Electric Double Layer Capacitance on Hierarchical Porous Carbons in an Organic Electrolyte

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Abstract

Nanoporous carbons were prepared by using colloidal crystal as a template. Nitrogen adsorption/desorption isotherms and transmission electron microscope images revealed that the porous carbons exhibit hierarchical porous structures with meso/macropores and micropores. Electric-double-layer-capacitor performance of the porous carbons was investigated in an organic electrolyte of 1 M LiClO$_4$ in propylene carbonate and dimethoxyethane. The hierarchical porous carbons exhibited large specific double layer capacitance of c.a. 120 F g$^{-1}$ due to their large surface areas. In addition, the large capacitance was still obtained at a large current density up to 10 A g$^{-1}$, which satisfies demands from the high power application such as hybrid electric vehicles. Capacitance analysis of the hierarchical porous structures revealed the contribution of meso/macropores and micropore to the electric double layer capacitance to be 8.4 and 8.1 μF cm$^{-2}$, respectively. The results indicated electric double layer is formed even when solvated ions are larger than pore diameters.

Keywords: Porous carbon; Micropore; Electric double layer capacitor, Organic electrolyte; Hierarchical pore
1. Introduction

Electric double layer capacitors (EDLCs) with both large energy density and large power density have been strongly demanded especially from the viewpoint of application to auxiliary power sources of hybrid electric vehicles (HEVs) or fuel cell electric vehicles (FCEVs). Energy density of EDLCs is given by a formula: \( \frac{1}{2} CV^2 \) where \( C \) and \( V \) is capacitance and cell voltage, respectively. To increase the energy density, porous carbons with large surface, e.g., activated carbons, area are widely used. Double layer capacitance depends not only on surface areas of the porous carbons,[1] but also on pore sizes,[2-4] functional groups,[5] orientation of graphene layers,[6,7] and so on. Micropores (\( d < 2 \text{ nm} \), \( d \) being a pore diameter) exhibit large specific surface areas but ion transportation may be interfered by pore walls, which means poorer power density. And especially when pore diameters are smaller than solvated ions, the formation of double layer is supposed to be impossible. We have so far prepared porous carbons by a colloidal-crystal-templating method using phenolic resin as a carbon source.[8,9] This class of carbons attracts our interests not only because they possess large specific surface areas, but also because they exhibit two distinct pore size distributions: one is in the region of mesopores (\( 2 \text{ nm} \leq d < 50 \text{ nm} \)) or macropores (\( d \geq 50 \text{ nm} \)), which is originated from template SiO\(_2\) particles, and the other is in the region of micropores, which is derived from carbonization of the phenolic resin. In other words, these porous carbons possess a hierarchical porous structure: micropores exist on pore walls of
meso/macropores. For such a porous structure, electrolyte ions can be delivered smoothly through meso/macropores to micropore surfaces with large specific surface areas. The hierarchical porous structure is also of importance from a fundamental viewpoint because it allows us to analyze contribution to capacitance by meso/macropores and micropores, separately. In the case of activated carbons or typical mesoporous carbons,[10,11] pore sizes are continuously distributed from micropore to mesopore regions, which make the analysis complicated. In addition, electrolyte transport in their narrow pores cause the kinetic polarization and thus capacitance may be underestimated. In the previous study, we reported that in an aqueous electrolyte (1 M H₂SO₄), contribution of micropores to the double layer capacitance was almost negligible,[9] which may not be surprising because the solvated ions are as small as pore sizes.

Another way to achieve high-energy EDLCs is to employ organic electrolytes instead of aqueous electrolytes, because they exhibit wider potential windows,[12] and accordingly increase maximum cell voltage (V) from c.a. 1 V to c.a. 2–2.5 V. However, organic electrolytes exhibit some drawbacks in contrast to aqueous electrolytes: e.g., poorer electrolyte conductivity (typically ~10 mS cm⁻¹) due to their viscous solvent [12] and smaller specific double layer capacitance (2–10 μF cm⁻²).[13] Structure of double layer in organic electrolytes might be different from that in aqueous electrolytes. Chmiola et al. reported that micropores with diameters smaller than 1 nm exhibit larger specific capacitance than larger
pores in a non-aqueous electrolyte. The results are against the commonly accepted concept that such micropores hardly contribute to capacitance, and they proposed that the structure of double layer in very small pores are different from that in relatively large pores. In this context, we investigated the double layer capacitance of the hierarchical porous carbons in an organic electrolyte.

