Multifunctional, supramolecular, continuous artificial nacre fibres

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Nature has created amazing materials during the process of evolution, inspiring scientists to studiously mimic them. Nacre is of particular interest, and it has been studied for more than half-century for its strong, stiff, and tough attributes resulting from the recognized “brick-and-mortar” (B&M) layered structure comprised of inorganic aragonite platelets and biomacromolecules. The past two decades have witnessed great advances in nacre-mimetic composites, but they are solely limited in films with finite size (centimetre-scale). To realize the dream target of continuous nacre-mimics with perfect structures is still a great challenge unresolved. Here, we present a simple and scalable strategy to produce bio-mimic continuous fibres with B&M structures of alternating graphene sheets and hyperbranched polyglycerol (HPG) binders via wet-spinning assembly technology. The resulting macroscopic supramolecular fibres exhibit excellent mechanical properties comparable or even superior to nacre and bone, and possess fine electrical conductivity and outstanding corrosion-resistance.

Natural composites of seashell nacre and bones with hierarchical micro- and nano-structures have been studied for at least half-century due to their extraordinary mechanical properties, complemented by unique biological functionalities. The microscopic architecture of nacre is depicted as the classic “brick-and-mortar” (B&M) model, which is constituted of highly aligned rigid CaCO₃ platelets sub-micrometre thick surrounded by 10–50 nm elastic organic glue layers. Numerous efforts have been addressed to mimic the nacre structure since 1990, using inorganic tablets such as clay, Al₂O₃, and layered double hydroxides (LDH) and organic linear polymers such as polyvinyl alcohol (PVA), polyelectrolytes, chitosan, and poly(methyl methacrylate) (PMMA). Various strategies have also been developed to fabricate bio-inspired materials, including layer by layer (LbL) deposition, freeze-drying assembly, and vacuum filtration-assisted assembly. However, only films and papers with limited size (cm scale) of nacre mimics have been accessed so far, mainly confined by the substrate-assisted synthetic pathways and the relatively poor solubility/dispersibility of the constituent inorganic particles.

Thus, to achieve the goal of continuous processing for nacre mimics, the synthetic strategy is needed to be reformed, and simultaneously the solubility of inorganic platelets must be fundamentally improved, through either reducing their thickness or increasing their surface functional groups. Accordingly, graphene, the thinnest known sheet in the universe, would be an ideal candidate for the construction of macroscopic biomimetic materials. Besides, graphene possesses marvelous mechanical, electrical, and thermally conductive properties, promising wide applications in devices and high-performance composites. However, either pristine graphene made by mechanical exfoliation of natural graphite or chemically converted graphene (CCG) reduced from graphene oxide (GO) exhibits poor solubility in common solvents and aggregation tendency owing to strong π-π stacking interactions. To improve the solubility of graphene, laborious efforts on surface functionalization have been made, but highly soluble graphene that can be used in continuous solution-processing has rarely been accessed (the solubility ≤ 7 mg mL⁻¹).

Here we firstly choose graphene as rigid platelets and hyperbranched polyglycerol (HPG) as elastic glue to construct macroscopic biomimetic materials via industrially viable wet-spinning technology. Due to the high solubility as well as abundant functional groups of HPG, the resultant sandwich-like building blocks, HPG-enveloped graphene sheets (HPG-e-Gs), are highly soluble in common solvents such as N,N-dimethylformamide (DMF) and N-methyl pyrrolidone (NMP) (~50 mg mL⁻¹), and they also can form liquid crystals at high concentrations. The first artificial nacre fibres (ANFs) up to tens of metres in length, with perfect B&M structures of graphene sheets and HPG binders, are continuously spun from their concentrated liquid crystalline (LC) dope. The supramolecular fibres constructed by hydrogen-bonding arrays exhibit eminent properties such as high tensile strength comparable to nacre and bone, fine conductivity, and excellent corrosion-resistance.
Results

Synthesis and characterization of HPG-e-Gs. Figure 1 shows our protocol to make ANFs with three steps: (i) synthesis of building blocks, (ii) formation of LC dope, and (iii) wet-spinning of ANFs.

