FAM134B-RHD Protein Clustering Drives Spontaneous Budding of Asymmetric Membranes

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ABSTRACT: Living cells constantly remodel the shape of their lipid membranes. In the endoplasmic reticulum (ER), the reticulon homology domain (RHD) of the reticulophagy regulator 1 (RETR1/FAM134B) forms dense autophagic puncta that are associated with membrane removal by ER-phagy. In molecular dynamics (MD) simulations, we find that FAM134B-RHD spontaneously forms clusters, driven in part by curvature-mediated attractions. At a critical size, as in a nucleation process, the FAM134B-RHD clusters induce the formation of membrane buds. The kinetics of budding depends sensitively on protein concentration and bilayer asymmetry. Our MD simulations shed light on the role of FAM134B-RHD in ER-phagy and show that membrane asymmetry can be used to modulate the kinetic barrier for membrane remodeling.

Cellular membranes are shaped by curvature-inducing proteins, lipid composition, or external mechanical forces. Protein oligomerization and clustering have been associated with membrane curvature induction, budding, and scission. In the endoplasmic reticulum (ER), the reticulophagy regulator 1 (RETR1/FAM134B) localizes into the autophagic puncta associated with the control of ER size and the removal of aggregated proteins by ER-phagy. Molecular modeling and molecular dynamics (MD) simulations revealed that the reticulon homology domain (RHD) of FAM134B (Figure S1) is responsible for membrane curvature sensing and active curvature induction. The MD simulations also showed that membrane deformations are amplified by FAM134B-RHD clustering. However, the exact mechanism and dynamics of protein clustering leading to membrane budding and scission remained unclear.

We found that FAM134B-RHD proteins are capable of reshaping a membrane from a flat to a budded shape (Figure 1A), making Lx an excellent reporter on membrane shape changes. Figure 1B shows the initial flat membrane and final budded state in an MD simulation with n = 9 FAM134B-RHDs and a leaflet asymmetry of ∆N = 300. We had shown earlier that, without proteins, the POPC membrane remained trapped in the flat metastable state for over 7 μs in a system with ∆N/N = 0.304.
and $N = 1634$ (Figure S1A in ref 14), that is, at an asymmetry higher than any of the systems studied here.

We confirmed that membrane reshaping and budding are specific to FAM134B-RHD and not triggered by membrane inclusions alone (Figure 1). As a control, we performed simulations of proteins not associated with membrane remodeling: (i) the transmembrane helical peptide KALP15, whose hydrophobic mismatch resembles that of the transmembrane hairpins of FAM134B-RHD; and (ii) the seven-transmembrane helical G-protein coupled receptor opsin. In these control simulations, the bilayers remained nearly flat despite the high asymmetry of $\Delta N = 300$ (Figure 1C,D). As an additional control, we placed $n = 15$ FAM134B-RHDs into the membrane in reverse topology, that is, with the N and C termini on the lower side of the membrane. In effect, this inverts the sign of the membrane asymmetry to $\Delta N = -300$. Even $n = 15$ FAM134B-RHDs were not sufficient to induce budding at this unfavorable asymmetry (Figure 1E). The protein identity and the insertion topology are thus decisive factors for budding.

Higher asymmetry and protein concentration both accelerate the induction of membrane buds (Figure 2). In simulations with a low leaflet asymmetry, $\Delta N = 100$, we did not observe any major membrane shape changes (Figure 2) at any protein concentration. At this low value of leaflet asymmetry, the excess lipids are accommodated by a minor compression and expansion of the respective leaflets of a flat bilayer. As the FAM134B-RHDs diffused in the membrane plane and formed transient clusters, the associated membrane deformations remained local and did not induce any budding events (Figure 3). However, for $n > 5$ FAM134B-RHDs, we observed spontaneous membrane budding, indicating that a critical number of proteins is required (Figure 3; $\Delta N = 200, 300$). At a high leaflet asymmetry ($\Delta N = 400$), we observed the largest number of spontaneous budding transitions from the flat bilayer, indicating that the kinetic barrier was lowered significantly even at low protein concentrations (Figure 2). For $\Delta N = 400$, we observed a budding event already with a single FAM134B-RHD embedded in the bilayer.

