The sequential exchange of filament composition to increase filament curvature was proposed as a mechanism for how some biological polymers deform and cut membranes. The relationship between the filament composition and its mechanical effect is lacking. We develop a kinetic model for the assembly of composite filaments that includes protein–membrane adhesion, filament mechanics and membrane mechanics. We identify the physical conditions for such a membrane remodeling and show this mechanism of sequential polymer assembly lowers the energetic barrier for membrane deformation.

DOI: 10.1103/PhysRevLett.129.268101

Nanoscale filaments that can produce work to reshape membranes are key for cell traffic, motility, and healing. These include the third endosomal sorting complex required for transport (ESCRT-III) proteins, which form polymers that reshape and cut membrane surfaces [1,2] and are involved in a range of membrane remodeling events, including membrane budding, cell division, and membrane repair [3–7]. ESCRT-III proteins come in many variants that copolymerize to form membrane-associated composite filaments. These composite polymers dynamically exchange different monomer types with the surrounding cytoplasm. Monomer exchange is driven by the activity of an enzyme called Vps4, which extracts energy from the hydrolysis of ATP (adenosine triphosphate). Vps4 extracts certain monomers while new monomers are added to the polymers that remain [2,8,9].

Recent experiments [10,11] have suggested that the composition of ESCRT-III filaments changes with time in a precise sequence as monomers assemble, and they are removed and replaced by monomers of a different type, in a fixed order, as shown in Fig. 1(a). Additionally, although monomers of different types share a similar molecular structure [2,5], polymers of different ESCRT-III monomers have been observed to have different shapes and different membrane binding interfaces, depending on their precise monomer composition [5,12–17]. Therefore, as the copolymers change in composition, their geometry and mechanical properties are also expected to change, allowing the ESCRT-III to perform work on the membrane [8,18–21].

The time ordering of this staged assembly-disassembly process is robust and is believed to progressively increase membrane curvature to constrict the membrane neck.

Although great progress has been made on understanding this biophysical system, the physicochemical properties that enable different monomer types to assemble into dynamically shape-shifting polymers remain unclear. Here, we develop a kinetic model that can capture the recruitment and spontaneous unbinding of monomers with

FIG. 1. Shape-shifting by sequential monomer exchange. (a) In vitro experiments, tracking the presence of different monomer types (termed Snf7, Vps2, and Did2) in membrane-bound yeast ESCRT-III filaments, showed that the filament composition changes according to a precise sequence [11], which is accompanied by the membrane deformation. (b) The geometry of a membrane-bound filament has been proposed to change together with its composition, driving membrane remodeling for cargo transport.
different physical properties. We identify rules under which the mechanical and chemical differences between different monomer types alone can drive changes in polymer assembly-disassembly kinetics, and consequently in polymer structure. These design rules allow us to make predictions about physiological properties of ESCRT-III monomers that are difficult to obtain experimentally, reveal the yet unexplained purpose of intermediate monomer types as catalysts for membrane deformation, and can inform efforts toward the generation of artificial shape-shifting nanomaterials.

**Model.**—The filaments of ESCRT-III homologs across evolution involve between 3 and 11 different monomer types, as in archaea [3] and animal cells [4], respectively. In addition, in systems with a large number of ESCRT-III types, some monomer types always bind together, effectively decreasing the number of assemblies [11]. To model the process, we therefore choose to use three types of monomers (i = 0, 1, or 2), as in Fig. 2(a), which can bind to the membrane and each other, forming heterogeneous membrane-bound filaments [Fig. 2(b)]. The filament is treated as a lattice, where each site can host up to one monomer of each type, such that different monomer types can bind side by side on a site. Each monomer type is associated with an energy of adhesion $E_{\text{adh}}$, as well as a preferred curvature $c_i$, and a preferred effective pitch $p_i$ (which is the membrane deformation depth induced by a homogeneous filament of that type). Figure 2(a) shows the case in which the pure filament of type 0 is a completely flat spiral ($p_0 = 0$) of curvature $c_0$, whereas filaments of types 1 and 2 are helices with larger pitches and curvatures, such that $0 < p_1 < p_2$ and $c_0 < c_1 < c_2$. A heterogeneous filament, made of monomers of different types, is associated with an average curvature $C$ and pitch $P$, which are averages of the curvatures and pitches of all its monomers.

