



Systematic Characterization of DMPC/DHPC Self-Assemblies and Their Phase Behaviors in **Aqueous Solution**

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Abstract: Self-assemblies composed of 1,2-dimyristoyl-sn-glycero-3-phosphocholine (DMPC) and 1,2-dihexanoyl-sn-glycero-3-phosphocholine (DHPC) form several kinds of structures, such as vesicle, micelle, and bicelle. Their morphological properties have been studied widely, but their interfacial membrane properties have not been adequately investigated. Herein, we report a systematic characterization of DMPC/DHPC assemblies at 20 °C. To investigate the phase behavior, optical density OD_{500} , size (by dynamic light scattering), membrane fluidity $1/P_{\text{DPH}}$ (using 1,6-diphenyl-1,3,5-hexatriene), and membrane polarity GP_{340} (using 6-dodecanoyl-N,N-dimethyl-2-naphthylamine) were measured as a function of molar ratio of DHPC (X_{DHPC}). Based on structural properties (OD_{500} and size), large and small assemblies were categorized into Region (i) ($X_{DHPC} < 0.4$) and Region (ii) ($X_{DHPC} \ge 0.4$), respectively. The DMPC/DHPC assemblies with $0.33 \le X_{\text{DHPC}} \le 0.67$ (Region (ii-1)) showed gel-phase-like interfacial membrane properties, whereas DHPC-rich assemblies ($X_{\text{DHPC}} \ge 0.77$) showed disordered membrane properties (Region (ii-2)). Considering the structural and interfacial membrane properties, the DMPC/DHPC assemblies in Regions (i), (ii-1), and (ii-2) can be determined to be vesicle, bicelle, and micelle, respectively.

Keywords: phospholipid assembly; bicelle; membrane fluidity; membrane polarity; phase behavior

1. Introduction

Phospholipids form a self-assembled membrane in aqueous solution to reduce the exposure of hydrophobic acyl chain groups. 1,2-Dimyristoyl-sn-glycero-3-phosphocholine (DMPC, C14:0) molecules form a bilayer vesicle (liposome), whereas 1,2-dihexanoyl-sn-glycero-3-phosphocholine (DHPC, C6:0) molecules form a micelle at a total lipid concentration higher than the critical micelle concentration (cmc: 15 mM [1]). Depending on the chemical structure of the phospholipids, the self-assemblies form various morphologies such as vesicle, micelle, etc. [2,3]. In general, the self-assembly behavior of amphiphilic molecules is discussed based on the critical packing parameter, in which the head group area, length of hydrocarbon chain, and volume of molecules are dominant factors that determine the self-assembly structure. Owing to the hydrophobic region at the interior



of the membrane, phospholipid assemblies are utilized as a platform to integrate various kinds of nanomaterials [4]. The lipid bilayer structure is desirable to study membrane proteins; generally, discoidal self-assemblies, called "bicelles" [5,6], are applied to reconstruct membrane peptides and proteins [7–9]. Basically, a bicelle structure is composed of a mixture of lipid and detergent, as in DMPC/DHPC systems [10]. Since the stability of a bicelle is sensitive to the lipid composition and temperature [11–14], it is important to investigate the thermodynamic behavior of bicelles.

The cmc value of DHPC is ca. 15 mM [1,10], whereas the critical vesicle concentration (cvc) of DMPC is ca. 6 nM. Due to the difference in critical aggregation concentrations, the stability of a bicelle is also sensitive to the total lipid concentration [10–12,15,16]. Upon dilution of a small bicelle assembly, the transformation from bicelle to vesicle was observed. To avoid this, a chemically modified detergent can be utilized, making the bicelle structure stable toward heating or dilution [17,18]. To characterize bicelle properties, experiments have been carried out with high concentrations of lipid (1–10%, w/w); by employing such conditions, the structural properties of bicelles can be evaluated using cryo-electron microscopy (Cryo-EM) [10,15]. With regard to the rupture of bicelles in the dilution process, a direct deposition of lipids onto a solid support, wherein the bicelles are used as carriers for lipid sources, becomes more successful [19,20]. Statically, the phase behavior of lipid mixture systems (e.g., DMPC/DHPC) can be controlled by focusing on parameters such as lipid composition, lipid concentration, and temperature. Then, the self-assembly morphology can be controlled in a dynamic process, such as heating/cooling and dilution, based on the phase equilibrium. However, the phase behavior of DMPC/DHPC mixture systems at low lipid concentrations (total lipid < 20 mM) has not been adequately investigated.

