Direct Imaging of Micrometer Thick Interfaces in Salt-Salt Aqueous Biphasic Systems

Damien Degoulange\textsuperscript{1,2,3}, Raj Pandya\textsuperscript{4,5}, Michael Deschamps\textsuperscript{3,6}, Dhyllan A. Skiba\textsuperscript{7}, Betar M. Gallant\textsuperscript{7}, Sylvain Gigan\textsuperscript{4}, Hilton B. de Aguiar\textsuperscript{4}, Alexis Grimaud\textsuperscript{1,2,3,8*}

1. Chimie du Solide et de l’Energie, Collège de France, UMR 8260, 75231 Paris Cedex 05
2. Sorbonne Université, 75006 Paris, France
3. Réseau sur le Stockage Electrochimique de l’Energie (RS2E), CNRS FR3459, 33 rue Saint Leu, 80039 Amiens Cedex, France
4. Laboratoire Kastler Brossel, ENS-Université PSL, CNRS, Sorbonne Université, Collège de France, 24 rue Lhomond, 75005 Paris, France
5. Department of Physics, Cavendish Laboratory, University of Cambridge, JJ Thomson Avenue, Cambridge CB3 0HE, United Kingdom
6. CNRS, CEMHTI UPR3079, Univ. Orléans, 1D avenue de la Recherche Scientifique, 45071 Orléans, France
7. Department of Mechanical Engineering, Massachusetts Institute of Technology, 77 Massachusetts Avenue, Cambridge, MA 02139, USA
8. Department of Chemistry, Merkert Chemistry Center, Boston College, Chestnut Hill, MA 02467, USA

*Corresponding author: Alexis Grimaud alexis.grimaud@bc.edu

Keywords: aqueous biphasic system / liquid-liquid phase separation / liquid-liquid interface / Gibbs adsorption / Raman imaging
Abstract

Unlike the interface between two immiscible electrolyte solutions (ITIES) formed between water and polar solvents, molecular understanding of the liquid-liquid interface formed for aqueous biphasic systems (ABSs) is relatively limited and mostly relies on surface tension measurements and thermodynamic models. Here, high-resolution Raman imaging is used to provide spatial and chemical resolution of the interface of LiCl-LiTFSI-water and HCl-LiTFSI-water, prototypical salt-salt ABSs found in a range of electrochemical applications. The concentration profiles of both TFSI anions and water are found to be sigmoidal, in agreement with an increasing surface tension as a function of concentration, both being typical of a negative adsorption mechanism. More striking, however, is the length at which the concentration profiles extend, ranging from 11 to 2 μm with increasing concentrations, compared to a few nanometers for ITIES. We thus reveal that unlike ITIES, salt-salt ABSs do not have a molecularly sharp interface but rather form an interphase with a gradual change of environment from one phase to the other. This knowledge represents a major stepping-stone in the understanding of aqueous interfaces, key for mastering ion or electron transfer dynamics in a wide range of biological and technological settings including novel battery technologies such as membraneless redox flow and dual ion batteries.
Significance Statement

Phase separation between oil and water is a well-known process. However, recent findings have shown that phase separation can also take place between two aqueous phases composed of two different dissolved salts. Such aqueous biphasic systems (ABSs) have applications both in life sciences and industrial processes. Yet very little is known about the composition and the structure of their interfaces. Here, through direct Raman imaging, we provide visualization of ABSs interfaces and show that they are made of a concentration gradient along several microns. This large interphase contrasts with the nanometer size, molecularly sharp interface formed between water and oil. Our result lay the foundation for understanding water-water interfaces and mastering interfacial reactions relying on ion and/or electron transfer.
Introduction

Liquid-liquid phase separation (LLPS) lies at the heart of chemistry both in the natural world, e.g. the membraneless separation of dense and dilute phases of macromolecules inside cells\textsuperscript{1–3}, and in industrial systems, e.g. for chemical extraction and purification steps\textsuperscript{4}. One ubiquitous example of LLPS is that of aqueous biphasic systems (ABSs), also called aqueous two phase systems (ATPSs), with a plethora of applications ranging from biomedicine to metal recycling\textsuperscript{5–11}. Typically, ABSs consist of two coexisting immiscible phases composed of only one solvent (water) with the phase separation driven by solutes. Many solutes have been found to drive such phase separation including polymers\textsuperscript{6,11}, proteins\textsuperscript{12} and ionic liquids (IL)\textsuperscript{13}. For most solutes, the phase separation was explained by the hydrophobicity of the large molecules (polymers) or the ability of ions to disturb or structure the hydrogen bonding network in water, i.e. chaotropic and kosmotropic effects respectively\textsuperscript{9,14,15}. Recently, LiX-LiTFSI-water (with $X = \text{Cl}^-$, $\text{Br}^-$ and $\text{I}^-$) salt-salt ABSs, using common alkali cations but anions with large size asymmetry, were reported\textsuperscript{16,17}. Such salt-salt ABSs are regarded as promising alternatives to “classical” liquid electrolytes used in battery technologies. Indeed, they can tackle problematic chemical crosstalk mechanisms observed in Li-ion/sulfur batteries\textsuperscript{18} or even enable the assembly of promising novel battery systems such as membraneless redox flow batteries\textsuperscript{19–21} or dual ion batteries\textsuperscript{22}.

