Structure and drug binding of the SARS-CoV-2 envelope protein transmembrane domain in lipid bilayers

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An essential protein of the SARS-CoV-2 virus, the envelope protein E, forms a homopentameric cation channel that is important for virus pathogenicity. Here we report a 2.1-Å structure and the drug-binding site of E’s transmembrane domain (ETM), determined using solid-state NMR spectroscopy. In lipid bilayers that mimic the endoplasmic reticulum–Golgi intermediate compartment (ERGIC) membrane, ETM forms a five-helix bundle surrounding a narrow pore. The protein deviates from the ideal α-helical geometry due to three phenylalanine residues, which stack within each helix and between helices. Together with valine and leucine interdigitation, these cause a dehydrated pore compared with the viroporins of influenza viruses and HIV. Hexamethylene amidolide binds the polar amino-terminal lumen, whereas acidic pH affects the carboxy-terminal conformation. Thus, the N- and C-terminal halves of this bipartite channel may interact with other viral and host proteins semi-independently. The structure sets the stage for designing E inhibitors as antiviral drugs.

Despite its importance to SARS-CoV-2 pathogenesis, E’s high-resolution structure, particularly for the ion-conducting transmembrane (TM) domain (residues 8–38) (Fig. 1b)19, has been elusive. Sedimentation equilibrium and gel-electrophoresis data for the homologous SARS-CoV-1 E indicate that the TM domain assembles into a homopentamer in detergents such as sodium dodecyl sulfate (SDS) and perfluorooctanoic acid13,14. Although early X-ray scattering data have suggested a helical hairpin model for E19, subsequent solution NMR studies of E bound to several detergent micelles, including dodecylphosphocholine (DPC)19, SDS19 and lyso-myristoylphosphatidylglycerol (LMPG)19, consistently indicate a single-span TM helix. However, the pore-facing residues and the pentameric assembly are not well-established. Fourier-transform infrared dichroic data suggest that the ETM helix orientation in lipid bilayers may be sensitive to the presence or absence of charged residues at the two termini of the TM domain, and by inference, the membrane surface charge13,14.

Here, we use solid-state NMR to determine the structure of the SARS-CoV-2 ETM structure in phospholipid bilayers, to avoid potential structural distortion caused by detergents. The structure sets the stage for the design of E inhibitors as antiviral drugs.

Results

Backbone conformation of ETM in lipid bilayers. We reconstituted ETM into an ERGIC-mimetic lipid bilayer containing phosphatidylcholine, phosphatidylethanolamine, phosphatidylinositol, phosphatidylserine and cholesterol. For comparison, we also incorpo-rated the protein into a dimyristoylphosphocholine (DMPC): dimyristoylphosphoglycerol (DMPG) model membrane, abbreviated as DMPX below. ETM was expressed in Escherichia coli using a hexahistidine (His6)-small ubiquitin-like modifier (SUMO) fusion tag and purified first by nickel affinity column chromatography and then by reverse-phase HPLC after cleavage of the solubility tag (Extended Data Fig. 1).

One-dimensional (1D) 13C and 15N NMR spectra of the protein in ERGIC and DMPX membranes show temperature-insensitive high intensities (Extended Data Fig. 2a,b), indicating that the protein is immobilized in lipid bilayers at ambient temperature. Two-dimensional (2D) 13C-13C and 15N-15N correlation spectra show well-resolved peaks (Fig. 1c,d), with 1H and 15N linewidths of 0.5 ppm and 0.9 ppm, indicating that the protein conformation is highly homogeneous. We assigned the chemical shifts using three-dimensional (3D) correlation NMR experiments (Extended Data Fig. 3a). These chemical shifts indicate that residues 14–34 form the α-helical core of the TM domain (Extended Data Fig. 3b,c and Supplementary Table 1). Comparison of spectra between the two membranes and at different temperatures (Extended Data Fig. 2d–f)
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Fig. 1 | Function, amino acid sequence and fingerprint NMR spectra of the SARS-CoV-2 E protein. a, E forms a cation-selective ion channel and mediates SARS-CoV-2 budding and release from the host cell’s ERGIC lumen. b, Domain architecture of E and sequence alignment of E’s transmembrane segment among several human-infecting coronaviruses. Highly conserved polar residues are shown in red. C-terminal domain; N-terminal domain. c, d, 2D 15N-13C correlation spectrum (c) and 2D 13C-13C correlation spectrum (d) of ERGIC-membrane-bound ETM. The spectra, measured at ambient temperature, show high sensitivity and resolution, indicating that ETM is structurally homogeneous in lipid bilayers.

indicate that the N-terminal segment (residues Glu8–Ile13) is dynamic at high temperature but is mostly α-helical, whereas the C-terminal segment (residues Thr35–Arg38) is more rigid but displays temperature-dependent conformations. Acidic pH perturbed the chemical shifts of C-terminal residues Leu34 to Arg38 (Extended Data Fig. 4), supporting the conclusion that the C terminus is conformationally plastic.

Oligomeric structure and hydration of ETM. The overall temperature insensitivity of the protein spectra suggests that ETM is oligomerized in lipid bilayers. To determine the oligomeric structure, we prepared two mixed labeled samples to measure intermolecular contacts. An equimolar mixture of 13C-labeled protein and 4-19F-Phe-labeled protein (Extended Data Fig. 1e) was used to double-resonance (REDOR) technique19 (Fig. 2a). ETM contains residues display 2 resolved 19F chemical shifts with a roughly 2:1 intensity ratio, indicating that one of the residues has a distinct side chain conformation. A 2D 13C-19F correlation spectrum (Fig. 2c) shows a cross-peak between the −118 ppm 19F signal and Ala22 Cα, indicating that this −118 ppm peak is due to either Phe20 or Phe23. The −113 ppm 19F peak shows strong cross-peaks with aromatic and numerous aliphatic 13C chemical shifts. Since Phe20 and Phe26 are too far away from each other to form intermolecular contacts, the −118 ppm 19F peak must be assigned to Phe20, while the −113 ppm peak must be assigned to Phe23 and Phe26. To constrain the interhelical packing at the two termini of the TM domain, we prepared a sample with mixed 13C and 15N labels and measured 2D NHHC correlation spectra to identify exclusively intermolecular 13N-15C correlations (Fig. 2d). These experiments together yielded 35 interhelical 13C-19F distance restraints and 52 interhelical 15N-13C correlations, which are crucial for determining the oligomERIC structure of ETM.

