Surface crystallization of ionic liquid crystals


The evidence for surface crystallization in ionic liquids is scarce. The existing reports seem to be contradictory as for its driving forces, since in the two compounds investigated in the literature, the contribution of coulombic and van der Waals forces is very different. In this work 1-dodecyl-3-methylimidazolium tetrafluoroborate was studied and its surface crystallization characterized by surface tension, ellipsometry and optical microscopy. The results obtained seem to reconcile previous observations, and it was further shown, using the same techniques, that this phenomenon is prevalent in other ionic liquids. MD simulation results illustrate the different possibilities of organization, providing reasonable models to rationalize the experimental observations.

Introduction

Ionic liquid crystals (ILCs) combine the ionic character of ionic liquids (ILs) with the benefits of liquid crystal (LC) anisotropy. They are characterized by long alkyl side chains, relatively low melting points and at least one liquid-crystalline mesomorph. Due to their potential applications in diverse fields, this class of compounds has been extensively investigated. & applications involving ILCs rely on their interfaces, detailed knowledge and understanding of their surfaces thus being important. While the surface and surface properties of ionic liquids have been extensively studied, studies on the surface of ILCs are scarce. The near-surface structure of [C14C1im][FAP] was investigated through X-ray absorption spectroscopy and resonant soft X-ray reflectivity, by Mezger et al., revealing depth-decaying near-surface layering. The authors did not claim however, that the observed surface ordering was akin to a LC. Yet, later Mezger and co-workers studied [C22C1im][NTf2] and [C20C1im][NTf2] observing the formation of four-layered, rectangularly packed, crystalline domains covering 5–15% of the surface. As argued by the authors, the dense packing of alkyl chains, vertically aligned normal to the surface, induce the formation of a higher-ordered surface phase, different from that of the bulk.

Only two works have so far been reported on the surface crystallization of ionic liquids of a widely different nature. Jeon et al., studied a short-chain Coulomb-dominated IL ([C4C1im][PF6]) observing the formation of four-layered, rectangularly packed, crystalline domains covering 5–15% of the surface. Pontoni et al., using a long-chain vdW-dominated IL ([C20C1im][NTf2]) observed a hexagonal crystalline monolayer covering the whole surface. As argued by the authors, the dense packing of alkyl chains, vertically aligned normal to the surface, requires the presence of long-chain impurities.

This work aims at studying the surface of ILCs at temperatures from the melting point to above the clearing point.
this purpose, ILC 1-dodecyl-3-methylimidazolium tetrafluoroborat ([C12C1im][BF4]) was chosen as a model compound due to its relatively low melting and clearing temperatures, and its large and stable mesophase region.3,20–22 Its choice relied also on the intermediate nature of the interactions present, when compared with the ILs previously studied via surface freezing, in terms of the van der Waals interactions due to its medium size alkyl chain length, yet with strong coulombic interactions resulting from the [BF4]− anion. This ILC was shown to be an exceptional medium for the selective synthesis of monoacylglycerides23 and a promising nuclear magnetic resonance (NMR) alignment medium.24