However, for salt-salt ABSs to be efficiently implemented in electrochemical devices, mastering the ion transfer at liquid-liquid interface is critical as it directly controls the (dis)charging rate by inducing an interfacial overpotential if too slow or can induce self-discharge when not selective. Despite this key role, interfaces of ABSs are poorly understood, almost entirely \textit{via} surface
tension measurements. Classically, the surface tension between two phases $\alpha$ and $\beta$ is captured by the Gibbs adsorption equation:

$$\frac{d\gamma}{dV} = -\sum I_j d\mu_j$$  \hspace{1cm} \text{eq. 1}$$

with $\gamma$ being the surface tension between the two phases, $I_j$ the surface excess concentration compared to the Gibbs dividing surface and $\mu_j$ the chemical potential of the different species J in solution. Noting that $\mu_j = \mu_j^0 + RT \ln(a_j)$, the surface excess concentration can be expressed as:

$$I_S = -\frac{a_S}{RT} \left( \frac{\partial \gamma}{\partial a_S} \right)_{T,J \neq S}$$  \hspace{1cm} \text{eq. 2}$$

linking the surface tension to the adsorption of species at the interface. If by increasing the activity of species $S$ the surface tension decreases (i.e. $\left( \frac{\partial \gamma}{\partial a_S} \right) < 0$), then $I_S > 0$ and the species $S$ is accumulated at the interface (positive adsorption – Fig. 1a). However, if by increasing the activity of species $S$ the surface tension increases (i.e. $\left( \frac{\partial \gamma}{\partial a_S} \right) > 0$), then $I_S < 0$ and the species $S$ is depleted at the interface (negative adsorption – Fig. 1b). The thickness of the interface can be defined as the length across which the negative (or positive) adsorption takes place.
Figure 1. Description of the different types of liquid-liquid interfaces. a-b, Schematic depicting the concentration profile of a solute (S) from phase α ($C_s^\alpha$) to phase β ($C_s^\beta$) across an interface for positive (a) and negative (b) adsorption. Green and red represent, respectively, positive and negative integrals when compared to a step change of concentration. c-d, Cartoons of ITIES (c) ABSs (d) interfaces.

In contrast to ABSs, interfaces between two immiscible electrolyte solutions (ITIES), such as between water and nitrobenzene or other polar solvents, have been widely studied. They are known to be molecularly sharp with a positive adsorption of ions over only a few nanometers at the interface (Fig. 1c), as demonstrated by a combination of surface tension, capacitance and surface-sensitive spectroscopic methods.$^{28-31}$ This picture of the ITIES was essential towards developing new applications based on ion$^{32,33}$ or electron transfer$^{34,35}$ such as coupling reactions$^{36}$ or electrosynthesis$^{37}$ at the interface. Thus, for ABSs to reach their full potential, a deeper understanding of their interfaces is critical (Fig. 1d). Yet the requirement of nanoscale chemical sensitivity means, to the best of our knowledge, that no experimental visualization of any aqueous-aqueous interfaces has yet been reported, with information regarding their composition remaining generally sparse. In this work, we use a unique combination of variable temperature nuclear magnetic resonance (VT-NMR) and high-spatial-resolution Raman imaging to resolve and understand the interface of salt-salt ABSs. Studying the LiCl-LiTFSI-water and the HCl-LiTFSI-water systems, we provide the demonstration for a negative adsorption of ions and water at the interface. Strikingly, we reveal a continuous, microscale change of chemical environment, fundamentally different from the nanoscale, molecularly sharp ITIES. Our results call for the reinvestigation of previously known ABSs using chemically sensitive imaging/spectroscopic methods and to, more broadly, study the structure and dynamics of liquid-liquid interfaces.
Results

