NMRIipids Databank: Overlay Databank of Lipid Membrane Simulations Arising from Open Collaboration

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ABSTRACT

We present a databank of lipid bilayer simulations from the NMRlipids open collaboration project.

1 Introduction

The importance of sharing MD simulation data following the FAIR principles¹ has been widely recognized^{2–9}, and databanks are emerging^{9–16}. The relevance of quality evaluation of simulation trajectories in databanks regarding technical details of simulations and accuracy of the underlying physical description of the system (force field) has become evident^{3,7,10} and such quality evaluation has in some cases also been implemented^{10,12}. However, straightforward quality comparisons between individual simulations or force fields within these databanks remain challenging. While importance of such databanks for MD simulations is widely recognized^{2–9} and different kinds of approaches are emerging^{9–16}, generally accepted protocols and best practices are still under active development.

Here we present a solution for lipid bilayers based on overlay databank structure illustrated in Fig. 1. The concept of overlay databank is developed here to solve the practical challenges in generating databanks of MD simulation data enabling flexible analyses over large sets of simulation data, but it can be potentially used for wide range of situations. Particularly when storage of raw data requires significant resources and final outcomes or best practices are not yet clear, overlay databank approach lowers the barrier to start without compromizing the long term stability or scalability.

The practical relevance of the NMRlipids databank is exemplified by automatic quality evaluation and ranking of large amount of MD simulation data, data-driven analysis detecting correlations between properties of model cell membranes and analyses of rare phenomena that are beyond the scope of standard MD simulation studies. The NMRlipids databank provides new tools for researchers in wide range of fields in academia and industry from cell membrane biology to lipid nanoparticle formulations and data-driven computational chemistry and machine learning.

2 Results

2.1 Overlay structure of the NMRlipids databank

The three layer structure of the NMRlipids databank, illustrated in Fig. 1 a, enables an efficient way to share and upcycle large MD simulation trajectories. Layer 1 contains raw simulation data that can locate in any publicly available repository or server offering long term stability and permanent links to the data, such as digital object identifiers. Layer 2 contains all the relevant information on the simulations, including links to the raw data in layer 1 and relevant metadata describing the systems. Layer 3 is the application layer containing outputs from the databank, such as graphical user interface (http://www.databank.nmrlipids.fi/) and results from analyses desribed in sections below.

Layer 2 is the core of the databank. In addition to the metadata and links to raw simulation data, it contains universal naming conventions for the molecules and atoms in the databank, computer programs to generate and analyse the databank, basic properties calculated from all simulations (area per lipid, C-H bond order parameters, x-ray scattering form factors and membrane thickness), and quality evaluation of simulations against experimental data. In practise, this information is stored is the git repository which is currently available at https://github.com/NMRLipids/Databank. Detailed description on its content is in the supplementary information. Applications in layer 3 uses the information in layer 2 to access raw data in layer 1 to perform automatic analyses over large sets of simulations in the databank.

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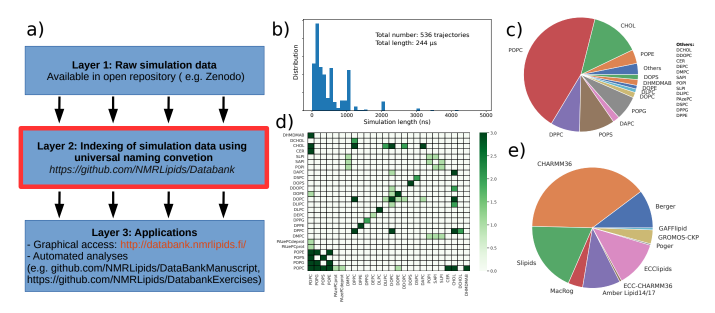


Figure 1. a) Structure of an overlay databank. More detailed structure of the layer 2 in the NMRlipids databank is illustrated in Fig. 5 in the SI. b) Distribution of the lengths of the trajectories, total number of trajectories and total length of the simulations in the NMRlipids databank. c) Distribution of lipids present in the trajectories in the NMRlipids databank. Lipids occurring in five or less simulations ('others') are listed in the right. d) Currently available binary mixtures in the NMRlipids databank. e) Distribution of force fields in the simulations in the NMRlipids databank. The figures and numbers are created on 9th of May 2022.

