Membrane Nanotubes Increase the Robustness of Giant Vesicles

Tripta Bhatia,† Jaime Agudo-Canalejo,‡§ Rumiana Dimova,† and Reinhard Lipowsky*,†‡

†Theory & Biosystems, Max Planck Institute of Colloids and Interfaces, 14424 Potsdam, Germany
‡Rudolf Peierls Centre for Theoretical Physics, University of Oxford, Oxford OX1 3NP, U.K.
§Department of Chemistry, The Pennsylvania State University, University Park, Pennsylvania 16802, United States

Supporting Information

ABSTRACT: Giant unilamellar vesicles (GUVs) provide a direct connection between the nano- and the microregime. On the one hand, these vesicles represent biomimetic compartments with linear dimensions of many micrometers. On the other hand, the vesicle walls are provided by single molecular bilayers that have a thickness of a few nanometers and respond sensitively to molecular interactions with small solutes, biopolymers, and nanoparticles. These nanoscopic responses are amplified by the GUVs and can then be studied on much larger scales. Therefore, GUVs are increasingly used as a versatile research tool for basic membrane science, bioengineering, and synthetic biology. Conventional GUVs have one major drawback, however: they have only a limited capability to cope with external perturbations such as osmotic inflation, adhesion, or micropipette aspiration that tend to rupture the membranes. In contrast, cell membranes tolerate the same kinds of mechanical perturbations without rupture because the latter membranes are coupled to reservoirs of membrane area. Here, we introduce GUVs with membrane nanotubes as model systems that include such area reservoirs. To demonstrate the increased robustness of these tubulated vesicles, we use micropipette aspiration and changes in the osmotic conditions applied to phospholipid membranes doped with the glycolipid GM1. A quantitative comparison between theory and experiment reveals that the response of the GUVs is governed by the membranes’ spontaneous tension, a curvature-elastic material parameter that describes the bilayer asymmetry on the nanoscale. Because of their increased robustness, GUVs with nanotubes represent improved research tools for membrane science, in general, with potential applications as storage and delivery systems and as cell-like microcompartments in bioengineering, pharmacology, and synthetic biology.

KEYWORDS: biomembrane, bilayer asymmetry, giant vesicle, spontaneous tubulation, micropipette aspiration, tube retraction, spontaneous tension

Giant unilamellar vesicles (GUVs) are formed by fluid bilayer membranes that separate interior aqueous microcompartments from the exterior aqueous environment. These membrane systems, which have been studied for some time,1 have attracted much recent interest, both in basic membrane science2–7 and in the context of artificial protocells.3–10 The multiresponsive behavior of the GUVs includes curvature generation by polymers6 and membrane proteins9,10 protein segregation in membrane–membrane interfaces,11 thermally driven transport of intramembrane domains,6 and membrane fission by protein crowding.12 The increased interest in GUVs is further demonstrated by the different microfluidic-based methods that have been developed during the last couple of years to produce large populations of monodisperse GUVs.10–16

In basic membrane science, one wants to stay in the regime of low membrane tension in order to mimic the behavior of cellular membranes. Likewise, when GUVs are used as microcompartments in bioengineering or synthetic biology, one would like to preserve the integrity of the membranes and to avoid membrane rupture. However, when the GUVs are osmotically inflated by hypotonic exterior solutions17 exposed to adhesive surfaces,18 or aspirated by micropipettes,19 their membranes easily rupture in an uncontrolled manner. In contrast, the plasma membranes of eukaryotic cells, which are coupled to reservoirs of membrane area20,21 can tolerate the same kinds of mechanical perturbations without rupture.

Here, we introduce GUVs with spontaneously formed membrane nanotubes as model systems that include such area reservoirs and demonstrate that these vesicles can resist even large mechanical perturbations without membrane rupture. We show that these GUVs tolerate several cycles of strong aspiration into and subsequent release from a micropipette as well as repeated inflation and deflation steps induced

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by changes in the osmotic conditions. In addition, a quantitative analysis of the micropipette experiments reveals that the tubulated vesicles behave, to a large extent, like liquid droplets with a variable surface area and an effective interfacial tension that is provided by the spontaneous tension introduced in ref 22. The latter tension is, in fact, a curvature-elastic material parameter that depends on the bilayer asymmetry and the associated spontaneous curvature but not on the vesicle geometry.

