Conductive graphene oxide-polyacrylic acid (GOPAA) binder for lithium-sulfur battery

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A B S T R A C T

Lithium-sulfur batteries have high cathode theoretical energy density, but the poor conductivity of sulfur and polysulfide shuttling result in serious polarization and low sulfur utilization. Moreover, the addition of insulating binder in the electrode increases the internal resistance, reducing specific capacity and rate-performance. Herein, we develop a composite binder with higher electronic conductivity, superior mechanical property and strong adsorption of polysulfides that imparts it some electrocatalytic activity. The reduced graphene oxide-polyacrylic acid (GOPAA) binder is prepared via a simple solution process. At constant loading fraction of 10 wt%, using GOPAA binder induces a 30% enhancement in the cathode capacity, better cycle life and rate capability compared to using PAA binder, reducing both the local charge-transfer resistance and the global electronic resistance before and after cycling. These are attributed to the enhanced binding strength and synergistic effect of reduced graphene oxide and PAA forming well-dispersed conductive bridges to promote rapid electron transfer. Additionally, GOPAA provides active sites for adsorption of lithium polysulfides and electrocatalytic activity, shifting redox peaks in cyclic voltammetry and improving roundtrip efficiency.

1. Introduction

Developing high-performance energy storage systems is of great importance for society and environment [1,2]. Among the various types of rechargeable batteries, lithium-sulfur (Li-S) batteries have attracted much attention due to their high cathode theoretical energy density (2600 Wh kg⁻¹), which is 3–5× that of commercialized cathodes, such as LiMn₂O₄, LiCoO₂, and LiFePO₄ [3,4]. However, there are two major problems that hinder the practical application of Li-S batteries: (1) The poor electronic conductivity of sulfur and its solid discharge products (Li₂S/Li₂S₂) leads to the low utilization of sulfur and reduces the practical cathode specific capacity to less than the theoretical capacity of 1675 mAh g⁻¹ [5,6,7]. (2) Soluble lithium polysulfide intermediates formed during the charge-discharge process can easily diffuse in the liquid electrolyte and cross over to the lithium anode, which causes rapid capacity fading [8–10]. Many efforts have been made to improve the electrochemical performance of Li-S batteries [11]. Commonly employed approach is to incorporate sulfur into nano-sized hosts such as carbon [12,13], polymer [14,15], and metal oxide [16] to form composite like KJC/S, a composite of sulfur and Ketjenblack carbon (KJC).

In electrodes, binder holds the active material and electron conductive agents together and intimately bonds them to the current collector [17]. Even though binder takes up only a small fraction of the dried electrode weight (usually 10 wt%), it plays a disproportionately important role, being the bridge that has to maintain connections. Conventional binders are usually electronically insulating polymers such as polyacrylic acid (PAA) [18], polyvinylidene fluoride (PVDF) [19], polyoxymethylene oxide (PEO) [20] and LA132 [21], which increase the internal resistance of the electrodes. So naturally there is desire to develop a new kind of binder with high electrical conductivity. Conductive polymers have been used as binders to prepare sulfur cathodes. Liu et al. [22] reported the conductive polymer poly(3,4-ethylenedioxythiophene) (PEDOT) as a conductive binder for Li-S batteries and demonstrated a increase in the electrochemical performance of sulfur cathodes. But it has been pointed out that most conducting polymers have unsatisfactory mechanical properties (adhesion strength), and limited conductivity [23,24], thus the balance between bonding strength and conductivity needs to be explored.

