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## Determination of orientations of aromatic groups in self-assembled peptide fibrils by polarised Raman spectroscopy

José C. Rodríguez-Pérez, † Ian W. Hamley and Adam M. Squires\*

In this paper we describe a novel combination of Raman spectroscopy, isotope editing and X-ray scattering as a powerful approach to give detailed structural information on aromatic side chains in peptide fibrils. The orientation of the tyrosine residues in fibrils of the peptide YTIAALLSPYS with respect to the fibril axis has been determined from a combination of polarised Raman spectroscopy and X-ray diffraction measurements. The Raman intensity of selected tyrosine bands collected at different polarisation geometries is related to the values and orientation of the Raman tensor for those specific vibrations. Using published Raman tensor values we solved the relevant expressions for both of the two tyrosine residues present in this peptide. Ring deuteration in one of the two tyrosine side chains allowed for the calculation to be performed individually for both, by virtue of the isotopic shift that eliminates band overlapping. Sample disorder was taken into account by obtaining the distribution of orientations of the samples from X-ray diffraction experiments. The results provide previously unavailable details about the molecular conformation of this peptide, and demonstrate the value of this approach for the study of amyloid fibrils.

### 1 Introduction

"Amyloid" fibrils form from self-assembly of a range of different proteins and peptides. Much of the research in this area has been driven by their association with diseases such as Alzheimer's, Parkinson's, and prion disorders;<sup>1,2</sup> in addition, fibrils may be formed *in vitro* by self-assembly of other proteins and designed peptides, leading to research into their potential use as nanomaterials.<sup>3,4</sup> X-ray fiber diffraction has demonstrated a common core "cross- $\beta$ " structure for many of these fibrils,<sup>5</sup> where strands are arranged perpendicular to the fibril axis, with the beta-sheet structure stabilised by hydrogen-bonding between adjacent strands.

It is very difficult to obtain more detailed structural information on polypeptide conformation within different fibrils, because in general they are not amenable to analysis by X-ray crystallography.<sup>6</sup> Recently there have been developments in this area using solidstate NMR (ssNMR),<sup>7,8</sup> which is capable of giving inter-atomic distances and dihedral angle constraints for molecular dynamics simulations; this has emerged as the main method to give high resolution 3D structures of peptide fibrils.

In this paper, we use a different approach based on polarised Raman spectroscopy to give detailed quantitative information on the orientation of aromatic residues within fibrils. This is particularly important, as it has been suggested that interactions between aromatic residues play an important role in amyloid fibril formation.<sup>9–12</sup> We envisage this approach as providing complementary data to ssNMR, and moreover, it has two key advantages: firstly, our approach uses much smaller quantities of peptide or protein (0.1 mg as compared with 1-10 mg for ssNMR); and secondly, in general it does not require the use of isotope editing, unless the peptide contains more than one copy of a particular aromatic residue. Isotope labeling is a barrier to the analysis of fibrils from larger proteins, where site-specific labelling usually cannot be done. In this paper, the peptide we chose has two tyrosine residues, and we use deuteration to analyse them independently; however, had we picked another example with a single tyrosine, no labelling would have been required-a significant advantage over ssNMR, which typically requires deuteration, <sup>13</sup>C, or <sup>15</sup>N labelling for structural analysis of peptide fibrils.<sup>8</sup>

Linear dichroism spectroscopy is a technique performed on aligned samples to give information on the orientation of

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Department of Chemistry, University of Reading, Reading, RG6 6AD, UK. E-mail: a.m.squires@reading.ac.uk

<sup>†</sup> Present address: NERC Space Geodesy Facility, East Sussex, BN27 1RN, UK.

specific groups relative to the axis of sample alignment by comparing spectra obtained with light polarised parallel and perpendicular to this axis.<sup>13</sup> Very few published examples exist of the application to amyloid fibrils of vibrational linear dichroism using polarised infrared<sup>6,14–16</sup> or Raman<sup>17</sup> spectroscopy, and these studies have only given broad, qualitative information on whether a particular part of the peptide is disordered, or aligned parallel or perpendicular to the fibril axis. Similar qualitative information has been obtained from linear dichroism using polarised UV/Visible spectroscopy.<sup>11,13,18–20</sup>

However, much more precise quantitative data on specific angles of orientation may in principle be obtained from polarised vibrational spectroscopy. In the case of Raman analysis, this requires knowledge of the shape of the Raman tensor for a particular normal mode of vibration, together with quantitative information on the degree of orientational sample disorder. Although such quantitative methods have not been applied to peptide fibrils before, pioneering work has been carried out by Tsuboi and co-workers on oriented samples of filamentous virus to determine the orientation of amino-acid side chains and DNA bases.<sup>21-24</sup> However, they had no direct information on the orientational distribution of virus filaments within their samples. For liquid samples oriented by shear flow, they therefore extrapolated the data to higher shear rates to estimate the polarised Raman intensity ratio expected from perfect alignment.<sup>25</sup> For solid samples, where this was not possible, they simply estimated the maximum error that this disorder would cause in the calculated orientation, and concluded that in their case it was small compared to other experimental uncertainties,<sup>26</sup> making the implicit assumption of perfect order. In contrast, in this paper we demonstrate that orientational disorder may be obtained directly using X-ray scattering data, allowing us to determine a meaningful geometry of aromatic residues relative to the fibril axis, rather than an average with respect to the macroscopic sample axis.

Deuteration of amino acid side chains is a useful strategy that has been employed to aid band assignment and structural interpretation of Raman spectra,<sup>22,27–31</sup> and represents a powerful advantage of polarised vibrational spectroscopy (infrared and Raman) over UV/Visible linear dichroism. The peptide studied in this article contains two tyrosine residues, and tyrosine ring deuteration on one of these enabled us to perform the calculations independently for each residue, since the labelled amino acid causes distinct peaks in the Raman spectra.

