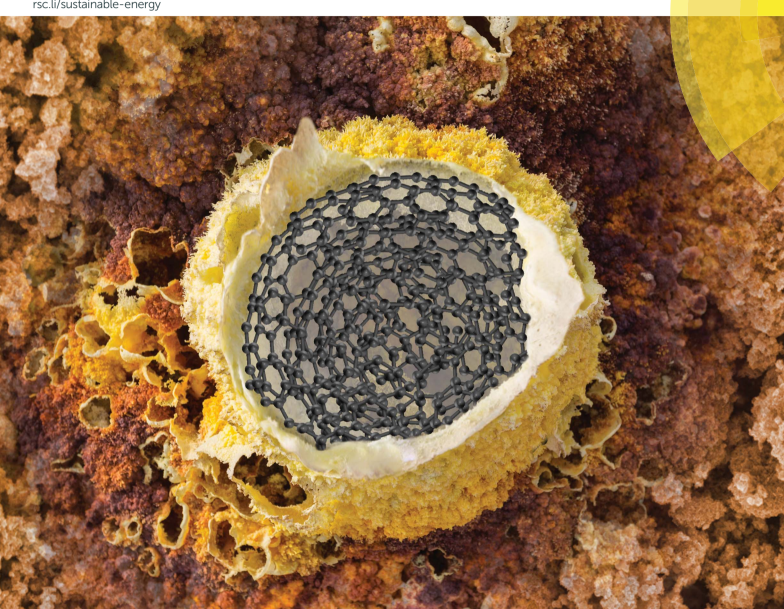
# Sustainable Energy & Fuels

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ISSN 2398-4902



# Sustainable **Energy & Fuels**



**PAPER** 

View Article Online



Cite this: Sustainable Energy Fuels, 2017, 1, 84

Received 2nd November 2016 Accepted 3rd January 2017

DOI: 10.1039/c6se00034g

rsc.li/sustainable-energy

# Carbon onion-sulfur hybrid cathodes for lithiumsulfur batteries†

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In this study, we explore carbon onions (diameter below 10 nm), for the first time, as a substrate material for lithium sulfur cathodes. We introduce several scalable synthesis routes to fabricate carbon onion-sulfur hybrids by adopting in situ and melt diffusion strategies with sulfur fractions up to 68 mass%. The conducting skeleton of agglomerated carbon onions proved to be responsible for keeping active sulfur always in close vicinity to the conducting matrix. Therefore, the hybrids are found to be efficient cathodes for Li-S batteries, yielding 97-98% Coulombic efficiency over 150 cycles with a slow fading of the specific capacity (ca. 660 mA h q<sup>-1</sup> after 150 cycles) in long term cycle test and rate capability experiments.

### Introduction

The increase in global energy demand and limited reserve of fossil fuel resources have led to intense research efforts for efficient and mobile energy storage technologies, especially rechargeable batteries.1 The further implementation of current lithium-ion battery technology is impeded by high costs, safety concerns, and the limited energy density associated with stateof-the-art insertion-compound electrodes like layered LiCoO<sub>2</sub>, spinel LiMn<sub>2</sub>O<sub>4</sub>, olivine LiFePO<sub>4</sub> cathodes, and graphite anodes.<sup>2,3</sup> The development of alternative low-cost, abundant electrode materials with high energy and power densities is crucial for widely employing rechargeable batteries for the storage of renewable energies and the transportation sectors. 4,5

Lithium-sulfur (Li-S) systems have emerged in recent years as a promising battery class.<sup>6-9</sup> In addition to the high specific capacity, using sulfur as a cathode material has the advantages of high natural abundance, low cost, and high environmental friendliness.7 Although the lithium-sulfur system operates at a relatively low average potential of 2.1 V against Li/Li<sup>+</sup>, it shows a very high theoretical specific capacity of 1675 mA h g<sup>-1</sup> and a high theoretical specific energy of 2600 W h kg<sup>-1</sup>.<sup>5,10</sup> A remaining hindrance to reach the theoretical limit is the low

sulfur utilization for Li-S rechargeable systems. Still, Li-S batteries suffer from the isolative behavior of the start (S<sub>8</sub>) and end product (Li2S), active material loss by dissolution of intermediate species (polysulfides) in the electrolyte, and shuttle mechanisms leading to anode passivation by the deposition of Li<sub>2</sub>S<sub>2</sub> and Li<sub>2</sub>S.<sup>11,12</sup>

Several studies have shown different carbons as a conductive substrate for insulating sulfur including a variety of carbons with a high surface area and conductivity. Among others, researchers have investigated multi-walled carbon nanotubes, 13-15 mesoporous carbon,16 graphene,14,17 carbide-derived carbon,18 biomass derived carbon, 19,20 three-dimensional bicontinuous gyroidal carbon,<sup>21,22</sup> carbon from polyacrylonitrile,<sup>23</sup> activated carbon fiber cloth,24 and hard25 or soft templated carbons26 as substrates for sulfur in lithium-sulfur batteries.

In this work, we explore carbon onions for lithium sulfur batteries. Carbon onions, also known as onion-like carbon (OLC), consist of several concentrically stacked graphitic carbon spheres.27 A facile and scalable synthesis route for carbon onions is thermal annealing of nanodiamond powder in an inert atmosphere or vacuum.28 During the annealing process at temperatures above 1300 °C, the sp<sup>3</sup>-hybridized nanodiamond is progressively converted to quasi-spherical sp<sup>2</sup>-carbon onions.29 The high electrical conductivity of carbon onion aggregates (ca. 4 S cm<sup>-1</sup>) provides a conductive network for sulfur, while the enhanced surface area (200-600  $\text{m}^2\text{ g}^{-1}$ ) and nanoscopic size (5-10 nm),30 hierarchical porous structure, and possibility to create a mechanically robust electrode structure are attractive for creating a large sulfur/electrolyte interface, mitigating polysulfide shuttling. Our work compared different synthesis approaches for obtaining carbon onion/sulfur electrodes, provides structural and chemical characterization, and benchmarks the resulting electrochemical performance.

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<sup>†</sup> Electronic supplementary information (ESI) available. See 10.1039/c6se00034g

## 2. Experimental description

#### Materials

Nanodiamond powder was purchased from NaBond Technologies. Battery-grade conductive carbon black (C65) was purchased from Imerys. Sodium thiosulfate pentahydrate (Na<sub>2</sub>S<sub>2</sub>O<sub>3</sub>·5H<sub>2</sub>O), elemental sulfur (S<sub>8</sub>), polyvinylidene fluoride (PVDF) powder (molecular mass *ca.* 534 000 g mol<sup>-1</sup>), *N*-methyl2-pyrrolidone (NMP), bis(trifluoromethane)sulfonimide lithium salt (LiTFSI), 1,2-dimethoxyethane (DME), 1,3-dioxolane (DOL), and lithium nitrate (LiNO<sub>3</sub>) were purchased from Sigma Aldrich, and used without any further purification. Nickel foil with a thickness of 20  $\mu$ m was procured from Schlenk Metallfolien. Electrochemical grade high purity (99.9%) lithium was purchased from PI-Kem. A trilayer porous polyolefine separator was obtained from Celgard and we used non-woven polypropylene separators from Freudenberg.

#### Synthesis of carbon onions

Carbon onions were synthesized by thermal annealing of the detonation nanodiamond with a diameter in the range of 4–6 nm (NaBond Technologies). The nanodiamond powder was annealed in an argon atmosphere in a graphite crucible using a water cooled high temperature furnace with a tungsten heater (Thermal Technology) at 1700 °C for 1 h (heating/cooling rate: 20 °C min<sup>-1</sup>). For more information, see ref. 28.

#### Preparation of carbon onion-sulfur composites

Herein, three different approaches were investigated (Fig. 1): (1) mechanical melt mixing of sulfur with carbon onions, (2) *in situ* formation of sulfur nanoparticles on the carbon surface, and (3) melting of the covering nanoparticles to form homogenous layers around the carbon onions. The melt mixing carbon onion–sulfur composite was prepared as follows: 65 mass% elemental sulfur ( $S_8$ ) were added to 35 mass% of carbon onions. It was primarily mixed in a mortar, and afterwards ball milled to get nearly uniform distribution of the sulfur. The carbon–sulfur

composite powder was thermally annealed at 155  $^{\circ}$ C for 5 h under an argon atmosphere. Using this treatment sulfur is fluid and attains minimum viscosity (ref. 16) so that uniform coverage of sulfur over all the available surface of carbon onion particles was obtained.

