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Cite this: Phys. Chem. Chem. Phys., 2024, 26, 19887

Methodological and force field effects in the molecular dynamics-based prediction of binding free energies of host-guest systems

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As a contribution to the understanding and rationalization of methodological and modeling effects in recent host-guest SAMPL challenges, using an alchemical molecular dynamics technique we have examined the impact of force field parameterization and ionic strength in connection with guest charge neutralization on computed dissociation free energies in two typical SAMPL heavily charged macrocyclic hosts encapsulating small protonated amines with disparate binding affinities. We have shown that the methodological treatment for host neutralization, with explicit ions or with the background neutralizing plasma in the context of alchemical calculations under periodic boundary conditions, has a moderate effect on the calculated affinities. On the other hand, we have shown that seemingly small differences in the force field parameterization in highly symmetric hosts can produce systematic effects on the structural features that can have a significant impact on the predicted binding affinities.

Received 30th April 2024, Accepted 26th June 2024

DOI: 10.1039/d4cp01804d

rsc.li/pccp

1 Introduction

The SAMPL challenge (statistical assessment of the modeling of proteins and ligands) is a long-standing international initiative, started in 2008, aimed at innovating and advancing reliable predictive tools for drug design.¹⁻⁷ The challenges over the last decade have been mostly focused on host-guest systems as "widely accepted model systems to validate computational binding affinity methods and to gain insight into the physical chemistry of molecular recognition."² Typical SAMPL hosts, such as cyclodextrins, octa-acid cavitands,⁸ or pillar[*n*]arenes,^{9,10} are characterized by a concave binding surface able to accommodate drugsize molecules with a wide range of affinities in aqueous solution. The latest SAMPL9 challenge overview⁷ reported on the results of the challenge for 8 cyclodextrin-phenothiazine systems and 13 ligands of the WP6 pillar[6]arene host. It was observed that "methods applying force fields [and molecular dynamics (MD) techniques] achieved in general better correlation with experiments for WP6 [as] opposed to the machine learning and docking models". This outcome is important as it justifies the use of such computationally intensive simulation techniques as the last screening step in a high-throughput virtual screening funnel-shaped pipeline for drug design. Nonetheless, MD-based alchemical methodologies¹¹⁻¹³ for absolute dissociation free energy (ADFE) calculations yielded predictions with significant discrepancies, even when using the same force field for the host and guest models.^{7,14} According to organizers, the "origins of such differences remain unclear".

Most of the recent SAMPL9 host-guest systems, despite being designed as an effective sketch in modeling the essential features of the protein-ligand recognition mechanism, present some additional challenges due to their peculiar chemicalphysical characteristics that are not common in protein-ligand interactions. The WP6 cavitand used in SAMPL9, for example, is a heavily charged host with the symmetry-related upper and lower rim (see Fig. 1) decorated by six deprotonated carboxylate moieties at the experimental pH where binding affinity measurements are taken. The quasi-D_{6h} symmetry makes this system extremely sensitive to tiny differences in the force field parameterization, whether related to electrostatics, dispersiverepulsive, or bonded terms. Indeed, the effect of a small error in a single symmetry-related parameter can be amplified in the modelization of the bound states, with a systematic impact on the calculated ADFEs.

MD-based approaches are also strongly affected by the way charge neutralization is imposed in the system. Finite-size artifacts in alchemical simulations due to the mainstream use of periodic boundary conditions are deemed to be an important source of systematic errors for free-energy calculations in systems that involve charge changes.^{15,16} Ionic strength modelization, whether imposed using explicit counter ions or a uniform neutralizing background plasma,¹⁷ may also be an important source of systematic errors in pillar[6]arenes such as WP6 bearing-12 charges, especially when MD boxes are



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Fig. 1 SP6 and WP6 host-guest systems.

relatively small as performed in most of the MD-based SAMPL9 submissions. $^{\rm 14,18-22}$

In this study, as a contribution to the understanding and rationalization of methodological and modeling effects in hostguest SAMPL challenge outcomes, we examine the impact of force field differences and ionic strength in connection with charge neutralization on computed dissociation free energies using the recently proposed virtual double system single box approach^{23,24} (vDSSB) in two typical SAMPL systems, namely, those involving the heavily charged WP6 host and the strictly related pillar[6]MaxQ²⁵ (SP6) system (see Fig. 1), with acetyl groups being replaced by sulfonic moieties, a modification that has yet to be featured in a SAMPL challenge. The latter choice was dictated by the fact that force field-based methods in SAMPL show particularly severe errors for sulfur-containing compounds.⁷

For the SP6 system, we show that, contrary to expectations, using a uniform neutralizing background or explicit ions for host-neutralization in the bound state has little effect on the calculated ADFEs. ADFEs computed with ions or a uniform background are found to be very well mutually correlated and in decent and similar agreement with the experimental counterparts, despite the presence of sulfur atoms on the host.

In the case of WP6, we show that the systematic overestimation of the ADFEs in the vDSSB ranked submission,²⁶ originally attributed to artifacts²⁷ due to the uniform neutralizing background in the bound-state leg of the alchemical cycle, can be notably reduced by using the GAFF force field²⁸ for the WP6 host which differ significantly from those obtained using the GAFF2 in the original submission in the modelization of a "single" torsional potential involving the phenoxyacetic moieties of the rim. We show that this apparently tiny difference, with the rest of the force field involving electrostatics, dispersive–repulsion and bonded terms remaining largely unchanged, is responsible for most of the systematic overestimation observed in ref. 26.

