentage increase in bending elastic moduli

of epoxy nanocomposite, produced by Protocol III-3, compared with literature, d) Electrical percolation curve of epoxy nanocomposite, produced by Protocol III-3, compared with literature.

## CONCLUSIONS

For the first time the direct *in-situ* exfoliation and dispersion of FLG/GNP into epoxy resins was demonstrated, without the need of any additives, solvents, compatibilisers or chemical treatments. This single step, top-down, scalable and high yield (100% conversion of natural graphite) process promises to alleviate the cost barrier which is currently preventing the industrial uptake of graphene in bulk applications like composites and adhesives. Good quality (low defects;  $I_D/I_G \sim 0.07$ ) FLG/GNP particles with an average aspect ratio greater than 300 and an average thickness of 5 nm were produced by fine tuning two important parameters: shear loading and temperature. Control over the first parameter resulted in an improved particle aspect ratio achieved by balancing the desirable reduction of particle thickness (delamination or exfoliation) with the inevitable particle break-down and reduction of lateral size (fracture). The second parameter was shown to improve the dispersion of the graphene nanoparticles while they are produced by mechanically exfoliation, due to a better matching of the surface energies of the graphite and liquid epoxy.

The above conclusions are interpreted in terms of simple energy balances and geometrical arguments, an analytical model of the TRM process and the Hansen solubility parameters.

The optimisation of the *in-situ* FLG/GNP particles morphology and ‘quality’ had a clear impact on macroscopic physical properties of the epoxy nanocomposites. An electrical percolation threshold of about 1 wt.% and an electrical conductivity exceeding  $10^{-2}$  S/m were measured for particles obtained with the best processing conditions found. Aspect ratio had also a significant effect on mechanical properties, with the bending elastic modulus increasing by ~160 % for 4 wt.% of optimised FLG flakes, corresponding to the highest mechanical reinforcement ever reported for epoxy/graphene nanocomposites.

## ASSOCIATED CONTENT

## Supporting Information

This material is available free of charge *via* the Internet at <http://pubs.acs.org>.

Supporting Information content: Modelling of the TRL calendaring process; Rheology of epoxy resin; Surface tension measurements; Morphology of particles; Review of graphitic nanoparticles and their composites properties; Summary of processing parameters.

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### Notes

The authors declare no competing financial interest.

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## Supporting Information

### *In-Situ* Exfoliation of Graphene in Epoxy Resins:

#### **a Facile Strategy to Efficient and Large Scale Graphene Nanocomposites**

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### Review of graphitic nanoparticles and their composites properties

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**Table S2.** A summary of the sizes of GNPs reported so far in commercial market.

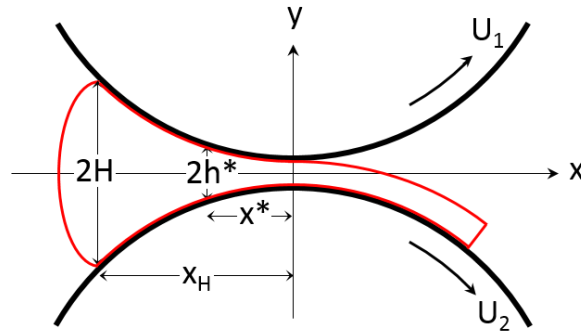
### Summary of processing parameters

**Table S3.** Processing parameters used in Three Roll Mill processing.

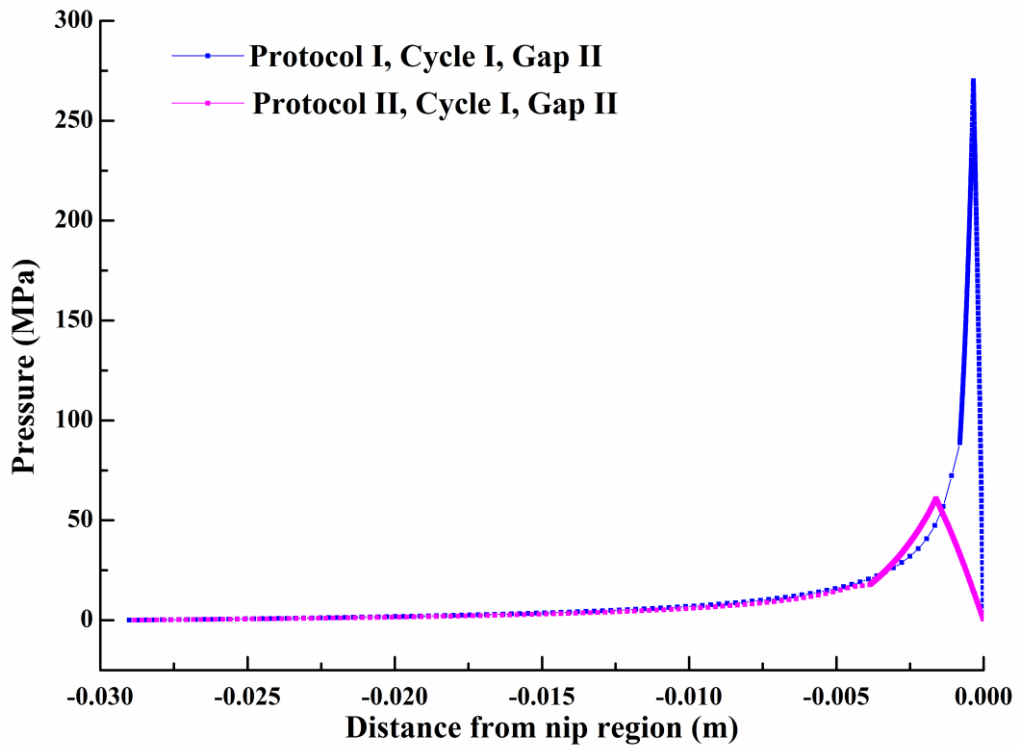
**Table S4.** Experimental trials of different processing parameters of TRM.

