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## Temperature-dependent tailoring of the pore structure based on MOF-derived carbon electrodes for electrochemical capacitors†

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The pore structures of carbon play a critical role in the charge storage process of electrochemical capacitors; however, the involvement of other varying characteristics, such as electrical conductivity and surface functionalities, complicate the research of the pore size effects on various electrochemical phenomena. In this study, by carbonizing MOF-5 at a selected temperature range of 500–700 °C, a series of MOF-derived carbon materials were obtained with pore size distribution concentrated in different size ranges while admitting similar results in the graphitization degree and surface functionalities. The related morphological changes of ZnO were systematically investigated by changing the carbonization temperature and dwelling time, demonstrating a “from thin to thick, from inside to outside” growth routine of ZnO crystals. With the pore size approximated as the sole variable, the assembled electrochemical capacitors present a linear relationship between the 1–10 nm pores and the impedance resistance, which for the first time demonstrate how 1–10 nm pores is beneficial to ion diffusion. The results of this study not only provide a useful approach to manipulating the pore structure in carbon electrodes but also pave the way to establish the numerical relationship between the pore structure and various phenomena in electrochemistry or other related areas.

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### 1. Introduction

Electrochemical capacitors (ECs), as an efficient tool for energy storage and management with high power density and reliable cyclability, are one of the promising solutions for achieving the stable and sustainable use of renewable energy sources.<sup>1,2</sup> Carbons (activated carbon, aerogels, xerogels, carbon nanotubes, and graphene) are the primary materials used to prepare EC electrodes owing to their chemical stability, natural abundance, and feasibility by engineering their structures/textures and surface chemistry to adapt to various application scenarios.<sup>2–5</sup> The electrochemical performance of carbon materials depends on various material characteristics, such as the pore size structure, structural crystallinity, and surface functionalities.<sup>2,3,6–9</sup> Adjusting the pore structure is a common and effective way to optimize the electrochemical performance of carbon-based ECs, especially electrochemical double-layer capacitors, where the properties of the as-formed double layer structure depend greatly on the pore size and curvature with respect to the chosen electrolytic ions.<sup>10–12</sup>

Template carbonization is one of the effective methods for controlling the pore size of carbon materials and is classified into hard- and soft-templating methods.<sup>6,13–15</sup> Metal-organic frameworks, composed of metal units and organic ligands with intrinsic porosity, are ideal soft-templating precursors for the construction of porous carbon materials,<sup>15–17</sup> eliminating the use of dangerous and toxic hydrogen fluoride usually required in the hard-templating methods.<sup>13,18</sup> Moreover, carbons produced by hard-templating methods are usually advanced with a narrow pore size distribution,<sup>6,13,18</sup> while in practice especially for industrialized products, carbon materials with a wide pore size distribution and hierarchical porous structure are the common case.<sup>19</sup>

So far, there has been extensive research on synthesizing porous carbons using MOFs as precursors ever since their first report in 2008 by Liu *et al.*,<sup>17</sup> utilizing Zn-based MOF-5 (consists of the metal unit Zn<sub>4</sub>O linked to the organic ligand *p*-benzenedioic acid),<sup>20</sup> to prepare nanoporous carbon at 530 °C with ZnO removed by acid washing, and at 1000 °C (boiling point of Zn metal 908 °C), showing excellent hydrogen absorption capacity and electrochemical properties. Further research by Zhang *et al.* revealed that the thermal decomposition of MOF-5 occurred at 400 °C and above involved breaking the carboxyl bridge between the benzene ring and the Zn<sub>4</sub>O cluster and producing amorphous carbon along with CO<sub>2</sub>, benzene, and ZnO crystals.<sup>21</sup> Effects of carbonization temperature on the properties of MOF-derived carbons have been repeatedly

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demonstrated where the resulting pore structure concentrated in the mesoporous range at temperature as low as 500 °C or higher than 700 °C;<sup>22,23</sup> a morphological change of ZnO was also observed upon increasing the carbonization temperature from 600 °C to 1000 °C,<sup>23,24</sup> showing that the conversion of Zn<sub>4</sub>O clusters to ZnO is quite sensitive to the carbonization temperature and demonstrating the possibility to control the pore size distribution through controlling the morphology of ZnO.

Herein, in order to locate the proper treatment conditions for tailoring the pore size distribution, the effects of carbonization temperature as well as the dwelling time have been systematically investigated using MOF-5 as the precursor. Varied pore size distributions were achieved with other critical features affecting the electrochemical performance changing little between 570 °C to 700 °C. Based on the obtained MOF-derived carbon with pore size distribution approximated as the sole variable, a linear relationship has been revealed between the 1–10 nm pores and the impedance resistance, which would offer fundamental insight into the pore size-ion diffusion relation for developing advanced electrochemical devices.

## 2. Experimental methods

### 2.1 Preparation of Zn-MOF-5

MOF-5 was synthesized following the method first reported by Yaghi *et al.*<sup>20</sup> 7.76 mmol of zinc acetate dihydrate (Zn(CH<sub>3</sub>-COO)<sub>2</sub>·2H<sub>2</sub>O, Alfa Aesar, 98%) was dissolved in a conical flask containing 50 ml of *N,N*-dimethylformamide (DMF, Alfa Aesar, 99.8%); 3.05 mmol of terephthalic acid (1,4-H<sub>2</sub>BDC, Alfa Aesar, 98%) was dissolved in 40 ml of DMF with 0.85 ml of triethylamine (TEA, Alfa Aesar, 99%) added afterwards. Then, the two solutions were mixed together under stirring at 600 rpm for 2.5 h. White precipitates of Zn-MOF-5 were washed three times with anhydrous ethanol (Sinopharm, AR grade), collected by centrifugation, and dried under vacuum at 80 °C for 3 hours.

### 2.2 Carbonization of Zn-MOF-5

The as-synthesized Zn-MOF-5 was loaded into a ceramic boat and placed in a quartz tube under Ar flow. The temperature was first raised from room temperature to 400 °C at a heating rate of 10 °C min<sup>-1</sup> and then dwelled for 1 h. Afterwards, the temperature was increased at a heating rate of 5 °C min<sup>-1</sup> to the set target temperature *T* and dwelling for a set period of time *t* (Table S1†). The carbonized MOFs were labeled as ZnO/C-*T<sub>x</sub>-t<sub>y</sub>* (where *x* is the target temperature, and *y* is the dwelling time). ZnO/C-*T<sub>x</sub>-t<sub>y</sub>* were then soaked in 6 M HCl (aq) at room temperature for 24 h to remove ZnO, followed by washing with copious deionized water until reaching neutral (pH ~ 7) and then dried at 100 °C for 12 h. The as-produced MOF-5-derived carbons were denoted as C-*T<sub>x</sub>-t<sub>y</sub>*.

### 2.3 Material characterization

Thermogravimetric analysis (TGA) was carried out using a TA instrument (HR-1) from room temperature to 1000 °C at a heating rate of 10 °C min<sup>-1</sup> under nitrogen protection. X-ray diffraction (XRD) data were collected on a Rigaku SmartLab

diffractometer (D/MAX- $\gamma$ A,  $\lambda$  = 1.54 Å). The scanning electron microscopic (SEM) images were obtained on a Zeiss Crossbeam 550, and the transmission electron microscopic (TEM) images, scanning transmission electron microscopic (STEM) images, and the corresponding elemental mapping results on a JEM-F200. Raman spectra were collected using a Renishaw InVia Reflex confocal microscope with an excitation light source of 532 nm. The laser power used was 0.15 mW to minimize burning the samples. X-ray photoelectron spectroscopy (XPS) spectra were collected on an ESCALAB 250Xi and calibrated by setting the C 1s photoemission peak for sp<sup>2</sup>-hybridized carbons to 284.8 eV. N<sub>2</sub> adsorption/desorption isotherms were acquired using a Quadasorb Instrument (Quantachrome, Autosorb IQ-MP-MP) at 77 K, where the surface area and pore size distributions were calculated by the quenched solid density functional theory (QSDFT) method.

