

Atomistic resolution structure and dynamics of lipid bilayers in simulations and experiments

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(Dated: November 5, 2015)

Recent progress in the analysis of lipid bilayer atomistic resolution structure and dynamics using combination of robust experimental data and molecular dynamics simulations is reviewed. The focus is on order parameters and spin relaxation times measured with NMR and on form factors measured with SAXS and SANS for phosphatidylcholine lipid bilayers. The experimental observables are chosen since these experiments are robust, well understood, highly reproducible and the connection between raw data and simulations is straightforward. Also the comparison between simulations and these observables is bidirectionally useful; it will quantitatively measure the quality of the simulation model respect to the reality, and if the quality is sufficient, the simulations give structural interpretation for the experimental data. Significant advance of molecular dynamics simulation models is that the same simulation model can be simultaneously compared to all of this parameters. If satisfactory agreement is found, it is highly likely that the model represents the reality due to the large amount of reproduced independent experimental parameters. In this case all the mentioned experiments would be simultaneously interpreted with the same model. Phosphatidylcholine lipids are chosen since large portion of model membrane studies have been focused on this lipid, producing enough experimental and simulation data to draw comprehensive picture on the level of understanding atomistic resolution structure and dynamics. We conclude that the acyl chain region structure and its changes are generally well described in simulations, in contrast to the glycerol backbone and choline. Also cation binding is significantly overestimated by several models.

INTRODUCTION

1. Citations missing Atomistic resolution structure and dynamics of lipid bilayers has been studied with wide range of techniques for many decades motivated mainly by their presence and important role in biological systems [?]. Lipid bilayers play direct or indirect role in several physiological and pathological molecular scale processes [?]. To fully understand these processes the atomistic and molecular level understanding of lipids is required. Since atomistic resolution studies are extremely difficult from biological samples, purified lipid systems are often used [?]. The biological relevance of these model systems is supported, e.g. by similar NMR order parameters measured from living cells, lipid extracts and model systems [?].

The most detailed information about lipid bilayer atomistic resolution structure and dynamics has been achieved with various Nuclear Magnetic Resonance (NMR) and scattering techniques [?]. The first one giving direct information on structures sampled by individual lipid molecules [?] and the latter one giving complementary information on average bilayer properties, like density and thickness [?]. Both techniques give robust, accurate and reproducible quantities related to the structure and dynamics. However, for structural and dynamical interpretation both techniques needs a model which reproduce the measured quantities [?].

On the other hand, remarkable progress in hardware and software has made possible to routinely perform classical atomistic resolution molecular dynamics (MD) simulations of lipid bilayer with duration of tens or hundreds nanoseconds. Ideally the molecules are sampling realistic conformations with realistic speed in these simulations. This can be verified by calculating directly measurable quantities from simulations and comparing these to experimental values. Here we

review such comparisons for different experimental observables: C-H bond order parameters, spin relaxation times and form factor. The first and second are measured with NMR thus they represent to structure and dynamics sampled by individual lipid molecules, respectively. The last is measured with scattering techniques and represents the bilayer average properties.

The order parameters and spin lattice relaxation times have been compared between simulations and experiments for validation and interpretation since the early days of lipid MD simulations [?]. On the other hand, scattering form factors for lipid bilayers have been replacing the comparisons of simulations to the experimental area per molecule during the last decade since form factor is directly measurable quantity while area per molecule value depends on model used to analyze the scattering data [?].

If an atomistic resolution model reproduces all the above mentioned experimental parameters, i.e. order parameters, spin relaxation rates and form factor, the simulation can be considered as an ultimate model giving interpretation for all these experiments simultaneously. In addition, it would be the correct atomistic resolution representation of the system with high probability since it reproduces large amount of independently measured experimental parameters simultaneously. Thus, the model usage of the for further applications would be well justified.

Here we review recent studies which have compared the order parameters, spin relaxation rates and scattering form factors between experiments and simulations in order to quantify the quality of simulation models and interpret the experiments. We focus on phosphatidylcholine lipid bilayers due to most comprehensive available datasets for both, simulations and experiments. However, the basic ideas of the approach is valid also for other lipids and surfactants and there is also data and literature available [?]. We also discuss the observed

structural changes of lipids induced by external conditions and their relation, e.g. to ion partition.

The general conclusion from the review is that the hydrophobic acyl chain region is well described in simulation models and the atomistic resolution interpretation of experiments has been successful for this region. However, the interfacial region is less well described in simulation models which may question their usability in studies where this region is important.

In addition to the structural lipid studies, the atomistic resolution lipid bilayer simulations are nowadays widely used to study for example membrane interactions with other biological molecules (e.g. proteins, drugs, ions etc.) [?]. Since the lipid structures and interactions with other molecules may be highly relevant in some of these applications, it is crucial to be aware the weak points of the models. One goal of this review is to ease the credibility evaluation of different simulation results in these applications.

C-H BOND ORDER PARAMETERS AS ATOMISTIC RESOLUTION STRUCTURAL MEASURE

2. References should be added, the text should be polished, things should be added and checked and figures should be improved. However, the main structure and idea of the section should be visible.

Definition and properties of C-H bond order parameter

In lipid bilayer systems the order parameter of a hydrocarbon C-H vector is typically defined as

$$S_{CH} = \frac{1}{2} \langle 3 \cos^2 \theta - 1 \rangle, \quad (1)$$

where the angle brackets denote an ensemble average over the sampled conformations, and θ is the angle between the C-H bond and the membrane normal. The numerical values of order parameters vary between $-\frac{1}{2} < S_{CH} < +1$ depending on the sampled θ distribution. The definition is motivated by its connection to the dipolar and quadrupolar splitting measured with ^1H - ^{13}C and ^2H NMR techniques, respectively. The functional form comes from the fundamental theory of interactions between spin systems which gives a connection between average molecules orientations and NMR measurables [1].

If the sampled distribution of θ for a C-H bond are known, the order parameter can be straightforwardly calculated from Eq. 1. However, the sampled θ distributions cannot be uniquely determined from the known order parameter. Thus the experimental order parameter values gives a set of conditions which structural molecular model (more specifically the C-H bond vectors of the model) has to fulfill but the experimental order parameters alone cannot be used to uniquely resolve the structure. The same applies practically to all experimental parameters used in biomolecular structure determination.

Atomistic resolution molecular dynamic simulations naturally produces the sampled structures from which the θ distributions can be calculated and used in Eq. 1 to calculate the order parameters. The sampled structures in the simulation can be considered to be realistic only if the experimental order parameters are reproduced. If this is the case, the simulation can be then considered as an atomistic resolution interpretation of experimental order parameters. Before MD simulations were feasible for such usage, other models have been used for this interpretation [2? ? -7]. It is important to note, however, that the sampled structures which reproduce the order parameters are not necessarily the correct ones since, in principle, several structural models can produce the same order parameters. Significant advance of the MD models compared to the traditional models is that the same MD structures can be straightforwardly compared to other experimental observables in addition to order parameters, like ^{31}P chemical shift anisotropy [8], ^{31}P - ^{13}C dipolar couplings [9] and scattering data [?]. This will significantly reduce the possibility of getting unrealistic structures reproducing correct order parameters.

The probability for unrealistic structures with correct order parameters is further reduced by the detailed and accurate experimental data available for order parameters. Order parameters are known with high quantitative accuracy for each C-H bond present in the lipid molecule for several lipid types [10-16]. The absolute values of order parameters can be measured two independent techniques by using either ^2H NMR [17] or ^1H - ^{13}C NMR [12, 13, 18, 19] and the sign can be measured with two different ^1H - ^{13}C NMR techniques [7, 18, 19]. It is also possible that two hydrogen bonds in the same carbon has different order parameters (*forking*) which can be detected with both ^2H NMR [20? , 21] and ^1H - ^{13}C NMR [?] techniques. As a result, for example for POPC lipid molecule in lipid bilayer there is ? order parameter numbers known experimentally and a structural model for this lipid should reproduce all of these. If some the order parameters are not reproduced, it is easy to detect the weak parts of the model since a single order parameter is very local quantity depending only on the position of two atoms (C-H pair). This is an advance over several other accurately measured NMR quantities, like ^{31}P chemical shift anisotropy [8] and ^{31}P - ^{13}C dipolar couplings [9] which depend on the position of several atoms, thus in the case of disagreement, it is more difficult to pinpoint the problems in the model.

Order parameters from ^2H NMR experiments

The absolute values of order parameters are connected to the quadrupolar splitting $\Delta\nu_Q$ (^2H NMR) measured in ^2H NMR experiments through the equation

$$|S_{CD}| = \frac{4}{3} \frac{h}{e^2 q Q} \Delta\nu_Q, \quad (2)$$

where e is the elementary charge, Q is the deuteron quadrupole moment and h is the Planck's constant. The parameter q is related to the largest electric field gradient and in practise its value is not known; therefore the static quadrupolar coupling constant $\frac{e^2qQ}{h}$ is defined, and its value measured for different compounds in their solid state ($\Delta\nu_Q$ measurement from the system where order parameter is known to be 1). In C-D order parameter measurements for lipids, it is typical to use the value measured for different alkenes, $\frac{e^2qQ}{h}=170$ kHz. The relation between order parameters and quadrupolar splittings then becomes $S_{CD} = 0.00784 \times \Delta\nu_Q$. This relation is useful as many publications report only the quadrupolar splittings. For a review and more accurate description see the work of Seelig [17].

Order parameters from ^{13}C NMR experiments

To determine order parameter with ^1H - ^{13}C NMR experiments, the dipolar splitting $\Delta\nu_{\text{CH}}$ is measured. This is then related to the effective dipolar coupling d_{CH} through a scaling factor depending on the pulse sequence used in the experiment [12, 13, 18, 19]. The effective dipolar coupling d_{CH} is connected to the absolute value of order parameter through equation

$$|S_{\text{CH}}| = \left(\frac{D_{\text{max}}}{2\pi}\right)^{-1} d_{\text{CH}}, \quad (3)$$

where $D_{\text{max}} = \frac{\hbar\mu_0\gamma_h\gamma_c}{4\pi(r_{\text{CH}}^3)}$. r_{CH} is the C-H distance, μ_0 is the vacuum permittivity, and γ_h and γ_c are the gyromagnetic constants for ^1H and ^{13}C nuclei. In contrast to Eq. 2, all the parameters in Eq. 3 are in principle known. However, for the internuclear distance only the average r_{CH} is known, not the third moment r_{CH}^3 . For this reason values between 20.2-22.7 kHz are used for $\frac{D_{\text{max}}}{2\pi}$ depending on the original authors [12, 13, 18, 19, 22, 23].

