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Investigating the relationship between changes of collagen fiber orientation during skin aging and collagen/water interactions by polarized-FTIR microimaging

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Abstract. Upon chronological aging, human skin undergoes structural and molecular modifications, especially at the level of type I collagen. This macromolecule is one of the main dermal structural proteins and presents several age-related alterations. It exhibits a triple helical structure and assembles itself to form fibrils and fibers. In addition, water plays an important role in stabilizing collagen triple helix by forming hydrogen-bonds between collagen residues. However, the influence of water on changes of dermal collagen fiber orientation with age is not been yet understood. Polarized-Fourier Transform Infrared (P-FTIR) imaging is an interesting biophotonic approach to determine in situ the orientation of type I collagen fibers, as we have recently shown by comparing skin samples of different ages. In this work, P-FTIR spectral imaging was performed on skin samples from two age groups (35- and 38-year-old on one hand, 60- and 66-year-old on the other hand); and our analyses were focused on the effect of H\textsubscript{2}O/D\textsubscript{2}O substitution. Spectral data were processed with fuzzy C-means (FCM) clustering in order to distinguish different orientations of collagen fibers. We demonstrated that the orientation was altered with aging, and that D\textsubscript{2}O treatment, affecting primarily highly bound water molecules, is more marked for youngest skin samples. Collagen-bound water-related spectral markers were also highlighted. Our results suggest a weakening of water/collagen interactions with age. This non-destructive and label-free methodology allows to understand better the importance of bound water in collagen fiber orientation alterations occurring with skin aging. Obtaining such structural information could find benefits in dermatology as well as in cosmetics.

Keywords: P-FTIR imaging, skin aging, collagen, water, hydrogen-deuterium exchange
Introduction

Skin is the largest organ of the human body and has many important functions such as protection against external aggressions and thermoregulation. This cutaneous structure is composed of three layers: the epidermis, the dermis and the hypodermis. The dermis is made up of two sub-layers: the superficial or papillary dermis, and the deep or reticular dermis. This dense conjunctive tissue is constituted of fibrous proteins like collagen and elastic fibers, and of non-fibrous components like proteoglycans (PGs) and cells. Type I collagen represents about 80% of skin dry mass and confers to the skin its strength and its resilience. Type I collagen fibers are responsible of tensile properties of the skin; they are thin and poorly organized in the papillary dermis while thicker and well organized in the reticular dermis\textsuperscript{1}.

During aging, skin undergoes several structural and molecular changes such as the appearance of wrinkles and the loss of elasticity. That is why it is the subject of much attention in dermatology as well as in cosmetology. Some studies reported an accelerated degradation and a decreased synthesis of type I collagen with age and that this loss of collagen leads to a dermal atrophy\textsuperscript{2,3,4}. Furthermore, it has been demonstrated that the elastic fibers network was altered with aging\textsuperscript{5,6}. A loss of hydration due to changes in PGs content and localization was observed upon aging\textsuperscript{7}.

Playing an important role in stabilizing collagen triple helix, water molecules form hydrogen-bonds between collagen residues\textsuperscript{8,9}. Xia et al. hypothesized that interactions between water molecules and PGs could influence the orientation of collagen fibers in the articular cartilage\textsuperscript{10}. Besides, several recent studies using magnetic resonance imaging demonstrated that the articular cartilage presents a well-distinct anisotropy closely connected to collagen ultrastructure and, also that the interaction between water molecules and PGs could modulate the orientation of collagen fibers via the influence of bound water modifications\textsuperscript{11,12}. Indeed, water bound to collagen behaves differently than the bulk water.
and is involved in the formation of hydrogen bonds and water bridges between collagen triple helix residues\textsuperscript{13-15}. Thus, it has been suggested that water molecules could have a crucial role in the stabilization of the conformation and the packing of collagen fibers in the articular cartilage\textsuperscript{16,17}.

