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Analysis of protein–protein and protein–membrane interactions by isotope-edited infrared spectroscopy

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The objective of this work is to highlight the power of isotope-edited Fourier transform infrared (FTIR) spectroscopy in resolving important problems encountered in biochemistry, biophysics, and biomedical research, focusing on protein–protein and protein membrane interactions that play key roles in practically all life processes. An overview of the effects of isotope substitutions in (bio)molecules on spectral frequencies and intensities is given. Data are presented demonstrating how isotope-labeled proteins and/or lipids can be used to elucidate enzymatic mechanisms, the mode of membrane binding of peripheral proteins, regulation of membrane protein function, protein aggregation, and local and global structural changes in proteins during functional transitions. The use of polarized attenuated total reflection FTIR spectroscopy to identify the spatial orientation and the secondary structure of a membrane-bound interfacial enzyme and the mode of lipid hydrolysis is described. Methods of production of site-directed, segmental, and domain-specific labeling of proteins by the synthetic, semisynthetic, and recombinant strategies, including advanced protein engineering technologies such as nonsense suppression and frameshift quadruplet codons are overviewed.

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1. Introduction

Protein–protein interactions play a crucial role in nearly all aspects of cell physiology, such as cell signaling, reception, immunology, metabolism, muscle contraction, cell adhesion *etc.*^{1–3} Moreover, defects in protein–protein interactions constitute the core cause of many diseases.^{4–6} On the other hand, interactions of proteins with cellular membranes are central to many vital processes such as transport and homeostasis of ions and molecules, endo- and exocytosis, interfacial enzymology, *etc.*^{7–9} Therefore, development of capable methods of analysis and characterization of such interactions is of pivotal importance for understanding the intricate molecular machinery driving life processes. Apart from the atomic- or near-atomic-resolution structural techniques like X-ray crystallography, nuclear magnetic resonance, cryo-electron microscopy and tomography,^{10–12} single molecule imaging,^{13,14} and advanced versions of atomic force microscopy,^{13–16} other methods that are conventionally considered low-resolution comprise incompletely appreciated structure-resolving power. One of such techniques is Fourier transform infrared (FTIR) spectroscopy, including its various versions such as polarized attenuated total reflection (ATR) and isotope-edited FTIR.

FTIR is vibrational spectroscopy that can be described with rather high fidelity using the harmonic oscillator (HO) approach (see the next section), according to which the vibrational frequency (ν) of a diatomic molecule of atomic masses of m_1 and m_2 is:¹⁷

$$\nu = \frac{1}{2\pi} \sqrt{\frac{k}{\mu}} = \frac{1}{2\pi} \sqrt{k \left(\frac{1}{m_1} + \frac{1}{m_2} \right)} \quad (1)$$

In eqn (1), μ is the reduced mass of the two atoms: $\mu = m_1 m_2 / (m_1 + m_2)$. According to eqn (1), ν is sensitive not only to the types of the two atoms with specific atomic masses, such as C and O, but also to the strength of the covalent bond between them (k), such as a single or a double bond. Moreover, hydrogen bonding (H-bonding) of the oxygen weakens the C=O covalent bond and thereby reduces the vibrational frequency, allowing identification of the presence and the strength of H-bonding that stabilizes protein structure. Furthermore, replacement of one or both atoms with a heavier isotope will generate a downshifted vibrational frequency, permitting analysis of a protein's local structure and dynamics, as described in the forthcoming sections.

The multiple vibrational modes of the amide group of proteins, *i.e.*, amide I through amide VII as well as the amide A and B modes, generate absorbance bands at specific mid-infrared frequencies that are sensitive to the protein's secondary and tertiary

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structure.¹⁸ The amide I mode (1700–1600 cm⁻¹) is mostly due to the C=O stretching vibration, with small contributions from the C–N, C–C–N, and N–H vibrations. Its frequency depends on the local geometry of the protein backbone and interactions between chemical groups such as H-bonding between the main-chain C=O and N–H groups and through space vibrational coupling effects. Hence, the amide I band is exquisitely sensitive to the protein secondary structure. Due to the spectral overlap between the H₂O bending mode and the protein amide I mode, a D₂O-based buffer is usually used in FTIR experiments. The use of D₂O results in a slight red shift of amide I frequencies caused by amide hydrogen/deuterium exchange (HX). The main secondary structures of a protein, such as various β- and γ-turns, α-helix, β-sheet, irregular structure, can be pinpointed based on their characteristic amide I wavenumbers ($W = 1/\lambda = \nu/c$, where λ is the wavelength and c is the speed of light). The approximate wavenumber ranges for proteins in H₂O (and D₂O, shown in parentheses) are as follows: 1700–1660 cm⁻¹ (1690–1650 cm⁻¹) for γ-turns, 1685–1655 cm⁻¹ (1675–1640 cm⁻¹) for β-turns, 1660–1648 cm⁻¹ (1655–1638 cm⁻¹) for α-helix, 1660–1652 cm⁻¹ (1648–1640 cm⁻¹) for irregular structure, and 1638–1630 cm⁻¹ (1635–1625 cm⁻¹) for β-sheet. Moreover, various subtypes of same secondary structure can be distinguished. For example, the α_{II}-helix, which has identical geometric parameters with the regular α-helix (rise per amino acid along the helical axis, number of amino acids per turn) but slightly tilted amide plane and hence weaker C=O···H–N H-bonding and stronger C=O and H–N covalent bonds, generates amide I signal at higher wavenumbers, ~1665 cm⁻¹, making FTIR a unique tool for detection of this structure. FTIR is also uniquely suited to distinguish between parallel and antiparallel and between intramolecular and intermolecular β-sheets as antiparallel β-sheets generate an additional (albeit weak) component at higher wavenumbers (around 1680 cm⁻¹) and the wavenumbers of intermolecular β-sheets, e.g. in aggregated proteins or amyloid peptides, are shifted towards lower wavenumbers (1625–1615 cm⁻¹).¹⁸

The amide II mode occurs around 1550 cm⁻¹ in H₂O and is mostly due to the amide N–H in-plane bending mode. In D₂O, the amide hydrogens undergo exchange with deuterium of the solvent, if exposed and not involved in strong intramolecular H-bonding, resulting in a ~90 cm⁻¹ downshift of the amide II mode. Thus, the extent and the kinetics of the amide HX, i.e. the rate of reduction of the ~1550 cm⁻¹ signal, is diagnostic for the extent of solvent exposure (tertiary structure) and the degree of involvement in H-bonding (secondary structure). Other amide modes are less sensitive to protein structure or dynamics, yet have been used for protein structural studies, as described in more detail elsewhere.¹⁸

The ATR-FTIR spectroscopy is a surface-sensitive version of FTIR that is based on creation of an evanescent field at the surface of an infrared-transparent internal reflection element (IRE), such as a germanium plate, during internal reflection of the infrared light within the IRE. As the evanescent wave decays with a characteristic length of 300–400 nm, this technique detects the sample, such as a lipid membrane with reconstituted protein(s) deposited on the surface of the IRE.

Plane-polarized infrared light is usually used to study not only the structural features of membrane proteins and lipids but also their spatial orientation.

The isotope-edited FTIR, which is the main topic of this article, is based on the spectral shift of the infrared signal due to replacement of atoms in biomolecules, such as H, C, N, by nonradioactive isotopes, such as ²H (D), ¹³C, ¹⁵N. For example, a lipid molecule with one unlabeled and one deuterated acyl chain generates infrared signals separated by ~730 cm⁻¹, which allows identification of selective cleavage of the *sn*-1 and *sn*-2 chains by phospholipase A₂ (PLA₂), reflecting the enzyme's mechanism.¹⁹ Combination of a ¹³C-labeled protein with an unlabeled protein results in amide I band splitting, allowing analysis of dynamic structural changes in each protein during intermolecular interactions.²⁰ Site directed or segmental ¹³C (or ¹³C, ¹⁵N or ¹³C=¹⁸O) labeling of a protein allows determination of the local conformational dynamics during functional transitions.^{21–23} The production of isotope-labeled proteins ranges from easy (e.g., uniform labeling by expression in bacteria grown a minimal medium containing a sole labeled nutrient) to moderate difficulty (e.g., chemical ligation of an unlabeled fragment of a protein with a labeled peptide or incorporation of a single labeled amino acid by the nonsense suppression method).

Apart from the advanced capabilities of FTIR spectroscopy, other features such as the relative simplicity of the experimental setup, the ease of the measurements and data analysis, the small amount of material per experiment (typically, around 50 μg of protein or lipid), and the wealth of the information on the structure of both protein and lipid (if present) from just one spectrum measured in 5 minutes, make the method attractive for broad applications, e.g. protein–protein and protein–lipid interactions and beyond.

2. Isotope effects on absorbance frequency and intensity

2.1. Model compounds

Replacement of an atom in a chemical group –X–Y by an isotope of different nuclear mass will shift the X–Y vibrational frequency. If Y has a much smaller mass than X, the X–Y stretching vibration can be considered as harmonic oscillation since Y will be essentially involved in the oscillation (like a ping pong ball attached to a bowling ball by a spring) and the frequency will be determined mainly by its mass ($\mu \approx m_Y$). Then, the frequency shift resulting from replacement of Y by an isotope (e.g., replacement of C–H with C–D or N–H with N–D) can be predicted using eqn (1). If the mass of Y is not very small, the HO approach is still valid when the X–Y bond is much stronger than the other bonds in a polyatomic molecule, such as the C=O double bond.¹⁷ Thus, the C–H to C–D, N–H to N–D, and ¹²C=¹⁶O to ¹³C=¹⁶O or to ¹³C=¹⁸O isotopic replacements, often used in protein and lipid FTIR spectroscopy, are expected to obey the HO rules, in particular the eqn (1), with acceptable accuracy.



The HO theory predicts that the C–H to C–D isotopic change will decrease the vibrational frequency by a factor of $\nu^j/\nu = 0.7338$ (ν^j and ν are the frequencies of the heavier isotopic and normal forms, respectively). This prediction is valid so long as the molecule is free from any kind of intermolecular interactions, ideally, when it is in gas phase.¹⁷ Vibrational coupling effects or interactions with the solvent or other solute molecules, e.g. via H-bonding, may drastically affect the resulting frequencies and intensities (see below). For example, the methylene asymmetric stretching wavenumbers of the CH₄ and CD₄ isotopomers of methane were 3019 cm⁻¹ and 2259 cm⁻¹, respectively.¹⁷ This corresponds to a factor of 0.7483, which agrees with the HO prediction with 2% accuracy. The respective wavenumber of ¹³CH₄ was 3010 cm⁻¹, as expected for a diatomic HO. The ¹³C=O stretching vibration of 3-¹³C-2,4-dimethyl-3-pentanone, 1-¹³C-2-methylpropionic acid, and C₆H₅¹³COOH (benzoic acid) dissolved in organic solvents was downshifted by 35–37 cm⁻¹ as compared to the wavenumbers of their normal (unlabeled) counterparts, in good agreement with the HO theory.¹⁷

Linear hydrocarbons in condensed phase produced strong infrared peaks in spectral regions 2926–2915 cm⁻¹ and 2855–2845 cm⁻¹ attributed to the antisymmetric and symmetric methylene stretching vibrations, and the respective wavenumbers of their perdeuterated (*i.e.*, totally deuterated) counterparts were red shifted by >700 cm⁻¹.^{24,25} For *n*-hexadecane, the spectral shifts were from 2926 cm⁻¹ to 2198 cm⁻¹ and from 2854 cm⁻¹ to 2095 cm⁻¹.²⁵ These data indicate that the spectral shift of the symmetric stretching vibration agrees with the diatomic HO prediction ($\nu^j/\nu = 0.7341$ as compared to the predicted value of 0.7338) but is less for the antisymmetric vibration ($\nu^j/\nu = 0.7512$), resulting in a wider separation between the perdeuterated peaks. The symmetric mode has been shown to be affected by intra- and intermolecular Fermi resonance interactions with the methylene bending mode, which results in secondary bands that contribute to the apparent intensity increase in the antisymmetric stretching region.²⁶ Such interactions are symmetry-forbidden for the antisymmetric stretching mode of an extended chain.

