Liquid Structure with Nano-Heterogeneity Promotes Cationic Transport in Concentrated Electrolytes


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*Supporting Information

ABSTRACT: Using molecular dynamics simulations, small-angle neutron scattering, and a variety of spectroscopic techniques, we evaluated the ion solvation and transport behaviors in aqueous electrolytes containing bis(trifluoromethanesulfonyl)imide. We discovered that, at high salt concentrations (from 10 to 21 mol/kg), a disproportion of cation solvation occurs, leading to a liquid structure of heterogeneous domains with a characteristic length scale of 1 to 2 nm. This unusual nano-heterogeneity effectively decouples cations from the Coulombic traps of anions and provides a 3D percolating lithium−water network, via which 40% of the lithium cations are liberated for fast ion transport even in concentration ranges traditionally considered too viscous. Due to such percolation networks, superconcentrated aqueous electrolytes are characterized by a high lithium-transference number (0.73), which is key to supporting an assortment of battery chemistries at high rate. The in-depth understanding of this transport mechanism establishes guiding principles to the tailored design of future superconcentrated electrolyte systems.

KEYWORDS: aqueous electrolytes, batteries, molecular dynamics simulations, spectroscopy, conductivity

The electrolyte serves as the indispensable ion conductor and electron-insulator between the cathode and anode in all electrochemical devices. It forces the charge transfer to occur at electrode/electrolyte interfaces instead of directly between the cathode (oxidant) and anode (reductant), and its ability to conduct ions determines how fast the cell chemistry can proceed.1 Traditionally, the design and formulation of electrolytes have centered around the salt concentrations where the maximum ion conductivity ($\sigma_{m}$) occurs. This optimum concentration ($C_{m}$) is the result of the compromise between solution viscosity and population of ionic carriers.2 For nonaqueous electrolytes, $\sigma_{m}$ always occurs in the dilute ranges near 1.0 m (m = mol/kg), while the higher solvating power of H$_2$O, as characterized by both high dielectric constant ($\varepsilon = 78$ at 25 °C) and acceptor and donor numbers (AN = 54.8, DN = 18), shifts $\sigma_{m}$ toward higher concentrations. Thus, in practical electrochemical devices, the salt concentrations are almost always around 1.0 M (M = mol/L) for nonaqueous electrolytes and below 5 m in aqueous electrolytes.

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By this tradition, excessively concentrated electrolytes were unfavorable due to undesirable properties such as lower ion conductivity, narrower liquid range, and higher viscosity. These disadvantages make it difficult for the electrolyte to fully access the porosity of both the electrodes and the separators and incapable of supporting the cell reactions with sufficient mass flow and severely restrict the service temperature of the devices. It should be mentioned that, by this “conductivity-centric” principle, only overall ion conductivity was considered, while the importance of ion-transference number ($t^+$ or $t^-$) was often overlooked. For a given cell chemistry, however, only the ionic current provided by a specific ionic species (such as Li$^+$ for lithium batteries) is meaningful, while those contributed by the migration of other ionic species is considered parasitic and responsible for concentration polarizations within the cell.

In recent years this “conductivity-centric” custom was breached by a series of efforts in the field of batteries when researchers sought “unusual” properties that were otherwise unavailable at lower salt concentrations to benefit the electrochemical performances of various battery chemistries. This trend was represented by the pioneering polymer-in-salt concept of Angell et al., who attempted to decouple ion transport from mechanical strength of polymer materials, followed by Abe and Ogumi et al., who found that the intercalation chemistry of graphite could be quite unexpectedly supported by propylene carbonate (PC) solvent as long as salt was used at sufficient concentrations ($\sim 5$ M), and Watanabe et al., who discovered that volatile ether-based electrolytes gained thermal stability at high salt concentrations, while demonstrating solubility/insolubility toward sulfur active materials.

These approaches were further refined by Yamada et al., who used unconventional solvents to achieve unusual interfacial properties on a graphite anode and high-voltage cathode when lithium salts were used at superconcentrations. Solvent concentrations also greatly influenced Al current collector corrosion, which was carried to its extreme when Suo et al. dissolved up to $21$ m lithium salt (lithium bis(trifluoromethane sulfonyl)imide, LiTFSI) and successfully expanded the electrochemical stability window of aqueous electrolytes from $1.23$ V to nearly $3.0$ V, leading to a paradigm shift in aqueous electrolytes, which have been demonstrated with an assortment of battery chemistries with exceptional performances by numerous research groups.

