Modulation of Cation Diffusion by Reversible Supramolecular Assemblies in Ionic Liquid-Based Nanocomposites

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ABSTRACT: Ionic liquid (IL) properties, such as high ionic conductivity under ambient conditions combined with nontoxicity and nonflammability, make them important materials for future technologies. Despite high ion conductivity desired for battery applications, cation transport numbers in ILs are not sufficient enough to attain high power density batteries. Thus, developing novel approaches directed toward improvement of cation transport properties is required for the application of ILs in energy-storing devices. In this effort, we used various experimental techniques to demonstrate that the strategy of mixing ILs with ultrasmall (1.8 nm) nanoparticles (NPs) resulted in melt-processable composites with improved transport numbers for cations at room temperature. This significant enhancement in the transport number was attributed to the specific chemistry of NPs exhibiting a weaker cation and stronger anion coordination at ambient temperature. At high temperature, significantly weakened NP−anion associations promoted a liquid-like behavior of composites, highlighting the melt-processability of these composites. These results show that designing a reversible dynamic noncovalent NP−anion association controlled by the temperature may constitute an effective strategy to control ion diffusion. Our studies provide fundamental insights into mechanisms driving the charge transport and offer practical guidance for the design of melt-processable composites with an improved cation transport number under ambient conditions.

KEYWORDS: nanocomposites, dynamic bonds, ionic conductivity, cation transport number, energy storage

INTRODUCTION

Ionic liquids (ILs) are materials of great importance for various energy applications because of their high ionic conductivity. However, a significant shortcoming of ILs is their relatively low cation transport number (\( t^+ \)) (typically \( \sim 0.4 \)), which impedes their practical application. For instance, a small contribution of the cation to conductivity creates a significant gradient in ion concentration that leads to increased cell polarization, thereby reducing the power capability of a battery. To advance the application of ILs in energy-harvesting and storage devices, the development of novel approaches leading to high cation transport numbers is desired.

The strategies toward increasing \( t^+ \) are related to anion immobilization through covalent or noncovalent interactions with polymers and/or various ceramics. Covalently attaching anions to polymers results in the transport number of cations close to unity, yielding single-ion-conductive materials. However, compared to ILs, the significant drop in the conductivity observed for these polymers in the dry state constitutes a major limitation for their application. Although more efforts are underway with solid polymer electrolytes, alternative commercially viable strategies include the development of single-ion-conducting gel polymer electrolytes, which show both high conductivity and high ion transport number. However, only a few examples of such materials are available in the literature.

The effect of covalent and noncovalent interactions between the cation and ceramics has been a topic of extensive research. One of the successful examples of covalently immobilized anion that have relatively high conductivity (\( 4 \times 10^{-4} \) S/cm at 343 K) and high cation transport number in the dry state has been reported for cografting bis-(trifluoromethanesulfonyl)imide (TFSI) anions and strands of polyethylene oxide (PEO) to aluminum and silica
nanoparticles (NPs). Furthermore, not only covalent attachment but also the physical blending of ionic materials with ceramics has been shown to significantly improve the cation transport number. For example, Croce et al. reported that depending on the acidity of the ceramics, the value of transport number of Li for a PEO/LiClO4-based polymer composite can be improved from 0.25 to 0.5–0.6 for acidic TiO2 particles while for less acidic Al2O3 particles, a modest increase from 0.25 to 0.3 is observed. Although the improvement of the cation transport number for polymer electrolytes in the presence of ceramics is supported by the literature, much less is known about the same effect in ILs. Nevertheless, the findings in the field of polymer electrolyte-based composites identify that the interfaces between the inert ceramic and polymer electrolytes might play a significant role in the improvement of the transport properties of the cation. In this respect, the utilization of smaller particles (below 100 nm) which have a large surface area might benefit the transport properties of the cation. However, the addition of NPs has raised concerns about material processability. For instance, in IL-based composites with nonconductive NPs, the high ionic conductivity is achieved only with NP concentration not exceeding 10 wt % while further increasing the concentration of NPs ultimately leads to a degradation in the mechanical and conductive properties. Given the performance trade-offs between ion conductivity, processability, and transport number in various polymer and composite materials, it is important to design a hybrid structure that will be easily processable and combine the high transport number of single-ion-conducting polymers with the high conductivity of ILs.

