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# PAPER

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# Unravelling the role of temperature in a redox supercapacitor composed of multifarious nanoporous carbon@hydroquinone†

Aditi Barua and Amit Paul 💷 \*

The intermittency of renewable energy sources has led to the invention of supercapacitors. The variation of temperature impacts their working capability particularly in the regions which are too hot or too cold. Herein, the effect of temperature on the double layer formation and the redox mechanism of hydroquinone adsorbed on multifarious nanoporous carbon (MNC) have been reported. The studies have been carried out in 2 M H<sub>2</sub>SO<sub>4</sub> electrolyte solution within a temperature range of -10 to 50 °C. The maximum specific capacitance of the composite material drops from 319 F g<sup>-1</sup> at 50 °C to 213 F g<sup>-1</sup> at -10 °C. An equivalent circuit model has been chosen to fit the EIS spectra at the double layer potential and formal potential. Subsequently, an Arrhenius type plot has been constructed to calculate the activation energy of the system which revealed 8.69 and 7.77 kJ mol<sup>-1</sup> activation energy at the formal potential and double layer potential respectively. The composite also shows excellent cyclability even at enhanced temperatures, which is a major requirement for the application of these supercapacitors in vehicles and other electrical equipment.

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# Introduction

It is a truth universally acknowledged that the ever growing population has accelerated the rate of consumption of fossil fuels due to the increasing exigency for energy. Focus has been placed on alternate sources of energy such as geothermal, sunlight, wind etc. However, the diurnal nature and seasonality of solar and wind power, along with the constraint of varying geography lead to intermittency of such renewable energy sources. Hence, to cope with the increasing demand for storing energy, research on electrochemical energy storage devices has gained momentum in the current era. As early as 1957, engineers at General Electric first began conducting experiments with activated charcoal for charge storage. Next, researchers at Standard Oil of Ohio (SOHIO) in 1966 accidentally came across the phenomenon of double layer capacitor while working on fuel cells. Yet it was not commercialized by them, and was instead licensed to NEC Corp. of Tokyo, who finally marketed this device as a 'supercapacitor' to be used in memory back-up power.1 Commercial supercapacitors are mainly of the electrical double layer type, called as electrical double layer capacitors (EDLCs) which employ the phenomenon of double layer formation at the electrode/electrolyte interface.<sup>2-4</sup> Combined with this, pseudocapacitors can boost the charge storage ability

utility in emergency shut-down support, ultrabatteries, voltage stabilizer, medical defibrillators, aeronautics, military equipments, hybrid electric vehicles, portable electronic devices to name a few.1,7-11 The key difference between batteries and supercapacitors lies in the fact that supercapacitors have a higher power density than that of batteries even though their energy density is much lower as compared to the same.12,13 The performance of supercapacitors is also influenced by the type of electrolyte used in the device.14,15 Ionic liquids and organic electrolytes are generally implemented because of their wider potential stability windows. Nonetheless, it has already been conceded that aqueous electrolytes have certain benefits in terms of their environmental friendliness, economic values, higher electrical conductivity etc.14 Ergo, understanding the role of the electrolyte and its impact on the electrode kinetics are of utmost significance. Since these devices majorly find use in electrical equipments, their performance in a high temperature environment caused by heating up of the equipment needs evaluation. Besides, due to varying weather conditions pertaining to different regions, the broad temperature effects of supercapacitors need further research. It is widely known that temperature plays a pivotal role in determining the stability of electrolytes, and working of the supercapacitors in general. Consequently, it is important to assess their functionality under harsh temperature conditions.

by incorporating redox reactions.5,6 Mostly, these devices find

The versatility of carbon based materials has rendered them a noteworthy status in the field of science and technology. Starting from batteries, supercapacitors,  $H_2$  storage to inks and

Department of Chemistry, Indian Institute of Science Education and Research (IISER), Bhopal, MP, 462066, India. E-mail: apaul@iiserb.ac.in

