

# Chapter 9. Molecular Structure and Dynamics of Nano-Confined Water: Computer Simulations of Aqueous Species in Clay, Cement, and Polymer Membranes

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## MOLECULAR STRUCTURE AND DYNAMICS OF NANO-CONFINED WATER:

## COMPUTER SIMULATIONS OF AQUEOUS SPECIES IN CLAY, CEMENT, AND POLYMER MEMBRANES

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Abstract. Molecular-level knowledge of the thermodynamic, structural, and transport properties of water confined by interfaces and nanopores of various materials is crucial for quantitative understanding and prediction of many natural and technological processes, including carbon sequestration, water desalination, nuclear waste storage, cement chemistry, fuel cell technology, etc. Computational molecular modeling is capable to significantly complement the experimental investigations of such systems by providing invaluable atomic-scale information leading to improved understanding of the specific effects of the substrate structure and composition on the structure, dynamics and reactivity of interfacial and nano-confined aqueous solutions. This paper offers a brief overview of recent efforts to quantify some of these effects for individual H<sub>2</sub>O molecules and hydrated ions confined at the interfaces and in nanopores of several typical hydrophilic and hydrophobic materials. The first molecular layer of aqueous solution at all substrates is often highly ordered, indicating reduced translational and orientational mobility of the H<sub>2</sub>O molecules. This ordering cannot be simply described as "ice-like", but rather resembles the behavior of supercooled water or amorphous ice, although with very significant substrate-specific variations.

## 1. INTRODUCTION

Most chemical reactions near the Earth's surface involve fluid (aqueous) phases and take place at fluid-solid interfaces or in confined spaces of mineral interlayers and nanopores. These reactions affect many important natural processes, including mineral weathering, adsorption or release of environmental contaminants in soil, drinking water quality, formation and behavior of ice nano-crystals and hydrated mineral nano-particles in the atmosphere, the fate of CO<sub>2</sub> in geologic carbon sequestration. The same interfacial phenomena are also at the core of many important technological processes related to water purification and desalination mambranes, fuel cells, production and utilization of cement, etc. All these phenomena are inherently multi-scale in time and space, and addressing the scientific and technological problems they pose is multidisciplinary. Molecular-level knowledge of the chemistry and physics of interfacial and confined aqueous solutions interacting with mineral surfaces and other substrates is essential to their understanding. Important questions center on how water and dissolved species behave at the surfaces and in nanoconfinement by different inorganic and organic substrates.

Current understanding of hydrated mineral-water interfaces is far from complete. Molecular-scale information is difficult to obtain experimentally due to the nature of the materials (e.g., interfacial and bulk structural disorder, the presence of molecular scale dynamical disorder with characteristic time scales that span many orders of magnitude, the physical limitations on the samples that can be effectively examined, and the difficulty of experiments under extreme conditions). Mineral surfaces themselves vary by the arrangements and reactivity of individual surface functional groups and the specific adsorption of inorganic and organic ions. Even the utility of macroscopic thermodynamic concepts, such as the dielectric constant and electric double-layer, remain open and are important questions driving the current research [1-3]. Modern experimental probes, including synchrotron light sources, high-resolution neutron scattering, time-resolved spectroscopy and diffraction, scanning force microscopy, and multi-dimensional NMR spectroscopy, are providing unprecedented, atomic-level and surface-specific information about mineral-fluid systems [3-5]. However, it is often difficult to interpret these experimental observations without parallel theoretical and computational studies [6-7]. In addition, the interaction with experimentalists is imperative for verifying theoretical models and guiding larger scale simulation efforts. Connecting computational and experimental results is crucial in addressing these questions and in bridging the gap between the atomic- and nanometer-scales and the micron- and larger scales that is essential for environmental, geochemical and energy applications of practical importance [8].

