Electric field inside a “Rossky cavity” in uniformly polarized water

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Electric field produced inside a solute by a uniformly polarized liquid is strongly affected by dipolar polarization of the liquid at the interface. We show, by numerical simulations, that the electric “cavity” field inside a hydrated non-polar solute does not follow the predictions of standard Maxwell’s electrostatics of dielectrics. Instead, the field inside the solute tends, with increasing solute size, to the limit predicted by the Lorentz virtual cavity. The standard paradigm fails because of its reliance on the surface charge density at the dielectric interface determined by the boundary conditions of the Maxwell dielectric. The interface of a polar liquid instead carries a preferential in-plane orientation of the surface dipoles thus producing virtually no surface charge. The resulting boundary conditions for electrostatic problems differ from the traditional recipes, affecting the microscopic and macroscopic fields based on them. We show that relatively small differences in cavity fields propagate into significant differences in the dielectric constant of an ideal mixture. The slope of the dielectric increment of the mixture versus the solute concentration depends strongly on which polarization scenario at the interface is realized. A much steeper slop found in the case of Lorentz interfacial polarization also implies a higher free energy penalty for polarizing such mixtures.

Keywords: Cavity field, surface polarization, hydration, polar response.

I. INTRODUCTION

When an interface is created in a dielectric, the surface dipoles change their preferential orientations relative to the dipoles in the bulk. The response of the dielectric to a weak external field is then a composite result of the response of these surface dipoles and the bulk dipoles. The question of whether the dielectric response of a material is sensitive to its surface structure is decided by the relative weights of these two contributions.

The Maxwell electrostatics of dielectrics neglects the structure of the interface and replaces the interfacial region of a finite microscopic dimension with an infinitesimally thin mathematical surface. This mathematical surface cuts through the dipoles of the medium creating a surface charge with the charge density \( \sigma_P \) (Fig. 1a). It is given by the projection of the dipolar polarization at the interface \( \mathbf{P}(r_S) \) on the outward normal \( \mathbf{n} \) to the surface bounding the dielectric, \( \sigma_P(r_S) = P_n(r_S) \).

The surface charge density produces the electric field of its own, which polarizes the dielectric to form an inhomogeneous polarization \( \mathbf{P}(r) \) near the interface. It decays to the uniform polarization field \( \mathbf{P} \), associated with a uniform external field \( \mathbf{E}_{\text{ext}} \), far from the interface. Since the spatial extent of the interface is neglected, the polarization \( \mathbf{P}(r) \) extends continuously up to the mathematical dividing surface. If the dielectric borders vacuum, the polarization field changes discontinuously from \( \mathbf{P}(r_S) \) at the dielectric side of the surface to zero at its vacuum side.

The sum of the electric field from the surface charge density and the external field is the Maxwell field \( \mathbf{E}(r) \). The Maxwell field, as well as the polarization \( \mathbf{P}(r) \), cannot be directly measured, but can be retrieved from electric fields inside cavities carved in the dielectric, as originally suggested by Thompson and Maxwell. For the simplest geometry of an empty spherical cavity, the solution of the Laplace equation with Maxwell’s boundary conditions gives the field at the cavity center (cavity field) \( E_M \)

\[
E_M = \frac{3}{2\epsilon + 1} E_{\text{ext}}, \tag{1}
\]

where \( \epsilon \) is the dielectric constant of the medium surrounding the cavity. The qualitative prediction of this result is that external field \( E_{\text{ext}} \) is diminished by a factor of \( 3/(2\epsilon) \) inside cavities created in highly polar dielectrics such as water. This scenario then describes a well-defined physical setup testable by laboratory or numerical experiment.

One wonders to what extent the mathematical formalism of Maxwell’s electrostatics applies to interfaces of polar liquids. The interface is obviously not a mathematical surface, but has a finite width (Laplace vs Poisson definition of the interface). In addition, liquid dipoles have the freedom to rotate and adjust to the lack of molecular interactions from the cavity side of the interface. For polar liquids, this restructuring results in preferential in-plane orientation of the liquid dipoles at free planar and closed interfaces. Unless the external field orients the dipoles off-plane, such orientational structure eliminates the surface charge since \( \sigma_P = P_n \approx 0 \) (Fig. 1b). The standard boundary conditions of Maxwell’s electrostatics do not apply, thus affecting the observable cavity field.