The peptide we chose to study in this article corresponds to residues 105–115 of the protein transthyretin, with sequence YTIAALLSPYS. Several naturally occurring mutations of the protein transthyretin are known to cause pathologies related to the formation of amyloid fibril precipitates.<sup>32</sup> A number of amyloidogenic peptides based on its sequence have been studied to obtain clues about their structure and mechanism of fibril formation.<sup>33–36</sup> The peptide YTIAALLSPYS has been the subject of numerous studies.<sup>35,37,38</sup> The backbone structure of the YTIAALLSPYS strands within the aggregated fibrils was determined by magic-angle spinning solid-state NMR experiments.<sup>39,40</sup> Fluorescence resonance energy transfer spectroscopy (FRET) was used to obtain details about the arrangement of the strands

forming the fibrils as well as its aggregation process.<sup>41</sup> The environment of the tyrosine residues of YTIAALLSPYS in solution was examined by UV resonance Raman, giving insights about the structure adopted by this peptide in the pre-aggregation stage.<sup>42</sup> Interstrand carbonyl distances were determined from magic angle spinning (MAS) NMR.<sup>43</sup> In previous ssNMR studies, the conformation of the tyrosine side chains in YTIAALLSPYS was not determined. Thus, in solving the molecular structure of the monomer, statistical data base restraints were used instead of experimental ones.<sup>40</sup> We present a polarised Raman spectroscopy study on this system, with the objective of determining the orientation of these residues within the fibril.

#### 1.1 Materials and methods

**1.1.1 YTIAALLSPYS.** YTIAALLSPYS peptide, corresponding to the sequence of the protein transthyretin from residues 105 to 115, was obtained from CS Bio (CA, USA) CS1652. The same peptide, labelled at position 1 with deuterated tyrosine (L-tyrosine-2,3,5,6- $d_4$ ), was obtained from CK Gas Products (Hampshire, UK), at 98% purity. We denote this latter sample as Y<sub>1</sub> $d_4$ -TIAALLSPYS. Amyloid fibril sample preparation was based on protocols reported in the literature.<sup>39</sup> The peptides were dissolved in purified water with 10% (v/v) acetonitrile, to a final concentration of 10 mg mL<sup>-1</sup>. The resultant solutions were incubated at 37 °C for 24 hours followed by incubation at room temperature for several weeks. After incubation, both samples formed a thick gel indicative of extensive fibril formation. Liquid nitrogen was used to freeze-thaw the samples and break the network of fibrils in order to facilitate alignment.

**1.1.2** X-ray diffraction. Dried stalk samples of labelled  $Y_1d_4^-$  TIAALLSPYS were prepared by leaving ~10 µL of 10 mg mL<sup>-1</sup> incubated peptide solution between the ends of two wax-coated tube capillaries, a standard procedure to induce alignment in fibrillar samples.<sup>44</sup> Samples were mounted vertically onto the four axis goniometer of a RAXIS IV++ X-ray diffractometer (Rigaku), equipped with a rotating anode generator and a Saturn 992 CCD camera (Biocentre, University of Reading). Two images were collected at different positions across the length of the samples to account for possible local variations in alignment.

1.1.3 Polarised Raman spectroscopy. Polarised Raman spectra of both YTIAALLSPYS and  $Y_1d_4$ -TIAALLSPYS were collected with a Renishaw inVia Reflex Raman microscope (Old Town, Wotton-Under-Edge, Gloucestershire, UK). The excitation radiation was provided by a 785 nm diode laser (300 mW power at source). The polarisation of the incident light was adjusted using a polariser filter, and that of the scattered radiation by using a second polariser with or without a half-wave plate. The laser was focused onto the surface of the samples with a  $20 \times$  objective (NA = 0.40, spot size in the diffraction limited case 2.4 µm), which also collected the Raman scattered radiation (180° backscattering geometry). The detector used was a Peltier-cooled CCD array. For the isotopically labelled peptide, data were obtained from four different positions along the stalk and the small differences encountered in relative peak intensities used for the error analysis. Spectra of the unlabelled sample were collected at a

single point, and used as a reference for the Raman bands shifted upon deuteration of the labelled tyrosine residue. The intensity response to different polarisations of the instrument was corrected by measuring the spectra of carbon tetrachloride.<sup>45</sup> The bands at 314 and 220 cm<sup>-1</sup>, with a known depolarisation ratio of 0.75, were used to estimate a correction factor that was applied to all the spectra.

By changing the polarisation of the incident and scattered light we collected two different spectra for each sample,  $I_{cc}$  and  $I_{bb}$ . The subscripts make reference to the direction of polarisation of the incident light and Raman scattering, where *c* is the direction of the sample/stalk axis and *b* perpendicular to it. No sample rotation was needed to collect the data, allowing us to keep the laser focused on the same position of the stalks when changing collection geometries. Exposure times were typically between 40 and 80 seconds at 10% laser power (30 mW at source). No fluorescence or sample heating artifacts were observed when operating at these conditions.

1.1.4 Data analysis. Leadbetter's expression relating the scattering intensity from a distribution of rod-shaped objects and their distribution of orientations was employed to analyse the X-ray diffraction data.<sup>46</sup> The distribution of orientation was calculated solving numerically the analytical solutions to Leadbetter's equation presented by Deutsch.<sup>47</sup> d-Spacings and azimuthal intensity integrations were carried out with the software Adxv.<sup>48</sup> The calculation of the orientation distribution function of the fibrils from X-ray data was performed with in-house programs written in Octave49 to solve the relevant integrals. Raman tensor rotations were performed with the computational algebra package Maxima.<sup>50</sup> The calculation of the average values of the Raman intensity from the experimental measurements, weighting by the orientation distribution function and  $I_{cc}/I_{bb}$  intensity ratio contour finding was done in Octave. Curve fitting of the C-D stretching bands was performed with the software fityk,<sup>51</sup> and rotation of the PDB structures with the package Avogadro.52

#### 2 Theoretical section

The intensity of a Raman band is dictated by the extent of polarisability change with the normal coordinate of the vibration. This is given by a rank two symmetric tensor unique for each vibration, the Raman tensor. The polarised Raman scattering intensity is given by:

$$I_{uv} = I_0 |u^{\mathrm{T}} \alpha_{abc} v|^2 \tag{1}$$

where  $\alpha_{abc}$  is the Raman tensor in the laboratory frame, u and v are unit vectors that indicate the direction of the electric field of the incident and scattered radiation,  $u^{T}$  is the transpose of u, and  $I_0$  is a constant.

