SUPPLEMENTARY INFORMATION FOR NMRlipids III: Lipid–Cholesterol Interactions in Atomistic Resolution Molecular Dynamics Simulations

Peter Heftberger,[†] Matti Javanainen,^{*,‡,¶} Jesper J. Madsen,^{§,||} Josef Melcr,[‡] Markus Miettinen,^{\perp ,#} O. H. Samuli Ollila,^{*,‡,¶} and Georg Pabst^{†,@}

†Institute of Molecular Biosciences, Biophysics Division, NAWI Graz, University of Graz, Graz 8010, Austria

‡Institute of Organic Chemistry and Biochemistry, Academy of Sciences of the Czech Republic, Prague 6, Czech Republic

 $\P{\it Institute~of~Biotechnology,~University~of~Helsinki}$

 $\S Department\ of\ Global\ Health,\ College\ of\ Public\ Health$

 $\parallel University \ of \ South \ Florida$

 $\perp Department$ of Chemistry, University of Bergen, Norway

#Computational Biology Unit, Department of Informatics, University of Bergen, Norway

@BioTechMed-Graz, Graz 8010, Austria

E-mail: matti.javanainen@helsinki.fi; samuli.ollila@helsinki.fi

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S1 Simulation Details

The force field specific simulation parameters used for each force field are listed in Table S1. The parameter input files (.mdp) are provided in the Zenodo portal (links in the main text). For all simulations, we used an integration time step of 2 fs and the leap-frog integrator of GROMACS. The isothermal–isobaric (NPT) ensemble was used with a temperature of 298 K and a pressure of 1 bar. All simulations were 1 µs long, and trajectories were written every 1 ns. The P-LINCS constraint algorithm^{1,2} was used for the bonds noted in Table S1. NMRlipids databank³ ID numbers of simulations are given in table S2.

Table S1: Simulation parameters used for different force fields.

Neighbour list type $$\operatorname{Verlet}^4$$ Long-range electrostatics $\operatorname{PME}^{5,6}$ LJ cut-off $1.2 \mathrm{nm}$ LJ modifier $-1.2 \mathrm{nm}$ Dispersion correction $-$ Thermostat $\operatorname{Nos\acute{e}-Hoover}^{8,9}$ Time constant (T) $1 \mathrm{ps}$ Coupling groups $-1 \mathrm{ps}$ Coupling type (P) $-1.2 \mathrm{nm}$ Barostat $-1.2 \mathrm{nm}$ Coupling type (P) $-1.2 \mathrm{nm}$ Semi-isotropic $-1.2 \mathrm{ps}$ Compressibility $-1.2 \mathrm{ps}$ Constraints $-1.2 \mathrm{nm}$ Bonds with H	Verlet ⁴ $PME^{5.6}$ 1.4 nm $-$ $Energy & pressure7$ $8.9 Stochastic rescaling10$	[- 건]	Verlet ⁴ PME ^{5,6} 1.0 nm - Energy & pressure ⁷ Stochastic rescaling ¹⁰ 0.1 ps
orrection $\operatorname{Ant}(T)$ $\operatorname{Sups}(P)$ $\operatorname{Ant}(P)$ $\operatorname{Inty}(P)$			1.0 nm Energy & pressure ⁷ Stochastic rescaling ¹⁰ 0.1 ps
orrection ant (T) be (P) ant (P) lity			Energy & pressure ⁷ Stochastic rescaling ¹⁰ 0.1 ps
orrection $\operatorname{ant}(T)$ $\operatorname{sups}(P)$ $\operatorname{ant}(P)$ $\operatorname{ant}(P)$			Energy & pressure ⁷ Stochastic rescaling ¹⁰ 0.1 ps
$\begin{array}{c} \operatorname{ant}\ (T) \\ \operatorname{sups} \\ \operatorname{pe}\ (P) \\ \operatorname{ant}\ (P) \\ \operatorname{lity} \end{array}$			Stochastic rescaling ¹⁰ 0.1 ps
	7 C	1 ps	0.1 ps
	ed 6.0		
	er Lipids & water	Lipids & water	Lipids & water
	nan ¹¹ Berendsen ¹²	Parrinello–Rahman ¹¹	Parrinello-Rahman ¹¹
	ic semi-isotropic	semi-isotropic	semi- $isotropic$
	10 ps	5 ps	4 ps
Bond	ar $4.5 \cdot 10^{-5} \text{ 1/bar}$	$4.5 \cdot 10^{-5} \text{ 1/bar}$	$4.5 \cdot 10^{-5} \text{ 1/bar}$
	H All bonds	Bonds with H	All bonds
Water model TIPS3P ¹³	$ ext{TIP}3 ext{P}^{14}$	${ m TIP3P^{14}}$	${ m TIP3P^{14}}$
FF Source CHARMM-GUI	UI CHARMM-GUI	CHARMM-GUI	Refs. 15 & 16
Parameter source CHARMM-GUI	UI Slipids website	CHARMM-GUI	Refs. 15 & 16

Table S2: NMR lipids databank IDs of performed simulations.