Water presents a particularly important test case for understanding the interfacial electrostatics of polar liquids. Strong hydrogen bonds between surface waters can

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homogeneous polarization in the case of Maxwell’s electrostatics (see below). The field accumulated by this uniform polarization results in the Lorentz field at the cavity center

$$E_L = \frac{\epsilon + 2}{3\epsilon} E_{\text{ext}}. \quad (2)$$

The main qualitative difference between this result and the standard cavity field given by Eq. 1 is that the Lorentz field does not decay to zero at $\epsilon \rightarrow \infty$ and instead levels off at a non-zero value of $E_L/E_{\text{ext}} \rightarrow 1/3$.

The model dipolar system studied previously by us has in principle shown that interfacial orientational structure of a polar liquid might be closer to the limiting scenario sketched in Fig. 1 than to the standard, solid-like picture sketched in Fig. 1b. The question of practicality of this observation remains however open. A vacuum hard-sphere cavity is a purely theoretical construct. Nevertheless, the scenario arising from this model can potentially describe non-polar particles solvated in real polar liquids.

In order to approach this more realistic situation, we study here the scenario that we have dubbed the “Rossky cavity”. The actual system is a Lennard-Jones (LJ) solute inserted in SPC/E water. As was originally shown by Rossky and co-workers and supported by many subsequent studies, water dipoles orient in-plane at interfaces with non-polar solutes. Although a non-polar solute studied here is certainly not a cavity, the orientational water structure is highly resilient to external perturbations and remains nearly intact for a broad range of solute-solvent attractions. The term “cavity” is then used to stress that it is a large energy of the water hydration $\Delta H \simeq 4 - 5 k_B T$ per bond for SPC/E water) that supports the interfacial orientational structure. Therefore, in contrast to the virtual Lorentz cavity, this physical cavity, like the traditional Maxwell cavity, is meant to represent realistic measurements of local fields inside dielectrics, in water in the present study. The Rossky cavity is meant to represent physical situations when the interfacial structure is dominated by in-plane orientations of water dipoles. We show that the boundary conditions imposed by molecular interfacial order of this cavity produce the electrostatic response approaching the conditions expected for the virtual Lorentz cavity and thus dramatically deviating from the Maxwell cavity field.

II. RESULTS

The model considered here consists of a single non-polar solute interacting with a large number of waters mimicking a typical solvation experiment. The solute-water interaction is modeled by a Kihara potential combining a hard-sphere core of the radius $R_{\text{HS}}$ with a surface

![Diagram](image-url)
solute, we adopt the radius of the closest approach $R_{0s}$ for the interfacial dielectric response in dielectric way to estimate this effect is to notice that only the ra-
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solution introduced in our present model. We are therefore

LJ layer of width $\sigma_{0s}$ (Fig. 2)

$$
\phi_{0s}(r) = 4\epsilon_{0s} \left[ \left( \frac{\sigma_{0s}}{r - R_{HS}} \right)^{12} - \left( \frac{\sigma_{0s}}{r - R_{HS}} \right)^{6} \right].
$$

Here, "0" and "s" are used to label the solute and solvent (water), respectively; $\epsilon_{0s}$ is the energy of solute-solvent LJ attraction. Further, in order to map hard-sphere cavities used in our previous studies on the "soft" Kihara solute, we adopt the radius of the closest approach

$$
R_{0s} = R_{HS} + \sigma_{0s},
$$

where $\sigma_{0s} = 3.0 \, \text{Å}$ has been used in all numerical calculations.

The Kihara potential is the only solute-solvent interaction introduced in our present model. We are therefore missing the effect of electronic polarizability of the solute always present even for non-polar particles. A crude way to estimate this effect is to notice that only the ratio of the dielectric constants of two interfacing media matters for the interfacial dielectric response in dielectric theories. While this prescription requires testing for microscopic liquid interfaces, the model situation of a fully non-polarizable solute can be mapped on real situations of solvated non-polar particles by rescaling the solvent dielectric constant with the dielectric constant $\epsilon_{0}$ of the solute, $\epsilon \rightarrow \epsilon/\epsilon_{0}$, if such property can be reasonably defined. We will therefore proceed with the present model using the term "cavity field" to describe the electric field at the center of the solute.

