Conformational mapping of the N-terminal segment of surfactant protein B in lipid using $^{13}$C-enhanced Fourier transform infrared spectroscopy

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Abstract: Synthetic peptides based on the N-terminal domain of human surfactant protein B (SP-B$_{1-25}$; 25 amino acid residues; NH$_2$-FPPIPLPYCWLCRALIKRIQAMIPKG) retain important lung activities of the full-length, 79-residue protein. Here, we used physical techniques to examine the secondary conformation of SP-B$_{1-25}$ in aqueous, lipid and structure-promoting environments. Circular dichroism and conventional, $^{12}$C-Fourier transform infrared (FTIR) spectroscopy each indicated a predominant $\alpha$-helical conformation for SP-B$_{1-25}$ in phosphate-buffered saline, liposomes of 1-palmitoyl-2-oleoyl phosphatidylglycerol and the structure-promoting solvent hexafluoropropanol; FTIR spectra also showed significant $\beta$- and random conformations for peptide in these three environments. In further experiments designed to map secondary structure to specific residues, isotope-enhanced FTIR spectroscopy was performed with 1-palmitoyl-2-oleoyl phosphatidylglycerol liposomes and a suite of SP-B$_{1-25}$ peptides labeled with $^{13}$C-carbonyl groups at either single or multiple sites. Combining these $^{13}$C-enhanced FTIR results with energy minimizations and molecular simulations indicated the following model for SP-B$_{1-25}$ in 1-palmitoyl-2-oleoyl phosphatidylglycerol: $\beta$-sheet (residues 1–6), $\alpha$-helix (residues 8–22) and random (residues 23–25) conformations. Analogous structural motifs are observed in the corresponding homologous N-terminal regions of several proteins that also share the 'saposin-like' (i.e. 5-helix bundle) folding pattern of full-length, human SP-B. In future studies, $^{13}$C-enhanced FTIR spectroscopy and energy minimizations may be of general use in defining backbone conformations at amino acid resolution, particularly for peptides or proteins in membrane environments.
The polypeptide motif of monomeric SP-B, characterized by six Cys residues paired as disulfides and periodic hydrophobic residues, belongs to a class of 'saposin-like' proteins that also includes saposins A–D, sulfated glyco-protein-I, acid sphingomyelinase, acyloxyacyl hydrolase, Entamoeba histolytica pore-forming peptides (amoebapores), plant aspartic proteases and NK-lysin. SP-B, as well as other proteins of the saposin family, uniformly exhibits high levels of α-helix (13, 14, 16–19), suggesting that this motif plays a critical structural role. Based on the crystallographic structure of the related hemerythrin (13, 20) and the 2D-NMR structure of NK-lysin (21), the saposin-fold is apparently characterized by amphipathic α-helices with hydrophilic residues facing the solvent, and hydrophobic side-chains forming a core stabilized by intramolecular disulfide bonds.

Given the critical in vivo role of SP-B in lung function, it is of interest that numerous actions of the full-length protein on surfactant lipids are mimicked by a synthetic peptide representing the N-terminal domain of SP-B (i.e. SP-B<sub>1-35</sub> residues 1–35; Table 1). Both SP-B and positively charged SP-B<sub>1-35</sub> increase the collapse pressure of surfactant lipid monolayers containing palmitic acid; this suggests that the cationic N-terminus of SP-B interacts with anionic lipids to remove the driving force for lipid squeeze-out from the surface film (22, 23). Another indication of specific interactions of proteins with lipids is the insertion of the protein into monolayers at varying surface pressures. The critical insertion pressure, an index of the degree of protein association with lipid films and membrane systems, is very high for both SP-B<sub>1-35</sub> and the parent SP-B (17, 24, 25). Furthermore, full-length SP-B and SP-B<sub>1-35</sub> each similarly induce a coexistence of buckled and flat monolayers when added to surfactant lipids, which may participate in the reduction of surface tension and the enhancement of respreadability of the surfactant monolayer (26). Finally, initial studies indicate that SP-B<sub>1-35</sub> has been shown to mediate lipid-vesicle mixing (27), as does the native SP-B (10, 28, 29).

In view of the likely participation of the N-terminal region of SP-B in key surfactant activities, it is important to elucidate the structure of this domain in lipid environments. Earlier theoretical analysis of the primary sequence of SP-B, using the hydrophobic moment algorithm or Chou-Fasman secondary structure predictions, suggested both a N-terminal β-turn (residues 6–9; Table 1) (30) and an α-helical segment (residues 10–23) (6, 24, 31, 32). Physical experiments on synthetic peptides based on the SP-B N-terminus have also been performed to more directly assign secondary
conformations. Previous circular dichroism (CD) and Fourier transform infrared (FTIR) spectroscopy on SP-B_{1-25} in lipids or solvents that mimic membranes [e.g. methanol, trifluor- oethanol (TFE)] indicated dominant helical content, with minor β- and disordered components [24, 33]. Upon repeating these experiments with a series of truncated SP-B peptides in methanol, a structural model was developed indicating that residues 8–25 (Table 1) were largely α-helix, while residues 1–8 were primarily β-sheet and β-turn [33]. An earlier proton 2D-NMR study of the truncated peptide SP-B_{1-25}, suspended in the membrane mimics methanol or sodium dodecyl sulfate (SDS) similarly demonstrated α-helix for residues 14-21 [34]. However, caution must be employed in extrapolating these structural findings using truncated N-terminal SP-B peptides to the corresponding conformation of SP-B_{1-25} in membrane lipids. Experiments using truncated peptides to localize conformations have inherent uncertainties, not only because shortened peptides may artificially fray at their unproctected N-terminal and C-terminal ends, but also because peptide conformations in membrane mimics (e.g. methanol, SDS) may not faithfully reflect structure in lipid environments. Accordingly, it is important to directly characterize the residue-specific conformation of SP-B_{1-25} in lipids using alternative methodologies.

Here, we study the structure of the N-terminal SP-B_{1-25} peptide in phosphate-buffered saline (PBS), the membrane mimic hexafluorisopropanol (HFIP), or the lipid 1-palmitoyl-2-oleoyl phosphatidylglycerol (POPG) using CD and conventional 13C-FTIR spectroscopy. Additional FTIR experiments were performed using site-directed, 13C-labeled SP-B_{1-25} to assess the residue-specific conformations of peptide bound to POPG liposomes. Previous 13C-labeled FTIR spectroscopic investigations have identified specific random, β-strand and β-turn structural domains in a soluble peptide [35], and also α-helical structure for discrete regions in the transmembrane domain of phospholamban [36]. Here, isotope-labeled SP-B_{1-25} peptides were prepared, in which 13C-carbonyl groups are introduced at various amino acids throughout the peptide. When suites of single and multiply 13C-enhanced SP-B_{1-25} peptides were incorporated into POPG liposomes, the resulting FTIR spectra permitted the mapping of secondary conformation to individual amino acids. POPG liposomes were chosen to provide a homogenous lipid environment for SP-B_{1-25} incorporation not only because POPG is found at relatively high levels (~22%) in lung surfactant preparations, such as Tanaka lipids [33], but also because SP-B_{1-25} is cationic and associates preferentially with the anionic lipids in surfactant dispersions [22, 24, 37].

