Fourier transform infrared conformational investigation of type I collagen aged by in vitro induced dehydration and non-enzymatic glycation treatments

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Abstract

The two main ageing-inducing events in the collagenous tissues are the water loss and the formation of intermolecular cross-links based on the reaction of collagen with matrix carbohydrates, following a mechanism known as non-enzymatic-glycation. With the aim to mimic the two deleterious processes for the protein structure, rat-tail collagen was submitted to hydration changes and allowed to interact with two sugars characterized by different reducing properties, D-glucose and D-ribose. Fourier transform infrared (FTIR) spectroscopy was employed to investigate the conformational changes induced in the protein by the two treatments by analyzing the subsequent spectra modifications. FTIR spectra monitored: i) the amplitude and position changes of the two characteristic absorption bands OH stretching and Amide I, in dependence on the humidity level; a significant hysteresis effect in the ν(OH) band (ν~3400 cm⁻¹) amplitude of the protein dehydrated and then rehydrated to the initial relative humidity (αw=0.92-0.06) may be related to the enhancement of the β-sheet fraction in the protein structure as revealed by the parallel modification in the Amide I band (ν~1650 cm⁻¹); ii) the area of the carbohydrate double band peaking at 1080 cm⁻¹ and 1031 cm⁻¹, associated to the accumulation of the glycation products, depending on the sugar concentration and incubation time. The association of both sugars to collagen only minimally affects the protein secondary structure as revealed by Amide I band Gaussian analysis. The whole set of results suggests hints to hypothesize a self-assembly model for collagen molecules induced by ageing.

Introduction

Collagen is the most abundant protein in mammals and accounts for 25-30% of their protein content. It is the main fibrous component of skin, bone, tendon and cartilage, accounting in particular for three-quarters of the dry weight of skin, and representing the most prevalent component of the extracellular matrix. Collagen molecule is a rigid rod like structure able to resist stretching. It is composed of three polypeptide left-handed alpha chains coiled around each other to form a typical right handed rope like triple helical rod (approximately 1.5 nm in diameter, 300 nm in length). The triple-helical sequences are comprised of Gly-X-Y repeats, X and Y being frequently proline and hydroxyproline, respectively. Stabilization of the triple helix is assured by a lot of structural conditions as the presence of glycine as every third residue, a high content of imino acids with a rigid cyclical structure (proline and hydroxyproline), and electrostatic interactions involving lysine and aspartate. However, a key role is undoubtedly played by water which provides inter-chain hydrogen bonds and water bridges, and coats triple helix by a cylinder-like of hydration. Water critically regulates chain flexibility and assures the water mediated H-bonds favoring fiber recognition and alignment. Water depletion, caused by tissue maturation, exposition to UV light and/or some pathological processes, like diabetes, triggers non-enzymatic glycation reactions between the protein and the complex carbohydrates present in the matrix, leading to the formation of oxidation products, known as Advanced Glycation End-products (AGE). AGEs have the effect to modify the collagen fibers physical properties, inducing an increase in stiffness and fiber diameter resulting in a loss of solubility and flexibility and brittleness of the collagenous tissues.

Several experimental techniques have been used in order to monitor and describe the changes in collagen structural properties as a consequence of ageing processes: high resolution X-ray diffraction, FTIR spectroscopy, nuclear magnetic resonance, circular dichroism, transmission electron microscopy, atomic force microscopy, and Raman spectroscopy.
fraction, Scanning Electron Microscopy (SEM), Fluorimetry and Raman spectroscopy.\textsuperscript{8-11} Among the spectroscopic techniques, Fourier Transform InfraRed Spectroscopy (FTIR) is recognized as one of the most sensitive for studying biological macromolecules, being able to monitor changes in the structural features in dependence on the physicochemical and environmental conditions.\textsuperscript{12} In the present work, with the aim to investigate the role of both the reactions on the conformational properties of collagen, and how they influence one each other, the protein was submitted to hydration changes and allowed to interact with two sugars characterized by different reducing properties, D-glucose and D-ribose.\textsuperscript{13} Fourier Transform InfraRed (FTIR) spectroscopy was employed to investigate the conformational changes induced in the protein by the two treatments by analyzing the subsequent spectra modifications.