For molecules in gas phase, the change in the integrated molar absorption intensity (*A*) is predicted to be proportional to the square of the change in frequency:¹⁷

$$\frac{A^i}{A} = \left(\frac{\nu^j}{\nu}\right)^2 \quad (2)$$

where the superscript *i* means the isotopic form. Accordingly, the *Aⁱ/A* ratio for CD₄ and CH₄ was 0.6260 as compared to $(\nu^j/\nu)^2 = 0.5599$, signifying a fairly good agreement with the theory.¹⁷

2.2. Peptides and proteins

For proteins or lipid bilayers, which are in the condensed phase and are involved in a multitude of through-space and through-bond intermolecular interactions, the HO predictions are not expected to strictly hold. In addition, the spectral shifts of the amide I signal due to the most frequently utilized ¹³C=O

labeling strategy cannot be described by a simple diatomic HO modeling as the amide I mode is not a pure C=O stretching vibration. Nonetheless, incorporation of just one or up to 5 consecutive backbone ¹³C=O-labeled amino acid residues in synthetic peptides produced a new amide I peak red shifted from that of the unsubstituted peptide by 37 ± 4 cm⁻¹, in agreement with the HO theory applied to an isolated C=O bond.^{27–34} Incorporation of ¹³C-labeled amino acids located consecutively or separated by one or more ¹²C units in a α -helical 25-mer peptide revealed a complex behavior of both the vibrational frequency and intensity.^{33,34} The frequency of a single ¹³C amide unit was downshifted by 36–37 cm⁻¹ (from 1632–1630 cm⁻¹ to 1596–1593 cm⁻¹) and increasing number of ¹³C units resulted in increase in both ¹²C and ¹³C frequencies, increase in ¹³C intensity, and decrease in ¹²C intensity.³⁴ Variably spaced ¹³C units produced a non-monotonous (“damped sinusoidal”) dependence of the downshifted frequency on the number of intervening ¹²C units. Strikingly, the presence of one or two ¹²C units between two ¹³C units produced amide I signal of reduced intensity downshifted by additional ~ 10 cm⁻¹.^{33,34} These effects were interpreted in terms of distinct distance dependencies of the through-covalent-bonding mechanical coupling and through-space electrostatic and transition dipole couplings as well as relative dipolar orientations of the amide oscillators at defined positions along the spiraling helical chain.^{21,34} The increased frequency and reduced intensity of the ¹²C signal upon incorporation of more ¹³C units is thought to be caused by vibrational decoupling between the ¹²C amide oscillators.³⁴

The amide I intensity depends on the secondary structure and its continuity and is different for ¹²C=O and ¹³C=O peptide segments. For example, the molar absorption coefficient at the maximum of the amide I band (ϵ_{\max}) was 640 M⁻¹ cm⁻¹ at 1 °C (mostly helical structure) and 392 M⁻¹ cm⁻¹ at 91 °C (mostly unordered) for the unlabeled (¹²C=O) peptide³⁰ (here, M⁻¹ means per M of amino acid units). The respective values of ϵ_{\max} were smaller for the ¹³C-labeled segments by a factor of 0.7–0.8. This difference in molar absorptivity was explained by higher helicity and longer chains of the unlabeled segments rather than by intrinsic, isotope-specific differences³⁰ echoing with the concept of disruption of the ¹²C=O/¹²C=O coupling by intervening ¹³C=O units mentioned above.³⁴ This is reminiscent of the up to 2-fold stronger amide I signal from the ¹³C-labeled 4-residue segment located centrally compared to that when located at the N- or C-termini of a 20-mer helical peptide.^{28,35}

The agreement between the observed and diatomic HO-predicted frequency shift of ~ 37 cm⁻¹ upon a \geq ¹²C=O \rightarrow \geq ¹³C=O substitution may be fortuitous due to compensation of diverse effects on the resulting vibrational frequency. When other amide I components such as the C–N stretching mode are included in the computation, the ¹³C-induced spectral shift is around 43 cm⁻¹ rather than 37 cm⁻¹.³⁶ Uniform ¹³C-labeling of 40–42 amino acid residue peptides reduced the amide I frequency from 1658 cm⁻¹ to 1617 cm⁻¹ in α -helical state and from 1633 cm⁻¹ to 1590 cm⁻¹ in β -sheet state, *i.e.* a 41–43 cm⁻¹ downshift as predicted by the more accurate simulation.³⁷



Likewise, other studies on chemically synthesized or recombinantly produced proteins reported 40–44 cm^{-1} downshift of the amide I peak due to singly, segmentally, or uniform $^{13}\text{C}=\text{O}$ labeling,^{38–41} while the spectral shift due to uniform ^{13}C , ^{15}N labeling of recombinant proteins was 45–55 cm^{-1} .^{42,43}

Apart from intramolecular vibrational couplings, the amide I frequency also depends on H-bonding with the solvent. Thus, for totally hydrated α -helical peptides that form H-bonding with water, the frequency is reduced by as much as 17 cm^{-1} , e.g., from $\sim 1650 \text{ cm}^{-1}$ down to $\sim 1633 \text{ cm}^{-1}$.²¹ Peptides in dry state (dried from hexafluoroisopropanol solution)^{37,44} or embedded in lipid bilayers^{40,45} show amide I peaks around 1661–1657 cm^{-1} as compared to $1633 \pm 4 \text{ cm}^{-1}$ displayed by peptides totally hydrated in a D_2O buffer.^{28,30,33,34} A similar effect has been reported for a β -sheet peptide.⁴⁶

Studies on peptides in β -sheet conformation revealed position-dependent ^{13}C isotopic shifts in spectral frequencies and absorbance intensities.^{29,46–49} Incorporation of one or two $^{13}\text{C}=\text{O}$ amino acids in a synthetic, antiparallel β -sheet forming tetradecapeptide produced a spectrally downshifted (from 1634–1628 cm^{-1} to 1611–1606 cm^{-1}) amide I component of anomalously high intensity.²⁹ The integrated molar absorptivity of the downshifted amide I component of a single $^{13}\text{C}=\text{O}$ label was more than doubled relative to that of the main $^{12}\text{C}=\text{O}$ part, i.e. constituted around 17% of the total amide I contour as opposed to the expected $1/14 \approx 7\%$. The peptide with two consecutive labels generated a similar amide I band. However, when the two $^{13}\text{C}=\text{O}$ labels were separated by one $^{12}\text{C}=\text{O}$ unit, the absorptivity increased nearly 2-fold compared to the species with two consecutive $^{13}\text{C}=\text{O}$ labels. These anomalous intensities were interpreted by through-space transition dipole coupling and through covalent bonding and through H-bonding interactions between the $^{13}\text{C}=\text{O}$ and $^{12}\text{C}=\text{O}$ amide oscillators. In case of single or sequential double labeled peptides, the ^{13}C -substituted oscillators borrow intensity from the unlabeled oscillators of both opposite (through H-bonding) and same (through covalent bonding) strands whereas the separated oscillators do the same but pick up more intensity from neighboring $^{12}\text{C}=\text{O}$ units through covalent bonding. Interestingly, in case of an α -helical peptide, a single $^{13}\text{C}=\text{O}$ substitution produced an amide I component downshifted by $\sim 38 \text{ cm}^{-1}$ as expected for an isolated oscillator, suggesting the absence of (or weaker) through-H-bond interactions in α -helix compared to β -sheet, consistent with a shorter $\text{N}\cdots\text{O}$ distance by 0.034 Å in the latter case.²⁹ In addition to this difference, Huang *et al.*³³ reported that in α -helical structure the isotopically substituted groups are coupled essentially to each other but not to the unlabeled units, in contrast to β -sheet structure.

An important feature observed in β -sheet peptides, which has helped distinguish between parallel and antiparallel β -sheets, is that the spectral downshift of the $^{13}\text{C}=\text{O}$ substituted units that are involved in coupling with each other is larger by $\sim 10 \text{ cm}^{-1}$ compared to that caused by $^{13}\text{C}=\text{O}/^{12}\text{C}=\text{O}$ coupling (1594–1591 cm^{-1} and 1604–1601 cm^{-1} , respectively).^{46,47,50,51} The effects of vibrational couplings between

same or different isotope units on vibrational frequencies and intensities has been demonstrated in studies on a dodecamer that forms a β -hairpin structure with amide I peak at 1633 cm^{-1} .⁴⁸ Incorporation of two $^{13}\text{C}=\text{O}$ units in opposing strands produced a new component that was downshifted more when the labeled $^{13}\text{C}=\text{O}$ oscillators were spatially (not along the chain) closer. In addition, the more downshifted signal (closer spacing) was less intense than the less downshifted signal generated by more remotely spaced units. These features were interpreted by $^{13}\text{C}=\text{O}/^{13}\text{C}=\text{O}$ and $^{12}\text{C}=\text{O}/^{13}\text{C}=\text{O}$ vibrational coupling effects: closely spaced $^{13}\text{C}=\text{O}$ oscillators couple stronger resulting in a larger downshift but do not mix with the $^{12}\text{C}=\text{O}$ modes whereas those spaced more distantly (more $^{12}\text{C}=\text{O}$ units in between) experience less $^{13}\text{C}=\text{O}/^{13}\text{C}=\text{O}$ coupling (higher frequency) and more $^{12}\text{C}=\text{O}/^{13}\text{C}=\text{O}$ coupling (higher intensity through mixing). These data underscore the dependence of spectral frequencies and intensities caused by ^{13}C -labeling on the number and location of labeled residues. Upon insertion of 3 backbone $^{13}\text{C}=\text{O}$ units in a 3-stranded antiparallel β -sheet peptide (one per strand), the $^{12}\text{C}=\text{O}$ frequency of the main β -sheet rose to 1642 cm^{-1} , resulting in a 54 cm^{-1} spectral gap between labeled and unlabeled signals.⁴⁹ This effect is explained by reduction of the strand length and subsequent disruption of coupling between $^{12}\text{C}=\text{O}$ oscillators,⁴⁹ like in case of α -helix discussed above.³⁴

Double isotopic labeling of the amide carbonyl with ^{13}C and ^{18}O provides a better resolution of the local structure of proteins or peptides.⁴⁰ Labeling one out of 25 residues of an α -helical peptide with $^{13}\text{C}=\text{O}/^{18}\text{O}$ resulted in an amide I component shifted to 1595 cm^{-1} from the main (unlabeled) component centered at 1659 cm^{-1} .⁴⁰ This 64 cm^{-1} shift is less than the 78 cm^{-1} predicted by diatomic HO calculations but is consistent with a more rigorous Hartree–Fock theory. The robust spectral resolution permitted determination of the local structure and orientation of the transmembrane domains of membrane proteins by ATR-FTIR^{52,53} or transmission FTIR⁵⁴ on oriented peptide/lipid films. 2D infrared studies on another helical peptide $^{13}\text{C}=\text{O}/^{18}\text{O}$ labeled at 1 position at a time spanning the central 11 residues, mixed with the unlabeled peptide at various proportions in lipid vesicles, revealed excitonic through-space vibrational coupling (delocalization) between the labeled oscillators, or the absence of such coupling at higher dilutions, which allowed construction of the structure of the transmembrane helical dimer.⁵⁵

For peptides forming cross- β -sheet amyloid fibrils, such as the amyloid $\beta(1-40)$ peptide, a main amide I peak was detected at 1625 cm^{-1} and a downshifted component in the 1585–1575 cm^{-1} range depending on the position of the $^{13}\text{C}=\text{O}/^{18}\text{O}$ substituted residue.⁵⁶ The spectral shift was stronger for the sample composed of the labeled peptide due to vibrational coupling along the fibril axis and was blue shifted by $\sim 20 \text{ cm}^{-1}$ (from $\sim 1575 \text{ cm}^{-1}$ to $\sim 1595 \text{ cm}^{-1}$) when combined with a large molar excess of the unlabeled peptide. In contrast to α -helix, where a single label may not produce a diagnostic signal because of the absence of coupling with another labeled residue, the isotopic substitution of a single amino acid in



parallel in-register fibril forming peptides produces a strongly delocalized and red shifted normal mode indicative of inter-strand coupling along the fibril axis.^{57,58} A heptadecapeptide with $^{13}\text{C}=\text{O}$ substituted alanine at position 5 or 15 displayed amide I spectra with a major peak at 1632 or 1629 cm^{-1} and a minor (isotope-induced) peak at 1587 or 1591 cm^{-1} , signifying a 45 cm^{-1} or 38 cm^{-1} red shift for the N- and C-terminal segments that were deduced to be in ordered cross- β and unordered conformations, respectively.⁵⁹ Thus, the $^{13}\text{C}=\text{O}$ isotope-induced frequency shift in β -sheet peptides was significantly less ($\sim 45\text{--}50\text{ cm}^{-1}$)^{56,59} than that in α -helices ($\sim 65\text{ cm}^{-1}$).^{40,52-54}

