Fourier Transform Infrared Spectroscopic Analysis of Protein Secondary Structures

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Abstract

Infrared spectroscopy is one of the oldest and well established experimental techniques for the analysis of secondary structure of polypeptides and proteins. It is convenient, non-destructive, requires less sample preparation, and can be used under a wide variety of conditions. This review introduces the recent developments in Fourier transform infrared (FTIR) spectroscopy technique and its applications to protein structural studies. The experimental skills, data analysis, and correlations between the FTIR spectroscopic bands and protein secondary structure components are discussed. The applications of FTIR to the secondary structure analysis, conformational changes, structural dynamics and stability studies of proteins are also discussed.

Keywords

FTIR; protein structure; protein dynamic

Infrared (IR) spectroscopy is one of the oldest and well established experimental techniques for the analysis of secondary structure of polypeptides and proteins [1−5]. The use of stable and powerful laser has led to the development of the Fourier transform (FT) method for IR data acquisition and reliable digital subtraction. The availability of modern computers has enabled the rapid and powerful FTIR data processing and conversion. FTIR spectroscopy is recognized as a valuable tool for the examination of protein conformation in H2O-based solution, as well as in deuterated forms and dried states, resulting in a greatly expanded use in studies of protein secondary structure and protein dynamics in the past decade [4−17]. Although X-ray crystallography provides the most detailed information concerning positions of individual atoms in the protein structure, it is not, however, possible for all proteins to form a quality crystal for such analysis. In addition, the crystallographic data of a protein cannot be easily extrapolated to the dynamic properties of the proteins in solutions. Nuclear magnetic resonance spectroscopy can be an alternative to X-ray crystallography in solution, but the interpretation of nuclear magnetic resonance spectra of a large protein is a very cumbersome process [18]. Thus, the vibrational spectroscopies, such as FTIR and circular dichroism (CD), are still important and commonly used techniques for protein structure and dynamics studies.

FTIR spectroscopy is a measurement of wavelength and intensity of the absorption of IR radiation by a sample. The IR spectral data of high polymers are usually interpreted in terms of the vibrations of a structural repeat unit [2,3,19]. The polypeptide and protein repeat units give rise to nine characteristic IR absorption bands, namely, amide A, B, and I-VII. Of these, the amide I and II bands are the two most prominent vibrational bands of the protein backbone [3−5]. The most sensitive spectral region to the protein secondary structural components is the amide I band (1700−1600 cm⁻¹), which is due almost entirely to the C=O stretch vibrations of the peptide linkages (approximately 80%). The frequencies of the amide I band components are found to be correlated closely to each secondary structural element of the proteins. The amide II band, in contrast, derives mainly from in-plane NH bending (40−60% of the potential energy) and from the CN stretching vibration (18−40%) [3], showing much less protein conformational sensitivity than its amide I coun-
Table 1 Characteristic infrared bands of peptide linkage

<table>
<thead>
<tr>
<th>Designation</th>
<th>Approximate frequency (cm(^{-1}))</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Amide A</td>
<td>3300</td>
<td>NH stretching</td>
</tr>
<tr>
<td>Amide B</td>
<td>3100</td>
<td>NH stretching</td>
</tr>
<tr>
<td>Amide I</td>
<td>1600–1690</td>
<td>C=O stretching</td>
</tr>
<tr>
<td>Amide II</td>
<td>1480–1575</td>
<td>CN stretching, NH bending</td>
</tr>
<tr>
<td>Amide III</td>
<td>1229–1301</td>
<td>CN stretching, NH bending</td>
</tr>
<tr>
<td>Amide IV</td>
<td>625–767</td>
<td>OCN bending</td>
</tr>
<tr>
<td>Amide V</td>
<td>640–800</td>
<td>Out-of-plane NH bending</td>
</tr>
<tr>
<td>Amide VI</td>
<td>537–606</td>
<td>Out-of-plane C=O bending</td>
</tr>
<tr>
<td>Amide VII</td>
<td>200</td>
<td>Skeletal torsion</td>
</tr>
</tbody>
</table>

Data are from Elliott and Ambrose [2], Krimm and Bandekar [3], Banker [20] and Miyazawa et al. [21].

