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dve for lithium storage†



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Surficial nanoporous carbon with high pyridinic/

pyrrolic N-Doping from sp<sup>3</sup>/sp<sup>2</sup>-N-rich azaacene

Rationally designed pyridinic/pyrrolic N-doping of anodic carbonaceous materials is significant but rare for lithium storage materials. Herein, two highly pyridinic/pyrrolic N-doped carbon materials *i.e.* the pristine material (NC, N content 10.9 wt%) and template (NPC, N content 12.6 wt%) are predictively achieved by the direct pyrolyzation of the azaacene dye with a high sp $^3$ /sp $^2$ -N content (13.6 wt%). Their total amount of pyridinic and pyrrolic content is as high as 7 at% (NC)/6.98 at% (NPC), which is close to the theoretical value (9.1 at%) with less/no graphitic N. Both of them present higher capacities of 1094.79 mA h g $^{-1}$  and 411.3 mA h g $^{-1}$ , respectively, by the 3<sup>rd</sup> cycle at 100 mA g $^{-1}$ . Featured by surficial bumps and hollows of NPC particles, the NPC electrode not only exhibits a first reversible specific discharge capacity as high as 1178.9 mA h g $^{-1}$  at a 100 mA g $^{-1}$  current density but is also stabilized at 614 mA h g $^{-1}$  after 200 cycles at a 200 mA g $^{-1}$  current density. Our results indicate that the sp $^3$ /sp $^2$ -N-rich azaacene dye can be a useful highly pyridinic/pyrrolic N-doped carbon source for high performance anodic materials.

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## Introduction

Carbon-based anode materials have been explored as one of the most key active power components for portable electronic devices due to their high storage capacity, cycling stability, and reversibility in the fields of lithium–ion batteries, <sup>1-3</sup> supercapacitors, <sup>4-6</sup> oxygen reduction reactions, <sup>7</sup> and H<sub>2</sub> storage. <sup>8</sup> Furthermore, the intrinsic properties, including low theoretical and actual capacities, and limited application ranges <sup>4,9</sup> of conventional anodic graphite can no longer meet the requirements for high-efficient and convenient power consumption. Moreover, two main strategies *i.e.* enlargement of the electrolyte–electrode interfacial area and increasing the redox sites with pre-embedded heteroatoms are applied to optimize the electrochemical properties of carbon-based anode materials.

Template precursors and alkaline activators<sup>10</sup> that tend to form various well-defined micro/nanostructures<sup>11-13</sup> have been applied to enlarge the electrolyte–electrode interfacial area.<sup>14-16</sup> Furthermore, sufficient and effective lithiation sites<sup>17</sup> between porous electrodes and liquid electrolytes could deeply enhance the chemical stability and electrochemical performance of carbon-based anodes,<sup>18-21</sup> and overcome the expansion and other defects of Si and Sn materials.<sup>2,22</sup>

Various approaches including pre-templating,23-26 graphitization,27 pressure-assisted carbonization,28 activating agents,96,29,30 and alternative carbon sources have been widely applied to improve the specific electrolyte-electrode interface<sup>2b,23,31</sup> and especially increase the hetero-doping ratio<sup>7d,16,32</sup> in carbon materials to achieve extraordinary electrochemical properties.<sup>24,33,34</sup> Since pyridinic/pyrrolic N-doped carbons possess a higher Li atom average adsorption energy than the Li-Li binding energy, 17b it is generally acknowledged that a much more higher ratio pyridinic/pyrrolic N-doping would enhance the insertion/deinsertion ability of the loaded Li<sup>+</sup> ions in carbonaceous materials. 19,24,35,36 Special carbon sources such as biomass, 10,37-39 polymers including conjugated polymers,40,41 polyaniline,42 and polyamide, 6,43,44 melamine-resorcinol-formaldehyde, 45 metal organic frameworks, 18,46 and complexes 47,48 containing sp2-N and sp<sup>3</sup>-N could produce versatile N-doped porous carbons.

