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N-Enriched carbon nanofibers for high energy density supercapacitors and Li-ion batteries†

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Nitrogen enriched carbon nanofibers have been obtained by one-step carbonization/activation of PAN-based nanofibers with various concentrations of melamine at 800 °C under a N₂ atmosphere. As synthesised carbon nanofibers were directly used as electrodes for symmetric supercapacitors. The obtained PAN-MEL fibers with 5% melamine stabilised at 280 °C and carbonized at 800 °C under a nitrogen atmosphere showed excellent electrochemical performance with a specific capacitance of up to 166 F g⁻¹ at a current density of 1A g⁻¹ using 6 M KOH electrolyte and a capacity retention of 109.7% after 3000 cycles. It shows a 48% increase as compared to pristine carbon nanofibers. Two electrode systems of the CNFM5 sample showed high energy densities of 23.72 to 12.50 W h kg⁻¹ at power densities from 400 to 30 000 W kg⁻¹. When used as an anode for Li-ion battery application the CNFM5 sample showed a high specific capacity up to 435.47 mA h g⁻¹ at 20 mA g⁻¹, good rate capacity and excellent cycling performance (365 mA h g⁻¹ specific capacity even after 200 cycles at 100 mA g⁻¹). The specific capacity obtained for these nitrogen enriched carbon nanofibers is higher than that for pristine carbon nano-fibers.

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1 Introduction

Cost-effective and eco-friendly energy storage systems are the need of the hour due to the limited availability of petroleum resources and escalating energy utilization.¹ Supercapacitors and Li-ion batteries have attracted a great deal of attention because of their simple principles, high durability, high power density, fast charging–discharging mechanism and long cycle life.^{2–6} They have a wide range of applications in digital equipment, electrical cars and pulsing techniques.^{7–9} Carbon based materials are well known as potential candidates for energy storage applications due to their low cost, high conductivity, and ready availability.^{10a–d} Carbon nanotubes, mesoporous carbon, and graphene are widely explored as electrodes for supercapacitors and Li-ion battery applications.^{11,12} These materials show good potential for energy storage but the

tedious synthesis processes and difficulty in scaling up their production restricts their application. Hence, carbon nanofibres produced by electrospinning are a better alternative to these materials.

Electrospinning is a very popular and well known technique used for polymer fibres with varying diameters. The fibres can be electrospun into yarn, well arranged fibrous arrays, or self-standing tapes.^{13–17} Polyacrylonitrile (PAN) gives good quality carbon nanofibres after stabilization and carbonization process. The porosity, presence of hetero-atoms and conductivity of PAN based carbon fibres depends on these stabilization and carbonization processes. Chemical or physical activation of PAN-based carbon materials is well reported previously.^{18–22} When ZnCl₂ or KOH, are used for chemical activation, a lot of impurities remain in the fibers as by-products.¹⁸ Many a times it leads to removal of most of the heteroatom functionalities such as functional groups containing N, S and O in due course of activation.²¹ Lower reaction temperature may be useful for obtaining satisfactory surface structure, but it leads to poor conductivity of materials. As a result, these materials cannot be used as a potential candidate for high power density devices. In case of PAN based materials the biggest advantage of activation process is, nitrogenated and oxygenated functionalities responsible for pseudo-capacitive behaviour last during this process.

In this study, PAN-based carbon fibres were obtained by electrospinning with addition of various concentrations of melamine for N doping. The electrospun fibres were then

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stabilised at 280 °C and carbonized at 800 °C in nitrogen atmosphere. The series of electrodes are prepared and electrochemically characterized in 6 M KOH as electrolyte. PAN-based carbon fibre electrodes show high power density supercapacitors and an anode for Li-ion battery application due to different nitrogen containing functionalities present in it as a result of addition of melamine.

