A practical guide to biologically-relevant molecular simulations with charge scaling for electronic polarization

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Molecular simulations can elucidate atomistic-level mechanisms of key biological processes, which are often hardly accessible to experiment. However, the results of the simulations can only be as trustworthy as the underlying simulation model. In many of these processes, interactions between charged moieties play a critical role, and current empirical force fields tend to overestimate such interactions, often in a dramatic way when polyvalent ions are involved. The source of this shortcoming is the missing electronic polarization in these models. Given the importance of such biomolecular systems, there is a great interest in fixing this deficiency in a computationally inexpensive way, that is without employing explicitly polarizable force fields. Here, we review the electronic continuum correction (ECC) approach, which accounts for electronic polarization in a mean field way, focusing on its charge scaling variant. We show that by pragmatically scaling only the charged molecular groups, we not only qualitatively improve the charge–charge interactions without extra computational costs, but also benefit from decades of force field development on biomolecular force fields.

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I. INTRODUCTION

Calcium signaling affects a plethora of cellular processes with specific proteins designed by nature as Ca$^{2+}$ binders. A necessary condition for calcium to act as a signaling agent is that the interaction with its binding partner has to be weak enough (i.e., below ten $k_B T$) so that it can bind and unbind at biologically relevant timescales. This is an imperative condition that needs to be satisfied by any molecular simulation that aspires to faithfully model processes involving calcium signalling. However, the widely used classical non-polarizable force fields tend to dramatically overbind calcium and other ions of high charge density (such as magnesium or lithium) not only to proteins and peptides, but also to phospholipids and nucleic acids. Similar overbinding is observed for other ions too, and the issues are further manifested in overestimated salt bridge strength between charged protein residues. Attempts to cure the overbinding problem by introducing an \textit{ad hoc} steric repulsion term between the binding partners may improve the situation, but provide at best the right result for an often unphysical reason. Physically, the overbinding originates from the lack of electronic polarization. The fast electrons should always be at equilibrium with nuclei positions and hence effectively screen the ion–ion Coulomb interaction. This missing screening causes the observed disbalance between ion–water and ion–counterion interactions in non-polarizable force fields, which can in principle be fixed by explicitly introducing electronic polarizability. This way, one might reproduce realistic behavior by incorporating missing physics in the model. However, due to issues connected with parametrization, numerical stability (i.e., the so called polarization catastrophe), and increased computational costs, biomolecular simulations with explicitly polarizable force fields are still rare.

Here, we address the question whether one can get a satisfactorily accurate behavior for (almost) correct reasons without extra computational cost. As demonstrated below, the answer is \textit{yes}, namely by including electronic polarization effects in a mean field way via charge scaling. In this Perspective, we aim to give an overview of recent efforts and strategies towards the development of scaled-charge force fields within the electronic continuum correction (ECC) approach, with a focus on applications in systems of biological interest. We want to provide a cookbook on how one can develop, test, and use such force fields, as well as highlight the potential pitfalls, open questions, and limitations of such an approach. We will first review the various existing strategies used to correct ion binding artefacts and
ion pairing in simulations, including the ECC strategy. We will then present the different steps towards the application of ECC framework to increasingly complex biosystems, starting from simple salt solutions all the way up to protein- and lipid- containing systems. While we will do our best to mention all contributions to this field, a particular focus will be put on the work performed in our group.

II. STRATEGIES TO IMPROVE ION BINDING AND PAIRING IN MOLECULAR DYNAMICS SIMULATIONS

A. Explicit treatments of electronic polarization

One strategy to improve the aforementioned overbinding artefacts is to explicitly include electronic polarization into classical force fields. This can be done in different ways, the most common approaches being the induced point dipole, the fluctuating charge, and the Drude oscillator models. These approaches and their applications in the simulation of electrolytes, ionic liquids, or biomolecular systems have been reviewed elsewhere and a detailed description thereof is outside the scope of the present review. Polarizable force fields have allowed for more realistic simulations of ionic liquids and electrolytes. Focusing on the interactions of ions with biomolecules, they have been successfully used to investigate, for instance, the geometry and relative stability of Mg$^{2+}$ binding modes to phosphate-based species, and the cation specificity in proteins. Polarizable force fields have also been shown to better capture the strength of salt bridges.

However, despite these significant advances and recent developments, polarizable force fields are not yet broadly used by the biosimulation community. The vast majority of simulations of complex biomolecular systems are still performed with non-polarizable force fields, even if they suffer from well identified artefacts. The reasons are that due to a higher number of parameters, complex molecules are in general tedious to parametrize for polarizable force fields, which often require the use of specialized software with a smaller user base. Additionally, their slower computational performance hampers their application to large biomolecular systems. Fortunately, significant advances have been recently made to overcome some of these limitations.
B. Implicit strategies

The current limitations in the explicit treatment of the electronic polarizability forced the simulation community to look for simpler, easy-to-use, and readily available methods to improve ion binding and pairing in widely used simulations. Several approaches have thus been developed to compensate for the artefacts arising from the lack of electronic polarization in such simulations within the existing interaction terms of non-polarizable force fields. We will briefly mention in the following the main strategies adopted in the literature, before focusing more closely on the electronic continuum correction approach.

One strategy to take into account induced polarization in a mean field approach has been to bypass the commonly used combination rules for Lennard-Jones parameters. Ad hoc pair-specific Lennard-Jones parameters (for example, the NBFIX corrections in the CHARMM force field\(^\text{32}\)) are defined for specific interacting pairs of ions or charged groups in biosystems\(^\text{33–36}\) in order to improve their pairing properties.\(^\text{7,36–38}\) There are several potential limitations to these approaches. The parametrization of each charge pair of interest has to be done independently, which can be tedious. Furthermore, experimental data required as a parametrization target are not always available and are often difficult to obtain. This is of course a problem for any parametrization effort, but it can often be partially overcome by relying on combination rules instead of parametrizing each interaction pair separately. The result is that the efforts relying on pair-specific interaction terms only focus on the interactions between a few atom types at a time. It is also unclear how the introduction of these specific parameters affects the balance of the interactions of other (unmodified) pairs and whether the corrections developed in different studies are mutually compatible and transferable to other force fields. Finally, in situations where the binding of a charged moiety is critical, the strategy of simply pushing away charges to decrease their interaction results in an inaccurate description of the interactions. Both polycoordination, \(e.g.,\) Ca\(^{2+}\) binding sites in proteins, or interactions where directionality is fundamental, \(e.g.,\) Zn\(^{2+}\) binding sites, are problematic.