### 2.4 Electrochemical tests

85 wt% C-*T<sub>x</sub>-t<sub>y</sub>*, 6 wt% carbon black, and 9 wt% polyvinylidene fluoride (PVDF, Arkema-HSV900, 99%) binder were mixed and ground with a mortar and pestle in *N*-methylpyrrolidone (NMP, Alfa Aesar, 99%). The slurry was then coated onto circular disks of aluminum foil (14 mm in diameter). The mass loading of each electrode was around 1 mg. ECs were assembled in a coin cell (CR2032-SUS316L, Japan). A Whatman glass microfiber filter was used as the separator immersed in 1 M tetraethylammonium tetrafluoroborate (Alfa Aesar, 99%)-polycarbonate (Alfa Aesar, 99%) (TEABF<sub>4</sub>/PC) as the organic electrolyte.

The cyclic voltammetry (CV) and electrochemical impedance spectroscopy (EIS) measurements were performed on a Solartron 1470E electrochemical testing system. CV was performed at scan rates ranging from 2 to 500 mV s<sup>-1</sup> within a voltage window of 0 to 2 V. EIS was recorded at 0 V over a frequency range from 100 kHz to 10 mHz with a bias amplitude of 5 mV. Galvanostatic charge-discharge (GCD) was measured using an Arbin instrument (LBT-21084) from 0 to 2 V at a series of current densities from 10 to 1000 mA g<sup>-1</sup>. The specific capacitances *C* (F g<sup>-1</sup>) based on CV tests were calculated using the following equation:

$$C = \frac{\int I dV}{\nu m \Delta V} \quad (1)$$

where *I* (A) is the current, *ν* (V s<sup>-1</sup>) is the scan rate, *m* (g) is the weight per electrode, and  $\Delta V$  (V) is the applied voltage window (2 V).

The specific capacitance based on the GCD tests was calculated based on the following equation:

$$C = \frac{2I\Delta t}{m\Delta V'} \quad (2)$$

where *m* (g) is the weight per electrode, *I* (A) is the discharge current,  $\Delta t$  (s) is the discharge time, and  $\Delta V'$  (V) is the potential change within  $\Delta t$  excluding the voltage drop.

## 3. Results and discussion

Fig. 1a schematically presents the synthesis procedure of MOF-5-derived carbon. In order to obtain porous carbon with

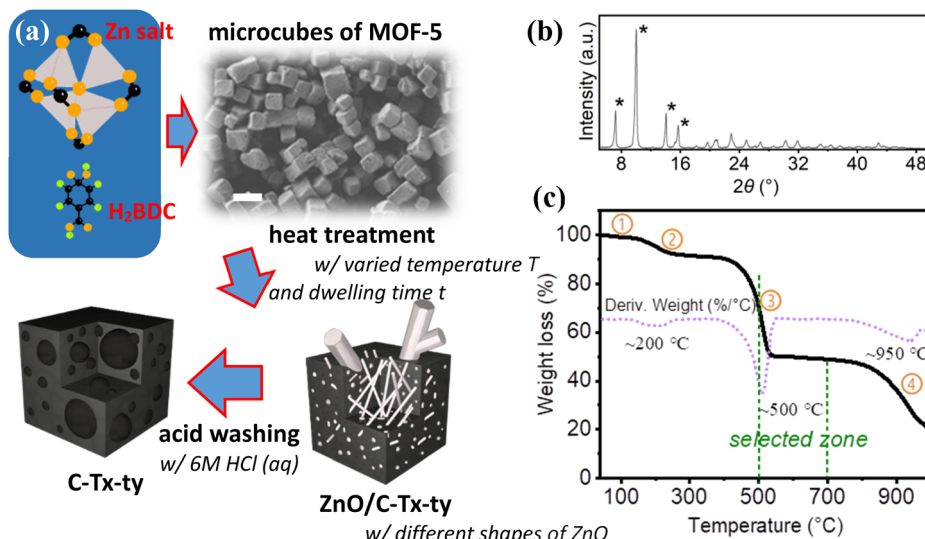


Fig. 1 (a) The schematic illustration of the synthesis of MOF-derived carbon  $C-T_x-t_y$ . (b) XRD pattern with four characteristic peaks at  $6.9^\circ$ ,  $9.8^\circ$ ,  $13.8^\circ$ , and  $15.4^\circ$  corresponding to (200), (220), (400), and (420) of Zn-MOF-5, respectively.<sup>21,23</sup> (c) TGA curve of Zn-MOF-5 under nitrogen atmosphere (the purple dotted line indicates the differential result of TGA curve).

different pore size distributions, the synthesized microcubes of Zn-MOF-5 (identified with XRD result, Fig. 1b) were treated with different carbonization temperatures  $T$  and dwelling time  $t$  to control the morphology of the ZnO formed. The temperature was chosen based on TGA results. As shown in Fig. 1c, the weight loss of Zn-MOF-5 during the heating from room temperature to  $1000^\circ\text{C}$  could be divided into four main stages: (1) the first stage (weight loss:  $\sim 1\%$ ) as located below  $\sim 110^\circ\text{C}$  could be ascribed to the removal of water adsorbed on the surface of the material;<sup>21,24,25</sup> (2) the second stage (weight loss:  $\sim 9\%$ ) between  $\sim (120\text{--}300)^\circ\text{C}$  was mainly due to the dissipation of the residual solvent molecules (DMF here) incorporated in the frameworks;<sup>24,25</sup> (3) significant weight loss ( $\sim 40\%$ ) occurred around  $\sim (400\text{--}550)^\circ\text{C}$  of the third stage, which should be caused by the thermal decomposition of the Zn-MOF-5 skeleton. During this stage, carboxylic bridges between benzene rings and  $\text{Zn}_n\text{O}$  clusters broke, releasing  $\text{CO}_2$  and benzene, and Zn-MOF-5 crystals were pyrolyzed into a composite of ZnO and carbon;<sup>21,22,24</sup> (4) as the temperature continued to reach around  $750^\circ\text{C}$ , carbonaceous materials started to deoxidize ZnO to Zn with itself being oxidized mainly into  $\text{CO}_2$  and  $\text{CO}$ ;<sup>22,24</sup> when reaching the boiling point of the zinc metal monomer ( $908^\circ\text{C}$ ), the gaseous zinc would begin to evaporate under the nitrogen current flow. This reaction was the most intense at  $950^\circ\text{C}$ ,<sup>22,24</sup> and the weight loss of this stage was  $\sim 28\%$ . Therefore, the temperature zone to tune the ZnO morphology was selected between  $(500\text{--}700)^\circ\text{C}$ , where carbon was produced from the decomposition of organic components but little consumption of carbon to reduce ZnO.