2 Experimental

2.1 Synthesis of PAN/Mel based carbon nanofibres

To fabricate polyacrylonitrile/melamine (PAN/Mel)-based carbon nanofibres by electrospinning, 1% melamine was dispersed in *N,N*-dimethylformamide (DMF). 10 Weight% PAN was added to melamine dispersed DMF solution with continuous stirring and heating to ensure formation of homogeneous solution. This polymeric solution was filled and ejected through a syringe (feed rate was kept at 4 ml h⁻¹) by applying a 25 kV bias between a rotating collector and the syringe needle to form nanofibres. The prepared electrospun PAN/Mel nanofibres were stabilized in air at 280 °C.²³ The stabilized nanofibres were further heated at 800 °C under N₂ atmosphere at a heating rate of 5 °C min⁻¹ for carbonization to obtain the final product (CNFM1). The various weight percent of melamine such as 3% (CNFM3), 5% (CNFM5) and 10% (CNFM10) were added to study its effect on carbon nanofibres. Sample CNF800 was prepared without melamine addition.

2.2 Materials characterisations

The polyacrylonitrile/melamine (PAN/Mel)-based carbon nanofiber's morphology was analyzed by Field Emission Scanning Electron Microscopy (FESEM, Hitachi, S-4700) and Transmission Electron Microscopy (TEM, (Phillips, TECHNAI-F20)). The elemental surface composition of the carbon nanofibers was analyzed by X-ray Photoelectron Spectroscopy (XPS, Thermo Scientific, ESCALAB 250). The chemical structure of the carbon nanofibers was studied by Raman Spectroscopy (NANOBASE, XperRam200, laser line: 532 nm) and X-ray Diffraction (XRD-D8, Advance, Bruker-AXS) with Cu K α radiation.

2.3 Electrochemical measurements

For the electrochemical measurements of supercapacitor application the self standing, flexible tapes of carbon nanofibers (CNF800, CNFM5) without any binding and conductive materials were used as electrode material. These self standing, flexible tapes were cut into 1.5 × 1.5 cm² and were used to form a symmetric supercapacitor device. Filter paper and nickel foil were used as separator and current collectors respectively. 6 M KOH was chosen for the aqueous electrolyte. A multichannel potentiostat/galvanostat (WBCS 3000 battery cycler system, Won-A Tech. Co., Korea) was used to determine the electrochemical properties. A voltage window of 0–1.0 V was used for GCD measurement of the specific capacitance at various current densities ranging between 1 to 30 A g⁻¹. Electrochemical impedance spectroscopy (EIS) study was done by using

impedance analyzer (IM6e, Zahner Elektrik) (frequency range 10 mHz to 100 kHz).

To perform electrochemical measurements for Li-ion battery application 2032 type coin cells were fabricated using the self standing, flexible tapes of carbon nanofibre CNFM5 without any binding and conductive materials as working electrode. CNFM5 tape was cut into circular disk of diameter 16 mm. Quartz filter paper and metallic lithium foil (thickness 75 μ m) were used as separator and counter electrode respectively. 1 M LiPF₆ in dimethyl carbonate (DMC) : ethylene carbonate (EC) (1 : 1 v/v) (BASF) was used as an electrolyte. The half cells were tested using cyclic voltammetry on the potentiostat/galvanostat (Metrohm Autolab) between 0 to 3 V. The galvanostatic charge-discharge behaviour was tested on MTI battery analyzer (vs. Li/Li⁺). Electrochemical impedance spectroscopy (EIS) study was done in the frequency range 0.1 Hz to 1 MHz and amplitude 5 mV.

3 Results and discussion

Fig. 1(a) shows the XRD patterns of the carbon nanofibers synthesized from polyacrylonitrile/melamine (PAN/Mel) at 800 °C. The broad peak around 2 θ ; 23.3° is an indication of presence of graphitic carbon, with disorder. The Raman spectroscopy is used to critically clarify the nature of carbon. Raman spectra for the carbon nanofibers synthesized from polyacrylonitrile/melamine (PAN/Mel) at 800 °C can be seen in Fig. 1(b). The characteristic D and G bands of carbon can be seen around 1320 cm⁻¹ and 1590 cm⁻¹ which can be assigned to the breathing mode of *k*-point phonons of A_{1g} symmetry and E_{2g} phonon of sp² carbon atoms, respectively.²⁴ The I_D : I_G ratio of CNF800, CNFM5 is 0.9564, 0.9353 respectively.