Interfacial membrane properties such as membrane fluidity and membrane polarity, which are evaluated using fluorescent probes, are also important factors to characterize self-assemblies [21–23]. In lipid bilayer vesicles, the interfacial properties can be used to determine the phase states [22,24]; the solid-ordered phase (gel phase) and liquid-disordered phase (liquid-crystalline phase) show quite different membrane fluidity and polarity. These properties reflect the microscopic phase state and phase separation behavior [24]. Saturated phospholipids are in gel phase below the phase transition temperature ($T_{\rm m}$), whereas the membrane turns into liquid-crystalline phase at temperatures above the $T_{\rm m}$. To date, bicelles (such as DMPC/DHPC mixture systems) have been hardly studied based on their interfacial properties, especially at low lipid concentrations (less than 1% lipids (ca. 20 mM). In fluorescent probe studies, the interfacial membrane properties can be investigated at a total lipid concentration of 100 μ M [21,24]. Therefore, this could be suitable to evaluate the interfacial membrane properties and structural properties of DMPC/DHPC mixture systems.

The packing state of a lipid membrane is dominantly related to the function of the membrane: for example, molecular permeability and interaction of proteins [25,26]. In our previous study, significant chiral selective amino acid adsorption onto a liposome membrane was observed in a gel phase [27]. The state of a lipid bilayer can be classified as gel phase (most ordered state, like a solid), ripple-gel phase (ordered state), and liquid-crystalline phase (disordered state). Considering the membrane fluidity and membrane polarity of micelles [28], they can be categorized as a disordered membrane. In gel phase, the surface pressure is up to ~50 mN/m in monolayer systems (planar membrane (no curvature)), whereas it decreases to ~30 mN/m in vesicular bilayer systems [29]. This may be due to the membrane curvature of the vesicles, which could decrease lipid packing. It is hence worthwhile to investigate the interfacial properties of such a flat bilayer system.

The aim of this study was to clarify the self-assembly behavior of DMPC/DHPC mixtures containing various total lipid concentrations and fractions of lipids. The DMPC/DHPC assemblies were prepared with different molar fractions of DHPC (X_{DHPC}). Analyses of the turbidity (optical density, OD_{500}) and size distribution (using dynamic light scattering (DLS)), and of the interfacial membrane properties, using fluorescent probes, 1,6-diphenyl-1,3,5-hexatriene (DPH) and 6-dodecanoyl-*N*,*N*-dimethyl-2-naphthylamine (Laurdan), were systematically carried out.

The obtained results are summarized as a diagram to show the relationship between the lipid concentrations and morphologies of the DMPC/DHPC self-assemblies.

2. Materials and Methods

2.1. Materials

DHPC and DMPC ($T_m = 23 \,^{\circ}$ C) were purchased from Avanti Polar Lipids, Inc. (Alabaster, AL, USA). DPH and Laurdan were purchased from Sigma-Aldrich (St. Louis, MO, USA). Sodium dihydrogenphosphate (anhydrous) and disodium hydrogenphosphate were purchased from Wako Pure Chemical (Osaka, Japan), and were used to prepare phosphate buffer (50 mM, pH 7.0). Ultrapure water was prepared with the Millipore Milli-Q system (EMD Millipore Co., Darmstadt, Germany). Other chemicals were used without further purification.

2.2. Vesicle Preparation

DMPC vesicles were prepared as a reference (standard self-assembly), based on literature [24]. Briefly, a chloroform solution of phospholipids was dried in a round-bottom flask by rotary evaporation under vacuum. The obtained lipid films were dissolved in chloroform once more, and the solvent was evaporated. This operation was repeated at least twice. The obtained lipid thin film was kept under high vacuum for at least 3 h, and then hydrated with phosphate buffer at room temperature. The obtained DMPC suspension was frozen at -80 °C and then thawed at 40 °C (over the $T_{\rm m}$ of DMPC); this freeze-thaw cycle was repeated five times. Notably, the DMPC vesicle suspensions were extruded through 2 layers of polycarbonate membranes with a mean pore diameter of 100 nm, using as extruding device (Liposofast; Avestin Inc., Ottawa, ON, Canada). The prepared unilamellar vesicles were kept at 4 °C until use.

2.3. Preparation of DMPC/DHPC Self-Assembly in an Aqueous Solution.

Lipid thin films of DMPC/DHPC mixtures were prepared by the mothed described on the above. Firstly, the self-assembly solution was prepared at the total lipid concentration of 20 mM, with various DHPC fractions, X_{DHPC} : 0.17–0.95. The obtained lipid thin films were hydrated with phosphate buffer at 20 °C. No mechanical treatments (sonication, extrusion) were applied for DMPC/DHPC assemblies, to keep their spontaneous structures. In dilution of the DHPC/DMPC solution (20 mM), an aliquot amount of buffer solution was gently added to adjust the total lipid concentration, wherein the final lipid concentrations were set at 15 mM, 10 mM, 9 mM, and 8 mM. The obtained self-assembly solutions were applied for monitoring the optical density (see Section 2.4), and dynamic light scattering (see Section 2.5). If necessary, the extrusion (see Section 2.2) was performed before fluorescent spectroscopic measurements (see Sections 2.6 and 2.7).