As shown in Fig. 2a-g, the carbonized MOFs (named  $\text{ZnO/C-T}_x-t_y$ , where  $x$  denotes the carbonization temperature and  $y$  the dwelling time) exhibited a morphological transition of ZnO crystals (as demonstrated with the XRD results, Fig. 2h) upon increasing the temperature and the dwelling time. Besides, the

carbon cubes presented different morphologies as well. All  $\text{ZnO/C-T}_x-t_y$  samples retained the cubic skeleton without collapse. Differences in the ZnO crystals are reflected not only in the shape and size but also in the location.  $\text{ZnO/C-T}_{500}-t_{0.5}$  treated at the lowest  $500^\circ\text{C}$  showed a sponge-like porous structure (Fig. 2a, the SEM) with fine ZnO nanoparticles ( $\sim 10\text{--}20\text{ nm}$  in diameter) and nanoneedles ( $\sim 10\text{--}30\text{ nm}$  in diameter and  $\sim 300\text{--}500\text{ nm}$  in length) embedded inside the cube and some protruding out (Fig. 2a, TEM) since the lower temperature is prone to the formation of fine ZnO nanoparticles/nanoneedles.<sup>26</sup> On prolonging the treatment time to 2 h,  $\text{ZnO/C-T}_{500}-t_2$  exhibited a much smoother surface with some ZnO aggregates grown onto the external surface (Fig. 2b, SEM); nanoneedles disappeared and were replaced by nanospheres ( $\sim 10\text{--}20\text{ nm}$  in diameter, Fig. 2b, TEM).<sup>27</sup>  $\text{ZnO/C-T}_{570}-t_{0.5}$  with the higher temperature at  $570^\circ\text{C}$  but a shorter dwelling time of 0.5 h presented a sponge-like porous structure similar to that of  $\text{ZnO/C-T}_{500}-t_{0.5}$ , and were filled with ZnO nanoparticles ( $\sim 10\text{--}20\text{ nm}$  in diameter) and nanoneedles as well ( $\sim 10\text{--}30\text{ nm}$  in diameter and  $\sim 300\text{--}500\text{ nm}$  in length, Fig. 2c); further increasing the dwelling time at this temperature to 2 h and 4 h lead to  $\text{ZnO/C-T}_{570}-t_2$  and  $\text{ZnO/C-T}_{570}-t_4$ , respectively, both exhibiting similar morphology to that of  $\text{ZnO/C-T}_{500}-t_2$  enriched with ZnO nanospheres ( $\sim 30\text{--}60\text{ nm}$ , Fig. 2d and e). As the temperature was further raised to  $700^\circ\text{C}$ , nearly all ZnO was extracted out of the cubic frameworks and grew into large ZnO rods ( $\sim 100\text{--}200\text{ nm}$  in diameter), possibly as a result of that temperature up to  $700^\circ\text{C}$  would melt the nanospheres and favor the growth of nanorods due to the carbon reduction of ZnO, where the resulting gaseous zinc reacted with the fine unreduced ZnO nanoneedles/nanospheres within the cubes to form ZnO rods that extended outwards.<sup>23,24,28</sup>  $\text{ZnO/C-T}_{700}-t_{0.5}$  and  $\text{ZnO/C-T}_{700}-t_2$  both presented clean structures inside and smooth surfaces except for some large holes probably marked



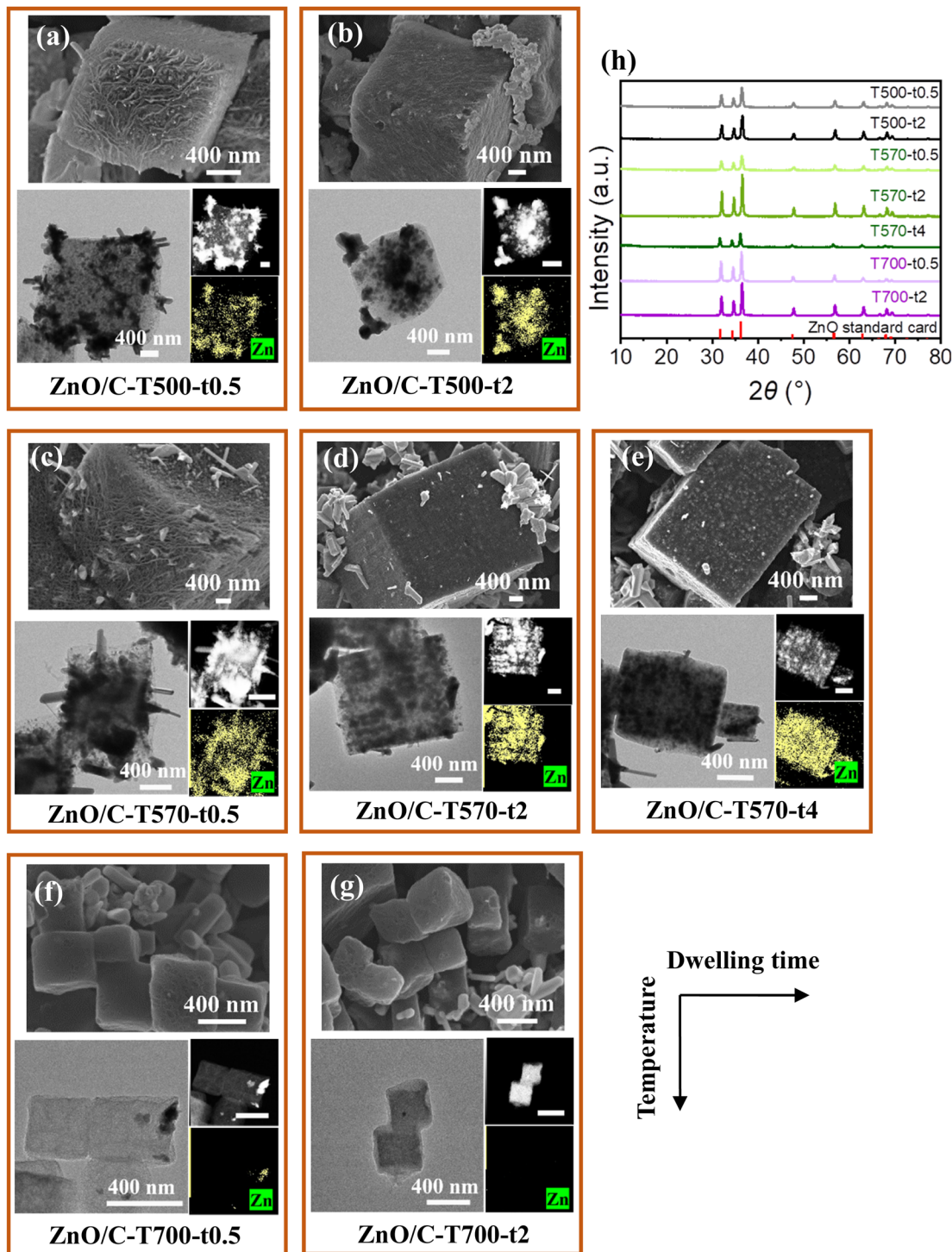


Fig. 2 SEM, TEM, and STEM images with elemental mapping results of (a) ZnO/C-T<sub>500</sub>-t<sub>0.5</sub>, (b) ZnO/C-T<sub>500</sub>-t<sub>2</sub>, (c) ZnO/C-T<sub>570</sub>-t<sub>0.5</sub>, (d) ZnO/C-T<sub>570</sub>-t<sub>2</sub>, (e) ZnO/C-T<sub>570</sub>-t<sub>4</sub>, (f) ZnO/C-T<sub>700</sub>-t<sub>0.5</sub>, and (g) ZnO/C-T<sub>700</sub>-t<sub>2</sub>. (h) XRD patterns of ZnO/C-T<sub>x</sub>-t<sub>y</sub> show distinct peaks at 31.6°, 34.5°, 36.2°, 47.5°, 56.6°, 62.8°, 66.3°, 67.9° and 69°, representing ZnO.<sup>23,27</sup> Note: the scale bar in the STEM images represents 500 nm.

by the ZnO rods (Fig. 2f and g). Conclusively, within the (500–700) °C temperature range, the morphology of ZnO could be controlled from nanoneedles to nanospheres, and eventually diffusing out of the cubic frameworks and forming rod-like ZnO crystals by increasing the carbonization temperature and adjusting the dwelling time.