### Experimental Procedures

#### Materials

Peptide synthesis reagents including Fmoc amino acids and coupling solvents were obtained from Applied Biosystems. Deuterium was supplied by Aldrich Chemical Co. Fmoc 13C-carbonyl leucine, isoleucine, alanine and glycine were purchased from Cambridge Isotope Laboratories. 13C-carbonyl phenylalanine (Cambridge Isotope Laboratories) was converted to the Fmoc derivative by AnaSpec. POPG was obtained from Avanti Polar Lipids. All organic solvents used for sample synthesis, purification and preparation were HPLC grade or better.

#### Solid-phase peptide synthesis, purification and characterization

The native 25 amino acid, N-terminal domain of human surfactant protein B (SP-B_{1-25}; Table 1), and also the 13C-enhanced SP-B_{1-25} analogs (Tables 1 and 2), were prepared using an ABI 431A peptide synthesizer, and purified as described previously [33]. The expected molecular masses of SP-B_{1-25} and SP-B_{1-25} analogs were obtained by fast-atom bombardment and electrospray mass spectrometry (UCLA Center for Molecular and Medical Sciences Mass Spectrometry). Quantitative amino acid compositions for the peptides were determined at the UCLA Protein Microsequencing Facility.
Rationale for $^{13}$C-site directed SP-B$_{1-25}$ substitutions

To probe the secondary conformations within SP-B$_{1-25}$, FTIR spectroscopy was conducted here using two categories of site-directed, isotope-enhanced peptides. The first class involves the incorporation of $^{13}$C-carbonyl groups into multiple, neighboring amino acid residues of synthetic SP-B$_{1-25}$ peptides (Table 1). Separate peptides were synthesized with ‘cassettes’ of multiply $^{13}$C-enhanced substitutions that were staggered to sequentially cover the peptide (Table 1). In earlier studies [35, 36], cassettes of similarly $^{13}$C-enhanced peptides allowed local domain mapping of various secondary conformations (e.g. α-helix, β-sheet, β-turn, random). The rationale behind such experiments is that the secondary structure within a peptide usually extends over more than several adjacent residues [38]. The second class involves peptides substituted with $^{13}$C-carbonyl groups at single amino acid residues (Table 2). FTIR spectroscopic analysis of single-site $^{13}$C-enhanced peptides may assign the secondary conformation to individual residues [36], in cases in which the labeled amino acid exhibits considerable flexibility, use of single-site $^{13}$C-enhanced peptides may indicate multiple conformations that coexist simultaneously.

Most of the isotopically enhanced SP-B$_{1-25}$ peptides in Table 1 [i.e. SP-B$_{1-25}$ [F1/I3/L5], SP-B$_{1-25}$ [L10/A13/L14] and SP-B$_{1-25}$[I18/A20/I21]] and Table 2 [i.e. SP-B$_{1-25}$ [I15], SP-B$_{1-25}$ [I18], SP-B$_{1-25}$ [A20] and SP-B$_{1-25}$ [G25]] share the same chemical composition as the native peptide. Since $^{13}$C-carbonyl Cys is not available commercially, the local region of SP-B$_{1-25}$ at Cys$^8$ and Cys$^{11}$ was probed by substituting these residues with $^{13}$C-carbonyl Ala [i.e. SP-B$_{1-25}$ [A8/A11]] in Table 1 and SP-B$_{1-25}$ [A8] and SP-B$_{1-25}$ [A11] in Table 2. Earlier studies [24] with a SP-B$_{1-25}$ variant, in which Cys$^8$ and Cys$^{11}$ were replaced with alanine residues, showed that these substitutions did not perturb either peptide–lipid binding or the secondary conformation of the peptide when bound to lipids.

Liposome preparation

Large unilamellar vesicles (LUV) of POPG in PBS (500 nmoles lipid/mL) used for CD and FTIR spectroscopic studies were prepared by extrusion through polycarbonate membrane filters [39]. A dry lipid film was hydrated with buffer-saline solution, followed by vortexing of the dispersion to form multilamellar vesicles. The suspension was freeze–thawed five times, and then extruded through 100-nm pore polycarbonate filters (Nuclepore Corp., Pleasanton, CA, USA) five times using a Lipofast device (Avestin Inc., Ottawa, ON, Canada). The size distribution of extruded unilamellar vesicles was determined by dynamic light scattering with a Microtrac 9230 UPA ultrafine particle analyzer (Leeds and Northrup, St. Petersburg, FL, USA) [40]. Extrusion through the 100-nm pore filters yielded a single population of POPG vesicles with a mean diameter of 95.8 nm and a standard deviation of 20.3 nm.

CD Spectroscopy

CD measurements were carried out using an AVIV 62DS spectropolarimeter (AVIV Associates), fitted with a thermoelectric temperature controller that maintained the sample temperature at 25°C [33, 41]. Peptide solutions or peptide/liposome suspensions were measured in 0.1-mm light path demountable cells scanned from 250 to 185 nm at a rate of 10 nm/min and a sample interval of 0.2 nm. The instrument was routinely calibrated with (+)-10-camphorsulfonic acid (1 mg/mL) and a 1-mm path length cell (42), and the ellipticity expressed as the mean residue ellipticity, [θ]MRE (deg cm$^2$/dmol). Peptide sample concentrations were determined from quantitative amino acid analysis (UCLA Microsequencing Facility).

The α-helical structure of SP-B peptides in solvent or lipid environments was determined from CD spectra using the following methodology. The percentage of α-helix conformation in the peptide was first estimated using the
formalism of Chen et al. (43). This approach assumes that the maximum theoretical ellipticity for a given peptide or protein at 222 nm may be derived from the number of amino acid residues (n) and the ellipticity at 222 nm of a helix of infinite length described by equation 1:

\[
\% \text{ helix} = \frac{[\Theta]_{222} \text{ deg cm}^2 \text{ dmol}^{-1} \text{ dmol}^{-1}}{[39 \times 500 (1 - 2.57/n)]} \text{ deg cm}^2 \text{ dmol}^{-1} \text{ dmol}^{-1}
\]

\[(1)\]

