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Molecular electrometer and binding of cations to phospholipid bilayers †

Andrea Catte,^{*a*[‡]} Mykhailo Girych,^{*b*} Matti Javanainen,^{*c,d*} Claire Loison,^{*e*} Josef Melcr,^{*f*} Markus S. Miettinen,^{*g,h*} Luca Monticelli,^{*i*} Jukka Määttä,^{*j*} Vasily S. Oganesyan,^{*a*} O. H. Samuli Ollila,^{*sb*} Joona Tynkkynen,^{*c*} and Sergey Vilov,^{*e*}

Despite the vast amount of experimental and theoretical studies on the binding affinity of cations — especially the biologically relevant Na⁺ and Ca²⁺ — for phospholipid bilayers, there is no consensus in the literature. Here we show that by interpreting changes in the choline headgroup order parameters according to the 'molecular electrometer' concept [Seelig *et al., Biochemistry*, 1987, **26**, 7535], one can directly compare the ion binding affinities between simulations and experiments. Our findings strongly support the view that in contrast to Ca²⁺ and other multivalent ions, Na⁺ and other monovalent ions (except Li⁺) do not specifically bind to phosphatidylcholine lipid bilayers at sub-molar concentrations. However, the Na⁺ binding affinity was overestimated by several molecular dynamics simulation models, resulting in artificially positively charged bilayers and exaggerated structural effects in the lipid headgroups. While qualitatively correct headgroup order parameter response was observed with Ca²⁺ binding in all the tested models, no model had sufficient quantitative accuracy to interpret the Ca²⁺:lipid stoichiometry or the induced atomistic resolution structural changes. All scientific contributions to this open collaboration work were made publicly, using nmrlipids.blogspot.fi as the main communication platform.

1 Introduction

Due to its high physiological importance — nerve cell signalling being the prime example — interaction of cations with phospholipid membranes has been widely studied via theory, simulations, and experiments. The relative ion binding affinities are generally agreed to follow the Hofmeister series $^{1-9}$, however, consensus on the quantitative affinities is currently lacking. Until 1990, the consensus (documented in two extensive reviews^{2,3}) was that while multivalent cations interact significantly with phospholipid bilayers, for monovalent cations (with the exception of Li^+) the interactions are weak. This conclusion has since been strengthened by further studies showing that bilayer properties remain unaltered upon the addition of sub-molar concentrations of monovalent salt^{4,10,11}. Since 2000, however, another view has emerged, suggesting much stronger interactions between phospholipids and monovalent cations, and strong Na⁺ binding in particular^{6–9,12–18}.

The pre-2000 view has the experimental support that (in contrast to the significant effects caused by any multivalent cations) sub-molar concentrations of NaCl have a negligible effect on phospholipid infrared spectra⁴, area per molecule¹⁰, dipole potential¹⁹, lateral diffusion¹¹, and choline head group order parameters²⁰; in addition, the water sorption isotherm of a NaClphospholipid system is highly similar to that of a pure NaCl solution — indicating that the ion–lipid interaction is very weak⁴.

The post-2000 'strong binding' view rests on experimental and above all simulational findings. At sub-molar NaCl concentrations, the rotational and translational dynamics of membrane-embedded fluorescent probes decreased^{7,9,12}, and atomic force microscopy (AFM) experiments showed changes in bilayer hard-



 ^a School of Chemistry, University of East Anglia, Norwich, NR4 7TJ, United Kingdom
 ^b Department of Neuroscience and Biomedical Engineering, Aalto University, Espoo, Finland

^c Tampere University of Technology, Tampere, Finland

 $^{^{}d}$ University of Helsinki, Helsinki, Finland

^e Univ Lyon, Université Claude Bernard Lyon 1, CNRS, Institut Lumiére Matiére, F-69622, LYON, France

^f Institute of Organic Chemistry and Biochemistry, Czech Academy of Sciences, Flemingovo nám. 2, 16610 Prague 6, Czech Republic, Charles University in Prague, Faculty of Mathematics and Physics, Ke Karlovu 3, 121 16 Prague 2, Czech Republic ^g Fachbereich Physik, Freie Universität Berlin, Berlin, Germany

^h Max Planck Institute of Colloids and Interfaces, Department of Theory and Bio-Systems, Potsdam, Germany

ⁱ Institut de Biologie et Chimie des Protéines (IBCP), CNRS UMR 5086, Lyon, France ^j Aalto University, Espoo, Finland

^{*}Author to whom correspondence may be addressed. E-mail: samuli.ollila@aalto.fi.