For *in situ* formation of sulfur nanoparticles on the carbon onion surface, 1.55 g of sodium thiosulfate was first dissolved in 40 mL of water. Afterwards, 100 mg of carbon onions was added with subsequent sonication for 30 min under ice-cold water. In the next step, 2.5 mL of 5 M HCl was added dropwise to the system. By this time, sulfur nanoparticles were formed in the mixture. The mixture was left to react for 2 h and was filtered through a 0.2  $\mu$ m hydrophilic PVDF filter. It was washed with copious amounts of water until pH 7 was achieved. The carbon onion/sulfur mixture was dried at 80 °C under vacuum for 12 h.

For the third OLC-S hybrid, the carbon onions/sulfur nanoparticle mixture was thermally annealed at 155  $^{\circ}$ C following the same protocol maintained for the first set of samples.

For comparison, electrodes of the same thickness were made with sulfur nanoparticles along with 10 mass% conductive additive (Super C65) and 8 mass% PVDF binder. The slurry was cast onto nickel foil following the same way as done for carbon onion/sulfur hybrids. This electrode is designated as SNP (sulfur nanoparticle) electrode.

#### **Electrode** preparation

The three sets of carbon–sulfur composite powders were individually mixed with 8 mass% of PVDF binder in NMP to attain the desired slurry viscosity. The slurry was cast on a nickel current collector, achieving a 300  $\mu$ m wet electrode thickness, and afterwards dried in an oven operated under vacuum at 60 °C for 12 h to get rid of any traces of solvent. In the dry state, each electrode has a mass loading of 3–4 mg cm<sup>-2</sup> (corresponding sulfur loading is 2–3 mg cm<sup>-2</sup>) which is at the level to attain the required areal capacity desired for applications in electrically driven vehicles.<sup>31</sup>

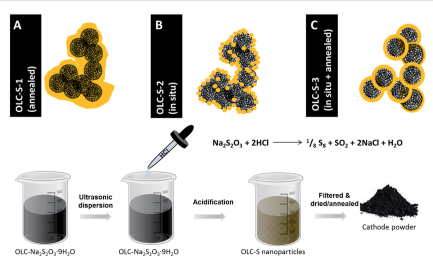


Fig. 1 Schematic representation of fabrication of OLC-S hybrids following three different approaches.

Conductivity measurements

The same slurries for each set of samples for preparing electrodes were again cast over  $50~\mu m$  thin polyimide foil for conductivity measurements. Sheet resistance measurements were made with a custom-built spring-loaded four-point probe with blunt gold contacts (tip diameter: 1.5 mm, tip distance: 3 mm).

#### Raman spectroscopy

Raman spectra were recorded with a Renishaw inVia Raman Microscope employing an Nd:YAG laser with an excitation wavelength of 532 nm or HeNe laser operating at 633 nm. A grating with 2400 lines per mm and a  $50\times$  objective were used to reach a spectral resolution of about 1.2 cm $^{-1}$ . The laser spot on the sample was about 2  $\mu m$  in diameter at a power of 0.2 mW. The acquisition time of each spectrum was 30 s, and 10 accumulations were averaged.

#### X-ray diffraction

The crystalline structure of the carbon–sulfur hybrids was analyzed by X-ray diffraction employing a D8 Advance diffractometer (Bruker AXS) with a copper X-ray source ( $Cu_{K\alpha}$ , 40 kV, 40 mA), a Göbel mirror, and a 2D area detector (VANTEC-500) that covers about 25°  $2\theta$ . All samples were measured in a range from 10 to 60°  $2\theta$  in 3 steps, with a step duration of 24 min.

#### Gas sorption measurements

Porosity analysis was carried out using an Autosorb iQ nitrogen gas sorption system (Quantachrome). The carbon onion powder was first outgassed at 300 °C for 10 h under vacuum ( $10^2$  Pa). The nitrogen gas sorption analysis was performed at a temperature of -196 °C in the relative pressure range from  $5 \times 10^{-7}$  to 1.0 in 68 steps. The pore size distribution (PSD) was calculated using the quenched-solid density functional theory (QSDFT)<sup>32,33</sup> supplied by Quantachrome assuming a slit-shape pore geometry. The BET SSA after Brunauer–Emmett–Teller<sup>34</sup> was calculated in the linear regime of the measured isotherms between 0.1 and  $0.3P/P_0$ . Carbon onion sulfur hybrids were not measured using gas sorption analysis due to the evaporation of sulfur and the possible contamination of the system.

#### Thermogravimetric analysis

To estimate the loading of sulfur in each carbon–sulfur hybrid, thermogravimetric analysis (TGA) was adopted. As sulfur sublimes when heated to 500  $^{\circ}$ C, TGA of the carbon–sulfur hybrids was performed using a Netzsch Libra TG 209 F1 in the temperature range of 30–500  $^{\circ}$ C with a heating rate of 10  $^{\circ}$ C min<sup>-1</sup> under continuous flow of argon.

#### CHNS elemental analysis

The chemical composition was further investigated with CHNS elemental analysis with a Vario Micro Cube system (Elementar Analysensysteme). After combustion, the samples were measured under oxygen at  $1150~^{\circ}$ C in a tin holder. The CHNS analyzer was

calibrated with sulfanilic acid using different masses (41.6 mass% C, 4.1 mass% H, 8.1 mass% N, 18.5 mass% S).

#### **Energy filtered transmission electron microscopy**

The specimen was prepared by dispersing the powders in ethanol followed by placing a drop on a lacey carbon film TEM grid (Plano GmbH). The TEM grid was dried at 50 °C to remove traces of ethanol before inserting into the TEM instrument. Transmission electron micrographs were recorded with a JEOL 2100F microscope at 200 kV. Energy filtered imaging (EFTEM) was conducted on a Libra 120 transmission electron microscope (Carl Zeiss) operated at 120 keV.

#### **Electrochemical benchmarking**

The coin-cell type cathodes were cut as discs of 14.2 mm diameter from the entire piece and 2032 coin cells were assembled in an argon filled glovebox (O2 & H2O: <1 ppm) with carbon onion/sulfur hybrid cathodes, lithium metal discs as the anode, and a porous trilayer PP-PE-PP membrane from Celgard and Freudenberg non-woven PP mat together as the separator (PP = polypropylene; PE = polyethylene). Each electrode has a sulfur loading of ca. 60 mass% (excluding binder). The separators were soaked with 100 µL of 1 M LiTFSI in 1:1 (by volume; v/v) 1,2-dimethoxyethane (DME) and 1,3-dioxolane (DOL) along with 0.25 M LiNO<sub>3</sub> serving as the electrolyte. Cyclic voltammetry was performed with a Biologic VPM-300 potentiostat-galvanostat within the potential range of 1 V to 3 V versus Li/Li<sup>+</sup> at a scan rate of 100 μV s<sup>-1</sup> (equivalent to 0.2C). Galvanostatic charge/ discharge tests were carried out with an Astrol BatSmall battery analyzer at a constant current density of 336 mA  $g^{-1}$  (=0.2C) for charging and 168 mA g<sup>-1</sup> (=0.1C) for discharging within the potential window of 1.8 V to 2.6 V versus Li/Li<sup>+</sup>. The rate capability experiments were done at different rates from 0.1C (discharge)/0.2C (charge) for 20 cycles, 0.2C (discharge)/0.4C (charge) for the next 10 cycles, 0.5C (discharge)/1C (charge) for further 10 cycles, 1C (discharge)/2C (charge) and then brought back to 0.1C (discharge)/0.2C (charge), and allowed to run for the next 20 charging-discharging cycles. For high rate cycle stability tests, the carbon onion hybrid cathode containing cells were subjected to discharge at 1C and charged at 2C for 500 cycles. In all our galvanostatic charge-discharge experiments, we set the charging rate at two times the discharge rate to simulate the actual usage of battery where a faster charging is required.21

## 3. Results and discussion

Carbon onions (OLC) were synthesized by thermal annealing of nanodiamonds (ND) at 1700  $^{\circ}$ C in argon.  $^{29,30}$  Using these conditions, ND particles with a mean size of  $\sim$ 5 nm completely transform to sp<sup>2</sup>-hybridized carbon onions. During this process, the diamond structure with a lattice spacing of 0.21 nm rearranges to a graphitic multi-shell carbon onion with an outer lattice spacing of 0.34 nm (Fig. S1, ESI†). Between the single carbon onions with a diameter between 5 and 10 nm, we also

see few-layer graphene nanoribbons as a result of the annealing process (Fig. S1, ESI†).28

