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# Methanol Accelerates DMPC Flip-Flop and Transfer: A SANS Study on Lipid Dynamics

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ABSTRACT Methanol is a common solubilizing agent used to study transmembrane proteins/peptides in biological and synthetic membranes. Using small angle neutron scattering and a strategic contrast-matching scheme, we show that methanol has a major impact on lipid dynamics. Under increasing methanol concentrations, isotopically distinct 1,2-dimyristoyl-sn-glycero-3phosphocholine large unilamellar vesicle populations exhibit increased mixing. Specifically, 1,2-dimyristoyl-sn-glycero-3phosphocholine transfer and flip-flop kinetics display linear and exponential rate enhancements, respectively. Ultimately, methanol is capable of influencing the structure-function relationship associated with bilayer composition (e.g., lipid asymmetry). The use of methanol as a carrier solvent, despite better simulating some biological conditions (e.g., antimicrobial attack), can help misconstrue lipid scrambling as the action of proteins or peptides, when in actuality it is a combination of solvent and biological agent. As bilayer compositional stability is crucial to cell survival and protein reconstitution, these results highlight the importance of methanol, and solvents in general, in biomembrane and proteolipid studies.

Lipid bilayers form the structural backbone of cellular membranes and possess marked lateral and transversal organization of lipids. This strict lipid organization has implications in vital cellular processes, including protein function and localization (1), vesicle fusion and budding (2), and apoptosis (3). Lipids undergo three types of spontaneous, diffusive motion: they 1) exchange between bilayers (interbilayer transfer/exchange), 2) translocate between bilayer leaflets (transverse lipid diffusion/flip-flop), and 3) laterally diffuse within the plane of the membrane surface. Herein, this study will focus on the former two because interbilayer exchange is linked to how lipids arrive, remain in, and leave cellular membranes, whereas lipid flip-flop disrupts the energy-driven maintenance of membrane asymmetry (i.e., the compositional difference between leaflets) in living cells. In essence, both dynamical actions are intrinsically linked to bilayers and their compositional stability. Previous studies have externally induced reorganization of lipids by outside factors as seen in model phospholipid membranes upon the addition of cations (4), detergents (5), and peptides

\*Correspondence: drew.marquardt@uwindsor.ca Editor: Tommy Nylander. https://doi.org/10.1016/j.bpj.2019.01.021 © 2019 Biophysical Society. (6–9). Here, we focus on how common organic solvents also impact these dynamics.

As membranes do not exist in isolation, they are in constant contact with compounds or solutions. Alcohols can be commonly found in the external environment of membranes. This proximity potentiates alcohol-membrane interactions, which can cause changes in membrane composition (10). For instance, methanol, the simplest alcohol, composed of a single methyl attached to a hydroxyl group, is ubiquitously utilized as a fuel source and chemical precursor in cells and in proteolipidic studies as an organic solvent of lipids and proteins/peptides. Such alcohols have been shown to alter lipid bilayer properties (11), making their study of great importance. Despite this importance, studies on lipid dynamics under the influence of alcohols are severely lacking. Several computational studies have examined the interactions between short-chain alcohols and lipid bilayers on a physical and molecular level (12,13) but none on lipid exchange and flip-flop in detail. To date, experimental data on these effects are also lacking, and the limited experimental reports have come into question because of the use of fluorescent probes (14,15). These probes can disrupt physicochemical bilayer properties and have also demonstrated unreliable flip-flop rates, differing depending on the type of fluorophore used, even within the same lipid system (16). We surmise this issue to extend



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to interbilayer exchange as well. To our knowledge, no such dynamical study has been conducted that examines methanol and model bilayers in a probe-free manner. This study overcomes these pitfalls by using small angle neutron scattering (SANS) as a noninvasive and probe-free technique to quantify lipid dynamics in the presence of methanol. SANS can temporally and spatially monitor the molecular organization within samples and has proven to be a powerful tool in allowing the simultaneous measurement of lipid flip-flop and exchange rates (17–20).