Our model considers three energy contributions: adhesion energy $-E_{\text{adh}}$, filament bending energy $E_{\text{fil}}$, and membrane energy $E_{\text{mem}}$. The energy gained by adhesion is

$$-E_{\text{adh}} = -\sum_{m,i} n_{m,i} E_{\text{adh}} < 0$$

(where $n_{m,i} = 1$ if site $m$ contains a type-$i$ monomer, and 0 otherwise) and always favors monomer binding. A heterogeneous filament has a nonzero bending energy, which results from monomers of different preferred curvatures binding to the same site or to neighboring sites, causing local elastic frustration. Assuming that sections of filament bend as elastic rods, we estimate the filament bending energy as

$$E_{\text{fil}} = \sum_{m,i} n_{m,i} \frac{\kappa}{2} (C_m - c_i)^2,$$

where $\kappa$ is the filament rigidity, and $C_m$ is the local filament curvature averaged over the eight nearest neighbors of site $m$. This is based on the observation that individual monomers in ESCRT-III assemblies interact with numerous (four to eight) other neighboring monomers via direct contact [17,24]. Finally, we estimate the membrane bending energy using the Helfrich model [25]:

$$E_{\text{mem}} = \frac{1}{2} k_{\text{mem}} \int_{\text{cap/tube}} C^2 dA,$$

where $k_{\text{mem}}$ is the membrane bending rigidity, and $C$ is the membrane curvature. We approximate the deformation of the membrane as a spherical cap or a cylinder with a spherical cap; thus, only two quantities ($C$ and $P$) are needed to describe the membrane shape [Fig. 2(b)]. Further details are provided in the Supplemental Material (SM) [26].

Overall, the adhesion term promotes binding, the filament elasticity term disfavors the neighboring binding of monomers with different curvatures, and the membrane term penalizes the binding of monomers with nonzero pitch and high curvature, which can only become part of the polymer if the membrane bends.

We investigate the dynamics of the system using the Gillespie algorithm, which is a variant of dynamic Monte Carlo. Monomers bind or unbind according to rates $r_{\text{on}}$ and $r_{\text{off}}$. We assume that monomer concentrations in the cytoplasm are constant, as in in vitro experiments [11], and equal for different monomers; and we assume that binding is diffusion limited so that $r_{\text{on}}$ is constant and sets the
timescale of the simulations. The probability of a given unbinding event (i.e., a given monomer detaching from a given site) depends exponentially on the energy change associated with it: \( r_{\text{off}} = r_{\text{on}} e^{(\Delta E_{\text{adh}} + \Delta E_{\text{mem}})} \). These energies depend on the composition of the filament; hence, the unbinding rates are not just monomer type and lattice-site dependent but also depend on the environment of the monomer, which changes in time. We tested that all our conclusions are robust against the choice of \( r_{\text{on}} \) and \( r_{\text{off}} \) (Fig. S3), as long as the ratio \( r_{\text{on}}/r_{\text{off}} \) is preserved via standard thermodynamics.

The length scale for our system is set by the reported curvature of an ESCRT-III monomer found in yeast that forms spirals with a radius of \( \sim \text{curvature of an ESCRT-III monomer found in yeast} \) that represents monomer 0 in our model. Membrane rigidity is set to 20\( k_B T \) [27]. Polymer rigidity is compared with indirect experimental estimates of physiological persistence lengths [13,28]. Polymer curvatures, pitches, and adhesion energies are parameters that we vary within a range of physiologically plausible values (Table S1). Our results hold for any value of polymer length within a physically relevant range (Fig. S3); here, we show results for a 42-site-long lattice.

**Minimal design rules.**—In our efforts to explain the experimental data shown in Fig. 1(a) [11], our first task is to find the conditions that enable monomers to bind and unbind sequentially. To do so, we explore a range of values for the adhesion energies \( E_{\text{adh}}^i \), curvatures \( c_i \), and pitches \( p_i \) of the three monomer types. We quantify how much of each monomer type is present on average in the filament as a function of time, similar to the experimental curves [Fig. 1(a)].