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paints, pencil tips, metal smelting etc., the usage of carbon is prodigious.<sup>16-20</sup> In supercapacitor technology particularly, carbon based materials are prominent as their high surface area leads to substantial charge storage properties in the form of a double layer.<sup>21-23</sup> Their performance can be further augmented by activation with oxidants such as NaOH, KOH, H<sub>3</sub>PO<sub>4</sub> etc.<sup>24-26</sup> KOH activation of a non-porous carbon led to a whopping 250 times increase in the specific capacitance as reported by Singh et al.26 The next obvious step of research on carbon based materials as supercapacitors would be the impact of temperature variation on their working principle. Literature reports are aplenty on the thermal characteristics of materials such as activated carbon,<sup>27-30</sup> onion-like carbons,<sup>31</sup> carbon nanotubes,30-32 graphene,33-37 and metal oxides like hydrous RuO2,38 MnO<sub>2</sub>,<sup>39-43</sup> α-Fe<sub>2</sub>O<sub>3</sub>,<sup>44</sup> La<sub>2</sub>O<sub>3</sub> nanosheet decorated reduced graphene oxide45 etc. Masarapu et al. reported the effect of temperature on the kinetics and diffusion mechanism of SWCNTs in an organic electrolyte and showed outstanding stability of the material after repeated heating and cooling followed by an additional 224 000 charge-discharge cycles.32 Iwama et al. investigated the characterizations of a Batscap SC for low temperature applications. Electrochemical impedance spectroscopic (EIS) studies in the above cases also revealed the change in various electrode processes upon systematic increase or decrease in temperature. Typically, such EIS studies are mostly carried out for double layer formation, and literature reports on the effect of temperature on the redox properties of

the active material are limited.<sup>30,32,46</sup> Additionally, since temperature dependence investigations are mostly performed in organic electrolytes and ILs, detailed reports on the workings of a supercapacitor in aqueous electrolytes under temperature variations are minimal.<sup>29,40</sup>

Further increase in the charge storage properties of supercapacitors can be brought about by the inclusion of redox active materials. Meyer and co-workers emphasized the effect of oxide sites on a glassy carbon working electrode to transform the irreversibility of catechol into reversibility.47 Singh et al. reported a maximum specific capacitance of 200 F  $g^{-1}$  in activated charcoal with hydroquinone physisorbed on its surface.48 In this work, we have examined the consequences of temperature alteration on a multifarious nanoporous carbon (MNC) with surface-adsorbed hydroguinone (H<sub>2</sub>Q) acting as the redox probe wherein H<sub>2</sub>SO<sub>4</sub> has been used as electrolyte. The ease of synthesis of the composite MNC-H<sub>2</sub>Q along with the relatively economical nature of the starting materials and their ecologically green genre make them appealing for future commercialization. Cyclic voltammetry (CV), electrochemical impedance spectroscopy (EIS), galvanostatic charge/discharge (GCD) experiments were employed extensively to analyse the performance of the supercapacitor at various temperatures. Particularly, EIS studies have been performed at both the double layer and at the formal potential to account for their differing responses under temperature fluctuations.



Fig. 1 Schematic representation of the preparation of MNC from micro/mesoporous carbon (MMC) and subsequent formation of MNC-H<sub>2</sub>Q from MNC.

# **Experimental section**

## Chemicals

Micro/mesoporous carbon (100 mesh, MMC), potassium hydroxide (KOH), hydroquinone ( $H_2Q$ ), *N*-methyl pyrrolidone (NMP), poly(vinylidene fluoride) (PVDF) have been purchased from Sigma-Aldrich. Concentrated sulphuric acid has been purchased from Merck Emplura. Acetylene black, platinum (Pt) foil and Pt wire electrodes were purchased from Alfa Aesar. A saturated calomel electrode (SCE) was purchased from CH instruments.

## KOH activation of MMC

Following a previous report,<sup>26</sup> micro/mesoporous carbon (MMC) was activated using KOH as the oxidant. 1 g of MMC and 5 g of KOH were ground together using a mortar and pestle and then transferred to an alumina crucible. The crucible was then inserted into a temperature controlled tube furnace, followed by carbonization under continuous N<sub>2</sub> flow (Fig. 1). The sample was heated to a temperature of 750 °C at a ramp rate of 5 °C min<sup>-1</sup> and subsequently held at the same temperature for 1 h. It was allowed to cool naturally under N<sub>2</sub> flow and then washed thoroughly with 5 N HCl to eliminate different mineral contents, followed by water, which in turn eliminated the chloride ions. The KOH activated MMC will henceforth be termed as MNC (Multifarious Nanoporous Carbon) in the rest of the discussion.

#### Preparation of active material (MNC-H<sub>2</sub>Q)

Previous literature report from our group has been followed to prepare the active material.<sup>48</sup> Briefly, 500 mg of MNC and 275 mg of hydroquinone (H<sub>2</sub>Q) were shaken in 50 mL distilled water in a conical flask at a temperature of 50 °C for 24 h at a rate of 90 rpm (Fig. 1). It was then filtered and dried in a hot air oven at 60 °C for 24 h. This active material has been termed as MNC-H<sub>2</sub>Q and the explanation behind the nomenclature has been provided in results and discussion.

