Preparation of Biocarbon Derived from Sugarcane Bagasse and Its Composites for Application in Supercapacitors

(サトウキビバガス由来のバイオカーボンとその複合材料の創製およびスーパーキャパシタへの応用)

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Abstract

Supercapacitors, as one of the most promising energy storage devices, due to their good energy density, superior power density, fast charge/discharge rate, and long cycle life, which make it exhibit promising applications in portable devices, renewable energy and transportation. Based on the different charge storage mechanisms, supercapacitors can generally be divided into two types: the electrical double-layer capacitors (EDLC) and the pseudocapacitors. Carbon materials, such as active carbon, graphene and carbon nanotube are usually used in EDLC electrode, but their low energy density, limited their further application, as well as most of methods to prepare of the above carbon materials often requires complicated preparation conditions or special facilities, leading to the high cost. In the present work, first, low cost and uncomplicated method of allium-giganteum-like biocarbon (KWB) were derived from sugarcane bagasse, and then different kinds of carbon-based composite materials including manganese dioxide (MnO$_2$)/KWB, polyaniline (PANI)/KWB and iron oxide (Fe$_3$O$_4$)/KWB were obtained. All of the above materials were used as electrode material of supercapacitors, and morphology, chemical composition and electrochemical performance were investigated. Besides, due to the unique structure of KWB, all the composites show the superior performance in supercapacitors.

In Chapter 1, the research backgrounds, research significance, research purpose and the construction of this thesis are particularly described.

In Chapter 2, the properties of experimental materials used in this thesis are presented. The experimental methods and characterizations are also particularly described in this chapter.

In chapter 3, KWB were derived from sugarcane bagasse via one-step carbonization and activation method. The results show that, both the mass ratio of potassium hydroxide (KOH) to sugarcane bagasse and the temperature of carbonization could impact on the morphology and chemical composition of KWB, thereby further affect
the electrochemical performance of KWB. Under the optimum conditions, the SEM of KWB shows the excellent 3D hierarchical porous network structure with allium-giganteum-like, which also has the best specific surface area. Its promise the excellent electrochemical performance when used as electrode materials for supercapacitors. What’s more, KWB also could be carbon-base to prepare other composites in the further research.

In chapter 4, MnO$_2$/KWB (KWBM) composites have been prepared by a simple method. The MnO$_2$ nanosheets anchored on the surface of biocarbon and obtained the KWBM, and KWB was used as both scaffolds and reducing agents for MnO$_2$ growth. The analysis results demonstrated that chemical composition and microstructure of KWBM were closely related to the mass ratio of KMnO$_4$/KWB, and which further impacted electrochemical performance of KWBM profoundly. Especially, the KWBM-4 exhibited a higher specific capacitance of 402 F g$^{-1}$ at a current density of 1 A g$^{-1}$ in three-electrode system. The asymmetric two-electrode system with outstanding energy density was assembled by employing the KWBM-4 as the positive electrode and the KWB as the negative electrode. The two-electrode system displays a high energy density of 25.9 Wh kg$^{-1}$ at a power density of 750 W kg$^{-1}$ within a potential range of 0-1.5 V. Furthermore, the system exhibited high cycle stability with only 5.8 % loss of its initial capacitance after 2000 cycles.

In chapter 5, a coral-like polyaniline/KWB (PANI/KWB) composites was prepared via in-site polymerization method. The PANI/KWB composites were obtained using KWB as the scaffolds for PANI growth and [CMMIm]Cl ionic liquid as dopant. The different mass ratio of aniline monomer (An) and KWB be researched. The results indicated that chemical composition and microstructure of PANI/KWB were closely related to the mass ratio of aniline monomer (An) and KWB, and which further impacted electrochemical performance of PANI/KWB profoundly. Especially, the PANI/KWB-4 exhibited a highest specific capacitance of 447 F g$^{-1}$ at a current density of 0.5 A g$^{-1}$. Furthermore, the asymmetric two-electrode system based on PANI/KWB-4 as positive electrode and KWB as negative electrode is successfully assembled with a voltage window of 0~1.6 V, exhibiting high energy density (27.3 Wh kg$^{-1}$) and power
density (800 W kg\(^{-1}\)), and excellent cycling stability (87% capacitance retention after 2000 cycles).

In chapter 6, Fe\(_3\)O\(_4\)/KWB nanocomposites (KBFe) have been prepared via a chemical coprecipitation method under different mass ratio of Fe\(_3\)O\(_4\) to KWB. As a result, the chemical composition and microstructure of the KBFe were intimately related to the mass ratio of Fe\(_3\)O\(_4\) and KWB, which impacted the electrochemical performance of the KBFe profoundly. Especially, the KBFe-4 displayed the highest specific capacitance of 342 F g\(^{-1}\) at a current density of 1 A g\(^{-1}\). Furthermore, the asymmetric two-electrode system based on KWBM composite as positive electrode and KBFe-4 as negative electrode was successfully assembled with a voltage window of 0~1.6 V, and exhibited high energy density (29.1 Wh kg\(^{-1}\)) and power density (800 W kg\(^{-1}\)), along with excellent cycling stability (89.5% capacitance retention after 1000 cycles).

In chapter 7, general conclusions of the study are made.
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Chapter 1 Introduction

1.1 Background

Since the beginning of the 21st century, the rapid development of the global economy and the explosive growth of the population have led to an increasing demand for energy. However, the non-renewable fossil fuels, which are the main energy sources, not only have the problem of insufficient stock supply, but also the greenhouse effect brought by the subsequent product carbon dioxide (CO$_2$) is threatening the human living environment, such as greenhouse gas emissions, air pollution and water pollution. For the future, health and sustainable development of our society’s ecosystem, energy have become the most important issue affecting human survival and development. Therefore, all global organizations agree that energy is the main challenge of this century that our habitat must overcome.

Nowadays, developing clean energy sources and their associated technologies are considered a practical way to solve this problem. Therefore, many researchers focus on the renewable energies, for instance, wind [1], seawater [2], geothermal [3], solar [4] [5], biomass [6, 7], and so on. However, most renewable clean energy sources are highly dependent on the time of day and regional weather conditions. For example, the solar cell can produce sufficient and improved efficiency during the daytime, but energy also needs to be stored during daytime for using at night. Development of related energy conversion and energy storage devices is therefore required to effectively harvest these intermittent energy sources. About this matter, electrochemical supercapacitors (ESs) are recognized as one of the most promising electrochemical energy storage/conversion devices, and have been widely applied in consumer electronics, transport, telecommunications, medicine, back-up memory systems and other sectors [8].
1.2 Supercapacitors

Supercapacitors, also are named as ultracapacitors or electrochemical capacitors. Supercapacitors with high power density (> 10 kW kg\(^{-1}\)), super-long cycle life, rapid charge/discharge rate, simple principle and small environment impact hold a great promise in the field of electrical energy storage [10-13]. Generally, due to the different of energy storage principle mechanism, supercapacitors can be divided into two categories: one is based on electrostatic interactions, known as electrical double layers capacitors (EDLCs) and the other involves chemical reactions which named pseudocapacitor [12,14]. In order to distinguish, we explain the two energy storage mechanism will be described separately, though they usually function together in a supercapacitor. Especially in recent years, a new type supercapacitors have been developed, hybrid capacitors, which refer to ones employing both the electrical double-layer (EDL) and pseudocapacitive mechanisms to store charges. With the rapid developments of this area in decade, some new battery-type hybrid devices have been developed, for example lithium ion capacitor and carbon//PbO\(_2\) [8]. These devices are
regarded as hybrid ESs ascribing they have one EDL or pseudocapacitive electrode combined with the other rechargeable battery-type electrode. Based on the difference between the electrode material and the electrode combination, the hybrid ESs reviewed include: (a) those based on composites electrodes made from both EDL capacitive materials and pseudocapacitive materials [15]; (b) those of asymmetric design with one EDL electrode and the other pseudocapacitive or battery-type electrode [16]; as well as, (c) those of the asymmetric structure with one pseudocapacitive electrode and the other rechargeable battery-type electrode [17]. These new hybrid capacitors have been demonstrated to be capable of delivering higher energy density and power density, as well as a broader range of applications [18, 19].

Table 1.1 Summary of the different types of supercapacitor devices [20].

<table>
<thead>
<tr>
<th>Type of supercapacitor</th>
<th>Electrode material</th>
<th>Charge storage mechanism</th>
<th>Merits/shortcomings</th>
</tr>
</thead>
<tbody>
<tr>
<td>Electrochemical double layer capacitor (EDLC)</td>
<td>Carbon</td>
<td>Electrochemical double layer (EDL), non- Faradaic process</td>
<td>Good cycling stability, good rate capability, low specific capacitance, low energy density</td>
</tr>
<tr>
<td>Pseudocapacitor</td>
<td>Redox metal oxide or redox polymer</td>
<td>Redox reaction, Faradaic process</td>
<td>High specific capacitance, relatively high energy density, relatively high power density, relatively low rate capability</td>
</tr>
<tr>
<td>Hybrid capacitor</td>
<td>Asymmetric hybrid</td>
<td>Anode: pseudocapacitance materials, cathode: carbon</td>
<td>High energy density, high power density and good cyclability</td>
</tr>
<tr>
<td>Hybrid capacitor</td>
<td>Symmetric composite hybrid</td>
<td>Redox metal oxide/carbon or redox polymer/carbon</td>
<td>High energy density, moderate cost and moderate stability</td>
</tr>
<tr>
<td>Hybrid capacitor</td>
<td>Battery-like hybrid</td>
<td>Anode: Li-insertion material, cathode: carbon</td>
<td>High energy density, high cost and requires electrode material capacity match</td>
</tr>
</tbody>
</table>

1.2.1 The storage energy principle of EDLCs

The main energy storage mechanism of EDLCs in supercapacitor arises from the reversible electrostatic accumulation of ions on the surface of the porous carbon. EDLCs have the ability to rapid storage and release of energy and storing much energy due to the large interfacial area and the atomic range of charge separation distances [13]. The construct of the double layer has been studied by Helmholtz whose first developed and modelled the double layer concept in investigations on colloidal suspensions in the 19th century [21]. As illustrated in Fig. 1.2a, the Helmholtz model give a description of the charge separation at the electrode/electrolyte interface when an electrode of surface
area is polarized [13, 22]. Because the Helmholtz model does not consider the diffusion of ions in the solution and the interaction between the dipole moment of the solvent and the electrode, for this, Gouy and Chapman put forward a diffuse model of the EDL in which the potential decreases exponentially from the electrode surface to the fluid bulk (Fig. 1.2b) [23, 24]. However, the Gouy-Chapman model is deficient for highly charged double-layers, and in 1924, Stern [25] combined the Helmholtz and Gouy-Chapman models by accounting for the hydrodynamic motion of the ionic species in the diffuse layer and the accumulation of ions close to the electrode surface (Fig. 1.2c). These two layers are equivalent to two capacitors in series, CH (Helmholtz layer) and CD (diffuse layer), and the total capacitance of the electrode (CDL) is given by Equation (1.1):

$$\frac{1}{C_{DL}} = \frac{1}{C_H} + \frac{1}{C_D}$$  \hspace{1cm} (Eq. 1.1)

where CH and CD are the capacitance of compact double layer and diffusion region, respectively.

Fig. 1.2 (a) Helmholtz model, (b) the Gouy-Chapman model, and (c) the Stern model of the electrical double-layer formed at a positively charged electrode in an aqueous
An EDLCs consists of two electrodes, generally, the material of the electrode is active carbon (AC). The material of the electrode is direct contact with the current collector, and the electrode separated by a porous film impregnated with an electrolyte solution (as shown in Fig. 1.3). When an electric potential difference is applied between the electrodes, the negative charge carriers, electrons, in the negatively polarized electrode are balanced by an equal number of positive cations at the electrode/electrolyte interface, while at the positively polarized electrode are electrically balanced by anions [22, 26, 27]. Therefore, the total capacitance can be controlled by the electrode with the smallest value according to the equation Eq. 1.2:

$$\frac{1}{C} = \frac{1}{C_1} + \frac{1}{C_2}$$ (Eq. 1.2)

where $C$, $C^+$ and $C^-$ are the capacitance of the whole supercapacitors, the positive electrode and the negative electrode, respectively. In case of a symmetric system, both electrodes employ by the equal in materials, mass and size, the Eq. 1.3 can be simplified to Eq. 1.3 [18]:

Fig. 1.3 Representation of EDLC in its charged state [28].
\[ C_{sp} = 4 \cdot \frac{C}{m_{AM}} \]  
(Eq. 1.3)

where \( C_{E} \) and \( m_{AM} \) are the capacitance of electrode and the total mass of active materials.

1.2.2 The storage energy principle of pseudocapacitors

Pseudocapacitors, also be named faradaic supercapacitor due to the energy storage formed via faradaic charge-storage phenomenon. Different from the purely electrostatic phenomena of EDLCs, pseudocapacitors have a chemically reactive environment between the electrode material, and the electrolyte, charge transfers are involved between them at the interface, that is, faradaic interactions take place. These charge transfers provoke chemical changes (oxidation-state change) in the electrode materials, and thus, charges are stored chemically at some oxidation states.

During discharge, this chemical reaction takes place in reverse, and the electrode materials are restored to the previous oxidation states and the stored charges are released [14, 20, 29] (as shown in Fig. 1.4). The chemical reactions are not completely reversible ever for the best effort reversible combination of electrode and electrolyte. Hence, even after complete discharge, pseudocapacitors are always had some residual electrode materials cannot take part in the reversible chemical reaction process [14]. This phenomenon issue in the loss of active materials, which further reduces the value of the maximum specific capacitance with increasing the cycle numbers, thus the cycle stability of the pseudocapacitors also reduces.

Furthermore, pseudocapacitors have the highest charge-transfer rate at a special potential. But sometime pseudocapacitors have instantaneous electrostatic responses on account of the delayed response of pseudocapacitors relative to EDLCs, which need a delay to overcome this potential barrier [14, 29]. Although, pseudocapacitors have the dispute of cycle stability and response-time, they are still aroused researcher's interesting owing to their high ability of storage energy based on the chemical reaction. Hence, overcome the above mentioned problems have been the topic of scientific investigation in recent years.
1.3 The electrode materials of supercapacitors

A typical supercapacitor contains four components: current collector, separators, electrolyte, and electrode. Electrode material is one of the core components, which directly determines the capacitance, delivery rate and efficiency of a supercapacitor.

1.3.1 Carbon-based electrode materials

The 21st century has predominantly become the carbon age, as carbon (graphite, hard carbon, glassy carbon, carbon black, as shown in Fig. 1.5) have previously been utilized in almost all new energy devices, including supercapacitors, Li-ion batteries, ultracapacitors, and hydrogen-storage devices [9]. Due to the sp²-hybridized solid carbon and the enormous adaptability to different interface processes of carbon element, carbon-based materials are the most prospective candidates for the supercapacitor negative electrode material arises from the unique chemical and physical properties, such as low cost, abundance, nontoxicity, and environmentally friendly nature, processability and compatibility in composite materials, as well as high electronic conductivity and outstanding temperature stability [27, 30]. Carbon materials generally
work as EDLCs, wherefore the high surface area range (1 to > 2000 m$^2$g$^{-1}$) and conductivity are critical factors for them as the supercapacitors electrodes. Furthermore, the surface area of carbon materials will be deeply impacted by the porous structures. Thus, a tunable porous structure and carbon surface chemistry, as well as large surface area and high electrical conductivity are desired attributes in tailoring electrode properties to achieve optimum performance.

Fig. 1.5 Schematic illustrations of several important carbon nanostructures, (a) fullerene, (b) graphene, (c) carbon nanotube, and (d) carbon nanofiber [30].

1) Activated carbon (AC)

Among the various carbon-based materials, activated carbon (AC) has been the first choice from research and commercial perspectives over the last 40 years due to their merits of low cost, large theoretical surface area (≈3000 m$^2$ g$^{-1}$) and a broad pore-size tenability [9, 13, 31]. In general, AC is produced from physical (thermal) and/or chemical activation of various types of carbonaceous source materials, such as polymer (dopamine [32], polypyrrole [33], protein [34]), chemical reagent (zeolitic imidazolate [35]), biomass (sugarcane bagasse [36], pine cones [37], bamboo [38], waste coffee [31]), and etc. Depending on the activation methods as well as the carbon precursors used, AC possessing various physicochemical properties with well-developed surface areas as high as 3000 m$^2$g$^{-1}$ have been produced, good electrochemical properties have
been studied, as well as the porous structure could be designed [39-41]. The International Union of Pure and Applied Chemistry (IUPAC) defines porous according to their size as follows [42]:

- **Macrospores**: larger than 50 nm
- **Mesopores**: between 2 and 50 nm
- **Micropores**: smaller than 2 nm

On account of the capacitive mechanism for AC is mainly physical adsorption/desorption. Therefore, the very important factor is the porous structure (as shown in Fig. 1.6). In other words, the capacity of AC depends on several mutually interlinked parameters, that is, total specific surface area, normally referred to as the Brunauer–Emmet–Teller (BET) surface area; the nature of the porosity of the electrode; the pore size distribution; and the conductivity of the electrode material [14, 43]. The porous structure of AC can be controlled through different experimental conditions, such as the carbonization temperature in physical activation, and the dosage of activating agents in chemical activation, while to obtain the perfect porous structure for electrode [27]. Someone successful prepared the AC with the specific surface area reach up to 3000 m²g⁻¹, but the corresponding specific capacitance < 10 μF cm⁻², much smaller than the theoretical value (15-25 μF cm⁻²), that means not all pores are benefited for charge accumulation [13]. Actually, a large portion of the micropores remains inaccessible to the electrolyte because of the incompatible pore size with electrolyte ions, which significantly drops the usable surface area and the specific capacitance [44].

In recent year, some researches show that a maximum EDLC could be observed in AC with pores of the size similar to that of electrolyte ions, while pores whose size is largely deviated from ion size can result in significant loss of the capacitance [45, 46]. Hence, it is imperative to control the pore-size distribution in ACs to maximize the energy/power density. Hierarchically porous AC with 3D network framework has been developed for high rate electrochemical capacitive energy storage [47-49]. This kind of 3D AC is characterized by porous of all different dimensions, including micropores, mesopores and macropores. What’s more, through research people have found that pores with different pore sizes have different effect. Micropores contribute mostly to
the specific surface area and accessible to hydrated ions [50, 51]; mesopores can allow the rapid transport of ions from the electrolyte to the entire surface of the electrodes and make them quickly available for electric charge exchange [30]; macropores and localized graphitic structure can serve as ion-buffering reservoirs and enhances the electrical conductivity, respectively [30]. Especially, the macro and mesopores are a prerequisite for the transport of the electrolyte to the micropores in many ACs (as shown in Fig. 1.7) [30, 52, 53].

![Fig. 1.6 Representation of charged state of a symmetric EDLCs using porous material and of the corresponding equivalent circuit [27].](image)

![Fig. 1.7 Schema for adsorption of electrolyte ions with or without solvation to the surface of pores with different sizes [53].](image)
However, it remains limited to obtain the maximum capacitance of EDLCs through control the surface area and pore size of the AC, as well as high energy density supercapacitors, require more excellent electrochemical performance of AC electrode materials. An effective means of improving electrochemical performance of AC material has been proved, that is introduced pseudocapacitive phenomena at the electrode and electrolyte interface via substituting some atoms in AC by heteroatoms (such as N [49], O [54], P [55], B [56]). Nevertheless, we note that the electronic properties of AC depend strongly on size and chirality, so the properties cannot be easily tailored [57, 58]. B or N usually be choice to doping in AC, due to B and N are the nearest neighbors of the C in the periodic table and atomic radius of B and N are similar to C [59, 60]. Also, Co-doped with two heteroatoms are also used, and this is a benefit for their application in electronic devices [56, 57, 61].

2) Carbon nanotubes (CNTs)

Despite the great use of AC materials as supercapacitors electrodes, but the inaccessibility of the electrolyte ions into their micropores and/or interior atoms at higher scan rates, as well as the poor electrical conductivity leads to a higher internal resistance still remains the central issue that limits their being used in high-power-density supercapacitors [9]. Carbon nanotubes (CNTs), its discovery has significantly advanced the science and engineering of carbon materials, and have been considered suitable as a supercapacitor electrode material [62]. In the past decade, CNTs have been widely researched for supercapacitors attribute to the high conductivity, unique mechanical properties, high chemical stability, high aspect ratios and high activated surface areas [63, 64]. As technology advances, various forms of CNTs have been generating in assorted conformations and crystalline order by control manipulation of various parameters in preparation process [65]. At the molecular level, CNTs can be viewed as a graphene sheet rolled up into a nanoscale tube form to produce a single-walled carbon nanotube (SWCNT) (as shown in Fig. 1.8). There may be additional graphene coaxial tubes around the SWCNT core to form a multiwalled carbon nanotube
(MWCNT) [66, 67].

However, CNTs even behave moderately small theoretical surface areas (≈50-1315 m² g⁻¹), but they performed higher capacitances over other AC. Because of their unique tubular structures and the high density of mesopores allow fast charge transport and large accessibility of electrolyte ions [9, 68]. As a supercapacitors electrode, the initial work on CNTs was mainly carried out with randomly oriented of MWCNTs, preforming a specific capacitor in the 80-130 F g⁻¹ [62, 65]. Compared to MWCNTs, SWCNTs exhibited better electrochemical performances attributed to their large specific surface area (≈1600 m² g⁻¹), high aspect ratio, and better accessibility to the electrolyte ions [67, 69]. Aligned CNTs are present unbundled structures, providing more mesopores and accessible surfaces. Moreover, they provide lower contact resistance during the charging–discharging process, bring about large energy and power densities [9, 70].

![Schematic representation of the CNTs formation](image)

**Fig. 1.8** Schematic representation of the CNTs formation by rolling up a 2D graphene sheet of lattice vectors a1 and a2, the roll-up chiral vector \( C_h = na_1 + ma_2 \), and the chiral angle \( \theta \) between \( C_h \) and \( a_1 \). When the graphene sheet is rolled up to form the cylindrical part of the nanotube, the chiral vector forms the circumference of nanotube’s circular cross section with its ends meeting each other. The chiral vector \( (n, m) \) defines the tube helicity. If \( n - m \neq 3q \) (n, m, and q are integers), the SWCNT is semiconducting; if \( n - m = 3q \), the SWCNT is metallic. Reproduced with permission [67].


3) Graphene

Graphene, the one-atom-thick 2D sheet of sp2-bonded carbon that is densely packed in a honeycomb crystal lattice, is a rising star in carbon material and nanoscience [71]. The unique structure of graphene is endowed its various superior properties such as high electrical and thermal conductivities, good transparency, great mechanical strength, inherent flexibility and huge specific surface area [72, 73]. Thence, during the past decade graphene has attracted a great deal of attention in the fields of microelectronic and optoelectronic devices, energy storage materials, electrocatalysts, polymer composites, and ultrastrong paper-like materials [74-76]. The first piece of graphene was obtained by Andre Geim and Konstantin Novoselov via manual mechanical cleavage of graphite with Scotch tape in 2004 [67, 71]. However, this production method is not applicable on a large scale. Currently, the most comment method is reduced graphite oxide method, which exfoliation GO followed by reduction to give the final product (as shown in Fig. 1.9) [14, 71, 77, 78]. Moreover, the other methods are assembling graphene sheets into graphene-based materials with ordered 2D or 3D microstructures and separating graphene sheets with other nanomaterials [73, 79].

![Fig. 1.9 Preparation of reduced graphene oxide by Hummer method [71].](image)

Unlike the current electrochemical capacitors fabricated with AC and CNT, the high specific surface area of graphene does not depend on the distribution of pores. Single or few layered structure of graphene irreversibly aggregate due to the π-π interactions and van der Waals forces, which have a lower degree of agglomeration [80]. What’s
more, the excellent electric conductivity and mechanical flexibility of graphene enable its electrodes to undergo fast electron transfer without any significant volume expansion/shrinking during charging-discharging [81]. Hence, many of reports directly utilize the 2D structure of graphene for supercapacitors. A. Yu et al. prepared an ultrathin film (e.g., 25 nm) with a moderate specific capacitor at 111 F g\(^{-1}\), and this value was further decreased by increasing its thickness [82]. In addition, Q. Du reported a method through thermal reduction of graphene oxide powder at a high temperature (1050 °C) rapidly generated a large number of gaseous species and exfoliated r-GO sheets into a porous structure with the high specific surface area, while the porous functionalized graphene with the specific capacitor of 233 F g\(^{-1}\) [83].