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[‡] The authors are listed in alphabetical order.

ness $^{14-18}$; in atomistic molecular dynamics (MD) simulations, phospholipid bilayers consistently bound Na⁺, although the binding strength depended on the model used $^{12,13,21-26}$.

Some observables have been interpreted in favour of both views. For example, as the effect of monovalent ions (except Li⁺) on the phase transition temperature is tiny (compared to the effect of multivalent ions), it was initially interpreted as an indication that only multivalent ions and Li⁺ specifically bind to phospholipid bilayers²; however, such a small effect in calorimetric measurements was later interpreted to indicate that also Na⁺ binds^{8,12}. Similarly, the lack of significant positive electrophoretic mobility of phosphatidylcholine (PC) vesicles in the presence of NaCl (again in contrast to multivalent ions and Li⁺) suggested weak binding of Na^{+1,8,14,15,27}; however, these data were also explained by a countering effect of the Cl^{-} ions^{22,28}. Furthermore, to reduce the area per lipid in scattering experiments, molar concentrations of NaCl were required ¹⁰, indicating weak ion-lipid interaction; in MD simulations, however, already orders of magnitude lower concentrations resulted in Na⁺ binding and a clear reduction of area per lipid 12,23. Finally, lipid lateral diffusion was unaltered by NaCl in noninvasive NMR experiments¹¹; however, as it was reduced upon Na⁺ binding in simulations, the reduced lateral diffusion of fluorescent probes ^{7,9,12} has been interpreted to support the post-2000 'strong binding' view.

In this paper, we set out to solve the apparent contradictions between the pre-2000 and post-2000 views. To this end, we employ the 'molecular electrometer' concept, according to which the changes in the C–H order parameters of the α and β carbons in the phospholipid head group (see Fig. 1) can be used to measure the ion affinity for a PC lipid bilayer $^{20,29-32}$. As the order parameters can be accurately measured in experiments and directly compared to simulations³³, applying the molecular electrometer as a function of cation concentration allows the comparison of binding affinity between simulations and experiments. In addition to demonstrating the usefulness of this general concept, we show that the response of the α and β order parameters to penetrating cations is qualitatively correct in MD simulations, but that in several models the affinity of Na⁺ for PC bilayers is grossly overestimated. Moreover, we show that the accuracy of lipid-Ca²⁺ interactions in current models is not enough for atomistic resolution interpretation of NMR experiments.

This work was done as an Open Collaboration at nmrlipids.blogspot.fi; all the related files³⁴ and almost all the simulation data (https://zenodo.org/collection/user-nmrlipids) are openly available.

2 Results and Discussion

2.1 Background: Molecular electrometer in experiments

The basis for the molecular electrometer is the experimental observation that binding of any charged objects (ions, peptides, anesthetics, amphihiles) on a PC bilayer interface induced systematic changes in the choline α and β segment C–H order parameters^{20,29–32,35–40}. Being systematic, these changes could be employed for determining the binding affinities of the charged objects in question. Originally the molecular electrometer was de-

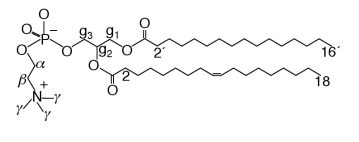


Fig. 1 Chemical structure of 1-palmitoyl-2-oleoylphosphatidylcholine (POPC), and the definition of γ , β , α , g_1 , g_2 and g_3 segments.

vised for cations^{20,29,30}, but further experimental quantification with various positively and negatively charged molecules showed that the choline order parameters S_{CH}^{α} and S_{CH}^{β} in general vary linearly with small amount of bound charge per lipid ^{30–32,35–40}. Let now $S_{CH}^{i}(0)$, where *i* refers to either α or β , denote the order parameter in the absence of bound charge; the empirically observed linear relation can then be written as⁴¹

$$\Delta S_{\rm CH}^{i} = S_{\rm CH}^{i}(X^{\pm}) - S_{\rm CH}^{i}(0) = \frac{4m_{i}}{3\chi}X^{\pm}.$$
 (1)

Here X^{\pm} is the amount of bound charge per lipid, m_i an empirical constant depending on the valency and position of bound charge, and the value of the quadrupole coupling constant $\chi \approx 167$ kHz.

