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Water confinement in nanoporous silica materials

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The influence of the surface polarity of cylindrical silica nanopores and the presence of Na⁺ ions as compensating charges on the structure and dynamics of confined water has been investigated by molecular dynamics simulations. A comparison between three different matrices has been included: a protonated nanopore (PP, with SiOH groups), a deprotonated material (DP, with negatively charged surface groups), and a compensated-charge framework (CC, with sodium cations compensating the negative surface charge). The structure of water inside the different pores shows significant differences in terms of layer organization and hydrogen bonding network. Inside the CC pore the innermost layer is lost to be replaced by a quasi bulk phase. The electrostatic field generated by the DP pore is felt from the surface to the centre of pore leading to a strong orientation of water molecules even in the central part of the pore. Water dynamics inside both the PP and DP pores shows significant differences with respect to the CC pore in which the sub-diffusive regime of water is lost for a superdiffusive regime. © 2014 AIP Publishing LLC. [http://dx.doi.org/10.1063/1.4862648]

I. INTRODUCTION

Water is vital for all forms of life and also plays a key role in all types of industrial development. Despite its simple chemical composition, this essential molecule in nature exhibits several unique properties such as large surface tension and high boiling temperature due to its ability to form hydrogen bond networks.

In recent years, a large body of research work has focused on the properties of water adjacent to a surface due to their important implications in a various fields such as evaporation, crystallization, or confined fluids.1 Notably, understanding the behavior of water trapped within pores with dimensions in the nanometer range is of considerable interest, given that it is relevant to systems and phenomena in biology, such as protein stability and permeability of ion channels, geology, such as transport through porous rocks, chemistry and chemical engineering through molecular sieves and nanofluidic devices.2

At the nanometric scale the dimensions of the system are comparable to the range of intermolecular interactions, and thus, liquid-liquid and liquid-pore wall interactions have to be considered explicitly in order to understand the behaviour of confined water.1 Many factors are likely to impact the structure and dynamics of confined liquids, such as the size, the shape, and the nature (hydrophilic or hydrophobic) of the confining medium. Confined liquids have been studied in both hydrophobic and hydrophilic media from several experimental techniques such as nuclear magnetic resonance,4 differential scanning calorimetry,5 neutron diffraction,6 etc. However, many fundamental properties of confined water have yet to be fully understood and the difficulty in coming to a full understanding of the structure of confined water via purely experimental data highlights the need for complementary atomistic-level simulations.7 Indeed, molecular simulations can explicitly account for the molecular nature of water, and they can also treat surfaces with atomically detailed precision.8 They have proved to be a powerful method to investigate confinement9–12 and interfacial effects,13,14 notably by describing changes in hydrogen bonding, molecular orientation, transitional and rotational dynamics of confined fluids.15

Investigating the impact of confinement on the physical properties of water trapped in silica nanopores is of particular interest since these latter can be readily fabricated with well-controlled geometry and highly ordered directions, with broad implications in catalysis, biology, nanofluidics, etc.16 Consequently, in recent years molecular simulations have been employed to investigate structural, dynamics, and dielectric properties of water confined in silica nanopores of various sizes, geometries, and hydrophilic/hydrophobic features.2,3,16–24

However, most of these studies only reported results for aqueous solutions confined inside uncharged systems. In contrast, only few works have focused on the properties of aqueous solutions confined into charged silica nanopores. Argyris et al.8 and Ho et al.25 have carried out molecular dynamics (MD) simulations of aqueous electrolyte solutions confined within slit-shaped pores of crystalline silica with various degrees of protonation. Zhu et al.26 have investigated the dynamics properties of ions confined within both uncharged and negatively charged cylindrical pores of amorphous silica. In the three studies mentioned above, the surface charge was balanced by redistributing the appropriate counter-charge on the atoms of the silica framework so as to fulfill the condition.

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of overall electroneutrality of the system. Alternatively, Bonnau et al. have investigated the structural and dynamics properties of water confined within negatively charged slit-shaped silica pores by adding calcium ions in solution so as to compensate the negative surface charge. Pore widths ≤20 Å have been considered in Refs. 8, 25, and 27. Under such extreme confinements, characterized by the absence of bulk-like water even at the pore centre, the properties of water are particularly badly understood and become highly sensitive to the pore geometry and the surface properties.

