Orientational order as the origin of the long-range hydrophobic effect

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The long range attractive force between two hydrophobic surfaces immersed in water is observed to decrease exponentially with their separation – this distance-dependence of effective force is known as the hydrophobic force law (HFL). We explore the microscopic origin of HFL by studying distance-dependent attraction between two parallel rods immersed in 2D Mercedes Benz model of water. This model is found to exhibit a well-defined HFL. Although the phenomenon is conventionally explained by density-dependent theories, we identify orientation, rather than density, as the relevant order parameter. The range of density variation is noticeably shorter than that of orientational heterogeneity. The latter is comparable to the observed distances of hydrophobic force. At large separation, attraction between the rods arises primarily from a destructive interference among the inwardly propagating oppositely oriented heterogeneity generated in water by the two rods. As the rods are brought closer, the interference increases leading to a decrease in heterogeneity and concomitant decrease in free energy of the system, giving rise to the effective attraction. We notice formation of hexagonal ice-like structures at the onset of attractive region which suggests that metastable free energy minimum may play a role in the origin of HFL.

I. INTRODUCTION

When water is confined between two large hydrophobic surfaces, one finds the emergence of a surprisingly long ranged effective attraction between the two hydrophobes. The attractive force grows exponentially as the distance between the surfaces is lowered. This exponential separation dependence of hydrophobic attraction has become widely known as the hydrophobic force law.

The first direct measurement of the interaction between two cylindrically curved hydrophobic surfaces in aqueous electrolyte solutions was performed by Israelachvili in a series of landmark experiments using specially designed surface force measurement apparatus. [1-5] These experiments observed that the attractive force between the surfaces is “long-ranged” – being detectable even as far as 80 Å to 100 Å separation. It is neither sensitive to the type and concentration of the electrolyte, nor to the pH of the solution. [5] The hydrophobic force \( F \) measured in experiments can be empirically fitted to the following form

\[
\frac{F}{R} = C \exp \left( -\frac{d}{\xi} \right) \tag{1}
\]

where \( R \) is the radius of the cylindrical curve, \( C \) is an arbitrary constant and \( \xi \) is the correlation length (or, decay length). Accurate estimate of the range of this hydrophobic force went through a period of uncertainty. Different experimental force-measuring techniques and different methods of hydrophobization apparently recognized substantial diversity in length scales. There are reports of a measurable attractive force at separation as large as 3000 Å. [6, 7] However, recent experiments have led to believe that the attraction at separations greater than 200 Å is due to a variety of system-dependent, nonhydrophobic (or only indirectly hydrophobic-dependent) effects. [8, 9]

The attractive forces at separations less than 100 Å represent the truly hydrophobic interaction. The exponential decay of the attractive force between hydrophobic surfaces (or, the hydrophobic force law), however, remains an integral feature of all experimental results.

Despite a great deal of research over the last three decades, understanding of certain aspects of the origin and nature of interaction between hydrophobic surfaces have remained unsatisfactory. Theoretical studies have shown that size of the hydrophobic surface, degree of hydrophobicity and temperature together determine the critical separation [10, 11] as well as the rate of evaporation of confined water. [12] The phenomenon is mainly attributed to the formation of a liquid-vapor interface induced by the surface, leading to cavitation at some critical intersurface separation. [13] Such cavitation is also observed near vapor-liquid coexistence for a Lennard-Jones fluid confined between hard walls. [14] This de-wetting induced collapse is generally described by density fields. In this context, Lum-Chandler-Weeks (LCW) theory [15] has been one of the most successful approaches. It involves the drying interface concept, and includes effects of orientation of the water molecules only implicitly in the Hamiltonian using prior knowledge of the radial distribution function. LCW theory predicts a critical intersurface separation of 50 Å for two parallel hard plates.

