Optimized synthesis of ultrahigh-surface-area and oxygen-doped carbon nanobelts for high cycle-stability lithium-sulfur batteries

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Abstract:
Hierarchical clews of carbon nanobelts (CsCNBs) with ultrahigh specific surface area (2300 m² g⁻¹) and large pore volume (up to 1.29 cm³ g⁻¹) has been successfully fabricated through carbonization and KOH activation of phenolic resin based nanobelts. The product possesses hierarchically porous structure, three-dimensional conductive network framework, and polar oxygen-rich groups, which are very befitting to load sulfur leading to excellent cycling stability of lithium-sulfur batteries. The composites of CsCNBs/sulfur exhibit an ultrahigh initial discharge capacity of 1245 mA h g⁻¹ and ultralow capacity decay rate as low as 0.162% per cycle after 200 cycles at 0.1 C. Even at high current rate of 4 C, the cells still display a high initial discharge capacity (621 mA h g⁻¹) and ultralow capacity decay rate (only 0.039% per cycle) after 1000 cycles. These encouraging results indicate that polar oxygen-containing functional groups are important for improving the electrochemical performance of carbons. The oxygen-doped carbon nanobelts have excellent energy storage potential in the field of energy storage.
Introduction

Lithium batteries are widely used in energy storage power systems such as hydropower, firepower, wind power and solar power plants, electric bicycles, aerospace and many other fields. [1-2]. However, the traditional Lithium batteries cannot satisfy the high requirement of more integrated devices with high energy-consumption as well as the environmental protection concept [3-5]. Lithium-sulfur batteries (LiSBs) are notable for their ultrahigh theoretical energy density (2600 W h kg\(^{-1}\)) that doubles the energy density of lithium-ion batteries [6-8], cheap raw materials and little pollution to the environment [9-10]. Therefore, LiSBs are very promising batteries for energy storage applications. However, there are three major drawbacks for LiSBs which hamper their development in energy storage fields. First, elemental sulfur has poor electrical conductivity, and final reaction products (Li\(_2\)S\(_2\) and Li\(_2\)S) are also electronic insulators, which are not beneficial to the high-rate performance of batteries [11-14]. Second, lithium polysulfide compounds (Li\(_2\)S\(_8\), Li\(_2\)S\(_6\) and Li\(_2\)S\(_4\)) are easily soluble in the electrolyte, resulting in a shuttle effect which easily destroys the electrolyte interface film (SEI film) [12, 15]. Third, sulfur will undergo a volumetric expansion of ~80% during charging and discharging process, which causes the pulverization of the electrode materials and thus damages the batteries [16-17].

In recent ten years, in order to solve the problems in LiSBs, researchers have conducted a lot of research on the modification of composite cathode materials. One of the most important, porous carbons have received great attention and research for its high electrical conductivity, high specific surface area (SSA) [18], high porosity, high chemical and mechanical stability, and ease of preparation. According to its structure, porous carbon is divided into one-dimensional materials (such as carbon nanotubes), two-dimensional materials (such as graphene) and three-dimensional materials (such as hollow carbon spheres). In a variety of porous carbon, hierarchical porous carbons (HPCs) have been more widely researched due to the co-existence of micropore (< 2 nm),
mesopore (2-50 nm), and macropore (> 50 nm) [19-22]. Hierarchical pores can make a synergistic effect of in LiSBs. Micropores provide anchoring sites for binding lithium polysulfide. Mesopores are commonly used to enhance the physical properties of microporous or macroporous networks. The smaller mesopores along with the micropores store sulfur and adsorb polysulfides, further slowing down the decay of capacity. The larger mesopores along with macropores promote charge transport and ensure electrolyte infiltration, which thereby improves sulfur utilization of active materials. The macroporous network serves to contain and transport the electrolyte to ensure good electrolyte impregnation. The physisorption characteristic of the pores can effectively improve the cycling stability of LiSBs.

In addition, Cui groups [15, 23] studied the interaction and binding energies of various functional groups (such as carbon groups and oxygen-rich groups) and lithium sulfide (LiS) species using density functional theory based on the vinyl polymer. As a result, carbon-based groups, such as alkanes, demonstrate lower binding energy with LiS species (0.30 eV) and oxygen-rich groups containing esters, amides, ketones and ethers, demonstrate higher binding energies between 1.01 and 1.26 eV. Oxygen-containing functional groups have been found to improve the combination with lithium polysulfides. In short, the chemisorption between LiS species and oxygen-rich groups has a potential improvement effect on the cycle-stability of LiSBs. Cheng and colleagues [24] found that the strong chemical combination between sulfur and oxygen-containing functional groups (mainly -OH and C=O) can significantly reduce the dissolution/diffusion of lithium polysulfides on reduced graphene oxide. Further, the cycle stability of the lithium-sulfur battery is improved.