With bound positive charge, the absolute value of the β segment order parameter increases and the α segment order parameter decreases (and *vice versa* for negative charge)^{20,29–32,35,40}. However, as $S_{CH}^{\beta}(0) < 0$ while $S_{CH}^{\alpha}(0) > 0^{42-44}$, both ΔS_{CH}^{β} and ΔS_{CH}^{α} in fact decrease with bound positive charge (and increase with bound negative charge). Consequently, values of m_i are negative for bound positive charges; for Ca²⁺ binding to POPC bilayer (in the presence of 100 mM NaCl), combination of atomic absorption spectra and ²H NMR experiments gave $m_{\alpha} = -20.5$ and $m_{\beta} = -10.0^{30}$. This decrease can be rationalised by electrostatically induced tilting of the choline P–N dipole^{31,32,46} — also seen in simulations^{23,24,47,48} — and is in line with the order parameter increase related to the P–N vector tilting more parallel to the membrane plane seen with decreasing hydration levels⁴⁵.

Quantification of $\Delta S_{\rm CH}^{\alpha}$ and $\Delta S_{\rm CH}^{\beta}$ for a wide range of different cations (aqueous cations, cationic peptides, cationic anesthetics) has revealed that $\Delta S_{\rm CH}^{\beta} / \Delta S_{\rm CH}^{\alpha} \approx 0.5^{-38,40}$. More specifically, the relation $\Delta S_{\rm CH}^{\beta} = 0.43 \Delta S_{\rm CH}^{\alpha}$ was found to hold for DPPC bilayers at various CaCl₂ concentrations²⁰.

2.2 Molecular electrometer in MD simulations

The black curves in Fig. 2 show how the headgroup order parameters for DPPC and POPC bilayers change in H² NMR experiments as a function of salt solution concentration^{20,30}: Only mi-

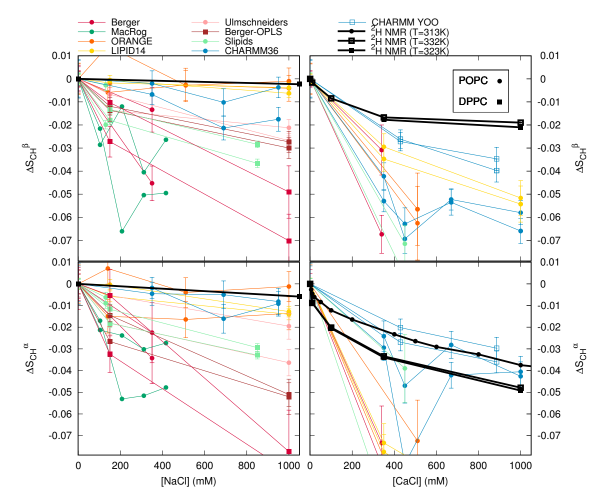


Fig. 2 Changes in the PC lipid headgroup β (top row) and α (bottom) segment order parameters in response to NaCl (left column) or CaCl₂ (right) salt solution concentration increase. Comparison between simulations (Table 1) and experiments (DPPCs from Ref. 20, POPC from Ref. 30). The signs of the experimental values, from experiments without ions^{42–44}, can be assumed unchanged at these salt concentrations^{30,33}. We stress that none of the models reproduces the order parameters without salt within experimental error, indicating structural inaccuracies of varying severity in all of them⁴⁵. Note that the relatively large drop in CHARMM36 at 450 mM CaCl₂ arose from more equilibrated binding due to a very long simulation time, see ESI[†].

nor changes are seen as a function of [NaCl], but the effect of $[CaCl_2]$ is an order of magnitude larger. Thus, according to the molecular electrometer, the monovalent Na⁺ ions have negligible affinity for PC lipid bilayers at concentrations up to 1 M, while binding of Ca²⁺ ions at the same concentration is significant^{20,30}.

Figure 2 also reports order parameter changes calculated from MD simulations of DPPC and POPC lipid bilayers as a function of NaCl or CaCl₂ initial concentrations in solution (for details of the simulated systems see Table 1 and ESI[†]). Note that although none of these MD models reproduces within experimental uncertainty the order parameters for a pure PC bilayer without ions (Fig. 2 in Ref. 45), which indicates structural inaccuracies of varying severity in all models⁴⁵, all the models qualitatively reproduce the experimentally observed headgroup order parameter increase with dehydration⁴⁵. Similarly here (Fig. 2) the presence of cations led to the decrease of S_{CH}^{α} and S_{CH}^{β} , in qualitative agreement with experiments. The changes were, however, overestimated by most models, which according to the molecular electrometer indicates overbinding of cations in most MD simulations.

