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Modulating the graphitic domains of hard carbons via tuning resin crosslinking degree to achieve high rate and stable sodium storage

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How to cite this article: Lu Z, Yin X, Ji Y, Li C, Wang R, Li S, Yuan T, Zhao Y. Modulating the graphitic domains of hard carbons via tuning resin crosslinking degree to achieve high rate and stable sodium storage. Energy Mater 2024;4:400038. https://dx. doi.org/10.20517/energymater.2023.117

Received: 26 Dec 2023 First Decision: 30 Jan 2024 Revised: 3 Apr 2024 Accepted: 3 Apr 2024 Published: 22 Apr 2024

Academic Editor: Yuhui Chen Copy Editor: Pei-Yun Wang Production Editor: Pei-Yun Wang

Abstract

Sodium-ion batteries (SIBs) are regarded as an outstanding alternative to lithium-ion batteries (LIBs) due to abundant sodium sources and their similar chemistry. As a most promising anode of SIBs, hard carbons (HCs) receive extensive attention because of their low potential and low cost, but their rational design for commercial SIBs is restricted by their variable and complicated microstructure, which is analogous to that of graphite in LIBs. Herein, a series of controllable HC materials derived from 3-aminophenol formaldehyde resin (AFR) were designed and fabricated. We discover that the optimized HC features expanded graphite regions, highly developed nanopores, and reduced defect content, contributing to the enhanced Na⁺ storage. This optimization is achieved by adjusting the resin crosslinking degree of the precursor. Specifically, a resin precursor with a higher crosslinking degree can produce HC with a larger interlayer distance, relatively higher crystallinity, and a lower specific surface area. Encouragingly, the as-optimized AFR-HC electrode manifests superior electrochemical performance in the aspect of high capacity (383 mAh·g⁻¹ at 0.05 A·g⁻¹), better rate capability (140 mAh·g⁻¹ at 20 A·g⁻¹), and high initial coulombic efficiency (82%) than other contrast samples. Moreover, the as-constructed full cell coupled with a Na₃V₂(PO₄)₃ cathode shows an energy density of 250 Wh·kg⁻¹. Together with the simple synthesis, cost-efficiency of the precursors and superior electrochemical performance, AFR-HCs are promising for the commercial application.

Keywords: Sodium-ion batteries, hard carbon, microstructures, 3-aminophenol formaldehyde resin, high rate



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INTRODUCTION

In the past 30 years, lithium-ion batteries (LIBs) have been the state-of-the-art rechargeable batteries and have been extensively used in electric vehicles and mobile electronics^[1]. However, the low reserves and high cost of lithium, along with the unsafety, impede their usage in large-scale energy storage systems^[2]. Sodiumion batteries (SIBs), which show similar chemistry to LIBs while being much cheaper, are deemed one of the most promising next-generation energy storage techniques to replace LIBs^[3]. Among the various advanced SIB anode materials, hard carbons (HCs) stand out as promising candidates for commercial SIBs by virtue of the abundant active sites for Na⁺ storage^[1], wide sources, and low cost^[2]. It has been demonstrated in the literature that a large reversible capacity ranging from 250 to 350 mA·h·g⁻¹ can be achieved with HC directly derived from resins, biomass, and pitch^[4]. Unfortunately, their reliability toward practical application has been critically questioned, evident in the consistency of electrochemical properties and HC microstructure^[1,5]. This could be attributed to the variable and complicated microstructure of HCs.

To this end, various methods for designing HC microstructures are widely employed to address the aforementioned challenges. Specifically, these include (i) expanding the interlayer distance of graphite for accommodating Na+ with large-radius^[1]; (ii) adjusting the porous structure to enhance Na+/Na cluster storage within appropriate pores; (iii) introducing heteroatoms^[6,7] or defects into the carbon lattice to trigger capacitive adsorption or reactions^[6,8,9]; (iv) constructing a favorable carbon form or structure to enhance ion and electron transfer efficiency. Furthermore, the microstructure of HC, including pore structure, defects, and layer spacing, can be controlled by carefully choosing appropriate precursors and adjusting the synthesis and pyrolysis processes. Various types of materials, including phenolic resin (PR)^[10,11], pitch^[12], anthracite, and biomass^[13], are investigated as precursors of HC anodes for SIBs. Among them, PR, which features distinct amorphous graphitic structures and possesses high-purity carbon content compared with other HC precursors, is taken as the most promising SIB anode precursor for real usage^[14]. Some pioneers prepared HC spheres from PR precursors with tunable structure in a wide temperature range, in which HC carbonized at 1,900 °C achieved an excellent low-voltage plateau capacity, accounting for 84% of the reversible capacity^[15]. However, the design of PR-based HC anodes encounters challenges due to the complicated structure of the precursor and the fussy preparation process. Therefore, the key to achieving high-performance HC anodes for practical applications lies in exploring strategies to regulate the microstructures of HCs and selecting the optimal carbon precursor.