The above picture basically describes water in terms of inward propagating density inhomogeneity originating from the two surfaces. However, orientational ordering induced by the surfaces is ignored. Indeed, as early as in 1959, Kauzmann introduced the concept of “clathrate-structure” of water surrounding the hydrophobic solutes to understand the hydrophobic effect in terms of entropic order parameter. The range of density variation is noticeably shorter than that of orientational heterogeneity. The latter is comparable to the observed distances of hydrophobic force. At large separation, attraction between the rods arises primarily from a destructive interference among the inwardly propagating oppositely oriented heterogeneity generated in water by the two rods. As the rods are brought closer, the interference increases leading to a decrease in heterogeneity and concomitant decrease in free energy of the system, giving rise to the effective attraction. We notice formation of hexagonal ice-like structures at the onset of attractive region which suggests that metastable free energy minimum may play a role in the origin of HFL.

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stability. [16] Formation of such structures emphasizes the role of orientational ordering in water. Subsequent computer simulations have also found evidence of significant orientational ordering among water molecules around hydrophobic residues of protein molecules. [17] In a recent study, Berry and coworkers have discussed how oppositely directed polarization from water dipoles can give rise to a long range attraction. [18]

In view of the above, the one-order-parameter (density) description of Lum et al. may not be adequate to describe ordering of water around a hydrophobic surface. Hydrophobic surfaces can induce orientational inhomogeneity by disrupting the natural orientational correlation among water molecules. This orientational ordering in turn may alter many fundamental properties of water, including density. It is thus expected to play a key role in describing the metastable or unstable state, if any, between the two hydrophobic surfaces. Comprehensive understanding of the long range hydrophobic attraction is a challenging task, particularly because of subtle nature of the hydrophobic effect itself and its interplay with an array of other forces.

Computational approaches have proved to be very useful in this regard as one can explore the microscopic details and try to understand their manifestation in the macroscopic behavior. Accurate and detailed models evidently gives improved accuracy in computer simulations. However, simplified models often give insights that are not obtainable at relevant time scales from complex models. They usually capture the essence of underlying physics while being more comprehensible – providing insights and illuminating concepts, and they do not require big computer resources.

In the present work, we employ the Mercedes-Benz (MB) model of water to study hydrophobic force law. This model was introduced by Ben-Naim [19, 20], and later parameterized by Dill and co-workers to mimic micelle-like properties. [21–25] The merit of this model lies in the simple form of the interaction potential and the reduced dimensionality. A schematic representation of the model is shown in Fig. 1 and details are given in Materials and Methods. Several previous studies have demonstrated that MB model generally reproduces most of the properties of water including the density anomaly, the minimum in isothermal compressibility as a function of temperature, and the thermodynamic properties of non-polar solvation. This model has also been used to study the thermodynamic properties and structural aspects of confined water. [26–29] To study the hydrophobic force, we have introduced rigid parallel hydrophobic rods made of non-interacting 2D Lennard-Jones (LJ) circles in MB water. We have performed molecular dynamics (MD) simulations in NVT ensemble to investigate the structure and dynamics of the system.

We find that this model system exhibits a clear attractive hydrophobic force law, with a correlation length $\xi$ of $\sim 4\sigma$, where $\sigma$ is the diameter of the MB circle. The force is found to decay to 90% of the contact value at around $\sim 10\sigma$. Our results and analysis provide a molecular level explanation of the origin of long-range attractive forces between the two hydrophobic surfaces, prior to the eventual cavitation induced collapse. We introduced and calculated appropriate orientational order parameters. We find that the correlation length of density heterogeneity is significantly shorter than that of the bond orientational order parameter, while the latter is comparable to the correlation length of the hydrophobic force. Destructive interference between orientational heterogeneities propagating inwards from the two surfaces lead to a lowering of imposed order, and hence lowering of free energy. As this destructive interference increases with decreasing separation between the two rods, there appears an effective attraction between the two surfaces.

**II. HYDROPHOBIC FORCE LAW IN MB WATER**

The hydrophobic force law has been inspected in computer simulations studies of 3D water models, but the validity of the same in reduced dimension, such as in the MB model, has not been looked at. Recently, Dill and coworkers have studied the behavior of confined MB water in enclosed cavities using NVT and $\mu$VT Monte Carlo simulation. [26] The force between the walls was estimated from the oscillating density profiles and was found to be attractive. The exponential distance-dependence and the correlation length were, however, not examined. In this work we calculate the pressure $P_{\text{cav}}$ in the confined region between the two hydrophobic rods by using the virial expression,

$$P_{\text{cav}} = \frac{Nk_BT}{V} + \frac{\sum_i r_{ij} \cdot f_i}{DV}$$  \hspace{1cm} (2)
Figure 2. Pressure $P$ on each individual hydrophobic LJ rods suspended in MB water increases exponentially as the distance between the rods ($d$) is increased. Simulation results are shown for different rod lengths, given by the number of LJ particles ($n_{rod}$) constituting the rod. The solid lines are the exponential fittings [Eq. 5] with global correlation length $\xi = 3.8 \pm 0.1$.