The spontaneous tubulation of vesicles has been observed previously for a variety of lipid compositions.2,23 Here, we use binary mixtures of the phospholipid 1-palmitoyl-2-oleoyl-sn-glycero-3-phosphocholine (POPC) and the ganglioside GM1, together with a small amount of fluorescently labeled lipids, which provide another example for bilayer membranes that undergo spontaneous tubulation. The glycolipid GM1 has attracted much recent interest because it is abundant in all mammalian neurons24 and plays an important role in many neuronal processes and diseases.25 Furthermore, GM1 acts as a membrane anchor for various toxins, bacteria, and viruses such as the simian virus 40.26 Our combined experimental and theoretical study implies, however, that the increased robustness of tubulated vesicles depends primarily on the bilayer asymmetry of these membranes. Therefore, any membrane with a sufficiently large asymmetry will exhibit the same robustness as the lipid membranes considered here. Recently, tubulated vesicles have also been studied by Monte Carlo simulations.27

We aspirate the tubulated GUVs by micropipettes and study the initial aspiration process as defined in Figure 1 in a systematic manner. To the best of our knowledge, such an experimental study has not been performed previously. A quantitative comparison between theory and experiment reveals that the response of the tubulated GUVs is governed by the spontaneous tension of their membranes, a quantity that is directly measured here by analyzing the evolution of the vesicle shape during initial aspiration. Our experimental data confirm the theoretical prediction22 that the presence of membrane nanotubes leads to a mechanical tension that is several orders of magnitude smaller than the spontaneous tension. Indeed, for the bilayer membranes studied here, the spontaneous tension is on the order of 10 pN/μm or 10⁻² mN/m, whereas the absolute value of the mechanical tension is smaller by 2 orders of magnitude. In this way, our results provide insight into the intricate and composite nature of membrane tension. Because of their increased robustness, tubulated GUVs represent improved research tools for membrane science, in general, with potential applications as storage and delivery systems as well as cell-like microcompartments in bioengineering, pharmacology, and synthetic biology.

RESULTS AND DISCUSSION

Spontaneous Tubulation of Giant Vesicles Doped with GM1. We prepared GUVs composed of the phospholipid POPC, a few mol % of the glycolipid (or ganglioside) GM1, and 0.1 mol % of the fluorescently labeled lipid Texas-Red-DHPE (dihexadecanoylglycerophosphoethanolamine) by electrotformation28 as described in the Methods section. The same preparation method has been previously used in ref 29. However, in contrast to the latter study, we subsequently diluted the vesicle suspension ten- to eleven-fold with native buffer, which led to the formation of stable nanotubes inside the GUVs, reflecting the larger GM1 concentration within the inner bilayer leaflets. This bilayer asymmetry is caused by the desorption of the GM1 molecules from the outer leaflets into the exterior solution, which is depleted of GM1 molecules because of the dilution step. After tubulation, the excess membrane area stored in the tubes was typically between 20 and 50% of the total membrane area. At room temperature, the binary lipid mixture of POPC and GM1 undergoes phase separation when the mole fraction of GM1 exceeds about 5 mol %.29 Here, we focused on lipid compositions with 2 and 4 mol % of GM1 for which we did not detect any intramembrane domains.

Examples for tubulated vesicles with these two lipid compositions are displayed in Figure 2. The nanotubes were only visible in the presence of the fluorescently labeled lipids, in

Figure 1. Different stages for the aspiration process of a tubulated GUV (red) by a micropipette (gray) of radius \( R_{pip} \). (a) Spherical mother vesicle comes into contact with the pipette. (b) With increasing suction pressure, some of the nanotubes are retracted, and the mother vesicle develops a tongue that has the form of a spherical cap. (c) When the suction pressure reaches a critical value, the cap-like tongue becomes a hemisphere with radius \( R_{pip} \) and the vesicle membrane starts to flow into the micropipette. (d) Depending on the membrane area stored in the nanotubes, the vesicle motion stops as soon as all nanotubes have been retracted (d, top) or continues until the vesicle is completely aspirated into the pipette (d, bottom).

Figure 2. Confocal cross sections of membrane nanotubes protruding from the mother vesicles (large red circles) into the vesicle interior. The vesicle membranes consist of the phospholipid POPC doped with the fluorescently labeled lipid Texas-Red-DHPE as well as 2 mol % of GM1 in (a) and 4 mol % of GM1 in (b). The nanotubes are only visible in the presence of the fluorescently labeled lipids (red), in agreement with the theoretical analysis which implies that the nanotubes have a width on the order of 100 nm. Scale bars: 10 μm.
agreement with the theoretical analysis described below, which implies that the nanotubes have a width on the order of 100 nm, that is, below optical resolution. For the vesicle in Figure 2b, a three-dimensional stack of confocal xy cross sections is displayed in Movie 1 of the Supporting Information (SI), which reveals the three-dimensional conformations of the nanotubes. Inspection of this movie shows that these tubes are quite long and attain complex and highly curved conformations. Below, we will compare this experimentally observed flexibility with theoretical estimates for the persistence length of the tubes. This comparison indicates that the tubes have a necklace-like morphology.

