Lipid Simulations: A Perspective on Lipids in Action

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In this article, we provide an overview of lipid simulations, describing how a computer can be used as a laboratory for lipid research. We briefly discuss the methodology of lipid simulations followed by a number of topical applications that show the benefit of computer modeling for complementing experiments. In particular, we show examples of cases in which simulations have made predictions of novel phenomena that have later been confirmed by experimental studies. Overall, the applications discussed in this article focus on the most recent state of the art and aim to provide a perspective of where the field of lipid simulations stands at the moment.

Lipids are very diverse in their structures and functions (Sackmann 1995; Mouritsen 2005; van Meer 2005). They are a crucial component of numerous biological entities such as membranes, lipoprotein particles, and lipid droplets, and they are involved in numerous cellular functions related to, for example, signaling and energy storage. Importantly, as lipids also compartmentalize biological membranes by creating membrane domains with different physical properties, lipids also affect or even govern membrane proteins and their functionality (McIntosh and Simon 2006; Lingwood and Simons 2010).

Although experiments are the cornerstone of lipid research, they are limited in resolution, permitting one to unravel biological phenomena only to a limited extent. Especially difficult to deal with are molecular scales with an objective to probe phenomena in the nanometer regime over timescales less than a microsecond. Molecular simulations, on the other hand, have no such limits with regard to resolution. Validated simulation models can be used to consider all sorts of phenomena, ranging from selectivity of ion channels to interactions of lipids with membrane proteins, and further to nonequilibrium lipid trafficking and domain as well as pore formation (Bjelkmar 2009; Bucher et al. 2010; Fan et al. 2010a,b).

The first simulations of lipid systems were performed in the early 1980s (Kox et al. 1980; van der Ploeg and Berendsen 1982, 1983). Starting from those times when solvent-free membranes composed of 32 lipids were simulated for about 80 picoseconds (van der Ploeg and
Berendsen 1982), the field of lipid simulations has matured to a stage in which the scales simulated in atomistic detail cover tens of nanometers (about 10^5–10^6 atoms) and several microseconds (Bjelkmar 2009; Dror 2009). This progress in atomistic simulations has been supported by the development of coarse-grained models and multiscale simulation techniques able to elucidate phenomena over scales much larger than the molecular ones (Ayton and Voth 2009; Murtola et al. 2009). The currently used particle-based coarse-grained models are appropriate for studies of systems of millions of particles over timescales of the order of 10–100 microseconds (Reynwar et al. 2007; Apajalahti 2010), and the situation continuously improves. Today, simulations can provide a great deal of insight into a variety of phenomena that are not tractable by experiments. Simulations are no longer used as tools for confirming what has been found in experiments; instead they have predictive power, guiding experiments to focus on novel phenomena. Current aims to bridge molecular simulations with computational systems biology foster the field further, coupling molecular and cellular phenomena to one another.

In this article, we provide an overview of lipid simulations, describing how a computer can be used as a laboratory for lipid research. We briefly discuss the methodology of lipid simulations followed by a number of topical applications that show the benefit of simulations. The applications given here as examples of simulations’ role for lipid research focus on the most recent state of the art and aim to provide a perspective of where we stand at the moment. A brief discussion of the prospects of lipid simulations closes this article.

SIMULATION METHODOLOGY IN A NUTSHELL

Extensive and detailed descriptions of computer simulation techniques as applied to molecular systems are many and available elsewhere (Tieleman et al. 1997; Frenkel and Smit 2002; Praprotnik et al. 2007; Murtola et al. 2009; Senn and Thiel 2009); thus, here we focus only on the essentials. The starting point of every simulation is to decide the questions one wants to clarify, because they determine the relevant length and timescales, and they in turn determine the technique that is most appropriate for a given problem. If the phenomenon one wishes to understand deals with electronic degrees of freedom, such as a chemical reaction because of lipases acting on a lipid, then the method of choice is quantum mechanics. For physical processes in which electronic properties are not important but which take place over molecular scales, the ideal technique is usually classical atomistic molecular dynamics (MD). At scales much larger than the molecular size, atomistic techniques are no longer reasonable, and one has to resort to coarse-grained (CG) models that usually represent the system as coarse-grained particles or with fields.

