Carbon–sulfur composites for Li–S batteries: status and prospects

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We review the development of carbon–sulfur composites and the application for Li–S batteries. Discussions are devoted to the synthesis approach of the various carbon–sulfur composites, the structural transformation of sulfur, the carbon–sulfur interaction and the impacts on electrochemical performances. Perspectives are summarized regarding the synthesis chemistry, electrochemistry and industrial production with particular emphasis on the structural optimization of carbon–sulfur composites.

Introduction

Carbon materials are commonly used electrode materials for their diversity, robust chemistry, conductivity, stability and abundance. Such materials, including activated carbon, nanoporous carbon, carbon nanotubes, graphene sheets and many other forms, have been widely used in supercapacitors,18 lithium ion batteries (anode materials),6,8–11 and lithium-air/ oxygen batteries (gas diffusion cathodes).14 Carbon materials have excellent conductivity and good mechanical resilience, and therefore have shown remarkable usefulness in facilitating the performance of insulating or semiconducting cathode materials (mainly lithium-transition metal oxides).10–13

A recent boom in carbon-related energy research has been triggered by the emerging focus on Li–S batteries.15 Li–S batteries utilize a lithium metal anode and a sulfur cathode. The multi-electron-transfer cathode reaction of \( S_{8} + 16 Li^{+} + 16 e^{-} \rightarrow 8 Li_{2}S \) offers an extremely high theoretical capacity of 1672 mA h g\(^{-1}\), while the lithium anode provides a theoretical capacity of 3842 mA h g\(^{-1}\). The average potential of a Li–S cell is 2.15 V with respect to Li\(^{+}/Li\), which is relatively low compared to graphite–LiMO\(_{2}\) batteries (>3 V). The specific energy of Li–S battery is still very large, however, the medium voltage is offset by the very high capacity and reaches a theoretical value of 2567 Wh kg\(^{-1}\). The use of lithium metal in a Li–S cell is far from

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optimal, as it is potentially risky due to dendrite formation.\textsuperscript{16} Many efforts have been devoted to lithium-metal free sulfur-based batteries.\textsuperscript{17–20} Unfortunately, the lithium metal anode and the lithium sulfide anode are both moisture sensitive. Large scale fabrication of battery anodes from these materials must be performed in a dry and inert environment, and this could potentially impact the industrial production of Li-S batteries.

The other obstacle that stands in the way of the large-scale uptake of Li-S technology is relevant to the sulfur cathode. Sulfur is a promising cathode material due to its advantages such as low cost, nontoxicity and virtually unlimited supply. However, several practical problems have delayed widespread application of sulfur. Sulfur itself is insulating ($5 \times 10^{-30}$ S cm$^{-1}$ at 25 $^\circ$C) and is thus unusable as an electrode material. This fact necessitates the use of carbon as a conducting additive in the sulfur cathode of Li-S batteries. The redox chemistry of sulfur in the cathode relies on a solid (cyclo-S$_8$)–liquid (chain-polysulfides (PS, S$_{4-n}^{2-n}$))–solid (Li$_2$S$_2$/LiI$_2$S) reaction with a systematic decrease in the ion chain length. The reduction of S$_8$ forms S$_2^{2-}$ at 2.39 V vs. Li$^0$/Li$^+$, which is successively reduced to S$_6^{2-}$ at 2.37 V vs. Li$^0$/Li$^+$ and then to S$_4^{2-}$ at 2.24 V vs. Li$^0$/Li$^+$.\textsuperscript{21} Because the PS ions are soluble, the reaction is fast (Fig. 1, first plateau). This process corresponds to a theoretical capacity of 418 mA h g$^{-1}$. Continuing discharge will reduce the PS ion to Li$_2$S$_2$/LiI$_2$S which are insoluble. Due to the much slower reaction kinetics at the second plateau and the tail, which correspond to solid state reactions, the theoretical value usually cannot be achieved and a lower capacity of 1256 mA h g$^{-1}$ is more realistic (corresponding to 1.5 e S$^{-1}$).