To further constrain the architecture of ETM self-assembly, we measured residue-specific water accessibilities using water-edited 2D 1H-13C correlation experiments (Fig. 2e and Extended Data Fig. 5d)20,21. Water 1H magnetization transfer is the highest to the N-terminal residues, is the least to the central residues Leu17 to Ala32 and is moderate to the C terminus (Fig. 2f). Thus, the hydration gradient of the protein is primarily along the bilayer normal. The preferential hydration of the N terminus is especially manifested by the high water-transferred intensity of Leu19 compared with that of Thr30, despite favorable chemical exchange to the Thr side chain22-24. For the dehydrated center of the TM domain,
Fig. 2 | Measurement of interhelical distances and water accessibility of membrane-bound ETM. a, Schematic of mixed [19F]ETM and [13C]ETM in a five-helix bundle. b, 2D 13Cα-F REDOR spectra of ERGIC-membrane-bound ETM. The control spectrum (S0, black) shows the signals of all residues, whereas the difference spectrum (ΔS, red) shows the signals of residues that are close to fluorinated Phe on a neighboring helix. c, 2D 13C-19F correlation spectrum allows assignment of the −118 ppm peak to Phe20 due to a cross-peak with Ala22, whereas the −113 ppm peak is assigned to Phe23 and Phe26 on the basis of correlations with Phe23 and Phe26 and with Val24 and Val25. A 1D 1H-19F cross-polarization (CP) spectrum is shown on the left. d, 2D NHHC correlation spectrum of mixed [13C]ETM and [15N]ETM, measured using 0.5 ms (red) and 1 ms (black) 1H mixing. All peaks arise from interhelical contacts. Selected assignments are given. e, Residue-specific water accessibilities of ERGIC-bound ETM obtained from the intensity ratios of water-edited spectra measured with 9 ms and 100 ms 1H mixing. Higher values (blue) indicate greater water accessibility. f, Water-edited intensities of ETM (black) and influenza BM2 (blue) obtained as the peak intensity ratios of the 9-ms and 100-ms spectra. Closed and open symbols indicate resolved and overlapped peaks, respectively. Error bars indicate the random uncertainty, which is propagated from the signal-to-noise ratios of the two spectra. ETM shows lower water-edited intensities than does BM2, indicating that the ETM pore is drier than the closed BM2 pore. g, Water- and lipid-edited 13C spectra of membrane-bound ETM. The Phe signals are high in the lipid-edited spectra but very low in the water-edited spectra, indicating that the three Phe residues are poorly hydrated and point to the lipids or the helix–helix interface.
Leu28 and Val25 show higher hydration than do their neighboring residues, suggesting that these two residues face the pore. A complementary lipid-edited experiment (Fig. 2g) showed much higher intensities for the Phe side chain carbons than their corresponding lipid-transferred intensities, indicating that the Phe residues are largely lipid-facing. The ERGIC-bound ETM shows twofold lower water accessibility than that of the closed state of the influenza BM2 at the same pH25 (Fig. 2f).

Structure calculation of ETM in ERGIC membranes. We calculated the structure of ETM using the above 56 (ϕ, ψ) torsion angles, 87 interhelical distance restraints (Supplementary Tables 2 and 3) and 196 intra-helical 13C-13C contacts obtained from 250-ms 2D 13C spin diffusion spectra (Extended Data Fig. 6)26. Initial calculation using directionally ambiguous interhelical contacts where the rotamers are not precisely defined, especially for side chains well away from the central three Phe residues (Extended Data Fig. 8b).

CSPs of HMA are larger than those of AMT and are consistent with the observed CSPs. The CSPs in lipid bilayers were measured under only fourfold drug excess, while in micelles, the smaller CSPs were measured under higher drug excesses of 10- to 31-fold16-18. This trend is consistent with the micelle data10,16, but the CSPs in lipid bilayers are much larger, with the N-terminal TGD11–14 triplet giving per-residue CSPs of 0.35–0.70 ppm. Moreover, the CSPs in lipid bilayers were measured under only fourfold drug excess, while in micelles, the smaller CSPs were measured under higher drug excesses of 10- to 31-fold16-18.

The higher sensitivity of ETM to HMA in lipid bilayers strongly suggests that the bilayer-bound protein conformation is more native. A docking pose based on these CSPs found that HMA intercalates shallowly into the N-terminal lumen with a distribution of orientations (Fig. 4d and Extended Data Fig. 9), suggesting a dynamic binding mode wherein HMA exchanges between multiple helices and inhibits cation conduction by steric occlusion of the pore. Within the ensemble of docked structures, more HMA molecules point the guanidinium into the pore and the hexamethylene ring towards the lipid headgroups than in the reverse orientation. AMT caused smaller CSPs than HMA (Fig. 4c and Extended Data Fig. 10a,b), but the binding site remains at the N terminus. Using the 3-19F label on adamantane, we measured protein-drug proximities using 13C-19F REDOR. The spectra showed modest dephasing for the N-terminal Asn15 and C-terminal Ile33 (Extended Data Fig. 10c–e), in qualitative agreement with the observed CSPs. The CSPs of HMA are larger than those of AMT and are consistent with the stronger affinity of HMA than AMT for SARS-CoV E, as well as with the micromolar half-maximal effective concentration (EC50) reported for HMA against other human coronavirus E proteins18.

ETM interactions with hexamethylene amiloride and amantadine. To investigate how ETM interacts with drugs, we measured the chemical shifts of the protein in the presence of HMA and [3-19F] amantadine. At a drug-protein molar ratio of 4:1, HMA caused significant chemical shift perturbations (CSPs) to N-terminal residues, including Thr9, Gly10, Thr11, Ile13 and Ser16, followed by more modest CSPs for the C-terminal Ala36 and Leu37 (Fig. 4a–c). This trend is consistent with the micelle data10,16, but the CSPs in lipid bilayers were measured under only fourfold drug excess, while in micelles, the smaller CSPs were measured under higher drug excesses of 10- to 31-fold16-18.

