

Solid-state NMR and simulation studies of equinatoxin II N-terminus interaction with lipid bilayers

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ABSTRACT

The interaction with model membranes of a peptide, EqtII_{1–32}, corresponding to the N-terminal region of the pore-forming toxin equinatoxin II (EqII) has been studied using solid-state NMR and molecular dynamics (MD) simulations. The distances between specifically labeled nuclei in [¹⁹F-para]Phe16-[1-¹³C]Leu19 and [¹⁹F-para]Phe16-[¹⁵N]Leu23 analogs of EqtII_{1–32} measured by REDOR in lyophilized peptide were in agreement with published crystal and solution structures. However, in both DMPC and mixed DMPC:SM membrane environments, significant changes in the distances between the labeled amino acid pairs were observed, suggesting changes in helical content around the experimentally studied region, 16–23, in the presence of bilayers. ¹⁹F-³¹P REDOR experiments indicated that the aromatic ring of Phe16 is in contact with lipid headgroups in both membrane environments. For the DMPC:SM mixed bilayers, a closer interaction between Phe16 side chains and lipid headgroups was observed, but an increase in distances was observed for both labeled amino acid pairs compared with those measured for EqtII_{1–32} in pure DMPC bilayers. The observed differences between DMPC and DMPC:SM bilayers may be due to the greater affinity of EqII for the latter. MD simulations of EqtII_{1–32} in water, on a pure DMPC bilayer, and on a mixed DMPC:SM bilayer indicate significant peptide secondary structural differences in the different environments, with the DMPC-bound peptide adopting helical formations at residues 16–24, whereas the DMPC:SM-bound peptide exhibits a longer helical stretch, which may contribute to its enhanced activity against PC:SM compared with pure PC bilayers.

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Key words: membranes; binding; phosphatidylcholine; sphingomyelin; surface properties.

INTRODUCTION

Equinatoxin II (EqII) is a 179-residue, 19.8 kDa cytolytic protein isolated from the Mediterranean anemone *Actinia equina* L.¹ EqII is a member of the family of actinoporins, which is characterized by a high pI, affinity for sphingomyelin, and permeabilizing activity of model and cell membranes.^{2–6} EqII is highly soluble in water, and its structure has been well characterized by X-ray crystallography and solution state NMR.^{7,8} The structure consists of two short helices packed against opposite faces of a β-sandwich structure formed by a pair of five-stranded β-sheets. The core of the protein has extensive hydrophobic interfaces formed by residues projecting from the internal faces of the two β-sheets.

The N-terminal 32 residues, including an amphiphilic helix, adopt a different structure upon interaction with membrane bilayers,⁹ but without disrupting the overall fold of the rest of the protein.^{10,11} A cluster of exposed aromatic amino acid residues on the surface of EqII, consisting of Tyr108, Trp112, Tyr113, Trp116, Tyr133, Tyr137, and Tyr138, is likely to have a role in membrane recognition and binding since aromatic residues have been shown to have affinity for the lipid–water interface.^{12–14} CD and FTIR studies show small increases in β-sheet and α-helical content at the expense of random structure in the presence of unilamellar vesicles.^{10,15–17} In principle, at least two steps are involved in formation of the EqII pore, with initial binding of monomers to the membrane being followed by subsequent oligomerization into a functional pore. Site-directed mutagenesis has been used to show that at least

Abbreviations: DMPC, dimyristoylphosphatidylcholine; EqII, equinatoxin II; MD, molecular dynamics; NMR, nuclear magnetic resonance; REDOR, rotational echo double resonance; SM, sphingomyelin; SS, solid-state.

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two regions of EqtlI became embedded in lipid membranes, the N-terminal region (residues 10–28)¹⁸ and the surface aromatic cluster including Trp112 and Trp116.^{10,11,15,19} A model of the membrane-bound monomer envisages that the N-terminal helix involving residues 11–33, the hydrophobic region (residues 105–120), and Arg144, Ser160, and nearby residues, interact with the lipid membrane.¹⁰ In the next step, the N-terminal helix translocates from the surface of the β -sandwich to the membrane, and finally traverses the lipid bilayer to form a cation-selective pore.^{5,6,15}

Although these models suggest which regions of EqtlI might interact with lipid membranes and undergo conformational changes to form the pore, little direct experimental evidence has been obtained concerning the conformational changes EqtlI undergoes in reaching an oligomeric state, or during its incorporation into the target membrane. Recently, a series of peptides corresponding to the N-terminal region of EqtlI has been studied and changes in secondary structure upon interaction with dodecylphosphocholine (DPC) micelles detected.⁹ These studies indicate that the N-terminal peptide is sensitive to its environment, especially when a membrane surface (or mimetic) is present.

In this study, solid-state nuclear magnetic resonance (SS NMR) has been used to investigate the N-terminal domain of EqtlI (EqtlI_{1–32}) and its interaction with DMPC bilayers and DMPC:SM mixed bilayers. These serve as model systems to help elucidate the initial stages of membrane permeabilization by actinoporins. Molecular dynamics (MD) simulations have been applied to provide further atomic-level insight into the interactions between the EqtlI_{1–32} peptide, starting from a number of different initial conformations, and membrane bilayer models (DMPC and a mixed DMPC:sphingomyelin bilayer). MD simulations were also performed to study the interactions between EqtlI placed initially in the water phase and a DMPC bilayer as a basis for comparison with our results on the N-terminal fragment peptide. Our computational simulations complement experimental studies; whereas, SS NMR data elucidate structural features of the final state of EqtlI_{1–32} interactions with DMPC and PC:SM membranes, MD simulations elucidate the structure, dynamics, and lipid contacts for the peptide and protein in the initial stages of interaction with the membranes. Additionally, a constrained simulation of the N-terminal fragment peptide on a DMPC bilayer with distance restraints derived from SS NMR provides further insights into the structure of the peptide.

MATERIALS AND METHODS

Materials

[¹⁹F-para]Phe was purchased from Sigma-Aldrich (St Louis, MO) and [1-¹³C]Leu and [¹⁵N]Leu from

Cambridge Isotopes (Andover, MA). [¹⁹F-para]Phe16, [1-¹³C]Leu19, and [¹⁵N]Leu23 labeled N-terminal peptide of EqtlI (EqtlI_{1–32}) were supplied by Mimotopes (Melbourne, Australia) with purity >95% as determined by HPLC and MS. SM and DMPC were purchased from Avanti Polar Lipids (Alabaster, AL).

Sample preparation

Lipids (DMPC or DMPC:SM) were mixed in a chloroform/methanol (2:1; vol:vol) at 10 mg/mL in a round bottom flask. The solvent was removed using a rotary evaporator and the lipid mixtures were dried under high vacuum for 3–5 h. Lyophilized EqtlI_{1–32} was dissolved in water at a concentration of 5 mg/mL (pH ~6.5) and appropriate amounts used to give the desired toxin-lipid mole ratio (DMPC:SM:EqtlI_{1–32}, 30:30:1 or DMPC:EqtlI_{1–32}, 30:1). Each sample was subsequently put through five cycles of rapid freezing in liquid nitrogen and equilibration at 50°C. The multilamellar vesicle (MLV) suspensions were then concentrated in a Beckman benchtop centrifuge (28,000 rpm, 1 h, 4°C). The pellet (~100 μ L) was loaded directly into a 4-mm NMR zirconia rotor for the magic angle spinning (MAS) NMR experiments.

Solid-state NMR spectroscopy

Heteronuclear distances were measured using REDOR^{20,21} SS NMR on a CMX500 spectrometer (Chemagnetics, Varian, Palo Alto, CA). Samples were packed in 4-mm rotors and measured using a T₃ HFX MAS probe (Chemagnetics, Varian, Palo Alto, CA).

REDOR experiments were carried out in two steps: once with rotor-synchronized dephasing pulses (S_r) and once without (S_0). The dephasing pulse changes the sign of the heteronuclear dipolar coupling, and this interferes with the spatial averaging resulting from the motion of the rotor.^{20,21} The difference in signal intensity ($\Delta S = S_0 - S_r$) for the observed spin is directly related to the corresponding distance to the dephasing spin. NMR data were analyzed using SIMPSON.²² Heteronuclear distance measurements were carried out at room temperature on a lyophilized powder and at low temperature (–60°C) on membrane-bound peptide at 8 kHz spinning speed. Parameters used for ¹³C-¹⁹F REDOR experiments were as follows: recycle delay 14 s, 90° ¹H pulse width of 3.2 μ s, ¹⁹F 180° pulse 6.2 μ s, ¹³C 180° pulse 6.8 μ s, 82 kHz TPPM decoupling power, dephasing time 0.5–20 ms, and the number of scans was 4–18 k and the XY-32 pulse phase cycling scheme was used. ¹³C chemical shifts were referenced to hexamethylbenzene (HMB) at 17.36 ppm, and ¹⁹F chemical shifts referenced to trifluoroethanol (TFE) at 0 ppm. For ³¹P-¹⁹F REDOR experiments, the parameters used were similar but with ¹⁹F 180° pulse 7.1 μ s and ³¹P 180° pulse 7.5 μ s. ³¹P chemical shifts were referenced to 85% H₃PO₄ (0 ppm). For ¹⁵N-¹⁹F REDOR

experiments, similar parameters were used but with a 90° ^1H pulse width of 2.93 μs , ^{19}F 180° pulse 6.6 μs , and ^{15}N 180° pulse 8.65 μs . ^{15}N chemical shifts were referenced to saturated $^{15}\text{NH}_4\text{Cl}$ at 0 ppm.