1.3.2 Transition metal oxides-based electrode materials

Although carbon-based materials hold great potential as negative electrodes, their intrinsically low specific capacitor and low energy density are major drawbacks. Transition metal oxides, exhibit higher energy density for supercapacitors than that of carbon materials due to the intrinsically fast and reversible redox reactions [73]. As we all know, the pseudocapacitance of transition metal oxides arises from both faradic redox reactions and the electrochemical adsorption/desorption of ions at the electrode/electrolyte interface [84]. The normal conditions for transition metal oxides in supercapacitors application are [85]:

- the metal can exist in two or more oxidation states that coexist over a continuous range with no phase changes involving irreversible modifications of a 3D structure;
- the protons can freely intercalate into the oxide lattice on reduction and out of the lattice on oxidation, allowing facile interconversion of O\(^2^-\) ↔ OH\(^-\);
- the oxide should be electronically conductive.

In current, the common transition metal oxides are ruthenium oxide, manganese oxide, cobalt oxide, iron oxide and nickel oxide.
1) Ruthenium dioxide (RuO₂)

RuO₂ is a promising electrode material, and first studied as charge storage material by Conway et al. in 1975, then the concept of a supercapacitor was introduced [86]. RuO₂ has been widely investigated due to its highly reversible redox reactions, three distinct oxidation states (Ru²⁺, Ru³⁺, Ru⁴⁺), high proton conductivity, remarkably high specific capacitance and high rate capability. The pseudocapacitive behaviours of RuO₂ in acid and alkali electrolyte exhibit different reactions. In acid electrolyte, the mechanism was suggested to be as follow (Eq. 1.4 and 1.5):

\[
\text{RuO}_2 + x\text{H}^+ + xe^- \leftrightarrow \text{RuO}_{2-x}(\text{OH})_x \quad 0 \leq x \leq 2 \quad \text{(Eq. 1.4)}
\]

Or

\[
\text{RuO}_{4}(\text{OH})_y + \delta\text{H}^+ + \delta e^- \leftrightarrow \text{RuO}_{x-\delta}(\text{OH})_{y+\delta} \quad \text{(Eq. 1.5)}
\]

Four steps were proposed to determine the capacitive performance of RuOₓ·nH₂O:
(i) electron hopping within RuOₓ·nH₂O particles; (ii) electron hopping between particles; (iii) electron hopping between electrode materials and current collectors; and (iv) the proton diffusion within RuOₓ·nH₂O particles [87]. It is worth noting, the crystallinity of RuO₂ is the key parameter in determining the value of specific capacitance. Although crystallization of RuO₂ can reduce the intraparticle electron hopping resistance of RuOₓ·nH₂O particles, while it results in an increase in the diffusion barrier of proton within crystalline RuO₂ and a loss of active sites [88]. Accordingly, nanocrystalline RuOₓ·nH₂O annealed at temperatures close to its crystalline temperatures usually exhibits the highest specific capacitor value [89]. Although amorphous RuO₂ can provide excellent capacitive performance, its drawbacks, such as the high cost and environment impact, hinder it from being used in the commercialization of supercapacitors [85].

2) Manganese dioxide (MnO₂)

Manganese oxides have been regarded a potential electrode material for pseudocapacitor applications owing to its low cost, low toxicity, environment safety and high theoretical capacities, and also have acted as the promising alternative class of materials for RuO₂ [85]. Lee and Goodenough first reported the amorphous MnO₂
could be used as an excellent electrode material for the faradaic electrochemical capacitor in 1999 [90]. After that, aiming the development and application of manganese oxides have attracted major attention. Manganese oxides also have pluralism valence states, including Mn(0), Mn(II), Mn(III), Mn(IV), Mn(V), Mn(VI), and Mn(VII). The capacitance of manganese oxides originates mainly from pseudocapacitive energy storage mechanism, which is attributed to reversible redox transitions involving the exchange of protons and/or cations on the surface and in the bulk of the electrode with the electrolyte, as well as the transitions between Mn(III)/Mn(II), Mn(IV)/Mn(III), and Mn(VI)/Mn(IV) [85, 91]. The surface Faradaic reaction involves the surface adsorption of electrolyte cations (C\(^+\) = H\(^+\), Li\(^+\), Na\(^+\) and K\(^+\)) on the manganese oxide [92, 93]:

\[
(MnO_2)_{\text{surface}} + C^+ + e^+ \leftrightarrow (\text{MnOOC})_{\text{surface}} \quad \text{(Eq. 1.6)}
\]

The bulk Faradaic reaction relies on the intercalation or deintercalation of electrolyte cations in the bulk of the manganese oxide [92, 93]:

\[
\text{MnO}_2 + C^+ + e \leftrightarrow \text{MnOOC} \quad \text{(Eq. 1.7)}
\]

It is noted that, in both charge storage mechanisms, a redox reaction between the III and IV oxidation states of Mn ions occurs.
Different from RuO$_2$, the pseudocapacitive behavior of manganese oxides is affected by both physical and chemical factors. Cycling stability is mostly controlled by the microstructure, however, the specific capacitance is controlled mostly by the chemical hydrated state [91]. Similar to RuO$_2$, manganese oxides also exist a variety of crystalline structures, including α-, β-, γ-, δ- and λ-MnO$_2$. Among them, α-, β-, and δ-MnO$_2$ have a tunnel structure (2 × 2 octahedral units for α-MnO$_2$, the relatively large 1D tunnel structured phase; 1 × 1 octahedral units for β-MnO$_2$, the more compact and dense 1D tunnel structured phase), and δ- MnO$_2$ has a relatively open 2D layered structure (as shown in Fig. 11 by birnessite) [85, 94].

Among of them, 2D layered structure of δ- MnO$_2$ exhibited outstanding capacitive performance, because the interlayer separation of δ- MnO$_2$ is about 7 Å, this size can allow mount of water and stabilizing cations such as Na$^+$ or K$^+$ between the sheets [95]. Due to this reason, several methods for synthesizing δ- MnO$_2$ have been developed, such as hydrothermal synthesis, redox reaction synthesis, sol–gel method, template method and so on [28] [96, 97]. However, the partial dissolution of MnO$_2$ causes the
capacitance degradation during cycling, as well as the low specific surface area and poor electronic conductivity also limit the application of MnO₂ in supercapacitors [85, 98].

3) **Iron oxide (Fe₃O₄)**

As one of transition-metal oxides, magnetite (Fe₃O₄) have two valence states (Fe²⁺, Fe³⁺) which make a promise to be applied as supercapacitor material. Furthermore, this material has advantages of low cost and environmentally affinity [99]. However, the intrinsic problem of Fe₃O₄ supercapacitor electrode is lack of electrical conductivity (the specific capacitance of pure Fe₃O₄ is 60-80 F g⁻¹) [100]. In particular, the capacitance of the Fe₃O₄ was found to be sensitive to the anion species but not to either alkaline cations or electrolyte pH<11. These behaviors suggest a capacitance mechanism is different from that of either RuO₂ or MnO₂ [87].

In addition, some experimental results indicate that, in Na₂SO₃, the capacitive current of magnetite electrode originates from the combination of electric double layer capacitance (EDLC) and the pseudocapacitance that involves successive reduction of the specifically adsorbed sulfite anions, from SO₃²⁻ through S²⁻, and vice versa. In Na₂SO₄, the current is due entirely to EDLC. The lowest capacitance of magnetite observed in KOH is attributed to the formation of an insulating layer on the magnetite surface. To solve this problem, the common countermeasure is the preparation of hybrid nanostructured material by integrating Fe₃O₄ with a carbon host, where the carbon host acts as the conductive network [101, 102].

1.3.3 **Conducting polymer-based electrode materials**

Since 1977, American scientists Heeger, MacDiarmid and their Japanese colleague Shirakawa discovered that a synthetic polymer (polyacetylene) could be conductive under certain conditions, conducting polymer materials have attracted more and more
attention by scholars [103, 104]. For the revolutionized discovery and development of conducting polymers, the Nobel Prize in Chemistry for 2000 was rewarded to the above scientists [105-107]. Conducting polymers, alternate single-double bonds along the backbone of the polymer is a key property for conjugated conducting polymers, which differentiates them from other polymers. The “sigma” (σ) electrons that form strong covalent bonds between the carbon atoms are localized, whereas the “pi” (π) electrons can be delocalized upon certain operation [108]. For this reason, they can render conductive through the conjugated bond system along the polymer backbone [109]. The molecular structures of some typical conjugated conducting polymers in their neutral form are given in Fig. 1.11.

Most of the conducting polymers show poorly conductive and unstable in the neutral state. However, their conductivity can improve by the formation of charge carriers upon oxidizing (p-doping) or reducing (n-doping) their conjugated backbone [110]. The
simplified reaction (Eq. 1.8 and 1.9) for these tow charging processes are as follows [109]:

\[
\begin{align*}
\text{Cp} & \rightarrow \text{Cp}^{n+}(A^-)n + n\text{e}^- \quad \text{(p-doping)} \quad \text{(Eq. 1.8)} \\
\text{Cp} + n\text{e}^- & \rightarrow (C^+)n\text{Cp}^{n-} \quad \text{(n-doping)} \quad \text{(Eq. 1.9)}
\end{align*}
\]

The typical n-doping conducting polymer are PA and poly(paraphenylene), and these polymers have high impedances. Polypyrrole (PPy) and polyaniline (PANI), they are the typical p-doping conducting polymers [111]. Depending upon the oxidation state, PANI has three basic structures (Fig. 1.12): (a) a fully reduced leucoemeral- dine base (LEB); (b) a fully oxidized pernigraniline base (PNB); and (c) a half oxidized/half reduce demeraldine base (EB) state. The protonated or doped EB form is highly conducting, while LEB and PNB are mainly insulating in nature. When EB (Fig. 1.12a) is doped with different acids a polaron is formed through the successive formation of bi- positive species (Fig. 1.12b), bipolaron structure (Fig. 1.12c) and more stable polaron structure (Fig. 1.12d). This polaron structure is responsible for electrical conduction through the hopping mechanism in the crystalline region and hopping may be intra-chain or inter-chain [112]. This process leads to an internal redox reaction converting the semiconducting form of PANI (emeraldine base) to a metallic form (emeraldine salt). The reaction between the different forms of PANI is shown in the Fig. 1.13 [113].
Fig. 1.1 Sketch of EB of polyaniline before protonation (a), after 50% protonation (b) and formation of bipolaron (c) and polaron (d) [112]

Fig. 1.12 Sketch of EB of polyaniline before protonation (a), after 50% protonation (b) and formation of bipolaron (c) and polaron (d) [112]

Conducting polymers are suitable materials for supercapacitors, due to the advantage of high conductivity in a doped state, high storage capacity/reversibility, adjustable redox activity through chemical modification, and low cost/environment impact [114, 115].
The common conducting polymer in supercapacitor applications is PANI [116], PPy [117], polythiophene [118] and their derivatives. PANI and PPy often use as cathode materials due to they can only be p-doped.

There are many reports on applying the above conducting polymer to supercapacitors. For example, Y. Shi et al. have reported a polypyrrole hydrogels exhibited a high specific capacitance and which reach up to 380 F g\(^{-1}\) [119]. A kind of 3D polyaniline has prepared by Y. Gawli et al. by inorganic and organic acid co-doping, and it possesses a nearly constant high specific capacitance of 350 F g\(^{-1}\) at the current density range of 1~ 40 A g\(^{-1}\) [106]. Even, Y. Huang et al. have prepared a kind of polypyrrole from the molecular ordering, and this kind of polypyrrole exhibited the excellent cycle stability (the capacitance retentions of over 97, 91 and 86 % after 15000, 50000 and 100000 charging/discharging cycles, respectively), and can sustain over 230000 charging/discharging cycles with still about half of initial capacitance retained [120].

Unfortunately, swelling and shrinking of conducting polymer unavoidable occur during the charging/discharging process (due to the intercalating/deintercalating function). This problem will further lead to mechanical degradation of the electrode and fading electrochemical performance during cycling, and then limit the conducting polymer applied to supercapacitors.

### 1.4 Contribution from this dissertation

Supercapacitors, as one of the most promising energy storage devices, due to their good energy density, superior power density, fast charge/discharge rate, and long cycle life, which make it exhibit promising applications in portable devices, renewable energy and transportation. Based on the different charge storage mechanisms, supercapacitors can generally be divided into two types: the electrical double-layer capacitors (EDLC) and the pseudocapacitors. Carbon materials, such as active carbon, graphene and carbon nanotube are usually used in EDLC electrode, but their low energy density, limited their further application, as well as most of methods to prepare of the above carbon materials often requires complicated preparation conditions or special facilities, leading to the
high cost. In the present work, first, low cost and uncomplicated method of allium-giganteum-like biocarbon (KWB) were derived from sugarcane bagasse, and then different kinds of carbon-based composite materials including manganese dioxide (MnO₂)/KWB, polyaniline (PANI)/KWB and iron oxide (Fe₃O₄)/KWB were obtained. All of the above materials were used as electrode material of supercapacitors, and morphology, chemical composition and electrochemical performance were investigated. Besides, due to the unique structure of KWB, all the composites show the superior performance in supercapacitors.

In Chapter 1, the research backgrounds, research significance, research purpose and the construction of this thesis are particularly described.

In Chapter 2, the properties of experimental materials used in this thesis are presented. The experimental methods and characterizations are also particularly described in this chapter.

In chapter 3, KWB were derived from sugarcane bagasse via one-step carbonization and activation method. The results show that, both the mass ratio of potassium hydroxide (KOH) to sugarcane bagasse and the temperature of carbonization could impact on the morphology and chemical composition of KWB, thereby further affect the electrochemical performance of KWB. Under the optimum conditions, the SEM of KWB shows the excellent 3D hierarchical porous network structure with allium-giganteum-like, which also has the best specific surface area. Its promise the excellent electrochemical performance when used as electrode materials for supercapacitors. What’s more, KWB also could be carbon-base to prepare other composites for the further research.

In chapter 4, MnO₂/KWB (KWBM) composites have been prepared by a simple method. The MnO₂ nanosheets anchored on the surface of biocarbon and obtained the KWBM, and KWB was used as both scaffolds and reducing agents for MnO₂ growth. The analysis results demonstrated that chemical composition and microstructure of KWBM were closely related to the mass ratio of KMnO₄/KWB, and which further impacted electrochemical performance of KWBM profoundly. Especially, the KWBM-4 exhibited a higher specific capacitance of 402 F g⁻¹ at a current density of 1 A g⁻¹ in
three-electrode system. The asymmetric two-electrode system with outstanding energy density was assembled by employing the KWBM-4 as the positive electrode and the KWB as the negative electrode. The two-electrode system displays a high energy density of 25.9 Wh kg\(^{-1}\) at a power density of 750 W kg\(^{-1}\) within a potential range of 0-1.5 V. Furthermore, the system exhibited high cycle stability with only 5.8 % loss of its initial capacitance after 2000 cycles.

In chapter 5, a coral-like polyaniline/KWB (PANI/KWB) composites were prepared via in-site polymerization method. The PANI/KWB composites were obtained using KWB as the scaffolds for PANI growth and [CMMIm]Cl ionic liquid as the dopant. The different mass ratio of aniline monomer (An) and KWB be researched. The results indicated that chemical composition and microstructure of PANI/KWB were closely related to the mass ratio of aniline monomer (An) and KWB, and which further impacted electrochemical performance of PANI/KWB profoundly. Especially, the PANI/KWB-4 exhibited a highest specific capacitance of 447 F g\(^{-1}\) at a current density of 0.5 A g\(^{-1}\). Furthermore, the asymmetric two-electrode system based on PANI/KWB-4 as positive electrode and KWB as negative electrode is successfully assembled with a voltage window of 0~1.6 V, exhibiting high energy density (27.3 Wh kg\(^{-1}\)) and power density (800 W kg\(^{-1}\)), and excellent cycling stability (87% capacitance retention after 2000 cycles).

In chapter 6, Fe\(_3\)O\(_4\)/KWB nanocomposites (KBFe) have been prepared via a chemical coprecipitation method under different mass ratio of Fe\(_3\)O\(_4\) to KWB. As a result, the chemical composition and microstructure of the KBFe were intimately related to the mass ratio of Fe\(_3\)O\(_4\) and KWB, which impacted the electrochemical performance of the KBFe profoundly. Especially, the KBFe-4 displayed the highest specific capacitance of 342 F g\(^{-1}\) at a current density of 1 A g\(^{-1}\). Furthermore, the asymmetric two-electrode system based on KWBM composite as positive electrode and KBFe-4 as negative electrode was successfully assembled with a voltage window of 0~1.6 V, and exhibited high energy density (29.1 Wh kg\(^{-1}\)) and power density (800 W kg\(^{-1}\)), along with excellent cycling stability (89.5% capacitance retention after 1000 cycles).
In chapter 7, general conclusions of the study are made.
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Chapter 2 Materials, experimental and characterizations

2.1 Materials

2.1.1 Chemical reagents

1) Sugarcane bagasse

Sugarcane bagasse is the remaining biomass waste after the sugarcane has been squeezed. Due to the amount of fiber inside is about 50%, sugarcane bagasse mainly used for paper and other industries [1]. However, in this thesis, sugarcane bagasse as a biomass waste is chosen to use as a carbon source for preparing the biomass activated carbon. Before used as the material, the sugarcane bagasse was cut into small pieces (opening size approximately 2 mm) by family-use grinder and washed with distilled water for 8 h at 90 oC, and then dried at 80 oC for 24 h. The sugarcane bagasse was bought from Nanning (Guangxi, China).

![Digital photograph of sugarcane bagasse (a), after crushing treatment (b) and SEM image of sugarcane bagasse after crushing treatment (c).]

Fig. 2.1 Digital photograph of sugarcane bagasse (a), after crushing treatment (b) and SEM image of sugarcane bagasse after crushing treatment (c).

2) Aniline

Aniline is a six-membered heterocyclic aromatic organic compound with the formula...
C₆H₇N. It is a colorless liquid and the color will become darker when it is exposed to air. Usually, it is purified by distillation immediately prior to use. In this thesis, aniline was used as a monomer to prepare polyaniline by the oxidation of aniline. Aniline (analytical grade reagent) was purchased from Nacalai Tesque, Inc. (Kyoto, Japan) and freshly distilled under pressure before use.

Fig. 2.2 Structure of aniline.

3) 1-methyl-3-alkylcarboxylic acid imidazolium chloride ([CMMIm]Cl) ionic liquid

Fig. 2.3 Structure of [CMMIm]Cl IL

Ionic liquids (ILs) also named room temperature molten salt, are a special class of salts that have melting points at or close to ambient temperature, which molecular structures were synthesized to compose bounded anions and usually organic cations with long chain [2]. Due to the advantages of good thermal stability, low flammability, high electrochemical stability, high ionic and thermal conductivity, ILs have been extensive applied as solvents, separation media, electrolytes, and lubricants in many fields.

In this thesis, we used [CMMIm]Cl IL to prepared PANI/KWB composites, and the [CMMIm]Cl IL was synthesis in lab. [CMMIm]Cl is a brønsted acidic ILs, which has acidic groups to provide a proton acid environmental for doped state PANI. The cation
of [CMMIm]Cl IL is ([CMMIm])+, has imidazole ring) will enter the main chain by charge balance, which is benefit for the regularity of the PANI chain, and further has a beneficial effect on the electrochemical performance of PANI. In addition, the [CMMIm]Cl IL is also a surfactant, and its addition can promote the dispersion of An monomer on the surface of KWB.

4) Other materials

<table>
<thead>
<tr>
<th>Name</th>
<th>Molecular formula</th>
<th>Structure formula</th>
<th>Molecular weight</th>
<th>Density (g cm-3)</th>
<th>Purity</th>
</tr>
</thead>
<tbody>
<tr>
<td>Potassium hydroxide</td>
<td>KOH</td>
<td>--</td>
<td>56.10</td>
<td>2.044</td>
<td>85%</td>
</tr>
<tr>
<td>Hydrochloric Acid</td>
<td>HCl</td>
<td>--</td>
<td>36.47</td>
<td>1.18</td>
<td>35-37%</td>
</tr>
<tr>
<td>Potassium permanganate</td>
<td>KMnO₄</td>
<td>--</td>
<td>158.03</td>
<td>1.01</td>
<td>99%</td>
</tr>
<tr>
<td>ferrous chloride</td>
<td>FeCl₂·4H₂O</td>
<td>--</td>
<td>198.75</td>
<td>3.16</td>
<td>99%</td>
</tr>
<tr>
<td>ferric chloride</td>
<td>FeCl₃·6H₂O</td>
<td>--</td>
<td>270.20</td>
<td>2.90</td>
<td>99%</td>
</tr>
<tr>
<td>Ethanol</td>
<td>C₂H₆O</td>
<td>CH</td>
<td>46.07</td>
<td>789</td>
<td>99.5%</td>
</tr>
<tr>
<td>Acetonitrile</td>
<td>C₂H₃N</td>
<td>--</td>
<td>41.05</td>
<td>0.79</td>
<td>99.5%</td>
</tr>
<tr>
<td>Chloroacetic acid</td>
<td>C₂H₃ClO₂</td>
<td></td>
<td>94.49</td>
<td>1.58</td>
<td>97%</td>
</tr>
<tr>
<td>1-Methylimidazole</td>
<td>C₄H₈N₂</td>
<td></td>
<td>82.10</td>
<td>1.03</td>
<td>99%</td>
</tr>
<tr>
<td>Ammonium persulphate (APS)</td>
<td>(NH₄)₂S₂O₈</td>
<td></td>
<td>228.20</td>
<td>1.98</td>
<td>98%</td>
</tr>
</tbody>
</table>

All the reagents were purchased from Nacalai Tesque, Inc. (Kyoto, Japan) and used without further purification. And all solutions were prepared with distilled water.
2.1.2 Materials of electrochemical test

For electrochemical testing, stainless mesh (300 mesh) and nickel foam are chosen as the current collector. Before use, stainless mesh or nickel foam was trimmed to rectangular strips with 3×1 cm, after that the strips were immersed in ethanol with sonication for 1 h. Finally, the strips were dried under vacuum at 60 °C for 12 h. The polytetrafluoroethylene (PTFE) were used as the binder between active materials and current collector, and the acetylene black were chosen as the conductive agent for preparing the testing samples [3, 4].

2.2 Experimental methods

2.2.1 Preparation of KOH-activated biomass carbon (KWB) derived from sugarcane bagasse

KWB was obtained by KOH activation method and the typical process as follows: first, sugarcane bagasse was cut into small pieces (opening size approximately 2 mm) and washed with distilled water for 8 h at 90 °C, and then dried at 80 °C for 24 h. Second, 1 g cleanly sugarcane bagasse and 0.6 KOH were poured into 12 ml absolute ethanol with stirring and heated at 60 °C until all of the ethanol was evaporated, and then dried at 70 °C for 12 h. Third, the remnant mixture was pyrolyzed in a tubular furnace in N₂ atmosphere at 800 °C (heating rate is 10 °C min⁻¹) for 2 h. Finally, the obtained carbon material was washed by 1 M HCl solution and deionized water till the filtrate became neutral and collected after vacuum dry.

Fig. 2.4 The process of high temperature carbonization.
2.2.2 Preparation of MnO$_2$/KWB composites (KWBM)

The KWBM composites were prepared by directly reacting KMnO$_4$ with KWB. In a typical process, 0.05 g KMnO$_4$ dissolved in 100 ml deionized water, and then 0.1 g KWB (the mass ratio of KMnO$_4$/KWB is 0.5:1) added the above solution. Subsequently, the mixture was stirred under water bath at 70 ºC until the color of solution changed from modena to tawny. The products removed from solution and washed with deionized water for several times, and dried in vacuum at 80 ºC for 12 h to obtain the KWBM-1 composite. The other products were denoted as KWBM-2, KWBM-3 and KWBM-4 for the mass ratio of KMnO$_4$/KWB is 1:1, 1.5:1 and 2:1, respectively.