2.4. Turbidity Measurements of Self-Assembly Solution

To assess the morphological insights for DMPC/DHPC assemblies, the turbidities of DMPC/DHPC mixtures at 500 nm (OD_{500}) were monitored by UV-1800 Spectrophotometer (Shimazu, Kyoto, Japan), in various total lipid concentrations. A thin quart cell (light path length: 1 mm) was employed to record the varied turbidity. An increased turbidity in the suspension could be evidence for the growth of self-assembly in size [30–32]. Thus, the morphological transition from bicelles to vesicles could be assessed by a jump-up of the OD_{500} values, in dilution.

2.5. Dynamic Light Scattering

The apparent sizes of mixtures (total lipid concentration: 20 mM) were determined by dynamic light scattering (DLS). Measurements were performed with the particle size analyzer (Zetasizer Nano ZS, Malvern Panalytical, Grovewood Rd, UK). The average diameters were calculated based on a number-average diameter.

2.6. Evaluation of Membrane Fluidity

As interfacial properties, the membrane fluidity was investigated by using DPH, based on previously described methods [21,33]. DPH was added to DMPC/DHPC assemblies with a molar ratio of lipid/DPH = 250/1. After incubation for 30 min, the fluorescence polarization of DPH was measured using a fluorescence spectrophotometer (FP-8500, Jasco, Tokyo, Japan) (Ex. = 360 nm, Em. = 430 nm). Fluorescence polarizers were set on the excitation and emission light pathways. With the emission polarizer angle of 0°, the fluorescence intensities obtained with the emission polarizer angle 0° and 90° were defined as I_{\parallel} and I_{\perp} , respectively. With the emission polarizer angle of 90°, the fluorescence intensities obtained with the emission polarizer angle of 90° were defined as i_{\perp} and i_{\parallel} , respectively. The polarization (P_{DPH}) of DPH was then calculated by using the following equations:

$$P_{DPH} = \left(I_{\parallel} - GI_{\perp}\right) / \left(I_{\parallel} + GI_{\perp}\right)$$
 $G = i_{\perp} / i_{\parallel},$

where *G* is the correction factor. The membrane fluidity was evaluated based on the reciprocal of polarization, $1/P_{\text{DPH}}$.

2.7. Evaluation of Membrane Polarity

Fluorescent probe Laurdan is sensitive to the polarity around itself, which allows the local polarity in lipid membranes to be determined [34,35]. Laurdan emission spectra exhibit a red shift caused by solvent attack and by solvent relaxation, thus its emission spectrum reflects the polarity (hydration state) of the self-assembly membrane. The Laurdan emission spectra were measured with an excitation wavelength of 340 nm, and the general polarization (GP_{340}), the membrane polarity, was calculated as follows:

$$GP_{340} = (I_{440} - I_{490}) / (I_{440} + I_{490}),$$

where I_{440} and I_{490} are the emission intensities of Laurdan excited at 340 nm. The obtained emission spectrum was furthermore analyzed by using Peakfit software (v.4.12, Systat Software Inc., San Jose, CA, USA) [28]. After deconvolution, the area fraction of each component was compared.

3. Results and Discussion

3.1. Systematic Characterization of DMPC/DHPC Assemblies Focusing on Turbidity, Size, Membrane Fluidity, and Membrane Polarity.

For DMPC/DHPC suspensions at a total lipid concentration of 20 mM, the solution turbidity varied depending on the X_{DHPC} . The transparency of the suspension was dependent on the size of the assemblies; the vesicles were relatively large (diameter > 30 nm: Region (i)), whereas the micelles or bicelles were small (diameter < 30 nm: Region (ii)) (Figure 1a). By employing fluorescent probes (DPH and Laurdan), the membrane fluidity $(1/P_{DPH})$ and membrane polarity (GP_{340}) values were determined. The DHPC micelle was categorized as disordered membrane $(1/P_{DPH} > 6, GP_{340} < 0.3)$. At 20 °C (< $T_{\rm m}$ of DMPC); some small assemblies (in Region (ii)) as well as the gel-phase DMPC vesicle showed rather ordered interfacial membrane properties (Figure 1b). The group of Region (ii) can be further divided into "ordered phase (ii-1)" and "disordered phase (ii-2)." The bicelle of DMPC/DHPC probably consisted of a micelle-like edge (rich in DHPC) and bilayer (rich in DMPC); therefore, the ordered phase found in Region (ii-1) could have been derived from the bilayer region of the bicelle. To classify each self-assembly, the threshold values of size, OD_{500} , $1/P_{DPH}$, and GP_{340} are summarized in Table 1. Based on the systematic characterization, a phase diagram of DMPC/DHPC at 20 mM is described in Figure 1c. The assembled states of DMPC/DHPC were confirmed as vesicle (Region (ii)), bicelle (Region (ii-1)), and micelle (Region (ii-2)).