We also prepared vesicles with nanotubes pointing toward the exterior solution by adding the GM1 molecules to the GUV dispersion without the dilution step (see Figure 3). In the presence of such out-tubes, the vesicles have a fuzzy surface and an ill-defined size. Thus, in the following, we will focus on GUVs with in-tubes that have a smooth surface and a well-defined size as in Figure 2.

The latter GUVs were aspirated by micropipettes (see the Methods section) in order to expose them to a tunable mechanical tension and to retract the tubes in a controlled and reversible manner. During initial aspiration, as shown in Figure 1a–c and Figure 4a,b, the emerging membrane tongue forms a spherical cap with a radius that decreases from the initial radius of the mother vesicle to the radius of the micropipette. The quantitative analysis of the shape evolution during the initial aspiration process allows us to directly measure the spontaneous tension of the GUV membrane, which is intimately related to the bilayer asymmetry and the associated spontaneous curvature of this membrane. To the best of our knowledge, initial aspiration of GUVs by micropipettes has not been studied previously in a systematic and quantitative manner.

**Initial Aspiration: Theory and Experiment.** In the spontaneous curvature model,\textsuperscript{40,51} the vesicle membrane is described by two fluid-elastic parameters, the spontaneous curvature $m$ and the bending rigidity $\kappa$. These two parameters can be combined into an intrinsic tension scale, the spontaneous tension $\sigma \equiv 2km^2$, which varies quadratically with the spontaneous curvature $m$ and represents the bending energy density of a weakly curved membrane segment that has a relatively large spontaneous curvature but cannot adapt its local mean curvature to this spontaneous curvature.\textsuperscript{22} The total membrane tension $\Sigma$ can then be decomposed according to

$$\Sigma = \Sigma_m + \sigma = \Sigma_m + 2km^2$$

where the mechanical tension $\Sigma_m$ depends, in general, on the size and shape of the vesicle as well as on external constraints and forces acting on the vesicle. As shown in ref 22, the spontaneous curvature model\textsuperscript{40,51} leads to a Euler—Lagrange or shape equation that depends explicitly on the total membrane tension $\Sigma$, which is equal to the superposition of the mechanical and the spontaneous tension as in eq 1. In the SI, we utilize this shape equation for spherical and cylindrical membrane segments to derive explicit expressions for the mechanical tension of the tubulated GUVs.

Let us now consider a giant vesicle with nanotubes protruding into the vesicle interior, as shown in Figure 2. The presence of the nanotubes reveals a large bilayer asymmetry or spontaneous curvature $m$. Because the nanotubes point toward the vesicle interior, the spontaneous curvature $m$ is negative, and the tube radius $R_{nt}$ is equal to $1/(2m\ell)$ or $1/lm\ell$ depending on whether the tubes are cylindrical or necklace-like.\textsuperscript{30,52} Tubulated GUVs are characterized by a large separation of length scales: the spherical mother vesicle has a radius $R_{mv}$ on the order of 10 $\mu$m, whereas the nanotubes have a radius $R_{nt}$ that is close to or below optical resolution. For the vesicles

Figure 3. Confocal cross sections of membrane nanotubes protruding from the mother vesicles (red circles) toward the exterior solution. Confocal cross section of two GUVs prepared from POPC and 4 mol % of GM1 that formed out-tubes after the addition of GM1 to the vesicle suspension. The out-tubes have a width below the optical resolution and are labeled by the same fluorophore (red) as in Figure 2. Scale bars: 10 $\mu$m.

Figure 4. Shape evolution of aspirated GUV as observed with fluorescence microscopy. (a,b) Initial aspiration of a tubulated vesicle composed of POPC and 4 mol % of GM1, with 0.1 mol % of the fluorescently labeled lipid Texas-Red-DHPE (red). (a) Mother vesicle (large dotted circle) has the radius $R_{mv}$ and forms a small tongue that has the shape of a spherical cap (small dashed circle). The radius $R_{nt}$ of this cap is smaller than $R_{mv}$ and larger than the pipette radius $R_{pip}$. (b) When the suction pressure reaches a critical value, the tongue attains a hemispherical shape with radius $R_{nt} = R_{mv}$, and the vesicle membrane starts to flow into the pipette. (c) Snapshot of a flowing vesicle just before it is completely aspirated into the pipette. The detailed time evolution of this latter process is displayed in Movie 2 of the SI. Scale bars in (a–c): 10 $\mu$m.
studied here, the tube radius $R_{nt}$ was on the order of 100 nm and, thus, two orders of magnitude smaller than the radius $R_{mv}$ of the mother vesicle, with the ratio $R_{nt}/R_{mv} \lesssim 0.01$. Furthermore, the length of the nanotubes was much larger than the tube width. This separation of length scales implies that the absolute value $|\Sigma|$ of the mechanical tension behaves as

