Brownian dynamics of electrostatically adhering small vesicles on a membrane surface induces domains and probes viscosity

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Abstract

Using single particle tracking, we investigate the interaction of small unilamellar vesicles (SUVs) that are electrostatically tethered to the freestanding membrane of a giant unilamellar vesicle (GUV). We find that the surface mobility of the GUV-riding SUVs is Brownian, insensitive to the bulk viscosity, vesicle size, and vesicle fluidity, but it is strongly altered by the viscosity of the underlying membrane. Analyzing the diffusional behavior of SUVs within the Saffman-Delbrück model for the dynamics of membrane inclusions, supports the notion that the mobility of the small vesicles is coupled to that of dynamically induced lipid clusters within the target GUV membrane. The reversible binding also offers a non-perturbative means for measuring the viscosity of bio-membranes, which is an important parameter in cell physiology and function.
Introduction

A single lipid membrane – a flexible, quasi-two-dimensional biomolecular surface composed of two apposing layers of amphiphilic lipids\(^1\)–fluctuates freely in water.\(^2\) As two membranes come into close proximity, their thermally-excited, out-of-plane fluctuations become suppressed giving rise to a long-range repulsive force that tends to drive the membranes apart.\(^3\) Overcoming this entropic repulsion – such as through biospecific ligand-receptor binding or nonspecific electrostatic attraction between oppositely charged membranes – can bring membranes together creating distinct adhesion configurations or intermembrane junctions.\(^4\) Such junctions are not uncommon in biology. They represent an integral part of intercellular signaling strategies used by multicellular organisms,\(^5\) among which perhaps the best known example is that of immunological synapse. Here, single T-cells come into a junction with single antigen-presenting cells characterized by a molecular pattern of adhesive bonds consisting of central clusters of T-cell receptors surrounded by a ring of adhesion molecules.\(^6\)

Beyond suppression of thermal undulations, the appearance of intermembrane adhesive states introduce additional physical perturbations to the interacting membranes\(^1\) with important ramifications. Previous efforts, focused dominantly on the adhesion of membranes of comparable dimensions, document processes of adhesion-induced changes in membrane physical properties. A variety of features – including adhesion-induced changes in membrane tension, lateral fluidity, spatial distributions of membrane molecules\(^2\) and chemically-differentiated domains\(^8\) within single membranes as well as exchange of lipids between interacting membranes\(^9\) – have all been reported. Between membranes interacting through weak electrostatic forces, a notable example is the emergence of adhesion gradient through molecular
redistributions of charged amphiphiles. Here, the induced charge gradient acts to propel the vesicle along the gradient reconstituting the so-called haptotaxis observed in living cells.\textsuperscript{10}

In this same vein, studies aimed purportedly at characterizing intermembrane junctions involving membranes of vastly different dimensions are much more limited. In living cells, such junctions appear transiently as pre-fusion or post-division docking states during endo- and exocytosis, morphogenesis of transporter vesicles, as well as during viral budding and egress.\textsuperscript{11}

