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A theoretical study of the hydration of Li⁺ by Monte Carlo simulations with refined ab initio based model potentials

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Abstract Four water models that have the same analytical potential but different degrees of freedom were used to examine the hydration of Li⁺: (a) a polarizable and flexible molecule with constraints that account for the quantal nature of the vibration, (b) a polarizable and classically flexible molecule, (c) a polarizable and rigid molecule, and finally (d) a nonpolarizable and rigid molecule. The goal was to determine how individual molecular properties affect the correct description of the hydration of ions by comparing the structural and thermodynamic predictions for the aqueous solution as made by the different models, which ranged from a very refined one to a simple effective potential. The length of the Monte Carlo runs was large enough to ensure convergence and provide statistically meaningful results; the four models attained good agreement with the experimental data available for the hydration of Li⁺, as well as with the results of the most refined

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simulations. A well-defined first hydration shell was found. It had four water molecules whose dipoles were not aligned to the electric field of the ion because of their hydrogen-bonding with water molecules in outer shells. In the case of the most refined water model, the results showed this pattern clearly. On the other hand, the rigid nonpolarizable version produced a slightly higher hydration number and an almost complete alignment of the dipoles to the ion's electric field. Moreover, a detailed analysis of a microscopic molecular model of hydration showed that the average intramolecular geometry of the water molecules in the first hydration shell was the same as the one for those in the bulk, whereas the electric field of the ion induced a dipole 0.2 D higher in the water molecules of the first hydration shell. The value of the bulk was recovered at the second shell, which explains the good performance of the simplest model. Thus, despite the differences found in the description of the first hydration shell between the polarizable and the nonpolarizable models, the major effect on the polarization of the water molecules resulted from the waterwater interaction.

Keywords Lithium ion hydration \cdot Polarizable force fields \cdot Monte Carlo simulation

1 Introduction

Nowadays, numerical simulations are a powerful tool for the study of physicochemical systems at the molecular level, in particular those involving small molecules like the aqueous solutions of atomic ions. It is now clear that, with the use of refined potentials, simulations can yield results that are in good agreement with the reported experimental data. This gives confidence to observations that are not amenable to experimental determination, such as the molecular details of the hydration of single ions. Single hydration of atomic ions can be readily done in simulations, whereas experimental data are obtained from the solution of salts and then extrapolated to the hypothetical standard state of infinite dilution, where no ion-ion interactions take place [51].

The hydration of the lithium ion has received considerable attention given the many technological applications of Li⁺ in batteries, alloys and lubricants, as well as its use in the treatment of psychiatric disorders and viral diseases – the applications of lithium have been recently reviewed by Birch [7] . On the other hand, the study of the lithium ion is also very attractive from the academic viewpoint; because of its size, it permits the use of a very refined quantum mechanical (QM) level to study its interaction with small molecules such as water [21,22,25]. This has led to multiple theoretical studies regarding the structural and energetic aspects of its hydration [16, 17, 45, 48, 63, 71, 81, 82, 84, 86]. Experimental studies of this phenomenon have also been undertaken, although there are discrepancies in the reported data. Specifically, the hydration number (i.e., the number of water molecules in the first hydration shell of the ion), has ranged from 3.3 to 6.5 depending on the concentration of LiCl in neutron scattering experiments [35,61], where the low values have been observed only at very large concentrations. Diffraction experiments [58,59,64,85] have reported a value of 4 similar to that of other lithium salts [11, 49], regardless of concentration. Raman spectroscopy further adds to disagreement by favoring the four molecule coordination.

Numerical simulations would be expected to help solve the problem, mainly in the low concentration limit. But simulations have also predicted values ranging from 4 to 6. There is a general trend indicating that pairwise simple potentials favour the larger limit [16, 17, 84, 86], while more recent and refined methods such as Car-Parrinello molecular dynamics (CPMD) [48], QM/MM (molecular mechanical) molecular dynamics (MD) [45], and the quasi-chemical theory of liquids (QCTL) [71] favour a hydration of four, even for the infinite-dilution limit. Spångberg and Hermansson studied the hydration of Li⁺ in a recent work [81,82], with potentials ranging from the pairwise expression to the inclusion of nonadditivity and polarizability. Their results show a clear trend from the upper to the lower reported hydration numbers in correlation with the successive complexity of the models. A value of four is found with the potentials that include nonadditive effects and polarizability, the importance of which has been shown in various studies of the ion-water interaction [1,2,6,9,27,28,41,42,53–55,60,65,66].

This work presents a Monte Carlo (MC) study of the Li⁺ hydration at high dilution (one ion and 999 waters). It uses a very refined potential whose results are in good agreement with the experimental observations of water under various different thermodynamic conditions [33,74,76] as well as of aqueous solutions of ions [1,2,9,10]. Simulations are also presented using the same model, though intramolecular geometry and/or polarization were restricted to the average values produced by the model in simulations of liquid water [75]. The modelling of the Li⁺–H₂O interaction is validated by comparison to the available experimental data in the gasphase and in the aqueous solution. This is followed by a detailed analysis of the microscopic structure of the hydrated ion.

2 Methods

Numerical simulations were performed using the MC method. As in the previous study [9] of Na⁺ and K⁺, the MCDHO [74] analytical potential was used for the water-water interactions and the same type of model was employed for the water-cation interactions. This model represents the electron cloud of a molecule as a negative mobile charge density with radial exponential decay attached to a positively charged point core by a harmonic oscillator. The functional form has been presented elsewhere [9,75], and it allows the restriction of intramolecular flexibility and polarizability while simultaneously keeping the same set of parameters; thus, it is possible to separately study the effects that the inclusion of those degrees of freedom has on the results of the simulation [1,2,75]. To this end, MC simulations were performed with the four versions of MCDHO reported in Refs. [75] and [34], viz. MCDHO $_{ff}$, $MCDHO_{fc}$, $MCDHO_r$ and MCD.