Other authors suggested\(^\text{39}\) adding a \(1/r^4\) term to the standard 12-6 Lennard-Jones potential to account for charge-induced dipole interactions that are neglected in standard force fields.\(^\text{39}\) A force field with the resulting 12-6-4 Lennard-Jones-type non-bonded model has been developed for a series of monovalent\(^\text{40}\) and multivalent\(^\text{41}\) ions in combination with dif-
ferent water models. This model was shown to be able to correctly capture hydration free energies, ion–water oxygen distances, and coordination numbers simultaneously, and has given promising results on ion pairing properties. However, for divalent cations such as Mg\(^{2+}\) or Zn\(^{2+}\), further tuning of their pairwise interaction with specific RNA groups, \textit{e.g.}, its phosphate backbone or the guanosine and adenosine N7 atom, was found necessary to reproduce experimental binding free energies.\(^{42}\)

Instead of modifying the parameters of the van der Waals dispersion interaction, different approaches have suggested to directly tune the electrostatic properties of the solute through the modification of atomic charges. In this spirit, refinements\(^{43,44}\) of standard force fields have been suggested to reduce the strength of protein salt bridges that are strongly overestimated in most standard force fields.\(^{15,16}\) In particular, the ff15ipq force field,\(^{45}\) recently developed with the implicitly polarized charge method (IPolQ), uses non-polarizable atomic charges that were rederived to implicitly capture the energy of polarization by the presence of a solvent, namely water in biological systems. This force field, which keeps full integer charges on charged residues but changes the charge distribution compared to the original force field, has been shown to significantly improve the description of the strength of salt bridges in proteins.

In another approach, researchers have attempted to scale the total charge of ions or charged moieties to improve their pairing properties. Charge scaling has been for a long time prevalent in the context of ionic liquid simulations,\(^{24,46,47}\) where a scaling factor of 0.7–0.8 has been shown empirically to give reasonable thermodynamic properties.\(^{48}\) Charge scaling is not an alien concept in biological simulations either,\(^{49,50}\) though its adaptation has been so far scarce and mostly limited to issues involving ion channels\(^{51}\) or enzymatic catalysis.\(^{52}\) In recent years, the charge scaling strategy has been revived and physically grounded in a series of papers by Leontyev and Stuchebrukhov\(^{53–56}\) who rationalized this approach in terms of a mean-field theory called here the electronic continuum correction, which we will focus on from here on.

C. Electronic continuum correction (ECC)

A physically well-justified approach to decrease the binding between charged moieties and groups in non-polarizable force field simulations is to introduce electronic polarizability
within a mean-field approach via ECC. Note that the missing electronic polarizability in non-polarizable force fields leads to insufficient screening of interactions between charges. As the electronic polarizability is fairly constant in biological systems, one can incorporate it as a dielectric continuum, which can in principle be done in two ways—either by immersing the system in a continuum medium with the electronic polarizability equal to $\epsilon_{el} (ECC \epsilon)^{57}$ or by scaling charges of an uncorrected force field. Both approaches are theoretically equivalent. In practice, the uniform weakening of all Coulombic interaction by a background $\epsilon_{el}$ is impractical for biomolecular simulations, as such a global scaling of the charges may introduce undesired changes (overscaling) in carefully parametrized interactions. Features such as dihedral angular terms in molecules, sensitive to changes in 1–4 charge interactions, or the area per lipid in a membrane, sensitive on modifications in the overall lipid–lipid interactions, are often carefully fitted in standard force fields to reproduce experimental data, and should thus be left untouched. Alternatively, one may develop an entirely new force field, but this is an enormous task considering the complexity of biomolecules and their environments. For some simple systems, $ECC\epsilon$ has been applied with promising results.\textsuperscript{57} Moreover, $ECC\epsilon$ seems to resolve some issues related to charge scaling.\textsuperscript{58} The charge scaling ECC variant is less intrusive, as it relies on the fact that only heavily charged groups or atoms contribute significantly to charge–charge interactions. This means that partial charges on weakly polarized groups can be safely left untouched.

The possibility to effectively account for electronic polarization via charge scaling has been around since the 1980s. Pioneer work by Jönsson, Edholm, and Teleman stated explicitly: “One very simple way to remedy the lack of dielectric screening in the water potential is to scale ionic interactions with the high frequency dielectric constant…”.\textsuperscript{49} However, later in the same paper the authors scaled the ionic charges by a factor of 0.5, which from the present point of view seems like overscaling. This ambiguity may explain why systematic charge scaling has not caught up until about a decade ago. Only then the appropriate scaling factor for ionic charges—equal to the inverse square root of the high frequency dielectric constant of water (\textit{i.e.} 0.75)—was introduced.\textsuperscript{53} In the meantime, nevertheless, charge scaling has been applied as an empirical correction to several problems, including studies on the structural and dynamic properties of ionic liquids,\textsuperscript{59} charge transfer from catalytic ions in enzymes,\textsuperscript{60} and ion channels.\textsuperscript{51}

The seminal contribution of Leontyev and Stuchebrukhov lies in developing charge scal-
ing on a firm physical basis. By a simple manipulation of Coulomb’s law, one can show that scaling the charges by the inverse square root of the high frequency dielectric constant is mathematically equivalent to immersing the charges into a medium with the respective relative permittivity. Indeed, following the derivation of classical electrostatics, one can show that such charge scaling is rigorous and represents a way to preserve the symmetry between the “source” and “test” charges in a dielectric medium. Moreover, this mean field approach is physically well justified for systems which are roughly homogeneous in their electronic responses, i.e., which do not possess spatial discontinuities in their high frequency dielectric constants. Fortunately, most of biologically relevant molecular environments composed of moieties such as water, proteins, lipids, and nucleic acids are quasi-homogeneous in this respect with a high frequency dielectric constant of about 2 within the relevant temperature range (see Table I).