2.2 Content of the databank

Currently the databank is composed of approximately 500 trajectories with the total length of approximately 231 microseconds of which most are contributed for the previous publications from the NMRlipids open collaboration^{17–20}. The distribution of lipids, force fields, length of the trjajectories and available binary mixtures are shown in Fig. 1.

2.3 Quality evaluation of force fields

Quality of membrane simulations with different force fields have been evaluated against experimental data during parameterization and in separate comparison studies ^{17,18,21–24}, but universal quality measure for membrane simulations is not defined and controversial results are often reported from simulations²⁵. The lack of universal quality measure for membrane simulations complicates the selection of proper simulation parameters and the estimation of reliability in simulations, thereby being a major obstacle in many applications of membrane MD simulations. To enable the rapid quality evaluation of membrane MD simulations, a quantitative quality measure based on C-H bond order parameters from NMR experiments is defined in the NMRlipids databank. This is complemented by comparing the locations of x-ray scattering form factor minima against experiments. Both of these are robust experimental measurables that can be directly connected to the simulation data²¹. The used quality measures are defined in detail in the supplementary information.

Simulations with the highest scores based on order parameter evaluation for all simulations are shown in Fig. 2 A and for POPE lipids in Fig. 2 B. The direct comparison with experiments for the best simulation, POPC with OPLS3e, is shown in Fig. 2 C, where discrepancies with experiments are observed only for the third carbon in the *sn*-1 chain, and carbons 8 and 9 close to the double bond in *sn*-2 chain. The best overall quality for POPE is found in GROMOS-CKP simulation, for which the direct comparison in Fig. 2 D show minor differences in acyl chains, but major discrepancies in glycerol backbone and in the first carbon of *sn*-2 chain. These parts are better described for POPE by CHARMM36 parameters (direct comparison shown in Fig. 2 E), but its overall quality suffers from the overestimated order in acyl chains.

2.4 Finding the best models for PC and PE lipid mixtures

The power of the NMRlipids databank in finding the best models for a certain lipid bilayer system is exemplified here for mixtures of PE and PC lipids. Area per lipids from available POPC and POPE bilayer simulations in the databank from different force fields at 310 K are shown in Fig. 3. Because the area per lipid strongly correlates with the average order of *sn*-1 chain (Fig. 2 F), we use order parameter qualities to evaluate which simulations predict the best area per lipids for POPC and POPE. Among the force fields with area per lipids shown in Fig. 3, Slipids ranks 1st for POPC with a clear difference to others, and 2nd

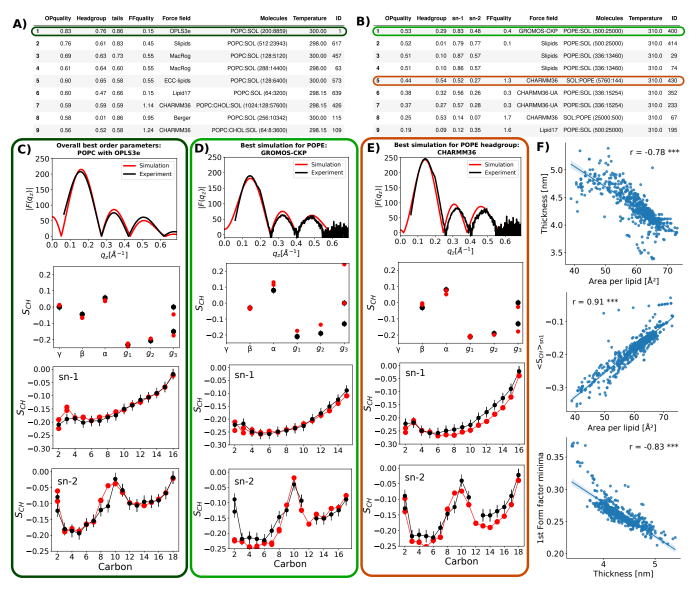


Figure 2. A) Best simulations ranked based on overall order parameter quality. B) Best simulations ranked based on the overall order parameter quality of POPE lipid. C) Scatter plots and Pearson correlation coefficients for the area per lipid, thickness, and experimental observables. All correlation coefficients have p-value below 0.001. For more correlations see Fig. S1. D)-F) Evaluation against experimental data exemplified for a simulation with the best overall order parameter quality (D), the best quality for POPE lipid (E), and the headgroup quality for POPE (F).