In this paper, we pursue the strategy of adding ultrasmall octaaminophenyl silsesquioxane (OAPS) NPs (D ≈ 1.8 nm) to the diocetyl diethy lammonium bis-(trifluoromethanesulfonyl)imide (N2288 TFSI) IL to design a melt-processable composite with an improved transport number of the cation. The selection of these NPs is driven by their large surface area and unique surface chemistry capable to form noncovalent interaction with the ions in ILs. Additionally, their size was selected based on the literature reports where the decoupling between charge transport and structural relaxation has been found in low-molecular-weight (MW) aprotic ILs. Understanding and nourishing this mechanism in IL composites might be effective in designing the new generation of batteries.

**EXPERIMENTAL DETAILS**

**Materials and Methods.** OAPS NPs were purchased from Mayatex. The N2288 TFSI IL was purchased from IoLiTec. All materials were used as received, that is, without further purification.

The OAPS particle and the IL were individually dissolved in tetrahydrofuran (THF). Based on the weight ratio, the OAPS/THF solution was added dropwise to the IL/THF solution while stirring. The mixture was stirred for more than an hour in a closed container, followed by the slow solvent evaporation after the lid was removed. The samples were dried in an oven at 110 °C for 24 h to remove the residual solvent and water before the measurements. The volume fractions of OAPS in nanocomposites were obtained by thermogravimetric analysis (TGA, performed with a Q50 device from TA Instruments using a cooling rate of 20 K/min, under air) from the weight loss between 403 and 1073 K.

**DSC Measurements.** The calorimetric tests of the studied samples were performed using a Mettler Toledo DSC1STAR System equipped with a liquid nitrogen cooling accessory and an HSS8 ceramic sensor (a heat flux sensor with 120 thermocouples). The sample was measured in aluminum pans with a 40 μL volume. Before the measurements, the samples were annealed for 15 min at 373 K. Then, heating from 143 to 373 K at a rate of 10 K/min was performed. During the experiments, the flow of nitrogen was maintained at 60 mL/min.

**Broadband Dielectric Spectroscopy.** The dielectric spectra were collected in the frequency range of 1010 to 107 Hz using a Novocontrol system that included an Alpha-A Impedance Analyzer and a Quattro Cryosystem temperature control unit. The samples were measured using a parallel-plate dielectric cell made of invar and sapphire. The separation between the electrodes was maintained at 47 μm which yielded a geometrical capacitance of 21 pF. The measurements were performed from high to low temperatures. The dielectric cell was placed inside the cryostat with a nitrogen atmosphere and equilibrated at 380 K for at least 1 h. The experimental results were reproducible, which indicated the samples did not degrade upon heating and contained no residual solvent. Before acquiring the dielectric spectra, the samples were equilibrated for at least 15 min to achieve thermal stabilization within 0.2 K after each temperature step. Additional technical details regarding the measurements with BDS can be found in ref 38.

**Measurements of Density.** The mass of the samples was measured using a weight balance (Mettler Toledo NewClassic MF model MS105DU) with an accuracy of 0.01 mg. The gas pycnometer (Micromeritics Accupyc II 1340) measured the volume of penetrable helium gas within the sample and sample cup (total volume 0.1 cm3) until the pressure did not vary more than 0.005 psig/min during the equilibration period. The accuracy of the volume measurement is 0.0001 cm3. This cycle was repeated five times for reproducibility. The density was then determined from the mass-to-volume ratio, and the error of this measurement is less than 0.3%. All the density data were obtained at atmospheric pressure and T = 293 K.

