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Curvature Sensing by a Viral Scission Protein

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Supporting Information

ABSTRACT: Membrane scission is the final step in all budding processes wherein a membrane neck is sufficiently constricted so as to allow for fission and the release of the budded particle. For influenza viruses, membrane scission is mediated by an amphipathic helix (AH) domain in the viral M2 protein. While it is known that the M2AH alters membrane curvature, it is not known how the protein is localized to the center neck of budding virions where it would be able to cause membrane scission. Here, we use molecular dynamics simulations on buckled lipid bilayers to show that the M2AH senses membrane curvature and preferentially localizes to regions of high membrane curvature, comparable to that seen at the center neck of budding influenza viruses. These results were then validated using in vitro binding assays to show that the M2AH senses membrane curvature by detecting lipid packing defects in the membrane. Our results show that the M2AH senses membrane curvature and suggest that the AH domain may localize the protein at the viral neck where it can then mediate membrane scission and the release of budding viruses.

Influenza virus budding requires a precise stepwise alteration of membrane curvature leading to the formation of a membrane bud that is attached to the host cell plasma membrane through a small membrane neck. Release of the budding virion requires membrane scission, which is mediated by an amphipathic helix (AH) domain in the influenza virus M2 protein. The influenza virus M2 protein is a 97-amino acid homotetrameric protein that contains a membrane proximal AH with an extended cytoplasmic tail.

Membrane insertion of the M2AH is sufficient to alter membrane curvature and cause budding in vitro and in vivo. However, to mediate membrane scission, the M2AH needs to specifically bind and insert into the membrane at the constricted neck of the budding virus.

Influenza viruses bud from lipid raft domains on the apical plasma membrane of infected cells. This budding localization is thought to be mediated by the intrinsic raft localization of the viral surface proteins, Hemagglutinin and Neuraminidase. Because of the length of its transmembrane domain, the M2 protein preferentially sorts to the bulk plasma membrane domain and is recruited to the periphery of the lipid raft budding domains only through interactions with the viral M1 matrix protein. This interaction places M2 near the constricted neck of the budding virion; however, scission requires more precise localization that would place the M2AH at the center of the membrane neck. This study investigates the mechanism of M2AH localization and shows that the M2AH senses membrane curvature and is preferentially located at the highly curved center of a constricted membrane neck where subsequent insertion of the AH would be sufficient to cause membrane scission.

The first 16 amino acids of the M2 cytoplasmic tail form a membrane-parallel AH domain that inserts into the membrane (Figure 1).

During virus budding, the M2 protein is recruited to the periphery of the assembly sites. However, to cause membrane scission, M2 needs to localize to the midpoint of the cytoplasmic face of the highly curved budding viral neck, a region that can be topologically depicted as a catenoid (Figure 2A).

Therefore, to determine the specific localization of M2AH, we performed course-grained molecular dynamics (MD) simulations on buckled lipid bilayers. These are planar lipid bilayers that are compressed in one dimension until the bilayers deform. These bilayers show regions of positive, negative, and neutral curvature following the arc length parameter $S$, allowing for the determination of any M2AH curvature binding preference (Figure 2B–D).

The simulations show that the M2AH rapidly binds to membranes and inserts at or below the lipid headgroups (Figure 2D). The lipid-bound M2AH then preferentially sorts.

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Figure 2. MD simulation of M2AH on buckled lipid bilayers. (A) Catenoid structure representative of the neck of a budding virion. (B) Representative top-view (top) and side-view (bottom) snapshots of the MD simulation of the M2AH on lipid bilayers, buckled in the Lx dimension. Lipid headgroups are colored pink and acyl chains gray. Lipid tail ends are shown as orange spheres, and the M2AH is colored red. Indicated vectors represent the orientation of the M2AH on the bilayer. The side view also shows the Euler buckling profile (blue hashed line) fitted to the bilayer midplane and the corresponding arc length parameter S ∈ [0, 1], which parametrizes the position along the curved membrane midplane (C), with s = 0.5 being the most positively curved position. (D) Side-view snapshot of one location of the M2AH (red) in the buckled bilayer, with the lipid headgroup phosphates shown as black dots.

to regions of high positive membrane curvature (Figure 3A,B and Movies S1 and S2), such as those seen at the narrowest point of the budding neck. While the peptide associated with a range of positively curved membrane domains, the association was the strongest with increased curvature (Figure 3B).