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A critical issue associated with the PS ions ($S_{4-8}^{2-}$) is their solubility in an electrolyte. The dissolved PS ions easily diffuse from the cathode to anode driven by a concentration gradient (Fig. 1, inset). In fact, even sulfur can be weakly dissolved in an aprotic electrolyte. Reduction of the dissolved PS is as yet unclear, but is possible that it occurs through (1) ion chain scission, (2) disproportionation, and (3) reduction by the Li anode. The insoluble Li$_2$S$_2$/Li$_2$S deposits on the Li anode can react with PS ions, yielding soluble medium-chain ions, which diffuse back to the sulfur cathode. The whole process is known as the “PS shuttle mechanism”. This shuttle phenomenon results in the following drawbacks: (1) the active mass loss from the cathode, (2) reduction of Coulombic efficiency, and (3) capacity decay upon cycling. Additionally the insoluble Li$_2$S$_2$/Li$_2$S can also deposit on the cathode and is insulating. This makes the cathode electrochemically inaccessible due to the hindered ion and electron transport. A further issue associated with the sulfur cathode is the volume variation. The density of sulfur is $2.03 \text{ g cm}^{-3}$ while that of Li$_2$S is $1.67 \text{ g cm}^{-3}$, which results in an 80% volume expansion. The current limitations of sulfur cathodes and the already proposed strategies to address these issues are summarized in Table 1. Many recent studies on carbon–sulfur composites are summarized in Table 2, where the synthesis method and the performance evaluation, typically the cyclic stability, are compared.

### Sulfur forms

Thermal annealing of sulfur with carbon is still the most general synthesis technique for the carbon–sulfur composite so far, and is also the best choice for industrial production. In this context, the thermal behaviour of sulfur is of interest. Elemental sulfur has a variety of crystalline and molecular/polymeric forms. The identification of these allotropic forms has been the subject of study over decades, and is summarized in Fig. 2. Ordinary sulfur ($\alpha$-S), i.e. the rhombic stacking of $S_{8}$, is the stable room-temperature form. It slowly converts to monoclinic sulfur ($\beta$-S) from 95.5 °C. Both rhombic and monoclinic sulfur are soluble in CS$_2$. The melting temperature of monoclinic sulfur is 118.7 °C and the boiling temperature of sulfur is 444.6 °C. Liquid sulfur is predominantly $S_8$ at the melting point, where cyclo-$S_8$ and catena-$S_8$ co-exist. $S_8$ rings undergo thermal scission to form linear sulfenyln diradicals at 159 °C. Between 159 °C and 444.6 °C, sulfur first polymerizes and then depolymerizes, accompanied with a viscosity change which reaches a maximum at 186 to 188 °C. $S_8$ is predominant at 444.6 °C, but is dissociated into short chain sulfur at higher temperatures ($S_6$, $S_4$: vapour at 600 to 800 °C, $S_2$: vapour above 850 °C, and $S$: vapour above 1800 °C).26

### Carbon–sulfur composites

#### Mesoporous carbon–sulfur composites

Wang and co-workers reported the first use of a worm-like mesoporous carbon as a host material for sulfur.27 In their approach, sucrose was chosen as the carbon source and sodium silicate as the silica template source. The mixture of sulfur and mesoporous carbon was heated at 200 °C for 6 h to let the sulfur melt diffuse into the mesopores of the carbon host, and then the temperature was raised to and held at 300 °C for another $3 \text{ h}$ to vaporize the superficial sulfur outside the carbon particles.27 An ionic liquid, 1-ethyl-3-methylimidazolium bis[(trifluoromethyl)sulfonyl]imide (EMITFSI), was used as the electrolyte solvent, in which lithium bis(trifluoromethanesulfonimidate (LiTFSI) was dissolved to form a 1 M LiTFSI–EMITFSI electrolyte. Their results showed much improved cycling stability of the mesoporous carbon–sulfur composite. Two reasons were proposed to explain the better performance: (1) the physical adsorption of polysulfides by the mesopores ($10-30 \text{ nm}$), and (2) the uniform distribution of sulfur in the mesopores, which assists with electron conduction by the composite cathode. A result noticed in Wang’s work was that the stability of the same material in an organic electrolyte was not as good as that in an ionic liquid electrolyte, and the cause was suggested to be the reduced solubility of PS in the ionic liquid. Meanwhile, the reduction potentials from sulfur to PS and to Li$_2$S$_2$/Li$_2$S were found to depend on the electrolyte, which may imply a PS solubility-related mechanism.