While the molecular electrometer is well established in experi-

ments (see Sec. 2.1 above), it is not *a priori* clear that it works in simulations. The overestimated order parameter decrease could, in principle, arise from an exaggerated response of the choline headgroups to the binding cations, instead of overbinding. Therefore, to evaluate the usability of the molecular electrometer in MD simulations, we analysed the relation between cation binding and choline order parameter decrease in simulations.

According to the molecular electrometer, the order parameter changes are linearly proportional to the amount of bound cations (Eq. (1)). Figure 3 shows this proportionality in MD simulations (see ESI[†] for the definition of bound ions); in keeping with the molecular electrometer, a roughly linear correlation between bound cation charge and order parameter change was found in all the eight models. Note that quantitative comparison of the proportionality constants (i.e. slopes in Fig. 3) between different models and experimental slopes ($m_{\alpha} = -20.5$ and $m_{\beta} = -10.0$ for Ca²⁺ binding in DPPC bilayer in the presence of 100mM NaCl³⁰) is not straightforward since the simulation slopes depend on the definition used for bound ions (see ESI[†]).

We note that the quantitative comparison of order parameter

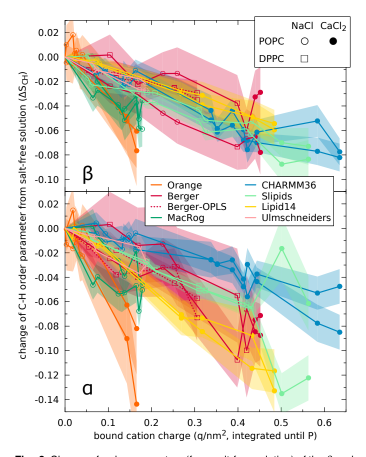


Fig. 3 Change of order parameters (from salt-free solution) of the β and α segments, $\Delta S^{\beta}_{\rm CH}$ and $\Delta S^{\alpha}_{\rm CH}$, as a function of bound cation charge. Eight MD simulation models compared; the two lines per model denote to the two hydrogens per carbon. The order parameters as well as the bound charge calculated separately for each leaflet; cations residing between the bilayer centre and the density maximum of Phosphorus considered bound; error bars (shaded) show standard error of mean over lipids.

changes in response to bound charge should be more straightforward for systems with charged amphiphiles fully associated in the bilayer, as the amount of bound charge is then explicitly known in both simulations and experiments. In such a comparison between experiments^{32,49} and previously published Berger-modelbased simulations⁵⁰, we could not rule out overestimation of order parameter response to bound cations (slopes m_{α} and m_{β}), see ESI[†]. This might, in principle, explain the overestimated order parameter response of the Berger model to CaCl₂, but not to NaCl (see discussion in ESI[†]). Since simulation data with charged amphiphiles are not available for other models, an extended comparison with different models is left for further studies.

Figure 3 shows that the decrease of order parameters clearly correlated with the amount of bound cations in simulations. This is also evident from Fig. 4, which shows the Na^+ density profiles of the MD models ordered according to the order parameter change (in Fig. 2) from the smallest (top) to the largest (bottom). The general trend in the figure is that the Na^+ density peaks are larger for models with larger changes in order parameters, in line with the observed correlation between cation binding and order parameter decrease in Fig. 3.

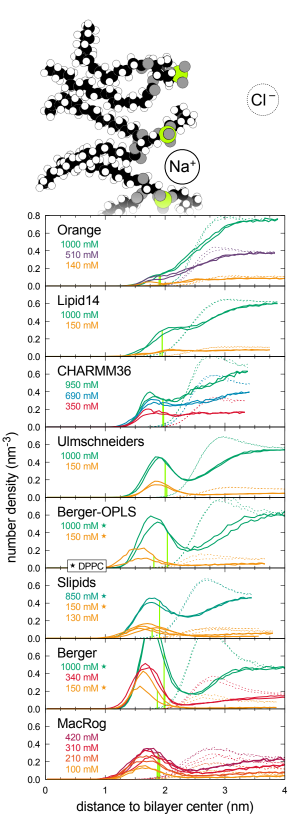


Fig. 4 Na⁺ (solid line) and Cl⁻ (dashed) distributions along the lipid bilayer normal from MD simulations at several NaCl concentrations. The eight MD models are ordered according to their strength of order parameter change in response to NaCl (Fig. 2) from the weakest (top panel) to the strongest (bottom). The light green vertical lines indicate the locations of the Phosphorus maxima, used to define bound cations in Fig. 3.

