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Solvent-mediated interactions between nanostructures: From water to Lennard-Jones liquid

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Solvent-mediated interactions emerge from complex mechanisms that depend on the solute structure, its wetting properties, and the nature of the liquid. While numerous studies have focused on the first two influences, here, we compare the results from water and Lennard-Jones liquid in order to reveal to what extent solvent-mediated interactions are universal with respect to the nature of the liquid. Besides the influence of the liquid, the results were obtained with classical density functional theory and brute-force molecular dynamics simulations which allow us to contrast these two numerical techniques. Published by AIP Publishing. https://doi.org/10.1063/1.5037571

I. INTRODUCTION

Interactions between two solids are usually well characterized by their intrinsic physical and chemical properties. However, in the presence of a liquid solvent, additional interactions emerge and can become dominant when the solids are also electrically neutral. This so-called solvent-mediated interaction is involved in various phenomena including self-assembly, ligand unbinding, and protein folding.

Numerous studies have examined solvent-mediated forces in terms of range, strength, and sign using both numerical and experimental techniques. In particular, it was found that the sign of solvent-mediated forces is controlled by the equilibrium contact angle. On the one hand, when the solids are solvophilic, solvent molecules are attached to the solid surface. Bringing the two solutes together leads to a perturbation, and the removal of this favorable structure causes a strong repulsive hydration pressure. On the other hand, when the solids are solvophobic, solvent molecules which are located between the two solids are expelled and a more stable vapor cavity emerges thus reducing the overall free energy. This so-called capillary evaporation has been the subject of numerous studies and leads to a solvophobic attraction. Along with the wetting properties of the solid, the role of its geometrical structure was also covered in several studies. For example, Jabes et al. recently demonstrated that using the same solid composition and size, qualitatively different solvent-mediated forces can be obtained only by changing the solid shape between fullerenes, nanotubes, and graphene-like structures.

Altogether, this work is integrated to a general mean-field theory of hydrophobicity developed by Lum, Chandler, and Weeks and further refined in more recent publications.

When modeling liquids using numerical simulations, two approaches are commonly employed. On the one hand, because water plays a crucial role in most applications and especially in biological systems, atomistic models for water molecules have been developed in order to mimic its thermodynamic properties and some special features including strong hydrogen bonding and ice polymorphism. On the other hand, generic model systems such as hard spheres and Lennard-Jones (LJ) potential are also often used for modeling fluids. Once the model is chosen, solvent-mediated forces can be computed using various types of numerical methods. Monte-Carlo and molecular dynamics (MD) simulations are widely employed especially for water modeling while classical density functional theory (DFT) in which molecules are treated as a density field can grant access directly to liquid density profiles and the corresponding free energy. DFT is less numerically expensive and avoids using free energy calculation techniques such as thermodynamic integration, transition path sampling, and umbrella sampling. However, DFT for water is not as highly developed as for simple fluids.

While numerous authors have suggested the ability of the Lennard-Jones liquid to reproduce behaviors similar to water regarding solvent-mediated effects, there is no detailed comparison of solvent-mediated forces obtained with atomistic simulations of extended simple point charge (SPC/E) model water and with DFT calculations of Lennard-Jones (LJ) particles. In this work, we make a direct comparison of solvent-mediated forces obtained from molecular dynamics simulations of water and DFT calculations of LJ using a very generic system made of two nanometric crystalline slabs immersed in a liquid. Free energy is computed as a function of the interslab distance, and we study thoroughly the influence of wettability and of the slab geometrical structure. Our work identifies differences and similarities between atomistic simulations of water and DFT calculations of LJ. Moreover, our results contribute to the overall understanding of solvent-mediated forces. We also discuss more generally to what extent molecular properties of water make it special in comparison to simple fluid models.