Biophotonic techniques are potential tools to assess the orientation of collagen fibers. Polarized light microscopy (PLM) is one of the first developed techniques allowing morphology studies on collagen-rich tissues\textsuperscript{18-20}. Some studies on skin have employed polarization sensitive-optical coherent tomography (PS-OCT) to reveal the molecular orientation of collagen fibers exhibiting some anisotropic properties such as birefringence\textsuperscript{21-24}. Second harmonic generation (SHG) technique is another emerging technique to investigate collagenous tissues by exploiting the specificity of collagen SHG signal. In addition, polarization-resolved imaging measurements permitted to determine the spatial distribution of the orientation of collagen fibers\textsuperscript{25-27}. Fourier Transform infrared (FTIR) spectroscopy is a vibrational technique which allows obtaining some structural information on oriented molecules like type I collagen when this biophotonic approach is combined to polarization measurements. Compared to previous techniques, FTIR has the advantage to probe the entire molecular composition of a complex sample such as cutaneous tissue. Indeed, some studies have shown the efficiency of this technique to compare collagen fiber orientation in normal and pathological articular tissue\textsuperscript{28-33}. Recently, our group highlighted some changes in the orientation of dermal collagen with age by using polarized-FTIR (P-FTIR) imaging\textsuperscript{34}. However, to the best of our knowledge, no studies have yet investigated the importance of water in collagen fiber orientation with skin aging.

In this work, skin samples from thirty- and sixty-years-old women were treated with deuterium oxide (D\textsubscript{2}O) to determine the influence of the bound water molecules in changes of collagen fiber orientation. Untreated and treated skin cryosections were analyzed by P-FTIR
imaging. Spectral data were processed by fuzzy C-means (FCM) clustering to define the different layers of the skin and the different types of collagen fiber orientation within the dermis. Moreover, discriminant vibrations related to collagen/water interactions were highlighted for the two different-age skin samples by using discriminant wavenumber selection algorithm.

Materials & Methods

Sample preparation

Frozen abdominal skin samples were obtained from female Caucasian subjects in their thirties (35- and 38-year-old) and in their sixties (60- and 66-year-old), through BIOPREDIC International (Rennes, France) and were treated as described by Zhang et al. for H$_2$O/D$_2$O substitution$^9$. Briefly, skin samples were immersed into 1 mL of distilled water for 24h and then into 1 mL of deuterium oxide (D$_2$O, 99.9% isotopic purity, Sigma-Aldrich, France) for 24h. The first period in water aims at saturating all the interaction sites of the skin samples. Skin samples from the same age groups were also immersed into distilled water for 48h as controls. Control and deuterated samples were embedded in Tissue-Tek O.C.T. compound (Sakura Finetek, France) and were immediately snap frozen in liquid nitrogen. Five thin serial tissue cryosections (5 µm) were deposited on calcium fluoride (CaF$_2$) windows. Hematoxylin and eosin (H&E) staining was used as standard staining on adjacent skin sections (5 µm) to those dedicated to FTIR analysis. Three 5 µm-thick sections of rat-tail tendons (Sprague-Dawley rats aged 2 months) were prepared according to the same protocol than skin samples.

Polarized-FTIR imaging
Spectral data were collected by using a Vertex 70 FTIR spectrometer coupled to a HYPERION 3000 FTIR microscope (Bruker Optics, Ettlingen, Germany) controlled by the Opus 7.2 software and equipped with a 64x64 pixel Focal Plane Array (FPA) detector. FTIR spectral acquisitions were realized in transmission mode with a x15 Cassegrain objective and matching condenser lens. Cryosections of rat’s tail tendon were aligned along the direction of x-axis of the motorized plate and spectral maps of 175x175 µm² were acquired in non-polarized mode and with IR radiation polarized perpendicular to the tendon orientation (i.e., x-axis direction). In such material, the collagen fibers are arranged and aligned longitudinally to the tendon length. For each skin cryosections, a final measuring area of 175x350 µm² mapped from the stratum corneum to the reticular dermis with a resolution of 2.7 µm²/pixel. It took about 3 minutes to record 8192 spectra. Polarization measurements were performed by inserting a polarizer in the incident IR beam. FPA imaging maps (n=5) were acquired for each tissue section whose the skin surface was oriented parallel to the x-axis of the motorized plate. To determine the spatial distribution of the collagen orientation, the polarization was chosen perpendicular to the skin surface. Each spectrum was recorded with 64 scans at a resolution of 4 cm⁻¹, on the 900-3800 cm⁻¹ spectral range. A background spectrum was recorded with 128 scans of accumulation on a blank part of the same CaF₂ window prior to spectral acquisitions. Each spectrum was rationed to the background spectrum in order to reduce any atmospheric effects and to obtain chemical features from the analyzed sample.