Molecular dynamics simulation methodology

Molecular dynamics (MD) simulations were performed on EqII₁₋₃₂ in the vicinity of DMPC and 1:1 ratio mixed DMPC and sphingomyelin (SM18:0) bilayers, with the systems solvated with SPC²³ water molecules and counterions included to ensure overall charge neutrality. A simulation was also performed on full-length EqII in the vicinity of a DMPC bilayer as a comparison with the peptide SS NMR and MD simulation results. All MD simulations were performed using the program GROMACS version 3.1.4^{24,25} under constant particle number, pressure, and temperature (NPT) conditions. The GROMOS96²⁴ force field with parameter set 43a1 were employed for all simulations. Lipid parameters for DMPC were adopted from Berger *et al.*²⁶ and those for sphingomyelin adopted from Chiu *et al.*²⁷ Temperature and pressure coupling were performed using the scheme described by Berendsen *et al.*²⁸ A constant pressure of 1 bar and pressure coupling constant of 1.0 ps were applied uniformly for the pure water systems, whereas for the bilayer simulations, separate pressure coupling was applied to the *xy*-plane and *z*-directions (i.e., the bilayer plane and normal, respectively). Water, DMPC, SM (for the mixed bilayer simulation), and protein were coupled separately to a temperature bath at 310 K with a temperature coupling constant of 0.1 ps. Electrostatic interactions were evaluated using the particle-mesh Ewald (PME) method²⁹ with van der Waals interactions truncated at 10 Å. Integration time steps of 2 fs were used. Bond lengths were constrained using the LINCS algorithm.³⁰ Analyses of MD trajectories were performed using the GROMACS suite of programs. Secondary structure analysis employed DSSP.³¹ Visualization of system geometries and interactions and evaluation of protein secondary structure were performed using VMD.³²

The simulation systems were set up as follows: for the full-length protein (labeled **prot-DMPC**), the atomic coordinates of the first monomer of the structure included in PDB file 1IAZ,⁷ simulated in pure SPC water for 10 ns (no significant structural changes were evident; data not shown), were used as initial coordinates. The protein was placed initially ~ 20 Å above the headgroup region of the 288 lipid DMPC bilayer [initial structure in Fig. 6(g)] and simulated for 50 ns. For the EqII₁₋₃₂/DMPC simulations, we used two different starting conformations and performed two separate unbiased simulations (i.e., no noncovalent distance constraints were applied). The first peptide simulation used the structure contained in BioMagResBank^{33,34} with accession number 6808, labeled **pep-exp-DMPC** [Fig. 6(a)] and was

simulated for 80 ns. The second peptide simulation used a structure obtained from a 10 ns simulation of the experimentally obtained structure in pure SPC water (data not shown), which produced significant loss of helicity at both termini [Fig. 6(d)]. This simulation is labeled **pep-wat-DMPC** and was also simulated for 80 ns. In both EqII₁₋₃₂/DMPC simulations, the peptide was initially placed ~ 35 Å above a 128 lipid DMPC bilayer. Thus, two simulations with significantly different starting conformations were performed. Simulations using different starting conformations were used to enhance conformational sampling and improve the statistical relevance of any structural details and peptide-bilayer contacts revealed through the course of the simulations. The **pep-exp** structure was chosen because it is the best known approximation of the structure of the peptide in a lipid-rich environment, whereas the **pep-wat** structure was chosen as an approximation of the peptide structure before contact with bilayers. Simulation of the latter structure enables examination of the binding mechanism of an initially fully solvated peptide to a bilayer surface. We also performed a distance-constrained simulation of EqII₁₋₃₂, in which the initial structure was taken from the **pep-pdb-DMPC** simulation at ~ 50 ns at which the best fits to experimental distance measurements were obtained. Throughout the 10 ns simulation, distance constraints of 5 and 8.2 Å were enforced between Phe19 (sidechain H ζ) and both Leu19 (backbone C) and Leu23 (backbone N), respectively. This simulation is labeled **pep-cons-DMPC** and was simulated for 15 ns. Applying constraints to an already equilibrated (i.e., stable) structure which possesses experimentally consistent inter-residue distances enables the study of the peptide structure that corresponds best to the SS NMR-determined structure in DMPC, while preventing introduction of large artificial biasing forces that might cause unphysical distortion of the peptide. This approach also enables examination of regions of the peptide other than those for which experimental measurements are available, while ensuring consistency with those same measurements.

For the EqII₁₋₃₂/DMPC:SM bilayer simulation (labeled **pep-exp-mixed** simulated for 70 ns), the BMRB-6808 peptide structure was used as the starting structure, placed initially ~ 35 Å above a mixed DMPC:SM bilayer. The mixed bilayer is composed of 32 molecules of each lipid type on each bilayer leaflet and is generated from a pure DMPC bilayer by random replacement of DMPC with SM molecules. A subsequent 10 ns simulation run of the peptide-free mixed bilayer was performed, after which the final bilayer structure was acquired for the peptide-interaction simulation. An initial 100 ps solvent equilibration run with protein and lipid atoms positionally restrained for all simulations, followed by 100 ps equilibration with protein atoms only restrained. This was followed between 50 (**prot-DMPC**) and 80 ns

(peptide simulations) of restraint-free (collection) simulations, the results of which we discuss in the following sections.

RESULTS AND DISCUSSION

Solid-state NMR

Heteronuclear distance measurements for lyophilized EqtlI_{1–32}

Figure 1(a) shows the structure of EqtlI and Figure 1(b) shows the structure of the N-terminus of EqtlI_{1–32} in DPC micelles, which is thought to represent a structure involved in an early step of membrane–protein interaction. Intermolecular distances for [¹⁹F-para]Phe16–[1-¹³C]Leu19 EqtlI_{1–32} and [¹⁹F-para]Phe16–[¹⁵N]Leu23 EqtlI_{1–32} pairs were determined by REDOR NMR. Figure 2(a–f) shows the ¹⁹F–¹³C REDOR of both dephasing and nondephasing spectra at rotor cycles 20, 40, 60, 80, and 90 of EqtlI_{1–32}. Although there was only one ¹⁹F label in the EqtlI_{1–32}, spinning sidebands were observed at 8 kHz as ¹⁹F has a relatively large chemical shift anisotropy (CSA) (~40 kHz). Figure 2(g) shows a dephasing curve of $\Delta S/S_0$ obtained with a best fit to the data. The [¹⁹F-para]Phe16–[1-¹³C]Leu19 distance obtained in lyophilized powder is in agreement with that measured in the crystal and solution structures of EqtlI^{7,8} (Table I). The distance for the pair [¹⁹F-para]Phe16–[¹⁵N]Leu23 in the EqtlI structures (14.6 Å in the crystal and 12.9 Å in solution) is beyond the REDOR distance measurement limit.

Intermolecular distance measurements in DMPC membrane bilayers

Labeled EqtlI_{1–32} was added to DMPC bilayers at a molar ratio of 1 peptide to 30 lipids, with a hydration level of 50% w/v. To confirm that the sample was in a bilayer lamellar phase, a ³¹P NMR spectrum with proton decoupling was obtained at 25°C. The spectrum (not shown) had CSA of –45 ppm, including a small isotropic component which was due to EqtlI_{1–32} disruption of the membrane bilayers. A similar result was reported previously for full-length EqtlI.³⁷

REDOR experiments were carried out at –60°C to eliminate the fast axial rotation of the peptide. Figure 3(a–c) shows ¹⁹F–¹³C REDOR spectra of [¹⁹F-para]Phe16–[1-¹³C]Leu19 EqtlI_{1–32} associated with DMPC bilayers. The signal decayed more rapidly than in the lyophilized form [Fig. 3(d)], implying that there was a significant decrease in the internuclear distance, which was determined to be 10.8 Å in the lyophilized form and 5.0 Å in DMPC bilayers (Table I). The distances between [¹⁹F-para]Phe16 and [¹⁵N]Leu23 EqtlI are 14.6 and 12.9 Å in the X-ray and solution structures and are, therefore, beyond measurement by REDOR. The differences between distances in the crystal and solution structures may be due to the sensitivity of the conformation of the N-terminus

