

Modeling of Interaction of Ions with Ether- and Ester-linked phospholipids

by

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A thesis submitted in partial fulfillment
of the requirements for the degree of
Master of Science in Biology
with a concentration in Cell & Molecular Biology
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Date of Approval:
October 23rd, 2019

Keywords: Molecular Dynamics, Gouy-Chapman Theory

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Abstract

Phospholipids are present in all parts of cells and are used in many signalling and structural roles. As structural molecules they act as the main component of cellular membranes. Bilayer properties are heavily influenced by the structure of their component polar lipids, and different lipids are found in different organisms. A distinguishing feature of Archaeal plasma membranes is that their phospholipids contain ether-links, as opposed to bacterial and eukaryotic plasma membranes where phospholipids primarily contain ester-links. In our work we examine the effects of salt on bilayer structure in the case of both ester- and ether-linked lipid bilayers. We use molecular dynamics simulations and compare equilibrium properties of two model lipid bilayers in NaCl salt solution — POPC and its ether-linked analog that we refer to as HOPC. We make the following key observations. The headgroup region of HOPC “adsorbs” fewer ions compared to the headgroup region of POPC. Consistent with this, we note that the Debye screening length in the HOPC system is $\sim 10\%$ shorter than that in the POPC system. Herein, we introduce a protocol to identify the lipid-water interfacial boundary that reproduces the bulk salt distribution consistent with Gouy-Chapman theory. We also note that the HOPC bilayer has excess solvent in the headgroup region when compared to POPC, coinciding with a trough in the electrostatic potential. Waters in this region have longer autocorrelation times and smaller lateral diffusion rates compared to the corresponding region in the POPC bilayer, suggesting that the waters in HOPC are more strongly coordinated to the lipid headgroups. Furthermore, we note that it is this region of tightly coordinated waters in the HOPC system that has a lower density of Na^+ ions. Based on these observations we conclude that an ether-linked lipid bilayer has a lower binding affinity for Na^+ compared to an ester-linked lipid bilayer.

1 Introduction¹

1.1 Ether- and Ester-linked Phospholipids

Phospholipids are a diverse species of biological molecules used in many roles in life.[?] These molecules are amphipathic, with a headgroup that is hydrophilic attached to hydrophobic acyl chains by a glycerol backbone.^{?,?} These molecules, when above a critical concentration in aqueous solution, self assemble into various aggregate structures in order to reduce the exposure of the hydrophobic chains to polar solvent — these structures include micelles, vesicles, and bilayers along with others.[?] Phospholipids tend to form bilayers, where the hydrophilic headgroups face the solvent and the hydrophobic acyl chains face into the center of the structure.^{?,?} In living cells, lipid bilayers help to delimit the cell and the environment, acting as one of the major constituents of the cellular plasma membrane.[?] The plasma membrane consists of a mixture of many lipid species, as well as proteins and lipid-like molecules, and the mixture is tuned by the cell to adapt the structure and flexibility of the plasma membrane to different environments.^{?,?,?} Furthermore, the cellular membrane is the interface for exchange of nutrients with the environment, and also the interface for the transmission of signals to and from the outside of the cell. This exchange is done through passive diffusion, proteins, and other molecules embedded into the membrane, and through the formation of vesicles and buds out of the lipid bilayer.[?] The behavior and function of the plasma membrane is heavily dependent on the structure of the bilayer, which again can be influenced directly by the mixture of phospholipids constituting the membrane.^{?,?}

Phospholipids can be modified in a variety of ways to change their chemical construction, and the structure of the individual molecules manifest significant changes in the structure

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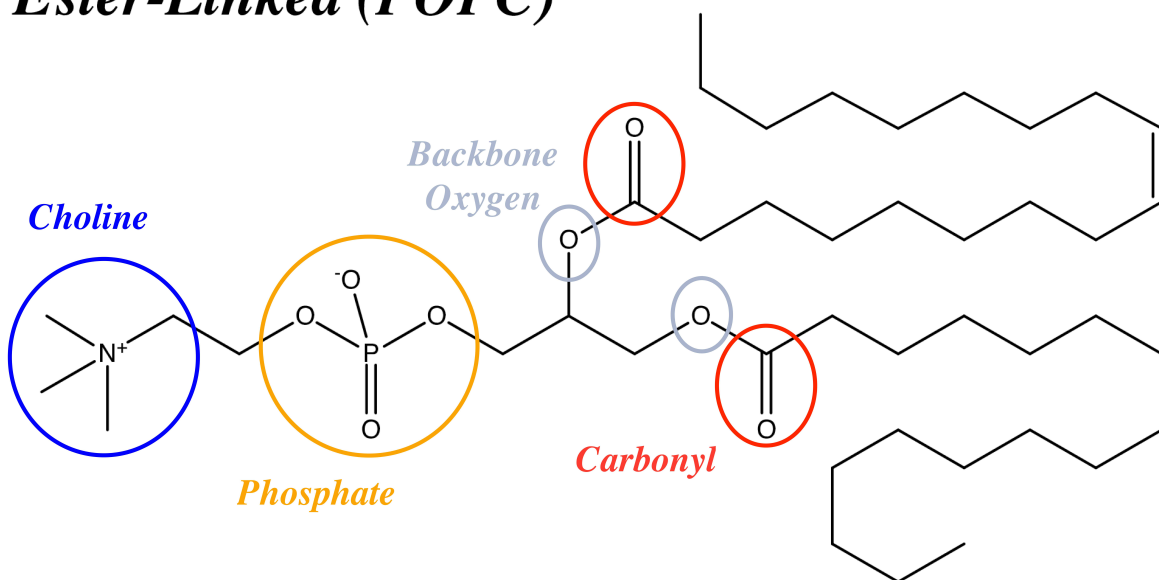
of aggregates, like lipid bilayers or cell membranes. Polar headgroups can be charged or zwitterionic.[?] The fatty acid chains can be saturated or unsaturated, and can have many lengths depending on the needs of the cell — differing the acyl chains influences the phase behavior of the bilayer, which can help cells to adapt to changes in temperature.^{?,?}

Phospholipids can also use several types of chemical links between the glycerol backbone and the fatty acid chains.^{?,?} Phospholipids containing ether-links represent one of the many major evolutionary divides — while they make up much of the phospholipids of archaeal membranes, they are present in only small fractions in eukarya and bacteria.^{?,?,?} These phospholipids differ from the more common ester-linked phospholipids of bacterial and eukaryotic membranes in that they lack fatty acid carbonyl groups (see figure ??).

The chemical difference between ester- and ether-linked lipids has been shown to influence bilayer properties including both structure and permeability.^{?,?,?,?,?} Experiments show that model bilayers composed exclusively of ether-linked lipids have smaller dipole potentials compared to bilayers composed of their ester-linked analogs;[?] a result also predicted by a molecular dynamics (MD) study done by our group.[?] The interfacial water in ether-linked lipid bilayers is also more structured compared to that in ester-linked lipid bilayers, as observed in NMR experiments in the form a larger quadrupolar splitting constants,[?] and as also reproduced in our simulations.[?] Two independent experimental studies also show that ether-linked lipid bilayers are less permeable to water compared to bilayers composed of their ester-linked analogs.^{?,?}

Nevertheless, all studies aimed at examining differences between ester- and ether-linked lipid bilayers have been carried out in pure water, except for the one recent study by Leonard *et al.* where an all-atom force field was parameterized for a different model ether-linked lipid.[?] In the work by Leonard *et al.*, the primary analysis given was for systems without salt, and did not treat the system with salt beyond structural measurements.[?] The questions of how ions bind differently to bilayers of ether- and ester-linked lipids as well as how they interact with the bilayers electrostatically are still open to investigation. The inclusion of ions in a

Ester-Linked (POPC)



Ether-Linked (HOPC)

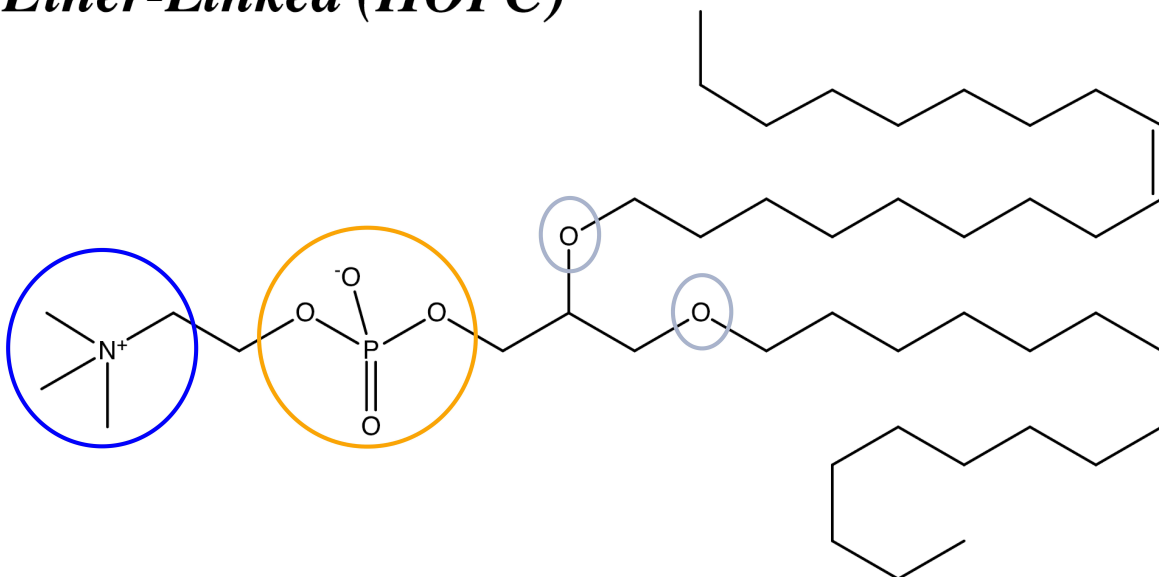


Figure 1.1: Chemical structures of representative phospholipids highlighting the difference between ether- and ester-links. POPC (1-palmitoyl-2-oleoyl-sn-glycero-3-phosphatidylcholine) is a diester lipid and HOPC (1-hexadecyl-2-(9-octadecenyl)-sn-glycero-3-phosphatidylcholine) is its diether analog that lacks fatty acid carbonyl groups.

bilayer system is an important step to achieving a more physiologically relevant system — biological membranes are always exposed to ions.[?] In biology, the affects of ions on bilayer structure cannot be ignored. As such, several studies show that salt significantly affects the properties of ester-linked lipid bilayers.^{?, ?, ?, ?, ?, ?} For example, in our previous works on the interactions of monovalent and divalent ions on zwitterionic ester-linked lipid bilayers^{?, ?} we found that monovalent cations insert into headgroup region. We reported reduction in area per lipid, increase in bilayer thickness, increased range of ordered solvent at the bilayer surface, and an increase in the electrostatic dipole potential of the bilayer. Despite the work done on these lipids in the past, the question of how ions interact with bilayers of ether-linked phospholipids compared to ester-linked remains a topic of interest.