$$|\Sigma| \leq \frac{R_{nt}}{R_{mv}} \sigma \ll \sigma = 2 \kappa m^3 \text{ for } R_{mv} \gg R_{nt} \approx \frac{1}{|\ln l|}$$

(2)

and is therefore several orders of magnitude smaller than the spontaneous tension. As a consequence, we can ignore the mechanical tension, which depends on the size and shape of the GUV, compared to the spontaneous tension, which represents a curvature-elastic material parameter. The inequality in eq 2, which follows from the mechanical equilibrium between the highly curved nanotubes and the weakly curved membrane of the mother vesicle, has been previously derived for cylindrical nanotubes.\(^{22}\) As shown in the SI, the same inequality applies to necklace-like nanotubes, as well.

During the initial aspiration of the tubulated vesicle, the mother vesicle forms a membrane tongue that has the form of a small spherical cap with curvature radius $R_{mv}$ (see Figure 1b and Figure 4a). For such a piece-wise spherical shape, minimization of the membrane’s bending energy leads to the pressure balance

$$\Delta P \equiv P_{ex} - P_{pip} = P_{el}(R_{nt}, R_{mv})$$

(3)

between (i) the suction pressure $\Delta P$, which depends on the pressures $P_{ex}$ and $P_{pip}$ in the exterior aqueous solution and in the pipette, respectively, and (ii) the elastic counter pressure

$$P_{el}(x, y) \equiv \left( \frac{1}{x} - \frac{1}{y} \right) \left( 2 \Sigma + 2 \sigma - 4 \kappa m \left( \frac{1}{x} + \frac{1}{y} \right) \right)$$

(4)

which depends on the radii $x = R_{nt}$ and $y = R_{mv}$ and, thus, on the vesicle geometry. The relationships 3 and 4 are explicitly derived in the SI. These relationships describe the mechanical balance between the suction pressure $\Delta P$, which represents an external control parameter, and the counter pressure $P_{el}$ arising from the elasticity of the membrane. A suction pressure $\Delta P > 0$ pulls the vesicle into the pipette, whereas a counter pressure $P_{el} > 0$ pushes the vesicle out of the pipette.

It follows from eq 4 that the elastic counter pressure $P_{el}$ depends, in general, both on the mechanical tension $\Sigma$ that stretches the membrane and on the spontaneous tension $\sigma$ that increases the bending energy of the weakly curved membrane segments. However, for the tubulated vesicles considered here, the mechanical tension can be ignored compared to the spontaneous tension, as shown in the SI and summarized by the relation 2. Therefore, the elastic counter pressure as given by eq 4 has the asymptotic form

$$P_{el}(R_{nt}, R_{mv}) \approx 2\sigma \left( \frac{1}{R_{nt}} - \frac{1}{R_{mv}} \right) \text{ for } R_{nt} \gg \frac{1}{|\ln l|}$$

(5)

An analogous equation applies to the micropipette aspiration of liquid droplets,\(^{32-34}\) for which the counter pressure arises from capillary forces, and the spontaneous tension $\sigma$ of the GUV is replaced by the interfacial tension of the droplet.

In the micropipette experiments, the vesicles were aspirated by a tunable suction pressure using the hydraulic pressure system described in the Methods section. The initial aspiration regime could be explored by increasing the suction pressure up to 3 Pa (see Figure 5). Using epifluorescence microscopy, we measured the radius $R_{nt}$ of the emerging tongue and the radius $R_{mv}$ of the mother vesicle. In Figure 5, we plot the suction pressure $\Delta P$ as a function of the inverse length

$$\Delta \kappa \equiv \frac{1}{R_{to}} - \frac{1}{R_{mv}}$$

(6)

for two different values of the overall GM1 concentration. Fitting the experimental data in Figure 5 to the pressure balance in eq 3 with the counter pressure $P_{el}$ as given by eq 5, we obtain the spontaneous tensions $\sigma = 8.33 \pm 0.76$ and $\sigma = 22.2 \pm 1.5$ Pa $\mu$m for 2 and 4 mol % of GM1, respectively. The values of the spontaneous curvature $m$ then follow from $m = -\sqrt{\sigma/(2\kappa)}$, where the minus sign takes into account that the GUVs studied here formed in-tubes.

