Chirality and liquid crystals are both widely expressed in nature and biology. Helical assembly of mesophasic molecules and colloids may produce intriguing chiral liquid crystals. To date, chiral liquid crystals of 2D colloids have not been explored. As a typical 2D colloid, graphene is now receiving unprecedented attention. However, making macroscopic graphene fibres is hindered by the poor dispersibility of graphene and by the lack of an assembly method. Here we report that soluble, chemically oxidized graphene or graphene oxide sheets can form chiral liquid crystals in a twist-grain-boundary phase-like model with simultaneous lamellar ordering and long-range helical frustrations. Aqueous graphene oxide liquid crystals were continuously spun into metres of macroscopic graphene oxide fibres; subsequent chemical reduction gave the first macroscopic neat graphene fibres with high conductivity and good mechanical performance. The flexible, strong graphene fibres were knitted into designed patterns and into directionally conductive textiles.

Chirality and liquid crystals are both widely expressed in nature and biology. Helical assembly of mesophasic molecules and colloids may produce intriguing chiral liquid crystals. To date, chiral liquid crystals of 2D colloids have not been explored. As a typical 2D colloid, graphene is now receiving unprecedented attention. However, making macroscopic graphene fibres is hindered by the poor dispersibility of graphene and by the lack of an assembly method. Here we report that soluble, chemically oxidized graphene or graphene oxide sheets can form chiral liquid crystals in a twist-grain-boundary phase-like model with simultaneous lamellar ordering and long-range helical frustrations. Aqueous graphene oxide liquid crystals were continuously spun into metres of macroscopic graphene oxide fibres; subsequent chemical reduction gave the first macroscopic neat graphene fibres with high conductivity and good mechanical performance. The flexible, strong graphene fibres were knitted into designed patterns and into directionally conductive textiles.
Liquid crystalline colloids are of great interest due to their significance in understanding colloid physics and accessing fluid-phase assembly. The chiral liquid crystals (CLCs) of one-dimensional (1D) colloids such as DNA₁, bacteriophage Φ₂ virus² and imogolite³ have been reported. Various conventional liquid crystalline mesophases of two-dimensional (2D) colloids—nematic, columnar and lamellar—have been reported⁴–¹⁰. However, the CLC phase of 2D colloids has not been reported previously, possibly because no one knows whether achiral flake particles can form long-range helical systems.

As a typical 2D colloid, graphene, a single atomic layer of graphite, consists of sp²-hybridized carbon atoms in a honeycomb lattice⁴¹. It has possibly become the fastest growing topic in science and technology because of its marvellous properties such as the strongest mechanical strength ever measured, giant intrinsic carrier mobility, tunable band gap or electrical conductivity and record thermal conductivity¹²–¹⁵. Advances have been achieved in the area of graphene-based devices and composites¹⁵–¹⁹, but to effectively translate the properties of graphene into macroscopic-ordered materials, such as continuous fibres, challenges must be overcome in terms of its limited dispersibility in common solvents as the lack of scalable assembly methods⁲⁰. Alternatively, chemically oxidized graphene or graphene oxide (GO), the most accessible precursor of graphene, possesses good solubility in water and polar organic solvents, and can form lyotropic nematic liquid crystals (LC)²¹,²², a typical mesophase of highly anisotropic colloids⁴–¹⁵. This conventional phase behaviour of GO sheets could lead to the discovery of novel mesophases. Furthermore, GO LC with regular ordering promises a brand-new methodology to fabricate high-performance carbonaceous fibres with versatile functionalities from natural graphite.

**Results**

**Textures of GO CLCs.** The GO sheets we used are single-layered, narrowly polydispersed and individually dispersed in water (Fig. 1a,b; Supplementary Fig. S1). The 2D morphology and width distribution of GO sheets are demonstrated in Figure 1a,b, respectively. Their average thickness (t) is 0.8 nm and the average lateral width (w) is 0.81 μm with a narrow polydispersity of 13% (much lower than 83% of the crude GO sample without size fractionation²³). The GO aqueous dispersions were observed by polarized optical microscopy (POM) between crossed polarizers (Fig. 1c–h). Upon increasing the volume fraction (φ) to 0.23%, GO dispersions evolve from blank to Schlieren texture (Fig. 1c), a typical birefringent texture of nematic LCs²⁴–²⁶. According to the model for polydispersed infinitely thin platelets, this concentration of isotropic–nematic phase transition (0.23%) is very close to the theoretical value (0.25%)²⁵. In the stable nematic phase, the black brushes represent the disclinations of orientational orders, corresponding to the neighbouring crossed direct vectors (Fig. 1d).