Here we present the results of MD simulations of water-filled cylindrical hydrophilic silica nanopores (pore diameter: 12 Å) considering a protonated nanopore, a deprotonated framework, and a silica material where the deprotonation of SiOH groups has been compensated by Na⁺ ions. This comparison allowed us to emphasize the impact of the presence of ions on both the structure and dynamics of confined water.

An important and non-trivial issue in performing simulations of liquids under nanoconfinement is the density used in the simulations. In this study, the water density inside pores was determined by performing MD simulations in the Isothermal–Isosurface–Isobaric statistical ensemble since this alternative method to Grand Canonical Monte Carlo simulations has been shown efficient to compute the density of liquids such as methanol and water inside mesoporous silica and the MIL-53(Cr) metal organic framework. Special care was taken to model the confined liquid phase under realistic conditions, i.e., considering nanopores in contact with water-filled external reservoirs.

Our motivation in performing this new set of simulations was to provide insight into the impact of the surface polarity and of the presence of Na⁺ ions on the structure and the dynamics of water under severe cylindrical confinement in hydrophilic media.

II. MODEL AND COMPUTATIONAL DETAILS

Water was modelled from the rigid TIP4P/2005 model of Abascal and Vega. The silica cylindrical pore was created from a cubic cell using a procedure proposed by Brodka and Zerda. Thus, the inner surface coverage of silanol (SiOH) groups has been compensated by Na⁺ ions. This comparison allowed us to emphasize the impact of the presence of ions on both the structure and dynamics of confined water.

TABLE I. Partial charges of atoms of the three pores. Partial charges on Si, Ob, Obh, and H of the PP substrate were taken in Ref. 31. Charges for the DP pore were compared with those obtained from DFT calculations. From their coordination numbers, we distinguished bridging oxygen Ob bonded to two silicon atoms from non bridging oxygen Obn bonded to only one silicon and bonded to one hydrogen atom.

<table>
<thead>
<tr>
<th>Charges (u.e.)</th>
<th>Si</th>
<th>Ob</th>
<th>Obn</th>
<th>Hb</th>
<th>Na</th>
</tr>
</thead>
<tbody>
<tr>
<td>PP pore</td>
<td>1.275</td>
<td>−0.637</td>
<td>−0.533</td>
<td>0.206</td>
<td>…</td>
</tr>
<tr>
<td>DP pore</td>
<td>1.284</td>
<td>−0.628</td>
<td>−0.533</td>
<td>…</td>
<td>…</td>
</tr>
<tr>
<td>CC pore</td>
<td>1.269</td>
<td>−0.652</td>
<td>−0.533</td>
<td>1.000</td>
<td>…</td>
</tr>
<tr>
<td>DFT charges in DP pore</td>
<td>1.422</td>
<td>−0.705</td>
<td>−0.454</td>
<td>…</td>
<td>…</td>
</tr>
</tbody>
</table>