The methods of atomistic computer simulations [9], often coupled to surface-specific experimental results, are already mature enough to provide otherwise unobtainable perspective into the structure, dynamics and energetics of fluid-mineral systems [10]. Computational approaches range from traditional quantum chemical methods to classical molecular dynamics (MD) and Monte Carlo (MC) simulations using semi-empirical interaction potentials to emerging methods including quantum molecular dynamics, metadynamics, and reactive force fields for molecular-scale modeling [11-35]. Critical questions currently revolve around the effect of solid surfaces on the properties of near-surface and

nano-confined fluids, structural and chemical modification of the surface by the fluid, the pathways of chemical reactions at surfaces and in fluids, the connections between these reactions and fluid structure and dynamics, and the consequent effects on the pathways of dissolution and precipitation. This paper provided a brief overview of our group's recent efforts to quantify some of these properties and processes at the interfaces and in nano-confinement with several typical hydrophilic and hydrophobic materials by classical MD simulations using the CLAYFF force field [12].

#### 2. COMPUTATIONAL MODELS AND METHODS

We compare the molecular structure of several qualitatively different substrate-water interfaces. Crystals of metal hydroxides, line **brucite**, **Mg(OH)**<sub>2</sub>, or **portlandite**, **Ca(OH)**<sub>2</sub>, can be naturally cleaved parallel to their (001) crystallographic plane, resulting in an electrostatically neutral surface fully covered by OH groups oriented away from the surface. They can easily form hydrogen bonds with interfacial H<sub>2</sub>O molecules. Therefore, we will use the surface of portlandite as a reference model of a hydrophilic interface [11,20]. A similar hydrophilic surface can be created by cleaving the crystal structure of **quartz**, **SiO**<sub>2</sub>, parallel to the (001) plane. In the presence of water and at relatively low pH, this surface will also be fully covered by OH groups [36]. In the structure of **hydrocalumite**, **Ca<sub>2</sub>Al(OH)**<sub>6</sub>]**Cl·2H<sub>2</sub>O**, 1/3 of the divalent Ca cations are isomorphically substituted by trivalent Al, resulting in a positive structural charge compensated by the presence of Cl<sup>-</sup> or some other anions in the interlayer space and on the surfaces of this material, an anionic clay [11,20]. Similar to its neutral portlandite analog, its cleaved (001) surface is also fully covered by OH groups and can be considered hydrophilic.

The structure of talc,  $Mg_3Si_4O_{10}(OH)_2$ , consists of so-called TOT layer (an octahedral layer of  $[MgO_4(OH)_2]$  sandwiched between two tetrahedral layers of  $[SiO_4]$ ) stacked along the crystallographic Z direction [18,21,27]. All tetrahedral sites are occupied by Si forming  $[SiO_4]$  structural units, and all octahedral sites are occupied by Mg forming  $[MgO_4(OH)_2]$  units. The TOT layers are, thus, electrostatically neutral and interact with each other only by weak by van der Waals forces, reflecting the hydrophobic nature of the basal siloxane Si-O-Si (001) surface. The TOT layered structure is typical for many clay minerals, which can develop various degrees of negative structural charge due to the isomorphic substitutions possible for both tetrahedral (e.g., Al for Si) and octahedral (e.g., Li for Mg) layers. The emergence of negative layer charge is balanced by the presence of charge-balancing cations in the interlayer space and on the surfaces of these minerals and makes them more hydrophilic. Their interaction with water have already been extensively investigated [12,25,26,29,30,32,37] and we focus here only on the behavior of neutral talc, as a reference hydrophobic surface.

It is known experimentally (e.g., [38]) that mineral surfaces rarely exist in nature in the atomically smooth ideal form as the ones listed above. To illustrate the effects of nanoscale surface roughness of the substrate on the structural and dynamic properties of the interfacial aqueous solution, the surface of **tobermorite**, **CasSi6O16(OH)2**, is taken here for comparison. Tobermorite is also known as one of the so-called C-S-H (calcium silicate hydrate) phases in cement chemistry and its surface properties control behavior of confined water in the nanopores of cement and concrete [11,20].

The disordered interfaces of water filtration membranes are also typically characterized by the surface roughness on the nanoscale and above. In this case, the diversity of the surface adsorption sites can be very large and it is not always possible to identify and investigate all of them individually of a truly atomistic level. Nevertheless, computer simulations can still provide invaluable information about the molecular mechanisms of the processes of membrane filtration. The effects of various metal cations and their interactions with dissolved natural organic matter on the process of bio-fouling of **polyethersulfone** (**PES**) **desalination membrane** is taken here to illustrate the case [39].

