Local Structure Formation in Alkyl-imidazolium-Based Ionic Liquids as Revealed by Linear and Nonlinear Raman Spectroscopy

KOICHI IWATA,† HAJIME OKAJIMA,‡ SATYEN SAHA,§ AND HIRO-O HAMAGUCHI* ,‡,†

ABSTRACT

We show several pieces of Raman spectroscopic evidence that are indicative of local structure formation in imidazolium-based ionic liquids. Low-frequency Raman spectra of C₄mimX, where C₄mim stands for 1-alkyl(C₃H₇)₃-3-methylimidazolium cation and X represents the anion, exhibit broad bands assignable to collective modes of local structures. Spatial distributions of coherent anti-Stokes Raman scattering (CARS) signals from C₄mim[BF₄] are consistent with local structures whose size increases with increasing n. Picosecond Raman spectra of S₁ trans-stilbene as a "picosecond Raman thermometer" show microscopic thermal inhomogeneity ascribable to local structure formation in C₄mim[BF₄] and C₄mim[PF₆]. We also describe two novel phenomena that we believe are relevant to extraordinary nanoenvironments generated by local structures in a magnetic ionic liquid C₄mim[FeCl₄].

Introduction

Liquids are much less understood than gases and crystals. The structure of molecules in the gas phase can be accurately determined by electron diffraction or high-resolution rotationally resolved spectroscopy. Molecular structures and their three-dimensional arrangements in crystals can be determined by X-ray or neutron diffraction. The diffraction and spectroscopic methods are also applicable to liquids, but with only limited resolution. Diffraction patterns become diffuse and rotational structures lost in the liquid phase. Structural information available for liquids is thus dilute. Molecules are disordered in gases, but they are ordered in crystals. What happens in liquids?

Recent research focus on ionic liquids is casting a new light upon this problem. Ionic liquids are composed solely of ions. Unlike ordinary molecular liquids, Coulomb interaction may play a major role in these liquids. The long-range nature of Coulomb interaction may give rise to structure and dynamics that are unique to ionic liquids but are not associated with molecular liquids. The many novel features of ionic liquids are likely to originate from these unique structures and dynamics. Ionic liquids are extreme examples of liquids that are highly informative and suitable for investigating possible molecular orderings in liquids.

Raman spectroscopy provides vibrational Raman spectra that are often referred to as “molecular fingerprints.” From these fingerprints, we are able to learn in detail the structure and dynamics of molecules (ions). The versatility of Raman spectroscopy enables us to measure Raman spectra and hence comparatively study molecules in different phases or states. In fact, we found the rotational isomerism in 1-butyl-3-methyl-imidazolium (C₄mim)- based ionic liquids by comparing the Raman spectra of supercooled liquid C₄mimCl and those of the two polymorphs of crystalline C₄mimCl, for which crystall and molecular structures are solved. In C₄mim-based ionic liquids, at least two rotational isomers of the C₄mim cation coexist, one with a trans and the other with a gauche conformation with respect to the C₇-C₈ bond of the alkyl group, where C₇ is the second carbon atom of the alkyl group attached to the imidazolium ring. The trans isomer is more stable than the gauche as is the case in nonpolar organic molecular liquids. However, the measured enthalpy differences in C₄mim[BF₄] (n = 3–10) are much smaller than the common value for neat alkane liquids at room temperature. It has also been found that the enthalpy difference increases with increasing the chain length n. These findings indicate that the alkyl chains in liquid C₄mim[BF₄] are not free to change conformation but are subject to strong intermolecular interactions and that these interactions become stronger in longer alkyl chains. Based on these Raman spectroscopic implications,
together with a wide-angle X-ray diffraction result,\(^8\) we have built a working hypothesis that these ionic liquids are not homogeneous at the microscopic level but that they form specific local structures.\(^1\) The possibility of local structure formation in ionic liquids has been discussed from various viewpoints: X-ray diffraction,\(^9,10\) diffusion measurement,\(^11,12\) neutron diffraction,\(^13\) ultrafast photoexcited dynamics,\(^14–18\) and molecular dynamics simulations.\(^19–22\)

