Accurate binding of calcium to phospholipid bilayers by effective inclusion of electronic polarization

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1.Abstract directly from Joe's conference abstracts. To be rewritten. Classical molecular dynamics simulations give detailed information about membrane structure and dynamics. However, there is still a room for improvements in current force fields it is known from the literature, that the binding of ions, especially cations, to phopholipid membranes is overestimated in all classical models [1]. We suggest that the membrane-ion interactions can be corrected by including implicit electronic polarizability into the lipid models through the electronic continuum correction (ECC) [2], which was already applied to monovalent and divalent ions yielding models that feature correct ion pairing [3]. Using the electrometer concept [3, 4] and x-ray scattering form factors, our simulations point out that our hypothesis is correct and ECC is indeed a missing important contribution in current classical lipid models. Moreover, the solid physical principles behind ECC are found not to hamper other relevant properties of a phospholipid bilayer. The new lipid model, "ECC-lipids", shows accurate binding affinity to sodium and calcium cations and headgroup order parameter response to bound charge. We also provide for the first time a realistic stochiometry of bound calcium cations to a POPC membrane, and their binding sites. This work will continue as an open collaboration project NMRlipids IV (http://nmrlipids.blogspot.fi).

I. INTRODUCTION

Cation interactions with cellular membranes play a key role in several biological processes, like in signal propagation in neurons and vesicle fusion. Cation interactions with lipid bilayers serving as a model for cellular membranes are thus widely studied with experimental [1? -4] and theoretical methods [5?]. General conclusion from experimental studies has been that multivalent ions and lithium have weak specific binding in phopholipid bilayers, while other monovalent ions do not essentially bind [1-3, 6]. The presence of anoinic lipids, like PS or PG, increase the concentration close to the bilayer and thus the amount of bound ions, but do not affect the specific binding constant [6]. The binding details, like binding sites and stoichiometry are not yet fully resolved but interpretation of NMR and scattering experiments suggest that one Ca2+ interacts mainly with the choline groups [7–9] of two phospholipid molecules [10].

Classical molecular dynamics simulations have potential to solve the ion binding details in lipid bilayers and reveal its relevance to various biological problems [11?, 12]. However, the available molecular dynamics simulation models have a strong tendency to overestimate cation binding on zwitterionic bilayers and none of them can reproduce the experimental data with the accuracy required for the interpretation of experiments [13]. The overestimated specific cation binding also makes zwitterionic bilayers effectively positively charged, which could potentially lead to significant artefacts in applications. Thus, there is a vast demand for classical molecular dynamics simulation model which correctly reproduces cation binding in lipid bilayers.

The lack of electronic polarizability from the classical MD simulation models is a potential source of artefact, which could lead to overbinding of cations. The issue has been considered higly relevant since the early days of lipid bilayer simulations and pioneering simulation studies scaled the partial charges to effectively include electronic polarizability [14?]. The explicit inclusion of electronic polarizability has been, however, turned to be practically complicated and implementations for lipids are rare [15].

In this work we show that the cation overbinding in classical MD simulations can be corrected by including electronic polarizability by using effective continuum correction (ECC) [16] for polar region of zwitterionic lipid molecules. This is essentially physicalle well justified version of partial charge scaling implemented in early days of lipid and surfictant simulations [14?]. The approach has been previously shown in to improve bulk performace of ion models against neutron scattering data [17?, 18]. The better bulk behaviour was not, however, sufficient to correct binding in lipid bilayers [13].

II. METHODS

A. Electronic continuum correction for lipid bilayers

According to the electronic continuum correction (ECC) the electronic polarizability can be effectively included in MD simulations by scaling the partial charges with a constant factor $f_q = \epsilon_{el}^{-1/2}$, where ϵ_{el} is the electronic part of the dielectric constant of media [16]. Measurements of high frequency dielectric constant gives values of approximately $\epsilon_{el} \approx 2$ for almost any biomaterial [16?]. The value measured for water, $\epsilon_{el} = 1.78$, gives a scaling factor of $f_q = 0.75$ [16?], which has been successfully used to improve the performance of ion

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force fields [17?, 18].

