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Electrochemical energy storage and adsorptive dye removal of platanus fruit derived porous carbon

Xin Wang, a Zhiyong Gao, a Jiuli Chang, a Dapeng Wu, a,b Xiaorui Wang, a Fang Xu, a,b Yuming Guo, a,b Kai Jiang a,b

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Activated platanus fruit carbon (a-PFC) was synthesized by pyrolysis carbonization and alkali activation treatment of a easily available biomass, platanus fruit (PF). The carbonization yields platanus fruit carbon (PFC) sample with partially graphitized phase, and the followed KOH activation creates highly porous texture containing a large fraction of micropores and therefore a high specific surface area. Both of which are beneficial for surface relevant applications such as electrode material for electrochemical capacitor and adsorbent for sucking of organic dyes. The supercapacitor based on a-PFC3, which was synthesized with KOH-to-PFC activation ratio of 3, offers a high gravimetric capacitance of 216 F g−1 at 1 A g−1, a high rate performance and cycling stability, highlighting the potential of a-PFC3 in electrochemical energy storage. The high specific surface area also makes a-PFC3 an efficient adsorbent for sucking of methylene blue (MB) from aqueous solution.

Introduction

Energy and environment are two major topics that have attracted widespread concerns in scientific community. With the ever-growing consumption of fossil energy and the concomitant environmental pressure, it is necessary to develop reliable and sustainable energy sources without pollution emission. Supercapacitor is an attractive electrochemical power source owning superior power density, cycling stability, charging-discharging duration and operational safety over other rechargeable batteries, as well as the promptly increased capacitance and energy output.1,2 hence it is regarded as a promising energy storage and supply device for portable electric products, electric vehicles, etc. Electrode material is the key component of supercapacitor, an ideal electrode material requires highly accessible surface area to electrolyte, well conductivity, structural stability and chemical inertness against electrolyte. On the other hand, organic dyes are important aquatic pollutants discharged from printing, dyeing, textile and painting industries, which seriously destroy the aquatic environment and may cause some threats to health. Because some dyes are difficult to be degraded, the adsorptive treatment is a feasible pathway to remove and even recycle them. The adsorptive treatment of organics necessitates highly porous adsorbent with high surface area and structural stability. Porous carbon can well meet the requirements of the abovementioned capacitive material and adsorbent. As a result, great efforts were devoted to the preparation of porous carbon materials and the applications in supercapacitor and organic dyes removal.

So far, various porous carbon materials including carbon nanotubes,3,4 graphene5,6 and hetero-elements doped carbon7,8,10 were synthesized as electrode materials for supercapacitors and demonstrated satisfactory performances. Meanwhile, these types of materials also showed high adsorption performances for removal of various dyes.11-13 Albeit excellent performances were achieved from the abovementioned materials, the relatively high costs and somewhat tedious synthesis processes limit their widespread applications. More facile and cost-effective synthesis of porous carbon materials in bulk will promote the more extensive applications. Direct pyrolysis and activation of carbonaceous substances is a competitive approach to produce porous carbons in large scale. Biomass, especially vegetations represent a big class of cheap, easily available and renewable carbonaceous substances widely distributed on our planet that sustains the global water and carbon cycles. Due to the presence of rich capillaries for transportation of water and nourishments to sustain the growth, rich pores exist in vegetations, which make them ideal feedstocks for preparation of porous carbons with high surface areas. Pyrolysis is the most direct and facile approach to transform biomass into porous carbons. Due to the abundant resource with low costs, various natural biomass, including nut shell,14,15 plant leaves,16,17 fruit peels,18,19 fish scales,20 feathers,21 hemp fiber22 and wood sawdust23,24 were widely employed as precursors for carbonization to generate porous carbon materials, and the afforded high capacitances or adsorption abilities highlighted the huge potential of biomass derived porous carbons in energy and environment related applications.

Platanus is a widely planted street tree for its strong adsorption ability toward dust and poisonous exhausts in urban atmosphere, excellent resistibility against contaminants, as well as its ultra-strong climate adaptability. The spheric fruit of platanus, namely PF contains numerous tiny fluffs, which is commonly undesirable and be discarded as waste owing to its allergy to certain susceptible population. Considering the possible high surface area associates with the slim and light fluffs, the carbonization can accomplish the resource utilization of PF and affords carbon materials for surface related applications, so the capacitive and adsorptive performances of PFC are expectable.