All MD simulations described here were performed at ambient conditions in the *NVT* statistical ensemble (constant number of atoms, constant volume, and temperature T = 300 K) after a pre-equilibration in the *NPT* ensemble at T = 300 K and P = 0.1 MPa using standard MD algorithms [9]. The mineral surfaces were constructed by cleaving the bulk crystal structures parallel to the (001) plane. The irregular polymer surface was created in the course of preliminary MD runs of condensing a polyethersulfone polymer chain consisting of 60 monomer units [39]. Each complete interfacial MD simulation cell consisted of a substrate slab about 2 to 3 nm thick, and a layer of liquid water about 3 to 7 nm thick placed in contact with them, as illustrated in Fig.1. The water was pre-equilibrated at 300 K and 0.1 MPa. Periodic boundary conditions [9] were then applied in all three dimensions to produce models of the interfaces formed by infinite (in x- and y directions) flat substrate layers interspersed with layers of water. In all cases, the thickness of the water layer and the size of the cell in the z-direction (normal to the surface) were sufficiently large to effectively exclude direct interactions between two different substrate/water interfaces created due to the periodicity of the system. The number of H<sub>2</sub>O molecules in the layer was chosen to reproduce the density of bulk water under ambient conditions (~1 g/cm³). The final x-, y-, and z-dimensions of the simulation supercells were approximately z = z-.

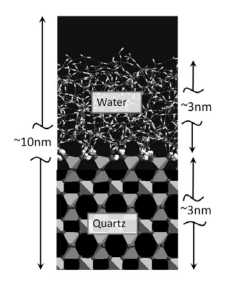


Figure 1: A schematic view of the molecular dynamics periodic simulation cell for the modeling of a substrate-water interface. Bulk quartz SiO<sub>2</sub> crystal is shown as tetrahedral (Si) and dark balls (O). White (H) and dark (O) balls represent the O-H groups of the fully hydroxylated surface. Approximately half of the surface O-H groups are bent to make hydrogen bonds among themselves, while the other half – donate H-bonds to the interfacial H<sub>2</sub>O molecules, which are shown as white (H) and dark (O) cylinders above the surface. H-bonds are indicated as thin dashed lines.

The MD time step was 1.0 fs, and each model system was allowed to equilibrate for at least 500 ps of MD before an equilibrium MD trajectory for each model was accumulated at 10 fs intervals during additional 500 to 1000 ps of MD simulation. The details of the MD simulation procedures and parameter settings for mineral-water systems are described in previous publications [11,12,18,20,27,37,39-41]. The structural analysis of the near-surface water films was undertaken using atomic density profiles in the direction perpendicular to the substrate surface and atomic density maps within defined slices of the water film parallel to the surface. These properties were all calculated by averaging over the last 500 ps equilibrium MD trajectory of each system. The position of the mineral surface (z = 0) was defined by the average position of the top-most surface oxygen atoms or of the corresponding OH groups, depending on the substrate.

## 3. SIMULATED RESULTS AND DISCUSSION

## 3.1. Interfacial water structure

Fig. 2 shows the time-averaged atomic density contour map of  $O_w$  in the first molecular layer of water on the (001) surface of quartz. It is important to note, that unlike the surfaces of brucite [18] and portlandite [11], only about 50% of the surface hydroxyl groups on quartz are oriented away from the surface and easily donate H-bonds to the interfacial  $H_2O$  molecules. The other 50% of the surface OH groups are oriented parallel to the surface of quartz, form H-bonds to other surface hydroxyls, and accept H-bonds from  $H_2O$  molecules (Figs1,2). The surface ordering of the  $H_2O$  molecules is clearly observed in the time-averaged patterns of the  $O_w$  distribution. Surface water molecules are significantly immobilized by each accepting an H-bond from one surface OH group and simultaneously donating one H-bond to another neighboring surface OH group. A similar pattern of  $H_2O$  ordering is also observed at the neutral hydroxylated surfaces of brucite [18] and portlandite [11]. However, the charged hydroxylated surface of hydrocalumite exhibits a completely different behavior where no H-bonds donation is possible for the interfacial water molecules [11].

