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Official URL: https://doi.org/10.1016/j.electacta.2018.11.122

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Investigation of ion transport in chemically tuned pillared graphene materials through electrochemical impedance analysis

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ABSTRACT

Chemically tuned pillared graphene structures show ability to limit restacking of graphene sheets for electrochemical energy storage in SCs. A comprehensive electrochemical characterization using various ion sizes allowed identification of ion sieving in the cross linked galleries of reduced pillared graphene materials (RPs). The access to the cross linked galleries, which provide additional ion sorption sites, offered slightly increased capacitances in RPs compared to completely restacked sheets in reduced graphene oxide (RGO). We performed electrochemical impedance analyses on RPs and RGO to understand the ion transport inside the cross linked graphene galleries. RGO adsorbs ions in the inter particle micro/meso pores and the ion access to such sites from the bulk electrolyte occurs with relative ease. RPs sieve ions into their inter layer gallery pores based on effective ion sizes and the ion transport process is resistive compared to RGO. A control study using 3D pillared graphene hydrogel with improved macro porosity assigns this resistive behavior and the moderate capacitances to limited ion access to the active sites due to excess number of pillars. The obtained results on the ion transport dynamics between graphene layers provide perspectives towards further optimization of these graphene materials for SCs.

Keywords:
Graphene
Grafting
Ion transport
Electrochemical impedance
Spectroscopy

1. Introduction

Supercapacitors (SCs) or electrochemical double layer capacitors are electrochemical energy storage devices able to deliver high power densities with excellent cycle life [1,2]. These characteristics are made possible by the charge storage process involving a charge separation at the electrode electrolyte interface via fast adsorption of electrolytic ions at the electrode surface. However, these systems are also characterized by limited energy densities and thus are widely studied as complementary to batteries [3]. To improve the energy densities, researchers are working on new systems with porous carbons, or the development of pseudocapacitive materials with additional faradaic processes, or hybrid systems that operate between batteries and SCs [4–6].

Among various carbons, activated carbons (ACs) have been widely studied for SCs applications owing to their high specific surface areas (SSA) and low material costs [4]. Interest in ACs also lies in their tunable porous structures that allowed analysis of the impact of pore sizes on specific capacitances [7,8]. These studies revealed that in addition to high surface areas, matching pore sizes to electrolytic ion sizes also enable high capacitances [9–11]. Extensive analyses on ion sieving in these pores have discovered that the high capacitances are a result of a partial desolvation and confinement of ions in these pores [12–14]. Among other carbon derivatives tested, graphene has been studied as an electrode material candidate [15–17]. The derived research paths have mainly focused on tuning graphene sample porosity [18,19], for example, by creating holes inside the graphene sheets, or by forming graphene aerogels displaying porosities adapted for efficient electrolyte ion transfer [20,21]. In parallel, many research paths involve the use of graphene as a support for a faradic or non faradic component to improve the overall capacity of the system [22,23].
Recently, our group proposed pillared graphene structures with organic molecules as pillars to prevent the significant restacking of graphene sheets during chemical reduction. We predict that such pillared structures would offer higher specific capacitances with more electrochemical active sites available for ion sorption. Hence, an interesting class of materials with varied inter graphene layer separation (d spacing), named as reduced pillared materials (RPs), were readily synthesized by varying the size of the pillar (alkyldiamine with 5, 6, 8 Cs) used [24–26]. In the course of related electrochemical studies, ion sieving behavior was observed in these graphene derived materials [24]. A clear dependency between d spacing of the inter layer galleries and electrolytic ion sizes was noted: ions with sizes smaller than d spacing can access gallery pores whereas larger ions cannot access the surface area inside the galleries [24]. The cross linked galleries in these materials sieve ions based on their size and hence demonstrate a potential for enhanced capacitances through optimization of d spacing/ion size couple [27]. While this phenomenon has been clearly identified and explained using cyclic voltammetry (CV) experiments, additional electrochemical impedance spectroscopy (EIS) studies are presented in this paper with further insights on the electrolytic ion transport in between the graphene layers. This analysis of ion dynamics inside the graphene galleries gives new perspectives and is of direct interest towards optimizing graphene materials for SCs. So, herein, pillared graphene materials with a d spacing of 0.78 and 0.80 nm—obtained with pillars of 1.5 diaminopentane and 1.6 diaminohexane respectively—were tested in various tetraalkylammonium tetrafluoroborate salt solutions in acetonitrile. Impedance analyses were performed under positive and negative polarizations to obtain the distinct anion and cation responses in these materials. Reduced graphene oxide (RGO), which only dis plays inter particle porosity but does not have any inter layer cross linked galleries, has been studied as a control material. Pillared graphene hydrogel using 1.6 diaminohexane (6 GH) with 3D porous network is also synthesized as a control material to investigate the role of porosity in 6 RP. EIS data of the materials have been analyzed using conventional Nyquist and Bode plots and a new set of analyses (evolution of imaginary part of capacitances vs. real impedances) is also proposed to investigate their ion transport characteristics.