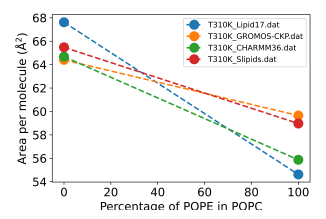


Figure 3. Differences in area per lipids of POPC and POPE bilayers between different force fields.

for POPE with only marginally lower quality than GROMOS-CKP (Figs. 2 A and B). The direct comparison with experiments in Fig. S3 in the supplementary information reveals that GROMOS-CKP and CHARMM36 predicts too ordered (too low area per lipid) bilayer for POPC, while the order parameters in Lipid17 are lower than in experiments indicating overestimated area per lipid. For POPE, order parameters in Slipids and GROMOS-CKP are close to experiments suggesting that their are per lipids are most reasonable, while CHARMM36 and Lipid17 are too packed thereby overestimating the order parameters.

In conclusion, the quality evaluation based on the NMRlipids databank suggests that the Slipids parameters would give the most reliable description for membrane packing of POPC:POPE mixtures. Slipids was a relatively good model also for POPC:POPS mixtures in similar comparisons utilizing the preliminary version of the NMRlipids databank²⁵, although charged membranes are more complicated as the counterion binding affinity plays a significant role in membrane packing¹⁹. On the other hand, CHARMM36 overcomes Slipids when the focus is on conformational ensembles of headgroups in different types of lipids because the accuracy of glycerol backbone is low in Slipids²⁰. Altogether, these examples demonstrate how the NMRlipids databank can be used to navigate in the complex landscape of lipid force field quality.

2.5 Water diffusion anisotropy in membrane systems

The anisotropic water diffusion through and along of membranes plays a role in the drug translocation through biological material, particularly in skin^{?,26–29}, and in MRI imaging[?]. Therefore, dependence of water permeation through membranes and translational diffusion at membrane surfaces on membrane properties has been studied with different methods, yet consensus has not been reached despite of several available models^{?,?,?,30,31}. MD have been also used in these endeavours, but systematic studies are limited by the accessible amount of data as only few water permeation events are typically observed in a single MD simulation trajectory[?]. Therefore, this serves as an excellent example where the large amount of available MD simulations in the NMRlipids databank can be utilized to answer important biological questions.

To this end, we first calculated the water permeability through membranes from all the simulations in the NMRlipids databank. Resulting permeabilities are shown as a function of temperature, membrane thickness, area per lipid, and acyl chain order in Figs. 4 B-E. The values for water permeabilities from simulations vary between 0.3 and 322 μ m/s with the mean and median of 14 μ m/s and 8 μ m/s, respectively. These values have the same order of magnitude as the experimentally determined diffusive permeability coefficients, but are on average below the values reported for PC lipids in liquid crystalline phase, 190-330 μ m/s³². To analyze how water permeability through membranes depend on these properties on average, we calculated the average permeabilities over all systems with the value of temperature, thickness, area per lipid, or acyl chain order within a fixed range. These averaged permeabilities also shown in Figs. 4 with black dots. As expected, the permeability increases with the temperature, giving $17 \pm 4 \, k_b T$ for the average energy barrier for the water permeation from the Arrhenius plot in Fig. B. Water permeability decreases with increasing membrane packing, i.e., with decreasing area per lipid and increasing thickness and acyl chain order. Linear dependence of permeability on area per lipid and thickness is previously reported from simulations and experiments for some systems, but not for all^{2, 2}. Our results suggest on average linear dependence for thicknesses above $\sim 3.9 \, \text{nm}$ and area per lipids below $\sim 69 \, \text{Å}^2$ (insets in Figs. 4 C-D), yet stronger dependence for more loosely packed membranes. Clear dependencies of permeability on charged lipids, cholesterol, POPE, or hydration level were not observed in Fig. S4 in the supplementary information.

