Insertion and Expulsion Mechanism of an Amphiphile in a Membrane Mimetic

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Supporting Information

ABSTRACT: A theoretical study, validated with experimental data, on the mechanism of insertion/expulsion of an amphiphile from its aggregate is presented. Using molecular dynamics (MD), the equilibrium of tetradecyltrimethylammonium (TTA+) between the aqueous phase and a bilayer, with TTA+ as the main component, was simulated. In order to be inserted, the first interaction of TTA+ with the bilayer is with the hydrophobic chain, and more importantly, before any interaction, TTA+ has to adopt a stretched conformation along with a perpendicular orientation with respect to the aggregate surface. Otherwise, insertion fails and the amphiphile goes back into the bulk water phase. This conformation and orientation remains unmodified during the insertion and expulsion processes. Thus, this mechanism can be called the "dive in mechanism".

INTRODUCTION

The aggregation process of head/tail amphiphilic molecules in water has been deeply studied. Nevertheless, because of experimental difficulties, the equilibrium of amphiphiles between the aqueous phase and its dissolved aggregate is much less studied. In this regard, there are several studies about unimer equilibrium exchange kinetics between micelles. From the Halperin and Alexander theory, it is concluded that the equilibrium is mainly achieved through insertion and expulsion mechanisms of single unimers. To obtain atomistic details on the self-assembly process of amphiphilic molecules in water, many simulation studies have been performed. Although the insertion process of amphiphilic molecules into its aggregates has been studied, the mechanism steps for insertion and expulsion of unimers with a single chain carbon from its aggregate remains unknown.

How an amphiphile, similar to the ones used in drug delivery systems, as templates for synthesizing nano- and mesostructured materials, or in membrane mimetics, gets in or out from its aggregate is a basic question whose answer opens a new way for understanding the equilibrium of amphiphiles between aggregates and bulk solution, thereby giving new insights on how to manipulate the aggregation process and the stability of bilayers in general.

Nowadays exists a broad range of membrane mimetics to select according to the needs. The aggregates occupied in this study are lyotropic nematic systems, developed in our laboratory, that classify as magnetically alignable bicelles, sometimes used as membrane mimetics. They are formed by aggregates of amphiphiles that adopt an discotic average geometry, called bicelles. They are nematic because the order between the aggregates is only orientational (Figure 1). These systems have the advantage of being composed of bicelles, so the local morphology resembles those of biomembranes (flat bilayer). Additionally, because of their geometry, bicelles have an anisotropic diamagnetic susceptibility, which makes them align with a magnetic field, enabling the obtention of valuable information such as quadrupole splittings. They provide information regarding order and dynamics.

EXPERIMENTAL SECTION

Mesophase Preparation. The system was prepared by dissolving 0.3448 g of tetradecyltrimethylammonium chloride (TTAC), 0.1094 g of NaCl, and 0.0730 g of decanol (DeOH) in 0.995 mL of H2O. Fully deuterated sodium dodecylsulfate (0.012 g, SDS-d25) and deuterated water (5 μL, D2O) were added per mL of water to be used as 2H NMR probes. The system was equilibrated 48 h at 300 K before any measurement.

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Figure 1. Representation of lyotropic nematic systems composed of magnetically alignable bicelles. $\vec{B}$ represents the magnetic field.
SDS-\textit{d}_{25}, TTAC, and DeOH were purchased from Aldrich and used as received. Water of HPLC grade and NaCl at the highest purity available were purchased from Merck and used as received.

\textbf{\textsuperscript{2}H NMR Spectra.} \textsuperscript{2}H NMR spectra were obtained in a Bruker Avance 400 NMR spectrometer, located at the Universidad de Santiago de Chile, using a broadband probe tuned to 61.425 MHz, at 300 K. The \textsuperscript{2}H 90° pulse was 19 \textmu s long, and more than 1000 transients from a spectral window of 40 kHz were accumulated in 32 kB files.

\textbf{Polarized Light Microscopy (PLM).} A Motic series B microscope equipped with crossed polarizers was employed to observe the liquid crystal textures. The samples were placed in a concave slide with a depth of 1 mm and were allowed to orient in a 0.84 T magnet for 15 min. Photos were taken with a 20-fold magnification, from the center of the concavity, avoiding variations in the thickness of the sample due to the curvature of the slide, at 300 K.

\textbf{Computational Methods.} GROMACS package 4.5.3 was used, along with the GROMOS53A6 force field.\textsuperscript{15} For the Lennard-Jones interactions of the aliphatic chains, the force field developed by Berger et al. was used.\textsuperscript{16} Ryckaert–Bellemans potential was used in the dihedral potential to reproduce the frequency of the trans/gauche transition of aliphatic chains.\textsuperscript{17} These combinations of force fields have been demonstrated to successfully reproduce similar systems to the study here.\textsuperscript{18–21}

Before the single independent simulation, the steepest descent energy minimization was applied. The initial velocities were randomly generated according with Boltzmann distribution.