2 Electrostatics in the MD alchemical simulation of highly charged systems

2.1 Ionic strength under periodic boundary conditions

In most of the host-guest SAMPL challenges, participants using MD-based methods relying on the alchemical protocol¹² (often scoring the best performance) adopted periodic boundary conditions with lattice sums²⁹⁻³¹ (PBC-LS) for the treatment of electrostatic interactions. Limiting ourselves to the host-guest challenges SAMPL7, SAMPL8, and SAMPL9, where participants were obliged to submit just one ranked prediction "intended to be the single entry each participant expects to be best performing",³² we note that all MD-based submissions were done using the PBC-LS protocol in combination with the particle mesh Ewald (PME) with tinfoil boundary conditions.³¹ The merits and virtues of PBC-LS compared to other methods such as those based on spherical boundary conditions (SBCs) and the reaction field have been authoritatively discussed by Sagui and coworkers in a seminal 2014 review paper.³³ Tinfoil (*i.e.* zero surface dipole²⁹) PBC-LS/ PME has become de facto a standard in the MD simulation of biological systems, supported in all the most popular MD packages such as GROMACS,³⁴ AMBER,³⁵ CHARMM,³⁶ NAMD,³⁷ LAMMPS,38 and OPENMM.39

For highly charged hosts, such as SP6 and WP6 examined in this study, a non-secondary methodological issue is related to how charge neutralization in the bound state of the host–guest system is handled. When using PBC-LS with PME, charge neutralization can be enforced either using explicit ions or using the uniform neutralizing background plasma implied in the Ewald Sum. The two approaches yield a significant difference in the nominal ionic strength of the MD box. Experimental measurements in SAMPL host–guest challenges are done in general in PBS buffer at pH = 7.4 with an ionic strength of the order of 0.2 M,^{7,40} neglecting the contributions of the host salt whose concentration is at least 3 orders of magnitude smaller than that of the buffer. The MD simulation of a single molecular system using PBC-LS, whether the guest, the host or the host–guest complex, is aimed at mimicking the conditions of an infinitely dilute solution, irrespective of the nominal concentration set by the (cubic) box of volume L^3 , provided that L is chosen large enough. This is because the use of PBC, by design, sets a minimal "invariant" distance L between the images in the lattice and no molecular encounter between images is allowed, while L is chosen large enough so that the local solvent density $\rho(r)$ at L/2 from any atom of the single-molecule system is approximately equal to the solvent density ρ . As a consequence of this fact, for "uncharged" small molecules, hydration energies computed using alchemical techniques in explicit solvents are found to be remarkably stable with varying box sizes even when using boxes with a small L (*i.e.* \simeq 12–15 Å),⁴¹ quickly reaching the limiting (infinite dilution) value for a large L.

The infinite dilution assumption and the negligible effect of the box size on solvation free energy still hold when the molecular system bears a net charge, provided that finite-size corrections due to the extra charge are accounted for ref. 17, 42 and 43 as we shall discuss further on.

The "effective ionic strength" (EIS) in an MD simulation under PBC-LS of a single molecular system is hence mainly due to the explicit ions included in the simulation box. In general, for *n* ions of a unitary charge in a cubic box with a side-length *L* (expressed in Å), the ionic strength (in M units) is given by:

$$I = \frac{1}{2} \frac{n}{L^3} 1661$$
 (1)

with $V_0 = 1661 \text{ Å}^3$ being the standard volume. Typically, SAMPL host-guest systems are simulated in a box with $L \simeq 40 \text{ Å}$, yielding an ionic strength of $\simeq 0.2$ M for the WP6 or SP6 systems with n = 12 in good agreement with the PBS experimental conditions.

Most of the MD-based submissions in the latest SAMPL7-9 challenges are done using alchemical techniques (see ref. 12 for a review on alchemical methods for ADFE calculations), whereby the guest is decoupled in a bulk solvent with the volume L_u^3 and in the solvated bound state with the volume L_b^3 . As the two decoupling processes should be done under the same thermodynamic conditions, the EIS in the two legs of the alchemical thermodynamic cycle should be the same, a fact that is often overlooked in SAMPL submissions of highly charged hosts. If the volume in the bulk simulation of the guest is L_g^3 , then the corresponding number n_g of unitary positive and negative charge pairs that must be inserted in the MD box in the bulk state to approximately match the ionic strength of the bound state for WP6 and SP6 with $L = L_{hg}$ is given by:

$$n_{\rm g} = \rm NINT \left(\frac{nL_{\rm g}^3}{2L_{\rm hg}^3} \right)$$
(2)

Given that n = 12 in both WP6 and SP6 systems, we obtain for typical box sizes of $L_g = 25-30$ Å and $L_{hg} = 40$ Å, $n_g = 2$, *i.e.* two positive and two negative monovalent ions are needed for the solvated guest in a typical MD box to match the ionic strength of the host–guest complex.

When the neutralization of the highly charged host is imposed using the uniform background, the EIS

$$I = \lim_{\Delta q \to 0} I = \Delta q Q V_0 / L_{\text{hg}}^3 = 0$$
(3)

is always zero, independent of the charge Q of the single molecular systems, whether the host-guest or guest. For this reason, neutralizing ions are generally considered the best practice¹⁶ as the use of the PBC-LS/PME neutralizing background, implying zero EIS irrespective of the host or guest charges, is considered to be inappropriate in alchemical free-energy calculations of complex heterogeneous systems.^{16,44} On the other hand, neutralizing with few monovalent counter ions can also be counterproductive since the convergence of the ion density around the charged host or host-guest system is a slow process, impeded by the ion diffusion and by the strength of the electrostatic interaction between the ions and highly charged moieties such as COO⁻ in some of the SAMPL charged hosts. These persistent interactions, if the MD simulations are not fully converged, can artificially compete with the host-guest electrostatic interaction with a systematic effect on the computed decoupling free energy of the guest molecule. Moreover, large EIS due to the use of explicit ions by hampering solvent polarization significantly reduces the dielectric constant of the solution,⁴⁵ thereby amplifying finite-size effects.^{17,30,42}