Figure 1. Systematic characterization of DMPC/DHPC assemblies at 20 °C, total lipid concentration of 20 mM. As a function of DHPC fraction (X_{DHPC}), (**a**) turbidity (OD_{500}) and size distribution. Region (i), relatively large assemblies ($OD_{500} > 1.0$, size > 30 nm); Region (ii), relatively small assemblies ($OD_{500} > 1.0$, size > 30 nm); Region (ii), relatively small assemblies ($OD_{500} < 0.1$, size < 30 nm); (**b**) membrane fluidity ($1/P_{DPH}$) and membrane polarity (GP_{340}). Region (ii-1), ordered assemblies ($1/P_{DPH} < 6$, $GP_{340} > 0.3$); Region (ii-2), disordered assemblies ($1/P_{DPH} > 6$, $GP_{340} < 0.3$). (**c**) Phase diagram of DMPC/DHPC at total lipid concentration of 20 mM. Region (i), *closed circle*; Region (ii-1), *open circle*; Region (ii-2), half-tone circle.

	Size (DLS)	Turbidity, OD ₅₀₀	Membrane Fluidity, 1/P _{DPH}	Membrane Polarity, GP ₃₄₀
Region (i)	>30 nm	≥ 0.1	<6	>0.3
Region (ii-1)	\leq 30 nm	<0.1	<6	>0.3
Region (ii-2)	$\leq 9 \text{ nm}$	<0.1	>6	<0.3

Table 1. Summary of the results for self-assemblies prepared at different regions.

3.2. Structural Properties of DMPC/DHPC Assemblies in Dilute Conditions

The turbidities of DMPC/DHPC suspensions are dependent on the morphology. Micelle solutions are transparent, whereas vesicle, cylindrical micelle, and holey lamellar suspensions are turbid [10]. The DHPC micelle solution, wherein the average micelle size was 3–4 nm, was transparent. This suggests a rough correlation between turbidity and size of the self-assembly; therefore, the OD_{500} (Figure 2) and size distribution (Figure 3) were investigated at different total lipid concentrations.

3.2.1. Optical Density (OD₅₀₀)

Figure 2 shows the OD_{500} values for DHPC/DMPC assemblies with different X_{DHPC} ratios, and at various total lipid concentrations. As a trend, a higher OD_{500} value was obtained with an X_{DHPC} value lower than 0.4, in the total lipid concentration range of 8 to 20 mM. A similar tendency was observed at a total lipid concentration of 1 mM, although the OD_{500} value was relatively low. At any total lipid concentration, a critical X_{DHPC} value, with an increase in the OD_{500} value, was observed. Herein, the X_{DHPC} range for "turbid" suspensions is shown as Region (i), and that for "transparent" ones as Region (ii). For the samples with high total lipid concentration, the value shifted to a higher DHPC fraction;

the critical X_{DHPC} values for 10, 8, and 1 mM systems were around 0.4, 0.7, and 0.9, respectively. It was therefore considered that an increase in the ratio of long-chained DMPC molecules (decrease in X_{DHPC}) caused the formation of larger self-assemblies. Consequently, the morphology of DMPC/DHPC assemblies is roughly indicated by the turbidity; the formation of small-sized self-assemblies can be confirmed using the transparency of the suspension ($X_{\text{DHPC}} > 0.6$). It was thus shown that there were two types of regions from the viewpoint of turbidity of the solution.



Figure 2. Turbidities (OD_{500}) of DHPC/DMPC solution at different DHPC fraction (X_{DHPC}) in various concentrations of total lipid concentration at 20 mM (*circle*), 15 mM (*square*), 10 mM (*diamond*), 8 mM (*triangle*), and 1 mM (*cross*) at 20 °C.

3.2.2. Average Size Estimated Using DLS Analysis

The size distribution of the self-assemblies formed in the DMPC/DHPC mixtures was confirmed using DLS. As an example, the size distribution of the self-assembly at a total lipid concentration of 8 mM is shown in Figure 3a. In all the conditions tested, a mono-dispersed distribution was obtained. The average size of the self-assembly was affected by the variation in the X_{DHPC} value. As shown in Figure 3b, the average size of the DHPC/DMPC assembly was plotted against the DHPC fraction at various concentrations of total lipids. Self-assemblies with larger sizes (>1 µm) were obtained for lower DHPC fractions ($X_{DHPC} < 0.3-0.4$) at high total lipid concentrations (10–20 mM). In contrast, the average sizes were 10–20 nm when the X_{DHPC} was higher than 0.4. A similar tendency was observed for a total lipid concentration of 8 mM, except for the self-assembly with $X_{DHPC} = 0.4$. With a threshold value of 100 nm, self-assemblies were size-dependently classified: larger-sized assemblies (>100 nm) as Region (i) and smaller-sized assemblies (10–20 nm) as Region (ii). The critical X_{DHPC} values are denoted as dotted lines in Figure 3b, and a similar tendency was observed in OD_{500} analysis (Figure 2). Particularly, in Region (ii), a self-assembly with a much smaller size (less than 10 nm) was observed at a high DHPC fraction ($X_{DHPC} : ~0.95$).

