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Conformational and thermal characterization of a synthetic peptidic fragment inspired from human tropoelastin: Signature of the amyloid fibers

Caractérisation conformationnelle et thermique d’un fragment peptidique synthétique inspiré de la tropolélastine humaine : signature des fibres amyloïdes

J. Dandurand, V. Samouillan, M.H. Lacoste-Ferre, C. Lacabanne, B. Bochicchio, A. Pepe

Physique des Polymères, CIRIMAT UMR 5085, Institut Carnot, Université Paul Sabatier, 118, route de Narbonne, 31062 Toulouse cedex, France

Department of Sciences, University of Basilicata, Via Ateneo Lucano, 10, 85100 Potenza, Italy

Abstract

Objectives. – This work deals with the conformational and thermal characterization of a synthetic peptide (S4) released during the proteolysis of human tropoelastin by the matrix metalloproteinase-12 that was shown to form amyloid-like fibres under certain conditions.

Materials and methods. – S4 peptides were synthesized by solid-phase methodology and aggregated in solution at 80 °C. Fourier transform-infrared spectroscopy (FT–IR) was used to access the secondary structure. Thermal characterization was performed by thermogravimetric analysis (TGA) and differential scanning calorimetry (DSC).

Results. – The DSC study of the soluble linear peptide S4 in solution in TBS reveals the irreversible aggregation into amyloid fibres. FT–IR, DSC and TGA analyses performed on freeze-dried samples evidence differences between the linear peptide and its associated amyloid-like fibres, both on the conformation and the physical structure. When S4 peptides are aggregated, the prominent conformation scanned by FT–IR is the cross β-structure, corresponding to TGA to an increase of the thermal stability. Moreover, the DSC thermograms of S4 fibres are characteristic of a highly ordered structure, in contrast to the DSC thermograms of S4 linear peptides, characteristic of an amorphous structure. Finally, the DSC analysis of differently hydrated S4 fibres brings to the fore the specific thermal answer of the wet interfaces of the cross β-fibres.

Discussion. – FT–IR and thermal techniques are well suited to evidence conformational and structural differences between the soluble peptide and its amyloid form.

Résumé

Objectifs. – Cette étude concerne la caractérisation physique et conformationnelle d’un peptide synthétique (S4) libéré lors de la protéolyse de la tropoélastine humaine par la métalloprotéase-12 et qui peut former des fibres amyloïdes sous certaines conditions.

Matériaux et méthodes. – Les peptides S4 sont synthétisés en phase solide et agrégés en solution à 80 °C. La spectroscopie infrarouge à transformée de Fourier (IR–TF) est utilisée pour accéder à la structure secondaire. La caractérisation thermique est effectuée par analyse thermogravimétrique (ATG) et par analyse calorimétrique diatherme (ACD).

Résultats. – L’analyse par ACD du peptide soluble et linéaire en solution dans le TBS révèle l’agrégation irréversible en fibres amyloïdes. Les analyses par IR–TF, ACD et ATG sur les échantillons lyophilisés...
1. Introduction

Elastin, the protein responsible for elasticity of tissues, such as lung, skin and arterial walls consists of a three-dimensional network whose turnover is almost absent under physiological conditions. However, under pathological conditions, the elastolysis by matrix metalloproteinase-12 (MMP-12) can produce biological active fragments as it was shown in photo-damaged skin [1], arteries of atherosclerotic patients [2] and in lung where the elastotic material contains amyloid-like fibres [3].

Moreover, an ex-vivo study evidenced amyloid-like fibres only reacting with a polyclonal antibody to human elastin within atherosclerosis plaques of cerebral arteries of senile patients [4], suggesting the involvement of elastin in the amyloidogenic process. Interestingly, the extensive investigation by mass spectrometry of the cleavage sites of MMP-12 in human skin elastin [5] showed that polypeptide sequences encoded by exon 30 were present in the elastin lysate. Given these finding, we hypothesize that elastin-derived soluble peptides released from elastin by elastolysis might aggregate into amyloid structures in the vicinity of the degraded elastin fibres, that could explain the “elastotic” material described in numerous reports.

