# Influence of membrane-cortex linkers on the extrusion of membrane tubes

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The cell membrane is an inhomogeneous system composed of phospholipids, sterols, carbohydrates, and proteins that can be directly attached to underlying cytoskeleton. The protein linkers between the membrane and the cytoskeleton are believed to have a profound effect on the mechanical properties of the cell membrane and its ability to reshape. Here we investigate the role of membranecortex linkers on the extrusion of membrane tubes using computer simulations and experiments. In simulations we find that the force for tube extrusion has a non-linear dependence on the density of membrane-cortex attachments: at a wide range of low and intermediate linker densities the force is not significantly influenced by the presence of the membrane-cortex attachments and resembles that of the bare membrane. For large concentrations of linkers however the force substantially increases compared to the bare membrane. In both cases the linkers provided membrane tubes with increased stability against coalescence. We then pulled tubes from HEK cells using optical-tweezers for varying expression levels of the membrane-cortex attachment protein Ezrin. In line with simulations, we observed that overexpression of Ezrin led to an increased extrusion force, while Ezrin depletion had negligible effect on the force. Our results shed light on the importance of local protein rearrangements for membrane reshaping at the nanoscopic scales.

### I. INTRODUCTION

Biological membranes can dynamically change their shape as a response to an external mechanical stress. The application of a localised force to a membrane leads to the formation of membrane tubes or tethers which are thin elongated cylindrical structures [1]. Membrane tubes are involved in a wide variety of cellular processes such as motility, mechanosensing, signalling, inter-cellular and intra-cellular trafficking [2–4]. Tubes can be artificially extracted through hydrodynamic flow experiments, atomic-force microscopy, optical or magnetic traps, both from reconstituted vesicles as well as from live cells [5-11]. In the case of giant vesicles and bare membranes the force-extension curve of the extrusion process has been well characterised both in theoretical models and in experiments [12–18]. The shape of the tube and characteristic extrusion force were used to directly infer the mechanical properties of the bare membrane, where the force was found to

depend on the membrane bending rigidity  $\kappa$  and surface tension  $\gamma$  as  $F \sim \sqrt{\kappa \gamma}$ , while the radius of the tube scaled as  $R \sim \sqrt{\kappa/\gamma}$ .

Plasma membrane of eukaryotic cells is connected to the underlying cytoskeleton via linking proteins such as ERM (Ezrin, Radixin and Moesin) and myosin 1 proteins [19, 20]. The influence of these linkers on the force required to extrude a tube is still a source of discussion. The attachment to the actomyosin skeleton has been suggested to play a role in the extension of membrane tethers, by providing resistance to the flow of the membrane towards the tube [21]. It was also shown that reducing the attachment of the plasma membrane to the actomyosin cortex, via myosin 1b or Ezrin depletion, reduces the static force required to extrude a tether in zebrafish progenitor cells [22]. A similar result was obtained with myosin 1g in lymphocytes [23]. Furthermore, Ezrin phosphorylation that increases membrane-cortex attachment was shown to increase the tether force in lymphoblasts [24].

Taken together, this data has been interpreted as evidence that membrane-cortex attachments effectively change the membrane tension. In this

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FIG. 1: The membrane is extruded into a tubular structure. (a) The membrane is modelled as a uniform thin sheet made of a single layer of phospholipids (grey beads). The membrane is pulled at a central bead and a tubular structure is extruded. b) The force-extension curve for a homogeneous membrane (in the absence of any cortex linkers) shows a peak at an extension of approximately  $l = 50\sigma$ , which corresponds to a transition in the protrusion's aspect from a truncated cone to a thin elongated tube ( $\sigma$  is the simulation unit length that corresponds to ~ 10nm). The curve eventually reaches a plateau after which the extrusion proceeds at the same force. The shaded area represents the standard error for measurements from 10 different simulation seeds.

interpretation the tension is increased by the strength of the membrane-cortex adhesion per unit area, which in turn also increases the force needed to extrude a membrane protrusion [25]. However, alternative pictures have also been put forward, in which the membrane flows around the membrane-cytoskeleton attachments into the tether, such that the attachments do not contribute to the extrusion force, resulting in a force that resembles that of the bare membrane [26, 27].