The bending rigidity $\kappa$ of POPC membranes doped with GM1 has been previously measured in ref 29. In the latter study, the GUVs were prepared without the additional dilution step and thus without the GM1 desorption from the outer leaflet that acts to reduce the final GM1 concentration in the membranes. In the regime of low GM1 fractions, the bending rigidity $\kappa$ was found to be on the order of $10^{-19}$ J. When we combine this $\kappa$ value with the measured values of the spontaneous tension $\sigma$, the relation $m = -\sqrt{\sigma/(2\kappa)}$ leads to the spontaneous curvatures $m \approx -1/(155 \text{ nm})$ and $m \approx -1/(95 \text{ nm})$ for 2 and 4 mol % of GM1, with an accuracy of about $\pm 20\%$. In order to adapt their shape to these spontaneous curvatures, the membranes form necklace-like or cylindrical nanotubes with a radius of $1/\text{ml}$ or $1/(2\text{ml})$, respectively. Therefore, we conclude that the tubes displayed in Figure 2 have a diameter below or close to optical resolution for both 2 and 4 mol % of GM1.

The deduced values of the spontaneous curvatures allow us to distinguish necklace-like and cylindrical nanotubes by estimating the persistence lengths of both types of tubes and then comparing these estimates with the thermally excited undulations of the tubes as observed experimentally. As shown in ref 2, the persistence length of a cylindrical nanotube is $\pi \kappa/(1
which was kept at constant aspiration pressure until the vesicle was completely sucked up and disappeared into the pipette, as visible in the last frame of Movie 2.

Essentially the same mechanical instability has been observed for liquid droplets, neutrophils (i.e., certain types of white blood cells) and fibroblasts. For liquid droplets, the parameter that replaces the spontaneous tension $\sigma$ in the cortex tension generated by the actomyosin cortex. For fibroblasts, this parameter was decomposed into the cortical tension, and another tension that remains after the cortex has been disassembled chemically by cytochalasin D. It is interesting to note that the membrane tension measured in ref 36 after the disassembly of the actin cortex is on the same order of magnitude as the spontaneous tensions obtained here.

The membrane flow arising from the mechanical instability of tubulated vesicles can lead to two different final states, depending on the membrane area initially stored in the nanotubes. If this area is relatively small, all of these tubes will be retracted before the whole vesicle has been sucked into the pipette, and the vesicle reaches a new equilibrium state, as shown by the top cartoon in Figure 1d. Further aspiration then leads to membrane stretching until membrane rupture and vesicle bursting, as demonstrated in Figure 6 for a GUV membrane without GM1.

On the other hand, if the membrane area stored in the tubes is sufficiently large, the vesicle will be aspirated completely into the micropipette with some remaining area stored in tubes, see Figure 1d, bottom and Movie 2 in the SI. The tube retraction is completely reversible: as long as the vesicle has not been sucked up completely by the micropipette, we can recover the nanotubes by reversing the changes in aspiration pressure at each stage of the aspiration process. In addition, the tubulated GUVs are sufficiently robust to tolerate repeated cycles of tube retraction coupled to in-flow of the membrane tongue and subsequent tube elongation coupled to out-flow of this tongue (see Movie 3 in the SI). The latter movie displays the response of one tubulated GUV exposed to several changes in the suction pressure and represents a sequence of five subsequent recordings including three full cycles of strong aspiration into the micropipette and subsequent release from this pipette. The total recording time of Movie 3 exceeded 1 h. On this time scale, we did not detect any significant changes in the behavior of the tubes, which indicates that the bilayer asymmetry and the associated spontaneous tension of the GUV membrane were not affected by the recurrent mechanical stress generated by the

$m k_0 T$, whereas the same length of a necklace-like tube is expected to be on the order of $2/lm$. For a bending rigidity $\kappa \approx 10^{-10}$ and a spontaneous curvature of about $1/(100 \text{ nm})$, we then obtain a persistence length of about $200 \text{ nm}$ and $6 \text{ } \mu \text{ m}$ for necklace-like and cylindrical nanotubes, respectively. Inspection of Figure 2 and Movie 1 reveals many sharp bends on the necklace-like and cylindrical nanotubes, respectively. Inspection of Figure 2 and Movie 1 reveals many sharp bends on the nanotubes, which indicates that the tubes are necklace-like.