2.2.3 Synthesis of 1-methyl-3-alkylcarboxylic acid imidazolium chloride ([CMMIm]Cl) ionic liquid

The [CMMIm]Cl was prepared as Fei reported [5]. In a typical fabrication of [CMMIm]Cl, 0.17 mol chloroacetic acid solid was added spoonful to 0.15 mol N-methylimidazole liquid with stirring in 1 h at 15-20 ºC. When the mixed solution uniform, put it into the water bath kettle to heat up to 70 ºC and run the reaction for 3-5 h. After that, the solution was used for suction filtration and washed the material with 30 ml of methyl cyanide for twice. Finally, the product was dried in vacuum at 40 ºC for 12 h to obtain the white solid of [CMMIm]Cl.

2.2.4 Synthesis of pure PANI without IL

For composite with the PANI via [CMMIm]Cl brønsted acidic IL as dopant, the pure were prepared by the same method while using sulfamic acid (SA) as the dopant. Typically, 5 g SA dissolved in 100 ml deionized water, then 1 ml An monomer added the above solution. Subsequently, the mixture was ultrasonication for 0.5 h, and the APS solution (1.5 g APS dissolved in 10 ml of hydrochloric acid solution (1 M)) was by drop to the mixture to start the polymerization. The reaction was performed with stirring at room temperature for 2 h. Finally, the product was removed from solution,
washed with distilled water until pH=6-7, and dried in vacuum at 80 °C for 12 h to obtain the pure PANI as black green powder.

2.2.5 Preparation of polyaniline/biomass carbon composites (PANI/KWB)

The PANI/KWB composites were prepared by in-site polymerization method. 5 g [CMMIm]Cl IL dissolved in 100 ml deionized water. Then add 1 ml An monomer and 0.1 g KWB (the mass ratio of An/KWB is 10:1) to the above solution. Subsequently, the mixture was sonicated for 0.5 h, and the APS solution (1.5 g APS dissolved in 10 ml of hydrochloric acid solution (1 M)) was added drop by drop to the mixture to start the polymerization. The reaction was performed with stirring at room temperature for 2 h. Finally, the product was removed from solution, washed with distilled water until pH=6-7, and dried in vacuum at 80 °C for 12 h to obtain the PANI/KWB-1 as black green powder. The other products were denoted as PANI/KWB-2, PANI/KWB-3, PANI/KWB-4 and PANI/KWB-5 for the mass ratios of An/KWB was 20:1, 30:1, 40:1 and 50:1 (the mass ratio of An monomer to IL and APS was kept at 1:5:1.5), respectively. In addition, we also prepared pure PANI as control.

2.2.6 Preparation of Fe₃O₄/KWB composites (KBFe)

KBFe composites were prepared via a coprecipitation method. In a typical process, 0.24 g FeCl₃·6H₂O and 0.10 g FeCl₂·4H₂O (the molar ratio of Fe³⁺/Fe²⁺ is 1.8:1) were dissolved in 100 ml deionized water, and then 0.2 g KWB was added with a vigorous stirring for 10 min. Subsequently, the NaOH solution (1 M) was added drop by drop to the mixture until the pH value of mixture reached to 11. Then the mixture was heated at 90 °C for 1 h with a vigorous stirring. Finally, the product was removed from the mixture solution by suction and washed several times with deionized water. The black solid product was dried in vacuum at 80 °C for 12 h to obtain the KBFe-1 composite (the theoretical value of mass ratio of Fe₃O₄ to KWB is 0.25:1). The other products were denoted as KBFe-2, KBFe-3, KBFe-4 and KBFe-5 for the theoretical value of mass ratio of Fe₃O₄ to KWB is 0.5:1, 0.75:1, 1:1 and 1.5:1, respectively.
2.3 Characterization

2.3.1 Morphological, structure, thermal and component characterizations

1) Scanning electron microscope (SEM)

The morphology and microstructure of samples were observed by scanning electron microscope (SEM, S-4300, Hitachi Co., Ltd., Tokyo, Japan). The sample was sputter-coated by gold with Ion sputter (E-1030, Hitachi Co., Ltd., Tokyo, Japan) for 2 min to provide enhanced conductivity. The test voltage was 5 KV or 15 KV, and an electric current was 10 μA.

2) Transmission electron microscopy (TEM)

The morphologies of the samples were also observed using a transmission electron microscope (TEM, TF20, FEI, Co., Ltd., USA). For TEM observations, samples were dispersed in ethanol and then a small drop of the suspension was spread onto a 400 mesh copper grid. The chemical composition was investigated by energy-dispersive X-ray spectroscopy (EDX).

3) Fourier transforms infrared spectroscopy (FTIR)

The chemical structures of the samples were confirmed by Fourier transform infrared spectroscopy (FTIR, IRT-7000, JASCO, Japan) with the KBr pellet method. The wavelength range was between 4000 and 600 cm\(^{-1}\), and the resolution was 4 cm\(^{-1}\). Fifty scans were averaged for each sample.

4) X-ray diffraction (XRD)

The crystallographic structure of samples was tested by X-ray diffraction system (XRD, XRD-6000, Shimadzu Co., Ltd., Kyoto, Japan) in range of \(2\theta = 5\text{–}90^\circ\) by step scanning with a diffract meter. Nickel-filter Cu K\(\alpha\) radiation (\(\lambda=0.15417\) nm) was used.
with a generator voltage of 40 kV and a current of 30 mA.

5) Nitrogen adsorption-desorption isotherms

The specific surface area and porosity of samples were proved using nitrogen adsorption-desorption isotherms which were performed by Brunauer-Emmett-Teller (BET) method of nitrogen sorption at 77 K using ASAP 2020 analyzer (Micromeritics, USA). All samples were outgassed under vacuum at 80 °C overnight prior to measurement. The specific surface area was calculated according to the Brunauer-Emmett-Teller (BET) equation, and the pore size distribution was derived from the adsorption branches of the isotherms.

6) X-ray photoelectron spectroscopy (XPS)

The chemical composition of samples was identified by X-ray photoelectron spectroscopy (PHI-5000, Perkin-Elmer Corp., USA) in a range of 0~1000 eV.

2.3.2 Electrochemical performance characterization

1) Preparation of working electrode

The fabrication of working electrodes was carried out as follows. For KWB materials, the sample, polytetrafluoroethylene (PTFE, the mass ratio of the sample and PTFE is 95:5), and ethanol, and grinding adequately to obtain a paste [6]. For KWBM, PANI/KWB, and KBFe composites, the sample, acetylene black, PTFE (the mass ratio of the sample, acetylene black and PTFE is 80:15:5) and ethanol, and grinding adequately to obtain a paste [7, 8]. Then the paste was coated onto the stainless mesh or nickel foam, which was followed by drying under vacuum at 80 °C for 24 h and then compressed at 10 MPa for 5 min. The mass loading of active material in an electrode is about 8 mg.
2) Electrochemical tests

The electrochemical tests were measured with an electrochemical workstation (CHI 660D, Shanghai Chenhua, China; and Models 1287, Solartron Analytical, UK) in the three or two-electrode system in electrolyte. In three-electrode system, the sample test electrodes were used as working electrode, a saturated calomel electrolyte as the reference electrolyte, and a platinum-wire electrodes as counter electrolyte. Cyclic voltammetry (CV) and galvanostatic charge/discharge (GCD) were conducted in a potential window. Electrochemical impedance spectroscopy (EIS) was accomplished at open circuit potential with perturbation of 5 mV in the frequency range from $10^2$-$10^{-2}$ Hz. For asymmetric two-electrode system, the system was assembled with two test electrodes as working electrodes. The CV and GCD tests were tested at different voltage windows.

The standard calomel reference electrode (SCE) based on GCD curves was evaluated according to the following equation [9, 10] (Eq. 1):

$$C_m = \frac{I \cdot t}{\Delta V \cdot m} \quad (\text{Eq. 2.1})$$

where $C_m$ (F g$^{-1}$) is the specific capacitance (SC), $I$ (A) is discharge current, $t$ (s) is the discharge time, $\Delta V$ (V) is the voltage window, and $m$ (g) is the total mass of active material on working electrode in a two-electrode system or three-electrode system.

The energy density ($E$, Wh kg$^{-1}$) and power density ($P$, W kg$^{-1}$) of the asymmetric supercapacitors were calculated from the discharge curves according to the following equations [11, 12]:

$$E = \frac{1}{2} C_m (\Delta V)^2 \quad (\text{Eq. 2.2})$$

$$P = \frac{E}{t} \quad (\text{Eq. 2.3})$$

where $C_m$ (F g$^{-1}$) is SC, $t$ (h) is the discharge time, and $\Delta V$ (V) is the potential window from GCD curves of two-electrode system, respectively.
References


Chapter 3 Biocarbon from sugarcane bagasse with superior performance in supercapacitors

3.1 Introduction

Since entering the 21st century, because of the exhausted fossil resource and population growth, energy shortage and environmental crisis become increasingly serious. This phenomenon propels people to find novel energy materials that can be commercialized in the near future, as well as the produce of materials are facile, environmentally friendly, low-cost, and nontoxic routes [1, 2]. Amongst various materials, carbon materials, especially structural carbon materials have attracted more and more attention of scholars, due to its potential applications in catalyst supports, adsorbents, carbon fixation, electrode, carbon fuel cells, cell biology, and gas storage [3-7]. Recently, with the rapid growth of nanotechnology, nanostructured carbon materials, such as carbon nanotubes (CNTs), graphene and porous carbons have been developed, all of them have the advantage of high surface area, electrical conductivity, chemical stability, fast energy storage-releasing, which are suitable as electrode materials to apply in supercapacitors and battery [8-10].

Supercapacitors is a new energy storage devices with considerable promise for a wide range of applications, including uninterruptable power sources, portable electronics, braking systems and hybrid electric vehicles [11-13]. According to the distinguishing energy storage mechanism, supercapacitors can be classified to electrical double-layer capacitors (EDLC) and pseudocapacitors. The EDLC is based on charge separation at the electrode/electrolyte interface, whereas the latter is based on redox reaction occurring within the active materials [14-17]. Currently, carbon-based EDLC materials are the most employed electrode material for commercially available supercapacitors due to their good cycle lifetime and high maximum [3, 18, 19].
At present, some important carbon materials also achieved high specific capacitance (Cs) and energy density, such as templated carbon [20], graphene [21] and carbon nanotube [22]. However, most of methods to prepare of the above carbon materials often require complicated preparation conditions or special facilities, leading to the high cost [23]. Therefore, a new research hotspot for preparation of high performance carbon materials are using cheap raw materials and simple method [24]. Recently, more and more scholars concentrate on the method of direct pyrolysis, as well as used the biomass or biowaste as carbon precursors [25, 26]. By virtue of low cost, renewability, availability, and environmental friendliness, make biomass become ideal candidates for producing structure carbon material with good capacitive performance [27, 28]. Many biocarbon have been prepared, such as pine cones [29], bamboo [30], waste coffee [31], oil palm [32] and so on. Sugarcane bagasse is a kind of typical biomass waste, which sometimes being processed by incineration or direct disposal. Obviously, this way not only wasted its use value, but also brought secondary pollution [32]. The discarded sugarcane bagasse waste residue is rich in lignin and cellulose, it can be used as the carbon source to make structure carbon. This is an efficient way to utilize sugarcane bagasse resources.

Hence, in this work, we report a simple method for preparation biocarbon (KWB) as the electrode materials for supercapacitors. KWB was obtained by direct pyrolysis of sugarcane bagasse after pretreatment by potassium hydroxide (KOH). The carbonization temperature and the mass ratio of KOH to sugarcane bagasse were researched as the influencing factors. We found that the two factors had a profound effect on the specific surface area of KWB, and further affect their electrochemical performance. The highest specific capacitance of KWB-3-800 reached 196.3 F g⁻¹ at a current density of 1 A g⁻¹, and this ensures KWB-3-800 can be used as high performance supercapacitors electrodes.
3.2 Experimental

3.2.1 Materials

Sugarcane bagasse was crushed by family-use grinder before use. The other reagents, such as ethanol, potassium hydroxide (KOH), and hydrochloric acid (HCl), all were purchased from Nacalai Tesque, Inc. (Kyoto, Japan) and of analytical grade, were used without further purification.

3.2.2 Preparation of KOH-activated biocarbon (KWB) derived from sugarcane bagasse

KWB was obtained by the KOH activation method and the typical process as follows: first, sugarcane bagasse was cut into small pieces (opening size approximately 2 mm) and washed with distilled water for 8 h at 90 °C, after that rinsed 2-3 times with deionized water and filtered, and then dried at the 80 °C for 24 h. Second, 1 g cleanly sugarcane bagasse and 0.5 g KOH were poured into 12 ml absolute ethanol with stirring and heated at 60 °C until all of the ethanol was evaporated, and then dried at 70 °C for 12 h. Third, the remnant mixture was pyrolyzed in a tubular furnace in N₂ atmosphere at 600 °C (heating rate is 10 °C min⁻¹) for 2 h. Finally, the obtained carbon material was washed by 1 M HCl solution for 2h, after that rinsed several times with deionized water till the filtrate became neutral and filtered, as well as collected after vacuum dry at the 40 °C for 12 h.

The carbonization temperature and the mass ratio of KOH to sugarcane bagasse were researched as the influencing factors. Firstly, the products with different mass ratio of KOH to sugarcane bagasse were be prepared at 800 °C, and denoted as KWB-1, KWB-2, KWB-3, KWB-4, and KWB-5 for the mass ratio of KOH to sugarcane bagasse is 0.2:1:0, 0.4:1, 0.6:1, 0.8:1, and 1:1, respectively. For comparison, the biocarbon without KOH activated was also prepared, which was denoted KWB-0. After that, on the basis of the best mass ratio of KOH to sugarcane bagasse, the different carbonation temperature was researched. The products was denoted as KWB-3-600, KWB-3-700,
KWB-3-800, and KWB-3-900 for the carbonization temperature is 600, 700, 800, and 900 °C, respectively.

Table 3.1 the preparation condition of KWB

<table>
<thead>
<tr>
<th>Samples</th>
<th>Mass ratio of KOH to bagasse</th>
<th>Carbonization temperature (°C)</th>
</tr>
</thead>
<tbody>
<tr>
<td>KWB-0</td>
<td>0:1</td>
<td>800</td>
</tr>
<tr>
<td>KWB-1</td>
<td>0.2:1</td>
<td>800</td>
</tr>
<tr>
<td>KWB-2</td>
<td>0.4:1</td>
<td>800</td>
</tr>
<tr>
<td>KWB-3</td>
<td>0.6:1</td>
<td>800</td>
</tr>
<tr>
<td>KWB-4</td>
<td>0.8:1</td>
<td>800</td>
</tr>
<tr>
<td>KWB-5</td>
<td>1:1</td>
<td>800</td>
</tr>
<tr>
<td>KWB-3-600</td>
<td>0.6:1</td>
<td>600</td>
</tr>
<tr>
<td>KWB-3-700</td>
<td>0.6:1</td>
<td>700</td>
</tr>
<tr>
<td>KWB-3-800</td>
<td>0.6:1</td>
<td>800</td>
</tr>
<tr>
<td>KWB-3-900</td>
<td>0.6:1</td>
<td>900</td>
</tr>
</tbody>
</table>

3.2.3 Structure characterization

The morphology and microstructure of samples were observed by scanning electron microscopy (SEM) at S-4300 SEM instrument (Hitachi, Japan). The specific surface areas of the samples were determined by Brunauer-Emmett-Teller (BET) method of nitrogen sorption at 77 K using ASAP 2020 analyzer (Micromeritics, USA). The crystallographic structure was carried out via X-ray diffraction (XRD) measurements at XRD-6000 instrument (Shimadzu, Japan) with Cu-Kα radiation (λ=0.154 nm) at a speed of 5° min⁻¹ from 2θ = 5° to 90°. The surface states of samples were analyzed via X-ray photoelectron spectroscopy (XPS) using a PHI Quantum 5000 instrument (ULVAC-PHI, Japan) equipped with Al Kα radiation source.

3.2.4 Electrochemical testing

Conventional three-electrolyte system and asymmetric two-electrode system were used in supercapacitors to measure the electrochemical characterization of KWB using Models 1287 electrochemical workstations (Solartron Analytical, UK). The test
electrodes were first prepared by mixing the KWB with Polytetrafluoroethylene (PTFE) at a mass ratio of 95:5, and ethanol, then the mixing were adequately ground to obtain a homogeneous paste for coating onto the stainless mesh to forming a current collector. After that, the test electrodes were dried in vacuum at 80 °C for 24 h, then compressed at 10 MPa for 5 min. In three-electrode system, the KWB test electrodes was used as working electrode, a saturated calomel electrolyte as the reference electrolyte, and a platinum-wire electrodes as counter electrolyte. Cyclic voltammetry (CV) and galvanostatic charge/discharge (GCD) were conducted in a potential window from -0.4 to 0 V for KWB. Electrochemical impedance spectroscopy (EIS) was accomplished at open circuit potential with perturbation of 5 mV in the frequency range from $10^2$-$10^2$ Hz. A 1M H$_2$SO$_4$ solution was used as an electrolyte for all electrochemical measurements.

The standard calomel reference electrode (SCE) based on GCD curves was evaluated according to the following equation [33, 34] (Eq. 1):

$$C_m = \frac{I \cdot t}{\Delta V \cdot m} \quad \text{(Eq. 1)}$$

where $C_m$ (F g$^{-1}$) is the specific capacitance (SC), I (A) is discharge current, t (s) is the discharge time, $\Delta V$ (V) is the voltage window, and m (g) is the total mass of active material on working electrode in three-electrode system.

### 3.3 Results and discussion

#### 3.3.1 Microstructural characterization

Fig. 3.1 illustrates the preparation process of KWB frameworks. At first, the KWB was obtained by carbonized and activated sugarcane bagasse at one step (as shown in Fig. 4.1a) [35]. As shown in Fig. 4.1b, the sugarcane bagasse exhibited a typical bundle-microtube-like structure of the biological tissue. While after carbonization and activation, the special structure was destroyed and an allium-giganteum-like structure with high interconnected frameworks which is favorable for the ion diffusion and suitable for preparation of carbon-based composites , can be clearly found in the as-
resulted KWB (as shown in Fig. 4.1e and f) [36]. The process of KOH activating carbon at high temperature be researched by some scholars [37, 38].

Fig. 3.1 Schematic illustration of the preparation process of KWB samples. Digital photographs of sugarcane bagasse (a), KWB (b); SEM images with different magnification of the sugarcane bagasse (c and d), and KWB (e and f).

In short, this process through three aspects: firstly, the formation of H$_2$O and CO$_2$ can contribute to porosity development by carbon gasification; then, the formation of K$_2$O and K$_2$CO$_3$ can etch the carbon framework by redox reactions, which is beneficial to form the pore structure; finally, the formation of metallic K is an important factor for developing the pore structure by inserting into the graphitic layers. What’s more, the reaction occurring between KOH and the carbon materials are as follows [39]:

\[
2\text{KOH} \rightarrow \text{K}_2\text{O} + \text{H}_2\text{O} \quad (1) \\
\text{H}_2\text{O} + \text{C} \rightarrow \text{CO} + \text{H}_2 \quad (2) \\
\text{CO} + \text{H}_2\text{O} \rightarrow \text{CO}_2 + \text{H}_2 \quad (3) \\
\text{CO}_2 + \text{K}_2\text{O} \rightarrow \text{K}_2\text{CO}_3 \quad (4) \\
\text{K}_2\text{CO}_3 \rightarrow \text{K}_2\text{O} + \text{CO}_2 \quad (5) \\
\text{CO}_2 + \text{C} \rightarrow 2\text{CO} \quad (6) \\
\text{K}_2\text{O} + 2\text{C} \rightarrow \text{K} + 2\text{CO} \quad (7) \\
\text{K}_2\text{CO}_3 + \text{C} \rightarrow 2\text{K} + 3\text{CO} \quad (8)
\]
The morphology and microstructure of the prepared KWB samples were investigated by SEM, and Fig. 3.2 shows the samples with different mass ratios of KOH/bagasse at 800 °C. The KWB-0 is the sample without KOH activating, and it can be clearly seen that the surface of KWB-0 is smooth without porous (as shown in Fig. 3a). However, the other KWB samples including from KWB-1 to KWB-5, all of them with a lot of porous on the surface can be clearly observed. Moreover, it is noteworthy that the mass ratio of KOH/bagasse has an important increasing the porous’ morphology and number is very different. When the mass ratio of KOH/bagasse less than 0.6:1, the number and diameter of the surface porous increase with the increase of the mass ratio. For KWB-1 (Fig.3.2 (b)), there are only a little of porous with the diameter at 1μm on the surface, while KWB-2 (Fig.3.2 (c)) can be directly observed the more porous with larger diameter than the porous structure of KWB-1. Furthermore, a large number of secondary porous with a diameter of about 0.5μm are present on the inner wall of the primary porous having a diameter of 2 mm, it means that KOH pretreatment makes KWB forming hierarchically porous structure biocarbon material with interconnected frameworks. When the mass ratio of KOH/bagasse more than 0.6:1, the excessive activation of KOH destroyed the interconnected frameworks structure of biocarbon material. For KWB-5 (Fig.3.2 (f)) the mass ratio of KOH/bagasse is 1:1, the surface of the primary porous of carbon skeleton has been completely collapsed, and only could
see a lot of debris and secondary porous with the diameter at 0.5 secondary hole. In summary, the SEM image of KWB samples shows the important influence of the mass ratio of KOH/bagasse for KWB samples, and KWB-3 (Fig.3.2 (d)) shows the perfect interconnected frameworks structure with multi-stage porous, which means the best activated degree of KOH activation agent is the mass ratio of KOH/bagasse at 0.6:1. In addition, the structure of carbon material may profoundly impact electrochemical performance of KWB samples.

Fig. 3.3 SEM images of KWB-3-600 (a), KWB-3-700 (b), KWB-3-800 (c), KWB-3-900 (d), simples.

The morphology of KWB samples with different carbonization temperature were also investigated by SEM. All the KWB samples have a lot of porous on the surface of samples, while the porous structures have a little different. It can be clearly observed that follow the increasing of carbonization temperature the diameter and number of porous were increasing, and KWB-3-800 shown the excellent interconnected frameworks structure with multi-stage porous at the 800 °C of carbonization temperature. In addition, the interconnected frameworks structure of KWB-3-900
shown more serious damaged, that may be due to the too high temperature to preserve the multi-stage porous. Hence, the best carbonization temperature of sugarcane bagasse is 800 °C, and the above phenomena may impact electrochemical performance of KWB samples profoundly.

Fig. 3.4 X-ray diffraction patterns of the prepared KWB-1, KWB-2, KWB-3, KWB-4 and KWB-5 samples (a), and KWB-3-600, KWB-3-700, KWB-3-800, and KWB-3-900 samples.

The crystallinity of the samples was examined by X-ray diffraction analysis. Fig. 3.4 shows the XRD patterns of KWB samples. All of KWB samples can be observed two diffraction peaks. A board diffraction peak centered at 25° correspond to the (002) plane of graphite, indicating the amorphous nature [36]. Additionally, a weak diffraction peak centered at 43.4° can be attributed to the (101) plane of graphite, which means of the complete transformation of sugarcane bagasse to another kind of activated carbon [40]. However, every patterns have a little different. In Fig. 3.4(a), the intensity of board peak at 25° decreased following the increasing of mass ratio of KOH/bagasse, and same effect also appeared with the increasing of carbonization temperature. This phenomenon may be due to the reducing concentration of nonparallel single layers carbon when more activation agent or higher carbonization temperature roles in the raw materials [38].
Fig. 3.5 N\textsubscript{2} adsorption-desorption isotherms (a, and b) and pore size distribution by BJH method (c, and d) of the prepared KWB-1, KWB-2, KWB-3, KWB-4, KWB-5, KWB-3-600, KWB-3-700, KWB-3-800, and KWB-3-900 samples.

