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MetalWalls: Simulating electrochemical interfaces between polarizable electrolytes and metallic electrodes

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Electrochemistry is central to many applications, ranging from biology to energy science. Studies now involve a wide range of techniques, both experimental and theoretical. Modelling and simulations methods, such as density functional theory or molecular dynamics, provide key information on the structural and dynamic properties of the systems. Of particular importance are polarization effects the electrode/electrolyte interface, which are difficult to simulate accurately. Here we show how these electrostatic interactions are taken into account in the framework of the Ewald summation method. We discuss, in particular, the formal set up for calculations that enforce periodic boundary conditions in two directions, a geometry that more closely reflects the characteristics of typical electrolyte/electrode systems and presents some differences with respect to the more common case of periodic boundary conditions in three dimensions. These formal developments are implemented and tested in MetalWalls, a molecular dynamics software which captures the polarization of the electrolyte and allows the simulation of electrodes maintained at a constant potential. We also discuss the technical aspects involved in the calculation of two sets of coupled degrees of freedom, namely the induced dipoles and the electrode charges. We validate the implementation, first on simple systems, then on the well-known interface between graphite electrodes and a room-temperature ionic liquid. We finally illustrate the capabilities of MetalWalls by studying the adsorption of a complex functionalized electrolyte on a graphite electrode.

I. INTRODUCTION

The simulation of electrochemical systems has become an important topic in chemical physics. This is mainly driven by the need for understanding the molecular mechanisms at play in devices such as batteries,1 electrocatalysts,2–5 supercapacitors,6 etc. Such simulations concern bulk electrode materials7 and electrolytes,8 whose properties need to be well characterized and rationalized, but more and more efforts are devoted to the description of electrode/electrolyte interfaces.9 Until recently, very little was known of the latter at the molecular scale and the only available picture was provided by theories such as the Gouy-Chapman-Stern one.6 However, these theories are limited to simple systems,10 and the use of concentrated electrolytes (e.g. ionic liquids) and complex electrode materials in current electrochemical devices requires much more sophisticated techniques, among which molecular dynamics (MD) simulation is the most widespread. Depending on the target properties, the flavor of the MD can be adapted: Classical MD, in which the forces are derived from an analytical interaction potential, is preferred when the objective is to obtain the structural, thermodynamic and transport properties of chemically inert systems, while ab initio MD, based on electronic structure calculations, most often in the framework of Density Functional Theory, for the evaluation of the interactions, is more adapted for studying systems undergoing chemical reactions such as chemisorption.11

Among the various methods used to simulate electrochemical interfaces,9 one of the most popular was introduced by Siepmann and Sprik.12 It consists in representing the electrode atoms using Gaussian charges, whose magnitude fluctuates in order to satisfy a constant potential condition within a given electrode, while the charge distribution within the electrolyte is represented

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using point charges. A difficulty when studying interfaces in MD is the handling of periodic boundary conditions (PBCs), in particular for long-ranged interactions. Usual implementations in MD codes correspond to 3D PBCs, so that interfacial systems become an infinite collection of slabs. This effect can be canceled using corrections such as the one introduced by Yeh and Berkowitz, and introducing a wide layer of vacuum between periodic images. Another approach is to derive explicitly the equations for 2D-periodic systems. Reed and Madden have thus formulated Ewald-type expressions for electrochemical systems made of point charge electrolytes and Gaussian charge electrodes. Their equations were later corrected by Gingrich and Wilson. This constant potential method was first implemented with 3D PBCs in the popular MD code LAMMPS, and technical improvements were recently proposed to improve the efficiency of the simulations, such as the use of particle-particle-particle-mesh solver approach or the use of a doubled cell approach.

However, the inclusion of polarization effects often improves the accuracy of the description of liquid electrolytes. In particular the dynamic properties of concentrated electrolytes, such as the diffusion coefficients, are better reproduced when including polarization. The charge-dipole and dipole-dipole interactions are also long-ranged, so one needs to treat them appropriately. In the presence of electrodes with Gaussian charges, the only derivation available was proposed by Pounds in his PhD thesis. A second difficulty lies in the self-consistent calculation of the electrode charges and of the induced dipoles for a given configuration of the electrolyte. In general, the two problems are solved by using conjugate gradient minimization techniques (even though in the case of electrode atoms, their position is often fixed which allows the use of more efficient methods such as matrix inversion or mass-zero constrained dynamics, but now the two problems are coupled: the charges depend on the dipoles and vice versa. In a first attempt to include both effects in simulations, we have used a recursive approach in which the two minimizations are performed one after each other until the two problems are converged. However, this increases considerably the computational time, limiting the ability to simulate large systems so that only molten salts could be studied. Borodin and co-workers proposed to update the electrode charges much less frequently than the atomic positions (typically every 250 fs), but they did not discuss the use of appropriate multiple time step techniques to achieve this.

MetalWalls is a MD software dedicated to the simulation of electrochemical interfaces. Here we introduce the recent implementations allowing for the inclusion of the interactions between polarizable electrolytes and metallic electrodes that are represented with fluctuating Gaussian charges. In the first section, we provide the derivation of the energies associated with these interactions within a 2D-Ewald framework. The second section discusses the optimization problem, for which we show that Jacobi preconditioning allows an efficient use of the conjugate gradient method. The implementation is then validated by simulating simple systems for which we can obtain reference results. Finally, we provide some results for two electrochemical systems. Firstly, we simulate a typical ionic liquid (1-ethyl-3-methylimidazolium bis(trifluoromethylsulfonyl)imide, EMIM-TFSI) between graphite electrodes, a system for which accurate non-polarizable force fields are available for comparison. Secondly, we study the adsorption of redox-functionalized ionic species, again on graphite, using a polarizable interaction potential recently parameterized in our group.

II. INTERACTION MODEL

The model implemented in MetalWalls includes different electrostatic components. The electrolyte is modeled as a set of \( N_p \) point charges and \( N_d \) point dipoles at positions \( \{ r_i \}_{i=1}^{N_p} \) and \( \{ r_j \}_{j=1}^{N_d} \), respectively. Denoting by \( \delta^3(r-r_i) \) the three-dimensional Dirac's delta centered at the point \( r_i \), the charge distribution of a point charge is simply given by

\[
\rho_i^q(r) = q_i \delta^3(r-r_i) \tag{1}
\]

so that, given the definition of the electric potential

\[
V(r) = \int \frac{d^3r'}{|r-r'|} \rho(r') \tag{2}
\]

the set of point charges generates a potential in space given by

\[
V^q(r) = \sum_{i=1}^{N_p} \frac{q_i}{|r-r_i|} \tag{3}
\]

Dipoles can be modeled as two opposite point charges connected by a rigid rod of length \( \delta \), so that the charge distribution can be written as

\[
\rho_j^\mu(r) = -q_j \delta^3(r-r_j + \delta/2) + q_j \delta^3(r-r_j - \delta/2) \tag{4}
\]

Using again Eq. (2) and taking the limit of \( \delta \to 0 \) (point-dipole approximation), the potential generated by the set of point dipoles is given by

\[
V^\mu(r) = \sum_{J=1}^{N_d} \frac{\mu_J \cdot (r-r_J)}{|r-r_J|^3} \tag{5}
\]

where \( \mu_J \) is the point dipole associated to the particle \( J \).