Phase diagram, surface tension and temperature behavior of the LiCl-LiTFSI-water system

To first understand the composition demixing for the LiCl-LiTFSI-water system, the phase diagram was obtained via the cloud point titration technique (see Materials and Methods). As shown in Fig. 2a, the phase diagram shows a binodal curve separating the monophasic region (purple) from the biphasic one (red). The interfacial tension for the LiCl-LiTFSI-water systems were measured from the drop-weight method (see Materials and Methods) using Tate’s law with Harkins and Brown correction (Fig. 2b). For the 5 m (mol.kg\(^{-1}\)) LiTFSI – 10 m LiCl biphasic system, the interfacial tension is estimated as ≈ 3 mN.m\(^{-1}\). This value is similar to IL-salt ABSs (0.4 to 2.3 mN.m\(^{-1}\)) but ten times lower than values reported for ITIES (20 to 30 mN.m\(^{-1}\)) and higher than that reported for polymer-polymer ABSs (0.01 to 500 μN.m\(^{-1}\)). Decreasing the molalities to 3 m LiTFSI – 6 m LiCl, the interfacial tension decreases down to values ≈ 200 μN.m\(^{-1}\). When plotting the interfacial tension as a function of the difference between salts total molality from these systems and the binodal curve at 2.6 m LiTFSI – 5.2 m LiCl (Δm, Fig. 2b), the surface tension tends to zero when reaching the binodal curve, i.e. for Δm = 0. This result agrees with the system becoming monophasic on crossing the binodal curve, hence having no interfacial tension. According to eq. 2, this decreasing interfacial tension with increasing molality suggests that salts have a negative adsorption at the interface, as schematized in Fig. 1b.
Figure 2. LiCl-LiTFSI-water system description. a, Phase diagram of the LiCl-LiTFSI-water system with the monophasic region in purple and the biphasic region in red. X m LiTFSI - Y m LiCl systems are studied along the line 2X=Y, as indicated with blue dots. Along this line, the binodal curve is crossed at 2.6 m LiTFSI – 5.2 m LiCl. b, Interfacial tensions for the LiCl-LiTFSI-water system as function of the salts total molality differential to the binodal curve in mol per kg (Δm) i.e. Δm=(X+Y)-(2.6+5.2). c-d, Infra-red spectra recorded for the 5m LiTFSI - 10m LiCl system for TFSI vibrations (c) and OH vibrations (d) with top phase in green and the bottom phase in orange. e-f, Infra-red spectra for the 2.6m LiTFSI - 5.2m LiCl system for TFSI vibrations (e) and OH vibrations (f) with the top phase in green and the bottom phase in orange. g, Variable temperature nuclear magnetic resonance analysis of the 2.6m LiTFSI - 5.2m LiCl systems at 1°C (blue), 20°C (yellow) and 35°C (red) for the nucleus 19F (top), 7Li (middle) and 1H (bottom) along the height of the NMR tube from left (top of the tube) to right (bottom of the tube).

The top and bottom phases collected from the 5 m LiTFSI – 10 m LiCl biphasic system were characterized by infrared spectroscopy (Fig. 2c, 2d). By looking at the TFSI peaks at 1340, 1190, 1125 and 1050 cm⁻¹ (Fig. 2c), the top phase is found to be nearly depleted in TFSI anions while the bottom phase is rich in TFSI. In Fig. 2d, the components at 3365 cm⁻¹ and 3230 cm⁻¹ corresponding to OH vibrations for water molecules with strong hydrogen bonding network
(OH\textsubscript{strong-HB}) are only observed for the top phase\textsuperscript{40,41}. In contrast, the bottom phase shows a band at 3570 cm\textsuperscript{-1} previously ascribed to OH vibrations for water with weaker hydrogen bonding network resembling that of the so-called water-in-salt systems, \textit{i.e.} a TFSI-rich environment (OH\textsubscript{weak-HB})\textsuperscript{41,42}. The bottom phase is thus richer in TFSI when compared to the top one, indicating a high ion partition coefficient. By decreasing the concentration close to the binodal curve and studying the 2.6 m LiTFSI – 5.2 m LiCl system (Fig. 2e, 2f), vibrations associated with the top and the bottom phases are found qualitatively similar, however with the intensities of the TFSI peaks still being modestly higher in the bottom phase, thus resulting in a low ion partition coefficient (Fig. 2e). Moreover, Fig. 2f shows that the intensity of the OH\textsubscript{weak-HB} vibration is constant for both phases. Only a slight decrease in OH\textsubscript{strong-HB} vibrations intensity, linked to a weakening of the water-water hydrogen bonding network, is found in the bottom phase. Overall, IR spectroscopy indicates that when moving away from the binodal curve, \textit{i.e.} as the interfacial tension increases, there is greater ion partition between the two immiscible phases.