For the first time, some polypeptide sequences encoded from exon 30 (the entire polypeptidic sequence encoded by exon 30) was previously demonstrated to be amyloidogenic [6], mimicking the elastin fragments produced by elastases were synthesized and studied both at the molecular and supramolecular levels by various spectroscopies, MD simulations and AFM microscopy [7]. Among these peptides, the longest one (S4 peptide) was shown to give rise to amyloid fibres.

The aim of this work is to get an insight into the thermal properties of the soluble, linear peptide S4 (labelled S4 peptides) and its aggregated form (labelled S4 fibres) mainly using thermal analysis (TGA and DSC). DSC is a well-suited technique for the analysis of biological materials both in solution [8–10] and in the condensed state [11–14], the insolvibility of some fibres being not an obstacle to the analysis. Combined with TGA, DSC allows to distinguish and to estimate the amount of the different types of water in proteins. Moreover, these techniques reveal ordered and amorphous structures at the mesoscopic level, connecting the molecular level (accessible by vibrational techniques) and the microscopic/macromolecular level (accessible by microscopy and turbidimetry), which were already explored in the previous work on S4 [7]. In Table 1, we summarize the different techniques used in this study focusing on the specific contribution of each of them.

It has long been recognized that amyloid fibrils contain a cross-β spine, with β-strands perpendicular to the fibril axis [15,16]. Atomic structures have been determined by X-ray analysis [17] and by molecular dynamics simulations [18–20] for some of these cross-β spines, revealing a pair of β-sheets mated closely together by intermending side chains in which has been termed a steric zipper [17]. The first level in the aggregation of amyloid structures is the rapid and reversible formation of β-sheets [16]. The slower second step, accompanied by the decrease of entropy, consists in the dehydration and interdigitation of the mating sheet, allowing the formation of the dry amide-stacking hydrogen bonds.

From a medical point of view, thermodynamic and solvation structural properties of amyloidogenic polypeptide sequences could help our understanding on the mechanism of aggregation and the very high insolubility of amyloid fibres also involved in neurodegenerative diseases [21,22]. From a material point of view, the excellent thermal stability and enhanced mechanical properties of amyloid fibres make these structures promising candidates for biomedical development [23], and their structure-function relationship deserves to be clarified.

2. Materials and methods

2.1. Peptide synthesis and purification

The S4 peptides (LIVGAGAGLGGLVGVLGPGVG, molecular weight: 1734 g/mol) were synthesized by solid-phase methodology using an automatic synthesizer Applied Biosystems model 431 A. Fmoc/DCM/HOBt chemistry was used, starting from (0.25 mM) Wang resin (Nova Biochem, Laufelfingen, Switzerland). The Fmoc-amino acids were purchased from Nova Biochem and from Inbios (Pozzuoli, Italy). The cleavage of the peptides from the resin was achieved by using an aqueous mixture of 95% trifluoroacetic acid. The S4 peptide were freeze-dried and purified by reversed-phase high performance liquid chromatography and their purity was assessed by electrospray mass spectrometry.

2.2. Formation of S4 fibres

According to previous turbidimetry experiments showing the self-aggregation of S4 into amyloid-like structures [7], S4 peptides were dissolved in TBS buffer (Tris (50 mM), NaCl (1.5 M), and CaCl2 (1.0 mM), pH 7) with a concentration of 30 mg/mL (17 mM); the solution was stirred at 80 °C for 3 h in order to precipitate S4 peptides into S4 fibres. Once formed, S4 fibres were separated from the solution by 3 cycles of centrifugation (15,000 rpm, 5 min) and washed in pure distilled water prior to the freeze-drying process.

2.3. Fourier transform–infrared spectroscopy (FT-IR)

Fourier transform–infrared spectroscopy/attenuated total reflectance (FT-IR/ATR) spectra were collected using a Nicomp 5700 FTIR (THERMO FISHER SCIENTIFIC, Waltham, MA) equipped in ATR device equipped with a KBr beam splitter and a MCT/R detector. Spectra were recorded over the region of 4000–450 cm−1 with a spectral resolution of 4 cm−1 and 64 accumulations. The ATR accessory used was a Smart Orbit equipped with a type IIA diamond crystal (refractive index 2.4). A single-beam background spectrum was collected from the clean diamond crystal before each experiment and this background was subtracted from the spectra.
Fourier-self-deconvolution (FSD) of the infrared spectra that allows resolution of several overlapping bands [24] was performed in the amide I/II region using Omnic software (THERMO FISHER SCIENTIFIC, Waltham, MA) to extract the peak maxima. The decomposition of the amide I/II bands on the renormalized spectrum was then performed with a Gaussian curve fitting procedure.

