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Isotope-edited IR spectroscopy for the study of membrane proteins
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Fourier transform infrared (FTIR) spectroscopy has long been a powerful tool for structural analysis of membrane proteins. However, because of difficulties in resolving contributions from individual residues, most of the derived measurements tend to yield average properties for the system under study. Isotope editing, through its ability to resolve individual vibrations, establishes FTIR as a method that is capable of yielding accurate structural data on individual sites in a protein.

Introduction

The electromagnetic spectrum between visible and microwave radiation is known as infrared light. Its wavelength range (750 nm–1 mm) corresponds to energies that overlap the vibrational transitions in molecules. These transitions, in turn, provide a wealth of information about the sample in question — for example, couplings between vibrational modes due to proximity; frequency shifts caused by hydrogen bonding or the particular electrostatic properties of the environment; and the orientation of a particular group relative to the laboratory frame. This, together with the sensitivity of modern spectrometers makes Fourier transform infrared (FTIR) spectroscopy a popular tool for molecular analysis.

In applying FTIR spectroscopy to the study of biological samples two problems are frequently encountered. The first is the water stretching and bending modes at ca. 3200–3700 cm⁻¹ and 1600–1630 cm⁻¹, respectively, which dominate the IR spectra of any dilute biological sample. When studying water-soluble proteins, it is possible to overcome this problem partially through the use of highly concentrated specimens, short sample path lengths, and/or use of D₂O. In contrast, membrane proteins are particularly suited for examination by FTIR spectroscopy. Here, water opacity is much less of a problem because the lipid bilayer is the dominant solvent of the protein. Hence, after the removal of bulk water, membrane proteins can readily be studied by FTIR in their native environment — the lipid membrane.

The second problem that limits the usefulness of IR spectroscopy as a tool for the structural study of biological macromolecules is spectral resolution. In other words, the width of a single vibrational mode is often significantly larger than the differences between individual vibrations. Therefore, traditional IR spectroscopy yields information that pertains to the average properties of a protein or that which is obtained because of coupling between numerous modes. The advent of isotope editing in IR spectroscopy of proteins has alleviated this problem to an extent, because of the ability of isotopomers to shift the vibrational frequency to a 'transparent' region of the spectrum. If we assume that the vibrational mode between atoms a and b behaves as a harmonic oscillator, its frequency, $\omega_{a,b}$, will change upon isotope editing of atom a to c and b to d as follows:

$$\frac{\omega_{a,b}}{\omega_{c,d}} = \sqrt{\frac{m_a m_b (m_a + m_b)}{m_c m_d (m_a + m_b)}}$$

whereby $m_a$, $m_b$, $m_c$ and $m_d$ are the masses of atoms a, b, c and d. Atoms a ↔ c and b ↔ d are isotopomeric pairs.

Note that the above equation indicates that the frequency shift is highly dependent on relative, not absolute, mass change. For example, the amide A vibrational mode (mostly the N–H stretch) at ca. 3300 cm⁻¹ will shift –890 cm⁻¹ upon deuteration ($^{14}$N–$^1$H → $^{14}$N–$^2$H) but only –7 cm⁻¹ upon isotope labeling of the nitrogen ($^{14}$N–$^2$H → $^{15}$N–$^2$H). This is due to a relative change of 100% with regard to the hydrogen atom mass in the first case, versus a relative change of 7% with regard to the nitrogen atom mass, in the second case. Thus, because of its ability to isolate an individual vibration, isotope-edited FTIR spectroscopy is a powerful tool that can yield precise information regarding the labeled site. Moreover, it does so with minimal interference, because isopomer species are normally completely silent with respect to the chemical nature (e.g., structural perturbation) of the examined protein.
Here, I review some of the recent results from studies that used isotope-edited infrared spectroscopy of membrane proteins. Because of length restrictions I am unable to cover all the relevant studies. The data are presented such that studies that utilize similar spectroscopic tools are grouped together, rather than those that study similar biological systems.

Secondary structure
In proteins, the most prominent absorptions of infrared light are those of the peptidic backbone, and are therefore called the amide modes. The most dominant amongst these is the amide I mode, which has been studied extensively [1] and shown to be dominated by the C=O stretching motion. The absorption of the amide I mode occurs between 1600 and 1700 cm⁻¹ (6.25–5.88 μm), and its frequency has long been used as a marker for the secondary structure of the protein in question: modes centered around 1655 cm⁻¹ correspond mostly to α-helical structures, whereas a central mode centered around 1630 cm⁻¹ and a minor mode at ca. 1690 cm⁻¹ correspond mostly to β-sheet secondary structure elements (for reviews, see [1–4]).