Electrodes are modeled using the so-called fluctuating charge model, which considers the metallic electrodes composed by \( N_g \) Gaussian-distributed charges

\[
\rho_{\alpha}^G(r) = Q_\alpha \left( \frac{\eta_\alpha^2}{\pi} \right)^{3/2} \exp\left[-\eta_\alpha^2 (r-R_\alpha)^2\right] \tag{6}
\]
placed at fixed sites \( \{R_{\alpha}\}_{\alpha=1}^{N_{\eta}} \), with \( \eta_{\alpha} \) being a model parameter (which can be changed to tune the electrode metallicity\(^{30}\)). The potential generated by this set of charges is given by

\[
V^Q(r) = \sum_{\alpha=1}^{N_{\eta}} \frac{Q_{\alpha}}{|r - R_{\alpha}|} \text{erf}(\eta_{\alpha}|r - R_{\alpha}|) \tag{7}
\]

The interaction energy for a system of Gaussian-distributed charges, point charges and point dipoles is composed by the six contributions

\[
U_{\text{elec}} = U^\mu\mu + U^{QQ} + U^{Qq} + 2U^q\mu + 2U^{qq} + 2U^{q\mu} \tag{8}
\]

where the superscripts indicate the specific interactions, so, for example, \( U^{Qq} \) is the interaction between the electrolyte’s point charges and the point dipoles. These energies are derived from the charge distribution density and from the electrostatic potential of the particular electrostatic components involved through the relation

\[
U_{\text{elec}} = \frac{1}{2} \int_{\mathbb{R}^3} d^3r d^3r' \frac{\varrho(r)\varrho(r')}{|r - r'|} = \frac{1}{2} \int_{\mathbb{R}^3} d^3r \varrho(r) V(r) \tag{9}
\]

where in going from the second to the third equality, the definition of the electrostatic potential given in Eq. (2) has been recognized. In what follows we explicitly report the expressions for the 2D Ewald decomposition for all the energy terms appearing in Eq. (8). Some of these terms are already reported in the literature (see for example Ref. 15 for \( U^{qq} \), \( U^{QQ} \) and \( U^{Qq} \)) and others are only discussed in the case of 3D PBC (\( U^\mu\mu \) and \( U^{q\mu} \) in Ref. 37). To the authors knowledge, no derivation or Ewald decomposition has ever been reported in the literature for the term \( U^{Qq} \). We report a complete derivation for energies and forces for this term in the Supplementary Material.

### A. 2D-PBC Ewald decomposition for Energies

Here we summarize the Ewald decomposition in the case of 2D-PBC for all the energy terms involved in the model of a system of interacting point-charges, point-dipoles and Gaussian charges. This is relevant because, in electrochemical systems, PBC are usually enforced only in the directions parallel to the electrodes. Assuming the electrodes extending on the \( xy \)-plane, the presence of replicated simulation boxes means that, for every particle in the system, there are an infinite number of other replicas at relative positions \( m = (n_x L_x, n_y L_y, n_z L_z) \) where \( L = (L_x, L_y, L_z) \) are the dimensions of the primary simulation box and, in the chosen geometry, \( n_x, n_y \in \mathbb{Z} \) while \( n_z = 0 \). To compute the energy of the system, a sum over all the replicas (i.e., over all boxes) needs to be performed. This sum will be noted in what follows by \( \sum_n \) and it runs over all the integers from \(-\infty\) to \(+\infty\) in all the components involved in the PBC. As commonly done in the Ewald decomposition, the conditional convergence of the Coulomb interaction is solved for each term of Eq. (8) by dividing the infinite summation over periodic boxes in a short-range part in real space (denoted by \( U_w \)) and a long-range in reciprocal space (denoted by \( U_b \)). Eventual artifacts arising from overcounting are taken into account through a self-energy term (denoted by \( U_{\text{self}} \)). The reciprocal space is characterized by the wavevectors indexes \( k \in \mathbb{Z}^3 \). The size of the system defines the reciprocal space wavevectors through the relation \( h = (2\pi k_x/L_x, 2\pi k_y/L_y, 2\pi k_z/L_z) \), where, once again in the chosen geometry, \( k_x, k_y \in \mathbb{Z} \) and \( k_z = 0 \). Due to the 2D PBC, the long-range term is to be divided in turn in the contribution from \( k = 0 \) and \( k \neq 0 \), which are denoted by \( U_{l=0} \) and \( U_{l>0} \), respectively. In the following we assume the periodic part of the system to be embedded in a medium with infinite dielectric constant (the so-called “tinfoil boundary conditions”\(^{38}\) together with global electroneutrality of the system.
1. **Point charge-point charge interaction**: $U^{qq}$

The expression is written as a sum of four terms $U^{qq} = U^{qq}_{sr} + U^{qq}_{ir,0} + U^{qq}_{ir,*} + U^{qq}_{self}$,

$$U^{qq}_{sr} = \frac{1}{2} \sum_{i=1}^{N_x} \sum_{j=1}^{N_p} \sum_{n} q_i q_j \frac{\text{erfc}[\alpha|\mathbf{r}_{ij} + \mathbf{m}|]}{\sqrt{\pi}}$$  \hspace{1cm} (10a)

$$U^{qq}_{ir,0} = -\frac{\sqrt{\pi}}{L_x L_y} \sum_{i=1}^{N_x} \sum_{j=1}^{N_p} q_i q_j \left( \exp\left[ -\alpha^2 \frac{z_{ij}^2}{\alpha} \right] + \sqrt{\pi} z_{ij} \text{erf}[\alpha z_{ij}] \right)$$  \hspace{1cm} (10b)

$$U^{qq}_{ir,*} = \frac{1}{L_x L_y} \sum_{i=1}^{N_x} \sum_{j=1}^{N_p} \int_{-\infty}^{\infty} du \sum_{k \neq 0} q_i q_j \frac{\exp\left[ -\frac{h^2 + u^2}{4\alpha^2} \right]}{h^2 + u^2} \exp[i(h \cdot \mathbf{r}_{ij} + u z_{ij})]$$  \hspace{1cm} (10c)

$$U^{qq}_{self} = -\frac{\alpha}{\sqrt{\pi}} \sum_{i=1}^{N_p} q_i^2$$  \hspace{1cm} (10d)

In Eq. (10a) we have introduced the primed sum over the replicated boxes to indicate that for $n = (0, 0, 0)$ the case $i = j$ is excluded. With some differences in notation these terms have already been reported in Ref. 15.

2. **Point charge-Gaussian charge interaction**: $U^{qQ}$

No self interaction has to be considered in this case as particles are of different nature. This fact is also highlighted by the extended notation $\mathbf{r}_i - \mathbf{R}_\alpha$ in place of the corresponding $\mathbf{r}_{ij}$ of the previous set. We can then write $U^{qQ} = U^{qQ}_{sr} + U^{qQ}_{ir,0} + U^{qQ}_{ir,*}$, with the following expressions for the various terms:

$$U^{qQ}_{sr} = \frac{1}{2} \sum_{i=1}^{N_x} \sum_{\alpha=1}^{N_p} \sum_{n} q_i q_\alpha \frac{\text{erfc}[\alpha|\mathbf{r}_i - \mathbf{R}_\alpha + \mathbf{m}|]}{\sqrt{\pi}} - \text{erfc}[\eta_\alpha|\mathbf{r}_i - \mathbf{R}_\alpha + \mathbf{m}|]$$  \hspace{1cm} (11a)

$$U^{qQ}_{ir,0} = -\frac{\sqrt{\pi}}{L_x L_y} \sum_{i=1}^{N_x} \sum_{\alpha=1}^{N_p} q_i q_\alpha \left( \exp\left[ -\alpha^2 \frac{(z_{i\alpha} - Z_\alpha)^2}{\alpha} \right] + \sqrt{\pi} (z_{i\alpha} - Z_\alpha) \text{erf}[\alpha (z_{i\alpha} - Z_\alpha)] \right)$$  \hspace{1cm} (11b)

$$U^{qQ}_{ir,*} = \frac{1}{L_x L_y} \sum_{i=1}^{N_x} \sum_{\alpha=1}^{N_p} \sum_{k \neq 0} q_i q_\alpha \exp\left[ -\frac{h^2 + u^2}{4\alpha^2} \right] \exp[i(h \cdot (\mathbf{R}_\alpha - \mathbf{r}_i) + u(Z_\alpha - z_{i\alpha})]$$  \hspace{1cm} (11c)

Note also that long-range interactions have the same form as in the case of the point-charge to point-charge case, with the integrated Gaussian charge $Q_\alpha$ in place of the point charge $q_i$. Also these terms were defined in Ref. 15 with some differences in notation.