Inspired by the above studies, using 3-aminophenol formaldehyde resin (AFR) as a precursor, we designed and produced various HC materials featuring regular spherical morphology and adjustable microstructure via a simple solution-based routine. Interestingly, we find that the obtained performance of HC shows linear relationship with the crosslinking degree of precursors. In detail, the resin precursor with a higher crosslinking degree could lead to the HC with higher carbon yield, larger d_{002} , fewer defects, and lower specific surface area (SSA). The as-optimized AFR-HC material with the highest crosslinking degree features well-regulated nanopores (0.5~2.0 nm), fewer defects [lower ratio of D- to G-band (I_D/I_G) of 0.95], decent SSA (11 m²·g⁻¹), and larger d_{002} (0.388 nm). These features meet the requirements for Na¹ intercalation and Na clustering during charge/discharge processes. As a result, the as-optimized AFR-HC electrode demonstrates a superior reversible capacity of 383 mAh·g⁻¹ at 0.05 A·g⁻¹, improved rate capability of 140 mAh·g⁻¹ at 20 A·g⁻¹, and enhanced initial coulombic efficiency (ICE) of 82%, surpassing the control samples. This simple and eco-friendly approach for producing HC could be scalable for regulating different kinds of PR carbons.

EXPERIMENTAL

Materials and methods

First, 4 g of 3-aminophenol is dissolved in a mixture of ethanol and water. Then, formaldehyde (37 wt%) is added. The mixed solution was stirred at 70 °C for 1.5 h and then dried at 80 °C for 24 h. Finally, the dried AFR powder was calcined under argon at 1,300 °C to obtain AFR-HCs; the prepared samples were labeled as 0.46-AFR-HC, 0.90-AFR-HC, and 1.80-AFR-HC, respectively.

Materials characterization

The sample morphologies can be observed using scanning electron microscopy (SEM, Hitachi S-4800). High-resolution transmission electron microscope (HRTEM) and selected area electron diffraction (SAED) images were obtained employing the JEM-2010F transmission electron microscope at 200 kV. X-ray diffraction (XRD) (18 kW D/MAX2500) with Cu K α radiation (λ = 0.15418 nm) was used to characterize the sample structure; the tube voltage was 40 kV, and the current was 40 mA. The molecular vibration and rotation information of the carbon material was qualitatively analyzed by confocal Raman microscopy (Nanofinder30) with a 532 nm laser excitation. The elemental composition on the surface of the material was measured by the K-alpha X-ray photoelectron spectrometer (XPS) of the German Thermosphere company (voltage:15 kV, current: 15 mA). The elemental analyzer (Vario EL) quantitatively determined the N and O elements in the sample. The SSA and pore information of the material was obtained by Micrometrics ASAP2020 at 77 K.

Electrochemical measurement

The electrochemical performance of AFR-HCs was tested in 2032-type coin cells using 1 mol/L NaPF $_6$ in diglyme electrolyte, which were assembled in a glove box. The active materials, AFRs, polyvinylidene fluoride (PVDF) and super P with mass ratios of 8:1:1, were then mixed in N-methyl-2-pyrrolidone (NMP). The electrode slurry was coated on a Cu foil. The electrolyte of a full cell is the same as that of a half cell. The capacity ratio of anode/cathode electrodes was 1.2, and the ratio of mass loading between the anode and the cathode was maintained at 1:3; the capacity of the full cell was calculated based on the anode weight loading. The charge-discharge and galvanostatic intermittent titration technique (GITT) test was performed on the Neware CT-4000Q. The diffusivity coefficient of sodium ions (D_{Na^+}) is calculated, according to an equation of Fick's second law, by:

$$D = \frac{4}{\pi \tau} \left(\frac{m_B V_M}{M_B S}\right)^2 \left(\frac{\Delta E_S}{\Delta E_T}\right)^2 \tag{1}$$

Cyclic voltammetry (CV) curve and electrochemical impedance spectroscopy (EIS) tests were carried out using a CHI660 electrochemical workstation.

The specific capacity (C_i) , specific energy (E), and specific power (P) of full cells are calculated based on the supplementary equations (2)-(4), where E (Wh·kg⁻¹) represents the specific energy. T (h) is the total time of the discharge. C_i is the specific capacity of cells. I (mA) is the current of the cell. P (W·kg⁻¹) is the specific power of the full cell. U_a (V) is the average discharge voltage of the cell, which is calculated from the ratio of specific energy (E) to specific capacity (C_i) . m_c and m_a (kg) are the mass of Na₃V₂(PO₄)₃ (NVP) in the cathode and HC in the anode, respectively.