Here we apply the same approach to lipid bilayers to accurately describe the lipid headgroup response to Na+ and Ca2+ concentrations [13]. We use the Lipid14 [19] force field parameters as a starting point, because they give the most realistic headgroup response with added cations and relatively realistic glycerol backbone and headgroup structures when compared with other state of the art lipid models [13, 20]. The partial atomic charges in Lipid14 were derived by fitting the electrostatic potential to its model quantum chemistry representation in vacuum. 2.I do not fully understand the end of the paragraph If such charges are obtained in an implicit solvent, they vary the most for the polar moieties [?]. In order to represent this difference in polarity in a model with fixed charges, we shall use the average charges from both environments (IpolQ charges) [?]. By taking the charges of oxygen atoms in vacuum and implicit water solvent for POPC from [21], we represent the IpolQ charges in the electronic continuum correction of the Lipid14 model by increasing the scaling factor for charges to $f_q = 0.8$. The newly presented ECC-POPC model is based on the POPC model from

Hydrocarbon chains in Lipid14 and other lipids models are highly optimized and give generally a good description for hydrophobic part of lipid bilayers in various conditions [22], in contrast to glycerol backbone and headgroup regions which require some improvement in all available lipid models [20]. To minimize the intercalation with the highly optimized hydrocarbon parameters, we decided to apply ECC correction only to the headgroup, glycerol backbone and carbonyl in acyl chains. These regions are also most polar parts in lipids and are expected to most affect on cation binding.

The scaling of these partial charges by a factor of 0.8 reduce the area per molecule to ??, which is significantly smaller than the experimental and original Lipid 14 values. The decrease of area was speculated 3.Did you analyze the effect of this to hydration or did we only speculate? to arise from reduced hydration of headgroup due to the lower polarity of molecules with scaled charges. The hydration can be increased by decreasing the radius of atoms by reducing σ term in Lennart-Jones potential, which allows water molecules approach closer to lipid atoms and have stonger electrostatic interactions with them. 4.This effect may have an official name. In that case we should mention it. Indeed, the scaling of σ increased the area per molecule and the results close to experimental and original Lipid14 values were achieved with scaling factor of ?? for σ and 0.8 for partial charges.

B. Comparison to experimental data

Structures sampled by individual lipid molecules were compared to experimental data by using C-H bond order parameters [22]

$$S_{\rm CH} = \frac{3}{2} \langle \cos^2 \theta - 1 \rangle,\tag{1}$$

where θ is the angle between C-H bond and membrane normal and average is taken over all sampled configurations. Bilayer

dimensions were compared to experiments by using the scattering form factor [22] 5.Add equation.

Ion binding was compared between experiments and simulations by using lipid headgroup order parameters and "electrometer concept" introduced by Seelig et al. [13, 23]. The concept is based on the experimental observation that the order parameters of α and β carbons in lipid headgroup (see Fig. ??) are proportional to the amount of bound charge in lipid bilayer [23]. More recent analysis included also the order parameter signs and concluded that the order parameters decrease with bound positive charge and increase with bound negative charge [13, 22]. The observations are rationalized as a change of lipid headgroup dipole tilt to more vertical orientation with bound positive charge and *vice versa* for negative [23].

6.ongoing,Actual concentration of cations in simulation has yet to be estimated. If it varies too much from the nominal concentration, I may need to tweak the scaling factors, f_q or only f_σ , to accomodate it. However, it is very unlikely, response to the surfactant DHMDMAB is OK. Big patches with loads of solvent are running at the moment to guide me on the possible finite-size errors and this conc-error. SAMULI: I thought about this a little bit. My current conclusion is that we can take the number concentration from simulations with large box having a clear bulk concentration region and calculate the bulk concentration in mol/I unit like this

$$C_{eq}[mol/l] = \frac{C_{plateau}[mM]}{0.602}$$
 (2)

C. Simulation details

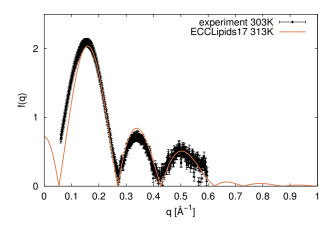
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III. RESULTS AND DISCUSSION

A. Lipid bilayers without ions

The scattering form factors, NMR order parameters and area per lipids calculated from the ECC corrected lipid model for POPC are compared to experiments and original Lipid14 results in Fig. 1 and in Table I. The structural quality is comparable to the state of art lipid models available in literature [22], thus we conclude that the ECC corrected lipid model reproduces the lipid bilayer structure in liquid disordered phase with similar accuracy than other available models 8.Discussion to be finished when we have all the results in the figure. 11.finalize figure (NMR headgr. OPs + SAXS, continue in the discussion.