After acid washing, Zn-MOF-5 derived carbons (Fig. 3a–g, denoted as C-T<sub>x</sub>-t<sub>y</sub>) were easily obtained. C-T<sub>x</sub>-t<sub>y</sub> inherited the cubic shape and the morphology of the precedent ZnO/C-T<sub>x</sub>-t<sub>y</sub> with ZnO being completely removed (as evidenced with only broad XRD peaks at ~23° and ~44° for amorphous carbon, Fig. 3h).<sup>17</sup> C-T<sub>500</sub>-t<sub>0.5</sub> and C-T<sub>570</sub>-t<sub>0.5</sub> exhibited sponge-like

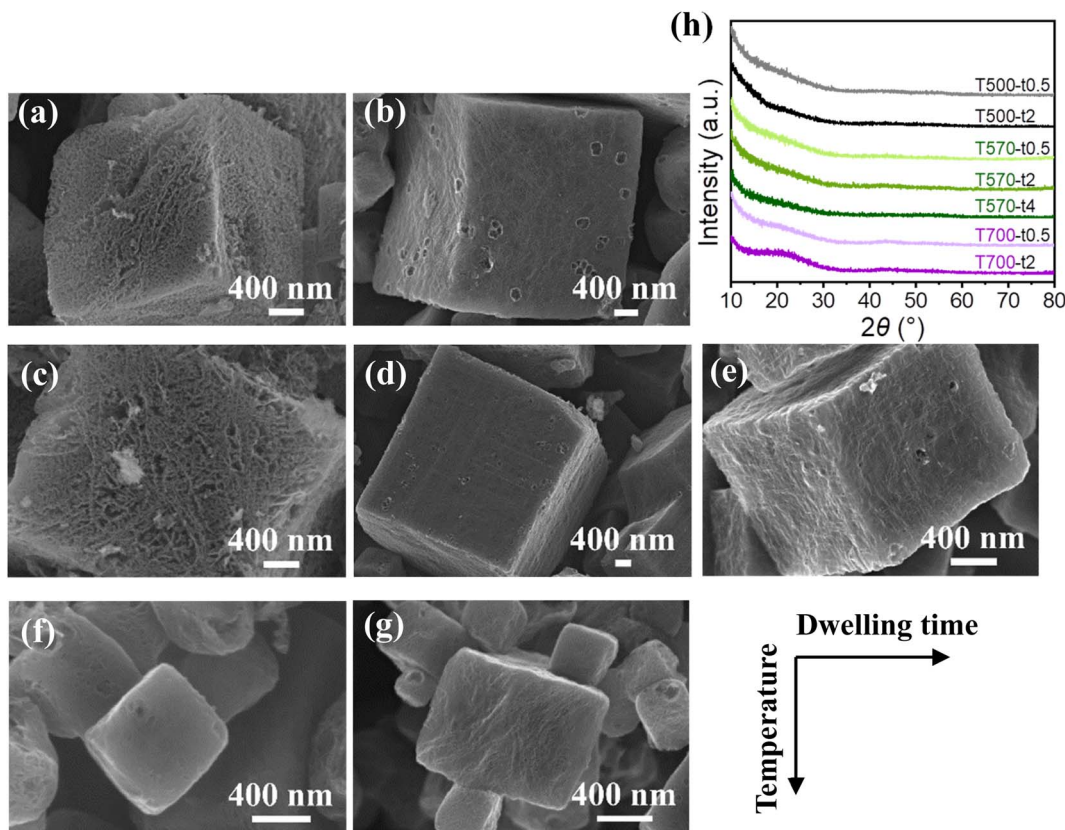


Fig. 3 SEM images of (a)  $C-T_{500-t0.5}$ , (b)  $C-T_{500-t2}$ , (c)  $C-T_{570-t0.5}$ , (d)  $C-T_{570-t2}$ , (e)  $C-T_{570-t4}$ , (f)  $C-T_{700-t0.5}$ , and (g)  $C-T_{700-t2}$ . (h) XRD patterns of  $C-T_x-t_y$ .

porous structures while the rest showed a relatively smoother surface. TEM images of the  $C-T_x-t_y$  presented distinct pore structures, where  $C-T_{500-t0.5}$  and  $C-T_{570-t0.5}$  were filled with large pores (up to 20–30 nm in diameter) while pores in  $C-T_{700-t0.5}$  and  $C-T_{700-t2}$  were too small to be appreciable (Fig. S2†).

Further analysis with  $N_2$  adsorption–desorption measurement demonstrated that the varied pore size distribution was achieved by controlling the morphology of ZnO. The QSDFT model was applied as an effective method for the pore size distribution calculation in micro-mesoporous carbons with heterogeneous structures.<sup>29–31</sup> The  $N_2$  adsorption–desorption isotherms and the corresponding pore size distribution are given in Fig. 4a–f.  $C-T_{500-t0.5}$  and  $C-T_{570-t0.5}$  not only presented similar textures of carbon microcubes (Fig. 3a and c) but also admitted similar isotherms (as a combination of type I (micropore), type IV (mesopore), and type II (macropore) isotherms, Fig. 4a and b), suggesting a hierarchical porous structure with a wide pore size distribution (Fig. 4d and e).<sup>32</sup> The sharp uptakes at a low relative pressure ( $p/p_0 < 0.1$ ) indicated the existence of micropores and the hysteresis loop in the region ( $p/p_0 \sim 0.5$  to 0.9) the presence of mesopores; the sharp uptakes at a high relative pressure ( $p/p_0 > 0.9$ ) indicated the presence of a large amount of macropores.<sup>33,34</sup> The rest five  $C-T_x-t_y$  treated either at higher carbonization temperature or longer dwelling time also possessed hierarchical porous structures as indicated by a combination of type I and type IV isotherms (Fig. 4a–c),

where the upward tail at a high relative pressure ( $p/p_0 > 0.9$ ) disappeared except for a small tail of  $C-T_{700-t2}$ . Hence, within the temperature range (500–700) °C, the higher the carbonization temperature or a longer period of dwelling time, the fewer macropores existed; while at the upper-temperature limit of 700 °C, with the longer dwelling time of 2 h, macropores started to show up again.<sup>23</sup>

The analyzed textural characteristics of the  $C-T_x-t_y$  carbons are summarized in Tables 1 and 2, where the carbon materials presented a significant increase in specific surface area (SSA) as compared to that of the Zn-MOF-5 (only 897  $m^2 g^{-1}$ , Fig. S3 and Table S3†). To further analyze the porous structure with respect to its effects on electrochemical performance, the pores were divided into three categories based on the electrolyte used in the following study (1 M TEABF<sub>4</sub>/PC, where the ion size of TEA<sup>+</sup> is 0.68 nm, and 1.30 nm with the solvated shell, and the ion size of BF<sub>4</sub><sup>–</sup> is 0.48 nm and 1.16 nm with the solvated shell,<sup>11,35</sup>), which were small pores of (<1 nm), medium pores of (1–10) nm, and large pores of (>10 nm). Desolvation or partial desolvation is required for electrolytic ions to diffuse into the small pores;<sup>11,35</sup> for the medium pores the pore curvature is significant while for the large pores the pore wall can be approximated as a planar surface with respect to the ions.<sup>12</sup>

A further comparison of the pore size distribution based on the above three categories (Tables 1 and 2) showed that  $C-T_{500-t0.5}$  and  $C-T_{570-t0.5}$  both held a considerable amount of large