$$C_t = \frac{I \times t}{m_c + m_q} \tag{2}$$

$$E = C_t \times U_a \tag{3}$$

$$P = \frac{E}{t} \tag{4}$$

RESULTS AND DISCUSSION

Materials characterization

In this work, the novel AFR sphere was fabricated by the polymerization between 3-aminophenol and formaldehyde; then, a series of HC materials with tailorable microstructures were designed and successfully produced using AFR as a precursor [Figure 1A][3]. The 3-aminophenol molecule contains both phenol and amine groups^[15], which reacts with formaldehyde to form benzoxazines and hydroxymethyl derivatives^[16]. Next, the benzoxazine and hydroxymethyl derivatives undergo condensation and dehydration to form dimers, which further evolve into polymer chains. Finally, the polymer chains would self-assemble and aggregate to form AFR microspheres. In the reaction, formaldehyde, serving as a curing agent, can interact with 3-aminophenol, and its dosage could further affect the polymerization kinetics of benzoxazine and hydroxymethyl derivatives, thus regulating the crosslinking degree of AFR. Therefore, the crosslinking degree enhancement of AFR can be achieved by increasing the polymerization kinetics induced by the content of formaldehyde, which is realized just by reducing the molar ratio of 3-aminophenol to formaldehyde. The thermogravimetric analysis (TG) was used to evaluate the crosslinking degree of AFR precursors [Figure 1B][10], and the residual carbon rates are 40% for 1.8-AFR-HC, 48% for 0.9-AFR-HC and 58% for 0.46-AFR-HC, respectively. This indicates that a lower molar ratio of aminophenol to formaldehyde leads to a higher crosslinking degree of the resin (0.46-AFR-precursor), eventually resulting in a higher carbon residue rate[11]. It has been reported that the carbon precursors with a higher crosslinking degree could hinder the rapid formation of conjugated carbon network during carbon pyrolysis, thus forming HC with a higher carbon residue rate^[14]. The crosslinking degrees for different samples were further analyzed through infrared spectroscopy (IR)[17,18]. As shown in Figure 1C, similar infrared characteristic peaks are illustrated on 0.46-AFR-precursor, 0.90-AFR-precursor and 1.80-AFR-precursor materials, all being pure AFR precursor infrared spectra without redundant peaks. Notably, the main peaks of C=C at 1,610 cm⁻¹ and methylene peaks at 1,460 cm⁻¹ and 1,480 cm⁻¹ are different for 0.46-AFR-precursor, 0.90-AFR-precursor and 1.80-AFR-precursor; the content of methylene and C=C in 0.46-AFR-precursor is higher than both 1.80-AFR-HC and 0.90-AFR-HC^[19,20], verifying that the crosslinking degree of AFR precursors can be tuned by the molar ratio of aminophenol to formaldehyde. Further, the peak of C-O-C ether bonds is not obvious, which indicates the polycondensation of 3-aminophenol and formaldehyde is mainly formed by methylene rather than ether bonds^[15]. Additionally, the crosslinking degree of the AFR precursor can be further evaluated using a long-term leaching method via a Soxhlet extractor. The productivity of AFR precursors is 92% for 0.46-AFR-precursor, 89% for 0.9-AFR-precursor and 83% for 1.8-AFR-precursor, respectively, which agrees well with the TG and IR results [Figure 1D]. The crosslinking degree of the AFR precursor plays a crucial role in shaping the morphology and microstructure of AFR-HC, suggesting the possibility of ultimately influencing its electrochemical performance [Figure 1E].

As mentioned, the sample morphologies are observable via SEM. As shown in Figure 2A-C, the morphologies of 0.46-AFR-HC, 0.90-AFR-HC and 1.80-AFR-HC samples are all microspheres with a size of around 500 to 900 nm^[16]. Notably, the agglomeration degree of HC microspheres gradually increased with rising crosslinking degrees of the resin precursor from 1.80-AFR-HC to 0.46-AFR-HC [Figure 2D-F and Supplementary Figure 1]. This observation supports the influence of the crosslinking degree of resin precursors on the morphology regulation. HRTEM images further display the microstructures of the asprepared samples [Figure 2G and H]; the 0.46-AFR-HC consists of a short-range ordered graphite domain features with a graphite domain length (L_a) of ~2.64 nm, a graphitic crystallite thickness (L_c) of ~1.40 nm, and a higher interlayer distance (d_{002}) of 0.388 nm. These characteristics are deemed favorable for the intercalation of Na⁺ ions^[2,16]. Additionally, the 0.90-AFR-HC and 1.80-AFR-HC samples illustrate straight short carbon layers featuring the average size of d_{002} (0.384 and 0.372 nm, respectively), the L_a (2.59 and