2. Experimental

Porous carbons were synthesized by following the same procedure as reported. First, colloidal crystal templates were fabricated by centrifugation of mono-dispersed SiO₂ colloid (particle diameter, \(d\): 8, 17, 45, 110 nm, Catalysis & Chemicals Co. Ltd.). Then, a mixture of phenol, formaldehyde (37%), and a small portion of hydrochloric acid was impregnated into interstices of the SiO₂ particles. After polymerization of phenol and formaldehyde and subsequent carbonization of the phenolic resin at 1,000 °C in Ar flow for 5 h, nanocomposites of SiO₂ and carbon was obtained. Finally, SiO₂ was removed by a hydrofluoric acid (46 wt-%) and meso/macroporous carbons were obtained. Hereafter, the porous carbons thus obtained is referred to as \(C[d_{SiO2}]\) where \(d_{SiO2}\) is the particle size of the template SiO₂ colloid. In this study, two specimens were fabricated for each \(d_{SiO2}\) in order to check the reproducibility. They are distinguished with a suffix “-1” or “-2”.

The residual SiO₂ in the nanoporous carbons was confirmed by thermalgravimetry and
differential thermal analysis (TG-DTA) on a TG/DTA6300 (Seiko Instruments Inc., heating rate: 10 K min$^{-1}$ in an air flow). To investigate carbon structure, X-ray diffraction (XRD) patterns were recorded on a RINT-2200 (Rigaku, with irradiation of Ni-filtered CuKα) and Raman spectra were measured on an RMP-210 (JASCO, with a 532-nm laser). The porous structure of carbons was observed by transmission electron microscopy (TEM, JEOL JEM-2010). Pore parameters were obtained by the analysis of nitrogen adsorption-desorption isotherms recorded at 77 K on a BELSORP-mini (BEL Japan, Inc.). Total specific surface area ($S_{\text{total}}$) was determined by analyzing $\alpha_{\text{SPE}}$-plots based on the subtracting pore effect (SPE) method. The specific surface area of meso- and macropores ($S_{\text{meso}}$) was analyzed by using $t$-plots. The specific surface area of micropores ($S_{\text{micro}}$) was estimated by subtracting $S_{\text{meso}}$ from $S_{\text{total}}$. Pore size distribution was obtained from the adsorption branches of the isotherms using the Barrett-Joyner-Halenda (BJH) method for mesopores and the Horvath-Kawazoe (HK) method for micropores.

Electrochemical properties were examined in a 1 M LiClO$_4$ solution in a mixed solvent of propylene carbonate (PC) and dimethoxy ethane (DME) (1:1 by volume) in a potential range from 2.0 to 4.0 V vs. Li/Li$^+$ at 25 °C by using three-electrode cells. Cyclic voltammetry and galvanostatic charge/discharge tests conducted on electrochemical analyzers S1470E (Solartron Analytical Inc.) and HJ-SM8 (Hokuto Denko Corp.), respectively. The charge/discharge tests employed the constant-current and subsequent constant-voltage
(CC-CV) mode. Electric double layer capacitance ($C_{DL}$) of nanoporous carbons was obtained as the differential capacitance from the slope of the galvanostatic charge/discharge tests. For the electrochemical measurements, mixtures of the porous carbons and poly(tetrafluoroethylene) (10:1 by mass) that were pressed onto Ni nets served as working electrodes. Both reference and counter electrodes employed metallic lithium pressed on Ni nets. The reference electrode was placed about 0.5-1.0 mm away from the working electrode by using Luggin-capillary. The working electrodes were immersed in the electrolyte in vacuo beforehand. The cell was assembled in an Ar-filled glove box in which $H_2O$ concentration was kept below 1 ppm.