By experimental determination or as a result of a calculation, the Raman tensor shape and orientation of its principal axes for specific vibrations can be obtained. However, to use eqn (1), the appropriate transformations of the Raman tensor from the molecular frame to the laboratory coordinate system have to be performed.<sup>53</sup> Tensors in one coordinate system are expressed in another reference frame using the rotation matrix R whose elements are the direction cosines relating both coordinate systems:<sup>54</sup>

$$\alpha_{abc} = \mathbf{R}^1 \alpha_{x'y'z'} \mathbf{R} \tag{2}$$

where  $\mathbf{R}^{\mathrm{T}}$  is the transpose matrix of  $\mathbf{R}$ .

We define three systems of axes to describe our experimental layout: the laboratory space-fixed axes, the fibril axes, and the molecular axes, as depicted in Fig. 1. These coordinate systems are related to each other by the Euler angles, which we defined using the same convention as Wilson, Decius and Cross.<sup>55</sup> The laboratory and sample axes are *abc*, with the laser beam passing in the *a* direction; both incident and scattered light may be polarised in the b or c direction. The samples are aligned along the axis c, containing fibrils which are uniaxially aligned about the stalk axis. Each individual fibril makes a variable angle  $\beta$  with the stalk axis (coincident with c). We assume that the system has cylindrical symmetry about this axis, so the fibrils have no preferred orientation in the plane *ab*. The principal axes of the Raman tensors of the tyrosine side chains are x'y'z', where the aromatic ring lies in the x'y' plane, and the x' axis is parallel to the phenyl C–O bond, as shown in Fig. 1. The x'y'z' axes (molecular frame) are defined with respect to the fibril axes XYZ by the angles  $\theta$  and  $\chi$ . We also assume cylindrical symmetry within each fibril about the fibril axis Z, so the tyrosine side-chains have no preferred orientation in the XY plane. Because of the axial symmetry of the fibrils it is not possible to obtain information about the third Eulerian



**Fig. 1** Euler angles relating the three coordinate systems that describe our setup. The axes *abc* define the laboratory frame. The stalk samples are aligned in the direction of the *c* axis. Individual fibrils within the stalk are distributed with cylindrical symmetry about *c*, making an angle  $\beta$  with it. The angles  $\theta$  and  $\chi$  define the orientation of the principal axes of the Raman tensor (x'y'z') with respect to the fibril axes (*XYZ*). Figure adapted from ref. 22.

angle that would be required to fully characterise the orientation of the x'y'z' axes.

If a perfect or very high degree of molecular alignment is obtained, as is the case with single crystals, it would be possible to relate the Raman tensor in the space fixed and molecular frames directly, assuming that the fibril and laboratory axes are coincident. In our case, the Raman tensor is first transformed from the molecular frame to the fibril axes, and afterwards to the laboratory frame. The expression obtained depends on the angle  $\beta$ , which reflects the imperfect alignment of the fibrils about the axis *c*. This expression is afterwards integrated for all the possible orientations about the fibril axis *Z* and the laboratory axis *c* because of the axial symmetry of the stalks and the fibrils. Each of the angular rotations needed to perform the full transformation is expressed by simple rotation matrices, and their product gives the direction cosine matrix **R** needed for eqn (2).<sup>54</sup>

The fibrils within the stalk samples will not be oriented at a single-valued angle  $\beta$  from the director; instead, they will present a distribution of orientations that has to be taken into account to relate the Raman intensities and the molecular orientations. We performed wide-angle X-ray scattering (WAXS) measurements to obtain the orientation distribution function of the fibrils within our samples, and explicitly included it in the calculations. With the orientation distribution of the fibrils, and making use of eqn (1) and (2), the polarised Raman intensity is obtained from the following expression:

$$I_{uv} = \frac{I_0}{4\pi^2} \int_0^{\pi/2} \int_0^{2\pi} \int_0^{2\pi} f(\beta) \left[ u^{\mathrm{T}} (\boldsymbol{R}^{\mathrm{T}} \alpha_{x'y'z'} \boldsymbol{R})^2 v \right] \sin \beta \, \mathrm{d}\omega \, \mathrm{d}\sigma \, \mathrm{d}\beta$$
(3)

where  $f(\beta)$  represents the orientation distribution function (ODF) of the fibrils about the sample axis. The integration steps with respect to  $\omega$  and  $\sigma$  correspond to averaging about, respectively, the fibril and the laboratory axes. After integration, eqn (3) depends on the components of the Raman tensor in the molecular frame,  $r_1$  and  $r_2$ , and on the angular variables  $\theta$  and  $\chi$ . The intensities from two different polarisation geometries,  $I_{cc}$ (exciting and scattered light polarised parallel to axis *c*) and  $I_{bb}$ (exciting and scattered light polarised perpendicular to *c*) can be divided to remove the constants:

$$I_{cc}/I_{bb} = F(r_1, r_2, \theta, \chi).$$
 (4)

As explained elsewhere,<sup>21</sup> if tensors for more than one Raman band for the same moiety are available, a graphical solution for the angular variables can be obtained from the experimental intensity ratios of those bands. The intersection point of the contour lines in ( $\chi$ ,  $\theta$ ) space that results from solving eqn (4) gives the angles that satisfy all the experimental intensities.

### 3 Results

#### 3.1 X-ray diffraction

The X-ray diffraction pattern of  $Y_1d_4$ -TIAALLSPYS, shown in Fig. 2, presents the same characteristics observed previously for

YTIAALLSPYS peptide fibrils.3 Meridional reflections at 4.7 Å arise from the repeating pattern of hydrogen bonded strands within the fibrils. The strands run perpendicular to the long fibril axis, forming sheets that extend alongside it. Several sheets stack together to form a fibril, at a regular spacing that gives rise to the equatorial reflection at 8.8 Å. These two reflections constitute the so called cross-ß pattern, a structural characteristic shared among fibrils formed from different amyloidogenic proteins and peptides.<sup>5</sup> We assume that the polypeptide strands that form the fibrils are perpendicular to the long fibril axis, and that therefore the order of the strands within the sample is equivalent to the order of the fibrils themselves, *i.e.* we assume that the distribution of orientations of the strands accurately represents the distribution of orientations of the fibrils. With this assumption, we used the meridional 4.7 Å arc to calculate the orientational distribution function of the fibrils.

