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Thermal and vibrational characterization of human skin

Influence of the freezing process

Rong Tang¹ · Valerie Samouillan¹ · Jany Dandurand¹ · Colette Lacabanne¹ · Florence Nadal-Wollbold² · Christiane Casas² · Anne-Marie Schmitt²

Abstract For a better understanding of the molecular and organizational changes in human dermis, biophysical methods were tested. The aim of this study was to find suitable and reproducible biomarkers for further clinical studies on intrinsic and extrinsic aging of dermis. Thermoporometry, hydric organization and thermal transitions of fresh and frozen skins were determined by differential scanning calorimetry (DSC). Fourier transform infrared spectroscopy (FTIR) was used to identify the absorption bands of the dermis especially in the 1800–1000 cm⁻¹ zone and to discriminate between the different secondary structures of proteins. A widening of the pore size distribution is evidenced with freezing, but there is no significant difference between the hydric organization and the endothermic collagen denaturation of fresh and frozen skins. The global FTIR spectra and the second derivative spectra in the scanned zone are also identical in fresh and frozen dermis, validating the storage protocol. DSC and FTIR are well-suited techniques to characterize human skin, giving accurate results with high reproducibility. The acquisition of thermal and vibrational biomarkers of the skin at the mesoscale and nanoscale contributes to its better knowledge and is promising for further studies on skin aging.

Keywords Collagen denaturation · DSC · Fourier transform infrared spectroscopy · Human skin · Hydration · Secondary structures of proteins

Introduction

With a growing elderly population, interest in general skin health and skin homeostasis has been increasing. Understanding hydration in skin is of primary importance to the medical and cosmetic communities. In particular, the ability to differentiate protein bound from bulk water is of interest in studies ranging from the evaluation of therapeutics for wound healing to the hydration efficacy of skin care products. Skin is the largest body organ and is easily accessible. A great number of clinical studies have been performed on skin biopsies to help the understanding on mechanisms involved in intrinsic and extrinsic (photo-) aging [1, 2] using histological and ultrastructural approaches. Unfortunately, the complexity and slowness of the aging processes leading to subtle changes of biological function (including biochemical, morphological, physical aspects) provide the aging research with enormous difficulties.

The skin consists of the epidermis and dermis, and the latter is thought to be more important in determining the physical properties of skin. Among the various components of the dermis, extracellular matrix consisting of collagens, elastic fibers and amorphous ground substances is strongly involved in the process of cutaneous aging [3], in close correlation with changes in hydration [4, 5]. Collagen, the most abundant component in the dermis, comprises 80 % of the tissue total dry mass and 90 % of all dermal proteins. If at least half of the twenty eight types of collagens are present in skin, the fibril-forming collagens I, III and V are

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the most abundant. Collagens I and III represent close to 90 and 10%, respectively, in the composition of dermal collagen fibrils, and collagen V is present as a minor fraction of about 2%. Altogether, the fibrils with their associated proteins confer tensile strength to the skin and are pivotal for the general organization and stability of the dermal extracellular matrix among other functions [6]. The dermis can also be divided into two layers: the papillary dermis and the reticular dermis. The papillary dermis layer contains much thinner and less packed of collagen type III fibers, and it is more fibroblastic than the reticular dermis layer [7]. It is also shown that the papillary dermis contains much more components of glycosaminoglycans such as hyaluronic acid and dermatan sulfate [8]. In the dermis, collagen bundles form a three-dimensional mesh with other fibrillar proteins and interstitial cells, including a network of contractile fibroblasts. The structure is cemented by a hyaluronic acid and proteoglycan gel that allows for viscous flux and mediates interchange of water nutrients and metabolites between the blood, the lymph, the interstitial cell network and the epidermal layers [9]. The recent studies propose a refined skin-associated adipocytes as dermal adipose tissue, historically termed subcutaneous adipose, hypodermis and subcutis, which underlie the reticular dermis. The major hygroscopic agents, glycerol and hyaluronic acid play a key role in skin hydration and consequently prevent skin aging [10]. Furthermore, the cutaneous aging could be accelerated by the decrease of overall collagen types after menopause [11].

New information on whole skin and dermis can be reached at the molecular level through near-infrared diffuse reflection (NIR-DR) [12, 13]. Fourier transform infrared spectroscopy/attenuated total reflectance (FTIR/ATR) [12, 14, 15], Raman spectroscopy [16–19], nuclear magnetic resonance/magnetic resonance imaging (NMR/MRI) [15, 20, 21], dynamic vapor sorption (DVS) [19], changes in collagen hydration with age yield somewhat conflicting results, which could be attributed to the preparation of samples.