The pressure is found to increase exponentially with the distance of separation between the two rods. We obtain the pressure at infinite separation $P_\infty$ by fitting it to the equation,

$$P_{cav} = P_\infty + A \exp \left( -\frac{d}{B} \right)$$

where $d$ is the inter-rod distance, $A$ and $B$ are fitting parameters. The effective pressure on the rods is subsequently obtained as

$$P = P_{cav} - P_\infty$$

In Fig. 2, we show the effective pressure on the rods as a function of inter-rod separation distance $d$. The length of the rod is given by the number of LJ particles ($n_{rod}$) constituting the rod. The effective pressure shows negligible dependence on the rod length, and decays exponentially with inter-rod distance. This is consistent with the experiments of Israelachvili et al. [2, 5] and can be fitted to the equation

$$P = C \exp \left( -\frac{d}{\xi} \right)$$

where $N$ is the number of atoms in the system, $k_B$ is the Boltzmann constant, $T$ is the temperature, $D$ is the dimensionality of the system, $V$ is the system volume, $\mathbf{r}_i$ is the position vector of $i^{th}$ particle, and $f_i$ is the total internal force (arising from the interatomic interactions) on $i^{th}$ particle. The second term in the above Eq. 2 is the virial.

The negative pressure indicates effective attraction between the rods, induced by the intervening water. As noted by the dashed lines in Fig. 2, the pressure in the confined region reaches 90% of the bulk at $d \approx 10$. Thus, there exists a tangible finite force of attraction between the two rods even when they are separated by 10 water diameters.

Table I. Fitted values of the correlation length and prefactor of the hydrophobic force law (see Eq. 5) observed between LJ rods in a system of MB particles

| $n_{rod}$ | $C$ | $\xi$ |
|----------|-----|------|
| 5        | $-0.21 \pm 0.01$ | $3.8 \pm 0.1$ |
| 10       | $-0.25 \pm 0.01$ |
| 15       | $-0.26 \pm 0.01$ |
| 20       | $-0.27 \pm 0.01$ |

which is analogous to the hydrophobic force law Eq. 1. The solid lines in Fig. 2 show the global fitting with constant $\xi$. The correlation length is found to be $\xi = 3.8 \pm 0.1$.

III. MICROSCOPIC ORIGIN OF THE HYDROPHOBIC FORCE: ORIENTATION AND DENSITY MAPS

To understand the origin of long range hydrophobic force, we seek to visualize the underlying control parameters – position-dependent density and orientation – of MB water in the region confined by the two parallel hydrophobic rods.

We map the orientation of MB particles at different inter-rod separations in the left panel of Fig. 3 and compare with the corresponding density maps in the right panel. The angles are measured from the $z$-axis, i.e. the axis normal to the rods. The angle zero represents an arm pointing away from the left rod. Conversely, the angle 180° represents an arm pointing away from the right rod. The MB particles (with 3 arms) near the wall prefer a particular orientation, with one of the arms pointing towards the wall. Thus, particles near the left wall prefers 60°, 180° and 300° angles, while particles near the right wall prefers 0°, 120° and 240°. It means that the particles near the left wall has a dangling, unsatisfied H-bond pointing towards the wall, and vice versa. The preference is due to
Figure 3. Orientation maps (left panel) and density maps (right panel) of MB water in the region confined between the two hydrophobic rods, at different inter-rod separation distance \( (d) \). Here, \( \theta \) is the angle made by the arms of the MB particles with the \( z \)-axis, i.e. the axis normal to the rods. The left rod is at \( z = 0 \). The \( y \)-axis is along the length of the rods, with the bottom of the rods at \( y = 0 \). The orientation maps show the probability distribution of \( \theta \) along \( z \). The density maps show the distribution of density in the \( y - z \) plane.

