

Cite this: *Chem. Sci.*, 2022, 13, 7560

All publication charges for this article have been paid for by the Royal Society of Chemistry

Received 26th April 2022
Accepted 7th June 2022DOI: 10.1039/d2sc02342c
rsc.li/chemical-science

Origin of low melting point of ionic liquids: dominant role of entropy†

Takatsugu Endo,^{ID}*^a Kouki Sunada,^b Hiroki Sumida^b and Yoshifumi Kimura^{ID}^{ab}

Ionic liquids (ILs) are salts with an extremely low melting point. Substantial efforts have been made to address their low melting point from the enthalpic standpoint (*i.e.* interionic interactions). However, this question is still open. In this study, we report our findings that entropic (large fusion entropy), rather than enthalpic, contributions are primarily responsible for lowering the melting point in many cases, based on a large thermodynamic dataset. We have established a computational protocol using molecular dynamics simulations to decompose fusion entropy into kinetic (translational, rotational, and intramolecular vibrational) and structural (conformational and configurational) terms and successfully applied this approach for two representatives of ILs and NaCl. It is revealed that large structural contribution, particularly configurational entropy in the liquid state, plays a deterministic role in the large fusion entropy and consequently the low melting point of the ILs.

Introduction

Salts are defined as materials composed solely of ions. Since interactions between cations and anions are governed by strong coulombic interactions, with the order of magnitudes of several hundred kJ mol⁻¹, melting points of salts tend to be extremely high. The typical example is table salt (NaCl), whose melting point is 1073 K. However, in the late 1990s, scientists started to recognize salts with extremely low melting points corresponding to room temperature or even below. Nowadays, these salts are called room temperature ionic liquids or simply ionic liquids (ILs). ILs have attracted considerable attention because they have several substantial properties, such as negligible vapor pressure and flammability, high thermal/chemical/electrochemical stabilities, high ionic/electric conductivity, and unique solubilities.^{1–3} These properties mark ILs as promising candidates in a wide range of applications, for example, green solvents, electrolytes, CO₂ absorbents, and cellulose solvents.^{1–3}

The melting point is considered the most important physical property of ILs since it distinguishes them from other salts. Many investigations have been intensively conducted to answer why ILs have such a low melting point; moreover, various solutions have been proposed.^{1,4–9} Thermodynamically, contributions to the melting point are divided into two terms, *i.e.*,

enthalpic and entropic terms. Although the importance of the entropic contributions (*e.g.*, conformational entropy stemming from flexible side groups) were pointed out,^{1,4,6,8,9} the mainstream discussion thus far has been made on enthalpic contributions.^{1,4–9} Namely, coulombic interaction among ions and lattice energy are weakened in ILs due to bulky, asymmetric, and sometimes charge-delocalized ions. In this work, we demonstrate that this is not the right approach and that the low melting point is caused due to entropic contribution in the majority of ILs.

Results and discussion

Alkali halides *versus* ionic liquids

From a thermodynamical standpoint, melting point (T_m) of material is the ratio of fusion enthalpy ($\Delta_{\text{fus}}H$) and fusion entropy ($\Delta_{\text{fus}}S$):

$$T_m = \frac{\Delta_{\text{fus}}H}{\Delta_{\text{fus}}S} \quad (1)$$

This rigorous expression shows that the melting point is low when $\Delta_{\text{fus}}H$ is small and/or $\Delta_{\text{fus}}S$ is large. Stating that the melting point of ILs is “low” requires a reference material for comparison. NaCl with a melting point of 1073 K is frequently selected. For statistical comparison purposes, 20 alkali halides were selected in this study, and their thermodynamic quantities¹⁰ were compared with those of 257 ILs.^{11,12} As the name suggests, the average T_m of ILs is ~ 3 times lower than that of alkali halides (Fig. 1A). An average value of $\Delta_{\text{fus}}H$ for ILs, associated with interionic interactions in the liquid and crystal states, is only 0.85 times the $\Delta_{\text{fus}}H$ value of alkali halides

^aDepartment of Molecular Chemistry and Biochemistry, Faculty of Science and Engineering, Doshisha University, 1-3 Tatara Miyakodani, Kyotanabe, Kyoto 610-0394, Japan. E-mail: taendo@mail.doshisha.ac.jp

^bDepartment of Applied Chemistry, Graduate School of Science and Engineering, Doshisha University, 1-3 Tatara Miyakodani, Kyotanabe, Kyoto 610-0394, Japan