Herein, PF was employed as raw feedstock for the synthesis of porous carbon by pyrolysis and the subsequent alkali activation. The highly porous a-PFC demonstrated high BET surface area with overwhelming fraction of micropores, so the application performances in capacitive energy storage and adsorptive dye removal were tested and analyzed.
Experimental

Preparation of a-PFCs

All reagents in this work are of analytical grade and be used directly without further purification. Mature PF was collected after they fell from the platanus trees. The spheric PF was initially rinsed with deionized water to get rid of the adsorbed dust. After being dried and exfoliated, the inner fluff was used as crude material for carbonization and activation to afford a-PFC. In brief, the clean fluff was placed in a tube furnace and heated to 600 °C at a heating rate of 5 °C min⁻¹ under N₂ atmosphere and then maintained at this temperature for 3 h. In the followed activation process, KOH was mixed with the obtained black PFC with mass ratio of 2, 3 or 4. Then the mixture was heated at 800 °C under N₂ atmosphere for 1 h for activation treatment. After being cooled naturally, the afforded a-PFC was washed copiously with diluted HCl and deionized water to eliminate the alkali and inorganics. The afforded a-PFC sample was denoted as a-PFCₙ (n = 2, 3 or 4, according to the KOH-to-PFC mass ratio).

Characterizations

The morphologies and microstructures of the samples were characterized by scanning electron microscopy (SEM, JEOL JSM-6390), high resolution transmission electronic microscopy (HRTEM, JEOL JEM-2100), X-ray powder diffraction (XRD, Bruker D diffractometer with Cu Ka radiation), Fourier Transform Infrared Spectroscopy (FTIR, Bio-Rad FTS-40) and X-ray photoelectron spectroscopy (XPS, Thermal Fisher ESCALab 250 X-ray photoelectron spectroscopy with Al Ka radiation). The adsorption-desorption isotherms were measured on a Micromeritics Gemini 2380 surface area analyzer at 77 K. The specific surface area was calculated according to multiple point Brunauer-Emmett-Teller (BET) method. The micropore size distribution was estimated according to Horvath-Kawazoe (HK) theory. All the samples were degassed at 200 °C for 3 h prior to measurement.

Electrochemical measurements

Cycling voltammetry (CV) measurements of the electrodes were performed on a 660D workstation within potential range of 0–1 V in 1 M H₂SO₄. The three-electrode test system includes a stainless steel mesh work electrode coated with paste containing 85 wt% active material, 10 wt% acetylene black and 5 wt% polytetrafluoroethylene binder, a platinum wire counter electrode and a Ag/CuAg reference electrode. Galvanostatic charging-discharging curves of the symmetric capacitors (on Land CT2001A cell test system, Wuhan China) were performed in double-electrode mode in 1 M H₂SO₄ electrolyte over the voltage range of -1~1 V. The weight of active materials on each electrode was in the range of 1.5–2 mg, and the precise weights were measured accurately for gravimetric capacitances calculation. Electrochemical impedance spectroscopy (EIS) measurements of the electrodes were also performed on the same CHI 660D electrochemical workstation in three-electrode test system in 1 M H₂SO₄ electrolyte, impedances were recorded over the frequency range of 10⁻²–0.01 Hz with an ac perturbation of 5 mV.

The specific capacitance (Cₛ, F g⁻¹), energy density (E, Wh kg⁻¹) and power density (P, W kg⁻¹) of the symmetric capacitors were calculated from galvanostatic charging-discharging measurements according to the following equations:

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

\[
E = C_s (\Delta V - IR)^2/8 \quad \text{(Eq. 2)}
\]

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

Where I (A) represents the discharge current, \(\Delta V\) (V) is the voltage change within the discharge duration \(\Delta t\) (s), IR is the voltage drop owing to inner resistance at the initial stage of discharge process, \(m\) (g) is the total mass of active material on each electrode.

