Quantitative prediction and measurement of Piezo's membrane footprint

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Piezo proteins are mechanosensitive ion channels that can locally curve the membrane into a dome shape [Y. R. Guo, R. MacKinnon, eLife 6, e33660 (2017)]. The curved shape of the Piezo dome is expected to deform the surrounding lipid bilayer membrane into a membrane footprint, which may serve to amplify Piezo's sensitivity to applied forces [C. A. Haselwandter, R. MacKinnon, eLife 7, e41968 (2018)]. If Piezo proteins are embedded in lipid bilayer vesicles, the membrane shape deformations induced by the Piezo dome depend on the vesicle size. We employ here membrane elasticity theory to predict, with no free parameters, the shape of such Piezo vesicles outside the Piezo dome, and show that the predicted vesicle shapes agree quantitatively with the corresponding measured vesicle shapes obtained through cryoelectron tomography, for a range of vesicle sizes [W. Helfrich, Z. Naturforsch. C 28, 693-703 (1973)]. On this basis, we explore the coupling between Piezo and membrane shape and demonstrate that the features of the Piezo dome affecting Piezo's membrane footprint approximately follow a spherical cap geometry. Our work puts into place the foundation for deducing key elastic properties of the Piezo dome from membrane shape measurements and provides a general framework for quantifying how proteins deform bilayer membranes.

Piezo ion channels | mechanosensation | mechanical gating | membrane mechanics

In common with other organisms, vertebrates possess a variety of senses that respond to mechanical stimuli (1). Despite intense efforts, the molecules and physical mechanisms underlying vertebrate mechanosensation have long remained elusive. In 2010, Piezo proteins-Piezo 1 and Piezo 2 in mammals-were discovered, which has led to stunning progress in the elucidation of the molecular basis for mechanosensation (2). Piezo channels are mechanosensitive ion channels that open in response to mechanical force. They contain a central pore and three long arms that extend away from the protein center (3-5). The extended arms are made of transmembrane helices and do not lie in a plane in their closed conformation, which induces the lipid bilayer membrane to curve in between the arms (3). We refer to the Piezo ion channel protein plus the curved lipid bilayer contained within the channel's approximate perimeter as the Piezo dome, with the lipid membrane connecting smoothly across the Piezo dome boundary. Piezo channels are thus expected to deform the lipid membrane outside the Piezo dome (6). We refer to the region of deformed lipid bilayer membrane outside the Piezo dome as Piezo's membrane footprint (7). The mechanical properties of the membrane footprint may have interesting consequences for how Piezo responds to mechanical stimuli and may serve to amplify Piezo's sensitivity.

The theoretical study of lipid bilayer-protein interactions has a long and rich history, suggesting that protein-induced bilayer shape deformations can be captured by membrane elasticity theory (7-12). Membrane elasticity theory should, therefore, be able to account quantitatively for the membrane shape deformations induced by the Piezo dome. In particular, if Piezo proteins are embedded in lipid bilayer vesicles, membrane elasticity theory should be able to predict the shape of such Piezo vesicles outside the Piezo dome. The aim of this paper is threefold. First, we provide quantitative comparisons between the predicted and observed shapes of Piezo vesicles. We thus test the continuum elasticity theory predicting the shape of Piezo's membrane footprint, and develop a general framework for quantifying how proteins deform bilayer membranes. Second, we employ our experimental measurements and theoretical predictions of Piezo vesicle shape to explore the coupling between Piezo and membrane shape. In particular, we investigate to what extent the geometry of the Piezo dome can be described by a spherical cap, as proposed previously based on cryoelectron microscopy (cryo-EM) and high-speed atomic force microscopy (HS-AFM) experiments (3, 13). Third, a quantitative understanding of Piezo vesicle shape can be used to provide insight into the elastic properties of the Piezo dome underlying its mechanosensory function, which we explore in the companion paper (14).

Significance

Studies over the past five decades have yielded a beautiful physical theory of how membrane proteins deform lipid bilayers, akin to the bending of a thin elastic shell. The resulting membrane footprints of proteins have been proposed to play important biological roles in protein regulation, membrane remodeling, and membrane organization. But it has been challenging to test and further develop the membrane elasticity theory of bilayer-protein interactions through direct comparison of predicted and measured membrane footprints. We quantitatively measure how Piezo proteins deform lipid bilayer vesicles, and show that the shape of these vesicles can be predicted accurately through membrane elasticity theory. We provide a general approach, of potentially wide applicability, for studying how proteins couple to membrane shape.

Author contributions: C.A.H. and R.M. developed the theory to analyze Piezo vesicles; Y.R.G. and Z.F. produced Piezo vesicles, collected tomograms, and digitized the vesicle profiles; and C.A.H. and R.M. applied the theory and wrote the paper.

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This study uses mathematical concepts and methods that might not be familiar to some readers. While the complete derivations for equations are presented in *SI Appendix*, we still outline the mathematical concepts at each step in the text. The paper is written, we hope, so that the logic can be followed and appreciated even if mastering some details might require further study (15–18).

Results

Measuring and Parameterizing Piezo Vesicles. Piezo 1 channels were reconstituted into 1-palmitoyl-2-oleoyl-sn-glycero-3-phosphocholine (POPC):1,2-dioleoyl-sn-glycero-3-phospho-L-serine (DOPS):cholesterol (8:1:1) vesicles, frozen in vitrified ice, and analyzed using cryo-EM tomography. Due to the intrinsic curvature of Piezo, the channels nearly exclusively reconstitute with their extracellular surface pointed toward the inside of the vesicle. For tomographic analysis, we identified Piezo vesicles containing Piezo near the outer edge of the vesicle, as viewed in projection in electron microscopy (EM) images (Fig. 1A). Piezo vesicles are identifiable because the Piezo C-terminal extracellular domain (CED) (Fig. 1A) is visible adjacent to the membrane on the inside of the vesicle. Images were then collected while tilting the specimen stage between $+42^{\circ}$ and -42° (Fig. 1*B*). When threedimensional (3D) tomographic reconstructions were generated, Piezo vesicles appeared to contain holes at opposite sides of the vesicle due to the limited range of tilt angles. Because we initially selected Piezo vesicles with Piezo near the outer edge of the vesicle as viewed in projection in EM images, we could define a cross-section of the vesicle that included Piezo and a band of membrane encircling the vesicle's long axis. In other words, if we

