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A Budding-Defective M2 Mutant Exhibits Reduced Membrane Interaction, Insensitivity to Cholesterol, and Perturbed Interdomain Coupling

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Abstract

Influenza A M2 is a membrane-associated protein with a C-terminal amphipathic helix that plays a cholesterol-dependent role in viral budding. An M2 mutant with alanine substitutions in the C-terminal amphipathic helix is deficient in viral scission. With the goal of providing atomic-level understanding of how the wild-type protein functions, we used a multipronged site-directed spin labeling electron paramagnetic resonance spectroscopy (SDSL-EPR) approach to characterize the conformational properties of the alanine mutant. We spin-labeled sites in the transmembrane (TM) domain and the C-terminal amphipathic helix (AH) of wild-type (WT) and mutant M2, and collected information on line shapes, relaxation rates, membrane topology, and distances within the homotetramer in membranes with and without cholesterol. Our results identify marked differences in the conformation and dynamics between the WT and the alanine mutant. Compared to WT, the dominant population of the mutant AH is more dynamic, shallower in the membrane, and has altered quaternary arrangement of the C-terminal domain. While the AH becomes more dynamic, the dominant population of the TM domain of the mutant is immobilized. The presence of cholesterol changes the conformation and dynamics of the WT protein, while the alanine mutant is insensitive to cholesterol. These findings provide new insight into how M2 may facilitate budding. We propose the AH–membrane interaction modulates the arrangement of the TM helices, effectively stabilizing a conformational state that enables M2 to facilitate viral budding.

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ASSOCIATED CONTENT

Supporting Information

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Extended Materials and Methods; saturation recovery curves of M2-WT and M2-5Ala site 57; T1 and jx values obtained from fits of SR data; PS EPR oxygen accessibility data; hypothetical DEER distance distribution of a homo-tetrameric protein; DEER data of M2-WT and M2-5Ala site 57; comparison of SSNMR model (PDB 2L0J) with experimental DEER data; CD spectra and secondary structure analysis; CW EPR spectra of additional AH sites in + cholesterol membranes; saturation recovery curves of M2-WT and M2-5Ala site 59 in + cholesterol membranes (PDF)

The authors declare no competing financial interest.
Antagonizing the properties of the AH that enable interdomain coupling within M2 may therefore present a novel strategy for anti-influenza drug design.

**Abstract**

The influenza A virus is a persistent source of morbidity and mortality worldwide.\(^1,2\) An atomic-level understanding of how the virus assembles and buds from the host cell plasma membrane could serve as the basis for new antiviral strategies inhibiting influenza infectivity.\(^3\) Recent studies suggest that the viral coat and core proteins work in concert to promote budding.\(^3\) Here, we examine the conformational properties of one member of the budding machinery, the M2 protein, as well as a mutant deficient in viral budding.

M2 is a homotetrameric, 97-residue membrane-bound protein that orchestrates several essential events in the influenza A infection cycle.\(^3,4\) The determinants for the separate functions of M2 are encoded in different protein domains (Figure 1A). Residues 25–46 associate into a homotetrameric, \(\alpha\)-helical transmembrane (TM) domain with ion channel activity necessary for viral uncoating.\(^4\) Less is known about the structure of the N-terminal ectodomain (residues 1–25) and C-terminal cytoplasmic tail (residues 46–97). However, numerous recent studies have explored the role of the C-terminal domain in influenza morphology and infectivity.\(^5–10\)

In particular, recent research efforts have examined the role the M2 cytoplasmic domain plays in viral budding.\(^12–14\) In vitro assays revealed that the full-length M2 protein, as well as just the region spanning residues 46–62, is sufficient to generate the type of curvature necessary for constricting the budding viral neck\(^15\) and induce budding in giant unilamellar vesicles.\(^12\) Several biophysical techniques, including solution NMR,\(^16,17\) solid-state NMR\(^18,19\) (SSNMR), and electron paramagnetic resonance\(^20,21\) (EPR), have demonstrated that M2 residues 47–62 form an amphipathic helix associated with the membrane surface.