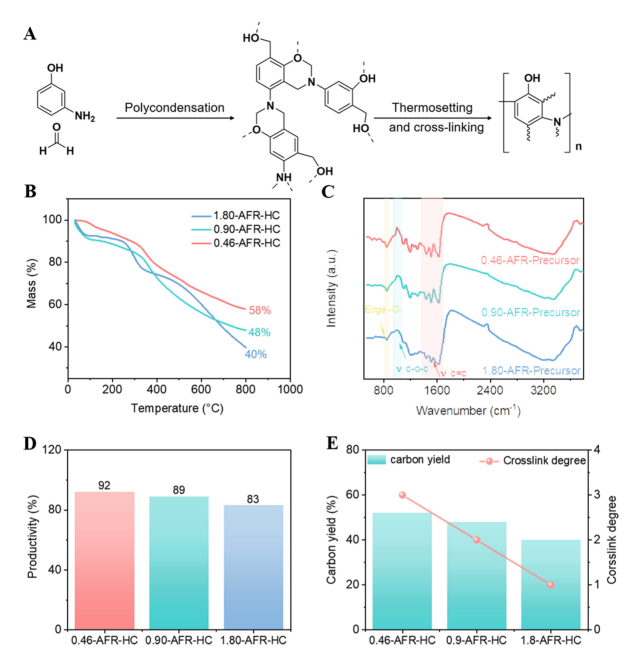


Figure 1. (A) Synthesis process of AFR; (B) TG results; (C) IR spectroscopy; (D) The yield of different samples was tested by a long-term leaching method; (E) Comparison of carbon yield and crosslinking degree. AFR: Aminophenol formaldehyde resin; TG: thermogravimetric analysis; IR: infrared spectroscopy.

2.56 nm, respectively), and the L_c (1.36 and 1.32 nm, respectively) [Figure 2H and I, Supplementary Table 1]. Elemental mapping by energy-dispersive X-ray spectroscopy (EDS) of 0.46-AFR-HC, 0.90-AFR-HC and 1.80-AFR-HC samples [Figure 2J, Supplementary Figure 2A and B] also show a uniform distribution of C, N and O.

The XRD can be performed to evaluate the graphitization degree and the microstructure parameters of HC materials^[21-24]. The XRD patterns of all AFR-HC materials show typical HC characteristic peaks at \approx 23° and \approx 42°, which are attributed to the (002) and (100) planes of HC, respectively [Figure 3A and Supplementary

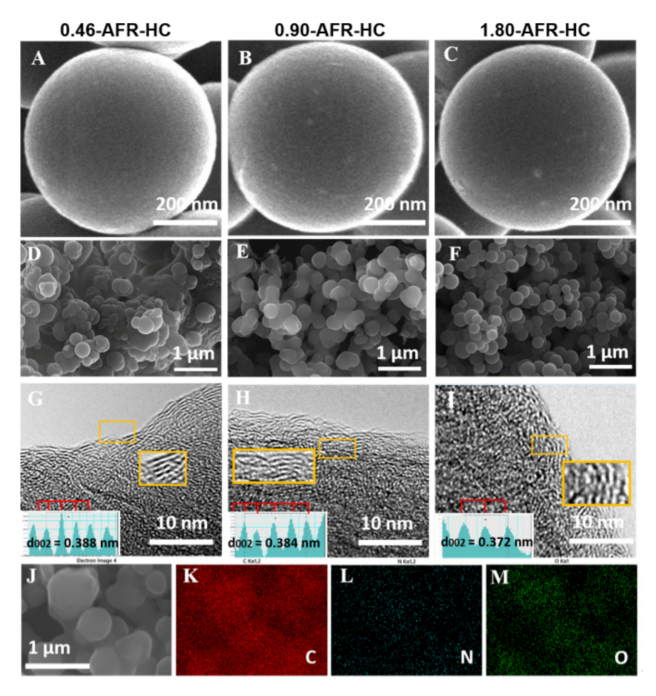


Figure 2. (A-F) SEM images; (G-I) HRTEM images; (J-M) EDS mappings of 0.46-AFR-HC, the (K), (L), and (M) figures belong to the EDS mappings of C, N and O elements respectively. SEM: Scanning electron microscopy; HRTEM: high-resolution transmission electron microscope; EDS: Energy dispersive spectrometer; AFR: aminophenol formaldehyde resin; HC: hard carbon.

Figure 3]^[25]. Surprisingly, the calculated d_{oo2} is 0.401 nm for 0.46-AFR-HC, 0.389 nm for 0.9-AFR-HC, and 0.374 nm for 1.8-AFR-HC, respectively. Additionally, the L_c gradually shifts toward higher values and L_a shifts toward lower values with an increasing resin precursor crosslinking degree. For example, 0.46-AFR-HC, 0.90-AFR-HC, and 1.80-AFR-HC samples feature with the L_a (3.01, 2.89, and 2.88 nm, respectively) and the L_c (1.46, 1.42, and 1.33 nm, respectively) are summarized in Supplementary Table 1. All these results are consistent with the TEM results [Figure 2G-I].