KOH activation is a very promising method for the preparation of HPCs, because of its low activation temperature, high carbon yield, well-defined pore size distribution and ultrahigh specific surface area [18]. Using biomass waste (soybean residue) as precursor, Zhou et al. [25] successfully prepared a new type of honeycomb nitrogen-oxygen co-doped porous carbon using KOH activation.
The derived porous carbon has a special multi-stage pore structure, a large specific surface area (SSA) (2690.3 m$^2$ g$^{-1}$), a high pore volume (1.34 cm$^3$ g$^{-1}$), and a suitable N, O co-doping. The carbon/sulfur composite [26-31] prepared by the porous carbon (sulfur content of 64.5%) exhibits an excellent cycle stability in LiSBs, and the capacity decay rate after cycling 600 times at a current rate of 1 C is as low as 0.063% per cycle.

Therefore, we prepared the hierarchical clews of carbon nanobelts (CsCNBs) by KOH activation for LiSBs (Scheme 1) in this work. KOH activation is employed to increase the SSA, pore volume, and number of polar oxygen-containing functional groups. By controlling the mass ratio of KOH/C, the CsCNBs-800*4 has obtained the highest SSA of 2300 m$^2$ g$^{-1}$ and the biggest pore volume of 1.29 cm$^3$ g$^{-1}$. Fundamentally, the CsCNBs with hierarchically porous structure, three-dimensional conductive network framework, and polar oxygen-rich groups are very befitting to load sulfur for the improvement of cycle-stability of LiSBs. Hence, CsCNBs/S composites were prepared. CsCNBs-800*4/S cells have an ultrahigh capacity at low current rates. The initial discharge/charge capacity is 1245/1187 mA h g$^{-1}$ at 0.1 C. The capacity decay rate is as low as 0.162% per cycle for 200 cycles (voltage window is 1.8-2.8 V). At high current rates (>xxxC), the cells still exhibit an excellent cycle stability. At 4 C, the initial discharge/charge capacity is 621/518 mA h g$^{-1}$. After 1000 cycles, the capacity decay rate (only 0.039% per cycle) is extremely low.
Scheme 1. A schematic illustration of the preparation of carbon nanobelts.

**Experimental section**

**Materials**

Hydroquinone [Analytical reagent (AR), 99%), formaldehyde (AR, 37 wt%), hydrochloric acid (AR, 37wt%), sulfur (AR, 99.9%), CS$_2$ (AR), KOH (GR, 95%) and tetrahydrofuran (THF, AR) were purchased from Shanghai Macklin Company (P. R. China).

**Synthesis of CsPNBs**

CsPNBs were synthesized according to ref. 22. In general, 1.65 g of hydroquinone, 2.5 mL of 37 wt% formaldehyde and 115 mL of 10 wt% hydrochloric acid are thoroughly mixed in a 200 mL autoclave with a polytetrafluoroethylene liner. Then, the autoclave was sealed and heated in an oven at 180 °C for 12 hours. The resulting black spongy product was filtered and washed with deionized water. The filter cake was dried under vacuum at 60 °C for 6 h. Finally, 2.0 g of light and dark brown powder was collected. The yield is close to the quantitative value (based on hydroquinone).

**Fabrication of CsCNBs**

CsPNBs (1.80 g) was heated to 800 °C (with a heating rate of 5 °C min$^{-1}$) under a flow of argon (50 mL min$^{-1}$) and then cooled down. This sample was called CsCNBs-800. CsCNBs-800 was chemically activated with KOH. The mixture, consisting of CsCNBs-800 and a certain amount of
KOH, was activated under a flow of argon (50 mL min$^{-1}$) at 400 °C for 0.5 h and then at 800 °C for 1 h at a heating rate of 5 °C min$^{-1}$. The weight ratios of KOH/C were 3, 4 and 5. Therefore, the samples were denoted according to the mass ratio as CsCNBs-800*3, CsCNBs-800*4 and CsCNBs-800*5, respectively. CsCNBs-950-40 was physically activated with CO$_2$ according to ref. 22. The activation time was 40 min.

