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Article

Hydrothermal Carbonization of Cellulose with Ammonium Sulfate and Thiourea for Production of Supercapacitor Carbon

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Abstract: Hydrothermal carbonization of cellulose was examined at 240 °C for 1 h. Ammonium sulfate and thiourea were selected as the doping sources of inorganic nitrogen and organic nitrogen for the preparation of supercapacitor carbon. The effect of boric acid on the properties of the resulting hydrochar after KOH activation was examined. The results showed that the proportion of functional groups and the specific surface area of the activated hydrochar would be reduced after the addition of boric acid, and the pore-forming process of the micropores would be inhibited. The hydrochar obtained from the reaction of cellulose and organic nitrogen compounds had better pore size distribution and electrochemical properties after activation. The largest specific surface area (952.27 m² g⁻¹) was obtained when only thiourea was used as the only doping source. In the three-electrode system, the specific capacitance resulting activated hydrochar reached 236.25 F g⁻¹ at a current density of 1 A g⁻¹. After 20,000 cycles of charging and discharging at a current density of 10 A g⁻¹, the capacitance retention rate reached 99.96%. Therefore, this study proves that the supercapacitor carbon with good electrochemical properties could be obtained by the direct reaction of cellulose with organic nitrogen compounds.

Keywords: hydrothermal carbonization; cellulose; ammonium sulfate; thiourea; doping; supercapacitor carbon

1. Introduction

In human life activities and industrial production, fossil fuels occupy a large amount of share. According to the World Energy Council, the world will need twice as much energy in 2050 as it consumes today.[1] Therefore, there is a need for a new energy source that can replace fossil fuels. Replacing conventional energy systems by implementing renewable energy systems is considered a crucial step towards sustainable development.[2,3]

Renewable energy requires an environmentally friendly and efficient energy storage device. The energy technologies that can be selected include solar energy, wind energy, tidal energy, geothermal energy, etc. These new energy sources must consider the influence of factors such as geographical location and climatic conditions.[4–6] After decades of development, supercapacitors have attracted extensive attention due to their high power density, fast charge and discharge speed, and long-term cycle stability compared with traditional secondary batteries.[7] These outstanding properties make supercapacitors a promising carrier for use in portable electronics, hybrid electric vehicles, etc., helping to build renewable energy systems.[8–11]

However, a significant reason currently limiting supercapacitors is their low energy density.[12–14] Some researchers have pointed out that using new materials, such as nanoscale metal oxides,[15,16] conductive polymers,[17] and heteroatom doping (B, N, S, P, O) carbon, etc.[18–22] could improve the energy density. For example, nitrogen enrichment is an effective way to achieve the N-doping of supercapacitor carbon materials. The nitrogen functional group in the supercapacitor carbon materials can improve the electron transfer rate and stability, reduce the equivalent series

resistance, and thus improve the electrochemical performance of the material.[23,24] Many reports have also pointed out that supercapacitor carbon materials have considerable advantages in many industrial applications due to their large specific surface area and porous structure. High porosity is an important parameter that can provide adsorption sites for ion storage and promote ion transport.[25–30]

There are many kinds of raw materials for the preparation of supercapacitor carbon materials, among which biomass has become a hot spot for its advantages of high yield, renewable, low pollution, zero carbon dioxide emission and so on. At present, the conversion of biomass to supercapacitor carbon materials is mainly achieved by a thermochemical method [31–35], among which hydrothermal carbonization(HTC) shows much promise as it can directly use wet biomass and produce high economic benefits[36]. During the HTC process, the hydrogen bond force between water molecules is weakened, and both the ionization constant and the concentration of H^+ in the liquid phase will increase.[37] At the same time, the pH value of the liquid phase will also increase with the addition of organic acids, which will promote the hydrolysis of oligosaccharides and other substances, thereby further promoting the growth of oxygen-containing functional groups on the surface of the resulting hydrochar.[38] However, the doping sources or the organic acids produced during the HTC process of the biomass are limited, and additional organic acids need to be added.

There have been a large number of reports on improving the performance of carbon materials prepared in various aspects by adding additives. For example, Susanti et al. added a small amount of citric acid in the HTC of salacca peel to increase the specific surface area of carbon materials.[39] Deng et al. added an N source to activated carbon to prepare electrodes with high spectral surface area and dense nanopores.[40] Guo et al. removed heavy metals through HTC by adding urea as a nitrogen source.[41] Huang et al. prepared carbon materials for supercapacitor use by adding melamine as a nitrogen source during the HTC of bamboo shoot shells[42]. Wu et al. used urea as a nitrogen source during the HTC of carboxymethyl cellulose to prepare spherical carbonaceous adsorbents (CSn) with micro-porosity used for carbon dioxide capture capacity.[43]

The type of doping source has a great influence on the electrochemical properties of the material. Among the doping sources, boric acid is widely used as an important doping source of B, which could improve the electrochemical performance of supercapacitor carbon, and the co-doping of B and N could further improve the electrochemical properties of the material. Zhao et al. proposed a low-cost and simplified strategy for the preparation of B,N-codoped porous C using freeze-drying chitosan-boric acid aerogel beads followed by a tube furnace carbonization. The B doping significantly affected the distribution of N species in the carbon material. The resulting doped carbon presented a good wettability and is conducive to the diffusion of electrolytes and the transport of ions.[44] Liang et al. developed an efficient strategy to carbonize the chitosan in the presence of boric acid, and the resulting carbon showed a high specific capacitance, large operation voltage and excellent cycle stability for supercapacitors.[45]

Therefore, in this present study, HTC of cellulose was examined with the addition of an inorganic N source (ammonium sulfate) or organic N source (thiourea) for the production of supercapacitor carbon. The effect of B addition on the specific surface area, pore structure, and electrochemical performance of the resulting carbon materials prepared by adding organic and inorganic nitrogen sources was elaborated in detail. Finally, the electrochemical performances of the activated carbon material were characterized by cyclic voltammetry (CV), galvanostatic charge-discharge (GCD) test, electrochemical impedance spectroscopy (EIS) and cycle life measurement.