Amphipathic helices (AHs) promote membrane remodeling in several systems,\(^22–24\) presumably by inserting into the membrane and reorienting lipids to induce curvature. Several groups have shown that replacing five hydrophobic residues of the M2 AH with alanine—F47A, F48A, I51A, Y52A, and F55A (Figure 1C)—abolishes M2 curvature-generation and budding activity in vitro and leads to a budding-defective morphology in vivo without affecting M2 ion channel function.\(^12–15,25,26\) The penta-alanine mutant, which we designate “M2–5Ala”, offers a valuable opportunity to characterize the structural and dynamic properties of M2 necessary for viral budding.
Several constructs of the M2 protein have been studied by a range of high-resolution biophysical techniques, including X-ray crystallography,\textsuperscript{27,28} \textit{solution NMR,\textsuperscript{16,17}} and \textit{SSNMR.\textsuperscript{18,29–32}} The method we employ here, site-directed spin labeling EPR spectroscopy (SDSL-EPR), is a powerful tool for characterizing membrane protein structure and dynamics\textsuperscript{33,34} and offers complementary information to previously published structural work on the M2 protein. A strength of SDSL-EPR is the ability to study membrane proteins embedded in lipid bilayers, which is particularly important for characterizing the membrane-remodeling behavior of M2. SDSL-EPR methods can detect mobility and conformational exchange events on time scales not accessible to other spectroscopic techniques,\textsuperscript{35} distances in the range of ~12–60 Å,\textsuperscript{34} and topology of a protein with respect to the membrane.\textsuperscript{33} Our group has used SDSL-EPR to characterize the sensitivity of the M2 TM and AH domains to the surrounding environment.\textsuperscript{20,21,36–39} We demonstrated that the M2 AH exists in conformational equilibrium and undergoes cholesterol-dependent conformational exchange.\textsuperscript{39} We hypothesized that this conformational exchange is relevant for M2 scission function at the edge of the viral budozon,\textsuperscript{12,13} which is enriched in cholesterol.\textsuperscript{40,41}

Here, we seek a structural understanding of the M2-5Ala mutant with the goal of gaining insight into how the wild-type (M2-WT) protein functions. In principle, a detailed understanding of the viral budding process could aid in the development of novel strategies for anti-influenza drug design.

To investigate how the mutations within M2-5Ala impact conformation and dynamics, we spin-labeled sites in both the C-terminal TM and AH domains, and collected SDSL-EPR data. We demonstrate that compared to M2-WT, the dominant population of M2-5Ala is more dynamic, shallower in the membrane, and has an altered quaternary arrangement of the C-terminal domain. Whereas M2-WT experiences notable changes in conformation, dynamics, and membrane topology upon the addition of cholesterol, M2-5Ala remains insensitive to the presence of cholesterol. Our data are consistent with the AHs sensing the properties of the membrane and regulating the arrangement of the TM helices, effectively stabilizing an overall shape of the protein optimal for viral budding.

\section*{MATERIALS AND METHODS}

\subsection*{Purification, Spin Labeling, and Membrane Reconstitution of Full-Length M2-WT and M2–5Ala Cysteine Mutants.}

Cysteine substitutions were introduced into a cysteineless background plasmid based on the A/Udorn/72 sequence,\textsuperscript{42} referred to as “M2-WT”, using site-directed mutagenesis. The F47A, F48A, I51A, Y52A, and F55A mutations were introduced into the M2-WT plasmid to generate the “M2-5Ala” plasmid, which was used for further cysteine mutagenesis. Verified single-cysteine plasmids were transformed into C43 competent cells for protein expression. Expression, purification, and spin labeling were performed according to previously published protocols\textsuperscript{21} with minor modifications described in the Supporting Information Materials and Methods. The purity of protein was verified using sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE).
We spin-labeled sites in the C-terminal TM and AH domain of M2-WT and M2-5Ala (Figure 1). To probe properties of the TM, we labeled site L43. This residue faces away from the pore of the channel, and previous electrophysiology studies of full-length M2 demonstrated that cysteine mutagenesis at this site does not significantly affect ion channel behavior.43,44 To probe the properties of the AH, we labeled a site on each face of the helix (H57 and L59).

Protein was reconstituted into 4:1 1-palmitoyl-2-oleoyl-sn-glycero-3-phosphocholine-/palmitoyl-2-oleoyl-sn-glycero-3-phospho-1-rac-glycerol (POPC/POPG; “− cholesterol”) and 56:14:30 POPC/POPG/cholesterol (“+ cholesterol”) lipid bilayers as previously described.21 This membrane system was chosen for its use in in vitro budding assays12 and previous SDSL-EPR studies of the M2 protein.20,21,37,39

EPR Spectroscopy and Data Analysis.

