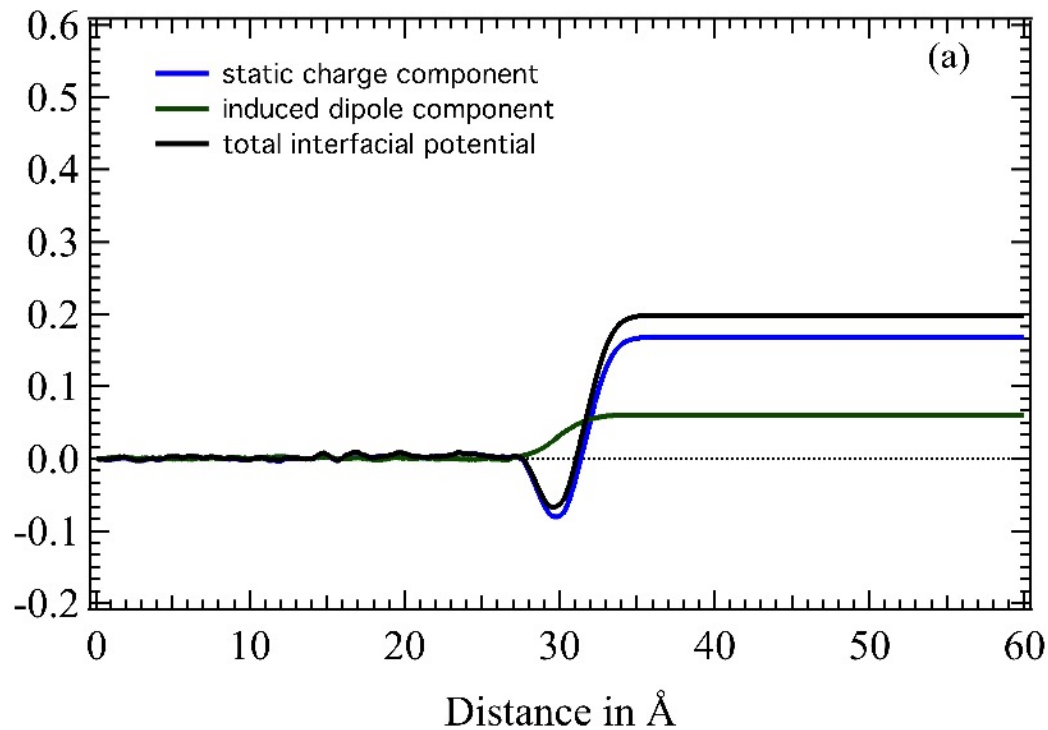
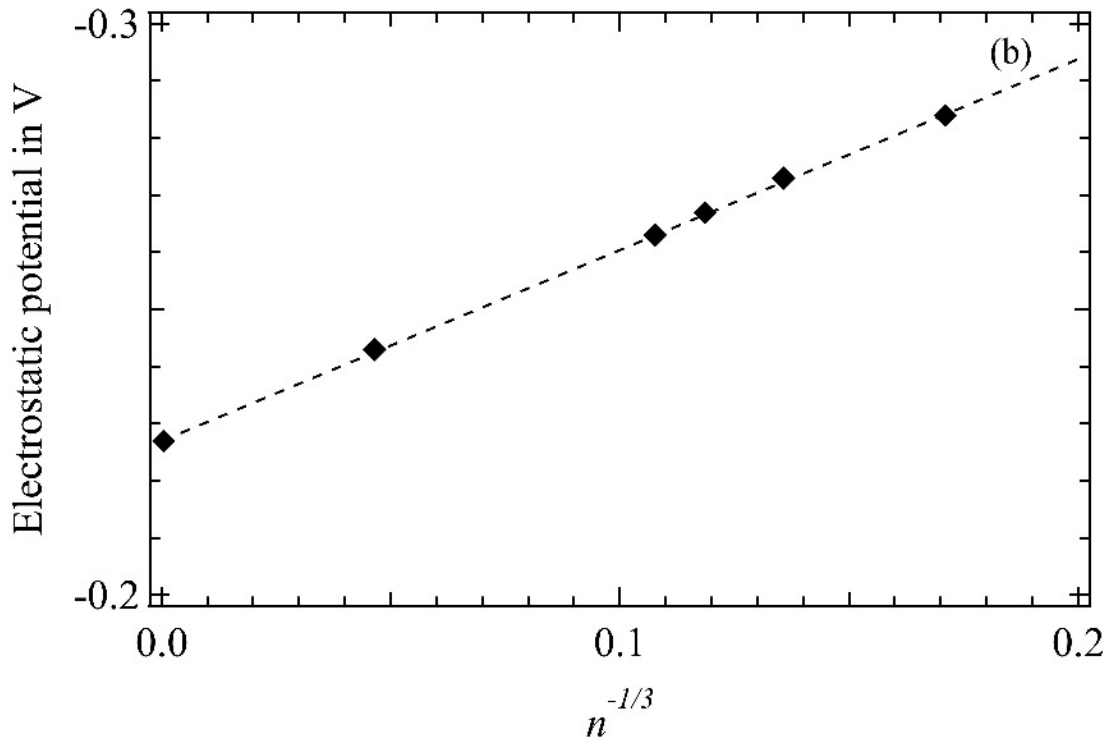


Electrostatic potential in V





Ion hydration free energies and water surface potential in water nano drops. The cluster pair approximation and the proton hydration Gibbs free energy in solution.

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Abstract

We estimate both single ion hydration Gibbs free energies in water droplets, comprising from 50 to 1000 molecules, and water/vacuum surface potentials in pure water droplets comprising up to 10 000 molecules. We consider four ions, namely Li^+ , NH_4^+ , F^- and Cl^- and we model their hydration process and water/water interactions using polarizable force fields based on an induced point dipole approach. We show both ion hydration Gibbs free energies and water surface potentials to obey linear functions of the droplet radius as soon as droplets comprising a few hundred water molecules. Moreover we also show that the differences in anion/cation hydration Gibbs free energies in droplets obey a different regime in large droplets than in small clusters comprising no more than six water molecules, in line with earlier results computed from standard additive point charge force fields. Hence point charge and more sophisticated induced point dipole molecular modeling approaches both suggest that methods considering only the thermodynamical properties of small ion/water clusters to estimate the absolute proton hydration Gibbs free energy in solution are questionable. In particular taking into account the data of large ion/water droplets may yield a proton hydration Gibbs free energy in solution value to be shifted by several $k_B T$ units compared to small clusters-based approaches.

19 Introduction

20 Ion hydration plays a pivotal role in many research fields from physics (like new high density
21 batteries¹), to chemistry (to understand pollution phenomena² and climate change,³ for
22 instance), and biology.^{4,5} Despite decades of research, estimating ion hydration Gibbs free
23 energies is far from being a routine activity. This arises from the difficulties with today
24 experimental techniques to investigate the hydration process of a single ion in aqueous phase
25 and from the necessity to consider theoretical models able to describe both short-range
26 ion/water interactions, water structural perturbation arising from the ion presence and long-
27 range ion/water interactions at an equal level of accuracy. This explains why until now the
28 most accepted thermodynamical properties of the hydrated proton were computed from
29 experimental and theoretical quantum data regarding only small ion/water clusters in gas
30 phase (and whose size n is no larger than 6 water molecules) and by assuming the Cluster
31 Pair Assumption, CPA, *i.e.* the binding Gibbs free energies to add further water molecules
32 to large enough anionic or cationic hydrated clusters become equal.^{6–8}

33 Among long-range ion/water effects, we may quote the effect of water surface on ion
34 hydration. The molecular organization at water boundaries can yield an unbalanced charge
35 distribution and thus a difference in the electrostatic potential $\Delta\Phi$ between the water system
36 core and its chemical environment (usually vacuum). $\Delta\Phi$ within a homogeneous phase is
37 constant. It thus has no effect on ion dynamics within the bulk phase, however, it may
38 favor/disfavor ion hydration by more or less largely shifting the ion hydration Gibbs free
39 energy by a constant value $q\Delta\Phi$ (q being the ion charge). First of all we may note a large
40 dispersion in the reported values of the water surface potential leading to a $q\Delta\Phi$ contribution
41 to the hydration Gibbs free energy of an elementary charge e ranging from -20 up to + 140
42 $k_B T$ at ambient conditions.^{9–13} Moreover only a few studies investigating the behavior of
43 water surface potential in water droplets have been reported, using standard additive water
44 point charge models^{14,15} or a more sophisticated one, the AMOEBA approach, which includes
45 dynamic polarization effects.¹⁶ Among these latter studies, we may quote in particular that

46 reported by Vlcek and co-authors who computed the absolute hydration enthalpies for a
47 large set of monovalent and monoatomic ions dissolved in water droplets whose molecular
48 sizes n were 64, 125, 296 and 1000 and by means of additive point charge force fields.¹⁵
49 Interestingly the latter authors concluded from their simulations that it is far from being
50 obvious to consider small clusters-based methods to estimate the absolute proton hydration
51 Gibbs free energy in solution as reliable and that significant differences may be expected from
52 values computed when considering the thermodynamical properties of large sized ion/water
53 droplets.

54 In the present study, we investigated the reliability of small clusters-based methods to
55 estimate the proton hydration Gibbs free energy in solution by computing the absolute
56 hydration Gibbs free energies of the monovalent ions Li^+ , NH_4^+ , F^- and Cl^- in aqueous
57 phase as well as in isolated and finite size droplets whose molecular size as large as 1 000 by
58 means of molecular dynamics simulations based on polarizable force fields. We also estimated
59 the surface potential in pure water droplets whose molecular size is as large as 10 000. The
60 present manuscript is organized as follows. First we present the polarizable force fields, the
61 numerical protocols used to compute the water surface potential and the absolute single ion
62 hydration Gibbs free energies. Then we discuss the convergence of the water surface potential
63 from droplets to bulk phase and the accuracy of our force fields to compute accurate absolute
64 ion hydration Gibbs free energies from both finite and bulk phase ion/water systems. Lastly
65 we discuss the relation between the difference in anion/cation hydration Gibbs free energies
66 in droplets and the droplet surface potential and the consequence of that relation on the
67 reliability of small clusters-based methods to estimate the absolute proton hydration Gibbs
68 free energy in solution.

