# Investigating Nanostructuring within Imidazolium Ionic Liquids: A Thermodynamic Study Using Photochromic Molecular Probes

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Molecular photoswitches have been used to investigate the possibility of nanostructured polar and nonpolar domains in ionic liquids (ILs). Two photochromic compounds, spiropyran (BSP) and spirooxazine (SO) were added to imidazolium based ionic liquids containing the anion [NTf2], and their photochromic behavior was monitored with increasing side chain length ( $C_2-C_{12}$ ) of the imidazolium cation. Increasing side chain length was found to have only minor effects on the rate of thermal relaxation of the merocyanine form of spiropyran (MC<sub>BSP</sub>) and spirooxazine (MC<sub>SO</sub>) to BSP and SO, respectively. BSP was found to be a suitable optical probe, as linear correlations in parameters were observed for this compound. This is believed to be because BSP-IL interactions are based on hydrogen bonding between the MC<sub>BSP</sub> and the ionic liquid cations, compared to  $MC_{SO}$ , which is limited to electrostatic interactions. Hence, the sensitivity of  $MC_{BSP}$  is enhanced in the charged polar regions of the IL. Increasing the side chain of the cation results in slight increases in MC<sub>BSP</sub> to BSP relaxation activation energy from 96.93 kJ⋅mol<sup>-1</sup> in [C<sub>4</sub>mIm][NTf<sub>2</sub>] to 105.27 kJ⋅mol<sup>-1</sup> in  $[C_{12}\text{mIm}][NTf_2]$ .  $MC_{BSP}$  to BSP relaxation  $\Delta S^{\dagger}$  and  $\Delta H^{\dagger}$  values also increase with increasing side chain length. The ability for spirocyclic compounds to switch between polar and nonpolar forms appears to allow polar and nonpolar regions in ILs to be probed dynamically using a single probe dye. It appears that the value of the ground state equilibrium constant,  $K_e$ , is dominated by the nonpolar regions of the IL while the equilibrium constant of activation,  $K^{\ddagger}$ , is dominated by the polar regions. A correlation of side chain length to equilibrium constant of activation is believed to be because polar regions are possibly expanding due to increasing influence of nonpolar side chain interactions and compound insertion upon the overall solvent structure. The result of such reordering and dispersion of polar regions reduces solvent-solute interactions which increases the rate of  $MC_{BSP}$  to BSP relaxation.

#### Introduction

Ionic liquids (ILs) are receiving increasing attention due to the large range of ions available, adding a "designer" aspect to the liquids, meaning that such liquids could provide solvents with specifically tailored properties. ILs consist entirely of ions in the liquid state under 100 °C. Their proposed applications include recyclable solvents¹ and replacements to molecular solvents in catalysis,² electrochemistry,³ synthesis,⁴ and elemental analysis.⁵ Several reviews have provided detailed insight into ILs and have promoted their implementation as common laboratory solvents.⁶ However, the lack of detailed knowledge of the physicochemical properties has resulted in their full potential not being completely realized.

Estimates of IL physicochemical properties has led to the re-evaluation of solvent polarity as a macroscopic property that assumes the entire solvent system acts as a homogeneous medium rather than a medium that may possess nanostructured regions with dramatically different characteristics. Reichardt's dye, can provide information about the polarity of a conventional solvent because of the absorption shift which accompanies the interaction of solvent molecules

with the zwitterionic sites of the dye, as these interactions affect the excitation band gap of the probe molecule. To compliment such single parameter probe studies, the combination of three dyes (including Reichardt's) by Kamlet and Taft enabled hydrogen bonding behavior of solvents to be examined. 9-11 Both single and multiparameter probe studies have seen considerable usage in evaluating ILs in an attempt to understand trends in their physicochemical properties, but it appears ILs are much more complex solvent systems capable of undergoing many types of interactions. Characterizing them with a single "polarity" term fails to describe the type and magnitude of individual interactions that make each IL unique. 4,12-14

Photochromic materials based on spirocyclic compounds form merocyanines (**MC**) when exposed to UV irradiation; these zwitterions are similar in structure to those of betaine's used in Reichardt's studies. It is proposed that such compounds could possibly act as multiparameter probes based on solvatochromic shifts of the **MC**, rates of thermal relaxations of **MC** to the spiro form, and equilibrium constants  $K_e$ . Photoswitching of spirocyclic compounds (Figure 1) involves the use of light to induce cleavage of the sp<sub>3</sub> hybridized carbon of the indoline ring known as  $C_{\rm spiro}$ . Initially, both **BSP** and **SO** are uncharged molecules with the indoline fragment (**A**) orthogonal to that of its respective pyran or oxazine fragment (**B**). Cleavage of this bond can occur by irradiation with UV light (typically around 365–375 nm) to form the merocyanine (**MC**) isomer.