To analyze the anisotropic diffusion of water in membrane environment, we also calculated the water translational diffusion along the membrane surface from all simulations in the NMRlipids databank. The results at different hydration levels are shown

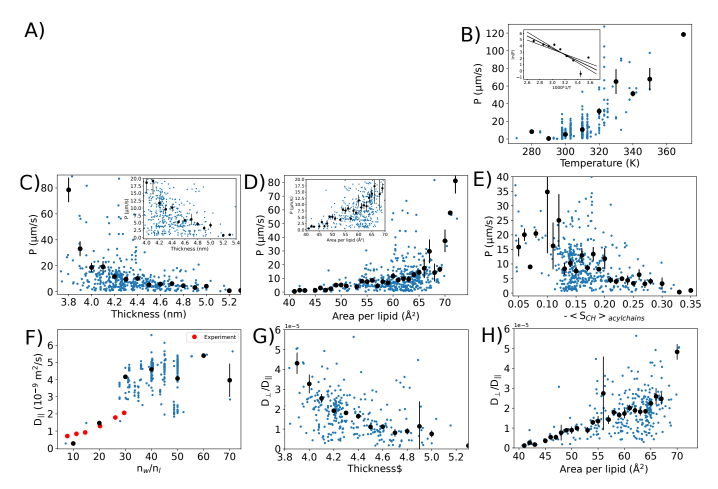


Figure 4. A **B-E** Water permeation through membranes analyzed from the databank as a function of temperature, thickness, area per lipid, and acyl chain order. Values from simulations with non-zero permeation values are shown with blue dots. Averages over fixed range of x-axis values are shown with black dots. Insert in B) shows the Arrhenius plot of permeation (ln (P) vs. 1/T) that gives $17 \pm 4 \, k_b T$ for the average activation energy for water permeation through lipid bilayer. Inserts in C) and D) show the region where the dependence could be considered approximately linear. **F** Lateral diffusion of water as a function of hydration level. Experimental points for DMPC bilayers at 313 K at different hydration levels are shown³³. **G-H** Diffusion anistoropy of water as a function of thickness and area per lipid.

in Fig. 4 F together with the experimental data³³. Simulation results are close to the experimental values at low hydration levels, but increase to the values of approximately two times higher than experimental bulk water diffusion value $(3.1 \cdot 10^{-9} \text{ m}^2/\text{s})$ at high hydration levels. This is not surprising as the most common water model used in membrane simulations, TIP3P, overestimates the bulk water diffusion? The water translational diffusion incereased with the temperature, but dependencies on membrane area per lipid, thickness, or fraction of charged lipids were not observed in Fig. S5 in the supplementary information. In order to estimate the diffusion anistoropy of water in multilamellar membrane system, we translated the water permeation coefficients through membranes to diffusion coefficients through multilamellar stack using the Tanner equation? The resulting diffusion coefficients through the membranes are approximately five orders of magnitude slower than along the membrane (Figs. 4 G and H), which is at the upper limits of the anisotropic estimated from the experimental data²⁸. Relatively high anisotropy is understandable as simulations give slightly slower permeation rates and higher lateral diffusion rates than experiments. Significant increase in the diffusion anisotropy with membrane packing is observed, as $D_{\perp}/D_{||}$ drifts away from one with decreasing area per lipid and increasing thickness in Figs. 4 G and H. This follows from decreasing water permeability with membrane packing (Figs 4 C and D), while lateral diffusion remains approximately constant (Fig. S5 A and C).

3 Discussion

, thereby not requiring large resources to handle and maintain. This lowers the barrier for starting such databank as well as for long term storage. The NMRlipids databank is essentially a git repository containing information on the location of raw data