The manifested advantages of mesoporous carbon were later highlighted by Ji and co-workers.28 The major progress relied on the optimal use of the ordered mesoporous carbon (CMK-3) synthesized by using the SBA-15 silica template. This CMK-3 carbon possessed uniform and narrow mesopores (3 nm) and large pore volume ($2.1 \text{ cm}^3 \text{ g}^{-1}$) (Fig. 3a). The sulfur content in the CMK-3-sulfur composite can be as high as 70 wt%. This value is lower than the theoretical value (79 wt%) because it is good for buffering the volume change during lithiation and delithiation of sulfur confined in the mesopores.24 As a result of the uniform distribution of sulfur in the mesopores, the conductivity of the composite is comparable to that of CMK-3 within errors ($0.21 \text{ S cm}^{-1}$ vs. $0.2 \text{ S cm}^{-1}$, $\pm 0.02 \text{ S cm}^{-1}$). This gave rise to a very high Coulombic efficiency (99.94%) in the first discharge–charge cycle, which indicated an extremely low fraction of PS dissolved in the electrolyte. A typical discharge–charge profile of the CMK-3–S composite shows the well resolved plateaus corresponding to the redox chemistry of sulfur, PS and lithium sulfides (Fig. 3b). It is notable that an organic electrolyte was used in this study. Ji et al. also demonstrated the enhanced stability of a PEG-coated CMK-3–sulfur composite,24 introducing the concept of carbon–sulfur–polymer ternary composites for Li–S batteries.

Many other types of mesoporous carbon–sulfur composites have subsequently been developed and examined for their efficacy in promoting sulfur stability in Li–S batteries.29–34 Bimodal mesoporous carbon with 2.0 nm and 5.6 nm
Table 2  Characteristics of various carbon–sulfur composites