Protein FTIR Data Analysis and Band Assignment

Data analysis

High sensitivity to small variations in molecular geometry and hydrogen bonding patterns makes the amide I band uniquely useful for the analysis of protein secondary structural composition and conformational changes [4, 11]. In the amide I region (1700–1600 cm\(^{-1}\)), each type of secondary structure gives rise to a somewhat different C=O stretching frequency due to unique molecular geometry and hydrogen bonding pattern. However, other than the distinctive absorbance maxima, the observed amide I bands of proteins are usually featureless, due to the extensive overlap of the broad underlying component bands, which lie in close proximity to one another and are instrumentally unresolvable. Thus, mathematical methods such as resolution-enhancement technique are necessary to resolve the individual band component corresponding to specific secondary structure. Fig. 1(A) is the IR spectrum of cAMP receptor protein (CRP) in buffer. The individual underlying
components can not be visualized without resolution enhancement.

Mathematical data analysis methods can be used to “enhance” the resolution of the protein spectrum, allowing the intrinsically broad components to be narrowed and separated beyond the instrument resolution [1,22]. The mathematical band-narrowing process does not actually increase the instrumental resolution, but rather increases the degree of separation by narrowing the half-bandwidth of individual components for easier visualization. This band-narrowing process is achieved at the expense of spectral quality of the original band, which leads to a degradation of signal-to-noise ratio. Several methods have been developed to estimate quantitatively the relative contributions of different types of secondary structures in proteins from their IR amide I spectra in solution, including FSD-curve fitting [7,22], second derivative analysis [4,7,11], partial least-squares analysis [23], and data basis analysis [24]. The FSD-curve fitting and second derivative analysis are the two most popularly used methods.

The theoretical backgrounds of IR data handling techniques, including FSD and second derivative, have been discussed in detail by Susi and Byler [4,6]. The FSD method was also reviewed by Surewicz and Mantsch [18]. The key meaningfulness of the FSD method is to select the conditions that achieve the maximum band narrowing while keeping the increase in noise and the appearance of side-lobes at minimum [6,14,22]. This method is based on the assumption that a spectrum of single bands (each narrow band is characteristic of a secondary structure) is broadened in the liquid or solid state. Therefore, the bands overlap and can not be distinguished within the amide envelope. An alternative approach to “enhance” the resolution of overlapping IR bands is based on the generation of nth order derivative band files. This can be carried out in the frequency (wavenumber) domain of the spectrum. Second derivative spectra allow the identification of various secondary structures present in the protein [6]. Most of the peak positions are easily found in the second derivative spectra. An improved method for carrying out second derivative analysis has established the utility of the method for obtaining quantitative as well as qualitative determination of α-helix, β-sheet, random and turn structures [11]. A curve fitting procedure can be applied to calculate quantitatively the area of each component representing a type of secondary structure [4,8−11]. Fig. 1(B) shows the second derivative spectrum of CRP, which was obtained according to the methods of Dong et al. [11]. Both FSD and second derivative methods were included in BOMEM GRAMS/32 software. Second derivative analysis was used in this work. Manipulation procedures were carried out in accordance with Dong et al. [11].

The overlap of secondary structural components is significant in the amide I region, even after mathematical resolution enhancement. Some investigators considered that the valley between two adjacent peaks of equal intensity must be 20% lower than the peak tops that could be resolved [1,25].

**Band assignment**

Quantitative analysis of protein secondary structure is based on the assumption that protein can be considered as a linear sum of a few fundamental secondary structural elements. Comparisons of IR spectra with high-resolution X-ray crystal structures of proteins could establish necessary spectra-structure correlations. Over the years, many correlations between IR spectra and particular protein
structure components have been established. The amide I band components can be assigned by studying their frequency behavior in which the protein secondary structure is known by other techniques. As mentioned above, the amide I band (1700–1600 cm⁻¹) is due mainly to the C=O stretching vibration (approximately 80%) of the amide groups coupled with little in-plane NH bending (<20%) [3]. The extract frequency of this vibration band depends on the nature of hydrogen bonding involving the C=O and NH moieties [3]. In turn, this is determined by the secondary structure adopted by the polypeptide chain, reflecting the backbone conformation and hydrogen-bonding pattern. Strictly speaking, the observed amide I band contours of proteins or polypeptides consist of overlapping component bands, representing α-helices, β-sheets, turns and random structures. It is needed to establish a correspondence between IR spectra and the various types of protein secondary structure. Assignments of the amide I band component to each secondary structure element are available for proteins in both D₂O and H₂O media [4–16].

