Attraction between like-charge surfaces in polar mixtures

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Abstract – We examine the force between two charged surfaces immersed in aqueous mixtures having a coexistence curve. For a homogeneous water-poor phase, as the distance between the surfaces is decreased, a water-rich phase condenses at a distance $D_c$ in the range 1–100 nm. At this distance the osmotic pressure can become negative leading to a long-range attraction between the surfaces. The osmotic pressure vanishes at a distance $D_c < D_t$, representing a very deep metastable or globally stable energetic state. We give analytical and numerical results for $D_t$ and $D_c$ on the Poisson-Boltzmann level.

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The forces between charged objects in electrolyte solutions are of fundamental importance in biology and colloidal science. Within the Poisson-Boltzmann (PB) mean-field theory, in a single pure solvent, the interaction between symmetrically charged colloids is always repulsive [1]. Experiments, on the other hand, have shown that highly charged colloids can attract each other when multivalent ions are present. This discrepancy has been explained theoretically by counterion correlations [2], fluctuation-induced forces [3] and other non-electrostatic interactions [4]. More recently, the PB theory was generalized by strong coupling theory [5,6], which predicts attractive forces appear between similar charged colloids [7,8], in the presence of ions[15]. Experiments, on the other hand, have shown that high charged colloids can attract each other when multivalent ions are present. This discrepancy has been explained theoretically by counterion correlations [5], fluctuation-induced forces [3] and other non-electrostatic interactions [4]. More recently, the PB theory was generalized by strong coupling theory [5,6], which predicts attractive forces appear between similar charged colloids [7,8].

In this letter we use the PB theory to show that strong attractive forces occur in mixtures of polar solvents. The colloids are modeled as flat surfaces located at $z = ±D/2$ and uniformly charged with a charge density $eσ$ per unit area, where $e$ is the elementary charge. A small amount of monovalent ions and weakly charged surfaces are assumed and therefore the coupling parameter is small and the PB theory is applicable [6]. The partially miscible solvents have a coexistence curve below the mixture critical temperature $T_c$ in the absence of ions. This coexistence curve is further modified by the presence of ions [15].

The grand potential density is given by [14,23]

$$\frac{\omega}{T} = \phi_b(\phi) + \frac{C}{2} |\nabla\phi|^2 + \frac{1}{2T} \varepsilon(\phi)(\nabla \psi)^2 + n^+(\log(v_0 n^+ - 1) + n^- (\log(v_0 n^- - 1) - (\Delta u^+ n^+ + \Delta u^- n^-) \phi - \lambda^+ n^+ - \lambda^- n^- - \mu \phi. \ (1)$$

Here the Boltzmann constant is set to unity, $T$ is the thermal energy and $C$ is a positive constant. $\phi$ is the volume fraction of water ($0 \leq \phi \leq 1$); far away from the surfaces the mixture is homogeneous and water-poor with composition $\phi_0 < 1/2$ and ion densities $n_0$. We use a regular solution form for the free energy of a binary mixture $\phi_b(\phi) = \phi \log(\phi) + (1 - \phi) \log(1 - \phi) + x\phi(1 - \phi)$. [24], where $\chi \sim 1/T$ is the Flory parameter and $v_0 = a^3$ is the molecular volume of both liquids. The third term in eq (1) is the

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electrostatic energy of the mixture, where \( \psi \) is the electrostatic potential. The mixture dielectric constant depends on the composition through a linear relation: 

\[
\varepsilon(\phi) = \varepsilon_c + (\varepsilon_w - \varepsilon_c)\phi,
\]

where \( \varepsilon_c \) and \( \varepsilon_w \) are the pure co solvent and water dielectric constants, respectively. The second line of eq. (1) is the ideal gas entropy of point-like ions, where \( n^+ \) and \( n^- \) are the positive and negative ion density, respectively. The first term on the third line is the bilinear solubility interaction of the ions and the solvent: the parameters \( \Delta u^+ \) and \( \Delta u^- \) measure the affinity of the positive and negative ions toward the water environment, respectively. We use the common case where both ions are hydrophilic and assume the symmetric interaction \( \Delta u^+ = \Delta u^- = \Delta u > 0 \). Lastly, \( \lambda^\pm \) and \( \mu \) are the Lagrange multipliers (chemical potentials) of the positive and negative ions and water composition, respectively.