Headgroup order parameter response to bound charge was evaluated against experimental data measured with cationic surfactant (dihexadecyldimethylammonium bromide, $C_{12}Cl_{16}^{+}N2C_{1}Br^{-}$) [25]. The exact amount of bound charge in the membrane is known these systems, because practically all cationic surfactant molecules are embedded in lipid bilayer due to their amphiphilic nature. Thus, the headgroup order parameter changes as a function of mole fraction of cationic surfactants in Fig. 2 gives also the order parameter changes as a function of bound cations. 13.I think that we should make a separate figures for this one and ion concentrations. In the current figure it would be



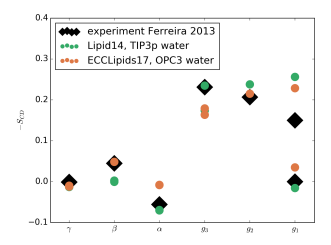


FIG. 1: X-ray scattering form factors from experiments [?] and simulations using Lipid14 and ECCLipids17 models. Headgroup and glycerol backbone order parameters from simulations with Lipid14 [19] and EECLipid17 models compared with experimental order parameters from [24].

9.Add acyl chain order parameters, POPC chemical structure 10.Should we add the results from original lipid14?

TABLE I: Area per lipid from different models for POPC without ions

model	$A(^2)$	Temperature [K]
Lipid14 (literature)	$65.6 \!\pm 0.5$	303
Lipid14ecc0.80+sigma0.875		313
GMX small patch	64.9	
GMX 4xbig patch	65.5	
oMM small patch	63.65	
oMM 4xbig patch	63.7	
experiment [?] 12.REF	62.7	293
experiment	64.3	303
experiment	67.3	323
experiment	68.1	333
experiment POPE	56.6	303

difficult to show this with a reasonable scale. Currently Lipid14 results are cut out. thus the scale should be increased. This would, however, make the changes with CaCl too small. The change of headgroup tilt could be incorporated in the same figure with order parameter response to cationic surfactant. The headgroup order parameter response to bound cation concentration is approximately linear up to ~ 0.3 mole fraction in experiments [25]. The linearity is also observed in simulations with original Lipid14 and with ECC correction. Quantitative comparison, however, reveals that the response is overestimated in original Lipid14 for both segments, while the ECC corrected model gives a good agreement for the change of α segment order parameter, but slightly underestimates the β segment order parameter change. The overestimation of order parameter changes with original Lipid14 model suggests that the overestimated response with CaCl₂ concentration in [13] can be partly explained by the high sensitivity of the headgroup order parameter response to bound charge.

Secondly, the S^{α}/S^{β} ratio of the response is found to be slightly larger than in experiments (VALexp vs VALsim 14.Add values of S^{α}/S^{β} response for sim and experiment SAMULI: Maybe these could be put in the same figure with the order parameter response and P-N vector angle change with surfactants.). This property is kept also by the newly derived model, ECC-lipid, and, hence, it shows a perfect agreement in the response of the headgroup order parameter S^{α} but underestimates slightly the response of S^{β} .

B. Cation binding in POPC bilayer

Headgroup order parameter response to increasing CaCl₂ concentration from experiments, original Lipid14 model and ECC corrected model are shown in Fig. 2. The order parameter response is significantly overestimated in original Lipid14 model, while results from ECC corrected model are in good agreement with experiments. This is a significant improvement over previously available models, which always overstimate the order parameter response to CaCl₂ concentration [13]. The good agreement with experiments indicate that the binding details of Ca2+ are realistic in the ECC corrected model and it can be thus used to study lipid-ion interaction details.

Ion density profiles between different simulation models are compared in Fig. 3. Density profiles from simulations with original Lipid14 and Dang ions [30–32] show a pronounced peak in the position of the phosphate moieties of POPC. The use of a ECC-ion model [17, 18] along with original Lipid14 does not significantly change it 16.If we show density profile for this, we should show also the order parameter changes. The new ECC-lipid model with scaled ions exhibits on the other hand smaller density in this region suggesting overall weaker binding of cations (Fig. 2). This demonstrates that cation binding in zwitterionic phospholipid bilayer can be accurately described with classical MD simulation model with effective included of electronic polarizability.

One possible example is to adress the discourse on the stoichiometry of Ca^{2+} binding to POPC [?] 19.REFs. In line with the early experimental finding [?], we find that our data fit well a ternary complex model, which assumes 2 POPC

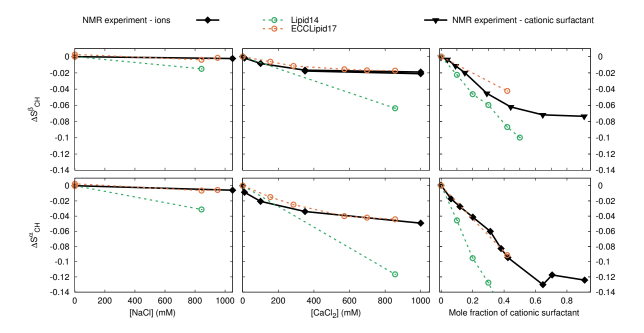


FIG. 2: Headgroup order parameter changes as a function of NaCl, $CaCl_2$ concentration and cationic surfactant (dihexadecyldimethylammonium bromide, $C_{12}Cl_{16}^+N2C_1Br^-$) from simulations and experiments (DPPC [26], POPC [10], surfactant [25]). Simulations with Lipid14 and qvist ion model from [13, 27–29].