Along with the increase in the number of budding events, the waiting times for budding decreased at increased FAM134B-RHD concentration (Figure 2). Importantly, though, in all cases there was a time lag between the start of the simulations and budding, indicating that even at the

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Figure 1. FAM134B-RHD specifically induces membrane budding in asymmetric membranes. (A) Time traces of the box width $L_x$ for one exemplary budding event with nine FAM134B-RHD (dark green) and three control simulations with nine opsins (purple), 15 KALP15 (red), and 15 FAM134B-RHD in reverse topology (light green). Corresponding snapshots (top and side views) at times 0 and 12 $\mu$s are shown in (B−E). All systems had an asymmetry of $\Delta N = 300$. (B) Nine FAM134 proteins embedded in an asymmetric bilayer induce the formation of a membrane bud within 2 $\mu$s. Simulation snapshots of (C) nine opsin proteins, (D) 15 KALP15 peptides, and (E) 15 FAM134B-RHDs (reverse topology), which do not induce budding of the asymmetric membrane. Embedded proteins are shown in surface representation, lipid phosphate groups in blue, and lipid tails in white. Water and ions are omitted for clarity.

Figure 2. Box width as the reporter of membrane shape changes. Time traces of the box width $L_x$ are shown for MD simulations of $n = 1$ to 13 FAM134B-RHDs (top to bottom) in POPC membranes with leaflet asymmetries of $\Delta N = 100, 200, 300$, and 400 (left to right). The three replicates are distinguished by color.
highest asymmetry and protein concentration budding had to overcome a kinetic barrier.

The shape of the buds and the width of their neck depend both on membrane asymmetry and the number of proteins on the bud (Figures 3, S2, and S3). The tubular shape of the buds formed at a lower asymmetry ($\Delta N = 200$) changed into a more spherical shape with a tighter neck at higher asymmetries ($\Delta N = 300, 400$). The neck tightened in a similar manner as more proteins packed onto the bud, likely compressed as a result of the highly asymmetric membrane footprint of FAM134B-RHD (Figure S5). These membrane shape changes are reflected in the differences in box width after budding (Figure 3B). With increasing asymmetry, the box length decreases more strongly, indicating that a larger membrane area is absorbed into the bud. At the highest asymmetry ($\Delta N = 400$), the bud size is likely limited by self-interactions of the nascent bud across the periodic boundaries of the simulation box (see also Figures S2 and S3).

At a high leaflet asymmetry ($\Delta N = 400$), we observed an alternate route to alleviate membrane stress. In a few replicates, mostly at low protein concentrations, the excess lipids of the dense upper leaflet folded onto themselves to create a bicelle-like protrusion attached to the otherwise flat bilayer (Figure 4C). The resulting drop in the box width was less pronounced than with actual budding (Figures 2 and 4A,B). FAM134B-RHDs localized to the connection between the protrusion and the membrane but did not sort onto the protrusion, unlike the vesicle buds.

FAM134B-RHD clusters act cooperatively to induce budding, as illustrated in Figure 3 for systems with $n = 9$ FAM134B-RHDs and different leaflet asymmetries $\Delta N$. By monitoring the vertical displacement $z$ of the center of mass of individual proteins with respect to the lowest lipid headgroup in the lower leaflet (Figure 3C), we found that seven, five, and three of the nine proteins were directly involved in the budding events at $\Delta N = 200, 300$, and $400$, respectively. The sharp increase in the $z$-positions of these proteins correlates with the increase in membrane height and the contraction of the box (Figure 3B,C). We then noticed that these proteins formed a distinct cluster, whose size had increased until just before budding (Figure 3D). Small lipid number asymmetries, $\Delta N = 200$, required a larger cluster of $\sim 15$ FAM134B-RHD transmembrane hairpins than larger asymmetries, $\Delta N = 300$, where $\sim 10$ hairpins sufficed. For $\Delta N = 400$, already a single FAM134B-RHD could induce budding. The observations that cluster formation preceded budding and that the required cluster size increases with decreasing energetic driving force together indicate that FAM134B-RHD cluster formation is critical for budding.

Interprotein contact maps of FAM134B-RHDs are dominated by strong interactions between transmembrane hairpins, in particular, between their luminal loops (Figure S4). These
interprotein interactions are similar to the intraprotein interactions reported in Bhaskara et al.,7 as seen also in inverted-pyramid-like RHD clusters. The consistently strong interactions between the luminal loops of the transmembrane hairpins promote the formation and stabilization of protein clusters across different protein concentrations and asymmetries.