The superconcentrated aqueous electrolyte reported therein, along with its improved variations reported later on, also presents an unexplored frontier, where little is known about how ions interact with solvent molecules and with each other or how they participate in transport. In this work, combining computational and complementary characterization techniques, we explore this frontier and describe how a discovered nanoscale heterogeneity influences a range of structural and dynamic properties in this class of aqueous electrolytes.

**RESULTS AND DISCUSSION**

**Ion Solvation.** It has been well established that cations are well solvated while anions remain relatively free in diluted nonaqueous electrolytes. This is partially caused by the fact that the formal charges are usually more delocalized on the anion rather than on the cation, but more importantly, the typical nonaqueous solvent molecules, regardless of chem-

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**Figure 1. Solvation:** (a) Fraction of the solvent-separated and free TFSI$^-$ anions as a function of salt concentration as derived from the $\mathrm{S=O}$ band stretch vibrational mode in FTIR and MD simulations using two different criteria for SSIP formation in MD simulations: (1) SSIP is formed if there is no Li$^+$ within a range of $5$ Å from N(TFSI), (2) SSIP is formed if there is no Li within $2.7$ Å from any of O(TFSI); a fraction of the solvent-separated Li$^+$ is also shown; (b) Li$^+$ cation coordination numbers from MD simulations using a $2.7$ Å cutoff for the first shell; and (c) probability of finding a specific number of oxygens from TFSI, $\mathrm{H_2O}$, or nitrogen atoms from acetonitrile (AN) in the Li$^+$ first coordination shell for LiTFSI in $\mathrm{H_2O}$ and LiTFSI in AN from MD simulations at solvent Li = 2.67. (d) Isosurfaces of water oxygen (red solid, $\rho/\rho_{\text{bulk}} = 5$, where $\rho$ is local density and $\rho_{\text{bulk}}$ is bulk average density), oxygen of TFSI (green mesh, $\rho/\rho_{\text{bulk}} = 2.5$), fluorine of TFSI (cyan mesh, $\rho/\rho_{\text{bulk}} = 12$), and Li$^+$ (purple solid, $\rho/\rho_{\text{bulk}} = 12$) for $21$ m LiTFSI in $\mathrm{H_2O}$ at $298$ K. Average atom positions of the flexible TFSI are shown, resulting in a somewhat collapsed structure of the $\mathrm{-SO_2-CF_3}$ group. TFSI colors: N (blue), O (red), F (cyan), S and C (brown).
istry—ethers, esters, sulfones, or nitriles—are better electron donors than acceptors, as structurally their negative termini are more readily accessible than positive termini (Figure S1a). However, this “cation preference” would be significantly weakened in aqueous solutions, as H2O molecules are bipolar in nature and solvate both cations and anions effectively (Figure S1b). According to quantum chemistry (QC) calculations, the total binding energy for Li’(H2O)4 −103 kcal/mol,15 is smaller than for bipolar solvents: Li’(AN)4 with a binding energy of −117.5 kcal/mol28 and −122 kcal/mol for Li’ coordinated by cyclic carbonates, as shown in Figure S2.29 Yet, salt dissociation is higher in H2O due to its bipolar character and the more effective solvation of anions. On the other hand, the interaction between cations and anions depends critically on the salt concentration, as the number of solvent molecules available to solvate cations or anions determines whether a counterion enters into the primary solvation sheath of each other, forming the so-called contact ion pair (CIP). As the concentration of LiTFSI increases from 1.0 m to 21 m, the H2O-to-salt molar ratio decreases from 5.6 to 2.6, resulting in the coexistence of various ionic species from solvent-separated ion pairs (SSIPs) to CIPs and ionic aggregates (AGGs). Fourier-transformed infrared (FTIR) measurements conducted on the LiTFSI−H2O system track such changes in the stretching vibrational mode of the S=O band (Figure S3). Applying principal component analysis, we were able to deconvolute the three major components (free TFSI anion, SSIPs, and AGGs consisting of Li’ and TFSI−), whose molar distribution is shown as a function of LiTFSI concentration (Figure 1a). A similar distribution of ionic species was also obtained from the symmetric stretching of the S−N bond, which includes in-plane (1132−1139 cm−1) and the out-of-plane (1143−1149 cm−1), or the antisymmetric stretching of the S−O bond, which includes in-plane (1352−1355 cm−1) and out-plane (1330−1336 cm−1), each of which were measured and analyzed (Figures S3 and S4). Based on this analysis, minimal CIP formation is observed below 5 m, while highly concentrated solutions yield a high fraction of CIPs and AGGs. This intensified association between Li’ and TFSI− constitutes the foundation for the formation of a LiF-rich interphase on the anode above 1.5 V vs Li/Li’ as a result of the salt reduction and is essential for expanding the electrochemical stability window.15,30 MD simulations also predicted little CIP formation at 5 m and a similar decrease in the fraction of SSIPs with increasing salt concentrations. Two definitions of the Li’−TFSI bond formation are utilized here: the first one uses the distance of the minima after the first Li’−N(TFSI) peak of 5.0 Å, while the second uses the position of the first Li’−O(TFSI) peak and counts the Li’−O(TFSI) ion pair being formed if any of the TFSI−oxygen is bound to a Li’ (see Figures S5 for the radial distribution functions). The Li’−O(TFSI)-based definition does not include the Li’−TFSI CIPs, where the Li’ cation is bound to the CF3 groups (see Figure S6a,b); thus it overestimates the solvent-separated TFSI anions. The Li’−N(TFSI) definition, on the other hand, counts all the Li’−TFSI bonds including most of the Li’−F(TFSI) contacts but in some cases incorrectly counts the compact SSIPs as CIPs, as shown in Figure S6c. Thus, the latter definitions provide a lower bound estimate of the fraction of the solvent-separated TFSI anions shown in Figure 1a. The experimentally determined fraction of the solvent-separated TFSI− shown in Figure 1a is in excellent agreement with the MD simulation predictions, as the two definitions used for the solvent-separated TFSI anions envelop the experimentally derived curve. Unlike FTIR, which can only detect the TFSI− anion, MD also probes the association behaviors of the Li’ cations, as shown in Figure 1a. An important insight generated by MD that will be of high significance to ion transport is that, according to the MD predictions, the fraction of SSIP Li’ is significantly higher than the fraction of SSIP TFSI−. This result suggests that a surprisingly high fraction of Li’ remains “free,” perhaps in the form of Li’(H2O)4 and that these clusters are complemented by negative anion-rich domains such as Li’(TFSI−)x with x > 1.