Although the OD_{500} values cannot provide precise information about self-assembly size, an increase (or decrease) in the OD_{500} value is suitable evidence to confirm the transformation of one state to another, especially in a continuous measurement (heating/cooling and dilution). Hence, the self-assembly that appeared in Region (i) was considered a vesicle, and that in Region (ii) as a micelle or micelle-like small assembly (i.e., bicelle). In the case of the DMPC/DHPC bicelle, a jump in the OD_{500} value was observed upon heating [7], and can be applied to judge the existence of the bicelle. Among the self-assemblies categorized in Region (ii), two samples were picked to investigate dynamic changes in the OD_{500} values upon heating (Figure S1 in the Supporting Information). At a total lipid concentration of 20 mM, the DMPC/DHPC assembly with $X_{DHPC} = 0.4$ showed a significant increase

in the OD_{500} value at 30 °C. At $X_{DHPC} = 0.95$, the OD_{500} value did not significantly vary upon heating and cooling, showing that the assembly state of the DMPC/DHPC assembly at $X_{DHPC} = 0.95$ was stable, as a micelle. It has been reported that a bicelle in an aqueous solution is not very stable; it transforms into a vesicle-like structure upon heating, and reversibly transforms into a bicelle upon cooling [36].



Figure 3. (a) size distributions of DMPC/DHPC self-assemblies (8mM); 8 mM $X_{DHPC} = 0.17$ (a-1), 0.4 (a-2), 0.67 (a-3), and 0.95 (a-4); (b) average size of the self-assembly in the solution at different DHPC fractions; total lipid concentration 20 mM (circle), 15 mM (square), 10 mM (diamond), and 8 mM (triangle). A line between the arrows to show the region (i) and (ii), described in the top of figure, indicate the critical DHPC fraction that give a change in the turbidity of the solution, where the turbid solution was changed to the transparent one with increasing DHPC ratio, and region (i) shifts rightward with the dilution.

In summary, it was estimated that a vesicle could form at Region (i), whereas a micelle or bicelle could form at Region (ii). Although it is difficult to distinguish a bicelle and micelle based on their sizes, a variation in the OD_{500} can be a clue to identify bicelle formation. Considering these findings, the self-assemblies in Region (ii) were further categorized into Region (ii-1) (i.e., bicellar assembly) and Region (ii-2) (i.e., micellar assembly). To obtain more appropriate evidence, the interfacial properties can be helpful.

3.3. Interfacial Membrane Properties of DMPC/DHPC Assemblies

DPH and Laurdan are widely used as membrane-bound fluorescent probes, and indicate membrane fluidity and membrane polarity, respectively. The combined use of DPH and Laurdan has been successful in investigating the membrane properties of various kinds of self-assemblies, except those of bicelles [24,35,37]. It is hence worthwhile to investigate the membrane fluidity and polarity of DMPC/DHPC assemblies of various compositions and concentrations.

3.3.1. Membrane Fluidity $(1/P_{\text{DPH}})$

DPH is a non-polar molecule, which could be inserted into both the DHPC micelle and the DMPC vesicle. By monitoring the fluorescence polarization of DPH as a parameter $(1/P_{\text{DPH}})$, the disordered

state (liquid-crystalline phase, $1/P_{\text{DPH}} > 6$) and gel phase ($1/P_{\text{DPH}} < 6$) can be distinguished [24]. The membrane fluidities ($1/P_{\text{DPH}}$) of DMPC/DHPC assemblies were investigated as shown in Figure 4. At a total lipid concentration of 20 mM, an increase in $1/P_{\text{DPH}}$ values (>6) was observed at high DHPC fractions ($X_{\text{DHPC}} > 0.77$). At a total lipid concentration less than 15 mM, an increase in the values was observed at DHPC fractions higher than 0.91. Region (ii) could be further divided into two regions: Region (ii-1) with $1/P_{\text{DPH}} < 6$ and Region (ii-2) with $1/P_{\text{DPH}} > 6$. The pure micelle of DHPC was categorized as Region (ii-2), revealing the disordered state of the micelle membrane. In Region (ii-1), the self-assemblies are small, and their membranes are in ordered states. Such ordered states could be due to the enrichment of DMPC because only DMPC molecules can possibly form a gel-phase bilayer at this composition. Thus, in Region (ii-1), the DMPC/DHPC assemblies were shown to form ordered phases such as a gel phase, showing that the DMPC/DHPC bicelle was consistent with the ordered bilayer. The critical value of DHPC fraction to distinguish the two regions gradually shifted from $X_{\text{DHPC}} = 0.7$ (15–20 mM in total lipid concentration) to 0.9 (1 mM). DPH prefers to bind to the ordered phase [24]; therefore, the DMPC/DHPC assemblies with $1/P_{\text{DPH}}$ value higher than 6 were completely in a disordered state, similar to the liquid-crystalline phase.