2. Materials and Methods

2.1. Materials

α -Cellulose powder (CL) was purchased from Shanghai Macklin Biochemical Technology Co., Ltd.(Shanghai, China). Thiourea (TU), ammonium sulfate (AS), and boric acid (BA) were purchased from Tianjin Damao Chemical Reagent Factory (Tianjin, China). Deionized water was prepared in the lab, which has a conductivity of less than 10 μ s/cm. Potassium hydroxide (KOH), hydrochloric

acid (HCL), and acetylene black BP2000 were purchased from Tianjin Kernel Chemical Reagent Co. Ltd.(Tianjin, China). The polytetrafluoroethylene (PTFE) binder was provided by Shanghai McLean Biochemical Technology Co. Ltd.(Shanghai, China).

The HTC reaction was performed using a mechanical reactor MSG100-P5-T3-SS1-SV-R (Anhui CHEM^N Instrument Co., Ltd., China) with a volume of 100 mL. This reactor is equipped with a stirrer. The heating power of the reactor was 1.1 kW. The rated temperature and pressure of the reactor are 300°C and 20.7 MPa, respectively.

2.2. HTC

5 g CL, 0.5 g TU or AS, 0.5 g BA(when needed), and 40 mL deionized water were loaded into a 100 mL beaker and mixed uniformly to form a paste. This paste was transferred into the 100 mL stainless-steel reactor and sealed. The switch was turned on to start the heating. About 30 min later(depending on the reaction temperature), the temperature inside the reactor reached 240 °C, and the reaction time was counted at this point. After 1 h reaction time was completed, the reactor was placed into a water bath. Once the temperature inside the reactor reached room temperature, it was taken out of the water bath. If there were gas products formed in the reactor, the valve needed to be opened to release the gas products first. The solid (hydrochar) and the aqueous phase were transferred into a büchner funnel for separation. The hydrochar was dried in an oven at 105 °C for 12 h. The yield of hydrochar was equal to the mass of hydrochar divided by the total mass of the material initially added into the reactor. The hydrochar (HC) produced from HTC of CL, CL+TU, CL+AS, CL+TU+ BA, CL+AS+BA were named HC-CL, HC-CL-TU, HC-CL-AS, HC-CL-TU-BA, and HC-CL-AS-BA, respectively.

2.3. Activation of HC

0.5 g HC and 1.5 g KOH were loaded into a 250 mL beaker, and 200 mL deionized water was added into the beaker and subjected to ultrasonication for 20 min. The HC was soaked in the KOH solution for 24 h. The soaked HC was separated by a büchner funnel and dried at 105 °C for 12 h. The soaked HC was loaded into a porcelain boat and placed into a tube furnace, which was heated to 800°C at a heating rate of 3°C min⁻¹ under an Ar environment at a flow rate of 20 mL min⁻¹ and stayed at 800°C for 1 h. After the activation, the activated hydrochar (AHC) was cooled to room temperature under an Ar atmosphere and soaked in 100 mL 1M HCL solution for 12 h. Subsequently, the AHC was washed with deionized water until the pH of the filtrate reached 7 and dried at 105 °C for 12 h. The activated hydrochar (AHC) produced HC-CL, HC-CL-TU, HC-CL-AS, HC-CL-TU-BA, and HC-CL-AS-BA were named AHC-CL, AHC-CL-TU, AHC-CL-AS, AHC-CL-TU-BA, and AHC-CL-AS-BA, respectively.

2.4. Characterization of AHC

The specific surface area (SSA) and pore size distribution of AHCs were determined by N₂ adsorption-desorption isotherm at 77 K through an JW-BK100 equipment (Beijing JWGB Instruments Co., Ltd.). The morphology and microstructure of samples were obtained by a Gemini SEM 500 scanning electron microscopy (Carl Zeiss AG, Oberkochen, Germany). X-ray diffraction was obtained by XRD-6100(Shimadzu Corporation, Nakagyo-ku, Kyoto, Japan). The infrared spectrum was obtained by Nicolet iS50(ThermoFisher Scientific, Waltham, MA, USA). The thermogravimetric analysis was obtained by STA449F5 (Netzsch, Selb, Germany). X-ray photoelectron spectroscopy (XPS) was acquired on ESCALAB 250Xi (Thermo Scientific, USA). Elemental analysis was obtained by Vario EL cube (Elementar Americas Inc., Ronkonkoma, NY, USA).