3. **Point charge-point dipole interaction**: $U^{q\mu}$

The interaction energy between point charges and point dipoles has been previously computed, for example, in Ref. 24 where 3D PBC are assumed. The extension to the 2D PBC case does not present particular difficulties and we report it in what follows. Note that, once again, particles involved in the sums belong to different set. Even though they both refer to electrolyte particles, we could imagine to have point-charge particles which are non-polarizable and also the opposite situation is possible. To emphasize this aspect we use the extended notation $\mathbf{r}_i - \mathbf{r}_j$ in place of $\mathbf{r}_{ij}$ also in this case. For the same reason, no self-interaction term is present and the interaction energy can be split as
\[ U^{Q\mu} = U_{st}^{Q\mu} + U_{lr,0}^{Q\mu} + U_{lr,*}^{Q\mu} \] The single terms are written as

\[ U_{st}^{Q\mu} = \frac{1}{2} \sum_{\alpha=1}^{N_{\alpha}} \sum_{\beta=1}^{N_{\beta}} \sum_{n} Q_{\alpha} Q_{\beta} \left( \text{erfc}[\alpha R_{\alpha} - r_{j} + m] + \frac{2\alpha}{\sqrt{\pi}} |R_{\alpha} - r_{j} + m| \exp[-\alpha^2 |R_{\alpha} - r_{j} + m|^2] \right) \]

(12a)

\[ U_{lr,0}^{Q\mu} = \frac{\pi}{L_{x} L_{y}} \sum_{\alpha=1}^{N_{\alpha}} \sum_{j=1}^{N_{j}} q_{\alpha} \mu_{j} \text{erf}[\alpha(z_{i} - z_{j})] \]

(12b)

\[ U_{lr,*}^{Q\mu} = -\frac{i}{L_{x} L_{y}} \sum_{\alpha=1}^{N_{\alpha}} \sum_{j=1}^{N_{j}} \int_{-\infty}^{\infty} \int_{-\infty}^{\infty} du \sum_{k \neq 0} q_{\alpha} (\mu_{j} \cdot h + \mu_{j} u) \exp\left[-\frac{h^2 + u^2}{4\alpha^2}\right] \exp\left[i(h \cdot (r_{i} - r_{j}) + u(z_{i} - z_{j}))\right] \]

(12c)

4. Gaussian charge-Gaussian charge interaction: \( U^{QQ} \)

The mutual interaction between Gaussian-distributed charges is also given in Ref. 15. Writing, as always, \( U^{QQ} = U_{st}^{QQ} + U_{lr,0}^{QQ} + U_{lr,*}^{QQ} + U_{self}^{QQ} \), we have

\[ U_{st}^{QQ} = \frac{1}{2} \sum_{\alpha=1}^{N_{\alpha}} \sum_{\beta=1}^{N_{\beta}} \sum_{n} Q_{\alpha} Q_{\beta} \left( \text{erfc}[\alpha(R_{\alpha} - r_{j} + m)] - \text{erfc}[\eta_{\alpha} \beta |R_{\alpha} - r_{j} + m|] \right) \]

(13a)

\[ U_{lr,0}^{QQ} = -\frac{\sqrt{\pi}}{L_{x} L_{y}} \sum_{\alpha=1}^{N_{\alpha}} \sum_{\beta=1}^{N_{\beta}} \sum_{n} Q_{\alpha} Q_{\beta} \left( \frac{\exp[-\alpha^2 Z_{\alpha}^2]}{\alpha} + \sqrt{\pi} Z_{\alpha} \text{erf}[\alpha Z_{\alpha}] \right) \]

(13b)

\[ U_{lr,*}^{QQ} = \frac{1}{L_{x} L_{y}} \sum_{\alpha=1}^{N_{\alpha}} \sum_{\beta=1}^{N_{\beta}} \sum_{n} Q_{\alpha} Q_{\beta} \exp\left[-\frac{h^2 + u^2}{4\alpha^2}\right] \exp\left[i(h \cdot R_{\alpha} + u Z_{\alpha})\right] \]

(13c)

\[ U_{self}^{QQ} = \left( \frac{\eta_{\alpha}}{\sqrt{2\pi}} - \frac{\alpha}{\sqrt{\pi}} \right) \sum_{\alpha=1}^{N_{\alpha}} Q_{\alpha}^2 \]

(13d)

In Eq. (13a) we defined \( \eta_{\alpha \beta} \equiv \frac{\eta_{\alpha} \eta_{\beta}}{\sqrt{\eta_{\alpha}^2 + \eta_{\beta}^2}} \). Also here, note that long-range interactions have the same form as in the case of the mutual interaction between point charges, with the integrated Gaussian charges in place of the point charges.

5. Gaussian charge-point dipole interaction: \( U^{Q\mu} \)

As mentioned above, this term has never been reported in the literature. Its form is therefore derived in detail in the SM. Here we write \( U^{Q\mu} = U_{st}^{Q\mu} + U_{lr,0}^{Q\mu} + U_{lr,*}^{Q\mu} \) with

\[ U_{st}^{Q\mu} = \frac{1}{2} \sum_{\alpha=1}^{N_{\alpha}} \sum_{j=1}^{N_{j}} Q_{\alpha} \mu_{j} \cdot (R_{\alpha} - r_{j} + m) \]

\[ \times \left( \text{erfc}[\alpha R_{\alpha} - r_{j} + m] + \frac{2\alpha}{\sqrt{\pi}} |R_{\alpha} - r_{j} + m| \exp[-\alpha^2 |R_{\alpha} - r_{j} + m|^2] + \right. \]

\[ \left. \left( \text{erfc}[\eta_{\alpha} R_{\alpha} - r_{j} + m] + \frac{2\eta_{\alpha}}{\sqrt{\pi}} |R_{\alpha} - r_{j} + m| \exp[-\eta_{\alpha}^2 |R_{\alpha} - r_{j} + m|^2] \right) \right] \]

(14a)

\[ U_{lr,0}^{Q\mu} = \frac{\pi}{L_{x} L_{y}} \sum_{\alpha=1}^{N_{\alpha}} \sum_{j=1}^{N_{j}} Q_{\alpha} \mu_{j} \text{erf}[\alpha(Z_{\alpha} - z_{j})] \]

(14b)

\[ U_{lr,*}^{Q\mu} = -\frac{i}{L_{x} L_{y}} \sum_{\alpha=1}^{N_{\alpha}} \sum_{j=1}^{N_{j}} \int_{-\infty}^{\infty} \int_{-\infty}^{\infty} du \sum_{k \neq 0} Q_{\alpha} (\mu_{j} \cdot h + \mu_{j} u) \exp\left[-\frac{h^2 + u^2}{4\alpha^2}\right] \exp\left[i(h \cdot (R_{\alpha} - r_{j}) + u(Z_{\alpha} - z_{j}))\right] \]

(14c)

Once again, Gaussian charges behave, in the long-range, as point charges equal to the integrated Gaussian charge (compare Eqs. (12b) and (12c) with Eqs. (14b) and (14c)).
6. Point dipole-point dipole interaction: $U^\mu\mu$