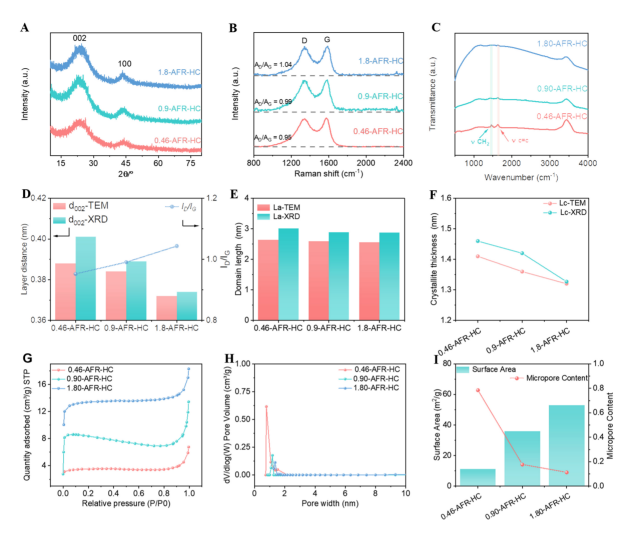


Figure 3. (A) XRD patterns; (B) Raman spectra; (C) FTIR spectroscopy; (D) Comparison of d_{002} and I_0/I_G ; (E and F) Comparison of L_a and L_c ; (G) N_2 adsorption-desorption isotherms at 77 K; (H) Pore diameter distributions based on N_2 adsorption-desorption isotherms; (I) Comparison of surface area and micropore content. XRD: X-ray diffraction; FTIR: Fourier transform infrared spectroscopy.

The microcrystalline variations of different samples are also confirmed by the Raman spectrum^[26]. As shown in Figure 3B, the Raman plots of 0.66-AFR-HC, 0.90-AFR-HC and 1.80-AFR-HC all contain two characteristic bands of 1,350 cm⁻¹ (D band) and 1,590 cm⁻¹ (G band), which correspond to the characteristic peaks of disordered or defective graphite (sp³) and crystalline graphite (sp²), respectively^[25,27]. Specifically, the Raman band can be fitted into four peaks located at 1,240, 1,350, 1,500, and 1,590 cm⁻¹, identified as D1, D, D2, and G, respectively [Supplementary Figure 4]. The D1 band (~1,240 cm⁻¹) is assigned to the disordered graphitic lattice of the stretching vibrations of a mixture of sp²-sp³ or sp³ bonds^[21]. The D2 band (~1,500 cm⁻¹) represents the amorphous carbon^[21]. It should be noted that the I_D/I_G represents the defect degree of HC, and its values are decreased in the order of 1.80-AFR-HC (1.043), 0.90-AFR-HC (0.99), and 0.46-AFR-HC (0.95) [Figure 3B and Supplementary Table 2], revealing that the defect degree of AFR-HC decreases slightly with the crosslinking degree of a precursor. The as-optimized 0.46-AFR-HC with a short-range ordered structure could be considered favorable for higher ICE. Fourier transform IR (FTIR) presents a similar trend to the Raman spectrum [Figure 3C]^[4]; intensity of methylene (1,460)

and 1,480 cm⁻¹) and C=C (1,610 cm⁻¹) peaks in 0.46-AFR-HC is higher than both 1.80-AFR-HC and 0.90-AFR-HC. This indicates higher crystallinity HCs can be obtained by increasing the crosslinking degree of the precursor. The microstructure parameters for all the samples were further summarized in Figure 3D-F, Supplementary Tables 1 and 2.

 N_2 adsorption was used to detect pore structure and SSA of HC [Figure 3G and H, Supplementary Table 3]^[28,29]. Interestingly, the crosslinking degrees of AFR-precursors directly affect the SSA and the nanopore structure of HC^[25]. The SSA values are 11.18 m²·g⁻¹ for 0.46-AFR-HC, 35.73 m²·g⁻¹ for 0.9-AFR-HC, and 52.89 m²·g⁻¹ for 1.8-AFR-HC, respectively. This indicates that the SSA of AFR-derived HC decreases as the precursor crosslinking degree increases. Notably, the micropore sizes of 0.46-AFR-HC mainly distributed from 0.5 to 2.0 nm while exhibiting a lower surface area [Figure 3H and Supplementary Table 3]. This indicates that as the crosslinking degree of the precursor increases, a substantial portion of the open pores in 0.46-AFR-HC becomes covered, leading to a higher proportion of closed pores and a lower surface area^[14,16]. The moderate SSA and increased number of closed pores contribute to the higher ICE and capacity of electrodes^[16,29].

XPS was employed to characterize the chemical composition and elemental state of the samples [Supplementary Figures 5-8]^[3]. The full spectra demonstrate that the samples of 0.46-AFR-HC, 0.90-AFR-HC, and 1.80-AFR-HC are mainly composed of C, O, and N elements [Supplementary Figure 5], however, with varying contents. It is noteworthy that 0.46-AFR-HC shows an increase in carbon content (93.29 at%) compared to 0.90-AFR-HC (92.63 at%) and 1.8-AFR-HC (92.07%) samples [Supplementary Table 4], indicating that the carbon content and the compaction density of HC could be regulated by varying the crosslinking degree of the precursor [Supplementary Table 5]. It has been proven that anode materials with a higher compaction density would benefit the construction of high energy density electrodes^[28-31].