The typical IR spectrum of a protein is dominated by the Amide bands, due to the vibrations of the peptide bond. Amide I band, ranging between 1720 and 1610 cm\textsuperscript{-1}, mainly due to contributions from the C=O stretching vibration of the peptide backbone, is particularly sensitive to the secondary structure composition of the protein. To predict the protein conformation, Amide I band was decomposed into individual components, corresponding to the different types of secondary structure.\textsuperscript{14} Water structured around proteins displays strong absorption bands in the IR frequency range.\textsuperscript{15-17} A broad absorption band in the range 3700-3000 cm\textsuperscript{-1} due to the hydroxyl stretching vibrations (ν(OH)), and a relatively sharp but weaker one, caused by the bending mode δ(OH), peaking around 1640 cm\textsuperscript{-1}. The ν(OH) profile and amplitude modification, in particular, as a consequence of the hydration fluctuations may be related to the reshaping of the water layer surrounding the bio-macromolecule and was correlated to the protein conformational changes.

Glycation compound formation was assessed by means of the identification in the IR spectrum of the protein/carbohydrate complex of an absorption band characteristic of the interaction between collagen and reducing sugars, such as Glucose and Ribose.\textsuperscript{11,17} Ribose and Glucose IR spectra are characterized by a lot of absorption bands largely superimposed to the main protein features, however the spectra of both the sugars are characterized by a huge absorption band in the 1150-950 cm\textsuperscript{-1} due to the glucid C-O-C, C-O, C-O and C-C vibrations. As a consequence, glycated collagen displays significant spectral differences with respect to the native protein, in correspondence of the wave number range spanning such region (1150-950 cm\textsuperscript{-1}). This band was assumed as a marker of the protein/carbohydrate interaction, as discussed in literature,\textsuperscript{11} and the modifications in the amplitude and area band were correlated to the glycation amount produced.

In the natural ageing processes dehydration and glycation may be considered as subsequent steps, where carbohydrates act as water substitutes, being rich in OH moieties, against the damages produced by moisture subtraction, but the structure rearrangements produced in collagen by the two reactions are not yet clarified.\textsuperscript{18,19} The purpose of the present preliminary study is to separately investigate the modifications produced in collagen structure by the two processes, dehydration and non-enzymatic glycation, in order to contribute to the elucidation and the spectroscopic characterization of both the ageing processes.

Materials and Methods

Samples
Collagen from rat tail (MW 102610) was purchased from Sigma-Aldrich and used without further chemical purification. It was solubilized in acid acetate aqueous solution (0.5M). Glucose and Ribose were purchased from Sigma-Aldrich.

The samples for FTIR measurements were thin films obtained by depositing a drop of protein solution on CaF\textsubscript{2} platelets and allowing them to dry in air. The sample was vertically positioned in such a way as to allow for transmission measurements.

Hydration-dehydration procedure
Different hydration degrees of the samples were obtained by equilibrating them for two days directly in the inner ambient of a home-made dry box over suitable hydrating saturated salt solutions put in a vessel placed into the box and assembled together with the sample. In this way the relative humidity was made to vary in the activity range \(a_w=0.06-0.92\), being \(a_w=0.06\) provided by NaOH, and \(a_w=0.92\) by KNO\textsubscript{3} saturated salt solutions.\textsuperscript{15}

Glycation reaction
It was accomplished by incubating collagen under sterile conditions in Glucose and Ribose 5 and 50 mM in phosphate buffer solutions (0.15 mM, pH 7.4) at 37°C. The reaction development was checked after 3, 10 and 15 days by taking aliquots of solution and smearing them on the CaF\textsubscript{2} platelets. The air dried films were equilibrated in the dry box at \(a_w=0.92\) as described above.