In Fig. 3(a), we first verify that monomers of equal properties bind together and reach an equilibrium lattice coverage for which the value is determined simply by the ratio of the binding and unbinding rates. Figure 3(b) shows that sequential monomer binding emerges when monomers are assigned different pitches, irrespective of their curvatures and adhesion energies; flat monomer 0 (blue) binds before low-pitch monomer 1 (purple), which binds before high-pitch monomer 2 (pink). At time 0, the membrane is flat so that binding of high-pitch monomers is penalized. Importantly, this initial state of the membrane creates a time asymmetry that dictates the ordering of binding. The reaction progresses until the energy of membrane deformation, induced by late-binding helix-shaped monomers, is balanced out by monomer adhesion energy. In summary, for the sequential binding to occur in the order 0 \( \rightarrow 1 \rightarrow 2 \), \( p_2 > p_1 > p_0 \) must be satisfied. Although these conditions result in sequential binding, they are not sufficient to significantly deform the membrane, which requires the unbinding of the low-pitch monomers.

Sequential unbinding in the model is a consequence of monomer frustration within a filament. If the preferred shape of a particular monomer is very different from the local average filament shape, the monomer pays a substantial cost in elastic energy and will likely spontaneously unbind. For this effect to cause asymmetric unbinding, one monomer type must be favored. This is the case if the monomers also have different adhesion energies, as shown in Fig. 3(c). The monomer type with highest

![FIG. 3. Geometry and adhesion control sequentiality. The effect of the pitches, curvatures, and adhesion energies for the three types of monomers on the filament composition (percent of lattice coverage). When monomers are all equal, (a) the filament stabilizes as a composite mixture of the three monomers in equal parts. When the monomer types have different pitches, (b) the binding of high-pitch monomers is delayed. Differences in curvature and adhesion energy together (c) result in the unbinding of weak-binding monomers, due to bending frustration. (d) Differences in pitch, curvature, and adhesion energy are all three needed to achieve sequential assembly and disassembly. Parameters are as follows: equal pitches [(a) and (c)], \( p_0 = p_1 = p_2 = 0 \text{ nm} \); otherwise, [(b) and (d)], \( p_0 = 0 \text{ nm}, p_1 = 30 \text{ nm}, \text{ and } p_2 = 45 \text{ nm} \). Equal curvatures [(a) and (b)], \( c_0 = c_1 = c_2 = (15 \text{ nm})^{-1} \); and \( E_{\text{adh}}^0 = E_{\text{adh}}^1 = E_{\text{adh}}^2 = 6.5 k_B T \); otherwise, [(c) and (d)], \( c_0 = (30 \text{ nm})^{-1}, c_1/c_0 = 2.5, c_2/c_0 = 8, \text{ and } E_{\text{adh}}^0 = 6.5 k_B T, E_{\text{adh}}^0/E_{\text{adh}}^2 = 3, E_{\text{adh}}^2/E_{\text{adh}}^0 = 12 \). Time is in units of \( 10^3 \) simulation steps.](image-url)
membrane-adhesion energy ends up dominating and eventually forcing the other two off the membrane. Conversely, the first monomer to unbind is the one with the smallest adhesion energy and a curvature most different from the first monomer to bind. Hence, if unbinding to occur in the order 0 → 1 → 2, \( c_2 > c_1 > c_0 \) and \( E_2^{\text{adh}} > E_1^{\text{adh}} > E_0^{\text{adh}} \) must be satisfied. The sequential disassembly fails if the monomers’ curvatures are too dissimilar; in which case, prohibitive filament frustration causes monomers to unbind as soon as they bind.

Taken together, this analysis suggests that sequential subunit exchange requires increasing pitch, increasing adhesion energy, and increasing curvature with time [see Fig. 3(d)]. A failure to satisfy any one of these three criteria hinders the sequence of binding or unbinding events. The robustness of this behaviour against changes in curvature and adhesion energies is further quantified in Fig. S4.

Sequential binding-unbinding.—To explore why the sequential assembly-disassembly is needed for membrane remodeling, we next analyze different possible pathways to the final membrane-wrapped tight helix configuration. Using the parameters that result in the most pronounced sequential assembly and disassembly of the monomers [Fig. 3(d)], we calculate the average energy of the system as a function of monomer 2 content [Fig. 4(a), turquoise curve]. We then compute the same quantity by sampling configurations in which we force monomers of type 2 to bind on their own, without preceding subunits (gray curve). The binding of monomers that prefer to form tight helices (type 2) exhibits a high energy barrier, which is completely erased when the system evolves in a stepwise fashion from 0 to 1 to 2. The transient presences of lower-curvature monomers 0 and 1 act as mechanical catalysts and deform the membrane enough to ensure the tight helical monomers can bind without having to dramatically bend the membrane from flat to highly indented in one step. This conclusion is further confirmed by measurements of the complete energy landscape (Fig. S7). As a consequence, the sequential assembly allows the process to occur in less time (Fig. S5).

