Three-Dimensional Sulfur/Graphene Multifunctional Hybrid Sponges for Lithium-Sulfur Batteries with Large Areal Mass Loading

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In this communication, we introduce the concept of three dimensional (3D) battery electrodes to enhance the capacity per footprint area for lithium-sulfur battery. In such a battery, 3D electrode of sulfur embedded into porous graphene sponges (S-GS) was directly used as the cathode with large areal mass loading of sulfur (12 mg cm\textsuperscript{-2}), approximately 6–12 times larger than that of most reports. The graphene sponges (GS) worked as a framework that can provide high electronic conductive network, abilities to absorb the polysulfides intermediate, and meanwhile mechanical support to accommodate the volume changes during charge and discharge. As a result, the S-GS electrode with 80 wt.% sulfur can deliver an extremely high areal specific capacitance of 6.0 mAh cm\textsuperscript{-2} of the 11th cycle, and maintain 4.2 mAh cm\textsuperscript{-2} after 300 charge-discharge cycles at a rate of 0.1C, representing an extremely low decay rate (0.08% per cycle after 300 cycles), which could be the highest areal specific capacity with comparable cycle stability among the rechargeable Li/S batteries reported ever.

In recent years, durable, affordable and high energy advanced rechargeable batteries are urgently required because the state-of-the-art energy storage technologies, especially lithium-ion batteries, are critical enabling technologies for the development of advanced, portable electronic devices, electric vehicles, and sustainable energy generation systems\textsuperscript{1,2}. Among these promising energy storage systems for next-generation, lithium-sulfur batteries have received special attentions due to the high theoretical specific capacity (1675 mAh g\textsuperscript{-1}) and high specific energy (2600 Wh kg\textsuperscript{-1}) at a moderate voltage of 2.2 V vs. Li/Li\textsuperscript{+}. In addition, the cathode material elemental sulfur also has other advantages, including its natural abundance, low cost (about $150 per ton), and low environmental impact\textsuperscript{2,5–11}. However, despite of these advantages, their commercialization are hindered due to the poor cycle performance, which is caused by the natural insulating of sulfur and Li\textsubscript{2}S, the dissolution and transportation of polysulfide intermediates in the electrolyte, as well as mechanical instability of the electrode caused by volume changes during charging/discharging cycles\textsuperscript{2,5–15}.

Over the past few decades, research and development efforts to address these issues are focused on the improvement of the conductivity and prevention of the polysulfide dissolution and shuttling. Of notable successes, carbon materials (such as graphene\textsuperscript{9,12,16–19}, carbon nanotubes\textsuperscript{20,21}, carbon nanofibers\textsuperscript{10,22}, carbon sphere)\textsuperscript{6,23}, conductive polymers (such as PEDOT: PSS\textsuperscript{24}) or metal oxides (such as TiO\textsubscript{2}\textsuperscript{11}) are used to modified the sulfur-based cathodes. As a result, high specific capacities exceeding 800 mAh g\textsuperscript{-1} in terms of sulfur have been achieved by many research groups. However, the commercialized application of the Li-S system hasn’t been realized till now, because of that, most of the as-prepared electrodes have extremely low areal mass loading (<2 mg cm\textsuperscript{-2})\textsuperscript{9,11,12,24}, which will greatly decrease the real capacity of the battery systems and cannot comparable with the commercialized lithium ion batteries, while the typical mass for the lithium ion battery electrode is as high as ~20 mg cm\textsuperscript{-2}.

To the best of our knowledge, the highest sulfur loading is reported by Zhang et al., in his paper, the sulfur loading is as high as 12 mg cm\textsuperscript{-2}\textsuperscript{26}. However, a heavy current collector (carbon cloth, ~14 mg cm\textsuperscript{-2}) was used and the electrochemical performance is really very poor with fast fading rates and low capacity at a current rate of 0.5 mAh cm\textsuperscript{-2}. Most recently, a great deal of work have been done to increase the areal mass loading, including introducing 3D structure electrodes\textsuperscript{27}. In particular, Zhou et al., report a 3D self-assembled graphene-sulfur (G-S) hybrid with good electrical conductivity, showing an outstanding electrochemical properties. It is believed that
developing the 3D structures of graphene will play a huge role on lithium sulfur batteries. Nonetheless, toxic CS$_2$ were used and they did not show how the results will go with increasing of the areal mass loading. Hence, developing facile methods to impregnating sulfur into 3D graphene with high areal mass loading and high capacity is very promising.

