

Molecular Dynamics Simulations of the Interaction of Beta Cyclodextrin with a Lipid Bilayer

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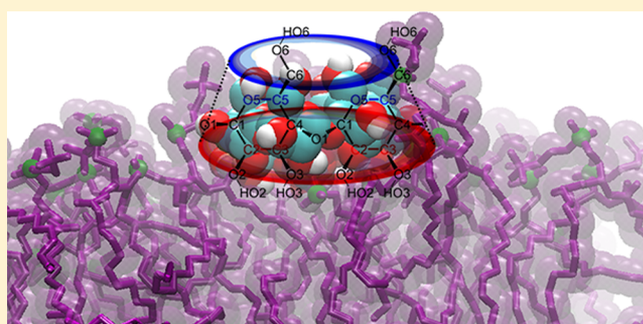
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S Supporting Information

ABSTRACT: Beta cyclodextrin (β CD) is well-known as a potent drug carrier improving drug solubility, stability, and bioavailability. The water layer adjacent to the membrane surface and lipophilic domain itself are a controlling barrier for drug transport. However, the molecular details of the interaction between β CD and the lipid membrane has not yet been clearly explained. Here, molecular dynamics simulations were performed to visualize the interaction process of the β CD molecule with the lipid bilayer for six microseconds in total. Our results show that β CD passively diffuses into the lipid bilayer by pointing its open secondary rim toward the lipid polar groups and then remains at the phosphate and glycerol-ester groups with hydrogen bond formation. The information obtained from this study may suggest that the association of β CD at the cellular membrane plays an important role for the transfer of drug and the extraction of cholesterol.



■ INTRODUCTION

Drug delivery systems (DDS) have been continuously developed with the aims to enhance therapeutic drug efficiency and consequently to reduce toxicity by the direct transportation to specific targets. Cyclodextrins (CDs) are used as potential drug carriers.¹ Natural CDs are α CD, β CD, and γ CD, composed of six, seven, and eight units of D-glucopyranose, respectively, with α -1,4 linkages (Figure 1). β CD is widely used in pharmaceutical and food industries because of low cost, easy synthetic accessibility, and suitable cavity size (0.60–0.65 nm) for the inclusion of small- and medium-sized drugs.² β CD and some derivatives have been used in inclusion complexation with drugs, and they have been approved and marketed especially for oral drug delivery.³ The structure of the β CD molecule is that of a truncated cone with a hydrophilic outer surface and a relatively hydrophobic nanocavity. Therefore, a poorly water-soluble molecule can be inserted into the CD's cavity leading to an increased solubility and consequently to a higher biological activity.

Based on biopharmaceutical drug classification, there are two main factors for efficient drug transport ability, the solubility and the permeation of the drug through biological membranes.⁴ Many experimental and theoretical studies have demonstrated that the aqueous solubility enhancement of lipophilic drugs by

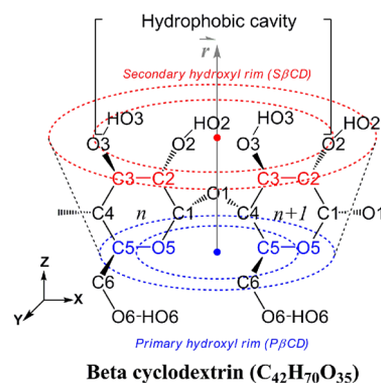


Figure 1. 2D structure of β CD showing the arrangement of glucose monomers. The atoms on the glucose ring representing the primary rim and the secondary rim are labeled in blue and red, respectively.

CDs inclusion leads to the increased drug efficacy.^{5–10} The β CD supported drug delivery may occur via two mechanisms. First, the drug dissociates from the β CD and then adsorbs into the membrane surface. Second, the drug- β CD complex

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directly transfers into the membrane in which the association and permeability of β CD complexes are possibly controlled by the barrier of the unstirred water layer (UWL) at the membrane surface.^{11,12} The drug- β CD complexes permeate through the UWL to reach the membrane surface via passive diffusion but may not get into the membrane interior due to their size and other physicochemical properties.¹¹⁻¹³ Actually, the binding of β CDs at the water-lipid interface has been experimentally proven by the extraction of cholesterol by β CDs.^{14,15} To date, there is no detailed study as to how the unstirred water layer and the membrane itself influence the association of the drug- β CD complex. As a very important prestep for the understanding of drug delivery at the membrane, the interactions of β CDs with the membrane have to be investigated.