Here we use minimal coarse-grained computer simulations to study the role of the cytosleketonmembrane attachments on the extrusion of membrane tubes in isolation from other effects that might be present in cells. Our simulations do not explicitly assume driving forces, but only assume the ingredients - a simple model of a fluid membrane that is sparsely attached to the underlying surface. We then carry out computational experiments where we extrude protrusions and measure the membrane response for a wide range of membrane-cortex attachment densities. We complement our observations with optical-tweezersbased tether pulling on HEK293T cells adhering on a substrate to measure the impact of the ERM protein Ezrin on membrane mechanics.

Our results indicate that the two previously polarised views can in fact co-exist: we find that at low membrane-cortex linker densities the extrusion force is not influenced by the cortex attachments and is equal to that of the bare membrane, while at high attachment densities the force strongly increases compared to the bare membrane. At low linker densities the membrane flows around the linkers and the linkers avoid entering the tube to minimise the energetic cost of extrusion, hence the membrane-cortex attachment does not contribute to the force. At high linker densities, however, the linkers cannot avoid entering the tube composition any more and necessarily start detaching, causing the substantial increase in the extrusion force accompanied by a slow tube relaxation dynamics. In agreement with the simulations findings, our experiments show that for cells that display moderate level of cortex-linker attachments, depleting ERM protein Ezrin has negligible influence on the extrusion force but impedes the tube relaxation, while overexpressing Ezrin leads to a significant increase in the force. Our results point to the importance of local effects – the redistribution of the membrane-cortex attachments around the protrusion – when mechanically probing cellular membranes at the nanoscale.

### II. RESULTS

The force profile for the bare membrane. We use a Monte Carlo model in which the fluid membrane is described as a dynamically triangulated elastic sheet, while the cytoskeletal linkers are modelled by attaching some of the membrane sheet vertices to the surface underneath it by finite harmonic springs. This model reproduces well the mechanical properties of biological membranes, such as bending rigidity and fluidity, but does not take into account the explicit bilayer structure and the details of the membrane and protein structures. We also repeated our key results using a self-assembled (non-triangulated) one-particle-thick membrane model within molecular dynamics simulations, as discussed in SI.

We first make sure that our model reproduces previously measured force-extension profiles for non-attached membranes. To that end we apply a vertical pulling force on a small membrane patch



FIG. 2: Force-extension curve in the presence of membrane linkers. (a) Side-view of a tubular structure pulled in the presence of membrane linkers (magenta beads). The linkers are restrained on the vertical plane with a harmonic potential. (b) Force extension curves at small and intermediate linker densities are not significantly affected by the presence of linkers. The peak and plateau forces remain constant independent of protein density (legend indicates the protein number density,  $\sigma$  is the simulation unit of length). c) The force on the tube tip measured for the target extension of  $60\sigma$ , averaged between  $1 \cdot 10^6$  and  $2 \cdot 10^6$  MC steps. At high expression levels of the proteins the force increases substantially. Inset shows the fraction of proteins in the composition of the tube. The error bars and shaded regions in (b)-(d) represent the standard error of the measurements over 5 different simulation seeds. d) The tube radius decreases as the tube is extruded. The linker density modulates the radius.

in the membrane centre. This leads, at first, to the extrusion of a cone-like structure (Fig. 1a). The force-extension profile shows a linear elastic response until a critical threshold is reached. The force then drops by around 8% and the protrusion undergoes a shape transition from the catenoid to a thin tubular structure as shown in Fig. 1b and the extrusion force eventually reaches a plateau. This extension profile is consistent with experimental and theoretical studies of tether extrusion on free membranes that show the familiar profile of a barrier followed by a plateau [12]. It has been shown that the height of the barrier is influenced by various factors such as membrane composition, speed of pulling and the area of contact between the membrane and the pulling object, while the plateau is typically determined only by the membrane bending rigidity and surface tension [14, 27–29]. We next explored how this forceextension curve is influenced by the addition of proteins linking the cytoskeleton to the membrane.

Extrusion force has a non-linear dependence on the density of cortex linkers. Cytoskeletal attachments, or linkers, were introduced into the system by connecting a fraction of membrane beads to a virtual plane beneath the membrane by a generic, finite harmonic, potential (Fig. 2a). Such a bond can be broken if sufficiently extended. Although some membrane-cortex linkers can exhibit more complex behaviour such as catchbond [30], this simple generic form of binding is believed to represent well the ERM proteins [31].