Fig. 2 shows typical scanning electron microscopy (SEM) images of the electrospun PAN/Mel carbon nanofibers with 5% of melamine after stabilization at 280 °C and carbonisation at 800 °C in nitrogen. The average diameter of the PAN/Mel carbon nanofibers with 5% melamine (Fig. 2(a)) is about 450 nm, after stabilisation at 280 °C these electrospun fibers have irregular edges and rough surface which can be seen in SEM images. After carbonisation at 800 °C in nitrogen (Fig. 2(b)), a decrease in diameter (300 nm) was observed. After carbonization the edges of the PAN/Mel fibers had become even and surface appeared smooth, which was also confirmed using transmission electron microscopy (TEM) analysis (Fig. 2(c)).

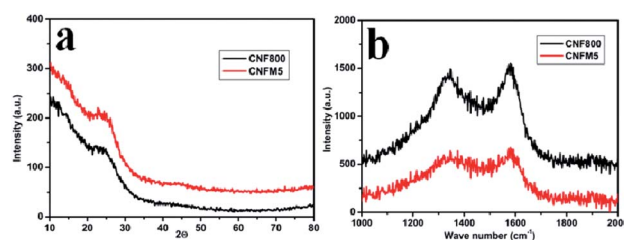


Fig. 1 (a) XRD patterns of PAN/Mel carbon nanofibers without melamine (CNF800) and with 5% melamine (CNFM5), (b) Raman spectra of CNF800 and with 5% melamine (CNFM-5).



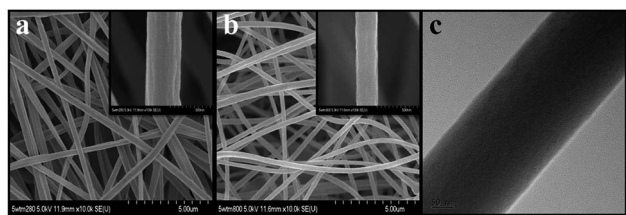


Fig. 2 Scanning electron microscopy (SEM) images of the electrospun PAN/Mel carbon nanofibers with 5% of melamine after stabilization at 280 °C (a), after carbonisation at 800 °C (b) and transmission electron microscopy (TEM) images (c) respectively.

Fine XPS scanning on different element region, is shown in Fig. 3. The C1s region at PAN/Mel carbon nanofibers without melamine (CNF800) (Fig. 3(a)) and 5% melamine (CNFM5) (Fig. 3(b)) could be deconvoluted into three peaks by peak fitting (284.6 eV, 285.83 eV, 287.77 eV for CNF800 and 284.6 eV, 286.5 eV, 288.0 eV for CNFM5), which correspond to sp^2 hybrid carbon (C–C), epoxy/hydroxyl (C–O) and sp^2 bonded carbon in N–C=N group, respectively.²⁵ Based on the XPS data, the oxygen containing functional groups are decreased by 51% while 44% increase in nitrogen containing functional groups can be observed due to addition of 5% melamine.

The N1s curve of the CNF800 sample (Fig. 3(c)) consists of two peaks: 397.9 eV (C–N=C) and 400.5 eV (N–H).²⁶ Compared with pristine CNF800, the CNFM5 also shows (Fig. 3(d)) two peaks at 398.04 eV and 400.2 eV along with two more peaks at 401.1 eV (quaternary nitrogen) and 402.80 eV (NO, oxidised nitrogen) which justifies the overall increase (56%) in amount of nitrogen which can be attributed to addition of 5% melamine during synthesis.

