ELECTROCHEMICAL PERFORMANCE OF ORDERED MESOPOROUS CARBON MODIFIED BY OXIDATIVE TREATMENT WITH AQUEOUS NITRIC ACID

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Abstract

In this study, ordered mesoporous carbon (OMC) was prepared via nano-casting method by using Santa Barbara Amorphous (SBA)-15 as a template and sucrose as a carbon precursor. The OMC was subsequently oxidized with aqueous nitric acid and referred as MOMC. The physicochemical properties of OMC and MOMC were determined using nitrogen adsorption–desorption analyser, field emission scanning electron microscopy (FESEM), transmission electron microscopy (TEM), and Fourier transform infrared spectroscopy (FT-IR). The results proved that the carbon replication process was successful. The electrochemical performance tests were carried out using cyclic voltammetry (CV) and galvanostatic charge–discharge (GCD) in 1 M KOH electrolyte for 1000 cycles. After oxidative treatment, the specific surface area and pore volume of OMC decreased but the specific capacitance of the electrode material has significantly increased from 117 F g⁻¹ to 344 F g⁻¹ at a scan rate of 10 mV s⁻¹.

Keywords: Ordered mesoporous carbon; SBA-15; template; oxidative treatment; electrochemical

Abstrak

Dalam kajian ini, karbon mesoliang bertertib (OMC) telah disediakan melalui kaedah penuangan-nano dengan menggunakan Santa Barbara Amorphous (SBA)-15 sebagai templat dan sukrosa sebagai pelopor karbon. OMC kemudiannya dioksida dengan asid nitrik akues dan dirujuk sebagai MOMC. Ciri-ciri fizikokimia OMC dan MOMC ditentukan menggunakan pengesan penjerapan–nyahjerapan nitrogen, mikroskop elektron pengimbas pancaran medan (FESEM), mikroskop pancaran elektron (TEM), dan spektroskopi inframerah transformasi Fourier (FT-IR). Keputusan kajian membuktikan bahawa proses replikasi karbon berjaya. Ujian prestasi elektrokimia telah dilaksanakan menggunakan voltammetri kitaran (CV) dan galvanostatik (GCD) di dalam elektrolit KOH 1 M untuk 1000 kitaran. Selepas rawatan oksidatif, luas permukaan spesifik dan isi padu liang OMC berkurang namun kapasitan spesifik bahan elektrod tersebut meningkat dengan ketara dari 117 F g⁻¹ kepada 344 F g⁻¹ pada kadar imbasan 10 mV s⁻¹.

Kata kunci: Karbon mesoliang bertertib; SBA-15; template; rawatan oksidatif; elektrokimia

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1.0 INTRODUCTION

There has been great interest in the development of ordered mesoporous carbon (OMC) material owing to its ordered pore structure, high specific surface area, large pore volume, chemical inertness, high mechanical stability and conductivity [1–2]. OMC can be prepared via nano-casting method which allows control of structure and morphology [3]. Generally, nano-casting method is a process in which a template with relevant nano-scale structure is impregnated with carbon precursor, and the initial template is subsequently removed [4]. The resulting carbon exhibits as mentioned criteria with variable three dimensional framework structures such as cubic and hexagonal, depending on the structure of the template. In other words, OMC prepared is usually a negative replicate of the used template. This method mainly used ordered mesoporous silica as template and sucrose or phenol resin as carbon precursor [5–6].

OMC material has been used in energy storage and electrochemistry applications due to its micropores which is essential for ions accumulation and well-aligned mesopores system (2–50 nm) which acts as highway for smooth electron and ions transportation [7]. However, the use of OMC materials as an electrode material for ideal electrochemical capacitor is mainly due to contribution of electrical double-layer capacitance, and the specific capacitance was less than 200 F g⁻¹ [8].

Previous reports demonstrated that the application of this material is strongly influenced not only by its structural properties, but also by its surface functionalities. Oxygen surface functionalities (e.g., carboxylic acid, carbonyl, and hydroxyl) can be introduced by liquid phase oxidative treatment with nitric acid, hydrogen peroxide, or iron (III) nitrate. These oxygen surface functionalities are able to generate additional capacitance through faradaic reaction and recover the carbon surface hydrophilicity in aqueous electrolyte thus lead to improvement in electrochemical capacitive behaviour [9–10]. Therefore, it is of great interest to verify the influence of oxygen surface functionalities on the electrochemical performance of OMC materials.

