The Dynamic Orientation of Membrane-Bound Peptides: Bridging Simulations and Experiments

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ABSTRACT The structural organization in a peptide/membrane supramolecular complex is best described by knowledge of the peptide orientation plus its time-dependent and spatial fluctuations. The static orientation, defined by the peptide tilt and a rotation about its molecular axis, is accessible through a number of spectroscopic methods. However, peptide dynamics, although relevant to understand the functionality of these systems, remains largely unexplored. Here, we describe the orientation and dynamics of Trp-flanked and Lys-flanked hydrophobic peptides in a lipid bilayer from molecular dynamics simulations. A novel view is revealed, where collective nontrivial distributions of time-evolving and ensemble peptide orientations closely represent the systems as studied experimentally. Such global distributions are broad and unveil the existence of orientational states, which depend on the anchoring mode of interfacial residues. We show that this dynamics modulates ²H quadrupolar splittings and introduces ambiguity in the analysis of NMR data. These findings demonstrate that structural descriptions of peptide/membrane complexes are incomplete, and in cases even imprecise, without knowledge of dynamics.

INTRODUCTION

Lipid membranes, in their physiologically relevant liquidcrystal phase, are characterized by the partial order and high fluidity of constituent amphiphilic rodlike molecules, defining a lamellar arrangement with a centered hydrophobic layer, flanked by hydrated polar regions (1). This complex organization conditions the structure, orientation, and dynamics of polypeptides there embedded, which are restrained with respect to the membrane plane while still being free to diffuse within the lipid bilayer (2).

From a practical point of view, a restricted environment reduces the number of degrees of freedom and simplifies the structure problem in protein-membrane complexes. Thus, despite the difficulties to obtain high resolution information from these systems with classical techniques (x-ray crystallography and high resolution NMR), alternative spectroscopic methods have yielded new insights into their molecular organization in the form of orientation parameters (3–8). These methods are applied to complexes of α -helical peptides with membranes, and give mainly the tilt of the peptide molecule and the rotation about its principal axis. The peptide tilt is profusely measured, as it relates directly to the mechanisms of membrane-active peptides, like those forming pores or promoting membrane fusion. Additionally, the tilt is also important to define the response of the peptide to interactions with the membrane, according to ideas illustrated by the mattress model and explained under the concept of hydrophobic mismatch (9,10). On the other hand, the peptide rotation, despite being more scarcely studied (8,11–14), is relevant to fix the position of anchoring residues with respect

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to the membrane interface and to display the helix face that can be used for possible intermolecular interactions.

Notwithstanding the importance of the above static structural descriptions, a complete understanding of the molecular organization of peptide membrane complexes necessitates knowledge about their dynamics. Recent relaxation studies by solid-state NMR experiments show transmembrane polypeptides undergoing both axial diffusion and small amplitude off-axis reorientation, in the nanosecond and microsecond, respectively, time regimes (15,16). Including the later movements in the models used to interpret raw spectroscopic data can be expected to improve the end orientation results. However, this would require a detailed understanding of the mechanisms of subjacent dynamic processes. Molecular dynamics (MD) simulations have the potential to provide such a description (17). While still limited by the quality of the underlying models and accessible time- and length-scales, MD methods are increasingly applied to biomembrane systems (18–29). Thus, pure lipid bilayers and protein-membrane complexes can now be studied at atomic detail for up to hundreds of nanoseconds, reproducing critical physicochemical properties when compared with experiments. This allows the investigation of relevant phenomena, like self-assembly of lipid aggregates (19–21), phase transitions (22,23), formation and closure of pores (24), insertion and folding of membrane peptides (25,26,29), or hydrophobic-mismatch adaptations (27,28).

In this work, we study the orientation and dynamics of hydrophobic peptides in a dimyristoylphosphatidylcholine (DMPC) bilayer by means of MD simulations. The peptides belong to series of well-characterized transmembrane systems, introduced by Davis and co-workers (30), and profusely studied by the group of Killian $(4,7,11,31-35)$; namely, we use acetyl-GW₂L₁₇W₂A-ethanolamine (WLP23)

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and acetyl- $GK₂L₁₇K₂A$ -ethanolamine (KLP23) (33). The investigation of multiple replica simulations allows accounting explicitly for time-evolving and ensemble orientation distributions of the peptides. This provides a comprehensive description of dynamic peptide-membrane complexes, consisting of fluctuating orientational states. Finally, we show that the use of these explicit orientation distributions yields a coherent interpretation of the ²H-NMR data available for the same systems.

METHODS

Simulations

Software and general simulation conditions

We use the GROMACS package for all our simulations (36). The united-atom lipid parameters were adapted from the work of Berger and co-workers (37) and the peptides used the GROMOS force field. A time step of 4 fs was used (38). To ensure that the lipids were in the fluid phase, the temperature was set to 308 K, coupled to a Berendsen thermostat (39) with a coupling constant of 0.1 ps. The pressure was coupled semiisotropically to a Berendsen barostat (39), with a coupling constant of 1.0 ps. Both, the short-range electrostatic and van der Waals interactions were calculated using a cutoff of 1.0 nm, while for long-range electrostatics we used a PME algorithm (40).