As the membrane tube is extruded from such a pinned membrane, we observe that the linkers tend to disperse away from the tube since there is a high energetic cost associated with opposing the pulling force and entering the tether. This causes the force required for extrusion not to be significantly affected by the linker attachments for low protein fractions, as shown in Fig. 2b. In the early stages of the tube formation, the force



FIG. 3: The linkers are excluded from the tube at intermediate linker densities. (a) Top-view of the membrane before and after the extrusion of the tube. The linkers avoid the tube and often to segregate into small clusters. (b) The average separation between the linkers decreases as the tube is extruded from the membrane. The linkers cluster size (dark purple line, inset) increases as the extrusion progresses (linkers are considered in the same cluster if the distance between them is less than  $3\sigma$ ). The data is obtained from averaging over 20 simulations where the tube is extruded to a target length of  $l_0 = 60\sigma$ . The shading represents the standard error of the measurements over 5 different simulation seeds. (c) Heatmaps showing the number of linkers per unit of membrane area at low (5%) and high (50%) linker densities. The squares are weighted according to the membrane area they contain. The central region, where the tube is extruded, (marked with a red 'X' symbol) is depleted of linkers at low densities, showing that proteins prefer to diffuse away from the tube. At high densities, the proteins are distributed uniformly with a significant fraction trapped inside the tube.

required to reach the same extension is slightly greater at higher protein concentrations [22], but the plateau force does not show a significant difference at low to moderate linker concentrations. However, if the linker density is high enough, the growth of a protrusion requires a large scale concerted reorganisation of the linkers in order to facilitate the transport of sufficient material into the tubular structure. In this case the linkers cannot effectively exclude themselves from the tube and will experience a pulling force, at the same time slowing down the dynamics of tube extrusion.

The growth of a protrusion hence becomes expensive, resulting in an exponential increase in the pulling force for higher protein concentrations, as seen in Fig. 2c, eventually fully preventing tube extrusion. Taken together, our model shows that the extrusion force can have non-linear dependence on the linker densities: at low densities the force will not be influenced by the linkers, while at higher linker densities the force will substantially increase.

To make sure that our results are not influenced by the details of our model – namely the Monte Carlo routine that might not capture the correct dynamics – we repeated our results using a nontriangulated spherical membrane within a molecular dynamics (MD) framework. As shown in the SI Section II H and Fig.S10–12, our results remain qualitatively unchanged; the molecular dynamics model also shows a non-linear dependence of the force on the linker density (Fig. S12). The only difference is that the critical linker density at which the force increases is shifted to lower values compared to the Monte Carlo model. These results indicate that the non-linear dependence of the force on the linker density is a robust result, which is not model dependent. However, the point at which the behaviour in the force changes from the flat curve to the exponential increase will depend on the details of the model and the system, namely on the strength of the interactions of the proteins with the cortex and the membrane fluidity.

Finally, we observe that presence of linkers also causes a slight reduction in the tube radius (Fig. 2d): as the linker density increases, there is less material freely available for the tube growth, resulting in a decrease of the tube diameter (see also Fig. S11 for the MD results ).

Linkers are excluded from the tube at intermediate densities. Fig. 3a further depicts how the linker proteins dynamically rearrange around the tube during tube extrusion in our simulations. To minimise the strain imposed on the membrane the linkers avoid the region around the tube neck, giving rise to a concentration gradient in proteins around the tube. To minimise the perturbation to the membrane the proteins also tend to segregate into small protein-rich clusters (Fig. 3b and Fig. S10). These results indicate that pulling the tube promotes the local aggregation of proteins by depleting the available phospholipid membrane area. The tube itself can con-



FIG. 4: Measuring plateau force for varying Ezrin levels. (a) Force regimes observed during tether pulling from cells corresponding to the steps depicted on the right. A bead at equilibrium position (1) is brought into contact with a cell (2). When the stage is pulled away from the trap, an initial force peak required to detach the membrane from the underlying actin cortex (3) is quickly followed by a stable force  $F_{tether}$  (4). (b) Plateau forces in HEK293T cells overexpressing GFP (n=19), GFP-EzrinWT (n=28), GFP-EzrinTD (n=21) or treated with control siRNA (sicontrol, n=36), Ezrin siRNA (siEzrin, n=23), or ERM siRNA mix (siERM, n=18). mean $\pm$ S.D(\*\*p = 0.008; \*p = 0.03). The depletion of Ezrin does not change the tether force compared to the control, while the expression of recombinant Ezrin-WT and Ezrin-TD proteins increases the force compared to control.

tain small concentrations of trapped proteins (Fig. 3c). We observed that the partitioning of proteins between the tube and the membrane is influenced by the linker binding energy, with higher binding energies resulting in stronger linker exclusion from the tube and smaller tube radii (Fig. S4 and Fig. S5).