The ion densities obey the Boltzmann distribution

\[
n^\pm = v_0^{-1}e^{\lambda^\pm \varepsilon(z)\psi/T + \Delta u\phi}.
\]

In a salt reservoir we have \( \lambda^\pm = \log(v_0n_0) - \Delta u\phi_0 \). Alternatively, when the mixture contains only negative counterions, \( \lambda^\pm \) is determined self-consistently by charge conservation: \( \int n^- dz = 2\sigma \).

The electrostatic potential is determined by the Poisson equation

\[
\nabla \cdot (\varepsilon(\phi)\nabla \psi) = -\varepsilon(n^+ - n^-)
\]

supplemented by the boundary condition on the surfaces

\[-n^\pm \nabla \psi(\pm D/2) = e\sigma/\varepsilon(\phi),\]

where \( n^\pm \) is the outward unit normal to the surface. Finally, the Euler-Lagrange equation for \( \phi \) reads

\[
C\nabla^2\phi = \frac{\partial f_b}{\partial \phi} - \frac{1}{2T} \frac{d\varepsilon}{d\phi}(\nabla \psi)^2 - \Delta u(n^+ + n^-) - \mu.
\]

In order to isolate electrostatic effects from the regular wetting-like behavior, we assume zero short-range chemical or long-range van der Waals interactions with the surfaces, leading to the boundary condition \( n^\pm \nabla \phi(z = \pm D/2) = 0 \).

The net force exerted on the surface by the liquid is given by the osmotic pressure \( \Pi = P_{zz} - P_b \), where \( P_b \) is the bulk pressure: \( P_b/T = \phi_0\partial f_b(\phi_0)/\partial \phi - f_b(\phi_0) + 2n_0(1 - \Delta u\phi_0) \), \( P_{zz} \) is the zz component of the Maxwell stress tensor [17,25]:

\[
P_{zz} = \frac{C}{2} \left( \frac{d\phi}{dz} \right)^2 - C\phi \frac{d^2\phi}{dz^2} + \phi \frac{d^2f_b}{d\phi^2} - f_b + (1 - \Delta u\phi)(n^+ + n^-) - \frac{1}{2T} \left( \frac{d\xi}{d\phi} + \varepsilon \right) \left( \frac{d\psi}{dz} \right)^2.
\]

\( P_{zz} \) is independent of \( z \) and can be calculated at the midplane \( (z = 0) \) where by symmetry \( \phi(d\phi/dz = d\varepsilon/dz = 0) \). Multiplying eq. (4) by \( \phi \) and inserting into eq. (5) we obtain

\[
\Pi = Tn_m - T\omega_b(\phi_m) - P_b,
\]

where \( \omega_b = f_b - \mu\phi \) and \( \phi_m \) and \( n_m \) are the composition and total ion density at the midplane, respectively.

In aqueous mixtures, water is drawn towards the walls by field gradients (dielectrophoretic) and solvent-induced (electrophoretic) forces [17,23,26]. These forces are interdependent as is evident from eqs. (3) and (4). From the solution of the governing equations we find two possible scenarios, distinguished according to the composition profiles at infinite separation. In the first scenario, when the surfaces are far apart, \( \phi(z) \) near each surface is composed of three layers: a thin water-rich layer \( (\phi > 1/2) \) of width \( t \) at the surfaces, a second layer with width of the bulk correlation length \( \xi \) adjacent to it, where the composition \( \phi \approx \phi_0 \). As the surface separation decreases down to \( D = D_t \), the general features of this profile do not change. In particular, only the width of the \( \phi \approx \phi_0 \) region changes if screening is strong, that is, when \( l_D \ll D \) where \( l_D \approx n_0^{-1/2} \) is the Debye length calculated at \( \varepsilon = \varepsilon(\phi_0) \). The dashed curve in fig. 1(a) shows \( \phi(z) \) just before \( D = D_t \) for this case.