As φ increases up to 0.39, 0.60 and 0.76%, the textures turn into aligned, vivid bands, which are similar to the fingerprint texture of cholesteric phase²⁷ and Grandjean texture of TGB phase²⁸ (Fig. 1e–g), additional fingerprint textures see Supplementary Fig. S2). As the incompatibility of helical director field and smectic-layered feature in a continuous system results in frustrated structures containing a regular screw lattice of grain boundaries in TGB phase²⁸, we can speculate that the textures of our samples result from the helical arrangement of GO sheets with vectors map between neighbouring bands as indicated in Figure 1h. The width of these aligned bands could be related to the pitch length, and becomes thinner with φ increasing from 0.39% (Fig. 1e) to 0.60% (Fig. 1f) and 0.76% (Fig. 1g). Such aligned banded structures have been confirmed by

---

**Figure 1** Atomic force microscopy image of graphene oxide sheets and POM observations of GO aqueous dispersions. (a) Typical atomic force microscopy image of GO sheets. (b) The width distribution of GO sheets obtained from Supplementary Figure S1. (c–g) POM images of GO LCs with successively increased volume fractions (φs) of 0.23, 0.38, 0.60, 0.76%. (d,h) Director fields of the disclination in the nematic phase (rectangle in c) and the fingerprint texture of chiral phase (e–g). The director vector (n) is defined as the normal direction of GO sheet plane. The crosses indicate ns into the paper and the arrows show ns on the paper in d and h. Scale bars, 2 μm (a) and 200 μm (c–g).
the inverse volume fraction (1/φ) is concluded in Figure 2d insert (detailed data see Supplementary Table S1). At 0.5% < φ < 2.3%, the evolution displays a linear relation, where d is proportional to 1/φ (d = t/φ, t is the thickness of 2D colloids), following the typical 1D swelling behaviour of lamellar 2D colloids LCs. The fitting line slope reflecting the experimental thickness of GO sheets is 0.46±0.03 nm, near to the theoretical thickness of a monolayer graphene sheet (0.34 nm), which proves that GO sheets are perfectly exfoliated to individual ones in their LCs. The linear region ends as φ approaches 0.5%, around the transition concentration (φ = 0.38%) from nematic to lamellar phase observed in POM investigations (Fig. 1c,e). For dilute dispersions with lower φ, d value is deviated from the first linear region, possibly suggesting the isotropic volume swelling behaviour in the nematic phase.

CD activity of GO CLCs. For a CLC, circular dichroism (CD) is a useful tool to characterize the selective reflection coefficients for right- and left-handed circularly polarized waves. To examine the chirality of our GO CLCs, we obtained their CD spectra, as shown in Figure 3a. The CD activity is very weak for the dilute GO dispersions, and becomes stronger upon increasing φ, for instance to 0.38% (a value close to nematic–CLC phase transition). Very strong CD signal (up to 2,400 millidegrees) is detected as φ rises to 0.76% (a value of GO CLC with stable lamellar structure). The
characteristic CD spectrum spans around 420 nm from 380 to 800 nm that covers a wide range from near ultraviolet and visible light to near infrared, implying a gradient helical arrangement of GO sheets. GO CLC with such a wide CD response can be used as potential optical components. By contrast, no obvious CD activity was detected for the filtered transparent GO solid films, in which GO sheets are aligned parallel along the film surface. These results validate that the CD activity of GO CLC is generated by the helical features of GO sheets (zeta potential of GO dispersions is $-61 \pm 5 $mV), the neighbouring blocks repel each other, forming the rotated conformation by turning to a certain angle (calculated as $7.0^\circ$, $\phi = 360 w/p$ at $\phi = 0.98\%$) to minimize the free energy (bottom right, Fig. 5). Along the helical axis, the blocks rotate successively in the same direction (clockwise or anticlockwise) to establish the helical arrangements (centre, Fig. 5). In other words, the negatively charged grain boundaries of GO sheets produce helical dislocations in their lamellar fluids and thus set the orderly frustrated structures in their chiral phases. In fact, we can recognize two twisted neighbouring lamellar blocks under the high magnification of cryo-SEM (Fig. 4c). Our twist-lamellar-block model for CLCs of 2D colloids can be analogous to the reported TGB phase of conventional small molecules.