Experiments: Influence of Ezrin expression on tube extrusion in cells. In order to qualitatively compare simulation results to experiments, we pulled tethers from adherent cells as depicted in Figure 4a with a typical length of 10  $\mu m$ . As previously shown, the initial force overshoot depends on the interaction area between the bead and the plasma membrane [14] and is not well controlled under experimental conditions (Figure 4a – step 3 and Fig. S16a). Upon tube formation the force next decreases and rapidly reaches a plateau. To measure the static tether force, we waited 10 seconds to let the force equilibrate (Figure 4a – step 4, and Figure S16a). In agreement with previous reports [32], we confirmed that the tether force does not depend on tether length over a broad range (Fig. S16b). In addition, special care was taken to ensure the measurement of forces for single tethers only.

We measured a tether force of  $16.0 \pm 2.7$  pN in HEK293T cells, which is at the lower end of values reported for adherent cells [33] (Figure S14). We checked that the tether force is unchanged when the membrane receptor EphB2 is expressed (HEK-EphB2). It is similar in HeLa cells, yet higher in RPE-1 cells (Figure S14), highlighting that the static tether force varies depending on the cell type. Next we studied the effect of the manipulation of the expression level of ERM on tether force. Since the transfection process can perturb the cells, we compare the measurements after depletion or over-expression with control cells transfected according the same protocol, but without plasmid. GFP-Ezrin is binding actin in a phosphorylation-dependent manner, whereas the phosphomimetic mutant GFP-EzrinTD permanently links actin to the membrane [34] (Figure S15b).

We found that the depletion of Ezrin or ERM using siRNA [35] (Figure S15c) did not change the tether force compared to the control (Figure 4b), while the expression of recombinant Ezrin-WT and Ezrin-TD proteins significantly increases the force in HEK cells as compared to control (Figure 4b and Figure S15). These results favourably compare to our simulations where we found that the force to extrude tethers increases only when the expression level of linkers is substantially increased. Moreover, depleting Ezrin/ERM enabled faster tube relaxation (Fig. 5b), in agreement with what has been observed in simulations (Fig. 5a). These tube relaxation results indicate that the presence of linkers imposes a barrier for the diffusion of membrane components, effectively increasing the local friction [27, 36].

Linkers stabilise single isolated membrane tubes. Membrane tubes pulled from bare membranes have been previously shown to merge in each other's proximity, as the free energy change involved in this process is always favourable [12, 37]. To test the influence of the linkers on the interactions between protrusions we go back to our simulations. We pulled two tubes at an initial separation of  $\Delta L = 20\sigma$  and allow them to diffuse freely. For bare membranes we observe that the two tubes grow and eventually merge through the basal truncated cone-like structure into a charac-



FIG. 5: Tether extrusion dynamics depends on the density of linkers. a) Simulations: As the tube is extruded the force acting on the protrusion's tip decays to a constant non-zero value. The dynamics of this relaxation is slower at higher linker density. Each curve represents an average over 5 simulation repeats. b) Experiment: upon fast tube extrusion the force relaxes to a constant non-zero value within a time-scale that is faster upon depletion of ERM/Ezrin.

teristic Y-shape as shown in Fig. 6a. This finding is in qualitative agreement with the structures observed in double tube experiments [37]. Interestingly, when the linkers are introduced, the probability of tube fusion decreased substantially and is lower at higher linker densities (Fig. 6c).

