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OPEN Hydrophilic Graphene Preparation from Gallic Acid Modified Graphene **Oxide in Magnesium Self-Propagating High Temperature Synthesis Process**

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Hydrophilic graphene sheets were synthesized from a mixture of magnesium and gallic acid (GA) modified graphene oxide (GO) in a self-propagating high-temperature synthesis (SHS) process, and hydrophilic graphene sheets displayed the higher C/O ratio (16.36), outstanding conductivity (~88900 S/m) and excellent water-solubility. GO sheets were connected together by GA, and GA was captured to darn GO structure defects through the formation of hydrogen bonds and ester bonds. In SHS process, the most oxygen ions of GO reacted with magnesium to prevent the escape of carbon dioxide and carbon monoxide to from the structure defects associated with vacancies, and GA could take place the high-temperature carbonization, during which a large-area graphene sheets formed with a part of the structure defects being repaired. When only GO was reduced by magnesium in SHS process, and the reduced GO (rGO) exhibited the smaller sheets, the lower C/O ratio (15.26), the weaker conductivity (4200 S/m) and the poor water-solubility because rGO inevitably left behind carbon vacancies and topological defects. Therefore, the larger sheet, less edge defects and free structure defects associated with vacancies play a key role for graphene sheets good dispersion in water.

Graphene is a novel 2-D carbon nano-material, and it is made up of conjugated sp² carbons and arranged in a honeycomb structure, which makes graphene to display the excellent electrical, mechanical, thermal and optical properties¹⁻⁷. Graphene can be prepared by physical or chemical methods, such as chemical vapor deposition (CVD), ultrasonic exfoliation of graphite, epitaxial growth and graphene oxide (GO) reduction⁸⁻¹⁵. The epitaxial growth and CVD methods can be used to provide the large area and almost free-defect graphene sheets for applications in electronics⁷, but those free-defect graphene sheets were difficult to be mass-produced.

Known as GO rich in oxygen containing groups such as hydroxyl, carbonyl and epoxy group, and the chemical reduction of GO is the most efficient method for low-cost and large-scale production of graphene¹²⁻¹⁵. Hydrazine¹³, NaBH₄¹⁴ and ascorbic acid¹⁵ et al. had been used to reduce GO, and hydrazine and NaBH₄ were effective in removing oxygen of GO. When N2H4 was used as reducing agent, N can be doped in the as-prepared graphene. Although NaBH₄ showed excellent ability in GO reduction, the hydrolysis property made it difficult to get a stable NaBH₄ aqueous solution¹⁶. Furthermore, it is usually considered that it is difficult to form large area graphene or free-defect sheets though chemical reduction¹⁷. The high electrical conductivity (56500 S/m) of graphene was obtained via thermal exfoliation of GO at low temperature and then annealing at 1900 °C under vacuum conditions¹⁸. Graphene also can be prepared by magnesium thermal reduction of GO with annealing at 650 °C for 6 h in Ar atmosphere¹⁹, but the high temperature reduction of GO under inert atmosphere condition leads to the number of CO or CO₂ elimination from the carbon lattice to form the atomic vacancies or graphene fragment, which cause the structural deterioration of as-prepared graphene²⁰. Most importantly, graphene prepared from GO reduction and annealed at high temperature all exhibited the high hydrophobic property.

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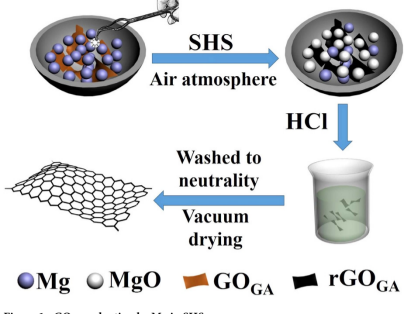


Figure 1. GO_{GA} reduction by Mg in SHS process.