1.2 Using Molecular Dynamics to Study Biological Membranes

In order to explore the question of how ions interact with bilayers of ether- and ester-linked lipids, we utilize Molecular Dynamics (MD) simulations.

Computational methods like molecular dynamics must strike a balance between computational cost and accuracy. The most expensive computations using *ab initio* methods with minimal approximations are the most accurate, but are only feasible for very small systems.[?] For biologically relevant systems, a classical approximation is able to reproduce most important phenomena.^{?, ?} MD makes this classical approximation, expressing particles as points in space, with masses centered at the nucleus.[?] Functional forms and functional parameters needed to calculate the hamiltonian of systems are defined in a force-field. Bonds and bond angles are modeled as harmonic potentials, and torsional parameters within molecules are modeled using periodic functions. Non-bonded interactions are modeled using simple pairwise-additive potentials and generally do not treat many-body interactions.^{?, ?} Van der Waals (VdW) dispersion and hard-shell repulsion are usually treated using a Lennard-Jones 6-12 potential, and short-range electrostatics are represented by a coulomb potential.^{?, ?} The potential terms are added together over the whole system of particles to get the overall potential function of the system, which is then used to evolve the system over time by integrating

Newton’s laws of motion.[?] An example of the total potential can be seen in equation ??.[?]

$$\begin{aligned}
V_{tot} = & \sum_{\text{bonds}} K_b(r - r_0)^2 \\
& + \sum_{\text{angles}} K_\theta(\theta - \theta_0)^2 \\
& + \sum_{\text{improper}} K_\Phi(\Phi - \Phi_0)^2 \\
& + \sum_{\text{torision}} K_\phi[1 - \cos(n\phi - \phi_0)] \\
& + \sum_{\text{Estatic}} \frac{q_i q_j}{r_{ij}} \\
& + \sum_{VdW} \frac{A_{ij}}{r_{ij}^{12}} - \frac{B_{ij}}{r_{ij}^6}
\end{aligned} \tag{1.1}$$

Most force-fields for molecular dynamics treat charges on particles in the system as an average picture of overall electronic behavior, as the timescale of electronic fluctuations is much smaller than the timescale of MD simulations; this means bonds are permanent, and average charge densities are fixed to particles.[?] Some force fields treat electronic polarizability as an explicit term in the potential function;^{?,?} however most MD force-fields do not include this while still giving a good reproduction of experimental results.^{?,?,?}

To ensure accurate reproduction of experimental results from complex systems, force-fields are developed and optimized to directly reproduce experimental data in simple model systems. To develop force-fields, systems of small molecules can be optimized to reproduce experimental heats and enthalpies of vaporization, molecular volumes, and densities of condensed phases.^{?,?,?,?} For systems of whole lipid bilayers, experimental methods such as deuterium NMR, small angle x-ray scattering, and small angle neutron scattering can be used to get ensemble average values for bilayer ordering and structure and are often used to validate lipid simulations.^{?,?,?,?,?,?,?} We can also look at electrostatic properties such as the bilayer electrostatic dipole potentials,^{?,?} or mechanical properties like bilayer compressibility or surface tension.^{?,?,?} For optimization of force-fields using small molecules, experimental

heats and enthalpies of vaporization, molecular volumes, and densities of condensed phases can be used.^{?, ?, ?, ?}

The earliest lipid simulations often used a united atom representation of the lipids, which simplified lipid chains by expressing carbon groups as beads rather than explicitly defining hydrogens.^{?, ?, ?} These simulations focused on validation with experimental values for bilayer structure, such as area per lipid and bilayer thickness that are taken from SAXS and SANS measurements. Later development of force-fields included calculating partial charges for different parts of the lipid molecules using *ab initio* quantum calculations.^{?, ?} Chiu *et al.* calculated charges in this way for the GROMOS 43-A1 force-field of DPPC in 1995.[?]

Berger *et al.* sought to further improve the densities given by the GROMOS force-field in 1997 by optimizing VdW parameters using molecular volumes and heats of vaporization in small hydrocarbon molecules to improve the acyl chains of DPPC,[?] work that was continued by Chiu *et al.*,^{?, ?} Later, further optimization was carried out by Chiu *et al.* in 2009 to improve torsional and VdW parameters using experimental heats of vaporization and densities, and validated the overall result by predicting small-angle x-ray scattering results for model lipid bilayers to create the GROMOS 43-A1S3 lipid force field that we use for this work, which gave excellent reproduction of SAXS form-factors for different lipids.[?] Parameters for the ether-links were added to the force-field by Kruczek *et al.* in 2017,[?] and the force-field was demonstrated to work well with ion parameters from Joung and Cheatham III[?] by reproducing ion binding constants in the same year.[?]

2 Interaction of salt with ether- and ester-linked phospholipid bilayers¹

In this work we seek to answer the general questions of how salt interacts with ether-linked lipid bilayers, and how ether-linked lipid bilayers differ from ester-linked lipid bilayers in the presence of salt. We addressed these questions by carrying out a molecular dynamics simulation of 1-hexadecyl-2-(9-octadecenyl)-sn-glycero-3-phosphatidylcholine lipid bilayer (HOPC) in NaCl salt. We also compared results of this simulation against our previous simulation of 1-palmitoyl-2-oleoyl-sn-glycero-3-phosphatidylcholine lipid bilayer (POPC) in NaCl.[?] Additionally, to gain insight into how salt modulates differences between ether- and ester-linked lipid bilayers, we also compared these simulations against those of ether- and ester-linked lipid bilayers in pure water.[?]

2.1 Methods

The trajectories for the POPC bilayer in 200 mM NaCl were taken from our previous work,[?] and were re-analyzed to address the specific goals of this project. The trajectories for the HOPC bilayer were generated as described below, following the same protocol we used for the POPC bilayer. Analysis was carried out using GROMACS^{?,?,?,?} utility tools, and in-house software developed using GROMACS API.

2.1.1 HOPC bilayer construction

To construct the HOPC bilayer, we first place 100 lipids on 10×10 nm grid to form a bilayer leaflet, and then reflect the leaflet to create the second leaflet of the bilayer. We then introduce 30,000 water molecules into the space outside the bilayer, which gives a 150:1 waters to lipid ratio. We then replace 216 randomly-selected waters with 108 Na^+ and 108

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Cl⁻ ions to set an initial NaCl concentration of 200 mM. Note that the high water to lipid ratio is necessary to get a good representation of bulk solvent, as the addition of salt increases the distance of solvent ordering from the bilayer surface.[?] We then remove bad contacts by energy minimizing using the steepest descent approach. Finally, we subject the system to one cycle of annealing. Heating is done under constant pressure to 500K, but in steps of 100K to avoid excess kinetic energy divergence. During heating, each step is simulated for 10 ps. Cooling is also performed under constant pressure, but in smaller decrements – cooling from 500K to 400K is done in 5 steps, and cooling from 400K to 300K in steps of 10 steps. Each cooling step is simulated for 50 ps. The total annealing time is 850 ps.

2.1.2 Simulation Details

MD simulation is performed using GROMACS version 5.1.5.^{?,?,?,?} Water is described using the SPC/E model,[?] and we use the united atom Gromos 43A1-S3 parameters for lipids and lipid-water cross terms developed by our group.^{?,?} These parameters have been verified in previous works without salt against experimental SAXS data to ensure accurate reproduction of bilayer structure.[?] Parameters for HOPC are derived from those for ether-linkages used for di-hexadecyl phosphatidylcholine developed for previous work without ions.[?] Ion-ion and ion-water interactions are described using parameters developed by Joung and Cheatham III,[?] and the ion-lipid cross terms are calculated explicitly using Lorentz–Berthelot rules.[?] These ion parameters were verified in previous work for binding constants as well as electrostatic potential, and we reported good reproduction of experimental results.[?] Temperature is held constant at 300K using the Nosé-Hoover thermostat,[?] and a coupling constant of 0.5 ps. Pressure is maintained at 1 bar using the Parrinello-Rahman semisotropic barostat,[?] and a pressure coupling constant of 1.5 ps. All bonds are constrained using the P-LINCS algorithm.[?] Neighbor-lists are updated every other time step, and integration is carried out using the leap-frog Verlet scheme. Long-range electrostatics beyond 16 Å are computed using the smooth particle mesh Ewald algorithm,[?] and Lennard-Jones interactions are calculated

with a cutoff of 16 Å. As before,[?] a continuous MD simulation is run for 0.5 μ s.

2.2 Results and Discussion

2.2.1 Bilayer Structure

We first compare mass densities of the different chemical components of HOPC against those of POPC (see figure ??).

We find that the headgroup region of HOPC has a higher density of solvent compared to POPC – this is visible as a distinct peak in the solvent density profile in the HOPC system. We verified that this additional peak does not result from accumulation of ions by comparing mass density profiles of only water molecules. This peak is similar to observations of a smaller peak made in our previous simulations in the absence of salt, and may be a direct result of the ether-linked lipids.[?]