† Electronic supplementary information (ESI) available. See <https://doi.org/10.1039/d2sc02342c>



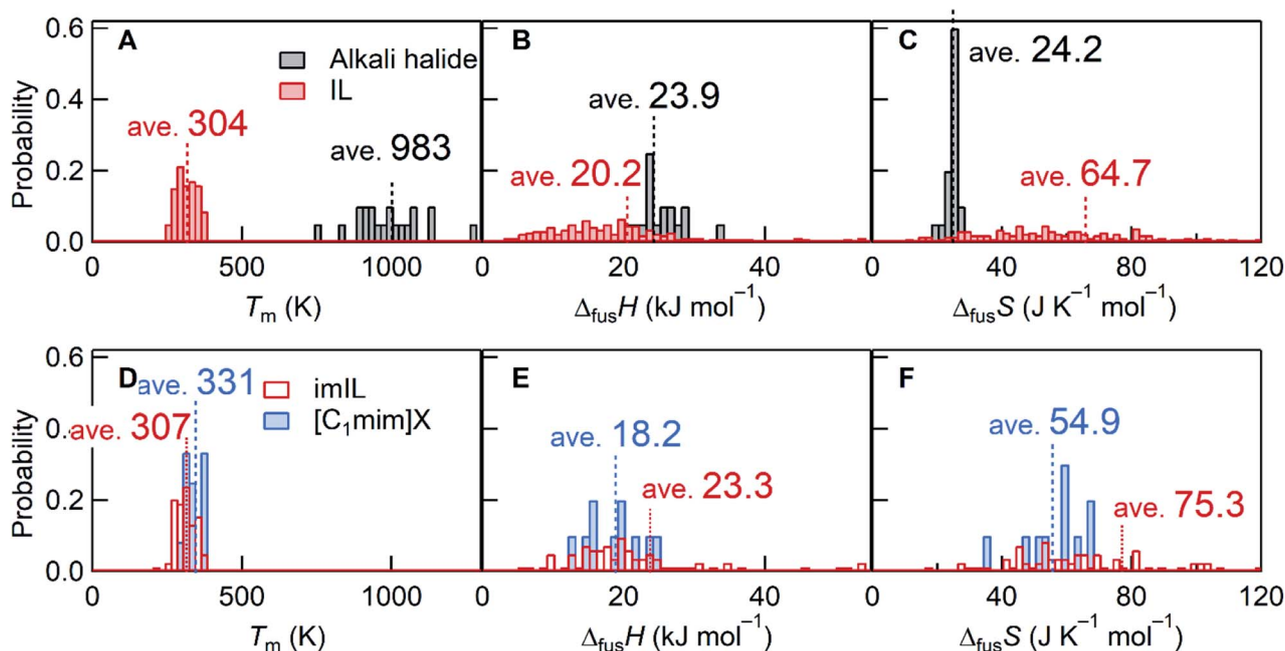


Fig. 1 Histogram comparisons of T_m , $\Delta_{fus}H$, and $\Delta_{fus}S$. (A–C) 20 alkali halides¹⁰ versus 257 ILs^{11,12} and (D–F) 84 imILs versus 12 [C₁mim]X. For [C₁mim]X, this work reported 10 data and the two additional data were taken from the ref. 41 and 42 (Table S3†). Bin widths for the T_m , $\Delta_{fus}H$, and $\Delta_{fus}S$ histograms were 20 K, 1 kJ mol⁻¹, and 2 J K⁻¹ mol⁻¹, respectively. For ILs and imILs, several data of $\Delta_{fus}H$ and $\Delta_{fus}S$ are out of range of the figures.

(Fig. 1B). Meanwhile, the average $\Delta_{fus}S$ of ILs is 2.67 times larger than that of alkali halides (Fig. 1C), which indicates that, in general, the large $\Delta_{fus}S$ plays a more critical role than the small $\Delta_{fus}H$ for lowering T_m of ILs.

