Squeezout phenomena and boundary layer formation of a model ionic liquid under confinement and charging

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Electrical charging of parallel plates confining a model ionic liquid down to nanoscale distances yields a variety of charge-induced changes in the structural features of the confined film. That includes even-odd switching of the structural layering and charging-induced solidification and melting, with important changes of local ordering between and within layers, and of squeezout behavior. By means of molecular dynamics simulations, we explore this variety of phenomena in the simplest charged Lennard-Jones coarse-grained model including or excluding the effect a neutral tail giving an anisotropic shape to one of the model ions. Using these models and open conditions permitting the flow of ions in and out of the interplate gap, we simulate the liquid squeezout to obtain the distance dependent structure and forces between the plates during their adiabatic approach under load. Simulations at fixed applied force illustrate an effective electrical pumping of the ionic liquid, from a thick nearly solid film that withstands the interplate pressure for high plate charge to complete squeezout following melting near zero charge. Effective enthalpy curves obtained by integration of interplate forces versus distance show the local minima that correspond to layering and predict the switching between one minimum and another under squeezing and charging. © 2015 AIP Publishing LLC. [http://dx.doi.org/10.1063/1.4907747]

I. INTRODUCTION

Ionic liquids (ILs), ionic salts based on organic molecules whose large size, amphiphilicity and anion-cation asymmetry generally yield melting below room temperature, encompass an enormous chemical variety,1 interesting physical properties,2 and numerous applications.3 The structural and electrical properties of ILs near solid interfaces have been investigated experimentally4,5 and theoretically.6,7 Data show that under force-induced boundary confinement, the liquid undergoes strong layering.8-11 Squeezout takes place for increasing force in bilayer steps, so that ions of opposite sign exit together, keeping overall neutrality. The electrical charge of the lubricant molecules and their layering under confinement naturally suggest the possibility to influence the layering, and the squeezout behavior, by means of plate charging or by otherwise applied fields. It has recently been shown, by AFM force spectroscopy measurements and MD simulations,12 that when a voltage is applied between a graphite substrate and a tip, the interfacial boundary layer may undergo an orientation transition. For what concerns nanotribology, the charging of a gold surface in an IL filled narrow AFM gap indeed showed important frictional changes as a function of the applied voltage,9 while different effects of charging have been reported in other experiments with different ILs.11 That confirms on one hand that charging is indeed an important parameter influencing structure, squeezout, and friction, and on the other hand that the details of these charging effects generally depend upon the specific nature of the IL and of plates. It seems at this point important from a more general viewpoint to conduct a wider theoretical and simulation inquiry, exploring possible charging induced phenomena based on less specific, more generic, reasonably simple models. These models of course do not replace the much more specific systems currently being simulated to describe specific ILs and their confined behavior.13,14 Simplicity of the model on the other hand is essential to allow a wide-angle view of some among the possible squeezout effects and scenarios provided by charging, either static or time-dependent, of the confining surfaces.

Within this broad exploratory program, and without aiming at modeling a specific case, we investigate first of all the scenario of ILs confined between two neutrals or nearly neutral, then charged, solid plates. This study encompasses a variety of charging-related static phenomena, preparatory to the dynamical study of frictional behaviour which will be considered in subsequent work.

As in the early approach of Fedorov and Kornyshev, we begin with a simple charged Lennard-Jones (LJ) system, but then we extend it by the minimal addition of a neutral “tail” making one of the ions considerably asymmetric, as is common in many real IL ions. Using an open geometry, where ions can escape or re-enter the interplate gap during slow, adiabatic...
changes of the interplate gap width \( D \), we observe in the simulation the formation of partly ordered layers under squeezing. In analogy to experiments, partially ordered layers have been seen in experiments. \(^5\)\(^6\) Squeezout is found to take place by successive transitions through plateaus at well defined interplate distances, through force-induced gap changes corresponding to one IL bilayer, clearly the neutral entity here. This transverse layering structure is accompanied by some degree of lateral, planar ordering inside each IL molecular layer. Planar order, even if probably overemphasized by our simple models relative to much more complex real ILs (where in-plane spatial order is never observed experimentally) is nonetheless a relevant feature that helps understanding both structure and dynamical behaviors. We characterize planar ordering of our confined IL by means of a k-space resolved structure factor, analogous to a two-dimensional, z-resolved diffraction scattering amplitude. Some level of frustrated plate- and confinement-induced 3D crystallization is also found, whose details are, as we show, model-dependent. We extract by means of force-distance integration a very instructive effective interplate interaction free energy, where the different plateaus appear as local minima, so that the squeezing transitions between them is described and in fact predicted by a Maxwell construction. The effect of charging is subsequently shown to cause structural changes which can in turn be related to changes of the interplate equilibrium free energy characteristics.