Table 1 | NMR and refinement statistics

| SARS-CoV-2 ETM (BMRB 30795, PDB 7K3G) | NMR and refinement statistics |
|----------------------------------------|-------------------------------|
| **NMR distance and dihedral constraints** |                                |
| Distance constraints                   |                               |
| Total NOE                              | 283×5                         |
| Inter-residue                          | 283×4                         |
| Sequential (|i|−|j| = 1)               | 125×5                         |
| Medium range (2 ≤ |i|−|j| ≤ 4)           | 71×5                          |
| Long range (|i|−|j| ≥ 5)             | 0×5                           |
| Intermolecular                         | 87×5                          |
| Hydrogen bonds                         | 11×5                          |
| Total dihedral-angle restraints        |                               |
| ϕ                                      | 28×5                          |
| ψ                                      | 28×5                          |
| **Structure statistics**               |                                |
| Violations (mean ± s.d.)               |                               |
| Distance constraints (Å)               | 0.17±0.06                     |
| Dihedral-angle constraints (°)         | 0.68±0.34                     |
| Max dihedral-angle violation (°)      | 5.54                          |
| Max distance-constraint violation (Å)  | 1.74                          |
| Deviations from idealized geometry     |                               |
| Bond lengths (Å)                       | 0.004±0.001                   |
| Bond angles (°)                        | 0.54±0.06                     |
| Improvers (°)                          | 0.42±0.04                     |
| Average pairwise r.m.s.d. (Å)          |                               |
| Heavy                                  | 2.06±0.58                     |
| Backbone                               | 1.67±0.65                     |

*Pairwise r.m.s.d. was calculated among the ten lowest-energy refined structures.

The lowest-energy structure ensemble, calculated using XPLOR-NIH (Supplementary Table 5 and Table 1), shows a long and tight 5-helix bundle with a vertical length of ~35 Å for residues Val14–Leu34. The structure resolution is higher for the middle of the TM domain, where 13C-19F REDOR distance restraints are available, and lower for the two termini, where fewer distance restraints are available (Fig. 3a and Extended Data Fig. 8a,b). The side chain rotamers are not precisely defined, especially for side chains well away from the central three Phe residues (Extended Data Fig. 8b).

The channel diameter, represented by backbone Cα-Cα distances between helices i and i+2 for pore-facing residues, varies from 11 Å to 14 Å. The helix is tilted by a small angle of 5–10° from the bilayer normal (Fig. 3b), but the orientation is not uniform along the length of the peptide, because the helix is non-ideal but exhibits a rotation angle change, or twist, between residues Phe20 and Phe23 (refs. 16-18). Consistent with the small tilt angle, the helical bundle does not display a strong handedness. The pore of the channel is occupied by predominantly hydrophobic residues, including Asn15, Leu18, Leu21, Val25, Leu28, Ala32 and Thr35 (Fig. 3b,c and Extended Data Fig. 8a,b), explaining the poor hydration of the protein. The N-terminal pore is constricted by Asn15, which forms interhelical side chain hydrogen bonds (Fig. 3g). The pore-facing positions of Asn15 and Val25 are consistent with single-channel conductance data showing that p.N15A and p.V25F abolish cation conductance27. The helix–helix interface is stabilized by aromatic stacking of Phe23 and Phe26 (Fig. 3e,f) and van der Waals packing among methyl-rich residues such as the Val29–Leu31–Ile33 triad (Fig. 3f). These extensive hydrophobic interactions give rise to a tighter helical bundle than do the viroporins influenza BM2 and HIV-1 Vpu (Extended Data Fig. 8d).

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Discussion

The current lipid-bilayer-based structural model of SARS-CoV-2 ETM has similarities with, but also considerable differences from, micelle-derived structural models (PDB 5X29)\(^1\). In LMPG micelles, the TM domain of a longer E construct (residues 8–65) also displays a kinked helix and a disordered N terminus, but the helical bundle is right-handed\(^1\), and the helices are more tilted and loosely packed (Extended Data Fig. 8c). In comparison, the bilayer-based ETM structural model does not have a strong handedness, consistent with the small helical tilt angle, and both reflect the measured interhelical distance restraints (Supplementary Tables 2 and 3).

The heavy-atom r.m.s. deviation (r.m.s.d.) for residues 14–34 between the 2 structural models is 6.1 Å, and the positions of various important residues differ. For example, in the LMPG-derived structural model, Phe26 is pore-facing and Thr30 is interhelical\(^1\), whereas in the bilayer-derived structure model, both residues point to lipids. The lipid-facing position of Thr30 in the current model is supported by single-channel conductance data showing that mutations of residues such as Thr30 and Thr11 to Ala do not affect the channel activity\(^1\). Another structural model of ETM determined in DPC micelles\(^1\) showed a left-handed and coiled helical bundle that differs qualitatively from the LMPG-bound model. These structural differences likely result from a combination of insufficient experimental restraints as well as an inherent conformational plasticity of the ETM. The LMPG-based structural model was obtained from ten unambiguous interhelical distances but no orientational restraints\(^1\), whereas the DPC-based structural model was built with orientational restraints but no unambiguous interhelical distance restraints\(^1\). For comparison, the current bilayer-derived ETM structure model was calculated from 87 interhelical distance constraints (Table 1).

Apart from experimental limitations, ETM’s oligomeric structure may be intrinsically sensitive to the membrane environment\(^2\) because the highly hydrophobic nature of the long central portion of the TM segment makes interhelical interactions non-specific. Indeed, SARS-CoV viruses with a p.V25F mutation develop escape mutants p.L27S, p.L19A, p.T30I and p.L37R in mice, implying that E’s channel activity is restored by these compensatory double mutations\(^12\). We speculate this could result from moderate changes of the helix rotation angle to give rise to alternate packing of the helical bundle. Future studies of E mutants are required to elucidate the structural basis for the loss and restoration of ion-channel activity.

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**Fig. 3 | Structure of SARS-CoV-2 envelope protein’s transmembrane domain in ERGIC-mimetic lipid bilayers.**

- **a**, Ensemble of ten lowest-energy structures.
- **b**, Side view of the most representative structure together with the HOLE-calculated pore water (gray). Pore-lining residues are shown as sticks.
- **c**, Simplified two-helix view with the pore-facing residues and their distributions in the lowest-energy ensemble.
- **d**, Pore radius of ETM obtained from the HOLE program.
- **e–h**, Additional snapshots from the most representative structure.
- **e**, Lipid-facing and helix-interface positions of the three Phe residues.
- **f**, Two clusters of methyl-interdigitating Leu, Ile and Val residues, stabilizing the helix-helix interface.
- **g**, Top views of the N-terminal Glu8, the pore-facing Asn15, and the three Phe residues.
- **h**, Surface plots of the pentamer, showing the N- and C-terminal vestibules where Asn15 and Lys28 are the first pore-facing residue.
How does the SARS-CoV-2 ETM structure compare with the structures of equivalent viroporins of influenza and HIV-1 viruses in lipid bilayers? The ETM helical bundle is compact and rigid, while AM2 and BM2’s TM domains, which have a higher percentage of polar residues such as His and Ser, form wider and more hydrated pores (Extended Data Fig. 8d)\(^2\). The HIV-1 Vpu TM domain has a high percentage of hydrophobic residues, similarly to SARS-CoV-2 E, but forms a shorter (~20˚ vertical length) pentameric helical bundle with more tilted helices (~20˚)\(^{30,31}\). The ETM helical bundle is more immobilized than M2 and Vpu helical bundles\(^{13}\), and does not undergo rigid-body fast uniaxial rotation at high temperatures (Extended Data Fig. 2d)\(^{19}\). This immobilization suggests that ETM may interact extensively with lipids\(^2\). Finally, the helix distortion at residues Phe20–Phe23 may cause the two halves of the protein to respond semi-independently to environmental factors such as pH, charge, membrane composition and other viral and host proteins.

