# RSC Advances



# PAPER

Cite this: RSC Adv., 2017, 7, 54626

Received 4th October 2017 Accepted 16th November 2017

DOI: 10.1039/c7ra10954g

rsc.li/rsc-advances

# Introduction

Electrochemical capacitors (ECs), also known as ultracapacitors or supercapacitors<sup>1</sup> belong to the class of electrochemical energy storage (EES) devices that find applications in electric vehicles<sup>2</sup> and several small to medium power electronic systems.<sup>3-5</sup> Based on the charge storage mechanisms, ECs can be categorized into two types: (1) the electrochemical double layer capacitors (EDLC), which are capable of storing energy electrostatically by the simple non-faradaic adsorption of ions from the electrolyte onto the large surface area electrodes; (2) the pseudocapacitors that benefit from the phenomenon of electrostatic charge storage, like EDLC, accompanied by a sequence of fast, reversible redox processes, electrosorption or intercalation.<sup>2</sup> In general, the performance of ECs largely

# Template-free synthesis of nitrogen doped carbon materials from an organic ionic dye (murexide) for supercapacitor application†

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The present study describes a template-free single step carbonization route to prepare hierarchically structured nitrogen-doped carbon materials (NCMs) by using an organic ionic dye (OID), ammonium purpurate (murexide). These NCMs exhibited moderate specific surface area (307  $m^2$  g<sup>-1</sup> for NCM<sub>(MDE)</sub>-900) and hierarchical macro/meso/microporous structures with abundant nitrogenous functionalities that contributed towards the high specific capacitance displayed by pseudocapacitive contribution. In particular, the NCM<sub>(MDE)</sub>-800 displayed superior specific capacitance (222 F g<sup>-1</sup> at 3 A g<sup>-1</sup>) and excellent cyclic stability of  $\sim$ 10 000 cycles at 10 A g<sup>-1</sup> in 1 M 1.0 M H<sub>2</sub>SO<sub>4(aq)</sub>. The material prepared at higher temperature, viz. NCM<sub>(MDE)</sub>-900, exhibited high rate capability at 50 A g<sup>-1</sup>, which is 92% of their specific capacitances at 10 A  $g^{-1}$ . Further systematic investigations of NCM<sub>(MDE)</sub>-800 in three different electrolytes, viz. 1.0 M H<sub>2</sub>SO<sub>4(aq)</sub>, 6 M KOH<sub>(aq)</sub> and 0.5 M Na<sub>2</sub>SO<sub>4(aq)</sub>, revealed that 1.0 M H<sub>2</sub>SO<sub>4(aq)</sub> is a promising electrolyte for achieving good specific capacitance, high capacitive retention and long term cyclic stability, which might have its origins in the formation of ionic interactions between the active protons in 1.0 M H<sub>2</sub>SO<sub>4(aq)</sub> and the alpha carbon atoms with Lewis basic character next to the nitrogen in these NCMs. **PAPER**<br> **Template-free synthesis of nitrogen doped carbs**<br> **Template-free synthesis of nitrogen doped carbs**<br> **CREATE THE CONSTRANT CONSTRANT CONSTRANT CONSTRANT CONSTRANT CONSTRANT CONSTRANT CONSTRANT CONSTRANT CONSTRAN** 

depends on the physical and chemical nature of the active electrode material.<sup>6</sup> For EDLC applications, different varieties of carbon materials such as carbon black, graphene,<sup>7</sup> carbon nanotubes,<sup>8</sup> and carbon onions<sup>9</sup> have been extensively exploited due to their high electrical conductivity, high surface area and excellent cyclic stability.<sup>2</sup>

One possible way to achieve high energy density and power density along with high specific capacitance and long cyclic life in EDLCs is to introduce a foreign heteroatom functionality such as nitrogen,<sup>10</sup> oxygen,<sup>11</sup> sulphur, boron and phosphorous<sup>12</sup> into the carbon framework.<sup>11,13</sup> Typically, nitrogen is found to be the most suitable heteroatom that upon doping in the carbon framework leads to superior capacitance, electronic conductivity,<sup>14</sup> surface wettability<sup>15</sup> and cycle life, due to its pseudo-capacitive contribution.<sup>16,17</sup> Though the actual role of doped nitrogen content and its types (pyridinic-N, pyrrolic-N or graphitic-N) for the capacitive performance of nitrogen doped carbon materials (NCMs) is still unclear, a recent study by J. Nakamura and co-workers<sup>18</sup> describes how the conjugation between the nitrogen lone pair and the  $\pi$ -bonding network of the carbon lattice dramatically change the physiochemical properties of the NCMs. Both the amount of doped nitrogen content and the type of nitrogen functionality can make remarkable improvements in achieving superior capacitive performance and better cyclic stability.<sup>19</sup> Moreover, the

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<sup>†</sup> Electronic supplementary information (ESI) available. See DOI: 10.1039/c7ra10954g

Lewis basicity of carbon atoms adjacent to the nitrogen and the lone pair of electrons on the doped nitrogen atoms offer active sites to develop ionic interactions in the electrolyte, which therefore induce reversible fast redox reactions during potential cycling and make pseudocapacitive contributions to the overall charge storage by these materials.<sup>20,21</sup> For high energy and high power density applications, the supercapacitor electrode material must contain hierarchical porous structures with abundant micropores (accumulate charge effectively), interconnected mesopores (reduce the ion diffusion distance) and appropriate macropores (serve as ion buffering reservoirs for electrolyte ions) to facilitate fast traf ficking of ions and their interactions with the active sites of the electrode material.<sup>22,23</sup>