Combined with thermogravimetric analysis (TG) measurements, differential scanning calorimetry (DSC) is a standard method to evaluate freezeable and unfreezable water [22–25]. DSC is also an appropriate method for assessing protein thermal stability and conformational changes. It is peculiarly well suited to evaluate the thermal stability of purified collagens in solution or in their aggregated form [26], or directly in native tissues [26–34] and biomaterials [28–30]. It has been successfully applied to characterize collagen both in animal and human skins [35–37]; nevertheless, few DSC data are available on human dermis, despite the ability of this technique to evidence evolution of the major proteins of dermis under physical or pathological factors, such as collagen alteration in irradiated human dermis [37], increased collagen stability with glycation induced by diabetes [38] or modification of collagen I/III in scars and keloids [39].

If punch biopsies from living patients are routinely taken for histologic assessment in vitro, few biophysical studies have been performed on such samples. The first objective of this work is to evaluate the effect of a quenching/storage protocol on fresh human skin explants for the scheduled clinical study on a large panel of young and aged patients. Simultaneously, it is essential to validate the combined use of DSC and FTIR techniques to obtain reproducible data from these small biopsies removed from living patients and to establish a routine protocol to analyze an important number of samples. The final aim of this work was to extract suitable biomarkers of the hydric organization and biomacromolecules integrity from the thermal and vibrational fingerprints of safe skin.

**Materials and Methods**

**Materials**

Fresh human abdominal skin disks (22 mm by diameter) were collected by BIOPREDIC Society (BIOPREDIC INTERNATIONAL, SAINT GREGOIRE, FRANCE) from 5 female patients (Caucasian type, 25–50 years old) undergoing abdominoplasty. The skin samples were first rinsed with PBS. Then, the adipose tissue/hypodermis was removed. The handling of the fresh skin was done under hood (natural atmosphere) to minimize contamination. Fresh portions were stored at 4°C for 24 h, and explants (4 mm by diameter, 5–7 mg by mass) were removed from these portions using a biopsy punch. Twenty explants could be extracted for a skin disk. A series of explants (fresh explants) were immediately analyzed. Another series of explants (stored explants) were stored in Eppendorf tubes and then quenched in liquid nitrogen for several minutes and stored at −20°C for 8 days. Before further analyses, these frozen samples were thawed at 4°C for 1 h.

**Differential scanning calorimetry (DSC)**

DSC measurements were performed with a DSC Pyris calorimeter (PERKIN ELMER, Waltham, MA) using an empty pan as reference. The calorimeter was calibrated using the manufacturer’s instructions with pure water, cyclohexane and indium as standards, resulting in a temperature accuracy of ±0.1°C and an enthalpy accuracy of ±2.10⁻⁴ J kg⁻¹. Temperature calibration was undertaken at each scanning rate. Fresh and frozen explants (5–7 mg by mass) were sealed in hermetic aluminum pan. Samples were cooled at 10°C min⁻¹ to −30°C (1st cooling scan),
then held 10 min to freeze water. For thermoporometry measurements (TPM), scans were conducted from −30 to 20 °C at a low scanning rate, 0.5 °C min⁻¹, to avoid thermal and time delays in the DSC curve.

To determine freezeable water amount and intrinsic thermal transitions, the samples were heated at 10 °C min⁻¹ to 85 °C (1st heating scan), then cooled at −10 °C min⁻¹ to −30 °C (2nd cooling scan) and then heated at 10 °C min⁻¹ to 85 °C (2nd heating scan).

After completing the DSC measurements, pans were reweighted to check that they had been correctly sealed. The sample pans were pierced and dried to constant mass at 195 °C for 10 min to determine the sample dry mass and the total water mass.

Fourier transform infrared analysis (FTIR)

Fourier transform infrared spectroscopy/attenuated total reflectance (FTIR/ATR) spectra were acquired using a Nicolet 5700 FTIR (THERMO FISHER SCIENTIFIC, Waltham, MA) equipped in ATR device with a KBr beam splitter and a MCT/B detector. Explants (dermis face) were directly laid on the ATR accessory (Smart Orbit with a type II A diamond crystal, refractive index 2.4) and covered by a hermetic cap with an ‘O’ ring to avoid dehydration of the IIA diamond crystal, and then held 10 min to freeze water. For thermoporometry measurements (TPM), scans were conducted from −30 to 20 °C at a low scanning rate, 0.5 °C min⁻¹, to avoid thermal and time delays in the DSC curve.