Molecular understanding of the complex biological systems have been achieved by molecular dynamics (MD) simulations, which provided reliable and reasonable results comparable to experiments.¹⁶⁻¹⁹ The simulations in the lipid phase well described the interaction of small molecules with the lipid membrane.^{16,20-24} Wei et al. showed the agreement of computational and experimental studies for the translocation of ribose and its two diastereomers into 1-palmitoyl-2-oleoyl-*sn*-glycero-3-phosphocholine (POPC).²⁵ Moreover, the MD simulations have been extended to investigate physicochemical properties, host-guest interaction, and self-aggregation of sugar oligomer as cyclodextrins in both gas and solution phases.²⁶⁻³⁴ An understanding of CDs-based DDS as a drug loading can be carried out by the estimation of the binding affinity of drug to CD and complexity itself. The structure stability and phase solubility studies suggested that the hydrophobic molecule is bound properly in the cavity of CDs by 1:1,³⁵⁻³⁸ 2:1,^{37,39} and 2:2^{40,41} stoichiometry. The study of the complex architecture of β CD with hydrochlorothiazide (HCT) concluded that the formation of the HCT- β CD complex was enthalpy driven and the inclusion mode of HCT was highly dependent on its ionization state.⁴² On the other hand, the drug releasing may be described with the interactions of drug, CD, and the drug-CD complex with the lipid membrane.¹¹ The studies of the exposing cells or model membranes to CDs have shown that the membrane cholesterol was removed from the lipid rafts.^{11,14,15,43-49} The MD simulation also showed that β CD can rapidly adsorb on the membrane with a consequence of cholesterol extraction. The ability of cholesterol extraction by head-to-head dimer β CD depended on the orientation, the distribution of β CD on the membrane surface, and the concentrations of cholesterol in the membrane.^{48,49} The mechanism to remove cholesterol from the lipid membrane was described by Lopez et al.^{48,49} The proper conformation for cholesterol extraction is that the hydroxyl rim of the β CD dimer bound to the membrane surface with hydrogen bonds. Moreover, the extraction energy was significantly reduced at low concentration of cholesterol because the lipids prevented the β CD-cholesterol interactions.⁴⁹ Although the extraction of cholesterol by CDs has been studied to some extent,^{48,49} drug delivery and CD-lipid interaction are still not completely resolved. Therefore, a deeper understanding on β CD permeability in the biological membrane at the molecular level is required for the further development of CD-based drug delivery and drug release control. Here, we performed all-atom MD simulations with various starting geometries of β CD with regards to the POPC lipid bilayer model. The permeation behavior of β CD and its structure properties were of concern.

METHODOLOGY

System Preparation. MD simulations were performed with five different initial configurations as shown in Figure 2(a)-

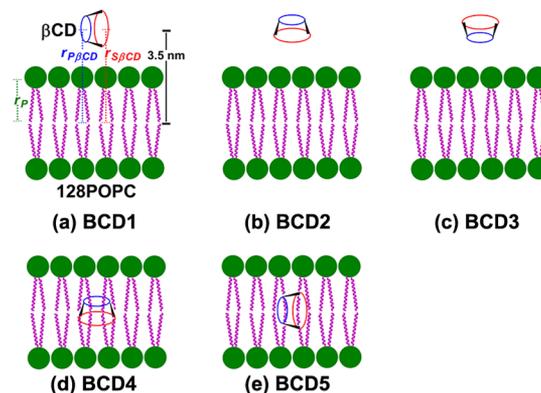


Figure 2. (a)-(c): Starting geometries of various orientations β CD at the membrane surface. (d)-(e): Starting geometries of various orientations β CD inside the membrane.

(e). In the BCD1-BCD3 systems, the β CD molecule with different orientations was located in the water phase at a distance of 3.5 nm in the z direction from the center of the bilayer. On the other hand, BCD4 and BCD5 represent the starting geometries when the β CD molecule is placed at the center of the lipid bilayer with parallel and perpendicular orientations, respectively. These setups allow us to study the behavior of β CD in the bilayer, and this technique has been widely used in the simulations of molecular transportation into the membrane.⁵⁰⁻⁵² In the case of BCD5, the hydroxyl rims of β CD closely interacted with lipid head groups. Note that this initial configuration may have an influence on the trajectory. A symmetric structure of the β CD molecule was taken from our previous study,⁵³ while the initial coordinates of the equilibrated 128 POPC bilayer were received from Tieleman's group (<http://wcm.ucalgary.ca/tieleman/downloads/popc128b.pdb>). All simulations were performed by using the GROMACS package version 4.5.5.⁵⁴ The POPC membrane was simulated by the modified Berger et al. parameters,^{19,55} and the β CD molecule was described by GROMOS 53A6⁵⁶ (the carbohydrate force field is equivalent to GROMOS 45A4⁵⁷). Note that a validation of the force field for β CD was carried out. The structural properties of β CD in water were compared with the result from the X-ray crystallography and also other MD simulations.⁵⁸⁻⁶¹ The general perspectives of β CD structural properties are in agreement with the previous studies as shown in Supporting Information Table S1. The POPC lipid bilayers were fully solvated by 7122 molecules of single point charge (SPC) water⁶² in the simulation box size of $6.42 \times 6.45 \times 9.19$ nm³.

