Extraordinary Slowing of Structural Dynamics in Thin Films of a Room Temperature Ionic Liquid

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Supporting Information

ABSTRACT: The role that interfaces play in the dynamics of liquids is a fundamental scientific problem with vast importance in technological applications. From material science to biology, e.g., batteries to cell membranes, liquid properties at interfaces are frequently determinant in the nature of chemical processes. For most liquids, like water, the influence of an interface falls off on a ~1 nm distance scale. Room temperature ionic liquids (RTILs) are a vast class of unusual liquids composed of complex cations and anions that are liquid salts at room temperature. They are unusual liquids with properties that can be finely tuned by selecting the structure of the cation and anion. RTILs are being used or developed in applications such as batteries, CO$_2$ capture, and biological processes. Here, it is demonstrated quantitatively that the influence of an interface on RTIL properties is profoundly different from that observed in other classes of liquids. The dynamics of planar thin films of the room temperature ionic liquid, 1-butyl-3-methylimidazolium bis(trifluoromethylsulfonyl)imide (BmimNTf$_2$), were investigated using two-dimensional infrared spectroscopy (2D IR) with the CN stretch of SeCN$^-$ as the vibrational probe. The structural dynamics (spectral diffusion) of the thin films with controlled nanometer thicknesses were measured and compared to the dynamics of the bulk liquid. The samples were prepared by spin coating the RTIL, together with the vibrational probe, onto a surface functionalized with an ionic monolayer that mimics the structure of the BmimNTf$_2$-Near-Brewster’s angle reflection pump-probe geometry 2D IR facilitated the detection of the exceedingly small signals from the films, some of which were only 14 nm thick. Even in quarter micron (250 nm) thick films, the observed dynamics were much slower than those of the bulk liquid. Using a new theoretical description, the correlation length (exponential falloff of the influence of the interfaces) was found to be 28 ± 5 nm. This very long correlation length, ~30 times greater than that of water, has major implications for the use of RTILs in devices and other applications.

I. INTRODUCTION

Liquids at interfaces occur in a wide variety of chemical and biological systems. Molecules in a liquid in contact with an interface will have distinct properties from those in the bulk liquid. The structure and dynamics of liquid molecules at an interface and for some distance beyond the layer in direct contact with an interface are distinct from those of the bulk liquid. The structural and dynamical properties of molecules at an active interface, e.g., a battery electrode or a cell membrane, can be intimately involved in chemical processes.

Molecules in an interfacial layer will see the interface material on one side and the liquid molecules on the other. The molecules at the interface, which will not have bulk properties, i.e., structural organization and dynamics, will affect the next layer of molecules further from the interface, which in turn will affect the next layer. Generally, the influence of an interface dies off rapidly. After several liquid layers, typically on a distance scale of one to several nanometers, the influence of the interface is no longer present. Molecules at interfaces typically exhibit slower structural dynamics than those in bulk liquids. Thus, the measurement of the dynamical time scales provides a useful approach for quantifying the influence of an interface and the length scale over which the interfacial perturbation propagates. For example, in the case of water molecules confined in sodium bis(2-ethylhexyl) sulfosuccinate (AOT) reverse micelles, the dynamics of water molecules located within ~2 nm from the interfaces are significantly altered, slowing substantially; however, water molecules further away from the interface behave like those in bulk water.

Similar length scales have been reported in other liquid/ interface systems. Room temperature ionic liquids (RTILs) are generally composed of organic cations and inorganic anions. RTILs can be highly structured, even on mesoscopic distance scales, due to the Coulomb, van der Waals, and hydrogen-bonding interactions among the bulky ions. RTILs are being used or investigated for a wide variety of applications. In many of these applications, such as electrolytes in batteries, solvents for CO$_2$ capture in supported ionic liquid membranes,
and media for biological studies, the RTILs will be in contact with interfaces or confined on nanometer to submicron length scales. Recently there have been qualitative reports that indicate RTILs may display the effect of interfaces on distance scales much longer than those found in other types of liquids, with estimated length scales varying from 60 to 2000 nm depending on the types of the RTILs and the interface. A molecular dynamics (MD) simulation by Voth and co-workers indeed showed long-range structural ordering of alkylmethylimidazolium-based ionic liquids at the liquid/vacuum interface. Particularly for the longer-chain cations, the interfacial structural ordering was found to persist beyond ~10 nm from the interface. A recent MD simulation by Margulis and co-workers also revealed that 1-methyl-3-octylimidazolium octylsulfate can form a lamellar structure at the liquid/vacuum interface, and the structure extends to the full length scale of their simulation box (9 nm). These measurements clearly demonstrate that the effects of the interfaces on dynamics propagate over much longer distance scales in the RTILs than in other liquids. However, the very broad distribution of the pore sizes (100–500 nm) and the complicated topography of the pores hindered the quantitative evaluation of distance scales.