Data analysis

Spectral data were preprocessed by an automated method based on Extended Multiplicative Signal Correction (EMSC) algorithm²⁵ implemented in the MATLAB 8.3 software (The Mathworks, USA). Briefly, EMSC allows to eliminate low signal/noise ratio spectra and to realize a baseline correction and normalization in order to compare spectra each other²⁶,²⁷.
Subsequently, FCM clustering was used as statistical multivariate processing. FCM allows to assign each spectrum to every clusters with associated membership values comprised between 0 (no class membership) and 1 (highest degree of cluster membership)\textsuperscript{38}. Consequently, a spectrum could belong to several clusters contrary to “hard” clustering techniques such as K-Means\textsuperscript{39}. FCM images were constructed with redundancy-based algorithm (RBA) that presents the advantage to determine automatically the number of clusters\textsuperscript{38-40}. FCM clustering were performed using home-made algorithms written in the MATLAB 8.3 software.

The randfeatures function was used to identify the most discriminant wavenumbers between 2 groups of spectral data\textsuperscript{42-44}. This algorithm, based on a random selection of a pool of 15 wavenumbers, reduces the spectra dimension (751 wavenumbers in our case) to the 15 wavenumbers previously selected. Then, each spectrum is mean-centered by the subtraction of the mean spectrum of its group. A linear and supervised classification algorithm is then applied and returns, for each spectrum, a membership value towards each spectral group. A spectrum will be considered as well classified if its biggest membership value is attributed to the good category and if this value is over a Confidence Threshold (fixed in our case to 95%). The last step deals with the number of spectra well classified, according to the previous conditions. If this classification is over a Performance Threshold (fixed here to 95%), the subset of 15 wavenumbers randomly selected is considered as efficient for the distinction between the two spectral groups, and thus a corresponding score is incremented by one for each wavenumber. This procedure is then repeated with another random selection of 15 wavenumbers. It must be noted that a wavenumber can be chosen several times. The randfeatures function stops when 1000 subsets are retained. Finally, the scores of every wavenumbers can be analyzed by the user. Higher is a score, the more discriminant is the
corresponding wavenumber. This discriminant wavenumber selection algorithm was written in the MATLAB 8.3 software.

Results

P-FTIR spectral data analysis on rat-tail tendons

As previously realized in other studies\textsuperscript{18, 34}, rat-tail tendon was used as a reference sample to measure the effect of polarization on type I collagen IR signal. Reconstructed images and mean spectra revealed a change in the Amide I/Amide II ratio as a function of the polarization mode used. The Amide I/Amide II ratio presented a value of 1.4 without polarization, and between 1.6 and 2 when the polarized IR radiation was perpendicular to the collagen fiber orientation probing mainly stretching vibration of C=O bands as shown in Fig. S1 (Supporting Information). Indeed, the C=O stretching, the major vibration of the Amide I band, displays a transition moment preferentially oriented perpendicular to the collagen fiber long axis. Also, the C-N stretching and the N-H bending, which contribute to the Amide II band, possess a transition moment preferentially oriented parallel to the collagen fiber long axis. Thus, the Amide I/Amide II ratio is a potential indicator of the orientation of collagen fibers, as probed by P-FTIR, and was used for next analyses on skin samples.

P-FTIR spectral data analysis on human skin samples

Fig. 1(a) and 1(b) display H&E stained histological sections of the 35-year-old human skin sample, untreated and treated with D\textsubscript{2}O respectively. These morphological images permitted to visualize the main skin structures which are the epidermis and the dermis. Adjacent sections were analyzed by FTIR imaging in a label-free manner. The presence of the 1338 cm\textsuperscript{-1} vibration referred to the collagen integrity, showed that collagen were not degraded during treatment for all skin samples (data not shown)\textsuperscript{45, 46}. Corresponding spectral FTIR images
were constructed using the Amide I/Amide II integrated intensities ratio for the non-polarized [Fig. 1(c) and 1(d)] and for the perpendicular polarization [Fig. 1(e) and 1(f)]. By comparing the images collected in non-polarized mode with those acquired in perpendicular polarization, it is noteworthy that the polarization of IR signal increased markedly the Amide I/Amide II ratio, permitting to enhance the image contrast and to reveal an orientation of the peptide bonds of the protein components (i.e., mainly collagen) of the skin. Compared to the values obtained from tendon analyses, the color-scale was adjusted from 1 (blue) to 2 (red). Three categories of orientation can be distinguished: Amide I/Amide II ratio ≥ 1.6 corresponds to collagen fibers oriented parallel to the skin surface, and ≤ 1.3 to those oriented perpendicular to the skin surface. A value between 1.3 and 1.6 could be associated to an oblique orientation of collagen fibers. The same methodology was also applied for the 38-, 60- and 66-year-old human skin samples, untreated [Fig. S2(a, c, e), Fig. S3(a, c, e) and Fig. S4(a, c, e)] and treated with D$_2$O [Fig. S2(b, d, f), Fig. S3(b, d, f) and Fig. S4(b, d, f)] (Supporting Information).