2.2 Finite size effects in the decoupling of charged guests in alchemical simulations with LS-PBC

In alchemical simulations of charged guests as those shown in Fig. 1, the formal total charge of the MD box at the initial and final end states differs. In reality, when using LS-PBC/PME, neutralization holds at any stage of the discharging or recharging process if the uniform neutralizing background is correctly accounted for (ref. 42) in the PME implementation, regardless of whether explicit ions have been included or not. As a consequence of this fact, as discussed in ref. 17 "the ion charging free energy is remarkably invariant to the system size". In ref. 42 it was shown that "even for systems with only 16 water molecules, it is possible to obtain accurate estimates of the solvation free energy of the sodium ion [via alchemical simulations]." This is because during the alchemical process, the changing uniform background is interacting with the guest as well as with the solvent molecules and with explicit ions (if present). The effect of the background on the guest is hence screened by the dielectric environment and the finite size effects on the solvation free energy are small provided that L is large enough and solvent polarization is strong.⁴⁶

Finite-size effects in the discharging of charged molecules or ions during alchemical simulations under PBC-LS have been thoroughly discussed in several seminal papers from the end of the past century.^{17,30,42,43,46} Briefly, the energy (in kcal mol⁻¹) computed in PBC-LS/PME of a single ion to bring its charge (in *e* units) from q_{beg} to q_{end} in a cubic MD box with side-length *L* (in Å) is given by the so-called Wigner energy:¹⁷

$$\Delta G_{\rm ew} = 332 \left(q_{\rm beg}^2 - q_{\rm end}^2 \right) \times \frac{\xi}{2L} \tag{4}$$

with ξ = 2.837297. Note that this strongly size-dependent energy is positive for the discharging process and negative when the charge of the ion is increased.

In PME, this size-dependent Wigner energy is given by a reciprocal lattice contribution and a direct lattice contribution. The former, ΔG_{recpr} , is automatically included in the standard PME implementation¹⁷ while the latter is sometimes added *a posteriori*^{19,20} and is given by

$$\Delta G_{\rm dir} = 332 \frac{(q_{\rm beg}^2 - q_{\rm end}^2)\pi}{2\alpha^2 L^3}$$
 (5)

where $\boldsymbol{\alpha}$ is the Ewald convergence parameter.

The free energy of discharging or recharging an ion *in vacuo*, however, must be equal to zero. To account for this fact, Figuerido *et al.*⁴³ and Hummer *et al.*⁴⁶ proposed to add an empirical "thermodynamic" size-dependent correction when computing the ion solvation free energy in an explicit solvent of a dielectric constant ε such that

$$\Delta G_{\text{therm}} = 332 \left(q_{\text{beg}}^2 - q_{\text{end}}^2 \right) \times \left[-\frac{\xi}{2\varepsilon L} + \pi \frac{(\varepsilon - 1)}{\varepsilon} \frac{R^2}{3L^3} \right] \quad (6)$$

where R is the ion radius. By adding eqn (6) and (4), we obtain the Hummer finite-size correction due to the charge change for PBC-LS alchemical simulations with the Ewald sum as follows:

$$\Delta G_{\text{finite-size}} = 332 \frac{(\varepsilon - 1)}{\varepsilon} (q_{\text{beg}}^2 - q_{\text{end}}^2) \times \left[\frac{\zeta}{2L} - \pi \frac{R^2}{3L^3}\right] \quad (7)$$

Note that, when $\varepsilon = 1$, $\Delta G_{\text{finite-size}}$ vanishes as it should be when changing the charge of an ion *in vacuo*. For highly polar solvents such that $(\varepsilon - 1)/\varepsilon \simeq 1$ and for small ions compared to *L*, eqn (7) is essentially equal to eqn (4). As remarked by Hummer,⁴² "this explains the success of using simply the bare eqn (4) implied in the Ewald sum for the free energy of charging an ion in a highly polar environment, without further finite-size correction that takes the ion size or the dielectric constant of the solvent into account."

The problem of the finite-size effects to the calculation of the charging free energy in the context of alchemical simulations of complex host-guest or ligand-protein systems (rather than simple monoatomic ions) has been re-examined in many instances in the past decades,^{15,16,47-50} proposing a plethora of L-dependent correction terms based again on the continuum model, sometimes requiring costly extra calculations. Strikingly, in the latest of these series of studies,¹⁶ dealing with alchemical MD simulations for systems as complex as watersolvated buckyball derivatives hosting charged molecular guests, finite size free energy corrections in charged simulation boxes with or without explicit salt ions were found to be systematically small (less than 1 kcal mol^{-1}) even for small boxes (i.e. with a minimal distance of any solute atom from the wall box of the order of 10 Å). The L-invariance of alchemical free energies under PBC-LS including only the Wigner term eqn (4) in host-guest systems involving charge changes was also numerically verified in SAMPL submissions.19,21

3 Force field effects on the ADFE of host-guest systems

The force field effect on ADFE calculations using alchemical simulations is a long-standing and complicated issue not only in host–guest SAMPL challenges⁷ but in general in computational drug design.^{51–54} Limiting ourselves on the host–guest MD-based modelization, the commonly adopted force fields by participants^{28,39,55–57} are based on an analytical expression of the form:

$$V = \sum_{\text{Bonds}} K_{\rm r} (r - r_0)^2 + \sum_{\text{Angles}} K_{\theta} (\theta - \theta_0)^2 + \sum_{\text{Dihedrals}} V_{\phi} [1 + \cos(n\phi - \gamma)] + \sum_i \sum_{j>i} \left[\frac{q_i q_j}{r_{ij}} + v_{\rm LJ} (r_{ij}) \right]$$
(8)

where the first three terms refer to the bonded potential, including stretching, bending, and torsional terms, and the last term is the non-bonded contribution to the force field, given by a sum on all non-bonded atom–atom pairs of dispersive–repulsive and electrostatic interactions, with the latter treated using PBC-LS. The parameters in eqn (8) are based on the definition of atomic classes depending on the chemical environment and atom hybridization and are derived from *ab initio* calculations and experimental data,^{51,58} most recently in combination with machine-learning techniques.^{59,60}

With a few exceptions, the atomic charges in the MD-based SAMPL submissions are fixed, derived using a variety of ab initio or empirical schemes.⁶¹⁻⁶³ In the real world, the electron density on a molecule is constantly modulated by charge reorganization due to the conformational activity of the host, guest, their mutual polarization and the polarization of the solvent. Fixed atomic charges are meant to account for the environment polarizability and for the charge reorganization due to conformational changes in a mean-field spirit.^{64,65} This approach in modeling electrostatic interactions is widely considered a decent compromise between accuracy and efficiency as the explicit accounts for atomic charge modulation would imply the use of polarizable force fields involving manybody effects. In some cases,^{21,22,66} the polarizable force field AMOEBA⁶⁷ has been used in SAMPL challenges, often with remarkable results.^{5,7} However, polarizable force fields and charge fluctuating simulations are much more demanding than the fixed charge MD protocol, casting doubts on the benefit-cost ratio of these techniques as the last screening stage in the HTVS pipeline for drug design.