The human islet amyloid polypeptide in both α -helical and β -sheet conformations, incorporating 1 or 2 adjacent $^{13}\text{C}=\text{O}$ labeled residues, was studied by 2D infrared spectroscopy.⁵⁸ Incorporation of a single label into the α -helical or β -sheet structure caused a $\sim 55\text{ cm}^{-1}$ and $32\text{--}42\text{ cm}^{-1}$ red shift, respectively, in qualitative agreement with the above assessment. The respective red shifts caused by two adjacent labels were $\sim 45\text{ cm}^{-1}$ and $\sim 52\text{ cm}^{-1}$ however, reversing the disparity between α -helical and β -sheet structures. This effect was consistent with normal mode analysis which highlighted differences between spectral shifts due to single vs. double (adjacent) labels in the two structures: two adjacent isotope labels cause a red shift from the uncoupled local mode frequency of the isotope substituted group for a β -sheet structure (through intra- and interstrand couplings) and a blue shift for α -helix (through covalent bond coupling).⁵⁸

The above discussions underscore the complexity of the behavior of vibrational frequencies and intensities and the involved isotopic effects. The frequencies and intensities of labeled vibrations depend on the secondary structure, the number of the labels and their sequentiality. The isotope labeling strategy should be selected taking into account the targeted secondary structure. For example, spectral features of a single isotope in an α -helix will produce only a local mode bearing little structural information but in a parallel in-register β -sheet structure or a cross- β -sheet structure a single label will be strongly coupled along and across the sheet plane producing the diagnostic amide I signal.^{49,58}

Interpretation of the shifted frequency in terms of secondary structure is not straightforward. For example, the $\sim 1633\text{ cm}^{-1}$ signal can be generated by β -sheet^{48,49} or by hydrated and totally amide deuterated α -helix.^{21,30-32,34} Additional methods such as circular dichroism or NMR are required to identify whether the spectral shift, e.g. from $\sim 1655\text{ cm}^{-1}$ to $\sim 1635\text{ cm}^{-1}$, reflects an $\alpha \rightarrow \beta$ transition or hydration of the protein's backbone carbonyls plus amide HX.^{29,30,49} The amide I signal in the spectrum of a $^{13}\text{C}=\text{O}$ or $^{13}\text{C}=\text{O}$ labeled peptide or protein in the $1600\text{--}1565\text{ cm}^{-1}$ region should be treated cautiously as it may be produced by either the labeled amino acid or the side chains of certain amino acids such as arginine or glutamic acid.^{30,48,56} Finally, possible artifacts stemming from the admixture of ^{13}C due to natural abundance ($\sim 1.1\%$) should be kept in mind to avoid misinterpretation of the intensity of the signal generated by the labeled site.

2.3. Lipids

Artificial membranes composed of isotopically labeled lipids have been used to study thermal phase transitions, lateral segregation, lipid transfer, lipid-protein interactions, and the influence thereof on the function of membrane proteins.⁶⁰⁻⁶⁴ Mostly, the acyl chain deuterated or carbonyl $^{13}\text{C}=\text{O}$ substituted lipids have been used to gain structural information on the membrane core or the polar headgroup region. The acyl chain methylene groups generate antisymmetric and symmetric stretching vibrations centered around 2920 cm^{-1} and 2850 cm^{-1} in protiated (CH_2) form while chain perdeuteration shifts these wavenumbers down to $\sim 2192\text{ cm}^{-1}$ and $\sim 2090\text{ cm}^{-1}$, respectively.^{25,60,65,66} The methylene stretching modes of perdeuterated chains are usually less intense in terms of peak height and the spectral spacing between the symmetric and antisymmetric peaks is larger ($\sim 100\text{ cm}^{-1}$) compared to the unlabeled lipids ($\sim 70\text{ cm}^{-1}$), similar to the features detected for *n*-alkanes discussed above. Analysis of ATR-FTIR spectra of unlabeled and acyl chain perdeuterated phospholipids produced nearly equal integrated molar absorption intensities: $A^i/A \approx 1.0$, indicating no isotope-induced changes in intrinsic absorptivities.⁶⁰

Temperature-induced gel to liquid crystal phase transitions of bilayer membranes composed of unlabeled and acyl chain perdeuterated lipids cause increase in the stretching wavenumbers by $2\text{--}5\text{ cm}^{-1}$.^{63,65-68} This spectral shift is interpreted in terms of decreased vibrational interactions between adjacent methylene units due to increasing ratio of *gauche/trans* conformational isomers.^{62,65,67} Similar methylene stretching vibrational shifts have been detected upon isotopic dilution of *n*-alkanes as well as phospholipids without involvement of conformational changes, explained by alterations in intermolecular Fermi resonance between like isotopic species.²⁵ Use of lipids with perdeuterated acyl chains has permitted analysis of thermal phase transitions of individual membrane components composed of binary or ternary lipid mixtures.^{62,68,69} Other vibrational modes of lipids such as the methylene rocking and scissoring modes and their spectral shifts upon deuteration have been analyzed to reveal membrane depth-dependent fractions of *gauche* rotamers and to detect lipid microdomain formation.^{70,71}

The carbonyl stretching mode of glycerophospholipids in aqueous suspensions generates a band with a peak wavenumber in the $1740\text{--}1730\text{ cm}^{-1}$ region which, upon resolution enhancement, reveals two components around 1742 cm^{-1} and 1727 cm^{-1} .^{67,72,73} These two components have been assigned to the carbonyls at *sn*-1 and *sn*-2 ester groups, respectively, and the splitting was ascribed to their distinct conformations.⁶⁷ Further studies on lipids selectively $^{13}\text{C}=\text{O}$ labeled at either *sn*-1 or *sn*-2 carbonyls resulted in two bands, the one generated by the labeled carbonyl red shifted by $40\text{--}43\text{ cm}^{-1}$.⁷²⁻⁷⁵ Strikingly, both the unlabeled and the $^{13}\text{C}=\text{O}$ labeled (and downshifted) bands were in turn composed of at least two components of their own separated by the same gap, i.e. $11\text{--}17\text{ cm}^{-1}$.^{72,74,75} It was concluded that the higher and lower frequency components of the carbonyl stretching vibrations are generated by dehydrated and hydrated fractions of the carbonyl groups



rather than structural difference that could account for a smaller (3–4 cm⁻¹) spectral difference.^{72,74} In general, an increasing polarity and H-bonding capability of the solvent results in lower frequency and wider vibrational bands.^{72,75} The red shift upon hydration can be explained by weakening of the >C=O double bond (parameter *k* in eqn (1)) due to strong >C=O...H-O-H hydrogen bonding.

FTIR studies on phosphatidylcholine selectively ¹³C=O labeled at one of the two ester carbonyls suggested that cholesterol forms H-bonding with the *sn*-2 but not *sn*-1 carbonyl group.⁷⁶ Polarized ATR-FTIR studies on substrate-supported multilayers of selectively ¹³C=O labeled phosphatidylcholines reported a ~40 cm⁻¹ red shift of the C=O double bond and a ~20 cm⁻¹ red shift of the ester CO-O single bond stretching vibrations, allowing determination of similar orientations of the C=O bonds of both chains (≥60°) and radically different orientations of the CO-O bonds relative to the membrane normal.⁷³

Both the methylene and the carbonyl stretching modes of lipids have been used to assess protein–lipid interactions and activities of membrane proteins. FTIR data on acyl chain perdeuterated phospholipids combined with unlabeled diacylglycerol identified domains of phosphatidylserine/diacylglycerol complexes that facilitated activation of protein kinase C.⁶² Membrane binding of human 5-lipoxygenase caused a shift of the carbonyl stretching band to higher wavenumbers, suggesting membrane dehydration.⁷⁷ Fluorescence data on lipid vesicles containing a surface polarity-sensitive probe implied membrane dehydration upon PLA₂ binding, however the lower and higher frequency components of the lipid carbonyl stretching bands (1728 cm⁻¹ and 1742 cm⁻¹) became stronger and weaker, respectively, indicating H-bonding between the protein and membrane lipids.⁷⁸ The mode of membrane binding and selective hydrolysis of membrane lipids by PLA₂ are described in detail in the forthcoming sections.

3. Protein–membrane interactions

3.1. Hierarchy of membrane proteins and modes of interactions with membranes

Around one-third of all gene products are membrane proteins,^{79,80} clearly implying that protein–membrane interactions constitute one of the major pillars of the temple of life. These proteins employ distinct mechanisms of membrane binding. While the integral proteins possess well-defined nonpolar stretches that interact with membrane lipids *via* hydrophobic contacts, the lipid-tethered proteins utilize a covalently attached fatty acid (most frequently, palmitic or myristic), glycosylphosphatidylinositol, or isoprenoid (farnesyl or geranylgeranyl) moiety for membrane binding.^{81–91} Polytopic proteins constitute the most abundant subclass of membrane proteins (>25%)⁸ and may contain 2 to 16 transmembrane helices.⁸¹ A smaller group of integral proteins utilizes an antiparallel transmembrane β-sheet structure arranged in a β-barrel-like architecture to serve as a transport passageway for relatively

large molecules or as receptors, cell adhesion and/or signaling molecules in mitochondria of animal cells, chloroplasts of plant cells and plasma membranes of prokaryotes such as the bacterial toxin α-hemolysin.^{8,82–84}

The peripheral proteins have been estimated to constitute ~9% of the human proteome and perform their function through transient association with plasma or organellar membranes.^{7–9} Some of these proteins are enzymes targeting membrane lipids for phosphorylation (*e.g.*, diacylglycerol kinase, phosphatidylinositol 3-kinase), oxygenation (lipoxygenases), or hydrolysis (*e.g.*, sphingomyelinase or phospholipases A₁, A₂, B, C, D). Typically, the peripheral proteins bind to membranes using an interfacial binding site (IBS).⁹ As the IBS is not always a clearly defined motif or domain, identification of the mode of membrane interaction of peripheral proteins remains a challenge. Analysis of 1328 experimentally determined and 1194 simulated (AlphaFold) peripheral protein structures identified some common features of the IBS, *i.e.*, protrusions rich in nonpolar amino acids (protruding hydrophobes)⁹² and lysine, which support membrane binding through hydrophobic and ionic contacts, respectively.⁹ In addition, there are many membrane binding modules specifically recognizing individual lipids or their assemblies. For example, most isoforms of protein kinase C become activated through binding to membrane-residing diacylglycerol using the C1 domain. Around 150 human proteins, including protein kinase C, contain one or more C2 domains, which mediate membrane binding, often in a calcium-dependent manner.⁹³ A range of signaling proteins bind to phosphatidylinositol phosphorylated at certain sites through the pleckstrin homology or FYVE domain, which recruit the respective proteins to distinct membranes where their function is most physiologically relevant.⁷ Other specialized lipid binding domain families are described elsewhere.^{9,93}

Peripheral proteins, including enzymes such as lipoxygenases, annexins, lipid kinases, phosphatases, and hydrolases, rely on binding to membranes with certain lipid composition, at a defined mode, including the depth of membrane insertion and the spatial orientation. The mode of membrane binding is crucial for the function of these proteins,⁹⁴ yet the “characterization of these fundamentally important lipid–protein interactions with experimental techniques is...prohibitively challenging”.⁹⁵ The forthcoming text will illustrate an example of how isotope-edited FTIR spectroscopy can determine the precise mode of membrane binding and thereby provide insight into the mechanism of action of a typical interfacial enzyme, a secreted PLA₂.