**Continuous Wave EPR Spectra.**—Continuous wave EPR (CW EPR) spectra were recorded at room temperature on an X-band Bruker EMX spectrometer equipped with an ER4123S resonator. Proteoliposomes were prepared at a peptide/lipid ratio of 1:500 in 50 mM Tris, pH 7.8, 100 mM KCl, 1 mM EDTA buffer. Samples were placed in 0.6 mm internal diameter, 0.9 mm length gas permeable capillary tubes (L&M EPR Supplies, Milwaukee, WI). CW EPR spectra were acquired using 2 mW incident microwave power, 1 G field modulation amplitude at 100 kHz, and 150 G sweep width.

**Paramagnetic Accessibility Measurements.**—Power saturation experiments were used to detect the oxygen accessibility of spin labels.45 Experiments were run at room temperature on an X-band Bruker EMX spectrometer equipped with an ER4123S resonator. Proteoliposomes were prepared at a peptide/lipid ratio of 1:500 in 50 mM Tris, pH 7.8, 100 mM KCl, 1 mM EDTA buffer. Samples were placed in 0.6 mm internal diameter by 0.9 mm length gas permeable capillary tubes (L&M EPR Supplies, Milwaukee, WI). Power saturation experiments were performed under nitrogen gas or equilibrated with ambient air. EPR spectra were collected over eight power levels for nitrogen power saturation experiments. For oxygen power saturation experiments, spectra were collected over 16 or 19 power levels for sites with moderate or high accessibility to oxygen, respectively. Data were fit to obtain ΔP1/2 parameters as described previously.20

**Saturation Recovery Measurements.**—Saturation recovery EPR (SR EPR) experiments were carried out at the National Biomedical EPR Center (Milwaukee, WI) on a home-built instrument operating at X-band (9.4 GHz) equipped with a loop-gap resonator, as previously described.39 The resulting SR signals were fitted with single- and biexponential decay functions using Origin software (OriginLab, Northampton, MA). Bimolecular collision rates with oxygen, Wx(O2), were calculated from spin–lattice relaxation rates (We) according to the relationship Wx(O2) = We(O2) − We(N2), where We = 1/2T1.

**Double Electron–Electron Resonance Measurements.**—Double electron–electron resonance (DEER) experiments were carried out at the Ohio Advanced EPR Laboratory at Miami University, Ohio, on a Bruker ELEXSYS E580 spectrometer equipped with a
SuperQ-FT pulse Q-band system with a 10 W amplifier and EN5107D2 resonator. DEER samples were prepared as lipodisq nanoparticles with a peptide/lipid ratio of 1:300, as previously described. All DEER samples were prepared at a spin label concentration of 100−130 μM. Glycerol (30% w/w) was used as a cryoprotectant. The sample was loaded into a 1.1 mm inner diameter quartz capillary (Wilmad LabGlass, Buena, NJ) and mounted into the sample holder inserted into the resonator. DEER data were collected using the standard four-pulse sequence \[ \frac{\pi}{2} \nu_1 - \tau_1 - (\pi) \nu_2 - (\tau_1 + \tau_2 - t) - (\pi) \nu_1 - \tau_2 - \text{echo} \] at Q-band with a probe pulse width of 10/20 ns, pump pulse width of 24 ns, 80 MHz of frequency difference between probe and pump pulse, shot repetition time determined by spin–lattice relaxation rate (T_1), 100 echoes/point, and 2-step phase cycling at 80 K collected out to ~2.0 μs for overnight data acquisition time (~12 h). DEER data were analyzed using DEER Analysis 2015. The distance distributions, P(r), were obtained by Tikhonov regularization in the distance domain, incorporating the constraint P(r) > 0. A homogeneous two-dimensional model was used for background correction. The regularization parameter in the L curve was optimized by examining the fit of the time domain.

Circular Dichroism Spectroscopy and Data Analysis.

CD spectra were collected on an Aviv model 435 circular dichroism spectrometer at 25 °C. Samples had a protein/lipid ratio of 1:200 and were reconstituted into bicelles. Bicelles with a q = 0.5 were prepared through the addition of 1,2-dihexanoyl-sn-glycero-3-phosphocholine (DHPC) to protein reconstituted in POPC/POPG 4:1 liposomes. The buffer used to collect CD data was 10 mM Tris, pH 7.8, 20 mM KCl, 1 mM EDTA. A 35 μL aliquot of sample was loaded into 0.011 cm path length demountable quartz cuvette. Spectra were collected from 280 to 190 nm with a 1.0 nm step size. The averaging time at each point was 10.0 s, and the bandwidth was 1.0 nm. Three scans were collected and averaged. A bicelle spectrum collected in the absence of protein was used to correct for background due to lipids and buffer. Spectral data collected directly from the instrument were converted from units of millidegrees to per residue molar absorption units (Δε, in units of M$^{-1}$ cm$^{-1}$). Secondary structure estimates were obtained from the DichroWeb server using the CDSSTR analysis program and a reference set appropriate for membrane proteins.