At the morphological level, they are perhaps best represented by closely apposed nanoscopic, colloidal (~50-100 nm in dia.) small unilamellar vesicles (SUVs) and a target, 10-50 \( \mu \)m diameter giant unilamellar vesicle (GUV). In this situation, a population of small vesicles adhering to and surfing on (in the case of weakly adhering membranes) a membrane produces a dynamic array of intermembrane junctions creating local and mobile “hotspots.” Similar localized hotspots, created through hybridization of single DNA tethers between lipids of SUVs and a solid supported membranes, have been previously shown to significantly impact diffusion.\textsuperscript{12} They reduce the lateral diffusivities of the surfing SUVs and thus the DNA-lipids in supported membranes by three- to five-fold, which was speculated to reflect changes in local environment of tethering DNA lipids, which effectively increases the size of the diffusing component in the supported bilayer. This multiple tethering effect can be expected to be much more pronounced when SUVs adhere onto GUV membranes electrostatically. In that situation a large adhesion zone, determined by the size of the SUVs, can be expected to cluster a number of underlying lipids in the GUV membrane affecting both the SUV mobilities and splitting the lipids in the GUV between two sub-populations: free lipids moving individually and clusters or discs of lipids, whose concerted mobilities determine the translational diffusivities of the riding SUVs.
In the work reported here, we track the motion of individual electrostatically adhering SUVs riding onto the GUV surface, which serve as a probe to measure the viscosity of the underlying membrane. This membrane viscosity has been proven difficult to measure, and various techniques have been developed in the literature. An interesting recent technique uses a relation, which is calibrated in bulk, between the viscosity and the emission of fluorescently labeled molecules incorporated into the membrane. Another novel approach measures the shear-induced, large-scale circulation within the membrane of a GUV attached to the wall of a flow chamber. Most methods however rely on measuring the diffusivity of tracer particles, embedded in or adhering to the membrane, and invoking a fluid mechanics model to translate the diffusivity to viscosity. For instance the mobility of membrane lipids or membrane proteins have been measured using fluorescence recovery after photo-bleaching, or fluorescence correlation spectroscopy. Translating diffusivity to viscosity however, using fluid mechanics models, generally works better for larger particles, than individual molecules. Therefore, in an attempt to accurately determine the viscosity of lipid bilayers, single particle tracking has been used to measure the diffusivity of large membrane inclusions, such as phase-separated lipid domains or peripherally bound tracer particles, using covalent bonds. This latter approach is complicated by the unknown effective size of the diffusing objects, owing to an uncontrollable number of bonds and a possible deformation of the underlying membrane. Here we circumvent these problems, by using a weak electrostatic force to bind small vesicles to the membrane. We will show below, that the vesicles associate to a cluster of membrane lipids, and that the radius of the diffusing cluster correlates well with the electrostatic adhesion zone, permitting a straightforward determination of the membrane viscosity.
Figure 1. (a) A 3D reconstruction from a stack of confocal cross sections of a GUV consisting of binary lipid mixture composed of DOPC and DOEPC (9:1) covered with SUVs (radius~ 60nm) composed of DOPC and DOPS (95:5) and doped with Rh-PE (1 mol%). (b) Schematic representation of electrostatically adhering small vesicles on the outer surface of a giant vesicle. The weak electrostatic tethering reversibly divides the membrane into two distinct sub-populations, i.e. free lipids and lipids in clusters (red regions) that are bound by the SUVs.

**Materials and Methods**

**GUV preparation.** Giant unilamellar vesicles (GUVs) were prepared using the electroformation method.\textsuperscript{23, 24} Briefly, stock solutions of lipid mixtures (mol:mol) were prepared at 1 mg/ml in chloroform (all lipids were purchased from Avanti Polar Lipids). 20 µL of the stock solution were spread onto the conductive side of ITO-coated slides within an area delimited by a O-ring and allowed to dry in vacuum for at least 1 h. Electroformation was performed with a 300 mM sucrose solution by using a commercial Vesicle Prep Pro (Nanion, Munich, Germany). Specifically, DOPC (1,2-dioleoyl-sn-glycero-3-phosphocholine):DOEPC [1,2-dioleoyl-sn-glycero-3-ethylphosphocholine (chloride salt)] (9:1), DOPC:cholesterol:DOEPC (5:4:1), and DMPC (1,2-dimyristoyl-sn-glycero-3-phosphocholine):DOEPC (9:1) GUVs were electroformed by applying an AC current at 500 Hz, 3 V and 45°C (above the gel–fluid transition temperatures of the lipid mixtures) for 120 min. Then the GUVs were diluted in a 300 mM glucose solution. Experiments involving DOPC:DOEPC GUVs were also performed in a more viscous solution.
For this purpose 25 % (v/v) glycerol was added to the external bath, in which the GUVs were
diluted. Upon glycerol addition, the GUV underwent shrinkage due to an osmotic imbalance.
However since the permeability of the membrane to glycerol is relatively high ($\sim 2 \times 10^{-6}$ cm/s),
the GUV regained its initial spherical shape a few min later, after yielding iso-osmotic
conditions. Therefore, SUVs were added 30 min after addition of glycerol to ensure complete
equilibrium.