Figures of the water-solute mixture were produced by Molecular Dynamics (MD) simulations of a single solute inserted in a box of SPC/E waters at standard conditions (Supplementary Material (SM)). The ratio of the cavity and external fields is obtained in the linear response approximation as the correlator of the electric field $E_{c}$ produced by water at the solute center with the water dipole moment $M_{s}$

$$
E_{c}/E_{ext} = 1 + \langle \beta/3 \rangle \delta E_{c} \cdot \delta M_{s} - E_{corr}.
$$

Here, $\delta E_{c}$ and $\delta M_{s}$ denote deviations from the corresponding average values and $\beta = 1/(k_{B}T)$ is the inverse temperature. In addition, the correction term $E_{corr}$ in the rhs of Eq. 5 accounts for the cutoff of the electrostatic interactions specific to a given simulation protocol. The result for Ewald sums is given in the SM.

Equation 5 was used to calculate the field inside the solutes of varying size $R_{HS}$ of the Kihara hard-sphere core, and the results are plotted in Fig. 3. In order to assess the effect of a uniform solute-solvent attraction on the cavity field we have simulated the configurations at two values of the LJ attraction, $\epsilon_{0s} = 0.65 \, \text{kJ/mol}$ (circles) and 20.0 kJ/mol (squares). The results for the Kihara solutes are plotted against the radius of closest approach $R_{0s}/\sigma_{s}$ defined by Eq. 1 ($\sigma_{s} = 2.87 \, \text{Å}$). Open points in the main panel and the inset refer to sets of data obtained for varied reduced dipole moment ($m^{*}$) $= \beta m^{2}/\sigma_{s}^{4}$ of the dipolar hard-sphere fluids: 0.5 (diamonds), 1.0 (triangles), and 3.0 (squares). The results for hard-sphere cavities in dipolar fluids are plotted against the radius of the closest hard-sphere solute-solvent approach $R_{0s}/\sigma_{s}$; $\sigma_{s}$ is the diameter of the solvent hard spheres. The dashed horizontal lines in the main panel and in the inset refer to the Lorentz result (L) of Eq. 2. The dash-dotted horizontal line in the inset refers to the Maxwell result (M) of Eq. 4.
the corresponding peak of the solvent-solvent distribution function. On the contrary, the larger value of the solute-solvent LJ attraction produced a substantial increase of the surface density of water as judged from the solute-solvent pair distribution function (Fig. 4). Furthermore, the radial distribution function is nearly zero between the first and second hydration shells indicating layering of water at the interface.

In order to characterize the orientational structure of the interfacial water dipoles, we define a series of distribution functions recognizing the radial symmetry of the problem and producing increasing symmetry orders of the water dipoles \( \hat{\mathbf{m}}_j \) in projection on the outward normal to the spherical solute surface \( \hat{\mathbf{r}}_j \) at the position \( \mathbf{r}_j \) of a water molecule (oxygen coordinates are used for the center of mass). The radial distribution functions are then defined in terms of \( \ell \)-order Legendre polynomials \( P_\ell(\cos \theta_{mj}) \), \( \cos \theta_{mj} = \hat{\mathbf{m}}_j \cdot \hat{\mathbf{r}}_j \) as follows

\[
g^{(\ell)}_0(r) = \frac{V}{N} \sum_j P_\ell(\cos \theta_{mj}) \delta(\mathbf{r}_j - \mathbf{r}). \tag{6}
\]

Here, \( V \) and \( N \) are the system volume and the number of particles, respectively. The zeroth-order distribution function is then the standard solute-oxygen radial distribution \( g^{(0)}_0(r) = g_{0s}(r) \). In addition to these radial distributions shown in Fig. 4 we also calculate the order parameters of the dipoles in the first hydration layer by integrating the radial functions over the volume \( V^l \) defined by the condition \( 0 < r < R_{0s} + \sigma_s/2 \)

\[
p^l = (N^l)^{-1} \rho_s \int V^l g^{(\ell)}_0(r)dr. \tag{7}
\]

Here, \( \rho_s \) is the water number density and the integrated radial function is normalized to the number of waters in the first hydration layer

\[
N^l = \rho_s \int V^l g^{(0)}_0(r)dr. \tag{8}
\]

The results for first-order, \( p^l_1 \), and second-order, \( p^l_2 \), order parameters vs the solute size are plotted in Fig. 5.