The electrochemical properties in the context of super-capacitor application of the carbon nanofibers synthesized from polyacrylonitrile/melamine (PAN/Mel) with 1 to 10% melamine have been investigated in two-electrode systems (Fig. 4(a)). Considering the ideal-like CV of the sample having 5% melamine (CNFM5), it has been studied in detail. Further, the specific capacitance of each electrode was calculated using

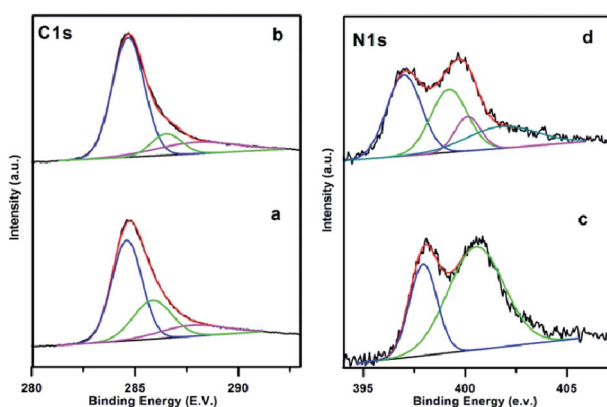


Fig. 3 XPS patterns of the electrospun PAN/Mel carbon nanofibers (a and c) without (CNF800) and (b and d) with 5% melamine (CNFM5) respectively.

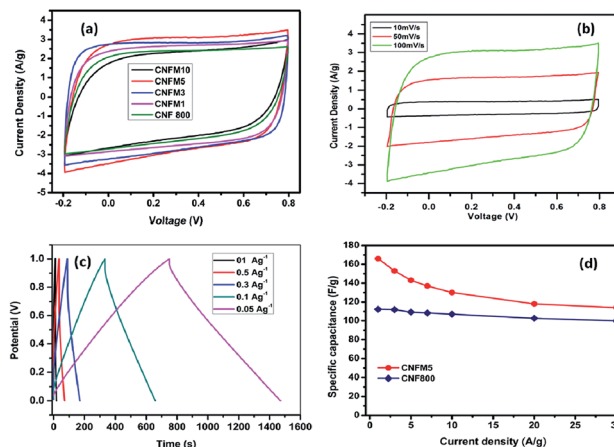


Fig. 4 (a) Cyclic Voltammetry (CV) measurements of CNF-800, CNFM-1, CNFM-3, CNFM-5, CNFM-10 at scan rate 100 mV s^{-1} in 6 M KOH and (b) Cyclic Voltammetry (CV) measurements of CNFM5 at various scan rates 10, 50, 100 mV s^{-1} in 6 M KOH (c) GCD profile of CNFM5 at different current densities. (d) Specific Capacitance Vs Current Density.

following equation; $C_g = 4(I \times \Delta t)/(m \times \Delta V)$, where, I is the discharge current, Δt is the discharge time, ΔV is the voltage change compared to the initial voltage, m is the total weight of the electrodes.

Fig. 4(b) shows the cyclic voltammograms obtained for the CNFM5 in 6 M KOH as electrolyte at various scan rates such as 10 mV s^{-1} , 50 mV s^{-1} and 100 mV s^{-1} . Loops of quasi-rectangular shape are seen without any oxidation or reduction peaks. This indicates formation of a perfect electrical double layer due to the elevated specific surface area of the material.²⁷

At different current densities ranging from 1 to 30 A g^{-1} , the calculated specific capacitance values are shown in also given in Table 1. A highest specific capacitance value of 166 F g^{-1} is obtained for CNFM5 sample at a current density of 1 A g^{-1} which shows 48% increase as compared to carbon nanofibers without melamine (CNF800). The specific capacitance decreases with increasing current density. This takes place on electrode surface due to the partial diffusion of the active ions during fast charging. At elevated current density the electrolyte cannot access all the micro pores on the electrode surface hence, the relative capacitance decreases.