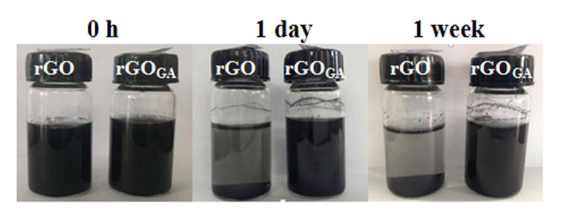


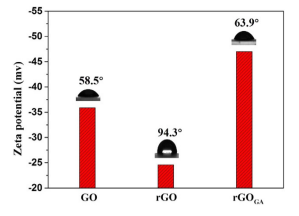
Figure 2. Dispersions of rGO and rGO_{GA} in aqueous solution at different time intervals (~1mg/mL).

SHS represents an attractive alternative to conventional methods for materials synthesis^{21–23}. This technique, which involves self-propagating reactions of either solid-solid or gas-solid type, is characterized by the fact that, once the starting mixture is ignited by means of external thermal sources for relatively short times, highly exothermic reactions propagate through the mixture in the form of self-sustained combustion wave, progressively leading to final products without requiring additional energy. Reaction characteristics include the high rates of the combustion front $(10^{-4}-10^{-3} \text{ m/s})$, generation of high temperatures $(1700~2000 \,^{\circ}\text{C})$, even to 4000 $^{\circ}\text{C}$), and rapid heating $(10^{3} - 10^{4} \text{ K/s})$. The SHS technique is also characterized by process simplicity, short reaction time, easy-to-build equipment, low energy requirements, and the possibility of obtaining complex or metastable phases. GA has three hydroxyl and one carboxyl groups in its benzene ring which has the same oxygen-containing groups with GO. The network holes of GO can be darn with GA by hydrogen bonds and ester bonds, and GO sheets can also be connected together by gallic acid. In this paper, GO was modified by GA to form GO_{GA}, and GO_{GA} was further reduced by magnesium in SHS process under air atmosphere condition (Fig. 1). This environmentally friendly technology is easy to achieve the higher C/O ratio, high conductivity and excellent water-solubility graphene sheets.

Results

In order to discuss the effects of addition amount of GA on sheet area, smooth, edge defects and hydrophobic property of graphene sheets, 1.0 g, 1.5 g and 2.0 g GA were loaded into equal mass of GO solution, and then reduced by magnesium with the same methods. The resulting materials are denoted as rGO_{GA1} , rGO_{GA} and rGO_{GA2} , respectively.

The water solubility of rGO and rGO_{GA} were investigated in deionized water at different time intervals. It can be seen from Fig. 2, rGO has lower dispersibility in aqueous solution, but rGO_{GA} can be uniformly dispersed in aqueous solution for a long time. Zeta potential is a direct parameter on reflecting the interaction





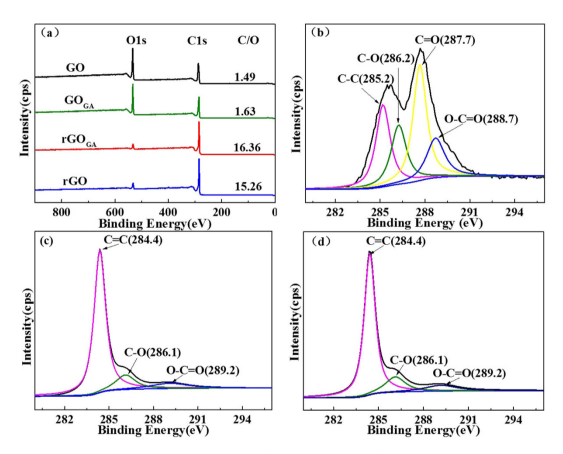


Figure 4. XPS survey spectra (a), C1s XPS spectra of GO (b), rGO_{GA} (c) and rGO (d).

among suspended particles. The zeta potentials of aqueous GO, rGO and rGO_{GA} were -35.9 mV, -24.6 mV and -47.0 mV, respectively (Fig. 3). Particles with zeta potentials more negative than -30 mV are considered to form stable suspension due to interparticle electrostatic repulsion^{24,25}. Therefore, rGO precipitated within one day, and rGO_{GA} dispersed in aqueous solution to form a stable colloidal for long time, indicating the excellent water-solubility of rGO_{GA}. But beyond that, contact angle measurements were used to further prove the hydrophilic of rGO_{GA}. It was noticed that the contact angle of the GO, rGO, and rGO_{GA} were 58.5°, 94.3° and 63.9°, respectively. And compared rGO_{GA}, the contact angle of rGO_{GA1} and rGO_{GA2} were 76.4° and 96.3° (Figure S1). The test results proved once again that the excellent hydrophilicity of rGO_{GA}.