The azimuthal intensity profile of the 4.7 Å reflections was background subtracted with the average of the intensity profiles immediately adjacent to it,<sup>56</sup> and afterwards used as our input data. The calculated orientation distribution function is shown in Fig. 3. Also shown is the predicted intensity from the calculated ODF obtained from Leadbetter's expression.<sup>46</sup> The predicted intensity fits the experimental diffraction profile accurately, indicating the adequacy of the numerical routines employed here and, more importantly, the robustness of the



**Fig. 2** Wide angle X-ray diffraction pattern of  $Y_1d_4$ -TIAALLSPYS. The stalk sample is aligned in the vertical direction. The intense meridional reflections correspond to the regular spacing between adjacent strands in the fibril axis.



**Fig. 3** Experimental and predicted intensity (left) and calculated ODF (right) for the 4.7 Å reflection of the wide angle X-ray diffraction pattern of a  $Y_1d_4$ -YTIAALLSPYS stalk sample.

approach of calculating  $f(\beta)$  directly from the analytical solution. In terms of the order parameters usually quoted to describe sample orientation, the values obtained indicate a high degree of uniaxial alignment ( $P_2 = 0.66$ ,  $P_4 = 0.3$  and  $P_6 = 0.07$  average values). The average tilt angle calculated from the distribution is 29°, and integration of  $f(\beta)$  shows that 50% of the scatterers are within 0° <  $\beta$  < 29° from the director axis *c*, and that 90% are within 0° <  $\beta$  < 45°.

#### 3.2 Raman spectroscopy

**3.2.1 Amide I.** The polarised Raman spectra of YTIAALL-SPYS and  $Y_1d_4$ -TIAALLSPYS are displayed in Fig. 4. The amide I band appears at the characteristic frequency for peptides in an extended conformation (~ 1667 cm<sup>-1</sup>).<sup>57</sup> This peak is the most intense signal in the  $I_{cc}$  spectra, as observed in previous work on aligned amyloid fibrils using polarised Raman spectroscopy.<sup>17</sup> As expected, both samples show a very high degree of orientation, as judged from the great intensity ratio  $(I_{cc}/I_{bb})^{1667}$  of this peak (7.3 for  $Y_1d_4$ -TIAALLSPYS). The amide I band originates mainly from a carbonyl stretching vibration, whose maximum polarisability oscillation is oriented at a small angle from the C–O bond direction in the amide bond plane.<sup>53,58</sup> Therefore, in a cross- $\beta$  structure this band is expected to be much more



**Fig. 4** Polarised Raman spectra of peptides YTIAALLSPYS (top) and  $Y_1d_{4^-}$ TIAALLSPYS (bottom). Black traces:  $l_{cc}$  spectra. Grey traces:  $l_{bb}$  spectra. The spectra are offset vertically for convenience. For visualisation purposes, both unlabelled YTIAALLSPYS spectra have been scaled up to approximately match the intensity of their labelled counterparts, taking the 1445 cm<sup>-1</sup> peak in the spectrum of Y<sub>1</sub>d<sub>4</sub>-TIAALLSPYS  $l_{cc}$  as a reference.

intense when both the incident and scattered light are polarised parallel to the fibril axis.

In principle, an estimation of the orientation of the fibrils could be obtained from the intensity ratio from this band, applying the expressions derived by Bower.<sup>65</sup> Although this method has been employed extensively for a number of systems,<sup>66–69</sup> certain assumptions on the symmetry of the Raman tensor are needed in order to solve the equations involved. Moreover, the information polarised Raman scattering experiments can give about  $f(\beta)$ is limited to the first two moments of its Legendre expansion  $(P_2 \text{ and } P_4)$ .<sup>70</sup> We have tested the error introduced by this latter limitation by performing our calculations with the truncated orientation distribution resulting from using  $P_2$  and  $P_4$ from our X-ray data, and also using the maximum entropy distribution function<sup>70</sup> from these two moments. Our conclusion is that the final results are only slightly affected by this approximation (results not shown). However, the assumptions about the amide I Raman tensor are not easily justifiable, since experimental studies on two different systems concluded that this tensor is not cylindrically symmetric.53,58 Indeed, in a recent polarised Raman study on the orientation of silk it was found that assuming an effective cylindrical amide I Raman tensor leads to substantial errors in the value of  $P_4$ , as opposed to the value obtained when using the (non cylindrical) local tensor from other systems.<sup>71</sup> Also, the disparities observed in the orientation distribution functions calculated from X-ray diffraction and polarised Raman spectroscopy support the conclusion that transferring the amide I effective tensor from other systems is only accurate for the first order parameter,  $P_2$ .<sup>72</sup>

**3.2.2** Tyrosine bands. The most prominent bands originating from the tyrosine residues are located at 1613, 1207, 1171, 851, 827 and 641 cm<sup>-1</sup> (see assignments in Table 1). Comparing the spectra of YTIAALLSPYS with that of  $Y_1d_4$ -TIAALLSPYS, we observe that the bands at 1613, 1171, 827 and 641 cm<sup>-1</sup> in the isotopically labelled sample have a lower intensity than in the unlabelled one. The decreased intensity indicates that the labelled tyrosine residues do not contribute to these bands due to a significant red-shift in the frequency of the corresponding vibrations.

Raman tensors for several vibrations of L-tyrosine have been determined by Tsuboi and co-workers from polarised Raman spectroscopy measurements on single crystals of this amino acid.<sup>60</sup> In principle, the polarised intensity ratio from all these bands could be used to determine the orientation of the tyrosine side chains in our samples. In practice, some of these tensors may not be transferable from the single crystal amino acid samples to the amyloid fibrils, since the chemical environment in both systems is likely to be different. The effect on the polarisability tensor that these differences may cause is difficult to predict. Thus, we solved our equations making use of all the available tensors and assessed the orientation information obtained on the basis of the agreement amongst the different results.

**3.2.3** Deuterated tyrosine bands. Red-shifted peaks arising from the deuterated tyrosine residue are easily identifiable in the spectra, as is the case with the bands at 1590, 1573, 790 and 622 cm<sup>-1</sup>, which are absent in the spectra of YTIAALLSPYS.