#### Characterizations

**Powder X-ray diffraction (PXRD).** Powder X-ray diffraction (PXRD) patterns were recorded by a Bruker AXS D8 Advance with Cu K $\alpha$  radiation (1.54 Å) with a step size of 0.02° in a 2 $\theta$  range of 0–80°.

**Brunauer–Emmett–Teller (BET).** N<sub>2</sub> sorption isotherms were performed using a Quantachrome Autosorb QUA211011 equipment. The sample was degassed using high vacuum at 60 °C for 20 h prior to gas sorption studies.

**Scanning electron microscopy (SEM).** The samples were dried at first and then spread over carbon tape, followed by gold coating for 120 s. SEM experiments were performed using a Carl ZEISS (ultraplus) FE-SEM at a working voltage of 20 kV.

**Transmission electron microscopy (TEM).** TEM (FEI TALOS 200S) experiments were carried out under an accelerating voltage of 200 kV. Samples were prepared by drop casting a suspension of powdered sample in isopropanol on a carbon-coated copper grid and dried at 60  $^{\circ}$ C overnight.

Thermogravimetric analysis (TGA). TGA was performed in a PerkinElmer TGA 4000 instrument between temperature ranges of 30 to  $350 \,^{\circ}$ C at a scan rate of  $5 \,^{\circ}$ C min<sup>-1</sup> with a N<sub>2</sub> flow rate of 20 mL min<sup>-1</sup>.

**Differential scanning calorimetry (DSC).** DSC was conducted in a PerkinElmer DSC 6000. MNC-H<sub>2</sub>Q samples were enclosed in an aluminium pen and temperature was varied between 30 to 350 °C at a scan rate of 5 °C min<sup>-1</sup> and N<sub>2</sub> flow rate of 20 mL min<sup>-1</sup>.

## **Electrochemical measurements**

Electrochemical experiments were conducted using a BioLogic potentiostat (SP-240). Temperature dependent studies were performed in an ESPEC SH-222 temperature and humidity chamber. Electrochemical experiments were performed at -10, 0, 10, 20,30, 40 and 50 °C. For the three electrode supercapacitor studies, a saturated calomel electrode (SCE), active material coated Pt foil electrode, and Pt wire were used as reference electrode, working electrode, and counter electrode respectively. The active material, acetylene black and PVDF were taken in a ratio of 10:2:2 and then added to 1 mL NMP solution. It was then stirred for 6 h to ensure proper homogenization. 100 µL of this solution was then drop-casted onto cleaned Pt foil working electrodes, covering 1 cm<sup>2</sup> area, and subsequently dried in a hot air oven at 100 °C for 14 h. Mass of material deposited on each electrode was 1.2 mg cm<sup>-2</sup>. Cyclic voltammetry (CV) studies were carried out in a potential range of -0.15 to +0.75 V vs. SCE at scan rates of 1, 5, 10, 20, 50 and 100 mV s<sup>-1</sup> in 2 M H<sub>2</sub>SO<sub>4</sub>. Galvanostatic charge/ discharge (GCD) studies were performed in the same potential window at current densities of 1, 1.5, 2, 5 and 10 A  $g^{-1}$ . Electrochemical impedance spectroscopy (EIS) experiments were performed at 0 V vs. SCE and at the formal potentials determined at different temperatures over a frequency range of  $10^{-2}$  to  $10^{5}$  Hz. The specific capacitance  $(C_{sp})$  values from CVs were calculated from eqn (1) and that from GCD were determined from eqn (2),

$$C_{\rm sp} = \frac{\int I dE}{2mE\nu} \tag{1}$$

$$C_{\rm sp} = \frac{It}{2mE} \tag{2}$$

wherein, *I*, *m*, *E*,  $\nu$ , and *t* denote the current, mass of deposited material, potential window, scan rate and time taken for the charge/discharge process respectively. The inverse of the maximum frequency  $f_0$ , known as the dielectric relaxation time constant,  $\tau_0$  has been calculated from the imaginary capacitance,  $C''(\omega)$  vs. frequency plot from eqn (3).<sup>49</sup>

$$C''(\omega) = \frac{-Z'(\omega)}{\omega \left| Z(\omega)^2 \right|} \tag{3}$$

wherein,  $Z'(\omega)$  is the real component of impedance,  $Z(\omega)$  is the total impedance and  $\omega$  is the frequency.

