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## Processing-properties-performance triad relationship in a *Washingtonia robusta* mesoporous carbon materials-based supercapacitor device†

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Two-electrode electrochemical tests provide a close performance approximation to that of an actual supercapacitor device. This study presents mesoporous carbon materials successfully derived from *Washingtonia robusta* bark (Mexican fan palm) and their electrical performance in a 2-electrode supercapacitor device. The triad relationship among carbon materials "processing, properties, and performance" was comprehensively investigated. X-ray diffraction reveal that amorphousness increases with activating KOH ratio and decreases with both activation time and temperature. Raman spectroscopy shows an increase in structural defects and degree of graphitization with an increase in KOH ratio, temperature and time while transmission electron microscopy shows conversion of aggregated particles to materials with interconnected porosity and subsequent destruction of porosity with an increase in KOH ratio. A nitrogen-sorption study reveals varying trends between BET, micro and mesopore surface areas, however, pore size and volume and hysteresis loop size decreases with KOH ratio and temperature. Electrochemical studies on the other hand reveal that both the specific capacitance and charge-discharge time increase with KOH ratio, temperature and time while both charge transfer and Warburg resistances decrease and the phase angles increases towards the ideal  $-90^\circ$  with an increase in KOH ratio, temperature and time. The device fabricated with the HHPB sample prepared at  $700^\circ\text{C}$ , KOH ratio 3 for 60 min attained a specific capacitance of 179.3 and  $169\text{ F g}^{-1}$  at a scan rate of  $5\text{ mV s}^{-1}$  and current density of  $0.5\text{ A g}^{-1}$ , respectively, good cycling stability with 95% capacitance retention and 100% coulombic efficiency when cycled 5000 times at a current density of  $2\text{ A g}^{-1}$ . HHPB electrodes reveal perfect EDLC behavior with an energy density of  $20\text{ W h kg}^{-1}$  and power density of  $2000\text{ W kg}^{-1}$  when used in a symmetric coin supercapacitor cell with 6 M KOH solution. These findings show the potential of fan palm bark as electrode materials with good stability and high-rate capability for supercapacitor application.

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## Introduction

Supercapacitors have gained considerable attention as a result of the increase in the usage of portable electronic devices. In addition, their unparalleled attributes of longer cycling ability, faster charging, and high power density compared to batteries

has projected them as a superior alternative.<sup>1</sup> Typically there are two main forms of supercapacitors based on the mechanism of charge storage: pseudocapacitors which store charge through faradaic redox reactions, and the electric double layer capacitor (EDLC) which stores charge through electrostatic interactions in the electrical double layer.<sup>1,2</sup> Various materials including conducting polymers, carbon-based materials and metal oxides have been studied as supercapacitor electrode materials.<sup>3</sup> Transition metal oxides are reported to offer high specific capacitance due to faradaic reactions compared to carbon based materials which are based on EDL formation.<sup>2</sup> Although pseudocapacitive based materials have been reported to have high specific capacitance, their life cycling stability is poor thus making carbon based materials the potential materials for supercapacitors. Apart from good life cycle, good conductivity, chemical stability, high specific surface area, and availability have made carbon-based materials such as activated carbon,

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graphene, carbon nanotubes and carbon aerogels promising for EDLC based supercapacitors.

Activated carbon is most preferred carbon material for application in commercial supercapacitors owing to its availability, high specific surface area, and low production cost.<sup>4</sup> However, these materials have poor electronic conductivity, low specific capacitance and low energy density which hinder their application in real life energy storage applications. Thus strategies to synthesize activated carbon from natural or synthetic sources with improved performance are highly needed. It is reported that activated carbon from synthetic sources, such as, polymers, can attain outstanding properties and physical structures for supercapacitors,<sup>5</sup> however, their high production cost, hampers their large scale production for commercialization. Thus carbon materials synthesized from natural sources are mostly preferred for commercial production of activated carbon for various applications due to their outstanding properties and low cost.<sup>1</sup> For many years activated carbons have been produced from coal, wood, petroleum residues, peat, and lignite but these precursors are nonrenewable and unsustainable. Therefore, due to sustainability and renewability credentials, biomass waste has recently been used for large scale production of activated carbon and used as electrode materials.<sup>6</sup> To date, biomass waste such as willow wood,<sup>7</sup> biogas slurry,<sup>8</sup> fish bladder,<sup>9</sup> walnut shells<sup>3</sup> and many other precursors as reviewed by Shanmuga Priya *et al.*,<sup>10</sup> Enock *et al.*,<sup>11</sup> and Lin *et al.*,<sup>12</sup> have been studied as supercapacitor electrode materials.

Additionally, most studied biomass based carbon for supercapacitor utilized three electrode system which is reported to be suitable for studying electrochemical properties of electrode materials rather than the performance of a supercapacitor device. Two-electrode test cell is thus reported to simulates what is actually happening in a real supercapacitor device in terms of the charge transfer, physical configuration and internal voltages and finally demonstrates the performance of the device.<sup>13</sup> The performance of materials in two electrode system still faces a challenge of stability at high current densities. Thus developing appropriate carbon materials with high capacitance and long cycling life for supercapacitor device from renewable and sustainable precursor is significant. This study therefore, utilized porous carbon derived from *Washingtonia robusta* to make supercapacitor electrodes that exhibited high specific capacitance, stable life cycle and good rate capability when tested in two electrode system. Further, the study comprehensively investigated the influence of hydrothermal pre-treatment and activation on the textural, internal structure, morphological, and electrochemical properties and hence performance of the supercapacitor device and revealed an intimate processing-properties-performance triad relationship. *Washingtonia robusta* commonly known as Washington palm, Mexican Washington palm or Mexican fan palm is one among common palms called fan palms. Its fronds has been used for various activities including making furniture and in construction works to replace steel in concrete.<sup>14</sup> It is composed of 46.82, 25.70, 22.46 and 5.02% cellulose, hemicellulose, lignin and extractives, respectively,<sup>14</sup> comparable to that of wood thus suggesting its potential for synthesizing activated carbon for various

applications. To the best of our knowledge there is no literature that reports utilization of carbon derived from dried palm barks for supercapacitor applications. The current study therefore, explored the possibility of synthesizing carbon from dried fan palm barks for supercapacitor applications. It further scrutinized the effect of carbon precursor hydrothermal pretreatment, activating agent ratio, temperature and time on BET, micro/mesopores surface areas, pore size and volume (textural properties); structural defects, degree of graphitization and crystallinity (internal structure properties); particle size and shape (morphology), phase angle, charge transfer and Warburg resistances (electrochemical properties) and on the specific capacitance, cycling stability, coulombic efficiency, capacitance retention, energy and power density (performance) of a 2-electrode supercapacitor device. Utilization of palm bark as carbon precursor addresses environmental pollution by reducing waste and the global energy demand by converting environmentally friendly biomass waste to functional materials with high power and energy density in supercapacitor applications.

## Methodology

### Chemicals

Chemicals including polyvinylidene fluoride (PVDF), Kuray active carbon for supercapacitor electrode (Model YP-50F) both from MTI cooperation, acetone, DMF, H<sub>2</sub>SO<sub>4</sub>, KOH and HCl were used without any purification. Commercial activated charcoal (AC115) from Alpha Chemica was also used.

### Synthesis of carbon

Dried Mexicana palm barks were cut from the garden around Botswana International University of Science and Technology (BIUST) and thorns were removed. Palm barks were cut into small pieces, washed with tap water to remove any adhering dirt and thereafter ashed with distilled water before being oven dried overnight at 100 °C. The dried pieces were then grounded with a compaction module, pulverized and then sieved to the size of 1.27 mm. Six (6) g of the pulverized sample was mixed with 100 mL of 2 M H<sub>2</sub>SO<sub>4</sub> and placed into a 120 mL Teflon lined autoclave, tightened and heated at 180 °C in oven for 24 h. After that, the autoclave was left to cool to room temperature and the resulting sample was filtered and washed several times with distilled water until filtrates became clear. The obtained sample was oven dried overnight at 100 °C, named HHPB and stored in closed containers for further use. One (1) g of HHPB was mixed with KOH pellets at different ratios varying from 1 to 3 and ground thoroughly with an agate mortar. The HHPB and KOH mixture was then activated in MTI OTF-1200X-60 series tube furnace under flow of nitrogen gas at 650, 700 and 750 °C for 60, 90 and 120 min. The heating ramp rate and nitrogen mass flow was set to 10 °C min<sup>-1</sup> and 20 SCCM, respectively. The sample was allowed to cool naturally in the furnace to room temperature under flow of nitrogen gas. The sample was then soaked in 1 M HCl for several hours to neutralize the remaining KOH and also remove ash after which it was washed with distilled water until the effluent pH reach 6.5 to 7. The washed sample was



oven dried at 100 °C overnight and labeled as HHPB<sub>x</sub>-<sub>z</sub> where  $x$  denotes HHPB to KOH ratio, while  $y$  and  $z$  denotes activation temperature and time, respectively. HHPB samples carbonized without activation are hereafter referred to as CHHPB<sub>x</sub>-<sub>y</sub> where  $x$  and  $y$  denote carbonization temperature and time, correspondingly.

### Material characterization

The morphology of the synthesized materials was studied by a JEOL JEM-2100F electron transmission electron microscopy (HR-TEM) operating at 200 kV accelerating voltage and Tescan Vega 3 LMH scanning electron microscope (SEM) while nitrogen sorption studies were used to determine pore size distribution and specific surface area employing Barret-Joyner-Halenda (BJH) desorption isotherm and Brunauer-Emmett-Teller (BET) method, respectively. Raman spectroscopy (Lab-RAM HR800 Raman spectrometer, Horiba JobinYvon, exciting samples with 532 nm Nd-YAG laser) and X-ray powder diffraction (XRD) (Bruker D8 Advance powder diffractometer with a Cu tube X-ray source with  $\lambda = 1.54056$  nm operating at 40 kV, 40 mA and with a LynxEye XE energy-dispersive strip detector) were used to characterize crystal structure changes in carbon materials prepared at different carbonization and activation conditions.