The last energy contribution for the model discussed in this paper is the mutual interaction between point dipoles. The expression for the 2D PBC case can be derived as the expression in the 3D PBC case given in Ref. 24. The interaction energy can be divided as $U^\mu\mu = U^\mu\mu_{st} + U^\mu\mu_{ir,0} + U^\mu\mu_{ir,s} + U^\mu\mu_{self}$ and the single contributions are given by

$$U^\mu\mu_{st} = \frac{1}{2} \sum_{l=1}^{N_u} \sum_{l'=1}^{N_u} \sum_{j=1}^{N_d} \sum_{j'=1}^{N_d} \sum_{m} \frac{\mu_j \cdot S_{l,l'} \cdot \mu_{j'}}{|r_{l,l'} + m|^3} \left[ \text{erfc} \left[ \frac{\alpha}{\sqrt{2}} |r_{l,l'} + m| \right] + \frac{2\alpha}{\sqrt{\pi}} |r_{l,l'} + m| \exp \left[ -\alpha^2 |r_{l,l'} + m|^2 \right] \right] +$$

$$- \frac{\mu_j \cdot (r_{l,l'} + m)}{|r_{l,l'} + m^2|} \left[ \mu_{j'} \cdot (r_{l,l'} + m) \right] \frac{4\alpha^3}{\sqrt{\pi}} \exp \left[ -\alpha^2 |r_{l,l'} + m|^2 \right]$$

$$U^\mu\mu_{ir,0} = \frac{2\alpha\sqrt{\pi}}{L_z L_y} \sum_{l=1}^{N_u} \sum_{j=1}^{N_d} \mu_j^2 \exp \left[ -\alpha^2 z_{l,j}^2 \right]$$

$$U^\mu\mu_{ir,s} = \frac{1}{L_z L_y} \int_{-\infty}^{\infty} du \int_{-\infty}^{\infty} \sum_{k \neq 0}^{} \left( \mu_j \cdot h + \mu_j^* u \right) \left( \mu_j \cdot h + \mu_j^* u \right) \exp \left[ -\frac{h^2 + u^2}{4\alpha^2} \right] \exp \left[ i(h \cdot r_{l,j} + u z_{l,j}) \right]$$

$$U^\mu\mu_{self} = -\frac{2\alpha^3}{3\sqrt{\pi}} \sum_{l=1}^{N_u} \sum_{j=1}^{N_d} \mu_j^2$$

where the matrix $S_{l,l'}$ is defined by $S_{l,l'}^\xi = \delta_{\xi_x} - \frac{3(\xi_x + m^2)(\chi_l + \chi_{l'})}{|r_{l,l'} + m|^2}$ and $\xi, \chi_l \in \{x, y, z\}$. $\delta_{\xi_x}$ is the Kronecker’s delta function, which is 1 if $\xi = \chi$ and 0 otherwise.

B. Implementation details

The long-range, $k \neq 0$ terms introduced in the previous subsection are in general complex number due to the presence of the complex exponentials. Nonetheless, these can be transformed in purely real expressions exploiting the Euler’s formula, i.e., $\exp[i\phi] = \cos(\phi) + i\sin(\phi)$. Let us note first that the problematic long-range terms can be grouped in two sets based on similarities in their structure. The first set consists of the terms $U^\mu\mu_{ir,s}$, $U^{QQ}_{ir,s}$, $U^{QQ}_{ir,s}$ and $U^\mu\mu_{ir,s}$. The only difference among these terms arises from the different “charges” (in some cases the point charge $q_i$, in others the integrated Gaussian charge $Q_o$) or the scalar product between the dipole moment and the wavevector $\mu_j \cdot h + \mu_j^* u$ and the specific definitions of the distances at the exponent appearing in their expressions. A similar observation applies to the second set, composed by the terms $U^\mu\mu_{ir,s}$, $U^{QQ}_{ir,s}$. We discuss these two sets separately. About the former, exploiting the definition of the complex exponential, the parity of some of the trigonometrical functions with respect to the summation and integration interval and the trigonometrical expression of a difference, we obtain

$$U^{QQ}_{ir,s} = \frac{1}{L_z L_y} \int_{-\infty}^{\infty} du \sum_{k \neq 0} \exp \left[ -\frac{h^2 + u^2}{4\alpha^2} \right] \left[ Q_o \cos(h \cdot R_\alpha + u z_\alpha) \right]^2 + \sum_{\alpha=1}^{N_o} Q_\alpha \sin(h \cdot R_\beta + u Z_\beta)^2$$

$$U^{QQ}_{ir,s} = \frac{1}{L_z L_y} \int_{-\infty}^{\infty} du \sum_{k \neq 0} \exp \left[ -\frac{h^2 + u^2}{4\alpha^2} \right] \left[ \sum_{\alpha=1}^{N_o} Q_\alpha \cos(h \cdot R_\alpha + u Z_\alpha) \right]^2 + \sum_{\beta=1}^{N_o} Q_\beta \sin(h \cdot R_\beta + u Z_\beta)^2$$

$$U^{QQ}_{ir,s} = \frac{1}{L_z L_y} \int_{-\infty}^{\infty} du \sum_{k \neq 0} \exp \left[ -\frac{h^2 + u^2}{4\alpha^2} \right] \left[ \sum_{\alpha=1}^{N_o} Q_\alpha \cos(h \cdot R_\alpha + u Z_\alpha) \right]^2 + \sum_{\beta=1}^{N_o} Q_\beta \sin(h \cdot R_\beta + u Z_\beta)^2$$

$$U^{QQ}_{ir,s} = \frac{1}{L_z L_y} \int_{-\infty}^{\infty} du \sum_{k \neq 0} \exp \left[ -\frac{h^2 + u^2}{4\alpha^2} \right] \left[ \sum_{l=1}^{N_d} (\mu_j \cdot h + \mu_j^* u) \cos(h \cdot r_{l,j} + u z_{l,j}) \right]^2 + \sum_{J=1}^{N_d} (\mu_J \cdot h + \mu_J^* u) \sin(h \cdot r_{J,j} + u z_{J,j})$$

(16d)
The same kind of procedure can be performed on the second set noting, in addition, that $i \exp[ix] = \exp[i(x + \frac{\pi}{2})] = \cos(x + \frac{\pi}{2}) + i \sin(x + \frac{\pi}{2}) = i \cos(x) - \sin(x)$. In this way we obtain

\[
U^q_{\text{lr},*} = \frac{1}{L_x L_y} \int_{-\infty}^{\infty} du \sum_{k \neq 0} \exp\left[ -\frac{h^2 + u^2}{2u^2} \right] \left[ \sum_{i=1}^{N_p} q_i \sin(h \cdot r_i + uz_i) \right] \left[ \sum_{J=1}^{N_d} (\mu_j \cdot h + \mu_j^* u) \cos(h \cdot r_J + uz_J) \right] +
\]

\[
- \sum_{i=1}^{N_p} q_i \cos(h \cdot r_i + uz_i) \right] \left[ \sum_{J=1}^{N_d} (\mu_j \cdot h + \mu_j^* u) \sin(h \cdot r_J + uz_J) \right] \right] \right] (17a)
\]

\[
U^q_{\text{lr},*} = \frac{1}{L_x L_y} \int_{-\infty}^{\infty} du \sum_{k \neq 0} \exp\left[ -\frac{h^2 + u^2}{2u^2} \right] \left[ \sum_{\alpha=1}^{N_g} Q_\alpha \sin(h \cdot R_\alpha + uZ_\alpha) \right] \left[ \sum_{J=1}^{N_d} (\mu_j \cdot h + \mu_j^* u) \cos(h \cdot r_J + uz_J) \right] +
\]

\[
- \sum_{\alpha=1}^{N_g} Q_\alpha \cos(h \cdot R_\alpha + uZ_\alpha) \right] \left[ \sum_{J=1}^{N_d} (\mu_j \cdot h + \mu_j^* u) \sin(h \cdot r_J + uz_J) \right] \right] \right] (17b)
\]

The expressions in Eqs. (16) and (17) are purely real and can be handle with standard numerical libraries and operators.