from a cubic cell using a procedure proposed by Brodka and Zerda. Thus, the inner surface coverage of silanol (SiOH) was 7.5 nm², which corresponds to highly hydrated silica MCM-41. From their coordination numbers, we distinguished bridging oxygen Ob bonded to two silicon atoms from non bridging oxygen Obn bonded to only one silicon and bonded to one hydrogen atom. This matrix corresponds to the protonated pore labelled PP. The pore radius was set at 6 Å in order to explore water under ultra confinement. To explore the influence of the surface polarity two additional pores were created: a deprotonated pore (DP) and a compensated-charge pore (CC). In the PP pore the O–H distances and the Si–O–H angles were maintained from a harmonic bond and valence potential. The complete force field used for the silica pores is reported in Table I and the different sets of charges are gathered in Table II. Partial charges of Si, Ob, Obh, and H atoms of PP pore type were taken from Ref. 31. An illustration of the three matrixes is given in Figure 1. To build the DP matrix hydrogen atoms of SiOH in PP framework were removed while the residual negative charge was redistributed among the atoms of the silica material. To ensure this “ad hoc” charge redistribution, we extracted the partial charges from a DFT calculation (see Table II). In this purpose a small cluster consisting of interfacial silanol groups was used. The cleaved bonds were saturated with methyl groups so as to maintain a standard hybridization. A snapshot of this cluster is provided in Figure 2. To extract the partial charges of DP matrix the DFT calculation was performed with the DMOL package from Accelrys Inc. In this calculation the polarized all electron atomic base DNP was used. The main conclusion was that the charges of silicon and oxygen atoms are close in both cases and the deviation is around 10%. Eventually
the CC framework was created using a method described by Bonnaud et al.\textsuperscript{27} To compensate the removal of hydrogen atoms the silanol groups, sodium ions were introduced. A non-polarisable model was considered to model Na\textsuperscript{+}.\textsuperscript{28} In contrast with Bonnaud et al.\textsuperscript{27} the partial charge of the oxygen atom of the silanol groups was conserved in the three frameworks to quantify the exact influence of the deprotonation and the presence of Na\textsuperscript{+} on water structure and dynamics. Finally, Na\textsuperscript{+} ions were added randomly within the pore. Molecular Dynamics simulations (MD) were carried out in the Isothermal–Isosurface–Isobaric (III) statistical ensemble\textsuperscript{10} where the volume is only allowed to vary axially. As shown in Figure 3, two reservoirs were added so as to study the bulk and confined properties of water from a single simulation. MD simulations were performed with DL\_POLY package.\textsuperscript{33} Long-range interactions were computed using the Smooth Particle Mesh Ewald sum method.\textsuperscript{34} Lennard-Jones potential was considered to allow for the van der Waals interactions. Crossed interactions were computed from the Lorentz-Berthelot rule. The temperature (300 K) and pressure (1 bar) were kept constant using the Berendsen thermostat and barostat with the relaxation constants of 0.5 fs and 0.1 fs, respectively.\textsuperscript{35} The integration algorithm was the Velocity Verlet algorithm with a time step of 2 fs.\textsuperscript{36} The simulations ran during 10 ns with 3 ns of equilibration.

\section{III. RESULTS AND DISCUSSION}

\subsection{A. Structure}

We report in Figure 4(a) the radial water density inside the three kinds of pores. The density profiles shown in Figure 4(a) were computed by considering the centre of mass of water molecules. For the PP matrix there are two local maxima at $r = 1 \text{ Å}$ and $r = 4 \text{ Å}$ suggesting a layering organization due to the excluded volume.\textsuperscript{28} The period of oscillation is about $3-4 \text{ Å}$ that basically corresponds to the molecular diameter of water molecules. Inside the deprotonated pore (DP), two water layers are still observed, with density maxima located at $r = 1.5$ and $4.5 \text{ Å}$ (the distance between the two layers being similar to what was observed inside the PP pore). Additionally layers are shifted toward the pore surface with respect to water confined into the PP pore. This shift is caused by the absence of the interfacial protons of the DP pore. Note that the different orientation of water molecules in PP and DP pores (see later) also contributes to shift water layers a bit closer to the surface inside the DP pore. Interestingly, the water density profile is very different inside the CC pore. A water layer is still observed close to the pore surface (approximately at the same position as inside the PP matrix). On the other hand, the layering organization of water molecules vanishes in the inner zone of the CC pore (i.e., the central part of the pore, see Figure 4(b)) for which the density of bulk water is recovered, i.e., $\sim 1000 \text{ kg m}^{-3}$. This particular structure inside the CC pore can be imputed to the presence of the sodium cations compensating the surface charge. We report in
FIG. 5. (a) Radial density profiles of sodium ions and water in the CC pore. Water density can be read on the right axis while number of Na\(^+\) is read on the left axis. (b) Radial water density in PP and CC pores of 12 Å in radius.

Figure 5(a) the radial density profile of Na\(^+\) in the CC pore. Figure 5(a) shows that the Na\(^+\) ions are located close to the silica interface and are absent from the inner region of the pore. This result cannot be compared with the radial distribution of Na\(^+\) ions obtained by confining a NaCl solution in the PP pore.\(^{28}\) Indeed, in CC pore Na\(^+\) ions interact with the oxygen atoms of the negatively charged SiO groups so as to compensate the negative charge on the framework. Thus the usual physics ruling the distance between a protonated wall and the monovalent ions cannot be discussed here. Interestingly, let us note that the hydration of Na\(^+\) ions share the interfacial zone with the water molecules. Thus, 75 Na\(^+\) ions were found inside the CC pore after equilibration, which corresponds to an ion concentration about 15 M. This huge amount of charge-compensating ions completely modifies the organization of water molecules and removes the layering organization in the inner zone. In other words, charge-compensating ions inside highly charged nanopores tend to screen confinement effects and to remove surface effects on water structure. Interestingly, the first water layer close to the interface is still present inside the CC pore, which can be attributed to excluded volume effect. The impact of charge-compensating ions on the water structure inside pores (decrease in confinement effect) was also observed in a CC pore of 12 Å in radius for which the third layer (inner zone) is removed (see Figure 5(b)).