The Li’−solvation sheath details are analyzed from MD simulations. The Li’ primary sheath has a radius of 2.7 Å, which corresponds to the location of the minimum after the first peak in the Li’−O(H2O) radial distribution function (see Figure S5). We found that in solutions with lower concentration (<5 m) Li’ is mainly solvated by four H2O molecules in its first coordination shell (Figure 1b), which is consistent with the neutron scattering results for other “dilute” salts.31,32 Contact ion pairs start to form within the 5−10 m concentration range, and the trend continues with the increase in salt concentrations, resulting in a very aggregated solution at 21 m with each Li’ coordinated by approximately two oxygens from TFSI and two oxygens from H2O in its first solvation sheath. However, the average coordination numbers do not give the full picture of the distribution of H2O and TFSI around Li’. A closer look is revealed by the analysis of the distributions of various environments shown in Figure 1c. Even though each Li’ is coordinated by 1.75 TFSI oxygens on average, a very high fraction (40%) of the Li’ actually exist as SSIPs with no TFSI in their first coordination shell at all. Specifically, the Li’ ions prefer either (1) to be fully solvated by four H2O molecules or (2) to participate in the formation of the LiTFSI aggregates so that each Li’ is surrounded by four oxygens of TFSI, rather than an even division of H2O molecules and TFSI around each Li’.

Such a solvation disproportionation is uncommon in nonaqueous electrolytes such as the LiTFSI−acetonitrile (AN) system, which was selected as a representative nonaqueous electrolyte due to multiple similarities between AN and H2O that include large dipole moments (3.92 D for AN vs 1.8546 D for H2O) and high dielectric constant (36.8 for AN vs 78 for H2O). Despite similar average coordination numbers (1.78 AN and 2.26 O(TFSI) per Li’) with 21 m LiTFSI−H2O, the primary Li’−solvation sheath in LiTFSI−AN consists of two solvents and two O(TFSI) as the most probable configuration, with only a small fraction (10%) of SSIP Li’. This even distribution of AN molecules and TFSI leads to a more homogeneous Li’ environment, where the cation and anion are expected to be intimately coupled. In sharp contrast, the “disproportionation” in solvation behaviors by H2O and TFSI would certainly create a rather inhomogeneous local Li’ environment, and the difference in solvation sheath structures is expected to result in dramatically different ion transport as well as interphasial behaviors.