3. Results and Discussions

TG analysis demonstrated that residual SiO$_2$ in the porous carbons was less than 3 %. According to elemental analysis, atomic ratio of H to C of the porous carbons was about 0.1-0.15, and was much smaller than that of original phenolic resin (0.93), which indicates the progress of the carbonization. In Raman spectra (not shown), two peaks were observed around 1,340 and 1,580 cm$^{-1}$ for all the porous carbons, which are the so-called D-band and G-band of carbonaceous materials, respectively. In XRD profiles, very broad peaks appeared around 25º and 44º due to the poor crystallinity, which are typical of so-called hard carbons.

Figure 1 shows that the $N_2$ adsorption/desorption isotherms of the porous carbons $C[d_{SiO2}]$. 
All the isotherms exhibited steep increases of \( N_2 \) uptake at relative pressures above 0.6, and the onset pressure of the increase shifted towards higher relative pressure with increasing \( d_{\text{SiO}_2} \), which indicates that the specimens possess meso/macropore and the pore diameter increases with increasing \( d_{\text{SiO}_2} \). The mesopore size distributions of the porous carbons are plotted in Fig. 2(a), which clearly demonstrated that the pore diameters corresponded to the template \( \text{SiO}_2 \) sizes. TEM images also confirmed that spherical meso/macropores with the same diameter as \( d_{\text{SiO}_2} \) were successfully formed as shown in Fig. 3(a)-(c). Figure 2(b) exhibits the micropore size distributions, which exhibited a peak around 0.7 nm. This peak pore size was independent of \( d_{\text{SiO}_2} \). In a high resolution TEM image (Fig. 3(d)), the micropores smaller than 1 nm were confirmed on meso/macropore walls. These micropores possibly result from the carbonization of the phenolic resin as mentioned in the introduction. Based on these results, the hierarchical porous structure is schematically illustrated in Fig. 4. Specific surface areas of the porous carbons are summarized in table 1. One may expect that the meso/macropore surface area, \( S_{\text{meso}} \), increase with decreasing \( d_{\text{SiO}_2} \), if the carbons were synthesized with a template of closest-packed spherical \( \text{SiO}_2 \) particles. No clear trend of \( S_{\text{meso}} \), however, was obtained, which indicates the disordered meso/macroporous structures for small \( d_{\text{SiO}_2} \), which were observed in TEM images (Fig. 3(c,d)).

Figure 5 displays cyclic voltammograms of the hierarchical porous carbons. The differential capacitance was almost rectangular and independent of electrode potential,
whereas voltammograms in aqueous electrolyte exhibited a peak around potential of zero charge ($E_{pzc}$). This means that no faradaic reaction and no potential dependent ion penetration occur in an organic electrolyte.

Figure 6 displays the charge/discharge curves of the porous carbons obtained by CC-CV mode. For all the porous carbons, almost straight lines were obtained when constant currents were applied. Table 1 summarizes electric double layer capacitance, $C_{DL}$, obtained from slopes of charge curves (2.0 V to 4.0 V vs. Li/Li$^+$). Large $C_{DL}$ of 119 F g$^{-1}$ at 0.05 A g$^{-1}$ was obtained for the porous carbons with $S_{total}$ of 1455 m$^2$ g$^{-1}$ (C[17]-2), which followed the relationship reported by Lozano-Castelló et al.[13] Larger $C_{DL}$ would be expected for a porous carbons with large surface area which would be obtained by optimizing the fabrication procedure. As summarized in table 1, series resistance, $R_s$, which was extrapolated from the charge curves at 5 A g$^{-1}$ increased with decreasing $d_{SiO2}$. $R_s$ is mainly attributed to the electrolyte resistance through the pores. The charge/discharge curves exhibited almost the same slopes, indicating the effective surface areas in double layer capacitance didn’t decreased even at a large current density of 10 A g$^{-1}$. $C_{DL}$ at 5 A g$^{-1}$ was almost the same as that at 0.05 A g$^{-1}$ (see table 1). To the contrary, for typical activated carbons, capacitance decreased to 20-60% of the initial capacitance when current density was increased to 2 A g$^{-1}$.[15] The very good rate capability of the hierarchical porous carbons is attributed to their meso/macropores which facilitate ion transport to whole surfaces. The rate capability of the
EDLC performance of the hierarchical porous carbons will be discussed in detail in a future article.