define Piezo's location as the Piezo vesicle's north pole, the tomogram defines a continuous strip of membrane around the Piezo vesicle that contains both the vesicle north and south poles, the south pole being that region on the vesicle surface farthest from Piezo (Fig. 1*B*). From this reconstruction, we cut out a plane that contains the long axis of the Piezo vesicle intersecting the north and south poles. This cutout is called the oriented Piezo vesicle image. For each Piezo vesicle, the tomographic analysis used here yields one oriented Piezo vesicle image. To be clear, the shape of the oriented Piezo vesicle image is generally not identical to the shape of the Piezo vesicle in projection in the sample image. The shapes would be the same if the Piezo channel was exactly on the edge of the Piezo vesicle in the projected EM image. The tomogram ensures that it is, within a small error, on the edge in the oriented image.

Piezo is a trimer of identical subunits with threefold rotational symmetry (Fig. 1C). The lipids used in the present study form spherical vesicles in the absence of Piezo (3). Thus, we expect that within a short distance away from Piezo, excess free membrane folds will be minimized, and the threefold symmetry of the Piezo channel will give rise to "smooth-rotational symmetry" (i.e., order infinity) in the pure lipid membrane part of the vesicle. This assertion is corroborated by Piezo vesicles that appear circular in projection in EM images when viewed along Piezo's threefold axis. With this idea in mind, we generated the complete 3D shapes of Piezo vesicles from the oriented images as follows (Fig. 2). The midmembrane contour of oriented Piezo vesicle images was digitized manually and converted to a continuous, interpolated contour as shown. The two sides of the interpolated contour, east and west, were reflected about the long vesicle axis connecting the vesicle north



Fig. 1. Measuring Piezo vesicles. (A) Cryo-EM image of a single Piezo vesicle with the Piezo CED marked with a yellow pointer. (Scale bar, 25 nm.) (B) Schematic of Piezo vesicle data collection and tomographic image reconstruction. (B, Left) Tilt series images are collected on the detector while tilting the sample relative to the electron beam. (B, Center) A 3D tomogram is reconstructed from the tilt-series images. The gray background and yellow interior correspond to one plane of the tomographic reconstruction. Part of the reconstructed vesicle is shown in green, and the Piezo CED is colored yellow. The dashed white curve marks the intersection of the vesicle and a plane that includes the Piezo CED and the opposite "south" pole of the vesicle. (B, Right) The oriented Piezo vesicle image shows the Piezo vesicle contour cut out from the intersecting plane, with the Piezo CED marked with a yellow pointer. (C) Top-down view of the Piezo dome. The approximate position of the curved mid-bilayer surface of the Piezo dome is indicated in gray, and its intersection with the Piezo protein is indicated in cyan. The ribbon model of Piezo1 (PDB ID code 6B3R) is colored yellow.



Fig. 2. Quantifying Piezo vesicle profiles. Oriented Piezo vesicle image obtained by cryo-EM tomography (*Left*). To quantify the Piezo vesicle shape, we manually trace the midmembrane vesicle profile on the oriented Piezo vesicle image (cyan dots). (Scale bar, 26 nm.) (*Right, i*) We take the line of maximal length from Piezo's CED to the interpolated vesicle profile to correspond to the vesicle symmetry axis. Left-right averaging of the interpolated vesicle profile about the vesicle symmetry axis yields the symmetrized Piezo vesicle profile (red curve). (*Right, ii*) We determine the Piezo dome boundary (green dots) by integrating out a vesicle surface area equal to the Piezo dome area, $A_P \approx 450 \text{ nm}^2$, starting at the intersection of the vesicle symmetry axis and the Piezo dome (the vesicle north pole). (*Right, iii*) As inputs for the membrane elasticity theory of Piezo vesicle shape, we extract from the measured (symmetrized) vesicle profiles the free vesicle area A_F , the in-plane radius of the Piezo dome r_b , and the height of the Piezo dome boundary above the vesicle south pole opposite the Piezo dome h_b . For a given Piezo vesicle radius R_v , A_F is determined by $A_F = 4\pi R_V^2 - A_P$, while the values of r_b and h_b follow directly from the observed positions of the Piezo dome boundary and of the vesicle south pole, respectively. (See also SI Appendix, Figs. S1 and S2.)

and south poles. The sides were averaged to produce a symmetrized Piezo vesicle profile, which we call the Piezo vesicle profile, and which defines the 3D surface of the Piezo vesicle through rotation about the central axis connecting the north and south poles. The border between the Piezo dome and the vesicle free membrane was defined by equating the surface area of the membrane region surrounding the vesicle north pole to the membrane surface area contained within Piezo's approximate perimeter, A_P , estimated from the channel's atomic coordinates (Protein Data Bank [PDB] ID code 6B3R). This is the surface area of the Piezo dome, which includes the membranefilled space in between Piezo's extended arms. To account for uncertainty-both due to inherent experimental limitations and due to the approximations used here-in defining the border separating the threefold symmetric channel and the smooth-rotationally symmetric free membrane, in the accompanying paper, we consider variations in the Piezo dome area over the range from 410 to 490 nm² (14). We call the surface area of the vesicle membrane outside the Piezo dome border A_F , for "free" membrane area. We specify the size of a vesicle using the variable R_v , the radius of a hypothetical sphere comprising the Piezo dome plus the free membrane: $A_P + A_F = 4\pi R_v^2$.