2.1 The fitting procedure

The parameters for the Li⁺-water interaction were fitted to reproduce the experimental polarizability of the ion [51], $\alpha = 0.0317 \, \text{Å}^3$, the potential energy surface (PES) built with ab initio calculations of the pair interactions, and the three- and four-body nonadditive contributions to the interaction energy, computed with ab initio calculations of trimers and tetramers.

The calculations of the pair-interaction were performed at the MP2(CP)/6-311++G(3df, 3p) level of the theory; that is, the second order many-body perturbation theory with counterpoise correction [8]. This was also chosen previously [9] because it yields the right polarizability and dipole moment of H₂O; it also yields the right polarizability of Li⁺. All quantum calculations in this work were done with the GAUSSIAN98 [24] program.

The Li⁺-H₂O interaction has been studied extensively by Feller et al. [21,22] and Glendening and Feller [25] who estimated the complete-basis-set limit (CBS) at the fourth-order many-body perturbation theory (MP4) level of the interaction energy to be $\Delta E = -35.0 \, \text{kcal mol}^{-1}$. The calculations of Spångberg and Hermansson [81], on the other hand, yield $\Delta E = -33.5 \, \text{kcal mol}^{-1}$ at the MP2(CP)/TZV++(3df, 3p) level. The calculations performed here resulted in $\Delta E = -33.0 \, \text{kcal mol}^{-1}$.

The final values of the parameters are presented in Table 1, using the notation of Ref. [9]; only nonzero values are shown. It should be stressed that this set of parameters is not unique; other values could produce a similar adjustment to the ab initio data, and more parameters could also be used. The additional criteria for the selection of the values in this work were the following: (1) to use integers for the core and the mobile charges of Li⁺; once q was chosen, the spring constant was determined from the polarizability as $k = q^2/\alpha$; (2) to keep the 12-6 exponents in the Lennard–Jones terms between the mobile charge of the ion and that

of the water molecule, and (3) to use the minimum possible number of parameters to reproduce the ab initio computed energies. There are thus four independent free parameters: the mobile charge of the ion, q, the decay length λ' , and the Lennard–Jones coefficients α_{Li-O} and β_{Li-O} . The parameters q and λ' were initially chosen to yield the best possible reproduction of the interaction energy and the Li⁺-O distance of the optimal dimer, and α_{Li-O} and β_{Li-O} were used to improve the agreement with the ab initio values. This set of parameters was used to generate $Li^+(H_2O)_n$ clusters, with $n = 1, 2, \dots, 6$, from which the three- and four-body nonadditive contributions were calculated at the HF(CP)/6-311++G(3df, 3pd) level of the theory. The correlation energy was neglected in these calculations because it is mostly additive [9,46]. The parameters were then readjusted to a total of 746 points in the pair interaction PES, and 146 three-body and 246 four-body nonadditive contributions to the energy. These points included a sample along the line of optimal Li⁺–H₂O orientation, and configurations with deformed water molecules; thus the intramolecular energy was considered in the many-body expansion of the interaction energy [9,68].

The plot in Fig. 1a shows the behavior of the model along the line of optimal Li^+ – H_2O orientation, compared to the ab initio values; the geometry of the water molecule was fully optimized for each Li^+ –O distance. As previously reported

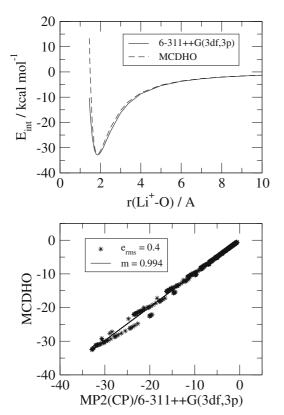


Fig. 1 Comparison of the model to the ab initio pair interaction energies. *Top*: Along the line of optimal ${\rm Li}^+ - {\rm H_2O}$ approach. *Bottom*: To the 746 points in the PES

[9], after 5 Å the interaction is purely electrostatic. For distances shorter than 1.6 Å, the model exaggerates the repulsion due to the short-range Lennard-Jones term. However, at that distance the ab initio energy is already 10 kcal mol⁻¹ above the minimum and the slope of the curve is large enough as to assume that this region will not be importantly sampled in a simulation of the aqueous solution under ambient conditions. On the other hand, the comparison to the 746 points in the ab initio pair interaction PES, at distances longer than 1.7 Å, is shown in Fig. 1b. The slope of the linear regression, m = 0.994, and the mean-square-error attained, $e_{\rm rms} =$ 0.4 kcal mol⁻¹, were deemed satisfactory. Further attempts to add Lennard-Jones terms to the Li+-H interaction did not result in any significant improvement, nor did the use of exponents other than 12-6, nor the use of a Buckingham short-range term.

In regard to the three- and four-body nonadditive contributions, the agreement of the model with the ab initio calculations was less satisfactory: the model overestimated the three- and the four-body nonadditivities by 20% and 55% respectively. The four-body contributions are small (~ 0.5 kcalmol⁻¹) and the three-body are reasonably well reproduced. Because Li⁺ is practically nonpolarizable, it has been shown that the nonadditive contributions to the interaction stem mainly from the polarizability of the water molecule [43]; hence a better agreement could not only be obtained by modifying the water model. The comparison of the model's nonadditive contributions to the ab initio values is shown in Fig. 2a, b. The slope of the linear regression in Fig. 2a is m = 1.1917 with a standard error of $e_{rms} = 0.6$ kcal mol^{-1} , whereas the corresponding values in Fig. 2b are m =1.545 and $e_{rms} = 0.14 \text{ kcal mol}^{-1}$. Though exaggerated, the three- and four-body nonadditive contributions to the energy produced by the model are in agreement with the ab initio