In this paper we present two-dimensional infrared (2D IR) spectroscopy studies on planar thin films of the RTIL 1-butyl-3-methylimidazolium bis(trifluoromethylsulfonyl)imide (BmimNTf2). The CN stretch of SeCN−, introduced into the RTIL as BmimSeCN, was used as the vibrational probe because of its strong transition dipole moment and long vibrational lifetime. The recently developed method of 2D IR spectroscopy in the near-Brewster’s angle reflection pump–probe geometry facilitated the detection of the small signals from the thin films. Films with the controlled thicknesses from 14 to 278 nm were investigated. In contrast to the experiments in the PES membranes, the films have well-defined thicknesses and topography, making it possible to obtain a quantitative evaluation of the distances over which the interfaces influence the dynamics of the RTIL. The 2D IR experiments showed that even quarter micron thick films exhibited dynamics that were significantly slower than the bulk liquid. As the films were made thinner, the dynamics slowed dramatically.
II. PREPARATION OF THE RTIL THIN FILMS

Several publications have reported the coating or deposition of ionic liquids on flat dielectric and metallic surfaces. In many of these studies, the focus was on the formation of nanodroplets of RTILs rather than planar films of RTILs of interest here. Successful preparation of the ionic liquid thin films has been reported using ultrahigh vacuum physical vapor deposition (UHV-PVD) methods, requiring complex apparatus that is not widely available.

Spin coating is a well-established method for preparing thin films. Spin coating BmimNTf2 RTIL directly onto an SiO2 surface using the RTIL dissolved in methanol yielded droplets rather than a planar film, as seen in the microscopy optical image shown in Figure S1 (see the Supporting Information). This result showed that the affinity between the BmimNTf2 RTIL and the SiO2 surface is not high enough to overcome the surface tension, preventing the formation of an extended planar film.

To enhance the affinity between the BmimNTf2 RTIL and the interface, the SiO2 surface was functionalized with an ionic monolayer that mimics the structure of BmimNTf2 (Figure 1A) by adapting a procedure reported by Xin and Hao. Spin coating of BmimNTf2 RTIL on the functionalized surface yielded a film with high optical quality, as seen in the microscope images (Figure 1B) acquired with a Raman microscope (Horiba XploRA). Micro-Raman spectra acquired at spatial points along the green line in Figure 1B clearly show the strong NS stretching mode originating from NTf2− anion (Figure 1C). The intensity of the NS stretch band at each point is plotted in Figure 1D, demonstrating that the thickness of the functionalized SiO2 ionic monolayer (~278, 112, and 39 nm) is significantly thinner than any of the RTIL films (~100 nm).

The thickness of the spin coated films was controlled by the concentration of the RTIL in the precursor methanol solution. To introduce the SeCN− anion as the vibrational probe for the 2D IR spectroscopy, BmimSeCN was mixed with BmimNTf2 in a 1:10 molar ratio. BmimSeCN only minutely perturbs the dynamics of BmimNTf2 RTIL (Figure S2). The thicknesses of the spin coated films were characterized with FTIR absorption spectra by comparing the peak heights of the CH stretching modes of the Bmim+ cation with those of the bulk BmimNTf2 liquid with a known thickness (Figure S3). Further details on the sample preparation and characterization can be found in the Supporting Information. All spin coating was conducted in a nitrogen-filled glovebox. The samples were then sealed in airtight sample cells in the glovebox. Thus, the samples were not exposed to either atmospheric humidity or oxygen.