1 Methods and computational details

1.1 The force field

We use the rigid water model TCPE/2013¹⁷ and the ion/water force fields detailed in our former studies dealing about the hydration of NH_4^+ ¹⁸ and of halide anions.¹⁹ Regarding the ion Li^+ we consider the *ab-initio* based force field detailed in Supplementary Material (see Section 1). The total potential energy ΔU corresponding to our force fields measures the total intermolecular interaction energy with respect to, *wrt*, individual unbound gas-phase chemical species. ΔU is a sum of five energy components

$$\Delta U = U^{rep} + U^{qq'} + U^{disp} + U^{pol} + U^{mbp}. \quad (1)$$

For a system of N atoms, the repulsive U^{rep} , Coulombic $U^{qq'}$ and dispersion U^{disp} terms are defined as:

$$U^{rep} + U^{qq'} + U^{disp} = \sum_{i=1}^N \sum_{j,j>i}^{N^*} \left[A_{ij} \exp(-B_{ij}r_{ij}) + \frac{q_i q_j}{4\pi\epsilon_0 r_{ij}} - \left(\frac{\sigma_{ij}}{r_{ij}} \right)^6 \right], \quad (2)$$

here, r_{ij} is the distance between atoms i and j , $\{q_i\}$ are the static charges located on the atomic centers, and $(A_{ij}, B_{ij}, \sigma_{ij})$ are adjustable parameters. Only cation/water interactions are modeled using dispersion.¹⁸ The superscript $*$ indicates the sum to include only atom pairs separated by more than two chemical bonds.

Polarization effects are modeled using induced dipole moments \mathbf{p}_i that obey

$$\mathbf{p}_i = \alpha_i \cdot \left(\mathbf{E}_i^q + \sum_{j=1}^{N_\mu^*} \mathbf{T}_{ij} \cdot \mathbf{p}_j \right). \quad (3)$$

All the N_μ non hydrogen atoms are polarizable centers, *i.e.* a single point polarizability is located on each non-hydrogen atomic center. Their isotropic polarizability is α_i . \mathbf{E}_i^q is the static electric field generated on the polarizable center i by the surrounding charges q_j and

\mathbf{T}_{ij} is dipolar tensor (they both include short-range Thole's-like damping functions^{17,20,21}).

The polarization energy is

$$U^{pol} = \frac{1}{2} \sum_{i=1}^{N_\mu} \frac{\mathbf{p}_i^2}{\alpha_i} - \sum_{i=1}^{N_\mu} \mathbf{p}_i \cdot \mathbf{E}_i^q - \frac{1}{2} \sum_{i=1}^{N_\mu} \sum_{j=1}^{N_\mu^*} \mathbf{p}_i \mathbf{T}_{ij} \mathbf{p}_j. \quad (4)$$

The terms U^{mbp} is a short-range anisotropic many-body energy term introduced to improve the description of water/water hydrogen bond (HB) networks and of halide X^- /water strong hydrogen bonds (SHB):

$$U^{mbp} = \sum f(r)g(\Theta). \quad (5)$$

The sum runs over all HB/SHB pairs, r and r_e are the HB/SHB length and its equilibrium value, respectively. Θ is a set of specific intermolecular angles ψ^l like the SHB angle $\angle X^- \cdots H - O$,^{17,19} for instance. The functions f and g are defined as

$$f(r)g(\Theta\{\psi^l\}) = D_e \exp\left(-\frac{(r - r_e)^2}{\gamma_r}\right) \times \prod_l \exp\left(-\frac{(\psi^l - \psi_e^l)^2}{\gamma_\psi^l}\right). \quad (6)$$

Here r_e and $\{\psi_e^l\}$ are the equilibrium values of the geometrical parameters r and $\{\psi^l\}$ as defined in molecular dimers optimized in gas phase using quantum *ab initio* methods. γ_r and γ_ψ^l are adjustable parameters. The intensity of U^{mbp} for a given HB/SHB is modulated by the chemical environment of the water molecule (or the halide) accepting the water hydrogen. This is achieved by taking D_e as a linear function of the local density n_b of the water oxygen (HB) or of the O – H bonds (SHB) in the vicinity of hydrogen acceptor species.^{17,19} No dispersion term is used to model HBs/SHBs. Lastly we also consider an additional three-body term to model the hydration of Cl^- that we recently propose to further improve the description of water/water interactions at the vicinity of an anion²²

$$U^{3b} = \sum A^{3b} \exp\left(-B^{3b} \times r_{O_i O_j}\right) f(r_{Cl O_i}) f(r_{Cl O_j}). \quad (7)$$

Here the sum runs on all the water oxygen pairs (i, j) located at vicinity of anion and the

function f is defined as above for U^{mbp} . A^{3b} and B^{3b} are adjustable parameters that do not depend on the anion nature.

All the above force-field adjustable parameters allowing to model ion/water interactions are assigned to reproduce geometrical and energetic properties of small clusters as computed from high end quantum computations extrapolated to the complete basis set limit (see Refs^{18,19,22} as well as Supplementary Material, Section 1). Note in particular our force fields for halides yield no overpolarization effect on the anion center as compared to the quantum CPMD approach.²¹ Regarding cations, we only consider small clusters where all the water oxygens interact directly with the cationic center.

1.2 Molecular dynamics

Our MD protocols correspond to those used in our former studies.^{18,23–25} Bulk phase systems (comprising all 1 000 water molecules) are simulated in the NPT ensemble using periodic boundary conditions and the Smooth Particle Mesh Ewald, SPME, scheme devoted to polarizable force fields based on an induced dipole moment approach.²⁶ To investigate the surface potential at the liquid water/vacuum interface, we simulate water slabs comprising 2 000 molecules using periodic boundary conditions and the latter SPME scheme, but in the NVT ensemble. Droplet systems (comprising 50, 100, 150, 200, 400, 600, 800 and 1 000 water molecules) are simulated in the NVT ensemble. To prevent evaporation phenomena, droplets are embedded in a large enough cavity and the interactions between the cavity wall and water molecules have no effect of the dynamics on the main droplet region that includes the water/vacuum interface.²⁴ The water and NH_4^+ intramolecular degrees of freedom are constrained during the simulations. All simulations are performed at the 10 ns scale. Their last 9 ns segments are sampled each 1 ps to generate the statistical ensemble from which the simulation averages are computed. More details regarding our MD protocols are provided as Supplementary Material (see Section 4).

1.3 Surface potential

The electrostatic potentials Φ at water-vacuum surfaces are computed by solving the Poisson's equation and by considering summations on atomic charge and dipole contributions (the so-called "P-convention" scheme)

$$\nabla^2\Phi = -\frac{1}{\epsilon_0} \sum_{i=1}^N (q_i + \mathbf{p}_i \cdot \nabla_i) \delta(\mathbf{r} - \mathbf{r}_i), \quad (8)$$

here, $(\mathbf{r}_i, q_i, \mathbf{p}_i)$ are the atomic position, static charge and induced dipole moment of atom i . δ is the volume delta function. As all the sized- n droplets discussed here are quasi-spherical (see our former studies¹⁸ and below) we integrate the above equation in the droplet case as

$$\Phi_d^n(r) = -\frac{1}{\epsilon_0} \int_0^r \frac{C_q^n(r) + C_p^n(r)}{r^2} dr. \quad (9)$$

The distance r is measured from the droplet center of mass (COM). $C_q^n(r)$ is the temporal mean sum of the static charges included in a sphere of radius r and $C_p^n(r)$ is the temporal mean density at distance r of the projection of the induced dipoles in the direction orthogonal to the surface. In the liquid water-vacuum interface case, the above integral is rewritten as

$$\Phi_{\text{bulk}}(z) = -\frac{1}{S\epsilon_0} \int_0^z (C_q(z) + C_p(z)) dz \quad (10)$$

here z is the distance from the simulated slab center in the direction orthogonal to the liquid water-vacuum interface. The functions C_q and C_p are computed by considering a rectangular domain orthogonal to the interface (whose surface S is constant along NVT simulations). Both the above integrals are computed discretely by setting dr and dz to 0.1 Å.

To compute accurate droplet surface potentials $\Phi_d^n(r)$, we performed 50 independent MD simulations (each of 10 ns and corresponding to a different set of initial atomic velocities) of pure water droplets comprising 200, 400, 600 and 800 molecules. The liquid water-vacuum surface potential $\Phi_{\text{bulk}}(z)$ is computed from 10 independent MD simulations (each also of

10 ns) of a periodic slab system comprising 2 000 water molecules. The functions $\bar{C}_{q/p}$ are the mean $C_{q/p}$ ones averaged on the independent MD trajectories and the $\delta C_{q/p}$ s are their corresponding root mean square deviation among the latter trajectories. We assume the $\delta C_{q/p}$ s to be a measure of the $\bar{C}_{q/p}$ uncertainties. They are smaller than 1%, regardless of the water system considered. We also estimated the surface potential of a large water droplet made of 10 000 molecules, however from a single trajectory generated as detailed in Ref.²⁴

Note that we reported an erroneous $\Delta\Phi_{\text{bulk}}$ value for the water model TCPE/2013 in our earlier study.¹⁹ The error arises from an erroneous factor 2 introduced in the computations of the function $C_q(z)$. We check our corrected computational protocol by reproducing the $\Delta\Phi_{\text{bulk}}$ value for the water model TIP3P (about 0.5 V,²⁷ see Supplementary Material, Figure 14).

We also computed the TCPE/2013 dipolar contribution $\Delta\Phi_{\text{bulk}}^{\text{D}}$ to the water surface potential by means of the so-called "M-convention" scheme, *i.e.* by considering the total molecular dipole vector as hosted at the water oxygen to solve the Poisson's equation. We checked the sum of the dipolar contribution $\Delta\Phi_{\text{bulk}}^{\text{D}}$ and of the Bethe's potential contribution $\Delta\Phi_{\text{bulk}}^{\text{Q}}$ arising from the water molecular quadrupole tensor to well reproduce the total water surface potential $\Delta\Phi_{\text{bulk}}$ (see Supplementary Material, Figure 14).

1.4 Droplet and bulk Gibbs free energies

To compute the single ion hydration Gibbs free energy, we consider the thermodynamic cycle **TC** shown in Figure 1. Based on the ion charge sign, we consider two different virtual, neutral and non polarizable atoms **V** whose first hydration shell structure in water is close that of a cation (*i.e.* **V** interacts directly with water oxygens) or of an anion (**V** interacts directly with water hydrogens). We denote these two kinds of virtual atoms as **V_c** and **V_a** when needed. The water/**V** interaction potentials were built to favor their presence within the droplet core along droplet NVT trajectories. The corresponding force-field parameters are provided in Supplementary Material (see Section 7, Table 3) and the propensity of both

virtual atoms for the core of any size water droplets is shown by the centripetal character of their potential of mean force, PMF, as interacting with water droplets computed using a standard Umbrella Sampling scheme (see Supplementary Material, Section 7 and Figures 6 and 7).

According to the thermodynamic cycle **TC**, the hydration Gibbs free energy of an ion **X** at infinite dilution is

$$\mu_{\text{hyd}}^{\infty}(\mathbf{X}) = \mu_{\text{hyd}}^{\infty}(\mathbf{V}) - \Delta G_n + (\mu_{\text{hyd}}(\mathbf{X}, n) - \mu_{\text{hyd}}(\mathbf{V}, n)). \quad (11)$$

Here ΔG_n is the Gibbs free energy cost corresponding to the alchemical reaction $\mathbf{X}_{\text{hyd}} \rightarrow \mathbf{V}_{\text{hyd}}$ in a sized- n water droplet, $\mu_{\text{hyd}}^{\infty}(\mathbf{A})$ and $\mu_{\text{hyd}}(\mathbf{A}, n)$ are the absolute bulk hydration Gibbs free energies of the entity **A** in bulk water and of **A** embedded in sized- n water droplet, respectively.