**SUV preparation.** Small unilamellar vesicles (SUVs) were made by the extrusion method.
Briefly, a mixture of DOPC, and DOPS [1,2-dioleoyl-sn-glycero-3-phospho-L-serine (sodium
salt)] (5 mol%) in chloroform was prepared at a total lipid concentration a solution of 5 mg/ml.
1 wt % rhodamine-DOPE (1,2-dioleoyl-sn-glycero-3-phosphoethanolamine-N-(lissamine
rhodamine B sulfonyl) in chloroform was also included (all lipids were purchased from Avanti
Polar Lipids). The lipid solution was first dried using a flow of nitrogen. The dried lipid film was
stored in vacuum for 3 h, after which it was rehydrated with buffer (Tris 10 mM, NaCl 150 mM
and pH 7.5). After vortex mixing of the solution of hydrated lipids, unilamellar vesicles were
made with an Avanti Mini-Extruder (Avanti Polar Lipids) using a polycarbonate membrane (100
nm pore size, Avanti Polar Lipids). The vesicle size distribution was measured by the Nano-
Sight particle tracking technique (NanoSight, U.K.).

**GUV and SUV mixing.** After electroformation, 10 µl of the GUV solution was mixed with 200
µl of a 300 mM glucose solution in the µ-Plate 96-well (Ibidi, GmbH, Germany) for microscopy
imaging. SUVs (5 mg/ml) were pre-diluted to 0.1 mg/ml in buffer (Tris 10 mM and NaCl 150
mM, pH 7.5) and 4 µL of this solution was added to the 200 µl GUV solution in the µ-Plate 96-
well, such that the SUVs sparsely decorate the GUVs. The pH and the ionic strength (NaCl) in
the final solution were about 7.5 and 4 mM, respectively.

**Fluorescence microscopy.** Spinning disk confocal microscopy measurements were carried out
using an inverted Eclipse TE 2000 microscope (Nikon) fitted with X-Light spinning-disk
confocal unit (CrestOptics, Rome, Italy) and an Andor iXon+ EMCCD camera (Andor
Technology, Belfast, Northern Ireland). For measurements, we used a 60× oil immersion
objective (NA 1.49), Rhod-DOPE (Ex/Em; 560/583) was exposed with a 50 mW 561 laser line.
Time-lapse images were acquired at 50 frames/sec. At least 5 GUVs were imaged for each
experiment.

**Results and Discussion**

Our experimental design involves the creation of a pericentric SUV-GUV interface stabilized by
electrostatics (Figure 1a). We used negatively charged SUVs (~120 nm dia.) consisting of 95
mol% zwitterionic DOPC doped with 5 mol% negatively charged DOPS and positively charged
GUVs, consisting of 90 mol% zwitterionic DOPC doped with 10 mol% positively charged
DOEPC. At these low charge densities, the SUV-GUV interface is stable showing little tendency
for intermembrane fusion, consistent with previous studies, which have shown that the lower
charge density threshold for fusion is roughly 10 mol%. Moreover, to minimize the
complicating effects of any molecular exchange and creation of adhesion gradients at the SUV-
GUV interface, we limit our measurements of SUV diffusion to the first several minutes after
SUV adsorption, during which molecular exchanges are known to be minimal. During these
experimental time scales, we do not observe noticeable changes in the number of adhering
vesicles, which confirms the irrelevance of intermembrane lipid mixing and/or fusion.
The diffusivity of vesicles was measured using a particle tracking method.\textsuperscript{22} Previously, this approach has been used successfully to characterize the motion of nanoparticles,\textsuperscript{30, 31} peptides\textsuperscript{32} and viruses\textsuperscript{33} at the membrane interface. To trace the 2D diffusive motions of vesicles, the equatorial plane of a GUV was imaged using confocal microscopy (Figure 2a). Using in-house image analysis software (developed in MATLAB), the GUV rim (dashed circle in Figure 2a) and the SUVs (marked by circles in Figure 2a) were detected and vesicle trajectories along the equatorial GUV rim were constructed from a sequence of typically $10^4$ images. After a vesicle moves into the confocal plane, its motion can be tracked only for a short while (typically one second), before it randomly moves out of the plane. Typically $\sim100$ SUVs reside on the GUV surface (diameter $\sim20\ \mu$m), which is equivalent to an average distance between the SUVs of $\sim4$ $\mu$m. Since this distance is much larger than the SUV diameter ($\sim100$ nm) and the Debye length ($\sim1$ nm), steric and electrostatic interactions between the SUVs are negligible. It is noted that for illustration purposes, the GUV in Figure 1a had a relatively large SUV coverage, while the SUV coverage in the diffusivity experiments was much lower.