II. MODEL AND SIMULATION GEOMETRY

ILs are molten salts usually made up of large-size anions and cations, generally organic, and with asymmetric irregular shapes often including long alkyl chains. The irregularity is important as it gives the molecule a larger gyration radius in the liquid state, while effectively preventing low temperature order and crystallization by replacing it with a glassy state. Dropping most of these complications, we restrict to a much simpler model, crudely including just the rough features of a generic IL. We will therefore build on the basic model already exploited in previous studies, namely a charged LJ system where anions and cations have different radii. We compare this model with its extension, obtained by attaching a neutral “tail” to the cation. In the extended model, the anion consists of a negatively charged, large-sized spherical LJ classical particle, the cation of a dimer made of a positively charged, small-sized LJ head, rigidly bound to an equal size, and neutral particle, the tail (see Fig. 1). A similar model with anisotropic ions has been used for the study of interfacial structure of ionic liquids at charged interfaces, providing insights into the nature of the camel shape of the voltage dependence of the double layer capacitance. The tail simultaneously achieves several goals. First, it enhances the liquid gyration radius, the solid-liquid density jump, and the general tendency to form a glass rather than a crystal at low temperatures. Second, it improves the plate wetting behavior, which without tails is unrealistically weak, while the wetting by real IL of surfaces such as mica is only partial. Partial wetting is plausibly associated, at least in some cases, with a first monolayer of cations whose outward pointing tails may provide a “phobic” gap against further layer-by-layer film growth. That is realized in our “tailed” model (TM) but not in the simpler charged LJ “salt-like” (SM) model.

Finally, the tail also introduces rotational entropy effects that must be present in real systems, but are missing in simple SMs.\(^1\)\(^5\)\(^6\)\(^7\)\(^8\) Building on the work of Fedorov and Kornyshev,\(^1\)\(^5\) we thus assume the anion (A), the positive head of the cation (C), and plate “atoms” (P) to interact electrostatically and via a LJ potential,

\[
V(r_{ij}) = 4\epsilon_{ij}\left[\left(\frac{\sigma_{ij}}{r_{ij}}\right)^{12} - \left(\frac{\sigma_{ij}}{r_{ij}}\right)^{6}\right] + \frac{e_i e_j}{4\pi \epsilon_0 \sigma_{r_{ij}}}
\]

with \( i, j = A, C, P \). The tails (T) interact with all other species through a repulsive potential,

\[
V^{T}(r_{iT}) = 4\epsilon_{iT}\left[\left(\frac{\sigma_{iT}}{r_{iT}}\right)^{12}\right]
\]

with \( i = A, C, P, T \). The full list of parameters is reported in Table I. We adopt the LJ potential strengths \( \epsilon_{AP} = \epsilon_{CP} = K_{B}T_{room}/2 = 13.3 \) meV as our energy unit and take ion-plate radii slightly smaller than those between ions in order to provide a reasonable wetting habit of the plates. The SM consists of the same model liquid simply without the tail, as used in Ref. 15. To account for the dielectric screening of real systems, the two ionic species of charge \( +e \) and \( -e \) the

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**Table I. List of LJ parameters used in the simulations.**

| \( \sigma \) (nm) | \( \epsilon \) (meV) |
|-------------------|---------------------|
| AA                | 1                   |
| CC                | 0.5                 |
| AC                | 0.75                |
| AP                | 0.7                 |
| CP                | 0.35                |
| TT                | 0.5                 |
| TA                | 0.75                |
| TC                | 0.5                 |
| TP                | 0.35                |

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electron charge) are immersed in a uniform average dielectric with \( \varepsilon_r = 2.0 \).

The liquid is confined between two negatively charged plates with a modest surface charge density \( q = -4 \, \mu C/cm^2 \), unless differently specified, chosen to break the charge symmetry. The plates, somewhat similar to mica surfaces used in Surface Force Apparatus (SFA) experiments, are made of LJ sites arranged in a rigid close-packed triangular lattice with spacing 0.52 nm.