Adsorption measurements

The adsorption capability of a-PFCₙ adsorbent was evaluated using MB as adsorbate. Briefly, 10 mg of a-PFCₙ was sonically dispersed in the MB aqueous solution (350 mL, 20 mg L⁻¹) in dark to form a uniform dispersion. After that, the mixture was stirred at 20 °C. At different intervals, 2 mL of dispersion was sampled and centrifuged to afford supernatant, which was diluted to double volume for monitoring the adsorption kinetics and capacity via absorbances. In the adsorption isotherms measurements, 10 mg of a-PFCₙ was dispersed in 250, 275, 300, 325, and 350 mL of MB solution, respectively for adsorption treatment at 20 °C, and the relationship between equilibrium adsorption amounts (\(Q_e\)) and equilibrium concentrations (\(C_0\)) remaining in solutions was used for isothermal model fitting after the adsorption systems reach to equilibrium.

Results and discussion

Morphologies and microstructures
Fig. 1 presents the morphologies and microstructure of PFC and a-PFC. PFC (Fig. 1a) exhibits tubular outline with outer diameter of up to 40 µm and length in millimeter scale, which resembles to the original appearance of raw PF fluff (not shown), implying the morphological dependence on the original PF source. KOH is a commonly employed porogen for carbon materials such as graphene, carbon nanotubes etc, due to the corrosive alkali can react with carbon framework to create rich pores and enhance the specific surface area.\textsuperscript{7,22,25} The widely accepted mechanism of alkali activation follows the equation: \[ 6\text{KOH} + 2\text{C} = 2\text{K} + 3\text{H}_2 + 2\text{K}_2\text{CO}_3, \] and the produced potassium and its carbonate can further intercalate into graphitic carbon layers and etch the carbon skeleton to continue producing more open pores.\textsuperscript{7,16,26} After being activated and etched by molten KOH at high temperature, the tubular PFC are damaged to different extents, as for a-PFC activated with alkali-to-PFC ratio of 3, the long tubes are broken into segments with length shorter than 100 µm, the orifices at the tip clearly evidences the tubular shape. The short tube is beneficial for easier diffusion of electrolyte ions or organics into the inner surface with short channel length, hence the improved capacitive performance and adsorption capability are favorable.\textsuperscript{27,28} TEM images (Fig. 1b and Fig. 1e) more clearly manifest the morphological change by alkali activation. PFC demonstrates 1-dimensional shape, the overall tube is dark, which is mainly due to the large diameter and the thick tube wall, electron beams can not penetrate the tube and results in the seemingly rod-like shape (Fig. 1b). After alkali activation (Fig. 1e), the enhanced transparency of the tube 70 indicates the much hollowed interior of tube wall, which validates that KOH can etch PFC and create a substantial amount of mesopores and micropores that are homogeneously distributed throughout the tube wall. More detailed observation reveals that the tube wall is composed of 1-dimensionally arranged mesopores (narrower than 50 nm) along the longitude direction, indicates that a-PFC\textsubscript{3} macrotube is actually composed of many primary nanotubes embedded in the tube wall. HRTEM image (Fig. 1e) of a PFC fragment shows alternative dark and light spots, revealing the presence of high density micropores in the tube wall, the sinuous fringes at the rim can also be easily observed, which is typical for carbon materials, indicating the graphitization of PFC. After alkali activation, the TEM still shows rich micropores (Fig. 1f). The wave shaped and incontinuous fringes (Inset) evidence that the high density of micropores destroy the continuity of graphitization regions throughout a-PFC. In this unique porous network with mesopores and micropores embedded into the tube wall of a-PFC, macropores and mesopores serve as main channel for electrolyte diffusion, the shorter length of a-PFC, by activation accelerate this process due to the short geometric distance, whereas the micropores provide major contribution to a high specific surface area, therefore electrolyte and organic molecules can readily infiltrate into the deep inner voids with huge interface area. Hence, a-PFC is expected to be an efficient porous material in surface dependent applications e.g. electrode materials of capacitor and adsorbent for water treatment. Controlled samples with different KOH-to-PFC ratios are also characterized, a-PFC\textsubscript{2} shows longer macrotubes, while the length of a-PFC\textsubscript{4}, meanwhile, more severe fragmentization occurs (Fig. S1), indicates the higher KOH dosage not only etches the carbon framework of PFC, but also causes the fracture and fragmentation of the carbon tubes. According to these results, the morphology, surface area, and therefore the application performances of a-PFC can be tuned by altering KOH activation dosage.