RESULTS

Alanine Mutations Lead to Large Changes in Mobility.

TM is Immobilized in M2−5Ala Relative to M2-WT.—To probe the impact of the C-terminal alanine mutations on the TM domain, we spin-labeled a site located at the end of the TM helix just prior to the start of the AH (L43; Figure 1). The spectral line shapes of spin-labeled residues provide insight into conformational dynamics that reflect both backbone motion as well as secondary and tertiary contacts.

In Figure 2 (top), we show an overlay of the CW EPR spectra of the TM site 43 in both M2-WT (black) and M2−5Ala (red). The spectra are a superposition of a broad, immobilized component (i), and a sharper, mobile component (m). In a previous study of M2, we characterized the multicomponent nature of M2-WT CW line shapes and established that
they arose from an equilibrium involving two different conformational substates of the protein.\textsuperscript{39} The M2-5Ala spectrum shown in Figure 2 exhibits a more prominent immobile (i) peak than the corresponding M2-WT site, indicating that the equilibrium of the mutant TM helices is shifted markedly toward an immobile population.

**AH Is More Dynamic in M2-5Ala Relative to M2-WT.**—To probe the properties of the AH region of M2-5Ala, we spin-labeled a site on each face of the AH (H57 and L59; Figure 1). In Figure 2, we show an overlay of the CW EPR spectra of M2-WT and M2-5Ala for AH sites 57 and 59. For both these AH sites, the mobile spectral component (m) is considerably more pronounced in M2-5Ala. Thus, the equilibria of the TM and AH in M2-5Ala shift in opposite directions relative to M2-WT: while the immobile component is more populated in the M2-5Ala TM, in the M2-5Ala AH, the immobile component is less populated.

**Both M2-5Ala and M2-WT are in Conformational Equilibrium.**

In a previous study of M2, we used SR EPR to demonstrate that multicomponent CW line shapes, similar to those shown in Figure 2, arose from an equilibrium involving two different conformational substates of the M2 protein.\textsuperscript{39} In our earlier work, we used a truncated construct, M2-WT(23–60). To confirm that conformational exchange is indeed a property of the full-length protein, and not an artifact of using a truncation, we collected SR data for both full-length M2-WT as well as M2-5Ala. In an SR experiment, an intense saturating pulse is applied at the frequency of the central resonance line of the nitroxide. In a spin label system undergoing fast rotameric exchange, the signal recovery is monoexponential. In contrast, spin labels in conformational exchange undergo a biexponential recovery.\textsuperscript{35,39}

As shown in Figure 3, the SR curves for both M2-WT (Figure 3A) and M2-5Ala (Figure 3B) are best characterized by a biexponential recovery. Furthermore, the $T_1$ values obtained from the fits are consistent with protein conformational exchange (Table S1).\textsuperscript{35} Biexponential recovery consistent with protein conformational exchange was also obtained for site 57 in both M2-WT and M2-5Ala (Figure S1; Table S1).

**The Dominant Population of the M2-5Ala AH Is further from the Membrane Surface.**

It has been hypothesized that the loss of budding function for M2-5Ala results from decreased interaction of the AH with the membrane surface due to the reduction in the hydrophobic moment of the AH helix (Figure 1B,C).\textsuperscript{12,13} To test this hypothesis, we probed the membrane depth of the M2-5Ala AH using two complementary EPR methods, SR and power saturation.

The spin–lattice relaxation rates measured using SR techniques can be modulated by the presence of paramagnetic relaxation agents such as oxygen. The relaxation rate of the nitroxide spin label, $W_\nu$, is proportional to the collision rate of the label with the relaxation agent.\textsuperscript{35} Oxygen is a lipophilic species that partitions in a gradient across the membrane, with the highest $O_2$ concentration at the center of the bilayer. An accessibility constant, $j_x$, that reports on the membrane accessibility of the spin label can be obtained by measuring $W_\nu$ at several different $O_2$ concentrations and calculating the slope of a linear best-fit line.\textsuperscript{35}
The insets of Figures 3 and S1 show the dependence of the spin–lattice relaxation rates on the concentration of oxygen for M2-WT and M2-5Ala. The $J_\chi$ value of the M2-5Ala fast component is notably smaller than that of M2-WT (Figure 3C; Table S1), indicating that the M2-5Ala mobile conformation has AHs shallower in the membrane than the corresponding M2-WT conformation. The $J_\chi$ values of the M2-WT and M2-5Ala slow components are the same within error (Figure 3C; Table S1), indicating that the M2-WT and M2-5Ala immobile conformations share similar membrane topology.