15.Add Lipid14-Aquist data. Lipid14/qvist data to be added from

https://github.com/NMRLipids/lipid_ionINTERACTION/blob/master/Data/POPC/CaCl/LIPID14/LIPID14caclCONSchange.dat. Joe: I'm not sure what is meant – I think Aquist data are actually plotted there. I recently changed it to L14+Dang ions (both data and plot). I think just one ion model is sufficient (and ECC-ions are based on Dang model). Samuli: I think that we could add also the Aqvist data. This is in the NMRlipids II publication so it might make easier to follow for people who have red it publication.

molecules per 1 Ca²⁺. In contrast to this experiment, however, we possess also a complete atomistic detail of this phenomenon. Hence we can support this empirical finding with a direct observation of transient complexes of Ca²⁺ and POPC. The simulation suggests that the stoichiometry 2 POPC:1 Ca²⁺ is the most common complex. In addition, there also is a nonnegligable probability of one Ca²⁺ cross-bridging three POPC molecules. This cannot be observed in experiments without atomistic detail as it makes only a small perturabation to the binding isotherm that assumes 2 POPC per 1 Ca²⁺. 20.Analyze stoichiometry for Na⁺, Ca²⁺, their interaction energies with the lipid membrane, etc, and finalize the discussion after these results.

It is also suggested that the addition of NaCl to the solution of $CaCl_2$ enhances the hedgroup order parameter response compared to the solution with only $CaCl_2$. [?] 21.Simulate this effect and discuss it further

22.The difference between DPPC and POPC – simulate and compare with experiment

Discussion:

23.It might be worth acknowledging each experimental finding in [?] and observations in [?].

other lipids: charged lipids? – ongoing research in our lab. I'm currently working on POPE with Aniket for curved (and flat) membranes.

Samuli: From the point of view of this paper, the most relevant other lipid to study would be DPPC to see if we can

reproduce the difference between POPC and DPPC in experiments

Joe: In addition, there are more experimental OP data on DPPC. However, it is not necessary and we can leave it to the community project.

Role of water model: we use OPC3 (current best), it would be worth giving an estimate how results change when we use say SPCE or even TIP3p at least in the SI (so that the reader knows what errors to expect comming from these sub-optimal models). In addition, there is protein force field Amber15-FB, which uses water close to OPC3, TIP3pFB 24.REFs. On the other hand, it might be the best if we used a model that doesn't have the dielectric constant from nuclei 78, but rather 44 – TIP4p2005 is the closest. [? ?]

questionable charge distribution from RESP – a leeway for further tweaks of the FF. It is not obvious that RESP charges provide the best description, especially due to its non-uniqe solution. Shall we solve RESP fitting with the constraint of full charges and then scale down, or shall we rather solve the fitting with a scaled total charge target?

acheivable accuracy of the MD engines themselves (mainly Hector's worry) is another limiting factor in fine tuning parameters – solid physical ground helps.

application of the correction to other lipid models: The rule looks general, however, it depends on how accurate the original model was. From preliminary simulations with POPE, it

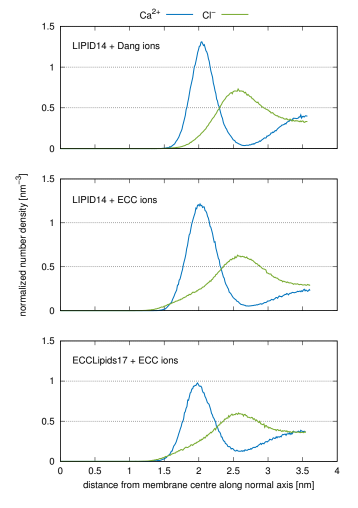


FIG. 3: Density profiles of Ca^{2+} and Cl^- for Lipid14 model with Aquist parameters and with ECC ions and ECCLipids17 with ECC ions.

17.We should add the location of bilayer here somehow. In NMRlipids II the location of phosphate was shown with green vertical line. 18.I would add Aqvist data from NMRlipids II in here as well.