**FTIR Spectroscopy**

Infrared spectra were recorded at 25°C using a Mattson Series FTIR spectrometer with a DTGS detector, averaged over 256 scans at a gain of 4 and a resolution of 2 cm⁻¹ (33, 44). No nonlinear baseline corrections, water vapor subtractions, deconvolutions (except for the Fourier self-deconvolutions as noted below in Fig. 3) or smoothing operations were performed on the original spectra. For FTIR spectra of SP-B₁₋₃₅ originally in solvents, peptide self-films were prepared by air drying peptide solutions [in 100% HFIP or aqueous 10 mM phosphate (pH 7.4)] onto 50 × 20 × 2 mm 45° ATR crystal (Spectral Solutions). For measurements with peptides in a lipid environment, SP-B₁₋₃₅ peptides were added to LUV of POPG from the HFIP solution, and allowed to incubate with the liposomes for 1 h. FTIR spectra were measured on the above unchromatographed peptide/liposome mixtures, and also on these peptide/lipid samples after passing through a Sephadex G-50 column to remove nonliposome-associated peptide (45).
Chromatographed and unchromatographed lipid–peptide samples were dried onto the ATR crystal. Both peptide–self-film and lipid–peptide samples were then hydrated for 2 h by passing nitrogen–D₂O vapor through ports in the cell body. Martin et al. [45] noted earlier that the peptide concentration is proportional to the area \( S_{\text{amide}} \) of the amide I band (1680–1600 cm⁻¹), while the lipid concentration is proportional to the area \( S_v\text{[C=O]lipid} \) of the lipid ν(C=O) band (1770–1700 cm⁻¹). Accordingly, the peptide/lipid ratio is proportional to the following: \( S_{\text{amide}}/S_v\text{[C=O]lipid} \).

The amide I bands of conventional \( ^{13}\text{C}-\text{FTIR} \) spectra of peptide self-films and chromatographed lipid/peptide samples were analyzed for the various SP-B₁₋₃₅ conformations [33]. For determinations with peptide/liposomes, the spectrum of the lipid film without peptide (i.e. the band centered at 1730 cm⁻¹) was subtracted from that of samples with peptide associated with lipid. The amounts of α-helix, β-turn, β-sheet and disordered conformations were determined by Fourier self-deconvolutions for band narrowing and area calculations of component peaks determined using curve fitting software supplied by Mattson and based on procedures described by Kauppinen et al. [46]. The frequency limits for the different structures were as follows: α-helix (1662–1645 cm⁻¹), β-sheet (1637–1613 cm⁻¹ and 1689–1682 cm⁻¹), β-turns (1632–1662 cm⁻¹) and disordered or random (1650–1637 cm⁻¹) [47–49].

Enhancement of SP-B₁₋₃₅ peptides with site-specific \( ^{13}\text{C} \)-carbonyl groups permits the direct determination of those amino acid residues participating in secondary conformations. Since the stretching frequencies of the peptide backbone carbonyl groups are sensitive to local conformations, replacement of \( ^{13}\text{C} \) with \( ^{13}\text{C} \) should reduce the stretching frequency of an isolated carbonyl oscillator by \( \approx 37 \text{ cm}^{-1} \) [35, 50, 51]. Specifically, the α-helix band should be lowered to 1635–1608 cm⁻¹, the β-sheet to 1600–1576 cm⁻¹ and 1652–1645 cm⁻¹, β-turns to 1645–1635 cm⁻¹ and disordered or random to 1613–1600 cm⁻¹. These spectral shifts were detected by measuring FTIR spectra of the natural abundance and \( ^{13}\text{C} \)-enhanced peptides in various environments [35, 56], subtle spectral shifts were also detected with difference FTIR spectra, obtained by subtracting the natural abundance spectrum from that of the isotopically enhanced peptide [35]. The difference FTIR spectra should show a negative peak at the original position of the conformational band, and a positive peak shifted from the original by \( \approx 37 \text{ cm}^{-1} \). Unless distorted by noise, the negative peak for a given conformational band should be symmetrical to the positive peak about an inflection point.

### Molecular modeling

The N-terminal peptide of human SP-B, SP-B₁₋₃₅, was modeled using Insight/Discover 97.0 software (Molecular Simulations, San Diego, CA, USA) running on a Silicon Graphics Indigo-2 R10000 High Impact workstation (Beckman Research Institute City of Hope Core Facility). Energy minimizations and molecular dynamics were conducted using the CVFF force field [52, 53] within the Discover software environment. Ramachandran plots were determined for the SP-B₁₋₃₅ structures derived from simulated annealing. The final refined model geometry parameters were evaluated by PROCHECK [54].

### Coordinates

The coordinates for the 10 lowest energy structures of SP-B₁₋₃₅ in POPG, together with a full list of restraints, have been deposited in the Protein Data Bank under the accession code 1DFW.
Results

CD Spectroscopy

To assess the overall conformation of the N-terminal sequence of SP-B (SP-B$_{1-35}$) in a variety of environments, CD spectra were recorded for peptide in an aqueous buffer, a membrane-mimic solvent or a liposome suspension. For example, Fig. 1A shows representative CD spectra of SP-B$_{1-35}$ suspended in either an aqueous environment (i.e. PBS, pH 7.4) or a membrane-mimetic solvent (i.e. HFIP/CH$_2$O with 10 mm phosphate, pH 7.4). The PBS and HFIP spectra in Fig. 1A each demonstrate a characteristic double minimum at 208 and 222 nm, suggesting a considerable $\alpha$-helical content for SP-B$_{1-35}$. The corresponding CD spectrum (Fig. 1B) of SP-B$_{1-35}$ with POPG liposomes in PBS, pH 7.4, after chromatography to remove unbound peptide, similarly indicates a broad double minimum at 208 and 222 nm that is characteristic of high $\alpha$-helix.

It is worthwhile quantifying the SP-B$_{1-35}$ conformation with further analysis of the CD spectra (Fig. 1). The percentage of $\alpha$-helix conformation for the peptide was estimated according to equation 1, as described above (43). Similar elevated proportions of $\alpha$-helix (i.e. 50–66%) were determined for SP-B$_{1-35}$ in the respective PBS, HFIP and POPG environments (Table 3). The relatively high noise for the PBS spectrum in Fig. 1A was attributed to both the limited solubility of amphipathic peptide in aqueous medium and increased light scattering due to the formation of SP-B$_{1-35}$ microaggregates. Previous ESR studies (33) of spin-labeled SP-B$_{1-35}$ in PBS indicated low solubility and elevated aggregation at high peptide concentrations (> 100 $\mu$m), similar to those used here in CD measurements (Fig. 1A). The elevated noise in the POPG spectrum (Fig. 1B) might be a consequence of enhanced light scattering due toSP-B$_{1-35}$-promoted liposome aggregation (27).