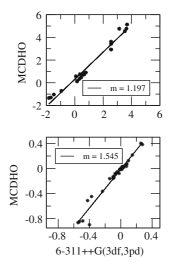


Fig. 2 Comparison of the model to the ab initio non-additive contributions to the interaction energy. *Top*: Three-body contributions. *Bottom*: Four-body contributions. All energy in kcal mol⁻¹

Table 1 Parameters for the Li⁺–H₂O interaction in a.u. Only nonzero parameters are presented

Z	3.0
q	-2.0
k	18.691316
λ'	0.773426
$lpha_{ ext{Li-O}}$	14000.0
$\beta_{ ext{Li-O}}$	-13.0

The definition of each parameter corresponds to the analytical potential of Refs. [9] and [75]

data within the limits of thermal fluctuations at T = 298 K. Thus, it was decided to use the model with the parameters of Table 1.

3 Results

3.1 Small clusters

To test the reliability of the models and their parametrization, their predictions of the interaction energy for various different $Li^+(H_2O)_n$ clusters (n = 1, 2, ..., 6) were compared to ab initio calculations [21,22,25] and gas-phase experimental data [18,72]. In the case of the dimer (n = 1), the geometrical parameters were also compared along with the water dipole moment (Table 2). It is worth mentioning that there is no unique way of assigning electrical properties to individual molecules from ab initio calculations of clusters. However, the quantum theory of atoms in molecules [3], based on the analysis of the topological properties of the electron density, provides a reasonable framework to assign charges to different sites. This method was used in this case to estimate the water dipole moment from the MP2(CP)/6-311++G(3df,3p) calculation, as was the case in previous studies [74].

Table 2 presents a comparison between the predictions for the $\text{Li}^+\text{-H}_2\text{O}$ dimer at the best ab initio level [21], the one computed in this work at the MP2(CP)/6-311++G(3df, 3p) level, and those predicted by the classical models used in this work.

Of course, the ab initio interaction and the Li⁺–O distance computed in this work were well reproduced by the MCDHO model since they were used in the fitting procedure. But the intramolecular geometry of the water molecule in the dimer and its dipole moment were not fitted. The flexible model for water produced a longer r(O-H) bond than the ab initio calculation and a somewhat larger dipole. The rigid polarizable MCDHO $_r$ model reproduced the energy and the separation quite well, with an even larger dipole for water. The nonpolarizable MCD model yielded a slightly longer separation and underestimated the interaction energy by 6%. Because the parameters of the MCDHO models were not

readjusted in this work, the longer bonds and narrower bond angle produced by them can be ascribed to some extent to the parametrization with a different water monomer deformation PES [74]. This different geometry, in turn, affects the dipole moment. Even if the differences found with the MCDHO $_{fc}$ model amounted to only 2% in the bond length and less than 1% in the bond angle, the difference in the dipole moment amounted to 15%.

The interaction energies of larger clusters are shown in Table 3, where the geometries are represented in terms of the number of water molecules in the first and second hydration shells, n_1 and n_2 , respectively, and they correspond to Fig. 1 of Ref. [25]. The ab initio results presented in this work and those of the classical models were compared to the best ab initio values [25]. The configurations were optimized with each version of the MCDHO model in all cases, whereas it was only possible to make full ab initio optimizations for clusters with up to n = 4 water molecules. For the larger clusters single-point ab initio calculations were performed with the configurations optimized with MCDHO f_c , and the CP correction was applied to obtain the interaction energies. As it should be expected from the comparison with the dimer, all MP2(CP)/6-311++G(3df, 3p) interactions turned out smaller than the best ab initio value [25]: for n = 1, 2the difference is roughly $1.5 \,\mathrm{kcal} \,\mathrm{mol}^{-1}$, and it increases up to 4 kcal mol^{-1} for n = 3, 4. The larger discrepancies for n = 5, 6 stem from using the optimal geometries produced by MCDHO f_c , which do not correspond to the optimal ab initio geometries.

In regard to the performance of the model, the four versions were able to produce the right configurations as stable local minima in their respective energy surfaces, albeit with somewhat different geometric parameters (distances and angles). Generally speaking, there was a fairly good agreement with the ab initio values obtained in this study, again smaller than the best calculations. However, as the number of water molecules increases, the water—water hydrogen-bonding becomes more important, in the absence of direct interaction with the ion. This is reflected in a smaller difference relative to the best calculations in the interaction energy for the six-water cluster than would be expected from the Li⁺—H₂O pair. Even the nonpolarizable MCD model kept some of this nonadditivity, as explained in Ref. [75].

A further validation of the models can be made by comparing their predictions of the enthalpies of succesively larger clusters at $T=298\,\mathrm{K}$, $\Delta H_{\mathrm{hyd}}^{298}$, to the theoretical and experimental data reported in the literature. This was done by performing MC simulations of the clusters, and using the ideal-gas formula [1,15,81]

$$\Delta H = \Delta U + p\Delta V = \Delta U - nRT,\tag{1}$$

where p=1 atm is the pressure, n is the number of water molecules in the cluster, R is the gas constant, and T=298 K the temperature. The results are presented in Table 4, showing

Table 2 Comparison to ab initio data of the interaction energy and the geometrical parameters of the optimal Li⁺-H₂O dimer produced by the models

	Est. CBS ^a	This work ^b	$\mathrm{MCDHO}^{\mathrm{c}}_{fc}$	$MCDHO_r$	MCD
Eint	-35.2	-33.0	-33.3	-33.2	-31.0
$r(Li^+-O)$	1.847	1.8593	1.8525	1.8523	1.8931
r(O–H)	0.962	0.9631	0.9832	0.9840	0.9840
∠HOH	105.0	105.5	104.9	102.7	102.7
$\mu_{ m H_2O}$		3.11 ^d	3.60	3.65	2.96