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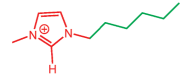
Figure 1. (a) Basic structure of spirocyclic compounds. Indoline (A) and pyran/oxazine (B) ring, where X = C (pyran), N (oxazine). (b) Spirocyclic compounds used in our studies and their photoswitchable states.

The MC isomer arises from cis—trans isomerization of the molecule, which results in a planar molecule with the charges largely delocalized across the molecule. The reorientation of the molecule results in the formation of several isomers; however, it is the cis—trans—cis (CTC) form which is most stable and produces the longest lifetime for solvent—solute interaction and thus use as a photochromic probe. The process is completely reversible by irradiation with white light. These photochromic compounds also revert spontaneously to their more stable "ring closed" isomer through thermal reversion, in which the MC uses heat from its surroundings to supply sufficient energy for the reorientation to occur. This process follows first order kinetics.

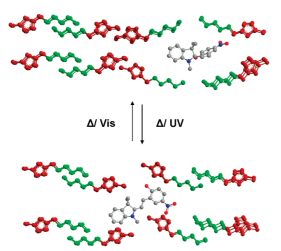
The MC isomer is also sensitive to its immediate molecular environment, and these corresponding specific and nonspecific interactions that occur mediate the compound's equilibrium between both isomers. 15-18 In molecular solvents, the polaritykinetic relationship was upheld for both BSP and SO with increasing rate constants observed with decreasing polarity. 19,20 Kamlet-Taft parameters determined that this is due to decreasing levels of hydrogen bonding between solvent and spirocyclic compounds.<sup>21</sup> MC<sub>BSP</sub> is known to form hydrogen bonds, therefore inhibiting its rate of thermal relaxation. Similar trends to that observed in molecular solvents implies that BSP can act as a solvent sensitive probe in ionic liquids. We recently reported the photo- and solvatochromic properties of BSP in ILs containing [NTf<sub>2</sub>]<sup>-</sup>. It was found that the kinetics and thermodynamics of the BSP-MC<sub>BSP</sub> equilibrium was sensitive to the nature of the cation. It was also observed that the cation,

[emim] $^+$ , can even form a through-space orbital interaction with the merocyanine isomer, rather than a simple electrostatic interaction, thus inhibiting the merocyanine conversion back to the BSP isomer. We also found that the MC isomer's spectroscopic properties could be significantly altered by varying the anion in phosphonium based ILs. $^{22}$  The same relationship was investigated in ILs using SO, but similar trends were not apparent. It is believed that the bulky nature of the MCsO and its substituents restricted interactions with the IL ions to electrostatic/nonspecific interactions. SO was therefore chosen to act as a reference compound for comparing solvent structure, as the relatively weak interactions mean that its process of thermal relaxation would be somewhat independent of the solvent system used.

Recent studies have proposed that imidazolium based ILs may form ordered liquid systems resembling crystalline environments based on stacking of mutual charges (aggregation) or ordered association of cation to surrounding anions and vice versa.  $^{23,24}$  Lopes et al. examined the formation of nanostructured domains in ILs containing the imidazolium cation; it was found that certain imidazolium cations in ILs can be divided into two specific regions: a polar headgroup where the ion charge resides and a nonpolar region where side groups extend into space (Figure 2). These polar head groups appear to interact preferentially with one another to form aggregates by three dimentional  $\pi$ -stacking and mutual association of the charged imidazolium rings with anion species to form polar regions. Alkyl side chains extend away from these regions and through



**Figure 2.** Imidazolium cation C<sub>6</sub>mIm showing polar (red) and nonpolar (green) regions of the molecule based on findings and convention of identification by Lopes et al.



**Figure 3.** Schematic of proposed 3D ordering of imidazolium cations with **BSP** probe in  $[C_6mIm][NTf_2]$  showing preferred residence of the MC (lower) and SP (upper) forms of the probe in polar and nonpolar regions, respectively.

side chain/van der Waals interactions form a complex network of nonpolar regions.