The quantity $\mu_{\text{hyd}}(\mathbf{A}, n)$ is the sum of three components: the Gibbs free energy cost $\mu_{\text{hyd}}^{\text{cav}}(\mathbf{A}, n)$ to create a cavity within liquid water and corresponding to the sized- n cluster, the electrostatic cost $\mu_{\text{hyd}}^{\text{elec}}(\mathbf{A}, n)$ to embed the cluster in the latter cavity and the repulsion/hydrogen bond contribution $\mu_{\text{hyd}}^{\text{surface}}(\mathbf{A}, n)$ corresponding to water/water local interactions at the cluster/cavity interface. We show that ion and **V** droplet systems are all quasi-spherical and the difference in their volume between ion and **V** systems is negligible for the droplet systems that we consider (see Supplementary Material, Figure 5). The difference in the $\mu_{\text{hyd}}^{\text{cav}}(\mathbf{A}, n)$ and $\mu_{\text{hyd}}^{\text{surface}}(\mathbf{A}, n)$ contributions between ion and **V** droplets is thus assumed to vanish as $n \rightarrow \infty$ and Equation 11 may be rewritten for large enough droplets as

$$\mu_{\text{hyd}}^{\infty}(\mathbf{X}) = \mu_{\text{hyd}}^{\infty}(\mathbf{V}) - \Delta G_n + (\mu_{\text{hyd}}^{\text{elec}}(\mathbf{X}, n) - \mu_{\text{hyd}}^{\text{elec}}(\mathbf{V}, n)). \quad (12)$$

Considering the charge distribution corresponding to an entity **A** embedded in a spherical

cavity (of radius a_n) within a continuum medium, the component $\mu_{\text{hyd}}^{\text{elec}}(\mathbf{A}, n)$ should obey²⁸

$$\mu_{\text{hyd}}^{\text{elec}}(\mathbf{A}, n) = \frac{1}{8\pi\epsilon_0} \sum_{l=0}^{\infty} \frac{(l+1)(1-\epsilon)}{(l+1)\epsilon + l} \left(\frac{M_l}{a_n^{2l+1}} \right), \quad (13)$$

here M_l is the electric moment of order l of the charge distribution. That relation is still valid when considering a distribution of induced dipole moments as they can be modeled by a specific set of charges. However the above relation implicitly corresponds to an abrupt transition in the dielectric constant between the solute cavity and the solvent. For an ion solvated in aqueous phase, the water local dielectric permittivity $\epsilon(\mathbf{r})$ behavior is shown to be strongly oscillatory until a distance of 12 Å from the ion.²⁹ From the work of Beveridge and Schnuelle,²⁸ that yields the $\mu_{\text{hyd}}^{\text{elec}}(\mathbf{A}, n)$ expansion for an ion to also include an a_n^{-2} term.

We showed the molecular density within pure water droplets to be converged to the bulk value within less than 2 %.¹⁸ The radius a_n of a sized- n droplet may be thus assumed to be proportional to $n^{-1/3}$ and from all the above, the gas-phase cluster ΔG_n may be written as a power law function of the droplet size n

$$\Delta G_n = \Delta G_{\infty} + \frac{\gamma_1}{n^{1/3}} + \frac{\gamma_2}{n^{2/3}} + O\left(\frac{1}{n}\right), \quad (14)$$

and according to Equation 11 the quantity ΔG_{∞} is

$$\Delta G_{\infty} = \lim_{n \rightarrow \infty} \Delta G_n = \mu_{\text{hyd}}^{\infty}(\mathbf{V}) - \mu_{\text{hyd}}^{\infty}(\mathbf{X}). \quad (15)$$

1.5 Computing Gibbs free energies

The bulk hydration Gibbs free energies $\mu_{\text{hyd}}^{\infty}(\mathbf{A})$ are estimated from MD simulations in bulk phase using periodic boundary conditions and the Thermodynamic Integration (TI) scheme by progressively decoupling solute/solvent interactions. For ionic species, we compute their $\mu_{\text{hyd}}^{\infty}(\mathbf{X})$ in two steps, first by decoupling the electrostatic and polarization ion \mathbf{X} /water interactions and then by decoupling the remaining interactions between the uncharged and

214 non-polarizable entity \mathbf{X}^0 and water (*i.e.* interactions corresponding to the energy terms
215 U^{rep} , U^{disp} and U^{slh}).

216 We systematically consider a 20 windows TI scheme. Each window corresponds to a 10
217 ns MD simulation where the solute \mathbf{A} /water solvent hamiltonian is scaled by a constant λ
218 regularly spaced between 1 and 0. Each 50 MD steps, we compute the derivative of the system
219 total potential energy ΔU wrt the scaling parameter λ using a finite difference method. From
220 the mean derivative values and their corresponding root mean square deviations, we compute
221 both the integrals providing $\mu_{\text{hyd}}^{\infty}(\mathbf{A})$ and their uncertainties assuming that the derivatives
222 computed along each MD windows are temporally uncorrelated. Because of the large size
223 of the derivative data sets, the latter uncertainties are negligible, at most about ± 0.05 kcal
224 mol^{-1} . Regarding the virtual atoms \mathbf{V} , their quantities $\mu_{\text{hyd}}^{\infty}(\mathbf{V})$ computed as above are -12.5
225 (\mathbf{V}_c) and -9.0 (\mathbf{V}_a) ± 0.05 kcal mol^{-1} . Note here that these values agree with those that
226 may be estimated from the virtual atom/water droplet PMFs provided as Supplementary
227 Material (see Figures 6 and 7).

228 To remove periodic artifacts arising from accounting for ion images in our simulation
229 protocol, we subtract the self-contribution $\delta\mu_{\text{self}}^{\text{bulk}}$ from our bulk hydration Gibbs free energies
230 $\mu_{\text{hyd}}^{\infty}(\mathbf{X})$. As the solute static charges are scaled by the parameter λ in our implementation
231 of the TI scheme, $\delta\mu_{\text{self}}^{\text{bulk}}$ obeys

$$\delta\mu_{\text{self}}^{\text{bulk}} = \frac{\xi q^2}{8\pi\epsilon_0\epsilon\bar{L}}. \quad (16)$$

232 ϵ is the liquid water dielectric constant predicted by the water model TCPE/2013 at 300
233 K (78.3¹⁷) and \bar{L} is the average simulation cell dimension along the NPT bulk trajectories,
234 about 31 Å, and $\xi = 2.837297$. That yields $\delta\mu_{\text{self}}^{\text{bulk}} = 0.2$ kcal mol^{-1} . A priori we should also
235 account for other periodic effects arising from the inappropriate orientational polarization of
236 the solvent during the simulation, for instance. These corrections, corresponding to higher
237 order terms of \bar{L}^{-1} , are here fully negligible because of the size of our simulation boxes.³⁰

238 Regarding clusters, we compute their quantity ΔG_n from NVT droplet simulations and
239 by means of the same TI scheme detailed above for bulk systems. We tested two approaches:

they correspond (1) to alchemically transform the charged ion \mathbf{X} in its corresponding virtual atom \mathbf{V} in one step, and (2) to uncouple first the electrostatic and polarization ion/water interactions and then to alchemically transform the uncharged and non-polarizable entity \mathbf{X}^0 in its corresponding virtual atom \mathbf{V} . Both approaches provide equal results regarding the hydration Gibbs free energy in droplet, within less than $0.1 \text{ kcal mol}^{-1}$.

1.6 Further corrections to simulation-based estimate of ion hydration Gibbs free energy

To estimate reliable single ion bulk hydration Gibbs free energy from our simulation data, two physical contributions have also to be considered. The first, δG_{ref} , arises from the identical ion concentration in gas phase and in liquid water in our bulk phase simulation protocol, while experimentally the ion concentration in gas phase correspond to that of 1 mole of an ideal gas and to 1 M in liquid water. We thus estimate δG_{ref} as the amount of energy to compress 1 mole of an ideal gas to reach a concentration of 1 mol l^{-1} . At ambient conditions, $\delta G_{\text{ref}} = RT \ln 22.4 = +1.89 \text{ kcal mol}^{-1}$.

The second, $\delta G_{\text{vib}}^{g \rightarrow l}$, arises from the use of constraints in our MD simulations, *i.e.* intramolecular degrees of freedom of the non-monoatomic species NH_4^+ and H_2O are frozen along our MD trajectories. However, quantum computations show large shifts in the intramolecular harmonic vibrational frequencies of NH_4^+ between its gas-phase isolated state and hydrated clusters. For the four hydrated NH_4^+ cluster, these shifts are on average of -230 cm^{-1} for the cation harmonic stretching frequencies and of $+60 \text{ cm}^{-1}$ for the bending ones.³¹

Liquid water intramolecular vibrational frequencies may be also altered when water molecules interact at short range from an ion. For instance the mean stretching frequency ν_{OD} in liquid HDO is experimentally reported to be shifted by about -70 and $+60 \pm 10 \text{ cm}^{-1}$ compared to bulk as HDO molecules lie in the first hydration shell of Li^+ and F^- ,³² respectively, in agreement for Li^+ with a quantum-based theoretical estimate.³³ For both the anion

Cl⁻ and the cation K⁺, the latter experimental shift is small about $+20 \pm 10 \text{ cm}^{-1}$ (cf. the data discussed in Ref.³²). Regarding Cl⁻, we may quote also an *ab initio* molecular dynamics study³⁴ showing the infrared stretching vibration spectrum of water molecules lying at its vicinity to be more dissymmetric than in the bulk (with a slightly larger weight of weak ν_{OH} frequencies). In all that suggests a very weak effect of the Cl⁻/K⁺ presence on the liquid water intramolecular vibrational properties. As the hydration properties of NH₄⁺ are close to K⁺, we assume as negligible the enthalpic $\delta H_{\text{vib}}^{g \rightarrow l}$ and Gibbs free energy $\delta G_{\text{vib}}^{g \rightarrow l}$ costs arising from the perturbation of the liquid water intramolecular vibrational spectrum due to the Cl⁻/NH₄⁺ presence.

We estimate the magnitude of $\delta G_{\text{vib}}^{g \rightarrow l}$ (and of its enthalpic $\delta H_{\text{vib}}^{g \rightarrow l}$ counterparts) arising from the solute or solvent intramolecular frequency shifts from standard polyatomic ideal gas formula (see Supplementary Material, Section 5). The enthalpy cost $\delta H_{\text{vib}}^{g \rightarrow l}$ is estimated to be -0.8 and -1.6 kcal mol⁻¹ for Li⁺ and NH₄⁺, respectively. The magnitude of the Gibbs free energy cost $\delta G_{\text{vib}}^{g \rightarrow l}$ is slightly weaker, about -0.5 and -1.0 kcal mol⁻¹ for the latter two ions, respectively. The order of magnitude of these contributions supports the reliability of simulating ion hydration by constraining intramolecular degrees of freedom.