Which structural features of this ETM helical bundle might be responsible for cation conduction? We hypothesize that the N terminus, which contains a (E/D/R)\(\times\)G(A/V)\(\times\)X(h/N/Q)\(\times\) motif (Fig. 1b), where h is a hydrophobic residue, contains the cation selectivity filter. In this conserved motif, the most exposed residue, Glu8, belongs to a dynamic N terminus whose residues (for example Thr9 and Gly10) manifest intensities only at high temperature (Extended Data Fig. 2d–f). The Glu8 side chain carboxyl is deprotonated at neutral pH and protonated at acidic pH, as manifested by \(^{13}C\) chemical shifts (Extended Data Fig. 2c). We speculate that the protonation equilibria of this loose ring of Glu8 pentamer, together with the anionic lipids in the ERGIC membrane, may regulate the ion selectivity of ETM at the channel entrance. Such a ring of negatively charged Glu residues has been observed as selectivity filters in the hexameric Ca\(^{2+}\)-selective Orai channels\(^{33}\) and designed K\(^{+}\) channels\(^{34}\). The third residue of the motif (G/A/V) is conserved among coronaviruses to be small and flexible (Fig. 1b), which might permit N-terminus motion and/or prevent occlusion of the channel lumen. The last residue of the motif is conserved to be either Asn or Gln, whose polar sidechains can coordinate ions and participate in interhelical hydrogen bonds to stabilize the channel\(^{27}\). At the C-terminal end of the TM segment, the conserved small residues Ala32 and Thr35 provide an open cavity for ions. In contrast to these small polar residues, the central portion of the TM domain contains four layers of hydrophobic residues, Leu18, Leu21, Val25 and Leu28, which narrow the pore radius to ~2 Å (Fig. 3d). This narrow pore can permit only a single file of water molecules, thus partially dehydrating any ions that move through the pore. Therefore, the structure determined here may represent the closed state of SARS-CoV-2 E, while the open state might have a larger and more hydrated pore. Narrow pores with multiple hydrophobic layers have also been observed in larger ion channels, including the tetrameric K\(^{+}\) channel TMEM175 (ref. \(^{35}\)) and the pentameric bestrophin channels\(^{36,37}\). Thus, it is possible to achieve charge stabilization and ion selectivity in such
a hydrophobic environment, although the detailed mechanisms remain to be understood.

The present membrane-bound ETM structure suggests that small-molecule drugs should have high-affinity binding to both the acidic Glu8 and the polar Asn15 in order to occlude the N-terminal entrance of the protein. The membrane topology of SARS-CoV-2 E is now recognized to be Nterm-αC-term, on the basis of antibody-detected selective permeabilization assays38 and glycosylation data39. This orientation may prime the protein to conduct Ca2+ out of the ERGIC lumen to activate the host inflammasome. Thus, small-molecule drugs should ideally be targeted and delivered to the Golgi and ERGIC of host cells to maximally inhibit SARS-CoV-2 E’s channel activity40.

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Methods

Cloning of recombinant ETM (8–38). The gene encoding full-length SARS-CoV-2 E protein (NCBI reference sequence YP_009724392.1, residues 1–75) was purchased from Genewiz. The gene encoding the TM domain (residues 8–38, EFGTTIYNSVLFFELAWNLTVLTAHLLL) was isolated using PCR and cloned into a Champion pET-SUMO plasmid (Invitrogen). The plasmid was transfected into E. coli BL21 (DE3) cells (Invitrogen) to express the SUMO–ETM fusion protein containing an N-terminal His6 tag (Extended Data Fig. 1a). The construct's DNA sequence was verified by Sanger sequencing (Genewiz).

Expression and purification of [13C,15N]ETM. A glycerol cell swamp was stored at −70°C was used to start a 10 ml LB culture containing 50 μg ml−1 kanamycin. The starter culture was used to inoculate 2 l of LB medium. Cells were grown at 37°C until an optical density at 600 nm (OD600) of 0.6–0.8 was reached, and was collected by centrifugation at 10 min at 20°C and 4,400g. These LB cells were resuspended in 11 of M9 medium (pH 7.8, 48 mM NaH2PO4, 22 mM KH2PO4, 8.6 mM NaCl, 4 mM MgSO4, 0.2 mM CaCl2, 50 mg kanamycin) containing 1 g/L [15N]NH4Cl. The cells were incubated in M9 media for 30 min at 15°C, then 1 g/L [13C]glucose dissolved in 5 ml sterile H2O and 3 ml 100 mM Tris-HCl buffer (pH 8.0, 50 mM Tris-HCl, 100 mM NaCl, 0.1% DDM, 250 mM imidazole) was added. The cells were grown for another 30 min, then protein expression was induced by addition of 0.4 mM IPTG along with 2 g/l [15N]glucose (10 mg l−1 of M9 medium). Labeling efficiency was ~94%, as estimated by MALDI mass spectrometry (Extended Data Fig. 1d). [13C]-ETM and [15N]-ETM were purified using preparative HPLC on a Varian ProStar 210 System using an Agilent C3 column (5-μm particle size, 21.2 mm × 150 mm). The protein was eluted using a linear gradient of 3 M urea) was used in place of wash buffer I.

Radiofrequency (RF) field strengths on the 3.2-mm probes were 50–91 kHz for 1H, 50–63 kHz for 13C and 33–42 kHz for 15N. RF field strengths on the 1.9-mm MAS probe were 83–130 kHz for 1H, 62.5 kHz for 13C and 71 kHz for 19F. Sample temperatures are direct readings from the probe thermocouple, whereas actual sample temperatures are 5–15 K higher at the MAS frequencies employed. 1H chemical shifts are reported on the tetramethylsilane scale using the adamantane chemical shift as an external standard. 13C chemical shifts are reported on the liquid ammonia scale using the N-acetylcyesteine peak at 122.0 ppm as an external standard.