### Critical Suction Pressure and Mechanical Instability

The initial aspiration process continues until the vesicle tongue attains a hemispherical shape. For this shape, the tongue radius $R_0$ is equal to the pipette radius $R_{pip}$ and the mother vesicle has a certain size $R_{mv} = R_{mv}^\ast$. The pressure balance then becomes

$$\Delta P = P_{el}^\ast \equiv P_0(R_{pip}, R_{mv}^\ast)$$

which defines the critical pressure $P_{el}^\ast$. The latter pressure has the simple form

$$P_{el}^\ast \approx 2\sigma \left( \frac{1}{R_{pip}} - \frac{1}{R_{mv}^\ast} \right) \text{ for } R_{pip} \ll \frac{1}{\mu \text{ m}}$$

The radius $R_{mv}^\ast$ of the mother vesicle at the critical pressure depends primarily on the initial size of the vesicle and on the pipette radius.

The pressure balance between the suction pressure and the elastic counter pressure as described by eq 3 leads to a sequence of stable equilibrium states during initial aspiration, that is, as long as $R_0 > R_{pip}$ and $\Delta P < P_{el}^\ast$. Thus, consider a state of mechanical equilibrium with $\Delta P = P_{el}(R_{mv}) < P_{el}^\ast$ and a perturbation of this state that moves the tongue further into the pipette, thereby reducing the tongue radius to $R_{0\,'} < R_0$ for fixed suction pressure $\Delta P$. The resulting counter pressure $P_{el}(R_{mv})$ exceeds the equilibrium value $P_{el}(R_{mv}) = \Delta P$ as follows from eq 5, and the positive pressure difference $P_{el}(R_{mv}) - \Delta P$ pushes the tongue back toward its equilibrium state.

In contrast, a perturbation that moves the tongue further into the pipette for the critical suction pressure $\Delta P = P_{el}^\ast$ now leads to a smaller radius $R_{mv}\,' < R_{mv}^\ast$ of the mother vesicle, whereas the tongue radius $R_0 = R_{pip}$ remains unchanged. It then follows from eq 5 that the counter pressure $P_{el}(R_{mv}^\ast)$ is smaller than the suction pressure $\Delta P$, which implies that the tongue is pushed even further into the pipette. As a consequence, as soon as the suction pressure has reached the critical value $P_{el}^\ast$, the vesicle membrane starts to flow into the micropipette, as observed experimentally; see Movie 2 in the SI. This movie displays a sequence of confocal $xy$ cross sections of the flowing vesicle

The mechanical tension arising from the suction pressure $U_{suc}$ for $1 \mu \text{ m}$ diameter GUVs can be estimated to be

$$U_{suc} \approx \frac{1}{2} \sigma A_{mic} \left( 1 - \frac{R_0}{R_{pip}} \right)$$

where $A_{mic}$ is the cross-sectional area of the micropipette.

The above expression 5 for the elastic counter pressure is the interfacial tension of the liquid–liquid interface. For neutrophils, the corresponding parameter was proposed to be the cortical tension generated by the actomyosin cortex. For fibroblasts, this parameter was decomposed into the cortical tension, and another tension that remains after the cortex has been disassembled chemically by cytochalasin D. It is interesting to note that the membrane tension measured in ref 36 after the disassembly of the actin cortex is on the same order of magnitude as the spontaneous tensions obtained here.

The membrane flow arising from the mechanical instability of tubulated vesicles can lead to two different final states, depending on the membrane area initially stored in the nanotubes. If this area is relatively small, all of these tubes will be retracted before the whole vesicle has been sucked into the pipette, and the vesicle reaches a new equilibrium state, as shown by the top cartoon in Figure 1d. Further aspiration then leads to membrane stretching until membrane rupture and vesicle bursting, as demonstrated in Figure 6 for a GUV membrane without GM1.

On the other hand, if the membrane area stored in the tubes is sufficiently large, the vesicle will be aspirated completely into the micropipette with some remaining area stored in tubes, see Figure 1d, bottom and Movie 2 in the SI. The tube retraction is completely reversible: as long as the vesicle has not been sucked up completely by the micropipette, we can recover the nanotubes by reversing the changes in aspiration pressure at each stage of the aspiration process. In addition, the tubulated GUVs are sufficiently robust to tolerate repeated cycles of tube retraction coupled to in-flow of the membrane tongue and subsequent tube elongation coupled to out-flow of this tongue (see Movie 3 in the SI). The latter movie displays the response of one tubulated GUV exposed to several changes in the suction pressure and represents a sequence of five subsequent recordings including three full cycles of strong aspiration into the micropipette and subsequent release from this pipette. The total recording time of Movie 3 exceeded 1 h. On this time scale, we did not detect any significant changes in the behavior of the tubes, which indicates that the bilayer asymmetry and the associated spontaneous tension of the GUV membrane were not affected by the recurrent mechanical stress generated by the
micropipette. Another example for the increased robustness of tubulated vesicles is provided by their response to changes in the osmotic conditions, as illustrated in Figure 7. In this figure, a tubulated vesicle is first inflated osmotically until essentially all nanotubes have been retracted. When the latter vesicle is subsequently deflated, the resulting membrane excess area is again stored in membrane nanotubes.