In addition to improving the obvious global electronic conductivity (K JC)


\[ R_{global}, \text{there is also another local electrocatalytic aspect of the binder that may need to be considered. Because binder bridges the sulfur-containing active material with electron donor, they are crucial for controlling the charge transfer resistance (R_{ct}), that characterizes thefacileness of local redox reactions whereby the active material receives an electron (thus may have to cross the “bridge” – the binder), and simultaneously captures a solvated Li}^+ \text{ from the liquid electrolyte, breaking its solvation shell and accomplishing redox of the sulfide in the process. These are molecular scale processes, so minute details of the binder surface (such as how well the liquid electrolyte wets it, and how well it adsorbs lithium sulfides) may matter. One can use electrochemical impedance spectroscopy (EIS) to measure } R_{global} \text{ and } R_{ct} \text{ of the electrode before and after cycling. Also, the electrocatalytic activity will show up in cyclic voltammetry (CV) tests.}

Graphene has the advantages of high specific area, superior electrical conductivity, and mechanical flexibility [25,26]. It is feasible to combine the flexible and conductive graphene with the insulating polymer binder to obtain a bi-function binder. Oh et al.[27] applied a highly conductive graphene as a nanofiller for PVDF binder and directly used the nanocomposite as a conductive binder for lithium titanate oxide (LTO) electrodes. However, the cycling performance of Li-S batteries based on PVDF binder is poorer than that of PAA binder [18]. PAA, a type of glue-like binder, has strong adhesion strength and can better restrain the electrode material desquamation due to large volume change and stress evolution in conversion electrodes.

In this work, we develop a conductive binder which consists of reduced graphene oxide (RGO) and PAA, termed GOPAA. The amount of RGO in GOPAA is really small, only 1/10th of the GOPAA weight. As GOPAA occupies 10 wt% of the cathode, the total amount of RGO in the cathode is only 1 wt%. But huge improvements in performance and cycle life are seen compared to using PAA binder alone at 10 wt%, indicating the disproportional importance of the binder.

2. Experimental section

2.1. Synthesis of reduced graphene oxide, GOPAA binder and KJC/S composite

Graphene oxide (GO) was synthesized by a modified Hummers’ method, which used natural flake graphite as the precursor [28]. Then, the obtained graphene oxide was reduced using the saturated vapor of dimethylhydrazine and further reduced by annealing in nitrogen for 5 h at 200 °C [29]. Finally, the material was cooled to room temperature and reduced graphene oxide (RGO) was obtained.

To prepare the GOPAA binder, 4 mg RGO was added into 3.2 mL N-methyl pyrrolidinone (NMP) and sonicated for 2 h to form a uniform dispersion. Then, 36 mg PAA was incorporated into the above solution with vigorous stirring under 70 °C for 24 h in a sealed system. The obtained composite conductive binder was named GOPAA.

To synthesize the KJC/S composite, sulfur and Ketjenblack carbon (KJC) were mixed at a mass ratio of 7:3. Then, the mixture was heated above solution with vigorous stirring under 70 °C for 24 h in a sealed filling glove box using the sulfur electrode as working electrode, lithium metal foil as the counter electrode and porous polypropylene (PP) membrane as the separator. The electrolyte was the mixing of 1, 3-dioxolane (DOL) and 1, 2-dimethoxyethane (DME) (volume ratio 1:1) with 1 M LiTFSI and 0.1 M LiNO\(_3\). The amount of the electrolyte used in the cells was ~20 μL. The galvanostatic charge-discharge tests were measured on a CT2001A cell test instrument (LAND Electronic Co.) between 1.7 and 3.0 V (vs. Li/Li\(^+\)). The cyclic voltammetry (CV) was conducted on a CHI660D electrochemical workstation (Shanghai Chenhua, China). Electrochemical impedance measurements (EIS) were conducted in the frequency range of 10\(^{-2}\)–10\(^5\) Hz with the amplitude of 5 mV.