Figure 2b shows the detected positions of the SUVs, expressed using the angle ($\theta$ in Figure 2a) along the equatorial GUV rim, as functions of time $t$. Since the SUVs continuously move in and out of the confocal plane, the trajectories in Figure 2a appear fragmented. Analysis of the mean square displacement (MSD) of the trajectories confirmed the Brownian character of the diffusion of the surface bound vesicles (Figure 2c). Figure 2d presents a histogram of the diffusion coefficients for the surface-bound vesicles (red bars) as well as for vesicles freely floating in the bulk. The latter is obtained from vesicle tracking in the bulk using nano-particle tracking analysis NTA; see Supporting Information Figure S2. The large spread in the diffusivity has many possible causes, e.g., measurement noise, distribution of vesicle sizes and charge densities, and
statistical fluctuations due to the stochastic nature of the diffusional motion and the relatively short time that this motion can be sampled, before the SUV moves out of the focal plane. We observe that the logarithm of the diffusivity is normally distributed (dashed lines), which means that the diffusivity itself is log-normally distributed.

The diffusion coefficient $D$ for surface bound vesicles is determined as $1.3 \pm 1.0 \, \mu m^2 s^{-1}$. With a radius $a = 57 \pm 18 \, nm$, which was measured using NTA, (See Supporting Information, Figure S2), the diffusivity of the freely floating vesicle equals $3.6 \pm 1.1 \, \mu m^2 s^{-1}$, which is a three times as large as that for the adhering SUVs. The smaller diffusivity for the adhering SUVs reflects a larger friction. To characterize the nature of this friction, we conduct a series of control experiments, where we study the influence of various membrane and solvent properties on SUV diffusivity on the GUV surface. The reader is referred to Table S1 and Figure S1 in the Supporting Information, which summarizes the SUV diffusivity and the displacement statistics for these control experiments.
Figure 2. Mobility analysis of single small unilamellar vesicles (SUVs) diffusing on the surface of a giant unilamellar vesicle (GUV). (a) Image processing including detection of the GUV edge at the equatorial cross-section (indicated by the dashed circle) and bound SUVs which appeared as bright spots on the GUV rim (marked by circles). An angle $\theta$ was assigned to each vesicle position at each frame. (b) Detected trajectories of bound SUVs on (angle, time) - plane at a temporal resolution of 20 ms. The inset shows a vesicle trajectory spanning about two seconds or hundred frames. (c) Displacement $x$ probability $p$ of surface-bound vesicles, as a function of the normalized displacement $x/(4Dt)^{1/2}$, where $D$ is the diffusivity and $t$ is time. The dashed line is a Guassian function. The inset shows that the mean square displacement (MSD) increases linearly with time, which indicates that the motion is Brownian. (d) Histograms of the diffusion coefficient $D$ for surface-bound vesicles (red bars) and freely floating vesicles (blue bars). Note the logarithmic $x$-axis.