In order to investigate the reasons for excellent water-solubility of rGO_{GA} , XPS was used to investigate the chemical structure and composition of the prepared samples. Only C1s and O1s peaks can be observed in the XPS survey scan (Fig. 4a), and no hetero atoms (including Mg) were doped into graphene. As for rGO_{GA} and rGO, the intensity of the O1s band dramatically decreases and the intensity of the C1s band significantly increases, suggested that oxygen-containing groups had been effectively removed. The C/O atomic ratio of GO and GO_{GA}

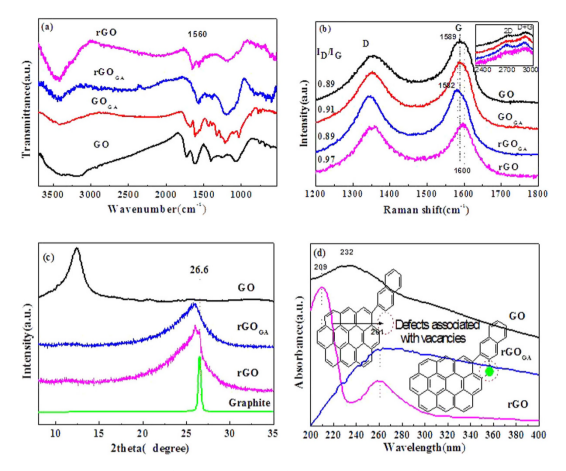


Figure 5. (a) ATR-IR spectra of GO, GO_{GA} , rGO_{GA} , and rGO. (b) Raman spectra of GO, GO_{GA} , rGO_{GA} , and rGO. (c) XRD patterns of graphite, GO, rGO_{GA} and rGO powder. The 2 θ angles of the XRD peaks (d-spacing) of GO and rGO_{GA} shifted from 12.4° (d-spacing ~0.8 nm) to 25.8° (d-spacing ~0.34 nm) after reduction. (d) UV-vis absorption spectra of GO, rGO_{GA} and rGO.

are 1.49 and 1.63. However, the C/O atomic ratio of rGO_{GA} and rGO increased to 16.36 and 15.26, indicating the full reduction of GO_{GA} . Most importantly, C/O atomic ratio of rGO_{GA} and rGO is much higher than that of GO reduced by other chemicals, such as hydrazine (10.3)¹³, NaBH₄ (8.6)¹⁴ and LiAlH₄ (12)²⁶. C1s XPS spectra of GO (Fig. 4b) showed a peak at 285.2 eV that corresponded to C-C bonds of carbon atoms in a conjugated honey-comb lattice²⁷, and other three different peaks centered at 286.2 eV, 287.7 eV and 288.7 eV are also observed, corresponding to single or double bonds in aromatic rings, i.e. C-O (epoxy and alkoxy), C=O, and O-C=O groups, respectively. Figure 4c,d showed that C=O groups completely disappeared in rGO_{GA} and rGO, indicating that the delocalized π -conjugated structure was well restored in rGO_{GA} and rGO²⁸. Importantly, the C/O atomic ratio of rGO_{GA} was higher than that of rGO, but rGO_{GA} displayed much better hydrophilic than rGO. Those results are far beyond our expectations. A hypothesis was proposed that graphene composition is not the main factor to influence the hydrophilic or hydrophobic property, but its structure characteristics must play a key role on its hydrophilic or hydrophobic property.