To further probe the membrane topology of the AH domains, we performed power saturation (PS) EPR experiments. The PS method measures changes in the relaxation efficiency of a spin label in the presence of paramagnetic reagents. Because oxygen partitions into the hydrophobic environment of the bilayer, the collision rate of the spin label with $O_2$, quantified by the $\Delta P_{1/2}(O_2)$ parameter, serves as a measurement of membrane accessibility complementary to SR. A comparison of the PS profiles of M2-WT and M2-5Ala sites is shown in Figure S2. Consistent with SR EPR results showing that the most populated conformation of M2-5Ala is shallower in the membrane than M2-WT, the M2-5Ala sites exhibit a lower oxygen accessibility than the corresponding M2-WT residues.

M2-5Ala Exhibits Altered Quaternary Arrangement within the Homotetramer.

The data presented above demonstrate that the AHs of M2-WT and M2-5Ala differ both in dynamics and membrane depth. We next explored whether M2-WT and M2-5Ala differ in the arrangement of AHs within the homotetramer.

DEER EPR can measure distances from ~20–60 Å and has been successfully applied to membrane proteins, although resolving distances in homo-oligomeric proteins like M2 has proven to be challenging. A single labeled site in a homotetramer with a single conformation would lead to two measurable distances resulting from subunits that are adjacent and diagonal in the tetramer (Figure S3). With the two distinct conformations of M2-WT identified by SR EPR measurements, we can thus predict four unique distances. However, because of the similar values of distances between symmetry-related sites within the homotetramer as predicted from published M2 structures (Figure S4), a broad overlapped distribution may be expected.

We collected DEER data for M2-WT and M2-5Ala AH sites 57 and 59. As shown in Figure 4 (black lines), the distance distributions of site 59 in M2-WT and M2-5Ala exhibit striking differences. The maximum of the distance distribution shifts from ~47 Å in M2-WT to ~27 Å in M2-5Ala, indicating a large rearrangement of subunits and/or rotation of the AHs within the homotetramer. The lack of well-defined oscillations within the time domain data, typical for membrane proteins in lipid bilayers, prevents the extraction of the four well-defined distances expected for two conformations of a homotetramer. However, site 57 similarly exhibits a shift in DEER distance distributions (from a maximum of ~48 Å in M2-WT to ~34 Å in M2-5Ala; Figure S5), further supporting the existence of a large difference between M2-WT and M2-5Ala in the orientation of their C-terminal domains.

It is instructive to compare our experimental DEER data to a published SSNMR model M2-WT in lipid bilayers (PDB 2L0J). We used mtslWizard to generate conformational
ensembles of the site 57 and 59 spin labels in silico and simulate a distance distribution between the four symmetry-related sites (Figure S4). Overlays of the simulated distance distributions with experimental DEER data reveal marked differences. The simulated distribution for site 59 yields a broad peak at ~20 Å, which is a much shorter distance than the experimentally observed bimodal peaks at ~32 and ~47 Å. One possible reason for the discrepancy between our measured DEER distances and those simulated from the published SSNMR model is that the SSNMR model used a truncated construct M2(22–62), whereas our DEER data are for full-length M2. However, we also observed discrepancies between the SSNMR model and DEER distances in our previously published study on M2-WT(23–60). 39

**Cholesterol Immobilizes M2-WT while M2-5Ala is Insensitive to Cholesterol.**

The M2 protein is proposed to bind cholesterol, 13, 40 but biochemical studies have shown that M2-5Ala does not coimmunoprecipitate with cholesterol in vivo. 13 We compared how M2-WT and M2-5Ala respond in terms of conformation and dynamics to the addition of cholesterol.