3.2. Selective hydrolysis of lipids by PLA₂

Secreted PLA₂s are medium-sized (13–16 kDa) extracellular enzymes that hydrolyze the *sn*-2 acyl chain of glycerophospholipids of cell membranes and liberate unsaturated fatty acids, in particular arachidonic acid, and lyso-phospholipid. These products serve as precursors for the biosynthesis of various eicosanoids or platelet-activating factor, respectively, that possess multiple biological activities, including cell signaling, apoptosis, inflammation, allergy, and tumorigenesis. Secreted



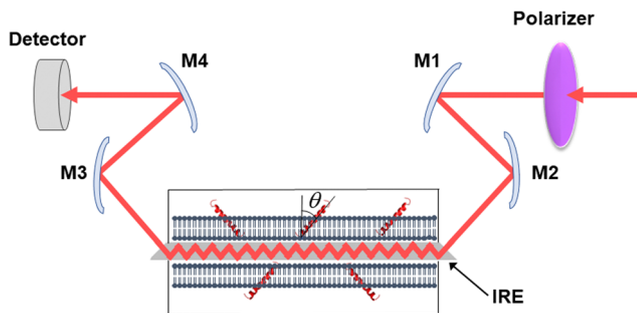


Fig. 1 Schematic presentation of the ATR-FTIR setting. The infrared light is plane-polarized via a polarizer and directed into the internal reflection element by mirrors M1 and M2. The incident light performs a series of internal reflections, exits the IRE and is directed to the detector by mirrors M3 and M4. The lipid membrane with embedded protein, shown as a single helix, is deposited at both surfaces of the IRE. Reproduced from ref. 94, with permission.

PLA₂s do not have special membrane binding or activation domains like many other interfacial enzymes, although an interfacial adsorption surface has been identified for each isoform. They bind to the cellular plasma membrane in a unique, “productive” mode that results in enzyme activation.⁹⁴ One of the factors that facilitate this process of interfacial activation of cationic PLA₂s is the presence of anionic lipids in the membrane.⁹⁶ To understand whether the elevated enzyme activity results from more efficient binding of the cationic protein to the negatively charged membrane surface or from higher intrinsic activity of the enzyme towards the anionic lipid, ATR-FTIR experiments have been conducted on lipid bilayers composed of an equimolar mixture of a zwitterionic lipid dipalmitoylphosphatidylcholine (DPPC) with uniformly perdeuterated acyl chains, DP(d₆₂)PC, and an unlabeled anionic lipid dipalmitoylphosphatidylglycerol (DPPG). In an ATR-FTIR experiment, the sample, in this case a lipid bilayer, is deposited on a germanium plate (Fig. 1). The incident light is directed into the germanium IRE at one edge with the aid of a set of mirrors and travels to the other edge by means of a series of internal reflections. At each internal reflection, an evanescent wave is created above the surface of the IRE that is absorbed by the sample deposited on it.

As described in Section 2.3, deuteration of the lipid acyl chains of DP(d₆₂)PC results in a $>700\text{ cm}^{-1}$ downshift of the methylene stretching vibrational frequency thereby producing FTIR spectra with well-resolved signals from the acyl chains of both unlabeled and deuterated lipids (Fig. 2a). The CD₂ absorbance band displays lower intensity and larger width compared to the CH₂ band, consistent with earlier observations.⁶⁰ Addition of PLA₂ results in hydrolysis of both lipids and removal of a fraction of the reaction products (free fatty acid and lyso-phospholipid) from the membrane, as evidenced by the reduction of the methylene stretching absorbance intensities. This is consistent with removal of $>30\%$ of the bilayer material from membranes caused by lipid hydrolysis by snake venom PLA₂s.^{97–99} A plot of the normalized intensities of the methylene stretching bands versus PLA₂ concentration indicates that both the zwitterionic and the acidic lipids are hydrolyzed to similar extents at a given PLA₂ concentration (Fig. 2b). Thus, the result of this isotope-edited ATR-FTIR experiment implies that the higher enzymatic activity of PLA₂ towards membranes containing negatively charged lipid(s) stems from a stronger electrostatic attraction between the enzyme and the membrane rather than from the intrinsic preference of the enzyme for the anionic lipid.¹⁹

PLA₂ activity depends on the presence of the reaction products, *i.e.* the free fatty acid and the lysolipid, in the membrane. The membrane binding affinity as well as the burst activation of PLA₂ following the initial lag period positively correlated with the presence of lipid hydrolysis products.^{100–103} Some studies suggested retention of the reaction products in the membrane^{100,101} while others demonstrated removal of a significant fraction of both products from the membrane upon hydrolysis by PLA₂.^{97–99} Isotope-edited FTIR has been used to determine whether both products of lipid hydrolysis dissociate from the membrane after lipid hydrolysis to equal extents or one of them preferentially accumulates in the membrane and plays a prevalent role in PLA₂ activity. PLA₂ was applied to supported membranes of DPPC with perdeuterated *sn*-1 chain and unlabeled *sn*-2 chain (Fig. 3a). In this case, the $>700\text{ cm}^{-1}$ separation of the methylene stretching vibrational signals

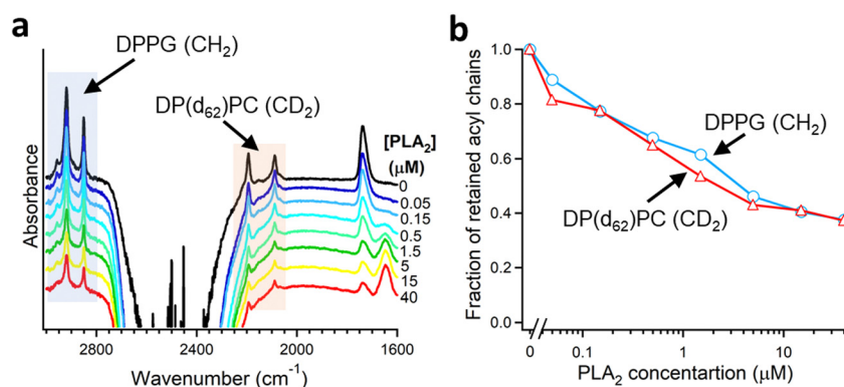


Fig. 2 Hydrolysis of the two components of a binary lipid bilayer by PLA₂. (a) ATR-FTIR spectra of a lipid bilayer composed of an equimolar combination of DPPG and chain-perdeuterated DP(d₆₂)PC at different concentrations of PLA₂, as indicated. (b) Dependence of the integrated methylene absorbance intensities of both lipids on PLA₂ concentration. Adapted from ref. 19, with permission.



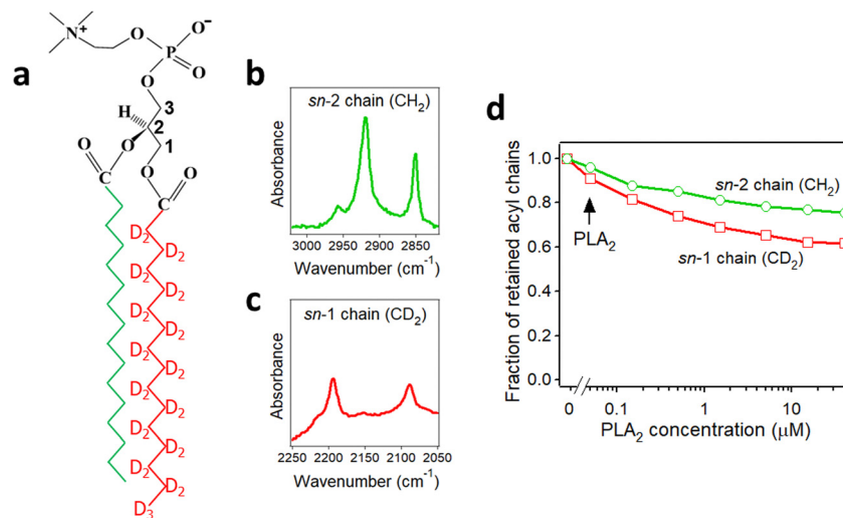


Fig. 3 Hydrolysis of the acyl chains of DPPC with deuterated *sn*-1 chain and unlabeled *sn*-2 chain by PLA₂. (a) Structure of DPPC with deuterated *sn*-1 chain (red) and unlabeled *sn*-2 chain (green). (b) and (c) The methylene stretching bands of the unlabeled (b) and deuterated (c) acyl chains of the lipid. (d) Dependence of the integrated methylene absorbance intensities of the unlabeled (green) and deuterated (red) acyl chains on PLA₂ concentration. Adapted from ref. 19, with permission.

provided spectral resolution of both acyl chains and thereby allowed monitoring the removal of each chain from the membrane upon lipid hydrolysis (Fig. 3b and c). As shown in Fig. 3d, while both CH₂ and CD₂ signals monotonously decrease with increasing PLA₂ concentration, the reduction in the CD₂ signal, which represents the lyso-phospholipid containing the *sn*-1 chain, prevails that of the CH₂ signals, which represents the free fatty acid. This result implies that the free fatty acid preferentially accumulates in the membrane following hydrolysis by PLA₂ and may modulate the enzyme's function by affecting the membranes surface charge and the structure.

3.3. Positioning of PLA₂ with respect to the membrane

As mentioned above, interfacial enzymes are activated by binding to biomembranes and forming a protein–membrane complex of highly specific configuration, including the depth of membrane insertion and the geometric orientation of the protein with respect to the membrane. For enzymes that use a membrane lipid as substrate, such as various PLA₂ isoforms, this specific configuration is a prerequisite for efficient enzyme function as it determines the strength of membrane binding and the mode of substrate accession and product release, *i.e.* the most important steps of the whole catalytic cycle.¹⁰⁴ Determination of the positioning of a secreted PLA₂ on a lipid membrane has been achieved by combination of protein engineering, isotope-edited FTIR, and fluorescence spectroscopy, as described below.

As shown in Fig. 1, the orientation of a molecule that has a rotational axis of symmetry, such as a helix, can be determined by polarized ATR-FTIR spectroscopy. Briefly, the spectra obtained with infrared light polarized coplanar and orthogonal to the plane of incidence are used to determine the dichroic ratio, *i.e.* the ratio of absorbances at two polarizations. The measured dichroic ratio, together with known parameters such

as the angle of the helical amide I transition dipole with respect to the helical axis ($39^\circ \pm 1^\circ$), the Cartesian electric vector components of the evanescent wave, and the refractive indices of the IRE, the membrane, and the bathing solution, allow determination of the helical order parameter which directly provides the average orientation of the helix relative to the membrane (angle θ in Fig. 1).^{18,105} Determination of the orientation of a protein of irregular shape is not trivial, however, as the protein may have multiple helices of arbitrary orientations. It has been shown that determination of the orientation of the whole protein requires the orientations of at least two different helices.¹⁰⁶ Human pancreatic PLA₂ has been chosen to solve the problem because it is a relatively simple protein with three α -helices, an N-terminal helix (residues 1–10) and two internal helices. A semisynthetic, segmentally ¹³C-labeled PLA₂ was produced where an unlabeled synthetic peptide corresponding to the N-terminal helix, with a thioester functional group at the C-terminus, was ligated to a uniformly ¹³C-labeled construct corresponding to the rest of the protein.¹⁰⁶ The residue 11 (the first residue of the ¹³C-labeled construct) is a cysteine and is ideally located for thioester-to-cysteine peptide ligation (Fig. 4). Analysis of the protein structure showed that the two internal helices were nearly antiparallel, so, in terms of orientation, the protein could be treated as a two-helix system (in ATR-FTIR, parallel and antiparallel orientations are equivalent). ATR-FTIR spectra with coplanar and orthogonal polarizations of the incident light produced multicomponent amide I bands, with the components generated by the unlabeled N-terminal helix and the two ¹³C-labeled internal helices located at 1656 and 1616 cm⁻¹, respectively (Fig. 5a and b). These data, along with specially developed analytic geometry algorithms, allowed determination of the orientations of the helices and hence of the whole protein relative to the membrane. The depth of membrane insertion was determined by experiments of