**Helical structures in GO CLCs.** The helical structural details of GO CLC have been directly visualized by cryo-scanning electron microscopy (SEM), as shown in Figure 4a–f. The freeze-fracture morphology of the GO CLC confined to a circular cavity shows annual ring-like structures comprising of undulating bands with distinct boundaries (Fig. 4a). At low salt concentration ([NaCl] < $1 \times 10^{-4}$ M), the ellipticity of GO CLC is strong, and decays with increasing [NaCl] from $1 \times 10^{-4}$ to $5 \times 10^{-3}$ M. Eventually, the ellipticity falls to zero at [NaCl] > $1 \times 10^{-2}$ M. These measurements clarify that the helical arrangement in GO CLCs is dominated by the electrostatic repulsion between GO sheets.

**TGB-like model for GO CLC.** On the basis of above experimental results, we constructed a model in one-pitch range for the CLC of 2D colloidal GO sheets that holds both continuous lamellar and helical features (Fig. 5). GO sheets assemble into lamellar blocks with a local vector (n) in their normal direction, and the $d$ between adjacent graphene sheets deceases with the increasing of $\phi$ (top left, Fig. 5). Because of negative charges at boundaries and on surfaces of GO sheets (zeta potential of GO dispersions is $-61 \pm 5 $mV), the neighbouring blocks repel each other, forming the rotated conformation by turning to a certain angle (calculated as $7.0^\circ$, $\phi = 360 w/p$ at $\phi = 0.98\%$) to minimize the free energy (bottom right, Fig. 5). Along the helical axis, the blocks rotate successively in the same direction (clockwise or anticlockwise) to establish the helical arrangements (centre, Fig. 5). In other words, the negatively charged grain boundaries of GO sheets produce helical dislocations in their lamellar fluids and thus set the orderly frustrated structures in their chiral phases. In fact, we can recognize two twisted neighbouring lamellar blocks under the high magnification of cryo-SEM (Fig. 4c). Our twist-lamellar-block model for CLCs of 2D colloids can be analogous to the reported TGB phase of conventional small molecules.

**Chiral silica gels by GO CLC templating.** The fluid GO CLC can be fixed by hydrolysis of tetraethyl orthosilicate via a sol–gel method, affording a transparent, self-standing gel film (Fig. 6a,b), with the parent structure of GO CLC. The central and fringe domains of the film show Schlieren (Fig. 6c) and vivid banded textures under POM (Fig. 6d), respectively. The difference between textures is likely ascribed to the tensile force of the fringe during cross-linking. Importantly, the film shows considerable CD activity, demonstrating that the gel holds the helical structure of original CLC (Supplementary Fig. S8). The resultant gel films with specific chirality can be practically applied to optical apparatus, and GO CLC could serve as templates to fabricate chiral porous materials.

**Continuous graphene fibres by wet-spinning.** To construct macroscopic graphene fibres is a big challenge that remains unsolved. Clearly, wet-spinning is the only viable approach because of the high-temperature stability of graphene (melt-spinning is not an option), as is the case for CNTs. Our discovery of GO LCs with lamellar structures allows to disperse GO at high-enough concentrations, suitable for efficient alignment and effective coagulation. For the first time, we have fabricated neat GO fibres by spinning the aqueous LCs in a coagulation bath of 5 wt% NaOH/methanol solution (spinning apparatuses are sketched in Supplementary Fig. S9). Direct spinning of GO LCs at concentrations of $\phi = 0.76$ and 2.0% resulted in brittle fibres and collapsed belts, respectively, (Supplementary Fig. S10a). From the concentrated LC dispersion (for example, $\phi = 5.7\%$), we obtained continuous fibres successfully. Metres of GO fibres have been readily produced in ~10 min, with