The structure of the TFSI− shell is examined in Figure 1d by plotting the isosurfaces of 3D distributions of species around the TFSI− anion. The Li’ cation primarily binds to TFSI− oxygens and is coordinated by oxygens from the other TFSI or water, as also shown in Figure S12. Water oxygens often coordinate the TFSI− anion on the side opposite the Li’ cation in addition to coordinating the Li’ cations. The Li’ cation binding to F(TFSI) atoms is less probable than binding to
of H₂O and the excellent solubility of LiTFSI in it, enabled by nucleophilic. The combination of this excellent solvation power with ethers, sulfones, or nitriles, which are polar but are usually only compared with the nonaqueous solvent molecules of esters, which even the nonaqueous electrolytes fail to rival.

approaches the −e typically results in a much higher salts therein. The bipolar nature of H₂O enables it to be very aqueous and nonaqueous electrolytes rests with the solubility of species, with or without charge. An apparent di aggregation results in a decrease of available free ions that can participate in conducting current, while increasing Coulombic frictions among cation, anion, and solvent molecules simultaneously creates additional viscous drag to any moving species, with or without charge. An apparent difference between aqueous and nonaqueous electrolytes rests with the solubility of salts therein. The bipolar nature of H₂O enables it to be very effective at coordinating with both cations and anions, which typically results in a much higher as compared with the nonaqueous solvent molecules of esters, ethers, sulfones, or nitriles, which are polar but are usually only nucleophilic. The combination of this excellent solvation power of H₂O and the excellent solubility of LiTFSI in it, enabled by its well-delocalized anion structure, makes it possible to create an extremely concentrated aqueous electrolyte, where the salt concentration can reach 21 m. In this superconcentrated LiTFSI–H₂O system, despite its high viscosity (32 mPa s), the ionic conductivity (∼8 mS/cm at 25 °C) is comparable to what typical nonaqueous electrolytes can offer (e.g., ∼10 mS/cm at 25 °C for LiPF₆ in EC/EMC, 30:70). For comparison, at 60 °C, the LiTFSI–H₂O conductivity is also about a factor of 2 higher than conductivity for LiTFSI–AN, one of the most conductive nonaqueous systems. Numerous aqueous cell chemistries have been found to be effectively supported by this electrolyte or its variations at high rates, some of which even the nonaqueous electrolytes fail to rival.

In a simple comparison between the reciprocal temperature dependences of ion conductivities (Figure S7), one would immediately conclude that the LiTFSI–H₂O system is characterized by strong Arrhenius behavior even at the extreme salt concentration of 21 m, while LiTFSI–AN displayed typically a non-Arrhenius feature that is well described by the empirical Vogel–Fulcher–Tamman (VFT) relation, as most nonaqueous electrolytes investigated thus far. The deviations from the Arrhenius behavior only occurred in the LiTFSI–H₂O system around 0 and 20 °C, when partial crystallization of either H₂O or LiTFSI leads to two-phase coexistence. Surprisingly, significant conductivity is still maintained in the two-phase region at lower temperatures, making this electrolyte rather durable for supporting battery operations even below its melting point.

Detailed analyses were carried out to identify the activation energies associated with ion transport in each electrolyte. Due to the nonlinear nature of the nonaqueous system, the temperature dependence of conductivity could be divided into three temperature regions, i.e., from −20 to ∼0 °C, from 0 to ∼30 °C, and from 30 to ∼60 °C (Figure S7). Each section displays strongly concentration-dependent activation energies, as shown in Figure 2a, with the concentrated electrolytes associated with the highest activation energy (30–50 kJ/mol), an indication of how difficult it is to move cations and anions free from their individual Coulombic cages. Thus, we speculate that in nonaqueous media at high salt concentrations, Li⁺ movement is expected to be intimately coupled with TFSI⁻. Similar increase of the activation energy with concentration has been observed in 1,3-dioxolane (DOL), dimethoxyethane (DME) doped with LiTFSI, glymes-LiTFSI, AN doped with LiTFSI, LiPF₆, LiClO₄, LiBF₄, and LiCF₃CO₂, and mixtures of carbonates with LiPF₆ or LiBF₄ salt. In sharp