All these observations illuminate the connection between the structures of the precursor and the local microstructure of the obtained HC. Notably, the optimized 0.46-AFR-HC exhibits short-order carbon layers with a larger d_{002} value, appropriate crystallinity, well-defined micropores, and lower SSA. These characteristics are deemed favorable for the Na⁺ filling in micropores and intercalation in graphite layers [Figure 3I]^[32-35].

Sodium storage performance

The electrochemical performances of the AFR-HCs were tested using Na||AFR-HC half cells in diglyme-based electrolytes^[31,32]. Obviously, the 0.46-AFR-HC electrode demonstrates a salient enhancement in the reversible capacity, ICE and rate capability. The galvanostatic charge/discharge curves (GCDs) of the first cycle for AFR-HCs show a representative charge and discharge process of HC, which display a slope region above 0.1 V and a plateau region below 0.1 V [Figure 4A]^[10,36-38]. During the first cycle, the discharge/charge capacities are 383/547, 348/440, and 271/363 mAh·g¹ at 0.05 A·g⁻¹ for 0.46-AFR-HC, 0.90-AFR-HC, and 1.80-AFR-HC electrodes, causing the ICE of 82%, 79%, and 73%, respectively [Figure 4A]^[37]. The superior capacity and ICE could be related to the well-regulated microstructure and smallest surface area of 0.46-AFR-HC.

Moreover, rate performance was also conducted [Figure 4B], and the corresponding GCD curves at different rates are shown in Supplementary Figure 9. Clearly, 0.46-AFR-HC displays reversible capacities of 383, 253, and 203 mAh·g⁻¹ at current densities of 0.1, 5.0, and 10.0 A·g⁻¹, respectively, far exceeding 0.90-AFR-HC and 1.80-AFR-HC electrodes. Even at a large current density as high as 20 A·g⁻¹, 0.46-AFR-HC still can deliver the discharge capacity of 140 mAh·g⁻¹, indicating that the well-regulated 0.46-AFR-HC is

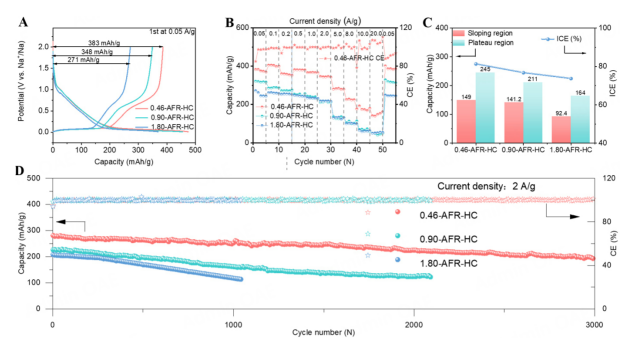


Figure 4. (A) The discharge-charge curves at the first cycle; (B) Rate performances; (C) Comparison of ICE and plateau/sloping capacities; (D) Cycling stability. ICE: Initial coulombic efficiency.

beneficial for fast Na⁺ reaction kinetics. Importantly, the plateau capacity ratio of 0.46-AFR-HC is retained by 62% (245 mAh·g⁻¹) at 0.05 A·g⁻¹, while the proportions are retained by 59% and 56% in 0.90-AFR-HC (211 mAh·g⁻¹) and 1.80-AFR-HC (164 mAh·g⁻¹) [Figure 4C], and the larger plateau region ratio would be advantageous for the construction of the full cell^[35]. Furthermore, after 1,000 cycles, the 0.46-AFR-HC electrode still exhibits a specific capacity of 268.4 mAh·g⁻¹ at 2 A·g⁻¹ with a high capacity retention of 94% [Figure 4D], correlating well with the SEM and TEM images for 0.46-AFR-HC after cycles [Supplementary Figures 10 and 11]. This is significantly improved compared with 0.90-AFR-HC for 70% (158.2 mAh·g⁻¹ at 2 A·g⁻¹) and 1.80-AFR-HC for 54% (115.3 mAh·g⁻¹ at 2 A·g⁻¹) after 1,000 cycles. To investigate the superiority of the AFR-HC on sodium storage behavior, the other types of half cells were also assembled, using pitch-based HC^[7,12] and biomass-based HC^[35,36] as anodes and sodium metal foil as counter electrodes. Among these materials, 0.46-AFR-HC possesses the highest rate capacity of 253 mAh·g⁻¹ under 5 A·g⁻¹ compared with biomass-based HC (54 mAh·g⁻¹ at 5 A·g⁻¹). The high capacity, rate capability, cycling stability, and ICE exhibited by 0.46-AFR-HC outperform the control samples and other types of HCs, indicating that 0.46-AFR-HC is a promising anode material for SIBs^[37].