However, there are few reports<sup>17,20</sup> on the rational design of sp<sup>3</sup>/sp<sup>2</sup>-N-rich aromatics as a source of high ratio pyridinic/pyrrolic N-doped carbon-based anode materials with N-doping ratios over 10 wt%,<sup>3,20</sup> large specific surface areas, and high Li<sup>+</sup> ion storage capacity.<sup>28,49,50</sup> Therefore, it is highly desirable to

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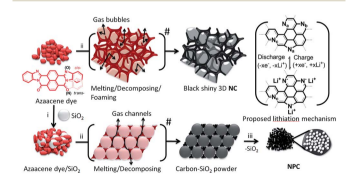
explore this type of new carbon source due to its advantages of low-cost, simple synthesis, and high output as high-performing carbon-based anodic lithium storage materials.

Herein, we successfully synthesized two novel amorphous carbon materials, i.e. N-doped porous carbon (NPC) and pristine N-doped carbon (NC), by pyrolyzing the low-cost, easilysynthesized N-rich azaacene dye (mixture of trans isomeric Pigment Orange 43 and cis isomeric Pigment Red 194)51 without/with a micro-SiO2 template under an inert gas atmosphere. NPC possesses the special characteristic of surficial nanopores, whereas NC has no surficial pores. The novel Ndoped carbons are systematically characterized via elemental analysis, X-ray photoelectron spectroscopy (XPS), X-ray powder diffraction (XRD), Raman spectroscopy, Brunauer-Emmett-Teller surface area measurements (BET), field emission scanning electron microscopy (FE-SEM), transmission electron microscopy (TEM), high-resolution transmission electron microscopy (HR-TEM), and electrochemical tests.

### Results and discussion

### **Synthesis**

The  $\pi$ -conjugated azaacene dye carbon precursor was synthesized via a facile cyclocondensation route in high yield.52 As shown in Scheme 1, NC was prepared by the direct carbonization of the azaacene dye at high temperatures for 2 h under an Ar atmosphere. Considering that a low temperature will result in incomplete carbonization, whereas a high temperature will lead to a low heteroatom doping content, 17d we have finally chosen 700 °C as the ideal temperature. As the temperature increased, the dye powder melted at 497-504 °C (ref. 52e) and decomposed accompanied by swelling. The high melting point led to a high carbon yield. Subsequently, the melted azaacene dye combusted and carbonized into intact, bulky, crisp, and black shiny 3D-like self-supported carbon due to the self-sealed pressure from the released waste gas. For NPC, the  $\pi$ -conjugated azaacene dye was granulated with micro-SiO<sub>2</sub> for 15 min and then carbonized under the former conditions. Then, the



Scheme 1 Synthetic route for the preparation of NC and NPC.  $N_a$ ,  $N_b$ , and N<sub>c</sub> represent pyridinic N, pyrrolic N, and graphitic N (less/none), respectively. Azaacene dye: 1,4,5,8-naphthalenetetracarboxylic dianhydride, o-benzenediamine, DMF, 150 °C, 4 h, 65% yield, red powder. (i) Micro-SiO<sub>2</sub>, granulation for 15 min; (ii) 700 °C, Ar; and (iii) NH<sub>4</sub>HF<sub>2</sub> (ag.), NaOH (ag.), deionized water. Bracket: proposed lithiation mechanism.

organic powder melted on the surface of the neighbouring micro-SiO<sub>2</sub> particles and then decomposed with waste gas released through the capillary space between micro-SiO2 particles. This led to the formation of an NPC-SiO2 mixture as a dark black powder, which is different from the shiny black 3D-like appearance (Scheme S1†) of NC. Then, the as-prepared NPC-SiO<sub>2</sub> mixture was immersed in excess aqueous NH<sub>4</sub>HF<sub>2</sub> and sodium hydroxide in sequence to get rid of SiO2. Finally, NPC was obtained as a black powder with a large surface area.