The energy is in kcal mol⁻¹, the distances are in Å, the angle in degrees and the dipole moment in Debye

Table 3 Comparison to ab initio data of the interaction energy of the optimal Li^+ – $(H_2O)_n$ clusters produced by the models (n = 1, ..., 6)

\overline{n}	Geometry	Ref. [25] ^a	This work ^b	$MCDHO^{c}_{fc}$	$MCDHO_r$	MCD
1	$1+0 (C_{2v})$	-34.5	-33.0	-33.3	-33.2	-31.0
2	$1+1(C_s)$	-51.5	-49.8	-52.6	-51.7	-46.4
	$2+0(D_{2d})$	-64.4	-63.1	-61.0	-60.7	-58.4
3	$2+1(C_s)$		-76.8	-78.0	-77.2	-72.6
	$2+1(C_2)$	-80.2		-76.9	-76.3	-74.0
	$3+0(D_3)$	-87.5	-84.0	-81.0	-80.6	-79.8
4	$3+1(C_2)$	-103.1	-97.8	-97.7	-97.0	-92.4
	$4+0(S_4)$	-104.1	-100.4	-96.4	-95.6	-94.8
5	$4+1(C_2)$	-118.5	-111.4^{d}	-112.3	-111.5	-109.4
6	$4+2(C_s)$	-130.9	-120.7^{d}	-127.1	-126.0	-121.5

The geometries are represented as $n_1 + n_2$, where n_1 is the number of molecules in the first hydration shell and n_2 the number of molecules in the second hydration shell. The energies are in kcal mol⁻¹, and the geometries are those of Ref. [25]

Table 4 Comparison to experimental data (second and third columns) and theoretical ab initio estimates (fourth column) of the cumulative enthalpy of the Li⁺-(H₂O)_n clusters produced by the models in gas-phase simulations at T = 298.15 K (n = 1, ..., 6)

n	HPMS ^a	CIDb	Ab initio ^c	$MCDHO_{fc}$	$MCDHO^{d}_{ff}$	$MCDHO_r$	MCD
1	-34.0	-32.7	-34.2	-33.1	-32.9	-33.0	-30.6
2	-59.8	-60.0	-61.7	-60.2	-60.0	-60.0	-56.9
3	-80.8	-82.5	-83.4	-79.4	-79.5	-79.0	-77.7
4	-96.9	-99.4	-98.6	-93.0	-92.8	-93.1	-91.7
5	-110.8	-113.8	-111.7	-104.2	-104.5	-104.2	-102.0
6	-122.9	-128.8	-122.4	-116.1	-115.0	-114.5	-110.1

The enthalpies were estimated with the ideal gas formula [15,1,81] $\Delta H = \Delta U + p\Delta V = \Delta U - nRT$, where *n* is the number of water molecules in the cluster. The values are in kcal mol⁻¹

the same behavior as in Table 3. For a more refined analysis, the enthalpies of the detachment reactions

$$\text{Li}^+(\text{H}_2\text{O})_n \to \text{Li}^+(\text{H}_2\text{O})_{n-1} + \text{H}_2\text{O}$$
 (2)

are plotted in Fig. 3. Though the models' values were systematically lower than the experimental data, the qualitative behavior of MCDHO $_{fc}$ was the same as that of the collision-induced dissociation data (CID) since the enthalpy change

from n = 6 to n = 5 is higher than from n = 5 to n = 4. A general feature found with the four models was that, in agreement with the ab initio data, they favoured a four-membered first hydration shell, given that the optimal configurations of larger clusters all have $n_1 = 4$ (Table 3).

The previous comparison led to the conclusion that the model was suitable for numerical simulations of the aqueous solution under ambient conditions.

^aEstimated complete basis set limit MP2 from Refs. [21,22]

^bOptimized with MP2(CP)/6-311++G(3df, 3p)

^cThe optimal dimer is the same for MCDHO ff

^dComputed from the charges obtained with the atoms-in-molecules method [3]

^aCounterpoise corrected values MP2(CP)/6-31+G*//RHF/6-31+G*

^bComputed with MP2(CP)/6-311++G(3df, 3p) on configurations optimized with the same level of the theory for n = 1, 2, 3, 4, and on the configurations optimized with MCDHO_{fc} for n = 5, 6

^cThe optimal clusters are the same for $MCDHO_{ff}$

^dGeometries optimized with the MCDHO fc model

^aHigh-pressure mass spectrometric measurements [18]

^bCollision-induced dissociation data [72]

^cAb initio calculations with large correlation-corrected basis sets [21]

^d An energy of (3/2)RT per molecule was added to correct for the classical sampling of the intramolecular degrees of freedom

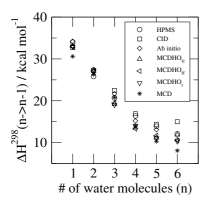


Fig. 3 Enthalpies of the detachment reactions $\text{Li}^+(\text{H}_2\text{O})_n \to \text{Li}^+(\text{H}_2\text{O})_{n-1} + \text{H}_2\text{O}, n = 1, 2, \dots, 6$, in kcal mol⁻¹. *HPMS*: High-pressure mass spectroscopy [18]; *CID*: Collision-induced dissociation [72]; ab initio data from Ref. [25]