**Rheology Measurements.** The rheological measurements of IL–OAPS nanocomposites were performed on an Anton Paar MCR302 with a CTD600 environmental chamber and an EVU20 evaporation unit. The temperature control had an accuracy of ±0.1 K. Small amplitude oscillatory shear (SAOS) with a strain amplitude, γ, of 0.1% has been performed at angular frequencies between 103 and 104 rad/s on a pair of parallel plates with a diameter of 8 mm. For each sample, we measured the SAOS spectra at temperatures from the glass transition temperature Tg up to Tg + 100 K. The linear viscoelastic spectra master curves over a wide frequency range were constructed by employing the frequency–temperature superposition. To directly determine the steady-state (zero) shear viscosity of the IL...
and composites, creep experiments were carried out in the same geometry with an applied stress of 100 Pa at high temperatures and up to 10,000 Pa at temperatures close to \( T_g \).

**FTIR Spectroscopy.** Temperature effects on infrared adsorption were measured using an Agilent Cary 680 FTIR spectrometer equipped with a heated attenuated total reflection stage (Specac, GoldenGate). The interferometer mirror speed was 5 kHz, the resolution was set to 4 cm\(^{-1}\), and 258 scans were integrated for the sample and backgrounds. The sample was allowed to equilibrate for 15 min at the given temperature before measurement. The temperature was scanned in both directions and the spectra showed less than 5% difference between the scans.

**PFG-NMR Measurements.** The PFG spin-echo sequence was applied to measure the individual diffusion coefficients of \(^1\)H and \(^3\)F nuclei. The measurements were carried out at 298, 313, 333, 353, and 373 K using a Varian Direct Digital Drive NMR spectrometer operating at 300 MHz for proton and 283 MHz for \(^3\)F and using a DOTY Z-spec diffusion probe and gradient driver with a maximum field gradient of 1200 G/cm. The duration of the gradient pulse \( \delta \) ranged from 1–4 to 100 ms, respectively. The gradient strength, \( g \), was varied for each experiment (32 values, linear increase, \( g = 0–1200\) G/cm). The self-diffusion coefficients, \( D \), were then extracted by fitting the decay echo signal with the Stejskal–Tanner equation:

\[
I = I_0 \exp[-D0^2g^2y^4(\Delta - \delta/3)]
\]

where \( I \) is the amplitude of the attenuated echo signal, \( I_0 \) is the initial intensity, and \( y \) is the gyromagnetic ratio.

### RESULTS

The chemical structures of the IL and OAPS are presented in Figure 1 while their physicochemical characteristics are collected in Table 1.

![Figure 1. Chemical structures of OAPS NPs and the IL.](image)

**Thermal Characteristics of the Studied Materials.**

DSC was used to assess the glass transition temperature of the composites. The corresponding thermograms are presented in Figure 2 with the \( T_g \) values summarized in Table 1. As can be seen, the increase in OAPS concentration leads to an increase in \( T_g \) (Table 1) and to a substantial broadening of the glass transition step (inset of Figure 2). The latter is associated with the increasing distribution of segmental relaxation times of the IL in the composite that arises because of heterogeneity that scales with an increase in particle concentration.

**Conductive and Viscoelastic Properties of Studied Samples.** The dielectric data of pure IL and IL–OAPS composites were gathered over a wide temperature range. The results obtained for IL–OAPS \(^{30}\) wt % are presented in conductivity and modulus terms in Figure 3A,B, respectively.

![Figure 3. (A) Real part of the conductivity spectra for IL–OAPS \(^{30}\) wt %. The plateau level was used to estimate the dc conductivity \( \sigma_{dc}(T) \). (B) Imaginary part of the dielectric modulus, with the peak position providing the characteristic time of ion relaxation.](image)