Four HPG samples with different molecular weights (4.4, 7.2, 87.6, and 125.8 kDa) were used as the organic layers, and the corresponding sandwich building blocks were denoted as HPG1-e-G, HPG2-e-G, HPG3-e-G, and HPG4-e-G, respectively. The fractions of HPG attached on graphene are 16, 34, 49, and 57 wt% calculated from thermal gravimetric analysis (TGA) curves (Supplementary Fig. S1), dependent on the molecular weights. The corresponding hydroxyl densities are 2.11, 4.54, 6.57, and 7.75 mmol g\(^{-1}\), or 35.9, 97.9, 183.3, and 260.5 hydroxyl groups per 1000 carbons of graphene (Table 1), at least twice as high as the functional group density of functionalized graphene reported previously \(30\). Thus, all the HPG-e-Gs exhibit excellent solubility in polar solvents such as DMF and NMP (\(5\) m g m \(^{-1}\), Supplementary Fig. S2) due to the strong interactions between the attached functional groups and solvent molecules.

The HPG-e-Gs are totally dispersed as individual nanosheets in solution, as proved by atomic force microscopy (AFM) measurements. Each side of graphene sheet is evenly coated with a layer of HPG protuberances. As a result, the thickness of HPG-e-Gs increases to 2–8 nm from 0.8 nm of GO, which depends upon the molecular weight of HPG (Fig. 2a and Supplementary Fig. S3). The height of HPG-e-Gs reveals the single-layer nature of the coated HPG, which is confirmed by molecular dynamic (MD) simulations. For HPG2-e-G, the simulated diametre of HPG2 unimolecule is 1.04–1.10 nm (Supplementary Fig. S4). The height of HPG2-e-G nanosheets measured by AFM is 3 nm, which is almost equal to the height of GO sheets (0.8 nm) plus the height of two monolayers of HPG2 (2.08–2.20 nm). So HPG2 should be distributed as single layer on both sides of graphene sheets. The high-resolution AFM image shows that the graphene sheet is decorated with HPG molecules uniformly and continuously, with sparsely unoccupied holes sub-100 nm in size (Fig. 2b). Scanning electron microscopy (SEM) images display that the morphology of freeze-dried HPG-e-Gs changes from grooved/screwed belts tens of micrometres long gradually into individual few-m\(^{\text{m}}\)''bowls'' with increasing the molecular weight of HPG, which is distinct from the relatively stretched morphology of neat GO sheets. This is likely originated from strong cooperative hydrogen bonding among adjacent HPGs upon different thickness of HPG-e-Gs (Supplementary Fig. S5).

| Table 1 | Selected Molecular Data and Electrical Conductivities of HPG-e-Gs Papers |
|---------|---------------------------------------------------------------|
| Samples | \(M_n\)           | PDI | \(\ell_{\text{w}0}\) (%) | \(\ell_{\text{w}1}\) (%) | \(\ell_{\text{w}2}\) (%) | concn\(^{-1}\) (mmol \(-\text{OH}/g\)) | \(d\) (nm) | \(d_{\text{car}}\) (nm) | \(d_{\text{car}}\) (nm) | Conductivity (S m\(^{-1}\)) |
| HPG1-e-G | 4.4 k            | 1.32 | 29.4          | 13.77          | 15.63          | 2.11          | 35.9          | 27.86          | 1.41          | 8.3          |
| HPG2-e-G | 7.2 k            | 1.75 | 44.4          | 10.84          | 33.56          | 4.54          | 97.9          | 10.21          | 3.85          | 0.18         |
| HPG3-e-G | 87.6 k           | 1.32 | 57.0          | 8.39           | 48.61          | 6.57          | 183.3         | 5.46           | 7.14          | 0.38         |
| HPG4-e-G | 125.8 k          | 1.24 | 64.3          | 6.96           | 57.34          | 7.75          | 260.5         | 3.84           | 10           | 12.5\(^*\)  |