Here, we focus on two techniques that are most often applied to molecular simulations of lipid systems: atomistic and CG molecular dynamics. In both cases, there are three main steps that need to be accomplished to simulate a lipid system. First, given the molecular structures of lipids, one has to construct the initial configuration of a system one aims to study, such as a hydrated lipid bilayer. Second, to describe intramolecular and intermolecular interactions one needs the Hamiltonian (force field), whose parameters are often determined from quantum-mechanical simulations and experiments. Finally, to generate the evolution of the system in time, one has to integrate Newton’s equations of motion for the particle positions and velocities a sufficient number of times.

The heart of molecular modeling is the force field. The interactions are described by a potential energy function typically having the form presented in Figure 1. There are two types of interactions: bonded and nonbonded ones. Bonded interactions include bonding, bending, and dihedral (torsional) terms, whereas nonbonded interactions typically include electrostatic and Lennard-Jones (van der Waals) interactions. The set of these interactions, together with their parameter values, constitute the force field. There are many force fields used
in simulations of biomolecular systems, the most important ones being Amber, Charm, OPLS, and GROMOS (van Gunsteren et al. 2006). These force fields differ in how their parameters are derived, and they often have a different number of terms with different forms in the potential energy function.

For practical purposes, a number of more advanced algorithms are used to complement the above-mentioned three key items. For instance, to minimize artifacts because of a finite system size, one typically employs periodic boundary conditions that effectively allow one to consider a system of infinite size. Further, to model the system under conditions that mimic experimental ones, one often uses thermostats and barostats to maintain the temperature and pressure at desired values.

Although classical MD simulations can provide a great deal of insight into lipid systems, they obviously also have a number of limitations. First, as one is dealing with a classical method, one can only consider physical processes because chemical reactions cannot be investigated without accounting for electronic degrees of freedom. For the same reason, there are currently no effective and commonly accepted methods for simulations in constant-pH conditions because a classical description of protons is an issue. Instead, one models the effect of pH by fixing the protonation state of lipids and membrane proteins to the one that is consistent with a given pH. Another important limitation is the length and timescale accessible to atomistic and coarse-grained MD simulations. The scales currently accessible to atomistic simulations are about 20 nm × 20 nm × 20 nm and 1 μs. In CG descriptions, the scales depend on the level of coarse graining, that is, how much one has simplified the underlying atomistic representation, but some flavor of typical scales is gained by assuming that if the system size is the same as in atomistic models, then the timescale can usually be multiplied by a factor of 10^3 to 10^6 (Marrink et al. 2004; Murtola 2009; Ayton and Voth 2009). If these scales sound small, it is worthwhile to recall that only 10 years ago simulations of small lipid bilayers covered about 10 ns, and protein simulations only about 1 ns. The progress in biomolecular simulations has been truly impressive.

Nonetheless, to speed up atomistic lipid simulations even further, an obvious way is to design more efficient algorithms and their implementations. Particularly considerable progress has been made recently in the development and implementation of efficient schemes for parallel simulations. In this manner, the efficiency of the commonly used simulation engines such as GROMACS (Hess et al. 2008) and NAMD (Phillips et al. 2005) has been improved significantly, and currently both of these simulation packages allow one to use 500 to 1000 processors efficiently. Another
possibility is to develop algorithms that are computationally less expensive but still of high accuracy. A good example in this context is the Particle Mesh Ewald technique (Essman et al. 1995), which computes electrostatic interactions much faster than the original formulation based on the Ewald summation scheme.

In the field of coarse graining, the progress during the last decade has been quite impressive. For given CG representations of molecules, several techniques to determine the effective interactions have been developed. Two often-used schemes are based on matching either forces (Yanting et al. 2009) or structural properties (Murtola et al. 2009) between the coarse-grained and the underlying atomistic systems. An alternative technique is to consider the thermodynamic free energy for partitioning in a water-like or oil-like environment, and to parameterize the coarse-grained model in this spirit. The MARTINI model (Marrink et al. 2004, 2007; Monticelli et al. 2008; Lopez et al. 2009), using this idea, has turned out to be very successful in lipid simulations.

