S, N co-doped pitch-based composite carbon nanofibers with enlarged interlayer distance as a superior potassium ion batteries anode

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Abstract. Potassium ion batteries (PIBs), an alternative to traditional lithium ion batteries to large-scale energy storage device, have attracted tremendous attention, due to abundant reserves of potassium resources and low cost. However, it still remains challenge to fabricate suitable anode materials with high K storage capabilities. In this work, facile S/N co-doped pitch based composite carbon nanofibers has been fabricated by electrospinning of coal tar pitch and polyacrylonitrile, and followed by carbonization under H₂/S/Ar atmosphere. The formation of -C-S-C- bond effectively increased S utilization, and enlarged carbon interlayer distance to some degree. As anode for PIBs, the S/N co-doped carbon displayed enhancement of capacity, rate capability and cycle stability. This work would shed a light on the fabrication of S/N co-doped materials for both battery, supercapacitor and electrocatalytic electrodes.

1. Introduction

Lithium ion batteries (LIBs) have been widely used in portable electronic equipment and electric vehicles. However, the large-scale utilization of lithium ion battery will inevitably push up the price of lithium resources, and the high production cost of lithium ion battery makes it unable to meet the requirements of large-scale energy storage devices in the future [1, 2]. Thus, as an alternative to LIBs, it is imperative to develop secondary batteries with rich resources and stable performance for the future large-scale energy storage system.

In recent years, potassium ion batteries (PIBs) have aroused tremendous attention as one of the most promising energy storage systems [3]. Potassium shows a lower redox potential (-2.936 V vs. K/K⁺) than that of sodium (-2.714 V vs. Na/Na⁺) and similar to that of lithium (-3.040 V vs. Li/Li⁺) [4]. Therefore, PIBs are expected to present an advantage in high voltage output as LIBs. Besides, because of weaker Lewis acidity of K⁺, the solvated K⁺ are smaller than Li⁺ and Na⁺, which enables potassium a faster migration rate in the electrolyte [4]. In addition, compared with lithium and sodium based electrolyte, there are fewer side reactions on the surface of anode materials in potassium-based electrolyte [5]. Graphite, general anode material used in LIBs, can form stage-one intercalation compounds of KC₈, however, the narrowed interlayer distance leading to a relatively low theoretical capacity of only 279 mAh g⁻¹. Moreover, the fully-potassiated graphite causes ~60% volumetric expansion, which would deteriorate cycle stability. Therefore, the fabrication of novel materials focused on large radius K⁺ storage is the key issue to realize the application of PIBs.

A series of anode materials have been conducted to for PIBs, for example, alloys [6, 7], metal oxide/sulfide [8, 9], and carbonaceous materials. Though Non-carbon materials generally present high capacity, however, the violent volume expansion inevitably causes poor cycle stability. Carbonaceous materials, such as soft carbon [10, 11], hard carbon [12, 13], graphene [14, 15], have been explored. However, the relatively low capacity still cannot meet the requirement of high-performance electrode. Heteroatom-doped (N, P, S, B) carbonaceous have attracted great attention because of the reinforcement of electrical conductivity, adsorption to K⁺, and enhancement of specific surface area. N has been confirmed efficient to further enhance the K storage of carbon materials by its higher electronegativity, induced extra defect, and enlarged interlayer distance [15, 16]. In addition, S doping with larger atomic radium and lower electronegativity would be beneficial for further expanding the interlayer space, and provide more active sites. As for PIBs anode, only a few work have been carried out so far, such as sulfur-doped carbon microspheres [13] and nanosheets [12]. In order to realize an extend interlayer distance and enhanced conductivity, synergistic effect of binary (S/N) doping is worth in consideration. Yang et al. fabricated S-doped N-rich carbon nanosheets as SIBs anodes with expanded interlayer distance. they found that the unique S priority substitute pyrrolic N, yielding a superior Na storage property [17]. However, only a few literatures up to now focus on S/N carbon materials for PIBs [18-20]. Thus, the...
exploration of novel strategies for S/N co-doped carbon for PIBs anode still remains challenging.

Herein, facile S/N co-doped pitch based composite carbon nanofibers has been fabricated by electrospinning of coal tar pitch and polyacrylonitrile, and followed by H2S/Ar carbonization. The enlarged interlayer distance could efficiently provide more active sites and fast K+ migration. Hence N/S doped carbon nanofibers showed high capacity, excellent rate and cycle performance.