C. Additional degrees of freedom

In the interaction potential, the induced dipoles represent the instantaneous polarization of a molecule or an ion, which fulfill the following expression at each timestep:

\[
\mu_j = \alpha_j E_j = -\alpha_j \nabla_j V(r) \quad (18)
\]

where $\alpha_j$ is the polarizability of atom $J$ and $E_j$ the electric field felt by the atom. The latter depends not only on the charges of the system but also on the other induced dipoles. In practice, instead of solving self-consistently the set of equations 18, most of the implementations are made by adding a polarization term to the energy, which is given by

\[
U_{\text{pol}} = \sum_{J=1}^{N_d} \frac{\mu_j^2}{2\alpha_j} \quad (19)
\]

In the absence of electrodes, the induced dipoles are then computed by minimizing $U_1 = U_{\text{elec}} + U_{\text{pol}}$. However, in MetalWalls we also have to account for the fact that electrodes are set at constant potential. This condition reads

\[
V(r) = \Psi_{\Omega_{\pm}} \quad (20)
\]

where $r$ is a point in the regions $\Omega_{\pm}$ occupied by the positive or negative electrode and $\Psi_{\Omega_{\pm}}$ is the potential set in the corresponding electrode. The condition can be reformulated in terms of the Gaussian charges as

\[
\frac{\partial U_{\text{elec}}}{\partial Q_\alpha} = \Psi_\alpha. \quad (21)
\]

with $\Psi_\alpha = \Psi_{\Omega_{\pm}}$ depending on the electrode in which $\alpha$ is located. In practice, in the absence of induced dipoles, this problem is solved by minimizing $U_2 = U_{\text{elec}} - \sum_{\alpha=1}^{N_g} \Psi_\alpha Q_\alpha$.

Now the two systems of additional degrees of freedom are coupled: Since $U_{\text{elec}}$ depends on both $\{\mu_j\}_{j=1}^{N_d}$ and $\{Q_\alpha\}_{\alpha=1}^{N_g}$, the Gaussian charges values will depend on the induced dipoles and vice versa. The total quantity which is minimized is $U = U_{\text{elec}} + U_{\text{pol}} - \sum_{\alpha=1}^{N_g} \Psi_\alpha Q_\alpha$.

III. COMPUTATION OF INDUCED DIPOLES AND ELECTRODE CHARGES: THE PRECONDITIONED CONJUGATE GRADIENT

Induced dipoles and electrode charges are computed in the Born-Oppenheimer approximation: Their value is determined through a minimization procedure of the energy function with respect to the additional dynamical variables modeling these quantities. Contrary to existing literature, in which either the induced dipoles or the electrode charges were to be computed, in this paper the Born-Oppenheimer condition has to be satisfied for the set of additional dynamical variables which includes both the induced dipoles $\mu$ and electrode charges $Q$. In other words, given the interaction energy $U(\mu, Q|r, R)$ as a function of $\mu$ and $Q$ and which depends parametrically on the electrolyte particles positions $r$ and on the geometry of the electrodes placed at positions $R$, the values of the dipoles and the electrode charges are found, for a given value of the parameters $r$ and $R$, as the solution of the equation

\[
\arg \min_{\mu, Q} U(\mu, Q|R, r) \quad (22)
\]

This optimization problem, which is now formulated in the space of dimension $3N_d + N_g$, can be very demanding from a computational point of view and many different approaches have been proposed to tackle its solu-
For models such as those considered in this paper, in which the dependence of the energy function on the additional variables is quadratic, the conjugate gradient method provides a good balance between accuracy and efficiency and it is the preferred choice nowadays. However, as the minimization procedure usually takes 90% of the simulation time in a typical Born-Oppenheimer simulation, methods and algorithms to improve the speed of convergence of the conjugate gradient procedure (or even replace it by a less costly method) are still object of investigation.

For the minimization of quadratic functions of the form \( U = \frac{1}{2} x^T A x + b^T x + c \) — which corresponds to finding the solution of the linear system of equations \( A x + b = 0 \) — preconditioning the symmetric, positive-definite matrix \( A \) has proven to be an effective tool for improving the speed of convergence of the conjugate gradient algorithm. Indeed, the convergence of the algorithm depends mainly on the condition number of the matrix \( A \) (see Ref. 43) defined as the ratio between its maximum and minimum eigenvalues. The preconditioning procedure is a technique in which the optimization problem is transformed in an equivalent one, i.e., one that has the same solution, but where a matrix with a lower condition number is involved. In practice, this is done by noting that the solution of the equation \( A x + b = 0 \) is the same of \( P^{-1} (A x + b) = 0 \) where the preconditioner \( P \) is a matrix of the same dimensions of \( A \). If the condition number of the matrix \( P^{-1} A \) is lower than that of \( A \) the algorithm will be more efficient in solving the problem.

Unfortunately, a general strategy for finding a good preconditioner does not exist and the choice of an effective matrix \( P \) is highly system dependent. A quite common approach — even though sometimes ineffective — is to choose \( P = \text{diag} A \), the so-called Jacobi preconditioner. This choice has the double advantage that, on the one hand, it is not necessary to store an additional entire matrix, but just a vector and, on the other, that the computation of \( P^{-1} \) is trivial. If effective, the Jacobi preconditioner is particularly well-suited for minimization problems in high-dimensional spaces. Other choices, sometimes more effective but at the same time more demanding numerically and in terms of required memory, are for example based on approximate inverses, (incomplete) Cholesky factorizations and domain decompositions.

To investigate the effectiveness of the Jacobi preconditioner for electrochemical problems we study the properties of the matrix associated to the system described in Ref. 30. This system consists of constant potential aluminium electrodes in contact with a molten salt (pure LiCl) in which the electrolyte ions are represented using a polarizable force field. While being very simple, this system, which can be effectively handled numerically, captures all the important features of electrochemical systems. A thorough study of the matrix requires its storage and analysis, a process that usually requires an order \( N^3 \) of operations, where \( N \times N \) (with \( N = 3 N_d + N_g \)) is the size of the \( A \) matrix. This is affordable for our chosen system.

For electrochemical systems in which either the electrode charges and the induced dipoles are considered as dynamical variables, the solution vector \( x \) can be ordered as

\[
x = (\mu_1^z, \mu_2^z, \ldots, \mu_{N_d-1}^z, \mu_{N_d}^z, Q_1, \ldots, Q_N)
\]

With this choice, the matrix \( A \) is represented in block form, as

\[
A = \begin{pmatrix}
\nabla^2_{\mu_1^z} U & \ldots & \nabla^2_{\mu_{N_d}^z} U \\
\nabla^2_{Q_1} U & \ddots & \nabla^2_{Q_N} U
\end{pmatrix}
\]

where \( U \) is the interaction energy given in Eq. (8). The diagonal of this matrix, which represents the Jacobi preconditioner will then be given by

\[
P = \text{diag} A
\]

\[
= (\nabla^2_{\mu_1^z} U, \ldots, \nabla^2_{\mu_{N_d}^z} U, \nabla^2_{Q_1} U, \ldots, \nabla^2_{Q_N} U)
\]

The analytic expression for \( P \) can be obtained starting from the Ewald decompositions given in the previous section. Assuming nearest-image convention, we then have

\[
\nabla^2_{\mu_j^z} U = \frac{2}{L_x L_y} \sum_{k \neq 0} \int_{-\infty}^{\infty} \text{du} \exp \left( -\frac{h^2 + u^2}{4\alpha^2} \right) (\kappa \xi)^2
\]

\[
+ \delta \xi \frac{4 \sqrt{\pi}}{L_x L_y} - \frac{4 \alpha^3}{3 \sqrt{\pi}} + \frac{1}{\alpha_j^2}
\]

\[
\nabla^2_{Q_{\alpha} U} = \frac{2}{L_x L_y} \sum_{k \neq 0} \int_{-\infty}^{\infty} \text{du} \exp \left( -\frac{h^2 + u^2}{4\alpha^2} \right)
\]

\[
- \frac{2}{\alpha} \frac{\sqrt{\pi}}{L_x L_y} + \frac{2}{\sqrt{\pi}} \left( \frac{\eta_\alpha}{\sqrt{\alpha}} - \alpha \right)
\]

for \( i = 1, \ldots, N_d, \xi \in \{x, y, z\} \) and \( \alpha = 1, \ldots, N_g \). In the previous equation we have also defined the vector \( \kappa = (2\pi L_x, 2\pi L_y, u) \) and we remark the difference between the Ewald smearing parameter \( \alpha \), the polarizability \( \alpha_j \), the electrolyte polarizable atom \( j \) (defined in Eq. (18)) and the index \( \alpha \) of the Gaussian-distributed charge as in \( Q_\alpha \) and \( \eta_\alpha \). Also, \( \delta \xi \) is the Kronecker delta. In Eq. (26), it is possible to recognize in both expressions the two terms arising from the long-range part of the Ewald decomposition \( (k = 0 \text{ and } k \neq 0) \) and the term arising from the self-interaction term. In the expression relative to the induced dipoles, Eq. (26a), also the contribution due to the so-called “self-polarization” term is present.