In order to analyze the effects of the surface on the water layers inside the different pores, the orientation of the water molecules was further investigated. Figures 6(a) and 6(b) report the distribution of the angle \(\theta\) between the vector between the dipolar moment of water molecules and the normal to the pore surface for both the inner and outer zones (see Figure 4(b) for illustration of these zones). Far from the interface, all angles were fairly sampled in both the PP and CC matrixes. This suggests that no orientation is privileged and that water molecules do not feel the surface. To the contrary, at the centre of the DP framework the maximum of the angular distribution is located around 180°. This suggests that the oxygen atom of water molecules is pointing at the centre of the pore while the hydrogen atoms are in direction of the surface. As shown in Figure 6(b) this effect is also observed close to the interface. This strong orientation is ruled by the interfacial negative charges. Figure 6(b) depicts a preferential orientation of water molecules around 180° close to the interface of the CC pore. As for the DP pore this orientation is probably ruled by the negative charge surface. As shown in
Figure 7 shows the radial distribution functions between the hydrogen atom of water (Hw) and the interfacial oxygen atoms of silica (O_{int}). RDFs were corrected for the excluded volume.\(^{22,23}\) Thus, it seems that the interfacial organisation (between \(R = 4\) Å and \(R = 6\) Å) is controlled by the surface/water interactions. However, the hydrogen bonding network would be different between DP and CC pores given the location of Na\(^+\) close to the interface.

In Figures 8(a) and 8(b), and 8(d) we report the radial profile of the average number of hydrogen bonds (nHB) per water molecule between water-water, water-substrate and the total HB number (water-water + water-substrate), respectively. The geometric criteria established by Chandler\(^{21}\) were used to identify both the hydrogen bonds between the pore and the water molecules and between water molecules themselves. For the PP matrix, Figure 8(a) shows that the nHB at the centre of pore is similar with the value in the bulk value (3.6). That suggests that the hydrogen bonding network is not modified at the centre of the PP pore. The number of HB per water molecule decreases from \(r = 3.1\) Å and falls down to 2.2 close to the surface. This decrease in interfacial nHB is due to the finite size effect. Indeed, since the pore radius is 6 Å and the molecular diameter of water is about 3 Å the number of coordination in the first hydration shell decreases for \(r > \sim 3\) Å. This tendency is accentuated inside the CC pore where nHB falls down from \(r = 2\) Å to reach a value of 1.4 close to the surface. Indeed, interfacial Na\(^+\) ions lead a local organisation of water molecules around cations breaking the hydrogen bonding network. For the DP framework a smaller nHB is observed in the central part of the pore (nHB = 3.2) with respect to both the PP and CC pores. This decrease in nHB at the centre of pore results from the impact of the surface charge which orientates water molecules from the surface to the centre of the pore (as shown in Figures 6(a) and 6(b)). This strong orientation prevents the setting of the usual tetrahedral organisation of the HB network and leads to a decrease in nHB. Interestingly, some discrepancies are also observed between the three pores regarding the radial profile of the number of HB formed between water molecules and the surface. Indeed, Figure 8(b) shows that water molecules form more HB with the surface of the DP than with the surface of PP and CC pores, which can be understood since water molecules can approach closer to (and so interact more easily with) the surface oxygen atoms in the case of the DP pore. This is corroborated by the radial density of Ob and Onb atoms reported in Figure 8(c) where the distribution of Ob is non-null at \(r = 6.2\) Å. This accessibility to the Ob sites allows an increase in nHB (water-surface) in the DP matrix leading to a constant value of total nHB(r) over the whole pore cross-section (Figure 8(d)). To get an accurate standpoint of the hydrogen bonding network we computed its “rigidity” which is defined as the ratio between the HB number per water molecule and the average number of neighbours.\(^{21}\) Figure 8(e) shows that the average number of neighbours in the centre of the three matrixes is similar and is slightly lower than in the bulk phase. Interestingly, we observe that the radial profile of the average number of neighbours is also correlated to the radial profile of HB number (Figure 8(a)). As shown in Figure 8(f) the “rigidity” term is higher close to the surface. This could be due to a faster interfacial decrease in the number of neighbours caused by excluded volume effect. Additionally, we observe
that the rigidity of HB network in the DP pore is weaker than in the PP and CC pores. This can be correlated with the decrease in HB number in the DP substrate (Figure 8(a)). We also explored the distortion of the HB network by calculating the distribution of the angle between the intramolecular bond O–H and the intermolecular O–Ob vector for the three matrixes. In this calculation we considered the angle between two water molecules such that the distance between two water molecules connected by a hydrogen bond satisfied rOH < 2.5. As shown in Figure 9(a) the maximum is located at 1011(b) and 40W. We adjusted the charge on internal structure of HB. We report in Figure 9(b) the distribution of the angle between the intramolecular bond O–H and the intermolecular O–Ob vector (Ob is the bridging oxygen between two silicon atoms; see Figure 2). Figure 9(b) shows two possible orientations of water close to the surface of the PP pore, at 20° and 90°. In contrast with the PP pore the CC and DP substrates do not present a maximum at 90°. We can conclude that this preferential orientation is probably due to the OH group of silanol.