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<th>Classification</th>
<th>Characteristics</th>
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<th>Cycle performance (1C = 1675 mA g⁻¹)</th>
<th>Electrolyte/voltage window</th>
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<td>Sucrose-derived hydrothermal carbon nanosphere</td>
<td>Sulfur loaded in 0.7 nm micropores</td>
<td>Sulfur melt adsorption</td>
<td>42%</td>
<td>650 mA h g⁻¹/400/1000th cycle</td>
<td>1 M LiPF₆, PC-EC-DEC, 1.0–3.0 V</td>
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<tr>
<td>α-Glucose-derived carbon coating on CNT</td>
<td>Sulfur loaded in 0.6 nm micropores</td>
<td>Sulfur melt adsorption</td>
<td>40.2%</td>
<td>1142 mA h g⁻¹/0.1C/200th cycle</td>
<td>1 M LiPF₆ in EC/DC (1 : 1 in wt%), 1.0–3.0 V</td>
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<tr>
<td>Phenolic resin-derived porous carbon</td>
<td>Sulfur loaded in &lt;1 nm micropores</td>
<td>Sulfur melt adsorption + extraction process</td>
<td>16%</td>
<td>200 mA h g⁻¹/3000/800th cycle</td>
<td>1 M LiPF₆ in (EC/DMC/EMC, 1 : 1 : 1 vol.), 1.5–2.8 V</td>
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<td>Sucrose-derived carbon with sodium silicate as template</td>
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<td>500 mA h g⁻¹/50/40th cycle</td>
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<td>PPy-derived carbon with colloidal silica as template</td>
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<td>613 mA h g⁻¹/0.1C/50th cycle</td>
<td>1 M LiTFSI/DOL-DME (1 : 1), 1.0–3.0 V</td>
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<td>Ordered mesoporous carbon (CMK-3)</td>
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<td>1100 mA h g⁻¹/168/200th cycle</td>
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<td>Phenol formaldehyde-derived carbon with TEOS and F127 as templates</td>
<td>2.0 nm and 5.6 nm</td>
<td>Sulfur melt adsorption</td>
<td>40%</td>
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<td>1 M LiTFSI/DOL-DME (1 : 1), 1.5–3.0 V</td>
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<tr>
<td>Phenol formaldehyde-derived carbon with TEOS and F127 as templates</td>
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<td>50%</td>
<td>730 mA h g⁻¹/1C/100th cycle</td>
<td>1 M LiTFSI/DOL-DME (1 : 1), 1.5–3.0 V</td>
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<td>Hierarchical porous carbon</td>
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<tr>
<td>Sucrose-derived carbon with silica and copolymer latex as dual-template</td>
<td>Macropore size-300 nm</td>
<td>Sulfur melt adsorption</td>
<td>50%</td>
<td>884 mA h g⁻¹/0.1C/50th cycle</td>
<td>1 M LiTFSI/DOL-DME (1 : 1), 0.1 M LiNO₃, 1.0–3.0 V</td>
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<td>Hollow carbon sphere</td>
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<tr>
<td>Petroleum pitch with silica as template</td>
<td>Diameter 200 nm</td>
<td>Vapour phase infusion</td>
<td>70%</td>
<td>974 mA h g⁻¹/0.5C/100th cycle</td>
<td>1 M LiTFSI in tetraglyme, 1.7–3.1 V</td>
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<tr>
<td>Glucose derived carbon with SnO₂ sphere as template</td>
<td>Diameter 300 nm</td>
<td>400 °C heat treatment</td>
<td>64%</td>
<td>690 mA h g⁻¹/0.1C/100th cycle</td>
<td>1 M LiTFSI in tetraglyme, 1.5–3 V</td>
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<td>CNT/CNF</td>
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<tr>
<td>Carbon nanotube array</td>
<td>CNT diameter from 7–30 nm</td>
<td>Solvent evaporation</td>
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<td>900 mA h g⁻¹/13/40th cycle</td>
<td>1 M LiTFSI/DOL-DME (1 : 1), 0.25 M LiNO₃, 1.0–3.0 V</td>
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<tr>
<td>Disordered carbon nanotubes</td>
<td>CNT diameter 200 nm</td>
<td>High temperature (500 °C) treatment</td>
<td>40%</td>
<td>700 mA h g⁻¹/200/100th cycle</td>
<td>1 M LiTFSI in TEGDME, 1.5–3 V</td>
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<tr>
<td>Styrene derived carbon hollow fiber with AAO as template</td>
<td>Hollow fiber</td>
<td>Sulfur melt adsorption</td>
<td>75%</td>
<td>730 mA h g⁻¹/0.2C/150th cycle</td>
<td>1 M LiTFSI/DOL-DME (1 : 1), 0.1 M LiNO₃, 1.7–2.6 V</td>
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<td>Graphene</td>
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<td>PEG modified GO-carbon black</td>
<td>Emulsion wrap 1 μm sulfur particles</td>
<td>Emulsion: Na₂S₂O₃ + Triton-X100 + HCl</td>
<td>70% (before annealing)</td>
<td>520 mA h g⁻¹/0.2C/100th cycle</td>
<td>1 M LiTFSI/DOL-DME (1 : 1), 1.7–2.6 V</td>
</tr>
<tr>
<td>KOH activated graphene</td>
<td>Amorphous sulfur mesoporous graphene composites</td>
<td>Sulfur melt adsorption</td>
<td>67%</td>
<td>1007 mA h g⁻¹/0.2C/60th cycle</td>
<td>1 M LiTFSI/DOL-DME (1 : 1), 1.0–3.0 V</td>
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</table>
mesopores was used with the intention of regulating ion conduction in the mesoporous framework for high-rate performance. The small mesopores were located in the walls of the large mesopores without harming the structural integrity of the mesoporous framework. By screening over various mesoporous carbons with different mesopore sizes and pore volumes, Li et al. concluded that (1) large pore volume is good for increasing sulfur percentage and higher capacity of the composite (total mass of carbon and sulfur), (2) empty pore space, i.e. partial sulfur loading, is crucial for facile supply of lithium ions and adsorption of PS, and (3) polymer coating of mesoporous carbon is important for better cathode performance. It is important to emphasize that the empty space in C-S composites is necessary for buffering the volume change during lithiation–delithiation of the cathode. By using a post-activation treatment, Liang et al. showed the generation of micropores on the walls of mesoporous carbon. These micropores were expected to accommodate sulfur while the mesopores can transfer lithium ions. However, the cycling stability of this carbon–sulfur composite was not as good as the CMK-3–sulfur composite. The relatively larger mesopores (7.3 nm) as well as the destroyed pore walls may be the causes.
were used.\textsuperscript{33,34} Interestingly, the introduction of nitrogen as a dopant in mesoporous carbon appeared to be beneficial for performance enhancement.\textsuperscript{35}

Microporous carbon–sulfur composites

Zhang and co-workers reported the exceptional stability of sulfur confined in microporous carbon spheres.\textsuperscript{36} The carbon spheres were synthesized by refluxing sucrose (5 wt% solution) in a 6 M sulfuric acid at 120 °C for 10 h and carbonizing the black product at 1000 °C for 2 h under flowing Ar gas. The mixture of microporous carbon and sulfur was heated at 149 °C for 6 h and then the temperature raised to 300 °C and kept for 2 h to vaporize the excess sulfur. The microporous carbon spheres ranged in size from 200 to 300 nm. The micropore size was about 0.7 nm and no mesopores were detected. The sulfur loading percentage was seen to be important for high capacity. The total pore volume (0.474 cm\textsuperscript{3} g\textsuperscript{-1}) allowed a theoretical sulfur loading of 49.5 wt%. A loading content of 51 wt% showed much smaller capacity (~300 mA h g\textsuperscript{-1}) than a 42 wt% loading (~900 mA h g\textsuperscript{-1}). This is likely related to the issues of lower conductivity and severer volume expansion at high sulfur loading. This composite was stable for 500 cycles, which was related to the strong adsorption of PS by the narrow micropores.\textsuperscript{36} Zhang \textit{et al.} observed lithium potential hysteresis during the discharge process which reflects the additional energy required to overcome the adsorption energy of the short-chain sulfur in micropores (Fig. 4).\textsuperscript{36} Because of the strong confinement effect, low molecular sulfur in micropores can be only fully discharged at a potential of 1 V vs. Li/Li\textsuperscript{+}, compared with 1.5 V vs. Li/Li\textsuperscript{+} for sulfur in mesopores. Similar cathode behaviour of sulfur in microporous carbon was reported by Wang \textit{et al.}\textsuperscript{37} and Xin \textit{et al.}\textsuperscript{36}