In Figure 5A, we show CW EPR spectra of M2-WT and M2-5Ala in 4:1 POPC/POPG bilayers with (red) and without (black) cholesterol. In M2-WT, the addition of cholesterol to the membrane increases the relative intensity of the immobile component for sites 43 and 57. The site 59 line shape in M2-WT shows only minor changes between samples with and without cholesterol. We also spin-labeled three additional M2-WT AH sites (51, 55, and 60) and observed the general pattern that the growth in the immobile population is greatest for C-terminal sites close to the TM domain and becomes attenuated at distal residues (Figure S6). In a previous study using truncated M2-WT(23−−60), we also observed a shift toward a more immobile population of the AH in the presence of cholesterol. 39

M2-5Ala line shapes both with and without cholesterol are shown in Figure 5B. Intriguingly, the 5Ala site 43 line shape is nearly superimposable in the presence and absence of cholesterol. For sites 57 and 59, the intensity of the immobile peak increases only slightly upon the addition of cholesterol, and the mobile population is still predominant.

We performed SR EPR experiments to assess whether the multicomponent nature of the CW line shapes in Figure 5 arises from conformational equilibria. The saturation curves for both M2-WT and M2-5Ala AHs in membranes with cholesterol are best characterized by biexponential fits with two unique spin–lattice relaxation rates (Figure S7), consistent with the existence of conformational exchange.

**M2-5Ala Membrane Topology, Quaternary Arrangement, and CD Profiles are Insensitive to Cholesterol.**

Unlike M2-WT, CW line shapes of M2-5Ala demonstrated no striking changes upon the addition of cholesterol. We next compared M2-WT and M2-5Ala responses to cholesterol in terms of membrane topology, quaternary arrangement, and CD profiles.

In a previous study using truncated M2-WT(23−60), we demonstrated that the AH shifts toward a conformation that is less membrane-buried in the presence of cholesterol. 39 Here
we tested whether this observation was true for full-length M2-WT. As seen for truncated M2, all three of our full-length M2-WT sites (43, 57, and 59) exhibit decreased accessibility to oxygen in the presence of cholesterol (Figure S8). In marked contrast, the oxygen accessibilities of the same sites in M2-5Ala do not greatly change upon the addition of cholesterol (Figure S8), consistent with the highly attenuated ability of M2-5Ala to sense and respond to the presence of cholesterol.

Previously we demonstrated that the AHs in truncated M2-WT(23–60) become more tightly packed within the homotetramer in the presence of cholesterol. The DEER data we collected here demonstrate that full-length M2-WT also becomes more tightly packed in the presence of cholesterol. Figures 4A (site 57) and S5 (site 59) show the distance distributions of M2-WT AH sites in membranes with (red lines) and without (black lines) cholesterol. Adding cholesterol to the membrane results in a shift toward shorter distances for M2-WT, consistent with the population shifting toward the more tightly packed conformation. In contrast, the addition of cholesterol causes only minor changes in the DEER distribution for M2-5Ala (Figures 4B and S5), which provides further evidence of the inability of M2-5Ala to sense and respond to the presence of cholesterol.

The CD profiles of M2-WT and M2-5Ala with and without cholesterol are shown in Figure S9. The CD spectra of both M2-WT and M2-5Ala without cholesterol exhibit two minima centered at 207 and 222 nm, which are characteristic of α-helical secondary structure. Differences in the molar absorption between the M2-WT and M2-5Ala are subtle throughout the range investigated, suggesting that the five alanine mutations do not grossly alter the global secondary structure of the protein. Furthermore, changes in the molar absorption upon the addition of cholesterol to both M2-WT and M2-5Ala are minor, suggesting that global secondary structure does not change in the presence of cholesterol. CD spectral deconvolution shows no marked changes in helical character for either M2-WT and M2-5Ala upon the addition of cholesterol (Table S2).

**DISCUSSION**

M2-5Ala is deficient in viral budding. The inability of M2-5Ala to facilitate viral budding has been hypothesized to arise from the loss of AH–membrane interaction, altered sensitivity to cholesterol, and impaired “cross-talk” between different protein domains. To our knowledge, we have for the first time experimentally tested each of these hypotheses through a detailed atomic-level comparison of the conformation, dynamics, and membrane topology of the full-length M2-WT and M2-5Ala proteins. We discuss each of these hypotheses in turn below. Finally, we describe a model for M2-facilitated viral budding wherein the membrane interaction of the AHs helps stabilize an arrangement of the TM helices necessary for proper budding.

**Reduced Interaction of M2-5Ala with the Membrane Surface.**

A common hypothesis for the loss of function observed with M2-5Ala is decreased interaction of the AH with the membrane surface. Replacing five hydrophobic residues of the AH with alanine reduces the hydrophobic moment of the helix (Figure 1B,C). A recent study demonstrated that a 16-residue M2-5Ala AH peptide did not insert as deeply into the
membrane as the corresponding M2-WT sequence. Results presented here for the full-length protein similarly show that the most highly populated conformation of M2-5Ala is not as closely associated with the membrane surface as M2-WT.