2. Experimental methods

The pillared graphene based materials studied here were prepared using graphene oxide as a precursor with alkylamines as pillars [24]. The amine groups react with oxygen containing functional groups such as epoxide and carboxylic acid on GO and bridge GO sheets together (Scheme 1) [25]. Graphene oxide (GO), used as a precursor for pillared graphene materials, was synthesized using a modified Hummers Offeman’s method [28,29]. GO sheets were first reacted with respective diamines to form cross linked graphene materials which are then chemically reduced with hydrazine to form reduced pillared materials (RPs) (Scheme 1) [30,31]. Depending on the diamine used in the first step, materials were named as 5 RP (1,5 diaminopentane) and 6 RP (1,6 diaminohexane). RGO was synthesized by a direct chemical reduction of GO with hydrazine hydrate. 6 GH was synthesized by treating GO and 1,6 diaminohexane at 180 °C in a sealed autoclave. The detailed syntheses protocols are published elsewhere and a brief description can be found in supplementary information [24].

Electrodes were fabricated by preparing an ink using a N methyl 2 pyrroldione (NMP) slurry of the synthesized materials (80%) with polyvinylidene fluoride binder (10%) and carbon black (10%) conductive additive. This ink is drop casted on stainless steel disks and dried at 65 °C under air for 6 h followed by 80 °C under vacuum overnight. The dried electrode specific weights were around 3–4 mg/cm².

Electrochemical analyses of the materials were performed in three electrode cell configuration using a silver wire as the quasi reference electrode, an overcapacitive (10 times by weight) YP50 porous carbon pellet as counter electrode and 20 μm cellulose piece as the separator in Swagelok® cells. 1 M tetraalkylammonium tetrafluoroborate (TABF₄) salt solutions in acetonitrile with constant anion size and varying cation sizes; ethyl (TEABF₄), propyl (TPABF₄), butyl (TBABF₄) and hexyl (THABF₄) were used as electrolytes. The unsolvated sizes of the electrolyte ions tested are TEA⁺—0.68 nm, TPA⁺—0.76 nm, TBA⁺—0.82 nm, THA⁺—0.95 nm and BF₄⁻—0.48 nm. Anhydrous acetonitrile and electrochemical grade electrolyte salts for the study were purchased from Sigma Aldrich. Electrochemical cells were assembled inside an Ar filled glovebox and once sealed have been electrochemically tested outside using a VMP 3 potentiostat device from Bio Logic. Prior to EIS measurements, the cells have been cycled at 50 mV/s scan rates in CVs until the electrolyte wets the whole porous material volume and the electrochemical signature reached gets a steady state—the area under the curve stabilizes (~100 cycles). For EIS measurements, cells were held at +1 V or -1 V vs. Ag in this 3 electrode configuration until the current is almost null and then studied with sinusoidal voltage amplitude of 5 mV within a frequency range spanning from 100 kHz to 10 mHz. Obtained impedance values

Scheme 1. Schematic representation of reduced graphene oxide (RGO) and reduced pillared graphene materials (RPs) showing the specific surface areas and d-spacing values determined from XRD. XRD patterns adapted with permission from Chem. Mater. 30, 2018, 3040-3047. Copyright (2018) American Chemical Society [24].
were normalized with electrode weights for plotting.