---

**Figure 3 | CD activity of GO CLCs.** (a) CD spectra of GO aqueous dispersions and GO solid film. (b) Normalized CD activity of GO CLC ($\phi = 1.0\%$) as a function of salt concentration.
Figure 4 | Cryo-SEM images and POM textures for GO CLCs. (a–e) Top-view SEM images show the fracture morphology of GO CLCs at volume fraction $\phi \approx 0.98\%$ confined in a circular cavity. The red crosses denote vectors ($\mathbf{n}_s$) into the paper and the orange arrows on the paper. (b) The central domain in (a). (c) The screw dislocation of neighbouring GO lamellar blocks with twist vectors. (f) Side-view SEM image of assembled GO blocks. (g–i) POM images between crossed polarizers of GO CLCs in central (g) and lateral (h, i) domains. The regular ring textures (g–i) are in accordance with the annual-ring undulating fracture morphology (a–e) of GO CLC. Scale bars, 200 $\mu$m (a), 25 $\mu$m (b–f) and 300 $\mu$m (g–i).

Figure 5 | Proposed model for one pitch of GO CLCs. The vectors ($\mathbf{n}_s$) of the lamellar blocks rotate anticlockwise along the helical axis. Top left: the lamellar structure of the organized GO block with an identical $n$ and a regular spacing $d$; the semitransparent blue blocks indicate interlayer water, and GO sheets are indicated as pink nets (double bonds in the basal planes and pedant functional groups have been omitted for clarity). Bottom right: the screw dislocation between two neighbouring grain boundaries of GO lamellar blocks (the red line denotes GO sheet boundaries). The negative charges (blue minus) spread the surface and boundaries of GO sheets and make two blocks repulse each other, with an angle $\theta$ between two vectors ($\mathbf{n}_1$ and $\mathbf{n}_2$).

Figure 6 | Chiral gel film templated from GO CLCs. (a, b) Respective photographs of the chiral gels under natural light and between crossed polarizers. The diameter of the film is 3 cm. (c, d) The respective POM images in the central zoom and fringe of the chiral film. Scale bars, 200 $\mu$m.
tunable diameters (50–100 μm), by adjusting the size of nozzle and the drawing speed (Fig. 7a,b). Typical knots are made and the fibres do not break as the knots are tightened (Fig. 7c; Supplementary Fig. S10b). The curved structure through 360° demonstrates the excellent flexibility and high resistance to torsion of GO fibres compared with the classical carbon fibres. Through SEM kept in high magnitude (Supplementary Fig. S10c), we identified that GO sheets aligned in their planar directions with crumbled surfaces.

Subsequently, we prepared neat graphene fibres by chemical reduction of GO fibres in hydroiodic acid (40%)\(^3\). The reduced graphene fibre shrinks in diameter and exhibits grey metallic lustre. The X-ray diffraction studies (Fig. 8a) indicate that the interlayer spacing decreases from 8.9 Å (20 = 9.86°) for GO fibres to 3.7 Å (20 = 24.25°) for graphene fibres, a value close to the interlayer spacing (3.35 Å) of graphite, due to the elimination of pendant functional groups of GO sheets during the chemical reduction. The reduction was further confirmed with Raman spectra of fibres (Supplementary Fig. S11). The reduced graphene fibre shows high electrical conductivity up to ~2.5×10^4 S m\(^{-1}\), even higher than the filtrated graphene paper reduced by hydrazine hydrate (~7.2×10^3 S m\(^{-1}\))\(^3\).

The continuous GO fibres exhibit a typical plastic deformation at room temperature (Fig. 8b). This behaviour possibly arises from the stretching of the crumbled GO sheets and by the displacements of GO sheets (see the fracture morphology in Fig 7d; Supplementary Fig. 11d–f). In the elastic region, the fibres have a typical Young's modulus of 5.4 GPa at small deformation. Importantly, the fibres show fracture elongations of ~6.8–10.1%, an order of magnitude greater than that of filtrated GO papers (~0.4%)\(^1\). The fracture strength of our GO fibres is measured to be 102 MPa, comparable to that (70–130 MPa) of filtrated GO papers\(^3\).