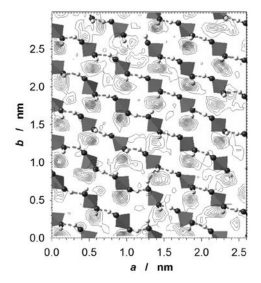


Figure 2: Contour maps of the atomic density of oxygens of  $H_2O$  ( $O_w$ ) within the first molecular layer of water at the fully hydroxylated (001) surface of quartz, which is shown below the contours as tetrahedra (Si) and dark (O) and white (H) balls . See Fig. 1for the detailed notations.

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This is clearly visible from the comparison of the atomic density profiles for O<sub>w</sub> and H<sub>w</sub> of these surfaces, which provide important additional insight into the ordering of water molecules at the interface. Two orientations of H<sub>2</sub>O molecules are energetically most favorable for the formation of stable hydrogen bonds to the hydroxylated surfaces of portlandite and quartz (Fig.3a,b). In one of them, water molecules have one H-atom directed toward an O-atom of the surface hydroxyl, thus donating an H-bond to the surface. This is reflected in the presence of a strong peak at ~0.2 nm on the  $H_w$  density profiles. In this orientation, the second H-atom contributes to the peak at ~0.35 nm. However, a stronger contribution to the 0.35 nm peak in the H<sub>w</sub> distribution arises from the other favorable H<sub>2</sub>O orientation, in which it accepts an H-bond donated by the surface hydroxyl groups. In this case, both H<sub>H2O</sub> of the surface water molecule contribute to the density profile peak at 0.35 nm. In both preferred orientations of the surface water molecules, their O-atoms are located approximately at the same distance from the surface (~0.25-0.28 nm). The ability of the surface water molecules to both donate and accept H-bonds to the substrate surface creates a very well developed Hbonding network across the interface with the structure strongly resembling that of bulk liquid water. O-O and O-H radial distribution functions of liquid water are shown as thick dashed lines in Fig. 3 to illustrate this similarity. The  $g_{\rm OH}(r)$ , peak at ~0.2 nm is a definitive signature of donated hydrogen bonds in the system, while the  $g_{\rm OO}(r)$  peak at ~0.45 nm clearly indicates that the local ordering of the H-bonding network is close to tetrahedral - a typical characteristic of bulk liquid water structure.

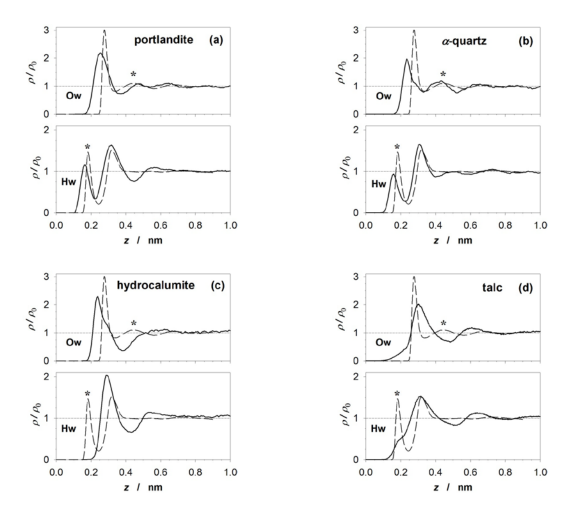


Figure 3: Density profiles of  $O_w$  and  $H_w$  atoms of the interfacial  $H_2O$  molecules as functions of the distance from the surface (z) of four different mineral substrates. Origin of z is defined here for all surfaces as the average Z-coordinate of the oxygen atoms at the substrate surface. The local density  $(\rho)$  is normalized with respect to the corresponding atomic density in bulk liquid water  $(\rho_0)$  and the radial distribution functions  $g_{OO}(r)$  and  $g_{OH}(r)$  of bulk liquid water are shown as thick dashed likes for comparison. Stars mark the characteristic H-bonding features of these functions.

For hydrocalumite (Fig.3c) the computed near-surface water structure is strikingly different. Due to the positive structural charge of this substrate, the H-bond donation to the surface by H<sub>2</sub>O molecules is prevented and the

well-interconnected H-bonding network is not formed in the interfacial region [11], despite the fact that the  $H_2O$  molecules are strongly attracted to this surface and it is clearly hydrophilic.