### Morphology

The shiny black 3D-like appearance of the self-supported carbon NC (Fig. S1a†) matched well with the smooth surface morphology shown in the FE-SEM images of the ground NC particles. The size of these particles ranged from several to tens of micrometers without well-defined structures, as shown in the magnified images (Fig. 1a inset). Moreover, NPC is presented as unshaped particles (Fig. 1b). From another magnified image (Fig. 1b inset), it was found that the surface of the NPC particles had large quantities of bumps and hollows in the range of tens of nanometers acting as effective Li<sup>+</sup> intercalation sites at the liquid-solid interface. The XRD patterns of both NC and NPC are inserted as white curves in Fig. 1a and b, which exhibit broad and low diffraction bands at the  $2\theta$  values of around 26-28° and 41–46°, representing the interplanar spacings of  $(d_{002})$ and  $(d_{100})$ , respectively. These bands suggest a low

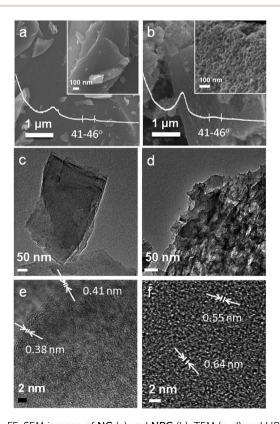


Fig. 1 FE-SEM images of NC (a) and NPC (b), TEM (c, d) and HR-TEM (e, f) images of NC (c, e) and NPC (d, f). Inset images: magnified surficial morphologies of NC and NPC. Inset white curves: XRD patterns of NC and NPC. White arrows and bars: distances of the crystal lattices.

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graphitization degree for both NC and NPC. According to the Bragg's Law  $2d\sin\theta=n\lambda$  ( $\lambda=1.5405$  Å), the values of the weak  $d_{002}$  and  $d_{100}$  bands are 0.34–0.32 nm (26–28°) and 0.22–0.20 nm (41–46°), respectively, indicating the low crystallinity<sup>53,54</sup> of these two new carbon materials.

TEM and HR-TEM analyses were further conducted to investigate the microstructures of the as-prepared carbon materials. In Fig. 1c and d, NPC displays a similar disordering of its inner crystalline lattice as NC; this indicates that their structural defects are favourable for Li<sup>+</sup> ion diffusion and effective contact between active materials and electrolyte.55 The HR-TEM images, as shown in Fig. 1e and f, also reveal that both NC and NPC are amorphous carbon materials without a long range order and straight crystalline lattices. However, a few large and distinguishable short lattice fragments were observed, which were marked by white arrows and bars. The measured width of the lattice fragments of NC was as large as 0.38/ 0.41 nm, and that for NPC was as large as 0.55/0.64 nm, which was much rougher than NC. It was proposed that the surface pores originated from the interfacial etching reactions among the surficial corrosive matters and groups (-OH, -COOH, etc) on the template. Moreover, for NPC, the micro-SiO2 template interdicted macroscale 3D-like cross-linkage and thermal agglomeration of the melting and combusting azaacene dye during the pyrolysis process.

Therefore,  $N_2$  adsorption-desorption isotherms were obtained, as shown in Fig. 2. From the analysis of the BET data shown in Fig. 2a, the specific surface area of **NPC** was found to be 456.8 m<sup>2</sup> g<sup>-1</sup>, which was almost 264 times that of **NC** 

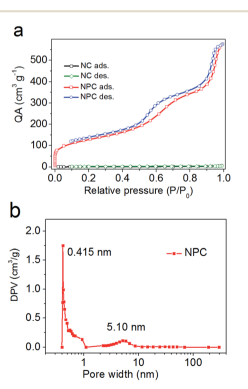


Fig. 2  $N_2$  adsorption–desorption isotherms (a) and pore size distributions (b) of NPC. QA: quantity adsorbed and DPV: differential pore volume.

(1.73 m<sup>2</sup> g<sup>-1</sup>). Moreover, **NPC** possesses both micropores (peaked at 0.415 nm) and mesopores (peaked at 5.10 nm) (Fig. 2b). The pores of **NPC** supply a number of commodious reservoirs for potential Li<sup>+</sup> ion transport and accumulation under electrochemical test conditions.