Analysis of the free energy of binding showed that the energy required for binding decreases as the curvature increases (Figure 3C). The binding approached an energy minimum at a radius of curvature comparable to that seen in a 20 nm diameter vesicle; however, absolute quantification would likely require an all-atom simulation. Interestingly, the simulations did not reveal a clear preferred orientation (θ) of the M2AH in the plane of the buckled bilayer surface (Figure 3D). This resembles earlier results for the antimicrobial peptide magainin ⋆ and suggests that the M2AH may be an isotropic curvature sensor, capable of identifying curved membranes regardless of the orientation of the helix relative to the direction of curvature. Isotropic curvature sensing would be of particular benefit for M2 protein localization as each AH domain in the full length tetrameric protein is oriented at a 90° angle from each other (Figure 1A).

To confirm the MD simulation data, we next performed a liposome binding assay using a fluorescent M2AH peptide. Unilamellar vesicles were made in a range of sizes, which were verified by dynamic light scattering (Figure S1). Vesicles were made without anionic lipids to ensure that charge interactions do not mask potential curvature sensing, though this masking was not observed in the simulations. As shown in Figure 4A, the M2AH binds to vesicles that have a diameter between 35 and 3135 nm. The level of binding is slightly reduced for larger vesicles >150 nm in diameter (which possess correspondingly less positive membrane curvature) and significantly enhanced for small vesicles 35 nm in diameter (which have more positive membrane curvature) (Figure 4A).

There is good correlation between the binding results and the MD simulations, both of which show enhancement of binding with increasing membrane curvature (Figures 3B and 4A). In addition, recent bicelle NMR experiments have shown that the M2AH domain can associate with highly curved bicelle edges, further supporting the ability of the M2AH to sense membrane curvature.

Binding results were confirmed using circular dichroism spectroscopy (CD) analysis of peptide secondary structure. The M2AH forms an α-helix upon membrane binding, and thus, the amount of α-helix observed in the structure is related to membrane binding activity. CD analysis showed an increasing percentage of α-helix formation as vesicle size decreased, reaching a maximum when the domain bound to 35 nm vesicles, though in all cases the M2AH was at least 70% α-helix, indicating that enhanced curvature facilitates but is not necessary for structuring of the AH domain (Figure 4B). Together, these results indicate that the M2AH strongly associates with highly curved membranes, such as would be seen at the neck of budding viruses.

Many AHs, such as those found in N-Bin-Amphiphysin-Rvs (N-BAR) domains and amphipathic lipid packing sensor (ALPS) motifs, are capable of sensing membrane curvature. ⋆, 7,9 ALPS motifs contain bulky hydrophobic residues on one face of

Figure 3. M2AH senses membrane curvature. (A) Heat map of the center of mass distribution of the M2AH during the MD simulation, superimposed on the lipid headgroup density (blue level curves). (B) Localization of the M2AH to regions of different curvature during the simulation, shown as a probability distribution and reflecting the arc length parameter S. (C) Binding free energy at the peptide center of mass on buckled bilayers as a function of the radius of curvature at each given point. (D) Distributions of the in-plane orientation (θ) of single M2AH peptides on bilayers. Because of symmetry in the system, the distributions are periodic (180°). Minor asymmetry arises from statistical fluctuations due to finite sampling, but no additional directional bias is present.

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Thus, these results suggest that the M2AH identifies membrane curvature by sensing lipid packing defects, similar to ALPS motifs.

Narrow preconstricted membrane necks occur at the junction between a budding influenza virion and the host cell, and infection of cells with influenza viruses that do not express the M2 protein results in the formation of stalled membrane buds that fail to undergo membrane scission. This suggests that M2 is responsible for completing scission at the narrow neck that forms at the site of viral budding. TEM analysis of the stalled buds, formed on the plasma membrane during a ΔM2 influenza virus infection, shows a constricted membrane neck with an average radius of curvature at the midpoint of the neck of 19.84 ± 7.91 nm (Figure 5).