CONCLUSIONS

In this paper, we introduced GUVs with spontaneously formed membrane nanotubes as model systems that mimic cellular membranes with reservoirs for membrane area and demonstrated that these systems resist even strong mechanical perturbations without rupture. A quantitative comparison between theory and experiment revealed that tubulated GUVs respond to mechanical perturbations in close analogy to liquid droplets, with the spontaneous tension of the membranes playing the same role for tubulated GUVs as the interfacial tension for liquid droplets. In fact, the spontaneous tension \( \sigma = 2km^2 \) represents a curvature-elastic material parameter that determines both the initial aspiration process (see eqs 3–5) and the critical suction pressure as given by eq 8. For the lipid compositions studied here, the numerical values of the spontaneous tension \( \sigma \) have been determined from the shape evolution of the membrane tongue during initial aspiration (Figure 4a,b and Figure 5). To the best of our knowledge, this initial aspiration process has not been studied previously in a systematic and quantitative manner. When we increase the GM1 concentration from 2 to 4 mol \%, the spontaneous tension is increased from \( \sigma \approx 8.3 \) to 22 pN/\(\mu m\), and the spontaneous curvature from \( m \approx -1/(155 \text{ nm}) \) to \(-1/(95 \text{ nm})\), respectively.

In the present study, we used a specific lipid composition to produce giant vesicles with nanotubes, but such vesicles can also be produced for other types of lipid compositions. One example is provided by ternary lipid mixtures of DOPC, DPPC, and cholesterol which form nanotubes both in their liquid-ordered and in their liquid-disordered phase when the two leaflets of the membranes are exposed to different PEG concentrations. In fact, any lipid membrane with a sufficiently strong bilayer asymmetry can form such tubes. We demonstrated the increased robustness of tubulated GUVs by micropipette aspiration with a tunable suction pressure (Figure 3 and Movie 2 in the SI) and by changes in the osmotic conditions (Figure 7). The micropipette experiments directly imply that GUVs with nanotubes can pass through small blood vessels or microfluidic channels without rupture even if the diameter of these vessels and channels is much smaller than the vesicle size. In the latter context, the tubulated vesicles could be used as storage and delivery systems. One example is provided by nonadhesive nanoparticles that are confined within necklace-like nanotubes. Tube retraction in response to external perturbations will then lead to the partial release of the stored particles. Another example is the enrichment of adhesive Janus particles within the nanotubes as a result of curvature-induced forces that move such particles toward the highly curved tubes after they have been adsorbed onto the weakly curved mother vesicle.

It is instructive to compare the approach described here, based on tubulated GUVs, with other methods that have been developed to increase the stability and robustness of giant vesicles. To increase the rupture (or lysis) tension of lipid bilayers, two classical approaches are to use polymerizable lipids\(^{46}\) or to add additional molecular components such as cholesterol.\(^{39}\) One may also replace the lipids by diblock copolymers to form polymeric GUVs or polymersomes,\(^{40,41}\) which have an increased rupture tension, as well.\(^{62}\) In the context of whole-GUV patch clamping, the contact area between the giant vesicle and the inner surface of the micropipette can be stabilized by \(\beta\)-casein that prevents the vesicle membrane from spreading further into the pipette.\(^{45}\) A general strategy to mechanically stabilize GUVs that has been introduced quite recently is to assemble these vesicles within microfluidic emulsion droplets and to use the droplet surfaces as mechanical scaffolds for the vesicles.\(^{10}\) Compared to these alternative strategies, the tubulated GUVs described here have a number of significant advantages: (i) The lipid membranes considered here exhibit the usual fluidity of phospholipid membranes, which mimics the fluidity of cellular membranes. (ii) These lipid membranes are biocompatible and provide the most natural environment for membrane proteins. (iii) The tubulated GUVs are fully immersed in water and are accessible to solutes and biomolecules from both the exterior and the interior aqueous solution. (iv) The increased robustness of tubulated GUVs arises from internal membrane reservoirs, in close analogy to cellular membranes, which implies that these
giant vesicles exhibit cell-like responses to mechanical perturbations.