### Reference

## Modelling of the TRM calendaring process



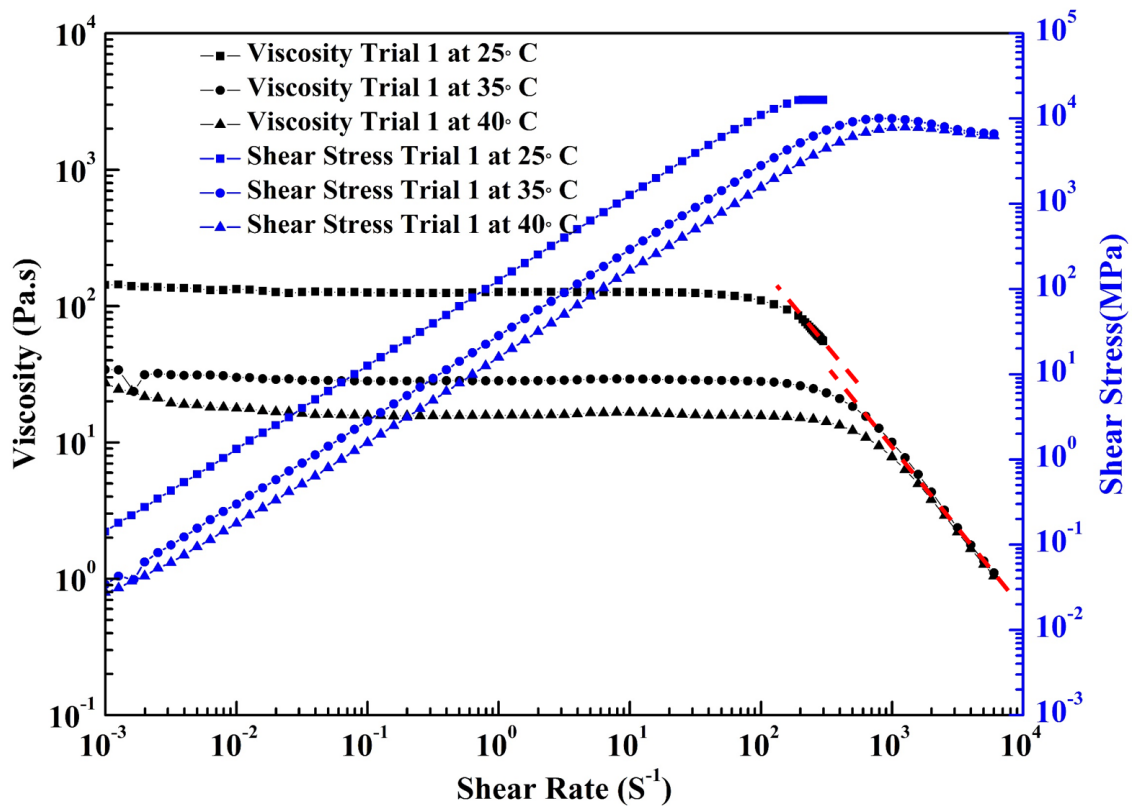
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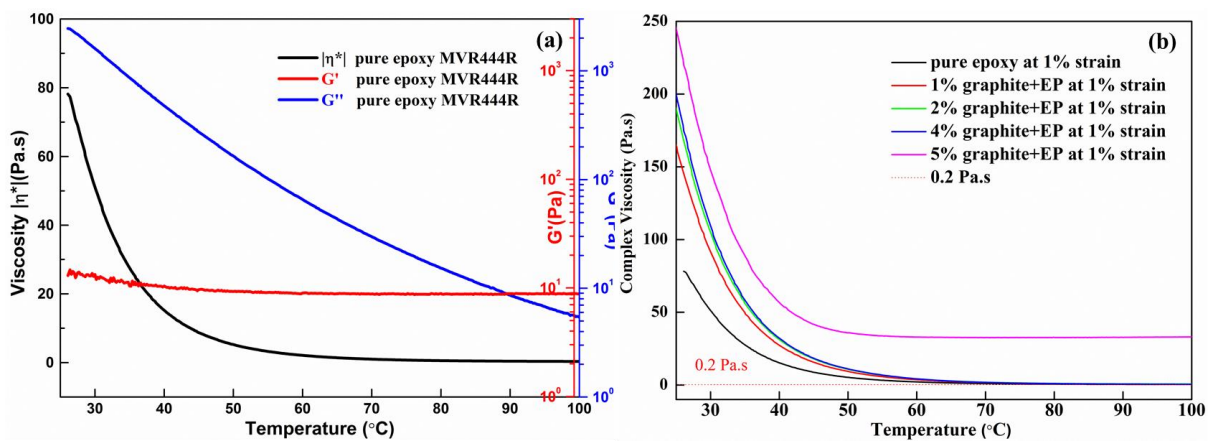
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## Rheology of epoxy resin