In principle, the electronic continuum correction (ECC) accounts for electronic polarization in a mean field way as follows: (i) Scale any charged group so that its total charge is 0.75 times its nominal charge. This takes care of electronic polarization by effectively putting the charged moiety in an environment with a high frequency dielectric constant of water $\epsilon_{el} \sim 1.78$ (corresponding to a refraction index of $n = \sqrt{\epsilon_{el}} \sim 1.34$). The charge scaling factor is then $\frac{1}{\sqrt{n}} = 0.75$, which is to a good approximation valid also for any aqueous environment or complex biological system. (ii) Run a standard classical molecular dynamics which recovers naturally the nuclear polarization. At least, this is how it should be done in the ideal world where the water model itself is fully compatible with the ECC approach, i.e., its dielectric constant only contains the nuclear polarization with $\epsilon_{nuc} = 45$. In reality, the commonly used non-polarizable water models exhibit dielectric constants between 50 and 95. This means that within their parameterization, the ECC concept has been implicitly taken into account only to some degree. Therefore, it is legitimate to consider scaling factors for ions lying between 0.75 and 1 depending on the employed water model, so that one avoids overscaling due to partial double-counting for the electronic polarization effects both within the ion and water models. Similar problems arise when considering charged groups of amino acids, lipids, nucleic acid, and other readily parametrized components of biological systems. Depending on the particular force field parameterization, electronic polarization effects are already implicitly included to varying degrees and one should again avoid overscaling. In our case, the strategy so far has been to make only minimal modifica-
tions to the existing biomolecular force fields, focusing the scaling of the partial charges to the charged (or zwitterionic) groups, with a factor of 0.75 or slightly higher.$^{4-6,10,67}$

Some quantities, such as the solvation free energy often used in force field parameterization or validation, are not so straightforwardly obtained with scaled charge ECC force fields, and care must be taken in their evaluation. Indeed, as pointed out and discussed early on by Stuchebrukhov and coworkers,$^{53}$ the solvation free energy $\Delta G_{\text{solv}}$ in the ECC framework should be calculated in two steps—the usual nuclear part usually obtained with alchemical free energy simulations, $\Delta G_{\text{MD}}$, and the electronic solvation energy part, $\Delta G_{\text{el}}$, which can be obtained by solving the Poisson equation or, for simple ions, estimated with a Born solvation model. The total solvation free energy is finally obtained as the sum of these two contributions $\Delta G_{\text{solv}} = \Delta G_{\text{MD}} + \Delta G_{\text{el}}$ which are of comparable magnitude.

Strictly speaking, the applicability of the ECC approach is limited to systems with an essentially uniform high-frequency dielectric constant across the system. Hence, simulations of a solvent–air or solvent–metal interface are problematic.$^{68}$ In such cases, on the solvent side, the ECC scaling factor should be applied, whereas on the other side a different scaling factor should be applied, e.g., a scaling factor of 1 (full charges) for air. Interestingly though, the application of ECC in molecular dynamics simulations of solid–liquid interfaces disregarding the limitation seems to improve the interaction of charges with highly polar surfaces.$^{69-73}$

### III. MONOATOMIC IONS / SIMPLE SALT SOLUTIONS

The ECC framework has been first applied to simulations of simple salt solutions made of monoatomic ions. Following the original prescription of the ECC theory, the total ionic charge is fixed at 0.75 times its nominal value (i.e., $\pm 0.75$ for monovalent ions and $\pm 1.5$ for divalent ions), so that only the two parameters of the 12–6 Lennard-Jones potential remain to be determined for each ion, i.e., the depth of the potential energy well, $\epsilon$, and the distance at which the potential is zero, $\sigma$. These parameters need to be slightly adjusted compared to full charge force fields because the charge reduction affects, among others, the ion–water interaction and thus the hydration structure. As a general rule, a 5–10% decrease in the ion radius allows to recover the correct hydration structure, however, the exact value is water model dependent. Because of the charge neutrality of salt solutions, it is impossible to obtain
experimental data for a single cation or anion. To overcome this difficulty, one strategy is to start with simple salts that are known to exhibit very little pairing in solution (at least at not too high concentrations), and to use local experimental probes to parametrize the ions. In this spirit, force fields for simple monovalent cations and anions such as Na\(^+\), Li\(^+\), and Cl\(^-\) were developed using data on the local ion hydration structure obtained from neutron scattering with isotopic substitution. This strategy improved the ion pairing behavior in such ionic solutions and was also shown to successfully capture the affinity of halide anions to a water–oil interface.

While in principle the charge scaling factor is fixed in the ECC theory, several groups considered it as an additional fitting parameter, which may reflect the limitations of water force fields. This resulted in suggested scaling factors lying in the 0.85–0.94 interval, depending on the employed water force field and experimental observables targeted within the fitting procedure. As a result, several scaled charge force fields for NaCl in solution are now available.

Scaled-charge force fields were shown to provide a faithful description of simple electrolytes over a broad range of concentrations. Moreover, compared to standard non-polarizable force fields, they significantly improved description of a number of structural and dynamic properties. For instance, the Madrid-2019 scaled charge force field for a series of simple monovalent ions was shown to properly capture a number of thermodynamic (activity coefficients, solubility) as well as dynamic (viscosity, diffusion coefficients) properties. It also improved the description of the salting out effect in NaCl solutions. In general, scaled charge force fields have significantly improved the description of ion pairing, as well as diffusion, viscosity of electrolyte solutions, and water reorientation dynamics in the ionic hydration shell.

In the light of these promising results, the ECC scaled charge strategy was then extended towards the more challenging cases of non-transition metal divalent cations that exhibit very strong binding artefacts in simulations. The first cation under study was Ca\(^{2+}\), which is highly relevant for many biological processes. The ECC calcium force field was parameterized against experimental neutron scattering data on concentrated calcium chloride solutions which are very sensitive to calcium–water distances. The ECC force field was shown to properly describe the structure, viscosity, and diffusion of calcium chloride solutions. One should note here that the use of scaled charge force field does not always reduce the amount of
ion pairing, which is determined by the balance between ion–ion and ion–water interactions. In the case of CaCl$_2$, the use of ECC force fields indeed leads to a slightly reduced amount of ion pairs in solution. Our ECC calcium force field was later further refined to yield better agreement with both experiments—Extended X-Ray Absorption Fine Structure (EXAFS) and neutron scattering—and benchmark computations ($ab$ initio molecular dynamics).$^{84}$ This approach has also led to a scaled charge Mg$^{2+}$ force field,$^{85}$ which was thoroughly checked against other Mg$^{2+}$ models recently.$^{86}$ A similar strategy was also attempted for Zn$^{2+}$.$^{85}$ Even though Zn$^{2+}$ is not formally considered to be a transition metal, it does form partially covalent interactions with solvent molecules, counterions, and other ligands, which are beyond what ECC (or even explicitly polarizable polarizable force fields) are able to describe. Attempts were made to treat such cases using specific interaction-sites potentials.$^{87}$

Scaled charge force fields were also recently proposed$^{88}$ for a wide range of polyvalent metal cations, including high valence transition metals such as Y$^{3+}$, Zr$^{4+}$, and Hf$^{4+}$, using the hydration free-energy and ion–water oxygen distance as target properties. Even if such quantities can be reasonably well reproduced, one should be cautious when modeling highly charged ions with classical force fields, as their specific chemical interactions may not be adequately captured.