To quantify this effect we measured the relative separation between the tips of the two pulled tubes during the course of the simulation (Fig. 6b). The average separation between the tubes decreases more slowly at greater protein densities, indicating that the linkers act as obstacles in the merging process. The fusion requires the neck of the tubes to be depleted of linkers before it can take place. The proteins thus act as a barrier for the fusion process, stabilising isolated tubes. Our findings are in an excellent agreement with the study of Nambiar et al [38], where the authors found the membrane tethers coalesce more frequently when the membrane-cortex attachment is decreased by depleting myosin 1 proteins, just like in our simulations. It can thus be concluded that the membrane linkers can serve to stabilise protrusive structures involved in physiological processes. Along the same lines, Datar et al. found that the membrane tethers move more easily on the surface of axons when the cytoskeleton is disrupted [36].

## **III. DISCUSSION AND CONCLUSIONS**

Using coarse-grained simulations, where lipid components and membrane-cortex attachments are modelled explicitly, we found that the force needed to extrude a protrusion has a non-linear dependence on the density of the membrane-cortex attachments. At low and moderate densities of the attachments the amplitude of the force needed to extrude a protrusion is unchanged compared to the bare membrane. In our model this is caused by i) the ability of the membrane to flow around the attachments, effectively excluding the attachments from the tube and ii) the ability of the attachments to diffuse away from the tube rather than to oppose the tube extrusion. In contrast, when the linker density is increased above a certain threshold, the force needed for tube extrusion increases significantly, as the linker exclusion out of the tube becomes prohibitively slow, as also demonstrated by a slower relaxation of the force to the equilibrium value.

The linkers' diffusion in our model effectively captures dynamic ERM attachment/detachment from the cortex [39]. To test the influence of the linkers' diffusion on our key results on the nonlinear dependence of the extrusion force on the linker density, we repeated these measurements for the case where the linkers cannot rearrange and diffuse in the membrane plane, but can still detach. As shown in Fig S9, the non-linear behaviour of the force versus linker density remains unchanged, although the point of cross-over was again shifted to lower linkers' density.

Our simulations indicate that great care should be taken when considering the membrane-cortex attachments in an effective way, for instance by describing them as a uniform membrane-surface adhesion or rescaled membrane tension. Here we show that the local effects, such as the exclusion of linkers from the tube, can have non-trivial effects on the resulting extrusion force, as well as on the protrusion radius.

Our experimental results align well with the non-linear effects observed in simulations and show that depletion of Ezrin/ERM does not have sig-



FIG. 6: **Protrusion coalescence is hindered by linkers.** (a) Tubes coalesce into a Y-shaped structure that transforms into a single long tube. (b) The distance between the pulled points on two tubes decays quickly at low protein densities, showing the tendency of adjacent tubes to coalesce. At higher densities, the linkers will maintain a stable distance, as they are prevented from fusion. The shading represents the standard error of the measurements over 20 different simulation seeds. (c) The percentage of tubes that fuse into a single structure decreases at greater linker densities.

nificant influence on the plateau force, while the over-expression does. Such a response could indicate that the density of linkers in HEK cells might lie close to the transition from a plateau to exponential part of the force versus linker density curve.

Previous studies based on static tether force measurements have reported effects of Ezrin and myosin 1 isoforms on effective membrane tension in non-adherent cells such as lymphocytes [23], or in cells with high blebbing propensity [40]. By interfering with the membrane-cortex attachment Diz-Muñoz et al. showed that a decrease in the attachment led to a decrease in the force required to pull a membrane tether from developing zebrafish embryo [22]. Recently, Berget et al. reported that incorporating a synthetic linker that increases the membrane-cortex attachment in embryonic mouse cells led to an increase in the tether-extrusion force [41]. Similarly, de Belly et al. [42] observed that mouse embryonic stem cells that display membrane blebbing, which is believed to be a sign of low membrane-to-cortex attachment, require lower force for tether extrusion than non-blebbing cells. In all of the above cases the forces required for tube extrusion from embryonic cells ( $\sim 50 \text{ pN}$ ) were substantially higher than in the case of HEK cells probed here ( $\sim 10$  pN). It is hence likely that the expression level of membrane-cortex attachments in embryonic cells is naturally higher than in epithelial cells studied here, which would cause the experiments to probe the right-most part of the non-linear curve presented in Fig. 2c, where the force severely depends on the linker concentration.