As we all know that the surface area and the pore-size distribution are the two key factors for the electrochemical performance of electrode materials. The specific surface area and pore-size distribution of KWB samples were characterized by nitrogen adsorption/desorption measurements (as shown in Fig. 4.6a, b, c, and d). It is clear that all KWB shows the typical combined I/IV type sorption isotherms, indicating the polyporous structure [41]. However, the KWB-3-800 shows the best volume absorbed and pore volume, due to the presence of more nanoscale porous than other KWB samples (Fig. 3.5d). As given in Table 2, the $S_{BET}$ of KWB-3-800 is up to 1717.91 m\textsuperscript{2} g\textsuperscript{-1}, which results in KWB-3-800 provides more reaction sites for efficient electrochemical interactions/reactions. What’s more, it also can be used as carbon-based material for preparing composite materials.
Table 3.2 Specific surface area, pore size of KWB samples. ($S_{BET}$ - the specific surface area; $P$ - the average pore size; $V_t$ - the total pore volume)

<table>
<thead>
<tr>
<th>Sample</th>
<th>$S_{BET}$ (m$^2$ g$^{-1}$)</th>
<th>$V_t$ (cm$^3$ g$^{-1}$)</th>
<th>$P$ (nm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>KWB-1</td>
<td>396.62</td>
<td>0.232</td>
<td>4.67</td>
</tr>
<tr>
<td>KWB-2</td>
<td>997.27</td>
<td>0.564</td>
<td>4.23</td>
</tr>
<tr>
<td>KWB-3</td>
<td>1717.91</td>
<td>0.975</td>
<td>2.58</td>
</tr>
<tr>
<td>KWB-4</td>
<td>1433.06</td>
<td>0.755</td>
<td>3.03</td>
</tr>
<tr>
<td>KWB-5</td>
<td>991.90</td>
<td>0.539</td>
<td>4.22</td>
</tr>
<tr>
<td>KWB-3-600</td>
<td>784.55</td>
<td>0.409</td>
<td>2.71</td>
</tr>
<tr>
<td>KWB-3-700</td>
<td>1079.07</td>
<td>0.588</td>
<td>2.63</td>
</tr>
<tr>
<td>KWB-3-800</td>
<td>1717.91</td>
<td>0.975</td>
<td>2.58</td>
</tr>
<tr>
<td>KWB-3-900</td>
<td>1303.74</td>
<td>0.729</td>
<td>2.85</td>
</tr>
</tbody>
</table>

Moreover, the specific surface area and pore distribution of KWB samples can be controlled by adjust the mass ratio of KOH/bagasse and carbonization temperature. As shown in Table 3.2, the specific surface area and pore volume of KWB was significantly increased firstly and then decreased with the mass ratio of KOH/bagasse increased, and during the change of carbonization shown the similar trend. On the contrary, the average pore size of KWB was significantly decreased firstly and then increased with the mass ratio of KOH/bagasse and carbonization temperature increased. This result further confirms that excessive activation of KOH may further react with graphite crystallite structure around the pore, leading to excessive burn-off of carbon surface, which will result in breakage in porous carbon structure, and thereby both surface area and pore volume decrease substantially [42]. On the other hand, the carbonization temperature also plays a vital role in determining the BET surface area, but excess higher temperature may destroy the already formed interconnected frameworks structure and reduce the specific surface area of the material, which would reduce the relative area that can actually participate in the electrochemical reaction and has an effect on the electrochemical performance of this material as electrode material in applied. Overall, at optimum mass ratio of KOH/bagasse and carbonization temperature, the BET surface area is of maximum, which is achieved at 0.6:1 and carbonization at 800 °C. What’s
more, this result is also confirmed with SEM.

Fig. 3.6 XPS survey spectra of KWB-1, KWB-2, KWB-3, KWB-4 and KWB-5 samples (a), KWB-3-600, KWB-3-700, KWB-3-800, and KWB-3-900 samples (b); high resolution C1s (c) and O1s (e) XPS spectra of KWB-3-800 sample.

The information of the surface chemical composition of samples was obtained from XPS spectra. The full scan spectra of samples are shown in Fig. 4.6a and b. All of the KWB have the signal of C and O element at the around of 535 and 284 eV. More information about the change of surface chemistry has been obtained by the deconvoluted high-resolution XPS spectra of three regions (C, and O as shown in Fig. 4.6c and d). The C1s spectra of KWB-3-800 could be deconvoluted into three peaks (as shown in Fig. 5.8c) with binding energies of 531.5, 533.2 and 534.8 eV. These peaks could be assigned to aromatic or other sp2-hyridised carbon atoms bound to neighboring carbon atoms (C-C, 284.4 eV), carbon in C-O (285.8 eV) and C=O (288.8 eV), respectively [43]. The O1s spectra of KWB-3-800 in Fig. 4.6d can be deconvoluted
into three components. The peaks of 531.5, 533.2 and 534.8 eV can be assigned to –OH/C-O, C=O and C-O-H groups, respectively [44].

<table>
<thead>
<tr>
<th>Sample</th>
<th>C at. %</th>
<th>O at. %</th>
</tr>
</thead>
<tbody>
<tr>
<td>KWB-1</td>
<td>89.63</td>
<td>10.37</td>
</tr>
<tr>
<td>KWB-2</td>
<td>89.88</td>
<td>10.02</td>
</tr>
<tr>
<td>KWB-3</td>
<td>90.95</td>
<td>9.05</td>
</tr>
<tr>
<td>KWB-4</td>
<td>91.89</td>
<td>8.11</td>
</tr>
<tr>
<td>KWB-5</td>
<td>88.87</td>
<td>11.13</td>
</tr>
<tr>
<td>KWB-3-600</td>
<td>92.33</td>
<td>7.67</td>
</tr>
<tr>
<td>KWB-3-700</td>
<td>89.59</td>
<td>10.41</td>
</tr>
<tr>
<td>KWB-3-800</td>
<td>90.95</td>
<td>9.05</td>
</tr>
<tr>
<td>KWB-3-900</td>
<td>88.95</td>
<td>11.05</td>
</tr>
</tbody>
</table>

The content of elements of every KWB samples are listed in Table 3.3. It is easy to find that all the KWB samples have approximately 10% of oxygen elements. These oxygen elements are mainly derived from cellulose, hemicellulose and lignin in bagasse, which are not involved in the activation reaction under high temperature, and formed C-O or C=O band with adjacent carbon elements. Many of published literatures have demonstrate that the addition of heteroatoms can significantly reduce the resistance of charge transfer. On the other hand, carbonaceous materials are evidenced to reduce the pathway for the electron transfer from active electrode materials to the electrolyte and facilitate fast diffusion of electrolyte ions [27, 45, 46]. The above phenomena may impact electrochemical performance of KWB samples profoundly.

3.3.2 Electrochemical performance of KWB samples

In order to evaluate the electrochemical characteristics of KWB samples, CV, GCD, and EIS measurements were employed to characterize the capacitive properties in 1 M H$_2$SO$_4$ aqueous solution in a three-electrode configuration. Fig. 3.7 shows the CV curves of KWB samples for different mass ratio of KOH/bagasse (Fig. 3.7a) and
different carbonization temperature (Fig. 3.7b) at a scan rate of 0.01 V s\(^{-1}\). The CV curve of all KWB samples shows nearly symmetric rectangular shape, indicating an ideal double-layer [47]. The offset from the standard rectangle is attribute to the reaction of a small amount of oxygen-containing groups in the material [45]. Obviously, the area enclosed by the CV curve, which is proportional to the capacitance, increases firstly and then decreased with the increase of the mass ratio of KOH/bagasse and carbonization temperature. However, KWB-3-800 shows the larger area enclosed by the CV curve than that of the KWB samples, revealing the highest SC.

![Fig. 3.7 CV curves of KWB-1, KWB-2, KWB-3, KWB-4 and KWB-5 samples (a), KWB-3-600, KWB-3-700, KWB-3-800, and KWB-3-900 samples (b).](image)

![Fig. 3.8 CV curves of KWB-1, KWB-2, KWB-3, KWB-4 and KWB-5 samples (a), KWB-3-600, KWB-3-700, KWB-3-800, and KWB-3-900 samples (b).](image)
The electrochemical performance of as-prepared samples was further analyzed by the GCD method and the results are shown in Fig. 3.8. The discharge time of KWB-3-800 is markedly longer than those of other samples, suggesting that the KWB-3-800 possesses further more excellent electrochemical capacitive performance than the other samples. We can get the specific capacitance by these curves [48]. From Eq. 3.1, the specific capacitances of KWB-1, KWB-2, KWB-3, KWB-4, KWB-5, KWB-3-600, KWB-3-700, KWB-3-800, and KWB-3-900 are 85, 135.6, 196.3, 161.4, 150.4, 62.2, 88.1, 196.3 and 132.1 F g\(^{-1}\), respectively, at a current density is 1 A g\(^{-1}\).

![Fig. 3.9 Specific capacitance versus current density of KWB samples.](image)

Fig. 3.9 Specific capacitance versus current density of KWB-1, KWB-2, KWB-3, KWB-4 and KWB-5 samples (a), KWB-3-600, KWB-3-700, KWB-3-800, and KWB-3-900 samples (b).

Furthermore, the SC of KWB (as shown in Fig. 3.9) increases first and then decreases with the increase of the mass ratio of KOH/bagasse and carbonization temperature. These results are in accordance with those deduced from CV curves. Furthermore, more interesting is that in the KWB samples, it appears a sudden potential drop (IR drop), and the IR drop increases with the increase of KOH/bagasse mass ratio. This can be attributed to the resistance of electrolyte and the inner resistance of ion diffusion in hierarchically porous carbon [49, 50].

To further study the capacitive behavior samples, EIS was used to measure the charge transport and ion diffusion. The Nyquist plots of KWB samples are showed in Fig. 3.9.
In EIS curve, the internal resistance (Rs, including the intrinsic resistance of the sample, the contact resistance between the sample and current collector, and the resistance of bulk electrolyte) can be obtained from the point intersecting with the real axis in the region of high frequency. The semicircle is due to the charge transfer and which will affect capacitive behavior at high current loading [51]. The inset shows that Rs of KWB-1, KWB-2, KWB-3, KWB-4, KWB-5, KWB-3-600, KWB-3-700, KWB-3-800, and KWB-3-900 are 1.30, 1.12, 0.70, 1.35, 1.60, 1.67, 1.19, 0.70 and 1.34 Ω, respectively. It is worth noting that Rs of KWB-3-800 is lower than those of KWB samples. The semicircle radius of KWB-3-800 is lower than the other samples, means that the KWB-3-800 possesses the lowest charge resistance among KWB samples. Moreover, the nearly vertical line at lower frequency of KWB-3-800 indicates the pure capacitor behavior as an ideal capacitor [46, 47]. In short, the analyses of EIS reveal that KWB-3-800 sample possesses an excellent capacitive performance, and this result agrees with the analyses of GCD and CV.

Fig. 3.10 EIS curves of KWB-1, KWB-2, KWB-3, KWB-4 and KWB-5 samples (a), KWB-3-600, KWB-3-700, KWB-3-800, and KWB-3-900 samples (b).

Fig. 3.11a shows the CV curves of KWB-3-800 samples at different scan rates ranging from 10 to 50 mV s⁻¹. Generally, the response current of KWB-3-800 gradually increases when increasing the scan rate. The CV curve has maintained the rectangular shape, and a rapid current response on voltage reversal at each end potential at the scan
rates ranging from 10 to 50 mV s\(^{-1}\). These reveal its good electrochemical capacitive performance at high scan rates. Also note that, the pair of weak peaks were also clearly observe at the scan rate of 50 mV s\(^{-1}\), which may be because oxygen-groups still participate in the reaction under high current, indicating that the material has good stability. Fig. 3.11b shows the specific capacitances of KWB-3-800 at different current densities from 0.5 to 10 A g\(^{-1}\). Clearly, the downward trend of specific capacitance of KWB-3-800 is nonlinear, which suggests that the pseudocapacitive behavior happens at the electrode-electrolyte interface [52]. From Eq. 3.1, the specific capacitances reach 196.3 and 123 F g\(^{-1}\) at the current density of 1 and 15 A g\(^{-1}\), respectively, and the capacitance retention of KWB-3-800 is 62.7 %. These results indicate that the KWB-3-800 possesses the fast charge-discharge properties, which is very important for supercapacitors application.

Fig. 3.11 CV curves of KWB-3-800 at different scan rates from 10 to 50 mVs-1 (a); GCD curves of KWB-3-800 at different current density from 0.5 to 10 A g-1(b)

3.4 Conclusions

In summary, a simple method for preparation KWB as the electrode materials for supercapacitors. KWB was obtained by direct pyrolysis of sugarcane bagasse after pretreatment by potassium hydroxide (KOH). The electrochemical performance of KWB was closely related to the mass ratio of KOH/bagasse and carbonization temperature. KWB-3-800 showed the best electrochemical performance when the mass
ratio of KOH/bagasse was 0.6 and the carbonization temperature is 800 °C in three-electrode system. In addition, the specific capacitances reach 196.3 and 123 F g⁻¹ at the current density of 1 and 15 A g⁻¹, respectively, and the capacitance retention of KWB-3-800 is 62.7 %. The preparation of KWB samples with a facile and low-cost method shows an excellent electrochemical performance, marking it a promising candidate in energy storage devices.
Reference


[10] Y. Liu, J. Liu, M. Hou, L. Fan, Y. Wang, Y. Xia, Carbon-coated Li4Ti5O12 nanoparticles with high electrochemical performance as anode material in sodium-


[41] K.S.W. Sing, Reporting physisorption data for gas/solid systems with special reference to the determination of surface area and porosity (Recommendations


Chapter 4 Manganese dioxide/biocarbon composites with superior performance in supercapacitors

4.1 Introduction

Supercapacitors, is one of the most promising energy storage devices with fast charge-discharge rates, superior power density and long cycle lifetime compared to the other chemical energy storage devices [1, 2]. According to the energy storage mechanism, supercapacitors are classified as electrical double-layer capacitors (EDLC) and pseudocapacitors. The EDLC is based on charge separation at the electrode/electrolyte interface, whereas the latter is based on redox reaction occurring within the active materials [3-5]. Currently, carbon materials are often used for EDLC due to their high electrical conductivity and fast energy storage-releasing, while the transition metal oxides often used for pseudocapacitors because of multiple oxidation states at the charge-discharge process [6, 7].

Among the multifarious transform metal oxides, MnO₂ is one of the most promising electrode materials for low cost, high theoretical capacity and environmental friendly supercapacitors devices [8]. However, the performances of single-phased MnO₂ are still insufficient to meet growing demands in supercapacitors due to its intrinsic properties, such as low conductivity and poor mechanical stability [9, 10]. To solve these problems, one-dimensional (1D) nanostructure MnO₂ (such as nanosheets MnO₂) has been found to be good candidate for its low density and excellent permeation [11, 12]. Nevertheless, 1D MnO₂ still does not achieve the desired electrochemical performance due to its electrochemical dissolution during cycling [13]. Therefore, development of new type MnO₂ materials to meet the desired electrochemical performance is urgently desirable.

To respond the above requirement, carbon materials have been select as a component for preparing MnO₂-based composites to effectively improve electrochemical
performance [14]. At present, many kinds of carbon materials, such as activated carbon (AC) [15], graphene [16] and carbon nanotube [17] are often used as the carbon content in MnO$_2$-based composites because their high surface area, excellent electronic conductivity, unique internal structure and superior thermal stability [18]. However, most of methods for prepare of the above carbon materials often require complicated preparation conditions or special facilities, leading to the high cost [19]. For example, as a rising star of graphene, it is often obtained by reduction of graphene oxide (rGO), and graphene oxide (GO) often obtained from natural graphite by Hummers method [20]. The method is a complex process, and contains four steps: preoxidation, further oxidation, decomposition of excessive oxidant and removed metal ions [21]. Therefore, one preponderant method of preparation of carbon materials is direct pyrolysis of carbon precursors, such as pyrolysis of biomass or biowaste [22]. This method has been noticed by many researchers, and many biocarbon derived from biomass such as corncobs [23], cocoons [24] and watermelon rind [25] have been prepared. All the biocarbon materials described above present high specific surface area and abundant pore structure. Therefore, the biocarbon materials are very suitable for preparation of MnO$_2$-based composites.

Hence, in this work, we report an efficient method for preparation of MnO$_2$/biocarbon composites (KWBM) as the electrode materials for supercapacitors. The biocarbon (KWB) was obtained by direct pyrolysis of sugarcane bagasse, and then the KWBM was obtained by redox reaction between KMnO$_4$ and KWB under different mass ratios of KMnO$_4$/KWB. We found that the chemical component and specific surface area of KWBM had a profound effect on electrochemical performance of KWBM. In addition, an asymmetric two-electrode system was fabricated by employing KWBM-4 and KWB as positive and negative electrode, respectively. We also found that the asymmetric system possessed the high energy density and high cycle stability.
4.2 Experimental

4.2.1 Materials

Sugarcane bagasse was crushed by family-use grinder before use. The other reagents, such as ethanol, potassium hydroxide (KOH), hydrochloric acid (HCl) and potassium permanganate (KMnO₄), all were purchased from Nacalai Tesque, Inc. (Kyoto, Japan) and of analytical grade, were used without further purification.

4.2.2 Preparation of KOH-activated biocarbon (KWB) derived from sugarcane bagasse

KWB was obtained by KOH activation method and the typical process as follows: first, sugarcane bagasse was cut into small pieces (opening size approximately 2 mm) and washed with distilled water for 8 h at 90 °C, and then dried at 80 °C for 24 h. Second, 1 g cleanly sugarcane bagasse and 0.6 g KOH were poured into 12 ml absolute ethanol with stirring and heated at 60 °C until all of the ethanol was evaporated, and then dried at 70 °C for 12 h. Third, the remnant mixture was pyrolyzed in a tubular furnace in N₂ atmosphere at 800 °C (heating rate is 10 °C min⁻¹) for 2 h. Finally, the obtained carbon material was washed by 1 M HCl solution and deionized water till the filtrate became neutral and collected after vacuum dry.

4.2.3 Preparation of MnO₂/KWB composites (KWBM)

The KWBM composites were prepared by directly reacting KMnO₄ with KWB. In a typical process, 0.05 g KMnO₄ dissolved in 100 ml deionized water, and then 0.1 g KWB (the mass ratio of KMnO₄/KWB is 0.5:1) added the above solution. Subsequently, the mixture was stirred under water bath at 70 °C until the color of solution changed from modena to tawny. The products removed from solution and washed with deionized water for several times, and dried in vacuum at 80 °C for 12 h to obtain of the KWBM-1 composite. The other products were denoted as KWBM-2, KWBM-3 and KWBM-4.
for the mass ratio of KMnO$_4$/KWB is 1:1, 1.5:1 and 2:1, respectively.

### 4.2.4 Structure characterization

The morphology of samples were observed by transmission electron microscopy (TEM, H-8110, Hitachi Co., Ltd. Tokyo, Japan) and scanning electron microscopy (SEM, S-4300, Hitachi Co., Ltd. Tokyo, Japan). Nitrogen adsorption-desorption isotherms were measured at –196 °C on a Micromeritics ASAP 2020 apparatus. The specific surface area was calculated according to the Brunauer-Emmett-Teller (BET) equation. All samples were outgassed under vacuum at 80 °C overnight prior to measurement. The crystallographic structure of the samples was observed by X-ray diffraction (XRD, XRD-6000, Shimadzu Co., Ltd. Kyoto, Japan) equipped with Cu Ka radiation. X-ray photoelectron spectroscopy (XPS) analysis was performed on the PHI Quantum 5000 equipped with an Al Ka radiation source.

### 4.2.5 Electrochemical test

The electrochemical tests were measured with an electrochemical workstation (CHI 660D, Shanghai Chenhua, China) in the two or three-electrode system in 1M KOH electrolyte. For three-electrode system, saturated calomel electrolyte and platinum-wire electrode were used as reference and counter electrode, respectively. The test electrodes were first prepared by mixing the KWB and polytetrafluoroethylene (PTFE) at a mass ratio of 95:5, and mixing KWBM with carbon black (CB) and PTFE at a mass ratio of 80:15:5. After that, the mixing were grinding adequately to obtained a homogeneous paste, that was coated onto the Ni foam for current collector and dried under vacuum at 80 °C for 24 h, and then compressed at 10 MPa for 5 min. Cyclic voltammetry (CV), and galvanostatic charge/discharge (GCD) were conducted in a potential window from -0.8 to 0 V for KWB biocarbon, and 0 to 0.7 V for KWBM composites respectively. Electrochemical impedance spectroscopy (EIS) was carried out in the frequency range from $10^5$ to $10^{-2}$ Hz at open circuit potential with as ac perturbation of 5 mV. The symmetric two-electrode system was assembled with two almost identical working
electrodes (the active material was KWBM-4 composites sample, and the counter electrode was KWB biocarbon sample). The CV and GCD tests were run at different voltage windows.

The standard calomel reference electrode (SCE) based on GCD curves was evaluated according to the following equation (Eq. 1):

\[ C_m = \frac{I \times t}{\Delta V \times m} \]  

(Eq. 1)

where \( I \) is discharge current (A), \( t \) is the discharge time (s), \( \Delta V \) is the voltage window (V), and \( m \) is the mass of active material on working electrode in three-electrode system, in two-electrode system is the total mass of active material on two working electrodes [26].

4.3 Results and discussion

4.3.1 Microstructural characterization

Fig. 4.1 illustrates the preparation process of KWB frameworks and KWBM composites. At first, the KWB was obtained by carbonized and activated sugarcane bagasse at one step (as shown in Fig. 4.1a) [3]. As shown in Fig. 4.1d and e, the sugarcane bagasse exhibited a typical bundle-microtube-like structure of the biological

![Fig. 4.1 Schematic illustration of the preparation process of KWBM composites. Digital photographs of sugarcane bagasse (a), KWB (b) and KWBM-3 (c). SEM images with different magnification of the sugarcane bagasse (d and e), KWB (f and g) and KWBM-3 composite (h).](image)
tissue. While after carbonization and activation, the special structure was destroyed and an allium-giganteum-like structure with high interconnected frameworks which is favorable for the ion diffusion and suitable for preparation of MnO2-based composites.
[27], can be clearly found in the as-resulted KWB (as shown in Fig. 4.1f and g). Therefore, KWB was used as both scaffolds and reducing agents for MnO$_2$ growth in this study. MnO$_2$ nanosheets grew on the surface of KWB based on the self-limiting reaction (Re. 1) between KMnO$_4$ and KWB in neutral aqueous solution [28, 29].

$$4\text{MnO}_4^- + 3\text{C} + \text{H}_2\text{O} \rightarrow 4\text{MnO}_2 + \text{CO}_3^{2-} + 2\text{HCO}_3^- \quad \text{(Re. 1)}$$

The morphology and microstructure of the prepared KWBM composites are shown in Fig. 4.2. The surface and interconnected frameworks of KWBM were coated by MnO$_2$ nanosheets in all samples. Moreover, it is noteworthy that a little of uncoated KWB was observed in the red circle of Fig. 4.2a (KWBM-1) and c (KWBM-2). The uncoated KWB gradually decreased with the mass ratio of KMnO$_4$/KWB increased till, KWB was completely coated by MnO$_2$ nanosheets when the mass ratio of KMnO$_4$/KWB is 2 (as shown in Fig. 4.2g). In addition, the high resolution SEM image of samples (Fig. 4.2b, d, f and h) showed the size of MnO$_2$ nanosheets gradually increased with the mass ratio of KMnO$_4$/KWB increased. The above phenomena may impact electrochemical performance of KWBM composites profoundly.

Fig. 4.3 TEM images of the KWBM-4 composite (a, b); the inset in a shows the corresponding SAED pattern taken from the edge of MnO$_2$ nanosheets; HRTEM image of the birnessite-type MnO$_2$ nanosheets (b); the EDX mapping images of the KWBM-4 composite (c).
The morphology of KWBM-4 can be further investigated by TEM. It can be clearly seen that the surface of KWB is coated by MnO₂ nanosheets (as shown in Fig. 4.3a). In addition, the nanosheets of MnO₂ structure is confirmed by HRTEM image (as shown in Fig. 4.3b). The interplanar spacing of MnO₂ nanosheets is 0.69 nm, corresponding to the typical interplanar spacing of the (001) plane of δ-MnO₂ which prefers forming sheet structures [28]. Fig. 4.3c is the energy-dispersive X-ray (EDX) mapping results of KWBM-4. The results reveal the co-existence and homogenous distribution of C, Mn and O across the core-shell nanostructure of KWBM-4. Moreover, the EDX mappings of other KWB composites are showed in Fig. 4.4, and the chemical composition of KWBM can be obtained by EDX analysis (as shown in Table 4.1). For KWBM composites, the content of C and O gradually decreased and increased, respectively, with the mass ratio of KMnO₄/KWB increased. The ratio of Mn and C content also increased with the mass ratio of KMnO₄/KWB increased. The chemical composition of samples will be further analyzed by XPS spectra.