The MC isomer of the two probe molecules provides a highly polar zwitterionic system which can interact with the opposing charges of the polar head groups of the solvent system, whereas the spiro isomer is known to be a neutral nonpolar molecule and should reside in the nonpolar region of the IL or at least at the interface of the region interfaces due to a marked reduction in solvent-solute interactions. As thermal relaxation occurs, the molecules would be expected to alternate their interactions between primarily polar and nonpolar regions as the charged MC isomer reverts to the uncharged SP form and solvent reorganization occurs. Previous spectroscopic studies of spiropyran in solutions containing lipid bilayers have exhibited diffusional behavior believed to be due to the differing associations of each form of the compound.<sup>26</sup> The similarity between this ordering and the proposed ordering of ILs infers the possibility that similar mechanisms may be present with ILs which would allow the photochromic compounds to preferentially interact with specific regions, thus probing the existence of such structuring. The spirocyclic compounds themselves are bulky molecules which are quite large when compared to the size of the solvent molecules. The introduction of such molecules would therefore be expected to somewhat disrupt the ordering of the IL structure. However, the resulting reordering of the solvent molecules around the probe and the thermodynamic stability of the solvent molecules to physically accommodate the probe molecules will provide insight into the solvent system and the extent to which solvent/intramolecular ordering occurs. Increasing the length of the nonpolar side chain of the imidazolium ion (Figure 4) should affect the ordering of the IL at the molecular level due to the increased volume of nonpolar regions, and related increased dispersion and corresponding expansion of polar regions arising from dissociation of imidazolium head groups from one another. Since the MC isomer

R = ethyl: C<sub>2</sub>mIm butyl: C<sub>4</sub>mIm hexyl: C<sub>6</sub>mIm octyl: C<sub>8</sub>mIm decyl: C<sub>10</sub>mIm dodecyl: C<sub>12</sub>mIm

**Figure 4.** Cations and anion used in this study: 1-alkyl-3-methylimidazolium  $[C_m \text{mIm}]^+$  and bis(trifluoromethanesulfonyl) amide  $[\text{NTf}_2]^-$ .

preferentially resides in polar regions, it is anticipated that increasing the surrounding nonpolar regions could possibly influence the equilibrium by favoring the **SP** form of the molecule. If structured polar domains exist, then their stabilizing influence on **MC** should reduce the effect of these nonpolar regions and subsequently provide rates of thermal relaxation much slower than that expected for long-chain, nonpolar molecules. If nanostructuring exists in ILs, we should observe a nonlinear relationship between  $K_e$  and thermal relaxation of the MC, due to the different molecular environments observed.

Thermodynamic and kinetic studies were carried out to examine the extent of structuring in imidazolium based ILs based on equilibrium effects, rates of thermal relaxation, and the effect upon solvent ordering due to the introduction of a bulky probe molecule.

#### **Experimental Section**

ILs were synthesized and purified in-house with salts obtained from Sigma-Aldrich using previously reported techniques.<sup>27</sup> The ILs produced were stored under argon to exclude uptake of water. Spectrometric studies were carried out using a Perkin-Elmer Lambda 900 spectrometer (Foss, Ireland) with a Perkin-Elmer PTP-1 temperature controller. Samples were irradiated with UV light at 375 nm using an in-house fabricated array based on 375 nm UV LEDs (Roithner Lasertechnik, Vienna, Austria). Reichardt's dye 30 (Sigma-Aldrich chemicals) and 6-nitro-1',3',3'-trimethylspiro[2H-1-benzopyran-2,2'-indolin] 1',3'-dihdro-1',3',3'-trimethyl-6-nitrospiro (BSP) (Sigma-Aldrich chemicals) were used as purchased with no further purification. 1,3,3-Trimethyl-5'-(2-benzothiazolyl)-spiroindoline-2,3'-naphtho(2,1-b)(1,4) oxazine (**SO**) was previously synthesized and supplied by Dr. Minkovska. All samples were prepared at room temperature.

### **Results and Discussion**

Polarity and Solvatochromic Effects. All ILs were examined using the ET<sub>30</sub> scale as described previously.<sup>21</sup> Spectroscopic shifts relating to solvent-dye interactions were found to have a linear response in ET<sub>30</sub> with increasing side chain length. The linear decrease in polarity was however minor when compared to that of molecular solvents of similar chain lengths. ET<sub>30</sub> values of 52.6 kcal·mol<sup>-1</sup> for [C<sub>2</sub>mIm][NTf<sub>2</sub>] and 53.2 kcal⋅mol<sup>-1</sup> for [C<sub>6</sub>mIm][NTf<sub>2</sub>] were observed, while similar chain lengths in molecular solvents with a polar region and extending nonpolar side chain (e.g., alcohols) had ET<sub>30</sub> values of 51.9 and 48.8 kcal·mol<sup>-1</sup> for ethanol (C<sub>2</sub>) and 1-hexanol (C<sub>6</sub>), respectively. As with ET<sub>30</sub> values, the observed solvatochromic shifts were subtle compared to those observed in molecular solvents. Such subtleties may be due to the fact that the polar regions of the ILs may somewhat retain their stabilizing influence upon the MC even with increasing chain length, thus

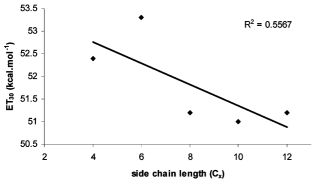


Figure 5. ET<sub>30</sub> values of ILs with increasing cation side chain length.