The conductivity spectra exhibit typical behavior for ionic materials with a frequency-dependent ac conductivity at higher frequencies, followed by a dc conductivity plateau \( \sigma_{dc} \) and further by a large decrease because of electrode polarization effects at lower frequencies.\(^{39}\) The AC–DC transition is visible in the imaginary part of the electric modulus (\( M' \)) representation as a well-resolved peak with the maximum

| sample                  | \( \rho \), g/cm\(^3\) | \( T_g \), K | \( \sigma_{dc} \) at \( T_g \), S/cm | wt % OAPS | \( \sigma_{dc} \) at 333 K S/cm | \( t_1/\tau_0 \) (333 K) |
|-------------------------|-------------------------|-------------|-------------------------------------|------------|-------------------------------|------------------------|
| pure IL                 | 1.20                    | 196         | 1 x 10\(^{-15}\)                    | 0          | 8.55 x 10\(^{-4}\)           | 0.44/0.56              |
| IL–OAPS \(^{10}\) wt %  | 1.22                    | 200         | 1 x 10\(^{-15}\)                    | 10         | 4.3 x 10\(^{-4}\)           | 0.48/0.52              |
| IL–OAPS \(^{30}\) wt %  | 1.30                    | 215         | 2.5 x 10\(^{-14}\)                  | 30         | 7.7 x 10\(^{-3}\)           | 0.59/0.41              |
| IL–OAPS \(^{50}\) wt %  | 1.51                    | 247         | 4 x 10\(^{-13}\)                    | 50         | 2.3 x 10\(^{-3}\)           | n/a                    |

\(^{39}\)TGA was used to estimate the weight fraction of OAPS. \( T_g \) was measured by DSC. BDS was used to measure the dc conductivity, and the results of PFG-NMR were used to estimate the transport number of cations/anions (\( t_1/\tau_0 \)).
defining the conductivity relaxation time, \( \tau_\sigma = 1/2\pi f_{\text{max}} \). Upon decreasing the temperature, the \( M^*(f) \) peak shifts toward lower frequencies and moves out of the experimental frequency window below the calorimetric \( T_g \).

In ILs, the (dipolar) structural relaxation cannot be accessed via BDS because of the high ion conductivity; therefore, rheology was employed to extract the structural relaxation time and the viscosity of the studied systems. The mechanical data of IL–OAPS30 wt % in the form of master plots are presented in Figure 4. The horizontal shift factors employed to construct the master plot of the shear modulus components \( G'(\omega) \) and \( G''(\omega) \) are presented in the inset of Figure 4A. In the following, the structural relaxation time \( \tau_s \) refers to the characteristic time of the high-frequency loss peak observed for \( G''(\omega) \), as shown in Figure 4A, reaching the calorimetric \( T_g \) value of about 100 s. The zero shear viscosity was extracted from the creep compliance \( J(t) \) data (Figure 4B). The slope of unity at a long time indicates the long-range flow regime of IL–OAPS30 wt %. The IL–OAPS30 wt % sample was measured only with the creep compliance, \( J(t) \), while the other (modulus) rheological measurements were difficult to implement because of the high stiffness of this material. Nevertheless, the \( J(t) \) measured at the constant stress contains the same set of dynamic information as the linear viscoelastic spectra of the storage modulus measured from the SAOS.

**Diffusion Measurements.** PFG-NMR measurements were used to study the diffusion of ions in pure IL, IL–OAPS10 wt %, and IL–OAPS30 wt %. The diffusion coefficient of ions in the IL–OAPS50 wt % sample, as well as that of OAPS NPs, could not be measured because of their low mobility. The results of NMR measurements for the diffusion coefficients of \( ^1H \) and \( ^19F \) nuclei, a representative for cations and anions, respectively, are presented in Figure 5 and discussed below.

**DISCUSSION**

**Charge Transport in the Supercooled Liquid State.** The dc conductivity data of all studied materials are presented in Figure 5A. As recognized here, there are two characteristic regions separated by \( T_g \), and the conductivity at \( T_g \) is marked with colored horizontal arrows. Above \( T_g \), the dc conductivity follows the Vogel–Fulcher–Tamman (VFT) behavior, while below \( T_g \), it can be described by the Arrhenius equation. Overall, under any constant \( T \)-condition, the dc conductivity decreases with the increase in OAPS concentration, that is, it becomes more and more separated from \( \sigma_{dc} \), corresponding to the pure IL.

