Impact of Nanoscale Hindrances on the Relationship between Lipid Packing and Diffusion in Model Membranes

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ABSTRACT: Membrane models have allowed for precise study of the plasma membrane’s biophysical properties, helping to unravel both structural and dynamic motifs within cell biology. Free-standing and supported bilayer systems are popular models to reconstitute membrane-related processes. Although it is well-known that each have their advantages and limitations, comprehensive comparison of their biophysical properties is still lacking. Here, we compare the diffusion and lipid packing in giant unilamellar vesicles, planar and spherical supported membranes, and cell-derived giant plasma membrane vesicles. We apply fluorescence correlation spectroscopy (FCS), spectral imaging, and super-resolution stimulated emission depletion FCS to study the diffusivity, lipid packing, and nanoscale architecture of these membrane systems, respectively. Our data show that lipid packing and diffusivity is tightly correlated in freestanding bilayers. However, nanoscale interactions in the supported bilayers cause deviation from this correlation. These data are essential to develop accurate theoretical models of the plasma membrane and will serve as a guideline for suitable model selection in future studies to reconstitute biological processes.

INTRODUCTION

The cascades for signal transduction usually begin at the cell surface, and for this reason the plasma membrane can be considered as the main hub for cellular signaling.1 However, drawing conclusions about membrane behavior and architecture proves challenging, not least because poorly understood or still unknown processes influence its dynamics.2,3 Our current knowledge shows that plasma membrane is a vastly complex and intricate system.4 Therefore, to truly appreciate and understand the finesse behind membrane dynamics, a “bottom-up” approach to discern different processes can prove useful.5 Several systems address this, employing a basic skeleton of only the essential biological components of the plasma membrane but engineered to allow systematic incorporation of complexity.6 Such reductionist systems can not only mimic membranes but also allow membrane-associated events to be systematically broken down to reveal their key contributing species owing to their controllable compositional complexity.7 Popular models include freestanding bilayers of synthetic lipids such as giant unilamellar vesicles (GUVs)8 or membrane blebs of live cells known as giant plasma membrane vesicles (GPMVs).5,10 However, the development of solid substrates to support bilayers has also shown promise, with two prominent constructs being the planar substrate/supported lipid bilayers (SLBs)11,12 and spherical bead supported lipid bilayers (BSLBs)13,14 (also termed spherical supported lipid bilayers, SSLBs).

It is certain that membrane models will continue to aid our understanding of the dynamics that underlie cellular signaling. Though, it is worth noting that each model brings its advantages and limitations, and caution should be employed while choosing appropriate model systems for given biological processes. It is, therefore, imperative to understand how each model influences bilayer behavior, not only to best select appropriate models for future research but to also avoid drawing misleading conclusions. This necessitates comprehensive comparison between models.

Here, we directly compare the biophysical properties of GUVs, SLBs, BSLBs, and cell-derived GPMVs. We apply fluorescence correlation spectroscopy (FCS),15 spectral imag-
ing, and super-resolution stimulated emission depletion (STED) spectroscopy to study the diffusivity, lipid packing, and nanoscale architecture of these membrane systems, respectively. We observed slower diffusion for SLBs and BSLBs compared to GUVs as reported in the literature.

While spectral analysis revealed no differences in lipid packing within these systems, STED combined with FCS showed nanoscale hindrances within SLBs and BSLBs that would explain their comparatively slower diffusion rates despite their similar lipid packing. Moreover, we showed that changes in lipid packing and diffusion in GUVs in response to compositional changes are predictable, while support has significant influence on this relationship. This work highlights the necessity of carefully comparing membrane models to progress research in membrane biology.

**METHODS**

**Cell Lines, Lipids, and Dyes.** Chinese hamster ovary (CHO) cells were cultured in DMEM/F12 (DMEM = Dulbecco’s Modified Eagle’s Medium) media supplemented with 10% fetal bovine serum (FBS) and 1% l-glutamine. Cells were prepared 2 d prior to experiments. Lipids stocks were obtained from Avanti Polar Lipids. GUV, SLB, and BSLB bilayers were prepared to contain 1% 1,2-dioleoyl-sn-glycero-3-[(N-(5-amo)-1-carboxypentyl)iminodiacetic acid] succinyl (DGS-Ni-NTA) with 1 mg/mL lipid stocks of 1,2-dioleoyl-sn-glycero-3-phosphocholine (DOPC), 1-palmitoyl-2-oleoyl-sn-glycero-3-phosphocholine (POPC), POPC:cholesterol (of varying concentrations), and 1,2-dipalmitoyl-sn-glycero-3-phosphocholine (DPPC):cholesterol (1:1), all in chloroform. Lipid stocks were stored under nitrogen at 4 °C.