## **Results and discussions**

Powder X-ray diffraction (PXRD) was performed for MNC and MNC-H<sub>2</sub>Q. Peaks at  $\sim 24^{\circ}$  and  $44^{\circ}$  were observed for both the

samples, characteristic of (002) and (100) diffractions of graphitic carbon (Fig. S1, ESI<sup>†</sup>).<sup>19</sup> Brunauer-Emmett-Teller (BET) N<sub>2</sub> adsorption/desorption analysis of MNC-H<sub>2</sub>Q has been depicted in Fig. 2a. The BET of MNC-H<sub>2</sub>Q exhibited mixed Type-I and Type-IV isotherm profile with a hysteresis loop of H4 type according to IUPAC classification. The surface area of MNC is 3090 m<sup>2</sup> g<sup>-1</sup> having pores centred at 0.56, 1.3, 2.6, 3.5 and 5.2 nm and with a total pore volume of 2.3 cm<sup>3</sup> g<sup>-1</sup> (Fig. S2, ESI<sup>†</sup>).<sup>26</sup> Since, the material had a range of nanopores, hence it has been termed as multifarious nanoporous carbon (MNC) in this work. After adsorption of H<sub>2</sub>Q, the surface area of the resultant MNC-H<sub>2</sub>Q declined to 610 m<sup>2</sup> g<sup>-1</sup> (Fig. 2a) and it has pores centred at 0.62, 1.4, 2.6, 3.8 and 7.4 nm respectively which was determined using non-local density functional theory (NLDFT) (Fig. 2b). This reduction in surface area is due to incorporation of H<sub>2</sub>Q inside the pores of MNC, which is further supported by a reduction in the total pore volume to 0.58 cm<sup>3</sup>  $g^{-1}$  (Fig. 2c). The amount of H<sub>2</sub>Q adsorbed on MNC-H<sub>2</sub>Q was determined from thermo-gravimetric analysis (TGA) which revealed a 12% weight loss in the temperature range of 130 to 250 °C (Fig. S3, ESI<sup>†</sup>). Since, the melting point of H<sub>2</sub>Q is 172 °C, the 12% weight loss can be attributed to the desorption of H<sub>2</sub>Q from MNC-H<sub>2</sub>Q. The weight loss between 50 and 130 °C is presumably due to the evaporation of water molecules present inside the pores, which has been previously observed also.48 It is discernible from BET analysis that the pores in MNC-H<sub>2</sub>Q have expanded slightly in accordance with the structural adjustment that the system had to undergo in order to make the

hydroquinone adsorption process entropically favourable, thereby gaining thermodynamic feasibility according to previous literature report from our group (vide infra).48 This is also evident from the SEM images in Fig. 3a and b wherein it can be clearly noticed that the adsorption of H<sub>2</sub>Q in MNC-H<sub>2</sub>Q resulted in morphological expansion compared to the precursor material (MNC). Differential scanning calorimetry (DSC) reveals that H<sub>2</sub>Q desorbed from MNC-H<sub>2</sub>Q at a temperature of 211 °C (Fig. S4, ESI<sup>†</sup>). Due to strong H-bonds between H<sub>2</sub>Q and MNC and presence of ultramicropores/micropores, the desorption temperature for H<sub>2</sub>Q rose from 172 to 211 °C. The enthalpy change ( $\Delta H$ ) for this melting process was (-)7.53 kJ mol<sup>-1</sup> which implies exothermicity of the process. Therefore, the process of adsorption must be endothermic in nature. Hence, for the spontaneity of the adsorption process, *i.e.* to achieve a negative Gibbs free energy ( $\Delta G$ ), the change in entropy ( $\Delta S$ ) must be positive. The morphological changes after the adsorption of H<sub>2</sub>Q can be directly related to this increase in entropy, which was caused by the enhanced randomness at the adsorbent/adsorbate interface.50

Fig. 3c and d compare the high resolution transmission electron microscopy (HR-TEM) micrographs of MNC before and after physisorption of  $H_2Q$  respectively. Fig. 3c reveals the highly amorphous texture and disordered nature of MNC with a porous morphology that is consistent with the results from BET. The porosity of the successor material was maintained after the adsorption of  $H_2Q$  (Fig. 3d) and it is apparent that the pores in MNC- $H_2Q$  are somewhat larger than that of MNC



Fig. 2 Characterizations of MNC-H<sub>2</sub>Q using BET. (a)  $N_2$  adsorption/desorption isotherms, (b) pore size distribution and (c) cumulative pore volume.



Fig. 3 Morphological characterizations of MNC and MNC-H<sub>2</sub>Q. SEM and HR-TEM images of MNC are shown in (a and c) and of MNC-H<sub>2</sub>Q in (b and d) respectively.

which contributed to the decrease in surface area of MNC after inclusion of  $H_2Q$  which is again consonant with the BET results.