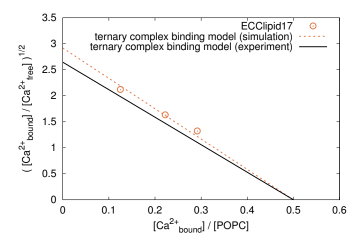


FIG. 4: Binding isotherm assuming stoichiometry of 2 POPC:1 Ca²⁺ as used in [?] fits the simulation data nicely.

looks that the rule works for Lipid14 FF, at least for zwitterionic headgroups.

IV. CONCLUSION

We present models of POPC and DPPC lipids that for the first time exhibit accurate headgroup order parameter response to cation binding. The models are derived from Lipid14 model [19] by applying the electronic continuum correction. The models were optimized to represent correct membrane structure, and they were validated with the use of electrometer concept [23, 33?]. Compared to the phenomenological observations of calcium:POPC stoichiometry in the experimental work [?], our simulations reveal the same stoichiometry of 2 POPC molecules per 1 calcium but through direct observation. This tells us on the subtle effects of calcium in phospholipid membranes possibly leading to a better understanding of their physical properties (elasticity etc.). 25.Improve this concluding-discussion with actual insights.

This will be a foundation stone of a new open-collaboration project NPRlipid 6 in nmrlipids.blogspot.fi....

Acknowledgments

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9. Add acyl chain order parameters, POPC chemical

10. Should we add the results from original lipid14? . .

12. put original references, not Slipids param. paper. .

SUPPLEMENTARY INFORMATION

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[14]	Ollila, et al., Phys. Chem. Chem. Phys. 18 (2016). B. Jonsson, O. Edholm, and O. Teleman, J. Chem. Phys. 85 , 2259 (1986).	2. I do not fully understand the end of the paragraph
[15]	J. Chowdhary, E. Harder, P. E. M. Lopes, L. Huang, A. D. MacKerell, and B. Roux, J. Phys. Chem. B 117, 9142 (2013).	4. This effect may have an official name. In that case
[16]	I. Leontyev and A. Stuchebrukhov, Phys. Chem. Chem. Phys. 13, 2613 (2011).	5. Add equation
	M. Kohagen, P. E. Mason, and P. Jungwirth, J. Phys. Chem. B 118 , 7902 (2014).	6. ongoing, Actual concentration of cations in simulation has yet to be estimated. If it varies too much
[18]	M. Kohagen, P. E. Mason, and P. Jungwirth, J. Phys. Chem. B 120 , 1454 (2016).	from the nominal concentration, I may need to tweak the scaling factors, f_q or only f_{σ} , to accommodate it.
[19]	C. J. Dickson, B. D. Madej, . A. Skjevik, R. M. Betz, K. Teigen, I. R. Gould, and R. C. Walker, J. Chem. Theory Comput. 10, 865 (2014).	However, it is very unlikely, response to the surfactant DHMDMAB is OK. Big patches with loads of solvent
[20]	A. Botan, F. Favela-Rosales, P. F. J. Fuchs, M. Javanainen, M. Kanduč, W. Kulig, A. Lamberg, C. Loison, A. Lyubartsev,	are running at the moment to guide me on the possible finite-size errors and this conc-error. SAMULI: I thought about this a little bit. My current conclusion is
[21]	M. S. Miettinen, et al., J. Phys. Chem. B 119 , 15075 (2015). A. Maciejewski, M. Pasenkiewicz-Gierula, O. Cramariuc, I. Vattulainen, and T. Rog, J. Phys. Chem. B 118 , 4571 (2014).	that we can take the number concentration from simulations with large box having a clear bulk concentration
[22]	O. S. Ollila and G. Pabst, Atomistic resolution structure and dynamics of lipid bilayers in simulations and experiments	region and calculate the bulk concentration in mol/l unit like this
	(2016), in Press, URL http://dx.doi.org/10.1016/j.bbamem.2016.01.019.	7. To be written
	J. Seelig, P. M. MacDonald, and P. G. Scherer, Biochemistry 26 , 7535 (1987).	in the figure
[24]	T. M. Ferreira, F. Coreta-Gomes, O. H. S. Ollila, M. J. Moreno, W. L. C. Vaz, and D. Topgaard, Phys. Chem. Chem. Phys. 15 ,	11. finalize figure (NMR headgr. OPs + SAXS, continue in the discussion

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should be increased. This would, however, make the	21. Simulate this effect and discuss it further 4			
changes with CaCl too small. The change of headgroup	21. Simulate this effect and disease it farther			
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figure with the order parameter response and P-N vector	23. It might be worth acknowledging each experimental			
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so it might make easier to follow for people who have	25. Improve this concluding-discussion with actual in-			
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