Table ?? compares the structural properties of HOPC and POPC bilayers. Lipid volumes $V_l = V_c + V_{hg}$, where V_c and V_{hg} are the volumes of lipid chains (tails) and headgroups. These are computed using the approach given in Petrache et al.[?] In this approach, we first compute the number densities $n_i(z)$ of the various system components as a function of the bilayer normal (z). We then use these $n_i(z)$ to determine their partial molecular volumes, v_i , by optimizing the objective function,

$$\Omega(v_i) = \sum_{z_j}^{n_s} \left(1 - \sum_{i=1}^{N_{\text{groups}}} (n_i(z_j) v_i) \right)^2. \quad (2.1)$$

Here, $N_{\text{groups}} = 7$ are the different system components including methyls (v_{CH_3}), methines (v_{CH_2}), methylenes (v_{CH}), headgroup carbons, headgroup and backbone oxygens, headgroup phosphate and nitrogen, and water (v_{H_2O}). Using these partial molecular volumes, we compute $V_c = 2v_{CH_3} + 2v_{CH_1} + 28v_{CH_2}$ and $V_{hg} = 10v_{hgC} + 2v_{P\&N} + 8v_{hgO}$. We find that the headgroup volume of HOPC is 27.1 Å³ smaller than POPC, due likely to the absence of carbonyl groups. As expected for lipids with the same hydrocarbon chains, the chain volumes are similar in the two lipids.

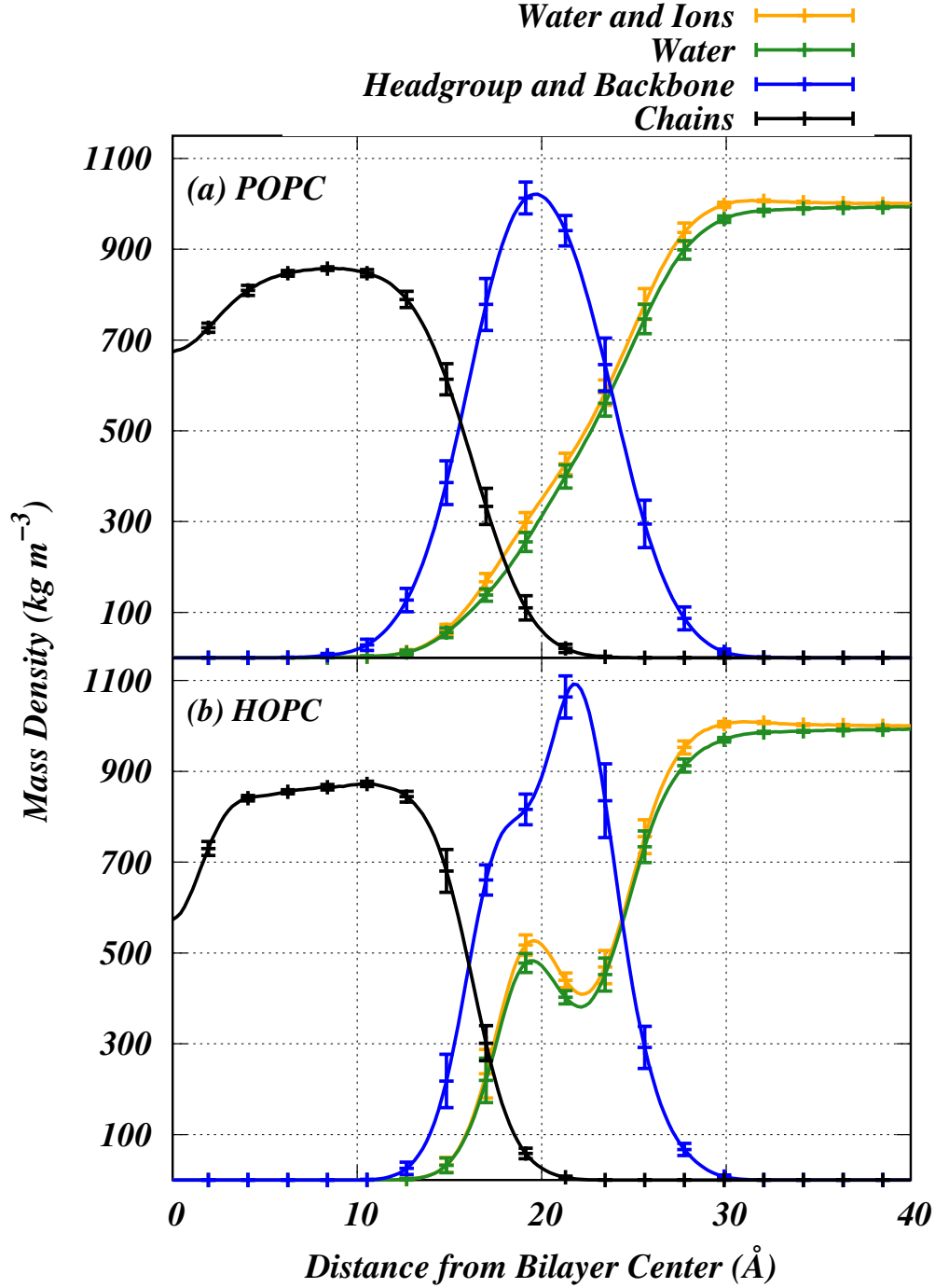


Figure 2.1: Comparison of component mass densities between the HOPC and POPC systems. Densities are computed by dividing the box dimension along the membrane normal into 2000 slices. For clarity of presentation, error bars are indicated for every 10th slice. Mass densities of solvent and ions, water, lipid headgroup and backbone, and lipid chains are shown. The plot for HOPC shows a peak in the water density inside the headgroup region that is not present in POPC, similar to what was seen in our previous work with DHPC.[?]

Table 2.1: Comparison of HOPC and POPC bilayer structure properties: V_l is the lipid volume; V_c and V_h are the partial volumes of lipids chains and headgroups, respectively; $2D_c$ and D_b are chain and bilayer thickness, respectively; $A_l = V_c/D_c$ is the area per lipid. N_w is the number of perturbed waters in the system, defined as those behind the *hydration boundary*. $\Delta\nu$ is the quadrupolar splitting constant of each system.

	POPC with NaCl	HOPC with NaCl
V_c (\AA^3)	898.4 ± 1.2	900.0 ± 1.0
V_h (\AA^3)	318.9 ± 0.9	290.2 ± 0.9
V_l (\AA^3)	1217.3 ± 0.5	1190.2 ± 0.9
$2D_c$ (\AA)	30.95 ± 0.32	32.13 ± 0.25
D_b (\AA)	46.22 ± 0.49	46.05 ± 0.34
A_l (\AA^2)	58.07 ± 0.63	56.03 ± 0.45
D_{hh} (\AA)	40.24 ± 0.75	42.62 ± 0.88
N_w	22.6	22.3
$\Delta\nu$ (Hz)	194.99	1756.08

We also use the number densities computed above ($n_i(z)$) to determine lipid chain thickness ($2D_c$) and bilayer thickness (D_b). These results are found experimentally from x-ray and neutron scattering.[?] In our simulations these parameters are found by first computing the probability of finding the i -th group in each slice of the simulation box, $P_i(z) = n_i(z)/\sum_i n_i(z)$. We define $2D_c$ as the distance between the Gibbs' surfaces on the interfacial probability densities of lipid chains. Each of the two surfaces is computed by identifying the point on the curve where the integrals of probability densities in the interfacial regions above and below the surface are equal. We compute D_b in a similarly, but from solvent probability densities. We find that both POPC and HOPC have similar membrane thicknesses (D_b); however a comparison of $2D_c$ indicates that the hydrophobic chains in HOPC are slightly thicker than those in POPC, reflecting tighter chain packing. This is also evident from comparison of electron densities (see figure ??).

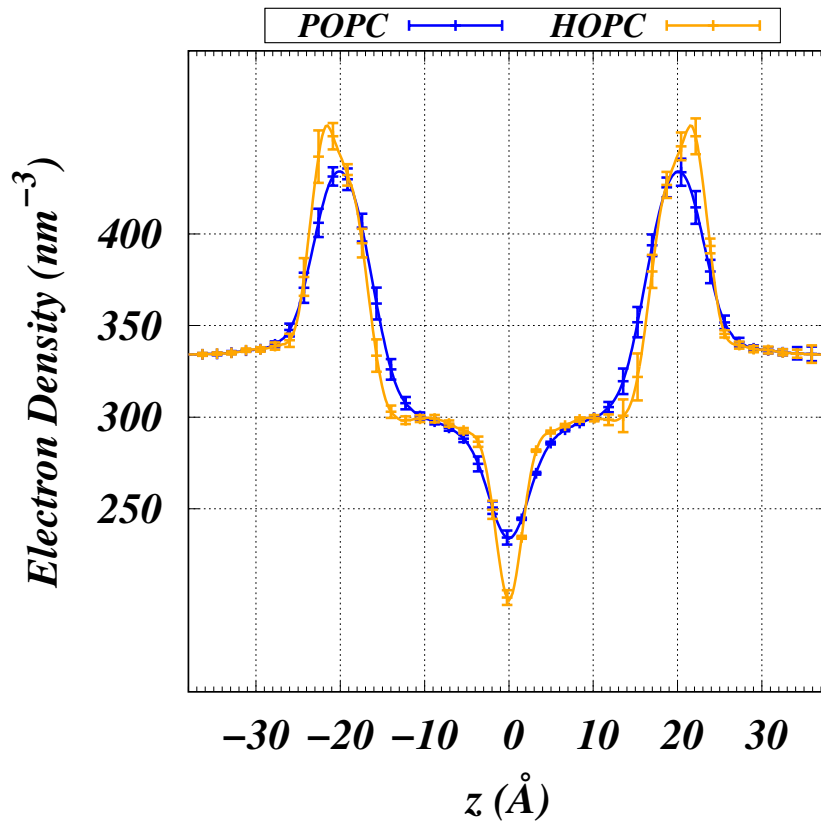


Figure 2.2: Electron Densities of Simulated Systems. Symmetrized electron density distribution of each simulated system as a function of z . We can see the distribution for HOPC is broader, with a longer peak-to-peak distance than in the POPC distribution. Furthermore, we see a deeper trough in the middle of the HOPC distribution. This plot was made by using the gromacs densities tool, dividing the system into 2000 slices. This data was calculated for every five nanoseconds, and averaged over the last 150ns of trajectories. Errorbars are shown for every 10th data point.

Electron density distributions are experimentally measured using x-ray scattering, by taking the reverse-fourier transform of the scattering form-factor.^{?,?,?} Since the experimental x-ray scattering is dependent on the model used to compute the reverse-transform, we often calculate the form-factor of our simulated bilayer to compare to experiment by transforming the electron density distribution.^{?,?,?} Since such data is not available for our current lipid system as far as we know, we cannot do this comparison here.