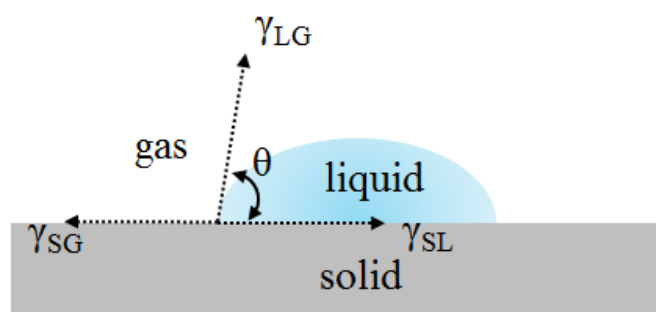
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## Surface tension

Drop shape analyser was used to measure the interfacial tension between the liquid epoxy and glass substrate. Surface energies were calculated from contact angle data of sessile drops. Base line and sessile droplet fitting were included for comparison. The most complicated, but also the theoretically most exact method for calculating the contact angle is the Young-Laplace equation <sup>1, 2</sup>. A given system of solid, liquid, and gas at a given temperature and pressure has a unique equilibrium contact angle. Indices S, L and G stand for “solid”, “liquid” and “gas”; the symbols  $\gamma_{SG}$  and  $\gamma_{LG}$  describe the surface tension of two phases (solid-liquid and liquid-gas, respectively); and  $\theta$  stands for the contact angle, corresponding to the angle between vectors  $\gamma_{LG}$  and  $\gamma_{SL}$ .



**Figure S5.** Schematic of a liquid drop showing the quantities according to the Young's equation.

The shape of a liquid/gas interface is determined by the Young-Laplace equation, with the contact angle playing the role of a boundary condition via Young's equation:

$$\gamma_{SG} = \gamma_{SL} + \cos\theta * \gamma_{LG} \quad (1)$$

During the experiment, we use the same glass substrate to keep the same surface roughness, and try to avoid potential contamination, or influence of possibly varying ambient conditions. In this method the complete drop contour is evaluated; the contour fitting includes a correction which takes into account the fact that it is not just interfacial effects which produce the drop shape, but that the drop is also distorted by the weight of the liquid it contains. After the successful fitting of the Young-Laplace Equation the contact angle is determined as the slope of the contour line at the three phases contact point. However, the calculation is only reliable for contact angles above 30°. Moreover, this model assumes a symmetric drop shape.

In order to make experiments easier, we choose a solvent, ethylene glycol, as a reference, whose surface tension is 47.70 N/m at 20 °C, with the boiling point at 197.3 °C (difficult to evaporate at 20 °C during the experiment procedure). Several  $\mu\text{l}$  ethylene glycol can maintain a good axisymmetric droplet profile on the glass substrate. With the measured volume, contact angle, the interfacial tension between the droplet and glass substrate can be calculated. According to Equation (1), the equilibrium condition can be described as follows,

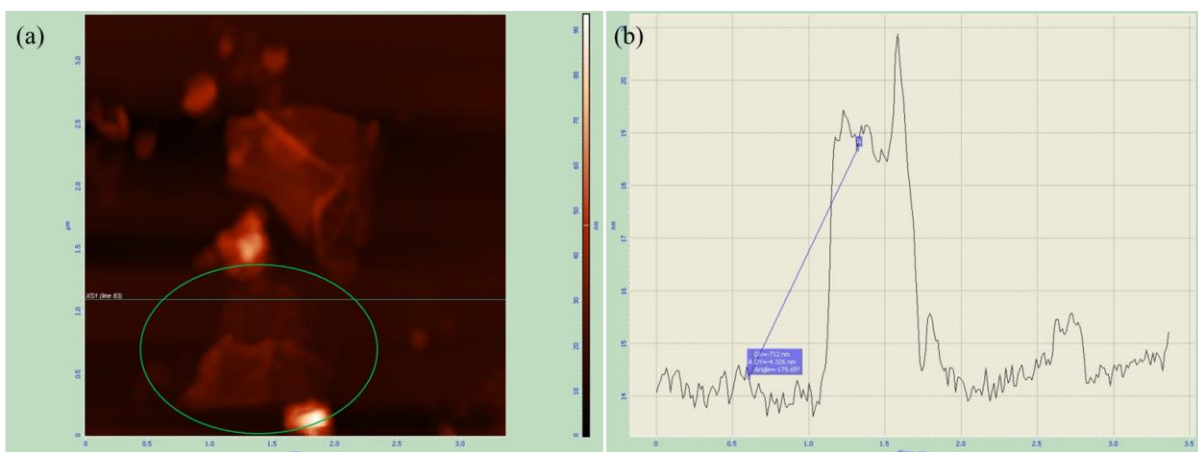
$$\gamma_{\text{Glass}} = \gamma_{\text{IFT1(Glass,Ethylene glycol)}} + \cos\theta_1 * \gamma_{\text{Ethylene glycol}} \quad (2)$$

$$\gamma_{\text{Glass}} = \gamma_{\text{IFT2(Glass,Epoxy)}} + \cos\theta_2 * \gamma_{\text{Epoxy}} \quad (3)$$