2.4. Computational modeling

Theoretical calculations of lithium sulfides adsorption were carried out by using Vienna ab initio simulation package (VASP) [30], which was based on density functional theory (DFT) [31] and the projected augmented wave (PAW) method [32]. Generalized gradient approximation (GGA) in the form of Perdew-Burke-Ernzerhof (PBE) [33] was chosen as exchange correlation potential. The kinetic energy cutoff was set to 500 eV. All structures were fully optimized and the maximum force on each atom was less than 0.01 eV Å\(^{-1}\). Various initial adsorption configurations have been considered, and the most stable optimized configuration is presented in Fig. 1. The adsorption energy (\(E_a\)) is defined as the difference between the total energy (\(E_{\text{tot}}\)) of Li\(_2\)S (Li\(_2\)S\(^-\))–PAA compound, and the energy sum of Li\(_2\)S (Li\(_2\)S\(^-\)) and PAA (\(E_{\text{PAA}}\)): \(E_a = E_{\text{Li}_2\text{S/Li}_2\text{S}^-}\text{PAA} - E_{\text{Li}_2\text{S}^-}\text{PAA} - E_{\text{PAA}}\).

3. Results and discussion

PAA can chemically adsorb lithium polysulfides by the Li–O bond (Fig. 1a, c). The lengths of the Li-O bond formed with Li\(_2\)S and Li\(_2\)S\(^*\) were calculated to be 1.84 and 1.93 Å, respectively. The coterminous oxygen in PAA also has an interaction with lithium polysulfides (the dotted line in Fig. 1a, c), thereby further increasing the adsorption strength. The adsorption energies of Li\(_2\)S and Li\(_2\)S\(^*\) on PAA were 1.44 and 1.31 eV, respectively. PAA has a higher binding energy with lithium polysulfides compared with PVDF (Table S1). Therefore, the cycling performance of Li-S batteries based on PAA is better than that based on PVDF [10,18]. To confirm the bond formation, the differential charge densities around Li\(_2\)S and Li\(_2\)S\(^*\) were calculated from the difference of Li\(_2\)S (Li\(_2\)S\(^-\))–PAA compound charge density and the sum of Li\(_2\)S (Li\(_2\)S\(^-\)) charge density and PAA charge density (Fig. 1b, d). The blue color represents the charge density decrease, while the green-yellow and orange colors correspond to charge density increase. The
Fig. 1. Schematic structures of (a) Li$_2$S and (c) Li-S* adsorbed on PAA. The differential charge densities around (b) Li$_2$S and (d) LiS*. (e) Schematic illustration for the formation of hydrogen bonding between RGO and PAA. (f) Fourier transform infrared spectroscopy (FT-IR) spectra of PAA, RGO and GOPAA.

Fig. 2. Scanning electron microscope (SEM) images of (a, c) KJC/S PAA electrodes and (b, d) KJC/S GOPAA electrodes.
result shows the decrease in the electron density between Li and S atoms and the increase between Li and O atoms, implying the Li atoms were strongly bound to O atoms.

GOPAA was prepared by a simple solution process of mixing PAA with RGO. The free carboxylic acid in PAA and the O-H, −COOH and C-O-C sites in RGO can form hydrogen bonds (Fig. 1e). In the FT-IR spectrum of PAA (Fig. 1f), two absorption bands at 1730 and 1447 cm\(^{-1}\) correspond to the stretching vibrations of C=O and O-H from −COOH in PAA[34]. In the FT-IR spectrum of RGO, the bands at 3444, 1596 and 1067 cm\(^{-1}\) are attributed to the O-H, −COOH and C-O-C stretching vibrations, respectively[35,36]. In the FT-IR spectra of PAA and GOPAA, the bands from 3700 to 2800 cm\(^{-1}\) correspond to the hydrogen bond. The band around 3200–3500 cm\(^{-1}\) in GOPAA shifts to a lower wavenumber compared to PAA, which result from the dissociation of the hydrogen bonding among −COOH groups in PAA[37,38]. These results demonstrate that hydrogen bonding exists between PAA and RGO, which contributes to the homogeneous dispersion of RGO. The as-prepared RGO shows a laminar wrinkle structure (Fig. S2a) and there is no obvious morphology change in GOPAA (Fig. S2B), thus the high conductive property of RGO should be maintained. The uniformly distributed RGO not only act as electron transport channels but also provide large active sites for the reversible conversion between S\(_8\)/S\(_8\)^2 and Li\(_2\)S/Li\(_2\)S.

