Electrospray molecular dynamics simulations using an octree-based Coulomb interaction method

Neil A. Mehta and Deborah A. Levin
Phys. Rev. E 99, 033302 — Published 11 March 2019
DOI: 10.1103/PhysRevE.99.033302
A new octree-based Coulomb interaction model is developed to model the electrospray of ionic liquids (ILs) in molecular dynamics (MD). Using an octree-based method, Coulomb interactions are categorized as intra- and inter-leaf Coulomb interactions based on a criterion related to the Bjerrum length of the IL. The octree-based method is found capable of reproducing Coulomb energy in agreement with established and computationally more expensive models, such as the direct Coulomb and the Damped Shifted Force (DSF) method in the absence of an external electric field. In the presence of an external electric field, the octree-based method produces distinctly different results compared to that obtained by the direct Coulomb method. The time required to form Taylor's cone was shorter for the octree method compared to the direct Coulomb approach. While no emission larger than monomers was observed from the direct Coulomb simulation, emission of larger species such as dimers and trimers was observed when the octree-based Coulomb interaction model was used. Furthermore, the octree-based model forms a smaller ion emission cone compared to that from the direct Coulomb method.

I. INTRODUCTION

Electrosprays have been used extensively in the field of micro-fabrication for controlled deposition [1], in particular, microfilm deposition [2], microcircuit manufacturing [3], and ion beam lithography [4]. More fundamentally, because of the ability to generate ions from macro-molecules, electrosprays have been used to analyze the properties of ILs using electrospray ionization mass spectroscopy (ESI-MS) [5]. Similarly, the ability to generate energetic ionic emission has made it possible to use electrosprays with ILs as propellants to serve as a propulsion device for nano-satellite station keeping and micro-thrust adjustments [6–9]. The electrospraying process starts with the formation of a Taylor’s cone [10], whereby a liquid surface deforms in the presence of high electric fields and ion clusters are then emitted from the tip of this cone towards the extractor ring. While traveling towards the extractor ring, the emitted ion clusters undergo further evolution due to the external electric fields. While electrosprays have been in use for more than 30 years, the underlying mechanism of cone formation and post-emission droplet evolution at an atomistic level is only possible by performing MD simulations [11, 12].

From these simulations, it is understood that the process of electrospraying is primarily dominated by non-covalent forces and in particular, contributions from the Coulomb interactions are key for any accurate MD simulations of electrosprays at the atomistic level. However, unlike van der Waals forces, Coulomb interactions decay as the inverse of inter-atomic radius, making these forces important over long distances, and therefore, computationally the most expensive part for MD simulations of charged systems. Traditionally, electrostatic interactions are divided between short and long-ranges. Interactions are deemed short-range if the inter-atomic distance is less than the user-defined cut-off radius. The short-range interactions are calculated using Coulomb’s law while those over longer-ranges are computed using long-range Coulomb interaction methods, such as the Ewald summation [13]. In periodic charge dense systems, a cut-off radius of 10-15 Å is considered adequate to sub-divide interactions into short- and long-ranges. However, for simulation domains with nanometer length scales, such as those used to model electrosprays, the spatial non-homogeneity of ion-pair distributions lead to sparse charge regions which makes it necessary to use accurate long-range Coulomb models.

The choice of the long-range Coulomb model and the associated Coulomb cut-off radius significantly affects the outcome of MD electrospray simulations. From our previous electrospray MD simulations of EMIM-BF$_4$, it was found that the accuracy of conventional long-range Coulomb interaction models strongly depend on the Coulomb cut-off radius and the domain size [12]. In that work, we found that previous electrospray MD simulations of coarse-grained 1-ethyl-3-methylimidazolium tetrafluoroborate (EMIM-BF$_4$) [11] were performed using incorrect long-range Coulomb interaction parameters and the force shifted Coulomb sum (FSCS) method [14]. This, in turn, led to a spurious weak-
ening of Coulomb interactions and predicted abnormally high ion emissions.

Since MD simulations typically scale as \( O(N^2) \), where \( N \) is the number of atoms used in the simulation, it is not feasible to use an infinitely large cut-off radius, as the simulations would become prohibitively expensive. We face a two-fold problem of performing simulations with a cut-off radius small enough to keep the computation cost low for the short-range Coulomb interactions as well as having an accurate long-range model appropriate for the given MD simulations. For large simulations (> 10,000 atoms), the cut-off radius required to achieve convergence in the system’s Coulomb energy also increases. There are a number of methods available to calculate the long-range part of electrostatic interactions that vary in their complexity of implementation. However, many of these methods require the domain to be periodic in all three dimensions to work efficiently. While long-range methods, such as particle-particle particle-mesh (PPPM) \[15\] methods have been reliable in MD simulations of bulk ionic liquids with periodicity in at least two dimensions, they are found to be inaccurate for simulations with domains that have sparse density distributions, non-periodicity, and charge non-homogeneity in domains with high aspect ratios \[12\]. For these reasons, we propose a novel Coulomb interaction model based on the Barnes-Hut algorithm \[16\] to simulate such systems.

Hierarchical algorithms, such as the Barnes-Hut algorithm \[17\] and the fast multipole method (FMM) \[18\] have been used to compute Coulomb forces in MD simulations \[19, 20\]. The standard steps involved in these algorithms are the construction of an octree structure, traversing the octree from the leaves to the root to calculate the Coulomb energies of atoms in each leaf due to atoms belonging to other leaves (outer expansion), traversing the octree from the root to leaves to calculate Coulomb energies inside the leaf due to atoms outside the leaf (inner expansion), which is then followed by the accumulation of forces on each particle. To access the parent-child information between leaves at the multiple levels of the octree, point-based octree approaches have been implemented. The need to have a pointer-based approach, however, makes FMM challenging to implement and its efficiency depends on how the particles are distributed in the domain \[21\]. Furthermore, the accuracy of the FMM method depends on the order of expansion. For astrophysical applications, which is one of the main areas of use, three expansion coefficients are considered adequate since they provide approximately three significant digits of accuracy \[22\]. For MD simulations, typically, five digits of accuracy are needed, which is possible only when seven expansion coefficients are used \[23\]. Although considerable work has been done to parallelize the FMM method on computers with distributed memory with good efficiency \[24–27\], it generally scales poorly with increasing number of processors for expansion coefficients greater than three. To circumvent problems associated with pointer-based octree methods, we present a Barnes-Hut based algorithm, which relies on a linearized octree structure \[28\], similar to the approach used by Jambunathan and Levin \[29\] for their work on particle-in-cell methods. The linearization of the octree makes it possible to implement this method with the pre-existing LAMMPS framework \[30\].

We present here a Barnes-Hut based Coulomb interaction model with the specific intent of studying the evolution of ionic liquid systems with a non-homogeneous charge distribution in the domain. In this work, we perform all-atom MD simulations of ethylammonium nitrate (EAN) using the new Coulomb interaction method. EAN is categorized as a room temperature ionic liquid and is one of the oldest known ILs \[32\] with the chemical structure and atomic partial charge shown in Fig. 1. The electrochemical properties of EAN have been analyzed in detail using both, experiments \[33\] and MD simulations \[31, 34, 35\]. Lenguito, et al. \[36\] and Alonso-Matilla, et al. \[37\] have experimentally shown that
EAN can be successfully electrosprayed.

To demonstrate the capability of the octree-based Coulomb interaction model, two categories of simulations have been performed. The first set of simulations are used to study the fluctuations and energy stability of the octree-based method during the evolution of an isolated droplet of EAN. In the second set of simulations, the extrusion of EAN is simulated using a larger domain to model the entire electrospray process, including the initial time-steps when the Taylor’s cone is formed. To validate the Coulomb energies and total energy predicted from the octree-based Coulomb interaction model, the same set of simulations have also been performed using Coulomb’s law directly, but with a large cut-off radius of 40 Å. We discuss the improvement in accuracy when the octree-based method is used and look at the differences in the Coulomb energy and emission species and currents that result from these two methods.