Carbon onion/sulfur (OLC-S) hybrids were synthesized using three different approaches, as described in the Experimental section. The overall distribution of sulfur on the carbon onion surface and in the interparticle voids was surveyed with energy filtered transmission electron microscopy (EFTEM). The sulfur signal was confirmed by comparing energy filtered images below (153 eV) and above (200 eV) the sulfur ionization edge (165 eV). As seen from Fig. 2B, the in situ deposition of sulfur on OLC (OLC-S-2) shows the most uniform sulfur distribution, whereas the melt-mixing (OLC-S-1) leads to a larger degree of localized sulfur clustering (Fig. 2A). Further annealing of in situ deposited sulfur on OLC (OLC-S-3) resulted in a low degree of sulfur clustering. Carbon onions act as nucleating surfaces for sulfur nanoparticle deposition, whereas the melt diffusion strategy results in non-uniform spreading of molten sulfur in a random manner. However, the in situ sulfur nanoparticle deposition on carbon onion surfaces followed by thermal annealing leads to a low extent of sulfur clustering compared to the mechanical melt mixing pathway, as evident from the carbon-sulfur colored overlay images (Fig. 2C).

The nitrogen gas sorption measurement of carbon onions shows a surface area of 425 m<sup>2</sup> g<sup>-1</sup> (BET) and 402 m<sup>2</sup> g<sup>-1</sup> (DFT) with a pore volume of 1.21 cm<sup>3</sup> g<sup>-1</sup>. The pore size distribution pattern can be differentiated into three regimes: pores related to particles in direct contact (for pores <1 nm), particles separated by spacer particles (at 1 nm pore size and in the mesopore range), and pores between particle agglomerates (Fig. 3A). These spaces among the particle interstices were utilized to embed insulating sulfur to attain a high degree of intimate contact with the carbon onion surfaces. Additionally, the pore hierarchy also plays a crucial role in functioning of lithiumsulfur batteries by accommodating a large quantity of sulfur mass as well as electrolyte wetting together with lowering the tendency of polysulfide migration out of the cathode to the bulk electrolyte.15,35-39

Thermogravimetric analysis of pristine carbon onions undergoes a very little mass loss of ca. 1% up to 500 °C. This mass loss is attributed to the decomposition of functional groups on the surface of carbon onions. The thermogravimetric measurements of OLC-S hybrids demonstrate one step mass loss at around 300 °C for all samples with a mass loss of 68% OLC-S-1 and 65% for OLC-S-2 and OLC-S-3 (Fig. 3B). The respective mass loss corresponds to the sulfur content from CHNS elemental analysis (Table 1). The absence of any other step in the thermogram indicates that sulfur is only coated onto the carbon onions and does not penetrate the inside of the particles, irrespective of the deposition process. For contrast, hierarchical porous carbons exhibit more than one step of mass loss during thermogravimetry. 40,41 It is reported in the literature that the sulfur confined in micropores requires a higher thermal energy and is released at much higher temperature than the sulfur present outside the micropores. 40,41

The structure of carbon onions was not changed by the synthesis procedures as indicated by Raman spectroscopy (Fig. 3C, Table 2, and Fig. S2, ESI†) and XRD (Fig. 3D). Independent of the treatment, carbon onions are characterized by their typical Raman spectrum with the carbon D-mode at 1338 cm<sup>-1</sup>, the G-mode at 1577 cm<sup>-1</sup>, and a distinct second order spectrum (Fig. S2, ESI†). The G-mode, representative for sp<sup>2</sup>-hybridized carbon, is measured at 1577 cm<sup>-1</sup>, indicating nanocrystalline and partially amorphous carbon. Amorphous carbon is mostly located between the particles (Fig. S1B, ESI†), leading to partially connected particles and a strong degree of agglomeration. Carbon onions present a rather high degree of

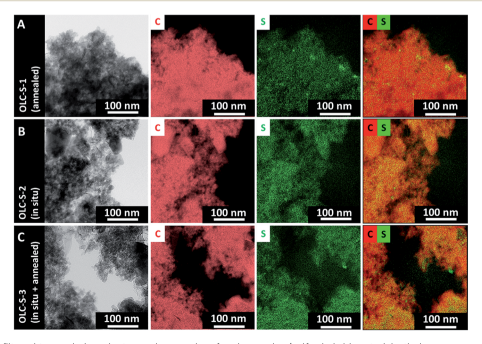


Fig. 2 (A-C) Energy filtered transmission electron micrographs of carbon onion/sulfur hybrid materials, their corresponding elemental maps, and the overlay of carbon and sulfur elemental mappings.

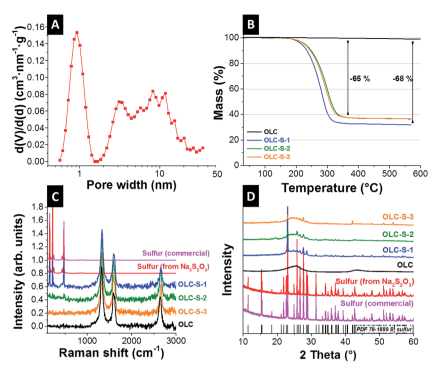


Fig. 3 (A) Pore size distribution of OLC calculated from QSDFT assuming slit pores, (B) sulfur contents from TGA, (C) Raman spectra (633 nm excitation wavelength) and (D) X-ray diffractograms of carbon onions and carbon onion/sulfur hybrids.

Table 1 Estimation of sulfur contents from TGA and CHNS elemental analysis

|         | Sulfur (mass%) |          |  |
|---------|----------------|----------|--|
| Sample  | TGA            | CHNS     |  |
| OLC-S-1 | 68             | $68\pm1$ |  |
| OLC-S-2 | 65             | $62\pm5$ |  |
| OLC-S-3 | 65             | $64\pm1$ |  |

ordering as demonstrated by the sharp G-mode (FWHM of 44 cm<sup>-1</sup>) and the distinct second order peaks at 2675 cm<sup>-1</sup>. By virtue of the instability under laser irradiation, we did not detect any sulfur when using a laser wavelength of 532 nm. To avoid such degradation, we conducted also experiments at 633 nm excitation wavelength (Fig. 3C). Commercial sulfur and sulfur derived from sodium thiosulfate were also characterized by Raman spectroscopy. Sulfur shows three distinct peaks at 156, 219, and 473 cm<sup>-1</sup>, which are related to the vibrations of the S-S bonds.14,19,42 When measuring the OLC-S hybrids by Raman

spectroscopy, OLC-S-2 and OLC-S-3 showed very weak sulfur signals, whereas OLC-S-1 demonstrated very clearly all three characteristic sulfur peaks. This indicates the presence of nonuniform distributions of sulfur onto the carbon onion surfaces whereas the two other hybrids claim to have nearly homogeneous coverage of sulfur on OLC. As a result, the Raman signals from carbon predominate over sulfur signals although all three hybrids contain ca. 65 mass% of sulfur (relative to the carbon mass).

The incompletely crystalline structure of carbon onions was further assessed by XRD (Fig. 3D). The broadening of the (002) graphitic carbon peak at  $\sim 26^{\circ}$   $2\theta$  is a consequence of the nanoscopic carbon particle size of 5-7 nm. In all three OLC-S hybrids, all crystalline peaks of sulfur are detected in alignment with PDF 78-1889. This confirms the growth of crystalline sulfur domains on the carbon onion substrates and ensures the allocation of sulfur in interparticle voids. When carbon with a large inner porosity is used as the host for sulfur, molten sulfur or nanoparticle precursor solutions tend to infiltrate into the pores driven by capillary forces. 16,24 In such a case, the imbibed sulfur in the pores does not display the well-resolved crystalline

Table 2 Raman spectroscopy results of OLC and OLC-S hybrids

| Sample  | Position D-band (cm <sup>-1</sup> ) | Position G-band (cm <sup>-1</sup> ) | FWHM D-band (cm <sup>-1</sup> ) | FWHM G-band (cm <sup>-1</sup> ) | $I_{ m D}/I_{ m G}$ |
|---------|-------------------------------------|-------------------------------------|---------------------------------|---------------------------------|---------------------|
| OLC     | 1338                                | 1580                                | 57                              | 44                              | 1.43                |
| OLC-S-1 | 1339                                | 1579                                | 67                              | 48                              | 1.79                |
| OLC-S-2 | 1339                                | 1577                                | 62                              | 46                              | 1.70                |
| OLC-S-3 | 1338                                | 1576                                | 62                              | 48                              | 1.69                |

peaks. 16,24 In the OLC-S hybrids, sulfur remains at the exterior surfaces of OLC particles. For this reason, no significant differences were visible in the XRD pattern of three different hybrid types.