In the present Account, we discuss our recent Raman spectroscopic results that further support our working hypothesis. Low-frequency Raman spectra of liquid C\(_n\)mim\([PF_6]\) \((n = 4, 6, 8)\) are shown to exhibit broad features assignable to collective modes of local structures with different size. Spatial distributions of coherent anti-Stokes Raman scattering (CARS) signals from C\(_n\)mim\([PF_6]\) show characteristic broadening ascribed to local structures whose average size is larger for larger \(n\). Picosecond time-resolved Raman results are discussed with a focus placed on the cooling process of photoexcited trans-stilbene in C\(_n\)mim\(\text{TF}_2\text{N}\) and C\(_n\)mim\(\text{BTf}_2\text{N}\). The observed cooling rates are not correlated with macroscopic thermal diffusivity, contrary to what we observed for molecular liquids, suggesting that these ionic liquids are thermally inhomogeneous. Finally, two newly found phenomena are described, which are indicative of extraordinary nanoenvironments in a magnetic ionic liquid, C\(_n\)mim\([\text{FeCl}_4]\). Though information carried by these experimental results is rather indirect, as is always the case with structural studies of liquids, they all point to the same direction of local structure formation in imidazolium-based ionic liquids.

**Low-Frequency Raman Spectra and Collective Cationic Motions in C\(_n\)mim\([PF_6]\)**

In low-frequency Raman spectra of liquids, various collective modes are expected to appear in addition to intramolecular vibrational modes.\(^23,24\) We are able to study those collective modes, which must bear key information for understanding ionic liquid structures, by measuring low-frequency Raman spectra. Though similar information is obtainable with the ultrafast optical Kerr effect (OKE) method\(^25–32\) and also by far-infrared/terahertz spectroscopy,\(^33,34\) Raman spectroscopy is more advantageous for its versatility. It would be difficult to measure by OKE or far-infrared/terahertz spectroscopy both the liquid and solid samples under the same experimental conditions.

A typical Raman spectrum of C\(_4\)mim\([PF_6]\) is shown in Figure 1a. A shoulder-like Raman band is observed in the low-frequency region below 100 cm\(^{-1}\). In order to discuss the low-frequency spectra quantitatively, we reduce the observed spectra by the following formula:\(^35\)

\[
I_{\text{red}}(\nu) = \left(1 - e^{-\frac{\nu}{kT}}\right)(\nu_0 - \nu)^{-3}I_{\text{obs}}(\nu)
\]

(1)

Here \(\nu_0\) is the wavenumber of Stokes Raman shift, \(\nu_0\) is the wavenumber of the incident laser light, \(c\) is the velocity of light, \(k\) is the Boltzmann constant, and \(T\) is the absolute temperature. The resultant reduced spectrum of C\(_4\)mim\([PF_6]\) is shown in Figure 1b. The intensities in the reduced spectrum are proportional to the Raman scattering cross sections of the \(\nu = 1 - \nu = 0\) transitions, where \(\nu\) is the vibrational quantum number. The physical meaning of reducing a Raman spectrum is to cancel out the effect of thermal excitation and make the spectrum comparable to calculated Raman intensities. Note that the negative feature in the anti-Stokes region in the reduced spectrum is the inverted mirror image of the Stokes region.

The reduced Raman spectra of C\(_n\)mim\(X\) ionic liquids are compared in Figure 2. We compare C\(_4\)mim\([\text{BF}_4]\) and C\(_4\)mim\([PF_6]\) for examining the anion dependence and C\(_4\)mim\([PF_6]\), C\(_6\)mim\([PF_6]\), and C\(_8\)mim\([PF_6]\) for investigating the dependence on the alkyl chain length of the cation. For the sake of quantitative discussion, these spectra are fitted by simple model functions assuming two independent components. One is a Gaussian band that is related to an inhomogeneously broadened vibrational band. The other is a relaxation band that corresponds to a Debye-type relaxational motion. The observed spectra are very well fitted with the model functions corresponding to the two components. The best-fitted curves are shown in Figure 2 by the solid lines. The peak position \(\Omega\) and the width \(\Gamma\) of the Gaussian component obtained

\[\text{FIGURE 1.} \ \text{The observed (a) and reduced (b) Raman spectra of C}_4\text{mim \([PF_6]\).}\]