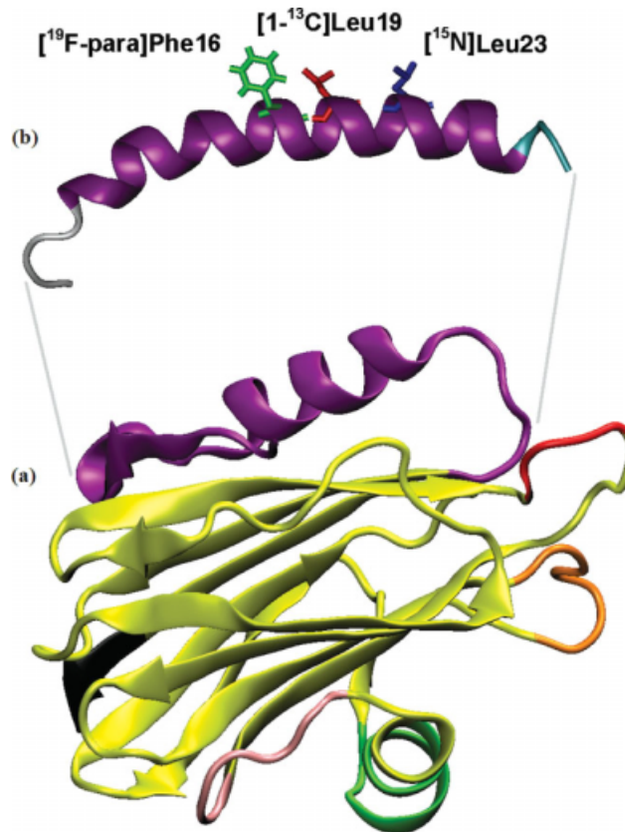


Figure 1

(a) Crystal structure of EqtlI (PDB access code 1KD6) with two short helices ($\alpha 1$, residues 15–26, and $\alpha 2$, residues 129–134) packed against opposite faces of a β -sandwich type structure formed by a pair of five-stranded β -sheets, Arg79–81 are labeled in red, Tyr 108–113 in orange; Lys123–Ala128 in pink; Glu134–Pro142 in green; Lys178–Ala179 in black, and (b) N-terminal domain EqtlI_{1–32} in DPC micelles (BMRB accession number 6808) with isotope labeling scheme as indicated. [Color figure can be viewed in the online issue, which is available at www.interscience.wiley.com.]

region of EqtlI to pH and temperature. The pH was 4.6 and the temperature 100 K for the X-ray structure and 3.9 and 303 K for solution NMR.⁸ When in the DMPC membrane environment, a notable change in the [¹⁹F-para]Phe16–[¹⁵N]Leu23 distance from ~14.6 Å to 8.2 Å was observed by REDOR. A similar reduction in the [¹⁹F-para]Phe16–[1-¹³C]Leu19 distance (from ~10.9 Å to 5.0 Å) was also seen, indicating that the region from Phe16–Leu23 underwent a conformational change when the peptide interacts with DMPC bilayer membranes. This is possibly due to changes in the orientation of the Phe16 side chain or a change in secondary structure around this region. We have employed MD simulations to obtain further insight into these changes, as described below.

EqtlI_{1–32}–DMPC headgroup contacts

Previous findings suggested that at least two regions of EqtlI became embedded in lipid membranes, the

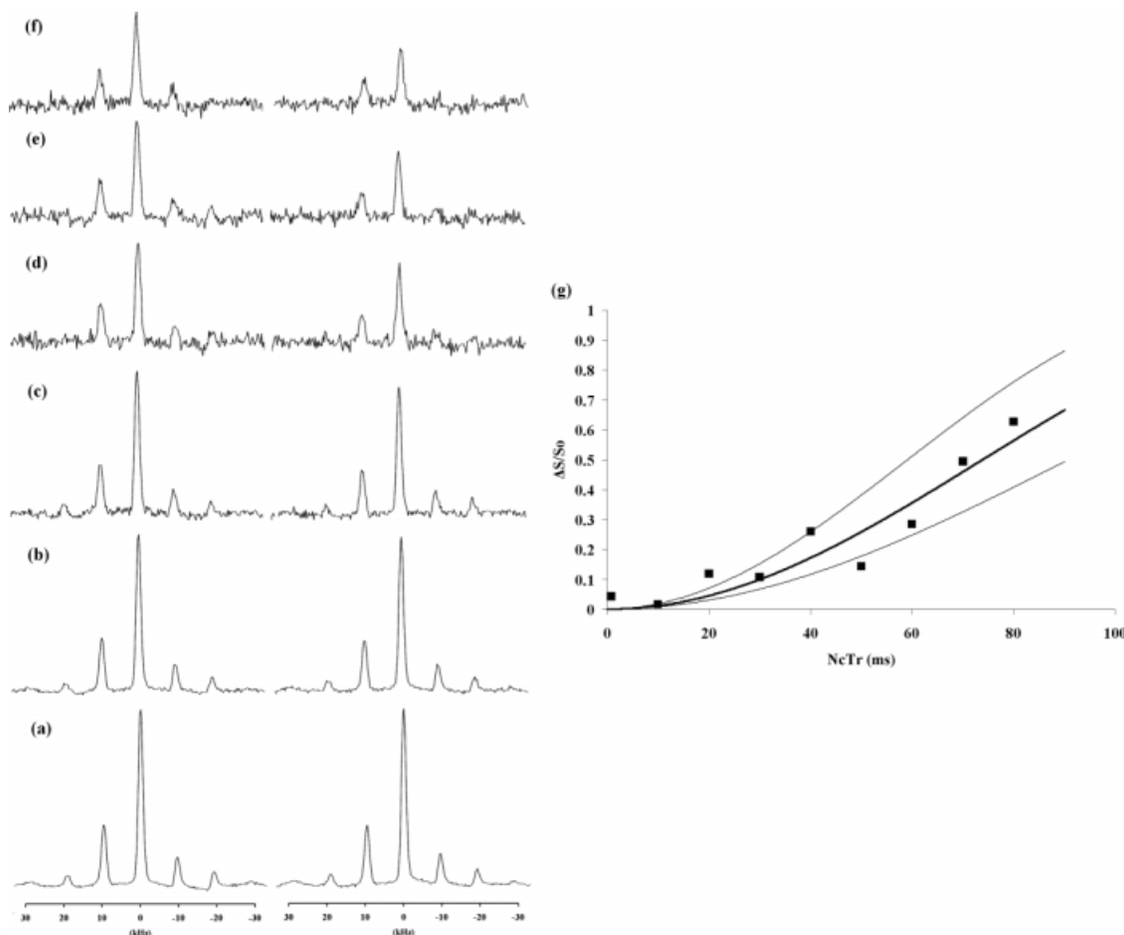


Figure 2

470.54 MHz ^{19}F - ^{13}C REDOR full echo (S_0 ; left) and difference (S_D) spectra of 5 mg [^{19}F -para]Phe16- ^{13}C Leu19 EqtII $_{1-32}$ as lyophilized powder after: (a) 20, (b) 30, (c) 40, (d) 60, (e) 80, and (f) 90 rotor cycles of dipolar evolution with MAS at 10 kHz. Each spectrum was the result of the accumulation of 4096 scans; and (g) ^{19}F - ^{13}C REDOR dephasing ($\Delta S/S_0$) data are represented by the squares. The solid line shows the calculated dephasing curve for an isolated ^{19}F - ^{13}C pair with dipolar couplings of 21.4 Hz.

N-terminal region (residues 10–28) and the surface aromatic cluster including Trp112 and 116.^{15,18,38} To gain insight into the relative positions of the Phe side chains with respect to the phospholipid headgroups, ^{19}F - ^{31}P REDOR experiments were performed on [^{19}F -para]Phe16 EqtII $_{1-32}$ - ^{31}P]DMPC. Rapid dephasing of the ^{19}F peak resulted from close proximity ^{19}F - ^{31}P contacts between the Phe side chain and the phospholipid headgroups. This provides direct evidence for the proximity of the Phe16 side chain of EqtII $_{1-32}$ to the bilayer phospholipid headgroups.

The precise conformation and orientation of the peptide relative to the bilayer, either on or penetrating into the bilayer, is currently unclear. Therefore, the orientation relative to the membrane normal was investigated using static ^{19}F NMR spectra with proton decoupling, for which an isotropic peak was observed with a linewidth of 500 Hz, whereas ^{19}F MAS spectra of peptide as

a lyophilized powder exhibited a linewidth of ~ 1500 Hz (Fig. 4). These linewidths indicate that axial rotation of the Phe16 sidechain is rapid and suggests that the benzene ring of Phe16 is on the water side of the water-bilayer interface, rather than buried inside the bilayer, in which case a slower axial rotation would have been observed.

Internuclear distance measurements in DMPC:SM bilayers

SM in the membrane creates conditions for irreversible insertion and pore formation by EqtII.^{5,6} More recent studies have shown that the coexistence of liquid ordered and disordered lipid phases favor membrane insertion of EqtII.^{39–42} Here, EqtII $_{1-32}$ was incorporated into a mixture of DMPC:SM (1:1 molar ratio), in which maximum leakage by EqtII has been demonstrated.³⁹ An increase in

Table I

Intermolecular Distances (± 0.5 Å for SS NMR Measurements³⁵) Involving Phe16 of EqtlI from X-Ray Crystallography,⁷ Solution,⁸ and Solid-State NMR Measurements and MD Simulations

Sample	XRD EqtlI	Solution NMR EqtlI _{1–32}	SS NMR lyophilized powder	SS NMR in DMPC	SS NMR in PC:SM	MD ⁺ EqtlI _{1–32}	MD ⁺ EqtlI in DMPC
[¹⁹ F-para]Phe16-[1- ¹³ C]Leu19	10.8	10.8	11.0	5.0	6.4	8 ± 2 (e-PC) 8 ± 2 (w-PC) 10 ± 2 (PC:SM)	<u>7.0 ± 2</u>
[¹⁹ F-para]Phe16-[¹⁵ N]Leu23	14.6	12.9	>	8.2	>	12 ± 2 (e-PC) 10 ± 2 (w-PC) 14 ± 2 (PC:SM)	<u>10 ± 2</u>
[¹⁹ F-para]Phe16-[³¹ P]lipid	N/A	N/A	N/A	7.0	6.0	9 ± 3 (e-PC) 15 ± 3 (w-PC) 11 ± 3 (PC:SM)	<u>9 ± 2</u>

All units in Å.

