Organic solvent-based graphene oxide liquid crystals: A facile route toward the next generation of self-assembled layer-by-layer multifunctional 3D architectures

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Keywords
generation, self, assembled, crystals, organic, solvent, facile, graphene, oxide, layer, liquid, multifunctional, architectures, 3d, route, toward, next

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Organic Solvent-Based Graphene Oxide Liquid Crystals: A Facile Route Towards Next Generation of Self-Assembled Layer-by-Layer Multifunctional 3D Architectures

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ABSTRACT. We introduce soft self-assembly of ultra-large liquid crystalline (LC) graphene oxide (GO) sheets in a wide range of organic solvents overcoming the practical limitations imposed on LC GO processing in water. This expands the number of known solvents which can support amphiphilic self-assembly to ethanol, acetone, tetrahydrofuran, N-dimethylformamide,
N-cyclohexyl-2-pyrrolidone and a number of other organic solvents, many of which were not known to afford solvophobic self-assembly prior to this report. The LC behavior of the as-prepared GO sheets in organic solvents has enabled us to disperse and organize substantial amounts of aggregate-free single walled carbon nanotubes (SWNTs, up to 10 wt. %) without compromise in LC properties. The as-prepared LC GO-SWNTs dispersions were employed to achieve self-assembled layer-by-layer multifunctional 3D hybrid architectures comprising of SWNTs and GO with unrivalled superior mechanical properties (Young’s modulus in excess of 50 GPa and tensile strength of more than 500 MPa).

KEYWORDS: Graphene Oxide, Liquid crystal, Liquid crystal in organic solvent, Self-assembled composite, Layer-by-layer composite.

Lyotropic liquid crystalline (LC) phases in dispersions containing two dimensional graphene and graphene oxide (GO) sheets have added a new dimension to soft self-assembly science. 1-9 Soft self-assembly of materials, which is the route for engineering of amphiphilic molecules into different supermolecular assemblies in one, two, or three dimensions, has been of interest for decades. 10-12 Fields such as nano-medicine, biocatalysis, bio-active delivery systems, self-assembled composite and solar cells have taken advantage of the self-organization of amphiphilic molecules in recent years. 13-16 Recently, a series of graphene based macroscopic structures including paper and fibers have been fabricated employing the novel amphiphilic soft self-assembly route. 1, 4, 17 The much expected enhancement in properties, self-assembly and alignment of GO might be achieved if the simultaneous dispersion of functional materials with
GO in the liquid crystalline media is realized. To date, the application of graphene-based liquid crystals as a promising building block in different fields has not yet been realized mainly because of the practical limitations induced by water, the only medium in which the formation of LC GO has been reported. However, there are limitations for aqueous media to disperse many nanomaterials that would be expected to introduce enhanced properties to LC GO. For example, the introduction of highly debundled, isolated and self-oriented carbon nanotubes (CNTs) which has only been observed in CHP and never in water with LC GO would be expected to enhance electrochemical and mechanical properties. Understanding and manipulating the forces involved in amphiphilic self-assembly and expanding the range of solvents in which such phenomena can be exploited, will enable the development of new composites based on LC GO.

Here we report the ability to support GO lyotropic LC phase formation in a wide range of organic solvents through the use of ultra-large GO sheets. This approach enables the exploitation of LC order of GO sheets in organic solvents to organize and align single-walled carbon nanotubes (SWNTs). This work has enabled self-assembly of ultra-stiff, ultra-strong three dimensional (3D) GO-SWNT architectures with high elongation-at-break.