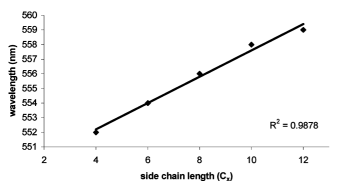


Figure 6. MC<sub>BSP</sub> solvatochromic shift versus cation side chain length.

reducing the overall solvatochromic effect of the increasing alkyl chain length of the solvent. In addition to the proposed reduction of the polar region influence, the increasing chain length size also results in increasing size of nonpolar domains and corresponding separation of polar regions from one another. Such increases would therefore produce a liquid system that would appear more nonpolar throughout to spirocyclic compounds and thus result in solvatochromic shifts corresponding to such environments.

**BSP**, due to its nitro group, exhibits solvatochromic effects due to a wavelength shift of the MC<sub>BSP</sub> absorption maximum. Linear bathochromic (red) shifts were observed with increasing cation side chain lengths as previously established in molecular solvents of decreasing polarity.<sup>28</sup> Similar effects were not observed for SO, as the equivalent MC<sub>SO</sub> does not exhibit solvatochromic behavior.

**Kinetic Parameters.** Samples were placed in a thermostatically controlled UV—vis spectrometer and were irradiated in situ using an LED array to induce ring-opening and MC formation. Upon removal of the UV light source, the thermal relaxation first order decay curves were then examined using eq 1.

$$\ln\frac{[A]}{[A_0]} = -kt \tag{1}$$

The rates of thermal relaxation were recorded at 298 K and summarized in Table 1. Slight increases in rates of relaxation were observed with increasing chain length when **BSP** was added to the ILs. Increasing chain length resulted in rates of thermal relaxation doubling from  $1.0 \times 10^{-3}$  to  $2.0 \times 10^{-3}$  s<sup>-1</sup> (Figure 7a). Since lengthening of the side chains would be expected to promote more interchain interactions, it is believed that such interactions place a strain upon the mutual interactions of the polar head groups. This strain in turn may cause the polar

head groups to separate further from each other, increasing the size of the polar region while reducing the overall charge density within the region to accommodate the solute molecules.<sup>29</sup> Early investigations into IL structuring observed deviations from traditional correlations between viscosity and diffusion for the diffusion coefficients of solutes within "wet" or hydrated ILs.<sup>30</sup> Further studies concluded that this could be explained through the IL system swelling to accommodate the solvent molecules, as had been observed previously in studies involving IL/ molecular solvent binary mixtures, such as [bmIm][PF<sub>6</sub>] and naphthalene.31 We propose that the addition of spirocyclic compounds, particularly those when predominantly in the charged/polar MC form, are integrated into the IL structure in a similar manner in addition to the size of the **BSP** physically disrupting the regions when added to the ILs. If separation of head groups occurs upon such integration, then the distance between the imidazolium ring charge and the C2 proton of the MC isomer would be lengthened. This would imply a reduction in the ability of the MC molecules to stabilize polar guests. Rates of thermal relaxation of MCBSP were found to increase with increasing chain length, and we believe this is due to a reduction in the effectiveness of interactions in polar regions, while the impact of nonpolar regions is enhanced, favoring the conversion to BSP.

SO was found to have similar relaxation rates of approximately  $2.3 \times 10^{-2} \, \text{s}^{-1}$  within all ILs (Figure 7b), with the increasing chain length having no apparent effect. This agreed with our previous findings and is believed to be due to the solvent-solute interactions being predominantly electrostatic, and its influence was therefore similar in each IL.21 Further support for this hypothesis comes from the rates of thermal relaxation, which were 10 times faster than that observed for the **BSP** system. This would be expected, due to considerably weaker interactive forces stabilizing the  $MC_{SO}$ . The relatively weak electrostatic forces also mean that SO/MCso molecules are relatively free to move within the IL due to the absence of any strong direct interactions in polar or nonpolar nanoregions in the IL. SO molecules may therefore migrate to an intermediate region between the two regions and where variations in side chain length and headgroup interaction would have less impact.

