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Solvation effects in phase transitions in soft matter

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Abstract
Phase transitions in polar binary mixtures can be drastically altered by even a small amount of salt. This is because the preferential solvation strongly depends on the ambient composition. Together with a summary of our research on this problem, we present some detailed results on the role of antagonistic salt composed of hydrophilic and hydrophobic ions. These ions tend to segregate at liquid–liquid interfaces and selectively couple to water-rich and oil-rich composition fluctuations, leading to mesophase formation. In our two-dimensional simulation, the coarsening of the domain structures can be stopped or slowed down, depending on the interaction parameter (or the temperature) and the salt density. We realize stripe patterns at the critical composition and droplet patterns at off-critical compositions. In the latter case, charged droplets emerge with considerable size dispersity in a percolated region. We also give the structure factors among the ions, accounting for the Coulomb interaction and the solvation interaction mediated by the composition fluctuations.

(Some figures in this article are in colour only in the electronic version)

1. Introduction
In many soft materials, phase transitions and structure formation occur in the presence of charged objects including small ions, charged colloids, and polyelectrolytes in polar fluids (mostly in water) [1–3]. Much attention has been paid to the consequences of the long-range Coulomb interaction, but not enough attention has yet been paid to the solvation interaction among charged objects and polar molecules [4–7]. In the theoretical physics literature, the solvation interaction has been mostly neglected in analytic theories and numerical simulations. In real aqueous systems, a number of molecular interactions are operative at short distances under the influence of a long-range electric potential [4]. In particular, hydrogen bonding is of primary importance in the phase behavior in aqueous systems, as studied in neutral polymer solutions [8]. However, small ions like Na\(^{+}\) form a solvation shell composed of several water molecules and are hydrophilic, around which the energy of the ion–dipole interaction typically exceeds the energy of the surrounding hydrogen bonds. Contrary to this, some neutral or ionized particles dislike contact with water molecules and become hydrophobic. Hydrophilic and hydrophobic particles deform the surrounding hydrogen bonding in different ways [4]. Thus solvation is highly complex in aqueous mixtures. In this paper we will show that the preferential (or selective) solvation of ions in a mixture solvent can drastically influence the phase transition behavior.

Nabutovskii et al. [9] discovered the possibility of mesophases in electrolytes in the presence of a coupling between the composition and charge density in the free energy. In our theory of electrolytes [10, 11], such a coupling originates from solvation in polar fluids and its magnitude is exceedingly high in many real systems. Recently, several theoretical groups have examined ion effects, including the solvation effect in mixture solvents, in electrolytes [10–19], polyelectrolytes [20, 21], and ionic surfactants [22] using phenomenological models. A review of the static properties along these lines was presented in [23]. A number of problems in the dynamics remain less well studied [16, 19]. On the other hand, a large number of microscopic simulations have been performed, for example to calculate the ion distributions near water–air interfaces [24] and solid surfaces [25, 26].

In this paper we are particularly interested in the statics and dynamics of phase transitions in aqueous mixtures with an antagonistic salt, which is composed of hydrophilic and hydrophobic ions. While the surface tension of liquid–liquid (and water–air) interfaces is known to be slightly increased by most ion pairs [27, 28], it can be dramatically decreased
by an antagonistic salt as observed in [29, 30]. Hydrophilic and hydrophobic ions segregate at liquid–liquid interfaces on the scale of the Debye screening length $\kappa^{-1}$, resulting in a large electric double layer [11, 12, 16]. They also interact differently with water-rich and oil-rich composition fluctuations. Mesophases (charge density waves) then appear near the solvent criticality for a sufficiently large solvation asymmetry between the cations and the anions.

We mention recent experiments in which an antagonistic salt was used: (i) Luo et al observed a unique interfacial ion distribution in an x-ray reflectivity experiment [30]. (ii) Sadakane et al [31] added a small amount of sodium tetraphenylborate NaBPh$_4$ to a near-critical mixture of D$_2$O and trimethylpyridine (3MP). This salt dissociates into hydrophilic Na$^+$ and hydrophobic BPh$_4^-$. The latter anion consists of four phenyl rings bonded to an ionized boron. They found a peak at an intermediate wavenumber $q_m (\sim 0.1 \ \text{Å}^{-1} \sim \kappa)$ in the intensity of small-angle neutron scattering. The peak height was much enhanced with formation of periodic structures. Moreover, they observed multi-lamellar (onion) structures at small volume fractions of 3MP (in D$_2$O-rich solvent) far from the critical point [32], where BPh$_4^-$ and solvating 3MP form charged lamellae. These findings demonstrate very strong hydrophobicity of BPh$_4^-$, which was effective close to the solvent criticality [31] and far from it [32]. (iii) Another interesting phenomenon is spontaneous emulsification (formation of small water droplets) at a water–nitrobenzene (NB) interface [33, 34]. This was observed when a large water droplet was pushed into a cell containing NB and an antagonistic salt (tetraalkylammonium chloride). This instability was caused by ion transport through the interface.

The organization of this paper is as follows. In section 2, we will present a summary of the solvation effects, particularly focusing on preferential solvation. In section 3, the Ginsburg–Landau model and dynamic equations for such systems will be given. As a new result, the surface tension of binary mixtures was shown to decrease to zero with increasing density. In section 4, we will numerically examine mesophase formation with an antagonistic salt in two dimensions (2D).

2. Solvation effects

This section is an introduction to preferential solvation. It provides a starting point for our theory in section 3.

2.1. Solvation shell and the Born theory

In polar liquids, a number of polar molecules are attached to a hydrophilic ion such as Na$^+$, Li$^+$, or Ca$^{2+}$ to form a solvation shell due to the ion–dipole interaction [4, 35] (see figure 1). However, hydrophobic particles interact with water molecules repulsively, so they tend to form aggregates in water and are more soluble in oil than in water. We may consider mixture solvents composed of polar and less polar components, where hydrophilic (hydrophobic) ions are surrounded by molecules of the more (less) polar component [5, 7] (see the right panel of figure 1). That is, solvation occurs preferentially or selectively in mixture solvents. There can be a variety of mixture solvents and, in this paper, we call the two components simply water and oil. Note that solvation is also called hydration for water (or heavy water) and for aqueous mixtures containing water.