3.2 Aqueous solution

Following the procedure described in Ref. [9], MC simulations with the Metropolis algorithm [29,57] were performed on the NVT ensemble at $T = 298.15 \,\mathrm{K}$, for a system of 999 water molecules and one Li+ ion in a cubic cell of $V = 29915 \,\text{Å}^3$, thus a density of $\rho = 0.999 \,\text{g cm}^{-3}$. A simulation of the same system was performed with each of the four versions of the MCDHO model. In all cases the lithium ion was considered polarizable, although its polarizability is very low. To explore a larger region of the configurations' space, each of the four simulations was distributed in five CPU's, as described in Ref. [75]. A spherical cutoff radius of $R_c = 10.85 \,\text{Å}$ was used for the intermolecular interactions, as well as periodic boundary conditions with Ewald sums [20] to account for long-range electrostatics. The adiabatic nuclear and electronic sampling (ANES) [12,52] scheme was used to treat the polarizability of MCDHO $_{ff}$ and of MCDHO $_r$, as well as the polarizability and flexibility of MCDHO fc (using T_{am} =0.05 K and a sampling ratio quantum:classical of 10:1). The blocking method of Flyvbjerg and Petersen [23] was used to make sure convergence was attained and statistical sampling was significant. The hydration enthalpy ΔH_{hyd} was computed as the difference between the average of the total interaction energy of the system, $\langle E_S \rangle$, and n times the average energy of a single water molecule, $\langle E_{\rm H_2O} \rangle$, i. e.,

$$\Delta H_{\text{hyd}} = \langle E_{\text{S}} \rangle - n \langle E_{\text{H}_2\text{O}} \rangle, \tag{3}$$

where n=999 is the number of water molecules in the system. This is a small difference between two large values, $\Delta H_{\rm hyd}$ being only a hundredth of the total energy of the system, and therefore very sensitive to the choice of $< E_{\rm H_2O} >$. To ensure that the reference value had exactly the same conditions as the aqueous solution, it was chosen as the average energy of the water molecules located at distances longer than 7 Å within the same simulation of the ion, instead of the average obtained from previous simulations of pure water [75]. This procedure has been applied successfully in previous studies of various different solutes [31,32,77]. Besides,

the discrepancy turned out to be less than 0.1 kcal mol⁻¹ per molecule for the four models.

The graph at the top of Fig. 4 shows the cumulative average of the predicted hydration enthalpy as a function of the number of MC cycles (1 $N_{\rm cycle} = 10^3$ configurations), after an equilibration period of 2.5×10^5 N_{cycles}. The bottom graph of Fig.4 shows that a plateau was reached for the standard deviation of each model, making the sample statistically meaningful.

The energetic and structural parameters of the Li⁺ hydration obtained from the simulations in this work are compared to experimental data in Table 5, and to the results of other simulations and theoretical calculations in Table 6. As discussed by Grossfield [26], the experimental estimates of the hydration enthalpy vary as much as 20 kcal mol⁻¹ from one another, because they depend on the partition scheme used to distribute the hydration enthalpy of the salt among the resulting solvated ions. Yet, the values obtained from the simulations in this study lie within the extremes and can thus be considered in as good agreement with the experimental data as the most refined theoretical results [45,48,71,81,82].

Another issue of discussion regards the hydration number [86]. As previously mentioned, the estimates made from experimental data range from 3.3 to 6.5 and seem to depend strongly on the salt concentration [35,61], though X-ray diffraction [59,85] and Raman spectroscopy [38,73] yield a hydration number of four. This is also the case of the most recent and refined numerical simulations and theoretical

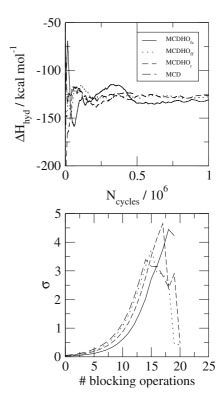


Fig. 4 *Top*: Cumulative solvation energy versus number of Monte Carlo cycles. *Bottom*: The corresponding standard deviations versus the blocking operations

Table 5 Li⁺ hydration energetic and structural properties obtained from the simulations in this work, compared to the experimental data reported in the literature

	$n_{ m H_2O}/n_{ m Li}$	$-\Delta H_{ m hyd}$	< n >	r _{MO}	$r_{ m MH}$	φ
CISDa		134				
NDIS ^b	5.6		3.3(5)	1.95(2)	2.50(2)	52(5)
NDIS ^b	15.6		5.5(5)	1.95(2)	2.55(2)	40(5)
$MTDH^{c}$		127	5.2	` ,		
$CISD^d$		138				
NDIS ^e	4.0		3.2(2)	1.96(2)	2.52(2)	
NDISe	15.4		6.0(4)	1.95(2)	2.52(2)	
NDIS ^e	55.6		6.5(2)	1.96(2)	2.52(2)	
$CISD^f$		138				
$MTDH^g$		121		1.94-2.28		
FEISh			5.1			
$MCDHO_{fc}$	999	129	4.0	1.97	2.54	40
$MCDHO_{ff}$	999	128	4.0	1.97	2.56	0
$MCDHO_{rc}^{jj}$	999	127	4.0	1.97	2.54	0
MCD	999	125	4.3	1.99	2.65	0

The hydration enthalpy $\Delta H_{\rm hyd}$ is in kcal mol⁻¹, and the distances in Å; $r_{\rm MO}$ is the location of the first peak of the radial distribution function $g_{\rm Li-O}(r)$. The first maximum of $g_{\rm Li-H}(r)$ is located at $r_{\rm MH}$. ϕ is the angle (in degrees) between the dipole of the water molecule and the Li-O axis. The numbers in parentheses indicate the uncertainty in the last reported digit

Table 6 Li⁺ hydration energetic and structural properties obtained from the simulations in this work, compared to the theoretical data reported in the literature