Quantitative accuracy of experimental order parameter values

It must be stressed that ^2H NMR and ^{13}C NMR are fully independent experiments since the deuterium quadrupolar splitting $\Delta\nu_Q$ and the dipolar splitting d_{CH} are different physical observables. In addition, the prefactors connecting the observables to the order parameter (Eqs. 2 and 3) are independently measured. Further independent experiments are performed With ^{13}C NMR by measuring the ^1H - ^{13}C dipolar couplings using different pulse sequences [12, 13, 18, 19] when the connection between dipolar splitting d_{CH} and effective dipolar coupling d_{CH} is different.

The measurements of quadrupole $\Delta\nu_Q$ and dipolar d_{CH} splittings are relatively accurate, especially for quadrupolar splitting. **3.How accurate exactly?** Thus the quantitative accuracy of measured order parameters is mainly determined by the prefactors connecting the measured splittings to the order parameters

(Eqs. 2 and 3). Since the prefactors are independently determined for the ^2H and ^{13}C NMR measurements, the quantitative accuracy is best estimated by comparing the independently measured order parameter values.

The comparisons between order parameters measured with ^2H NMR and ^{13}C NMR by several authors shows very good agreement [12, 13, 16, 19]. Botan et al. collected literature values for PC lipid choline headgroup and glycerol backbone order parameters and concluded that order parameters would be known with the accuracy of ± 0.02 for these segments in purified PC lipid bilayer samples [16] which agrees with the estimate of Gross et al [19]. The lower order parameter reported in some studies [7, 18, 24] were suggested to arise from lower experimental accuracy. The values collected by Botan et al. and the suggested sweet spots where choline and glycerol backbone order parameters should fall in the simulation models are shown in Fig. 1 A).

Acyl chain order parameters from different techniques are compared in Table 1 by Gross et al. [19], Dvinskikh et al. [12] and Ferreira et al. [13]. The comparison by Ferreira et al. [13] is also shown in Fig. 1 C). Generally good agreement between different methods is seen also for acyl chain order parameters, however, for some segments the 0.02 accuracy might not be achieved. **4.Maybe specify to which ones?**

Qualitative accuracy of experimental order parameter values

When order parameters are measured as function of changing condition (e.g. temperature, hydration level, ion concentration, etc.), the prefactors connecting the order parameter and the experimentally measured couplings can be considered to be unchanged. Therefore, accuracy of the measured change is determined by the accuracy of the splitting measurement in contrast to the previous section. Here we refer to this as a qualitative accuracy. Due to the high resolution of splitting measurements, especially in ^2H NMR, the qualitative accuracy is much higher than the quantitative accuracy discussed in previous section.

The high qualitative accuracy of ^2H NMR experiments is exemplified in Fig. 2 by using the the classical experiment by Akutsu and Seelig [25], where the effect of different ions on the quadrupolar splittings of choline headgroup α and β segments was measured, [118] see Fig. ??). The effects of different ions on the quadrupole splittings are clearly differentiable with the experimental accuracy in Fig. ?? A). However, when transformed to the order parameter units, these changes correspond only changes below 0.03 units for β and 0.05 for α as shown in Fig. ?? B).

As another example of the high qualitative accuracy of order parameter experiments, the measured order parameters for β and α carbons as a function of hydration level for different PC lipids are shown in Fig. 3. Quantitative numbers from different experiments show slight variation between different temperatures and lipid compositions. However, in all experiments the order parameters increase with decreasing hy-

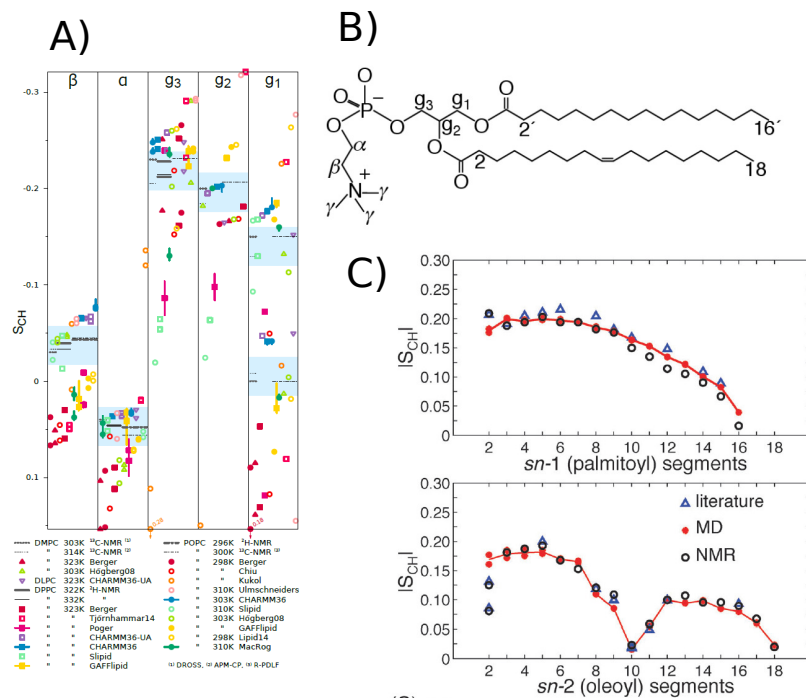


FIG. 1: A) Order parameters from simulations and experimental values from literature for glycerol and choline groups collected by Botan et al. [16]. The experimental values were taken from the following publications: DMPC 303 K from [19], DMPC 314 K from [12], DPPC 322 K from [2], DPPC 323 K from [25], POPC 296 K from [26], and POPC 300 K from [13]. The original citations for the force fields used in the simulations are Berger [27], Hogberg08 [28], Poger [29], Ulmschneiders [30], Kukol [31], Chiu [32], CHARMM36 [33], GAFFlipid [34], Slipid [35], MacRog [36], Tjörnhammar14 [37], Lipid14 [38], CHARMM36-UA [39]. The vertical bars shown for some of the computational values are not error bars, but demonstrate that for these systems we had at least two data sets (see Table 1 in Botan et al. [16]); the ends of the bars mark the extreme values from the sets, and the dot marks their measurement-time-weighted average. The interactive version of this figure is available at <https://plot.ly/~HubertSantuz/72/lipid-force-field-comparison/>. B) Chemical structure of 1-palmitoyl-2-oleoylphosphatidylcholine (POPC). C) Picture adapted from [13]. Order parameter magnitude $|S_{CH}|$ vs. carbon segment number for the sn-1 and sn-2 acyl chains of POPC (A and B respectively). Data from fully hydrated POPC at 300 K obtained with $^1H^{13}C$ solid-state NMR (black dots) [13] and MD simulations (red dots) [13], as well as data from $2H$ NMR (blue triangles) (sn-1 [4] and sn-2 [4, 40] at 300 K).

dration. [119] The smallest order parameter change is only slightly above 0.01 units, measured by Dvinskikh et al. with ^{13}C NMR [41], demonstrating that high qualitative accuracy can be also achieved with ^{13}C NMR.

In conclusion, the experimental qualitative accuracy order parameter measurements is very high. It is much higher than the accuracy achieved with state of the art MD simulations. Thus, the accuracy order parameter change comparison between simulations and experiments is limited by the simulation accuracy, not experimental.

On the other hand, it should be noted that since even very small splitting changes can be detected experimentally one

should always connect these to the changes in real molecules to avoid overinterpretation. For example, it has been measured that cholesterol induces a measurable change (less than 2 kHz) in quadrupolar splitting [43]. This may tempt to conclude that cholesterol affects to the choline structure, however, this quadrupolar splitting corresponds 0.016 unit change in the order parameter which indicates almost negligible conformational change [16].