Fig. 3 shows the 3D mid-bilayer surface of the free lipid membrane in an idealized Piezo vesicle (Fig. 3*A*) and its profile on a graph with the vesicle south pole at the origin (Fig. 3*B*). We denote the arclength along the vesicle profile by *s*, with s = 0 at the south pole and $s = s_b > 0$ at the boundary where the free membrane meets the Piezo dome. In the analysis to follow, the free membrane region of a Piezo vesicle profile is represented parametrically with the horizontal r(s) and vertical h(s) coordinates given as functions of arclength, *s*, as shown in Fig. 3*B*. The coordinate system is thereby oriented so that r(s) corresponds to the radial distance from the long vesicle axis to a particular point on the vesicle surface and h(s) corresponds to the radial.

Strategy to Predict Free Membrane Shape. Next, we focus our attention on predicting the shape of the free membrane: that is, the part of the vesicle corresponding to $0 \le s \le s_b$ in Fig. 3*B*. You might wonder, if we can simply measure the shape experimentally, why would we want to predict it? The answer has

two parts. First, our aim in this study is ultimately to deduce forces on Piezo from experimentally measured shapes of vesicles, which we pursue in the companion paper (14). This is possible only if we understand why the Piezo vesicles are shaped the way they are, and prediction from theory is our working definition of understanding. Second, a membrane elasticity theory has been developed and tested over the past fifty years to successfully account for the experimentally observed shapes in pure lipid membrane systems, such as giant unilamellar vesicles (19, 20). However, in our case, we are studying frozen small unilamellar vesicles with a protein channel incorporated into



Fig. 3. Membrane elasticity theory of Piezo vesicle shape. (A) 3D illustration of Piezo vesicles. The curved shape of the midmembrane surface of the Piezo dome (indicated in gray) deforms the free lipid bilayer membrane in Piezo vesicles (indicated in blue). (B) Schematic of a Piezo vesicle profile in the vesicle symmetry plane containing Piezo's central pore axis. We denote the arclength along the vesicle profile by s, with s = 0 at the vesicle south pole opposite the Piezo dome and $s = s_b > 0$ at the boundary (green dots) between the Piezo dome (gray curve) and the free lipid bilayer membrane outside the Piezo dome (blue curve). We represent the symmetry axis of the Piezo vesicle by the h axis, the (in-plane) radial coordinate perpendicular to the *h* axis by *r*, and the angle between the tangent to the vesicle profile and the *r* axis (in the direction of increasing *r*) by $\psi = \psi(s)$. For a given Piezo vesicle, the free membrane shape minimizing the Helfrich energy equation, Eq. 1, is determined by the values of three parameters, which are indicated in blue: the free membrane area A_F , the in-plane Piezo dome radius r_b , and the Piezo dome contact angle $\alpha = \pi - \psi(s_b)$. We have the cord length (radius of curvature) $R_P = \frac{r_b}{\sin \alpha}$ at the Piezo dome boundary. As illustrated by the triangle relating the differentials ds, dr, and dh, the shape variables satisfy the geometric relations $\dot{r} = \cos \psi$ and $\dot{h} = \sin \psi$.

them. Thus, we need to see how well the theory works in this circumstance.

Before presenting the mathematical approach to membrane shape prediction, we describe our basic strategy in words. We observe Piezo vesicles that contain one part protein and another part free lipid membrane. We have a theory for the shape energy of free lipid membranes (21). This theory gives us an energy equation for the free membrane that lets us put in the free membrane shape and get out an energy. Note that we do not have such a theory for the protein. In fact, this is ultimately what we seek. So, for the free membrane part, how do we begin? We consider the protein as an object in the vesicle that exerts specific geometric constraints on the free membrane (i.e., boundary conditions), which we determine from experiments. Then, we focus our attention on the free membrane part of the vesicle and ask, given the boundary conditions and free membrane area, what free membrane shape is associated with the lowest energy? The core assumption here is that the lowestenergy shape will correspond to the observed shape. In principle, one could try out all possible free membrane shapes that satisfy the boundary conditions by putting them into the energy equation and selecting the one that gives the lowest energy, but there are an infinite number of possible shapes. In practice, there is a mathematical approach (starting with the energy equation, the free membrane area, and the boundary conditions) to derive the lowest-energy shape (15-17). This is what we mean by predicting the free membrane shape from the energy equation.

The Energy Equation. The energy equation for the free membrane shape, due to Helfrich, is

$$G_M = \frac{K_b}{2} \int dA \, (c_1 + c_2)^2, \qquad [1]$$

where the constant K_b is the lipid bilayer bending modulus and c_1 and c_2 are the principal curvatures of the midmembrane surface, which are functions of position on the surface (21, 22). The integral is carried out over the entire free membrane vesicle surface. In words, every point on the free membrane surface is associated with a curvature number equal to the sum of the principal curvatures at that point. Eq. 1 says that the elastic energy to bend the entire free membrane from a plane into its curved shape, G_M , is computed by squaring the curvature number at each point, adding the values up, and multiplying the result by a constant, $\frac{1}{2}K_b$. Because the lipids used in the present study do not produce membranes with discernable spontaneous curvature, Eq. 1 does not involve a term for "intrinsic" curvature (i.e., its value is zero) (21). Nor does it have a term for the Gaussian curvature, which is directly proportional to $c_1 c_2$, because, as mandated by the Gauss-Bonnet theorem, the Gaussian curvature contribution to Eq. 1 can be expressed as a topological constant and a boundary term along the Piezo dome perimeter. This boundary term is cancelled by a corresponding boundary term due to the interior of the Piezo dome for an identical Gaussian curvature elastic modulus of the lipid bilayer inside and outside the Piezo dome and, thus, it should not affect the free membrane vesicle shape (22). Also, note that the Piezo vesicles were formed under conditions that preclude a pressure gradient across the membrane. Thus, vesicle volume is unconstrained, and the free membrane is under negligible tension.