The H-bond energy gained by the particles from forming “good” H-bonds with their remaining two arms. Interestingly, this constraint induces an orientational pattern in the system. Oppositely oriented patterns are created by the two hydrophobic walls. The inward propagation of the opposing orientational heterogeneity causes an interference as the opposite “waves” meet near the center of the confined region. This results in a frustration of orientation near the center, and the orientational pattern survives for large inter-rod distances – even when the rods are separated by 10 water diameters.

The density maps of MB water in the enclosed regions are plotted alongside in the right panel of Fig. 3 for comparison. Uniformity along the \( y \)-axis ensure that the orientational patterns of the MB particles are in dynamical equilibrium. Visual inspection shows that oscillations in density at the center of the inter-rod region becomes bulk-like at \( d = 8.0 \), while the induced orientational pattern exists even at \( d = 10.0 \) and beyond. The emergence of such exotic patterns both in orientation and density in such simple systems is remarkable and fascinating. They provide a new insight to the origin of long range hydrophobic force.

IV. DISTANCE DEPENDENCE OF DENSITY AND ORIENTATIONAL INHOMOGENEITY: QUANTITATIVE ANALYSIS

One primarily employs density-dependent descriptions, such as the LCW theory to understand hydrophobic force law. However, as noted in the introduction, this one-order-parameter description might not be enough to describe a complex liquid like water, especially due to the decrease in stability of the liquid confined within the hydrophobic surfaces. Orientational inhomogeneity may propagate longer distances than density variation, as we have visualised in the preceding section. Here, we obtain and compare the range of density and orientational inhomogeneity in terms of correlation lengths of the corresponding order parameters.

A. Length scale and amplitude of density heterogeneity

Density profiles of MB particles within an enclosed cavity were studied by Dill et al. [26]. In the present work, the particles located in between the rods are in dynamic equilibrium with the bulk. Here, we calculate the distance-dependent density within the cavity using the same standard method,

\[
\rho_{\text{avg}}(z) = \frac{N}{Lh} \tag{6}
\]

where \( N \) is the average number of particles in an interval of width \( h \) at distance \( z \) from the left rod and \( L \) is the length of the rods \([L = (n_{\text{rod}} - 1)\sigma_{LJ}]\). The density profile
within the cavity is normalized by the density of the bulk \( \rho_{\text{bulk}} \). In Fig. 4, we show the computed profiles of water density \( \rho_{cav}(z)/\rho_{\text{bulk}} \) between the two parallel rods, at different inter-rod separations \( d \). The profiles are consistent with the earlier work of Dill et al. [26] The results show that the water molecules are structured into multiple layers between the two rods, and hence the density oscillates along the \( z \)-direction. We note that similar results were also obtained for three dimensional water models, studied by Choudhury and Pettitt. [10]

The density \( \rho_{cav} \) of MB water in the region enclosed by the two parallel hydrophobic rods gives the scalar density order parameter. It is obtained by time averaging over the simulation trajectory, and is normalized by the bulk density \( \rho_{\text{bulk}} \) of MB water. In Fig. 5 we show the variation of \( \rho_{cav}/\rho_{\text{bulk}} \) as a function of the inter-rod separation \( d \), while each rod consists of 15 LJ particles \( (n_{\text{rod}} = 15) \). The results from simulation are shown by solid squares. The variation is found to be exponential, and can be fitted to the equation

\[
\rho_{cav}/\rho_{\text{bulk}} = 1 + A \exp\left(-\frac{d}{\zeta}\right)
\]

where \( \zeta \) is the correlation length of the scalar density order parameter. The fitting is shown by the solid line in Fig. 5. We obtain \( A = -1.86 \) and \( \zeta = 2.4 \). Thus, the correlation length for density is less than the correlation length of the force between the two rods.