**Generation of Model Membranes.** GUVs were prepared by electroformation. With this approach unilamellar vesicles between 10 and 100 μm diameters in size are produced. Lipid stock was spread onto two parallel platinum wires attached to a custom-built Teflon-coated chamber and left briefly to evaporate solvent. Wires were passed under nitrogen gas before submersion in 300 mM sucrose. Ten hertz AC current was applied to wires for 1 h to trigger vesicles swelling, followed by 2 Hz for 30 min.

GPMVs were prepared as described previously by Sezgin et al. Briefly, CHO cells were grown to 60% confluence, washed three times with GPMV buffer (10 mM 4-(2-hydroxyethyl)-1-piperazinethanesulonic acid (HEPES), 2 mM CaCl₂, 150 mM NaCl, pH 7.4), and then incubated at 37 °C in GPMV buffer with Parafomaldehyde (PFA) and dithiothreitol (DTT) (∼10 mM HEPES, 2 mM CaCl₂, 150 mM NaCl, 25 mM PFA, and 2 mM DTT) for 1–2 h to stimulate membrane blebbing. The supernatant containing GPMVs was then extracted.

SLBs were prepared by spin-coating. Briefly, glass coverslips of 0.25 mm and #1.5 thickness were first cleaned in piranha-solution (sulfuric acid (95–98%): hydrogen peroxide (30%), 3:1) for 1 h. Cleaned coverslips were then repeatedly washed and then stored in distilled water for no longer than one week. Twenty-five microliters of 1 mg/mL of lipid stock was pipetted onto the center of a dried coverslip and immediately spun for 30 s at 3500 rpm. The coated coverslip was then placed into a metal chamber and rehydrated with 1 mL of SLB buffer to form a bilayer (150 mM NaCl, 10 mM HEPES, pH 7.4).

BSLBs were prepared from spontaneous fusion of liposomes of lipid stock with 5 μm silica beads obtained from Bangs Laboratories. Liposomes were prepared by tip sonication. Lipid stock was placed under nitrogen gas to evaporate solvent completely leaving a dry, thin lipid film. Tris buffer saline solution (50 mM Tris HCl, 150 mM NaCl, pH 8.0) (500 μL) was added to lipid residue as liposome buffer. Lipid solution was then transferred to ice and sonicated at 55 Amp for 15 min, with 10 s pulse periods separated by 10 s rest intervals. Silica beads were washed with 1 mL of phosphate-buffered solution (PBS) before centrifuging for 30 s at 2000 rpm. Supernatant was removed with residual left to prevent beads from drying, and washing was repeated twice. Beads were mixed with liposomes (1:5) and then shaken for 20 min at 1200 rpm to form BSLBs. After they were centrifuged at 2000 rpm for 30 s, BSLBs were then washed twice with PBS, with ~500 μL solution reserved from the final wash.

**Confocal Imaging.** Membranes were imaged with Zeiss LSM 780 or 880 microscopes. For labeling, 20–100 ng/mL (final concentration) AbiSTR-PE was added to GUVs and GPMVs and incubated for 15 min. For SLBs and BSLBs, fluorescent lipid analogue was added to the lipid mixture with the ratio of 1:2000 (labeled lipid to total lipid ratio). With this way, the fluorescent probe localizes to the outer leaflet of GUVs but both leaflets of SLBs and BSLBs, and therefore the diffusion of the probe represents both leaflets for all the model systems (as our GUVs are symmetric). GUVs, SLBs, and GPMVs were transferred to Ibidi 8-well plastic chambers of #1.5 thickness. Wells were previously treated with 1 mg/mL bovine serum albumin (BSA), left for 1 h, and then washed three times with PBS or GPMV buffer before transfer. GUVs and BSLBs were then suspended in PBS. GPMVs were left unsuspended for 1 h to allow for settling of imaging. Laser (633 nm) was focused onto bilayers by 40X water immersion objective (NA 1.2) for excitation of AbiSTR-PE fluorophore.

**Confocal and STED Fluorescence Correlation Spectroscopy (FCS).** FCS was used to measure and compare the diffusion of AbiSTR-PE through GUV, SLB, and BSLB models prepared with POPC, POPC:cholesterol (of varying concentrations), and DPPC:cholesterol compositions. Models were incubated with 0.05 μg/mL AbiSTR-PE as previously described. GUVs, SLBs, and GPMVs were measured in Ibidi glass chambers of #1.5 thickness prepared as before. SLBs were measured on #25 mm and #1.5 thickness glass cover slides.

