Physical and Electrochemical Characterization of Palm Kernel Shell Biochar (PKSB) as Supercapacitor

Wan Azlina Wan Ab Karim Ghani1, Praveen Shawn Fernandez1, Mohamad Qayyum Halele1, Shafreeza Sobri1, Jasronita jasn2

1Department of Chemical and environmental Engineering, Faculty of Engineering, Universiti Putra Malaysia, Serdang, Selangor, MALAYSIA
2Department of Electrical and Electronic Engineering, Faculty of Engineering, Universiti Putra Malaysia, Serdang, Selangor, MALAYSIA

Abstract. A potential low cost and environmentally friendly supercapacitor has been prepared from Palm Kernel Shell Biochar (PKSB). In this study, physical and electrochemical properties of raw, activated and chemical treated (potassium hydroxide (KOH)) as supercapacitors such as high carbon content, high charge storage capacity and stable were evaluated. For physical analyses, the scanning electron microscopy (SEM) was used to study the surface morphology and surface area and porosity were measured using Brunauer-Emmert-Teller (BET). The chemical treated PKSB shows the highest surface area values of 55.15 m²/g as compared to raw and activated samples with surface area are 0.17 m²/g and 19.32 m²/g, respectively. This is verified by in enhancement of capacitance achieved from 1.76 x10⁻³ Fg⁻¹ for the activated biochar and 1.87x10⁻⁶ Fg⁻¹ for untreated PKSB showed by Raman spectroscopy. This enhancement reflected the charge storage capacity is attributed to the creation of broad distribution in pore size and a larger surface area. In addition, this phenomenon also supported by the electrochemical profiles through cyclic voltammogram (CV) measured by Potentiostat-Gavanostat (EIS). CV of the treated PKSB gave better square shape than the activated and raw biochar samples. These characterizations conclude that the raw palm kernel biochar need further treatment to become supercapacitor electrodes to replace activated carbon.

1 Introduction

In response to the changing global landscape, energy has become a primary focus of the major world powers and scientific community. There has been great interest in developing and refining more efficient energy storage devices. One such device, the supercapacitor, has matured significantly over the last decade and emerged with the potential to facilitate major advances in energy storage.

Supercapacitors, also known as ultracapacitors are electrical energy storage devices. Compare to other electrical energy storage devices such as conventional capacitors, batteries, and fuel cells, supercapacitors can store high capacity electrical energy at respectively high power density [1, 2, 3] as illustrated in Figure 1. The charge-storage process for supercapacitor is highly reversible, resulting in longer cycle-life compare to other power storage devices [3]. Advanced carbon materials, such as graphene, carbon nanotubes and carbon onions from metal carbide have shown high capacitance and high power density in electrochemical supercapacitors and hybrid supercapacitors. However, those advanced carbon materials will be significantly limited by high cost [4]. In addition, most conventional synthesis methods for these innovative carbon materials involve application of toxic strong oxidants, such as chlorine, potassium permanganate and hydrazine [5]. Therefore, hierarchical carbon materials, prepared from sustainable precursors with scalable processes, would be of great benefit to large scale application of supercapacitor in energy storage and conservation.

![Ragone plot of energy storage devices](image_url)

Figure 1. Ragone plot of energy storage devices [6]
2 Materials and Methodologies

For this study, three forms of Biochar samples for each organic material are prepared, namely; control BC (raw) (not activated and not treated), activated BC (activated but not treated), and treated and activated BC (activated and treated). Biochar was prepared through pyrolysis of a PKS in a Nitrogen gas atmosphere by applying a temperature program to a MTI GSL-1100X tube furnace. The temperature was increased from room temperature to 200°C and then held at this temperature for 1 h to remove moisture. This was followed by increasing temperature to 750°C at around 3°C/min and dwelling at this temperature for 1 h to remove volatile organics and to carbonize solid residues. The resulting Biochar chunk was removed from the furnace for characterization and evaluation after the temperature was cooled down to room temperature. The BC chunk was cut into small portions. Those with weights of approximately 1 mg were directly used as supercapacitor electrodes without introducing any organic binder.