Because of its ability to derive secondary structure information, traditional FTIR spectroscopy is an alternative to circular dichroism (CD) spectroscopy. FTIR is particularly well suited to the study of membrane proteins because it does not suffer from the differential absorption flattening observed in CD spectra of membrane proteins [5]. Furthermore, the advent of site-specific isotopic incorporation in peptides has enabled the first rapid site-specific secondary structure measurement [6].

The procedure for using isotope-edited IR spectroscopy to determine secondary structure involves labeling the protein of interest with an amino acid bearing an isotope-labeled carboxylic group (¹³C=¹⁸O or ¹³C=¹⁶O), measuring the FTIR spectrum of the resultant protein in the amide I region (1560–1700 cm⁻¹), and determining the frequency of the isotope-edited peak. The ~40 cm⁻¹ shift that results from ¹³C=¹⁶O incorporation yields an isotope-edited mode present as a shoulder on the natural (unlabeled) ¹²C=¹⁶O peak (see Figure 1). Furthermore, the natural abundance of ¹³C (1.1%) maintains that in any protein with 90 amino acids or more, the contribution of non-labeled ¹³C is as much, or more than that of the label. To overcome the above shortcomings, we often utilize the ¹³C=¹⁸O label [7,8]. The frequency shift of ~65 cm⁻¹ places the isotope-edited absorption band so that it is nearly base-line resolved from the much larger ¹²C=¹⁶O mode, and the natural abundance of the ¹³C=¹⁸O label (1.1% × 0.2% = 0.0026%) is negligible.

Based on the nature of the label, one determines the frequency from which it ‘originated’, and its corresponding secondary structure. For example, a ¹³C=¹⁶O isotope-edited peak at 1615 cm⁻¹ indicates that the site is helical, because without isotopic labeling it would resonate at 1655 cm⁻¹, which is typical of α-helical segments [1,2]. In large proteins, it is useful to compare the results with spectra of an unlabeled protein to verify that the suspected peak originates from the isotopically labeled amino acid. Finally, we note that in the above procedure, any coupling differences between isotope and non-isotope bands are neglected, thereby assuming similar frequency shifts regardless of the secondary structure.

Examples
Site-specific analysis of secondary structures is particularly useful in proteins that are composed from different secondary structure elements. For example, work from two groups on the HIV-1 gp41 fusion protein has been able to identify topological elements in the protein that might change upon interaction with the lipid bilayer. Because the fusion protein has been shown to contain
both α-helical and β-sheet elements, both Waring and co-workers [9,10] and Shai and co-workers [11] have labeled the amino terminal residues of the protein with $^{13}$C–$^{18}$O. This enabled both groups to assign a β-sheet secondary structure to the N-terminal domain of the protein, although there is still some disagreement as to whether the β-sheet is composed of parallel [11] or anti-parallel [9] strands.

Axelsen and co-workers have used internal reflection spectroscopy to analyze a hexapeptide containing $^{13}$C–$^{18}$O labels, that partitions into the lipid bilayer [12]. Excitonic model simulations and comparisons with the measured spectra enabled the authors to show conclusively that the peptide forms an anti-parallel β-sheet in the membrane, and that the spectra are modulated by the vibrational couplings both in individual strands and between them. Barry and co-workers [13], on the other hand, have analyzed a significantly larger protein: the lactose permease (LacY), which contains 417 amino acid residues. The authors performed a ‘reaction-induced difference FTIR’, in which a reaction is initiated by changing the sample conditions and the resulting difference in the FTIR spectra is recorded. By using LacY uniformly labeled with $^{13}$C and $^{15}$N the authors were able to assign peaks in the difference spectra at 1765–1730 cm$^{-1}$ to carboxylic groups as well as to assign additional spectral features to contributions of other side chain moieties such as lysine, arginine, histidine and the amide residues. Future studies employing site-specific labeling strategies may enable the exact pinpointing of residues responsible for the derived spectral features in the reaction-induced difference spectra.

Structure and orientation

The absorption of light is proportional to the squared scalar product of the electric field vector and the dimensionless transition dipole moment of the vibrational mode. Thus, polarized spectroscopy of a macroscopically aligned sample can determine the orientation of a particular chromophore if the geometry of the transition dipole moment in relation to the chromophore’s principal axis is known. Because it is very easy to macroscopically align membrane proteins by depositing them in membrane stacks, this methodology has been used extensively in infrared spectroscopy to determine the average tilt angle of helices based on a simple linear dichroism experiment (see Figure 2 for a schematic representation). Specifically, the measurement of the absorption ratio between parallel and perpendicular polarized light (relative to the membrane plane) can yield the orientation of the molecule in question (for review see [14–16]).

One limitation complicating the above approach is sample disorder, which results in the fact that the measured dichroism is no longer solely a function of the orientation of the sample, but also of its level of order.