The critical density of linkers after which the force increases is expected to depend on the details of the system as well as on the pulling procedure. It is important to note that, due to the simplicity of our model and the lack of experimental information on the interactions and local densities, it was not our intention to provide quantitative mapping to an exact experimental system. In fact, our simulations clearly show that when using different models that assume different protein-cortex interactions and different degrees of the membrane and linkers' diffusivities, the value of the critical linker density can substantially change. However, our central result – the non-linear nature of the force versus linker density curve – remains unchanged.

Our simulations and experiments were carried out in a quasi-static way, which is comparable to the rate of filopodia growth [43], and allows enough time for the linkers to exclude themselves out of the tube for low and moderate linker densities. Such a pulling is however still not infinitely slow, and for high linker densities the force is not necessarily relaxed. Hence it can be expected that the critical linker density for which the force increases will be shifted to higher values for even slower pulling and longer equilibration times, both in simulations and experiments. Likewise, in fast dynamic pulling experiments the crossover might be shifted to lower linker densities, or the initial plateau of constant force might be completely absent. Interestingly, non-linear effects similar to those reported in here were observed in dynamic tether pulling experiments via hydrodynamic flows in red blood cells [8]. The extrusion velocity of the tube was observed to depend on the extrusion force in a nonlinear fashion, which was also attributed to the dynamical feature of membranecytoskeleton attachments.

Numerous other effects can be present in live cells that contribute to the membrane-pulling force when the level of ERM proteins is changed. This can for instance include changes in cell adhesion, and changes in the cytoskeleton architecture and flow, which all might have an influence on the global tension and/or local force. However, even in the minimal system presented in our simulations it is apparent that the role of membrane-cortex attachments might be non-trivial and that their contribution cannot be necessarily considered as an effective adhesion or an effective tension.

Finally, in simulations we observed that the linkers have a substantial effect on the stability of protrusions, preventing their diffusion and rendering them more stable against coalescence with other tubes, in line with previous experimental observations of the increased tube coalescence upon depletion of myosin 1 proteins [38]. We hope that our simulations and experiments will renew discussions on the mature but still unsolved question of the influence of membrane-cortex attachments on membrane tension and membrane reshaping, and stimulate the investigation of local effects and protein dynamics in the formation of membrane protrusions.

#### Author contributions

A.P. performed and analysed the Monte Carlo simulations. C.V.C. developed and analysed the molecular dynamics simulations. A.Š. conceived and supervised the research. T.L., E.C. and P.B designed, performed, and analysed the membrane pulling experiments. The manuscript was written by A.P., P.B., and A.Š. All authors revised and approved the manuscript.

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## Simulation details

The membrane was modeled as a dynamically triangulated sheet with hard sphere beads situated at every vertex of a triangular lattice [44–46]. The system contains two types of beads, one type representing the membrane phospholipid patches and another corresponding to ERM (Ezrin, Radixin or Moesin) proteins. These proteins are responsible for linking the membrane phospholipids and transmembrane proteins to the underlying cytoskeletal components [19]. The dynamics of the system are governed by a Monte Carlo algorithm that involves two types of moves: vertex displacement moves and bond flip moves (Fig. S1). The vertex displacement moves mimic the lateral diffusion of lipids and proteins and allows for vertical membrane fluctuations. The bond flip moves ensure that the membrane preserves its fluidity by dynamically rearranging the connectivity. This membrane model captures correct mechanical properties of biological membranes – bending rigidity, fluidity, and the response to external forces. The model however does not describe the bilayer explicitly and the effects such as interleaflet friction are not included. The model also does not allow for topology changes such as poration or tube scission.

A tube is drawn from the membrane by applying an external point force to a predetermined lipid bead. The simulation methodology aims to mimic a slow pulling procedure such as optical tweezers or an atomic force microscopy experiment. The system's energy is calculated using the Hamiltonian described in equation (1). This takes into account the energy required for bending a membrane patch, the energy required to create new surface area as well as the harmonic potential imposed to extrude the tube. The membrane to cortex attachment energy is controlled by a series of harmonic springs that can unbind at distances sufficiently far from the vertical equilibrium position.