Fig. 2a presents the XRD patterns of PFC and a-PFCs, two broad diffraction bands can be observed at 24° and 43° for PFC, which correspond to (002) and (110) facets of graphite-type carbon, the low intensity and broadness of the diffraction bands indicate the low crystallinity of PFC. After activation, the (002) diffraction band shift to lower angle, which is presumably due to the enlargement in interlayer distance of adjacent graphitized carbon layer by the introduced oxygen functional groups as spacer. In contrast, the intensity of (110) facet increases, which means higher integrity of in-plane graphitic carbon skeleton. From this phenomenon, the graphitization degree is actually increased by alkali activation at a higher temperature, although a high density of pores are created in carbon framework. As a whole, both of the diffraction peaks are still low, suggesting the largely amorphous nature the samples. Moreover, the high baseline in the low angle region for a-PFCs probably derives from the high density of micropores in carbon framework.
Fig. 2b shows the FTIR spectra of PFC and a-PFCs, the vibration peaks at 3420 cm⁻¹ (stretching vibration mode) and 45
1635 cm⁻¹ (bending vibration mode) indicate the presence of -OH groups in PFC and a-PFCs, the enhancement in intensity of both
peaks for a-PFCs evidences that the activation treatment mainly
produces plenty of hydroxyl groups in carbon framework. The vibration at 1726 cm⁻¹ and 1680 cm⁻¹ are attributed to -C=O in 50
carboxyl and carbonyl motifs. All of these vibrations manifest the
presence of hydroxyl, carbonyl and carboxyl in all the samples,
thus offer essential hydrophilicity and polarity, allowing the
accessibility of active surface to electrolyte ions and organics,
which is also an indispensable factor for application as capacitive 55
material and adsorbent for polar organics.

The surface chemical composition of PFC and a-PFCs are revealed by XPS analysis. From Fig. 2c, the survey spectrum of
PFC (black curve) shows three peaks with binding energy at
284.8, 532.4 and 400.2 eV, which are characteristics of C 1s, O 1s 60
and N 1s orbital, respectively, all these peaks indicate the surface
composition of PFC comprises carbon, oxygen and nitrogen
elemental. After alkali activation, the peak belonging to N 1s
is not observed in a-PFC3 (red curve), suggesting the elimination of nitrogen elemental by activation treatment. Compared with PFC, 65
the enhanced intensity of O 1s peak relative to C 1s in a-PFC3 indicates the introduction of oxygen functional groups by alkali
activation. The deconvoluted C 1s peak of a-PFC3 shows the
presence of C-C bond in graphite domains (284.8 eV), C-OH
(285.7 eV) and C=O (286.5 eV) groups (inset), further verifies 70
the partial graphitization of carbon framework and the presence of hydroxyl and carbonyl groups on a-PFC3, which basically coincides with the XRD and FTIR data.

The pore properties of PFC and a-PFCs were analyzed by N₂ sorption isothermal measurements. As shown in Fig. 2d, both of 75
the two samples demonstrate type I (Langmuir) isotherm coupled with a faint hysteresis loop characteristic of type IV isotherm
according to IUPAC classification. Micropore filling occurs and
quickly reaches to a saturated plateau at relative pressure lower
than 0.15. The pronounced adsorption in this range evidences the 80
presence of overwhelming micropores in both samples. These
micropores are presumably derived from the intrinsic voids by
evaporation of less stable substances inside PF during pyrolysis
and activation treatment, which agree well with the TEM and
XRD results. At relative pressure of 0.4-1, the low nitrogen
adsorption amount and a weak but discernable hysteresis loop
verify that only limited fraction of mesopores exists in both
samples. The approximately H1 type hysteresis loop characterized by the parallel trend between adsorption and
desorption branches indicates the slit-like geometry of mesopores,
which associates with the embedded mesopores or capillary in
tube wall of a-PFC3 (Fig. 1e). The slight increment in adsorption
amount at relative pressure approximate to 1 suggests the
presence of a tiny fraction of macropores, which may be ascribed to the large internal diameter of PFC (or a-PFC) tubes. BET
surface area and overall pore volume of a-PFC3 are 1215 m² g⁻¹
(1139 m² g⁻¹ for micropores) and 0.65 cm³ g⁻¹ (0.58 m³ g⁻¹ for
micropores) (Table S1). Both parameters are five times higher than the counterparts of PFC: BET surface area of 243 m² g⁻¹
(216 m² g⁻¹ for micropores) and pore volume of 0.12 cm³ g⁻¹
(0.09 cm³ g⁻¹ for micropores). Relative to PFC, micropores volumetric ratio of a-PFC3 increases from 75 % to 88 %, manifesting the creation of more micropores by KOH activation, which may be due to the etching of carbon framework or the opening of dead pores in PFC by alkali activation. Moreover, HK pore size distribution calculated from adsorption branch shows enhanced probability below 2 nm, also verifies the increased micropores in a-PFC3. No discernable pore size distribution over
2 nm can be observed, also confirms the much lower fraction of mesopores and macropores in a-PFC3. The high BET surface area and versatile pore feature guarantee a highly accessible surface to electrolyte ions and organic molecules, facilitating the adsorption of more electrolyte ions and organics onto the entire surface of a-PFC3, which will further result in a high electric-double-layer (EDL) capacitance and adsorption capacity for organics. KOH activation dosage dependent BET surface area and pore volume are also investigated (Fig.S2 and table S1), the specific surface areas of a-PFC2 and a-PFC3 are 954 and 1513 m² g⁻¹, respectively, indicates the higher alkali dosage is really beneficial for a high BET surface. Whereas the micropore volume ratio slightly decreases from a-PFC2 to a-PFC3, and the probability at larger