Thermodynamic Parameters. The dependence of the rate of thermal relaxation with temperature was investigated using eqs 2 and 3 to find the activation energy  $(E_a)$ , entropy of activation  $(\Delta S^{\ddagger})$ , enthalpy of activation  $(\Delta H^{\ddagger})$ , and Gibbs energy of activation  $(\Delta G^{\ddagger})$ . An alternative form of the Eyring equation (eq 4) was also employed to derive the equilibrium of the activated complex of the transition state theory. The thermodynamic parameters found are summarized in Table 1.

$$ln k = E_a/RT + ln A$$
(2)

$$\ln(k/T) = -\Delta H^{\dagger}/RT + \ln(k_{\rm B}/h) + \Delta S^{\dagger}/R$$
 (3)

$$k = (k_{\rm B}T/h)K^{\ddagger} \tag{4}$$

where R = gas constant, h = Planck's constant,  $k_b = \text{Boltzmann}$  constant, and  $\Delta S^{\ddagger}$  values were positive and increased with chain length (Table 1). Changes in this parameter were obvious for **BSP** with entropies from 13.79  $J \cdot K^{-1} \cdot \text{mol}^{-1}$  in  $[C_4 \text{mIm}][NTf_2]$  to 46.15  $J \cdot K^{-1} \cdot \text{mol}^{-1}$  in  $[C_{12} \text{mIm}][NTf_2]$ . Entropy of activation is a measure of the amount of reorientation of the system during the thermal relaxation from **MC** to **BSP** within the IL, and it is

TABLE 1: Physicochemical Properties of Spirocyclic Compounds in Molecular Solvents and Ionic Liquids (Reference Values in Parentheses)

SO										
					Ar	rhenius	Eyring			
IL	λmax MC (nm)	$(\times 10^{-2} \text{ s}^{-1})$	$ET_{30} \\ (kcal \cdot mol^{-1})$	$K_{\rm e} \times 10^{-3}$	$\frac{E_{a}}{(kJ \cdot mol^{-1})}$	A	$\frac{\Delta S^{\ddagger}}{(J \cdot K^{-1} \cdot \text{mol}^{-1})}$	$\frac{\Delta H^{\ddagger}}{(kJ \cdot mol^{-1})}$	$\Delta G^{\dagger}_{25} $ (kJ·mol <sup>-1</sup> )	$K^{\ddagger}$ (×10 <sup>-15</sup> )
ethanol <sup>21</sup>	642	(3.8)	(52.1)		(89.89)	$(2.037 \times 10^{13})$	(6.4)	(87.45)	(81.05)	(5.23)
acetone <sup>21</sup>	642	(5.1)	(42.5)		(69.51)	$(4.929 \times 10^{10})$	(-44.73)	(67.08)	(80.41)	(8.83)
C <sub>2</sub> mIm NTf <sub>2</sub>	642	2.3	52.6 (52.6) <sup>35</sup>	6.62	95.23	$1.11 \times 10^{15}$	34.92	92.8	82.39	3.68
C <sub>4</sub> mIm NTf <sub>2</sub>	642	2.5	$52.4 (50.0)^{36}$	4.38	95.18	$1.14 \times 10^{15}$	35.2	92.75	82.26	3.95
C <sub>6</sub> mIm NTf <sub>2</sub>	644	2.2	53.3 (51.9) <sup>35</sup>	3.84	92.35	$3.16 \times 10^{14}$	24.52	89.91	82.61	3.51
C <sub>8</sub> mIm NTf <sub>2</sub>	646	2.5	51.2 (51.1) <sup>14</sup>	4.97	90.79	$1.91 \times 10^{14}$	20.31	88.35	82.3	3.82
C <sub>10</sub> mIm NTf <sub>2</sub>	644	2.3	51.6 (51.0) <sup>36</sup>	3.77	92.87	$4.39 \times 10^{14}$	27.23	90.44	82.33	3.68
C <sub>12</sub> mIm NTf <sub>2</sub>	646	2.6	51.2	5.03	95.32	$1.34 \times 10^{15}$	36.48	92.89	82.02	4.2
					BSP					