To probe the influence of temperature on salt-salt ABSs interface, the 2.6 m LiTFSI – 5.2 m LiCl system was studied between 0°C and 35°C. Upon cooling from room temperature to a temperature close to 0°C, the system reversibly switches from biphasic to monophasic, forming a so-called lower critical temperature (T\textsubscript{LC}) system, as confirmed by reaction microcalorimetry (Fig. S1 and Supplementary Discussion 1)\textsuperscript{27,43}. VT-NMR was used to provide spatially resolved information across the ABS interface (z-axis, Fig. S2) and to assess the ion partition as function of temperature (Fig. 2g). At 35°C, the intensity profile for \textsuperscript{19}F shows a sharp difference across the interface, with more TFSI found in the bottom phase than in the top phase (Fig. 2g - right panels in red), in agreement with the infrared spectroscopy data (Fig. 1e, 1f). Similarly, the intensity
profiles for $^7\text{Li}$ and $^1\text{H}$ show the top phase, i.e. the TFSI-poor one, to be richer in lithium and protons. By lowering the temperature to 20°C, the partitions for Li$^+$ and TFSI ions and for water decrease (Fig. 2g - middle panels in yellow). Finally, at 1°C, the system is monophasic and the solution is found to be homogeneous (Fig. 2g - left panels in blue). Therefore, the ion partition decreases when decreasing the temperature, eventually crossing the binodal curve and forming a monophasic system at low temperature. For systems far from the binodal curve, i.e. 5 m LiTFSI – 10 m LiCl, much larger ion and water partitions are recorded at room temperature, as seen in Fig. 2c and 2d, and no transition from biphasic to monophasic was observed between -15°C and 80°C.

**Physical properties of the interface in the LiCl-LiTFSI-water system**

In order to gain a microscopic understanding of the interface composition, Raman imaging was performed (lateral resolution ≈300 nm; axial resolution ≈800 nm, spectral resolution ≈15 cm$^{-1}$), taking advantage of the different phase-dependent vibrations for TFSI anions and water. Fig. 3a shows the heat map of the integrated intensity of the TFSI peak at ≈740 cm$^{-1}$ for the 2.6 m LiTFSI – 5.2 m LiCl system, with increasing TFSI concentration shown from blue to red. Comparing the Raman spectra collected across the interface (Fig. 3b) reveals a continuous change of the spectra, indicating a gradient of chemical compositions and solvation environments (behavior also seen in Fig. 3c for OH vibrations).