Redundancy-based algorithm (RBA)-fuzzy C-means (FCM) clustering on FTIR spectral images

In order to distinguish more precisely the different regions of the dermis, RBA-FCM clustering was performed to the perpendicular polarized FTIR data acquired from 35-year-old human skin sections, untreated and treated with D$_2$O (Fig. 2). The same processing was applied on 38-, 60- and 66-year-old human skin sections, treated or not with D$_2$O (Fig. S5, S6 and S7 in Supporting Information). With this automatic clustering, the cluster #1 represents the epidermis and clusters #2, 3, 4 and 5 are associated to the dermis. By comparing these latter images with those created from the Amide I/Amide II ratio, it appears that the clusters corresponding to dermis match quite well to different categories of collagen orientation. Thus,
the cluster #2 can be associated to collagen fibers oriented parallel to the skin surface and, conversely, collagen fibers oriented perpendicular to the skin surface correspond to the cluster #4. Collagen fibers with an oblique orientation are found in the cluster #3. Presenting a marked spatial dispersion of pixels for all skin sections, having a low signal-to-noise ratio and being minority with regard to other clusters, the cluster #5 was not considered in the following results.

Table 1 summarizes the spatial repartition (in terms of relative proportion of pixels) of each orientation (parallel, perpendicular and oblique) of collagen fibers for different-age human skin samples, treated or not with D₂O. Values were averaged on 5 cryosections for each skin sample. For the 35- and 38-year-old skin samples, results show an increase (from 25.1% to 34.3%) of the proportion of perpendicular orientations of collagen fibers (cluster #4), a decreasing trend (38.2% to 27.5%) of the proportion of oblique orientations (cluster #3), and no significant changes for parallel orientations (cluster #2) with D₂O treatment. For the 60- and 66-year-old skin samples, no significant changes associated to the D₂O treatment are noticed for any orientation of collagen fibers.

Collagen-bound water-related spectral markers from different-age skin samples

In order to identify vibrations associated to changes of collagen fiber orientation, extracted spectra from the cluster #4 (perpendicular oriented collagen fibers) of different-age and – treatment of skin samples were compared using the randfeatures algorithm. Due to significant differences as previously presented in Table 1, only the cluster #4 was considered in this analysis. A selection of 3000 spectra, extracted from the 5 different cryosections for each skin sample and equally distributed between the untreated and the D₂O-treated skin samples, was performed. Fixing a score > 30, 11 and 8 most discriminant wavenumbers were retained respectively for the 35-year-old [Fig. 4(a)] and for the 66-year-old skin samples [Fig. 4(b)],
and were arranged in the form of spectral color barcode according to the discrimination level of the scores. The same procedure was also applied on 38- and 60-year-old human skin sections, giving similar results (Fig. S8 in Supporting Information). The color codes were used as follows: *orange* for the less discriminant wavenumbers (scores between 30 and 33); *brown* for the intermediate discriminant (scores between 34 and 36); and *red* for the most discriminant (scores ≥ 37). It can be noted that 8 wavenumbers in 1650 cm⁻¹, 1555 cm⁻¹, 1455 cm⁻¹, 1390 cm⁻¹, 1330 cm⁻¹ and between 1300 and 1200 cm⁻¹, were found common to the skin types. They correspond mainly to the vibrations specific of proteins. Besides, three additional discriminant wavenumbers in 1120 cm⁻¹, 1080 cm⁻¹ and 1050 cm⁻¹, corresponding to carbohydrates vibrations, were found exclusively in the young skin sample. Their peak position and molecular attribution are summarized in Table 2.