Figure 4. Membrane fluidities of DMPC/DHPC assemblies. Total lipid concentration 20 mM (*circle*), 15 mM (*square*), 10 mM (*diamond*), 8 mM (*triangle*), and 1 mM (*cross*). The drastic change of each value shows transition of lipid assembling state at $1/P_{\text{DPH}} = 6$. The vertical dotted line in the figures displays the phase transition region (ii-1)/(ii-2) at each concentration: (ii-1) ordered phase, (ii-2) disordered phase.

3.3.2. Membrane Polarity (*GP*₃₄₀)

In DMPC/DHPC assemblies with $X_{\text{DHPC}} > 0.77$ (at 20 mM), the DMPC molecules did not form a gel phase (Figure 1b). In other words, the micellar structures were not perturbed by the presence of relatively longer acyl chain molecules (DMPC) at X_{DHPC} values > 0.77, suggesting that DMPC molecules, having longer acyl chains as compared to DHPC, are able to diffuse in micelle membranes freely. Laurdan, having a C12 hydrocarbon chain, can also diffuse in the self-assembly membrane freely, and its emission spectrum can be used to monitor the polar environment of each DMPC/DHPC assembly.

According to the packing of the lipid membrane, the emission properties of Laurdan can vary; thus, Laurdan is widely used to investigate the microscopic polarity at the lipid membrane

interface [35]. At any concentration of total lipids, the GP_{340} values did not change at low DHPC fractions ($X_{\text{DHPC}} < 0.4$) (Figure 5). However, the GP_{340} values varied depending on the total lipid concentration. In this study, the GP₃₄₀ values of DMPC vesicle and DHPC micelle were used as the standard: DMPC vesicle, $GP_{340} = 0.40$ (gel phase); DHPC, $GP_{340} = -0.20$ (disordered phase). With high total lipid concentrations (15 and 20 mM), the GP_{340} values slightly increased at the range of 0 $< X_{\text{DHPC}} < 0.5$, suggesting a more ordered state (Region (ii-1), bicelle). The GP₃₄₀ values gradually decreased in proportion to the DHPC fraction at $X_{DHPC} > 0.6$. At the middle concentration levels (8–10 mM), the GP_{340} values were constant at 0 < X_{DHPC} < 0.8, and varied at X_{DHPC} > 0.8. The results of the *GP*₃₄₀ analysis were also helpful to identify the bicelle and micelle of DMPC/DHPC assemblies at total lipid concentrations less than 20 mM. The coexistence of DHPC could disturb the ordered properties of the DMPC vesicle [38]. As compared to the pure DMPC vesicle, a slight increase in GP_{340} values was observed for DMPC/DHPC assemblies at X_{DHPC} of 0.33 or 0.4, suggesting more ordered (tightly packed) states of these membranes. The Laurdan emission spectrum was further analyzed by deconvolution (Figure S2 in the Supporting Information). At X_{DHPC} values of 0.33 and 0.4, the area fraction of λ_2 (440 nm) was slightly higher than that of pure DMPC, revealing a more stable gel phase in the bilayer region of the DMPC/DHPC assembly.



Figure 5. Membrane polarities of DMPC/DHPCassemblies; total lipid concentration 20 mM (*circle*), 15 mM (*square*), 10 mM (*diamond*), 8 mM (*triangle*), and 1 mM (*cross*). The drastic change of each value shows transition of lipid assembling state at $GP_{340} = 0.3$. The vertical dotted line in the figures displays the phase transition region (ii-1)/(ii-2) at each concentration: (ii-1) ordered phase, (ii-2) disordered phase.

3.3.3. Comparison of Interfacial Membrane Properties in the Cartesian Diagram

By combining the information obtained from each fluorescent probe (DPH and Laurdan), the phase state and phase separation behavior of the membrane can be systematically discussed [22–24,28]. The phase state (ordered phase or disordered phase) of the membrane can be investigated based on the positioning of the obtained data in the Cartesian diagram, where the quadrants are set using the transition of the membrane properties (polarity (*y*-axis) and membrane fluidity (*x*-axis)) [24]. By plotting the data obtained from each self-assembly system, the interfacial membrane properties of various kinds of self-assemblies can be directly compared [23,28]. The details are summarized in our previous work (see the supporting materials of Bui et al. [22]). In Figure 6, the DHPC membranes show a significantly higher membrane fluidity and a low GP_{340} value, which appeared in the boundary

region of the first (partially disordered state) and fourth quadrants (fully disordered state). This clearly reveals that the DHPC micelle membrane is in a disordered state, showing higher fluidity.