III. RESULTS AND DISCUSSION

A. Linear Spectra. The normalized FTIR spectra of the CN stretch of SeCN− in bulk BmimNTf2 and in films with thickness of 278, 112, and 39 nm are shown in Figure 2. The 39 nm film, for example, has a CN absorbance of only 1 mOD. The peak center positions and the line widths for Gaussian fits are given in Figure 2. All of the bands have the same center position within experimental error, 2062.5 ± 0.5 cm−1. The bulk and 278 nm film spectra have the same fwhm within error, 22.6 cm−1. The fits give an increasing fwhm for the thinner films, 24.8 and 27.2 cm−1, for the 112 and 39 nm films, respectively. Imperfect baseline subtractions can produce systematic error, which might be responsible for the different line widths. The same measurement was applied to the thinnest 14 nm film as well. Though the data are not included in Figure 2, the fit to the 14 nm film yielded a fwhm width of ~25.5 cm−1. Though the line width may be increasing as the film thickness is reduced (by up to 20% in the 39 nm film compared with the bulk liquid), overall the peak positions and line widths in the films are not substantially altered from those of the bulk liquid. As discussed below, in spite of the lack of substantial differences in the linear absorption spectra, the dynamics of the films are very different from those of the bulk liquid and depend strongly on thickness.

B. 2D IR Spectroscopy on the RTIL Thin Films. As seen in Figure 2, the infrared absorption band of the SeCN− anions has a finite bandwidth. The width, i.e., the variation of the CN stretch vibrational frequency (inhomogeneous broadening), arises from different RTIL structural configurations, which cause distinct interactions between the SeCN− anions and their surroundings. The RTIL undergoes structural fluctuations, inducing the CN frequencies to change with time. At sufficiently long time, each CN will sample all frequencies in the range of frequencies. Thus, measuring the time evolution of the vibrational frequencies (spectral diffusion) reports directly on the RTIL structural dynamics.

Two-dimensional IR spectroscopy characterizes the spectral diffusion by making a correlation plot between the initial and the final frequencies. The measurements of the initial and final frequencies are separated by a waiting time T (Figure 3). As seen in Figure 3A,B, 2D IR spectroscopy involves three input pulses. The first two “pump” pulses together label the initial frequencies (2D spectrum horizontal axis ω) and after the time T, the third “probe” pulse initiates the emission of
The first two "pump" pulses label the initial frequencies, and after a waiting time $T_w$, the third "probe" pulse induces the emission of the vibrational echo signal, which reads out the final frequencies. The emitted signal field interferes with the transmitted probe pulse as a local oscillator, and the amplitude of the interference is recorded as the signal. (B) Two-dimensional IR spectroscopy implemented in the near-Brewster angle reflection geometry for ionic liquid thin films. The two pump pulses and the probe pulse play the identical roles as in the standard geometry. The emitted signal field interferes with the heavily attenuated reflected probe field, which enhances the signal to local oscillator ratio, greatly improving the detection. (C) Two-dimensional IR spectra for the bulk liquid (RTIL) and thin films (RTIL Thin Film) acquired at the waiting time $T_w = 100$ ps. More correlated band shapes in the thinner films indicate slower structural dynamics.

The vibrational echo pulse, which reads out the final frequencies (vertical axis, $\omega_m$). If $T_w$ is small compared with the dynamical time scales of the system, the initial and the final frequencies will be similar and yield a diagonally elongated band shape. In contrast, when $T_w$ is sufficiently large, the structural fluctuations during the waiting time substantially randomize the vibrational frequencies, and the initial and final frequencies lose correlation, yielding a more rounded band shape.

Figure 3C shows 2D IR spectra for three samples at a single waiting time, $T_w = 100$ ps. The signs of the 2D IR band shapes are different between the bulk sample and the thin films because they were measured in different geometries as shown in Figure 3A (the transmission geometry for the bulk liquid sample) and Figure 3B (the reflection geometry for films). The bulk spectrum is nearly round, indicating the structural fluctuations in the bulk RTIL during the 100 ps between the labeling and reading have almost randomized the vibrational frequency. The nearly round shape shows that almost all of the structural configurations that give rise to the inhomogeneously broadened line have been sampled in 100 ps. In contrast, the spectrum of the 112 nm film is elongated along the diagonal, and the 39 nm sample is substantially more elongated. That is, during the 100 ps, the vibrational frequencies were not significantly altered, demonstrating that the structural fluctuations in the thin films occur on much slower time scales than that of the bulk RTIL. It is qualitatively clear that the thinner films exhibit slower dynamics.