To derive the lowest-energy free membrane shape (that is, to minimize the energy equation), we follow the path developed for the study of axisymmetric vesicles (19, 23–25). We begin

by expressing Eq. 1 using variables corresponding to the arclength parameterization of vesicle profiles (Fig. 3*B*). In addition to r(s) and h(s), for convenience we introduce a third variable $\psi(s)$, which is the angle between a line tangent to the vesicle profile and the *r* axis (Fig. 3*B*). The variables $\psi(s)$, r(s), and h(s), which we will call the shape variables, are related to each other through the geometric relations $\dot{r} = \cos \psi$ and $\dot{h} = \sin \psi$; it is understood that ψ , *r*, and *h* are functions of the arclength parameter *s*, and the dot notation means derivative with respect to *s* (Fig. 3*B*). Eq. 1 then becomes

$$G_M = \pi K_b \int_0^{s_b} ds \ r \left(\dot{\psi} + \frac{\sin \psi}{r} \right)^2, \qquad [2]$$

where $\dot{\psi}$ and $\frac{\sin\psi}{r}$ are the principal curvatures, c_1 and c_2 , in the arclength parameterization. When minimizing Eq. 2, certain general conditions must be met. First, the geometric relations $\dot{r} = \cos\psi$ and $\dot{h} = \sin\psi$ must be satisfied at every point on the free vesicle profile, and second, the total free membrane area must equal A_F , the value of which will be determined from experiment. These conditions are enforced by introducing constraints into Eq. 2 through the mathematical method of Lagrange multipliers, to produce a constrained energy equation,

$$\widetilde{G}_{M} = \int_{0}^{s_{b}} ds \left[\pi K_{b} r \left(\dot{\psi} + \frac{\sin \psi}{r} \right)^{2} + 2\pi \lambda_{a} r + \lambda_{r}(s) \left(\dot{r} - \cos \psi \right) + \lambda_{b}(s) \left(\dot{h} - \sin \psi \right) \right] - \lambda_{a} A_{F}, \quad [3]$$

which is a function of the Lagrange multipliers $\lambda_r(s)$, $\lambda_b(s)$, and λ_a as well as the shape variables and (or) their derivatives with respect to s. The coefficients $\lambda_r(s)$ and $\lambda_b(s)$ in Eq. **3** associated with the geometric constraints depend on s, and λ_a is a constant coefficient for the term $2\pi r$, which integrates to give the free membrane area and must satisfy $2\pi \int_0^{s_b} ds r = A_F$. The utility of the Lagrange multipliers is this: the values or functional forms of the Lagrange multipliers are selected so that the constraints are met after Eq. **3** is minimized (16). But notably, when the constraints are met (i.e., when $\dot{r} = \cos\psi$, $\dot{h} = \sin\psi$, and $2\pi \int_0^{s_b} ds r = A_F$), the terms multiplied by the Lagrange multipliers vanish. This means that minimization of Eq. **3** will yield what we seek: the minimum of our original energy equation, Eq. **2** or **1**, for a vesicle in which the constraints are satisfied.

The Hamilton Equations. For what comes next, we rewrite Eq. 3 as

$$\widetilde{G}_M = \int_0^{s_b} ds \ L(\psi, \dot{\psi}, r, \dot{r}, \dot{b}),$$
[4]

where *L*, called the Lagrangian, stands for the integrand in Eq. **3** and, without loss of generality, we omit the constant $-\lambda_a A_F$ in Eq. **3** (17). If the right-hand side of Eq. **4** was an ordinary function, say f(x, y, ...), at a minimum point its value would not change when $x \to x + \delta x$, $y \to y + \delta y$, ..., and we would find the values $x = x_{\min}$, $y = y_{\min}$, ... associated with the minimum of f(x, y, ...)by solving the derivative equations $\frac{\partial f(x, y, ...)}{\partial x} = 0$, $\frac{\partial f(x, y, ...)}{\partial y} = 0$, ... for x, y, ... Similarly, the minimum of Eq. **4** occurs when small changes in the shape variables (i.e., $\psi \to \psi + \delta \psi$, $r \to r + \delta r$, and $h \to h + \delta h$) do not change the energy. By analogy to solving the derivative equations of an ordinary function, we could find the minimum energy form of the shape variables in Eq. **4** by solving "functional derivative" equations called Euler–Lagrange equations, $\frac{\partial L}{\partial \psi} - \frac{d}{ds} \frac{\partial L}{\partial \psi} = 0$, $\frac{\partial L}{\partial r} - \frac{d}{ds} \frac{\partial L}{\partial r} = 0$, and $\frac{\partial L}{\partial b} - \frac{d}{ds} \frac{\partial L}{\partial b} = 0$, subject to specific boundary conditions at s = 0 and $s = s_b$ (17). As a mathematically equivalent alternative to solving the Euler–Lagrange equations directly, we derive the minimum energy form of the shape variables by solving the Hamilton equations associated with Eq. 4 (18). The Hamilton equations facilitate the numerical calculation of the free membrane shape of Piezo vesicles, and have the advantage that the boundary conditions encoding the key physical properties underlying Piezo vesicle shape can be imposed in a particularly transparent manner.

As outlined in *SI Appendix*, section S2, the Hamilton equations associated with Eq. 4 are derived by switching from $\dot{\psi}$, \dot{r} , and \dot{h} to $p_{\psi}(s) \equiv \frac{\partial I}{\partial \dot{\psi}} = 2\pi K_b r \left(\dot{\psi} + \frac{\sin\psi}{r}\right)$, $p_r(s) \equiv \frac{\partial I}{\partial \dot{r}} = \lambda_r$, and $p_b(s) \equiv \frac{\partial I}{\partial \dot{b}} = \lambda_h$ in the Euler–Lagrange equations through the Legendre transformation $H \equiv p_{\psi}\dot{\psi} + p_r\dot{r} + p_b\dot{h} - L$, where *H* is known as the Hamiltonian (17, 18). p_{ψ} , p_r , and p_b are called generalized momenta, in reference to the physical meaning that these terms carry when describing dynamical systems in which the independent variable is time instead of arclength (17, 18). Following this Legendre transformation, rather than having one Euler–Lagrange equation for each shape variable ψ , *r*, and *h*, we have two first-order differential equations for each: one describing how its generalized momentum changes with *s*. The Hamilton equations associated with Eq. 4 are

$$\dot{\psi} = \frac{1}{r} \left(\frac{p_{\psi}}{2\pi K_b} - \sin \psi \right),$$
 [5]

$$\dot{r} = \cos\psi,$$
 [6]

$$\dot{h} = \sin \psi,$$
 [7]

$$\dot{p}_{\psi} = \left(\frac{p_{\psi}}{r} - p_{b}\right)\cos\psi + p_{r}\sin\psi, \qquad [8]$$

$$\dot{p}_r = \frac{p_{\psi}}{r^2} \left(\frac{p_{\psi}}{4\pi K_b} - \sin \psi \right) + 2\pi \lambda_a, \qquad [9]$$

$$\dot{p}_{h} = 0.$$
 [10]