In this paper we introduce a solvation chemical potential $\mu_{\text{sol}}(\phi)$ for each hydrophilic or hydrophobic charged particle, where $i$ represents the particle species and $\phi$ is the water composition. It is a thermodynamic quantity obtained after statistical averaging over the thermal fluctuations. With formation of a well-defined solvation shell, the typical magnitude of $\mu_{\text{sol}}$ much exceeds the thermal energy $k_B T$. In his original theory, Born took into account the polarization of polar molecules around a spherical ion using a continuum theory of electrostatics [36]. It is the space integral of the electrostatic energy $\varepsilon E^2/8\pi$, where $E = -\nabla \Phi$ in terms of the potential $\Phi$. Here $\Phi = Z_i e/\varepsilon r$ depends on the distance $r$ from the ion center with $Z_i e$ being the charge. The classic Born formula thus obtained is written as

$$\mu_{\text{sol}}(\phi) = Z_i^2 e^2/2 R_{\text{ion}} e \varepsilon = k_B T Z_i^2 \ell_B^2/2 R_{\text{ion}}^2, \tag{2.1}$$

which is applicable for hydrophilic small ions. Here the linear dielectric constant $\varepsilon$ is used, though the electric field in the vicinity of a small ion is strong beyond the linear response level (leading to dielectric saturation). The space integral is in the region $R_{\text{ion}} < r < \infty$, where the lower cut-off $R_{\text{ion}}$ is called the Born radius of order 1 Å for small metallic ions [4, 37]. In terms of the Bjerrum length $\ell_B = e^2/\varepsilon k_B T$, the Born form $\mu_{\text{sol}}(\phi)$ itself considerably exceeds the thermal energy $k_B T$ for $R_{\text{ion}} < \ell_B$. Here $\ell_B$ is 7 Å for room-temperature water and is longer for less polar solvents. The Born formula is in accord with the fact that solvation is stronger for smaller ions. It also indicates that solvation is much intensified for multivalent ions such as Ca$^{2+}$ or Al$^{3+}$. For mixture solvents, the dielectric constant $\varepsilon = \varepsilon(\phi)$ depends on the composition $\phi$ of the more polar component A [38], changing from $\varepsilon_B$ of the less polar component B to $\varepsilon_A$ of the more polar component A. Since the Born expression is inversely proportional to $\varepsilon(\phi)$, this indicates strong preferential solvation. However, the Born formula is known to be very crude, neglecting the formation of the shell structure, the density and composition changes (electrostriction), and the nonlinear dielectric effect. Moreover, the solvation should be strongly coupled with the surrounding hydrogen bonds in aqueous mixtures.

![Figure 1](image.png) Illustration of hydration of Na$^+$ surrounded by a shell composed of water molecules in (a) pure water and (b) water–nitrobenzene.
We give a simple argument on the condition for the formation of a well-defined solvation shell composed of several polar molecules. Let the solvent molecular size $a = v_0^{1/3}$ be longer than the ion radius $\sim R_{\text{ion}}$ (where $a \sim 3$ Å for water). For single component polar liquids, we compare the electrostatic energy in the region $r \sim a$ and the thermal energy $k_B T$ to find the criterion

$$A_0 \equiv Z^2 \epsilon_1^2 / 8\pi \epsilon k_B T a \gtrsim 1.$$  \hfill (2.2)

For mixture solvents, a phenomenological theory yields the criterion [10]

$$A \equiv Z^2 \epsilon_1^2 / 8\pi \epsilon k_B T a \gtrsim 1,$$  \hfill (2.3)

for the formation of a well-defined preferential shell. Here the molecular volumes of the two components are assumed to be of the same order $v_0$ and we define

$$\epsilon_1 = \partial \epsilon(\phi) / \partial \phi,$$  \hfill (2.4)

where the derivative is at fixed $T$ and $p$. In these relations, we have estimated the binding energy $\epsilon_b$ of a polar molecule to an ion to be $Ak_B T$ or $A_k k_B T$. Since $\epsilon_b$ is a microscopic energy, these estimations should be very crude (as well as the Born formula). In this theory, the strong solvation regime is realized for $A_0 \gg 1$ or for $A \gg 1$. If the influence on the hydrogen bonding network is neglected, the radius $R_{\text{shell}}$ of a solvation shell grows as $A_0^{1/2} a$ or as $A^{1/4} a$ from the decay of the electrostatic energy density $\epsilon \propto r^{-4}$. Here it is worth noting that the shell radius $R_{\text{shell}}$ plays the role of the hydrodynamic radius $R_0^i$ in the diffusion constant $D_i = k_B T / 6\pi \eta R_0^i$ with $\eta$ being the shear viscosity. As a result, ions with smaller molecular radii ($\sim R_{\text{ion}}^i$) have larger hydrodynamic radii, which is explained from the relation $R_0^i \equiv R_{\text{shell}}$ [39]. For example, Li$^+$ ions are smaller than Na$^+$ ions, but the diffusion constant of Li$^+$ is smaller than that of Na$^+$ in water.

In electrochemistry, much attention has been paid to ion distribution and the electric potential difference across a liquid–liquid interface [39, 40]. Remarkably, water molecules solvating a hydrophilic ion in the water-rich region remain attached to the ion even when the ion crosses an interface and moves into the oil-rich region. Osakai et al [41] measured the number of water molecules extracted together with hydrophilic ions in a NB-rich phase with a small water composition ($\sim 0.168$ M). In NB–water at room temperature the number of coextracted water molecules in a NB-rich phase was estimated to be 4 for Na$^+$, 6 for Li$^+$, and 15 for Ca$^{2+}$. Furthermore, using proton NMR spectroscopy, Osakai et al [42] studied successive formation of complex structures of anions (such as Cl$^-$ and Br$^-$) and water molecules by gradually increasing the water composition in NB.

These experiments demonstrate that the solvation shell composed of water molecules is stable even when the ambient water composition is much decreased. This is a remarkable result with a number of important consequences. For example, with a small number of hydrophilic ions in oil, let us gradually increase the water composition $\phi$ from zero. If the binding energy $\epsilon_b$ of a water molecule to such an ion is much larger than $k_B T$, we may argue that a hydration shell is formed for [21]

$$\phi > \phi_{\text{sol}} \sim \exp(-\epsilon_b / k_B T),$$  \hfill (2.5)

where the crossover composition $\phi_{\text{sol}}$ is very small for strongly hydrophilic ions. The solvation chemical potential $\mu_{\text{sol}}^i(\phi)$ of hydrophilic ions in water–oil should decrease considerably in the narrow dilute range $0 < \phi < \phi_{\text{sol}}$. Its variation should be milder in the range given in (2.5). The solubility of such ions should also increase abruptly in the same narrow range of $\phi$ with addition of water to oil. Therefore, solubility measurements of hydrophilic ions are informative at small water compositions.