The Raman spectra were deconvoluted into their respective components to obtain qualitative chemical insight into the changes occurring across the interface. Based on previous studies$^{42,44}$--
the TFSI band can be decomposed into three components (Fig. 3d, 3e and S3, S4). The TFSI-rich phase is mostly composed of components corresponding to mono- (contact ion pairs (CIP), 738 cm\(^{-1}\)) and bi-coordinated (aggregates (Agg), 743 cm\(^{-1}\)) TFSI ions with a limited contribution from “free” TFSI ions (725 cm\(^{-1}\)). Moving from the TFSI-rich phase (Fig. 3d) to the TFSI-poor phase (Fig. 3e), the intensity of the bi-coordinated component gradually drops alongside that of the free TFSI component, while the intensity of the mono-coordinated TFSI component is still present but at a lower intensity (Fig. S4). Similarly, the OH band can be decomposed into three components\(^40\text{-}42\) (Fig. 3f, 3g and S3, S4): two at 3365 cm\(^{-1}\) and at 3230 cm\(^{-1}\) corresponding to OH\(_{\text{strong-HB}}\) interactions in the water hydrogen bonding network and one component at 3570 cm\(^{-1}\) corresponding to OH\(_{\text{weak-HB}}\) interactions from a TFSI-rich environment. When crossing from the TFSI-rich (Fig. 3f) to the TFSI-poor phase (Fig. 3g), the intensity of the components associated with a strong hydrogen bonding between water molecules increases (OH\(_{\text{strong-HB}}\), low wavenumber components), while the intensity of the component associated with weaker water interactions remains approximately constant (OH\(_{\text{weak-HB}}\), high wavenumber component) (Fig. S4). Together, the Raman peak deconvolutions reveal that the interface is composed of a continuum of solvation environments gradually switching from a solvation structure previously ascribed to water-in-salt electrolytes (TFSI-rich phase with weak hydrogen bonding between water molecules and with TFSI aggregates) to that of a more classical salt-in-water electrolyte (TFSI-poor phase with a stronger hydrogen bonding network between water molecules and without TFSI aggregates). We note that as the polarization of our pump and detected Raman light is poorly defined (owing to imaging with high numerical apertures) we refrain from more quantitative analysis.
By plotting the normalized integrated Raman intensity variation and peak decomposition parameters across the interface for TFSI and OH vibrations, a sigmoidal curve is obtained (Fig. 3h and Fig. S4). This observation confirms that both the solvent, *i.e.* water, and one of the solutes, *i.e.* TFSI anions, have a negative adsorption at salt-salt ABSs interface (eq. 2 and Fig. 1b). More importantly, the thickness of the ABS interface is deduced by measuring the length over which the negative adsorption takes place, which is estimated by the distance necessary to increase from 10% to 90% of the normalized integrated Raman intensity. For the 2.6 m LiTFSI – 5.2 m LiCl system, the thickness is estimated to be 9.1 μm and 11.3 μm from the OH and the TFSI vibrations, respectively. The estimated thickness was found to be stable over hours and between multiple samples, reflecting a steady state (Fig. S5). Furthermore, the curvature of the interfaces is far larger (ten times) than the axial resolution of our laser system (≈0.8 μm) as shown in Fig. S6 where a 3D reconstruction of the different phase regions is built. Consequently, the interfaces can be approximated as a vertical slab with thickness effects ignored.
Figure 3. Raman imaging across the interface of the 2.6m LiTFSI – 5.2m LiCl and 3m LiTFSI – 6m LiCl systems. a, Raman imaging of the 2.6 m LiTFSI – 5.2 m LiCl system generated by integration of the TFSI vibrations. A slight movement of the liquid during imaging results in a partial shift in the image (≈50 μm from bottom); the top and bottom parts of the image are hence not compared. b-c, Raman spectra collected across the interface for TFSI vibrations (b) and for OH vibrations (c) of the 2.6 m LiTFSI – 5.2 m LiCl system. The color of plots matches that in (a). d-e, Peak decomposition of the TFSI vibration in the TFSI-rich phase (d) and TFSI-poor phase (e) of the 2.6 m LiTFSI – 5.2 m LiCl system. f-g, Peak decomposition of the OH vibration in the TFSI-rich phase (f) and TFSI-poor phase (g) of the 2.6 m LiTFSI – 5.2 m LiCl system. h, The normalized intensity of OH (green) and TFSI (orange) vibrations across the interface of the 2.6 m LiTFSI – 5.2 m LiCl system. i, The normalized intensity of OH (green) and TFSI (orange) vibrations across the interface of the 3 m LiTFSI – 6 m LiCl system.

Having established the negative adsorption of TFSI anions and water molecules at the interface of a system close to the binodal curve, similar studies were carried out for systems with greater concentrations. For a 3 m LiTFSI – 6 m LiCl system (Fig. 3i and S7), a gradual change in solvation environment is observed when crossing the interface, similar to that of the 2.6 m LiTFSI – 5.2 m LiCl system. The sigmoidal curve observed for the integrated intensity (Fig. 3i) confirms the negative adsorption of both TFSI and water even at higher molalities. The thickness of the
Interface was estimated to be 2.7 μm and 4.3 μm from the OH and the TFSI vibrations, respectively, smaller than previously measured for the 2.6 m LiTFSI – 5.2 m LiCl system (Fig. 3h). Systems with greater concentrations are reported in Fig. S7, showing an interface of about 3 μm and 2.5 μm for the 4 m LiTFSI – 8 m LiCl and 5 m LiTFSI – 10 m LiCl systems, respectively. Overall, we establish that for a system with a T_{LC}, the interface gets thinner with increasing salt molalities, i.e. moving away from the binodal curve, but remains on the order of few micrometers.