2.1 Activation of biochar electrodes

Biochar electrodes were further activated by immersing them into Potassium Hydroxide (KOH) solution at room temperature overnight to increase surface hydrophilicity and the coverage of surface oxy-gen groups. They were rinsed using deionized before use. The activation process was carried out for three hours at 950 °C in a muffle furnace. The rate at which the muffle furnace heating is set is 5°C/min. Once the activation was completed, the activated BC was cooled within the muffle furnace in the same Nitrogen gas environment until the temperature drops to room temperature, which takes approximately nine hours. Then, the activated biochar sample is rinsed with 0.1mol/L hydrochloric acid (HCl) at 100 °C using a spiral condenser. Once that is done, the activated BC is again rinsed this time with deionized water to pH 7 (neutral). The reason for using deionized water specifically is to avoid contact of external ions on the surface of the activated BC which might sabotage the experimental results. Lastly, the activated BC sample is dried at 105 °C for the overnight under vacuum.

2.2 Treatment of BC electrodes

The activated BC samples were then further modified with 4M HNO3. This is to facilitate for the third control sample, which is the activated and treated BC sample. The activated BC samples were left to soak in 20mL of 4 mol/dm3HNO3 solution sealed in Polytetrafluoroethylene (PTFE) reactor for 48 hours at 150 °C. Once soaking was done, the treated BC was then cooled and filtered. The BC electrodes were rinsed with deionized water until the filter liquid equalled pH 7. Lastly, the filter cake of treated BC was collected and left to dry overnight in a conventional oven at temperature 110 °C.

2.3 Physical characterization

Surface morphology and elemental composition of the electrodes were characterized using a field-emission environmental scanning electron microscope operating at 15 keV equipped with a light-element energy-dispersive spectroscope. BET surface areas of BC samples were measured using a Machine (Micromeritics Gemini VII 2390 V1.03) surface area/pore volume analyzer. Raman measurements were performed using a high-resolution research-grade (Horiba LabRAM HR 3D-capable) Raman spectroscopy imaging system that is optimized for the visible-to-NIR spectral range. A 532 nm wavelength laser was used in this study. DC electrical conductivity of BC was determined from Ohmic resistance, cross-section area, length and porosity of a small Palm Kernel Shell chunk based on Pouillet’s law. The Ohmic resistance corresponds to the slope of linear potential–current curve measured by applying potential scan to two ends of the BC electrode chunk.

2.4 Electrochemical measurements

Voltammetric and constant current charge–discharge studies of a single BC electrode were carried out in 0.5 mol solution with a coil of 1 mm diameter Platinum wire and a silver/silver chloride (Ag/AgCl) as the counter electrode and the reference electrode, respectively. The
electrical contact of a BC electrode was obtained by wrapping fine gold wire around the PKS portion to minimize contact resistance. The geometric surface area is around 0.08 with 2 mm in width and 4 mm in length. The capacitive behavior of a BC supercapacitor constructed by putting two BC electrodes in similar size separated by a sheet of glassy filter-paper in 0.5 mol was evaluated by cyclic voltammetry and chronopotentiometry. Their preliminary durability was evaluated via successive potential cycling.

3 Results and Discussion

3.1 Physical characterization of Palm Kernell Shell biochar.

Figure 2 shows the result SEM of original PKS biochar, activated PKS biochar and treated PKS biochar, respectively. In these images, light region correspond to carbon wall and black region correspond to pores. The PKS biochar developed high porosity, presenting longitudinal pores with sizes ranging from micro to macro pores (10-200μm) [15]. It is clearly shows that the morphology of treated PKS biochar was more porous compared to original PKS biochar. The porous structures of the treated PKS Biochar sample indicate clearly that the potassium hydroxide, KOH activation process did in fact slice through the pores causing a larger pore size structure to be formed, and also the pore distribution is seen to be more frequent in an orderly manner. The positive results obtained from the difference in surface porosity is clear visible here via SEM thus verifying the reasons for samples to undergo Activation process using potassium hydroxide and Treatment using nitric acid, HNO₃.

Table 1 shows the overall Raman spectrum comparison of PKS Biochar samples. The G-band peaks recorded for Original, Activated, and Treated samples are 1583.14 cm⁻¹, 1578.82 cm⁻¹, and 1582.81 cm⁻¹ respectively. The peaks that appear for the D-band are 1354.39 cm⁻¹, 1354.06 cm⁻¹ and 1354.72 cm⁻¹ for the same samples of Biochar.