Setup protocol

A total of 10 simulations were performed: one for a purely lipid system, conducted up to 50 ns; five for a WLP23-membrane complex, run for 200 ns each; and four for KLP23-membrane complex, up to 300-ns each. In all cases the lipid bilayer consisted of 128 DMPC lipids (64 per leaflet), in presence of 3655 water molecules. Chloride ions were included in simulations with the KLP23 peptide to ensure a electrically neutral simulation cell. The starting DMPC bilayer was downloaded from http://moose.bio.ucalgary.ca/ downloads.

Peptides were generated as ideal α -helices using the software Swiss PDB viewer (41) (http://www.expasy.org/spdbv), with the N-termini acetylated and the C-termini amidated. They were solvated with SPC water molecules in a cubic box of 5.0 nm, energy minimized and simulated during 5 ns with position restraints on the peptide backbone. A single relaxed peptide was inserted symmetrically across the bilayer, with its long axis aligned with the membrane normal and the system was solvated in a bath of water. After energy minimization, preequilibrium simulations were performed for 2 ns, with position restraints on the peptide backbone atoms, before the production runs.

Data analysis

Definition and calculation of membrane-peptide orientation

The static orientation of a rodlike α -helical peptide in a lipid membrane is defined by the tilt (τ) of the helix long-axis (H) with respect to the membrane normal (N, here corresponding to the z axis), and the polarity or azimuthal rotation (ρ) of the helix about its molecular long-axis. The azimuthal rotation ρ is found in the plane perpendicular to the helix axis, as the angle between a vector t pointing in the direction of the tilt and a reference vector r pointing into the C^{α} of an arbitrary residue, here Gly¹ (Fig. 1).

The helix axis was calculated for each frame of the simulations by using the backbone atoms of the 17 central Leu residues. The tilt vector is calculated as $t = h \times (h \times n)$, with h and n being unit vectors parallel to the helix axis (H) and the membrane normal (N) , respectively (Fig. 1). In practice, throughout the simulations the peptide structure diverged slightly from ideal. This was found not to affect significantly the calculated helix axis and the values of τ , but it can affect ρ , as it may change the relative angular position of the

FIGURE 1 Static orientation of a helical peptide, bound to a lipid bilayer. A pair of angles, tilt (τ) and rotation (ρ) , is sufficient to define the peptide orientation. The value τ is the angle formed between the molecular long axis (*H*) of the helix and the membrane normal (*N*). The value ρ is the angle between the direction of the peptide tilt (t) and a vector r perpendicular to H , pointing to the C^{α} carbon of a reference residue, here Gly¹.

reference Gly¹. Thus, representative values of ρ were calculated as averages of rotations referred to residues in the center of the peptide $(Leu^{11}-Leu^{14})$ and translated into the virtual ideal position of Gly¹, using a pitch of 100 $^{\circ}$ between contiguous residues.

Experiment-like orientations

Reported experimental τ - and ρ -values for WLP23 and KLP23 (33) have been determined from ²H-NMR quadrupole splittings ($\Delta \nu$) of the $-C^{\beta}D_3$ group in deuterated Ala, substituting, one by one, four central Leu residues (11–14) of the helix. To make a fair comparison of these experimental values with our simulations, we calculated virtual (experiment-like) ${}^{2}H$ splittings that would correspond to the orientation of C_{α} -C_β bonds in residues 11–14, from which we obtained experiment-like τ - and ρ -angles by using a fitting procedure similar to the GALA (geometric analysis of labeled alanines) (11). Briefly, assuming that the membrane normal is aligned with the magnetic field, instantaneous (Δv_{ins}) static splittings for each residue in a particular frame of the trajectory are first calculated as

$$
\Delta \nu_{\rm ins} = 3/4 (e^2 Q q / h) (3 \cos^2 \theta - 1), \tag{1}
$$

where (e^2Qq/h) is the nuclear quadrupole coupling constant, taken as 168.0 kHz for a C-D group (7,11), θ is the orientation of the C^{α}–C^{β} bond (virtually the C^{α}-C^{β}D₃ vector) with respect to the membrane normal. $\Delta \nu_{ins}$ values are multiplied by one-third to account for fast motional averaging about the C^{α} - $C^{B}D_{3}$ bond in virtual methylene deuteron splittings. These are then averaged over the full trajectory, for each individual simulation, or over a complete set

of trajectories, for a global ensemble corresponding to each peptide, and the absolute value of the average (Δv_{aver}) is considered for further treatments. Finally, experiment-like orientations $\{\tau, \rho\}$ are obtained for each set of four experiment-like splittings from a fit of the theoretical equation (11,33),

$$
\Delta \nu_{\text{theo}} = 3/4K \{ 3\cos^2 \varepsilon_{\parallel} [\cos \tau - \sin \tau \cos \tau \cos \tau - (\rho + \varepsilon_{\perp} + \varphi) \tan \varepsilon_{\parallel}]^2 - 1 \}, \tag{2}
$$

where $K = 1/3(e^2Qq/h)S$, with the factor 1/3 associated with fast rotation about the C^{α} - $C^{\beta}D_3$ bond and S being an order parameter accounting for peptide dynamics. The angle ε_{\parallel} is that defined between the $C^{\alpha}-C^{\beta}$ bond vector of the residue under consideration and the helix axis, ε_{\perp} is an angle between the C^{α} - C^{β} bond and a vector from the helix axis to the C^{α} , both projected onto a plane perpendicular to the helix axis, and φ is the rotation pitch angle between the C^{α} of the reference Gly¹ and the C^{α} of the residue considered for each particular Δv_{aver} splitting. As in the experimental studies we want to compare with Özdirekcan et al. (33), we use $S = 0.875$, giving $K = 49$ kHz, a pitch of 100° between contiguous residues, corresponding to an ideal α -helix, and a constant estimated value of -43.3° for the angle ε_{\perp} . Then, values of τ , ρ , and ε_{\parallel} are optimized during the fitting procedure, while minimizing root mean-squared deviations between Δv_{aver} and Δv_{theo} .