2.2. Gibbs transfer free energy and solvation chemical potentials

We consider a liquid–liquid interface between a polar phase $\alpha$ and a less polar phase $\beta$ with bulk compositions $\phi_{\alpha}$ and $\phi_{\beta}$, across which there arises a difference in $\mu_{\text{sol}}^i$ due to its composition dependence:

$$\Delta \mu_{\text{sol}}^i = \mu_{\text{sol}}^{i\alpha} - \mu_{\text{sol}}^{i\beta}. \hfill (2.6)$$

where $\mu_{\text{sol}}^{i\alpha}$ $(K = \alpha, \beta)$ are the bulk values of the solvent chemical potential of species $i$ in the two phases. In electrochemistry [40, 41], the difference of the solvation free energies $\Delta G_{\text{sol}}^i$ between two phases has been called the standard Gibbs transfer energy. (Since $\Delta G_{\text{sol}}^i$ is usually measured in units of kJ per mole, dividing it by the Avogadro number gives $\Delta \mu_{\text{sol}}^i$ per ion.) It is well known that if there are differences among $\Delta \mu_{\text{sol}}^i$ ($i = 1, 2, \ldots$), an electric double layer emerges at the interface, giving rise to an electric potential jump $\Delta \Phi = \Phi_{\alpha} - \Phi_{\beta}$, across the interface in equilibrium (called the Galvani potential difference). Similar potential differences also appear at liquid–solid interfaces (electrodes) [39]. It is also worth noting that there can be an electric potential difference between two liquids due to orientation of molecular dipoles even without added ions, for which see [43] for water–hexane.

To examine the above effect, let us suppose only two species of ions ($i = 1, 2$) with charges $Z_i e$ and $Z_i e$ ($Z_1 > 0$ and $Z_2 < 0$). At sufficiently low ion densities, the ion chemical potentials are expressed as [39, 40]

$$\mu_i = k_B T \ln(n_i / \lambda_i^Z) + Z_i e \Phi + \mu_{\text{sol}}^i(\phi), \hfill (2.7)$$

for $i = 1, 2$. $\lambda_i$ is the thermal de Broglie length (but is an irrelevant constant in the following). From the charge neutrality in the bulk regions, the bulk ion densities $n_{1\alpha}$, $n_{1\beta}$, $n_{2\alpha}$, and $n_{2\beta}$ satisfy

$$Z_1 n_{1\alpha} + Z_2 n_{2\alpha} = 0, \quad Z_1 n_{1\beta} + Z_2 n_{2\beta} = 0. \hfill (2.8)$$

The continuity of $\mu_i$ across the interface yields

$$k_B T \ln(n_{i\alpha} / n_{i\beta}) + Z_i e \Delta \Phi + \Delta \mu_{\text{sol}}^i = 0, \hfill (2.9)$$
for \(i = 1, 2\). Then the Galvani potential difference is expressed as
\[
\Delta \Phi = \frac{\Delta \mu^1_{\alpha \beta} - \Delta \mu^2_{\alpha \beta}}{\varepsilon (Z_1 + |Z_2|)}. \tag{2.10}
\]
The ion densities in the two phases in the bulk are related by
\[
\ln \left( \frac{n_{\alpha \alpha}}{n_{\beta \beta}} \right) = \ln \left( \frac{n_{\alpha \beta}}{n_{\beta \alpha}} \right) = \frac{|Z_2| \Delta \mu^1_{\alpha \beta} + Z_1 \Delta \mu^2_{\alpha \beta}}{(Z_1 + |Z_2|) k_B T}. \tag{2.11}
\]

Let us take the \(z\) axis perpendicular to a liquid–liquid interface. While the interface thickness is given by the correlation length \(\xi\), the potential \(\Phi(z)\) changes on the scale of the Debye–Hückel screening length, \(\kappa_\text{sp}\), spatial scale of the Debye–Hückel screening length,
\[
Z_k = \kappa_\text{sp} \sqrt{\frac{\varepsilon}{n_k}}.
\]
For monovalent ions, \(\kappa_\text{sp}\) is very small in one of the two phases [11]. Remarkably, \(\Delta \Phi\) in equation (2.10) is independent of the ion densities. It is typically of the order \(10 \kappa_\text{sp} T/e (= 258 \text{ mV})\) for \(\Delta \mu^1_{\alpha \beta} \neq \Delta \mu^2_{\alpha \beta}\). It vanishes for symmetric ion pairs with \(\Delta \mu^1_{\alpha \beta} = \Delta \mu^2_{\alpha \beta}\). As equation (2.11) indicates, the ion densities in the two phases are very different in most cases. For example, if \(\Delta \mu^1_{\alpha \beta} / k_B T = \Delta \mu^2_{\alpha \beta} / k_B T = 10\) in the monovalent case, the common ratio \(n_{\alpha \beta} / n_{\alpha \alpha} = n_{\beta \beta} / n_{\beta \beta}\) becomes \(e^{-10} = 4.5 \times 10^{-5}\). If \(\kappa_{\alpha \beta}\) is macroscopic (as in contact of air with salt water), \(\Delta \Phi\) may not be observable.

In the literature, data for \(\Delta G^1_{\alpha \beta}\) on water–NB at room temperatures are available [40, 41], where water and NB are strongly segregated but a considerable amount water is present in the NB-rich phase (\(\sim 0.168 \text{ M}\)). For aqueous mixtures with \(\alpha\) being the water-rich phase, \(\Delta G^1_{\alpha \beta}\) is negative for hydrophilic ions and positive for hydrophobic ions. For water–NB [40], we have \(\Delta \mu^1_{\alpha \beta} / k_B T = -13.6\) for \(\text{Na}^+\), \(-15.3\) for \(\text{Li}^+\), \(-27.1\) for \(\text{Ca}^{2+}\), \(-11.3\) for \(\text{Br}^{-}\), and \(-7.4\) for \(\text{I}^{-}\) as examples of hydrophilic ions, while we have \(\Delta \mu^2_{\alpha \beta} / k_B T = 14.4\) for hydrophobic \(\text{BPh}_3^-\). In these experiments, a hydration shell should have been formed around each hydrophilic ion even in the water-poor phase \(\beta\). Equation (2.5) then suggests \(\phi_0 > \phi_{\text{water}}\). Though more data are needed, those of \(\Delta G^1_{\alpha \beta}\) for NB–water indicate the composition dependence of \(\mu^1(\phi)\) quantitatively, which is very strong even after the formation of solution shells.

### 3. Ginzburg–Landau theory

We will first give a short summary of our previous results in the Ginzburg–Landau scheme in the presence of preferential solvation in sections 3.1–3.3. New results will then follow on the structure factors among ions, the surface tension behavior with antagonistic salt, and the phase diagram. We treat two species of monovalent ions. However, for three species of ions, we obtain more complex ionic distributions near liquid–liquid interfaces [12, 30] and solid surfaces [26].

