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Cholesterol dependent conformational exchange of the C-terminal domain of the influenza A M2 protein

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Abstract

The C-terminal amphipathic helix of the influenza A M2 protein plays a critical cholesterol dependent role in viral budding. To provide atomic-level detail on the impact cholesterol has on the conformation of M2 protein, we spin-labeled sites right before and within the C-terminal amphipathic helix of the M2 protein. We studied the spin-labeled M2 proteins in membranes both with and without cholesterol. We used a multipronged site-directed spin-label electron paramagnetic resonance (SDSL-EPR) approach and collected data on line shapes, relaxation rates, accessibility of sites to the membrane, and distances between symmetry related sites within the tetrameric protein. We demonstrate that the C-terminal amphipathic helix of M2 populates at least two conformations in POPC/POPG 4:1 bilayers. Furthermore, we show that the conformational state that becomes more populated in the presence of cholesterol is less dynamic, less membrane buried, and more tightly packed than the other state. Cholesterol dependent changes in M2 could be attributed to the changes cholesterol induces in bilayer properties and/or direct binding of cholesterol to the protein. We propose a model consistent with all our experimental data that suggests that the predominant conformation we observe in the presence of cholesterol is relevant for the understanding of viral budding.

Influenza A viruses cause recurrent seasonal epidemics and global pandemics.^{1,2} The threat of future pandemics coupled with growing resistance to current antivirals makes the development of new antiviral influenza drugs a national healthcare priority.³ A series of recently published papers provides new insight into how influenza virus buds at the plasma membrane of infected cells.^{4–6} An atomic-level understanding of the budding process could lead to strategies to inhibit the replication of viruses and new tactics for inhibiting viral infectivity.⁷ One of the proteins found on the viral coat of influenza, the M2 protein, has been shown to be critical to viral budding^{5,6,8} and is the focus of this paper.

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SUPPORTING INFORMATION AVAILABLE. Figures S1–S6. This material may be accessed free of charge online at <http://pubs.acs.org>.

The M2 protein is a multifunctional membrane-bound protein that acts at several stages of the virus lifecycle.⁹ Each 97-amino-acid monomer within the homotetrameric protein consists of a N-terminal ectodomain, a transmembrane domain and a C-terminal cytoplasmic domain.⁹ The most extensively studied function of the M2 protein is its proton channel activity, mediated by the transmembrane domain, that is crucial for the uncoating of virions when viruses enter cells. More recently, the C-terminal domain of M2 has been shown to play a critical role in viral budding, which is an essential step in the replication cycle of influenza.^{8,10}

Deletion mutants provided initial evidence that the C-terminal region of the M2 protein is critical for infectivity and virus morphology.¹¹ Further deletion studies and alanine-scanning experiments by several groups identified residues ~46–61 as being critical to viral budding.^{5,6,12} An *in vitro* assay has been used to probe the 46–61 region and demonstrated that both the full-length protein as well as just the 46–61 region are capable of inducing budding in giant unilamellar vesicles.⁸

Several groups, including our own, have demonstrated that the 46–60 residue region forms an amphipathic membrane-surface helix.^{13,14} Amphipathic membrane-surface helices have been shown to promote curvature in several systems,^{15–17} presumably due to their ability to cause lipid-packing defects that induce curvature. As a homotetramer, the M2 protein contains four amphipathic helices, but it is not understood how the four helices are arranged with respect to each other and how they work in concert to promote viral budding.

Site-directed spin label EPR spectroscopy (SDSL-EPR) has proven to be a powerful method for studying membrane proteins reconstituted into lipid bilayers¹⁸ and can provide conformational insight not accessible from other biophysical techniques. Structural parameters measured in SDSL-EPR experiments include distances from ~12–60 Å, the topology of a membrane-protein with respect to a lipid bilayer and mobility information on multiple timescales.¹⁹ Our group has previously published several SDSL-EPR studies on M2.^{13,20–23} As part of this work, we demonstrated that the conformation and dynamics of the C-terminal amphipathic helix of M2 changes in response to pH¹³ and the addition of adamantane drugs.²⁰

Influenza budding studies have shown that viral budding is cholesterol dependent.⁴ It has been hypothesized that cholesterol is necessary for guiding M2 to the edge of an infected cell's cholesterol-rich budzone and for M2's ability to promote curvature for viral budding.²⁴ Despite extensive study and speculation, atomic-level details of the roles cholesterol might play in viral budding are not understood.^{24,25} In this paper, we extend our previous SDSL-EPR work on the C-terminal amphipathic helix of M2 and present data on the impact cholesterol has on the conformation and dynamics of this region with the goal of providing insight into the cholesterol dependence of M2's role in viral budding.

For this paper, we studied spin-labeled sites right before and within the C-terminal amphipathic helix. Utilizing a multipronged SDSL-EPR approach, we demonstrate that the C-terminal amphipathic helix can exist in at least two conformational states that differ in the depth of the helices in the membrane and how the four helices within the homotetrameric

protein are arranged with respect to each other. The populations of these two conformational states shift upon addition of cholesterol to the membrane lipid bilayer.

MATERIALS AND METHODS

Synthesis, spin labeling and membrane reconstitution of M2 peptides

Peptide synthesis and purification was performed as described previously¹³ and the sequences of M2TMC 23–60 peptides are provided in Figure 1B. This 38-residue peptide construct has been used in a large number of structural studies on M2,^{13,20,26–27} and has been shown to generate the membrane curvature necessary for viral budding.²⁸ Furthermore the C-terminal helix in the M2TMC 23–60 peptide has been shown to have the same conformation as the same region in the full-length M2 protein.²¹ Peptides were nitroxide spin-labeled using cysteine-specific 1-oxy-2,2,5,5-tetramethyl-3-pyrroline-3-methyl methanethiosulfonate (MTSL) spin labels.¹³

For line shape analysis and accessibility studies, M2 peptides were reconstituted into lipid bilayers consisting of 1-palmitoyl-2-oleyl-*sn*-glycero-phosphocholine (POPC) and 1-palmitoyl-2-oleyl-*sn*-glycero-3-[phospho-*rac*-(1-glycerol)] (POPG), POPC/POPG 4:1 molar ratio. The peptide:lipid molar ratio was 1:200. The reconstitution protocol was described in detail previously.¹³ We chose POPC/POPG 4:1 bilayers because an *in vitro* budding assay has demonstrated that the M2 protein can induce curvature and initiate budding in vesicles made of POPC/POPG 4:1.⁸ Furthermore, a number of previously published biophysical studies on M2 have utilized POPC/POPG 4:1 bilayers and provide valuable context for this study.^{13,20–23} The sample buffer was 50 mM Tris pH 7.8, 100 mM KCl, 1 mM EDTA.

For double electron-electron resonance (DEER) EPR samples, M2 peptides were reconstituted into lipid bilayers of POPC/POPG 4:1 molar ratio at a peptide:lipid 1:300 molar ratio, micelles consisting of 1,2-dihexanoyl-*sn*-glycero-3-phosphocholine (DHPC) at peptide:detergent 1:500 molar or bicelles consisting of POPC/POPG/DHPC at a peptide:lipid 1:500 molar ratio. M2 peptides were also reconstituted into Lipodisq nanoparticles comprised of POPC/POPG lipid bilayers as previously described.²⁹ The DEER sample buffer was 50 mM Tris pH 7.8, 100 mM KCl, 1 mM EDTA in 30% (v/v) glycerol.

For saturation-recovery EPR (SR-EPR) samples, M2 peptides were reconstituted into POPC/POPG 4:1 molar ratio lipid bilayers at a peptide:lipid 1:500 molar ratio. The samples were spin-diluted by adding unlabeled M2TMC C50S at an unlabeled:labeled 4:1 molar ratio.

EPR Spectroscopy and Data Analysis

Continuous wave (CW) EPR spectra were recorded at room temperature on an X-band Bruker EMX spectrometer equipped with an ER4123D resonator. Samples used for analysis of spectral line shapes were placed in glass capillary tubes and EPR spectra were acquired using 2 mW incident microwave power, 1 G field modulation amplitude at 100 kHz, and 150 G sweep width. For comparison of line shapes, each spectrum was double integrated and normalized to the same number of spins.

