Interface enhanced well-dispersed Co₉S₈ nanocrystals as an efficient polysulfide host in lithium–sulfur batteries

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The high specific capacity and energy density of lithium–sulfur batteries have attracted strong considerations on their fundamental mechanism and energy applications. However, polysulfide shuttle is still the key issue that impedes the development of Li–S batteries. Exploring nanocrystal hosts for polysulfide immobilization and conversion is a promising way. In this contribution, we have investigated well-dispersed Co₉S₈ nanocrystals grown on graphene oxide (GO) nanosheets with different degrees of dispersion as cathode host materials for Li–S batteries. The Co₉S₈–GO composite with 1 wt% GO (GCS1) has an average crystal size of 76 nm and shows the strongest adsorption capability toward lithium polysulfides. When used as the host material for the cathode of Li–S batteries, the GCS1–sulfur composite exhibits an initial specific capacity of ~1000 mAh g⁻¹ at 0.5 C and shows an average decay rate of 0.11% for 500 cycles. This work on the dispersion control of Co₉S₈ nanocrystals may inspire more investigations on well-dispersed nanocrystal based hosts for Li–S batteries.

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1. Introduction

Lithium-ion batteries (LIBs) have been a mature energy storage technique for electrochemical energy storage after more than 30 years’ research and development (R&D). However, its relatively low specific capacity and energy density have caused range anxiety in electric vehicles [1,2]. Lithium-sulfur (Li–S) batteries, which have a theoretical specific capacity of 1675 mAh g⁻¹, are considered to be one of the most promising energy storage systems for high-energy-consumption equipments [3–6]. For example, through the utilization of lithium-sulfur batteries as the power source, the goal of 500-mile range of electric cars in a single charge is highly expected. However, there are many bottlenecks blocking the practical use of Li–S batteries [7–13]. On the cathode part, there exist three main classic problems. The first issue is the polysulfide shuttling process, during which soluble polysulfides shuttle to the lithium anode and react with it and then shuttle back to the cathode. The second aspect is the low conductivity of sulfur and its solid product after lithiation. The third one is the volume changes of sulfur during lithiation and delithiation.

Many solutions have been put forward to investigate the fundamental mechanism and electrochemical performance optimization approaches of Li–S batteries. Various sulfur host materials have been proposed, mainly in two categories, non-polar and polar ones. The former is generally the diversified nanostructured carbon materials, such as mesoporous carbon [14,15], graphene [16,17], carbon nanotubes [18,19] and so on, which have weak binding energies with lithium polysulfides. The latter are mainly doped carbon materials [20,21], conductive oxygen-deficient titanium oxides [22–25], metallic cobalt sulfides [26–29], catalytic MnO₂ [30–32], bifunctional perovskites [33], metal nitrides [34–36], heterostructures [37–39] and so on [40,41]. Based on the previous researches, many properties of the host materials influencing the performance of Li–S batteries are summarized [42], such as the adsorption

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capability of polysulfides, electronic conductivity, surface area, porosity volume and distribution, catalytic conversion of lithium polysulfides/sulfides, etc. However, little attention has been paid to the influence of dispersion of supported nanocrystals on the electrochemical performances of Li-S batteries. Recently, the “polysulfide catalysis” concept is gaining more and more interest in Li-S batteries. In fact, polysulfide conversion is a process where nanocatalysts are needed as mediators [9,43]. Dispersion is a concept of much importance in heterogeneous catalysis because only exposed surface atoms can contribute to the catalytic reactions [44]. In heterogeneous catalysis, dispersion often comes together with particle sizes and increases with the decrease of particle sizes. Previous studies showed that metal catalysts such as Pt-graphene can effectively reduce the overpotential of the polysulfide reduction [45]. So it is of much significance to investigate nanocrystal based host materials with a proper degree of dispersion in Li-S batteries.