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II. METHODS

A. Studied system

Our calculations make use of two types of molecules: slab molecules and liquid molecules. The slabs are composed of rigid arrangements of solid molecules while the liquid is treated dynamically. We hold constant temperature and density of the liquid while varying the solid properties. The slabs are made of three square layers of $40 + 41 + 40 = 121$ atoms which are kept fixed in a face centered cubic (FCC) structure with the (100) face exposed and with the lattice spacing, $a$. The interaction between the slab and liquid particles is modeled via a Lennard-Jones potential parametrized by its length scale, $\sigma_{\text{wall}}$ and well depth, $\epsilon_{\text{wall}}$. When varying $a$, $\sigma_{\text{wall}}$ is also modified using $\sigma_{\text{wall}} = \frac{a}{a_0} \sigma_{\text{liq}}$, where $a_0$ is the zero-temperature FCC equilibrium lattice spacing equal to 1.5424 $\sigma_{\text{liq}}$. With such a model, wetting properties as defined by the contact angle are driven by the ratio between the liquid/liquid and liquid/solid attractions. In practice, we varied $\epsilon_{\text{wall}}$ while keeping the liquid properties constant and measured the corresponding contact angle, $\theta$. We note that additional complexities which also influence the surface solvophobicity including functionalization and polarity effects cannot be captured with our present model.\textsuperscript{36}

Finally, the results are shown in physical units. For water, energy is displayed in kcal/mol and distances are shown in angstroms. When computing $a$, we used $\sigma_{\text{liq}} = 2.75$ Å as it is the approximate size of a water molecule. For LJ, the potential parameters are denoted $\epsilon_{\text{LJ}}$ and $\sigma_{\text{LJ}}$. We worked at a temperature of $k_B T = 0.8 \epsilon_{\text{LJ}}$ and we chose $k_B T = 0.593$ kcal/mol at 300 K in order to rescale $\epsilon_{\text{LJ}}$ to real units. Concerning the distances, we imposed $\sigma_{\text{LJ}} = 2.75$ Å as well.

B. Molecular dynamics simulation of water

The extended simple point charge (SPC/E) model is used for water.\textsuperscript{62} Bonds in water molecules are constrained using the SHAKE algorithm, and long-range Coulombic interactions are computed with the Particle-Particle-Particle-Mesh solver with a precision tolerance equal to $10^{-4}$ and a real space cutoff equal to 9.8 Å. At the initialization step, water molecules are arranged on a simple cubic lattice structure with a lattice spacing equal to 3.1 Å. Solids are modeled with rigid molecules made of three face-centered cubic layers. Solid molecules and oxygen atoms of water interact via a truncated and shifted Lennard-Jones (LJ) potential with a cutoff equal to 9.8 Å. The Large-scale Atomic/Molecular Massively Parallel Simulator (LAMMPS) package\textsuperscript{63} is used for all the simulations. From there, two types of calculations are performed: (i) droplet equilibration to measure the wetting properties of the solid and (ii) solvent-mediated forces between two slabs.

1. Droplet equilibration and contact angle

A hemisphere of water with a radius equal to 50 Å is initially deposited onto the solid surface. On top of the spherical cap, a cage made of fixed atoms is also placed to help the droplet equilibration and prevent it from leaving the solid surface at the initialization stage. These atoms only interact with oxygen atoms via a LJ potential ($\epsilon_{\text{cage}} = 1$ kcal/mol and $\sigma_{\text{cage}} = 3.166$ Å). The entire simulation box measures 300 Å × 300 Å × 200 Å which is large enough to avoid the influence of periodic images. For the equilibration protocol, the time step is set equal to 0.5 fs. NVE simulations are performed during 5 ps; then, NVT simulations are performed during another 5 ps at 300 K. From there, cage atoms are removed to allow for droplet shape relaxation and the time step is changed to 2 fs. After an equilibration run during 500 ps, snapshots are taken every 2.5 ps during 500 ps. The density profiles of oxygen atoms are averaged through time (see Fig. 1). From the density profiles, a liquid/gas interface is obtained, as depicted in Fig. 1. A linear fit with all the points located below 8 Å is then used to compute the contact angle (see Fig. 1). Uncertainties are evaluated by the standard deviation measured every 100 ps for 5 independent runs; then, an additional factor of two is incorporated to account for error in the method for contact angle extraction (see Fig. 2). In addition, the duration of the

![FIG. 1. Water density profiles for an equilibrated water droplet on top of a wall with $\epsilon_{\text{wall}}$ equal to 0.1 kcal/mol (a), 0.2 kcal/mol (b), and 0.3 kcal/mol (c). Red dots represent the liquid-gas interface which is used to compute the contact angle, and red lines are the associated linear fit. The image size is equal to 30 Å × 60 Å.](image-url)
simulation is sufficient to reach equilibration, as observed in Fig. 4(a).