Figure 5a showed the FT-IR spectra of GO, GO_{GA} , rGO_{GA} and rGO. A series of absorption bands ranging from 1000 to 1750 cm⁻¹ exhibited the presence of C=O bonds (at 1720 cm⁻¹), O-H bending (at 1621 cm⁻¹), O-H deformation (at 1406 cm⁻¹), C-O groups in epoxy (at 1221 cm⁻¹) and C-O groups in alkoxy (at 1057 cm⁻¹)^{29,30}. Hydroxyl (OH) groups have an intensive absorption band between 3200 and 3700 cm⁻¹, due to the presence of moisture intercalated within hydrophilic GO sheets³⁰. After modified with gallic acid (GO_{GA}), the characteristic bands for oxygen containing groups did not change remarkably, but some new FT-IR absorptions of GO_{GA} appeared in the range of 1490–1190 cm⁻¹, which indicated that gallic acid molecules were adsorbed or darned on GO²⁵. It is noteworthy that the C=O band of GO_{GA} is shifted by 44 cm⁻¹ from 1720 cm⁻¹ to 1676 cm⁻¹ compared to GO, corresponding to the formation of O=C-O-C groups, which demonstrates that a portion of gallic acid are combined with GO by esterification. After samples undergo magnesium self-propagating high temperature synthesis process, the characteristic bands for oxygen-containing groups of rGO and rGO_{GA} almost completely disappeared, but the bands of aromatic carbon (smaller π -conjugated C=C structure) at 1575 cm⁻¹ and (Larger π -conjugated C=C structure) at 1560 cm⁻¹ appeared. These observations suggested that GO and GO_{GA} were well reduced, and the larger-area π -conjugated structure was established in rGO_{GA}.

GO, GO_{GA}, rGO_{GA} and rGO were further characterized by Raman spectroscopy. As shown in Fig. 5b, there are two significant peaks for these four samples, assigned to D band (\sim 1350 cm⁻¹) and G band (\sim 1585 cm⁻¹), which are E2g vibration mode in-plane and A1g breathing mode³¹. The appearance of a distinct D band in the spectrum

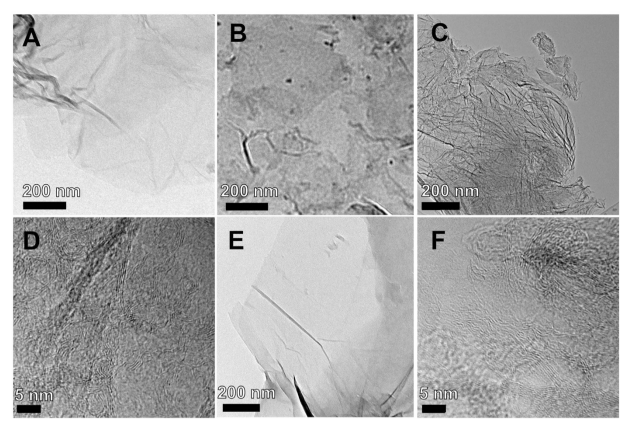


Figure 6. TEM images of GO (**A**), $GO_{GA}(\mathbf{B})$, rGO (**C**) and rGO_{GA} (**E**). High resolution TEM images of rGO (**D**) and rGO_{GA} (**F**).

is an indication of disorder in graphene originating from the defects associated with vacancies and grain boundaries³². As compared to the G band of GO, that of rGO_{GA} was shifted by 7 cm⁻¹ from 1589 to 1582 cm⁻¹, displaying a high reduction degree³³. Previous reports exhibited that the size of the defect-free sp² cluster regions is the inverse of the ratio of the D and the G band integrated intensities $(I_D/I_G)^{32,34,35}$. In the case of GO, the I_D/I_G ratio was 0.89, and the I_D/I_G ratio of GO_{GA} increased to 0.91. However, I_D/I_G ratio of rGO was 0.97, rGO_{GA} I_D/I_G ratio declined to 0.89. From those results, the following conclusions can be drawn: (1) the most oxygen ions of GO reacted with magnesium to form MgO, reducing the number of carbon atoms vacancies of the carbon lattice; (2) the average size of the sp² domains of rGO_{GA} was larger than rGO slightly; (3) gallic acid can repair the structure defects of graphene during magnesium thermal reaction. Apart from D band and G band, there are two Raman bands with weaker intensity called 2D and D + G located at 2680–3000 cm⁻¹ as shown in insets of Fig. 5b. Zhan *et al.*²⁸ discovered that it could be utilized to distinguish the electronic conjugation of GO and rGO_{GA} by comparing these two bands. Compared with rGO (or GO), the enhancement of I_{2D}/I_{D+G} ratio of rGO_{GA}.