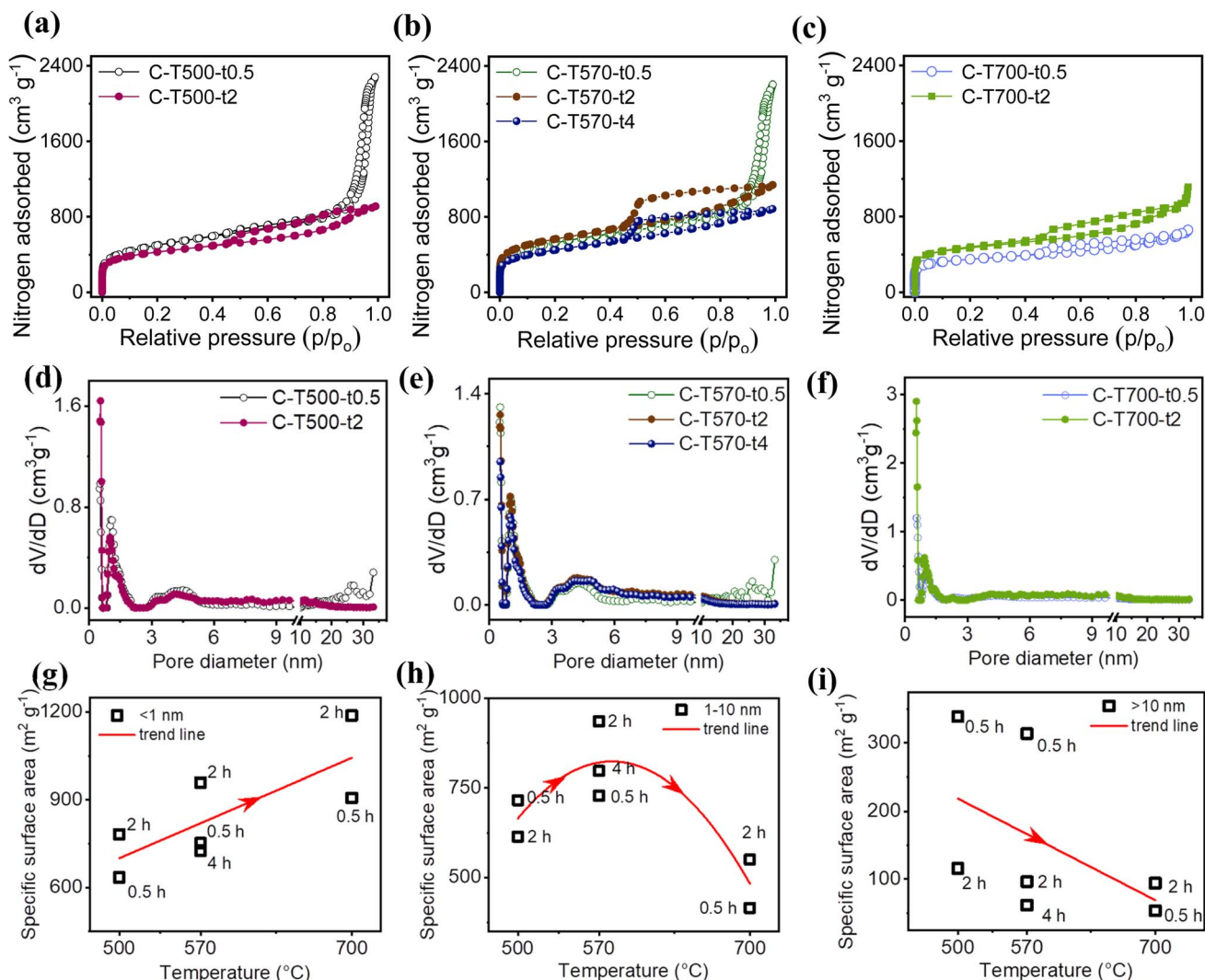


Fig. 4 N<sub>2</sub> adsorption–desorption isotherms and pore size distribution of (a and d) C-T<sub>500</sub>-t<sub>0.5</sub> and C-T<sub>500</sub>-t<sub>2</sub>, (b and e) C-T<sub>570</sub>-t<sub>0.5</sub>, C-T<sub>570</sub>-t<sub>2</sub> and C-T<sub>570</sub>-t<sub>4</sub>, and (c and f) C-T<sub>700</sub>-t<sub>0.5</sub>, C-T<sub>700</sub>-t<sub>2</sub>. Relationship between carbonization temperature and the partial SSA contributed by (g) (<1 nm) pores, (h) (1–10) nm pores, and (i) (>10 nm) pores in C-T<sub>x</sub>-t<sub>y</sub> materials with trend line.

pores of (>10 nm) while C-T<sub>700</sub>-t<sub>0.5</sub> and C-T<sub>700</sub>-t<sub>2</sub> possessed a fairly large amount of small pores of (<1 nm). Large pores of (>10 nm) took up a pore volume ratio (the partial pore volume  $V_{>10 \text{ nm}}$  contributed by large pores divided by the total pore

volume  $V_{\text{total}}$ ) of 71.8% and 67.5% in C-T<sub>500</sub>-t<sub>0.5</sub> and C-T<sub>570</sub>-t<sub>0.5</sub> (Table 1), hence C-T<sub>500</sub>-t<sub>0.5</sub> and C-T<sub>570</sub>-t<sub>0.5</sub> could be considered predominantly composed of large pores. Small pores of (<1 nm) took up an SSA ratio (the partial SSA  $\text{SSA}_{<1 \text{ nm}}$  contributed by

Table 1 The partial pore volume ( $V_{<1 \text{ nm}}$ ,  $V_{1-10 \text{ nm}}$ , and  $V_{>10 \text{ nm}}$ ) and the corresponding pore volume ratios with respect to three types of pores<sup>a</sup>

Sample	$V_{\text{total}}$ ( $\text{cm}^3 \text{g}^{-1}$ )	$V_{<1 \text{ nm}}$ ( $\text{cm}^3 \text{g}^{-1}$ )	$V_{1-10 \text{ nm}}$ ( $\text{cm}^3 \text{g}^{-1}$ )	$V_{>10 \text{ nm}}$ ( $\text{cm}^3 \text{g}^{-1}$ )	$V_{<1 \text{ nm}}$ (%)	$V_{1-10 \text{ nm}}$ (%)	$V_{>10 \text{ nm}}$ (%)
C-T <sub>500</sub> -t <sub>0.5</sub>	2.84	0.19	0.61	2.04	6.6	21.6	71.8
C-T <sub>500</sub> -t <sub>2</sub>	1.29	0.24	0.63	0.42	18.6	48.8	32.6
C-T <sub>570</sub> -t <sub>0.5</sub>	2.69	0.23	0.64	1.82	8.5	24.0	67.5
C-T <sub>570</sub> -t <sub>2</sub>	1.62	0.29	0.97	0.36	17.9	59.9	22.2
C-T <sub>570</sub> -t <sub>4</sub>	1.27	0.21	0.83	0.23	16.9	65.0	18.1
C-T <sub>700</sub> -t <sub>0.5</sub>	0.90	0.27	0.42	0.21	29.8	46.6	23.6
C-T <sub>700</sub> -t <sub>2</sub>	1.36	0.36	0.64	0.36	26.5	46.9	26.6

<sup>a</sup>  $V_{\text{total}}$  is the total pore volume.