In what follows, we show results for a 10 ps simulation of the system of aluminum electrodes and polarizable LiCl. The polarizability of the lithium ion is set to \( \alpha_{Li} = 0.13 \text{Å}^3 \), while the one of chloride is set to \( \alpha_{Cl} = 2.96 \text{Å}^3 \) as reported in previous work. Along the
simulation, we sample the configuration of the electrolyte every 100 fs and we compute the condition number with \((P^{-1}A)\) and without \((A)\) preconditioning. Results are shown in Figure 1. Note how the Jacobi preconditioner reduces the condition number of the problem by one order of magnitude. It is also interesting to check the behavior of the conjugate gradient algorithm for the configurations for which we computed the condition number of the matrix. Figure 2 shows the number of iterations needed for convergence of the algorithm for these configurations, again without and with preconditioning. As expected, Figures 1 and 2 confirm that to a lower condition number of the matrix corresponds a lower number of iterations of the conjugate gradient algorithm to reach convergence.

For larger systems such as the one discussed in Section V, where the study of the condition number is unfeasible due to the high dimensions of the matrix, the effectiveness of the Jacobi preconditioner can still be investigated by looking at the number of iterations to reach convergence. In Figure 3 we present this indicator for the LiCl-Al system, demonstrating that the Jacobi preconditioner reduces the number of iterations needed for convergence of the algorithm by roughly 50%.

IV. VALIDATION ON SIMPLE SYSTEMS

A. Single dipole between single atom electrodes

A difficulty to validate the implementation of the expressions provided in Section II is that no other molecular dynamics code can be used to provide reference results. However, most of the terms were already derived and validated for 2D \(U_{qq}, U_{qQ}, U_{QQ}\) or 3D \(U_{qq}, U_{qQ}, U_{q\mu}, U_{QQ}\) and \(U_{\mu\mu}\) PBCs. The main validation test therefore
concerns the $U^{Q\mu}$ term for which no results are available. To this end, we take advantage from the equivalence between an explicit dipoles and a set of two charges in the following.

In a first step, two systems composed of two single-atom electrodes placed in a box of dimension $L_x = L_y = 26.46 \, \text{Å}$ and $L_z = 52.92 \, \text{Å}$ were setup. In both cases, the first electrode is located at coordinates $\left( \frac{L_x}{2}; \frac{L_y}{2}; 0 \right)$, while the second one is at $\left( \frac{L_x}{2}; \frac{L_y}{2}; L_z \right)$. These electrodes atoms have a Gaussian charge distribution width $\eta = 0.4 \, \text{Å}^{-1}$, and the electrodes are held at a constant potential difference of $1 \, \text{V}$. In the first system, which is referred to as a “point dipole system” in the following, a single atom is located in between the electrodes at coordinates $\left( \frac{L_x}{2}; \frac{L_y}{2}; z \right)$. This atom carries no charge but can be polarized, its polarizability being set to a value of $0.15 \, \text{Å}^3$. The energy and the force acting on this atom are computed for different $z$ coordinates. The second system mimics the first one, by introducing two central atoms $d_\mu = 0.53 \, \text{Å}$ apart. They carry opposite charges, thus forming an explicit dipole which is placed in between the two electrodes. This system is thus called a “two-charges system”. The charges on the atoms are taken from the value of the dipole moment $\mu$ obtained from the point dipole system at a similar $z$ position. Charges $+q_\mu$ and $-q_\mu$ are assigned to the atoms of coordinates of $\left( \frac{L_x}{2}; \frac{L_y}{2}; z - \frac{d_\mu}{2} \right)$ and $\left( \frac{L_x}{2}; \frac{L_y}{2}; z + \frac{d_\mu}{2} \right)$, respectively. These atoms are not interacting with each other, so that from a physical point of view, the two systems are identical, but in the first setup the interaction potential only contains $U^{QQ}$ and $U^{Q\mu}$ terms, while the second one contains $U^{QQ}$ and $U^{\mu q}$. The latter were already validated in numerous studies, so this allows us to directly test the new implementation.

For the Ewald summation, the cut-off distance for real space interactions is set to $r_{\text{cut}} = 12.70 \, \text{Å}$. The relative error below which real-space and reciprocal-space terms are not included in the summation are set to $10^{-12}$. Electroneutrality of the systems is enforced when computing the partial charges on the electrodes as described in Ref. 41. To compute energies and forces, the preconditioned conjugate gradient described in the previous section is used. The iterative process stops when: [residual] $< \text{tolerance} \times \sqrt{\text{N} \text{atoms}}$, where tolerance is set at $10^{-12}$. 151 calculations are performed with the $z$ coordinate varying from $2.96 \, \text{Å}$ to $49.96 \, \text{Å}$, with a step length of $0.31 \, \text{Å}$ to sample a large variation of the induced dipole moment.

Figure 4 shows the evolution of the energy (top) and of the force along $z$ (bottom) for the point dipole system (black curve) and the two-charges system (orange dots). In the latter case, the force is obtained by summing the individual forces acting on each atom. The results for the two systems agree very well, with relative errors lower than $10^{-5}$ for the energies and lower than $2.5 \times 10^{-2}$ for the forces. Additional tests were performed with varying values for the cut-off distance for the real space interactions, showing similar level of agreement.

B. Single dipole between 100 atoms electrodes

In order to compare more precisely the charge distribution on the electrodes, another couple of systems were built in which the electrodes are made of a single plane of 100 atoms placed on a square lattice with a cell parameter of $2.94 \, \text{Å}$. There again, the point dipole and the two-charges systems were set up for comparison. The parameters (electrode atoms Gaussian width, applied potential, Ewald real space cut-off, convergence of the conjugate gradient) were set similarly as for the single atom electrode tests. Figure 5 shows the charges on the electrode atoms for the two systems. Here also, a very good agreement is obtained, with a relative error lower than $10^{-5}$, which confirms the validity of the expression derived for $U^{Q\mu}$ and its implementation in MetalWalls.
FIG. 5. Comparison of the charge on each atom of the electrodes obtained for the point dipole and the two-charges system between 100 atoms electrodes with an applied potential difference of 1 V. Electrode atoms are labeled by their index, with atoms 1 to 100 belonging to the first electrode and atoms 101 to 200 belonging to the second one. The maximum of the relative error obtained is lower than $10^{-8}$.

V. VALIDATION ON A REALISTIC CAPACITOR

A. Simulation setup

Once the model has been validated for static calculations on model systems, it is also important to test the software on realistic cases. We have therefore chosen an already well-characterized system, composed of an ionic liquid between two graphite electrodes. This choice was made because, for such a system, it is possible to account in a mean-field way for polarization effects without including induced dipoles, by using reduced charges for all the ionic liquid atoms.\(^{21}\) Such an approach was shown to yield correct structural properties, which provides us with a reference to compare with our polarizable force field simulations (other properties, in particular dynamic ones, may vary much more than the structural ones). We study an ionic liquid consisting of 322 ion pairs of 1-ethyl-3-methylimidazolium bis(trifluoromethylsulfonyl)imide (EMIM-TFSI). The two electrodes are at a distance of 10.9 nm and made of three parallel graphene sheets separated by 0.335 nm. The total dimensions of the box are $L_x = 3.408$ nm, $L_y = 3.689$ nm and $L_z = 11.38$ nm. Periodic boundary conditions have been applied in the $x$ and $y$ direction. A typical snapshot of the system is shown in Figure 6.