A useful measure to quantify the effect of the walls on the water structure is to find the separation at which the density reaches 90% of the bulk density. This is achieved at \( d \approx 8.0 \) in the present case, as shown by the dashed lines in Fig. 5. Thus, the density of MB water in the confined region saturates to bulk density even though there persists a significant finite force of attraction between the rods. While the exponential decay of density may be correlated to the exponential decay of the force at short length scales, the origin of the attractive force beyond a certain length scale (where cavity density saturates to bulk density) cannot be correlated with the density order parameter.

B. Length scale and amplitude of orientational heterogeneity

In order to quantify the extent of orientational order present in the confined region, one must define an orientational order parameter. As we have already noted, orientations of the MB particles are opposite to each other...
near the two walls. Hence they induce opposing patterns, which propagate towards the center. Towards this goal, we introduce the orientational order parameter \( Q = \langle \cos 3\theta \rangle \) to quantify the orientational order propagation. Here \( \theta \) is the angle made by the arms of the MB particles, relative to the \( z \)-axis, and \( \langle ... \rangle \) denotes the averaging. The value of \( \langle \cos 3\theta \rangle \) can range from \(-1\), when all particles are perfectly ordered with arms pointing away from the left wall, to \(+1\) when all particles are perfectly ordered with arms pointing away from the right wall.

The order parameter \( \langle \cos 3\theta \rangle \) has been calculated for MB water within small intervals of width \( h \) at a distance \( z \) from the left rod. The distance-dependent propagation of \( Q \) is shown in Fig. 6 at different inter-rod separations. We see interference of the orientational order induced by the two rods. Strong oscillations are observed in the central region. Adjacent positive and negative values indicate reversal of orientations, due to interference among the inwardly propagating oppositely oriented heterogeneity. It is interesting to note that fluctuations in \( Q \) persist for longer distance than that in density profile.

C. Decay of global orientational order

A global order parameter within the cavity could be helpful for comparison with the scalar density order parameter and formulation of a phenomenological theory. Here we use the six-fold bond orientational order \( \psi_6 \), which is defined as

\[
\psi_6 = \frac{1}{N} \left| \sum_{k=1}^{N} \left( \frac{1}{N_{nk}} \sum_{j=1}^{N_{nk}} e^{i6\theta_{kj}} \right) \right| \tag{8}
\]

where \( N_{nk} \) is the number of nearest neighbors of a tagged particle, \( \theta_{kj} \) is the angle of the bond that the \( j \)-th neighbor makes with the tagged particle and \( N \) is the total number of MB particles in the confined region. We had earlier shown that \( \psi_6 \) is an effective order parameter to distinguish the honeycomb solid and liquid state of the MB liquid [30].

In Fig. 7, we show the distributions of \( \psi_6 \) in the confined region, at different inter-rod separations and compare it with that of bulk MB liquid. The long tails of the distributions reveal the ordering in the confined region. While bulk-like behavior is approached with increasing inter-rod separation, it is interesting to note the significant ordering in the confined region even at \( d = 20.0 \). In Fig. 8 we show the variation of \( \psi_6 \) as function of inter-rod separation \( d \). The results from simulation are shown by solid squares.

\[
\psi_6 - \psi_\infty = A \exp \left( -\frac{d}{\gamma} \right) \tag{9}
\]

where \( \gamma \) is the correlation length of the orientational order. The fitting is shown as a solid line in Fig. 8. We obtain, \( A = 0.5, \gamma = 3.4 \) and \( \psi_\infty = 0.04 \). It is interesting to note that the correlation length for orientational order is much longer ranged than the density order (\( \zeta = 2.4 \)), and comparable to that of the hydrophobic force (\( \xi = 3.8 \)).