### Electrochemical studies

Cyclic voltammetry (CV), electrochemical impedance spectroscopy (EIS) and cyclic charge and discharge (CCD) was used to study the electrochemical properties/performance of the electrodes at room temperature using Gamry potentiostat/galvanostat reference 3000 in two electrode system. A symmetrical two-electrode supercapacitor system was assembled using EQ-HSTC split-able test cell with Ni foam as current collector and Whatman filter paper as separator. The Whatman filter was soaked in 6 M KOH overnight before being used as a separator. The two working electrodes were prepared by mixing 80% of the synthesized carbon material, 10% of PVDF and 10% Kuray active carbon for supercapacitor in a solution of acetone and DMF (1 : 2 volume ratios) to make viscous slurry which was pasted on nickel foam pieces of 3.14 cm<sup>2</sup>. The electrodes were oven dried at 60 °C overnight to evaporate the solvent. The total mass of active material on single electrode was calculated by taking the mass difference between the nickel foam before and after coating. The total weight of active materials on both electrodes varied from 12–16 mg. EIS measurements were done at open circuit potential with the amplitude of 5 mV at a frequency range of 100 kHz to 10 mHz.

The specific capacitance ( $C_s$ ) in F g<sup>-1</sup> from both CV and CCD for two electrode system was calculated using eqn (1) and (2), respectively.<sup>15,16</sup>

$$C_s = \frac{2}{m \times \nu \times \Delta V} \int_{V_i}^{V_f} I dV \quad (1)$$

$$C_s = \frac{4I\Delta t}{m\Delta V} \quad (2)$$

where  $\int_{V_i}^{V_f} I dV$  symbolizes the area under the CV curve,  $m$  is the total mass (g) of active materials in both electrode,  $\nu$  is the scan rate (V s<sup>-1</sup>),  $\Delta V$  is the potential window (V),  $V_f$  and  $V_i$  (V) are final and initial potential limit of the CV tests,  $I$  is the discharge current (A) and  $\Delta t$  is the discharge time (s).

The energy density ( $E$ ) and power density ( $P$ ) in W h kg<sup>-1</sup> and W kg<sup>-1</sup>, respectively, were calculated using eqn (3) and (4).<sup>16–19</sup>

$$E = \frac{C_s \times \Delta V^2}{2 \times 3.6} \quad (3)$$

$$P = \frac{E}{\Delta t} \times 3600 \quad (4)$$

where,  $C_s$  is the specific capacitance in F g<sup>-1</sup>,  $\Delta V$  is the potential window and  $\Delta t$  is the discharge time in seconds.

## Results and discussion

### Characterization

Fig. 1a–c shows the XRD patterns of HHPB samples at different activation conditions. All samples demonstrate two broad diffraction peaks at about 22 and 44°,  $2\theta$ , corresponding to (002) and (100) graphitic planes, respectively. The broad and asymmetric peaks signify that the carbon samples were turbostratic in nature.<sup>20</sup> The (002) peak intensity was found to decrease with an increase in KOH ratio from 1 to 3 during activation, Fig. 1a. This was probably caused by amorphization of the samples as the KOH etched out inorganics and accelerated volatilization of extractives in the carbon precursor. Contrary, activation temperature and time led to an increase in peaks intensity, particularly (002), Fig. 1b and c. This is indicative of an increase in crystallinity of the samples, albeit within the turbostratic range, caused by carbon microstructure coarsening and crystallites growth during graphitization process at high temperatures.<sup>21</sup>

Fig. 2 represents SEM images of HHPB samples at different activation conditions. It can be seen that heterogeneous morphology composed of microspheres and irregular monoliths after hydrothermal treatment and carbonization at 700 °C for 1 h (Fig. 2a). The microsphere morphology was preserved after activation with KOH ratio 1 at 700 °C for 1 h (Fig. 2b). However, the microspheres disappeared as KOH ratio increased to 2 and 3 (Fig. 2c and d) and faceted particles with more interparticles pores, cavities and more rough surfaces were formed. This might be due to the fact that the microspheres were destroyed by the strong etching effect of increased KOH ratio during activation process as previously reported by Romero-Anaya *et al.*<sup>22</sup> Upon varying activation temperature from 650 to 750 °C, KOH ratio 1, the carbon particles, though still agglomerated and reduced in size, (Fig. S1 in the ESI†). Moreover, TEM was used to study the morphological and structural changes that occurred as the samples were activated at different KOH ratio, temperature, and time. The carbonized sample showed agglomerated particles that are relatively small and thick (Fig. 3a and e) compared to its activated counterparts, Fig. 3b–d and f–h. Upon activation, the agglomerated thick



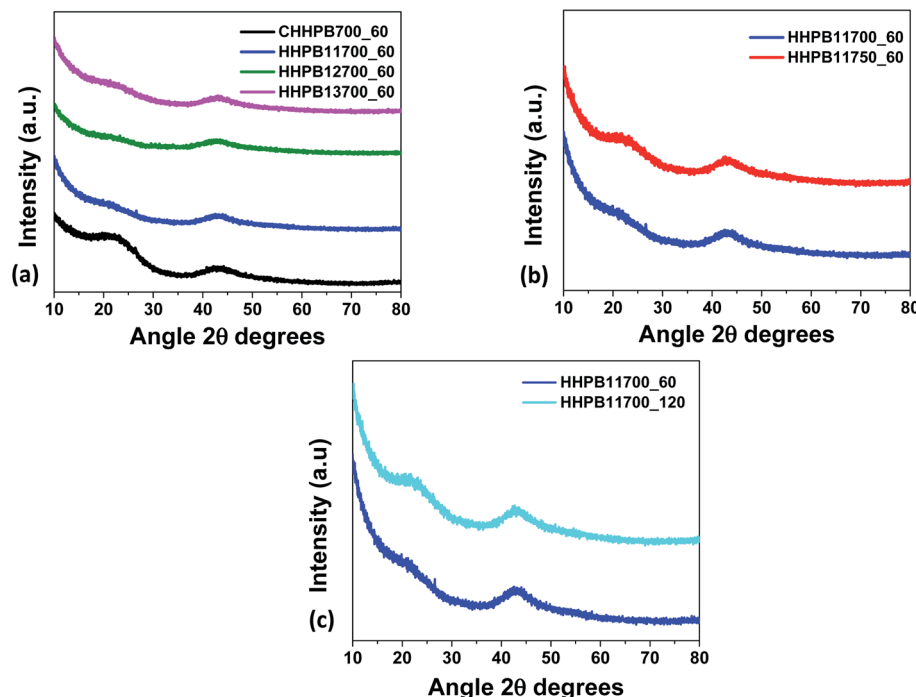


Fig. 1 XRD of HHPB at different activation (a) ratio (b) temperature (c) time.

particles were converted to carbon materials with highly interconnected porosity at KOH ratio of 1, Fig. 3b and f. At this ratio, KOH is suggested to have optimally etched inorganics and

accelerated volatilization of extractibles in the carbon precursor leaving behind fixed carbon with interconnected porous structure. Increasing KOH ratio to 2 significantly reduced the

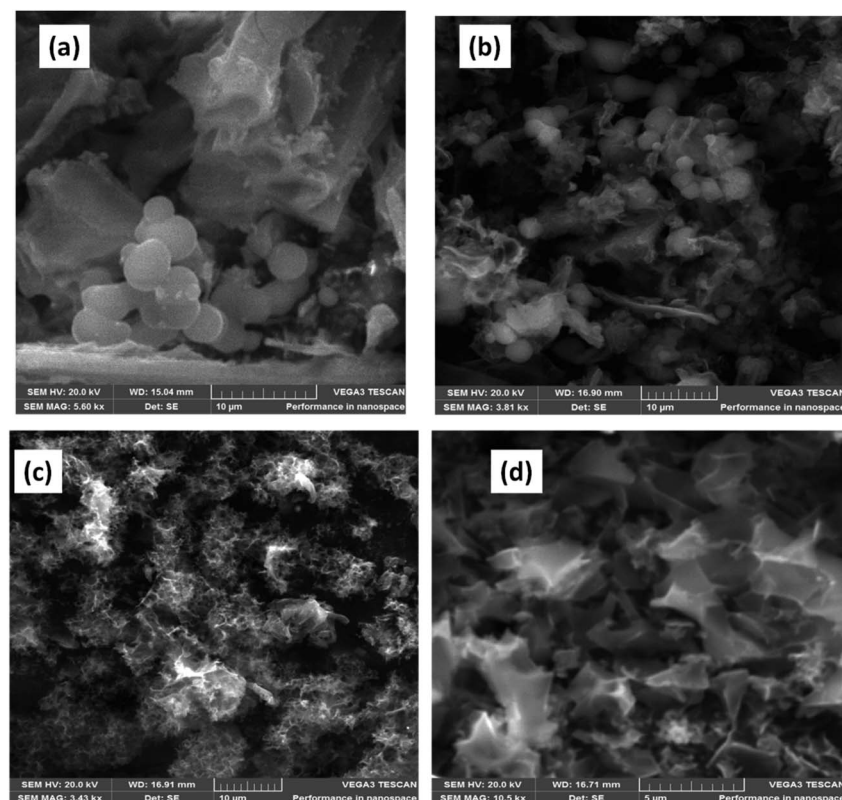


Fig. 2 SEM image of (a) CHHPB700\_60 (b) HHPB11700\_60 (c) HHPB12700\_60 (d) HHPB13700\_60.