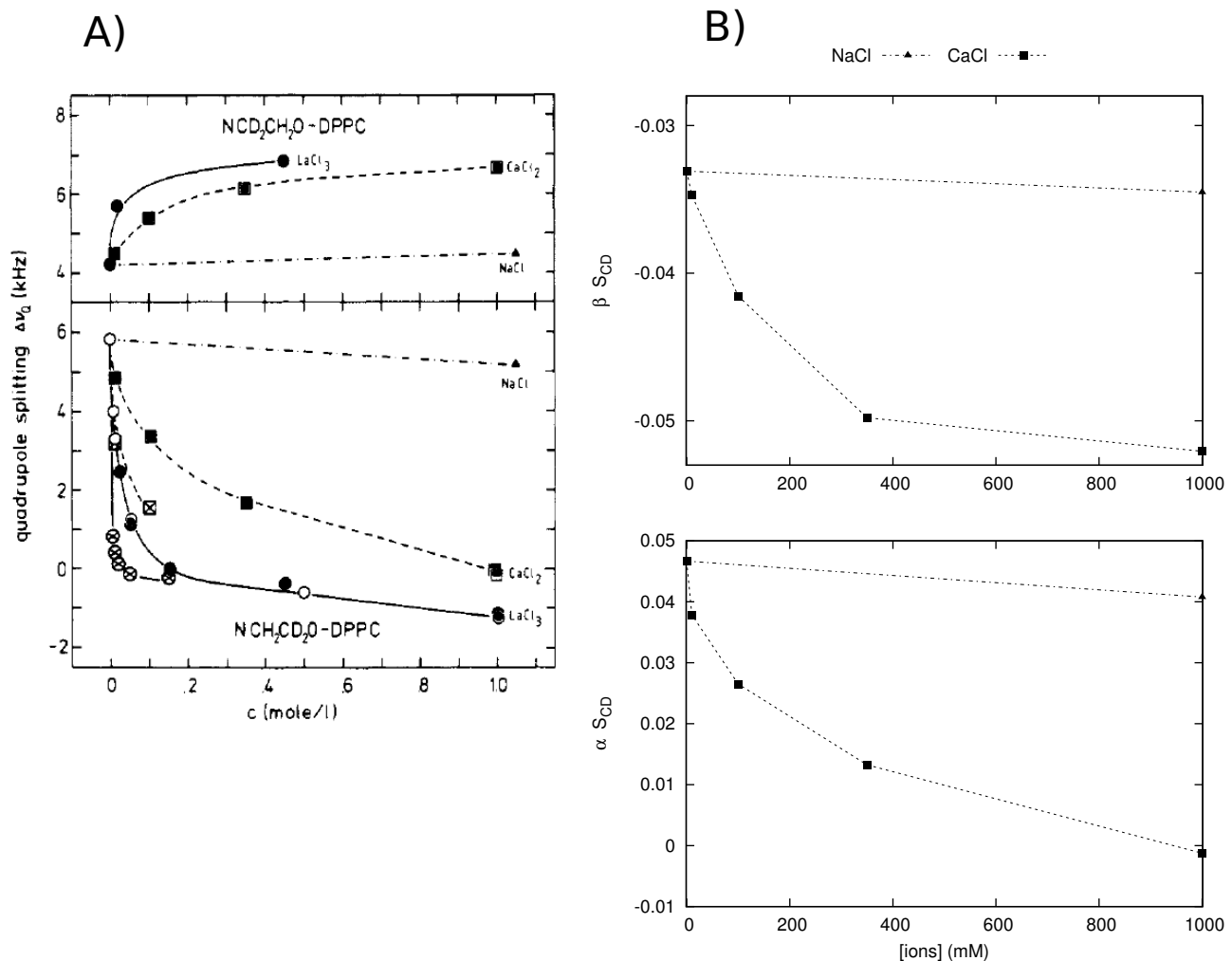


FIG. 2: A) Quadrupolar splittings as function of different ion concentrations measured by Akutsu and Seelig with ^2H NMR [25]. B) The data from A) translated to order parameters ($S_{CD} = 0.00784 \times \Delta\nu_Q$). Also the sign of β order parameter is put negative according to more recent experiments [7, 18, 19] (see also Ref. [16] and Section 4).

Signs of order parameters

While only the absolute values of order parameters are accessible with ^2H NMR, two different ^1H - ^{13}C NMR techniques allow also the measurement of the sign: first Hong et al. first measured order parameter signs for eggPC [18] and DMPC [7]; then later Gross et al. used a different NMR technique to measure signs for DMPC [19]. All the experiments report negative order parameters for almost all the segments, only α and γ are positive.

Furthermore, the signs [7, 18, 19] and magnitudes [13, 16, 20] of choline headgroup and glycerol backbone order parameters are practically unaffected by the acyl tail contents of the bilayers. Thus, it can be fairly assumed that the order parameter signs for these segments are the same in all PC lipids in bilayer. On the other hand, the positive signs for g_1 , g_3 and C_2 has been reported by Aussenac et al. [44] which has led to some confusion in comparison between simulations and

experiments [28]. However, these signs are not directly measured but extracted from the model used to interpret ^2H NMR order parameters from DMPC bicelles. Thus, it is reasonable to conclude that order parameters are negative for all segments except for α and γ , as directly measured with ^1H - ^{13}C NMR [7, 18, 19].

Even though the sign was not measurable with ^2H NMR, the sign was believed to be negative for acyl chains because θ was expected to fluctuate around 90° leading to negative order parameters [17]. This was later confirmed by using ^{13}C NMR measurements [18]. Also MD simulations always produce negative order parameters for acyl chains.

Typically when the response of order parameters to varying conditions (ions, dehydration and cholesterol) is measured, only the absolute values are reported [13, 25, 26, 41, 42, 45]. Where clear responses are observed, like with multivalent ions [25, 45] and dehydration [26, 41, 42], the experiments are done by gradually changing the conditions and the order parameter response is systematic, see Figs. 2 and 3. Thus, it

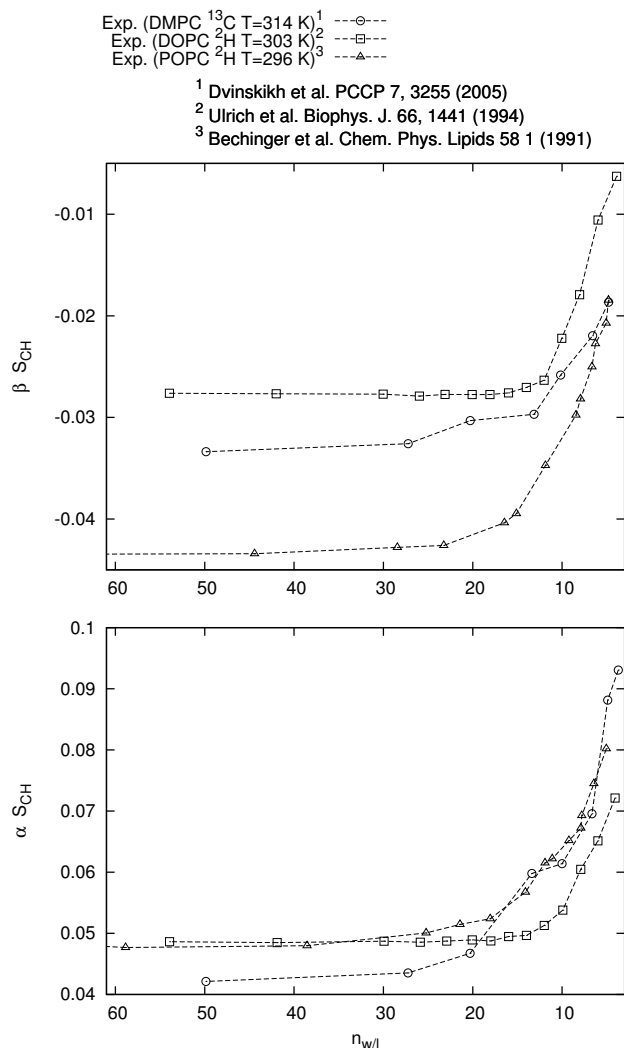


FIG. 3: Dehydration changes on α and β order parameters measured with different methods. The data taken from Dvinskikh et al. [41], Ulrich et al. [42] and Bechinger et al. [26] Also the sign of β order parameter is put negative according to more recent experiments [7, 18, 19] (see also Ref. [16] and Section 4).

is reasonable to assume that also the signs are not suddenly changing. However, it seems that the sign of the α carbon order parameter does change in response to a large amount of bound charge, such as multivalent ions. In this case, the absolute value of the order parameter first decreases to zero and then starts to increase again [45, 46], as seen from the nicely illustrated spectra shown Fig. 4 by Altenbach et al. [45].

Forking of order parameters

It is possible that the CH_2 segment is sampling such orientations that the order parameters for different hydrogens in the

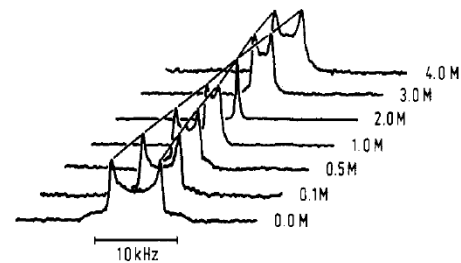


FIGURE 1: ^2H NMR spectra of coarse dispersions of POPC bilayers at various CaCl_2 concentrations (no NaCl). The lipid was deuterated at the α -segment ($-\text{NCH}_2\text{CD}_2\text{OP}-$). Measuring temperature, 40 °C.

FIG. 4: Quadrupolar splitting $\Delta\nu_Q$ of α of POPC as a function of CaCl_2 concentration, related to the order parameter as $S_{\text{CD}} = 0.00784 \times \Delta\nu_Q$. We know nowadays that the order parameter of α in the absence of CaCl_2 is positive [7, 18, 19]. Thus, the most obvious interpretation for the result is that the α order parameter decreases to zero when CaCl_2 concentration reaches 2.0M, and above these concentrations becomes increasingly negative with further addition of CaCl_2 . Reprinted with permission from Altenbach and Seelig, Biochemistry, 23, 3913 (1984). Copyright 1984 American Chemical Society.

same carbon have different values. We call this phenomena as *forking*, as done also previously to avoid confusion with splittings measured with NMR. The forking is detected with both ^2H NMR [4, 20?, 21] and ^1H - ^{13}C NMR techniques [?] as two different quadrupolar or dipolar splitting values, respectively, related to the same CH_2 segment.

The forking is observed for g_1 , g_3 , and C_2 carbon in the *sn*-2 chain segments in a fluid PC lipid bilayer, for other CH_2 segments the equal order parameters are observed for both hydrogens are equal [3, 4, 13, 20? ?]. The forking has been studied in detail with ^2H NMR techniques by separately deuterating the R or S position in CH_2 segment to assign order parameters to correct hydrogens [20]. **5.Has this been done for the C_2 carbon in the *sn*-2 chain?**

^2H NMR studies also show that the forking really arises from differently sampled orientations of the two C–H bonds, not from two separate populations of lipid conformations [20, 21]. This means that realistic atomistic resolution molecular models has to reproduce the forking correctly. Thus, the simulated order parameters has to be calculated separately for each hydrogen by taking into account the isomeric position for the structural comparison to the experimentals. Recent work by Botan et al. shows that several simulation models has problems in this respect [16] (see also Fig. 1).

Order parameter values in the literature

For ^2H NMR measurements the CH_2 segments has to be labeled with deuterium. This can be done specifically for a certain segment or for the several segments simultaneously [?]. In the first case, it is known that the measured order param-

eter (quadrupolar splitting) is related to the labeled segment. In the latter case several order parameters (quadrupolar splittings) are measured which arise from all the labeled segments, however, it is not known which order parameter belongs to which segment. Majority of the 2H NMR data in the literature is measured from samples with perdeuterated acyl chain [10] since the synthesis of specifically deuterated lipids is complicated [?] and they are expensive. From this kind of data alone one cannot know which order parameter is assigned to which CH_2 segment. On the other hand, also order parameter data from specifically deuterated lipids are available from several lipid types in various conditions [3, 4, 20? ?].