**Results and discussion**

**Formation of LC GO in various organic solvents**

To date, water is recognized as the practical medium for the self-assembly of GO. In the pursuit of rationally designed lyotropic LC GO dispersions in various organic solvents, we dispersed ultra-large GO sheets in a number of common solvents including: water, N-N-dimethylformamide (DMF), N-Cyclohexyl-2-pyrrolidone (CHP), Tetrahydrofuran (THF), acetone, ethanol and a number of other organic solvents, many of which were not known to afford solvophobic self-assembly prior to this report. The organic solvents, successfully used
here, can support dissolution or dispersion of a wide range of materials. This means that self-assembly of a variety of compositions in the solvent media that support LC GO formation would be possible. For example, DMF and THF are the most common solvents for processing polymers. Therefore, dispersing GO in these solvents and achieving LC GO would provide unique opportunities in the production of self-assembled, fully ordered and novel LC GO-based polymer composites. LC GO in THF might also be used as an ordered template for the synthesis and self-assembly of metallic nano-particles such as boron or magnesium, which are water and air sensitive. LC GO in CHP, being the best known solvent for debundling CNTs, might promote fabrication of fully ordered self-assembled CNT-GO composites containing highly debundled CNTs. Ethanol and acetone are general purpose solvents which are commonly used as building blocks in organic chemistry. Acetone is also the solvent of choice for a wide range of epoxy families. Therefore, attaining LC GO in acetone would open an avenue for novel self-assembled epoxy based nano-composites. Other solvents in which we observed lyotropic LC formation of GO include ethylene glycol, methanol, acetonitrile, isopropanol, N-methyl pyrrolidone (NMP) and dimethyl acetamide (DMAc). All of the non-polar solvents were ineffective in dispersing GO and therefore could not afford any LC phase as expected. Representative polarized optical microscopy (POM) micrographs of the representative solvents are given in Figure 1. POM micrographs clearly show the birefringent lyotropic LC behavior of GO in water and all of the above-mentioned organic solvents (see also Figure S1 to S6). The transition concentration from isotropic to the nematic phase was experimentally found to be ~0.25 mg ml⁻¹ for water, DMF, CHP and ethanol, and ~0.50 mg ml⁻¹ for acetone and THF (Table 1). At higher concentrations, the nematic phase formed in all of the solvents spontaneously. Depending on the concentration, as-prepared organic LC GO can be either stable for months or
undergo what is stated as degradation. It should be noted that these concentrations, although the lowest filler content ever reported for the formation of liquid crystals from any colloid, are still higher than the theoretical biphasic region between 0.05 to 0.09 mg ml\(^{-1}\) calculated for rigid platelets (see supporting information for details). This discrepancy can be attributed to the flexible nature of the monolayer GO sheets and their tendency of wrinkling especially in the presence of attached functional groups.

Figure 1. Representative crossed polarized optical microscopy (POM) of LC GO in various organic solvents at a GO concentration of 2.5 mg ml\(^{-1}\).
Table 1. LC formation concentration and GO sheet properties in different solvents.

<table>
<thead>
<tr>
<th>Solvent</th>
<th>LC formation concentration (mg ml⁻¹)</th>
<th>Sheet thickness* (nm)</th>
<th>d-spacing** (nm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Water</td>
<td>0.25</td>
<td>0.83</td>
<td>0.825</td>
</tr>
<tr>
<td>DMF</td>
<td>0.25</td>
<td>1.1</td>
<td>1.05</td>
</tr>
<tr>
<td>CHP</td>
<td>0.25</td>
<td>1.0</td>
<td>0.101</td>
</tr>
<tr>
<td>THF</td>
<td>0.50</td>
<td>0.92</td>
<td>0.937</td>
</tr>
<tr>
<td>Acetone</td>
<td>0.50</td>
<td>0.86</td>
<td>0.846</td>
</tr>
<tr>
<td>Ethanol</td>
<td>0.25</td>
<td>0.83</td>
<td>0.820</td>
</tr>
</tbody>
</table>

*: Sheet thickness obtained from AFM images

**: d-spacing obtained from XRD patterns

Characterization of LC GO

Atomic force microscopy was employed to assess the number of layers and quality of GO sheets in the organic solvent-based LC dispersions, (Figure 2). No aggregation or restacking of GO sheets was observed in any solvent investigated here. As-prepared GO dispersions in all of the organic solvents contained GO sheets that are predominantly in the size of more than tens of micrometres. Step height measurements performed on the samples indicated that all of the samples contained single layer GO sheets; as the mean measured height was between 0.8 to 1.2 nm depending on the solvent used. The thickness of a monolayer of graphene is about 0.34 nm, however, GO has functional groups that act as pillars giving rise to the larger measured height. Some solvent molecules are also expected to bond with the surface of GO and remain even after drying. Hydrogen bonding of each solvent molecule with GO resulted in different apparent sheet thickness, depending on the solvent molecular size and arrangement on the surface of GO sheets.
Moreover, the crumpling of GO sheets that occurs in the case of THF (Figure 2) might also result in lower effective aspect ratio which manifests itself in the observation of different phase boundaries presented in Table 1.

**Figure 2.** AFM images of GO sheets prepared from various organic solvent based LC GO dispersions. Marked line in each image shows the measured thickness of the sheet.