	$n_{ m H_2O}/n_{ m Li}$	$-\Delta H_{ m hyd}$	< n >	$r_{ m MO}$	$g_{\text{Li-O}}(r_M)$	r_m	$r_{ m MH}$	ϕ
MD-TIP4P ^a	150	125	5.9(3)	2.09(2)			2.57(2)	
MC-TIP4P ^b	124		5.8(4)	2.20	9.5	3.11	2.78	26
QCTL ^c		128	4.0	2.00				
$CPMD^d$	32		4.0	1.96	8.5		2.58	40-50
MD-IMC ^e	256	100	4.0	1.96	7.0			
MD-SPC/E ^f	255		4.4	1.97	13.8	2.56		0
QM/MM-HF ^g	499		4.2	1.95	9.9	2.97		30
QM/MM-DFT ^g	499		4.1	1.93	9.9	2.71		40
MD-RWK2h	230	116	4.6	1.93	9.5			10-15
MD-3BP-SPC/E ⁱ	512	120	4.0	1.96	8.8	2.66	2.62	0
QCSM ^j	89		4.0	2.05		2.78	2.71	
$MCDHO_{fc}$	999	129	4.0	1.97	9.9	2.61	2.54	40
$MCDHO_{ff}$	999	128	4.0	1.97	9.7	2.64	2.56	0
$MCDHO_{rc}$	999	127	4.0	1.97	9.7	2.66	2.54	0
MCD	999	125	4.3	1.99	9.2	2.85	2.65	0

The hydration enthalpy $\Delta H_{\rm hyd}$ is in kcal mol⁻¹, and the distances in Å; $r_{\rm MO}$ is the location of the first peak of the radial distribution function $g_{\rm Li-O}(r)$, and r_m the following minimum. The first maximum of $g_{\rm Li-H}(r)$ is located at $r_{\rm MH}$. ϕ is the angle (in degrees) between the plane of the water molecule and the Li–O axis. The numbers in parentheses indicate the uncertainty in the last reported digit

^aCISD Cluster ion solvation data, Ref. [40]

^bNDIS Neutron diffraction and isotopic substitution, Ref. [61]

^cMTDH Model for the thermodynamics of hydration, Refs. [50,51]

^dCISD Cluster ion solvation data, Ref. [13]

^eNDIS Neutron diffraction and isotopic substitution, Ref. [35]

^f CISD Cluster ion solvation data, Ref. [83]

^gMTDH Model for the thermodynamics of hydration, Ref. [78]

^hFEIS Field evaporation of ions from solutions, Ref. [39]

^aMolecular dynamics (MD) with the TIP4P model of water [37], Ref. [84]

^bMonte Carlo (MC) simulation with the TIP4P model of water [37], Ref. [16]

^cQuasi-chemical theory of liquids, Ref. [71]. The free energy $\Delta_{hyd}G$ is shown, instead of the enthalpy

^dCar-Parrinello molecular dynamics simulation CPMD, Ref. [48]

^eMolecular dynamics with the SPC model of water [5] and an effective potential for the Li⁺–H₂O interaction, obtained with the inverse MC (IMC) method [47], Ref. [48]. The free energy $\Delta_{\text{hvd}}G$ is shown, instead of the enthalpy

^f Molecular dynamics with the SPC/E model of water [4] and a Li⁺-H₂O model of Pettitt and Rossky [69], Ref. [86]

^gQuantum-mechanical molecular-mechanical (QM/MM) MD simulations, Ref. [45]

^hMolecular dynamics with the RWK2 model of water [70], Ref. [17]

¹Molecular dynamics with an effective three-body potential (3BP) based on the SPC/E model for water [4], Refs. [81,82]

^jCombined quantum-chemical statistical-mechanical simulation, Ref. [63]

methods [45,48,71,81,82]. As shown below, the results of this work are in agreement with a hydration number of four.

The Li⁺–O radial distribution functions (rdf's) are shown in the top graph of Fig. 5. The four models employed produced very similar results, with the small differences shown in the bottom graph. MCD produced the largest departure from MCDHO f_c . This was mainly due to the occurrence of its first maximum of $g_{Li-O}(r)$ at a longer distance than those of the other models, which is consistent with the larger Li⁺-OH₂ separation found previously for the pair (Table 2). In contrast to other singly-charged cations [9], a depletion zone similar to that produced by doubly-charged cations [6] was found. This was due to lithium's shorter hydration radius. The Li⁺-H rdf also showed a first peak (Fig.6), suggesting a well defined first hydration shell, but without the depletion zone that is produced for instance by Mg²⁺ [6]. In this regard, Li⁺ proved similar to Ca²⁺ [6], a result that is consistent with the mean lifetime of a water molecule in the first hydration shell of each of the three cations (see e.g. Fig. 1 of Ref. [30]) and with the fact that the classical "solventberg" model is adequate to describe the mobility of the Li⁺ ion in water [43]. After the first hydration shell, a second one could be appreciated in both rdf's. There was even a third one gauging from the third peak of $g_{Li-H}(r)$.

An estimate of the hydration number can be made from integrating $g_{\text{Li-O}}(r)$ up to the first minimum. The result of this procedure is presented in Fig. 7a, where it can be seen that the three polarizable models yielded the same value of

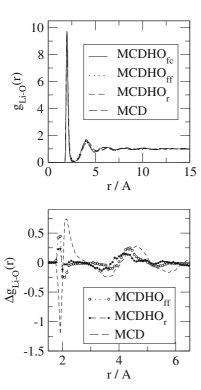


Fig. 5 *Top*: Radial distribution functions $g_{\text{Li-O}}(r)$ produced by the four models. *Bottom*: difference relative to the results obtained with MCDHO f_C

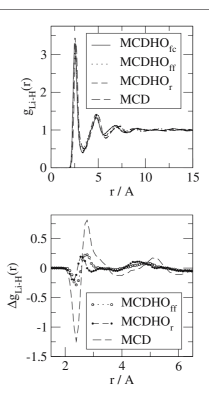


Fig. 6 *Top*: Radial distribution functions $g_{\text{Li-H}}(r)$ produced by the four models. *Bottom*: Difference relative to the results obtained with MCDHO f_c

four, while MCD produced a somewhat higher prediction, 4.3. This discrepancy can be explained with the histogram in Fig. 7b: the three polarizable models kept four molecules in the first hydration shell for more than 90% of the simulation, with small departures to three and to five. Alternatively, in the case of the simulation with MCD a, first hydration shell of four amounted to less than 50% and hydration numbers of three, five or six also occurred. The results of the polarizable models were in good agreement with results from QM-MM-MD [45], with the QCTL [71] and with earlier predictions derived from ab initio calculations [21].