Where,  $\gamma_{\text{Glass}}$ ,  $\gamma_{\text{Ethylene glycol}}$ ,  $\gamma_{\text{Epoxy}}$  represent the surface tension of the glass substrate, ethylene glycol and epoxy resin MVR444R, respectively.  $\gamma_{\text{IFT1(Glass,Ethylene glycol)}}$  and  $\gamma_{\text{IFT2(Glass,Epoxy)}}$  respectively, represent the interfacial tension of the ethylene glycol droplet and epoxy resin MVR444R with the glass substrate.  $\theta_1$  and  $\theta_2$  are the contact angles of ethylene glycol and epoxy resin with glass substrate under equilibrium condition. The surface tension of epoxy resin MVR444R is calculated as follows,

$$\gamma_{\text{Epoxy}} = \frac{(\gamma_{\text{IFT1}} - \gamma_{\text{IFT2}}) + \cos\theta_1 * \gamma_{\text{Ethylene glycol}}}{\cos\theta_2}, \quad \gamma_{\text{Epoxy}} = \frac{(\gamma_{\text{IFT1}} - \gamma_{\text{IFT2}}) + 47.7 * \cos\theta_1}{\cos\theta_2} \quad (4)$$

## Morphology of particle



**Figure S6.** (a) Semicontact mode AFM image of GNP particle obtained from Proccol III (35 °C). (b) Thickness measurement of the obtained GNP particle obtained from Proccol III (35 °C) (thickness  $t = 4.326$  nm).

## Review of graphitic nanoparticles and their composites properties

**Table S1.** A summary of the sizes of GNPs reported so far in scientific literature.

Matrix/Substrate	Carbon source	After Exfoliation Particle Dimension	Fabrication of the Filler	Ref
NMP	Graphite flakes	6-12 layers, ~ 1.0-3.5 $\mu\text{m}$	Bath /Probe sonication	3
NMP	Graphite powder	63 mg/mL, 3 layers, ~1.0* 0.5 $\mu\text{m}$	Bath sonication/Probe sonication	4
90 wt.% water/[BMIm]Cl electrolyte	Graphite	Carbon nanoribbons (10 nm* (60 $\pm$ 20) nm) In water-rich ILs, the size of the carbon nanoparticles is larger (8-10 nm); In pure ILs, carbon nanoparticles are 2-4 nm.	Ionic liquid-assisted electrochemical exfoliation	5
Ni film on a SiO <sub>2</sub> /Si substrate	Methane -CH <sub>4</sub>	1 to ~12 graphene layers.	CVD on polycrystalline Ni films	6
DMF	Expanded graphite (EG)	Yield of 4–5 wt.% , thickness of graphene layer, decreases from 6–7 nm to 0.75–1.07 nm	Ultra sonication and centrifugation	7
DMF	Highly oriented pyrolytic graphite (HOPG)	Lateral size ~ several hundred nm, thickness: several nm, low yield	Bath sonication and centrifugation	8
potassium permanganate, sodium nitrate, and sulfuric acid	HOPG	Lateral size ~ 10 $\mu\text{m}$ , 100% monolayer , thickness 0.96 nm	Chemical exfoliation by Hummers method	8
H <sub>2</sub> SO <sub>4</sub> solution	HOPG	Lateral size ~ 1.0-2.0 $\mu\text{m}$ , thickness 2.1 nm	Electrochemical expansion and exfoliation	8
Ionic liquid and water as electrolyte,	Graphite Rod	Several hundred nm, thickness: 1.1 nm	Electrochemical exfoliation	9
LiClO <sub>4</sub> and propylene carbonate as electrolyte, -15 $\pm$ 5 V	Graphite powder or HOPG	Thickness 1.5 nm, lateral size 1-2 $\mu\text{m}$	Electrochemical exfoliation assisted by >10 h	10

			sonication	
0.48 g/L H <sub>2</sub> SO <sub>4</sub> applying DC bias from -10 V to +10 V	Natural graphite flakes or HOPG	Thickness 1.5 nm, lateral size several μm	Electrochemical exfoliation	11
0.1 MSDS aqueous solution, 12 h from -1 V to 2 V.	Graphite Rod	Thickness 1.0 nm, lateral size ~ several hundred nm	Electrochemical exfoliation	12
1 M HClO <sub>4</sub> solution, 20 min from -1.6 V to 2 V	Laminated graphite foil	Lateral size several μm	Electrochemical exfoliation	13
Potassium permanganate, sodium nitrate, and sulfuric acid	Natural graphite flakes	Thickness 1.2 nm, lateral size ~ several hundred μm	Chemical exfoliation by Hummers method	14
Potassium permanganate, sodium nitrate, and sulfuric acid	Natural graphite particles or HOPG	Thickness 0.93 nm, lateral size 10-20 μm	Chemical exfoliation by Hummers method	15
Potassium permanganate, sodium nitrate, and sulfuric acid	Acid intercalation graphite flakes	Thickness 0.94 nm, lateral size 11-14 μm	Chemical exfoliation by Hummers method, microwave assisted expansion	16
NMP	Graphite powder	Thickness 3 layers, lateral size : several hundred nm, 4.0 wt.% monolayer	Bath sonication.	17
2 wt.% sodium cholate aqueous solution	Graphite flakes	thickness 1-2 nm, lateral size 100 nm	Horn sonication	18
Water with 2 wt.% surfactant Sodium dodecylbenzene sulfonate (SDBS)	Graphite powder	>40% of these flakes had <5 layers, ~3% of flakes consisting of monolayers, thickness 1 nm, lateral size 250 nm	Bath sonication	19
Organic solvents such as N-methyl-pyrrolidone	Graphite	1 wt.% monolayer	Bath sonication	20
Water/acetone mixtures	Graphite	0.21 mg/ ml ~50% of the nanosheets < 1 nm thick	Mild sonication for 12 h	21
DMF	Multi-layered graphite nanosheets	0.8-1.8 nm	Wet ball milling	22