Confocal FCS was performed on Zeiss LSM 780 microscope with 40X NA 1.2 water immersion objective. Before measurement, focal spots were calibrated using a mixture of 10 nM Alexa 488 and 647. FCS measurements were recorded with a 633 nm laser at 0.1% power (~2 μW). Laser focusing was completed by finding axial positions of maximum fluorescence intensity at the bilayer. Correlation curves were obtained over 5 s periods with five repeats per area studied. Curves were then fitted with the freely available FoCuS-point software to extract diffusion coefficients. All FCS data were fitted with a two-dimensional diffusion model that incorporated an initial triplet state that describes a fixed 5 μs relaxation period. STED-FCS was performed with Leica SP8 microscope using 100X NA 1.4 oil immersion objective. All FCS measurements were done at the bottom membrane of GUVs, SLBs, GPMVs, and BSLBs to avoid refractive index mismatch. Laser focusing and data acquisition were performed as previously described.
Spectral Imaging. Spectral imaging was used to measure and compare packing within bilayers. GUVs, BSLBs, SLBs, and GPMVs were incubated with 1 μM C-Laurdan for 10 min. GUVs, BSLBs, and GPMVs were then transferred to Ibidi plastic bottom chambers prepared as previously described. Imaging was performed by Zeiss LSM 780 microscope equipped with a 32-channel GaAsP detector array and a polarizer that minimizes photoselection effect. Laser light at 405 nm was selected for C-Laurdan excitation, and the λ-detection range was set between 415 and 691 nm. Images were analyzed with the custom generalized polarization (GP) plugin of FIJI software as described previously.16 GP was calculated with the formula:

\[ GP = \frac{I_{440} - I_{490}}{I_{440} + I_{490}} \]

RESULTS AND DISCUSSION

There is already substantial evidence of a support’s influence on bilayer diffusion.25–28 To further assess the influence of support on other pivotal biophysical parameters and the relationships between them, we selected four models that broadly represent the spectrum of designs extensively utilized. Our chosen models span from freestanding GUVs to supported planar SLB and spherical BSLB constructs and to the more cellular inspired bilayer model of GPMVs (Figure 1).

GUV Models Confirm a Relationship between Lipid Packing and Diffusivity within the Bilayer. We first considered the freestanding bilayer model of GUVs. GUVs are by definition free from a support influence and thus can reveal unbiased relationships between biophysical parameters of the bilayer. Here, we systematically altered the composition of GUVs in the form of glycerophospholipid species and cholesterol concentration with the intent of incrementally increasing bilayer ordering and seeing how this correlates with diffusion. C-Laurdan was incorporated into bilayers to report on lipid ordering. Spectral imaging confirmed increased ordering as compositions progressed from comprising fully unsaturated lipid (DOPC, 18:1/18:1) to monounsaturated (POPC, 16:0/18:1) and to saturated lipid (DPPC, 16:0/16:0) (Figure 2A) and increased too alongside cholesterol concentration, as expected.32,33 Extracting GP scores that quantify ordering within images revealed a monotonic increase in ordering for GUVs comprised with higher concentrations of...
cholesterol (Figure 2B, violin plots). We then gauged for the influence of lipid composition on bilayer diffusivity. To this end, we employed point FCS to obtain the lateral diffusion coefficients of Abberior Star Red-labeled phosphatidylethanolamine analogue incorporated within GUVs of the same compositions (Figure 2C). Diffusion measurements for GUVs agree with earlier reports for other similarly structured fluorophores (Figure 2B, box-and-whisker plots).\(^{34}\) Diffusion coefficients decreased with lipid saturation and also as a function of cholesterol concentration, agreeing with previous studies, and describe an approximately linear correlation.\(^{35}\) Overlaying respective diffusion coefficients and GP scores revealed a negative correlation between the two measures and so suggests a distinct relationship between bilayer ordering and diffusivity in GUV models (Figure 2B).