For full charge models of ions, Li and Merz demonstrated that Lennard-Jones parameters optimized for the ions within a standard 12-6 interaction potential were able to reproduce either the solvation free energies or the ion–water oxygen distances.$^{39}$ With the extension to 12-6-4 potential form these two measurables could be reproduced simultaneously, with an open question concerning the description of ion pairing. In this context, it is worth asking whether ECC, which was developed to reproduce structural properties of electrolyte solutions, also describes ion–water energetics accurately. To this end, we evaluated (Fig. 1) for some of our cation ECC force fields the solvation free energy $\Delta G_{\text{solv}}$, following the ECC prescriptions (see Section II C). The two contributions to the solvation free energy, $\Delta G_{\text{MD}}$ and $\Delta G_{\text{el}}$, are comparable in magnitude. The total $\Delta G_{\text{solv}}$ was found to be in excellent agreement with experimental data, with similar accuracy as standard force fields, even though it never entered in the parameterization procedure. While such calculations represent a useful validation of the force field, one should keep in mind that the value of $\Delta G_{\text{el}}$, which accounts for half the total free energy, is highly sensitive to the details of the cavity used for its estimation and thus has a large uncertainty. Hence, the total solvation
free energy could hardly be employed as a target property for parametrization.

FIG. 1. Example solvation free energies of ions into SPC/E water. The full charge ion models are from Ref. 89 (Ca$^{2+}$ & K$^+$), Ref. 90 (Na$^+$), and Ref. 91 (Mg$^{2+}$). For these models, $\Delta G_{\text{MD}}$ agrees well with the experimental values. For the ECC ions (Ca$^{2+}$ (the Ca$_{2s}$ version) is from Refs. 83 & 84, K$^+$ is from Ref. 93, and Mg$^{2+}$ (the “small” version) from Ref. 85), the electronic solvation energy $\Delta G_{\text{el}}$ is estimated with the Born solvation model with Born radii, and the sum of $\Delta G_{\text{MD}}$ and $\Delta G_{\text{el}}$ is in great agreement with experiment. The $\Delta G_{\text{MD}}$ values contain the correction for the neutralization of the charge during the decoupling of electrostatic interactions. These values are somewhat water model dependent, with the values obtained with four commonly used models (SPC/E, TIP3P, TIP4P/2005, and OPC) differing by up to 7%.

The description of salt solutions at the air–water interface is problematic within the ECC framework due to the discontinuity in the high frequency dielectric constant, as discussed above. As a result, the experimental dependence of the surface tension on the electrolyte concentration is not properly captured using ECC ions, see Fig. 2. Here, the surface tensions of both tested water models, SPC/E and 4-point OPC show too weak dependence on the solution molality. The vertical shift between the models stems from the ability of OPC to better describe the surface tension of the pure water–air interface. This limitation of ECC in describing interfacial behavior has to be taken into account when modeling phenomena involving the air–water interface. It is fortunate that such interfaces are rare in biological systems.
FIG. 2. Surface tension of NaCl and CaCl\textsubscript{2} solutions calculated from molecular dynamics simulations using the ECC ions and two water models; SPC/E that was used in the original ECC ion papers\textsuperscript{74,83} as well as the 4-point OPC model which provides the best description out of commonly used and rigid water models of surface tension of pure water.\textsuperscript{96} Data are also shown for full charge ion models by Dang and coworkers (“Full”)\textsuperscript{89,90} with the SPC/E water model. The surface tension values were calculated from the asymmetry of the pressure tensor, with the error estimates being smaller than the symbols employed in the graphs. ECC fails quantitatively to capture the slopes of the experimental curves regardless of the water model employed. Full charge models, in contrast, tend to overestimate the effect of ions on surface tension. Experimental data are from Refs. 97 (CaCl\textsubscript{2}) and 98 (NaCl).

IV. POLYATOMIC IONS

Since ECC prescribes the total charge of the ions in simulations with implicit electronic parametrization, only the two parameters of the Lennard-Jones potential remain to be optimized for monoatomic ions. The situation is significantly more complex for polyatomic ions, as there are many more parameters to determine. To a first approximation, all bonded terms (bonds, angles, and dihedrals) can be adapted without change from regular force fields. In some cases, however, the dihedral terms may have to be adjusted as scaling the charges affects the interactions of atoms separated by 3 bonds (i.e., the 1–4 interactions). Only the total ion charge is prescribed by ECC, which leaves the partial charges and Lennard-Jones parameters of the individual atoms to be determined.

Many common force fields, including those of the Amber family, use partial charges
which are determined by fitting the electrostatic potential obtained by quantum chemical calculations under certain restraints. This provides a charge distribution that reproduces the electrostatics potential around the molecule of interest. The idea behind ECC (see above and the recent review\textsuperscript{62}) is that the electronic polarization effectively reduces this electrostatic potential. Hence, in a most straightforward way ECC partial charges may be obtained by simply scaling the charges fitted to the electrostatic potential by a factor of $1/\sqrt{\epsilon_{\text{el}}} \simeq 0.75$.

The Lennard-Jones parameters still remain to be determined. For monoatomic ions, this has been done such as to reproduce the location of the first peak of the ion–water oxygen radial distribution function, obtained from neutron scattering or EXAFS. For complex ions, such experimental radial distribution functions are rarely known. While standard force fields often use the solvation free energy as a target property, this is not practical for scaled charge models, as discussed above. Alternative targets can be used, such as osmotic coefficients or radial distribution functions calculated from \textit{ab initio} simulations, but there is \textit{a priori} no unique recipe. In a first approximation and in view of the scarcity of the available data, many scaled charge force fields for complex ions simply did not modify the Lennard-Jones parameters.

Adopting this strategy for carbonate, nitrate\textsuperscript{93} and sulfate\textsuperscript{99} allowed for capturing successfully the structure of concentrated KNO\textsubscript{3}, K\textsubscript{2}CO\textsubscript{3} and Na\textsubscript{2}SO\textsubscript{4} solutions, in which ions exhibit excessive pairing and aggregation when described with non-polarizable force fields. A similar strategy was more recently adopted also for the acetate anion.\textsuperscript{84} In this case, it turned out that already after charge scaling, the radial distribution functions between the acetate oxygen atoms and water were in good agreement with those obtained from \textit{ab initio} molecular dynamics simulations. Thus, further scaling of the Lennard-Jones parameter $\sigma$ was not required. In addition, this scaled charge acetate combined with the scaled charge calcium developed previously, were shown to describe well the structure of concentrated calcium acetate solutions as validated against neutron scattering measurements.\textsuperscript{84} Force fields with either full charges (or with only the cation charges scaled) strongly overestimate ion pairing and aggregation in the solution.\textsuperscript{84} However, it is worth noting that a careful comparison with various experimental data suggests that, while full charge force fields indeed strongly overestimate the binding, the ECC force field with a scaling factor of 0.75 appears to underestimate the binding strength.\textsuperscript{100} A force field based on a scaling factor of 0.8 was
found to be in a better agreement with experimental data, in line with previous suggestions discussed earlier ref part of the discussion.