In this study, we reported an approach to assemble sulfur-graphene sponge (S-GS) with sulfur uniformly distributed into the pores of graphene sponge that can be directly used as cathode for Li-S batteries with significantly improved areal mass loading and stability. In our approach, self-supporting graphene sponges were firstly obtained using hydrothermal reduction and then sulfur were impregnated into pores of the as-prepared graphene sponges through a heat treatment. In our well-designed structure, porous graphene sponges will not only to improve the overall conductance of the cathode, but also to absorb the soluble polysulfides intermediate.

### Results and Discussion

Figure 1a shows the photos of the typical graphene hydrogel easily prepared through hydrothermal reaction. As can be seen from figure 1b, three GO hydrogel can easily support 500 g weight with little deformation, showing comparable mechanical properties to those of chemically cross-linked polymer hydrogels. In addition, the electrical conductivity was measured by four-probe method. With a conductivity of about 42 S/m, it showed an attractive property for many practical applications. The interior microstructures of GO sponges before and after sulfur impregnated were imaged by scanning electron microscope (SEM) of the freeze-dried samples. As it can be seen from figure 1c, the GO sponges shows a well-defined and interconnected 3D porous network and the sizes of the pores are about 3–10 μm. After the emerged of sulfur, showing in figure 1d, the pores are filled with sulfur while the graphene sponge worked as the framework that provide mechanical support and electronic conductive. Moreover, the absorptivity and the surface area of the S-GS were tested, the results showed that the S-GS can absorb electrolyte more than 18 times its weight and the surface areas decreased from 215.12 m$^2$ g$^{-1}$ to 6.54 m$^2$ g$^{-1}$, which indicates that the cavity in the electrode is enough to absorb the electrolyte and soluble polysulfides intermediate.

![Figure 1](image1.png)

**Figure 1** | (a) Photographs of a 3.5 mg mL$^{-1}$ homogeneous GO aqueous dispersion before and after hydrothermal reduction at 180 °C for 18 h; (b) photograph of three strong GO hydrogel allowing supporting weight; (c) SEM image of the interior microstructures of the GO sponges and (d) SEM image of interior microstructures of the S-GS with 80 wt.% sulfur.

X-ray diffraction (XRD) patterns of the GO, GS and S-GS with 80 wt.% sulfur are shown in Figure 2. For GO, a peak appears at 10.9°, which corresponds to an interlayer spacing of 0.81 nm. After the hydrothermal reaction for 18 h, a new peak appears at 26.3° was appeared and instead of this peak, indicating the decreasing of the spacing, which exhibits that after the hydrothermal reaction, the GO was reduced at some degree. For the S-GS, no obvious change of this peak was found when compare with GS and the sulfur showed sharp and strong peaks, indicating that the sulfur was in a well-defined crystal structure, which is also been detonated by the IR analyst, shown in figure 3, after the impregnation of sulfur, no evidence of the C-S bonds were found (υc-s at 987 cm$^{-1}$), confirming that the sulfur was absorbed by the GO sponges, which agree well with the XRD results.

In order to clearly identify the possible structural advantages of the graphene sponges on the cycle stability of sulfur, electrochemical performance of the S-CG were evaluated. Figure 4 shows the charge–discharge voltage profiles of the 11th cycle measured during galvanostatic cycled at 0.1C (where 1C corresponds to a current density of 1675 mA g$^{-1}$) within 1.5 ~ 3.0 V (vs Li$^+$/Li). As can be seen, two plateaus at 2.35 V and 2.05 V were clearly observed during the discharge process, which correspond to the formation of long-chain lithium polysulfides (Li$_2$S$_x$, 4 ≤ x ≤ 8) and short-chain lithium polysulfides (such as Li$_2$S$_2$ and Li$_2$S), respectively. Such discharge profile is typical for sulfur cathodes. On the basis of the discharge results, the S-GS electrode was able to deliver 6.0 mAh cm$^{-2}$ of the electrode and 625 mAh g$^{-1}$ of sulfur in the 11th discharge process. The obtained low gravimetric specific capacity is because of the large areal mass loading of sulfur. As with the previous reported for lithium ion batteries, thicker electrode cells will be discharged at...
a higher current density, thus tending to the lower impedance and thicker electrode also places more of a strain on the transport of lithium ions in the electrolyte, thus possibly decrease the gravimetric specific capacity.