Since the discussion would depend on the ions constituting the IL, we focus on imidazolium-based ILs (imILs) as the most representative IL series. Analyzing the available data on 84 imILs,^{11,12} the importance of the large $\Delta_{fus}S$ is emphasized as seen in Fig. 1D–F. While $\Delta_{fus}H$ values are comparable, their $\Delta_{fus}S$ values are 3.11 times larger than those corresponding to alkali halides. Considering the origin of the large $\Delta_{fus}S$ of ILs (or imILs), a prominent contribution may come from conformational entropy as described above.^{1,4,6,8,9} For example, most IL cations contain flexible alkyl chains that produce multiple conformations in the liquid state. This increases the entropy of the liquid state and consequently $\Delta_{fus}S$. An approach to thermodynamically elucidate the role of conformational entropy is to compare $\Delta_{fus}S$ of imILs with flexible side chains to those without flexible groups. However, such data is rare. Therefore, we synthesized ten 1,3-dialkylimidazolium salts ([C₁mim]X) with no conformational entropy in the cation and the anion and measured T_m , $\Delta_{fus}H$, and $\Delta_{fus}S$. The difference in T_m between [C₁mim]X and imILs is rather low, whereas both $\Delta_{fus}H$ and $\Delta_{fus}S$ of [C₁mim]X are smaller than those of imILs (Fig. 1D–F). The small $\Delta_{fus}H$ and $\Delta_{fus}S$ values would be primarily due to the losses of inter-chain interactions^{1,6–9} and (as expected) conformational entropy, respectively. However, even without conformational entropy, not the small $\Delta_{fus}H$ (18.2 kJ mol⁻¹) but the large $\Delta_{fus}S$ (54.9 J K⁻¹ mol⁻¹) seems to prevail in lowering T_m of

ILs, compared to alkali halides ($\Delta_{fus}H = 23.9$ kJ mol⁻¹ and $\Delta_{fus}S = 24.2$ J K⁻¹ mol⁻¹).

Fusion entropy decomposition

To quantitatively unravel ambiguous entropic contributions, entropy needs to be decomposed into individual components associated with molecular-level properties. The decompositions were conducted based on the free energy landscape model^{13,14} (Fig. 2). Absolute entropy (S) can be expressed as the sum of

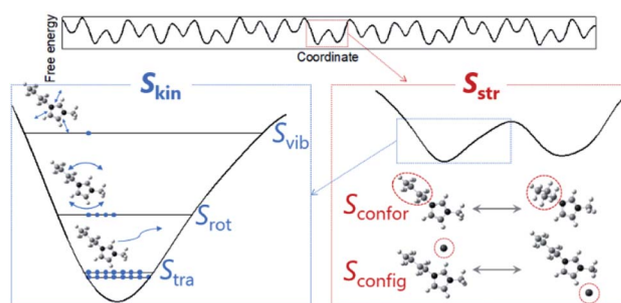


Fig. 2 Schematic of entropies based on the free energy landscape. Absolute entropy is expressed as the sum of kinetic (S_{kin}) and structural (S_{str}) contributions. Kinetic entropy corresponds to the number of distinguishable states where a particle of interest occupies (blue symbols) in a potential well. Structural entropy, corresponding to the number of potential wells, is expressed as the sum of conformational (or intramolecular) (S_{confor}) and configurational (or intermolecular) (S_{config}) parts. S_{config} is displayed based on an ion-pair model where an anion is drawn as a monoatomic ion.



kinetic (S_{kin}) and structural (S_{str}) entropies. The former is further divided into translational (S_{tra}), rotational (S_{rot}), and intramolecular vibrational (S_{vib}) terms and the latter is composed of conformational/intramolecular (S_{confor}) and configurational/intermolecular (S_{config}) entropies. Since we focus on the difference in entropy between liquids and crystals at the melting point, by assuming individual terms are independent, appropriate equations are as follows:

$$\Delta_{\text{fus}}S = \Delta_{\text{kin}}S + \Delta_{\text{str}}S \quad (2)$$

$$\Delta_{\text{kin}}S = \Delta_{\text{tra}}S + \Delta_{\text{rot}}S + \Delta_{\text{vib}}S \quad (3)$$

$$\Delta_{\text{str}}S = \Delta_{\text{confor}}S + \Delta_{\text{config}}S \quad (4)$$