### Experimental

#### Chemicals

The ionic liquid crystal 1-dodecyl-3-methylimidazolium tetrafluoroborat ([C12C1im][BF4]), CAS 244193-56-4, M = 338.24 g mol−1 was purchased from Iolitec with a purity of >98% (according to the supplier). Prior to each measurement and in order to reduce impurities, like water and volatile compounds, individual samples of [C12C1im][BF4] were dried under vacuum (0.1 Pa), moderate temperature (313.15 K) and constant stirring, for a minimum period of 48 h. The water content of the dried samples was determined using a Metrohm 831 Karl Fischer coulometer, with the anlyte Hydranal – Coulomat AG obtained from Riedel-de Haën, and was found to be below 100 ppm for all samples. The chemical structure of this compound is present in Fig. 1. In addition, the following compounds were also used: 1-methyl-3-tetradecylimidazolium tetrafluoroborate ([C14C1im][BF4]), CAS 244193-61-1, M = 266.28 g mol−1, Iolitec, 98%), 1-dodecyl-3-methylimidazolium chloride ([C12C1im]Cl), CAS 114569-84-5, M = 286.89 g mol−1, Iolitec, >98%), 1-methyl-3-tetradecylimidazolium chloride ([C14C1im]Cl), CAS 171058-21-2, M = 314.94 g mol−1, Iolitec, 98%), 1-methyl-3-tetradecylimidazolium bromide ([C14C1im]Br, CAS 471907-87-6, M = 359.39 g mol−1) and dimethylidietradecylammonium chloride ([N111414]Cl, CAS 10108-91-5, M = 474.29 g mol−1). [C12C1im]Br and [N111414]Cl were synthesized in our laboratory.25

#### Methods

**Differential scanning calorimetry (DSC)**

The thermal events of the pure ILC were determined using a Hitachi DSC7000X model working at atmospheric pressure and coupled to a liquid nitrogen cooling system. Samples of approximately 5 mg, tightly sealed in aluminum pans, were submitted to a heating–cooling–heating regime at least 3 times. The heating and cooling rates were 1 K min−1 and 2 K min−1, respectively. The melting temperatures were taken as the peak temperatures. The equipment was previously calibrated with anthracene, benzoic acid, caffeine, decane, diphenylacetic acid, heptane, indium, lead, naphthalene, 4-nitrotoluene, potassium nitrate, tin, water and zinc; all with weight fraction purities higher than 99%.

**Density and viscosity**

Density measurements of the pure ILC [C12C1im][BF4] were carried out at atmospheric pressure and in the 303.16 to 363.14 K temperature range, using an Anton Paar GmbH 4500 vibrating-tube densimeter (Graz, Austria). Two integrated Pt 100 platinum thermometers provided good precision of the internal temperature control (±0.01 K) and the densimeter includes an automatic correction for the viscosity of the sample. The overall uncertainty of the measurements was estimated to be better than 5 × 10−5 g cm−3. Viscosities were carried out using an automated SVM 3000 Anton Paar rotational Stabinger viscometer–densimeter (temperature uncertainty: 0.02 K; dynamic viscosity relative uncertainty: 0.35%), working at atmospheric pressure. Measurements were performed in the temperature range from 319.75 to 373.15 K.

**Interfacial tension**

Three different setups were used: the interfacial tension setup: a Dataphysics contact angle system OCA-20 was used to determine the surface tension of the ILC [C12C1im][BF4] through the analysis of the shape of a pendant drop. A Hamilton DS 500/GT syringe connected with a Teflon coated needle was used to form the drops with volumes of (9 ± 1) µL. The needle was placed inside an aluminium air chamber able to maintain the temperature control within 0.1 K. The temperature was attained by circulating water in a double-jacketed aluminum cell by means of a Julabo F-25 water bath. This temperature was measured with a Pt100 placed at a distance of approximately 1 cm from the liquid drop. After temperature stabilization, drops were formed and allowed to thermally stabilize for 30 minutes prior to the surface tension determination. To ensure a dry environment, silica gel was placed inside the closed aluminum chamber and the cell was kept sealed during the measurements. For each temperature, at least 2 drops were formed and for each drop, an average of 200 images was captured. The analysis of the drop shape was performed using the software module SCA 20 where the gravitational acceleration (g = 9.8018 m s−2) and latitude (lat = 40°) were used according to the location of the assay. The density values required for the calculation of the surface tensions were measured in this work. Furthermore, details on the equipment and its validity to measure surface tensions of ILs can be found in previous publications.26,27 The high temperature interfacial tension setup: surface tensions of [C12C1im]Cl and [C14C1im][BF4] were determined using the pendant drop method in the 378 to 410 K temperature range. Drops (≈7 µL) were produced using a micrometric syringe (Gilmont Instruments)
connected to a Teflon tube, which was introduced inside a Ramé-Hart chamber (model 100-07-00). At high temperatures, the micrometric syringe was heated using a heating tape. The liquid drops were held between the parallel faces of an aluminium block kept inside the chamber at a constant temperature (±0.05 K) using a PID controller. The temperature of the liquid drop was additionally monitored using a differential copper constantan thermocouple. The ambient chamber was dried using silica gel and the chamber flushed using dry nitrogen. A camera (Al CV-A50) coupled to a Wild M3Z microscope was used to analyze the shape of the pendant drops and their analysis performed using the ADSA-P software (Axisymmetric Drop Shape Analysis, Applied Surface Thermodynamics Research Associates, Toronto, Canada). The estimated overall uncertainty in the temperature measurements is 0.5 K. At least 5 drops were formed at each temperature and 5 images were captured for each drop. Additional details can be found elsewhere.38,29