Fig. 3 a) CVs of PFC and a-PFC3 electrodes at scan rate of 100 mV s⁻¹ in 1 M H₂SO₄ electrolyte. b) CVs of a-PFC3 electrode at different scan rates. c)
Galvanostatic charge-discharge curves of symmetric capacitor in two-electrode mode based on a-PFC3 at different current densities, dashline: PFC capacitor at 1 A g⁻¹. d) Specific capacitances of PFC and a-PFC3 based capacitors at different current densities. e) Cycling stabilities of PFC and a-PFC3 based capacitors at 1 A g⁻¹. f) Nyquist plots of PFC and a-PFC3 electrodes, inset: magnified impedance at high frequency region.
pore size elevates simultaneously, both of these tendencies suggest that low dosage of KOH and the related corrosive species mainly intercalate into carbon layers and etch the carbon skeleton to create predominant micropores in a-PFC, if the KOH dosage is increased, the excessive corrosive species can further etch the rim of micropores and transform some of them into meso- or macropores. From these results, the specific surface area, pore volume and pore size can be roughly tuned by altering alkali dosage.

**Electrochemical properties**

The porous characteristics of a-PFCs with high surface area and partial graphitization make them good candidates as electrode materials for supercapacitor. To evaluate the capacitive performance of a-PFC, CVs of PFC and a-PFC electrodes were measured at first in three-electrode system within potential range of 0–1 V at a scan rate of 100 mV s⁻¹. From Fig. 3a, both electrodes exhibit quasi-rectangular loop without obvious redox peaks, characteristic of typical EDL capacitive behavior. The dramatically higher plateau current and loop area of a-PFC relative to PFC indicates the much higher capacitance of a-PFC, which is mainly attributed to the dramatic increment in BET surface area and therefore larger electrode/electrolyte interface area for ions accumulation. Additionally, the increased oxygen functional groups by activation enhance the wettability of a-PFC, to aqueous electrolyte, which also favors the readily infiltration and access to the deep inner voids, improving the surface utilization ratio of active material.

Fig. 3b presents the CVs of a-PFC electrode at different scan rates. It is obvious that the plateau current increases accordingly with scan rate, and the quasi-rectangular loop can be largely retained without apparent distortion even at higher scan rate, evidencing the small inner resistance and fast electrolyte diffusion kinetics even at high scan rates, which further contributes to a high rate capability.