Because the dc conductivity is directly related to the number of ions and their mobility, the observations coming from the BDS technique should be also reflected in ion diffusion. As depicted in Figure 5B, the ion diffusivity measured by PFG-NMR slows down with an increase in OAPS concentration. Additionally, for IL–OAPS50 wt %, in the temperature range above 350 K, cations diffuse slower than anions, resembling the diffusive behavior of ions in the pure IL. The slower diffusion of the \( [N_{22268}]^+ \) cation can be assigned to its large size further influenced by the ability of the alkyl tail (Figure 1) to form self-assembled structures.43,42 In the temperature regime below 350 K, the diffusivity of anions in IL–OAPS50 wt % drops below that of cations. Such an unexpected switch in the diffusion coefficient reflects some changes in the diffusive environment with the increase in the OAPS concentration. Indeed, some evidence of heterogeneity growth in the composites with an increase in the concentration of OAPS exists in broadening of the glass transition temperature step (Figure 2) observed from DSC.

Overall, our analysis of conductivity and diffusivity shows that they both decrease with the increase in the OAPS concentration. To understand the underlying mechanism of this decrease, we examine the effect of the glass transition temperature on the conductive behavior. This dynamic parameter affects the conductivity in the systems in which the structural relaxation directly controls ion transport. Such a coupling between conductivity and structural relaxation has been reported for many aprotic ILs.43–45 As shown in the inset of Figure 5A, the addition of OAPS NPs leads to an increase in \( T_g \). From scaling the conductivity plot in Figure 5A by \( T_g \) (Figure 6A), it can be seen that the conductivity data at high \( T \) nearly coincide with each other, suggesting that, indeed, in this temperature range, the \( T_g \) parameter is the major driver of the conductivity changes.

**Ion Dynamics in the Vicinity of Liquid–Glass Transition.** The same \( T_g \)-based argument, however, cannot be used to explain behavior in the region near \( T_g \) (see Figure 6A), where the conductivity data for the composites strongly...
deviate from the IL curve. This is especially noticeable for the IL−OAPS30 wt% and IL−OAPS50 wt% composites.

For the studied composites, different values of \( \sigma_{dc} \) are observed at \( T_g \) (Figure 6A). Specifically, \( \sigma_{dc}(T_g) \) is \( \sim 4 \times 10^{-13} \) S/cm for IL−OAPS30 wt% and IL−OAPS50 wt%, \( \sim 1 \times 10^{-15} \) S/cm for IL−OAPS10 wt%, and pure IL, respectively. According to the literature, \( \sigma_{dc}(T_g) \sim 10^{-15} \) S/cm indicates that the charge transport is fully coupled to structural relaxation in an ionic system. This situation is observed in the case of pure IL and IL−OAPS10 wt%. The remaining composites exhibit \( \sigma_{dc} \) at \( T_g \) higher than that of fully coupled systems. This indicates that the samples with high OAPS concentration exhibit different degrees of decoupling of ion transport from structural dynamics. To quantify the magnitude of such a phenomenon, we use the decoupling index defined as \( R_d(T_g) = 15 + \log \sigma_{dc}(T_g) \). This simple approach is often used to estimate the degree of decoupling for various ionic systems without a direct viscoelastic characterization of the materials. For our systems, the largest degree of decoupling (with \( R_d(T_g) = 6 \)) is found for IL−OAPS50 wt% for IL−OAPS30 wt%, this value is smaller and equal to 1.4, while pure IL and IL−OAPS10 wt% practically follow the behavior of fully coupled systems. An alternative method to quantify the decoupling phenomenon is based on the joint analysis of conducting and viscoelastic properties of a given system in terms of the Walden plot. A significant advantage of this approach is the possibility to investigate the decoupling between structural dynamics and ion transport not only at \( T_g \) as is the case for \( R_d(T_g) \), but also over a broad temperature range.

