Multifunctional graphene woven fabrics

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Tailoring and assembling graphene into functional macrostructures with well-defined configuration are key for many promising applications. We report on a graphene-based woven fabric (GWF) prepared by interlacing two sets of graphene micron-ribbons where the ribbons pass each other essentially at right angles. By using a woven copper mesh as the template, the GWF grown from chemical vapour deposition retains the network configuration of the copper mesh. Embedded into polymer matrices, it has significant flexibility and strength gains compared with CVD grown graphene films. The GWFs display both good dimensional stability in both the warp and the weft directions and the combination of film transparency and conductivity could be optimized by tuning the ribbon packing density. The GWF creates a platform to integrate a large variety of applications, e.g., composites, strain sensors and solar cells, by taking advantages of the special structure and properties of graphene.

The creation and modulation of the nanomaterials’ structure and function are of great scientific interest and importance. Nanomaterials (especially one-dimensional, 1D) are important building blocks for nanoscale electronics1,2, as they can function as miniaturized devices as well as electrical interconnects3,4. A typical example of macrostructures from self-assembly is metamaterial5. Metamaterials are artificial materials engineered to have unique properties gained from structure rather than composition, using small inhomogeneities to create effective macroscopic behavior6,7. Another example is woven or non-woven fabrics of nanomaterials, including organic electronic fibres for logic circuits8, oxide nanowires for photoswitches and memories9, electrospun nanofibers for composites10,11, “smart” textiles for sensors and displays12, woven electrodes for photovoltaics13, self-organized silicon nanowire network for highly-integrated crossbar nanocircuits14.

Regarding carbon nanomaterials, carbon nanotube (CNT) 1D long fibers, two-dimensional (2D) thin films and three dimensional (3D) arrays and sponges have been widely investigated15. Cross-stacked CNT films have been used for transparent and stretchable conductors16,17. Recently, graphene macrostructures have also attracted a lot of attention, e.g., foam16,17, mesh18, pattern19, ribbons20, which take the advantages of the outstanding properties of graphene to make full use of its macroscale assembly.

In this work, we report a simple route to realize a graphene-based woven fabric (GWF), that is assembled by intersecting graphene micron-ribbons (GMRs). To our knowledge, this is the first report on the woven fabric structure made from aligned GMRs. The structural and characteristic features of our GWFs are: (i) It possesses high structural integrity and much better mechanical performance compared with polycrystalline graphene films; (ii) In contrast to monolayer graphene, it has good gas/liquid permeability (due to the presence of micron holes); (iii) Upon external loading, it shows an interesting exponential resistive response which is favourable for high sensitivity strain sensing. Based on the unique structural features and properties, we demonstrate the following applications of our GWFs: (i) GWF/polymer composites, which are well aligned GMRs embedded in polydimethylsiloxane (PDMS) to produce 2D network with multi-joint conductive channels, or deposited on PDMS as strain sensors. (ii) GWF/semiconductor solar cells, in which the GWF functions as the transparent electrode in Schottky junction to provide enhanced photovoltaic conversion. Most importantly, the periodic voids of GWFs can be filled with other functional materials, or served as a permeable membrane of electrolytes for solar cell applications.

Results

Structure and morphology of the GWF. GWFs were grown by atmospheric CVD using copper meshes as substrates (see Methods section). Fig. 1a shows the three main steps used in the fabrication of GWFs: (i) CVD growth of graphene on copper mesh, (ii) removal of copper wires with FeCl3/HCl aqueous solution, and (iii) collapse of graphene to form double layered GMRs. The CVD grown GWF retains the network configuration of the copper mesh. For the cases described here, the copper mesh consists of copper wires (≅60 µm in diameter),
arranged in a crisscross pattern. Fig. 1b shows the optical and plan-view scanning electron microscope (SEM) images of a copper mesh before and after graphene growth. In the next step, the GWFs can be collected from the liquid surface with desired target substrates. The details are described in the Methods section. Fig. 1c shows the as-obtained GWF films floating on water and deposited on glass and PET, clearly demonstrating the well-aligned arrays of GMR lines, obtained GWF films floating on water and deposited on glass and PET. Scale bars, 200 μm. (c) Optical images of GWF films floating on water and deposited on glass and PET. Scale bars, 5 mm. (d) TEM image of a GMR and selected area electron diffraction pattern from the region marked with a yellow box. Scale bars, 50 nm (left), 5 (1/nm) (right).