2.5. Electrochemical performance test of AHC

The electrochemical performances of individual electrodes were evaluated by an electrochemical workstation model CHI1030B (Shanghai Chenhua Instrument Co. Ltd., Shanghai, China). A three-

electrode device was used as the test system under an ambient atmosphere, where the platinum sheet electrode served as the counter electrode and Hg/HgO for the reference electrode.

The AHC, acetylene black, and PTFE were mixed at a mass ratio of 16:3:1 and dissolved in ethanol. The mixture was subjected to sonication at a power of 120 W for 20 min and dried at 105°C for 5 h to totally remove the ethanol. The dried mixture was pressed into flakes at uniform mass under a pressure of 2 MPa and cut into circular flakes with a diameter of 10 mm. Cut two pieces of nickel foam into circular slices with a diameter of 15 mm. The 10 mm diameter electrode disc and a nickel sheet were placed between the two pieces of nickel foam discs and extruded by a tablet press at a pressure of 4 Mpa to make a working electrode. The electrolyte used in the testing process was 6 M KOH. Electrochemical tests included cyclic voltammetry (CV) at scan rates ranging from 10 to 200 mV s⁻¹ for CV curves, galvanostatic charge/discharge (GCD) at current densities between 1 and 20 A g⁻¹ for GCD curves, and electrochemical impedance spectroscopy (EIS) at scan frequencies between 0.01 Hz and 100 kHz. On the Land cell test program, the cycling stability was examined through 20,000 cycles at 10 A g⁻¹. As shown in Equation (1), the specific capacitance based on GCD measurements was determined.

$$C = \frac{I\Delta t}{m\Delta V} \quad (1)$$

where C is the gravimetric specific capacitance of electrodes (F g⁻¹), I is the discharge current (A), Δt is the discharge time (s), m is the mass of the active material (g), ΔV is the voltage range (V).

3. Results and Discussions

3.1. Yield of the HC and elemental analysis of the HC and AHC

The HC yield obtained by direct HTC of CL was 52.8 wt%. Whether AS or TU was added, the HC yield obtained after HTC of CL was slightly increased, and was maintained at about 53.5 wt%. The reactions occurred between CL and AS or TU, and the addition of BA slightly reduced the hydrochar yield to about 53 wt%. Figure 1 shows the reaction path of CL carbonization under hydrothermal conditions.[46] The aqueous solution of BA is acidic, which is not conducive to the carbonization reaction of CL or its hydrolysates. In addition, an acidic condition also facilitated the hydrolysis of oligosaccharides into organic acids and the decomposition of HMF into acids/aldehydes and phenols, and thus reduce the hydrochar yield.

Table 1 lists the elemental composition of HC and AHC produced from different reaction feedstocks. Compared with CL, the hydrochar obtained from the HTC of CL has higher C content (71.14 wt%) and lower H(4.35 wt%) and O(24.07 wt%) contents, indicating that the dehydration reaction occurred during the carbonization process of CL, as described in the reaction path in Figure 1. This is also illustrated by the low H/C and O/C of the hydrochar obtained from the HTC of CL. The addition of AS or TU promoted the HTC of CL because the resulting hydrochar has a higher C content than that of pure CL. Compared with CL, AS and TU have a higher H content, so the resulting hydrochar has a higher H content after adding AS and TU. The reaction between CL and AS or TU resulted in higher contents of N and S in the hydrochar, and the higher the N and S in the additive, the higher the N and S in the hydrochar. It should be noted that the S content of hydrochar obtained by the reaction of CL and AS is only 0.35 wt%, while the content of hydrochar and S obtained by TU is 4.54 wt%, indicating that organic sulfur is more likely to react with CL. Since the contents of C, H, N and S of the obtained hydrochar increased after adding AS or TU, the responding O content decreased. The addition of BA promoted the reaction of CL with AS or TU, which further increased the N and S contents of the obtained hydrochar. The H/C of hydrochar is reduced, which also shows that BA promoted the carbonization reaction of CL with AS or TU. At the same time, BA also participated in the reaction, resulting in an increased O content of hydrochar and a small amount of boron.

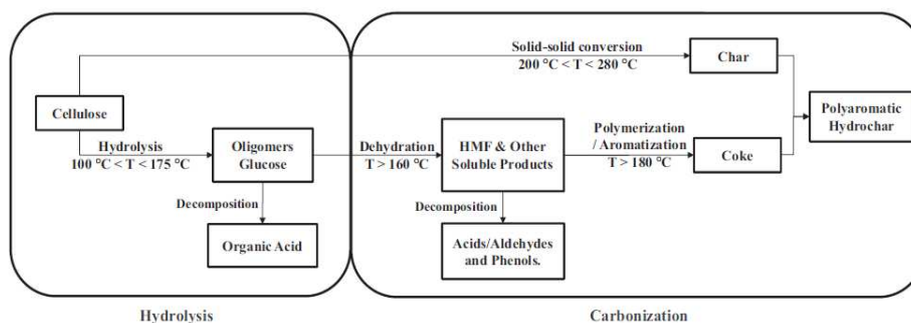


Figure 1. The reaction path of CL carbonization under hydrothermal conditions.

Since the activation was carried out in a high temperature and inert atmosphere, all activated hydrochar have a higher C content and a lower H and O contents than hydrochar. The H/C ratio of all activated hydrochar was also as low as 0.26-0.29. The contents of N and S in the activation process of hydrochar also has different degrees of loss, resulting in the contents of N and S in the activated hydrochar were lower than that of hydrochar.