II. NUMERICAL FRAMEWORK FOR THE OCTREE-BASED COULOMB MODEL

To improve computational efficiency without compromising the accuracy of the Coulomb energy, an octree-based Coulomb interaction model is proposed to calculate the short- and long-range Coulomb energies. An octree is a hierarchical structure which encompasses a domain that will undergo recursive subdivision into eight equal parts until a user-defined criterion is satisfied. In this structure, the three-dimensional simulation domain encompassing all the atoms in the system forms the first internal node, also known as the root or the queen. The root is divided into octants or children, which are the first generation of nodes. Based on the two limiting criteria, namely, (1) the number of particles present in the children and (2) its dimension, the children nodes will undergo further recursive subdivision, until one of the two conditions are met. The final nodes that do not undergo sub-division are known as the leaf nodes. Note that a leaf is analogous to a cell in grid based methods. A schematic of a two-dimensional octree is shown using arbitrary notional particles, representing the partial charges, instead of actual ion-pairs for simplicity, in Fig. 2(a). The red outermost frame, which encompasses the domain is known as the root of this octree. During the MD simulation, the root then undergoes recursive subdivision, every time step, to form leaves, which are analogous to cells of a uniform grid. In this example, the octree root has 10 leaves.

The electrostatic interaction energy between the partial charges of atoms $i$ and $j$ is given by Coulomb’s law as,

$$A_{ij} = C \frac{q_i q_j}{r_{ij}}$$  \hspace{1cm} (1)

where, $r_{ij}$ is the distance between atoms $i$ and $j$, $C$ is Coulomb’s constant, and $q_i, q_j$ are the user defined partial charges on atoms $i$ and $j$, respectively, as shown in Fig. 1. For a group of $n$ atoms, the
Coulomb energy is given as,

$$E_{\text{Coul}} = \sum_{i=1}^{n-1} \sum_{j=i+1}^{n} A_{ij}$$  \hspace{1cm} (2)

These electrostatic interactions scale as $1/r$, which leads to large fluctuations in energy for small changes in $r_{ij}$ at distance less than 15 Å but the fluctuations decay rapidly over larger distances ($> 60$ Å). When charge interactions are calculated by summing up the atomic partial charges over large inter-atomic distances, using Eq. 2 becomes computationally expensive. To overcome this limitation, the electrostatic energy calculations are split into short- and long-range Coulomb interactions, based on a user specified cut-off distance, $R_c$, and the electrostatic energy is now calculated as,

$$E_{\text{Coul}} = E_{\text{short-range}} + E_{\text{long-range}}$$  \hspace{1cm} (3)

where, the first term, $E_{\text{short-range}}$, takes into account all interactions with radius less than $R_c$ and is commonly known as $E_{\text{direct}}$ [38],

$$E_{\text{short-range}} = E_{\text{direct}} = \sum_{i=1}^{n-1} \sum_{j=i+1}^{n} A_{ij} \text{ for } r_{ij} < R_c.\hspace{1cm} (4)$$

$$B_{ij} = C \kappa \left[ \frac{\text{erfc}(\alpha_d \chi)}{\chi} - \frac{\text{erfc}(\alpha_d R_S)}{R_S} + \left( \frac{\text{erfc}(\alpha_d R_S)}{R_S^2} + \frac{2\alpha_d}{\sqrt{\pi}} \frac{\exp(-\alpha_d^2 R_S^2)}{R_S^2} \right) \right] \left( \chi - R_S \right) \text{ for } \chi \leq R_S \hspace{1cm} (5)$$

Using Eq. 2 with a cut-off radius leads to a major drawback, because the charges contained within the cut-off radius must be charge neutral. However, as the atomic partial charges move back and forth across the cut-off radius in subsequent time steps due to fluctuations in inter-atomic forces, the region inside the cut-off radius is no longer charge neutral and can cause a kinetic energy drift or a spurious energy gain within the system. Also the Coulomb energy calculations are no longer be considered “stable” or energy conserved [39]. These fluctuations in the inter-atomic forces are inherent to all MD simulations and one way of addressing them is by shifting and dampening the Coulomb interactions, such that they are close to zero at the cut-off radius. Using the DSF method of Fennell and Gezelter [14], the $A_{ij}$ term in Eq. 2 can be modified by introducing the shift radius, $R_S$, and dampening parameter, $\alpha_d$, as shown in Eq. 5, where, $\kappa = q_i q_j$ and $\chi = r_{ij}$. The complementary error function, erfc, is introduced to reproduce the effective screening used by the Ewald summation method and the dampening parameter accelerates the convergence of the Coulomb energies by dampening the Coulomb energy fluctuations around the shift radius [40, 41]. It can be observed that Eq. 5 will simplify to Eq. 2 if the dampening parameter, $\alpha_d = 0$ and shift radius, $R_S$ tends to infinity. The $R_S$ and $\alpha_d$ parameters are inter-dependent and the choice of $R_S$ used for the simulations of this work will be discussed in Sec. III. Equation 5 has been specifically used to obtain Coulomb energies for charged systems in a continuous bulk media. Typically, for simulations of bulk media, a smaller shift radius and consequently, a larger dampening parameter, is adequate as there are sufficient charges that contribute to the Coulomb energy within the spherical volume of radius equal to the shift radius. However, simulations of electrosprays represent a domain with non-homogeneous, sparsely distributed charges which will require very large ($\sim 100$ Å) shift radius and a small dampening parameter to accurately account for charges distributed over longer ranges. Since a larger shift radius could negate the benefits obtained by using Eq. 5, some modification of the DSF method will be required for the octree-based method.

For the octree based Coulomb interaction method, the Coulomb interactions are divided into the intra- and inter-leaf Coulomb energies,

$$E_{\text{Coul}} = E_{\text{intra-leaf}} + E_{\text{inter-leaf}}$$  \hspace{1cm} (6)

where, $E_{\text{intra-leaf}}$ is calculated as the sum of Coulomb interactions between all particles belonging to that octree leaf using,

$$E_{\text{intra-leaf}} = \sum_{m=1}^{l} \sum_{i=1}^{n_m-1} \sum_{j=i+1}^{n_m} B_{i,m;j,m} \hspace{1cm} (7)$$

where, $l$ is the total number of leaves in the octree, $n_m$ is the number of atomic partial charges in octree leaf $m$. For the schematic of an octree shown in Fig. 2(a), the intra-leaf Coulomb interactions are evaluated using Eq. 7 over all ten leaves. Figure 2(b) shows that the $E_{\text{intra-leaf}}$ contribution from leaf $m = 10$ will have Coulomb interactions from only the four notional particles belonging to that leaf.