2.Experimental section

2.1. Fabrication of S/N co-doped pitch-based carbon nanofibers

Pitch-based nanofibers was fabricated by electrospinning of coal tar pitch and polyacrylonitrile (PAN) [21]. Typically, 2 g PAN (Mw=150,000) was first dissolved in dimethylformamide at 10 wt%. Then an equal amount of coal tar pitch (purchased from Ansteel Group Corporation) was added to the solution, followed by stirring at 60 °C for 4 h. The dark brown suspension was conducted in our home-made electrospinning device. The voltage, needle-to-collector distance, and inner diameter of the needle was 18 kV, 20 cm, and 0.84 mm, respectively. Then, the as-spun nanofibers were vacuum dried over night to remove the residue solvent at 60 oC. The nanofiber membrane was first stabilized in air at 280 °C for 2 h with 1°C min-1, then carbonized in H2S/Ar (5/95 (v/v)) mix gas at 800 °C for 2 h with 3°C min-1. The obtained S/N co-doped pitch-based carbon was donated as S-NC. For comparison, the stabilized nanofibers were carbonized in Ar under the same conditions as mentioned, termed as NC. The experimental process can be seen in Fig. 1.

2.2. Material characterizations

transmission electron microscope, (TEM, FEI Tecnai F30, 300 kV), scanning electron microscope, (SEM, FEI, Nova450), and X-ray diffraction (XRD, Rigaku D/Max2400) were employed to measure the morphology and structure of both two samples. X-ray photoelectron spectroscopy (XPS, Thermo ESCALAB 250) was used to analyze surface composition. Raman (Thermo Fisher DXR) and FT-IR (Thermo Fisher 6700) spectra were conducted to analyze the microstructure and groups. Elemental analyses were measured by Vario EL analyzer.

2.3. Electrochemical measurements

S-NC and NC membranes were cut into disks with 1.2 cm in diameter, and used as anodes directly. 2016-type half cells were assembled in glovebox filled with Ar, followed by using 0.8 mol L−1 KPF6 (in EC/DEC (v:v=1:1)) as the electrolyte, and K foil as counter electrode. The galvanostatic discharge/charge tests were measured by using Neware CT-4008 at 0.01~2.5 V (vs. K/K+). Electrochemical impedance spectroscopy (EIS) and Cyclic voltammetry (CV) measurements were tested on a VMP-300 electrochemical work-station, from 0.01~2.5 V (vs. K/K+) and 100 kHz to 10 mHz, respectively.

3. Results and discussion

The composition of coal tar pitch was first analyzed. Fig. 2a shows the FT-IR spectrum of coal tar pitch. The characteristic peak at 3028 cm-1 corresponds to the stretching vibration of unsaturated C-H bond in the condensed aromatics, and the characteristic peaks at 2920 and 2852 cm-1 refer to the stretching vibration of saturated C-H bond, indicating that a small amount of alkyl side chains exist in the aromatic molecules of coal tar pitch. Elemental analysis shows that the mass ratio of carbon, hydrogen, nitrogen and oxygen of coal tar pitch is 92.4%, 4.5%, 1.0% and 2.1%, respectively.

![Fig. 1. Fabrication procedure of S-NC.](image)

![Fig. 2. (a) FT-IR spectrum of coal tar pitch. (b, c) SEM image, (d) diameter distribution, (e) TEM image, and (f) EDS elemental mapping of S-NC.](image)
As shown in Fig. 2b, S-NC membrane is weaved with non-oriented carbon nanofibers. The one-dimensional nanostructure is conducive to shortening the potassium ion diffusion distance, and the three-dimensional cross-linked network is favorable to reducing the electron transport resistance. The non-woven fiber membrane maintains good flexibility and structural stability. From the magnified image of Fig. 2c to 2e, the diameter mainly ranges from 280 to 320 nm with rough surface. In addition, EDS elemental mapping indicates homogeneous distribution of element C, S, and N in the entire area of fibers (Fig. 2f).

From the XRD patterns of Fig. 3a, both S-NC and NC show broad peaks at around 25°, indicating their amorphous nature. S-NC shows an obvious lower 2 theta degree, which suggests that S-NC has a larger d002 (0.367 nm) than that of NC (0.354 nm), revealing that the doped S effectively extends the interlayer distance by its larger atomic radius. The enlarged interlayer distance would provide sufficient buffer to huge volume expansion, as well as abundant ion storage sites and reduced ion diffusion distance. In Raman spectra (Fig. 3b), both S-NC and NC present two broad bands near 1350 cm⁻¹ (D band) and 1580 cm⁻¹ (G band), which respectively correspond to amorphous region and in-plane vibration of sp² hybridized carbon. Generally, the intensity ratio (ID/IG) of D band and G band is used to approximately evaluate the order degree of nano-crystallinity. S-NC has a larger ID/IG (1.04) than that of NC (0.99), indicating a higher structural distortion induced by S doping.

As shown in Table 1, NC has a relatively high nitrogen content (8.9 wt%) but negligible sulfur. In contrast, after H2S heat treatment, S-NC present a markedly elevated sulfur content (6.3 wt%). In Fig. 3c, carbon, nitrogen, oxygen can be detected for both NC and S-NC, while, a pair of peaks centered at around 163 eV and 228 eV respectively respond to S 2p and S 2s in S-NC. The element content change is similar to that of elemental analysis (Table 2). Fig. 3d shows three peaks for sulfur at 163.5, 164.7, and 167.5 eV, which can be respectively assigned to S 2p3/2, S 2p1/2, and oxidized S, indicating the formation of -C-S-C- bond in the S-NC. Besides, it is notable that, an obvious decrease of nitrogen content occurs with the raising Sulfur content in both elemental and XPS analyses, which may related to the substitution of sulfur for pyrrolic-N, forming thiophene-like structure [17].