Note that the Lagrange multipliers $\lambda_r(s)$ and $\lambda_b(s)$ were eliminated in the derivation of the Hamilton equations, but that the associated geometric constraints on the free membrane are preserved in Eqs. **6** and **7**. The Lagrange multiplier for the free membrane area, λ_a , enters the Hamilton equations through Eq. **9**; it is a parameter whose value must be determined from experiments through measurement of A_F . To apply the specific free membrane area constraint to each vesicle under study, we add to the system of Eqs. **5–10** a differential equation for the membrane area up to an arclength *s* from the south pole, A(s):

$$A = 2\pi r.$$
 [11]

To predict the free membrane shapes of Piezo vesicles, we solve the six Hamilton equations, Eqs. 5–10, and the area equation, Eq. 11, subject to the boundary conditions at s = 0 and $s = s_b$ associated with Piezo vesicles.

Boundary Conditions and Input Parameters. For a Piezo vesicle with the south pole at the origin of the r - h parametric plot, we have, from geometry, $\psi(s) = 0$ and r(s) = 0 at s = 0 (Fig. 3*B*). As shown in *SI Appendix*, section S3, $\psi(0) = 0$ and r(0) = 0 imply that $p_{\psi}(0) = 0$. Since *h* does not appear explicitly in *L*, we are free to define h(0) = 0. Furthermore, the projected perimeter radius of the Piezo dome is given by $r(s_b) = r_b$

(Fig. 3B). We assume that, at $s = s_b$, the free membrane and Piezo dome surfaces are tangent to each other, because the lipid membrane connects smoothly across the Piezo dome boundary. Note that, at $s = s_b$, a cord of length $R_P = \frac{r_b}{\sin \alpha}$, drawn perpendicular to the membrane so that it connects the long vesicle axis to the Piezo dome boundary, forms an angle α with the *h* axis (Fig. 3*B*). We call α the contact angle. Therefore, from geometry, $\psi(s_b) = \pi - \alpha$. Note that the values of $\psi(s_b)$ and $r(s_b)$ are fixed in a Piezo vesicle by the shape of the Piezo dome. By contrast, $h(s_b)$ is not fixed by Piezo because the length $h(s_b) - h(0)$ can freely vary when G_M is minimized [i.e., constrained physically by only the fixed values of $\psi(s_b)$, $r(s_b)$, and the free vesicle area, the vesicle could become shorter or longer, so its height is determined by the minimization of G_M]. Thus, we have what is called a zero-force or natural boundary condition on h(s) at $s = s_b$ (15, 17). As shown in *SI Appendix*, section S3, this means that $p_b(s_b) = 0$. However, we already saw that $\dot{p}_h(s) = 0$, Eq. 10, which is to say that $p_h(s)$ is unchanged over $0 \le s \le s_b$, and therefore $p_b(0) = 0$.

Another important aspect of the minimization of Eq. 4 is that the integration domain length (i.e., the arclength s_b) is not fixed (23). Beginning at the south pole, the free membrane profile could take a circuitous path in getting to the edge of the Piezo dome as long as the boundary conditions already mentioned are satisfied and the free membrane area equals A_F . As shown in *SI Appendix*, section S3, the variable integration domain length, together with the fact that *s* does not appear explicitly in *L*, leads to $p_r(0) = 0$. For the membrane area, we have A(0) = 0 and $A(s_b) = A_F$. Thus, in total, at s = 0 we have the seven "initial" conditions

$$\psi(0) = 0, r(0) = 0, h(0) = 0, p_{\psi}(0) = 0,$$

 $p_r(0) = 0, p_h(0) = 0, A(0) = 0,$
[12]

with which to solve the six Hamilton equations, Eqs. 5-10, and the area equation, Eq. 11. However, these equations are associated with three parameters that are a priori unknown. We already saw that λ_a , the Lagrange multiplier on the free membrane area, and s_b , the upper limit of the integration domain, must be determined as part of the minimization of G_M . In addition, the vesicle profile curvature at the south pole, $\dot{\psi}(0)$, must be entered to solve Eqs. 5-11. To see this, note that the right-hand side of Eq. 5 evaluates to $\dot{\psi}(0)$ as $s \to 0$, and thus, $\dot{\psi}(0)$ must be specified to make Eq. 5 meaningful at s = 0. While $\dot{\psi}(0)$ could in principle be estimated from the observed vesicle profiles, it is difficult to measure curvature accurately. For this reason, we set $\dot{\psi}(0) \equiv u_0$ (24) with the value of u_0 , along with the values of λ_a and s_b , to be determined from experimental constraints on the shape of each measured Piezo vesicle.