Fourier transform infrared spectroscopy
Infrared spectra were collected with a Jasco 420 FTIR spectrophotometer operating at RT within the 4000-600 cm\textsuperscript{-1} range (128 scans, spectral resolution 2 cm\textsuperscript{-1}). Gaussian functions were used to fit Amide I and ν(OH) bands. Amide I band (1720 – 1610 cm\textsuperscript{-1}) was decomposed in sub-bands whose position and area were related respectively to the different types and to the amount of secondary structures.\textsuperscript{14} The sum of the areas of the peaks represents the total amount of secondary structure in the protein.

OH stretching band (wave number range 4000-3000 cm\textsuperscript{-1}) was analyzed after subtraction of the spectrum of the same sample pre-heated to 80°C for 4 h assumed as dry sample. In this way any contribution to the band of Amide \(\Lambda (N-H\) stretching, ν~3300 cm\textsuperscript{-1}) and aliphatic CH\textsubscript{2}/CH\textsubscript{3} stretching bands (2960-2850 cm\textsuperscript{-1}), occurring in the same wave number range investigated, was systematically eliminated. The cleaned OH stretching band was decomposed into components whose frequencies were related to different O-H bond lengths, i.e. to water molecule sets different in the vibrational energies as a consequence of the various interaction strengths, mutual and over the binding sites offered by the protein surface.

The area of the glycation band, the twin peak ranging between 1150-950 cm\textsuperscript{-1}, was evaluated after baseline correction.

Results and Discussion
In Figure 1, FTIR spectra of pure collagen (a) and collagen glycated with ribose (50 mM, 15 days) (b) films, prepared at the highest attained water activity \((a_w=0.92)\) and recorded in the wave number range 4000 – 400 cm\textsuperscript{-1}, are displayed and compared. In the past 25 years infrared spectroscopy has produced a significant upgrade in structural studies of collagen:\textsuperscript{20-25} Although the IR spectra of the protein has already been shown and discussed in several papers in the past,\textsuperscript{21} the characteristic features are summa-
rized here, for the sake of clarity. In the region between 3700 and 2600 cm⁻¹ ν(OH), ν(NH) and ν(CH) absorption bands characteristic for bound water (~3400 cm⁻¹), Amide A (3300 cm⁻¹) and CH₃, CH₂ groups of alkyl/acyl chains (2950-2860 cm⁻¹). The skeletal functionality region, below 2000 cm⁻¹, is dominated by the three main Amide bands, Amide I (1650 cm⁻¹), Amide II (1550 cm⁻¹) and Amide III (1340-1220 cm⁻¹). The main IR-active groups are indicated in Figure 1 by arrows. The two spectra, a and b, are similar in the main features except for the low wave number range, between 1150-950 cm⁻¹ where the typical twin band due to the carbohydrate moiety vibration modes shows, in the b spectrum, a very larger amplitude with respect to the native collagen one (a spectrum).

**Hydration-dehydration treatments**

In order to study the role of water interacting with collagen, the FTIR spectrum was analyzed in the OH stretching region by changing the hydration content of the sample following the method described in the Experimental Section. The composition in Gaussian sub-bands of the ν(OH) feature can be correlated to the hydrogen bond network around the protein each component being assigned to different sets of H₂O molecules each one related by O-H bonds of comparable lengths. Figure 2A displays the absorption spectrum of the water bound to collagen prepared at aₘ=0.92 (as described in Materials and Methods). The band was resolved, following the procedure of the second derivative analysis, into four Gaussian components (R²=0.998) peaking at 3610, 3513, 3360 and 3180 cm⁻¹. They can be related to four classes of water molecules bound to the protein, different in vibrational energies and each one characterized by a single average H-bond distance (H•••O-H) of the protein helix: 0.303, 0.293, 0.283, 0.276 nm, respectively.