In this study, OMC was prepared via nano-casting method by using SBA-15 as the template, followed by oxidative treatment with aqueous nitric acid. The as-synthesize OMC and oxidized OMC (MOMC) were characterized with respect to their morphological and structural properties and the electrochemical performance was investigated. The objectives of this study are to identify the differences observed in electrochemical behaviour of OMC and MOMC and clarify them in accordance to the role of oxygen surface functionalities.

2.0 EXPERIMENTAL

2.1 Materials

The Pluronic 123, tetraethyl orthosilicate (TEOS) and sucrose were purchased from Sigma–Aldrich, Germany. All other chemicals used for materials preparation and electrochemical testing were purchased from QReC™, Merck, Sigma–Aldrich and RCI Labscan Ltd., and used without further purification.

2.2 Synthesis of OMC and MOMC

SBA-15 template was prepared according to the procedure described by Zhao et al. [11]. Then, OMC was prepared via nano-casting method according to procedure described by Ryoo et al. [12]. The as-synthesize OMC was oxidized with 2 M HNO₃ aqueous solution at 80 °C for 2 h by reflux and the product obtained was referred as MOMC. For electrochemical performance testing, OMC and MOMC electrodes were prepared by mixing the samples (70 wt.%) with carbon (Super P, 15 wt.%) and PVDF binder (15 wt.%) by using NMP as a solvent. The mixture was stirred for 24 h and coated onto a pre-cleaned nickel foam substrate, heated at 60 °C for 24 h and finally pressed at 5 MPa.

2.3 Physicochemical Characterization of OMC and MOMC

The textural properties of OMC and MOMC such as specific surface area, pore volume, and pore size distribution were evaluated from nitrogen adsorption–desorption isotherms by using Micromeritics ASAP 2010 volumetric adsorption analyser. The specific surface area was determined from Brunauer–Emmet–Teller (BET) plot and pore size distribution was estimated by Barrett–Joyner–Halenda (BJH) method from the desorption branch. TEM and FESEM images were obtained using JOEL microscopic instruments. The chemical species present after oxidative treatment was identified by Perkin-Elmer series 1600 FT-IR spectrometer.

2.4 Electrochemical Performance Tests

Cyclic voltammograms were obtained in the potential range of -1.0–0.0 V at various scan rates using three–electrode system using a potentiostat (Autolab PGSTAT 30, Eco Chemie B.V., The Netherlands). The gravimetric specific capacitance \( C_s \) of OMC and MOMC was calculated using the following equation:

\[
C_s = \frac{1}{2mn} \int_{E_1}^{E_2} \frac{i(E) \ dE}{E_2 - E_1}
\]

(1)
where \( m, v, E_2, E_1 \) and \( i(E) \) are sample mass, potential scan rate, higher potential cutoff, lower potential cutoff, and current, respectively. Galvanostatic charge–discharge (GCD) was obtained by cycling the potential from -1.0 to 0.0 V with different current density. The specific capacitance \( (C_S) \) was also calculated from charge–discharge cycling using the equation:

\[
C_S = \frac{it}{m\Delta V}
\]  

where \( i, t, m, \) and \( \Delta V \) are constant discharge current, discharge time, sample mass, and open potential, respectively. All of the electrochemical measurements were performed using platinum as the counter electrode and Ag/AgCl as the reference electrode. Electrochemical performance of OMC and MOMC electrodes was compared in 1 M KOH electrolyte.