The chemically reduced graphene fibres we obtained were strong and flexible. Compared with GO fibres, graphene fibres show enhanced Young's modulus (7.7 GPa) and fracture strength (140 MPa), while keeping considerable fracture elongation (~5.8%; Fig. 8b). This enhanced strength could be ascribed to the stronger interactions between graphene sheets resulting from the more compact stacking (small interlayer distance from Fig. 8a) for the graphene fibre. Although our fibres are far weaker than the individual graphene sheets (Young's modulus 1.0 TPa and fracture strength 130 GPa\(^1\)), we believe that the mechanical property can be greatly improved by optimization of the spinning process and post annealing to decrease the voids in fibres and then enhance interactions between the graphene sheets. Similar to GO fibres, the crumbled graphene sheets can be stretched under tensile stress, leading to the high fracture elongation for the graphene fibre (see the surface and fracture morphology in Fig. 7c; Supplementary Fig. S12). No breakage occurs when the fibre is curved to a tightened knot (Fig. 7f), showing the flexibility of neat graphene fibres. In addition, we used the flexible graphene fibres to embroider patterns in other textiles (for example, a Chinese character, ZHONG, in Fig. 7g), and to wave pure graphene textiles or mixed ones with commercial

---

**Figure 7** | Macroscopic neat GO fibres and chemically reduced graphene fibres. (a) Four-metre-long GO fibre wound on a Teflon drum (diameter, 2 cm). SEM image of the fibre (b), and its typical tighten knots (c). (d) The fracture morphology of GO fibre after tensile tests. The surface wrinkled morphology (e) and the tighten knot (f) of graphene fibre. (g) A Chinese character ("Zhong") pattern knitted in the cotton network (white) using two graphene fibres (black). (h) A mat of graphene fibres (horizontal) woven together with cotton threads (vertical). Scale bars, 50 μm (b–f) and 2 mm (g, h).

**Figure 8** | X-ray diffraction patterns and typical mechanical measurements under tensile loading of GO and graphene fibres. X-ray diffraction patterns (a) and typical mechanical measurements under tensile loading (b) of GO and graphene fibres. The number 1 and 2 denote graphene and GO fibres, respectively. The strain rate is 5% min\(^{-1}\). The Young's moduli of GO and graphene fibres at low deformation are about 5.4 and 7.7 GPa, respectively.
threads (Fig. 7h). These graphene-based fibres and textiles should have promising applications in electromagnetic shields, antennas and batteries.

Discussion

Recently, some exotic phase behaviours have been observed in the liquid crystalline system of asymmetrical 2D colloids, such as in empty liquids31 and in biaxial LCs32, indicating the fluid physics of 2D colloids is still growing with various unknown mesophases despite the past 70 years3. Herein, we found that 2D colloids can form a CLC phase by using the prototype of GO sheets. With the help of synchrotron SAXS measurements, we have disclosed two different arrangements in GO dispersions, depending on the concentration: orientational order and lamellar attributes, which are assigned to the nematic and chiral mesophases, respectively.

Theoretically, Landau–Peierls instability in the system with the 1D density wave in 2D liquid medium exhibits a quasi-long-range order in smectic phases, rather than in the true-long-range order, caused by the thermal fluctuation37. Experimental results have demonstrated that the positional correlation function decay algebraically as some power of the distance in smectic-A mesophase38–40. To further explain the orderings in GO CLCs, Calliée lineshape analyses (see Supplementary Note) were carried out to the first scattering peak \( q_0 \) in SAXS profiles. We observed that the scattering intensity also obeys the power-law decay as a function of scattering vectors \( \langle S(0,0,q) - (q - q_0)^2 \rangle \), \( \eta \) the Calliée exponent, implying the lamellar structures of GO CLCs. In a single GO sheet, all the atoms involved are covalently bonded, and thus the sheet has an extremely high Young’s modulus (experimental value for the chemically converted graphene 0.25 TPa39). We estimate that the Young’s modulus multiplies by the average area of GO sheets \((w = 0.81 \pm 0.13 \mu m)\) to make the value of elastic modulus \( (K = 1.6 \pm 0.5 \times 10^4 N/m) \), about five orders of magnitude higher than that of self-assembled bilayers without any intermolecular covalent bonding in the conventional smectic phase39. By fitting the scattering profiles, we obtained the Calliée exponents \( \eta \) of GO CLCs with gradient 0.6 from 2.12 to 0.38. Derived from the equations (1) and \( (2) \),