Molecular Dynamics Simulation. After the energy minimization with the steepest descent algorithm for 10000 steps, classical MD simulations were carried out with NPT ensemble (the particle number, pressure, and temperature were kept constant) over one microsecond, except for the BCD5 system, where the simulation was extended to two microseconds. At least equilibrated trajectories over 500 ns were used for analysis. The integration time step was set at 2 fs, and the trajectories were sampled every 2 ps. The periodic boundary was applied in all directions. β CD, lipid, and water molecules were separately thermostated at 298 K by the Parrinello-Bussi

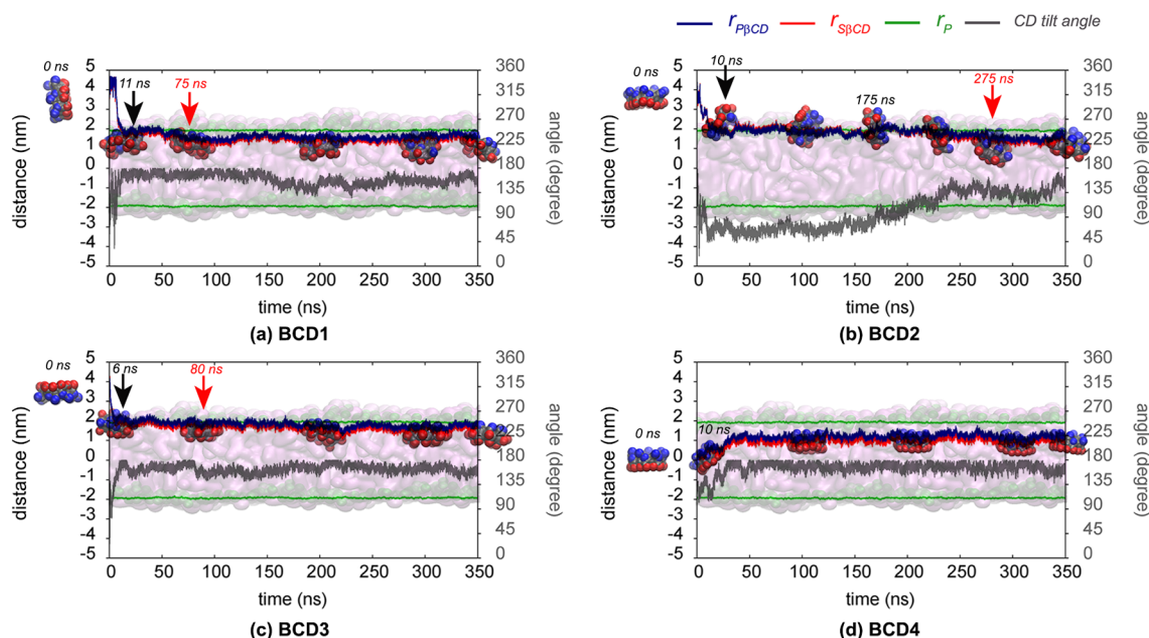


Figure 3. Closed-up for the first 350 ns of the permeation of β CD into the lipid bilayer for four simulations ((a) BCD1–(d) BCD4) using different initial structures. The distances as a function of time between the centers of the bilayer and both rims of the β CD molecule i.e. primary rim; $r_{P\beta CD}$ (blue) and secondary rim; $r_{S\beta CD}$ (red) are depicted. The β CD tilt angle and the distances of phosphate groups from the center of the bilayer for both leaflets (r_P) are plotted in gray and green, respectively. The snapshots of β CD at different simulation times are enclosed in the plots where the blue and red balls represent the atoms of the primary and secondary rims, respectively. The lipids of the bilayer are shown by purple rods, while their phosphorus atoms are depicted in green balls. The black and red arrows indicate the approaching and permeating time of β CD to the bilayer, respectively. The complete distances of the simulations are collected in Figure S1 in the [Supporting Information](#).