N/A, not applicable due to lack of lipid in XRD crystal, solution NMR in aqueous solution and solid-state NMR in lyophilized powder.

>, out of range, maximum range for ¹⁹F-¹⁵N distance measurement is ~8 Å³⁶.

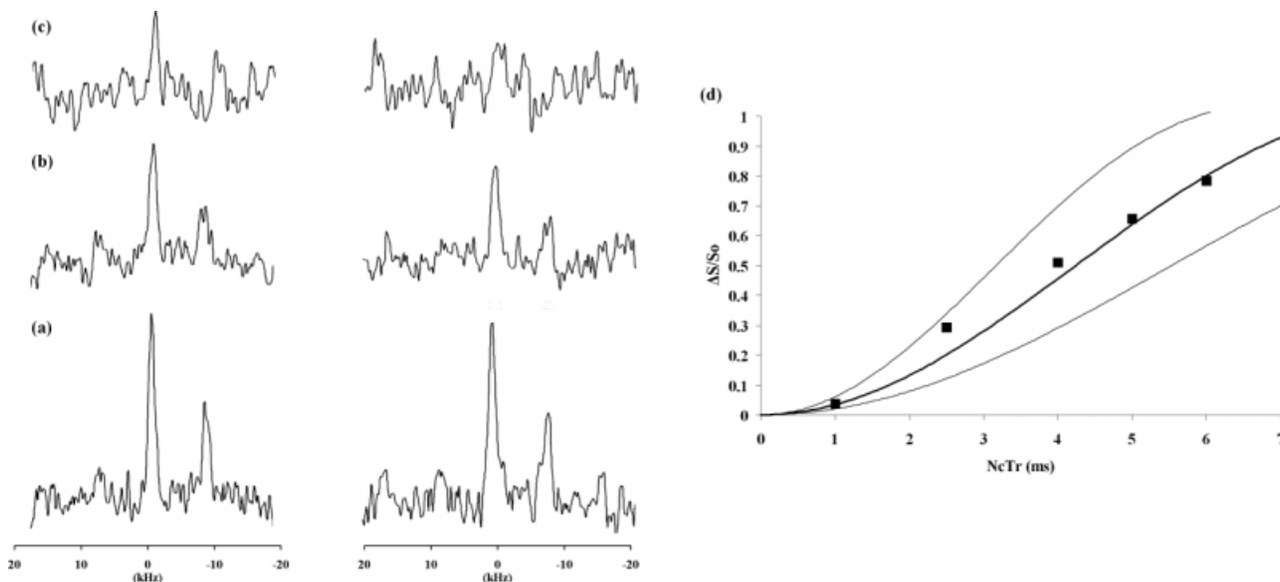
+, averaged over final 20 ns of simulation trajectories. “e-PC” and “w-PC” indicate “pep-exp-DMPC” and “pep-wat-DMPC” systems, respectively, whereas “PC:SM” indicate “pep-exp-mixed” system (see text for details). Uncertainty given as ± standard deviation. MD data which fit within current SS NMR experimental uncertainty are underlined.

the [¹⁹F-para]Phe16-[1-¹³C]Leu19 distance from 5.0 to 6.4 Å was seen when the peptide was incorporated into DMPC/SM bilayers compared with DMPC bilayers (Table I). An increase in the [¹⁹F-para]Phe16-[¹⁵N]Leu23 distance to a more extended structure (out of the REDOR measurable range) was also observed when the peptide was incorporated into DMPC/SM compared with that in DMPC, that is, 8.2 Å (Table I). This suggests that the presence of SM in this system affects the secondary structure of the peptide. Distances within the segments 16–23

and 19–23 are consistent with helical structure, but an induced kink brings these distances closer when the peptide interacts with a membrane bilayer. This is discussed in the section below (“Molecular dynamics simulations”).

EqtlI_{1–32}-phospholipid headgroup contacts

In DMPC:SM mixed bilayers, two types of phospholipid headgroups are present. A ³¹P proton-decoupled MAS spectrum was acquired in an attempt to resolve

**Figure 3**

470.54 MHz ¹⁹F-¹³C REDOR full echo (S_0 ; left) and difference (S_D) spectra of 1 mg [¹⁹F-para]Phe16-[1-¹³C]Leu19 EqtlI_{1–32} incorporated in DMPC at lipid: peptide molar ratio of 30:1 after: (a) 4, (b) 12, and (c) 32 rotor cycles of dipolar evolution with MAS of 8 kHz and temperature -60°C ; and (d) ¹⁹F-¹³C REDOR dephasing ($\Delta S/S_0$) data represented by the squares. The solid line shows the calculated dephasing curve for an isolated ¹⁹F-¹³C pair with dipolar couplings of 227.5 Hz.

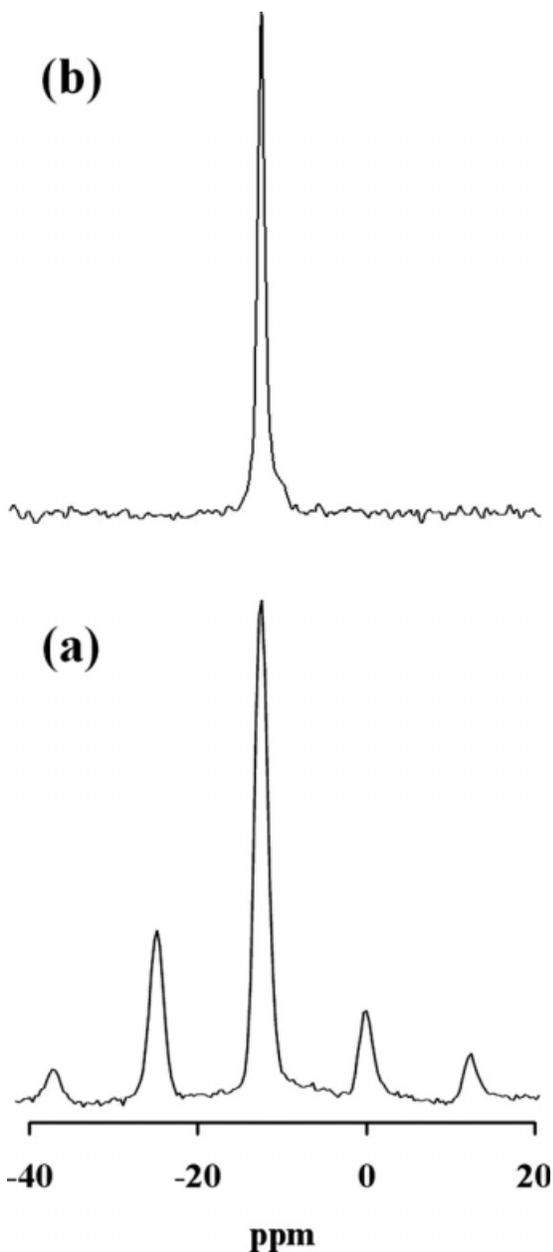


Figure 4

^{19}F spectra of $[^{19}\text{F}\text{-para}]\text{Phe16 EqII}_{1-32}$ at 25°C as: (a) a lyophilized powder with MAS of 10 kHz, and (b) without spinning when incorporated into fully hydrated DMPC.

these two components, but at low temperature (-60°C), the linewidth of the ^{31}P peak increased and only one phosphorus component was seen. A $^{19}\text{F}\text{-}^{31}\text{P}$ REDOR experiment was performed on $[^{19}\text{F}\text{-para}]\text{Phe16 EqII}_{1-32}\text{-}[^{31}\text{P}]\text{DMPC:SM}$, in which a reduction in distance between the peptide and phospholipid headgroups from 7.0 Å in DMPC to 6.0 Å in DMPC:SM was observed (Table I). Thus, the presence of SM brings the peptide and lipid closer together than in pure DMPC. Static ^{19}F