LIPIDS IN ACTION

Lips and Peptides and Membrane Proteins

Membrane proteins constitute a large and important class of proteins. As about 30% of the genome codes for membrane proteins, and because studies of their dynamics are a challenge for any experimental technique, it is not surprising that MD simulations have become one of the most versatile techniques to explore their structure and dynamics (Kandt et al. 1998; Khalili-Araghi et al. 2009; Klepeis et al. 2009). Membrane proteins come in two varieties: peripheral proteins loosely associated with the membrane–water interface and integral proteins spanning the membrane. Aside from these two, there are also peptides associated with membranes.

Peptides provide an excellent opportunity to show how MD simulations complement experiments because the timescales related to peptide dynamics are much shorter compared to the dynamics of more complex proteins. Simulations of peptides provide a means to explore in atomistic detail the molecular phenomena that take place in 1–1000 nanoseconds. To this end, let us consider one of the biologically relevant functions of peptides: the formation of pores across lipid membranes (Fig. 2). When pores are formed, they facilitate the translocation of ions and small molecules through a membrane. When that takes place, cells go through lysis and die. Antimicrobial peptides use this mechanism to kill cells.

Peptides interacting with membranes are typically amphipatic, that is, composed of hydrophobic and hydrophilic (usually lysine and...
arginine) amino acids. A good example is magainin 2, an antimicrobial peptide originating from the skin of an African frog *Xenopus laevis* (Cruciani et al. 1992). The helical structure of magainin is characterized by its apolar residues facing the membrane, whereas the polar residues are exposed to water. The cooperative self-assembly of five to six magainin molecules leads to the formation of a toroidal membrane-spanning pore. The first computational model of the pore constructed nearly 10 years ago showed that the peptide-induced pore complex stabilized itself in nanoseconds (Pasenkiewicz-Gierula et al. 2000; Murzyn et al. 2004). The mechanism by which the pore is formed was shown in a more recent study (Leontiadou et al. 2006). It was found that pore formation is a comparatively rapid process, taking place in about 10–100 ns, provided that there are a sufficient number of peptides nearby. Pores were found to be clearly permeable for water and lipids, but they were also observed to be rather disordered, showing greater variability in terms of peptide conformation and orientation than it was believed earlier. Similar studies were recently performed on another antimicrobial peptide, melittin, which also showed the disordered nature of pore formation (Sengupta et al. 2008). Overall, simulations have been able to shed light on the complex mechanisms by which pores form and function, providing atomistic and molecular-scale insight that is inaccessible to experiments.

Meanwhile, MD simulations of membrane proteins are computationally much more challenging compared to peptide simulations. This is largely because of the larger scales in time and space needed to simulate membrane proteins. Typical system sizes and timescales that are currently feasible are about 1,000,000 atoms simulated over a millisecond (Bjelkmar et al. 2009; Khalili-Araghi et al. 2009; Klepeis et al. 2009). The most commonly studied proteins are membrane channels including aquaporins, ion channels, and mechanosensitive channels. Simulations of these channels provide detailed information about the mechanisms associated with their conductance. For aquaporins, MD simulations were able to show a collective transport mechanism by which up to ten water molecules migrated through the channel in a concerted fashion; the dynamics of protein side chains played an important role in this process (Jensen et al. 2008). For the Na\(^+\)/H\(^+\) antiporter, MD simulations showed a detailed patch of Na\(^+\) ions and identified the key residues inside the channel that are responsible for transport (Arkin et al. 2007). These findings were confirmed in mutagenesis studies. A similar detailed picture has very recently been obtained for some of the K\(^+\) (Jensen et al. 2010) and mechanosensitive channels (Vásquez et al. 2008).

Although the membrane in the above-mentioned studies served only as an environment for the protein to carry out its function, it is known that lipids play an active role in protein functioning, which in turn is associated with specific lipid–protein interactions. This is highlighted by a recent simulation of rhodopsin in a three-component lipid bilayer composed of phosphatidylincholines, phosphatidylethanolamines, and cholesterol (Khelashvili et al. 2009). The simulation showed the distribution of cholesterol around rhodopsin to be nonuniform. Cholesterol was located next to the highly conserved GPCR family motive of the protein, stabilizing a specific conformation in this region (Khelashvili et al. 2009). In the same spirit for the Kv1.2 ion channel, one has found preferential binding of negatively charged phosphatidylglycerols (PGs): This enrichment of PGs was driven by interactions with the positively charged residues at the protein surface (Bjelkmar et al. 2009).