#### 3.1. Electrostatic and solvation interactions

We consider a polar binary mixture (water–oil) containing a small amount of monovalent salt \((Z_1 = 1\) and \(Z_2 = -1\)). The ions are sufficiently dilute and their volume fraction is negligible. Hereafter the Boltzmann constant will be set equal to unity. Neglecting the image interaction, we assume the free energy density [10, 11]:
\[
f = f_0(\phi, T) + \frac{T}{2} C |\nabla \phi|^2 + \frac{\varepsilon}{8 \pi} E^2 + T \sum_i \left[ \ln (n_i \lambda_i^{\phi}) - 1 - g_i \phi |n_i| \right]. \tag{3.1}
\]

The first term \(f_0\) is the chemical part. In our simulation we use the Bragg–Williams form
\[
\frac{v_0}{T} f_0 = \chi \phi \ln (\phi) + (1 - \phi) \ln (1 - \phi) + \chi \phi (1 - \phi), \tag{3.2}
\]
where \(v_0\) is the solvent molecular volume common to the two components and \(\chi = \chi (T)\) depends on \(T\) (at constant pressure). Here \(\chi = 2\) at the solvent critical temperature \(T = T_c\) without ions and \(\chi = 2 / \alpha \approx T - T_c\) for small \(T - T_c\). The second term is the gradient part with \(\chi\) being a constant. The third term is the electrostatic free energy, where \(E = -\nabla \Phi\) is the electric field and the potential \(\Phi\) is produced by the charge density \(\rho = \varepsilon (n_1 - n_2)\) via the Poisson equation
\[
\nabla \cdot \varepsilon (\phi) \nabla \Phi = 4 \pi \rho. \tag{3.3}
\]
The dielectric constant \(\varepsilon (\phi)\) depend on the composition \(\phi\) [38]. In our previous work a linear composition dependence
\[
\varepsilon (\phi) = \varepsilon_0 + \varepsilon_1 \phi \tag{3.4}
\]
has been assumed, where \(\varepsilon_0\) and \(\varepsilon_1\) are constants. The last term on the right-hand side of equation (3.1) consists of the entropic part and the solvation part of the ions, with \(\lambda_i\) being the thermal de Broglie length. The solvation terms \((\chi g_i)\) follow if \(\mu^1(\phi)\) depend on \(\phi\) linearly as
\[
\mu^1_{\phi}(\phi) = \text{const.} - T g_i \phi, \tag{3.5}
\]
where the constant term gives rise to a contribution linear in \(n_i\) in \(f\) and is irrelevant, while the second term \((\chi g_i)\) yields the solvation coupling in \(f\). In this approximation, the parameter \(g_i\) represents the strength of the preferential solvation of ion species \(i\). The difference of the solvation chemical potential in two-phase coexistence in equation (2.6) is given by
\[
\Delta \mu_{\phi}^1 = T g_i \Delta \phi, \tag{3.6}
\]
Figure 2. Normalized potential $\Phi(z)/\Delta\Phi$ and composition $\phi(z)$ (upper panel), and normalized ion densities $v_0 n_1(z)$ and $v_0 n_2(z)$ (lower panel) (taken from [11]), where $\chi = 3, g_1 = g_2 = 10$, and $v_0 n_{1a} = v_0 n_{1b} = 2 \times 10^{-4}$. Here $A = 4$ means $e^2/\epsilon_a T = 8.5$. Hydrophilic cations (hydrophobic anions) tend to be on the left (right) near the interface, resulting in a large electric double layer. The asymmetry of the two curves of $n_1$ and $n_2$ is due to the $\phi$ dependence of $\epsilon(\phi)$ ($\epsilon_1/\epsilon_0 = 4/3$ here).

Figure 3. Surface quantities $\sigma, 2\sigma_a, 2\sigma_b,$ and $\sigma_0 - T\Gamma + \sigma_1$ in units of $T a^{-2}$ versus $v_0 n_\alpha$ for a monovalent, antagonistic salt with $g_1 = -g_2 = 10$ at $\chi = 2.1$. Here $2\sigma_a$ and $\sigma_0 - T\Gamma + \sigma_1$ nearly coincide to support equation (3.36).

3.2. Dynamic equations

We present the dynamic equations for $\phi, n_1, n_2,$ and the velocity field $v$ [16, 44]. The fluid is incompressible and isothermal, so we assume $\nabla \cdot v = 0$ and treat the mass density $\rho_0$ and the temperature $T$ as constants. Then $\phi$ and $n_i$ obey

$$\frac{\partial \phi}{\partial t} + \nabla \cdot (\phi v) = -\frac{L_0}{T} \nabla^2 \delta F,$$  \hspace{1cm} (3.7)

$$\frac{\partial n_i}{\partial t} + \nabla \cdot (n_i v) = \frac{D_i}{T} \nabla^2 n_i - g_i n_i \nabla \phi,$$ \hspace{1.1cm} (3.8)

The kinetic coefficient $L_0$ and the ion diffusion constants $D_1$ and $D_2$ are constants. The symbol $\pm$ in the second line of equation (3.8) denotes $-$ for the cations ($i = 1$) and $+$ for the anions ($i = 2$). The momentum equation is expressed as

$$\frac{\partial \rho_0}{\partial t} = -\nabla p_1 + \eta \nabla^2 v - \nabla \cdot \Pi,$$ \hspace{1.1cm} (3.9)

where $p_1$ ensures $\nabla \cdot v = 0$, $\eta$ is the shear viscosity, and $\Pi = \{\Pi_{\alpha\beta}\} (\alpha, \beta = x, y, z)$ is the reversible stress tensor of the form

$$\Pi_{\alpha\beta} = T C \nabla_{\alpha} \phi \nabla_{\beta} \phi - \frac{\delta}{4\pi} E_{\alpha} E_{\beta},$$ \hspace{1.1cm} (3.10)

with $\nabla_{\alpha} = \partial/\partial x_{\alpha}$. The relation $\nabla \cdot \Pi = \phi \nabla (\delta F/\delta \phi) + \sum_n n_i \nabla (\delta F/\delta n_i)$ follows, which is needed for the self-consistency of our model dynamic equations [16, 44].

In numerically integrating the dynamic equations, we may use the following two approximations [16]. First, in the Stokes approximation we set the right-hand side of equation (3.9) equal to zero. Then $v$ is expressed in terms of the Oseen tensor as inversely proportional to the viscosity $\eta$ [44]. Second, in the quasi-static approximation, we assume that the characteristic ion diffusion time is much faster than the relaxation time of the composition. This is justified near the solvent critical point and in describing slow domain growth. In such cases, the ion

$\chi = 3$ (see [11] for more details), where we can see a marked electric double layer. The bulk cation densities in the two sides, $n_{1a}$ and $n_{1b}$, coincide from equation (2.8) and are set equal to $2 \times 10^{-4} v_0^{-1}$. The interface thickness $\xi$ is of order $5a$, while $\Phi(z)$ changes on the scale of $\kappa_a^{-1} \sim \kappa_{\beta}^{-1} \sim 10a$. The surface tension change due to salt is $\Delta \sigma = \sigma - \sigma_0 = -0.0417/a^2$ (where $\sigma_0 = 0.497T/a^2$ without salt) and the surface adsorption of ions is $\Gamma = 0.014a^{-2}$ (see equation (3.33)). If $g_1 = -g_2$, the electric double layer is increasingly enhanced with further increasing $g_1$ and/or $n_{1a}$, essentially leading to vanishing of the surface tension (see figure 3). We can understand these features by solving the nonlinear Poisson–Boltzmann equation [12] (see also a similar recent calculation of the interface profiles [19]). The ionic interfacial distribution was measured for a ternary, antagonistic salt in water–NB in the experiment by Luo et al [30] and was theoretically explained in our previous paper [12].
distributions are expressed in terms of $\phi$ and $\Phi$ in the modified Poisson–Boltzmann relation [11]

$$n_i = n_i^0 \exp[g_i \phi \mp e \Phi/T].$$  \hspace{1cm} (3.11)

