Orientation determination of membrane-disruptive proteins using powder samples and rotational diffusion: A simple solid-state NMR approach

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Abstract

The orientation of membrane proteins undergoing fast uniaxial rotation around the bilayer normal can be determined without macroscopic alignment. We show that the motionally averaged powder spectra exhibit their 0° frequency, \( \delta_{0} \), at the same position as the peak of an aligned sample with the alignment axis parallel to the magnetic field. This equivalence is exploited to determine the orientation of a \( \beta \)-sheet antimicrobial peptide not amenable to macroscopic alignment, using \( ^{13} \)CO and \( ^{15} \)N chemical shifts from powder spectra. This powder sample approach permits orientation determination of naturally membrane-disruptive proteins in diverse environments and under magic-angle spinning.

1. Introduction

The orientation of membrane proteins has been tradition-ally determined in solid-state NMR by means of macroscopically aligned samples [1]. When the membrane is uniaxially aligned so that the bilayer normal is parallel to the magnetic field (\( B_{0} \)), the NMR spectrum of a single site collapses into a single line at a frequency that reflects the orientation of the protein with respect to the bilayer normal.

However, aligning lipid membranes mechanically on glass plates or magnetically in bicelles is generally difficult. Many proteins cannot be aligned due to their inherent membrane-disruptive or curvature-inducing nature [2]. Usually only certain membranes are amenable to alignment for a specific protein. Alignment becomes more difficult as the protein size and concentration increase. For glass-plate samples, it is often difficult to control pH, ion concentration, or other parameters that may be relevant for the function of the membrane protein.

Therefore, it is desirable to determine membrane protein orientation without using macroscopic alignment. In fact, it has been realized that for molecules undergoing fast uniaxial rotational diffusion around the bilayer normal, orientation information can be obtained from unoriented samples [3,4]. Uniaxial mobility is present for membrane proteins in liquid-crystalline bilayers as long as the protein is not too large [5], and has been reported for many membrane peptides and proteins such as gramicidin A [6], protegrin-1 [7], KL14 and h\( \Phi \)19W [8]. Here we show a simple way of deriving the equivalence between aligned and powder samples, and apply this principle to \( \beta \)-sheet antimicrobial peptides, which are less well understood than \( \alpha \)-helices. \( ^{13} \)CO and \( ^{15} \)N chemical shift constraints obtained from powder samples are used to determine the orientation of a \( \beta \)-sheet antimicrobial peptide that has been resistant to macroscopic alignment.

2. Materials and methods

Protegrin-1 (PG-1) and tachyplesin-I were synthesized by Fmoc solid-phase methods as described before [7,9].
Unoriented proteoliposome samples were prepared by codissolving the peptide and lipids in chloroform and TFE, lyophilization, and rehydration to 35% water by mass. Aligned samples were prepared as described before [7]; the codissolved peptide and lipid organic solution was spread evenly on ~80 μm thick glass plates. The sample was vacuum dried thoroughly, and rehydrated at >95% humidity over a saturated salt solution for several days. The plates were stacked, wrapped in parafilm and sealed for measurements.

All spectra were acquired on a Bruker DSX-400 spectrometer (9.4 T) using a static probe. For aligned samples, a home-built rectangular radiofrequency (rf) coil was used, typical 1H decoupling field strengths and CP field strengths were 50 kHz.

3. Results and discussion

We first derive the equivalence between the frequency of an immobile and uniaxially aligned sample with the alignment axis parallel to the magnetic field $B_0$ (0°-aligned samples), and the 0° frequency, $\delta_0$, of a mobile unoriented sample. The frequency of an immobile 0°-aligned sample depends on the polar ($\theta$) and azimuthal ($\phi$) angles of $B_0$ in the principal axis system (PAS) of the relevant interaction tensor (Fig. 1a):

$$\omega_{0 \text{-aligned}} = \frac{1}{2} \delta(3 \cos^2 \theta - 1 - \eta \sin^2 \theta \cos 2\phi) + \omega_{iso}. \quad (1)$$

Here $\delta$ and $\eta$ are the anisotropy and asymmetry parameters, respectively, of the rigid-limit interaction tensor. Since $B_0$ is parallel to the alignment axis, ($\theta, \phi$) are also the polar coordinates of the bilayer normal in the PAS.