Fig. 3c shows the galvanostatic charging-discharging curves of symmetric capacitors based on a-PFC at various current densities in double-electrode mode. The approximately triangular charge-discharge branches with good symmetry and nearly linear discharging trend is indicative of typical EDL capacitance with rapid charging-discharging process. No discernable voltage drop can be observed at the beginning of the discharge stage even at current density of 5 A g⁻¹, validating the low intrinsic resistance of a-PFC, which may be attributed to the well conductivity of partially graphitized a-PFC after high temperature carbonization and activation treatment, as well as the rapid electrolyte diffusion kinetics owing to the versatile porous network with the coexistence of micro-, meso- and macropores. The discharging duration of a-PFC at 1 A g⁻¹ (red curve) is apparently longer than that of PFC (black dashline), implies the advantageous Cₑ of the former. From the discharging branch, Cₛ of a-PFC cell according to Eq. 1 are calculated to be 223, 216, 201 and 194 F g⁻¹, respectively at current density of 0.5, 1, 2, and 5 A g⁻¹, all of which are drastically higher than that of PFC based cell (Fig. 3d). It should be noted that the decrement in Cₑ at higher current density for a-PFC is mainly caused by the insufficient diffusion of electrolyte into deep micropores at high current density. Albeit this, a 87% Cₑ retention can still be achieved in the range of 0.5–5 A g⁻¹ (Fig. 3d), revealing the high rate capability of a-PFC. We attribute to the high EDL capacitance and the rate capability to the unique electrolyte diffusion channel with the embedding of prominent micropores and small fraction of mesopores into the tube wall of a-PFC, which enables efficient infiltration and transfer of electrolyte ions into the deep inner void of porous electrode, maximizing the charge accumulation. Additionally, the partial graphitization of a-PFC benefits the rapid charge transfer during charging-discharging process. Both of these factors are essential for a high EDL capacitive performance.

Cycling performance of PFC and a-PFC based capacitor were evaluated by consecutive galvanostatic charging-discharging measurements at 1 A g⁻¹ (Fig. 3e). Cₛ of a-PFC capacitor retains 91.2% of its initial value after 2000 cycles, indicating a high degree of reversibility in the repetitive charging-discharging cycles. The high cycling performance can be attributed to the versatile pores and the well structural stability of a-PFC. The macropores and mesopores, although in limited fraction, benefit the shuttling of electrolytes, alleviating the over-accumulation of electrolyte in micropores and therefore the volumetric variation during charging-discharging cycles. Besides, the structural stability and chemical inertness of a-PFC also help enduring the successive volumetric changes free of damage, both of which contribute to the high cycling stability. Although PFC possesses better cycling stability, the much lower Cₑ relative to a-PFC limits its practical significance in capacitor applications.

To further understand the electrochemical behavior of a-PFC, EIS of PFC and a-PFC electrodes were measured and compared, the Nyquist plots of both electrodes show similar shape, which comprise an unconspicuous semicircle in high-frequency region and a low-frequency straight line (Fig. 3f). The intercept at higher frequency end in real axis represents the series resistance (Rₛ), which includes the bulk electrolyte resistance, intrinsic active material resistance and the contact resistance between electrode and collector. At lower frequencies, no distinct Warburg impedance can be observed for both samples (the line with slope near 45°), which suggests the versatile pore network facilitates the rapid electrolyte diffusion kinetics. The more vertical straight line of a-PFC at low frequency is indicative of a better EDL capacitive behavior with rapid ion diffusion kinetics. In short, by alkali activation at a higher temperature, lower functional groups owing to low alkali dosage during activation, a substantial improvement in capacitive performance.

The effect of PFC-to-KOH activation ratio on the capacitive performance was also investigated (Fig. S3). As for a-PFC, due to longer tubes with lower BET surface area (Fig. S1 and Fig. S2), as well as the possibly smaller amount of hydrophilic oxygen functional groups owing to low alkali dosage during activation, a lower Cₑ of 202 A g⁻¹ is obtained at 1 A g⁻¹. Cₛ of a-PFC based capacitor (206 A g⁻¹) is elevated yet still lower than that of a-PFC based cell, which is presumably due to the over hollowed texture by higher alkali activation dosage. The too high pore volume limits the conductive channel of porous electrode, which can be evidenced by the higher Rₛ, whereas lower Rₛ in impedance plots (green plots in Fig. S3d). In this sense, too high
or too low surface area is not necessarily beneficial for capacitance, an appropriate alkali dosage is essential to balance the contributions from conductivity and BET surface area. In a-PFC, the balanced factors result in a maximized capacitance. The rate performances of a-PFC$_2$ and a-PFC$_4$ are both slightly lower than a-PFC$_3$, further evidences the significance of alkali activation dosage on capacitive performance. Ragone plots shows that a-PFC$_3$ based capacitor offers a high energy density 30.4 Wh kg$^{-1}$ at power density 970 W kg$^{-1}$ (Fig. S3e), both parameters are apparently higher than that of a-PFC$_2$ and a-PFC$_4$. The ultrahigh energy density herein is mainly attributed to the high $C_s$ at low current density in aqueous electrolyte and the wide voltage window during discharging process. Hence, in our case, a-PFC$_3$ is a preferential electrode material with high capacitive performance.