3. Results and discussion

3.1. Reduced pillared materials: characteristics

In our previous work, pillared graphene structures in RPs were analyzed using a combination of spectroscopic and electrochemical characterization methods (Fig. S1) [24–26]. X-ray diffraction (XRD) patterns of RGO, synthesized as a control, showed a broad peak around 28° corresponding to the graphite like arrangement (002) with an inter graphene sheet distance of 0.37 nm (Fig. S1a). This graphite like arrangement suggested an efficient reduction of GO leading to a partial restacking of the reduced graphene sheets [32]. Whereas, the XRD patterns of RPs displayed two peaks corresponding to partial graphitic restacking (−25°) and diamond induced pillaring of graphene layers (−15°). Thus, RPs demonstrate pillared structures characterized with d spacing of 0.78 nm in 5 RP and 0.80 nm in 6 RP. The N₂ gas sorption analyses have assigned a high SSA of 330 m²/g for RGO and lower values of ~330 m²/g for RPs. Lower surface areas in RPs compared to RGO could arise from excessive inactive weight of alkyldiamines in RPs, structural changes during degassing and the diamine pillars may also sterically hinder the gas molecules and render the inter layer gallery pores inaccessible [33,34]. The pore size distributions (PSDs), calculated using 2D non local density functional theory (2D NLDFT), attribute high volume of micropores and a significant contribution of mesopores for RGO and a high mesopore content with some micropores in RPs (Fig. S1c) [35].

The electrochemical responses of RGO and RPs in various electrolytes with increasing cation sizes and constant anion size investigated the ion sieving effect in pillared graphene materials (Fig. 1). CVs of RGO and RPs showed similar rectangular quasi ideal capacitive behavior in TEA BF₄ but, an increase in cation size from TPABF₄ to TBABF₄ and THABF₄ resulted in current decay under negative polarizations for RPs (Fig. 1 & Fig. S2a). Thus, the access to inter layer gallery pores in RPs is regulated by the d spacing/ion size couple whereas RGO doesn’t have any pillared structures and hence, exhibits similar curves in different electrolytes. The calculated capacitances values were lower in RPs when the inter layer graphene galleries have a smaller d spacing than the ion sizes, implying that the galleries restrict larger ions (Fig. S2b). For instance, 5 RP shows a specific capacitance of 130 F/g in TEA BF₄ but when larger cations were used, a steep decline from 115 F/g (TPABF₄) to 42 F/g (THABF₄) is noted. Whereas, RGO, which adsorbs ions only at the surface of graphene layers (SSA 330 m²/g), exhibits similar capacitances of ~110 F/g in all the electrolytes studied. In short, this electrochemical study revealed ion sieving based on ion sizes inside the pillared graphene materials [24].

3.2. Impedance studies: Nyquist plots

To further characterize this ion sieving and also to gain insights into the ion transport inside the graphene galleries, EIS measurements were performed on RGO and RPs under negative and positive polarizations. The normalized complex plane Nyquist plots are displayed in Fig. 2. These plots can be divided into three regions [36–38]. At high frequencies, only the electrical conductivity of the electrode (ionic and electronic) can be seen; thus, SC electrodes behave like a resistor (Z<sub>ι</sub> = 0). The sharp increase of the imaginary part of the impedance (Z′) in the low frequency region where the plot is nearly parallel to the y axis is related to a capacitive behavior. The intermediate 45° region is associated to the ion transport in the porous network of the electrode that corresponds to the electrolyte resistance in the pores.

At 1 V vs. Ag, the electrodes are negatively polarized with respect to open circuit voltages (OCV) and cations are predominantly adsorbed at the electrode surfaces [39]. The Nyquist plots show differences between RGO and RPs in all the electrolytes tested (Fig. 2a–c). Firstly, in each electrolyte, as compared to RGO, RPs show a larger electrolyte resistance in the porous electrode network that extend into lower frequencies. Due to this resistance, the low frequency capacitive behavior in RPs shifts into higher resistances. Secondly, with increasing cation size, the vertical line indicating capacitance in each RP gradually diverges away from the ideal 90°. The extent of this deviation from the ideal capacitance can be estimated by replacing the capacitance C with a Constant Phase Element (CPE, see Equation (1)) using a parameter [40]. α values are obtained by measuring the slopes in equation (1) at low frequencies.

\[ \log \left| Z'(\omega) \right| = \alpha \log(C_{0}) \quad 0 < \alpha < 1 \] (1)

A value of 1 for α indicates a fully capacitive behavior of the material. 6 RP shows decline of α values from 0.87 in TEA BF₄ to 0.77 in TBABF₄ while, 5 RP shows low value of ~0.80 in all the electrolytes indicating a non ideal capacitive behavior (Table S1). Thus, there is an important influence of the cation size on the frequency responses of RPs under negative polarization. RGO exhibits α values close to 1 from TEA BF₄ (0.95) to TBABF₄ (0.93) electrolytes, evidencing a faster ion transport in the pores.