\(\text{Number-average molecular weight of HPG.}\) \(\text{Weight loss fraction before 500°C in TGA curve.}\) \(\text{The fraction of residual functional groups of chemically reduced graphene after functionalization.}\) \(\text{The true content of HPG \(\ell_{\text{w}2} = \ell_{\text{w}1} - \ell_{\text{w}0}\).}\) \(\text{The concentration of hydroxyl groups per gram of HPG-e-Gs.}\) \(\text{The average numbers of hydroxyl groups per 1000 carbons of graphene occupied.}\) \(\text{The average numbers of carbon atoms of graphene per hydroxyl group occupied.}\) \(\text{The density of hydrogen bonds.}\) \(\text{Electrical conductivities of free-standing papers of HPG-e-Gs.}\) The \(\text{''*'' indicates that the conductivity is recorded on a thin film on polypropylene.}\)
Stability of HPG-e-Gs. The HPG macromolecules are ultra-stable on graphene sheets, which can not be removed by repeated washing with acid (1 M), base (1 M), and LiCl-containing organic solvents, due to the cooperative interactions of hydrogen bonds, ester bonds, and possible mechanical interlocking between HPG and graphene (Supplementary Fig. S6). Such a robust character can be further extended to chemical reactions. The hydroxyl groups of HPG-e-Gs were modified with succinic anhydride to obtain carboxyl ones, which could be used to grow superfine platinum nanoparticles 1–3 nm (Supplementary Fig. S6). Such a robust character can be further extended to chemical reactions. The hydroxyl groups of HPG-e-Gs were modified with succinic anhydride to obtain carboxyl ones, which could be used to grow superfine platinum nanoparticles 1–3 nm (Supplementary Fig. S6). This modified HPG-e-Gs was stable in DMF, showing HPG unimolecules anchored uniformly on graphene sheets, the inset in a shows the height of HPG4-e-G increased to 8 nm. (c) POM image of HPG2-e-G in DMF (1 mg mL⁻¹) stayed for 30 days without precipitate. (e) HPG2-e-G in DMF (−20 mg mL⁻¹). (f) HPG2-e-G in DMF (−30 mg mL⁻¹), forming highly viscous semi-gel. (g) HPG2-e-G LC spinning gel (−50 mg mL⁻¹). Upon increasing the concentration, HPG2-e-G dispersions changed from viscous homogeneous liquid (e) to semi-gel with limited mobility (f), then to free-standing gel (g) without precipitates, showing the true solution behavior of HPG2-e-G in a good solvent. Free-standing papers of HPG2-e-G (h, i) and HPG3-e-G (j) made by the classic filtration-assisted assembly method. (k) Stress-strain curves of HPG2-e-G (1) and HPG3-e-G papers (2). (l) Representative SEM image of side-view of HPG2-e-G paper, with perfect layered structures. Scale bars are 2 μm (a), 500 nm (the inset in a), 100 nm (b), 100 μm (c) and 1 μm (l).

Continuous ANFs made by wet-spinning. Wet-spinning technology, which is simple, efficient, scalable, economical, and green, has been widely used in industry for more than 50 years to make tens of kinds of fibres with millions of tons annually, and it has also been utilized to fabricate carbon nanotube (CNT) fibres36–38, Kevlar40, chitosan41, and polyaniline42 fibres. However, this technology has never been tried to spin nacre-mimic fibres, probably because of the poor solubility of inorganic particles and the phase separation between inorganic and organic components in their dispersions. For the first time, we use wet-spinning strategy to capture nacre-mimic fibres from LC dopes of HPG-e-Gs (Supplementary Fig. S9). As the spinning dopes were injected into a coagulation bath (e.g., saturated solution of NaOH in methanol, ethyl acetate, acetone, or their mixture), HPG2-e-Gs procreated flexible and continuous fibres up to tens of metres (Fig. 3a), whereas HPG1-e-Gs and HPG4-e-Gs brought brittle fibres just in centimetre-scale length and HPG3-e-Gs shaped slightly flexible fibres in limited length. The fine flexibility of HPG2-e-G fibres promised us to make knots in μm-dimension (Fig. 3b), additionally burdening 3 g clips without breakage (Supplementary Fig. S10). Considering the continuity of the spun fibres, we found that the methanol solution of NaOH was the most favoring bath for spinning HPG2-e-Gs. Upon immersing in the coagulation bath, the wet fibre in gel state consisting of aligned
HPG2-e-Gs coagulated, and the solvent (DMF) rapidly diffused out of the fibre into the methanol with simultaneous back-diffusing of methanol into the fibre. The flow-induced alignment could be maintained by the NaOH solution that allowed the nanosheets to rapidly stack together as they were coming out of the nozzle. Continuous fibres with 10–100 μm in diameters were then collected onto the spinning drum outside the bath, followed by washing with methanol to remove the residual NaOH. The actual diametres of fibres mainly depended on the nozzle size, injection rate, flow conditions, and coagulation procedure.