Uniaxial rotation around the bilayer normal in the 0°-aligned sample does not change the frequency, since this rotation is also around the magnetic field and thus does not change ($\theta, \phi$) (Fig. 1b). Thus, Eq. (1) also applies to mobile oriented samples.

For a mobile but unoriented sample, the motional axis is generally not parallel to $B_0$, thus the NMR spectra depend on the anisotropy parameter, $\delta$, of the motional averaged tensor (Fig. 1d). Since $\delta$ is the frequency difference from the isotropic frequency observed when $B_0$ is along the unique axis of the averaged tensor, which is the motional axis (Fig. 1c),

$$\delta = \omega_{0 \text{-aligned}} - \omega_{iso} = \frac{1}{2} \delta(3 \cos^2 \theta - 1 - \eta \sin^2 \theta \cos 2\phi). \quad (2)$$

This averaged anisotropy parameter, together with the averaged asymmetry parameter $\eta$ of 0, completely determine the powder lineshape of the protein. The 0°-edge of this powder pattern, which results from bilayer normals parallel to the magnetic field, appears at

$$\delta_{0} = \delta + \omega_{iso} = \omega_{0 \text{-aligned}}.$$ \quad (3)

In other words, the $\delta_{0}$ edge of the motionally averaged powder spectrum is identical to the frequency of the 0°-aligned sample. Thus, one can determine the orientation of membrane proteins using powder samples provided the protein undergoes uniaxial rotation faster than the interaction strength. Moreover, in the static spectra, one can determine $\delta_{0}$ from the high-intensity 90° peak, $\tilde{\delta}_{0}$, since the two are related by:

$$\delta_{0} = \tilde{\delta}_{0} - \omega_{iso}.$$ \quad (4)

Fig. 2 shows calculated motionally averaged 13CO and 15N powder spectra for several orientations of an ideal β-sheet peptide. The peptide was constructed with torsion angles ($\phi, \psi, \omega$) of (−139°, +135°, +178°), and exhibits little twist for the short length considered. Thus a single 13CO and 15N label was used to represent the overall orientation. The 13CO and 15N chemical shifts were calculated as a function of the polar coordinates of $B_0$ in a molecule-fixed frame defined by the β-strand axis and β-sheet plane. For the 13CO tensor, rigid-limit principle values of 248, 170, 100 ppm were used in the calculation, the $\sigma_{22}$ axis is parallel to the C=O bond, and the $\sigma_{33}$ axis is perpendicular to the peptide plane [10]. For the 15N tensor, the principal values are 217, 77, 64 ppm, the $\sigma_{11}$ axis is 17° from the N–H bond, and the $\sigma_{33}$ axis is 25° from the peptide plane [11]. Four orientations, defined by the tilt angle ($\tau$) of the β-strand axis from the bilayer normal and the rotation angle ($\rho$) of the β-sheet plane around the strand axis, were considered. Fig. 2 shows that the motionally averaged 13CO and 15N powder spectra are exquisite sensitive to the β-sheet orientation. For example, when the β-strand axis is perpendicular to the bilayer normal but the β-sheet plane is parallel to it ($\tau = 90°$ and $\rho = 0°$), the 15N powder spectrum has nearly rigid-limit CSA while the 13CO spectrum is extremely narrow (Fig. 2a and e). These result from the fact that the 15N $\sigma_{33}$ axis and the 13CO $\sigma_{22}$ axis are parallel to