					Arrh	enius	Eyring			
IL	λmax MC (nm)	$(\times 10^{-3} \text{ s}^{-1})$	$ET_{30}$ (kcal·mol <sup>-1</sup> )	$K_{\rm e} \times 10^{-3}$	$\frac{E_{\rm a}}{(\rm kJ \cdot mol^{-1})}$	A	$\frac{\Delta S^{\ddagger}}{(J \cdot K^{-1} \cdot \text{mol}^{-1})}$	$\Delta H^{\ddagger} (kJ \cdot mol^{-1})$	$\Delta G^{\dagger}_{25} \atop (\text{kJ} \cdot \text{mol}^{-1})$	$K^{\ddagger}$ (×10 <sup>-16</sup> )
ethanol	539	0.5	$(52.1)^{21}$		102.01	$4.71 \times 10^{14}$	27.84	99.57	91.27	0.87
acetone	560	5.3	$(42.5)^{21}$		104.50	$1.26 \times 10^{16}$	55.18	102.07	85.63	8.62
C <sub>2</sub> mIm NTf <sub>2</sub>	552	1.2	52.6 (52.6) <sup>35</sup>	8.66	106.69	$6.43 \times 10^{15}$	49.55	104.26	89.49	2.02
C <sub>4</sub> mIm NTf <sub>2</sub>	552	1	52.4 (51.6) <sup>36</sup>	7.02	96.93	$8.71 \times 10^{13}$	13.79	94.49	90.38	1.65
C <sub>6</sub> mIm NTf <sub>2</sub>	554	1	53.3 (51.9) <sup>35</sup>	11.6	98.84	$2.67 \times 10^{14}$	21.48	96.41	90.01	1.65
C <sub>8</sub> mIm NTf <sub>2</sub>	556	1.7	51.2 (51.1)14	5.31	103.34	$2.18 \times 10^{15}$	40.55	100.9	88.82	2.71
C <sub>10</sub> mIm NTf <sub>2</sub>	558	1.8	51.6 (51.0) <sup>36</sup>	7.94	106.68	$1.02 \times 10^{16}$	52.42	104.24	88.62	2.96
$C_{12}mIm\ NTf_2$	559	2	51.2	7.63	105.27	$5.73 \times 10^{15}$	46.15	102.14	88.39	3.22

related to the rigidity of the solvent and the overall thermal stability of the MC<sub>BSP</sub>-BSP system. Positive values imply that the IL-solute system undergoes significant reorientation during thermal relaxation from MC<sub>BSP</sub> to BSP. The observed increase in values may be due to increasing ordering of nonpolar regions, as increasing chain length will strengthen interactions in the nonpolar regions. Steric effects from solvent-solvent and solvent-MC<sub>BSP</sub> interactions could result in expansion of polar regions, allowing an easier transfer of the probe molecule during MC<sub>BSP</sub>-BSP conversion. At short chain lengths (C<sub>2</sub>mIm), it would be expected that little ordering exists and the IL ions are in random motion much like molecular solvents. Since ordering is enhanced as chain lengths increase, the IL may adopt a more structured form at the nanoscale, with polar and nonpolar regions between which the MCBSP and BSP molecules can move dynamically as the equilibrium shifts. The introduction of the spirocyclic compound and its subsequent transition from MC<sub>BSP</sub> (charged) to BSP (uncharged) should also result in considerable reorientation of the IL ions around the guest molecules. Furthermore, the physical size of the probe molecule itself is

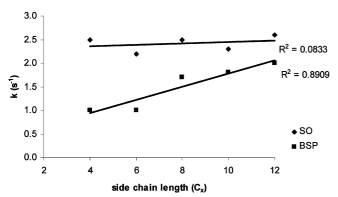
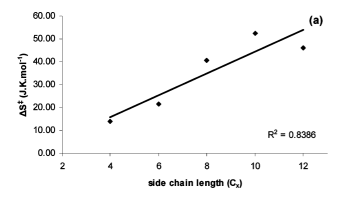


Figure 7. Rate of thermal relaxation of MC isomer to BSP/SO versus cation side chain length.

expected to significantly disrupt the established IL structure. Movement of the probe molecules during their transition from the MC form to their respective BSP/SO forms should also involve molecular reorientation to accommodate the probe molecules, thus increasing the entropy of the system. Positive entropies also imply that the interactions between the IL ions are weak enough for the spirocyclic compound to disrupt the system. This means that the IL structure itself is "fluid" enough to facilitate dynamic movement of the probe molecules between the IL polar/nonpolar regions. SO once again showed no clear correlation between the entropies of activation and chain length, which suggests that for the MCso-SO thermal relaxation in the various ILs is relatively independent of the nanostructure of the IL.

The related parameters of activation energies  $(E_a)$  and enthalpies of activation ( $\Delta H^{\dagger}$ ) were also found to slightly increase with increasing chain length. Initially, this appears to be contradictory, as increasing energy barriers would be expected to yield slower rates of relaxation. However, this could arise due to compensation for the increased separation of charged sites as the chain length is increased. Such changes in the solvent system would be expected to produce an environment that is generally less polar than that of shorter chain cations, and therefore less favorable to MC stability. Since imidazolium cations are known to interact through charge association (coordinated by anions in the polar region), ion pairing, hydrogen bonding, and, to a lesser extent, pi-stacking, it would be expected that *intra*molecular interactions would dominate. <sup>23,33</sup> The expansion of such systems therefore reduces these interactions, and may in fact facilitate intermolecular interactions (MC-IL). Such interactions would result in an increased stability of the MC form of the probe molecule, and thus increase the energy barrier required for thermal relaxation to occur. However, the thermal relaxation process provided far greater energy than this energy barrier, and thus, although the activation energy and the enthalpies of activation increase, the