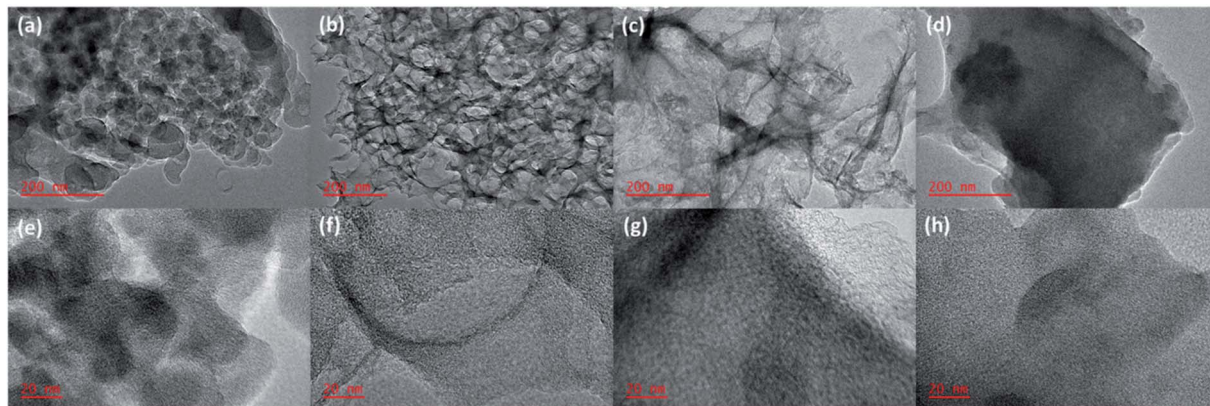


Fig. 3 (a-d) low magnification TEM images of CHHPB700\_60, HHPB11700\_60, HHPB12700\_60, and HHPB13700\_60, respectively, (e-h) high magnification TEM images of (a-d) in that order.

interconnected porous structure leading to formation of relatively large but thinner particles, Fig. 3c and g. At KOH ratio of 3, the interconnected porous structure was completely lost and large relatively thick particles were formed, Fig. 3d and h. The reduction of and complete loss of the interconnected porosity at KOH ratios of 2 and 3, respectively, was probably due to the destruction of walls separating adjacent pores by the KOH. The

TEM results showing highly porous carbon structure at KOH ratio 1 is in harmony with  $N_2$  sorption studies, Fig. 5, which show lack of hysteresis loop for none activated carbon materials (CHHPB700\_60), largest hysteresis loop and a sharp pore size distribution peak centered at around 4 nm with highest pore volume for carbon materials activated at KOH ratio of 1 (HHPB11700\_60). Carbon microstructure coarsening was

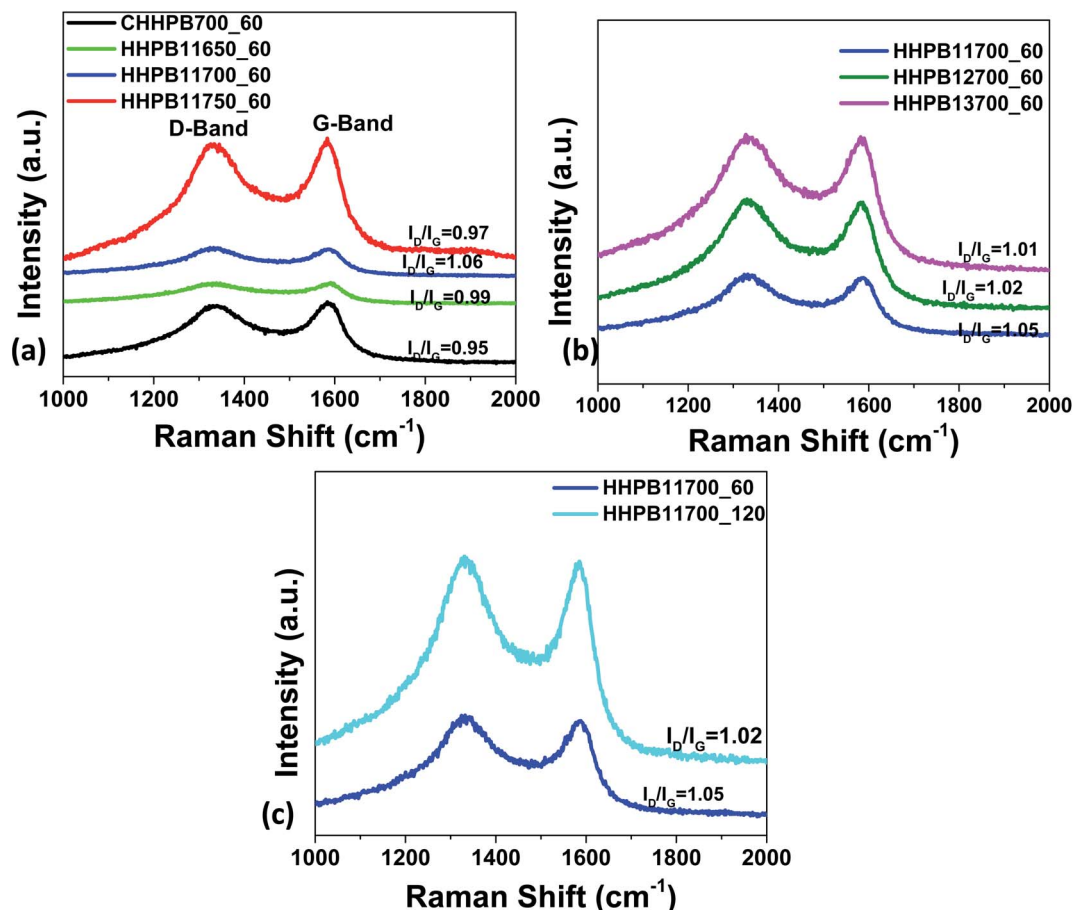


Fig. 4 Raman spectra of HHPB at different activation (a) temperature (b) ratio (c) time.



Table 1 Textural properties of the synthesized HHPB samples

Sample	BET <sup>a</sup> (m <sup>2</sup> g <sup>-1</sup> )	$A_{\text{micro}}^b$ (m <sup>2</sup> g <sup>-1</sup> )	$A_{\text{meso}}^c$ (m <sup>2</sup> g <sup>-1</sup> )	$V_{\text{micro}}^d$ (cm <sup>3</sup> g <sup>-1</sup> )	$V_{\text{meso}}^e$ (cm <sup>3</sup> g <sup>-1</sup> )	$V_{\text{total}}^f$ (cm <sup>3</sup> g <sup>-1</sup> )	Pore size (nm)
<b>Activation ratio</b>							
CHHPB700_60	532.50	218.65	313.85	0.10	0.36	0.46	7.27
HHPB11700_60	1830.61	577.15	1253.46	0.25	0.85	1.09	5.58
HHPB12700_60	1478.11	231.58	1246.54	0.09	0.55	0.64	4.48
HHPB13700_60	1509.79	1340.59	169.20	0.62	0.30	0.91	2.42
<b>Activation temperature</b>							
HHPB11650_60	3572.03	2527.28	1044.75	1.10	1.06	2.16	2.42
HHPB11700_60	1830.61	577.15	1253.46	0.25	0.85	1.09	5.58
HHPB11750_60	2453.56	142.30	2311.26	0.03	1.76	1.79	4.48
<b>Activation time</b>							
HHPB11700_60	1830.61	577.15	1253.46	0.25	0.85	1.09	5.58
HHPB11700_120	1899.31	1612.51	286.80	0.68	0.23	0.91	1.91
HHPB11700_150	1746.26	156.91	1589.35	0.06	0.84	0.90	3.25

<sup>a</sup> BET surface area. <sup>b</sup> Micropore area. <sup>c</sup> Mesopore area. <sup>d</sup> Micropore volume. <sup>e</sup> Mesopore volume. <sup>f</sup> Total pore volume.

observed with increase in activation temperature revealed by formation of small and scattered semi-crystalline domains at 700 and 750 °C, Fig. S2 in the ESI.† This was due to carbon crystallites grow with temperature. These images further supports XRD results, Fig. 1b, that depict turbostratic nature of

carbon samples and an increase in XRD peaks intensity with increase in temperature.