To solve the six Hamilton equations and the area equation, Eqs. 5–11, associated with the three unknown parameters λ_a , s_b , and u_0 , we supplement the seven initial conditions in Eq. 12 with three endpoint boundary conditions measured from the Piezo vesicle profiles. As seen above, $A(s_b)$ must equal the measured free membrane area A_F , $r(s_b)$ must equal the measured in-plane Piezo dome radius r_b , and $\psi(s_b)$ must equal π minus the Piezo dome contact angle α . In practice, because obtaining the value of α involves taking the ratio of derivatives of the observed vesicle profile, we cannot measure α as accurately as a position on the vesicle profile. Thus, we use the measured position of $h(s_b) = h_b$ in place of $\psi(s_b) = \pi - \alpha$ when predicting the free vesicle shapes. Using an experimental value for $h(s_b)$ in this way does not violate the zero-force boundary condition on h(s) at $s = s_b$, which led to the initial condition $p_b(0) = 0$ in Eq. 12. Employing a so-called shooting method (*SI Appendix*, section S4), we repeatedly solve Eqs. 5–11 subject to Eq. 12 to numerically adjust the values of λ_a , s_b , and u_0 so as to satisfy the experimental constraints

$$h(s_b) = h_b, \ r(s_b) = r_b, \ A(s_b) = A_F$$
 [13]

for each measured Piezo vesicle. Thus, for each Piezo vesicle, we use the measured values of the three experimental input parameters h_b (in place of α), r_b , and A_F to predict the free membrane shape through minimization of the Helfrich energy equation, without any free parameters.

Free Membrane Shapes. Tomograms were collected on seven Piezo vesicles ranging in radius, R_{ν} , from 12.1 to 36.2 nm. The vesicle profiles are shown in red (Fig. 4). The dome boundary, in green, marks the point where the Piezo dome meets the free membrane as defined by a Piezo dome area $A_P = 450$ nm². The coordinates of the dome boundary, r_b and h_b , together with the free membrane area, A_F , were used to solve the Hamilton equations and the area equation, Eqs. **5–11**, subject to the initial conditions in Eq. **12** with the constraints in Eq. **13**. The calculated free membrane profiles, obtained with no free parameters, are shown in blue. The calculated profiles conform to the measured shapes of the Piezo vesicles, and thus, the

Helfrich energy equation appears to successfully predict the free membrane shape.

There are several shortcomings of the approach used here that may limit the agreement between experimental and theoretical results. In particular, the predicted free vesicle shapes depend crucially on the measured values of h_b , r_b , and A_F , which experiments can only yield approximately. Furthermore, the tomograms of Piezo vesicles do not produce a continuous vesicle surface around the long vesicle axis. As a result, we expect the measured, oriented Piezo vesicle images to be tilted slightly out of the planes defined by the vesicle cross-sections and hence, to correspond only approximately to the predicted vesicle cross-sections. Moreover, the location of the Piezo dome boundary can only be defined approximately in experiments, and the threefold symmetry of the Piezo protein is expected to induce (slight) variations in the Piezo dome boundary conditions around the long vesicle axis. In contrast, our theoretical description of Piezo vesicles assumes perfect, smooth rotational symmetry of the free membrane shape. Considering these limitations, as well as the inevitable errors incurred when defining the mid-bilayer surface in the oriented Piezo vesicle images, the agreement between the predicted and measured free vesicle shapes in Fig. 4 appears to be remarkably good.

Up to this point, we have described the mathematical approach taken as one to find the minimum energy shape of the free membrane subject to geometric constraints arising from the



Fig. 4. Quantitative prediction of Piezo vesicle shape. Symmetrized (measured) Piezo vesicle profiles (red curves) and predicted free membrane profiles (blue curves) for Piezo vesicles with radii (1) $R_v \approx 12.1$ nm, (2) $R_v \approx 13.7$ nm, (3) $R_v \approx 24.7$ nm, (4) $R_v \approx 25.1$ nm, (5) $R_v \approx 27.6$ nm, (6), $R_v \approx 35.0$ nm, and (7) $R_v \approx 36.2$ nm. The approximate rmsds of the measured and predicted vesicle profiles are given by (1) 0.29, (2) 0.35, (3) 0.28, (4) 0.49, (5) 0.24, (6) 0.25, and (7) 0.44 nm. To calculate the blue curves we used, as inputs for the membrane elasticity theory of Piezo vesicle shape, the values of A_F , r_b , and h_b measured for each vesicle. The calculated free membrane profiles were obtained from the Helfrich energy equation, Eq. 1, without any free parameters. The oriented Piezo vesicle image in Fig. 2, *Left* corresponds to vesicle 3. (Scale bars, 5 nm.)

Piezo dome shape and the vesicle size. In fact, the approach potentially identifies any shape that renders the free membrane energy unchanged under small perturbations of the shape variables (15, 17). This could include a local instead of a global minimum of the Helfrich energy equation, a maximum, or any other stationary point. However, because our mathematical solutions match the experimentally observed shapes of Piezo vesicles that self-assembled under equilibrium conditions, it seems likely that the predicted free vesicle shapes indeed correspond to the global energy minimum of the free membrane shape.

Much consideration has been given to the concept that certain membrane proteins might distort the shape of their surrounding lipid bilayer membrane, creating a membrane footprint (6-12). Based on the membrane elasticity theory of bilayer-protein interactions, it has been proposed that the membrane footprint of proteins may affect membrane protein function, assist with membrane remodeling, yield bilayer-mediated protein interactions, and induce protein cooperativity. But it has been challenging to measure quantitatively and predict how proteins deform bilayer membranes, and hence to directly compare predicted and observed membrane footprints. The free membrane shapes of Piezo vesicles described here represent a direct measure of a protein's membrane footprint, in the context of lipid bilayer vesicles, and show that membrane elasticity theory can be used to predict Piezo's membrane footprint, thus linking membrane and protein shape.

Piezo Dome Shapes. We next turn our attention to the parts of Piezo vesicles that contain Piezo. The Piezo dome in profile, shown in red between the green dome boundary marks (Figs. 4 and 5*A*), has a somewhat irregular shape. This is not surprising since the Piezo dome contains the Piezo protein that is viewed perpendicular to its randomly oriented threefold axis. Shape irregularities notwithstanding, systematic differences in the Piezo dome shape as a function of vesicle radius, R_v , are evident: in smaller vesicles, the Piezo dome is more curved, and in larger vesicles, it is flatter.