2D 13C–13C correlation experiments were conducted using combined-driven (CORD) mixing for 13C spin diffusion. 2D and 3D 13C–1N correlation spectra, namely, NCA, NDCDX and CONCAW were measured on the 900-MHz spectrometer. These experiments used spectrally induced filtering in combination with cross-polarization (SPECIPOL) for heteronuclear polarization transfer. Water-edited 2D 31P–13C correlation spectra were measured under 11.8-kHz MAS velocity 20 h mixing times of 9 ms and 100 ms. 2D 13C–13C correlation spectra were measured using an out-and-back transferred-echo double resonance (TREDOR) pulse sequence on the 800 MHz NMR. Inter-molecular 2D NHHC correlation spectra were measured using 0.5 ms and 1 ms 1H–1H mixing. 1D and 2D 13C–15N REDOR experiments were used to measure distances between 13C–15N labeled and 13C–15N unlabeled ETM. Details for the solid-state NMR experiments are given in Supplementary Table 6. Details for the 13C–15N REDOR simulations and fitting are given in Supplementary Notes.

NMR spectral analysis. NMR spectra were measured on a Bruker AVANCE NEO 900 MHz (21.1 T) spectrometer and an Avance II 800 MHz (18.8 T) spectrometer using 3.2 mm HCN probes. [13C,15N]REDOR experiments were conducted on an Avance III HD 600 MHz (14.1 T) spectrometer using a 1.9 mm HFX probe. Magic-angle-spinning (MAS) frequencies were 14.5 kHz for 900-MHz experiments and 14 kHz for the 800- and 600-MHz experiments. Radiofrequency (RF) field strengths on the 3.2-mm probes were 50–91 kHz for 1H, 50–63 kHz for 13C and 33–42 kHz for 15N. RF field strengths on the 1.9-mm MAS probe were 83–130 kHz for 1H, 62.5 kHz for 13C and 71 kHz for 19F. Sample temperatures are direct readings from the probe thermocouple, whereas actual sample temperatures are 5–15 K higher at the MAS frequencies employed. 1H chemical shifts are reported on the tetramethylsilane scale using the adamantane chemical shift as an external standard. 13C chemical shifts are reported on the liquid ammonia scale using the N-acetylcyesteine peak at 122.0 ppm as an external standard.

3D 1H–15N correlation experiments were conducted using combined-driven (CORD) mixing for 1H spin diffusion. 2D and 3D 1H–15N correlation spectra, namely, NCA, NDCDX and CONCAW were measured on the 900-MHz spectrometer. These experiments used spectrally induced filtering in combination with cross-polarization (SPECIPOL) for heteronuclear polarization transfer. Water-edited 2D 31P–13C correlation spectra were measured under 11.8-kHz MAS velocity 20 h mixing times of 9 ms and 100 ms. 2D 13C–13C correlation spectra were measured using an out-and-back transferred-echo double resonance (TREDOR) pulse sequence on the 800 MHz NMR. Inter-molecular 2D NHHC correlation spectra were measured using 0.5 ms and 1 ms 1H–1H mixing. 1D and 2D 13C–15N REDOR experiments were used to measure distances between 13C–15N labeled and 13C–15N unlabeled ETM. Details for the solid-state NMR experiments are given in Supplementary Table 6. Details for the 13C–15N REDOR simulations and fitting are given in Supplementary Notes.

NMR spectral analysis. NMR spectra were measured on the TopSpin software and chemical shifts were assigned in Sparky.10 TALOS-N was used to calculate torsion angles (φ, ψ) after computing the chemical shifts with the torsion angles. Residue-specific chemical shift differences (Δδ) between drug-bound and apo samples were calculated from the measured 13C and 15N chemical shifts (6) according to:

$$\Delta \delta = \sqrt{\sum_{i} (\delta_{\text{drug}} - \delta_{\text{apo}})^2 + (\delta_{\text{1H}} - \delta_{\text{1H}})^2}$$

(1)

2D heatmaps of normalized water-edited 2D NCA spectra were generated using an in-house Python script that removes spectral noise while calculating integrated intensities. The intensities were divided and scaled by the number of scans to obtain a 2D contour map that reflects the peak intensity ratios between the 9 ms and 100 ms spectra.

The water accessibility data for the high-pH influenza RM2 proton channel (Fig. 2f) were originally measured in 2D 13C–13C correlation spectra with 4 ms (S) and 100 ms (S) H spin diffusion. To allow comparison with the ETM spectra measured at 9 ms and 100 ms mixing, we scaled the RM2 S (4 ms)/S (100 ms) ratios by the intensity of the aliphatic intensity ratio of 1.976 between the 9 ms and 100 ms spectra. This scaling factor was verified to be accurate for two resolved sites, Thr24 and Gly26, in the 13C spectra.

XPLOR-NIH structure calculations and analysis. Initial structure calculation using ambiguous interhelical restraints, where each helix can contact both
neighboring helices, did not converge. Thus, we generated parallel pentamer models to specify the direction of $^{13}$C–$^{19}$F and NHHC distance restraints where possible. The models take into account the water- and lipid-edited spectra to qualitatively identify the pore- versus lipid-facing residues. The best-case ideal helix model (Extended Data Fig. 7a), with 3.5 residues per helical turn, places Asn15 at the pore-facing $d$ position and Phe20 at the lipid-facing $b$ position, in agreement with the water-edited spectra. However, the model conflicts substantially with other data, for example, Thr35($c$) (Thr35 at position $c$) and Leu31($f$) are lipid-facing in this model, which contradicts the water-edited spectra; Val29($d$) and Phe26($a$) are pore-facing, which contradicts the water- and lipid-edited spectra. The arc of Phe20($b$), Phe23($e$) and Phe6($a$) on the helical wheel makes it unlikely to establish interhelical Phe–Phe contacts, thus contradicting the $^{13}$C–$^{19}$F distance data.