Interfacial elasticity setup: measurements on the interfacial tension were carried out on a pendant drop tensiometer (Teclis Tracker, IT Concept) using dynamic oscillatory tests at different temperatures (316–326 K). The ILC was placed in a vertical syringe and drops were formed and aged for 5 minutes, creating an ILC–air interface. Sinusoidal deformations around the interfacial area with oscillation frequencies of 0.05, 0.1, and 0.2 Hz were then performed. No frequency influence was observed in the results. The first few drops formed were discarded and the experiments were performed in triplicate. The profile of each drop formed was scanned by a CCD camera and treated by a personal computer. The equipment software calculates the dynamic interfacial tension by fitting the drop profile with the Young–Laplace equation. The total viscoelastic modulus – and its elastic and viscous components – of the film were derived from the dilatational rheological experiments. Due to the high viscosity of the ILC sample, the drop creation and the dynamic tests were performed with rigid control of the drop area and volume and oscillation amplitude, aiming at a greater reproducibility of the results. In addition, the best volume values and drop area which would produce results with lower noise in the determined properties were evaluated.

The total dilatation modulus (or elasticity) gives values of resistance to the creation of an interfacial tension gradient. The interfacial dilatational module, \( \varepsilon \), is defined as the increase of the interfacial tension per unit of increment of surface area (A), being mathematically defined as30

\[
\varepsilon = \frac{d\gamma}{d\ln A}
\]

(1)

where \( \gamma \) is the interfacial tension (mN m\(^{-1}\)) and A is the interfacial surface (mm\(^2\)). When an oscillatory deformation is applied, the elasticity presents an imaginary component (\( \varepsilon'' \)) and a real component (\( \varepsilon' \)) defined by an oscillatory area, where the first term is equal to the elastic contribution and the second is proportional to the viscous contribution:

\[
\varepsilon = \varepsilon' + \varepsilon''i
\]

(2)

Further details on experimental conditions can be found elsewhere.31,32

**Light-polarized optical microscopy**

The melting profile of the ionic liquid crystal \([\text{C}_{12}\text{C}_{1}\text{im}]\text{BF}_4\) was evaluated using an Olympus BX51 light polarized microscope (Olympus Co., Tokyo, Japan) coupled to a Linkam LTS120 temperature-controlled peltier system (Linkam Scientific Instruments, Ltd, Tadworth, U.K.) with a precision of ±0.05 K. The optical textures of \([\text{C}_{14}\text{C}_{1}\text{im}]\text{BF}_4\), \([\text{C}_{12}\text{C}_{1}\text{im}]\text{Cl}\), \([\text{C}_{14}\text{C}_{1}\text{im}]\text{Cl}\), \([\text{C}_{14}\text{C}_{1}\text{im}]\text{Br}\) and \([\text{N}_{111414}]\text{Cl}\) were captured using an optical microscope Leica DM 2700 M with a temperature controller Linkam LTS 420. Samples of around 1 mg were placed in a conclave glass slide, covered with a lamella and sealed with thermal glue in order to avoid the adsorption of water. Different heating and cooling rates were applied, and are described and analyzed in the Results section.