			${f Slipids}$		
CHARMM36			System		ID
System		ID	Slipids_POPC_CHOL20	L	691
1024POPC_128CHOL_57600SOL_	298K	426	Slipids_POPC_CHOL47	_M	712
64POPC_3200SOL_298K		546	Slipids_POPC_CHOL29	_S	693
1024POPC_896CHOL_96000SOL_	298K	543	Slipids_POPC_CHOL20	_S	692
64POPC_40CHOL_5200SOL_29	8K	620	Slipids_POPC_CHOL20	_M	687
64POPC_56CHOL_6000SOL_29	8K	91	Slipids_POPC_CHOL11	_S	672
1024POPC_416CHOL_72000SOL_	298K	412	Slipids_POPC_CHOL11	_M	681
1024POPC_256CHOL_64000SOL_	298K	88	$Slipids_POPC_L$		696
256POPC_64CHOL_16000SOL_2	98K	72	Slipids_POPC_CHOL38	_S	661
64POPC_8CHOL_3600SOL_298K		109	$Slipids_POPC_M$		708
256POPC_104CHOL_18000SOL_298K		298	Slipids_POPC_CHOL38	_L	670
64POPC_16CHOL_4000SOL_29	8K	525	$Slipids_POPC_S$		664
64POPC_26CHOL_4500SOL_29	8K	393	Slipids_POPC_CHOL29	_L	709
1024POPC_640CHOL_83200SOL_	298K	550	Slipids_POPC_CHOL29	_M	659
256POPC_160CHOL_20800SOL_2	298K	275	Slipids_POPC_CHOL11	_L	697
256POPC_224CHOL_24000SOL_2	298K	166	Slipids_POPC_CHOL38	$_{ m M}$	668
256POPC_32CHOL_14400SOL_298K		119	Slipids_POPC_CHOL47	_S	703
			Slipids_POPC_CHOL47	_L	682
Lipid17			${f MacRog}$		_
\mathbf{System}	ID		\mathbf{System}	ID	
Lipid17_POPC_CHOL29_M	700	Mac	Rog_POPC_CHOL38_S	716	_
Lipid17_POPC_CHOL47_L	717	Mac	Rog_POPC_CHOL29_L	705	
Lipid17_POPC_CHOL11_L	663	MacI	Rog_POPC_CHOL47_M	699	
Lipid17_POPC_CHOL29_L	683	Mac	Rog_POPC_CHOL47_L	671	
Lipid17_POPC_CHOL20_S	662	Mac	Rog_POPC_CHOL20_L	677	
Lipid17_POPC_CHOL47_M	714		MacRog_POPC_S	674	

System	ID	System	ID
Lipid17_POPC_CHOL29_M	700	MacRog_POPC_CHOL38_S	716
Lipid17_POPC_CHOL47_L	717	MacRog_POPC_CHOL29_L	705
Lipid17_POPC_CHOL11_L	663	MacRog_POPC_CHOL47_M	699
Lipid17_POPC_CHOL29_L	683	MacRog_POPC_CHOL47_L	671
Lipid17_POPC_CHOL20_S	662	MacRog_POPC_CHOL20_L	677
Lipid17_POPC_CHOL47_M	714	MacRog_POPC_S	674
Lipid17_POPC_CHOL38_S	688	MacRog_POPC_CHOL20_M	706
Lipid17_POPC_L	684	MacRog_POPC_CHOL38_L	698
Lipid17_POPC_S	715	MacRog_POPC_CHOL38_M	679
Lipid17_POPC_CHOL20_M	666	$MacRog_POPC_M$	675
Lipid17_POPC_M	657	$MacRog_POPC_L$	658
Lipid17_POPC_CHOL29_S	686	$MacRog_POPC_CHOL29_M$	655
Lipid17_POPC_CHOL38_L	673	MacRog_POPC_CHOL11_L	660
Lipid17_POPC_CHOL47_S	707	MacRog_POPC_CHOL11_S	665
Lipid17_POPC_CHOL11_S	680	$MacRog_POPC_CHOL29_S$	690
Lipid17_POPC_CHOL20_L	667	$MacRog_POPC_CHOL11_M$	695
Lipid17_POPC_CHOL38_M	694	$MacRog_POPC_CHOL47_S$	704
Lipid17_POPC_CHOL11_M	689	$MacRog_POPC_CHOL20_S$	702