The surface of dried fibres shows the orientation of nanosheets along the fibre axis (Fig. 3c), and anode contour is observed in the cross-section (Fig. 3d), caused by the shrinking of dope during the coagulation procedure. Inside the fibres, HPG-e-G hybrid building blocks are found stacking together to form densely ordered lamellar microstructures, resembling the B&M structure of nacre (Fig. 3e,f, Supplementary Fig. S11). So we name it artificial nacre fibres (ANFs).

Under low magnification of SEM, we observed curled and folded lamellae, which were ascribed to the dislocation domains in the LC dope (Fig. 3e). To confirm the incorporation of graphene in ANFs, the fibres were further characterized by transmission electron microscopy (TEM) and Raman spectroscopy. Similar to the SEM observations, layered structures of black graphene sheets are also identified (Supplementary Fig. S12). Raman spectra of ANFs show the characteristic peaks of graphene: the defect-induced D band at ~1,353 cm⁻¹, and G band at ~1,602 cm⁻¹ that is related to in-plane vibration of sp² carbon atoms in a 2D hexagonal lattice (Supplementary Fig. S13). The intensity ratio of D to G band (I_D/I_G) for ANFs (~1.43), which is related to the size of the sp² domains, is close to that (~1.59) of HPG2-e-G building blocks. These results indicate that the chemical structure of graphene hardly changes during the wet-spinning process.

Tensile tests reveal that σ, E, and ultimate strain (ε) of ANFs are 125±10 MPa, 8.2±2.2 GPa, and ~3.7%, respectively (Fig. 4a). The tensile strength is comparable to those of nacre (130 MPa) and osteons in bone (102–125 MPa), while the ultimate strain is about 3 folds higher than that of nacre, implying superior toughness of our ANFs. The much better flexibility of ANFs is attributed to the much thinner hard inorganic platelets and self-adapting ability of supramolecular interactions between soft HPG layers during sliding of HPG-e-G building blocks under tension. From the polymer point of view, although HPG2 has null tensile strength as it is a viscous fluid at room temperature with glass transition temperature (T_g) of ~48.4 °C (Supplementary Fig. S14), its composite fibres demonstrate much greater values of both σ and ε than nacre-like films (~76–105 MPa, ε ~0.6–0.97%) containing extremely stronger organic layers (~30–50 MPa) of well-known linear chitosan and PVA. This result, together with the aforementioned case of HGP-e-G films, declares that dendritic polymers, even without any strength, can also be employed to construct high strength composites.

The good mechanical performance of our fibres is ascribed to their similar hierarchical structures to nacre and the efficient load transfer between graphene sheets and HPG moieties. At the nanometre scale, protuberances of HPG unimolecules on the graphene sheets (Fig. 2a,b) closely resemble the nanograins on tablets of nacre. At the microscale, the wrinkling morphology of graphene sheets is analogous to the waviness of CaCO₃ platelets in nacre. In higher level architecture, the alternatively organized graphene and HPG layers corresponds with the classic B&M structure of nacre. Such hierarchical structures conduce to the improvement of
mechanical strength. Moreover, the HPG interlayers favor the load transfer between two graphene sheets, spreading the load all over the fibre to increase the tensile strength. This is in accordance with the previous report on composite papers of GO and PVA, which demonstrates the cooperative intersheet hydrogen bonding network between GO and PVA can enhance the load transfer efficiency and increase the mechanical property. The effective load transfer makes the attractive mechanical properties of graphene sheets “survive” in the composite fibres.