All molecular dynamics (MD) simulations were performed using the LAMMPS code.\(^{21}\) Canonical ensemble configurations were sampled by means of a Langevin thermostat directly applied to the lubricant molecules. As in previous MD simulations,\(^ {22}\) under similar operative conditions, the details of the adopted dissipation scheme are not expected to change the essence of the system response. The plates were treated as rigid bodies, the lower one fixed and the upper one either subjected to a \( z \)-directed force \( F_n \) (the load) as shown in Fig. 1, or else driven along \( z \), inwards or outwards, at some very small constant velocity. While the total number of simulated ions is constant, the finite plate width along \( x \) ending with free vacuum on both sides in the simulation cell geometry (Fig. 1), permits, unlike other IL simulations, otherwise very realistic,\(^ {13,14}\) particle squeezeout with formation of two lateral ionic liquid drops. The drops serve as liquid reservoirs so that the number of ions effectively confined inside the gap can dynamically change depending on the loading conditions, realizing an effectively grand-canonical situation.

III. BULK MELTING AND SOLIDIFICATION

A basic property of an IL is its melting habit. We investigate and compare the bulk melting temperature of the TM and SM models by performing a slow-varying temperature loop MD simulation of both models with \( N = 2000 \) ions and full 3D periodic boundary conditions. Starting from a high temperature \( T_m \), where both SM and TM are in the liquid state, we first decrease the temperature to a much lower \( T_m \) with a rate of 1.7 K/ns, producing solidification; then increase it again back to \( T_m \), to produce melting. Fig. 2(a) shows the internal energy, \( E_{int} \) of the SM in the temperature loop. On cooling, \( E_{int} \) linearly decreases until a sharp drop at \( T_1 = 195 \) K corresponding to a liquid-solid transition. On heating back the solid, a sharp rise of \( E_{int} \) is found at \( T_2 = 285 \) K, corresponding to the reverse solid-liquid transformation. The large magnitude of jumps reflects strong crystalline order in the SM solid. The equilibrium melting temperature is to be found between those of internal energy drop and rise, whose difference reflects the hysteresis normally encountered in the absence of nucleation centers, such as surfaces. The TM behaves similarly, but displays smoother transitions, reflecting a much poorer ordering in the solid, and lower transition temperatures \( T_1 = 127.5 \) K and \( T_2 = 165 \) K, as shown in Fig. 2(b). In both SM and strong and TM, these transitions are accompanied by correspondingly strong changes of diffusion coefficient (not shown). If we assume arbitrarily that the melting temperature is the average between \( T_1 \) and \( T_2 \), we obtain \( T_m = 150 \pm 20 \) K for TM and \( T_m = 240 \pm 40 \) K for SM, respectively. The decreased density of TM relative to SM reduces the cohesive energy and thus the melting temperature. The tails in TM also prevent crystallization in the rocksalt structure typical of SM as shown in Fig. 3. Unless otherwise specified, all simulations reported in the following were done at temperature \( T_{TM} = 225 \) K and \( T_{SM} = 300 \) K for the TM and SM, respectively, safely above not too far above the bulk melting temperatures, as in real experimental cases. In our simulations, therefore, both ILs are in a fully liquid state as long as the gap width between the two plates is sufficiently large. Both ILs may solidify, as we will show, when the confining gap between plates is reduced down to the molecular thicknesses typical of boundary lubrication conditions.

![FIG. 2. Bulk internal energy \( E_{int} \) of the SM (upper panel) and TM (lower panel) liquids as a function of simulation time, during a linear back and forth temperature ramping of rate 1.7 K/ns.](image)

![FIG. 3. Snapshot of (a) SM and (b) TM bulk solids below their respective melting temperatures. The presence of tails in TM prevents crystallization in the rocksalt structure as in SM.](image)
IV. WETTING PROPERTIES

The wetting of plate surfaces such as mica is known to be partial by at least some ILs. Squeezout phenomena and layering necessarily involve intimate features of the liquid-solid plate interface, not disconnected with those determining the plate wettability. Thus, it is important to monitor the plate wetting habit of our IL models. To do that, we simulate for both models the behaviour of a liquid droplet consisting of $N = 2000$ ions deposited on a neutral and then on a lightly charged plate. The adhesion parameters of the liquids should be chosen in a reasonable way and in order to reproduce qualitatively the wetting properties. We find (Fig. 4) that the TM wetting of the neutral plate is poor, forming a wetting angle $\theta > 90^\circ$. On the contrary, the SM completely wets the substrate. Only in the TM, the cations provide outwards pointing tails, which give rise to a “phobic” gap, responsible for the incomplete wetting, see Fig. 4(a). Upon charging of the plates, one expects wetting to be generally enhanced for either sign of charge, over the neutral case. For moderately negative charging, we find that the TM liquid indeed wets better than in the neutral case, yielding a smaller wetting angle $\theta \simeq 30^\circ$. The charging evidently reduces the effect of tails, and their effective screening of electrostatic forces. For a moderately positive charge on the other hand, the TM wetting angle $\theta$ is slightly larger than negatively charged substrate. This asymmetry is due to the different sizes and shapes of ions.