The structure of surface waters is a “squeezed” hexagonal ice lattice in which two oxygen sublayers of hexagonal sheets are brought into one plane of the closest solute-solvent approach corresponding to the first peak of the solute-oxygen radial distribution function (Fig. 4). The hydrogens of first-shell waters are randomly distributed for weak solute-solvent attraction of a hydrophobic surface (Fig. 4), but get ordered with increasing the attractive pull of the solute (Fig. 4b). The breaking of the sublayers of the hexagonal ice sheets results in buckling of the O–H–O bond from the straight line found in hexagonal ice to preferential angles of ~ 10° (more populated) and ~ 70° (less populated) (see the SM). This distribution of hydrogen-bond angles is also quite insensitive to the strength of the LJ attraction. Further, consistent with a broad distribution of the first-shell hydrogens shown by the dotted lines in Fig. 4, the water dipoles are broadly distributed for weak solute-solvent attraction, but become more ordered when LJ attraction becomes stronger (Fig. 4b). In all cases, however, the in-plane orientation of the water dipoles is preferred, and is highly resilient to the changes in the solute-solvent attraction.
The distributions of projections of water O–H bonds and the normal to the H$_2$O plane (Fig. 2) are shown in Fig. 6(a),c. Both are consistent with the picture of the in-plane orientation of the water dipoles, with the first-shell waters populating variably the states of H$_2$O plane perpendicular to the surface normal and slightly tilted, at ~10° in respect to the normal. These two states are populated in the ratio of about 1:2 for the weak attraction, but the tilted state becomes a dominant one at the stronger attraction. This observation is consistent with the general tendency of interfacial waters to increasingly occupy one particular orientational state with increasing strength of attraction of either protons or oxygens to the solute. In contrast, the hydrogen disorder generally increases when a combination of two types of attractions are present at the interface, a situation typical of hydrated proteins.

III. DISCUSSION

The results for the electric field inside the Rossky cavity in water are shown by filled points in Fig. 3. They clearly tend to the Lorentz field limit. We also plot in Fig. 3(open points) the results for the electric field inside hard-sphere cavities in dipolar fluids. Although the two situations are physically distinct, a non-polar solute in the former case vs an actual cavity in the latter, the phenomenology of the polar interfacial response is generic for both cases. The common output of these two sets of simulations suggests that the picture of non-polarized interface represents the response of free and non-polar interfaces of polar liquids in general.

It is instructive to see how the Maxwell cavity field and the Lorentz field appear based on the assumptions regarding the surface charge density [Fig. 1(a,b)]. We first assume a constant value for the surface charge density $\sigma_P = \sigma_0$. The Maxwell electric field in the medium is then the sum of the external field and the radial field propagating from a uniformly charged interface

$$E(r) = \frac{1}{\epsilon}E_{ext} + \frac{4\pi R_0^2 \sigma_0}{\epsilon r^3}r.$$  

This electric field will polarize the dielectric yielding the polarization field $P(r) = (\epsilon - 1)E(r)/(4\pi)$. This polarization field is inhomogeneous close to the interface and decays to the uniform polarization $P$ in the bulk. The polarization of the dielectric will in turn produce its own electric field, which, combined with the external field, gives the cavity field

$$E_c = E_{ext} + \int_\Omega T(r) \cdot P(r)dr.$$  

Here, $T(r) = \nabla \nabla^{-1}$ is the dipolar interaction tensor and the integral is over the dielectric occupying the volume $\Omega$ outside the dielectric cavity of radius $R_0$. The radius $R_0$ does not need to be specified since the calculation results do not depend on its value.

The radial polarization arising from the second summand in Eq. (9) gives zero contribution to the cavity field and the result of Eqs. (9) and (10) is the Lorentz field given by Eq. (2). A constant, angular-independent surface charge density makes therefore no contribution to the cavity field. It however contributes to a non-zero, spatially constant electrostatic potential inside the cavity.