Here, we apply SANS to monitor lipid mixing of two distinct 1,2-dimyristoyl-sn-glycero-3-phosphocholine (DMPC) populations, one chain perdeuterated DMPC (d-DMPC) and the other fully protiated DMPC (h-DMPC), in the presence of an increasing deuterated methanol (d-methanol) concentration. This measurement is achieved by setting the ratio of H<sub>2</sub>O and D<sub>2</sub>O (here 45% D<sub>2</sub>O) such that the water solvent neutron scattering length density is matched to uniformly mixed d-DMPC/h-DMPC vesicles. Unmixed vesicles will thus display contrast versus the water solvent, resulting in heightened scattering intensity, whereas fully mixed samples will display scattering intensities akin to the solvent background (i.e., a flat and featureless curve). Thus, as h-DMPC and d-DMPC large unilamellar vesicles begin to mix via lipid monomers transferring within and between bilayers, the measured intensity will decay and eventually reach an intensity baseline, corresponding to a single population of completely mixed vesicles (shown in Fig. 2a). The experimental scheme can be seen in Fig. 1, whereas a more detailed protocol can be found in the Supporting Materials and Methods. A normalized intensity decay was calculated from the collective scattering curves of each sample and analyzed with a model for exchange/flip-flop by Nakano et al. (Fig. 2 b) (17). With this experimental setup, we are able to quantify both DMPC flip-flop  $(k_f)$ and exchange  $(k_e)$  rates under the influence of methanol.

Increasing methanol concentrations had a profound effect on the kinetics of DMPC monomers in free-floating large unilamellar vesicles. Despite differences in vesicle size and investigative techniques, our unperturbed DMPC flipflop and transfer rates are in excellent agreement with many values previously found (17,21-24). In a closely related study, Gerelli et al. used neutron reflectometry to measure DMPC flip-flop and exchange between vesicle dispersions and adsorbed planar bilayers (25). The exchange half-times for fluid-phase DMPC coincide with values found here (timescale of hours), whereas flip-flop was magnitudes faster ( $\leq 2.5$  min). As recently shown (16), the incomplete surface coverage of planar bilayers led to microscopic defects, which can facilitate lipid flip-flop and thus result in flip-flop rates on the order of seconds to minutes. This study, with fully sealed vesicles, bypasses such issues. More significantly, our results also reveal that methanol accelerates both fluid-phase DMPC flip-flop and transfer rates (Table 1). The flip-flop rate increases exponentially



FIGURE 1 Schematic of the contrast matching scheme used. Vesicles composed solely of d-DMPC (d54-DMPC) and h-DMPC are placed together in a  $H_2O/D_2O$  (55/45) mixture, contrast matched to an neutron scattering length density equal to fully mixed vesicles of d-DMPC and h-DMPC. Over time, because of lipid exchange and flip-flop, intensity loss can be monitored as vesicles mix and near the contrast match point.

(Fig. 2 c), whereas the exchange rate increases linearly under the studied concentrations. Our flip-flop finding is in line with Schwichtenhövel and co-workers, who found that radioactive and fluorescent lipid probes in human erythrocytes demonstrated exponential acceleration of inward flipping rates in the presence of 1-alkanols  $(C_2-C_8)$  and alkyl diols (14). Methanol has by far the weakest hydrophobic character in the short-chain alcohol group yet seems to perturb the membrane through the same or similar fashion as longer chained alcohols and alkyl diols. Although it has been shown that other short-chain alcohols affect inward flipping rates, we provide new, to our knowledge, insights with regards to both flip-flop and transfer rates in the presence of methanol. In general, at low methanol levels, DMPC undergoes slower flip-flop than transfer, but at concentrations above 2% (v/v) methanol, the situation is reversed. Interestingly, these observations suggest that methanol affects the two dynamical processes in distinct ways.