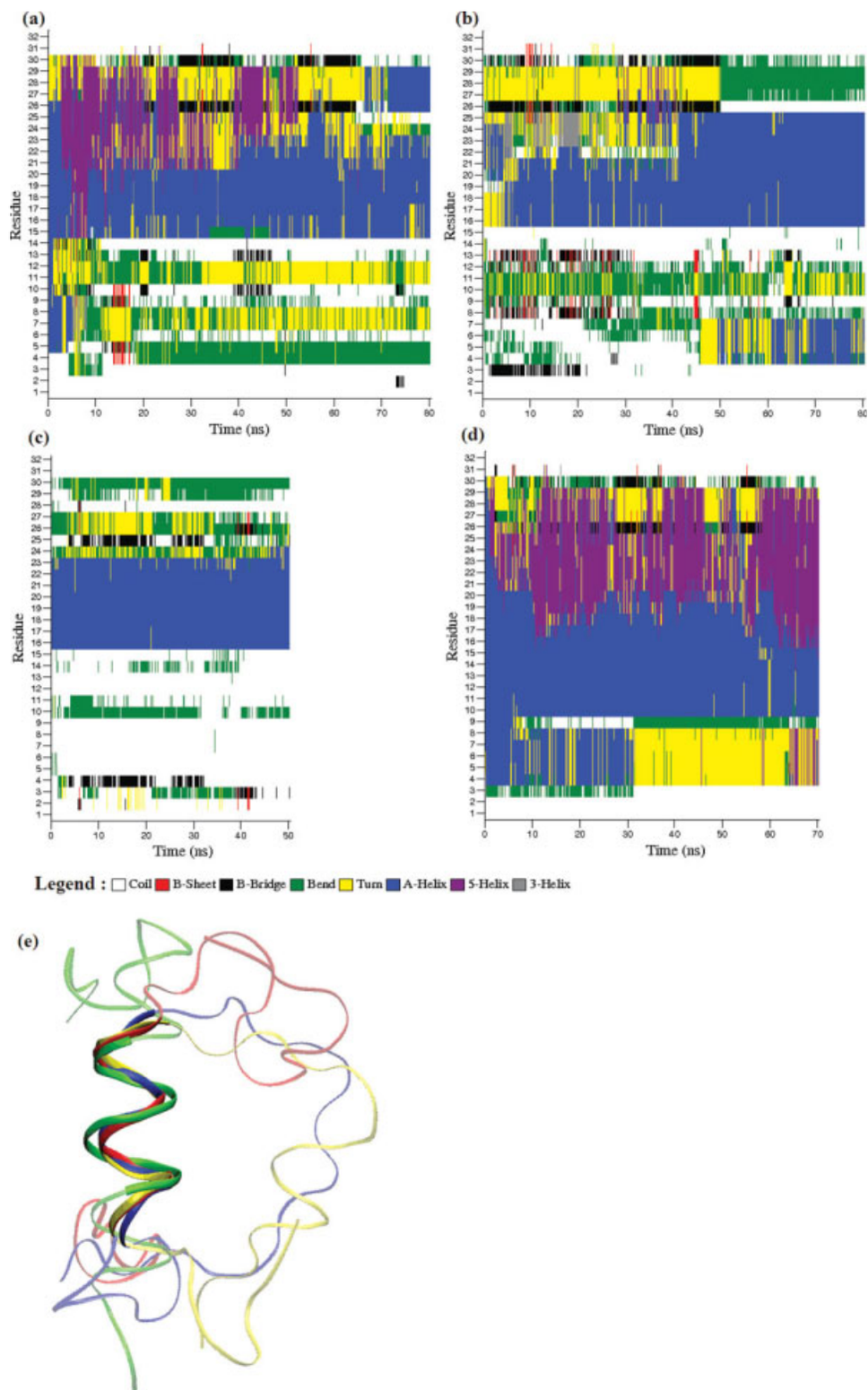
NMR spectra of $[^{19}\text{F}\text{-para}]\text{Phe16 EqII}_{1-32}$ incorporated into DMPC:SM at 25°C were similar to those in DMPC (data not shown), indicating that the Phe16 side chain is still located at the water interface.

Molecular dynamics simulations

EqII₁₋₃₂ and EqII structures and distances: comparisons with experiments

We have performed computational simulations on EqII_{1-32} and EqII in the presence of lipid bilayers. For both the peptide and protein, we have also obtained structures by MD simulation in pure water. Simulation of EqII_{1-32} in water up to 10 ns revealed that the peptide in water adopts flexible loop and turn structures, in contrast to the helical structure seen in DPC micelles.⁹ Inspection of the initial peptide conformation [Fig. 6(d)] for the **pep-wat-DMPC** simulation and the DSSP plot for this system at 0 ns [Fig. 5(b)] show that α -helicity is retained between residues 21–24, suggesting $\sim 83\%$ loss of the α -helical content compared with the DPC structure. This is consistent with previous NMR studies, which indicated that EqII_{1-32} in aqueous solution does not adopt an ordered conformation.⁹ For full-length EqII , there are no significant secondary structural changes during 10 ns simulation in water, although a higher degree of flexibility is observed in the N-terminal region (residues 1–15) relative to the rest of the protein (data not shown). This is consistent with previous NMR studies of EqII ,⁸ which indicate the relative flexibility of the N-terminus.

For EqII_{1-32} and EqII simulated in the presence of bilayers, average interatomic distances for Phe16-Leu19, Phe16-Leu23, and Phe16-P (calculated over the final 20 ns of the simulated trajectories, during which the peptide or protein maintains close contact with the bilayer surface throughout) and the associated standard deviations are given alongside experimentally determined values in Table I. Of the simulation data, the best fit to the SS NMR data are those of EqII on DMPC (**prot-DMPC**), for which the lower limit of both Phe16-Leu19 and Phe16-Leu23 intramolecular interatomic distances fall within the experimentally determined range. All simulations (peptide and protein) produced Phe16-Leu19 distances that are statistically lower than those of the crystal and solution structures and lyophilized-power SS NMR experiments, in qualitative agreement with our current SS NMR experiments, which indicate a lowering of these values as a result of peptide interaction with DMPC. The Phe16-Leu23 distance for **prot-DMPC** and **pep-wat-DMPC** simulations are statistically lower than that obtained from the crystal and solution structures, also in qualitative agreement with the trend obtained from SS NMR. The **pep-exp-DMPC** simulation obtains a value lower than that of XRD but only marginally lower

**Figure 5**

DSSP representations of secondary structure with respect to MD simulation time for: (a) pep-exp-DMPC, (b) pep-wat-DMPC, (c) prot-DMPC (residues 1–32), and (d) pep-exp-mixed. Secondary structures are differentiated by color, as defined in the figure legend. (e) Ribbon structures of residues 1–32 of the protein (blue), pep-exp-DMPC (red), pep-wat-DMPC (yellow), and pep-exp-mixed (green) obtained from the final snapshots of their respective simulations, overlaid via residues 16–24, which are shown as thicker ribbons.

than that obtained by solution NMR. Additionally, the Phe16-Leu19 distance for the peptide simulation on a mixed DMPC:SM bilayer (**pep-exp-mixed**) is higher than that obtained on a pure DMPC bilayer, with ~ 10 Å and ~ 8 Å, respectively. This is in qualitative agreement with SS NMR data, which indicate a longer Phe16-Leu19 distance (by 1.4 Å) in DMPC:SM compared with DMPC.

Overall, these distance data indicate that for the DMPC bilayer, the structure of EqtII_{1–32} resembles the structure of residues 1–32 in the full-length protein, and thus lends some support to the N-terminal peptide as a valid model of the N-terminus of the full-length protein in membrane interaction studies. Furthermore, the simulations are consistent with our SS NMR results in so far as the Phe16-Leu19 and Phe16-Leu23 distances are reduced compared with the crystal and solution structures. However, quantitative agreement between constraint-free simulation and SS NMR for EqtII_{1–32} was not obtained, possibly because of the disparity in the time-scales between simulation (80 ns) and experiment (orders of magnitude longer). Nevertheless, the RMSD of the peptide backbone of all simulations plateaued at ~ 20 ns (data not shown), indicating that the simulations had reached equilibrium (i.e., were in local energy minima); furthermore, several distinct trends in peptide secondary structure and the mechanisms of initial bilayer interactions and subsequent binding were manifested in all of the simulations performed, discussed in subsequent sections. To obtain insights into the structure of the peptide while taking into account SS NMR-acquired distances, a distance-constrained simulation of EqtII_{1–32} was performed in which Phe16(H_ε) was distance-constrained at 5.0 and 8.2 Å to Leu19(C) and Leu23(N), respectively. Structural evolution for all systems simulated are discussed below.

Secondary structures

The secondary structural evolutions with respect to MD simulation time for EqtII_{1–32} and EqtII are represented by DSSP plots (Fig. 5). For both **pep-exp-DMPC** and **pep-wat-DMPC** systems, the secondary structure feature that emerges is persistent helicity in the C-terminal half of the peptide (residues 16–24), with largely random-coil structures in the N-terminal half (residues 1 to 15). The emergence of similar secondary structural motifs after ~ 30 ns simulations for two peptide systems starting from different initial structures lends support to the notion that although the peptide is mainly unstructured in water (as observed in 10 ns simulation), the binding of the peptide to a DMPC bilayer produces structural bias toward helical forms in the C-terminal segment. This is consistent with experimental results for EqtII_{1–32} in DPC micelles.⁹ Inspection of the DSSP plot for residues 1–32 of **prot-DMPC** [Fig. 5(c)] indicates stability of secondary structural features throughout the 50 ns

simulation. In agreement with the EqtII_{1–32} simulations described earlier, residues 16–24 remain α -helical, whereas residues 1–15 are unstructured. Agreement between the secondary structure for EqtII_{1–32} and the N-terminal segment of EqtII provides further support for the validity of using peptide fragments as experimental models of full-length actinoporin proteins. For the distance-constrained simulation **pep-cons-DMPC**, for which distance constraints were applied to maintain interatomic distances of 5 and 8.2 Å of Phe16(H_ε) to Leu19(C) and Leu23(N), respectively, we find that the secondary structure does not change appreciably from that of the 50 ns “snapshot” of the **pep-exp-DMPC** structure, from which the initial **pep-cons-DMPC** structure was derived. This structure retains an α -helical segment between residues 16–23, a distorted helix between residues 24–29, and random coil between residues 1–15 [Fig. 6(l)], throughout the 15 ns simulation. The stability of the helical segments of the peptide in the presence of experimentally derived distance constraints lends further support to our (unbiased, constraint-free) MD simulations which indicate that the SS NMR-measured region of the peptide consists mainly of helix segments. It also indicates that the experimentally measured distances are consistent with a stable peptide structure.