Fig. 1. Schematics showing the equivalence between the 0°-aligned spectra and unoriented spectra. (a) Rigid 0°-aligned sample. (b and c) Mobile aligned sample. In (c), the motionally averaged tensor has the unique axis along the bilayer normal. (d) Mobile unoriented sample.
the bilayer normal at this orientation. When the β-sheet lies in the plane of the bilayer (\(\tau = 90^\circ\) and \(\rho = 90^\circ\)), the \(^{15}\)N CSA is reduced to half the rigid-limit value and inverted in sign (Fig. 2f), while the \(^{13}\)CO spectrum has the \(\delta_l\) edge close to \(\sigma_{33}\) (Fig. 2b). For all orientations, the \(0^\circ\)-aligned spectrum shows the same frequency as the \(\delta_l\) edge of the powder pattern. An example of the equivalence between the powder spectra and the oriented spectra for uniaxially mobile molecules is given by PG-1, a β-sheet membrane peptide [12]. Fig. 3 shows the \(^{13}\)CO spectra of Val 16\(^{13}\)CO-labeled PG-1. The \(0^\circ\)-aligned spectrum of PG-1 (a) exhibits a \(^{13}\)CO chemical shift of 216 ppm [7]. The unoriented sample gives an axially symmetric powder pattern with \(\delta_l = 216\) ppm and \(\delta_d = 151\) ppm (b), much narrower than the rigid-limit CO lineshape (Fig. 4f). The powder pattern, obtained with \(^1\)H–\(^{13}\)C CP, exhibits a sharp lipid signal at \(\approx 173\) ppm. After subtracting the lipid background signal using a single-pulse \(^{13}\)C spectrum, the difference spectrum of the peptide shows a ‘magic-angle hole’ at the isotropic shift (c). This is characteristic of the CP spectra of uniaxially mobile molecules, where the chemical shift and the dipolar coupling tensors are collinear with the motional axis. When the bilayer normal is \(54.7^\circ\) from \(B_0\), the averaged \(^{13}\)C CSA and the \(^1\)H–\(^{13}\)C dipolar coupling both vanish, thus abolishing CP at the isotropic shift. The \(\delta_l\) singularity of the powder spectrum is identical to the frequency of the \(0^\circ\)-aligned spectrum (d) obtained by tilting the glass plates to make the alignment axis perpendicular to \(B_0\). The \(\delta_d\) (216 ppm), isotropic shift (173 ppm), and \(\delta_l\) frequencies (151 ppm) are related by Eq. (4) as expected for uniaxial tensors.

This motionally endowed favorable frequency equivalence has been exploited indirectly in bicelle-bound membrane proteins [13]. When bicelles are aligned magnetically with the alignment axis perpendicular to \(B_0\), the fast uniaxial rotation of the protein–bicelle complex...
yields well-resolved $^{15}$N spectra whose $\delta_1$ frequencies are related to the $\delta_0$ frequencies of the $0^\circ$/C176-aligned glass-plate samples according to Eq. (4). We generalize this frequency equivalence to any orientation of the membrane, thus it is not necessary even to prepare bicelles.

It is important to note that the rotational diffusion of membrane proteins differs from that of lipids: most proteins are internally rigid, while the conformational flexibility of lipid molecules prohibits orientation determination even in the presence of global rotational diffusion [14].

We use this powder sample approach to determine the orientation of a $\beta$-sheet membrane peptide that has not been amenable to macroscopic alignment so far. Tachyplesin-I is a disulfide-linked $\beta$-hairpin antimicrobial peptide found in the hemocytes of the horseshoe crab, Tachyplesus tridentatus [15]. Fig. 4 shows the static $^{15}$N (a and b) and $^{13}$CO spectra (d and e) of $^{15}$N–Phe4 and $^{13}$CO–Val6 labeled TP-I in unoriented DLPC membrane. In the Lα-phase (a), the $^{15}$N spectrum shows a uniaxial lineshape, reduced anisotropy ($\delta_0 = 195$ ppm), and a magic-angle hole, indicating that TP-I undergoes fast uniaxial rotation. Cooling the peptide to below the phase-transition temperature