FIG. 3. Chemical structure of the guanidinium–acetate ion pair together with the arginine–aspartate salt bridge. The guanidinium nitrogen atoms and acetate hydrogen atoms substituted in the neutron scattering experiment\textsuperscript{101} are colored in blue and green, respectively.

The same ECC acetate was used together with a scaled imidazolium force field to investigate ion pairing in imidazolium–acetate solutions.\textsuperscript{101} These two moieties represent a simple model of interacting groups involved in salt bridges in proteins—acetate is a proxy for carboxylate groups in aspartate or glutamate residues, and the imidazolium cation is a proxy for the arginine side chain (Figure 3). While the original full charge force fields overestimated ion pairing, charge scaling led to a structure of the solution being in a very good agreement with results of neutron diffraction experiments with double isotopic substitution, which provide a sensitive measure of the degree of ion pairing in the solution. The reliability of the ECC force field being thus validated, additional information about the geometry of the ion pair was obtained, such as the existence of an out-of-plane binding mode involving the hydrophobic group of acetate in addition to the standard in-plane geometry. Analysis of the Protein Data Bank revealed that such an out-of-plane interaction motif, although less common than the in-plane one, is found in 25–55% of protein structures. This suggests that despite the apparent simplicity of such model salt solutions, valuable insights can be gained into the strength and geometrical aspects of biologically relevant interactions thanks to the use of appropriately scaled force fields. Furthermore, this study confirmed that the strength
of salt bridges is strongly overestimated in standard non-polarizable force fields.\textsuperscript{15,16}

Other polyatomic biologically relevant ions such as multivalent phosphates\textsuperscript{102} or hydrogenoxalate and oxalate\textsuperscript{103} anions were also recently successfully improved using the ECC framework.

V. BIOMOLECULES

Many biomolecules such as lipids, proteins, or DNA/RNA are either charged or contain charged groups. The ECC approach for monoatomic and polyatomic aqueous solutions can be extended to more complex solutions and biomaterials\textsuperscript{17} as the variation in $\epsilon_{el}$ is small, see Table I. As a result, the charge scaling factor, although strictly speaking environment-dependent, can be well approximated as being constant in these biomolecular systems. This approximation allows the usage of a single scaling factor in the development of force fields where molecular interactions generated for diverse molecules are expected to be mutually compatible.

<table>
<thead>
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<th>Material</th>
<th>T(K)</th>
<th>$n$</th>
<th>$\epsilon_{el}$</th>
<th>$s$</th>
<th>Ref</th>
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<tr>
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<td>0.67</td>
<td>104</td>
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<td>0.68</td>
<td>104</td>
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<td>1.4394</td>
<td>2.07</td>
<td>0.69</td>
<td>104</td>
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<tr>
<td>Hexadecane</td>
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<td>1.430</td>
<td>2.04</td>
<td>0.70</td>
<td>104</td>
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<tr>
<td>Cornea (human)</td>
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<td>1.89-1.96</td>
<td>0.71-0.73</td>
<td>105</td>
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<tr>
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<td>1.369</td>
<td>1.87</td>
<td>0.73</td>
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<tr>
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<td>0.73</td>
<td>104</td>
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<td>1.360</td>
<td>1.85</td>
<td>0.74</td>
<td>107</td>
</tr>
<tr>
<td>20% Glycerol in water</td>
<td>293</td>
<td>1.3572</td>
<td>1.84</td>
<td>0.74</td>
<td>104</td>
</tr>
<tr>
<td>water</td>
<td>303</td>
<td>1.333</td>
<td>1.77</td>
<td>0.75</td>
<td>104</td>
</tr>
<tr>
<td>colorectal mucosa (human)</td>
<td>310</td>
<td>1.315</td>
<td>1.73</td>
<td>0.76</td>
<td>63</td>
</tr>
</tbody>
</table>
A. Lipids

All cells, organelles, and vesicular bodies are encapsulated by membranes. Their basic constituents are amphipathic lipid molecules. Independently of their net charge, most lipid moieties and their derivatives contain positively and negatively charged groups as well as some highly polar groups. These charged groups include for example (methylated) ammonium, choline, carboxylate, and phosphate groups. As with simple molecules containing these motifs, lipids display excessive binding when interacting with other charged groups when described with standard non-polarizable force fields. In fact, the interaction of lipids with charged moieties is a conundrum of great relevance.\(^\text{108}\)

Until recently, the consensus was that cations, even monovalent ones such as Na\(^+\), interact strongly with the membranes. This idea originated from molecular dynamics simulations with standard non-polarizable force fields, within which most biologically relevant cations accumulated at and often overcharged even membranes made of only zwitterionic lipids with zero net charge.\(^\text{109}\) Yet, the conformation of the lipid headgroup region, as characterized by NMR-derived order parameters, showed no major changes during titration experiments with monovalent cations.\(^\text{110}\) A more recent study employing electrophoresis and isothermal titration calorimetry concluded that these simulations exaggerated Na\(^+\) adsorption onto bilayers composed of zwitterionic phosphatidylcholine lipids.\(^\text{111}\) For divalent ions, the situation is even more dramatic—in standard simulations, Ca\(^{2+}\) ions bind irreversibly to membranes consisting of phosphatidylcholine lipids and their strong accumulation at the membrane surface leads to overcharging, which contradicts with most of the available experimental data.\(^\text{9}\) This excessive binding of cations to membranes in all standard non-polarizable force field simulations has been thoroughly addressed recently.\(^\text{32,112}\)

To address the overbinding problem, the developers of the CHARMM36 force field introduced a specific pair-wise non-bonded repulsive potential between certain charged groups to prevent their close approach.\(^\text{32}\) While this potential results in somewhat extended distances between these groups, it leads to a better agreement with several other experimental measurables. It effectively reduces the interaction and helps to address the overbinding issue. However, this approach, which lacks a clear physical justification, has its drawbacks. Processes that involve polycoordination of ions with several lipids are not described properly. This is especially important for divalent cations such Ca\(^{2+}\) that can mediate binding...
between like charge regions of signaling proteins and membrane lipids. An example of such a case is the membrane-binding C2 domain of protein kinase C α (PKCα-C2), which is bridged by Ca\(^{2+}\) to a negatively charged phosphatidylserine lipid (see Fig. 4). Modeling such situations is beyond the capabilities of the original CHARMM36 force field and its NBFIX modification. As shown in the upper left panel of Fig. 4, a membrane consisting of phosphatidylcholine and phosphatidylserine lipids gets overcharged by Ca\(^{2+}\), and PKCα-C2 does not attach to the membrane when the CHARMM36 force field is used without NBFIX. When NBFIX terms are included, PKCα-C2 binds to the membrane. However, the binding is not bridged by Ca\(^{2+}\) and the protein orientation does not agree with one in the crystal structure.