### Raman spectroscopy

The Raman spectra in Fig. 3a exhibit two distinct peaks attributed to the D band (ca. 1353/1367 cm<sup>-1</sup>) and G band (ca. 1600/1571 cm<sup>-1</sup>) for NC and NPC, respectively. The intensive G bands indicate that the degree of graphitization decreases from NC to NPC after the addition of the micro-SiO<sub>2</sub> template. Compared with that of NC, the intensity ratio of the D-band versus G-band ( $I_D/I_G$ ) in NPC was calculated to be 1.02, which was larger than 0.86 calculated for NC and suggested much more structural defects and disorder<sup>56-58</sup> for NPC. This was further confirmed by its widely distributed surficial pores (Fig. 1b inset).

### X-ray photoelectron spectroscopy (XPS)

XPS was used to analyse the elemental distribution and the related bonding species in NC and NPC. As shown in Fig. 3b, both NC and NPC mainly contain C, N, and O dopants with three characteristic peaks at  $\sim\!285$  eV,  $\sim\!400$  eV, and  $\sim\!532\pm1$  eV, corresponding to C 1s, N 1s, and O 1s. For NC, the total content of C, N, and O elements was 85.07 at%, 7.48 at%, and 7.45 at%, respectively. For NPC, the content of C, N, and O elements was 85.6 at%, 7 at%, and 5.57 at%, respectively. Moreover, a certain amount of residual fluorine (1.64 at%) and Si (0.17 at%) in NPC was detected, which most likely originated from the F $^-$  of NH<sub>4</sub>HF<sub>2</sub> and SiO<sub>2</sub> particles, respectively, buried and trapped in the amorphous carbon during the de-SiO<sub>2</sub> process.

In Fig. 3c, the C 1s spectra for NC and NPC can be deconvoluted into several individual peaks. The fitted peaks at 284.6 eV, 286.1 eV, and 288.5 eV are related to the bonding configurations of C=C, C=N/C-O, and C=O, respectively.59 It should be noted that the ratios of sp<sup>2</sup>-Cs in NC and NPC occupied a large percentage (61.58 at% and 58.99 at%, respectively). The other types of C elements occupied 23.49 at% for NC and 26.61 at% for NPC. In Fig. 3c, the three deconvoluted peaks for N element in NC at 398.3 eV, 400.8 eV, and 402.8 eV belong to pyrrolic-N, pyridinic-N, and graphitic N/oxidized N species, occupying 3.67 at%, 3.31 at%, and 0.5 at%, respectively.<sup>29,60,61</sup> However, in Fig. 3d, only pyrrolic-N and pyridinic-N peaks in NPC were detected, which were 3.41 at% and 3.59 at%, and they originated from the pre-embedded sp<sup>2</sup>-N and sp<sup>3</sup>-N, respectively. As shown in Fig. S3b,† the spectra of O element in NC were also deconvoluted into two peaks at 531.5 eV and 532.8 eV, occupying 5.11 at% and 2.34 at% from O=C and O-C, respectively. For NPC, the corresponding two fitted peaks at 531.5 eV and 532.8 eV of O element occupied 3.55 at% and 2.02 at%, respectively. The FE-SEM mapping images exhibit the homogenous distribution of C, N, and O for both the NC (Fig. S2d-S2f†) and NPC (Fig. S2h-S2j†) particle samples. Elemental analysis disclosed that the N content weight ratios in both NC and NPC were as high as 10.9 wt% and 12.6 wt%,

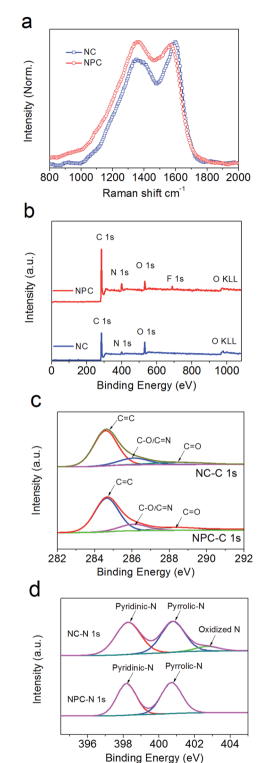


Fig. 3 Raman spectra (a) of NC and NPC synthesized from  $\pi$ -conjugated azaacene dye at 700 °C. C 1s (c) and N 1s (d) XPS spectra (b) with the deconvoluted curves of NC and NPC.