Table 1. Elemental analysis(wt%) of the HC and AHC.

Sample	C	H	N	S	O	H/C	O/C
CL	66.61	4.44	0.00	0.00	28.55	0.06	0.43
HC-CL	70.14	4.35	0.00	0.00	24.07	0.06	0.34
HC-CL-AS	66.98	5.12	3.63	0.35	15.36	0.10	0.66
HC-CL-TU	52.34	6.20	4.67	2.92	12.94	0.08	0.37
HC-CL-AS-BA	63.21	4.66	4.92	4.54	15.81	0.07	0.34
HC-CL-TU-BA	67.54	4.81	4.31	0.37	16.14	0.07	0.35
AHC-CL	76.11	1.27	0.00	0.00	21.91	0.02	0.29
AHC-CL-AS	78.59	1.88	2.84	0.32	11.69	0.02	0.15
AHC-CL-TU	70.32	1.94	3.25	2.23	21.46	0.03	0.31
AHC-CL-AS-BA	78.56	1.74	4.02	3.97	10.27	0.02	0.13
AHC-CL-TU-BA	79.34	1.79	3.77	0.32	12.73	0.02	0.16

3.2. BET Analysis of the AHC

The porosity of AHC-CL, AHC-CL+AS, AHC-CL+TU, AHC-CL+AS+BA, AHC-CL+TU+BA was investigated using N₂ adsorption/desorption isotherms at 77 K. From Figure 2(a), it can be seen that none of the isotherms of AHC-CL+TU has obvious hysteresis loops, showing typical I-type behavior. But AHC-CL+AS+BA, AHC-CL+TU+BA, and AHC-CL+AS showed obvious hysteresis loops. At lower relative pressures P/P_0 ($P/P_0 < 0.1$), nitrogen uptake increased dramatically, indicating that the as-prepared AHC has typical microporous properties. When the relative pressure P/P_0 was increased to 0.2, the uptake of nitrogen gradually increased, which was partly due to the micro/mesoporosity. Compared with AHC-CL+AS+BA and AHC-CL+TU+BA, AHC-

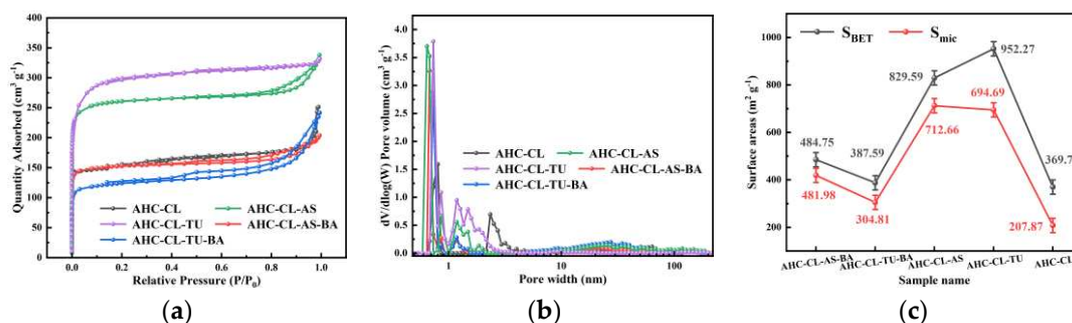
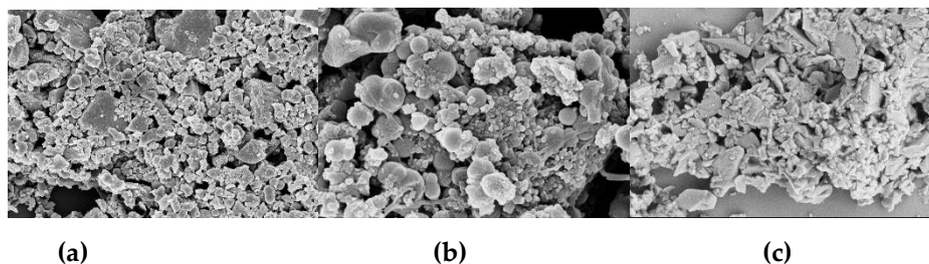


Figure 2. (a). Adsorption/desorption curves of AHC-CL, AHC-CL+AS, AHC-CL+TU, AHC-CL+AS+BA, AHC-CL+TU+BA under N_2 environment; (b). Pore size distribution curves of AHC-CL, AHC-CL+AS, AHC-CL+TU, AHC-CL+AS+BA, AHC-CL+TU+BA; (c). Specific surface area and micropore area of AHC-CL, AHC-CL+AS, AHC-CL+TU, AHC-CL+AS+BA, AHC-CL+TU+BA.

CL+AS and AHC-CL+TU obviously have better adsorption capacity, which indicates that AAM and AAT have a higher SSA. This may be because the boron element hinders the generation of pore structure during the HTC process, resulting in the reduction of SSA. At the same time, it can be seen that the effect of boron doping on doping TU is significantly higher than that on AS. The addition of BA would reduce the specific surface area of AHC doped with TU and AS, and the specific surface area of the experimental group doped with TU would be more obvious. It can be seen from Figure 2(b) that the number of micropores in the AHC after doping with boron is significantly reduced, which further illustrates that the doping of boron occupied the pores of the AHC, resulting in a decrease in the SSA of the AHC. As shown in Figure 3, the SEM images show that AHC-CL+TU and AHC-CL+AS have more pore structures than those of AHC-CL+TU-BA and AHC-CL+AS-BA.