The change in the apparent thickness of GO sheets as a result of bonding with different solvents can also be confirmed by measuring the interlayer $d$-spacing of GO sheets in cast dried films made from various solvent based GO (Table 1). The $d$-spacing represents the interlayer distance between individual GO sheets in the paper material that are oriented perpendicularly to the diffraction plane. XRD measurements were performed to evaluate the effect of the solvents on the interlayer $d$-spacing of GO films (Figure 3a). The process of making GO in water or organic solvents is accompanied by an increase in the $d$-spacing between the graphene layers.
from about 0.34 nm to ≈ 0.8–1.1 nm, which is related to the degree of oxidation and the hydration level (in the case of GO prepared in water) or the bonding of other solvent molecules to graphene sheets.\textsuperscript{1,23-26} Therefore, the peak in the XRD patterns of our as-prepared GO films corresponds to the layer-by-layer distance ($d$-spacing) of each sample prepared in each organic solvent according to the Bragg’s law. The position of the peak and correspondingly the $d$-spacing values observed in the XRD patterns varied with the solvent used. The differences on $d$-spacing values could be attributed to the confinement of organic solvent molecules in the lamellar GO sheet layers, as in the case of water for aqueous LC GO dispersions. This result could be illustrated by the larger $d$-spacings for GO sheets when dispersed in acetone, THF, CHP and DMF, which had intersheet distances of 0.978 nm, 1.01 nm and 1.17 nm, respectively, than that of GO in water (0.846 nm). A very small shift ($d$-spacing 0.820 nm) was observed in ethanol-based LC GO consistent with the similarity of size with water. The measured $d$-spacing values were also in good agreement with the measured GO sheet thickness from AFM results (Table 1). Therefore, we suggest that the individual GO platelets are interlinked via a non-uniform network of hydrogen bonds mediated by oxygenated functional groups and solvent molecules.

Thermal gravimetric analysis (TGA) was also performed on GO papers (Figure 3b). GO papers which were cast from organic solvents with high boiling point temperatures (154 °C and 153 °C for CHP and DMF, respectively) lost considerably more weight at higher temperatures compared to GO recovered from water. In contrast, GO dispersed in volatile solvents (acetone, THF and ethanol) started to lose weight at much lower temperatures. This behavior can be attributed to the confinement of solvent molecules between GO sheets during film formation. The variation between the final weight losses was due to the different amount of solvents associated with each
GO sample. The observed significant weight loss at around 180 °C to 200 °C is attributed to the partial reduction of GO.

Figure 3. a) X-ray diffraction patterns (XRD) and b) Thermal gravimetric analysis (TGA) of GO films as a function of solvents.

Insights into the formation of LC GO

The self-assembly of amphiphiles in water is generally driven by hydrophobic interactions, \(^{23-25}\) which is an important component of a larger solvophobic effect. \(^{23, 26-28}\) Studies into the
thermodynamic driving force for the self-assembly of amphiphiles into LC phases have highlighted that the solvophobic force is almost always dominated by entropic contribution. This behavior is very similar to hydrophobic forces in water where the nature of these interactions is entropic in origin as the enthalpy change is actually unfavorable in most cases. Therefore, the process of self-assembly is an interplay of entropy and enthalpy terms in the free energy as given in Equation (1).

\[ \Delta G^\circ_{\text{self-assembly}} = \Delta H^\circ_{\text{self-assembly}} - T\Delta S^\circ_{\text{self-assembly}}. \]  

Other contributing factors are hydrogen bonding, van der Waals interactions and electrostatic interactions. Graphene oxide is typically considered as a negatively charged amphiphilic molecule and as a result, the \( \Delta G^\circ_{\text{self-assembly}} \) in this case can be represented as the sum of the free energy change associated with hydrogen bonding, \( \Delta G_{\text{hydrogen bond}} \), and electrostatic interactions, \( \Delta G_{\text{electrostatic}} \) (see Equation (2)).

\[ \Delta G^\circ_{\text{self-assembly}} = \Delta G_{\text{hydrogen bond}} + \Delta G_{\text{electrostatic}} \]  

With water as the self-assembly medium, both factors contribute to the free energy change term. Only a few other solvents other than water are capable of supporting amphiphilic self-organization. These solvents include a very limited range of multifunctional alcohols (such as ethylene glycol) and amides and a wide range of protic ionic liquids widely known as PILs. With PILs, where the solvent itself is an ion, the surface charge screening results in negligible electrostatic contribution, which is in contrast with water or organic solvents where the electrostatic charges play an important role in the free energy associated with ordering. In the case of LC GO, the solvent confined between the charged GO sheets adopts a more structural arrangement to balance the steric and repulsive forces. In this sense, GO can be considered as a self-assembling material due to the fact that it involves supermolecular and supramolecular
interactions such as electrostatic repulsion between the adjacent sheets and hydrogen bonding
with the confined solvent molecules, which compensate for the loss of rotational entropy during
the self-assembly process. Therefore, in the case of organic solvents, it is the interplay between
the ability to form multiple hydrogen bonds and electrostatic charges, manifested through
Gordon parameter that governs the process of self-assembly.