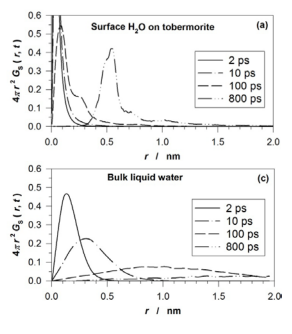
The atomic density profiles of the surface water on talc (Fig.3d) show that  $H_2O$  molecules are located somewhat farther away than from the previously discussed hydrophilic surfaces (Figs.3a-c) and the  $H_2O$  molecular orientation is predominately parallel to the surface (the peaks of  $O_w$  and  $H_w$  are approximately at the same distance of 0.3-0.32 nm), in accordance with the hydrophobic character of talc. However, donation of weak H-bonds is still possible to this surface, as is clearly evident from the shoulders of the  $O_w$  and  $H_w$  distributions around 0.2 nm. Thus, talc surface exhibits a weak degree of hydrophilicity under ambient conditions, which can be most pronounced at very low relative humidity [33], or can even be turned into a strongly hydrophilic surface by the application of very high pressures and temperatures typical for the Earth's mantle conditions [40,41].

## 3.2. Interfacial water dynamics

The simulated diffusional dynamics of surface water is in agreement with these observations: the  $H_2O$  molecules have noticeably higher mobility at the hydrophobic surface of talc and significantly lower mobility at all hydrophilic surfaces [18,20]. Longer-time-scale MD simulations provide an additional opportunity to quantify these relatively slow diffusional motions of interfacial by means of the Van Hove self-correlation function (VHSCF) [9],

$$G_{S}(r,t) = \frac{1}{N} \left\langle \sum_{i=1}^{N} \delta[r + r_{i}(0) - r_{i}(t)] \right\rangle, \tag{1}$$

where r is the coordinate of the molecule, t is time, and N is the total number of molecules. This relationship describes the correlation in the positions of the same atom at different times, such that  $4\pi r^2 G_S(r,t) dr$  is the probability of finding an atom at distance r after a time t if the position of this atom was at the origin r=0 at the initial time t=0. The Fourier transform of the VHSCF represents the incoherent or self-intermediate scattering function (SISF), which can be directly measured in incoherent quasielastic neutron scattering experiments and contains detailed information concerning the single-molecule dynamics both in time and space domain.



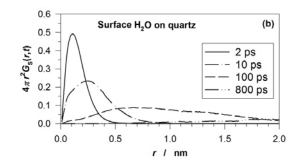


Figure 4: Van Hove self-correlation functions for diffusing  $H_2O$  molecules at the surface of tobermorite (a) and quartz. The VHSCF behavior of  $H_2O$  molecules in bulk liquid water (c) is also shown for comparison.

The calculation of this self-correlation function for the water molecules strongly bound to the surface of tobermorite, which exhibits nanoscale surface roughness strongly affecting the first two monolayers of  $H_2O$ , shows a dramatic difference in the dynamic behavior of adsorbed  $H_2O$  (Fig. 4a), as compared with the diffusional dynamics in bulk liquid water (Fig. 4c), and yields a characteristic time scale of the diffusional process of the order of  $\tau_m$ ~0.8 ns, in excellent agreement with NMR measurements, indicating a mean time between diffusional jumps on the surface of ~0.8-1.0 ns [43]. The calculated long-time-scale VHSCF also points to a hopping diffusional mechanism with a length scale of  $l_m$ ~5.5 Å, i.e., close the characteristic lattice dimension of crystalline ice between two neighboring strong H-bonding sites. From the two-dimensional Einstein equation,  $D = \langle |\mathbf{r}(t) - \mathbf{r}(0)|^2 \rangle / 4\tau$ , the mean time for jumps between

surface sites from the NMR results yields a diffusion coefficient of  $0.9\times10^{-10}$  m²/s, which is also in remarkable agreement with the average diffusion coefficient for all surface-associated H<sub>2</sub>O molecules obtained from MD simulations  $(1.0\times10^{-10}$  m²/s). Here,  $\bf r$  is the mean jump displacement (assumed to be 5.5 Å, from the results of the VHSCF calculations, Fig.4a) and  $\tau$  is the mean jump time,  $\tau \sim \tau_m \sim 0.8$  ns. This level of agreement definitely provides strong support for the interpretation of the experimental results and strong encouragement for further application of atomistic computational modeling techniques to study the molecular scale properties of nano-confined water.