The time of simulation was 1.2 \mu s. The integrator was the leapfrog algorithm with an integration time step size of 2 fs. van der Waals interactions were simulated using the Lennard-Jones potential (LJ), and long-range electrostatic interactions were calculated using the Particle Mesh Ewald (PME) method.\textsuperscript{22} A cutoff of 1.0 nm was employed for the LJ and the electrostatic interactions. A linear constraint solver was used to restrain the bond lengths. Simulations were carried out in an NPT ensemble at 328 K and 1 atm, coupled to a modified Berendsen thermostat (\textit{v}-rescale) and a Parrinello–Rahman barostat, with time constants of 1 and 2 ps for temperature and pressure, respectively. This temperature was necessary to reproduce the experimental data. Simulations were performed with periodic boundary conditions in all directions of space. Simple point charge (SPC) water model was employed.

\section{RESULTS AND DISCUSSION}

\textsuperscript{2}H NMR spectra of the studied system along with their respective PLM textures were obtained and are displayed in Figure 2. The spectral profile and the Schlieren textures observed in PLM textures confirms that the studied system corresponds to a lyotropic nematic phase.\textsuperscript{23–27} The main component TTAC is in a concentration of 1.1809 M, largely above its CMC of 0.0047 M.\textsuperscript{28} The signal at the center, near 0 Hz, is from D_{2}O, and the rest are from the deuterium of SDS-\textit{d}_{25}. The separation in Hz between two signals coming from the same deuterium is the quadrupole splitting (\textit{\Delta}\nu_{Q}), see Figure 2.

From the theory of magnetic resonance, \textit{\Delta}\nu_{Q} is related to the so-called “order parameter” according to eq 1. In our case, the order parameter is from the bond between carbon and deuterium (\textit{S}_{CD}).
computational box is displayed in Figure 3. The computational box contained 60 TTA+, 12 DeOH, eight DS situated in the hydrophobic core were removed. The ensemble to stabilize the pressure. The simulation time was 1.2 ns, and an equilibrium was rapidly achieved, registered by the plateau in the potential energy and density plots of the system (Figures 3S and 4S). A representation of the simulated plateau in the potential energy and density plots of the system is displayed in Figure 3.

To validate the simulations, the experimental $\Delta \nu_{Q}$ (obtained from the spectrum in Figure 2) was satisfactorily reproduced, see Figure 4. To avoid the transient state, the simulated order parameters were obtained as an average of all the SD$^-$ between 10 ns and 1.2 $\mu$s. To obtain the simulated $\Delta \nu_{Q}$, GROMACS calculates the bond order parameter with respect to the three Cartesian axes of the box. Because bicelles orient with their principal symmetry axis perpendicular to the external magnetic field (see Figure 1), the magnetic field corresponds to $x$ and $y$ axes in the simulation. Thus, an average bond order parameter with respect to the $x$ and $y$ axes is used in eq 1 to calculate $\Delta \nu_{Q}$. From Figure 4 can be observed that $\Delta \nu_{Q}$ decreases as the carbons of SDS- $d_{15}$ get closer to the hydrophobic core; this because the orientational order, given by the interface interaction, decreases toward the interior of the aggregate. Reproduction of the experimental $\Delta \nu_{Q}$ could be achieved by raising the temperature to 328 K and halving the SD$^-$ charges. Reducing the charges is sometimes used in molecular dynamics, principally to compensate the absence of polarizability, which in conjunction with other factors overestimates electrostatic interactions.20,30

After an extensive literature search, no study regarding a simultaneous membrane that reproduces the equilibrium between amphiphiles in the bulk solution and amphiphiles in the aggregate was found. It is important to mention that even above the CMC some amphiphiles always remain in the bulk water solution.

Surprisingly, during the simulation, some TTA’s spontaneously got out of the bilayer and remained in bulk solution; however, after a few nanoseconds, they got back into the aggregate. To be sure this was not an artifact, a total of 1.2 $\mu$s was simulated. During this time, an equilibrium between TTA’s in the aggregate and in the bulk solution was achieved, see Figure 5.