Figure 2. Ion transport: (a) Comparison of activation energies of LiTFSI–AN and LiTFSI–H₂O as a function of salt concentrations from experiments and for LiTFSI–H₂O from MD simulations; (b) ion and H₂O self-diffusion coefficients for the LiTFSI–H₂O electrolyte from MD simulations at 25 °C and pfg-NMR experiments at 20 °C; (c) the Li⁺ transference number as measured by pfg-NMR for the electrolytes with one primary solvation group: for LiTFSI in H₂O at 20 °C, LiPF₆ in PC, EC/EMC (1:1 wt %), and LiTFSI–AN at 25 °C; data from Takeuchi et al. for PC-LiPF₆ at 25 °C are also shown; (d) conductivity (σ) and the Li⁺ contribution to conductivities (σli⁺) for LiTFSI–H₂O, LiTFSI–AN, and LiPF₆–EC:EMC (1:1 wt/wt).
contrast, over the entire concentration range, the aqueous electrolytes demonstrated a concentration independence of the activation energies, which remain at a constant low value of \( \sim 20 \text{ kJ/mol} \) even at superconcentration. As predicted in Figure 1c, a very substantial fraction of the solvent-separated \( \text{Li}^+\text{(H}_2\text{O)}_4 \) is expected to exist, which facilitates the \( \text{Li}^+ \) diffusion through a framework consisting of TFSI and yields a noticeably different transport mechanism in LiTFSI–H\(_2\)O. It is this “solvation disproportionation” in ion aggregation that is responsible for the fast ion conduction even at very high salt concentration. Interestingly, analysis of ion transport in LiTFSI–H\(_2\)O using the so-called Walden rule, as shown in Figure S8 as the dependence of molar ion conductivities on viscosity and decreasing viscosity, indicated that increasing viscosity and decreasing fluidity brought data slightly closer to the ideal line, indicating decoupling of ionic motion as salt concentration increases.

The diffusivities of \( ^7\text{Li} \), \( ^{19}\text{F} \), and \( ^{19}\text{F} \) were measured separately by p\text{g}-NMR as a function of salt concentration, which correspond to the movement of solvent molecules (H\(_2\)O), cations (Li\(^+\)), and anions (TFSI\(^-\)), respectively. Three representative nonaqueous electrolytes, LiTFSI–AN, LiPF\(_6\)–PC, and LiPF\(_6\)–ethylene carbonate/ethylmethyl carbonate (EC/EMC) (1:1), were selected for comparison. Figure 2b compares these diffusivities in LiTFSI–H\(_2\)O at 20 °C, along with a data set provided by MD simulations at 25 °C, and the apparent transference number of \( \text{Li}^+ \) (\( \text{t}^+ \)) is evaluated according to

\[
\text{t}^+ = \frac{D_{\text{Li}^+}}{D_{\text{Li}^+} + D_{\text{TFSI}^-}}
\]

(1)

where \( D_{\text{Li}^+} \) and \( D_{\text{TFSI}^-} \) represent cation and anion diffusivities, respectively. Figure 2c summarizes the concentration dependence of \( \text{t}^+ \) for LiTFSI–H\(_2\)O along with these nonaqueous counterparts. The superconcentrated LiTFSI–H\(_2\)O at 21 m stands out with a \( \text{t}^+ \) of 0.70 and 0.73 at 20 and 30 °C, respectively, as shown in Tables S2 and S3, far above what typical nonaqueous electrolytes could offer. On the other hand, a comparable \( \text{t}^+ \) of 0.6–0.7 has been also described in a few superconcentrated electrolytes based on ethers, including LiTFSI–triglyme or LiTFSI–1,3-dioxalane/dimethoxyethane\(^9\) and sulfolane/ethyl acetate/LiBF\(_4\) electrolytes.\(^{42}\) After converting overall ion conductivities (\( \sigma \)) into \( \text{Li}^+ \) conductivities (\( \sigma_{\text{Li}^+} \)), the superconcentrated LiTFSI–H\(_2\)O at 21 m is clearly a much better electrolyte to support Li-ion chemistries (Figure 2d), with a \( \sigma_{\text{Li}^+} \) of 7.3 mS/cm in comparison with 3.7 mS/cm for the typical nonaqueous electrolyte 1.0 m LiPF\(_6\)–EC/EMC (50:50), despite the similar overall conductivities for both (\( \sim 10 \) mS/cm).