**Ellipsometry**

The spectroscopic ellipsometry measurements were carried out using an AutoSE ellipsometer (HORIBA Scientific) with a total of 250 points in the wavelength interval of 450–850 nm, an incidence angle (\( \theta_i \)) of 69.8° and a signal to noise ratio of 30. A measurement spot area of 250 × 250 \( \mu \)m\(^2\) was used. Three measurements were performed for each temperature. The temperature of the sample was controlled by using a Peltier system. The refractive index values were calculated assuming a three-layered structure model consisting of one layer for the \([\text{C}_{12}\text{C}_{1}\text{im}]\text{BF}_4\) ILC bulk material, a surface layer, and a layer of air as an ambient medium. The data were minimized using the Simplex algorithm, and the dispersion curves were determined using the Cauchy absorbent model, given by:

\[
\eta(\lambda) = A + \frac{B}{\lambda^2} + \frac{C}{\lambda^4}
\]

(3)

where \( n \) is the refractive index, \( \lambda \) is the wavelength (nm), and A, B and C are constants.

**Refractive Index**

Refractive index (\( \theta_0 \)) measurements were performed at 589.3 nm using an automated Abbemat 500 Anton Paar refractometer (maximum temperature deviation: ±0.01 K; maximum refractive index uncertainty: 2 × 10^-5). Measurements were carried out in the temperature range from (308 to 322) K and at atmospheric pressure. The device uses reflected light to measure the refractive index where the sample on the top of the measuring prism is irradiated from different angles by using a light-emitting diode.

**Results and discussion**

On heating, \([\text{C}_{12}\text{C}_{1}\text{im}]\text{BF}_4\) first melts to a viscous smectic A mesophase,19,20–22 which then turns into an isotropic liquid. The solid–liquid crystal–isotropic liquid phase transitions (hereby called melting and clearing points, respectively) of the tetrafluoroborate-based ILC were measured by DSC and these results, presented in Table 1, are in good agreement with the literature.20–22 The clearing point was further confirmed via polarizing optical microscopy (POM). At times, this compound
displayed a reversible crystal–crystal transition at $T_{\text{sn}} = 300.62 \pm 0.34$ K, in the second and/or third heating cycle (see Fig. S1 of the ESI†); a similar thermotropic polymorphic behavior was already reported for 1-alkyl-3-methylimidazolium tetrachloropalladate(II) salts.33

To fully characterize the [C12C1im][BF4] bulk properties, the density, viscosity, and refraction index were measured and are reported in the ESI†. Densities and viscosities of pure [C12C1im][BF4] were measured at several temperatures and at atmospheric pressure, and are listed in Table S1 and presented in Fig. S2 and S3 (ESI†). The experimental refractive indices for pure [C12C1im][BF4] are listed in Table S2 (ESI†). To the best of our knowledge, there are no data available for comparison in the open literature. The densities and refractive indexes display clear transitions at around 320 K which are in agreement with the clearing point measured. Due to the density, there is a clear variation on the isobaric thermal expansion coefficient (Fig. S2, ESI†) corresponding to a change in behavior due to an increase in the disorganization of the liquid phase. The refractive index also presents a sudden decrease as the liquid becomes isotropic. Although the viscosity also presents a significant change around the clearing point it was not possible to measure the viscosity of [C12C1im][BF4] in the liquid crystal phase because it was outside the range of our apparatus.

To study the surface of the [C12C1im][BF4] ILC, measurements of the surface tension of the liquid crystal and isotropic liquid were carried out. Due to the high viscosity of the liquid phase, surface tension measurements were only possible above 317 K, as shown in Fig. 2a. The results reveal a clear change of slope at around 322 K. This change of slope was shown to be reversible. Although this behavior was not reported by Law and Watson,34 upon close inspection of their data for [C12C1im][BF4], an inversion of slope seems to be present at the lowest temperatures, although within the experimental uncertainty of the data.