We begin by analyzing the effect of the GUV membrane fluidity on the SUV diffusivity. For this purpose, we measure SUV diffusivity on GUVs consisting of a mixture of 50 mol% DOPC, 10 mol% DOEPC and 40 mol% cholesterol. Upon adding cholesterol to the underlying membrane, the SUV diffusivity is observed to decrease by a factor of two from $1.3 \pm 1.0 \, \mu m^2/s$ to $0.6 \pm 0.4 \, \mu m^2/s$ (Figure 3a). This relative decrease is of similar magnitude as observed upon adding 40
mol% cholesterol to an egg phosphatidylcholine (PC) GUV membrane, i.e. from $3.5 \ \mu m^2 s^{-1}$ to $1.5 \ \mu m^2 s^{-1}$.\textsuperscript{34} This agreement suggests a correlation between the diffusivity of the SUV and that of the underlying membrane. Another possible cause for the observations in Figure 3a would be the presence of cholesterol-enriched domains within the GUV membrane, that act as diffusion barriers for the SUVs. In systems where a high-$T_m$ (liquid crystal-crystalline phase transition temperature) and a low-$T_m$ lipid are mixed with cholesterol (e.g. POPC/sphingomyelin/cholesterol), macroscopic phase separations are readily observable.\textsuperscript{35} However, observations of phase separation for a binary phosphocholine/cholesterol system (which is more similar to our study) appear to be ambiguous and dependent on the techniques one employs and contrasting results have been reported. For instance coexisting fluid domains have not been detected using fluorescence microscopy,\textsuperscript{36} while they have been observed using EPR\textsuperscript{37}. Hence, to elucidate if phase separation occurs in the 40 mol% cholesterol sample is largely outside the scope of the current study.

To further elucidate the role of the diffusivity of the underlying membrane, we conducted an experiment in which the zwitterionic DOPC lipid in the GUV membrane is replaced with the 1,2-dimyristoyl-sn-glycero-3-phosphocholine (DMPC) lipid. As for DMPC $T_m = 24^\circ C$,\textsuperscript{38} individual lipids in the GUV membrane are essentially immobile in this experiment ($22^\circ C$). Figure 3b shows the SUV trajectories on the DMPC membrane, defined as the SUV angular positions along the rim ($\theta$ in Figure 2c) as functions of time. As the trajectories appear as straight horizontal lines, we conclude that the SUVs are immobile on the DMPC surface. This observation provides strong evidence that SUV mobility is linked to the lipid mobility in the GUV membrane, or more specifically, as demonstrated below, to the viscosity of the GUV membrane.
Figure 3. Effect of membrane properties on SUV diffusivity. (a) Histograms of diffusion coefficients $D$ for surface-bound SUVs before (red bars) and after (blue bars) adding 40 mol% cholesterol to the underlying GUV membrane. The addition of cholesterol is seen to slow down the diffusional motion of the vesicles. Note the logarithmic x-axis. (b) Trajectories of SUVs on a gel-phase GUV. The horizontal lines on the (angle, time) - plane indicate that the SUVs are not moving.

Our single-vesicle tracking method allows studying the effect of the vesicle size on SUV diffusivity on the membrane surface. Assuming that the number of fluorescence dye (Rho-PE) molecules is proportional to the SUV surface area, we determine the size of an individual vesicle from the square root of the fluorescence emitted by the vesicle. Figure 4a shows the correlation between the adhering vesicle size and its diffusivity by means of the joint probability density function. The observed symmetry of this function (with a cross correlation of -0.07) indicates a very weak size dependence of the diffusivity of adhering vesicles, which is markedly different from the diffusivity in the bulk, which depends inversely on the size, as given by the Stokes - Einstein relation for particle diffusion in three dimensional (3D) fluids:

$$D = \frac{kT}{6\pi\alpha \eta},$$

where $\eta$ is the fluid viscosity, $\alpha$ is the particle radius and $kT$ is the Boltzmann energy. In contrast to hydrodynamics in 3D fluids, the hydrodynamics in (quasi) two-dimensional (2D) fluids, such
as bilayers, is relatively size independent, as predicted by the Saffman-Delbrück model for the
diffusivity of membrane inclusions, that is based on the hydrodynamic coupling between the
membrane and the surrounding bulk:\textsuperscript{30}

\[ D = \frac{kT}{4\pi \eta_m} \left( \log\left( \frac{\eta_m}{\eta_a} \right) - \gamma \right). \quad (2) \]