D. Spatio-temporal correlation of orientational relaxation

The orientational relaxation timescale provides further insight to the dynamics of orientation. The orientational relaxation of the MB particles is characterized by the ori-
entational correlation function $C(t)$,

$$C(t) = \left\langle \sum_{i=0}^{N} \cos(\theta_i(t) - \theta_i(0)) \right\rangle$$  \hspace{1cm} (10)$$

where $\theta(t)$ is the angle made by the arms of the MB particles, relative to the $z$-axis at time $t$, $N$ is the number of particles in the region considered, and $\langle \ldots \rangle$ denotes the ensemble average. We calculate $C(t)$ for the MB particles within small intervals of width $h$ at a distance $z$ from the left rod. We have considered time average instead of the ensemble average, since they are equivalent because of ergodicity. The orientational relaxation is found to depend on $z$. We fit it to a biexponential form

$$C(t) = a_1 \exp\left(-\frac{t}{\tau_1}\right) + (1-a_1) \exp\left(-\frac{t}{\tau_2}\right)$$  \hspace{1cm} (11)$$

where the time constants for the decay of the orientational correlation are given by $\tau_1$ and $\tau_2$, with amplitudes $a_1$ and $(1-a_1)$ respectively. The average relaxation time $\langle \tau \rangle$ is subsequently obtained as $\langle \tau \rangle = a_1 \tau_1 + a_2 \tau_2$ where $a_2 = (1-a_1)$.

Figure 7. Probability distribution of six-fold bond orientational order $\psi_6$ of MB liquid confined between the two hydrophobic plates. The distribution at different inter-rod separation $d$ are color coded, as shown in the legend. The long tail of the distributions of confined MB liquid, as compared to the distribution in bulk, indicate occurrence of ordered liquid within the cavity.

Figure 8. Average six-fold bond orientational order $\psi_6$ of MB water in the region between the two parallel hydrophobic rods, as a function of distance $d$ between the two rods. The solid squares are the simulation results. The solid line is the fitting to Eq. 9, showing that the correlation length $\gamma = 3.4$ propagates longer distance than density order parameter, and is comparable to that of the hydrophobic force (see Fig. 2).

Figure 9. Plots of average orientational relaxation time $\langle \tau \rangle$ of MB water along the $z$-axis of separation of the two parallel hydrophobic rods at different inter-rod separations $d$. Results from simulations are the points, whereas the solid lines are merely visual aid.
surface, while it reduces as we approach the bulk-like liquid near center of the confined region. For large inter rod separations, the average relaxation time saturates to the bulk value of $\tau_{\text{bulk}} = 11.3$ near the center.

V. FREE ENERGY SURFACE FOR CONFINEMENT INDUCED TRANSITION: POSSIBLE ROLE OF A METASTABLE PHASE

As the density of the liquid in the confined region decreases below the bulk value and ice-like orientational correlation increases (see Fig. 6–9), additional contribution to the stability of these low density molecular arrangements may come from the presence of nearby metastable phases. [31, 32] In the present problem, the low density metastable phase is the hexagonal 2D ice (honeycomb). For the 3D water, nearby metastable phases are ice 1h and the low density liquid(LDL)-like arrangement. For MB particles, there is an additional metastable phase which is oblique phase. [30, 31, 33] However, oblique phase is quite far off in the parameter space and we do not see any evidence of oblique phase. However, we see strong evidence of the presence of ice-like configurations (see Fig. 10). As discussed elsewhere [31], metastable minima in free energy can lower the surface energy and thereby play an important but “hidden” role in the stability of the confined liquid, and hence provide additional attraction at some separation.

As mentioned earlier, existing theories of hydrophobic force are based primarily on number density as the sole order parameter that varies across the system as we travel from one surface to another. As a result, they can miss important physical constraints coming explicitly from orientation of the water molecules, such as the clathrate-like structures with pronounced orientational ordering. [16, 17, 34] It becomes particularly important when the separation is somewhat large, may be of the order of 8–10 molecular diameters wide in the present MB system. At such separations, the density inhomogeneity becomes rather small but orientational heterogeneity may still be significant, as we indeed find in our simulations. This suggests that orientational heterogeneity is important and needs to be taken into account. The crucial role of orientation was earlier pointed out by Tanaka [35], in case of crystallization, quasi-crystal formation, glass transition, etc. in 3D liquids. In fact, for water-like systems, orientation could be more important of the two as density is slaved to orientation imposed by H-bonding.