NMRlipids databank

https://github.com/NMRLipids/Databank/

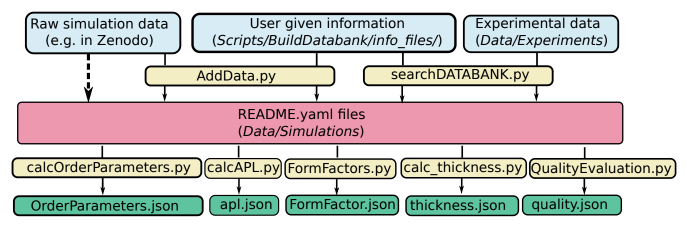


Figure 5. Structure of the NMRlipids databank. Manually added input data (blue boxes) includes basic information on the simulation, permanent links to the raw data, and experimental data if available. The databank entries (red box) and analysis results (green boxes), locating at

https://github.com/NMRLipids/Databank/tree/main/Data/Simulations are automatically generated by the computer programs included in the NMRlipids databank (yellow boxes). Because raw data are not permanently stored but can be accessed based on the information in the databank, this connection is marked with the dashed line.

and its indexing with universal naming convention. In addition to all computers where the databank is developed and used, the NMRlipids databank git is stored to Zenodo server, thereby enabling a very cost effective long term storage for the databank.

Quality measure and automatic quality evaluation of lipid bilayer MD simulations introduced in the NMRlipids databank enables rapid ranking of available simulation models agains experimental NMR and x-ray scattering data. This provides a tool for researches to rapidly evaluate the credibility of MD simulations of their own and reported by other groups. Because such tool and quality measure has not been available, this will elaborate the quality of published MD simulations of lipid bilyers and reduce potentially misleading results²⁵. The power of NMRlipids databank to select the best models for particular applications has been demontrated for PC/PE lipid mixtures (Fig. 2), PC/PS lipid mixtures²⁵, and lipid headgroups²⁰.

The increasing amount of MD simulation data with programmatic access in the NMRlipids databank opens up possibilities for wide range of applications utilizing the large set of accessible data. Extend of the data in the NMRlipids databank in terms of quantity (e.g., simulation length and number of conformations), content (e.g, lipid compositions and ion concentrations) and quality enables analyses that are not possible to conduct from MD simulation data produced by a single research group. Applications of the NMRlipids databank to understand how diffusion of water through and along cellular membrane depends on its physical properties are demonstrated in Fig. ??. Permeation of water through membranes resembles the permeation of also other hydrophilic molecules, such as drugs, and detailed understanding of water dynamics through and along membranes is potentially useful for the development of MRI imaging methods³⁵. These examples demonstrate the practical applications of NMRlipids databank on problems in the biological and biomedical sciences.

Building accessible databanks of molecular dynamics simulation data has been challenging due to the required long term support for hardware and software maintenance. In the overlay model used in the NMRlipids databank, the demand of hardware can be distributed and open collaboration model reduces the risk for ending software maintenance. Furthermore, the open collaboration model used in the NMRlipids project credits contributors by offering authorship in published articles, thereby creating an incentive for contributions. This model could be extended also to other fields where similar barriers to establish publicly accessible databanks exist. Emerging applications of machine learning are increasing the impact of such databanks. For example, the existing Protein Databank (PDB)² containing experimentally determined protein structures with programmatic access has enabled the development of machine learning based tools building on the data collected in the databank over the years². NMRlipids and other databanks with open programmatic access have potential to lead similar unforeseen applications in the future.

4 Methods

4.1 Structure of the databank

Structure of the NMRlipids databank is illustrated in Fig. 5. The required input information to create an entry into the NMRlipids databank are listed in table 1. While the raw simulation data is not directly stored in the NMRlipids databank, permanent links from where the raw data can be accessed have to be given and are then stored in the README.yaml files at https://github.com/NMRLipids/Databank/tree/main/Data/Simulations. These files contain all the essential information listed in table 1 on each simulation entry that are needed for further applications. The raw MD simulation data can locate in any stable publicly available repository, although all the current data locates in Zenodo www.zenodo.org.

In order to evaluate the quality of simulations, sets of C-H bond order parameters from NMR and from factors from x-ray scattering are included in the NMRlipids databank (https://github.com/NMRLipids/Databank/tree/main/Data/experiments). The required information for an experimental dataset are listed in table 2. A simulation is connected to a experimental data set if molar concentrations of all molecules are within ± 5 percentage units, charged lipids have the same counterions, and temperature is within ± 2 degrees. In such cases, a simulation and experimental data are paired by adding the path to the experimental data into the simulation README.yaml file.