We also use the electron density distribution to calculate the peak-to-peak distance D_{hh} , which is used another measure of the bilayer thickness. This is roughly reflective of the distance between the electron-dense phosphate groups on either bilayer leaflet.

D_{hh} values for POPC and HOPC indicate that the ether-linked bilayer is $\sim 2\text{\AA}$ thicker on average than the ester-linked bilayer. This discrepancy between the values of D_b and D_{hh} is likely due to the irregular shape of the distribution of solvent in the headgroup of HOPC — the extra region of solvent accumulation may move the Gibbs’ surface used for finding D_b into the bilayer surface.

The difference in bilayer thickness can also be shown directly from comparison of chain order parameters (See figure ??). As lipid chains become more ordered on average, bilayer thickness increases.[?]

We determine chain order parameters from the chain order tensor $S_{\alpha\beta}$ defined as

$$S_{\alpha\beta} = \frac{1}{2} \left\langle 3 \cdot \cos(\theta_\alpha) \cdot \cos(\theta_\beta) - \delta_{\alpha\beta} \right\rangle, \quad (2.2)$$

where θ_α and θ_β are the angles made by the molecular axes with α and β as either x , y , or z , and $\delta_{\alpha\beta}$ is the Kroneker-delta function. For the saturated bonds[?]

$$-S_{CD}^{Sat} = \frac{2}{3} S_{xx} + \frac{1}{3} S_{yy}, \quad (2.3)$$

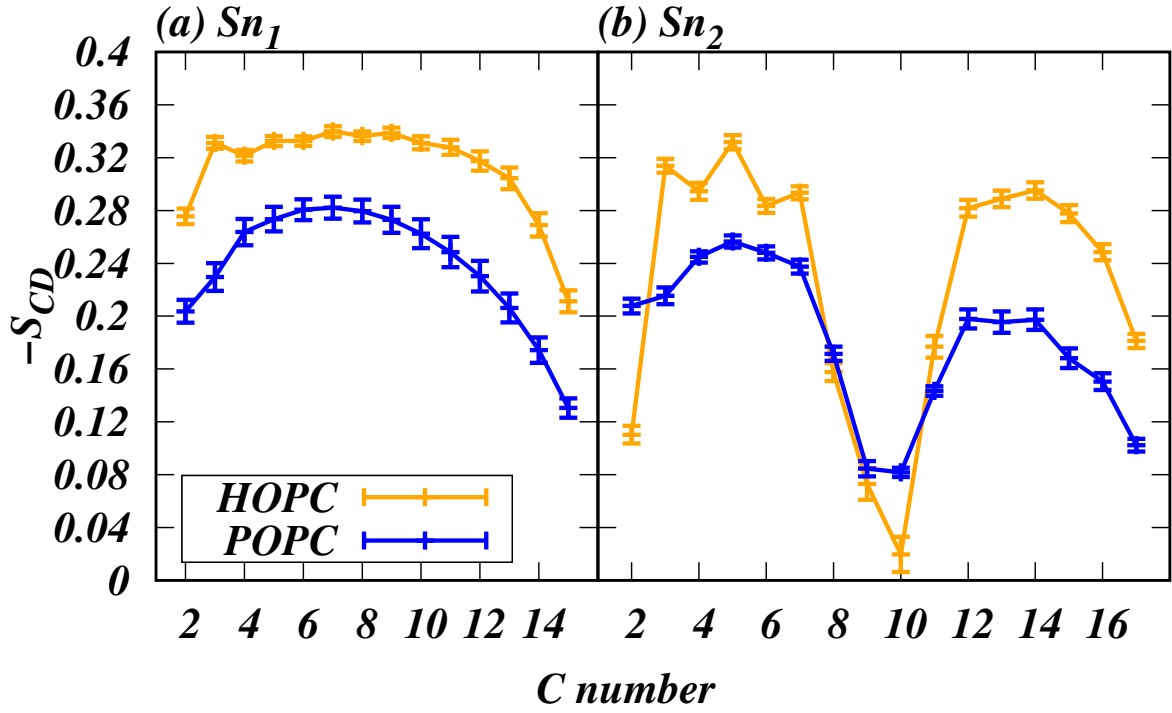


Figure 2.3: Order parameters of lipid chains. Sn_1 and Sn_2 refer to the two lipid tails. The carbon numbering starts from the polar carbon at the ester bond; however, we do not calculate an S_{CD} for the first carbon as this carbon would not have a hydrogen attached in an ester-linked lipid. This convention was maintained for the ether-linked lipid. The unsaturated bond is between carbons 9 and 10 on the Sn_2 chain. In both chains, HOPC shows higher ordering, except for the unsaturated carbons on the 9-octadecene chain. This ordering corresponds with the lower area per lipid of the HOPC bilayer, values of which can be seen in table ??.

and for unsaturated bonds?

$$-S_{CD}^{Unsat} = \frac{1}{4}S_{zz} + \frac{3}{4}S_{yy} \mp \frac{\sqrt{3}}{2}S_{yz} \quad (2.4)$$

We then use V_c and D_c to calculate areas per lipid, $A_l = V_c/D_c$. We find that the lateral surface area of HOPC is slightly smaller than that of the POPC bilayer.

2.2.2 Membrane-salt interactions

Ions prefer to associate or coordinate with specific sites in the headgroup regions of the lipid bilayers (See figure ??) — Na^+ ions near phosphates and Cl^- ions near cholines. Here we note that ion peaks are positioned similarly in both HOPC and POPC systems. Cl^- ions do not enter the bilayer headgroup region, while Na^+ ions move far into this region of the bilayer surface. However, the Na^+ densities trail off deeper into the POPC bilayer by ~ 5 Å (visible in the inset of figure ??). In fact, we find that more ions localize in the POPC bilayer — Averaged over the last 150 ns we see 74.45 ± 0.56 Na^+ ions bound to POPC, compared to 62.27 ± 2.00 Na^+ ions bound to HOPC. We consider an ion to be bound to the membrane when half or fewer of the inner-shell coordination partners of the ion are waters. The inner shell of Na^+ ions is defined using a cutoff of 3.15 Å based on radial distribution functions of Na^+ with the water oxygens.[?] The Cl^- ions do not lose waters and do not seem to adsorb on the bilayer, hence we have not analyzed the binding behavior of Cl^- in this work. Figure ?? shows the inner-shell coordination environment of Na^+ ions as a function of their positions in the bilayer. For this, we used a cutoff of 3.3 Å, based on the radial distribution functions of Na^+ with the various lipid oxygens (data not shown). When Na^+ ions are bound, they are primarily coordinated by phosphate oxygens — this holds for both HOPC and POPC. The main difference between POPC and HOPC is in how carbonyl oxygens, backbone oxygens, and water oxygens coordinate with ions. In POPC, ions are partially coordinated by both carbonyl oxygens and water, but not by backbone oxygens;

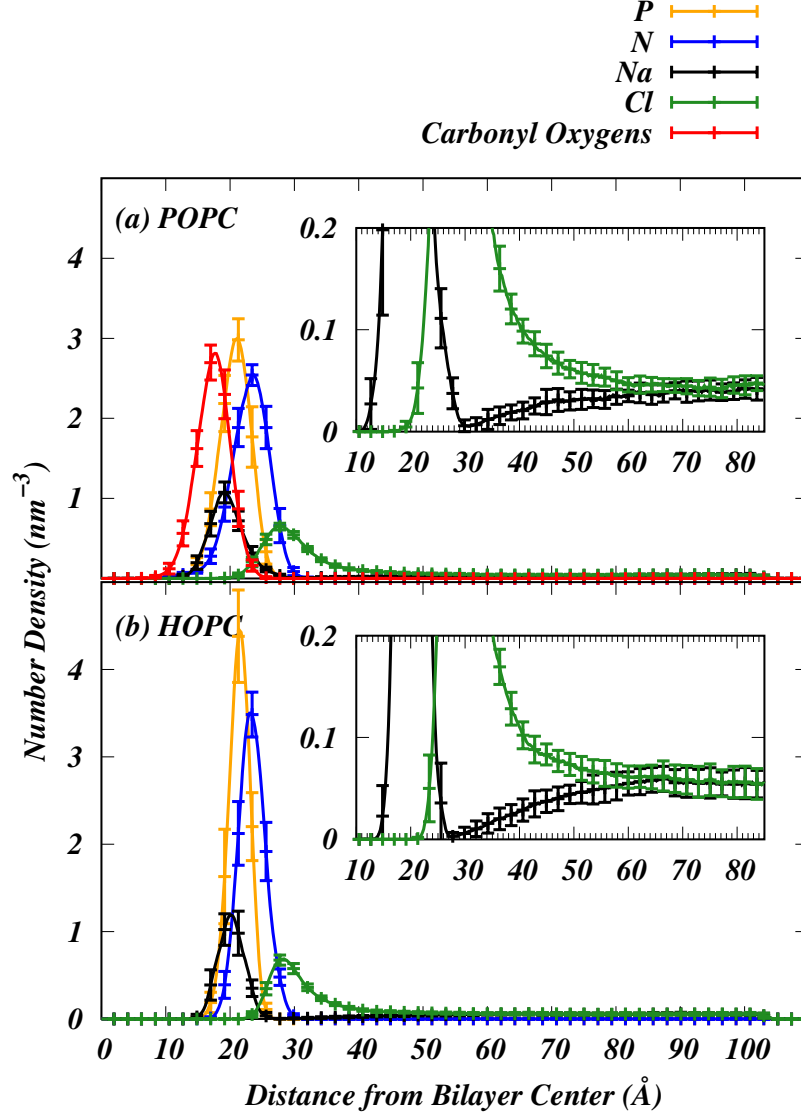


Figure 2.4: Comparison of headgroup component and ion number densities between the HOPC and POPC systems. The inset zooms in on the ordinate axis to visualize the extent of ion adsorption into bilayer as well as ion densities in the bulk. Densities are computed by dividing the box dimension along the membrane normal into 2000 slices. Error bars are indicated for every 10th slice. We note that sodium ions have a broader distribution in POPC than in HOPC. The peak of the Na^+ distribution in POPC is also closer to the center of the bilayer than in HOPC. This may be due to the interaction of ions with the carbonyl oxygens (shown in red) that are present in POPC but absent in HOPC. In both bilayers, the chloride ions gather near the positively charged choline trimethylammonium, shown in blue. The inset shows the distribution of ions in the transitional region from the bilayer surface to bulk solvent. As one looks further out from the bilayer, the density of Na^+ and Cl^- equalize, as is expected in bulk solvent.