20 Adsorption properties

Because of the high specific surface area and the porous feature with rich oxygen functional groups on the surface, a-PFC is thought to be an efficient adsorbent for organic dyes in aquatic environmental. Herein, the dye uptake capability of a-PFC$_3$ adsorbent was evaluated using MB, MO (methyl orange), NG (naphthol green), CR (congo red) and RhB (Rhodamine B) dyes as model adsorbates. The dye uptake capacity of a-PFC$_3$ can be easily estimated via monitoring the variation in absorbance. When 10 mg of a-PFC$_3$ was dispersed in 180 mL of different dye solutions (each is 20 mg L$^{-1}$), the adsorptive decoloration of dyes occurred to different extents within 24 h (Fig. S4). As shown in Fig. 4, the uptake ratios of a-PFC$_3$ toward dyes differ apparently, almost 100% of MB is removed from solution, which is higher than other dyes to different extents, revealing the well adsorption ability toward MB. The high MB adsorption ability may be attributed to the strong $\pi-\pi$ stacking interaction between aromatic rings in MB molecules and the partially graphitized carbon skeleton of a-PFC$_3$, the anion-cation attraction between the positive charge of MB and the oxygen functional groups on a-PFC$_3$, hydrogen bonds, relatively lower steric hindrance of MB owing to the linear aligned aromatic rings, as well as the structural merits such as high BET surface area and versatile pore feature of a-PFC$_3$.

Due to the high uptake ratio for MB, the adsorption capacity for MB was evaluated. $Q_{eq}$ (mg g$^{-1}$) for MB can be calculated based on the following mass conservation equation:$^{33}$

$$Q_{eq} = \frac{(C_0 - C_{eq})V}{m}$$  \hspace{1cm} (Eq. 4)

Where $C_0$ and $C_{eq}$ are the initial and equilibrium concentrations of MB (mg L$^{-1}$) in solution, $V$ is the volume of MB solution (L), and $m$ is the mass of adsorbent (g). Fig. 5a shows the plots of adsorption amount as a function of time in MB solution (20 mg L$^{-1}$, 350 mL), $Q_{eq}$ for MB is estimated to be 550 mg g$^{-1}$. The value of $Q_{eq}$ is higher over or comparable with the previous reported porous carbon materials,$^{11,34-37}$ revealing that a-PFC$_3$ is a competitive adsorbent for MB uptake. However, our value is still lower than that of other porous carbon or graphene due to the relatively lower surface area or the limited fraction of mesopores.$^{38-40}$

The adsorption kinetics for MB was investigated by monitoring the decoloration rate relative to its initial absorbance. From Fig. 5a, it is clear that the adsorption rate is rather high in the initial 2 h and then gradually slows down and reaches a saturation platform within 24 h. The long adsorption duration herein is mainly due to the much higher volume of MB solution (350 mL) relative to a-PFC$_3$ dosage (10 mg), which restrains the diffusion and sufficient contact to a-PFC$_3$. The adsorption kinetic data was analyzed using a pseudo-second order kinetic model which is based on the assumption that chemisorption is the rate determining step. The pseudo-second order kinetic model can be described as follows:

$$t/Q_{eq} = 1/k_2 Q_{eq}^2 + t/Q_{eq}$$ \hspace{1cm} (Eq. 5)

Where $Q_{eq}$ (mg g$^{-1}$) is the MB adsorption amounts at arbitrary time $t$, $k_2$ is the pseudo-second-order rate constant (g mg$^{-1}$ min$^{-1}$). The plots of $t/Q_{eq}$ versus $t$ are shown in Fig. 5b, the plots strictly obey the fitted straight line with a correlation coefficient of 0.9986, which is apparently higher than that fitted by pseudo-first-order kinetics equation (correlation coefficient: 0.9150, Fig. S5), verifies that the adsorption behavior follows more by pseudo-second-order kinetics, and $k_2$ is calculated to be $5.79 \times 10^9$ g mg$^{-1}$ min$^{-1}$. According to this pseudo-second-order kinetics, the adsorption rate is dependent on both concentrations of absorbent and absorbate, a higher a-PFC$_3$ concentration in the dispersion should accelerate this chemisorption process and therefore shortens the adsorption duration.