Hence adding the latter two corrections to $\mu_{\text{hyd}}^{\infty}(\mathbf{X})$ yields the absolute single ion bulk hydration Gibbs free energy value $\tilde{\mu}_{\text{hyd}}^{\infty}(\mathbf{X})$

$$\tilde{\mu}_{\text{hyd}}^{\infty}(\mathbf{X}) = \mu_{\text{hyd}}^{\infty}(\mathbf{X}) + \delta G_{\text{vib}}^{g \rightarrow l} + \delta G_{\text{ref}}. \quad (17)$$

2 Results and discussion

2.1 Convergence of the water droplet surface potential to the bulk limit

The plots of the n -sized droplet surface potential profiles $\Phi_d^n(r)$ are provided as Supplementary Material (see Figures 10 to 13). Agreeing with earlier results,¹¹ their main features are

similar to the bulk-vacuum profile $\Phi_{\text{bulk}}(z)$ shown in Figure 2(a): $\Phi_d^n(r)$ are constant outside of the droplets and within their cores, with a rapid transition at the droplet surfaces. We denote hereafter $\Delta\Phi_d^n/\Delta\Phi_{\text{bulk}}$ the difference in the $\Phi_d^n(r)/\Phi_{\text{bulk}}(z)$ values between vacuum and the aqueous core. As shown in Figure 2(b), $\Delta\Phi_d^n$ obeys a decreasing linear function of $n^{-1/3}$

$$\Delta\Phi_d^n = \Delta\Phi_{\text{bulk}} - \frac{\phi}{n^{1/3}}. \quad (18)$$

It converges towards the bulk limit $\Delta\Phi_d^\infty = -0.227$ V that matches our computed $\Delta\Phi_{\text{bulk}}$ value within less than 0.1 % (the slope $-\phi$ is here -0.331 V). Hence all the droplet values $\Delta\Phi_d^n$ are larger in magnitude than the bulk one, by 20 % for the 200 sized droplet, for instance.

The linear dependence on $n^{-1/3}$ of our droplet $\Delta\Phi_d^n$ data agrees with earlier results reported by Kastenholtz and Hünnerberger¹⁴ and by Vlcek and co-authors¹⁵ from simulations performed using the point charge water models SPC and SPC/E, respectively. However we may quote that Pollard and Beck¹⁶ showed the $\Delta\Phi_d^n$ values estimated from ion hydration enthalpy data in small droplets ($n < 200$) and the AMOEBA force field to exhibit also a decreasing but non $n^{-1/3}$ linear behavior.

The bulk surface potential value $\Delta\Phi_{\text{bulk}}$ predicted by the water model TCPE/2013 is less than half the value reported for most of the available water models¹¹ based on point charges (point dipoles) approaches, but twice as large as the point-charge water model TIP5P.³⁵ However the TCPE/2013 dipolar $\Delta\Phi_{\text{bulk}}^D$ contribution is clearly larger than that corresponding to available water models including TIP5P and closer to the quantum DFT estimates,³⁵ +0.38 and +0.48 V, respectively. From the discussions reported by Remsing *et al* (in particular regarding their data summarized in the Table 2 in Ref.³⁵) the latter result suggests TCPE/2013 to predict a molecular organization at the water surface close to that predicted by quantum DFT-based simulations. On the other hand and as all point charge-based models, TCPE/2013 yields an overall large but negative Bethe's potential contribution $\Delta\Phi_{\text{bulk}}^Q = -0.61$ V, whereas quantum DFT simulations predict a very large and positive contribution, about +4 V.^{35,36}

315 2.2 Mean stepwise water binding free energy in large ion/water 316 droplets

317 We define the mean stepwise water binding free energy $\delta\tilde{\mu}_{m \rightarrow n}$ as the free energy cost to
318 further add a water molecule to ion \mathbf{X} /water droplets whose size is included between m and
319 n :

$$\delta\tilde{\mu}_{m \rightarrow n} = \frac{\mu_{\text{hyd}}(\mathbf{X}, n) - \mu_{\text{hyd}}(\mathbf{X}, m)}{n - m}. \quad (19)$$

320 As shown by the plots of Figure 3, that quantity is about equal for our four ions no sooner
321 than $m = 400$. The specific nature of a monovalent ion has thus still a noticeable effect on
322 ion/water interactions in aqueous droplets until a distance from the ion of about $R_{\delta\mu} = 14 \text{ \AA}$,
323 a distance that matches the R_{ϵ} one for which the oscillatory behavior of the water permittivity
324 induced by ion presence vanishes.²⁹ As the two distances $R_{\delta\mu}$ and R_{ϵ} have been estimated
325 using two different kinds of force field (in particular, a classical and a polarizable one), their
326 agreement shows the influence of a specific ion on water to extend far beyond the usual cut
327 off distance used by protocols based on the Quasi Chemical Theory (about 6 \AA ³⁷) to define
328 the "inner-shell" term that is shown to produce alone accurate estimates of the full bulk
329 ion hydration Gibbs free energy. However we may quote an experimental study that shows
330 monovalent ions (namely I^- and Na^+) dissolved in $n \approx 250$ water droplets to not affect the
331 water hydrogen bond network behind the ion second hydration shell.³⁸

332 2.3 Single ion hydration Gibbs free energies : from droplets to 333 bulk

334 As our force-field total potential energy ΔU is a sum of five components, we may decompose
335 ΔG_n into five contributions

$$\Delta G_n = \Delta G_n^{\text{rep}} + \Delta G_n^{\text{disp}} + \Delta G_n^{\text{mbp}} + \Delta G_n^{\text{qq}'} + \Delta G_n^{\text{pol}}. \quad (20)$$

336 The first three contributions correspond to interaction energy terms that are short ranged. As
337 shown by our TI computations, they converge more rapidly to their bulk limit value than the
338 last two contributions. The free energy components corresponding to the alchemical reaction
339 $[\text{NH}_4^0(\text{Li}^0)/(\text{H}_2\text{O})_n] \rightarrow [\mathbf{V}_c, (\text{H}_2\text{O})_n]$ or to the sum of components $\Delta G_n^{\text{rep}} + \Delta G_n^{\text{disp}} + \Delta G_n^{\text{mbp}}$ for
340 the reaction $[\text{Cl}^-, (\text{H}_2\text{O})_n] \rightarrow [\mathbf{V}_a, (\text{H}_2\text{O})_n]$ are already converged within $0.25 \text{ kcal mol}^{-1}$ on
341 average as soon as $n = 400$ and within $0.1 \text{ kcal mol}^{-1}$ at $n = 1000$, regardless of the ion. We
342 denote both the latter quantities as ΔG_n^{sr} and their converged $\Delta G_\infty^{\text{sr}}$ values are summarized
343 in Table 1.

344 We thus extrapolated only the sum of the non-zero electrostatic and polarization contri-
345 butions $\Delta G_n^{\text{elec}} = \Delta G_n^{\text{qq}'} + \Delta G_n^{\text{pol}}$ using the power law function shown in Equation (14) for
346 our four ions. The ΔG_n^{elec} quantities are plotted as functions of $n^{-1/3}$ in Figure 4 and the
347 extrapolated $\Delta G_\infty^{\text{elec}}$ values are summarized in Table 1. The uncertainty affecting the $\Delta G_\infty^{\text{elec}}$
348 values arising from the fitting process is $\leq 0.3 \text{ kcal mol}^{-1}$. Accounting also for the uncer-
349 tainties regarding the energy derivative averages computed along the TI MD simulations and
350 the $\Delta G_\infty^{\text{sr}}$ values yields a total error for the values $\Delta G_\infty = \Delta G_\infty^{\text{sr}} + \Delta G_\infty^{\text{elec}}$ of about $\epsilon_{\text{err}} \approx 0.6$
351 kcal mol^{-1} . This supports the reliability our droplet-based protocol to compute single ion
352 hydration thermodynamic properties.

353 In Table 1 we summarize the bulk limit values ΔG_∞ extrapolated from droplet data, the
354 single ion bulk hydration Gibbs free energy $\mu_{\text{hyd}}^\infty(\mathbf{X})$ values computed from bulk simulations,
355 the differences $\mu_{\text{hyd}}^\infty(\mathbf{X}) - \mu_{\text{hyd}}^\infty(\mathbf{V})$ in the bulk hydration Gibbs free energy between ions
356 and their corresponding virtual atoms. As expected from Equation (12), the difference
357 in the quantities ΔG_∞ and $[\mu_{\text{hyd}}^\infty(\mathbf{V}) - \mu_{\text{hyd}}^\infty(\mathbf{X})]$ ($5.2 \pm 0.5 \text{ kcal mol}^{-1}$ on average and in
358 absolute values) for all the ions equals the surface potential contribution $q\Delta\Phi_{\text{bulk}}$ within our
359 computational protocol uncertainty.

360 We also estimated single ion bulk hydration Gibbs free energies from droplet data using
361 different extrapolation schemes than above. For instance, we extrapolated the total Gibbs

362 free energy $\Delta G_n = \Delta G_n^{\text{elec}} + \Delta G_n^{\text{sr}}$ using the power law function

$$\Delta G_n = \Delta G_\infty^\circ + \frac{\gamma_1^\circ}{n^{1/3}} + \frac{\gamma_2^\circ}{n^{2/3}} + \frac{\gamma_3^\circ}{n}. \quad (21)$$

363 The last term of the right hand equation is introduced to account for short range non elec-
364 trostatic ion/water interactions. It is a priori well suited to account for ion/water dispersion
365 as modeled in our force-field for cations Li^+ and NH_4^+ . The adjusted parameters ΔG_∞° and
366 $\gamma_{1,2,3}^\circ$ are provided as Supplementary Material (see Table 4). The new single ion bulk hy-
367 dration Gibbs free energy estimates $\Delta G_n^{\infty,\circ}$ differ from those computed based on the Table
368 1 data by about $\pm 1 \text{ kcal mol}^{-1}$, a difference in line with the uncertainty ϵ_{err} of computed
369 Gibbs free energy data from our TI simulations.