EPR lineshapes were simulated using Multicomponent, a LabView program written by Christian Altenbach. The program fits slow-motion EPR data using a microscopic order macroscopic disorder (MOMD) model.³⁰ The data were well fit using two components with distinct rotational properties. Initial values for magnetic tensors were taken from previously published data of MTSL dynamics on a solvent-exposed membrane protein alpha-helix site.³¹

For power saturation measurements, samples were collected in gas-permeable TPX capillary tubes. Power saturation data were obtained under three sets of conditions: equilibrated with nitrogen gas, equilibrated with ambient air, and equilibrated with nitrogen in the presence of 20 mM nickel(II) ethylenediaminediacetate (NiEDDA). For experiments with low accessibility to fast relaxing paramagnetic reagents, EPR spectra were measured at 8 power levels. For experiments with high accessibility to paramagnetic reagents, 16 power levels were studied as the saturation effect was sometimes significantly mitigated and thus more data points were required to achieve a good fit. Data were analyzed and $P_{1/2}$ parameters reflecting side chain accessibility to paramagnetic reagents were determined as described previously.¹³

Saturation recovery EPR experiments (SR-EPR) were carried out at the National Biomedical EPR Center on a home-built instrument operating at X-band (9.4 GHz) equipped with a loop-gap resonator.³² Briefly, samples (5 μ l) contained in a gas-permeable TPX capillary were placed under a continuous flow of N₂ mixed with air to give the desired final oxygen concentration. A 1 μ sec saturating microwave pulse was applied at the absorption maximum of the central ($M_I = 0$) line, followed by signal acquisition with sampling intervals ranging from 2 – 20 nsec per point for 2048 points, and an overall repetition rate of 40 – 100 kHz. Total accumulation times varied from 5 – 60 minutes. The resulting SR signals were fitted with single and double exponential decay functions using Origin software (OriginLab, Northampton, MA). Bimolecular collision rates with oxygen, $W_x(O_2)$, were calculated from spin-lattice relaxation rates (W_e) according to the relationship^{19,33}, $W_x(O_2) = W_e(O_2) - W_e(N_2)$ where $W_e = 1/2T_1$.

Double electron-electron resonance (DEER) EPR 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.²⁹ DEER data were collected using the standard four-pulse sequence³⁴ $[(\pi/2)_{\nu_1} - \tau_1 - (\pi)_{\nu_1} - t - (\pi)_{\nu_2} - (\tau_1 + \tau_2 - t) - (\pi)_{\nu_1} - \tau_2 - \text{echo}]$ 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 for overnight data acquisition time (12 hours). The DEER data were analyzed using DEER Analysis 2015,³⁵ and the distance distributions were extracted by Tikhonov Regularization.³⁶ The regularization parameter was optimized by L-curve calculation using a modified algorithm provided by Dr. Clark Hyde at the University of Chicago.³⁷

RESULTS

M2 conformation and dynamics is sensitive to choice of membrane mimic

Structural studies of the influenza A M2 protein have been published using membrane mimics ranging from detergent micelles to complex bilayers.^{26,38–40} In an effort to reconcile the differences seen in published structures, we collected EPR spectra of M2 in a range of different membrane mimics.

EPR experiments are well suited to study the impact of membrane environment since the time scale of the spectroscopic measurement allows a wide range of membrane mimics to be employed. In Figure 2 we show EPR spectra of a spin-labeled M2 peptide in a range of different environments used in previously published work. DHPC micelles were used for solution nuclear magnetic resonance spectroscopy (NMR) studies,²⁶ POPC/POPG bilayers were employed for site-directed spin-label EPR (SDSL-EPR) experiments,^{13,20–23} and viral mimetic membranes (SM:DPPC:DPPE:Chol:PEG-PE 23:21:21:30:1) were used in solid state NMR (SSNMR) experiments.⁴¹ POPC/POPG bilayers were also used in an *in vitro* budding assay, which demonstrated that adding cholesterol to POPC/POPG bilayers modulates the ability of the M2 protein to initiate budding.⁸

A striking observation from the stack plot of spectra in Figure 2 is that the CW EPR spectra appear to be a superposition of two different spectral components, whose relative weights shift as the membrane mimetic moves from detergent micelles toward bilayers with increasing amounts of cholesterol. The different spectral components are highlighted in gray. The darker gray box indicates a broader, more immobilized component (I) that increases in intensity from top to bottom in Figure 2. The more mobile component (M) indicated by a lighter gray box decreases in intensity from top to bottom.

Considering the impact that cholesterol has on the ability of M2 to initiate viral budding, we were particularly intrigued by the differences seen in the EPR line shapes in POPC/POPG bilayers upon the addition of cholesterol (Figure 2). We thus designed experiments to more fully characterize cholesterol-induced change in M2 conformation and dynamics.

Line shapes for sites within the C-terminal domain are a weighted average of two different components

In Figure 2, one spin-labeled site (I42C) is used to analyze the impact of the environment on line shape. In Figure 3 we show EPR spectra for five different sites located before and within the C-terminal amphipathic helix. The goal of this experiment was to rule out the possibility that the two-component nature of the M2TMC-I42SL spectra shown in Figure 2 was simply a localized effect for site 42.

As demonstrated in Figure 3, the EPR spectra for all five sites exhibit a two-component nature. The two components present are labeled as mobile (M) and immobile (I). As the cholesterol content of the lipid bilayer changes, the relative intensities of these two components change. With increasing cholesterol concentration, the peak corresponding to the more immobile component increases in intensity as this conformation becomes more

populated. The observation that this trend is seen for all five sites is consistent with a concerted cholesterol-dependent change occurring in this region.

Shown in Figure 4A are EPR spectra for site 55 and fits using an EPR simulation package based on the MOMD approach³⁰ (see Experimental Procedures for details). Simulations of these spectra required two motional components to obtain a reasonable fit, but were not improved by including a third motional component. The mobile and immobile components have correlation times of 2.43 and 8.55 ns, respectively. In the cholesterol-free, 20% cholesterol, and 30% cholesterol environments, the relative population of the immobile component is 45%, 57%, and 68%, respectively.

The work presented in this paper uses a M2 construct (M2TMC 23–60) that does not include the last 37 residues of the C-terminal domain. It is conceivable that the multicomponent nature of the EPR line shapes seen in Figures 2 and 3 could be attributed to the use of a truncation of the full-length protein. However, it has recently been demonstrated that the multicomponent EPR line shapes we observe for sites within the amphipathic helix of M2 are very similar to the line shapes observed for the same sites on the full-length M2 protein.²¹ The secondary structure and membrane depth of the amphipathic helix in M2TMC 23–60 is also strikingly similar to the same region in the full-length M2 protein.²¹ These findings suggest that the multicomponent nature we report here is an intrinsic property of the M2 protein.

Saturation recovery data consistent with conformational exchange of C-terminal amphipathic helix

The two-component EPR spectra shown in Figure 3 can arise from an equilibrium involving either different conformational substates of the M2 protein or different rotamers of the MTSL spin-label.¹⁹ It is important to be able to distinguish between the origins of multicomponent line shapes, and the technique of saturation recovery EPR (SR-EPR) is well suited for this measurement.¹⁹ In SR-EPR, an intense saturation pulse is delivered at a frequency corresponding to the central line of the nitroxide. The return of spectral intensity is monitored with a weak CW observing microwave field at the same frequency and occurs with a characteristic time constant, T_1 . For a system with spin-label rotameric exchange in the absence of conformational exchange, a single exponential recovery is expected. For a system undergoing conformational exchange, the relaxation to equilibrium is generally bi-exponential with relaxation rate constants that are a function of the intrinsic T_1 values of the two components and the exchange rate. Rotamer exchange for a MTSL spin label lies in the ~0.1-1 μ s time range, while conformational exchange is at least an order of magnitude slower.¹⁹

SR-EPR data were collected for sites 42, 51 and 55. SR-EPR data for site 55 is presented in Figure 5. As shown in Figure 5, the saturation recovery curve for site 55 is best fit by a double exponential, which corresponds to two unique spin-lattice relaxation rates, consistent with conformational exchange. SR-EPR data for sites 42 and 51 (Figure S1) are also characterized by double-exponential recoveries. The observation of double exponential recoveries for all three sites (42, 51 and 55) is consistent with conformational exchange of the entire C-terminal amphipathic helix and not with localized effects at a single residue.