Metal ions-crosslinked ANFs. Previous investigations on nano-structures of nacre demonstrated that mineral bridges embedded in the “mortar” layer were not only able to control crack propagation in nacre, but also increase the mechanical performance of the interfaces in the natural layered materials.8,10,11 Divalent ions such as Mg2+ and Ca2+ had also been utilized to cross-link GO sheets for making strong GO papers.14 Thus, to further enhance the strength and stiffness of ANFs, we introduced Mg2+ to bridge the adjacent “bricks” by coordinate interactions, mimicking the mineral bridges in nacre. The hydroxyl groups of HPG in HPG-e-G building blocks were firstly converted into carboxyl groups, followed by wet-spinning and immersing in the aqueous ionic solution to afford metal ions-modified ANFs. X-ray energy dispersive analysis proved the uniform distribution of magnesium element in the cross-section of fibres (Supplementary Fig. S15). The resulting Mg2+-modified ANFs show considerably enhanced mechanical strength (σ 145±18 MPa) and stiffness (E 10.0±2.6 GPa), accompanying with fine flexibility (ɛ ~1.5%) (Fig. 4a). The enhancing effect is resulted from two main factors. First, the coordinate interactions between Mg2+ and carboxyl groups in Mg2+-modified ANFs are stronger than the hydrogen bonds in the unmodified ANFs. Second, the “cavities” between neighboring HPG molecules and “defects” at the wrinkles and LC dislocation domains in unmodified ANFs could be filled up with Mg2+ during the ion cross-linking process, further enhancing the mechanical strength of the resulting fibres. Notably, the Mg2+-modified ANFs present superior mechanical performance to GO papers cross-linked by divalent ions of Mg2+ and Ca2+ (σ 80.6–125.8 MPa, ɛ 0.33–0.50%),14 and poly(allylamine)-crosslinked GO papers (σ 91.9 MPa, ɛ 0.32%)50 (Fig. 4b), mainly due to crumbling and twisting of graphene building blocks in fibres that can bring additional friction force under tension but rarely occurred on papers of well stacked, flattened GO sheets. Our fibres also exceed nacre (σ 130 MPa, ɛ 0.9%)1 and artificial nacre films composed of clay and poly(diallyldimethylammonium) chloride via LbL assembly (σ 95–109 MPa).14 ANFs with ~34 wt% polymer are even ahead of neat graphene materials in tensile strength. For instance, the tensile strength of Mg2+-modified ANFs is comparable or superior to neat GO fibres (102 MPa)51 and papers (63.6–130 MPa)52,53, as well as reduced graphene fibres (140 MPa)54 and graphene papers (150 MPa)55 (Fig. 4b). In addition, the E of our composite fibres is higher than those of neat GO fibres (5.4 GPa) and reduced graphene fibres (7.7 GPa).56

Corrosion-resistance and electrical conductivity of ANFs. The chemical resistance of graphene17 and the compact stacked structure of ANFs endure the composites with good corrosion-resisting property. After being immersed in hydrochloric acid (1 M), NaOH (1 M), and saturated LiCl solution in DMF for three days, the ANFs still kept excellent mechanical strength and flexibility (σ ~90–100 MPa, ɛ ~1–5%) (Fig. 5a), and they were knotted without breakage after being immersed for two weeks (Fig. 5b,c). On the contrary, bubbles were released when nacre was immersed into a hydrochloric acid aqueous solution due to the reaction between CaCO3 tablets and acid (Fig. 5d), and the hard nacre completely disappeared after around one day with a little residue of soft protein membrane (Fig. 5e). Other bio-fibres such as silk and spider silk mostly composed of protein were broken into failed pieces after being immersed in a NaOH aqueous solution for at most three days (Fig. 5f–i).

Apart from the extraordinary mechanical performance and corrosion-resistance, our fibres are electrically conductive, with conductivity of ~0.24 S m−1 for ANFs and ~4.88 S m−1 for Mg2+-modified ANFs. As expected, the electrical conductivity is comparable to small molecules-functionalized graphene (0.1–30 S m−1)57 and linear polymer-functionalized graphene (0.84 S m−1)58, but inferior to neat reduced GO papers (170–8100 S m−1)59 and graphene fibres (~2.5×103 S m−1). The Mg2+ ions induced the cross-linking of neighboring graphene sheets and partially restored their conjugated networks, thus the conductivity was improved. In Raman spectra of ANFs (Supplementary Fig. S13), the 1/L0 decreases from ~1.43 to ~1.10 after the introduction of Mg2+, suggesting a possible increase in the average size of sp2 domains of graphene sheets.60

Discussion

As shown in Fig. 4a, ANFs exhibit an elastic behavior with small deformation (0–0.5%). At high deformation before breakage (0.5–2.3%), ANFs show a plastic behavior with a relative plateau in the