Results and discussion

Thermal characterization

Thermoporometry measurements (TPM)

DSC curves of fresh and stored explants are plotted in Fig. 1. In both curves, two distinct endothermic peaks were recorded in the zone of ice melting, what is an evidence for the existence of pores filled by water in the explants. It is well known that structure and connectivity of pores in skin influence the heat and mass transport processes associated with the thermoregulatory function of the organ [40]. As commonly observed in mesoporous hydrophilic materials, the low temperature peaks are assigned to the melting of the ice confined in the pores while the second ones result from the melting of the ice outside the pores (excess water) [41–44].

According to the Gibbs–Thomson effect, Brun et al. [41] established that the radius \( r_p \) of hypothetical cylindrical pores filled by water and the differential pore volume \( dV_p/\text{dr}_p \) could be calculated from the melting point depression \( \Delta T \) and heat flow \( dQ/dt \). The following equations are as follows:

\[
\begin{align*}
 r_p (\text{nm}) &= -32.33 \frac{T}{T_f} + 0.68 \\
 &\text{with } \Delta T = T - T_f \quad \text{and} \quad -40 < \Delta T < 0 \\
 \frac{dV_p}{dr_p} &= \frac{dQ}{dt} \times \frac{dr}{d(\Delta T)} \times \frac{d(\Delta T)}{dr_p} \times \frac{1}{m \Delta H_f(T) \rho(T)}
\end{align*}
\]

where \( d(\Delta T)/dt \) is the scanning rate of the DSC experiments, \( m \) is the mass of dry material, and \( \Delta H_f \) and \( \rho \) the heat of fusion and density of water, respectively. The

![Fig. 1](https://via.placeholder.com/150)

Statistical analysis

All experiments were performed with at least three replicates for each group and for at least three independent repeats. Quantitative values are shown as mean ± SEM. In statistics, we practiced one-way analysis of variance (one-way ANOVA) to compare the means of the samples. The \( p \)-value was also used to determine the significant difference between groups. It was considered statistically significant threshold of \( p \)-value less than 0.05.
following values were used: \( \Delta H_2(T) = (334.1 + 2.1197 - 0.00783T^2) \times 10^{-3} \) (J kg\(^{-1}\)) and \( \rho(T) = 917 (1.032 - 1.17 \times 10^{-7}T) \) (kg m\(^{-3}\)) [45].

For the present study, pore radius was calculated on the assumption of cylindrical pore shape. This is unlikely to be absolutely correct, but is believed to be a reasonable approximation for the elongated voids expected within skin, and it has been successfully used for fibrous materials [40, 46, 47]. Using Eq. (1), the temperature axis of TPM measurements can be converted into a pore size scale; in this study, \( T_0 \) was determined experimentally as the onset temperature of the second endothermic peak. After a baseline subtraction step that effectively removes the underlying heat capacity contribution to the DSC signal, the heat flow curve, \( dQ/dt \), was converted into a differential pore volume using Eq. (2). The resulting pore size distributions (PSD) determined for fresh and stored explants are reported in Fig. 2. From Fig. 2, we can note a relatively narrow size distribution for the fresh skin; since the radius of pores range from 0 to 20 nm, these pores can be classified as mesopores. The storage protocol (quenching and storage at \(-20^\circ C\) during 8 days) leads to a widening of the pore size distribution, in peculiar enhancing the formation of macropores with radius from 20 to 60 nm.

Even if the scans at 0.5 \(^{\circ}C\) min\(^{-1}\) can be used to quantify total freezable water, the main objective of this study was to establish a robust protocol to determine with a single DSC scan, preferentially low time-consuming, both the hydration and the intrinsic transitions of dermal proteins. Comparison of DSC curves performed at different scanning rates between 0.5 and 20 \(^{\circ}C\) min\(^{-1}\) (not shown here), evidenced that a scanning rate of 10 \(^{\circ}C\) min\(^{-1}\) was the optimal choice to reduce the experimental time and to measure reproducible and well-defined ice melting and protein denaturation in skin biopsies.

**Water quantification and collagen denaturation**

In Fig. 3 are reported the DSC curves (normalized to the initial mass) of fresh and stored explants corresponding to the first cooling between 20 and \(-30^\circ C\) and the successive heating at 10 \(^{\circ}C\) min\(^{-1}\) between \(-30\) and 30 \(^{\circ}C\); for the sake of clarity, only one curve for each type of sample is reported and an enlargement is made in the temperature zone of ice crystallization/melting. The discussion has focused on melting peaks only, because the ice crystallization temperature is usually not reproducible for a super cooled liquid [42]. As already observed in TPM measurements at 0.5 \(^{\circ}C\) min\(^{-1}\), the endothermic peaks recorded in this temperature zone correspond to the melting of freezable water, even if in this case the resolution into separated peaks is not achieved because of the scanning rate. This extrinsic transition is widely used to quantify the amount of total freezable water in hydrated proteins and tissues (by dividing the area of the measured endothermic peak by 0.334 J kg\(^{-1}\), corresponding to the melting enthalpy of pure ice at 0 \(^{\circ}C\)) [23, 48], and completes thermogravimetric analyses giving the total amount of water. The amount of unfreezable water can be calculated by a simple difference. This quantification was performed for all the samples and reported in Fig. 4.