The contribution of meso/macropores and micropores to $C_{DL}$ were analyzed by using the following equation:[1,9]

$$C_{DL} = c_{dl,meso} \cdot S_{meso} + c_{dl,micro} \cdot S_{micro} \quad (1)$$

or

$$\frac{C_{DL}}{S_{micro}} = c_{dl,meso} \cdot \frac{S_{meso}}{S_{micro}} + c_{dl,micro} \quad (2)$$

where $c_{dl,meso}$ and $c_{dl,micro}$ are specific electric double layer capacitance on meso/macropores and micropores, respectively. In Fig. 7, which plots $C_{DL}/S_{micro}$ versus $S_{meso}/S_{micro}$, a linear relationship was clearly observed. By comparing the line and Eq. (2), $c_{dl,meso}$ and $c_{dl,micro}$ were estimated to be 8.4 $\mu$F cm$^{-2}$ and 8.1 $\mu$F cm$^{-2}$, respectively. It should be noted that $c_{dl,micro}$ is not negligible in the organic electrolyte, and even comparable to $c_{dl,meso}$. On the other hand, in an aqueous electrolyte of 1 M $\text{H}_2\text{SO}_4$, $c_{dl,meso}$ was c.a. 22 and $c_{dl,micro}$ that depended on the electrode potential, $E$, was 5-10 $\mu$F cm$^{-2}$ around $E_{pzc}$ and 0-5 $\mu$F cm$^{-2}$ off $E_{pzc}$.[9] The difference in the capacitance contribution between the aqueous electrolyte and the organic electrolyte might indicate the different electric double layer structures: the thickness of the Helmholtz layer and the diffusion layer, the viscosity of the solvent, and the wettability of surfaces, and so on.

Let us now discuss the size of solvated ions and pores. In the cyclic voltammograms (Fig.
the differential capacitance was slightly higher for potential range $E > E_{pzc}$ ($\sim 3.0$ V vs. Li/Li⁺) than $E < E_{pzc}$. For the former potential range, perchlorate ions are mainly accumulated in double layer whereas for the latter, lithium ions are attracted to electrode surfaces. One might suppose that this is caused by the ion sieving effect by micropores,[16] i.e., the Stokes’ radius of lithium ions (0.41 nm in PC) is larger and that of perchlorate ions (0.26 nm) is smaller than the peak radius of the micropores (0.35 nm),[17] and thus, lithium ions cannot form double layers in micropores. In this case, the capacitance would be related to micropore ratio ($S_{\text{micro}}/S_{\text{total}}$). However, this model is denied by the fact that $S_{\text{micro}}/S_{\text{total}}$ was distributed in the range from 25 to 60% for all the porous carbons, whereas the capacitance difference $C_{DL}(E < E_{pzc})$ and $C_{DL}(E > E_{pzc})$ was c.a. 25% and less dependent on the specimens. The capacitance difference would be simply explained by the thickness of the Helmholtz layer: the Helmholtz layer is thinner and specific capacitance is larger for $\text{ClO}_4^-$ than for Li⁺. This means the ions can be accumulated even in the micropores that are smaller than their sizes. These results agree with that reported by Chmiola et al: in small pores the compressed electric double layer is formed with distorted ions.[14] According to them, $C_{DL}$ for such compressed double layer exhibit larger values than in larger pores without double layer compression, although $c_{dl,\text{micro}}$ was comparable to $c_{dl,\text{meso}}$ in this work. There may be two plausible origins of the distorted ions: distorted structure of solvated Li⁺ ion, or partially desolvated Li⁺ ion. Electrochemical impedance spectra (EIS, not shown) support the distorted structure of
solvated Li\(^+\). According to Ue, Li\(^+\) in PC is supposed to form solvated ion with surrounding PC molecules due to its large charge density, whereas solvation of ClO\(_4\)\(^-\) is almost negligible.[17] In addition, it is reported that partial desolvation of lithium ions require an additional process.[18,19] However, EIS obtained above and below \(E_{pzc}\) (corresponding to accumulation of ClO\(_4\)\(^-\) and Li\(^+\), respectively) did not exhibited obvious difference, which means that no additional process occurs when Li\(^+\) enters micropores. On the other hand, as for aqueous solution, partial desolvation is reported.[20] However, water is different from organic solvents because water molecules tend to form a highly ordered structure in the carbon nanospaces.[21] Capacitance in compressed double layer would be further investigated in a future study by using different electrolytes with various ion sizes or by using porous carbons with different micropore sizes.