Information on the surface excess entropy can be taken from the temperature slope of the surface tension9,35 using $\Delta S_{\text{surf}} = d\gamma/dT = -(S_a - S_b)$, where $S_a$ and $S_b$ are the entropies for the surface and the bulk, respectively. A negative temperature slope indicates that the molecules on the surface are less constrained than those in the bulk ($S_a > S_b$). The ordering of the liquid surface results in a surface entropy reduction and thus $S_a < S_b$ and $d\gamma/dT > 0$. A change in slope for the surface tension is thus a strong indicator of an ordering transition at the surface. This behavior, observed here for [C12C1im][BF4], and previously reported for the n-alkanes, liquid metal SDS/water solutions,9,35 and in liquid crystals around the nematic to isotropic transition,36 is a clear indication of the formation of a liquid crystalline10–15,36 or solid layer4,7,9,35 at the surface of the isotropic liquid. First order transitions are characterized by small negative slopes for the liquid surface and large positive ones on the order crystalline surface as observed for alkanes.9 This is translated by the entropy change upon formation of the surface layer, $\Delta S_{\text{fus}} = (\gamma_{\text{surface}} - \gamma_{\text{liquid}})/T$ calculated as the difference between the slope of the surface tension below and above the layer formation temperature. For [C12C1im][BF4], $\Delta S_{\text{fus}}$ has a value of 0.066 mN m$^{-1}$ K$^{-1}$, considerably

### Table 1 Phase transition temperatures and enthalpies of \([C_{12}C_{1}im][BF_4]\) and comparison with literature data

<table>
<thead>
<tr>
<th></th>
<th>Solid → liquid crystal</th>
<th>Liquid crystal → liquid</th>
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<tbody>
<tr>
<td></td>
<td>Heating</td>
<td>Cooling</td>
</tr>
<tr>
<td>$T/K$</td>
<td></td>
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<tr>
<td>302.94 ± 1.87</td>
<td>286.15 ± 1.39</td>
<td>320.20 ± 0.27</td>
</tr>
<tr>
<td>299.55$^a$</td>
<td>280.55$^a$</td>
<td>311.65$^a$</td>
</tr>
<tr>
<td>285.37$^b$</td>
<td>287.33$^b$</td>
<td>323.17$^b$</td>
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<tr>
<td>305.75$^c$</td>
<td>283.25$^c$</td>
<td>322.45$^c$</td>
</tr>
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$^{a}$ Holbrey and Seddon, 1999.

Fig. 2 Temperature dependence of the surface tension, $\gamma$, of [C12C1im][BF4], [C14C1im][BF4] and [C12C1im][Cl] measured using two different setups (description available in the ESI†). The lines are guides for the eyes.
lower than the values of $\Delta S_s$ presented for $n$-alkanes (for example: for hexadecane $n = 16$, and $\Delta S_s = 0.896 \text{ mN m}^{-1} \text{ K}^{-1}$). In addition, it is possible to calculate the entropy change in the bulk liquid-rotator transition or bulk freezing, $\Delta S_b = d\varepsilon/dT\text{rotator} - d\varepsilon/dT\text{liquid}$, which gives information about the similarity between the ordered monolayer and that of the bulk rotator phase. $d\varepsilon/dT\text{rotator}$ can be approximated from latent heat measurements using the melting properties presented in Table 1. Thus, by taking an average area per molecule of $45 \text{ Å}^2$, one can calculate $\Delta S_b = 0.359 \text{ mN m}^{-1} \text{ K}^{-1}$. As expected, the entropy change upon bulk freezing is higher than the entropy change upon surface freezing, given the more organized nature of the anisotropic liquid phase on top of which the surface crystal develops.