Figure 6. Comparison of interfacial membrane properties in the Cartesian diagram. T = 20 °C, DMPC/DHPC at 20 mM (*filled circle*), 10 mM (*diamond*), and 1 mM (*cross*). The value in the bracket neighboring to the data point show the DHPC fractions X_{DHPC} . The position plotted in the Cartesian diagram indicates the phase state of each membrane: the first and fourth quadrants for disordered phase, and the second quadrant for ordered (gel) phase. The reference values of various liposomes were plotted on the diagram (empty circle); s_0 : solid ordered phase, l_0 : liquid ordered phase, l_d : liquid disordered phase. The broken line crossing at $1/P_{DPH} = 6$ and $GP_{340} = 0.3$ is a boundary of ordered phase and partially-disordered phase.

For DMPC/DHPC assemblies with various X_{DHPC} and total lipid concentrations, the obtained GP_{340} and $1/P_{\text{DPH}}$ values were plotted in the Cartesian diagram, together with some standard data (obtained from liposomes, in our previous work). The data for the DMPC/DHPC mixtures (20 mM) at rather low DHPC fractions ($X_{DHPC} = 0.4$ and 0.67) were plotted in the second quadrant, suggesting that these self-assemblies had an ordered bilayer as well as DMPC vesicles in the gel phase. In contrast, at $X_{\text{DHPC}} = 0.77$, the data position was in the first quadrant. This indicates that the membrane fluidity was higher than that of the DHPC micelle, whereas the membrane was less hydrated as compared to the micelle. Since DHPC can behave as a detergent (surfactant), a mixed micelle [39,40] might be formed in DMPC/DHPC mixture systems, but it is still difficult to compare the difference between a bicelle and mixed micelle. The variation in the membrane properties at $X_{DHPC} = 0.7$ suggests that the assemblies were in disordered phases. It was difficult to distinguish the bicelle and micelle based on their sizes; in contrast, their interfacial membrane properties were clearly different. Hence, the DMPC/DHPC assemblies with $X_{DHPC} \ge 0.77$ were in a disordered phase, wherein DMPC molecules did not form (ordered) a bilayer. In addition to the unique structural characteristics of the bicelle, the interfacial properties also contribute to the investigation of the "ordered state" of the bicelle composed of DMPC/DHPC mixtures.

Limited to DMPC/DHPC systems, the bicelle assemblies required the interfacial membrane properties of the gel phase. By defining these assemblies as "gel-bicelles," a micelle (Region (ii-2)) and gel-bicelle (Region (ii-1)) can be clearly distinguished based on their interfacial properties. Similarly, a systematic characterization was carried out for the DMPC/DHPC assemblies at total lipid concentrations of 1-10 mM. While the $1/P_{\text{DPH}}$ and GP_{340} values at $X_{\text{DHPC}} = 0.77$ slightly changed,

the interfacial membrane properties of the assemblies with X_{DHPC} of 0.4 and 0.67 remained constant. In vesicle systems, the GP_{340} values varied depending on the temperature: $GP_{340} > 0.3$ in gel phase $(T < T_{\text{m}})$, and $GP_{340} > -0.2$ in fluid phase $(T > T_{\text{m}})$ [4]. The GP_{340} value gradually decreases at temperatures slightly higher than the T_{m} . When the GP_{340} value is lower than 0.3, the membrane is heterogeneous (under phase transition, coexistence of two phases). These results imply that ordered membranes $(1/P_{\text{DPH}} < 6, GP_{340} > 0.3)$ were maintained at X_{DHPC} of both 0.4 and 0.67, at low total lipid concentrations. This is the first investigation of the interfacial membrane properties of DMPC/DHPC

3.4. Discussion on Phase Behavior of DMPC/DHPC Assemblies

DMPC/DHPC self-assemblies were classified into three categories: Region (i), Region (ii-1), and (ii-2). The assemblies of Region (ii-1) were small micelles, which exhibited an ordered phase similar to that of the gel-phase vesicles. Hence, they can be considered as gel-phase bicelles. The phase behavior of the DMPC/DHPC assemblies at 20 °C is shown as a diagram (Figure 7). The self-assemblies at Region (i) were observed, typically, with excess DMPC and less DHPC, due to which the DMPC/DHPC mixtures could form vesicles independent of the total lipid concentration. On the contrary, the transparent solutions including the self-assemblies of small size (less than 10 nm) were observed in the condition with excess DMPC and less DMPC (Region (ii-2)). These concentration levels correspond well with the well-known and usual conditions to prepare the DMPC liposome and DHPC micelle.

assemblies. We found that the bicelle membranes (Region (ii-1)) are in ordered states; the bilayer

membranes are tightly packed as well as or better than the pure gel phase of DMPC vesicles.



Figure 7. Phase diagram for DMPC/DHPC assemblies at 20 °C, investigated by systematic characterization. The bold filled squares showed the increasing turbidities each DHPC fractions as the morphological change. The phase boundary concentrations of $X_{DHPC} = 0.77$, 0.4, and 0.33, are 9 mM, 12 mM, and 16 mM, respectively (described in total lipid concentration). Blue and Orange circles are transparent assemblies due to forming small assemblies. These each had membrane properties; (blue) ordered phase, (orange) disordered phase.