With continued cycling, as shown in figure 5a, the electrode exhibited well-overlapped and flat plateaus, suggesting good stability and reversibility of the electrode. Figure 5b compares the capacity of the electrode as a function of cycle numbers. Although the initial discharge capacity is relatively low (4.93 mAh cm$^{-2}$ of the electrode and 513 mAh g$^{-1}$ of sulfur), it can continued to increase until the 11th cycle (6.0 mAh cm$^{-2}$ of the electrode and 625 mAh g$^{-1}$ of sulfur). This behavior can be contributed to the activation step of the S-GS electrode because the surface area of the as-prepared S-GS is low, and therefore, it takes some time for the electrolyte to flood the internal surfaces of the GO sponges. Only under this condition, the deeply buried sulfur and disulfide bonds can contact with the electrolyte and become electrochemically active. Subsequently, the capacities almost stabilized and demonstrated little fading upon extended cycling. And as a result, the electrode can exhibit a reversible and comparable specific capacity. Of the electrode, the decay rate was as low as 0.08% per cycle for 300 cycles. At the same time, as can be seen from the inserted map in figure 5b, the coulombic efficiency remained at around 98%. On the basis of such superior cyclic stability, it is reasonable to conclude that the graphene sponges framework could effectively improve the cycle stability of the lithium sulfur batteries, likely through absorption and immobilization of the polysulfides intermediate and provide better mechanical support to accommodate the volume changes during charge and discharge.

**Conclusion**

In summary, we have synthesized a 3D electrode of sulfur embedded into porous graphene sponges for lithium sulfur batteries by a heat treatment, which is simple, highly efficient, and scalable. This 3D architecture electrode was able to demonstrate high areal specific capacity and high retention ratio even at a large areal mass loading of $\sim$12 mg sulfur/cm$^2$, approximately 6–12 times higher than that of most reports. The observed high areal specific capacity of the electrode (4.53 mAh cm$^{-2}$ after 300 cycles) and slow decay rate at 0.1C (0.08% per cycle after 300 cycles) represents a significant step forward for the application of Li–S batteries.

**Methods**

**Synthesis of GO**. The graphene oxide was synthesized from natural flake graphite by Hummers’ method. The concentration of the GO suspension obtained was $\sim$3.5 mg ml$^{-1}$, which was determined by drying 20 mL the suspension at 70 °C under vacuum for 72 h and then weighing the dried GO.

**Synthesis of GO Sponges (GS)**. The GO Sponges was prepared by hydrothermal reduction-assembly of homogeneous GO suspension. Briefly, 10 mL of the GO suspension was sealed in a 15-mL Teflon-lined stainless steel autoclave and maintained at 180 °C for 18 h. After been cooled to room temperature, black GS hydrogel can be obtained. For GO sponges preparation, the as-prepared graphene hydrogel was freeze-dried to remove absorbed water.

**Synthesis of S-GS**. The GO sponge was cut and shaped into a circular disc with a diameter of 10 mm (weight of 2.34 mg). Subsequently, appropriate amount of pure sulfur was evenly put on the GO sponge disc and then the sample were put in quartz tubes that were sealed under vacuum. The sulfur impregnation was further carried out by heating the sample in the vacuum-sealed quartz tube under 155 °C for 10 h. the weight of the prepared S-GS was 11.68 mg, corresponding the sulfur content in the total cathode is 80% and the areal mass loading of sulfur is $\sim$11.90 mg cm$^{-2}$.

**Materials characterization**. The composites were characterized by X-ray diffraction (XRD) with Cu-Kα irradiation, Fourier transform infrared spectra (FT-IR), and four-probe resistivity measurement system. And field emission scanning electron microscope (FESEM) was applied to observe the morphology of the synthesized composite material. Total sulfur loading ($\sim$9.34 mg) in the final electrode sample was calculated by weighing the sample before and after sulfur infusion (using a Mettler Toledo MS105DU Semi Micro Balance, 0.01 mg readability and ±0.02 mg repeatability).

![Figure 4](image-url) **Figure 4** | The charge–discharge voltage profiles of the 11th cycle for the S-GS cathode measured during galvanostatic cycled at 0.1C.

![Figure 5](image-url) **Figure 5** | (a) Charge-discharge profiles at different cycle numbers as labeled; (b) Cyclic performance and coulombic efficiency of the S-GS cathode for Li-S battery at a current density of 0.1C for 300 cycle.
Electrochemical measurement. The electrochemical experimental methods used in this work were similar to those in our previous study. The S-GS circular disc was used as the cathode directly. 2032 type coin cells were assembled in an argon-filled glove box with lithium foil as the anode. The separator was purchased from Celgard (model 2400). The electrolyte was 0.1 M lithium nitrate and 1.0 M lithium bis-trifluoromethane sulfonylimide in 1, 3-dioxolane and 1, 2-dimethoxyethane (volume ratio 1:1) (Zhangjiagang Guotai-Huarrong New Chemical Materials Co., Ltd). Galvanostatic measurements were conducted using a LAND CT2001A battery test system between 1.5 V and 3.0 V (vs Li/Li+).