Table 2 The partial SSA ( $S_{<1\text{ nm}}$ ,  $S_{1\text{--}10\text{ nm}}$ , and  $S_{>10\text{ nm}}$ ) and the corresponding SSA ratios with respect to three types of pores

Sample	SSA ( $\text{m}^2\text{ g}^{-1}$ )	SSA <sub>&lt;1 nm</sub> ( $\text{m}^2\text{ g}^{-1}$ )	SSA <sub>1–10 nm</sub> ( $\text{m}^2\text{ g}^{-1}$ )	SSA <sub>&gt;10 nm</sub> ( $\text{m}^2\text{ g}^{-1}$ )	SSA <sub>&lt;1 nm</sub> (%)	SSA <sub>1–10 nm</sub> (%)	SSA <sub>&gt;10 nm</sub> (%)
C- $T_{500-t_{0.5}}$	1689	634	716	339	37.5	42.4	20.1
C- $T_{500-t_2}$	1512	782	614	116	51.7	40.6	7.7
C- $T_{570-t_{0.5}}$	1794	752	728	314	41.9	40.6	17.5
C- $T_{570-t_2}$	1991	959	936	96	48.2	47.0	4.8
C- $T_{570-t_4}$	1585	726	797	62	45.8	50.3	3.9
C- $T_{700-t_{0.5}}$	1376	907	415	54	65.9	30.2	3.9
C- $T_{700-t_2}$	1832	1188	550	94	64.8	30.0	5.2

small pores divided by the total SSA) of 65.9% and 64.8% in C- $T_{700-t_{0.5}}$  and C- $T_{700-t_2}$ , respectively (Table 2), hence C- $T_{700-t_{0.5}}$  and C- $T_{700-t_2}$  could be considered predominantly composed of small pores. As for C- $T_{570-t_4}$ , the medium pores of (1–10) nm took up relatively higher ratios by 50.3% in SSA ratio and 65.0% in pore volume ratio. C- $T_{500-t_2}$  and C- $T_{570-t_2}$  did not present a major type of pores but a relatively evenly distributed combination of different-sized pores. More interestingly, as vividly shown by the trend lines (Fig. 4g–i), SSA<sub><1 nm</sub> increased with the carbonization temperature, while SSA<sub>>10 nm</sub> decreased with the carbonization temperature, and SSA<sub>1–10 nm</sub> reached a maximum at 570 °C. Hence, though there was no clear relation existed between the total SSA and the carbonization temperature, partial SSA did relate to the temperature; based on this, it is possible to realize fine-tuning to desirable pore size distributions by designing the treatment conditions within the temperature range of (500–700) °C.

Besides the pore structures, Raman, and XPS were applied to further analyze the effects of carbonization temperature and dwelling time on the chemical properties of C- $T_x-t_y$ . As shown in Fig. 5a, the Raman spectra of all the C- $T_x-t_y$  materials were fitted by four Gaussian peaks: TPA peak (at  $\sim 1200\text{ cm}^{-1}$ , attributed to heteroatoms), D peak (at  $\sim 1350\text{ cm}^{-1}$ , corresponding to A<sub>1g</sub> vibrations of sp<sup>3</sup> defects), A peak (at  $\sim 1490\text{ cm}^{-1}$ , from defects in graphite layer stacks), and G peak (at  $\sim 1595\text{ cm}^{-1}$ , the vibrations of sp<sup>2</sup> hybridized carbon structures).<sup>36,37</sup> The smaller the intensity ratio of  $I_D/I_G$  (the ratio of the deconvoluted peak areas), the higher the degree of graphitization.<sup>16,37</sup> C- $T_{500-t_{0.5}}$  showed the largest  $I_D/I_G$  of 2.33, indicating the low degree of graphitization probably due to the low carbonization temperature.<sup>38</sup> Increasing the dwelling time to 2 h, the  $I_D/I_G$  of C- $T_{500-t_2}$  decreased to 1.42. The rest of  $I_D/I_G$  for higher carbonization temperatures of 570 °C and 700 °C were all around  $\sim 1.2$ , suggesting no significant change in the crystallinity of the MOF-derived carbons when the temperature fell between 570 °C and 700 °C.

Chemical identification of C- $T_x-t_y$  carbons was addressed by XPS. All survey scans showed a dominant peak around 284.8 eV for C 1s and one around 532.9 eV for O 1s,<sup>16,39</sup> while no peaks were observed for the Zn element (Fig. S4†), again demonstrating that ZnO was completely removed after acid washing. Fitting results of high-resolution scans gave the same combination of deconvoluted peaks (Fig. S5 and S6†). Taking C- $T_{570-t_2}$  as an example (Fig. 5b and c), the high-resolution C 1s region

could be deconvoluted into four peaks: sp<sup>2</sup> carbon ( $\sim 284.8\text{ eV}$ ), sp<sup>3</sup> carbon ( $285.4 \pm 0.3\text{ eV}$ ), C–O(C–OH) ( $286.6 \pm 0.3\text{ eV}$ ) and O=C–O(H<sub>2</sub>O<sub>ads</sub>) ( $289.6 \pm 0.3\text{ eV}$ ),<sup>16,36</sup> while the high-resolution O 1s region could be deconvoluted into two peaks: C–O(C–OH) ( $532.6 \pm 0.3\text{ eV}$ ) and O=C–O(H<sub>2</sub>O<sub>ads</sub>) ( $533.6 \pm 0.3\text{ eV}$ ).<sup>16,39,40</sup> Comparing the calculated atomic ratios of different carbons (Fig. 5d and Table S4†), the ratio of sp<sup>2</sup> carbon, sp<sup>3</sup> carbon, and the functional groups varied little after 570 °C, demonstrating similar chemical compositions. Taking sp<sup>2</sup> carbon as an example, C- $T_{500-t_{0.5}}$  and C- $T_{500-t_2}$  treated at 500 °C admitted lower ratios of only 52.62 at% and 53.33 at%, respectively. After increasing the carbonization temperature to 570 °C, the ratio increased to  $\sim 64$  at% where prolonging the dwelling time from 0.5 h to 4 h affected the ratio little. The sp<sup>2</sup>-C ratio decreased a little to  $\sim 57$  at% after further increasing the carbonization temperature to 700 °C, probably due to the reduction of ZnO that might have taken place at this temperature.

Conclusively, control of the pore size distribution can be achieved by tuning the carbonization temperature and the dwelling time; moreover, as the chemical properties varied little within the temperature range (570–700) °C, pore size distribution could be approximated as a single variable to investigate its effects on various applications.

The electrochemical properties of C- $T_x-t_y$  materials were characterized for EDLC applications using the organic electrolyte 1 M TEABF<sub>4</sub>/PC. As shown in Fig. 6a, C- $T_{500-t_{0.5}}$  showed poor electrochemical performance with significantly distorted CV curves even at a moderate scan rate of 10 mV s<sup>-1</sup>, and the specific capacitance was only 2.1 F g<sup>-1</sup> at 500 mV s<sup>-1</sup>, exhibiting low rate capability (Table S5†). After increasing the dwelling time to 2 h, the CV curves were approaching rectangular with only slight distortion, and the specific capacitance was raised to 15.1 F g<sup>-1</sup> at 500 mV s<sup>-1</sup> with a rate capability of 31.9% for C- $T_{500-t_2}$  (Fig. 6b, and Table S5†). Optimal performance was obtained for C- $T_{570-t_2}$  (Fig. 6d), admitting a specific capacitance of 48.1 F g<sup>-1</sup> at 500 mV s<sup>-1</sup> with a rate capability of 60.2%, while further increasing the carbonization temperature and the dwelling time resulted in a slight reduction of the as-measured specific capacitance (Table S5†). The improved electrochemical performance could be ascribed to the enhanced crystallinity and electrical conductivity at temperature  $\geq 570$  °C (Fig. 6h). The shape of all the CV curves presented a quasi-rectangular geometry (Fig. 6c–g), corroborating the enhanced crystalline structure ( $I_D/I_G \sim 1.2$ ) and chemical composition (sp<sup>2</sup> carbon  $\sim 60$  at%) within the range of