To obtain the desired properties in the nitrogen doped carbon materials, most of the strategies involve using hard or soft templates to achieve high surface area and highly porous structures. The use of any of these template methods has its disadvantages, which therefore make the overall process of synthesis quite complicated, with the final product containing small fractions of impurities coming from the template itself.19,24,25 In some cases, the removal of template requires harsh experimental conditions (use of HF to remove  $SiO<sub>2</sub>$ template) and trace amounts of template are found to have significant impact on the properties of such materials prepared by template assisted routes.<sup>26</sup> More recently, Watanabe and co-workers reported a new strategy of using protic ionic salt (PIS) based precursors containing aromatic heteroatoms or  $\pi$ -conjugated systems that impart special thermal stability to the PIS. This high thermal stability plays a major role in producing highly graphitized nitrogen rich carbon networks during high temperature carbonization.<sup>27</sup> Like PISs, organic ionic dyes (OIDs), commonly used in the pigment industry, are also highly conjugated carbon networks with a variety of structurally bound nitrogen atoms. These OIDs are cheap and commercially available in large quantities, due to their simple synthesis route making them potential candidates to synthesize NCMs for electrochemical device applications.<sup>28</sup> Moreover, the use of OIDs as a precursor for NCMs is in its infancy and further investigation of these systems is required to produce NCMs with desired properties for supercapacitor application. Paper<br>
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The present work describes a simple template-free route for the synthesis of nitrogen doped carbon materials (NCMs) by using the OID (ammonium purpurate aka murexide with formula  $\text{[NH}_{4}^+ \text{][C}_{8} \text{H}_{4} \text{N}_5 \text{O}_6 {}^- \text{]).}$  The unique chemical nature of this dye (ammonium cation and purpurate anion) and the molecular structure of the organic anion (abundant carbonyl groups) make this precursor a suitable candidate for high temperature carbonization, since the in situ release of gaseous products  $(CO<sub>2</sub>$  and NH<sub>3</sub>) may act as an internal porogen to form hierarchical porous structures in the final NCMs. The detailed investigation of the newly synthesized NCMs, particularly  $NCM<sub>(MDE)</sub>$ -800, reveals the superior specific capacitance (222 F  $g^{-1}$  at 3 A  $g^{-1}$ ), rate capability and cyclic stability (10 000 cycles at 10 A  $g^{-1}$ ) of this material, outperforming several known NCMs reported to date.<sup>4,29-32</sup>

# Materials and methods

### Synthesis of  $NCM$ <sub>(MDE)</sub>-X

Nitrogen-doped carbon materials (NCMs) were synthesized from murexide (ammonium purpurate,  $\text{[NH}_4^+ \text{][C}_8\text{H}_4\text{N}_5\text{O}_6^- \text{])}$  by a simple one step carbonization method in a vacuum furnace (MTI model: VBF-1200X). In brief, 10 g of murexide were placed into the furnace in a ceramic crucible covered with a lid. The material was annealed under inert Ar environment (flow rate of 150  $\text{cm}^3 \text{ min}^{-1}$ ) at 700–900 °C (heating rate of 10 °C min<sup>-1</sup>) for 2 h. Afterwards, the fine carbon powder formed was first washed with dilute hydrochloric acid (0.1 M) to remove any possible impurities formed during the carbonization process, followed by washing with water to neutralize the pH. The pyrolysis products formed at  $X^{\circ}$ C were denoted as NCM<sub>(MDE)</sub>-X, where X represents the pyrolysis temperature (being  $X = 700$ , 800 or 900). Finally, the synthesized NCMs were dried overnight in a vacuum oven at 80 $\degree$ C.

#### Physiochemical characterization

Thermogravimetric analysis (TGA) was performed on the pure murexide to examine the thermal degradation of the precursor using a thermogravimetric analyser (Linseis STA PT-1000) at a 10 °C min<sup>-1</sup> temperature ramp under Ar flow. For morphological characterization, a Quanta FEG 450 field emission scanning electron microscope (FE-SEM) and JEOL 3010 transmission electron microscope (TEM) operating at 300 kV, and JEOL 2100F (200 kV) were used. The X-ray powder diffraction (XRD) patterns were recorded using a STOE Theta/theta X-ray Diffractometer System with monochromatic Cu Ka radiation of wavelength,  $\lambda = 1.54060$  nm operating at 20 kV and 5 mA within 20–60 (2 $\theta$ ). Raman spectra were recorded for NCMs with a Renishaw in Via Reflex system with a He-Ne laser (632.8 nm) to measure the extent of graphitization, and ATR-FTIR spectra were recorded (Bruker-Tensor II) to examine the functional groups attached to the synthesized NCMs. An X-ray photoelectron spectrometer (PSP Electron Energy Analyzer) with Mg Ka radiation ( $hv = 1253.6$  eV, 12 kV, 144 W) under a base pressure of  $\leq 2 \times 10^{-10}$  mbar with H<sub>2</sub> as the main residual gas was employed for surface elemental compositional analysis of carbon and other heteroatoms present in the newly synthesized NCMs. The specific surface area of the catalysts was determined from  $N_2$  adsorption-desorption data at 77 K by using a Micromeritics Tristar II 3020 surface area and porosity analyzer. For each analysis, 0.3 g of sample was degassed at 300  $\degree$ C for 3 h to remove any possible moisture content from the NCMs surface as well as other adsorbed gases. The pore size distribution (PSD) plot was recorded based on the Barret–Joyner–Halenda (BJH) model.

#### Electrochemical studies

The working electrodes for electrochemical characterization were prepared by mixing 80% of the as synthesized carbon material NCM<sub>(MDE)</sub>-X with 10% carbon black (Super C 65, TIMCAL) and 10% polyvinylidene fluoride (PVDF) in N-methylpyrrolidinone to form a homogeneous ink by ultrasonication for

1.5 h. This ink was coated onto a monolith substrate  $(1 \text{ cm}^2)$  and dried under an infrared lamp; the electrodes were dried at 80 °C in an oven overnight. The mass loading of active material on each current collector was 2 mg.

The electrochemical behaviour of  $NCM_{(MDE)}-X$  materials was investigated using three electrode systems on a potentiostat/ galvanostat (VMP Biologic) at room temperature (25  $^{\circ}$ C). The working electrode was fabricated as described above. Ag/ AgCl<sub>(saturated)</sub> was used as the reference electrode and Pt as the counter electrode. Aqueous 1 M  $H<sub>2</sub>SO<sub>4</sub>$ , 6 M KOH and 0.5 M  $Na<sub>2</sub>SO<sub>4</sub>$  solutions were used as electrolytes in this study. Prior to electrochemical studies, all electrolytes were thoroughly purged with highly pure nitrogen for half an hour to remove oxygen, while a nitrogen blanket was kept above the solution during measurements. Cyclic voltammetry (CV) studies were performed at scan rate of 5–200 mV  $s^{-1}$  in the -0.8 V to +0.2 V voltage window. The current densities applied in the galvanostatic charge–discharge (CD) experiments were between 1– 50 A  $g^{-1}$ . Electrochemical impedance spectroscopy (EIS) was performed in the 10 mHz to 100 kHz frequency range under the influence of 5 mV AC frequency amplitude. **ESC** Advances Article in the interaction and a modelli methods were alternative constrained to the interaction of the electrodechical constrained on the creative of MAS noncomputer interactions are common common a proper

The specific or gravimetric capacitance  $(C_s)$  was calculated from CD measurements, using the following equation:

Specific capacitance, 
$$
C_s = \frac{I \times t}{m \times V}
$$
 (1)

in the above expression,  $I$  represents the current,  $t$  is discharge time,  $m$  is the mass of active material and  $V$  is the total voltage window.