Spin-labels in two different conformational states can have different accessibilities to paramagnetic relaxation agents (such as oxygen) that will modulate the intrinsic T_1 values of the two states. O_2 is a hydrophobic molecule and preferentially partitions into lipid bilayers, with the highest concentration of O_2 in the middle of the bilayer. Figure 5A (inset) shows the dependence of the spin-lattice relaxation rates on the concentration of oxygen. The data shown indicate the slow component (blue line) is less dependent on changes in oxygen concentration than the fast component (red line), which is consistent with the slow component being less accessible to oxygen and thus not as deep in the hydrophobic membrane. The finding that the less mobile component is less deeply buried in the membrane may at first seem surprising. For many membrane proteins, regions that are more deeply buried in the membrane are immobilized relative to regions that are on the surface of the membrane.¹⁸ However, as we will discuss below, both protein-protein interactions and cholesterol binding are also means for immobilization of the M2 protein.

Accessibility studies show that the conformational state of the C-terminal helix that becomes more populated upon the addition of cholesterol is not as deep in the membrane as the other conformation

The saturation recovery data described above indicate that the two conformations of the C-terminal helix of M2 have different accessibilities to membrane-soluble paramagnetic oxygen. To further explore this finding we used another complementary method that also probes membrane accessibility (power saturation) and collected data for four sites within the C-terminal amphipathic helix (46, 48, 51 and 55).

Power saturation EPR experiments have been widely used in SDSL-EPR to determine a residue's accessibility to paramagnetic reagents.⁴² In our experiments, accessibility to two paramagnetic reagents, NiEDDA and O_2 , were measured by probing changes in the relaxation efficiency of the spin label in the presence of these compounds. As mentioned above, O_2 is a hydrophobic molecule that preferentially partitions into lipid bilayers. Conversely, NiEDDA is a hydrophilic compound that remains in the aqueous phase. Accessibility to NiEDDA and O_2 provide complementary data and as such, the trends seen should be self-consistent.

All the sites within the amphipathic helix (46, 48, 51, and 55) have an increased accessibility to NiEDDA (Figure 6A) and a decreased accessibility to oxygen (Figure 6B) upon addition of cholesterol. The data are consistent with the amphipathic helix becoming less buried in the membrane upon addition of cholesterol. Although we can say with confidence that the helix is on average deeper in the membrane in the absence of cholesterol, there are significant challenges to assigning specific distances to the changes in membrane depths. The power saturation studies report on data from both conformational states present. Reporting quantitative distances near the membrane surface for our membrane-bound M2 protein is complicated by a lack of distance calibration points above the glycerol backbone.⁴³ Therefore we do not report a quantitative change in depth due to these limitations. We can say, however, that the changes in oxygen accessibility we observe here are substantially larger than what was reported for changes in oxygen accessibility for the

M2 C-terminal amphipathic helix resulting from pH-triggered opening of the M2 proton channel.¹³

Studies on other proteins have demonstrated that cholesterol addition to membranes decreases the ability of membrane associating helices to penetrate into the lipid head group region.⁴⁴⁻⁴⁶ As explained in more detail in the Discussion, we postulate that the observed movement of the amphipathic helices away from the membrane surface could result from the condensing effect cholesterol induces on liquid-disordered phases which decreases the ability of membrane associating helices to penetrate into the lipid head group region.⁴⁷

Distance dependent dipolar couplings demonstrate the C-terminal amphipathic helix is more closely packed in the presence of cholesterol

The data presented in Figure 6A,B clearly demonstrate that the C-terminal amphipathic helix of M2 changes in membrane depth upon the addition of cholesterol. We next explored whether there was a change in the arrangement of the C-terminal amphipathic helices within the homotetramer upon the addition of cholesterol. These experiments would also be a test of the hypothesis that increased intersubunit interactions between the amphipathic helices might account for our observed immobilization of M2 in the presence of cholesterol.

Broadening of CW-X-band line shapes are observed when two spin labels are less than ~20 Å apart.⁴² Dipolar coupling leads to peak broadening and a decrease in the central peak intensity. It is possible to quantify dipolar coupling by comparing central peak intensities between a fully-labeled sample, which has four spin labeled M2 monomers per tetramer, and a dilute labeled sample, which has only one spin labeled M2 monomer per tetramer. An Ω value greater than 1 indicates that there is significant broadening due to dipolar coupling of sites separated by less than ~20 Å. Since CW EPR gives an averaged spectrum, Ω reflects the effect of dipolar coupling for both M2 C-terminal conformational states.

Ω values are shown in Figure 6C for M2 both with and without cholesterol. The addition of cholesterol increases the measured value of Ω for all four sites. Ω values result from dipolar coupling which is inversely proportional to inter-label distance. Thus the measured increases in Ω values for sites 46, 48, 51 and 55 indicate the addition of cholesterol leads to the closer approach of M2 subunits within the tetrameric bundle.

It is challenging to quantitatively interpret Ω values for this system. Our measured Ω values reflect dipolar coupling from both conformational states present, which have overlapping central peaks. It is non-trivial to extract reliable quantitative distances from dipolar broadened CW line shapes when more than one conformation is present.^{18,48}

DEER data reveal distances consistent with more than one conformation

Double Electron-Electron Resonance (DEER) EPR can resolve distance distributions within a population as well as measure distances longer than the ~20 Å limit measurable from broadening of CW line shapes.⁴⁹⁻⁵⁰ Furthermore, DEER has been successfully applied to membrane proteins,^{29,51} although resolving distances in homo-oligomeric proteins has proven challenging.³⁷ For example, a single labeled site in a homotetrameric protein with a single conformation would lead to two measurable distances, one resulting from subunits

that are next to each other within the tetrameric bundle (proximal) and another distance resulting from monomers that are diagonally related across the tetrameric bundle (Figure S3). With two distinct conformations of a homotetramer, we thus theoretically expect four unique distances (Figure S3).

DEER data for sites 51 and 55 are shown in Figure 7, with corresponding L-curves shown in Figure S4. A model-free Tikhonov Regularization fit was used to determine an approximate distance distribution.³⁵ The Tikhonov Regularization of site 51 ($\lambda=40$) reveals local maxima at approximately 24, 31, and 40 Å. The Tikhonov Regularization of site 55 ($\lambda=25$) reveals local maxima at approximately 19, 33, 37, and 47 Å. The distance distributions shown in Figure 7 include more than the two distances predicted for a single conformation of a homotetramer and are consistent with the presence of more than one conformation. However, our time domain data does not show particularly well-defined oscillations, which suggests we interpret the data with caution.

The reliability of our distance distributions is supported by analyses of both sites 51 and 55 in different hydrophobic environments. Distance distributions for each of these two sites reveal local maxima at similar distances regardless of the membrane mimic employed to solubilize the protein. (Figure S5). Although there are similarities between the unique DEER distances, there are significant differences in the intensities of the peaks. This indicates that the multiple distances observed in Figure 7 likely correspond to unique protein conformations that are not artifacts of data analysis or a particular hydrophobic environment, but exist in different membrane mimics with varying populations.