Since the ideal-helix geometry cannot agree with all experimental data, we sought another model by including slight deviations from an ideal helix. We turned to the measured chemical shifts to determine where such a deviation is most likely to occur. The Cβ chemical shift of L21 is 1.4 ppm downfield from the average of all other helical Leu residues (Extended Data Fig. 3b), suggesting that the helix is disordered between Phe20 and Phe23. Indeed, such a disorder was already noted in previous solution NMR data. We generated four alternative pentamer models with varying positions and degrees of helix disorder (Extended Data Fig. 7b–e and Supplementary Table 4). Only one model (model 5), generated by a small rotation angle advance of $\sim 50^\circ$ at Phe23, adequately reproduces all key features of the experimental data. This model places Asn15($d$) and Val25($d$) at the solvent-exposed and the three aromatic residues at the helix core, Phe20($e$), Phe23($f$) and Phe26($a$). This model was then used to disambiguate the NHHC and $^{13}$C–$^{19}$F distance restraints (Supplementary Tables 2 and 3) by mainly considering only residues that are fewer than four residues away in the primary sequence and that are in close proximity between two helical wheels. With this approach, 42 of the 87 interhelical restraints were set to be unambiguous. In principle, the handedness of the helical bundle can be determined from the determinate families of interhelical contacts if the position of interfacial residues are known. However, remaining $^{13}$C and $^{15}$N chemical-shift overlap among the many hydrophobic residues precluded unequivocal determination of the handedness of the helical bundle. Orthogonal experimental constraints, such as backbone N-H bond orientations, which would directly probe the helix tilt angle, will be needed to obtain a higher-resolution structure.

As has been previously described, the ETM structure was calculated using XPLOR-NIH $^\dagger$ hosted on the NMRbox. The calculation contained two stages. In the first stage, five extended ETM monomers were placed in a parallel pentamer geometry, with each monomer located 20 Å from the center of the pentamer. A total of 120 independent simulated annealing runs were performed with 5,000 steps of torsion angle dynamics at 1,000 K followed by energy minimizations in torsion angle and Cartesian coordinates. The first stage, five extended ETM monomers were placed in a parallel pentamer geometry, with each monomer located 20 Å from the center of the pentamer. A total of 120 independent simulated annealing runs were performed with 5,000 steps of torsion angle dynamics at 5,000 K, followed by annealing to 20 K in decrements of 20 K with 100 steps at each temperature. After the annealing, energy minimizations in torsion angle and Cartesian coordinates were carried out. The five monomers were restrained to be identical in the annealing step using the non-crystallographic symmetry term DistSymmPot and the translation symmetry term DistSymmPot. Chemical-shift-derived torsion angles (φ, ϕ) predicted by TALOS-N were implemented with the dihedral-angle restraint term CDIH, with ranges set to the higher value between twice the TALOS-N predicted uncertainty and 20°. Measured interhelical distance restraints were implemented using the NOE potential with an upper limit set to 2.9 Å and 11.5 Å for 500 μs and 1,000 μs of $^{1}H$-$^{1}H$ mixing for the NHHC constraints. Negative REDOR contacts, that is, $^{13}$C sites without dephasing, were implemented as two NOE's: one performed with 5,000 steps of torsion angle dynamics at 1,000 K followed by energy minimizations in torsion angle and Cartesian coordinates. A total of 64 potential terms were used as independent inputs for structure refinement. A total of 64 potential terms were used as independent inputs for structure refinement. A total of 64 potential terms were used as independent inputs for structure refinement. A total of 64 potential terms were used as independent inputs for structure refinement. A total of 64 potential terms were used as independent inputs for structure refinement. A total of 64 potential terms were used as independent inputs for structure refinement. All the potentials employed in annealing were also used during refinement, with the exception of the HBDB potential term HBDB were implemented during annealing to favor the formation of the helical wheel makes it unlikely to establish interhelical Phe–Phe contacts, thus contradicting the $^{13}$C–$^{19}$F distance data. Since the ideal-helix geometry cannot agree with all experimental data, we sought another model by including slight deviations from an ideal helix. We turned to the measured chemical shifts to determine where such a deviation is most likely to occur. The Cβ chemical shift of L21 is 1.4 ppm downfield from the average of all other helical Leu residues (Extended Data Fig. 3b), suggesting that the helix is disordered between Phe20 and Phe23. Indeed, such a disorder was already noted in previous solution NMR data. We generated four alternative pentamer models with varying positions and degrees of helix disorder (Extended Data Fig. 7b–e and Supplementary Table 4). Only one model (model 5), generated by a small rotation angle advance of $\sim 50^\circ$ at Phe23, adequately reproduces all key features of the experimental data. This model places Asn15($d$) and Val25($d$) at the solvent-exposed and the three aromatic residues at the helix core, Phe20($e$), Phe23($f$) and Phe26($a$). This model was then used to disambiguate the NHHC and $^{13}$C–$^{19}$F distance restraints (Supplementary Tables 2 and 3) by mainly considering only residues that are fewer than four residues away in the primary sequence and that are in close proximity between two helical wheels. With this approach, 42 of the 87 interhelical restraints were set to be unambiguous. In principle, the handedness of the helical bundle can be determined from the determinate families of interhelical contacts if the position of interfacial residues are known. However, remaining $^{13}$C and $^{15}$N chemical-shift overlap among the many hydrophobic residues precluded unequivocal determination of the handedness of the helical bundle. Orthogonal experimental constraints, such as backbone N-H bond orientations, which would directly probe the helix tilt angle, will be needed to obtain a higher-resolution structure. As has been previously described, the ETM structure was calculated using XPLOR-NIH $^\dagger$ hosted on the NMRbox. The calculation contained two stages. In the first stage, five extended ETM monomers were placed in a parallel pentamer geometry, with each monomer located 20 Å from the center of the pentamer. A total of 120 independent simulated annealing runs were performed with 5,000 steps of torsion angle dynamics at 5,000 K, followed by annealing to 20 K in decrements of 20 K with 100 steps at each temperature. After the annealing, energy minimizations in torsion angle and Cartesian coordinates were carried out. The five monomers were restrained to be identical in the annealing step using the non-crystallographic symmetry term DistSymmPot and the translation symmetry term DistSymmPot. Chemical-shift-derived torsion angles (φ, ϕ) predicted by TALOS-N were implemented with the dihedral-angle restraint term CDIH, with ranges set to the higher value between twice the TALOS-N predicted uncertainty and 20°. Measured interhelical distance restraints were implemented using the NOE potential with an upper limit set to 2.9 Å and 11.5 Å for 500 μs and 1,000 μs of $^{1}H$-$^{1}H$ mixing for the NHHC constraints. Negative REDOR contacts, that is, $^{13}$C sites without dephasing, were implemented as two NOE's: one to each neighboring helix. Implicit hydrogen bonds using the hydrogen-bonding database potential term HBDB were implemented during annealing to favor the formation of the α-helical conformation. Finally, standard XPLOR potentials were used to restrain the torsion angles using a structural database with the term TorsionDB, and standard bond angles and lengths were set with terms BOND, ANGL, IMPR, Repel and Repel14 potentials, while excluding PosDiffPot, DistSymmPot and TorsionDB potentials. The ten structures with the lowest energies across the specified potentials were included in the final structural ensemble. Where lipid-contact structures are shown, the most representative conformer, selected as the model with the lowest average r.m.s.d. for residues 10–36 with respect to all the other structural models, is shown. The Ramachandran plot statistics for the final structure ensemble are as follows: 93% of residues are in favored regions, 5% of residues are in allowed regions and 2% of residues are in disallowed regions. The only outlier is Leu37, which is outside the TM helix, near the C terminus.