Given axial symmetry of the problem, surface charge density can be expanded in Legendre polynomials, $\sigma_P(\theta) = \sum \sigma_L L(\cos \theta)$, where $\theta$ is the polar angle with the direction of the external field. Since the zero-order term $\sigma_0$ does not contribute to the cavity field, one can take the first-order term and, neglecting the quadrupolar and higher moments, have the dipolar approximation $\sigma_P(\theta) = \sigma_1 \cos \theta$. The dipole moment created by the solute interface is then $M_0 = \sigma_1 \Omega_0$. This is the situation sketched in Fig. 1(b). The direct solution of the Laplace equation with Maxwell’s boundary conditions results in the interfacial dipole

$$M_0 = -P \frac{32\sigma_0}{2\epsilon + 1}.$$  

where $P = (\epsilon - 1)/(4\pi\epsilon)E_{ext}$. The negative sign here indicates that the interface dipole $M_0$ orients oppositely to the external field, as is also clear from Fig. 1(b).

The dipolar surface charge density following from Eq. (11) is shown by the solid line in Fig. 7. This surface
charge density creates a dipolar field, which adds to the external field to produce the Maxwell field

$$E(r) = \frac{1}{\epsilon} E_{\text{ext}} + T(r) \cdot M_0. \quad (12)$$

When the polarization field $P(r)$ is calculated from Eq. \ref{eq:12} and substituted into Eq. \ref{eq:10}, the cavity field becomes

$$\frac{E_c}{E_{\text{ext}}} = \frac{\epsilon + 2}{3\epsilon} + \frac{8\pi(\epsilon - 1)}{9} \chi_1, \quad (13)$$

where

$$\chi_1 = \frac{M_0}{\Omega_0 E_{\text{ext}}} \quad (14)$$

is the dipolar response function of the solute interface to the polarization external field. The Maxwell result for $M_0$ in Eq. \ref{eq:14} gives the standard cavity field of Eq. \ref{eq:11}, while $\chi_1 = M_0 = 0$ leads to the Lorentz field of Eq. \ref{eq:2}.

Surface charge density $\sigma_P(\theta)$ provides a convenient mathematical idealization of the polarization of the interface. Its direct calculation from real-space, finite-size simulations presents, however, a significant challenge. The problem is illustrated in Fig. \ref{fig:7} which shows $\sigma_P(\theta)$ calculated at the solute surface from MD trajectories using the linear response approximation (see the SM). The result is a clearly angular-independent function, producing $\sigma_1 \simeq 0$ and $E_c \simeq E_L$. This outcome is, however, not entirely consistent with a slightly positive $\sigma_1$ following from the substitution of the simulated $E_c$ into Eq. \ref{eq:14} (dashed line in Fig. \ref{fig:7}b).

The reason of possible uncertainties in the attempts to calculate $\sigma_P(\theta)$ from sampling surface dipole directions from MD trajectories is illustrated in Fig. \ref{fig:7}. The definition of a mathematical dividing surface is uncertain when the dipolar response is strongly varied at the interface, as is the case here. We show the dielectric constant of the polar layer of radius $r$ from the cavity center obtained from the linear response approximation as $\sigma(r) = 1 + (4\pi\beta/3\Omega(r)) \delta M_s(r) \cdot \delta M_s$, where $M_s(r)$ and $\Omega(r)$ are, respectively, the dipole moment and volume of the solvent within the $r$-surface. The layer dielectric constant peaks near the interface and then slowly decays to the bulk dielectric constant $\epsilon$. Any surface drawn next to the interface will therefore reflect a different variance of the surface dipole, with a different outcome for $\sigma_P(\theta)$. How much the functional form of $\sigma_P(\theta)$ changes depending on the surface definition cannot be answered here. Attempts to obtain $\sigma_P(\theta)$ at surfaces deeper into the bulk did not produce converged functions and therefore are not shown here.