Information on the surface excess entropy can be taken from the temperature slope of the surface tension using $d\varepsilon/dT = -(S_s - S_b)$, where $S_s$ and $S_b$ are the entropies for the surface and the bulk, respectively. A negative temperature slope indicates that the molecules on the surface are less constrained than those in the bulk ($S_s > S_b$). The ordering of the liquid surface results in a surface entropy reduction and thus $S_s < S_b$ and $d\varepsilon/dT > 0$. A slope change in the surface tension is thus a strong indicator of an ordering transition at the surface. This behavior, observed here for $[\text{C}_{12}\text{C}_{1}\text{im}][\text{BF}_4]$, and previously reported for the $n$-alkanes, liquid metal SDS/water solutions and in liquid crystals around the nematic to isotropic transition, is a clear indication of the formation of a liquid crystalline or solid layer at the surface of the isotropic liquid. First order transitions are characterized by small negative slopes in the liquid surface and large positive ones in the order crystalline surface as observed for alkanes. This is translated by the entropy change upon formation of the surface layer, $\Delta S_s = d\varepsilon/dT\text{surface} - d\varepsilon/dT\text{liquid}$, calculated as the difference between the slope of the surface tension below and above the layer formation temperature. For $[\text{C}_{12}\text{C}_{1}\text{im}][\text{BF}_4]$, $\Delta S_s$ has a value of 0.066 mN m$^{-1}$ K$^{-1}$, considerably lower than the values of $\Delta S_s$ presented for $n$-alkanes (for example: for hexadecane $n = 16$, $\Delta S_s = 0.896 \text{ mN m}^{-1} \text{ K}^{-1}$). In addition, it is possible to calculate the entropy change in the bulk liquid-rotator transition or bulk freezing, $\Delta S_b = d\varepsilon/dT\text{rotator} - d\varepsilon/dT\text{liquid}$, which gives information about the similarity between the ordered monolayer and that of the bulk rotator phase. $d\varepsilon/dT\text{rotator}$ can be approximated from latent heat measurements using the melting properties presented in Table 1. Thus, by taking an average area per molecule of $45 \text{ Å}^2$, one can calculate $\Delta S_b = 0.359 \text{ mN m}^{-1} \text{ K}^{-1}$. As expected, the entropy change upon bulk freezing is higher than the entropy change upon surface freezing, given the more organized nature of the anisotropic liquid phase on top of which the surface crystal develops.

Having identified the existence of a surface ordering phenomenon at the surface of the smectic $[\text{C}_{12}\text{C}_{1}\text{im}][\text{BF}_4]$ ILC, and two other ILCs, and aiming to further probe the interfacial reorganization, the interfacial elasticity ($\varepsilon$), and its elastic ($\varepsilon'$) and viscous ($\varepsilon''$) moduli were evaluated for $[\text{C}_{12}\text{C}_{1}\text{im}][\text{BF}_4]$ at different temperatures, as depicted in Fig. 3. Above 323 K the values of $\varepsilon$ are close to zero, suggesting no interfacial reorganization during the oscillations and thus, that the system presents the behavior of an isotropic liquid. At lower temperatures an increase in the total interfacial elasticity is observed, indicating that reorganization takes place at the interface. Furthermore, a clear dependence of $\varepsilon$ on temperature suggests a continuous increase in the interfacial activity, with the behavior depicted in Fig. 3b indicating that this interface behavior is almost purely elastic. Despite the high bulk viscosity of the compound at these temperatures, the viscous and elastic moduli are related to the interface features. While the viscous modulus associated to the liquid-like structural characteristics of the interface remains constant and close to zero, the essentially elastic behavior of the surface denotes a solid-like structure, showing that we are in the presence of a surface freezing phenomenon similar to that observed by Pontoni et al.