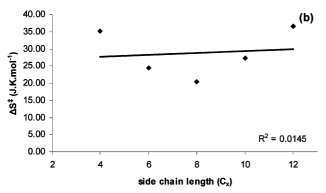
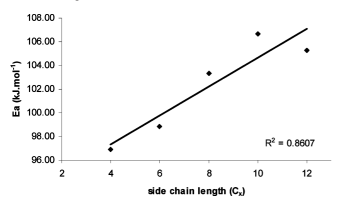


Figure 8. Entropies of activation of BSP (a) and SO (b) versus cation side chain length.



**Figure 9.** Activation energy of MC-BSP thermal relaxation versus cation side chain length.

overall equilibrium has a dominating influence on the process and so increasing rates of relaxation with increasing side chain length is the overall effect observed.

In contrast, for SO, the thermodynamic parameters showed little or no variance with increasing chain length. Since SO is believed to interact with the ILs primarily through electrostatic interactions, it is proposed that the process of thermal relaxation is also somewhat independent of the nature of the IL, provided the charged center remains unchanged, as is the case in this study. The bulky nature of the SO substituents may provide sufficient steric hindrance to prevent hydrogen bonding which would influence the relaxation process. The ability of the SO and  $MC_{SO}$  molecules to move somewhat freely between the IL nanostructured regions also means that the  $MC_{SO} \rightarrow SO$  conversion could occur with relatively little interaction with the IL ions. As a result, the process for thermal relaxation is similar in all ILs tested, and is independent of chain length.

The ground state equilibrium,  $K_e$ , and transition state equilibrium,  $K^{\ddagger}$ , were also examined to determine the effects of chain

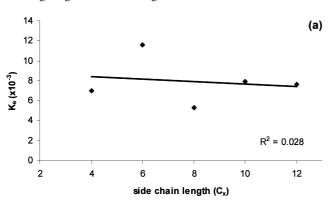
length and IL nanostructure on the  $MC_{SO}$ -SO and  $MC_{BSP}$ -BSP equilibria. Ground state equilibria were determined using eq 5

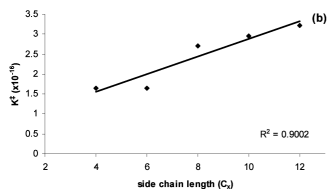
$$K_{\rm e} = \frac{[{\rm MC}]}{[{\rm BSP}]} = \frac{A}{\varepsilon \times C - A}$$
 (5)

where C = concentration of BSP/SO and  $\varepsilon = \text{extinction coefficient } [3.5 \times 10^4 \,\text{M}^{-1} \,\text{cm}^{-1} \,\text{for BSP}^{34} \,\text{and } 7.85 \times 10^4 \,\text{M}^{-1} \,\text{cm}^{-1} \,\text{for SO} \,\,\text{(obtained experimentally by Dr. Minkovska)}].$ 

Ground state equilibrium constants were found to be consistent in relation to increasing IL cation side chain length for both **BSP** and **SO**.  $K_e$  values around  $7 \times 10^{-3}$  for **BSP** and  $5 \times 10^{-3}$  $10^{-3}$  for **SO** were observed in each of the ILs. This implies that the spirocyclic compounds remained in their closed (BSP/ **SO**) form in preference to the respective **MC** form in each of the ILs. Since the closed, spiro forms of the compounds have a lower energy level than that of the respective MC form, it would be expected that they would be primarily in the closed form at equilibrium. However, polar regions within the ILs may shift the equilibrium toward the MC form. Examination of the shorter chain (more polar) ILs showed little change in equilibrium constants compared to that of the longer chain cations. This would appear to reinforce the existence of domains within the solvent system, as increasing the chain length of the IL cation should favor the formation of the spiro form. If the spirocyclic compounds are residing in the nonpolar regions in their closed form, then it would appear that changing the size of such regions has little effect upon the equilibrium. Since spirocyclic compounds thermodynamically favor residing in such regions due to the uncharged nature of their closed form (and its lower thermodynamic energy level), intuitively, it would be expected that the contribution of the nonpolar regions of each IL to the equilibrium constant would be similar or equivalent in each of