## Results and discussion

A brief overview of the present literature on the developments of heteroatom-doped carbon materials, particularly NCMs, shows that most of the efforts remain focused on using templating strategies to achieve high surface areas in these materials.<sup>19,33-35</sup> However, the use of either a hard template such as nanosized silica particles,<sup>19</sup> or soft templates such as use of molten salts,<sup>36</sup> have their own drawbacks<sup>37</sup> and there is a huge controversy about calling such doped carbon materials true metal free systems.<sup>26</sup> In the present work, an ammonium based organic ionic dye  $([\text{NH}_4^+][\text{C}_8\text{H}_4\text{N}_5\text{O}_6^-])$  is used to exploit the benefits of the *in situ* release of gaseous products  $(CO_2$  and  $NH_3)$  that produce highly porous carbon structures without involving a foreign additive as a template.

Ammonium purpurate (ammonium 2,6-dioxo-5-[(2,4,6-trioxo-5-hexahydropyrimidinylidene)amino]-3H-pyrimidin-4-olate), commonly known as murexide, is a commercially available OID with molecular formula  $C_8H_8N_6O_6$  as shown in Fig. 1. It is frequently used as an indicator for complexometric titrations in water hardness measurements as well as a colorant/pigment in the dye industry. Fig. S1 (ESI†) displays the typical trace for thermogravimetric analysis (TGA) of murexide that was recorded from room temperature to 1000 °C under Ar atmosphere. The initial weight loss of about 6–7% up to 230  $\degree$ C can be attributed to the loss of moisture in this material. The major weight loss (up to 90%) occurred in two steps from 230  $\degree$ C to



Fig. 1 Structural formula of murexide

440 °C, leaving 10% carbon residue at temperatures over 600 °C. The high thermal stability of the ammonium purpurate has been reported to be due to several solid–solid phase transitions arising from the mobile ammonium ion, which forms hydrogen bonding networks in different ways.<sup>38</sup> All NCMs were therefore prepared at temperatures over 600 °C (*i.e.* 700, 800 and 900 °C) to ensure the complete carbonization of the template free precursor.

Fig. 2 shows Raman spectra collected for the newly synthesized NCMs with increasing annealing temperature from 700  $\mathrm{^{\circ}C}$ to 800 °C to 900 °C (NCM<sub>(MDE)</sub>-700, NCM<sub>(MDE)</sub>-800 and NCM(MDE)-900, respectively). All samples have two distinct peaks at 1358  $\text{cm}^{-1}$  and 1590  $\text{cm}^{-1}$ , designated as D and G bands, respectively. In general, the D band arises from the breathing mode of  $sp<sup>2</sup>$  carbons on edges or defects indicating a degree of disorder, whereas the stretching motion of  $sp^2$ carbons generates the graphitic G band.39,40 The level of disorder in the carbon structure due to the incorporation of dopant heteroatoms can be determined from the intensity ratio of D and G bands.<sup>41</sup>

The Raman spectra of these NCMs showed a typical trend of decreasing  $I_D/I_G$  intensity ratio from NCM<sub>(MDE)</sub>-700  $\rightarrow$  $NCM_{(MDE)}-800 \rightarrow NCM_{(MDE)}-900$ . This in turn suggests that the NCMs synthesized at higher carbonization temperatures achieve higher degree of graphitization as indicated by the lower  $I_D$  $I_{\rm G}$  intensity ratio<sup>42</sup> values as shown in Table 1.

The formation of graphitic phases and the level of disorder in the carbon structure introduced by the nitrogen heteroatom



Fig. 2 Raman spectra of NCM<sub>(MDE)</sub>-X annealed at  $X = 700$  °C, 800 °C and  $900 °C$ 

Table 1  $I_D/I_G$  ratios calculated for NCM<sub>(MDE)</sub>-700, NCM<sub>(MDE)</sub>-800 and  $NCM<sub>(MDE)</sub> - 900$ 



Fig. 3 XRD patterns of NCM<sub>(MDE)</sub>-700, NCM<sub>(MDE)</sub>-800 and NCM<sub>(MDE)</sub>-900.

were also investigated by XRD. From Fig. 3, it can be seen that the XRD patterns of all three NCMs (i.e.,  $NCM_{\text{[MDE]}}$ -700, NCM(MDE)-800 and NCM(MDE)-900) display a diffuse peak centred around *ca*.  $2\theta = 26^{\circ}$  and two sharp peaks *ca*.  $2\theta = 44^{\circ}$ , and  $2\theta = 64.2^{\circ}$ . In the case of the former, the diffuse peak at *ca*.  $2\theta =$  $26^\circ$  corresponds to the (002) plane, while the sharp XRD reflection at  $2\theta = 44^{\circ}$  is consistent with the (100/101) set of planes, and the  $2\theta = 64.2^{\circ}$  peak is consistent with the (103) plane of hexagonal graphitic carbon.<sup>43,44</sup> The presence of the broad (002) peak together with the sharp peaks (100/101) and (103) in the XRD patterns of these NCMs indicates the coexistence of amorphous and crystalline phases. The former phase is related to the disordered structure, manifesting heteroatom (nitrogen) doping, while the latter phase confirms the level of ordered structure maintained by these NCMs even at high nitrogen content. The existence of a crystalline zone is further evidenced by SAED patterns (Fig. S3†). It has been reported that the 002 peak broadens upon high nitrogen doping in the graphitic carbon framework.<sup>45</sup> The full width at half maximum (FWHM) of this peak (5.9 for  $NCM<sub>(MDE)</sub>$ -700, 5.3 for  $NCM<sub>(MDE)</sub>$ -800 and 4.8 for  $NCM<sub>(MDE)</sub>$ -900) decreases with the increase in annealing temperature from 700  $^{\circ}$ C to 800  $^{\circ}$ C to 900 °C. This observation suggests that the high temperature carbonization leads to a high degree of graphitization (due to low nitrogen content).<sup>4</sup> A number of previously reported heteroatom doped carbon materials (such as nitrogen-doped,<sup>46</sup> nitrogen/sulphur co-doped<sup>47</sup> and nitrogen/phosphorous codoped<sup>19</sup>) display XRD patterns with only broad diffuse peaks, indicating highly disordered structures with amorphous carbon as the dominant phase. It is important to note that the NCMs synthesized in the present study were washed with dilute HCl, thus excluding any possibility of the presence of metallic components in them. In this scenario, the emergence of

crystalline phases indicates a very distinct feature of these materials, which might have an influence on the superior electrochemical performances displayed by these NCMs as discussed later.