The hopping surface diffusion mechanism is not evident for the water molecules at an atomically smooth surface, such as that of quartz (Fig.4b): the corresponding VHSCFs show only slight differences with the diffusional dynamics in bulk liquid water (Fig.4c).

## 3.3. Effect of metal cations on the membrane fouling

Prediction of ultrafiltration membrane performance is not generally possible without performing pilot-scale tests because membrane fouling is related to membrane material and solution chemistry. The inadequate understanding of membrane fouling has in many cases hindered wider adoption of membrane processes in large-scale drinking water treatment plants. Computational molecular modeling of the model membrane-solution interfaces allowed detailed probing of the fouling process due to the interaction of dissolved natural organic matter (NOM) with various metal cations [39]. It was observed that divalent ions (Ca<sup>2+</sup> and Mg<sup>2+</sup>) may cause membrane fouling not by forming "ionic bridges" between the negatively charged functional groups on the membrane surface and the negatively charged functional groups of NOM, but rather by promoting the aggregation of NOM molecules in solution.

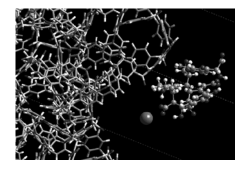


Figure 5: A fragment of the MD simulation box containing the polyethersulfone membrane surface (cylinder representation), a calcium ion (large dark ball), and a molecule of natural organic matter (ball and stick representation) coordinating to Ca<sup>2+</sup> by its deprotonated carboxylic group. All interfacial water molecules are hidden for clarity.

Thus, even though the presence of truly dissolved NOM is not important in the fouling process, it can contribute to fouling after being aggregated by divalent ions. The partially neutralized Ca<sup>2+</sup>-NOM complex (Fig. 5) may be more easily adsorbed at the membrane surface than a negatively charged NOM molecule. However, it still remains unclear whether fouling is caused by the cation-mediated NOM aggregation in solution or by stronger NOM-surface interactions in the presence of divalent ions. Since only one NOM molecule was used in the present model, NOM aggregation could not be quantitatively studied in this work. More extensive MD simulations using multiple NOM molecules, employing much larger size of the simulated systems, and much longer simulation times (at least on the order of tens of nanoseconds) are necessary to quantitatively address these problems.

## 4. CONCLUSIONS

The methods of computational molecular modeling in application to the mineral-fluid systems have advanced dramatically in the past decade and have already emerged as very powerful quantitative tools in the studies of the structure and properties of interfacial and nano-confined aqueous solutions. There is a growing consensus, however, that the idealized mineral-fluid interfaces typically used in the present-day molecular simulations and the relatively small assemblages used in these models and in quantum calculations are at least an order of magnitude too small in characteristic length (three orders of magnitude in volume) to effectively address many critically important issues. A typical size of the systems currently simulated is on the order of a few nanometers, but can reach up to 20 nm [26] for classical force-field-based MD simulations which can probe the systems evolution over tens of nanoseconds. However, the application of quantum MD is still limited to the sizes < 1.5-2 nm and time periods ~10 ps. The inability to effectively model realistic compositional and structural complexity is significantly impeding further progress. For instance, idealized, defect-free mineral surfaces are of little use in predicting local charge development, which is key to understanding surface reactivity in real systems [13,16]. Particle edges, surface defects, the surface roughness on length scales of 10 to 100 nm play central roles in determining the energetics and kinetics of dissolution, precipitation, sorption and catalytic reactions at mineral interfaces, but these phenomena are currently modeled only very rarely and on a limited scale [19,21,34,35]. There is no doubt, however, that further rapid development of supercomputing capabilities

- will allow such molecular simulations to achieve truly realistic system sizes and time scales, thus significantly
- 236 increasing their practical value.
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