If the adhesion energy is high enough, tight helical monomers could in principle overcome the binding barrier and deform the membrane without the need for intermediate filaments. However, we expect there to be an upper limit to the adhesion energy, as in any supramolecular system, due both to the physical-chemical constraints of the membrane interaction and to the fact that active disassemblers (e.g., Vps4) need to be able to disassemble the filament to trigger neck fission using the finite energy provided by ATP hydrolysis.

Figure 4(a) shows that the presence of multiple monomer types renders the binding of a highly membrane-deforming filament possible through lowering the energy barrier, but it does not explain why earlier monomer types also need to unbind. Figure 4(b) shows the depth of membrane indentation for four different cases where monomer unbinding is suppressed. In all these cases, the final composition of the filament is different from the target one (type-2 monomers only); as a consequence, the extent of membrane indentation is reduced. Thus, the unbinding of monomers of lower curvatures and pitches optimizes the mechanical action of the filament on the membrane and is expected to be needed for membrane neck scission [29].

Role of activity.—All the parameters in our model are physiologically realistic (Table S1), except for the bending rigidity of the filament. To reproduce the unbinding sequence observed in experiment, we find that our filament rigidity needs to be between three- and sevenfold larger than previous estimates [13,28]. This apparent mismatch is due to the fact that our passive model does not account for...
active energy supply; whereas in experiments, dynamic disassembly is only observed upon addition of ATPase Vps4, which unfolds monomers and extracts them from the filament [9,10,20,30–34]. This led us to explore whether a Vps4-like activity that promotes stress-dependent polymer disassembly could aid the process. We therefore extended the model so that a monomer with a bending stress above a certain threshold received an additional energy penalty, with a multiple of $E_{\text{ATP}} \approx 13k_B T$, making it more likely to unbind (details are provided in the SM [26]). Figure 4(c) shows that maximal membrane indentation can be recovered for realistic values of the filament rigidities ($< 0.8k_B T \mu m$) if enough energy is provided to frustrated monomers. The amount of energy needed is of the order of $3E_{\text{ATP}} \sim 39k_B T$, which is consistent with the energy consumed by ATPases such as Vps4 [35]. Our findings are hence in line with the hypothesis that frustrated monomers in heterogeneous polymers could be more favorably exposed to Vps4, which in turn preferentially extracts them from the polymer [18].

Discussion.—In this Letter, we have shown how the staged assembly and disassembly of nanoscale filaments can deform membranes. Such polymers function as a dynamic shape-shifting metamaterial: a filament changes its composition, and hence its properties, over time to perform a function that each of its parts cannot achieve when acting alone. The overall effect is a membrane-bound elastic structure that becomes more curved with time [21]. Over time, the recruitment of high-pitch monomers is favored, which in return facilitates the unbinding of low-pitch monomers, generating an increasingly deformed membrane. The dynamics of the structure is governed by the properties of the composite monomers and by membrane mechanics. To generate a deep indentation of the membrane, subunits must have increasing adhesion energies, increasing preferred curvatures and increasing preferred membrane indentations, which together lower the energy barriers for binding.

Here, we have considered only three types of monomers. However, it is likely that the inclusion of additional monomer types (as seen in many experimental systems) can further lower the minimum membrane-adhesion energy required to drive the process to completion. Nevertheless, adding more monomer exchanges costs additional time and adds complexity. It is to be expected that there is an optimal number of subunits that minimizes the overall time for membrane deformation by minimizing the energy barriers and the number of steps in tandem.

Within our framework, the unbinding of earlier subunits is caused by filament frustration. This frustration could be realized in different ways. It could be a result of monomers bound together having different curvatures (as shown above), different torsion rigidities (explored in Fig. S8), or a combination of both. In addition, there is mounting evidence that the mechanical properties of the membrane also modulate ESCRT-III function by affecting the binding dynamics of the monomers [36–39]. Our model quantitatively supports this hypothesis because monomers are more likely to bind to a membrane that is already deformed according to the filament’s preferred morphology. Finally, these insights may inform the design of dynamic cell reshaping nanomachines, such as membrane-attached DNA-origami structures [40,41].