It is worth pointing out that even the simple MCD model yielded reasonably good energetic and structural results. Previously such performance had only been recorded by the effective pairwise potential of Zhou et al. [86], and by SPC/E [4] combined with the nonadditive ion—water potentials of Dang [15].

The distribution of the angle ∠O-Li-O for water molecules in the first hydration shell is shown in Fig. 8. In agreement with CPMD [48] and QM-MM-MD [45], the three polarizable models produced a maximum at 109.4°, evidence of a rather stable tetrahedral arrangement, with a tail toward wider angles, indicating the disruption of the structure by exchange reactions that altered the hydration number temporarily. On the other hand, the MCD model yielded a wider distribution with a maximum at 100.0° and a significant population around 180°, indicating the presence of planar arrangements. This last result is similar to the one obtained

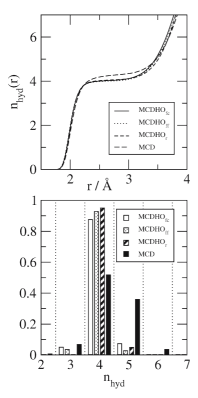
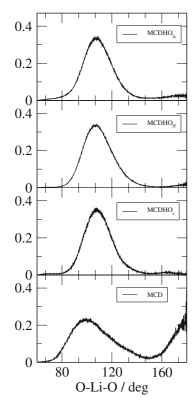


Fig. 7 *Top*: Cumulative number of water molecules $n_{\text{hyd}}(r)$ obtained from integrating $g_{\text{Li}-\text{O}}(r)$. *Bottom*: Histogram of the number of water molecules in the first hydration shell



 $\begin{tabular}{ll} Fig.~8 & Distribution of the angle O-Li-O among water molecules in the first hydration shell \\ \end{tabular}$

by Duan and Zhang [17] with the flexible RWK2 model for water [70].

In order to determine the orientation induced on the water molecules by the ion's electric field, the angle ϕ between the position vector of the water's oxygen relative to the Li⁺ ion, ${\bf r}_{\rm Li-O}$, and the dipole moment of the water molecule, $\mu_{\rm H_2O}$, was computed. In the case of MCD this corresponds to the usually quoted tilt angle [48,61] between the \mathbf{r}_{Li-O} vector and the bisector of the ∠H–O–H angle (some authors use the supplementary angle [15, 17, 45, 86]). The distributions of ϕ for each of the four studied models and each of the four regions (three hydration shells and the bulk) are shown in Fig. 9. MCDHO_{fc} yields a maximum at $\phi \approx 40^{\circ}$, in good agreement with neutron diffraction data [61] and simulations with QM force-fields [45,48]. MCDHO_r had a local maximum at the same value of ϕ , whereas MCDHO_{ff} only had a noticeable shoulder, a feature shared with less intensity by MCD. The behavior of the latter, nonpolarizable model resembled that of effective pair potentials, as in the case of Zhou et al. [86] – even if combined with nonadditive ion—water potentials [15].

Among the first attempts to explain the deviation from the alignment to the electric field, a specific interaction of Li⁺ with a "lone-pair" of the water molecule was proposed [61]. The pair interaction obtained from ab initio calculations [21] has proved that interpretation to be wrong, as the optimal configuration corresponds to $\phi = 0$. Instead, the deviation can be ascribed to enhanced hydrogen-bonding between water molecules in the first and second hydration shells [9].

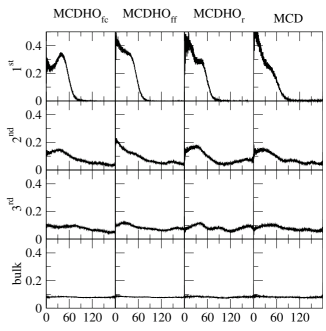


Fig. 9 Distribution of the angle between the electric field of the ion and the dipole moment of each water molecule in the first, second and third hydration shells, and the bulk, from *top* to *bottom*.

It is worth mentioning that in a recent interpretation of neutron diffraction data on aqueous solutions of hydroxides, at solute concentrations ranging from 1 solute per 12 water molecules to 1 solute per 3 water molecules, by means of Soper's empirical potential structure refinement (EPSR) technique [79,80], Imberti et al. [36] concluded that there was a rather strong alignment induced on the dipole moments of first-shell water molecules by the cation's electric field. While this happened with Li⁺ and Na⁺, in the case of K⁺ they found the aforementioned 45° deviation. Moreover, they claim that their results on this alignment are in agreement with the CPMD simulations of Lyubartsev et al. [48], in spite of the latter authors' mention of a 40–50° deviation. Another discrepancy between Imberti's results and those from refined simulations, including those presented here, lies in the predicted structure of the first hydration shell: the first minimum of their Li⁺-O rdf does not reach a value low enough to correspond to a depletion layer. This is in disagreement with previous experimental results on the residence time of a water molecule in the first hydration shell [30]. Though Imberti et al. [36] did not report the hydration number of any of the cations they studied, it is likely to be larger than four in the case of Li⁺, gauging from their rdf's.