Physical picture becomes more interesting as we move to larger $d$ due to enhanced population of 6-membered rings. In Fig. 10, we show some representative snapshots of the simulations at different inter-rod separations. We find an increase in the presence of hexagonal ice-like rings at intermediate separation distances ($d = 8–12$) between the rods. As already discussed, this corroborates with the argument that the system experiences the metastable free energy minimum due to the ice phase. Clearly, such a minimum can help lowering surface free energy if the system permits orientational arrangement close to ice. Cooperativity of water molecules induced by the hydrophobic surface has been observed in 3D water models as well. [17, 34]

In Fig. 11 we show the calculated free energy surface corresponding to the melting of MB system at $P = 0.19$ and $T = 0.15$. The minimum at high $\rho$ and low $\psi_6$ corresponds to the liquid phase, whereas the honeycomb phase (akin to low density ice) creates another minimum at lower $\rho$ and higher $\psi_6$. The increase in orientational order and decrease in density, as observed in our simulations and visualized in the snaps (Fig. 10) may correspond to this metastable minimum. While LCW theory considers the cavitation and gas-liquid transition for collapse of large hydrophobic surfaces, this metastable minimum should be considered for longer ranged attraction.

The above conclusions are substantiated by the sharp rise in fluctuation of density and orientational order as shown in Fig. 12. We show the variance of density and orientation order parameters at different inter-separation distances in Fig. 12a. The crossover at $d = 4.0$ suggests increased fluctuation and a corresponding free energy minimum, which might be due to metastable hexagonal phase. Another interesting way to explore the underlying free energy is to study the susceptibility for the different prop-
Microscopic analysis of the structure and orientation of water. The simplicity of the present model and the crudity of interactions experienced by real water molecules, it also crucial to phenomena such as mineral flotation, wetting, coagulation, and many processes that involve surfactant aggregation, e.g., detergency. Deciphering the hydrophobic interactions can also have profound implications for drug design and delivery to specific target proteins in a cell. [40]

VI. CONCLUDING REMARKS

Long range hydrophobic attraction plays a key role in several biological self-organizing processes, such as protein folding, assembly of membrane structures, ligand binding to hydrophobic patches on target proteins, micellar aggregation, lipid bilayer formation, etc. [36–39] It is also crucial to phenomena such as mineral flotation, wetting, coagulation, and many processes that involve surfactant aggregation, e.g., detergency. Deciphering the hydrophobic interactions can also have profound implications for drug design and delivery to specific target proteins in a cell. [40]

The present work demonstrates that a significant, attractive force can exist between two hydrophobic rods in a simple 2D molecular liquid. The system exhibits an exponential increase in the attractive force as the two rods are brought closer together. In addition, even the estimate of the range of the force is in rough agreement with experimentally observed range.

Although the MB model does not possess the complexity of interactions experienced by real water molecules, it mimics many of the important anomalies and properties of water. The simplicity of the present model and the ease of visualization can therefore be harnessed to gain important insights needed to understand the puzzling (and often controversial) aspects of water. These are sometimes obscured by the complexity of the 3D models.

Microscopic analysis of the structure and orientation of the system reveals several important issues. The length
scale of density fluctuations is found to be shorter in range than the length scale of hydrophobic force between the rods. In contrast, orientational heterogeneity propagates to longer lengths with a correlation length that compares well with that of the hydrophobic force law. Hydrophobic force law is found to be a consequence of the minimum frustration of the extended H-bond network – the frustration being imposed by the hydrophobic surface. Destru
tive interference between the orientational heterogeneity induced by the two surfaces creates a metastable ice-like phase in the central region, thereby lowering the free energy of the system.

With the insight gained from the MB model, it would be interesting to see an extension of this work to 3D water models with specific attention to orientation of the water molecules, which has largely been overlooked in past investigations. A detailed theoretical analysis with the orientational order parameter may be rewarding, and could complement our present understanding. On a slightly different note, pattern formation in nature has been a fascinating and demanding field of research and the interfering orientational patterns appearing in this system (see Fig. 3) could be an excellent model to study. Role of orientation in the hydrophobic force law indicates that such interference as observed here may play an important role in directing protein association and self-assembly where attraction between extended hydrophobic surfaces is believed to be involved.