To compute the $E_{\text{inter-leaf}}$ term, which includes
the Coulomb interactions between atomic partial charges belonging to different leaves, the center-of-mass of each leaf, $\vec{R}_p$, is first calculated and assigned a charge equivalent to the sum of charges of all particles belonging to that leaf, i.e.,

$$\vec{R}_p = \frac{\sum_{i=1}^{n_p} m_i \vec{r}_{i,p}}{\sum_{i=1}^{n_p} m_i}$$

where, $p$ is the leaf index, $n_p$ is the total number of atoms belonging to leaf $p$, and $\vec{r}_{i,p}$ and $m_i$ are the position vector and mass of the $i$th particle belonging to leaf $p$, respectively. The charge, $Q_p$, associated with the leaf $p$ is calculated as,

$$Q_p = \sum_{i=1}^{n_p} q_{i,p}.$$

The inter-leaf Coulomb energy is then calculated using,

$$E_{inter-leaf} = \sum_{m=1}^{l-1} \sum_{p=m+1}^{l} \left\{ \sum_{i=1}^{n_m} \sum_{j=1}^{n_p} B_{i,m;j,p} \text{ for } r_{i,p} \leq R_{inner} \cdot \theta \right\}$$

$$= \left\{ \sum_{i=1}^{n_m} \sum_{j=1}^{n_p} B_{i,m;j,p} \text{ for } r_{i,p} \leq R_{inner} \cdot \theta \right\} + \left\{ \sum_{i=1}^{n_m} \sum_{j=1}^{n_p} B_{i,m;j,p} \text{ for } r_{i,p} > R_{inner} \cdot \theta \right\}$$

where, for $B_{i,m;j,p}$, $\kappa$ is now calculated as the product of $q_{i,m}$ and $Q_p$, $\chi = |\vec{r}_{j} - \vec{R}_p|$ is the distance between the $i$th atomic partial charge belonging to leaf $m$ and the center-of-mass of leaf $p$. Note that the intra- and inter-leaf energy terms are analogous to the conventionally used terminologies of the short- and long-range interactions but with some important differences. In the octree-based Coulomb method, the short-range Coulomb interactions are now calculated using the intra-leaf term by Eq. 7 and the first part of the inter-leaf term by Eq. 10.

The demarcation between the short- and long-range Coulomb interactions, $R_{inner}$ and $\theta$, is not decided arbitrarily using a Coulomb cut-off radius. The inner radius, $R_{inner}$, corresponds to the dimensions of the smallest leaves in the octree and the accuracy parameter, $\theta$, determines the extent of the short-range Coulomb interaction. The product of the inner radius and the accuracy parameter becomes the effective cut-off radius over which Coulomb interactions are deemed short-range. For example, in the octree schematic constructed using the $R_{inner}$ criterion of 20 A, shown in Fig. 2(a), assuming that $\theta = 2$, the inter-octree Coulomb energy between notional particles A and B is computed using the first row of Eq. 10, since the distance between notional particle A and the center-of-mass of leaf $m = 2$, which contains notional particle B, is less than $R_{inner} \cdot \theta$. The distances between notional particle A and the center-of-mass of leaves $m = 7$, 8, and 10 are greater than $R_{inner} \cdot \theta$. Therefore, to calculate the Coulomb interaction between notional particles A and C, D, E, the second row of Eq. 10 is used.

The schematic shown in Fig. 2(a), also explains the origin of fluctuations that will also be observed in energy conservation for the octree-based Coulomb interaction method. If we consider the interaction between notional particles B and E, where, notional particle E lies at the boundary of leaves $m = 8$ and 5, it is clear that due to small fluctuations in its positions generated by interactions with other notional particles or kinetic energy, notional particle E may lie either in leaf $m = 7$ or 5 in subsequent time steps during the simulation. When notional particle E is present in leaf $m = 5$, its Coulomb interaction energy will be calculated using the first expression of Eq. 10 as the distance between the centers-of-mass of leaves $m = 2$ and 5 is less than $R_{inner} \cdot \theta$. However, when notional particle E jumps to leaf $m = 7$ in the next time step, the partial charge on notional particle E will now be approximated as a coarsened charge located at the center-of-mass of leaf $m = 7$ because the distance between the centers-of-mass of leaves $m = 2$ and 7 is greater than $R_{inner} \cdot \theta$. In the absence of dampening and force-shifting, this would lead to differences in the Coulomb energy calculated in subsequent time steps that would cause a spurious gain in kinetic energy of the system.

The octree algorithm proposed by Barnes-Hut [16] does not specify a lower limit on the minimum dimensions of the leaf, $R_{inner}$, but instead requires that each octree leaf should not contain more than one particle. However, this approach would not be computationally practical in regions with high number densities of atoms or for accurately maintaining the long-range order of the liquid during the simulation. Therefore, the accuracy parameter, $\theta$, and inner radius, $R_{inner}$, are selected based on physical considerations. For the ionic liquid simulations, we propose that $R_{inner}$ must be adequately large to capture all the co-ordination shell radii or peaks of the radial density function (RDF) of the bulk IL system under consideration. The average location of individual ions in a bulk solution is expressed in terms of a radial density function, $g(r)$. This function represents the probability of finding another ion at a par-
particular distance, \( r \), from an individual ion. Similar to the crystal structure of solids, there are distances at which ions can be found with a higher probability relative to other ions in ionic liquids. The distances at which higher probability of ions is observed is unique to each ionic liquid and is referred to as the long-range order, which can be verified using x-ray diffraction studies and MD simulations. For the nitrogen atoms of the ammonia and nitrate groups, representing the cation and anion parts of EAN, it is observed that the long-range order or atom-atom correlation persists to approximately 12 Å, as shown in Fig. 3. Therefore, a value of \( R_{\text{inner}} = 10 \) Å is implemented as the smallest leaf size for the octree-based Coulomb interaction model to perform MD simulations of EAN.

The Debye-Hückel-Bjerrum theory is used to obtain the accuracy parameter, \( \theta \), from the Bjerrum length, \( \lambda_B \). The Bjerrum length is defined as the distance at which the electrostatic interaction between two ions of opposite charges is comparable in magnitude to thermal energy, \( k_B T \), of the medium, which in our study is the energy equivalent to the temperature of the ionic liquid [42, 43]. To obtain the Bjerrum length, we use the Poisson-Boltzmann equation and the number density of the medium, \( \hat{n} \), to derive the inverse Debye screening length,

\[
\kappa_0 = \sqrt{4\pi \lambda_B \hat{n}(r)} = \sqrt{\frac{\hat{n}(r)e^2}{\epsilon_0 \epsilon_n k_B T}}, \tag{11}
\]

such that, the Bjerrum length, \( \lambda_B \) is,

\[
\lambda_B = \frac{e^2}{4\pi \epsilon_0 \epsilon_n k_B T}. \tag{12}
\]

Using \( \epsilon_n = 25.1 \) as the dielectric constant [33], in Eq. 12, the Bjerrum length of EAN is calculated to be, \( \lambda_B = 22.19 \) Å. The Bjerrum length represents the distance over which the Coulomb interaction is significantly more important for a given charge dense system compared to contributions from covalent bonds, angles, dihedrals, van der Waals potential, and kinetic energy. Thus, a larger value of \( R_{\text{inner}}, \theta = 30 \) Å is a reasonable criterion to demarcate the boundary between short- or long-range interactions for EAN simulations. Since the value of \( R_{\text{inner}} \) for EAN is 10 Å obtained from the radial density function, an accuracy parameter \( \theta = 3 \) is obtained. Using this hypothesis, we expect to obtain converged Coulomb energy values for the EAN simulations when using \( R_{\text{inner}} = 10 \) Å and \( \theta = 3 \), as will be demonstrated in Sec. III.

Finally, because the octree is an adaptive grid, the grid generation process is dynamic. During the MD simulations, the octree structure from the previous time step is destroyed and recreated every time step to ensure that the number of atomic partial charges in each leaf satisfy the recursive sub-division criterion at any given time. To construct the octree grid, the root is recursively subdivided using the \( R_{\text{inner}} \) and a minimum number of particles criteria. First, the number of ion-pairs (or partial atomic charges) in each leaf is counted. If there are more than two ion-pairs (\( \sim 40 \) atoms) inside a leaf, that leaf is subdivided further. This process is performed until either the number of ion-pairs in each leaf is a maximum of two, or the leaf size is equal to \( R_{\text{inner}} = 10 \) Å. During the simulation, atomic partial charges move from one leaf to another, which may cause the number of atomic partial charges in a leaf to be higher than the criteria used for the recursive subdivision. This is not problematic when calculating intra-leaf Coulomb interactions using Eq. 7 because all atomic partial charges mapped to each leaf are considered. However, when additional atomic partial charges enter into a previously formed octree leaf, the center-of-mass, \( \vec{R}_P \), calculated using Eq. 8, as well as the charge associated with that leaf, \( Q_p \), are affected. During the inter-leaf Coulomb evaluations, changes in center-of-mass and charges associated to the leaves when the octree structure itself has not changed would result in Coulomb interactions being calculated using the second expression of Eq. 10 instead of the first, which may lead to a spurious gain.
in kinetic energy.