The total hydration of fresh skin is 66.2 \(\pm\) 1 %, distributed in 41.7 \(\pm\) 0.5 % of freezable water and 24.5 \(\pm\) 1.1 % of unfreezable water. It must be pointed out that the determination of freezable water from the area of the endotherms recorded at 0.0083 \(^{\circ}C\) s\(^{-1}\) leads to similar values.

**Fig. 2** Representative differential pore size distributions from TPM measurements for a matched set of fresh and stored skin explants.

**Fig. 3** Representative DSC curves for a matched set of fresh and stored skin explants [cooling and successive heating, scanning rate 10 \(^{\circ}C\) min\(^{-1}\) and enlargement in the \((-30; 30^\circ C)\) window]. Whole curves in the inset.
The total hydration of fresh skin is consistent with literature data, [10, 36], giving an average value of 65–70 % for the hydration of human dermis and epidermis; this remarkably high value, corresponding to about 2 kg of water per kg of dry matter, is due to a combination of physical and chemical factors including the presence of interconnected pores, the peculiar hygroscopic nature of hyaluronic acid and the hydrophilicity of collagen. If a well-documented literature is present for the total hydration of stratum corneum, epidermis and dermis measurable by different biophysical techniques [5], few data, obtained from Raman spectroscopy [19, 49], DVS [19] and NMR [21] are reported on the quantification of the different kinds of water (namely free and bound waters) in human skin. The present DSC study shows that the freezable water roughly represents two-third of the total water in human skin. The term freezable water is preferred to the term ‘free’ water—rather reserved to vibrational or relaxational techniques—since it covers bulk water in excess but also confined water in mesopores, as previously shown by TPM. In accordance with previous NMR, DVS and Raman results, this work evidences that bulk water is in relatively large quantity in safe human skin.

One-third of the total water of human skin is unfreezable water. To avoid confusion, in this study, the term ‘unfreezable water’ is preferred to the term ‘bound’ water, which refers to water directly bound to protein via hydrogen bonds in the first hydration shell, but that can also be extended to the multilayer hydration shell and identified as a slowing down of the dynamics in dielectric and NMR techniques [24, 50–54]. Besides the idea of monolayer, multilayer and free water as defined for S-shaped sorption isotherms, the types of water can be also divided by their behavior during freezing and thawing, and in this case bound water is sometimes divided in ‘unfreezable bound water’—strongly associated with hydrophilic groups—and ‘freezable bound water’—less closely bound and immobilized in DSC experiments [55, 56]. In our case, unfreezable water is ‘unfreezable bound water’ and it corresponds to the filling of the first hydration shell of proteins and other hydrophilic components such as HA; the value of 24.5 % of unfreezable water, corresponding to 0.72 kg of unfreezable water per kg of dry matter, is consistent with literature data on biological tissues [22, 57]. Usually, about 0.3–0.7 kg of unfreezable water remains associated per kilogram of a ‘dry’ protein [58]. Assuming that the average molar mass of one amino acid in collagen is 0.1 kg mol⁻¹, a roughly approximation leads to a value of 4 mol of water per moles of amino acid of collagen, which is slightly higher than the value determined by DVS [59] or X-ray diffraction for adsorbed multilayer water in collagen and collagen peptides [60], certainly because of the presence of HA in dermis.

The statistical analysis do not indicate any significant difference between the amounts of total, freezable and unfreezable waters in fresh and stored skins, evidencing that the quenching/storage protocol of the skin explants, even if it induces a widening of size pore distribution, does not induce quantitative modifications of the global hydric organization of the samples. The first hydration shell of the proteins is well conserved during the storage procedure.

In Fig. 5, the DSC curves of fresh and frozen skins recorded at 10 °C min⁻¹ between −30 and 85 °C in the 60–85 °C zone are reported. Superimposed on this figure is the DSC curve of a decellularized bovine pericardium in the native state, which is a relatively simple and uniform tissue mainly constituted by type I collagen and representative of the thermal answer of hydrated collagen.