In D₂O solution, it has been revealed that the broad protein amide I band contours can be decomposed into a number of components. A component centered between approximately 1658 and 1650 cm⁻¹ has been assigned to the α-helix, which is consistent with both theoretical calculation [3] and the observation of bands in the spectra of α-helical proteins [4,26]. Bands near 1663 cm⁻¹ are assigned to 3₁₀ helices [3,4,11], although this structure is rarely found in proteins. More than one β-component has been observed in the spectra of many β-sheet proteins. Bands in the regions of 1640–1620 cm⁻¹ and 1695–1690 cm⁻¹ have been assigned to β-sheet by many authors [3,4,23]. Theoretical calculation of β-sheets also predicts an IR active mode between approximately 1695 and 1670 cm⁻¹ [3]. These α-components are often complicated by the presence of more than one band above 1670 cm⁻¹. The assignment of bands around 1670, 1683, 1688 and 1694 cm⁻¹ to β-turns has been proposed [3]. Turns are also associated with a characteristic band around 1665 cm⁻¹. The unordered conformation (usually referred to as random coil) is usually associated with the IR band between 1640 and 1648 cm⁻¹ [3]. The distinguishing feature of random coil is that it is non-repetitive. Such assignment is supported by the position of a prominent band in the spectra of apparently order-less proteins [7].

H₂O as a solvent is much more preferable than D₂O for studying protein structure [8–11]. D₂O changes the protein properties somewhat in comparison with the native ones. In H₂O solution, the bands between 1654 and 1658 cm⁻¹ are assigned to α-helix, which is supported by human hemoglobin A and bovine Myoglobin proteins, and are expected for proteins with α-helical structure [11]. The bands between 1642 and 1624 cm⁻¹ are assigned to β-sheets components through the IR spectra of immunoglobulin G and concanavalin A, which contain more than 60–70% β-sheet structures and almost no α-helix. In globular protein, it has been observed that approximately 30% of amino acid residues reside in β-turn conformations [11,28]. The bands located at 1688, 1680, 1672, and 1666 cm⁻¹ are assigned to β-turn structures. The characteristic band for random coil conformation can be assigned to the band located at 1648±2 cm⁻¹.

Some minor or rare structures might interfere with the band assignments discussed above. For example, the β-turn band at approximately 1665 cm⁻¹ is near the characteristic IR band representing 3₁₀-helices. Vibrations of some amino acid side chains might make small contributions to the intensity of characteristic protein amide bands. In addition, the experimental procedure might also bring spectral error. All these complications indicate that there is no simple correlation between the IR spectra and secondary structural components. Caution has to be exercised in the interpretation of IR spectra of proteins. Table 2 shows the deconvoluted amide I band frequencies and assignments to secondary structure for proteins in D₂O and H₂O.

| Subtraction of background |

The accuracy of subtraction of large H₂O bands is

<table>
<thead>
<tr>
<th>Table 2</th>
<th>Deconvoluted amide I band frequencies and assignments to secondary structure for protein in D₂O and H₂O media</th>
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<tbody>
<tr>
<td>H₂O†</td>
<td>Mean frequencies</td>
</tr>
<tr>
<td>1624±1.0</td>
<td>β-sheet</td>
</tr>
<tr>
<td>1627±2.0</td>
<td>β-sheet</td>
</tr>
<tr>
<td>1633±2.0</td>
<td>β-sheet</td>
</tr>
<tr>
<td>1638±2.0</td>
<td>β-sheet</td>
</tr>
<tr>
<td>1642±1.0</td>
<td>β-sheet</td>
</tr>
<tr>
<td>1648±2.0</td>
<td>Random</td>
</tr>
<tr>
<td>1656±2.0</td>
<td>3₁₀ Helix</td>
</tr>
<tr>
<td>1667±1.0</td>
<td>β-Turn</td>
</tr>
<tr>
<td>1675±1.0</td>
<td>β-Turn</td>
</tr>
<tr>
<td>1680±2.0</td>
<td>β-Turn</td>
</tr>
<tr>
<td>1685±2.0</td>
<td>β-Turn</td>
</tr>
<tr>
<td>1691±2.0</td>
<td>β-sheet</td>
</tr>
<tr>
<td>1696±2.0</td>
<td>β-sheet</td>
</tr>
</tbody>
</table>