Wang \textit{et al.} designed a two-step adsorption–extraction method to differentiate the differing cathode properties of sulfur confined in micropores and large pores.\textsuperscript{37} Their work clearly demonstrated the excellent efficacy of micropores in enhancing the stability of sulfur (up to 800 cycles at high rate).\textsuperscript{37} They also observed the higher binding energy of Li–S bond confined in micropores, which is likely because of the strong adsorption.\textsuperscript{37} They proposed a desolvation effect to interpret the unusual stability of sulfur in microporous carbon.\textsuperscript{37} Since the observation of ion desolvation in carbide-derived carbon, it has been widely acknowledged that solvated ions tend to be desolvated in micropores with size close to ion size.\textsuperscript{36} As generally accepted, the sulfur reduction is a solid (S)–liquid (PS ions)–solid (Li\textsubscript{2}S\textsubscript{2}/Li\textsubscript{2}S) process, and the mesopores stabilize sulfur through the adsorption of dissolved PS ions (Fig. 5, left). According to the ion–desolvation theory, the PS ions in micropores which are devoid of solvent react with desolvated ions (Fig. 5, right). This suggests a quasi-solid-state reaction of the sulfur under solvent-deficient conditions. The low Li\textsuperscript{+} conductivity in S and solid sulfides could also explain the retarded lithiation of sulfur in micropores. Xin \textit{et al.} fabricated microporous carbon nanofibers in which carbon nanotubes were the backbone.\textsuperscript{38} Much improved cathode stability of sulfur was also observed, as well as potential hysteresis.\textsuperscript{38} These three works\textsuperscript{36–38} shared the same properties of microporous carbon–sulfur composites: (1) lithiation potential hysteresis and (2) excellent...
stability. Xin and co-workers considered the lithiation of small sulfur molecules. On account of the tiny 0.5–0.6 nm micropores, Xin et al. proposed that cyclo-S$_4$–S8 molecules with at least two dimensions larger than 0.5 nm cannot exist inside, while small S$_2$–S4 molecules with at least one dimension less than 0.5 nm can be hosted. However, characterization of the molecular structure of amorphous sulfur confined in micropores is still a great challenge.

Hierarchical porous carbon–sulfur composites

Recent efforts have been devoted to designing hierarchical porous carbon with the aim of combining the advantages of microporous texture in regards to stability with large pores for rapid ion transport. An ordered hierarchical porous carbon was fabricated by a dual-template self-assembly approach. Typically, 350 nm colloidal polymer particles were self-assembled with 9 nm silica spheres to form a hierarchical periodic template. Sucrose was infiltrated as the carbon source together with sulfuric acid as the carbonization catalyst. The product has a pore volume of 1.4 cm$^3$ g$^{-1}$, bimodal macropores (120 nm, 300 nm), mesopores (10 nm) and micropores. With a 50 wt% loading of sulfur, the composite could be cycled for 50 times at 167.5 mA g$^{-1}$ with a capacity around 800 mA h g$^{-1}$. Fish scales were used as an alternative bio-derived carbon source to produce hierarchical porous carbon. The natural hierarchical texture of the fish scale was inherited in the final carbon product giving rise to the hierarchical porous texture. This bio-derived carbon has a very high specific surface area (2441 m$^2$ g$^{-1}$) and a good pore volume (1.69 cm$^3$ g$^{-1}$). Benefiting from the hierarchical porosity, the initial discharge capacity of sulfur in this carbon material can be as high as 1039 mA h g$^{-1}$ at 1675 mA g$^{-1}$ and remain as high as 1023 mA h g$^{-1}$ after 70 cycles.