Figure 1 | Fabrication of GWFs by CVD using copper wire meshes as substrates. (a) Schematic of steps for GWF preparation. (b) Macroscopic optical images (left), top-view SEM images (right) of copper meshes before (top) and after (bottom) graphene growth. Scale bars, 200 μm. (c) Optical images of GWF films floating on water and deposited on glass and PET. Scale bars, 200 μm.

GWF/polymer composites. We then show the fabrication of GWF/polymer composite films (bonding with inert, transparent polymer, e.g., PDMS). To obtain the GWF/PDMS composite mesh, before etching away the copper skeleton by FeCl₃/HCl solution, a thin layer of PDMS with controlled thickness was deposited on the surfaces of the graphene-coated copper wires, followed by cross-linking and curing the PDMS as a support to prevent graphene networks from collapsing during copper etching. As shown in Fig. 2d, real free-standing GWF/PDMS composite films can be obtained after copper etching. In spite of the thin layer of PDMS coating, this composite structure still has good optical and electrical properties. As revealed in the insets of Fig. 2d, the free-standing GWF/PDMS composite film can be twisted by tweezers without fracture. The use of the PDMS support layer was critical for preparing the free-standing GWF composite film. This method can keep the 3D structure of GWFs (see bottom inset of Fig. 2d). As further revealed in Fig. 3, the composite film is woven by interconnected graphene/PDMS micron-tubes, forming a multi-joint channel system. During CVD growth, two interconnected copper wires will melt and join together at the crossing point at 1000°C, resulting in a monolithic structure of the copper mesh.
Carbon was introduced by decomposing methane, and graphene was then grown on the surface of the interconnected copper wires. In the resulting composite (Fig. S4), graphene serves as a lining of the PDMS hollow structure and could be used as protective coating, reinforcement phase, electrical and thermal conducting channels in the composites.

To form fully filled GWF/PDMS composites (Fig. S3a and Fig. S4), a large amount of PDMS has been applied. The transparency of the composite film is high as the GWF is semi-transparent, and the GWF mesh was planarized and stabilized by filling the spacing between GMRs with PDMS.

Tunable morphology and optoelectronic properties. The diameter of the copper wire and the mesh density directly determine the structure of GWFs. The three frames of Fig. 3a show optical images of three GWFs produced using copper meshes woven with 100 μm thick wires of different packing density. The corresponding representative small-area SEM views of these three samples are shown in Fig. 3b. Visual inspection of the samples and high-resolution images collected from them indicate excellent spatial uniformity and low defect density. The packing density of GMRs is also tunable by post manipulation (Fig. S5), for instance, controlling the liquid flow during film transfer.

Due to the unique configuration, the GWF becomes inherently strong (compared with polycrystalline graphene films) and conductive. The optoelectronic performance in terms of sheet resistance \( R_s \) and transparency of GWF strongly relies on the geometrical parameters, such as GMR width, spacing (related to graphene coverage density). As shown in Fig. 3c,d, the film transparency is varied owing to the cavity of the GWF. The \( R_s \) of 500–2500 Ω/sq and 200–1200 Ω/sq after HNO₃ treatment at the corresponding transmittances of 50%–90% are obtained. The resistance values still cannot meet the criteria for transparent conducting applications that may replace the conventional indium tin oxide (ITO). The combined conductivity and transparency property of the GWF is also not advantageous due to its polycrystalline structure and the presence of surface wrinkles which will greatly reduce the conductivity and transparency of GMRs, in turn, GWFs. The conductivities of the GWFs are expected to be improved by synthesis process optimization and post acid doping.