Lipid order parameters

The order of hydrocarbon chains in lipid bilayers is characterized by the deuterium order parameter S_{CD} measured through 2 H-NMR experiments. If θ_{i} is the angle between a C-D bond of a methylene group, i , and the bilayer normal, aligned with the direction of the applied magnetic field, the order parameter of that particular group is defined as

$$
S_{\rm CD}(i) = 1/2 \langle 3\cos^2 \theta_i - 1 \rangle, \tag{3}
$$

where the brackets denote time and ensemble average. The absolute value of the order parameters of the methylene segments is reported. Since we employed a united-atom force field, the order parameters were calculated from the positions of the carbon atoms along the chain (42).

Peptide structure analysis

Profiles of secondary structure of peptides along the simulation trajectories were obtained with the help of standard GROMACS analysis tools, which use the DSSP algorithm (43).

RESULTS AND DISCUSSION

Orientation and dynamics of the membrane peptides

In the axially ordered lamellar phase of a lipid membrane system, the orientation of an embedded helical-peptide is well described through the pair of angles $\{\tau, \rho\}$, giving, respectively, the helix tilt with respect to the membrane normal and the helix rotation about its molecular axis (see Methods and Fig. 1). However, membrane peptides are far from being rigid rodlike molecules at a unique and fixed orientation. In the fluidlike membrane, peptides experience complex wholebody movements, varying their tilt or their rotation and introducing time- and ensemble-dependent orientational diversity. Additionally, internal peptide dynamics and distortions from often assumed ideal α -helices, may affect the direction of the molecular axis and the reference for determining the ρ -angle.

In all our simulations (see a list in Table 1), the peptides were found to maintain an α -helical structure along the full trajectories (shown in Fig. 1 of the Supplementary Material). Thus, for the analysis and discussions below we will focus mainly on whole-body time-dependent fluctuations and variability of orientation states within a molecular ensemble.

Peptide tilt

After a quick displacement away from the starting orientation, where the helix axis was aligned with the membrane normal, τ oscillates with variable amplitude and frequency throughout the complete simulation. Some representative cases are shown in Fig. 2, graphs A and B . Due to the slow relaxation of peptide reorientation (15,16) we do not expect reaching equilibrium within the simulated times. Thus, we will use here the term ''stabilization time'' as the approximate time needed

TABLE 1 Summary of simulations of peptide/membrane complexes with parameters defining peptide orientation

Peptide	Simulation number	Total simulation time (ns)	Orientational parameters*					
			Direct (averaged angles)		Indirect (from calculated ${}^{2}H$ splittings)			
			τ (°) ^{\dagger}	ρ (°) ^t	τ (°)	ρ (°)	ε_{\parallel} (°)	$RMSD‡$ (KHz)
WLP23		200	40 ± 5	171 ± 18	37	169	55.3	1.10
	2	200	16 ± 10	115 ± 86	10	108	56.0	0.35
	3	200	39 ± 5	177 ± 14	40	181	55.9	1.70
	4	200	36 ± 5	122 ± 17	38	124	57.7	0.76
	5	200	24 ± 6	235 ± 29	25	245	56.6	0.20
	Global	1000	31 ± 12	173 ± 58	19	167	56.1	0.50
	Experiment				8	176	58.7	0.40
KLP23		300	35 ± 7	347 ± 22	36	347	56.4	1.11
	2	300	12 ± 7	204 ± 107	4	281	58.2	0.48
	3	300	15 ± 6	200 ± 63	11	221	57.6	1.48
	4	300	20 ± 7	188 ± 48	16	199	57.6	0.22
	Global	1200	20 ± 11	206 ± 104	6	260	58.7	1.52
	Experiment ¹				8	265	59.3	0.70

*The first 50 ns of the trajectory were discarded.

[†]Error intervals are given as mean \pm SD.

[‡]Root mean-squared deviations with respect to theoretical ²H quadrupole splitting values (given by Eq. 2) in the fitting procedure.

[¶]Experimental values from Özdirekcan et al. (33).

for the system to evolve outside the starting state, into a relatively stable characteristic orientation, for which we will perform our analysis. This process and the characteristic tilts are rather heterogeneous among the two peptides and corresponding replicas. For WLP23 (Fig. 2 A), simulations 1, 3, and 4 appear to stabilize slowly (within \sim 50 ns) and passing through successive stages of small to intermediate tilts maintained for a few nanoseconds, before reaching relatively large (\sim 40°) and stable tilts (Fig. 2 A, black line). Simulations 2 and 5, however, evolve more rapidly outside the initial state (within \sim 10 ns), remaining stable at smaller tilt angles (\sim 10° and \sim 20°, respectively). After 150 ns, and for the rest of the explored time, the peptide tilt of simulation 2 increases up to \sim 30 $^{\circ}$ (Fig. 2 A, *red line*). For KLP23 (Fig. 2 B), stabilization occurs within the first 10 ns with characteristic tilts generally smaller than in the case of WLP23, except for simulation 1, which oscillates within the first \sim 50 ns and stabilizes at values close to 35° (Fig. 2 B, black line). The mean tilt angles for each simulation were calculated after discarding the first 50 ns (longest stabilization time), and they are given in Table 1.