To perform the $\Delta_{\text{fus}}S$ decomposition, we have developed a computational protocol using classical molecular dynamics (MD) simulations (see the ESI† for details). $\Delta_{\text{fus}}S$ and $\Delta_{\text{confor}}S$ were estimated from the thermodynamic integration^{15,16} and conformational analyses, respectively. We employed the two-phase thermodynamic (2PT) approach, initially developed by Lin *et al.*, for the kinetic entropy estimations.^{17–19} Conventionally, the entropy of vibrational motions for molecules in a solid state has been estimated through the density of states functions $g(\nu)$ by assuming that all vibrational motions are harmonic oscillators. However, this technique cannot treat diffusive motions existing in a liquid state. The 2PT approach enables the estimation of entropy in a liquid state by dividing $g(\nu)$ in the liquid state into solid-like (for harmonic oscillators) and gas-like (for diffusive motions) components. The remaining $\Delta_{\text{config}}S$ was obtained based on eqn (2)–(4) by the deduction of the other terms. For reliable discussion on results obtained from classical MD simulations, a selection of force fields is critical. We selected the force fields developed by Maginn *et al.* for two prototype imILs as 1-ethyl-3-methylimidazolium hexafluorophosphate ([C₂mim]PF₆) and 1-butyl-3-methylimidazolium hexafluorophosphate ([C₄mim]PF₆).²⁰ Partial charges in these force fields were determined by *ab initio* MD in the crystal state with periodic boundary conditions so that the charge transfer and polarization in the condensed state are implicitly included. Furthermore, MD simulations on NaCl with the Tosi–Fumi potential²¹ were conducted for comparison. Simulated T_{m} , $\Delta_{\text{fus}}H$, and $\Delta_{\text{fus}}S$ were in good agreement with the experimental values (Table S7†). It is noted that the absolute entropies also satisfactorily reproduced experimental data (Table S10†).

The results of the $\Delta_{\text{fus}}S$ decompositions for NaCl and the two ILs are presented in Fig. 3. In NaCl that is composed of monoatomic ions, the following stands: $\Delta_{\text{fus}}S \approx \Delta_{\text{kin}}S = \Delta_{\text{tra}}S$. For the ILs, $\Delta_{\text{kin}}S$ is smaller than that of NaCl, even though the ILs have larger absolute entropy than NaCl in any state of matter, mainly due to the presence of S_{rot} and S_{vib} (Fig. S14†). There are three reasons for this counter-intuitive behavior. First, $\Delta_{\text{vib}}S$ of the ILs does not contribute to $\Delta_{\text{kin}}S$, or they are even slightly negative. This would result from the fact that intramolecular vibrations are mostly unchanged by melting, as suggested by previous reports.^{22,23} The second reason is related to the difference in

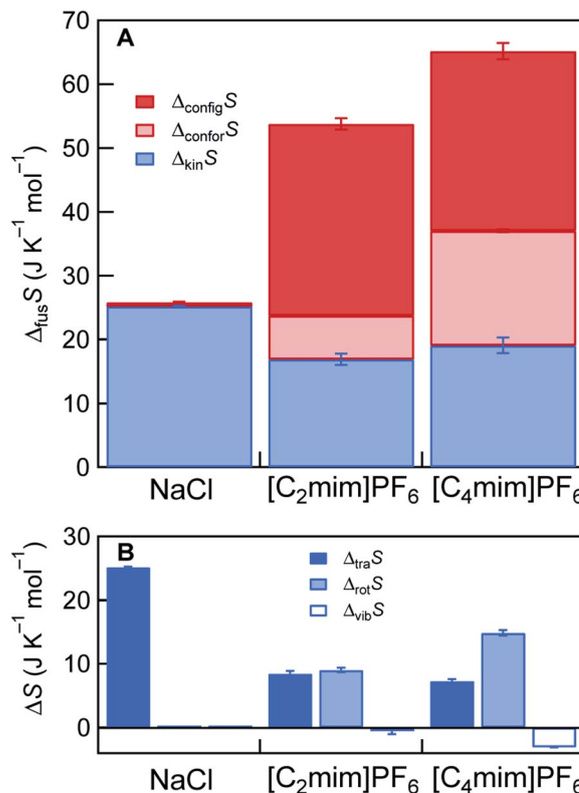


Fig. 3 Decomposition of simulated $\Delta_{\text{fus}}S$. For [C₂mim]PF₆ and [C₄mim]PF₆, $\Delta_{\text{vib}}S$ shows slight negative values, hence $\Delta_{\text{kin}}S (= \Delta_{\text{tra}}S + \Delta_{\text{rot}}S + \Delta_{\text{vib}}S)$ is displayed in (A). Individual $\Delta_{\text{tra}}S$, $\Delta_{\text{rot}}S$, and $\Delta_{\text{vib}}S$ are shown in (B).

melting point. Since the melting points of the ILs are intrinsically lower than that of NaCl, diffusive motions (translational + rotational) represented by $g(0)$ (density of states at zero frequency) are not activated in the ILs, compared to NaCl (Fig. S11–S13 and Table S12†). The slow diffusive motions are also expected for other ILs since their viscosity is generally as high as several tens or more mPa s at the melting point. Third, the spherical PF₆⁻ anions already gain rotational diffusivity in the crystal state (Fig. S12C and S13C†), consistent with previous NMR observations,²⁴ resulting in lowering $\Delta_{\text{rot}}S$.