Figure 6B represents the Walden plot where the “ideal” line corresponds to the case of complete coupling of ion conductivity to structural relaxation and a full dissociation of all ion pairs. It has the slope equal to 1 that corresponds to the case of the molar conductivity \( \Lambda \propto \eta^{-1} \) (the steady-state viscosity), and it is well-obeyed by the conductivity of \( K' \) in dilute aqueous solutions (dot-dashed line in Figure 6B). The data above this line correspond to a “superionic” regime, with ionic motions faster than structural relaxation, while the data below the ideal line usually indicate a poor dissociation of ion pairs. In the case of the Walden plot analysis, the decoupling between ion conductivity and structural relaxation is characterized by the exponent \( \alpha \) estimated from the power-law behavior \( \Lambda \propto \eta^{-\alpha} \), where \( \alpha \) is the slope in the corresponding double-logarithmic plot which is connected to the decoupling exponent as \( \epsilon = 1 - \alpha \). Our data demonstrate that in IL and IL−OAPS10 wt%, the conductivity is coupled to structural relaxation, as the slope is very similar to that of the ideal case (\( \epsilon = 0 \)). Such a behavior is in line with the behavior of most aprotic ILs which show vehicle-type conduction. On the other hand, a high concentration of OAPS renders the corresponding systems strongly decoupled over a broad temperature range, in which the charge transport becomes faster than structural dynamics. The decoupling is the strongest and extends over the broadest temperature range in IL−OAPS50 wt%. The decoupling exponent, \( \epsilon \), reached 0.5 for IL−OAPS50 wt%, which, to our knowledge, represents the strongest decoupling reported for IL-containing materials. For comparison, the decoupling exponent of ILs reported in refs 35 and 36 is only \( \sim 0.1 \). Interestingly, for the case of IL−OAPS30 wt%, the conductivity is decoupled over a broad temperature range but at temperatures above \( \sim 350 \) K, the system becomes coupled again as it crosses the ideal line.

As we mentioned above, only a few reports revealed a decoupling phenomenon in classical aprotic ILs. The appearance of decoupling in IL−OAPS resembles more the behavior of polymerized ILs (PolyILs). In these materials, one type of ion is covalently attached to the polymer chains, which makes them significantly less mobile than the ion of the opposite charge sign. Several studies dealing with the underlying mechanisms of decoupling in PolyILs demonstrated the critical effect of the MW and the chain-like dynamics on the decoupling. Specifically, it was shown that the degree of decoupling increases with the increase in MW of the polymer, while the charge transport of monomers and dimers remains closely coupled to the structural relaxation. However, one critical difference between the PolyILs and ILs is that pure ILs do not display suprastructural dynamics. Thus, the questions arise: why does the addition of OAPS make ion diffusion faster than the structure fluctuations? Could it be that the addition of OAPS initiates a suprastructural arrangement in the low-MW IL?

Reversible Gelation and Molecular Origin of the Decoupling Phenomenon in Nanocomposites. To address this point, we turn to the analysis of rheological spectra that are presented in Figure 7.

As seen here, the mechanical complexity of composites increases with the increase in the OAPS concentration. At low frequencies or high temperatures, the presence of the terminal relaxation mode manifesting itself as \( G'(\omega) \sim \omega^\alpha \) and \( G''(\omega) \sim \omega^\beta \) is observed for all composites presented in Figure 7A. Although IL−OAPS50 wt% \( G' \) and \( G'' \) could not be monitored because of the high stiffness of the sample, the creep compliance, \( J(t) \), presented in the inset of Figure 7A demonstrates that even in this composite, the final flow regime can be reached as the slope at a longer time is nearly unity. The presence of this terminal flow regime at high \( T \) reveals that the present composites are heat-processable. This property differentiates them from the conventional polymer electrolyte composites with bigger filler sizes.