Sodium storage behavior exploration and the microstructural parameters-performance

The better electrochemical performances regarding rate performance, capacity and cyclic stability make 0.46-AFR-HC electrodes promising for practical SIBs and, meanwhile, prompt us to explore their kinetics behavior. CV tests were conducted to reveal the Na storage behavior of 0.46-AFR-HC, 0.90-AFR-HC and 1.80-AFR-HC samples within charge/discharge processes $(0.01\sim2.0\text{ V})^{[15]}$. In Supplementary Figure 13, a distinctive sharp peak at \approx 0.1 V observed at 0.2 mV·s⁻¹ belongs to the plateau region (< 0.1 V)^[39]. This can be attributed to the Na⁺ filling in micropores and intercalation in graphite layers^[1]. The enhanced rate capability of the AFR-0.46-HC electrode is evident from the CV curves^[24]. Notably, the decision step of kinetic behavior for the interface reaction was assessed by applying the equation I = av^b to fit the b values, a commonly used approach for kinetic evaluation^[24,40]. In Supplementary Figures 14 and 15, excellent linear fit

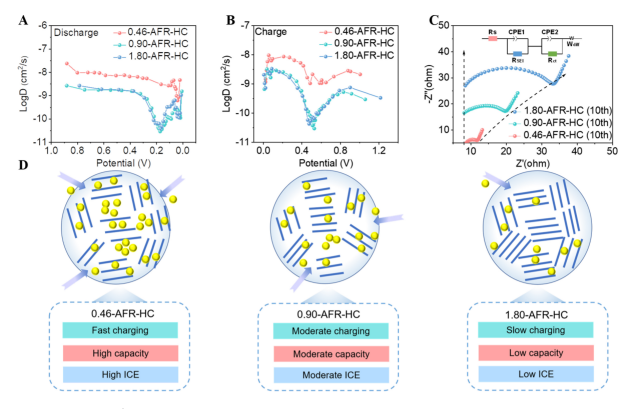


Figure 5. (A and B) Na⁺ diffusion coefficients calculated from the GITT potential profiles under discharge/charge processes; (C) EIS analysis after ten cycles; (D) Sodium storage diagram of AFR-HCs. GITT: Galvanostatic intermittent titration technique; EIS: electrochemical impedance spectroscopy; AFR: aminophenol formaldehyde resin; HCs: hard carbons.

results are observed for all three samples; the fitted b value for 0.46-AFR-HC (0.58) is larger than that for 0.90-AFR-HC (0.56) and 1.80-AFR-HC (0.55) [Supplementary Figure 15A-C]. The b value is between 0.5 and 1.0. This means that both diffusion control and capacitance processes may contribute to the charge storage of 0.46-AFR-HC^[24,30].

Furthermore, a GITT test was performed to further analyze sodium ion diffusion coefficients (D_{Na}^{+}) in 0.46-AFR-HC, 0.90-AFR-HC and 1.80-AFR-HC electrodes [36,39]. It was found that the D_{Na}^{+} values of the 0.46-AFR-HC electrode (10^{-7,81} to 10^{-9,50}) are higher than the 0.90-AFR-HC (10^{-8,75} to 10^{-10,54}) and 1.80-AFR-HC (10^{-8,64} to 10^{-10,47}) at all potentials, especially in the low voltage region (10^{-9,50} vs. 10^{-10,54} vs. 10^{-10,47}) [Figure 5A and B]. The incredible enhancement of D_{Na}^{+} in 0.46-AFR-HC at all potentials could also explain its better rate performance [3]. Such superior electrochemical performance could be derived from the well-regulated 0.46-AFR-HC microstructure. To gain insight into the interfacial reaction kinetics of 0.46-AFR-HC, 0.90-AFR-HC, and 1.80-AFR-HC electrodes, the EIS measurements were performed [37,41]. Selected Nyquist plots for these samples are presented in Figure 5C [37]. These results indicate that the resistances originating from the solid electrolyte interphase (SEI) layer (R_{SEI}) and charge transfer (R_{CI}) in 0.46-AFR-HC electrodes are both lower than those in 0.90-AFR-HC and 1.80-AFR-HC electrodes [36,39,41], that is, 2.6 Ω vs. 3.9 Ω vs. 7.3 Ω for R_{SEI} and 3.6 Ω vs. 7.9 Ω vs. 11.8 Ω for R_{CI} , respectively [Supplementary Table 8].