## Electrochemical results of MNC-H<sub>2</sub>Q

All the electrochemical experiments were carried out in 2 M  $H_2SO_4$  electrolyte solution. The cyclic voltammograms (CVs) of



Fig. 4 Proton-coupled electron transfer mechanism for MNC-H<sub>2</sub>Q.

Table 1 Specific capacitances of MNC-H<sub>2</sub>Q at different scan rates and different temperatures

Scan rate (mV $s^{-1}$ )	Specific capacitance (F $g^{-1}$ )								
	$-10\ ^\circ C$	0 °C	$10\ ^{\circ}\mathrm{C}$	20 °C	30 °C	$40~^{\circ}C$	50 °C		
100	191	197	208	222	226	229	234		
50	195	202	211	226	233	234	242		
20	200	206	216	232	240	242	253		
10	204	209	218	238	244	247	260		
5	206	212	221	243	250	255	268		
1	213	222	241	259	269	280	319		

MNC-H<sub>2</sub>Q were rectangular in shape with a sharp redox peak at  $\sim$ 0.46 V vs. SCE which is ascribed to the redox reaction between benzoquinone (BO) and hydroquinone (H<sub>2</sub>O) at all temperatures (Fig. 5a). We have previously reported that H-bonds formation takes place between the oxide sites of nanoporous carbon and H<sub>2</sub>Q (Fig. 4).<sup>48</sup> The H-bond between the oxide groups of MNC and H<sub>2</sub>Q leads to a proton shuttling mechanism wherein the  $H^+$  moves between the –OH functionalities of  $H_2Q$ and the carbonyl oxygen of MNC and simultaneously two electron transfers occur between the electrode and MNC-H<sub>2</sub>Q (Fig. 4).

This H-bonding, in turn, also imparts stability to the H<sub>2</sub>Q/BQ system (vide infra). The specific capacitance of the composite material at 1 mV s<sup>-1</sup> was determined to be 259 F g<sup>-1</sup> at 20  $^{\circ}$ C (Table 1). This value nearly remains unaltered even after incorporation of H<sub>2</sub>Q in the system, (Fig. S5 and Table S1, ESI<sup>†</sup>) since H<sub>2</sub>Q blocks a large number of pores of MNC which presumably reduces double layer charging. Nevertheless, because of the inclusion of pseudocapacitance which resulted

in a sharp peak in the CV, it was possible to probe the temperature dependence of redox reactions which is the main objective of this study.

The influence of temperature on the performance of the supercapacitor was examined within a range of -10 to 50 °C. From the CV curves collected at  $1 \text{ mV s}^{-1}$  scan rate in Fig. 5a, the increase in the integral area with increasing temperature is apparent. Table 1 reveals that the specific capacitances of MNC-H<sub>2</sub>Q increases with increasing temperature at all scan rates since the ionic mobility of the electrolyte ions increases with rise in temperature, resulting in faster movement of ions towards the electrode surface for the formation of the electrical double layer. Indeed, the highest specific capacitance at 1 mV  $\rm s^{-1}$  at 50 °C was 319 F  $\rm g^{-1}$  which decreased to 213 F  $\rm g^{-1}$  at -10 °C (Table 1). Thus, there was a 33% drop in the specific capacitance at the lowest scan rate with decreasing operational temperature (Fig. 5c and Table 1). However, the specific capacitance drop at a higher scan rate of 100 mV s<sup>-1</sup> was only 18% (Fig. 5c, Table 1 and Fig. S6h, ESI†). This is due to the fact that at a faster time scale, only a limited number of ions will be able to contribute to the formation of double layer and subsequent faradaic reaction and hence the temperature variation will affect the kinetics to a much smaller extent at a faster scan rate as compared to that at a slower scan rate.

Galvanostatic charge/discharge curves at 1 A  $g^{-1}$  for MNC-H<sub>2</sub>Q have been shown in Fig. 5b and it also reveals that specific capacitance increases with increasing temperature. The discharging time of the supercapacitor was longer than the charging time at low current density (1 A  $g^{-1}$ ), *i.e.* coulombic efficiency (C.E.) was greater than one, which was presumably due to slight onset of hydrogen evolution reaction (HER) occurred at the lowest potential window (between -0.1 to



Fig. 5 Electrochemical characterizations of MNC-H<sub>2</sub>Q in 2 M H<sub>2</sub>SO<sub>4</sub> electrolyte solution. (a) Overlay of CVs at 1 mV s<sup>-1</sup> in the temperature range of -10 to 50 °C. (b) GCD overlay at 1 A g<sup>-1</sup> in the temperature range of -10 to 50 °C. (c) Specific capacitance vs. scan rate in the same temperature range. Long term cyclic tests of the system at (d) 20 °C for 10 000 cycles, and at (e) 50 °C for 5000 cycles.