370 As discussed in Section 2.3, the nature of a monovalent ion has an effect on water in
371 aqueous droplets up to a distance of about 14 Å from the ion center. Such a distance
372 corresponds to the radius of a $n = 400$ -sized water droplet, a droplet size for which all
373 the non electrostatic contributions to the single ion hydration Gibbs free energy in droplets
374 are converged to their bulk limit, like ΔG_n^{sr} and $\mu_{\text{hyd}}(\mathbf{V}, n)$. Hence for $n \geq 400$, we may
375 approximate a water droplet to a continuous medium. Assuming the dielectric constant of
376 water droplets corresponding to $n \geq 400$ as already large enough (> 10 , see Ref.³⁹) and from
377 the Born equation, the electrostatic free energy contribution $\mu_{\text{hyd}}^{\text{elec}}(\mathbf{X}, n)$ and thus the free
378 energy ΔG_n should then obey a linear $n^{-1/3}$ regime

$$\Delta G_n = \Delta \tilde{G}_\infty + \frac{\tilde{\gamma}}{n^{1/3}} \quad (22)$$

379 The corresponding adjusted parameters are listed in Table 2. Note first the linear regression
380 coefficients to be all > 0.99 and even > 0.999 for Li^+ , F^- and Cl^- supporting the assumption
381 regarding the linear $n^{-1/3}$ regime to which the ΔG_n values (and thus the single ion hydration
382 Gibbs free energies in droplets μ_{hyd}^n) should obey in large droplets. Moreover and as their
383 $\Delta G_n^{\infty,\circ}$ counterparts, the $\Delta \tilde{G}_\infty$ values agree with those that may be computed from the data

384 listed in Table 1 also within ± 1 kcal mol⁻¹ on average.

385 2.4 Single ion and proton bulk hydration Gibbs free energy

386 In Table 3 we summarize and compare to data available in literature the absolute single ion
387 hydration Gibbs free energies $\tilde{\mu}_{\text{hyd}}^{\infty}$ computed from the droplet data of Table 1. Our anion
388 $\tilde{\mu}_{\text{hyd}}^{\infty}$ values agree reasonably well with the data of Tissandier *et al*⁷ and differ slightly more
389 from the data of Kelly *et al*.⁸ Regarding cations our $\tilde{\mu}_{\text{hyd}}^{\infty}$ values are clearly underestimated
390 by about 10 kcal mol⁻¹ compared to both the latter sets of data. That difference in the
391 $\tilde{\mu}_{\text{hyd}}^{\infty}$ result quality between anions and cations yields also a clear difference in the absolute
392 proton hydration Gibbs free energy $\mu_{\text{hyd}}^{\infty}(\text{H}^+)$ when estimated for both kind of ions. If we
393 consider the Tissandier *et al*⁷ set of *conventional* ion free energies $\mu_{\text{hyd}}^{\infty, \text{con}}$ that differ at most
394 by 0.1 kcal mol⁻¹ from the set of Fawcett *et al*⁴⁰ and that are tied to the ion and proton
395 $\mu_{\text{hyd}}^{\infty}$ values according to

$$\mu_{\text{hyd}}^{\infty, \text{con}}(\mathbf{X}^{ze}) = \tilde{\mu}_{\text{hyd}}^{\infty}(\mathbf{X}^{ze}) - z\mu_{\text{hyd}}^{\infty}(\text{H}^+). \quad (23)$$

396 we get an anion-based estimate $\mu_{\text{hyd}}^{\infty}(\text{H}^+)$ of 269.2 ± 0.1 kcal mol⁻¹ that is overestimated by
397 a few kcal mol⁻¹ compared to the Tissandier's⁷ and Kelly's⁸ values, 263.9 and 265.9 kcal
398 mol⁻¹, respectively, whereas our cation-based estimate is 255.3 ± 0.1 kcal mol⁻¹, a value
399 that deviates more largely from the Tissandier's and Kelly's estimates, also by about 10 kcal
400 mol⁻¹.

401 We may note first that both our cation and anion-based proton values overall agree with
402 the results reported by earlier authors that range from 249.5 to 264 kcal mol⁻¹.^{11,15,41-43} We
403 may also note that the Tissandier's and Kelly's $\mu_{\text{hyd}}^{\infty}(\text{H}^+)$ values are derived by averaging
404 data corresponding to hundreds of cation/anion pairs and that their $\mu_{\text{hyd}}^{\infty}(\text{H}^+)$ uncertainty
405 is no less than 2 kcal mol⁻¹.^{7,8} It is thus not obvious to draw definitive conclusions from the
406 data of our small ion set to assess the reliability of our computational approach. However,

we identified a cation/water force field artifact arising from an inaccurate description of the water pair interactions among molecules belonging to the cation first and second hydration shells : the interaction energy of such water pairs is underestimated by about 1 kcal mol⁻¹ by our force-field (see Supplementary Material, Section 2). As there are about 10 first/second hydration shell water pairs in bulk phase at the vicinity of both Li⁺ and NH₄⁺, improving the description of these kind of water pair interactions should yield a priori a better agreement in the proton hydration Gibbs free energies computed from our cation and anion simulated data. Note the force fields we use for cations are built to reproduce precisely the interaction energies of cations interacting only with water molecules in their first hydration sphere and no larger cluster.

We made an attempt to remediate that cation/water force field artifact using a correction scheme similar to the one we recently proposed for anions as using our polarizable force fields. Here we reinforce the water/water interactions for molecules belonging respectively to the first and the second hydration sphere of Li⁺ using a three body energy function whose analytical form is close to the hydrogen bonded energy term U^{lh} of the water model TCPE/2013 (see Supplementary Material, Section 3). Using such a function we computed again the absolute single ion hydration Gibbs free energy $\tilde{\mu}_{\text{hyd}}^{\infty}(\text{Li}^+)$ from bulk simulations. It is then shifted to 123.4 kcal mol⁻¹, in better agreement with the Tissandier's and Kelly's data. Even if the validity of such a correction scheme has to be assessed on a wider set of monovalent cations, the latter result supports the above assumption regarding the origin of the above disagreement between our computed and well accepted estimates regarding cation $\tilde{\mu}_{\text{hyd}}^{\infty}(\mathbf{X})$ data. We may thus consider that the cation force field artifact has no incidence on cation/water long range interactions and thus on the conclusions drawn above regarding the quantities $\delta\tilde{\mu}_{m \rightarrow n}$ for instance.

A priori the above cation force field artifact is not tied to the inability of our force fields to accurately model cation/water interactions but it arises mainly from a water model TCPE/2013 artifact that has no effect on modeling the properties of pure water systems.

Hence the accurate modeling of ion hydration thermodynamic properties doesn't depend only on the quality of the description of ion/water short and medium range interactions⁴⁴ but also on the quality of the description of the water/water interactions at the vicinity of ionic centers.

2.5 The net water interface potential

A series of single ion hydration Gibbs free energy values have been reported^{42,45-47} that measure the ion bulk hydration Gibbs free energy contribution disentangled from surface effects at both the aqueous-gas phase interface and at the boundary of the local cavity hosting the ion within water. These *no surface* contributions $\mu_{\text{hyd}}^{\infty,ns}$ are tied to absolute ion hydration Gibbs free energies according to

$$\tilde{\mu}_{\text{hyd}}^{\infty}(\mathbf{X}) = \mu_{\text{hyd}}^{\infty,ns}(\mathbf{X}) + q_{\mathbf{X}}\Phi_{np}, \quad (24)$$

here $q_{\mathbf{X}}$ is the charge of the ion \mathbf{X} . Φ_{np} is the net interface potential, the sum of a local potential Φ_{lp} and of the water surface potential $\Delta\Phi_{bulk}$ defined in Section 1.3. Let us consider an aqueous medium (bulk phase or a large enough finite size droplet) as a continuum medium that ignores the electrostatic charge dispersion within a water molecule. The local potential Φ_{lp} may be then interpreted as the electrostatic potential shift when crossing the boundary of the small cavity hosting an ion within the aqueous medium, not accounted for by the latter water continuum representation but arising from the water molecular charge dispersion. A priori Φ_{lp} depends on the ion nature.

As Pollard and Beck,¹⁶ we estimated a mean net potential $\tilde{\Phi}_{np}$ from the *no surface* data of Marcus^{45,46} and from the differences in our own absolute single ion bulk hydration Gibbs free energies $\tilde{\mu}_{\text{hyd}}^{\infty}$ for cation/anion (\mathbf{C},\mathbf{A}) pairs

$$\tilde{\Phi}_{np} = \frac{(\tilde{\mu}_{\text{hyd}}^{\infty}(\mathbf{C}) - \tilde{\mu}_{\text{hyd}}^{\infty}(\mathbf{A})) - (\mu_{\text{hyd}}^{\infty,ns}(\mathbf{C}) - \mu_{\text{hyd}}^{\infty,ns}(\mathbf{A}))}{2e}. \quad (25)$$

We get $\tilde{\Phi}_{np}$ values included within -0.30 (Li^+/Cl^-) and -0.41 ($\text{NH}_4^+, \text{F}^-$) V. If we consider the corrected value $\tilde{\mu}_{\text{hyd}}^\infty$ for Li^+ , 123.4 kcal mol⁻¹, from our attempt to remediate the cation/water force field artifact, we get then $\tilde{\Phi}_{np} = -0.43$ (Li^+/Cl^-) and -0.50 (Li^+, F^-) V. All these values agree with earlier estimates ranging from -0.40 to -0.50 V (see discussions and the references cited in^{16,37}) but clearly differs from the recent quantum estimates predicting a positive value for water net potential, ranging from 0.05 to 0.25 V.^{35,36}

2.6 The Cluster Pair Approximation for large ion/water droplets and the proton bulk hydration Gibbs free energy

The Cluster Pair Approximation, CPA, is the assumption according to which adding a new water molecule to a large enough ion/water cluster leads to a change in the ion hydration Gibbs free energy that does not depend on the ion nature. Let us consider again an ion immersed in an aqueous medium large enough so that it can be modeled as a continuum medium. The Gibbs free energy increment to add new water molecules to that system is proportional to the square of the ion charge q in line with CPA. That suggests the difference $\Delta_{\text{AC}}\mu_{\text{hyd}}^n$ in the hydration Gibbs free energies between a cation and an anion to be constant as soon as these ions are solvated in large enough water droplets.