The usefulness of the carbon onions as a host for embedding sulfur was further tested by sheet resistance measurements of the 80–100  $\mu m$  (dry) thick electrodes coated on polyimide foil (Table 3). Using a four-point-probe, the measured electrical conductivity of all samples showed little difference, with values around 0.2 S cm $^{-1}$ .

Electrochemical characterization including cyclic voltammetry (three-electrode setup) and galvanostatic charge-discharge measurements (two-electrode setup) was performed by using the OLC-S composite electrode as the working electrode (cathode) and lithium as the counter electrode (anode)

Table 3 Conductivity measurements *via* the four-point-probe of OLC-S hybrids

| Sample            | Conductivity (S cm <sup>-1</sup> ) |
|-------------------|------------------------------------|
| OLC-S-1 electrode | $19.8 	imes 10^{-2}$               |
| OLC-S-2 electrode | $23.3\times10^{-2}$                |
| OLC-S-3 electrode | $20.8\times10^{-2}$                |

and reference electrode. The cyclic voltammograms show the characteristic shape of sulfur oxidation and reduction (Fig. 4A, C and E). During discharge, the characteristic two reduction peaks at around 2.4 V and 2.0 V vs. Li appear (Fig. 4A, C and E). These peaks correspond to the ring opening reduction of sulfur to  $\text{Li}_2S_n$  (n > 4) and the subsequent reduction of higher order polysulfides to short chain polysulfide species, respectively.<sup>43</sup> The anodic sweep shows a characteristic single peak at around 2.4 V vs. Li. The relatively narrow peaks for OLC-S-2 and stable peak positions for oxidation and reduction indicate good reaction kinetics for reduction and oxidation reactions in the first cycle and run in a comparable range for all cycles. The two other hybrids OLC-S-1 and OLC-S-3 have much broader oxidation and reduction peaks which are indicative of relatively slow and inferior reaction kinetics compared to OLC-S-2. Such broadening of redox peaks arises from uneven distribution of sulfur in the OLC matrix. Thus, the reaction steps are overlapping and yield broader redox peaks, which indicate poor performances such as low energy efficiency and slow kinetics. The poorer performance can be related to larger amounts of inactive sulfur insufficiently connected to the conductive carbon onion substrate. In the case of OLC-S-3, due to the thermal treatment after nanoparticle formation, the molten sulfur moves to some extent from the surface and reorganizes as larger sulfur

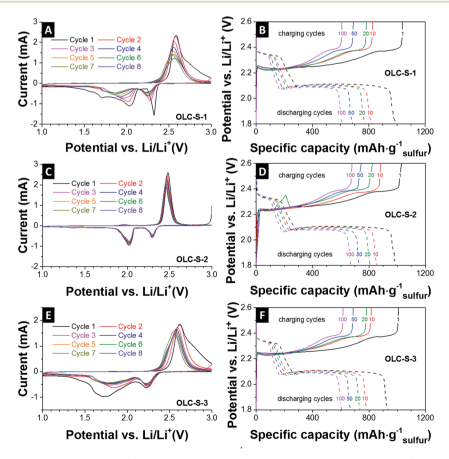


Fig. 4 Cyclic voltammograms at a rate of  $0.1 \, \text{mV} \, \text{s}^{-1}$  and galvanostatic charge—discharge curves at 336 mA  $g^{-1}$  (0.2C) for charging and 168 mA  $g^{-1}$  (0.1C) for discharging within the potential window of 1.8 V to 2.6 V *versus* Li/Li<sup>+</sup> of (A and B) OLC-S-1, (C and D) OLC-S-2, and (E and F) OLC-S-3.

particles, which are not ideally covering the carbon onion surfaces anymore. Those agglomerated parts remain electrochemically inactive, thus demonstrating poorer performance compared to OLC-S-2.

The reversible galvanostatic charge–discharge tests of the OLC-S hybrids were carried out in 2032 coin cells at 0.1C discharge rate (167 mA g $^{-1}$ ) and 0.2C charging rate (334 mA g $^{-1}$ ). The plateaus in the voltage profiles (Fig. 4B, D and F) match the peak voltages of the cyclic voltammograms for reduction and oxidation (Fig. 4A, C and E). We identify two plateaus in the discharge process, as summarized by Ji *et al.* (ref. 16):

- The first one contributes a minor part to the overall capacity from 2.4 V to ca. 2.1 V vs. Li. This plateau corresponds to the ring opening conversion of elemental sulfur (S<sub>8</sub>) to Li polysulfide anions (Li<sub>2</sub>S<sub>x</sub>; where x is typically 4–5).<sup>6,7,11</sup> The kinetics of this step are fast.
- The second plateau appears at around 2.1 V vs. Li due to the conversion reaction of higher order polysulfides to Li<sub>2</sub>S<sub>2</sub> and then to Li<sub>2</sub>S. This reaction occurs at a much slower rate and contributes to the large fraction of the overall capacity.

The similarity of voltage profiles at different cycles is indicative of slower capacity fading with prolonged cycling.

Additionally, the voltage plateaus of all samples that appear at the same potential during discharge and charge reflect the mitigation of active mass loss per charging-discharging cycles.

The corresponding specific capacities for charging and discharging are represented normalized to the sulfur mass (Fig. 5A). The specific capacities of all hybrids present values of 900–1200 mA h  $g_{sulfur}^{-1}$  in the initial cycles (up to 5 cycles). After 10 charge-discharge cycles, OLC-S-2 and OLC-S-3 hybrids showed only a slow capacity fading per cycles. The capacity stabilized at 800 mA h  $g_{sulfur}^{-1}$  and 600 mA h  $g_{sulfur}^{-1}$  up to 150 charging-discharging cycles for OLC-S-2 and OLC-S-3 hybrids, respectively. Cathodes containing carbon onions/sulfur hybrids, where the melt diffusion route was followed, demonstrated severe capacity fading within 50 cycles and remained with only 400 mA h  ${
m g_{sulfur}}^{-1}$  at the end of 150 cycles of charge– discharge. The electrode prepared from sulfur nanoparticles (SNPs) with 10 mass% conductive additive showed a low specific capacity in the initial cycles. After 20 cycles, the specific capacity was stabilized at 200 mA h g<sub>sulfur</sub><sup>-1</sup> for the following 150 cycles. The relatively low specific capacity values are because of an insufficient amount of conducting carbon with high surface area around the sulfur nanoparticles.

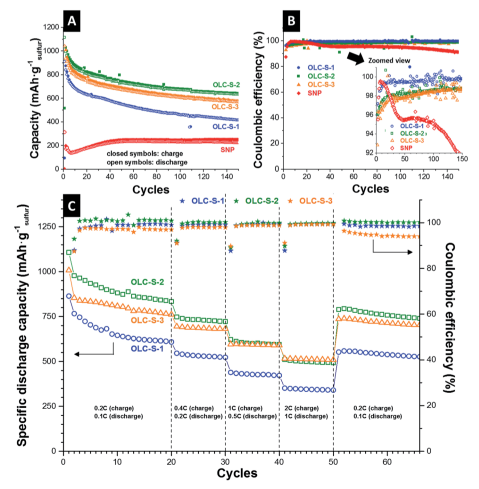


Fig. 5 Cycling performance (0.2C charge, 0.1C discharge) for the (A) specific capacity and (B) Coulombic efficiency (inset shows the zoomed range of 92–100%). (C) Rate capability study of the carbon onion/sulfur hybrid and sulfur nanoparticle (SNP) cathodes.