Although it is possible to model this disorder using different distribution functions [17–19], it remains essential to experimentally determine the sample disorder to deconvolve the orientational information from the measured dichroism. Without disorder deconvolution, FTIR dichroism will only yield an order parameter, which is of little use for high-resolution structural analysis.

Recent studies have used X-ray reflectivity to independently measure the membrane mosaicity, and as such have presented a simple route for deriving accurate orientational information directly from FTIR dichroism measurements [20]. The above methods, in combination with isotopic labeling, enable one to derive accurate orientational restraints for membrane proteins in its native environment [21,22]. These restraints can later be used to obtain highly reliable structural models of the protein in question (e.g. [7,23–26,27]).

Examples

Two studies from the Arkin group [26,27] demonstrate the utility of site-specific FTIR spectroscopy in deriving accurate orientational restraints for membrane proteins. In analyzing the transmembrane helical bundle formed by the T-cell receptor CD3-ζ component, the authors employed 11 different peptides, each containing a $^{13}$C–$^{18}$O label in a unique position. The site-specific dichroism for each of the peptides was used to derive an orientational restraint for each of the respective carbonyl groups. These restraints were then implemented as orientational refinement energy terms in molecular dynamics simulations, leading to a structure for the
protein complex that correlated well with evolutionary conservation information [28]. The data were sufficiently accurate to detect a kink in the transmembrane helices that resulted in asymmetrical water accessibility to the amino and carboxy termini of the helical bundle. This pattern of water accessibility was later confirmed by 2D IR studies, as elaborated below [29**].

In a similar effort, 10 different peptides, each with a uniquely inserted $^{13}$C$\equiv^{18}$O label, were used to study the trimerizing MHC class II-associated invariant chain transmembrane domain structure [26]. The dichroisms of the isotope-edited modes were obtained for each of the labels and were used to derive a structure of the transmembrane helical bundle as in the case of CD3-$\zeta$ (see above). Once again, the structure correlated well with experimental data, placing key residues such as Gln47 and Thr50 in the core of the bundle.

A different type of study was recently conducted [30**] to experimentally measure the strength of the glycine Ca-H\cdots O=C hydrogen bond predicted to be present in transmembrane helices [31] (see Figure 3). The study employed two different peptides, corresponding to the dimeric and the (mutant) monomeric transmembrane domains of the human glycoporphin A [32]. Gly79 was exchanged for a glycine in which the two Ca hydrogens were replaced with deuteriums. This enabled the authors to selectively measure the glycine CD$_2$ stretching frequency and to determine the effects of hydrogen bond formation thereupon. The results of the study show a 6 cm$^{-1}$ difference between the peak of the monomeric and the dimeric glycine CD$_2$ stretching mode, indicating that in the dimeric form the methylene group is hydrogen bonded to the C=O group in the opposing helix, as predicted [31]. Using an empirical correlation obtained from a large number of compounds [33], the authors were able to derive the enthalpy of the Ca-H\cdots O=C bond as 0.88 kcal/mol [30**].

**Mechanistic insights**

FTIR spectroscopy has long been a particularly powerful tool for analyzing protein function and mechanism of action. This is because vibrational modes may be influenced by factors such as hydrogen bonding, solvation and bond deformation, thereby serving as excellent probes of the local structure in areas of interest. The ability to spectroscopically select a certain site in the protein to observe, through isotopic labeling, makes this technique particular powerful. This normally means that the most challenging part in the experimental scheme is the introduction of the label at the location of choice (see Conclusions section).

**Examples**

The light-driven H$^+$ pump bacteriorhodopsin (BR) has been a popular system for biophysical investigation. Its photocycle in particular has been studied in detail, in the context of the high-resolution structure of the protein [34]. Recent studies from the Braiman [35*], Kandori [36*] and Rothschild [37**] groups (amongst others) utilizing isotope-edited FTIR demonstrate the advantageousness of the method. Rothschild and co-workers have studied the influence of incorporating selenomethionine in a cell-free expression system on the spectral properties of BR [37**]. To show that the particular modes at 1284 cm$^{-1}$ were due to the incorporated selenomethionine and not to tyrosine, they compared the spectra with those obtained from proteins that were labeled with selenomethionine and L-tyrosine-[ring]-d$_4$. This enabled the authors to show conclusively that the introduced selenomethionine is responsible for the modes at 1284 cm$^{-1}$.