† Data are from Dong et al. [12,13]; ‡ Data from Susi et al. [4,7].
always a concern. H$_2$O has strong IR absorbance with three prominent bands around 3400 (O-H stretching), 2125 (water association), and 1645 cm$^{-1}$ (H-O-H bending). The amide I vibration for proteins absorbs between 1600 and 1700 cm$^{-1}$, overlapping directly with the H$_2$O bending vibrational band at 1645 cm$^{-1}$. The intensity of the water absorbance at 1645 cm$^{-1}$ is approximately an order of magnitude higher than the amide I absorbance of proteins. For IR study of protein in H$_2$O solution, water absorption in the 1600–1700 cm$^{-1}$ region might be the biggest problem. It is much easier if the spectroscopic study is carried out in D$_2$O solution because there is no absorption spectrum of D$_2$O in the region where the amide I and II bands are observed. A large path-length (i.e. 50 µm) IR cell can be used and results in a higher signal-to-noise ratio, a factor of particular importance for proteins of low solubility. However, the amide I band frequencies are strongly affected by the H-D exchanges in the peptide linkages [4, 7]. The effect of these exchanges on protein structural properties is not fully understood, especially under incomplete H-D exchange conditions. Furthermore, because the exchange of D for H can affect the strength and length of hydrogen bonds, it is possible that protein secondary structures might be altered by the replacement of H$_2$O by D$_2$O. Therefore, H$_2$O-based media have the advantage of providing a more native environment.

The greater sensitivity of the FTIR instrument and new H$_2$O substraction program make it possible to obtain a good protein FTIR spectrum in H$_2$O solution [23,29,30]. The problem of water absorption can also be limited by using an IR cell of sufficiently small path-length (6–10 µm) to permit IR radiation passing through the material under observation. By using the same cell to record both the reference spectrum and the spectrum of protein solution under identical scan conditions, the aqueous water contribution can be removed from the spectrum of the protein solution by digital subtraction. Furthermore, progress in the development of methods for spectral data analysis makes it easier to distinguish the individual components within the intrinsically overlapped amide I band contours [11,12]. Nevertheless, the accurate measurement of frequency and intensity of the amide bands is elusive [12]. The short cell path-length makes it more difficult to match the pathlengths of sample and reference cells [12,31]. Typically, the same cell was used to record the spectra of both sample and reference, although the IR cell drying and reloading steps might slightly alter the cell path-length. In order to get a successful subtraction of absorption bands due to liquid water and gaseous water in the atmosphere, one must have a criterion for determining whether or not the absorption by water is correctly compensated, that is, the region of the spectrum where no absorption by the sample but absorption by water is present [31]. Two criteria have been established to judge whether the spectrum is good or not [12]. First, the bands originating from water vapor must be subtracted accurately from the protein spectrum between 1800 and 1500 cm$^{-1}$. Second, a straight baseline must be obtained from 2000 to 1750 cm$^{-1}$. Many investigators have used a straight baseline between 2000 and 1750 cm$^{-1}$ as the standard for judging the success of water subtraction to obtain protein spectra [12,32,33]. By using two criteria, the average experimental errors of the spectra at the amide I and II band maxima could be less than 3% and 1.5%, respectively [8].

Empirically, in order to obtain a high quality protein spectrum, high protein concentration (>10 mg/ml in H$_2$O) [34] and small cell path-length (6–10 µm) are needed. The identical scan conditions are used to record buffer reference spectrum and the spectrum of protein solution. In addition, water vapor, as well as water (or buffer) itself, must be subtracted accurately from the protein spectrum. As mentioned above, using a straight baseline between 2000 and 1750 cm$^{-1}$ as the standard to judge the successfulness of water subtraction would lead to a higher quality of protein spectra.

**Application of FTIR to Protein Secondary Structure**

FTIR spectroscopy has been used to study the secondary structure composition, structural dynamics, conformational changes (effects of ligand binding, temperature, pH and pressure), structural stability and aggregation of proteins.

**Estimation of protein secondary structure**

Proteins are frequently referred to as having a certain fraction of structural components (α-helix, β-sheet, etc.). The secondary structural composition is some of the most important information for a structure-unknown protein. Therefore estimation of protein secondary structure is one of the major applications of the FTIR technique.