Fig. 4(a) displays the comparison of typical X-ray photoelectron spectroscopy (XPS) survey scans for  $NCM_{(MDE)}$ -700,  $NCM<sub>(MDE)</sub>$ -800 and  $NCM<sub>(MDE)</sub>$ -900, indicating the presence of C 1s, N 1s, and O 1s in these newly synthesized NCMs.

Table 2 shows the amount of nitrogen, carbon and oxygen as determined from the core level XPS spectra of  $NCM<sub>(MDE)</sub>$ -700,  $NCM<sub>(MDE)</sub>$ -800 and  $NCM<sub>(MDE)</sub>$ -900. As expected, the atomic percentage of doped nitrogen content decreased consistently from 12.17 to 9.40 to 4.72% as temperature increased for  $NCM<sub>(MDE)</sub>$ -700 to  $NCM<sub>(MDE)</sub>$ -800 to  $NCM<sub>(MDE)</sub>$ -900 materials, respectively. The elemental analysis given in Table S1 (in the ESI†) is also in agreement with this trend. This clearly indicates that high annealing temperature has a detrimental effect on the content of nitrogen doped in these materials. To identify the nature of nitrogen functionalities, the N 1s peak region in the XPS survey spectra of all NCMs was deconvoluted and fitted with three component peaks, *ca.* (398.27  $\pm$  0.1 eV), (399.93  $\pm$  0.1 eV) and (401.00  $\pm$  0.1 eV), which are ascribed to pyridinic-N (N-6), pyrrolic-N (N-5) and quaternary-N (N-Q), respectively for  $NCM<sub>(MDE)</sub>$ -800 (Fig. 4(b-d)).<sup>48,49</sup> These nitrogen functionalities are involved in different types of chemical bonding with their neighbouring carbon atoms, which play a major role in the superior electrochemical performances of these materials. Table 3 shows that the  $NCM<sub>(MDE)</sub>$ -700 has the highest amount of pyridinic nitrogen (N-6) content (50.19%), compared to NCM(MDE)-800 (46.33%) and NCM(MDE)-900 (41.06%). An opposite trend is observed for pyrrolic-N (N-5) content in these NCMs with increasing carbonization temperature, where the XPS data in Table 3 shows that the  $NCM<sub>(MDE)</sub>$ -900 contains the highest proportion of pyrrolic-N (N-5) (39.88%) compared to  $NCM<sub>(MDE)</sub>$ 800 (31.46%) and NCM(MDE)-700 (30.34%). Interestingly, the change in carbonization temperature does not show much impact on the amount of quaternary-N (N-Q) in these NCMs. The XPS data in Table 3 also reveals that the  $NCM<sub>(MDE)</sub>$ -700 and  $NCM<sub>(MDE)</sub>$ -900 have nearly the same proportion of quaternary-N (N-Q) content (19.47  $\pm$  0.05% and 19.07  $\pm$  0.05%, respectively) and the highest content of quaternary-N (N-Q) is present in the  $NCM<sub>(MDE)</sub>$ -800 sample (22.2%). It has been previously reported that the pyridinic-N (N-6) and pyrrolic-N (N-5) type nitrogen functionalities can deliver free electrons to the conduction band, which contribute in the pseudocapacitive faradaic reactions.<sup>50,51</sup> The most stable nitrogen configuration compared to other nitrogen functionalities, quaternary-N (N-Q), accounts for the graphitic component intrinsically displaying high electronic conductivity and enhancing capacitance by creating active sites and helping electron transportation.<sup>19,52,53</sup> The NCM<sub>(MDE)</sub>-800 material, having the highest quaternary-N (N-Q) content in addition to a reasonable proportion of the pyridinic-N (N-6) content, presents superior performance for supercapacitor application. Puper<br>
Total 1.6, 2005 cacuated for HCP<sub>Pack</sub>-700 HCP<sub>Pack-</sub>900 and capabilities, which registeres are the teachers are<br>  $\frac{1}{2}$   $\frac{1}{2}$   $\frac{1}{2}$   $\frac{1}{2}$   $\frac{1}{2}$   $\frac{1}{2}$   $\frac{1}{2}$   $\frac{1}{2}$   $\frac{1}{2}$   $\frac{1}{2}$ 

> Field emission scanning electron microscopy (FE-SEM) was used to examine the morphology of the synthesized  $NCM_{(MDE)}-X$ materials. The SEM micrographs at low and high magnification



Fig. 4 (a) XPS survey spectra of synthesized NCM<sub>(MDE)</sub>-700, NCM<sub>(MDE)</sub>-800 and NCM<sub>(MDE)</sub>-900; (b) N 1s deconvoluted spectrum for NCM<sub>(MDE)</sub>-700; (c) N 1s deconvoluted spectrum for NCM<sub>(MDE)</sub>-800; (d) N 1s deconvoluted spectrum for NCM<sub>(MDE)</sub>-900.

Table 2 The amount of N 1s, C 1s and O 1s in NCMs as determined from the core-level XPS spectra of NCM<sub>(MDE)</sub>-700, NCM<sub>(MDE)</sub>-800 and  $NCM<sub>(MDF)</sub> - 900$ 

Sample ID	N 1s (%)	C 1s (%)	O 1s (%)
$NCM(MDE)$ -700	12.17	73.14	14.69
$NCM(MDE)$ -800	9.40	75.63	14.97
$NCM(MDE)$ -900	4.72	79.42	15.86

Table 3 The values of pyridinic-N (N-6), pyrrolic-N-5 and quaternary-N (N-Q) in atomic % as determined from the deconvolution of N 1s component in the XPS spectra of  $NCM_{(MDE)}$ -700,  $NCM_{(MDE)}$ -800 and  $NCM<sub>(MDE)</sub> - 900$ 



in Fig. 5 show loosely stacked graphitic sheet-like morphology for all materials; in addition to sheets, some globular structures were also formed. Fig. S2 (in the ESI†) displays the energydispersive X-ray spectroscopy elemental mapping of  $NCM<sub>(MDE)</sub>$ -800, indicating the homogeneous distribution of nitrogen doping throughout the material. For charge transfer on the large electrode/electrolyte interface, the nanosheet architecture offers minimum resistance to mass transport, which is an important parameter for supercapacitor materials because shorter diffusion traveling distances ease ion

transport.<sup>54-56,56-58</sup> HRTEM images for NCM<sub>(MDE)</sub>-800 show that the material has nano graphitic sheets as shown in Fig. 6. The images also show the rough texture due to the porosity of the material, as confirmed by the surface area and porosity analysis described later. Selected area electron diffraction (SAED) images show the crystalline and amorphous nature of the material, depending on the region (Fig. S3(b and d) in ESI†). The SAED pattern of the crystalline phase presents defined rings with bright spots from the diffraction of the lattice. The average d-spacing values calculated from the SAED pattern are 0.22 nm (101), 0.15 nm and 0.12 nm. The conductive graphitic layers provide paths for fast electron transport while nanopores serve as active sites for ions.<sup>59</sup> All these characteristics are important for excellent supercapacitive electrode materials.