Fig. 4a–c show Raman spectra of different HHPB samples with D peak related to the defect or disorder-induced scattering signal at around 1330 cm<sup>-1</sup> and a G peak assigned to the

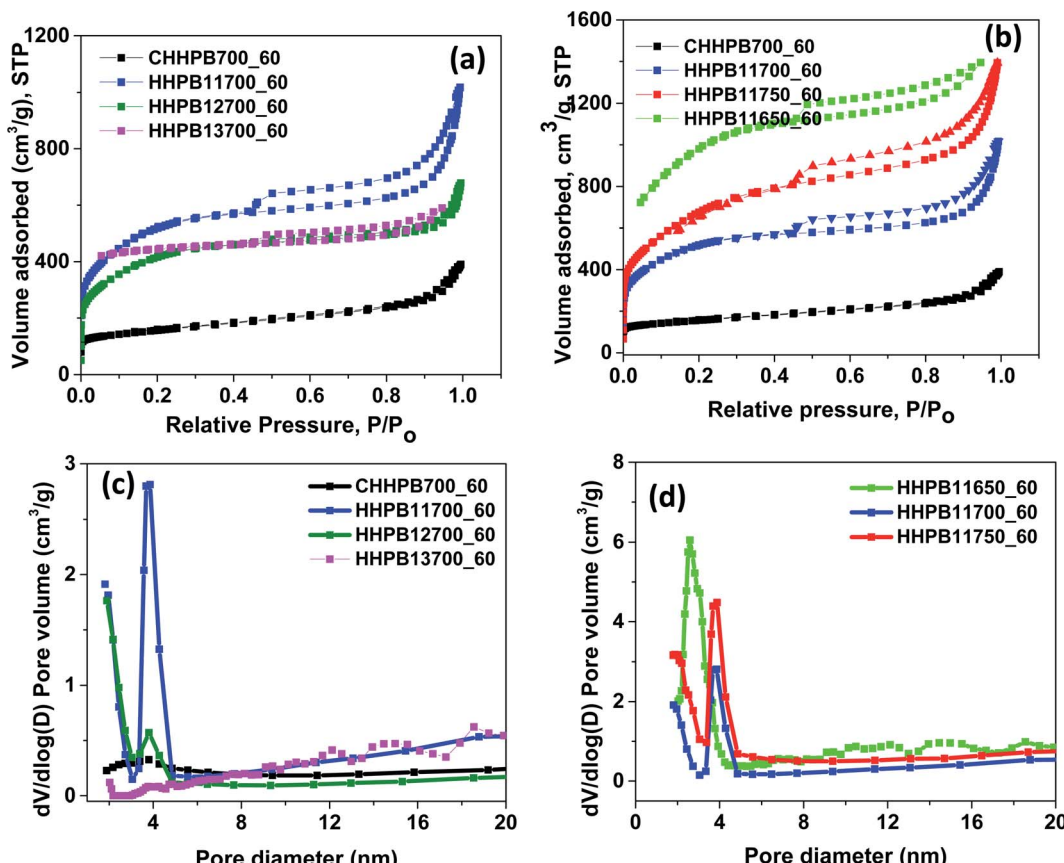


Fig. 5 (a and b) Nitrogen adsorption–desorption isotherms (c and d) pore size distribution of HHPB samples at different activation KOH ratios and temperatures.



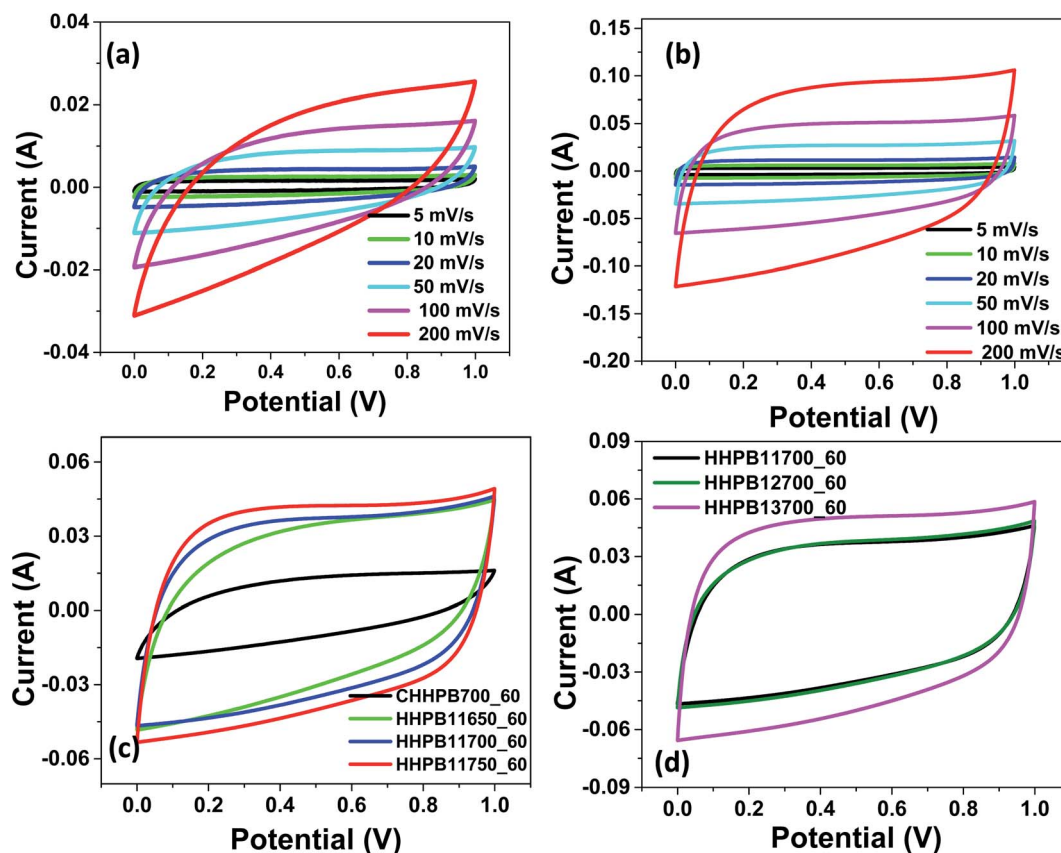


Fig. 6 CV curves of (a) CHHPB700\_60 (b) HHPB13700\_60 at different scan rates, (c) and (d) different electrodes prepared from HHPB samples activated different temperature and KOH ratios, at  $5 \text{ mV s}^{-1}$ .

vibration signal of  $\text{sp}^2$ -bonded carbon atoms at around  $1590 \text{ cm}^{-1}$ . The intensity ratio of the D to G peak ( $I_{\text{D}}/I_{\text{G}}$ ) in carbon materials represents the degree of graphitization and the structural disorder of the carbon framework. The higher the  $I_{\text{D}}/I_{\text{G}}$  value the more the defects in carbon materials. These defects have been shown to favor charge storage in supercapacitors.<sup>23</sup> The values of  $I_{\text{D}}/I_{\text{G}}$  increased from 0.99 to 1.06 as activation temperature increased from 650 to 700 °C. However, the intensity ratio dropped to 0.97 as activation temperature increased to 750 °C (Fig. 4a) as a result of rearrangement in the lamellar structure of the carbon when the sample was treated under high activation temperature. Similarly,  $I_{\text{D}}/I_{\text{G}}$  increased from 0.95 to 1.01 as activating agent (KOH) ratio increased from 0 to 3, Fig. 4b. This was due to the formation of structure defects in the carbon framework as the KOH etched out inorganics and accelerated volatilization of extractables in the carbon precursor and formed  $-\text{OH}$  and  $-\text{COOH}$  functional groups on the fixed carbon framework. These defects in the HHPB samples increased active sites and active surface area and are thus beneficial for the enhancement of the capacitance as previously reported by Hou *et al.*<sup>24</sup> Likewise, activation time was found to have a significant influence on the degree of graphitization and disorder since an increase in activation time led to an increase in the intensity of both the D and G bands, Fig. 4c.

Nitrogen adsorption studies were used to study the BET surface area, pore volume, t-plot micropore area and pore width of HHPB samples and the results are summarized in Table 1. From Table 1, it can be seen that the none activated sample, CHHPB700\_60 showed the lowest BET, micropore, and mesopore surface areas and this could explain its poor electrochemical performance. BET surface area of the activated HHPB samples and total pore volume increased from 532.50 to  $1830.61 \text{ m}^2 \text{ g}^{-1}$  and 0.46 to  $0.91 \text{ cm}^3 \text{ g}^{-1}$ , respectively, after introduction of activating agent due to generation of large amount of pores in the course of the activation process. The BET surface area, however, decreased to  $1509.79 \text{ m}^2 \text{ g}^{-1}$  as activating agent ratio increased to 3, probably due to pore blocking effects as previously reported by Kennedy *et al.*<sup>25</sup> In addition, the deterioration of porous structure resulting from destruction of walls between adjacent pores by the etching effect of KOH might also have contributed to the reduction in BET surface area. Previous reports suggest that the overall porosity of a chemically activated carbon materials increase with KOH concentration.<sup>26</sup> However, this trend was not observed in our case. For instance as KOH ratio increased from 0 to 1, mesopore surface area and volume increased from  $21\ 313.85$  to  $1253.46 \text{ m}^2 \text{ g}^{-1}$  and from  $0.36$  to  $0.85 \text{ cm}^3 \text{ g}^{-1}$  respectively. As KOH ratio increases further to 3, the mesopores surface area and volume decreased to  $169.3 \text{ m}^2 \text{ g}^{-1}$  and  $0.3 \text{ cm}^3$