\[
\eta = \frac{q^2 k_B T}{8 \pi (BK)^{1/2}} \\
\xi = (Kd^2)/(B)^{1/4}
\]

where \( k_B \) is the Boltzmann constant \( (1.38 \times 10^{-23} \text{ J/K}) \), \( B \) the compression modulus, \( T \) the thermodynamic temperature \( (298 \text{ degree Kelvin for our samples}) \), \( \xi \) a correlation length for the fluctuation \( (\text{for the distance longer than } \xi \text{, the layer height fluctuations are coherent from layer to layer, whereas for distance smaller than } \xi \text{, fluctuations are single layer and incoherent}) \), we further obtained the corresponding correlation lengths \((\xi)\) of GO CLCs that range from 161.4 ± 25.7 to 589.5 ± 93.0 \( \mu m \) (detail fitting and quantitative analysis are given in Supplementary Note and Table S2). These calculated correlation lengths are giant, around four orders of magnitude longer than that in the conventional smectic phase of small molecules41, implying that there are little thermal fluctuation in GO CLCs. This effect is mainly attributed to the rigid, solid-like structure of individual GO sheet. The lamellar structures have also been observed previously in the suspensions of solid-like, single colloidal platelets with very little thermal fluctuation42. It is worth pointing out that more effort is needed to put on the specific theory model and computer simulation on LCs of 2D colloids, as the solid flakes have essentially different natures from the conventional bilayers noncovalently assembled from small molecules or lipids.

Importantly, we have detected that GO LCs begin to exhibit CD activity from \( \phi = 0.38\% \), and the optical activity is significantly enhanced with increasing \( \phi \). In the combination of CD inactivity in the filtrated GO films with uniform lamellar structure (Fig. 3a), we speculated that the observed optical activity originates from the twist dislocations of the lamellar structure in GO CLCs. To assess the major role of electrostatic repulsion in the formation of CLC, we have investigated both structures (by SAXS) and CD activities of GO dispersions with gradient salt concentrations. With enhancing the screening effect caused by the stronger ionic strength of higher salt concentration, the weakened repulsive interaction between the basal planes of GO sheets brought the stepwise compression of the lamellar structure (with decreasing interlayer spacing), and the weakened repulsive interaction between GO sheets boundaries induced the faded optical activity (Figs 2e and 3b). Therefore, the electrostatic interaction in GO CLCs is the dominate force and contributes to this new phase behaviour.

Owing to the similar POM textures between our GO CLC and TGB phase, we offered a twist-lamellar-block model to explain the natures of the CLC of 2D colloids, holding lamellar and helical structural attributes simultaneously (Fig. 5). Moreover, the fingerprint textures (Fig. 4g–i) and annual ring-like fracture structures (Fig. 4a–f) are in accordance with this model. Following with the first exposal of CLC for 2D colloids, more CLCs of 2D colloids (for example, clays) will be probably unveiled in the future, to enrich our understandings on colloidal LC regime, which will in turn boost the creation of new materials. In addition, this discovery also calls for the construction of theoretical model for CLC of 2D colloids.

It is known that one of the most important applications of lyotropic LCs is to make macroscopic fibres by wet-spinning technology. Up to date, linear, rigid polymers such as aromatic polyamides as well as 1D nanowires, such as CNTs, have been successfully spun into strong and continuous fibres from their LC dopes42–45, due to the noncovalent chemical bonds (for example, van der Waals forces and hydrogen bonding) and strong mechanical chain entanglements or interconnection among different tubes. Even though LCs of 2D colloids have been reported for several decades, macroscopic fibres made from neat 2D plates have never been achieved previously, partly because the interactions between solid-like flakes are weak, and partly because the solubility of colloids is not high enough for continuous spinning. We have demonstrated that highly soluble graphene sheets, a typical model of 2D colloids, can be readily spun into continuous and robust fibres from their LC dispersions. The sheet alignment inherited from the intrinsic lamellar order of LCs provides strong interactions between contacted sheets that are responsible for the strong mechanical strength of graphene fibres, and the locally crumbled structures of individual sheets make the graphene fibres flexible. The integration of high conductivity, excellent mechanical property and unique attributes associated with graphene makes the fibres highly attractive in many applications such as functional textiles and chemical sensors. The industrially viable wet-spinning technique provides a novel approach to fabrication of carbon fibres from natural graphite. In our near term, particularly promising applications of GO LCs as useful hosts are macroscopic graphene-based fibres with diversiform guests such as polymers, nanocrystals and CNTs; such new materials would be widely useful in the fields of high-performance fibres, energy storage and fuel cells. Inspired by the concept of continuous fibres made from graphene sheets, we believe that the possible LCs of other 2D colloids (such as montmorillonite4, metallic sheets43, MoS2, BN and other 2D nanosheets44) can be wet-spun into macroscopic fibres with unique properties for many promising applications.