A variety of organic solvents	Graphite nanosheets	Thickness 0.8~1.8 nm, lateral size 100–200 nm	Ball-milling	23
Polystyrene	Graphite nanoplatelets	Mono- and few-layer graphene, ~ 1.74 nm	Ball-milling	24
PVC dispersed in dioctyl phthalate DOP (adhesive)	Natural graphite	1.13-1.41 nm	Three Roll Mill	25
silicone polymer	Graphite nanoplatelets	Thickness ~ 5 to 35 nm	Three Roll Mill	26
Sylgard184SiliconeElastomer	graphite nanoplatelets	Thickness 20-200 nm, lateral size 5 $\mu$ m	Dual asymmetric centrifuge mixing, Speed Mixer	27

**Table S2.** A summary of the sizes of GNPs reported so far in commercial market.

Graphene producer	Graphene product	Details/Quality	
Graphene Platform	Silver coated graphene	Silver Decorated Graphene with 30 wt.% , Particle size : 4.5 $\mu$ m	
		Silver Decorated Graphene with 70 wt.% , Particle size : 7.2 $\mu$ m	
	3D graphene	Grown on Cu/Ni Foam, continuous layer with few small multilayer islands coverage exceeding 95%.	
	Graphene dispersion	in NMP with non-ionic dispersant in NMP no surfactant	Different concentration 0.1,1.0,10,50,100 mg/ml, Purity : >99%, 1~10 Layers : >70%, >30 Layers : <5%
in water with non-ionic dispersant		Different concentration 0.1,1.0,10 mg/ml, Purity : >99%, 1~10 Layers : >70%, >30 Layers : <5%	
Thomas Swan Advanced Materials	Elicarb® Graphene	Graphene powder	few-layer graphene flakes with an average of 5-7 layers.
		Graphene Dispersion	A water/surfactant dispersed GNP at 1g/l.
ACS Material	Graphene Series	Single Layer Graphene, surface area (g/m <sup>2</sup> ): 400~1000; Electrical resistivity ( $\Omega$ ·cm) $\leq$ 0.30	
		Nitrogen-doped Graphene 1-5 atomic layer , Lateral size : 0.5-5 $\mu$ m; surface area (g/m <sup>2</sup> ): 500~700 ; Conductivity (S/m) >1000	
		Industrial-Quality Graphene, Thickness (nm) $\leq$ 3.0; surface area (g/m <sup>2</sup> ): ~600; Electrical resistivity ( $\Omega$ ·cm) $\leq$ 0.30	

		Carboxyl Graphene, Diameter 1~5 $\mu\text{m}$ , thickness 0.8~1.2 nm, Carboxy ratio ~ 5.0%, Purity ~ 99%
		Carboxyl Graphene, Carboxyl Graphene Water Dispersion Diameter 1~5 $\mu\text{m}$ , thickness 0.8~1.2 nm, Carboxy ratio ~ 5.0% Purity ~ 99%
		Graphene Oxide Diameter 1~5 $\mu\text{m}$ , thickness 0.8~1.2 nm, single layer ratio ~ 99%. Purity~ 99%. Diameter 1~15 $\mu\text{m}$ , thickness 0.8-1.2 nm
		Graphene Oxide, High Surface Area Graphene Oxide Diameter 1~5 $\mu\text{m}$ , thickness 0.8~1.2 nm, single layer ratio ~ 99%. Purity~ 99%.
		Single Layer, Oxide Ethanol Dispersion, Flake size: 0.5-2.0 $\mu\text{m}$ ; thickness: 0.6-1.2 nm; Single-layer Ratio: >80%
ACS Material	Graphene Series	Single Layer Graphene Oxide Water Dispersion (1) 10 mg/ml, 100 ml (1 g), Flake size: 0.5-2.0 $\mu\text{m}$ ; Thickness: 0.6-1.2 nm; Single-layer Ratio: >80% (2) 10 mg/ml, 100 ml (0.5 g), Flake size: 500 nm; Thickness: 0.6-1.2 nm; Single-layer Ratio: >80%
		Diameter: ~5 $\mu\text{m}$ ; Thickness: 2-10 nm; surface area ( $\text{g}/\text{m}^2$ ): 20-40 , Conductivity: 80000 S/m
		Graphene Film-Super Paper, Diameter: 40 mm, thickness: 20 $\mu\text{m}$ , Conductivity: 2000 S/m
		Graphene Oxide Film, Diameter: 40 mm, thickness: 20 $\mu\text{m}$ ; Non-conductive, $8 \times 10^{-2}$ S/m
		Aminated Graphene, Conductivity: 6.36 S/m
	CVD Graphene	Trivial Transfer Graphene, Predominantly single-layer graphene; Transparency: >95%
		3D Graphene Foam, Sheet Resistance: <600 $\Omega/\text{sq}$
		Graphene on Copper Foil, Sheet Resistance: <600 $\Omega/\text{sq}$
		Graphene on Si 1) Super large size graphene on copper foil up to 30 cm x 20 cm; 2) Double or multi-layer graphene; 3) transferred onto silicon substrate; Sheet Resistance: <600 $\Omega/\text{sq}$ ; Transparency: >95%
		Graphene on $\text{SiO}_2$ 1) Super large size graphene on copper foil up to 30 cm x 20 cm; 2) Double or multi-layer graphene; 3) transferred onto silicon dioxide substrate; Sheet Resistance: <600 $\Omega/\text{sq}$ ;