In Figure 10 we provide the radial distribution functions between the oxygen atoms of water molecules in bulk phase and inside the different nanopores. Usually, the tetrahedral structure is highlighted from the position of the second hydration shell.58 Clearly, we observe a decrease in intensity of the main peak due to the excluded volume while the second shell vanishes in both DP and CC pores. In DP pore its absence can be imputed to the surface polarity while in the CC material its absence is due to the huge ion concentration.

**B. Dynamics**

The translational dynamics of water molecules was investigated from their mean square displacement (MSD). We report in Figure 11 the radial and the axial components of the MSD Mean Square displacement (MSD) of water molecules in the three matrixes in both inner and outer regions. The MSD components were calculated from Eqs. (1) and (2).

\[
MSD_{||} = \left\langle \sum_{i=1}^{N} \left[ z_{com,i}(t + t_0) - z_{com,i}(t_0) \right]^2 \right\rangle, \tag{1}
\]

\[
MSD_{\perp} = \left\langle \sum_{i=1}^{N} \left[ r_{com,i}(t + t_0) - r_{com,i}(t_0) \right]^2 \right\rangle, \tag{2}
\]

with \( r_{com} = (x_{com}^2 + y_{com}^2 + z_{com}^2)^{1/2} \).

The subscript \( com \) indicates that the centre of mass of water molecules was considered in this calculation. The brackets in both cases denote an average over the number of water molecules considered in the corresponding areas and over the different configurations. Axially (Figures 11(a) and 11(b)) water molecules in both the inner and outer zones of the pores do not reach the diffusive regime, i.e., the MSD\(_{||}\) is not proportional to time and has a long time dependence of the form

\[
MSD = at^\alpha.
\]

As discussed by Gallo and Rovere anomalous diffusion corresponds to \( \alpha \neq 1 \) such that \( \alpha > 1 \) is named superdiffusion and \( \alpha < 1 \) subdiffusion.39,40 We adjusted the MSD\(_{||}\) from the function \( at^\alpha \) to extract \( \alpha \). We report in Table III the values of \( \alpha \). Interestingly we observe that at the centre of pores \( \alpha < 1 \) which indicates a subdiffusive behaviour. For the CC substrate \( \alpha \) is higher than for the two others pores, which shows a less pronounced “anomalous” water behaviour under confinement. Close to the pore surface (outer zone) this subdiffusive regime is also observed into the PP and DP matrixes while in the CC framework a superdiffusive regime is highlighted. Indeed, we found \( \alpha = 1.21 \) in the CC pore. This result suggests a subdiffusive to superdiffusive regime transition along the radial position from the centre to the interface of the CC pore. The radial components of MSD...
TABLE III. Hydrogen bond life time and $\alpha$ coefficient obtained from a fit of the axial MSD (MSD$_{\parallel}$) from MSD$_{\parallel} = \alpha t^\beta$.