We conclude that the enhanced electrochemical performance of 0.46-AFR-HC can be attributed to its ideal microstructure. Firstly, there is a certain linear relationship between the resin crosslinking degree of a precursor and the microstructure of as-obtained HC; that is, the greater the degree of resin crosslinking, the higher carbon yield, the higher graphitization degree, the larger d_{002} and lower SSA. Promisingly, the

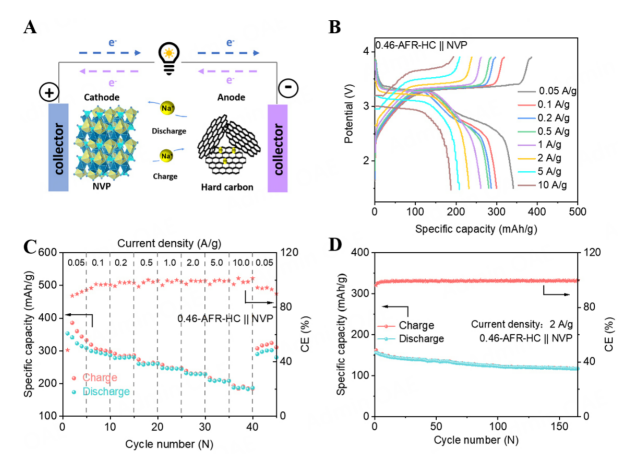


Figure 6. (A) Schematic of sodium-ion full-cell; (B) The GCD profiles under different rates; (C) Rate performance; (D) Cycling stability. GCD: Galvanostatic charge/discharge curve.

optimized 0.46-AFR-HC material features well-regulated microstructures (0.5~2.0 nm), less defect (lower I_D/I_G of 0.95), lower SSA (11 m²·g⁻¹), and larger d_{oo2} (0.388 nm). These characteristics fulfill the requirements for the intercalation and Na clustering of Na⁺ during charge and discharge processes [Figure 5D].

Electrochemical performance of AFR-HC||NVP full cell

Full cells were assembled to further verify the practical application of 0.46-AFR-HC material; the full-cell configuration is shown in Figure 6A, and NVP is selected as a cathode^[36]. Before assembling the full cell, the 0.46-AFR-HC anode first underwent pre-sodium through an electrochemical process to facilitate the formation of the SEI^[42,43]. The full cell shows a high charge capacity of 365 mAh·g⁻¹ at 0.05 A·g⁻¹ (based on the active mass of the anode material, Figure 6B)^[1], with a specific energy density of 256 Wh·kg⁻¹. Furthermore, it delivers good rate performance, maintaining a capacity of 158 mAh·g⁻¹ at 10 A·g⁻¹ [Figure 6C]. This corresponds to a power density of 6.43 kW·kg⁻¹. Additionally, it exhibits stable capacity over 200 cycles, maintaining approximately 110 mAh·g⁻¹ at 2 A·g⁻¹ (80% capacity retention) [Figure 6D]. As a result, the superior sodium storage properties of 0.46-AFR-HC in half/full cells illustrate that it is a promising carbon material and can be used in SIBs.

CONCLUSIONS

In summary, we have fabricated a AFR-based HC material with better performance, higher ICE and capacity than other control samples by adjusting the crosslinking degree of the AFR precursor. The 0.46-

AFR-HC electrode presents higher reversible capacity (383 mAh·g⁻¹ at 0.05 A·g⁻¹), ICE (82%) and good rate capability (140 mAh·g⁻¹ at 20 A·g⁻¹) compared to the control samples and other types of HCs. Further, the full cell with 0.46-AFR-HC as an anode and NVP as a cathode demonstrates high energy/power densities of 250 Wh·kg⁻¹ and 6.43 kW·kg⁻¹, respectively. The superior electrochemical performance observed in the experiment can be attributed to the consistency between the crosslinking degree of the precursor and the microstructure parameters of HC. Specifically, a resin precursor with a higher crosslinking degree tends to produce the optimized 0.46-AFR-HC with higher carbon yield, a larger d_{002} , fewer defects, and a lower surface area. These characteristics are deemed promising for constructing high-performance anode materials. This work proposes a reliable design for high performance HCs and provides a simple and ecofriendly method to prepare HCs with an ideal microstructure in large quantities.

DECLARATIONS

Authors' contributions

Wrote the article: Lu Z

Made substantial contributions to conception and design of the study and performed data analysis and interpretation: Lu Z, Yin X

Performed experiment: Ji Y, Li C, Wang R, Li S, Yuan T

Performed data acquisition and provided administrative, technical, and material support: Zhao Y

Availability of data and materials

Supporting information is available from the authors.

Financial support and sponsorship

This work is supported by the National Natural Science Foundation of China (22179077), National Natural Science Foundation Youth Fund (22209104), Shanghai Science and Technology Commission's "2020 Science and Technology Innovation Action Plan" (20511104003), and Natural Science Foundation in Shanghai (21ZR1424200).

Conflicts of interest

All authors declared that there are no conflicts of interest.

Ethical approval and consent to participate

Not applicable.

Consent for publication

Not applicable.

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