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 Table 4
 Reported values for selected carbon/sulfur electrodes in comparison to this work. In all reports, metallic Li was used as the anode. Used abbreviations: PC = polyethylene glycol dimethyl carbonate; PC = polyethylene glycol dimethyl carbonate; PC = polyethylene glycol dimethyl
  $ether; \ \mathsf{PYR}_{14} = 1-butyl-1-methylpyrrolidinium; \ \mathsf{TEGDME} = \mathsf{tetraethylene} \ \mathsf{glycol} \ \mathsf{dimethyl} \ \mathsf{ether}; \ (\mathsf{w} = \mathsf{mass}; \ \mathsf{v} = \mathsf{volume})$ 

| Surface area   Surface area   Sulface lading   Separator   Electrolyte   Separator   Electrolyte   Separator   Electrolyte   Separator   Separator |   |                              |                                    |                                | '  |                      |                       |                             |                     |  |      |
|--|---|------------------------------|------------------------------------|--------------------------------|--|----------------------|-----------------------|-----------------------------|---------------------|--|------|
| Surface area         Sulfut loading         Ekectrolyte         Cycle 50         Cycle 50         Cycle 50           ca. 400         2–3         Polypotefin         1 M LiffSI in 1:1 (v/v) DME/DOL + ca. 950         ca. 700         ca. 300           ca. 65         1.5         Polypotefine         1 M LiffSI in 1:1 (v/v) DME/DOL + ca. 950         ca. 500           ca. 800         0.81         Unknown         0.5 M LiNO <sub>3</sub> ca. 107         ca. 770         ca. 700           ca. 107         Unknown         Celgard         0.5 M LiNO <sub>3</sub> ca. 110         ca. 100         ca. 270           ca. 150         Unknown         Celgard and of the LiffSI in 1:1 (v/v) DME/DOL + ca. 1100         ca. 100         ca. 100           ca. 150         Unknown         Celgard and the LiffSI in 1:1 (v/v) DME/DOL + ca. 290         ca. 550         ca. 750           ca. 156         0.45         Unknown         I M LiffSI in 1:1 (v/v) DME/DOL + ca. 290         ca. 100           ca. 156         0.25         Celgard         1 M LiffSI in 1:1 (v/v) DME/DOL + ca. 290         ca. 550           ca. 156         0.25         Celgard         1 M LiffSI in 1:1 (v/v) DME/DOL + ca. 290         ca. 550           ca. 500         ca. 500         ca. 500         ca. 800           ca. 51000         ca  |   | e<br>i                       | :                                  |                                |  | Specific cap         | oacity (mA h          | $\mathrm{g_{sulfur}}^{-1})$ | :                   |  |      |
| ca. 400         2-3         Polyolefin         1 M LITESI in 1:1 (w) DME/DOL + ca. 950         ca. 500         ca. 730           ca. 65         1.5         Polypropylene         1 M LIPE, PCIEC/DEC 1:4:5 (w)         ca. 600         ca. 500         ca. 500           ca. 800         0.81         Unknown         0.5 M LINO, CF, in 8:2 (w) DME/DOL + ca. 1100         ca. 1000         ca. 500         ca. 500           ca. 150         Unknown         Celgard and ca. 1 M LITESI in 1:1 (w) DME/DOL + ca. 1100         ca. 1000         ca. 1000         ca. 1000           ca. 150         Unknown         Polypropylene of LITESI in 1:1 (w) DME/DOL + ca. 250         ca. 1500         ca. 1500         ca. 500         ca. 500           ca. 156         0.45         Not mentioned of M LITESI in 1:1 (w) DME/DOL + ca. 250         ca. 500         ca. 500           ca. 1566         0.25         Celgard of M LITESI in 1:1 (w) DME/DOL + ca. 700         ca. 500         ca. 500           ca. 1566         0.25         Unknown         1 M LITESI in 1:1 (w) DME/DOL + ca. 700         ca. 500           ca. 300         ca. 300         ca. 400         ca. 800         ca. 800           ca. 378         ca. 300         ca. 400         ca. 800           ca. 379         ca. 250         ca. 250         ca. 250  | Carbon for cathode                        | Surface area $(m^2 g^{-1})$  | Sulfur loading $({ m mg~cm}^{-2})$ | Separator                      | Electrolyte  | Cycle 5              | Cycle 50              | Cycle 100                   | Potential<br>window | C-Rate                                   | Ref. |
| ca. 65         1.5         Polypropylene         1 M LiPE, PC/EC/DEC 1:4:5 (v/v) DME/ or 6.00         ca. 600         ca. 500           ca. 800         0.81         Unknown         0.5 M LiSO, CF, in 8:2 (v/v) DME/ or 2.100         ca. 770         ca. 700           -         Unknown         Celgard         0.5 M LiNO; in 1:1 (v/v) DME/DOL+         ca. 1100         ca. 1000           -         Unknown         Celgard and of 3 M LiTFSI in 1:1 (v/v) DME/DOL+         ca. 1300         ca. 1000         ca. 1000           ca. 150         1.36         Unknown         Polypropylene         1 M LiTFSI in 1:1 (v/v) DME/DOL         ca. 1300         ca. 1300           ca. 1305         Unknown         Polypropylene         1 M LiTFSI in 1:1 (v/v) DME/DOL         ca. 1000         ca. 500           ca. 1306         0.25         Celgard         1 M LiTFSI in 1:1 (v/v) DME/DOL         ca. 1000         ca. 500           ca. 300         0.4-0.6         Unknown         1 M LiTFSI in 1:1 (v/v) DME/DOL         ca. 780         ca. 580           ca. 300         0.4-0.6         Unknown         1 M LiTFSI in 1:1 (v/v) DME/DOL         ca. 780         ca. 580           ca. 300         0.4-0.6         Unknown         Celgard         1 M LiTFSI in 1:1 (v/v) DME/DOL         ca. 1000         ca. 590           <  | This work                                 | ca. 400                      | 2–3                                | Polyolefin<br>separators       | 1 M LiTFSI in 1 : 1 (v/v) DME/DOL + $0.25$ M LiNO <sub>3</sub>                         | ca. 950<br>(OLC-2)   | ca. 730<br>(OLC-2)    | ca. 670<br>(OLC-2)          | 1.8-2.6 V           | 0.1C<br>(discharge),<br>0.2C (charge)    | 1    |
| ca. 800         0.81         Unknown         0.5 M LiSO <sub>3</sub> CF <sub>3</sub> in 8 : 2 (w) DME/         cz. 770         cz. 700           ca. 107         Unknown         Celgard         0.5 M LiTSSI in 1: 1 (w) DME/DOL         cz. 1100         cz. 1000           ca. 150         c. clgard and glass fiber mat         0.25 M LiNO <sub>3</sub> cz. 170         cz. 1100         cz. 1000           ca. 150         1.36         Unknown         Polypropylen         1 M LiTSSI in 1: 1 (w) DME/DOL         cz. 150         cz. 150           ca. 1566         0.25         Celgard         1 M LiTSSI in 1: 1 (w/) DME/DOL         cz. 790         cz. 650           ca. 1566         0.25         Celgard         1 M LiTSSI in 1: 1 (w/) DME/DOL         cz. 790         cz. 650           ca. 1566         0.25         Celgard         1 M LiTSSI in 1: 1 (w/) DME/DOL         cz. 780         cz. 500           ca. 300         Unknown         1 M LiTSSI in 1: 1 (w/) DME/DOL         cz. 780         cz. 580           ca. 578         Unknown         Celgard         1 M LiTSSI in 1: 1 (w/) DME/DOL         cz. 780         cz. 580           ca. 578         cz. 580         cz. 580         cz. 580         cz. 580         cz. 580           ca. 578         cz. 580         cz. 580         cz. 580         cz. 580 <td>Acetylene black</td> <td>ca. 65</td> <td>1.5</td> <td>Polypropylene<br/>separator</td> <td>1 M LiPF<sub>6</sub> PC/EC/DEC 1:4:5 (v/v)</td> <td>ca. 600</td> <td>ca. 500</td> <td>I</td> <td>1.8-2.8 V</td> <td>0.02C</td> <td>45</td>  | Acetylene black                           | ca. 65                       | 1.5                                | Polypropylene<br>separator     | 1 M LiPF <sub>6</sub> PC/EC/DEC 1:4:5 (v/v)  | ca. 600              | ca. 500               | I                           | 1.8-2.8 V           | 0.02C                                    | 45   |