![Fig. 4](image_url)  
**Fig. 4** Hydric organization of fresh and stored skin explants

![Fig. 5](image_url)  
**Fig. 5** Representative DSC curves for a matched set of fresh and stored skin explants and pericardium tissue [first and second heating, scanning rate 10 °C min⁻¹ and enlargement in the (60–85 °C) window]
In a similar way as in bovine pericardium, in skin biopsies an endothermic phenomenon is recorded in this temperature zone during the first heating and associated with the irreversible thermal denaturation of collagen (not observed on successive scans). Small reversible endothermic events can be sometimes detected on the first and second scan and attributed to the reversible phase transition of lipids [61, 62]. Nevertheless, when the hypodermis is correctly removed, these peaks can be neglected and do not interfere with the collagen answer.

In Fig. 5, the main characteristics of the denaturation peak T onset, T max and T mid and the area under the endotherm peak are shown. T mid represents the temperature at which 50% of collagen is denatured. Whole integral area of the heat flow is reported in the inset of Fig. 5, allowing the determination of T mid at the temperature midpoint. T onset, T max and T mid are alternatively taken as the denaturation temperature according to the previous studies [34, 37, 39, 63, 64] while ΔH once normalized to the dry mass gives a measure of the specific heat/enthalpy of denaturation. It is well known that the characteristic feature of type I and type III collagens is the triple helical structure of three left-handed polyproline type helices twisted into a right-handed super helix. The formation of such a structure is due to the repeating sequence Gly-XY, where X and Y are often proline and hydroxyproline, respectively, and hydrogen bonding takes place between chains within the triple helix. On heating, the triple helix unfolds to produce random chains of gelatin that can remain covalently linked to each other or not depending on the degree of heating [65, 66]. The collagen denaturation—distinct from degradation—is a thermally activated process that involves rupture of hydrogen bonds coupling the three α-chains and a rearrangement of the triple helix into a random chain configuration [26]. If collagen denaturation was firstly described like a ‘polymer melting’ [67], detailed calorimetric studies are clearly consistent with the ‘polymer in a box’ mechanism, in which the molecule is thermally stabilized by confinement [27].

Differences in thermodynamic parameters of denaturation for proteins are believed to reflect the differences in their native states [68]. The denaturation temperature and the associated denaturation enthalpy are sensitive to the level of hydration, cross-links within and between molecules and to the amount and character of side chains exposed to the surrounding medium during unfolding [32]. Lowering of the expected denaturation temperature can indicate collagen degradation, fragmentation or expansion of the collagen lattice structure [33, 36, 37], while an increase of this temperature can be due to the formation of heat-stable cross-links, as observed in aged skins [34] or in chemically cross-linked collagenous biomaterials [28]. A change in enthalpy can indicate molecular degradation, denaturation or modification of the heat-labile bonding in the collagen molecules [33, 36, 37]. Since collagen denaturation in skin was shown to be multi-component [34–36], it can be interesting to calculate the mean reciprocal temperature (1/\(T_{\text{mean}}\)) averaged over the whole endotherm and more representative of the thermal stability of collagen as proposed by Miles et al. [36].

\[
\frac{1}{T} = \frac{1}{T_1} + \frac{1}{T_2} - \frac{1}{T_{\text{mid}}} \Delta T
\]

where \(T_1\) and \(T_2\) are the lower and upper temperature limits of the endotherms.

The different thermal parameters of collagen denaturation in skin explants are reported in Fig. 6 for fresh and stored samples. For a better comparison, the reciprocal of (1/\(T_{\text{mean}}\)) was converted into °C and displayed in the same figure. It must be pointed out that \(T_{\text{max}}\) and the reciprocal of (1/\(T_{\text{mean}}\)) are displayed with a very small standard error for the studied samples, what validates the use of DSC to quantify the denaturation phenomenon.

In the case of fresh skins, the \(T_{\text{max}}\) temperature, which is generally ascribed to the denaturation temperature is 71.6 ± 0.5 °C; it is in the same line of order than the thermal parameters of hydrated rich collagenic tissues reported in literature and measured with the same scanning rate such as rat tail tendon [27], pericardium tissue [57] and skins from animals [36]. It was previously shown that the denaturation temperature drastically increases as the hydration decreases [27, 57], due to the decrease of interfibrillar water content compacting the packing of fibers, and to the replacement of protein–protein hydrogen bonds by protein–water hydrogen bonds. This denaturation temperature is independent on hydration for tissues hydrated above 1 kg of water per kg of dry tissue [37] or 30 mol of water per three residues [27] (which corresponds to 1.8 kg of water per kg of dry matter). It must be
pointed out that this condition is checked for the studied samples, with a total hydration of 2 kg per kg of dry matter. In this case, the temperature range of the denaturation phenomenon can be considered as an intrinsic characteristic of collagens in the skin.