S2 Additional Results

S2.1 Scattering Intensities

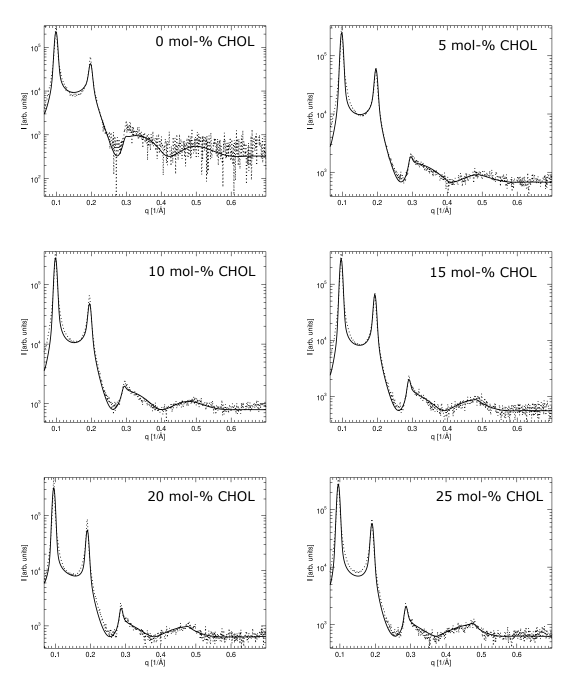


Figure S1: Scattering intensities from X-ray scattering experiments with various concentrations of cholesterol. More data are shown in the next figure.

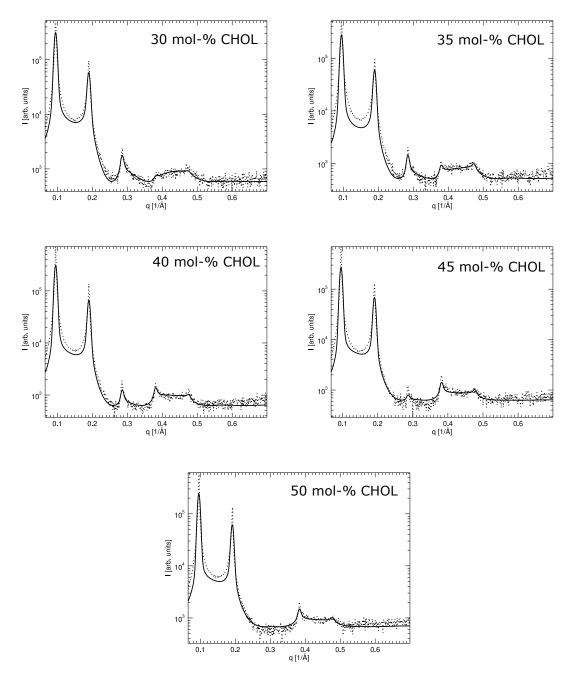


Figure S2: Scattering intensities from X-ray scattering experiments with various concentrations of cholesterol. More data are shown in the previous figure.

S2.2 Form Factors & Electron Density Profiles

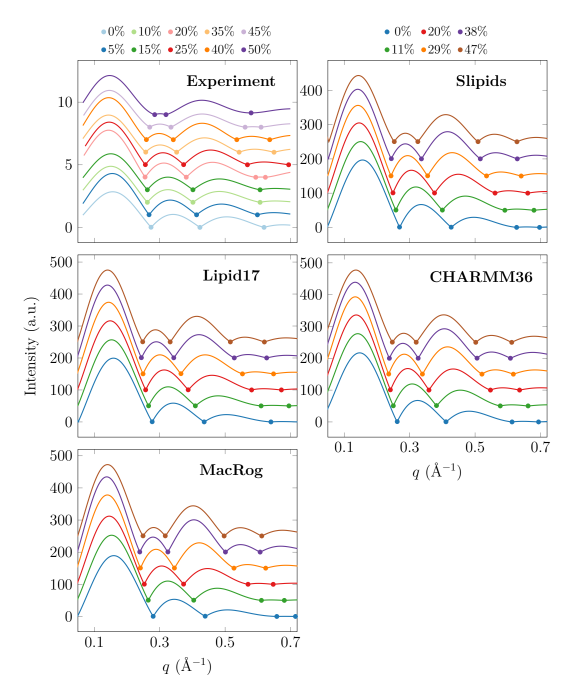


Figure S3: Scattering intensities as a function of scattering vector from experiments and simulations. Each of the profiles is shifted vertically with respect to the previous one, by 1 for the experimental profiles and by 50 for the computational ones. The minima are marked by filled circles to guide the eye.