GWF/PDMS hybrid films. GWFs were transferred on PDMS substrates to make GWF/PDMS hybrid films, and their mechanical properties (in tension) were studied. The interlacing (or crossing) points are the major locations where interactions between GMRs take place, through which the GWF forms an interlocked structure. We demonstrate that conductive GWF/PDMS hybrid films can act as electromechanical sensors under uniaxial tension. Tensile test accompanied by simultaneous electrical measurement shows a significant increase of resistance during the tensile cycle. The strain dependent resistance variations of GWF/PDMS films were investigated and summarized in Fig. 4a. As shown in the schematics (left panel), the X direction is defined as the direction of GMR lines and the XY direction is defined as the diagonal direction. The electromechanical response differs along different directions and under different strains. The hybrid film shows certain flexibility and can be stretched within a relatively low strain range (＜5%) and the resistance returns to the initial value upon unloading (Fig. 4a). Irreversible tensile strain and resistance changes occur at relatively high strain levels. The hybrid film shows a significant jump of resistance (by several orders of magnitude, up to 10^4 times) when it was
stretched to a strain of 12% (Fig. 4a), due to the fracture occurring along the boundaries within the polycrystalline GMRs (Fig. S6). When the hybrid film is returned to 0% strain, the resistance increases to 3~5 times of its original value. In comparison, when stretched along the XY direction, the quadrilateral GMR network is self-adjusting. The stress applied on GMRs is smaller than that when the same strain is applied along X direction. If the PDMS substrate was pre-stretched before GWF deposition, the resistance variation becomes more gently. As shown in Fig. 4a, the resistance change is only 3 times when a pre-stretched sample was stretched to 10% along XY direction. Similar resistance change has been observed in several other graphene/PDMS25 and CNT/PDMS26 composites under stretching. Our GWF/PDMS hybrid film shows reversible resistance change (ΔR) relative to the original value (R₀) over many cycles and maintains a non-linear relationship versus strain in every cycle. Fig. 4b shows the results at maximum stains of 2% and 5% for a GWF/PDMS sample along X direction. The resistance shows an increase of 25 times under 2% strain and 230 times under 5% strain. Based on this result, the GWF can not only be used as a strain sensor for composites, if high resolution and large image fields of view could be obtained, the resistance variation and optical diffraction from optical images might provide information for the deformation of the composite. As a demonstration, the GWF/PDMS hybrid film was further fixed to an index finger to perform as a stretchable sensor. The resistance was detected at different finger motion. As shown in Fig. 4c, taking about 45° as a step, the resistance of the hybrid film increases with finger bending, which is similar to the results in the stretching experiments.

GWF/Si Schottky solar cells. Alternative approaches to achieve transparent electrodes have investigated the use of conductive nanostructures such as graphene, CNTs and metallic nanowires, or conductive polymers such as poly(3,4-ethylenedioxythiophene)/poly(styrene sulfonic acid) (PEDOT:PSS). Though the GWF is not a good window electrode material for transparent conductor applications, the periodic voids can be filled with PEDOT, or served as permeable membrane of electrolytes for photovoltaic applications. Our techniques have provided transparency on the order of 70~80%, with conductivities in the 1000 Ω/sq range. GWF/Si solar cells were first fabricated from the GWF transparent electrodes and compared with similar devices fabricated using graphene film28. The solar cells with GWF electrodes outperformed devices having graphene transparent electrodes in efficiency (η) (>2% and up to 3.8%, see Fig. S7). While these parameters are sufficient for some flexible electronics applications, the resistive losses remain too high for many scaled applications. The discontinuity of the GWF films also leads to other issues impacting device performance. As shown in Fig. 5a, the GWF-Si solar cell was doped by HNO₃ fume. Acid infiltration of GWF networks boosts the cell efficiency by reducing the internal resistance to improve the fill factor (FF) and by forming photoelectrochemical units that enhance charge separation and transport29,30. In this way, the efficiency of
solar cells could reach 6.1% (2-fold increase), with $V_{oc}$ of 0.53 V, short-circuit current density ($J_{sc}$) of 15.6 mA/cm², and fill factor (FF) of 58.4% at AM 1.5 (80 mW/cm²).

Based on the GWF/Si solar cell, PEDOT was further added and filled in the cavity of the GWF. PEDOT was first spin coated and then annealed at 140°C for 15 min, producing a GWF-PEDOT composite film. The introduction of PEDOT, commonly used as a buffer layer, facilitates the hole injection and extraction (Fig. 5b). The work function of GWF is 4.8 eV and the occupied molecular orbital of PEDOT is 5.1 eV. This indicates a large hole-injection barrier between the composite film and Si with a conduction band of 5.2 eV, thus resulting in higher $V_{oc}$ and FF and leading to a 40% increase in efficiency. Furthermore, as the GWF networks provides many gaps, redox electrolyte containing 40% HBr and 3% Br₂ was dropped on the GWF side, forming a hybrid solar cell which consists of two types of photovoltaic systems, including a Schottky junction cell and a photoelectrochemical cell connected in parallel (see the corresponding energy diagram in the inset of Fig. 5c), cooperating with each other\(^\text{11,32}\). By this treatment, we obtain a 30% increase in efficiency (Fig. 5c).