XRD was used to characterize graphite, GO, rGO_{GA} and rGO. Figure 5c shows the 2 θ peak of graphite powder was at 26.6° (d-spacing ~0.34 nm) and a broad peak near 12.4° (d-spacing ~0.80 nm) was observed for GO, indicating that the graphite was fully oxidized. Compared with the parent GO, the peak of rGO_{GA} shows an obvious shift to higher 2 θ angles (25.8°; d-spacing ~0.35 nm), suggesting that rGO_{GA} was well ordered with two-dimensional sheets and well removal of surface functional groups¹¹. Compared to rGO_{GA} , a weak shoulder peak at 26.5° of rGO was found, which revealed a small portion of the reduced graphene sheets agglomerated together or formed graphite structure³⁶.

Samples were also characterized by UV-vis absorption, as shown in Fig. 5d, GO has a intensive absorption band at 235 nm, which was due to the $\pi \rightarrow \pi^*$ transition of aromatic C=C bond. In addition, a weaker absorption shoulder at around 303 nm, which was attributed to the $n \rightarrow \pi^*$ transition of the C=O bond³⁷⁻⁴⁰. The $\pi \rightarrow \pi^*$ transition band of rGO_{GA} and rGO red shifted to 264 nm owing to the restoration of the conjugated structure, which indicated the removal of oxygen-containing functional groups and the disappearance of $n \rightarrow \pi^*$ transition band^{41,42}. However, some $\pi \rightarrow \pi^*$ transition band of rGO shifted to 209 nm, indicating the defects associated with vacancies. Those results further proved that GA took place the high-temperature carbonization to repair the structural defects during SHS process, therefore, a large-area graphene sheets of rGO_{GA} formed with a part of the structure defects being repaired during a SHS process.

Figure 6A,B showed the TEM images of GO and GO_{GA} . By comparison, GA was adsorbed on the surface and edge of GO by the interaction between GA molecule and defects of GO (Fig. 6B). Figure 6C,E showed the TEM images of few-layer rGO and rGO_{GA}. There are some wrinkles and scrolls in rGO sheets like the reported work²⁵, and the TEM analysis of rGO showed the edge of rGO sheets generated irregular defect with the formation of

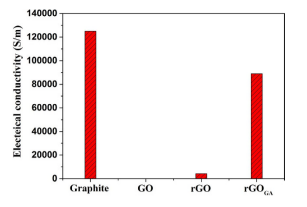


Figure 7. The electrical conductivity of different samples.

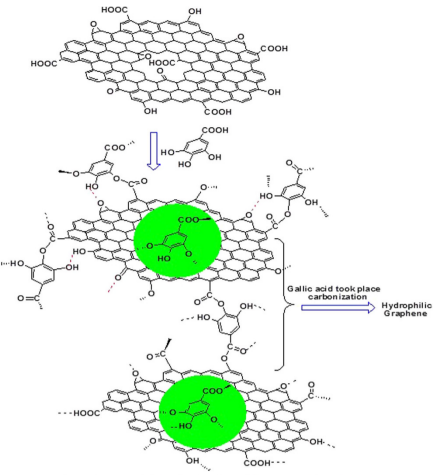
small fragment. However, rGO_{GA} showed a larger sheet, nearly no wrinkles and scrolls, and less edge defects. In addition, Figure S2A,b showed the TEM images of few-layer rGO_{GA1} and rGO_{GA2}. Due to little addition amount of GA, it still can be seen that small number of the edge and inside defects from Figure S2A. It can be shown that only parts of graphene defects were restored. In contrast to rGO_{GAL} Figure S2A showed that there were more carbon impurities which were obtained by magnesium thermal reduction of excess GA on the surface of graphene sheets. The high-resolution TEM (Fig. 6D,F) clearly exhibits the signature image of the few-layer graphene with the number of layers ranging from 3 to 8. The measured lattice space of this material is about 3.45 Å, which is in good agreement with the thickness of a monolayer graphene. The biggest difference between Fig. 6D,F are that the small layers of rGO are messy accumulation, but the large layers of rGO_{GA} are in an ordered arrangement because of the restoration of GA. These differences are attributed to the generation of atomic vacancies in the rGO sheets, and the atomic vacancies had a high mobility at high temperature. The amount of generated atomic vacancies was so large that rGO sheets disintegrated and caused a significant structural deterioration at high temperature²⁰. Several experimental and modeling studies have focused on exploiting surface roughness to engineer super hydrophobicity^{43,44}. It is generally accepted that roughness-induced hydrophobicity. rGO_{GA} has a highly polished surface because of the larger sheet, smooth, no fold, less edge defects. So it shows the more hydrophobic property. Based on those characterizations, a conclusion can be drawn that the wrinkles, scrolls, irregular structure defects were associated with vacancies and irregular grain boundaries rather than the C/O atomic ratio to determine the hydrophilic or hydrophobic property of graphene sheets. In summary, the larger sheet, smooth, no fold, less edge defects and the structure defects associated with vacancies play a key role for graphene sheets good dispersion in water media.