The higher capacitance observed in CNFM5 sample as the amount of nitrogen containing functional moieties are highly active in basic medium such as KOH. It is well known that the nitrogen containing functional groups do not show much activity in solutions of lower pH. Hence, it approves the choice of the electrolyte. From XPS data (Fig. 3(d)), we can further calculate the different nitrogen contributions by deconvolution of the N1s peak. The amount of C–N=C and N–H type of nitrogen present at the edges of graphitic layers is 37% and 30% respectively. While the quaternary nitrogen present inside the graphitic layer and oxidised nitrogen is 15% and 16% respectively. Both these, C–N=C and N–H type of nitrogen present in an ample amount (67%) contribute to pseudo-capacitive behavior. This justifies the increase in capacitance observed



Table 1 Specific capacitance calculated at various current densities

Current density	Specific capacitance, F g ⁻¹	
	CNF800	CNFM5
1	112	166
3	112	153
5	109	143
7	108	137
10	107	130
20	103	118
30	100	114

in CNFM5 sample due to addition of melamine which contributes in this increase in N containing functional groups than that of CNF800. CNF800 can acquire N containing functional groups only from PAN while CNFM5 can get it from both PAN and melamine added during synthesis.

Fig. 5(a) shows the Nyquist plots of CNF800 and CNFM5 for two-electrode capacitors in KOH electrolyte. A 45° straight line at low frequency region corresponds to the Warburg impedance (*W*), that can be linked to the kinetic leakage processes associated with faradaic reactions during diffusion. This region is expanded for the CNFM5 sample than CNF800 which confirms a boost in the pseudo-capacitive input. The straight line nature of the *W* region in the Nyquist plot signifies the complete wettability of porous electrode surface by electrolyte ions.²⁷ EIS measurement signifies that the porous structure of the PAN/MEL fiber carbon electrode helps in quicker diffusion of electrolyte ions with decreased ionic resistance leading to almost ideal capacitive performance.

Fig. 5(b) shows Ragone plot which can be used for further analysis of the performance of PAN/MEL carbon fibers. It is generally observed that as power density increases the energy density decreases.^{28–32} In case of CNFM5 no significant decrease is observed in energy density with increasing power density unlike CNF800. This higher value of stability is due to conductivity and surface texture of the PAN/MEL carbon fibers. The micro porous structure supports the easy migration of ions to the active surface which makes the transfer of charge smoother.³³

No fading in capacitance can be seen even after 3000 cycles at a high current density of 1 A g⁻¹ (Fig. 6). This excellent capacity retention and high energy density obtained for CNFM5

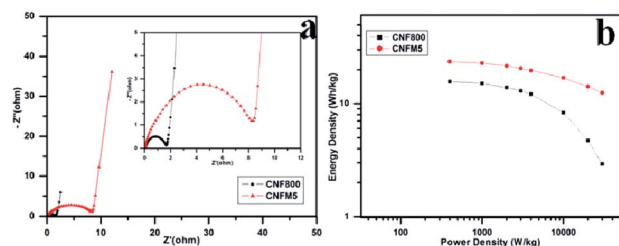


Fig. 5 (a) Nyquist plot in the range of 10 mHz to 100 kHz, (b) Ragone plot in KOH electrolyte of CNF800, CNFM5.

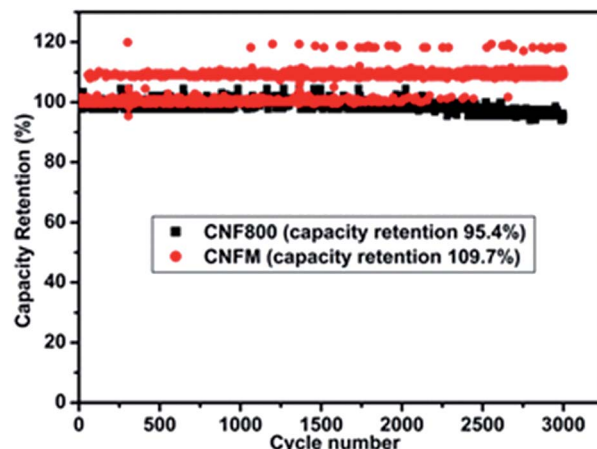


Fig. 6 Percentage retention of capacitance measured at 1 A g⁻¹ current density of CNF800 and CNFM5 sample in 6 M KOH electrolyte.

further encouraged us to check its applicability as electrode material for Li-ion battery.