Eq. (2) adequately describes the diffusion of proteins\textsuperscript{41} as well as phase separated domains\textsuperscript{18}
within the lipid bilayer, as well as that of lipid clusters associated to externally adhering particles\textsuperscript{22}. In Eq. (2), \( \eta \) and \( \eta_m \) are the viscosities of the solvent and the membrane, respectively, \( a \) is the
inclusion radius and \( \gamma \approx 0.58 \) is Euler's constant. Eq. (2) can be regarded as the (pseudo) 2D
analog to the Stokes - Einstein relation for particles diffusing in 3D fluids [Eq. (1)], where \( \eta_m \)
takes over the role of \( \eta \). In contrast to Eq. (1), where we have an inverse dependence on particle
size, Eq. (2) reveals a weak, logarithmic dependence on size. Note also that the membrane
viscosity \( \eta_m \) is closely related to the membrane diffusivity; a large membrane viscosity
corresponds to a low membrane diffusivity and conversely.

An unnoticeable size dependence (Figure 4a) supports that the diffusivity of GUV-riding
SUVs is dominated by the membrane viscosity and not by the bulk viscosity. To further verify
the inferior role of the bulk viscosity, we increase this quantity by a factor two from 1.0 to 2.3 g
m\textsuperscript{-1} s\textsuperscript{-1} by adding 25\% (v/v) glycerol to the solvent, which corresponds to a concentration of 3.4
M. Upon adding glycerol, the average diffusivity decreases only 25\% from 1.3 ± 1.0 to 1.0 ± 0.6,
which is small compared to the two-fold increase in the viscosity. This confirms that the solvent
viscosity plays an inferior role in the diffusivity of adhering vesicles, which, as argued above, is
mainly controlled by the viscosity of the underlying membrane. Figure 4b illustrates the contrast
in diffusivity between vesicles on the membrane surface and vesicles in the bulk, by showing the
measured vesicle diffusivity as a function of the bulk friction coefficient \( \eta_a \). For freely floating
vesicles [Eq. (1)] the diffusivity is inversely proportional to this parameter, while the relation for surface adhering vesicles is much weaker.

![Figure 4](image)

Figure 4. Effect of SUV size, solvent viscosity and SUV charge density on the vesicle diffusivity. (a) The joint probability density function of vesicle diffusivity $D$ and radius $a$. The size is reconstructed from the fluorescence intensity. Symmetry of this function indicates that size and diffusivity are uncorrelated. (b) Vesicle diffusivity $D$ versus bulk friction coefficient $\eta a$ for freely floating vesicles (blue) and surface adhering vesicles (red). The dashed lines correspond to theory for freely floating vesicles [Eq. (1); blue] and membrane inclusions [Eq. (2); red]. (c) Diffusivity $D$ for membrane-adhering SUsVs containing 1 mol% and 5 mol% negatively charged lipids.

The experimental evidence presented above, strongly suggests that, instead of the bulk viscosity, the SUV diffusivity is governed by the viscosity of the underlying GUV membrane $\eta_M$. To validate this hypothesis, we follow the approach presented by Hormel et al. and calculate $\eta_M$ by inserting the measured $D$ into the equation for the diffusivity for membrane inclusions [Eq. (2)]. In order to apply this model, we assume that the mobility of the SUV is coupled to that of a cluster of bound lipids within the GUV membrane. The cluster moves in a surrounding membrane of DOPC:DOEPC (9:1). Due to electrostatic attraction, it is conceivable, that the charged DOEPC lipids concentrate in the clusters, which may lower the DOEPC concentration in the surrounding membrane. Given the extremely low coverage of the clusters (they cover a fraction of $\sim 10^{-7}$ of the GUV surface), this effect is negligible and the clusters are assumed to move in a surrounding membrane, that is unaffected by the presence of the clusters. As the cluster is electrostatically bound to the SUV, we suppose that it is within one Debye length of the
For spherically shaped SUVs (radius \( a \)) and in the limit \( a/\lambda \gg 1 \), the radius \( a_c \) of this contact area equals (see Supplemental Materials):

\[
a_c = \sqrt{2a\lambda}.
\]  

In our system we have \( a = 58 \text{ nm} \) (Figure S2a) and \( \lambda = 4.8 \text{ nm} \) (4 mM NaCl), which gives \( a_c = 23 \text{ nm} \). Inserting this, together with the measured SUV diffusivity \( D = 1.3 \pm 1.0 \mu\text{m}^2\text{s}^{-1} \) into Eq. (2) we then obtain a membrane viscosity of: \( \eta_M = (7 \pm 5) \times 10^{-10} \text{ kg s}^{-1} \).