\( T_{\text{mid}} \) and \( T_{\text{onset}} \) have also been computed for this study, and in this case also there is no significant difference between the temperature values from fresh and stored explants; \( T_{\text{onset}} \) is of course located in a lower zone than \( T_{\text{max}} \). and error bars of \( T_{\text{onset}} \) are a little wider than \( T_{\text{mid}} \). \( T_{\text{mid}} \) is in the same temperature range as \( T_{\text{max}} \) and the reciprocal of \( (1/T)_{\text{mean}} \).

The reciprocal of \( (1/T)_{\text{mean}} \), which is assumed to be equal to \( T_{\text{max}} \) for narrow and symmetric endotherm peak, is in this case slightly upper than \( T_{\text{max}} \), highlighting the asymmetry of the endotherm in skin samples, as already observed on other collagenic tissues [36]. The standard error is very weak for this parameter. Since it takes into account the proportion of the heat-stable cross-links in the denaturation phenomenon (in contrast with \( T_{\text{max}} \), \( T_{\text{onset}} \) and \( T_{\text{mid}} \)), it can be considered as the preferential biomarker of the thermal stability of collagens in skin for further studies.

The denaturation enthalpy of human fresh skin is evaluated at 0.0128 J kg \(^{-1} \) of dry matter, with a standard error of about 10 %. This value that can be considered rather low when compared to pure collagenic tissues such as tendon or animal skins (\( \Delta H \) comprised between 0.03 and 0.06 J kg \(^{-1} \)); first it must be reminded that some analyses are normalized to the real content of collagen (and not to the dry tissue). Moreover, in the present samples, neither epidermis nor cellular components were removed, decreasing the amount of collagen in the samples. Only few data are available on the denaturation enthalpy of fresh human skins [35, 39], and our values perfectly fit with Wiegand et al. ones, with a denaturation enthalpy of 0.0045 J kg \(^{-1} \) of fresh tissues for intact female skin (which corresponds to 0.013 J kg \(^{-1} \) of dry tissue). Moreover, it was shown that in a similar way as the denaturation temperature, the denaturation enthalpy was widely affected by hydration. At very low hydration, the enthalpy is one-fifth of the enthalpy of the fully hydrated triple helix, which implies that, compared with the fully hydrated state, the triple helix at very low hydrations is held together by one-fifth of its original number of hydrogen bonds. It reaches a constant value above 30 mol of water per three residues [27], and this condition of hydration is checked for the present skin samples. So \( \Delta H \) can be used as a suitable biomarker for the hydrogen bonds network of hydrated collagen in skin.

No significant difference at the 0.05 level were found between the mean values of \( T_{\text{max}} \), \( 1/(1/T)_{\text{mean}} \) and \( \Delta H \) of fresh and stored explants, evidencing that the storage protocol do not impair the thermal stability of collagens in human skin: The hydrogen bonds network, the packing of the fibers and the triple helical domain are completely conserved with a quenching at \(-196 ^\circ\text{C}\) and a consecutive storage at \(-20 ^\circ\text{C}\).

**Vibrational characterization**

FTIR technique in the transmission mode was first devoted to the analysis of lyophilized biological tissues because of the main absorption of water hiding all the other vibration modes; in the ATR mode, native biological tissues can be directly set on the crystal without any other preparation and kept at a constant hydration. This technique is widely applied to investigate in vivo or in vitro the outermost layer of human skin, i.e., the stratum corneum [61, 62, 69–73], but few data are available on the internal layer of human skin, i.e., reticular dermis. The penetration depth of the evanescent wave can be estimated with the refraction index of the crystal and the tissue, the frequency and incident angle of the radiation [73]. The measuring depth of ATR-FTIR in the skin is typically a few microns over the wave number window 4000–650 cm \(^{-1} \) [74].

In Fig. 7, the mean representative FTIR spectra of fresh and stored explants (corresponding to the reticular dermis) in the 2000–950 cm \(^{-1} \) zone are reported; to assist the assignment of the different vibration bands of dermis, the FTIR spectra of hydrated type I collagen (hydrated in the same range as dermis samples, namely containing 65–70 % of water) and purified triglycerides are superimposed on this figure.

Absorption bands in fresh and stored and dermis samples were identified using FTIR data on peptides, protein [75–79] and biological tissues [80–85], peculiarly collagens [63, 80, 86–89] and dermis [17, 90, 91] and probable bands assignments are listed on Table 1.