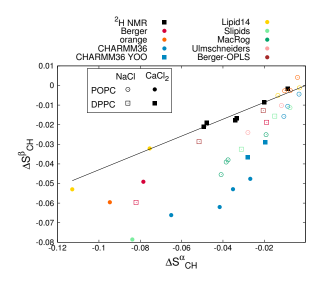


Fig. 5 Relation between ΔS^{β}_{CH} and ΔS^{α}_{CH} from experiments²⁰ and different simulation models. Solid line is $\Delta S^{\beta}_{CH} = 0.43 \Delta S^{\alpha}_{CH}$ determined for DPPC bilayer from ²H NMR experiment with various CaCl₂ concentrations²⁰.

Figure 5 compares the relation between ΔS_{CH}^{β} and ΔS_{CH}^{α} in experiments²⁰ and in MD models. Only Lipid14 gave $\Delta S_{CH}^{\beta} / \Delta S_{CH}^{\alpha}$ ratio in agreement with the experimental ratio; all other models underestimated the α segment order parameter decrease with bound cations with respect to the β segment decrease.

In conclusion, a clear correlation between bound cations and order parameter decrease was observed for all simulation models. Consequently, the molecular electrometer can be used to compare the cation binding affinity between experiments and simulations. However, we found that quantitatively the response of α and β segment order parameters to bound cations in simulations did not generally agree with the experiments; e.g., the $\Delta S^{\beta}_{CH}/\Delta S^{\alpha}_{CH}$ ratio agreed with experiments only in the Lipid14 model (Fig. 5). Thus, the observed overestimation of the order parameter changes with salt concentrations could, in principle, arise from overbinding of cations or from an oversensitive lipid headgroup response to the bound cations (see also discussion in ESI[†]). A careful analysis with current lipid models is performed in the next section.

2.3 Cation binding in different simulation models

The order parameter changes (Fig. 2) and density distributions (Fig. 4) demonstrated significantly different Na⁺ binding affinities in different simulation models. The best agreement with experiments (lowest ΔS^{α}_{CH} and ΔS^{β}_{CH}) was observed for the three models (Orange, Lipid14, CHARMM36; see Fig. 2) that predicted the lowest Na⁺ densities near the bilayer (Fig. 4). All the other models clearly overestimated the choline order parameter responses to NaCl (Fig. 2) — and notably the strength of the overestimation was clearly linked to the strength of the Na⁺ binding affinity (compare Figs. 2 and 4), which leads us to conclude that Na⁺ binding affinity was overestimated in all these models.

As in the best three models the order parameter changes with NaCl were small (< 0.02), the achieved statistical accuracy did not allow us to conclude which of the three had the most realistic Na⁺

binding affinity, especially at physiological NaCl concentrations (\sim 150 mM) relevant for most applications. The overestimated binding in the other models raises questions concerning the quality of predictions from these models when NaCl is present. Especially interactions between charged molecules and the bilayer might be significantly affected by the strong Na⁺ binding, which gives the otherwise neutral bilayer an effective positive charge.

Significant Ca²⁺ binding affinity for phosphatidylcholine bilayers at sub-molar concentrations is agreed on in the literature ^{2,3,20,30}, however, several details remain under discussion. Simulations suggest that Ca²⁺ binds to lipid carbonyl oxygens with a coordination number of 4.2¹³, while interpretation of NMR and scattering experiments suggest that one Ca²⁺ interacts mainly with the choline groups ^{106–108} of two phospholipid molecules ³⁰. A simulation model correctly reproducing the order parameter changes would resolve the discussion by giving atomistic resolution interpretation for the experiments.

As a function of CaCl₂ concentration, all models but one (CHARMM36 with the recent heptahydrated Ca^{2+} by Yoo et al.⁷⁶) overestimated the order parameter decrease (Fig. 2), which according to the molecular electrometer indicates too strong Ca²⁺ binding. (We note that while this is the most likely scenario for the models that overestimated changes in both order parameters, for CaCl₂ it is possible also that the headgroup response is oversensitive to bound cations, see ESI[†].) In CHARMM36 with the heptahydrated Ca²⁺ by Yoo et al. ⁷⁶, ΔS^{β}_{CH} was overestimated but ΔS^{α}_{CH} underestimated (Fig. 2), in line with the $\Delta S^{\beta}_{CH}/\Delta S^{\alpha}_{CH}$ ratio in CHARMM36 being larger than in experiments (Fig. 5). As we do not know whether ΔS_{CH}^{β} or ΔS_{CH}^{α} was more realistic, we cannot conclude whether Ca^{2+} binding was too strong or too weak in CHARMM36. This could be resolved by comparing against experimental data with a known amount of bound charge (e.g., amphiphilic cations^{32,49}), however, such simulation data are not currently available.

