# Quantitative quality of lipid-cholesterol interactions in atomistic resolution molecular dynamics simulations

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The quantitative quality of lipid-cholesterol interactions in atomistic resolution models will be determined against NMR and scattering data.

### INTRODUCTION

The order parameter changes as a function of cholesterol for each segment are shown in Fig. 2 (currently only sn-1).

The quantitative quality of lipid-cholesterol interactions in atomistic resolution models will be determined against NMR and scattering data.

### **METHODS**

### X-ray scattering experiments

This folder contains SAXS data on POPC multilamellar vesicles (MLVs) at various cholesterol concentrations. Data have been obtained at the EMBL BioSAXS beamline (Hamburg) using 20 keV photons, T = 27C. Data were analyzed in terms of the SDP-GAP model described in Heftberger et al., J. Appl. Cryst. 2013 and Heftberger et al. Biophys. J. 2015. Data from MLVs are a convolute of structure factor (the crystalline lattice) and form factor. By fitting the scattered intensity data we obtain both contributions. Here we posted only form factors (ASCII format). For information on the quality of the fit we also give plots of the fitted intensity data. The electron density profile has been modelled in terms of the SDP model (see papers by Kucerka and coworkers), that is volume distribution functions are modelled by individual Gaussians or error functions. Cholesterol is also accounted for by two Gaussians. This model has been proposed by Jianjun Pan (USF, Tampa, FL), but is to the best of our knowledge not published (see also PhD Thesis by Peter Heftberger). Additional figures show the volume distribution functions and the resulting electron density profiles.

Authors to consult and potentially include in publications using this data: Peter Heftberger (peter.heftberger@gmx.at), Georg Pabst (georg.pabst@uni-graz.at)

## Molecular dynamics simulations

# COMPARISON OF ACYL CHAIN ORDER PARAMETERS BETWEEN EXPERIMENTS AND SIMULATIONS

Order parameters from simulations and experiments for acyl chains of 1-palmitoyl-2-oleoylphosphatidylcholine (POPC) are shown in Fig. 1.

TABLE I: Simulated lipid bilayers containing cholesterol. The simulation file data sets marked with  $^*$  include also part of the trajectory.  $^a$  The number of lipid molecules  $^b$  The number of cholesterol molecules  $^c$  Cholesterol concentration (mol%)  $^d$  The number of water molecules  $^e$  Simulation temperature  $^f$  The total simulation time  $^g$  Time frames used in the analysis  $^h$  Reference link for the downloadable simulation files  $^i$  Reference for the full simulation details