respectively, which were higher than those in other N-contained carbonaceous materials for LIBs.  $^{19,62,63}$ 

The investigation on NC and NPC suggested that employment of effective templates and heteroatom pre-embedded

precursors is an efficient strategy to produce highly N-doped porous carbon materials, 33,64,65 which are expected to display effective Li<sup>+</sup> ion insertion/de-insertion behaviour according to the proposed lithiation mechanism (Scheme 1).<sup>17</sup>

### **Electrochemical performance**

Fig. 4a and S4b† present the first three voltage curves of the NC and NPC electrodes measured through cyclic voltammogram (CV) under ambient conditions between 0.005 and 3.0 V at a scan rate of 0.5 mV  $\rm s^{-1}$ . The first discharge semi-cycle did not overlap the subsequent cycles. The strong but broad peaks at 0.4–1 V were mainly contributed by the newly generated solid electrolyte interphase (SEI) film,  $^{66-71}$  especially for the enhanced peak at 0.4 V for NPC with an improved surface area.

In Fig. 4b, the NC and NPC electrodes present rate performance curves at different current densities ranging from 100, 200, 400, 800, and 1600 to 3200 mA  $g^{-1}$ . At a current density of 100 mA g<sup>-1</sup>, the initial irreversible discharge capacity of the NPC electrode reached as high as 2283 mA h g<sup>-1</sup>, which was thrice the corresponding value of 767 mA h g<sup>-1</sup> for the NC electrode. It should be noted that these values are almost double of their initial reversible charge capacities of 1107 mA h  $g^{-1}$  and 448 mA h  $g^{-1}$ . The irreversibility is most likely due to the SEI effect and unidirectional Li<sup>+</sup> ion trapping or consumable reaction. However, in the first reversible cycle, the specific discharge capacity of the NPC electrode reached as high as 1178.9 mA h  $g^{-1}$  and stabilized at 1014 mA h  $g^{-1}$  by the end of the 11th cycle, which were both nearly 2.7 times the corresponding value of 432 mA h  $\rm g^{-1}$ , and stabilized at 387 mA h  $\rm g^{-1}$ for the NC electrode. At a higher specific rate of 1.6 A  $g^{-1}$ , the capacity was still stabilized at  $\sim$ 336.0 mA h g<sup>-1</sup>, which was close to the theoretical value of graphite (372 mA h  $g^{-1}$ ). Even at the strongest current density of 3.2 A g<sup>-1</sup>, the NC and NPCbased electrodes still delivered 155 and 207 mA h  $\rm g^{-1}$  at the 57  $^{\rm th}$ cycle. Significantly, when the current density returned to 100 mA g<sup>-1</sup> in the 70<sup>th</sup> cycle, the discharge capacity was still over 976 mA h g<sup>-1</sup> and 389 mA h g<sup>-1</sup> for **NPC** and **NC**, respectively, which recovered to the initial value although slightly lower than that in the first cycle at a current density of 100 mA  $g^{-1}$ .

Fig. 4c shows the cyclability and coulombic efficiency (CE) at a current density of 200 mA  $g^{-1}$  for the NC and NPC electrodes. The initial irreversible discharge capacity of the NPC electrode reached as high as 1648 mA h g<sup>-1</sup>. The NPC electrode exhibited good and stable cyclability with a specific capacity ranging from 700 to 614 mA h  $g^{-1}$  up to the 200<sup>th</sup> cycle. In contrast, the NC electrode presented a much lower specific capacity ranging from 325 to 364 mA h g<sup>-1</sup> up to the 200<sup>th</sup> cycle. However, both their coulombic efficiencies from the second cycle up to the 200<sup>th</sup> cycle were almost over 93%. In this situation, the corresponding galvanostatic charge/discharge profiles (Fig. 4d) of the better performing NPC electrode were obtained at six different current densities ranging from 100, 200, 400, 800, 1600, to  $3200 \text{ mA g}^{-1}$  at the  $10^{\text{th}}$ ,  $20^{\text{th}}$ ,  $30^{\text{th}}$ ,  $40^{\text{th}}$ ,  $50^{\text{th}}$ , and  $60^{\text{th}}$  cycles, respectively, which indicated that the NPC electrode exhibited matching electrochemical properties, almost coinciding with the charge/discharge voltage-capacity curves.