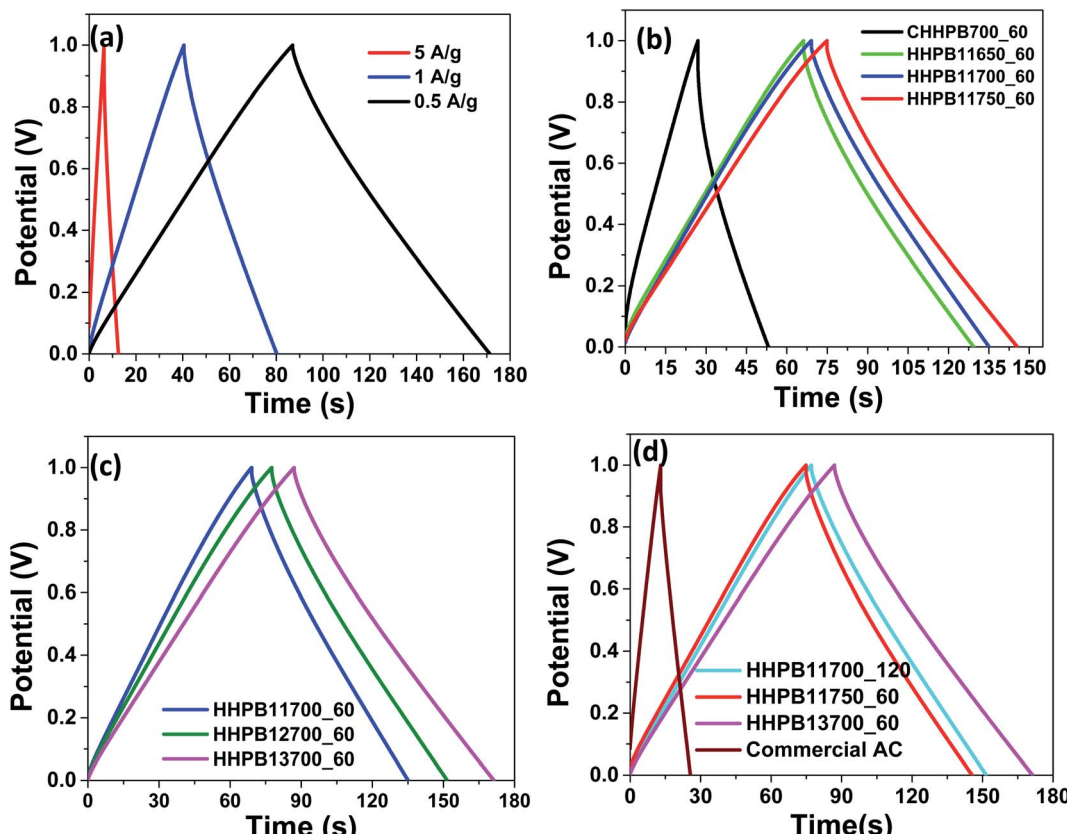


Fig. 7 CCD curves of (a) HHPB13700\_60 at different current densities (b and c) HHPB samples prepared at different activation temperature and KOH ratios at 0.5 A g<sup>-1</sup> (d) commercial AC compared with different HHPB samples at current density of 0.5 A g<sup>-1</sup> in two electrode systems.

$\text{g}^{-1}$  respectively. These results demonstrate the strong influence of KOH amount on the textural properties of the HHPB samples especially in the formation of mesopores. The formation mechanism of porosity in carbon at different KOH concentrations is a very complex process which follows several steps as depicted by eqn (5)–(11). Normally, the KOH etchant reacts with carbon generating the alkali metal (K), alkali compounds ( $\text{K}_2\text{O}$  and  $\text{K}_2\text{CO}_3$ ), and  $\text{H}_2$  gas in several steps as per eqn (5) and (6). The whole process can be explained as follows: at lower temperature around 400 °C dehydration of KOH to  $\text{K}_2\text{O}$  occurs (eqn (6)). Water ( $\text{H}_2\text{O}$ ) then react with C and CO to form  $\text{H}_2$  as per eqn (7) and (8) followed by carbonate formation, eqn (9). At temperatures higher than 700 °C, the carbonates decompose into  $\text{CO}_2$  and  $\text{K}_2\text{O}$ , eqn (10). The  $\text{K}_2\text{O}$  formed is then reduced by carbon to form metallic potassium (K), eqn (11), which intercalate between carbon layers. During washing with HCl and water, the intercalated metallic K and other compounds containing K are removed resulting into expanded carbon lattices that cannot return to their earlier structure therefore pores that are essential for charge storage are formed.<sup>27–29</sup> Moreover, the surface area decreased from 3572.03 to 2453.56  $\text{m}^2 \text{ g}^{-1}$  as temperature increased from 650 to 750 °C owing to carbon microstructure coarsening and crystallites growth during graphitization process at high temperatures.<sup>21</sup> The pore sizes of all HHPB samples varied between 1.91–7.27 nm which

represent mesoporous materials according to IUPAC pore size definition.

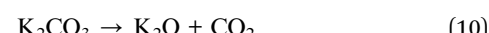
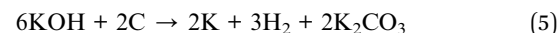


Fig. 5a and b presents the nitrogen adsorption–desorption isotherms at 77 K for HHPB samples. The shape of the isotherms looks different before and after activation at different KOH ratios and temperatures. The none activated sample, CHHPB700\_60 afforded type II isotherm representing un hindered monolayer–multilayer adsorption quintessential for non-porous materials. Its activated counterparts, samples HHPB11650\_60, HHPB11700\_60, HHPB12700\_60 HHPB13700\_60 and HHPB11750\_60 samples showed type IV



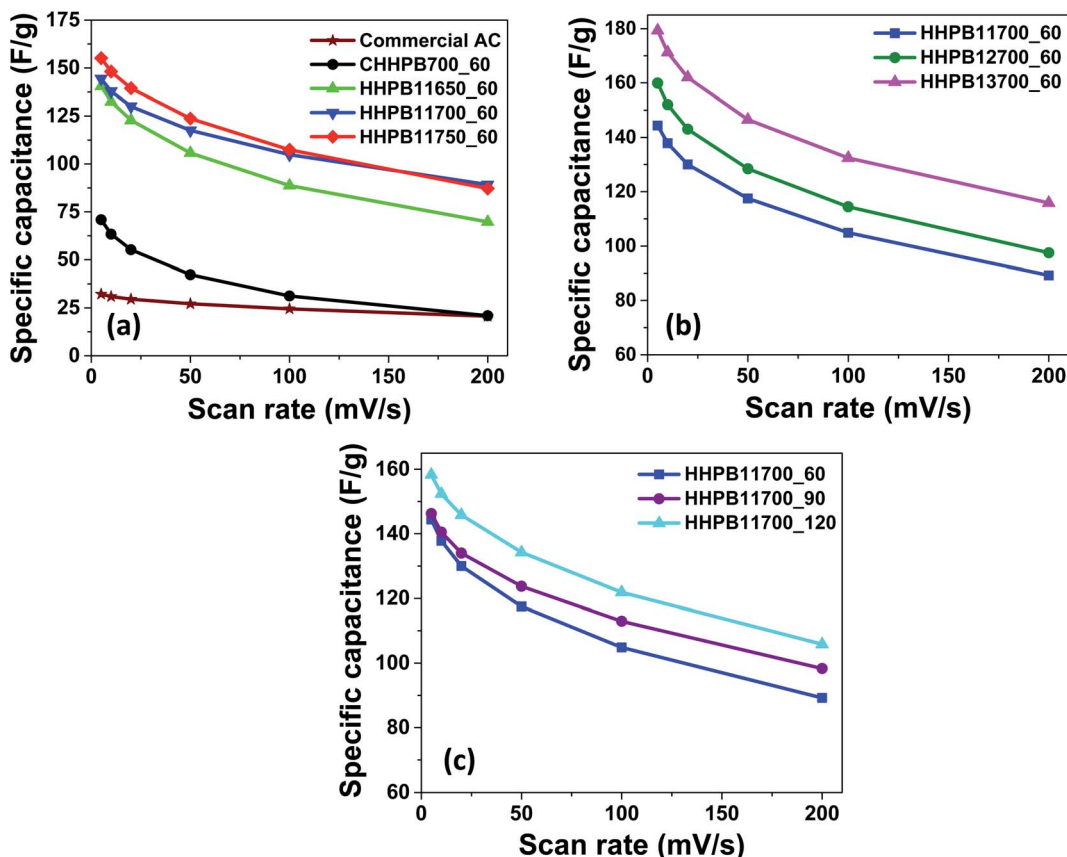


Fig. 8 Specific capacitance of HHPB samples and commercial AC at different scan rates calculated from CV by varying activation (a) temperature (b) KOH ratio (c) time.

isotherms typical for mesoporous materials with  $H_4$  type hysteresis loops between 0.4 and 1.0 relative pressure,<sup>30</sup> which is associated with the slit-shaped pores or particles with internal voids of irregular shape and broad size distribution.<sup>31</sup>

The pore size distributions of HHPB samples were analyzed by and BJH (Barrett–Joyner–Halenda) method and presented in Fig. 5c and d. It was observed that non-activated sample, CHHPB700\_60 afforded very low differential volume and almost a flat pore size distribution curve with a small hump around 4 nm, Fig. 5c. This together with the fact that the sample did not show any hysteresis loop (Fig. 5a) is indicative of its non-porous nature. At KOH activation ratio of 1, a sharp pore size distribution peak centered at around 4 nm and high differential volume was observed, Fig. 5c, which is in agreement with the relatively large hysteresis loop size observed for the same sample. Though the pore size distribution peak remained at around 4 nm, increasing KOH ratio to 2 dramatically reduced the differential volume (Fig. 5c) and the size of the hysteresis loop (Fig. 5a). Further increase in KOH ratio to 3 led to an almost flat pore size distribution curve with the lowest differential pore volume in the entire pore size range, Fig. 5c. The highest differential volume and relatively large hysteresis loop at KOH ratio of 1 show that the removal of inorganics and volatilization of extractables from the carbon precursor by the KOH thereby creating a high number of mesopores was successful. The

reduction of differential pore volume and pore size distribution peak with increase in KOH ratio implies loss of mesoporosity by the destruction of walls separating adjacent pores by the KOH. Similarly, an increase in activation temperature decreased the differential pore volume and shifted pore distribution peak to high pore diameter, Fig. 5d. This is probably due to carbon microstructure coarsening and crystallites growth during graphitization process at high temperatures leading to fusion of adjacent mesopores thereby forming bigger pores.