III. MD SIMULATION SET-UP AND VALIDATION OF OCTREE-BASED COULOMB INTERACTION MODEL

To validate and assess the accuracy and efficiency of the octree-based Coulomb interaction method, two sets of MD simulations of EAN were performed using both, direct Coulomb as well as the octree-based Coulomb interaction models. Since, the EAN molecule consists of a relatively small cation and anion species, compared to more complex aprotic and other protic ILs, it is a good candidate to study the effects of Coulomb interaction models in the presence of an external electric field. For the simulations discussed in this section, 2,000 ion-pairs of EAN were arranged in a spherical geometry, with an approximate radius of 40.81 Å. The primary droplet, for this case, represents a large droplet that could be emitted in an electrospray and is large enough to analyze the effects of the Coulomb model but is also computationally tractable. The spherical arrangement of EAN ion-pairs is henceforth referred to as the primary droplet to distinguish it from the secondary emissions that occur when a droplet is placed in the presence of an external electric field. The primary droplet was placed in the center of the simulation domain with dimensions 160 x 160 x 1280 Å. However, since it is necessary to have a cubic domain shape to generate the octree root, the simulation domain was divided into eight equal roots, each of size 160 x 160 x 160 Å. Initially, the ion-pairs were arranged randomly within a sphere, which was followed by potential energy minimization step to eliminate un-realistic ion arrangements and atom/bond overlaps generated while arranging the ion-pairs. A canonical ensemble (NVT) simulation was performed for 250 ps using the Berendsen thermostat to lower and maintain the EAN primary droplet at 5.0 K.

All MD simulations in this work were performed using the LAMMPS toolkit [30]. The OPLS-AA inter-atomic potential used for the EAN simulations was obtained from the work of Jorgensen, et al. [44] and Umebayashi, et al [34]. A cut-off radius of 12.0 Å was used for the Lennard-Jones forces between the atoms. Each EAN ion-pair was modeled using nine covalent bonds and angles simulated using harmonic potentials. Each ion-pair was also modeled with four dihedral angles simulated using the OPLS potential. The covalent and non-covalent interaction parameters used in this work are provided as supplementary materials. A time step value of 0.5 fs was used for all the simulations. In addition to the octree-based Coulomb interaction method, all direct Coulomb and DSF simulations of the primary droplet were performed using a Coulomb cut-off radius of 40 Å. Note that a Coulomb cut-off radius of this size is not possible in the larger electrospray simulations performed in the presence of an external electric field that will be discussed in Sec. IV. Therefore for those larger simulations, the Coulomb cut-off radius will be reduced to 20 Å.

A. Selection of octree-based Coulomb interaction model parameters

There are three independent variables, namely, \( R_S \), \( R_{inner} \), and \( \alpha_d \) that are needed in the MD simulations with the octree-based Coulomb interaction model. To understand the sensitivity of the MD simulations to these parameters, the direct Coulomb method (Eq. 4), the DSF method (Eq. 5), and the octree-based Coulomb method (Eqs. 7 and 10) were used to calculate the Coulomb energy of a single snapshot composed of 2,000 ion-pair EAN droplets held at 5.0 K, as a function of the aforementioned parameters. Wolf, et al. [39, 45] and Demontis, et al. [46] recommend that the shift radius used in the MD simulations should be \( R_S \geq 5d_{ij} \), where, \( d_{ij} \) is the distance between the cation and anion in the first co-ordination shell. From the radial distribution function of the nitrogen atom of the ammonia and nitrate groups of EAN, shown in Fig. 3, the first co-ordinate shell radius is \( d_{ij} = 4.0 \) Å, which leads to an \( R_S = 20 \) Å. Demontis, et al. [46] also suggest that the optimal dampening parameter should be \( \alpha_d = 2/R_S \). These recommendations for \( R_S \) and \( \alpha_d \) were made for MD simulations of systems of 432 MgO molecules and aluminosilicate-water, respectively. These, however, are not adequate for MD simulations of ILs because the long-range order in liquids extends to longer distances compared to solids. Furthermore, as stated previously, when force shifting is used, all interactions beyond \( R_S \) are ignored. Therefore, for the octree-based method, a larger shift radius is used. The purpose of using the octree-based method is to account for Coulomb interactions over distances greater than 60 Å and a smaller shift radius would not allow us to account for these interactions. Since the EAN primary droplet has a diameter of 80 Å, a shift radii of 160 (two times the diameter) and 320 Å (four times the diameter) are used in the MD simulations. Using the expression given by Demontis, et al. [46], the associated \( \alpha_d \) values for \( R_S = 160 \) and 320 Å are 0.01 and 0.005,
FIG. 4: Comparison of Coulomb energy convergence for a single snapshot taken at a time of 35 ps of atom positions of 2,000 ion-pairs representing a EAN primary droplet at a temperature of 5.0 K. Direct Coulomb and DSF method ($R_S=320$ Å, $\alpha_d=0.005$) comparison shown in left and Octree-based Coulomb interaction method shown in right.

The comparison of the system Coulomb energy obtained from different Coulomb models is shown in Fig. 4. As shown in Fig. 4(a), using the dampening and force shifting for the DSF method with $R_S = 320$ Å and $\alpha_d = 0.005$ leads to a reduction (less negative) in the Coulomb energy of the system due to charge-screening and dampening of the Coulomb energies compared to that obtained from the DC method. When the DC or DSF method is used, it generates large fluctuations in Coulomb energies for small changes in the cut-off radius and provides converged results only beyond large cut-off distances greater than 40 Å. In comparison, as shown in Fig. 4(b), the octree-based Coulomb interaction model implemented using Eqs. 7 and 10 provides converged (non-fluctuating) Coulomb energy for an accuracy parameter as small as $\theta = 1.75$ ($R_{inner} \cdot \theta = 17.5$ Å) for all the selected $R_S$ and $\alpha_d$ values but due the dampening and shifting, the Coulomb energies are lower (less negative) than that from the direct method. The octree-based method allows one to account for electrostatic contributions over distances greater than traditional cut-off radii used for the DC method, which leads to better convergence when calculating the Coulomb energy. It is observed that decreasing the shift radius or increasing the dampening parameter leads to lower (less negative) Coulomb energy indicating a weakening of cohesion of the droplet structure. The Coulomb energy calculated by the DSF and the octree-based method with $R_S = 320$ Å and $\alpha_d = 0.005$ are in close agreement and therefore these $R_S$ and $\alpha_d$ values will be considered as the baseline values.