Ion sorption in RGO occurs in the micro/meso inter particle pores and thus the TEA<sup>+</sup>, TPA<sup>+</sup> and TBA<sup>+</sup> cations access them with relative ease. Therefore, the impedance measurements of RGO show ideal capacitive like behavior and do not vary a lot in these electrolytes. In RPs, ion sorption occurs significantly in the inter layer porous network in addition to the inter particle pores. Access to the inter layer pores in RPs is regulated by d spacing/ion size couple. Therefore, the frequency responses of RPs are

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Fig. 1. Normalized CVs of RGO and RPs obtained at a scan rate of 10 mV/s in the voltage range of −1 V to 1 V vs. Ag in 1 M solutions of (a) TEA BF₄, (b) TPABF₄ and (c) TBABF₄ in acetonitrile. Normalization was performed with respective currents observed at open circuit voltages (~0.1 V).
dependent on their d spacing and the effective size of the electrolyte ions. 6 RP, with a d spacing of 0.80 nm, has a performance similar to that of RGO in TEABF₄ but extended diffusion regions are seen with larger cations. Impedance data in THABF₄ electrolyte further supports the different sorption mechanisms for RGO and RPs (Fig. S3a). In RPs, THA⁺ cation (0.95 nm) is larger than the d spacing values, like with TBA⁺, and hence even larger impedances were recorded. Notably, RGO possesses a major contribution from inter particle pores with a diameter of ~0.9 nm and thus only when THA⁺ cations are used, a sudden increase in the Warburg region and decrease in α parameter (0.83) are observed (Table S1).

At +1 V vs. Ag, the materials are positively polarized with respect to open circuit voltages and BF₄⁻ ions are predominantly adsorbed on the electrode surfaces. The Nyquist plots of RGO and RPs show similar shapes among themselves in the three different electrolytes tested (Fig. 2d–f). The BF₄⁻ ions are smaller than the gallery pores in RPs and hence display capacitive like behavior similar to RGO. However, the vertical part in the low frequency region slightly shifts away from the ideal behavior from TEABF₄ to TBABF₄, α values for RGO range from 0.97 to 0.95 whereas RPs range from 0.93 to 0.86 in TEABF₄ to TBABF₄ (Table S1). This interesting shift from ideal behavior despite analyzing the same BF₄⁻ ions in different electrolytes hints at a possible role of cations even under positive polarization, such as previously observed [41]. In summary, RGO and RPs show similar responses in all electrolytes under positive potentials and an increasing influence of larger cations, in addition to anions, is also noted. This observation was described as an ion exchange mechanism in earlier reports [41–43]. With the largest cation in THABF₄, α values of RGO and RPs decrease to 0.90 and ~0.76 respectively, indicating a greater cation exchange effect.

3.3. Impedance studies: complex capacitance analysis

In order to further explore the frequency responses of the synthesized materials, we performed a complex capacitance analysis that was proposed by Taberna et al. and derived from Cole and Cole [44–46]. In the proposed analysis, complex capacitance is studied in order to focus on the capacitive behavior and ionic resistance within the active material. The closest equivalent circuit representation of the capacitor in this analysis has a series resistance, Rₛ, together with an angular frequency dependant RₛCₛ bridge: a combination of resistance, Rₛ(ω), in parallel with a capacitance, Cₛ(ω) (Fig. S5).

This complex analysis allowed determination of the real capacitance, C'(ω), which represents the deliverable capacitance of the cell under DC or low frequency AC conditions. C'(ω) is
calculated using equation (2), where \(Z''(w)\) is the imaginary part of impedance \(Z\) and \(w\) is the frequency.

\[
C'(w) = \frac{\dot{Z}'(w)}{w|Z(w)|^2}
\]  

(2)

The evolution of \(C'\) with respect to frequency for each sample at 1 V and -1 V vs. Ag was plotted accordingly (Fig. 3). At higher frequencies, the system acts as a pure resistor with capacitance values close to zero and at lower frequencies, a pure capacitive behavior is seen as a plateau if the maximal \(C'\) occurs in the tested frequency range. The intermediate zone gives information about electrolyte access to the electrode porosity (gallery and inter particle pores) [44].