The self-standing, flexible carbon fibre tape of CNFM5 was used as working electrode in Li-ion battery (2032-type coin cell) and its electrochemical performance was studied. Fig. 7(a) shows the CV curves of CNFM5 electrode at first three cycles. The electrochemical reaction kinetics of Li-ion insertion and de-insertion can be studied from the shape of the curves obtained. In the first cycle, a small oxidation peak can be seen at ~0.5 V due to the insertion of Li-ion into pores between carbon nanofibre network.³⁴ The irreversible reduction peak can be seen at ~0.3 V due to decomposition of electrolyte. The reduction peak due to formation of a solid electrolyte interface (SEI) layer can be seen at ~0.3 V of 2nd cycle.³⁵ The shapes of CV curves obtained at 2nd and 3rd cycles are almost coincidental which confirms the CNFM5 electrode demonstrates a stable Li-ion insertion and de-insertion mechanism.

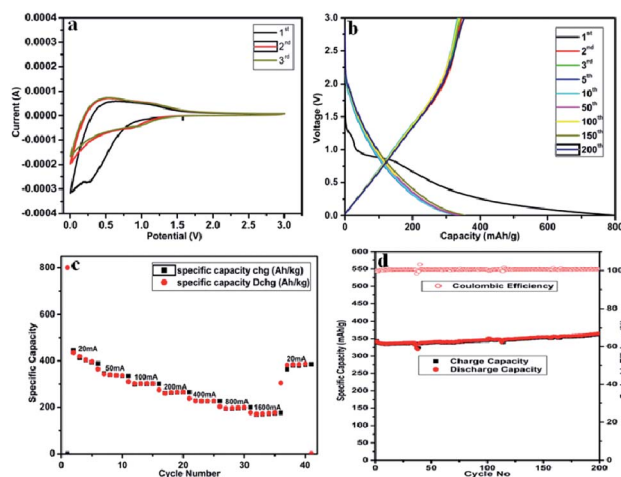


Fig. 7 (a) Electrochemical performance of Li-ion battery CV curves of between 0.01–3.0 V first three cycles (b) charge/discharge curves of CNFM5 at current density 100 mA g⁻¹ (c) specific capacities of CNFM5 at different current rates, (d) cyclic performance and corresponding coulombic efficiencies of CNFM5 at current density of 100 mA g⁻¹.



Fig. 7(b) shows the charge–discharge curves of the self-standing, flexible carbon fibre tape of CNFM-5 for the 1st, 2nd, 3rd, 5th, 10th, 50th, 100th, 150th, 200th cycles at current density 100 mA g^{−1} within potential window of 0.01–3.0 V. The first discharge curve show a sharp decrease followed by a plateau around 0.8 V, which is mainly due to rapid generation of SEI layer due to decomposition of electrolyte. CNFM5 shows a superior initial discharge capacity of 800 mA h g^{−1} which is attributed to the surface microstructure and graphitic upper surface layer which gives ease of insertion and de-insertion to Li⁺ ion.³⁶ The sloping capacity can be due to lowering of inter-layer spacing and the interactions between lithium and heteroatoms. The elevated irreversible capacity between first discharge and charge may be due to the fast generation of SEI film. No noticeable plateau is observed for further cycles which indicate that the sites of Li⁺-ions into the CNFM5 are electronically as well as geometrically discrete.^{37,38} Similar behaviour is observed in other CNFs synthesized by electrospinning using other starting materials.^{39,40} The electrode made from self-standing, flexible carbon fibre tape of CNFM5 shows 365 mA h g^{−1} after 200 cycles which confirms stable Li-ion insertion/de-insertion mechanism which leads to excellent repeatability.