\[\text{FIGURE 2.} \ \text{Reduced Raman spectra for C}_n\text{mim}X \text{ionic liquids: (a) C}_4\text{mim}\text{[BF}_4\text{]; (b) C}_4\text{mim}\text{[PF}_6\text{]; (c) C}_6\text{mim}\text{[PF}_6\text{]; (d) C}_8\text{mim}\text{[PF}_6\text{].} \ \text{The solid lines indicate the fitting results. The vertical bars indicate the calculated results for the C}_4\text{mim cation.}\]
from the best fit are listed in Table 1. The relaxational components have similar widths in all measured ionic liquids. They are around 20 cm$^{-1}$ (corresponding to about 1 ps damping time constant) and correctly reproduce the baselines of the spectra in the 200–600 cm$^{-1}$ region.

The peak position $\Omega$ of the Gaussian component shows no appreciable anion dependence, but it decreases systematically with increasing chain length, $n$, of the cation. It is therefore considered that the Gaussian vibrational bands are due to cationic motions. However, these bands are not assignable to intracationic vibrations, because their area intensities are much higher than those of the other intracationic bands. In fact, a quantum chemical calculation of an isolated C$_4$ mim$^{+}$ cation at the B3LYP/6-31+G** level$^{36}$ predicts only very weak bands around 80 cm$^{-1}$ (vertical bars in Figure 2b). We therefore assign these Gaussian components to inhomogeneously broadened bands due to collective motions of cations involved in local structures. We note that a similar band in an OKE spectrum has been assigned to a librational mode.$^{26}$ The unusually large widths, 35–40 cm$^{-1}$, are consistent with local structures having different size.

In the 200–400 cm$^{-1}$ region, we observe a few bands assignable to intracationic modes. It is known that the accordion vibration of alkyl chain (in-phase C–C–C bend) exists in this frequency region.$^{37}$ According to the quantum chemical calculation, the 330 cm$^{-1}$ band of C$_4$mim[Pf$_6$] is assignable to the accordion vibration from the butyl chain, coupled with the out-of-plane vibration of the imidazolium cation. If the chain length of the cation becomes longer, the frequency of the accordion vibration shifts to lower frequency. Corresponding to this expectation, we find broad features in the 200–300 cm$^{-1}$ region of the spectra of C$_4$mim[Pf$_6$] and C$_6$mim[Pf$_6$]. It seems that the accordion band becomes broader as well with increasing $n$. This observation is consistent with stronger interchain interactions in liquid C$_4$mim[Pf$_6$] with larger $n$. The peak position $\Omega$ of the Gaussian component and the width of the accordion band both change more markedly between $n = 4$ and $n = 6$ than between $n = 6$ and $n = 8$. It seems that the liquid structure changes more on going from C$_4$mim[Pf$_6$] to C$_6$mim[Pf$_6$] than from C$_4$mim[Pf$_6$] to C$_8$mim[Pf$_6$].

### Table 1. The Peak Position $\Omega$ and Width $\Gamma$ of the Gaussian Component in the Low-Frequency Raman Spectrum of C$_n$mimX

<table>
<thead>
<tr>
<th>C$_n$mimX</th>
<th>$\Omega$, cm$^{-1}$</th>
<th>$\Gamma$, cm$^{-1}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>C$_4$mim[BF$_4$]</td>
<td>69 ± 2</td>
<td>40 ± 2</td>
</tr>
<tr>
<td>C$_4$mim[Pf$_6$]</td>
<td>71 ± 2</td>
<td>36 ± 2</td>
</tr>
<tr>
<td>C$_6$mim[Pf$_6$]</td>
<td>64 ± 2</td>
<td>37 ± 2</td>
</tr>
<tr>
<td>C$_8$mim[Pf$_6$]</td>
<td>62 ± 2</td>
<td>37 ± 2</td>
</tr>
</tbody>
</table>

Local Structures in C$_n$mim[Pf$_6$] as Revealed by Spatial Distribution of Coherent Anti-Stokes Raman Scattering (CARS)