It is well known that cation-transference numbers in typical nonaqueous electrolytes are seldom above 0.50\(^1\)–\(^2\),\(^44\)–\(^45\) and are often even lower for nonconcentrated aqueous electrolytes: within 0.2 to 0.45 for the LiCl–, LiClO\(_4–\), and LiCF\(_3\)SO\(_3–\)-based aqueous electrolytes.\(^{46}\) This under-representation of the cation in ion transport current implies the inability of electrolytes to maintain sufficient mass flow to support a cell chemistry involving the cation, especially under high current, where severe concentration polarization occurs. A typical example is the Li-ion battery, which suffers from a series of harmful parasitic reactions at high C-rates, such as Li metal deposition due to Li\(^+\) depletion at electrolyte/electrode surfaces and excessive electrolyte decompositions. In this sense, a preferred cation transport would significantly improve the electrochemical performances of batteries by minimizing the movement of the ions that do not participate in the intercalation or conversion reactions at electrodes.

Closer examination of measured diffusivities reveals that, across the entire concentration range, H\(_2\)O molecules are the most mobile species, followed by Li\(^+\), while TFSI\(^-\) are the least mobile (Figure 2b). At concentrations higher than 5 m, the ratio among the diffusivities of H\(_2\)O, Li\(^+\), and TFSI\(^-\) becomes approximately constant, indicating that a connection exists between the solvent and ion movements. A combination of p\text{g}-NMR and conductivity measurements allows us to determine the so-called ionicity using eqs 2 and 3, which defines \( \alpha_b \), which reflects the degree of uncorrelated ionic motion:

\[
\alpha_d = \frac{\sigma}{\sigma_{\text{uncorr}}}
\]

(2)

\[
\sigma_{\text{uncorr}} = \frac{\epsilon^2}{k_bT} (n_eD_e + n_D_D)
\]

(3)

where \( \epsilon \), \( n_e \), and \( n_D \), \( k_b \), and \( T \) are electron charge, number densities of Li\(^+\) and TFSI\(^-\) in the material, Boltzmann constant, and temperature, respectively. Interestingly, in accord with the Walden plot, ionicity was found to remain relatively steady in the region of 0.64 to 0.76 as salt concentration increases (see Figure S9), indicating that even in the very concentrated regime
where one should expect significant cation–anion interactions, the movements of \( \text{Li}^+ \) and \( \text{TFSI}^- \) are only weakly correlated. Similar values (0.6 to 0.7) were also observed in superconcentrated \( \text{LiTFSI}^- \)-triglyme.\(^ {41}\) Temperature dependence of \( \alpha_d \) as derived from MD simulations showed that it decreases slightly from 0.7 to 0.45 as the temperature increased from 298 K to 363 K for 21 m, implying that at this high salt concentration the ion dynamic correlation would increase with increasing temperature. However, at lower salt concentrations, \( \alpha_d \) would remain largely unchanged.

**Structure of Nano-heterogeneity.** The solvation disproportionation, the uneven distribution of \( \text{H}_2\text{O} \) and TFSI with each \( \text{Li}^+ \), the nearly constant activation energy, the preferred \( \text{Li}^+ \) transport, and the degree of uncorrelated cation and anion movements suggest that an unexpected liquid structure may have formed in the superconcentrated aqueous electrolytes. This structure, differing from more conventional diluted aqueous/nonaqueous analogues, immobilizes the anion in some manner and frees a significant portion of cations from the Coulombic traps of their anions. The promoted fast cation transport effectively supported various cell chemistries at high C-rates.\(^ {15,17,24}\) Although researchers started suspecting unusual liquid structures arising in the superconcentration realm of nonaqueous electrolytes,\(^ {47}\) the salt concentrations therein (often <5 m) were not high enough to make these behaviors pronounced enough to be unambiguous, and direct evidence was never presented.

Based on MD simulation results presented above, an overall picture of matter distribution can be generated on a molecular level. We have observed that in the 21 m \( \text{LiTFSI}^- \)-\( \text{H}_2\text{O} \) solution 40% of \( \text{Li}^+ \) does not have any oxygen of TFSI in its first coordination shell, while 25% of \( \text{Li}^+ \) does not have any \( \text{H}_2\text{O} \) in its primary solvation sheath (Figure 1c). The system does not phase-separate on a microscopic scale; however, \( \text{H}_2\text{O} \) is distributed nonhomogeneously and forms the interconnecting clusters shown in red in Figure 3a in a 3D rendering. The “anionic framework” as represented by wireframe regions appears to be structurally dissimilar to what exists in superionic glass/ceramics due to its long-range disorder and highly dynamic nature, as will be shown below. In a certain sense, these superconcentrated electrolytes should be viewed as an intermediate state transitioning from a diluted solution into an ionic liquid. Those 40% of \( \text{Li}^+ \), entirely coordinated by \( \text{H}_2\text{O} \) molecules, are expected to transport \( \text{Li}^+ \) through the 3D percolating network shown in Figure 3a and are likely responsible for the unusually high \( \text{Li}^+ \)-transference numbers. Importantly, the above nano-heterogeneity was not only observed in MD simulations but also confirmed experimentally with small-angle neutron scattering (SANS), in which \( \text{D}_2\text{O} \) was used in place of \( \text{H}_2\text{O} \) to enhance contrast due to a much larger coherent scattering length for deuterium than hydrogen. Figure 3b shows a structure factor from SANS that is precisely at the same position and has the same shape as predicted from MD simulations. A small low-Q peak in the structure factor around
0.45 to 0.50 Å⁻¹ indicates that a nanostructure is present on the scale of ~1.4 nm.