### 3.0 RESULTS AND DISCUSSION

#### 3.1 Physicochemical Characterization of OMC and MOMC

The nitrogen adsorption–desorption isotherms of OMC and MOMC, as shown in Figure 1 (a) described typical type IV profile which is a characteristic of mesoporous materials [13]. This phenomenon is associated with capillary condensation in mesopores commenced at relative pressure \( P/P_o > 0.42 \). After oxidative treatment, BET isotherm studies showed decrease in specific surface area (\( S_{BET} \)) from 899.30 m\(^2\) g\(^{-1}\) (OMC) to 551.10 m\(^2\) g\(^{-1}\) (MOMC) and the pore volume decreased from 0.61 cm\(^3\) g\(^{-1}\) (OMC) to 0.40 cm\(^3\) g\(^{-1}\) (MOMC) for mesopores and decreased from 0.32 cm\(^3\) g\(^{-1}\) to 0.21 cm\(^3\) g\(^{-1}\) for micropores (seen in Table 1). The decrease in \( S_{BET} \) and pore volumes of MOMC may attributed to the introduction of oxygen surface functionalities over its pore wall and surface that caused pore disorientation or/and destruction on MOMC. Similar phenomenon has also reported by Cheah and co-workers on carbon coated monolith [14]. Besides, OMC shows H3 hysteresis loop, which does not exhibit any limiting adsorption at high \( P/P_o \), representing aggregates of carbon particles with slit-shape pores [13]. After oxidative treatment, the MOMC shows similar loop shape but with narrower size.

Different structural parameters of OMC and MOMC were also observed from their pore size distributions calculated from BJH method, as presented in Figure 1 (b). The pore volume broad peaks centred at 4.10 nm and 4.45 nm for OMC and MOMC, respectively, suggests that the pore volumes are predominantly composed of mesopores and concentrated in a relatively wide size range. However, SBA-15 (data not shown) has larger pore size and better pore uniformity based on a much narrow and sharp capillary condensation step. This phenomenon occurred as OMC polymers undergo a substantial volume contraction and pore narrowing/broadening during high temperature carbonization (800 °C) [15]. Nevertheless, the replication process of OMC from SBA-15 template was successful.

![Figure 1](image-url)  

**Figure 1** Nitrogen adsorption–desorption isotherms (a) and pore size distributions (b) of OMC and MOMC.

<table>
<thead>
<tr>
<th>Sample</th>
<th>( S_{BET} ) (m(^2) g(^{-1}))</th>
<th>( V_{MESO} ) (cm(^3) g(^{-1}))</th>
<th>( V_{MICRO} ) (cm(^3) g(^{-1}))</th>
<th>( d_{BJH} ) (nm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>OMC</td>
<td>899.30</td>
<td>0.61</td>
<td>0.32</td>
<td>4.10</td>
</tr>
<tr>
<td>MOMC</td>
<td>551.10</td>
<td>0.40</td>
<td>0.21</td>
<td>4.45</td>
</tr>
</tbody>
</table>

\( a \) Surface area determined from multi point BET analysis.  
\( b \) From the difference between the pore volume at \( P/P_o = 0.97 \) and \( V_{MICRO} \).  
\( c \) From \( t \) plot (Harkins–Jura equation).  
\( d \) Calculated from desorption branch of the isotherm by BJH method.

![Figure 2](image-url)  

**Figure 2** FESEM images of OMC (a) and MOMC (b); TEM images of OMC (c) and MOMC (d)
particle morphology of OMC and MOMC with average length of 600–700 nm and 200–300 nm width. The obtained OMC had successfully retained the rod-like shape particle morphology of SBA-15 template, similar with previous study [6,16]. In addition Figure 2 (c) and (d), the TEM images of OMC and MOMC, respectively, exposed the presence of well-aligned mesopores channels with average width of less than 6 nm. From FESEM and TEM observation, we know that the ordered mesostructure structure of OMC was preserved after oxidative treatment and it is constant with nitrogen adsorption–desorption analysis data.

![FT-IR spectra of OMC and MOMC](image)

FT-IR spectra of OMC and MOMC are shown in Figure 3. Both samples exhibited two identical characteristic peaks at 3400 cm\(^{-1}\) and 1600 cm\(^{-1}\) corresponding to O–H group stretching and C=O stretching vibrations in graphene sheets, respectively. After oxidative treatment, two significant peaks appeared at 1720 cm\(^{-1}\) (C=O stretching vibration) and 1230 cm\(^{-1}\) (C–O vibration) indicated the existence of carboxyl groups. In addition, due to the usage of HNO\(_3\), a weak absorption peak corresponding to NO\(_2\) bending vibrations is observed around 1340 cm\(^{-1}\). The –C=O and –C–O functional groups play an important role in enhancement of the carbon surface hydrophilicity as well as the electrochemical performance.