Peptide rotation

The time evolution of the instantaneous rotation angle is shown in Fig. 2, C and D , for representative simulations. Because, by definition (Fig. 1), ρ depends on the direction of the tilt, and the peptide can be assumed to incline initially at random, the azimuthal rotation may start at any value. For the same reason, a small and fluctuating τ , as occurring often for short simulation times, may easily originate abrupt fluctuations of ρ . After the consensus 50-ns stabilization time (defined above from the evolution of τ), the helix rotation tends to equilibrate, although with persisting fluctuations

(Fig. 2, C and D). These are of small amplitude ($\pm 20^{\circ}$) for $\tau > 20^{\circ}$, or large jumplike transitions, for $\tau < 20^{\circ}$, the latter case being most typical for KLP23. As for the peptide tilt, the mean azimuthal angle for each replica is given in Table 1.

As observed before in other MD studies (25,28), the average tilts of the simulations are systematically larger than values derived experimentally from 2 H-NMR (33). With respect to the azimuthal rotations, although they tend to approach the experimental angles, there are also some notable discrepancies (see Table 1). However, as we shall see, the explicit dynamics of the system, in the form of a large spread of angular values and distinguishable orientational states, has an impact on the interpretation of experimental data.

Distributions of orientations

Time-dependent distributions

The diversity and relative weight of peptide orientations is better represented by frequency distributions of tilt and rotation angles, as depicted in Fig. 3. In general, the distributions of τ for individual simulations approximate to unimodal, Gaussian-like probability density functions, with characteristic standard deviations of $\sim 10^{\circ}$, similar to other reported distributions from MD simulations (25). Exceptions are simulation 2 of WLP23 (Fig. 3 A), which shows a principal τ -mode at \sim 12°, and a second less populated mode at \sim 32°, and simulation 2 of KLP23 (Fig. 3 B), with an asymmetric peak of maximum frequency at \sim 7° and a tail toward long τ -values.

The individual distributions of ρ are more broad, with standard deviations larger than 20°. Simulations of WLP23 give a set of partially overlapping and fairly symmetric, single bell-shaped peaks between \sim 100 $^{\circ}$ and 260 $^{\circ}$, although

FIGURE 2 Time evolution of peptide orientation. The graphs represent instantaneous values of the angles τ (A and B) and ρ (C and D) along the time coordinate for representative case examples of WLP23 (A and C) and KLP23 (B and D). In panels A and C , data from simulations 1, 2, and 5 are represented with colors black, red, and orange, respectively. In panel B, data from simulations 1, 2, 3, and 4 are drawn black, red, green, and blue, and in panel D data from simulations 2, 3, and 4 are red, green, and blue, respectively.

FIGURE 3 Frequency distributions of orientational parameters. Shown are probability densities of τ (A and B) and ρ (C and D) from peptides WLP23 (A and C) and KLP23 (B and D). The graphs show data from simulations 1 (black, continuous line), 2 (black, dashed line), 3 (black, dashed-dotted line), 4 (shaded, dashed line), and 5 (shaded, dashed-dotted line). Global distributions of orientations found in the complete set of simulations, for each peptide, are drawn as a shaded continuous line.

runs 2 and 5 show significant intensity outside this interval (Fig. 3 C). In contrast, for KLP23 (Fig. 3 D), only simulation 1 is unimodal and centered around \sim 350°, while ρ distributions from runs 3 and 4 are overlaps of partially resolved peaks around \sim 195 $^{\circ}$ and simulation 2 shows a very large spread of values, with resolved broad peaks in both the 350° and 195° regions.

Although all simulations display considerable orientational diversity, a comparison among them suggests that each one represents only part of the total dynamics of the system. One may think of increasing the available configurational space by increasing the simulation time, although this does not appear feasible within a reasonable limit (note that our simulations are 200–300-ns long). We may also enlarge this space through a global treatment of a set of replicas, as if they represented a small molecular ensemble. As we show below, such a global analysis suggests a complex orientational landscape, with a broad structural diversity and the emergence of different states.

Global distributions

For each of the two peptides the collective distribution of τ gives an asymmetric profile with a large spread of angles. In the case of WLP23, this shows a preference for large tilts, with a most prominent peak at \sim 38° and a mean at 31° \pm 12° (Fig. 3 A). However, for KLP23, the principal peak seats at relatively small tilts (around 15°), although coexisting with a resolved peak at larger tilts (at \sim 37°), all together averaging at $20^{\circ} \pm 11^{\circ}$ (Fig. 3 B).