| az. 107         Unknown         Celgard         0.5 M LiTPSI in 1:1 (v/v) DME/DOL + ca 1100         ca 1000           —         Unknown         Celgard and glass fiber mat glass fiber mat ca. 150         1.36         Unknown         Celgard and 1 M LiTPSI in 1:1 (v/v) DME/DOL + ca. 250         ca. 750         ca. 750           ca. 150         1.36         Unknown         Polypropylene of 1 M LiTPSI in 1:1 (v/v) DME/DOL + ca. 790         ca. 1100         ca. 1100           2445         0.45         Nor mentioned of 1 M LiTPSI in 1:1 (v/v) DME/DOL + ca. 700         ca. 1500         ca. 650           ca. 300         0.25         Celgard         1 M LiTPSI in 1:1 (v/v) DME/DOL + ca. 700         ca. 650           ca. 300         Unknown         1 M LiTPSI in 1:1 (v/v) DME/DOL + ca. 780         ca. 800           ca. 300         Unknown         1 M LiTPSI in 1:1 (v/v) DME/DOL + ca. 780         ca. 800           ca. 578         Unknown         Unknown         1 M LiTPSI in 1:1 (v/v) DME/DOL + ca. 780         ca. 580           ca. 578         Unknown         Unknown         1 M LiTPSI in 1:1 (v/v) DME/DOL + ca. 950         ca. 950           ca. 578         Celgard         0.35 M LiTPSI in 1:1 (v/v) DME/DOL + ca. 1550         ca. 650         ca. 650           ca. 578         Celgard         0.35 M LiTPSI in 1:1 (v/v) DME/DOL + ca. 1550         ca   | Ketjen black                              | ca. 800                      | 0.81                               | Unknown                        | 0.5 M LiSO <sub>3</sub> CF <sub>3</sub> in $8:2$ (v/v) DME/<br>DOL + imidazolium salts | ca. 770              | ca. 700               | ca. 600                     | 1.8-2.8 V           | 0.5C                                     | 46   |
|  | Multi-walled carbon<br>nanotube           | ca. 107                      | Unknown                            | Celgard                        | 0.5 M LiTFSI in 1 : 1 (v/v) DME/DOL + 0.5 M LiNO <sub>3</sub>                          |                      | ca. 1000              | ca. 950                     | 1.7-2.7 V           | 0.18C                                    | 35   |
| a. 136         Unknown         1 M LiTFSI in 1:1 (v/v) DME/DOL         ca 1250         ca 1100           a. 1395         Unknown         Polypropylene separator         1 M LiTFSI in 1:1 (v/v) DME/DOL         ca 1300         ca 650           2445         0.45         Not mentioned         1 M LiTFSI in 1:1 (v/v) DME/DOL         ca 500         ca 650           2a. 1566         0.25         Celgard         1 M LiTFSI in 5: 40 (v/v) DME/DOL         ca 500         ca 300           -         0.4-0.6         Unknown         1 M LiTFSI in 1:1 (v/v) DME/DOL         ca 500         ca 800           -         Unknown         Celgard         LiTFSI in 1:1 (v/v) DME/DOL         ca 500         ca 900           -         Unknown         Unknown         1 M LiTFSI in 1:1 (v/v) DME/DOL         ca 690         ca 800           2000         6.5         Celgard         1 M LiTFSI in 1:1 (v/v) DME/DOL         ca 690         ca 800           2000         6.5         Celgard         1 M LiTFSI in 1:1 (v/v) DME/DOL         ca 650         ca 600           2000         6.5         Celgard         1 M LiTFSI in 1:1 (v/v) DME/DOL         ca 650         ca 600           2000         2-3         Polyolefin         1 M LiTFSI in 1:1 (v/v) DME/DOL         ca 650         ca 600 <td>Vertically aligned CNTs</td> <td>I</td> <td>Unknown</td> <td>Celgard and<br/>glass fiber mat</td> <td>1 M LiTFSI in 1 : 1 (v/v) DME/DOL + 0.25 M LiNO<sub>3</sub></td> <td>ca. 850</td> <td>ca. 750<br/>(cycle 40)</td> <td>1</td> <td>1-3 V</td> <td>0.08C</td> <td>47</td>  | Vertically aligned CNTs                   | I                            | Unknown                            | Celgard and<br>glass fiber mat | 1 M LiTFSI in 1 : 1 (v/v) DME/DOL + 0.25 M LiNO <sub>3</sub>                           | ca. 850              | ca. 750<br>(cycle 40) | 1                           | 1-3 V               | 0.08C                                    | 47   |
| a. 1395         Unknown         Polypropylene         1 M LiTFSI in 1:1 (v/v) DME/DOL         ca. 790         ca. 650           2445         0.45         Not mentioned separator         0.1 M LiNO <sub>3</sub> ca. 100         ca. 650         ca. 600           2445         0.25         Celgard         1 M LiTFSI in 55:40 (v/v) DME/DOL         ca. 500         ca. 300           -         0.4-0.6         Unknown         1 M LiTFSI in 1:1 (v/v) DME/DOL         ca. 500         ca. 300           -         0.4-0.6         Unknown         1 M LiTFSI in 1:1 (v/v) DME/DOL         ca. 780         ca. 800           -         Unknown         Celgard         LiTFSI in PYR <sub>1</sub> /TFSI/PEGDME         ca. 780         ca. 580           -         Unknown         Unknown         1 M LiTFSI in 1:1 (v/v) DME/DOL         ca. 690         ca. 500           2000         6.5         Celgard         1 M LiTFSI in 1:1 (v/v) DME/DOL         ca. 600         ca. 800           2000         6.5         Celgard         1 M LiTFSI in 1:1 (v/v) DME/DOL         ca. 650         ca. 600           2-3         Polyolefin         1 M LiTFSI in 1:1 (v/v) DME/DOL         ca. 650         ca. 600           ca. 200         2-3         Polyolefin         1 M LiTFSI in 1:1 (v/v) DME/DOL         ca. 650  | Small CNTs confined inside a large CNT    | ca. 150                      | 1.36                               | Unknown                        | 1 M LiTFSI in 1 : 1 (v/v) DME/DOL  | ca. 1250             | ca. 1100              | ca. 1000                    | 1.5-3 V             | 0.1C                                     | 43   |
| 2445         0.45         Not mentioned         1 M LiTFSI in 1:1 (v/v) DME/DOL         ca. 1000         ca. 900           ca. 1566         0.25         Celgard         1 M LiTFSI in 55:40 (v/v) DME/DOL         ca. 500         ca. 300           ca. 300         Unknown         1 M LiTFSI in 1:1 (v/v) DME/DOL         ca. 780         ca. 800           ca. 300         Unknown         1 M LiTFSI in 1:1 (v/v) DME/DOL         ca. 780         ca. 800           ca. 578         Unknown         LiTFSI in PYR <sub>14</sub> TFSI/PEGDME (1:1)         ca. 900         ca. 520           ca. 578         Unknown         LiTFSI in TEGDME         ca. 690         ca. 520           ca. 578         Unknown         1 M LiTFSI in 1:1 (v/v) DME/DOL         ca. 950         ca. 800           2000         6.5         Celgard         1 M LiTFSI in 1:1 (v/v) DME/DOL         ca. 1000         ca. 600           ca. 1300         2-3         Polyolefin         1 M LiTFSI in 1:1 (v/v) DME/DOL         ca. 650         ca. 600           ca. 2000         2-3         Polyolefin         1 M LiTFSI in 1:0 (v/v) DME/DOL         ca. 650         ca. 600           ca. 2000         2-3         Polyolefin         1 M LiTFSI in 1:0 (v/v) DME/DOL         ca. 650         ca. 600           ca. 2000         0.8   | Carbon aerogel                            | ca. 1395                     | Unknown                            | Polypropylene<br>separator     | 1 M LiTFSI in 1 : 1 (v/v) DME/DOL + 0.1 M LiNO <sub>3</sub>                            | ca. 790              | ca. 650               | ca. 600                     | 1.8-2.8 V           | 0.1C                                     | 40   |
| ca. 1566         0.25         Celgard         1 M LiTFSI in 55 : 40 (v/v) DME/DOL         ca. 500         ca. 300           -         0.4-0.6         Unknown         1 M LiTFSI in 1:1 (v/v) DME/DOL         ca. 780         ca. 300           ca. 300         ca. 300         Unknown         1 M LiTFSI in 1:1 (v/v) DME/DOL         ca. 780         ca. 580           -         Unknown         Celgard         LiTFSI in PYR <sub>14</sub> TFSI/PEGDME (1:1         ca. 900         ca. 900         ca. 500           2000         6.5         Celgard         1 M LiTFSI in TEGDME         ca. 950         ca. 950         ca. 800           2000         6.5         Celgard         1 M LiTFSI in 1:1 (v/v) DME/DOL         ca. 950         ca. 800           2000         6.5         Celgard         1 M LiTFSI in 1:1 (v/v) DME/DOL         ca. 950         ca. 800           2000         2-3         Polyolefin         1 M LiTFSI in 1:1 (v/v) DME/DOL         ca. 650         ca. 650           ca. 900         2-3         Polyolefin         1 M LiTFSI in 1:1 (v/v) DME/DOL         ca. 650         ca. 600           ca. 2000         2-3         Polyolefin         1 M LiTFSI in TEGDME + 0.15 M         ca. 650         ca. 600           ca. 2000         0.8         Celgard         1 M LiTFSI in TEGD  | Ordered mesoporous carbon                 | 2445                         | 0.45                               | Not mentioned                  | 1 M LiTFSI in 1 : 1 (v/v) DME/DOL  | ca. 1000             | ca. 900               | ca. 800                     | 1.5-3 V             | 1C                                       | 48   |