**Synthesis of CsCNBs/S composites**

CsCNBs (0.08 g) and sulfur (0.12 g) were dispersed in CS$_2$ (1 ml) and stirred till totally dry. To ensure a complete infiltration of sulfur into CsCNBs, the mixture was heated at 155 °C for 12 h. Therefore, all composites contain CsCNBs and sulfur are denoted as CsCNBs-800/S, CsCNBs-800*3/S, CsCNBs-800*4/S, CsCNBs-800*5/S and CsCNBs-950-40/S. Meanwhile, the mass fraction of sulfur in each composite was 60%. Similarly, composite (90 wt% of sulfur) was also prepared and named CsCNBs-800*4/S90.

**Characterization**

Field emission scanning electron microscopy (FESEM) images were collected via a field emission SU-70 microscope. High-resolution transmission electron microscopy (HRTEM) images were obtained with a JEOL JEM2010 electron microscope. High-resolution X-ray diffraction (HR-XRD) patterns were recorded on a Bruker D8 Advance diffractometer using Cu-K$_\alpha$ radiation (40 kV and 200 mA). Raman spectra were measured using a laser confocal Raman spectrometer quipped with an argon-ion laser (514.5 nm). Measurement of the Brunauer–Emmett–Teller (BET) specific surface areas was performed at 77 K using a low-temperature nitrogen adsorption surface area analyser (ASAP 2020, Micromeritics Ins, USA). The total pore volume and pore size distribution were calculated by a density functional theory (DFT) model. Thermogravimetric analysis (TGA) was performed on a TGA-Q55 thermal gravimetric analyzer. X-ray photoelectron spectroscopy (XPS) experiments were carried out on a Microlab 350 spectrometer using a
monochromatic Mg-Ka X-ray source.

**Cell fabrication and measurements**

According to the Ref. 22, the lithium-sulfur cells (CR2016) were fabricated in a glove box (O2<0.1 ppm; H2O < 0.1 ppm) to test the electrochemical performances of CsCNBs/S. The cathode slurry was prepared by mixing 80% of the composite, 10% Super-P and 10% of PVDF binder in 1-methyl-2-pyrrolidinone (NMP). The slurry was blade cast onto carbon-coated aluminum foil and dried at 60 °C for 12 h under vacuum. For composites with 60% sulfur, the sulfur loading for each cathode foil was about 1.0 mg cm$^{-2}$. The sulfur loading for CsCNBs-800$^*$4/S90 was approximately 1.8 mg cm$^{-2}$. The diameter of the cathode was 15 mm. The electrolyte consisted of 1 M LiTFSI and 0.1 M LiNO$_3$ in DME/DOL (1,2-dimethoxyethane/1,3-dioxolane)(1:1 by volume). The ratio of electrolyte to sulfur was 20 µL mg$^{-1}$. The cyclic voltammetry (CV) measurements were carried out on an electrochemical workstation (CHI660E, Chenhua Instrument Company, Shanghai, China) over the potential range of 1.8-2.8 V vs. Li$^+$/Li at a scan rate of 0.2 mV s$^{-1}$. The charge/discharge experiments were conducted on NEWARE (Shenzhen, China) instruments (model 5 V-10 mA and 5 V-50 mA) with a voltage window of 1.8-2.8 V vs. Li$^+$/Li.

**Results and discussion**

The FESEM images of carbonized sample (CsCNBs-800) and activated samples (CsCNBs-800$^*$4 and CsCNBs-950-40) obtained by different methods are shown in Fig. 1. As shown in Fig. 1a, 1c and 1e, it can be clearly seen that the diameter of clews is in the range of 5-30 µm. Meanwhile, there is no obvious change in the morphology of clews after activation. In Fig. 1b, 1d and 1f, it can be found that the clews are disorderly intertwined by a large number of very thin nanobelts. In addition, the nanobelt width of the activated samples is narrower than that of the carbonized sample.
The HRTEM images of CsCNBs-800, CsCNBs-800*4 and CsCNBs-950-40 samples are shown in Fig. 2. In Fig. 2a, 2c and 2e, it can be clearly observed that three samples display same structure, which indicates that the nanobelts remain unchanged after carbonization and activation. It can be seen that the nanobelts are narrow and thin, consistent with the FESEM images. In Fig. 2b, 2d and 2f, there are a lot of disordered black stripes, which correspond to the (002) facet of amorphous carbon [32-34]. High density white spots can also be seen, which are mainly characterized by mesopores.
Fig. 2. HRTEM images of (a, b) CsCNBs-800, (c, d) CsCNBs-800*4, (e, f) CsCNBs-950-40.