2.4. Thermogravimetry analysis (TGA)

Analyses were performed in triplicate on freeze-dried samples (initial weight 5 mg) set in alumina pans between 25 and 650 °C at 10 °C/min under N2 atmosphere using a TGA Q50 (TA INSTRUMENTS, New Castle, DE).

2.5. Water sorption

Initial freeze-dried samples were equilibrated on MgNO3:6H2O saturated solutions at 25 °C (corresponding to a 53% relative humidity). Samples were weighted with a Mettler microbalance after different times of equilibration. After measurements samples were completely dehydrated over P2O5 overnight at 105 °C and weighted again to determine the mass of the dry samples and the mass of water by difference. The hydration level was defined as the mass of water divided by the mass of the hydrated sample.

2.6. Differential scanning calorimetry (DSC)

Analyses were performed using a DSC Pyris calorimeter (PERKIN ELMER, Waltham, MA). The calorimeter was calibrated using cyclohexane and indium as standards, resulting in a temperature accuracy of ± 0.1 °C and an enthalpy accuracy of ± 0.2 J/g. The sample cell was purged with helium and the cooling and heating scan rates were 10 or 20 °C/min depending on the experiments.

An amount of 10 µL of S4 peptide solutions [2 mM in solution in TBS buffer Tris (50 mM), NaCl (1.5 M), and CaCl2 (1.0 mM), pH 7] were sealed in 15 µL aluminium hermetic pans. Experiments, done in triplicate, were performed between 10 and 90 °C in the heating and cooling mode at 10 °C/min.

Freeze-dried S4 peptides and freeze-dried S4 fibres, 5 mg in weight, were sealed into non-hermetic aluminium pans. Experiments, done in triplicate, were performed between 5 and 225 °C with a heating rate of 20 °C/min. Hydrated S4 fibres, 2–10 mg in weight, were sealed into hermetic aluminium pans. Samples were first cooled at −50 °C at 10 °C/min and equilibrated at −50 °C during 5 min. Experiments, done in triplicate, were recorded in the heating mode between −50 and 30−70 °C (depending on the hydration level) with a heating rate of 20 °C/min.

3. Results

3.1. DSC analysis of S4 peptides in solution

During the first heating, the thermogram of S4 peptides in solution (Fig. 1) is characterized by the ΔCp heat capacity step at 37 °C and two sharp exothermic peaks at 61 °C and 78 °C, superimposed on ΔCp heat capacity steps; in contrast, the thermogram recorded during cooling does not present any transition in the 80–60 °C temperature zone (successive heating scans, not shown here, are also devoid of such sharp exothermic transitions), demonstrating the irreversibility of these two transitions. The only event recorded during the cooling run is a well-defined ΔCp heat capacity step at T = 42 °C, reproducible in successive heating/cooling cycles.

3.2. FT–IR spectroscopy of S4 peptides and S4 fibres in the freeze-dried state

The FT–IR spectra of S4 peptides and S4 fibres (Fig. 2) are typical of polypeptides and proteins, exhibiting the amide A (free and H-bonded O–H stretching and N–H stretching), amide I (C=O stretching), amide II (C–N stretching and N–H bending), and amide III (N–H in plane deformation) bands. Some distinct features can be observed on these global spectra, particularly the narrowing and the shift (from 3289 to 3281 cm\(^{-1}\)) of the main band of amide A band in S4 fibres as well as the narrowing and the shift (from 1644 to 1624 cm\(^{-1}\)) of the amide I band.

The amide A band includes the free and bonded absorptions of OH and NH groups [25]. As already observed in α-elasticin, the shift toward lower wave numbers of the amide A band in S4 fibres bands is linked to the increase in the strength of the hydrogen bonds and phase structure [25]. Since the amide I–II region consists of several bands strongly dependent on secondary conformations, it was subjected to curve fitting in order to resolve the various underlying and overlapping spectral features that contribute to this complex region [26–30]. The decomposed FT–IR spectra of the amide I–II region are presented in Fig. 3.