As indicated in Figure 2(c), we can see that the SSA and micropore area of AHC are suppressed when boron is doped, especially when BA and TU were co-carbonized, the SSA and micropore area of the obtained AHC are significantly inhibited. When only TU acts, the SSA of the AHC reached $952.27 m^2 g^{-1}$, and the micropore area reaches $694.69 m^2 g^{-1}$. Under the action of only CL, the SSA of the AHC reached $829.59 m^2 g^{-1}$, and the micropore area reaches $712.66 m^2 g^{-1}$. When boron was added, the specific surface area of AHC-CL+AS+BA was reduced to $484.76 m^2 g^{-1}$, and its micropore area was reduced to $418.99 m^2 g^{-1}$; the specific surface area of AHC-CL+TU+BA was reduced to $397.59 m^2 g^{-1}$, and its micropore area was reduced to $304.81 m^2 g^{-1}$. This may be because the addition of BA occupied the micropore sites during the HTC process, resulting in a decrease in the micropore area of the HC sample. It can be seen that after boron doping, both the SSA and the micropore area were affected, and the most affected group was the group doped with TU. And when no boron is doped, the specific surface area of the AHC added with AS was much smaller than that of the AHC added with TU, and the prepared micropore area was almost the same, which proved that the addition of AS had a better ability to manufacture micropores ability. While removing the interference of boron, the AHC with AS addition also showed a stronger ability to create micropores.



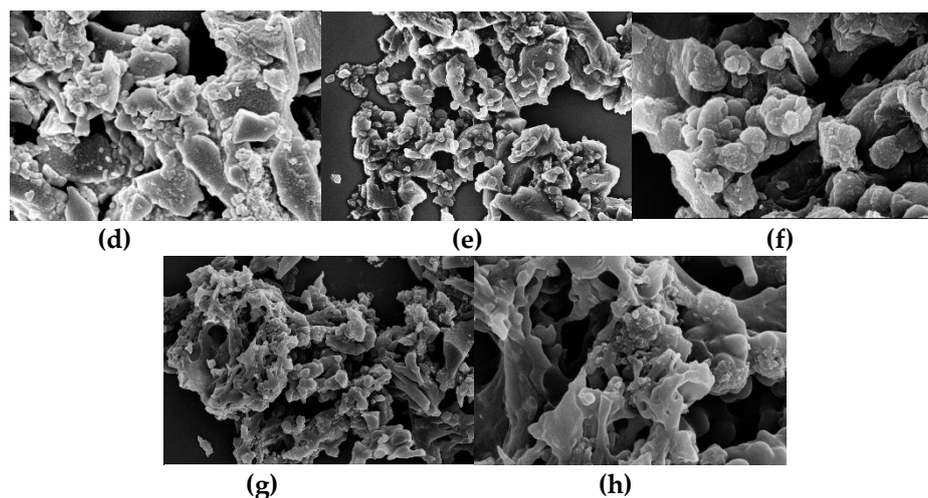
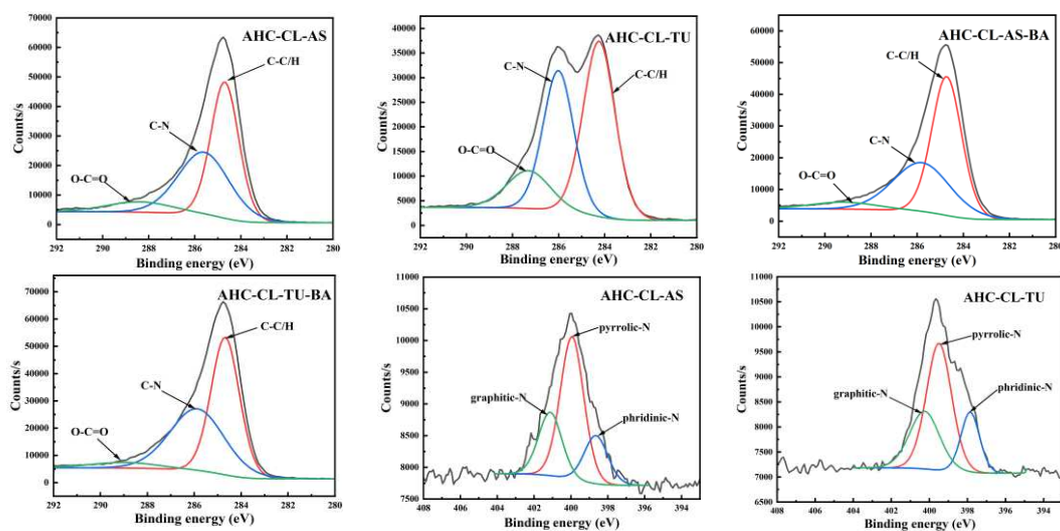


Figure 3. SEM images of (a, b). AAM; (c, d); AABM; (e, f). AAT; (g, h). AABT.