Adsorption isotherm can be employed to elucidate the interactions between adsorbates and absorbent. Fig. 5c shows the adsorption isotherm of MB on a-PFC$_3$, $Q_{eq}$ increased accordingly with $C_{eq}$. The experimental data was fitted by Langmuir isotherm, which assumes the homogeneous monolayer chemisorption of adsorbates on identical sites and the uniform adsorption energy

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![Fig. 4](image-url) Uptake ratios of different dyes by a-PFC$_3$.

![Fig. 5](image-url) a) Time dependent adsorption amount of MB on a-PFC$_3$. b) Pseudo-second-order adsorption kinetics. c) Adsorption isotherm of MB on a-PFC$_3$ at 20 °C. d) Plots of $C_0/Q_{eq}$ against $C_{eq}$ based on Langmuir isotherm model.
due to the high affinity between adsorbate and adsorbent, the interactions between adsorbate molecules can be neglected. The Langmuir isotherm can be expressed as follows:

$$Q_{eq} = \frac{Q_{max} \cdot K_{L} \cdot C_{eq}}{1 + Q_{max} \cdot K_{L}}$$  

(6.2)

Where $C_{eq}$ (mg L$^{-1}$) and $Q_{eq}$ (mg g$^{-1}$) are the equilibrium concentration and adsorption capacity of MB, respectively, $Q_{max}$ (mg g$^{-1}$) is the maximized MB adsorption amount, and $K_{L}$ (L mg$^{-1}$) is the Langmuir adsorption constant. Fig. 5c displays the plots of $Q_{eq}$ vs $C_{eq}$ at different initial concentrations, showing high adsorption capacity for uptake of MB in aqueous solution. The well capacitive and adsorptive performances manifest the huge potential of a3PFC as energy and environmental materials.

In summary, a3PFC exhibits well linear relationship with correlation coefficient of 0.9831 (Fig. 5d), the value is higher than that fitted by Freundlich adsorption mode in our case follows more by Langmuir isotherm (correlation coefficient: 0.8699, Fig. S6), indicating the desired capacitive performances including high specific capacitance, rate capability and cycling stability when employed due to the high affinity between adsorbate and adsorbent, the interactions between adsorbate molecules can be neglected. The Langmuir isotherm can be expressed as follows:

$$Q_{eq} = \frac{Q_{max} \cdot K_{L} \cdot C_{eq}}{1 + Q_{max} \cdot K_{L}}$$  

(6.2)

Where $C_{eq}$ (mg L$^{-1}$) and $Q_{eq}$ (mg g$^{-1}$) are the equilibrium concentration and adsorption capacity of MB, respectively, $Q_{max}$ (mg g$^{-1}$) is the maximized MB adsorption amount, and $K_{L}$ (L mg$^{-1}$) is the Langmuir adsorption constant. Fig. 5c displays the plots of $Q_{eq}$ vs $C_{eq}$ at different initial concentrations, showing high adsorption capacity for uptake of MB in aqueous solution. The well capacitive and adsorptive performances manifest the huge potential of a3PFC as energy and environmental materials.

Conclusions

In summary, a-pFCC was facilely synthesized by carbonization and alkaline activation of PF. The as-prepared product demonstrated highly porous network with a large fraction of micropores, high BET surface area and partial graphitization, all of which lead to 90% desired capacitive performances including high specific capacitance, rate capability and cycling stability when employed as electrode material for supercapacitor. Moreover, a-pFCC also showed high adsorption capacity for uptake of MB in aqueous solution. The well capacitive and adsorptive performances manifest the huge potential of a-pFCC as energy and environmental materials.

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Notes and references

1. School of Chemistry and Chemical Engineering, Henan Normal University, Xinxiang 453007, P.R. China. E-mail address: zyguo512@163.com (Z. Guo); kjiang512@163.com (K. Jiang).

2. Collaborative Innovation Center of Motive Power & Key Materials, Henan, 453007, P.R. China.