Nanostructured conductive metal sulfide (such as Co$_9$S$_8$) catalysts can effectively adsorb polysulfides and accelerate the conversion between soluble lithium polysulfides and solid lithium sulfides [27,28,46,47]. Herein, we investigated well-dispersed Co$_9$S$_8$ nanocrystals grown on graphene oxide nanosheets with different degrees of dispersion as cathode host materials for Li-S batteries. With graphene oxides (GO) incorporated in the composites increasing its contents, Co$_9$S$_8$ nanocrystals exhibited higher dispersity and smaller sizes. Fig. 1 illustrates the composite and its efficient function in regulating polysulfide adsorption and conversion in Li-S batteries. Compared to bulk Co$_9$S$_8$ with the same weight, the Co$_9$S$_8$-GO composite has more individual Co$_9$S$_8$ nanocrystals and shows more active sites toward lithium polysulfides. However, when used as polysulfide host materials in Li-S batteries, well-dispersed Co$_9$S$_8$ nanocrystals with a proper content of graphene oxides delivered the best electrochemical performances. The Co$_9$S$_8$-GO composite with 1 wt% GO (GCS1) and an average crystal size of 76 nm showed the best cycling stability in Li-S batteries among the series.

2. Results and discussion

The powder X-ray diffraction (XRD) patterns (Fig. 2(a)) of the as-prepared Co$_9$S$_8$ and Co$_9$S$_8$-GO composites with different amounts of GO confirm that Co$_9$S$_8$ exists in the composites with the cobalt pentlandite crystallographic phase which can be indexed to cubic Co$_9$S$_8$ (JCPDS Card No. 03-065-1765, space group Fm-3m). The two most intensive peaks at 29.8° and 52.1° correspond to the diffractions of (311) and (440) planes respectively. No obvious peak can be observed in the patterns around the 26° because the low content of graphene and relatively low diffraction intensity of graphene. Based on the weight ratios of GO to Co(NO$_3$)$_2$·6H$_2$O when synthesizing the composites, they were named GCS0 (0 wt% GO), GCS1 (1 wt% GO) and GCS4 (4.1 wt% GO). In order to compare the nanocrystal sizes of GCS1 and GCS4, we used Scherrer formula to calculate the average crystallite size by selecting the (311) facet diffraction peak as a representative. As shown in Table S1, on average, the crystallite size of GCS1 and GCS4 are 38 and 33 nm, respectively. This indicates that graphene oxide can promote the nucleation of Co$_9$S$_8$ nanocrystals and reduce its average sizes.

The surface area and pore distribution of these materials were measured by N$_2$ adsorption/desorption isotherm analysis. The

![Fig. 1. Illustration of the Co$_9$S$_8$-GO composite in contributing to the Li-S electrochemical cycling. Dispersed on graphene oxide nanosheet, Co$_9$S$_8$ nanocrystals can efficiently catalyze the redox reaction in Li-S batteries.](image)