4 Methods

4.1 Simulation setup and parameters

The SP6 host has yet to be examined in SAMPL challenges. SP6 guests are shown in Fig. 1. We applied the GAFF2 force field for the host and guest modelization, using the PrimaDORAC online interface⁶⁸ for atomic type assignment. Atomic RESP^{62,63} charges on the host and guests were computed using the



Fig. 2 Left: Highlighted quadruplets for the O-C-C-O torsions in WP6. Right: Torsional energy for the O-C-C-O torsions in GAFF, GAFF2, and in the GAFF2 combination (see the text).

Antechamber AMBER suite.⁶⁹ MD simulations were conducted in the NPT ensemble under standard conditions using an isotropic Parrinello-Rahman Lagrangian⁷⁰ and a series of Nosé thermostats⁷¹ for pressure and temperature control, respectively. For the simulation of the bound (host-guest complex) and unbound (guest in bulk) states, cubic boxes of a standard size were used with $L \simeq 38 \text{ Å}^3$ and $L \simeq 25 \text{ Å}^3$ (after equilibration of 100 ps), respectively, filled with OPC3⁷² water molecules. Bond constraints were imposed on X-H bonds only, where X is a heavy atom. Charge neutralization was imposed using the PBC/LS/PME neutralizing background (zero EIS) and using explicit ions (EIS = 0.2 M). In the latter case, in the bound state, we added 12 Na⁺ to neutralize the charge of the host, while in the unbound state two Na⁺ ions and two Cl⁻ ions were added to approximately match the same EIS as that of the bound state (see Section 2.1). The mass of the ions, irrespective of their type, was set to 4 amu to enhance the passive diffusion of the ions with the scope of accelerating the convergence of the ionic distribution in the bound and unbound states.

For the WP6 host-guest system, we selected the guests (reported in Fig. 1) that gave the largest systematic overestimation compared to the experimental ADFEs in our SAMPL9 submission.²⁶ We used GAFF2⁵⁵ with AM1-BCC charges for the guests as in ref. 26. For the WP6 host, we used instead the GAFF parameterization as obtained from PrimaDORAC⁶⁸ with AM1-BCC charges. Both GAFF and GAFF2 for the WP6 host are based on eqn (8) and have identical AM1-BCC charges, atom types, and very similar bonded and non-bonded parameters. They differ significantly only in the parameters of the O-C-C-O torsion involving the quadruplet connecting the -CH₂COO⁻ charged groups to the phenyl rings⁷³ (see Fig. 2). In GAFF, this torsion is zero, while in GAFF2 this torsion has three non-zero terms with multiplicity 1, 2, and 3. The resulting torsional potentials for GAFF and GAFF2 are shown in Fig. 2. For each O-C-C-O, there are two O-C-C-O combinations and the total number (12) of -CH₂COO⁻ groups lead to 24 O-C-C-O combinations. The two combinations for a given -CH₂COO⁻ group differ by a phase factor of 180°. The torsional energy resulting from the combination of the two O–C–C–O GAFF2 torsions is shown in green in the plot of Fig. 2 with a barrier of 2.5 kcal mol^{-1} between the two minima at 0 and 180° . We hence expect the distribution of the host O–C–C–O dihedral angle in GAFF2 to be markedly different compared to that observed with GAFF where the O–C–C–O rotation is unrestrained.

The MD protocol (equilibration, thermodynamic conditions, constraints, and water model) is the same as that used for SP6 host–guest batch.

4.2 The vDSSB approach for absolute dissociation free energies

The vDSSB methodology has been thoroughly described in ref. 23, 24, 26 and 74. Briefly, the method consists of two massively parallel computational steps independently applied to the two legs of the alchemical cycle, namely the HREM stage and the nonequilibrium alchemical stage. The HREM stage is aimed at collecting the equilibrium enhanced sampling of the host-guest bound state (with the ligand at full coupling). The initial configurations for the unbound state are obtained by combining the HREM-sampled gas-phase (decoupled) ligand configurations with a pre-equilibrated box filled with explicit water. The HREM technical specification is identical to those adopted in ref. 26. For the NE stage, starting from the HREM sampling of the end states, we launched independently two swarms of 500 and 250 NE alchemical trajectories where the ligand was decoupled in the bound state and recoupled in the unbound state, respectively. The bound state and unbound state annihilation trajectories lasted for 720 ps and 360 ps, respectively. The resulting NE work distributions were combined to produce a sample of 125 000 work values, yielding the final highly resolved vDSSB convolution of the two bound and unbound distributions. The vDSSB work convolution refers to a NE "unidirectional" process, corresponding to the host-guest dissociation, where the ligand is annihilated in the bound state while materializing in the far distant bulk solvent, with the possibility, by design, of calculating independently the decoupling and recoupling contributions. The dissociation free energy is recovered from the work distribution

produced by the convolution using well-known nonequilibrium theorems.^{75,76}

4.3 Further technical details

All calculations were done using the open-source ORAC program,⁷⁷ freely available at the site **https://www.chim1.unifi.it/orac**. All data, including MD input parameters for the HREM and NE computational stages, enhanced sampling PDB trajectories, work distributions, and software for input preparation and for post-processing the work data, along with essential documentation, are available at the general-purpose open-access repository Zenodo (https://zenodo.org/records/11074240) to enable anyone willing to do so to reproduce our data.