$$\mathcal{H} = \frac{\kappa}{2} \sum_{\langle ij \rangle} (1 - \mathbf{n}_i \mathbf{n}_j) + \gamma A + k_{\text{pulling}} (l - l_0)^2 / 2 + \sum_k k_{\text{linking}} (z_i - z_0)^2 / 2 + \mathcal{H}_{\text{steric}} \quad (1)$$

where  $\kappa$  is the membrane's bending rigidity, i, j are pairs of triangles sharing an edge,  $\mathbf{n}_i$  and  $\mathbf{n}_j$  are the normal vectors of the two triangles,  $\gamma$  is the membrane surface tension and A is the total membrane area.  $k_{\text{pulling}}$  is the pulling constant that controls the tube extrusion, l is the vertical position of the pulled membrane bead and  $l_0$  is its target equilibrium position, which is varied between  $20\sigma$  and  $100\sigma$  to obtain the force-extension curves. The linkers are attached to the membrane through an harmonic potential controlled by the spring constant  $k_{\text{linking}}$  (Fig. 2a). The vertical position of each linker is denoted by  $z_i$  with  $z_0$  being its equilibrium vertical position.  $\mathcal{H}_{\text{steric}}$  is a term that accounts for the hard sphere repulsion between beads, forbidding any two beads to be closer than  $\sigma$ . The beads are not allowed to be at distances greater than  $3\sigma$  to avoid the formation of holes and pores.

A single Monte Carlo sweep consists of  $N_{\text{vertices}}$ attempts at displacing a vertex and  $N_{\rm vertices}/2$ bond flip attempts, where  $N_{\text{vertices}}$  is equal to the total number of membrane beads. In this case  $N_{\text{vertices}} = 3600$  and the initial vertical side lengths  $L_x = L_y = 60\sigma$ . The acceptance probability of a Monte Carlo move is controlled by a simple Metropolis-Hastings criterion p = $\min[1, \exp(-\Delta E/k_B T)]$ , where  $\Delta E$  is the energy change after a potential Monte Carlo move. The simulation is run in the  $NA_pT$  ensemble, where  $A_p$  is the membrane's projected area. The bending rigidity is set to  $\kappa = 20k_BT$ , and the surface tension is kept constant at  $\gamma = 3k_BT/\sigma^2$ . The pulling constant,  $k_{\text{pulling}} = 1k_B T/\sigma^2$ , is chosen to ensure that the extrusion is slow enough such that the membrane beads can diffuse and rearrange on the timescale of the tube extrusion. The resulting force matches those in the experiments ( $\sim 10 \text{pN}$ ) when by taking the MC unit length to be  $\sigma = 10 \text{ nm}$ (which also matches the experimentally observed tube radii of tens of nm.)

The linkers are subjected to an additional harmonic potential with constant  $k_{\text{linking}} = 3k_B T/\sigma^2$ , but they are allowed to freely diffuse in the membrane's plane. The linkers are allowed to unbind if they are displaced from their vertical equilibrium position of  $z_0 = -3\sigma$  by more than  $5\sigma$ , but also to rebind to maintain the simulation's detailed balance. The choice of the equilibrium position and attachment constant did not significantly affect the results. The membrane is equilibrated for 300,000 Monte Carlo steps and then the simulations are run for 2 million steps in total. During equilibration, additional volume change moves are implemented to relax the membrane surface tension. After equilibration, the spheres on the membrane's edge are fixed in place and the box size is kept fixed during the extrusion. We checked that the system has enough excess membrane area such that the boundary conditions do not affect our results (Fig. S3).

The force profile is recorded by running a set of simulations in which a single randomly chosen central membrane bead is pulled to incrementally increasing equilibrium positions. The membrane will thus exert a restoring force on that bead. The force profile is then reconstructed from the average values of the force at every extension. This procedure allows the reconstruction of the forceextension curve obtained in a static pulling experiment. The force experienced by the protrusion's tip decreases as the protrusion grows. The force eventually converges to a finite nonzero value as the protrusion reaches its maximum extension (Fig. S2). The measured pulling force is calculated as  $F_{\text{pulling}} = k_{\text{pulling}} \Delta z$ , where  $k_{\text{pulling}}$  is the pulling force constant and  $\Delta z$  is the distance between the point being pulled and its vertical equilibrium position. As such, the force will not reach a zero value and the protrusion will grow until it reaches an extension around which it will fluctuate. We measured the force by averaging it over the course of the 1 million Monte Carlo steps, ensuring that it reached convergence. Each forceextension curve is obtained from averaging over five different random seeds, unless otherwise specified.

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