C. Center Line Slope (CLS) Decays and Spectral Diffusion Time Constants. The dynamical time constants are obtained by evaluating the evolving band shapes of the 2D spectra with increasing $T_w$. The $T_w$-dependent changes in the band shapes can be analyzed quantitatively using the center line slope (CLS) method. The CLS($T_w$) decay is the normalized frequency-frequency correlation function (FFCF). Figure 4 shows the CLS decays for each sample. As the sample becomes thinner, there is a dramatic slowing of the decay, reflecting a slowing of the structural dynamics of the sample.

The solid curves through the data are multieponential fits, with the fitting time constants given in Table 1. Both the bulk sample and the 278 nm film were fit well with the same functional form (a triexponential), and both decay close enough to zero and have sufficient slope that it is reasonable to assume that there is no slower component that cannot be observed in the 200 ps limit on the data collection due to the vibrational lifetime.

For the bulk sample and the 278 nm films, the three time constants, $\tau_1$, $\tau_2$, and $\tau_3$, are given in Table 1 and can be directly compared. The three time constants reflect time scales for different types of structural dynamics. The dynamics on all three time scales slow substantially, by factors of ~2 to ~3, in the 278 nm films compared with the bulk liquid. These results show the dramatic slowing of the dynamics in the film even though the film is approximately a quarter of a micron thick.

While the bulk data and the 278 nm film data fit extremely well to triexponential fits, fits to the CLS decays for the other three film thicknesses (112, 39, and 14 nm) are biexponentials. For fitting purposes, it was assumed that the slower component of each curve decays to zero (see further discussion below). The fit time constants are $\tau_1$ and $\tau_2$ in Table 1. In the biexponential fits to these three thinner films, the slower time constants ($\tau_2$) were slower than $\tau_1$ in the bulk liquid and 278 nm film.
Table 1. CLS Parameters from Fits

| sample          | $T_w = 0$ valuea | $\tau_1$ (ps) | $\tau_2$ (ps) | $\tau_3$ (ps) | $\tau_4$ (ps) | normalized CLSb at $T_w = 150$ ps |
|-----------------|------------------|--------------|--------------|--------------|--------------|----------------------------------|
| bulk liquid     | 0.76             | 2.5 ± 1      | 15 ± 2       | 62 ± 3       | 30 ± 1       | 0.03                             |
| 278 nm          | 0.84             | 5.0 ± 0.7    | 46 ± 17      | 136 ± 8      | 115 ± 7      | 0.27                             |
| 112 nm          | 0.88             | 2.0 ± 0.6    | 136 ± 2      | 132 ± 2      | 0.32         |
| 39 nm           | 0.90             | 3.8 ± 2      | 278 ± 7      | 270 ± 7      | 0.57         |
| 14 nm           | 0.91             | 4.0 ± 1.2    | 484 ± 14     | 479 ± 10     | 0.73         |

aThe difference between the $T_w = 0$ value and 1 is directly related to the homogeneous line width. The closer the value is to 1, the narrower the homogeneous line width. The value of the CLS at 150 ps after the curves have been normalized to 1 at $T_w = 0$ ps to eliminate the differences caused by the varying homogeneous line widths.