To further investigate the surface freezing, polarized optical microscopy observations of the phase transitions were carried out, revealing some very interesting textures at a length scale accessible to direct observation, as shown in Fig. 4. On heating it is possible to see the disappearance of the liquid crystals together with some specific structures with a diameter of around 5 μm. These structures decrease in number and size with increasing temperature, but persist after the clarification of the bulk liquid crystal. The same phenomenon is observed during cooling, where an initial appearance of the micrometric structure and its reorganization at the surface of the isotropic
A liquid is observed, followed by the appearance of the bulk liquid crystals (a video is available in the ESI†). The same behavior was also observed for two other ILCs, namely [C_{14}C_{1}im][BF_{4}] and [C_{12}C_{1}im]Cl, for which the surface tension is reported here, along with the surface tension reported for [C_{14}C_{1}im]Cl, [C_{14}C_{1}im]Br and [N_{111414}]Cl as shown in Fig. 5.

Although a more detailed characterization of these structures was not possible with the experimental techniques available, our interpretation of these observations is based on the experiments of Jeon et al.\textsuperscript{18} in the presence of crystalline domains at the surface of an isotropic ionic liquid. Pontoni et al.\textsuperscript{19} distinguish the surface crystallization with a full surface coverage for [C_{20}C_{1}im][NTf_{2}] with vdW-dominated interactions, from the crystalline domains observed on Coulomb-dominated [C_{4}C_{1}im][PF_{6}]. The intermediate chain length of the ILs studied here, combined with the strongly charged anions Cl\textsuperscript{-} and [BF_{4}]\textsuperscript{-}, induce a behavior that bridges the two types of surface freezing on ILs observed previously: from the crystalline domains and partial coverage on top of an isotropic liquid at high temperatures above the clearing point, to a full coverage on top of a liquid crystalline phase at temperatures below the clearing point.

Aiming to further characterize the ordered surface of [C_{12}C_{1}im][BF_{4}] and to support the interpretation of the results reported above, the [C_{12}C_{1}im][BF_{4}] refractive index, dispersion curves and thickness were measured via spectroscopic ellipsometry as a function of temperature (see the ESI† for detailed results in Fig. S4 and S5). As can be seen in Fig. 6, there is a discontinuous transition in the refractive index measured using both refractometry and ellipsometry techniques. This behavior confirms the appearance of an ordered surface. In addition, it was possible to measure the thickness of the surface layer that continuously decreases with increasing temperature. The layer disappears around 323 K, somewhat above the clarification point in agreement with the transitions observed with the other techniques used here. One slope change can be seen at the liquid crystal–liquid phase transition. This behavior previously observed before for LCs was described as a wetting phenomenon.\textsuperscript{11}

In order to obtain molecular level insight into the studied system, the surface of bulk phases of the [C_{12}C_{1}im][BF_{4}] IL, the isotropic liquid and the smectic liquid crystal at different temperatures, was also studied by atomistic molecular dynamic simulations. As will be shown, the simulation results illustrate different possibilities of surface organization, thus contributing to the interpretation of the experimental observations and an overall understanding of the phenomenon.
The CL&P force field was used to model the ionic liquid.\textsuperscript{39} The simulation of the isotropic liquid started from a random low density configuration of 1000 ion pairs and was equilibrated in the \textit{NVT} ensemble in an elongated box with a volume corresponding to a state of liquid–vapour coexistence. The liquid crystalline phase was simulated following the approach proposed and validated by Bruce \textit{et al.}\textsuperscript{40} and extending the simulation box in the direction perpendicular to the smectic layers, in order to explicitly obtain two liquid–vacuum interfaces. The liquid crystalline structure was then frozen by gradually lowering the temperature until a structure very close to the crystalline solid was recovered. The full simulation details are presented in the ESL\textsuperscript{†}

The interfacial structures were analysed by obtaining the atomic density profiles, in the direction normal to the liquid–vacuum interface, from the simulation trajectories. The density profiles for the anion, the charged part of the cation and the terminal group of the cation side chain are shown in Fig. 7, together with simulation snapshots of the corresponding structures.