It is instructive to compare the distances shown in Figure 7 to published models for M2. Published models that used a M2 construct similar to the M2 peptide employed in this paper include a model based on solid-state NMR (ssNMR) in bilayers consisting of 1,2-dioleoyl-sn-glycero-3-phosphocholine (DOPC) and 1,2-dioleoyl-sn-glycero-3-phosphoethanolamine (DOPE), DOPC/DOPE 4:1 (PDB 2LOJ)⁴⁰ and a model based on solution-state NMR in DHPC micelles (PDB: 2RLF).²⁶ DEER derived distances for both I51 and F55 are similar to some of the distances measured from the ssNMR model collected in DOPC/DOPE lipid bilayers (Figure S6).⁴⁰ Distances were measured between $H_{C,\gamma}$ of corresponding sites from the published ssNMR model (PDB 2LOJ).

The DEER data for site 51 includes a distance at ~ 24 Å, which is similar to the diagonal distance between $H_{C,\gamma}$ (~ 24 Å) observed in the solid-state NMR model of M2 in bilayers.⁴⁰ The proximal distance for this conformation (~ 17 Å) would not be observed in the DEER data because it is below the ~ 20 Å detection limit of the method. The DEER distances for site 51 at ~ 31 and 40 Å likely correspond to the proximal and diagonal distances of a previously uncharacterized less packed conformational state. The ~ 31 and 40 Å DEER distances have a ratio close to 1:1.4, which is expected from the proximal:diagonal distances in a symmetric homotetramer.

The DEER data for site 55 includes a distance at ~ 19 Å, which is similar to the diagonal distance between $H_{C,\gamma}$ (~ 19 Å) observed in the ssNMR model of M2 in bilayers.⁴⁰ The proximal distance for site 55 for this conformation (~ 15 Å) would not be observed in the

DEER data because it is below the ~20 Å detection limit of the method. The site 55 DEER distances of ~33 Å and 47 Å have a ratio of 1:1.4. These two distances could correspond to the proximal and diagonal distances of a previously uncharacterized less packed conformational state. The fact that the existence of these pairs of distances for both sites 51 and 55 do not correspond to distances in previously published structures provides strong evidence that these longer distances are from one of two conformational states we have detected using line shape analysis and saturation recovery.

The proximal and diagonal distances measured from the M2 model in micelles,²⁶ are 12 and 16.5 Å for I51 and 4.7 and 7.3 Å for F55. All four of these distances are below the ~20 Å detection limit of DEER. However, a recent solution NMR study compared the model of M2 in DHPC micelles to M2 in DMPC/DHPC bicelles¹⁴ and unlike the micelle model, the bicelle structure has significant similarities with the published ssNMR structure in bilayers.

DISCUSSION

Data from line shape analyses, relaxation rates, accessibility parameters and the measurement of distance-dependent dipolar couplings provide rich complementary information of the conformations of the C-terminal amphipathic helix of M2 protein. We have been able to demonstrate that the C-terminal amphipathic helix populates at least two conformations in POPC/POPG 4:1 bilayers, a lipid environment in which M2 induced functional budding assays have been carried out.⁸ Furthermore, we have been able to show that the conformational state of the C-terminal amphipathic helix that becomes more populated in the presence of cholesterol is less dynamic, less membrane buried, and more tightly packed than the other state. As described below, the intriguing result that the conformation that is less membrane buried is also less mobile could be attributed to the changes cholesterol induces in bilayer properties and/or direct binding of cholesterol to the protein.

M2 protein as a “shape shifter”

The hypothesis that the M2 protein has multiple low-energy conformational states is not new.^{23,52–58} It has been proposed that the various structural models proposed to date provide snapshots of distinct conformational states sampled by the protein during the viral life cycle. The M2 protein has at least two functions, the proton channel activity crucial for uncoating of virions⁹ and the generation of curvature essential for viral budding.¹⁰ These two distinct functions could conceivably require two different conformations.

Several studies have specifically addressed the potential for the C-terminal amphipathic helices of M2 to adopt different conformations. An early paper postulated that the C-terminal region might adopt either a water-soluble helical extension of the transmembrane helix or that each amphipathic helix of the extension could individually make a tight, 90° turn, and then bind hydrophobically to the membrane.⁵⁷ SDSL-EPR data has demonstrated that the C-terminal amphipathic helix of M2 in POPC/POPG 4:1 lipid bilayers lies on the membrane surface, but changes in membrane depth and helix packing in response to pH,¹³ the addition of adamantane drugs,²⁰ and the lipid composition of the bilayer.²² Solution NMR work has shown that the conformation of the C-terminal amphipathic helix changes

depending on solubilizing conditions and that models based on data from DHPC micelles²⁶ differ from models based on data collected in DMPC/DHPC bicelles.¹⁴ ssNMR studies have also shown that the conformation of the M2 protein changes depending on the composition of the bilayer.⁵⁸

Characteristics of M2 that promote conformational exchange

The inherent flexibility that allows M2 to access multiple conformational states was predicted by the observation that a remarkably large fraction of M2 variants with mutations at the transmembrane helix-helix interfaces maintain the ability of the protein to tetramerize.⁵⁴ The ability of the M2 protein to tolerate mutations along the helix-helix interfaces is consistent with structural plasticity that may be required for functional requirements for conformational exchange.

Beyond M2, it has been observed that stabilizing interactions have gone untapped in many other transmembrane proteins.⁵⁹ The lack of stability optimization has been hypothesized to result from the fact that the surrounding lipid bilayer provides enough constraints on membrane proteins for sufficient stability for cell viability. Furthermore, flexibility of membrane proteins could facilitate exchange between different states critical to the function of many membrane proteins, including proteins that act as gated channels and receptors that transduce signals across membranes.⁶⁰ SDSL-EPR has been demonstrated to be particularly well-suited to detect conformational substates of membrane proteins.⁶¹

In addition to the inherent plasticity in the packing of the four helices in the transmembrane domain in M2, it is likely that the loop region between the transmembrane domain and C-terminal domain could act as a flexible hinge. Movement about this flexible hinge could allow for different packing arrangements and different membrane depths of the four C-terminal amphipathic helices.

Model for how cholesterol leads to the conformational shift seen in our EPR data

The impact of cholesterol on the M2 protein could be due to changes cholesterol induces in the lipids surrounding the protein, direct binding of cholesterol to the protein, or a combination of both effects. Cholesterol is a polycyclic amphipathic molecule.⁶² The apolar section of cholesterol buried in the membrane has an asymmetric structure with two distinct faces that could form numerous van der Waals interactions with either lipids or membrane associated proteins. The polar section of cholesterol is a single hydroxyl (OH), which could form a hydrogen bond with either a lipid or membrane-associated protein.

Cholesterol can have a range of impacts on lipid bilayer properties depending of the lipid composition of the membrane and the temperature.⁶³ The POPC:POPG 4:1 lipid bilayers used in this study are in the liquid disordered phase at room temperature.⁶⁴ Cholesterol exerts a condensing effect on liquid-disordered phases and increases the thickness of the bilayer.⁴⁷ It has been previously demonstrated that increasing the thickness of the bilayer decreases the tilt of helices within the transmembrane domain of M2 leading to a more tightly packed homotetramer.²³ A number of studies have also shown that the condensing effect of cholesterol on membranes decreases the ability of membrane associating helices to penetrate into the lipid head group region.⁴⁴⁻⁴⁶ This effect could explain our finding that the

C-terminal amphipathic helix of M2 moves away from the membrane surface upon the addition of cholesterol.

Cholesterol binding has been characterized for a range of different membrane proteins.^{65–67} The observation that M2 localizes to the edge the cholesterol-rich budzone on the surface of an infected cell has been pointed to as evidence that there is a cholesterol binding site on the M2 protein.²⁴ The M2 protein contains a possible cholesterol recognition/interaction amino acid consensus (CRAC) sequence,^{8,24} but the looseness of the CRAC definition ((L/V)-X_{1–5}-(Y)-X_{1–5}-(K/R)) has raised some skepticism about its predictive value.⁶⁶ However, in the case of M2, the observation that the removal of a mandatory tyrosine within the proposed CRAC domain of M2 disrupts the association of the M2 protein with cholesterol-containing membrane surfaces lends support to the existence of a cholesterol binding site.⁶⁸

The direct binding of cholesterol to M2 could explain why the addition of cholesterol shifts the conformational equilibrium towards a more immobilized state. The pairing of a transmembrane helix surface with a rigid cholesterol molecule could be expected to be entropically advantageous compared with association with more flexible lipids.⁶⁷ It has also been previously suggested for other membrane proteins that the apolar faces of cholesterol could interact with neighboring transmembrane domains and exert a condensing effect on the whole protein.⁶² This cholesterol-induced condensing effect could not only account for the observed immobilization, but also explain the observed closer approach of the C-terminal helices within the tetrameric bundle.