**Interface for upper critical solution temperature: the HCl-LiTFSI-water system**

The above results were then compared with those for HCl-LiTFSI systems (Fig. S8) where VT-NMR (Fig. S9) and reaction microcalorimetry (Fig. S10) measurements reveal an upper critical temperature (T_{UC})^{27,43}, i.e. the system becomes miscible upon heating. As for the LiCl-LiTFSI-water systems, Raman imaging carried out on flat interfaces reveals that the solvation environment changes gradually over a few microns from one phase to the other (Fig. 4). Going from the TFSI-rich to the TFSI-poor phase, the spectral components at lower frequency, corresponding to strong hydrogen bonding, are found to increase for OH vibrations (Fig. 4c and Fig. S11) along with an increase of the high frequency components corresponding to mono- and bi-coordinate TFSI anions (Fig. 4b and Fig. S11). The concentration-position profile for TFSI anions and water across the interface are both found to be sigmoidal in shape (Fig. 4d). Hence, independent of whether a given system has a T_{LC} or a T_{UC}, a negative adsorption is found for the solute and the solvent. Repeating measurements at different concentrations, it is found that the interfacial thickness decreases with increasing molality (≈ 5 μm for 5 m LiTFSI – 10 m HCl system, Fig. 4d and ≈ 2 μm
for 6 m LiTFSI – 12 m HCl system, Fig. S12), as for the LiCl-LiTFSI-water system.

Figure 4. Raman imaging across the interface of the 5m LiTFSI – 10m HCl system. a, Raman imaging of the 5 m LiTFSI – 10 m HCl system generated by integration of the TFSI vibrations. b-c, Raman spectra collected across the interface for the TFSI vibration (b) and for OH vibrations (c) The color of plots matches that in (a). d, The normalized intensity of OH (green) and TFSI (orange) vibrations across the interface.

Our results show that, unlike ITIES, salt-salt ABSs interfaces cannot be considered as molecularly sharp interface. Rather, they are microns-thick interphases forming a continuous gradient between both liquid phases. Given that the interfacial tension of salt-salt ABSs (about few mN.m\(^{-1}\)) lies between that of ITIES (tens of mN.m\(^{-1}\)) and polymer-polymer ABSs interfaces (hundreds of μN.m\(^{-1}\)) which both have interfaces on the sub-300 nm scale, our results show that salt-salt ABSs are outliers. Based on our findings, models that have been developed for ITIES using molecular isotherms, which link molecular interactions, surface tension and interfacial thickness hence do not appear appropriate for describing ABS. One way to qualitatively reconcile our observations is via the Gibbs adsorption equation (eq. 1) which does not take interfacial thickness or molecular interactions into account. Given the negative adsorption experimentally revealed in this study, i.e. \( \Gamma_j < 0 \), the increasing interfacial tension as function of increasing molality is the result of an
increase of the chemical potentials of ions and/or water. However, when considering solutions far from the dilute regime, as in this study, the activity of water decreases when increasing the salts concentration unlike the activities of ions that are increasing, as previously shown. One can thus conclude that the formation of the salt-salt ABSs interface is driven by the activity of ions rather than by that of water.

Summarizing the different systems studied in this work, we find that the thickness of the interface is decreasing with both total molality (Fig. 5a and Fig. S13) and surface tension (Fig. 5b). For large molality, \textit{i.e.} for high surface tension (Fig. 5c), we observe large ion partitioning while Raman imaging reveals thin interfaces. Decreasing the molality, \textit{i.e.} for lower surface tension, the ion partitioning is smaller but the thickness of the interface increases (Fig. 5d). However, a more quantitative interpretation of the Gibbs adsorption equation, relating surface excess to surface tension, requires a simultaneous knowledge of the concentration profiles for all species, currently out of reach for ions such as Cl\textsuperscript{-} or Li\textsuperscript{+} using Raman imaging.
Figure 5. Evolution of interfacial thickness as function of molality and surface tension. a, Comparison of the integrated intensity for TFSI vibrations for different LiCl-LiTFSI-water systems. b, Interfacial thickness as function of the interfacial tension for the LiCl-LiTFSI-water and HCl-LiTFSI-water systems studied in this work. The green line is only a guide to the eye. c-d, Schematic depicting the concentration profile of solutes across an interface for high (c) and low (d) surface tension. Green and red represent, respectively, positive and negative integrals when compared to a step change of concentration.