In fabricating the electrodes, the weight ratio of KJC/S, acetylene black and binder was 85: 5: 10, with nearly 60 wt% sulfur in the electrode (Fig. S3b). The average adhesion strength of PAA and GOPAA electrode measured from peel test is 111.36 and 158.55 mN mm\(^{-1}\), respectively. The enhancement in peel strength of GOPAA electrode
results from the better-dispersed binder in the cathode, despite the slightly lower mass loading of PAA (9 wt% in GOPAA electrode vs 10 wt% in PAA electrode). The morphology of KJC/S electrodes with different binders is shown in Fig. 2. The KJC/S PAA electrodes clearly exhibit a compact structure with the aggregation of small particles (Fig. 2a, c). The KJC/S GOPAA electrodes, in contrast, show abundant pores with RGO nanosheets (red circle in Fig. 2b), favorable for liquid electrolyte infiltration and rapid ion diffusion. Many pores with diameter of ~200 nm are distributed on the surface of the GOPAA electrodes (red arrow in Fig. 2d) and the RGO nanosheets act as bridges penetrating the active materials, facilitating electron transport and electrocatalysis.

The typical CV curves of GOPAA and PAA electrodes at a scan rate of 0.1 mV s\(^{-1}\) are shown in Fig. 3a and b, respectively. For the PAA electrode, only one reduction peak can be observed at 2.21 V in the first cycle, indicating a serious polarization. After the first cycle, two reduction peaks appear at 2.24 and 1.95 V, corresponding to the transformation of elemental sulfur (S\(_8\)) to soluble lithium polysulfides (Li\(_2\)S\(_n\), 4\(\leq\)n\(\leq\)8), and then to insoluble Li\(_2\)S\(_2\)/Li\(_2\)S, respectively. The oxidation peak at ~2.39 V corresponds to the conversion of Li\(_2\)S\(_2\)/Li\(_2\)S to the long-chain lithium polysulfide (Li\(_2\)S\(_6\)) [39]. The GOPAA electrode exhibits two reduction peaks at ~2.29 and 2.04 V and two oxidation peaks at ~2.36 and 2.31 V, which shows significantly better roundtrip efficiency than PAA electrode. Importantly, the position and shape of these peaks obtained from different cycles (including the first cycle) are almost identical, demonstrating that the introduction of GOPAA conductive binder has electrocatalytic effects and improves the electrochemical reversibility.

The charge-discharge profiles of PAA and GOPAA electrodes at various rates are shown in Fig. 3c and d. The GOPAA electrodes still maintained the ~2.0 V lower discharge plateau at high rates, such as 2 C. Moreover, the voltage difference \(\Delta U\) between charge and discharge plateaus of GOPAA electrodes was significantly smaller than that of PAA electrodes at the increased rates, indicating lower polarization and better roundtrip energy efficiency. At a current density of 0.2 C (1 C = 1675 mA g\(^{-1}\)), the discharge specific capacity of GOPAA electrodes reached 1000 mAh g\(^{-1}\) (Fig. 3e, specific capacities are calculated based on the mass of sulfur). Even at a high current density of 3 C, the GOPAA electrode showed a specific capacity of 464 mAh g\(^{-1}\), demonstrating excellent rate performance. The GOPAA electrodes exhibited a 30% higher specific capacity than PAA electrodes at 0.5 C (Fig. 3f). The initial specific capacity of GOPAA electrode was ~820 mAh g\(^{-1}\). Note the capacity gradually increased from the second to the fifth cycle, which can be attributed to the more porous structure of the GOPAA electrode that requires the activation steps to increase the utilization of active materials. After 100 cycles, the specific capacity of GOPAA electrode still remained at 635 mAh g\(^{-1}\). For PAA electrode, the initial discharge specific capacity was about 607 mAh g\(^{-1}\) and decreased to 448 mAh g\(^{-1}\) after 100 cycles. The capacity decay rate of GOPAA (0.22%/cycle) was lower than that of PAA (0.26%/cycle).