Discussion
The easy operation in transferring is partly due to the favourable mechanical properties of the woven structure and the multilayer fishnet stacks. As the GWF is composed of GMRs, both the adhesion and friction between the two sets of GMRs (the warp and the weft) caused by the hairness of the ribbon surface play important roles in determining the fabric strength. When the fabric is being stretched in the warp direction, the weft ribbon would shrink more and intensify the friction between the two sets of GMRs. Similar to a traditional woven fabric, the clamping action of the weft on the warp would reduce the unevenness of the warp GMRs, resulting in enhanced warp ribbon strength. On the other hand, the weft GMRs might share some of the external load. The absence of visible cracking or tearing suggests high levels of physical toughness and mechanical flexibility. On the basis of a conventional theory\(^\text{10}\), the strength of the GWF has been predicted (see Fig. S8 and calculation details in Supplementary Information). Although such structures are mechanically robust, handling or otherwise manually manipulating them is difficult owing to their polycrystalline feature. As a result, similar to CVD grown graphene films, the strength of a GWF cannot be measured by direct tensile test. Practical applications still require a supporting substrate of some type (either liquid or solid).

As of the electromechanical properties of the GWF/PDMS hybrid film, the repeatable results of stretching in two different ways reveal that the GWF/PDMS resistance is sensitive to the elongation rather than the shear strain. The step-by-step crack formation in GWF upon external loading provides non-linear resistive response which is essential for highly sensitive strain sensing. Since a strain gauge always gives a linear response on
elongation, the exponential response on elongation shown here might implicate a subtle metal-insulator transition\textsuperscript{34} by strain engineering due to the macroscopic motion of graphene flakes in polycrystalline GMRs, which requires further investigations to understand the underlying physics.

The greater electrical conductivity of the GWF film (as thin as 1 nm) would make it possible to retain adequate lateral electrical conductivity while reducing the thickness of, or eliminating entirely, the traditional semitransparent conductive layer. The GWF film is coated conformally onto a patterned substrate to form heterojunction with the underlying semiconductor (e.g. Si, CdSe, GaAs with band gaps of 1.0–2.5 eV). As shown in the energy band diagrams (Fig. 5), our calculations indicate that carbon makes a Schottky junction with silicon\textsuperscript{30}, which is favorable for producing a relatively large built-in field and charge separation. A space charge region is formed in the silicon near the interface. This indicates that the GWF film not only serves as a transparent electrode for light illumination, but also an active layer for electron/hole separation and hole transport. During this process, light absorption by the silicon produces photo-generated carriers (electron/hole pairs); ii) photogenerated holes (h\textsuperscript{+}) and electrons (e\textsuperscript{-}) are separated and accelerated by the built-in electric field; iii) the holes are driven into the GWF film and the electrons are driven into the silicon, respectively to produce electricity. The use of GWF films in this type of solar device is a particularly attractive approach for several reasons: the extremely high surface area offers a tremendous opportunity for exciton dissociation; in addition to enhancing photovoltaic conversion efficiency, the incorporation of mesh structure can potentially improve mechanical and environmental stability.

In summary, transparent, flexible electrodes using GMRs woven in a mesh configuration are prepared by the CVD approach. The GMRs are assembled into a fabric mesh having 100–200 μm spacings in orthogonally directed layouts. The GWFs are mechanically flexible because of their 2D flat morphology, with lengths greater than centimeters in some cases. The GWFs can be further optimized for conductivity and transparency through controlling of the micron-ribbon ratios and spacing. The ease of handling and the low fabrication cost have made fabrics attractive for structural and functional applications (e.g. composites and solar cells). The GWFs are compatible with plastic substrates and scalable to high throughput roll-to-roll or printing processes, thus showing the potential to

**Figure 5 | GWF-based solar cells.** Device schematics, energy band structure diagrams and J-V curves of (a) GWF/Si solar cell, (b) PEDOT filled GWF/Si solar cell and (c), hybrid Schottky and PEC GWF/Si solar cell.
impact future flexible electronics technology. Because of the presence of crystallographic defects like grain boundaries, the GMRs are not yet very strong, far below their ideal mechanical strength.

