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Chasing Aqueous Biphasic Systems from Simple Salts by Exploring the LiTFSI/LiCl/H$_2$O Phase Diagram

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Supporting Information

ABSTRACT: Aqueous biphasic systems (ABSs), in which two aqueous phases with different compositions coexist as separate liquids, were first reported more than a century ago with polymer solutions. Recent observations of ABS forming from concentrated mixtures of inorganic salts and ionic liquids raise the fundamental question of how “different” the components of such mixtures should be for a liquid—liquid phase separation to occur. Here we show that even two monovalent salts sharing a common cation (lithium) but with different anions, namely, LiCl and lithium bis(trifluoromethanesulfonfyl)imide (LiTFSI), may result in the formation of ABSs over a wide range of compositions at room temperature. Using a combination of experimental techniques and molecular simulations, we analyze the coexistence diagram and the mechanism driving the phase separation, arising from the different anion sizes. The understanding and control of ABS may provide new avenues for aqueous-based battery systems.

The development of aqueous Li-ion batteries has recently led to the exploration of highly concentrated aqueous electrolyte solutions using organic lithium salts to slow the kinetics of water splitting in the vicinity of the electrodes and therefore enable higher operating voltages.$^{3,5}$ In an attempt to decrease the amount of expensive organic salt used in these aqueous electrolytes,$^4$ we investigated mixtures with an inexpensive lithium salt and could observe an unusual phenomenon, namely the formation of an aqueous biphasic system (ABS). When two liquids with very different physicochemical properties, such as water and oil, contact each other, they usually coexist as separate phases with a sharp interface. This fact has long been exploited, e.g., for the separation of aqueous solutes via liquid—liquid extraction using organic phases or ionic liquids.$^{5,6}$ While the first ABSs based on polymers were discovered in the late 19th century,$^8$ the coexistence of two aqueous solutions of different salts has been reported only much more recently and the range of such ABSs remains comparatively limited.$^{7–11}$ Their composition usually involves a concentrated inorganic salt and an ionic liquid. The resulting phases both contain a sizable amount of water (hence their name), and the origin of the phase separation is thought to result from the simultaneous presence of water hydrogen bond network structuring (kosmotropic) or disordering (chaotropic) ions.$^{10}$ This raises the fundamental question of how “different” the components of the mixture should be for such a liquid—liquid phase separation to occur, i.e., in the aqueous case, for an ABS to form. Here we show that even mixtures of two monovalent salts sharing a common simple cation (lithium, Li$^+$) but with different anions result in the

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presence of water in the coexistence of two aqueous liquid phases over a wide range of compositions at room temperature.

Specifically, we investigate aqueous mixtures of lithium bis(trifluoromethanesulfonyl)imide (LiTFSI), which can form so-called “water-in-salt”\(^1\)–\(^3\) electrolytes (with a salt:solvent ratio of >1 in mass and volume) relevant, e.g., for battery applications, and the more standard lithium chloride (LiCl) salt (see Figure 1A). We show in Figure 1B that a global composition of 12m (moles per kilogram of water) LiCl and 5m LiTFSI results in approximately equal volumes of two coexisting liquids with a sharp interface. The formation of an interface between these two liquids exchanging matter is further evidenced in the Supporting Information by images showing the evolution of a drop of 5m LiTFSI in a saturated (18m) LiCl solution as well as by the measurement of a finite surface tension (5.7 mN/m for the 12m LiCl–5m LiTFSI, approximately an order of magnitude smaller than for the water–air interface).

Nuclear magnetic resonance (NMR) then provides spatially resolved specific information about the chemical environment of the various species. The evolution of the chemical shifts across the interface (Figure 1C) indicates that both Li\(^+\) cations and water molecules adopt distinct local environments in the two liquid phases, while the change in the dielectric constant results in a distortion at the interface between the two phases. In addition, Fourier transform infrared (FTIR) spectroscopy allows us to probe the local environment of water in both phases, which in turn provides indirect information about their composition. As shown in Figure 1D, water molecules from the top phase exhibit O–H bond stretching modes (broad signal around 3400 cm\(^{-1}\)) similar to those in an aqueous 18m LiCl solution, while weak absorbance peaks around 1250 cm\(^{-1}\) reveal traces of TFSI in solution.\(^1\)\(^2\) A similar resemblance between the bottom phase and aqueous 20m LiTFSI can also be deduced from their FTIR spectra.