The mechanism for the self-assembly process in organic solvents is considered to involve the
solvophobic effect which is linked to the solvent cohesiveness. 26 The Gordon parameter
\( G = \gamma / V_m^{1/3} \) is a direct measure of the solvent cohesiveness where \( \gamma \) is the surface tension and \( V_m \)
is the molar volume. 35 High Gordon parameter represents firstly a higher chance to achieve self-
assembled liquid crystalline phases and secondly a higher thermal stability. 36 However, the ultra
large size of GO sheets used in this study may push the boundaries toward organic solvents with
Gordon parameters which are normally considered to be too low to be able to support
solvophobic self-assembly. So far, the lowest reported Gordon parameter that supports
amphiphilic self-assembly was \( G = 0.576 \) J m\(^{-3}\) for a protic ionic liquid (EAB), 34 which is far
beyond the limit predicted by Evans (\( G \geq 1-1.2 \) J m\(^{-3}\)). 37 However, as the Gordon parameter is
directly dominated by surface tension and surface tension is a direct result of electrostatic
charges and hydrogen bonding, in the case of PILs, the use of Gordon parameter might be
irrelevant as the only dominating force is the ability of the liquid to form extensive hydrogen
bonding network. Recently, Drummond et al. also discovered that some low molecular weight
amides can be utilized as self-assembly media with a Gordon value as low as 0.53 J m\(^{-3}\), which is
the direct result of the similarities between the chemical structure of PILs and the amides. 26
However, apart from these two groups of solvents, no other organic polar solvent with lower
Gordon parameter less than 1.3 J m\(^{-3}\) is yet found to act as an amphiphilic self-assembly medium.
However, employing ultra large GO sheets has enabled us to achieve LC GO dispersions in a wide range of organic solvents with Gordon values previously deemed too low to support self-assembly. The list of solvents we have shown to support LC GO, along with Hansen parameter and Gordon parameter, are given in Table 2.

Table 2. Hansen parameters and Gordon parameter for the solvents could support LC GO.

<table>
<thead>
<tr>
<th>Solvent</th>
<th>LC formation concentration (mg ml$^{-1}$)</th>
<th>Hansen Parameter for solvents</th>
<th>Surface tension (mN m$^{-1}$)</th>
<th>Gordon parameter (J m$^{-3}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>dispersive</td>
<td>polar</td>
<td>hydrogen</td>
</tr>
<tr>
<td>Water</td>
<td>0.25</td>
<td>15.5</td>
<td>16.0</td>
<td>42.3</td>
</tr>
<tr>
<td>ethylene glycol</td>
<td>0.25</td>
<td>17.0</td>
<td>11.0</td>
<td>26.0</td>
</tr>
<tr>
<td>N-methyl pyrroldone</td>
<td>0.25</td>
<td>18.0</td>
<td>12.3</td>
<td>7.20</td>
</tr>
<tr>
<td>DMF</td>
<td>0.25</td>
<td>17.4</td>
<td>13.7</td>
<td>11.3</td>
</tr>
<tr>
<td>dimethyl acetamide</td>
<td>0.25</td>
<td>16.8</td>
<td>11.5</td>
<td>10.2</td>
</tr>
<tr>
<td>CHP</td>
<td>0.25</td>
<td>18.2</td>
<td>6.80</td>
<td>6.50</td>
</tr>
<tr>
<td>methanol</td>
<td>0.25</td>
<td>15.1</td>
<td>12.3</td>
<td>22.3</td>
</tr>
<tr>
<td>THF</td>
<td>0.50</td>
<td>16.8</td>
<td>5.70</td>
<td>8.00</td>
</tr>
<tr>
<td>Acetone</td>
<td>0.50</td>
<td>15.5</td>
<td>10.4</td>
<td>7.00</td>
</tr>
<tr>
<td>Ethanol</td>
<td>0.25</td>
<td>15.8</td>
<td>8.80</td>
<td>19.4</td>
</tr>
<tr>
<td>isopropanol</td>
<td>0.25</td>
<td>15.8</td>
<td>6.10</td>
<td>16.4</td>
</tr>
</tbody>
</table>