Methods

Preparation of narrowly dispersed GO samples. GO sheets were prepared by the protocol described previously41. The isopinic differential centrifugation was performed on the GO dispersion \( (\Phi = 0.075\%) \). After the centrifugation at 5,000 r.p.m. for 2h, the bottom gel is the classified GO sample, and the supernatant was processed for the successive centrifugation. Through repeating process at respective speeds of 10,000 and 15,000 r.p.m. (25,300 g) for 2h, we obtained two corresponding classified samples and took the sample (at 15,000 r.p.m.) with the smallest
width for investigations in this study. GO dispersions (5 ml) were freeze-dried for 2 days and then further dried at room temperature under vacuum for 5 days and their mass fractions ($f_{g}$) were obtained. The volume fraction ($\phi$) is calculated by:

$$\phi = \frac{f_{g}}{\rho_{GO}}$$

where $\rho_{GO}$ is the GO density that is estimated to be 1.32 g cm$^{-3}$.

**Silica-based gel film.** The silica-based gel film was prepared by cross-linking of tetrathydroxy orthosilicate under weak basic surrounding, using CLO of GC as the template. We introduced 1 ml tetrathydroxy orthosilicate into 2 ml methanol/water solution of GO (vol:alcohol:water = 9:1, $\phi$ = 0.38%), and stirred for 12 h to get a homogenous solution. After injecting few drops of ammonia solution (pH = 11) and stirring, the mixture was loaded into a circular disc with a diameter of 3 cm, and kept at room temperature for 24 h. Finally, we obtained the self-standing, silica-based gel film with CD activity.

**Preparation of GO and graphene fibres.** During the-spinner process for GO fibres (Supplementary Fig. S9), the GO dispersions were loaded into glass syringes and injected into the NaOH/methanol solution under 1.5 MPa held by N$_2$. The silica fibres were casted into the cavity, loaded into liquid nitrogen, transferred into the cryotransfer and the sample was cut by a blade. The fracture was sublimed for 1 h before imaging. SAXS characterizations. This work is funded by the National Natural Science Foundation of China (2010R10021), Fundamental Research Funds for the Central Universities (2009JQ044), National Key Basic Research Program of China (2009Q044), Qianjiang Talent Foundation of Zhejiang Province (2010R10021), Fundamental Research Funds for the Central Universities (2009QNA040), Research Fund for the Doctoral Program of Higher Education of China (20100110110049) and Zhejiang Provincial Natural Science Foundation of China (20100110110049).

**Acknowledgements**

We thank Professor S.F. Chen, Dr H.Q. Zhao, Dr Y.Z. Jin and Dr M. Hanfi for the helpful discussions and the staffs in the Shanghai Synchrotron Radiation Facility for SAXS characterizations. This work is funded by the National Natural Science Foundation China (20097403 and 11374162), National Key Basic Research Program of China (2007CB711603), National Science Foundation of China (2010101110049).
Author contributions
Z.X. and C.G. conceived and designed the research, analysed the experimental data and wrote the paper; Z.X. conducted the experiments; C.G. supervised and directed the project.

Additional information
Supplementary Information accompanies this paper at http://www.nature.com/naturecommunications

Competing financial interests: The authors declare no competing financial interests.

Reprints and permission information is available online at http://npg.nature.com/reprintsandpermissions/

How to cite this article: Xu, Z. & Gao, C. Graphene chiral liquid crystals and macroscopic assembled fibres. Nat. Commun. 2:571 doi: 10.1038/ncomms1583 (2011).

License: This work is licensed under a Creative Commons Attribution-NonCommercial-Share Alike 3.0 Unported License. To view a copy of this license, visit http://creativecommons.org/licenses/by-nc-sa/3.0/