| —         0.4-0.6         Unknown         1 M LiTFSI in 1:1 (v/v) DME/DOL + (az 1000) (az 800 0.14 M LiNO₃           aa. 300         Unknown         1 M LiTFSI in 1:1 (v/v) DME/DOL (az 780 ca 580 0.14 M LiNo₃)         ca. 580 ca 580 ca 580 ca 580 ca 580 ca 580 ca 578           —         Unknown         Unknown         1 M LiTFSI in TEGDME (1:1 ca 900 ca 520 ca 800 ca 578 0.14 M LiTFSI in 1:1 (v/v) DME/DOL (az 950 ca 800 ca 520 ca 800 ca 1300 ca 1300 ca 1300 ca 1400 ca   | Ordered mesoporous carbon (KOH activated) | ca. 1566                     | 0.25                               | Celgard                        | 1 M LiTFSI in 55 : 40 (v/v) DME/DOL  | ca. 500<br>(51.5% S) | са. 300               | 1                           | 1-3.6 V             | 1.5C                                     | 37   |
| ca. 300         Unknown         Unknown         1 M LiTFSI in 1:1 (v/v) DME/DOL         ca. 780         ca. 580           —         Unknown         LiTFSI in PYR14TFSI/PEGDME (1:1 a. 900         ca. 900         ca. 900         ca. 520           2000         6.5         Celgard         1 M LiTFSI in 1:1 (v/v) DME/DOL         ca. 690         ca. 520           2000         6.5         Celgard         0.35 M LiTFSI in 1:1 (v/v) DME/DOL         ca. 950         ca. 400           ca. 1300         2-3         Polyolefin         1 M LiTFSI in 1:1 (v/v) DME/DOL         ca. 1650         ca. 400           ca. 900         2-3         Polyolefin         1 M LiTFSI in 1:1 (v/v) DME/DOL         ca. 650         ca. 600           ca. 2000         0.8         Celgard         1 M LiTFSI in TEGDME + 0.15 M         ca. 650         ca. 600   | Sulphur-TiO <sub>2</sub> yolk-shell       | 1                            | 0.4-0.6                            | Unknown                        | 1 M LiTFSI in 1 : 1 (v/v) DME/DOL + 0.14 M LiNO <sub>3</sub>                           | ca. 1000             |                       | са. 900                     | 1.7-2.6 V           | 0.5C                                     | 49   |
| oxide         —         Unknown         LiTFSI in PYR14TFSI/PEGDME (1:1 a. 900         ca. 900         ca. 900           d sulfur         —         Unknown         Unknown         1 M LiTFSI in TEGDME         ca. 690         ca. 520           wich         ca. 578         Unknown         Celgard         1 M LiTFSI in 1:1 (v/v) DME/DOL         ca. 950         ca. 800           ts         creloth         2000         6.5         Celgard         0.35 M LiTFSI in 1:1 (v/v) DME/DOL         ca. 1000         ca. 950           t         ca. 1300         2-3         Polyolefin         1 M LiTFSI in 1:1 (v/v) DME/DOL         ca. 1650         ca. 400           ca. 900         2-3         Polyolefin         1 M LiTFSI in 1:1 (v/v) DME/DOL         ca. 650         ca. 600           separator         0.25 M LiNO <sub>3</sub> 1 M LiTFSI in TEGDME + 0.15 M         ca. 650         ca. 600           ca. 2000         0.8         Celgard         1 M LiTFSI in TEGDME + 0.15 M         ca. 1100         ca. 1000  |   |                              |                                    | Unknown                        | 1 M LiTFSI in 1 : 1 (v/v) DME/DOL  | ca. 780              | ca. 580               | 1                           | 1.5-3.2 V           | 1C                                       | 20   |
| d sulfur         —         Unknown         Unknown         1 M LiTFSI in TEGDME         ca. 690         ca. 520           wich         ca. 578         Unknown         Celgard         1 M LiTFSI in 1:1 (v/v) DME/DOL         ca. 950         ca. 800           er cloth         2000         6.5         Celgard         0.35 M LiTFSI in 1:1 (v/v) DME/DOL         ca. 1000         ca. 950           er cloth         2-3         Polyolefin         1 M LiTFSI in 1:1 (v/v) DME/DOL         ca. 1650         ca. 400           separator         0.25 M LiNO <sub>3</sub> A LiNO <sub>3</sub> ca. 650         ca. 650         ca. 600           ca. 2000         0.8         Celgard         1 M LiTFSI in TEGDME + 0.15 M         ca. 1100         ca. 1000  | Reduced graphene oxide                    | I                            | Unknown                            | Celgard                        | Litesi in $PYR_{14}TFSI/PEGDME$ (1 : 1 w/w)  | ca. 900              | ca. 900               | I                           | 1-3 V               | 0.02C (3 cycles) then 0.1C for 47 cycles | 51   |
| tes er cloth 2000 6.5 Celgard 0.35 M LiTFSI in 1:1 (v/v) DME/DOL ca. 1000 ca. 950 1 ca. 1300 2-3 Polyolefin 1 M LiTFSI in 1:1 (v/v) DME/DOL + ca. 1650 ca. 400 2-3 Polyolefin 1 M LiTFSI in 1:1 (v/v) DME/DOL + ca. 1650 ca. 600 2-3 Polyolefin 1 M LiTFSI in 1:1 (v/v) DME/DOL ca. 650 ca. 600 2-3 Robinstaror ca. 2000 0.8 Celgard 1 M LiTFSI in TEGDME + 0.15 M ca. 1100 ca. 1000 LiNO <sub>3</sub>   | Graphene-enveloped sulfur                 | — 53<br>— 24<br>— 27<br>— 27 | Unknown                            | Unknown                        | 1 M LITESI in TEGDME   | ca. 690              | ca. 520               | 1                           | 1.5-3 V             | 0.2C                                     | 17   |
| er cloth 2000 6.5 Celgard 0.35 M LiTFSI in 1:1 (v/v) DME/DOL ca. 1000 ca. 950 +0.29 M LiNO <sub>3</sub> ca. 1300 2-3 Polyolefin 1 M LiTFSI in 1:1 (v/v) DME/DOL + ca. 1650 ca. 400 separator 0.25 M LiNO <sub>3</sub> ca. 900 2-3 Polyolefin 1 M LiTFSI in 1:1 (v/v) DME/DOL ca. 650 ca. 600 separator ca. 2000 0.8 Celgard 1 M LiTFSI in TEGDME + 0.15 M ca. 1100 ca. 1000 LiNO <sub>3</sub>  | type graphene sheets                      | cu. 3/8                      | UIIKIIOWII                         | Ceigaid                        | 1 M LIIFSI III 1 : 1 (V/V) DME/DOL   |                      |                       | I                           | v c-1               | 0.10                                     | 76   |
| ca. 1300 2–3 Polyolefin 1 M LiTFSI in 1:1 (v/v) DME/DOL + ca. 1650 ca. 400 separator 0.25 M LiNO <sub>3</sub> ca. 900 2–3 Polyolefin 1 M LiTFSI in 1:1 (v/v) DME/DOL ca. 650 ca. 600 separator  ca. 2000 0.8 Celgard 1 M LiTFSI in TEGDME + 0.15 M ca. 1100 ca. 1000 LiNO <sub>3</sub>   | Activated carbon fiber cloth              | 2000                         | 6.5                                | Celgard                        | 0.35 M LiTFSI in 1 : 1 (v/v) DME/DOL + 0.29 M LiNO <sub>3</sub>                        |                      |                       |                             | 1.7-2.5 V           | 0.09C                                    | 24   |
| ca. 900       2-3       Polyolefin       1 M LiTFSI in 1:1 (v/v) DME/DOL       ca. 650       ca. 600         separator       separator       1 M LiTFSI in TEGDME + 0.15 M       ca. 1100       ca. 1000   | Inverse opal carbon                       | ca. 1300                     | 2-3                                | Polyolefin<br>separator        | 1 M LiTFSI in 1 : 1 (v/v) DME/DOL + 0.25 M LiNO <sub>3</sub>                           | ca. 1650             | ca. 400               |                             | 1.5-2.8 V           | 0.1C                                     | 26   |
| $\it ca.$ 2000 0.8 Celgard 1 M LİTFSI in TEGDME + 0.15 M $\it ca.$ 1100 $\it ca.$ 1000 LİNO <sub>3</sub>   | 3D gyroidal carbon                        | ca. 900                      | 2-3                                | Polyolefin<br>separator        | 1 M LiTFSI in $1:1$ (v/v) DME/DOL  | ca. 650              | <i>ca.</i> 600        | ca. 520                     | 1.5-2.8 V           | 0.1C (discharge), 0.2C (charge)          | 21   |
|  | 3D gyroidal carbon<br>(activated)         | ca. 2000                     | 0.8                                | Celgard                        | 1 M LiTFSI in TEGDME + 0.15 M LiNO $_3$  | ca. 1100             | ca. 1000              | ca. 830                     | 1.5-2.8 V           | 0.1C                                     | 22   |