In the second scenario, dielectrophoretic and solvation forces are not strong enough and at large surface separations \( \phi \) is always close to the bulk value \( \phi_0 \). As \( D \) decreases from infinity the water composition between the surfaces increases continuously. In particular, in the ideal gas regime, defined by \( b \gg D \), where \( b \propto \sigma^{-1} \) is the Gouy-Chapman length, \( \phi(z) \) is nearly uniform. Figure 1(b) shows such profiles for a mixture containing only counterions. Here, the ion profile \( n(z) \) and electric field are nearly uniform leading to small composition gradients.

In both scenarios, when \( D < D_t \) the whole space between the surfaces becomes rich in water (see the solid curves in fig. 1). The transition occurs at a distance in the range \( D_t = 1–100 \) nm depending on the parameter values.
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Fig. 2: (Colour on-line) (a) Osmotic pressure $\Pi$ vs. surface separation $D$ in the strong screening limit for different values of $n_0$ (other parameters as in fig. 1(a)). Inset: excess grand potential for different values of $\Delta u$. (b) $\Pi$ in the ideal gas regime for different values of $\Delta u$ (other parameters as in fig. 1(b)).

Inset: the function $-\omega_b(\phi)$. Filled square and circle markers correspond to $\phi_0$ and the binodal compositions, respectively. In (a) and (b) open markers on solid lines correspond to distances $D$ with the same symbols in fig. 1(a) and (b).

and is accompanied by a decrease in the osmotic pressure. This can lead to a negative osmotic pressure, such that a surface separation $D_e < D_t$ exists at which $\Pi = 0$. $\phi(z)$ at $D = D_e$ is shown by the dash-dot curves in fig. 1.

The excess grand potential per unit area relative to infinite separation is $\Delta \Omega(D) = -\int_0^D \Pi(D')dD'$. $\Delta \Omega(D)$ has a cusp at $D = D_t$ and this results in a discontinuity of the pressure. Close to $D_t$, both a water-poor and a water-rich profiles are solutions of the Euler-Lagrange equations but only one of these is a stable solution. At $D > D_t$ the water-poor phase is stable, at $D > D_t$ the water-rich phase is stable, and at $D = D_t$ the grand potentials of the two phases are equal. Furthermore, $\Delta \Omega$ has a minimum at $D_e$ ($D_e < D_t$) corresponding to mechanical equilibrium. An example of $\Delta \Omega(D)$ is shown in the inset of fig. 2(a), where the dashed line is $\Delta \Omega$ of metastable solutions.

The transition to a water-rich phase has a different physical origin in the two limiting cases. In the strong screening regime, the transition is promoted by the energy gained when the interface vanishes, as in capillary condensation. The thin water-rich layer near the walls remains nearly independent of $D$. In the ideal gas regime, on the other hand, when the composition approaches the coexistence composition, preferential solvation promotes a transition to a water-rich phase. For intermediate cases, both mechanisms play a role in the transition. These effects are enhanced but are not limited to the vicinity of the critical temperature where the differences between the phases become smaller.

In fig. 2 we plot the osmotic pressure as a function of surface spacing; when $\Pi > 0$ the surfaces repel each other while for $\Pi < 0$ they attract. $\Pi$ is discontinuous at $D = D_t$. In fig. 2(b) we plot $\Pi$ for different values of $\Delta u$ in the strong screening limit. An increase in $n_0$ decreases $\Pi$ at large distances but increases it at small distances (entropy loss of the ions).

The negative jump in $\Pi$ at $D = D_t$ increases with decreasing $n_0$. This can be explained by the interplay between the first two terms in eq. (6). At $D = D_t$ the increase in $\phi_m$ dominates the decrease in $\Pi$, an effect more pronounced for smaller values of $n_0$ since the positive ideal gas term is proportional to $n_m \times n_0$. Recall that in a pure solvent only the ideal gas term exists and hence $\Pi$ is always positive. Furthermore, this entropic repulsion will eventually dominate $\Pi$ leading to $\Pi = 0$ at a separation $D_e$.