Flip-flop has a large energy barrier resulting from the transport of a polar, and often charged, lipid headgroup through the hydrophobic core. This unfavorable process can be augmented via intercalation of polar alcohol molecules within the membrane, which can cause short-lived transient pores and/or increase in the membrane's dielectric constant; both can justify the enhanced flip-flop observed here, but the latter is unlikely to be the major driving force as shown in past work (26). During lipid transfer, it is thermodynamically unfavorable for the hydrophobic tails to



FIGURE 2 (a) SANS curve of d-DMPC and h-DMPC vesicles with 3% (v/v) d-methanol solvent. Periodic measurements were conducted at  $37^{\circ}$ C over 38 h. (b) Normalized contrast decay curves of increasing d-methanol presence; continuous lines indicate fitted curves used to derive flip-flop and lipid exchange rate constants. Each data point represents the normalized integrated intensity of a single SANS curve similar to those found in (a). (c) Plot of measured flip-flop and lipid exchange rate constants as a function of d-methanol percentage concentration. Solid lines represent curves of best fit.

pass through both the polar headgroup region and aqueous phase. As exemplified by De Cuyper et al. (27), an increase in acyl chain length reduces the transfer of lipid monomers, presumably because of a greater exposed hydrophobic moiety. They also saw that incorporation of polyalcohols into these lipid structures caused an increase in transfer. In view of these results, alcohols can play a significant role in masking a lipid's hydrophobic character within a polar environment. Therefore, increasing methanol, an organic solvent that is fully miscible in water, should lower the energy barrier associated with lipid transfer.

Fluorescence techniques and freeze-fracture electron microscopy have demonstrated that short-chain alcohols promote vesicle hemifusion and complete fusion, hypothetically by way of outer leaflet disruption (28,29). This fusing of individual membrane vesicles provides sites of enhanced lipid exchange via fusion pores, which allows lateral diffusion of lipids to the adjacent leaflet, yielding faster than expected lipid dynamics (30). As a result, dynamic light scattering measurements of size and polydispersity were taken before and after incubation at multiple methanol concentrations, neither of which revealed significant changes, maintaining a vesicle diameter of  $\sim$ 140 nm and a polydispersity index of 0.15  $\pm$  0.02. Significantly, these results suggest fusion events did not occur as an increase in mean particle size and a polydispersity index would have been observed.

TABLE 1 DMPC Flip-Flop and Exchange Half-Times and Rate Constants

d-Methanol	Flip-Flop		Exchange	
(v/v%)	<i>t</i> <sub>1/2</sub> (min)	$k_f(\times 10^{-3} \text{ min}^{-1})$	<i>t</i> <sub>1/2</sub> (min)	$k_t (\times 10^{-3}{ m min}^{-1})$
0	87.2 ± 1.1	$8.0 \pm 0.1$	$62.8 \pm 0.3$	$11.0 \pm 0.05$
0.5	$78.1~\pm~2.6$	$8.9 \pm 0.3$	$54.8~\pm~0.4$	$12.6 \pm 0.1$
1.0	$58.3~\pm~1.4$	$11.9 \pm 0.3$	$53.0~\pm~0.4$	$13.1 \pm 0.09$
2.0	$47.8~\pm~1.3$	$14.5 \pm 0.4$	$44.3~\pm~0.3$	$15.6 \pm 0.1$
3.0	$16.7~\pm~1.3$	$41.6 \pm 3.3$	$40.8~\pm~0.2$	$17.0~\pm~0.1$

Further biophysical studies, using elastic SANS and small angle x-ray scattering (SAXS), were applied. SANS and SAXS are complementary techniques used to probe sample structure and are known to be extremely sensitive to membrane lamellarity and lipid bilayer structure. For example, multilamellarity can be signified by the appearance of Bragg peaks in the scattering data. The fact that both SANS and SAXS yielded curves that displayed diffuse scattering (i.e., no detectable sharp Bragg peak in Fig. S1) proves that multilamellar bilayers did not evolve under the presence of d-methanol. Collectively, these qualitative findings indicate that methanol did not alter the vesicles' morphological structure.