Fig. 3a shows the XRD patterns of carbonized sample (CsCNBs-800) and activated samples (CsCNBs-800*3, CsCNBs-800*4, CsCNBs-800*5 and CsCNBs-950-40), which exhibit two diffraction peaks like a steamed bread at approximately 24° and 43° [35-36], corresponding to the diffraction of (002) and (100) planes of graphite, indicating that this microporous carbon consists of small domains of ordered graphene sheets, typical for amorphous carbons with non-crystalline structures [37-39]. After activation, the diffraction peaks of samples are much weaker, especially CsCNBs-950-40, indicating that activation is very effective in increasing disorder and defects. The Raman spectra of all samples are shown in Fig. 3b, which display two characteristic peaks unique to carbon at 1360 cm\(^{-1}\) and 1586 cm\(^{-1}\) [40]. The characteristic peak at 1586 cm\(^{-1}\) named G band is produced by the stretching movement of all sp\(^2\) atoms in the carboatomic rings or carbon chains.
The characteristic peak at 1360 cm\(^{-1}\) named D band is generated by the breathing vibration mode of sp\(^2\) atoms in the hexagonal carbocycles, and induced by disorder and defects. Generally, the R value (intensity ratio of I\(_D\)/I\(_G\)) represents the degree of graphitization and disorder of carbons [41-43]. In comparison to CsCNBs-800, the R values of samples after activation are higher, especially CsCNBs-950-40, which is consistent with the analytical result of the XRD patterns. Especially, there is a 2D characteristic band at 2700 cm\(^{-1}\) in the Raman spectrum of samples after KOH activation, indicating the presence of few layer graphene [44-47]. Fig. 3c shows the nitrogen adsorption-desorption isotherms of CsCNBs-800, CsCNBs-800*3, CsCNBs-800*4, CsCNBs-800*5 and CsCNBs-950-40, which characterize the properties of the surface and pores. According to the IUPAC classification, the isotherm of all CsCNBs samples display the characteristic of type I/IV isotherms, indicating the existence of a large number of micropores and mesopores. At a relative pressure bigger than 0.9, the degree of nitrogen adsorption isotherms rapidly rises, indicating that macropores are ubiquitous in all samples. The pore size distribution (PSD) curve based on the DFT theory also verifies this conclusion. In Fig. 3d, the PSD curve of CsCNBs-800 shows that micropores are mainly located at about 0.54, 0.80 and 1.18 nm, and there is a small amount of mesoporous. The PSD curves of CsCNBs after activation show that mesopores are located at about 2.5 nm and micropores at 0.54, 0.80, 1.18 and 1.48 nm. The existence of numerous micropores and mesopores also confirms the results of HRTEM. The textural parameters of CsCNBs are shown in Table S1. Calculated by BET model, compared with other activated samples, CsCNBs-950-40 has a maximum SSA (2610 m\(^2\) g\(^{-1}\)). However, the DFT approach is more accurate than the BET model to calculate the SSA for microporous carbon. For KOH activation, the SSA obtained by DFT approach increases with increasing degree of activation. When the KOH/C mass ratio is 5, the SSA is significantly decreased, indicating that the amount of KOH is excessive at this time. CsCNBs-800*4 has the same SSA as CsCNBs-950-40. CsCNBs-800*4 and CsCNBs-950-40 have
the same excellent characteristic parameters, such as, higher DFT specific surface area values and pore volume, and the rational distribution of micropores and mesopores, which are better than those of other samples. It is worth noting that the carbonization yield of CsCNBs-800*4 is 1.7 times that of CsCNBs-950-40. The inherent three-dimensional conductive network structure of the CsCNBs material ensures high accessibility and utilization of the nanobelt surface for ion adsorption. In addition, with the three-dimensional conductive network structure, the direct exposure of the highly porous nanobelt to the electrolyte can effectively shorten the transmission path and reduce the resistance of ion diffusion. These structural features can give CsCNBs an excellent potential for the uses in high performance LiSBS.

Fig. 3. (a) XRD patterns and (b) Raman spectra of the five carbon samples; (c) nitrogen adsorption-desorption isotherms and (d) PSD curves of the five carbon samples, calculated from the nitrogen adsorption isotherms using the DFT method.