Because README.yaml contains all the essential information on each simulation, arbitrary analyses can be automatically performed over all the simulations in the NMRlipids databank. In addition to the order parameters for each C-bonds and x-ray scattering form factors used in the quality evaluation, the NMRlipids databank contains the area per lipid and thickness calculated from all simulations in the databank. These results are stored in the same folders as the README.yaml files in https://github.com/NMRLipids/Databank/tree/main/Data/Simulations.

4.2 Molecule and atom naming convention

Because universal convention for lipid molecules and atoms therein has not been defined, the naming conventions vary between authors and force fields. To enable automatic analyses over large sets of simulation in the NMRlipids databank, we have defined unique naming conventions for lipid molecules and atoms. The abbreviations of molecule names used in the NMRlipids databank are listed in table 3. The unique atom names for each molecule and corresponding names in each simulation are defined using mapping files introduced in the NMRlipids project (https://nmrlipids.blogspot.com/2022/04/new-yaml-format-of-mapping-files.html).

Molecule and atom names in each simulation are connected to the unique naming convention with the COMPOSITION dictionary in the README.yaml file. Upon addition of a new entry in the databank, the molecule names in the simulation corresponding the unique names and mapping files (available at https://github.com/NMRLipids/Databank/tree/main/Scripts/BuildDatabank/mapping_files) are defined in the dictionary. The numbers of molecules in the system are then automatically calculated by the NMRlipids databank codes (AddData.py in Fig. 5) and stored in the README.yaml together with other content of the COMPOSITION dictionary. This information can be then used to find each molecule and atom from each simulation in the analysis codes.

4.3 Quality evaluation

4.3.1 Conformational ensembles using C-H bond order parameters

The quality of conformational ensembles of individual lipid molecules in simulations are evaluated using the C-H bond order parameters which can be directly compared with robust experimental data²¹. The order parameters are defined as

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

where θ is the angle between the C–H bond and the membrane normal and angular brackets denote the ensemble average, i.e. average over all sampled configurations of all lipids in a simulation. Because conformational ensembles of individual lipids are independent in a simulation of a fluid lipid bilayer, the number of independent sample points for each C-H bond, n, equals the number of lipids in a simulation. Denoting the S_{CH} from Eq. 1 as the sample mean and s its variance calculated over individual lipids, $\frac{S_{\text{CH}} - \mu}{s/\sqrt{n}}$ has a Student's t-distribution with n-1 degrees of freedom, where μ is the real mean of the order parameter. Because lipid bilayer simulations contain at least dozens of lipids, the Student's t-distribution could be safely approximated with a normal distribution. However, when calculating the probability for a simulation to locate within experimental order parameters, the normal distribution gives values below the numerical accuracy of computers for simulations values far from experiments. To avoid such numerical instability, we use the first order Student's t-distribution. Therefore, we estimate the

¹For example, if you upload 100-200 ns part of total 200 ns simulation, this should value should be 100.

²For example, if you upload 0-200 ns part of total 200 ns simulation where the first 100 ns should be considered as an equilibration, this value should be 100.