Apart from water which has a Gordon parameter higher than 1.2 J m$^{-3}$, DMF (which is an amide) shares a degree of structural similarity with PILs and can form extensive hydrogen bonded networks similar to water. Therefore, although the Gordon parameter is very low, DMF could afford formation of LC GO. For the case of other solvents, it is evident that the
ability to support LC GO is largely governed by the capability of the solvents to form hydrogen bonds. As an example, although the Gordon value of isopropanol and ethanol is very low ($G = 0.541$ J m$^{-3}$ and $G = 0.569$ J m$^{-3}$, respectively), their ability to form multiple hydrogen bonds which is even much higher than DMF, can overcome the low cohesive energy density of the solvent. On the other hand, the lower ability of acetone and THF to form hydrogen bonds resulted in an increased GO concentration required for LC formation (0.50 mg ml$^{-1}$) compared to ethanol (0.25 mg ml$^{-1}$), which has slightly lower Gordon parameter but much higher hydrogen bonding ability. As a result, it is safe to assume that the ability to form an extensive hydrogen bonding network is the most important parameter to govern the self-organization process in the case of LC GO. Consequently, the interplay between the hydrogen bonding and the Gordon parameter can effectively determine: (i) whether an organic solvent can induce the self-assembly process, and (ii) the lowest possible concentration in which spontaneous self-organization can occur.

**Exploitation of the self-assembly nature of LC GO**

The intrinsic self-assembly nature of LC materials can be used to exploit them as versatile templates for the synthesis and alignment of nanoparticles. We utilized LC GO to induce liquid crystallinity to SWNTs dispersion through the addition of LC GO to the SWNTs dispersion. Here we have investigated the effect of introducing SWNTs to LC GO dispersion. POM micrographs of LC GO-SWNTs are presented at Figures 4a and 4b which show the nematic LC behavior of as-prepared mixtures in CHP and DMF, respectively. Although SWNTs and rod shaped particles under some specific conditions can form LC phases, this requires modification of the SWNTs surface by bio-polymers, functionalization (with a subsequent compromise in electronic properties), or the use of superacids. Here we have dispersed
SWNTs in CHP at concentrations as high as surfactant assisted dispersions (1 mg ml\(^{-1}\)). Subsequent mixing of this SWNT dispersion with CHP- or DMF-based lyotropic LC GO resulted in birefringence yet preserves the fundamental properties of SWNTs (no dispersant was added). It has been shown that organic solvent stabilized SWNT dispersions are sensitive to the presence of water whereby the addition of very small amounts of water will cause the dispersion to agglomerate and crash out of solution. The water free nature of our as-prepared LC GO in organic solvents is demonstrated by the fact that SWNTs do not agglomerate and crash out of the dispersion when they are added to the as-prepared LC GO in organic solvents. The quality of the dispersion even after addition of the GO is evidenced by the UV Vis spectra, (Figure 4c). The well-resolved inter-band transitions in the UV/Vis-near-IR spectra of the SWNT dispersion before and after the addition of LC GO indicate that SWNT sizes are preserved in the composite formulation. According to Smalley and Hague, UV/Vis-near-IR spectroscopy is the most reliable method to determine SWNTs size distribution based on Van Hove singularities. As it is evident from UV/Vis-near-IR spectra, the size of our nanotubes is predominantly in the range of 1 nm and 0.95 nm corresponding to the wavelength of ~1300 nm and ~1150 nm in the as-prepared SWNTs dispersions, respectively. The quality of SWNTs dispersion (bundle size) was also preserved during the combination with LC GO.

It is pertinent to mention that although many particles can disturb the liquid crystal director field depending on the particle size, shape and surface interaction with liquid crystal media, GO in this regard enjoys a unique benefit. This unique benefit is due to minimizing the exert forces or torques in the direction perpendicular to the director to promote a configuration in which the distortion is minimal, leading to the in-plane alignment of anisotropic particles (the ordering of particles on GO planes) such as SWNTs; as evident from the strength of our as-prepared
composite films (see next section). Prior to this work, the only successful method of increasing the CNT concentration in lyotropic LCs has been through using extensive amounts of surfactants (which can adversely affect CNTs performance). Only trace amounts of CNTs have been dispersed in thermotropic LCs in order to avoid CNT aggregation. However, in our case, we have been able to disperse and organize substantial amounts of SWNTs (up to 10 wt%) in GO LC without losing birefringence properties and observing any aggregation. Therefore, this LC formulation was utilized for a facile fabrication of self-assembled layer-by-layer LC GO-SWNT 3D assemblies.
Figure 4. Representative POM micrographs of a) LC GO-SWNTs/CHP (90:10 at ~1 mg ml⁻¹), b) LC GO-SWNTs/DMF (90:10 at ~1 mg ml⁻¹). c) UV/Vis-near-IR spectra of SWNTs and LC GO dispersions before and after mixing together.
Self-assembled layer-by-layer multifunctional composite