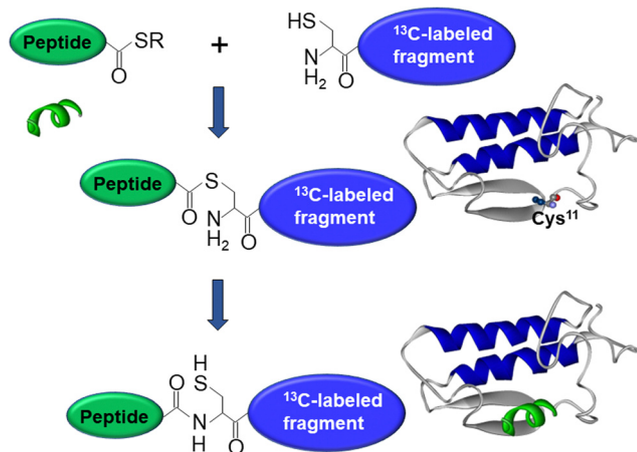


Fig. 4 Graphical description of the thioester–cysteine chemical ligation. The synthetic unlabeled peptide (green) with a C-terminal thioester group reacts with the thiol of the N-terminal cysteine of a recombinant, uniformly ^{13}C -labeled protein fragment (blue) followed by S-to-N acyl transfer and formation of a natural peptide bond, resulting in a semisynthetic, segmentally isotope-labeled protein. Adapted from ref. 94, with permission.

quenching of the fluorescence of the single tryptophan-3 of pancreatic PLA₂ by lipids brominated at various positions along the acyl chains. Isotope-edited polarized ATR-FTIR and fluorescence data thus produced the structure of the protein–membrane complex with a unique positioning of the protein on the membrane in terms of geometric configuration and depth of membrane insertion (Fig. 5c).¹⁰⁶

In this structure, the *z* coordinate of each atom of the protein is its distance from the membrane center so the location of each amino acid can be easily pinpointed. The protein was found to significantly penetrate into the membrane hydrocarbon core, with tryptophan-3 located at 9 ± 1 Å from the membrane center and providing a hydrophobic anchor for membrane docking of PLA₂ together with phenylalanine-19 and leucine-20. Penetration of a cobra venom PLA₂ into phosphatidylcholine bilayers 5–7 Å

below the lipid headgroup plane has been reported,⁹⁹ in good agreement with this finding. In addition to hydrophobic interactions, several cationic amino acid residues were involved in H-bonding with lipid carbonyl oxygens or in ionic interactions with lipid phosphate groups. Inspection of the structure shown in Fig. 5c indicates that the opening of the substrate-binding cleft faces the membrane surface so that the catalytic histidine-48 is located at ~ 7 Å from the *sn*-2 ester group of the glycerophospholipid allowing the lipid to travel a shorter distance to reach the catalytic site compared to ~ 15 Å proposed earlier.^{107,108} Thus, this technique, where isotope-edited FTIR plays a central role, provides an efficient tool to elucidate the mode of membrane binding of a peripheral protein and the mechanism of its action.

4. Protein–protein interactions

4.1. Overview and importance of protein–protein interactions

Protein–protein interactions play crucial roles in cell function by supporting vitally important biomolecular structures or processes. In fact, from 300 000 to 650 000 promiscuous protein–protein interactions are involved in human physiology.² Protein assemblies can be divided into two classes, *i.e.*, stable multiprotein structures that serve specific functions and transient complexes that are mostly involved in the dynamic events of cell signaling.^{1,3} Interactions between protein subunits are facilitated by a combination of nonspecific (hydrophobic, ionic, H-bonding) and specific (*e.g.*, shape-complementary, hand-in-glove- or lock-and-key-type) contacts between domains and motifs that often contain critically important “hot spot” amino acids.^{1,3,81,109–111} As an antithesis of the lock-and-key interactions between (relatively) rigid domains, the “induced fit” or “folding-upon-binding” interactions between initially flexible protein modules such as antibodies or receptors or their ligands that acquire defined shape upon interaction result in promiscuity and polyreactivity effects of profound biological importance.^{3,112–116}

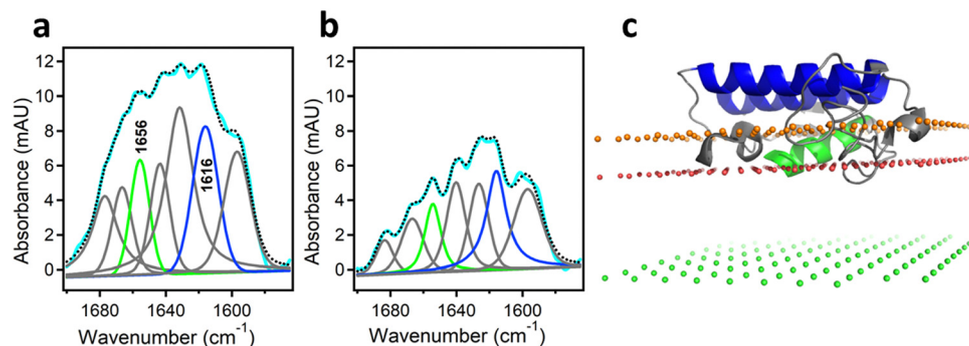


Fig. 5 Determination of the orientation of a membrane-bound protein by polarized ATR-FTIR spectroscopy. (a) and (b) ATR-FTIR spectra of membrane-bound human pancreatic PLA₂ with unlabeled N-terminal helix and ^{13}C -labeled fragment at coplanar (a) and orthogonal (b) polarizations of the incident light with respect to the plane of incidence. The measured spectrum is shown in cyan and the sum of all amide I components is shown as dotted black line. The amide I components generated by the unlabeled and ^{13}C -labeled helices are shown in green and blue, respectively. (c) The structure of the protein bound to the membrane with its N-terminal and internal helices shown in green and blue, respectively. The three planes indicate the locations of the membrane center (green), the *sn*-1 carbonyl oxygens (red) and the phosphate atoms (orange) of the lipid. Adapted from ref. 106, with permission.



The transient complexes between proteins are formed through highly specific interactions of moderate affinity (typically, nanomolar to micromolar dissociation constants).^{117–119} Important examples of transient protein–protein complexes are the enzyme–substrate,^{117–120} enzyme–inhibitor,^{121,122} ligand–receptor,^{123–125} and antigen–antibody interactions.^{3,111,126} More than 100 signaling proteins in human genome contain one or more Src homology 2 domain(s) that mediate binding to phosphorylated tyrosine involved in signal transduction.¹²⁷ Other protein modules such as the PDZ domain, breast cancer susceptibility protein 1, forkhead-associated domain, mad homology domain 2, interferon regulatory factor 3, polo-box domain unique to the polo-like kinases, the FF, WW, and WD40 motifs containing the respective dipeptides mediate the transient binding of signaling molecules to phosphoserine or phosphothreonine.^{2,128} The 14-3-3 family proteins bind to and regulate around 1200 partners involved in a plethora of cellular functions and are regulated themselves by caspase-mediated proteolysis, lysine acetylation, or serine/threonine phosphorylation.^{2,129} The Src homology 3 (SH3) domain, on the other hand, binds proline-rich protein motifs that fit into a groove on the surface of the SH3 domain.¹³⁰ Finally, regulatory (effector) proteins up- or down-regulate the activity of enzymes or switch from one to another function of multidomain modular proteins *via* interacting with an allosteric site.¹³¹

A special type of protein–protein interactions is aggregation of misfolded proteins and formation of amyloid fibrils that are involved in many pathologies. Some prominent examples are involvement of the amyloid β ($A\beta$) and tau proteins in Alzheimer's disease, α -synuclein in Parkinson's disease, prion protein in Creutzfeldt–Jakob disease, insulin in iatrogenic amyloidosis, immunoglobulin light chain and transthyretin in cardiac amyloidosis and polyneuropathy, islet amyloid polypeptide (amylin) in type II diabetes, hemoglobin in sickle cell anemia *etc.*^{132–141} Amyloidogenesis occurs with or without genetic, inheritable mutations in the respective protein, and the fibrils display remarkable structural similarities, such as formation of the cross- β -sheet structure, despite the absence of sequence similarities.^{137,140}

The importance of protein–protein interactions in cell physiology has led the researchers to develop multiple methods to analyze protein complex formation and the structure and function of the complexes. An incomplete list of these methods includes the yeast two-hybrid system, the protein–fragment complementation assay, fluorescence resonance energy transfer, co-immunoprecipitation, tandem affinity purification coupled to mass spectrometry, (in-cell) cross-linking mass spectrometry, (in-cell) single-molecule tracking, stable isotope labeling by amino acids in cell culture, and computational (*in silico*) methods.^{110,142–149} Described in the following sections are the use of isotope-edited FTIR to analyze the structural aspects of $A\beta$ fibrillogenesis and the regulation of sarcoplasmic reticulum (SR) Ca^{2+} -ATPase by phospholamban (PLB).

4.2. Protein aggregation

The $A\beta$ peptide aggregates constitute a histopathological hallmark of the Alzheimer's disease (AD).¹⁵⁰ $A\beta$ is derived from the

amyloid precursor protein, a bitopic protein in neuronal membranes, and represents mostly its α -helical transmembrane part. Upon enzymatic cleavage and exposure to aqueous medium, the peptide undergoes transition to β -sheet structure followed by fibrillogenesis. $A\beta$ occurs in various forms in human brain in terms of the number of amino acids and enzymatic modifications; in addition to the prevalent 40- and 42-residue species ($A\beta_{1-40}$, $A\beta_{1-42}$), N-terminally truncated and pyroglutamylated forms ($A\beta_{pE3-40}$, $A\beta_{pE3-42}$) are present as well and are hypertoxic.³⁷ Since in human brain these peptides are intermixed and undergo aggregation into hetero-complexes, it is of interest to understand the structural changes in each species during aggregation and the mutual structural effects. Isotope-edited FTIR has been used to achieve this goal.

The synthetic, lyophilized, uniformly ^{13}C -labeled $A\beta_{1-42}$ and unlabeled $A\beta_{pE3-42}$ peptides were dissolved in hexafluoroisopropanol to disperse possible aggregates and were dried on a CaF_2 FTIR window. These samples generated amide I spectra with peaks around 1617 cm^{-1} and 1658 cm^{-1} and amide II peaks around 1533 cm^{-1} and 1544 cm^{-1} , respectively (Fig. 6a and c, black lines). In view of the $\sim 40\text{ cm}^{-1}$ downshift due to ^{13}C -labeling, these data indicate α -helical structure for both peptides before hydration. To observe the kinetics of α -helix to β -sheet transition upon hydration as well as the intermediate oligomeric forms that are the main toxic species, the structural change was slowed down by pumping D_2O -saturated nitrogen gas into the chamber of the spectrometer allowing measurements at distinct stages of the transition. Hydration with D_2O vapor resulted in gradual shift of the α -helical peaks to 1588 cm^{-1} for ^{13}C - $A\beta_{1-42}$ and to 1632 cm^{-1} for $A\beta_{pE3-42}$, indicating α -helix to β -sheet transition of both proteins (Fig. 6a and c). Concomitantly, the intensity in the initial (non-deuterated) amide II band declined because of the downshift of the deuterated amide II mode by $\sim 90\text{ cm}^{-1}$ (see Introduction). The kinetics of the HX, *i.e.* the time-dependence of amide II intensity, and of the α -helix- β -sheet transition, *i.e.* the ratio of absorbance intensities of the β -sheet and α -helical amide I signals, are shown in Fig. 6b and d. ^{13}C - $A\beta_{1-42}$ readily underwent amide HX and α -helix to β -sheet transition once exposed to D_2O vapor (Fig. 6b) but the peptide was not entirely amide-deuterated after 5 hours of exposure to D_2O vapor as indicated by the residual amide II intensity (red line in Fig. 6a and blue line in Fig. 6b), indicating partial solvent protection possibly due to a compact tertiary structure. $A\beta_{pE3-42}$ behaved differently. Despite the significant decrease in the amide II intensity during the first 1.0–1.5 hours of deuteration, indicative of extensive HX, the peptide initially experienced little structural changes (Fig. 6d). Following the lag period, the peptide underwent α - β transition in a sigmoidal manner, yet a significant fraction of the initial α -helix as well as β -turn structure persisted after the transition leveled off (shoulder around 1677 – 1658 cm^{-1} of the red spectrum of Fig. 6c). In contrast to ^{13}C - $A\beta_{1-42}$, nearly all amide protons of $A\beta_{pE3-42}$ exchanged for deuterium, as indicated by the reduction of the amide II signal to zero (Fig. 6c), suggesting a more open, solvent-exposed tertiary fold. This result is consistent with recently published