This form was also used in analysis of experimental data [30] and in that of molecular dynamics simulation [25]. The coefficients $n_i^0$ are determined from the conservation of the ion numbers

$$\langle n_i \rangle = \int d\mathbf{r} n_i(\mathbf{r}) = n_0,$$  \hspace{1cm} (3.12)

where $\langle \cdots \rangle = V^{-1} \int d\mathbf{r} \langle \cdots \rangle$ denotes the space average with $V$ being the cell volume. The average density $n_0 = \langle n_1 \rangle = \langle n_2 \rangle$ is a given constant. It will be measured in units of $v_0^{-1}$ or the normalized density $v_0 n_0$ will be used with $v_0$ appearing in equation (3.2).

3.3. Structure factors and effective ion–ion interaction in one-phase states

We suppose small deviations $\delta \phi(\mathbf{r}) = \phi(\mathbf{r}) - \langle \phi \rangle$ and $\delta n_i(\mathbf{r}) = n_i(\mathbf{r}) - n_0$ of the composition and the ion densities in one-phase states. Their equilibrium distributions are treated to be Gaussian in the mean field theory. Then, from the free energy (equation (3.1)), we may readily calculate the following structure factors:

$$S(q) = \langle (\phi q_i)^2 \rangle_e, \quad G_{ij}(q) = \langle n_{iq} n_{jq}^* \rangle_e / n_0,$$
$$G(q) = \langle (\rho q_i)^2 \rangle_e / e^2 n_0,$$  \hspace{1cm} (3.13)

where $\phi_q$, $n_{iq}$, and $\rho_q$ are the Fourier components of the composition $\phi$, the ion densities $n_i$, and the charge density $\rho = e(n_1 - n_2)$ with wavevector $q$ and $\langle \cdots \rangle_e$ denotes the thermal average. From (3.13) we have the following relation:

$$G(q) = G_{11}(q) + G_{22}(q) - 2G_{12}(q).$$  \hspace{1cm} (3.14)

The $\phi$ dependence of the dielectric constant $\varepsilon$ is negligible for small fluctuations. We introduce the Bjerrum length $\ell_B$ and the Debye wavenumber $\kappa$ by

$$\ell_B = e^2 / \varepsilon T, \quad \kappa = (8 \pi \ell_B n_0)^{1/2}.$$  \hspace{1cm} (3.15)

After eliminating the ion fluctuations, the second-order contribution of the free energy is written as $\delta F^{(2)} = T \sum_q |\phi_q|^2 / 2V S(q)$. The inverse of $S(q)$ is written as [10, 11]

$$\frac{1}{S(q)} = \tilde{r} - (g_1 + g_2)^2 n_0 / 2 + C q^2 \left[ 1 - \frac{\gamma_p^2 \kappa^2}{q^2 + \kappa^2} \right],$$  \hspace{1cm} (3.16)

where $\tilde{r}$ is related to the second derivative of $f_0$ and use of equation (3.2) yields

$$\tilde{r} = f_0'' / T = v_0^{-1} [1/\phi(1 - \phi) - 2\chi].$$  \hspace{1cm} (3.17)

The parameter $\gamma_p$ represents asymmetry of the solvation of the two ion species and is defined by

$$\gamma_p = (16 \pi C \ell_B)^{-1/2} |g_1 - g_2|. $$  \hspace{1cm} (3.18)

Remarkably, $\gamma_p$ is independent of the salt density. Note that the structure factor for weakly ionized polyelectrolytes has the same form [2, 3], where the solvation has been neglected, however. Recently, we have presented a generalized form of the structure factor for polyelectrolytes including the solvation interaction and the ionization fluctuations [20].

We explain implications of the structure factor (equation (3.16)) as follows. (i) The second term gives rise to a shift of the spinodal curve. For example, if the cations and anions are hydrophilic and $g_1 \sim g_2 \sim 15$, the shift term is of order $-500 n_0$ and its magnitude can be appreciable even for $v_0 n_0 \ll 1$. It is well known that the coexistence curve and the spinodal curve of aqueous mixtures are much shifted even by a small number of hydrophilic ions such as Na$^+$ and Br$^-$ [45]. (ii) On the other hand, $\gamma_p$ can be increased for antagonistic salt [10, 11, 31]. From the last term in equation (3.16) a Lifshitz point appears at $\gamma_p = 1$. (In figure 2, $\gamma_p = 5/4\sqrt{3} < 1$.) That is, while $S(q)$ is maximum at $q = 0$ for $\gamma_p < 1$, it exhibits a peak at an intermediate wavenumber $q_m = (\gamma_p - 1)^{1/2} \kappa$ for $\gamma_p > 1$. The peak height is given by $S(q_m) = 1 / (\tilde{r} - r_m)$, where

$$r_m = (g_1 + g_2)^2 n_0 / 2 - C(\gamma_p - 1)^2 \kappa^2.$$  \hspace{1cm} (3.19)

For $r < r_m$, mesophase formation should take place (see figure 4). For $r > r_m$, two-phase states with a planar interface can be stable or metastable even for $\gamma_p > 1$. (iii) If $\gamma_p < 1$, the criticality and the macroscopic phase separation can remain in the mean field theory. That is, for $q \ll \kappa$, we obtain the Ornstein–Zernike form

$$S(q) \cong 1 / [\tilde{r} - (g_1 + g_2)^2 n_0 / 2 + C(1 - \gamma_p^2) q^2].$$  \hspace{1cm} (3.20)
The coefficient in front of $q^2$ is reduced by the factor $1 - \gamma_0^2$, which leads to enhancement of $S(q)$ in the range $\xi^{-1} < q < \kappa$. When $\kappa > \xi^{-1}$, the correlation length $\xi$ should be redefined by

$$\xi^2 = C(1 - \gamma_0^2)/(F - (g_1 + g_2)^2) n_0. \quad (3.21)$$

Next, retaining the ion densities, we eliminate the composition fluctuations in $F$ by setting

$$(\vec{r} + C q^2) \phi_q = \sum_i g_i n_{i_q}, \quad (3.22)$$

where the terms nonlinear in $\delta \phi$ are neglected. We then obtain effective interactions among the ions mediated by the composition fluctuations. The resultant free energy of ions is written as

$$F_{\text{ion}} = \int d\vec{r} \sum_i T n_i \ln(n_i \lambda_i^3) + \frac{1}{2} \int d\vec{r} \int d\vec{r}' \sum_{ij} V_{ij}(\vec{r} - \vec{r}') \delta n_i(\vec{r}) \delta n_j(\vec{r}'), \quad (3.23)$$

where the terms linear in $n_i$ are not written explicitly. Here the deviations $\delta n_i = n_i - \langle n_i \rangle$ need not be very small and the logarithmic terms are not expanded with respect to them. The effective interaction potentials $V_{ij}(\vec{r})$ are expressed as [11]

$$V_{ij}(r) = \frac{Z_i Z_j e^2}{4 \pi \kappa C r} \exp(-r/\xi), \quad (3.24)$$

where $Z_i e$ and $Z_j e$ are the ion charges (being $\pm e$ in the monovalent case) and $\xi = (C/\kappa)^{1/2}$ is the correlation length without ions (different from $\xi$ in equation (3.21)).