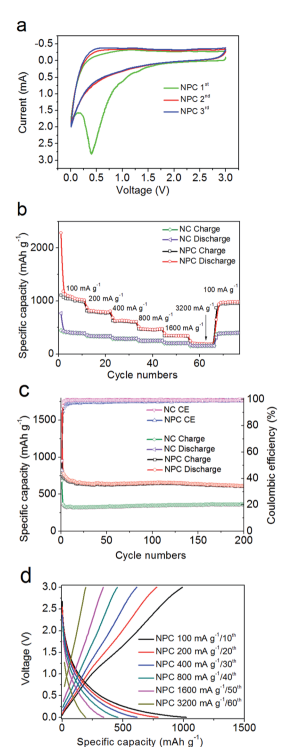


Fig. 4 (a) Electrochemical performance of the NPC electrode: the first, second, and third CV profiles at a scan rate of 0.5 mV s<sup>-1</sup> over the potential window of 0.005-3 V (vs. Li/Li<sup>+</sup>). (b) Rate performance at different current densities from 100 to 3200 mA  $\dot{g}^{-1}$ . (c) Cycling performance and CE at 200 mA g<sup>-1</sup> d<sup>-1</sup>) Galvanostatic chargedischarge curves of NPC at different current densities ranging from 100 to 3200 mA  $g^{-1}$  in the  $10^{th}$  –  $60^{th}$  cycles.

To further understand the charge-transfer ability and illustrate the advantages of the NPC electrode over the NC electrode, EIS tests on NPC and NC electrodes were conducted under the

same conditions. The charge transfer resistances are depicted in Fig. S5.† It can be seen that both electrodes present highfrequency semicircles, representing the charge-transfer process. The EIS plots in the low-frequency region suggest that the NPC electrode possesses a lower Li<sup>+</sup> ion diffusion resistance than the NC electrode. From the CV measurement and calculation, the charge-transfer resistance of the NPC electrode was calculated to be about 42.5  $\Omega$ , smaller than that of NC (55.7  $\Omega$ ), indicating its better conductivity.

Compared to the NC electrode, the improved high capacity and rate capability of the NPC electrode could be explained by its special porous structure and high-level N-doping. As abovementioned, the carbonaceous NPC electrode plays a key role as conductive channels1,9b,18 for electron transport, whereas its large number of pores possibly act as Li<sup>+</sup> ion storage cisterns.<sup>71</sup> Furthermore, the enlarged electrode/electrolyte interface of the NPC electrode promotes the rapid absorption and release of Li ions with a fast charge-transfer process.

## Conclusions

In summary, two novel pyridinic/pyrrolic N-doped carbon materials have been rationally designed and successfully obtained via the pristine and template pyrolyzation of an sp<sup>2</sup>/sp<sup>3</sup>-N-rich azaacene dye with a complete  $\pi$ -conjugated framework and high melting point. The anodic NPC electrode exhibited high Li<sup>+</sup> ions storage performance due to its high pyridinic/ pyrrolic N-content and electrolyte and liquid interface. Its specific capacity was as high as 614 mA h g<sup>-1</sup> after 200 cycles even at 200 mA  $g^{-1}$ . Our results indicate that the pyrolyzation of azaacene dye rich in sp<sup>3</sup>/sp<sup>2</sup>-N is an effective strategy to achieve more highly monospecies or concomitant pyridinic/pyrrolic Ndoped carbon materials.