The two peptides show also distinguishable global ρ distributions (Fig. 3, C and D; violet lines). In this case, due to their complexity, the global analysis is more clearly made from representations of tilt vectors (see Fig. 1), which can be plotted directly over an Edmundson's helical-wheel of the peptides (Fig. 4, A and B). Making the modulus of each tilt vector proportional to the occurrence of its corresponding ρ -angle, the average ρ is determined from the direction of the resultant vector. For WLP23, the tilt vectors group densely in a sector between 100° and 260° , with their sum forming an angle $\rho = 173^\circ$ with respect to the reference vector (Fig. 4 A). In contrast, the complex global distribution for KLP23 (Fig. 3D, shaded continuous) corresponds to tilt vectors populating two sectors (Fig. 4 D), with a global average defining a $\rho =$ 206 $^{\circ}$. These global ensemble averages of ρ approximate to values determined from experiments (33) with differences within the standard deviation. However, the global tilt angles are considerably larger in the simulations (Table 1). The origin of these discrepancies is discussed later with more detail.

Orientational states

The global distributions appear generally asymmetric and/or multimodal (Fig. 3). Although this could be due to incomplete sampling of possible orientations, it suggests the existence of characteristic populations or states within the ensemble. A plot of $\{\tau, \rho\}$ pairs against their frequency (Fig. 5) permits defining these possible states. In the case of WLP23 there is a most probable state with τ around 38° and well-defined values of ρ close to 180°. Additionally, there are much less probable orientations for smaller tilts, which for $\tau < 20^{\circ}$ correspond to a very wide distribution of ρ -values (Fig. 5 A). Thus, a representative structure of this peptide in DMPC is that defined by the pair of angles $\{\tau = 38^{\circ}, \rho = 173^{\circ}\}\,$, depicted in Fig. 6 A. On the other hand, KLP23 is clearly more dynamic $(Fig. 5B)$. For this peptide the small tilts again correspond to a broad peak in the ρ -dimension, which is most intense at \sim 195°, while the large tilts correspond to a well-resolved

FIGURE 4 Vectorial representation of peptide rotations. Tips of tilt vectors, depicted as open circles, are drawn over an Edmundson helical wheel of the peptides WLP23 (A) and KLP23 (B) . Only most important residues are represented: reference $Gly¹$ and anchoring Trp (A) or Lys (B), with solid representation for residues at the N-terminus and shaded representation for residues at the C-terminus. The ρ -angles are defined with respect to the reference vector pointing into Gly^1 , fixed at the horizontal axis (see Fig. 1 for detailed definitions). Being the modulus of each tilt vector proportional to the occurrence of the corresponding rotation, the resultant tilt vector ($arrow$) marks the average ρ .

 ρ -peak around 350 $^{\circ}$. We can then define two representative states for KLP23, with pairs of most probable angles $\{\tau = 18^{\circ},\}$ $\rho = 195^{\circ}$, depicted in Fig. 6 B, and $\{\tau = 37^{\circ}, \rho = 350^{\circ}\}\,$ depicted in Fig. 6 C.

FIGURE 5 Global distributions of pairs of peptide orientational angles. (A) WLP23. (B) KLP23.

The just-described large orientational variability, including the existence of alternative states, stable within at least a few hundreds of nanoseconds, contrasts with views of a static and well-defined membrane peptide orientation. Membrane peptides require relatively long simulation times to equilibrate (28) which, for whole-body orientational properties, due to their long correlation times (15,16), might be up to a few microseconds. Here, simulations needed up to several tens of nanoseconds before acquiring characteristic orientations, which, despite still outside equilibrium, may be close to possible stable states. Note that, although each simulation samples principally one state (Fig. 3), the large overlap among the individual distributions shows some level of transition between states within the 200–300-ns time range. Moreover, for small tilt values, transitions between ρ -states are clearly facilitated (as in simulations 2 of both peptides).

Implications for the analysis of experimental data

The systems under study are among the best-characterized peptide-lipid complexes. In particular, it has been reported from solid-state ²H-NMR that peptides of the WLP/WALP and KLP/KALP families possess well-defined pairs of τ and ρ in DMPC lipid bilayers (11,33). Special attention has been paid to the tilt angles, since the values determined through ²H-NMR methods are in general surprisingly small, even under positive hydrophobic mismatch, and compared with other experimental tilts from transmembrane peptides (31,44–46). Although dimerization has been claimed among the possible causes for the small tilts, no proof for oligomers at relevant conditions has been documented (34). With respect to the rotational angles, the data are more scarce and in general show that ρ adopts characteristic values that depend on the type of interfacial residues (33).

Influence of rotational dynamics on ²H NMR observables

We have seen above that simulated KLP23 and WLP23 peptides in membranes mainly present tilt angles larger than derived from ²H-NMR experiments, while the helix azimuthal rotations are closer to the reported experimental data, specially for the global ensemble averages (Table 1). We should note, however, that experimentally determined orientations are, as well, calculated values, obtained on the basis of an assumed model where the unknown peptide dynamics is largely simplified. For instance, internal motions, wobbling and fluctuations of the helix rotation are, at most, collectively imputed as an order parameter factor. To test the impact of explicit peptide dynamics in the structural interpretation of ²H-NMR data, we calculate orientation angles from the simulations by applying the same model and mathematical framework as it was used for the experiments (7,11,33). Thus, static ²H quadrupolar splittings corresponding to the four residues labeled for the NMR experiments (residues 11–14) were first back-calculated from each frame of the simulations,