The solvation-induced interaction is effective in the range $a \lesssim r \lesssim \xi$ and can be increasingly important on approaching the solvent criticality (for $\xi \gg a$) which is explained in the following. (i) It is attractive among the ions of the same species ($i = j$) and dominates over the Coulomb repulsion for

$$g_i^2 > 4 \pi \kappa C \epsilon_B. \quad (3.25)$$

Under this condition there should be a tendency of ion aggregation of the same species. (ii) In the antagonistic case ($g_1 g_2 < 0$), the cations and anions can repel one another in the range $a \lesssim r \lesssim \xi$ for

$$|g_1 g_2| > 4 \pi \kappa C \epsilon_B, \quad (3.26)$$

which triggers charge density waves near solvent criticality.

The ion structure factors $G_{ij}(q)$ in equation (3.13) can be calculated from $F_{\text{ion}}$ in equation (3.23). The inverse of the $2 \times 2$ matrix $G_{ij}(q)$ is written as

$$G_{ij}(q) = \delta_{ij} + n_0 V_{ij}(q)/\Gamma, \quad (3.27)$$

where $V_{ij}(q)$ is the Fourier transformation of $V_{ij}(r)$ in equation (3.24). In the monovalent case, some calculations give the desired results:

$$\frac{G_{11}(q)}{G_{00}(q)} = 1 + n_0 S(q) \left[ g_1^2 + g_2^2 - \frac{(g_1 - g_2)^2}{2(u + 1)} - \frac{2g_2^2u}{2u + 1} \right],$$

$$G_{12}(q) = \frac{1}{2} + \frac{n_0}{4} S(q)(g_1 + g_2)^2 - \frac{1}{4} C(q),$$

$$C(q) = \frac{2u}{u + 1} + n_0 S(q) \frac{(g_1 - g_2)^2}{(u + 1)^2} u^2, \quad (3.28)$$

where $u = q^2/\kappa^2$ and

$$G_0(q) = \frac{u + 1/2}{u + 1} = \frac{q^2 + \kappa^2/2}{q^2 + \kappa^2} \quad (3.29)$$

is the structure factor for the cations (or for the anions) in the absence of solvation. The solvation parts in equation (3.28) are all proportional to $n_0 S(q)$, where $S(q)$ is given by equation (3.16), so they diverge on approaching the spinodal discussed below equation (3.18). The Coulomb interaction suppresses large-scale charge density fluctuations, so $C(q)$ tends to zero as $q \to 0$.

### 3.4. Surface tension

Addition of an antagonistic salt can dramatically reduce the surface tension $\sigma$ of liquid–liquid interfaces even at small content [12]. This trend was in fact observed [29, 30]. Here the behavior of $\sigma$ will be examined in some detail as a continuation of [11], but the image interaction will be neglected.

We suppose a planar liquid–liquid interface at $z = 0$ taking the $z$ axis in its perpendicular direction. We impose the boundary condition $\phi \to \phi_0$ in phase $\alpha$ and $\phi \to \phi_p$ in phase $\beta$ far from the interface and minimize the grand potential $\Omega = \int d\vec{r} \omega$. The grand potential density $\omega$ reads

$$\omega = f - h\phi = \sum_i \mu_i \epsilon_i, \quad (3.30)$$

where $f$ is the free energy density in equation (3.1) and the chemical potentials $\mu_i = \delta F/\delta \phi$ and $\epsilon_i = \delta F/\delta n_i$ are homogeneous constants. The relation $d\omega/dz = 2C\phi/\phi'' - d(\rho\Phi)/dz$ is then derived and is integrated to give

$$\omega = C\phi^2 - \rho\Phi + \omega_\infty. \quad (3.31)$$

where $\Phi' = d\phi'/dz$ and $\phi'' = d^2\phi'/dz^2$. Since $\phi'$ and $\rho$ tend to zero far from the interface, $\omega(z)$ tends to a common value $\omega_\infty$ as $z \to \pm \infty$. The surface tension is written as

$$\sigma = \int dz [\omega(z) - \omega_\infty] = \int dz \left[ C\phi^2 - \frac{\xi}{4\pi} E^2 \right]. \quad (3.32)$$

In our previous work [11, 12], we expanded $\sigma$ with respect to the salt density as

$$\sigma \cong \sigma_0 - T \Gamma = \int dz \frac{\xi}{8\pi} E^2, \quad (3.33)$$

where $\sigma_0$ is the surface tension without solute and $\Gamma$ is the surface adsorption of ions. In terms of the total ion density $n = n_1 + n_2$, it is defined by

$$\Gamma = \int dz \left[ n - n_0 - \frac{\Delta n}{\Delta \phi} (\phi - \phi_0) \right], \quad (3.34)$$

where $n_K = n_1 + n_2$ ($K = \alpha$ or $\beta$) and $\Delta n = n_\alpha - n_\beta$, so the integrand tends to zero as $z \to \pm \infty$. The expansion (equation (3.33)) is valid up to linear order in the salt density. For a dilute neutral solute, it becomes the well-known Gibbs formula $\sigma \cong \sigma_0 - T \Gamma$ [46]. The electrostatic contribution to the surface tension has been neglected in the Gibbs formula.
but it can be crucial for an antagonistic salt [12] and for ionic surfactants [22].

We have numerically examined these relations. To this end, we introduce the areal densities of the electrostatic energy and the gradient free energy as

\[ \sigma_e = \int dz \varepsilon E^2 / 8\pi, \quad \sigma_g = \int dz C\phi^2 / 2. \]  

(3.35)

Then equations (3.32) and (3.33) yield \( \sigma = 2\sigma_g - 2\sigma_e \) and

\[ 2\sigma_g \equiv \sigma_0 - T\Gamma + \sigma_e. \]  

(3.36)

In figure 3, we show the surface quantities \( 2\sigma_e, 2\sigma_g, \sigma = 2\sigma_g - 2\sigma_e \), and the combination \( \sigma_0 - T\Gamma + \sigma_e \) as functions of the bulk ion density \( n_\alpha \), where \( g_1 = -g_2 = 10, \chi = 2.1, \) and \( \chi_p = 3.455 \). Here \( n_\alpha = n_{1\beta} \) for \( g_1 = -g_2 \). The surface tension vanishes at \( \nu_0 n_\alpha = 2.5 \times 10^{-4} \), where \( \sigma_g = \sigma_e \). The approximate relation (equation (3.36)) holds excellently here, where its deviation is at most of the order \( 10^{-3}T/\mu^3 \).