![Fig 7](image_url)
The classical absorption bands of proteins (amides I, II and III) are found on the spectra of fresh and stored derman, and their positions are very close to the absorption bands of pure type I collagen. In peculiar, collagen absorption features in the fingerprint region (the specific triplet of bands at 1204, 1238 and 1280 cm⁻¹ [81, 92] as well as the specific band at 1338 cm⁻¹) [15, 81] are found in dermis samples, and the correlation coefficient between collagen and dermis spectra is more than 0.95 in this zone.

The very weak absorption in the 1740–1720 cm⁻¹ zone, corresponding to the specific absorption of the carbonyl stretching of ester bond of phospholipids, unsaturated triglycerides and cholesterol esters [93, 94] confirms the clean-up of hypodermis in these dermis samples.

In order to enhance the resolution, the second derivative spectra in the amide I/amide II, which are sensitive to protein secondary structures, were displayed in Fig. 8. The different minima detected in this zone were clearly identified as constant in dermis spectra and not considered as artefacts. The comparison of the second derivative spectra confirms the close correlation between collagen and fresh/stored dermis, with the main minima at of the amide I at 1694 cm⁻¹ (anti-parallel β sheets), 1682 cm⁻¹ (anti-parallel β sheets and β turns), 1667 cm⁻¹ (turns, non H bonded loops), 1659 cm⁻¹ (α helices), 1651 (random helices), 1640–1636 cm⁻¹ (intermolecular water), 1627 cm⁻¹ (intramolecular β-sheets) and 1618 cm⁻¹ (intermolecular β-sheets). According to previous studies on cardiovascular tissues [95], the spectral band at 1659 cm⁻¹ attributed to α-helical structure is particularly representative of the exclusive α-helical structure of collagen [87].

The main minima of the amide II, even if the assignation to peculiar secondary structures is less straightforward in the case of complex tissues [95], are located at 1555, 1543 and 1534 cm⁻¹ in collagen and dermis explants. In a wide class of tissues, the vibration bands at around 1550 cm⁻¹ are generally addressed to β-sheet conformations and the bands in the 1546–1540 cm⁻¹ are attributed to helical conformations [81]; in pure freeze-dried collagen, a band at 1549 cm⁻¹ is associated with triple helix structure cm⁻¹ while a band at lower wavenumber (1530 cm⁻¹) is ascribed to disorder collagen [88]. Side-chain absorptions are well detected at 1510 cm⁻¹ (Phe, Tyr) [88], 1527 cm⁻¹ (CH₂, CH₃) mainly from proteins and lipids.

**Table 1** FTIR bands assignment of fresh and stored skin dermis

<table>
<thead>
<tr>
<th>Band position/cm⁻¹</th>
<th>Assignment</th>
</tr>
</thead>
<tbody>
<tr>
<td>1744</td>
<td>v(C=O) triglycerides, cholesterol esters, phospholipids</td>
</tr>
<tr>
<td>1694–1630</td>
<td>v(C=O) amide I, δ(O–H) water</td>
</tr>
<tr>
<td>1558–1542</td>
<td>v(C–N), δ (N–H) amide II</td>
</tr>
<tr>
<td>1520, 1515, 1507</td>
<td>Side chains (tyrosine, phenylalanine)</td>
</tr>
<tr>
<td>1452</td>
<td>δ(CH₂, CH₃) mainly from proteins and lipids</td>
</tr>
<tr>
<td>1401–1393</td>
<td>v₃(COO–) free amino acids, fatty acid</td>
</tr>
<tr>
<td>1338</td>
<td>CH₂ side-chain rotation (proline) specific band of collagen</td>
</tr>
<tr>
<td>1312</td>
<td>v(C–N), and δ(N–H) amide III</td>
</tr>
<tr>
<td>1282</td>
<td>δ(CH₃) specific of collagen</td>
</tr>
<tr>
<td>1239</td>
<td>Amide III, v₃(PO₃⁻) nucleic acid, phospholipids</td>
</tr>
<tr>
<td>1204</td>
<td>v(C–OH) specific of collagen</td>
</tr>
<tr>
<td>1167–1156</td>
<td>v₃(C=O–C) esters de cholesterol, phospholipids</td>
</tr>
<tr>
<td>1086–1082</td>
<td>v₃(PO₃) nucleic acids, phospholipids, phosphorylated proteins</td>
</tr>
<tr>
<td>1032</td>
<td>v₃(C–O) oligosaccharides, glycolipids</td>
</tr>
<tr>
<td></td>
<td>v(C–O) and v(C–O–C) of the carbohydrate moieties in collagen</td>
</tr>
<tr>
<td></td>
<td>v₃(C=O–OC) polysaccharides, collagen</td>
</tr>
</tbody>
</table>
important difference is the depletion of the band at
secondary structures with the DSC denaturation: the most
or FSD spectra highlights the evolution of the proteins
Fig. 9 Representative mean normalized FTIR spectra in the
spectra in the 2100–950 cm\textsuperscript{-1} zone of fresh and heat-denatured dermis samples.
Inset: Mean representative second derivative spectra and FSD of fresh
and heat-denatured dermis samples in the amide I zone
(Lysine), 1576 cm\textsuperscript{-1} (Asp, Trp) and 1609 cm\textsuperscript{-1} (Gln) [75].
It is noteworthy that the position of the vibration bands in
the global FTIR spectra and in the second derivative
spectra in the 2100–950 cm\textsuperscript{-1} zone are identical in fresh and stored dermis, validating the conservation of the pro-
tein secondary structures with storage and thus validating
this protocol.