These TM results—monolayer coating, partial wetting by the drop—are in qualitative agreement with experimental data for 1-butyl-3-methyl-imidazolium bis-(triuoromethyl sulfonyl)imide, [BMIM] [TFSI], on mica, which show a precursor layer forming ahead of a macroscopic drop with $\theta \approx 30^\circ$. This partial wetting scenario is absent and replaced by complete wetting in the SM. Thus, in such simplified modeling, the introduction of neutral repulsive tails, mimicking, e.g., alkyl chains in real IL systems, turns to be important for the correct wetting properties of the IL. It is highly reasonable to presume that this important element of realism will be important in future studies of sliding friction.

V. SQUEEZOUT SIMULATIONS AND LAYERING

Liquids close to hard surfaces display, both experimentally and theoretically, spatial structuring phenomena with respect to their uniform bulk density. When a liquid film is confined in a planar gap between hard plates a few molecular diameters wide, it develops quite generally a layered density profile, with increasing solid-like properties that eventually enable it to resist squeezout and to support static friction. The squeezout of these layers one at a time under increasing load is well studied theoretically and experimentally. Similar to common liquids, ILs generally give rise to partially ordered molecular-size layers and enhanced viscosity at interfaces with solid surfaces such as mica, silica, and gold. For reasons of charge neutrality and compatibility with ion asymmetry, ILs organize themselves in alternating positive and negative layers not too far from the plates. The squeezout between approaching plates occurs in this case by neutral entities, that is by pairs of layers, as seen in all experiments, as also described by a recent theoretical model. Depending on the specific IL, its melting temperature and its relationship to the confining surfaces, experimental squeezouts by sharp AFM tips may require relatively large applied forces from some nN to several tens of nN, which for an area in the order of 1-10 nm$^2$ gives an indication of the strength of interactions involved.

The available simulations of confined ILs are carried out under unrealistic sealed conditions, via periodic boundary constraints, which do not permit squeezout. Experimental SFA and AFM setups constitute eminently open geometries, allowing dynamical liquid squeezout and suck-in. For our simulations, we therefore adopt an open geometry, albeit a very simple and schematic one. As shown in Fig. 1, it consists of two rigid plates that are infinite along $x$, but finite along $y$, with the IL filling the gap between them. When the plates are moved closer to one another, the liquid is squeezed out of the gap, flowing sideways to form two side drops that constitute IL reservoirs. A simulation geometry similar to ours was used by Landman’s group, whereas a different open geometry allowing for squeezout was adopted by Tartaglino et al. Open simulations were also recently used to investigate squeezout and phase transition of an argon film between two solid surfaces. In our initial protocol, the plates were very slowly approached, and the resulting average force on the...
top plate, \( F \), was monitored as a function of the diminishing inter-plate distance \( D \). Beginning, for example, with a TM bulk liquid confined between plates and closing gradually the gap width \( D \), \( F \) remains initially zero while the liquid is readily expelled, until a critical distance is reached between plates, \( D_c \simeq 4 \) nm, amounting to a few bilayers, where the force-displacement curve departs significantly from zero and begins to grow. The growing force \( F(D) \) between the plates shows layering oscillations as in Fig. 5. Each force dip corresponds to the complete squeeze-out of a pair of layers, one positive and one negative, as in experiment\(^5\) and simulations.\(^7\) If the plate motion is inverted, now increasing the gap width with time, the squeezeout is replaced by "suck-in" of the IL. The black and the blue force curves of squeezeout and suck-in are not identical at a given distance and form a narrow hysteresis loop. The area enclosed in the force-displacement loop measures the dissipation work implied by our plate to-and-from motion, exerted at small but infinitesimal speed. Where the plate speed tends to zero (quasi-static limit), dissipation would vanish, and the two force-distance curves would collapse onto a single adiabatic force \( F(D) \) at all finite temperatures. In these conditions, the adiabatic work \( W(D) = \int_{D_0}^{D} F(z)dz \) provides an accurate measure of the interaction free energy between the plates. Layering oscillations of this distance-dependent free energy was earlier demonstrated in simulations of simple liquids.\(^{28}\)

Changing from fixed distance to fixed force, where a normal load, \( F_n \), is applied to the top plate, and the interplate distance \( D \) is observed to vary as a consequence, these free energy curves establish a relation between \( F_n \) and the consequent number of confined layers. For example, when \( F_n > 3.5 \) nN, only two metastable states with 3 or 5 layers are allowed, while at \( F_n \approx 0 \) all states are allowed. That concept was recently underlined by Hoth et al.\(^{36}\)