The amide I region (corresponding to the stretching vibrations of the C=O group of the peptide backbone) of S4 peptides contains a main component at 1633 cm\(^{-1}\), ascribed to bonded β turns/pleated β-sheets. This component together with the additional ones observed at 1694 and 1680 are indicative of anti-parallel β-sheet conformations and β turns, respectively [7,27]. Since the bands comprised between 1640 and 1660 are generally associated with unordered conformations and water absorption [7,24,29], the large component detected at 1651 cm\(^{-1}\) is certainly due to an overlapping of these contributions. It is also noteworthy that non-hydrogen bonded C=O groups absorb in the 1667–1661 cm\(^{-1}\) range [7], confirming the presence of PPII conformation in S4 peptide as already revealed in solution by CD [7,29]. The decomposition of the amide II band yields two main components.

![Fig. 1. DSC thermograms of S4 peptides in solution (2 mM) (heating and cooling runs).](image1)

![Fig. 2. FT–IR/ATR spectra of freeze-dried S4 peptides and S4 fibers.](image2)
at 1552 and 1518 cm\(^{-1}\), which are considered representative of the presence of the \(\beta\)-sheet conformation [7]. The band at 1537 cm\(^{-1}\) could be tentatively assigned to the unordered component [31]. We propose for S4 peptides in the freeze-dried state a dominance of \(\beta\)-structures together with unordered and PPII conformations.

The amide I region of S4 fibres is very distinct from the amide I region of S4 peptides, with a prominent component at 1627 cm\(^{-1}\) ascribed to intermolecular \(\beta\)-sheets in cross \(\beta\)-structures [32], together with a band of increasing intensity at 1697 cm\(^{-1}\), which is typical of the anti-parallel \(\beta\)-sheet conformation. The two minor bands at 1651 and 1669 cm\(^{-1}\) can be attributed to unordered conformations/water absorption [7,29] and \(\beta\) turns [26], respectively. The predominance of the cross \(\beta\)-structure in S4 fibres is attested by the presence of two main components in the amide II region at 1552 and 1525 cm\(^{-1}\), that are representative of the \(\beta\)-sheet conformation [7,32].

### 3.3. Thermogravimetric analysis of S4 peptides and S4 fibres in the freeze-dried state

Thermogravimetric (TG) and temperature/derivative (DTG) plots of freeze-dried S4 peptides and freeze-dried S4 fibres are shown in Fig. 4. The mass decrement during the heating process was determined from TG curves and the temperature of the maximum speed of the process \((T_{\text{max}})\) was determined from the maximum of the DTG curves. The global trend of the two TG plots corresponds to the classical thermal behaviour of freeze-dried proteins [33,34]. The first stage (between 25 and 200 °C) is generally connected with the evaporation of water absorbed to the protein, and corresponds to 8.5 and 1.36% of the total mass for S4 peptides and S4 fibres, respectively, meaning that freeze-drying of S4 fibres is more efficient than freeze-drying of S4 peptides.

The second stage occurring between 200 and 500 °C is associated with the degradation of the sample, namely the progressive deamination, decarboxylation and depolymerization arising from the breaking of polypeptide bonds. The degradation of S4 peptides consists of two well-marked steps as shown by two maxima on the DTG plots at \(T_{\text{max 1}} = 235\) and \(T_{\text{max 2}} = 299\) °C, in contrast with the degradation of S4 fibres that occurs in a narrow temperature range with a single maximum at \(T_{\text{max}} = 305.3\) °C.

### 3.4. Water sorption experiments on S4 peptides and S4 fibres in the freeze-dried state

Water sorption experiments at 53% RH were performed at 25 °C (Fig. 5) in order to obtain information on the water/polypeptides affinity first revealed by TGA.

Under these conditions of humidity, we can notice that S4 fibres can adsorb up to 85% of water in contrast to S4 peptides that cannot adsorb more than 19% of water. As revealed by TGA, it is noteworthy that the freeze-drying procedure is more efficient on S4 fibres than on S4 peptides, leading to initial freeze-dried states with 2% and 8% of water, respectively.
3.5. DSC analysis of S4 peptides and S4 fibres in the freeze-dried state

Thermograms of freeze-dried S4 peptides and freeze-dried S4 fibres were recorded in the heating mode between 5 and 120 °C (first run) and between 5 and 225 °C on the successive runs (Fig. 6).