B. Validation of energy conservation using droplet simulations

For MD simulations of charge dense systems, it is vital to ensure that the selected Coulomb interaction method does not introduce large, spurious energy gains in the system due to numerical artifacts. One method of testing the presence of spurious energy gains is to perform microcanonical ensemble (NVE) MD simulations of charged systems using different Coulomb interaction methods, during which, the potential and kinetic energy of the atoms are monitored. As mentioned previously, while, the use of any limiting cut-off condition in treating Coulomb interactions precludes strict energy conservation, the gain or loss of energy should be small ($< 1\%$) in comparison to the sum of the kinetic and potential energies. To test the energy conservation of the
FIG. 5: Comparison of Coulomb energy and total (potential + kinetic) energy of an isolated primary droplet. Coulomb energy calculated using direct (Eq. 4), DSF (Eq. 5), and octree-based method (Eqs. 7 and 10 with \( R_S = \infty \) and \( \alpha_d = 0.0 \)) shown in (a). Coulomb energy calculated using DSF and octree-based methods (Eqs. 7 and 10) shown in (b). Total energy of simulations using direct, DSF, and octree-based method \( (R_S = \infty \) and \( \alpha_d = 0.0 \)) shown in (c). Total energy of simulations using octree-based methods shown in (d).

octree-based Coulomb interaction model, the primary droplet maintained at 5.0 K is simulated for an additional 25 ps using the Berendsen thermostat, following which, the thermostats are removed. The droplet is then simulated using a microcanonical ensemble (NVE), where any spurious changes of kinetic or potential energy are monitored. The primary droplet at 5.0 K has intentionally very low kinetic energy so that even small gains in energy can be easily observed. At higher temperatures, the gain in the kinetic energy would be a very small fraction of the total energy and would only be detected af-
ter considerably longer simulation time compared to that required for a 5.0 K primary droplet simulation.

When the Coulomb energy from the non-damped and non-shifted octree-based Coulomb model \( (R_S = \infty, \alpha_d = 0.0) \) is compared with that obtained for the direct Coulomb method using \( R_S = 40 \, \text{Å} \) (Eq. 4), large differences are observed in the potential energy, as shown in Fig. 5(a). In the presence of a thermostat, during the NVT phase of the simulation, the Coulomb energies from the non-damped octree-based and direct method agree within 1.0% (-10,400 eV versus -10,300 eV, respectively). However, in the absence of a thermostat for the non-damped octree-based simulation, the thermal motion of the atoms, back and forth across the leaf boundaries at distances equal to the effective cut-off radius \( (R_{inner} \cdot \theta) \) leads to large gain in kinetic energy of the atoms, which is then reflected by an increase in the Coulomb energies in the subsequent time steps, as shown in Fig. 5(a). A similar analysis of the isolated primary droplet is also performed using the DSF method (Eq. 5) with a shift radius of \( R_S = 320 \, \text{Å} \), and the dampening parameter, \( \alpha_d = 0.005 \). It is observed from Fig. 5(a) that the Coulomb energy obtained from the DSF method is 3.0 % lower than that obtained from the direct method at all simulation times (-10,050 eV versus -10,300 eV, respectively). Another noticeable difference between the two methods is reduced fluctuations in the calculated Coulomb energy of the system when the DSF method is used compared to the direct method. Shifting the potential truncates the electrostatic interactions and neglects all interactions beyond the shift radius. However, because the shift radius of 320 Å is sufficiently large to encompass the entire droplet geometry, the effects of this shift are minimal. In parallel, as discussed previously, the use of dampening parameter allows charge screening at the shift radius, reducing the influence of atoms moving back and forth at this radius. This leads to lower Coulomb energy fluctuations in the DSF Coulomb energies shown in Fig. 5(a).

Because shifting and dampening are also used in the octree-based method, the results from the DSF simulations are used as the baseline for comparison. The octree-based Coulomb model (Eqs. 7 and 10) are used to perform simulations for different combinations of \( R_S \) and \( \alpha_d \). From Fig. 5(b), it can be observed that using the octree-based method with shift radius and dampening parameter eliminates the spurious gain previously observed for the non-damped octree-based case shown in Fig. 5(a). Furthermore, compared to the Coulomb energies obtained from DSF method, the octree-based method results are free of fluctuations. For the DSF and octree-based simulations performed using the same \( R_S = 320 \, \text{Å} \) and \( \alpha_d = 0.005 \), the Coulomb energies from the octree method is only 1.0 % lower than those obtained from the DSF case (-9,950 eV versus -10,050 eV). When the DSF method is used, all Coulomb interactions within the shift radius are treated using Eq. 5. But when the octree-based Coulomb interaction method is used, interactions between atomic partial charges for distances greater than the \( R_{inner} \cdot \theta \) criterion are computed using the center-of-mass of the leaf and its associated charge (second part of Eq. 10). This coarsening of charges allows computational saving albeit with a small loss of accuracy, as indicated by a relative difference of only 1.0 % in the computed Coulomb energies. As previously discussed in Sec: III, the recommended dampening parameters for shift radii of 160 and 320 Å are 0.01 and 0.005, respectively. However, to analyze the effect of varying the dampening parameter, simulations are also performed using \( R_S = 160 \, \text{Å} \) with \( \alpha_d = 0.005 \) as well as using \( R_S = 320 \, \text{Å} \) and a \( \alpha_d = 0.01 \). It was found that using a larger dampening parameter or decreasing the shift radius adversely affects the calculation of Coulomb interactions by weakening the Coulomb energies of the system, as shown in Fig. 5(b).

A comparison of the total (sum of potential and kinetic) energy of the system for these simulations are shown in Figs. 5(c) and 5(d). As shown in Fig. 5(c), while the total energy from the direct and DSF simulations remain steady, the non-damped octree-based simulation shows a significant increase in the total energy. Since the Coulomb energy is the dominant contributor to the potential energy, the rise of the total energy for the non-damped octree-based method shown in Fig. 5(c) compared with that from the Coulomb energy in Fig. 5(a) 25 ps onwards shows that the total energy increases more rapidly than the Coulomb energy. This indicates that the kinetic energy of the system increases more rapidly than the Coulomb energy. This rapid increase of kinetic energy suggests that even small changes in Coulomb energy will lead to a large increase in the kinetic energy of the system. Although the total energy obtained using the DSF and octree method with \( R_S = 320 \, \text{Å} \) and \( \alpha_d = 0.005 \) can be seen in Fig. 5(b) to be in agreement, the octree-based method shows comparatively higher fluctuations in total energy. While the Coulomb energy obtained using the octree-based method, as shown in Fig. 5(b), is comparatively smoother compared to the DSF method because of the coarsening of charges beyond \( R_{inner} \cdot \theta \), coarsening also leads to loss of accuracy in calculating
FIG. 6: Snapshots of the simulation with constrained cation moving away from the primary droplet surface. Distance between cation and the droplet surface in (a), (b), (c), (d) is 0, 10, 30, and 60 Å respectively.

the electrostatic potential between any two particles at these distances. Considering the example octree shown in Fig. 2(a), there is a small difference in the electrostatic potential energy generated on particle D due to particle A, which is not equal to that generated on particle A by particle D due to the use of coarsening. However, it should be noted that over multiple time steps these small differences are canceled out resulting in a steady Coulomb energy calculations, devoid of spurious gains, as shown in Fig. 5(d). Consistent with the lower Coulomb energy (less negative) calculated using the octree-based method for $R_S = 160$ and $α_d = 0.01$, the total energy calculated for this simulation is also lower (less negative) compared to the DSF and the non-damped octree-based simulation.

Even though there are differences in the Coulomb energy of the system when either the Coulomb model or the model parameters are changed, the RDFs of the ammonia and nitrate nitrogen atoms for the DSF and the octree-based methods were found to coincide. The co-ordinate shell locations and their intensities obtained from all Coulomb models were the same, which suggests that the RDF and as a consequence, the EAN mass density, is not affected by the shift radii and dampening parameters used in this work.