It has recently been shown by us that the local structures in liquids and solutions can be sensitively probed by measuring the spatial distribution of coherent anti-Stokes Raman scattering (CARS) signals.$^{38,39}$ In CARS, two laser beams with $\omega_1$ and $\omega_2$ angular frequencies and with $k_1$ and $k_2$ wavenumber vectors, respectively, are incident on a sample medium. If the frequency difference $\omega_1 - \omega_2$ coincides with the frequency $\Omega$ of a Raman active transition ($\omega_1 - \omega_2 = \Omega$, Raman resonance), a coherent CARS signal is emitted in the phase-matched direction determined as $2k_1 - k_2 = k$ (Figure 3). This phase-matching condition is derived on the assumption that the interaction length $L$ of the incident radiations with the medium is much larger than their wavelengths, $\lambda_1$ and $\lambda_2$ ($L \gg \lambda_1$ and $\lambda_2$). In other words, the sample medium must be optically homogeneous (invariant refractive index) over a range much larger than $\lambda_1$ and $\lambda_2$. In reality, however, the sample medium can be optically inhomogeneous (distributed refractive index) for many reasons. If, for example, the sample liquid has mesoscopic local structures, the refractive index of the medium varies microscopically, even though it is transparent and seemingly homogeneous, and the interaction length $L$ is limited due to this optical inhomogeneity. Then, the phase-matching condition is relaxed and a portion of CARS signal is emitted off from the phase-matched direction ($\theta > 0$ in Figure 3). This phase-mismatched CARS signal at $\theta = 90^\circ$ was already measured and studied as partially coherent anti-Stokes Raman scattering (PCARS).$^{40,41}$ The correlation of the PCARS intensities with the optical inhomogeneity in binary liquid mixtures was discussed with reference to Rayleigh scattering intensities.$^{41}$ If the spatial distribution of the phase-mismatched CARS signal is measured as a function of $\theta$, it gives more-detailed information on the optical inhomogeneity in the medium. We recently used polystyrene beads of different size (100 and 350 nm in diameter) dispersed in water as a model system and quantitatively examined the CARS signal spatial distribution.$^{38}$ The CARS signal from the ring breathing vibration of polystyrene was measured as a function of $\theta$. The observed spatial distribution showed clear dependence on the bead size, with larger beads giving less broadened distribution. This size dependence was successfully reproduced by a theoretical simulation assuming a microscopic distribution of refractive index that corresponds to random location of polystyrene spheres in water.$^{38}$ Thus, measurement of the CARS signal spatial distribution has been shown to provide quantitative information on the size or extent of the optical inhomogeneities existing in liquids.

If local structures are formed in ionic liquids, they give rise to optical inhomogeneities, and they can be probed by CARS. Figure 4 shows the measured CARS signal spatial distributions of the totally symmetric PF stretch mode of Pf$_6^-$ in liquid C$_4$mim[Pf$_6$], C$_6$mim[Pf$_6$], and C$_8$mim[Pf$_6$].$^{39}$ The spatial distribution becomes narrower on going from C$_4$mim[Pf$_6$] to C$_8$mim[Pf$_6$], unequivocally...
indicating that these three ionic liquids are optically inhomogeneous at the microscopic level and generate significant phase-mismatched CARS signals. It also indicates that the inhomogeneity is less extensive in C_{4}mim[PF_{6}] than in C_{6}mim[PF_{6}] and C_{8}mim[PF_{6}]. In other words, there exist local structures giving rise to optical inhomogeneity, and these local structures are more extensive in ionic liquids with longer alkyl chains. Since we are looking at the CARS signal from the PF_{6}^{-} anion, only the inhomogeneities in the anion distribution are detected, but they must be accompanied with the inhomogeneities of the cation distribution. More extensive local structures in C_{n}mim[PF_{6}] with larger n are consistent with the results discussed in the preceding section, since stronger interactions in longer alkyl chains are likely to generate local structures with larger size. Furthermore, the trend gives plausible explanation to the fact that the shear viscosity increases rapidly with increasing n: C_{4}mim[PF_{6}], 371 cP; C_{6}mim[PF_{6}], 680 cP; C_{8}mim[PF_{6}], 866 cP. Here again, we see that changes are larger on going from n = 4 to n = 6 than on going from n = 6 to n = 8.

The size of the detected local structures of C_{n}mim[PF_{6}] is not clear yet. It is certain that the size is smaller than the wavelength of visible light, because they are all transparent. Considering the fact that the polystyrene beads with 100 nm radius give similar width of CARS transparent, it is not clear yet. It is certain that the size is smaller than a few tens of nanometers and a few hundred nanometers. However, this estimation needs to be refined by comparison of the experimental spatial distribution with theory, in particular, with molecular dynamics simulations. We note that recent molecular dynamics simulation studies have suggested much smaller size (a few to several nanometers) of local structures.