![Fig. 3.](image)

**Fig. 3.** (a) XRD patterns, (b) Raman spectra, and (c) XPS spectra of NC and S-NC, respectively. (d) XPS spectrum of S 2p in S-NC.

Elemental and XPS analyses were conducted to evaluate the element content and chemical state of surface atom. As shown in Table 1, NC has a relatively high nitrogen content (8.9 wt%) but negligible sulfur. In contrast, after H2S heat treatment, S-NC present a markedly elevated sulfur content (6.3 wt%). In Fig. 3c, carbon, nitrogen, oxygen can be detected for both NC and S-NC, while, a pair of peaks centered at around 163 eV and 228 eV respectively respond to S 2p and S 2s in S-NC. The element content change is similar to that of elemental analysis (Table 2). Fig. 3d shows three peaks for sulfur at 163.5, 164.7, and 167.5 eV, which can be respectively assigned to S 2p3/2, S 2p1/2, and oxidized S, indicating the formation of -C-S-C- bond in the S-NC. Besides, it is notable that, an obvious decrease of nitrogen content occurs with the raising Sulfur content in both elemental and XPS analyses, which may related to the substitution of sulfur for pyrrolic-N, forming thiophene-like structure [17].

**Table 1.** Elemental analysis of NC and S-NC

| Sample | C       | H       | N       | S       | O* |
|--------|---------|---------|---------|---------|----|
| NC     | 80.3    | 1.2     | 8.9     | 9.6     |    |
| S-NC   | 80.5    | 1.1     | 3.5     | 6.3     | 8.6|

*: by difference

**Table 2.** XPS analysis of NC and S-NC

| Sample | C  | N  | S  | O  |
|--------|----|----|----|----|
| NC     | 80.9| 5.6| 2.0| 4.4|
| S-NC   | 84.6| 2.0| 4.4| 9.0|

Fig. 4a shows the initial five galvanostatic discharge/charge profiles of S-NC at 0.05 A g⁻¹. The initial discharge and charge capacity is 564 and 299 mAh g⁻¹, respectively, with an initial Columbic efficiency of 53.0%. The irreversible capacity can be related to the formation of SEI film on the electrode surface and other side reactions. The capacity almost overlaps during the 2nd to 5th cycle, indicative of its excellent cycle stability. In dQ/dV profiles (Fig. 4b) of both S-NC and NC at the 5th cycle, an obvious reduction peak for S-NC at around 1.0 V can be assigned to the reaction between S and K, and there is a corresponding oxidation peak at around 1.6 V. Whereas, the peak couple are different from those of K-S batteries, where three couples of redox peaks appears. It can be concluded that the reduction peak at 1.0 V was related to the reaction between K⁺ and S bonded carbon framework, and the C-S-C bond is stable during repeated potassiation and depotassiation.

Benefit from the enlarged interlayer induced by sulfur doping, S-NC shows the reversible capacity of 293, 245, 222, 202, and 180 mAh g⁻¹ at the current density of 0.05, 0.1, 0.2, 0.5, and 1 A g⁻¹ (Fig. 4c). Even at a high current density of 2 A g⁻¹, it still retains 145 mAh g⁻¹. When the current density restores to 0.05 A g⁻¹, 291 mAh g⁻¹ is recovered, indicative of a stable cycle stability. In Fig. 4d, both S-NC and NC show Nyquist plots with a semicircle in the high frequency and a sloping line in low frequency. S-NC displays similar charge transfer resistance, but an obvious larger slope, indicating a higher solid-state ion migration. Fig. 4e displays an outstanding cycle stability of both S-NC and NC. After 1000 cycles at 1 A g⁻¹, S-NC can still retain a capacity of 152 mAh g⁻¹ with a capacity retention of 93.9%. While NC shows a passable cycling with a low capacity of 85 mAh g⁻¹ and 88.9% retention.
4. Conclusions

In summary, a facile S/N co-doped pitch based composite carbon nanofibers has been fabricated by electrospinning of coal tar pitch and PAN, and followed by H2S/Ar carbonization. The formation of -C-S-C- bond effectively extends the interlayer distance of carbon material by its larger atomic radius. As anode materials for PIBs, S-NC exhibits superior capacity of 293 mAh g⁻¹ and remarkable rate performance with 93.9% capacity retention at 1 A g⁻¹ after 1000 cycles. This work would shed light on the pervasive fabrication of S/N co-doped materials for both battery, supercapacitor and electrocatalytic electrodes.

Acknowledgments

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