It seems worth emphasizing some critical differences between both (Maxwell and Lorentz) continuum results and the molecular picture offered by the present numerical simulations. The continuum cavity field does not depend on the cavity size, but only on the cavity shape (e.g., on the length/diameter ratio for a cylindrical cavity). This is of course the consequence of neglecting the actual spread of the interfacial region relative to the size of the solute, which is behind the definition of the interface as a mathematical dividing surface. Our simulations allow us to estimate the solute size at which this approximation becomes valid. Figure \ref{fig:3} shows that the cavity field starts leveling off to a size-independent limit at $R_0/\sigma_s \simeq 2 - 2.5$. It gives an estimate of the solute size at which the “Lorentz continuum” starts to take hold: the solute needs to be 4 - 5 times larger than the solvent molecule. The “Maxwell continuum” is, however, never reached in our simulations.

\section*{IV. EXPERIMENTAL OBSERVABLES}

The surface polarization effects discussed here have a number of consequences observable in a macroscopic laboratory experiment. The total free energy of polarizing the dielectric is measured by the dielectric experiment and is affected by the dipole accumulated at the interface with a non-polar solute. The total dipole of a mixture sample $M_{\text{mix}}$ is reduced relative to the homogeneous solvent by the volume excluded by the solute and, in addition, is affected by the dipole of the interface

$$M_{\text{mix}} = P\Omega - (23)(\epsilon - 1)M_0N_0. \quad (15)$$

Here, $N_0$ is the number of the solutes in the solution and $\Omega$, as above, is the solvent volume. The solutes are assumed to be non-interacting, although the theory can be extended to non-ideal solutions requiring the solute-solute structure factor as an additional input.

The standard arguments of the theory of dielectrics then suggest the equation for the dielectric constant of
FIG. 8. Relative dielectric constant increment $\Delta \epsilon/\epsilon$, $\Delta \epsilon = \epsilon_{\text{mix}} - \epsilon$ for several aqueous solutions: lysozyme (filled squares), dioxane (open circles), glucose (open diamonds), and maltotriose (open triangles). The solid line (M) shows the result of Eq. (18), while the dashed line (L) is the dielectric increment for a solute with no surface charge density, $\sigma_1 = 0$, and therefore Lorentz result for the cavity field [Eq. (19)]. Experimental and calculation results used to convert the experimentally reported solute concentrations to volume fractions can be found in the SM. The dash-dotted lines connect the experimental points for two saccharides.

The mixture of the solvents and cavity fields inside solutes both give experimental routes to measure the dipolar response function of the interface $\chi_1$. Combining two equations together, one gets

$$\frac{\epsilon}{\epsilon_{\text{mix}}} = 1 + 3\eta_0 \left[ \frac{E_c}{E_{\text{ext}}} - 1 \right]$$

(16)

where $\eta_0 = N_0Q_0/(N_0Q_0 + \Omega)$ is the volume fraction of the solutes. Equations (16) and (18), taken together, suggest that dielectric constants of low-concentration solutions and cavity fields inside solutes both give experimental routes to measure the dipolar response function of the interface $\chi_1$. Combining two equations together, one gets

$$\frac{\epsilon}{\epsilon_{\text{mix}}} = 1 + 3\eta_0 \left[ \frac{E_c}{E_{\text{ext}}} - 1 \right]$$

(17)

For the cavity field $E_c = E_\text{M}$ given by Maxwell’s form in Eq. (11) one arrives at the result (15,27):

$$\frac{\epsilon}{\epsilon_{\text{mix}}} = 1 + \eta_0 \frac{3(\epsilon - 1)}{2\epsilon + 1}.$$ 

(18)

This equation is consistent with the Maxwell-Wagner theory in the limit of small volume fraction $\eta_0$.

Figure 8 illustrates the application of Eqs. (16) and (18) to solutions of ionic and hydrogen-bonding substances in water. The closed squares in the figure show the relative dielectric constant increment $\Delta \epsilon/\epsilon$, $\Delta \epsilon = \epsilon_{\text{mix}} - \epsilon$ for hydrated lysozyme. At the pH of the measurements the protein carries the total charge of $+10$ electron units. Open circles in the figure show the results for the dioxane-water mixture. Dioxane offers strong hydrogen-bond acceptor sites. Finally, diamonds and triangles show, respectively, hydrated glucose and maltotriose both offering multiple hydrogen-bonding sites. The results for lysozyme and for two saccharides refer to dielectric increments of the loss peak of the water component of the solution. This approach has allowed us to eliminate the contribution of the permanent dipole moment of the solute, characterized by a much longer relaxation time. The increment of the water loss peak reflects only the interfacial polarization described by the present model. The dipole moment of dioxane is zero and the increment of the static dielectric constant was used in that case.