The component band at the highest wave number region is corresponding to H-bond distances characteristic of vapor-like state. Similar features, detected in different biological macromolecules were assigned to the non-H-bonded or weakly H-bonded O-H groups. It is reasonable to suppose that in our sample, prepared at very low hydration levels, they could be originated by the most external water molecules sitting on the outer hydration layer coating the macromolecule. The inability for these molecules to establish active H-bonds with the surroundings, as in bulk water, accounts for the high mobility and the large vibrational energy comparable with those of the water molecules in the vapor state. They represent about the 5% of the total amount of water hydrating collagen at the relative humidity settled for the experiment, being the percentages calculated as the ratio A/Aₜ of the single component area (A) with respect to the total band area (Aₜ). The two component bands peaking at the intermediate wave numbers may be related to H₂O molecules coordinated by two or three more or less distorted or strained H bonds. They may be identified with H₂O molecules deeply located inside the protein helix, acting as water bridges within a single peptide chain and/or interconnecting the different α-chains in the triple helix. They constitute the 19% and the 31%, respectively, of the total water amount. The corresponding H-bond distances suggest they could represent the solvation molecules involving C=O groups belonging to Glycine and Hydroxyproline moieties. The component at 3180 cm⁻¹ may be attributed to water molecules located near to the protein surface and hydrogen bonded to polar and charged groups exposed to the macromolecule surface: they originate some ice-like tetrahedral structures more or less distorted in dependence on the orientation of the surface chemical groups. The corresponding average H bond length is in fact characteristic for arrangement of H₂O molecules as in small solid water clusters, assuming the length O–O of hydrogen bonds in ice as 0.276 nm. The broad profile of such a band testifies for the large distribution of water molecules in a wide con-

![Figure 1. Fourier transform infrared spectra of (a) native collagen from rat tail tendon (SIGMA) and (b) collagen glycated for 15 days with glucose (50 mM). Measurements were carried out in transmission geometry on dried films smeared on CaF₂ slabs. To facilitate the comparison, the spectra were normalized with respect to the amount of protein in each sample.](image1)

**Figure 2. Gaussian deconvolution of ν(OH) stretching band for native collagen hydrated at aₘ=0.92, after subtraction of the dry sample spectrum (A).** ν(OH) stretching band for collagen hydrated at aₘ=0.92 (line a) and after dehydration at aₘ=0.06 and subsequent rehydration at aₘ=0.92 (curve b). Curve c represents the difference spectrum (B). Amide I band deconvolution for native collagen at aₘ=0.92 (C). Amide I band deconvolution for native collagen dehydrated at aₘ=0.06 and thereafter rehydrated at aₘ=0.92 (D). The sum of the fitted curves is shown by the thin grey lines, closely overlapping the experimental data trace, which is shown as a full heavy line. Blue lines in panels C and D indicate the component bands mainly modified by the dehydration-rehydration treatment.
figuration variety, concurring to the feature. Such a sub-band represents the main hydration fraction at the settled humidity, corresponding to the 45% of the total amount of hydration water.

The dehydration treatment at $a_w=0.06$ followed by a subsequent rehydration up to the original relative humidity ($a_w=0.92$) causes a pronounced hysteresis effect in the $\nu$(OH) band amplitude recover and a remarkable modification of the profile (Figure 2B, curve b). In Figure 2B, curve c deserves particular note: it was obtained as the difference between curve a, corresponding to the hydrated sample, and curve b, representing the rehydrated one, after dehydration. It can be considered as the spectrum of water that can no longer be drawn back into the sample after the dehydration treatment. The broad band is peaking at $\nu \sim 3375$ cm$^{-1}$, therefore it is roughly filling the position of the third component of the original hydration band (a). Moreover it exhibits a shoulder on the lower wave number side ($\nu \sim 3100$ cm$^{-1}$). This finding reveals that dehydration treatment gives rise in collagen to the loss of a considerable amount of H$_2$O molecules involved in the internal water bridges and of a fraction of the molecules coating the polypeptide surface. These water molecules once desorbed, only partially can be re-adsorbed. It accounts for the 21% of the total water amount initially hydrating the sample.