Compared with the above simulations, different secondary structural behavior is observed for the 1:1 mixed DMPC:SM simulation, **pep-exp-mixed**, in which the initial structure is mainly α -helical (similar to that of **pep-exp-DMPC**). It exhibits greater structural stability relative to the peptide bound to DMPC as evidenced by the persistence of helical elements throughout the peptide between residues 10 and 29 [Fig. 5(d)], which is a much longer helical stretch than that exhibited by the peptide in the presence of DMPC only. There is a conformational transition at the N-terminal end (residues 1–9) to an extended β -strand after ~ 30 ns; unlike the DMPC simulations, very little random-coil structure is formed. The relative persistence (compared with DMPC alone) of helical and β -stranded secondary structure may be due to residue-dependent preference for interaction with either PC or SM, coupled with the structure of the underlying mixed bilayer “template,” which forms distinct segregated nanometre-sized “domains” (i.e., small regions largely composed of either DMPC or sphingomyelin); the membrane thus imposes order on the peptide. More detailed discussions of peptide–lipid interactions are given in the next sections. The persistence of stable helicity throughout a larger portion of the peptide, together with the rapid formation of β -stranded N-terminus, may contribute to its enhanced activity against PC:SM bilayers compared with pure PC bilayers,⁴³ as the existing helix-strand motif may render the peptide more energetically favorable to bilayer permeation.

We note that that the N-terminal peptide structural characteristics are qualitatively consistent across all four

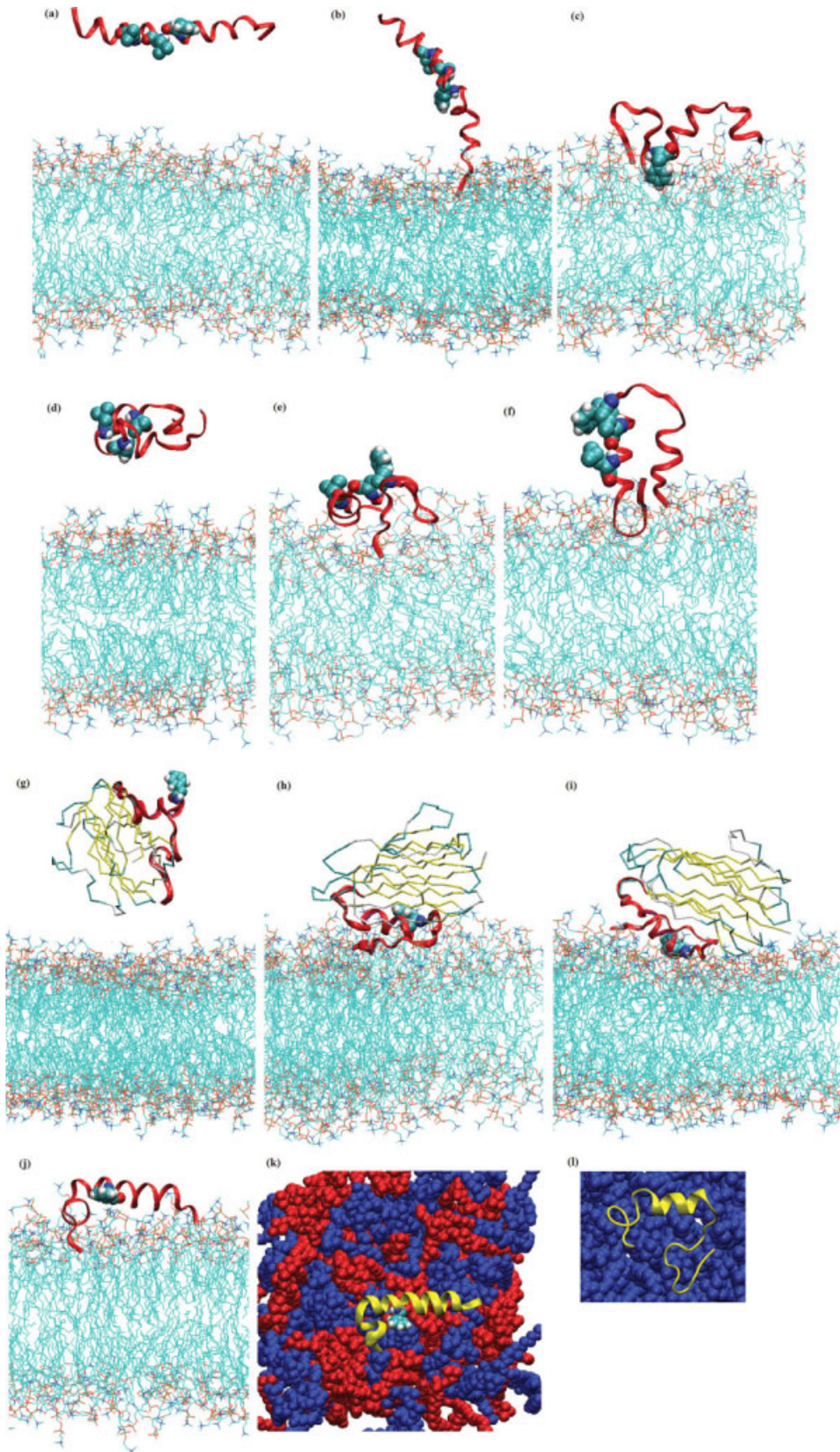


Figure 6

Representative structures of EqtII₁₋₃₂ and EqtII interactions with DMPC bilayers obtained as “snapshots” from MD simulation trajectories. Peptide backbone represented as red ribbons, Phe16, Leu19 and Leu23 represented in van der Waals’ spheres, and lipids represented as thin lines. Atoms are colored as follows: cyan carbon, brown phosphorus, blue nitrogen, and red oxygen. (a) pep-exp-DMPC at 0 ns, (b) 1 ns, (c) 80 ns; (d) pep-wat-DMPC at 0 ns, (e) 20 ns, (f) 80 ns; (g) prot-DMPC at 0 ns, (h) 20 ns, (i) 50 ns; and (j) pep-exp-mixed at 70 ns viewed along axis perpendicular and (k) parallel to bilayer normal, with DMPC lipids represented as blue spheres, SM represented as red spheres, and the peptide backbone colored in yellow; (l) pep-cons-DMPC at 15 ns, viewed along bilayer normal, with DMPC lipids as blue spheres. [Color figure can be viewed in the online issue, which is available at www.interscience.wiley.com.]

of our simulations, with α -helical structure obtained in all four simulations between residues 16 and 24, and extended or random-coil structures at residues 1–15 and 24–32. The qualitative structural similarities are illustrated in Figure 5(e), which show the final snapshots of the peptide from the respective trajectories, overlaid by structural alignment of residues 16–24.

Interactions of EqtII_{1–32} and EqtII with bilayers

Average interatomic distances for Phe16-P and the associated standard deviations for EqtII_{1–32} and EqtII simulated with bilayers are given alongside experimentally determined values in Table I, calculated over the final 20 ns of the simulated trajectories for which the peptide or protein maintains close contact with the bilayer surface throughout. For the DMPC bilayer, the best fits to the SS NMR data are **pep-exp-DMPC** and **prot-DMPC**, consistent with visual inspection of these trajectories which indicate close contact between the Phe16 sidechain and the lipid surface [Fig. 6(c,i) respectively]. Likewise, inspection of the **pep-exp-mixed** trajectory indicate interactions between Phe16 and the bilayer [Fig. 6(j)], although specific close contacts between Phe16(H_z) and phosphorus are scarce. However, **pep-wat-DMPC** exhibited few close contacts between Phe16 and the bilayer surface; visual inspection confirms that the sidechain does not make close contact with the lipid headgroups throughout the 80 ns trajectory [Fig. 6(e,f)].