Interaction of AHs with membrane surfaces is critical in regulating curvature-sensing and curvature-inducing activity in a range of systems. Mutations that disturb the hydrophobic properties of AHs have been shown to abolish membrane-remodeling activity. However, the curvature-generating behavior of amphipathic helices in transmembrane proteins has not been as extensively studied.

**Loss of Sensitivity of M2-5Ala to the Presence of Cholesterol.**

Previously, we showed that truncated M2-WT(23–60) undergoes cholesterol-dependent conformational exchange in lipid bilayers. M2-WT protein is proposed to bind cholesterol, but biochemical studies have shown that M2-5Ala does not coimmunoprecipitate with cholesterol in vivo. In this study, we sought to provide atomic-level conformational details to explain how M2-5Ala differs from M2-WT in its response to cholesterol.

We demonstrated that the addition of cholesterol to full-length M2-WT shifts the equilibrium toward the immobile component and changes the quaternary arrangement of the C-terminal domain. In contrast, M2-5Ala exhibits only minor changes in dynamics and quaternary arrangement when exposed to cholesterol.

The impact of cholesterol on M2-WT has been proposed to arise from changes in the membrane surrounding the protein and/or direct binding of cholesterol to the protein. Cholesterol can have a range of impacts on lipid bilayer properties. For example, a number of studies have shown that cholesterol has a condensing effect on membranes, which decreases the ability of membrane-associating helices to penetrate into the lipid headgroup region. In M2-5Ala, the amphipathic helix is further away from the membrane surface and therefore is not as sensitive to changes in bilayer properties. Furthermore, there is some evidence for a specific cholesterol-binding site within the amphipathic helix in the M2 protein. The alanine mutations could perturb the binding site and abolish the direct interaction of the protein with cholesterol.

**Impaired Cross-Talk between Different Domains of M2-5Ala.**

The AH domain of M2 alone can induce membrane curvature and cause budding in giant unilamellar vesicles. Full-length M2-WT, however, has an enhanced ability to generate curvature, consistent with the TM domain enhancing curvature generation through protein shape effects. Our structural work on the full-length M2 protein, both WT and a budding-deficient mutant, provides a unique opportunity to test the hypothesis that the domains of M2 work synergistically to induce curvature.

A cartoon schematic shown in Figure 6 summarizes a model that is consistent with our data. Shown on the left in Figure 6 are the most populated conformations of M2-WT and M2-5Ala in membranes without cholesterol. The M2-5Ala AHs (orange) are further from the membrane surface than the M2-WT AHs (red). Furthermore, M2-5Ala is more tightly packed than M2-WT. Although the mutations in M2-5Ala are in the C-terminal domain, the
differences in the TM domain could arise from the AHs anchoring the TM helices into the membrane surface, effectively stabilizing one of several possible conformational states sampled by the TM helical bundle. A number of studies have shown that M2 TM domain can sample multiple structural substates.29,36,69

Shown on the right in Figure 6 are the most populated conformations of M2-WT and M2-5Ala in the presence of cholesterol. Our model for full-length M2-WT in cholesterol-containing membranes (Figure 6, top right) favors a conformation that is more compact than M2-WT in noncholesterol membranes, consistent with our previously published model of a budding-relevant M2 conformation based on SDSL-EPR data of M2(23–60).39

Our model for the conformation for M2-5Ala in cholesterol-containing membranes (Figure 6, bottom right) is very similar to the conformation of M2-5Ala in membranes without cholesterol (Figure 6, bottom left). The lack of membrane interaction of the AHs of M2-5Ala eliminates interdomain coupling needed for the TM domain to respond to the presence of cholesterol.

We have traced in blue the membrane interaction domains of our models for M2-WT and M2-5A in the presence of cholesterol. M2-WT forms a conical, or wedge-like shape (see also Figure S4). In contrast to M2-WT, we propose M2-5Ala forms a cylindrical shape resulting from the lack of membrane interaction of the AHs. The ability to form a wedge shape was previously hypothesized to play a role in M2’s ability to generate curvature.15 When a transmembrane protein forms a wedge within a lipid bilayer, it can exclude different volumes in the two bilayer leaflets and dynamically induce membrane curvature.24

**SUMMARY**

We have provided new insight into the mechanism of M2-facilitated budding using a multipronged SDSL-EPR approach. Numerous studies have characterized the functional deficiencies of M2-5Ala12–15,25 but the unique contribution of this study is the detailed atomic-level comparison of the conformation, dynamics, and membrane topology of M2-WT and the budding-deficient M2 mutant (M2-5Ala). Our use of full-length protein in membranes has allowed us to test the hypothesis that the AH and TM domains of M2 work synergistically to induce membrane curvature. A model consistent with our data incorporates coupling between the AH and TM domains of M2. We propose interdomain coupling is necessary for the protein to respond to the presence of cholesterol and generate an overall protein shape that facilitates viral budding.15 M2 plays a key role in the replication cycle of the influenza virus. Antagonizing the dynamic properties of the AH that enable interdomain coupling may therefore represent a novel strategy for anti-influenza drug design.