The density distributions with CaCl₂ showed significant Ca²⁺ binding in all models (Fig. 6), however, some differences occurred in details. The Berger model predicted deeper penetration (density maximum at ~ 1.8 nm) compared to other models (~ 2 nm); the latter value is probably more realistic as ¹H NMR and neutron scattering data indicate that Ca²⁺ interacts mainly with the choline group^{2,106–108}. In CHARMM36 (but not in Slipids) practically all Ca²⁺ ions present in the simulation bound the bilayer within 2 μ s (Fig. 6 and ESI[†]), which hints that the Ca²⁺ binding affinity of CHARMM36 is among the strongest of these models.

The origin of inaccuracies in lipid–ion interactions and binding affinities is far from clear. Potential candidates are, e.g., discrepancies in the ion models^{109–111}, incomplete treatment of electronic polarizability¹¹², and inaccuracies in the lipid headgroup description⁴⁵.

Considering the ion models, Cordomi et al.²⁴ showed the Na⁺ binding affinity to decrease when ion radius is increased; however, in their DPPC bilayer simulations (with the OPLS-AA force field¹¹³) even the largest Na⁺ radii still resulted in significant binding. In our results, the Slipids force field gave essentially similar binding affinity with ion parameters from Refs. 88 and **Table 1** List of MD simulations. The salt concentrations calculated as [salt]= $N_c \times [water] / N_w$, where [water] = 55.5 M; these correspond the concentrations reported in the experiments by Akutsu et al.²⁰. The lipid force fields named as in our previous work⁴⁵.