Conclusion

In summary, with microscopic chemical resolution, we have directly imaged the interface formed for salt-salt ABSs as a function of concentration. Our study reveals, unlike for ITIES or polymer ABSs, very large interfacial thicknesses ranging from 1 μm to 10 μm. The thickness correlates inversely with the salt concentration and the interfacial tension. Using Raman spectroscopy, the molecular bonding environment across the interfaces was probed, demonstrating a gradual change in water structure from an environment in which the hydrogen bonding network is weak and anions aggregates exist to an environment resembling more of a dilute aqueous solution. In addition, using NMR and calorimetry methods, we dynamically probed the effect of temperature
on the miscibility of salt-salt ABSs. Comparing LiCl-LiTFSI-water and HCl-LiTFSI-water systems, we find that changing the cation from Li$^+$ to H$^+$ inverts the temperature-dependence, with the former becoming miscible upon cooling below room temperature while the latter becomes immiscible. Nevertheless, such opposing behavior does not dramatically change the thickness of the interface, which remains in the 1-10 $\mu$m range. Considering a negative adsorption for both water and solutes at the interface, we find that ion partitioning increases with salt concentration. We postulate that the formation of the interface is dictated by increased activity of solute ions with concentration. Following the insights provided by our findings, better theoretical understanding of ABSs is still needed. Indeed, the surface tension of salt-salt ABSs lies between that of ITIES and of polymer-polymer systems, yet they show much larger interfaces, beyond access for classical molecular dynamics simulations. Understanding further how the properties and size of the interfaces can be tuned is not only of fundamental interest but crucial if ABSs systems are to be used for electrochemical devices$^{18-22}$, chemical separation/purification steps$^{10}$, carbon capture$^{49}$ or in multiphase catalytic applications$^{50}$. Control over the phase diagram will allow new electrolytes with enhanced properties to be engineered, improving the efficiency of extraction, purification and capture reactions but also opening new avenue for designing electrochemical devices, e.g. systems that phase separate upon temperature increase such that thermal runaway processes can be shut-off$^{51,52}$. One important question nevertheless remaining relates to how the nature of the cations influences temperature dependence of the phase diagram for salt-salt ABSs. Generally, probing liquid-liquid interfaces in systems—be they electrochemical, biological or otherwise—has remained challenging. Our results demonstrate the power of Raman imaging, particularly over other popular optical imaging methods such as reflection/scattering.
microscopy, for providing label-free microscopic chemical information especially in material science applications. Combining such techniques with super-resolution\textsuperscript{53,54} or tip-enhanced\textsuperscript{55–57} approaches will allow access to the sub-100 nm resolution limit and broaden the set of systems that can be examined, \textit{e.g.} polymer ABSs and phase separations at electrified interfaces.

Combining more advanced Raman schemes (triple grating spectrometers\textsuperscript{58,59}, heterodyne optical Kerr effect\textsuperscript{60,61}) with other techniques such as neutron reflectometry (especially deep inelastic neutron scattering\textsuperscript{62–66} and sum frequency generation\textsuperscript{67–70} will be useful to gain an improved understanding of critical low-energy interactions such as hydrogen-bonds. The combination of these characterization techniques with novel chemistries will help unravel the complex ion transport properties across the ABSs interface, leading to new discoveries in the design of complex liquid electrolytes for membraneless electrochemical devices or phase separation.

Finally, comparing transport properties across liquid-liquid interfaces with that across more traditional solid-liquid interfaces\textsuperscript{71–74} ubiquitous in electrochemical devices will shine a new light on how (de)solvation can be modulated to fine tune ion transfer dynamics.
Materials and Methods

Materials. Lithium bis(trifluoromethanesulfonyl)imide (LiTFSI Extra dry, Solvionic, 99.9%), Lithium chloride (LiCl anhydrous, Alfa Aesar, 99%), Hydrochloric acid (HCl 30%, Sigma-Aldrich Suprapur) were weighted and MilliQ water was added to reach the desired molalities.

Phase diagram construction. Phase diagrams were measured using the cloud point titration method previously described\textsuperscript{16,17}. Briefly, starting from a known mass of a concentrated solution of one component, an aqueous solution of the second component is weighted while being added dropwise and vortexed until the solution turns cloudy. Then water is weighted while being added dropwise and vortexed until the solution turns clear again. These steps are repeated for each point of the diagram. The diagrams were made starting from different initial LiCl, LiTFSI and HCl concentrations especially for the HCl-LiTFSI-water system in order to prevent imprecision due to HCl evaporation.

Interfacial tension. Interfacial tensions were measured from the drop-weight method using Tate’s law with Harkins and Brown correction\textsuperscript{38}. Briefly, using a peristaltic pump (Watson-Marlow, 120U-DM2), drops of the highest density phase are slowly let fall into the lowest density phase. The average weighted of a drop is deduced and surface tension are calculated according to Harkins and Brown correction. Densities were measured using an electronic density meter (Anton Paar, DMA 35 Basic).

Infra-Red Spectra. The IR spectra for LiCl-LiTFSI-water system were taken using a Nicolet iS5 FTIR spectrometer, mounted with a diamond Attenuated Total Reflectance accessory (iD1 ATR). The
background correction was performed by measuring the ambient atmosphere and each spectra was acquired from 16 scans with a 4 cm\(^{-1}\) resolution from 4 000 to 500 cm\(^{-1}\).