The CL&P\(^{48,49}\) force-field is used for the non-polarizable simulation, with atomic charges scaled by a factor 0.8, as suggested in previous studies of the bulk liquid.\(^{50-52}\) The polarizable simulations use an upgrade of the force-field (CL&Pol),\(^{23,53,54}\) in which all atoms are polarizable except the hydrogen atoms. Note however that in the initial parameterization by Goloviznina et al.,\(^{23}\) the polarization effects were handled using Drude oscillators.\(^{55}\) The corresponding parameters were therefore replaced by a polarizability in order to be used in our induced dipoles scheme. Atomic polarizabilities were taken from Schröder\(^{56}\) following the CL&Pol methodology. In addition, short-range effects are handled a bit differently since we do not use Thole functions\(^{57}\) for damping dipole-dipole interactions at short range. Instead, we use the conventional OPLS\(^{58}\) rule for intramolecular charge-dipole and dipole-dipole interactions, \textit{i.e.} they are set to 0 for nearest and second-nearest neighbors, while they are scaled by a factor 0.5 for atoms separated by two bonds. Concerning the intermolecular interactions, the charge-dipole interactions are damped at short-range using Tang-Toennies functions\(^{59}\)

\[
g^{ij} (r_{ij}) = 1 - c_D^{ij} \exp(-b_D^{ij} r_{ij}) \sum_{k=0}^{4} \frac{(b_D^{ij} r_{ij})^k}{k!} \quad (27)
\]

where $b_D^{ij}$ sets the range of the damping effect and $c_D^{ij}$ the strength of the ion response.\(^{22}\) All atom pairs $ij$ have the same damping functions parameters, $b_D^{ij} = 1.06 \text{Å}^{-1}$ and $c_D^{ij} = 0.53 \text{Å}$. Simulations of the bulk liquid were performed in order to check that the so-modified polarizable force field yielded similar density, structural and diffusion properties as the original parameterization by Goloviznina et al.\(^{23,53}\) The Lennard Jones parameters for the electrodes atoms are set to $\epsilon = 0.23 \text{kJ/mol}$ and $\sigma = 3.37 \text{Å}$ as in previous works\(^{60}\). The Lennard-Jones interactions are cut at 14.8 Å. The cut-off distance for real space interactions in the Ewald summation is set at $r_{\text{cut}} = 17 \text{Å}$. The relative error below which real-space term and reciprocal-space are not included in the summation is set to $10^{-5}$.

The initial configuration was constructed using the PACKMOL software.\(^{61}\) In order to obtain the correct liquid density, the electrodes were allowed to move in the $z$ direction for few nanoseconds, by applying a pressure of 0 bar on both sides (free piston). When the electrode positions are stabilized, a bulk density of 1576 kg/m$^3$ is reached, which is close to the experimental value of 1526 kg/m$^3$ at 293.5 K.\(^{62}\) Starting from this setup, a long simulation was performed using the non-polarizable interaction potential, from which three starting points were extracted. Three independent production trajectories were then performed using both the polarizable and non-polarizable potentials. These simulations were carried out for 20 ns, with a timestep of 2 fs and a temperature of 398 K. The latter value was chosen in order to accelerate the dynamics of the system and to converge the structure of the liquid within shorter simulations, since the objective of this work is to validate the code and not to do a specific study of the system. The two electrodes potential were fixed to 0 V. The systems were simulated in the NVT ensemble, using a Nosé-Hoover thermostat chain of length 5 with a relaxation time of 500 fs. In the following, the results were obtained by averaging over the three simulations for both polarizable or non-polarizable force-field.

B. Results

![Graph showing density profiles](image)

FIG. 7. Density profile for simulations including or not the polarization effects. Dotted lines show the position of the electrodes. The top panel shows the total density profiles, while the bottom panels show the density of anions and cations at the interfaces.

Figure 7 shows the number density profiles along the $z$ direction for both force fields. For each ion type, the density is obtained by summing the averaged density of all atoms of the molecule, and normalized by the number of atoms of the ion. The top panel shows the total number of ions, and the bottom panels correspond to the EMIM$^+$ and TFSI$^-$ ions zoomed in at the interfaces. Electrode positions are shown as blue light vertical lines. Our results agree with the ones obtained in previous works, mainly with non-polarizable interaction potentials,\textsuperscript{63–68} except for one study.\textsuperscript{69} The structure is characterized by a strong ordering at the interface, with the formation of several layers in the vicinity of the electrode. We can conclude from these comparisons that our simulations with the polarizable force field yield the correct structure of the system. This indirect validation on a realistic system and the validations shown in the previous sections demonstrate the accuracy of the implementation of the polarization effects in conjunction with Gaussian charges in MetalWalls.

Despite yielding the same structure, it is likely that the different charge distributions in the two models will result in different electrostatic potentials at the interface. The latter can be computed as

$$\Psi(z) = \Psi_q(z) + \Psi_\mu(z)$$  \hspace{1cm} (28)$$

where $\Psi_q$ is the potential due to the charge distribution (from the electrolyte or electrode atoms) $\rho_q$, and $\Psi_\mu$ is the one due to the $z$-component of the induced dipole moments distribution $\rho_\mu$ at a given position $z$,

$$\Psi_q(z) = \Psi_q(z_0) - \frac{1}{\epsilon_0} \int_{z_0}^{z} dz' \int_{-\infty}^{z} dz'' \rho_q(z'')$$

$$\Psi_\mu(z) = \Psi_\mu(z_0) + \frac{1}{\epsilon_0} \int_{z_0}^{z} dz' \rho_\mu(z')$$  \hspace{1cm} (29)$$

where $\epsilon_0$ is the vacuum permittivity. The profiles are obtained by taking the reference point ($z_0$) inside the left electrode, so that $\Psi(z_0) = 0$. The variations of $\rho_q$, $\rho_\mu$, and $\Psi$ are shown for the non-polarizable and the polarizable models on Figure 8. The two charge distributions are of course very similar since the structure of the system does not differ much with the interaction potential. The main difference is the amplitude of the oscillations, which is reduced for the non-polarizable case due to the use of
scaled charges. The dipole density also displays some peaks, the most intense corresponding to the layer of liquid closest to the electrodes in which a net polarization is observed. Interestingly, the positions of the peaks do not coincide with the ones obtained for the charge density and the oscillations seem to extend towards a longer distance away from the electrode, despite the shorter range of interactions involving dipoles. This is probably a consequence of the layered structure, which results in local polarization inside the layers.

A consequence of polarization effects is thus that the potential profiles differ markedly between the two cases. Although the peaks appear at the same positions, they have very different intensities. Consequently, the two models have different potentials of zero charge (PZC). This quantity can in principle be measured experimentally; it is defined as the potential difference between the bulk and the electrodes when the latter carry no net charge:

$$\Delta \Psi_{PZC} = \Psi_{\text{electrode}} - \Psi_{\text{bulk}}$$

In constant potential simulations, the PZC is readily obtained from the bulk potential on the profiles shown on Figure 8. We obtain a value of -0.16 V for the non-polarizable case and -0.52 V for the polarizable case, respectively. Our results show that this quantity mostly reflects the charge density distribution of the model, and not the structure of the liquid. It should therefore not be used to test the ability of a potential to predict the double-layer structure of a system with respect to experiments.

A much more robust test is to compute the capacitance of the system from the fluctuations of the total charge of the electrode according to Ref 41. Note that the contribution to the capacitance of the empty capacitor (i.e., in the absence of liquid between the electrodes) is also taken into account. Figure 9 shows the probability to have a given total charge on the electrode for the two models. The distributions are very similar, which results in similar capacitances of 2.35 ± 0.79 µF/cm² for the non-polarizable case, and 2.03 ± 0.42 µF/cm² for the polarizable case. The fact that the same structure of the liquid yields similar results is a further (indirect) proof that the charges induced on the electrodes are correctly determined.