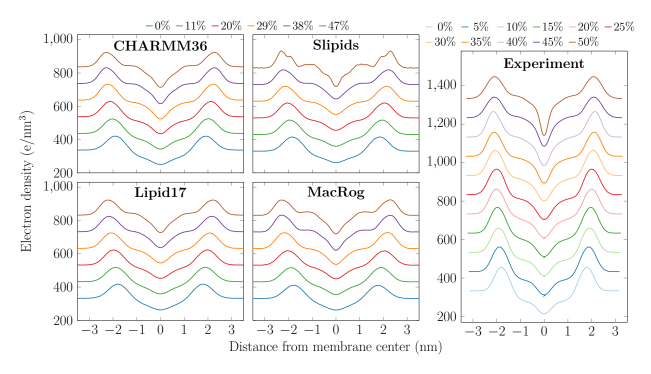


Figure S4: **Electron density profiles.** Each of the profiles is shifted vertically with respect to the previous one by 100 units.

S2.3 Order Parameters

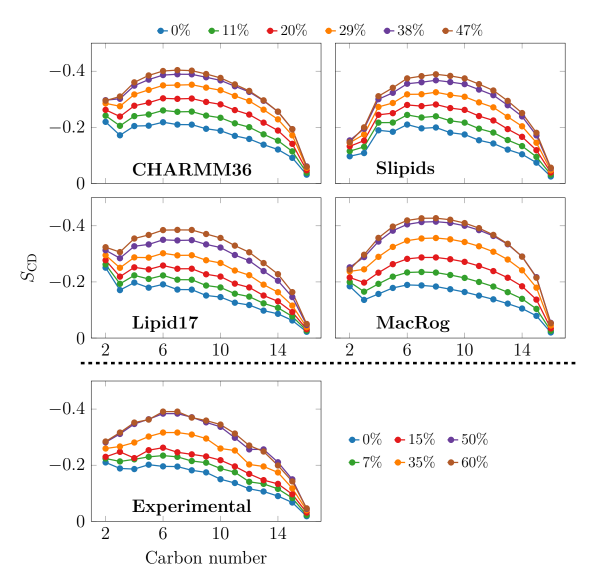


Figure S5: Effect of cholesterol on the acyl chain order parameters of the POPC sn-1 (palmitate) chain. The legend at the top corresponds to all simulations, and the one at the bottom to the experiments.

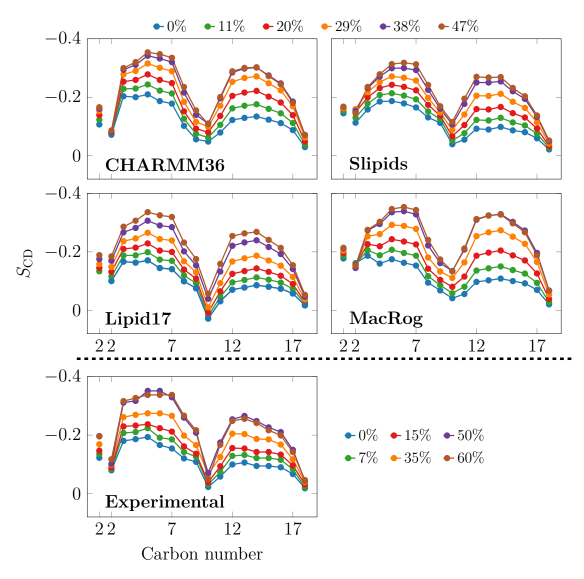


Figure S6: Effect of cholesterol on the acyl chain order parameters of the POPC sn-2 (oleate) chain. The legend at the top corresponds to all simulations, and the one at the bottom to the experiments. Since the order parameters measured for the two hydrogens bound to the C2 carbon differ, they are both shown in the plots.

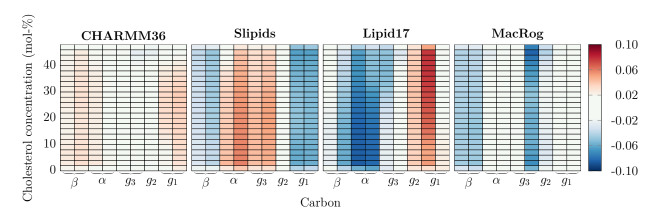


Figure S7: The deviation of POPC head group parameters from experimental values as a function of CHOL concentration.