In the case of freeze-dried S4 peptides, the broad endothermic peak observed on the first scan at 63.1 °C may be considered as the first order transition commonly observed in a broad class of hydrated biopolymers. By analogy with previous works [33,35,36], this endothermic event is attributed to the evaporation of bound water, and the value of the enthalpy (ΔH = 135 J/g) is coherent with an amount of bound water comprised between 8 and 10% as measured both by TGA and dehydration over P2O5 at 105 °C. On the second thermogram, the specific thermal answer of wholly dehydrated S4 peptides can be observed. The presence of a glass transition at 147.3 °C (ΔCp = 0.45 J/g/K), reversible on successive scans, underlines the fact that freeze-dried S4 peptides possess an amorphous region, with a lack of long-range order, like elastin [37,38]. As already observed by TGA, the degradation of S4 peptides begins above 210 °C.

In the case of freeze-dried S4 fibres, the endothermic event associated with the evaporation of water is weaker (ΔH = 27.9 J/g) than the event reported for S4 peptides. The ratio between enthalpies is roughly the same as the ratio between the total amounts of bound water measured by TGA and dehydration over P2O5 at 105 °C.

The main event detected on the second thermogram is the sharp endothermic peak at 145.6 °C associated to an order–disorder transition, which can be considered as to the thermal signature of amyloid fibres. The sharpness of this first order transition is indicative of a highly cooperative process. This transition, not observed on the third scan, is irreversible.

3.6. DSC analysis of S4 fibres in the hydrated state

Freeze-dried S4 fibres 5 mg in weight were equilibrated over MnO6.6 H2O saturated solutions at 25 °C (relative humidity 53%) until the water uptake reached a maximum. The total hydration level of the samples was evaluated by gravimetry, and was equal to 84.3% under these conditions of humidity. Different hydration levels were obtained equilibrating the fully hydrated samples over KOH saturated solutions at 25 °C (relative humidity 8%) during different times, allowing a gradual dehydration of the samples. Complete dehydration was obtained by a heating at 60 °C during 10 min. The thermograms of the differently hydrated samples recorded in the heating mode are shown in Fig. 7. For the sake of clarity, only one cooling thermogram (corresponding to the sample containing 84.3% of water) has been superimposed in this figure.

For high hydration levels, a part of water absorbed in the sample is able to crystallize as shown by the exothermic events at –14 °C and –39 °C on the first cooling thermogram of the Fig. 7. On the successive heating thermogram, the melting of this crystallized water gives rise to endothermic phenomena between –30 and 0 °C. This melting peak consists of two contributions for high hydration levels, suggesting the presence of two kinds of ice in this case. For lower hydration levels (70%, 58% and 29.8%), only the low temperature peak is observed; in previous studies on water insoluble biological polymers, this peak was attributed to freezing bound water [39], corresponding to loosely bound water adsorbed on polymer having a certain kind of crystalline structure which may be the same as the structure of natural ice [40]. The second important event detectable on the thermograms of differently hydrated S4 fibres is the endothermic peak occurring between 15 and 35 °C, for hydration levels comprised between 84.3 and 2.1%, vanishing for completely dehydrated samples, and indicative of the transition from an ordered to an unordered structure.

4. Discussion

4.1. Thermal behaviour of S4 peptides in solution – irreversible aggregation

In the case of S4 peptides, the first heating thermogram reported in Fig. 1 gets an insight into the possible aggregation mechanism into S4 amyloid fibres evidenced in a previous work by turbidimetry and biochemical tests [7]. Firstly, the positive ΔCp contribution at 37 °C is certainly associated with the progressive destabilization of the polyproline II (PPI) conformation from 0 to 60 °C as it was evidenced by circular dichroism [7], shifting the equilibrium between PPI, unordered structures and β turns towards β turns preferential conformation. It must be pointed out that PPI is a flexible structure devoid of intramolecular hydrogen bonds that can easily interchange with other conformations [7].