Specific labeling is not needed for order parameter measurement with ^{13}C NMR due the natural abundance of ^{13}C , however it could be used to enhance the signal for specific segment under interest [?]. Order parameter measurements with ^{13}C NMR are 2D experiments, the chemical shift being in the first dimension and dipolar coupling in the second [?]. The chemical shift depends on the local chemical environment and is different for each carbon segment. In the second dimension the dipolar coupling (order parameter) corresponding each chemical shift value is measured, and its value can be connected, in principle, to each carbon segment by using the chemical shift value. In practise, challenges occur in the acyl chain region where chemical shift values of different segments are very close to each others [12, 13, 15, 18, 19]. This issue has been addressed by filtering the spectra by using partially deuterated lipids [13] and using data from simulations and specifically deuterated experiments to help in the assignment [13, 15]. It should be noted that the overlapping spectra is a challenge only for acyl chain regions with several similar segments. On the other hand, the order parameters for hydrocarbon segments in choline, glycerol backbone, close to the double bonds, and in the beginning and the end of acyl chains can be straightforwardly measured from the natural abundance lipid samples.

The order parameters are typically measured from multilamellar samples which are good for comparison to MD simulations since it is a nearest experimental correspondence for a simulation box with periodic boundary conditions. In this work we do not discuss order parameters can be measured also from other type of samples, e.g. bicelles [44, 47, 48], or indirect measurements by using, e.g. relaxation data [49] since their comparison to simulations is less straightforward.

The available experimental data for order parameters is reviewed quite recently [10, 11]. These reviews are comprehensive especially for acyl chains in pure lipid bilayer samples. In addition to this, there is significant amount of order parameter data for glycerol backbone and headgroup segments [16], and also data as a function of changing conditions for all lipid segments [12, 13, 25, 50?]. The amount of data, especially from ^{13}C NMR, has been also increasing lately [13–15]. Especially the order parameter data of headgroup responses to varying conditions has a lot of unused potential since it can be used to measure, e.g. ion partitioning to lipid bilayer [25, 51?] and lipid protein interactions [14, 52, 53]. Simultaneously, it

gives accurate experimental parameters which can be directly compared to MD simulations [16, 51].

Order parameters from simulations

In atomistic resolution molecular dynamics simulations molecules are sampling the defined ensemble according to the used simulation parameters and the coordinates of each atom as a function of time are saved to the trajectory files. The coordinates in the trajectory file can be straightforwardly used to calculate order parameters directly from the definition in Eq. 1. The average is taken over the molecules and time.

For simulations with united atom models without explicit hydrogen atoms [16, 54, 55], the hydrogen positions can be generated post-simulationally from the positions of the heavy atoms and the known hydrocarbon geometries. This can be done explicitly by creating a trajectory with added hydrogens [16, 54], or by using equations which directly calculate order parameters from heavy atom positions [55, 56]. For C–H and C–H₂ segments without forking these two approaches gives essentially identical results when applied correctly. However, the latter is valid only for the cases with no forking, i.e. order parameters are equal for both hydrogens attached to the same carbon. Since this is not known *a priori* for the analyzed model, it is better to use the first approach with explicitly added hydrogens.

The difference in the forking analysis is most likely reason for different choline and glycerol backbone order parameters reported in the literature for the same model [16, 57]. Also different order parameters from the same model for C–H bonds has been reported in literature [54, 58] which most likely arises from incorrect implementation of widely used version of *g_order* program in the Gromacs package for this segment. Also, the *g_order* program prints $-S_{CH}$ which is most likely the reason to the reported positive order parameters for acyl chains in some studies [59]. When these issues are taken into account, the order parameters from the same models reported in the literature are generally in good agreement.

The statistical error estimates for order parameters in simulations are estimated by using the error of the mean calculated averaging over time blocks [54], over independent simulations [57] and over different lipids [16]. The maximum error bars given by all these approaches are $\sim \pm 0.01$.

It was recently pointed out that the sampling of individual dihedral angles might be very slow compared to the typical (100 ns) simulation timescales [60]. This result raises a question if typical simulation time scales are long enough to allow the molecules to sample the full phase space. On the other, another recent study showed that the slowest rotational autocorrelation function observed (for g_1 segment) in the Berger model reached a plateau (S_{CH}^2) after ~ 200 ns and its relaxation was significantly too slow compared to NMR relaxation experiments [23]. This indicates that if the typical simulation times are too short for the full sampling of the structures, then the dynamics is unrealistically slow in the simulation model.

Comparison between order parameters from simulations and experiments

Since the early days of lipid bilayer simulations the acyl chain order parameters are commonly calculated and compared to experiments when validating simulation studies [27, 56, 61–71]. Practically all the state of the art force field parametrization publications report these to be in good agreement with experiments [8, 27–39]. It is remarkable that the experimental order parameters for acyl chains in fully hydrated pure lipid bilayers can be reproduced within experimental error, see also Fig 1 C).

Exception is the C_2 segment in *sn*-2 chain in all PC lipids which is known to have measurable forking and lower magnitudes compared to other order parameters in the beginning of the acyl chain [?]. This important structural fingerprint is related to the different conformations between carboxyl segments in the beginning of chains [?]. This feature is, however, not analyzed or not reproduced for several lipid models [8, 28, 30–32, 34–37] while some models report the lower magnitude but the forking is not reproduced correctly or analyzed [8, 33, 38]. The united atom CHARMM36 is really close the experimental results [39].

In addition to the quantitatively good agreement in pure bilayers, the changes in acyl chain order parameters as a function of changing conditions are generally reasonable. For example, experimentally observed increase of order parameters as a function of cholesterol concentration [13, 55, 72, 73] and with dehydration [41, 50] are reproduced in simulations [13, 55, 74–79]. However, systematic and quantitative comparisons of these effects between experiments and simulations are rare [13, 79]. Comparison between widely used model (Berger lipids [27] and Höltje cholesterol model [80] [120]) for cholesterol containing lipid bilayer revealed that even though the acyl chain response is reasonable, the simulation model cannot be considered to agree with experiments with 34% and higher cholesterol concentrations. CHARMM36 model has been shown to slightly underestimate the cholesterol ordering effect in DMPC bilayer [77], while Slipids [78] and Lipid14 [79] models show satisfactory agreement. Lipid14 is also compared to the same extensive experimental data for POPC/cholesterol as Berge/Höltje model [13] and the agreement is significantly better [79], see Fig. 5. Also the orientation of cholesterol itself is reasonable in all models [13, 55, 77, 79], however, the cholesterol acyl chain has some issues in Höltje model [13] (too low order parameters) and in Lipid14 [79] (significant forking) while CHARMM36 reproduces experiments well [77].

The decrease of acyl chain order parameters due to the addition of double bonds is also generally reproduced by different simulation models [13, 33, 34, 38, 39, 54, 58, 78, 81–89]. Very good agreement for oleyl chain in POPC bilayer with one *cis* double bond is demonstrated in Fig. 1 taken from Ferreira et al. [13]. Also the effect of multiple double bonds (polyunsaturation) [54, 58, 81, 82, 84–86, 88, 89] (see Fig. 6) and difference between *cis* and *trans* double bonds can be repro-

duced in MD simulations [90].

In contrast to acyl chains, the order parameters for the glycerol backbone are not routinely reported in simulation literature and when reported, the agreement with experiments is concluded to be poor [16, 57, 75] or good [28, 75, 93] depending on the authors. The difference arises from different estimations for the accuracy of experimental and simulated order parameters, see also section 2. The NMRlipids collaboration recently carefully compared glycerol backbone and choline order parameters from 13 different models to the experiments and concluded that none of the available models reproduces these within experimental error, see Fig. ???. Also responses of glycerol backbone and choline order parameters to dehydration, cholesterol concentration and charge penetration were studied by the NMRlipids collaboration [16, 51]. Despite of the incorrect structures in simulation models the experimentally measured choline order parameter increase due to dehydration [16] and decrease due to penetrating ions [51] (see Fig. 7) were qualitatively reproduced. The comparison reveals, however, that the Na^+ penetration is significantly overestimated by many models [51]. Also the effect of cholesterol on glycerol backbone and choline was overestimated by the Berger/Höltje model while CHARMM36 and MacRog performed better [16].

In conclusion, the experimental order parameters for acyl chains and their changes are reasonably reproduced all state of the art lipid models (except for C_2 segment in *sn*-2). However, all models have difficulties with varying severity to reproduce the glycerol backbone and choline order parameters, and their changes.

Interplay between simulations and NMR order parameters: Validation and interpretation

As reviewed here, the order parameters can be measured with high accuracy for each hydrocarbon segment in lipid in bilayer and the values are available in the literature for wide range of different lipids in different conditions. Thus, the experimental order parameters give very detailed and local information about the orientations sampled by each C–H bonds in the lipid bilayer system. The order parameters can be also calculated from MD simulations with high accuracy and compared to the experiments. If the order parameters agree within experimental error, the simulated structures can be considered as an structural interpretation for order parameter experiments. On the other hand, if the agreement is not good, the simulation is sampling incorrect structures.

As discussed in the previous section, the order parameters for acyl chain region from MD simulations generally agree well with experiments (except for the C_2 segment in the *sn*-2 chain). Thus, the acyl chain structure is most likely realistic in simulations and they can be used for structural interpretation for this region. This is a significant advancement to the traditional structural models build based on the fittings to the order parameters [?]. The dynamical visualization of simula-

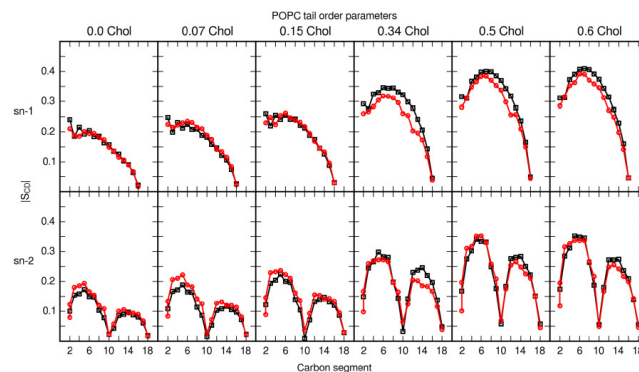


FIG. 5: Cholesterol effect on acyl chain order parameters compared between ? model [79] and experiments [13]. The agreement with experiments with this model is significantly better than Beger/Höltje based model compared by Ferreira et al. [13].

tion trajectory immediately reveals very dynamical nature of acyl chains, rapidly sampling large amount of different conformations (for dynamics see the Section ??). These videos published by several authors in supplementary information [?] gives significantly better intuitive understanding of dynamical nature of lipid bilayers compared to the static ones from traditional models. Since the lipid bilayers can be considered as a simplistic models for cell membranes and other biological lipid layers, this understanding has significant impact on biophysics and biochemistry.