Fig. 7(c) shows, the comparison of rate performance of CNFM5 electrode at different current densities from 20 to 1800 mA g^{−1} (Table 2). When the current density revisits 20 mA g^{−1}, the specific capacity observed as 387.32 mA h g^{−1} with stable cycling. Fig. 7(d) shows, the long cycling performance of CNFM5 investigated at current density 100 mA g^{−1}. It is clearly seen that the CNFM5 electrode maintains a stable discharge capacity of 365 mA h g^{−1} after 200 cycles. For the first 10 cycles a small decrease in the initial capacity of CNFM5 electrode can be observed, which is due to SEI layer formation. But after 10 cycles the capacity increases with nearly 100% coulombic efficiency. This remarkable increase in capacitance may be due to the additional specific capacity provided by a viscous layer formed on the surface of the CNFM5 electrode or a fresh SEI layer grown over uneven texture of the original one.^{41,42}

Fig. S1† shows the cyclic performance of pristine carbon nanofibre, which shows initial discharge capacity around 292 mA h g^{−1} at a current rate of 100 mA g^{−1}, which reduces to 285 at the end of 20th cycle. Herein, there is a drop in capacity

with cycling. However, nitrogen doped carbon fibre sample CNFM5 shows increase in capacity with cycling as discussed above.

Fig. 8 shows the Nyquist plots obtained by electrochemical impedance spectrometry of CNFM5 electrode after 200 cycles in charged and discharged states of the cell. It is observed to possess a composition of two depressed semicircles in high frequency region (shown in inset for clarity) followed by a polarization curve in lower frequency side. The obtained plots were fitted to the equivalent circuit given in inset. Here, R_s is the ohmic internal resistance of the cell. C_f and R_f are the capacitance and resistance of the passive film forming at the lithium anode and C_{dl} represents the double layer capacitance. In this model, the C_{dl} and CPE are combined into one CPE, labeled as C_{dl} . R_{ct} is the charge transfer resistance and W_c is a diffusional impedance.⁴³ The simulated data is shown as continuous lines overlayed with the original impedance data in Fig. 8. Also, values of fitting parameters of the equivalent circuit are given in Table 3. It may be noted that there is not much significant variation in cell resistance, while a significant difference in passivation film resistance and charge transfer resistance are encountered. The Warburg coefficient is found to be almost doubled indicating a resistance to diffusion of Li⁺ ions after discharge.

To find out the reason behind the excellent cycling performance of CNFM5 electrode and verify the morphological changes in the structure after 200 cycles, we disassemble the cell to carry out its FE-SEM imaging. Fig. 9(a–c) shows the FE-SEM images of CNFM5 electrode after 200 cycles at current density of 100 mA g^{−1}. When compared to previous images (Fig. 2(b)) it can be observed that there is not much change in shape, size and fibrous geometry which indicates the structural stability of CNFM5 sample. The surface of the fibres appears slightly rough due to the SEI layer which acts as a protective film to improve the performance of electrode but other than that no cracks or fractures were observed. Stable electrochemical

Table 2 Specific capacity of CNFM5 at different current densities

Current density, mA g ^{−1}	Specific capacity of CNFM5, mA h g ^{−1}
20	435.47
50	364.53
100	310.59
200	275.24
400	265.77
800	203.01
1600	176.82
20	387.32

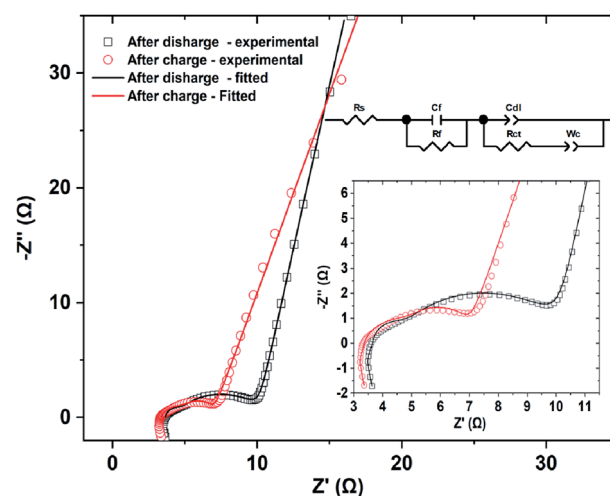


Fig. 8 Nyquist plots obtained by electrochemical impedance spectrometry of CNFM5 electrode after 200 cycles in charged and discharged states of the cell and equivalent circuit.