3.3. XPS Analysis of the AHC

To further probe the chemical state of the elements on the surface of the AHC, the X-ray photoelectron spectroscopy (XPS) of the AHC was used. Figure 3 shows all the results. The peaks around 286.1 eV and 287.6 eV represent C-N and O-C=O. Compared with AS, the C-N/C-O peak intensity of the TU-doped experimental group increased significantly. Moreover, the peak intensity of C=O in the experimental group doped with TU is obviously higher than that of doped with AS. It shows that compared with doped AS, AHC doped with TU would undergo a certain degree of decarboxylation reaction in the HTC reaction. It is further proved that the addition of TU could make the prepared AHC have more oxygen-containing functional group (OFGs), which could improve the wettability of AHC pores and make it easier for ions to enter the pores.

The main peaks appear at 399.1 eV and 398.3 eV correspond to pyrrole-N and pyridine-N, respectively. The increase in pyrrole-N in the AHC would prompt to generate more electrochemically active sites, thereby improving the faradaic capacitance by changing the electron distribution on the surface of the AHC. As shown in Table 1, TU-doped AHC has a higher N content, but the pyrrole-N peak intensity of AHC-CL+TU is smaller than that of AHC-CL+AS, indicating that not all N was converted into pyrrole-N during the HTC process. Instead, it existed in the AHC as other N form. AHC-CL+AS has more pyrrole-N, which indicates that this kind of carbon may has a better charge transfer performance and electron supply performance.



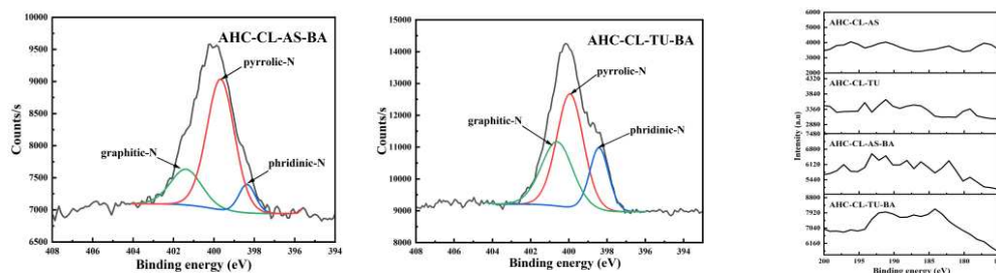


Figure 3. XPS spectra of AHC-CL+AS, AHC-CL+TU, AHC-CL+AS+BA, AHC-CL+TU+BA.

When the nitrogen and boron doping sources were added at the same time, the C-N peak intensity of AHC-CL+TU+BA is significantly higher than that of AHC-CL+AS+BA, indicating that B-assisted TU synergistic carbonization was more conducive to the formation of C-N. At the same time, the peak intensity of O-C=O in the B-assisted TU synergistic carbonization process was better than that in the B-assisted CL synergistic carbonization process, indicating that the B-assisted TU synergistic carbonization process could promote the decarboxylation reaction during the HTC process. During the process of CL/TU synergistic carbonization assisted by B addition, the increase of pyrrole-N content would induce more electrochemically active sites in the AHC, thereby improving the faradaic capacitance by changing the electron distribution on the surface of the AHC. The pyrrole-N peak intensity of AHC-CL+AS+BA is similar to that of AHC-CL+TU+BA. Although the AHC-CL+AS+BA and AHC-CL+TU+BA have similar N content, however, the intensity of the pyridine-N peak of AHC-CL+TU+BA was significantly higher than that of AHC-CL+AS+BA. During the process of adding TUs as the N source, after adding B as the doping source at the same time, the C-N peak intensity of the TU-doped experimental group was significantly reduced. It shows that the addition of B would inhibit the formation of C-N. Moreover, after adding B, the peak intensity of C=O also decreased significantly. It shows that the addition of B assists the process of TU synergistic carbonization, which would inhibit the decarboxylation reaction of this process. This would reduce the OFGs of AHC and reduce the wettability of AHC pores, and thus prevent ions from entering the pores. After adding B as a dopant source, the intensity of the pyrrole-N peak of AHC decreased. As indicated in Table 1, the N contents of the two groups of AHCs are similar, indicating that not all N were converted into pyrrole-N during the HTC or activation process, but exist in the AHC in the other form of N. The pyrrole-N content of AHC-CL+TU was higher, indicating that this sample may have better charge transfer performance and electron supply performance.

In the process of adding AS as N source, after adding B as doping source, the C-N peak intensity of the AS-doped experimental group was significantly reduced. It shows that the addition of B would inhibit the formation of C-N during the HTC and /or activation process. Moreover, after adding B, the peak intensity of C=O also decreased. It shows that the process of synergistic carbonization by adding B to assist TU will inhibit the decarboxylation reaction in this process, which is not conducive to the carbonization process. This process will reduce the OFGs of AHC, reduce the wettability of AHC pores, and thus prevent ions from entering the pores. After adding B as the doping source, the pyrrole-N peak intensity of AHC is similar. As shown in Table 1, the N content of AHC-CL+AS+BA is significantly higher than that of AHC-CL+ASA, indicating that not all N were converted into pyrrole-N during HTC and /or activation process, but existed in AHC in the other form of N.