![Fig. 4.4 The EDX mapping images of the KWB (a), KWBM-1 (b), KWBM-2 (c) and KWBM-3 (d) composites.](image_url)
Table 4.1 EDX analysis of chemical compositions

<table>
<thead>
<tr>
<th>Samples</th>
<th>C/at. %</th>
<th>O/at. %</th>
<th>Mn/at. %</th>
<th>Mn/C %</th>
</tr>
</thead>
<tbody>
<tr>
<td>KWB</td>
<td>89.8</td>
<td>10.2</td>
<td>--</td>
<td>--</td>
</tr>
<tr>
<td>KWBM-1</td>
<td>57.7</td>
<td>15.9</td>
<td>26.4</td>
<td>45.8</td>
</tr>
<tr>
<td>KWBM-2</td>
<td>52.6</td>
<td>21.8</td>
<td>25.6</td>
<td>48.7</td>
</tr>
<tr>
<td>KWBM-3</td>
<td>47.3</td>
<td>26.5</td>
<td>26.2</td>
<td>55.4</td>
</tr>
<tr>
<td>KWBM-4</td>
<td>44.9</td>
<td>27.7</td>
<td>27.4</td>
<td>61.0</td>
</tr>
</tbody>
</table>

Fig. 4.5 X-ray diffraction patterns of the prepared KWB biocarbon, KWBM-1, KWBM-2, KWBM-3 and KWBM-4 composites.

The crystallinity of the samples was examined by X-ray diffraction analysis. Fig. 4.5 shows the XRD patterns of KWB and KWBM. A broad diffraction peak centered at 23.1° correspond to the (002) plane of graphite, indicating the amorphous nature [27]. Additionally, a weak diffraction peak centered at 43.4° can be attributed to the (101) plane of graphite, which means of the complete transformation of sugarcane bagasse to another kind of activated carbon [26]. For KWBM samples, the other three diffraction peaks appearing at 12.1°, 36.6° and 65.5° are indexed to the diffraction of (001), (006) and (119) planes of δ-MnO₂, respectively (JCPDS no. 18-0802) [29]; the broad peaks suggest a poor crystallinity and almost amorphous nature of the MnO₂ nanosheets [19].
Table 4.2 Specific surface area, pore size and pore volume of samples.

<table>
<thead>
<tr>
<th>Samples</th>
<th>Pore Volume ($cm^3 g^{-1}$)</th>
<th>$S_{BET}$ ($m^2 g^{-1}$)</th>
<th>Average pore size (nm)</th>
<th>Ratio $^a$ (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>KWB</td>
<td>0.775</td>
<td>1433.06</td>
<td>3.03</td>
<td>46.1</td>
</tr>
<tr>
<td>KWBM-1</td>
<td>0.700</td>
<td>1132.53</td>
<td>4.49</td>
<td>38.8</td>
</tr>
<tr>
<td>KWBM-2</td>
<td>0.487</td>
<td>769.82</td>
<td>5.56</td>
<td>33.9</td>
</tr>
<tr>
<td>KWBM-3</td>
<td>0.398</td>
<td>601.17</td>
<td>6.01</td>
<td>29.3</td>
</tr>
<tr>
<td>KWBM-4</td>
<td>0.331</td>
<td>471.33</td>
<td>6.23</td>
<td>28.8</td>
</tr>
</tbody>
</table>

$^a$ Pore diameter less than 5 nm volume to total pore volume ratio.

As we all know that the surface area and the pore-size distribution are the two key factors for the electrochemical performance of electrode materials. The specific surface area and pore-size distribution of KWB and KWBM were characterized by nitrogen adsorption/desorption measurements (as shown in Fig. 4.6a and b). It is clear that the KWB shows the typical combined I/IV type sorption isotherms, indicating the poly porous structure [30]. The specific surface area of KWB is up to 1433.06 m² g⁻¹ (as shown in Table 4.2), which results KWB suit for acting as the scaffolds to combine with the active materials of MnO₂. For KWBM composites, the MnO₂ nanosheets homogenously anchor on the surface of KWB and will block the mesopores and micropores of KWB. Presumably, the specific surface area of KWBM will be lower
than pure KWB, and the pore distribution will be changed.

Moreover, the specific surface area and pore distribution of KWBM can be controlled by adjust the mass ratio of KMnO₄/KWB. As shown in Table 4.2, the specific surface area and pore volume of KWBM was significantly decreased with the mass ratio of KMnO₄/KWB increased. On the contrary, the average pore size of KWBM was significantly increased with the mass ratio of KMnO₄/KWB increased. In addition, the volume ratio (pore diameter less than 5 nm volume to total pore volume ratio) of KWBM gradually decreased from 46.1% (volume ratio of KWB) with mass ratio of KMnO₄/KWB increased, which indicating that the micropores and mesopores structure of KWB gradually disappeared. However, the volume ratio and specific surface area of the KWBM-4 can still reach up to 28.8% and 471.33 m² g⁻¹, respectively. These acceptable pore structure and specific surface area of KWBM-4 have an enough promise to be electrode material for supercapacitors application.

The information of the surface chemical composition of samples was obtained from

Fig. 4.7 XPS survey spectra of KWB biocarbon, KWBM-1, KWBM-2, KWBM-3 and KWBM-4 composites (a); high resolution C1s (b) and O1s (c) XPS spectra of KWB biocarbon; high resolution Mn2p (d), C1s (e) and O1s (f) XPS spectra of KWBM-4 composite.
XPS spectra. The full scan spectra of samples are shown in Fig. 4.7a. Compared with KWB, the XPS spectra of KWBM added a signal of Mn element. For KWBM composites, the content of C and O gradually decreased and increased, respectively, with the mass ratio of K\text{MnO}_4/KWB increased. Moreover, the ratio of Mn and C content were also increased with the mass ratio of K\text{MnO}_4/KWB increased. These phenomena were much accorded with EDX analysis. For Mn 2p spectra of KWBM-4 (as shown in Fig. 4.7d), two peaks located at 654 and 642.4 eV were attributed to the Mn 2p\textsubscript{1/2} and Mn 2p\textsubscript{3/2} with a spin-energy separation of 11.6 eV, confirmed the formation of \text{MnO}_2 [31].

<table>
<thead>
<tr>
<th>Samples</th>
<th>C/at. %</th>
<th>O/at. %</th>
<th>Mn/at. %</th>
<th>Mn/C (%)</th>
<th>O-Mn (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>KWB</td>
<td>84.76</td>
<td>15.24</td>
<td>--</td>
<td>--</td>
<td>--</td>
</tr>
<tr>
<td>KWBM-1</td>
<td>50.72</td>
<td>26.45</td>
<td>22.83</td>
<td>45.01</td>
<td>56.46</td>
</tr>
<tr>
<td>KWBM-2</td>
<td>46.13</td>
<td>29.47</td>
<td>24.40</td>
<td>52.89</td>
<td>63.02</td>
</tr>
<tr>
<td>KWBM-3</td>
<td>43.43</td>
<td>31.89</td>
<td>24.68</td>
<td>56.83</td>
<td>64.22</td>
</tr>
<tr>
<td>KWBM-4</td>
<td>38.34</td>
<td>38.26</td>
<td>23.40</td>
<td>61.03</td>
<td>69.37</td>
</tr>
</tbody>
</table>

Fig. 4.8 High resolution O1s XPS spectra of KWBM-1 (a), KWBM-2 (b) and KWBM-3 (c) composites.

More information about the change of surface chemistry has been obtained by the deconvoluted high-resolution XPS spectra of two regions (C and O, as shown in Fig. 4.7). The C1s spectra of KWB and KWBM-4 in Fig. 4.7b and e can be deconvoluted into three components. The peaks of 284.4, 285.8 and 288.8 eV can be assigned to aromatic or other sp2-hydridised carbon atoms bound to neighboring carbon atoms (C-C, 284.4 eV), carbon in C-O (285.8 eV) and C=O (288.8 eV), respectively [32]. The
O1s spectra of KWB could be deconvoluted into three peaks (as shown in Fig. 4.7c) with binding energies of 531.5, 533.2 and 534.8 eV. These peaks could be assigned to –OH/C-O, C=O and C-O-H groups, respectively [33]. Compared with KWB, except the above three peaks in O1s spectra of KWBM-4 (as shown in Fig. 4.7f), the other new peak appeared in 529.8 eV, which can be attributed to oxygen bonded with manganese (O-Mn) in MnO2 crystal lattice [34]. The content of oxygen atom in O-Mn of KWBM (which obtained by O1s spectra of KWBM-1, 2, 3 and 4, as shown in Fig. 4.7f and 4.8) is also listed in Table 4.3. It is easy to find that the content of O-Mn was significantly increased with the mass ratio of KMnO4/KWB increased. These transformation further evidences that the element composition of KWBM can be controlled by adjusting the mass ratio of KMnO4/KWB, and the mass ratio has a meaningful impact on electrochemical performance of KWBM composites.

4.3.2 Electrochemical performance of KWBM composites

To explore potential applications of the KWBM composites to supercapacitors, we first performed the CV, GCD and EIS measurements in a three-electrode configuration. Fig. 4.9 shows the CV curves of KWBM composites at a scan rate of 0.01 V s⁻¹.

![Fig. 4.9 CV curves of KWBM composites at scan rate of 0.01 V s⁻¹](image-url)
Different from CV curve of KWB (as shown in Fig. 3.11), all the CV curves of KWBM deviate from the rectangular shape, due to the charge storage process of MnO$_2$-based electrode. Some researchers have proposed that the process of charge storage was ascribed to the rapid intercalation of alkali metal cations such as K$^+$ intercalation into the electrode during reduction and deintercalation upon oxidation [35, 36], as shown in reaction (Re. 2):

\[
\text{MnO}_2 + \text{M}^+ + \text{e}^- \leftrightarrow \text{MnO}_\text{M}
\]  

(Re. 2)

where M$^+$ = Na$^+$, K$^+$, and Li$^+$. It should be noted that the process involved a redox reaction, and which can significantly improve the capacitive performance. Furthermore, KWBM-4 electrode exhibited a much larger current response than that of the other electrode, and presented an excellent electrochemical performance, attributing to the acceptable microstructure and element composition.

The electrochemical performance of KWBM was further analyzed by the GCD method and the results are shown in Fig. 4.10. All the charge curves of the KWBM were almost symmetrical to their corresponding discharge curves with slight deformation, indicating that the capacitance is in account of the combination of
electrical double-layer capacitance (EDLC) and pseudocapacitance corresponds to Re. 2. Moreover, the discharge time of KWBM-4 is longer than that of other samples, indicating that the KWBM-4 possesses the more excellent capacitive performance than the other samples. The specific capacitance (SC) of KWBM could be got by Eq. 1. The SC of KWBM-1, 2, 3 and 4 are 260, 346, 360 and 402 F g\(^{-1}\), respectively, at a current density of 1 A g\(^{-1}\). The SC of KWBM increases with the increase of the KMnO\(_4\)/KWB mass ratio. These results are further evidences for the inference form CV curves.

![Fig. 4.11 EIS curves of KWBM composites.](image)

The EIS measurements were performed in order to further explore the charge transport and ion diffusion between KWBM electrode and electrolyte. The Nyquist plots of KWBM are showed in Fig. 4.11. All samples present similar shapes, and are composed of three parts: a semi-circular line in the high frequency region (as shown in the inset of Fig. 4.11), a nearly 45\(^{\circ}\) diagonal line in the medium frequency region and a nearly vertical line in the low frequency region. These three parts correspond to charge transfer resistance, diffusion resistance and capacitive behavior, respectively [37]. From the inset, is easily found that the KWBM-4 exhibits the smaller charge transfer resistance than the other samples due to the fact that it has the least diameter of arc.
Furthermore, KWBM-4 shows the shortest 45° diagonal line and the largest slope of the straight line at the medium and low frequency region, respectively, indicating that KWBM-4 also possesses the lowest different resistance and best excellent capacitive performance than those of the other samples. In addition, the very high-frequency intercept at the real axis at the beginning of the arc represents the internal resistance (Rs), including the intrinsic resistance of the sample, the contact resistance between the sample and current collector, and the resistance of bulk electrolyte [38]. The inset shows that the Rs of KWBM-1, 2, 3 and 4 are 1.48, 1.39, 1.40 and 1.35 Ω, respectively. These above results indicate that KWBM-4 possesses an outstanding capacitive performance, and this conclusion agrees with the analyses of CV and GCD measurements.

![Graph showing specific capacitance versus current density for KWBM composites.](image)

**Fig. 4.12** Specific capacitance versus current density of KWBM composite

Fig. 4.12 shows the SC as a function of current density for KWBM composites. In general, the SC of all samples decreases with the increase of current density, this is may be due to the contribution of MnO₂ pseudocapacitance to total capacitance decrease with the current density increase. Moreover, it is noteworthy that the SC downtrend of KWBM-4 is larger than that of the other samples. This is mainly due to the specific surface area of KWBM-4 is smaller than that of the other samples, resulting in that the
contact area of KWBM-4 and electrolyte is smallest among all samples. However, the SC of KWBM-4 could still reach to 203 F g⁻¹ at the current density of 5 A g⁻¹, and this value is still larger than that of the other samples. This result suggests that the KWBM-4 possesses the fast charge-discharge performance, which is essential for supercapacitors application.

Fig. 4.13 CV curves of KWB and KWBM-4 in three-electrode asymmetric system at a scan rate of 0.01 V s⁻¹.

Fig. 4.14 CV (a, scan rate of 0.01 V s⁻¹) and GCD (b, current density of 1 A g⁻¹) curves of two-electrode asymmetric system (using KWBM-4 and KWB as the positive and negative electrode, respectively) at different potential windows.
CV curves of KWBM-4//KWB measured at scan rate of 0.01 V s⁻¹ exhibit rectangular-like shapes (as shown in Fig. 4.14a) even at the potential window up to 1.5 V, demonstrating KWBM-4//KWB present an ideal capacitive behavior. Moreover, a pair of weak redox peaks appears in all CV curves, suggesting that the pseudocapacitance and EDLC capacitance are simultaneously exist. In addition, there is no significant increase of anodic current at potential window of 0-1.5 V, which implies that the electrolyte is not decomposed, because that the storage of nascent hydrogen on the electrode is below the thermodynamic potential for water decomposition [40]. Fig. 4.14b shows the GCD curves of KWBM-4//KWB at current density of 2 A g⁻¹ under different potential windows. Similar to CV curves, all GCD curves show the equilateral triangle-like shape even at the potential window up to 1.5 V, which still implies the ideal capacitive behavior of KWBM-4//KWB. The SC of KWBM-4//KWB from these GCD curves can be obtained by Eq. 1. The SC is 63.4, 65.1, 66.5, 73.3 and 76.5 F g⁻¹ at 0-0.8, 0-1.0, 0-1.2, 0-1.4 and 0-1.5 V, respectively. Therefore, these above results confirm that the best potential window of KWBM-4//KWB is 0-1.5 V.

Fig. 4.15 CV (a) and GCD (b) curves of asymmetric system measured at different scan rates and current densities in a potential window of 1.5 V, respectively (using KWBM-4 and KWB as the positive and negative electrode, respectively).

Fig. 4.15a shows the CV curves of the KWBM-4//KWB measured at various scan rates of 0.01-0.1 V s⁻¹ during 0 and 1.5 V. The CV curves exhibit a nearly rectangular-
like shape, which also indicate the ideal capacitive behavior. The rectangular CV shape remains very well even at a high scan rate of 0.1 V s$^{-1}$, indicating KWBM-4//KWB possesses outstanding rate capability and meets high-power supercapacitors. GCD measurements were made at different current densities to further evaluate the capacitive performance of the KWBM-4//KWB at the potential window of 0-1.5 V, as shown in Fig. 4.15b. These GCD curves are highly linear and symmetrical, indicating a rapid response of current and voltage, and an excellent electrochemical reversibility [9]. The SC of KWBM-4//KWB is 82.9 F g$^{-1}$ at a current density of 1 A g$^{-1}$, and still remains at 51.3 F g$^{-1}$ when the current density increases to 10 A g$^{-1}$, further suggesting that the KWBM-4//KWB has excellent capacitive performance.

\[ \begin{align*}
E &= \frac{1}{2} C_m (\Delta V)^2 \quad (\text{Eq. 2}) \\
P &= \frac{E}{t} \quad (\text{Eq. 3})
\end{align*} \]

where $C_m$, $t$ and $\Delta V$ are the SC, discharge time (h) and potential window (V) of

Fig. 4.16 Ragone plot of asymmetric system.

Ragone plot, a performance indicator of power density against energy density, was drawn to characterize the electrochemical performance of KWBM-4//KWB [37]. The energy density ($E$, Wh kg$^{-1}$) and power density ($P$, W kg$^{-1}$) were calculated from the discharge curves according to the following equations [41]:
KWBM-4//KWB, respectively. Fig. 4.16 shows the ragone plot of KWBM-4//KWB. It is obviously that the KWBM-4//KWB shows the outstanding performance with 25.9 Wh kg\(^{-1}\) of the energy density at a power density of 750 W kg\(^{-1}\). The exciting performance is comparable or superior to those of MnO\(_2\)-based asymmetric two-electrode system reported in literature (Table 4.4).

Table 4.4 Performance of MnO\(_2\)-based electrodes in asymmetric two-electrode system.

<table>
<thead>
<tr>
<th>Positive electrode</th>
<th>Negative electrode</th>
<th>Cell voltage (V)</th>
<th>Energy density (Wh kg(^{-1}))</th>
<th>Power density</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>rGO/MnO(_2)/CB</td>
<td>rGO/CB</td>
<td>1.8</td>
<td>24.3</td>
<td>45 kW kg(^{-1})</td>
<td>[42]</td>
</tr>
<tr>
<td>MnO(_2)/CNF</td>
<td>Bi(_2)O(_3)/CNF</td>
<td>1.8</td>
<td>11.3</td>
<td>352.6 W kg(^{-1})</td>
<td>[43]</td>
</tr>
<tr>
<td>MnO(_2)/CNTs</td>
<td>AC</td>
<td>1.5</td>
<td>13.3</td>
<td>600 W kg(^{-1})</td>
<td>[44]</td>
</tr>
<tr>
<td>MnO(_2)</td>
<td>AC</td>
<td>1.5</td>
<td>7.4</td>
<td>400 W kg(^{-1})</td>
<td>[45]</td>
</tr>
<tr>
<td>NaMnO(_2)</td>
<td>AC</td>
<td>1.8</td>
<td>19.5</td>
<td>130 W kg(^{-1})</td>
<td>[46]</td>
</tr>
<tr>
<td>MnO(_2)/MnCO(_3)/rGO</td>
<td>rGO</td>
<td>1.8</td>
<td>17.8</td>
<td>400 W kg(^{-1})</td>
<td>[16]</td>
</tr>
<tr>
<td>MnO(_2)-nanorods</td>
<td>rGO</td>
<td>1.6</td>
<td>14.31</td>
<td>5.1 kW kg(^{-1})</td>
<td>[47]</td>
</tr>
<tr>
<td>MnO(_2)-nanowires</td>
<td>rGO</td>
<td>1.4</td>
<td>19.97</td>
<td>6.2 kW kg(^{-1})</td>
<td>[47]</td>
</tr>
<tr>
<td>MnO(_2)-nanoneedles</td>
<td>rGO</td>
<td>1.8</td>
<td>24.12</td>
<td>7.8 kW kg(^{-1})</td>
<td>[47]</td>
</tr>
<tr>
<td>MnO(_2)</td>
<td>PEDOT</td>
<td>1.8</td>
<td>13.5</td>
<td>120.1 kW kg(^{-1})</td>
<td>[48]</td>
</tr>
<tr>
<td>KWBM-4</td>
<td>KWB</td>
<td>1.5</td>
<td>25.9</td>
<td>750 W kg(^{-1})</td>
<td>This work</td>
</tr>
</tbody>
</table>

Fig. 4.17 Cycle stability of asymmetric system
The long-term cycling stability is an important criterion for supercapacitors applications. The cycle stability measurement 2000 cycles for KWBM-4//KWB was conducted using GCD test at 6 A g\(^{-1}\) and the results are shown in Fig. 4.17. The KWBM-4//KWB exhibits excellent cycle stability with 94.2 % SC retention after 2000 cycles. The inset in Fig. 4.17 represents GCD curves of the first three and last three cycles. The last three cycles remain the symmetric triangular shape, indicating KWBM-4//KWB still keep the ideal capacitive performance. Furthermore, cycling performance of KWBM-4/KWB is comparable to those of other MnO\(_2\)-based asymmetric two-electrode system reported in literature (Table 4.5).

<table>
<thead>
<tr>
<th>Positive electrode</th>
<th>Negative electrode</th>
<th>Cell voltage</th>
<th>Capacitance retention after cycling</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>MnO(_2)</td>
<td>CNT-CNf</td>
<td>2.0 V</td>
<td>92% after 2000 cycles</td>
<td>[11]</td>
</tr>
<tr>
<td>MnO(_2)/MWNT</td>
<td>MWNTs</td>
<td>2.45 V</td>
<td>72% after 300 cycles</td>
<td>[49]</td>
</tr>
<tr>
<td>MnO(_2)</td>
<td>FeOOH</td>
<td>1.85 V</td>
<td>85% after 300 cycles</td>
<td>[50]</td>
</tr>
<tr>
<td>CNTs/MnO(_2)</td>
<td>CNTs/SnO(_2)</td>
<td>1.7 V</td>
<td>92% after 1000 cycles</td>
<td>[51]</td>
</tr>
<tr>
<td>KWBM-4</td>
<td>KWB</td>
<td>1.5 V</td>
<td>94.2% after 2000 cycles</td>
<td>This work</td>
</tr>
</tbody>
</table>

### 4.4 Conclusions

In this work, we successful prepared the allium-giganteum-like biocarbon—KWB, which derived from the common biomass of sugarcane bagasse using one-step facile carbonization and activation method. The interconnected frameworks of KWB offered a fine platform for anchoring MnO\(_2\) nanosheets. Subsequently, MnO\(_2\) nanosheets were uniformly distributed on the surface of KWB by in-suit redox deposition and obtained the KWBM composites. The electrochemical performance of KWBM was closely related to the mass ratio of KMnO\(_4\)/KWB. KWBM-4 showed the best electrochemical performance when the mass ratio of KMnO\(_4\)/KWB was 2 in three-electrode system. When employing the KWBM-4 as the positive electrode and the KWB as the negative
electrode, the assembled asymmetric two-electrode system exhibited a remarkable energy density of 25.9 Wh kg\(^{-1}\), excellent rate capability and high cycle stability (SC retention is 94.2 % after 2000 cycles). Therefore, it is a promising strategy for employing low-cost and eco-friendly biomaterials to prepare advanced electrode materials for energy storage devices.
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Chapter 5 Polyaniline/sugarcane bagasse derived biocarbon composites with superior performance in supercapacitors

5.1 Introduction

Supercapacitors are one of the most promising energy storage devices due to their superior power density, fast charge/discharge rate, and long cycle life [1-4]. According to the energy storage mechanism, supercapacitors can be classified as pseudocapacitors and electrical double-layer capacitors (EDLC) [5, 6]. Polyaniline is a typical electrode material of pseudocapacitors, and it possesses many advantages, such as: ease of synthesis, high conductivity, highly reversibility and large storage capacity [7-9]. Unfortunately, its electrochemical performance tends to fade for swelling and shrinking during cycling (the insertion/deinsertion process of counter ions), which hinder its further application in supercapacitors [10].

To respond the above problems, some research focused on preparation of PANI/carbon composites. Until now, many kinds of carbon materials, such as carbon nanotube [11], graphene [12] and carbon nanofibers [13] are often used as the content of PANI/carbon composites because their superior electronic conductivity, high specific surface, special microstructure, reasonable chemical stability and excellent thermal stability [14]. However, the preparation processes of these carbon materials are complex, and these carbon materials usually need a pretreatment process for preparation of PANI/carbon composites, leading to high cost [15-17].

Therefore, simple and low cost preparation method of carbon materials such as direct pyrolysis of biomass or biowaste entered the researcher’s vision [18-21]. In recent years, many biocarbon materials have been prepared from biomass such as fallen leave [22], ramie [23], coffee endocarp [24] and cassava peel [25]. The above biocarbon materials present high specific surface area and abundant surface oxygen functional groups,
which makes them suitable for preparation of PANI/carbon composites. Especially, we found an Allium-giganteum-like microstructure in sugarcane bagasse derived biocarbon, it not only has the above advantages, but also presents an Allium-giganteum-like microstructure [26]. This particular microstructure is possibly beneficial to ion transport if this biocarbon is applied to supercapacitors.