The surface area and porosity characteristics of the synthesized NCM<sub>(MDE)</sub>-X were investigated by  $N_2$  adsorption-desorption isotherms Fig. S4 (in the ESI†). It can be seen that the shapes of the isotherms were nearly identical for all  $NCM<sub>(MDE)</sub> - X$ materials and can be categorized as type IV with a characteristic type H4 hysteresis loop at  $p/p^{\circ} > 0.4$ , which is indicative of the presence of micropores and mesopores.<sup>60,61</sup> The nearly linear middle portion in these isotherms signifies multilayer adsorption, while the hysteresis shows capillary condensation in the mesopores.<sup>62</sup> Interestingly, in NCM<sub>(MDE)</sub>-700, the steep region of the desorption branch reached a non-closure point at lower relative pressure, indicating irreversible interaction between  $N_2$ and the absorbent. However, at higher relative pressure, the almost linear uptake indicates the existence of macropores.<sup>62</sup> The BET data shown in Table S2 $\dagger$  reveal that the NCM<sub>(MDE)</sub>-700,  $NCM<sub>(MDE)</sub>$ -800,  $NCM<sub>(MDE)</sub>$ -900 exhibited a surface area  $\sim$ 128, 287 and 307  $m^2$   $g^{-1}$  and average pore diameters of 11.46, 10.09



Fig. 5 Low and high magnification FE-SEM images of NCM<sub>(MDE)</sub>-700 (a, d), NCM<sub>(MDE)</sub>-800 (b, e) and NCM<sub>(MDE)</sub>-900 (c, f)



Fig. 6 Low and high magnification HR-TEM images of  $NCM_{(MDE)}$ -800

and 9.39 nm, respectively. Fig. S5 (in the ESI†) shows the pore size distribution (PSD) profiles for  $NCM_{(MDE)}$ -700,  $NCM_{(MDE)}$ -800, NCM $_{(MDE)}$ -900, with the inset displaying the PSD profiles for these materials up to the 10 nm scale, indicating the porosity of all NCMs lying in the mesoporous region (pore diameter < 50 nm). Macropores and mesopores serve as ion buffering reservoirs and decrease ion transportation distance, while micropores contribute to energy storage.<sup>60,63,64</sup> The detailed porosity analysis results of NCM<sub>(MDE)</sub>-X are summarized in Table S2.<sup>†</sup>

The BET results (Table S2†) show that in going from NCM- $700 \rightarrow NCM-800$ , the increase in surface area is around 160 m<sup>2</sup>  $g^{-1}$ , whereas the reduction in particle size is around 26 nm. At further high annealing temperature, in going from

NCM-800  $\rightarrow$  NCM-900, the increase in the surface area was about 20  $\mathrm{m^2\,g^{-1}}$ , whereas the reduction in particle size was only 1.38 nm. This comparison suggests that the particle size reduction was smaller in going from NCM-800  $\rightarrow$  NCM-900 when compared to particle size reduction in going from NCM- $700 \rightarrow NCM-800$ . Table S2<sup>†</sup> also shows that the total pore volume shows a similar incremental change of 0.03 cm<sup>3</sup>  $g^{-1}$  in going from NCM-700  $\rightarrow$  NCM-800 and further from NCM-800  $\rightarrow$  NCM-900. From this observation, it can be concluded that the increase in pore volume in going from NCM-800  $\rightarrow$ NCM-900 contributed more to the increase in surface area of about 20  $m^2 g^{-1}$  in NCM-900 sample, which in other cases could not be explained based on the particle size reduction. There is another correlation that can be seen between the total pore volume and the  $I_D/I_G$  values from Raman studies, where an increase in annealing temperature causes a low  $I_D/I_G$  value and corresponding high total pore volume in these NCMs. The loss of entrapped nitrogen at elevated temperature causes increased porosity and improves the graphitic character as indicated by the low  $I_{\rm D}/I_{\rm G}$  value.

The electrochemical performances of the  $NCM<sub>(MDE)</sub>$ -700, NCM(MDE)-800 and NCM(MDE)-900 were tested as electrode materials for supercapacitor application in a three-electrode configuration cell using  $Ag/AgCl<sub>(saturated)</sub>$  as the reference electrode and Pt as the counter electrode. In brief, the working electrodes were cast with 2 mg of electroactive material loaded on a graphite monolith substrate (details in the Experimental section). The initial set of electrochemical studies was performed in 1.0 M  $H_2SO_{4(aq)}$  within the voltage window from  $-0.2$ and  $+0.8$  V vs. Ag/AgCl. Fig. 7(a) shows the CV profiles recorded for NCM<sub>(MDE)</sub>-700, NCM<sub>(MDE)</sub>-800 and NCM<sub>(MDE)</sub>-900 at the scan rate of 25 mV  $s^{-1}$ , where all the materials exhibited nearly rectangular shapes, which typically represent the contribution from electrical double layer capacitance (EDLC). In Fig. 7(a), the  $NCM<sub>(MDE)</sub>$ -700 and  $NCM<sub>(MDE)</sub>$ -800 exhibit slightly curved CV



Fig. 7 Comparative analysis of the electrochemical data for NCM<sub>(MDE)</sub>-700, NCM<sub>(MDE)</sub>-800 and NCM<sub>(MDE)</sub>-900 in 1.0 M H<sub>2</sub>SO<sub>4(aq)</sub> electrolyte: (a) cyclic voltammograms at 25 mV s<sup>-1</sup>; (b) galvanostatic charge–discharge cycles at 3 A g<sup>-1</sup>; electrochemical impedance spectra, and (c) Nyquist plot (inset: magnified region  $0-0.8 \Omega$ ).

profiles indicating redox reactions possibly coming from the interactions between the doped nitrogen functionalities and the  $H^+$  in the acidic electrolyte media.<sup>1</sup> From CV profiles, the current densities of  $NCM_{(MDE)}$ -X electrodes increase in the following order:  $NCM_{(MDE)}$ -800 >  $NCM_{(MDE)}$ -700 >  $NCM_{(MDE)}$ -900. The high current density displayed by NCM<sub>(MDE)</sub>-800 can be attributed to the overall high nitrogen content (9.40%) and moderate surface area (287  $\mathrm{m^2\,g^{-1}}$ ) of this material.

