Dynamics Simulation of Langmuir-Blodgett Films

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DYNAMICS SIMULATION OF LANGMUIR-BLODGETT FILMS

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Introduction

Langmuir-Blodgett film monolayers are formed when amphiphilic molecules in solution are deposited on water surface followed by compression. During the compression, the randomly oriented molecules reorient themselves with the hydrophilic head groups on the water surface and the hydrophobic tail standing away from the water surface. The organic ultra-thin films with highly ordered layer structure, have shown a wide range of potential applications in biochemistry, microelectronics, integrated optics, microlithography, and many other fields [1-4]. It is very important to get a better understanding of the relationship between the properties of the films and the micro-structure, for example, the molecular conformation, packing, orientation and motion in the films. For this purpose, within the past few years, different computer simulation methods were applied on LB film monolayers, including Monte Carlo methods [2,3] and molecular dynamics simulations [4-6]. In this paper, we present the results of a molecular dynamics simulation of a fatty acid (CH$_3$(CH$_2$)$_7$COOH) monolayer on a water surface. This fatty acid is a preliminary model for our next study object: CH$_3$(CH$_2$)$_{15}$C=C-C(CH$_3$)$_2$COOH.

Methodology

QUANTA 2.0 (CHARMM 21), a comprehensive, integrated molecular modelling package developed by Polygen Corporation is used in the present simulation. In this calculation, the Steepest Descent method [7] is employed for energy minimization and the Verlet algorithm is employed for the dynamics simulation. The hardware employed is a GS1000 Graphics Supercomputer from Stellar Computer Inc.

The simulation starts with 169 molecules, in a 13x13 matrix, parallel to each other standing vertically on a layer of water. During the dynamics simulation, the position of the water molecules are frozen. A 15x15 molecule matrix (225 molecules) without water is also studied.

To relax the conformation and remove bad contacts, the system undergoes energy minimization at zero K. The potential energy terms concerned are:

$$E_p = E_{\text{bond \ length}} + E_{\text{bond \ angle}} + E_{\text{cyclic \ angle}} + E_{\text{improper \ torsion}} + E_{\text{electrostatic}} + E_{\text{van \ der \ waals}}$$

The energy for hydrogen bonding is included in electrostatic and van der waals terms.

The optimized system is then heated to the desired temperature, i.e. 300K, by gradually assigning random velocities to each atom. The relationship between the atom velocity, the kinetic energy and temperature of the system is:

$$E_{\text{kinetic}} = \frac{1}{2} \sum_{i=1}^{N} m_i v_i^2 = \frac{3}{2} Nk_BT$$

where $N$ is the total number of atoms and $m_i$ and $v_i$ are the $i$th atom's mass and velocity, respectively; $k_B$ is Boltzmann's constant.

When the system is heated to 300K, it undergoes an equilibration period to stabilize average temperature and structure. After the system is equilibrated, the final molecular dynamics simulation starts. The coordinates of the molecules are recorded periodically. The sets of coordinates recorded during the dynamics simulation are then analyzed to extract information about molecular packing, orientation and motion.

Results and Discussion

Water strongly limits the mobility of the molecules, especially the polar head groups. In Table I, the vertical mobility of the two end-groups of the molecules is represented by the fluctuations in height. Without water, both end-groups have similar fluctuations as shown in Table I and Fig. 1a. When water is introduced into the system, the surface formed by the polar groups (the lower curve in Fig. 1b) is more even due to their hydrophilic interaction.

During the simulation the molecules tend to adjust the intermolecular distance to reach the minimum potential energy. For closely packed systems (i.e. with smaller area per molecule), the final molecular areas of 21.22 Å$^2$ (see Table I) are consistent with that for 'close-packed heads' phase of fatty acids monolayers [8]. For closely packed system, there is pseudohexagonal packing with defects (Fig. 2). Relatively loosely packed systems however have significantly less packing order.

Comparing Fig. 3a and 3b we can see that the center molecule, surrounded by other molecules, has low mobility and flexibility, represented by the tilt and molecular length respectively; the molecule in the center of the molecule matrix has more mobility and flexibility. When loosely packed, there is more space around each molecule and the cohesion is weaker, therefore the molecules will have more tilt (see Table I) and stronger edge effect as shown in Fig. 4: the tilts for central molecules are less than 10° and for molecules at the matrix corner can be as high as 60°. To correct for the edge effect, simulations with periodic boundary conditions are being made and will be discussed.

Conclusion

Simulations of the LB films without periodic boundaries show clearly the effect of water and packing condition on the behavior of molecules in the monolayer. It is instructive to see appearance of domain boundaries and lattice defects in the simulation process as result already seen experimentally but whose origin is not well understood. The strong edge effect makes it necessary to introduce periodic boundaries in future simulations.
References

Table 1. The height fluctuation [Å] of the end-groups, the change of molecular area [Å²/molecule] and the average tilt [*] in the dynamics simulations with and without water. The simulation time scale is pico second [ps]

<table>
<thead>
<tr>
<th>Simulation</th>
<th>15x15 molecules with water</th>
<th>15x15 molecules without water</th>
</tr>
</thead>
<tbody>
<tr>
<td>Start</td>
<td>16 36 25 36</td>
<td>21 25 22 27</td>
</tr>
<tr>
<td>End</td>
<td>21 25 22 27</td>
<td>21 25 22 27</td>
</tr>
<tr>
<td>Molecular Area [Å²/molecule]</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Head Group</td>
<td>2.52 4.18 1.14 0.86</td>
<td>2.93 4.16 3.83 3.54</td>
</tr>
<tr>
<td>Tail Group</td>
<td>2.52 4.18 1.14 0.86</td>
<td>2.93 4.16 3.83 3.54</td>
</tr>
<tr>
<td>Average Tilt [*]</td>
<td></td>
<td></td>
</tr>
<tr>
<td>First 30ps</td>
<td>9.37 14.65(17ps) 12.35 20.02</td>
<td>18.44 30.00</td>
</tr>
<tr>
<td>Last 30ps</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Fig. 1 The time averaged height of the mass center and the two end-groups of the molecules. (a) Without water (21 Å²/molecule). (b) With water (22Å²/molecule).

Fig. 2 The projection of average mass centers of the molecules (open circle) on a water surface with molecular area 22Å²/molecule (starting from 25Å²).

Fig. 3 The molecular length [Å] and tilt [*] as a function of dynamics simulation time (in pico seconds) for (a) a center molecule and (b) a corner molecule in a loosely packed system (27Å²/molecule). 0° tilt means the molecule is vertical to water surface.

Fig. 4 The average tilt angle [*] of molecules in a loosely packed system vs. the molecule position in two dimensions with molecular area 27Å² for 30 ps simulation.