During the simulation, 14 TTA’s spontaneously leave the bilayer and 12 return to it. Figure 5 shows the distance between the center of mass of the bilayer and the center of mass of each TTA that leaves the bilayer. Examining the trajectories, TTA’s always began insertion with the carbon of the tail, adopting different orientation and conformation (Figure 1S), but one of them is by large the most probable, corresponding to an orientation with the major principal axis perpendicular to the bilayer interface and a conformation with its chain fully extended. Figure 7 shows a picture of a process representative of the insertion and expulsion of a TTA$^+$ from the aggregate. The LJ energy of interaction between TTA’s and the bilayer together with the normalized projection of the major principal inertia moment in the $z$ axis ($I_{TTA}^{z}$) as a function of time were plotted (Figure 7). LJ energy of 0 means that TTA$^+$ is in the isotropic domain (bulk), and when it begins to decrease, in the analyzing context, TTA$^+$ started its insertion into the membrane. If $I_{TTA}^{z}$ is changing, meaning a free rotation, thus TTA$^+$ is in the bulk solution, and when this value is close to 1, the amphiphile is extended with a perpendicular orientation with respect to the plane of the bilayer. In Figure 7 (bottom), it can be observed that before TTA$^+$ gets in contact with the bilayer (LJ = 0) it adopts an orientation perpendicular to the bilayer with a fully extended conformation ($I_{TTA}^{z} = 1$) and remains that way during the entire insertion process.

Studying in detail the interaction between one TTA$^+$ and the bilayer while in the bulk domain, it is observed that TTA$^+$ has 99 independent contacts with the bilayer before insertion (LJ TTA$^+$/bilayer < 0), this during the 77 ns that TTA$^+$ is outside the bilayer (1.28 contacts/ns). The independence of each contact was established arbitrarily if it fulfills the condition of being 100 ps apart from another contact. A TTA$^+$ remains, in average, 68 ns in the bulk solution before its insertion; thus, one amphiphile has an average of 87 contacts (1.28 contacts/ns × 68 ns) before being inserted. Thus, the idea that an amphiphile has to be adsorbed by the interface and then immersed in the bilayer is not applicable in this case. Therefore, there must be a direct immersion into the hydrophobic core; otherwise, the insertion processes fails.

This behavior was expected mainly because the amphiphile head has the same charge as the bilayer surface, so by charge repulsion it should be the hydrophobic part of the molecule to first approach the interface. However, the requirement of having an extended chain and a perpendicular orientation, before any interaction, in order to be inserted is less expected. Even more, the carbon chain of the amphiphile in the bulk solution has a less stretched conformation (see Figure 6) and

![Figure 4](image4.png)

Figure 4. Simulated (circles) $\Delta \nu_{Q}$ and experimental (squares) $\Delta \nu_{Q}$ of SDS-$d_{15}$ as a function of carbon number.

![Figure 5](image5.png)

Figure 5. Distance between the center of mass of the bilayer and the center of mass of selected TTA’s as a function of time.
more mobility with respect to an amphiphile inserted into the aggregate (Table 1). To represent the decrease in the stretched conformation when the amphiphile is expelled from the aggregate, the moment of inertia around the principal, middle, and minor axes of a representative TTA+ is plotted as a function of time. To represent when the TTA+ is in the bilayer or in the bulk, the LJ energy between the TTA+ and the bilayer is also plotted, while TTA+ in the bilayer adopts a stretched conformation with a characteristic prolate profile observed in Figure 6. This profile is consistent with a moment of inertia around the major axis close to 0 and around the middle and minor axis with higher and similar values. When TTA+ is expelled (LJ = 0), the stretched conformation remains but with a significant decrease in its linearity, represented by a decrease of the inertia moment around the middle and minor axes and an increase of the moment of inertia around the principal axis.

The increase in TTA+ internal mobility when it leaves the bilayer is represented by the decrease of the dihedral order parameter \( S_{D}^{2} \) of the dihedral angles along the TTA+ carbon chain, see Table 1. \( S_{D}^{2} \) was calculated for the dihedral angles formed by four consecutive carbons conforming the TTA+ carbon chain, starting from carbon 1 (head) up to carbon 14 (tail). Each \( S_{D}^{2} \) was obtained from the dihedral autocorrelation function, from 20 ns of simulation with a time step of 2 fs.

To have a mental picture image, when Olympic divers are approaching water, they continuously change their orientation and conformation, but before entering the pool, they adopt the same conformation and orientation that TTA+ adopts before its insertion. Thus, we can call this mechanism the “dive in mechanism”.

## CONCLUSIONS

The mechanism of insertion of an amphiphile in its aggregate corresponds with the expulsion one. There is a strong correlation on the conformation and orientation of the amphiphile with its displacement in an axis perpendicular to the bilayer. A stretch conformation with an orientation perpendicular to the bilayer plane facilitates this displacement and allows the insertion or expulsion of the amphiphile from the bilayer aggregate. These observations point to establish that the mechanism of insertion or expulsion of an amphiphile from its aggregate has a geometrical director. With this in mind, comprehension of the aggregation process and linked equilibriums will be improved.

## ASSOCIATED CONTENT

### Supporting Information

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Notes

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