The quantitative estimation of protein secondary structure is based on the assumption that any protein can be considered as a linear sum of a few fundamental secondary structural elements, and the percentage of each element is only related to the spectral intensity (the molar absorptivity of C=O stretching vibration for each secondary structural element is essentially same). To analyze the amide I band

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component, FSD or second derivative spectra need to be curve fitted. The areas of FSD or second derivative spectra correspond to different types of secondary structure components. It is considered that the accuracy of measured band areas in the second derivative amide I spectra depends upon the correct positioning of the baseline [11]. For the globular protein, the correlation coefficient between IR and X-ray estimates of α-helices, β-sheets, β-turns and remainder was 0.98, 0.99, 0.90 and 0.92, respectively [8]. Dong and colleagues reported the distribution of secondary structures determined from the amide I spectra of globular proteins in aqueous solutions, which was nearly identical to the amount computed from crystallographic data [12].

Fig. 2 shows the analysis procedure of second derivative FTIR spectra of CRP. The percentage of α-helices was 41, calculated by the relative area at 1653 cm⁻¹. The percentage of β-sheets was 36, calculated by the relative area near 1637 cm⁻¹. The percentage of β-turns was 18, calculated by adding the areas of all β-turn bands between 1670 and 1690 cm⁻¹. The band area at 1648 cm⁻¹ was assigned to random coil. This data is consistent with the X-ray crystallography data. Table 3 lists some results calculated from the FTIR method compared with the X-ray crystallographic method, including the data of CRP and pyruvate kinase. From these data, the results from FTIR and X-ray crystallography methods match well.

**Kinetics of H-D exchange related to conformation**

Conformation and structural dynamics are essential to proteins for carrying out their proper biological function. The structural dynamics of a given conformation is likely to influence the activity of the protein. H-D exchange has been used extensively in studies of structural dynamics of...
proteins [36–38]. It has been established that the rates at which the amide proton exchange with solvent deuterium reflect the structural dynamics of proteins [39] and they are sensitive to the secondary structural composition and experimental conditions such as pH, temperature and pressure [36–39]. At constant experimental conditions, a more rapid rate of exchange implies a greater flexibility and motion in the structural region of the exchange.

FTIR spectroscopy has been used in connection with H-D exchange in polypeptides and proteins. The intensity changes of the amide I and II bands and the intensity change of the predominant secondary structural elements (α-helix and β-sheet) can be determined. It is suggested that it is more convenient to base H-D exchange investigations on apparent intensity changes of the amide II band [4], because it was not adversely interfered by the absorption bands of H2O, HOD, or D2O. Barksdale and Rosenburg suggested that the protein amide H-D exchange ratio could be presented as the fraction of unexchanged amide proton [40]. A calculation equation was established and used by some investigators [16,41]. Fig. 3 shows the H-D exchange ratio of pyruvate kinase in Tris buffer as a function of time, indicated by both original and second derivative FTIR spectra.

Most IR structural dynamics studies are focused on the H-D exchange of proteins on a global scale [16,36–38]. One may also focus on the exchange rate of secondary structure components. As a successful example, Dong et al. focused on the exchange rate of CRP secondary structural components, and obtained some useful information. CRP is a dimer with two functional domains in each subunit. There is a bias in the distribution of secondary structural elements between the two functional domains, namely extensive β-sheets in the cAMP binding domain and predominant α-helices in the DNA binding domain [42,43]. They took advantage of this feature and facilitated the interpretation of the ligand-induced conformational and structural dynamics changes [16].

Protein stability

Generally, protein stability studies can provide information on the folding/unfolding and structural stability of protein molecules. The advantage of FTIR amide I spectroscopy over other techniques is that the IR method can, in principle, monitor the folding/unfolding process of all parts of the protein simultaneously [44]. However, FTIR amide I spectroscopy has not been widely used as a method of choice in studies of protein folding. The reason is that obtaining good IR spectra of proteins in the presence of chemical denaturant, which must be used at high concentration to ensure protein unfolding, is not simple and has had only limited success [15]. First, IR cell with short path-lengths is required for preventing a saturation of IR detector by the absorption bands of urea or GdnHCl. Second, the major IR bands of denaturants (urea or GdnHCl) mask weaker amide I bands. Previous works on GdnHCl-induced denaturation of proteins have shown that accurate subtraction of GdnHCl absorption is difficult above 1660 cm\(^{-1}\) [46].\(^{13}\)C urea in D2O shifts the urea IR absorption to 1562 cm\(^{-1}\), leaving a clear window in the IR spectrum for observing the protein amide absorption.
The combination of IR cells with a short path-length and 13C-labeled urea allows us to measure the IR spectra of protein in the presence of high concentrations of chemical denaturant and gain useful information from the conformation-sensitive amide I band of protein [44,45].