The ratio between the D-band and G-band intensities (ID/IG) represents the scale of defect concentration. The higher the intensity ratio, the higher the defect concentration. The defect mentioned here corresponds to the proportion of sp³ carbon formation, which caused disruptions to the original sp² bonding, hence attributing to the structural defects being measure. In this case the values of (ID/IG) for Original, Activated and Treated samples are 0.856, 0.858, and 0.857 respectively. The increase in (ID/IG) after the introduction of nitric acid to treat the samples is due to the hydrothermal oxidation caused by nitric acid. This process generates oxygen containing functional groups on the surface of the sample thus increasing the number of sp³ hybridized carbon, causing increased surface defects.

Another obvious change in bands was at band 1362.58 cm⁻¹ which was not present in the Raw biochar, appeared in Activated and Treated biochar although it appears to be shifted slight in the Treated biochar bands at 1243.54 cm⁻¹. This range of bands can be assigned to the C-O stretching and OH bending of alcohol and carboxylic acids.

Figure 2. SEM images of (a) Original PKS Biochar, (b) activated PKS biochar and (c) treated PKS biochar.
In all, similar functional groups were observed on the surface of the activated carbons due to their exposure to similar chemical treatment and narrow variation of temperature of activation and treatment (see Table 2).

**Table 2. Peak Assignments for Raw, Activated and Treated Biochar Samples.**

<table>
<thead>
<tr>
<th>Peak Assignments</th>
<th>Original</th>
<th>Activated</th>
<th>Treated</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aromatic C-H (cm⁻¹)</td>
<td>3015.23</td>
<td>3012.36</td>
<td>3010.93</td>
</tr>
<tr>
<td>Aliphatic C-H (cm⁻¹)</td>
<td>2813.40</td>
<td>2812.74</td>
<td>2815.28</td>
</tr>
<tr>
<td>C=O stretching (cm⁻¹)</td>
<td>1704.86</td>
<td>1713.65</td>
<td>1721.86</td>
</tr>
<tr>
<td>C=C stretching (cm⁻¹)</td>
<td>1590.71</td>
<td>1591.39</td>
<td>1591.16</td>
</tr>
<tr>
<td>O-H stretching (cm⁻¹)</td>
<td>3717.22</td>
<td>3747.85</td>
<td>3648.66</td>
</tr>
<tr>
<td>C-N stretching (cm⁻¹)</td>
<td>1094.38</td>
<td>1018.83</td>
<td>1243.54</td>
</tr>
</tbody>
</table>

The values of BET specific surface area are 0.2982, 12.9034 and 53.8346 for original, activated and treated biochar, respectively. The average pore size of the samples also decreased from 338.19797 to 51.78124 nm after potassium hydroxide, KOH activation. The pore size for Treated biochar increased drastically to 7929.64 nm. This has been verified by the SEM results obtained that clearly shows the larger pore sizes on the biochar after treatment with nitric acid.

**3.2 Cyclic Voltammetry (CV)**

The suitability of the carbon obtained for electrodes in a supercapacitor was evaluated by using an electrochemical technique, cyclic voltammetry (CV). With the prepared Activated biochar as the electrodes, the electrochemical behavior of the biochar was characterized with the CV using 0.5 mol potassium hydroxide as the electrolyte and SCE as reference electrode. The CV response of the electrodes was measured at different scan rates varying from 10 to 200 mV s⁻¹ at potentials between 1 and 0 V. Figure 4 shows CV curve of the biochar samples in 0.5 mol potassium hydroxide.

![Figure 3. FTIR spectrum raw, activated and treated PKS biochars](image)

![Figure 4. CV curve of the biochar samples in 0.5 mol potassium hydroxide](image)
Figure 4. Cyclic Voltammometry (CV) (a) raw, (b) activated and (c) treated PKS biochar electrodes at scanning rate 0.05 mV/s.

From these figures, it is observed that the Activated biochar has a better closer resemblance to a square shape compared to the original biochar CV curve. According to this observation, the activated biochar shows better capacitor characteristics as compared to the original biochar. The treated biochar however exhibits a very scattered CV curve due to the high level of noise. This can be attributed to the uneven rough surface of the treated electrode when tested using CV.

4 Conclusion

Sustainable production of activated carbon was produced from Palm Kernel Shell using potassium hydroxide as the activation agent and nitric acid as the treatment medium, was developed and the results reported. A supercapacitor has been developed using 0.5 mol Potassium hydroxide as the electrolyte. Based on the test results albeit not perfect the palm kernel shell based biochar has the potential to be a lower cost substitute for the advanced, more expensive carbon materials currently being commercialized such as metal carbides, graphene or activated graphene. The process of synthesizing the palm kernel shell biochar is without artificial templates hazardous substances.

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References