key	description	type
DOI	DOI from where the raw data is found	user given (compulsory)
SOFTWARE	Software used to run the simulation (e.g. Gromacs, Amber, NAMD,	
	etc.)	
TRJ	Name of the trajectory file found from DOI	
TPR	Name of the topology file found from DOI (trp file in the case of	
	Gromacs)	
PREEQTIME	Pre-equilibrate time simulated before the uploaded trajectory in	
	nanoseconds. ¹	
TIMELEFTOUT	Equilibration period in the uploaded trajectory that should be dis-	
	carded in analyses. ²	
COMPOSITION	Molecules names used in the simulation and corresponding mapping	
	files (see section 4.2)	
DIR_WRK	Temporary working directory in your local computer.	
UNITEDATOM_DICT	Information for constucting hydrogens for united atom simulations,	
	empty for all atom simulations	
TYPEOFSYSTEM	Lipid bilayer or something else	
PUBLICATION	Give reference to a publication(s) related to the data.	User given (optional)
AUTHORS_CONTACT	Name and email of the main author(s) of the data.	
SYSTEM	System description on free text format	
SOFTWARE_VERSION	Version of the used software	
FF	Name of the used force field	
FF_SOURCE	Source of the force field parameters, e.g, CHARMM-GUI, webpage,	
	citation to a publication, etc.	
FF_DATE	Date when force field parameters were accessed on the gives source	
	(day/month/year).	
FFmolename	Molecule specific force field information, e.g., water model with	
CD-00	FFSOL and sodium parameters with FFSOD.	
CPT	Name of the Gromacs checkpoint file.	
LOG	Name of the Gromacs log file.	
TOP	Name of the Gromacs top file.	
GRO	Name of the Gromacs gro file.	
TRAJECTORY_SIZE	Size of the trajectory file in bytes	automatically extracted data
TRJLENGTH	Lenght of the trajectory (ps).	
TEMPERATURE	Temperature of the simulation.	
NUMBER_OF_ATOMS	Number of atoms in the simulation.	
DATEOFRUNNIG	Date when added into the databank	
EXPERIMENT	Potentially connected experimental data	
COMPOSITION	Numbers of lipid molecules (NPOPC, NPOPG, etc.) per membrane	
	leaflet are calculated by determining on which side of the center	
	of mass of the membrane the center of mass of the head group of	
	each lipid molecule is located. Numbers of other molecules such	
	as solvent and ions (NSOL, NPOT, NSOD, etc.) are read from the	
	topology file.	

Table 1. Keys stored in the README.yaml files of simulations.

key	description
DOI	DOI of the publication related to the experimental data.
TEMPERATURE	Temperature of the experiment.
MOLAR_FRACTIONS	Dictionary of molar fractions of bilayer components
ION_CONCENTRATIONS	Dictionary of ion concentrations of the system
TOTAL_LIPID_CONCENTRATION	Total concentration of lipid components. If exact concentration is not
	known, but experiments are performed in excess water, 'full hydration'
	can be given.
COUNTER_IONS	Type of counter ions if present.

Table 2. Keys stored in the README.yaml files of experiments.

Abbreviation	Molecule name	
POPC	1-palmitoyl-2-oleoyl-sn-glycero-3-phosphocholine	
POPG	1-palmitoyl-2-oleoyl-sn-glycero-3-phosphoglycerol	
POPS	1-palmitoyl-2-oleoyl-sn-glycero-3-phospho-L-serine	
POPE	1-palmitoyl-2-oleoyl-sn-glycero-3-phosphoethanolamine	
CHOL	cholesterol	
DHMDMAB	dihexadecyldimethylammonium	
POT	potassium ion	
SOD	sodium ion	
CLA	chloride ion	
CAL	calcium ion	
SOL	water	

Table 3. Abbreviations used in the databank

probability for the simulation order parameter to locate within experimental error bars from equation

$$P = f\left(\frac{S_{\text{CH}} - (S_{\text{exp}} + \Delta S_{\text{exp}})}{s/\sqrt{n}}\right) - f\left(\frac{S_{\text{CH}} - (S_{\text{exp}} - \Delta S_{\text{exp}})}{s/\sqrt{n}}\right),\tag{2}$$

where f(t) is the first order Student's t-distribution.

The error of $\Delta S_{\text{exp}} = 0.02$ is currently assumed for all experimental order parameters²¹. Because phospholipids sample their conformational ensemble within nanosecond timescale³⁶, all simulations in the databank would be sufficiently long to sample the realistic conformational phase of individual lipids. However, some force fields exhibit too slow dynamics which leads to large error bars in order parameter values³⁷. Because large error bars widen the Student's t-distribution in Eq. 2 thereby artificially increasing the probability to find the simulated value within experimental error bars, the order parameters with error bars larger than the experimental error 0.02 are not included in the quality evaluation.

The qualities of different fragments within each lipid type in a simulation, $P^{\text{frag}}[\text{lipid}]$, are then estimated by averaging the probabilities for individual C-H bond order parameters to locate within experimental errors according to Eq. 2, and dividing the average with the percentage of order parameters for which the quality is available within the fragment, $p_{\text{frag}}[\text{lipid}]$,

$$P^{\text{frag}}[\text{lipid}] = \frac{\langle P[\text{lipid}] \rangle_{\text{frag}}}{p_{\text{frag}}[\text{lipid}]},\tag{3}$$

where frag can be *sn*-1, *sn*-2, headgroup or total (all order parameters within a molecule). The overall quality of different fragments in a simulation are then defined as a molar fraction weighted average over different lipid components

$$P^{\text{frag}} = \sum_{\text{lipid}} \chi_{\text{lipid}} P^{\text{frag}}[\text{lipid}], \tag{4}$$

where χ_{lipid} is the molar fraction of a lipid in the bilayer.