2. Calculation of the solvent-mediated interactions

Two nanoslabs which are made of 40 + 41 + 40 = 121 atoms are positioned parallel to each other. The entire simulation box measures $52 \times 52 \times 60 \,\text{Å}$ and contains 5344 water molecules (see Fig. 5). The solids are first disposed on top of each other, and NVT simulations are performed at 300 K during 60 ps with a time step equal to 2 fs. After this equilibration procedure, the solids are instantaneously moved apart by 0.25 Å with the distance measured as the difference in height between the center of mass of both slabs. For each separation denoted $z$, the system is equilibrated for 50 ps and production run is done during another 50 ps. The free energy as a function of $z$ is then given by numerical integration of the forces

$$
\Delta F(z) = \int_{-\infty}^{\infty} \frac{\partial F}{\partial z'} dz' = \int_{-\infty}^{\infty} \langle \vec{f}_1 \cdot \vec{u}_z - \vec{f}_2 \cdot \vec{u}_z \rangle dz',
$$

where $\vec{f}_1$ and $\vec{f}_2$ are the forces between water molecules and solid atoms with 1 and 2 designating respectively the upper and the lower solids. $\vec{u}_z$ is a unit vector along the $z$ direction going upward. The difference in forces used in the integration scheme Eq. (1) is shown in Fig. 3. Duration of the simulation time is considered sufficient to reach equilibration, as assessed by Fig. 4(b). The error bars in this figure are computed as the standard deviation obtained with 5 independent runs.

C. Density functional theory calculation of Lennard-Jones

For this second method, liquid particles interact via a LJ potential with $\epsilon_{LJ}$ and $\sigma_{LJ}$ as energy and length parameters, respectively. The cutoff distance is equal to $3\sigma_{LJ}$. The density and the temperature are respectively $\rho_{LJ} = 0.7 \sigma_{LJ}^{-3}$ and $k_B T = 0.8 \epsilon_{LJ}$ which is located between the triple point and the critical temperature. This corresponds to the liquid density for a chemical potential supersaturation equal to $\Delta \mu = 0.27 k_B T$. While the value of $\Delta \mu$ quantitatively influences the solvent-mediated forces, the supersaturation is chosen in this work to match the ratio of pressure between water coexistence pressure and atmospheric pressure ($P_{coex} = 1/20 P_{atm}$). Interactions between liquid molecules and solid atoms are also modeled with LJ potential truncated at $3\sigma_{LJ}$. Within the DFT framework, free energy is expressed as a functional of the liquid density. For LJ interaction, the potential is separated in two parts, the repulsive part modeled with the White Bear functional and the attractive part treated in mean field. The density is computed on a discretized three-dimensional grid with 8 lattice points per unit of $\sigma_{LJ}$ and the free energy is obtained through minimization with respect to the density field. In order to match MD calculations that are made in the NVT ensemble, DFT calculations are also run with a fixed number of particles rather than a fixed chemical potential. The DFT method is described in greater details in our previous contributions. Accuracy of the DFT treatment is discussed in Ref. 54. Droplet equilibration results were taken from our previous work. For solvent-mediated interactions, we used the same system as with molecular dynamics simulation of water except that there is no equilibration protocol and the free energy is obtained directly through DFT.

In recent studies regarding solvent mediated forces, calculations are performed in $\mu$VT or NPT ensembles in order to supply particles during the drying transition. To evaluate if our system is large enough to cope with this issue, we also ran calculations in the $\mu$VT with $\Delta \mu = 0.27 k_B T$ [see Fig. 8(b)]. The results are not significantly different from those obtained in NVT thus justifying our approach.

III. RESULTS AND DISCUSSION

Figures 5 and 6 show the typical results obtained respectively for water and Lennard-Jones. In both cases, when walls are solvophilic, the gap between the two slabs is filled with
FIG. 5. Liquid density profiles for water molecules confined between two nanoslabs obtained for $a = a_0$ at different distances and degrees of solvophobicity. The contact angles of 35° and 120° are obtained with $\epsilon_{\text{wall}}$ respectively equal to 0.42 kcal/mol and 0.10 kcal/mol. Each image is a slice of width 1 Å passing through the middle of the solute and measuring $48 \times 48$ Å.