velocity rescale algorithm.^{63,64} Semi-isotropic pressure was applied by the Berendsen algorithm,⁶⁵ at an equilibrium pressure of 1 bar both in the xy -plane and in the z -direction (bilayer normal) with a time constant of 3.0 ps and a compressibility of $4.5 \times 10^{-5} \text{ bar}^{-1}$. The Lennard-Jones and the real-space parts of electrostatic interactions were cut off at 1.0 nm. The particle mesh Ewald (PME) method^{66–68} was used to compute long-range interactions with the reciprocal-space interactions evaluated on a 0.12 nm grid with cubic interpolation of order four. All bond lengths were constrained by the LINCS algorithm.⁶⁹ The used simulation protocol has been tested and validated to work well for lipid systems.^{70,71} Molecular visualizations were done using Visual Molecular Dynamics (VMD) software.⁷²

RESULTS AND DISCUSSION

The results of this study have been divided into three sections as follows: *I) The permeation and insertion of the β CD molecule into the lipid bilayer;* the β CD molecule permeation into the POPC lipid bilayer and the effect of the β CD molecule insertion on the bilayer were shown. *II) The interaction of the β CD molecule with the lipid bilayer;* the number of hydrogen bonds between the β CD molecule and the lipid bilayer were calculated to determine the binding interaction between β CD and lipids. In addition, the influence of β CD on lipid bilayer properties was analyzed in terms of a 2D-density map of β CD and POPC as well as local lipid thickness. In the last section, *III) The conformational change of the β CD molecule;* the conformation and structural stability of the β CD molecule at water, water–lipid interface, and lipid bilayer were considered.

I. The Permeation and Insertion of the β CD Molecule into the Lipid Bilayer. The β CD molecule was initially set far away from the bilayer center (around 3.5 nm) to avoid the bias

of the interactions between the β CD molecule and the lipid bilayer (as seen in Figure 2). This allowed the β CD to move and rotate freely in aqueous solution before approaching the bilayer surface. To study the permeation of the β CD molecule into the lipid bilayer, the distances in the z -axis of the β CD molecule ($r_{\beta CD}$) and its rims ($r_{P\beta CD}$ and $r_{S\beta CD}$ for primary and secondary hydroxyl sides, respectively) away from the lipid bilayer's center were determined as a function of time (as seen in Figure 3). The center of mass (COM) of all lipid molecules, the COM of all atoms in the β CD molecule, the COM of C5 and O5 atoms, and the COM of C2 and C3 atoms were defined as the bilayer center, the β CD center, the primary rim center, and the secondary rim center, respectively. The tilt angle of the β CD molecule is characterized by the angle between vector \vec{r} (the vector pointing from the COMs of the primary to the secondary rims, as seen in Figure 1) and the bilayer normal. The definitions of all mentioned parameters are illustrated in Figure 1 and Figure 2(a).

Figure 3 shows that the β CD molecule in the aqueous phase spontaneously moved toward the POPC bilayer and approached the bilayer surface within the first 15 ns. For the BCD1 and BCD3 systems, the β CD molecule attached to the bilayer with the secondary rim and readily permeated into the lipid bilayer within approximately 100 ns. The β CD remained underneath the phosphate group for the rest of the simulation time (Figure S1). For the BCD2 system, the β CD molecule first approached the bilayer surface with the primary rim, but it did not permeate into the bilayer over a hundred nanoseconds. After 175 ns simulation time, the β CD molecule rotated to have the secondary rim associated with the bilayer surface with the consequence of the β CD permeation within a few tens of nanoseconds as illustrated in Figure 3(b). After 500 ns, all β CD molecules are located in the region between phosphate and

glycerol-ester groups with the orientation of the secondary rim pointing toward the bilayer center, although starting from the various geometries (Figure 4). The averaged distances between

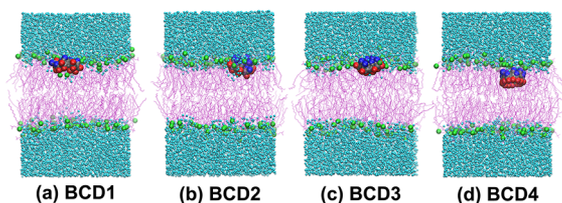


Figure 4. Last snapshots of (a) BCD1–(d) BCD4 simulations were extracted at 1 μ s. The primary rim of β CD, secondary rim of β CD, phosphorus atoms of lipid, and water molecules are presented by blue, red, green, and cyan spheres, respectively. The purple lines represent the POPC lipid molecules.