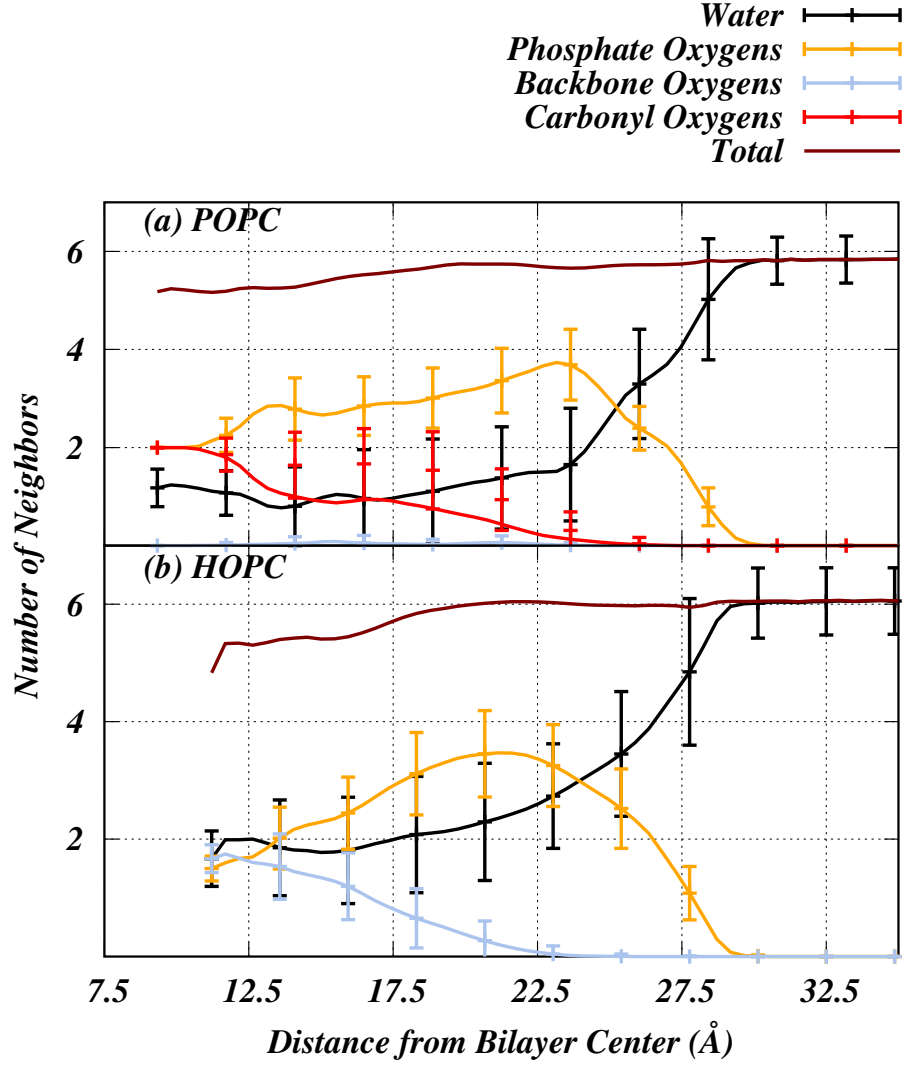


Figure 2.5: First shell coordination environment of Na^+ ions. The cutoff distance for the first shell is taken as 3.3 ångströms, based on the radial distribution function of Na^+ with the various lipid oxygens. This chart was made by dividing the box into 400 slices, and then symmetrizing around the bilayer center. Data is averaged over the last 100ns of simulation trajectory. Errorbars are shown for every tenth data point. Note that the coordination number of Na^+ drops from 6 to 5 as it interacts deeper in both bilayers, and this drop occurs sooner in the HOPC system than in POPC. No values are shown for at distances less than 7.5 Å from the bilayer center, due to the lack of ions beyond in this region.

however, in HOPC backbone oxygens do partially coordinate Na^+ . This may contribute to the deeper penetration of Na^+ into the POPC bilayer, and the greater number of ions adsorbed onto the POPC bilayer surface when compared to HOPC.

2.2.3 Water structure and dynamics

As we noted above in Figure ??, the headgroup region of HOPC has a higher density of water than POPC. We have also noted previously that in general salts tend to extend the region of structured water near the bilayer surface.[?] To gain further insight into how the absence of carbonyl groups in ether lipid bilayers increases water density, we systematically characterize the structure and dynamics of water.

Figure ?? compares the integrated radial distribution of water hydrogens around lipid oxygens. We note that there is very little water directly coordinating the backbone oxygens in the POPC bilayer ($\sim 1/3$ water per lipid). Compared to backbone oxygens, more water directly coordinates the ester carbonyl oxygens and the phosphate oxygens in POPC. We see a larger number of waters coordinating with the backbone oxygens in HOPC than in POPC, nearly 2 waters per lipid. Therefore, we attribute the higher density of water in HOPC to coordination with backbone oxygens.

Note that in POPC, carbonyl oxygens compete with phosphates for water coordination. Consequently, we expect waters to have a greater average ordering in the HOPC headgroup. We had observed this greater ordering in our previous comparative study of diether- and diester-lipid bilayers.[?]

In order to further explore solvent ordering by the bilayer surface, we compute order parameters of water O-H bonds. The first order parameter (P_1) is calculated by time-averaging the cosine of the angle θ that the OH bond makes with the bilayer normal, that is $P_1 = \langle \cos(\theta) \rangle$. Consequently, a positive value represents an orientation away the bilayer. The second parameter, which is defined as the second Legendre polynomial of $\cos(\theta)$, that is, $P_2 = \langle (3 \cdot \cos^2(\theta) - 1) \rangle / 2$. We calculate both P_1 and P_2 as a function of position along

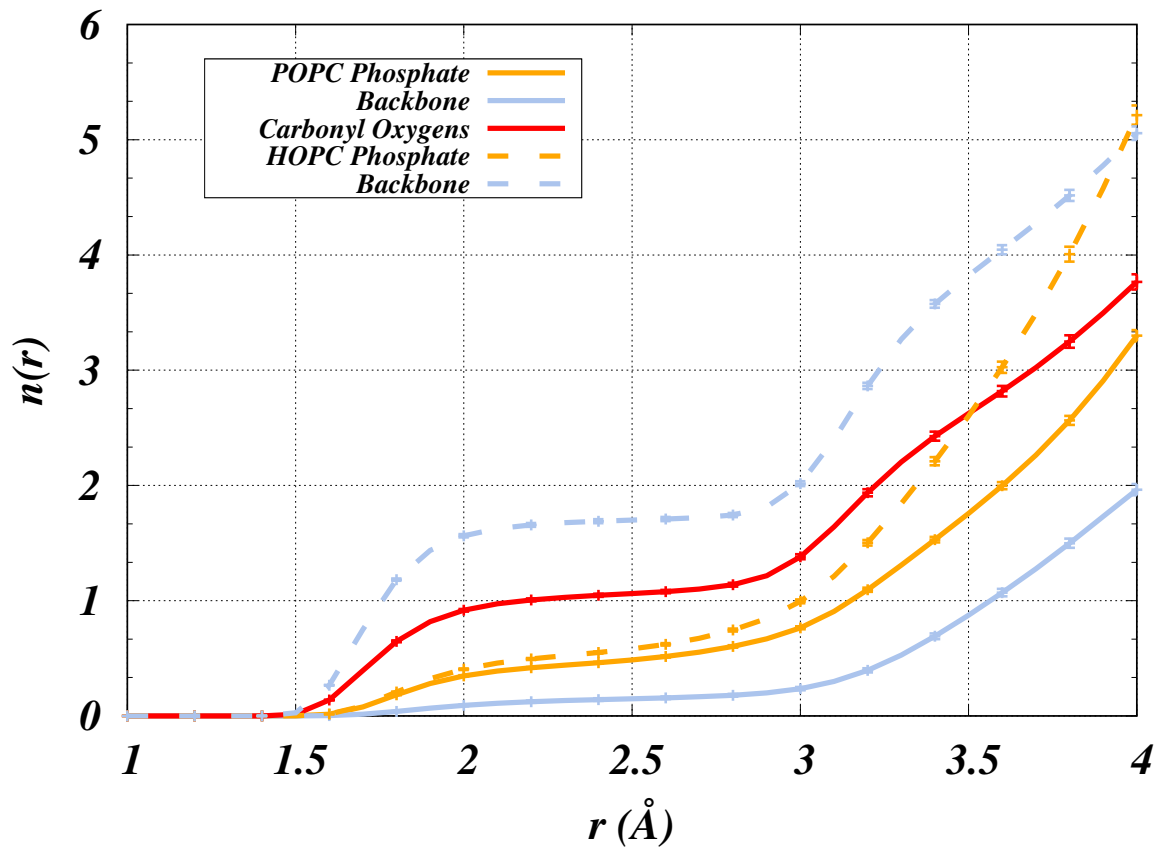


Figure 2.6: Cumulative radial distribution functions of water hydrogens around various lipid oxygens. The distributions for POPC are shown in solid lines, while HOPC is shown with dotted lines. The oxygens are grouped into categories as denoted in figure ??.

the bilayer normal, and they are shown in figure ??.

We find that the water in the headgroup region is clearly more ordered in the HOPC bilayer. Figure ??b shows the second order parameter. We note that the induced perturbations in the water layer are more prominent in HOPC than that of POPC. At the same time, however, we find that the numbers of water that are ordered in the two systems are similar. We count the average number of non-bulk (perturbed) waters as waters within the non-zero region of the second order parameter. We find this boundary by fitting an exponential function to the region starting at the peak in the first order parameter, and taking the inverse of the length-scale of the fitted function as the position of the surface. The region beyond this point is considered bulk water. We will refer to this surface as the *hydration boundary* for the remainder of the paper. For the purpose of counting the number of waters in the surface, we truncated the box at this boundary, to a length of 35 ångströms. Numbers of perturbed waters per lipid for each system can be seen in table ?? line 8.