### Electrochemical performance

Electrochemical studies were performed to assess the performance of HHPB based supercapacitor in two electrode system. Fig. 6a-d presents the CV curves of electrodes made from different HHPB samples at scan rate of 5 to 200  $\text{mV s}^{-1}$ . From Fig. 6 it can be observed that, all CV curves of HHPB electrodes exhibited relatively regular rectangular shapes at all scan rates, attesting to their ideal EDL capacitance behavior of the produced carbon. Moreover, there was no noticeable distortion (polarization) of the CV curves from the standard rectangular shape indicating that the electrodes still allow smooth and easy penetration of the electrolyte ions during charge and discharge cycles even at higher scan rates, Fig. 6b. It is worth noting that the electrodes prepared from HHPB11750\_60 and HHPB13700\_60 samples show high area under the CV curves at



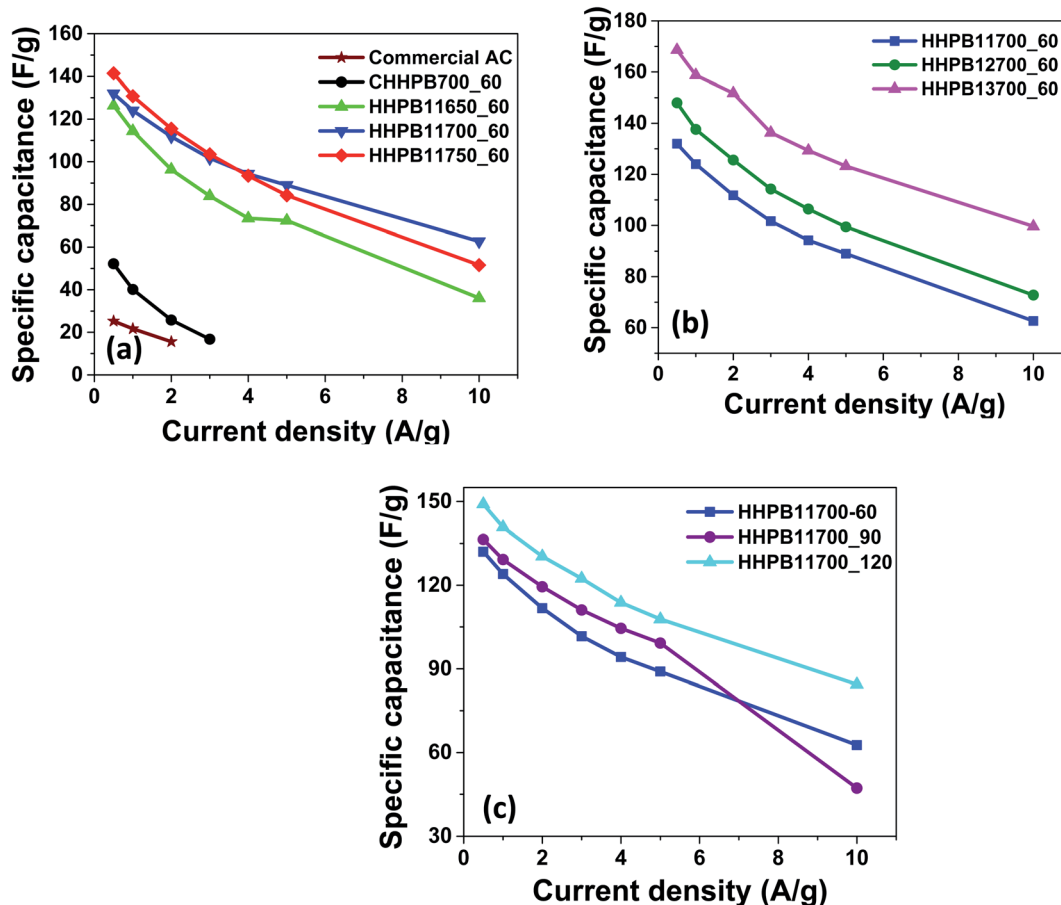


Fig. 9 Specific capacitance of HHPB samples and commercial AC at different current densities as calculated from CCD by varying activation (a) temperature (b) ratio (c) time.

scan rate of  $5 \text{ mV s}^{-1}$  when compared to non-activated sample, CHHPB700\_60. This is an indication that they possess large specific capacitance (Fig. 6c and d). We deduce this may be due to their high surface areas and structural defects compared to that of CHHPB700\_60.

Fig. 7a represents the CCD curves of the HHPB13700\_60 electrode at current density of  $0.5$  to  $5 \text{ A g}^{-1}$  while Fig. 7b and c shows CCD curves of electrodes from different HHPB samples prepared at different activation temperatures and KOH ratios at current density of  $0.5 \text{ A g}^{-1}$ , respectively. All CCD curves exhibit a regular triangular shape signifying their ideal EDL capacitance and good charge-discharge reversibility which are in good agreement with the CV plots presented earlier. The charge-discharge time for HHPB11750\_60 and HHPB11700\_60 electrodes is higher than that of the HHPB11650\_60 indicating that the specific capacitance increased with increase in activation temperature. It is also worth to note that, charge-discharge time for HHPB13700\_60 electrodes is larger than that of HHPB12700\_60 and HHPB11700\_60 showing that capacitance increased with increase in activating KOH ratio (Fig. 7c). This superior performance is attributed to its high BET surface area that provided a larger number of active site for the electrolyte ions. Commercial activated carbon (AC) posted substantially

larger IR drop and shorter charge-discharge time compared to all HHPB carbon based electrodes demonstrating the superior electrochemical performance of the carbon materials prepared in our study, Fig. 7d.

Fig. 8 and 9 illustrate the specific capacitances of different devices made from different HHPB samples at different scan rates and current densities as calculated using eqn (1) and (2), respectively. The specific capacitance of all HHPB based cells decreased with the increase in scan rate from  $5$  to  $200 \text{ mV s}^{-1}$  and current density from  $0.5$  to  $10 \text{ A g}^{-1}$  due to limited time of electrolyte to infiltrate to the pores of the electrode materials.<sup>32</sup>

#### Effect of combined activation conditions on carbon yield and electrochemical performance

Activating agent ratio is one among the factors that greatly influence physical properties and electrochemical performance of biomass-based carbon materials for supercapacitor applications. At  $650 \text{ }^{\circ}\text{C}$  and  $60 \text{ min}$  activation conditions, specific capacitance increased to a maximum at KOH ratio of  $2$  then dropped, Fig. 10a. This trend was the same for  $90$  and  $120 \text{ min}$  activation time, Fig. 10b and c. Increasing activating KOH ratio from  $1$  to  $3$  increased specific capacitance consistently for samples activated at  $700 \text{ }^{\circ}\text{C}$  for  $60$  and  $90 \text{ min}$  as shown in



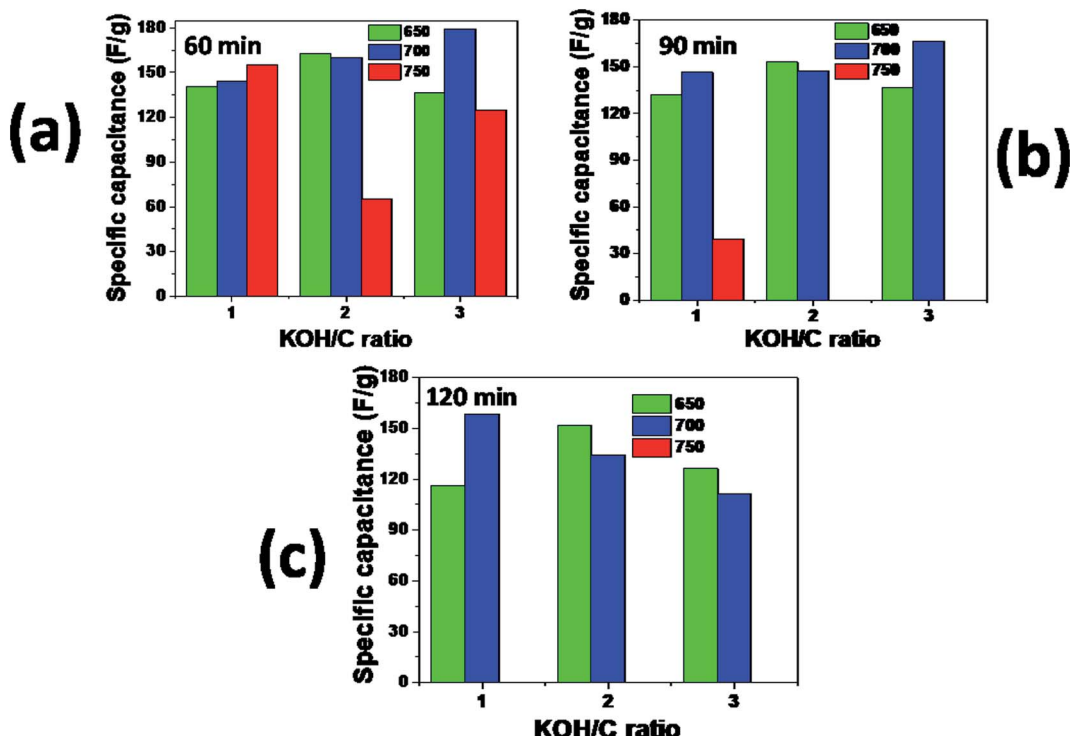


Fig. 10 Specific capacitance of HHPB samples activated at different KOH/C ratio, temperature and time as calculated from CV curves at scan rate of  $5 \text{ mV s}^{-1}$ , (a) 60 min, (b) 90 min, and (c) 120 min(a) 60 min, (b) 90 min, and (c) 120 min.