The source of the excessive interaction between ions and lipids in classical force fields is again the lack of electronic polarizability and the associated shielding. Therefore, the use of polarizable force fields should in principle solve the problem. While there is a significant progress in the accuracy of polarizable lipid force fields,\(^{21,114–117}\) their results are not always in better agreement with experiment than those of classical force fields.\(^{21}\) Combined with the limited availability of parameters for lipid moieties and a tedious parametrization process, the larger requirements of computational resources renders polarizable simulations of limited use in studies of complex biological processes. It is, therefore, tempting to instead correct the classical force fields within the ECC approach.

The very first, to the best of our knowledge, atomistic simulation of a lipid bilayer\(^{50}\) used charges reduced by 50% based on a pioneering work on surfactants by Jönsson, Edholm, and Teleman.\(^{49}\) However, the scaling factor of 0.5 lacks a firm physical basis. The first attempts to solve the overbinding problem within the ECC framework used scaled ions\(^{74,83}\) together with unmodified traditional lipid force fields.\(^{8,9}\) From these studies three main conclusions can be extracted. First, scaling only the ions is not sufficient to avoid the overbinding of cations to a phosphatidylcholine (PC) membrane.\(^{9}\) Second, the change in binding brought about by ECC ions depends on the lipid force field. As an example, ion scaling leads to a decrease in Na\(^{+}\) binding to POPC membranes with both Slipids\(^{118}\) and CHARMM36 force fields.\(^{119}\) However, for Ca\(^{2+}\), the expected decrease in binding with ECC ions is only observed with CHARMM36, where the binding is no longer irreversible at the simulation timescale. In contrast, the binding is still irreversible and (counterintuitively) enhanced when the Slipids force field\(^{120}\) is used. Third, anions such as Cl\(^{-}\) do not bind to POPC membranes.
FIG. 4. Top panel: characteristic snapshots of PKα-C2 (orange) after 500 ns of simulation with a membrane containing palmitoyloleoylphosphatidylcholine (green) and palmitoyloleoylphosphatidylserine (yellow) with CaCl$_2$ (cyan). Three force fields were used for the simulations: the standard (full charge) CHARMM36 with and without the NBFIX correction, and its scaled variant based on ECC. Only the ECC force field is able to capture the correct binding mode present in the crystal structure$^{113}$ (PDB:1DSY). This is further highlighted in the bottom left panel, which shows aligned structures of the PKα-C2 together with Ca$^{2+}$ and PS during 400 ns of simulation after the binding is established during the first 100 ns. The bottom right panel shows the histogram of the shortest distance between the phosphatidylserine lipids and the center of mass of the loops that bind Ca$^{2+}$. Only the simulation with scaled charges agrees reasonably well with the crystal structure (dashed line). The last 400 ns of simulation data are again used for the analysis.

Solving the overbinding problem within the ECC framework requires simultaneous scaling of ions and lipids. This includes the ubiquitous phosphatidylcholine lipids that contain zwitterionic headgroups but have a zero net charge. The ECC framework does not provide a recipe on how the scaling of such molecules should be done. One can scale only the charged and polar groups within the headgroup region, i.e., amine, choline, carboxyl, phosphate, and carbonyl groups, leaving any other apolar groups of the lipid untouched. Alternatively, the charges of the whole head group and the glycerol region could be scaled. Both approaches will ultimately decrease the charge separation and local dipoles within a lipid. The best scaling strategy depends on the distribution of partial charges in the original lipid force...
For lipid models of the Amber force field family, the net charges of charged groups are distributed all over the headgroup and glycerol region. This leads to a significantly smaller charge separation in the headgroup compared to CHARMM-based force fields. To some extent, such charge delocalization resembles the ECC scaling process and may explain why applying the ECC framework is not straightforward for the Amber force field family. Notably, one needs to avoid overscaling of the already (implicitly) scaled charges.

A uniform scaling of the entire headgroup and glycerol regions with a factor of 0.80 was used for the Lipid14 palmitoyloleoylphosphatidylcholine (POPC) model, which led to a dramatic improvement of the binding behavior of monovalent Na\(^+\) and divalent Ca\(^{2+}\) cations. This ECC-POPC model achieves a nearly quantitative agreement with NMR experiments. To achieve such a level of accuracy, the scaled atoms had their Lennard-Jones \(\sigma\) parameters scaled by a factor of 0.89. The same scaling strategy was applied to palmitoyloleoylphosphatidylserine (POPS) using the Amber Lipid17 force field as a template. Contrary to POPC, POPS is a charged lipid and thus the total charge of the ECC-POPS model was scaled to \(-0.75\). Again, the ECC scaling rendered the interactions between POPS and the cations more realistic than in the original force field.

The strategy of scaling the headgroup and the glycerol as a whole was also applied to the CHARMM36 POPC force field resulting in an improvement of the binding of monovalent and divalent ions. Note, however, that lipid models using the variants of the CHARMM force fields are modular, meaning that a molecule is composed of fragments usually involving functional groups with a net integer charge. In this case, a scaling strategy where only charged fragments are scaled seems more appropriate. Such an approach avoids undesired structural changes and minimizes the changes in the interactions with other membrane moieties with respect to the original model. Within this strategy, the nominal charge of each fragment after scaling is the original charge of the group times the ECC scaling factor. For example, a entire choline group with a total charge of \(+1\) e has a total charge of \(+0.75\) e after scaling. Yet, the partial charges of independent atoms do not need to follow this relation. In addition, some strong dipoles in very polar functional groups may need to be scaled, such as the carbonyls resulting from the esterification of the glycerol group and the acyl chains. This is usually done to keep the partial charges on these carbonyl oxygens below the values of their more polar counterparts such as phosphates and carboxylic groups after
their scaling.\textsuperscript{10}

Overall, adapting existing lipid force fields to the ECC framework dramatically improves the description of lipid–ion interaction.\textsuperscript{9,10,67} This is achieved without compromising the excellent performance of the original force fields concerning structural properties of the lipid bilayers. By limiting the ECC scaling only on charged (and possibly also highly polar) groups, accurate binding of lipids to charged moieties is achieved without the need to reparametrize the entire lipid model. The remaining question to address in the future is whether charge scaling also improves the binding of other charged biomolecules, such as proteins, to membranes.