We thank T. C. T. Michaels and J. Palacci for useful discussions. We thank Claudia Flandoli for the illustrations in Fig. 1(b) and Fig. 2. We acknowledge funding by the European Union’s Horizon 2020 Research and Innovation Programme under the Marie Skłodowska-Curie Grant Agreement No. 101034413 (I. P.), the Royal Society Grant No. UF160266 (A. Š.), the European Research Council under the European Union’s Horizon 2020 Research and Innovation Programme (Grant No. 802960; B. M., I. P., and A. Š.), and the Volkswagen Foundation Life Grant (B. B. and A. Š.).
G. Hummer, C. Bustamante, and J. H. Hurley, ATP-dependent force generation and membrane scission by ESCRT-III and Vps4, Science 362, 1423 (2018).

[11] A. K. Pfitzner, V. Mercier, X. Jiang, J. Moser von Filseck, B. Baum, A. Šarić, and A. Roux, An ESCRT-III Polymerization sequence drives membrane deformation and fission, Cell 182, 1140 (2020).

[12] W. M. Henne, N. J. Buchkovich, Y. Zhao, and S. D. Emr, The endosomal sorting complex ESCRT-II mediates the assembly and architecture of ESCRT-III helices, Cell 151, 356 (2012).

[13] Q. T. Shen, A. L. Schuh, Y. Zheng, K. Quinney, L. Wang, M. Hanna, J. C. Mitchell, M. S. Otegui, P. Ahlquist, Q. Cui, and A. Audhya, Structural analysis and modeling reveals new mechanisms governing ESCRT-III spiral filament assembly, J. Cell Biol. 206, 763 (2014).

[14] S. T. Huber, S. Mostafavi, S. A. Mortensen, and C. Sachse, Structure and assembly of ESCRT-III helical Vps24 filaments, Sci. Adv. 6, 34 (2020).

[15] J. Moser von Filseck, L. Barberi, N. Talledge, I. E. Johnson, A. Frost, M. Lenz, and A. Roux, Anisotropic ESCRT-III architecture governs helical membrane tube formation, Nat. Commun. 11, 1516 (2020).

[16] M. Alqabandi, N. De Franceschi, S. Maity, M. Bally, W. H. Roos, W. Weissenhorn, P. Bassereau, and S. Mangenot, The ESCRT-III isoforms CHMP2A and CHMP2B display different effects on membranes upon polymerization, BMC Biol. 19, 66 (2021).

[17] H. C. Nguyen, N. Talledge, J. McCullough, A. Sharma, F. R. Moss, J. H. Iwasa, M. D. Vershinin, W. I. Sundquist, and A. Frost, Membrane constriction and thinning by sequential ESCRT-III polymerization, Nat. Struct. Mol. Biol. 27, 392 (2020).

[18] A. K. Pfitzner, J. Moser von Filseck, and A. Roux, Principles of membrane remodeling by dynamic ESCRT-III polymers, Trends Cell Biol. 31, 856 (2021).

[19] J. McCullough and W. I. Sundquist, Membrane remodeling: ESCRT-III filaments as molecular garrotes, Curr. Biol. 30, R1425 (2020).

[20] F. Hurtig, T. C. Q. Burgers, A. Cezanne, X. Jiang, F. Nico Mol, J. Traparić, G. Tarrason-Risa, A. A. Pulshen, L. Harker-Kirschneck, A. Šarić et al., The patterned assembly and stepwise Vps4-mediated disassembly of composite ESCRT-III polymers drives archael cell division, bioRxiv 2022.09.16.508273 (2022).

[21] X. Jiang, L. Harker-Kirschneck, C. Vanhille-Campos, A.-K. Pfitzner, E. Lominadze, A. Roux, B. Baum, and A. Šarić, Modelling membrane reshaping by staged polymerization of ESCRT-III filaments, PLoS Comput. Biol. 18, 1 (2022).

[22] L. Harker-Kirschneck, B. Baum, and A. Šarić, Changes in ESCRT-III filament geometry drive membrane remodelling and fission in silico, BMC Biol. 17, 82 (2019).

[23] J. Schönberg, I. H. Lee, J. H. Iwasa, and J. H. Hurley, Reverse-topology membrane scission by the ESCRT proteins, Nat. Rev. Mol. Cell Biol. 18, 5 (2017).

[24] J. McCullough, A. K. Clippinger, N. Talledge, M. L. Skowrya, M. G. Saunders, T. V. Naismith, L. A. Colf, P. Afonine, C. Arthur, W. I. Sundquist, P. I. Hanson, and A. Frost, Structure and membrane remodeling activity of ESCRT-III helical polymers, Science 350, 1548 (2015).