These results agree with the results of our liposome binding experiments that showed efficient binding to 35 nm vesicles (possessing a 17.5 nm radius of curvature) and with our molecular dynamics simulations, which showed an increasing level of binding with decreasing vesicle size (Figures 3B and 4A).

This suggests that the M2AH sorts to regions of high membrane curvature similar to that seen at the center of the membrane neck formed during influenza virus budding and is consistent with the observed in vivo localization of the M2 protein during virus budding. When the M2 protein is present at the membrane neck during wild-type influenza virus budding, membrane scission is completed and the influenza virion is released.

The M2 protein is thought to localize to the boundary between the viral budding domain and the bulk plasma membrane, which would position the M2 protein near the neck of the budding virus, though it is not clear if this localization is mediated through M2–M1 interactions or because of intrinsic biophysical properties of the full length protein. However, to complete membrane scission, M2 needs to act at the midpoint of the constricted neck. Here we show that the M2AH senses membrane curvature (Figure 3B,C) by detecting lipid packing defects occurring at highly curved membranes (Figure 4A). This curvature sensing allows the level of M2AH–membrane binding to increase as the membrane radius of curvature decreases (Figure 3C). This allows for strong binding of the M2AH to membranes with a radius of curvature between 10 and 20 nm, similar to what is seen at the center neck of budding influenza virions (Figure 5).

Thus, M2AH curvature sensing may facilitate placement of the helix at the center of the preconstricted membrane neck and could ensure the maintenance of this localization during further constriction. At the neck midpoint, M2AH membrane insertion, and induction of curvature, may be sufficient to cause membrane scission. Thus, the release of influenza viruses likely
requires a specific combination of curvature sensing and curvature induction by the M2AH domain.

■ ASSOCIATED CONTENT

* Supporting Information
The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.biochem.6b00539.

Two figures and experimental methods (PDF)
Movie 1 (MP4)
Movie 2 (MP4)

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Notes
The authors declare no competing financial interest.

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■ REFERENCES

(1) Rossman, J. S., Jing, X., Leser, G. P., and Lamb, R. A. (2010) Cell 142, 902–913.
(2) Rossman, J. S., Jing, X., Leser, G. P., Balannik, V., Pinto, L. H., and Lamb, R. A. (2010) J. Virol. 84, 5078–5088.
(3) Schnell, J. R., and Chou, J. J. (2008) Nature 451, 591–595.
(4) Tian, C., Gao, P. F., Pinto, L. H., Lamb, R. A., and Cross, T. A. (2003) Protein Sci. 12, 2597–2605.
(5) Schmidt, N. W., Mishra, A., Wang, J., DeGrado, W. F., and Wong, G. C. (2013) J. Am. Chem. Soc. 135, 13710–13719.
(6) Gómez-Llobregat, J., Elias-Wolf, F., and Lindén, M. (2016) Biophys. J. 110, 197–204.
(7) Drin, G., and Antonny, B. (2010) FEBS Lett. 584, 1840–1847.
(8) Wang, T., and Hong, M. (2015) Biochemistry 54, 2214–2226.
(9) Bhatia, V. K., Madsen, K. L., Bolinger, P. Y., Kunding, A., Hedegard, P., Gether, U., and Stamou, D. (2009) EMBO J. 28, 3303–3314.
(10) Bigay, J., Casella, J. F., Drin, G., Mesmin, B., and Antonny, B. (2005) EMBO J. 24, 2244–2253.
(11) Nath, S., Dancourt, J., Shtein, V., Puente, G., Fong, W. M., Nag, S., Bewersdorf, J., Yamamoto, A., Antonny, B., and Melia, T. J. (2014) Nat. Cell Biol. 16, 415–424.
(12) Garcia-Saez, A. J., Chiantia, S., and Schwille, P. (2007) J. Biol. Chem. 282, 33537–33544.
(13) Rossman, J. S., and Lamb, R. A. (2011) Virology 411, 229–236.
(14) Schroder, C. (2010) Subcell. Biochem. 51, 77–108.
(15) Nayak, D. P., Balogun, R. A., Yamada, H., Zhou, Z. H., and Barman, S. (2009) Virus Res. 143, 147–161.