At 1 V vs. Ag, \(C'\) vs \(w\) plots for RGO approach a maximal \(C'\) plateau in TEABF\(_4\), TPABF\(_4\) and TBABF\(_4\) at the lowest frequency tested (10 mHz), revealing a fast transport of electrolytic ions in the electrode material (Fig. 3a–c). However, in THABF\(_4\), RGO displays lower \(C'\) values and a non ideal behavior as the cation size approaches the inter particle pore sizes in RGO (Fig. S4a). For 6 RP, \(C'\) is closely equivalent to that of RGO in TEABF\(_4\) and increasing cation sizes shifts the pure capacitive behavior to lower frequencies. 5 RP, even in TEABF\(_4\), does not approach a plateau and the ion transport in gradually limited in other electrolytes. Additionally, with larger cations in RPs, the maximal \(C'\) values and the slopes in the intermediate region gradually decrease.

This EIS data under negative polarization really shows a slower ion transport in RPs when larger cations are used. In RPs, steric effects when ion size is in the same range as the inter layer gallery size lead to limited ion transport and accessibility to adsorption sites. Thus, with a complex dynamic process, RPs do not reach a plateau with full capacitance in this frequency range and confirm resistive behavior. At +1 V vs. Ag, RGO and RPs all show curves with similar maximal \(C'\) values and slopes in the intermediary region (Fig. 3d–f). However, with increasing cation size, it can be noted that the plateaus of \(C'\) slightly drift towards lower frequencies. This observation, as noted in the Nyquist plots, could be seen as an influence of cations via the existence of an ion exchange mechanism [41]. Results obtained with THABF\(_4\) under positive potential show a steep fall of \(C'\) values in all the materials, supporting this effect (Fig. 5Sb). Moreover, an increase in the alkyl chain length from TEA\(^+\) to THA\(^+\) ions could give rise to stronger interactions between graphene sheets and alkyl carbons of the electrolytic ions, resulting in enhanced influence of cations.

The complex capacitance model also allows the determination of imaginary capacitance (\(C''\)) which corresponds to the energy loss arising from irreversible processes at the electrodes [44]. This \(C''(w)\) parameter was calculated from EIS data using equation (3), where \(Z''(w)\) is the real part of impedance \(Z\):

\[
C''(w) = \frac{\dot{Z}''(w)}{w|Z(w)|^2}
\]

(3)

The imaginary \(C''\) goes through a maximum at a frequency \(f_0\), which defines the transition between purely resistive and capacitive behaviors of a SC. This frequency, known as knee frequency, defines the time constant (\(\tau_0\)) as minimum time required to extract half of the capacitance from a SC [45]. \(C''\) vs \(w\) plots of RGO and RPs at +1 and -1 V vs. Ag show similar and dissimilar curves respectively, supporting the role of ion discussed above. (Figs. S6 and S7). The \(\tau_0\) values obtained from these curves assign better power capability for RGO than RPs (Table 1). Interestingly, the \(\tau_0\) values for

---

**Table 1**

<table>
<thead>
<tr>
<th>Material</th>
<th>TEABF(_4) +1 V</th>
<th>-1 V</th>
<th>TPABF(_4) +1 V</th>
<th>-1 V</th>
<th>TBABF(_4) +1 V</th>
<th>-1 V</th>
<th>THABF(_4) +1 V</th>
<th>-1 V</th>
</tr>
</thead>
<tbody>
<tr>
<td>RGO</td>
<td>6</td>
<td>6</td>
<td>6</td>
<td>6</td>
<td>6</td>
<td>6</td>
<td>6</td>
<td>12</td>
</tr>
<tr>
<td>5 RP</td>
<td>7</td>
<td>10</td>
<td>11</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>6 RP</td>
<td>10</td>
<td>15</td>
<td>25</td>
<td>20</td>
<td>32</td>
<td>41</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

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Fig. 3. Real capacitance (\(C'\)) vs frequency plots for RGO and RPs in 1 M solutions of (a,d) TEABF\(_4\), (b,e) TPABF\(_4\) and (c,f) TBABF\(_4\) in acetonitrile at (a,b,c) -1 V vs. Ag and (d,e,f) 1 V vs. Ag.
5 RP are lower than for 6 RP in all the electrolytes tested under positive polarization \[45\]. For example, \( \tau_0 \) of 5 and 6 RPs in TBABF4 are 11 and 20 s respectively. This difference between RPs is counterintuitive as one would expect 6 RP, with a higher density spacing, to facilitate better transport and lower \( \tau_0 \) value for BF4 anion. As \( \tau_0 \) values are affected by a combination of ionic, electrical and cell resistances, it seems important to determine the distinct impact of the material properties on ionic resistances.