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**Figure 4** Mechanical properties of ANFs. Typical stress-strain curves for ANFs (a). The strain rate is 5% per minute. The numbers 1 and 2 denote HPG2-e-G and Mg2+ enhanced HPG2-e-G fibres, respectively. Comparison of tensile strength and strain with a set of nanocomposites with layered structures (b).
stress-strain curve. This mechanical characteristic is in accordance with the shear lag model for nacre \(^1,3\). At the tensile stress less than 50 MPa, the interface between two HPG adlayers, starts to yield in shear and the HPG-e-G building blocks slide on one another, accompanying with the destruction and reformation of the hydrogen-bonding network among HPG macromolecules confined in the intersheet channel. Upon increasing the strain, the load transfers from the soft organic moieties to hard graphene sheets, then the shear spreads all over the fibre, followed by failure of fibre by pullout of HPG-e-G building blocks (Fig. 3g,h, Supplementary Fig. S16). Based on the "pull out" breaking mechanism, the total force in tension (\(F\)) could be expressed by:

\[ F = F_1 + F_2 \]

where \(F_1\) represents the force provided by the network of hydrogen-bonding between HPG interlayers (Fig. 3i), \(F_2\) denotes mechanical friction force between building blocks, which consists of \(f_1\) (friction force provided by the rough surface with protuberances of HPG unimolecules on graphene sheets), and \(f_2\) (friction force generated by the deformation of flat building blocks such as wrinkling, folding, and torsion at micro-scale). Accordingly, the excess mechanical strength of ANFs mainly originates from \(F_2\) (or \(f_2\), as compared with the previous nacre-mimetic films/papers built with relatively flat and thick inorganic tablets\(^8,10\).

On the hydrogen bonding network (\(F_1\)) between graphene sheets, Buehler and coworkers have made systematic analysis in a joint experimental theoretical and computational study with the example of GO-PVA paper system\(^46\). Since the chain entanglement between PVA molecules is not considered (because the degree of polymerization for PVA is 5) in simulations, it is reasonable to introduce the simulations results to our system of graphene and HPG (in which only hydrogen bonding works). The density of hydrogen bonds on graphene sheets in our ANFs is \(\sim 3.85\) hydrogen bonds nm\(^{-2}\) (Table 1). At the similar situation, the MD simulations show a modulus of 34 GPa for GO-PVA layered papers\(^46\), whereas the experimental modulus of our ANFs is only 8.2 ± 2.2 GPa. In fact, there are lots of joints and defects among HPG-e-G sheets in our ANFs that are not considered in the MD simulations, which will be adverse to the mechanical performance. In addition, the MD simulations indicated that ultrahigh tensile strength (\(\sim 1.7\) GPa) could be achieved for the GO-PVA papers\(^46\), which suggested that the mechanical performance of graphene-based nacre-mimics could be further improved by optimization of assembly conditions. Furthermore, \(F_3\) can be significantly improved by introduction of stronger supramolecular interactions\(^54,55\) such as coordination, electrostatic forces\(^6,9\) and covalent linkages\(^7,11\). On the other hand, the mechanical strength of ANFs can be improved by replacing dendritic polymers with strong linear or crystalline polymers such as PVA\(^7,8\), chitosan\(^10,12\), cellulose, nylon\(^56\), and aromatic polyamides\(^57\).

Given the hierarchical structures of ANFs, the mechanical performance can be highly improved by design and optimization of their primary (e.g., size of inorganic sheets and molecular structure of organic glue), secondary (e.g., strong occlusive\(^8\)/chain entangled/ crosslinked\(^11\) interlayer), tertiary (e.g., regular and compact cross section), and quarternary (e.g., twisting along the fibre axis) structures. Figure 6 shows three kinds of proposed secondary structures, aiming to enhance the interactions of organic molecules between adjacent inorganic layers. First, nano-asperities on the tablets mechanically occlude each other (Fig. 6a), increasing the mechanical friction force under tension. Such structures might be accessible by

![Figure 5](https://www.nature.com/scientificreports)

**Figure 5** | Corrosion-resistance of ANFs. Typical stress-strain curves for ANFs after being immersed in different solutions for 3 days (a). The numbers 1, 2 and 3 denote ANFs after being immersed in 1 M HCl, 1 M NaOH, and saturated solution of LiCl in DMF, respectively. Photographs of ANFs knots after being immersed in 1 M NaOH (b) and 1 M HCl (c) for two weeks. Photographs of a nacre immersed in 1 M HCl for 0 h (d) and 24 h (e), silk immersed in 1 M NaOH for 0 h (f) and 72 h (g), and spider silk in 1 M NaOH for 0 h (h) and 48 h (i). All of the natural materials have been completely destroyed after being immersed in either acid or basic solution, while our ANFs are still quite strong and robust after being treated with acid and basic aqueous solutions as well as highly polar organic solvent. Scale bars, 3 mm (b,c).
Figure 6 | Proposed secondary structures for stronger ANFs. Mechanically occlusive interaction resulted from the rigid organic teeth arrays between two tablets (a). Chain entanglements or crystallization of polymer brushes grafted on the tablet surfaces (b). Cross-linked interlayer polymer chains (c).