nm film, and as the thickness was reduced $\tau_2$ slowed substantially. This behavior is in accordance with the general behavior observed so far: i.e., the thinner films exhibit increasingly slower dynamics. In contrast, the faster component ($\tau_1$) is faster than the fastest component $\tau_1$ observed in the 278 nm film (see Table 1). One possibility that can explain this seemingly anomalous behavior is a substantial slowing of some of the dynamical components that are responsible for the ultrafast fluctuations that contribute to motional narrowing in the bulk liquid. Motional narrowing occurs when $\Delta \times \tau < 1$, where $\Delta$ is the range of the frequency fluctuations, and $\tau$ is the time for sampling these frequencies. The motionally narrowed dynamical component contributes to the Lorentzian-shaped homogeneous line width in the linear absorption spectrum, and reduces the CLS value at $T_w = 0$ ps from 1 in the 2D spectrum; the motionally narrowed component does not contribute to the CLS decay. As is evident from Figure 4, the CLS value at $T_w = 0$ ps increases as the thickness is reduced, indicating that the homogeneous line width has decreased. It is likely that some of the homogeneous line width is converted to inhomogeneous line width as $\tau$ slows, and consequently, $\Delta \times \tau < 1$ is no longer satisfied. The new inhomogeneous component will undergo fast spectral diffusion as originally it experienced faster fluctuations that caused motional narrowing. The result is the fast $\tau_1$s and smaller homogeneous line widths in the three thinnest films.

To compare all samples, in spite of the fact that they do not fit to the same functional form, the correlation times $\tau_i$ (the time integrals of the normalized decays) were calculated assuming the slowest component of each curve decayed to zero. Due to the observable time range, limited by the vibrational lifetime of the vibrational probe (~100 ps), we could not exclude the possibility that even slower dynamics exist in the thinner films. The correlation times for the thinner films listed in Table 1 and on Figure 4 should therefore be regarded as lower bounds.

D. Origin of the Observed Dynamical Slowing. Before discussing quantitatively the thickness dependence of the thin films’ structural dynamics, it is illuminating to consider the possible physical mechanisms of the observed dynamical slowing. An MD simulation by Del Pópolo and Voth provides important insights.45 Based on the physical picture from their MD simulation, the SeCN$^-$ anion vibrational probe is trapped in a “long-lived cage” formed by the surrounding cations and anions. The dynamics on faster time scales (<10 ps) may arise from very local motions of the cage and the SeCN$^-$ anions. The dynamics occurring on slower time scales (>100 ps) are associated with longer distance scale structural evolution that structurally relaxes the cages.

As seen in Figure 4 and Table 1, the dynamics of the thinner films are characterized by the emergence of extremely slow dynamics ($\gg$100 ps), demonstrating that the influence of the interfaces persists for much longer times. The interfaces impose global structural and/or dynamical ordering through long-range Coulomb interactions, which result in the extensive length scale of the “cages” surrounding individual cations and anions.

The dynamical slowing in supercooled liquids near glass transition temperatures ($T_g$) has often been discussed in a similar manner, i.e., the growing dynamical correlation lengths of cooperatively rearranging regions as the liquids are cooled toward $T_g$.46–50 In this sense, the dynamical behavior of ionic liquid molecules at the interfaces may resemble that of a supercooled liquid near $T_g$. The observations here might provide insights into the behavior of supercooled liquids under confinement.41,52

E. Quantifying the Correlation Length. A main interest here is the quantitative determination of the correlation length, i.e., the length scale over which the interfaces affect the dynamics of the RTIL molecules. Instead of the correlation times discussed above, which have ambiguity due to the limited observable time range, we define and evaluate the correlation length based on the values of the normalized CLS decays at $T_w = 150$ ps, which have well-defined values for all of the decays in Figure 4. The values are listed in Table 1 and plotted in Figure 5A. The decays were normalized to eliminate the influence of varying CLS values at $T_w = 0$ ps (given in Table 1) originating from varying homogeneous line widths. The value of $T_w = 150$ ps was selected because of the pronounced differences among the curves and good S/N ratios.

Figure 5B shows a schematic illustration of the RTIL film and the dynamical behavior at each location inside the film. The interfaces are separated by a distance, $d$. Far enough from the interface, the system will have bulk dynamics. Near the interface, the dynamics are slower. As a result, the frequency–frequency correlation function (FFCF) at $T_w = 150$ ps, $f(r, T_w = 150$ ps), varies within the film depending on the distance $r$ from the interface. We assume that the influence of the interface falls off exponentially with the distance, $r$, from the interface. Then the dependence is described by

\[ f(r, T_w) = \left[ f_0(T_w) - f_b(T_w) \right] \exp\left(-r/l\right) + f_b(T_w) \]  

where the waiting time $T_w$ here is 150 ps, and $f_b(T_w)$ and $f_b(T_w)$ are the frequency–frequency correlation functions of a thin slab at the interface ($r = 0$) and in the bulk region ($r \rightarrow \infty$), respectively. In this model, the characteristic length scale of the interfacial influence is specified by the correlation length $l$, which describes the exponential falloff of the interfacial effect as the distance $r$ from the interface is increased.