In the future, simulations are expected to provide further benefit for membrane science both in terms of unraveling phenomena associated with more complex proteins and in terms of considering more complex lipid compositions of membranes.

**Lipids and Proteins in Pharmacology**

When we are sick, a medical doctor makes a diagnosis and provides us with a prescription, and often the sickness is cured by the drugs we have been ordered.
Does the success in treating the sickness also imply that we understand the action mechanism of the drug? No. In many cases it is not known, for example, how the drug binds to other molecules thus inhibiting their function and causing (for instance) inflammation to recover. What atomistic simulations can do is provide insight into the phenomena associated with the actions of drugs. Here we discuss some possibilities in this context.

When drugs act on biological membranes, the existing drugs have usually been targeted to affect membrane proteins. However, this practice is changing. Recently, as the role of lipids in drug-induced effects has been noticed, more attention has been paid to lipids as significant targets of drugs (Lúcio et al. 2010). A good example of this progress is lipid therapy, in which a drug molecule acts on a membrane to change the lipids’ structure. In this way, they activate membrane receptors that in turn initiate a signaling pathway (Escriba 2006). Interestingly, the idea of lipid therapy is similar in spirit to the recent idea discussed by Cantor for the mechanism of general anesthetics (Cantor 1997; Jerabek et al. 2010). He proposed that anesthetic molecules could play a role in the activation of membrane proteins by changing the pressure profile of a lipid membrane. In this manner, lipids mediate interactions between anesthetics and proteins, as the force exerted on a protein by the membrane depends on the membrane composition. Another reason for the interest in drug–lipid interactions comes from the design of novel drug delivery systems based on lipids, such as liposomes and other lipid assemblies (Paasonen et al. 2007).

Computational methods have been used in drug design for a long time (Song et al. 2009). The most popular techniques have been structure–activity relationship and quantitative structure–activity relationship (Scior et al. 2009), as well as various docking methods based on scanning the protein surface and its cavities to find an optimal fit for the binding of a drug to the protein (Taylor et al. 2002). Currently, atomistic molecular dynamics simulations are not typically used in drug design because of the high computational cost. This is unfortunate because they can provide substantial insight into molecular mechanisms associated with the action of a drug and provide information about the interactions of a drug with other important biomolecules such as lipids.

To show the significant benefit of atomistic simulations, let us consider the MD simulation of β2-adrenergic receptor (β2-AR) with nebivolol (Fig. 3) (Kaszuba et al. 2010). β-ARs belong to the family of G-protein coupled receptors (GPCRs) that constitute about 80% of the known receptors for neurotransmitters, hormones, and neuromodulators, and about 5% of the genes in eukaryotic organisms (Vigh et al. 2005). A list of diseases related to GPCR is long and includes, among others, certain forms of blindness, obesity, inflammation, depression, cancer, and hypertension (Lúcio et al. 2010). Therefore, not surprisingly, these receptors are one of the main targets in pharmacology. β2-AR activity induces smooth muscle relaxation; thus, β2-AR blockade is commonly used in asthma inhalers (Taylor 2007). Meanwhile, nebivolol is a new, highly selective β1-AR blocker (Prisant et al. 2008) characterized by lack of side effects that are otherwise typical for β-blockers, and also lack of interactions with other drugs (Veverka et al. 2006). Nebivolol has four chiral centers and 10 stereoisomers, though only two of them are of pharmaceutical interest: the srrr-form and the rsss-form (Siebert et al. 2008).

Very recently, atomistic molecular dynamics simulations were used to unravel the issue of nebivolol selectivity toward one of the receptor’s subtypes, as well as to understand its stereo-specificity (Kaszuba et al. 2010). Simulations clearly showed that the srrr-form interacts more favorably with the receptor, demonstrating the capacity of atomistic MD simulations for determining stereo-specificity. Most interestingly, this effect was observed to arise from energetically favorable interactions with water molecules. This finding is profoundly interesting because it highlights the important role of hydration of the bonding pocket and, more generally, the importance of water in drug–protein interactions. This perspective is of...
general importance for drug design because most of the docking methods commonly used in computational studies of drug–protein complexes do not include water. Clearly, there is reason to revise the docking techniques to account for this aspect.