C. Energy fluctuations due to leaf jumping

The octree-based Coulomb interaction method must ensure the continuity of the Coulomb forces across the interface between the intra- and inter-leaf Coulomb interaction limits. This is tested by artificially moving a single cation away from the primary droplet-surface. The cation is moved 0.1 Å every time step and the simulation is stopped when the cation moves 100 Å from the primary droplet surface. Snapshots from this MD simulation are shown in Fig. 6. When the cation is at the primary droplet surface or 10 Å above it, as shown in Figs. 6(a) and 6(b), respectively, the cation is acted upon by the Coulomb interaction from half the primary droplet using Eq. 7 because the effective cut-off is, $R_{inner} \cdot θ = 30$ Å. When it is moved farther than 30 Å away from the droplet surface, as shown in Figs. 6(c) and 6(d), the Coulomb interactions of the ion with the droplet are calculated entirely using Eq. 10.

The Coulomb energy of the cation during its motion away from the droplet should be devoid of spurious gains or losses within reasonable limits. As the cation moves away from the droplet, the Coulomb energies generated between the cation and the droplet by the direct and DSF methods are in reasonable agreement, as shown in Fig. 7(a). The non-homogeneous distribution of atomic partial charges inside the primary droplet as the cation moves away leads to fluctuations in the Coulomb energy. For the DSF method, the electrostatic in-
FIG. 7: Comparison of Coulomb energy of the constrained cation with potential energy contribution from other sources shown in (a). Comparison of Coulomb energy of the constrained cation from damped octree-based Coulomb interaction model shown in (b).

interactions between the atoms of the cation and the rest of the primary droplet are calculated on a per-atom basis, whereas, for the octree-based method, the interactions between the cation and the primary droplet ion-pairs are evaluated as a combination of intra-leaf (within the \( R_{\text{inner}} \cdot \theta \) limit) and by using the center-of-mass weighted average of charges (beyond \( R_{\text{inner}} \cdot \theta \) distance). This leads to lower (less-negative) Coulomb energies calculated by the octree-based method compared to the DSF method for distances between the cation and droplet surface of less than 15 Å, as shown in Fig. 7(b). The one-to-one electrostatic interactions between all atoms also leads to noisier Coulomb energies for the DSF method compared to the results from the octree-based method. When the cation-droplet surface distance is more than 15 Å, the Coulomb energies from the octree-based method and the DSF are in reasonable agreement and the result from the octree-based Coulomb interaction method is markedly less noisy. For the cation-droplet surface distance greater than 15 Å, the majority of the Coulomb interactions between the cation and primary droplet atomic partial charges in the octree-based Coulomb method is calculated using charge coarsening, which leads to a smoother Coulomb energy. The difference between the Coulomb energy from the octree-based and the DSF method at distances less than 15 Å does not cause significant differences in the total Coulomb energy of the system, as seen in the good agreement shown previously in Fig. 5(b). It is also worth noting that the intra-molecular potential contributions from other sources, such as the covalent bonds, angle, and dihedral terms, are more dominant than the Coulomb energy for a single cation beyond a 15 Å distance from the primary droplet, as shown in Fig. 7(a). However, for a system of many charged particles in the presence of external fields, as is the case for electrosprays, Coulomb interactions do exert considerable influence on the emission characteristics, as discussed next.

IV. MD SIMULATIONS WITH THE OCTREE-BASED COULOMB METHOD IN AN EXTERNAL ELECTRIC FIELD

Coulomb interactions are significant in dense charge systems especially in the presence of an external electric field which inputs energy into the system to overcome cohesive Coulomb interactions between ion-pairs. To study the ability of the Octree Coulomb method to model such systems two types of cases are considered in this section. First, the primary droplet geometry considered in the previous section is now simulated assuming external radial and normal electric field strengths of 0.05 and
FIG. 8: Snapshots of the primary droplet simulation in the presence of an external electric field. This case is performed with damped octree-based Coulomb interaction method with $R_S = 320$ Å and $\alpha_d = 0.005$. Droplet structure in (a), (b), and (c) is at a simulation time of 2.5, 5.0, and 7.5 ps, respectively.

In a second, larger simulation, 19,810 ion-pairs of EAN were placed inside a capillary formed by placing 17,115 platinum atoms as a hollow cylinder open on both ends, forming a radius of 59 Å and a height of 290 Å. The capillary was formed by placing 17,115 platinum atoms as a hollow cylinder, open on both ends. The EAN ion-pairs were randomly placed inside the volume of the cylinder formed by the platinum atoms. Similar to the primary droplet simulations, the capillary filled with the EAN ion pairs was also potential energy minimized to remove spurious atom/bond overlaps. This was followed by 250 ps of a canonical ensemble simulation to equilibrate the IL within the capillary at 295 K. The extrusion simulations were then performed by superimposing the external electric field calculated using Laplace’s equation for an extrusion potential of -60 V for the tip boundary condition [12]. The EAN ion-pairs were ejected from the capillary using a moving potential wall to generate a mass flow rate of $2.50 \times 10^{-12}$ kg/s. The domain size for the extrusion simulation was 320 x 320 x 1280 Å, such that the bottom of the capillary was placed normal to the $x-y$ plane at $z = 0$. To quantify ion emission from the extrusion simulations, emission currents are obtained by tallying the emitted ions at the collection zone, which is located 1000 Å away from the tip of the capillary. The collection zone starts starts in the $x-y$ plane at $z = 1270$ Å and continues up to $z = 1280$ Å. The lower plane of the collection zone is known as the extrusion plane. During the simulations, emitted ion-species are tallied in the collection zone to calculate the emission currents. The height of the collection zone is deliberately set at 10 Å, such that it is adequate to tally the emission but small enough to avoid double counting which may occur before the emitted ion-species are removed from the domain.

A. Droplet evolution in the presence of an external electric field

Snapshots from the droplet evolution simulation for the octree-based Coulomb interaction model with $R_S = 320$ Å and $\alpha_d = 0.005$ are shown in Fig. 8.
FIG. 9: Octree structure generated by the damped octree-based method for the droplet simulation in the presence of external electric field at 7.5 ps (see Fig. 8(c)). Top and bottom: isometric and front view. The smallest leaves are of size equal to $R_{\text{inner}} = 10$ Å.

FIG. 10: Comparison of number of secondary monomers emitted from the primary droplet in the presence of external electric field.

After 2.5 ps of simulation time, the primary droplet starts emitting secondary monomers (cations) towards the positive Z direction and anions towards the negative Z direction, as shown in Fig. 8(a). Unlike the spherical shape held by the droplet at 2.5 ps, significant deformation of the primary droplet can be observed after 5.0 ps, which is accompanied by a higher emission of secondary ions, as shown in Fig. 8(b). The primary droplet loses its shape and is on the verge of complete fragmentation after 7.5 ps in the presence of the external fields, as shown in Fig. 8(c). The particular arrangement of atoms shown in Fig. 8(c) is a simulation domain with a discrete distribution of charged particles, which is where the octree-based Coulomb method performs especially well. The octree generated for this simulation domain at 7.5 ps is shown in Fig. 9. It can be observed that the octree leaves (cells) are more refined near the center region with higher charge density and becomes coarser near the $Z_{\text{min}}$ and $Z_{\text{max}}$ boundaries.