Despite the small $\Delta_{\text{kin}}S$ of the ILs, the presence of large $\Delta_{\text{str}}S (= \Delta_{\text{confor}}S + \Delta_{\text{config}}S)$ is the cause for the large $\Delta_{\text{fus}}S$. The presence of $\Delta_{\text{confor}}S$ has been previously pointed out by several groups.^{1,4,6,8,9} The calculated $\Delta_{\text{confor}}S$ values were 6.8 J K⁻¹ mol⁻¹ for [C₂mim]PF₆ and 18.0 J K⁻¹ mol⁻¹ for [C₄mim]PF₆. These values are slightly smaller than the previously reported S_{confor} in the gas state (8.3 J K⁻¹ mol⁻¹ for [C₂mim]PF₆ and 22.8 J K⁻¹ mol⁻¹ for [C₄mim]PF₆).²⁵ This would result from the fact that S_{confor} in the crystalline state was calculated to be non-zero due to the slight disordering of the alkyl group in the cation (Tables S8 and S9†). A significant feature in Fig. 3A is that $\Delta_{\text{config}}S$ plays a more prominent role in lowering T_{m} than $\Delta_{\text{confor}}S$. The $\Delta_{\text{config}}S (= \text{here } S_{\text{config}} \text{ in the liquid state})$ values of [C₂mim]PF₆ and [C₄mim]PF₆ were estimated to be 30.0 J K⁻¹ mol⁻¹ and 28.1 J K⁻¹ mol⁻¹, respectively. Assuming $S_{\text{config}} = R \ln W$, where W is the number of distinguishable microscopic states, at least 30–



40 potential energy basins exist for these ILs in the liquid state. Although multiple configurations in ILs were already pointed out, and they were associated with the low melting point in a few papers,^{8,26} we quantitatively demonstrate the dominant role of $\Delta_{\text{config}}S$ in the low melting point of ILs. The origin of multiple configurations is considered to come from the delocalized charges in the ion and the asymmetric ion structure.²⁶ It is worth noting that these characteristics of ILs have been previously considered to contribute to lowering the melting point enthalpically by reducing coulombic interactions and packing efficiency.^{1,6,8,9} Our findings indicate that the delocalized charge and the asymmetric ion structure can be entropically, rather than enthalpically, important for the low melting point. Since large $\Delta_{\text{kin}}S$ cannot be expected for general ILs, because of close-to-zero $\Delta_{\text{vib}}S$ and high viscosity (small $\Delta_{\text{tra}}S + \Delta_{\text{rot}}S$), large $\Delta_{\text{str}}S$ (particularly large $\Delta_{\text{config}}S$) is the key for the large $\Delta_{\text{fus}}S$ and the consequent low melting point of ILs.

Conclusions

In this work, we have discovered that the low melting point of ILs is mainly driven entropically, *i.e.*, the large $\Delta_{\text{fus}}S$ dominantly decreases the melting point of ILs. To unravel the origin of the large $\Delta_{\text{fus}}S$, imILs as the most representative IL series were focused. With the developed protocol based on MD simulations, $\Delta_{\text{fus}}S$ of the two imILs as well as NaCl were successfully decomposed into individual components with a clear physical origin. The simulations indicated that, somewhat counterintuitively, not the kinetic but the structural part, particularly $\Delta_{\text{config}}S$, was the main contributor to the low melting point of the ILs. Breaking the mainstream of the previous “enthalpic” discussion, our findings based on the “entropic” contributions can change the bottom concept for designing ILs and could consequently accelerate the production of a number of novel functional ILs.

Experimental

Selection of thermodynamic data of ILs

The thermodynamic data, T_m , $\Delta_{\text{fus}}H$, $\Delta_{\text{fus}}S$ for ILs were taken from the ILThermo database as summarized in Table S1.[†] (It is noted that T_m , $\Delta_{\text{fus}}H$, $\Delta_{\text{fus}}S$ for alkali halide are also summarized in Table S2[†]).¹⁰ The ILThermo database seems to contain some inappropriate data for discussion, *i.e.*, T_m is too high to be ILs, or $\Delta_{\text{fus}}H$ or $\Delta_{\text{fus}}S$ is too low to be considered as enthalpy/entropy difference on melting. In this work, we excluded the data with $T_m > 373.2$ K, $\Delta_{\text{fus}}H < 3$ kJ mol⁻¹, or $\Delta_{\text{fus}}S < 10$ J K⁻¹ mol⁻¹ from the discussion. In some cases, several different data were reported in one IL because of, *e.g.*, differences in experimental conditions, experimental error, and purities as well as the existence of polymorphism. Considering the purity and thermodynamic stability of polymorphs, we selected the one with the highest melting point.