**Discussion**

Understanding the interaction between dermal collagen network and its surrounding environment is of interest to gain insight on molecular and functional changes undergone by the cutaneous tissue during aging. Type I collagen is the main macromolecule of the dermis able to form a fibrous network and is strongly involved in the mechanical properties of the skin as well as in its hydration. Indeed, water plays a crucial role for maintaining the assembly formed by collagen macromolecules. Most of water molecules are tightly bound to proteins and consequently intervene in the skin microstructure. Apart these interacting water molecules, there are water molecules, called bulk or tetrahedron water, not bound to proteins but more likely bound to each other⁴⁷. Treatment by heavy water is a way to investigate the water/collagen interactions as previously used to observe changes of the collagen model peptides stability and to highlight hydration forces between collagen triple helices⁴⁸,⁴⁹. More recently, other groups realized H/D exchange on different tissues such as skin, pericardium
and bone in order to remove primarily water molecules bound to collagen fibers\textsuperscript{9, 50, 51}. Substitution of bound water molecules by D\textsubscript{2}O is possible due to the respective thermodynamic properties of water and heavy water. Indeed, although these two molecules have a similar size, deuterium bonds are stronger than hydrogen bonds with dissociation energies of 6.53 kJ mol\textsuperscript{-1} and 5.52 kJ mol\textsuperscript{-1}, respectively, according to theoretical models\textsuperscript{52, 53}. In our study, the effect of H\textsubscript{2}O/D\textsubscript{2}O substitution was considered by studying the collagen fibers orientation, structural information that can be probed within skin specimens by exploiting the polarization of the IR radiation\textsuperscript{34, 41}. Such changes of behavior of collagen network observed between skins of different ages are very likely to be linked with a degradation of the mechanical properties, whose one of the visible signs is the appearance of wrinkles. Our approach relies on a quantification of the effect of H\textsubscript{2}O/D\textsubscript{2}O substitution on collagen orientation by comparing skins from two different age groups.

A particular statistical clustering method (RBA-FCM) was used to overcome the inconvenient of more classical techniques such as K-Means which the result is highly dependent of the number of the clusters chosen by the operator. In RBA-FCM processing, the number of clusters is automatically determined facilitating the comparison of samples since it is not subject to the operator interpretation. Thus, distinct parts associated to different orientations of collagen fibers were highlighted in the dermis: parallel (cluster #2), oblique (cluster #3), and perpendicular (cluster #4) to the skin surface. It is worth noting that pixel spatial distributions of parallel and perpendicular orientations appear to be packed, whereas that of the oblique orientation seems to be sparse. Concerning the cluster associated to the parallel oriented collagen fibers, the quantitative results were obtained by taking into account both the papillary dermis and a part of the reticular dermis. The examination of the FCM images (cluster #2) showed that the part corresponding to the papillary dermis decreases in
the samples of the oldest group, which is consistent with the loss of papillary dermal compartment observed during aging\textsuperscript{54}.

Thereafter, regarding the impact of D\textsubscript{2}O treatment, in the youngest skins, the H/D exchange led to favor perpendicular orientation of collagen fibers at the expense of oblique orientation. On the contrary, the D\textsubscript{2}O treatment appears to not affect significantly the proportions of different orientations of collagen fibers in the old skin samples. These observations could reflect differences between the age groups in the number of water molecules bound to collagen and also in their interaction strength. It is hard to study in details the involved molecular mechanisms; additional experimental approaches would be required to assess thermodynamics behaviors of water molecules interacting with macromolecular proteins by using for example dynamic vapor sorption and scanning electron microscopy to study simultaneously the organization of collagen network at the nanoscale\textsuperscript{9, 55}. Here, our objective was humbly to show the sensitivity of FTIR imaging in highlighting specific behaviors for skin samples of different ages. We focused our interpretation on the collagen that is by far the main component of the dermis. We demonstrated from reference tendon samples that P-FTIR is a very sensitive to probe collagen fibers orientation. However, another macromolecular component, elastin, is known to be affected by ageing and also to be directly involved in the skin mechanical properties\textsuperscript{56} even though the elastic fiber network represents only 5\% of the dermis dry mass. By analyzing nuchal bovine ligament, we checked that FTIR signal of elastin appeared to be much less sensitive to the polarization of the incident beam that collagen (data not shown). Consequently, elastic fibers were not considered in our data interpretation but it would deserve to bring attention to it, possibly by investigating their spatial distribution thanks to their intrinsic fluorescent properties by using multiphoton microscopy\textsuperscript{57}. 
In addition to determine the relative proportion of the different collagen fiber orientations, infrared wavenumbers affected by the D$_2$O treatment were identified using statistical processing. Obviously, protein vibrations are pinpointed for the two age groups. But interestingly, vibrations were highlighted in the 1000-1150 cm$^{-1}$ range for the youngest skin samples. This latter spectral region is often associated to sugars originating from advanced glycation endproducts$^{40, 58}$, from PGs$^{29, 32}$, and in a much less extent to nucleic acids$^{59, 60}$. As PGs represent the major part of carbohydrates in the dermis, their interaction with collagen fibers is very probably altered during the H/D exchange. These results are consistent with the involvement of GAG in collagen hydration. Indeed, GAGs from PGs have a key-role in skin regarding to their ability to bind up to 1000 times their volume in water, due to their high polarity. Therefore, skin hydration is highly correlated to the content and distribution of dermal GAGs$^{61}$. Recently, Li et al. demonstrated that the size of GAG chain of decorin, the most abundant PGs in human dermis, was reduced in sun-protected old skin compared to young skin$^{62}$. Alterations of this GAG entity of decorin could explain the lack of interaction between water molecules and GAGs in old dermis skin and could contribute to skin fragility of elderly people. Playing a crucial role in the hydration of collagen fibers, PGs could therefore affect the structure but also the orientation of the collagen fibers in young skin. PGs may less interact with collagen with age and consequently lead to a decrease in the hydration of collagen fibers. This latter could be related to the changes of orientation of collagen fibers demonstrated in our study.