$\nu (\mathrm{cm}^{-1})$	Assignment	Ref.	
2289	$Y_1d_4$ symmetric C–D st.		
2274	$Y_1d_4$ symmetric C–D st.	22	
1667	Amide I (β-sheets)	59	
1613	Y C <sub>2</sub> –C <sub>3</sub> , C <sub>6</sub> –C <sub>5</sub> in-phase st.,	60, 61	
	shifts to $\sim 1590 \text{ cm}^{-1}$ on deuteration		
1595	Y C–C st.	60, 61	
1572	$Y_1d_4$ C–C st.	60	
1445	L, A, I, V C–H <sub>3</sub> b.	31	
1345	S, L, I C–H <sub>2</sub> tw./r., A C <sup><math>\alpha</math></sup> H b., C <sup><math>\alpha</math></sup> C st.	31	
1230	Amide III	59	
1207	Y $C^{\beta}-C^{\gamma}$ st.	60, 62	
1171	Y C–H ring b., shifts to 849 $\text{cm}^{-1}$	60	
	on ring deuteration		
1151	I, V CC st.	31	
1130	L, V, I C–C st.	31	
1090	$CH_3$ r., $C^{\alpha}-C^{\beta}$ st.	59	
1056	S $C^{\alpha}$ - $C^{\gamma}$ CH <sub>2</sub> st.	63	
1030	Y	30	
1008	Y ring b.	59	
900	P ring br.	64	
851	Y Fermi resonance	24, 59, 6	
	(br. mode and CCC out-of-plane b.)		
827	Y ring br. shifts to $\sim$ 795 cm <sup>-1</sup>	24, 59, 6	
	on ring deuteration		
790	$Y_1 d_4$ br.	60	
641	Y ring def. shifts to $\sim 620 \text{ cm}^{-1}$	60	
	on ring deuteration		
622	$Y_1 d_4$ def.	60	

At higher wavenumbers (~2275 cm<sup>-1</sup>), the C–D stretching vibrations of the labelled tyrosine residue appear in a spectral region free from overlapping with other bands (Fig. 5). The positions of these three peaks, at 2289, 2274 and 2256 cm<sup>-1</sup> are almost the same as those observed in L-tyrosine-2,3,5,6- $d_4$  single crystal,<sup>22</sup> although, of course, their relative intensities are different since the orientations are different in our sample.



**Fig. 5** Polarised Raman spectra of Y<sub>1</sub>d<sub>4</sub>-YTIAALLSPYS in the spectral region 1500–2800 cm<sup>-1</sup>. Black trace:  $I_{cc}$ . Grey trace:  $I_{bb}$ . The spectra are offset vertically for convenience. The C–D stretch vibrations appear at ~ 2275 cm<sup>-1</sup>, with a higher intensity in the  $I_{bb}$  spectrum. Insets: C–D stretching bands at both polarisations and sub-band decomposition and fitting with Lorentzian curves.

In the fibrils, these bands are slightly overlapping due to the more heterogeneous environment of amyloid fibril systems with respect to single crystals. The two stronger bands, at 2290 and 2272 cm<sup>-1</sup>, have been assigned to the symmetric stretching modes.<sup>22</sup> The weak peak at ~ 2250 cm<sup>-1</sup>, assigned to an antisymmetric stretching, is only visible as a small shoulder, more prominent in the  $I_{bb}$  spectrum.

Other peaks assigned to deuterated tyrosine, in particular those at 790 and  $622 \text{ cm}^{-1}$ , were not employed to perform the orientation calculation, because the Raman tensor is defined as the first derivative of the polarisability with respect to the normal coordinate, and therefore its specific value depends on the atomic displacements involved in each normal mode. Since atomic displacements depend on the nuclear masses of the atoms involved, tensors from non labelled molecules are generally non transferable to isotopically substituted ones.<sup>73</sup> We therefore did not consider it valid to assume that published tensors from vibrations in non-deuterated tyrosine could also be used for their deuterated analogues.

3.2.4 Intensity measurements. Intensities for the bands of the non labelled tyrosine residue at 1613, 1171, 827 and 641 cm<sup>-1</sup> were taken as the absolute heights at the peak centres after a linear baseline was subtracted. Area measurements would be problematic except for the vibration at  $641 \text{ cm}^{-1}$ , which is the only band isolated from other contributions. For the bands corresponding to the C-D stretching vibrations we employed a curve fitting procedure to resolve the contributions from the sub-bands present in that spectral region. The heights of the Lorentzian curves that fitted the data at 2290 and 2273 cm<sup>-1</sup> were taken as the intensities for these vibrations. Although these two bands plus that at 2250  $\text{cm}^{-1}$  are the most intense in this region, some extra functions were added to account for a curved baseline and other minor contributions such as the gaseous N<sub>2</sub> peak at  $\sim 2330$  cm<sup>-1</sup>. Both the peak heights and areas of the fitted functions gave very similar values for the ratio  $I_{cc}/I_{bb}$ . An example of the curve fitting results is shown in Fig. 5. The average intensity ratios at each collection geometry for all the tyrosine peaks used for the orientation calculation are detailed in Table 2.

#### 3.3 Orientation calculation

All the Raman tensors used in the present work have been determined by Tsuboi and co-workers from polarised Raman experiments on single crystals of tyrosine.<sup>22,60</sup> We make the assumption that these tensors are transferable to our fibril structures, and will assess the results obtained on the basis of consistency. The precise shape and orientation of these tensors in the molecular frame are shown in Fig. 6. Because of the high

**Table 2** Experimental intensity ratios  $I_{cc}/I_{bb}$  and standard errors for tyrosineresidues Y1 (deuterated) and Y10 in Y1d4-TIAALLSPYS fibrils

	Y10				$Y1(d_4)$	
Freq. $(cm^{-1})$	1613	1171	827	640	2289	2274
$I_{cc}/I_{bb}$	0.46	1.52	0.89	0.83	0.53	0.55
$\sigma_{ m mean}$	0.02	0.08	0.04	0.03	0.06	0.03



**Fig. 6** Raman tensors for the vibrational bands used in this work. These tensors have been determined by Tsuboi and co-workers from polarised Raman measurements on single crystals of L-tyrosine and L-tyrosine-2,3,5,6- $d_A$ .<sup>22,60</sup> The local coordinate axes are indicated for each tensor, with *x* and *y* in the plane of the paper and *z* perpendicular to it. All the tensors present the same orientation for the molecular system of axes, except in the case of the band at 642 cm<sup>-1</sup>, which is rotated by 45° in the plane of the phenolic ring. Adapted from Tsuboi *et al.*<sup>22,60</sup>

symmetry of the phenolic ring most of the tensors share the same orientation of their principal axes. This system of axes is defined as follows: the axis *x* is parallel to the line defined by atoms  $C_1-C_4$  of the tyrosine residue and the axis *y* is parallel to the line defined by atoms  $C_2-C_6$  and intersects the axis *x* at the centre of the aromatic ring (apart from the vibration at 642 cm<sup>-1</sup>, where *x* and *y* are rotated by 45° in the plane of the phenolic ring); the axis *z* is perpendicular to both *x* and *y* (see Fig. 6). These correspond to *x'*, *y'* and *z'*, respectively, in Fig. 1.