We also use these order parameters to compute the quadrupolar splitting of water as measured in NMR experiments. It is essentially a weighted average of the second order parameter, similar to the one derived by Kruczek *et al.*:[?]

$$\Delta\nu = \frac{3}{4}\chi\frac{1}{N_w}\sum_{z_i=0}^{z_0}n_w(z_i)P_2(z_i), \quad (2.5)$$

where the number of perturbed waters in the system is N_w , n_w is the number of waters in each slice of the box, and $P_2(z_i)$ is the value of the second order parameter in the i^{th} slice. We take the quadrupolar splitting constant of water $\chi = 220\text{KHz}$ from Åman *et al.*[?] The summation over slices is carried out from the bilayer center ($z = 0$) to the end of the box. We report that the quadrupolar splitting of water in the POPC system is 194.99 Hz, and that in the HOPC system is 1756.08 Hz (also shown in table ??, line 9). These values reflect a similar difference to what we observed in our earlier comparative study of ether- and ester-linked lipid bilayers in pure water.[?]

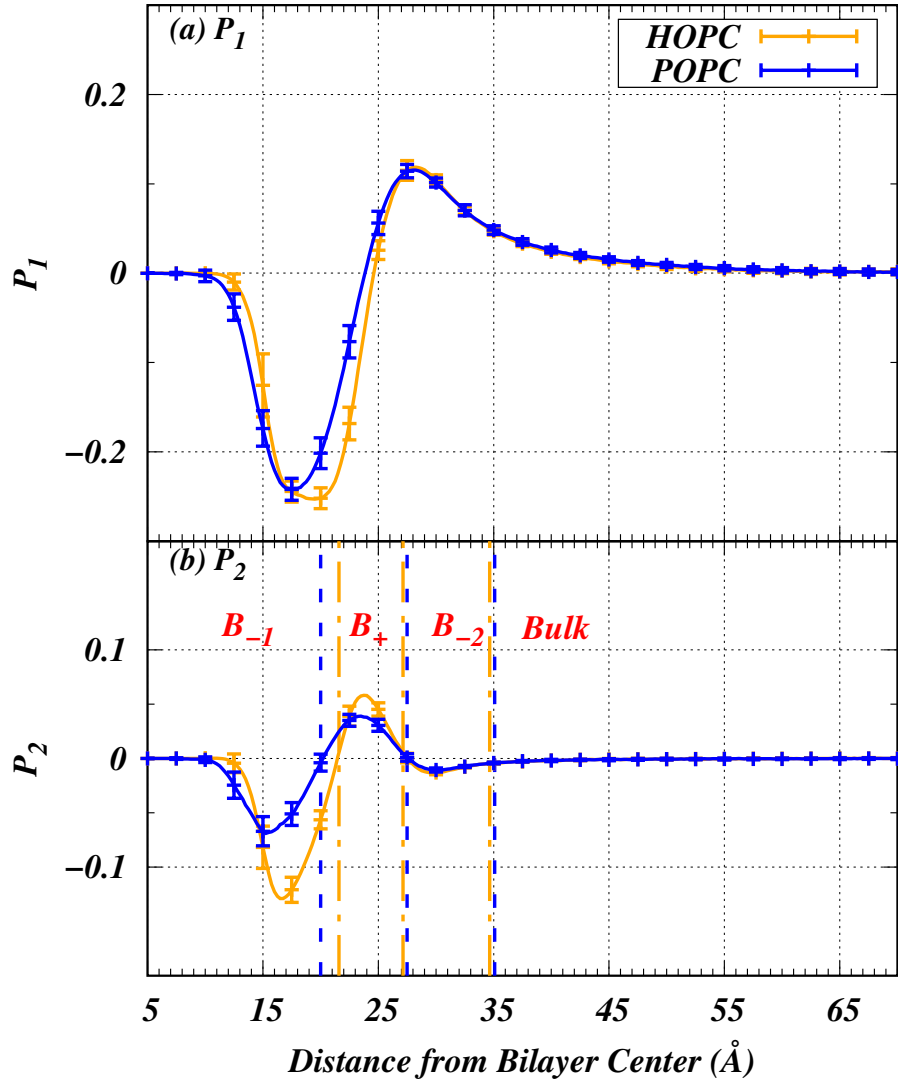


Figure 2.7: The first and second order parameters, P_1 and P_2 , of water O-H bonds. Each parameter is calculated by first dividing the simulation box into 2000 slices, and then averaging the data separately in each slice. The box is symmetrized around the bilayer center, and thus only half the box is shown. Standard deviations are calculated by dividing the trajectory into over 5ns blocks. The dashed vertical lines in the P_2 plot are demarkations for the different spatial regions, B_{-1} , B_+ , B_{-2} , and B_{bulk} , as discussed in the associated text. P_1 is the average of the first legendre polynomial of the cosine of the angle between the bilayer normal and the O-H bond vector of each water molecule per box slice. P_2 is the average of the second legendre polynomial. Looking to P_1 , both systems exhibit similar magnitudes of ordering. P_2 shows a more complicated distribution, with larger magnitudes of ordering in all regions in the HOPC system.

Next we compute the lateral diffusion coefficient of water. To gain detailed insight, we compute it separately for four different regions along the bilayer normal. We define these regions using the $P_2(z)$ profile: B_{-1} is the region closest to the bilayer center where $P_2(z)$ is negative; B_+ is the region where $P_2(z)$ is positive; B_{-2} is the region beyond the B_+ boundary where $P_2(z)$ is negative; and the remaining portion beyond B_{-2} we refer to as bulk. These regions are also labeled in figure ???. The diffusion coefficients for waters in these regions are included in table ??. They are calculated using Einstein’s relationship, and a protocol detailed in our previous study.⁷ Taking note of the overall pattern, the lateral diffusion decreases progressively as one moves deeper into the bilayer. Additionally, in the innermost regions B_{-1} and B_+ water diffuses more slowly in HOPC compared to POPC. This result is similar to what we observed previously⁷ in simulations without salt.

To characterize the orientational dynamics of waters, we compute autocorrelations of the O-H bonds,

$$C^{O-H}(t) = \left\langle \vec{v}_{O-H}(0) \cdot \vec{v}_{O-H}(t) \right\rangle. \quad (2.6)$$

We calculate expectation values separately for all the 2 Å slices along the bilayer normal by tracking individual water molecules for 500ps, and also averaging over all waters in the slice. We also weight each water by the fraction of duration it spends in each slice. This is done in 5 ns chunks, and then averaged over the trajectories. We then model these autocorrelations as a sum of three exponential terms,

$$C^{O-H}(t) = A_1 e^{-t/\tau_1} + A_2 e^{-t/\tau_2} + (1 - A_1 - A_2) e^{-t/\tau_3}, \quad (2.7)$$

where A_i are positive, and τ_i are correlation times. Fitting is carried out using the Marquardt-Levenberg least-squares fitting method. The correlation times are plotted in figure ??.

We find that τ_1 shows a profile similar to what we observed for ether- and ester-lipids simulated in pure water.⁷ Additionally, we find that the correlation times τ_1 and τ_2 in the HOPC bilayer are all longer than in POPC. The maximum correlation times of 1838 ps in

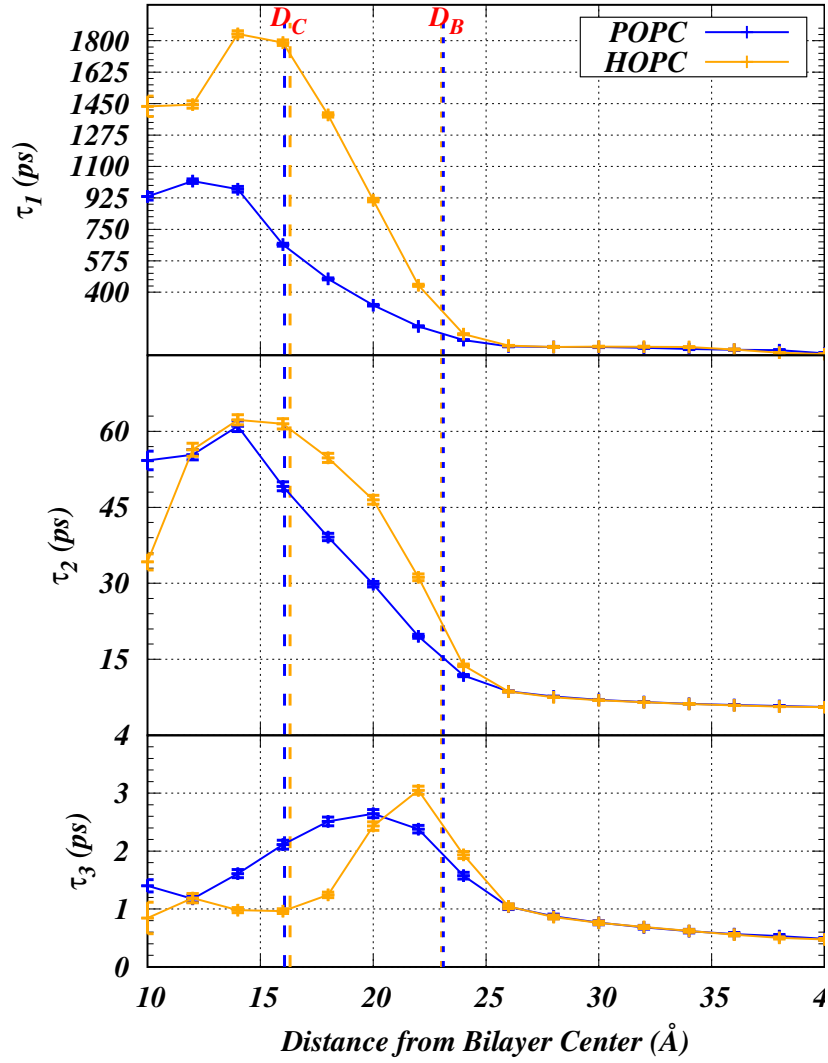


Figure 2.8: Orientational autocorrelation times of O-H bonds in water, as determined by modeling their autocorrelations using three-exponential fits. No values are shown for at distances less than 12 \AA from the bilayer center, as there are no waters in this region. The dashed vertical lines indicate chain boundaries D_C and bilayer thickness $D_b/2$, colored to indicate the lipid system.