IL syntheses and characterizations

The ILs [C₁mim]X were synthesized following standard procedures.¹ The detailed procedures were described in the ESI.[†] The

ILs were vacuum-dried with heating for several days (typically 353 K for 2 days) before use. The synthesized ILs were characterized with NMR spectroscopy (JEOL, JNM-ECA300W). No recognizable impurities were observed *via* NMR spectra except for water. The water content of the ILs was determined by Karl Fischer titration (MKC-501, Kyoto Electronics). When an IL was in the crystal state at room temperature, the water content measurement was conducted by dissolving it in anhydrous acetonitrile. Several ILs potentially contain a slight amount of alkali halides as a byproduct, the content of which was determined by Na⁺ meter (LAQUATwin Na-11, Horiba) or the Mohr's method.

DSC measurements

Differential scanning calorimetry (DSC) measurements were performed with a DSC7020 (Hitachi High-Tech Science). A dried IL was sealed in an Al pan in an Ar-atmosphere glovebox. In the measurements, the sample was first melted by heating, and data was subsequently collected by cooling down to 175 K and re-heating above the melting point at a scanning rate of 5 K min⁻¹. The obtained DSC traces are shown in Fig. S1,[†] and the numerical values are listed in Tables S3 and S4.[†]

Quantum chemical calculations

Quantum chemical calculations were performed to estimate absolute entropies of NaCl, [C₂mim]PF₆, and [C₄mim]PF₆ in the gas phase with Gaussian09²⁷ program package. B3LYP^{28–30} combined with GD3 (ref. 31) correction at 6-311++G(d,p) level were employed for the calculations. Full geometry optimization and subsequent vibrational frequency analyses were performed. The initial structure of the IL ion pairs for the geometry optimization was referred to the ones reported previously³² (Fig. S2[†]). It is noted that no imaginary frequency was obtained for the optimized structures, which confirms that they are local minima. The absolute entropies and the decomposed kinetic entropies (S_{tra} , S_{rot} , and S_{vib}) for NaCl, [C₂mim]PF₆, and [C₄mim]PF₆ are summarized in Table S5.[†]

Molecular dynamics simulations

Classical MD simulations were performed with Gromacs 2018.4 Software package^{33,34} (single precision) with the periodic boundary condition. Newton's equation of motion was solved in 1 fs time step with the leap-frog algorithm unless otherwise stated. Cut-off radii for both coulombic and LJ potentials were set to be 1.2 nm. Long-range dispersion corrections were applied. Long-range coulombic interactions were treated with the particle mesh Ewald summation method.^{35,36} Temperature and pressure were controlled with Nose–Hoover thermostat and Parrinello–Rahman barostat, respectively. No constraint was applied for intramolecular bonds and angles.

Ion pairs of 2048 (NaCl), 400 ([C₂mim]PF₆), and 250 ([C₄mim]PF₆) were used in MD simulations. The force field of NaCl was based on the Tosi–Fumi potential.²¹ The ones developed by Maginn *et al.*²⁰ with the aid of the general AMBER force field^{37,38} were used for the two ILs. A crystal structure with fcc lattice was employed for the NaCl crystal. Crystal structures of



[C₂mim]PF₆ (ref. 39) and [C₄mim]PF₆ (ref. 40) were taken from the references. For liquid state simulations, ions were randomly placed into a cubic cell. After energy minimizations with the steepest descent algorithm, an initial simulation box was first equilibrated in the NPT ensemble until system energy became constant, at least for 2 ns. If necessary, a subsequent NVT ensemble simulation was conducted with the equilibrated cell size. For statistical error estimation, a production run after the equilibration was equally divided into 4 blocks. The standard deviation based on 4 simulations was taken as the statistical error. It is noted that in the figures displayed in the main text and the ESI,[†] the average data of the 4 blocks are shown. Detailed procedures of estimations of T_m , $\Delta_{fus}H$, $\Delta_{fus}S$, and the entropies were described in the ESI.[†]

Data availability

All available data are included in the ESI.[†]

Author contributions

T. Endo conceived and designed this study. K. Sunada synthesized the compounds. T. Endo performed the molecular dynamics simulations. H. Sumida performed the quantum chemical calculations. T. Endo analyzed the data. T. Endo produced the manuscript. K. Sunada, H. Sumida and K. Kimura checked and modified the manuscript.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

We thank Dr Miguel Caro (Aalto University, Finland) for his helpful advice on the 2PT analyses. The work was supported by JSPS KAKENHI Grant Number JP19K05393.