Conventional $^{13}$C-FTIR spectroscopy

The secondary conformations of SP-B$_{1-35}$ in the above solvent and lipid systems were then investigated using conventional $^{13}$C-FTIR spectroscopy. Typical FTIR spectra of the amide I band are shown in Fig. 2, and were similar for the three environments. A dominant band occurs at 1658 cm$^{-1}$ for the PBS (Fig. 2A), HFIP (Fig. 2B) and POPG (Fig. 2C) systems, indicative of high $\alpha$-helical content for SP-B$_{1-35}$. Subsequent curve fitting of the POPG spectrum using the criteria of Byler & Susi (47) confirmed a high proportion of SP-B$_{1-35}$ as $\alpha$-helix, with much smaller contributions due to $\beta$-sheet, $\beta$-turn and disordered or random structure (Fig. 3, Table 3). For the 25 amino acid sequence SP-B$_{1-35}$ in POPG, the FTIR analysis in Table 3 suggests that this peptide has $\approx 13$ residues as $\alpha$-helix, $\approx 9$ residues as $\beta$-conformations and $\approx 3$ residues as random structure. Deconvolution of the FTIR spectra of SP-B$_{1-35}$ in either the PBS or the HFIP solvent systems indicated comparable proportions of secondary structure (Table 3).

The association of SP-B$_{1-35}$ with POPG liposomes was quantified by recording $^{13}$C-FTIR spectra, before (not shown) and after chromatographing peptide/lipid mixtures. For POPG incubated with SP-B$_{1-35}$ at an initial calculated peptide/lipid (P/L) ratio of 1:70, the $|S_{amide}|/|S_{4C=O/lipid}|$ ratio for unchromatographed and chromatographed samples indicated > 80% peptide uptake by liposomes. These results are consistent with the relatively low solubility of the peptide in PBS buffer, and the probable strong association between this cationic, amphipathic SP-B$_{1-35}$ and the anionic, polar headgroup region of POPG (see below) (33).

Isotopically enhanced $^{13}$C-FTIR spectroscopy of cassette-labeled SP-B$_{1-35}$ in POPG liposomes

Although the above $^{13}$C-FTIR spectroscopic findings indicate that SP-B$_{1-35}$ is conformationally heterogeneous when bound to POPG liposomes (Fig. 3, Table 3), it is not possible using only this data to assign the secondary conformations (i.e. $\alpha$-helix, $\beta$-turn, $\beta$-sheet, random) to specific residues. To probe local conformations within the peptide, isotopically enhanced, FTIR spectroscopy was performed with SP-B$_{1-35}$ labeled with $^{13}$C-carbonyl groups at multiple sites (Table 1). For peptides incorporated into POPG liposomes, Fig. 4 shows the natural abundance, $^{13}$C-FTIR spectrum of SP-B$_{1-35}$, and the isotopically enhanced, $^{13}$C-FTIR spectra of a suite of cassette-labeled SP-B$_{1-35}$. There are major differences between the native and cassette-labeled SP-B$_{1-35}$ spectra, which are attributed to the presence of $^{13}$C-carbonyl groups.

The putative $\beta$-conformation for the N-terminal region of SP-B$_{1-35}$, predicted theoretically (13, 30) and experimentally using truncated peptides (33), was tested here by comparing the FTIR spectra of native SP-B$_{1-35}$ with that of SP-B$_{1-35}$ [F$_1$/L$_3$/I$_5$] (Table 1). In the SP-B$_{1-35}$ [F$_1$/L$_3$/I$_5$] spectrum (Fig. 4), there is a large decrease in the area 1637–1613 cm$^{-1}$ corresponding to $\beta$-sheet, and a concurrent increase in the area 1550–1576 cm$^{-1}$, indicating an isotopic shift of $\approx 37$ cm$^{-1}$. The nature of this isotopic shift is visualized more readily in a difference FTIR spectrum, obtained by subtracting the native FTIR spectrum (dashed line in Fig. 4A) from that of the SP-B$_{1-35}$ [F$_1$/L$_3$/I$_5$] spectrum (solid line in Fig. 4A); the
Table 3. Proportions of secondary structure* for SP-B<sub>1-25</sub> in solvents and POPG lipid dispersions, as estimated from Fourier self-deconvolution of the FTIR spectra of the peptide amide I band and CD spectra

<table>
<thead>
<tr>
<th>System</th>
<th>% Conformation α-Helix</th>
<th>β-Sheet</th>
<th>β-Turn</th>
<th>Disordered</th>
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<tr>
<td>FTIR Spectra&lt;sup&gt;a&lt;/sup&gt;</td>
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<tr>
<td>PBS&lt;sup&gt;&lt;sub&gt;5&lt;/sub&gt;&lt;/sup&gt;</td>
<td>50</td>
<td>10</td>
<td>32</td>
<td>8</td>
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<tr>
<td>HFIP&lt;sup&gt;&lt;sub&gt;6&lt;/sub&gt;&lt;/sup&gt;</td>
<td>54</td>
<td>7</td>
<td>25</td>
<td>14</td>
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<tr>
<td>POPG&lt;sup&gt;&lt;sub&gt;7&lt;/sub&gt;&lt;/sup&gt;</td>
<td>52</td>
<td>12</td>
<td>24</td>
<td>12</td>
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<tr>
<td>CD Spectra&lt;sup&gt;a&lt;/sup&gt;</td>
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<td>PBS&lt;sup&gt;&lt;sub&gt;5&lt;/sub&gt;&lt;/sup&gt;</td>
<td>50</td>
<td>ND&lt;sup&gt;&lt;sub&gt;8&lt;/sub&gt;&lt;/sup&gt;</td>
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<tr>
<td>HFIP&lt;sup&gt;&lt;sub&gt;6&lt;/sub&gt;&lt;/sup&gt;</td>
<td>66</td>
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<td>ND</td>
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<tr>
<td>POPG&lt;sup&gt;&lt;sub&gt;7&lt;/sub&gt;&lt;/sup&gt;</td>
<td>53</td>
<td>ND</td>
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</table>

a. Data are the means of four separate determinations and have an SE ± 5% or better. b. FTIR spectra were deconvoluted as described in Experimental procedures (Fig. 3). c. SP-B<sub>1-25</sub> (341 µM) dried on to the ATR from PBS, pH 7.4. (Fig. 2A). d. SP-B<sub>1-25</sub> (141 µM) dried on to an ATR plate from hexafluoropropanol/HFIP/water (7:2:3, v/v) in 10 µM phosphate, pH 7.4. (Fig. 3B). e. Chromatographed SP-B<sub>1-25</sub>/POPG liposomes (initial peptide/lipid molar ratio 1:70) in PBS, pH 7.4, dried on to an ATR plate (Fig. 2C). f. CD spectra were analyzed with the method of Chen et al. (43) (Fig. 1). g. POPG<sub>1</sub> suspended in phosphate buffered saline, pH 7.4 at 225 µM (Fig. 1A). h. SP-B<sub>1-25</sub> in HFIP/water (7:3, v/v) with 10 µM phosphate, pH 7.4 at 173 µM (Fig. 1A). i. Chromatographed SP-B<sub>1-25</sub>/POPG (initial peptide/lipid molar ratio 1:70) in PBS, pH 7.4 (Fig. 1B). j. ND, not determined.