Lipoproteins as Carriers of Cholesterol
Cardiovascular diseases (CVD) are the primary cause of death in western countries (Mokhad et al. 2004), causing about one out of five deaths in the population. One of the main causes of this condition is atherosclerosis, that is, the lipid accumulation and plaque formation on arterial walls. This severe condition relates to lipoprotein particles that transport cholesterol in the body. High levels of low-density lipoprotein (LDL) particles in blood have been found to increase the risk of atherosclerosis (Castelli et al. 1986; Hevonoja et al. 2000), whereas high levels of high-density lipoprotein (HDL) are known to reduce the risk (Colvin and Parks 1999; Linsel-Nitschke and Tall 2005).

To understand the functions of HDL and LDL, one should first know their structures, but despite extensive studies, this issue has remained unclear. A number of different models have been proposed for HDL (Phillips et al. 1997; Segrest et al. 1999; Martin et al. 2006; Wu et al. 2009) and LDL (Hevonoja et al. 2000), but a large amount of uncertainty has remained because the nanoscale size of HDL and LDL and the thermally fluctuating nature of these nanoparticles have challenged the resolution of experimental approaches.

Meanwhile, the role of computer simulations in lipoprotein research has increased steadily, and a number of simulation studies about lipoproteins have been published recently. For example, the properties of the lipoprotein core have been studied both for cholesterol esters and triglycerides (Heikelä et al. 2006; Hall et al. 2008). HDL discs have also been paid considerable attention (Shih et al. 2005, 2008; Catte et al. 2006), and recent work on spheroidal HDL has been very illuminating (Catte et al. 2008; Koivuniemi et al. 2009). Especially, HDL simulations have been able to provide a great deal of insight into the atomistic structure of both the lipid droplet and the apolipoproteins ApoA-I surrounding the lipid part of HDL, and the important role of free cholesterol in stabilizing ApoA-I structure at the surface of HDL has become apparent.

A recent study (Yetukuri et al. 2010) shows the benefit of bridging molecular simulations to experiments particularly well. In that article, the authors considered two subject groups whose living habits were very different. One of...
the groups favoring a healthy diet and good living habits had a high level of HDL, whereas the other group with less healthy living habits had low HDL. High-throughput mass spectroscopy used to analyze the HDL lipid compositions revealed major differences, especially in the molar fractions of free cholesterol and triglycerides, illustrating that people with low HDL had reduced amounts of free cholesterol and larger amounts of triglycerides. The analyzed lipid compositions were employed in CG molecular simulations to compare the HDL structures of subjects in the two different groups. The results found via simulations were quite fascinating (Fig. 4): In low-HDL subjects, triglycerides, instead of being confined to the hydrophobic core of HDL, were observed to migrate to the HDL-water interface in which their concentration was markedly larger compared to the high-HDL case. At the surface region, the elevated triglyceride concentration was coupled to a reduced concentration of free cholesterol, implying that the likelihood of triglyceride-ApoA-I contacts increased at the expense of a reducing number of contacts between free cholesterol and ApoA-I. As ApoA-I has been found to prefer contacts with cholesterol, the excess triglyceride replacing cholesterol in low-HDL subjects has likely a role to play in the function of HDL.

Although more extensive studies are needed, it seems plausible that diet and living habits influence the structure of the lipid droplet in HDL. That, in turn, may have an effect on the structure of ApoA-I and, hence, the functions of HDL. In a more general context, the study (Yetukuri et al. 2010) illustrates the benefit of coupling clinical research to bioinformatics and systems biology, and further to molecular simulations.
Complex Concerted Dynamics in Membranes

The lipid raft model (Simons and Ikonen 1997) and the extensive research that has followed this proposition have provided a great deal of insight into the structural aspects of biological membranes. Meanwhile, the understanding of membrane properties is largely incomplete, as rather little is known of the dynamics associated with membrane domains, their formation through diffusion of lipids, and even the diffusion of lipids in model membranes.