Is there a simple way to understand the observed Piezo dome shapes? We adapted the theory developed above for the free membrane outside the Piezo dome to the interior of the Piezo dome. The objective here is to determine whether the same physical principle governing the free membrane shape outside the Piezo dome-namely, minimization of the Helfrich energy equation-can also be used to describe the Piezo dome shape. Specifically, we minimized Eq. 1 through the solution of Eqs. 5-11 with the initial conditions in Eq. 12 but now employing, as input parameter values, $A(s_b) = A_P$ and the values of α and r_b associated with the predicted free membrane shapes in Fig. 4. Note that s = 0 corresponds here to the vesicle north pole rather than the vesicle south pole. The resulting Piezo dome shapes, shown in blue, are seen to approximate the observed Piezo dome shapes (Fig. 5A). Curiously, these calculations suggest that the interior of the Piezo dome behaves somewhat like a flexible membrane, even though the Piezo dome contains the Piezo protein.

We also want to point out that the overall shape of the Piezo dome is not very far from a spherical cap (Fig. 5*A*). The shape of a spherical cap is conveniently specified through the cap radius of curvature R_P and the cap area A_{cap} (26). From the geometry of spherical caps, we see that R_P and A_{cap} can be used to define the Piezo dome properties affecting membrane shape deformations through $\alpha = \cos^{-1} \left(1 - \frac{A_{cap}}{2\pi R_P^2}\right)$ and $r_b = \left[\frac{A_{cap}}{\pi} \left(1 - \frac{A_{cap}}{4\pi R_P^2}\right)\right]^{1/2}$ (Fig. 5*B*). Previous cryo-EM and HS-AFM experiments suggest



Fig. 5. Coupling between Piezo and membrane shape. (A) Symmetrized (measured) Piezo dome profiles in Fig. 4 (red curves), Piezo dome shapes minimizing the Helfrich energy equation, Eq. 1, subject to $A(s_b) = A_P = 450 \text{ nm}^2$ and the values of α and r_b associated with the predicted free membrane shapes in Fig. 4 (blue curves), and Piezo dome profiles corresponding to a spherical cap with area $A_{cap} = A_P$ and the in-plane Piezo dome radius r_b associated with the predicted free membrane shapes in Fig. 4 (gray curves). As in Fig. 4, the Piezo dome profiles correspond to Piezo vesicles with radii (A, 1) $R_v \approx 12.1$ nm, (A, 2) $R_v \approx 13.7$ nm, (A, 3) $R_v \approx 24.7$ nm, (A, 4) $R_v \approx 25.1$ nm, (A, 5) $R_v \approx 27.6$ nm, (A, 6) $R_v \approx 35.0$ nm, and (A, 7) $R_v \approx 36.2$ nm. (Scale bars, 5 nm.) (B) Schematic of the spherical cap model of the Piezo dome. The in-plane cap radius r_b and the cap angle α or, alternatively, the cap radius of curvature R_P and the cap area A_{cap} completely define the geometric properties of the spherical cap. (C) Piezo dome contact angle α and (D) in-plane Piezo dome radius r_b vs. radius of curvature R_p for the predicted free membrane shapes of the Piezo vesicles in Fig. 4 (points) and for a spherical dome (cap) with fixed area $A_{cap} = A_P$ (solid curves). For the Piezo vesicles, the values of α and r_b associated with the predicted free membrane shapes in Fig. 4 determine, as illustrated in Fig. 3*B*, the radius of curvature at the Piezo dome boundary through $R_P = \frac{r_b}{\sin \alpha}$. For the spherical dome, the calculated curves follow from the geometric relations $\alpha = \cos^{-1} \left(1 - \frac{A_{cap}}{2\pi R_p^2} \right)$ and $r_b = \left[\frac{A_{cap}}{\pi} \left(1 - \frac{A_{cap}}{4\pi R_p^2} \right) \right]^{1/2}$ associated with spherical caps (26).

setting $A_{cap} = A_P$ in these equations, so that the shape of the Piezo dome is approximated by a spherical cap with fixed area A_P and variable radius of curvature R_P (3, 13). When we thus graph α and r_b as a function of R_P for the seven vesicles in Fig. 4, we find that α and r_b indeed vary as if the Piezo dome approximates a spherical cap of fixed area (Fig. 5 C and D), suggesting that the features of the Piezo dome affecting Piezo's membrane footprint can, approximately, be captured by a single parameter, the Piezo dome radius of curvature R_P . We use this result in our companion paper to study the mechanical response of the Piezo dome to the forces exerted by the surrounding lipid bilayer membrane (14).

Discussion

We have shown here that the Helfrich energy equation accurately predicts the observed shapes of the free membrane in Piezo vesicles over a range of vesicle sizes, without any free parameters. While it is not possible to directly measure Piezo's membrane footprint in infinite, asymptotically planar membranes, our results suggest that membrane elasticity theory accurately captures protein-induced lipid bilayer shape deformations from a length scale corresponding to molecular sizes to at least a length scale ~ 100 nm.

Continuum elasticity theory implies that the free membrane shape of a particular Piezo vesicle follows from the values of three input parameters, which can all be estimated directly from experiments. In particular, we found here that the values of the free membrane area A_F , the projected radius of the Piezo dome r_b , and the height of the Piezo dome boundary above the opposite vesicle pole h_b can be used to predict successfully the free vesicle shape. Other choices for the experimental input parameters could be used and give similar results.

Furthermore, we have shown here how Piezo changes its shape as a function of vesicle size, with the Piezo dome becoming less curved as the vesicle size is increased (13). We find that both the geometric properties of the Piezo dome at its boundary and the shape of the Piezo dome conform approximately to a spherical cap with fixed area. Minimization of the membrane bending energy within the Piezo dome boundary also yields shapes close to the measured Piezo dome shapes. This seems surprising, given that the Piezo dome is not a uniform elastic membrane but contains the Piezo protein, and raises the fascinating prospect that the Piezo protein may be similarly flexible as the surrounding lipid bilayer membrane.