3.4. Electrochemical Analysis of the AHC

In the three-electrode system, the electrochemical performance of AHC in 6 M KOH electrolyte was obtained by testing CV, GCD and EIS. In order to explore the change of electrochemical performance of AHC after the addition of B assisted TU and A S synergistically carbonized cellulose, the GCD and CV of AHC-CL+TU, AHC-CL+AS, AHC-CL+TU+BA, and AHC-CL+AS+BA were tested, respectively. As shown in Figure 4(a), all the AHC have good symmetrical triangular distributions, indicating good electric double layer capacitance and electrochemical reversibility. The AHC-CL+TU takes longer to charge and discharge. At a current density of 1 A g⁻¹, the mass specific

capacitance of AAT reaches 236.25 F g^{-1} . This indicates that AHC-CL+TU has a higher energy storage capacity.

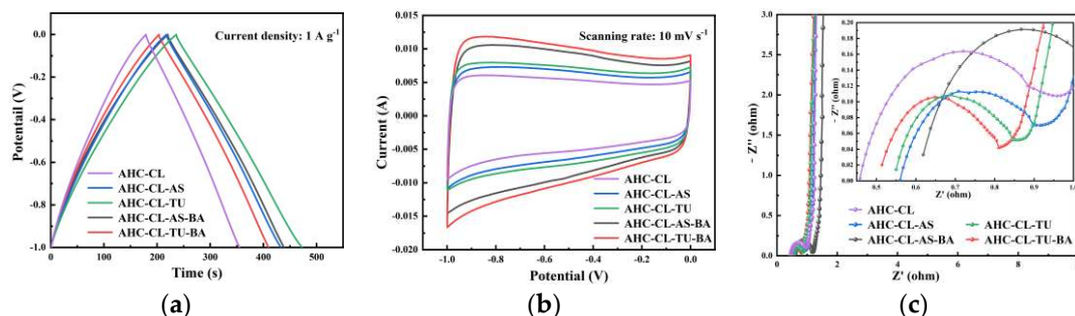


Figure 4. The electrochemical performance data of the three-electrode system using 6M potassium hydroxide solution as the electrolyte. (a). GCD curves of AHC-CL+TU, AHC-CL+AS, AHC-CL+TU+BA, and AHC-CL+AS+BA at current density of 1 A g^{-1} ; (b). CV curves of AHC-CL+TU, AHC-CL+AS, AHC-CL+TU+BA, and AHC-CL+AS+BA at a scan rate of 10 mV s^{-1} ; (c). Nyquist spectra of AHC-CL+TU, AHC-CL+AS, AHC-CL+TU+BA, and AHC-CL+AS+BA.

Figure 4(b) shows the CV curves of AHC-CL+TU, AHC-CL+AS, AHC-CL+TU+BA, and AHC-CL+AS+BA at a scan rate of 10 mV s^{-1} . It can be seen that the CV curves of AHC-CL+TU+BA and AHC-CL+AS+BA have a larger quasi-rectangular area, slightly protruding, and less deformation, indicating that the electric double layer capacitor sample contains a small amount of oxygen, which is the redox reaction of the surface functional groups during the rapid charge and discharge process caused behavior. AHC-CL+TU and AHC-CL+AS have more regular quasirectangular shapes, indicating that electrolyte ions undergo rapid ionic reactions in these two samples.

As shown in Figure 4(c), it is the electrochemical impedance spectroscopy (EIS) of AHC-CL+TU, AHC-CL+AS, AHC-CL+TU+BA, and AHC-CL+AS+BA. The Nyquist plots of the four groups of samples (AHC-CL+TU, AHC-CL+AS, AHC-CL+TU+BA, and AHC-CL+AS+BA) present a small quasi-semicircle in the high-frequency region, followed by a linear transition in the low-frequency region, which is characteristic of carbon materials. And all exhibit good electrical conductivity and ion diffusion properties. A semicircle on the real axis is approximately equal to the charge transfer resistance. And observing the semicircle section connected to the x-axis, it can be seen that with the addition of B, the charge transfer resistance of the electrode doped with AS as the N source increases. However, the charge transfer resistance of electrodes doped with TU as N source decreases. At the same time, the Nyquist curves of AHC-CL+TU, AHC-CL+AS, AHC-CL+TU+BA, and AHC-CL+AS+BA are almost straight lines in the low frequency region, and the slope is almost infinite, indicating that the ion diffusion resistance of the four electrode materials is high.

Figure 5(a, b, c, d) shows the CV curves of AHC-CL+TU, AHC-CL+AS, AHC-CL+TU+BA, and AHC-CL+AS+BA at scan rates of 20 mV s^{-1} , 50 mV s^{-1} , 100 mV s^{-1} and 200 mV s^{-1} , respectively. As shown in Figure 7(a), it can be seen that the CV curves of AHC-CL+TU+BA and AHC-CL+AS+BA have larger quasi-rectangular regions with slight protrusions and less deformation, indicating that the samples of electric double layer Behavior induced by redox reactions of surface functional groups during discharge. AAT and AHC-CL+AS have more regular quasi-rectangular shapes, indicating that electrolyte ions underwent rapid ionic reactions in these two samples. As the scan rate increases, the quasi-rectangular shape of the CV curve starts to change. At a high-speed scan rate of 100 mV s^{-1} , AHC-CL+AS and AHC-CL+TU still maintain a relatively stable quasi-rectangular structure. It shows that AHC has typical double-layer capacitance characteristics and good reversible capacitance characteristics.