![Fig. 2. (a) PXRD patterns of GCS0, GCS1 and GCS4 and the indexed PDF card. (b) N$_2$ adsorption/desorption isotherms of GCS0, GCS1 and GCS4, inset is the pore distribution curve of them. (c) Raman spectra of GCS0, GCS1, GCS4 and GCS10. TEM images of (d) GCS0, (e) GCS1 and (h) GCS4 and corresponding size distribution diagrams of (f) GCS1 and (g) GCS4.](image)
curves for GCS0, GCS1 and GCS4 and the corresponding pore distribution curves (inset) are shown in Fig. 2(b). GCS1 and GCS4 have a type IV physisorption isotherm, while it is type II for GCS0. It means that both GCS1 and GCS4 have mesopores while GCS0 has macropores. The hysteresis loops of GCS0, GCS1 and GCS4 are H3 type, which indicates that there exist slit type pores in these materials. The surface area of GCS0, GCS1 and GCS4 are 8.5, 8.5 and 16.5 m² g⁻¹, respectively (Table S1). Elemental analysis was conducted to examine the carbon and nitrogen content of GCS1 and GCS4 and the data are shown in Table S2. The carbon and nitrogen content of GCS1 is 2.8 wt% and 1.32 wt% with a C/N ratio of 2.1. For GCS4, the carbon content is 7.9 wt%, nitrogen content is 1.4 wt% and the C/N ratio is 5.6. The nitrogen should be from the solvent dimethylformamide which doped graphene with nitrogen atoms. The Raman spectra of GCS0, GCS1 and GCS4 are shown in Fig. 2(c). In the range of 100–1000 cm⁻¹, the most intense peaks are located at around 200, 486 and 696 cm⁻¹, which should be ascribed to the existence of Co₃S₄ [48,49]. In the range of 1000–2000 cm⁻¹, no D or G peaks related to the existence of graphene is observed. However, GCS10 exhibited a very different Raman spectrum compared to them. No intense characteristic signals of Co₃S₄ are found in GCS10 while D and G peaks are obvious (Fig. 2(c), purple). This indicates that the content of graphene is so high that GCS10 is more like a carbon material rather than Co₃S₄-containing material.

The transmission electron microscopy (TEM) images confirm that ultrafine nanocrystals of Co₃S₄ are well dispersed on graphene nanosheets. Fig. 2(d) reveals that GCS0 is made up of bulk Co₃S₄ solids. The size of every single Co₃S₄ particle is about one micrometer. Fig. 2(e) and (h) exhibits the morphology of GCS1 and GCS4, respectively. The Co₃S₄ nanocrystals with different sizes disperse and anchor on the wrinkled graphene nanosheets in GCS1 and GCS4. Fig. 2(f) and (g) is the corresponding nanocrystal size distribution diagrams. The size of nanocrystals of GCS1 is about 73 nm while it decreases to 26 nm in GCS4. The content of GO plays a vital role in the morphology of resulting samples. GO here has mainly two functions in the synthesis procedure. Firstly, GO can provide a platform for Co₃S₄ nanocrystals to nucleate and grow on the carbon nanosheet with oxygen-containing functional groups and defects. Secondly, the oxygen-containing functional groups on the carbon nanosheets of GO make it a good anionic surfactant, which can avoid the aggregation of Co₃S₄ nanocrystals. Without the existence of graphene oxide, Co₃S₄ grows freely in the dimethyl formamide (DMF) solvent and results in a nanosheets-constructed flower-like nanostructure (Fig. S1(a)).

Conductivity is crucial to the electron transfer process during the electrochemical process in Li–S batteries. Symmetrical cells comprised of GCCx (x = 0 or 1 or 4, the same below) electrodes and the Li₂S₈ electrolyte was constructed. Through electrochemical impedance spectroscopy (EIS) analysis, the cell with the GCS0 electrode showed the lowest impedance of about 4 kΩ. For cells with GCS1 and GCS4, it was around 5 kΩ, but the former was a little smaller than the latter (Fig. 3(a)). This result showed that the addition of graphene oxide decreased the overall electron conductivity of the composite materials. Moreover, the catalytic capability toward polysulfides reduction is also a vital parameter that decides the specific capacity of Li–S batteries. Cyclic voltammetry (CV) was used to differentiate the capability of GCS0, GCS1 and GCS4 in catalyzing polysulfides reduction. As shown in Fig. 3(b), symmetrical cells with GCS0, GCS1 and GCS4 show similar hysteresis loop-like curves. However, at the highest overpotential, GCS1 shows the highest current density and GCS4 exhibits the lowest. It indicates that GCS1 has the largest capability in catalyzing the conversion of polysulfides, which is also confirmed by the long-cycle performance of Li–S batteries afterwards. Polysulfide adsorption capability is also a very important parameter that determines the performance of a host material employed as the cathode material in Li–S batteries. The visualization experiment was conducted to compare the capability of polysulfides adsorption of GCS0, GCS1 and GCS4. Fig. 3(e) shows that the Li₂S₈ in DOL/DME solvent is a dark red solution. With the addition of GCS0, GCS1 and GCS4 (the same weight) into Li₂S₈ solutions in different test tubes, after a rest for 24 h, the color of the supernatant became distinctly different. GCS1 interacted with polysulfides strongly and the supernatant of the test tube with it became very clear. However, GCS0 showed weak adsorption capability toward polysulfides and the color of the supernatant of the test tube with it still kept dark red, a little lighter than the original Li₂S₈ solution. GCS4’s adsorption capability was in between GCS1 and GCS0 and the color of its supernatant was light dark red. This experiment shows that GCS1 is the strongest in polysulfides adsorption, GCS4 is the second and GCS0 is the weakest.