The use of a wide range concentration of denaturant GdnHCl in FTIR spectroscopy has been reported. Bowler and colleagues successfully monitored GdnHCl-induced cytochrome c denature by FTIR spectroscopy [15]. The first point to be considered regarding the spectra of denatured proteins in the presence of GdnHCl is the reliability of the subtraction procedure. Protein-bound water molecules do not seem to have an altered vibrational spectrum from unbound water molecules [8,9]. The effects of over- and under-subtraction are best evaluated after resolution enhancement by second derivative methods [15]. It is apparent that the major effects of inaccurate subtraction are limited to the wavelength region between 1690 and 1665 cm$^{-1}$ [15]. One important criterion is subtraction of GdnHCl must be done so as to produce a smooth amide I band shape with no discontinuities and to maintain the amide I to amide II peak intensity ratio close to that observed for the native protein. Improper subtraction is evidenced by a doubled peak in the amide I region or an amide I peak with a concave discontinuity [15]. When the amount of amide I and GdnHCl intensity becomes comparable during the subtraction, the spectrum shows two peaks, allowing the GdnHCl intensity to be removed with a good degree of certainty. Although one cannot be certain of the absolute accuracy of the denaturant subtraction, by using consistent methods a high degree of precision in subtraction can be achieved allowing comparison of differences in the spectra of denatured states of closely related proteins, even in the region between 1690 and 1665 cm$^{-1}$ [15]. The instrument saturates near 4.0 M GdnHCl at the cell path-length used, which is another limitation that cannot be neglected. The highest GdnHCl concentration that is actually used should be less than 3.5 M by FTIR spectroscopy. Fig. 4 shows the subtraction of GdnHCl. In the figure, A represents the absorbance spectrum of CRP with 0.8 M GdnHCl, B represents the absorbance spectrum of 0.8 M GdnHCl, and C represents the absorbance spectrum of CRP after proper subtraction. Compared with Fig. 1(A), this IR absorbance spectrum is smooth and continuous with a normal intensity ratio of the amide I over the amide II band, so the subtraction of 0.8 M GdnHCl is acceptable.

Unlike chemical induced unfolding, thermal induced protein unfolding followed by FTIR spectroscopy can avoid limitations in saturation and subtraction. This technique has been widely used in protein aggregation studies [46,47]. Current theoretical and experimental evidence for protein aggregation suggests that aggregates are formed from partially folded intermediates [48]. Aggregation of proteins is a problem with serious medical implications and economic importance. To develop strategies for preventing protein aggregation, the mechanism and pathways by which protein aggregate must be characterized. In contrast to far-ultraviolet CD, IR spectroscopy is insensitive to light scattering, thus providing a valuable tool for protein aggregation studies. The thermally-induced aggregation processes of the majority of proteins studied by FTIR can be described with a two-state model, the predominant secondary structural element (α-helix or β-sheet) decreases as a function of temperature and is concomitantly replaced by intermolecular β-sheet aggregates [47]. As a successful example, Dong et al. adopted FTIR to investigate protein aggregation by a combination of thermal and chemical denaturation, thus providing a means to populate and characterize aggregation intermediates. They suggested that this method is valuable for studying the aggregation processes of a wide range of proteins. The identification and characterization of aggregation intermediates might lead to new interdiction strategies for amyloidogenic human disease, as well as improvements in industrial processing, storage, and delivery of therapeutic proteins.
Secondary Structures of Proteins Adsorbed onto Aluminum Hydroxide

Recently, a new IR technique was developed for studying protein structure at low concentration in solution [50]. Adsorbing proteins onto alhydrogel provides a means of obtaining FTIR spectra to study secondary structure and conformational changes of proteins in aqueous solution at very low concentration. This new procedure effectively lowers the concentration requirement for FTIR studies of proteins in aqueous solutions by at least 40-fold, as compared with the conventional FTIR method. This technique permits FTIR study of proteins to be carried out in the same concentration range as those used for CD and fluorescence, making it possible to compare structural information obtained by three commonly used techniques in the biophysical characterization of proteins.