Hollow carbon–sulfur composites

Jayaprakash et al. reported the synthesis of hollow carbon capsules with a mesoporous shell and their use as a sulfur host (Fig. 6a). In this design, the hollow carbon was synthesized using a silica template and petroleum pitch as the carbon source. The resulting product has a 3 nm average pore size and can accommodate up to 70 wt% sulfur. The vapour-phase infusion technique allowed the encapsulation of sulfur in the interior and the porous shell of the carbon capsule, which sequesters the active mass. Several advantages of this hollow carbon host were suggested: (1) maximizing the amount of sulfur sequestered by the capsules, (2) minimizing PS dissolution and shuttling, (3) preserving fast transport of lithium ions, and (4) good conductivity. Because of these attributes, the as-derived carbon–sulfur composite can be cycled at a low rate (850 mA g$^{-1}$) over 100 cycles with a high capacity of ca. 1000 mA h g$^{-1}$. In another approach, complex double-shelled hollow carbon spheres were synthesized using hollow SnO$_2$ spheres as a template. Glucose was uniformly coated on the interior and exterior surfaces of the template and the complex hollow carbon was derived thereafter (Fig. 6b). The double-shelled structure was designed to host higher amounts of sulfur, suppress the outward diffusion of dissolved PS and buffer the volume change upon cycling. A satisfactory capacity of around 500 mA h g$^{-1}$ was obtained at a current density of 850 mA g$^{-1}$.

Carbon black–sulfur composites

Carbon black is a commercial conducting additive used widely in the battery industry. Synthesis of carbon–sulfur composites from this abundant commercial material is of crucial economic value. Wang and co-workers reported the synthesis of core–shell carbon–sulfur nanocomposite through an aqueous solution route. Controlled direct deposition of sulfur from the following reaction:

$$\text{SO}_2 + 2\text{S}^{2–} + 4\text{H}^+ \rightarrow 3\text{S} + 2\text{H}_2\text{O}$$  (1)

can be uniformly performed on the exterior surface of carbon black particles with a thickness of 10 nm (Fig. 7a). This coating approach led to a very high sulfur loading of 84.62 wt%. The nanocomposite was ball milled with carbon additives and poly(ethylene oxide) for 4 hours, which may affect the as-obtained core–shell structure but was not verified. A mixture of 1,3-dioxolane and diethylene glycol dimethyl ether was used as a solvent to obtain a 1 M LiClO$_4$ electrolyte solution. In spite of the lack of confinement of sulfur in this composite as well as the shielded electron conduction by the sulfur coating, the cathode stability was surprisingly good (around 400 mA h g$^{-1}$ at 800 mA g$^{-1}$) for 50 cycles and the discharge potential did not reduce greatly when current density was increased.
samples with sulfur prepared from the same procedure but without precise coating showed much worse performance in terms of the large impedance and low capacity (~650 mA h g⁻¹). A cooling-precipitation method was adopted to fabricate a homogeneous composite of sulfur with carbon black (Fig. 7b). In this method, dimethyl sulfoxide was used as the solvent to dissolve sulfur at temperatures above 115 °C and the precipitation of sulfur occurred together with carbon black at temperatures below 80 °C. With this approach, 64.74 wt% sulfur could be loaded. The use of carbon black can significantly reduce the size of sulfur precipitates and allow a homogeneous distribution throughout the composite, giving rise to enhanced cathode performance.

**Carbon nanotube/fibre–sulfur composites**

Sulfur-coating strategies similar to those above have been used for the synthesis of sulfur-coated carbon nanotube (CNT) nanocomposites. It was suggested that the low surface tension of sulfur plays an important role in wetting of the exterior surface of CNTs. The mixture of the CNTs and sulfur was first ball milled and then annealed at 155 °C for 24 h. The CNT–S composite showed improved capacity and stability compared to a sulfur–carbon black composite and the sulfur–CNT mixture without the coating treatment. Disordered CNTs (DCNTs) with less graphitic tube walls and bamboo-like hollow voids have also been used as sulfur hosts. The sulfur was introduced in a vapour form in a vacuum at 160, 300, and 500 °C. The thermal stability of the DCNT–S composite increased as the annealing temperature increased, and the loss due to sulfur sublimation reduced, which suggested enhanced absorption and the possibility of sulfur–carbon bonding or new compounds being produced as indicated by unknown XRD peaks. However, the reduction of PS ions at the first plateau gradually vanished from 160 °C to 500 °C, which was accompanied by a remarkable potential hysteresis similar to that observed in microporous carbon–sulfur composites (Fig. 4 and 8). Assemblies of CNTs in the forms of microspheres or an aligned forest were studied.