**Supplementary Material**

Refer to Web version on PubMed Central for supplementary material.

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ABBREVIATIONS

SDSL-EPR  site-directed spin labeling electron paramagnetic resonance spectroscopy
SSNMR  solid-state NMR
TM  transmembrane
AH  amphipathic helix
WT  wild-type
POPC  1-palmitoyl-2-oleoyl-sn-glycero-3-phosphocholine
POPG  1-palmitoyl-2-oleoyl-sn-glycero-3-phospho-1-rac-glycerol
CW EPR  continuous-wave electron paramagnetic resonance
SR EPR  saturation recovery electron paramagnetic resonance
DEER  double electron–electron resonance
CD  circular dichroism
PS  power saturation
DHPC  1,2-dihexanoyl-sn-glycero-3-phosphocholine

REFERENCES


Figure 1.
Domain structure of the M2 protein monomer studied here. In vivo, M2 functions as a homotetramer. (A) Each M2 monomer consists of an N-terminal ectodomain, a transmembrane domain, and a C-terminal cytoplasmic domain. The sequence of the 16 residues comprising the amphipathic helix (sites 47–62) is shown. Residues in red are mutated to alanine in M2-5Ala (F47A, F48A, I51A, Y52A, and F55A). Asterisks (*) represent sites (L43, H57, L59) selected for spin labeling studies. (B) Helical wheel diagrams and hydrophobic moments (μH) of the M2-WT and (C) M2-5Ala amphipathic helices created by HeliQuest. Residues charged at pH 7.8 are shown in black, and uncharged residues are shown in gray.
Figure 2.
EPR line shapes indicate TM site 43 is immobilized in M2-5Ala relative to M2-WT, while AH sites 57 and 59 are more mobile in M2-5Ala relative to M2-WT. X-band CW EPR spectra of M2-WT and M2-5Ala in 4:1 POPC/POPG bilayers. Each sample contains, on average, one spin-labeled monomer per tetramer. Inset shows the low-field immobile (i) and mobile (m) peak components.
Figure 3.
M2-WT and M2-5Ala exhibit two components in conformational exchange. (A) Representative saturation-recovery data and fit for site 59 of M2-WT and (B) M2-5Ala in 4:1 POPC/POPG bilayers under 40% air. Residuals for mono- and biexponential fits are shown below the curve. Inset shows the dependence of $W_e$ on the concentration of $O_2$ for the fast, mobile component (black) and slow, immobile component (red). (C) Oxygen accessibility determined from the slopes of the linear best fit lines of data shown in the insets.
Figure 4.
DEER reveals different distance distributions of spin-labeled site 59 of (A) M2-WT and (B) M2-5Ala reconstituted into 4:1 POPC/POPG bilayers without (−) or with (+) cholesterol. Background-subtracted dipolar evolution data are shown as black dots with the corresponding fits as solid lines. Overlaid distance distributions were normalized to the area under the curve. Distances below 20 Å fall below the reliable detection limit of DEER.
Figure 5.
M2-WT and M2-5Ala CW line shapes exhibit differing sensitivity to cholesterol. CW EPR spectra of (A) M2-WT and (B) M2-5Ala in bilayers without (−) or with (+) cholesterol. The low-field immobile (i) and mobile (m) component peaks are indicated.
Figure 6.
A cartoon model consistent with experimental data collected from the M2-WT and M2-5Ala AH and TM domains. For clarity only two of the four subunits of the M2 homotetramer are shown. Furthermore, only the most populated conformation for each set of conditions is shown. The addition of cholesterol to the membrane shifts the M2-WT AH population toward a more compact conformation. Coupling between the AH and TM causes M2-WT to adopt a wedge-like shape that promotes curvature, shaded in blue. The M2-5Ala AH do not interact closely with the membrane surface, and consequently the conformation of M2-5Ala is insensitive to bilayer properties.