The total number of close contacts (defined as interatomic contacts below 0.35 nm) between the peptide/protein and the lipid bilayer atoms as a function of simulation time is given in Figure 7. All of the EqtII_{1–32} simulations predict rapid association of the peptide with the bilayer surface, with **pep-exp-DMPC**, **pep-wat-DMPC**, and **pep-exp-mixed** forming peptide–bilayer contacts within 1 ns of simulations (Fig. 6). The contact-number plots reach a fluctuating plateau after ~40 ns, indicating saturation of peptide interactions with the bilayer surface (i.e., no further surface contacts are possible without significant peptide conformational changes). Visual inspection of the simulation trajectories reveals further details of the overall mechanism of interaction between EqtII_{1–32} and EqtII with lipid bilayers. Figure 6 consists of “snapshots” of the simulation trajectories which serve to illustrate the overall dynamical behavior of the peptides and protein with respect to the membrane for the simulated systems. For the **pep-exp-DMPC** system, the initial contact between the peptide and bilayer is made via the N-terminus [Fig. 6(b)] and occurs within 1 ns. The helical segment of the peptide (residues 15–23; see above discussion on Secondary Structures) and the C-terminus subsequently form contacts with the lipid surface, with the helical segment lying at an incline of ~30° to the membrane normal [Fig. 6(c)]. A similar mechanism of binding is obtained for **pep-wat-DMPC**, which also interacts

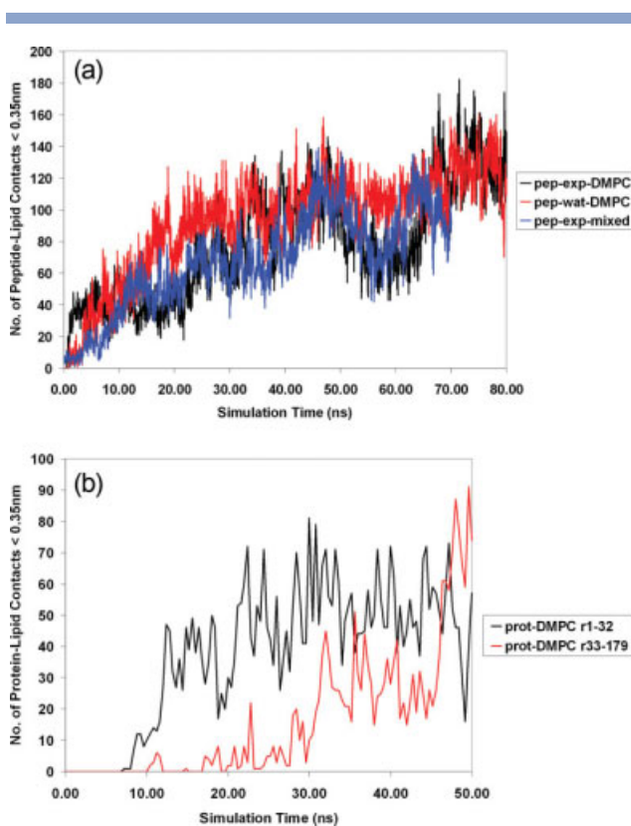


Figure 7

Total numbers of atomic contacts (<0.35 nm) between peptide and lipids with respect to MD simulation time for (a) pep-exp (black line), pep-wat (blue line) and pep-exp-mixed (red line) simulation systems, and (b) full-length EqtII, decomposed into contributions from residues 1 to 32 (black line) and 33 to 179 (red line). [Color figure can be viewed in the online issue, which is available at www.interscience.wiley.com.]

initially with, and inserts into the bilayer via the N-terminus, within 1 ns. The C-terminus also binds to and inserts into the membrane surface at ~20 ns [Fig. 6(e)]. Interestingly, the helical segment of the peptide between residues 16 and 25, which adopts an orientation approximately parallel to the membrane plane upto ~60 ns, reorients after this point to adopt an “upright” orientation, with the helical axis roughly perpendicular to the membrane plane until the end of the trajectory at 80 ns [Fig. 6(f)]. Such a conformation may be suggestive of a “preinsertion” state of the peptide, in which the (potentially) pore-spanning helical segments adopt an orientation (tilt angle) similar to that of the fully inserted peptide, in preparation for membrane permeabilization.

Examination of the total number of contacts between each residue of the peptide with the phosphate headgroup and lipid carbon tails (averaged over the final 20 ns of trajectory) provides further details of the peptide’s binding mechanism to the bilayer. Such “contact profile” bar graphs are shown in Figure 8. Figure 8(a) shows that significant contacts at both the N- and C-termini with lipid phosphate headgroups are made for both **pep-exp**

and **pep-wat-DMPC**, with the C-termini contacts maintained by the basic Lys30, Arg31, and Lys32. Basic sidechains are known to anchor membrane proteins to lipid bilayers via snorkeling interactions.⁴⁴ A number of additional contacts are formed for **pep-exp-DMPC**, at Ser13, Ser15, Phe16, and Lys20. Figure 8(b) shows the contact

profiles for the peptides and the lipid carbon-tail atoms. There are far less numbers of atomic contacts, as none of the simulated peptides (and protein) penetrated the bilayer to any significant extent. However, in addition to the N- and C-termini, Gly11, Ala12, and Phe16 maintain contacts with the upper regions of the hydrophobic region of the bilayer.

We note that the relatively large fluctuations of peptide–lipid contact numbers for the **pep-exp-DMPC** simulation may be indicative of a relatively lower stability of the peptide–lipid binding orientation for this system compared with that of **pep-wat-DMPC**. The latter system exhibits relatively minor fluctuations in Figure 7(a) and is suggestive of “tighter” binding to the membrane surface. However, a significant result is obtained when we examine the DMPC trajectories to explain the apparent differences in binding orientation stability between the two simulations. Specifically, we note that there are greater numbers of contacts between Ser1, Asn28, Lys30, and Arg31 and the bilayer surface for the lower fluctuation, “more stable” **pep-wat-DMPC** simulation [Fig. 8(a)]. Both Asn28 and Arg31 also penetrate deeply into the bilayer in **pep-wat-DMPC**, making contacts with hydrophobic lipid chains [Fig. 8(b)]. The “less stable” **pep-exp-DMPC** simulation, in contrast, exhibits no significant sidechain insertions into the hydrophobic region of the bilayer at the C-terminus [Fig. 8(b)]. This suggests that close contact between the C-terminal residues—in particular, Asn28 and Arg31—and the bilayer enhances the strength of interaction between the peptide and the membrane, effectively anchoring the peptide to the membrane surface and resulting in a more stable binding mode (as in **pep-wat-DMPC**), while lack of these interactions results in greater orientational mobility of the peptide on the bilayer surface, as manifested in the **pep-exp-DMPC** simulation. This is consistent with the known preferential location of Arg at the lipid–water interface region in membrane proteins, in which the chemical structure of its sidechain enables it to form “snorkeling” interactions (e.g., Ref. 44). Thus, despite the relatively large fluctuation in the peptide–lipid contact numbers time series for **pep-exp-DMPC**, the combined results of the **pep-exp** and **pep-wat-DMPC** simulations serve to highlight the potentially

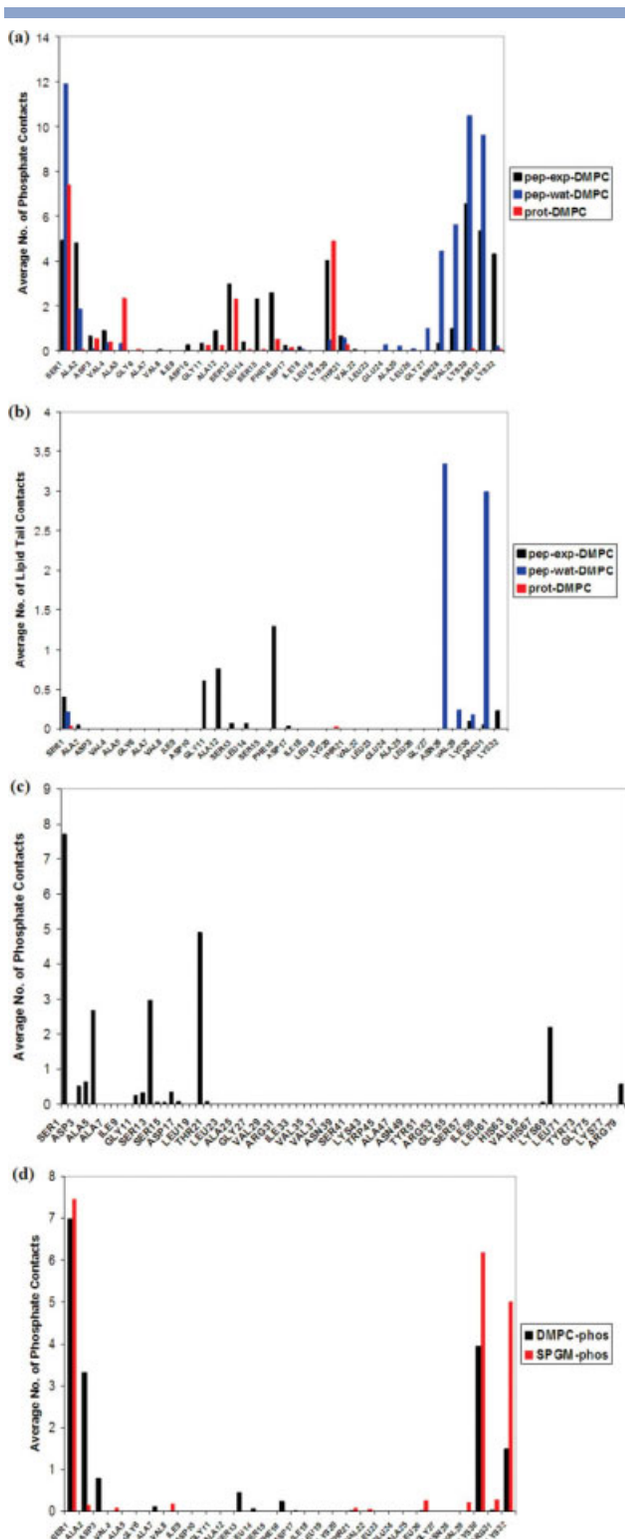


Figure 8

Numbers of atomic contacts (<0.35 nm) between each residue of EqII_{1–32} or EqII and lipids, averaged over the final 20 ns of simulation trajectories. Residue-specific lipid contact profiles for (a) lipid phosphate group atoms and peptide for **pep-exp** (black bars), **pep-wat** (blue bars), and residues 1–32 of full-length protein (red bars); (b) lipid carbon-tail atoms and peptide for **pep-exp** (black bars), **pep-wat** (blue bars), and residues 1–32 of full-length protein (red bars); (c) lipid phosphate group atoms and residues 1–79 of EqII (residues 80–179 exhibited negligible contacts); and (d) the EqII_{1–32} peptide residue-specific contacts with lipid phosphate group atoms from DMPC (black bars) and SM (red bars). [Color figure can be viewed in the online issue, which is available at www.interscience.wiley.com.]