Fig. 10a and b. Contrary, the specific capacitance decreased consistently as the KOH ratio increased from 1 to 3 at  $700 \text{ }^{\circ}\text{C}$  and 120 min, Fig. 10c. Increasing activation KOH ratio, time, and temperature above 1, 90 min, and  $700 \text{ }^{\circ}\text{C}$ , respectively, led to burning out of all the sample. On the other hand, specific capacitance increased with time only for samples prepared at KOH ratio 1 at  $700 \text{ }^{\circ}\text{C}$  activation temperature, Fig. 10a–c. Thus the optimum activation conditions for producing carbon materials with high specific capacitance from palm bark are KOH ratio 3,  $700 \text{ }^{\circ}\text{C}$ , and 60 min (HHPB13700\_60) which

attained the highest specific capacitance of  $179.3$  and  $169 \text{ F g}^{-1}$  from CV and CCD at scan rate of  $5 \text{ mV s}^{-1}$  and current density of  $0.5 \text{ A g}^{-1}$ , respectively, for a 2 electrodes supercapacitor device.

Moreover, it has been reported that mesoporous structure and large surface area facilitate fast electrolyte ions transfer thereby enhancing specific capacitance.<sup>33</sup> But this is not always the case. As shown in Table 2, sample HHPB13700\_60 with surface area of  $1509.79 \text{ m}^2 \text{ g}^{-1}$  attained the highest specific capacitance of  $169 \text{ F g}^{-1}$  while sample HHPB11650\_60 with surface area of  $3572.03 \text{ m}^2 \text{ g}^{-1}$  attain specific capacitance of  $148$

Table 2 Relationship between BET surface area,  $V_{\text{micro}}$  and  $V_{\text{meso}}$  with specific capacitance

Sample	BET ( $\text{m}^2 \text{ g}^{-1}$ )	$V_{\text{micro}}$ ( $\text{cm}^3 \text{ g}^{-1}$ )	$V_{\text{meso}}$ ( $\text{cm}^3 \text{ g}^{-1}$ )	$V_{\text{micro}}/V_{\text{meso}}^a$	SP <sup>b</sup> ( $\text{F g}^{-1}$ )
<b>Activating ratio</b>					
CHHPB700_60	532.50	0.10	0.35	0.29	52
HHPB11700_60	1830.61	0.25	0.85	0.29	132
HHPB12700_60	1478.11	0.09	0.55	0.16	148
HHPB13700_60	1509.79	0.62	0.30	2.07	169
<b>Activating temperature</b>					
HHPB11650_60	3572.03	1.10	1.06	1.04	148
HHPB11700_60	1830.61	0.25	0.85	0.29	132
HHPB11750_60	2453.56	0.03	1.76	0.02	141
<b>Activating time</b>					
HHPB11700_60	1830.61	0.25	0.85	0.29	132
HHPB11700_120	1899.31	0.68	0.23	2.96	149

<sup>a</sup> Ratio of micropore to mesopore volume. <sup>b</sup> Specific capacitance.

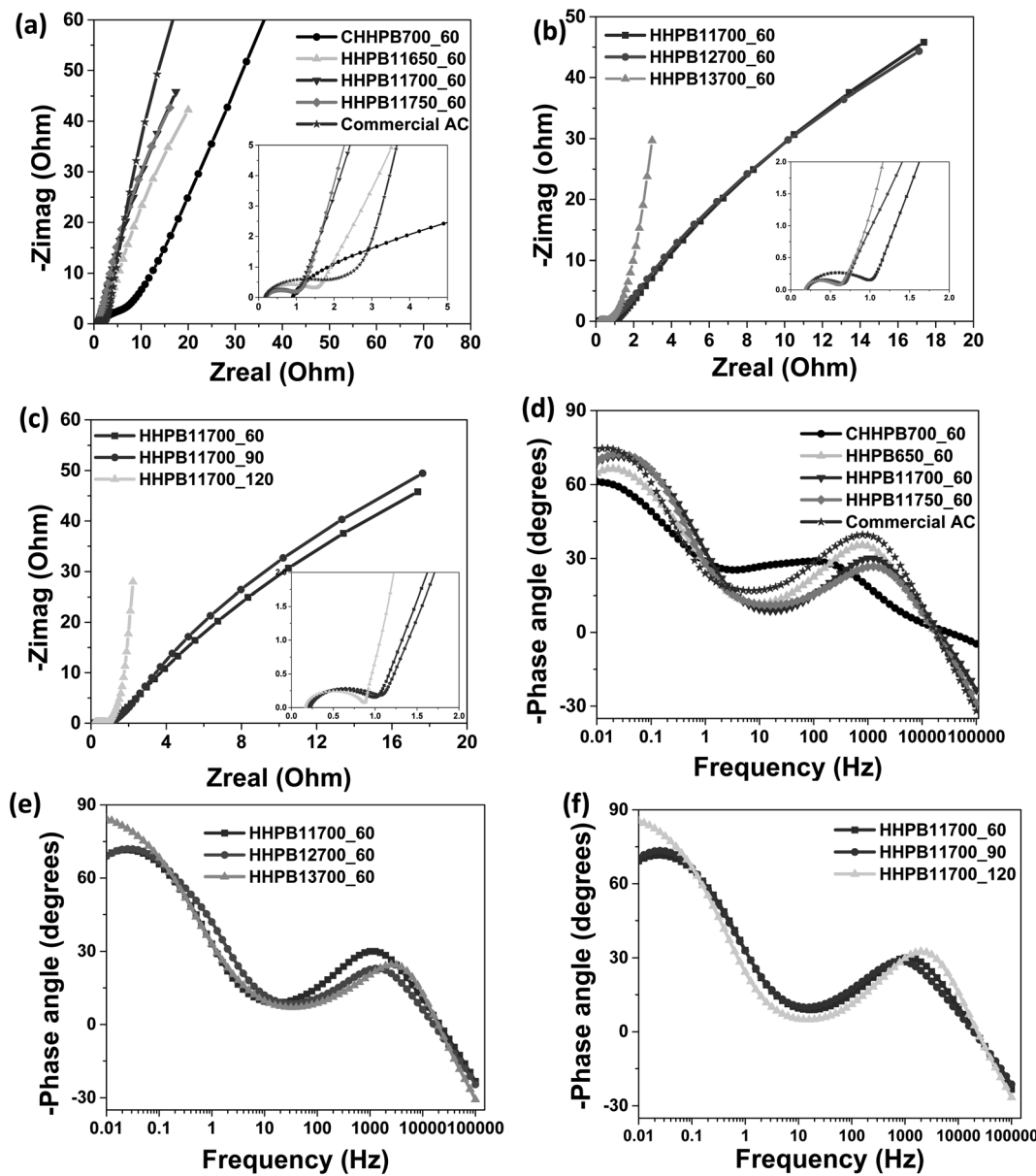


Fig. 11 (a–c) Nyquist and (d–f) Bode plots of HHPB samples at different activation temperatures, KOH ratios, and time.

$F\text{ g}^{-1}$ . This is indicative of the fact that BET surface area is not the main factor influencing the electrochemical performance. Further analysis of micro to mesopores volume ratio ( $V_{\text{micro}}/V_{\text{meso}}$ ) showed that for all the activation parameters: carbon/KOH ratio; temperature and time, the specific capacitance and hence the performance of the supercapacitor device was highest when  $V_{\text{micro}}/V_{\text{meso}}$  was largest, Table 2.

Electrochemical impedance spectroscopy (EIS) was performed and the impedance spectra of HHPB samples at different activation conditions are shown in Fig. 11. The Nyquist plots Fig. 11a–c consists of two main regions: the semi-circle at the high-frequency region where its diameter represent the charge transfer resistance of electrodes and the solution interface and the straight line in low frequency, showing the EDL

capacitance behavior. The charge transfer resistance decreased with increase in activation temperature from 650 to 750 °C (Fig. 11a), activation KOH ratio (Fig. 11b) and time (Fig. 11c). The intercept at the  $Z_{\text{real}}$  axis shows the equivalent series resistance (ESR) or internal resistance of the electrode. As seen in inserts of Fig. 11a–c representing the enlargement of the high-frequency region, all ESR values are around 0.25 Ω indicating good electrical conductivity and fast exchange of ions between the electrode and electrolyte interface. The semicircle of commercial AC is larger than that of HHPB samples implying its high charge transfer resistance and IR drop and thus its poor electrochemical performance in supercapacitor device in terms of specific capacitance, Fig. 9a, and power and energy density, Fig. 12f.