FIGURE 6 Characteristic structures of peptide-membrane complexes. The models were chosen out of the complete set of trajectories to match the most populated $\{\tau, \rho\}$ pairs (Fig. 5). (A) WLP23 at $\tau = 38^{\circ}$, $\rho = 173^{\circ}$. (*B*) KLP23 at $\tau = 18^{\circ}$, $\rho = 195^{\circ}$. (*C*) KLP23 at $\tau = 37^{\circ}$, $\rho = 350^{\circ}$. The lipid acyl tails are depicted in light gray and headgroup atoms are shown in red. The peptides are drawn as light blue ribbons, showing only side chains of anchoring residues, $Trp(A)$ and Lys (B, C) , in yellow and dark blue, respectively. The C^{α} of Gly¹, marking the reference for the rotation angle, is shown as a green sphere.

using Eq. 1, and subsequently averaged over a full trajectory. The tilt and rotation angles were then obtained by fitting a theoretical curve given by Eq. 2 (called quadrupolar wave (11)) to a set of experiment-like splittings. For most simulations of WLP23 and for simulation 1 of KLP23, the τ - and ρ -angles so obtained are very similar to the values determined directly from the corresponding time averages. However, for run 2 of WLP23 and runs 2–4 of KLP23, the experiment-like tilts are smaller than the values calculated directly (Table 1). Simulations of this second group are characterized by broad distributions of ρ (Fig. 4), which effectively reduce the calculated quadrupolar splittings (see Table 1 of Supplementary Material). Interestingly, this affects mainly the fitted tilts, which get close to the experimental values (Table 1). Thus, it appears that a large spreading of the ρ -angle leads to a reduction of the tilt angles obtained by this method.

As we would expect from a representative ensemble distribution of orientation states, the collective treatment of inferred quadrupolar splittings for each peptide yields experiment-like splittings which generally compare closely with values determined from solid-state ²H-NMR (Table 1 of the Supplementary Material). This shows that our limited sets of simulations are sufficient to reproduce an important part of the peptide orientational landscape. Consequently, the experiment-like $\{\tau, \rho\}$ pairs are in good agreement with the corresponding solid-state ²H-NMR values (Table 1). It is important to notice that although for some of the individual simulations the calculated splittings (and corresponding fitted angles) approach the experimental ones, the best results for each peptide were obtained when considering globally all the corresponding simulations. Thus, the dynamics exhibited by a limited number of replicas, within 200–300 ns time windows, appears a fair representation of the dynamics of the peptide/membrane complex. Moreover, the latter results stress that the rotational diversity defining these systems is contributed by the different orientational states, each of them exhibiting time-dependent fluctuations.

The findings described above solve a recurrent paradox about the orientation of WLP and KLP peptides studied by

 2 H-NMR, and offer an explanation (most likely extensible to WALP and KALP peptides) for the small tilts, barely reacting to mismatch, generally obtained in these cases (7,11,13,32,33). While a uniform rotation about the peptide long axis is to be discarded from the inequality of the splittings for different residues around the helix (7,11,33), such inequality is compatible with nonuniform broad fluctuations of ρ , as demonstrated here. This rotational dynamics, in a nanosecond timescale, can correspond to the axial diffusion with a 10^{-8} – 10^{-7} s correlation time reported from solid-state NMR relaxation experiments for peptides very similar to KLP23 (15,16). The same relaxation studies report small amplitude off-axis reorientations in the 10^{-6} – 10^{-5} s time regime, which may correspond to the dynamics of the tilt, including slow exchange between the tilt states inferred from the simulations.

Reinterpretation of experiments including dynamics

For real case experimental studies, in the absence of an explicit description of the underlying rotation dynamics, ideal distributions of ρ could be evaluated. Using a Gaussian distribution reduces the calculated quadrupolar splittings with only small variations of the phase of theoretical quadrupolar waves. Thus, the extracted value of τ will be smaller, with a very similar ρ -angle (the mean of the distribution). However, this allows multiple fittings of similar quality for a broad range of tilts, depending on an arbitrary choice of the standard deviation of the distribution. For example, the four experimental splittings measured for WLP23 in DMPC (33) can be fitted for τ -values of 8° , 20° , and 30° , using distributions with standard deviations of 0° , 71° , and 87° (Fig. 7). Although one might consider the system to be underdetermined with only four data points, all them in a helix turn, the use of eight experimental ²H splittings well distributed along the helix, as measured for WALP23 in DMPC (11), does not lift the ambiguity (see Fig. 2 of the Supplementary Material). Despite the unresolved τ , the ρ -angle is better determined, with values 176° , 185° , and 186° , respectively, for the three alternative fittings considered in Fig. 7. It should be realized, however,

FIGURE 7 Best fit of quadrupolar splitting waves. Theoretical waves calculated by Eq. 2 are fitted to experimental values for WLP23 in DMPC (solid circles) (33). (Solid line, $\tau = 8^\circ$, $\rho = 176^\circ$, SD = 0.0°, error = 0.4 kHz, and $\varepsilon_{\parallel} = 58.7^{\circ}$. Dashed line, $\tau = 20^{\circ}$, $\rho = 185^{\circ}$, SD = 71°, error = 0.25 kHz, and $\varepsilon_{\parallel} = 58.7^{\circ}$. Shaded line, $\tau = 30^{\circ}$, $\rho = 186^{\circ}$, SD = 87°, error = 0.17 kHz, and $\varepsilon_{\parallel} = 59.7^{\circ}$.)

that the ambiguity will also affect ρ in cases with multimodal distributions of this latter parameter, like in the Lys-flanked peptide (see below). A detailed systematic evaluation of the expected influence of orientational dynamics on ²H NMR data will be published elsewhere.