		Transparency: >95%
		Graphene on PET 1) Super large size graphene on copper foil up to 30 cm x 20 cm; 2) Double or multi-layer graphene; 3) Graphene transferred onto PET substrate
		Graphene on Plastic, Graphene transferred to Plastic substrate (a polymer mainly containing PET <10%)
		Graphene on Quartz, Single Layer Graphene on Quartz Substrate; Sheet Resistance: <600 Ω/sq; Transparency: >95%
		Multi-layer , Predominantly Double- or Multi-Layer Graphene; Sheet Resistance: <600 Ω/sq; Transparency: >95%
		PMMA-coated , Pretreated Graphene-PMMA Coated; Sheet Resistance: <600 Ω/sq; Transparency: >95%
	Graphene Quantum Dots	Aminated Graphene Quantum Dots, Solution, Colorless solution; PL peak: 440 nm; Particle Size: <5 nm; Concentration: 1 mg/ml (available up to 20 mg/ml);Solution: Water
		Blue Luminescent Quantum Dots, Quantum Dots Size 15 <nm, Thickness 0.5 ~ 2 nm, Purity ~ 80%, concentration 1mg/ml.
		Carboxylated Graphene Quantum Dots, Solution, Colorless solution; PL peak: 487 nm; Particle Size: <10 nm; Concentration: 1 mg/ml (available up to 20 mg/ml);Solution: Water
		Carboxylated Graphene Quantum Dots, pale yellow powder; PL peak: 487 nm; Particle Size: <10 nm.
		Chlorine Functionalized Graphene Quantum Dots, Solution, Colorless solution; PL peak: 452 nm; Particle Size: <6 nm. Concentration: 1 mg/ml (available up to 2 mg/ml), Solution: Water, Containing a little ethylene glycol
		Green Graphene Quantum Dots, Solution, Colorless solution; PL peak: 530 nm; Particle Size: <6 nm. Concentration: 1 mg/ml (available up to 2 mg/ml), Solution: Water, Containing a little DMF
		Hydroxylated Graphene Quantum Dots, Solution, Colorless solution; PL peak: 375 nm; Particle Size: <6 nm. Concentration: 1 mg/ml (available up to 2 mg/ml), Solution: Mixture of water and ethylene glycol
XG Science	xGnP bulk dry powder	Grade C, an average particle diameter of less than 2 microns. Average surface areas are 300, 500 and 750 g/m <sup>2</sup> .
		Grade H, a typical surface area of 60 to 80 g/m <sup>2</sup> , available with average particle diameters of 5, 15 or 25 μm.



		Grade M, ~ 6 to 8 nm , surface area of 120 to 150 g/m <sup>2</sup> , available with average particle diameters of 5, 15 or 25 μm.
	xGnP dispersions	Aqueous: xGnP® Graphene Nanoplatelets can be dispersed into water with probe sonication or high shear mixing.
		Organic solvents, Suggested solvents include NMP, DMF, THF, toluene, ethyl acetate, isopropanol, ethanol, acetone, methyl ethyl ketone (MEK) and chloroform, 2 amino-butane and other polar solvents.
		Resins and custom
Advanced Graphene Products	MONOLAYER Graphene	HSMG™ on PMMA. Transparent film, Optical transmittance at 550 nm: >97%; Coverage: >95%; 1 layer;
	MONOLAYER Graphene	Thickness (theoretical): ~0.345 nm; sheet resistance: 220-800 Ohm/sq; Grain size: Up to 1 mm
		HSMG™ monolayer on Si, Substrate: Si(B) (111) type p; Thickness 300 μm ; Single side polished; Res: 9-12 ohm/cm
	MULTILAYER Graphene	HSMG™ on PMMA., Transparent film, Optical transmittance at 550 nm: >85%; Coverage: >95%; 3-5 layers; sheet resistance < 800 Ohm/sq; Grain size: Up to 1 mm
		HSMG™ monolayer on Si, Substrate: Si(B) (111) type p; Thickness 300 μm ; Single side polished; Res: 9-12 ohm/cm
NanoXplore	NXE-Graphene	Grade A: Purity: 96% by weight, Average Specific surface area : 25-30 g/m <sup>2</sup> , 4-5 layers; Average sheet diameter : 5-20 μm. Highly OH edge functionalized
		Grade B: Purity: 96% Average Specific surface area : 10-15 g/m <sup>2</sup> , 2-3 layers; Average sheet diameter : 0.5-5 μm. Highly OH edge functionalized
		Grade C: Purity: 96% Average Specific surface area : 200 g/m <sup>2</sup> , 4-5 layers; Average sheet diameter : 5-20 μm,
		Grade D Purity: 96% Average Specific surface area : 10-15 g/m <sup>2</sup> , 2-3 layers; Average sheet diameter :0.5-5 μm,
	NXE-Graphite Composite	Grade E Purity: 96% by weight, Average Specific surface area :7-9 g/m <sup>2</sup> , 5-20 μm graphene sheets mixed with large natural graphite flakes
	NXE-Graphene Partially Oxidized	Grade F1: Purity: 96% by weight, Average Specific surface area : 100 g/m <sup>2</sup> , 2-3 layers, 200-500 nm, C content: 75% (with 20% Oxygen), Impurity: 2 wt.%, Humidity: 2 wt.%, Low defect density
	NXE-GO	Grade F2 Purity: 96% by weight, Average Specific surface area : 100 g/m <sup>2</sup> , 2-3 layers, 100-200 nm, C content: 60% (with 30% Oxygen),