In this study, various methodological approaches of sample preparation (heavy water substitution), of data acquisition (polarization mode) or of data processing (automatic clustering and discriminant wavenumbers selection), were applied to investigate alterations of the dermis associated to the chronological aging at the molecular level. Similar approaches could be used in other issues without complicated adaptation, such as in the context of photo-
aging where water structure and interactions with proteins and GAGs were demonstrated by Gniadecka et al. to be altered in photo-aged skins; and also in more medical issues such as age-related diseases like diabetes. In addition, they could be of interest the characterization of skin cancer, as shown in a recent study reporting that the in vivo determination of water content by using vibrational Raman spectroscopy may contribute to differentiate between tumor and healthy tissue.

**Conclusion**

This study exhibits the potential of P-FTIR microimaging associated to advanced statistical clustering methods to probe changes of collagen fibers orientations between skin samples of different ages. Recent devices equipped with FPA matricial detectors make feasible in routine use of the technique by reducing considerably the acquisition time of tissue samples. D₂O treatment affects significantly only the youngest skin samples, quite possible via PG involvement. This methodology is promising for a better insight on relationship between water molecules and collagen fiber organization upon chronological aging, bringing a potential added value as well in cosmetics as in dermatology.

**Acknowledgements**

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**References**


Figures captions

Figure 1: FTIR imaging on 35-year-old skin sections treated or not with D$_2$O. H&E stained adjacent tissue sections from untreated (a) and treated (b) skin sample. Corresponding Amide I/Amide II (integrated intensities: 1590-1720 cm$^{-1}$/1480-1590 cm$^{-1}$) ratio-based reconstructed FTIR images in the non-polarized mode (c, d) and with the perpendicular polarization (e, f). Scale bar = 50 µm.

Figure 2: FCM clustering on perpendicular polarized FTIR images from untreated (a) and treated with D$_2$O (b) 35-year-old skin section. Assignment of the clusters: cluster 1 (epidermis); clusters 2, 3, 4 and 5 (dermis).

Figure 3: Discriminant wavenumbers identified by using randfeatures algorithm in perpendicular oriented collagen fibers, (a) between D$_2$O-untreated (in red) and treated (in blue) 35-years-old dermal skin samples and (b) D$_2$O-untreated (in orange) and treated (in black) 66-years-old dermal skin samples. Respective scores are indicated above color-coded discriminant wavenumbers according to their discrimination level (30-33: orange; 34-36: brown; ≥ 37: red).

Figure S1: FTIR imaging on rat-tail tendon section. Amide I/Amide II (integrated intensities: 1590-1720 cm$^{-1}$/1480-1590 cm$^{-1}$) ratio-based reconstructed FTIR images and mean spectra in the non-polarized mode (a) and with the perpendicular polarization (b).

Figure S2: FTIR imaging on 38-year-old skin sections treated or not with D$_2$O. H&E stained adjacent tissue sections from untreated (a) and treated (b) skin sample. Corresponding Amide I/Amide II (integrated intensities: 1590-1720 cm$^{-1}$/1480-1590 cm$^{-1}$) ratio-based reconstructed
FTIR images in the non-polarized mode (c, d) and with the perpendicular polarization (e, f).
Scale bar = 50 µm.

**Figure S3:** FTIR imaging on 60-year-old skin sections treated or not with D$_2$O. H&E stained adjacent tissue sections from untreated (a) and treated (b) skin sample. Corresponding Amide I/Amide II (integrated intensities: 1590-1720 cm$^{-1}$/1480-1590 cm$^{-1}$) ratio-based reconstructed FTIR images in the non-polarized mode (c, d) and with the perpendicular polarization (e, f).
Scale bar = 50 µm.