Cell membranes bustle with dynamic phenomena at a wide range of characteristic timescales and length scales. Motion of lipids in the plane of the membrane—lateral diffusion—is the most widely studied of these, and yet the mechanism of lateral diffusion has been far from well understood. It has been thought rather commonly that lateral diffusion follows the jump-diffusion model in which diffusion consists of rapid “rattling” motion of lipids confined to cages formed by their neighbors, punctuated by nearly instantaneous, discrete jumps in which a whole lipid molecule moves out of its cage, moving a distance comparable to its size. Short-range techniques, such as quasi-elastic neutron scattering (QENS), have been thought to measure the rapid rattling-in-a-cage motion, whereas long-range techniques, such as fluorescence recovery after photobleaching, have been assumed to gauge the slower motion consisting of jumps between cages. Phenomenological free volume models based on the jump-diffusion model have been used to interpret the dependence of lateral diffusion coefficients on temperature and membrane composition (Galla et al. 1979; MacCarthy and Kozak 1982; O’Leary 1987; Almeida et al. 1992). However, no direct experimental evidence for the jumps as a dominating diffusion mechanism has been reported.

Recent atomistic and coarse-grained simulations have shed light on this issue (Falck et al. 2007; Apajalahti et al. 2010). Falck et al. considered single-component lipid bilayers and analyzed the trajectories of all lipids in detail. They did not find evidence for jump-like motion in which individual lipids would have rapidly moved a distance of their own size in the bilayer plane in a slowly changing environment. Instead, Falck et al. identified the motion of transient lipid clusters in which about ten lipids moved in unison as loosely defined clusters (Fig. 5). Over larger scales, they further observed transient dynamic correlations in which the motions of lipids were coupled to one another over scales of ~10 nm. In a more recent work (Apajalahti et al. 2010), coarse-grained simulations were used to consider similar phenomena in raft-like membranes over much larger scales in time and space. They observed the dynamical correlations in lateral motion to persist over a microsecond timescale. Recent simulation studies (Roark and Feller 2009) have also provided support to the concerted diffusion mechanism.

Importantly, the simulations by Falck et al. predicted that the lipid diffusion in membranes takes place through concerted motions of lipids as transient lipid clusters. Experimental data of these phenomena did not exist until early 2010 (Busch et al. 2010), when the Unruh group reported QENS data for lipid diffusion. Their data confirmed the predictions of simulations, highlighting the concerted motion of lipids as the mechanism of lipid diffusion.

Flow-like patterns have earlier been found in hydrodynamic two-dimensional systems in which the driving force has been the conservation of momentum. In lipid diffusion, momentum is not conserved within a membrane because lipids exchange momentum with the water phase. The origin of the dynamical correlations observed in membranes is therefore different: There are local and transient density waves that emerge spontaneously, giving rise to the observed large-scale motions. The findings highlight the importance of spontaneous collective fluctuations and suggest that similar phenomena are abundant also in other soft matter systems. Most recently, this has been observed for integral membrane proteins whose motion also takes place in a concerted manner with 50 to 100 lipids around it (Niemela et al. 2007).
This finding may have considerable significance to the dynamics and function of cell membranes, as it implies that the dynamics of proteins and lipids in cell membranes are not two separate issues but have to be considered together.

CONCLUDING REMARKS

Computer simulations of lipid systems have been performed since the 1980s. During these decades, the role of lipid simulations has increased significantly. Although in the early times simulations mainly confirmed what had been seen in experiments, the field of lipid simulations has now matured to a level in which it can also guide experiments by making predictions of novel phenomena not yet observed in experimental laboratories. For example, the concerted motions of diffusing molecules in lipid bilayers were first observed in atomistic simulations (Falck et al. 2007), and later confirmed in QENS experiments (Busch et al. 2010). Membrane protein structures can be resolved by crystallographic means, but the dynamics of the proteins can be investigated by computer simulations, providing a way to predict and clarify their activation mechanisms (Bjelkmar 2009). Simulations can also generate insight into phenomena that cannot be studied...
experimentally, such as in quantifying the magnitude and range of membrane perturbations induced by fluorescent lipid probes (Repakova 2005; Holttävuo 2008). Considering the benefit of bridging experiments with simulations, we are confident that this trend will be promoted, fostering better science.

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