Instead of following the theoretical approach employed here, we could have substituted the observed shapes of the (smoothed) free vesicle membrane into the Helfrich energy equation and hence, deduced the membrane deformation energy associated with the observed free vesicle shapes. While this procedure would yield similar results for the membrane deformation energy, such an approach would be unable to predict the free membrane shape. Importantly, the agreement between the predicted and measured free membrane shapes suggests that the measured shapes correspond to minima of the Helfrich energy equation. With such a physical understanding of Piezo vesicle shape in hand, we can then ask: what would be the free membrane shape if the Piezo dome geometry was perturbed, and what would be the corresponding energy of the free membrane? As we show in the companion paper, the answer to this question allows calculation of the forces exerted by the Piezo dome on the surrounding lipid membrane and, conversely, on the Piezo dome by the surrounding lipid membrane (14). From such a quantitative understanding of bilayer-protein interactions one can then deduce, based on the observed vesicle shapes, elastic properties of the Piezo dome underlying its mechanosensory function.

The simple version of the Helfrich energy equation employed here only involves a single term-the membrane bending energy-and only a single physical parameter-the bilayer bending rigidity, K_b —that can be measured directly in experiments. The predicted free vesicle shapes are independent of the value of K_b , and follow directly from minimization of the square of the sum of the two principal membrane curvatures across the free vesicle surface. The agreement between the predicted and measured free vesicle profiles suggests that, within experimental uncertainty, it is not necessary to account for effects other than those captured by minimization of the membrane bending energy, such as thermal fluctuations, bilayer coupling, or lipid tilt, to account for the observed Piezo vesicle shapes. Inclusion of such additional effects could further improve the agreement between observed and predicted vesicle shapes. Our results also suggest that the rapid freezing of Piezo vesicles for cryo-EM does not induce substantial perturbations of the free membrane shape.

The agreement between predicted and measured free membrane shapes may thus be taken as an indication that cryo-EM can be used not only to accurately measure protein structures, but also to quantify the 3D shapes of lipid bilayers deformed by membrane proteins.

Materials and Methods

Proteoliposome Sample Preparation. Full-length mouse Piezo1 protein was expressed using the BacMam method as previously described (3). mPiezo1 was purified with a 6xHis tag at the C-terminus by cobalt affinity chromatography and size exclusion chromatography. Proteoliposomes (Piezo vesicles) with the lipid composition POPC:DOPS:cholesterol at a ratio of 8:1:1 (wt:wt:wt) were prepared as previously described (13). To enrich the proteoliposomes for cryoe-lectron tomography, the liposome sample was subjected to cobalt affinity chromatography following reconstitution.

Freshly prepared proteoliposome samples were supplemented with 3 mM fluorinated fos-choline-8 (FFC-8) immediately before freezing. C-flat 200 mesh gold R1.2/1.3 holey carbon grids were glow-discharged prior to use. A first drop of 3.5 μ L of proteoliposome sample was applied to the grid, incubated for 15 to 60 s, and then manually removed using a filter paper. Then, a second drop of 3.5 μ L of proteoliposome sample was added, incubated for 15 s, and then blotted once for 1 s with -1 force and plunged into liquid ethane using a Vitrobot Mark IV (FEI Company, Hillsboro, Oregon) operated at 22 °C and 100% humidity. The grids were then stored in liquid nitrogen until imaging.

Tilt-Series Data Collection. Tilt series were collected using a Titan Krios (Thermo Fisher) with a Gatan Bioquantum energy filter (Gatan) and a direct detector Gatan K3 (Gatan) at 300 keV. Data were collected symmetrically with a tilt range of -42° to 42° in 3° increments using SerialEM with 400-ms frames for each tilt image at a nominal defocus of 4 μ m. The total dose per tilt series collected was 103 e⁻/Å², with dose rates of approximately 20 e⁻/pixel per second at a pixel size of 2.6 Å. Full-frame alignment was performed using Motion-Cor2 (27).

Tilt-Series Alignment. Tilt series were aligned using Appion-Protomo (28–30). Tilt series were coarsely aligned, manually aligned, and then refined using a set of alignment thicknesses. The best-aligned iteration was reconstructed for visual analysis using Tomo3D simultaneous iterative reconstruction technique (SIRT) (31, 32) after dose compensation using a previously described method (33). Contrast transfer function (CTF) correction was not performed.

Tomogram Image Processing. The reconstructed tomograms were visualized and analyzed in the 3DMOD. The coordinate of the Piezo1 C-terminal extracellular domain (CED) in every liposome was determined by fitting the PDB model of mPiezo1 (PDB ID code 6B3R). The distance between the CED and the outer membrane leaflet of the liposome was measured in the 3D reconstruction. The region of membrane associated with the longest distance away from the CED is defined as the Piezo vesicle south pole. The maximum projection plane (oriented Piezo vesicle image) was determined by drawing a plane intersecting the CED and the south pole.

The slice images of vesicles corresponding to the maximum projection plane were processed using an in-house script on the MATLAB R2021a platform (34) to annotate the mid-bilayer surface of the entire vesicle. Circles of diameter of 5.7 nm, the average distance between phospholipid head-group layers of our sample from previous measurements (13), were placed by visual inspection along the vesicle membrane defined by the head-group layers. The (*x*, *y*) coordinates of the centers of these circles defined the midmembrane contour of the vesicle within the maximum projection plane. A file of (*x*, *y*) coordinates was used to generate a vesicle profile using the Interpolation command (with InterpolationOrder \rightarrow 3) in Mathematica (35).

Prediction of Piezo Vesicle Shape. To predict the free membrane shapes of Piezo vesicles in Fig. 4, we numerically solved Eqs. **5-11** subject to Eqs. **12** and **13** using Mathematica (35). We proceeded similarly in Fig. 5*A* to calculate the Piezo dome shapes through minimization of Eq. **1**. A detailed description of mathematical derivations and methods can be found in *SI Appendix*, sections S2–S4.

Data, Materials, and Software Availability. The tomograms of Piezo vesicles are deposited in the EMDataBank (accession codes EMD-27569 (36) [vesicle 1], EMD-27571 (37) [vesicles 2 and 4], EMD-27568 (38) [vesicle 3], EMD-27567 (39) [vesicles 5 and 6], and EMD-27570 (40) [vesicle 7]). All other data are included in the manuscript and/or supporting information.

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