The liquid–liquid phase separation is further quantitatively examined in Figure 2, which compares the equilibrium density profiles for various elements probed by NMR imaging experiments and computed by molecular dynamics (MD) simulations (see Methods). Despite the difference in length scales probed by both approaches (~1 cm for NMR and ~10 nm for MD), they provide a consistent picture of a sharp interface. The compositions are in good agreement for the species that we could probe experimentally (all but Cl\(^-\) anions), which validates the MD simulations. In particular, MD simulations predict almost quantitatively the ratio of Li\(^+\) concentration in both phases as well as the virtual absence of TFSI\(^-\) in the Cl-rich phase. The predicted water content in the TFSI-rich phase is smaller than in the other, even though the ratio is slightly higher than the experimental one. The density profile for Cl\(^-\) is also readily available from MD and indicates that its concentration in the 

![Figure 1](image1.png)

**Figure 1.** (A) Constituents of the system: water molecule, Li\(^+\) cation, TFSI\(^-\) and Cl\(^-\) anions. (B) Stoichiometric amount of LiTFSI, LiCl, and water used to prepare the 12m LiCl–5m LiTFSI aqueous biphasic system (ABS). (C) Chemical shift imaging (CSI) of a nuclear magnetic resonance tube containing the 12m LiCl–5m LiTFSI ABS, revealing distinct local environments for Li\(^+\) cations and water in the two phases. The color indicates the intensity (from blue for low to red for high) as a function of chemical shift and position, while the shape of the peaks reflects the distortion of the magnetic field near the liquid–liquid and liquid–air interfaces. (D) Fourier transform infrared spectra of the top and bottom phases of the 12m LiCl–5m LiTFSI ABS compared with those of an 18m LiCl solution and a 20m LiTFSI solution.

![Figure 2](image2.png)

**Figure 2.** (A) Density profiles for \(^19\)F, \(^7\)Li, and \(^1\)H obtained by NMR for the 12m LiCl–5m LiCl ABS compared with (B) those obtained by molecular dynamics simulations for Cl\(^-\), TFSI\(^-\), Li\(^+\), and H\(_2\)O for the same system. On the top of panel B, a snapshot extracted from the simulation (Li\(^+\) ions are colored orange, Cl\(^-\) anions green, TFSI\(^-\) anions red, and water molecules are not shown) illustrates the presence of a sharp interface.
The formation of an ABS, i.e., phase separation between two liquid phases, depends on the global composition of the system. The boundary between the monophasic and biphasic regions of the phase diagram, determined by cloud point measurements (see Methods), is shown in Figure 3. With respect to the system (12m LiCl and 5m LiTFSI) mentioned above, we also observe a phase separation in the MD simulations for a global composition of 6m and 6m, while for a more dilute system (1m and 1m), the system remains monophasic (see Figure S2). At a global composition of 4m and 4m, the system is close to the transition between the one- and two-phase behavior (see density profiles for 4m and 4m and 6m and 6m in the Supporting Information). The resulting compositions of the coexisting liquids in the phase-separating cases, obtained from the MD density profiles (see the Supporting Information), are very close to the experimental phase boundaries. Such an agreement is remarkable considering the complexity of the system. This further supports the validity of the MD simulations, which in turn complements the experimental phase boundaries with the tie lines (which could in principle be obtained by separating and weighting both phases but would require larger volumes and therefore be much more expensive).

We finally turn to the physical origin of the phase separation. In contrast to previously reported ABSs, in the systems presented here the salts share a common cation. In agreement with previous MD studies of bulk WiSEs,

The driving force to form an ABS was previously proposed to result from the difference in interaction with water molecules between two salts, one structuring water (kosmotropic) and the other disordering the hydrogen bond network of water (chaotropic),

However, the concept of cha- and kosmotropicity, which underlines the effect of individual ions on water, should be taken with some caution at such high concentrations. This is particularly true for WiSEs in which the salt:water ratio is >1, because in this regime the effects of ion—water interactions are not additive.

From a thermodynamic point of view, a mixture of a solvent with two salts sharing a common ion can be considered as a ternary mixture, due to the constraint of electroneutrality. Using integral equations, Lo Celso et al. analyzed the phase behavior of such mixtures for neutral (solvent) and charged (ions) hard spheres and found that an asymmetry in the size of the counterions can be sufficient to induce a phase separation, for a range of compositions and temperatures. On the microscopic scale, this means that the liquid structures satisfying both the local electroneutrality and packing constraints with each counterion separately are more stable than for the case in which all ions are mixed. This is not to say that water does not play a role in the mixing free energy, which determines whether the phase separation occurs: It does contribute both energetically (via ion solvation and screening of electrostatic interactions) and entropically (configurational entropy via the composition of each phase). However, this contribution may not be the dominant one. This claim is further supported by the low solubility of alkali halides in conventional ionic liquids.

In this work, we have shown the formation of ABSs formed by mixing salts containing a common cation, namely, LiTFSI and LiCl. With the help of NMR and MD simulations, we demonstrated that for a wide range of compositions, the system segregates into two phases, one being rich in LiTFSI and containing some LiCl and one containing almost exclusively LiCl at high concentrations. While the cha- and kosmotropic phenomenological effect of ions was previously suggested, our findings support the idea that the anion size asymmetry is the driving force for the formation of an ABS in this system. While the first ABS-based flow cell battery was proposed recently,

Experimental methods and details of the MD computer simulation are described in the Supporting Information. No unexpected or unusually dangerous safety hazards were encountered.
 Experimental procedures and simulations details, photographs of a drop of LiTFSI in LiCl aqueous solutions, concentration profiles calculated from MD simulations with different global compositions, solvation of Li⁺ cations in the LiCl and LiTFSI-rich phases, viscosity measurements and Jones–Dole coefficients for LiTFSI, LiCl, and KCl aqueous solutions, and a dual-ion battery using the ABS as the electrolyte (PDF)

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