In this paper, we report an efficient method for preparation of PANI/biocarbon composites (PANI/KWB) which are going to be as electrode materials for supercapacitors. The biocarbon (KWB) was obtained by direct pyrolysis of sugarcane bagasse, and then PANI/KWB was obtained by in-site polymerization method under different mass ratios of An/KWB. During the polymerization process, [CMMIm]Cl ionic liquid (IL) was chosen as dopant is due to its proton acid and surfactant features [27, 28]. We found that chemical composition and microstructure of PANI/KWB were closely related to the mass ratio of aniline monomer (An) and KWB, and which further impacted electrochemical performance of PANI/KWB profoundly. In addition, an asymmetric two-electrode system was assembled by employing PANI/KWB and KWB as positive and negative electrode. We also found that the asymmetric system possessed the high energy density and high cycle stability. The results indicate a promising candidate for supercapacitors.

5.2 Experimental

5.2.1 Materials

Sugarcane bagasse was crushed by family-use grinder before use. Aniline monomer (An) was purified by distillation under reduced pressure and stored at 4 °C in dark. The ionic liquid (IL), 1-methyl-3-alkylcarboxylic acid imidazolium chloride ([CMMIm]Cl), was synthesized in our laboratory and the synthesis method described in supporting information. The other reagents, such as ethanol, potassium hydroxide (KOH), hydrochloric acid (HCl), and ammonium persulfate (APS), were all purchased from Nacalai Tesque, Inc. (Kyoto, Japan) and of analytical grade, being used without further
purification.

5.2.2 Synthesis of 1-methyl-3-alkylcarboxylic acid imidazolium chloride ([CMMIm]Cl) ionic liquid

The [CMMIm]Cl was prepared as Fei reported. [27] In a typical fabrication of [CMMIm]Cl, 0.17 mol chloroacetic acid solid was added spoonful to 0.15 mol N-Methylimidazole liquid with stirring in 1 h at 15-20 °C. When the mixed solution uniform, put it into water bath kettle to heat up to 70 °C and run the reaction for 3-5 h. After that, the solution was used for suction filtration and washed the material with 30 ml of methyl cyanide for twice. Finally, the product was dried in vacuum at 40 °C for 12 h to obtain the white solid of [CMMIm]Cl.

5.2.3 Preparation of KOH-activated biocarbon (KWB) derived from sugarcane bagasse

KWB was obtained by KOH activation method according to the previous reported work [26] and the typical process as follows: first, sugarcane bagasse was cut into small pieces (opening size approximately 2 mm) and immersed in distilled water for 8 h at 90 °C, and then dried at 80 °C for 24 h. Second, 1 g clean sugarcane bagasse and 0.6 g KOH were poured into 12 ml absolute ethanol with stirring and heated at 60 °C until all of the ethanol was evaporated, and then dried at 70 °C for 12 h. Third, the remnant mixture was pyrolyzed in a tubular furnace in N\textsubscript{2} atmosphere at 800 °C (heating rate is 10 °C min\textsuperscript{-1}) for 2 h. Finally, the obtained carbon material was washed by 1 M HCl solution and deionized water till the filtrate became neutral and collected after vacuum dry.

5.2.4 Preparation of polyaniline/ activated biocarbon composites (PANI/KWB)

The PANI/KWB composites were prepared by in-site polymerization method. 5 g [CMMIm]Cl IL dissolved in 100 ml deionized water. Then add 1 ml An monomer and
0.1 g KWB (the mass ratio of An/KWB is 10:1) to the above solution. Subsequently, the mixture was sonicated for 0.5 h, and the APS solution (1.5 g APS dissolved in 10 ml of hydrochloric acid solution (1 M)) was added drop by drop to the mixture to start the polymerization. The reaction was performed with stirring at room temperature for 2 h. Finally, the product was removed from solution, washed with distilled water until pH=6-7, and dried in vacuum at 80 °C for 12 h to obtained the PANI/KWB-1 as black green powder. The other products were denoted as PANI/KWB-2, PANI/KWB-3, PANI/KWB-4 and PANI/KWB-5 for the mass ratios of An/KWB was 20:1, 30:1, 40:1 and 50:1 (the mass ratio of An monomer to ILs and APS was kept at 1:5:1.5), respectively. In addition, we also prepared pure PANI as control.

5.2.5 Synthesis of pure PANI without IL

For composite with the PANI via [CMMIm]Cl brønsted acidic IL as dopant, the pure were prepared by the same method while used sulfamic acid (SA) as the dopant. Typically, 5 g SA dissolved in 100 ml deionized water, then 1 ml An monomer added the above solution. Subsequently, the mixture was ultrasonication for 0.5 h, and the APS solution (1.5 g APS dissolved in 10 ml of hydrochloric acid solution (1 M)) was by drop to the mixture to start the polymerization. The reaction was performed with stirring at room temperature for 2 h. Finally, the product was removed from solution, washed with distilled water until pH=6-7, and dried in vacuum at 80 °C for 12 h to obtained the pure PANI as black green powder.

5.2.6 Structure characterization

The morphology of samples were observed by transmission electron microscopy (TEM, H-8110, Hitachi Co., Ltd. Tokyo, Japan) and scanning electron microscopy (SEM, S-4300, Hitachi Co., Ltd. Tokyo, Japan). Nitrogen adsorption-desorption isotherms were measured at -196 °C on a Micromeritics ASAP 2020 apparatus. The specific surface area was calculated according to the Brunauer-Emmet-Teller (BET) equation. All samples were outgassed under vacuum at 80 °C overnight prior to
measurement. The chemical structures of the samples were confirmed by Fourier transform infrared spectroscopy (FTIR, IRT-7000, JASCO, Japan). The crystallographic structure of the samples was observed by X-ray diffraction (XRD, XRD-6000, Shimadzu Co., Ltd. Kyoto, Japan) equipped with Cu Kα radiation. X-ray photoelectron spectroscopy (XPS) analysis was performed on the PHI Quantum 5000 equipped with an Al Kα radiation source.

5.2.7 Electrochemical test

The electrochemical tests were measured with an electrochemical workstation (CHI 660D, Shanghai Chenhua, China) in the two or three-electrode system in 1 M Na₂SO₄ electrolyte. For three-electrode system, saturated calomel electrolyte and platinum-wire electrode were used as reference and counter electrode, respectively. The test electrodes were first prepared by mixing PANI/KWB composites with carbon black (CB) and PTFE at a mass ratio of 80:15:5. After that, the mixing were grinding adequately to obtained a homogeneous paste, that was coated onto the Ni foam for current collector and dried under vacuum at 80 °C for 24 h, and then compressed at 10 MPa for 5 min. The mass loading of active material in an electrode is about 8 mg. Cyclic voltammetry (CV), and galvanostatic charge/discharge (GCD) were conducted in a potential window from -0.8 to 0.2 V for AB, and -0.2 to 0.8 V for PANI/KWB composites, respectively. Electrochemical impedance spectroscopy (EIS) was carried out in the frequency range from 10⁵ to 10⁻² Hz at open circuit potential with as ac perturbation of 5 mV. The asymmetric two-electrode system was assembled with two almost identical working electrodes (the active material was PANI/KWB composites sample, and the counter electrode was KWB sample). The CV and GCD tests were run at different voltage windows.

The standard calomel reference electrode (SCE) based on GCD curves was evaluated according to the following equation (Eq. 1):

\[ C_m = \frac{I \times t}{(\Delta V \times m)} \]  

(Eq. 1)

where I is discharge current (A), t is the discharge time (s), \( \Delta V \) is the voltage window
(V), and m is the mass of active material on working electrode in three-electrode system, in two-electrode system is the total mass of active material on two working electrodes [29].

The energy density (E, Wh kg\(^{-1}\)) and power density (P, W kg\(^{-1}\)) of the asymmetric two-electrode system were calculated from the discharge curves according to the following equations [30]:

\[
E = \frac{1}{2} C_m (\Delta V)^2 \quad \text{(Eq. 2)}
\]

\[
P = \frac{E}{t} \quad \text{(Eq. 3)}
\]

where \(C_m\), \(t\), and \(\Delta V\) are the specific capacitance (F g\(^{-1}\)), discharge time (h) and potential window (V) form GCD curves in two-electrode system, respectively.

5.3 Results and discussion

5.3.1 Microstructural characterization

![SEM images of PANI (a), PANI/KWB-1 (b), PANI/KWB-2 (c), PANI/KWB-3 (d), PANI/KWB-4 (e) and PANI/KWB-5 (f) composites.](image)

Fig. 5.1 SEM images of PANI (a), PANI/KWB-1 (b), PANI/KWB-2 (c), PANI/KWB-3 (d), PANI/KWB-4 (e) and PANI/KWB-5 (f) composites.

The morphology and microstructure of the prepared samples are shown in Fig. 5.1. As shown in Fig. 5.1a, PANI nanoparticles exhibit a coral-like structure and with a large number of conical protrusions (as shown in Fig. 5.2c and d). Compared with pristine KWB, the clean surface and interconnected frameworks of KWB (as shown in Fig. 5.2a
and b) were gradually covered by PANI nanoparticles with the mass ratios of An/KWB increased. It is noteworthy that the surface and frameworks of KWB were completely covered by PANI nanoparticles (as shown in Fig. 5.1e) when the mass ratio of An/KWB is 40:1.

![Fig. 5.2 SEM images of AB (a-b) and PANI (c-d)](image)

For compare with PANI/KWB composites, the SEM of KWB and PANI shown in Fig. 5.2. The morphology of KWB is hierarchical porous structure (as shown in Fig. 5.2a and b). It is clearly observed that there are a large number of nano-scale pores on the inner wall of the micron-sized macropores, and the surface of KWB is smooth.

The morphology of PANI/KWB-4 was further investigated by TEM. It can be clearly seen that the surface of KWB is coated by PANI as shown in Fig. 5.3a. Moreover, the conical projection corresponding to SEM image of PANI was also observed (as shown in Fig. 5.3b). In addition, for PANI/KWB-5 (Fig. 5.1f), the porous structure of KWB completely disappeared, which means that the porous structure of KWB was filled by PANI nanoparticles. The above phenomena may impact electrochemical performance of PANI/KWB composites profoundly.
Fig. 5.3 TEM images of PANI/KWB-4 composites

Fig. 5.4 X-ray diffraction patterns of the prepared KWB, PANI and PANI/KWB composites.

The crystallinity of the samples was examined by X-ray diffraction analysis. Fig. 5.4 shows the XRD patterns of KWB, PANI and PANI/KWB composites. For KWB, a broad diffraction peak centered at 24.5° correspond to the (002) plane of graphite, arising from the amorphous nature of graphite [31]. Furthermore, a weak diffraction peak can be observed at 43.4° corresponding to graphite (100), which means of the complete transformation of sugarcane bagasse to another kind of activated carbon [32]. For pure PANI, three broad diffraction peaks at 15, 20.4 and 25° were correspond to the
(010), (100), and (110) plane of PANI, respectively [33], and these peaks suggest a poor crystallinity and partially crystalline of the PANI [34]. The characteristic peaks of PANI/KWB are exactly the same as those of pure PANI, indicating that PANI nanoparticles covered on the surface of KWB.

Fig. 5.5 FT-IR spectra of the prepared pure PANI and [CMMIm]Cl (a), and AB biocarbon, PANI and PANI/KWB (b) composites.

For [CMMIm]Cl sample (as shown in Fig. 5.5a), the peaks at 3090 cm\(^{-1}\) is attributed to the overlap peaks of the V=C-H stretching of the imidazole ring and the hydroxyl groups of carboxylic acid. The peaks at 2861 cm\(^{-1}\) correspond to the saturation C-H stretching mode. The peaks at 2580, 2480 cm\(^{-1}\) are attributed to the characteristic C-N and C=N stretching of the imidazole rings. The peak at 1730, 1650 and 1580 cm\(^{-1}\) are the characteristic of carbanyl group, the C-N and C=N of the imidazole rings, respectively. Band located around 1400 cm\(^{-1}\) are C=C and C-C stretching. The peak at 1190 cm\(^{-1}\) is the C-O stretching. The peak at 1090 cm\(^{-1}\) is attributed to O-H out-of-plane bending. In addition, Bands around 978 and 773 cm\(^{-1}\) are attributed to C–H in-plane and out-of-plane bending [27]. For pure PANI sample (as shown in Fig. 5.5a), the absorption band at 3440 cm\(^{-1}\) is the associating hydroxyl groups. The bands at 1580 and 1490 cm\(^{-1}\) are attributed to the characteristic N=Q=N stretching of the quinoid, N-B-N of the benzenoid, and band centered at 1300 cm\(^{-1}\) correspond to N-H bending and asymmetric C-N stretching modes of the benzenoid ring. Bands around 1130 and 798
cm$^{-1}$ are attributed to C–H in-plane and out-of-plane bending of aniline rings in protonation [36].

The chemical structure of KWB, PANI and PANI/KWB composites were characterized by FT-IR spectra (as shown in Fig. 5.5b). In the spectra of KWB, the peak at 1099 cm$^{-1}$ was for C–O group, and this group formed during the bagasse pyrolysis and activation process [35]. In the spectra of PANI, the peak at 1115 cm$^{-1}$ assigned to the C–H in-plane stretching of the quinoid. Other peaks are assigned as follows: saturation C–H stretching peak at 2921 cm$^{-1}$, C–N and C=N stretching of the imidazole ring peak at 2358 cm$^{-1}$, N=Q=N stretching of the quinoid peak at 1567 cm$^{-1}$, N–H bending of the benzenoid ring peak at 1261 cm$^{-1}$, and C–H in-plane bending of aniline ring peak at 796 cm$^{-1}$ [36, 37]. Once KWB compositied with PANI, the feature peak of KWB will overlap with the peak of PANI (peak at 1115 cm$^{-1}$), and the other feature peak of PANI will still existed. It further suggests that PANI nanoparticles covered on the surface of KWB.

![Fig. 5.6 N$_2$ adsorption-desorption isotherms (a) and pore size distribution by BJH method (b) of the prepared PANI and PANI/KWB-1, PANI/KWB-2, PANI/KWB-3, PANI/KWB-4 and PANI/KWB-5 composites.](image)

The nitrogen adsorption/desorption isotherm for the PANI and PANI/KWB composites and their corresponding pore size distribution are shown in Fig. 5.6. For PANI, its isotherm can be classified as a type III isotherm with a type H3 hysteresis,
suggesting the non-porous structure [38]. The KWB display type I isotherms with a type H2 hysteresis (as shown in Fig. 5.7a) and possess a particularly high specific surface area (1433.06 m² g⁻¹, as shown in Table 5.1), indicating the poly porous structure [39]. For PANI/KWB composites, the PANI nanoparticles covered on the surface of KWB and will block the porous structure of KWB. Possibly, the \( S_{\text{BET}} \) of PANI/KWB will be lower than KWB, and the \( P \) will be changed.

![Fig. 5.7 N\(_2\) adsorption-desorption isotherms (a) and pore size distribution by BJH method (b) of the prepared KWB.](image)

Table 5.1 Specific surface area, pore size of samples. (\( S_{\text{BET}} \) - the specific surface area; \( S_{\text{mi}} \) - the micropore specific surface area; \( P \) - the main pore size; \( V_t \) - the total pore volume)

<table>
<thead>
<tr>
<th>Sample</th>
<th>( S_{\text{mi}} ) (m² g⁻¹)</th>
<th>( S_{\text{BET}} ) (m² g⁻¹)</th>
<th>( V_t ) (cm³ g⁻¹)</th>
<th>( P ) (nm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>KWB</td>
<td>634.88</td>
<td>1433.06</td>
<td>0.775</td>
<td>3.03</td>
</tr>
<tr>
<td>PANI</td>
<td>--</td>
<td>22.58</td>
<td>0.093</td>
<td>20.58</td>
</tr>
<tr>
<td>PANI/KWB-1</td>
<td>5.10</td>
<td>36.82</td>
<td>0.130</td>
<td>23.94</td>
</tr>
<tr>
<td>PANI/KWB-2</td>
<td>--</td>
<td>25.659</td>
<td>0.109</td>
<td>19.03</td>
</tr>
<tr>
<td>PANI/KWB-3</td>
<td>--</td>
<td>33.58</td>
<td>0.123</td>
<td>19.6</td>
</tr>
<tr>
<td>PANI/KWB-4</td>
<td>--</td>
<td>54.47</td>
<td>0.234</td>
<td>18.58</td>
</tr>
<tr>
<td>PANI/KWB-5</td>
<td>--</td>
<td>28.57</td>
<td>0.117</td>
<td>19.32</td>
</tr>
</tbody>
</table>

The \( S_{\text{BET}} \) and \( P \) of PANI/KWB can be influences by the mass ratio of An/KWB. As shown in Table 1, the \( S_{\text{BET}} \) of PANI/KWB was increased first and then decreased with
the mass ratio of An/KWB increased. On the contrary, the $P$ of PANI/KWB was decreased first and then increased with the mass ratio of An/KWB increased. When the mass ratio of An/KWB reached to 40:1 (PANI/KWB-4), the $S_{BET}$ of PANI/KWB showed the maximum value of 54.47 m$^2$ g$^{-1}$. In addition, the $S_{BET}$ of PANI/KWB is higher than that of PANI and much smaller than that of KWB, which indicating that the polyporous structure of KWB is conducive to improving the $S_{BET}$ of electrode material. Moreover, only PANI/KWB-1 has a $S_{mi}$, which is due to part of surface and porous structure of KWB is covered by PANI nanoparticles when the mass ratio of An/KWB is 10. Soon afterwards, the surface and porous structure of KWB is completely covered by PANI nanoparticles with the increase of the mass ratio of An/KWB, which has been proved by SEM images. However, the $S_{BET}$ of KWB will be close to pure PANI and the support function of KWB will lose when the mass ratio of An/KWB is too large (more than 40), which is owing to KWB buried by a large number of PANI nanoparticles. Nevertheless, the $S_{BET}$ of PANI/KWB-4 can still reach up to 54.47 m$^2$ g$^{-1}$ (much higher than that of pure PANI). This acceptable $S_{BET}$ of PANI/KWB-4 has an enough promise to be electrode material for supercapacitors application.

<table>
<thead>
<tr>
<th>Sample</th>
<th>C/at. %</th>
<th>N/at. %</th>
<th>O/at. %</th>
</tr>
</thead>
<tbody>
<tr>
<td>KWB</td>
<td>84.76</td>
<td>-</td>
<td>15.24</td>
</tr>
<tr>
<td>PANI</td>
<td>81.11</td>
<td>8.28</td>
<td>10.61</td>
</tr>
<tr>
<td>PANI/KWB-1</td>
<td>80.58</td>
<td>8.13</td>
<td>11.28</td>
</tr>
<tr>
<td>PANI/KWB-2</td>
<td>81.24</td>
<td>8.95</td>
<td>9.81</td>
</tr>
<tr>
<td>PANI/KWB-3</td>
<td>80.98</td>
<td>8.75</td>
<td>10.27</td>
</tr>
<tr>
<td>PANI/KWB-4</td>
<td>80.63</td>
<td>8.87</td>
<td>10.50</td>
</tr>
<tr>
<td>PANI/KWB-5</td>
<td>81.08</td>
<td>8.04</td>
<td>10.88</td>
</tr>
</tbody>
</table>

The information of the surface chemical composition of samples was obtained from XPS spectra. The full scan spectra of samples are shown in Fig. 5.7a. Compared with KWB, the XPS spectra of PANI/KWB composites added a signal of N element. In addition, the signal of O element of PANI spectra indicates that the [CMMIIm]$^+$ (contain
-COOH) as the cationic of IL were doped into the molecular chain of PANI.

![Diagram](image)

Fig. 5.8 XPS survey spectra of KWB biocarbon, PANI and PANI/KWB composites (a); high resolution C1s (c) and O1s (e) XPS spectra of KWB biocarbon; high resolution N1s (b), C1s (d) and O1s (f) XPS spectra of PANI/KWB-4 composite.

More information about the change of surface chemistry has been obtained by the deconvoluted high-resolution XPS spectra of three regions (C, O and N, as shown in Fig. 5.8). The C1s spectra of KWB could be deconvoluted into three peaks (as shown...
in Fig. 5.8c) with binding energies of 531.5, 533.2 and 534.8 eV. These peaks could be assigned to aromatic or other sp²-hydridised carbon atoms bound to neighboring carbon atoms (C-C, 284.4 eV), carbon in C-O (285.8 eV) and C=O (288.8 eV), respectively [40]. Compared with KWB, except for the above three peaks in C1s spectra of PANI/KWB-4 (as shown in Fig. 5.8d), the other new peaks appeared in 286.2 and 284.8 eV, can be attributed to C=N+/-C-N+ and C-N/C=N, respectively [41]. The N1s spectra of PANI/KWB-4 (as shown in Fig. 5.8b) could be deconvoluted into three components. The peaks of 401.5, 399.7 and 398.7 eV can be attributed to cationic atoms (-N+-), amine (-NH-) and imine (=N-), respectively [42]. The O1s spectra of KWB and PANI/KWB-4 in Fig. 5.8e and f can be deconvoluted into three components. The peaks of 531.5, 533.2 and 534.8 eV can be assigned to –OH/C-O, C=O and C-O-OH groups, respectively [43]. Obviously, the content of C-O-OH group in PANI/KWB-4 is much higher than that of AB, which is also owing to the [CMMIm]⁺ doped into the molecular chain of PANI. The high content of C-O-OH group in PANI/KWB-4 has a meaningful impact on electrochemical performance of composites.

### 5.3.2 Electrochemical performance of PANI/KWB composites

To explore potential applications of the PANI/KWB composites to supercapacitors, we first performed the CV, GCD and EIS measurements in a three-electrode configuration. Fig. 5.9 shows the CV curves of PANI and PANI/KWB composites at a scan rate of 0.01 V s⁻¹. The CV curve of KWB shows nearly symmetric rectangular shape (as shown in Fig. 3.11), indicating an ideal double-layer capacitor behavior [44]. Different from CV curve of KWB, both the CV curves of PANI and PANI/KWB composites deviate from the rectangular shape, due to the charge storage process of PANI-based material. In the CV curves of PANI/KWB composites, the pairs of peaks P1/P2 (the weak peak of P1 as shown in the dark green circle of Fig. 5.9) are attributed to the redox of PANI molecules (leucoemeraldine and pernigraniline species), and the other pairs of peaks P3/P4 are ascribed to benzoquinone/hydroquinone redox transitions [44]. The PANI/KWB-4 shows the larger area enclosed by the CV curve than that of
the PANI and PANI/KWB, revealing the highest specific capacitance (SC).

![CV curves of PANI and PANI/KWB composites at scan rate of 0.01 V s\(^{-1}\)](image)

**Fig. 5.9** CV curves of PANI and PANI/KWB composites at scan rate of 0.01 V s\(^{-1}\)

![GCD curves (a) at current density of 1 A g\(^{-1}\) and specific capacitance (b) versus current density) of PANI and PANI/KWB composites.](image)

**Fig. 5.10** GCD curves (a, at current density of 1 A g\(^{-1}\)) and specific capacitance (b, versus current density) of PANI and PANI/KWB composites.

The enhancement of the SC is also confirmed by the GCD curves, as shown in Fig. 5.10a. The SC of the PANI and PANI/KWB at various current densities is calculated from the discharge curves and the results were shown in Fig. 5.10b. In general, the SC of all samples decreases with the increase of current density. The SC of PANI, PANI/KWB-1, 2, 3, 4 and 5 are 343, 236, 350, 375, 447 and 323 F g\(^{-1}\), respectively, at a current density of 0.5 A g\(^{-1}\). The SC of PANI/KWB composites first increases and
then decreases with the increase of the An/KWB mass ratio. These results are further evidences for the inference form CV curves. Furthermore, the SC of PANI/KWB-4 is higher than that of the other PANI/KWB composites at various current densities. This is due to $S_{BET}$ of PANI/KWB-4 is bigger than that of the other samples, resulting in the contact area of PANI/KWB-4 and electrolyte being biggest among all samples.