Graphical images depicting the structures were generated in PyMOL v2.3.4. The reported channel radii were calculated using the HOLE program $^\ddagger$, and represent the radii of the largest sphere that can be accommodated from exclusion of the van der Waals diameter of all atoms at each XY plane along the Z channel coordinate, which is collinear with the bilayer normal and the putative direction of ion permeation. The cutoff radius for the calculation was 5 Å. The HOLE output was visualized in PyMOL by setting the van der Waals radius of the HOLE-generated spheres ‘SPH’ to the B-factor values of the SPH output. Details of HMA docking to ETM are given in Supplementary Notes.

### Data availability
NMR chemical shifts, distance and torsion-angle restraints have been deposited in the Biological Magnetic Resonance Bank (BMRB) with ID numbers 30795. The structural coordinates for ETM have been deposited in the Protein Data Bank with the accession code 7KKG.

### Code availability
NMR pulse programs and in-house Python scripts used for structure calculation and data analysis such as water-edited spectral analysis are available upon request.

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**Author contributions**

M.H. designed the project and supervised experiments and data analysis. V.S.M., M.J.M. and A.A.S. cloned, expressed and purified the protein and conducted the solid-state NMR experiments. V.S.M. and M.J.M. assigned and analyzed the spectra. V.S.M. calculated the structure with contributions from A.J.D. and M.J.M. A.A.S. conducted $^{19}$F NMR experiments, simulations and docking. A.K. synthesized fluorinated amantadine. All authors discussed the results of the study and wrote the paper.

**Competing interests**

The authors declare no competing interests.

**Additional information**

Extended data is available for this paper at [https://doi.org/10.1038/s41594-020-00536-8](https://doi.org/10.1038/s41594-020-00536-8).

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Extended Data Fig. 1 | Cloning, purification and characterization of ETM. a, Amino acid sequence of SUMO-tagged ETM. b, SDS-PAGE gel showing purification of ETM by nickel affinity column chromatography. The flowthrough contains all soluble proteins that have low affinity for nickel. The column was washed with 30 mM imidazole, and SUMO-ETM (18 kDa band) was eluted at >90% purity with 250 mM imidazole. High molecular-weight SUMO-ETM oligomers are visible as a minor species. ETM was cleaved from the SUMO tag using SUMO protease. c, Preparative reverse-phase HPLC chromatogram after protease cleavage. ETM elutes at 37.5 min. d, MALDI mass spectrum of purified U-13C, 15N labeled ETM. e, MALDI mass spectrum of purified 4-19F-Phe labeled ETM. The measured masses are in good agreement with the theoretical masses. 83% of the 4-19F-Phe labeled ETM monomers have all three Phe residues fluorinated, indicating a per-site labeling efficiency of 94%.
Extended Data Fig. 2 | See next page for caption.
Extended Data Fig. 2 | Effects of temperature and membrane composition on ETM structure. a, $^{13}$C and $^{15}$N CP-MAS spectra of ERGIC-bound ETM. The spectra show high sensitivity and resolution, indicating a well ordered and rigid protein. b, $^{13}$C and $^{15}$N CP-MAS spectra of ETM in DMPX membranes from 303 K to 263 K. The spectral intensities and linewidths are insensitive to temperature, indicating that the protein is mostly immobilized at ambient temperature. c, $^{13}$C direct-polarization (DP) spectra of DMPX-bound ETM. The E8 sidechain carboxyl chemical shift changes between high and low pH, indicating that this residue is protonated at low pH. d-f, 2D $^{15}$N-$^{13}$C (left) and $^{13}$C-$^{13}$C (right) correlation spectra of ETM at high and low temperatures and in ERGIC versus DMPX membranes. Yellow rectangles highlight peaks with clear chemical shift or intensity changes. d, 2D spectra of ERGIC-bound ETM (orange) at 293 K and DMPX-bound ETM at 303 K (green). The chemical shifts are similar, indicating that the protein conformation is unaffected by the presence of POPS, POP1 and cholesterol. T11 and L12 signals are not detected in the ERGIC sample at this temperature, suggesting that the N-terminus is mobile under these conditions. e, 2D spectra of ERGIC-bound ETM at 293 K (orange) and 263 K (blue). Moderate chemical shift changes are observed for C-terminal residues from T35 to R38, while the I13 signal is not visible at low temperature. f, 2D spectra of DMPX-bound ETM at 303 K (green) and 263 K (purple). The C-terminal residues exhibit temperature-dependent chemical shifts, similar to the ERGIC-bound peptide. The N-terminal residues of T9 to I13 do not exhibit signals at 263 K, indicating that the N-terminus undergoes intermediate-timescale motion at this temperature. Thus, the C-terminal conformation is temperature-dependent while the N-terminus is dynamic at high temperature.
Extended Data Fig. 3 | Chemical shift assignment and secondary structure of ETM. 

**a.** Representative strips from 3D NCACX (magenta), CONCA (green) and NCOCX (blue) spectra of ERGIC-membrane bound ETM. These spectra allow full assignment of the $^{13}$C and $^{15}$N chemical shifts. 

**b.** $C\alpha$ (blue) and $C\beta$ (orange) secondary chemical shifts compared to random coil chemical shifts. Most residues show positive $C\alpha$ and negative $C\beta$ secondary shifts, indicating an $\alpha$-helical conformation.