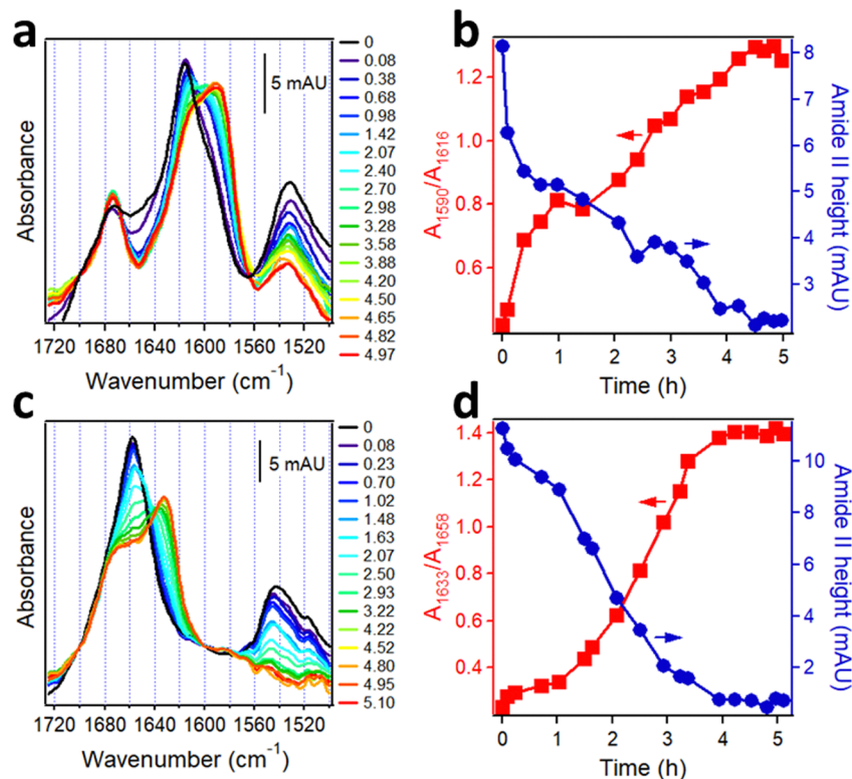


Fig. 6 Structural changes in Aβ peptides upon hydration with D₂O-saturated nitrogen gas. FTIR spectra of ¹³C-Aβ₁₋₄₂ (a) and Aβ_{pE3-42} (c) dried on a CaF₂ window (black lines) and upon injection of D₂O-saturated nitrogen for ~5 h (blue to red). Panels a and c are appended with columns indicating time of exposure to D₂O vapor (in hours). Panels (b) and (d) show the kinetics of β-sheet structure formation in ¹³C-Aβ₁₋₄₂ (b) and Aβ_{pE3-42} (d) (red) and change in amide II intensities (blue) as a function of time of exposure to D₂O vapor. Adapted from ref. 37, with permission.

fluorescence data showing solvent-inaccessibility of tyrosine-10 of Aβ₁₋₄₂ and solvent exposure of same residue of Aβ_{pE3-42}.¹⁵¹

Above data uncover distinct structural pathways of α-helix to β-sheet structural transitions of the unmodified and pyroglutamylated Aβ peptides, *i.e.*, resistance of the latter to transition to β-sheet structure and a less compact tertiary fold. However, in brain tissue and in the extracellular plaques various forms of Aβ are blended and may affect each other's structural behavior. Such mutual structural effects have been analyzed as follows. Aβ_{pE3-42} and ¹³C-Aβ₁₋₄₂ have been dissolved in hexafluoroisopropanol and deposited separately on the opposite sides of a CaF₂ FTIR window as well as mixed at 1:1 molar ratio and deposited on the same side of another CaF₂ window. These two samples were dried by desiccation and mounted on two mutually orthogonal vertical faces of a rotatable stage and placed in the spectrometer. Consecutive FTIR spectra of both separate and combined samples were measured alternately by turning the stage back and forth by 90 degrees about a 4-fold vertical axis while the chamber was purged with D₂O-saturated nitrogen. The difference between these spectra at a given time point, *i.e.* at a given level of amide deuteration, then revealed the mutual structural effects.³⁷

These samples generated complex spectra incorporating the features of both unlabeled and ¹³C-labeled peptides. In the isolated system, both ¹³C-Aβ₁₋₄₂ and Aβ_{pE3-42} were α-helical in dry state indicated by the amide I peaks around 1618 cm⁻¹ and

1658 cm⁻¹, respectively (black line in Fig. 7a). The weaker amide I intensity of ¹³C-Aβ₁₋₄₂ is consistent with observations of 20–30% lower amide I peak heights of ¹³C-labeled helical peptide segments compared to the unlabeled segments.³⁰ Hydration by D₂O vapor transformed the spectra and produced downshifted peaks at 1586 cm⁻¹ and 1628 cm⁻¹ with higher frequency shoulders located at 1597 cm⁻¹ and 1639 cm⁻¹ (red line in Fig. 7a). The strong signal at 1586 cm⁻¹ and 1628 cm⁻¹ is readily assigned to intermolecular β-sheet structure in ¹³C-Aβ₁₋₄₂ and Aβ_{pE3-42}, respectively, while the upshifted weaker signal at 1597 cm⁻¹ and 1639 cm⁻¹ indicates a smaller fraction of β-sheet with intramolecular H-bonding.^{18,152} An additional peak around 1673 cm⁻¹ can be assigned to β-turn structure in Aβ_{pE3-42}.

Spectra of the combined ¹³C-Aβ₁₋₄₂/Aβ_{pE3-42} sample hydrated by D₂O vapor displayed a stronger higher frequency signal assigned to intramolecular H-bonded β-sheet and a weaker lower frequency, intermolecular H-bonded β-sheet signal (red line in Fig. 7b). In fact, the signal at 1586 cm⁻¹ assigned to the intermolecular β-sheet of ¹³C-Aβ₁₋₄₂ was not present in the combined peptide sample at all. These spectral effects can be amplified by resolution enhancement techniques. The second derivative spectra shown in Fig. 8 identify strong signals of the isolated sample located at 1584 cm⁻¹ and 1627 cm⁻¹ and weaker signal around 1597 cm⁻¹ and 1537 cm⁻¹ (blue line of Fig. 8) signifying larger fractions of aggregates with intermolecular H-bonding for both



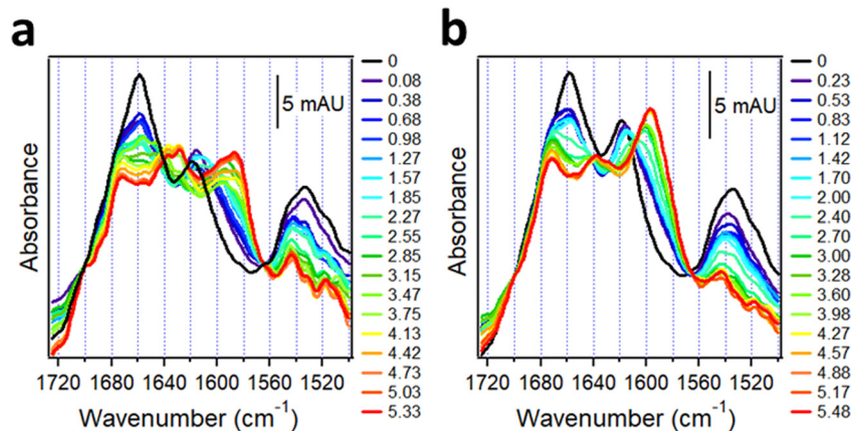


Fig. 7 Structural changes in ^{13}C - $\text{A}\beta_{1-42}$ and $\text{A}\beta_{\text{pE3-42}}$ upon hydration with D_2O -saturated nitrogen gas when physically separated (a) and combined at 1:1 molar ratio (b). Both panels are appended with columns indicating time of exposure to D_2O vapor (in hours). Reproduced from ref. 37, with permission.

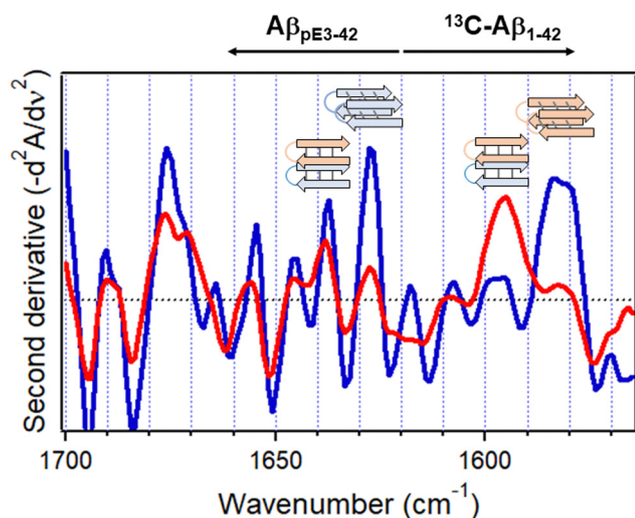


Fig. 8 Inverted second derivative amide I spectra of ^{13}C - $\text{A}\beta_{1-42}$ and $\text{A}\beta_{\text{pE3-42}}$ after 5 h hydration with D_2O -saturated nitrogen gas when physically separated (blue) and combined at 1:1 molar ratio (red). Cartoons for homo- or hetero-oligomers stabilized by intermolecular and intramolecular H-bonding, respectively, are shown above respective spectral features (^{13}C - $\text{A}\beta_{1-42}$, rose, and $\text{A}\beta_{\text{pE3-42}}$, blue). Reproduced from ref. 37, with permission.

^{13}C - $\text{A}\beta_{1-42}$ and $\text{A}\beta_{\text{pE3-42}}$. In the spectrum of the combined sample, the relative intensities of these signals were reversed (red line of Fig. 8), indicating conversion of the intermolecular H-bonded aggregates to β -sheet structures stabilized by intramolecular H-bonding, as schematically shown above the respective spectral features in Fig. 8. These data identify mutual structural effects of the unmodified and pyroglutamylated $\text{A}\beta$ peptides, *i.e.*, diversion of the aggregation process from fibril formation stabilized by intermolecular H-bonding to oligomeric structures of hairpin-like β -sheets that may exert a stronger cytotoxic effect as the oligomers are more toxic than the fibrils.^{37,141,153–155} This is an example of how isotope-edited FTIR spectroscopy can provide insight into the molecular mechanism of a disease such as AD.