Indeed, giant vesicles with membrane nanotubes will also tolerate other mechanical perturbations, arising, for example, from the adhesion and engulfment of large bodies, in close analogy to the cellular uptake via phagocytosis and pinocytosis, or in response to constriction forces that can lead to fission of the GUVs into smaller membrane compartments. The latter process of artificial cytokinesis is an important objective for the bottom-up assembly of artificial protocells.1,28 1-Palmitoyl-2-oleoyl-sn-glycero-3-phosphatidylcholine was purchased from Avanti Polar Lipids (dissolved in a chloroform solution to a concentration of 10 mg/mL). GM1 was purchased from Avanti Polar Lipids and dissolved in a dichloromethane/methanol (2:1) solution to a concentration of 3 mM. Two samples were prepared: (i) 3 mM POPC with 2 mol % of GM1 and (ii) 3 mM POPC with 4 mol % of GM1. For fluorescence imaging of the vesicles, additionally 0.1 mol % of Texas-Red-DHPE was added to samples 1 and 2. Two indium–tin oxide (ITO)-coated glass plates (Delta Technologies Ltd.) were preheated at 55 °C for 5–10 min, and an aliquot of 10 μL of the sample solution was spread onto the ITO surface to form a closed chamber. HEPES buffer (1 mM, pH 7.4, 0.5 Na HEPES; Sigma-Aldrich, St. Louis, MO) was preheated at 55 °C and filled into the chamber (2 mM HEPES buffer was used in the experiments displayed in Figure 7). A sinusoidal AC electric field at 10 Hz was applied for electroswelling the lipid films at 55 °C. In the first phase of the electroswelling process, the amplitude of the applied field was linearly increased from 0.1 V (peak to peak) to 0.8 V (peak to peak) over 30 min. Thereafter, the amplitude of the AC field was kept constant during 60–120 min to grow the vesicles. Finally, during the last 60 min, the GUVs were separated from the ITO surface by linearly decreasing the voltage and frequency to 0.5 V and 1 Hz. After electroswelling, GUVs were cooled to 23 °C in an oven at a rate of 0.1 °C/min. The resulting vesicles were found to be stable and could be transferred to an observation chamber for optical investigations. The latter chamber was closed using sealing paste (Korasilon from Carl Roth, #0856.1) to stop fluid flow and evaporation.

Formation of Out-Tubes after External Addition of GM1. GM1 was dissolved in a dichloromethane/methanol (2:1) solution to a concentration of 3 mM. The organic solvent was removed under vacuum at 55 °C. Milli-Q water was added to reach a GM1 concentration of 3 mM, and the sample was stirred. One microliter of this GM1 stock solution was added to 500 μL of GUV suspension to increase the concentration of GM1 in the outer leaflet by spontaneous insertion of the molecules from the surrounding concentrations. The chamber was closed with the same sealing paste. Immediately afterward, the vesicles were observed to exhibit out-tubes as in Figure 3. Micropipette Aspiration System. Micropipettes were prepared from glass capillaries (World Precision Instruments Inc.) that were pulled using a pipette puller (Sutter Instruments, Novato, CA). Pipette tips were cut using a microforge (Narishige, Tokyo, Japan) to obtain tips with an inner diameter between 4 and 6 μm. Adhesion of the membrane to the pipette was prevented by incubation of the pipette tips in 1 mg/mL aqueous solution of casein (Sigma). A new pipette was used for the aspiration of each GUV. After the pipette was inserted into the observation chamber, the zero pressure across the pipette tip was attained and calibrated by watching the flow of small particles within the tip. The aspiration pressure was controlled through adjustments in the height of a reservoir mounted on a linear translational stage (M-531.PD; Physik Instrumente, Germany); for further details see the Supporting Information of ref 44. This setup allowed the pressure to increase up to 2 kPa with a pressure resolution of 1 mPa. The pressure was changed by displacing the water reservoir at a speed of 0.01 mm/s. The displacement was then stopped for 2–3 min before any data were recorded.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.nano.8b00640.

Supplementary text with derivations of eqs 2–4 (PDF)
Movie captions (PDF)
Movie 1 (MPG)
Movie 2 (MPG)
Movie 3 (AVI)

AUTHOR INFORMATION

Corresponding Author
*E-mail: lipowsky@mpi.kg.mpg.de.

ORCID

Reinhard Lipowsky: 0000-0001-8417-8567

Notes

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