**c.** $(\phi, \psi)$ torsion angles calculated using TALOS-N. Residues G10 to L34 show $\alpha$-helical conformation. Error bars represent the precision of the TALOS-N prediction, defined as one standard deviation for the $(\phi, \psi)$ angles among the best-matched peptides for each residue.
Extended Data Fig. 4 | Effects of pH and ions on the chemical shifts of DMPX-membrane bound ETM. Where cations are present, the ion concentration is 5 mM. a. 2D $^{15}$N-$^1$H correlation spectra of high-pH ETM with 5 mM NaCl and low-pH ETM with 5 mM CaCl$_2$. Chemical shift changes are observed for C-terminal residues such as R38, L37 and L34 (yellow highlight). b. 2D $^{13}$C-$^{13}$C correlation spectra of low-pH ETM with CaCl$_2$ and high-pH ETM with NaCl. c. 2D $^{13}$C-$^{13}$C correlation spectrum of low-pH ETM with CaCl$_2$ and low-pH ETM without salt. These spectra indicate that the chemical shift changes mainly result from pH changes.
Extended Data Fig. 5 | Additional 13C-19F REDOR spectra and water-edited spectra to determine the interhelical packing of ETM. a, 2D 13C-13C correlation spectrum of mixed 4-19F-Phe labeled and U-13C,15N-labeled eTM (black). The 13C chemical shifts of most residues are similar to the 13C,15N-labeled protein (red), indicating that fluorination does not perturb the ETM conformation. F20/23 Cβ, F26 Cβ, and T30 Cγ2 show small chemical shift changes (blue) of 0.3–0.6 ppm. The spectra were measured at 293 K. b, 1D 13C-19F REDOR control (S₀), dephased (S), and difference (ΔS) spectra. The difference peaks result from carbons that are in close proximity to a fluorine in a neighboring helix. The broadband REDOR spectra (left) show both sidechain and backbone 13C signals whereas the Cα-selective REDOR spectra (right) detect only Cα signals. c, Representative 13C-19F REDOR dephasing curves for broadband and Cα-selective C-F REDOR spectra. The S/S₀ values have been corrected for the isotopic dilution factor (50%) and the peak-overlap factor. Best-fit distance curves are shown as solid lines, and lower and upper distance bounds are shown as dashed lines. Error bars represent random uncertainty of the measured S/S₀ values, which were propagated from the signal-to-noise ratios of the S₀ and S spectra. d, Water-edited 2D 15N-13Cα correlation spectra to detect well hydrated residues. The spectra were measured at 293 K under 11.8 kHz MAS using 1H-1H mixing times of 9 ms (red) and 100 ms (blue).
Extended Data Fig. 6 | Inter-residue correlations obtained from 250 ms 2D $^{13}$C spin diffusion spectra of ERGIC-membrane bound ETM. a, Representative strips from a well-resolved 3D NCACX spectrum recorded with 250 ms $^{13}$C spin diffusion. Inter-residue cross peaks are assigned in black and intra-residue resonances are marked in blue. b, Overlay of 2D $^{13}$C-$^{13}$C correlation spectra measured with 250 ms mixing (black) and 20 ms (orange). Representative inter-residue cross peaks are assigned in blue. All spectra were measured at 293 K under 11.8 kHz MAS.
Extended Data Fig. 7 | ETM pentameric models analyzed to disambiguate the direction of interhelical constraints used for structure calculation. For each model, the heptad repeat positions (abcdefg) of every residue from L12 to T35 is indicated on the helical wheel for at least one subunit. On the two neighboring helices, residue positions that violate measured $^{13}$C-$^{19}$F correlations are shown in pink, while residue positions that violate the water and lipid accessibility data are shown in green. The positions of Phe residues that satisfy the interhelical contacts are shown in blue.

**a**, Model 1 places N15 at heptad position d without a twist, and is thus an ideal helix model. **b**, Model 2 places N15 at d with a twist such that F23 moves from position e to c. **c**, Model 3 places N15 at position e with a twist such that F23 moves from f to b. **d**, Model 4 places N15 at position a with a twist such that F23 moves from b to c. **e**, Model 5 places N15 at position a with a twist such that F23 moves from b to f. Model 5 does not violate any experimental data and was thus chosen to disambiguate intermolecular contacts for structure calculation. To make the interhelical contacts explicit, model 5 shows the residue positions for three consecutive helices in the pentamer.
Extended Data Fig. 8 | See next page for caption.
Extended Data Fig. 8 | Lipid-bilayer bound SARS-CoV-2 ETM structure (PDB code: 7K3G) and its comparison with ETM structure solved in micelles and with other viroporin structures. a, N-terminal top views of various residues in the ETM pentamer. Most residues are hydrophobic, including both pore-facing and lipid-facing residues. The most representative structure of the lowest-energy ensemble is shown. b, Top views of representative pore-facing residues in the lowest-energy ensemble. The structure distribution is likely due to a combination of the sparseness of experimental restraints and true protein conformational disorder. c, Comparison of the ERGIC-membrane bound ETM structure model (slate and red) and the LMPG-micelle-bound ETM structure model (gray and salmon). Side view depicts differences in helix orientation and helical bundle handedness, while top view shows differences in pore radii. d, Structural comparison of the pentameric ETM channel, the closed tetrameric influenza BM2 proton channel, and the pentameric HIV-1 Vpu channel. The ETM pentamer is longer and tighter than the BM2 and Vpu helical bundles.
Extended Data Fig. 9 | Additional docking poses of HMA to SARS-CoV-2 E, shown in side view (left) and N-terminal top view (right). a, Structure with hexamethylene ring up and HMA vertical, obtained from docking in DMSO. b, Structure with hexamethylene ring down and HMA vertical, obtained from docking in DMSO. c, Structure with HMA across the channel entrance, bridging two helices, obtained from docking in water. The lipid-facing I13 and pore-occluding N15 are shown in sticks to guide the eye.
Extended Data Fig. 10 | Effects of amantadine binding on ETM. The peptide is reconstituted in DMPC: DMPG membranes with an AMT: ETM monomer molar ratio of 8:1.  

**a**, 2D $^{15}$N-$^{13}$C correlation spectra of apo (blue) and AMT-bound ETM (magenta). The spectra were measured at 305 K under 14 kHz MAS. Zoomed-in areas show peaks with significant CSPs.  

**b**, 2D $^{13}$C-$^{13}$C correlation spectra with 20 ms mixing of apo (blue) and AMT-bound ETM (magenta). The spectra were measured at 263 K. Zoomed-in areas shows peaks with significant CSPs. The perturbed residues are concentrated in the N- and C-termini of the protein.  

**c**, 1D $^{19}$F direct-polarization spectra of 3F-AMT with and without the peptide in DMPX membranes. The spectra were measured at 270 K under 14 kHz MAS.  

**d**, $^{13}$C$^{\alpha}$ selective $^{19}$F-dephased REDOR spectra of AMT-bound ETM in DMPC: DMPG membranes. The $\Delta S$ spectra show dephasing at 65.5 ppm, 63.6 ppm, 56 ppm and 54 ppm.  

**e**, Broadband $^{13}$C-$^{19}$F REDOR spectra. The $\Delta S$ spectra show $^{13}$C dephasing for sidechains that belong to residues that show $^{\alpha}$C dephasing in (d).
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