To determine if a defect-mediated mechanism can account for the rate enhancements, we examined pertinent bilayer properties. Previous simulation and experimental studies on model membranes revealed that the area per lipid  $(A_L)$ generally increases with increasing alcohol concentrations, irrespective of lipid saturation and chain length (12,31). The in silico study, in particular, found that this led to greater transient defects that provided lipid headgroups an opportunity to traverse the bilayer core and flip-flop (12). However, because of differences in lipid and alcohol concentrations between their systems and ours, we conducted our own structural analysis. A joint refinement of SANS and SAXS data was applied to robustly derive these structural features (32). As shown in Fig. S1, the scattering profiles of pure lipid and methanol-treated samples are indiscernible. In terms of relevant bilayer structural parameters, they are essentially unchanged (shown and explained in Fig. S1) and match well with previously reported DMPC data (33). These results do not outright disagree with previous studies. For example, Klacsová et al. observed changes in  $A_L$  and bilayer thickness  $(D_B)$  of a diunsaturated PC system hydrated with varying concentrations of aliphatic alcohols of different chain lengths,  $C_{8-18}$  (34); it was generally seen that shorter chain lengths imparted fewer effects on bilayer properties. With methanol having the smallest alkyl group, it makes sense that its effect on membrane structure is more difficult to detect. A plausible mechanism of methanol perturbation could be instead due to a decrease in chain order, often seen under increasing short-chain alcohol levels (35). However, an increase in  $A_L$  is also associated with such a change but, as seen here, the  $A_L$  remains static. Thus, the most likely explanation must involve methanol inducing short-range and perhaps short-lived defects, which are thus difficult to discern via methods that measure a global structural average, such as the biophysical techniques used in this study.

Traditionally, protein and peptide reconstitution have relied on either solvent addition or preincorporation with lipids before thin film hydration, the choice being dependent on the folding nature of the proteins/peptides. With the former, short-chain alcohols, including methanol, are popular carrier solvents used in studies on peptide activity (36,37), channel activity conductance (38), the critical structural motifs of proteins (39), and physical interactions between proteins and lipid bilayers (40). Our data suggest that even low concentrations of methanol, commonly used in these studies, have a profound effect on the compositional stability of membranes. In specialized cases involving lipid asymmetry, methanol-induced lipid scrambling can have unwanted consequences; lipid vesicles and likely other geometric setups exposed to methanol will have their asymmetric stability dramatically reduced manyfold. Such an effect limits the time of study and possible applicable techniques and assays. Because the response to methanol will vary depending on phospholipid composition, cholesterol content, and buffer of the studied membrane, ideal solvents that do not perturb membranes must be determined as well as the use of assays that can evaluate individual leaflet compositions and/or the degree of membrane asymmetry. These adaptations can help ensure experimental protocols have not altered the bilayer composition. Our findings further highlight an additional complication when adding proteins or peptides externally; for example, in studies on antimicrobial peptides (AMPs), though AMP attack is better simulated in this manner, these studies have the potential to incorrectly assign the cause of the enhanced lipid kinetics to the AMP, when in actuality it may be due to the carrier solvent or some combination of the two. Ultimately, this work highlights the importance of understanding the interplay between the system of interest and the carrier solvent on lipid mobility.

### SUPPORTING MATERIAL

Supporting Materials and Methods and two figures are available at http://www.biophysj.org/biophysj/supplemental/S0006-3495(19)30057-8.

#### **AUTHOR CONTRIBUTIONS**

M.H.L.N and D.M. designed the research. M.H.L.N, M.D., B.W.R., and D.M. carried out experimentation, and analyzed the data. E.G.K. and C.B.S. provided expert SANS help. M.H.L.N. wrote the article. M.H.L.N, M.D., B.W.R., C.B.S., E.G.K., and D.M. revised the manuscript.