A postulated model that incorporates our experimental data into the two observed conformations of M2 is shown in Figure 8. As compared to Conformation I (left in Figure 8A,B), the C-terminal helices of Conformation II (right in Figure 8A,B), are less mobile, less deep in the membrane and closer together within the homotetramer. In the absence of cholesterol, both Conformation I and Conformation II are present, but the equilibrium is shifted towards Conformation I. In the presence of cholesterol, both Conformation I and Conformation II are present, but cholesterol preferentially stabilizes Conformation II and the equilibrium is shifted towards Conformation II. Therefore, we postulate that Conformation II is predominant in the highly curved region of the neck of viral budding where cholesterol is concentrated (Figure 8C). The fact that viral budding is cholesterol dependent supports our proposal that Conformation II is relevant for budding. Furthermore, observations from DEER data with membrane mimics containing different degrees of curvature (Figure S5) indicate samples with less curved bilayered membranes include substantially longer distances representative of Conformation I compared to samples with highly curved membrane mimics (micelles and bicelles).²⁷

SUMMARY AND CONCLUSIONS

The C-terminal amphipathic helix of the influenza A M2 protein plays a critical cholesterol dependent role in viral budding. In principle, a detailed understanding of M2-mediated viral budding could assist in the development of antiviral drugs targeted to influenza. With the goal of providing atomic-level details on the role cholesterol plays in budding, we spin-

labeled sites right before and within the C-terminal amphipathic helix of the M2 protein and studied the proteins in membranes both with and without cholesterol. We used a multipronged SDSL-EPR approach and collected data on line shapes, relaxation rates, accessibility of residues to the membrane, and distances between symmetry related sites within the tetrameric protein. Our results demonstrate the existence of two conformational states of the C-terminal amphipathic helix that differ in mobility, subunit arrangement and membrane depth. The condensing effect cholesterol exerts on lipid bilayers and/or direct binding of cholesterol to M2 could help explain the observed shift in the populations of conformational states upon the addition of cholesterol, and suggests that the predominant conformation we observe in the presence of cholesterol is relevant for the understanding of viral budding.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgments

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ABBREVIATIONS

| | |
|-----------------|---|
| SDSL-EPR | site-directed spin labeling electron paramagnetic resonance |
| CW-EPR | continuous wave electron paramagnetic resonance |
| SR-EPR | saturation-recovery EPR |
| DEER | double electron-electron resonance |
| MTSL | 1-oxyl-2,2,5,5-tetramethylpyrro- line-3- methylmethanethiosulfonate |
| POPC | 1-palmitoyl-2-oleyl- <i>sn</i> -glycero-phosphocholine |
| POPG | 1-palmitoyl-2-oleyl- <i>sn</i> -glycero-3-[phospho- <i>rac</i> -(1-glycerol)] |
| DHPC | 1,2-dihexanoyl- <i>sn</i> -glycero- 3-phosphocholine |
| MOMD | microscopic order macroscopic disorder |
| ssNMR | solid-state NMR |
| NiEDDA | nickel(II) ethylenediaminediacetate |

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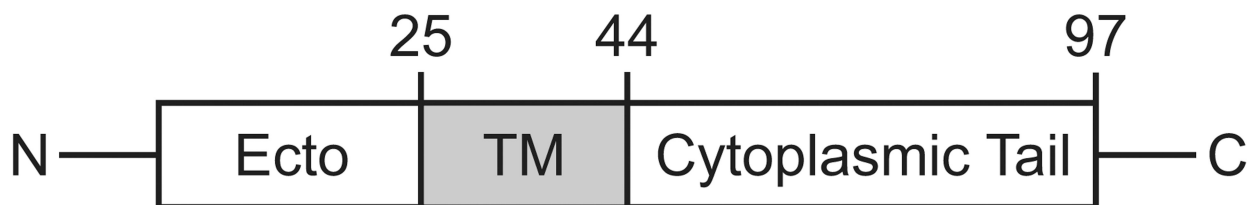
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A.



B.

| | 23 | 60 |
|------------|--|----|
| M2TMC-WT | SDPLVVAASIIGILHLILWILDRLFFKCIYRFFEHLGLK | |
| M2TMC-I42C | SDPLVVAASIIGILHLILWCLDRLFFKSIYRFFEHLGLK | |
| M2TMC-L46C | SDPLVVAASIIGILHLILWILDRCLFFKSIYRFFEHLGLK | |
| M2TMC-F48C | SDPLVVAASIIGILHLILWILDRLFCFSIYRFFEHLGLK | |
| M2TMC-I51C | SDPLVVAASIIGILHLILWILDRLFFKSCYRFFEHLGLK | |
| M2TMC-F55C | SDPLVVAASIIGILHLILWILDRLFFKSIYRFCEHLGLK | |
| M2TMC-C50S | SDPLVVAASIIGILHLILWILDRLFFKSIYRFFEHLGLK | |

Figure 1. Spin-labeled M2 peptides used in this study

A. Each monomer of the M2 protein is composed of a N-terminal ectodomain, a transmembrane domain (shaded in gray), and a C-terminal cytoplasmic domain. B. Wild-type sequence corresponds to the M2 protein from influenza strain A/Udorn/72 (H3N2). M2TMC (23–60) peptides were spin labeled at single cysteine sites (dark gray boxes). All M2TMC sequences other than the WT sequence have a C50S replacement in addition to the cysteine substitution necessary for spin labeling. The last cysteine-less sequence was used for spin dilution.