The corresponding FTIR spectral results of another cassette, SP-B<sub>1-25</sub> (I8/A20/L22), provide some evidence that the α-helix terminus at Ile<sup>22</sup>. For the SP-B<sub>1-25</sub> (I8/A20/L22) spectrum in Fig. 4D, the band at 1662–1645 cm<sup>-1</sup> is reduced with respect to the native spectrum, with a new band arising concurrently at 1630–1603 cm<sup>-1</sup>, indicating an isotopic shift of 42 cm<sup>-1</sup>. The corresponding FTIR spectrum for SP-B<sub>1-25</sub> (I8/A20/L22) in Fig. 5D confirms the presence of negative and positive bands centered at 1658 and 1615 cm<sup>-1</sup>, respectively. One possible interpretation is that only one of the <sup>13</sup>C-carbonyls of SP-B<sub>1-25</sub> (I8/A20/L22) participates in the α-helix; this would be consistent with the relatively modest decrease in the α-helical band at 1662–1645 cm<sup>-1</sup> and the much larger isotopic shift for the new peak (Fig. 4D) than those of SP-B<sub>1-25</sub> (L10/A13/L14) (Fig. 4C). Here, the positive band centered at 1615 cm<sup>-1</sup> (Figs 4D and 5D) would itself be a composite, because of the <sup>13</sup>C-carbonyls of Ile<sup>18</sup>, Ala<sup>20</sup> and Ile<sup>22</sup> of SP-B<sub>1-25</sub> (I8/A20/L22) assuming α-helical and other conformations. In this context, it is important to note that the helix-breaking Pro<sup>13</sup> prevents the carbonyl of Ala<sup>20</sup> hydrogen bonding to Lys<sup>94</sup> as an α-helix. Accordingly, the <sup>13</sup>C-carbonyl of Ile<sup>18</sup> may participate in the α-helix by bonding to the amide hydrogen of Ile<sup>22</sup>, whereas the corresponding carbonyls of Ala<sup>20</sup> and Ile<sup>22</sup> assume other conformations (Fig. 6). The only α-helical involvement for the Ala<sup>20</sup> and Ile<sup>22</sup> residues would be through donation of amide hydrogens to their respective carbonyl partners, Lys<sup>94</sup> and Ile<sup>18</sup> (see below).

Isotopically enhanced <sup>13</sup>C-FTIR spectroscopy of single residue-labeled SP-B<sub>1-25</sub> in POPG liposomes

The local conformations within the peptide were also probed using FTIR spectroscopy of SP-B<sub>1-25</sub> labeled with <sup>13</sup>C-carbonyl groups at single sites (Table 2). The use of such labeled peptides allows study of whether a given residue is flexible enough to exhibit multiple conformations. Single-site <sup>13</sup>C-carbonyl-labeled peptides are particularly advantageous in assessing α-helical domains. For a residue (N)

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whose carbonyl group forms an α-helical C=O···H–N hydrogen bonding with its residue partner (N + 4), the FTIR assignment of an α-helix to the 13C-carbonyl N-residue also confirms α-helical conformations for four consecutive residues: N + 1, N + 2, N + 3 and N + 4 [36].

The conformation of the C-terminus of SP-B1-35 bound to POPG liposomes was studied by comparing the FTIR spectrum of native SP-B1-35 with that of SP-B1-35 [G25; Table 2]. In the SP-B1-35 [G25] spectrum [Fig. 7C], there is a small decrease in the area (1650–1637 cm⁻¹) corresponding to random structure, as well as an increase in the area 1613–1600 cm⁻¹, indicating an isotopic shift of ~37 cm⁻¹. This isotopic shift is seen more clearly in the corresponding difference FTIR spectrum [Fig. 7D], which confirms negative and positive bands centered at 1647 cm⁻¹ and 1606 cm⁻¹, respectively. The results suggest that, in the native SP-B1-35 spectrum, residue Gly³⁵ contributes to the random structural band identified in the Fourier deconvoluted spectrum at 1650–1637 cm⁻¹ [Figs 3 and 6].

The α-helical structure for SP-B1-35, tentatively assigned in Fig 6 from FTIR spectroscopy of cassette 13C-labeled peptides, was examined further using single-site 13C-carbonyl-labeled SP-B1-35 peptides [Table 2]. Residue 8 of SP-B1-35 was investigated by recording the FTIR spectrum of SP-B1-35 [A8] incorporated into POPG liposomes. Both the FTIR spectrum [Fig. 8A] and the difference FTIR spectrum [Fig. 8B] of SP-B1-35 [A8] show that the major conformation (Fig. 7B) at position 8 (Cys⁸ in native peptide) is α-helical, as indicated by the additional positive band at 1618 cm⁻¹; the small, second positive band at 1637 cm⁻¹ in the difference FTIR spectrum [Fig. 8B] also reflects a minor β-turn element for position 8. Apparently, position 8 for SP-B1-35, is conformationally flexible, assuming primarily an α-helical conformation with a minor β-turn element [Fig. 6]. These experiments were repeated with additional single-site 13C-carbonyl-enhanced SP-B1-35 peptides to determine the conformation of positions 11, 15 and 18 [Table 2]. The respective FTIR spectra and difference FTIR spectra for
α-helical conformation for Ile^{18} (i.e. a positive peak at 1618 cm^{-1} in Fig. 9B), while the corresponding difference FTIR spectrum for SP-B_{1-25} [A20] reports β-turn and random conformations for Ala^{20} (i.e. peaks at 1627 and 1608 cm^{-1} in Fig. 9D). These results confirm our preliminary assignment that the FTIR spectrum of SP-B_{1-25} [I18/A20/I22] is a composite, arising from the carbonyl of Ile^{18} participating in an α-helix (through bonding with the amide H of Ile^{23}) with the carbonyls of Ala^{20} and Ile^{22} assuming other conformations. The finding of multiple conformations for the carbonyl groups at the ends of the α-helix for SP-B_{1-25} [Fig. 6] suggests some fraying of the helix at the N- and C-termini.

Molecular modeling

Although the above $^{13}$C-FTIR spectroscopic analysis assigned the conformation of the majority (> 75%) of the amino acids for SP-B_{1-25} in POPG liposomes, the conformations of a number of residues (i.e. Pro^{3}, Pro^{6}, Pro^{7}, Tyr^{7}, Pro^{15}, Lys^{24}) were not determined experimentally. Molecular modeling was conducted to provide structural information on these 'gap' residues, and also to assess the conformational flexibility of domains within SP-B_{1-25}. From the $^{13}$C-FTIR spectral results in Fig. 6, the initial structure was modeled as an α-helix, except for the terminal regions (residues 1–7 and 22–25), which were modeled as β-strands using Discover software. Ten cycles of simulated annealing were performed, using a protocol published earlier (55), to explore those conformations accessible to the terminal regions and helix side-chains. Throughout the simulation, constraints were applied to the helix backbone so that the structure would remain consistent with data derived from $^{13}$C-FTIR spectroscopic mapping studies. Using a time step of 0.5 fs, the system was equilibrated for 1000 iterations at 300 K. The temperature was then increased to 900 K in increments of 30 K over a period of 20 ps. Molecular dynamics was then carried out at 900 K for 30 ps, followed by annealing to a temperature of 300 K over 50 ps. The resulting structure was then energy minimized (52, 53) using 5000 steps, followed by 10 000 steps of conjugate gradients until the maximum derivative was < 0.001. Throughout the simulation, the backbone atoms of residues 8–22 were constrained using a force constant that increased by 100 [kCal/mol × Å²] during each heating step. A similar torsional constraint was applied to all peptide bonds to maintain them in the trans conformation.