X-ray photon spectroscopy (XPS) was used to detect the interactions between GCCx and lithium polysulfides (here is Li₂S₈). The mixtures were collected from the precipitates by centrifuging the GCCx-Li₂S₈ liquids. As shown in Fig. 3(c), the Co 2p spectrum exhibits two contributions from Co 2p₃/₂ and Co 2p₁/₂. For GCS0, the peaks at 780.88 and 797.28 eV should be assigned to the state of Co³⁺ and the peaks at 782.68 and 803.38 eV should be assigned to the state of Co⁴⁺[50]. In the case of the GCS0-Li₂S₈ mixture, peaks at 777.68 and 792.86 eV are due to the existence of Co³⁺ and peaks at 779.68 and 795.98 eV should be assigned to the state of Co⁴⁺. The XPS spectra of GCCS1-Li₂S₈ and GCS4-Li₂S₈ mixture are very similar and show characteristic peaks at nearly the same locations. For example, in the case of GCCS1-Li₂S₈ mixture, characteristic peaks of the state of Co³⁺ are located at 777.78 and 792.78 eV and Co²⁺ are at 779.78 and 795.78 eV. The peaks of Co 2p₃/₂ in GCS0 decreased about 3.1 eV compared to that in the mixture of GCCS1-Li₂S₈. However, there is ca. 0.1 eV change of the peaks between the GCCS0-Li₂S₈ and GCCS1-Li₂S₈ mixture. This result shows that the interactions between GCCx and polysulfides are very strong (up to 3.0 eV variation), which should be ascribed to the polar property of Co₃S₄. The similarities between the three GCCx-Li₂S₈ (x = 0, 1, 4) mixtures indicate that XPS is hard to differentiate the polysulfides adsorption capabilities of GCCx. Fig. 3(d) depicts the S 2p spectra of GCCS0 and GCCSx-Li₂S₈ mixtures. Following convention, only S 2p₃/₂ will be mentioned in the S 2p₃/₂/S 2p₁/₂ doublet. The spectrum of Li₂S₈ exhibit two 2p₃/₂ contributions at 162.88 and 163.48 eV, respectively. The former peak should be assigned to the state of bridging sulfur (S–S). The latter should be assigned to the terminal sulfur. There is a broad peak at about 165 eV and it can be ascribed to the partial oxidation of Li₂S₈ during transfer when testing. In the sample of the GCCS0-Li₂S₈ mixture, two 2p₁/₂ contributions are located at 160.38 and 161.78 eV, representing a ~2.5 and ~1.7 eV shift, respectively. GCCS1-Li₂S₈ and GCS4-Li₂S₈ mixtures show similar spectra with the GCCS0-Li₂S₈ mixture. No more than 0.2 eV shift exists in the peaks. This result indicates that the existence of Co₃S₄ changed the chemical environment of sulfur greatly. Also, like the Co 2p spectrum, the interactions of GCCx and polysulfides are hard to be distinguished from the S 2p spectrum. The Li 1s spectrums show that there is ca. 1 eV shift from Li₂S₈ to the GCCSx-Li₂S₈ sample (Fig. S2). It corroborates the strong interactions between GCCx and polysulfides. In order to uncover the strong interactions between Co₃S₄ and lithium polysulfides, we employed density functional theory (DFT) calculations to investigate the intrinsic origins. The Co₃S₄ (440) lattice plane with atomic composition ratio of Co:S = 1:1 on exposed surfaces is taken as the representative because it exhibits the strongest diffraction peak in the XRD pattern. Fig. 4(a) depicts the optimized structure of Co₃S₄. The adsorption energies of Li₂S₈, Li₂S₈ on the Co₃S₄(440) plane are −3.67, −4.04 eV (Fig. 4(b) and (c)), respectively.
Batteries with solid sulfur cathodes were tested through CV and electrochemical cycling to compare the effects of GCS0, GCS1 and GCS4 on the electrochemical performances of Li-S batteries. The active sulfur cathode material was made by melt-diffusing sulfur in the GCSx material at 155 °C. The powder XRD patterns are shown in Fig. S3. Characteristic peaks originated from sulfur are obvious while peaks from Co₉S₈ have disappeared. Thermogravimetric analysis was used to test the sulfur content in GCSx-S composites. As shown in Fig. S4, the sulfur content in GCS0-S, GCS1-S and GCS4-S composites are 71 wt%, 72 wt% and 72 wt%, respectively. The CV curves of GCS0, GCS1 and GCS4 were obtained through scanned at 0.1 mV s⁻¹ in the potential window of 1.7–2.8 V.
There are two cathodic peaks and one anodic peak for every kind of material. Following the method widely used in electrocatalysis, the onset potential for every peak was taken at a current density of 10 μA cm$^{-2}$ beyond the baseline current where the change of current is the smallest, i.e. dI/dV=0. For GCS1, the onset potential for the first cathodic peak is 2.366 V which is 27 mV lower than GCS0 but 2 mV higher than GCS4. It is 2.070 V for the second onset potential of GCS1, 28 mV lower than GCS0 and 137 mV higher than GCS4. For the anodic peaks, they are 2.157, 2.189 and 2.191 V for GCS0, GCS1 and GCS4, respectively. It indicates that GCS0 has the largest onset potential for the cathodic process and the lowest onset potential for the anodic process. It should be ascribed the highest electronic conductivity of GCS0. Moreover, the peak voltages were also investigated. They are 2.293, 2.251 and 2.241 V for the first cathodic peak of GCS0, GCS1 and GCS4, respectively. For the second cathodic peak, they are 2.030, 1.991 and 1.969 V for GCS0, GCS1 and GCS4, respectively. Because the anodic peak is a combined peak of several oxidation reactions of polysulfides, so it has no sense to compare the anodic peak voltages of different materials. Through analyzing the trend of change of cathodic peak voltages, we found that it had a similar tendency to the onset potential, which should also be ascribed to the intrinsic conductivity of GCS0, GCS1 and GCS4. Besides, the rate of lithium ion diffusion within the electrodes in Li–S batteries has...
an essential effect on the conversion process of sulfur redox. We took advantage of the classical Randles–Sevcik equation to calculate the lithium diffusion coefficient \( D_{Li} \) based on cathodic current peaks \( I_{1a} \) and \( I_{2a} \) and the highest anodic current peak \( I_p \) (Fig. 5(c) and Table S3) [51–53]. For the reduction process, \( D_{Li} \) coefficients of GC5 and GC4 are an order of magnitude higher than that of GC50. GC51 is a little higher than GC54. However, for the oxidation process, \( D_{Li} \) coefficients of GC50, GC51 and GC54 are at the same order of magnitude. GC51 still has the highest \( D_{Li} \) coefficient. This indicates that GC51 and GC54 showed a higher activity in catalyzing the reduction process of sulfur species compared to the oxidation process.