The capacitance performances of the  $NCM<sub>(MDE)</sub>$ -X materials were further investigated by carrying out the galvanostatic charge–discharge (GCD) experiments at different current densities in the voltage window between  $-0.2$  and  $+0.8$  V vs. Ag/ AgCl. Fig. 7(b) shows the GCD profiles for  $NCM<sub>(MDE)</sub>$ -700, NCM(MDE)-800 and NCM(MDE)-900 at the current density of 3 A  $\rm g^{-1}.$  These GCD curves display a negligible ohmic drop (iR), which is indicative of low internal resistance in these materials. According to eqn (1), the calculated specific capacitance  $(C_s)$ value at 3 A  $\rm g^{-1}$  for NCM $_{\rm (MDE)}$ -800 is 222 F  $\rm g^{-1},$  which is higher than 138, and 111 F  $g^{-1}$  for NCM<sub>(MDE)</sub>-700 and NCM<sub>(MDE)</sub>-900, respectively. There are a number of factors that contribute to the superior capacitive performance of  $NCM<sub>(MDE)</sub>$ -800 compared to  $NCM_{(MDE)}$ -700 and  $NCM_{(MDE)}$ -900; these can be broadly categorized as high surface area (287  $\mathrm{m}^2\ \mathrm{g}^{-1}$ ), appreciable pore volume (0.13  $\rm cm^3~g^{-1}$ ), high doped nitrogen content of 9.4% with highest quaternary nitrogen functionality, and the highest micropore to total pore volume ratio (0.85). All these factors have their own influence on the capacitive performance of the materials: high surface area offers more charge storage platform, high nitrogen content offers more pseudocapcitive

contribution in the overall charge storage, quaternary nitrogen functionality is the most stable and conductive type among other nitrogen functionalities, and lastly, the highest micropore to total pore volume ratio (0.85), which suggests ease of mass transport of ions from electrolyte through porous carbon network with greater access to ion-accessible surface area and number of electroactive sites. The supercapacitor performances of recently reported nitrogen doped carbon materials synthesized from different precursors are given in Table 4 for comparison purposes.

Electrochemical impedance spectroscopy (EIS) is a very powerful technique to investigate the electronic and electrochemical responses of an electrode in contact with an electrolyte. From Fig. 7(c), we can see that the Nyquist plots for all NCMs exhibit quasi-semicircles in the high-frequency region followed by linear traits at low-frequencies. The quasisemicircle at high-frequencies is visible for  $NCM<sub>(MDE)</sub>$ -700, and is significantly reduced in  $NCM<sub>(MDE)</sub>$ -800 and it is practically absent in the case of the  $NCM_{(MDE)}$ -900 plot. Meanwhile, the near-linear spike in the case of both  $NCM<sub>(MDE)</sub>$ -800 and  $NCM<sub>(MDE)</sub>$ -900, with slope approaching closer to 90 $^{\circ}$  in the low frequency region, suggests an improvement in the capacitive behaviour with fast charge carrier diffusion (high electronic conductivity) in these materials upon electrode polarization.<sup>65-67</sup> The lowest ESR value for  $NCM<sub>(MDE)</sub>$ -900 indicates high electronic conductivity and low overall cell resistance for this material, which might have its origin in the higher degree of graphitization due to low levels of heteroatom doping.<sup>44</sup> The Bode plots display the dependence of the phase angle on the





frequency. In general, the phase angle between  $90^\circ$  and  $45^\circ$  is caused by the capacitive behaviour of a system; values close to 90° are observed in the response of an ideal capacitor. Fig. S6<sup>†</sup> shows that the  $NCM<sub>(MDE)</sub>$ -800 electrode has the highest phase angle, almost  $86^\circ$  at 10 mHz AC frequency, close to the ideal capacitor behaviour.

Since high power demanding applications require the charging and discharging of electrode material at high current densities, high capacitance retention at high scan rates or at high current densities is key for practical applications. From Fig. 8(a), we can see that the CV curves for  $NCM<sub>(MDE)</sub>$ -800 in 1 M  $H<sub>2</sub>SO<sub>4(aq)</sub>$  exhibit a linear increase in current density with increasing scan rate from 5 mV  $s^{-1}$  to 200 mV  $s^{-1}$ . Fig. 8(b) displays the galvanostatic charge–discharge (CD) curves for NCM(MDE)-800 with increasing current density values from

3 A  $g^{-1}$  to 50 A  $g^{-1}$ . The charging and discharging segments of GCD curves display symmetrical triangles, even at high current density values, indicating fast charge–discharge dynamics and good capacitive performances. Fig. 8(c) displays the values of specific capacitance for all NCMs as a function of the current density. From Fig.  $8(c)$ , a rapid decrease in the specific capacitance of  $NCM<sub>(MDE)</sub>$ -700 can be observed with an increase in current density from 3 to 50 A  $g^{-1}$ . The poor micropore/total pore volume ratio for this material indicates fewer micropores in this material, which could be one reason for the decreasing specific capacitance at high current density. In general, fast electrolyte dynamics at high density require adequate levels of micropores in the material to allow facile trafficking of electrolyte ions and their interactions with the active sites at the material interface. The other high temperature materials, viz.



Fig. 8 (a) Cyclic voltammograms of NCM<sub>(MDE)</sub>-800 in 1.0 M H<sub>2</sub>SO<sub>4(aq)</sub> at different scan rates from 5 to 200 mV s<sup>-1</sup>. (b) Galvanostatic charge– discharge curves for NCM<sub>(MDE)</sub>-800 at current density values ranging from 3 to 50 A g<sup>-1</sup>. (c) Specific capacitances of NCM<sub>(MDE)</sub>-X at different current densities. (d) Cyclic stability test: specific capacitance calculated from discharge curves of 10 000 cycles at a current density of 10 A g $^{-1}\!$ . (inset) galvanostatic charge–discharge curves of the 1<sup>st</sup> and 10000<sup>th</sup> cycles for NCM<sub>(MDE)</sub>-800 at a current density of 10 A g<sup>-1</sup>.