COMs of the β CD molecule and the bilayer as well as the tilt angle of the β CD molecule with respect to the bilayer are presented in Table 1. In agreement with our results, in the simulations of cholesterol extraction from the lipid monolayer or the bilayer, the head-to-head β CD dimer adsorbed on the membrane surface, however; the dimer oriented its primary rim toward the bilayer surface instead.⁴⁹ Based on spectroscopic study and *ab initio* calculations, Mascetti et al. suggested that the β CD molecules preferred to stack in parallel to the pure cholesterol, mixed 1,2-dimyristoyl-*sn*-glycero-3-phosphoglycerol (DMPG)/cholesterol, and mixed 1,2-dimyristoyl-*sn*-glycero-3-phosphocholin (DMPC)/cholesterol lipid monolayers.¹⁵

Next, the behavior of the β CD molecule and the effect of β CD insertion inside the bilayer were studied. Therefore, β CD was placed at the bilayer's center with the r^- vector alignments in parallel (BCD4) and perpendicular (BCD5) to the bilayer normal. For the BCD4 system, the β CD molecule is able to form hydrogen bonds with the glycerol-ester groups of the lipid bilayer and moved toward the water–lipid interface within 10 ns (in Figure 3(d)). The equilibrium location of β CD was 1.0 nm from the bilayer center corresponding to the position of the glycerol-ester group. No bilayer deformation has been observed within a simulation time of one microsecond. Unlike the BCD5 system (Figure 5), the β CD molecule was initially oriented in perpendicular to the bilayer normal; therefore, hydroxyl groups of the β CD formed hydrogen bonds with the lipid heads of both bilayer leaflets. At a longer simulation time, the β CD molecule rotated to be in parallel to the bilayer normal, and a water pore across the lipid bilayer could be formed (Figure 5(b)).

II. The Interaction of the β CD Molecule with the Lipid bilayer. Hydrogen Bonding Interactions. The number of hydrogen bonds (H-bonds) between β CD, its rims, and the components of lipid head groups (phosphate and glycerol-ester moieties) were calculated using the usual geometric restrictions

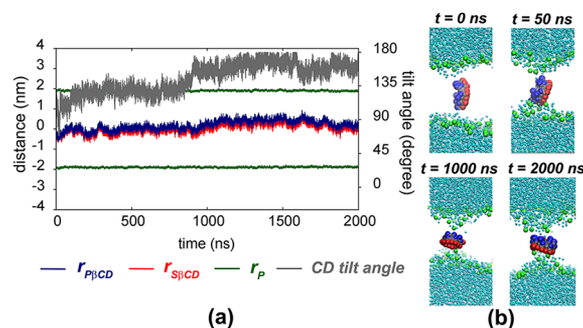


Figure 5. (a) Dynamics behavior of β CD starting from the perpendicular orientation in the lipid bilayer (BCD5). (b) Water pore formation induced by β CD at a longer simulation time, where the cyan sphere represents the water molecules, while the blue, red, and green spheres are symbolized as the atoms on primary and secondary rims of β CD and phosphorus of lipid head groups, respectively.

for hydrogen bonding. A hydrogen bond is defined by the distance between the donor and the acceptor (r_{HB}) < 0.35 nm and the deviation from the linearity < 30°. The distance value of 0.35 nm corresponds to the first minimum of the radial distribution function (RDF) of water. Figure 6 shows the

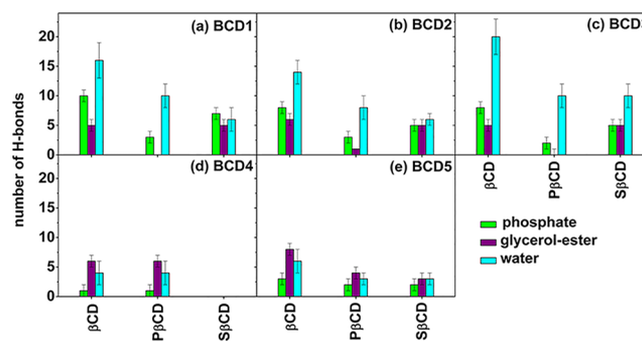


Figure 6. Hydrogen bond contribution of the interactions between the β CD molecule and phosphate (green) and glycerol-ester groups (purple) of the lipid and the solvating water molecules (cyan) for the systems (a) BCD1–(e) BCD5.