We are in the process of trying to develop these GWFs as a new class of flexible photovoltaic and composite materials. Our approach shows a possibility for full integration of flexible electronics within a woven smart textile. If properly tailored, hybrid electronic and optical properties of our GWFs will be utilized for future nano-optics and nano-optoelectronic engineering. The optimization and further development of this GWF technology will promise many promising applications, including heat insulation, ventilating but water and liquid proof, radiation resistant coatings, filters, masks, templates, logic circuits, TEM grids, catalyst carriers, etc.

**Methods**

**Synthesis of GWFs.** Copper meshes with different line width and spacing were used as scaffold templates for the CVD growth of GWFs. The mesh was cut into pieces of 5 × 8 cm² which depended on the space of the CVD reactor tube and then was ultrasonically treated with hydrochloric acid and acetone to clean their surfaces and eliminate the thin surface oxide layer. Then the mesh was heated to 1000 °C in a CVD furnace under Ar (500 mL/min) and H₂ (50 mL/min). After reaching 1000 °C, the Ar was turned down to 200 mL/min and H₂ to 20 mL/min. A small amount of CH₃OH (3–15 mL/min) was then introduced into the reactor at ambient pressure. After 10 min growth, the mesh was rapidly cooled down to room temperature. Like graphene grown on copper foils, GWFs from copper meshes are also polycrystalline.

**Characterizations.** The samples were characterized using optical microscope (OLYMPUS BX60), scanning electron microscope (SEM, LEO1530, 10 kV), transmission electron microscope (TEM, JEM2010, 200 kV), Raman spectroscopy (Renishaw, RM2000, He-Ne laser excitation line at 514 nm). To measure its transmission and sheet resistance, the GWF was transferred to a highly polished quartz substrate. Optical transmission spectra were taken by UV, Hitachi 3-1010 optical spectrometer.

**Polymer composites.** GWF-on-PDMS hybrid films: GWF was put into FeCl₃/HCl mixture (1 : 1 mol/L) to dissolve the copper. Fig. S9 illustrates the etching process of a single copper wire. The lower part of the copper wire immersed in the solution was etched away firstly. During etching, the graphene film would not spread since the upper graphene is still wrapping on the copper surface, thus keeping the folding state of lower graphene. Then the upper part exposed to air gradually sank into the solution and pulled the lower part. About 2 h later, the copper dissolved completely, and GWF collapsed to form the planar mesh structure floating on the solution. The width of the resulting GWF approximately equals to the diameter of the copper wire. Solidified PDMS (1 mm thick) film was used to collect the GWF.

**GWF-in-PDMS composites:** To avoid structural failure of the GWF during copper etching, the lower layer of PDMS was used as a support to reinforce the GWF structure. The copper mesh after graphene growth was drop-coated with PDMS solution, and then baked at 60 °C for 12 h. After the PDMS solidified completely, the composites were put into FeCl₃/HCl mixture (1 : 1 mol/L) to dissolve the copper at 80 °C. About 5 h later, the copper dissolved completely to give the free-standing GWF/PDMS composites.

**GWF/Si solar cells.** n-Si (100) wafers (doping density: 1.5–3 × 10¹⁷/cm³) with a 300 nm SiO₂ layer were patterned by photolithography and wet-etching of oxide (by hydrofluoric acid solution) to make square windows (0.1 cm²) where n-Si was used as scaffold templates for the CVD growth of GWFs. The mesh was cut into pieces of 5 × 8 cm² which depended on the space of the CVD reactor tube and then was ultrasonically treated with hydrofluoric acid solution to make square windows (0.1 cm²) where n-Si was used as scaffold templates for the CVD growth of GWFs. The mesh was cut into pieces of 5 × 8 cm² which depended on the space of the CVD reactor tube and then was ultrasonically treated with hydrofluoric acid and acetone to clean their surfaces and eliminate the thin surface oxide layer. Then the mesh was heated to 1000 °C in a CVD furnace under Ar (500 mL/min) and H₂ (50 mL/min). After reaching 1000 °C, the Ar was turned down to 200 mL/min and H₂ to 20 mL/min. A small amount of CH₃OH (3–15 mL/min) was then introduced into the reactor at ambient pressure. After 10 min growth, the mesh was rapidly cooled down to room temperature. Like graphene grown on copper foils, GWFs from copper meshes are also polycrystalline.

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Author contributions
H.W.Z. and X.L. conceived and designed the experiments. X.L., P.Z.S., L.L.F. performed the experiments. All authors interpreted the results. H.W.Z. and X.L. co-wrote the manuscript.

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

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