5 Results and discussion

5.1 The SP6 host-guest system: ionic strength effects

In this host-guest batch, all guests have a single positive charge on the amino moiety (see Fig. 1) with the host bearing 12 negative charges. We hence expect the finite size effects due to the treatment of the electrostatic interactions and ionic strength to be remarkable when using MD boxes of standard sizes (i.e. with $L = \simeq 38$ and $L = \simeq 25$ for the bound and unbound state, respectively). To ascertain the effect of the ionic strength, possibly intertwined with finite-size effects related to charge annihilation or creation, we performed two full vDSSB calculations with and without explicit ions. In the first calculation, no counter ions were inserted in both the bound and unbound states, with the neutralizing PME background plasma yielding an EIS of zero (see eqn (3)) irrespective of the total nominal charge in the MD box. In the second calculation, we inserted 12 Na⁺ host-neutralizing ions in the bound state and two Na⁺ and Cl⁻ ions in the unbound state to match the ionic strength (see eqn(1) in the two legs of the alchemical thermodynamic cycle.

Unlike in the WP6 batch, for the SP6 system, we noticed from the preliminary unrestrained HREM simulations of the Vinaprepared bound state that bulky guests tend to significantly populate poses where the center of mass (COM) of the guest molecule is mostly lingering outside the host toroidal cavity. To allow for the sampling of these poses (that are in any case included in a SAMPL isothermal titration calorimetry measurement⁴⁰), in the bound state, we used a weak harmonic restraint potential with a force constant of K = 0.05 kcal mol⁻¹ Å⁻² between the COM of the host and the guest, with a guest allowance volume of $V_{\text{restr}} = (2\pi RT/K)^{3/2} \simeq 600$ Å³. We then computed standardly the free energy correction to the annihilation free energy of the bound guest due to the restraint^{78,79} as $\Delta G_{\text{rest}} = RT \ln V_{\text{restr}}/V_0$ for all guests, as done in most of the recent alchemy-based SAMPL submissions.^{32,80,81}

Host charge neutralization with or without ions has profound and troubling implications in the electrostatic potential at the host center. In Fig. 3, we report the ionic radial distribution function from the COM of the host of the bound state of the SP6 host-guest systems, obtained from the HREM simulations with explicit ions and with the background (no counter ions). We note that $\simeq 9$ explicit ions on average are found



Fig. 3 Ionic radial distribution, g(r), from the COM of the SP6 host in the equilibrium bound state for all 11 SP6-guest systems of Fig. 1 with explicit ions. Dashed lines are the integral $n(r) = \frac{4\pi n}{L^3} \int_0^r g(u) u^2 du$ of the g(r). The thick black lines refer to the average g(r) and n(r) for all host–guest systems with explicit ions. The magenta line is the result obtained with the background only.

within a sphere of radius 18 Å (and volume $\simeq L^3/2$) centered on the COM of the host, with a high local density near the outer side of the SO₃-decorated rim (with a radius of $\simeq 5.5$ Å). The g(r)of the neutralizing background (magenta lines) is equivalent to a uniform distribution of -12 charges and is equal to 1 everywhere, with an integral reaching 6 units at $r = 18 \simeq L/2$ Å.

These striking differences in the ionic distribution in the two approaches for host neutralization (with ions or with the background) can indeed translate into significant discrepancies in the discharging free energy of the ligand in the bound state. In Table 1, we show the finite-size corrections calculated for the bound and unbound states according to eqn (4) (Wigner energy), eqn (5) (direct lattice Wigner correction) and eqn (6) (Hummer-Figuerido correction) with explicit ions and with the background. The latter correction was calculated with a mean gyration of the guest of R = 3.4 \pm 0.3 Å, and ε = 80. Taking into account that the reciprocal lattice contribution to the Wigner energy ΔG_{ew} is automatically included in vDSSB calculations when using PME and need not to be considered,¹⁷ the finitesize correction to the bound and unbound states for SP6 includes the a posteriori Wigner corrections in the direct lattice $\Delta G_{\rm dir}$, eqn (5), and the thermodynamic corrections $\Delta G_{\rm therm}$, eqn (6), in the two legs of the alchemical cycle, the latter leading

 $\label{eq:table_transform} \begin{array}{l} \textbf{Table 1} & \mbox{Finite-size corrections to the dissociation free energy (kcal mol^{-1})} \\ \mbox{in the SP6 host-guest systems with and without explicit ions} \end{array}$

	$\langle L \rangle$	α	Grid	Nions	ΔG_{ew}	$\Delta G_{ m dir}$	$\langle \Delta G_{ m therm} angle$
With ion	S						
Bound	37.0	0.37	1.17	12	12.40	0.06	0.0
Bulk	24.8	0.38	1.03	4	18.84	0.21	0.0
Total	n/a	n/a	n/a	_	-6.44	-0.15	0.0
With bac	kgroui	nd					
Bound	37.8	0.37	1.17		-285.1	-1.37	1.9 ± 0.3
Bulk	24.9	0.38	1.03		18.84	0.21	0.0 ± 0.04
Total	n/a	n/a	n/a		266.26	-1.16	$\textbf{1.9}\pm\textbf{0.3}$

Table 2 Dissociation free energies (kcal mol⁻¹) in the SP6 host-guest systems computed by way vDSSB with the explicit ions and with the PME neutralizing background. Errors were computed by bootstrapping with resampling on the bound and unbound distributions before performing the convolution. The experimental values are taken from ref. 25

		With counter ions	$\frac{\text{With background}}{\Delta G_{\text{vDSSB}}}$	
Guest	ΔG_{exp}	$\Delta G_{ m vDSSB}$		
L24	9.5	11.5 ± 0.5	11.3 ± 0.8	
L25	10.9	10.5 ± 0.9	9.3 ± 1.1	
L26	8.5	9.5 ± 0.6	8.1 ± 0.3	
L27	7.0	7.8 ± 0.9	7.8 ± 1.3	
L28	10.6	8.9 ± 0.9	7.8 ± 0.7	
L29	8.3	6.2 ± 0.6	4.9 ± 0.7	
L30	8.3	5.8 ± 1.8	4.4 ± 1.0	
L31	6.8	4.5 ± 0.6	4.6 ± 0.8	
L32	7.8	6.8 ± 1.3	3.5 ± 0.8	
L33	9.9	11.1 ± 0.7	10.8 ± 1.0	
L34	10.3	10.9 ± 0.7	11.6 ± 0.7	

to a negligible correction for the ADFE when explicit ions are included.