The LC properties of our as-prepared LC GO-SWNTs dispersions in organic solvents induce a spontaneous self-assembly into engineered long range ordered layer-by-layer 3D structures upon simple casting and drying as shown in Figure 5. The ease of synthesis, much shorter processing time and high scalability of this route in contrast with other layer-by-layer production methods (such as Langmuir–Blodgett deposition) offers the opportunity for facile fabrication of 3D frameworks with exceptional properties. The excluded volume generated by large GO sheets for SWNTs results in an entropic rearrangement to form long range ordering. Therefore, LC GO in CHP can be employed as a dispersing media to process SWNTs based on hydrophobic and π-π interactions as well as a medium in which SWNTs can be self-assembled. This can consequently result in the fabrication of highly aligned and macroscopically periodic self-assembled structures of GO-SWNTs hybrid material. The aggregate-free nature of our as-prepared LC GO-SWNTs enabled us to achieve ultra-stiff, ultra-strong layer-by-layer 3D architecture with high elongation-at-break which enjoy an average modulus of 51.3 GPa, tensile strength of 505 MPa and elongation at break of 9.8% (Figure 6), which is much higher than the parent GO architecture.
Figure 5. a) Representative photograph of a flexible free-standing paper of LC GO-SWNT made by cast drying method. b) SEM image of the cross-section of as-cast dried LC GO-SWNT paper. c) SEM image of the surface of the layer-by-layer composite which is marked as region (i) in (b). Some of the SWNTs are laid on the surface of the paper (white arrow) while others are placed between layers of GO sheet (black arrow). Transparency of the monolayer/few layers of GO sheets allows observing tubes sites in different layers. d-f) Cross-section of composite paper at different magnifications (marked as (ii) in (b)), confirmed the self-oriented nature of the composite as well as maintaining SWNTs debundled after the fabrication of composite.
Figure 6. a) Stress-strain curves of LC GO and LC GO-SWNTs self-assembled composite. b) Diagram of mechanical performance data for layer-by-layer nanocomposites and carbon-based neat papers and fibers in selected significant published reports and in this study (see also table S1).
The average strength, reported here, is much higher than those reported for bucky papers, GO and rGO papers (prepared by either filtration or casting strategies) and fibers (either as is or cross-linked), paper-like materials based on vermiculite, flexible graphite foil, neat SWNTs fibers, and even layer-by-layer assembled polymer, SWNTs, or MWNTs nanocomposites. As proposed by Ruoff et al. very high mechanical strength can be obtained if ordering and alignment of fibrils/macromolecules are achieved. In comparison with irregularly laid-down individual fibrils obtained via filtration, the LC route provided us with a self-mediated platform to organize and order SWNTs resulting in an extraordinarily high mechanical strength. Our average Modulus is also considerably higher than all those (Figure 6 b and Table S1) and just inferior to either SWNT fibers composites with polymers or extensively processed SWNTs. The as-prepared composite paper exhibited an exceptional conductivity of 1500 S m⁻¹.