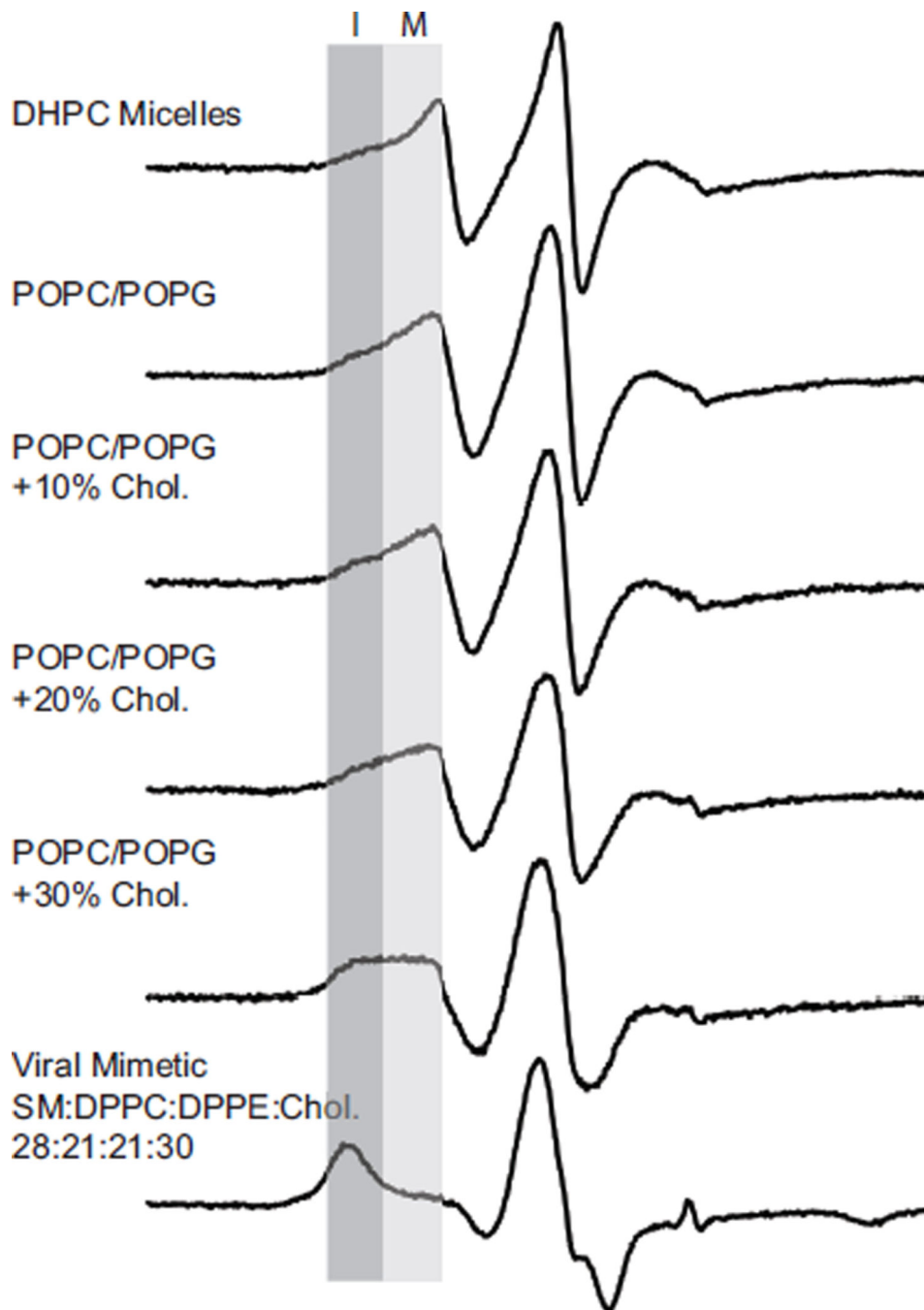


Figure 2. EPR line shape depends on choice of membrane mimic

Normalized X-band CW EPR spectra of M2TMC (23–60) spin labeled at I42C in DHPC micelles, POPC/POPG (4:1) bilayers with varying molar quantities of cholesterol, and a viral membrane mimic (SM:DPPC:DPPE:Chol:PEG-PE 23:21:21:30:1). Each sample was spin-diluted with unlabeled peptide (unlabeled:labeled 4:1) to eliminate broadening resulting from spin-coupling between monomers within the homotetrameric protein. The darker gray box indicates a broader, more immobilized component (I) that increases in intensity from

top to bottom. The more mobile component (M) indicated by a lighter gray box decreases in intensity from top to bottom.

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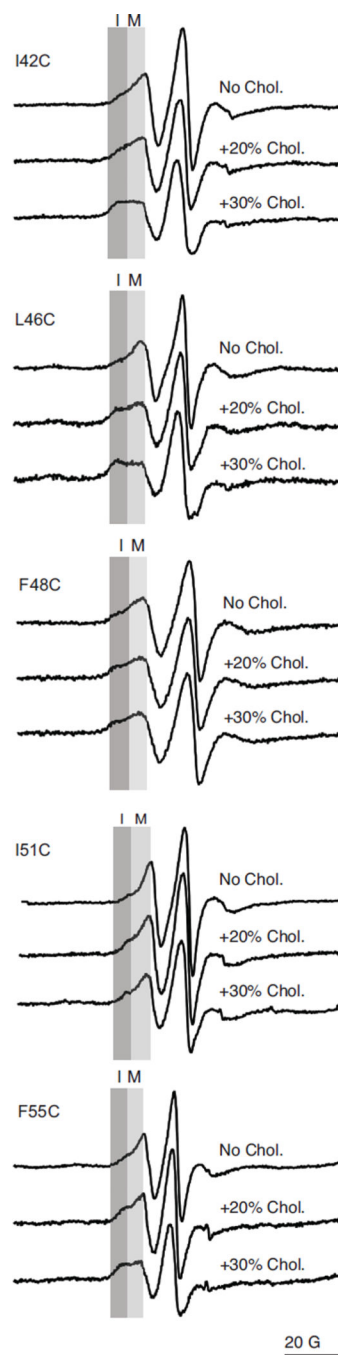


Figure 3. Normalized X-band CW EPR spectra of M2TMC (23–60) spin labeled at sites I42C, L46C, F48C, I51C, and F55C in POPC/POPG 4:1 bilayers in the absence and presence of cholesterol

Each sample was spin-diluted with unlabeled peptide (unlabeled:labeled 4:1) to eliminate broadening resulting from spin-coupling between monomers within the homo-tetrameric protein. The light grey box (M) and dark grey box (I) refer to the peak corresponding to the mobile and immobile conformations, respectively.

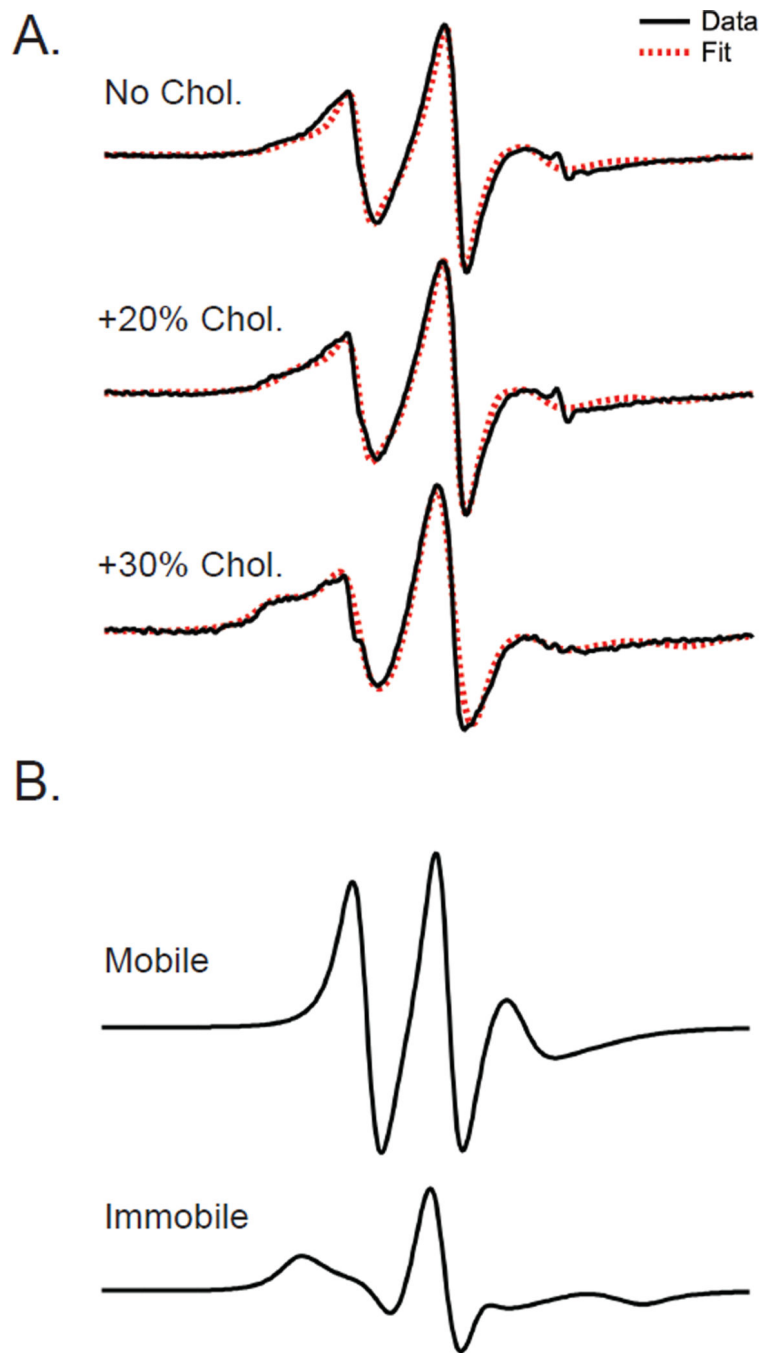


Figure 4. Simulation of line shapes for different amounts of cholesterol indicate a good fit to experimental data can be achieved using a weighted average of two different spectral components

A. EPR spectra for M2TMC spin-labeled at site F55C in POPC:POPG (4:1) bilayers with varying amounts of cholesterol (black lines) along with simulations (red dashed lines). B. Spectra corresponding to the mobile and immobile components used to fit lineshapes. The mobile and immobile components have correlation times of 2.43 and 8.55 ns, respectively. In the cholesterol-free, 20% cholesterol, and 30% cholesterol environments, the relative population of the immobile component is 45%, 57%, and 68%, respectively.