The critical role of FAM134B-RHD clusters is confirmed by their localization on the emerging bud. As illustrated in a detailed view of the time evolution of FAM134B-RHD clusters during a representative budding event for $\Delta N = 300$ and $n = 9$ (Figure 5), we found that budding is initiated at the site of a spontaneously formed FAM134B-RHD cluster (yellow) consisting of three proteins. Its merger with another cluster of two proteins (green; Figure 5A,B) then triggered the rapid emergence of a membrane bud. The FAM134B-RHD cluster occupied the cusp of the emerging bud and remained there during the entire transition, indicating that the active curvature induction of the emergent cluster lowered the kinetic barrier for budding.

FAM134B-RHD clustering is enhanced by curvature sensing. By tracking the cluster centers with respect to time, we found that FAM134B-RHD alone or in clusters of up to three proteins preferentially diffused toward the emerging bud. Even after the emergence of the bud, the remaining FAM134B-RHD clusters and individual proteins (orange, Figure 5B,C) tend to sort onto the highly curved membrane bud. This curvature sorting is observed in all analyzed systems (Figure 3C). As time progresses, all proteins eventually migrate onto the bud and remain there. Effective attractions between membrane-embedded proteins mediated by elastic deformations of the bilayer have been studied by continuum theoretical models, molecular simulations, and experiments. Protein-induced curvature fields can exert long-range attractive forces to enable self-organization into clusters.7,16

Sensors of membrane curvature and mechanical drivers of budding are the wedge-shaped transmembrane hairpins and their flanking amphipathic helices embedded in the upper leaflet.7 As a consequence of the resulting asymmetric shape, a membrane-inserted FAM134B-RHD displaces $\sim 20$ POPC lipids from the upper leaflet than from the lower leaflet (Figure S5). The wedging associated with this asymmetric
footprint appears to amplify the stress caused by membrane asymmetry in our simulations. As we could show, by combining these two effects and by concentrating them locally in the membrane at a FAM134B-RHD cluster, the kinetic barrier to budding can be overcome on an MD time scale. In particular, transmembrane hairpin clustering precedes the budding transitions (Figure 3D). In vitro deletion and liposome remodeling experiments confirmed the importance of interhairpin interactions in curvature induction and sensing.7

The curvature-sensing and curvature-inducing properties of FAM134B-RHD are instrumental to the organization of autophagic receptors in the peripheral ER. We found (i) that FAM134B-RHDS tend to cluster in highly curved regions of the membrane and (ii) that these clusters aid in the spontaneous formation and stabilization of large-scale membrane buds. Both observations are relevant to the biology of FAM134B-RHD in ER-phagy. First, our observation of curvature-mediated FAM134B-RHD clustering is consistent with its segregation into the curved regions of the ER.5,7 The formation of FAM134B-RHD clusters is also consistent with the localization of FAM134B-RHD in autophagic puncta during ER-phagy, as seen in immunofluorescence experiments.7 The reported oligomerization of FAM134B-RHD in ER-phagy7 is in the range of cluster sizes seen here to drive membrane remodeling. Second, our observation of FAM134B-RHD-induced membrane budding explains the “membrane shredding” action of FAM134B-RHD, which results in the formation of tiny vesicular structures.5,7 Highlighting the biological relevance of FAM134B-RHD in ER-phagy, we note that Zika and Dengue viruses proteolytically target FAM134B-RHD to evade host responses during infection.18

Other effects such as changes in membrane tension could also affect the kinetic barrier for budding. We found in our earlier study14 that negative tension (i.e., lateral compression) was effective in triggering a membrane budding from flat asymmetric membranes. On the basis of this finding and membrane elastic theory, we expect that positive in-plane tension will raise the barrier. Accordingly, cells might additionally modulate membrane tension locally within the ER, for example, through an attachment to cytoskeletal elements.

On the basis of our findings for FAM134B-RHD, we expect MD simulations with asymmetric membranes to be useful to screen also other proteins for a possible curvature induction and sensing properties. The power to modulate the energetic driving force and the kinetic barrier for membrane shape changes appears to be exploited also in living cells, where asymmetry-creating lipid flippases have been implicated in budding processes. The observed interplay between membrane asymmetry and curvature-inducing proteins has important biological implications on how cells can regulate and induce the formation of membrane buds.

■ ASSOCIATED CONTENT

* Supporting Information
The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.jpclett.1c00031.

Extended methods (PDF)

MD simulation trajectory of FAM134B-induced membrane budding event (MPG)

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Notes
The authors declare no competing financial interest.

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