4.3. In-membrane protein–protein interactions

Described in this section is the use of isotope-edited FTIR spectroscopy to analyze the regulatory interaction between two membrane proteins, *i.e.*, regulation of the activity of SR Ca^{2+} -ATPase by PLB that plays a central role in the contraction–relaxation cycle of heart muscle.¹⁵⁶ Excitation of the cardiomyocyte triggers Ca^{2+} influx through the voltage-sensitive Ca^{2+} channels, which in turn activates the type-2 ryanodine receptor Ca^{2+} channels in the SR membrane resulting in additional increase in the cytosolic Ca^{2+} level and contraction by the Ca^{2+} -sensitive contractile proteins. This is followed by relaxation that takes place mainly by pumping the cytosolic Ca^{2+} back into the SR lumen by the SR Ca^{2+} -ATPase. The activity of the Ca^{2+} -ATPase is regulated by PLB, a relatively small bitopic protein in the SR membrane, as schematically shown in Fig. 9a. Specifically, PLB moderates the enzymatic activity of the Ca^{2+} -ATPase and phosphorylation of PLB by cAMP- and/or Ca^{2+} /calmodulin-dependent protein kinases ceases the inhibitory effect.^{157–159} Various forms of cardiomyopathy in humans are associated with reduction of the levels of both the SR Ca^{2+} -ATPase and PLB at mRNA and protein levels.¹⁵⁶ The regulatory effect of PLB has been shown to involve direct molecular contact between the ATPase and PLB and several intermolecular interaction sites have been identified but the mechanism of regulation remains unclear.^{158–160}

Since PLB exerts similar regulatory effects on both the cardiac and skeletal muscle SR Ca^{2+} -ATPases and the latter maintains higher functional stability following purification, ATR-FTIR studies have been carried out on recombinant, uniformly ^{13}C -labeled PLB and skeletal muscle SR Ca^{2+} -ATPases co-reconstituted in supported membranes composed of SR-extracted lipids.¹⁵⁷ First, spectra of each membrane-embedded protein were measured separately. The Ca^{2+} -ATPase generated amide I and amide II bands centered at 1658 cm^{-1} and 1546 cm^{-1} , respectively (red line in Fig. 9b). The strong signal at 1658 cm^{-1} is consistent with the mostly α -helical secondary structure of the Ca^{2+} -ATPase (Fig. 9a). ^{13}C -PLB produced a wide



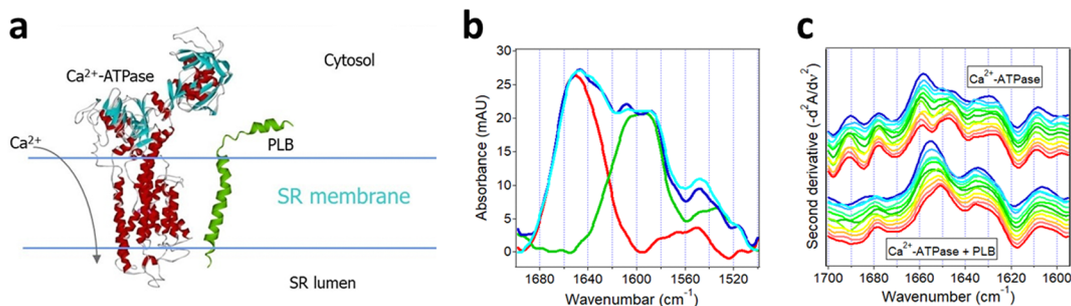


Fig. 9 Structural effects resulting from interaction of SR Ca^{2+} -ATPase with uniformly ^{13}C -labeled PLB. (a) Cartoon for the Ca^{2+} -ATPase and PLB in the SR membrane. (b) ATR-FTIR spectra of membrane-reconstituted Ca^{2+} -ATPase (red), uniformly ^{13}C -labeled PLB (green), the sum of the spectra of Ca^{2+} -ATPase and ^{13}C -PLB measured separately (blue), and the spectrum of the 1:11 molar combination of Ca^{2+} -ATPase and ^{13}C -PLB (cyan). (c) Inverted second derivative spectra of Ca^{2+} -ATPase alone (upper group of curves) and with ^{13}C -PLB at 11-fold molar excess (lower group of curves). Time of exposure to D_2O -based buffer was 2 min, 4 min, 9 min, 19 min, 36 min, 1 h, 1.4 h, 3 h, 4.4 h, 6 h, and 8 h (blue to red). Partially adopted from ref. 157, with permission.

amide I band composed of two components at 1608 cm^{-1} and 1585 cm^{-1} indicating α -helical and β -sheet structures, in accord with earlier FTIR studies,¹⁶¹ and an amide II band at 1525 cm^{-1} (green line in Fig. 9b). The 50 cm^{-1} spectral shift between the α -helical amide I modes of the two proteins is consistent with the expected isotopic effect.

The spectrum of the Ca^{2+} -ATPase and ^{13}C -PLB co-reconstituted at 11:1 PLB:ATPase molar ratio displayed the structural features of both proteins (cyan line in Fig. 9b). A resolution enhancement procedure, *i.e.* second derivative spectral analysis in the amide I region of Ca^{2+} -ATPase, was conducted to discern structural effects exerted by PLB. Time-dependent changes in the second derivative spectra over an 8 hour period unveiled interesting effects. In the initial stages of exposure of the proteins to D_2O , a strong component at 1658 cm^{-1} and weaker one at 1647 cm^{-1} were present in the spectrum of the Ca^{2+} -ATPase (Fig. 9c, upper family of curves), which can be readily assigned to non-deuterated and deuterated α -helices.¹⁸ By 8 hours of exposure to D_2O , the relative intensities of these components switched toward a larger fraction of the deuterated helices. The spectral features in the $1637\text{--}1625\text{ cm}^{-1}$ and $1700\text{--}1670\text{ cm}^{-1}$ regions represent β -sheet and turn structures while those below 1610 cm^{-1} are assigned to side chains. In the presence of PLB, the α -helical signal at 1658 cm^{-1} was downshifted by $3\text{--}4\text{ cm}^{-1}$ and its further downshift due to deuteration occurred much slower (Fig. 9c, lower family of curves). Both effects suggest stabilization of the α -helices of Ca^{2+} -ATPase upon interaction with PLB. More stable helices involve stronger intramolecular H-bonding and hence weaker $\text{C}=\text{O}$ covalent bonds, corresponding to lower vibrational frequency. The stronger $\text{C}=\text{O}\cdots\text{H}-\text{N}$ H-bonding, on the other hand, will slow down the amide deuteration, as detected. In addition, the relative intensities of non-deuterated *vs.* deuterated β -sheets of the Ca^{2+} -ATPase (signals around 1637 cm^{-1} and 1625 cm^{-1}) was shifted toward the former in the presence of PLB, in agreement with overall slower HX. Finally, the more stable structure and less effective HX of the Ca^{2+} -ATPase in the presence of PLB was confirmed by a stronger amide II band of the Ca^{2+} -ATPase around 1546 cm^{-1}

as compared to that in the spectral sum of the two proteins measured separately (*cf.* cyan and blue lines of Fig. 9b). The data thus indicate stabilization of the secondary structure of the Ca^{2+} -ATPase by PLB with no significant structural changes in PLB itself. These findings offer a mechanism for the inhibitory action of PLB on Ca^{2+} -ATPase through increased rigidity of the ATPase structure by PLB and hindrance of the conformational changes involved in the catalytic turnover.

5. Site-specific structure determination

5.1. Isotope labeling schemes

Data described above indicate the remarkable sensitivity of FTIR spectroscopy to the structural dynamics of proteins. However, FTIR usually provides the global rather than local structural information and hence is considered a “low-resolution” technique. Local, site-specific structure of proteins can be determined by isotope-edited FTIR spectroscopy using proteins that are isotopically labeled at selected amino acid residues. Since the conformation-sensitive amide I mode is produced mostly by the main chain $\text{C}=\text{O}$ stretching vibration (76%) as well as by the C–N stretching, C–C–N deformation, and N–H in-plane bending modes,¹⁸ the most common labeling schemes include carbonyl ^{13}C labeling, double ^{13}C , ^{15}N -labeling, or double $^{13}\text{C}=\text{O}$ -labeling of main chain atoms, which result in significant spectral shifts and resolution of the targeted site(s), as described in Section 2.2.

The incorporation of the labeled amino acids can be achieved by various methods, such as chemical synthesis^{162–165} or recombinant expression in an *E. coli* strain that is auxotrophic for a certain amino acid which is added into the growth medium in isotope-labeled form.^{166–168} For example, functionally active insulin has been produced by total chemical synthesis $^{13}\text{C}=\text{O}$ -labeled at a selected residue.¹⁶⁴ The synthetic method works so long as the synthesis of the whole sequence is practicable, and with the advent of the automated fast-flow peptide synthesis technology the limit of the number of amino acids in the synthetic



protein has been pushed beyond 200.^{169,170} The use of *E. coli* auxotrophs is a facile strategy as currently around 220 BL21(DE3)-based strains are available allowing production of proteins incorporating various combinations of isotopically labeled amino acids.¹⁶⁸ Native chemical ligation of synthetic or recombinantly expressed proteins is another strategy to generate relatively long protein chains incorporating isotopically labeled segments or individual amino acids; an example of a semisynthetic, segmentally isotope-labeled fully functional PLA₂ obtained by this method is described above (Section 3.3). FTIR analysis of the lens protein γ D-crystallin that was produced by thioester/cysteine chemical ligation of the unlabeled N-terminal domain with the ¹³C-labeled C-terminal domain allowed elucidation of the molecular mechanism of fibril formation.¹⁷¹ Transmembrane proteins, such as the cation-specific ion channel p7 of hepatitis C virus (63 amino acid residues), have been produced by thioester/cysteine or hydrazide-based chemical ligation of the p7(1–26) and p7(27–63) chains that were selectively labeled with ¹³C or ¹⁵N for FTIR and NMR studies.^{172,173} Other methods of incorporation of labeled amino acids in single or multiple sites, such as the use of modified tRNA to achieve nonsense suppression, are described below (Section 6.2).

5.2. Local and global structure of A β by segmental isotope labeling

The structure of the oligomers formed by the A β _{1–42} peptide and its truncated/pyroglutamylated counterpart, A β _{pE3–42}, has

been studied using synthetic peptides harboring uniformly ¹³C, ¹⁵N-labeled amino acids (Cambridge Isotope Laboratories). The labeled amino acids should constitute at least ~5–10% of the total number of residues to (a) produce a readily distinguishable amide I signal and (b) generate a normal mode vibration *via* inter-residue couplings that can be used for site-specific structure assessment. A β _{1–42} and A β _{pE3–42} were labeled at three or four consecutive amino acid residues, *i.e.* the K¹⁶L¹⁷V¹⁸ or V³⁶G³⁷G³⁸V³⁹ segments that play an important role in initiation of A β aggregation^{174,175} and were studied separately as well as combined at an equimolar ratio to gain insight into the structural pathways of fibrillogenesis. All peptide samples, incubated in aqueous buffer for 1 hour, featured a major amide I peak located between 1635 cm⁻¹ and 1630 cm⁻¹ that constituted 50 ± 5% of the total amide I band area (Fig. 10). These amide I wavenumbers indicate formation of oligomers of peptides in β -sheet structure with intramolecular H-bonding as the cross- β sheet fibrils generate amide I signal below 1630 cm⁻¹.^{50,56–58,176} Smaller fractions of turn structure (1677–1672 cm⁻¹), α -helix (1657–1652 cm⁻¹), and in some cases irregular (unordered) structure (~1644 cm⁻¹) were present as well.

The spectral component generated by the ¹³C, ¹⁵N-labeled stretches K¹⁶L¹⁷V¹⁸ and V³⁶G³⁷G³⁸V³⁹ were clearly manifested at 1603–1602 cm⁻¹ and 1601–1600 cm⁻¹ and constituted around 7% and 10% of the amide I area, respectively. These spectral

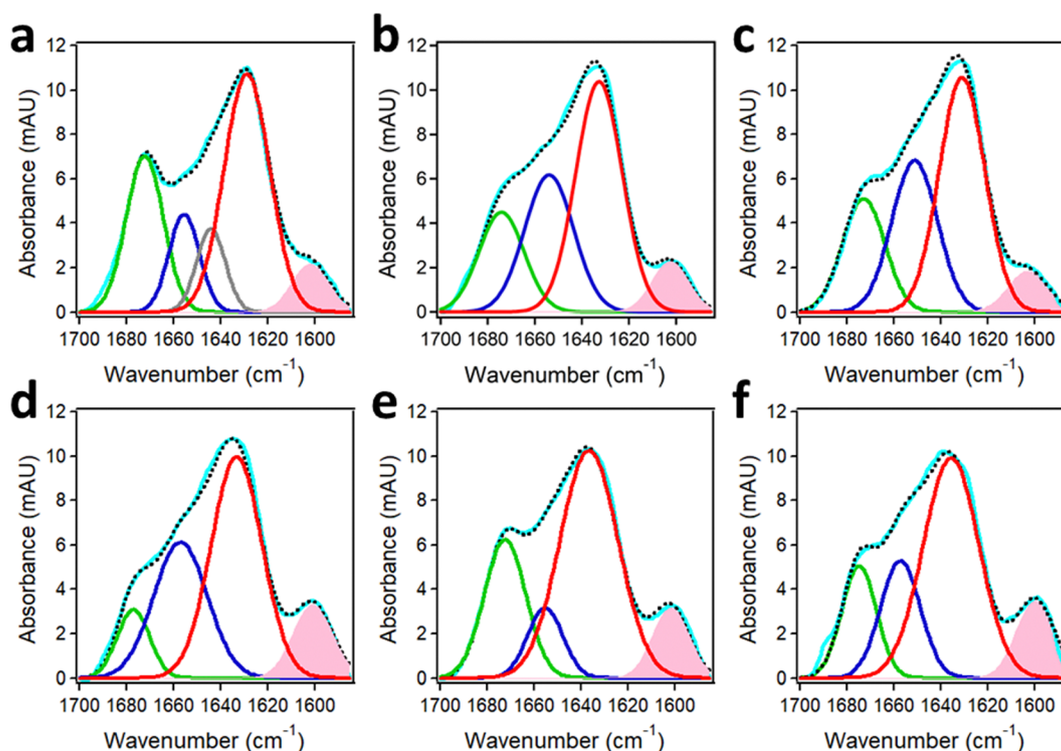


Fig. 10 Structures of A β _{1–42} and A β _{pE3–42} from FTIR spectra of segmentally ¹³C, ¹⁵N-labeled peptides. Peak-fitted FTIR spectra of K¹⁶L¹⁷V¹⁸-labeled A β _{1–42} (a), K¹⁶L¹⁷V¹⁸-labeled A β _{pE3–42} (b), their 1 : 1 combination (c), V³⁶G³⁷G³⁸V³⁹-labeled A β _{1–42} (d), V³⁶G³⁷G³⁸V³⁹-labeled A β _{pE3–42} (e), and their 1 : 1 combination (f) incubated in 50 mM NaCl + 50 mM Na,K-phosphate buffer (pD 7.2) for 1 hour. Color code: measured spectrum is cyan; the turn, α -helix, irregular, and β -sheet components are shown in green, blue, gray, and red lines, respectively. The component generated by the ¹³C, ¹⁵N-labeled segment is shaded pink. The sum of all components is shown in black dotted line. Total peptide concentration was 50 μ M. Adapted from ref. 44, with permission.