The experimental results are compared to Maxwell [Eq. (13), solid line] and Lorentz [Eq. (11), dashed line] scenarios. All solutes, potentially breaking the interfacial network of hydrogen bonds, are either consistent with the Maxwell interfacial polarization or fall between the Maxwell and Lorentz predictions.

The Lorentz interfacial interface predicts a much enhanced sensitivity of the dielectric constant to non-polar impurities, which also implies a significant free energy penalty for polarizing such mixtures. Experimental testing of the Lorentz scenario should therefore meet with obvious solubility difficulties. However, scenarios intermediate between the Lorentz and Maxwell limits clearly exist and the concept of an effective surface charge density $\sigma_P$ provides a useful conceptual framework for developing theories of polar response not restricted to Maxwell’s boundary conditions.

While uniform fields of the dielectric experiment measure the dipolar response of the entire solution, non-uniform external fields give access to the response function $\chi_1$ and therefore to the dipolar polarization of the interface. If the external field varies on a scale large compared to the dimension of the solute, the interfacial dipole couples to the field gradient. The result is a force acting on the solute

$$F_z = (\Omega_0/2)\chi_1 \nabla_z E_{\text{ext}}(z)^2,$$

(19)

where the $z$-axis is chosen along the external field. This phenomenon, known as dielectrophoresis, allows a direct access to the polarization of the interface. The Maxwell solution for $\chi_1$ [Eqs. (11) and (18)] leads to the standard dielectrophoresis coefficient $K \propto \chi_1$ used in the theory of colloidal suspensions. In these applications, $\epsilon$ is typically replaced with the ratio of the dielectric constants of the solvent and the solute, $\epsilon \rightarrow \epsilon/\epsilon_0$, with the result

$$K = \frac{\epsilon_0 - \epsilon}{\epsilon_0 + 2\epsilon}.$$ 

(20)

If the solute is more polar than the solvent, one gets positive dielectrophoresis ($K > 0$, $\chi_1 > 0$), and solute’s attraction to a stronger electric field. The opposite case of $K < 0$ ($\chi_1 < 0$) implies negative dielectrophoresis and thus repulsion of the solute from the field.
V. SUMMARY

This paper extends our previous studies\textsuperscript{14,15} of dipolar fluids interfacing hard-sphere cavities to attractive non-polar solutes hydrated by SPC/E water. In-plane orientational structure of the surface dipoles holds for interfaces of both dipolar liquids and water. The orientational distribution of the interfacial dipoles alters the boundary conditions of the polar response problem monitored here in terms of the electric field inside the solute. Like in previous studies of cavities in dipolar fluids, the field inside the solute does not follow the prediction of Maxwell’s electrostatics [Eq. (1)] and instead tends, with increasing solute size, to the limit established by the Lorentz field of a non-polarized interface [Eq. (2)].

We find that the deviation of the cavity field from the Lorentz result is given by the dipolar response function of the interface that also enters the dielectric constant of an ideal solution. Finding the slope of the dielectric increment of the low-concentration mixture vs the solute concentration [Eqs. (17) and (16)] thus provides a direct input into the cavity field [Eqs. (17) and (13)]. This statement also applies to the frequency-dependent response. As such, the dielectric increment of the fast water component of the solution was taken in Fig. 8 to avoid the effect of a slowly relaxing permanent dipole of the solute. Our formalism is therefore applicable to ionic and polar solutes when the frequency range exceeds the loss peak of the solute dipole. Along these lines, the formalism can also be used at much higher frequencies of UV/VIS light to find the refractive index corrections for rates of radiative decay of photoexcited chromophores and quantum dots in solution. According to our present results, this input should be sought from measuring the refractive index of corresponding mixtures [Eq. (17)].