The XPS survey spectra of CsCNBs-800, CsCNBs-800*4 and CsCNBs-950-40, including carbon and oxygen, are shown in Fig. 4. The carbon and oxygen contents are separately calculated from the peak areas of C 1s and O 1s peaks, which are consistent with the results of the elemental analysis test in Table S2. In Fig. 4a, 4d and 4g, the three samples possess the similar XPS survey
spectra. In Fig. 4b, the C 1s spectrum of CsCNBs-800 sample displays three different component peaks, ascribed to C−C/C=C (284.3 eV), C−O (285.9 eV) and C=O (287.2 eV). Correspondingly, the O1s spectrum of CsCNBs-800 clearly exhibits three oxygen-based groups, corresponding to C=O (531.7 eV), C−O (533.3 eV), and H2O (534.7 eV) in Fig. 4c. The spectra (C 1s spectrum in Fig. 4h and O 1s spectrum in Fig. 4i) of CsCNBs-950-40 sample is similar to CsCNBs-800 sample, excepting that the oxygen content is greatly reduced. After KOH activation, the C 1s spectrum of CsCNBs-800*4 (Fig. 4e) has an extra weak peak ascribed to O−C=O species (289.7 eV). The O 1s spectrum of CsCNBs-800*4 (Fig. 4f) additionally possesses a considerable peak corresponding to C−OH/C−O−C (536.1 eV). The C-OH/C-O-C functional group (hydroxyl/epoxide) is a typical polar functional group with a high surface energy, which has a strong binding effect on polysulfides and is a great benefit to LiSBs that can achieve high-rate and high cycling stability. In general, compared to CO2 activation, KOH activation facilitates enhancing the retention of oxygen in carbons and creating a new polar oxygen-containing functional group.
Due to its unique multi-level pore structure, three-dimensional high conductivity network, high surface area and pore volume, CsCNBs materials are expected to be used in various energy applications such as energy storage, adsorption and catalysis. As a proof of concept, CsCNBs are used for LiSBS after loading sulfur. For the consistency of the experimental variables, all CsCNBs samples were loaded with 60% sulfur. In order to study the electrochemical performance at high sulfur content, CsCNBs-800*4 is used to synthesize a carbon-sulfur composite (90 wt% of sulfur), due to its high specific surface area and high pore volume.

The FESEM images of all CsCNBs/S samples are shown in Figure 5. As is illustrated in the picture, the nanobelts in CsCNBs-800/S and CsCNBs-800*3/S are thicker than other three samples with a sulfur content of 60%, because there are some sulfur on the surface of nanobelts. However, CsCNBs-800*4/S, CsCNBs-800*5/S and CsCNBs-950-40/S have high pore volume, and almost all of the sulfur is adsorbed in the micropores and mesopores. As shown in Figure 5f, the sulfur is clearly coated on the surface of the CsCNBs-800*4/S90 nanobelts, which is attributed to the high pore volume of CsCNBs-800*4, which cannot fully adsorb high sulfur content. In Fig. 6, the carbon, nitrogen, oxygen and sulfur mappings of CsCNBs-800*4/S display a similar pattern, demonstrating that the homogeneous distribution of these elements. It is further proved that the sulfur is well embedded in CsCNBs-800*4/S.
Fig. 5. FESEM images of (a) CsCNBs-800/S, (b) CsCNBs-800*3/S, (c) CsCNBs-800*4/S, (d) CsCNBs-800*5/S, (e) CsCNBs-950-40, (f) CsCNBs-800*4/S90.
Fig. 6. The EDX images of CsCNBs-800*4/S composite.

Fig. 7. (a) XRD patterns, (b) Raman spectra and (c) TGA of all carbon-sulfur composites.

The XRD patterns of pure sulfur and all carbon/sulfur composites are shown in Fig. 7a. Obvious sulfur characteristic peaks can be seen in the XRD curves of CsCNBs-800/S and CsCNBs-800*3/S, which masks some characteristic peaks of carbon materials. On the XRD curves of CsCNBs-800*4/S and CsCNBs-950-40/S, the sulfur characteristic peaks disappear because the sulfur, dispersed in the micropores and mesopores, is inhibited by the pores and cannot be crystallized. However, there are a few sulfur characteristic peaks in the XRD curve of
CsCNBs-800*5/S, because the pore volume of CsCNBs-800*5 is not large enough to completely accommodate and adsorb sulfur. The sulfur characteristic peak in the XRD curve of the CsCNBs-800*4/S90 composite completely obscures the carbon characteristic peak because the pore structure in the CsCNBs-800*4 material cannot completely contain sulfur and the sulfur is distributed on the surface of the nanobelts. The Raman spectra of pure sulfur and all carbon/sulfur composites are shown in Fig. 7b. All CsCNBs/S composites with 60% sulfur content showed only D band and G band of amorphous carbon, and no sulfur characteristic peak was observed. However, a strong sulfur characteristic peak can be seen in the Raman spectrum of the CsCNBs-800*4/S90 composite due to its high sulfur content. Compared with the intensity ratio of the corresponding CsCNBs, the R values of all CsCNBs/S composites are slightly reduced, indicating that sulfur is present in the pore structure of all samples, thereby increasing the degree of order. Fig. 7c shows the TGA of all carbon/sulfur composites. The actual sulfur content of CsCNBs-800*4/S90 composite is 89.7%, and other composites are close to 60%. The sulfur contents of all composites have reached the expected setting. The weight loss curves of CsCNBs-800/S, CsCNBs-800*3/S and CsCNBs-800*4/S90 composites are divided into two stages. The weight loss steps in the range of 150~250 °C and 250~400 °C are attributed to the volatilization of the sulfur outside and inside micropores/mesopores, respectively. Since the pore volume of CsCNBs-800 is small (0.31 cm$^3$ g$^{-1}$), the sulfur inside the micropores/mesopores is less than 5% (based on the whole composite). Due to the smaller pore volume of CsCNBs-800*3 (0.60 cm$^3$ g$^{-1}$), the sulfur inside the micropores/mesopores is around 40% (based on the whole composite). During the temperature rises from 150 to 250 °C, about 70% (based on the entire complex) of sulfur is volatilized in the CsCNBs-800*4/S90 composite, which indicates that the high pore volume of the CsCNBs-800*4 is not sufficient to accommodate ultrahigh content of sulfur. However, the weight loss curves of CsCNBs-800*4/S, CsCNBs-800*5/S and CsCNBs-950-40/S composites have only one weight loss...
process, because the pore volume of CsCNBs-800*4, CsCNBs-800*5 and CsCNBs-950-40 is large enough to encapsulate most of the active substance sulfur. Table S3 shows the elemental analysis data of each CsCNBs/S composite, which is basically consistent with the TGA results.