However modeling water as a continuum medium implicitly neglects the energy cost for a microscopic charge to cross the water surface and the cavity boundary in which the ion is hosted and arising from non zero potentials $\Delta\Phi_{\text{bulk}}$ and Φ_{lp} . Considering *no surface* hydration Gibbs free energies as defined in the above section and the latter two potentials, the difference in the bulk hydration Gibbs free energy between a monovalent anion **A** and cation **C** is

$$\Delta_{\text{AC}}\mu_{\text{hyd}}^\infty = \Delta_{\text{AC}}\mu_{\text{hyd}}^{\infty,ns} - e(\Phi_{lp}^\infty(\mathbf{A}) - \Phi_{lp}^\infty(\mathbf{C})) - 2e \cdot \Delta\Phi_{\text{bulk}}, \quad (26)$$

here $\Phi_{lp}^\infty(\mathbf{X})$ is the local potential shift from the bulk water core to the center of the cavity

hosting ion \mathbf{X} . In the droplet case that relation can be rewritten as

$$\Delta_{AC}\mu_{\text{hyd}}^n = \Delta_{AC}\mu_{\text{hyd}}^{n,ns} - e(\Phi_{lp}^n(\mathbf{A}) - \Phi_{lp}^n(\mathbf{C})) - 2e \cdot \Delta\Phi_d^n, \quad (27)$$

here $\Phi_{lp}^n(\mathbf{X})$ in the local potential for droplets. For large enough ion/water droplets, the CPA assumption yields $\Delta_{AC}\mu_{\text{hyd}}^{\infty,ns} = \Delta_{AC}\mu_{\text{hyd}}^{n,ns}$ and combining the above two relations yields

$$\Delta_{AC}\mu_{\text{hyd}}^{\infty} = \Delta_{AC}\mu_{\text{hyd}}^n + e \sum_{\mathbf{X}=\mathbf{A},\mathbf{C}} z_{\mathbf{X}}(\Phi_{lp}^{\infty}(\mathbf{X}) - \Phi_{lp}^n(\mathbf{X})) - 2e(\Delta\Phi_{\text{bulk}} - \Delta\Phi_d^n). \quad (28)$$

Here the charge of the monovalent ion \mathbf{X} is $q = z_{\mathbf{X}}e$. That relation may be considered as a generalization of the formula to compute the CPA-based estimate of the water surface potential proposed by Vlcek and co-authors.¹⁵ For large enough droplets, we expect $\Phi_{lp}^{\infty}(\mathbf{X}) \approx \Phi_{lp}^n(\mathbf{X})$ regardless of ion \mathbf{X} yielding the fundamental relation

$$\Delta_{AC}\mu_{\text{hyd}}^{\infty} = \Delta_{AC}\mu_{\text{hyd}}^n - 2e(\Delta\Phi_{\text{bulk}} - \Delta\Phi_d^n). \quad (29)$$

From the plots of the electrostatic potentials at the vicinity of our virtual atoms $\mathbf{V}_{\mathbf{a}}$ and $\mathbf{V}_{\mathbf{c}}$ in droplets (see Supplementary Material, Section 9 and Figures 8 and 9), relation (29) may be considered as exact as soon as $n = 400$. Moreover according to Equation (18) the water model TCPE/2013 yields

$$2e(\Delta\Phi_{\text{bulk}} - \Delta\Phi_d^n) = \frac{2e\phi}{n^{1/3}}, \quad (30)$$

with $2e\phi = 15.3 \text{ kcal mol}^{-1}$, a large value suggesting the difference $|\Delta_{AC}\mu_{\text{hyd}}^{\infty} - \Delta_{AC}\mu_{\text{hyd}}^n|$ to be smaller than $0.1 \text{ kcal mol}^{-1}$ no sooner than $n = 3.7 \cdot 10^6$. That corresponds to a droplet whose radius is $\approx 300 \text{ \AA}$. This is also supported by the data reported by Kastenholtz and Hünenberger by means of the point charge water model SPC.¹⁴ The latter authors showed a quasi linear $n^{-1/3}$ behavior for the droplet surface potential corresponding to a $2e\phi$ value of about 20 kcal mol^{-1} . We may also quote Vlcek and co-authors¹⁵ who showed monovalent ion hydration enthalpies to decrease slowly with the droplet size and to be roughly proportional

496 to $n^{-1/3}$.

497 To check the validity of Equation (29) we may first consider the parameters $\tilde{\gamma}$ allowing
498 to estimate the absolute ion hydration Gibbs free energies in $n > 400$ droplets according to

$$\mu_{\text{hyd}}^{\infty}(\mathbf{X}) = \mu_{\text{hyd}}^n(\mathbf{X}) - \frac{\tilde{\gamma}_{\mathbf{X}}}{n^{1/3}}. \quad (31)$$

499 From the values summarized in Table 2 we note the anion parameters $\tilde{\gamma}$ to be larger than the
500 cation ones from 7 to 10 kcal mol⁻¹. However the uncertainty affecting these parameters is
501 large, about ± 2.9 kcal mol⁻¹ (see Supplementary Material, Section 8). The mean difference
502 in anion/cation $\tilde{\gamma}$ values is 8.4 ± 5.8 kcal mol⁻¹, a value a priori in line with our $2e\phi$
503 estimate, 15.3 kcal mol⁻¹, within the error bar. Moreover we also plot in Figure 5 the
504 quantity $\Delta\Delta_{\text{AC}}^{n,\infty} = \Delta_{\text{AC}}\mu_{\text{hyd}}^{\infty} - \Delta_{\text{AC}}\mu_{\text{hyd}}^n$ for all the anion/cation pairs that may be drawn
505 from our ion set and by considering the 4th order power law relation (21). All the quantities
506 $\Delta\Delta_{\text{AC}}^{n,\infty}$ are decreasing functions of $n^{-1/3}$ and they are upper bounded by $2e\phi \cdot n^{-1/3}$. That
507 also supports the fundamental relation (30).

508 We plot in Figure 6 our $\Delta_{\text{AC}}\mu_{\text{hyd}}^n$ values for droplets whose size n ranges from 50 to
509 1000 together with those corresponding to small ion/water clusters ($n \leq 6$) considered by
510 Tissandier's and co-authors and Kelly's and co-authors in their original papers^{7,8} for all the
511 pairs that may be built from our ions. For large droplets, we also reported on that Figure
512 $\Delta_{\text{AC}}\mu_{\text{hyd}}^n$ values for Li⁺ pairs shifted by $\delta G_{\text{C}} = 5.8$ kcal mol⁻¹, the correction value arising
513 from our attempt to remediate the cation force field artifact discussed in Section 2.4. The
514 plots clearly show the quantity $\Delta_{\text{AC}}\mu_{\text{hyd}}^n$ to obey two different regimes for all pairs : that
515 quantity increases in absolute value up to $n \approx 15$ and then it linearly decreases from $n \approx 40$
516 to $+\infty$. That linear behavior is very close for all the ion pairs. Lastly a transition region
517 exists for $15 \leq n \leq 40$, where the quantity $\Delta_{\text{AC}}\mu_{\text{hyd}}^n$ should present a maximum at about
518 $n = 25$ for all our ion pairs.

519 The proton absolute hydration Gibbs free energy $\mu_{\text{hyd}}^{\infty}(\text{H}^+)$ can be computed from the

differences in both anion/cation absolute $\Delta_{AC}\mu_{\text{hyd}}^{\infty}$ and *conventional* $\Delta_{AC}\mu_{\text{hyd}}^{\text{con},\infty}$ hydration
Gibbs free energies according to

$$\mu_{\text{hyd}}^{\infty}(\text{H}^+) = \frac{1}{2}(\Delta_{AC}\mu_{\text{hyd}}^{\text{con},\infty} - \Delta_{AC}\mu_{\text{hyd}}^{\infty}). \quad (32)$$

According to relations (29) and (30) that may be rewritten for large enough droplets as

$$\mu_{\text{hyd}}^{\infty}(\text{H}^+) = \frac{1}{2}(\Delta_{AC}\mu_{\text{hyd}}^{\text{con},\infty} - \Delta_{AC}\mu_{\text{hyd}}^n) + \frac{e\phi}{n^{1/3}}. \quad (33)$$

Both Tissandier and co-authors and Kelly and co-authors estimated the proton hydration
Gibbs free energy from small ion/water clusters data assuming

$$\mu_{\text{hyd}}^{\infty}(\text{H}^+) \approx \frac{1}{2}(\Delta_{AC}\mu_{\text{hyd}}^{\text{con},\infty} - \frac{1}{c_n}(\mu_{\text{hyd}}^n(\mathbf{A}) + \mu_{\text{hyd}}^n(\mathbf{C}))), \quad (34)$$

here c_n is a constant for ion/water clusters of a given size n . As both the water models
TCPE/2013 and SPC/E predict the term $e\phi/n^{1/3}$ to be far from negligible up to very large
droplets, it is clearly questionable to consider the latter relation as an accurate enough
approximation of relation (33). In line with the earlier conclusion of Vlcek and co-authors,¹⁵
this means that the proton bulk hydration Gibbs free energy estimated from large clusters, in
particular whose size $n \geq 400$, can differ significantly from well accepted estimates computed
from small cluster data.

However we may note that our ion/water polarizable force fields reproduce accurately
high level *ab initio* quantum data regarding the binding energies of small ion/water clusters
whose size is $n \leq 4$. As *ab initio* quantum approaches have been shown to be able to
reproduce accurately experimental results regarding the water stepwise binding energies in
small ion clusters (see among other Ref.,³¹ for instance), we may thus reasonably consider
that our polarizable force fields will provide a proton hydration Gibbs free energy value
based on small cluster data close to the Tissandier's and Kelly's estimates,^{7,8} *i.e.* ranging

539 from 264 to 266 kcal mol⁻¹. On the other hand the proton bulk hydration Gibbs free energy
540 estimated from our $n \geq 400$ droplet data for the pairs Li⁺/F⁻ and Li⁺/Cl⁻ (and accounting
541 for the cation correction δG_C) is about 265 ± 2 kcal mol⁻¹ on average. Even if more efforts
542 are needed to understand how the latter estimate is affected by possible force field artifacts,
543 that estimate suggests the order of magnitude of a possible difference in proton hydration
544 Gibbs free energies as estimated from small clusters and large droplets data to be no larger
545 than 2-3 kcal mol⁻¹, *i.e.* a few k_BT unit.

546 3 Conclusion

547 We computed the absolute single ion bulk hydration Gibbs free energies regarding Li⁺,
548 NH₄⁺, F⁻ and Cl⁻ from bulk simulations of an infinitely replicated water box in which ions
549 are embedded and by extrapolating to their bulk limit data corresponding to finite size
550 ion/water droplets whose water molecular size vary from $n = 50$ to 1000. We also computed
551 the surface potential in pure water droplets whose size ranges from $n = 200$ to 10 000. All
552 our simulations were performed using our own polarizable force fields, in particular for water
553 we consider the water model TCPE/2013.¹⁷ Our main results are

- 554 1. the TCPE/2013 surface potentials of pure water droplets obey a linear function of the
555 droplet "radius" $R_d = n^{1/3}$ as soon as $n \geq 200$;
- 556 2. the ion hydration Gibbs free energy increment arising from additional water molecules
557 added to a given ion/water droplet does not depend anymore from the ion nature no
558 sooner than $n = 400$ droplets;
- 559 3. for ion/water droplets corresponding to $n \geq 400$ and in line with the Born equation,
560 the absolute ion hydration Gibbs free energies μ_{hyd}^∞ and the differences in anion/cation
561 hydration Gibbs free energies $\Delta_{AC}\mu_{\text{hyd}}^n$ are both linear functions of the inverse of the
562 droplet radius $n^{1/3}$;

563 4. the differences $\Delta_{AC}\mu_{\text{hyd}}^n$ obey two different regimes in small cluster ($n \leq 15$) and large
564 droplet ($n \geq 40$) domains.