[25] W. Helfrich, Elastic properties of lipid bilayers: Theory and possible experiments, Z. Naturforsch. C 28, 693 (1973).

[26] See Supplemental Material at http://link.aps.org/supplemental/10.1103/PhysRevLett.129.268101 for a detailed description of the model and additional data.

[27] R. Dimova, Recent developments in the field of bending rigidity measurements on membranes, Adv. Colloid Interface Sci. 208, 225 (2014).

[28] N. Chiaruttini, L. Redondo-Morata, A. Colom, F. Humbert, M. Lenz, S. Scheuring, and A. Roux, Relaxation of loaded ESCRT-III spiral springs drives membrane deformation, Cell 163, 866 (2015).

[29] L. Harker-Kirschneck, A. E. Hafner, T. Yao, C. Vanhille-Campos, X. Jiang, A. Pulshen, F. Hurtig, D. Hryniuk, S. Culley, R. Henriques et al., Physical mechanisms of ESCRT-III-driven cell division, Proc. Natl. Acad. Sci. U.S.A. 119, e2107763119 (2022).

[30] S. Lata, G. Schoehn, A. Jain, P. Pires, J. Piehler, H. G. Götlinger, and W. Weissenhorn, Helical structures of ESCRT-III are disassembled by VPS4, Science 321, 1354 (2008).

[31] B. A. Davies, I. F. Azmi, J. Payne, A. Shestakova, B. F. Horazdovsky, M. Babst, and D. J. Katzmann, Coordination of substrate binding and ATP hydrolysis in Vps4-mediated ESCRT-III disassembly, Mol. Biol. Cell 21, 3396 (2010).

[32] B. Yang, G. Stjepanovic, Q. Shen, A. Martin, and J. H. Hurley, Vps4 disassembles an ESCRT-III filament by global unfolding and processive translocation, Nat. Struct. Mol. Biol. 22, 492 (2015).

[33] S. Maity, C. Caillat, N. Miguet, G. Sulbaran, G. Effantin, G. Schoehn, W. H. Roos, and W. Weissenhorn, VPS4 triggers constriction and cleavage of ESCRT-III helical filaments, Sci. Adv. 5, aau7198 (2019).

[34] A. K. Cada, M. R. Pavlin, J. P. Castillo, A. B. Tong, K. P. Larsen, X. Ren, A. L. Yokom, F.-C. Tsai, J. V. Shiah, P. M. Bassereau, C. J. Bustamante, and J. H. Hurley, Friction-driven membrane scission by the human ESCRT-III proteins CHMP1B and IST1, Proc. Natl. Acad. Sci. U.S.A. 119, e2204536119 (2022).

[35] J. Schönberg, M. R. Pavlin, S. Yan, M. Righini, I.-H. Lee, L.-A. Carlson, A. H. Bahrami, D. H. Goldman, X. Ren, G. Hummer et al., ATP-dependent force generation and membrane scission by ESCRT-III and Vps4, Science 362, 1423 (2018).

[36] V. Mercier, J. Larios, G. Molinarid, A. Goujon, S. Matile, J. Gruenberg, and A. Roux, Endosomal membrane tension regulates ESCRT-III-dependent intra-lumenal vesicle formation, Nat. Cell Biol. 22, 947 (2020).

[37] J. Lafaurie-Janvore, P. Maiuri, I. Wang, M. Pinot, J. B. Manneville, T. Betz, M. Ballard, and M. Piel, ESCRT-III assembly and cytokinetic abscission are induced by tension release in the intercellular bridge, Science 339, 1625 (2013).
[38] A. Booth, C. J. Marklew, B. Ciani, and P. A. Beales, In vitro membrane remodeling by ESCRT is regulated by negative feedback from membrane tension, iScience 15, 173 (2019).

[39] B. Nepal, A. Sepehri, and T. Lazaridis, Mechanisms of negative membrane curvature sensing and generation by ESCRT III subunit Snf7, Protein Sci. 29, 1473 (2020).

[40] H. G. Franquelim, A. Khmelinskaia, J.-P. Sobczak, H. Dietz, and P. Schwille, Membrane sculpting by curved DNA origami scaffolds, Nat. Commun. 9, 811 (2018).

[41] S. Dey, A. Dorey, L. Abraham, Y. Xing, I. Zhang, F. Zhang, S. Howorka, and H. Yan, A reversibly gated protein-transporting membrane channel made of DNA, Nat. Commun. 13, 2271 (2022).