When neutralization is handled via the background (i.e. without explicit ions), due to a charge change for all ligands of Fig. 1 from -11 to -12e in the bound state, the total direct lattice term is negative (as with explicit ions) and equal to $\simeq -1.2$ kcal mol⁻¹, whereas the total thermodynamic term yields an average ''positive'' correction term of 1.9 \pm 0.3 kcal mol to the ADFE. Had one used this correction in the SAMPL9 vDSSB submission²⁶ where a systematic overestimation of $\simeq 3$ kcal mol⁻¹ of the binding strength was observed, such systematic overestimation would have increased, significantly worsening the accuracy of the method. In ref. 26 it was hypothesized that the observed overestimation of the ADFE was due to the fact that the uniform background charge ρ_{BG} artificially causes²⁷ the guest with charge q_g to favor the lower dielectric environment region inside the toroidal cavity with dielectric constant ε_1 . Probably, this effect could be related to the observed significant improvement of the calculated binding affinity of continuum model approaches when adjusting the cavity dielectric ε_{in} in host-guest systems.⁸² Hub et al.²⁷ provided a simplified negative correction to the ADFE, based on the Poisson–Boltzmann model, for the free energy difference in the two dielectrics (low dielectric environment and bulk water), proportional to $q_g \rho_{BG} R^2 / \epsilon_1$. Additional and subtle corrections within PBC-LS/PME due to the finite size and shape of the ionic system in the final and initial states of the alchemical transitions¹⁶ may hence cancel out, in light of the remarkable insensitiveness of solvation free energies observed in a strongly polar solvent for both uncharged⁴¹ and charged species, whether monatomic or molecular ions,^{16,22,42,46} without any other correction but the Wigner energy eqn (4).

Therefore, in computing the host-guest ADFE for the SP6 system, as done in our study and other successful SAMPL submissions, $^{19-21,26}$ we only included the direct lattice finite-size Ewald corrections (eqn (5)) shown in bold in Table 1, neglecting altogether the thermodynamic empirical correction (eqn (6)) and other continuum-model corrections. 15,27,48,50

In Table 2, we report the results for the calculated ADFEs of the host–guest SP6 system obtained with the two vDSSB calculations, namely with counter ions and with the background. In Fig. 4, we show the corresponding correlation plots for the calculated vDSSB dissociation free energies with the experimental counterpart,²⁵ obtained with and without counter ions, and their mutual correlation. In general, we note that the correlation with the experiment, as measured by the Pearson correlation coefficient *R* and by the Kendall rank coefficient τ , is better when using explicit ions.

In both cases, vDSSB tends to underestimate the host–guest dissociation free energy, yielding mean signed errors (MSEs) of 1.26 and 0.50 kcal mol⁻¹ when using the uniform background and with counter ions, respectively. Strikingly, the mutual correlation of the two vDSSB free energies (see Fig. 4(c)) is excellent, resulting in R = 0.94, $\tau = 0.74$, and an MSE of 1.0 kcal, possibly due to unaccounted (and small) finite-size effects when using the uniform background. Incidentally, we note that the inclusion of the thermodynamic correction of $\simeq 1.9$ kcal mol⁻¹ when using the background plasma would have brought the ADFEs of the four bulky ligands L28–L32 closer to the experimental data, while further worsening the results for all the other ligands, yielding a correlation coefficient with the



Fig. 4 SP6 correlation plots: (a) background, (b) with counter ions and (c) counter ions vs. uniform background correlation. R: Pearson's correlation coefficient; MAE: mean average (unsigned) error in kcal mol⁻¹; MSE = $\langle \Delta G_{exp} - \Delta G_{vDSSB} \rangle$: mean signed error in kcal mol⁻¹, τ : Kendall's rank coefficient.

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experiment of R = 0.65 and an MSE in the opposite direction of -0.6 kcal mol⁻¹.

We were indeed surprised at the small differences, often within the confidence interval, between the two calculations with zero EIS (the background) and realistic EIS set by the explicit counter ions, notwithstanding the notable differences in the ionic distributions in the two cases (see Fig. 3). The small effect, on the dissociation free energies of host-guest systems computed with alchemical techniques, of the insertion of explicit counter ions with moderate EIS *in lieu* of the uniform background in host-guest systems was also recently observed in ref. 16, 21 and 22.

The results shown in Fig. 4 and Table 2, as well those reported in recent studies,^{16,21,22} can have important implications in the practical application of the alchemical simulation in drug design. Charge changes in ligand-protein systems, even when using the simple background plasma and no counter ions, are much less pronounced than those in the bound state of the SP6 or other SAMPL host-guest systems and MD boxes are significantly larger, so that finite-size effects account for corrections to the ADFE well below 1 kcal mol⁻¹.¹⁶ MD simulations in PBC-LS/PME for computing ADFEs are, after all, only a model for the real system where many physical effects are not and cannot be included if not devising corrections based on crude approximations (such as the continuum model) relying on a multitude of arbitrary and system-dependent parameters. The goodness of the model should be judged, in the spirit of the SAMPL challenge, by the quality of its predictions in connection with the computational cost, and by the simplicity and smoothness of its application.