With the residue-specific constraints based on $^{13}$C-FTIR measurements described above, 10 molecular models
Figure 6. Conformational map of the N-terminal peptide of SP-B (SP-B_{1-35}) in the lipid POPG, as estimated from the FTIR spectra of \(^{13}\text{C}\)-labeled peptides. Amino acids are represented by three-letter codes. Peptides, at a P/L ratio of 1:70, were added to POPG liposomes suspended in PBS. Codes (in parentheses) for peptide conformations are: \(\beta\)-sheet (\(\beta\)), \(\beta\)-turn (\(\beta\-T\)), \(\alpha\)-helix (\(\alpha\)) and random (\(R\)). Secondary conformations were determined for SP-B_{1-35} peptides labeled with \(^{13}\text{C}\) at multiple sites (Table 1) by recording the FTIR spectra of SP-B_{1-35} \([\text{F1/F5/L5}]\) in Fig. 4A, SP-B_{1-35} \([\text{A8/A11}]\) in Fig. 4B, SP-B_{1-35} \([\text{L10/A13/L14}]\) in Fig. 4C and SP-B_{1-35} \([\text{L16/A30/123}]\) in Fig. 4D. Secondary conformations were also determined for SP-B_{1-35} labeled with \(^{13}\text{C}\) at single sites (Table 3) by recording the FTIR spectra shown in Figs 7–9.

Representing possible lipid conformations of SP-B_{1-35} were generated by simulated annealing molecular dynamics. The superimposed conformers are shown as stick figure representations in Fig. 10A, whereas a ribbon representation of a selected conformer showing the location of each residue is indicated in Fig. 10B. The geometry parameters of the final conformer models were evaluated using PROCHECK [54]. The Ramachandran plot in Fig. 11 verifies that the backbone torsion angles for all 10 conformers fell within the allowed regions. Furthermore, the Ramachandran plot indicates high levels of backbone torsion angles lying in the \(\alpha\)-helix and \(\beta\)-structural zones, consistent with the structure of SP-B_{1-35} principally subdivided into \(\alpha\)-helix and \(\beta\)-conformations (Fig. 11). The superimposed conformers capture the major structural features indicated by our \(^{13}\text{C}\)-FTIR spectra, namely an \(\alpha\)-helix (residues 8–23) capped with \(\text{Pro}^6\) and \(\text{Pro}^{33}\), a \(\beta\)-conformation at the N-terminal sequence (residues 1–6) and random structure at the C-terminus (residues 23–25). Within the helix, several alternative packing arrangements are accessible to the side-chains. It is interesting to note that, in the modeled structures of Fig. 10A, the side-chain of Tyr\(^7\) often packs against the side-chains of the i + 3 residue [Leu\(^{10}\)], an interaction that is believed to be helix stabilizing [56]. In our models, Tyr\(^7\) is the residue that appears to initiate the helix, its carbonyl group being the first to participate in the helix hydrogen-bonding network. The \(\beta\)-conformation structure for the N-terminal sequence (residues 1–6) exhibits considerable flexibility in Fig. 10A, evincing some ability to fold on itself as an antiparallel \(\beta\)-sheet. Since the \(\alpha\)-helix was constrained as an initial condition throughout, one important limitation of this modeling exercise is that it is unable to demonstrate any fraying at the two ends of the \(\alpha\)-helix.

The pleiomorphism that was noted at the ends of the \(\alpha\)-helix in Fig. 6 may be due to the lack of hydrogen bonding that occurs at N- and C-terminal caps.

Discussion

The residue-specific conformational model for SP-B_{1-35} in lipid [Figs 6 and 10], developed here using \(^{13}\text{C}\)-FTIR spectroscopy and molecular modeling, is broadly in agreement with previous theoretical predictions and physical measurements on the full-length (35 residue) peptide. Consistent with earlier Chou–Fasman predictions of secondary structure [30, 32], \(\beta\)-conformations were concentrated in the N-terminal region with an \(\alpha\)-helix in the C-terminal domain [Figs 6 and 10]. It is also important to compare the \(\alpha\)-helical content reported here from CD spectroscopy of SP-B_{1-35} in POPG liposomes [Fig. 1B, Table 3] with that obtained from \(^{13}\text{C}\)-FTIR spectroscopy [Figs 6 and 10B]. Both CD and site-directed \(^{13}\text{C}\)-FTIR spectroscopic analyses indicate similar \(\alpha\)-helical content (i.e. 53 and 60%, respectively) for SP-B_{1-35} in POPG liposomes. These results are of particular interest, given that CD spectroscopy used continuously hydrated peptide/liposomes, whereas ATR-FTIR spectroscopy subjected samples to a drying and rehydration step. Our finding of similar total \(\alpha\)-helical content from these two techniques argues against the drying and rehydration step introducing significant experimental artifacts into the \(^{13}\text{C}\)-FTIR spectral analysis. In agreement with this are our observations that dried and rehydrated FTIR samples possess relatively high levels of hydration. Okamura et al. [57] previously showed that the hydration state of DPPC multilayers could be determined using FTIR spectroscopy, by noting that the
Figure 7. FTIR and difference FTIR spectra of the amide I band for \(^{13}\)C-carbonyl SP-B\(_{1-25}\) [native] and single \(^{13}\)C-carbonyl-enhanced SP-B\(_{1-25}\) peptides labeled at Ile\(^{15}\) [i.e. SP-B\(_{1-25}\) (I15)] or Gly\(^{15}\) [i.e. SP-B\(_{1-25}\) (G25)]. Peptides at an initial P/L ratio of 1:70 were added to POPG liposomes suspended in PBS, pH 7.4. Spectra were recorded at 25°C on chromatographed peptide/liposomes that were dried on the ATR and hydrated with D\(_2\)O for 3 h. (A) SP-B\(_{1-25}\) (I15) is the solid line and native SP-B\(_{1-25}\) is the dashed line. The minor peak at 1618 cm\(^{-1}\) in the SP-B\(_{1-25}\) (I15) spectrum indicates an \(\alpha\)-helix for Ile\(^{15}\). (B) SP-B\(_{1-25}\) spectrum [dashed line in A] from SP-B\(_{1-25}\) (I15) [solid line in A]. The peak at 1618 cm\(^{-1}\) [arrow] indicates an \(\alpha\)-helix for Ile\(^{15}\). (C) SP-B\(_{1-25}\) (G25) is the solid line and native SP-B\(_{1-25}\) is the dashed line. The minor peak at 1606 cm\(^{-1}\) in the SP-B\(_{1-25}\) (G25) spectrum indicates a random conformation for Gly\(^{15}\). (D) SP-B\(_{1-25}\) spectrum [dashed line in C] from SP-B\(_{1-25}\) (G15) [solid line in C]. The peak at 1606 cm\(^{-1}\) [arrow] indicates a random conformation for Gly\(^{15}\). See Table 1 for amino acid sequences of peptides. Spectra and peak assignments are representative of three independent experiments.