The electrical conductivity is considered a highly sensitive indicator of the extent to which electronic conjugation is restored after deoxygenation of GO. The electrical conductivity of compressed-powder samples of the pristine graphite, GO, rGO and rGO_{GA} were measured at room temperature (Fig. 7), and the electrical conductivity are 125000 S/m, 0.012 S/m, 4200 S/m and 88900 S/m, respectively. The conductivity of rGO_{GA} is much high than that of rGO, and closely approaches that of pristine graphite. In addition, the electrical conductivity of rGO_{GA1} and rGO_{GA2} are 29760 S/m and 59277 S/m, respectively (Figure S3). Conclusion can be drawn that the numerous defects associated with vacancies and carbon impurities in graphene sheets play the key role on the lower electrical conductivity of grapheme.

The following mechanism was suggested for our reduction process (Fig. 8). Gallic acid and GO can be connected together by forming a number of intermolecular hydrogen bonds. Importantly, Gallic acid and GO can take place the esterification reaction at 90 °C for 24 h. Therefore, gallic acid can act as the medium to connect together with adjacent GO sheets though ester bond and hydrogen bonds. After gallic acid took place the high-temperature carbonization during magnesium high temperature thermal reduction process, a part of pyrolytic carbon self-assemble into a graphene structure, and another portion of carbon filled inside of the graphene atomic vacancies to repair of structural defects. Magnesium thermal reaction is a classical reaction which is usually used to reduce metal oxide. Magnesium can capture oxygen atoms of metal oxide to generate corresponding metal element and magnesium oxide. Importantly, previous article reported that burning magnesium metal in dry ice resulted in few-layer nanosheets of graphene in high yields⁴⁵. In other words, Magnesium also can capture the oxygen atom of carbon dioxide. So during the SHS process, magnesium could capture the oxygen atom from GO, which also prevent the escape of carbon dioxide and carbon monoxide to from the structure defects associated with vacancies. Therefore, a large-area hydrophilic graphene sheets were formed by a SHS process.

Conclusion

In conclusion, a novel and facile approach to the synthesize graphene sheets based on magnesium SHS process is reported using GO modified by gallic acid (GO_{GA}) as precursor. The merit of this method is that the process carried out in the air atmosphere without external heating source, and gallic acid can repair the structure defects of graphene during SHS process. The obtained graphene sheets could be stably dispersed in water for more than one month. The wrinkles, scrolls, irregular structure defect associated with vacancies and grain boundaries (rather than C/O atomic ratio) determine the hydrophilic property. While the less edge defects and free structure defects associated with vacancies play a key role for graphene sheets good dispersion in H₂O. Moreover, the synthetic



GO can be connected together by gallic acid to form a number of intermolecular hydrogen bonds and ester bonds

Figure 8. Suggested mechanisms for the modification of gallic acid and the reduction of GO_{GA} with Gallic acid-assisted magnesium reduction of GO in a SHS process.

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process can be utilized to produce large-scale graphene by an environmentally friendly way. Obviously, these advantages will facilitate its industrial applications.

Materials and Methods

Materials. Natural graphite powder (80 mesh) was supplied by Tianjin Chemical Co., Ltd and used as received. NaNO₃ (99.8%), KMnO₄ (99.5%), H_2SO_4 (98%), HCl (35%), magnesium powder and magnesium ribbon *et al.* were purchased from Tianjin Kermel Chemical Reagent Co., Ltd.