Notes and references

- 1 D. R. MacFarlane, M. Kar and J. M. Pringle, *Fundamentals of Ionic Liquids: From Chemistry to Applications*, Wiley-VCH, 2017.
- 2 D. Rogers Robin and R. Seddon Kenneth, *Science*, 2003, **302**, 792–793.
- 3 N. V. Plechkova and K. R. Seddon, *Chem. Soc. Rev.*, 2008, **37**, 123–150.
- 4 I. Krossing, J. M. Slattery, C. Daguenet, P. J. Dyson, A. Oleinikova and H. Weingärtner, *J. Am. Chem. Soc.*, 2006, **128**, 13427–13434.
- 5 S. Zahn, F. Uhlig, J. Thar, C. Spickermann and B. Kirchner, *Angew. Chem., Int. Ed.*, 2008, **47**, 3639–3641.
- 6 P. M. Dean, J. M. Pringle and D. R. MacFarlane, *Phys. Chem. Chem. Phys.*, 2010, **12**, 9144–9153.
- 7 Y. V. Nelyubina, A. S. Shaplov, E. I. Lozinskaya, M. I. Buzin and Y. S. Vygodskii, *J. Am. Chem. Soc.*, 2016, **138**, 10076–10079.
- 8 E. I. Izgorodina, Z. L. Seeger, D. L. A. Scarborough and S. Y. S. Tan, *Chem. Rev.*, 2017, **117**, 6696–6754.
- 9 F. Philippi and T. Welton, *Phys. Chem. Chem. Phys.*, 2021, **23**, 6993–7021.
- 10 G. J. Janz, *Molten Salts Handbook*, Academic, 1967.
- 11 Q. Dong, C. D. Muzny, A. Kazakov, V. Diky, J. W. Magee, J. A. Widegren, R. D. Chirico, K. N. Marsh and M. Frenkel, *J. Chem. Eng. Data*, 2007, **52**, 1151–1159.
- 12 A. Kazakov, J. W. Magee, R. D. Chirico, E. Paulechka, V. Diky, C. D. Muzny, K. Kroenlein and M. Frenkel, *NIST Standard Reference Database 147: NIST Ionic Liquids Database – (ILThermo), Version 2.0*, National Institute of Standards and Technology, Gaithersburg MD, 2021, p. 20899, <http://ilthermo.boulder.nist.gov>.
- 13 L. Berthier, M. Ozawa and C. Scalliet, *J. Chem. Phys.*, 2019, **150**, 160902.
- 14 F. H. Stillinger, *Science*, 1995, **267**, 1935–1939.
- 15 D. M. Eike, J. F. Brennecke and E. J. Maginn, *J. Chem. Phys.*, 2005, **122**, 014115.
- 16 K. Bernardino, Y. Zhang, M. C. C. Ribeiro and E. J. Maginn, *J. Chem. Phys.*, 2020, **153**, 044504.
- 17 S. T. Lin, M. Blanco and W. A. Goddard III, *J. Chem. Phys.*, 2003, **119**, 11792–11805.
- 18 M. A. Caro, T. Laurila and O. Lopez-Acevedo, *J. Chem. Phys.*, 2016, **145**, 244504.
- 19 S. T. Lin, P. K. Maiti and W. A. Goddard III, *J. Phys. Chem. B*, 2010, **114**, 8191–8198.
- 20 Y. Zhang and E. J. Maginn, *Phys. Chem. Chem. Phys.*, 2012, **14**, 12157–12164.
- 21 J. L. Aragonés, E. Sanz, C. Valeriani and C. Vega, *J. Chem. Phys.*, 2012, **137**, 104507.
- 22 J. Dupont, *J. Braz. Chem. Soc.*, 2004, **15**, 341–350.
- 23 V. H. Paschoal, L. F. O. Faria and M. C. C. Ribeiro, *Chem. Rev.*, 2017, **117**, 7053–7112.
- 24 T. Endo, H. Murata, M. Imanari, N. Mizushima, H. Seki, S. Sen and K. Nishikawa, *J. Phys. Chem. B*, 2013, **117**, 326–332.
- 25 T. Endo, T. Higuchi and Y. Kimura, *Bull. Chem. Soc. Jpn.*, 2020, **93**, 720–729.
- 26 P. A. Hunt, *J. Phys. Chem. B*, 2007, **111**, 4844–4853.
- 27 M. J. Frisch, G. W. Trucks, H. B. Schlegel, G. E. Scuseria, M. A. Robb, J. R. Cheeseman, G. Scalmani, V. Barone, B. Mennucci, G. A. Petersson, H. Nakatsuji, M. Caricato, X. Li, H. P. Hratchian, A. F. Izmaylov, J. Bloino, G. Zheng, J. L. Sonnenberg, M. Hada, M. Ehara, K. Toyota, R. Fukuda, J. Hasegawa, M. Ishida, T. Nakajima, Y. Honda, O. Kitao, H. Nakai, T. Vreven, J. A. Montgomery Jr, J. E. Peralta, F. Ogliaro, M. Bearpark, J. J. Heyd, E. Brothers, K. N. Kudin, V. N. Staroverov, T. Keith, R. Kobayashi, J. Normand, K. Raghavachari, A. Rendell, J. C. Burant, S. S. Iyengar, J. Tomasi, M. Cossi, N. Rega, J. M. Millam, M. Klene, J. E. Knox, J. B. Cross, V. Bakken, C. Adamo, J. Jaramillo, R. Gomperts, R. E. Stratmann, O. Yazyev, A. J. Austin, R. Cammi, C. Pomelli, J. W. Ochterski, R. L. Martin, K. Morokuma, V. G. Zakrzewski, G. A. Voth, P. Salvador, J. J. Dannenberg, S. Dapprich, A. D. Daniels, O. Farkas, J. B. Foresman,