A comparison of a linear curve-fit obtained for the number of monomers emitted during the simulation for the different Coulomb models is shown.
FIG. 11: Comparison of Coulomb energy and total energy per atom of the primary droplet in the presence of external electric field shown in (a) and (b), respectively.

in Fig. 10. For all Coulomb models, it was observed that only monomers were emitted during the simulation. It is evident that the number of monomers emitted by the direct Coulomb method is markedly higher compared to that from the DSF and octree-based simulations, even though, in the absence of any external fields, the Coulomb interactions calculated by all three methods are in relative agreement of about 4.0%, as discussed in Sec. III.B. When the first monomer is emitted, the limited Coulomb cut-off used by the direct Coulomb method restricts the Coulomb force that is exerted by the ion-pair on the droplet beyond the cut-off distance resulting in a higher emission rate predicted by the direct Coulomb method. In contrast for the DSF model, a much larger shift radius of $R_S = 320 \, \text{Å}$ and consequently, a much smaller dampening parameter of $\alpha_d = 0.005$, increases the number of Coulomb interactions between ion-pairs in the droplet as it fragments in the electric field compared to the direct method. This, in turn, lowers the DSF emission rate compared to that of the DC method. The strengths of the octree method become apparent during the dynamic evolution of the droplet in the presence of an external electric field. For the same $R_S$ and $\alpha_d$, the emission rate for the octree-based method is slightly higher compared to the DSF method because of its charge coarsening. However, the use of charge coarsening is justified by the fact that the octree-based simulation is computationally two times faster compared to the DSF method and provides considerably good agreement. Predictably, when the shift radius, $R_S$ is lowered to 160 Å, the octree-based emission rate is higher.

As shown previously in Fig. 5(a), the direct Coulomb method generates the most negative or strongest cohesion in the absence of an external electric field. However, when the electric field is applied, the Coulomb interactions calculated using direct Coulomb method rapidly start decreasing (less negative), faster than other methods, as shown in Fig. 11(a). These trends occur because a more stable Coulomb energies are calculated over longer distances with the DSF and the octree-based method that limit the rise (decrease) of Coulomb energy. This is also reflected in the gradual increase in the total energy (sum of potential and kinetic) of the system, shown in Fig. 11(b). The octree-based Coulomb model provides the ability to account for long-range Coulomb forces that counteract the acceleration of monomers extracted by the external electric field. This limits the kinetic energy gained by the secondary in-pair emissions, thus, slowing the increase in the kinetic energy of the system.
FIG. 12: Snapshot of atomic partial charge positions at 30 ps (a) for the capillary electrospray simulation performed with damped octree-based Coulomb interaction method, and front (b) and isometric (c) views of the generated corresponding octree structure. Radius of the capillary is 59 Å.

B. Capillary electrospray at $2.5 \times 10^{-12}$ kg/s

Having validated the performance of the octree-based Coulomb method for the droplet geometry in the presence of an electric field, simulations of the electrospray process are performed to study the formation of Taylor’s cone and the extrusion process. The capillary simulations contain a comparatively large number of atoms (314,265) and therefore, 512 processors were used to perform the cap-
illary electrospray simulations. Although the DSF and the octree-based methods are demonstratively superior in modeling emission from the IL droplet in the presence of an external electric field, the former cannot be used for the capillary geometry since the computational cost required for such a simulation would be prohibitively high for an $R_S = 320$ Å. Therefore, to compare the emission rate and currents obtained from the octree-based Coulomb model, simulations were also performed using the direct Coulomb method with a comparatively smaller cut-off radius of 20 Å, because even a 40 Å value would be untenable. When using the octree-based method, initial computations with a shift radius of $R_S = 320$ Å and $\alpha_d = 0.005$ indicated kinetic heating of the ionic liquid within the capillary due to thermal motion of the atoms as they move through a distance greater than the shift radius. While the shift radius of 320 Å and an associated dampening parameter of 0.005 provided converged Coulomb energies for the droplet geometry, the columnar arrangement of IL bulk in the capillary gives rise to highly non-symmetric interactions between ion-pairs within the shift radius leading to spurious kinetic energy gains for the smaller dampening parameter. Therefore, the dampening parameter was increased to $\alpha_d = 0.1$ only for the inter-leaf Coulomb interactions, but, $\alpha_d = 0.005$ was used for the intra-leaf Coulomb component. The shift radius was unchanged for both octree components.

The octree structure for a snapshot of atomic partial charge positions at 30 ps is shown in Fig. 12. At 30 ps, the Taylor’s cone structure is almost fully formed and, as shown in Fig. 12(a), ion-species emission can be seen occurring from the sides as well as the tip of the Taylor cone. Monomers A-C and dimer D can be seen to be moving towards the collection zone. As shown in Fig. 12(b), the octree grid is refined in the regions containing these ion-species. The three-dimensional isometric view of the octree structure generated for the atomic partial charge positions at 30 ps is shown in Fig. 12(c).

Snapshots of the Taylor’s cone formation from the MD simulation using the octree-based method are shown in Fig. 13. The snapshot in Fig. 13(a), obtained at 2.5 ps, shows ion-pairs creating the upper meniscus of the IL inside the capillary within its tip. At 25 ps, the external electric field affects the shape of the bulk structure of IL pushed out of the capillary, which initiates the formation of the Taylor’s cone, as shown in Fig. 13(b). The fully formed Taylor’s cone is seen in Fig. 13(c) at the tip of the capillary at 35 ps. The process of formation of the Taylor’s cone is similar for both, the direct and octree-based Coulomb method. However, the slightly weaker Coulomb energies generated due to
FIG. 14: Comparison of Coulomb (a), total energy per atom (b), emission currents (c), and number of atoms present in the simulation (d) during the capillary extrusion simulations. Figures 14(a), 14(b), and 14(d) use MD data collected over the entire domain and include both anions and cations.

dampening in the octree-based method causes the Taylor’s cone structure to form approximately 15 ps earlier compared to the direct Coulomb method.

Similar to the droplet evolution case, the Coulomb energy per atom of the system from the octree-based method is within 3% agreement of that obtained from the direct Coulomb method, as shown in Fig. 14(a). The charge coarsening coupled with heavier dampening for inter-leaf terms ($\alpha_d = 0.1$) generates a system with marginally weaker Coulomb interactions but this weakening is small and in fact, predicts somewhat slower ion emission and atom loss rate, as seen in Figs. 14(c) and 14(d), respectively. The total energy (sum of potential and kinetic energies) comparison, shown in Fig. 14(b), reveals significant differences between the octree-based and the direct Coulomb methods. During the initial stages of the Taylor’s cone formation, there is a steady growth
FIG. 15: Comparison of the cation monomer (a) and dimer (b) emission currents as a function of distance from the capillary tip, obtained from the octree-based and direct Coulomb methods for 2,000 frames after 20 ps simulation time.

FIG. 16: Comparison of the cation trimer emission currents as a function of distance from the capillary tip, obtained from the octree-based and direct Coulomb methods for 2,000 frames after 20 ps simulation time.

in the total energy for both the methods. Since the Coulomb energies of both the simulations are relatively constant, as shown in Fig. 14(a), the rise in the total energy can be attributed primarily to the kinetic energy gain. In the octree-based method, as the Taylor’s cone is being formed, more accurate representation of long-range Coulomb interactions predicts the emission of larger ion-species, such as dimers and trimers as the Taylor’s cone is forming, which leads to a rise in the total energy of the system beyond the first 15 ps.

Similar to the droplet emission case, when the ion-species are tallied at the collection zone, only monomer emission is observed from both the simulations. Since the ion emission is not continuous, emission currents are presented as cumulative moving averages for the two methods in Fig. 14(c). Consistent with the gain in kinetic energy of the system, monomer emission current from the octree-based method is higher compared to that from the direct method. However, the time of first-emission is delayed for the octree-based Coulomb interaction method. Even though the emission currents from the octree-based simulation is higher, the long-range Coulomb interactions provide stability to the Taylor’s cone structure, ensuring that the emission of ion-species occurs towards the extrusion zone and not towards the sides. This is also reflected in the rate of loss of atoms from the simulation domain during the Taylor’s cone formation. As shown in Fig. 14(d), there is a continuous loss of atoms,
FIG. 17: Comparison of the normalized $X$ (a) and $Y$ (b) velocity distribution functions of the atomic partial charges emitted at the tip of the capillary, obtained from the octree-based and direct Coulomb methods for cations after 20 ps for 2,000 time frames.