Additionally, a comparison of specific capacities from the literature with different types of carbon substrates can be found in Table 4.

The capacity loss in the initial cycles can be attributed to the irreversible diffusion of polysulfides possibly from the dissociative sulfur on the surface of OLC particles. This irreversible reaction leads to slightly lower Coulombic efficiency close to 97-99% in reversible (Fig. 5B) or irreversible mode (Fig. 5C). For comparison, the SNP electrodes demonstrated a Coulombic efficiency below 95% after 30 cycles and after 100 cycles, the Coulombic efficiency dropped below 90%. This behavior can be explained by a lack of structural integrity and loss of electrical connectivity between the additive and sulfur nanoparticles. The relatively high specific capacity in the beginning and the slow capacity fading of OLC-S hybrids can be attributed to the benefits of the enhanced surface area of carbon onions and the accessible interparticle space. The interconnecting feature of carbon onion agglomerates provides a high electrical conductivity and is beneficial to ensure accessibility of the active material by the electrolyte and the Li<sup>+</sup> ions with prolonged cycling.

The rate capability tests of the carbon onion-sulfur hybrids were carried out in 2032 coin cells at multiple C-rates (Fig. 5C). The respective charge and discharge protocol is explained in the experimental part. After 20 cycles at a low rate (0.2C charge and 0.1C discharge) and 30 cycles at a high rate (up to 2C charge and 1C discharge) the specific capacity was recovered close to the level of the initial 20<sup>th</sup> cycles after multiple irreversible charging-discharging fluctuations. From the irreversible capacity retention ability, it can be concluded that the carbon onion-sulfur composite nanostructures were not ruptured and the sulfur distribution was restored even after such stringent charging-discharging conditions.

Finally, we investigated the long-term cycle performance of the OLC-S hybrids at very high C-rates, for instance, 2C (equivalent to 3350 mA g<sup>-1</sup> with respect to sulfur) for charging and 1C (equivalent to 1675 mA g<sup>-1</sup> with respect to sulfur) for discharging (Fig. 6). Among all three OLC-S hybrids, the onion-like carbon hybrid fabricated by the in situ routine followed by thermal annealing demonstrated an initial specific discharge capacity of 800 mA h g<sup>-1</sup>. After 25 cycles, the battery showed a very high stability over 500 cycles with a slow rate of capacity fading (Fig. 6). For the latter, we measured a remaining capacity of 400 mA h g<sup>-1</sup> after 500 cycles. The other in situ hybrid

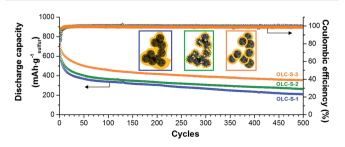


Fig. 6 Long term cycle stability at high C-rates (charging at 2C/discharging at 1C) study of the carbon onion/sulfur hybrids.

(i.e., without annealing) followed a similar trend, yielding 300 mA h  $g^{-1}$  after 500 cycles. The OLC-S hybrid fabricated *via* the melt diffusion strategy had been showing poorest performances amongst all three hybrids due to random distribution of sulfur in the hybrid. The better performance of the OLC-S-3 at higher rates relative to other hybrids was also observed in rate capability experiments (Fig. 5C). The slower capacity fading at such high specific currents is mainly ascribed to the excellent electrical conductivity of the OLC-S hybrids, which are capable of satisfying the requirement for such fast discharge and charge during repetitive cycling. 41,43 We attribute the better performance at higher charge/discharge rates of OLC-S-3 to the much more integrated hybrid nanostructure by melting of the in situ deposited sulfur on the surface of OLC. Our findings are also in good agreement with the published reports by Nazar et al. performed at high C rates (cut-off voltage limit 1.5-3 V vs. Li/Li<sup>+</sup>) exploiting bimodal porous carbon possessing over 2000 m<sup>2</sup> g<sup>-1</sup> surface area.44

#### 4. Conclusions

Our work introduces carbon onions for the fabrication of carbon/sulfur hybrid cathodes for lithium-sulfur batteries with sulfur loading up to 68 mass% and without any further addition of a conducting aid such as carbon black. Three different approaches were explored: (1) melt diffusion, (2) in situ generation of sulfur nanoparticles on the surface of onion-like carbon, and (3) in situ generation of sulfur nanoparticles on the surface of onion-like carbon followed by thermal annealing.

The electrochemical performance of these hybrid materials has been tested by electrochemical characterization. Carbon onion-sulfur hybrids prepared via the in situ approach (OLC-S-2 and OLC-S-3) had an optimum interaction between the hybrid components exhibiting a specific capacity of ca. 700 mA h g<sup>-1</sup> even after 150 charge-discharge cycles at 167 mA g<sup>-1</sup> specific current (corresponds to 0.1C discharge rate). In addition, the capacity restoration ability of OLC-S in situ hybrids right after running with a high current density of 1675 mA g<sup>-1</sup> (corresponds to 1C discharge rate) demonstrated a specific capacity of 740 mA h  $g^{-1}$  close to its earlier capacity at 0.1C rate (800 mA h g<sup>-1</sup>) when the current density was reduced to 167 mA g<sup>-1</sup> (corresponds to 0.1C discharge rate). This hybrid material showed enhanced performance compared to the other OLC-S hybrids studied at slow discharge rates. In addition, these hybrids demonstrate good cycle performances operated with very high C-rates (1C for discharge and 2C for charge). Amongst them, the hybrid fabricated via in situ generation of sulfur nanoparticles on the surface of onion-like carbon followed by thermal annealing (OLC-S-3) has been found to be superior  $(800 \text{ mA h g}^{-1} \text{ at cycle 1, } 600 \text{ mA h g}^{-1} \text{ at cycle 20, and } 400 \text{ mA})$ h  $g^{-1}$  at cycle 500).

The high electrochemical stability and attractive rate handling can be attributed to carbon onion aggregates which contribute a high electronic conductivity and provide an exclusively outer surface, which enables the synergy of high sulfur mass loading and high percentage of electrochemically active sulfur.

## Acknowledgements

V. P., S. C., and M. Z. gratefully acknowledge funding from the German Federal Ministry for Research and Education (BMBF) in support of the nanoEES<sup>3D</sup> project (award number 03EK3013) and L. B. in support of the Mechanocarb project (award number 03SF0498), both as part of the strategic funding initiative energy storage framework. This work was supported by the CREATe-Network Project, Horizon 2020 of the European Commission (RISE Project No. 644013). The authors thank Prof. Eduard Arzt (INM) for his continuing support.

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