Water deprivation has significant consequences on the macromolecule structure. They can be reflected by the changes in the low wave number region of the IR spectrum. The secondary structure analysis was carried out by analyzing the Amide I band in Gaussian components in the native sample and in the rehydrated one after dehydration. Qualitative and quantitative information about the conformational composition of the protein prepared in the different hydration states were obtained. The comparison of the spectra C and D in Figure 2 shows that the Amide I band amplitude of the rehydrated sample is reduced with respect to the freshly prepared one, as expected, due to the lacking of the water binding contribution in such wave number region ($\delta$(OH) ~ $1640$ cm$^{-1}$). The curve fitting procedure allowed determining the secondary structure composition in the freshly prepared sample ($a_w=0.92$, Figure 2C), as shown in Table 1: $\beta$-turn (1695 cm$^{-1}$), $\beta$-sheet (1680 cm$^{-1}$), $\alpha$-like helix (1660 cm$^{-1}$), unordered structure (1644 cm$^{-1}$), triple helix (1628 cm$^{-1}$), $\beta$-sheet (1615 cm$^{-1}$), side chains (1607 cm$^{-1}$). The multipeak decomposition of the band measured in the rehydrated sample (Figure 2D), revealed the modification occurred in the secondary structure. By normalizing the two peaks with respect to the maximum amplitude it was possible to compare the band shape. The procedure enabled the detection in the rehydrated sample of a band enlargement in the high wave number side. The most significant change in the conformational composition was in fact concerning the growth of a broad sub-band peaking at $\nu \sim 1670$ cm$^{-1}$ largely spanning the high wave number region of the Amide I band and the parallel decrease of the component bands at 1695 and 1660 cm$^{-1}$. Such changes can be explained by assuming the formation of chemical bonds among collagen molecules laterally associating by forming structures side by side interacting. The formation of such structures may be envisaged as responsible for a relative increase of $\beta$ components with respect to the $\alpha$-helix contribution in the Amide I envelope. Therefore, as a consequence of the removal of water, the association of the collagen molecules takes place and once water is excluded, the assembled fibers are no able to re-adsorb all the water restituted during the rehydration phase, due to the lacking of available H-bonding sites. This arrangement is the main responsible for the decrease in the percentage amount of hydration water in the rehydrated sample and of the hysteresis effect measured in the OH stretching band.

**Non-enzymatic glycation reaction**

The Infrared marker for sugar and protein-sugar compounds is represented by a characteristic twin band in the fingerprint region, between 1150-950 cm$^{-1}$, as shown by the b spectrum in Figure 1. The band is peaking at 1032 cm$^{-1}$ and at 1080 cm$^{-1}$, the two peaks corresponding to C-O, C-C stretching vibrations and to C-O-H and C-C-O bending vibrations, respectively and can be used for the discrimination between native and glycated collagen by means of the amplitude increase displayed as a consequence of glycation reaction.\(^{18}\) No frequency shift was monitored but a change in amplitude as a function of the sugar concentration and the incubation time. This change was employed for kinetic monitoring of glycation process and for roughly estimating the glycation reaction development in dependence on the amount and type of sugars. The area under the Amide I band was assumed as indicative of collagen content and it was used for normalization of sugar peak areas used for measure the glycation product accumulation. Figure 3A shows the plot of normalized absorbance vs. sugar concentration as measured after 3, 10 and 15 days from the start of the reaction. The amount of glycation products increases with increasing sugar concentration, although no linear dependence was detected, as previously shown in literature.\(^{26}\) The extent of glycated collagen formation is markedly enhanced with the initial addition of sugar, indicating the high propensity to the interaction of protein with carbohydrates. Formation of complex is strongly sugar type dependent, the ribose being more potent in the reaction yield, in accordance with the difference in the reductive properties of the two sugars.

Figure 3B displays the temporal kinetic of the glycation reactions promoted by the two sugars at different concentrations (5 and 50 mM). The accumulation of the reaction products is linearly dependent on the time course of the reaction, monitored for 15 days, and the formation rate is higher for the reaction of collagen with ribose than with glucose.