To complete the research of biomarkers of collagen
integrity, a selected set of biopsies was denatured by
heating at 10 °C min\textsuperscript{-1} from 20 to 85 °C in hermetic pans
according to previous DSC data. Global spectra of fresh
and denatured samples in the amide I/I zone are reported in Fig. 9. The comparison between FTIR spectra of fresh and DSC-denatured derma lead to two main dif-
ficulties: the first one is the enhancement of the vibration band at 1740 cm\textsuperscript{-1}, attributed to the stretching vibration of
carbonyl group of ester bonds. Such an enhancement has
been already detected in gelatin (denatured collagen)
[87, 88] but no hypothesis was proposed to explain it. We
may assume that it comes from a chemical alteration of
collagen in derma occurring during heating. The second
important discrepancy is the shift of the amide II band
toward low wavenumber in DSC-denatured derma. Such a
behavior has been already reported in previous work on
heat-denatured collagen films [88]. This shift toward low
wavenumber of the amide II band can be explained by the
depletion of the triple helical structure of collagen and the
enhancement of disordered structures [88].

The comparison between the second derivative spectra
or FSD spectra highlights the evolution of the proteins
secondary structures with the DSC denaturation: the most
important difference is the depletion of the band at
1659 cm\textsuperscript{-1}—previously addressed to α-helical structures
[95]—and the occurrence of a new band at 1654 cm\textsuperscript{-1} that
can be ascribed to destabilized helices or disordered
structures. This evolution with denaturation has already
been assumed in literature data for collagen [87, 88]. It
confirms the assignment of the absorption band at
1659 cm\textsuperscript{-1} to the vibrational signature of the native triple
helix of collagen. This absorption band can be considered
as a good biomarker of collagen integrity in dermis, while
the absorption band at 1654 cm\textsuperscript{-1} is ascribed to a marker
of altered collagen. On the other hand, the second deriva-
tive or FSD of amide II region shows less difference
between fresh and denatured skin sample compared to the
amide I region.

Conclusions
Through the combined use of thermal and vibrational
techniques, we can clearly find out that there is no sig-
nificant influence of the storage effect of explants on the
hydric organization, thermal characteristics of the denat-
uration endotherm and secondary structures of proteins in
human skins. Moreover, this works evidences that DSC is
well-suited technique to evaluate the hydric organization
of biological tissue like human skin, giving accurate
results with a high reproducibility. Combined with FTIR,
the use of DSC allows the identification of reproducible
and suitable biomarkers of the main protein in dermis,
namely collagen, from the nanometric to the mesoscopic
scale in biopsies 4 mm in diameter. The denaturation
temperature of collagen 1/(1/T)\textsubscript{mean}, including the asym-
metry of the endotherm, is a reliable biomarker of the
intrinsic thermal stability of collagens in the skin, inde-
dependent on the hydration of fresh skin. Moreover, the
denaturation enthalpy can be considered as another suit-
ble biomarker for the hydrogen bonds network of
hydrated collagen in skin. Finally, the assignment of the
different vibration bands of the amide I and II in human
skin explants has been corroborated thanks to the com-
parison of fresh and DSC-denatured skin explants,
allowing the quantification of the different secondary
structures of proteins and side chains in dermis trough the
decomposition the amide I and II peaks.

After the validation of this DSC/FTIR protocol, the
perspectives will be focused on its application and the
evolution of the evidenced biomarkers with enzymatic
degradation of collagen in dermis, which will be biologi-
cally relevant. Finally, these protocols will be applied in
the scheduled clinical study on young and aging biopsies
(from 30 female 20–30 years old and from 30 female
60–70 years old) for further investigations on intrinsic and
extrinsic aging of dermis.