<table>
<thead>
<tr>
<th></th>
<th>Inner zone</th>
<th>Outer zone</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\alpha$ coefficient</td>
<td>0.77</td>
<td>0.77</td>
</tr>
<tr>
<td>PP pore</td>
<td>0.65</td>
<td>0.84</td>
</tr>
<tr>
<td>DP pore</td>
<td>0.93</td>
<td>1.21</td>
</tr>
<tr>
<td>CC pore</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hydrogen bond life time (ps)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>PP pore</td>
<td>3.3</td>
<td>6.8</td>
</tr>
<tr>
<td>DP pore</td>
<td>2.3</td>
<td>6.2</td>
</tr>
<tr>
<td>CC pore</td>
<td>7.2</td>
<td>20.0</td>
</tr>
</tbody>
</table>

are shown in Figures 11(c) and 11(d). For the three pores three regimes seem to be drawn, a plateau of diffusion for $t > 100$ ps, a subdiffusive regime for $10 \text{ ps} < t < 100 \text{ ps}$ and a probable ballistic regime for $t < 10$ ps. The absence of diffusion in the confined direction was already reported in similar systems with different geometries.

The investigation of the rotational motion of the water molecules was based on the computation of the auto-correlation function $C(t)$ of the total dipole moment (Eq. (3)).

$$C(t) = \left( \frac{M(0)M(t)}{M(0)^2} \right), \quad (3)$$

where $M$ is the sum of the dipolar moment of water molecules. In Figure 12 we report the correlation function $C(t)$ as function of time. Figure 12 shows an absence of dipolar decorrelation in both DP and CC pores. In the DP pore this absence of decorrelation can be explained from the strong orientation of the water molecules leading a reduction in the rotational motion of the water molecules. For the CC matrix the correlation of the dipolar moment of water molecules is higher than with the DP pore and is caused by the orientation of water molecules around the Na$^+$ ions.

We report in Table III the hydrogen bond life time ($\tau_{\text{HB}}$) in inner and outer zones for the three matrixes. The life time is calculated from an exponential adjustment of the correlation function $C_{\text{HB}}(t) = \left( \frac{h(0)h(t)}{h(0)^2} \right)$ where $h$ is unity when two tagged molecules are hydrogen bonded at time $t$ and is zero otherwise. This function $C_{\text{HB}}(t)$ indicates the conditional probability that a HB remains intact at time $t$, given it was intact at time $t = 0$. $C_{\text{HB}}(t)$ does not consider breaking of HB at intermitten times between time zero and $t$. For all pores $\tau_{\text{HB}}$ is higher in the outer region than in the inner zone. This interfacial increase in $\tau_{\text{HB}}$ is due to a decrease in interfacial $n_{\text{HB}}$ leading to a decrease in HB exchanging increasing the HB life time in the outer zone. The longest HB life times are observed in the CC pore (both in the inner and outer zones). As shown in Table III $\tau_{\text{HB}} = 7.2$ ps and 20 ps in inner and outer zones, respectively. This increase is due to the dipolar saturation of water leading to a preferential orientation around of Na$^+$ ions and then a decrease in HB exchange.

IV. CONCLUDING REMARKS

Molecular dynamic simulations of water confined into cylindrical silica nanopores with different surface polarities have been performed and analysed. The structure and dynamics of confined water have been studied for three matrixes, namely, (1) a protonated matrix (PP) with silanol surface groups, (2) a deprotonated pore (DP) where the protons of the PP pore have been removed such that the residual charge has been redistributed in the framework, and (3) a deprotonated pore where the charge of the protons has been compensated by adding Na$^+$ cations inside the pore (CC).

Structure of water has been studied by means of radial distribution functions, number of hydrogen bonds, and angular distribution of the angle between the water dipolar vector and the normal to the surface. The results obtained with the three matrixes revealed significant difference in radial organisation of water molecules. Two water layers form into both PP and DP pores while only one layer has been obtained in the CC pore, the water density in central part of this pore being bulk like. In the DP matrix the electrostatic field, caused by the removing of the proton, is felt to the centre of pore leading a strong orientation of water molecules.

Translational and rotational dynamics have been studied from the calculation of mean square displacement and the dipolar correlation, respectively. The results obtained from the three matrixes show a drastic difference between the PP and DP pore and the CC matrix. Indeed, we found one superdiffusive regime of water with the CC framework while a subdiffusive regime was evidenced with PP and DP pores. Additionally, a sub- to superdiffusive regime transition has been evidenced in the CC pore.

These results show that a deprotonation of the surface can change the diffusion and the structure of the confined fluid which is likely to modify substantially the transport properties of electrolytes solutions through nanoporous membranes as those used, e.g., in desalination processes. This point will be addressed in a subsequent work.

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