Table 2.2: Diffusion coefficients of water measured in nm^2/s in different regions along the bilayer normal (see figure ?? and related text for the definitions of the different spatial regions. The two innermost-bilayer regions B_{-1} and B_{+} in HOPC contain waters that have a greatly lowered diffusion coefficient. In the regions B_{-2} and bulk, the bilayer composition does not seem to change the diffusion behavior. These values were calculated using the method outlined in previous work by our lab[?] The various regions are illustrated on figure ??.

	B_{-1} (nm^2/s)	B_{+} (nm^2/s)	B_{-2} (nm^2/s)	Bulk (nm^2/s)
HOPC	0.85 ± 0.59	1.783 ± 0.94	16.1 ± 5.24	26.67 ± 0.65
POPC	1.075 ± 0.62	3.078 ± 0.74	16.27 ± 5.19	27.30 ± 0.66

HOPC and 1019 ps in POPC are characteristic of rotationally immobile waters inside the bilayer surface and also of crystallographically-resolved waters in protein-protein interfaces.[?] Note that τ_3 , which is of the order of picoseconds, is close to the sampling time of our trajectories. Thus, we regard it simply as a fitting parameter.

Overall, we find that the headgroup water in HOPC is significantly more organized and also diffuses more slowly as compared to the headgroup water in POPC. This is similar to what we had observed[?] when we compared another diether lipid bilayer against its chemically analogous diester-lipid bilayer, but in the absence of NaCl. This leads us to conclude that in ether-linked lipid bilayers water forms a rigid and immobile layer within the headgroup region of the bilayer, and exposure to a moderate concentration of salt does not disrupt this layer.

2.2.4 Bilayer electrostatics

The dipole potential of a bilayer is the potential difference between the inside of the bilayer and bulk water. In our previous comparative study of ether- and ester-linked bilayers in pure water[?] we had found that the dipole potential of the ether-linked lipid bilayer was 56 ± 5 mV smaller than that of the ester-linked lipid bilayer, and this finding was consistent with experiment.[?] We also know that membrane association with salt can modify its dipole potential, with increases of around 200mV from that of a bilayer without salt reported

previously.^{?,?,?}

Figure ?? compares the electrostatic potential in HOPC and POPC as a function of distance along the bilayer normal. These are calculated from an integration of the one dimensional Poisson’s equation,

$$\phi(z) = -\frac{1}{\epsilon_0} \int_0^z \int_0^{z'} \rho(z) dz dz' + C_1 z + C_2, \quad (2.8)$$

where $\rho(z)$ is the charge density along the bilayer normal, and ϵ_0 is the vacuum permittivity. Note that we integrate this equation using two boundary conditions — the electric field in bulk water is zero, yielding $C_2 = 0$, and the potential at the simulation box boundary is also zero, which makes $C_1 = 0$.

The first observation we make is that the dipole potentials of both HOPC and POPC are similar; a behavior distinct from that observed in simulations in pure water. Nevertheless, the overall electrostatic potential profiles are similar to those observed in pure water. Firstly, a characteristic peak near the phosphate region still exists, and this peak is higher for the diether lipid bilayer. This could potentially serve as a higher permeation barrier in the diether-linked lipid system. This interpretation is consistent with our observation above that there are fewer membrane-associated ions in HOPC compared to POPC. Secondly, the diether-linked lipid system had a trough next to the peak, which is absent in the diester-linked lipid bilayer, and may serve as a trap for solvent molecules. In fact, the spatial location of the trough corresponds to the high water density region in the HOPC bilayer.

2.2.5 Salt distribution at the bilayer-solvent interface

The behavior of salt near bilayer surfaces is often modeled using Gouy-Chapman theory; a mean-field approximation that has its roots in Poisson-Boltzman theory.^{?,?} This theory also forms the basis for electrophoretic mobility studies.^{?,?} In this theory, water is modeled as a dielectric continuum in which ion particles behave as an uncorrelated gas. This means that the number density distribution of ions in the dielectric continuum only depends on the

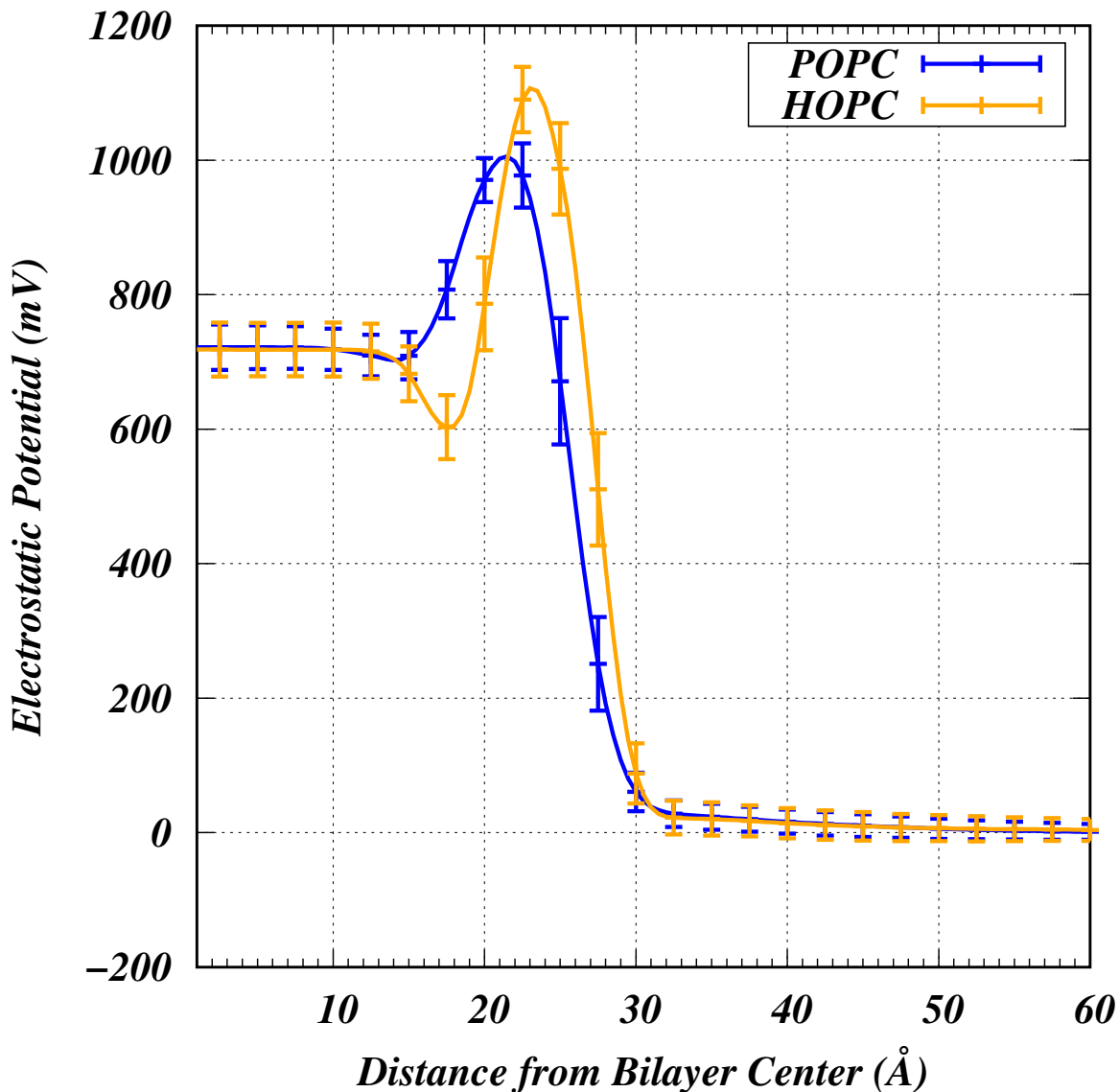


Figure 2.9: Comparison of electrostatic potentials between the HOPC and POPC systems. These distributions are calculated by integrating the charge density of our systems twice, assuming the potential goes to zero at the box edge and that the electric field of bulk solvent is zero. Looking to the region from the bilayer center to around 10 Å, one can see the bilayer dipole potential. We see very similar values for this in both systems. We also see a large peak in the HOPC system, and a large trough that is not present in POPC. The larger peak at the bilayer surface may provide a barrier to solvent and dissolved ions, and the trough may be a trap that causes the buildup of solvent in the bilayer headgroup region.

temperature and the electrostatic potential applied to the system.

So far, due to limitations from system size and ionic force field parameters, we have refrained from comparing or validating our simulations at the atomistic level against this theory. Here, we show convergence between these atomistic simulations and the Poisson-Boltzman theory.

Within Poisson-Boltzmann theory, the number density of ions near the bilayer surface is described as

$$\rho(z) = \rho_0 \exp(-\bar{z}e\beta\psi(z)), \quad (2.9)$$

where ρ_0 is the ion density in the bulk, \bar{z} is the valency of the ion, $\beta = (k_bT)^{-1}$, e is the charge on an electron, and $\psi(z)$ is the electrostatic potential. We define the interfaces as the *hydration boundaries* (see subsection ?? ‘Water structure and dynamics’). The lengths of the solvent occupied regions, D , in each of the two systems are listed in table ?. We set the center of this solvent occupied region as $z = 0$, which then implies that the interfaces are essentially at $z = \pm D/2$ nm, as illustrated in figure ?. We model $\psi(z)$ as a sum of two Debye-Huckel potentials,[?] each a reflection of the other around the center of the solvent occupied region:

$$\psi_1(z) = \psi_s \exp\left(-K\left(z + \frac{D}{2}\right)\right) \quad (2.10)$$

$$\psi_2(z) = \psi_s \exp\left(K\left(z - \frac{D}{2}\right)\right) \quad (2.11)$$

$$\psi(z) = \psi_1(z) + \psi_2(z) - (\psi_1(0) + \psi_2(0)) \quad (2.12)$$

Here $\psi_s = \sigma/\epsilon_0\epsilon K$ is the electrostatic potential at the bilayer surface, where σ is the surface charge density of the bilayer leaflet.[?] K is the inverse Debye length,

$$K = \sqrt{\sum_i \rho_{0,i} \bar{z}_i^2 \frac{e^2}{\epsilon_0 \epsilon k_b T}}, \quad (2.13)$$

where the sum of $\rho_{0,i}$ ’s is over all the included ions in our system. Note that in Equ. ??