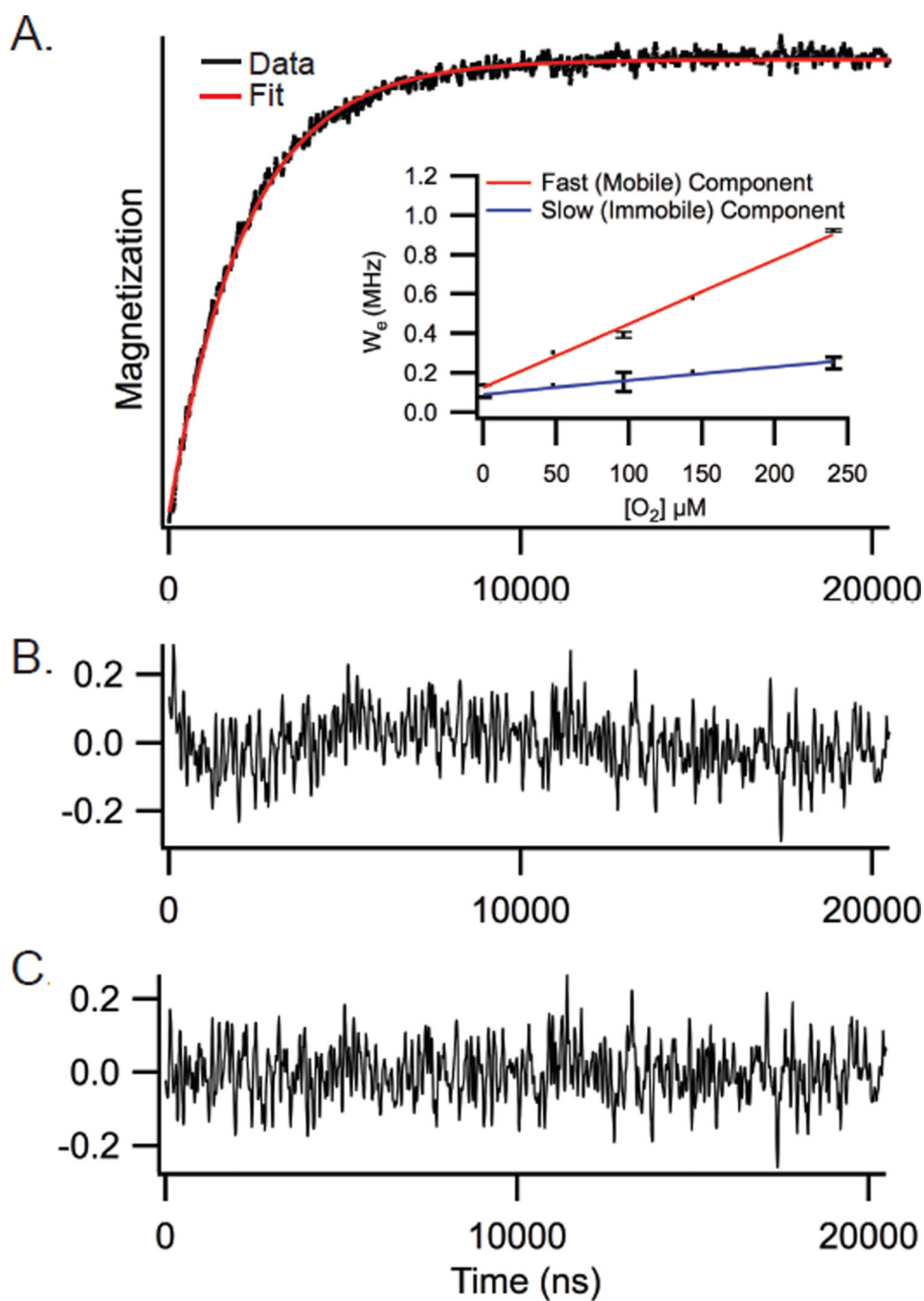


Figure 5. Saturation recovery data indicates two components in conformational exchange
 A. Representative saturation curve for M2TMC spin labeled at site F55C in POPC:POPG (4:1) under 20% air (black) with a double exponential fit (red). Inset shows the dependence of the spin-lattice relaxation rates on the concentration of oxygen for the fast, mobile (red) and slow, immobile (blue) saturation recovery components. B. Residual of a single exponential fit. C. Residual of a double exponential fit.

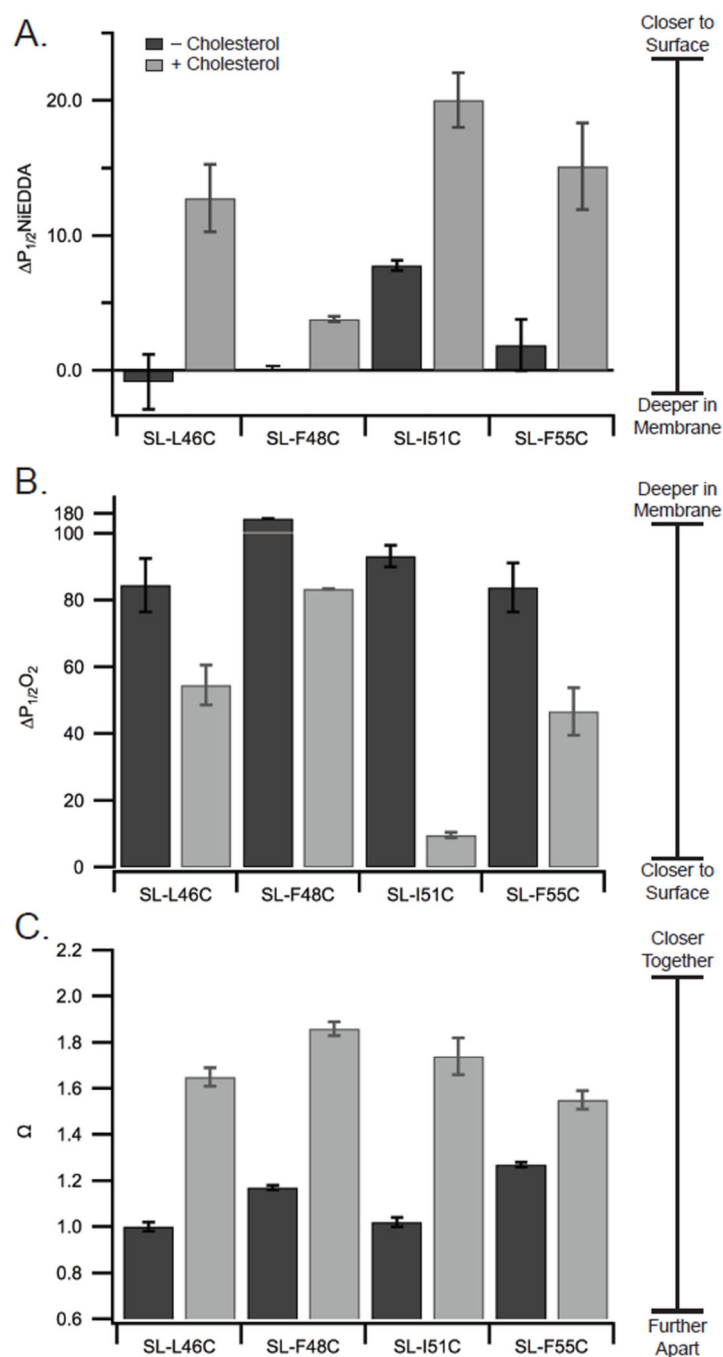


Figure 6. Addition of cholesterol changes the membrane topology of the C-terminal juxtamembrane region

Accessibility data to NiEDDA (A) and oxygen (B), and interaction parameter (Ω) data (C) for M2TMC labeled at the indicated sites in POPC:POPG (4:1) bilayers with 0% and 30% cholesterol. Ω is the ratio of the central line intensity for fully-labeled and spin-diluted samples. Spectra used to calculate Ω are shown in Figure S2.