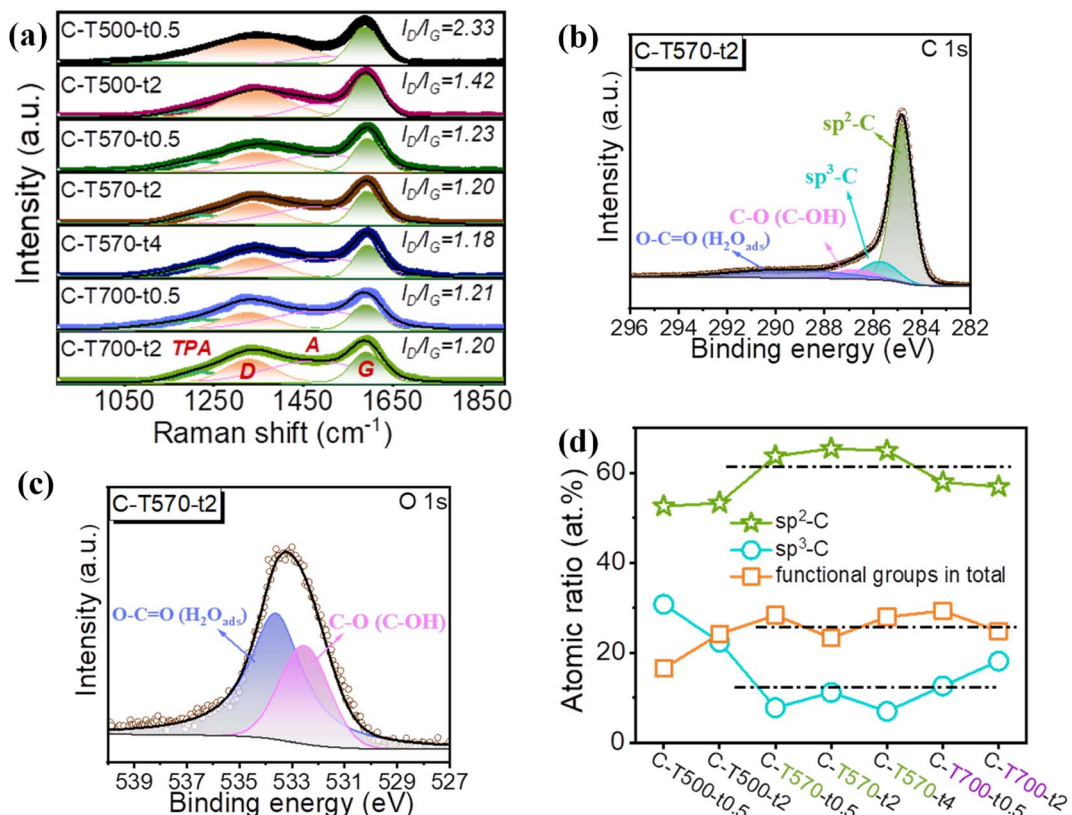


Fig. 5 (a) Raman spectra of  $C-T_x-t_y$ . High-resolution spectra of (b) C 1s, and (c) O 1s of  $C-T_{570}-t_2$ . (d) Comparison of atomic ratios of  $sp^2$  carbon,  $sp^3$  carbon, and functional groups as calculated from the fitting results of C 1s spectra of  $C-T_x-t_y$ .

(570–700) °C. Moreover, there was quite a difference in charge capacitance. Taking the scan rate of 2 mV s<sup>-1</sup> as an example, the specific capacitance admitted the highest 79.9 F g<sup>-1</sup> for  $C-T_{570}-t_2$  and the lowest 56.7 F g<sup>-1</sup> for  $C-T_{700}-t_{0.5}$ . As displayed in Fig. 6i (Fig. S9†), the specific capacitance increased with increasing SSA, hence SSA remained as one of the critical characteristics affecting the electrochemical energy storage in our carbon-based EDLCs mainly depending on the double-layer formation to store energy (as indicated with the nearly straight charge/discharge line, Fig. S10†);<sup>1,2</sup> however, as circled in Fig. 6i, three samples,  $C-T_{570}-t_{0.5}$ ,  $C-T_{570}-t_2$ , and  $C-T_{700}-t_2$  with similar SSAs of 1794 m<sup>2</sup> g<sup>-1</sup>, 1991 m<sup>2</sup> g<sup>-1</sup> and 1832 m<sup>2</sup> g<sup>-1</sup>, respectively, (with a difference ratio of 9.88%) showed an 18.99% difference in specific capacitance. Tracing back the structural characteristics of the three samples, it has been noticed that they had distinct pore size distributions (Tables 1 and 2), where  $C-T_{570}-t_{0.5}$  majored with large pores of a pore volume ratio up to 67.5%,  $C-T_{700}-t_2$  majored with small pores of an SSA ratio up to 64.8%, and  $C-T_{570}-t_2$  possessed a relatively evenly-distributed pore size distribution with the highest ratio of medium pores among the three. Therefore, it is reasonable to conjecture that varied pore size distributions contributed to the difference in specific capacitance and medium pores would be more beneficial for storing energy compared to the other two.<sup>6,8,10,41</sup>

To validate the above conjecture, EIS measurements were carried out from the viewpoint of ion diffusion kinetics. Nyquist

plots of all MOF-derived carbons are given in Fig. 7a, all composed of a semi-circle (or incomplete semi-circle for  $C-T_{500}-t_{0.5}$  and  $C-T_{500}-t_2$ ) at the high-frequency range, followed by a straight diffusion line at the low-frequency range related to the mass transfer process. All the  $C-T_x-t_y$  exhibited a nearly vertical straight line except that  $C-T_{500}-t_{0.5}$  presented a jumbled curve with a slope of ~45°, suggesting distinct and significantly poor ion diffusion resistance, possibly due to the low degree of graphitization (Fig. 5a).<sup>37,40</sup> All spectra as measured except for that of  $C-T_{500}-t_{0.5}$  (Fig. S11†) were fitted with the equivalent circuit consisting of two parallel circuits (Fig. 7b). The existence of the two circuits in series instead of one could be explained by a possible interference of surface functional groups (C-O and O-C=O, which would intervene between the electrolytic ions and the electrode/electrolyte interface and create variation in the electrostatic bonding strength), and will be explored in the future study.<sup>42</sup> The fitting parameters are summarized (Table S7†). Series resistance  $R_s$  is mainly contributed by the electrolyte, carbon electrodes, their interfacial resistance, and the contact resistance between the active material and the current collector, while the charge transfer resistance  $R_{ct}$  is mainly affected by the surface functionalities.<sup>40</sup>  $C-T_{500}-t_2$  admitted relatively higher series resistance  $R_s$ , higher charge transfer resistance  $R_{ct-1}$  and  $R_{ct-2}$  as well as substantially high diffusion impedance  $W_o-R$  and the diffusion interpretation  $W_o-T$ , demonstrating that the carbonization temperature of 500 °C,