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-0.15 V vs. SCE). In our experimental set-up, part of the Pt working electrode was exposed in the electrolyte solution and it is well known that Pt catalyses HER. The long term stability of the active material was evaluated by performing several CV cycles. It showed remarkable specific capacitance retention of 92% after 10 000 cycles at 20 °C (Fig. 5d). It must also be noted that even at a high temperature of 50 °C, 86% retention was observed after 5000 cycles (Fig. 5e). The decrease in cyclic stability of a supercapacitor at higher temperatures has been

observed previously.<sup>41,51</sup> The decrease in cyclic stability at higher temperature can be attributed to the enhancement of decomposition of  $H_2Q$  moieties during electrochemical cycling. Fig. S6, ESI† revealed that at room temperature, the CVs remained similar after 1000 cycles, however at 50 °C, a significant decay in the redox peak was observed and thus suggesting degradation of  $H_2Q$  moieties during electrochemical cycling.

The role of temperature in influencing the various electrochemical processes at the electrode/electrolyte interface has



Fig. 6 Electrochemical impedance spectroscopic (EIS) studies of  $MNC-H_2Q$ . Overlay of Nyquist plot of  $MNC-H_2Q$  at (a) formal potential and (b) double layer potential at all temperatures. Overlay of Nyquist plots at high frequency at (c) formal potential and (d) at double layer potential at all temperatures. Imaginary capacitance vs. frequency overlay plots at (e) formal potential and (f) double layer potential at -10, 20 and 50 °C.

been analysed further by electrochemical impedance spectroscopy (EIS). These EIS studies were carried out at a double layer potential, *i.e.*, 0 V vs. SCE and at the formal potential  $(E^{0'})$  of the H<sub>2</sub>Q/BQ couple at different temperatures (Fig. 6a-d). The equivalent circuit chosen to analyse the EIS data has been provided in Fig. 7.<sup>52</sup> In this circuit,  $R_s$  denotes the solution resistance or the equivalent series resistance and the current has to pass through this part of the circuit at all frequencies. The inte a c mo  $Q_{d1}$  utilizing eqn (4) wherein  $\omega$  and "a1" represent frequency corresponding to the maxima on the imaginary component of the impedance in the high frequency region and exponent of  $Q_{dl}$ respectively.

$$C_{\rm dl} = Q_{\rm dl} \times (\omega)^{a_1 - 1} \tag{4}$$

CPE behaves as a pure capacitor when its exponent, "a" equals to 1, and behaves like a pure resistor when "a" is 0. An exponent value of 0.5 signifies the Warburg element, characterized by diffusion limitation and all other values correspond to intermediate behaviour.<sup>54</sup> At higher frequencies, the current will pass through  $Q_{dl}$ . However, as the frequency decreases, impedance due to Q<sub>dl</sub> increases rapidly and hence does the current flow through  $R_{ct}$ , *i.e.*, the charge transfer resistance at the electrode/electrolyte interface and Q<sub>ms</sub>, *i.e.* the diffusion capacitance arising from mass transport of electrolyte ions inside the nanopores of the nanomaterial.  $R_{ads}$  represents the resistance due the adsorption of the electrolyte ions inside the pores of the active material. As the frequency further decreases, impedance through  $Q_{\rm ms}$  increases and hence the current passes through  $R_{ads}$ , and finally through  $Q_{ps}$ , the pseudocapacitive charging originating from the movement of electrons inside the material.

In the high frequency region, the first intersection on the real axis is the solution resistance  $R_s$  (Fig. 6c and d). The  $R_s$  values ble layer potential and formal potential e is not dependent on electrode potendl is the double layer capacitance just at e interface at a very fast time scale and it is interesting to note that a comparison of C<sub>dl</sub> values at the double layer potential and formal potential revealed a higher

 $Q_{\rm ps}$  (F s<sup>(a3-1)</sup>)

0.085

0.109

0.889

0.100

0.092

0.107

0.108

 $\tau_0$  (s)

1.2

1.1

0.82

0.80

0.76

0.70

0.60

аЗ

0.970

0.972

0.968

0.956

0.969

0.962

0.955

e double layer capacitance just at the electrode/electrolyte	are nearly same at dou
erface at a very fast time scale has been represented by	since solution resistant
onstant phase element (CPE, $Q_{dl}$ ) because of surface inho-	tial (Tables 2 and 3). Co
geneity of the electrode. ${}^{52,53}C_{dl}$ values can be calculated from	the electrode/electrolyte