To evaluate the electrochemical properties of these samples, all CsCNBs/S composites are assembled into a button-type LiSBs, in which a lithium metal sheet is used as a negative electrode. The assembling batteries are cycled at various current rates in a voltage window of 1.8-2.8 V [48-50]. Figure 8a shows the initial discharge/charge curves of each battery at 0.1 C [51]. The discharge curves of all samples display two platforms of discharge at 2.3 V and 2.1 V, corresponding to the characteristic discharge platforms of sulfur [52-53]. Comparing with other composites, both discharge platforms of CsCNBs-800*4/S90 are moving down because the ultrahigh content of sulfur reduces the conductivity of the entire composite. There is a small platform in the discharge curve of CsCNBs-800*X (X=3, 4, 5)/S, and CsCNBs-950-40/S at approximately 1.8 V, which is attributed to the consumption of LiNO₃ in the electrolyte [54-55]. The reduction of LiNO₃ may be catalyzed by the bare carbon surface with a high specific surface area at a small current rate. Fig.8b shows the cycle curve of each battery at 0.1C. The initial discharge specific capacities of CsCNBs-800/S, CsCNBs-800*X(X=3, 4, 5)/S, CsCNBs-950-40/S and CsCNBs-800*4/S90 are 839, 1064, 1245, 1170, 1371 and 785 mA h g⁻¹, respectively. It can be found that the larger the specific surface area and pore volume of the CsCNBs material, the higher the initial discharge specific capacity of the corresponding CsCNBs/S composite. For the CsCNBs-800*4/S90 composite, the sulfur content is 90%, which causes the conductivity of the positive electrode to decrease, resulting in most of the available sulfur not being fully utilized. Therefore, a lower but acceptable discharge specific capacity is exhibited. After 200 cycles of charge/discharge test, the capacity retention rates of CsCNBs-800/S, CsCNBs-800*X(X=3, 4, 5)/S, CsCNBs-950-40/S and CsCNBs-800*4/S90 are 65.2%, 58.8%, 67.5%, 62.9%, 44.0% and 70.7%,
respectively. The CsCNBs-800*4/S exhibits the best long-term cycling stability at 0.1 C with a capacity decay rate as low as 0.162% per cycle. When the sulfur content is as high as 90% (CsCNBs-800*4/S90), an excellent long-term cycling stability is still retained. However, the capacity decay rate of CsCNBs-950-40/S composite with almost specific surface area and pore volume is 0.280% per cycle. In short, the superior cycle stability of CsCNBs-800*4/S composite can be attributed to its rich polar oxygen-containing functional groups, which have a stronger adsorption effect on polysulfides and increase the utilization of sulfur [56-57].

**Fig. 8.** Electrochemical performance of all composites: (a) the initial discharge/charge curves and the cycle performances at (b) 0.1 C, (c) 0.5 C and (d) 1 C.