**Microscopic Energy Transfer and Local Structures in Ionic Liquids Probed with a “Picosecond Raman Thermometer”**

Energy transfer is a fundamental process in the condensed phase, in which a molecule exchanges its energy with the surrounding molecules constantly through intermolecular interactions. Monitoring the energy transfer process therefore gives quantitative information about the intermolecular interactions in the condensed phase. One of the most effective ways for studying energy transfer is to record the cooling kinetics of a probe molecule after instantaneous excitation of its vibrational states. Immediately after the excitation, the probe molecule starts cooling toward the thermal equilibrium by transferring the excess vibrational energy to the surrounding molecules. Because this vibrational cooling proceeds in a few picoseconds to a few hundreds of picoseconds, time-resolved spectroscopy with time resolution of a few picoseconds is required. More than a decade ago, we constructed a transform-limited picosecond time-resolved Raman spectrometer and started studying picosecond vibrational dynamics of molecules in liquids and solutions. In the course of these studies, we found that the C=C stretch Raman band of the first excited singlet (S_{1}) state of trans-stilbene (tSB) showed linear dependence on temperature and that it served as an excellent “picosecond Raman thermometer”. Recently, we have examined the cooling kinetics of S_{1} tSB in ionic liquids by using this picosecond Raman thermometer.

We measure picosecond time-resolved Raman spectra with the pump–probe method. The stilbene molecule is photoexcited to the S_{1} state with the pump pulse at 296 nm. The produced S_{1} state is monitored with the probe pulse at 592 nm, which is in resonance with the S_{n}\rightarrow S_{1} absorption. With this setting of the pumping wavelength, an excess energy of 3000 cm\(^{-1}\) is deposited to S_{1} tSB. Time resolution of the spectrometer, estimated by the cross-correlation function between the pump and probe pulses, is 1.7 ps.

A set of time-resolved Raman spectra of S_{1} tSB measured in a C_{n}mimTf_{2}N solution (5.0 \times 10^{-3} mol dm^{-3}) is shown in Figure 5. The Raman bands from the solvent C_{n}mimTf_{2}N have been subtracted in the spectra. It is clear from the figure that the C=C stretch band near 1570 cm\(^{-1}\) shifts toward higher wavenumbers with increasing time delay. This shift reflects the temperature decrease of the S_{1} tSB molecule during the cooling process. The cooling curve of S_{1} tSB, obtained by plotting the peak position of the 1570 cm\(^{-1}\) band against the time delay, is fitted well with a single exponential function. The obtained rate constants are summarized in Table 2.
We previously measured similar cooling kinetics of S1\textsubscript{t}st in ten molecular liquids.\textsuperscript{49} The obtained rate constants showed a good correlation with the thermal diffusivity $\kappa$ of the solvent. Thermal diffusivity characterizes the macroscopic heat conduction in the solvent, which is well represented by the diffusion equation of heat:

$$\frac{\partial U}{\partial t} = \kappa \Delta U \tag{2}$$

Here, $U$ is temperature and $t$ is time. The diffusion equation of heat is directly derived from the continuity equation and Fourier's law of heat conduction. Figure 6 shows the plot of the observed cooling rate of S1\textsubscript{t}st vs thermal diffusivity in ten molecular liquids and two ionic liquids. A clear relation between the cooling rate and the thermal diffusivity is observed for the molecular liquids. We have explained this observation by assuming the following scheme of thermal energy transfer. The excess energy first given to S1\textsubscript{t}st is shared in the first solvation shell within the time resolution of the experiment (a few picoseconds). The solvent–solvent energy transfer then follows. The whole cooling rate is controlled by the heat transfer process in the solvent, which is well characterized by thermal diffusivity. The cooling rates observed in the ionic liquids are, however, not explainable by the simple model that we proposed for molecular liquids. The reported values of thermal diffusivity\textsuperscript{49} for C\textsubscript{2}mimTF\textsubscript{2}N and C\textsubscript{4}mimTF\textsubscript{2}N, $(6.01 \pm 0.04) \times 10^{-8}$ m\textsuperscript{2} s\textsuperscript{-1} and $(6.0 \pm 0.1) \times 10^{-8}$ m\textsuperscript{2} s\textsuperscript{-1}, respectively, are much smaller than the values for heptane $(8.01 \times 10^{-8}$ m\textsuperscript{2} s\textsuperscript{-1}) or ethanol $(8.70 \times 10^{-8}$ m\textsuperscript{2} s\textsuperscript{-1}). Contrary to what is expected from these values of thermal diffusivity, the cooling rates observed in the ionic liquids are similar to that in ethanol and are larger than that in heptane (Figure 6). It seems that, for ionic liquids, thermal diffusivity does not represent the energy transfer process at the microscopic level.