![Fig. 4](image_url)

Fig. 4. $^{15}$N (a and b) and $^{13}$CO (d and e) static powder spectra of $^{15}$N–Phe4 and $^{13}$CO–Val6 labeled TP-I in DLPC membrane (1:15 molar ratio). $^{15}$N spectrum at 303 K (a) and 243 K (b) differ in the CSA. (c) Simulated rigid-limit $^{15}$N powder pattern. (d) $^{13}$CO spectrum of the peptide and the lipids at 303 K. (e) $^{13}$CO spectrum of the peptide after subtracting the lipid background signal. (f) Simulated rigid-limit $^{13}$CO spectrum.

![Fig. 5](image_url)

Fig. 5. TP-I orientation from $^{15}$N and $^{13}$CO chemical shifts. (a) Phe4$^{15}$N chemical shift as a function of ($\theta$, $\phi$) of the bilayer normal in the molecule-fixed PDB system. (b) Val6$^{13}$CO chemical shift as a function of ($\theta$, $\phi$). The measured chemical shifts with the associated uncertainty are colored. (c) The $^{13}$CO and $^{15}$N chemical shifts overlap at ($\theta$, $\phi$) = (78°, 155°). (d) TP-I orientation with the bilayer normal in the vertical direction. The peptide and DLPC bilayers are drawn to scale.
returned the rigid-limit 15N CSA (b and c). The 13CO spectrum at 303 K after subtracting the lipid background signal also shows a uniaxial lineshape, $\delta_{\parallel} = 155$ ppm and $\delta_{\perp} = 205$ ppm. The broadness of the $\delta_{\parallel}$ edge results from insufficient 1H decoupling on the hydrated membrane sample. But the well-defined $\delta_{\perp}$ singularity and Eq. (4) still yield the 0° frequency to ±5 ppm.

To determine TP-I orientation, we calculate the $\delta_{\parallel}$ chemical shifts as a function of $(\theta, \phi)$ of $B_0$ in the molecule-fixed PDB coordinate system [7]. Our recent study of the TP-I conformation indicates that the two strands of the hairpin adopt ideal $\beta$-sheet conformation [9] similar to its solution NMR structure in 60 mM DPC micelles [16]. The 15N–Phe4 and 13CO–Val6 chemical shift surfaces calculated with this conformation are shown in Fig. 5(a and b). The two experimental shifts overlap at a single position, $(\theta, \phi) = (78°, 155°)$, in the entire orientational space (c). This results in a $\beta$-hairpin that is tilted by ~20° from the membrane plane (Fig. 5d). The overall in-plane orientation is consistent with 13C–31P distance measurements indicating that Val6 in the N-terminal strand and Gly10 at the $\beta$-turn are equidistant from the phosphate headgroups, and 1H spin diffusion data indicating that the peptide is not close to the lipid acyl chains of the membrane [9].

4. Conclusion

We have shown by simulation and experiments that in the presence of fast uniaxial rotation, membrane protein orientation can be determined by using unoriented proteoliposomes. Using this approach, we found that the $\beta$-hairpin antimicrobial peptide TP-I is oriented roughly parallel to the plane of the D LPC bilayers.

The use of powder samples for orientation determination opens up many spectroscopic and biological possibilities inaccessible to macroscopically aligned samples. This is the only method for determining the orientation of un-alignable proteins such as curvature-inducing antimicrobial peptides. The removal of glass plates or the need for dilute bicelle solutions increases the sample amount in the rf coil, thus increasing sensitivity. The ease of preparing unoriented proteoliposomes allows direct studies of membrane protein orientation as a function of external parameters such as pH and membrane composition. Finally, with powder samples, one can access the large repertoire of magic-angle spinning (MAS) techniques for site-resolved orientation determination. For example, N–H dipolar couplings can be measured by 2D MAS experiments to determine helix orientation, as we will show elsewhere [17].

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