Also the order parameter changes with changing conditions are qualitatively reproduced in the acyl chain region, however the systematic quantitative comparison of changes is rare [13, 54, 79]. The MD simulations have been especially useful to explain the origin of order parameter decrease due to *cis* double bonds in the acyl chain [54, 58, 85, 86, 94–96]. The order parameter decrease might arise from reduced order of the chain or from the changed average θ angle in Eq. 1. From NMR experiments alone it was impossible to judge which is the correct explanation for the decreased order parameters due to double bonds [85, 86, 94–96]. Several simulation studies by different authors using different models has showed that the decreased order parameter due to *cis* double bonds can be reproduced by introducing proper dihedral potentials next to double bonds, and due to the flexibility of these dihedrals the polyunsaturated acyl chain becomes more flexible and the order is reduced [54, 58, 86, 94–96]. These studies concluded that order parameter decrease due to double bonds arises from genuine disorder of the chain, not from the changes in average angle [95, 96]. This is a prime example of the case where MD simulations have significant advance over more traditional modeling approaches [95].

The increase of acyl chain order parameters and related bilayer thickening due to addition of cholesterol is also qualitatively reproduced by simulations giving also intuitive visualizations for these effects [13, 79?]. However, quantitative comparison reviewed in previous section reveals that some models do not reproduce the acyl chain order parameters within experimental error in cholesterol mixtures [13, 77].

This indicates that these models are not yet accurate enough to give atomistic resolution interpretation to delicate lipid cholesterol interactions which are known to induce liquid-ordered and liquid-disordered phase coexistence [97]. The recent Amber model [79] seems promising in this respect, see also Fig. 5. Careful quantitative comparison of dehydration induced ordering effect has not been made but it is mentioned to be weaker in simulations compared to experiments [75].

Simulation studies have also predicted changes in the acyl chain region which are yet to be experimentally confirmed, e.g. order parameter decrease due to lipid oxidation and changes in order parameter sign in oxidized acyl chain [59].

As discussed in the previous section, simulations models are not able to reproduce the glycerol backbone and choline headgroup order parameters within experimental error [16] in contrast to acyl chains. Thus, even the state of the art simulation models are not able to resolve the sampled atomistic resolution structure of these segments which has been also tremendous challenge to the more traditional models [?]. Consequently, the conclusions made from MD simulations which depend on atomistic resolution structure of energetics of these segments should be taken with extreme caution. On the other hand, the qualitative response of choline α and β order parameters to the dehydration and penetrating ions (increase and decrease, respectively) was correctly reproduced by several models, despite of the incorrect structure in fully hydrated lipid bilayer [16, 51]. These changes could be related with the changes of P–N vector angle respect to the membrane normal as suggested previously in [?]: the tilting of P–N vector more parallel to membrane plane with dehydration leads to the increase of choline order parameters [16] and *vice versa* with penetrating ions [51]. **6.The analysis with ions not actually done yet!** The widely used Berger/Hiltje model significantly overestimates the structural effects due to cholesterol in the glycerol backbone and choline regions [13, 16] which seems to be the case also for Lipid14 [79].

The recent progress in the NMRlipids collaboration has shown that simulations confirm the electrometer concept suggested by Seelig et al. in a serie of classical publications [?

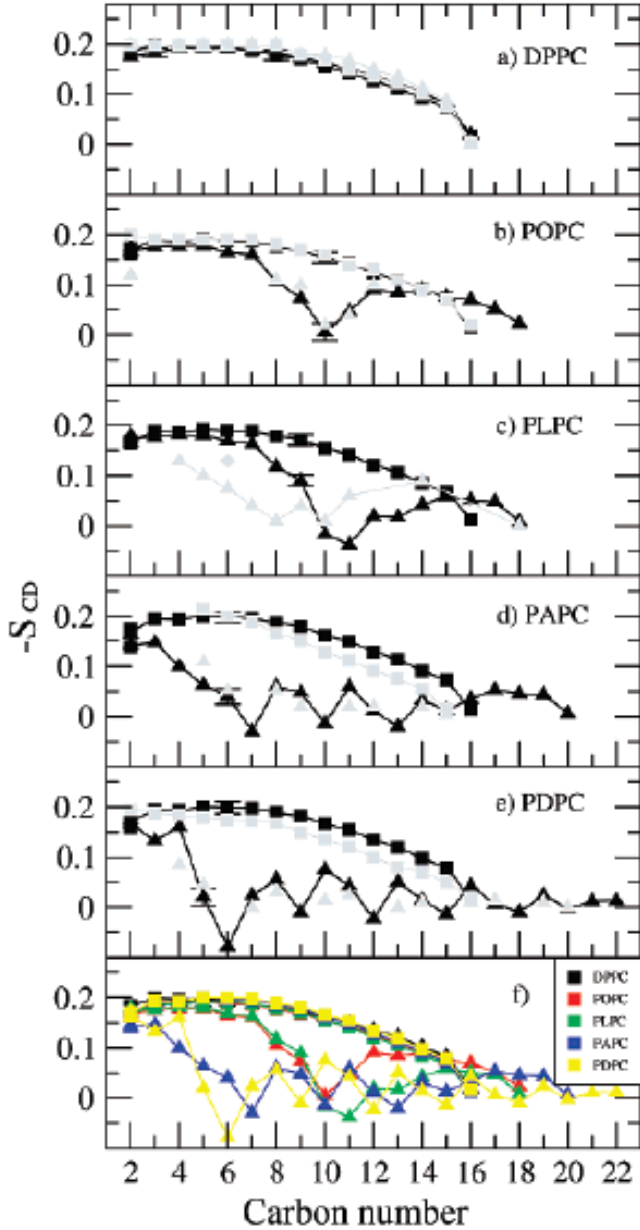


FIG. 6: Figure comparing order parameters in polyunsaturated acyl chains between simulations and experiments taken from [54]. Order parameters for the sn-1 (squares) and sn-2 (triangles) chains of (A) DPPC, (B) POPC, (C) PLPC, (D) PAPC, and (E) PDPC. Simulation results are shown in full black, and experimental results for comparison in gray. Additionally, part F summarizes the data for all bilayers from the simulations. Experimental order parameters were chosen for comparison as follows. The order parameters for DPPC ($T=323\text{K}$) are based on studies by Petrache et al. [91] whereas the experimental S_{CD} values for PDPC and for the sn-1 chain of POPC ($T=310\text{K}$) are based on studies by Huber et al. [85] For the sn-1 chain of PDPC, the data set at 310K is obtained by linearly interpolating between data at 303 and 323K , whereas for the sn-2 chain the data at 303K are presented [85]. Experimental values for the sn-2 chain of POPC are based on studies by Seelig et al. [4] A single experimental value is available also for the sn-2 chain of the PLPC bilayer at 313K (diamond) [6] to compare with our simulated order parameters for PLPC. Together with PLPC, there are also experimental results for PiLPC ($T=313\text{K}$) [6]. Experimental order parameters for the sn-1 and sn-2 chains of PAPC ($T=303\text{K}$) are based on quadrupole splittings measured by Rajamoorthi et al. [92]. For the sn-1 chain the monotonic decrease through the acyl chain is expected. For the sn-2 chain, values are fitted such that the agreement is as good as possible.

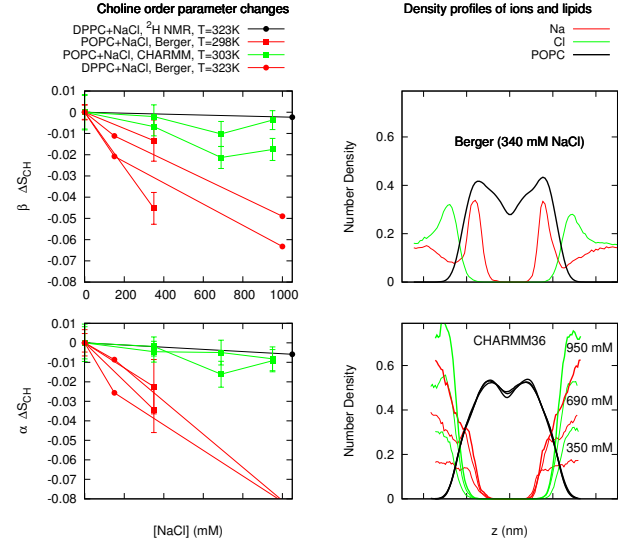


FIG. 7: Changes in order parameters as a function of Na^+ concentration (left column) and ion density distributions (right column). With Berger model significant order parameter reduction and ion partition is observed while with CHARMM36 modest order parameter change and ion partition observed. The results confirm the concept by Seelig et al. which allows the ion partition to be measured by changes in α and β order parameters. The results clearly show that Berger model has too strong Na partition. For more discussion see [51].

]: the decrease of α and β order parameters is a measure of charge penetrated in PC lipid bilayer [51], see also Fig. 7. The concept gives a direct and quantitative route to compare charge binding into PC lipid bilayers between simulations and experiments through the choline order parameters [51]. While experimental results show only very weak or negligible Na^+ binding, some simulation models predict significantly stronger binding, see [51] and Fig. 7. This is a serious artefact since specific cation binding makes a lipid bilayer positively charged, which has a potential to lead incorrect conclusion when interactions with charged objects are studied. Thus the numerous conclusions from simulation studies with strongly binding Na^+ has to be taken with extreme caution. In contrast to monovalent ions the Ca^{2+} and other multivalent ions significantly partition to PC bilayers [?]. This is also seen in simulations, however, the structural effects are overestimated at least in the Berger model [51]. The CHARMM36 behaves more realistically, however, more careful simulation studies are needed to conclude if some of the existing models in capable to correctly reproduce PC ion interactions [51].

In conclusion, the atomistic resolution MD simulations are invaluable in understanding the structural details and their changes in acyl chain region, especially for double bonds. However, serious artefacts are possible, or even likely, in simulations studies where choline or glycerol backbone structure, or cation binding are important.

C-H BOND ROTATIONAL DYNAMICS FROM SPIN RELAXATION RATES AND SIMULATIONS

7. This is the first sketch of this section. A lot of references should be added, the text should be polished, things should be added and checked and figures should be improved. However, the main structure and idea of the section should be visible.