Table 2 Equivalent circuit parameters of MNC-H<sub>2</sub>Q at double layer potential (0 V vs. SCE)

 $R_{\rm ct}(\Omega)$ 

0.476

0.386

0.258

0.228

0.211

0.179

0.170

 $R_{\mathrm{ads}}\left(\Omega\right)$ 

0.132

0.123

0.115

0.106

0.093

0.088

0.082

 $Q_{\rm ms}$  (F s<sup>(a2-1)</sup>)

2.885

1.770

1.435

0.901

0.875

0.695

0.162

 $a^2$ 

0.456

0.501

0.554

0.582

0.675

0.702

0.753

Table 3 Equivalent circuit parameters of MNC-H<sub>2</sub>Q at formal potentials

 $C_{dl}$  (mF)

3.17

3.33

3.57

3.91

4.07

4.44

5.05

Temperature (°C)	$R_{\rm s}(\Omega)$	$C_{\rm dl}$ (mF)	$R_{\rm ct}(\Omega)$	$R_{\rm ads}(\Omega)$	$Q_{\rm ms} ({\rm F}{\rm s}^{(a2-1)})$	a2	$Q_{\rm ps} ({\rm F}  {\rm s}^{(a3-1)})$	a3	$\tau_0$ (s)
	5()			uus ( )	(1115)( )		(ps ( )		
-10	1.47	0.70	0.262	0.381	2.57	0.308	0.414	0.958	5.8
0	1.27	1.11	0.254	0.372	1.66	0.315	0.309	0.957	4.1
10	1.01	1.32	0.219	0.331	1.43	0.345	0.414	0.959	4.0
20	0.916	1.42	0.131	0.237	1.41	0.364	0.425	0.957	3.8
30	0.845	1.46	0.121	0.175	1.07	0.387	0.417	0.960	3.3
40	0.713	1.62	0.071	0.166	0.860	0.465	0.452	0.961	3.0
50	0.631	1.81	0.062	0.158	0.680	0.534	0.423	0.953	2.5

Temperature

 $R_{\rm s}(\Omega)$ 

1.33

1.23

0.995

0.884

0.815

0.735

0.631

(°C)

-10

0

10

20

30

40

50



Fig. 7 Equivalent circuit diagram used for the analysis of EIS results.

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value at the double layer potential since the amount of H<sub>2</sub>Q adsorbed, the source of pseudocapacitance, is only 12% (Tables 2 and 3). On the other hand,  $R_{\rm ct}$  values at the double layer potential is higher than that of formal potential presumably due to the fact that with increasing electrode potential, the corresponding electric field created in the electrolyte solution will be stronger and thereby lessening the driving force for the charge/ electron transfer process taking place at the electrode/ electrolyte interface (Tables 2 and 3). Interestingly, R<sub>ads</sub> values follow a reverse trend compared to  $R_{ct}$  since at 0 V potential, the population of the electrolyte ions in proximity to the electrode will be less at double layer potential than that at formal potential (Tables 2 and 3). Diffusion capacitance  $(Q_{ms})$  revealed a higher capacitive behaviour at the double layer potential; a conclusion reached by comparing "a2" values in Tables 2 and 3 which could be due to significant electron transfer contribution at the formal potential. The  $Q_{ps}$  values are also significantly higher at the formal potentials in comparison to that at the double layer potential (Tables 2 and 3) due to the same concept described for  $Q_{\rm ms}$ . The plot of imaginary capacitance vs. frequency shows a maximum, and the inverse of this frequency is known as the dielectric relaxation constant ( $\tau_0$ ), which is the time required for the supercapacitor device to deliver 50% of its power<sup>49</sup> (Fig. 6e and f). At both the double layer potential and the formal potential, the  $\tau_0$  values have been determined and summarized in Tables 2 and 3. The  $\tau_0$  values were 4–5 times higher at formal potential compared to double layer potential since electron transfer processes require longer time scales than that of double layer formation due to electrostatic interaction. Furthermore, herein the electron transfer process also involves H<sup>+</sup> transfer between H<sub>2</sub>Q and MNC during redox reactions which may further increase time constants.