features can be used to determine the mode of peptide aggregation. As indicated in Section 2.2, $^{13}\text{C}=\text{O}/^{13}\text{C}=\text{O}$ vibrational dipolar coupling, which would occur in case of in-register cross- β -sheet formation, generates lower frequency signals ($1594\text{--}1591\text{ cm}^{-1}$) whereas the $^{13}\text{C}=\text{O}/^{12}\text{C}=\text{O}$ coupling produces signal in the $1604\text{--}1601\text{ cm}^{-1}$ range.^{46,47,50,51} This implies that the labeled stretches are involved in β -sheet structure as well, thus increasing to total β -sheet to around 60%. More importantly, none of the isotopically labeled segments, *i.e.* the 16–18 or the 36–39 stretches, are involved in coupling with labeled segments, excluding the parallel in-register β -sheet formation. Instead, both peptides form β -hairpin-like structures upstream of residue 16 stabilized by intramolecular H-bonding. The α -helix is most likely located at the N-terminus and the turn is most likely between the two β -strands.⁴⁴ The aggregation occurs *via* non-H-bonding interactions between these hairpin-like structures, which can exert the cytotoxic effect through various mechanisms including membrane insertion and permeabilization.^{151,177}

6. Conclusions and outlook

6.1. Strengths and limitations

Isotope-edited FTIR spectroscopy offers a variety of options to analyze protein–protein and protein–membrane interactions. One of the strengths of this approach is that isotope labeling, *i.e.* addition of one or two neutrons in the nucleus of an atom, preserves the chemical (and most of physical) properties of the amino acid as opposed to labeling with relatively bulky exogenous molecular probes such as fluorophores or paramagnetic agents often used in fluorescence, EPR, or NMR spectroscopy. Interactions between two proteins and the accompanying conformational changes in each protein can be probed by studies on binary samples composed of an unlabeled and a uniformly ^{13}C or $^{13}\text{C},^{15}\text{N}$ -labeled protein.^{20,43,157,178} Domain-specific, segmental, or single amino acid labeling of proteins with ^{13}C , $^{13}\text{C},^{15}\text{N}$, or $^{13}\text{C}=\text{O}$ allows for amide I spectral resolution and structural characterization of the labeled site or stretch.^{21–23,41,162–164,166,171,179} Spectral shifts in the amide II band provide an opportunity to assess the rates of the amide HX and to gain information about the solvent accessibility and the structural dynamics of the labeled and unlabeled parts of the protein. In addition, vibrational coupling effects between amide oscillators of distinct isotope content provide information on the packing of the labeled and unlabeled segments relative to each other and hence on the overall tertiary fold of the protein.^{44,48,49,57,58} The sensitivity of the vibrational frequency to the strength of the involved covalent bonds, and consequently to the strength of H-bonding, permits distinguishing between very similar secondary structures, such as α_{I} - and α_{II} -helices, which is difficult to accomplish by other methods.¹⁸⁰ Polarized ATR-FTIR studies on segmentally labeled proteins reconstituted in lipid bilayers or multilayers allow determination of the orientation of the labeled segment relative to the membrane, thereby providing clues to the mode of its interaction with the membrane.¹⁰⁶

Studies on isotope-labeled membrane lipids produce useful information on membrane structure. FTIR studies on glycerophospholipids ^{13}C -labeled at *sn*-1 or *sn*-2 ester carbons yielded information on the hydration properties of the membrane surface and the orientations of the carbonyl and other chemical groups of the lipid headgroup, as well as changes of these parameters during thermal phase transitions and interdigitated structure formation.^{72–74} Data on membranes composed of selectively deuterated lipids at the *sn*-1 or *sn*-2 hydrocarbon chains can be used to determine the details of the arrangement and packing of lipids in a bilayer structure.⁷⁴ As shown in Section 3.2, ATR-FTIR studies on supported membranes composed of an unlabeled and a chain-deuterated lipid can identify which component is preferentially hydrolyzed by PLA₂.¹⁹ On the other hand, selective deuteration of one of the chains allows deciphering which reaction product (fatty acid or lyso-phospholipid) is mostly removed from the membrane upon hydrolysis by PLA₂.¹⁹

Despite these advantages, isotope-edited FTIR spectroscopy has its limitations. One inherent issue is the spectral overlap between the amide I signals generated by the unlabeled and ^{13}C or $^{13}\text{C},^{15}\text{N}$ -labeled chains. $^{13}\text{C}=\text{O}$ labeling results in additional downshift of the amide I signal and thereby may offer a better spectral resolution (see Section 2.2). Single amino acid labeling may be problematic for several reasons: (a) technical difficulties of incorporation of a labeled amino acid in large proteins, (b) too weak signal compared to the whole amide I band, and (c) uncertainty regarding the secondary structure based on a local amide I mode generated by one amino acid. Solutions to (a) are described in the next section that describes nonsense suppression strategies. Even when the amide I signal of a single amino acid is reliably resolved, its spectral shift and its relation to the local secondary structure is not straightforward because the secondary structure is determined not by the frequency of an isolated amide oscillator (local mode) but by vibrational coupling with neighboring oscillators (normal mode).^{21,57,58} The $^{12}\text{C}/^{13}\text{C}$ vibrational coupling occurs in β -sheet but not in α -helix conformation. Therefore, the signal generated by a single ^{13}C -labeled amino acid will be diagnostic for β -sheet structure but cannot be used to characterize α -helix structure or orientation.^{46,181} Production of proteins isotopically labeled at single or multiple sites by the synthetic method or *via* nonsense suppression may present useful solutions to such problems, as discussed in the next section.

6.2. Future directions

Coupling of FTIR spectroscopy with atomic force microscopy (AFM-IR) is a powerful technique that provides spatial resolution of $\leq 10\text{ nm}$ and thereby allows secondary structural analysis of protein samples or segments of distinct morphologies as opposed to the averaged FTIR spectra. Formation of prefibrillar oligomers and mature fibrils of the A β peptide,^{182,183} α -synuclein,¹⁸⁴ or transthyretin¹⁸⁵ during the process of aggregation have been resolved by this method. Through improvements of the stability, sensitivity, and accuracy of AFM-IR,



single protein molecule resolution has been achieved.¹⁸⁶ This technique has been used to analyze the response of cancer cells to drugs at single cell level and at nanoscale resolution.¹⁸⁷ Further developments in this area may involve the combination of isotope-edited protein samples with AFM-IR. For example, the molecular compositions of protein aggregates or the structural transitions in individual proteins during functional interactions can be resolved at single molecule spatial resolution.

Isotopic labeling of selected domains of proteins using techniques such as native chemical ligation represents an attractive strategy to discern domain-specific structural changes during functional turnover.¹⁷¹ For proteins composed of disulfide bridged domains, such as the insulin receptor family proteins, the protein can be produced in unlabeled and uniformly ¹³C-labeled forms followed by disulfide reduction and re-association of the unlabeled hormone-binding α -subunit with the ¹³C-labeled tyrosine-kinase β -subunit.¹⁸⁸ Various cell-based or cell-free expression systems for functional mammalian proteins are available that are suitable for post-translational modifications such as glycosylation. These include HEK293 cells, insect cells, wheat germ cells and cell-free systems such as rabbit reticulocyte or HeLa cell extracts,^{189,190} which can be supplemented with isotopically labeled amino acids or labeled nutrients such as ¹³C₆-D-glucose. ATR-FTIR studies on the hormone-activation of a domain-specific isotope labeled membrane-reconstituted receptor will reveal the dynamic conformational changes in each domain reflecting the transmission of the hormone signal across the membrane. The peptide hormone itself may be produced in ¹³C=¹⁸O isotopic form to move its amide I signal further away towards lower frequencies.¹⁶⁴

Isotope labeling of one or several amino acids in a protein is crucial for obtaining site-specific structural information. A rapidly developing protein engineering strategy is the use of a genetically modified cell or cell-free expression system containing a stop (nonsense, termination) codon in a given gene, such as TAG (amber), at the site of interest and a nonsense suppressor tRNA aminoacylated with a labeled amino acid and harboring an anticodon complementary to the stop codon (CUA, in this case).¹⁹¹⁻¹⁹⁴ The selected stop codon as well as the suppressor tRNA/amino acyl tRNA synthetase (aaRS) pair should be orthogonal (heterologous) to the host organism or the cell-free system to prevent cross-reaction. In the semisynthetic method, the tRNA_{CUA} lacking the last two nucleotides is prepared by *in vitro* transcription with T7 RNA polymerase and the synthetic, labeled amino acid dinucleotide is appended to the 3' end with the T4 RNA ligase.^{192,195} A similar method uses flexible tRNA acylation ribozymes coupled with a translation system containing the activated, labeled or otherwise modified amino acid.^{196,197} More than one labeled amino acids can be incorporated in a protein by using other stop codons such as TAA (ochre) or TGA (opal) coupled with respective suppressor tRNAs, *i.e.* tRNA_{UUA} or tRNA_{UCA} provided the expression system is using a third stop codon.^{198,199} Artificial, such as quadruplet codons can be introduced alone or in tandem with an orthogonal stop codon resulting in double nonsense/frameshift suppression and incorporation of two labeled or unnatural

amino acids allowing for simultaneous targeting of two sites in a protein.^{191,193,200} Three or four noncanonical amino acids have been incorporated in one protein chain by using all three natural stop codons¹⁹⁴ or by engineering four suppressor tRNA/aaRS pairs targeting four frameshift quadruplet codons.²⁰¹ Future efforts will likely expand the scope of protein engineering allowing facile production of proteins labeled at multiple sites through orthogonal tRNA/aaRS pairs or chemically aminoacylated tRNAs that work efficiently in mammalian expression systems, which will enhance the ability of isotope-edited FTIR spectroscopy to gain higher resolution structural insight into the function of important proteins.

Abbreviations

aaRS	Amino acyl tRNA synthetase
A β	Amyloid β
AD	Alzheimer's disease
AFM	Atomic force microscopy
ATR	Attenuated total reflection
FTIR	Fourier transform infrared
DPPC	Dipalmitoylphosphatidylcholine
DPPG	Dipalmitoylphosphatidylglycerol
HO	Harmonic oscillator
HX	Hydrogen-deuterium exchange
IBS	Interfacial binding site
IR	Infrared
IRE	Internal reflection element
PLA ₂	Phospholipase A ₂
PLB	Phospholamban
SR	Sarcoplasmic reticulum

Data availability

No primary research results, software or code have been included and no new data were generated or analyzed as part of this review.

Conflicts of interest

The author declares no conflict of interest.

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