This value agrees well with previously found values for DOPC lipids based on fluorescence recovery after photobleaching: \( \eta_M = 2 \times 10^{-10} \text{ kg s}^{-1} \),\(^{15} \) and based on fluorescence lifetime imaging: \( \eta_M = 4 \times 10^{-10} \text{ kg s}^{-1} \).\(^{13} \) The agreement supports the notion that vesicle diffusivity \( D \) is described by theory for membrane inclusions. The applicability of this theory to the present experiment is illustrated in Figure 4b, where we compare Eq. (2) to the measured diffusivity as a function of the bulk friction coefficient \( \eta a \).

Applicability of Eq. (2) to the present system suggests that each SUV is coupled to a disk-like GUV domain, illustrated in Figure 1b by the red regions. The quasi 2D hydrodynamics of the membrane disk\(^{40} \) explains the observed size-insensitivity of the SUV diffusivity (Figure 4a). Here we present one final control experiment to support this insensitivity. For this purpose, we reduce the fraction of the negatively charged lipids (DOPS) in the SUV membrane five-fold, i.e. from 5 mol\% to 1 mol\%. As shown in Figure 4c, this change in the charge density does not appreciably affect the SUV diffusivity, which decreases insignificantly from 1.3 ± 1.0 to 1.1 ± 0.8 \( \mu\text{m}^2\text{s}^{-1} \) (Figure 4c). Insensitivity of the diffusivity with respect to the magnitude of the electrostatic attraction is in line with the weak (logarithmic) size dependence in Eq. (2).

However, since zwitterionic SUVs have a small but measurable negative surface potential\(^{43} \), it is possible that increasing the DOPS concentration from 1 mol\% to 5 mol\%
changes the overall surface potential only moderately. To elucidate this, another control experiment was done, using SUVs composed of pure zwitterionic DOPC, which were found not to adsorb onto the positively charged GUV surface. Therefore, the negative surface potential of zwitterionic SUVs is expected to be small compared to the potential at 1 mol% DOPS, and changing the DOPS concentration from 1 mol% to 5 mol% changes the overall surface potential substantially.

**Conclusions**

We use confocal microscopy to track the motion of electrostatically adhering SUVs riding onto the GUV surface. We find that the SUVs execute a two-dimensional Brownian motion, which is insensitive to the solvent viscosity, vesicle radius, vesicle charge density and vesicle fluidity, but is correlated with the fluidity of the underlying GUV membrane instead. Assuming that the adhesion zone defines the size of a disk-shaped inclusion in the GUV membrane – which moves in concert with the SUV – we extract the membrane viscosity within the Saffman - Delbrück framework. A good agreement with previous measurements of membrane viscosity supports the notion that the mobilities of the SUVs are electrostatically linked to that of bound lipid clusters within the target GUV membrane. A major implication of our work is that the electrostatic tethering of small vesicles to cell-sized giant vesicles can divide the diffusional properties of lipid membranes into separate, distinct populations, i.e. clusters of lipids and individual lipids, where the clusters are bound by the SUVs and diffuse slowly within their own milieu, (see Figure 1b). In this context it is noted that, while the Saffman – Delbrück model pertains to the diffusivity of rigid, disk-shaped membrane inclusion, the lipids in the cluster are not necessarily rigidly bound to the SUV, but are more likely in dynamic equilibrium with the free lipids in the membrane. Our work also demonstrates that the reversible binding of SUVs to target membranes
offers a simple non-perturbative means to measure membrane viscosity – an important parameter, whose measurement has long remained challenging.

ASSOCIATED CONTENT
Supporting Information
The Supporting Information is available free of charge on the ACS Publications website.
Image analysis and additional supporting figures (Figures S1-S2) and supporting Table 1. (PDF)

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Notes
The authors declare no competing financial interest.

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