Definition and properties of rotational autocorrelation function

The second order auto-correlation function for the reorientation of the C–H chemical bond axis is defined as [98]

$$g(\tau) = \langle P_2[\vec{\mu}(t) \cdot \vec{\mu}(t + \tau)] \rangle, \quad (4)$$

where P_2 denotes the second Legendre polynomial, $P_2(\xi) = 1/2(3\xi^2 - 1)$, $\vec{\mu}(t)$ is the unitary vector having the direction of the C–H bond at time t , and the angular brackets denote a time-average. This autocorrelation is usually chosen to describe the C–H bond rotational dynamics since it is connected to the experimentally measurable spin relaxation rates through its Fourier transformation called spectral density

$$j(\omega) = 2 \int_0^\infty \cos(\omega\tau) g(\tau) d\tau. \quad (5)$$

In this review we focus only on experiments measured from multilamellar samples with randomly oriented sheets, thus only the second order auto-correlation function is needed [?].

In randomly oriented multilamellar samples the auto-correlation function of bond orientations always decays to zero with long enough time scales. However, the relaxation timescales can be divided to two distinct timescales. First the relaxation processes shorter than microsecond timescales occurs when lipid molecules are reorienting in the lipid bilayer but are not essentially moving between lipid bilayer regions with different orientations. Then with larger than microsecond timescales the movement between differently oriented bilayer regions decays the rotational correlation function to zero. In addition, MAS experiments the sample spinning causes lead orientational relaxation in kHz region. The full auto-correlation decaying to zero is illustrated in Fig. 8. Due to the timescale separation the correlation function can be written as [99]

$$g(\tau) = g_f(\tau)g_s(\tau); \quad (6)$$

$g_f(\tau)$ describes the decay of $g(\tau)$ due to fast molecular motions and $g_s(\tau)$ contains the contribution from slower motions

$$g_s(\tau) = e^{-\frac{\tau}{\tau_s}} \left[\frac{2}{3} \cos(\omega_R \tau) + \frac{1}{3} \cos(2\omega_R \tau) \right], \quad (7)$$

where τ_s is a correlation time due to slower isotropic molecular motions originating from the diffusion between bilayers with different orientations of their principal symmetry axis,

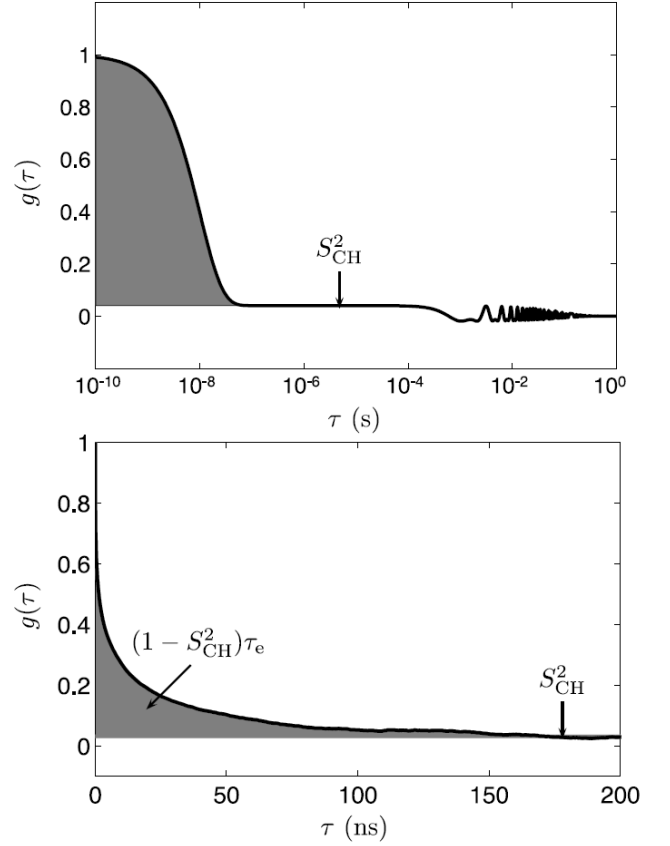


FIG. 8: (Top) Illustration of the auto-correlation function $g(\tau)$ and effective correlation time τ_e for a ^{13}CH bond in a lipid bilayer in MAS experiment (x-axis with logarithmic scale). Plateau after short timescale relaxation processes ($g(\tau)_f$) is shown between roughly 10^{-7}s and 10^{-4}s . After this timescale the slow relaxation processes ($g(\tau)_s$) and oscillation due to MAS (Eq. 7) are shown. (Bottom) Example of $g(\tau)$ from a united-atom MD simulation of a POPC bilayer in excess water, illustrating the decay towards S_{CH}^2 (x-axis with linear scale). This represents the $g(\tau)_f$ in Eq. 6 and decrease to the plateau in the top figure. The effective correlation time τ_e is equal to the area in gray scaled by $(1 - S_{CH}^2)$. Figure adapted from [23]

and the cosine terms are the contribution from magic angle spinning of the sample, rotating at $\omega_R/2\pi$ cycles per second [100], typically in the kHz frequency range.

The order parameter measurements with ^2H NMR and ^{13}C NMR measure the bond order after the relaxation of rotational motion inside the bilayer plane but before the relaxation between different bilayer orientations, as illustrated in Fig. 8. In typical molecular dynamics simulations with periodic boundary conditions the lipid molecules are restricted to single bilayer orientation and also the timescales are currently typically below microsecond. In these simulations the auto-correlation function in Eq. 4 decays to the square of order parameter in Eq. 1 in bilayers with planar symmetry, i.e. no microscopic phase separation of defects present. Also this is illustrated in Fig. 8.

The rotational correlation function describes how long does it take for a single molecule on average to sample the conformations. The effective correlation time

$$\tau_e := \int_0^\infty \frac{g_t(\tau) - S_{CH}^2}{1 - S_{CH}^2} d\tau \quad (8)$$

can be used as an intuitively useful single parameter to describe this time. The larger this parameter is, the longer it takes in average to sample the conformations related to the bond. With this definition the area between the correlation function and its plateau becomes $(1 - S_{CH}^2)\tau_e$.

Detecting C–H bond dynamics experimentally

The most used parameter to detect the C–H bond dynamics experimentally in time scales comparable to simulations are the spin-lattice relaxation rates R_1 from deuterium labels and ^{13}C . R_1^C measured from ^{13}C is connected to the spectral density (Eq. 5) through the equation

$$R_1^C = \frac{D_{\max}^2 N_H}{20} \left[j(\omega_H - \omega_C) + 3j(\omega_C) + 6j(\omega_C + \omega_H) \right], \quad (9)$$

where ω_C and ω_H are the Larmor angular frequencies of ^{13}C and ^1H respectively, N_H is the number of bound protons and $\frac{D_{\max}}{2\pi} \approx 22$ kHz as in section 2. R_1^D measured from ^2H is connected to the spectral density (Eq. 5) through the equation

$$R_1^D = \frac{12\pi^2}{40} \left(\frac{e^2 q Q}{h} \right)^2 \left[j(\omega_D) + 4j(2\omega_D) \right], \quad (10)$$

where $\frac{e^2 q Q}{h} = 170$ kHz as in the case of order parameters in section 2.

Also the model free approach to measure the effective correlation time (Eq. 8) was recently introduced [23]. The method is based on the combination of experimental order parameter S_{CH} , spin-lattice relaxation rates R_1 and the transverse magnetization under a spin lock pulse $R_{1\rho}$ with measured appropriate nutation frequency through equation

$$\tau_e \approx \frac{5R_{1\rho}^{\text{plateau}} - 3.82R_1^C}{D_{\max}^2 N_H (1 - S_{CH}^2)}. \quad (11)$$

Analyzing C–H bond dynamics from simulations

As in the case of order parameters, the auto-correlation function for each C–H bond can be calculated directly from simulations using the definition in Eq. 4 since the trajectories of each atom is known as a function of time. As in the case of order parameters the positions of hydrogens can be determined for united atom models based on heavy atom positions and assuming tetrahedral configurations [23, 101, 102]. Usually in the correlation function calculation all the available

time intervals from the simulation data are used and the average over those and all molecules is taken. However, since the amount of data decreases when the time interval approaches the total length of the simulation, usually the largest time interval used is the half of the total simulation length, for more details see [103].

To connect the auto-correlation functions from simulations to the experimentally measurable spin lattice relaxation times, the spectral density (Eq. 5) must be first calculated. In principle, this could be done using numerical Fourier transformations techniques, however this often leads to unnecessarily large fluctuations. Instead, commonly used approach is to fit analytical functional form to the calculated auto-correlation function and then use analytical Fourier transform of the fitted function [23, 101, 102, 104, 105]. Most commonly the sum of 4 or more exponentials is used as a fitting function [23, 54, 94, 104–106] but also stretched exponential has been used [101, 102]. Numerically the functional form of the fitting function should not matter as long as the fit is good, however, theoretically the correct correlation function form to describe the modes of physical motion can be debated [10, 102, 107–109]. It is clear from correlation functions from simulations that one exponential is not enough to produce a good fit while 4 gives a reasonable fit [94]. This is not surprising since more than one relaxation timescale is definitely expected to be present in lipids in bilayer [10, 104–106].

After the fitting the analytical form of the spectral density predicted by simulations is available. Then its values can be calculated at the required larmor frequency values and substituted to Eqs. 9 and 10 to get the R_1^C and R_1^D . The value of the effective correlation time can be calculated directly from the integrated area below the correlation function, see Fig. 8 or from Eq. 30 in Ref. [23].

Comparing C–H bond dynamics between simulations and NMR experiments

The spin lattice relaxation parameters R_1^C , R_1^D and $R_{1\rho}$ are considered as directly measurable experimental parameters and the effective correlation time τ_e can be derived directly from directly measurable parameters without further assumptions (see Eq. 11 and Ref. [23]). Spin lattice relaxation parameters R_1^C and R_1^D can be calculated from simulations by first calculating the auto-correlation function (Eq. 4), then calculating the spectral density from the Fourier transformation (Eq. 5) and finally substituting its values into Eqs. 9 and 10 (see also previous section). The effective correlation time can be calculated directly from integrated area and order parameter or from Eq. 30 in Ref. [23]. In practise, the $R_{1\rho}$ cannot be calculated from simulations directly since its value depends also on the slow relaxation dynamics ($g_s(t)$ in Eq. 6) which is not present in simulations. The same applies to the calculation of NOESY relaxations rates and in this case decay time of 170 ns was assumed for the $g_s(t)$ [110], while 4.2 ms was

measured by Ferreira et al. [23].