As shown in Fig. 8c, the initial discharge specific capacities of CsCNBs-800/S, CsCNBs-800*X(X=3, 4, 5)/S, CsCNBs-950-40/S and CsCNBs-800*4/S90 are 611, 895, 974, 977, 1035 and 572 mA h g⁻¹, respectively. After 400 cycles, the capacity retention rates are 69.0%, 67.2%, 72.4%, 71.5%, 61.0% and 67.7%, respectively. Each battery exhibits an excellent long-term cycling stability, especially for the battery made from the CsCNBs-800*4/S. The cyclic curve of each battery test at 1 C current rate is shown in Fig. 8d. The initial discharge specific capacities of
CsCNBs-800/S, CsCNBs-800*X(X=3, 4, 5)/S, CsCNBs-950-40/S and CsCNBs-800*4/S90 are 544, 611, 884, 922, 939 and 440 mA h g\(^{-1}\), respectively. After 600 cycles, the capacity retention rates are 80.0%, 75.1%, 75.0%, 61.8%, 65.0% and 63.4%, respectively. Discharged at high currents, CsCNBs-800*4/S composites still exhibit high capacity and excellent long-term cycle stability. Even with a sulfur content of 90%, the CsCNBs-800*4/S90 electrode obtains acceptable electrochemical performance.

![Fig. 9. The CV curves of (a) CsCNBs-800*4/S and (b) CsCNBs-950-40/S measured between 1.8 and 2.8 V versus Li\(^+\)/Li at a sweep rate of 0.2 mV s\(^{-1}\). Charge/discharge curves of (c) CsCNBs-800*4/S and (d) CsCNBs-950-40/S at 0.5 C.](image)

Table S4 is the electrochemical performance of the CsCNBs-800*4/S and CsCNBs-950-40/S in LiSBs. It is found that the CsCNBs-800*4/S composite exhibits a low initial discharge specific capacity, but the decay rate of per cycle capacity is much lower than that of CsCNBs-950-40/S composites, especially at low current cycles. Porous carbon materials, CsCNBs-800*4 and CsCNBs-950-40, have similar specific surface area and pore volume, three-dimensional conductive network, pore structure, etc. The biggest difference is the difference in oxygen content. As can be
seen from Fig. 4f, CsCNBs-800*4 sample has rich polar oxygen-containing functional groups (C-OH/C-O-C, hydroxyl/epoxide). These functional groups are favorable for adsorbing lithium polysulfide during the cycles of a lithium-sulfur battery, suppressing the shuttle effect, and improving the utilization rate of the active material sulfur. Adequate evidence for the excellent electrochemical performance is exhibited by CsCNBs-800*4 sample.

As mentioned above, the CsCNBs-800*4/S composite exhibits the best electrochemical performance. Therefore, the electrochemical performances of CsCNBs-800*4/S and CsCNBs-950-40/S composites are further investigated by CV testing [58-59]. As shown in Fig. 9a, CsCNBs-800*4/S composite displays two peaks at 2.25 and 1.95 V during the first voltage scan. During the second voltage sweep, the peak at 2.25 V is unchanged, while the peak at 1.95 V is moved to 2.00 V. During the fifth voltage sweep, the curves are substantially coincident and the peaks are unchanged, indicating that good electrochemical stability and reversibility. Fig. 9c shows the first, second, and fifth discharge/charge curves at 0.5 C, which show that the migration of the discharge platform is consistent with the CV curve and the capacity attenuation is small. Fig. 9b shows the CV curves of CsCNBs-800*4/S composite. During the first voltage scan, there are two peaks at 2.20 and 1.90 V. During the second and fifth voltage sweeps, the peak of high voltage discharge platform is essentially unchanged, and the peak of low voltage discharge platform migrates to 1.95 V. Fig. 9d shows the first, second, and fifth discharge/charge curves at 0.5 C, which show that the displacement of the discharge platform is consistent with the CV curve, and the capacity decays faster. The CV curves further indicate that CsCNBs-800*4/S composite has a small capacity attenuation and high cycle stability.
In order to visually characterize the rate cycling characteristics, the batteries of CsCNBs-950-40/S and CsCNBs-800*4/S composites are tested for charge/discharge cycles at various current rates [60]. Fig. 10a shows the rate performance of CsCNBs-950-40/S composite. At 0.1 C, the discharge specific capacity of CsCNBs-950-40/S rapidly decreased from 1348 mA h g\(^{-1}\) to 1158 mA h g\(^{-1}\). The capacities of CsCNBs-950-40/S at 0.2, 0.5, 1, and 2 C are 958, 832, 746, and 597 mA h g\(^{-1}\), respectively. When the current rate is switched back to 1 and 0.5 C, the capacities can be largely recovered. However, when the current rate is switched back to 0.2 and 0.1 C, the capacity loss is large. The rate performance of CsCNBs-800*4/S composite is shown in Fig. 10b. At 0.1 C, the discharge specific capacity of CsCNBs-950-40/S rapidly decreased from 1225 mA h g\(^{-1}\) to 1141 mA h g\(^{-1}\). The capacities of CsCNBs-800*4/S at 0.2, 0.5, 1, and 2 C are about 1005, 907, 835, and 761 mA h g\(^{-1}\), respectively. When the current rate is switched back to 1 C, 0.5 C, 0.2 C, and 0.1 C, the discharge specific capacity is greatly restored. At 0.1 C, discharge specific capacity is 1031 mA.
h g\(^{-1}\), indicating that the CsCNBs-800\(^*\)4/S composite has an excellent rate cycling performance.