In the 2D IR measurements, all of the molecules inside a film contribute to the signal, and therefore the data reflect the
The red curve in Figure 5A is the fit, which yields the correlation length \( l = 28 \pm 5 \) nm. The equivalent correlation length for water molecules confined in AOT reverse micelles has been estimated to be \( \sim 1 \) nm, and thus the correlation length obtained here for the RTIL films is roughly 30 times longer. The value of \( l \) is consistent with the fact that even a sample that is \( 278 \) nm thick displays dynamics that differ significantly from bulk dynamics. For \( d = 278 \) nm, the center of the sample, \( d/2 \), is \( \sim 139 \) nm from the interfaces, which is \( <1l \). Most of the sample volume will feel the influence of the interfaces.

Note that, in the theoretical model, the influences of the two different surfaces (ionic monolayer surface and 1 atm nitrogen gas) on the RTIL were assumed identical. While the solid surface is functionalized with cations, the previous molecular dynamics simulation demonstrated that RTIL/vacuum interfaces should also be cation-rich. Therefore, the correlation lengths imposed by these two interfaces might be similar. The single \( 28 \) nm correlation length obtained here should be regarded as the “average” correlation length originating from these two interfaces.

**F. Effect of Thickness Variation.** As discussed in the Supporting Information, measurements using Raman microscopy show that there are fluctuations in the thickness as a function of position across the film. For the thin samples, the standard deviation of the range of thicknesses can become large, and even for the thicker samples, the variation is not negligible. However, the variations occur over short distances. The laser spot size that gives rise to most of the signal is \( \sim 180 \) \( \mu \)m in diameter, which is large compared to the distances over which the thickness variations are observed (\( <10 \) \( \mu \)m). Therefore, the observed signal is the average over the thickness variation. It is important to address how much the variation in thickness within the laser spot will affect the observed results for various film average thicknesses, \( d_{ave} \).

To address the influence of thickness fluctuations within the laser spot area, we can begin with eq 3. It is necessary to integrate eq 3 over the variations in \( d \) for a particular average value of \( d_{ave} \). The probability distribution of the thickness, \( P(d) \), is taken to be Gaussian with a standard deviation \( \sigma \) about the average thickness, \( d_{ave} \), for a given sample. In addition, the signal varies linearly in \( d \). A thicker region of the sample within the laser spot will give more signal. To account for this, there is an additional multiplicative factor, \( d/d_{ave} \).

\[
f_{ave}(T_w) = \int_{-\infty}^{\infty} \left(2/d\right)f_b(T_w)\int_0^{d/2} \exp(-r/l) \, dr + \left(2/d\right)\int_0^{d/2} f_b(T_w) \, dr
\]

For simplicity, the influence of the two interfaces is taken to be the same, so the integral is performed from 0 to \( d/2 \). The term \( (2/d) \) is the normalization constant. The integration of eq 1 over \( r \) yields the observed FFCF, \( f_{obs}(d, T_w) \), for a film with thickness \( d \) given by

\[
f_{obs}(d, T_w) = (2/d)f_b(T_w)\int_0^{d/2} \exp(-r/l) \, dr + (2/d)\int_0^{d/2} f_b(T_w) \, dr
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\]

The result of the integration is

\[
f_{ave}(T_w) = \frac{2[lf_b(T_w) - f_b(T_w)]}{d_{ave}\sqrt{2\pi\sigma^2}} \int_0^{\infty} [1 - \exp(-d/2l)] \exp(\frac{-(d - d_{ave})^2}{2\sigma^2}) \, dd + f_b(T_w)
\]