<i>RS MINES</i>	Reduced GO (RSrGO) Paste	highly oxidised, highly conductive reduced graphene oxide paste, multiple uses including the enhancement of energy storage devices, conductive additive for polymers, and of course to make single to few layer graphene.	
Graphenea	Suspended Monolayer Graphene	on Cavities, Substrate size up to 1.5 x 1.5 cm, Substrate withstand 450 °C Temperature, Cavity size up to 30 µm, Minimum cavity depth: 500 nm, Film, transparent; transparency >97%, 1 layer, thickness :0.345 nm, Grain size: Up to 10 µm	
	Monolayer Graphene	on SiO <sub>2</sub> /Si or Cu or SiO <sub>2</sub> /S or PET or Quartz Film, Transparency: >97 %, Coverage: >95%, Thickness (theoretical): 0.345 nm, Grain size: Up to 10 µm	
	Bilayer Graphene	on SiO <sub>2</sub> /S, Transparency >94%, Appearance (Form): Film, Coverage >95%, 2 layer, Thickness : 0.69 nm, Grain size: Up to 10 µm	
	Trilayer Graphene	on SiO <sub>2</sub> /S, Transparency >92%, Appearance (Form): Film, Coverage >95%, 3 layer, Thickness : 1.035 nm, Grain size: Up to 10 µm	
	Suspended Monolayer Graphene	on TEM Grids (Quantifoil Gold) , Film, Transparency: >97 %, Coverage: >95%, Thickness : 0.345 nm, Grain size: Up to 10 µm	
	Graphene Oxide		Form: Dispersion of graphene oxide sheets, Sheet dimension: Variable, Colour: Yellow-brown, Odour: Odourless Dispersibility: Polar solvents, Solvent: Water, pH: 2,2 - 2,5 Concentration: 4 mg/mL, Monolayer content (measured in 0.5 mg/mL): >95% (*)
			Form: Dispersion of graphene oxide sheets, Solvent: Water, Concentration: 0.5 mg/mL, Monolayer content
	Reduced Graphene Oxide	Form: Powder, Sheet dimension: Variable, Colour: Black, Odour: Odourless, Solubility: Insoluble Dispersability: It can be dispersed at low concentrations (<0.1 mg/mL) in NMP, DMSO, DMF	
	GO Film	Diameter: 4 cm, Thickness: 12-15 µm, Non-conductive	
Angstrom Materials	Graphene and GO Dispersions	N002-PS-0.5 Graphene Oxide Solution Water Content (percent): ≥ 99.50, Average Z Dimension (nm): 1.0 – 1.2, Average X & Y Dimensions (um): 0.554	
		N002-PS-1.0 Graphene Oxide Solution Average Z Dimension: 1-1.2 nm , (Single Layer GO), Average X-Y Dimension: ~ 500 nm	
	Graphene and GO Powder	N002-PDE Graphene Oxide Powder Few Layer Graphene Oxide, 2-3 nm , lateral size ≤ 7 µm, Specific Surface	

		Area (g/m <sup>2</sup> ): ≥ 400
		N008-P-40 Polar Graphene Powder Average Z Dimension (nm): 50 – 100, Average X & Y Dimensions (um): ≤ 10, Specific Surface Area (g/m <sup>2</sup> ): 20-40
		N008-P-10 Polar Graphene Powder Average Z Dimension (nm): 50 – 100, Average X & Y Dimensions (um): ≤ 7, Specific Surface Area (g/m <sup>2</sup> ): ≤ 40
		N008-N Pristine Graphene Powder Average Z Dimension (nm): 50 – 100, Average X & Y Dimension (um): 5, Specific Surface Area (g/m <sup>2</sup> ): ≤ 30
		N006-P Polar Graphene Powder Average Z Dimension (nm): 10 – 20, Average X & Y Dimensions (um): 5, Specific Surface Area (m <sup>2</sup> /g): ≥ 15
		N002-PDR Few Layer Graphene Powder Less than 3 layers, Average X & Y Dimensions (um): ≤ 10, Specific Surface Area (m <sup>2</sup> /g): 400 – 800