The number of confined layers between identical plates is odd, as is demanded by symmetry. Confinement of ILs between mica plates offers a direct example of odd layering\(^{10}\) due to the symmetrically slightly negative charge of both plates, each of which attracts a similar coating of cations. The "transverse" density profile measured along the vertical, plate-to-plate \( z \) axis shows a clear oscillatory order with cations and anions populating different, well separated layers. In the TM in addition, the light brown line represents the density of tails (Figs. 6(a)–6(c)). The positions of cation and anion peaks closest to the plates (vertical blue dashed lines), are geometrically related to their radii. As the number of layers increase, all oscillations broaden and weaken, more so in the TM due to the disordering effect of tails. The SM liquid presents a qualitatively similar picture, where the absence of tails and related disorder allows much more pronounced layering order, occurring at a larger distance between the plates. The force needed to squeeze out a pair of SM layers is much larger, and the peaks of density profile along vertical axis more pronounced, as shown in Figs. 6(e)–6(g).

The bottom panels in Figs. 6(d) and 6(h) display, in the case of seven layers, the number of charges per layer, \( n_l = Q_{l}/e \), with \( Q_l \) total charge per layer and \( e \) electron charge. The total amount of charge, including the plates and the confined liquid, preserves electroneutrality. The TM shows a clear overscreening, with the charge of layers in contact with plates substantially larger than far away. Due to confinement, the SM liquid shows a much readier tendency to solidify. The charge plateau in Fig. 6(b) indicates precisely a crystalline region where a well defined number of ions per layer is required for a rocksalt structure with (111) planes parallel to the plates.

![FIG. 5. Force \( F \) as a function of distance \( D \) between plates. Black and blue curves correspond to downward and upward motion of the top plate, respectively. The force difference is due to the small but finite plate speed \( \nu_z = 0.11 \) m/s, and their average represents our best approximation to the adiabatic force \( F(D) \). The red curve is the free energy \( W(D) \) obtained by integration of the adiabatic force. Numbers indicated in figure denote the number of IL layers confined between the plates. The charge on both plates is \( q = -4\mu C/cm^2 \), a modest value compared with that of, e.g., mica.\(^8\)](image)

![FIG. 6. Density profile along \( z \) axis for the TM (left) and SM liquid (right) and increasing numbers of confined layers. Vertical blue dashed lines indicate the position of the plates at separation \( D \), light brown peaks in panels (a), (b), (c) show the density of tails. Black squares in panels (f) and (b) mark the single layers that are analyzed in Figs. 8(a) and 8(b). Lowest panels: number of charges per layer, \( n_l = Q_{l}/e \), with \( Q_l \) total charge per layer and \( e \) electron charge. The full circles at the edges of panels indicate the charge assumed on the plates.](image)
VI. LAYER-RESOLVED PLANAR STRUCTURE, 2D STRUCTURE FACTORS; 3D STRUCTURES, CRYSTALLIZATION PHENOMENA

The simulation temperatures, substantially but not excessively above the bulk melting temperature, \((T_{TM} = 225 \text{ K} \quad \text{and} \quad T_{SM} = 300 \text{ K})\) for TM and SM, respectively) preserve a reasonably large amount of short range order in the IL. That short range order naturally emerges near hard plates and yields both transverse and planar static orders in the layered structure of the liquid confined between the plates. That order is associated with some partial solidification, and indeed when the gap width \(D\) decreases below a distance corresponding to 10 molecular diameters, both SM and TM layered ILs increasingly resist squeezeout. Accompanying the layering, there must be some amount of planar ordering. Contrary to layering, well known and clearly reflected by squeezeout experiments, the planar order is so far experimentally undetected. Figs. 7(a) and 7(b) show top views of SM and TM liquids for \(D\) corresponding to five confined layers, revealing the nature and extent of planar order. The effect of tails on planar order is quite strong.

For a more precise characterization of planar order, we calculated the z-resolved structure factor \(S_{z0}(q)\) per layer, defined as

\[
S_{z0}(q) = \frac{1}{N_{z0}} \langle \sum_{j,k,z \in z_0} e^{-i\mathbf{q} \cdot (\mathbf{r}_j - \mathbf{r}_k)} \rangle,
\]

where \(\mathbf{r}\) is the \((x, y)\) position of particles in the confinement with \(z\) coordinate in a window close to \(z_0\), \(N_{z0}\) is the number of particles in that window, and \(\langle \ldots \rangle\) indicates time average.