Further examination of the rheological spectra presented in Figure 7A reveals that in both IL−OAPS10 wt% and IL−OAPS50 wt%, the structural relaxation peak is separated from the terminal relaxation mode by several orders of magnitude, implying the presence of supramolecular dynamics which contribute to the overall mechanical spectra of these materials.
To identify the presence and evolution of the hydrogen bonds in the composite, we probed the N-H stretching region (3150–3550 cm⁻¹) in the FTIR spectra versus temperature. Figure 8 shows the N-H stretching region of samples IL–OAPS₃₀ wt % (empty symbols) and OAPS (solid lines). The absorbance of samples is scaled with the concentration of OAPS.

In particular, for IL–OAPS₃₀ wt %, the intermediate relaxation regime spans between the terminal relaxation frequency 1/τₓ ~ 10⁻¹² rad/s and the structural relaxation frequency at 1/τₐ ~ 10⁻¹ rad/s.

The evolution of the rheological spectra of IL mixed with OAPS resembles that of PolyILs where the appearance of intermediate relaxation is associated with the chain-like relaxation governed by the polymer MW. However, because the neat IL does not exhibit any chain-like relaxation, the presently uncovered supramolecular relaxations should be associated with the development of long-relaxing structural features triggered by the addition of OAPS. To get some insights into the type of structural organization in the composites, we analyzed the spectral shape of the intermediate frequency regions. In the case of IL–OAPS₃₀ wt %, G' (ω) ~ ω¹ is observed at the intermediate frequency region which points out to the absence of gelation but suggests a presence of rather large slowly relaxing clusters. In IL–OAPS₃₀ wt %, intermediate frequencies G' and G'' display a weak power-law dependence G' (ω) ~ ω⁰.³³ and G'' (ω) ~ ω⁰.²⁸ that is the characteristic of gel-like dynamics. The presence of the region f(t) ~ t¹/₂ spanning over 3 decades in the intermediate time (inset Figure 7A) suggests a presence of supramolecular relaxation and gel-like dynamics also in IL–OAPS₃₀ wt %. The photographs of the gel-like structure of IL–OAPS₃₀ wt % made at room temperature (RT) are presented in Figure 7B. The presence of a terminal relaxation mode at high T for these samples manifesting as G' (ω) ~ ω¹ and G'' (ω) ~ ω for IL–OAPS₃₀ wt % and f(t) ~ t¹ for IL–OAPS₃₀ wt % reflects the reversible nature of the gel-like network in the intermediate frequency region. The photograph demonstrating that IL–OAPS₃₀ wt % flows at high temperature is presented in Figure 7C. Similar effects of the reversible gelation have been reported in the literature for various systems associating through noncovalent interactions.

In our case, we hypothesize that the transient supramolecular assemblies might originate from the hydrogen bond formation in composites.
Scheme 1. Possible Mechanisms of Immobilization of Anions by OAPS

**Effect of a Structural Network on the Transport Number and Decoupling Phenomenon.** As can be seen from Scheme 1, the hydrogen-bond-induced localization of TFSI in the cluster and/or as a part of the network could ultimately lead to weakening of the electrostatic interaction within the IL’s ionic pairs. This may affect both the ions’ dissociation and their diffusivities. In this context, the important question to address is how the structural network formation impacts the diffusion coefficients of the individual ions in ILs.

To address this question, we calculated the transport number for pure IL and mixtures with 10 and 30 wt % of OAPS NPs. The transport numbers of the two types of ions are calculated for cations and anions, respectively, both obtained from PFG-NMR measurements (Figure 5B). The temperature evolution of the transport number for N_{2288} cations (t_+) is presented in Figure 9.

In the temperature interval from 300 to 350 K, t_+ increases with the increase in OAPS concentration. In this temperature range, the stronger contribution of cations to the overall diffusion indicates that the network formation evidenced from rheology slows down (cages) the anions through the H-bonded aggregates, as depicted in Scheme 1 (Scenario 1). This effect is amplified by the increase in OAPS concentration (Scheme 1, Scenario 2), which makes anions even less mobile. This is the case for the IL−OAPS^{30 wt %} mixture where the slowing down of anions results in complete switching in the rate of the diffusion of the ions and significant enhancement in t_− compared to that the pure IL and IL−OAPS^{10 wt %}. Specifically, at ∼313 K, t_− in IL−OAPS^{30 wt %} reaches the value of 0.62, which is about 40% higher than that of the pure IL. The interaction between the anion and OAPS reduces not only the mobility of the anion but also possibly decreases electrostatic interaction in ion pair which leads to enhancement in the cation diffusion. The possible mechanism of diffusion of ions at low T is presented in Scheme 2.