Typical galvanostatic discharge/charge voltage curves of GC50, GC51 and GC54 at 0.5 C were also investigated in the potential range of 1.7–2.8 V. Fig. 5(b) shows that there are usually two plateaus in the discharge process and one plateau in the charge process for every specific material. GC50 showed a very sluggish kinetic discharge process. For example, there is a deep ditch (black arrow) at the nexus of liquid-liquid single-phase reduction and liquid-solid two-phase reduction, which was analyzed by a former review [54]. If the host material could not adsorb the polysulfides efficiently, the kinetic rate became very low and the potential drop would become obvious too. It corroborates the low efficiency of GC50 in adsorbing polysulfides because at this point polysulfides solution has the largest viscosity. It is in highly accordance with the former visualization test. As expected, GC51 has the highest kinetic rate in the discharge process. What is more, the voltage hysteresis between the discharge and charge process for GC51 is about 250 mV and is the lowest amongst the three materials. For practical use of Li–S batteries, we conducted a long-term cycling test of the Li–S batteries enabled by GC50, GC51 and GC54 (Fig. 5(d)). After activation for several cycles, GC51 showed an initial specific capacity of ~1000 mAh g\(^{-1}\) and kept a retention of ~58% after 200 cycles and of ~46% after 500 cycles. However, GC50 showed its initial specific capacity of ~800 mAh g\(^{-1}\) and a retention of ~57% after 200 cycles. For GC54, its first specific capacity was about 930 mAh g\(^{-1}\) and it retained ~50% after 200 cycles. The capacity fading rate for GC50, GC51 and GC54 in the 300 cycles are 0.16%, 0.14% and 0.18%, respectively. For GC51, its capacity fading rate in the 500 cycles is 0.11%.