averaged number of hydrogen bonds at the last 500 ns and 1 μ s for the systems of BCD1–BCD4 and BCD5, respectively. The β CD molecules in the BCD1–BCD3 systems interacted with the lipid headgroup with the hydroxyl groups at the secondary rim, and it was more preferred to form hydrogen bonds with phosphate groups than with glycerol-ester groups. On the other hand, this finding may suggest that the β CD interacted with the lipid surface stronger than the β CD dimer in the cholesterol extraction.⁴⁹ In the BCD4 and BCD5 systems, the primary rim of β CD is associated with the glycerol-ester group of the lipid bilayer. This is in good agreement with the results of the

Table 1. Average Values of the Focused Parameters on β CD–Lipid Interactions for All Simulations (BCD1–BCD5) over the MD Production Period

systems/properties	$r_{\beta CD}$ (nm)	$r_{P\beta CD}$ (nm)	$r_{S\beta CD}$ (nm)	tilt angle (degree)
BCD1 (500–1000 ns)	1.50 \pm 0.10	1.54 \pm 0.10	1.40 \pm 0.10	163.6 \pm 8.0
BCD2 (500–1000 ns)	1.55 \pm 0.13	1.61 \pm 0.13	1.49 \pm 0.13	148.8 \pm 7.6
BCD3 (500–1000 ns)	1.65 \pm 0.12	1.72 \pm 0.12	1.54 \pm 0.12	167.4 \pm 5.9
BCD4 (500–1000 ns)	1.08 \pm 0.13	1.17 \pm 0.13	0.96 \pm 0.13	157.0 \pm 11.1
BCD5 (1000–2000 ns)	0.30 \pm 0.14	0.37 \pm 0.15	0.20 \pm 0.12	161.4 \pm 7.4

simulation snapshots and the permeate depths of the β CD molecule into the bilayer. Interestingly, the number of hydrogen bonds of β CD with water molecules significantly decreased from 16 to 5 when the β CD molecule translocated from water–lipid interface into the bilayer. In conclusion, the results of the hydrogen bonds suggest the translocation of β CD deeply into the hydrophobic region of the lipid bilayer is unfavorable.

The Influence of the β CD on the Lipid Membrane Properties. To study the influence of β CD on the bilayer structure, the two dimension (2D) density on the xz -plane of the β CD molecule, some POPC components (i.e., phosphate groups and glycerol-ester groups), and water were plotted in Figure 7 as well as the plots on the xy -plane in Figure S2

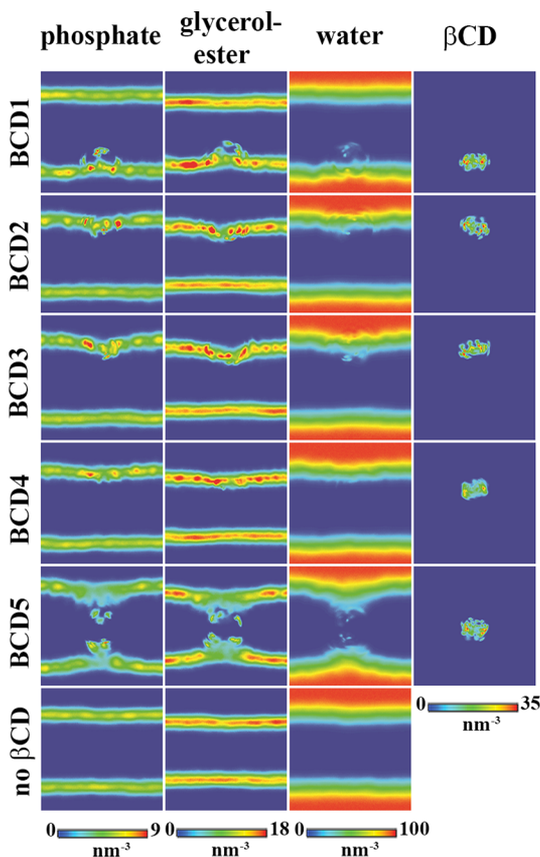


Figure 7. 2D-density maps on the xz -plane of the phosphate groups, glycerol-ester groups, water, and the β CD molecule for the BCD1-BCD5 systems, while those of the simulation of the POPC bilayer without β CD were given for comparison.