As seen from Eqs. 9 and 10, the numerical values of R_1^C and R_1^D depend also on the carbon ω_C , hydrogen ω_H and deuterium ω_D Larmor frequencies which, in turn, depend on the spectrometer external magnetic field. From simulations it is straightforward to calculate the spectral density with any Larmor frequency value to get the R_1^C and R_1^D as a function of external magnetic field. However, in experiments with standard spectrometers the external field cannot be changed, i.e. each spectrometer has their specific field strengths. Thus, to measure R_1^C or R_1^D values as a function of magnetic field one has to use several spectrometers which is tedious and spectrometers exist only with limited amount of magnetic field strengths. Despite of these challenges this kind of experiments have been done and the available data is reviewed by Leftin et al. [10]. Another approach to measure the magnetic field strength dependence of spin lattice relaxation times is the Field Cycling NMR [111, 112]. However, this is not yet feasible with standard spectrometer and only limited amount of data is available, mostly from ^{31}P NMR [111–113] but also from ^{13}C NMR [114].

The magnetic field strength dependence of spin relaxation times between simulations and experiments is compared in several studies [71, 101, 102, 104, 105, 108]. Examples of such comparison for acyl chain segments are shown in Fig. 9. On the other, some studies have compared R_1^C and R_1^D measured with one magnetic field strength to the value calculated from simulations [23, 54, 71, 86, 89, 94]. Example of such comparison between R_1^C from simulations and experiments is shown in Fig. 10. Comparisons seem to generally show a good agreement with large Larmor frequencies which is getting worse when the Larmor frequency decreases. On the other hand, the level of the agreement depends on the carbon segment and the type of relaxation used in the comparison; Berger model compared with ^2H NMR gives better agreement for C_7 segment compared to C_3 (Fig. 9 B)) while comparison to ^{13}C NMR relaxation with one frequency gives similar discrepancy for both segments (Fig. 10). For CHARMM the agreement seems better when going towards bilayer center, see Fig. 9 A). Due to the complicated connection between molecular dynamics and spin relaxation it is not straightforward to make conclusions about the dynamical differences between reality and simulations. This is, however, possible with careful analysis as discussed in the next section.

To ease the intuitive interpretation of experimentally measured rotational dynamics Ferreira et al. showed that the effective correlation time can be measured by combining directly measurable quantities without any assumptions on dynamical processes present in the fast relaxation regime [23]. The comparison of effective correlation times between experiments and simulations for different segments is shown in Fig. 10. From this result it is straightforward to conclude that acyl chain rotational dynamics is generally well described while in the interfacial region, especially in glycerol backbone, the dynamics is too slow in simulations. It should be noted, however, that the effective correlation describes the total relax-

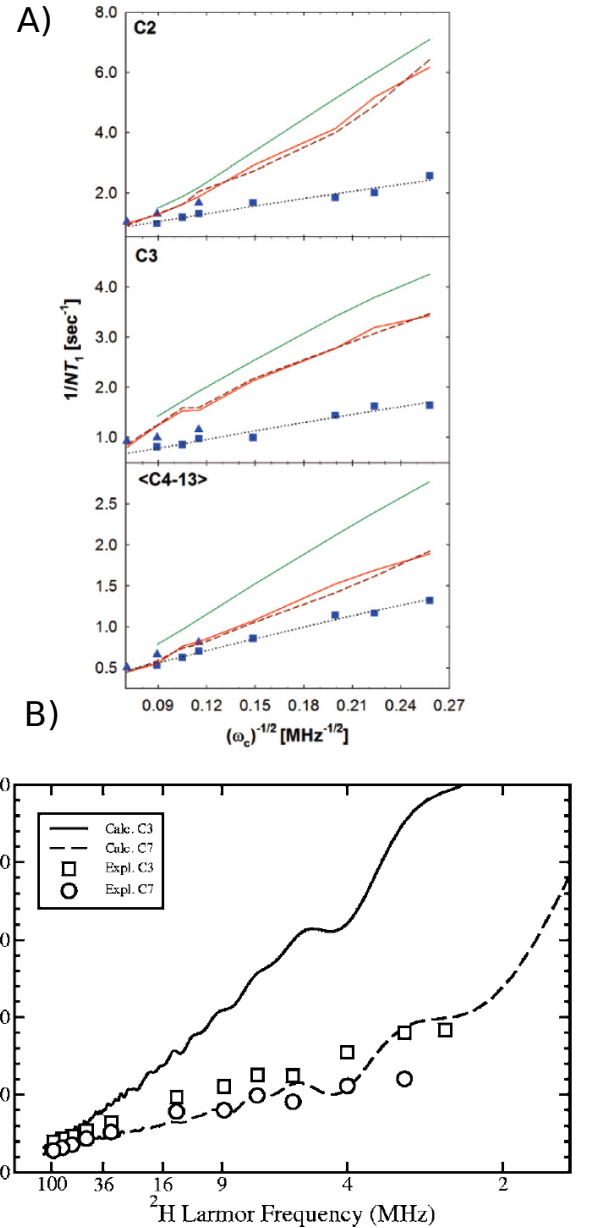


FIG. 9: A) Comparison of R_1^C dependence on magnetic field between experiments and CHARMM simulations for acyl chain carbons (DPPC bilayer in 323K) adapted from [108]. Experiments as points; MD simulations as solid and dashed lines; and a model-free fit to the vesicle data as dotted lines. B) Comparison of R_1^D dependence on magnetic field between experiments and Berger simulations for acyl chain carbons (DMPC bilayer in 300K) adapted from [102].

ation over all short timescales present in $g_f(t)$. Even if this is correct, the balance between different processes with different relaxation times may not be correct. This is actually seen in the acyl chains, where R_1 does not perfectly agree with experiments while τ_e does.

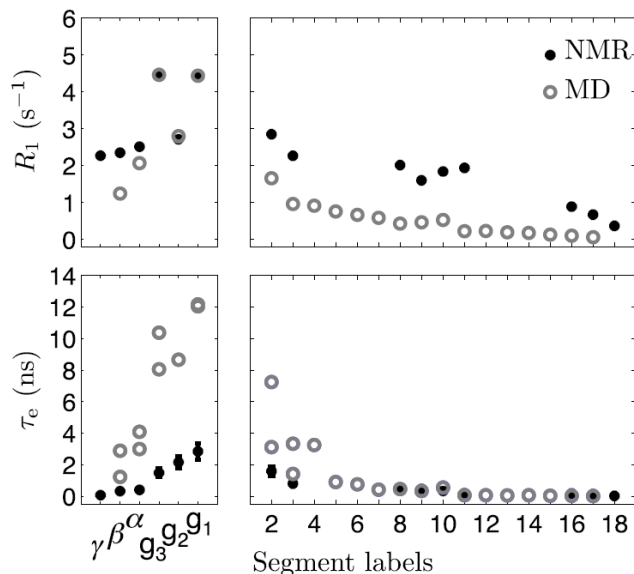


FIG. 10: (top) R_1^C for POPC bilayer in 298K calculated from simulations with Berger model compared to experimental results measured with field strength corresponding to the Larmor frequency of 125 MHz for ^{13}C . (bottom) Effective correlation times for the same system compared between simulations and experiments. Figure adapted from Ferreira et al. [23].

Interplay between simulations and NMR spin lattice relaxation times: Validation and interpretation of dynamics

By measuring single spin relaxation time values it is almost impossible to make any conclusions about molecular dynamics due to their complicated connection through the spectral density. Even changes of relaxation times cannot be directly related to molecular dynamics without further information since, e.g. faster dynamics may lead to the decrease or increase of spin relaxation times, depending on the other dynamics processes present and the used magnetic field strength as demonstrated, for example in [23].

Careful studies of spin relaxation times as a function of temperature and magnetic field to overcome this issue are recently reviewed by Leftin and Brown [10]. From compilation of different experimental data sets it was possible to conclude, for example, that the interfacial region of lipid molecules has slower dynamics than acyl chains [10]. Another successful approach to interpret the spin relaxation times has been to use the atomistic resolution molecular dynamics simulations to reproduce the measured changes and then analyze the dynamical changes from the simulation trajectory [86, 94, 115]. This approach has been especially useful in the studies of polyunsaturated acyl chain dynamics which concluded by combining the simulation and NMR relaxation data that the double bonds speed up the chain dynamics due to flexible dihedrals next to the double bonds [86, 94–96].

To ease the interpretation of the measured spin relaxation

times Ferreira et al. recently introduced a model free connection between effective correlation time and directly measurable spin relaxation rates [23]. The main advantages of effective correlation time is that its connection to molecular dynamics is straightforward in the sense that the general dynamics is faster when effective correlation time decreases and *vice versa* and that it can be quantitatively compared to simulations. As shown in Fig. 10 these experiments immediately show that the glycerol backbone has the slowest dynamics of lipid segments (in agreement with conclusions from the combination of several previous experimental sets [10]) and that the dynamics of these is significantly underestimated by the Berger model.

Most importantly, the rough agreement of spin relaxation rates and effective correlation times between simulations and experiments shows that the rotational dynamical processes present in simulations has the correct order of magnitude [23, 54, 71, 71, 86, 89, 94, 101, 102, 104, 105, 108]. Consequently, the dynamical visualizations of simulation trajectories (videos) can be considered as a realistic intuitive presentations of lipid bilayers. However, in more detailed studies it should be kept in mind that all the dynamical (and also structural) details are not exactly correct.

The lipid rotational dynamics has been often interpreted by using the so called wobble in the cone model [104, 105, 108, 109, 114]. The main idea of the model is that the whole lipid molecule is wobbling as a cone such that all the all the segments share the time scale for this wobbling. In addition, each segments have further timescales related to their dynamics inside the cone. The auto-correlation functions assuming the wobble in the cone model can be nicely fit to the simulation data and also some experimental results can be reproduced. However, also auto-correlation functions having different type of dynamics can be used to do the same. On the other hand, it has been recently pointed out that significant changes of structure and dynamics experienced in acyl chain region may not reflect to the headgroup [16, 116] indicating weak coupling between these segments. This idea is also in line with one possible interpretation of recent field cycling experiments [113]. In addition, the role of undulations in the relaxation data measured with low frequencies is under discussion [10, 107–109].