At this point, the question arises about possible consequences of extensive molecular motions on other experimental NMR observables, typically used to obtain orientational information, like ^{15}N chemical shifts and $^{15}N^{-1}H$ dipolar couplings. Both parameters are usually measured in twodimensional correlation spectra, known as polarization inversion spin exchange at the magic angle (PISEMA) (47), from which characteristic patterns, or polarization index of the slant angle (PISA) wheels, are obtained and assigned to welldefined peptide orientations (48,49). Unfortunately, there are no PISEMA spectra published so far for WLP/WALP or KLP/ KALP model peptides, which would provide an independent experimental measurement of orientation for those cases and allow comparison with our simulation results. However, because both the ^{15}N chemical shift anisotropy and $^{15}N^{-1}H$ dipolar interaction tensors align almost parallel to the axis of rotation (helix long axis), we expect this type of measurement to be much less influenced by rotational dynamics than the ${}^{2}H$ NMR splittings. Supporting this idea, a recent hydrophobic mismatch study of cell-signaling peptides in bilayers of different thickness, using both MD simulations and PISEMA experiments, found a very good agreement in the tilts obtained from the two methods (50). In fact, simulated PISEMA spectra, considering motions around a central ρ -value, predict only small effects on the corresponding PISA wheels (51). However, the same study also shows a significant influence of librational motions of the peptide planes and wobble motions about a central tilt. This may cause ambiguous results which can be alleviated by increasing the number of spectral as-

signments. Clearly, more extensive investigations, combining NMR experiments, MD simulations, and including dynamics for the interpretation of spectra, are needed to completely solve this issue.

What determines peptide orientation and dynamics?

The orientation of membrane peptides is an important structural quality to define the molecular organization in peptide/ membrane complexes. This is usually rationalized within the mattress model (9), which basically explains peptidemembrane interaction as a mutual adaptation dominated by rules of the hydrophobic mismatch (10), defined in turn as the difference between the hydrophobic length of the peptide and the hydrophobic width of the lipid bilayer. However, other intrinsic characteristics of the peptide and the bilayer lipids, like the nature of interfacial peptide residues and the lipid headgroup, may interplay together with hydrophobic matching to condition the type and properties of peptide-lipid interactions. Additionally, the dynamics of the membrane allows orientational fluctuations of embedded peptides, which may originate alternative peptide-membrane binding states and facilitate transitions between them. We will discuss now briefly how the dynamic peptide orientation is achieved through mutual adaptations of the peptide and the membrane.

Peptide adaptation to membrane binding

The apolar core of a DMPC bilayer is \sim 23.0 Å thick (52). In turn, the hydrophobic length of the two peptides used in this study can in principle be approximated to 25.5 Å , considering the central 17 Leu residues and a 1.5 Å rise per residue for an ideal α -helix. Therefore, we are under conditions of positive hydrophobic mismatch and the peptides are expected to tilt to alleviate unfavorable contacts. However, the response of the lipid bilayer and uncertainty, or variability, of anchoring interactions between residues at the peptide ends and groups in the membrane interface, make mismatch, a priori, difficult to determine. For instance, although according to the simple calculation above, WLP23 and KLP23 would exhibit the same hydrophobic length, the first one shows clearly larger tilts, both for individual simulations and global averages (Table 1). This demonstrates a different effect of the Trp and Lys residues on the effective peptide orientation, either directly, by a different contribution to the total hydrophobic length, or indirectly, through distinct localized interactions of each type of residue at the membrane interface. In practice, both effects can be observed. The large rings of Trp are embedded at the level of the glycerol group, partially penetrating the hydrophobic core of the membrane (Fig. 6 A). A similar localization has been found for tryptophan and tryptophan analogs by NMR and MD simulations (53–55). This relatively deep interfacial position is in agreement with the large free energies of transfer of Trp, both to the interface

 $(\Delta G_{\text{wif}} = -1.8 \text{ kcal/mol})$ and to *n*-octanol $(\Delta G_{\text{woct}} = -2.1$ kcal/mol) in whole-residue hydrophobicity scales, with a difference favorable for insertion in the membrane core $(\Delta G_{\text{word}} - \Delta G_{\text{wif}} = -0.3 \text{ kcal/mol})$ (56). Thus, the total effective hydrophobic length for WLP23 should account, at least partially, for the contribution of the flanking Trp residues, implying larger values of positive mismatch and peptide tilt. On the other hand, the positively charged Lys in KLP23 is more promiscuous, leading to alternative binding states. It shows a preference to interact with the polar phosphoryl groups of the phospholipids, which, in principle, restrains the peptide to relatively smaller tilt angles (Fig. 6 B). However, the long aliphatic chain with large conformational flexibility of Lys allows also a deeper binding of this residue, while its charged amino group can still reach the polar region via snorkeling. This latter possibility corresponds to a larger effective peptide hydrophobic-length and a larger tilt (Fig. 6 C).