The ultra-large nature of our as-prepared GO sheets (with lateral size in excess of 32 µm compared to typically few hundreds of nm in the case of other reports) provides us with a highly wrinkled topography, which can first contribute to an overall increase in strain, and second maximize the fraction of the surface area available for mechanical reinforcement and toughening. Therefore, in the case of ultra large GO sheets, the initial straightening due to wrinkled topography plays a crucial role in the observed increase in elongation-at-break of our as prepared GO compared to other reports. Furthermore, this enhanced wrinkled topography provides us with a unique platform to accommodate individual, separated SWNTs in between the GO sheets resulting in an overall enhancement in exploitation of the extraordinary mechanical properties of SWNTs. Also, the introduction of SWNTs can prevent the restacking of individual two-dimensional GO sheets, further enhancing the properties of the hybrid material.
Furthermore, our GO-SWNTs hybrid material exhibited a very high toughness in the order of 20 J g⁻¹, which is almost two orders of magnitude higher than GO paper, ⁵² three orders of magnitude higher than pristine bucky paper, ⁵¹, ⁶⁶ and flexible graphite foils and even 7 times higher than GO/rGO fibers. ¹⁷ The flat fracture surface of our ruptured paper is evidence of the good material homogeneity and layer-by-layer nature of both our as-prepared neat and hybrid papers. The strong interfacial bonding in the case of our hybrid paper is also evident from the straight fracture surface in contrast to the rupture of bucky paper (Figure 5). In regards to mechanical properties, SWNTs act as bridging components between individual GO sheets (see high magnification SEM micrographs in Figure 5d-f). As the GO sheets are not strongly attached together, reinforcing them with SWNTs can result in an overall increase in mechanical properties. SWNTs can bridge individual GO sheets and therefore increase the shear force between GO sheets. Furthermore, the most important limitation of using SWNTs as reinforcing agents is the intertube and interfacial slippage within bundles. The poor load transfer within bundles results in interfacial slippage as the effective moduli and strength for bundles are far below those expected for individual SWNTs. The individual nature of SWNTs reported in our study, as evident in high resolution SEM figures presented in Figure 5 and the well-resolved inter-band transitions in the UV/Vis-near-IR spectra of the SWNT dispersion before and after the addition of LC GO presented in Figure 4, ensures that the shear slippage of nanotubes within the bundle does not occur and the mechanical properties of SWNTs used in this study are preserved resulting in an overall enhancement of the final composite properties. The findings presented here will pave the way to versatile and highly scalable routes for the fabrication of a wide range of large-scale 3D graphene based architectures (including metallic or polymer based composites).
with extensive applications in multifunctional wearables, sensors, supercapacitor devices and electronic gadgets.

**Conclusion**

In summary, our discovery of lyotropic LC GO dispersions in a range of organic solvents contributes to the fundamental understanding of the solvophobic effect and the parameters affecting the self-assembly process. The ability of the solvents to promote self-assembly in GO is governed by their polarity and is linked to the ability to form extensive hydrogen bonding. The steric hindrance between the highly charged GO sheets is the factor that overcomes the unfavorable loss of rotational entropy associated with ordering. The expansion of the list of known solvents that can promote the self-assembly process and lyotropic liquid crystallinity has enabled us to tailor-make processable self-assembled, self-oriented SWNTs/GO hybrid composites with superior mechanical performances (Table S1). This discovery could provide practical solutions to the processability of a wide range of materials that require organic solvents because of solubility issues and/or water sensitivity (*i.e.* metal oxides, polymers and nanomaterials).

**Methods**

*LC GO synthesis:* In order to obtain fully oxidized graphite and preserve the high initial lateral sizes of graphite flakes, dry expandable graphite flakes (3772, Asbury Graphite Mills USA) were first thermally treated at 1050 °C for 15 sec. The resultant expanded graphite (EG) was used as the precursor for GO synthesis following previously described methods. 1, 65, 67 Briefly, 5 g of EG and 1 lit of sulfuric acid were mixed and stirred in a flask for 24h. 50 g of KMnO₄ was added to
the mixture dropwise. The mixture was transferred into an ice bath, and 1 lit of Milli-Q water and 250 mL of \( \text{H}_2\text{O}_2 \) were poured slowly into the mixture, realizing a colour change of the suspension to light brown. Having stirred for another 30 min, the GO particles were then washed and centrifuged with a HCl solution (9:1 water:HCl by volume), then centrifuged again and washed with Milli-Q water until the pH of the solution became about 5 to 6. The resultant ultra-large GO sheets were dispersed in deionized water by gentle shaking (i.e. without the aid of sonication process).

N-N-dimethylformamide (DMF), N-Cyclohexyl-2-pyrrolidone (CHP), tetrahydrofuran (THF), acetone, ethylene glycol, N-methyl pyrrolidone (NMP), dimethyl acetamide (DMAc), methanol, isopropanol and absolute ethanol, all from Sigma, have been chosen for investigation. LC GO dispersions in various organic solvents were prepared by extraction of water from the parent aqueous LC GO dispersion via repeated centrifugation-washing steps (6 times of 10 to 30 minutes at 11000 rpm, ProSciTech TG16WS) using the selected solvent. Briefly, 15 ml of the parent aqueous LC GO (2.5 mg ml\(^{-1}\)) was poured into a 50 ml centrifuge tube (Nalgene centrifuge tube) to which 20 ml of the selected solvent was added and then mixed vigorously by vortex shaking. After centrifugation, 30 ml of the supernatant was extracted, replaced with 30 ml of the selected solvent and then mixed vigorously by vortex shaking. This process was repeated 5 times to replace the water with the selected organic solvent.