In order to characterize the high current discharge performance, the batteries of CsCNBs-800\(^*\)4/S are tested at 2 C, 3 C and 4 C [11, 61, 62]. The initial discharge/charge curve is shown in Fig. 10c. All initial discharge curves show two sulfur discharge platforms at 2.20 V and 1.95 V. At 2 C, 3 C and 4 C, the initial discharge ratio capacities are 797, 612 and 621 mA h g\(^{-1}\), respectively. The coulombic efficiency decreased in turn 96.2%, 94.5% and 93.6%, respectively. The cycle performances and coulombic efficiencies of CsCNBs-800\(^*\)4/S batteries at 2 C, 3 C and 4 C are shown in Fig.10d. After 1000 cycles of charge and discharge, the discharge specific capacity of the battery at each current rate was reduced to 500, 425 and 381 mA h g\(^{-1}\), respectively. The capacity decay rate was as low as 0.037%, 0.030% and 0.039% per cycle, indicating a slow loss of sulfur during the cycle. Although the first coulombic efficiency of CsCNBs-800\(^*\)4/S composites at 2 C, 3 C and 4 C is lower, they are very close to 100% from the second to the 1000\(^{th}\) cycle, indicating that the shuttle effect of sulfides is effectively suppressed during cycles.

Compared with CsCNBs-950-40 material, CsCNBs-800\(^*\)4 material exhibits superior electrochemical performance in LiSBs, including high capacity, excellent rate performance and long-term high cycling stability, which can be ascribed to four main factors. First, the highly conductive three-dimensional network structure of the nanobelts promotes the rapid transmission of electrons and improves the overall conductivity of the positive electrode. The EIS results of CsCNBs-950-40 and CsCNBs-800\(^*\)4 have further investigated this point (supplementary materials Fig.6). Second, the nanobelts are very thin (the range of 10-30 nm), and the diffusion path of lithium ions can be reduced to half the thickness of the nanobelts (about 10 nm), resulting in rapid lithium ion kinetics. Third, the synergistic effect of multi-stage pores stores sulfur, has physical adsorption to lithium polysulfides, and inhibits the occurrence of shuttle effect. The foregoing three mechanism factors of CsCNBs-800\(^*\)4 material are basically similar to those of CsCNBs-950-40.
Fourth, the polar oxygen-containing (hydroxyl/epoxide) functional groups can strongly bind to lithium polysulfides and have a strong chemisorption to polysulfides, which can prevent polysulfides from dissolving into the electrolyte and inhibit the generation of shuttle effect. This is the core of CsCNBs-800*4 material with superior rate performance and cycle stability. In addition, the electrochemical performance of other related carbon/sulfur composite were exhibited in Supplementary materials Table S5, and these results further prove the outstanding electrochemical performance of our prepared carbon nanobelts in LiSBs.

**Conclusion**

In summary, CsCNBs-800*4 with ultrahigh SSA (2300 m² g⁻¹) and large pore volume (up to 1.29 cm³ g⁻¹) has been successfully fabricated from phenolic resin based nanobelts by carbonization and KOH activation. The product possesses hierarchically porous structure, three-dimensional conductive network framework, and polar oxygen-rich groups, which are very befitting to load sulfur for excellent cycle-stability of LiSBs. As a result, CsCNBs-800*4/S cells display an ultrahigh initial discharge capacity of 1245 mA h g⁻¹ and ultralow capacity decay rate as low as 0.162% per cycle after 200 cycles at 0.1 C. Even at high current rate of 4 C, the cells still exhibit a high initial discharge capacity (621 mA h g⁻¹) and ultralow capacity decay rate (only 0.039% per cycle) after 1000 cycles. These encouraging results indicate that polar oxygen-containing functional groups are important for improving the electrochemical performance of carbons. The oxygen-doped carbon nanobelts have an excellent energy storage potential in the field of energy storage.

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