The z-resolved \(S_{z0}(q)\) of the window enclosed in the black rectangle in Fig. 6(f) shows that anions approximately form a 2D triangular lattice, indicating that the SM liquid crystallizes in a rocksalt structure with a \((111)\) plane in contact with the plate (see Fig. 8(a)). The confined TM film behaves in this respect quite differently from SM. The layer structure factor \(S_{z0}(q)\) of the TM anion layer enclosed in the square of Fig. 6(b) displays a liquid-like structure factor, caused by random orientation of wall-like structures (see Fig. 8(b)). Away from the plates, the layering is less pronounced, and there the TM liquid shows a tendency to appear organized with a 2D triangular lattice symmetry. Fig. 7(c) shows a side view of the single vertical wall indicated by the red square in Fig. 7(b), where tails have been removed from the picture for clarity. The \((100)\) plane of rocksalt is easily recognized. Because of the random
orientation of tails, the liquid cannot solidify in a complete three-dimensional crystal, but it manages to arrange in vertical neutral rocksalt walls with tails segregated in the spaces between adjacent walls and oriented parallel to the plates.

These two examples suggest some more general understanding that may be valid beyond the two specific cases. First, plate-induced layering and charge neutrality are universal, as was to be expected. Second, the detailed nature of planar and of three-dimensional ordering, ranging from nearly crystalline to glassy, is variable and much more dependent upon the specific nature of the IL constituents.

VII. SYMMETRIC PLATE CHARGING

The properties of plate-confined ILs described so far for neutral or nearly neutral plates are strongly disturbed once the plates are electrically charged. We study first the structural and squeezeout behaviors under symmetrically charged plates, that is both plates with same charge density \( q \). In order to preserve the overall system neutrality in simulation, we simultaneously remove from the liquid a number of ions amounting to the compensating charge \( -2q \). Upon increasing negative \( q \) on both plates, the confined IL acquired a more solid-like structure, which is mechanically stronger and can bear higher loads before being squeezed out. Fig. 9(a) shows a force-distance curve for the TM liquid, with a charge density of \( q = -10 \mu\text{C/cm}^2 \) on both plates. The force-distance curve shows much higher peaks than for \( q = -4 \mu\text{C/cm}^2 \) as in Fig. 5, signaling the larger force needed for squeezeout from \( N \) to \( N - 2 \) layers.

The SM has a very similar behavior, with even higher peaks in the force-distance curve (not shown). Fig. 10(a) shows the density profile along the \( z \) direction for five TM confined layers between negative plates. Cations crowd up near the negatively charged plates, their tail protruding away from the plates. For totally neutral plates, however, in presence of the sole adhesion forces between ions and plates, the structure of the two model liquids differs rather drastically. The force-distance curve for the more realistic TM liquid displays just two peaks and no evidence for extended layering (see Fig. 9(b)).

With neutral plates, the IL cannot support the 1 nN load down to a gap width of \( D \approx 3 \text{ nm} \). A qualitatively similar picture also emerged in recent experimental investigations showing how surface charges affect the molecular structure and flow properties of ILs. The transverse density profile for \( q = 0 \) and \( D \approx 3 \text{ nm} \) is shown in Fig. 10(b) with formation of just two neutral layers (which we denote as 2n) where both anions and cations coexist in a single monolayer, which is bound to the plate by short-range adhesion. The cation tails point outward and form a weakly repulsive coating. Vertical dashed lines indicate the position of plates. (c) Structure factor \( S_2(q) \) for the layer enclosed in the red rectangle, showing that ions form a rocksalt (100) square monolayer.

FIG. 9. (a) Force-distance curve for the TM liquid and a charge density of \( q = -10 \mu\text{C/cm}^2 \) on both plates. (b) Force-distance curves with neutral plates for the TM (black) and SM (blue) liquid.

FIG. 10. (a) Density profile between plates for the TM liquid forming a stable five layer film between plates charged with \( q = -10 \mu\text{C/cm}^2 \), pressed together with a force \( F = 7 \text{ nN} \). (b) Density profile for the same system under the same force, but now with neutral plates. Cations and anions form two neutral layers, bound to the plates by short-range adhesion. The cation tails point outward and form a weakly repulsive coating. Vertical dashed lines indicate the position of plates. (c) Structure factor \( S_2(q) \) for the layer enclosed in the red rectangle, showing that ions form a rocksalt (100) square monolayer.
adhesive forces. In our case, the sole adhesion to neutral plates binds both cations and anions to the plate, so that they arrange in a rocksalt (100) monolayer with tails pointing outward. A negative plate instead selectively binds cations but not anions, giving rise to an alternation of positive and negative layers with tails parallel to the plates. A pictorial representation of these gives rise to an alternation of positive and negative layers with tails parallel to the plates. A pictorial representation of these structures, so does the electric electro-pumping of ions in and out of the gap, $N_{\text{part}}$, grow back to four layers with reversed charge order. Although four layers are not, as we shall show later, the lowest free enthalpy state, which are instead six layers, the metastable four layer state is nearly solid. In that state, the kinetics becomes very slow, so that the time needed to suck back in another bilayer becomes much longer than the simulation time. As shown in Figs. 12(a) and 12(b), the charging driven inter-plate breathing cycles continue periodically, and so does the effective electro-pumping of ions in and out of the gap. This kind of charging dependent phenomena based