lipid	$^a\mathrm{N_l}$	$^b\mathrm{N}_\mathrm{chol}$	$^{c}\mathrm{C}_{\mathrm{CHOL}}$	$^d\mathrm{N_w}$	eT(K)	$^f t_{\rm sim}({\rm ns})$	$g_{t_{anal}}$ (ns)	$^h\mathrm{Files}$	$^i$ Details
POPC	128	0	0%	7290	298	270	240	[2]*	[3]
POPC	120	8	6%	7290	298	100	80	[6]*	[5]
POPC	110	18	14%	8481	298	100	80	[7]*	[5]
POPC	84	44	34%	6794	298	100	80	[8]*	[5]
POPC	64	64	50%	10314	298	100	80	[9]*	[5]
POPC	50	78	61%	5782	298	100	80	[10]*	[5]
POPC	128	0	0%	5120	303	150	100	[13]*	[14]
POPC	100	24	19%	4960	303	200	100	[15]*	[14]
POPC	80	80	50%	4496	303	200	100	[16]*	[14]
POPC	128	0	0%	6400	310	400	200	[18]*	[14]
POPC	114	14	11%	6400	310	400	200	[18]*	[14]
POPC	72	56	44%	6400	310	400	200	[18]*	[14]
POPC	64	64	50%	6400	310	400	200	[18]*	[14]
POPC	56	72	56%	6400	310	400	200	[18]*	[14]
	POPC POPC POPC POPC POPC POPC POPC POPC	POPC 128  POPC 120  POPC 110  POPC 84  POPC 64  POPC 50  POPC 128	POPC 128 0 POPC 120 8 POPC 110 18 POPC 84 44 POPC 64 64 POPC 50 78 POPC 128 0 POPC 100 24 POPC 80 80 POPC 128 0 POPC 128 0 POPC 128 0 POPC 128 0 POPC 14 14 POPC 72 56 POPC 64 64	POPC 128 0 0%  POPC 120 8 6%  POPC 110 18 14%  POPC 84 44 34%  POPC 50 78 61%  POPC 128 0 0%  POPC 100 24 19%  POPC 80 80 50%  POPC 128 0 0%  POPC 128 0 0%  POPC 128 0 44%  POPC 50 56 44%  POPC 64 64 50%	POPC 128 0 0% 7290  POPC 120 8 6% 7290  POPC 110 18 14% 8481  POPC 84 44 34% 6794  POPC 64 64 50% 10314  POPC 50 78 61% 5782  POPC 128 0 0% 5120  POPC 100 24 19% 4960  POPC 80 80 50% 4496  POPC 128 0 0% 6400  POPC 114 14 11% 6400  POPC 72 56 44% 6400  POPC 64 64 50% 6400	POPC 128 0 0% 7290 298  POPC 120 8 6% 7290 298  POPC 110 18 14% 8481 298  POPC 84 44 34% 6794 298  POPC 64 64 50% 10314 298  POPC 50 78 61% 5782 298  POPC 128 0 0% 5120 303  POPC 100 24 19% 4960 303  POPC 80 80 50% 4496 303  POPC 128 0 0% 6400 310  POPC 114 14 11% 6400 310  POPC 72 56 44% 6400 310  POPC 64 64 50% 6400 310	POPC 128         0         0%         7290         298         270           POPC 120         8         6%         7290         298         100           POPC 110         18         14%         8481         298         100           POPC 84         44         34%         6794         298         100           POPC 64         64         50%         10314         298         100           POPC 50         78         61%         5782         298         100           POPC 128         0         0%         5120         303         150           POPC 100         24         19%         4960         303         200           POPC 80         80         50%         4496         303         200           POPC 128         0         0%         6400         310         400           POPC 114         14         11%         6400         310         400           POPC 72         56         44%         6400         310         400           POPC 64         64         50%         6400         310         400	POPC 128         0         0%         7290         298         270         240           POPC 120         8         6%         7290         298         100         80           POPC 110         18         14%         8481         298         100         80           POPC 84         44         34%         6794         298         100         80           POPC 64         64         50%         10314         298         100         80           POPC 50         78         61%         5782         298         100         80           POPC 128         0         0%         5120         303         150         100           POPC 100         24         19%         4960         303         200         100           POPC 80         80         50%         4496         303         200         100           POPC 128         0         0%         6400         310         400         200           POPC 114         14         11%         6400         310         400         200           POPC 72         56         44%         6400         310         400         200	POPC 128         0         0%         7290         298         270         240         [2]*           POPC 120         8         6%         7290         298         100         80         [6]*           POPC 110         18         14%         8481         298         100         80         [7]*           POPC 84         44         34%         6794         298         100         80         [8]*           POPC 64         64         50%         10314         298         100         80         [9]*           POPC 50         78         61%         5782         298         100         80         [10]*           POPC 128         0         0%         5120         303         150         100         [13]*           POPC 100         24         19%         4960         303         200         100         [15]*           POPC 80         80         50%         4496         303         200         100         [16]*           POPC 128         0         0%         6400         310         400         200         [18]*           POPC 72         56         44%         6400         310

# COMPARISON OF FORM FACTORS BETWEEN EXPERIMENTS AND SIMULATIONS

The form factors calculated from different simulations with different cholesterol content are shown in Fig. 3.

# CONCLUSIONS

# SUPPLEMENTARY INFORMATION

### CHARMM36 results from different simulation packages

The results from CHARMM36 model for lipid bilayers from different simulation packages have been reported to give different results in the literature [19, 20]. The results are mainly dependent on different Lennart-Jones cut-off settings, but all the details are not quite understood. In this work we use the results from Gromacs 5 with settings suggested to be optimal by Gromacs webpage. We also compared the results from Gromacs 5 with these settings to the results simulated with NAMD, OpenMM and literature values. The comparison is shown in Fig. 4