565 Regarding absolute single ion hydration Gibbs free energies and water surface potentials
566 in droplets, our results computed by means of point charge - point induced dipole polarizable
567 force fields agree with earlier ones derived from standard additive point charge force fields,
568 like those of Kastenholz and Hünenberger¹⁴ regarding water surface potentials and those of
569 Vlcek and co-authors¹⁵ regarding ion hydration enthalpies. In particular the latter authors
570 showed these differences to also obey two different regimes in small cluster and large droplet
571 domains.¹⁵

572 We identified an artifact regarding our cation/water force fields, arising a priori from an
573 inaccurate description of local interactions between water molecules belonging to monovalent
574 cation first and second hydration spheres respectively, and yielding underestimated cation
575 hydration Gibbs free energies in solution. This artifact can be remediated by adding a
576 short range three-body energy term to improve the modeling of interactions among cation
577 and its first/second hydration sphere water molecules. That energy term affects ion/water
578 interactions up to a distance of about 6 Å from the ionic center. Because of the size of all
579 our droplets (for instance the radius of the $n = 50$ droplet is already > 6 Å), we may thus
580 assume the reliability of all the above conclusions.

581 Our results, in particular regarding the two regimes to which the differences in an-
582 ion/cation hydration Gibbs free energy $\Delta_{AC}\mu_{\text{hyd}}^n$ obey in droplets, clearly suggest that
583 methods based on the Cluster Pair Assumption and by considering the thermodynamical
584 properties of small ion/water clusters (whose size $n \leq 6$), like those proposed by Klotz,⁶
585 Tissandier's and co-authors⁷ and Kelly and co-authors⁸ to estimate the absolute proton hy-
586 dration Gibbs free energy in solution $\mu_{\text{hyd}}^\infty(\text{H}^+)$ are questionable. Our present simulations
587 show that taking into account larger ion/water droplets may yield a $\mu_{\text{hyd}}^\infty(\text{H}^+)$ value shifted
588 by several $k_B T$ units compared to CPA/small clusters-based estimates.

Supplementary Material

A free trial version of our code POLARIS(MD) together with the input files allowing to perform the Gibbs free energy computations detailed in the present manuscript for the system NH_4^+ /100 water droplet is available via a link provided as Supplementary Material. They will be soon available on the POLARIS(MD) official website (<http://biodev.cea.fr/polaris/download.html>). Moreover, Supplementary Material materials provide also the details of our polarizable Li^+ /water force field, of the water slab system used to investigate the properties of the liquid water/air interface, the formula allowing one to estimate the entropic cost arising from the perturbation of solute/solvent intramolecular vibrational properties upon ion hydration, the plots of the droplet molecular densities, the raw data and the plots corresponding to the pure water droplet surface potentials, and the surface potential of liquid water computed from our TCPE/2013 model using the "M-convention" scheme and from the pairwise water model TIP3P using the "P-convention" scheme. This material also provides files summarizing the charge and dipole quantities computed along the independent MD droplet trajectories and allowing one to compute the droplet water-vacuum surface potential.

Acknowledgments

We would like to thank Maria Reif and the reviewers for helpful discussions regarding this work. The members of the PhLAM laboratory acknowledge support by the French government through the Program "Investissement d'avenir" (LABEX CaPPA / ANR-11-LABX-0005-01 and I- SITE ULNE / ANR-16-IDEX-0004 ULNE), as well as by the Ministry of Higher Education and Research, Hauts de France council and European Regional Development Fund (ERDF) through the Contrat de Projets Etat-Region (CPER CLIMIBIO). Furthermore, this work was granted access to the HPC resources of [CINES/IDRIS/TGCC] under the allocation 2016-2019 [x2016081859 and A0010801859 and A005030707] made by GENCI.

Table 1: Single ion hydration Gibbs free energy components in kcal mol⁻¹ of the ions $X = \text{Li}^+, \text{NH}_4^+, \text{F}^-$ and Cl^- . V is the virtual atom corresponding to anions or cations (see the scheme detailed in Figure 1). $\delta G_{vib}^{g \rightarrow l}$ is a correction accounting for intramolecular vibrational frequency shifts upon hydration (see Section 1.6). ΔG_∞ is the droplet free energy cost ΔG_n defined in Figure 1 extrapolated for $n \rightarrow \infty$. The ΔG_∞ components $\Delta G_\infty^{\text{sr}}$ and $\Delta G_\infty^{\text{elec}}$ are detailed in Section 2.3. The $\mu_{\text{hyd}}^\infty(X)$ and $\mu_{\text{hyd}}^\infty(V)$ are the single ion bulk hydration Gibbs free energy for the ions and the virtual atoms (for the cation and the anion virtual atoms, the $\mu_{\text{hyd}}^\infty(V)$ values are -12.5 and -9.0 kcal mol⁻¹, respectively). $q\Phi_{\text{bulk}}$ is the free energy cost for a point charge q to cross the liquid water/vapor interface, *i.e.* the water surface potential contribution to the hydration free energy of an ion of charge q . The quantity $\Delta\Delta\mu$ is defined as $\Delta G_\infty - [\mu_{\text{hyd}}^\infty(V) - \mu_{\text{hyd}}^\infty(X)]$. Note the uncertainty affecting the quantities $\Delta G_\infty^{\text{sr}}$, $\mu_{\text{hyd}}^\infty(X)$ and $\mu_{\text{hyd}}^\infty(V)$ is 0.1 kcal mol⁻¹, and the uncertainties affecting the quantities $\Delta G_\infty^{\text{elec}}$ and $\Delta\Delta\mu$ are 0.6 and 0.8 kcal mol⁻¹, respectively (see text for details).

Ion	$\delta G_{vib}^{g \rightarrow l}$	$\mu_{\text{hyd}}^\infty(X)$	$\Delta G_\infty^{\text{sr}}$	$\Delta G_\infty^{\text{elec}}$	ΔG_∞	$\mu_{\text{hyd}}^\infty(X) - \mu_{\text{hyd}}^\infty(V)$	$\Delta\Delta\mu$	$q\Phi_{\text{bulk}}$
Li^+	-0.5	-113.1	+9.1	-114.6	-105.5	-100.6	-4.9	-5.3
NH_4^+	-1.0	-69.2	+11.2	-73.0	-61.8	-56.7	-5.1	-5.3
F^-	+0.4	-103.5	+21.8	-110.6	-88.8	-103.7	+5.9	+5.3
Cl^-	0.0	-77.2	+27.0	-90.4	-63.2	-68.2	+5.0	+5.3

Table 2: Parameters of the function $\mu_{\text{hyd}}^n(\mathbf{X}) = \tilde{\mu}_{\text{hyd}}^\infty(\mathbf{X}) + \tilde{\gamma}_{\mathbf{X}} \cdot n^{-1/3}$ adjusted from droplet data corresponding to $n \geq 400$, all in kcal mol⁻¹. The parameter values are provided \pm their uncertainty arising from our TI computations. Reg.: regression coefficient of the linear fitting process to adjust the parameters. The $\tilde{\mu}_{\text{hyd}}^\infty(\mathbf{X})$ s account for the correction term $\delta G_{\text{vib}}^{g \rightarrow l}$, see Table 1.

Ion	$-\tilde{\mu}_{\text{hyd}}^\infty$	$\tilde{\gamma}$	Reg.
Li ⁺	118.6 \pm 0.5	70.5 \pm 2.9	0.999
NH ₄ ⁺	74.1 \pm 0.3	73.1 \pm 2.9	0.990
F ⁻	96.4 \pm 0.3	80.9 \pm 2.9	0.999
Cl ⁻	69.8 \pm 0.3	79.0 \pm 2.9	0.999

Table 3: Absolute single ion hydration Gibbs free energies in solution and at ambient conditions, in kcal mol⁻¹. The data from our work correspond to the droplet one. ϵ_{err} : mean uncertainty affecting the free energies. (*) : the value is a priori the same as reported by Tissandier *et al.*⁷

	F ⁻	Cl ⁻	Li ⁺	NH ₄ ⁺	ϵ_{err}
This work	97.1	70.7	117.3	74.4	0.6
Tissandier ⁷	102.4	72.7	126.5	84.1	1.9
Kelly ⁸	104.4	74.5	128.4	85.2	≈ 2
Hünnerberger-Reif ⁴⁸	103.5	73.7	125.4		1.2
AMOEBa ⁴⁹		84.6			0.1
DFT-MD ⁴⁴	113.4		119.8		0.7
Force-Matching-MD ⁵⁰	101.4-106.2	68.6-71.8			-
Cluster Theory ⁵¹	104.1	74.3	124.9		(*)
CC-QCT theory ⁵²			116.1		-

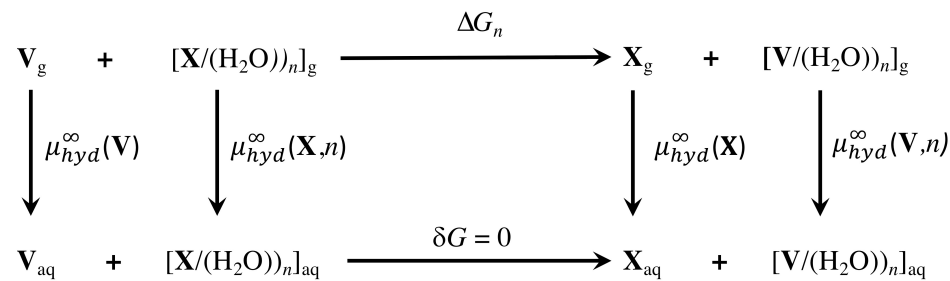


Figure 1: Thermodynamic cycle **TC** showing the link between bulk and finite size system single ion hydration Gibbs free energies.

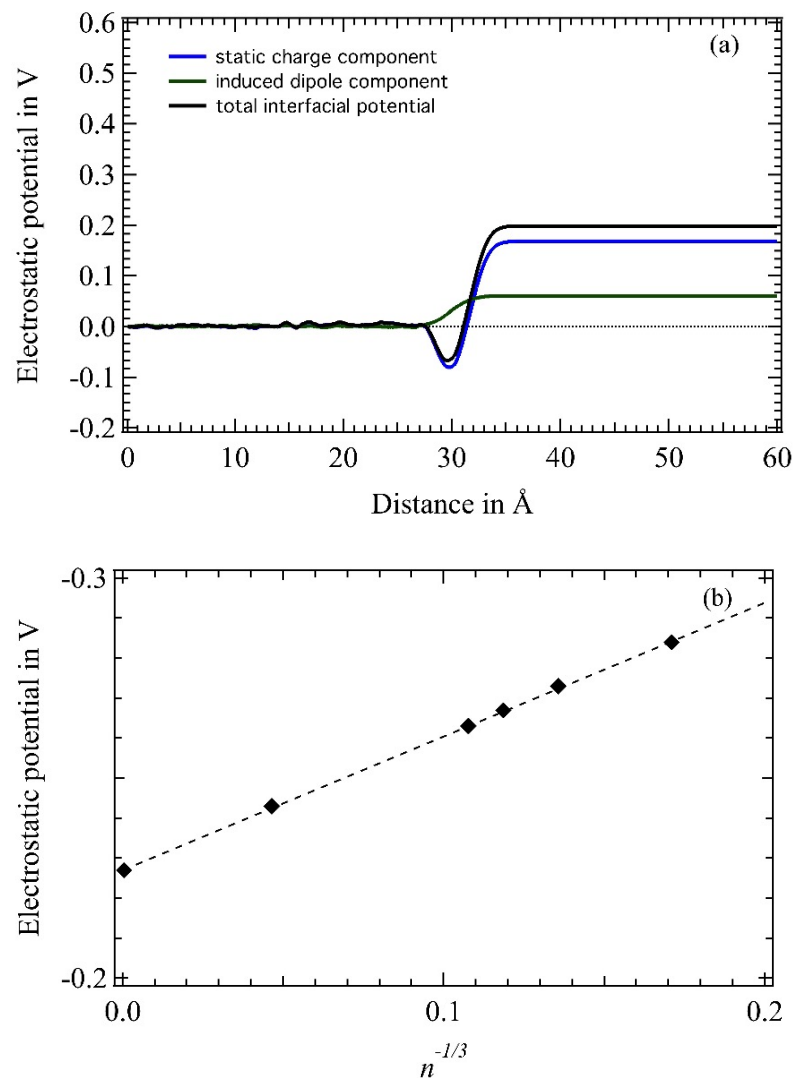


Figure 2: Water surface potential. (a): at the bulk water-vacuum interface, as a function of the distance z to the interface, together with its two components computed from the atomic static charges and the induced dipole moments. (b): surface potential at water droplet surfaces *wrt* to $n^{-1/3}$ and convergence to the bulk limit (shown at $n^{-1/3} = 0$). The uncertainty affecting the surface potential values is smaller than 1 % of the computed values. In dashed line, the linear regression fit, the linear regression coefficient is larger than 0.999.