**Figure S4:** FTIR imaging on 66-year-old skin sections treated or not with D$_2$O. H&E stained adjacent tissue sections from untreated (a) and treated (b) skin sample. Corresponding Amide I/Amide II (integrated intensities: 1590-1720 cm$^{-1}$/1480-1590 cm$^{-1}$) ratio-based reconstructed FTIR images in the non-polarized mode (c, d) and with the perpendicular polarization (e, f).
Scale bar = 50 µm.

**Figure S5:** FCM clustering on perpendicular polarized FTIR images from untreated (a) and treated with D$_2$O (b) 38-year-old skin section. Assignment of the clusters: cluster 1 (epidermis); clusters 2, 3, 4 and 5 (dermis).

**Figure S6:** FCM clustering on perpendicular polarized FTIR images from untreated (a) and treated with D$_2$O (b) 60-year-old skin section. Assignment of the clusters: cluster 1 (epidermis); clusters 2, 3, 4 and 5 (dermis).
**Figure S7:** FCM clustering on perpendicular polarized FTIR images from untreated (a) and treated with D$_2$O (b) 66-year-old skin section. Assignment of the clusters: cluster 1 (epidermis); clusters 2, 3, 4 and 5 (dermis).

**Figure S8:** Discriminant wavenumbers identified by using randfeatures algorithm in perpendicular oriented collagen fibers, (a) between D$_2$O-untreated (*in red*) and treated (*in blue*) 38-years-old dermal skin samples and (b) D$_2$O-untreated (*in orange*) and treated (*in black*) 60-years-old dermal skin samples. Respective scores are indicated above color-coded discriminant wavenumbers according to their discrimination level (30-33: orange; 34-36: brown; ≥ 37: red).

**Tables captions**

**Table 1:** Quantitative analysis of different orientations (parallel, oblique and perpendicular) of collagen fibers in the different-age skin samples, untreated (- D$_2$O) and treated (+ D$_2$O) with D$_2$O. The percentages were calculated from the number of pixels in the clusters associated to the different orientations. Standard deviations were obtained after averaging data collected on 5 analyzed sections for each skin sample.

**Table 2:** FTIR spectral peak attributions.
Figure 1: FTIR imaging on 35-year-old skin sections treated or not with D₂O. H&E stained adjacent tissue sections from untreated (a) and treated (b) skin sample. Corresponding Amide I/Amide II (integrated intensities: 1590-1720 cm⁻¹/1480-1590 cm⁻¹) ratio-based reconstructed FTIR images in the non-polarized mode (c,d) and with the perpendicular polarization (e,f). Scale bar = 50 µm.
Figure 2: FCM clustering on perpendicular polarized FTIR images from untreated (a) and treated with D$_2$O (b) 35-year-old skin section. Assignment of the clusters: cluster 1 (epidermis); clusters 2, 3, 4 and 5 (dermis).
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<table>
<thead>
<tr>
<th>Collagen fiber orientation</th>
<th>Young skin samples (35- and 38-year-old)</th>
<th>Old skin samples (60- and 66-year-old)</th>
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<tr>
<td></td>
<td>- D$_2$O</td>
<td>+ D$_2$O</td>
</tr>
<tr>
<td>Parallel</td>
<td>36.7 ± 1.6 %</td>
<td>38.2 ± 2.2 %</td>
</tr>
<tr>
<td>Oblique</td>
<td>38.2 ± 1.9 %</td>
<td>27.5 ± 1.2 %</td>
</tr>
<tr>
<td>Perpendicular</td>
<td>25.1 ± 2.5 %</td>
<td>34.3 ± 1.6 %</td>
</tr>
</tbody>
</table>
Figure 3: Discriminant wavenumbers identified by using randfeatures algorithm in perpendicular oriented collagen fibers, (a) between D$_2$O-untreated (in red) and treated (in blue) 35-years-old dermal skin samples and (b) D$_2$O-untreated (in orange) and treated (in black) 66-years-old dermal skin samples. Respective scores are indicated above color-coded discriminant wavenumbers according to their discrimination level (30-33: orange; 34-36: brown; ≥ 37: red).
Table 2: FTIR spectral peak attributions.