As seen in Fig. 7 (middle), the liquid crystalline smectic phase displays a well-defined layered structure formed by distinct ionic bilayers. Within each layer, the anions and the charged part of the cations alternate in a plane, forming a checkerboard distribution. The ionic bilayers alternate forming layers of dense, interdigitated, alkyl side-chains.

On cooling, as could be expected, the system clearly becomes denser and more organized (Fig. 7 (top)). The profile peaks become slightly narrower, reflecting the lower mobility of the particles and higher density. The interdigitated alkyl side-chains adopt a tilted conformation and distinct packing planes become visible. The structure is believed to be very close to a crystalline solid, although the profile peaks are slightly broader than those of a typical solid.

In the isotropic liquid phase (Fig. 7 (bottom)), the ionic bilayered fine structure of the liquid crystal is completely lost. Single peaks now appear in the density profiles of both the charge groups and alkyl side-chains, revealing a fully liquid-like organization. The alkyl side-chains lost the previous recognizable interdigitated structure.

Regarding the surfaces, the simulation results are again able to illustrate the different possibilities of organization, providing reasonable models to rationalize the experimental observations.

The isotropic liquid surface shows a layered structure that decays with the distance from the interface (and also with temperature). It is often argued\textsuperscript{7} that this “conventional” surface layering, which increases at lower temperatures, gradually propagates to the bulk of the liquid. At the clearing temperature the whole system will be liquid-crystalline. It is also apparent that the alkyl side-chains, as expected preferentially oriented towards the vacuum, are relatively disorganized. This results from the low density of side-chains at the interface that precludes an effective close-packing of the chains and an efficient coverage of the charged moieties.

The surface of the liquid crystalline mesophase is clearly more organized than that of the isotropic liquid and is compatible with a liquid crystal surface phase.\textsuperscript{19} This already displays

![Fig. 7](image-url) Simulation snapshots and surface normal atomic density profiles for the solid (top), liquid crystal (middle) and isotropic liquid (bottom) surfaces. Profiles: dark blue = charged part of cation (C atom between the N atoms), red = anion (B atom), light blue = terminal methyl carbon of the C12 chain. Snapshots: charged parts are represented by blue (cation) and red (anion) van der Waals spheres; the C12 side chain is represented by light blue and white wireframe.
the characteristic checkerboard plane configuration of the ionic parts. However, the alkyl side chains are still relatively disorganized.

On cooling, it is clear from Fig. 7 (bottom) and 8 that a highly organized structure is formed at the surface. The alkyl chains now adopt a highly tilted configuration that simultaneously allows close packing of the chains and an efficient coverage of the surface. This structure is compatible with a solid crystalline surface and in our opinion is a good model for the solid-like, highly organized surface evidenced by the experimental results. The simulation results also indicate that the formation of a stable organized solid-like structure at the surface of an ILC does not require the presence of an impurity particle to increase the density of chains (Fig. 8).

Conclusions

This work studied the surface of \([\text{C}_{12}\text{C}_{1}\text{im}]\text{[BF}_4]\) at temperatures from the melting point to above the clearing point. This compound was chosen due to its low melting and clearing temperatures and large and stable mesophase. The nature of the interactions present when compared with the ILS previously reported to show surface freezing allows it to link the two extreme types of surface behavior reported: formation of crystal domains in coulombic dominated ILS or full coverage for long chain, van der Waals dominated ILS. The experimental techniques used here support the presence of crystalline domains at the surface of the isotropic liquid, fully covering the surface of the liquid crystalline mesophase. The solid nature of this layer, demonstrated here by the elastic behavior revealed by the surface rheology, is in agreement with the results obtained by using the X-ray method by Pontoni et al. Surface tension measurements and optical microscopy suggest that this behaviour is a prevalent phenomenon on this type of compound that would deserve further investigation and characterization. MD simulation results illustrate the different possibilities of organization, providing reasonable models to rationalize the experimental observations.

Conflicts of interest

There are no conflicts to declare.

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