FIG. 18: Comparison of the normalized $Z$ (c) velocity distribution functions of the atomic partial charges emitted at the tip of the capillary, obtained from the octree-based and direct Coulomb methods for cations after 20 ps for 2,000 time frames.

There are quantifiable differences between the octree-based Coulomb and direct Coulomb simulations in terms of emission species and currents. Even though the Coulomb energy of the system predicted by the direct Coulomb method is more negative compared to the octree-based method, the limited region within which the Coulomb interactions are calculated for the direct Coulomb method leads to more emission of smaller ions rather than larger species. Also, only monomers were observed in the collection zone since any larger ion-species that were emitted from the Taylor’s cone underwent further fragmentation due to energy obtained from the external electric field while moving towards the collection zone. To better understand the nature of emissions from the Taylor cone, instead of only tallying ion-species at the fixed collection zone 1,000 Å away from the tip of the Taylor’s cone, the ion-species are tallied every 50 Å from the capillary tip. Comparisons of the monomer, dimer, and trimer emissions as a function of distance from the capillary tip, for the two simulations, are shown in Figs. 15(a), 15(b), and 16, respectively. While no species larger than monomers were observed from the direct Coulomb simulation at any
FIG. 19: Comparison of the normalized emission probability distribution functions of both cations and anions emitted between 20 and 45 ps, obtained from the octree-based and direct Coulomb methods. Capillary tip is located at \( z = 290 \text{ Å} \).

distance from the capillary tip, significant emission currents from the dimer and trimer were observed closer to the capillary tip from the octree-based simulation, as shown in Figs. 15(b) and 16, respectively. From our previous simulations [47], it was observed that larger ion-species are emitted only from a well-formed, coherent Taylor’s cone. Thus, it can be concluded that the octree-based Coulomb model promotes the formation of a more coherent Taylor cone structure compared to the direct Coulomb method and as a consequence, the latter method does not predict the emission of larger ion-species.

The lower emission current coupled with the higher loss of ion-pairs predicted by the direct Coulomb model (see Figs. 14(c) and 14(d), respectively) suggests that a majority of ion-pairs emitted from the capillary do not contribute to the emission currents tallied at the collection zone. These are lost or wasted when they leave from the lateral boundaries due to sideways or radial emission since they do not contribute to the emission currents after the Taylor cone is formed. The Taylor’s cone structure is also weaker and hence allows a higher number of ion-pairs to escape, leading to significant, albeit a wasteful loss of ion-pairs from the domain. This was measured by comparing the velocity distribution functions for all cations emitted from the capillary for both the simulations. A comparison of the \( X \) and \( Y \) lateral velocity components shows that the distribution is narrower for cations emitted from the octree-based simulation, as shown in Figs. 17(a) and 17(b). This indicates that when the direct Coulomb method is used, ion-pairs are emitted with higher lateral velocities compared to that observed when the octree-based method is used. Another significant difference between the two methods can be noted from the distribution of the axial (\( Z \) component of) velocity. As shown in Fig. 18, the distribution from the octree-based simulation is not only comparatively narrower but also the peak of the distribution is located at a lower axial velocity compared to that observed from the direct Coulomb method. This is again indicative of heavier cations such as dimers and trimers that are emitted when the octree-based Coulomb method is used.

While the Taylor’s cone was observed in both set of simulations, the stability provided to this structure due to long-range Coulomb interactions by the octree-based Coulomb method generates more focused, concentrated emission. This was analyzed by generating a probability distribution of distance from the capillary tip at which the ions leave the domain between 20 and 45 ps during the simulation, representing the time during which the Taylor’s cone is partially to fully formed. As shown in Fig. 19, the normalized probability distributions from the two methods are considerably different. The peak of both distributions lie within 100 Å from the capillary tip and is comprised of monomer anions that cannot travel towards the collection zone because the applied electric field is positive. That is, anions leave the domain from the lateral boundaries at distances less than 100 Å from the capillary tip. The secondary peak observed from the octree-based simulation represents the distance away from the capillary tip at which a large fraction of cations leave the domain. The wider velocity distribution from the direct Coulomb simulation indicates that the cone-emission is isotropic and the majority of the ions emitted from the capillary are lost laterally, soon after they are emitted.

**V. CONCLUSIONS**

In this work, we have used a modified Barnes-Hut algorithm to develop an octree-based Coulomb interaction model targeted towards simulations of ILs, in the presence of external electric fields and sparse, non-periodic domains. The Coulomb interactions are sub-divided into intra- and inter-leaf Coulomb interactions based on the Bjerrum length criterion,
thus associating the Coulomb cut-off criterion with a physical electro-chemical property of the simulated IL. The energy conservation validation of the octree-based method showed that attention must be paid to the selection of the shift radius and dampening parameters to avoid spurious gains in the kinetic energy of the system. To obtain agreement with the direct Coulomb and DSF method Coulomb energies, the octree method simulations were performed with $R_S = 320 \text{ Å}$ and $\alpha_d = 0.005$. Using a smaller shift radius and consequently, a larger dampening parameter would lead to weaker Coulomb energies of the system. It should be noted that the selection of $R_S$ and $\alpha_d$ was based on the characteristic dimensions of the simulation, such as, the radius of the primary droplet and the length of the capillary. It is advisable to use the largest characteristic length present in the simulation as a shift radius when employing the octree-based Coulomb interaction method to avoid the need to run simulations to estimate its value. The use of the octree-based Coulomb interaction model in the absence of an electric field does not change the physical properties of the simulated ILs, such as the radial density function.

While the Coulomb and total energy of the system obtained from the direct, DSF, and the octree-based methods are in reasonable agreement in the absence of external electric fields, the differences are significant between the direct and the octree-based method in the presence of an external electric field. For the droplet evolution test case, the ability to account for Coulomb interactions over longer distances limited the emission of ions when the octree-based Coulomb interaction model was used, even though, the Coulomb energies of the entire system are weaker (less-negative) compared to that predicted by the direct Coulomb method. For the same droplet geometry, the octree-based Coulomb method was found to be twice as fast compared to the DSF method, while providing a good agreement in Coulomb energies and ion emission rates.

For the capillary based simulations, the long-range Coulomb interactions captured by the interleaf Coulomb term in the octree-based method reduces the Taylor cone formation time compared to the direct Coulomb model. Furthermore, emission of larger species such as dimers and trimers was observed only from the simulation performed with the octree-based Coulomb interaction model. The lack of long-range interactions leads to larger tangential or cross-stream emission of ions and subsequent loss of atoms when the direct Coulomb method is used to model extrusion from the capillary geometry. This resulted in a significantly lower emission current compared to that obtained from the octree-based Coulomb interaction model. Finally, it was shown that the emission of ions from the Taylor’s cone was into a smaller angle when the octree-based method was used for the capillary extrusion case.

VI. ACKNOWLEDGMENT

Funding for this work was provided by the Air Force Office of Scientific Research (AFOSR) through Dr. Mittat Birkan under Grant No. AF FA9550-16-1-0193. We are thankful to XSEDE TACC for providing us with the vital computational resources on their STAMPEDE2 cluster.

[9] B. D. Prince, P. Tiruppathi, R. J. Bemish, Y.-H. Chiu, and E. J. Maginn, Molecular dynamics simulations of 1-ethyl-3-methylimidazolium...
A. Borner and D. A. Levin, Use of advanced particle methods in modeling space propulsion and its supersonic expansions (PHD Dissertations, PSU, 2014).


P. Walden, Ueber die molekulargrosse und elektrische leitfahigkeit einiger geschmolzenen salze, Bulletin de l’Academie Imperiale des Sciences de St.-Petersbourg 8, 405 (1914).


J. V. L. Beckers, C. P. Lowe, and S. W. De Leeuw, An iterative pppm method for simulating coulombic...