- J. V. Ortiz, J. Cioslowski and D. J. Fox, *Gaussian 09*, Gaussian, Inc., Wallingford CT, 2010.
- 28 A. D. Becke, *J. Chem. Phys.*, 1993, **98**, 5648–5652.
- 29 C. Lee, W. Yang and R. G. Parr, *Phys. Rev. B*, 1988, **37**, 785–789.
- 30 B. Miehlich, A. Savin, H. Stoll and H. Preuss, *Chem. Phys. Lett.*, 1989, **157**, 200–206.
- 31 S. Grimme, J. Antony, S. Ehrlich and H. Krieg, *J. Chem. Phys.*, 2010, **132**, 154104.
- 32 B. A. Marekha, O. N. Kalugin and A. Idrissi, *Phys. Chem. Chem. Phys.*, 2015, **17**, 16846–16857.
- 33 D. Van Der Spoel, E. Lindahl, B. Hess, G. Groenhof, A. E. Mark and H. J. C. Berendsen, *J. Comput. Chem.*, 2005, **26**, 1701–1718.
- 34 C. Kutzner, S. Páll, M. Fechner, A. Esztermann, B. L. de Groot and H. Grubmüller, *J. Comput. Chem.*, 2019, **40**, 2418–2431.
- 35 T. Darden, D. York and L. Pedersen, *J. Chem. Phys.*, 1993, **98**, 10089–10092.
- 36 U. Essmann, L. Perera, M. L. Berkowitz, T. Darden, H. Lee and L. G. Pedersen, *J. Chem. Phys.*, 1995, **103**, 8577–8593.
- 37 J. Wang, R. M. Wolf, J. W. Caldwell, P. A. Kollman and D. A. Case, *J. Comput. Chem.*, 2004, **25**, 1157–1174.
- 38 H. Chen, T. Yan and G. A. Voth, *J. Phys. Chem. A*, 2009, **113**, 4507–4517.
- 39 W. M. Reichert, J. D. Holbrey, R. P. Swatloski, K. E. Gutowski, A. E. Visser, M. Nieuwenhuyzen, K. R. Seddon and R. D. Rogers, *Cryst. Growth Des.*, 2007, **7**, 1106–1114.
- 40 A. R. Choudhury, N. Winterton, A. Steiner, A. I. Cooper and K. A. Johnson, *J. Am. Chem. Soc.*, 2005, **127**, 16792–16793.
- 41 T. Endo, T. Morita and K. Nishikawa, *Chem. Phys. Lett.*, 2011, **517**, 162–165.
- 42 U. Domańska, A. Pobudkowska and F. Eckert, *Green Chem.*, 2006, **8**, 268–276.