Expressions for the polarised intensities  $I_{cc}$  and  $I_{bb}$  were obtained by rotating the Raman tensor from the molecular to the laboratory frame, followed by an integration over the angular variables  $\omega$  and  $\sigma$ . The orientation distribution function shown in Fig. 3 was then used to weight the Raman intensities to account for sample disorder. The ratio  $I_{cc}/I_{bb}$ , needed to remove the constants present in the equations, represents the height of a surface in ( $\chi$ ,  $\theta$ ) space. The contours given by the ratio of the experimental intensities for each band were plotted; the intersection points from bands corresponding to each tyrosine residue represent the angular coordinates that satisfy all the experimental conditions.

**3.3.1** Tyrosine 1 ( $Y_1d_4$ ). As we have already argued, it is not valid to assume that Raman tensors from isotopically labelled samples are transferabe to non-labelled ones. Hence, there are only two different tensors that can be used to obtain the orientational information for this residue: those corresponding to the vibrations at 2290 and 2273 cm<sup>-1</sup>. The rest of the published tyrosine tensors belong to vibrations of the non deuterated tyrosine ring, and the other bands in our data that are unique to deuterated tyrosine do not have published Raman tensor values. In any case, a minimum of two vibrations sought. In an ideal situation, it is preferable where possible to use three or more bands to determine orientation, as we have with Tyr-10, to

uncover and quantify inconsistencies due to slight variation of polarizability tensors between crystal and fibril environments. Where two bands are used, as with Tyr-1, it is possible that a solution can be obtained even if the tensors are distorted. When more data become available on Raman tensors from additional vibrations in deuterated tyrosine, further validation will be possible for Tyr-1. At any rate, these vibrations are highly localised and therefore their Raman tensors are particularly well suited to transfer between different chemical species,<sup>22</sup> *i.e.* their values are less likely to be affected by changes in the chemical environment.

The plots in  $(\chi, \theta)$  space for  $Y_1(d_4)$  are shown in Fig. 7 for the range  $0^{\circ} < \chi < 180^{\circ}$  and  $0^{\circ} < \theta < 90^{\circ}$ . Because eqn (1) contains powers of trigonometric terms there is no unique solution for the angular variables. For these two bands, a total of four Eulerian coordinates satisfy the experimental data: ( $\chi_1$  =  $180^{\circ} - \theta_1$ ). Uncertainties in the intensity measurements of the Raman bands directly translate into uncertainties in the angular coordinates for each residue. We estimated these uncertainties by plotting the contours corresponding to the average values for the intensity ratios  $I_{cc}/I_{bb}$  plus and minus one standard deviation of the mean. The possible angular values determined within experimental uncertainty lie in the area limited by these contour lines. In numerical terms, for the first solution we find ( $\chi_1 = 26^\circ \pm 3$ ,  $\theta_1 = 61^\circ \pm 10$ ). The uncertainties for the other three solutions are identical to the first one.

**3.3.2 Tyrosine 10 (Y10).** For the unlabelled tyrosine at position 10 of  $Y_1d_4$ -TIAALLSPYS there are four Raman bands that can be used to determine its orientation. If we assume that the tensors for these vibrations are transferable from the single crystal system, the parametric contours generated from all these bands should intersect at a common point, giving the



**Fig. 7** Contour plots in  $(\chi, \theta)$  space for tyrosine residue Y1. The lines are the locus of points that satisfy the two experimental intensity ratios obtained. Solid black line: contour of the 2290 cm<sup>-1</sup> band; dashed line: contour of the 2273 cm<sup>-1</sup> band. The uncertainty arising from the intensity measurements is represented by the grey lines, corresponding to the average results plus/less one standard deviation of the mean. The coordinates of the two interception points displayed here are  $(\chi_1 = 26^\circ, \theta_1 = 61^\circ)$  and  $(180^\circ - \chi_1, \theta_1)$ .

orientation of the molecular system of axes of the tyrosine side rings. However, we found that there are no solutions that satisfy the intensity ratios for all the Raman tensors. Instead, there are three sets of points where three of the four traces intersect. The contour plots for these Raman bands are shown in Fig. 8 for the region  $0 < \chi < 180^{\circ}$  and  $0 < \theta < 90^{\circ}$ . The first possible set of solutions, labelled S<sub>1</sub> in the figure, are the two intersection points of the tensors corresponding to the vibrations at 1179, 827 and 640 cm<sup>-1</sup>, at ( $\chi_1 = 89^{\circ}$ ,  $\theta_1 = 55^{\circ}$ ) and ( $\chi_1$ ,  $180^{\circ} - \theta_1$ ). The second set of solutions, S<sub>2</sub>, originate from the intersection of the 1179, 1614 and 640 cm<sup>-1</sup> traces, at ( $\chi_2 = 113^{\circ}$ ,  $\theta_2 = 62^{\circ}$ ) and ( $\chi_2$ ,  $180^{\circ} - \theta_2$ ).

As discussed above, the principal axes of the Raman tensor for the vibration at 640  $\text{cm}^{-1}$  are rotated 45° in the plane of the tyrosine ring with respect to the principal axes for the other bands. In order to obtain the contour lines in common  $(\chi, \theta)$ space it was necessarily to increment the values of  $\chi$  by 45° for this trace.<sup>21</sup> Since the tyrosine ring can itself be rotated about the line defined by atoms  $C^{\delta}$  and  $C^{\zeta}$ , *i.e.* turned upside down, a further possible solution arises from this tensor, obtained from subtracting  $45^{\circ}$  to the calculated contour line. This additional solution, S<sub>3</sub>, is symmetric about  $\chi = 90^{\circ}$  with respect to  $(\chi_2, \theta_2)$ and thus have the angular coordinates ( $\chi_3 = 67^\circ$ ,  $\theta_3 = 62^\circ$ ) and  $(\chi_3, 180^\circ - \theta_3)$ . These three sets of solutions arise from the intersection of three different traces, making their individual uncertainties smaller than the solutions obtained for the labelled residue. By plotting the results obtained for the mean intensity ratio plus and less one standard deviation of the mean we conclude that the uncertainties for each solution do not exceed  $2^{\circ}$  in both  $\gamma$  and  $\theta$ .