FIG. 6. Liquid density profiles for Lennard-Jones particles confined between two nanoslabs obtained for $a = a_0$ at different distances and degrees of solvophobicity. The contact angles of 35° and 120° are obtained with $\epsilon_{\text{wall}}$ respectively equal to 0.4 $\epsilon_{\text{LJ}}$ and 0.2 $\epsilon_{\text{LJ}}$. Each image is a slice of width 0.275 Å passing through the middle of the solute and measuring $68.75 \times 68.75$ Å.

liquid even at small distances [see Figs. 5(b) and 6(b)]. For solvophobic walls, this happens only for large enough separations. In addition, structuring can be observed in the vicinity of the slabs especially when looking at the density profiles obtained by DFT of Lennard-Jones particles (see Fig. 6). The structure is more pronounced for solvophilic walls.

A. Influence of the wall lattice spacing

In this first study, we worked at a fixed value of $\epsilon_{\text{wall}}$ while changing the wall lattice spacing so that both the structure and the hydrophobicity are modified. In Figs. 7(a) and 7(b), excess free energy is plotted for a moderate value of $\epsilon_{\text{wall}}$ for which the contact angle is $\theta = 35^\circ$ [(a) and (b)]. Calculations are run at a value of $\epsilon_{\text{wall}}$ for which the contact angle is $\theta = 120^\circ$ [(a) and (b)] and at $\theta = 35^\circ$ [(c) and (d)] for $a = a_0$. The values of $\epsilon_{\text{wall}}$ are given in the captions of Figs. 5 and 6.

FIG. 7. Lattice spacing influence on the excess free energy obtained with SPC/E water [(a) and (c)] and with Lennard-Jones [(b) and (d)]. Calculations are run at a value of $\epsilon_{\text{wall}}$ for which the contact angle is $\theta = 120^\circ$ [(a) and (b)] and at $\theta = 35^\circ$ [(c) and (d)] for $a = a_0$. The values of $\epsilon_{\text{wall}}$ are given in the captions of Figs. 5 and 6.

denser and thus becomes less solvophobic. Therefore, both the range and the height of the solvent-mediated interaction are reduced. The results obtained with LJ DFT and with water MD are qualitatively similar, and we demonstrate that in this case, LJ can be used to reproduce water-mediated interactions. For solvophilic walls, the results of the comparison are not so close [see Figs. 7(c) and 7(d)]. In both systems, oscillations in the free energy are observed due to the layering of the liquid near the walls. However, the oscillation amplitudes vary significantly between water and the LJ fluid. This is likely due to the asymmetry of water molecules: as they pack together to form denser layers near the wall, their interlayer distance does not depend solely on their average size but rather on their size in some particular directions.

B. Influence of the wall energy

Solvent-mediated interactions are plotted at a fixed value of $a = a_0$ but for different values of $\epsilon_{\text{wall}}$ in Fig. 8. When comparing the results from MD water and DFT LJ, several similarities can be identified. First, when the walls are solvophobic, free energy monotonically increases as the nanoslabs are pulled apart with an almost linear behavior. Then, when the walls are solvophilic, damped oscillations are observed because of the emergence of structured layers near the wall. Also, the lowest energy state is always at contact, meaning that the nanoparticle would preferentially stay near the wall as long as it overcomes the intermediate free energy barrier. Finally, the range of the depletion force does not go beyond 20 Å which corresponds to approximately 7 liquid layers. These similarities were already raised in the literature, and our work allows for a more direct comparison as we studied the same system (i.e., two nanoslabs made of the same structure) while only changing the liquid nature.
FIG. 8. Wall energy influence on the excess free energy obtained at $a = a_0$ with SPC/E water (a) and with Lennard-Jones (b). The solvophobicity increases as coloring goes from blue to red with $\epsilon_{\text{wall}}$ going from 0.50 kcal/mol (0.65 $\epsilon_{\text{LJ}}$) to 0.05 kcal/mol (0.05 $\epsilon_{\text{LJ}}$) for water (for LJ). This corresponds to contact angle ranging from 0° to 180°. Each color is separated by 0.05 kcal/mol (by 0.1 $\epsilon_{\text{LJ}}$) for water (for LJ). The dashed lines correspond to the results obtained with LJ/DFT in the $\mu$VT ensemble using $\Delta \mu = 0.27 k_B T$.