With respect to the rotational angle, it is thought to be determined mainly by the position and type of the N- and C-terminal peptide-anchoring residues. For instance, ρ -values differing by \sim 100 $^{\circ}$ have been found experimentally for the WLP23 and KLP23 peptides (33). As the Trp and Lys interfacial residues occupy equivalent positions around the helix, the observed rotation angles were ascribed to differences in their mode of interaction with the bilayer headgroup region. From global average treatments of the simulations we arrive to ρ -values similar to the experimental ones (Table 1). However, the details of the distributions show that KLP23 is best described by two orientational states (Fig. 3, B and D, and Fig. 4 B): one associated with an average ρ at \sim 195°, similar to the characteristic rotation of WLP23 (173°), and a second with an average ρ at \sim 350°. Therefore, there seems to be a preferred azimuthal rotation characteristic of both peptides and determined by the position of the anchoring residues about the helix, where the tilt vector points between the two N-terminal Trp or Lys residues (Figs. 4 and 6, A and B). Additionally, the type of interfacial residue also matters. Thus, different from Trp, the large and flexible chain of Lys is involved in stabilizing alternative orientations with a tilt vector pointing toward Gly^1 (Fig. 4 and Fig. 6 C).

Membrane response to a TM peptide

Peptide binding across a membrane affects its structure and dynamics. Typical consequences are appreciable changes in the electron density profiles, measured by x-ray diffraction (57), and variations in the order parameters of lipid tails, determined by 2 H-NMR (31), both indicating changes in the membrane thickness. Molecular dynamics simulations can assess these membrane properties directly, allowing a useful comparison with experimental studies and a detailed evaluation of underlying biophysical phenomena.

The order parameters of lipid tails from simulations of a pure DMPC bilayer are in good agreement with those from 2 H-NMR experiments (58). If represented against the ali-

FIGURE 8 Lipid order parameters. Represented are absolute values of S for methylene groups of the $sn-2$ acyl chain of DMPC. Shaded lines correspond to pure lipid bilayers, analyzed by 2 H-NMR experiments (58) (stars, continuous line) or MD simulations (diamonds, dashed line). The solid lines are order parameters in the presence of peptides, determined from experiments (WALP19 (31), triangles, dashed-dotted line) or simulations (WLP23, circles, continuous line, and KLP23, squares, dashed line). Note that the experimental data correspond to carbons C3 to C14, while simulated data extend from C2 to C13.

phatic carbon position (Fig. 8), the shape of the curve is well reproduced by simulations, although the absolute values are slightly smaller for some methylene groups. In presence of WLP23 or KLP23, we observed an increase of the order parameters, averaged over the total set of simulations, for the 30 lipids nearest to the peptide (Fig. 8). Although there are no experimental order parameters available for membrane complexes with WLP23 or KLP23, a similar increase of order parameters has been reported in the presence of WALP19 in DMPC, at a peptide/lipid molar ratio of 1:30 (31). Nevertheless, we noticed that the increase of the bilayer width accompanying the changes in the order parameters is only marginal, as reported from experiments (57), and peptide tilting appears to be the principal adaptation upon peptide/ membrane interaction.

CONCLUSIONS

Despite the limited time window and small number of replicas, conforming to a reduced molecular ensemble, our observations can be considered an explicit representation of the complex orientational dynamics of *trans*-membrane peptides. The existence of fluctuating alternative $\{\tau, \rho\}$ states, arrestable within at least a few hundreds of nanoseconds, is the most relevant property. Such a view contrasts with often envisioned static, well-defined membrane peptide orientations and stresses the importance of dynamics to describe the molecular organization of these systems. Dynamic models should thus ideally accompany structural definitions in peptide/membrane complexes as a necessary ingredient to understand their functionality, especially in cases of intrinsically dynamic

processes like membrane insertion, pore formation, membrane fusion, and intermolecular peptide association.

We have demonstrated as well the importance of properly considering peptide dynamics to determine orientational parameters from spectroscopic ²H-NMR data, giving an explanation to counterintuitive small tilts in model WLP/WALP and KLP/KALP peptides. Although general implicit distributions are not easy to find, explicit dynamics representations obtained from a limited number of MD simulations can be used to guide the calculation and interpretation of ${}^{2}H$ NMR splittings from experimental data. Such rotational dynamics of the peptide is expected to have a more limited influence on $15N$ chemical shift and $15N-1H$ dipolar coupling NMR observables.

Finally, the fact that the orientation of the peptides, and particularly the rotational angles, are well reproduced by the MD simulations, implies that the principal features of the peptide-membrane interaction are well captured. This opens the possibility to analyze such interactions from atomicdetailed dynamic models, with particular emphasis in the membrane interface.

SUPPLEMENTARY MATERIAL

To view all of the supplemental files associated with this article, visit www.biophysj.org.

Note added in proof: While this article was under review, an independent MD study on the model peptide WALP23 has been completed (59). Similar to our work, this latter investigation also shows that the large fluctuation of the peptide rotational angle reduces back-calculated 2 H-NMR splittings, which can be the origin of abnormally small tilts when determined from experiments.

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