The EPSR technique involves refining a starting interatomic potential energy function in a way that produces the best possible agreement between the simulated and measured site-site partial structure factors [80]. But in order for EPSR to be successful the rdf's must uniquely determine the higher order correlation functions. This, in turn, can only occur in a pairwise additive system [19,80]. Thus, a possible explanation of the discrepancies mentioned above is the use of a rigid nonpolarizable model in the MC simulations of Imberti et al. [36]. As was previously mentioned, this study found that the MCD model does predict the alignment of the dipoles of first-shell water molecules to the ion's electric field.

Figure 9 shows that the water molecules in the second shell are still affected by the electric field of the ion, though to a much less extent. The orientation in the third shell is almost random, showing the effect to be short-ranged. Thus a completely random orientation was reached after 7 Å from the ion.

To analyze the effect of the ion's electric field on the individual water molecules, the per-molecule dipole moment was computed at each region. The corresponding distributions are presented in Fig. 10. It is clear that the induced dipole is 0.2–0.26 D higher for waters in the first hydration shell, as compared to the bulk, but the bulk value is recovered from the second shell onward. The effect of the ion's field on the intramolecular geometry is even lower. The graphs in Fig. 11 show that the r(O-H) bond-length and the HOH angle of the waters in the first hydration shell are only slightly different from those of bulk waters. Even the asymmetry between the two bonds shown in the rightmost graphs of Fig. 11 is identical in the first hydration shell to that in the bulk. This mild and short-ranged effect on the geometric properties of the water molecules explains the good performance of the rigid $MCDHO_r$ model.

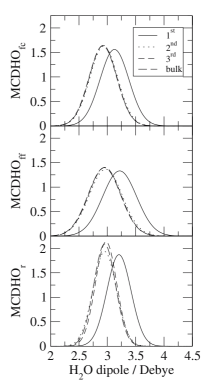


Fig. 10 Distributions of the dipole moment of each water molecule in the hydration shells and in the bulk as produced by the three polarizable models. The averages in the first shell are $\mu = 3.13$ D with MCDHO $_{fc}$, $\mu = 3.22$ D with MCDHO $_{ff}$, and $\mu = 3.21$ D with MCDHO $_{fc}$. The bulk values of Ref. [75] are fully recovered at the second shell

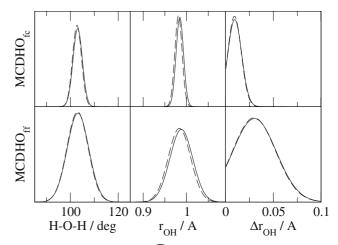


Fig. 11 Distributions of the \overrightarrow{HOH} bond-angle (left) and r_{OH} bond-lengths (middle) of the water molecules in the first hydration shell ($continuous\ line$), compared to the bulk ($broken\ line$). The $right\ column$ shows the distribution of the difference between the two r_{OH} bond-lengths of each molecule. The averages in the first hydration shell are slightly larger than in the bulk

4 Concluding remarks

For this study, $a \rightarrow c Li^+-H_2O$ analytical potential was fitted to ab initio calculations and to the experimentally determined

polarizability of the ion. This was employed with four versions of the MCDHO model for water [74,75] in numerical simulations of the diluted aqueous solution under ambient conditions. The good agreement with the experimental data validated the models, thus showing the reliability of their predictions. In particular, there is a clear convergence with other theoretical methods that predict a hydration number of four, even in much diluted solutions. Hence, the experimental determination of this value should be revised.

The interaction of Li⁺ with its first hydration shell produces a very stable and rigid tetrahedral coordination. Though not to the same extent as Mg²⁺, Li⁺ behaves as a hydrated ion. It is likely to be modeled as a single entity using the hydrated ion model [67], which would prevent the appearance of five- and six-coordination that would arise from using effective pairwise potentials. The bulk water geometry should probably be used for the molecules in the hydrated ion model instead of the optimal geometries for the gas-phase cluster, as suggested by the results shown in Fig. 11.

The suite of models presented here performed well when reproducing experimental data and were in agreement with refined simulations. The good execution of the rigid polarizable MCDHO_r model, which was also found in previous simulations [33], supports the view that bulk water geometry is quite stable. The nonpolarizable MCD model performs better than other simple pairwise potentials in regards to both energetic and structural predictions. In fact, its more than modest performance deserves some further comment, all the more so since the failure of effective pairwise potentials to describe aqueous solutions of ions has been well documented (see for instance [14]), especially on the subject of the hydration number. The results with the MCD model in this work also showed an overestimation that was, however, smaller than that of other models [81]. The other major discrepancies relative to the more complex models were the reduced stability of the tetrahedral arrangement of the firstshell water molecules and the enhanced alignment of their dipoles with the cation's electric field. The performance of the MCD model resembles that of the SPC/E [4] with the nonadditive ion-water potentials of Dang [15], which have been successfully applied to the study of aqueous solutions under different thermodynamic conditions [43,44,56,62]. Dang's potentials include explicit polarization and three-body terms to treat the nonadditive character of the water-ion-water interactions, and the SPC/E model takes into account the energetic cost of the polarization of the water molecules in the liquid phase. These features are intrinsically built in the MCDHO family of models [74], including MCD [75].

The failure of the nonpolarizable water models to correctly describe aqueous solutions of ions has been ascribed to their inability to respond to the ion's electric field, which should induce larger dipoles preferentially aligned to itself and thus produce repulsive dipole–dipole interactions [14]. This effect is usually minor for singly charged cations (see e.g. [9]). But in the case of Li⁺ it is enhanced because of the short ion–water distance. The results of Fig. 10 in this work are consistent with this interpretation, since they show

a significantly larger dipole moment of the first-shell water molecules in relation to those in the bulk.

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