To do so, herein, we studied C" with respect to Z'. We believed that such an analysis could enable facile quantitative determination of the effective ionic resistances (R\(_i\)) involved in ion transport. The R\(_i\) values directly obtained here could be seen as equivalent to the one third of the in pore ionic resistance calculated by employing the Transmission Line Model \[47\]. Similar to other plots, three different regions could be identified in such a curve (Fig. 4). At high frequencies the SC performs like a resistor and C" would be close to zero (from Eq. (3)). The mid frequency range is related to the ion transport inside the electrode porous structure hence, a steep in zero (from Eq. (3)). The Ri values were obtained from the real Z values at the start and end of the mid range frequencies.

The C" vs Z' plots at \(-1\) V vs. Ag and \(+1\) V vs. Ag show a clear difference between RGO and RPs in TEABF4, TPABF4 and TBABF4 (Fig. 4a-c). The responses of RGO and RPs in various frequency regions and their evolution in electrolytes could be distinctly identified here compared to Nyquist and Bode plots. RGO displays curves with a clear maxima of C" occurring at a low resistance and then rapidly decreasing along the x axis. The calculated Ri show close values of around 10 mOhm.g from TEABF4 to TBABF4 (Fig. 4g). 6RP Ri is closely equivalent to that of RGO in TEABF4 (Fig. 4a) but, with larger cations, the C" extends to even higher resistances and eventually shows a continuous plateau with TBABF4 (Fig. 4c). The plateau symbolizes an extended limitation of ion transport inside the porous electrode while the cell is not reaching a capacitor like behavior. This increasingly resistive behavior of 6 RP with larger cations is quantified with rise in Ri values from 20 mOhm.g in TEABF4 to 70 mOhm.g in TBABF4. 5 RP plot shows continuous plateau with no transition peak even with the smallest cation tested in TEABF4 and already shows a high Ri of 55 mOhm.g (Fig. 4a). Further increase in cation sizes lead to even higher Ri up to 172 mOhm.g in TBABF4 (Fig. 4c). This continuous loss of energy even at the low frequencies led us to examine the leakage currents for all cells. The measured leakage currents show similar values for RGO and RPs in all electrolytes and suggest that the observed continuous energy losses are characteristics of the materials (Table S2). At \(+1\) V vs. Ag, RGO and RPs show similar curves, however, occurring at different values of real Z (Fig. 4d–f). The C" values for RPs occur at greater resistances than RGO indicating greater irreversible energy losses. Among the two RPs, 6 RP has slightly larger ionic resistances than 5 RP in all the electrolytes.

![Fig. 4](image)

Fig. 4. Imaginary part of capacitances (C") vs. real impedances (Z') for RGO and RPs in 1 M solutions of (a,c) TEABF4, (b,d) TPABF4 and (c,f) TBABF4 in acetonitrile at (a,b,c) \(-1\) V vs. Ag and (d,e,f) \(-1\) V vs. Ag. The arrows in (a), (b) and (c) show increasingly larger ionic resistances (R\(_i\)) in 6 RP. The obtained Ri values for RGO and RPs at \(-1\) V vs. Ag and \(+1\) V vs. Ag.
This resistive behavior of 6 RP supports the earlier observed greater \( \tau \) values than 5 RP. We speculate that the conformation (trans & gauche) of the diamines inside the gallery could be the reason behind this behavior. It has been reported that with increasing alkyl chains lengths in the diamine the ratio of trans gauche conformations shifts towards higher trans content [25]. Hexanediamine (in 6 RP) may locally occupy higher volume in the gallery than pentanediarnine (in 5 RP) and thus have higher ionic resistances. Moreover, the odd and even number of carbons in the diamines used here could also affect the overall polarity and ionic sorption in the gallery. The cation sorption at 1 V, masks these fine details among RPs as the steric effects result in large \( R_i \) values. As the \( R_i \) values at +1 V are smaller, this subtle influence of the alkyl chain conformation on 5 and 6 RPs is seen distinctly. Overall, the proposed \( C' \) vs. \( Z' \) plots show an amplified difference between the materials compared to Nyquist, Bode plots or other representation proposed by Pickup et al. [48]. It is also noteworthy that we obtain effective ionic resistances directly from the experimental data without using TLM or PSD TLM models [47,49].