To further clarify the mechanisms of improved battery performance, EIS tests of the PAA and GOPAA electrodes were conducted before and after 100 cycles. The EIS spectra of the PAA and GOPAA electrodes at fresh state are compared in Fig. 4a. The Nyquist plots of the two electrodes consist of a depressed semicircle at high frequency region and an oblique line at medium frequency region. The diameter of the depressed semicircle represents the charge transfer resistance (\(R_{ct}\)). The GOPAA electrode shows a much lower \(R_{ct}\) than that of the PAA electrode. \(R_{ct}\) characterizes the local redox reactions whereby the sulfur-containing active material receives an electron from conductive agents (RGO and carbon black), and simultaneously captures a lithium ion from the liquid electrolyte, breaking its solvation shell and
accomplish redox of the sulfides in the process. \( R_{ct} \) is different conceptually from \( R_{global} \), the global (long-ranged) percolative transport of electrons, but \( R_{global} \) is also better in the GOPAA electrode as indicated by the smaller interelectrode distance due to the cathodes after 100 cycles, the Nyquist plots exhibit two depressed semicircles and a sloping line (Fig. 4b). The GOPAA electrode not only exhibits much lower \( R_{ct} \) (local), but also much lower \( R_{global} \) compared with the PAA electrode after cycling. This may be ascribed to the following reasons: (a) GOPAA is a better binder mechanically, as indicated by the stronger adhesion in peel test, bringing the active material closer to the conductive agents. RGO is also an active substrate for the reversible electrons, but during cycling, there is large volume change in the sulfur-containing active solid, which can induce self-stress and disrupt the global electron transport network. But due to its better mechanical properties, the GOPAA binder provides a more effective electronic conductive network and a more stable interface structure than using PAA. The characteristics of two different binder electrodes for Li\(S\) batteries are schematically shown in Fig. 4c and d. Compared with the PAA electrodes, the GOPAA electrodes increase the effective electrical contact area between the electrode materials, the current collector and the liquid electrolyte, as well as facilitating a stable long-range conductive network.

4. Conclusion

In summary, even though the binder takes up only a small fraction of the dried electrode weight (usually 10 wt%), it can play a disproportionally important role in battery performance, being the bridge that has to maintain connections between the active content, the electron source and ideally access to the liquid electrolyte. Because of its bridging role, it is also in an ideal position to mediate electrocatalysis involving multiple intermediate states of the lithium sulfides if it is conductive. So minute details of the binder surface (such as how well the liquid electrolyte wets it, and how well it adsorbs lithium sulfides) may matter. A novel conductive binder GOPAA was successfully prepared for sulfur cathodes, where we demonstrated ample hydrogen bonding between RGO and PAA, excellent dispersion, and maintenance of the conductivity and area of RGO. The GOPAA sulfur electrodes showed higher initial specific capacity, better rate performance, lower capacity decay rate and more favorable electrocatalytic kinetics. Inspired by these results, it might also be possible to disperse RGO by other chemical interactions, such as in-situ polymerization and covalent grafting, to obtain a multifunctional binder that gives high conductivity, superior peel strength, and low charge transfer resistance electrodes for lithium-ion batteries, sodium-ion batteries, lithium-air batteries, and lithium-sulfur batteries.

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Appendix A. Supplementary material

Supplementary data associated with this article can be found in the online version at [http://dx.doi.org/10.1016/j.nanoen.2016.12.002](http://dx.doi.org/10.1016/j.nanoen.2016.12.002).
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