**Diffusion in GUVs Differ from Supported Membrane Models.** Having observed a clear relationship between lipid ordering and mobility in GUVs, we sought to investigate whether this relationship extended to other models. As discussed, we selected models that together broadly capture current key design motifs; herein we investigated aforementioned GUVs alongside SLBs and BSLBs of supported planar and spherical designs, respectively, as well as freestanding GPMVs that reflect the complex composition of native bilayers. First, we established diffusion profiles of all models in identical conditions for comparison. FCS was performed in models, for the interest of simplicity, composed of single-component POPC. Confocal images confirmed homogeneous fluorescence signal throughout respective bilayers at the microscopic level (Figure 1). FCS curves for all models fitted well to a one-component two-dimensional diffusion model (Figure 3A). Expectedly, diffusion coefficients measured highest in GUVs, which demonstrated an approximate threefold increase in mean diffusion coefficient over SLBs and GPMVs, with an approximate fivefold increase over BSLBs (Figure 3B). Our results generally agree with previous studies that separately demonstrated an increase in diffusion speeds in GUVs compared with in SLBs\(^ {18}\) and GPMVs.\(^ {36,37}\)

**Lipid Packing and Diffusion Do Not Correlate across Different Supported Models.** Having revealed the trends in diffusivity across models, we then set to investigate their lipid packing and assess for a correlation as observed previously in GUVs. Spectral imaging expectedly reported homogeneous packing within all four models, that is, no notable microscopic inhomogeneity (Figure 3C). Although statistically significant, there was, on the one hand, an extremely small difference in packing between BSLBs, SLBs, and GUVs (Figure 3D). GPMVs, on the other hand, showed higher GP values compared to all model systems (Figure 3D), presumably due to their complex lipid composition, which includes saturated lipid components and cholesterol. When we overlaid GP data with diffusion coefficients, we observed no correlation between the two, suggesting the slow diffusion trends reported within supported models is not dictated by lipid packing (Figure 3E).

**STED-FCS Reveals Nanoscale Hindrances in the Architecture of Supported Models.** Until this point, diffusion measurements were performed on the diffraction-limited resolution scale, which cannot report on nanoscale dynamics. In contrast, super-resolution STED combined with FCS (STED-FCS) offers resolution down to 20–40 nm (Figure 4A), with which nanoscale dynamics can be addressed.\(^ {17,28}\) In this regard, STED can glean assessment of bilayer dynamics in the context of surface architecture, an appreciated consideration in supported model design.\(^ {27}\) For
instance, unwanted aggregates or surface defects could hinder diffusion of the fluorescent species as they travel through the observation spot and thus yield slow diffusion that is independent of the microscopic viscosity or lipid packing of the sample. Measuring the diffusion in a reduced focal volume with an STED laser allows us to distinguish free Brownian diffusion from hindered diffusion (Figure 4A). A simple measurement yielding the ratio of confocal ($\tau_C$) and STED ($\tau_S$) transit times through the focal spot ($\frac{\tau_S}{\tau_C}$) will reveal any differences (and hence unexpected hindrances) within the models. Nanoscale hindrances lead to higher $\tau_S$ values and thus to smaller $\frac{\tau_S}{\tau_C}$ ratios.

We performed confocal FCS on model membranes composed of single-component POPC as before, immediately followed by STED-FCS. The ratio of confocal and STED transit times ($\frac{\tau}{\tau_C}$) was calculated for all models (Figure 4B). GUVs demonstrated a $\frac{\tau}{\tau_C}$ value of 19.7 ± 3.7, while SLBs and BSLBs showed 15.2 ± 1.4 and 13.1 ± 2.3, respectively. This suggests nearly free diffusion, that is, minimal hindrance on lipid mobility in GUVs but hindrances in SLBs and BSLBs at the nanoscale resulting in higher $\tau_S$ values.

The Relationship between Lipid Ordering and Diffusion Is Less Pronounced in Supported Models. We showed that lipid packing is comparable in bilayers of our single-component POPC models (Figure 3D), but supported models exhibit altered diffusion profiles and hint toward a “broken” relationship between ordering and diffusion in such models (Figure 3E). We sought to demonstrate how this discrepancy is affected in more ordered membrane systems by incrementally increasing ordering in GUVs, SLBs, and BSLBs. We then measured GP and diffusion and overlaid them. GP analysis of spectral images confirmed an increase in ordering as bilayer compositions changed from POPC to POPC:chol (1:1), and DPPC:chol (1:1) (Figure 5A−C). Confocal FCS measurements for each model also showed an expected trend for all models, namely, slower diffusion for more saturated membranes. (Figure 5A−C).

To quantitatively assess how these two parameters are correlated and to have a more comprehensive picture of the changes in diffusion and GP at different compositions, we calculated the “fold change” in diffusion (mean diffusion coefficients in POPC divided by mean diffusion coefficient in POPC:chol or DPPC:chol). Similarly, we calculated unit change in GP (mean GP in POPC:chol or DPPC:chol minus mean GP in POPC) for all model systems (Figure 5D,E). As expected, more ordered membrane systems yielded slower diffusion, and this trend was maintained in all systems (Figure 5D,E) but, importantly, to different extents. In GUVs, the differences in diffusion as well as GP (between ordered and disordered membrane systems) is the highest. This suggests that the freestanding GUV system is very sensitive to

Figure 4. STED-FCS to reveal nanoscale hindrances in model systems. (A) Principle of STED-FCS. A super-resolved observation volume can be produced by a depletion beam (blue ring) designed with a zero-intensity center that effectively cancels surrounding emission signal from excited fluorophores. (B) Ratio of calculated confocal ($\tau_C$) and STED ($\tau_S$) transit times in model membranes composed of single-component POPC. A lower ratio of confocal/STED transit times suggests nanoscale hindrances.