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#### SUPPORTING CITATIONS

References (41-43) appear in the Supporting Material.

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# **Supplemental Information**

# Methanol Accelerates DMPC Flip-Flop and Transfer: A SANS Study on

# **Lipid Dynamics**

Michael H.L. Nguyen, Mitchell DiPasquale, Brett W. Rickeard, Christopher B. Stanley, Elizabeth G. Kelley, and Drew Marquardt

## **Materials & Methods**

1,2-dimyristoyl-sn-glycero-3-phosphocholine (14:0/14:0 PC, h-DMPC), 1,2-dimyristoyl-d54-sn-glycero-3-phosphocholine (14:0(d27)/14:0(d27), d-DMPC) 1,2-dimyristoyl-sn-glycero-3-phosphoglycerol (14:0/14:0 PG, DMPG) were purchased from Avanti Polar Lipids (Alabaster, AL). Lipids were received and used without further purification. Lipids in powdered form were dissolved in HPLC-grade chloroform prior to sample preparation. Deuterated methanol (d-methanol) was purchased from Cambridge Isotope Laboratories, Inc. (Andover, MA, USA).

# h-DMPC/d-DMPC Large Unilamellar Vesicle Sample Preparation

Exact quantities of lipids were mixed then placed under a constant stream of nitrogen and then dried in a vacuum oven at 50 °C to ensure organic solvent evaporation. Each sample included 5% DMPG to promote unilamellarity. The resulting lipid films were then hydrated in the pertinent D<sub>2</sub>O mixture solvent to produce samples of ca. 17 mg/mL. Subsequently, h-DMPC and d-DMPC samples were extruded separately through 100 nm pores in polycarbonate filters at 35 °C, above their phase transition temperature. Prior to any measurements, samples with d-methanol added were let stand for 1 hour to ensure equilibrium was reached before heating and measurements. Using a Wyatt DynaPro NanoStar, the mean particle diameter was determined to be ~140 nm, before methanol addition and after methanol incubation in small angle neutron scattering (SANS) banjo cells. Sizing measurements were taken at 30 °C.

# **Determining Bilayer Structure**



**Figure S1.** Small angle X-ray scattering (SAXS, open circles) and SANS (open circle inset) curves of pure h-DMPC LUVs in 100%  $D_2O$  (a) and in 3% d-methanol,  $D_2O$  solvent (b). Bold continuous lines represent fits using a 5-slab model which jointly analyzed both SANS and SAXS curves to robustly determine relevant bilayer parameters (2), as shown in the tables within the figure. All measurements shown here were conducted at 37 °C. Parameter uncertainty is estimated to be 2% according to Eicher et al. (2).



**Figure S2.** Electron density profiles (EDP) of DMPC in pure  $D_2O$  (black line) and 3% v/v dmethanol (teal dashes). EDPs were derived from the joint refinement of SANS and SAXS curves found in Fig. S1. The EDPs are essentially identical, highlighting the unchanged DMPC bilayer structure after d-methanol treatment.

# SANS and SAXS Structural Measurements

Elastic SANS curves were measured on the Extended Q-range SANS (EQ-SANS) instrument located in Oak Ridge National Laboratories (ORNL) (3). A sample-to-detector distance of 1.6 m was set, and a neutron beam with a wavelength band of 4 to 8 Å was used to produce a scattering vector, q, range of 0.02 Å<sup>-1</sup> to 0.8 Å<sup>-1</sup> where q represents the scattering vector, found by q =  $4\pi \sin\theta/\lambda$ , where 20 is the angle of scattered neutrons. The resulting scattered neutrons were detected via a 2D <sup>3</sup>He-based detector and radially averaged to produce a 1D I(q) scattering curve. Data processing was done using Mantid Software provided by ORNL where an absolute scale was established using a porous silica standard, as well as subsequent reductions accounting for sample transmission, pixel sensitivity, dark current and sample background. SANS curves can be found in Figure S1 as the red open circle plots in the foreground.