B. Proteins

Five out of the 20 essential amino acids—Glu, Asp, Lys, Arg, and His (when doubly protonated)—have charged side chains, and play a critical role in protein structure and function. Charged amino acids drive protein folding by promoting their exposure to the solvent. For example, protein stability and structure is dependent on pH, which changes the protonation state of these amino acids.\textsuperscript{121} Salt bridges, where a positive and a negative amino acid directly interact, constitute fundamental intra- and inter-protein interactions.\textsuperscript{122} They drive multiple phenomena, such as multimerization and the formation of complexes required to form functional assemblies. Charged amino acids are also fundamental in the binding of ligands and cofactors where they foster the recognition, selectivity, and strength of the binding.\textsuperscript{123} Furthermore, both monoatomic and polyatomic ions act themselves as protein ligands and cofactors.\textsuperscript{124}

Considering that the high frequency dielectric constant ($\epsilon_{el}$) of the protein environment is comparable to that of water (see Table I), it is not surprising that charge–charge interactions involving proteins are also overestimated when using standard non-polarizable molecular dynamics simulations. This issue was noticed early by people simulating ion channels, where the ions, especially divalent cations, usually get trapped in the protein’s selectivity filter.\textsuperscript{125} For this reason, charge scaling of the ions and even the charged groups in the protein’s selectivity filter is one of the strategies proposed to overcome such excessive interaction and to allow for realistic movement of ions through the channel.\textsuperscript{51} The potential of applying charge scaling to amino acids to improve overbinding problems was soon realized by Stuchebrukhov
et al.\cite{61,126} when developing the ECC framework. In particular, the description of the important salt bridges in proteins improved dramatically when using the ECC correction.\cite{61,101} ECC-scaled proteins were then adopted and proved extremely useful in studies on systems where the proper description of ion–protein interaction is central.\cite{4–6,127}

In particular, Ca\textsuperscript{2+} plays a key role in cellular signaling, one of its most important receptors being calmodulin. Calmodulin has four Ca\textsuperscript{2+} binding sites in its EF hands, each with a different but reversible binding affinity. Standard force fields lead to grossly overestimated binding energies of up to \(\sim -19\) kcal/mol whereas experiments predict \(\sim -5.4\) kcal/mol.\cite{4} Using an ECC Ca\textsuperscript{2+} ion model together with the original Amber protein force field decreased the largest binding free energy to \(-10\) kcal/mol, which is still twice the experimental value. Only when the negatively charged side chains were scaled, too, did the binding free energies drop close to the experimental values. Quantitatively, scaling has varying effects on each of the binding site. The binding behaviour of Ca\textsuperscript{2+} to calmodulin in an aqueous solution with 50 mM CaCl\textsubscript{2} with and without ECC scaling is illustrated in Fig. 5. It is of importance to highlight the overbinding of Ca\textsuperscript{2+} to calmodulin when the standard CHARMM36 force field without NBFIX is used. We observed that \(\sim 20\) Ca\textsuperscript{2+} were bound to calmodulin instead of the expected 4 (which corresponds to one per binding site). We saw that certain calmodulin binding loops embraced two Ca\textsuperscript{2+} ions. Furthermore, even a single carboxylate group occasionally bound two Ca\textsuperscript{2+}. This is clearly an artifact, as carboxylate groups are abundant in water-soluble proteins and not all of them bind Ca\textsuperscript{2+}. When using the same CHARMM36 force field with ECC parameters, the Ca\textsuperscript{2+} binding is reduced drastically to an average of 6 Ca\textsuperscript{2+} bound to calmodulin. This improvement was not without collateral problems, as the binding affinity to some loops became too low so that they unloaded during the simulations (see black loop on the left snapshot of Fig. 5). Despite this drawback, ECC rendered the binding profile more realistic, as the binding to carboxylate groups was reduced and a complete Ca\textsuperscript{2+}-binding loop was thus required for the interaction to take place.

For another Ca\textsuperscript{2+} binding protein, recoverin, ECC scaling not only improved the binding of Ca\textsuperscript{2+}, but also captured the overall protein conformational change upon ion binding. This conformational change is of biological interest as it is required for the activation of the myristoyl switch.\cite{127}

Intrinsically disordered proteins, especially highly charged ones, constitute another group of proteins for which ECC scaling provides a promising modeling approach. Most of the
FIG. 5. On top, characteristic snapshots of CaM in solution with 50 mM of CaCl$_2$. On the left side, the scaled CHARMM36 force field is used to account for ECC while on the right side, standard CHARMM36 force field is used. On both snapshots, the four calmodulin loops are highlighted with different colors (red, black, blue and green). On the lower part, the distribution of the contacts between Ca$^{2+}$ and COO$^-$ is reported for either the ECC force field (green) or the standard full charge force field (red).

commonly used protein force fields predict too stiff protein structures with little room for conformational changes. This results from the traditional need to preserve the crystallographic secondary and tertiary structures of proteins. While many factors may contribute to such stiffness, the interaction between neighbouring charged groups is an important factor in some common motifs. This interaction is in turn strongly dependent on the ionic envi-
While the CHARMM27 and Amber ff99SB-ILDN force fields resulted in collapsed structures, the CHARMM36 force field yielded a more extended configuration. OPLS force field provided an intermediate situation. The use of NBFIX always promoted more extended peptide structures by reducing the ions bound to the peptide. When applying ECC corrections instead, it was evident that the results—more extended or collapsed peptides—depended non-trivially on the original force field. Again, as with the lipids, the outcome of applying ECC scaling to the CHARMM36 force field is fairly predictable, resembling that of NBFIX. It reduces the number of counterion–peptide contacts. As bound ions shield the intra-peptide repulsion, ECC promotes more extended peptide structures. The overall repulsion between neighboring negatively charged groups is increased, although ECC weakens this interaction too. With the Amber force field, which implicitly accounts for electronic polarization to some extent with its delocalized partial charges, the results are system-dependent. In some cases, the binding of counterions to the peptide is even enhanced, resulting in more compact conformations. Therefore, ECC scaling may alter the balance between intra-peptide and peptide–ion interactions in largely unpredictable ways.