 $NCM<sub>(MDE)</sub>$ -800 and  $NCM<sub>(MDE)</sub>$ -900, display specific capacitances of 161 F  $g^{-1}$  and 106 F  $g^{-1}$  at a higher current density of 50 A  $\mathrm{g}^{-1}$ , which is 87% and 92% of their specific capacitances at 10 A  $g^{-1}$ , respectively. From Fig. S7 (ESI†), the CV profiles for  $NCM<sub>(MDE)</sub>$ -800 and  $NCM<sub>(MDE)</sub>$ -900  $NCM<sub>(MDE)</sub>$ -X also exhibit high rate capability, since no distortion was observed in the CV curves for these materials with increasing the scan rate from 5 to 200 mV s $^{-1}$ . High capacitance retention of NCM $_{\rm (MDE)}$ -800 and NCM(MDE)-900 might have its origin in the adequate size distribution of micro/mesopores in the material, providing better access for the electrolyte and allowing fast ion trafficking at high scan rates.<sup>68</sup> The long term cycling stability of  $NCM_{(MDE)}$ -800 was also tested by carrying out the GCD experiment up to 10 000 cycles at the current density of 10 A  $\rm g^{-1}$ . Fig. 8(d) shows the corresponding plot of specific capacitance versus cycle number for NCM<sub>(MDE)</sub>-800 within the  $-0.2$  V to 0.8 V potential window and exhibiting 100% capacitive retention for this material. From this plot, it is evident that the initial value of the specific capacitance of 195 F  $g^{-1}$  progressively increased with GCD cycling and reached a value of 203 F  $g^{-1}$  after 10 000 cycles. Moreover, the triangular shape of the  $10,000$ <sup>th</sup> GCD cycle, shown in Fig. 8(d) inset, also indicates that the  $NCM<sub>(MDE)</sub>$ -800 electrode has good charge propagation and exhibits stable performance over extended periods of time.<sup>69</sup>

In Fig. S8,† the EIS measurements on  $NCM<sub>(MDE)</sub>$ -800 before and after 10 000 GCD cycles shows an improvement in the capacitance as indicated by the increase in phase angle (Bode plot). Overall, it can be concluded that long term cycling serves as a means of electrochemical activation under the influence of an electric field, which may allow better penetration of the hydrated electrolyte ions into the micropores.<sup>75,76</sup>

As a potential electrode material for supercapacitor application, further electrochemical studies were carried out for  $NCM<sub>(MDE)</sub>$ -800 in three different electrolyte solutions to investigate the effect of solution pH and the nature of ions from the electrolyte. Fig. 9(a) displays the CV profiles for  $NCM<sub>(MDE)</sub>$ -800 recorded in 1 M H<sub>2</sub>SO<sub>4(aq)</sub>, 6 M KOH<sub>(aq)</sub> and 0.5 M Na<sub>2</sub>SO<sub>4(aq)</sub> electrolytes with respective pH values of 0.87, 13.8 and 7.9, at the scan rate of 25 mV  $s^{-1}$ . The CV profiles recorded in 1 M  $H<sub>2</sub>SO<sub>4(aq)</sub>$  and 6 M KO $H<sub>(aq)</sub>$  have the rectangular shaped curves with superimposed redox peaks due to surface functionalities. These redox peaks are associated with different types of chemical interactions formed between the surface nitrogen functionalities and the active ions in the electrolyte media. Fig. 10 shows an illustration of possible types of ionic interactions that can occur between the surface nitrogen functionalities (N6/N5) and the ions in the acidic and basic media during potential cycling, which in turn explain the pseudocapacitive contribution of the nitrogen moieties in these materials.<sup>19,61</sup> In contrast, the CV curve recorded for  $NCM_{(MDE)}$ -800 in neutral electrolyte  $(0.5 M Na<sub>2</sub>SO<sub>4(aq)</sub>)$  exhibits the double layer capacitance with the absence of redox peaks. Moreover, NCM<sub>(MDE)</sub>-800 achieves higher current density in 1 M  $H_2SO_{4(aq)}$  compared to 6 M KO $H_{(aq)}$ , while the lowest current density is observed in 0.5 M Na<sub>2</sub>SO<sub>4(aq)</sub> electrolyte. The redox peaks from the CV profiles of NCM<sub>(MDE)</sub>-800 in 1 M  $H_2SO_{4(aq)}$  and 6 M KOH<sub>(aq)</sub> indicate that the carbon atoms with Lewis basicity next to nitrogen and the lone pair of electrons on doped nitrogen itself likely develop coulombic interactions with the active proton  $(H<sup>+</sup>)$  in the acidic media or hydroxide  $(OH<sup>-</sup>)$  in the basic media at the initial stage. Later on, these ionic interactions may lead to reversible redox reactions, as evident from the CV profiles of RSC Advances<br>
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Fig. 9 The effect of pH of different electrolytes (i.e.; 1.0 M H<sub>2</sub>SO<sub>4</sub> (0.87 pH), 0.5 M Na<sub>2</sub>SO<sub>4</sub> (7.9 pH) and 6 M KOH (13.8 pH)) on NCM<sub>(MDF)</sub>-800. (a) Cyclic voltammograms at 25 mV s $^{-1}$ . (b) Galvanostatic charge–discharge curves at 3 A g $^{-1}$  current density. (c) Specific capacitances of NCM<sub>(MDE)</sub>-800 at different current densities and (d) % capacitance retention up to 1000 cycles at 5 A  $g^{-1}$  current density.



Fig. 10 Illustration of possible types of ionic interactions between the surface nitrogen functionalities (N6/N5) and the ions in the acidic and basic media during potential cycling.

 $NCM<sub>(MDE)</sub>$ -800 in acidic and basic media,<sup>26</sup> as shown in Fig. 10. The corresponding CV curves for these materials at the scan rates of 5, 50, 100 and 200 mV  $s^{-1}$  are shown in Fig. S9.†

In Fig. 9(b), the GCD curves recorded for  $NCM<sub>(MDE)</sub>$ -800 in these three different electrolyte media at current density  $\sim$ 3 A g<sup>-1</sup> also support the results from CV measurements with much longer discharging time observed in 1 M  $H_2SO_{4(aq)}$ , compared to 6 M KOH<sub>(aq)</sub> and 0.5 M Na<sub>2</sub>SO<sub>4(aq)</sub>. The values of specific capacitance calculated for NCM<sub>(MDE)</sub>-800 at 3 A  $g^{-1}$  in 1 M  $H_2SO_{4(aq)}$ , 6 M KO $H_{(aq)}$  and 0.5 M Na<sub>2</sub>SO<sub>4(aq)</sub> electrolytes are 222, 126 and 73 F  $g^{-1}$ , respectively.