The intensity ratio of the OH stretching band of water (at 3380 cm\(^{-1}\)) to the CH\(_2\) band of lipid (at 2938 cm\(^{-1}\)) is proportional to the hydration state (% weight). Following this approach, our FTIR samples rehydrated for 2 h indicated substantial D\(_2\)O content, with an intensity ratio of 2.5 for the OD stretching band of D\(_2\)O (at 2493 cm\(^{-1}\)) to that of the CH\(_2\) band of lipid (at 2925 cm\(^{-1}\)) (spectra not shown). Also in support of our view that the drying and rehydration step does not introduce errors, Goormaghtigh et al. [58] reported no differences for FTIR spectra of selected proteins recorded either by transmission in solution or by drying and rehydration on the ATR crystal.

It is also important to note that the proportions of secondary conformation determined by Fourier self-deconvolution of \(^{13}\)C-FTIR spectra of SP-B\(_{1-25}\) in POPG liposomes (i.e. \(\approx 13\) residues as \(\alpha\)-helix, \(\approx 9\) residues as \(\beta\)-conformations and \(\approx 3\) residues as random structure; Fig. 3; Table 3) are in good agreement with those determined from site-directed \(^{13}\)C-FTIR spectroscopy and modeling (\(\approx 13\) residues as \(\alpha\)-helix, \(\approx 7\) residues as \(\beta\)-conformations and \(\approx 3\) residues as random; Figs 6 and 10). Fourier self-deconvolution of FTIR spectra is a standard technique used to estimate the proportions of secondary structure in peptides and proteins [47, 49]. With this computer-fitting methodology, the integrated intensities of the component band are proportional to the populations of the conformational structures represented by these components. The relative percentages of secondary structures may be determined by adding the areas of all bands assigned to each of these structural elements, and then expressing the sum as a fraction of the total amide I band area (48). Since Fourier self-deconvolution assumes that the intrinsic absorption of each structural component will be similar in all peptides and proteins, Surewicz & Mantsch [48] believed that the percent conformations derived with this approach should be viewed as estimates only. Despite these uncertainties, however, good correlations were reported for a wide range of proteins between the secondary structures determined from X-ray diffraction and Fourier self-deconvolution of FTIR spectra [47, 49]. Here, the percent secondary structures were assessed for SP-B\(_{1-25}\) in POPG liposomes using both Fourier self-deconvolution (Fig. 3; Table 3) and \(^{13}\)C-FTIR spectroscopy (Figs 6 and 10B). Such comparisons are a further independent test of the Fourier self-deconvolution technique, because the experimental conditions for Fourier self-deconvolution and site-directed \(^{13}\)C-FTIR spectroscopy are virtually identical (except for the use of \(^{13}\)C-labeled peptides), and no assumptions are made in \(^{13}\)C-FTIR
analyses about the intrinsic absorptions of secondary structures. It is of interest, then, that the proportions of secondary conformation determined by Fourier self-deconvolution are in good agreement with those obtained from site-directed $^{13}$C-FTIR analysis (see above). Although our $^{13}$C-FTIR findings support the use of Fourier self-deconvolution in this one instance, future comparisons should be performed on a much wider suite of proteins and peptides. Also, it should be emphasized that, unlike Fourier self-deconvolution, site-directed $^{13}$C-FTIR measurements provide additional information on the actual location of secondary conformations.

The residue-specific, conformational model for SP-B$_{1-25}$ in POPG lipid (Figs 6 and 10) may be further compared with previous models based on truncated SP-B$_{1-25}$ peptides in membrane-mimic environments (33, 34). The local conformations for SP-B$_{1-25}$ in POPG may be summarized as follows: β-sheet (residues 1–6), α-helix (residues 8–22) and random (residues 23–25). Using difference $^{13}$C-FTIR spectroscopy of SP-B$_{1-25}$ labeled with $^{13}$C-carbonyl at single sites (Figs 8A, B and 9C, D), multiple conformations were also observed at each end of the α-helix (i.e., α-helix and β-turn for the Cys8 position, random and β-turn for Ala20; see Fig. 6); this demonstrates some fraying of the helix at the N- and C-termini. Additional conformational flexibility was noted in the β-sheet of residues 1–6, from examination of the conformers determined with energy minimization (Fig. 10A). Earlier physical measurements with truncated peptides (33, 34) confirm this overall pattern of secondary structure in SP-B$_{1-25}$. For example, prior CD and conventional $^{13}$C-FTIR spectroscopy on truncated SP-B$_{1-25}$ peptides [e.g., SP-B$_{1-20}$ and SP-B$_{1-21}$] in structure-promoting methanol, indicated elevated β-conformations in the N-terminal segment Phe1 to Cys8, with high α-helix in the C-terminal segment Cys8 to Gly25 (33). Moreover, 2D-NMR analysis of another shortened N-terminal peptide (SP-B$_{1-21}$) in methanol or SDS demonstrated α-helix for residues 14–21 (34). Although these physical studies with shortened peptides support the present $^{13}$C-FTIR results that the β-sheet localizes in the N-terminal portion and the α-helix in the C-terminal region, the two approaches show differences in the precise location of the major secondary elements. These
Figure 9. FTIR and difference FTIR spectra of the amide I band for $^{13}$C-carbonyl SP-B$_{1-35}$ (native) and single $^{13}$C-carbonyl enhanced SP-B$_{1-35}$ peptides labeled at Ile$^{18}$ [i.e. SP-B$_{1-35}$ (I18)] or Ala$^{20}$ [i.e. SP-B$_{1-35}$ (A20)]. Peptides at an initial P/L ratio of 1:70 were added to POPG liposomes suspended in PBS, pH 7.4. Spectra were recorded at 35°C on chromatographed peptide/liposomes that were dried on the ATR and hydrated with D$_2$O for 2 h. [A] SP-B$_{1-35}$ (I18) is the solid line and native SP-B$_{1-35}$ is the dashed line. The minor peak at 1618 cm$^{-1}$ in the SP-B$_{1-35}$ (I18) spectrum indicates an α-helix contribution for Ile$^{18}$. [B] SP-B$_{1-35}$ spectrum (dashed line in A) from SP-B$_{1-35}$ (I18) [solid line in A]. The peak at 1618 cm$^{-1}$ indicates an α-helix for Ile$^{18}$. [C] SP-B$_{1-35}$ (A20) is the solid line and native SP-B$_{1-35}$ is the dashed line. The minor peak at 1627 cm$^{-1}$ in the SP-B$_{1-35}$ (A20) spectrum indicates a β-turn conformation for Ala$^{20}$. [D] SP-B$_{1-35}$ spectrum (dashed line in C) from SP-B$_{1-35}$ (A20) [solid line in C]. The peaks at 1637 and 1608 cm$^{-1}$ (arrows) indicate that Ala$^{20}$ exhibits both β-turn and random conformations, the loss of some symmetry for the negative and positive bands in [D] is attributed to a relatively high noise, which distorted the shape of the negative band. See Table 2 for amino acid sequences of peptides. Spectra and peak assignments are representative of three independent experiments.