Table 3 Cell resistance, R_s , passivation film resistance, R_f , charge transfer resistance, R_{ct} and Warburg coefficient, W_c obtained by fitting of impedance plots to the equivalent circuit mentioned in Fig. 8

Element	Value after charge	Value after discharge
R_s (Ω)	3.64	3.36
R_f (Ω)	0.76	2.05
R_{ct} (Ω)	6.05	1.65
W_c ($\Omega^{-1} s^{-1}$)	28.66	52.24

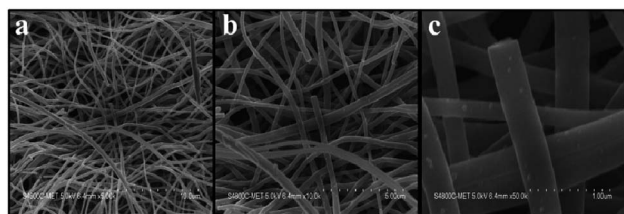


Fig. 9 FE-SEM images of CNFM5 electrode after 200 galvanostatic charge-discharge cycles at a current density of 100 mA g^{-1} .

performance of CNFM5 confirms its superior potential for Li-ion battery application. In both the energy devices, we found that nitrogen-doped carbon fibres showed excellent electrochemical performance as compared to their bare carbon nanofibres. It is reported that, incorporation of nitrogen in carbon accelerates the surface-dominated charge-storage.⁴⁴ Generally, any doping in the pristine material creates defects and same phenomenon is happening N-doped carbon, which increases the number of active sites for Li^+ storage. A stronger interaction between ions occurs due to higher electronegativity possessed by N-doping in carbon.^{45,46} Recently, Chen *et al.* have reported that N-doped hard carbon microspheres exhibited high-rate capability due to surface-driven charge storage for potassium ion batteries.⁴⁷ Also, N-doping influences the fast ion transport to the surface. The different nitrogen percent incorporation shows change in morphology as well as capacitance and specific capacity for Li-ion battery application. However, moderate nitrogen incorporation gives good electrochemical performance. It indicates that adequate N-doping induces desired surface modification of the carbon, *i.e.* nitrogen doping modifies the surface of the material which includes vacancy defects, as well as morphology which helps in achieving higher electronegativity for increased interaction between ions as discussed above.

4 Conclusions

In summary, novel PAN-based nitrogen enriched carbon fibers were synthesized by electrospinning with addition of various concentrations of melamine. The obtained PAN-MEL fibers with 5% melamine stabilised at 280°C and carbonized at 800°C in nitrogen atmosphere showed excellent electrochemical performance with a specific capacitance of up to 166 F g^{-1} at current density 1 A g^{-1} using 6 M KOH electrolyte and capacity retention

of 109.7% after 3000 cycles. This increase in capacitance observed in CNFM5 sample was found due to addition of melamine which contributes in increase in 'N' containing functional groups than that of CNF800. Two electrode systems of CNFM5 sample showed high energy densities of 23.72 to $12.50 \text{ W h kg}^{-1}$ and power densities from 400 to $30\,000 \text{ W kg}^{-1}$. CNFM5 sample used as an anode for Li-ion battery application showed a high specific capacity up to $435.47 \text{ mA h g}^{-1}$ at 20 mA g^{-1} , good rate capacity and excellent cycling performance (365 mA h g^{-1} specific capacity even after 200 cycles at 100 mA g^{-1}).

Conflicts of interest

There are no conflicts to declare.

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