5.2 The WP6 host-guest system: force field effects

We have seen in the previous section that the imposition of a realistic EIS in the SP6 host-guest system through explicit ions, while somewhat improving the agreement with the experiment, does not entail dramatic changes in the computed dissociation free energies (see Fig. 4(c)). The latter is found higher by $\simeq 1$ kcal mol⁻¹ when ions are included. It is not clear why binding, on average, is more favorable in the SP6 system when ions are included. Based on Tables 1 and 2 referring to SP6, we may nonetheless infer that adding explicit ions to the strictly related WP6 host-guest system, where dissociation free energies computed with the background were systematically overestimated,²⁶ would likely further worsen the accuracy of the method.

In ref. 73 it was shown that the structural and dynamical behaviors of the $-CH_2COO^-$ tails in the *apo* form of WP6 was dramatically altered when switching from the GAFF force field to the GAFF2 force field. As discussed in ref. 73 and in the "Methods" section, the only significant difference between the GAFF and GAFF2 modelization for the WP6 host involves just one torsion, *i.e.* the O-C-C-O torsion with the two terminal atoms referring to carbonyl and esteric oxygen atoms (see Fig. 2). Such a seemingly minor difference in the two WP6 parameterizations, amplified by the presence of twelve equivalent $-CH_2COO^-$ moieties, yielded a much open average configuration of the $-CH_2COO^-$ tails with the



Fig. 5 o-c-ct-os torsional distribution obtained with the GAFF and GAFF2 force fields from the HREM simulation of the *apo* form of WP6 in water.

GAFF force field compared to GAFF2, which is attributed⁷³ to the host–solvent and host–explicit ion interactions.

In Fig. 5, we show the effect of this difference on the overall distribution of the 24 O–C–C–O dihedral angles obtained with GAFF and GAFF2 from the HREM simulations of the *apo* form of WP6 under standard conditions without including explicit ions, *i.e.* using the same host neutralization protocol of the original vDSSB SAMPL9 submission.²⁶ The GAFF2 distribution exhibits peaks at 0° and $\pm 180^{\circ}$, whereas the GAFF distribution has maxima at $\pm 100^{\circ}$. The maxima of the GAFF2 dihedral probability density are in full accordance with the potential energy of the GAFF2 torsion reported in Fig. 2 exhibiting minima at 0° and $\pm 180^{\circ}$. In GAFF, the O–C–C–O torsional potential is zero and the unhindered rotation about the C–C bond is modulated by the other interacting terms inside the system resulting in a more uniform and spread-out distribution.

As observed in ref. 73 where WP6 was simulated including explicit ions, the absence of the O–C–C–O torsional potential in GAFF translates in a more open and disordered behavior of the



Fig. 6 OO-distance distribution obtained with GAFF and GAFF2 force fields from the HREM simulation of the *apo* form of WP6 in water.

Table 3 Dissociation free energies (kcal mol⁻¹) computed by way vDSSB using the GAFF force field and the GAFF2 force field for the WP6 host. Errors were computed by bootstrapping with resampling on the bound and unbound distributions before performing the convolution. GAFF2 ADFEs are taken from ref. 26, $\Delta G_{\rm Exp}$ values for G2–G10 are taken from ref. 83, and $\Delta G_{\rm Exp}$ for G14, G15, G16, and G17 are taken from ref. 84–86

Guest	$\Delta G_{ m exp}$	$\Delta G_{ m vDSSB}$ (GAFF2)	$\Delta G_{ m vDSSB}$ (GAFF)
G02	10.59	17.0 ± 0.4	14.0 ± 1.3
G03	8.03	11.5 ± 0.6	12.1 ± 0.6
G06	8.08	10.4 ± 1.3	11.5 ± 1.1
G07	7.07	12.2 ± 1.7	11.2 ± 1.6
G08	6.04	12.2 ± 1.6	9.4 ± 1.1
G09	6.32	12.5 ± 0.6	10.7 ± 1.1
G10	9.96	13.5 ± 1.1	12.9 ± 0.7
G14	9.68	17.2 ± 0.4	11.8 ± 0.4
G15	8.37	11.2 ± 1.6	8.0 ± 0.8
G16	10.59	14.5 ± 1.5	10.5 ± 0.9
G17	6.48	7.9 ± 0.8	$\textbf{7.1}\pm\textbf{0.9}$

 $-CH_2COO^-$ tails compared to GAFF2. This can be appreciated in Fig. 6 where we show the probability distribution of the distances between opposite oxygen atoms in the upper and lower rims obtained with the two force fields. The maximum of the distribution in GAFF2 occurs at a shorter distance (of $\simeq 2$ Å) than in GAFF. Correspondingly, by aligning the HREM sampled structures on the six connecting carbon atoms of the ring methylene groups of one of the rims, the resulting distribution of the $-CH_2COO^-$ tails (shown in the circles of Fig. 6) is much more disordered and stretched out in GAFF than that obtained with the GAFF2 force field.

This strikingly different behavior produced by the O–C–C–O torsion in GAFF and GAFF2 must likely give rise to a systematic impact on the calculated binding free energy. When the hydrophobic scaffold of a WP6 guest (see Fig. 1) enters the host cavity, we might infer that the inward and more rigid configuration of the –CH₂COO⁻ tails in GAFF2 could entail a decrease in the k_{off} , due to the stronger interaction between the carboxylate and the protonated amino groups of the guest pointing outside the WP6 torus. As the binding poses for most of the 11 guests are characterized by non-polar moieties trapped into the WP6 cavity and exposed amino groups interacting with the

 $-CH_2COO^-$ tails of the rim, this structural feature in GAFF2, with a tightly restrained guest, could eventually translate into higher dissociation free energies compared to GAFF.

To verify this hypothesis, we repeated here the vDSSB calculations presented in the SAMPL9 challenge^{7,26} on WP6 with the only difference that we adopted the GAFF force field for the host, while maintaining the GAFF2 force field for the guests and the OPC3 model for water. We emphasize that the atomic (AM1-BCC) charges for the host and the guests are identical in the present calculation and that of ref. 26 as well as the other methodological parameters connected to the use of the background neutralizing plasma, the standard volume correction, the HREM protocol for the bound state, the duration and decoupling/recoupling of the NE trajectories. The comparison of the two vDSSB ADFEs for the hostguest WP6 system is hence aimed at singling out the effect of the O–C–C–O torsion on the computed vDSSB dissociation free energy.