The specific capacitance values recorded for  $NCM<sub>(MDE)</sub>$ -800 in 1 M  $H_2SO_{4(aq)}$ , 6 M KO $H_{(aq)}$  and 0.5 M Na<sub>2</sub>SO<sub>4(aq)</sub> electrolytes at different current densities (from 3 to 20 A  $\rm g^{-1})$  are shown in Fig. 9(c). Interestingly, the  $NCM_{(MDE)}$ -800 exhibits a notable decrease in  $C_s$  from 3 A  $g^{-1}$  to 5 A  $g^{-1}$  in the case of both acidic (1 M  $H_2SO_{4(aq)}$ ) and basic (6 M KO $H_{(aq)}$ ) electrolytes, whereas a less rapid decrease in  $C_s$  was observed for the CD profiles recorded in 0.5 M  $\text{Na}_2\text{SO}_4$ <sub>(aq)</sub> electrolyte within the same current density range. However, if we compare the rate capability of NCM(MDE)-800 in the three electrolytes with current density between 5 and 20 A  $g^{-1}$ , this material achieved the highest capacitance retention of  $\sim$ 91% in 1 M H<sub>2</sub>SO<sub>4(aq)</sub>, compared to 78% in 0.5 M Na<sub>2</sub>SO<sub>4(aq)</sub> and 68% in 6 M KOH<sub>(aq)</sub>.

Fig. 9(d) displays the cyclic stability test for  $NCM_{(MDE)}$ -800 up to 1000 GCD cycles in all three electrolytes, viz. 1 M  $H_2SO_{4(aq)}$ , 6 M KOH<sub>(aq)</sub> and 0.5 M Na<sub>2</sub>SO<sub>4(aq)</sub> at a current density of 5 A g<sup>-</sup> . The results reveal that the  $NCM<sub>(MDE)</sub>$ -800 has excellent capacitive retention (98%) in 0.5 M  $Na<sub>2</sub>SO<sub>4</sub>$ , up to 1000 cycles. In contrast, a notable decay in  $C_s$  with almost 96% capacitance retention was observed after 200 cycles for the GCD measurements carried out in 6 M KOH. Interestingly, Fig. 9(d) shows a gradual improvement in the specific capacitance (194 to 220 F  $g^{-1}$ ) for NCM<sub>(MDE)</sub>-800 in 1 M H<sub>2</sub>SO<sub>4(aq)</sub>, which is nearly 114% of the value of the capacitance of this material after 1000

cycles. These long cycle stability tests indicate that the ionic strengths, electrolyte pH and the nature of ions involved in the charge storage mechanism, all contribute to the superior capacitive retention displayed by  $NCM_{(MDE)}$ -800.

The H<sup>+</sup> ion has the smallest hydrated ionic radius  $(1.0 \text{ Å})$  and highest ionic mobility in water ( $\mu = 36.23 \times 10^{-8}$  m<sup>2</sup> s<sup>-1</sup> V<sup>-1</sup> at 298 K) compared to  $K^+$  (ionic radius (3.31 Å) and ionic mobility  $(\mu = 7.62 \times 10^{-8} \text{ m}^2 \text{ s}^{-1} \text{ V}^{-1}$  at 298 K)) and Na<sup>+</sup> (ionic radius (3.58 Å) and ionic mobility ( $\mu = 5.19 \times 10^{-8} \text{ m}^2 \text{ s}^{-1} \text{ V}^{-1}$  at 298 K)).<sup>77,78</sup> Due to the small ionic size and ultra-fast mobility, the H<sup>+</sup> can penetrate deep into the pores of NCM<sub>(MDE)</sub>-800, thereby, contributing to the high rate capability of this material at all scan rates. In summary, the  $H_2SO_{4(aq)}$  electrolyte is a suitable choice for NCM type supercapacitor materials, due to the presence of active protons with small ionic radius and ultrafast mobility, in addition to the capability to develop ionic interactions with the carbon atoms with Lewis basicity adjacent to the nitrogen and the nitrogen itself.

## Conclusions

The present work has shown that without the requirement of an expensive template, murexide is a good precursor for the preparation of hierarchical porous nitrogen-doped carbon materials (NCMs), which can be used as active electrode materials for supercapacitor application. The  $NCM<sub>(MDE)</sub>$ -800 shows superior capacitive performance ( $C_s \sim 222$  F g<sup>-1</sup> at the current density of 3 A  $\rm g^{-1})$ , excellent cyclic stability (10 000 cycles at the current density of 10 A  $g^{-1}$ ) and good capacitance retention (91%) at current density as high as 20 A  $\rm g^{-1}$ . The comparison of capacitive performance for the three materials indicates that the high proportion of pyridinic-N and quaternary-N functionalities accompanied by reasonable surface area and hierarchical porosity are the major factors for achieving high specific capacitance by  $NCM<sub>(MDE)</sub>$ -800. The detailed investigation of this material in three different electrolytes (acidic, basic and neutral) with current density from 5 to 20 A  $g^{-1}$  reveal that the capacitance retention in 1 M  $H_2SO_{4(aq)}$  (91%) is significantly higher than the capacitive retention in 0.5 M  $Na<sub>2</sub>SO<sub>4(aq)</sub>$  (78%) and 6 M KOH $_{(aq)}$  (68%). The electrochemical analysis reveals that the 1 M  $H_2SO_{4(aq)}$  is the most suitable electrolyte for a supercapacitor based on N-doped carbon materials, due to the active proton with small ionic radius, ultra-fast mobility and the capability to develop ionic interactions with carbon atoms next to the nitrogen (with Lewis basicity) and the nitrogen itself.

# Conflicts of interest

There are no conflicts to declare.

## Acknowledgements

The authors acknowledge the Stephenson Institute for Renewable Energy, University of Liverpool, United Kingdom for offering their research facilities and Higher Education Commission of Pakistan for indigenous and overseas financial assistance Grants No. (112-37208-2PS1-414) and (IRSIP 30-PS-23). The authors

would also like to extend their sincere appreciation to the Deanship of Scientific Research at the King Saud University for its funding of this research through the Research Group, Project No. RGP-255. We acknowledge the support from Science without Borders Program, Ministério da Educação, Coordenação de Aperfeiçoamento de Pessoal de Nível Superior (CAPES); and the Engineering and Physical Sciences Research Council (EPRSC) under grant EP/N032888/1. We thank Dr R. Clowes for providing BET analyses, and the Nanoinvestigation center at Liverpool (NiCaL) for access to the TEM and SEM and scientific discussions with Dr V. Dhanak. FISC Articles. Properties Articles. Published on 28 November 2017. The Common Access Article is licensed to the Common Access Articles. Access Articles. Access Articles. Access Articles. Access Articles. Access Articles.

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