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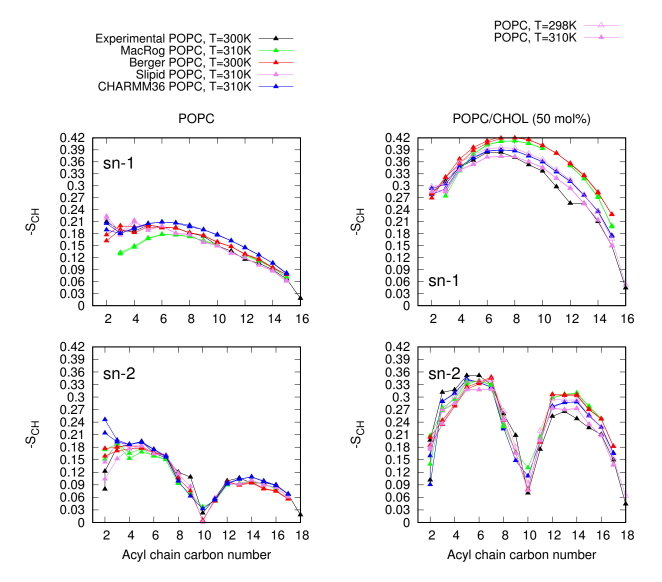


FIG. 1: Order parameters from simulations and experiments for acyl chains of 1-palmitoyl-2-oleoylphosphatidylcholine (POPC).

1. Why are the order parameters of CHARMM36 too large compared to simulations even without cholesterol? Discussion in

https://github.com/NMRLipids/NmrLipidsCholXray/issues/4 2.Why there is decrease in order parameters towards beginning of acyl chain in MacRog model without cholesterol?

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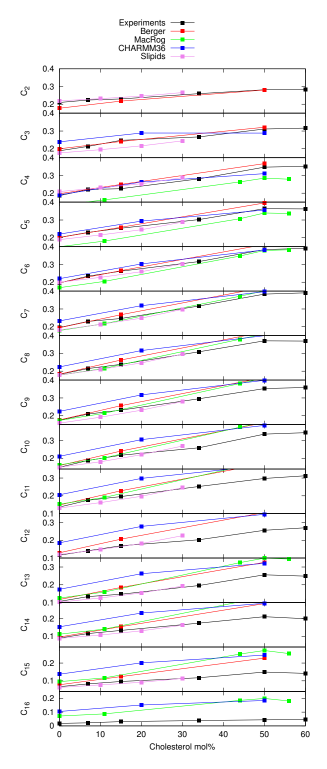


FIG. 2: Order parameter changes from simulations and experiments for each segment in sn-1 chain of 1-palmitoyl-2-oleoylphosphatidylcholine (POPC) as a function of cholesterol concentration.

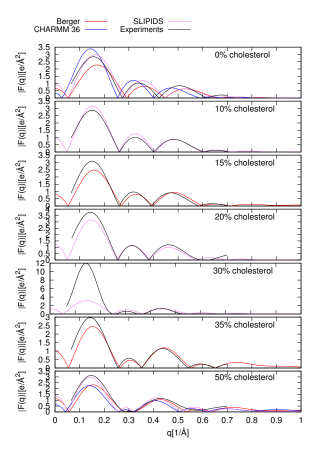


FIG. 3: Form factors from simulations and experiments.

4.Form factor calculation method should be double checked.

5.Experimental form factor amplitudes are not scaled to match with simulations, as done usually

6.Not all experimental and simulation data is here.

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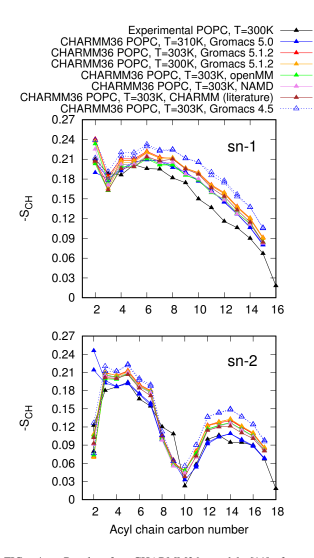


FIG. 4: Results for CHARMM36 model [11] from different simulation packages. Discussion going on at https://github.com/NMRLipids/NmrLipidsCholXray/issues/4.

Róg	(2015),	URL	{http://dx.doi.org/10.5281/
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### **ToDo**

1. Why are the order parameters of CHARMM36 too large compared to simula-	
tions even without cholesterol? Discussion in https://github.com/NMRLipids/NmrLipidsCholXray/issues/4 2. Why there is decrease in order parameters towards	3
beginning of acyl chain in MacRog model without cholesterol?	
3. Do the results suggests that condensation effect is too strong in MacRog and Berger models? Discussion in https://github.com/NMRLipids/NmrLipidsCholXray/issues/5	3
4. Form factor calculation method should be double checked	
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