Hollow carbon nanofibers (CNFs) were reported as a desirable host for a sulfur cathode mainly because of the limited diffusion pathways of PS ions (Fig. 9). This hollow design was believed to contribute the following ideal characteristics: (1) a closed structure for efficient PS containment, (2) limited surface area for sulfur–electrolyte contact, (3) sufficient space to accommodate sulfur volumetric expansion/shrinkage, and (4) a short electron and Li ion transport pathway. Ji et al. reported sulfur deposition onto PMMA-templated porous CNFs from a polysulfide solution. The reaction formula is as follows:

\[ \text{S}_x^{2-} + 2\text{H}^+ \rightarrow (x - 1)\text{S} + \text{H}_2\text{S} \]  

The as-obtained porous CNF–S composite was further annealed to ensure that the exterior sulfur can diffuse into the interior pores of the nanofibers.

**Graphene sheet–sulfur composites**

The feasibility of graphene sheets for stabilizing sulfur cathodes as well as the use of conducting additives were first reported by researchers using solvothermal-derived porous graphene sheets. Successive optimization of the synthesis methods resulted in diverse configurations of the graphene–sulfur composites: graphene-wrapped sulfur particles, sandwich-type composites, and sulfur coated graphene sheets. Disproportionation of thiosulfate as follows:

\[ \text{S}_2\text{O}_3^{2-} + 2\text{H}^+ \rightarrow \text{SO}_2 + \text{S} + \text{H}_2\text{O} \]
in acid was used to synthesize uniform sulfur particles which were coated by graphene sheets (Fig. 10a).24 HF acid etching of the graphene sheets can help the nucleation and growth of the sulfur precipitate from the reaction bath.25 A facile solvent (CS2)-evaporation method was aided by ultrasonication to coat the sulfur particles with hydrophilic graphene oxide sheets which were reduced by ammonia solution to graphene sheets.26 Trace residual nitrogen was present in this composite and might be beneficial for cathode performance.27 Sandwich-structured graphene–sulfur composites can also be synthesized by thermal annealing, and a Nafion polymer coating was observed to be good for better stability (Fig. 10b).28 Expanded graphite, which is composed of loosely stacked graphene sheets with a preserved layered structure, was used as an intercalation host for a sulfur cathode.58–60 Direct annealing of the graphene–sulfur mixture in air resulted in sulfur-coated graphene sheets.61 An interesting two-step method is reported to prepare uniform sulfur-coated graphene sheets.62 The reduction and synchronous sulfur-modification of graphene oxide by sulfide and sulfate were found to be beneficial for the homogeneous growth of a sulfur film on graphene sheets.62 Similar chemistry but using a hydrothermal technique has also been demonstrated.63 Graphene sheets are advantageous for wrapping sulfur because of their large lateral size, good conductivity and the flexible structure. Under ideal conditions, however, graphene sheets conduct ions along the lateral direction making ion conduction across the sheets very difficult. To overcome this problem, as well as to provide more sites for sulfur storage, activated graphene sheets have been synthesized and used as the sulfur hosts.64 KOH activation produced 3.8 nm mesopores on the graphene sheets which is large enough to transport Li ions while being small enough to hold PS ions.64 Compared with pristine graphene sheets, the activation was clearly good for rate performance and stability.64 Graphene-based hybrid materials are of special interest in energy storage. A unique graphene–single walled CNT (SWCNT) hybrid structure was developed by Zhao and co-workers.65 The graphene constituent in the hybrid formed a hexagonal hollow container for sulfur while the exterior SWCNT forests provided electron conduction pathways as well as conferring a mesoporous texture to adsorb PS ions (Fig. 11). The highly graphitic nature of this hybrid is of significance for the excellent high-rate performance.65

Apart from the diverse range of synthesis chemistry and approaches for producing graphene–sulfur composites, the interaction between graphene and sulfur has also been of interest. Residual oxygen functional groups attached on graphene sheets were found to enhance the adsorption of sulfur and PS ions, both of which contribute greatly to the composite stability.66,67

**Polycrylonitrile-derived carbon–sulfur composites**

Polycrylonitrile (PAN) was first used for the preparation of carbon–sulfur composites more than a decade ago.68 A simple process was adopted by directly heating the mixture of PAN powder and sublimed sulfur at 280 to 300 °C in an Ar gas environment for 6 h. Since elemental sulfur is an effective dehydrogenating reagent, the ratio of H to C atoms in the final product (1 : 3) was lower than that in pristine PAN (1 : 1) (Fig. 12a).68 The effective dehydrogenation of PAN by sulfur gave rise to a conducting main chain. NMR results suggested the formation of a carbon double bond in the final product and the cyclization of the –CN group, which was highly thermally stable. As such, this PAN-derived carbon–sulfur composite had very

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**Fig. 10** (a) Graphene-sheet wrapped sulfur particles,64 Reproduced from ref. 54. Copyright 2011 American Chemical Society. (b) Sandwich-type graphene–sulfur composite.67 Reproduced from ref. 57. Copyright 2010 Royal Society of Chemistry.