There is an extra region to obtain self-assembly with certain characteristics such as transparent solution, small size (10–20 nm), dependence of size on the DHPC fraction, rather ordered properties of surface (plotted at the second quadrant in the Cartesian diagram), and temperature-sensitive behavior of optical density. The gel phase is beneficial to form a planar bilayer; however, it is notable that the essential property of a bicelle should be magnetic orientation originating from its discoidal structure [5]. In the case of the bicelle or discoidal assembly in the TEM image [10], the self-assembly has two different patterns of shapes, such as face-on (sticky image) and edge-on (discoidal image). Together with the findings obtained in this study, a geometric consideration of the self-assembly, focusing on the molar ratio of DHPC and DMPC at Region (ii-1) (DMPC/DMPC = ~1:2–2:1), suggests that a bicelle-like structure could form. Figure 8a shows the TEM image of the DMPC/DHPC assembly with $X_{DHPC} = 0.5$ (total lipid concentration of 20 mM), revealing a non-spherical shape and rather a discoidal shape. As schematically shown in Figure 8b, since both shapes (face-on and edge-on) could be revealed, the DMPC/DHPC assembly that satisfies a small size (<30 nm) and "ordered" interfacial properties could be considered as a bicelle.



Figure 8. TEM images of DMPC/DHPC mixture; (**a**) X_{DHPC} = 0.4, 20 mM, and (**b**) the image of the bicelle-like structure.

4. Conclusions

We systematically characterized self-assemblies composed of DMPC/DHPC at total lipid concentrations less than 20 mM (<1 wt% lipid). The assemblies were categorized as Region (i), Region (ii-1), and Region (ii-2), in which the dominant state was large vesicle (>30 nm), gel-bicelle (<30 nm, "ordered" interfacial membrane properties), and micelle (<30 nm, "disordered" interfacial membrane properties), respectively. In particular, the bicelles observed in the DMPC/DHPC mixture systems exhibited both low membrane fluidity $(1/P_{DPH} < 6)$ and high membrane polarity ($GP_{340} > 0.3$). This reveals that the DMPC/DHPC bicelle structure could be observed only below 23 $^{\circ}$ C (*T* < *T*_m of DMPC) because the mean head group area of DMPC increases and the bilayer packing decreases in the disordered phase. The transparency of the suspension is a good indicator to roughly judge the size of the self-assembly [41]. In the conditions of fixed total lipid concentration, the self-assembly structure could be controlled using the DHPC fraction ratio: bicelle for $0.33 \le X_{DHPC} \le 0.67$ and micelle for $X_{\text{DHPC}} \leq 0.77$. In the conditions of fixed X_{DHPC} (especially at 0.33 < $X_{\text{DHPC}} < 0.4$), dilution leads to transformation from bicelle to vesicle, with significant increase in turbidity. The findings of this study are summarized in a diagram, which reveals the phase behavior of DMPC/DHPC assemblies at 20 °C. Bicelle assemblies can be used as lipid sources in supported lipid bilayer preparation [20]. Based on the phase equilibrium, DHPC molecules can be excluded from bicelle membranes by dilution. Utilizing this, bicelle assemblies can be applied for a continuous vesicle preparation process flow system.

Supplementary Materials: The following are available online at http://www.mdpi.com/2504-5377/2/4/73/s1, Figure S1: Temperature dependence on turbidity (OD_{500}) of 20 mM DMPC/DHPC self-assembly; $X_{DHPC} = 0.17$ (Region (i)), 0.4 (Region (ii-1)), and 0.95 (Region (ii-2)). This experiment was conducted in turn heating (20 °C (black) \rightarrow 30 °C (red)) to cooling (30 °C (red) \rightarrow 20 °C (blue)). Figure S2: Spectra of Laurdan in each 20 mM DMPC/DHPC membrane at 20 °C. (a) $X_{DHPC} = 0.17$ (Region (i)). (b) $X_{DHPC} = 0.4$ (Region (ii-1)). (c) $X_{DHPC} = 0.95$ (Region (ii-2)). The spectra were as follows: (d) $X_{DHPC} = 0.17$, (e) $X_{DHPC} = 0.4$ and (f) $X_{DHPC} = 0.95$ at 8 mM total lipid concentration.

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Abbreviations

DMPC	1,2-dimyristoyl-sn-glycero-3-phosphocholine		
DHPC	1,2-dihexanoyl-sn-glycero-3-phosphocholine		
Tm	phase transition temperature		
CMC	critical micelle concentration		
DLS	dynamic light scattering		
DPH	1,6-diphenyl-1,3,5-hexatriene		
Laurdan	6-dodecanoyl-N,N-dimethyl-2-naphthylamine		
GP	general polarization		
OD	optical density		

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