S2.4 Dynamic Properties

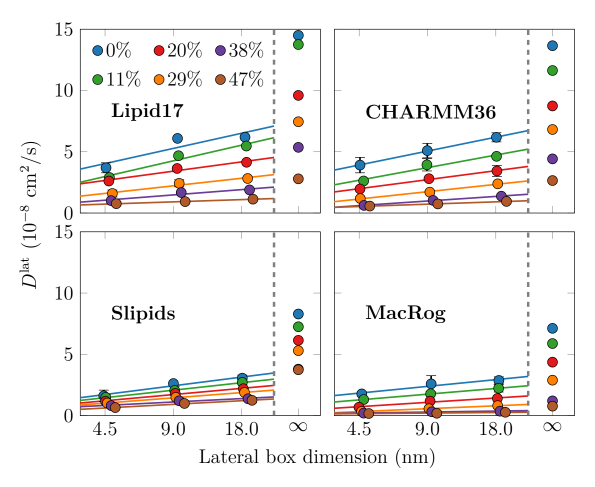


Figure S8: Dependence of lateral diffusion coefficients on simulation box size. The values calculated for the lipid centres of mass with gmx msd after eliminating leaflet drift. The values for the three system sizes are shown as markers together with fits of Eq. (2). The values extrapolated to infinite system sizes are also shown in the separate column.

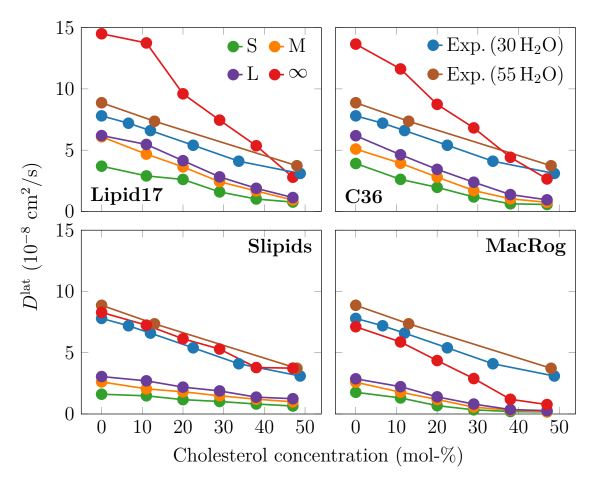


Figure S9: Dependence of lateral diffusion coefficients on cholesterol concentration. Data are shown for all system sizes; small (S), medium (M), and large (L). The values extrapolated to infinity are shown as well (∞). Experimental measured at two hydration levels, 30 m-% and 55-% of water. ^{17,18}

S2.5 Finite-Size Effects

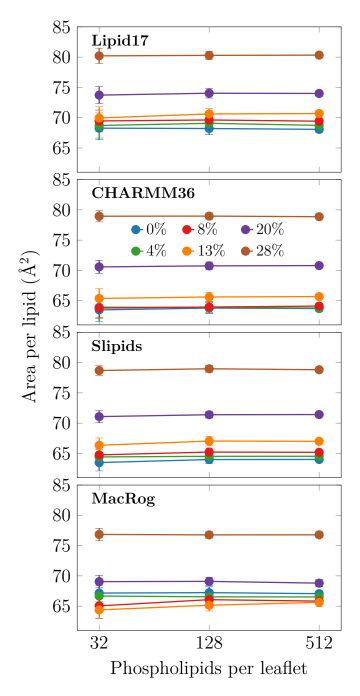


Figure S10: Dependence of area per lipid on simulation box size. Area per lipid is calculated by dividing the membrane area by the number of lipids in one leaflet. Error bars show standard error extracted using block averaging in gmx analyze.

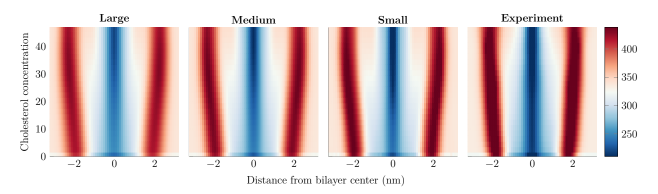


Figure S11: Effect of system size on the density profiles. Membrane undulations are larger in the larger systems, which leads to the smearing of the electron density profiles. Here, data are shown for CHARMM36 in the large (1024 POPC in total), medium (256 POPC in total), or small (64 POPC in total) systems. The experimental electron density profile is shown for comparison.

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