Discrepancies are attributed to the inherent uncertainties of working with truncated peptides that may fray at their unprotected N-terminal and C-terminal ends. Accurate assignment of SP-B$_{1-35}$ conformations using truncated peptides requires that the fragments retain the same conformation as in the full-length peptide, and this is probably valid only as a first approximation. The conformation model developed here for SP-B$_{1-35}$ in POPG lipid using $^{13}$C-FTIR spectroscopy and energy minimizations [Figs 6 and 10] is more accurate because measurements are made using the full-length SP-B$_{1-35}$ peptide containing nonperturbing, $^{13}$C-carbonyl substituted amino acids. Moreover, $^{13}$C-FTIR spectroscopy permits direct conformational measurements for peptide in actual membrane lipids, whereas the above studies with truncated peptides were performed using only membrane mimics (e.g. methanol and SDS).

It is worthwhile determining whether our structural motif for SP-B$_{1-35}$ (Figs 6 and 10) is also represented in the corresponding domains of homologous proteins. As noted above, sequence and structural homologies indicate that the 79-residue, full-length SP-B probably assumes a 'saposin-fold', characterized by amphipathic α-helices forming a core stabilized by multiple intramolecular disulfide bonds [13–15, 21]. The structural homologies for SP-B and other saposins may extend beyond the overall folding pattern to the local domain elements. For example, the N-terminal segment of SP-B$_{1-35}$ [residues 1–9], with multiple proline residues interspersed with hydrophobic residues, exhibits striking sequence similarities with the corresponding region of hemerythrin [13]. It should be noted, then, that the N-terminal domains of SP-B$_{1-35}$ [residues 1–6; Figs 6 and 10] and hemerythrin [residues 1–9] [13, 20] each form β-sheet conformations. However, X-ray crystallographic analysis of hemerythrin [20] indicates that the N-terminal domain [residues 1–9] folds as an antiparallel β-sheet, whereas molecular modeling of the corresponding N-terminal region of SP-B$_{1-35}$ [residues 1–6] shows an extended β-sheet with considerable flexibility, but only limited ability to form an antiparallel conformation. The N-terminal segment of SP-B$_{1-35}$ [residues 1–9] was earlier proposed to act as a sort of
'biochemical Velcro,' facilitating in vivo either the aggregation of SP-B or the interactions of SP-B with surfactant protein C (SP-C) [30]. The self-association of SP-B molecules forms protein-rich networks in lipids that may well be critical in mediating the surfactant activity of this protein [23, 26, 59]. Our structural model for SP-B_{1-25} (Fig. 10) raises the possibility that SP-B molecules cross-link through complementary hydrogen bonding of opposing N-terminal segments, each assuming an extended β-sheet conformation.

The α-helix observed here for SP-B_{1-25} (residues 8–22) [Figs 6 and 10] is probably a ubiquitous feature in the N-terminal region of proteins sharing the 'saposin-fold.' The 3D structure of NK-lysin has been elucidated in a recent 2D-NMR study, and established an overall saposin folding pattern for the soluble protein [21]. Based on the known sequence similarities between SP-B and NK-lysin and the 2D-NMR spectral finding of an α-helix for the N-terminus [residues 3–18] of NK-lysin, Liepinsh et al. [21] predicted that residues 7–22 of SP-B would also assume an α-helical conformation. Our experimental 13C-FTIR results indicating α-helix for residues 8–22 of SP-B_{1-25} (Figs 6 and 10) provide dramatic confirmation of this prediction and further suggest α-helical conformations for the corresponding N-terminal domains of other saposin proteins, such as saposins A–D, sulfated glycoprotein-I, acid sphingomyelinase, acyloxyacyl hydrolase and E. histolytica pore-forming peptides (amoebapore) [10, 13–15, 19, 21]. Although the functional role of this N-terminal α-helical domain remains to be determined in other saposin proteins, prior work has established that this amphipathic α-helical motif is an important participant in the surfactant activities of SP-B_{1-25} [24, 25, 32].
The results obtained here on SP-B_{1-25} confirm that the joint use of $^{13}$C-FTIR spectroscopy and energy minimizations will be of general utility in mapping the secondary structures of conformationally heterogeneous peptides and proteins. Past $^{13}$C-FTIR spectroscopic investigations have identified the sites of random, $\beta$-strand and $\beta$-turn structural domains in a soluble peptide (35), and also $\alpha$-helical structure at specific regions in the transmembrane domain of phospholamban (36). Here, the residue-specific structure of the conformationally rich SP-B_{1-25} peptide bound to lipids has been elucidated using $^{13}$C-FTIR spectroscopy and molecular simulations, such that the discrete locations of $\alpha$-helix, $\beta$-sheet, random conformations have been characterized (Fig. 10). Furthermore, FTIR spectroscopic analysis of peptides substituted at single residues with $^{13}$C-labeled carbonyls offers the potential to assess those conformationally flexible amino acids that assume multiple secondary structures (Fig. 6). We anticipate that future $^{13}$C-FTIR spectroscopic and molecular modeling experiments may well yield complete conformational maps for other membrane-bound proteins (or peptides), which have been difficult to analyze with more standard methodologies (i.e. X-ray crystallographic and 2D-NMR spectroscopy). In future $^{13}$C-FTIR spectroscopic investigations, it would be worthwhile studying whether the conformational properties of the SP-B_{1-25} peptide are the same as when this sequence is part of the full-length SP-B protein. Despite the similar surfactant activities reported for SP-B_{1-25} and full-length SP-B (22, 23, 26, 37, 59), it is unclear whether the isolated SP-B_{1-25} peptide accurately represents the structure of this domain in the full-length protein (10). This problem may be addressed by repeating the $^{13}$C-FTIR experiments presented here with our $^{13}$C-labeled SP-B_{1-25} peptides, each incorporated into the full-length (i.e. 79-residue), synthetic SP-B molecule (23, 59).

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