3.5. Phase diagram with an antagonistic salt

With the addition of an antagonistic salt, the consolute criticality of binary mixtures disappears due to the emergence of mesophases. However, we have not fully understood the phase behavior even in the mean field theory. So far we have found: (i) instability of one-phase states for \( \chi_p > 1 \) in the range \( \bar{r} < r_m \), as discussed around equation (3.19), and (ii) vanishing of the surface tension \( \sigma \) with increasing salt density as in figure 3 or on approaching the solvent criticality. Thus, in figure 4, mesophases should be formed in the area with \( \chi < 2.14 \) and \( r < r_m \) in the \( \phi-\chi \) plane. Here \( g_1 = -g_2 = 15 \) at \( \nu_0 n_\alpha = 4 \times 10^{-4} \).

4. Simulations in two dimensions with an antagonistic salt

We have already presented 2D simulation results of the phase separation of a binary mixture with a monovalent, antagonistic salt at the critical composition \( \langle \phi \rangle = 0.5 \) [16]. Here we present further results in two dimensions at \( \langle \phi \rangle = 0.5 \) and 0.4. The composition patterns are analogous to those well known for block copolymers, surfactant systems, and magnetic fluids [44, 47, 48]. In our ionic case, however, we shall encounter some unique features in the droplet patterns. We stress that three-dimensional simulations are needed to really understand the phase ordering in this system. For example, we should reproduce the observed onion structure [32].

4.1. Background to the simulation

We vary the interaction parameter \( \chi \) and set the average ion density \( n_0 = (n_1 + n_2) / 2 \) equal to 0.002\( v_0^{-1} \) or 0.003\( v_0^{-1} \). The other parameters are the same as in our previous work [16]. Namely, we set \( C = a^{-1}, g_1 = -g_2 = 15, \epsilon_1 = 0, \) and \( \epsilon_0 = 3a. \) Then we obtain \( \chi_p \approx 2.44 \) from equation (3.18). These values of \( g_1 \) and \( g_2 \) are realistic in view of the data for the Gibbs transfer free energy, as discussed below equation (2.11). On a 256 \( \times \) 256 square lattice, we integrated the dynamic equation (3.7) using the Bragg–Williams free energy density \( f_0 \) in equation (3.1), the Stokes approximation for the velocity field \( \nu \), and the modified Poisson–Boltzmann expressions (equation (3.12)) for the ion densities \( n_1 \) and \( n_2 \). At \( t = 0 \), we started with the initial condition \( \phi(r, 0) = \langle \phi \rangle \) with small random numbers superimposed. Subsequently, phase ordering took place for the parameter values given below. The space mesh size was \( a = v_0^{-1/3} \) and the time step of integration was 0.01\( t_0 \), where \( t_0 \) is related to the kinetic coefficient \( L_0 \) in equation (3.7), and the viscosity \( \eta \) in equation (3.9) as

\[ t_0 = a^5 / L_0, \quad L_0 = 0.16a^3 T / \eta. \]  

(4.1)

Time-evolution of the composition is measured in units of \( t_0 \). At late stages it is much slower than \( t_0 \).

4.2. Slow coarsening and patterns

In figure 5, we plot the (angle-averaged) structure factor \( S(q, t) \) of the composition versus \( q \) for \( \langle \phi \rangle = 0.4 \) at \( t = 10^4 t_0 \). It exhibits a sharp peak at \( q \approx q_m(t) \). In our previous work [16], a similar plot of \( S(q, t) \) was given for \( \langle \phi \rangle = 0.5 \). For each simulation run, we calculate the characteristic wavenumber \( q_m(t) \) by

\[ q_m(t) = \sum_q |q| \phi_q |^2 / \sum_q |\phi_q |^2. \]  

(4.2)

where \( \phi_q \) is the Fourier component of \( \phi(r, t) \). In figure 6, we plot the characteristic domain size \( 2\pi / q_m(t) \) versus \( t \). For all \( \chi \) at \( \langle \phi \rangle = 0.5 \) and for \( \chi = 2.0 \) and 2.1 at \( \langle \phi \rangle = 0.4 \), the coarsening is nearly stopped at long times. For \( \chi > 2.2 \) at \( \langle \phi \rangle = 0.4 \), it is considerably slowed down at long times and, if we write \( q_m \sim t^{\gamma} \), the exponent \( \gamma \) increases from zero up to about 0.1 at \( \chi = 2.4 \). Note that the typical domain size grows as \( t^{3/4} \) for a conserved order parameter without the hydrodynamic interaction and is faster with the hydrodynamic interaction [44, 49].

We next show composition patterns at \( t = 10^4 t_0 \). From figure 6, they are frozen or very slowly evolving. In figure 7,
they are stripe-like and bicontinuous for $\langle \phi \rangle = 0.5$. In figure 8, they are droplet-like for $\langle \phi \rangle = 0.4$. The gradation of each snapshot represents the local value of $\phi(r, t)$. For $\langle \phi \rangle = 0.5$ (or 0.4), the variance $\langle (\phi - \langle \phi \rangle)^2 \rangle^{1/2}$ is about 0.16 (or 0.13) at $\chi = 2$ and increases to 0.33 (or 0.31) at $\chi = 2.4$. Note that no phase separation occurs for $\chi \leq 2$ without ions. We recognize that the patterns are very sensitive to the values of $\chi$ and $v_0 n_0$. A marked feature is that the largest domain size increases with increasing $\chi$ and decreases with increasing $n_0$. More details are as follows. (i) At $\langle \phi \rangle = 0.5$ in figure 7, the patterns are lamellar-like for $\chi = 1.9$, while they resemble those in systems with competing interactions [47, 48]. (ii) In the droplet patterns at $\langle \phi \rangle = 0.4$, water-rich droplets in a percolated oil-rich region have considerable size dispersity. This is possible only when even small droplets have long life times. (Without ions, small droplets quickly disappear due to the surface tension effect.) Large and small droplets segregate exhibiting polycrystal-like medium-range order for $\chi = 2$, while larger droplets are growing and smaller ones are diminishing very slowly in random configurations for $\chi \geq 2.2$.