### 3.2 Electrochemical Studies

Figure 4 (a) and 4 (b) show CV profiles of OMC and MOMC electrodes at various scan rates (10, 25, 50, and 75 mV s\(^{-1}\)) in 1 M KOH electrolyte. The profiles comprise a series of approximately rectangular-like shape with a steep current change at the switching potential, which indicating electric double-layer capacitor behaviour. As the scan rates rises, both peak currents and peak-to-peak separation increased. The curves are able to preserve near rectangular shape as 75 mV s\(^{-1}\) is achieved, suggesting a highly reversible fast charging and discharging processes. After oxidative treatment, distortion of the CV curves becomes pronounce probably due to slow charging process induced by localization of electrons on the inhomogeneous MOMC surface [9]. The oxidative treatment also affects electronic resistance of the carbon materials [1,17].

![Cyclic voltammograms of OMC (a) and MOMC (b)](image)

The specific capacitance \(C_S\) of OMC and MOMC was measured at various scan rates (10–150 mV s\(^{-1}\)) and the values are presented in Figure 5 (a). It can be clearly seen that the \(C_S\) of both electrode materials are strongly dependent on the scan rate. Specifically, the \(C_S\) decreases with increase in scan rate. The highest \(C_S\) was obtained at 10 mV s\(^{-1}\). At all tested scan rate, MOMC exhibits higher \(C_S\) values compared to OMC. The increment in \(C_S\) is an indicative of the pseudo-capacitance presence happened on the MOMC electrode interface, due to redox reactions of oxygen surface functionalities (–C=O and –C–O groups). However, when the scan rate increased from 10 mV s\(^{-1}\) to 150 mV s\(^{-1}\), the \(C_S\) of OMC decreased 21% of its initial value. In contrast, the \(C_S\) of MOMC dropped by 35% from 344 F g\(^{-1}\) to 223 F g\(^{-1}\). The drop percentage is higher than OMC probably contributed by distributed-capacitance effect and impaired conductivity after the oxidative treatment [9]. Improved hydrophilicity of the MOMC surface has also contributed to the capacitance improvement by favouring the electrolyte ions transportation to the micropores [7].

Charge–discharge analysis was conducted to determine \(C_S\) retention of OMCs electrodes in order for the electrodes to work steadily and safely. Figure 5 (b) shows the charge–discharge curves of OMC and MOMC at 0.5 A g\(^{-1}\). The shape of the curves are not exactly linear, but show a typical triangle shape distribution which means that the materials have good double-layer electrochemical capability. MOMC experienced longer discharge time compared to OMC,
indicating a high charge–discharge propagation, low resistivity, and good reversible process [18]. The C₃ of OMC and MOMC electrodes were summarized in Table 2. Worth to mention that the C₃ of MOMC decreased to 146 F g⁻¹ as the current density increased to 15 A g⁻¹, which proves its mass transportation efficiency and good charge–discharge rate [7].

![Figure 5](image.png)

**Figure 5** The specific capacitance (Cₛ) as a function of scan rate (a) and charge–discharge curves at 0.5 A g⁻¹ (b) in 1 M KOH electrolyte

**Table 2** Specific capacitance (Cₛ) for OMC and MOMC electrodes at various current densities (0.5–15 A g⁻¹)

<table>
<thead>
<tr>
<th>Samples</th>
<th>Specific Capacitance, Cₛ (F g⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>0.5</td>
</tr>
<tr>
<td>OMC</td>
<td>153</td>
</tr>
<tr>
<td>MOMC</td>
<td>271</td>
</tr>
</tbody>
</table>

### 4.0 CONCLUSION

The electrochemical performance of OMC modified by oxidative treatment with aqueous nitric acid showed a significant improvement in specific capacitance (Cₛ) from 117 F g⁻¹ (OMC) to 344 F g⁻¹ (MOMC) due to contribution of pseudo-capacitance induced by faradic reaction of oxygen surface functionalities (–C=O and –C–O groups). The Cₛ achieved a maximum value of 344 F g⁻¹ at 10 mV s⁻¹ and still remained a value of 222 F g⁻¹ as the scan rate approaches 150 mV s⁻¹. The results proved that the oxidative treatment is suitable for OMC-based materials thus favouring its application as an electrode material for supercapacitor.

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**References**