Supporting Information. The local thickness of the POPC bilayer on the xy -plane was also analyzed by means of the Voronoi algorithm⁷³ as shown in Figure 8. All analyses were computed from the last 100 ns of trajectories after removing the motion of the β CD molecule. The adsorption of one β CD molecule on the lipid bilayer cannot deform the membrane structure as seen by the insignificant change of 2D-density maps and the bilayer thickness of the BCD1-BCD3 ($r_{\beta\text{CD}} \sim 1.50$ – 1.65 nm) relative to the simulation of the POPC bilayer without β CD. The association of β CD on the lipid surface slightly induced the decrease of membrane thickness at the β CD location. When the β CD stayed underneath the glycerol-ester groups in the BCD4 system (the distance from the bilayer

center, $r_{\beta\text{CD}}$ of ~ 1.08 nm) it caused less disturbance on the lipid bilayer. In contradictory, the tilted β CD in the BCD5 system could bind with the lipid head groups causing the dramatic change in lipid thickness and inducing a water pore formation. In addition, the 2D-density maps (Figure 7 and Figure S2) could confirm that the β CD molecule in the BCD1-BCD3 systems preferred to stay at the bilayer surface with the hydroxyl rim tilted toward the bilayer center. The density maps on the xy -plane (Figure S2) showed that a few head groups of POPC molecules occupied inside the β CD cavity and 7–9 POPC molecules bound outside the β CD cavity. The ability of the cyclodextrin to uptake lipid or cholesterol from the biomembrane had been intensively demonstrated by *in vitro*, *in vivo*, and *in silico* studies.^{47,74–76} This extraction may induce the pore formation or membrane change which had been reported by the leakage of potassium and hemoglobin at high concentration of cyclodextrins.⁴⁷

Moreover, to investigate the dynamical property of the β CD influence on the POPC bilayer, we calculated the diffusion coefficient of the β CD molecule. The mean squared displacement (MSD) for 100 ns at the last of the BCD1-BCD5 systems and also the simulation of β CD in water were plotted and then fitted based on Einstein's relation $\langle r^2 \rangle \sim 4Dt$, where D is the diffusion coefficient, and t is time. The calculation is presented in the average of the diffusion coefficient by fitting of MSD every 20 ns. The results showed that the β CD diffusion coefficient in the water of our simulation is $(0.280 \pm 0.170) \times 10^{-9}$ m²/s. This value is in the same magnitude as the experimental data.⁷⁷ The diffusion coefficient of β CD in the lipid phase, $(0.019 \pm 0.006) \times 10^{-9}$ m²/s, is 1 order of magnitude smaller than in water. The slow diffusion of β CD in the lipid phase is related to strong binding of β CD with the lipid molecules.

III. The Conformational Change of the β CD Molecule.

To investigate the conformational change of the β CD molecule, we determined the intramolecular hydrogen bonds obtained by the number of hydrogen bonds of the secondary rim between the adjacent glucoses. The intramolecular hydrogen bonds are related to the stability of the CD molecule.^{78,79} The number of intramolecular hydrogen bonds of the BCD1-BCD3 systems was decreased continuously until there were no bonds left when the β CD molecule was attaching to the bilayer surface. The loss of this interaction was due to the preferable hydrogen bond formation between β CD and lipid head groups as well as water molecules^{59,80–82} (Figure 6) and resulted in the structural deformation as shown in Figure 9. The root-mean-square displacement (RMSD) of the hydroxyl side chains on each rim and glycosidic oxygens (O1) with respect to its anhydrous structure (in Table 2) showed that the primary rim of β CD was more flexible than the other one.⁸⁰ The solvating water molecules affected the flexibility of the β CD structure for both side chain rims and the O1 core. In addition, the area of the β CD cavity was calculated by the following equation

$$A = \frac{\pi}{7} \sum_{i=1}^7 r_i^2 \quad (1)$$

where r_i is the distance between each hydroxyl group and the center of the O1 atoms, and the hydroxyl groups at 6- and 3-positions are used for representing the cavity area of primary and secondary rims, respectively. The cavity areas of the β CD molecule in water, water–lipid interface, and lipid phase are summarized in Table 2. As expected for the truncated cone