Growing polyaniline nanowires on GO sheets. Second, the polymer chains on adjacent tablets entangle each other (Fig. 6b), resulting in superhigh van der Waals forces. Third, covalent cross-linking of polymer chains between tablets could also produce ultra-strong and ultra-stiff composites (Fig. 6c). For example, the tensile strength of nacre-like films made by LbL assembly of PVA and clay was enhanced from ~150 to ~400 MPa by cross-linking with glutaraldehyde, and the corresponding modulus was increased from ~13 to ~106 GPa simultaneously.

In conclusion, we have advanced nacre-mimics from limited films to continuous fibres with wide range of promising applications. The macroscopic-assembled fibres possess perfect BKM structures constructed by graphene and HPG, and exhibit excellent mechanical property beyond nacre as well as extra corrosion-resistance and good electrical conductivity. We have demonstrated that null strength and mechanical properties of such strong materials through large-area supramolecular arrays, breaking a new path to advanced supramolecular materials. The compositions of bio-mimetic fibres can be extended to other 2D colloids (e.g., clay, LDH, WS2, MoS2, Al2O3, and metallic platelets) and dendritic/linear polymers, expanding the content of both nanocomposite fibres and bio-mimics.

Methods

Preparation of HPG-e-Gs. GO was synthesized from natural graphite powder (40 μm in size, Qingdao Henglide Graphite Co., Ltd.) using the same protocol reported previously. HPG was synthesized by ring-opening anionic polymerization with acetaldehyde, and the corresponding modulus was increased from ~13 to ~106 GPa simultaneously.

Preparation of HPG-e-Gs papers and fibres. The papers of HPG-e-Gs were prepared by vacuum-assisted filtration of DMF solution with concentration of 5–10 mg mL−1, followed by drying at 80 °C in vacuum for 12 h. For the spinning of HPG-e-G fibres (Supplementary Fig. S9), the HPG2-e-G dispersion in DMF (25–50 mg mL−1) was loaded into a 1 mL plastic syringe with a spinning nozzle (PEK tube with diameter of 60, 100, 160 μm), and injected into the NaOH/methanol solution by an injection pump (20 μL min−1). After coagulation for 10–15 minutes, the fibres were rolled onto the drum, washed by water to remove NaOH and dried for 24 h under 80 °C in vacuum. The spinning process of carboxyl abundant HPG2-e-G fibres was similar to the above procedure except that the coagulation bath was changed to the mixture of acetone and ethyl acetate (1:1 in vol).

Instruments. AFM images were taken in the tapping mode by carrying out on a NSK SPI3800 or a Nanoscope IIIa scanning probe microscope, by spin-coating sample solutions onto freshly cleaved mica substrates at 800 rpm. TGA was carried out on a Perkin-Elmer Pyris 6 TGA instrument with a heating rate of 20 °C min−1 under a nitrogen flow (30 mL min−1). SEM images were obtained on a Hitachi S4800 field-emission SEM system. TEM analysis was performed on a JEOL JEM1200EX electron microscope at 120 kV, or a FEI/Philips CM200 electron microscope operating at 200 kV. POM observations were performed with a Nikon E600POL and the liquid samples were loaded into planar cells for observations. FTIR spectra were recorded on a Bruker Vector 22 spectrometer. The tensile stress-strain tests were performed on a Microcomputer Control Electronic Universal Testing Machine made by REGER in China (RGWT-4000-20). The strain rate is 1 mm min−1 with the gauge length of two centimetres. The electrical conductivity of fibres was measured using a four-probe resistivity instrument (RTS-4, PROBES TECH). UV-visible spectra were obtained with a Varian Cary 300 Bio UV-visible spectrophotometer.

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