MD simulations allow us to further prove the above link between the low-Q peak and the proposed structure by artificially decreasing the fraction of Li⁺(H₂O)₄ and examining how it influences the size of the nanodomains. MD simulations utilizing such a modified force field resulted in reducing the fraction of Li⁺ SSIP at 21 m from 40% for the original force field to 30%. This very small decrease in the Li⁺ SSIP population renders the low-Q peak much less defined as shown in Figure 3b. Thus, a small change of the Li⁺→H₂O interactions and the fraction of solvent-separated Li⁺ in MD simulations induces a significant change in the intermediate range structure, and the existence of a high fraction of the fully solvated Li⁺(H₂O)₄ is important for creating a nanodomain structure in this electrolyte. Moreover, the sensitivity of the low-Q peak toward the fraction of the solvent-separated Li⁺ cation provides additional evidence that the fraction of Li⁺ SSIP observed in MD simulations is accurate and highlights the high quality of the many-body polarizable APPLE&P force field used in simulations that also accurately predicted ion self-diffusion coefficients, conductivity, and viscosity.

The evolution of the nano-heterogeneity length scale with salt concentration was also analyzed as shown in Figure 4a. The peak increases in size with decreasing salt concentration up to 9.3 m and seems to completely disappear at 5 m. In accord with changes of the low-Q peak, the nano-heterogeneity with a length scale of 1 to 2 nm seems to be more pronounced in the range of 9 to 14 m, while increasing the salt concentration to 21 m makes it less distinct. This trend becomes more visible in the snapshots of matter distribution (Figure 4b), where H₂O molecules and LiTFSI are shown in red and blue, respectively. Additional snapshots can also be seen in Figure S10. Analysis of the partial contributions to the structure factor shows that D₂O→D₂O correlations contribute the most to the low-Q peak, as shown in Figure S13. Figure S14 shows that the low-Q peak is largely absent in the deuterated AN–LiTFSI solutions because its solvation is dominated by Li⁺(AN)ₙ(TFSI)ₖ₋ₙ (n = 2, 3, 4) solvates that combine both AN and TFSI and a very small fraction of the Li⁺(AN)ₙ SSIP. Small and hydrophilic anions such as Br⁻ yielded structure factors without the low-Q peak. Poly(ethylene oxide) (PEO)–LiTFSI, on the other hand, formed the large-size Li⁺(ethylene oxide)ₙ (n = 5, 6) solvates that are often solvent separated from TFSI⁻, resulting in the large intermediate range peak at 0.6 Å⁻¹, as shown in Figure S14, and was previously observed in neutron diffraction experiments. In the TFSI-based room-temperature ionic liquids with small cations the low-Q peak was observed at larger Q values, >0.8 Å⁻¹, and was attributed to cation–cation and anion–anion correlations.