Comparing the special feature of the sharp exothermic peaks occurring at 61 and 78 °C with the DSC-monitored aggregation of insulin [41], β-lactoglobulin [42] and β2-microglobulin [43] into amyloid fibres, we propose to ascribe them to the formation of β-sheets and cross β-structures, leading to the irreversible
aggregation of the S4 peptides into S4 fibres. This multi-step, irreversible aggregation mainly driven by hydrophobic interactions involves both formation of hydrogen bonds into β-sheet structures [41,42], dipole–dipole interactions into the steric zipper and breakdown of hydrogen bonds (rearrangement of water around the peptide, expulsion of water at the β-sheet–β-sheet interface of the steric zipper); the total area of the exothermic peaks is estimated at ≈6 J/g (where the mass corresponds to the mass of dry S4 peptides); this net negative enthalpy is certainly due to a total increase of the global hydrogen bonds network density and the positive contribution of Van der Waals interactions at the dry interface of the steric zipper [16–18]. This assumption is well supported by turbidimetry experiments, suggesting for S4 peptides an irreversible inverse-temperature phase transition above 60 °C. The reorganization of water molecules around the β-sheets structure could explain the associated variations of the heat capacity, since the heat capacity change originates primarily from the hydrogen-bonding properties of water near the protein surface [44,45].

4.2. Conformation of S4 peptides and S4 fibres – relation with thermal stability

Our FT–IR results in good agreement with the structural data on amyloid fibrils; while S4 polypeptides adopt a wide distribution of conformations (PPII, unordered conformations, β-sheets and β turns), S4 fibres consist mainly of intermolecular β-sheets in cross β-structures, with an increasing strength of the involved hydrogen bonds.

The multiplicity of the thermal degradation evidenced by TGA on S4 peptides can be correlated with the multiplicity of the secondary structures revealed by FT-IR analysis, the unordered structure being less stable than the highly ordered, high hydrogen bonded β-sheets. Moreover, it is noteworthy that even the degradation of the more stable component in S4 peptides occurs at lower temperature (Tα ≈ 299 °C) than in S4 fibres. This thermal stabilization of β-sheets in the S4 fibres can be related to their organization into cross β-spines with steric zippers, with a densification of hydrogen bonds.

4.3. Physical structure of S4 peptides and S4 fibres in the freeze-dried state

The amorphous character of S4 peptides is evidenced by DSC with the presence of a glass transition without any endothermic peak that would be indicative of an ordered structure. This lack of long-range order is due to the coexistence of unordered conformations, labile β turns and PPII conformations as evidenced by FT–IR analysis in the dry state. It is noteworthy that the glass transition temperature of S4 peptides is close to the glass transition temperature of the polypeptide encoded by exon 10 of human tropoelastin (Tg = 144.2 °C, Mw = 2100.5 g/mol) [10], and lower than recombinant freeze-dried human tropoelastin (Tg = 190 °C) [26] or freeze-dried elastin (Tg = 200 °C) [30].

In agreement with vibrational and thermogravimetric results, the thermal transitions of S4 fibres are very distinct from the S4 peptides ones. The sharp endothermic peak at 145.6 °C, indicative of the cooperative melting of a long-range ordered structure can be addressed to the thermal signature of amyloid fibres, forming highly ordered cross β-structures with intra- and intermolecular β-sheets that extend throughout the entire length of the fibre. Such an endothermic event has already been observed by DSC in the hydrated state for the fibrils of the N47A mutant of the R-spectrin SH3 [13]. The temperature of this peak is indicative of an important stabilization of the ordered β-sheets structure when S4 forms fibres. It must be pointed out that the enthalpy of this endothermic event (ΔH = 4 J/g) is in the same range of order than freeze-dried collagen type I [33,46], which can be also considered as a “crystalline” biopolymer possessing a very long-range order.

4.4. Physical structure of S4 fibres in the presence of water

The first important point concerns the difference of solubility of S4 peptides and S4 fibers: if S4 peptides are soluble in water like a wide class of linear polypeptides, once formed S4 fibres are insoluble and cannot be studied in solution. That is the reason why S4 fibres were analyzed in this work at different degrees of hydration.