important roles of charged sidechains such as Lys and Arg in imparting stability to the binding orientation of the peptide to DMPC membrane surfaces. A similar argument may be applied to **pep-exp-mixed** (mixed composition DMPC:SM bilayer), whose peptide–lipid contact plot also exhibits large fluctuations between 40 and 70 ns relative to **pep-wat-DMPC**, but somewhat lower fluctuations compared with **pep-exp-DMPC**. **Pep-exp-mixed** exhibits no sidechain penetration into the hydrophobic regions of the bilayer, unlike **pep-wat-DMPC**; hence, the former’s lower binding stability and greater peptide–lipid contact number fluctuation. However, **pep-exp-mixed** exhibits greater numbers of contacts between Lys30/32 and headgroup phosphates compared with **pep-exp-DMPC**, which may result in marginally greater binding stability for the former. Thus, the binding stability of the peptide on the mixed bilayer is intermediate between that of the two DMPC simulation systems.

Protein–lipid contact plots for the EqtII protein are given in Figure 7(b), decomposed into contributions from residues 1–32 of the protein and residues 33–179 (i.e., the “remainder”) of the protein. The plot indicates rapid association of the N-terminal, r_{1-32} fragment of EqtII with the bilayer surface, with contacts formed with the membrane within ~ 5 ns (we note that the slower time to initial contact for the protein compared with the peptide is likely due to the slower diffusional motion of the former). The remainder of the protein (r_{33-179}) forms few, nonpersistent contacts between 20 and 30 ns, with more significant numbers of contacts formed only after 30 ns of simulations. This indicates that the r_{1-32} segment of the protein is strongly attracted to the bilayer surface, more so than other segments of EqtII. Indeed, visual inspection of the **prot-DMPC** trajectory [snapshots shown in Fig. 6(g–i)] reveals that, while the initial orientation of the protein is placed such that the long axis of the r_{1-32} segment lies nearly parallel to the membrane normal [Fig. 6(g)], the protein undergoes rapid rotational and translational motion which results in the placement of r_{1-32} directly in contact with the membrane, with the long axis aligned approximately perpendicular to the membrane normal [Fig. 6(h)]. Furthermore, between 40 and 50 ns, the helical segment at residues 14–23 is tilted to the bilayer normal at $\sim 70^\circ$ [Fig. 6(i)]. This coincides with the formation of new contacts between other regions (residues 69 and 79; see below) of the protein with the bilayer. Thus, **pep-exp-DMPC**, **pep-wat-DMPC**, and **prot-DMPC** all indicate that the short helical fragment around this region is oriented at an incline to the membrane surface.

A contact profile bar graph of the r_{1-32} segment of EqtII superimposed on the EqtII $_{1-32}$ simulations [Fig. 8(a), red bars] indicates some similarities in the profile of residue-dependent contact with the phosphate headgroup atoms of DMPC; similar to both **pep-exp-DMPC** and **pep-wat-DMPC**, the protein makes significant head-

group contacts at the N-terminus. There are also significant contacts between Lys20 and phosphates. However, unlike the peptides, there are no significant contacts between the region around Lys32 (i.e., the C-terminus of EqtII $_{1-32}$) and phosphates, since this segment is bound to the remainder of the full-length protein. Since the protein is known to exhibit greater biological activity than the N-terminal fragment peptide,^{2,3,9} significant binding between the (Lys32) C-terminus and the lipid headgroup region may be one factor which limits EqtII $_{1-32}$ membrane activity. We note that there are no significant contacts between any residue of the full-length protein and the hydrophobic region of DMPC [Fig. 8(b)]. This is in agreement with fluorescent studies, which suggest that EqtII does not insert deeply in the bilayer.¹⁹ An extended contact profile graph for EqtII over residues 1 to 80 is shown in Figure 8(c) (residues 81 to 179 exhibited no contacts with the bilayer during this simulation; data not shown), indicating significant bilayer interactions at the N-terminal (r_{1-32}) segment and isolated interactions via Lys69 and Arg79. The latter two interactions are formed later in the simulation (40–50 ns).

The plot of the total number of contacts between the EqtII $_{1-32}$ peptide and a mixed 1:1 DMPC:SM bilayer [Fig. 7(a)] shows rapid association between the peptide and the bilayer, in a similar manner and timescale as the peptide-DMPC simulations described earlier. Representative structures of EqtII $_{1-32}$ peptide after binding to a 1:1 mixed DMPC:SM bilayer are shown in Figure 6(j,k). Similar to the DMPC simulations, initial interactions are formed between the N-terminus of the peptide (within 1 ns) with the lipid bilayer, followed by the C-terminus (~ 20 ns). Figure 6(k) illustrates the system at 70 ns, viewed along the bilayer normal. DMPC molecules are shown as blue spheres, while sphingomyelin molecules are shown as red spheres. The graphic illustrates a degree of phase separation between the two lipid types, with nanometre-scale domains (“nanodomains”) composed primarily of DMPC or SM distributed throughout the simulated bilayer. The C-terminal part of the peptide exhibits preferential interaction with SM. This is more clearly manifested in a contact profile plot [Fig. 8(d)], which indicates that while the N-terminal side of the peptide interacts somewhat more favorably with DMPC phosphates (with contacts made via Ser1, Ala2, and Asp3, as opposed to only Ser1 for sphingomyelin), the C-terminal side (via Lys30 and Lys32) exhibits significantly greater numbers of contacts with sphingomyelin phosphates. Preferential interaction of the peptide with either DMPC or SM may contribute to its enhanced secondary structural persistence (discussed earlier) compared with that of a pure DMPC bilayer. The mixed bilayer thus imposes structural order on the peptide, due to its own intrinsic nanodomain order. The enhancement of structural order may in turn contribute to the enhanced activity of the peptide on PC:SM bilayers.⁴³

CONCLUSIONS

We have applied SS NMR and MD computational simulations to examine the structure and DMPC:SM bilayer interactions of the N-terminal peptide (residues 1–32) of the highly soluble toxin EqtII. For the peptide in solution, the distances between specifically labeled nuclei in [¹⁹F-para]Phe16-[¹³C]Leu19 and [¹⁹F-para]Phe16-[¹⁵N]Leu23 analogs of EqtII_{1–32} measured by REDOR in lyophilized powder were in agreement with published crystal and solution structures. However, in both DMPC and mixed DMPC:SM membrane environments, significant changes in the distances between the labeled amino acid pairs were observed, suggesting changes in helical content in this region in the presence of bilayers. Differences in REDOR-acquired peptide–lipid interatomic distances between pure DMPC and mixed DMPC:SM bilayers were also observed, which may be due to the greater affinity of EqtII for the latter. MD simulations of both EqtII_{1–32} and EqtII indicated that the regions around the N-terminus and the aromatic and positively charged residues are highly flexible and contribute to membrane binding as well as undergoing changes in secondary structure during interaction with bilayers. Comparison of the water-solvated peptide structure, which is largely disordered, indicates that binding to a DMPC surface induces helix formation at residues ~16–24. The simulation results also indicate strong interaction between the peptide and the mixed DMPC:SM bilayer, with the C-terminus exhibiting preferential interaction with SM. The large portion of the peptide which maintains helical structure, coupled with a tendency to form a β-stranded N-terminal tail, may be a contributing factor to the greater membrane activity of the peptide against mixed PC:SM bilayers.

The appearance of similar structural and bilayer-binding characteristics in simulations with widely varying initial conformations lends credence to the notion that these phenomena are genuine features of EqtII_{1–32}-bilayer interactions. Our MD simulation results indicate persistent interactions with the lipid bilayer of the N-terminal region, as well as the aromatic and positively charged regions. These regions are known from previous studies to play a role in membrane interactions of EqtII. Thus, the overall mechanism of EqtII peptide and protein initial interaction with the DMPC bilayer, based on the contacts with phosphates and acyl chains which appear to be shared in common, may be summarized as follows: the initial binding event, driven by electrostatic attraction, appears to be between the N-terminus and the phosphate headgroups. Thereafter, the N- and C-termini both anchor the peptide to the bilayer. Further electrostatic interactions between Lys residues (which are located at the center and C-terminal regions of the peptide) and phosphate headgroups may serve to drive the peptide toward a roughly parallel orientation. Hydrophobic

interactions may also be formed between various nonpolar residues and the upper segments of the acyl chains, such as Gly11 and Ala12, as suggested by **pep-wat-DMPC**. The latter interactions may, given a longer time-scale, disrupt the bilayer core sufficiently for full peptide and protein insertion. In summary, the results obtained from our SS NMR and MD studies of the interaction of the N-terminus of EqtII with phospholipid bilayers may be indicative of the first step in the conformational changes required for the permeation and subsequent disruption of the membrane due to the binding of the peptide.

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