### Summary of processing parameters

**Table S3.** Processing parameters used in Three Roll Mill processing.

Processing Parameters	Levels									
	1	2	3	4	5	6	7	8	9	10
I. Shear Rate	5μm, 30 rpm	5 μm, 60 rpm	5 μm, 90 rpm	5 μm, 150 rpm	5μm, 200 rpm	120/40μm, 200 rpm	60/20 μm, 200 rpm	30/10 μm, 200 rpm	15/5 μm, 200 rpm	5N/mm 200 rpm
II. Filler concentration	1.0 wt.%	2.0 wt.%	3.0 wt.%	4.0 wt.%	5.0 wt.%	-	-	-	-	-
III. Temperature	25 °C	30 °C	35 °C	40 °C	-	-	-	-	-	-
IV. Number of cycles	1	2	3	4	5	6	7	8	9	10
V. Direct (D) Vs Masterbatch (M) + Dilution (D)	D	M+D	-	-	-	-	-	-	-	-

**Table S4** Experimental trials of different processing parameters of TRM.

		I										II					III				IV										V			
		1	2	3	4	5	6	7	8	9	10	1	2	3	4	5	1	2	3	4	1	2	3	4	5	6	7	8	9	10	1	2		
I	1										X	X	X	X	X	X				X	X												X/O	
	2										X	X	X	X	X	X						X	X										X/O	
	3										X	X	X	X	X	X								X	X							X/O		
	4										X	X	X	X	X	X										X	X					X/O		
	5										X	X	X	X	X	X												X	X			X/O		
	6										X	X	X	X	X	X				O/T												X/O		
	7										X	X	X	X	X	X					O/T											X/O		
	8										X	X	X	X	X	X																	X/O	
	9										X	X	X	X	X	X																	X	
	10										X	X	X	X	X	X						O/T	O/T	O/T	O/T	O/T	O/T	O/T	O/T				X	
II	1					O	O			O						X/O				X/O	X/O	X/O	X/O	X/O	X/O	X/O	X/O	X/O	X	X		X/O		
	2					O	O			O						X/O				X/O	X/O	X/O	X/O	X/O	X/O	X/O	X/O	X/O	X	X		X/O		
	3					O/T	O/T			O/T						X/O	T	T	T	X/O	X/O	X/O	X/O	X/O	X/O	X/O	X/O	X/O	X	X		X/O		
	4					O/T	O/T			O/T						X/O	T	T	T	X/O	X/O	X/O	X/O	X/O	X/O	X/O	X/O	X/O	X	X		X/O		
	5					O/T	O/T			O/T						X/O	T	T	T	X/O	X/O	X/O	X/O	X/O	X/O	X/O	X/O	X/O	X	X		X/O	O	
III	1					O/T	O/T			O/T	X/O	X/O	O/T	O/T	O/T					O/T	O/T	O/T	O/T	O/T	O/T	O/T	O/T	O/T				O/T		
	2					O/T	O/T			O/T			O/T	O/T	O/T					O/T	O/T	O/T	O/T	O/T	O/T	O/T	O/T	O/T				O/T		
	3					O/T	O/T			O/T			O/T	O/T	O/T					O/T	O/T	O/T	O/T	O/T	O/T	O/T	O/T	O/T				O/T		
	4					O/T	O/T			O/T			O/T	O/T	O/T					O/T	O/T	O/T	O/T	O/T	O/T	O/T	O/T	O/T				O/T		
IV	1	X				O/T					X/O	X/O	X/O/T	X/O/T	X/O/T	X/O	O/T	O/T	O/T													X/O/T		
	2	X					O/T				X/O	X/O	X/O/T	X/O/T	X/O/T	X/O	O/T	O/T	O/T													X/O/T		
	3		X							O/T	X/O	X/O	X/O/T	X/O/T	X/O/T	X/O	O/T	O/T	O/T													X/O/T		
	4		X							O/T	X/O	X/O	X/O/T	X/O/T	X/O/T	X/O	O/T	O/T	O/T													X/O/T		
	5			X						O/T	X/O	X/O	X/O/T	X/O/T	X/O/T	X/O	O/T	O/T	O/T													X/O/T		
	6			X						O/T	X/O	X/O	X/O/T	X/O/T	X/O/T	X/O	O/T	O/T	O/T													X/O/T		
	7				X					O/T	X/O	X/O	X/O/T	X/O/T	X/O/T	X/O	O/T	O/T	O/T													X/O/T		
	8				X					O/T	X/O	X/O	X/O/T	X/O/T	X/O/T	X/O	O/T	O/T	O/T													X/O/T		
	9					X					X	X	X	X	X	X																X		
	10					X					X	X	X	X	X	X																X		
V	1	X	X	X	X	X	X	X	X	X	X/O	X/O	X/O	X/O	X/O	X/O	X/O	X/O	X/O	X/O	X/O	X/O	X/O	X/O	X/O	X/O	X/O	X/O	X	X				
	2									O	O					O/T	O/T	O/T	O/T	O/T														

X-Process I, O- process II, T-Temperature Controlled samples, using Process II.

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