In fig. 2(b) we plot $\Pi$ for different values of $\Delta u$ in the ideal gas regime. For $\Delta u = 1$ the pressure is purely repulsive since preferential solvation is not strong enough to induce a water-rich phase. For $\Delta u = 7$ the interaction is repulsive down to $D_t$ where $\Pi$ becomes negative. When $\Delta u$ increases to 11 the transition is at a larger distance and to a more negative pressure. The inset of fig. 2(b) shows the function $-\omega_b(\phi)$, showing a decrease in pressure at the transition (cf. eq (6)).

Figure 3 shows $\Pi$ in the weak screening regime where $l_p \approx D$ and for three different values of $\Delta u$. When $\Delta u = 0$, the dielectrophoretic force alone can induce a water-rich

![Fig. 3](image-url)
phase albeit the pressure is always repulsive [26]. The pressure can become attractive for $\Delta u = 1$ or 7. Corresponding curves of $\Delta \Omega$ are plotted in the inset of fig. 3. These show a metastable minimum at a finite value of $D$, $D = D_c$, for $\Delta u = 1$ and a global minimum for $\Delta u = 7$. The depth of the minimum for $\Delta u = 7$ is $\approx 440$ T for two colloids with an effective surface area of 100 nm $\times$ 100 nm.

For $\Delta u = 10$, a second metastable minimum can appear in the curve $\Delta \Omega(D)$ at a smaller value of $D$, see the dashed and dash-dot curves in fig. 4. A similar value of $\Delta u$ is cited in the literature for mixtures of water and 1-butanol containing NaCl at room temperature [12]. When $\sigma$ is further increased (solid curves), this minimum can become globally stable. For large enough $\sigma$, only the second minimum exists (not shown). In this large $\Delta u$ case, preferential solvation leads to the liquid between the surfaces being nearly pure water; $\phi_m$ is close to unity and $\omega_z$ diverges.

The distance $D_t$ can be obtained in the two limiting cases presented in fig. 1. In the strong screening limit we use the continuity of $\Delta \Omega$ at $D_t$ to obtain [27]

$$D_t \simeq 2t + 2 \frac{\xi P_b + \int_D^{D/2-t} \omega(\phi(z)) \, dz}{P_b - P_h},$$  \hfill (7)

where we used $\psi \approx 0$ at the midplane.

At $D = D_t$ the surfaces can be regarded as isolated and a zeroth-order approximation of $\psi$ can be obtained using the well-known result for a single surface [1] with a homogeneous composition $\phi \approx \phi_h$ and a modified Debye length $\xi(\phi_h) = (\varepsilon(\phi_h)/2\varepsilon_0\mu_0 \omega_h(\phi_h - \phi_0)\mu_0)T/\omega$. Since at $|z| < D/2 - t$ $e\psi/T \ll \Delta u(\phi_h - \phi_0)$, $t$ is obtained from $e\psi/T = 0.01\Delta u(\phi_h - \phi_0)$. In analogy with classical mean field theory [24], the Landau expansion of eq. (8) around $\phi_c = 1/2$ gives for $\xi$

$$\xi \approx n_0^{1/3} \frac{1}{\sqrt{1 - \frac{T}{T_c} - 2\Delta u n_0 \omega_h(\frac{\phi}{2} - \phi_0) + \frac{3}{2}}} \frac{1}{\Delta u n_0 \omega_h(\frac{\phi}{2} - \phi_0) + \frac{3}{2}}. \hfill (9)$$

The preferential solvation term in the root is comparable in magnitude to $1 - T/T_c$. Thus, the bulk correlation length is modified appreciably by the preferential solvation of the ions.

Figure 5(a) shows the comparison of $D_t$ from eq. (7) with numerical results for different values of $n_0$. The agreement is quite good despite the crude approximation. $D_c$ in fig. 5(a) decreases slightly when $n_0$ increases, its value being $\approx 5$ nm.