As mentioned in the previous section, in some cases ions can act as bridges for binding of proteins to lipids. Such a case is a demanding benchmark for the ECC approach, since it requires the correct balance between lipid–ion, lipid–protein, and ion–protein interactions. As shown in the upper right panel in Fig. 4, our ECC simulations of PKCo-C2 capture its binding to a membrane consisting of POPC and POPS lipids in the presence of CaCl$_2$. As shown in the bottom left panel of Fig. 4, the binding mode agrees well with the crystal structure, with Ca$^{2+}$ bridging the PKCo-C2 to a POPS lipid. The bottom right panel of Fig. 4 shows the histogram of minimum distances between the center of mass of the Ca$^{2+}$-binding loops of PKCo-C2 and membrane POPS. The ECC simulations are in much better agreement with experiment (gray dashed line) than full charge simulations using CHARMM36 with or without NBFIX. Still, the agreement is not perfect, as the orientation of a POPS lipid differs somewhat from the crystal structure, and for the majority of the simulation, only one Ca$^{2+}$ is involved in the bridging instead of the experimental value of two (PDB:1DSY). Moreover, the aforementioned PS–loops distance is slightly larger than in the crystal structure. These differences may have arisen from limited sampling in the
simulations or from the differences between the binding mode in the crystal structure and in the native environment. Finally, more work needs to be done to deal with other charged groups found in proteins such as those originating from phosphorylation.

C. Other biomolecules

Description of other highly charged biomolecules should also benefit from ECC scaling. Prime examples are sugars such as glycosaminoglycans and sialic acids in N-glycans, as well as DNA and RNA. Applying the ECC strategy to such biopolymers shares the problems discussed for polyatomic systems earlier. So far, there is no unique procedure to derive the scaled charges, which ultimately depend on how the unscaled partial charges were originally derived and distributed within the molecule. Nevertheless, promises of the ECC approach to describe phosphate-containing species have been demonstrated in a recent work on solutions containing divalent and trivalent phosphate, osmotic properties of which were greatly improved by the use of scaled charges.\textsuperscript{102} The extension of this approach to monovalent phosphates should be straightforward, but the challenge lies in its application to RNA and DNA. Indeed, contrary to proteins—where the charges are relatively far from the backbone that defines the secondary structure—in DNA, RNA, as well as in sugars, the charges are located directly on the backbone or sit next to the rings. Therefore, any change in partial charges is expected to have a large effect on the dihedral terms. These dihedral terms have been carefully optimised in full charge force fields to reproduce the complex conformational phase space of these biomolecules. Hence, extension of ECC to these biomolecules without compromising the description of their structural parameters remains challenging.

VI. CURRENT LIMITATIONS OF THE ECC FRAMEWORK

Despite the apparent simplicity of the ECC theory, which prescribes that partial charges of charged interacting moieties have to be adequately scaled, its practical implementation is not always that straightforward. First, an increasing number of studies suggests that the theoretical ECC scaling factor of 0.75 often results in too weak charge–charge interactions. Several groups have proposed scaling factors closer to one to resolve this issue,\textsuperscript{66} which can in part be justified by the use of non-optimal water models within the ECC framework.\textsuperscript{62}
Secondly, scaling of charges affects the interactions of the charged group with other moieties such as water, which were often fitted to reproduce experimental data (for instance ion–water oxygen radial distribution functions). Hence, even for simple salt solutions, charge scaling often needs to be accompanied with refining the Lennard-Jones parameters to restore full agreement with experiment in these properties. More generally, concrete implementation of the charge scaling depends on how the starting underlying full charges were originally developed for polyatomic groups. Note that there is not a unique set of charges compatible with the ECC framework, in the same way as different full charge force fields adopt different distributions of partial charges. From our experience, the charge distributions in several of the majorly used biomolecular force fields already implicitly include the effect electronic polarizability to a certain extent. In such cases, one should thus be careful not to overscale the charges. The water model and especially its dielectric constant also has a non-negligible effect on the ECC parametrization, with the same being also true for full charge models, as illustrated by the fact that many recent ion force fields are available in particular flavors adapted to each water model. In practice, simply scaling the charges of an existing biomolecular force field is often not sufficient to match experimental data for ion binding.

In addition, one should note that the outcome of charge scaling within the ECC framework, contrary to some other schemes developed to quench specific charge–charge interactions, is not always the weakening of ion pairing. In particular, because ion pairing is a property that depends on the balance between ion–ion and ion–water interactions, scaling down the ionic charges can sometimes (rarely in fact) result in enhanced ion pairing. In addition, charge scaling may require introducing correction terms for pressure calculation.

Finally, certain limitations of the ECC approach are inherent to its mean-field nature. For instance, ECC can hardly capture cross polarization between multiple charged groups, which may be important in specific binding environments. The mean-field nature of ECC also results in inherent issues when dealing with interfaces separating regions with different $\epsilon_{el}$ (see Fig. 2) or when simulating a system immersed in an external electric field.

**VII. SUMMARY AND FUTURE DIRECTIONS**

Force field molecular dynamics is an efficient simulation technique that has proven its value in providing understanding of molecular mechanisms. Nevertheless, there is increasing
amount of evidence suggesting that standard non-polarizable force fields overestimate interactions between charged groups, which are of paramount importance for many biological systems. The central problem is the lack of effective charge shielding due to the electronic polarizability induced by the surrounding environment. One solution to this problem is to use explicitly polarizable force fields. However, issues connected with parameterizations, as well as supra-microsecond time scales and large system sizes inherent in many biological problems represent a challenge for polarizable models due to their computational cost.

Several approaches have been developed to attenuate the binding between charged moieties and groups within the classical force field framework, such as tailoring specific pairwise interactions between charged moieties or including electronic polarization in a mean-field way within the ECC framework. The latter, which has been reviewed here, is a physically well justified approach. In its charge scaling variant, ECC is least intrusive, as only charges on atoms or groups contributing significantly to charge–charge interactions are scaled. This means that partial charges of weakly polarized groups can be safely left unmodified. This approach can be adopted with only minor modifications to existing biomolecular force fields and thus benefits from decades of their developments. We have shown here that, despite all current limitations, this approach has been very successful, allowing not only for a faithful description of electrolyte solutions, but also improving significantly the behavior of charged species in biosystems. Namely, it leads to correct description of ion binding to proteins and salt bridges, as well as to significant improvements in lipid–ion and lipid–protein interactions, which demonstrates the potential of the ECC approach for realistic description of complex biomolecular systems.

At present, there is no unique set of charges compatible with the ECC framework. Therefore, additional and non-standardized criteria are required. This calls for a unified protocol for force field development within the ECC framework, possibly going back to the drawing board to develop de novo a consistent and unified model of water, ions, peptides and proteins, lipids, sugars, and nucleic acids. If we find enough stamina for this immense task, this should be the focus of future development of the ECC “ecosystem”.
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