*Layer-by-layer self-assembly of the LC GO-SWNT composite:* SWNTs dispersion was prepared by adding 15 mg of HiPCO-SWNTs to 15 ml of CHP. This dispersion was subjected to a 3 hr of high-power tip sonication (SONICS Vibra Cell 500 W, 30 % amplitude) followed by a one day low-power bath sonication (Branson B5500R-DTH). The SWNTs/CHP (1 mg ml\(^{-1}\)) dispersion was then mixed with LC GO in DMF or in CHP (2.5 mg ml\(^{-1}\)) at the weight ratio of
(10:90) followed by 10 min of vortex mixing. A self-assembled layer-by-layer composite was fabricated by casting 2 ml of the composite formulation into a Teflon mold (2 cm X 2 cm) and then oven dried at 110 °C for 2 days. The resultant oven-dried GO-SWNT paper was washed several times by DMF, acetone and dried in air to remove the remaining solvent. GO paper, as control, was made using GO in DMF; however, SWNT in CHP was not form free standing paper using simple casting.

**Characterizations:** AFM analysis was carried out by first depositing GO sheets from their dispersions on pre-cleaned and silanized silicon wafer (300 nm SiO$_2$ layer). Silane solution was prepared by mixing 3-Aminopropyltriethoxysilane (Sigma) with water (1:9 vol) and one drop of hydrochloric acid (Sigma). Pre-cut silicon substrates were silanized by immersing them in aqueous silane solution for 30 min and then washed thoroughly with Millipore water. LC GO in each solvent was first diluted to ~50 µg ml$^{-1}$, then GO sheets were deposited onto silanized silicon substrates by immersing a silicon substrate into the GO dispersion for 5 seconds, then immersed in the solvent bath (the same solvent used to form the LC GO) for 30 seconds and then dried under nitrogen flow at room temperature. The CHP based sample was then heated up to 70°C to be dried due to its higher boiling point compared to the other solvents. Prior to AFM analysis, GO sheets were observed under an optical microscope to ensure uniform GO sheet deposition was achieved. Atomic force microscopy (MFP-3D AFM Asylum Research, CA) was carried out in tapping mode under ambient conditions. X-ray diffraction (XRD) studies were performed using a powder XRD system (Philips1825) with CuKα radiation (λ=0.154 nm) operating at 40 keV and with a cathode current of 20 mA. Thermal gravimetric analyses (TGA) were carried out in nitrogen atmosphere from room temperature to 250 °C at a temperature ramp rate of 1 °C min$^{-1}$. The birefringence of LC GO dispersions was examined by polarized optical
microscopy (POM, Leica CTR 6000) operated in transmission mode by looking at a drop of LC GO on a glass slide. UV/Vis-near-IR spectra were recorded using a Shimadzu UV-3600 spectrometer from 600 nm to 1400 nm. The conductivity of LC GO-SWNTs paper was measured using a JANDEL 4-point probe resistivity system (Model RM2) with a linear four-point probe head. The mechanical properties of the composite paper were measured using a Shimadzu tensile tester (EZ-S) at a strain rate of 0.5 % min⁻¹ parallel to the GO plain in the paper. Youngs modulus (Y), tensile strength (σ), breaking strain (ε), and breaking energy (toughness) were calculated, and the average reported for 10 samples. The thickness of the composite papers was ~10 µm and the papers are cut in 20 mm by 5 mm. The obtained strips were mounted on aperture cards with commercial superglue and allowed to air dry. Note, only 10 mm of each sample was exposed to the applied strain as 5 mm from each side is used for gluing to the aperture card.

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ABBREVIATIONS

liquid crystalline, LC; graphene oxide, GO; single walled carbon nanotubes, SWNTs; tetrahydrofuran, THF; N-dimethylformamide, DMF; N-cyclohexyl-2-pyrrolidone, CHP; N-methyl pyrrolidone, NMP; dimethyl acetamide; DMAc; polarized optical microscopy, POM; Atomic force microscopy, AFM; X-ray diffraction, XRD; Thermal gravimetric analysis, TGA.
ASSOCIATED CONTENT

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Supporting Information.

Calculation of the theoretical concentration for LC formation. Figure S1 to S6, representative polarized optical micrographs of LC GO dispersions in different solvents at various GO concentrations. Figure S7 XRD pattern of LC GO/SWNTs composite. Table S1, comparison of LC GO/ SWNTs mechanical performance data with layer-by-layer nanocomposites and carbon-
based neat papers and fibers in selected significant previous reports and in this study. This material is available free of charge via the Internet at http://pubs.acs.org.

REFERENCES