The cooling rate of a solute may not show a good correlation with macroscopic thermal diffusivity of the solvent if there are local structures formed in the solvent, as we expect for C\textsubscript{2}mimTF\textsubscript{2}N and C\textsubscript{4}mimTF\textsubscript{2}N. The pico-second cooling process is determined by the energy transfer within a group of solvents that are surrounding the solute. By the time the flow of the excess energy of 3000 cm\textsuperscript{-1} reaches the boundary of the local structure, it is already distributed to the many degrees of freedom of a large number of solvent molecules. The temperature may well be cooled back to the room temperature at that time. The whole cooling process is thus completed within the local structure. The macroscopic heat conduction, however, proceeds over the boundary between the local structures. If the energy transfer between a pair of local structures is less effective than the energy transfer within the local structures, then macroscopic heat conduction is controlled by the inter-local-structure energy transfer as shown in Figure 7. We consider that the apparent discrepancy between the thermal diffusivity dependence of the cooling rate in the molecular liquids and that in the ionic liquids indicates the presence of local structures in C\textsubscript{2}mimTF\textsubscript{2}N and C\textsubscript{4}mimTF\textsubscript{2}N.

<table>
<thead>
<tr>
<th>solvent</th>
<th>$k_{\text{vib}}$, ps\textsuperscript{-1}</th>
<th>$\kappa$, $10^{-8}$ m\textsuperscript{2} s\textsuperscript{-1}</th>
</tr>
</thead>
<tbody>
<tr>
<td>C\textsubscript{2}mimTF\textsubscript{2}N</td>
<td>0.11 ± 0.02</td>
<td>6.01 ± 0.04*</td>
</tr>
<tr>
<td>C\textsubscript{4}mimTF\textsubscript{2}N</td>
<td>0.12 ± 0.02</td>
<td>6.0 ± 0.1*</td>
</tr>
<tr>
<td>heptane</td>
<td>0.08*</td>
<td>8.01*</td>
</tr>
<tr>
<td>ethanol</td>
<td>0.13*</td>
<td>8.70*</td>
</tr>
</tbody>
</table>

\textsuperscript{*} Reference 45. \textsuperscript{b} Reference 47.
Extraordinary Nanoenvironments in a Magnetic Ionic Liquid, C₄mim[FeCl₄]

Local structures in imidazolium-based ionic liquids may well produce extraordinary molecular environments and hence lead to novel molecular phenomena that have not been found for molecular liquids. Here, we describe two newly found phenomena that are ascribable to such molecular environments in C₄mim[FeCl₄]. C₄mim[FeCl₄] is a prototype magnetic ionic liquid that shows strong response to a magnet. Raman spectroscopy has indicated that the cation structure in C₄mim[FeCl₄] is very similar to those in the other C₄mim-based ionic liquids discussed in the preceding sections.

In the course of Raman spectroscopic examination of liquid C₄mim[FeCl₄] under a microscope, we accidentally found that unknown crystals occasionally formed when the liquid was irradiated by the back illuminating light. It was later confirmed that the crystal formation was accelerated by ultraviolet irradiation at 365 nm. We suspect that a photochemical reaction of the FeCl₄⁻ anion results in this crystal formation (see discussion below).