## The role of temperature in capacitance behaviour

The temperature effect on the EIS at the two potentials was also investigated and the equivalent circuit parameters are mentioned in Tables 2 and 3 which showed considerable variations. The R<sub>s</sub> values decreased with increasing temperature at both double layer potential and formation potential (Tables 2 and 3) since ionic mobility increases with increasing temperature. Similarly, R<sub>ct</sub> and R<sub>ads</sub> decreased with increasing temperature (Tables 2 and 3) since enhanced ionic mobility assist ion penetration inside the nanopores in the midfrequency region which helps the charge/electron transfer process at the electrode/electrolyte interface  $(R_{ct})$  and adsorption of ions inside nanopores  $(R_{ads})$ . In this context, these phenomena can be alternatively explained that with increasing temperature, viscosity decreases which helps the ions to move faster and thus lowering the resistances. Diffusion of ions inside the pores of the material is also assessed by observing the linear rise of imaginary component of impedance in the mid frequency range of Nyquist plots. The exponent factor "a2" associated with  $Q_{\rm ms}$  decreases with decreasing temperature suggesting a higher resistive nature with decreasing temperature or an enhanced capacitive nature with increasing temperature (Tables 2 and 3). This observation was also due to

increase of ionic mobility with increasing temperature. Furthermore, the dielectric relaxation constants ( $\tau_0$ ) at both the potentials decreased with increasing temperature due to faster kinetics (Fig. 6e, f, Tables 2 and 3).

The " $\tau_0$ " is the supercapacitor factor of merit which provides the time required for delivering 50% of power and hence it captures different types of molecular or ionic interactions required to charge a supercapacitor. Undoubtedly, the most time consuming process or in other words kinetically most hindered process primarily dictates  $\tau_0$  values. Since, time constant is inversely proportional to the rate constants, an Arrhenius type equation presented in eqn (5) can be considered.

$$\tau_0 = A \, \exp\left(\frac{E_{\rm a}}{RT}\right) \tag{5}$$

In eqn (5), A,  $E_a$ , R, and T represent pre-exponential factor, activation energy, universal gas constant and absolute temperature respectively. In  $\tau_0 vs. 1/T$  curves have been plotted at both double layer potential and formal potential in Fig. 8 and good linear fittings have been obtained. The activation barriers  $(E_a)$ were found to be 8.69 and 7.77 kJ mol<sup>-1</sup> at formal potential and double layer potential respectively. These energy values are close to that of enthalpy changes related to weak H-bonds, and it can be speculated that at the double layer potential, there will be H-bond formation between the solvated ions as well which leads to 7.77 kJ mol<sup>-1</sup> activation barrier. At the formal potential, there will be additional repeated breaking and formation of Hbonds between H<sub>2</sub>O and the oxide groups of MNC and that contributes mainly to the activation barrier. These activation barrier values are close to that of previously reported for ionhopping or proton hopping, i.e. Grotthuss mechanism. 33,37,55 It is important to emphasize that even though the two activation energies obtained at two potentials are close to each other, still marginally higher activation energy at the formal potential presumably is a reflection of the potential-driven redox reaction wherein the repeated shuttling of the H<sup>+</sup> between the -OH group of the H<sub>2</sub>Q units and the oxide sites of MNC increases the energy barrier.



Fig. 8 Arrhenius plots of  $ln(\tau_0)$  vs. 1/T at formal and double layer potential.

# Conclusion

In conclusion, the temperature dependence of a supercapacitor and its effect on electrode kinetics at double layer potential and formal potential has been investigated for a MNC-H<sub>2</sub>Q system. At a high temperature of 50 °C, a highest specific capacitance of 319 F  $g^{-1}$  has been exhibited by this system, which decreases to 213 F g<sup>-1</sup> at a temperature of -10 °C. Electrochemical impedance spectroscopy reveals improved charge transfer and double layer capacitance upon increasing temperature at both the potentials. Moreover, the dielectric relaxation time constants  $(\tau_0)$  at different temperatures at the two potentials have also been determined and it unfolds an increase in their values upon lowering of temperature. The kinetic behaviour of the supercapacitor has been determined from Arrhenius type plots where ln  $\tau_0$  has been plotted *vs.* 1/T and from the slope, the activation energies have been determined and they were 8.69 and 7.77 kJ mol<sup>-1</sup> at formal potential and double layer potential respectively. The higher activation energy at the formal potential was due to site to site electron hopping within the nanomaterial. The cyclic stability at room temperature and at an elevated temperature of 50 °C were 92% after 10 000 cycles and 86% after 5000 cycles respectively, and thus revealing superior stability of the system. Finally, we speculate that the insights into the electrode processes at different potentials achieved herein utilizing electrochemical impedance spectroscopy may lead to opening up new pathways to optimize the working principle of supercapacitor devices.

# Conflicts of interest

There are no conflicts to declare.

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