force field for lipids / ions	lipid	salt	[salt] (mM)	$^{a}N_{1}$	^b N _w	^c N _c	^d T (K)	^e t _{sim} (ns)	^f t _{anal} (ns)	^g files
Berger-POPC-07 ⁵¹ / –	POPC	no	0	128	7290	0	298	270	50	52
Berger-POPC-07 ⁵¹ / ffgmx ⁵³	"	NaCl	340	"	7202	44	"	110	"	54
Berger-POPC-07 ⁵¹ / ffgmx ⁵³	"	$CaCl_2$	340	"	7157	"	"	108	58	55
Berger-DPPC-97 ⁵⁶ / –	DPPC	no	0	72	2880	0	323	60	50	57
Berger-DPPC-97 ⁵⁶ / ffgmx ⁵³	"	NaCl	150	"		8	"	120	60	58
Berger-DPPC-97 ⁵⁶ / ffgmx ⁵³	"	"	1000	"	2778	51	"	"	"	59
BergerOPLS-DPPC-06 ⁶⁰ / –	DPPC	no	0	72	2880	0	323	120	60	61
BergerOPLS-DPPC-06 ⁶⁰ / OPLS ⁶²	"	NaCl	150	, <u> </u>		8	"		"	63
BergerOPLS-DPPC-06 ⁶⁰ / OPLS ⁶²	"	"	1000	"	2778	51	"	"	"	64
CHARMM36 ⁶⁵ / –	POPC	no	0	128	5210	0	303	200	150	66
CHARMM36 ⁶⁵ / $-$	"	"	0	72	2242	"	"	30	20	67
CHARMM36 ⁶⁵ / CHARMM36 ⁶⁸	"	NaCl	350	, <u> </u>	2085	13	"	80	60	69
CHARMM36 ⁶⁵ / CHARMM36 ⁶⁸	"	"	690	"		26	"	73	"	70
CHARMM36 ⁶⁵ / CHARMM36 ⁶⁸	"	"	950	"	2168	37	"	80	"	71
CHARMM36 ⁶⁵ / CHARMM36	"	$CaCl_2$	350	128	6400	35	"	200	100	72
CHARMM36 ⁶⁵ / CHARMM36	"	"	450	200	9000	73	310	2000	"	73
CHARMM36 ⁶⁵ / CHARMM36	"	"	670	128	6400	67	303	200	120	74
CHARMM36 ⁶⁵ / CHARMM36	"	"	1000	"	"	100	"	"	100	75
CHARMM36 ⁶⁵ / –	DPPC	no	0	128	8000	0	323	170	150	_
CHARMM36 ⁶⁵ / Yoo ⁷⁶	"	$CaCl_2$	430	"	7760	60	"	200	170	_
CHARMM36 ⁶⁵ / Yoo ⁷⁶	"	"	890	"	7520	120	"	"	"	_
MacRog ⁷⁷ / –	POPC	no	0	128	6400	0	310	400	200	78
MacRog ⁷⁷ / –	"	"	0	288	14400	"	"	90	40	79
MacRog ⁷⁷ / OPLS ⁶²	"	NaCl	100	"	14554	27	"	"	50	80
MacRog ⁷⁷ / OPLS ⁶²	"	"	210	"	14500	54	"	"	"	"
MacRog ⁷⁷ / OPLS ⁶²	"	"	310	"	14446	81	"	"	"	"
MacRog ⁷⁷ / OPLS ⁶²	"	"	420	"	14392	108	"	"	"	"
Orange / –	POPC	no	0	72	2880	0	298	60	50	81
Orange / OPLS ⁶²	"	NaCl	140	"	2866	7	"	120	60	82
Orange / OPLS ⁶²	"	"	510	"	2802	26	"	"	100	83
Orange / OPLS ⁶²	"	"	1000	"	2780	50	"	"	80	84
Orange / OPLS	"	$CaCl_2$	510	"	2802	26	"	"	60	85
Slipids ⁸⁶ / –	POPC	no	0	128	5120	0	310	200	150	87
Slipids ⁸⁶ / AMBER ⁸⁸	"	NaCl	130	200	9000	21	"	105	100	89
Slipids ⁸⁶ / AMBER ⁶²	"	$CaCl_2$	450	"	"	73	"	2000	"	90
Slipids ⁹¹ / –	DPPC	no	0	128	3840	0	323	150	100	92
Slipids ⁹¹ / AMBER ^{93,94}	"	NaCl	150	600	18000	49	"	100	40	-
Slipids ⁹¹ / AMBER ^{93,94}	"	"	850	128	3726	57	"	205	200	95
Slipids ⁹¹ / AMBER ^{93,94}	"	"	1750	"	3612	114	"	105	100	"
Slipids ⁹¹ / AMBER ^{93,94}	"	"	2570	"	3514	163	"	"	"	"
Lipid14 ⁹⁶ / –	POPC	no	0	128	5120	0	298	205	200	97
Lipid14 ⁹⁶ / AMBER ⁶²	"	NaCl	150	"	"	12	"	"	"	98
Lipid14 ⁹⁶ / AMBER ⁶²	"	"	1000	"	"	77	"	"	"	99
Lipid14 ⁹⁶ / AMBER ⁶²	"	$CaCl_2$	350	"	6400	35	"	200	100	100
Lipid14 ⁹⁶ / AMBER ⁶²	"	"	1000	"	"	100	"	"	"	101
Ulmschneiders ¹⁰² / –	POPC	no	0	128	5120	0	298	2×205	2×200	103
Ulmschneiders $\frac{102}{100}$ / OPLS $\frac{62}{100}$	"	NaCl	150	"	"	12	"	205	200	104
Ulmschneiders ¹⁰² / OPLS ⁶²	"	"	1000	"	"	77	"	"	"	105

a Number of lipid molecules

b Number of water molecules

c Number of cations

d Simulation temperature

e Total simulation time

 $f {\rm Time}$ used for analysis

g Reference for simulation files

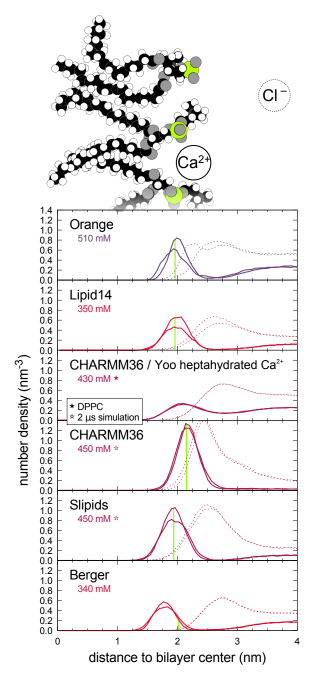


Fig. 6 Ca²⁺ (solid line) and Cl⁻ (dashed) distributions along the lipid bilayer normal from MD simulations. For clarity, only one CaCl₂ concentration per MD model is shown; see ESl[†] for a plot including all the available concentrations. The light green vertical lines indicate the locations of the Phosphorus maxima, used to define bound cations in Fig. 3.