4.3.2 Membrane dimensions with x-ray scattering form factors

While C-H bond order parameters relate to the conformational ensembles of individual lipids, the x-ray scattering form factors depend on membrane dimensions and density distribution²¹. Because experiments give form factors only in relative intensity

scale, they are typically scaled to the simulation data. Here we use the same scaling method as in SIMtoEXP program³⁸ where experimental form factor intensities are scaled by a factor defined as

$$k_e = \frac{\sum_{i=1}^{N_q} \frac{|F_s(q_i)||F_e(q_i)|}{(\Delta F_e(q_i))^2}}{\sum_{i=1}^{N_q} \frac{|F_e(q_i)|^2}{(\Delta F_e(q_i))^2}},$$
(5)

where F_s and F_e are form factors from a simulation and experiment, respectively, and summation goes over the experimentally available N_q points. Quality measure based on differences in simulated and experimental form factors accross available q-range is also defined in the SIMtoEXP program. However, as shown in Fig. S2, the lobe heights in the simulated form factors depend on the simulation box size, but locations of minima does not. Therefore, quality measure depending on lobe heights is also dependent on simulation box size. In order to define a quality measure which is independent on simulation box size, we measure the form factor quality based on the location of the first minima observed in experiments. As shown in Fig. 2, this minima correlates well with the thickness of a membrane. Although the correlation in second minima would be even stronger (Fig. S1), we have chosen to use the first minima because automatic definition of second minima is inaccurate in some experimental data. In practise, we first filter the fluctuations from the form factor data using Savitzky-Golay filter (window lenght 30 and polynomial order 1) and locate the first minima above $0.1 AA^2$ from both simulation and experimental data. The quality of a form factor is then defined as Eucledian distance between the minima in simulated and experimental form factors, $FF_q = |FF_{\min}^{\text{exp}}|$.

4.4 Analysing simulations in the NMRlipids databank

The README.yaml files contain all the essential information to perform arbitrary analyses of simulations in the NMRlipids databank, i.e., the permanent location of the original data and naming convention for all atoms and molecules in each system. In practise, the analyse codes contains a loop over all README.yaml files (i.e., simulations in the NMRlipids databank) which first downloads the raw simulation to a local computer and then uses the information about the atom and molecule naming conventions in README.yaml and mapping files to perform the desired analyses. For example, the code that calculates all C-H bond order parameters of all systems is available at https://github.com/NMRLipids/Databank/blob/main/Scripts/AnalyzeDatabank/calcOrderParameters.py and minimal example of a analysis code is available at https://github.com/NMRLipids/Databank/blob/main/Scripts/AnalyzeDatabank/template.ipynb.

While the order parameters, form factors, area per lipid and thickness are stored within the NMRlipids databank (https://github.com/NMRLipids/Databank/tree/main/Data/Simulations), further analyses can be conventiently stored in separate repositories with the same folder structure based on hash identities of trajectory and topology files. For example, results from further analyses performed here are stored in folders at https://github.com/NMRLipids/DataBankManuscript/tree/main/Data. Such organization of the data enables further upcycling of the analyzed data as similarly to the original NMRlipids databank repository.

For the permeation of water through membranes, the number of permeation events in each trajectory was first calculated using the code by Camilo et al.³⁹, available at https://github.com/crobertocamilo/MD-permeation. The permeation was then calculated from equation $P = r/2c_w$, where r is the rate per time and area, and c_w is the concentration of water in bulk with the value of 33.3679 (nm)⁻³?

References

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Author contributions statement

Must include all authors, identified by initials, for example: A.A. conceived the experiment(s), A.A. and B.A. conducted the experiment(s), C.A. and D.A. analysed the results. All authors reviewed the manuscript.

Additional information