4. Conclusions

In this paper, we investigated electric double layer capacitance of the hierarchical porous carbons in an organic electrolyte. The porous carbons exhibited large specific double layer capacitance of 120 F g\(^{-1}\) in an organic electrolyte of a 1 M LiClO\(_4\)/(PC+DME) due to their large surface areas, and superb rate capability resulting from interconnected hierarchical porous structures. The results satisfy the demands for EDLCs for high-power applications such as HEVs. The hierarchical porous structures also revealed the contribution of
meso/macropores and micropore to the electric double layer capacitance, separately. The specific capacitance of meso/macropores and micropores were 8.4 and 8.1 μF cm⁻², respectively. The results indicated electric double layer is formed even when solvated ions are larger than pore diameters.

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References


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Figure captions

Figure 1. N₂ adsorption/desorption isotherms at 77 K of the porous carbons. Solid and open symbols are of adsorption and desorption branches, respectively.

Figure 2. Pore size distribution of (a) mesopores and (b) micropores obtained by the BJH and HK method, respectively.

Figure 3. TEM images of the porous carbons: (a) C[110]-1, (b) C[17]-1 and (c) C[8]-1. (d) A high resolution image of C[17]-1.

Figure 4. Schematic hierarchical porous structure of the porous carbons synthesized by the colloidal-crystal-templating method.

Figure 5. Cyclic voltammograms of the hierarchical porous carbons.

Figure 6. Charge/discharge curves of the hierarchical porous carbons obtained by CC-CV mode at current densities of 0.05 to 10 A g⁻¹.

Figure 7. Plots of $C_{DL}/S_{micro}$ vs. $S_{meso}/S_{micro}$. 
Table 1. Specific surface areas and specific electric double layer capacitance ($C_{DL}$) of the porous carbons. (a) Total surface area, $S_{total}$, and meso-/macropore surface area, $S_{meso}$, were obtained from an $\alpha_{SPE}$-plot and t-plot, respectively. Micropore surface area, $S_{micro}$, was calculated by subtracting $S_{meso}$ from $S_{total}$. (b) $R_s$, electric series resistance, was estimated by extrapolation of the charge curves at 5 A g$^{-1}$. (c) $C_{DL}$, electric double layer capacitance, was obtained from the slope of the charge curves at 0.05 A g$^{-1}$ and 5 A g$^{-1}$.

<table>
<thead>
<tr>
<th></th>
<th>Specific surface area (m$^2$ g$^{-1}$)</th>
<th>$R_s$ (Ω g$^{-1}$)</th>
<th>$C_{DL}$ (F g$^{-1}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>$S_{total}$ $S_{meso}$ $S_{micro}$</td>
<td>$@5$ A g$^{-1}$</td>
<td>$@0.05$ A g$^{-1}$</td>
</tr>
<tr>
<td>C[110]-1</td>
<td>1230 456 774</td>
<td>0.031</td>
<td>107</td>
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<tr>
<td>C[45]-1</td>
<td>1409 1060 349</td>
<td>0.050</td>
<td>116</td>
</tr>
<tr>
<td>C[45]-2</td>
<td>1204 711 493</td>
<td>-</td>
<td>96</td>
</tr>
<tr>
<td>C[17]-1</td>
<td>1302 801 501</td>
<td>0.053</td>
<td>104</td>
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<tr>
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<td>1455 986 469</td>
<td>-</td>
<td>119</td>
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<td>95</td>
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<tr>
<td>C[8]-2</td>
<td>1089 791 298</td>
<td>-</td>
<td>94</td>
</tr>
</tbody>
</table>
Fig. 1

$N_2$ uptake (cm$^3$ g$^{-1}$ @STP)

Relative pressure, $p / p_0$

C[8]-1
C[17]-1
C[45]-1
C[110]-1
Fig. 2a
Fig. 2b
Fig. 3
Fig. 4
Fig. 5
Fig. 6