To examine the effect of polar regions of the ILs, the transition state equilibrium,  $K^{\dagger}$ , was determined. The equilibrium is based on the thermodynamic interconversion between MC and the respective **BSP/SO** forms, and the process of relaxation is based upon which is the preferred state of the molecule. The transition state equilibrium determines whether the process of relaxation preferentially occurs. At this transition state (MC<sup>‡</sup>), the compound is in the process of relaxation to BSP/SO. However, sufficient interaction with the IL system will increase the stability of the MC form which competes with one another during the process. Unlike the ground state equilibrium constants, where the compound is predominantly in its most stable, ground state (closed) form, the compound is now in its MC form which is inherently unstable, as such isomers are at a thermodynamically higher energy state. The spirocyclic compounds in their excited states therefore preferentially relax to their ground state, but the ability to do so is based upon the equilibrium of the system. The transition state equilibrium, in particular, examines which isomer of the compound is preferred at the transition state, which is related to the relative interactions of the MC form with the IL polar regions and the competing influence of the nonpolar IL regions favoring the closed form. The result is that any changes in the equilibrium are therefore sensitive to changes to the system which can only be observed when the spirocyclic compounds are in the process of thermal relaxation and migrating within the IL. Since the ground state (closed) form of the probe molecules requires energy to form their respective MC isomers, the ground state equilibrium,  $K_e$ , is somewhat biased toward the closed form and so the compound





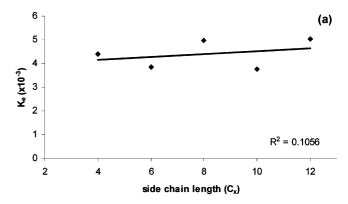
**Figure 10.** (a) Ground state equilibria and (b) equilibria of activation of **BSP** versus cation side chain length.

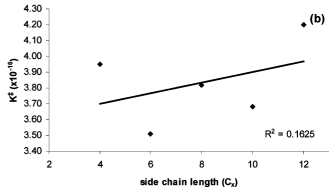
resides consistently in the nonpolar regions of the ILs. This may explain why there is little relationship between the rate constants and the ground state equilibrium constants. In the case of MC thermal relaxation and the transition state equilibrium, the opposite occurs, as the compound must now relax to the spiro form from its higher energetic state.

With increasing chain length, structuring within the solvent is proposed to disperse and weaken the relative interactions of the polar regions. Theoretical models by Lopes agreed with this convention, with  $[C_{12}\text{mIm}]^+$  based ILs showing a majority of nonpolar regions compared to that of  $[C_4\text{mIm}]^+$  ILs. Reduction in the strength of MC–IL interactions results in reduced stability of the MC form and further enhances the formation of the closed form which shifts the equilibrium toward the closed form of the compound. This is observed with increasing  $K^+$  values from  $1.65 \times 10^{-16}$  in  $[C4\text{mIm}]^+$  to  $3.22 \times 10^{-16}$  in  $[C12\text{mIm}]^+$ . For SO, the response in  $K^+$  values due to chain length was not as clear as that observed for BSP. This is believed to be due to the passive nature of  $MC_{SO}$ –IL interactions and the relatively independent nature of the relaxation process resulting from this.

### **Conclusions**

Imidazolium based ILs appear to present nanostructured polar and nonpolar domains. Spirocyclic compounds added to ILs with a nonpolar side chain of increasing length resulted in moderate changes to thermodynamic parameters possibly due to enhanced structuring of the nanodomains. **BSP** was found to be sufficiently sensitive to quantify thermodynamic and kinetic parameters due to its intimate interaction with the imidazolium cation when in its zwitterionic **MC** form. **SO** failed to achieve the same effect, and this was believed to be due to its inability to form hydrogen bonds and its restriction to electrostatic interactions with the IL. Probing of ILs with **BSP** suggests that the molecule may dynamically transfer between the polar and





**Figure 11.** (a) Ground state equilibria and (b) energy of activation of **SO** versus cation side chain length.

nonpolar nanostructured domains in the IL. Ground state equilibria values were similar regardless of the IL cation chain length, which suggests that stabilization effects are similar for each form of the probe molecule in each IL. This may have implied that the ratio of anion to cation was the same in each IL but that the structuring of the IL with increasing chain length resulted in strain and possible dispersion or expansion of such regions from  $[C_2mIm]^+$  to  $[C_{12}mIm]^+$ . The lack of significant variation in activated complex parameters reinforces this interpretation, as similar interactions would logically result in similar levels of stabilization of the respective forms of the spirocyclic compound. Interestingly, positive  $\Delta S^{\ddagger}$  values mean that the compound undergoes significant reordering within the solvent system during thermal relaxation. This further implies that the spirocyclic probe disrupts the IL nanostructure when it is introduced, and it migrates dynamically between the nanostructured regions. This interaction with the ILs may be a reason why rates of relaxation are found to be slow in ILs compared to molecular solvents.

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