In figure 9, at $\langle \phi \rangle = 0.4$, we display the composition $\phi(r, t)$, the ion densities $n_1(r, t)$ and $n_2(r, t)$, and the potential $\Phi(r, t)$ at $t = 10^3 t_0$ for the two cases $\chi = 2.1$ and 2.4. Similar snapshots at $\langle \phi \rangle = 0.5$ can be found in our previous paper [16]. The typical droplet radii are a few times larger for $\chi = 2.4$ than for $\chi = 2.1$, but small droplets (with diameter $\sim 5 \sigma$) still remain in the two cases, giving rise to sharp peaks in the cation distribution. Due to the strong selective solvation, the cations are within the droplets and anions are outside them. For $\chi = 2.1$, the typical droplet size is small and the cation density is highest at the droplet centers. For $\chi = 2.4$, the cations and anions are mostly localized in the electric double layers at the interfaces. In these cases, the water-rich droplets are positively charged and are repelling one another. However, we have not yet examined how this repulsion can influence the coarsening process.

4.3. Effective surface tension

In figure 10, we show the time-development of the effective surface tension defined by

$$\sigma_{\text{eff}} = \frac{1}{S} \int dr \left( C |\nabla \phi|^2 - \frac{\varepsilon}{4\pi} \right).$$  (4.3)

where $S$ is the total interface length calculated separately. This quantity becomes the usual surface tension for a planar interface in equilibrium from equation (3.32). (In our previous work [16], the space averages of the gradient free energy density and the electrostatic energy density were separately plotted for $\langle \phi \rangle = 0.5$ in figure 10 there.) We recognize that $\sigma_{\text{eff}}$ tends to zero for bicontinuous domain morphology but remains positive for droplet morphology. It is rather surprising that the coarsening in the droplet case is considerably slowed down even for positive $\sigma_{\text{eff}}$. To explain these results theoretically, we need a more systematic analysis of the problem.

5. Summary and remarks

In this work, we have demonstrated the crucial role of the preferential solvation of ions in phase transitions of polar binary mixtures. Together with a short summary of our previous work, we have presented some new results. (i) In equation (3.28), we have calculated the structure factors among the ion density fluctuations. (ii) In figure 3, we have examined how the surface tension tends to zero with increasing density of the antagonistic salt. (iii) In figure 4, we have shown a phase diagram of a binary mixture containing an antagonistic salt, where mesophases are realized near the solvent criticality. Note that the original criticality without ions disappears when an antagonistic salt with $\gamma_p > 1$ is added. (iv) In figures 5–10, we have numerically examined the phase ordering of a binary mixture containing an antagonistic salt for four values of $\chi$ in the bicontinuous case at $\langle \phi \rangle = 0.5$ and in the droplet case at $\langle \phi \rangle = 0.4$. The coarsening is markedly slowed down with decreasing $\chi$ to 2 and/or with increasing ion density $n_0$. The size dispersity of water-rich droplets is significant, because even small droplets have very long life times. In figure 9, we have found that water-rich droplets are positively charged with

![Graph](image-url)
Figure 7. Composition patterns of a binary mixture at $\langle \phi \rangle = 0.5$ containing an antagonistic salt with $v_0n_0 = 0.02$ (top) and 0.03 (bottom) at $t = 10^4 t_0$, where (a) $\chi = 1.9$, (b) 2.1, (c) 2.2, and (d) 2.4. These patterns are frozen in time.

Figure 8. Composition patterns of a binary mixture at $\langle \phi \rangle = 0.4$ containing an antagonistic salt with $v_0n_0 = 0.02$ (top) and 0.03 (bottom) at $t = 10^4 t_0$, where (a) $\chi = 2.0$, (b) 2.1, (c) 2.2, and (d) 2.3. Patterns are frozen for (a) and (b) and are very slowly evolving for (c) and (d). The size dispersity of droplets is conspicuous.

the anions in the percolated oil-rich region. In figure 10, the effective surface tension $\sigma_{\text{eff}}$ versus $t$ is plotted for $\langle \phi \rangle = 0.5$ and 0.4. It tends to zero for $\langle \phi \rangle = 0.5$ but remains positive for $\langle \phi \rangle = 0.4$.

In figures 7 and 8, the domain configurations exhibit some mesoscopic order (lamellar-like or polycrystal-like) for $\chi$ close to 2, while they remain disordered for larger $\chi$ in our simulation time. In this paper, we cannot conclude whether or not the domain growth has ultimately stopped or not. If thermal noise is present, the system might be slowly approaching lamellar or crystalline states at very long times.

In our theory, the molecular volumes of the two components are commonly $v_0$. However, the molecular volumes of D$_2$O and 3MP (the inverse densities of the pure components) are 28 and 168 Å$^3$, respectively, for example. Moreover, the coefficient $C$ of the gradient free energy is an arbitrary constant. Therefore, our theory remains very qualitative. It is not clear how $C$ can be determined in aqueous mixtures where the hydrogen bonding is of primary importance. For pure water, the observed surface tension outside the critical region $1 - T/T_c \gtrsim 0.1$ agrees fairly well with the calculated surface tension from the
van der Waals model with the gradient free energy density $10Ta^5|\nabla n|^2/2$ [52, 53], where $n$ is the density and $a = 3.1 \text{ Å}$ is the van der Waals radius.

There can be a number of dynamical effects in the presence of an antagonistic salt, including multi-exponential decays of the dynamic light scattering intensity [16] and singular rheological behavior with the appearance of mesophases [44]. We propose experiments to investigate the surface instability taking place for negative surface tension. That is, a water–oil interface becomes unstable with addition of an antagonistic salt above a critical amount (as suggested by figure 3) or on approaching the solvent criticality with a fixed density of antagonistic salt. In the introduction, we have already mentioned the experiments of spontaneous emulsification at a water–oil interface with antagonistic salt [33, 34].

Though this paper has mostly treated antagonistic salts, intriguing solvation effects can well be predicted for aqueous mixtures containing usual hydrophilic ion pairs. For example, many groups have detected long-lived heterogeneities by dynamic light scattering in one-phase states of various aqueous mixtures containing a hydrophilic salt [50]. Recently, we have shown that preferential solvation can stabilize water-rich domains enriched with ions even for $\chi < 2$ when

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**Figure 9.** Composition $\phi$, cation density $n_1$, anion density $n_2$, and potential $\Phi$ (from top to bottom) for $\chi = 2.1$ (left) and $\chi = 2.4$ (right) in a binary mixture at $\langle \phi \rangle = 0.4$ and $n_0v_0 = 0.002$. Snapshots in a quarter of the total system are displayed.
Also presented was which hydration-induced condensation occurs. A Ginzburg–Landau approach to this problem along the lines of was long been a topic of investigation in the literature. Around hydrophilic ions in metastable gas mixtures containing water is the minority component. We also mention the problem of preferential wetting of mixture solvents on ionizable surfaces, which can be either planar, spherical, or cylindrical. Around an ionizable rod, we already predicted a first-order wetting transition with discontinuous jumps in the preferential adsorption and the degree of ionization.

Finally, it is worth noting that ion-induced nucleation has recently been a topic of investigation in the literature. In such situations, we should take into account the solvation interaction among the charges (consisting of the ionized units on the surface and the counterions) and the polar molecules.

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