Complexity of FTIR Spectrum: Side-chain Absorption

The FTIR studies of peptides and proteins have made efforts in identifying characteristic frequencies and determining their relations to the structures of protein molecules. The data analysis depended on empirical correlations of the spectra of chemically similar molecules, and occasionally yielded significant insights into the dependence of the spectrum on the conformation of the polypeptide chains [3]. It is very important to recognize that there is no simple correlation between IR spectra and the secondary structure elements of proteins. FTIR spectra are complicated in that every single spectrum has its own characteristics due to different micro-circumstances. Different ambient circumstances, including side-chain absorbance, make it difficult to precisely assign the secondary structure and frequency. The band assignment shown in Table 2 is only empirical. The estimation of secondary structure with the FTIR method should be considered only as a good approximation [8].

The main difficulty in a quantitative study of the protein spectra is the estimation of side-chain absorption, which must be taken into account in the analysis of protein spectra. The contribution of side-chain in the globular protein spectra is 10%–30% of the overall absorption [8]. This absorption is superimposed on the peptide absorption and can somewhat change the amide I band spectral parameters. Some amino acid residues, especially arginine, asparagine, glutamine, aspartic and glutamic acids, lysine, tyrosine, histidine and phenylalanine have intensive absorption in the amide spectral region [8,49]. Venyaminov and Kalnin [8] and Chirgadze et al. [49] have established the absorption parameters of these residues in the amide spectral region in H2O solution (Table 4). The quantitative estimation of amino acid side-chain groups permits more refined analysis of the secondary structure of polypeptides and proteins by FTIR, allows the protein spectra to be investigated in detail and achieves a better interpretation of the observed spectral effects. This should be kept in mind when analyzing protein spectra, especially when the content of these residues is high [8].

Summary

FTIR spectroscopy is a well-established experimental technique for studying the secondary structural composition and structural dynamics of proteins. Armed with mathematic resolution enhancement techniques,

<table>
<thead>
<tr>
<th>Table 4</th>
<th>Spectral parameters for absorbance bands of backbone and side chain in the 1700–1600 cm⁻¹ range in H₂O⁺</th>
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</thead>
<tbody>
<tr>
<td>Absorbance band assignment</td>
<td>ν₀ (cm⁻¹)</td>
</tr>
<tr>
<td>Asparagine (amide)</td>
<td>C=O (CONH₂) stretching</td>
</tr>
<tr>
<td></td>
<td>NH (CONH₂) bending</td>
</tr>
<tr>
<td>Glutamine (amide)</td>
<td>C=O (CONH₂) stretching</td>
</tr>
<tr>
<td></td>
<td>NH (CONH₂) bending</td>
</tr>
<tr>
<td>Arginine (guanidine)</td>
<td>C=N (CN₃H₅⁺) asymmetric stretching</td>
</tr>
<tr>
<td></td>
<td>C=N (CN₃H₅⁺) symmetric stretching</td>
</tr>
<tr>
<td>Lysine (amine)</td>
<td>NH⁺ (NH₃⁺) asymmetric bending</td>
</tr>
<tr>
<td>Tyrosine (phenyl)</td>
<td>Ring vibration</td>
</tr>
<tr>
<td>N-terminal group (amine)</td>
<td>NH⁺ (NH₃⁺) asymmetric bending</td>
</tr>
</tbody>
</table>

† ν₀, band position; ε₀, absorptivity of mole residues. Data are from Venyaminov and Kalnin [9].

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various methods of FTIR data analysis have been well developed. The correlations between IR spectra and protein secondary structures have been established. The amide I band component assignments to protein secondary structure elements, such as α-helix, β-sheet, β-turn and random structures, are available for proteins in H₂O as well as D₂O media. One could adopt those techniques to study various protein systems for different purposes. Sample backgrounds have to be subtracted accurately from the protein spectrum with strict procedure(s). To obtain high-quality IR spectra, relatively high concentrations of proteins (e.g. >10 mg/ml) and small cell path-length (6–10 µm) are needed for proteins in H₂O solution unless protein was adsorbed onto alhydrogel.

The singular advantage of FTIR over other techniques is convenience. IR spectra can be obtained for proteins in a wide range of environments with a small amount of sample. Other than estimating the content of protein secondary structures, IR can also provide information on protein structural stability and dynamics. The FTIR spectrum is also complex, and some characteristic bands of secondary structure elements might overlap. The background subtraction procedure could also bring experimental error. Estimation of side-chain absorption must be taken into account in the analysis of protein spectra. All the complications indicate that caution has to be exercised in the interpretation of IR spectra of proteins. When estimating the percentage contents of protein secondary structures, a combination of FTIR and CD is recommended for increasing prediction accuracy.

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