Characterization. FT-IR was recorded from a TESOR 37 (BRUKER Corporation, Germany) and operated by attenuated total reflectance (ATR) in the wavenumber range of 4000–500 cm⁻¹. Raman spectra were obtained by using a micro-Raman system on (Renishaw, RW1000-In Via) with an excitation energy of 2.41 eV (l=514.5 nm). XRD patterns were measured on a Bruker XPS (Germany) using Cu K α radiation. UV-vis detection was carried out by UV/Vis Spectrometer (Thermofisher, USA). TEM and HRTEM images were obtained from a JEM-2100 transmission electron microscope (JEOL, Japan). Electrical conductivity of samples was measured on a ST2722 semiconductor powder resistivity tester (Suzhou, China). X-ray photoelectron spectroscopy (XPS) analysis was carried by (Thermofisher, K-alpha).

Sample preparation. Preparation of GO. GO was prepared by modified Hummers' method⁴⁵. 5.0 g graphite, 2.5 g NaNO₃, and 125 ml H₂SO₄ were loaded in a 1000 ml round-bottom flask with magnetic stirring. After being cooled with ice-water bath at 0 °C for 1 h, 17.5 g KMnO₄ was added into above reaction system. After reaction was kept at 0–5 °C for 1 h and then heated at 35 °C for 2 h, the mixture was diluted with 250 ml deionized water and reacted at 98 °C for 45 min. 200 ml deionized water and 50 ml H₂O₂ (30%) were added into above suspension under the condition of magnetic stirring. After centrifugation and dry, GO was obtained.

Preparation of GO_{GA} . Exfoliation of GO was obtained by ultrasonication in water for 2 h. GO_{GA} was prepared in three steps: (1) 1.5 g gallic acid was loaded into 500 ml GO (5 mg/ml) solution, and the mixture was dispersed by ultrasound for 40 min. (2) the mixed solution was allowed to stand at room temperature for 3 h. (3) The mixed

solution was dried at 90 °C under air atmosphere for 24 h to obtain the mixture of gallic acid-modified GO (GO_{GA}). As a control, different addition amount of gallic acid (1.0 g and 2.0 g) in equal mass of GO (were labeled as GO_{GA1} and GO_{GA2}) were prepared with the same method.

Preparation of rGO_{GA} , rGO_{GA2} , rGO_{GA2} and rGO. 3.0 g Magnesium powder and 1.5 g GO_{GA} powder were mixed evenly. The mixture was put in a crucible, and mixture was covered with a thin layer of magnesium powder on the surface of mixture. Mixture was ignited by magnesium ribbon under air atmosphere condition. After the reaction is over, the mixed products were transferred to a beaker containing 250 ml of 3 M HCl. Products were transferred into the acid solution to remove MgO and residual Mg. The resulting rGO_{GA} was collected with filtration, washed with deionized water until the filtrate turned out to be of neutral. Obtained rGO_{GA} was dried at 60 °C under high vacuum conditions for 10 h. As a control, rGO_{GA1} , rGO_{GA2} and rGO were prepared from GO_{GA1} , GO_{GA2} and GO reduction by magnesium with the same methods.

Contact angle measurement. The contact angle of the samples was measured on DSA 100 (KRUSS) contact angle system at ambient temperature. Before the measurement, 20 mg power samples were dispersed in 500 ul ethanol with ultrasonic treatment 1 h. In the following, the suspension was dropped onto a glass slide to form a uniform film.

Electrical conductivity measurement. Electrical conductivity of samples was measured on a ST2722 semiconductor powder resistivity tester (Suzhou, China). The powder sample was filled into a test slot, then applying a pressure about 18 MPa. The electrical resistivity tester would reflect the electrical resistivity of the sample directly. The electrical conductivity of samples was calculated through the following equations:

$$Electrical \ conductivity = 1/Electrical \ resistivity$$
(1)

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Author Contributions

L.C., Z.L. and B.C. have contributed to the design of the study and the critical revision of the article. L.C. and K.S. did the experiments, analyzed the data, prepared figures and drafted the article.

Additional Information

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