MATERIALS AND METHODS

A. Mercedes Benz model

In the Mercedes-Benz (MB) model, water molecules are represented as two dimensional disks with three symmetrically arranged arms, separated by an angle of 120°, as in the Mercedes Benz logo. Each molecule interacts with other through an isotropic Lennard-Jones (LJ) and an anisotropic hydrogen-bond (HB) term in a completely separable form as

\[ U(\mathbf{X}_i, \mathbf{X}_j) = U_{\text{LJ}}(r_{ij}) + U_{\text{HB}}(\mathbf{X}_i, \mathbf{X}_j) \]  

(14)

where \( \mathbf{X}_i \) denotes vectors representing both the coordinates and orientation of the \( i \)th particle and \( r_{ij} \) the distance between the centers of two particles. The notations are summarized in Fig. 1. The LJ term \( U_{\text{LJ}} \) is defined as

\[ U_{\text{LJ}}(r_{ij}) = 4\epsilon_{\text{LJ}} \left[ \left( \frac{\sigma_{\text{LJ}}}{r_{ij}} \right)^{12} - \left( \frac{\sigma_{\text{LJ}}}{r_{ij}} \right)^6 \right] \]  

(15)

where \( \epsilon_{\text{LJ}} \) and \( \sigma_{\text{LJ}} \) are the well-depth and diameter for the isotropic LJ interaction. The anisotropic HB part of potential, \( U_{\text{HB}} \) is defined as

\[ U_{\text{HB}}(\mathbf{X}_i, \mathbf{X}_j) = \epsilon_{\text{HB}} G(r_{ij} - r_{\text{HB}}) \sum_{k,l=1}^{3} \left( \mathbf{i}_k \cdot \hat{u}_{ij} - 1 \right) G(\mathbf{j}_l \cdot \hat{u}_{ij} + 1) \]  

(16)

where \( G(x) = \exp(-x^2/2\sigma^2) \). This form of the anisotropic potential ensures that neighboring water molecules form explicit H-bonds (have favorable energy) when the arm of one molecule aligns with the arm of another – the strongest bond occurs when they are perfectly collinear and are opposite in direction. The unit vector \( \mathbf{i}_k \) represents the \( k \)th arm of the \( i \)th particle (\( k = 1, 2, 3 \)) and \( \hat{u}_{ij} \) is the unit vector joining the center of particle \( i \) to the center of particle \( j \). We have used the optimal parameters of the MB model that are known to reproduce the properties of water, \( \epsilon_{\text{HB}} = -1.0, r_{\text{HB}} = 1.0, \epsilon_{\text{LJ}} = 0.1, \sigma_{\text{LJ}} = 0.7 \) and \( \sigma_{\text{HB}} = 0.085 \).

B. Simulation details

In this study, we use MB model of water. The rigid hydrophobic rods are made up of non-interacting 2D Lennard-Jones (LJ) particles. The LJ particles were separated by a distance of 0.7\( \sigma_{\text{HB}} \). For the interaction between the LJ particles and MB particles, we considered the same parameters as used by Debenedetti et al. [12] in their 3D simulation, but in reduced units, \( \epsilon_{\text{LJ-MB}} = 0.006 \) and \( \sigma_{\text{LJ-MB}} = 1.2 \). We incorporated the 2D MB force field in LAMMPS for carrying out Molecular Dynamics (MD) simulation. The forces and torques were calculated using similar techniques as used earlier [41] for the 3D model.

We generate an initial configuration of 2160 MB particles in a honeycomb lattice. We perform equilibration at temperature, \( T = 0.2 \) and pressure \( P = 0.19 \) to obtain a liquid configuration. We insert two rods at required distance by replacing any MB particles, if within contact distance. The system was equilibrated for 10\(^5\) steps at constant temperature and volume (NVT ensemble), with each time step \( \tau = 0.007 \). The production run was carried out for 10\(^7\) steps at constant temperature and volume. We used Nose-Hoover thermostat to keep the temperature bath at \( T = 0.2 \).

ACKNOWLEDGEMENTS

We thank Prof. P. G. Wolynes for his valuable comments on the manuscript and insight to the problem. BB thanks Prof. I. Ohmine for a useful discussion. This work was supported in parts by grants from BRNS (Mumbai) and DST (Delhi). We thank DST for a JC Bose Fellowship (to BB).

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