![Diagram of charged plate arrangements](image)

FIG. 11. (a) Side view of TM liquid in the state $2n$ with neutral plates. Only one layer is shown for clarity. (b) Top view of the single layer in panel (a) showing its square arrangement of oppositely charged ions. (c) Sketch of plate charging induced structural transition from a square to a wall-like arrangement.

there is enough adiabaticity to ensure that the effects of charge transport and heat dissipation are safely taken care of. Still, as we shall see, some notable non-adiabatic effects persist, connected with charge-induced solidification. By keeping the load constant at $F_n = 1 \text{ nN}$, the charging period $\tau = 72 \text{ ns}$ and magnitude $Q = 10 \mu C/cm^2$, we simulate the spontaneous evolution of the confined IL and obtain the results shown in Fig. 12.

Symmetry demands that the equilibrium layering structure for antisymmetric plate charging should occur with an even number of players, whereas with symmetric charging the layer number was odd. The simulation begins with a charge density of $|q(t = 0)| = 10 \mu C/cm^2$ on plates and the IL structured in six layers, which is the stable configuration under the fixed applied load force $F_n = 1 \text{ nN}$. As the plate charge slowly drops in time, the IL structure gradually changes and softens. Eventually, the IL becomes soft enough and squeezout suddenly takes place, despite the constant load. The interplate distance $D$ drops, the IL film structure reaching a single, charge neutral two-component layer branded $1\text{n}$, (see inset of Figs. 12(a) and 12(c)) when $q = 0$. That $1\text{n}$ structure is a history dependent state that forms asymmetrically depending upon the initial sign of the charge. As the charging grows again now with inverted sign, the IL is sucked back in and both the gap width $D$ and the number of confined particles in the gap, $N_{\text{part}}$, grow back to four layers with reversed charge order.

Noteworthy as they may be, these planar structure features seem presently inaccessible to experiments, and the development of techniques that could allow their study would be very interesting. Voltage-controlled phase transitions have been proven at electrode-electrolyte interfaces and investigated in recent MD simulations of ionic liquids, showing transitions from multilayer ion structure to ordered monolayer structures, with an ordering in the innermost ion layers dependent on the surface charge density.

VIII. ANTISYMMETRIC, TIME-DEPENDENT PLATE CHARGING

In this section, we finally study how the order in the TM liquid is affected by opposite charging of the two plates. The charging is given a slow sinusoidal variation with time, $q(t) = \pm Q \sin(2\pi t/\tau)$ such as would be caused by externally driven plate charging with a sufficiently low AC frequency to ensure that all the ionic motion caused inside the interplate gap occurs on a much faster time scale. In this respect, we have checked

![Diagram of particle arrangement](image)

FIG. 12. (a) Spontaneous change of interplate distance $D$ between plates as a function of time for a fixed interplate force $F_n = 1 \text{ nN}$ and a slowly oscillating charge with amplitude $Q = 10 \mu C/cm^2$ and period $\tau = 72 \text{ ns}$ (panel (c)). (b) Number of particles in the interplate gap, $N_{\text{part}}$, as a function of time, under variable charging. The time dependent charging causes the plate separation $D$ to open up at maxima (IL suck-in) and to collapse at minima, (IL squeezout) with a strong electropumping action. The inset in panel (a) shows a side view of the confined liquid structure for 4 layers ($q = 10 \mu C/cm^2$) and for a single neutral layer $1\text{n}$ ($q = 0$).
on spontaneous sucking in of the liquid, solidification at large charge with buildup of a solid-like resistance to squeezeout, followed by sudden squeezeout associated with melting of the confined layers upon charge reversal is very likely a more general feature that could occur in a larger class of ILs beyond our simple model. The charging induced transitions just observed in simulation can actually be predicted by the free enthalpy of the system, $W(D) + F_n D$, where $W(D)$ is calculated the same way as earlier by integrating the interplate force from infinity to the gap width $D$. Fig. 13(a) shows the free enthalpy curves so obtained as a function of $D$ for three values of the plate charge density $\pm q$, starting with six layers which is the lowest free enthalpy configuration at a charge density $|q| = 10 \mu C/cm^2$. Calculated curves show that upon decreasing $|q|$ the energy barriers between states at different numbers of layers decrease gradually, eventually pushing the metastable six-layer state leftward to a much smaller distance $D$ between plates. A dramatic squeezeout event takes place near $q \approx 0$ (see Fig. 12(a)), with a jump to the single layer state $1n$. The successive reversal and increase of plate charge provokes the opposite transition at $q \gtrsim 3|\mu C/cm^2$ where, as described in the previous section, the IL is sucked in, and structures up into four layers. As the charge rises again to $|q| = 10 \mu C/cm^2$, the IL never returns to the equilibrium six layer state, indicating that the thermal fluctuations and the simulation time are not sufficient to negotiate the four-to-six-layer free enthalpy barrier (see Fig. 13(a) black curve). The nearly solid four layer state effectively resists the insertion of the last bilayer and blocks the system in a metastable state which is very long-lived, at least on our simulation time scale.