The Raman spectrum of one of those crystals is given in Figure 8a. As shown in the inset of the figure, the formed crystal has sharp edges characteristic of a single crystal. The Raman spectrum shows two prominent bands at 3401 and 1619 cm⁻¹ and two weaker bands in the low-frequency region. This spectral pattern is quite different from those of C₄mim[FeCl₄]; neither the C₄mim⁺ nor the FeCl₄⁻ bands are observed. The frequencies 3401 and 1619 cm⁻¹ happen to be close to those of water. We therefore repeated the same experiment under a D₂O atmosphere with an expectation that water impurity absorbed from the atmosphere might play a role. The resultant Raman spectrum is shown in Figure 8b. A crystal similar to that in spectrum a was obtained. In spectrum b, in addition to the two peaks observed also in spectrum a, four peaks are observed at 2541, 2503, 1427, and 1192 cm⁻¹. These spectral changes are consistent with the assignments of the two prominent bands in spectrum a to water, 3401 cm⁻¹ (symmetric OH stretch) and 1619 cm⁻¹ (HOH bend). The new bands observed in spectrum b are assigned accordingly: 2541 cm⁻¹ (OD stretch of DHO), 2503 cm⁻¹ (symmetric OD stretch of D₂O), 1427 cm⁻¹ (HOD bend), and 1192 cm⁻¹ (DOD bend). The spectral change in the low frequency region can be ascribed to lattice vibrations including water molecules.

It is obvious that water molecules are contained with an extremely high concentration in the crystal formed in C₄mim[FeCl₄]. It is also clear that all the observed water bands show much smaller widths than those in ice and liquid water. In particular, the full width at half-maximum (fwhm) of the HOH bend band in the spectrum a is less than 9 cm⁻¹. As far as the authors are aware, such a sharp HOH band of water has never been reported. These unusually sharp bands are indicative of unusual structure of water molecules in the crystal. If the crystal contains water molecules only, it is a new form of ice at room temperature. We note this tempting possibility. We also note the possibility that some unknown species, which is silent in Raman spectroscopy, is contained in the crystal. For example, a clathrate hydrate structure, (H₂O)ₙ, is conceivable. Under UV irradiation, the FeCl₄⁻ anion can be photohydrated to FeCl₃(OH)⁻ and generate H⁺ and Cl⁻. Then, the H⁺ and Cl⁻ ions are likely to be present in the irradiated ionic liquid and play a role in the crystal formation.

Separation of the crystals formed in C₄mim[FeCl₄] has so far been unsuccessful, hindering its X-ray structural analysis. Whatever the chemical composition might be, the uniqueness of the crystal must be a reflection of extraordinary molecular environments specific to C₄mim[FeCl₄]. The relevance of magnetic interactions with this crystal formation is not clear yet.

The other new phenomenon is the magnetic-field-dependent fabrication of nanostructures of conducting polymers in C₄mim[FeCl₄]. We found that nanoparticles and rod-like structures of conducting polymers (polypyrrole, N-methyl-polypyrrole, polythiophene) formed by simply adding monomer liquids to C₄mim[FeCl₄]. It was also found that the formed nanostructures were dependent on the external magnetic field. Figure 9 shows the SEM (scanning electron microscope) images of polypyrrole synthesized by this method with and without an external magnetic field. Without an external magnetic field (Figure 9a), nanoscale particles with 60 ± 20 nm diameter were formed (see also inset a-1). With an external magnetic field (Figure 9b), the particles tend to be aligned in a particular direction and a rod-like structure is formed. It is highly

FIGURE 9. SEM images of the nanostructures of polypyrrole formed in C₄mim[FeCl₄] without (a) and with (b) an external magnetic field (1.3 T).
likely that local structures in C₄mim[FeCl₄] give rise to specific nanoenvironments that cause the formation of nanostructures and that these local structures and hence the nanoenvironments can be regulated by an external magnetic field. Nanostructure formation in nonmagnetic room-temperature ionic liquids has been reported.\textsuperscript{58,59}

### Conclusion

It seems beyond doubt that specific local structures are extremely complicated with a large number of shallow local minima.\textsuperscript{1} An ionic liquid system may have an unusually long thermal relaxation time on account of this complicated energy landscape. We cannot rule out the possibility that it is even longer than our lifetime! New physicochemical approaches and standpoints, both experimental and theoretical, must be taken in order to correctly address this problem. Elucidation of local structures in ionic liquids may alter the concept of thermal equilibrium and the definition of material phase based on it.

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**Note Added after ASAP Publication:** This manuscript was released ASAP on October 27, 2007 with an incorrect version of Figure 4. The correct version was posted on November 20, 2007.

### References


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