In the ideal gas regime $\phi(z) \approx$ const and the counterions density $n^-$ is uniform and equal to the average charge density $\omega_z$ [24]: $\langle n^- \rangle \approx 2n/D$. The composition equation reads

$$\frac{\partial f}{\partial \phi} - \frac{2\Delta u n^\sigma}{D} + \mu_0 = 0. \hfill (10)$$

Here preferential solvation merely shifts the chemical potential of the mixture. Hence, $D_t$ occurs when $\phi$ is the
for a mixture at a critical composition $\phi_0 = 1/2$ at three temperatures above $T_c$. For $T/T_c = 1.01$ (dash-dot curve) the interaction is purely repulsive. When the temperature is decreased the interaction turns attractive (dashed curve), becoming stronger closer to $T_c$ (solid curve). Here $n_0 = 0.01$ M, $\Delta u = 4$ and $\sigma = 1$ nm$^{-2}$. The walls have a surface area of $A = 0.01$ $\mu$m$^2$.

binodal composition at which $\partial f_b/\partial \phi = 0$. Thus,

$$D_\text{t} \simeq -2\sigma \Delta u / \mu_0,$$

which means that $D_\text{t} \to \infty$ when the bulk composition approaches the binodal ($\mu_0 \to 0$).

$D_c$ can be found by noting that $\Pi = 0$ in eq. (6) gives $n_c = \omega_b(\phi_c) + P_b/T$, where $\phi_c$ and $n_c$ are the composition and ion density at $D_c$, respectively. Inserting this into eq. (4) we obtain for $\phi_c$

$$\frac{\partial \omega_b(\phi_c)}{\partial \phi} = \Delta u (\omega_b(\phi_c) + P_b/T),$$

which is solved and then used to get $n_c$ and $D_c$ from the equations for $\Pi$ and ($n^-$), respectively. Comparison of the formulae for $D_t$ and $D_c$ with the full numerical results are presented in fig. 5(b). As expected, the agreement is good for small values of $\sigma$ where the ideal gas limit is valid. The dashed curve in fig. 5(b) marks the charge density above which a water-rich layer exists at infinite separation and the above approximations no longer hold.

We find that an attractive interaction is possible even at temperatures above $T_c$. Since at $T > T_c$ there is no miscibility gap, the first-order phase transition is missing and there is no discontinuity in the pressure. Instead, the pressure becomes smoothly attractive with decreasing plate distance. Figure 6 shows the interaction potential $U = \Delta \Omega \times A$ for two plates of surface $A$ immersed in a critical mixture at three different temperatures. As the critical temperature is approached, a purely repulsive potential (dash-dot curve) becomes attractive (dashed curve). The interaction is more attractive closer to $T_c$ (solid curve). The depth and location of the minimum in the curves and their dependence on the temperature are similar to those observed in recent experiments on the salt-dependent interaction of a charged particle suspended in a critical binary mixture near a charged wall [21]. Thus, the mechanism we describe may be important to capture correctly electrostatic effects in these experiments. We believe that it is possible to attribute some of the results of these experiments to a solvation related force [28] in addition to the critical Casimir force.

The qualitative difference in the interaction between charged colloids or macromolecules in mixtures compared to pure solvents stems from the contribution to $\Pi$ of the second term in eq. (6): $\omega_b(\phi_m$) – the midplane mixture grand potential. This term is absent in pure solvents. Thus, unlike similar ion-induced phase transitions in pure solvents [29] we observe a jump to a negative pressure (at $D_t \approx 1–100$ nm). Moreover, the nontrivial dependence of $\omega_b(\phi_m$) on the system parameters $(T, \phi_0, n_0, \sigma)$ through the governing equations leads to qualitatively different behavior compared to usual condensation transitions due to surface fields [27]. In addition, the attractive force is in many cases strong and long-range compared to the van der Waals force [1] as is evident by the values of $\Pi$ at $D = D_t$ (fig. 2 and fig. 3) and the energy minimum being deeper than $\sim 100$ T.

The mechanism we describe should be at play in the aggregation of charged colloids or macromolecules in mixtures near the coexistence temperature [18,19]. We believe it is directly relevant to the attraction seen between colloids and surfaces in critical mixtures and attributed to the critical Casimir force [20–22].

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