The six-layer equilibrium state should of course be recovered in a sufficiently slow charge dynamics. Similarly to the TM, the SM IL shows even more dramatic charge induced squeezeouts, followed by successive relaxations. The occurrence of dramatic electro-squeezeout occurring under constant load due to the charge-induced boundary solidification of the IL and its sudden melting when the charge is removed, is quite likely a general characteristic that should be explored in real ILs.

**IX. VOLTAGE BEHAVIOR BETWEEN PLATES**

The plate-confined IL also has interesting capacitive properties. The antisymmetric plate charging described in Sec. VIII corresponds to the application of a voltage between the plates. Assuming a uniform charge density in the $(x,y)$ plane and integrating Poisson’s equation along the vertical axis, we obtain an analytical expression for the voltage as a function of the coordinate $z$ between the plates

$$U(z) = -\int_0^z \frac{\rho(z')}{\epsilon_0 \epsilon_r} (z-z') dz' + \frac{q}{\epsilon_0 \epsilon_r} z,$$

where $\epsilon_0$ is the dielectric permeability of vacuum, $\rho(z)$ is the density along vertical axis $z$, and the second term is the contribution from surface charge density, $\pm q$, on the plates. Numerical values of $\rho(z)$ calculated from the simulation are integrated to obtain the voltage profiles, $U(z)$. Fig. 14(a) shows the variation of voltage for a charge density $q = \pm 10 \mu C/cm^2$ and six IL layers confined, in full thermodynamic equilibrium, between the plates. We find that while the potential drop in vacuum would be $\Delta U_V = 34$ V, in the presence of TM liquid $\Delta U_{TM} \approx 3.5$ V. Moreover, $\Delta U_{TM}$ remains nearly constant near the center of the film, showing that the TM liquid effectively screens the plate charge at short distances, as expected from the short Debye screening length of the bulk TM. Fig. 14(b) is the density profile along vertical axis, while panel Fig. 14(c) displays the average total charge $n_q$ per layer. The color of circles

![FIG. 13.](image1)

![FIG. 14.](image2)
refers to cations (red) and anions (blue), while the full circles indicate the plate charge in terms of electron charge \( q_A/e \), with \( A \) a area of plate. The total amount of charge, including the plates and confined liquid, is zero, preserving neutrality. The behavior \( n_q vs z \) shows a clear overscreening, a phenomenon well known in electrowetting\(^{15} \) with the charge of layers in contact with plates about twice larger than that on the plates.

X. DISCUSSION AND CONCLUSIONS

We have undertaken a detailed simulation study of the behavior of simple model ionic liquids confined between plates that are being pressed together producing squeezeout. We studied that without and with electrical charging of the plates, showing that charging influences both the structure of the liquid and its squeezeout behavior. Odd-number layering and near solidification is found between equally charged plates, from where squeezeout occurs by successive expulsion of neutral bilayers, as is commonly observed in experiments. The intimate structure of the confined film is analysed, showing that charging influences both the structure of the liquid and its squeezeout behavior. Odd-number layering and near solidification is found between equally charged plates, from where squeezeout occurs by successive expulsion of neutral bilayers, as is commonly observed in experiments.

Although the simplicity of the models permits only a limited connection with existing data, the present study demonstrates a variety of phenomena that can and will take place when real ionic liquids are confined under charged plates. The striking effectiveness of the charging induced solidification-melting process with consequent “electrosqueezing” discovered in simulation might have practical applications.

Finally, the strong control exerted on the IL nanostructure by the charging of plates suggests that the lubrication of plate sliding should be equally affected. That will be the subject of our forthcoming study.

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