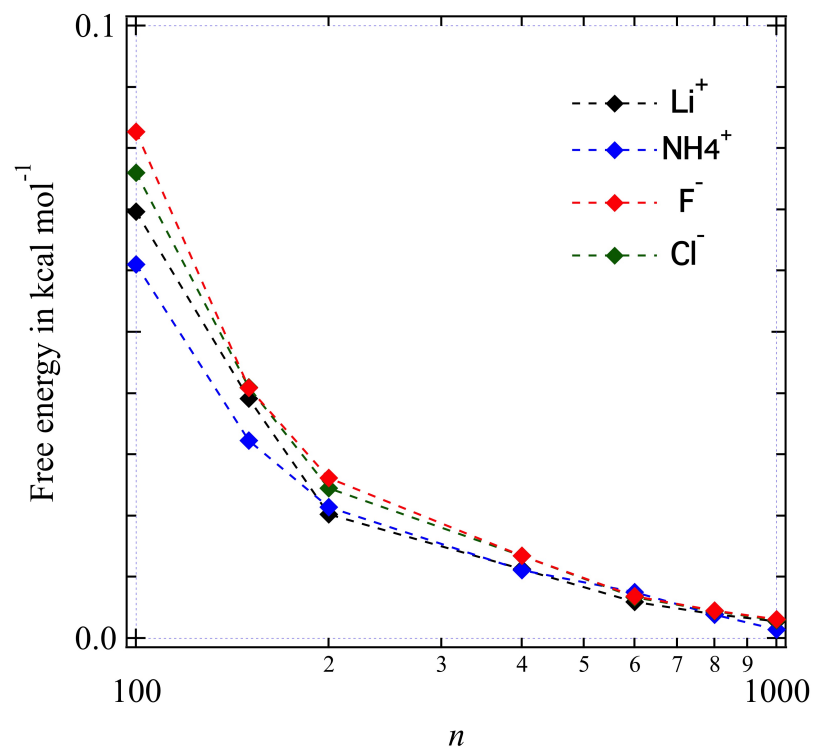


Figure 3: Evolution of the quantity $\delta\tilde{\mu}_{m\rightarrow n}$ as a function of the droplet size n . The uncertainty regarding the plotted values is $\leq 10^{-3}$ kcal mol $^{-1}$, regardless of n .

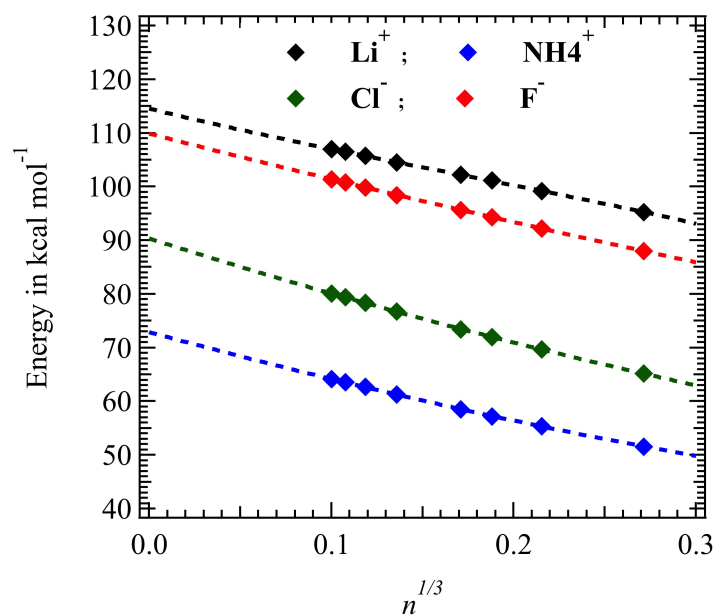


Figure 4: Droplet ΔG_n^{elec} values (diamonds) and the corresponding power law fitted functions (dashed lines). The ΔG_n^{elec} s are the electrostatic components of the free energy costs ΔG_n corresponding to the reaction $\text{V}_g + [\text{X}/(\text{H}_2\text{O})_n]_g \rightarrow \text{X}_g + [\text{V}/(\text{H}_2\text{O})_n]_g$, see Figure 1 and the definitions provided in Section 2.3. The uncertainty regarding the plotted values is $0.1 \text{ kcal mol}^{-1}$ (see text for details).

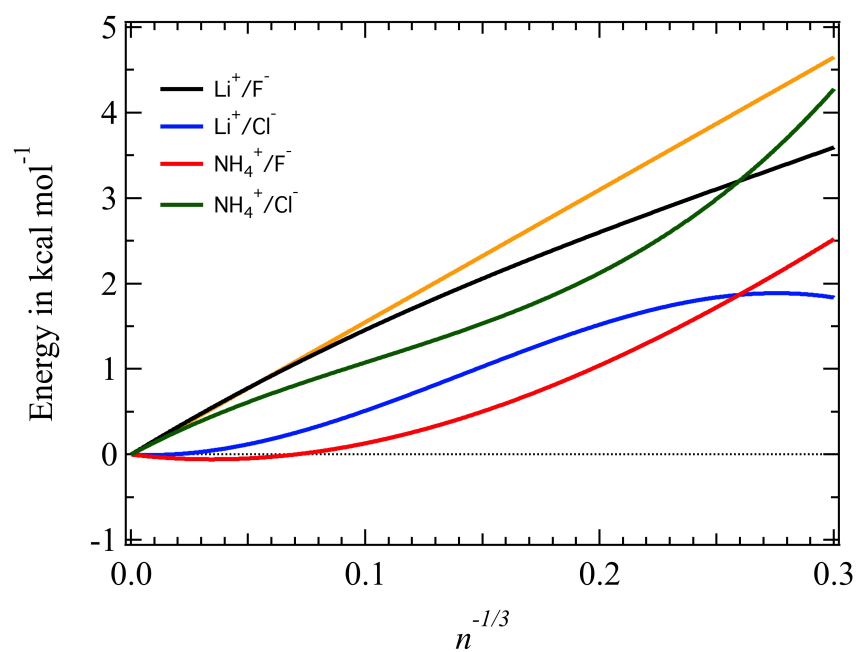


Figure 5: $\Delta\Delta_{\text{AC}}^{n,\infty}$ quantities for all our anion/cation pairs as computed from 4th order power law functions (21). In yellow, the linear function $2e\phi \cdot n^{-1/3}$.

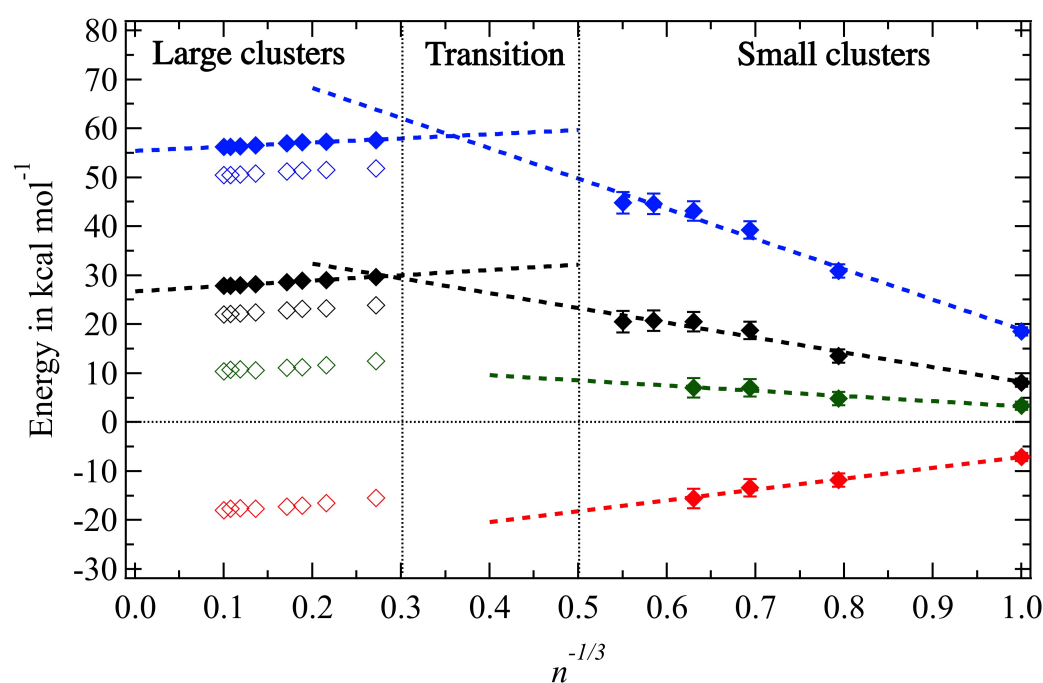


Figure 6: $\Delta_{AC}\mu_{\text{hyd}}^n$ quantities for all our anion/cation pairs from the raw data (empty diamonds) and the corrected ones to account for the cation force field artifact (full diamonds) as computed from our ion/water droplet simulations (*i.e.* data corresponding to $n^{-1/3} \leq 0.3$, the uncertainty affecting these data is 0.1 kcal mol⁻¹); and from the experimental data considered by Kelly and co-workers⁸ (for $n^{-1/3} \geq 0.5$). In dashed lines linear regression functions of the data corresponding to the small and large cluster domains. Same color labeling as in Figure 5.

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PLEASE CITE THIS ARTICLE AS DOI:10.1063/1.5109777

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