Table 2.3: Parameters in Gouy-Chapman theory. K was calculated using equation ???. This was done using the value for ρ_0 for each system, found by taking the average number density of Cl^- ions within the solvent-occupied region of the box. The charge density σ was calculated by integrating the charge of the ions from the bilayer center to the surface at 35 Å from the bilayer center for each system, shown to be the *hydration boundary* of both bilayers. D gives the length of the solvent-occupied region of each simulation box.

	$K(\text{nm}^{-1})$	$D \text{ nm}$	$\rho_0 (\text{nm}^{-3})$	$\sigma (e \text{ nm}^{-2})$
POPC	0.98 ± 0.021	13.167 ± 0.0071	0.043 ± 0.0018	0.13
HOPC	1.09 ± 0.037	13.752 ± 0.0068	0.060 ± 0.0041	0.128

we subtract out the value of the potential at the center of the solvent occupied region from $\psi(z)$. Note also that the form of ψ_s is valid only for small surface potentials, generally those smaller than 20 millivolts.[?] Our systems have a relatively small surface charges (See table ??), which will result in small surface potentials (See table ??). The surface charges are calculated by integrating charge densities of ions contained behind the surface. We chose to do this because all other parts of the simulated system are uncharged, and the surface is entirely charged by the adsorbed ions.

Figure ?? compares the potential and ion distribution obtained from theory against those estimated directly from simulations.

We note a very close match between the two. Additionally, we find that HOPC shows a slightly larger K than POPC, indicating a $\sim 10\%$ shorter screening length for the system. This means that a bilayer of HOPC is better shielded from environmental electric fields. The shorter screening length in the HOPC is also reflective of the smaller number of ions that adsorb onto the surface of the bilayer, as this directly results in a larger number density in bulk solvent due to the fixed number of ions in our system.

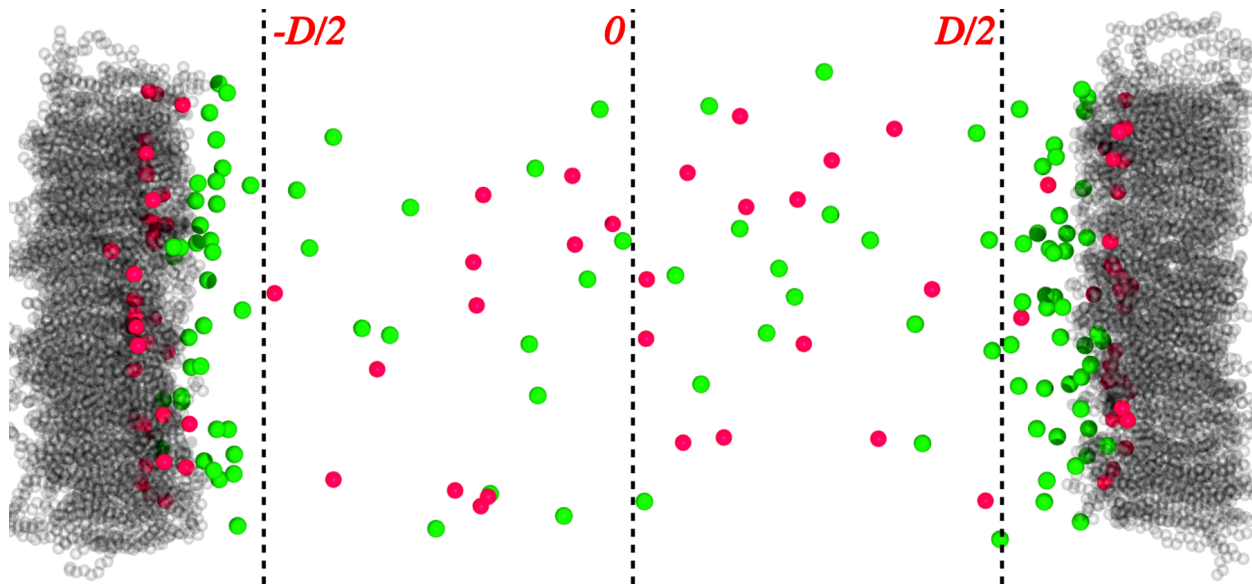


Figure 2.10: Illustration of bulk ion distribution, with surfaces used in Gouy-Chapman theory calculations. For the purpose of illustration, solvent has been hidden from this image. Cl^- ions have been colored in green, and Na^+ ions are shown in red. Surfaces set at the *hydration boundary* of each bilayer leaflet are shown with dotted lines. The center of the solvent occupied region of the box is set at zero.

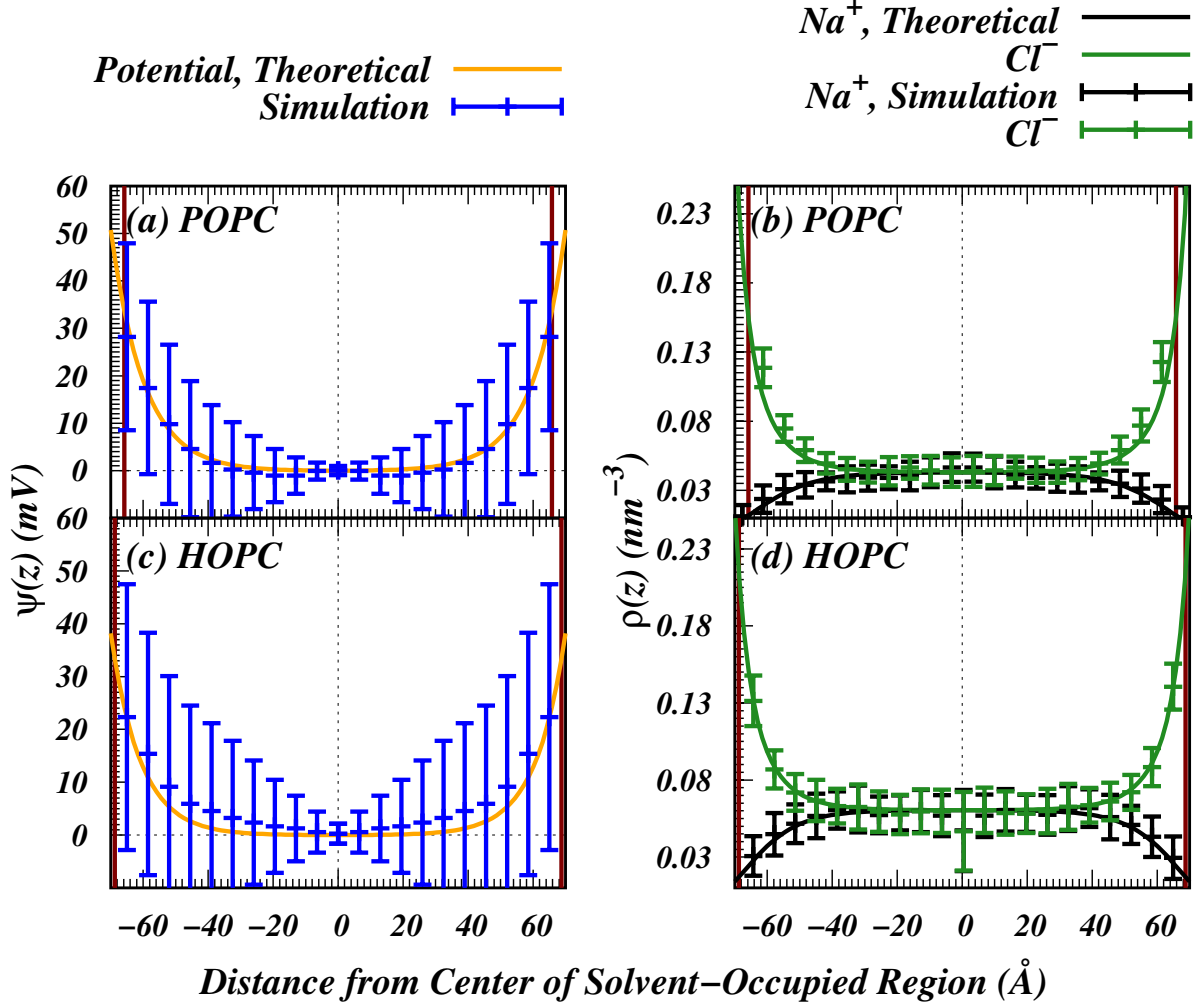


Figure 2.11: Comparison of Gouy-Chapman theory predictions and simulation results. Theoretical distributions are given by the solid lines, and data from our simulations are shown as points with error bars. (a) and (c) show the electrostatic potentials of each system. For clarity, we show error-bars for the simulation results for every 13th point. The Debye-Huckle potentials shown follow the form given in equation ?? . (b) and (d) show the number density distribution of ions in each system. Error-bars for the simulation results are shown for every 30th point. Note that both systems are translated to center around the solvent-occupied region of the box, giving an interspace region of ~ 13 nm in each system. Parameters used in Gouy-Chapman theory are in table ??

3 Conclusion

Phospholipids are an important part of all living things, acting as the major constituent of cellular plasma membranes. The great variety of phospholipid species allows organisms another parameter to fine-tune the behavior and structure of their membranes in order to adapt to a wide range of different environments.[?] This tunability makes lipid bilayers interesting to study, as small changes in lipid structure can result in large changes in the overall bilayer structure.[?] Classical molecular dynamics simulations are well suited to studying lipid bilayers,[?] and have been used extensively to study these systems over the last 25 years.

In our work, we have sought to model the interactions between dissolved ions and model ether- and ester-linked lipid bilayers. We found that ether-linked lipid bilayers show a distinct peak in solvent density that is not disrupted by the inclusion of a moderate concentration of salt. This region shows significantly reduced lateral diffusion, and longer autocorrelation times than the same region in an analogous ester-linked lipid bilayer. Furthermore, we find that the characteristic lower dipole potential of an ether-linked bilayer reported in previous work[?] is increased to match that of the ester-linked bilayer in the presence of salt. We also find that, with our new protocol for determining the interface boundary of the lipid bilayer, our simulations model ion distributions and electrostatic potentials from Gouy-Chapman theory well. The shorter screening length in the ether-linked system suggests better shielding from electric fields far from the bilayer compared to the ester-linked system. This is a result of the ether-linked lipid bilayer’s lower affinity for ions than the ester-linked analog.

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