Equation 6 is a useful result. It is similar to eq 3 except for the extra term in the exponential, \( (\sigma^2/8l^2) \). To determine the
influence of thickness variation within the laser spot, we need to compare the magnitude of $\sigma^2/8l^2$ to the other term in the exponential, $d_{AVE}/2l$. From the fit shown in Figure 5A, the correlation length is $l = 28$ nm. For the thickest sample, $d_{AVE} = 280$ nm, and $d_{AVE}/2l = 5$. For this sample thickness, the typical standard deviation is about 0.1 of the thickness, $\sigma = 0.1d_{AVE}$. Then the additional term in the exponential is 0.125, which is negligible compared to $d_{AVE}/2l = 5$. The thinnest sample has a much larger fractional variation. For the thinnest sample, $d_{AVE} = 14$ nm, and $d_{AVE}/2l = 0.25$. $\sigma \sim 0.4d_{AVE}$ (see the Supporting Information). Then the extra term in the exponential is $5 \times 10^{-3}$, which is negligible compared to 0.25. The net result is that the variation in thickness within the laser spot size does not affect the estimation of the correlation length which was based on the average thickness, $d_{AVE}$.

IV. CONCLUDING REMARKS

Liquid molecules in contact with an interface will always have properties that are different from those of the bulk liquid. The distinct intermolecular interactions between the interface and the liquid molecules and the topography of the interfacial layer will cause the liquid contact layer to have different structural and dynamical properties from those of the bulk liquid. For most liquids, after a few molecular diameters (1-2 nm) from the interface, the liquid’s intermolecular interactions will overcome the influence of the distinct interface–molecule interactions, and the liquid will have bulk properties. However, these trends may not apply when the liquids in question are room temperature ionic liquids. In this study we have shown quantitatively that an ionic liquid can have nonbulk properties far from an interface. The difference between the behavior of conventional liquids and RTILs is presumably caused by the strong Coulomb interactions among the complex cations and anions that comprise the RTIL.

To investigate the influence of interfaces on RTILs, we have presented a detailed study of thin films of BmimNTf2. The RTIL was spin coated onto a surface functionalized with an ionic monolayer, the structure of which resembles that of the Bmim’ cation. The structural dynamics of the films were studied by measuring spectral diffusion using 2D IR spectroscopy on a vibrational probe, SeCN−, dissolved in the films. The film thickness ranged from 14 to 278 nm. Measurements on the films were compared to measurements on the same RTIL bulk liquid.

The structural dynamics of the films were found to be substantially slower than those of the bulk liquid, even for the 278 nm thick films. Based on the correlation times obtained from the observed spectral diffusion dynamics, the thinnest 14 nm film exhibited dynamics that are at least 15 times slower than those of the bulk liquid. A model was developed to quantify the substantial slowing of the dynamics as the films become thinner. Fitting the data in Figure 5A with eq 3, the correlation length $l$ was found to be $28 \pm 5$ nm. The correlation length can be regarded as the length scale over which the influences of the interfaces fall off exponentially. Any interface will have interactions with the liquid molecules that are different from interactions among the molecules themselves. Here, the solid interface is functionalized with cations, and it is likely that the RTIL/air interface is also cation-rich based on MD simulation studies.22 With cation-rich interfaces, the anion solvation will be very different from the bulk liquid, substantially perturbing the ionic structure. This modification of the ion–ion arrangement will propagate far out from the interfaces because of the long-range Coulomb interactions.

The study presented here raises a large number of questions. What effect will the length of cation alkyl chains have on the correlation length? What is the role of the interface properties, e.g., replacing a cationic functionalized interfacial layer with a nonionic but polar layer? How will changing the ions influence the correlation length, for example, replacing NTf2− with BF4−? The slowing of the dynamics of the RTIL induced by a well-defined interface, as reported here, can serve as a benchmark for understanding interfacial effects on RTILs. The results also have implications for interfacial processes, such as the diffusion of ions to a battery electrode, and for possible control of interfacial boundary layer properties by tailoring the nature of the interface and the choice of RTIL.

ASSOCIATED CONTENT

* Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acscentsci.8b00353.

Sample preparation procedure, thickness characterization by FTIR absorption spectra, optical images and Raman microscopy, and ultrafast infrared laser system (PDF)

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Notes

The authors declare no competing financial interest.

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