**Fig. 11** (Left) illustration of the graphene–SWCNT hybrid as a sulfur host and (right) SEM image of the hybrid-sulfur composite.68 Reproduced from ref. 65. Copyright 2012 American Chemical Society.

**Fig. 12** (a) Reaction mechanism during heating sulfur with PAN and (b) discharge–charge curves of a PAN-derived carbon–sulfur composite.68 Reproduced from ref. 69. Copyright 2003 Wiley-VCH.
Flexible carbon–sulfur composites

In an innovative approach, activated carbon fibre (ACF) cloth has been recently demonstrated as a binder-free cathode support for sulfur. This ACF cloth is monolithic and holds the sulfur in the micro pores. The structural integrity of the monolithic ACF cloth is sufficient to withstand the volume variation during lithiation and delithiation. Meanwhile, the abundant micropores that can accommodate sulfur promised excellent resilience and physical–chemical attraction of polysulfide.

Tremendous progress in carbon–sulfur composites has been achieved in recent years. Substantial attention has been paid to the morphology, chemistry and electrochemistry of carbon–sulfur composites. In the future the adsorption, intercalation, and functionalization of carbon materials still require further investigation in terms of the cathode stabilization through the confinement and physical–chemical attraction of polysulfide.
thus the optimal loading of sulfur is a balance between the desire for maximum capacity and the need to allow for the volume change to ensure stability. Hollow carbons with rigid shells and large internal voids are desirable for this purpose. Similarly, graphene sheets with good flexibility are excellent for buffering volume change.

A summary of the efficacy of the different kinds of carbons for stability, capacity and conductivity is given in Table 3. It is clear that no one carbon form alone can meet the comprehensive performance criteria, but an effective combination of these materials is necessary to gain the best advantage from the properties of the individual building blocks.

Several critical factors are suggested here for the rational design of advanced carbon–sulfur composites:

(1) Pore size: micropores/small mesopores for strongly confining polysulfides.

(2) Large pore volume for maximum sulfur loading: carbon is not an active cathode material in Li–S batteries and its weight ratio is required to be as low as possible without harming the overall performance of sulfur–carbon cathodes. However, the very low conductivity of the composite when sulfur content increases is a critical problem to be solved.

(3) Graphitization level: a preferential content of graphitic carbon to facilitate electron conduction to the insulating sulfur/lithium sulfides.

(4) Electrolyte impregnation and lithium ion migration: a short pathway to preserve fast migration of lithium ions from the bulk electrolyte to active sulfur and the release of lithium ions from lithium sulfides. Voids are required in the final carbon–sulfur composites.

(5) Flexible or rigid carbon scaffold: volume change occurs during the discharge or charge of sulfur–carbon cathodes; the carbon host should buffer the stress-induced strain and survive over extended cycles.

(6) Low cost production and easy scale-up: templated ordered mesoporous carbon and CNTs/nanofibers are highly effective due to their periodic structure but are unlikely to be produced at an industrial scale for Li–S batteries owing to their high cost and unsatisfactory performance–cost ratio; novel continuous synthesis techniques, inclusive of but not limited to atomization carbonization and hydrothermal carbonization, are promising.

(7) An advanced technique to form carbon–sulfur composites: common methods are impregnation with the sulfur melt or sulfur organic solution, disproportionation reaction and in-situ encapsulation, and vapour diffusion. These are post-carbon-synthesis and complicate the industrial processing. A one-step method for fabrication of sulfur–carbon composites is necessary.

We emphasize here that the correlations between the carbon structure (porosity, surface chemistry, graphitic degree) and the sulfur structure need to be comprehensively studied and optimized (Fig. 14). Facile and low cost material fabrication techniques are also desired. An optimal carbon–sulfur composite will benefit from a combination of fundamental insights and an advanced synthesis approach. We also acknowledge that the sulfur cathode problem is not the only issue that impedes Li–S technology. Problems with the lithium anode, the electrolyte as well as engineering difficulties in fabricating lithium metal batteries are all great challenges to be faced, but recent dramatic progress in carbon–sulfur composites is likely to form the basis for future commercialization of Li–S batteries.

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Notes and references

Mechanisms of sulfur reactions

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