Figure 5. Relationship between lipid ordering and diffusion across lipid compositions. Direct comparison of GP (violin plots) and diffusion (box-and-whisker plots) within (A) GUV, (B) SLB, and (C) BSLB models of POPC, POPC:chol, and DPPC:chol. Dotted lines indicate trends in GP, while solid lines indicate trends in diffusion. (D) Calculated fold change in mean diffusion coefficient across compositions within models. (E) Calculated unit change in mean GP value across compositions within models.
compositional changes. However, SLBs and BSLBs did not react as well to the changes in saturation. For instance, the ΔDiffusion value for DPPC:chol (i.e., mean diffusion coefficient of fluorescent lipid in POPC divided by mean diffusion coefficient of fluorescent lipid in DPPC:chol) was ~6.5 for GUVs, while it was ~4 and ~3.4 for SLBs and BSLBs, respectively. In other words, diffusion in DPPC:chol GUVs is 6.5 times slower compared to POPC GUVs; however, diffusion in DPPC:chol SLBs is only 4 times slower compared to POPC SLBs (Figure 5D).

Similarly, ΔGP for DPPC:chol (i.e., mean GP in DPPC:chol minus mean GP in POPC) is ~0.65 units for GUVs and ~0.44 and ~0.57 for SLBs and BSLBs, respectively. In other words, GP in DPPC:chol GUVs is 0.65 GP units higher compared to POPC GUVs; however, GP in DPPC:chol SLBs is only 0.44 units higher compared to POPC SLBs (Figure SE). These data highlight a tight and near-linear relationship between ordering and diffusion in GUVs but not in supported membranes, which exhibit decreased sensitivity to compositional changes and deviation from the tight relationship between ordering and diffusion.

**CONCLUSIONS**

There is growing interest in reconstituting membrane-associated processes in vitro using model membrane systems given their proven potential to refine mechanisms and glean new hypotheses. However, few studies have ventured a comprehensive comparison of their biophysical properties as a function of the inherent parameters (a support presence, its material, model geometry, etc.). Indeed, a few studies have highlighted this concern by demonstrating altered protein functionality in different membrane models. We believe our comprehensive comparison of lipid diffusion and ordering between GUVs, planar SLBs, spherical BSLBs, and GPMVs can be a guideline for future studies that can use similar approaches for different biophysical properties.

Employing super-resolution STED-FCS, we confirmed the presence of nanoscale hindrances that seemingly influence the relationship between lipid ordering and diffusion in supported models, contrasting the linear relationship we observed in freestanding GUVs. Lipid ordering was comparable between GUVs, SLBs, and BSLBs of equal composition—at least at the microscopic scale, suggesting a more direct influence of support on diffusion. Indeed, the drag imparted by a lipid support is widely appreciated. However, our application of STED-FCS adds a spatial-temporal framework for these interactions and suggests repeated but brief interactions between support and constituting lipids. This importantly highlights the necessity of applying super-resolution techniques to better elucidate the bilayer structures and local physicochemical properties. Nanoscale surface perturbations due to the support material have also been previously appreciated to influence diffusion. Therefore, it is possible that nanoscale changes in surface topography of supported models effectively “slow” lateral diffusion. Our observed discrepancy with BSLBs could reflect a change of support material, in which application of other methodologies such as atomic force microscopy could prove most useful.

The influence that bilayer ordering poses over lateral mobility can essentially be explained by increased van der Waals efficiency between lipids. However, absent from previous diffusion studies is direct comparison with lipid ordering we employ here. When combined with the capacity of the GUVs to retain unhindered lipid dynamics, we were able to demonstrate a direct correlation between lipid ordering and diffusion. Indeed, this is an attractive property of GUVs; due to their lack of support, GUV diffusion is considered “free” or, more importantly, predictable. The potential of model membranes is impressive given their inherent skeletal framework, and indeed current applications prove this. Yet, our results highlight an unpredictability in model behavior that can accompany the incorporation of design motifs, in this case the presence of a support, and should not be overlooked in sensitive analysis.

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**Notes**

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