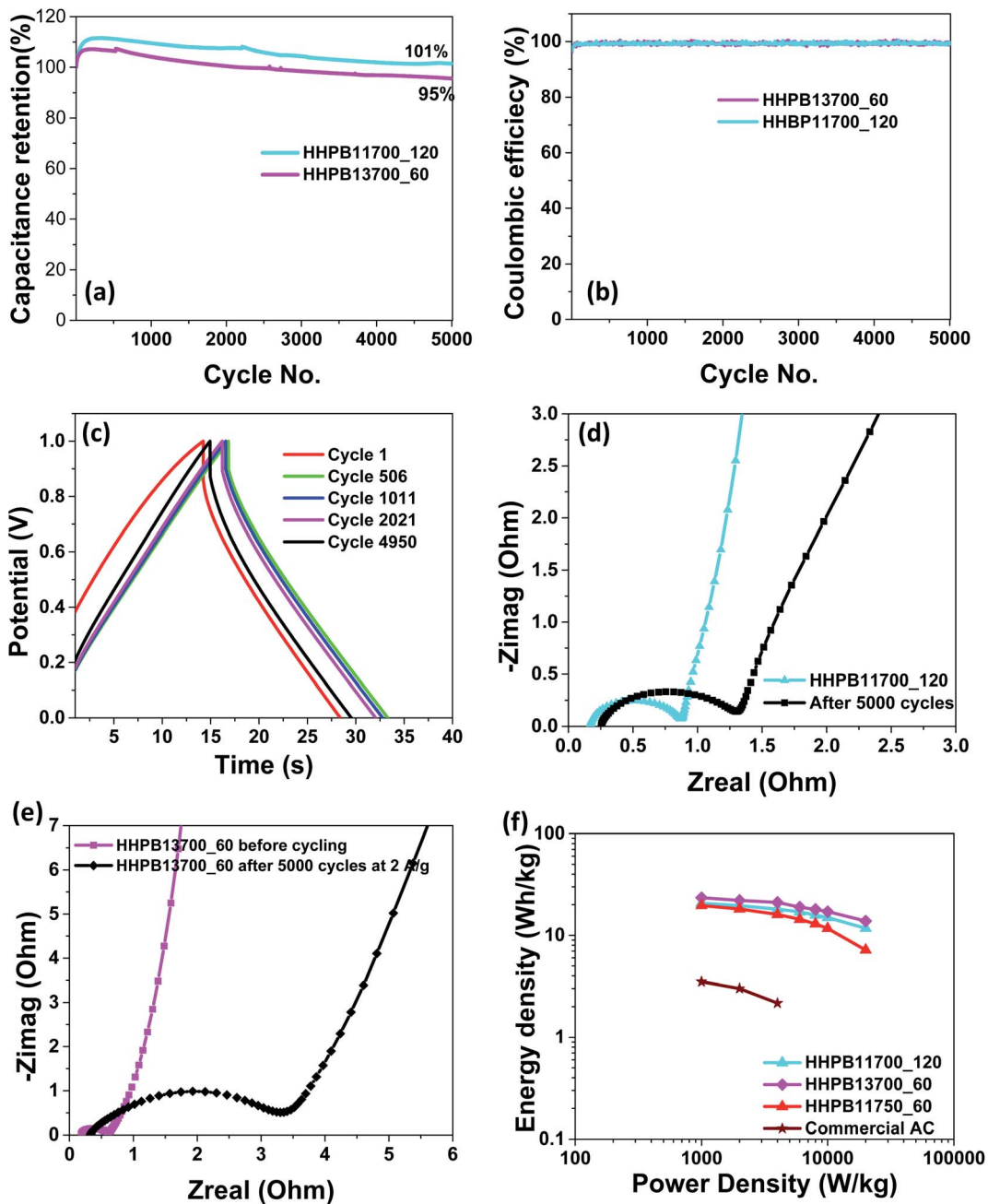


Fig. 12 (a) Capacitance retention of HHPB13700\_60 and HHPB11700\_120 at current density  $2 \text{ A g}^{-1}$  (b) coulombic efficiency of HHPB13700\_60 and HHPB11700\_120 at current density  $2 \text{ A g}^{-1}$  (c) CCD curves of HHPB13700\_60 at 1<sup>st</sup>, 506<sup>th</sup>, 1011<sup>th</sup>, 2021<sup>st</sup>, and 4950<sup>th</sup> cycle (d) Nyquist plot of HHPB11700\_120 before and after 5000 charge–discharge cycles at  $2 \text{ A g}^{-1}$  (e) Nyquist plot of HHPB13700\_60 before and after 5000 charge–discharge cycles at  $2 \text{ A g}^{-1}$  (f) Ragone plot of HHPB samples compared with commercial AC.

Fig. 11d-f represents the relationship between the phase angle and the frequency. Activation conditions were found to have different effects on the phase angle. For instance, increasing activation temperature from 650 to 750 °C at low frequency, the phase angle increased from  $-64$  to  $-70$ °, Fig. 11d. Introducing activating agent at different ratios from 0 to 3 caused the phase angle to increase from  $-61$ ° for non-activated sample, CHPB700\_60, to  $-84$ ° for HHPB13700\_60 which is very close to that of an ideal capacitor of  $-90$ , Fig. 11e.

This explains the good capacitive behavior (highest specific capacitance) demonstrated by these materials. Likewise the phase angle increased from  $-70$  to  $-85$ ° as activation time increased from 60 to 120 min, Fig. 11f.

Cycling stability of the electrode material is a significant parameter to consider in energy storage applications. Therefore, cycling stability of HHPB13700\_60 and HHPB11700\_120 in two electrode system was studied by measuring 5000 charge–discharge cycles at current density of  $2 \text{ A g}^{-1}$ . HHPB11700\_120

Table 3 Performance of different biomass-based carbon for supercapacitor applications<sup>a</sup>

Carbon source	Activation route	BET SSA (m <sup>2</sup> g <sup>-1</sup> )	Electrode configuration	Electrolyte	SPC* (F g <sup>-1</sup> )	CR* (%)	Ref.
Coconut kernel pulp (milk free)	KOH activation	1200	2 electrode system	1 M TEABF <sub>4</sub> in PC	56 at 0.25 A g <sup>-1</sup>	100% at 1 A g <sup>-1</sup> after 2000 cycles	35
Fallen leaves	Mixed KOH and K <sub>2</sub> CO <sub>3</sub>	1078	2 electrode	6 M KOH	223 at 0.5 A g <sup>-1</sup>	~100% at 1 A g <sup>-1</sup> after 2000 cycles	36
Sugar cane bagasse	ZnCl <sub>2</sub>	1155	2 electrode	1 M H <sub>2</sub> SO <sub>4</sub>	300	77 at 5 A g <sup>-1</sup>	37
Starch	Stabilization, carbonization followed by KOH activation	3251	2 electrode	6 M KOH	304 at 0.05 A g <sup>-1</sup>	98 at 40 A g <sup>-1</sup> after 10 000 cycles	38
Fructose corn syrup	Hydrothermal followed by physical self-activation	1223	2 electrode	6 M KOH	168 at 0.2 A g <sup>-1</sup>	~83 at 0.5 A g <sup>-1</sup> after 2000 cycles	39
Fan palm barks	KOH	1509.79	2 electrode	6 M KOH	169 at 0.5 A g <sup>-1</sup>	95 at 2 A g <sup>-1</sup> after 5000 cycles	This study

<sup>a</sup> SPC\* = specific capacitance, CR\* = capacitance retention.

and HHPB13700\_60 electrodes retain 101 and 95% of its capacitance, respectively, after 5000 CCD cycles with 100% coulomb efficiency, Fig. 12a and b, approving the suitability of the HHPB materials for supercapacitor device. The HHPB-based device shows excellent charge–discharge cycling behavior as demonstrated by the increase in charge–discharge time and reduction in IR drop in the first 2021 cycles, Fig. 12c, attributed to electrochemical activation of the electrodes.

Furthermore, EIS measurement of HHPB13700\_60 and HHPB11700\_120 devices were carried out after 5000 charge discharge cycles cycling at 2 A g<sup>-1</sup> and presented in Fig. 12d and e. It can be seen that both plots shows a semicircle and a straight line in the high and low frequency regions, respectively. The straight line in the low frequency region after cycling is more slanted than before cycling, indicating low ions diffusion and mass transport at electrode/electrolyte interface. Also, the diameter of the semicircle in the EIS spectra increased after 5000 charge–discharge cycles implying increase in charge transfer resistance arising from longer ion/electron transport pathway as also revealed by Agudosí *et al.*<sup>34</sup> From Nyquist plot, the charge transfer resistances of HHPB11700\_120 device increased from 0.70 Ω (before cycling) to 1.05 Ω (after cycling) while that of HHPB13700\_60 the device increase from 0.47 Ω (before cycling) to 3.8 Ω (after cycling).

The power and energy densities are crucial parameters to consider for supercapacitor applications and are commonly plotted on Ragone plots. Fig. 12f shows the Ragone plot of different HHPB electrodes whereby HHPB13700\_60 based supercapacitor device afforded the highest energy density compared to other HHPB samples and 7 times greater than that of commercial AC. The electrochemical performance of HHPB materials prepared in our study is relatively higher than that of other biomass based carbon materials presented in the literature, Table 3.

The result from this study shows that electrodes derived from palm barks have a good rate capability compared to some other biomass-based carbon presented in literature as summarized in Table 3 suggesting that it can be utilized in devices that need fast charging process.

## Conclusion

Plentiful and low cost *Washingtonia robusta* barks (Mexican fan palm) were used to successfully synthesize good carbon materials for 2 electrode supercapacitor device *via* by hydrothermal treatment followed by KOH chemical activation. The electrochemical studies shows that the specific capacitance of the supercapacitor device was affected by material preparation parameters including activation temperature, impregnation ratio and time. Both the charge transfer and Warburg resistance were found to decrease and phase angle increase towards the idea  $-90^\circ$  with increase in activation KOH ratio, temperature, and time. The symmetric supercapacitor device made from HHPB13700\_60 sample attained a specific capacitance of 179.3 and 169 F g<sup>-1</sup> from CV and CCD at scan rate of 5 mV s<sup>-1</sup> and current density of 0.5 A g<sup>-1</sup>, respectively. The device exhibits capacitance retention of 95 and 100% coulombic efficiency after 5000 charge–discharge cycles. The supercapacitor devices exhibited perfect EDLC behavior with approximate energy densities of 20 W h kg<sup>-1</sup> at power density of 2000 W kg<sup>-1</sup>. These findings show that carbon materials derived from palm barks have a huge potential in supercapacitor device applications.

## Author contributions

Joyce Elisadiki: conceptualization, methodology, investigation, validation, formal analysis, writing – original draft. Mavis K. Gabookolwe: investigation and writing–reviewing and editing. Oluwatayo R. Onisuru: investigation and writing–reviewing and editing. Reinout Meijboom: resources and writing–reviewing and editing. Cosmas Muiva: formal analysis and writing–reviewing and editing. Cecil K. King'ondu: supervision, conceptualization, formal analysis, methodology, resources, project administration, funding acquisition and writing–reviewing and editing.

## Conflicts of interest

The authors declare that they have no competing interests.



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