The inhomogeneous structure of the superconcentrated LiTFSI–H₂O system could be viewed as consisting of two independent but interpenetrating networks, one bearing excessive positive charge (i.e., Li⁺(H₂O)₄) and one bearing excessive negative charge (i.e., Li⁺(TFSI)⁻, domain). There is some loose analogy between this structure and the structure of the "soggy sand" electrolytes consisting of the solid–liquid electrolyte with anions adsorbed on the solid resulting in the increase of tᵣ. One critical question remaining is, are the domains equally dynamic and do Li⁺ ions move together with H₂O or by solvent exchange? To seek an answer, the mean-squared displacements (MSDs) were calculated for Li⁺(H₂O)₄ not coordinated by TFSI and the Li⁺ coordinated only by TFSI anions, respectively (Figure 4b). We found that the Li⁺ cations that have no TFSI around diffuse 3 times faster compared to the Li⁺ cations bound by two TFSI on the time scale of a few nanoseconds, which is the lifetime of these solvates. Much faster diffusion of Li⁺(H₂O)₄ suggests that these solvates are the dominant charge carriers in the solution. Furthermore, the vehicular motion of the Li⁺(H₂O)₄ is identified as the dominant mechanism, because the Li⁺ moves multiple H₂O sizes (from 5.1 to 6.4 Å depending on salt concentration) before it changes its hydration shell on the nanosecond time scale at 298 K. Thus, the H₂O-rich domain in the nano-heterogeneity is responsible for the fast ion transport as well as the high cation–transference number, with a significant contribution of the Li⁺ vehicular mechanism instead of the solvent exchange mechanism, contrary to the speculation of Yim et al. In contrast, in (AN)₂ₓLiTFSI electrolyte the solvent-exchange contribution becomes important at high concentrations as the Li⁺ exchanges each solvent on average as it moves a size of the solvent molecule, which is consistent with the previous discussions. This unusual and interesting transport mechanism in LiTFSI–H₂O correlates quite well with the observation of nearly constant activation energies for ion conductivity (Figure 2a), which in turn explains why in typical nonaqueous electrolytes there is such a dramatic change in activation energies with salt concentration due to participation of anion and solvent exchange in the latter. Undoubtedly, the structure and transport mechanisms identified in this work shed valuable light on the tailored design of future electrolyte systems, both aqueous and nonaqueous and either dilute or superconcentrated.

CONCLUSION

Combining electrochemical, scattering, spectral, and computational techniques, we thoroughly examined the ion solvation and transport in the LiTFSI–H₂O system in a wide concentration range from dilute to superconcentrated (21 m) and found that the presence of solvent-separated Li⁺ even at high salt concentrations induces solvation disproportionation of Li⁺ and TFSI⁻ by H₂O molecules, leading to the formation of a liquid structure with nano-heterogeneity. The inhomogeneous distribution of Li⁺ creates two interpenetrating but dynamic nanodomain networks. The TFSI-rich domain relatively immobilizes the anion movement, while the Li⁺(H₂O)₄ domain serves as a 3D percolating channel for fast Li⁺ transports with high transference number, in sharp contrast to the low transference numbers reported for dilute aqueous electrolytes. It is this ion transport mechanism achieved in superconcentrated aqueous electrolytes that supports diversified cell chemistries at high C-rates. The understanding of this liquid structure of nano-heterogeneity will undoubtedly benefit the future efforts of seeking improved electrochemical storage devices.

MATERIALS, METHODS, AND MODELING DETAILS

Lithium bis(trifluoromethane sulfonyl) imide (LiN(SO₂CF₃)₂, LiTFSI) (>99%, TCI), lithium hexafluorophosphate (>99.9%, BASF), acetonitrile (HPLC grade, Sigma-Aldrich), and carbonate solvents including ethylene carbonate (EC), ethylmethyl carbonate (EMC), and propylene carbonate (PC) (battery grade, BASF) were all used as received. Aqueous and nonaqueous electrolytes were prepared by mixing either LiTFSI or LiPF₆ with solvents at specified molalities, with the former being made in open ambient and the latter in an Ar-filled glovebox. They were denoted hereafter as LiTFSI–H₂O,
MD simulations were performed on the LiTFSI–H₂O system at 298, 333, and 363 K at four concentrations (5.05, 9.25, 13.88, and 20.8 m), which correspond to H₂O/LiTFSI ratios of 11, 6, 4, and 2.7, respectively. Rounded-up molalities of 5, 9.3, 14, and 21 m are used to refer to these systems in the discussion. Three replicas with different starting configurations were simulated for 21 m LiTFSI in order to establish convergence of MD simulation results with the slowest relaxation. MD simulations utilized a previously developed Apple~LiP6 many-body polarizable force field that was revised for Li”TFSI” and Li’H₂O along with one additional modification: the Li+ and TFSI− oxygen charges were reduced to make their overall charges 0.92 to enhance ion dynamics.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.nano.7b05664.

Description of IR experiments, MD simulations, conductivity data from experiments, and MD simulations, Walden plot, “ionicity” or dynamic degree of ion dissociation, simulation snapshots, neutron-weighted structure factors from MD simulations and SANS experiments, and the solvate binding energies from quantum chemistry (PDF)

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Notes

The authors declare no competing financial interest.

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