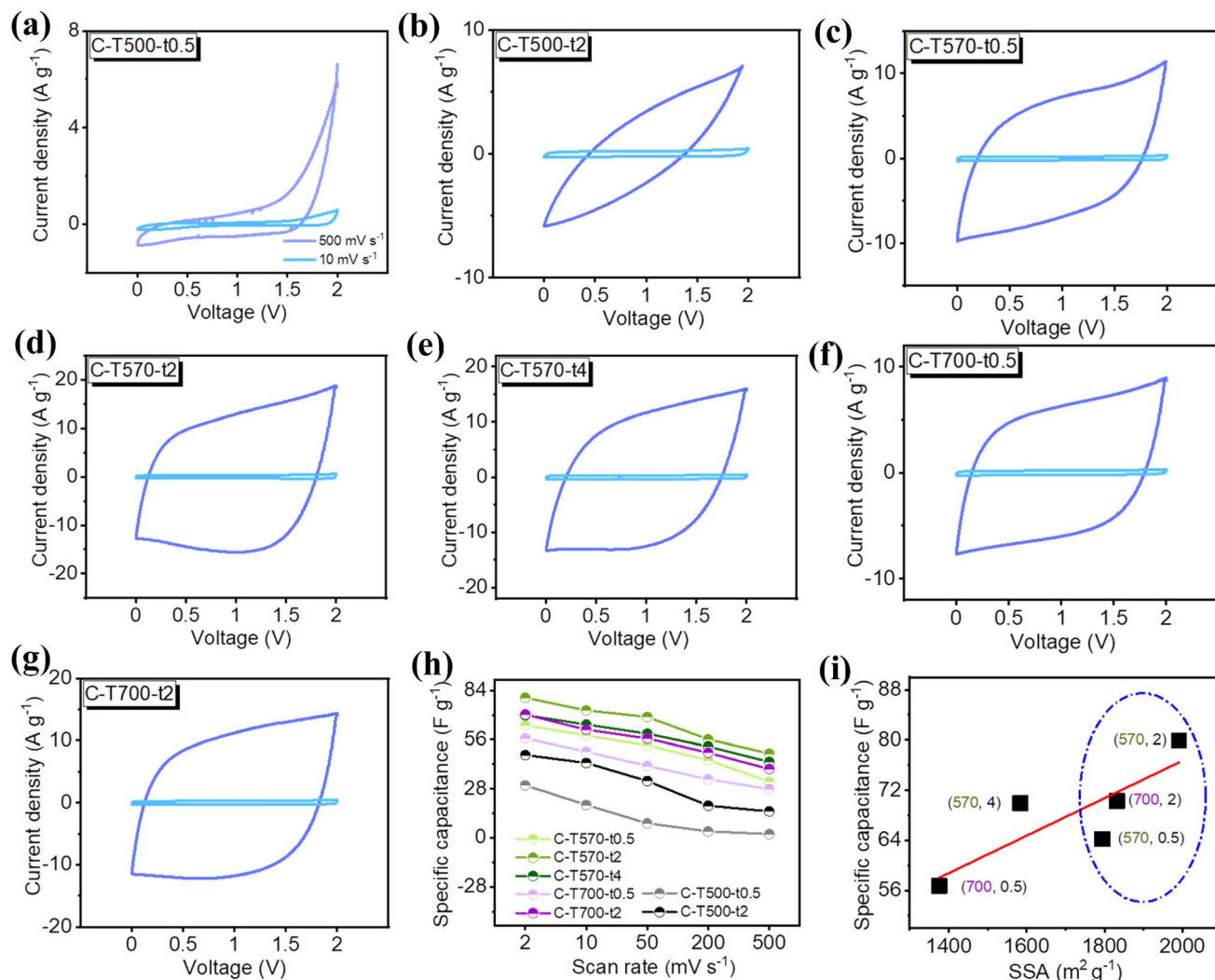


Fig. 6 (a-g) CV curves of  $C-T_x-t_y$  at the scan rates of 10 and  $500\text{ mV s}^{-1}$ . (h) The as-calculated specific capacitance values. (i) The specific capacitance of  $C-T_x-t_y$  at  $2\text{ mV s}^{-1}$  versus the specific surface area with a fitted trend line in red (with the correlation coefficient  $R^2 \sim 0.7$ ).

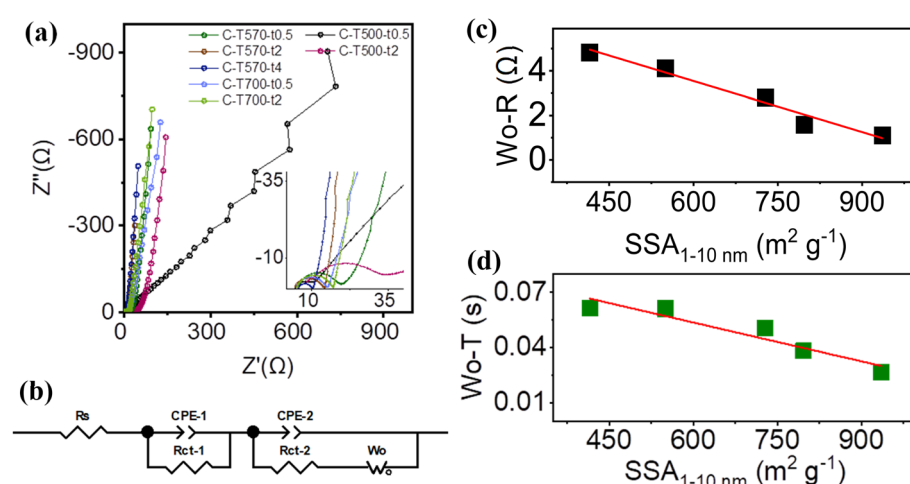


Fig. 7 (a) Nyquist plots of  $C-T_x-t_y$ , (b) the equivalent circuit for all  $C-T_x-t_y$  except  $C-T_{500}-t_{0.5}$ . (c)  $W_o-R$  and (d)  $W_o-T$  versus the partial  $SSA_{1-10\text{ nm}}$  with a fitted trend line in red (with the correlation coefficient  $R^2 \sim 0.97$  and 0.90, respectively).

although could successfully pyrolyze Zn-MOF-5, restricted it from the poor crystallinity and was not suitable for application as the electrode material. For the other five 570 °C and 700 °C treated samples, their series resistance  $R_s$  and charge transfer resistance  $R_{ct}$  varied but showed no significant difference, which was consistent with their similar ratios of  $I_D/I_G$ , and of  $sp^2$ -C,  $sp^3$ -C, and functional groups (except for the  $R_{ct}$ -1 of C-T<sub>570</sub>-t<sub>0.5</sub>, which might be related to the pore structure dominated by >10 nm pores). The effects of the size of pores that serve as ion transporting channels on the  $R_{ct}$  were also reflected by the elevated  $R_{ct}$ -2 of C-T<sub>700</sub>-t<sub>0.5</sub> and C-T<sub>700</sub>-t<sub>2</sub> majored with <1 nm pores. Interestingly,  $W_o$ -R ( $W_o$ -T) admitted the smallest values for C-T<sub>570</sub>-t<sub>2</sub> and C-T<sub>570</sub>-t<sub>4</sub> possessing the highest ratios of medium pores, and the largest values for C-T<sub>700</sub>-t<sub>0.5</sub> and C-T<sub>700</sub>-t<sub>2</sub> possessing the lowest ratios of medium pores. As illustrated in Fig. 7c and d (Fig. S12†),  $W_o$ -R ( $W_o$ -T) decreased nearly linearly upon increasing the partial surface area SSA<sub>1-10</sub> nm contributed by 1–10 nm pores, in both terms of absolute value and the relative ratio. This result directly proved that with respect to the electrolyte 1 M TEABF<sub>4</sub>/PC, 1–10 nm pores provided the most suitable transporting channel, as the pores can be accessed freely and quickly by electrolyte ions and facilitated the formation of more stable electrical double layer structures;<sup>41,43,44</sup> mostly importantly, it demonstrated the linear relationship between the structural characteristic and performance parameter by controlling the pore size distribution to be approximated as the sole variable. The roles of small pores or large pores were not presented in this study and could be studied in further research.

## 4. Conclusions

We reported the exploration of a synthesis route to prepare porous carbon materials with a pore size distribution approximated as a single variable, demonstrating that a temperature zone of (570–700) °C is suitable to carbonize Zn-MOF-5 with similar chemical properties. Studying the electrochemical performance of the MOF-derived carbons reveals a linear relationship between the 1–10 nm pores and the diffusion impedance  $W_o$ -R, which is the first numerical relation built empirically between the electrode structural feature and ion diffusion characteristic. The results of this study enrich the understanding of pore size effects on the electrochemical process within the porous medium, and the method to tailor the pore size as a single variable could be extended to various electrochemical or electrocatalysis-related areas to quantitatively evaluate the effects of pore size distribution.

## Conflicts of interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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