## **Experimental**

### **Materials**

Naphthalene-1,4,5,8-tetracarboxylic acid dianhydride purchased from Zhengzhou Keyulong Chemical Products Co., Ltd. o-Benzene-diamine was purchased from Energy Co., Ltd. Micro-SiO<sub>2</sub> (8  $\pm$  2  $\mu$ m  $\geq$ 80%) was obtained from highperformance thin layer chromatography plates (200  $\times$  200 mm) purchased from Yantai Jiangyou Guijiao Kaifa Co., Ltd. All other reagents and solvents were purchased from Wanging Co., Ltd. All solvents were used without further purification.

### Synthesis of NC and NPC

Azaacene was synthesized, filtered, washed, and dried as a red powder in 64% yield by refluxing DMF in o-benzene-diamine and naphthalene-1,4,5,8-tetracarboxylic acid dianhydride. The mixture of azaacene dye (1.5 g) with micro-SiO2 (3 g) was mixed and ground for 10 min. Azaacene (1 g) and the ground mixture were put into corresponding Al<sub>2</sub>O<sub>3</sub> crucibles, in an OTF-1200X tube furnace with SiO2 glass and protected under an Ar atmosphere and heated to 700 °C at a heating rate of 3 °C min<sup>-1</sup> and kept for 0.5 h under a nitrogen atmosphere. The activated mixture was then washed with the  $NH_4HF_2$  solution (1 mol  $L^{-1}$ ), NaOH solution (1 mol  $L^{-1}$ ), and deionized water in sequence until the filtrate became neutral. The sample was finally dried overnight at 100 °C in an oven. The carbonization yields of **NC** and **NPC** were 65% and 40%, respectively.

#### Methods

The morphology of the samples was observed via FE-SEM (Hitachi S-4800, Tokyo, Japan). TEM observation was carried out using a JEOL 2100F microscope operating at an accelerating voltage of 100 kV. HR-TEM observation was carried out using an FEI TECAI G2F20 microscope operating at an accelerating voltage of 200 kV. X-ray diffraction (XRD) patterns were obtained using a Bruker D8 X-ray diffractometer with Cu Ka radiation ( $\lambda = 1.5405 \text{ Å}$ ). Raman spectroscopy was carried out suing a WITec alpha300M+ micro-Raman confocal microscope. CHN elemental analysis was performed using a Thermo Scientific FLASH 2000 elemental analyzer. XPS measurements were performed using a Thermo Scientific ESCALAB 250XI system with a monochromatic Al Kα X-ray source. Nitrogen adsorption and desorption isotherms were determined by nitrogen physisorption at 77 K using a V-Sorb 2800P BET surface area and pore volume analyser.

#### **Electrochemical tests**

The electrochemical properties of NC and NPC were measured with 2032 coin cells using pure lithium foil as the counter electrode. The working electrodes were constructed by mixing the as-prepared azaacene dye samples of 80 wt% NC or NPC, 10 wt% polyvinylidene fluoride (PVDF), 10 wt% acetylene black, and N-methyl-2-pyrrolidone as the dispersing solvent to obtain slurries, which were coated homogeneously onto the copper foil and dried at 100 °C overnight in a vacuum oven. Circular electrodes (12 mm in diameter) were prepared using a punching machine and weighed with an electronic balance within 0.01 mg. The electrolyte was a 1.0 mol L<sup>-1</sup> LiPF<sub>6</sub> in ethylene carbonate and dimethyl carbonate solution (1:1 (w/w)). Celgard 2300 was applied as the separator. Coin cells with both electrodes were assembled in an inert Ar gas-filled glove box. Both the moisture and oxygen contents were kept below 0.1 ppm. The galvanostatic charge/discharge parameters of the cells were measured using a multichannel NEWARE batterytesting system with the voltage ranging from 0.005 to 3.0 V  $\nu s$ . Li<sup>+</sup>/Li at different rates. The calculated capacity values were determined based on the weight of the loaded carbon materials on the collectors. Galvanostatic cycling was performed using a NEWARE multichannel battery testing system. Alternative current (AC) impedance measurements were conducted using a CHI 760D electrochemical workstation (CH Instruments, Inc.) at room temperature. EIS was performed with an AC voltage of 5 mV amplitude in the frequency range from 100 kHz to 0.1 Hz.

# Conflicts of interest

There are no conflicts to declare.

## Acknowledgements

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