At temperatures above 350 K, the anion diffuses faster than the cation, resembling the diffusion behavior of the pure IL (Figure 5B). In this temperature interval, the terminal relaxation regimes that are present in the rheological spectra of all composites suggest that the H-bonded network is significantly weakened, which leads to dissociation of the anion from the network, enhancing its contribution to overall diffusion. The mechanism of diffusion for the high T region is also sketched in Scheme 2.

The observed effective “on” and “off” switching of the network via the temperature and the concentration of NPs permits modulation of the diffusion of ions and enables control over transport properties.

It is important to mention that the observed slowing down of the anions at the low-temperature limit also triggers the decoupling of ion transport from structural relaxation, and is especially noticeable in IL−OAPS^{30 wt %}. This is visible from both the Walden plot analysis (Figure 6B) and the behavior of the transport number (Figure 9). Indeed, the Walden plot of IL−OAPS^{30 wt %} crosses the ideal line at high T (at ∼350 K the steady-state viscosity ∼10^{-2} Pa·s) (Figure 6B), and the turning point in the behavior of the transport number for IL−OAPS^{30 wt %} also occurs at about 350 K. Therefore, below 350 K, the cation diffusion is decoupled from the viscosity while anions are slowed down by a supramolecular network. At temperatures above 350 K, the reorganization and disappearance of the network restore the diffusive pattern of the pure IL where diffusion is governed by (hence coupled to) the segmental relaxation which is highlighted by the Walden plot. In this sense, the effective incorporation of the anion into the network leads to (i) a decrease in its mobility and (ii) an improvement in ions dissociation by reducing an electrostatic interaction in ion-counterion pairs (Scheme 1). Both these

Figure 9. Temperature dependence of the transport number calculated for cations.
effects enhance the relative diffusion of the cation and promote its decoupling from the fluctuations of the suprastructure.

**SUMMARY**

Combining conductivity, rheology, FTIR, and NMR diffusivity results, we demonstrate that the addition of ultrasmall NPs to a IL results in an enhancement of the transport number of the cation by ~40% at 40 °C. We associate the observed increase with the surface chemistry of NPs that is capable of selective hydrogen bond formation with the anion, slowing down the diffusion of the latter close to RT. The anion–OAPS association is evident from the presence of supramolecular dynamics in rheology and FTIR measurements and the diffusive behavior monitored via PFG-NMR. In particular, at ambient temperature, selective enhancement of the cation diffusion through suppression of anion diffusion with the increase in OAPS concentrations were observed. At a given critical NP concentration, the cation diffusion coefficient exceeded that of the anion which varies from the diffusive behavior of the two ions in the pure IL. The crossover in diffusivity coincides with the onset of the dynamical decoupling between the charge transport and the structural relaxation, with the cations diffusing faster than the the timescale for structural fluctuations. At high temperatures, the anion–OAPS interaction weakens and the diffusion of anions became faster than that of cations, resembling the trivial diffusive behavior of the pure IL. The transient nature of the supramolecular network, observed from the diffusive behavior and reflected in the rheological measurements, suggests technology-desired processability of composites even for a high concentration of OAPS, thus distinguishing our system from conventional polymer composites with larger fillers’ sizes. Our results demonstrate that the formation of transient supramolecular networks can be an effective strategy toward designing melt-processable composites with improved transport number of cations at RT. Furthermore, because of the widespread use of TFSI-based ILs in modern energy-storage technologies, our approach provides a path to control ion diffusion which is important for battery technology.

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Notes
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