The existence of three different sets of points for Tyr-10 (and their symmetric counterparts at  $180^{\circ} - \theta$ ) may indicate that at



**Fig. 8** Contour plots in  $(\chi, \theta)$  space for tyrosine residue Y10. Solid black line: contour of the 1179 cm<sup>-1</sup> band; dotted line: contour of the 640 cm<sup>-1</sup> band; dashed line: contour of the 827 cm<sup>-1</sup> band; dotted-dashed line: contour of the 1614 cm<sup>-1</sup> band. The symmetric 640 cm<sup>-1</sup> trace which generates solution S<sub>3</sub> is not shown for clarity (see main text). Grey traces represent the uncertainty in the intensity measurements for each band. The coordinates of the three solutions found in this range are ( $\chi_1 = 89^\circ$ ,  $\theta_1 = 55^\circ$ ), ( $\chi_2 = 113^\circ$ ,  $\theta_2 = 62^\circ$ ) and ( $\chi_3 = 67^\circ$ ,  $\theta_3 = 62^\circ$ ).

least one of the tensors employed can not be transferred from the crystal structure to the chemical environment of our system. Unfortunately, we can not make a decision about which set of solutions is the correct one with the data available. In any case, the range of possible  $\theta$  values is greatly restricted, lying between  $55^{\circ} \pm 2$  and  $62^{\circ} \pm 2$ , and its symmetric counterpart region at  $180^{\circ} - \theta$ , *i.e.* between  $118^{\circ} \pm 2$  and  $125^{\circ} \pm 2$ . For the angle  $\chi$  the range is broader, centred at ~90° from  $67^{\circ} \pm 2$  to  $113^{\circ} \pm 2$ .

#### 4 Discussion

We have obtained several possible values for the orientation of the two tyrosine residues in YTIAALLSPYS amyloid fibrils. These sets of solutions are related but not entirely equivalent. However, from Raman spectroscopy measurements alone it is not possible to distinguish between them. Moreover, the axial symmetry of the samples precludes the determination of the angle  $\omega$  (see Fig. 1), impeding the full characterisation of the orientation of the tyrosine rings. However, these data offer new constraints that can be used to determine the position of the tyrosine residues in this molecule, since the possible orientations they can adopt are restricted by these results.

#### 4.1 Comparison with ssNMR data

Previous ssNMR studies on YTIAALLSPYS were performed without experimental restraints for the tyrosine side chains.<sup>40</sup> Instead, statistical database-derived data were used to limit the torsion angles of these amino acids to the values most commonly observed in proteins. A total of 20 structures were deposited in the protein data bank by Jaroniec and co-workers, representing the lowest energy solutions obtained from simulated annealing molecular dynamics calculations. For comparison purposes, we have calculated the angles  $\chi$  and  $\theta$  for the two tyrosine residues in the published structures of YTIAALLSPYS (PDB ID code 1RVS).<sup>40</sup> The coordinates of the five central residues of the strand were used to define the system of axes, considering the fibril axis to be parallel to the average orientation of the carbonyl bonds.

The angles  $\chi$  and  $\theta$  for the two tyrosine residues for each one of these structures are plotted in Fig. 9. It is clear from the figure that the possible coordinates adopted by the tyrosine rings in the fibril structure of YTIAALLSPYS do not randomly span the entirety of the solution space. Furthermore, the clusters of most frequently appearing coordinates are different for Tyr-1 and Tyr-10. The  $(\chi, \theta)$  coordinates for Tyr-1 are mainly concentrated in a region of low  $\chi$  values (<40°) and  $\theta$  angles between approximately  $40^{\circ}$  and  $60^{\circ}$ . On the other hand, the angular coordinates for Tyr-10 are more spread. Still, almost half of them appear in a region with  $\chi$  values centered around 90° and  $\theta$  angles between 132° and 160°. Compared to the polarised Raman data, the most frequent coordinates for Tyr-1 are in very good agreement with one of the four possible solutions consistent with our experiments. Models with tyrosine angular coordinates far off the ones we determined are in discrepancy with these data and are thus unlikely candidate structures.

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**Fig. 9** Angular  $\chi$  and  $\theta$  coordinates of the two tyrosine rings for the 20 minimum energy structures obtained by Jaroniec *et al.*<sup>40</sup> from ssNMR measurements and subsequent molecular dynamics simulations. Black dots: Tyr-1. Grey dots: Tyr-10. Dark grey circles and light grey circles: coordinates for Tyr-1 and Tyr-10, respectively, obtained in the present work from polarised Raman measurements.

In the case of Tyr-10, despite the multiplicity of the solutions obtained by spectroscopic means, these require the phenolic ring of this residue to be oriented in one of two restricted regions in ( $\chi$ ,  $\theta$ ) space. This is the case with the most frequent solutions obtained by Jaroniec *et al.*, although the  $\theta$  values for these points are 10° to 40° higher than our own. The absence of points in certain regions of the ( $\chi$ ,  $\theta$ ) plot indicate that some solutions obtained from Raman spectroscopy, although numerically consistent with the data, are not feasible given the restrictions imposed by the rest of the molecule on the orientation of the phenolic rings.

#### 4.2 Proposed coordinates from Raman spectroscopy

As discussed above, it is not possible to generate a unique solution for the orientation of the tyrosine residues from the Raman data alone. Besides, our data provide angular orientations rather than explicit cartesian coordinates, which depend on the position of the rest of the molecule. However, it is feasible to generate tyrosine coordinates consistent with our data from a starting structure. Based on the orientations obtained from the ssNMR structures, we performed rotations about the bonds  $C^{\alpha}-C^{\beta}$  and  $C^{\beta}-C^{\gamma}$  in Tyr-1 and Tyr-10, keeping the rest of the bond angles and atomic distances constant. Given the overall good agreement between the Raman data and the ssNMR structures, only minor rotations were necessary to generate coordinates compatible with our results.