<table>
<thead>
<tr>
<th>Peak position (cm⁻¹)</th>
<th>Possible vibrations</th>
<th>Assignments</th>
</tr>
</thead>
<tbody>
<tr>
<td>1650</td>
<td>Amide I (C=O stretch)</td>
<td>Proteins</td>
</tr>
<tr>
<td>1550</td>
<td>Amide II (C-N stretch, N-H bend)</td>
<td>Proteins</td>
</tr>
<tr>
<td>1455</td>
<td>C-H bend</td>
<td>Proteins</td>
</tr>
<tr>
<td>1390</td>
<td>C-H and N-H bend</td>
<td>Proteins</td>
</tr>
<tr>
<td>1340</td>
<td>CH₂ wagging</td>
<td>Proteins</td>
</tr>
<tr>
<td>1280</td>
<td>Amide III (C-N stretch, N-H bend)</td>
<td>Proteins</td>
</tr>
<tr>
<td>1240</td>
<td>Amide III (C-N stretch, N-H bend)</td>
<td>Proteins</td>
</tr>
<tr>
<td>1205</td>
<td>Amide III (C-N stretch, N-H bend)</td>
<td>Proteins</td>
</tr>
<tr>
<td>1115</td>
<td>C-O stretch, C-O bend</td>
<td>Carbohydrates</td>
</tr>
<tr>
<td>1080</td>
<td>C-O stretch, C-O bend</td>
<td>Carbohydrates</td>
</tr>
<tr>
<td>1030</td>
<td>C-O stretch, C-O bend</td>
<td>Carbohydrates</td>
</tr>
</tbody>
</table>

Table 2: FTIR spectral peak attributions.
Figure S1: FTIR imaging on rat-tail tendon section. Amide I/Amide II (integrated intensities: 1590-1720 cm⁻¹/1480-1590 cm⁻¹) ratio-based reconstructed FTIR images and mean spectra in the non-polarized mode (a) and with the perpendicular polarization (b).
**Figure S2:** FTIR imaging on 38-year-old skin sections treated or not with D$_2$O. H&E stained adjacent tissue sections from untreated (a) and treated (b) skin sample. Corresponding Amide I/Amide II (integrated intensities: 1590-1720 cm$^{-1}$/1480-1590 cm$^{-1}$) ratio-based reconstructed FTIR images in the non-polarized mode (c, d) and with the perpendicular polarization (e, f). Scale bar = 50 µm.
Figure S3: FTIR imaging on 60-year-old skin sections treated or not with D$_2$O. H&E stained adjacent tissue sections from untreated (a) and treated (b) skin sample. Corresponding Amide I/Amide II (integrated intensities: 1590-1720 cm$^{-1}$/1480-1590 cm$^{-1}$) ratio-based reconstructed FTIR images in the non-polarized mode (c, d) and with the perpendicular polarization (e, f). Scale bar = 50 µm.
Figure S4: FTIR imaging on 66-year-old skin sections treated or not with D$_2$O. H&E stained adjacent tissue sections from untreated (a) and treated (b) skin sample. Corresponding Amide I/Amide II (integrated intensities: 1590-1720 cm$^{-1}$/1480-1590 cm$^{-1}$) ratio-based reconstructed FTIR images in the non-polarized mode (c, d) and with the perpendicular polarization (e, f). Scale bar = 50 µm.
Figure S5: FCM clustering on perpendicular polarized FTIR images from untreated (a) and treated with D₂O (b) 38-year-old skin section. Assignment of the clusters: cluster 1 (epidermis); clusters 2, 3, 4 and 5 (dermis).
Figure S6: FCM clustering on perpendicular polarized FTIR images from untreated (a) and treated with D₂O (b) 60-year-old skin section. Assignment of the clusters: cluster 1 (epidermis); clusters 2, 3, 4 and 5 (dermis).
**Figure S7**: FCM clustering on perpendicular polarized FTIR images from untreated (a) and treated with D$_2$O (b) 66-year-old skin section. Assignment of the clusters: cluster 1 (epidermis); clusters 2, 3, 4 and 5 (dermis).
Figure S8: Discriminant wavenumbers identified by using randfeatures algorithm in perpendicular oriented collagen fibers, (a) between D$_2$O-untreated (in red) and treated (in blue) 38-years-old dermal skin samples and (b) D$_2$O-untreated (in orange) and treated (in black) 60-years-old dermal skin samples. Respective scores are indicated above color-coded discriminant wavenumbers according to their discrimination level (30-33: orange; 34-36: brown; ≥ 37: red).