Based on the above results, we can conclude that GC51 has the strongest capability of adsorption toward polysulfides. GC51 has more active sites (C\(_0\)S\(_2\) nanocrystals) than bulk C\(_0\)S\(_2\), which guarantees higher adsorption toward soluble lithium polysulfides (Li\(_2\)S\(_n\), 4 ≤ n ≤ 8) and more efficient conversion of them to solid Li\(_2\)S\(_2\) and Li\(_2\)S (Fig. 5(e)). Even though GC54 possesses smaller C\(_0\)S\(_2\) nanocrystals, the interfaces of C\(_0\)S\(_2\) and graphene oxide should be less than GC51 because it has a higher content of bare GO, which is not the best ratio in the composite. That is why GC51 can adsorb more lithium polysulfides than GC54. Moreover, although its conductivity is not the highest, it still exhibits the best electrochemical performance when employed as the cathode host material in Li–S batteries. Higher conductivity will contribute to higher onset potentials in the cathodic process and lower onset potentials in the anodic potentials. All in all, polysulfide adsorption is the most important parameter that determines the rate of discharge/charge reactions, specific capacity and cycling stability of Li–S batteries.

3. Conclusions

In summary, we have investigated the influence of well-dispersed metal sulfides (C\(_0\)S\(_2\)) grown on graphene nanosheets with different degrees of dispersion (GC51 and GC54) as host materials on the electrochemical performances of Li–S batteries. The visual polysulfide adsorption test showed that GC51 had the strongest capability toward the adsorption of lithium polysulfides. The electronic conductivity, lithium ion diffusion rate and catalytic contribution of polysulfide conversion of GC50, GC51 and GC54 have been compared too. GC51 exhibits the best electrochemical performance when employed as the cathode host material in Li–S batteries. It should be ascribed to its largest capability in polysulfides adsorption and good catalytic contribution of polysulfides conversion. What is more, GC51 facilitates the lithium ion diffusion most among the three materials. This work may inspire more research on supported nanocatalyst catalysts with different degrees of dispersion for polysulfides conversion and its contribution in improving practical uses of Li–S batteries.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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Supplementary materials

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References