In conclusion, the timescale of rotational dynamics is correct in simulations while the exact correlation times and relaxation processes are not fully reproduced. More experimental and simulation studies are needed to fully understand current quality of dynamics in current models. To interpretate the relaxation processes present in lipid bilayer, a model which can be shown to agree with experiments on all fast timescales is needed.

STRUCTURE FACTORS FROM SCATTERING AND SIMULATIONS

For this section I would be more than happy for some help

Form factor measured with X-ray or Neutron scattering

In scattering experiments the scattering intensity $I(q) \sim |F(q)|^2 S(q)$ is observed, where $F(q)$ is the form factor and $S(q)$ is the structure factor. In the case of lipid bilayer the structure factor describes the structure of lamellar sheets and the form factor describes the internal structure of lipid bilayers. The structure factor depends on the topological phase of lipids and can be considered to be practically constant for unilamellar vesicles while in multilamellar phase its form is known. Thus, in practise direct information about form factor is achieved from the measured scattering intensity.

Form factor for a lipid bilayer is defined as a Fourier transformation of the scattering length density $\rho_s(z)$ along the membrane normal coordinate

$$F(q) = \int_0^\infty \rho_s(z) \cos(qz) dz. \quad (12)$$

The X-rays are scattering mainly from electrons thus the scattering length density in this case is $\rho_s(z) = \rho_w(z) - \rho_l(z)$, where $\rho_w(z)$ and $\rho_l(z)$ are the water and lipid electron density profiles along the membrane normal, respectively. The neutrons are scattered from nuclei thus the scattering length density is **8. I do not have correct papers at hand now. I will do this later. 9. Discussion about related issues can be found at: <https://github.com/NMRLipids/NMRLipids.V-Review/issues/1>**

Form factors are quite accurate given that experimental care has been taken. A delicate issue can be subtraction of background (from water and capillary). **10. The discussion is going in at: <https://github.com/NMRLipids/NMRLipids.V-Review/issues/2>**

Key questions now:

Do we get independently measured Form Factors plotted to the same figure?

Are there good references for the accuracy?

Form factor calculation from simulations

To calculate the form factor measured with scattering from simulations the scattering length density profile needs to be first calculated and then substituted in Eq. 12. In the case of x-ray scattering this is the electron density profile along membrane normal. Since electrons are not explicitly present in classical MD simulations, those exact locations need to be placed when calculating the electron density profile. The most common practise is to assume that all the electrons belonging to each atom are pointwisely located at nucleus location in the MD simulation [?]. Electrons are also assumed to be distributed as gaussian [?] or ? functions [?] around the atom positions. Very popular SimtoExp program uses the last approach [?] while most molecular dynamics simulation modelling papers are using the first option [?]. However, the results are not expected to strongly depend on this choice since this affects only to very fine details of the electron density profile and the thermal motion of atoms in MD simulations will coarse the differences further [?].

The neutron scattering length density is typically calculated from atom positions using the neutron scattering lengths known for different atoms [?].

11. More discussion at:

<https://github.com/NMRLipids/NMRLipids.V-Review/issues/3>

and

<https://github.com/NMRLipids/NMRLipids.V-Review/issues/4>

The most important question is now to check the discussion about undulations in the literature

Interplay between simulations and scattering experiments: Validation and interpretation

Similarly to NMR order parameter, the form factor gives accurate information about lipid bilayer structure but the structure cannot be uniquely resolved from the experimental data only. To solve the structure different models have been introduced [?]. The general idea of these models is to introduce plausible functional form for the scattering length density profile arising from different molecular components of lipids and then fit the weights of these components such that the experimental form factor

is reproduced [?]. Also MD simulations have been used in structural interpretation [?]. Also combination of other structural models and MD simulations have been used [?].

The structural models based on pure fitting to scattering data are shown to easily produce unphysical structures arising from the fact that several scattering density distributions can produce correct form factors [?]. However, when the most obvious unphysicalities are removed, the models based on fitting produce form factors with better agreement with experiments than MD simulation models [?]. Careful comparison between structures from MD models and fittings to x-ray and neutron data shows that the structures from different approaches are very similar but not exactly same [?]. Differences are found especially in the lipid headgroup region which is in agreement with the NMR order parameter studies which has shown that glycerol backbone and choline structures are not reproduced within experimental error in MD simulations [16].

Ideally the MD simulations would correctly reproduce the experimental form factors in zero tension giving the atomistic resolution interpretation for the experiment and simultaneously confirming the quality of the used model. This comparison is nowadays routinely done in almost all force field parametrization publications instead of comparing area per molecule as previously done [?]. The agreement is often concluded to be good, however, almost always detectable differences between experiments and simulations can be seen [?]. The attempts to run simulations with fixed area per molecule (generating tension in bilayer) in order to interpret the experiments have shown that this can improve the agreement, however, the models based on fitting gives better agreement and slightly different structure [?]. On the other hand, different simulation models with different electron density profiles and form factors are concluded both to be in good agreement with experiments [?]. This is possible since the absolute electron density is not known from experiments and is thus fitted to the simulation results when comparing [?].

In conclusion, all the state of the art simulation models gives form factor very close to experimental data, however, it seems still the best agreement can be produced by fitting the scattering density models to the experimental data [?]. The problems to exactly reproduce the scattering data has been often related to the lipid headgroup structure which agrees with the conclusions from the NMR studies [?].

12. The more detailed discussion can be found at:

<https://github.com/NMRLipids/NMRLipids-V-Review/issues/5>

CONCLUSIONS

Recent studies quantitatively comparing atomistic resolution structure and dynamics of lipid bilayer between experiments using C–H bond order parameters, spin relaxation rates and scattering form factors are reviewed. The purpose of these studies is to quantify the atomistic resolution structural and dynamical quality of simulation models as well as to give an atomistic resolution interpretation for the experimental results. For interpretation of these experiments the atomistic resolution MD simulations reproducing all the experiments would be an ultimate tool since the same model can be used to interpret all the experiments. For the same reason, this atomistic resolution model would be most likely a realistic representation for atomistic resolution structure and dynamics of lipid bilayers. The main conclusion of this review is that the current MD simulations are not quite yet on the level to perform this task, however, in several aspects indicate that there is potential to improve the models to become truly realistic atomistic resolution representations of these systems.

More specific conclusions are the following:

- The order parameters for each C–H bond in lipid molecules in bilayers can be measured with high quantitative accuracy with both ²H NMR and ¹³C NMR and this data is available for wide range of lipids in different conditions. Comparison of this data to simulations gives a very detailed picture about the quality of sampled atomistic resolution structures in lipid bilayer and may also help in structural interpretation of experiments. The order parameter changes with varying conditions can be used to study and compare structural changes between simulations and experiments.
- The main conclusions from the comparison of order parameters between experiments and simulations are: 1) the acyl chain structures are generally described realistically for PC lipid bilayers in simulations, 2) the changes in acyl chain region are qualitatively correct but not always quantitatively, 3) the glycerol backbone and choline structures are not within experimental error for any available model and 4) the results depending on structure and energetics of these sections, like ion binding and lipid–cholesterol interactions should be taken with caution.
- The C–H bond rotational dynamics has correct timescale in simulations models for which comparison has been done. Most likely the same applies for all models. However, more careful comparison reveals that, e.g. the glycerol backbone dynamics is too slow in Berger model. Also for CHARMM model all relaxation process do not seem to be correctly described.
- The development in the scattering methodology has allowed the direct comparison of whole bilayer structure between experimental form factor and simulations. This is complementary to the NMR order parameter which are related to the sampled structures of individual molecules. These comparisons show that the structural lipid bilayer properties like thickness and area per molecule are close to real values, however, it seems that there is room for improvement especially in the lipid headgroup

region, in agreement with conclusions from NMR experiments.

When applying lipid bilayer simulations to study complicated biochemical systems it is crucial to recognize potential artefacts arising from the inaccuracies revealed by these comparisons. Hypothetical example of a situation where artificial conclusions might be difficult to avoid would be study of protein approaching PC lipid bilayer in physiological NaCl concentration. If the above mentioned issues are not carefully taken into account one might choose a model where the protein is approaching an effectively positively charged lipid bilayer due to artificial Na⁺ binding with incorrect choline structure. In addition to this the protein might be incorrectly folded already in the bulk water [?].

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- [118] These changes were later shown to be consistent with the addition of different charges into the bilayer, and the electrometer concept was introduced to measure the amount of charge incorporated in the bilayer interface [117].
- [119] This increase is related to the P-N vector tilting more parallel to the membrane plane [16] which is in agreement with electrometer concept suggesting that penetrating charge has opposite effect on headgroup tilt leading to decrease of order parameters [51, 117].
- [120] In this work CH₂/CH₃ groups in cholesterol were changed to LP₂/LP₃ groups to make it more consistent with the Berger parameters. This is not usually done in the studies done with these model.

ToDo

	P.
1. Citations missing	1
2. References should be added, the text should polished, things should be added and checked and figures should be improved. However, the main strucutre and idea of the section should be visible.	2
3. How accurate exactly?	3
4. Maybe specify to which ones?	3
5. Has this been done for the C ₂ carbon in the <i>sn</i> -2 chain?	6
6. The analysis with ions not actually done yet!	9
7. This is the first sketch of this section. A lot of references should be added, the text should polished, things should be added and checked and figures should be improved. However, the main strucutre and idea of the section should be visible.	11
8. I do not have correct papers at hand now. I will do this later	15
9. Discussion about related issues can be found at: https://github.com/NMRLipids/NMRLipids_V-Review/issues/1	15
10. The discussion is going in at: https://github.com/NMRLipids/NMRLipids_V-Review/issues/2	
Key questions now:	
Do we get independetly measured Form Factors plotted to the same figure?	
Are there good references for the accuracy?	15
11. More discussion at: https://github.com/NMRLipids/NMRLipids_V-Review/issues/3	
and https://github.com/NMRLipids/NMRLipids_V-Review/issues/4	
The most important question is now to check the discussion about undulations in the literature	15
12. The more detailed discussion can be found at: https://github.com/NMRLipids/NMRLipids_V-Review/issues/5	16