The specific capacity data of AHC-CL+TU, AHC-CL+AS, AHC-CL+TU+BA, and AHC-CL+AS+BA were calculated according to the GCD curves, as shown in Figure 7(e). As shown in Figure 7(f), the cycle durability of AHC-CL+TU was investigated by GCD measurement for 20,000 cycles at a charge-discharge current density of 10.0 A g^{-1} . AHC-CL+TU exhibits good cycling stability and high capacitance retention of 99.96%.

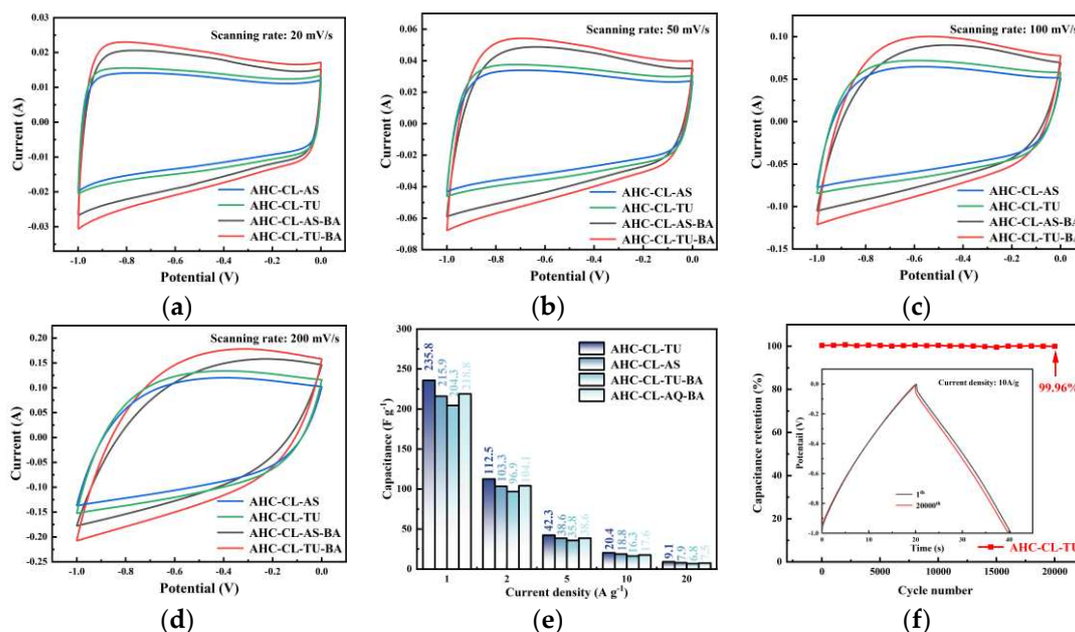


Figure 5. The electrochemical performance data of the three-electrode system using 6M potassium hydroxide solution as the electrolyte. (a). CV curves of AHC-CL+TU, AHC-CL+AS, AHC-CL+TU+BA, and AHC-CL+AS+BA at a scan rate of 20 mV s^{-1} ; (b). CV curves of ABM, ABT, AM, and AT at a scan rate of 50 mV s^{-1} ; (c). CV curves of AHC-CL+TU, AHC-CL+AS, AHC-CL+TU+BA, and AHC-CL+AS+BA at a scan rate of 100 mV s^{-1} ; (d). CV curves of AHC-CL+TU, AHC-CL+AS, AHC-CL+TU+BA, and AHC-CL+AS+BA at a scan rate of 200 mV s^{-1} ; (e). Histogram of specific capacitance at flow densities of 1, 2, 5, 10, 20 A g^{-1} respectively. (f). Based on AHC-CL+TU supercapacitors were tested for stability of 20,000 cycles at a current density of 5 A g^{-1} .

5. Conclusion

The doping of nitrogen and sulfur in the process of hydrothermal carbonization of cellulose could be realized. The doping effect of organic nitrogen and sulfur compounds was better than that of inorganic compounds. The addition of boric acid had a negative effect on the carbon materials prepared by the co-carbonization of both nitrogen sources and cellulose, most notably when thiourea acted as an organic nitrogen source. Boric acid will reduce the specific surface area of porous materials and inhibit the pore-forming process of micropores. The electrochemical performance of the doped thiourea sample was significantly reduced, and the influence of boric acid on the experimental sample could be eliminated by cycling. In the carbonization experiment of doping N and B elements at the same time, boric acid has a more obvious inhibitory effect. Compared with AHC-CL-AS ($829.59 \text{ m}^2 \text{ g}^{-1}$) and AHC-CL-TU ($952.27 \text{ m}^2 \text{ g}^{-1}$), the specific surface area of AHC-CL-AS-BA ($484.76 \text{ m}^2 \text{ g}^{-1}$) and AHC-CL-TU-BA ($397.59 \text{ m}^2 \text{ g}^{-1}$) has obvious decline. At a current density of 1 A g^{-1} , the specific capacitance of AHC-CL-AS-BA (219.3 F g^{-1}) and AHC-CL-AS (216.4 F g^{-1}) has no obvious difference, and AHC-CL-TU-BA (205.8 F g^{-1}) was compared with AHC-CL-TU (236.25 F g^{-1}) decreased significantly. Therefore, boric acid has a certain inhibitory effect on carbonization experiments doped with nitrogen. Nevertheless, we view that supercapacitor carbon materials with good properties can be prepared by hydrothermal carbonization of cellulose and nitrogen and sulfur compounds.

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