As an example, we generated the model with angular coordinates ( $\chi = 26^\circ$ ,  $\theta = 61^\circ$ ) for Tyr-1, and ( $\chi = 89^\circ$ ,  $\theta = 125^\circ$ ) for Tyr-10, which is the closest to the most frequent solutions obtained by Jaroniec and co-workers. In Fig. 10, we show the resultant model for YTIAALLSPYS amyloid fibrils obtained taking the first ssNMR structure as starting point, overlayed against the rest of unmodified structures. It has to be stressed that the



**Fig. 10** Ensemble of ssNMR structures of YTIAALLSPYS amyloid fibril strands from Jaroniec *et al.*<sup>40</sup> The highlighted model corresponds to the least energy structure from ssNMR with the tyrosine rings oriented according to the polarised Raman data obtained in this work.

**Table 3** Cartesian coordinates for the residues Tyr-1 and Tyr-10 in the monomer of YTIAALLSPYS amyloid fibrils. Obtained from the first ssNMR structure rotating about bonds  $C^{\alpha}-C^{\beta}$  and  $C^{\beta}-C^{\gamma}$  to satisfy the Raman data

	Tyrosine 1				Tyrosine 10			
	id	x	у	z	id	x	у	z
С	4	-15.298	-2.727	1.126	143	14.070	0.486	0.391
$\mathbf{C}$	5	-16.112	-2.798	2.399	144	12.925	1.408	0.749
$\mathbf{C}$	6	-15.627	-2.263	3.585	145	12.765	2.630	0.107
$\mathbf{C}$	7	-17.366	-3.399	2.415	146	12.006	1.058	1.731
$\mathbf{C}$	8	-16.365	-2.326	4.751	147	11.722	3.477	0.433
$\mathbf{C}$	9	-18.111	-3.465	3.577	148	10.960	1.898	2.062
$\mathbf{C}$	10	-17.607	-2.928	4.742	149	10.822	3.106	1.410
0	11	-18.343	-2.993	5.902	150	9.782	3.948	1.736
н	13	-15.367	-3.686	0.636	153	14.076	-0.325	1.104
Н	14	-14.269	-2.542	1.400	154	14.988	1.046	0.483
Н	15	-14.655	-1.792	3.589	155	13.471	2.917	-0.658
Н	16	-17.758	-3.820	1.499	156	12.116	0.111	2.239
Н	17	-15.969	-1.905	5.663	157	11.614	4.422	-0.078
Н	18	-19.082	-3.936	3.569	158	10.257	1.608	2.828
Н	19	-19.261	-2.787	5.712	159	9.083	3.447	2.165

id: residue index within the whole structure according to the PDB file. Only the coordinates of the atoms from the tyrosine residues with modified positions are shown, except atoms 4 and 143, which are unchanged with respect to the original PDB structure.

model shown in the figure does not represent the unique coordinates that the tyrosine residues can adopt to be in agreement with our data. It is, however, the one that required the minimum amount of rotations from this particular starting ssNMR geometry. At any rate, this is to be understood as an example of how our experiments can reduce the number of potential solutions by using these new angular restraints. New molecular dynamics calculations would be required to refine the coordinates of the whole structure taking into account these data.

For reference purposes, we include in Table 3 the new cartesian coordinates for the atoms in the tyrosine residues obtained from the rotations performed. The coordinates of Tyr-1 are the result of rotating  $-5^{\circ}$  about the  $C^{\alpha}-C^{\beta}$  bond and  $15.4^{\circ}$  about the  $C^{\beta}-C^{\gamma}$  bond. The coordinates of Tyr-10 were obtained by rotating 21.2° about the  $C^{\alpha}-C^{\beta}$  bond and  $-0.9^{\circ}$  about the  $C^{\beta}-C^{\gamma}$  bond.

#### 5 Conclusions

Complete structural determination of amyloid fibril systems is still a difficult task, which has only been achieved in a limited number of cases. A great array of techniques have been used to gain structural insights from amyloids, including a variety of spectroscopies, X-ray diffraction and X-ray crystallography. In the present work we have demonstrated, for the first time, the use of polarised Raman spectroscopy in combination with site-specific isotope labelling to resolve the orientation of two residues in an amyloid system. These data can be used as new restraints for molecular dynamics simulations and fine tune the molecular model of YTIAALLSPYS fibril structure.

In this paper, we have used these new information to validate and refine a pre-existing model of YTIAALLSPYS fibril structure. The amino acid residues we studied are of particular importance, since it is unclear what precise role aromatic side chains play in the process of aggregation.<sup>9,10</sup> Accurate knowledge of the aromatic rings' orientation should prove helpful to elucidate open questions such as the contribution towards overall fibril stability from these residues.

We envisage a number of ways that the technique can be employed. On its own, the polarised Raman data will typically have more than one unique orientational solution, as in the example here. However, it is likely that only one of these is energetically plausible, so computational methods can resolve this issue. Nonetheless, it is unlikely that a full peptide structure can be obtained using polarised Raman as the only experimental method; instead, it can provide previously unavailable experimental constraints, complementing other data obtained using methods such as solid state NMR and/or 2D infrared spectroscopy.

Some aspects of this technique that make it attractive include the limited amounts of material required (100  $\mu$ g peptide) as compared with ssNMR and a very simple sample preparation. Since sample disorder is accounted for by using X-ray diffraction data, no exceptional high levels of fibril alignment should be required. The use of isotopic labelling is not necessary in the case of peptides with a single instance of the residue of interest, or where other strategies such as sequence mutation are employed. This is in contrast with solid-state NMR, and also with the 2D-IR methods employed by Zanni and others.<sup>74–76</sup> This latter approach employs <sup>13</sup>C<sup>18</sup>O labels to investigate local peptide backbone conformation, complementary to the data obtained here, on orientation of aromatic groups relative to the fibril axis.

It must be noted that these methods are not limited to the characterisation of tyrosine side chains. As long as Raman tensors for the amino acids under study are known and the relevant vibrations produce spectral bands of enough quality, the procedures we employed can be applied to other systems. Raman tensors for several moieties are already available in the literature,<sup>77</sup> and where no experimental data exist computer calculations could provide a reasonable approximation. As such data become available, from experiments and theoretical calculations, and for a greater number of biologically important molecular fragments, we envisage our approach becoming an increasingly powerful tool for structural analysis of fibrillar biomaterials.

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