### 3.4. Investigating role of porosity

Earlier analysis of capacitance values from CVs have shown marginally higher values for RPs than for RGO with appropriately smaller ions (TEABF\(_4\)). However, even in the case of TEABF\(_4\), the impedance analysis has indicated that RPs exhibit sluggish ion transport relative to RGO. This could be explained by hindered mobility of the ions in the galleries due to unadapted porosity or the presence of alkyl diamine pillars. To investigate if the impeded ion transport is arising from unsuited porous network, we synthesized and analyzed a pillared graphene hydrogel (GH) that has a 3D percolating network using 1,6 hexanediamine as the pillar (6 GH) (details in SI). The 3D percolating network offers multiple ion pathways and is known to show enhanced ionic mobility characteristics in the hydrogels [22]. The \( N_2 \) gas sorption isotherms and corresponding PSDs characterize 6 GH with enhanced micro and meso pores compared to 6 RP (Fig. S9 a&b). The calculated SSAs indicate increase in the values from 130 m\(^2\)/g in 6 RP to 182 m\(^2\)/g in 6 GH (Table S3). Scanning electron microscopic images also reveal more open macro porosity in 6 GH than 6 RP (Fig. S9 c&d). Additionally, 6 GH shows greater d spacing of 0.86 nm compared to 0.80 nm of 6 RP probably due to the higher reaction conditions and diamond consumption during 6 GH synthesis. The TGA profiles of the two materials support this explanation with a greater weight loss due to decomposition of covalently attached diamines.

Electrochemical analysis of 6 GH in 1 M TEABF\(_4\), which has sufficiently smaller ions than the d spacing, shows slightly higher capacitances of 142 F/g than 6 RP (130 F/g) and RGO (117 F/g). This shows that the 3D assembly with improved SSAs, along with the cross linking at nanoscale, yields enhanced capacitances in these pillared materials. However, the impedance analysis under negative polarizations shows that the ion motion remains very much impeded in 6 GH despite its 3D structure and the larger d spacing (0.86 nm) (Fig. S5 a and b). In fact, the obtained \( R_i \) and \( \tau \) values show greater resistances in 6 GH with higher values of 57 mOhm.g and 23 s compared to 20 mOhm.g and 17 s for 6 RP (Table S3). These observations suggest that the alkyl diamine pillars inside the galleries play an important role in ion transport between the graphene layers. Moreover, the obtained capacitances of both 6 RP and 6 GH are only slightly better than RGO despite enhanced ion access to the cross linked galleries. This is possibly due to excessive filling of the galleries with diamine pillars leading to impeded ion transport to active sites. This understanding of the impact of pillars and the porosity provides new strategies to improve energy and power densities of pillared materials. Decreasing the number of pillars between the graphene layers could be a straightforward strategy for enhanced power and energy values. Integrating additional inter particle pores could be another strategy for enhancement of energy densities. Collectively, pillared materials with low pillar density and additional inter particle pores could offer both high energy and power densities.

### 4. Conclusions

Impedance measurements have been performed to investigate the ion transport dynamics in pillared graphene based materials. A range of electrolytes with increasing cation sizes and a constant anion size were studied under positive and negative electrode polarizations. Depending on the cation size, Nyquist and Bode plots of RPs exhibited large Warburg like regions and delayed capacitive behaviors whereas RGO responses remained similar. Ionic resistances obtained through original \( C' \) vs. \( Z' \) plots enabled a quantitative analysis of these observations and show that the ion transport in RPs is mainly controlled by graphene inter layer gallery sizes. As RPs sieve ions into the pillared graphene layers, steric effects impede the ion transport when ion size is in the same range as the inter layer gallery size. RGO adsorbs ions in the inter particle micro/meso pores and the ion access to the active sites is relatively easier. The observed high ionic resistances for RPs compared to RGO are thus in accordance with the earlier CV experiments that displayed poor power capability for RPs at high scan rates. This
knowledge from cyclic voltammetry and the dynamic ion transport between graphene layers materials opens up perspectives towards enhanced charge storage in graphene materials.

Acknowledgments

This work was supported by CEA (H. Banda) and by French ANR Program (SUGGEST ANR 15 CE05 0022). The access to mutualized equipment and PFNC Nano characterization Minatec platforms was appreciated by the authors. G. Bidan and D. Aradilla are also thanked for fruitful discussions and advice.

Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.lectacta.2018.11.122.