To further study the capacitive behavior samples, EIS was used to measure the charge transport and ion diffusion. The Nyquist plots of PANI and PANI/KWB are showed in Fig. 5.11. In EIS curve, the internal resistance (Rs, including the intrinsic resistance of the sample, the contact resistance between the sample and current collector, and the resistance of bulk electrolyte) can be obtained from the point intersecting with the real axis in the region of high frequency. The semicircle is due to the charge transfer and which will affect capacitive behavior at high current loading [45]. The inset shows that Rs of PANI/KWB-1, 2, 3, 4 and 5 are 1.19, 1.09, 1.10, 1.01 and 1.20 $\Omega$, respectively. It is worth noting that Rs of PANI/KWB-4 is lower than those of PANI/KWB composites. The semicircle radius of PANI/KWB-4 is lower than the other samples, means that the PANI/KWB-4 possesses the lowest charge resistance among PANI/KWB composites. Moreover, the nearly vertical line at lower frequency of PANI/KWB-4 indicates the
pure capacitor behavior as an ideal capacitor [46, 47]. In short, the analyses of EIS reveal that PANI/KWB-4 sample possesses an excellent capacitive performance, and this result agrees with the analyses of GCD and CV.

Fig. 5.12 CV curves of KWB and PANI/KWB-4 in three-electrode asymmetric system at a scan rate of 0.01 V s\(^{-1}\) (a)

In order to further evaluate the potential application of PANI/KWB in supercapacitors, an asymmetric two-electrode system with the PANI/KWB-4 as the positive electrode and the KWB as the negative electrode is assembled (the two-electrode system denoted as PANI/KWB-4//KWB). The voltage window is a key factor for two-electrode system, for it can impact the energy density of supercapacitors [48]. In three-electrode system, the stable potential window of KWB is -0.8~0.2 V, while PANI/KWB-4 is table between -0.2~0.8 V (as shown in Fig. 5.12). Accordingly, it is expected that the potential window of PANI/KWB-4//KWB can be extend up to 1.6 V.

The comparative CV curves of PANI/KWB-4//KWB at different voltage windows were measured at a scan rate of 0.01 V s\(^{-1}\) to evaluate the optimized voltage window (as shown in Fig. 5.13a). It can be found that all CV curves exhibit a rectangular shape when the voltages are below 1.6 V. There is no significant increase of anodic current even when voltage reach to 1.6 V, which mean that the electrolyte is not being decomposed due to the storage of nascent hydrogen on the electrode below the
thermodynamic potential for water decomposition [49]. Furthermore, a redox peak appears at 1.5 V in CV curve when voltage is 1.6 V, indicating that the electric double-layer capacitance and pseudocapacitance simultaneously exist. The SC of PANI/KWB-4//KWB calculated from GCD curves at 1 A g⁻¹ at different voltage windows are displayed in Fig. 7c. The SC increases from 57 to 77 F g⁻¹ as the voltage window increase from 1.0 to 1.6 V. The GCD curve still maintains nearly symmetric even at 1.6 V, suggesting the fast current response and small resistance [50]. These above data analyses show that the best voltage window of PANI/KWB//KWB is 0~1.6 V.

Fig. 5.13 CV (a, scan rate of 0.01 V s⁻¹) and GCD (b, current density of 1 A g⁻¹) curves of two-electrode asymmetric system (using PANI/KWB-4 and KWB as the positive and negative electrode, respectively) at different potential windows;

Fig. 5.14 CV (a) and GCD (b) curves of asymmetric system measured at different scan rates and current densities in a potential window of 1.6 V, respectively (using PANI/KWB-4 and KWB as the positive and negative electrode, respectively).
The CV curves of PANI/KWB-4//KWB at different scan rates (0.01~0.1 V s\(^{-1}\)) measured between 0~1.6 V are displayed in Fig. 5.14. The current increases as the scan rate increases, while its shape gradually deviates from the rectangular shape, which suggests that the PANI/KWB-4//KWB gradually deviates from the ideal capacitive behavior with the scan rate increase. GCD measurements were made at different current densities to further evaluate the capacitive performance of the PANI/KWB-4//KWB at the voltage window of 0~1.6 V, as shown in Fig. 5.14. The SC of the PANI/KWB-4//KWB is also obtained from GCD curves. The SC is 77 F g\(^{-1}\) at a current density of 1 A g\(^{-1}\), and still maintains at 55 F g\(^{-1}\) when the current density increases to 8 A g\(^{-1}\).

![Graph showing cycle stability](image)

Fig. 5.15 Cycle stability of asymmetric system (using PANI/KWB-4 and KWB as the positive and negative electrode, respectively).

The long-term cycling stability is an essential property of Supercapacitors. The cycle stability of the PANI/KWB-4//AB is evaluated by the GCD test in a voltage window of 0~1.6 V at a current density of 8 A g\(^{-1}\) for 2000 cycles (as shown in Fig. 8f). The PANI/KWB-4//KWB retains about 87% of the initial SC after 2000 cycles, which is comparable to those of other polymer-based asymmetric or symmetric two-electrode system, including PANI/MnO\(_2\)/carbon//activated grapheme (70% retention after 5000 cycles) [51], PANI/RGO//PANI/RGO (85% retention after 1000 cycles) [52],
graphene/RuO$_2$/graphene/PANI (70% retention after 2500 cycles) [53], RGO–PANi//RGO–PANi (80% retention after 1000 cycles) [54], manganese dioxide/reduced graphene oxide/indium tin oxide (MRI)//polypyrrole/reduced graphene oxide/indium tin oxide (PRI) (75% retention after 2000 cycles) [55], and 3D-hollow balls of graphene and polyaniline (HBGP)//3D-HBGP (86% retention after 500 cycles) [56].

Supercapacitors with excellent performance, the high power and energy density are expected. The energy and power density of the PANI/KWB-4//KWB are calculated from the GCD curves according to Eqs. (2) and (3), and the results of Ragone plots are shown in Fig. 8. Basically, the energy density of PANI/KWB-4//KWB decreases gradually with the power density increases. The maximum energy density is 27.29 Wh kg$^{-1}$ at a power density of 800 W kg$^{-1}$ and still maintains 19.38 Wh kg$^{-1}$ at a ‘power density of 6400 W kg$^{-1}$. These values are superior to the previous reported that of majority asymmetric or symmetric devices, such as RGC-RuO$_2$//RGO-PANi (26.3 Wh kg$^{-1}$, 150 W kg$^{-1}$) [54], RGC-PANi//RGO-PANi (13.9 Wh kg$^{-1}$, 150 W kg$^{-1}$) [54], PANI/CMK-3//CMK-3 (23.8 Wh kg$^{-1}$, 206 W kg$^{-1}$) [57], graphene-wrapped PANI
nanofibers/activated carbon (19.5 Wh kg\(^{-1}\), 738.9 W kg\(^{-1}\)) [58], layered PPy fibre/graphene film (20.6 Wh kg\(^{-1}\), 1.28 kW kg\(^{-1}\)) [50], CNT/MnO\(_2\)/GR//CNT/PANI (24.8 Wh kg\(^{-1}\), 200 W kg\(^{-1}\)) [59], XC-72//rGO/CNT/PPy (14.3 Wh kg\(^{-1}\), 6.62 kW kg\(^{-1}\)) [60], and Co(OH)\(_2\)/NEMG//PPy/rGO (24.9 Wh kg\(^{-1}\), 224 W kg\(^{-1}\)) [61]. These findings suggest the PANI/KWB-4//KWB holds a great promise in the practical application.

### 5.4 Conclusions

In this work, PANI/KWB composites were synthesized via in-site polymerization method. In this synthesis process, KWB (derived from the biomass waste of sugarcane bagasse) and [CMMIm]Cl IL acted as the scaffolds for PANI growth and paint, respectively. The electrochemical performance of PANI/KWB was closely related to the mass ratio of An/KWB. PANI/KWB-4 showed the best electrochemical performance when the mass ratio of An/KWB was 40 in three-electrode system. In addition, the assembled asymmetric two-electrode system based on PANI/KWB-4 and KWB exhibits high energy density (27.29 Wh kg\(^{-1}\)), high power density (800 W kg\(^{-1}\)) and excellent cycling stability of 87% SC retention after 2000 cycles within the voltage window of 0~1.6 V. The PANI/KWB composites synthesized with facile and low cost method shows an excellent electrochemical performance, marking it a promising candidate in energy storage devices.


References


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Chapter 6 Fe₃O₄ / biocarbon composites with superior performance in supercapacitors

6.1 Introduction

The global energy crisis and the accompanying environmental pollution has been becoming more and more critical. Therefore, the efficient energy storage conversion equipment and the efficient use of renewable resources have become an urgent requirement [1, 2]. Supercapacitors are novel energy storage devices, which exhibits fast charge/discharge rate and excellent cycle life. Recently, supercapacitors have attracted much more attention due to its superior energy density compared with batteries and other common capacitors [3-6]. Based on the different charge storage mechanisms, supercapacitors can generally be divided into two types: the electrical double-layer capacitors (EDLC) and the pseudocapacitors [7, 8]. In pseudocapacitors, the reversible multielectron redox faradaic reactions makes pseudocapacitors revealing much higher specific capacitance and energy density [9]. Metal oxides and conductive polymer are usually used as pseudocapacitors electrode materials, such as RuO₂, MnO₂, NiO, polyaniline, polypyrrole and PEDOT [10, 11]. In contrast to pseudocapacitors, the energy storing of EDLC arises from the accumulation of ionic charges at the interface between the electrode and the electrolyte. Carbon materials with high specific surface area and good conductivity are usually used in EDLC electrode, but their low energy density limited their further application [12-15].

In recent year, development and application of carbon materials have been becoming a central issue of materialogy [16]. Among multifarious carbon materials, biocarbon has been the most concerned due to its high specific surface area, abundant surface oxygen functional groups, low cost and simple preparation method [17-19]. Many biocarbon derived from biomass including rice husk [20], coconut fiber [21] and
tobacco stems [22] have been prepared. Sugarcane bagasse is a kind of typical biomass waste, and it is sometimes used as animal feed. Besides, it is always discarded and thus leads to many disposal problems. Fortunately, the sugarcane bagasse has abundant hollow and tubular morphology, which can be facilely converted into hierarchical porous biocarbon material via simple pyrolysis and alkali activated [23, 24]. Especially, compared with other biocarbon materials, the sugarcane bagasse derived biocarbon not only has the above advantages, but also presents an allium-giganteum-like microstructure (has been confirmed in our previous work [25]). This particular structure of sugarcane bagasse derived biocarbon is very beneficial to ion transport if this biocarbon is applied to supercapacitors.

As one of transition-metal oxides, magnetite (Fe$_3$O$_4$) have two valence states (Fe$^{2+}$, Fe$^{3+}$) which make a promise to be applied as supercapacitor material. Furthermore, this material has advantages of low cost and environmentally affinity [26]. However, the intrinsic problem of Fe$_3$O$_4$ supercapacitor electrode is lack of electrical conductivity (the specific capacitance of pure Fe$_3$O$_4$ is 60-80 F g$^{-1}$) [27]. To solve this problem, the common countermeasure is preparation of hybrid nanostructured material by integrating Fe$_3$O$_4$ with a carbon host, where the carbon host acts as the conductive network [28, 29]. Until now, many kind of Fe$_3$O$_4$/carbon composites have been prepared. Kim et al. [30] prepared Fe$_3$O$_4$/carbon nanotubes (CNTs) nanocomposites with a high specific capacitance of 165 F g$^{-1}$. The Fe$_3$O$_4$/reduced graphene oxide [31] and Fe$_3$O$_4$/active carbon [32] were also prepared, and both of them had high surface area/conductivity serve of carbon material.

Hence, in this paper, an efficient method for preparation of Fe$_3$O$_4$/biocarbon composites (KBFe) was developed. The biocarbon (KWB) was obtained by one-step direct pyrolysis of sugarcane bagasse, and then the KBFe composites were obtained by coprecipitation method under different mass ratios of Fe$_3$O$_4$ to KWB. As a result, the chemical component and specific surface area of KBFe had a profound effect on the electrochemical performance of KBFe, and the highest specific capacitance of KBFe reached 342 F g$^{-1}$ at a current density of 1 A g$^{-1}$. In addition, an asymmetric two-electrode system was fabricated by employing KWBM as positive electrode and KBFe-
4 as negative electrode. The asymmetric system also possessed a high energy density and a high cycle stability.

6.2 Experimental

6.2.1 Materials

Sugarcane bagasse was crushed by family-use grinder before use. Ethanol, potassium hydroxide (KOH), hydrochloric acid (HCl), iron dichloride tetrahydrate (FeCl$_2$·4H$_2$O), ferric chloride hexahydrate (FeCl$_3$·6H$_2$O), and sodium hydroxide (NaOH) were purchased from Nacalai Tesque, Inc. (Kyoto, Japan). All the chemicals were analytical grade and used without further purification.

6.2.2 Preparation of KOH-activated biocarbon (KWB) derived from sugarcane bagasse

KWB was obtained by KOH activation method and the typical process as follows [25]: first, sugarcane bagasse was cut into small pieces (opening size approximately 2 mm) and washed with distilled water for 8 h at 90 °C, and then dried at 80 °C for 24 h. Second, 1 g cleanly sugarcane bagasse and 0.6 g KOH were poured into 12 ml absolute ethanol with stirring and heated at 60 °C until all of the ethanol was evaporated, and then dried at 70 °C for 12 h. Third, the remnant mixture was pyrolyzed in a tubular furnace in N$_2$ atmosphere at 800 °C (heating rate is 10 °C min$^{-1}$) for 2 h. Finally, the obtained carbon material was washed by 1 M HCl solution and deionized water till the filtrate became neutral and collected after vacuum dry.

6.2.3 Preparation of Fe$_3$O$_4$/KWB composites (KBFe)

KBFe composites were prepared via a coprecipitation method. In a typical process, 0.24 g FeCl$_3$·6H$_2$O and 0.10 g FeCl$_2$·4H$_2$O (the molar ratio of Fe$^{3+}$/Fe$^{2+}$ is 1.8:1) were dissolved in 100 ml deionized water, and then 0.2 g KWB was added with a vigorous stirring for 10 min. Subsequently, the NaOH solution (1 M) was added drop by drop to
the mixture until the pH value of mixture reached to 11. Then the mixture was heated at 90 °C for 1 h with a vigorous stirring. Finally, the product was removed from the mixture solution by suction and washed several times by deionized water. The black solid product was dried in vacuum at 80 °C for 12 h to obtain the KBFe-1 composite (the theoretical value of mass ratio of Fe₃O₄ to KWB is 0.25:1). The other products were denoted as KBFe-2, KBFe-3, KBFe-4 and KBFe-5 for the products prepared with the theoretical value of mass ratio of Fe₃O₄ to KWB is 0.5:1, 0.75:1, 1:1 and 1.5:1, respectively.

6.2.4 Preparation of pure Fe₃O₄

Fe₃O₄ was prepared via a coprecipitation method. In a typical process, 0.24 g FeCl₃·6H₂O and 0.10 g FeCl₂·4H₂O (the molar ratio of Fe³⁺/Fe²⁺ is 1.8:1) were dissolved in 100 mL deionized water with a vigorous stirring for 10 min. Subsequently, the NaOH solution (1 M) was added drop by drop to the mixture until the pH value of mixture reached to 11. Then the mixture was heated at 90 °C for 1 h with a vigorous stirring. Finally, the product was removed from the mixture solution by suction and washed several times by deionized water. The black solid product was dried in vacuum at 80 °C for 12 h to obtain the Fe₃O₄.

6.2.5 Preparation of MnO₂/KWB composites (KWBM)

The KWBM composites were prepared by directly reacting KMnO₄ with KWB. In a typical process, 0.2 g KMnO₄ dissolved in 100 ml deionized water, and then 0.1 g KWB (the mass ratio of KMnO₄/KWB is 0.5:1) added the above solution. Subsequently, the mixture was stirred under water bath at 70 °C until the color of solution changed from modena to tawny. The products removed from solution and washed with deionized water for several times, and dried in vacuum at 80 °C for 12 h to obtain of the KWBM composite.
6.2.6 Structure characterization

The morphology and microstructure of samples were observed by scanning electron microscopy (SEM) and transmission electron microscopy (TEM). The SEM was performed by S-4300 instrument (Hitachi, Japan). Transmission electron microscopy (TEM) was carried out on a TF20 field emission electron microscope (FEI, USA). The chemical composition was investigated by energy-dispersive X-ray spectroscopy (EDX). The specific surface areas of samples were determined by Brunauer-Emmett-Teller (BET) method of nitrogen sorption at 77 K using ASAP 2020 analyzer (Micromeritics, USA). The crystallographic structure was carried out via X-ray diffraction (XRD) measurements at XRD-6000 instrument (Shimadzu, Japan) with Cu-Kα radiation (λ=0.154 nm) at a speed of 5° min⁻¹ from 2θ = 5° to 90°. The surface states of samples were analyzed via X-ray photoelectron spectroscopy (XPS) using a PHI Quantum 5000 instrument (ULVAC-PHI, Japan) equipped with Al Kα radiation source.

6.2.7 Electrochemical testing

Conventional three-electrolyte system and asymmetric two-electrode system were used in supercapacitors to measure the electrochemical characterization of KBFe using Models 1287 electrochemical workstations (Solartron Analytical, UK). The test electrodes were first prepared by mixing the KBFe or KWBM with carbon black (CB) and PTFE at a mass ratio of 80:15:5, then the mixing were adequately ground to obtain a homogeneous paste for coating onto the Ni foam to forming a current collector. After that, the test electrodes were dried in vacuum at 80 °C for 24 h, then compressed at 10 MPa for 5 min. In three-electrode system, the KBFe composites test electrodes was used as working electrode, a saturated calomel electrolyte as the reference electrolyte, and a platinum-wire electrodes as counter electrolyte. Cyclic voltammetry (CV) and galvanostatic charge/discharge (GCD) were conducted in a potential window from -1.2 to 0 V for KBFe composites and -0.3 to 0.4 V for KWBM composite respectively. Electrochemical impedance spectroscopy (EIS) was accomplished at open circuit potential with perturbation of 5 mV in the frequency range from 10⁵ to 10⁻² Hz. For
asymmetric two-electrode system, the system were assembled with KBFe-4 and KWBM test electrodes as working electrodes. The CV and GCD tests were tested at different voltage windows. A 1M KOH solution was used as an electrolyte for all electrochemical measurements.

The standard calomel reference electrode (SCE) based on GCD curves was evaluated according to the following equation [33] (Eq. 1):

\[
C_m = \frac{(I \cdot t)}{(\Delta V \cdot m)} \quad \text{(Eq. 1)}
\]

where \(C_m\) (F g\(^{-1}\)) is the specific capacitance (SC), \(I\) (A) is discharge current, \(t\) (s) is the discharge time, \(\Delta V\) (V) is the voltage window, and \(m\) (g) is the total mass of active material on working electrode in a two-electrode system or three-electrode system.

The energy density (E, Wh kg\(^{-1}\)) and power density (P, W kg\(^{-1}\)) of the asymmetric supercapacitors were calculated from the discharge curves according to the following equations [34]:

\[
E = \frac{1}{2}C_m(\Delta V)^2 \quad \text{(Eq. 2)}
\]

\[
P = \frac{E}{t} \quad \text{(Eq. 3)}
\]

where \(C_m\) (F g\(^{-1}\)) is SC, \(t\) (h) is the discharge time, and \(\Delta V\) (V) is the potential window from GCD curves of two-electrode system, respectively.

### 6.3 Results and discussion

#### 6.3.1 Microstructural characterization

As shown in Fig. 6.1, the KWB was obtained by carbonized and activated sugarcane bagasse at one step. It is clearly found that the KWB (SEM image as shown in Fig. 6.2a) presented an allium-giganteum-like structure with high interconnected frameworks which is favorable for the ion diffusion and suitable for preparation of Fe\(_3\)O\(_4\)-based composites [35]. And then, Fe\(^{3+}\) and Fe\(^{2+}\) were absorbed on the surface of KWB. Finally, the surface of KWB was uniformly covered by Fe\(_3\)O\(_4\) nanoparticles when NaOH solution was dropped slowly into the mixture. The Fe\(_3\)O\(_4\) nanoparticles grew on the surface of KWB because of the chemical reaction between Fe\(^{3+}\) and Fe\(^{2+}\) through
coprecipitation method (Re. 1) [30, 32]:

\[
2\text{Fe}^{3+} + \text{Fe}^{2+} + 8\text{OH}^- \rightarrow \text{Fe}_3\text{O}_4 + 4\text{H}_2\text{O} \quad (\text{Re.1})
\]

Fig. 6.1 Illustration of preparation procedure of KBFe composites

The morphology of the prepared samples were examined by SEM analysis (Fig. 6.2). An interconnected framework is clearly observed in the KWB (Fig. 6.2a) caused by a lot of secondary pores in the primary pores, which is favorable for the ion diffusion and is suitable for preparation of carbon-based composites. As shown in Fig. 6.2b, \(\text{Fe}_3\text{O}_4\) nanoparticles exhibit a globular structure forming agglomerates. The surface and interconnected frameworks of KWB in the KBFe composites were coated by \(\text{Fe}_3\text{O}_4\) nanoparticles. However, it is noteworthy that a little of uncoated KWB was observed in KBFe-1 (Fig. 6.2c and d). The uncoated KWB gradually decreased with the increase in the mass ratio of \(\text{Fe}_3\text{O}_4/\text{KWB}\) until the KWB was completely coated by \(\text{Fe}_3\text{O}_4\) nanoparticles when the mass ratio of \(\text{Fe}_3\text{O}_4/\text{KWB}\) is 1 (KBFe-4, Fig. 6.2i and j). In addition, the interconnected structure of KWB was filled by \(\text{Fe}_3\text{O}_4\) nanoparticles when the mass ratio of \(\text{Fe}_3\text{O}_4/\text{KWB}\) was 1.5 (KBFe-5, Fig. 6.2k and l). This phenomenon will affect the ion diffusion and thus the electrochemical performance of KBFe.
Fig. 6.2 SEM images of KWB (a), Fe$_3$O$_4$ (b), KBFe-1 (c, d), KBFe-2 (e, f), KBFe-3 (g, h), KBFe-4 (i, j) and KBFe-5 (k, l) composites.
Fig. 6.3 TEM images of the KBFe-4 nanocomposites (a, b); the inset shows the corresponding SAED pattern taken from the edge of Fe₃O₄ nanoparticles; EDX mapping image of C, O and Fe in KBFe-4 (c). EDX spectrum of the KBFe-4 nanocomposites (d).

The morphology of KBFe-4 was further investigated by TEM. As we all know, the diffraction rings of selected area electron diffraction pattern (SAED) correspond to polycrystals, and the diffraction points of SAED correspond to single crystals [36, 37]. For the inset of Fig. 6.3a, it can be clearly observed the diffraction rings, which correspond to the polycrystals of magnetite Fe₃O₄ (The polycrystals of Fe₃O₄ is including α-Fe₂O₃, γ-Fe₂O₃, FeO and so on [38, 39]). About the magnetite of Fe₃O₄, it will be further confirmed by XRD. In addition, for Fig. 6.3a, it can be clearly observed that only few Fe₃O₄ was aggregating, however, most Fe₃O₄ nanoparticles were uniformly loaded on the surface of KWB biocarbon with several nanometers size due
to the conductive of carbon better for dispersion of Fe$_3$O$_4$ [40]. Furthermore, EDX mapping image reveals the co-existence and homogenous distribution of C, Fe and O across the core-shell nanostructure of KBFe-4; also, the EDX spectrum of the KBFe-4 nanoparticles demonstrates the covering effect of Fe$_3$O$_4$ for the high content of Fe element.

Fig. 6.4 N$_2$ adsorption-desorption isotherms (a), pore size distribution by BJH method (b) of the prepared KWB biocarbon, KBFe-1, KBFe-2, KBFe-3, KBFe-4 and KBFe-5 composites.