Complementary investigations of the conformational modifications induced in the protein by the interaction with sugars was carried out once more by analyzing Amide I band in the spectra of collagen after incubation (15 days) with ribose and glucose (50 mM). The results of the analysis are displayed in Table 1 where the percentage amount of each secondary structure motif detected in the native collagen is compared with the values obtained for collagen glycated with the two carbohydrates.

No significant difference in the conformational motifs were detected except for a light increase in the random coil amount and in the $\beta$-turn content respectively in collagen-glucose and in collagen-ribose products. The result demonstrates that the overall conformation of the collagen is not altered by sugar attachment and the recorded modifications, involving only negligible portions of the polypeptide may be interpreted as due more to local rearrangements of the structure, perhaps as a consequence of the incubation procedure, than to structural modifications induced by the glycation reaction.

**Conclusions**

In conclusion, FTIR spectroscopy is confirmed to be a technique particularly valuable for analyzing the conformational conversion of collagen as a consequence of dehydration and non-enzymatic glycation reaction with reducing sugars, the two main ageing inducing agents in the collagenous biological tissues such as skin, tendon and cartilage. The study was performed by correlating the
Table 1. Secondary structure content in native collagen, collagen glycated with glucose, and collagen glycated with ribose, as evaluated from the Amide I band deconvolution.

<table>
<thead>
<tr>
<th>Assignments</th>
<th>Band position (cm⁻¹)</th>
<th>Collagen (%)</th>
<th>Collagen+glucose* (%)</th>
<th>Coll+ribose° (%)</th>
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<td>1</td>
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<tr>
<td>β sheet</td>
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<td>4</td>
<td>3</td>
<td>5</td>
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<tr>
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<td>17</td>
<td>16</td>
<td>16</td>
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<tr>
<td>Unord</td>
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<td>27</td>
<td>16</td>
</tr>
<tr>
<td>Helix/coil</td>
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<td>38</td>
<td>30</td>
<td>28</td>
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</tr>
<tr>
<td>β turn</td>
<td>1695</td>
<td>8</td>
<td>5</td>
<td>12</td>
</tr>
</tbody>
</table>

*Collagen glycated with glucose (50 mM) for 15 days; °collagen glycated with ribose (50 mM) for 15 days.

modifications in the Amide I band, reflecting the changes occurred in the secondary structure of the protein in dependence on the treatments to which it was submitted, with the two spectral features specifically monitoring the chemical groups directly involved in the reactions: ν(OH) band and the twin peak at 1080 and 1031 cm⁻¹.

The main conclusions can be summarized as follows.

First, the results clearly stress the key role played by the hydration level on the FTIR spectrum and the extent by which water removal modifies the assembly properties of collagen. Molecular stability of the collagen structure is assured by four water classes different in the vibrational properties: two of these are related to inner helix H₂O molecules, involving Gly and Hyp residues, and two to external water stratified in poly-molecular layers.

Second, dehydration–rehydration cycle induces lateral association of the fibers and shrinkage with a consequent squeezing out of a considerable fraction of external water unable to be re-adsorbed when the initial hydration level is restored. The effect is accompanied by an increase of β sheet amount. On these bases a simple model can be hypothesized describing one of the phases of the most dramatic changes in collagenous tissues with age. It consists in the alignment and assembly of the collagen molecules towards fibers polymerization through intermolecular cross-linking water excluding.

Third, the interaction collagen-sugars can be monitored by analyzing the absorption band ranging between 1150-950 cm⁻¹. Glycation reaction is dependent both on the concentration and the reducing properties of the sugar. In particular, the linear dependence of the glycation products amount on the incubation time may help to backtrack the early stage of the reaction in vivo and opens new research perspectives on the ageing of biological tissues providing improvements in many fields, for example in the dating of archaeological samples.²⁷

Fourth, conformational analysis performed through Amide I deconvolution shows that collagen-sugar blends only minimally affect collagen secondary structure, confirming the protective role of sugars on protein conformational changes induced by deleterious effects such as tissue water loss.²⁸

References