The very distinct water affinity of S4 peptides and S4 fibres in the freeze-dried state revealed by water sorption experiments is quite striking at first glance because of the same chemical composition of both samples. From a polymer point of view, it can appear paradoxical since S4 peptides possess unordered conformations in contrast to S4 fibres that are quasi-wholly organized in ordered cross β-structures. Interestingly, a large amount of water can be trapped in this highly ordered structure, as already observed for other systems, such as hydrogels [40], multi-lamellar vesicles [47] and silk-keratin composites [48] constituted of numerous interstices, bringing to the fore different states of water (freezing free water, freezing bound water and non-freezing bound water).

In the case of S4 fibres, the trapping of a large quantity of water must be connected to the definition of the wet interfaces between β-sheets of the outside faces of the pair of sheets revealed by structural analysis of amyloid fibres [16]. As a matter of fact, there are two distinctly different interfaces between β-sheets of the cross β-spine: a dry, dewetted interface between the paired sheets engaged into the steric zipper, in contrast to the highly hydrated external faces exposed to solvent [16].

The content of crystallized water into differently hydrated S4 fibres was estimated from the heating thermograms (Fig. 8) as the ratio of the melting enthalpy ΔH of the hydrated sample and the melting enthalpy of bulk water [49] (333.55 J/g) and reported in Fig. 8. By difference, the uncrystallized water amount was also evaluated and reported in Fig. 8. In contrast to the amount of crystallized water that increases with the hydration level, the content of uncrystallized water reaches a maximum for hydration levels comprised between 30 and 70%. In this range of hydration, approximatively 15% of water molecules are tightly bound to S4 fibres. We have also reported in Fig. 8 the enthalpy (ΔH fibres)

![Fig. 8. Amounts of crystallized and uncrystallized water in S4 hydrated fibers and enthalpy of the second endothermic peak as a function of the total hydration of S4 fibers computed from Fig. 7.](image-url)
normalized to the mass of dry fibres, corresponding to the second endothermic event detected between 15 and 35 °C on the thermograms of Fig. 7, associated to an ordered structure. The enthalpy of this event sharply increases between 0 and 5% of hydration, meaning that uncrystallized/bound water is essential for the stabilization of this structure. Interestingly, the enthalpy is maximum when the amount of uncrystallized water is maximum, confirming this assumption. It is clearly shown in Fig. 8 that an excess of crystallized water destabilizes this ordered structure, resulting also in a decrease of bound water. Water, in this case possesses a peculiar behaviour, very distinct from what observed for globular proteins [49], where the amount of uncrystallized water increases with increasing water content, reaching a maximum for high hydration contents. This endothermic event can be considered as the thermal answer of the wet interface of the cross β-fibres, mainly due to the disruption of hydrogen bonds between water and polypeptide chains when the temperature increases.

5. Conclusion

In contrast to tropoelastin that coevacates to give rise to native fibrillar structures, with an equilibrium at the conformational level between unorganized conformations, labile PPII and β turns, β-strands and α-helices, essential for its biological function, the synthetic derived-elastin peptide S4 mimicking the released fragment from elastolysis by MMP-12 self-assembles into β-structure-rich amyloid-like fibres [7], which shows a greater degree of conformational restriction than the native elastin structure. In this work, we attempted to get an insight into the physical structure and water affinity of this peptide in its linear state and its aggregate state, since few studies in the literature have reported thermodynamics of amyloid fibre formation and thermal characterization of amyloid fibres.

Firstly, it was shown that the formation of irreversible aggregates when temperature is raised up in S4 polypeptide solution could be monitored by DSC, corroborating the previous turbidimetry assays. This multi-step mechanism is associated with a net negative variation of enthalpy, due to a net positive increase of H-bonds and Van der Waals interactions. Further experiments performed at different concentrations and different scan rates will be necessary to quantitatively estimate the balances of changes in enthalpy and entropy. Moreover, it will be essential to evaluate the influence of the medium (in peculiar the role of calcium) on this aggregation phenomenon in order to correlate it with pathologic conditions.

In the condensed state, the thermal signature of S4 precipitated fibres is completely distinct from the one of the linear polypeptides; the former is highly stable and ordered, while the latter is less stable, without any long-range order. These radically distinct features at the ultrastructural level are well correlated with the major changes detected at the conformational level by FT–IR analysis.

Finally, DSC is well suited to evidence the peculiar thermal signature of the wet interfaces of the outside faces of the pair of sheets structure of amyloid fibres first proposed by Nelson et al. [16] on structural analysis of elongated microcrystals.

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