93,94 (Fig. 4). Further, compensation of missing electronic polarizability by scaling the ion charge ^{112,114} reduced Na⁺ binding in Berger, Berger-OPLS and Slipids, but not enough to reach agreement with experiments (ESI[†]). The charge-scaled Ca²⁺ model¹¹⁵ slightly reduced binding in CHARMM36, but did not have significant influence in Slipids (ESI[†]). The heptahydrated Ca²⁺ ions by Yoo et al.⁷⁶ significantly reduced Ca²⁺ binding in CHARMM36 (Fig. 6), however, the model must be further analysed to fully interpret the results.

The lipid models may also have a significant influence on ion binding behaviour. For example, the same ion model and nonbonded parameters are used in Orange and Berger-OPLS⁶⁰, but while Na⁺ ion binding affinity appeared realistic in Orange, it was significantly overestimated in Berger-OPLS (Fig. 4). However, realistic Na⁺ binding does not automatically imply realistic Ca²⁺ binding (see Orange, Lipid14, and CHARMM36 in Fig. 2) or realistic choline order parameter response to bound charge (see Orange and CHARMM36 in Fig. 5). It should also be noted that the low binding affinity of Na⁺ in CHARMM36 is due to the additional repulsion (NBFIX⁶⁸) added between the sodium ions and lipid oxygens (ESI^{\dagger}), and that in the Ca²⁺ model by Yoo et al.⁷⁶ the calcium is forced to be solvated solely by water. Altogether, our results indicate that probably both, lipid and ion force field parameters, need improvement to correctly predict the cation binding affinity, and the associated structural changes.

3 Conclusions

In accordance with the molecular electrometer^{20,29–32}, cation binding to lipid bilayers was accompanied with a decrease in the C–H order parameters of the PC head group α and β carbons in all the simulation models tested (Fig. 3) — despite of the known inaccuracies in the actual atomistic resolution structures⁴⁵. Hence, the molecular electrometer allowed a direct comparison of Na⁺ binding affinity between simulations and noninvasive NMR experiments. The comparison revealed that most models overestimated Na⁺ binding; only Orange, Lipid14, and CHARMM36 predicted realistic binding affinities. None of the tested models had the accuracy required to interpret the Ca²⁺:lipid stoichiometry or the induced structural changes with atomistic resolution.

Taken together, our results corroborate the pre-2000 view that at sub-molar concentrations, in contrast to Ca²⁺ and other multivalent ions $^{1\!-\!4,10,11,19,20,27,30},$ Na^+ and other monovalent ions (except Li⁺) do not specifically bind to phospholipid bilayers. Concerning the interpretation of existing experimental data, our work supports Cevc's view² that the observed small shift in phase transition temperature is not indicative of Na⁺ binding. Further, our findings are in line with the noninvasive NMR spectroscopy work of Filippov et al.¹¹ that proved the results of Refs. 7,9,12 to be explainable by direct interactions between Na⁺ ions and fluorescent probes. Finally, as spectroscopic methods are in general more sensitive to atomistic details in fluid-like environment than AFM, our work indirectly suggests that the ion binding reported from AFM experiments on fluid-like lipid bilayer systems¹⁴⁻¹⁸ might be confounded with other physical features of the system. Concerning contradictions in MD simulation results, we reinterpret the strong Na⁺ binding as an artefact of several simulation models, e.g., the Berger model used in Refs. 12,13.

The artificial specific Na⁺ binding in MD simulations may lead to doubtful results, as it effectively results in a positively charged phosphatidylcholine lipid bilayer even at physiological NaCl concentrations. Such a charged bilayer will have distinctly different interactions with charged objects than what a (more realistic) model without specific Na⁺ binding would predict. Furthermore, the overestimation of binding affinity may extend from ions to other positively charged objects, say, membrane protein segments. This would affect lipid–protein interactions and could explain, for example, certain contradicting results on electrostatic interactions between charged protein segments and lipid bilayers^{116,117}. In conclusion, more careful studies and model development on lipid bilayer–charged object interactions are urgently called for to make molecular dynamics simulations directly usable in a physiologically relevant electrolytic environment.

This work was done as a fully open collaboration, using nmrlipids.blogspot.fi as the communication platform. All the scientific contributions were communicated publicly through this blog or the GitHub repository³⁴. All the related content and data are available at Ref. 34.

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