The results are collected in Table 3. We notice that in general the ADFEs computed with GAFF for the host are smaller than those computed with GAFF2.

In Fig. 7, we show the corresponding correlation plots with the experimental data and the mutual correlation between the two vDSSB estimates. As inferred, the rigidity of the host $-CH_2COO^-$ tails with GAFF2 appears to favor binding in most of the cases. The size of the resulting overestimation is nearly halved, passing from -4.44 kcal mol⁻¹ with GAFF2 MSE to -2.54 kcal mol with GAFF. Ranking agreement with GAFF as measured by τ also increases, while we see a degradation of the *R* Pearson coefficient. Overall, these results neatly show that seemingly small differences in the force field parameters for highly symmetric hosts, like those referring to the single O–C–C–O WP6 torsion in GAFF and GAFF2 (see Fig. 2), may produce systematic errors in the estimates that are far larger than the errors related to methodological issues such as those examined in the preceding section for SP6.

6 Conclusions

We have analyzed the effect of the ionic strength and of the force field parameterization in the calculations of the absolute



Fig. 7 WP6 correlation plots: (a) with GAFF2, (b) with GAFF and (c) GAFF2 vs the GAFF correlation. R: Pearson's correlation coefficient; MAE: mean average (unsigned) error in kcal mol⁻¹; MSE = $\langle \Delta G_{exp} - \Delta G_{vDSSB} \rangle$: mean signed error in kcal mol⁻¹, τ : Kendall's rank coefficient.

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dissociation free energies of the highly charged host-guest systems (see Fig. 1), typically adopted in the SAMPL challenges, via an MD-based alchemical technique. We have shown that the methodological treatment for host neutralization (with explicit ions or with the background neutralizing plasma in the context of PBC-LS/PME) has a moderate effect on the calculated ADFEs, irrespective of whether finite-size corrections derived from continuum models and related to charge change of the guests are included or not. On the other hand, we have shown that seemingly small differences in the force field parameterization in some cases might be quite significant. In this regard, the systematic overestimation of ADFEs observed in ref. 26 for the WP6 host-guest system can be attributed in large part to a single torsional potential involving the -CH₂COO⁻ moieties of the host alone. Atomic charge parameterization may also play an important role, showing better agreement with those in the experiments observed in SP6, where we used RESP charges, compared to WP6, where we used AM1-BCC charges.

This study, in accordance with other SAMPL predictions, shows that the confidence intervals of MD-based alchemical ADFE estimates in host-guest systems, if judiciously implemented, are of the order of 2 kcal mol^{-1} and this is probably the best that we can currently perform with standard force fields of the form of eqn (8) adopted in most SAMPL submissions throughout the years. Explicit ions matching the ionic strength of the system should certainly be considered for improving the estimates. However, such a consensus on the methodological protocol does not appear as a decisive factor even in critical/ pathological systems such as highly charged SP6 and WP6. The 2 kcal mol⁻¹ uncertainty in the MD-based free energy translates into 1.5 pK_d units, with a correlation (R > 0.5) that is in general higher than that obtained with end point, docking, and machine learning approaches. However, this uncertainty and correlation are still not optimal for reliably using MD-based alchemical methodologies as the last screening step in the HTVS pipeline in industrial settings for drug design.

Our study, as well as the bulk of SAMPL challenge submissions in the last decade, shows that efforts should be focused on improving the force field parameterization. In ligand-protein systems, apparently minor deficiencies in the force field, even when connected with mismatches in the assignment and handling of the parameters from force field databases in automatic procedures,⁸⁷ can make big differences in the calculated ADFEs, most likely larger than those related to methodological aspects such as the imposition of the ionic strength via counter ions and/or finite-size correction in the PBC-LS context. Concerning this point, it should be noted that, in the latest SAMPL challenges, the "polarizable" AMOEBA force field⁶⁷ in combination with a standard MD-based alchemical methodology was consistently found among the top-performing methods. In AMOEBA, electrostatic interactions explicitly include polarization effects via distributed atomic polarizabilities. Polarization effects, related to induction phenomena or molecular charge reorganization due to conformational changes, can indeed be decisive in shaping the binding affinity, especially in highly charged host-guest systems such as WP6 and

SP6. The computational burden of the AMOEBA many-body protocol is still too high compared to the standard approach with fixed atomic charges, making the method unfit in HTVS pipelines for the time being. However, AMOEBA's success in SAMPL challenges shows, in our view, that the future endeavor in MD-based methodologies for improving the reliability of ADFE estimates in ligand-receptor systems should be mostly devoted to the development of efficient atomic charge fluctuating techniques that account in a realistic fashion for the polarizability feedback between drugs, receptors and solvents in the process of molecular recognition and induced fit.

Data availability

All data for reproducing the results presented in this study, including trajectory files, starting structures, force field parameters, template input files, and ancillary software for preparing the HREM and NE stages on HPC platforms are available at the public repository Zenodo https://zenodo.org/records/11074240.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

The computing resources and the related technical support used for this work have been mostly provided by CRESCO/ ENEAGRID High Performance Computing infrastructure and its staff. The CRESCO/ENEAGRID High Performance Computing infrastructure is funded by ENEA, the Italian National Agency for New Technologies, Energy and Sustainable Economic Development and by the Italian and European research programmes (see https://www.cresco.enea.it for information). Part of the calculations were performed on the LEONARDO HPC cluster, for which we acknowledge the CINECA award under the ISCRA initiative (HP10CV2G9I), for the availability of high-performance computing resources and support. PP acknowledges the National Recovery and Resilience Plan, Mission 4 Component 2 - Investment 1.4 - NATIONAL CENTER FOR HPC, BIG DATA AND QUANTUM COMPUTING - funded by the European Union - NextGenerationEU - CUP B83C22002830001.

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