**Variable temperature nuclear magnetic resonance.** Pulse-field gradient stimulated echo (PFG-STE) diffusion measurements were realized on a Bruker 200 MHz wide bore spectrometer, equipped with a Diff50 gradient system and a \(^{19}\)F-\(^{7}\)Li 8 mm insert inside a liquid-state Bruker probe. 1D profiles were recorded using the improf Bruker sequence (90°-gradient-170°-gradient-acquisition), with maximum field gradient strengths of 27, 29 and 30 G/cm respectively. The signals were acquired as full echoes (1-1.5 ms echo times) to avoid antiphase components and make the phasing easier. The maximum theoretical resolution was around 10-20 µm. Recycling delays of 7, 9 and 17 seconds were used, with 96, 16 and 16 transients acquired for \(^{1}\)H, \(^{19}\)F and \(^{7}\)Li respectively.

**Raman imaging.** The microscope was a standard layout of an epi-detected Raman microscope. A pump laser beam (wavelength = 532 nm, Newport Millenia eV) was spectrally cleaned up by a bandpass filter (FLH05532-4, Thor Labs), and its beam width was expanded to 7.2 mm before entering a home built inverted microscope. Additional waveplates (half-waveplate and quarter-waveplate for 532 nm, Foctek Photonics) precompensated the ellipticity introduced by the dichroic filter (F38-532_T1, AHF) and also generated circularly polarized light to alleviate effects due to molecular orientation. We used high numerical aperture (NA) oil-immersion objectives (Nikon 60X/1.4NA oil) to ensure high-resolution imaging and increase collection efficiency. Pump power before the objective was 150 mW, a power level that ensured no degradation of domains within the scanned region. The samples were scanned/descanned with a set of galvanometric mirrors (Thor Labs). The Raman inelastic backscattered light was collected by the same objective.
and focused with the microscope tube lens on the entrance slit of the detector which acted as
the effective pinhole for confocal detection. A notch filter blocked residual pump light (NF533-
17, Thor Labs) before guiding the signal to the spectrometer (Andor, Shamrock 303i, grating 300
l/mm), equipped with a high-sensitivity charge-coupled camera (Andor, iXon 897). All images
presented were taken with integration times/pixel in the 0.1 to 0.3 s settings range. Background
subtraction of Raman spectra was performed using a modified iterative polynomial smoothing
method. Peak decomposition was performed using a custom Python code and the lmfit toolbox.

**Titration Microcalorimetry.** The calorimetric titration measurements were completed
isothermally using a power compensation microcalorimeter (Thermal Hazard Technology, µRC).
Each injection was performed using an automated syringe pump tower attachment. The
reference samples were 1 mL of the respective ABSs solutions, and samples were stirred to allow
for uniform mixing. The partial enthalpy of mixing was found by integrating the resulting power
spike upon injection of one phase into the second phase (Supplementary Discussion 1).
Supporting Information

See the supporting information.

Acknowledgments

The authors acknowledge the French National Research Agency for its support through the Labex STORE-EX project (ANR-10LABX-76-01) and through the projects BALWISE (ANR-19-CE05-0014). R.P thanks Clare College, University of Cambridge for funding via a Junior Research Fellowship and Alpha A. Lee (Cambridge), Marcus Fletcher (Imperial College London) and Damien Laage (ENS) for useful discussions on the data.

Data availability

The data that support the findings of this study are available within the article and its supporting information. Additional data are available from the corresponding author upon reasonable request.

References


64. Kolesnikov, A. I. et al. Inelastic and deep inelastic neutron spectroscopy of water molecules under 

65. Schlossman, M. L. Liquid–liquid interfaces: studied by X-ray and neutron scattering. *Current 


for orientational flip-flop of water molecules at charged interfaces: A heterodyne-detected 

69. Nihonyanagi, S. *et al.* Unified Molecular View of the Air/Water Interface Based on Experimental and 
16880 (2011).

70. Yu, C.-C., Seki, T., Wang, Y., Bonn, M. & Nagata, Y. Polarization-Dependent Sum-Frequency 
Generation Spectroscopy for Ångstrom-Scale Depth Profiling of Molecules at Interfaces. *Phys. 

71. Cheng, Q. *et al.* Operando and three-dimensional visualization of anion depletion and lithium 

(2022).

73. Huang, L. *et al.* Negating Li+ transfer barrier at solid-liquid electrolyte interface in hybrid batteries. 