Complementary X-ray scattering data were measured using a Rigaku BioSAXS-2000 home source with a Pilatus 100 K detector. H-DMPC samples in pure  $D_2O$  or 3% d-methanol solutions were measured at a fixed sample-to-detector distance of 480 mm. As seen in Figure S1 (inset), resultant curves were averaged and subtracted via the relevant solvent backgrounds using ATSAS software (4).

In Fig. S1, several important bilayer parameters are shown:  $A_L$ ,  $D_B$ ,  $2D_C$ , and  $D_{HH}$ . AL represents the lateral area per lipid which is a good measure of lipid packing.  $D_B$  represents the bilayer (or Luzzati) thickness, as determined by the SANS portion of the fitting.  $2D_C$  is the hydrocarbon thickness.  $D_{HH}$  is another measure of bilayer thickness but is determined by the SAXS data and is representative of the distance between phosphate-phosphate groups. The parameter values for methanol-treated and untreated DMPC liposomes are almost unchanged, revealing that the statistical average of the vesicle structure is unchanged as well.

## Measuring Lipid Flip-Flop and Transfer Rates Using Small Angle Neutron Scattering

Dynamical lipid SANS measurements were conducted on the Very Small-Angle Neutron Scattering (VSANS) instrument located at the National Institute of Standards and Technology Center for Neutron Research (NIST-CNR). The white beam option was used on VSANS to maximize neutron count rates and minimize the required count times. A neutron wavelength ( $\lambda$ ) of 5.3 Å with a wavelength spread  $\Delta\lambda/\lambda$  of 40% was used with two detector carriages with sample-to-detector distances of 4 m and 19 m was used to accesses yielded a q-range of 0.003 Å<sup>-1</sup> to 0.12 Å<sup>-1</sup>. Data were collected using 3 min. acquisition times. Lipid samples were mixed and immediately measured on SANS using 1 mm or 2 mm path length quartz banjo cells at 37 °C. Each sample was run until the intensity decayed to a minimum and therefore reached the fully scrambled equilibrium. The total intensity was calculated using the Igor Pro reduction software and VSANS macros provided by the NIST-CNR by subtracting contributions from the external background, sample transmission, empty cell scattering as well as by correcting for detector pixel sensitivity. This resulted in well populated intensity vs. q curves after stitching.

Subsequent analysis follows the scheme used by Nakano et al. (1). Each sample possessed numerous intensity scattering curves, each representing a measurement taken at a single time point. All curves were converted into a single contrast decay curve per sample after normalization using  $\Delta\rho(t)/\Delta\rho(0) = (I(t)^{0.5}-I(\infty)^{0.5})/(I(0)^{0.5}-I(\infty)^{0.5})$ , where  $\Delta\rho(t)/\Delta\rho(0)$  represents the normalized total intensity decay, while I(t),  $I(\infty)$ , and I(0) are the scattering intensity at some time after mixing, scrambled infinity, and initial mixing time, respectively. Each sample was allowed to run until base-line, or full decay, was reached, signifying fully mixed vesicles and that our contrast match solvent was adequate. These final fully decayed curves provided an infinity point to use for normalization. The resultant decay curves were then fitted using

$$\frac{\Delta\rho(t)}{\Delta\rho(0)} = \left(0.5 - \frac{k_f}{X}\right)e^{\left(-\frac{k_e + 2k_f + X}{2}t\right)} + (0.5 + \frac{k_f}{X})e^{\left(-\frac{k_e + 2k_f - X}{2}t\right)}$$

where  $X = (4k_f^2 + k_e^2)^{0.5}$  to find DMPC flip-flop (k<sub>f</sub>) and transfer rates (k<sub>e</sub>), respectively. Half-times for these rates were calculated via  $t_{1/2} = (ln2)/k$ 

## Disclaimer

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# **Supporting References**

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