Table 6.1 Specific surface area and pore size of samples. ($S_{BET}$ - the specific surface area; $P$ - the main pore size; $V_t$ - the total pore volume)

<table>
<thead>
<tr>
<th>Sample</th>
<th>$V_t$ (cm$^3$ g$^{-1}$)</th>
<th>$S_{BET}$ (m$^2$ g$^{-1}$)</th>
<th>$P$ (nm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>KWB</td>
<td>0.755</td>
<td>1433.06</td>
<td>3.03</td>
</tr>
<tr>
<td>KBFe-1</td>
<td>0.656</td>
<td>936.64</td>
<td>4.19</td>
</tr>
<tr>
<td>KBFe-2</td>
<td>0.502</td>
<td>652.01</td>
<td>4.55</td>
</tr>
<tr>
<td>KBFe-3</td>
<td>0.488</td>
<td>586.51</td>
<td>4.94</td>
</tr>
<tr>
<td>KBFe-4</td>
<td>0.513</td>
<td>490.93</td>
<td>6.83</td>
</tr>
<tr>
<td>KBFe-5</td>
<td>0.508</td>
<td>346.86</td>
<td>9.37</td>
</tr>
</tbody>
</table>

The nitrogen adsorption/desorption isotherm and corresponding pore size distribution of samples were shown in Fig. 6.4a and b. Obviously, the KWB shows the
typically combined I/IV type sorption isotherms, indicating a poly porous structure [41, 42]. As given in Table 6.1, the $S_{BET}$ of KWB is up to 1433.06 m$^2$ g$^{-1}$, which results in KWB suitable for acting as the scaffolds to combine with the active materials of Fe$_3$O$_4$. For KBFe composites, the Fe$_3$O$_4$ nanoparticles were covered on the surface of KWB, which will block the mesopores and micropores of KWB. The $S_{BET}$, P and Vt of KBFe can be controlled by adjusting the mass ratio of Fe$_3$O$_4$/KWB. As shown in Table 1, the $S_{BET}$ and Vt of KBFe significantly decreased with the increase of mass ratio of Fe$_3$O$_4$/KWB. By contrary, the P of KBFe significantly increased when the mass ratio of Fe$_3$O$_4$/KWB increased. These results indicate that the micropores and mesopores structure of KWB gradually disappeared. However, the Vt and $S_{BET}$ of the KBFe-4 still increased to 0.513 cm$^3$ g$^{-1}$ and 490.93 m$^2$ g$^{-1}$, respectively. These acceptable pore structure and specific surface area of KBFe-4 have a great potential for electrode material in supercapacitors application.

Fig. 6.5 X-ray diffraction patterns of the prepared KWB biocarbon, KBFe-1, KBFe-2, KBFe-3, KBFe-4 and KBFe-5 composites.

The phase and crystallinity of samples were tested by XRD and the results were shown in Fig. 6.5. A broad diffraction peak centered at 23.1° can be attributed to the (002) plane of graphite resulted from its amorphous nature. In addition, a weak
diffraction peak at 43.4° corresponded to the (101) plane of graphite, which is owing to the complete transformation of sugarcane bagasse to a kind of activated carbon [43, 44]. For KBFe composites, the six diffraction peaks appearing at 30.2°, 35.5°, 43.3°, 53.8°, 57.2° and 62.7° are indexed to the diffraction of (220), (311), (400), (422), (511) and (440) planes of the Fe₃O₄ nanoparticles, respectively (JCPDS no. 19629) [45]. Additionally, the diffraction peak intensity of KWB (peak centered at 23.1°) decreased with the increased mass ratio of Fe₃O₄/KWB. This phenomenon further shows that the surface of KWB is gradually coated by Fe₃O₄ nanoparticles as the mass ratio of Fe₃O₄/KWB increased.

<table>
<thead>
<tr>
<th>Samples</th>
<th>C/at. %</th>
<th>O/at. %</th>
<th>Fe/at. %</th>
<th>Fe/C (%)</th>
<th>O-Fe (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>KWB</td>
<td>84.76</td>
<td>15.24</td>
<td>--</td>
<td>--</td>
<td>--</td>
</tr>
<tr>
<td>KWFe-1</td>
<td>67.63</td>
<td>22.67</td>
<td>9.70</td>
<td>14.34</td>
<td>30.81</td>
</tr>
<tr>
<td>KWFe-2</td>
<td>63.29</td>
<td>23.79</td>
<td>12.96</td>
<td>20.47</td>
<td>39.32</td>
</tr>
<tr>
<td>KWFe-3</td>
<td>55.61</td>
<td>29.61</td>
<td>14.78</td>
<td>26.58</td>
<td>46.39</td>
</tr>
<tr>
<td>KWFe-4</td>
<td>42.63</td>
<td>36.01</td>
<td>21.36</td>
<td>50.11</td>
<td>60.00</td>
</tr>
<tr>
<td>KWFe-5</td>
<td>42.82</td>
<td>35.77</td>
<td>21.41</td>
<td>50.00</td>
<td>60.15</td>
</tr>
</tbody>
</table>

The surface chemical composition of KWB and KBFe composites were examined by XPS analysis (Fig. 6.6). The survey spectra of samples are shown in Fig. 6.6a. Except for the signals of C and O elements, there is a signal of Fe element in the XPS spectra of KBFe when compared with KWB. Table 6.2 indicates that the contents of Fe and O progressively increased with the increase of mass ratio of Fe₃O₄/KWB; however, a reverse situation occurred in the content of C. More information about the change of surface chemistry was further acquired from the deconvoluted high-resolution XPS spectra. The Fe 2p spectra of KBFe-4 (Fig. 6.6b) presents two main peaks located at 711.4 and 725 eV due to Fe 2p₃/2 and Fe 2p₁/2 with a spin-energy separation of 13.6 eV, which confirms that the oxide in the sample is magnetite Fe₃O₄ [46]. The peak of Fe
2p_{3/2} was deconvoluted into two peaks at 710.8 and 712.8 eV, and that of Fe 2p_{1/2} was deconvoluted into two peaks at 724.6 and 726.3 eV. The peaks of 710.8 and 726.3 eV was attributed to the Fe^{3+} of Fe_{3}O_{4}, and the others (peak at 712.8 and 724.6 eV) was assigned to the Fe^{2+} of Fe_{3}O_{4} [47, 48].

![Fig. 6.6 XPS survey spectra of KWB biocarbon, KBFe-1, KBFe-2, KBFe-3, KBFe-4 and KBFe-5 composites (a); high resolution C1s (c) and O1s (e) XPS spectra of KWB high resolution Fe2p (b), C1s (d) and O1s (f) XPS spectra of KBFe-4 composite.](image)
Fig. 6.7. High resolution O1s XPS spectra of KBFe-1 (a), KBFe-2 (b), KBFe-3 (c) and KBFe-5 (d) composites.

The C1s spectra of KWB and KBFe-4 (Fig. 6.6c and e) were deconvoluted into three components. The peaks of 284.4, 285.8 and 288.8 eV were attributed to C–C, C–O, and C=C bonds resulted from the sp^2-hyridised carbon atoms bonded with neighboring atoms, respectively [49]. The O1s spectrum of KWB were deconvoluted into three peaks (Fig. 6.6e) with binding energies of 531.5, 533.2 and 534.8 eV, which were attributed to -OH/C–O, C=O and C-O-OH groups, respectively [50]. However, for the O1s spectrum of KBFe-4 (Fig.5f), a new peak appeared at 530.5 eV due to the oxygen bonded with iron (O–Fe) in Fe_3O_4 crystal lattice [47]. The content of O–Fe in KBFe (it is obtained by O1s spectra of KBFe-1, 2, 3, 4 and 5, and as shown in Fig. 6.6f and Fig. 6.7) is listed in Table 6.2. The content of O–Fe significantly increased along with the increase of the mass ratio of Fe_3O_4/KWB from 0.25 to 1), and then reached to constant when the mass ratio of Fe_3O_4/KWB was higher than 1. These results indicate that the
content of Fe$_3$O$_4$ in the KBFe composites is in a dominant position when the mass ratio of Fe$_3$O$_4$/KWB is greater than 1. Additionally, these transformations evidence that the element composition of KBFe can be adjusted by the mass ratio of Fe$_3$O$_4$/KWB. In summary, the mass ratio of Fe$_3$O$_4$/KWB has a significant impact on the $S_{BET}$ and element composition and hence the electrochemical performance of the KBFe.

6.3.2 Electrochemical performance of KBFe composites

Fig. 6.8 CV curves of KBFe composites at scan rate of 0.01 V s$^{-1}$.

To explore potential applications of the KBFe composites in supercapacitors, the CV, GCD and EIS measurements were performed firstly in a three-electrode configuration. Fig. 6a illustrates the CV curves of the KBFe at a scan rate of 0.01 V s$^{-1}$. Compared with CV curve of KWB (as shown in Fig.3.11), all the CV curves of KBFe show the redox peaks, indicating the existence of pseudocapacitive of Fe$_3$O$_4$. The pseudocapacitance was resulted from the redox reaction of Fe ions, i.e., the oxidation of Fe$^{2+}$ to Fe$^{3+}$ and the reduction of Fe$^{3+}$ back to Fe$^{2+}$ [51]. In addition, the pseudocapacitance reaction of Fe$_3$O$_4$ in KOH electrolyte was accompanied by intercalation of OH- balance the extra charge with the Fe$_3$O$_4$ layers, as shown in Re. 2 and 3 [29]:
\[
\begin{align*}
\text{FeO} + 2\text{OH}^- & \leftrightarrow \text{Fe(OH)}_2 + 2e^- \quad \text{(Re. 2)} \\
2\text{Fe}^{2+} + 2\text{OH}^- & \leftrightarrow (\text{Fe}^{3+}\text{O})^+(\text{OH}^-)_2(\text{Fe}^{3+}\text{O})^+ + 2e^- \quad \text{(Re. 3)}
\end{align*}
\]

KBFe-4 exhibited a much larger current response than the other composites, and presented an excellent electrochemical performance thanks to the element composition and acceptable $S_{BET}$.

![Fig. 6.9 GCD curves of KBFe composites at current density of 1 A g\(^{-1}\)](image)

Meanwhile, GCD method was also conducted to compare the capacitive performance of KBFe composites (Fig. 6.9). Overall, all the charge curves of KBFe were nearly symmetrical to their corresponding discharge curves except that the potential is within the range of -0.8 ~ -1.2 V, indicating that the capacitance is in account of the blend of EDLC and pseudocapacitance corresponding to Re. 2 and 3. Furthermore, the discharge time of KBFe-4 is longer than that of other samples, demonstrating that the KBFe-4 possesses more excellent capacitive performance among all the KBFe composites. According to Eq. 1, the SC of KBFe-1, 2, 3, 4 and 5 were calculated to 237, 288, 332, 342 and 258 F g\(^{-1}\), respectively, at a current density of 1 A g\(^{-1}\). Furthermore, the SC of KBFe increased firstly and then decreased with the mass ratio of the Fe\(_3\)O\(_4\)/KWB increasing, which are further evidenced by the inference from CV curves.
To further evaluate the capacitive behavior of KBFe, EIS was used to evaluate the ion diffusion and charge transport. The Nyquist plots of KBFe are showed in Fig. 6.10. In general, all the EIS curves consist of three parts: a semicircle in the high frequency region, a nearly diagonal line in the medium frequency region, and a nearly vertical line in the low frequency region, which correspond to the charge-transfer resistance, diffusion resistance between electrolyte and electrode, and capacitive behavior, respectively [26]. The KBFe-4 represents the shortest 45° diagonal line at medium frequency region and the largest slope of straight line at low frequency region, indicating that KBFe-4 possesses the lowest diffusion resistance as well as best capacitive performance among all the samples. From the inset, it is found that KBFe-4 exhibits the least diameter of arc at the high frequency region, which suggests that KBFe-4 has a smaller charge transfer resistance than others. Furthermore, the internal resistance (Rs) of samples can be denoted by the point intersecting of EIS curve with the real axis at high frequency region, including the contact resistance between the active material and current collector, the intrinsic resistance of the active material and the resistance of bulk electrolyte. The inset shows that the Rs of KBFe-1, 2, 3, 4 and 5 are 2.17, 2.04, 1.97, 1.83 and 2.08 Ω, respectively, which demonstrates that the Rs of
KBFe-4 is lower than the other samples. In summary, the results of EIS conclude that KBFe-4 possesses outstanding capacitive performance, which is in line with the CV and GCD measurements.

The SC plots of KBFe composites in different current density are shown in Fig. 6.11, indicating that the SC of all samples decreased with the increase of current density. The SC downtrend of KBFe-4 is larger than that of the others, especially when current density increased from 0.5 A g\(^{-1}\) to 1 A g\(^{-1}\) (decreased from 494 F g\(^{-1}\) to 342 F g\(^{-1}\)). This is because the \(S_{BET}\) of KBFe-4 is smaller than that of the others, and thus the contact area of KBFe-4 and electrolyte is smaller than the others. However, the SC of KBFe-4 could reach to 171 F g\(^{-1}\) at the current density of 10 A g\(^{-1}\), and this value is not only larger than that of the others, but also larger than that of reported previously [52, 53], which suggests that the KBFe-4 has the most excellent fast charge/discharge performance among all the KBFe composites.
To further evaluate the capacitive performance of KBFe-4, an asymmetric two-electrode system (KWBM//KBFe-4) was assembled by using the KWBM that was prepared in our previous report as a positive electrode and KBFe-4 as a negative electrode. For confirming the potential window of KWBM//KBFe-4, the CV measurements of KWBM and KBFe-4 were firstly implemented in three-electrode system (Fig. 7a). The KWBM and KBFe-4 electrodes have different potential windows in three-electrode system (their potential windows are -0.3 to 0.4 V and -1.2 to 0 V, respectively). It is expected that the potential window of KWBM//KBFe-4 can be extended up to 1.6 V.

The CV curves of KWBM//KBFe-4 in different potential windows were performed to determine the best potential window (Fig. 6.13a). Obviously, all CV curves were close to a rectangular shape at the potential window up to 1.6 V, revealing that the two-electrode system presents an ideal capacitive behavior. Furthermore, a redox peak appears in CV curve when potential window is 0~1.6 V, illustrating that the pseudocapacitance and EDLC capacitance simultaneously exist. Fig. 6.13b shows the GCD curves of KWBM//KBFe-4 calculated from GCD at curves density of 1 A g⁻¹.
under different voltage windows. The SC of KWBM//KBFe-4 under different potential windows can be obtained by Eq. 1, where the m is the total mass of active material on two electrodes. The calculated SC increased from 28 to 81.9 F g$^{-1}$ with the potential window increased from 0.8 to 1.6 V. Hence, the above results suggest that the best potential window of KWBM//KBFe-4 is 0~1.6 V.

Fig. 6.13 CV curves (a, scan rate of 0.01 V s$^{-1}$) and GCD curves (b, current density of 1 A g$^{-1}$) of two-electrode asymmetric system (using KWBM as the positive electrode and KBFe as the negative electrode) at different potential windows;

Fig. 6.14 CV (a) and GCD (b) curves of two-electrode asymmetric system (using KWBM as the positive electrode and KBFe-4 as the negative electrode) at different scan rates and current densities in a potential window of 1.6 V, respectively.
As shown in Fig. 6.14a, the CV curves of KWBM//KBFe-4 at different scan rates (0.01~0.1 V s^{-1}) were measured between 0~1.6 V. Following the increase of scan rate, the CV curves progressively deviated from the rectangular shape, which suggests that the KWBM//KBFe-4 gradually deviated from the ideal capacitive behavior as the scan rate increased. GCD measurements were conducted at different current densities to further evaluate the capacitive performance of the KWBM//KBFe-4 at the voltage window of 0~1.6 V (Fig. 6.14b). These GCD curves were close to triangular shape, indicative of an ideal capacitive behavior. The SC of the KWBM//KBFe-4 was also obtained from GCD curves based on Eq. 1. The SC was 81.9 F g^{-1} at a current density of 1 A g^{-1}, and still maintained at 35.6 F g^{-1} when the current density increased to 10 A g^{-1}, further suggesting that the KWBM//KBFe-4 has excellent capacitive performance.

![Graph showing cycle stability](image)

**Fig. 6.15 Cycle stability of asymmetric system.**

The long-term cycling stability is an indispensable property of supercapacitors. The repeating GCD measures was used to evaluated the cycle stability of KWBM//KBFe-4 at the current density of 5 A g^{-1} in 0~1.6 V (Fig. 6.15). The KWBM//KBFe-4 exhibits an outstanding cycle stability with 89.5% SC retention after 1000 cycles, which is
comparable to those of other Fe$_3$O$_4$-based asymmetric two-electrode system, such as Fe$_3$O$_4$/Fe-CNTs//AC (82.1% retention after 1000 cycles) [54], Fe$_3$O$_4$/AC (82% retention after 500 cycles) [28], and polyaniline-DBSA/Fe$_3$O$_4$/polyaniline-DBSA/Fe$_3$O$_4$ (65% retention after 1000 cycles) [57].

![Ragone plot of KWBM//KBFe-4 asymmetric system](image)

The energy and power density of the KWBM//KBFe-4 were calculated from the GCD curves according to Eq. 2 and 3, while the results were shown in Fig. 6.15. Basically, the energy density of KWBM//KBFe-4 gradually decreased with the increase of power density. The maximum energy density is 29.1 Wh kg$^{-1}$ at a power density of 800 W kg$^{-1}$. These values are superior to the previous reports, such as Fe$_3$O$_4$@Fe$_2$O$_3$/Fe$_3$O$_4$@MnO$_2$ (26.6 Wh kg$^{-1}$, 500 W kg$^{-1}$) [47], MnO$_2$/Fe$_3$O$_4$ (8.1 Wh kg$^{-1}$, 10200 kW kg$^{-1}$) [56], Fe$_3$O$_4$/graphene/Fe$_3$O$_4$/graphene (9 Wh kg$^{-1}$, 3000 W kg$^{-1}$) [57], Fe$_3$O$_4$ composite//porous carbon (18.3 Wh kg$^{-1}$, 351 W kg$^{-1}$) [40]. These finding suggest the KWBM//KBFe-4 holds a great promise in the practical application.
6.4 Conclusions

In summary, KBFe composites were synthesized via a coprecipitation method, where the KWB derived from sugarcane bagasse acted as the scaffolds for Fe$_3$O$_4$ deposition and growth. The electrochemical performance of KBFe was closely related to the mass ratio of Fe$_3$O$_4$/KWB. KBFe-4 showed the best electrochemical performance when the mass ratio of Fe$_3$O$_4$/KWB was 1 in three-electrode system. In addition, the assembled asymmetric two-electrode system based on KBFe-4 and KWBM exhibits high energy density (29.1 Wh kg$^{-1}$), high power density (800 W kg$^{-1}$) and an excellent cycling stability of 89.5% SC retention after 1000 cycles. The KBFe composites synthesized with a facile and low-cost method showed an excellent electrochemical performance, marking them promising candidate in energy storage devices.
References


[41] K.S.W. Sing, Reporting physisorption data for gas/solid systems with special
reference to the determination of surface area and porosity.


Chapter 7 Conclusions

Supercapacitors, as one of the most promising energy storage devices, due to their good energy density, superior power density, fast charge/discharge rate, and long cycle life, which make it exhibit promising applications in portable devices, renewable energy and transportation. Based on the different charge storage mechanisms, supercapacitors can generally be divided into two types: the electrical double-layer capacitors (EDLC) and the pseudocapacitors. Carbon materials, such as active carbon, graphene and carbon nanotube are usually used in EDLC electrode, but their low energy density, limited their further application, as well as most of methods to prepare of the above carbon materials often requires complicated preparation conditions or special facilities, leading to the high cost. In the present work, first, low cost and uncomplicated method of allium-giganteum-like biocarbon (KWB) were derived from sugarcane bagasse, and then different kinds of carbon-based composite materials including manganese dioxide (MnO$_2$)/KWB, polyaniline (PANI)/KWB and iron oxide (Fe$_3$O$_4$)/KWB were obtained. All of the above materials were used as electrode material of supercapacitors, and morphology, chemical composition and electrochemical performance were investigated. Besides, due to the unique structure of KWB, all the composites show the superior performance in supercapacitors.

In Chapter 1, the research backgrounds, research significance, research purpose and the construction of this thesis are particularly described.

In Chapter 2, the properties of experimental materials used in this thesis are presented. The experimental methods and characterizations are also particularly described in this chapter.

In chapter 3, KWB were derived from sugarcane bagasse via one-step carbonization and activation method. The results show that, both the mass ratio of potassium hydroxide (KOH) to sugarcane bagasse and the temperature of carbonization could impact on the morphology and chemical composition of KWB, thereby further affect the electrochemical performance of KWB. Under the optimum conditions, the SEM of
KWB shows the excellent 3-D hierarchical porous network structure with allium-giganteum-like, which also has the best specific surface area. It promises the excellent electrochemical performance when used as electrode materials for supercapacitors. What’s more, KWB also could be carbon-base to prepare other composites in the further research.

In chapter 4, MnO$_2$/KWB (KWBM) composites have been prepared by a simple method. The MnO$_2$ nanosheets anchored on the surface of biocarbon and obtained the KWBM, and KWB was used as both scaffolds and reducing agents for MnO$_2$ growth. The analysis results demonstrated that chemical composition and microstructure of KWBM were closely related to the mass ratio of KMnO$_4$/KWB, and which further impacted electrochemical performance of KWBM profoundly. Especially, the KWBM-4 exhibited a higher specific capacitance of 402 F g$^{-1}$ at a current density of 1 A g$^{-1}$ in three-electrode system. The asymmetric two-electrode system with outstanding energy density was assembled by employing the KWBM-4 as the positive electrode and the KWB as the negative electrode. The two-electrode system displays a high energy density of 25.9 Wh kg$^{-1}$ at a power density of 750 W kg$^{-1}$ within a potential range of 0-1.5 V. Furthermore, the system exhibited high cycle stability with only 5.8 % loss of its initial capacitance after 2000 cycles.

In chapter 5, a coral-like polyaniline/KWB (PANI/KWB) composites was prepared via in-site polymerization method. The PANI/KWB composites were obtained using KWB as the scaffolds for PANI growth and [CMMIm]Cl ionic liquid as dopant. The different mass ratio of aniline monomer (An) and KWB be researched. The results indicated that chemical composition and microstructure of PANI/KWB were closely related to the mass ratio of aniline monomer (An) and KWB, and which further impacted electrochemical performance of PANI/KWB profoundly. Especially, the PANI/KWB-4 exhibited a highest specific capacitance of 447 F g$^{-1}$ at a current density of 0.5 A g$^{-1}$. Furthermore, the asymmetric two-electrode system based on PANI/KWB-4 as positive electrode and KWB as negative electrode is successfully assembled with a voltage window of 0-1.6 V, exhibiting high energy density (27.3 Wh kg$^{-1}$) and power density (800 W kg$^{-1}$), and excellent cycling stability (87% capacitance retention after
2000 cycles).

In chapter 6, Fe$_3$O$_4$-KWB nanocomposites (KBFe) have been prepared via a chemical coprecipitation method under different mass ratio of Fe$_3$O$_4$ to KWB. As a result, the chemical composition and microstructure of the KBFe were intimately related to the mass ratio of Fe$_3$O$_4$ and KWB, which impacted the electrochemical performance of the KBFe profoundly. Especially, the KBFe-4 displayed the highest specific capacitance of 342 F g$^{-1}$ at a current density of 1 A g$^{-1}$. Furthermore, the asymmetric two-electrode system based on KWBM composite as positive electrode and KBFe-4 as negative electrode was successfully assembled with a voltage window of 0~1.6 V, and exhibited high energy density (29.1 Wh kg$^{-1}$) and power density (800 W kg$^{-1}$), along with excellent cycling stability (89.5% capacitance retention after 1000 cycles).
Publication

I. 審査付投稿論文


注：博士論文テーマ関連：3編（1）〜（3）, その他：1編（4）

II. 国際会議発表論文


注：博士論文テーマ関連：1件

III. 国内学会発表

[2] 陳姣、邱建輝、境英一、伊藤一志、王濱、馮輝霞、小宮山崇夫。スーパーキャパシタ適用のためのサトウキビバガスの水酸化カリウム活性化にによる炭素電極の創製。2016年9月日本機械学会東北支部第52期秋季講演会、秋田市。

[3] 陳姣、邱建輝、境英一、イオン液体を用いたin situ重合によるポリビロール/黒鉛複合材料の作製および性能評価。2015年9月日本機械学会東北支部第51期秋季講演会、いわき市。

[4] 王濱、邱建輝、境英一、伊藤一志、陳姣、馮輝霞、尾藤輝夫。カーボンナノワイヤー/二酸化マンガン複合材料の創製およびスーパーキャパシタの適用。2016年9月日本機械学会東北支部第52期秋季講演会、秋田市。

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