The groups of Kandori [35*] and Braiman [36*] have used arginine labels to study the BR photocycle. Kandori and co-workers labeled BR with $\nu$(1,2)-$^{15}$N-arginine and were able to identify perturbations in the arginine hydrogen bonding [35*]. Moreover, analysis of the different states in the BR photocycle enabled the authors to show that in the photocycle intermediate L this perturbation is relaxed. Braiman and co-workers used the same selective arginine label as well as a uniformly $^{15}$N-labeled arginine in their study [36*] and were able to assign a prominent mode at 1555 cm$^{-1}$ as the perturbed arginine C=N stretch in the BR photocycle intermediate M. Comparison with model compounds has indicated that this perturbed mode might occur due to light-induced deprotonation of the arginine guanidino moiety.
2D IR spectroscopy
One of the most exciting advances in the field of infrared spectroscopy has been the development of multidimensional methods (see for review [38]). In analogy to multidimensional NMR, 2D IR spectroscopy can be used to simplify the interpretation of complex single dimensional spectra, providing information about the coupling (and by inference, the distance and angles) between individual vibrations. In addition, 2D IR has the ability to probe the dynamics of molecules in time regimes that are inaccessible to most other methods, providing information on the dynamics of structures and environments down to the femtosecond regime (see Figure 4).

Examples
Because multidimensional IR spectroscopy is a very recent technique, its application to the study of proteins in general, and membrane proteins in particular has been limited. However, two recent studies from the Zanni group on the transmembrane domain of CD3-ζ demonstrate the promise that the approach possesses. In the first study of a membrane protein by 2D IR [39], the authors measured the homogeneous and inhomogeneous line-widths of the single $^{13}$C-$^{18}$O labeled site and compared that with the entire peptide amide I (arising from the $^{12}$C-$^{16}$O modes). This study was later extended to 11 different $^{13}$C-$^{18}$O residues along the entire length of the transmembrane peptide [29**]. The analysis pointed to the fact that although all sites exhibit the same homogeneous broadening, they differed substantially in their inhomogeneous linewidths. Comparison of experiments with simulations indicate water accessibility of the protein backbone as the primary cause that affects the dynamical behavior of the different sites. The differential water accessibility correlated well with the structural model of the protein [27**].

Exposure to solvent
Amide H+/D+ exchange is another example of the use of isotopes in FTIR spectroscopy. This method can readily determine the percentage of a membrane protein that is in direct contact with the aqueous solvent [40**]. The principle of the method is that amide H+/D+ exchange will not take place in the hydrophobic milieu of the lipid bilayer. Thus, monitoring the reduction of the amide II mode (mostly the amide N–H deformation mode) at ca. 1540 cm$^{-1}$, which is caused by exposure of the sample to D$_2$O, will yield an accurate assessment of the number of residues that are accessible to the solvent (see Figure 5 for an example).

Examples
Simple hydropathy algorithms predict the topology of a membrane protein, and by inference, the number of residues that span the membrane. However, in numerous membrane proteins, large aqueous cavities and vestibules result in many of the residues in the transmembrane...
helices undergoing H+/D+ exchange as well. Lancaster and co-workers [41] have recently studied Wolinella succinogenes quinol:fumarate reductase focusing on the role of a key carboxylic residue (Glu-C180) in the function of the protein. Using site-directed mutagenesis along with H+/D+ exchange the authors were able to show that Glu-C180, located in the transmembrane segment of the protein, undergoes protonation and deprotonation as evidenced by peak changes at 1733–1741 cm⁻¹. Goormaghtigh and co-workers [40] used H+/D+ exchange to study structural dynamics in the gastric H⁺,K⁺ ATPase. In their analysis, using attenuated total internal reflection spectroscopy, the authors showed that the H+/D+ exchange kinetics are intimately correlated with the molecular movements of the protein. Because these movements may be correlated with the catalytic cycle of the protein, they provide insight into the kinetics and structural transitions that the enzyme undergoes.

In a demonstration of technique sensitivity, Vogel and co-workers [42] have used microscope FTIR spectroscopy to observe nanogram quantities of the serotonin and acetylcholine receptors in fully hydrated protein films. From H+/D+ exchange analysis the authors were able to determine the secondary structure of the proteins and to show that the water-accessible percentage of both proteins is similar, as predicted in previous studies.

Conclusions

From the numerous examples of experiments now using isotope-edited FTIR, it is clear that it is a powerful technique that is particularly applicable to the study of membrane proteins, and that it can provide rich information on the structure and mechanism of the protein in question. One obvious limitation that must be considered is the ability to label the target protein. Two categories of proteins should therefore be considered for FTIR analysis; those that can, and those that cannot be made synthetically. Inserting a label into a synthetic protein is trivial — it is simply introduced at will during the synthesis. The problem with this approach is that ‘straight-forward’ chemical synthesis is limited to proteins smaller than 60–70 residues. However, chemical ligation can be used to extend this upper limit significantly [43], as shown by the recent total synthesis of functional mechanosensitive channels from Escherichia coli and Mycobacterium tuberculosis, containing 136 and 151 residues, respectively [44]. Isotopic labeling of proteins that cannot be made synthetically is much more of a challenge. Approaches that have been used to address this technical obstacle are metabolic labeling of unique amino acids [35,36,37] and tRNA engineering [45,46].

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