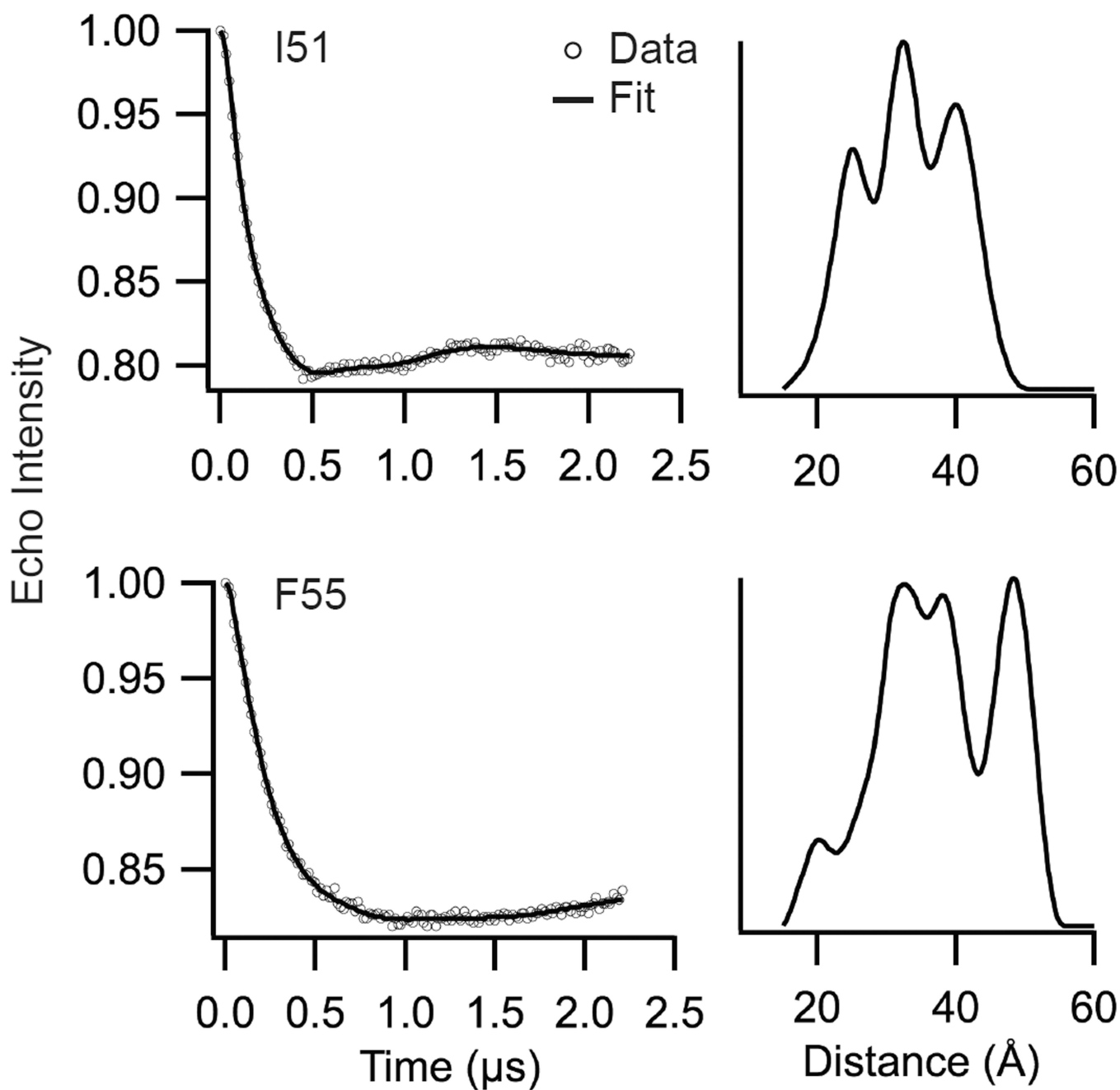


Figure 7. DEER reveals presence of multiple distances and protein conformations

Tikhonov Regularization and distance distributions of M2TMC-I51C ($\lambda=40$, top) and M2TMC-F55C ($\lambda=25$, bottom). Background-subtracted DEER data are shown as open circles and corresponding fits as black lines. L-curves are shown in Figure S4. M2 peptides were incorporated into 1,2-dihexanoyl-*sn*-glycero-3-phosphocholine (DHPC) at a peptide:DHPC molar ratio of 1:500.

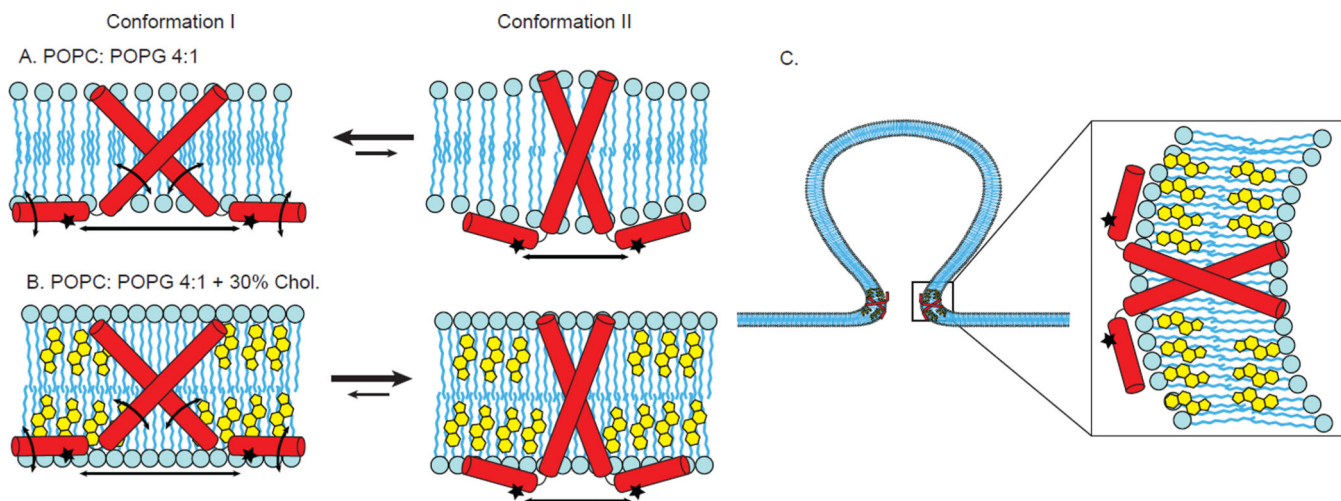


Figure 8. Cartoon model that is consistent with experimental data collected on the C-terminal amphipathic helices of M2
 Conformation I (left in Figure 8A,B), and Conformation II (right in Figure 8A,B). As compared to Conformation I, the C-terminal helices of Conformation II are less mobile, less deep in the membrane and closer together within the homotetramer. A. In the absence of cholesterol, both Conformation I and Conformation II are present, but the equilibrium is shifted towards Conformation I. B. In the presence of cholesterol, both Conformation I and Conformation II are present, but cholesterol preferentially stabilizes Conformation II and the equilibrium is shifted towards Conformation II. C. We postulate that in the highly curved region of the neck of viral budding where cholesterol is concentrated, Conformation II is predominant. For clarity, only two of the four subunits of a M2 homotetramer are shown in the cartoon models.