VI. ADSORPTION OF REDOX-FUNCTIONALIZED SPECIES ON GRAPHITE ELECTRODES

In this last section, we focus on an example for which polarization effects can only be represented via a suitable interaction potential. We simulate biredox ionic liquids, a new class of systems in which redox-active moieties are grafted to both the cationic and the anionic species. Such systems have shown very promising results when employed in supercapacitors since they allow to markedly increase the amount of electricity stored in...
the devices with respect to conventional electrolytes such as ionic liquids.\textsuperscript{71,72} However, their electricity storage mechanisms, in particular the interplay between capacitive and faradaic processes at the electrodes interfaces, remain to be solved.\textsuperscript{73} Molecular dynamics can provide a microscopic view of the double-layer in such systems, but no accurate force field was previously available in the literature. Recently, we have developed a polarizable interaction potential for the redox active groups TEMPO and anthraquinone (AQ) based on electronic structure calculations,\textsuperscript{35} which opens the way towards the simulation of biredox ionic liquids.

Here we simulate a system made of 10 biredox ionic liquid pairs, namely AQ-TFSI\textsuperscript{−} and TEMPO-EMIM\textsuperscript{+}, mixed with 50 BMIM\textsuperscript{+} TFSI\textsuperscript{−} ion pairs and 1624 acetonitrile molecules. The composition of the system was chosen according to previous experimental studies.\textsuperscript{71} Each simulation was performed for 3.3 ns under a constant potential difference of either 0, 1 or 2 V in the NVT ensemble (\(T = 298\) K). Two dimensional (2D) periodic boundary conditions were used, with box dimensions \(L_x = 34.0\) Å and \(L_y = 36.0\) Å, respectively. The electrodes separation has been fixed to 154.2 Å for which the density of the system in the middle of the box corresponds to the bulk value as in the previous section.

Although an extensive study of the system is out of the scope of the present work, we illustrate the software abilities by studying the adsorption mechanisms of redox-active molecules on the graphite electrodes, which impacts for example the kinetics of electron transfer. Firstly, we observe from the trajectory that the affinity of the ions for the electrode is similar for non-grafted and grafted species (within the statistical accuracy of the simulation, which is rather limited by the small number of ions in the simulation cells). However, as illustrated on the typical snapshots shown on Figure 10, the grafted ions tend to interact more strongly with the carbon surface through their redox-active moiety, leaving the ionic part of the molecule slightly behind the first plane. We can therefore expect a facilitated tunneling of the electrons when such liquids are used in supercapacitor devices.

In order to get a more quantitative analysis, we study the orientation of the species at the interface. For each ion, we define a structural coordinate to analyze its orientation at the electrode. For the cation we choose the normal to the plane of the imidazolium ring and for the anion the vector connecting the two sulfur atoms and the projection of the normal to the plane of the positive electrode at different potential difference for AQ-TFSI\textsuperscript{−} and TFSI\textsuperscript{−}. The error bars correspond to the standard deviation.

![Figure 10](image1.png)

**FIG. 10.** Snapshots of the simulation showing the adsorption of the functionalized ionic liquid anion (left) and cation (right).

![Figure 11](image2.png)

**FIG. 11.** Top: Angle between the normal to the plane of the imidazolium ring and the normal to the negative electrode at different potential difference for TEMPO-EMIM\textsuperscript{+} and BMIM\textsuperscript{+}. Bottom: Angle between the vector connecting the two sulfur atoms and the projection of the normal to the plane of the positive electrode at different potential difference for AQ-TFSI\textsuperscript{−} and TFSI\textsuperscript{−}. The error bars correspond to the standard deviation.
We can see that the redox functionalization of the ionic liquid changes the orientation at the electrode. In both cases, while the bare ions lie parallel to the surface, they become perpendicular in the biredox species. This effect reflects again the stronger affinity between graphite and redox moieties, which forces the ionic part of the molecules to adapt and change their orientation. Such an effect was not expected since the charged species are stabilized by the image charges in the metallic electrodes. The planar shape of the anthraquinone provides a lead for its affinity with graphite, since it may allow a more efficient packing, however it is not the case of the TEMPO group. Another possibility is that the polar C=O function of anthraquinone and N=O function of the TEMPO display a large interaction with the fluctuating charge of the metal. However, this would require costly additional simulations, and we leave this for future work.

VII. CONCLUSION

The simulation of electrochemical systems requires to account precisely for polarization effects at the electrode/electrolyte interface. By combining two sets of additional degrees of freedom, the induced dipoles in the liquid and the partial charges inside the electrode, MetalWalls now allows to use state-of-the-art force fields in such simulations. In this work, we have derived the equations enabling the simulation of systems including point charges, Gaussian charges and induced dipoles. Due to the long range nature of these interactions, it is necessary to use the Ewald summation method, which consists in splitting the interaction potential in two sums, one in the real space and the other in reciprocal space. The expressions are provided for 2D periodic boundary conditions, which corresponds to the common setup for simulating electrochemical systems. The additional degrees of freedom are computed through a conjugate gradient minimization procedure, whose computational cost can be decreased through the use of simple preconditioning techniques. Note that another common approach, which consists in using 3D PBCs with an additional (large) vacuum region and adding a correction term for the slab geometry is also implemented in MetalWalls. Although this allows to reduce the computational time, as noted in a recent work this setup should be used with caution when used for simulating constant potential electrodes.

We have then validated the implementation of the formulae for 2D PBC. This cannot be done through a simple comparison with other codes since the above features are not yet included elsewhere. We therefore focused first on a simple system consisting of a single dipole between electrodes, for which the dipole can be replaced by a couple of explicit charges, providing reference results. We then compare the results obtained on lengthy simulations of a realistic system, the EMIM-TFSI ionic liquid in contact with graphite electrodes. This system was chosen because it is possible to account effectively for the polarization of the electrolyte using rescaled charges. The simulations yield similar structure for the interfacial liquid and the capacitance as expected.

The capabilities of MetalWalls were then demonstrated by simulating a system for which no non-polarizable force field exists. It consists in a pair of redox-functionalized ions dissolved in a conventional electrolyte, again in contact with graphite electrodes. The structural analysis showed that the redox moieties impact the adsorption of molecules at the surface of the electrode, providing a basis for the understanding of the properties of these electrolytes in future works.

Although this work only focused on simple electrode geometries, the methods presented here are general and can be applied to any electrode geometry. MetalWalls can therefore be used to simulate complex systems, such as slat nanopores, carbide-derived carbons, etc. It is parallelized using MPI for conventional CPU-based high-performance computers and OpenACC for GPU, so that typical system sizes of 10,000 to 50,000 atoms can be simulated routinely. MetalWalls is therefore a valuable tool for the characterization of the structure of the electrolyte in many electrochemical devices, such as supercapacitors, batteries, electrocatalysts, etc. The next challenge will be to develop the adequate force fields for such systems. There are currently many developments aiming at accounting for the different metallicities of the electrode materials, but there is no systematic recipe for choosing the corresponding parameters. Approaches based on high-level ab initio calculations will certainly provide an important input towards this direction.

SUPPLEMENTARY MATERIAL

The supplementary material contains the complete derivation of 2D-PBC Ewald decomposition for energy and forces of Gaussian charges-point dipoles term and the Ewald decomposition for the forces.

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DATA AVAILABILITY

The MetalWalls code is openly available in the repository https://github.com/ampere2/metalwalls.


This is the author's peer reviewed, accepted manuscript. However, the online version of record will be different from this version once it has been copyedited and

Number of Iterations

Configuration

No Preconditioning

Jacobi Preconditioner
$\Psi_{bulk, polarizable} = 0.52 \text{ V}$

$\Psi_{bulk, non polarizable} = 0.16 \text{ V}$
Non polarizable

Polarizable