In order to quantitatively compare the results from water MD and LJ DFT, we define the following two positions: (i) $z = 5$ Å gives $\Delta F_{\text{contact}}$ and (ii) the position, denoted $z_{\text{mid}}$, at which the most solvophilic interaction reaches its maximum is used as an intermediate value called $\Delta F_{\text{mid}}$ (see Fig. 8). In Fig. 9, the results are reported for different contact angles that are determined after the equilibration of sessile drops. For the highest degrees of solvophilicity (and solvophobicity), droplets are not stable and the contact angle is trivially 0° (and 180°). Therefore, when reporting $\Delta F_{\text{mid}}$ and $\Delta F_{\text{contact}}$ as a function of the corresponding contact angle, not all the data from Fig. 8 are considered. As the contact angle is increased, $\Delta F_{\text{mid}}$ decreases almost linearly. Water MD and LJ DFT curves have similar slopes, and the constant difference between the two curves is roughly of 24 kcal/mol. However, the sign of $\Delta F_{\text{mid}}$ is different, which indicates that qualitatively different behaviors are expected. In the water case, this intermediate distance is less energetically favorable than having the nanoslabs far from each other. For $\Delta F_{\text{contact}}$, LJ DFT and water MD also lead to qualitatively different results. Indeed, while for LJ, $\Delta F_{\text{contact}}$, like $\Delta F_{\text{mid}}$, decreases linearly, for water, $\Delta F_{\text{contact}}$ is non-monotonic and peaks around 80°. As already raised in Sec. III A, LJ is not well-adapted to model water at contact because water has orientational order especially for solvophilic walls which cannot be seen with LJ. Furthermore, while there is an intermediate range of solvophobicity ($\theta \in [70°; 110°]$) where good agreement for $\Delta F_{\text{contact}}$ is found, the signs of $\Delta F_{\text{mid}}$ are different as raised above.

C. Hysteresis in solvent-mediated forces

Throughout this study, the results on the solvent-mediated forces were obtained as the two solutes are pulled apart from each other. Yet, another possibility concerns the case when the slabs are disposed far from each other and then brought together. This leads to the question of reversibility of the interaction. In Fig. 10, the solvent mediated forces are plotted in this second approach. In the case of water MD, qualitative agreement is found when comparing the results from Fig. 8 with solvophobic walls. However, strong repulsive interactions are observed with solvophilic walls. This results from water molecules that can be trapped between the two plates if they are brought together too rapidly. Furthermore, the capillary evaporation which is not observed when the solutes are pulled apart is not only driven by the solute interdistance and additional order parameters such as the solvent density between the solutes can be used.32–34 Essentially, the time for gas to nucleate between the two walls is so large that we cannot obtain the equilibrium state with brute-force molecular dynamics simulations.32 This apparent hysteresis is not found in LJ DFT since the technique enables to circumvent any of these kinetic

FIG. 9. Comparison between the results of LJ DFT and water MD using the contact angle dependence of $\Delta F_{\text{mid}}$ (a) and $\Delta F_{\text{contact}}$ (b). In gray, the results for $\Delta F_{\text{mid}}$ obtained with LJ DFT are vertically shifted in order to show that the difference with water MD is only a constant.

FIG. 10. Solvent-mediated forces at $a = a_0$ with SPC/E water (a) and with Lennard-Jones (b) obtained as the solutes are brought closer to each other. Color designations are described in Fig. 8.
issues and directly leads to the most stable state in which the gap between the walls is emptied of liquid.

IV. CONCLUSIONS

In summary, solvent-mediated forces were measured in a very generic case of two nanoslabs embedded in a liquid. Two different models for the liquid along with two different methods for measuring free energy were employed. Such a direct comparison of these two approaches allowed us to identify both similarities and differences. On the one hand, oscillations for solvophilic walls and a linear decrease for solvophobic walls were observed with the two liquids. In addition, the range of the depletion force and the presence of a minimum at contact are two additional features that seem to support the idea of a universal behavior of the solvent-mediated forces. On the other hand, no region of solvophobicity seems to show quantitative agreement between water and LJ. In particular, amplitudes of the depletion force and the presence of a minimum at contact were observed with the two liquids. In addition, the range of energy at contact does not have the same behavior as the contact angle is changed. Ultimately, using LJ or water in order to quantifies the error made if one wishes to use LJ instead of water.

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