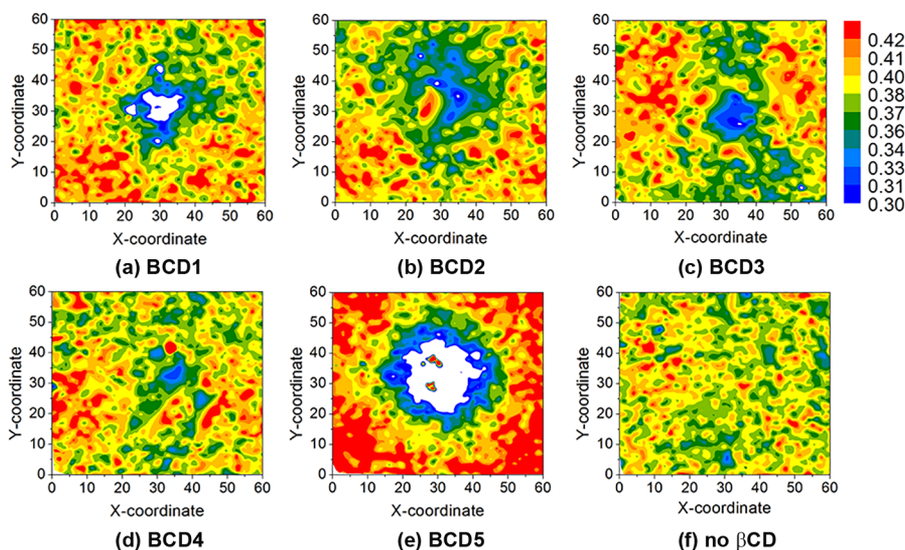


Figure 8. Local lipid thickness of the POPC bilayer in the systems with and without β CD ((a) BCD1–(e) BCD5 and (f) no β CD) where the β CD was located around the box center.

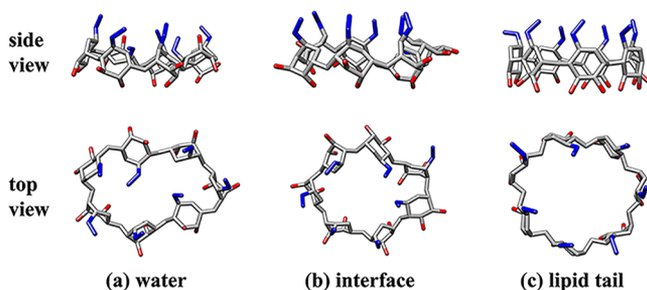


Figure 9. BCD in water was independently simulated for 500 ns, and the last MD snapshot was shown in (a). The last snapshots of β CD molecules obtained from the simulations of BCD3 and BCD4 for representing the β CD structure in lipid–water interface and lipid tail region as (b) and (c), respectively. The primary and secondary rims of the β CD molecule were labeled in blue and red, respectively. The backbone atoms of β CD molecules were represented in gray.

geometry of the β CD molecule, the cavity area of the secondary rim was larger than that of the primary rim about 1.5-fold in water and lipid phases, whereas the shape of the β CD molecule in the lipid tail resembled a cylinder. The area cavities of the secondary rim in the lipid phase were $\sim 15\%$ and $\sim 25\%$ smaller than in water and water–lipid interface, respectively. These findings may be related to the mechanism of drug release in which the open and closed states of the secondary rim of the β CD molecule at different phases may play an important role.

CONCLUSION

Based on the MD simulations of the β CD molecule with the lipid bilayer in the microsecond time scale, we found that the β CD molecule spontaneously permeated toward the lipid

surface but did not further proceed into the bilayer tail region in agreement with free energy calculations.⁴⁹ No structural distortion of the bilayer could be observed when the β CD molecule attached to the bilayer surface. In contrast, the water pore formation in the lipid bilayer possibly occurred when the β CD molecule stayed inside the bilayer. The simulation results also showed that the secondary rim of the β CD molecule mainly contributed to the β CD–lipid interactions in which 5–7 hydrogen bonds between the secondary rim and the lipid heads were always found for the whole length of simulation time. Interestingly, the conformations of the β CD molecule changed at various solvation phases (water, water–bilayer interface, and bilayer) because of the loss of the intramolecular hydrogen bonds. An enlargement of area cavity at the water–bilayer interface could be observed. These findings might be related to the mechanism of drug releasing; however, there is still more work to be done in order to understand the function of β CD as a potential drug carrier.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.jcim.5b00152.

Figures S1 and S2, Table S1, and references (PDF)

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Table 2. RMSD and Cavity Area of the β CD Molecule on Each Hydroxyl Rim

systems	$r_{\beta CD}$	RMSD (nm)			cavity area (nm ²)		
		O1	1st	2nd	O1	1st	2nd
in water		0.12 ± 0.02	0.33 ± 0.04	0.25 ± 0.02	0.93 ± 0.04	1.19 ± 0.12	1.56 ± 0.11
at interface	1.59 ± 0.10	0.10 ± 0.01	0.29 ± 0.01	0.23 ± 0.01	0.86 ± 0.03	1.36 ± 0.06	1.69 ± 0.03
in lipid tail	1.00 ± 0.10	0.06 ± 0.01	0.17 ± 0.02	0.08 ± 0.02	0.80 ± 0.02	1.35 ± 0.07	1.35 ± 0.04

Notes

The authors declare no competing financial interest.

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