We also find that small differences in the cavity field, which are hard to resolve within the current simulation protocol, propagate in very substantial differences in the dielectric response of a mixture. While this fact also spells out a thermodynamic difficulty of preparing solutes when the frequency range exceeds the loss peak of the solute dipole. Along these lines, the formalism can also be used at much higher frequencies of UV/VIS light to find the refractive index corrections for rates of radiative decay of photoexcited chromophores and quantum dots in solution. According to our present results, this input should be sought from measuring the refractive index of corresponding mixtures [Eq. (17)].

Returning to the question posed in the Introduction, the liquid interface assumes the structure that makes its contribution to the field inside and outside the solute essentially null. The whole solvent response to an external polarizing field is given by the uniform polarization of the bulk and does not include an inhomogeneous component due to the surface charge [second summand in Eq. (12)]. In retrospect, this outcome should have been expected. The standard practice of liquid state theories and corresponding numerical simulations suggests a short propagation length of perturbations in liquids. A cavity or a non-polar solute should therefore be “invisible” to an observable (e.g., Maxwell field) measured a few solvent diameters from the interface, as indeed our results show. This result of course goes against the concept of the surface charge inducing a long-ranged Coulomb perturbation in the solvent.

The original concept of electric polarization envisioned by Maxwell anticipates a limited, elastic response of medium’s electric charge to an external electric field. For molecular dipoles of a dielectric material, this concept implies limited small-amplitude reorientations of the dipoles, aligning them with a weak external field (linear response). This physical picture, which seems to match the problem of dipolar polarization of a free solid-like interface, was historically extended to liquid dielectrics which, in contrast, have more extended ability to respond by both changing the positions of their dipoles and producing large-amplitude dipolar reorientations. In other words, the translational and rotational mobility supported by the liquid phase allows the surface dipoles to react to the creation of the interface by rearranging their dipolar orientations in a way of diminishing the stress of a sharply varying density profile. This orientational order responds to a weak external field by the rules that do not require a surface charge and the induction of an interfacial dipole combining the negative and positive lobes of the surface charge density (Fig. 11). The boundary conditions applicable to these systems differ from the ones anticipated for Maxwell dielectrics.

By the way of yet another historical aberration, the standard theory of dielectrics has been mostly applied not to free liquid interfaces, for which Maxwell’s construct was put forward, but to highly hydrophilic and wetted interfaces covered with surface ions or to solvation of molecular ions (Born theory of ion solvation and its extensions). The agreement between observations done for these surfaces and solutions with the predictions of dielectric models is often used to support the conceptual framework of dielectric theories developed for free or non-wetted (small attraction) surfaces. Our present development cautions against this inconsistency and points out that the orientational structure of the interface defines boundary conditions of dielectric theories and with that the microscopic and macroscopic fields observed near dielectric interfaces.

The picture of in-plane dipolar polarization of the liquid interface is rather robust and insensitive to the interface curvature. It holds even for planar surfaces (filled diamond in Fig. 12) suggesting that the conclusions reached here for an admittedly narrow range of solute radii may extend to larger solutes of potentially meso-to-macroscopic size.

What matters for the boundary conditions entering the electrostatic response functions is the orientational distribution of surface waters. This can be altered by surface ions and polar groups, which potentially can create a non-zero surface charge density $\sigma_p \neq 0$ matching the standard conditions of Maxwell’s electrostatics. The standard prescriptions will apply to those cases. It follows directly from Coulomb’s law that the change of the
normal component of the electric field at the interface of a dielectric with vacuum is related to the surface charge density, $-\Delta E_n = 4\pi\sigma_p$. The near-zero $\sigma_p$ then implies the continuation of the normal component of the electric field, and, therefore, the continuation of the vector $E$ of the Maxwell field across a dielectric interface.

Liquids with large cohesive energy, network liquids, such as water, in particular, seem to be particularly relevant to this discussion. The strength of hydrogen bonds ($\sim 4 - 5 k_B T$ per bond) is so significant that in-plane dipolar pattern may withstand local electric fields of solute partial charges. The ratio of the characteristic strengths of the solute-solvent to solvent-solvent interactions will therefore determine the orientational distribution of the surface dipoles and, ultimately, the type of boundary conditions used in calculations of the electrostatic response.

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