Spatially resolved spectroscopic differentiation of hydrophilic and hydrophobic domains on individual insulin amyloid fibrils

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The formation of insoluble β-sheet-rich protein structures known as amyloid fibrils is associated with numerous neurodegenerative diseases, such as Alzheimer’s and Parkinson’s disease. A detailed understanding of the molecular structure of the fibril surface is of interest as the first contact with the physiological environment in vivo and plays a decisive role in biological activity and associated toxicity. Recent studies reveal that the inherent sensitivity and specificity of tip-enhanced Raman scattering (TERS) renders this technique a compelling method for fibril surface analysis at the single-particle level. Here, the reproducibility of TERS is demonstrated, indicating its relevance for detecting molecular variations. Consequently, individual fibrils are systematically investigated at nanometer spatial resolution. Spectral parameters were obtained by band-fitting, particularly focusing on the identification of the secondary structure via the amide III band and the differentiation of hydrophobic and hydrophilic domains on the surface. In addition multivariate data analysis, specifically the N-FINDR procedure, was employed to generate structure-specific maps. The ability of TERS to localize specific structural domains on fibril surfaces shows promise to the development of new fibril dissection strategies and can be generally applied to any (bio)chemical surface when structural variations at the nanometer level are of interest.

Protein misfolding often results in the formation of β-sheet-rich aggregates, known as amyloid fibrils. Many unsolved structural questions of great interest in biochemistry are associated with these protein aggregates. The development of efficient drugs for the treatment or prevention of amyloid fibril-related diseases such as Alzheimer’s or Parkinson’s would certainly benefit from a detailed surface characterization of fibrillar species. Various microscopic techniques, such as scanning or transmission electron microscopy (SEM and TEM) as well as atomic force microscopy (AFM), are commonly used to study protein aggregates. Although these approaches can supply morphological information at the nanometer scale, little chemical information for the analyzed sample can be obtained. Vibrational circular dichroism (VCD), solid-state NMR, IR and Raman spectroscopies have recently become valuable tools for fibril characterization but are limited by the requirement for a large sample volume and high protein concentration (~10 mM). This way, averaged information of the bulk volume of a sample is obtained but individual fibrils cannot be measured.

Amyloid fibrils typically exhibit high morphological and structural heterogeneity, a phenomenon known as fibril polymorphism. Insulin forms tape-like fibrils at pH below 2 and twisted aggregates at pH above this point. Despite significant tertiary and secondary structural rearrangements during fibrillation of insulin the disulfide bridges remain intact and are not cleaved. Using NMR coupled with hydrogen-deuterium exchange and deep UV resonance Raman spectroscopy, it was possible to demonstrate that insulin did not decompose. Moreover, the

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amino acid residues, that were not accessible to the solvent and consequently had to be located in the fibril core, could be determined. Thus, all other amino acid residues are then located on the fibril surface and are accessible to the solvent. It was observed that one half of these amino acids was hydrophilic and the other hydrophobic. However, no information about the spatial distribution of hydrophilic and hydrophobic domains on the surface of the individual insulin fibrils could be obtained.

Coupling Raman spectroscopy with scanning probe microscopy provides an access to both topography and molecular structure characterization in a single experiment with (sub-)nanometer resolution.

Therefore, this approach known as tip-enhanced Raman scattering (TERS) becomes extremely valuable for structural characterization of individual amyloid fibrils. TERS is a highly sensitive and specific tool in this regard and is most valuable for studies at the single-molecule level. So far it has been assumed that the spatial resolution of TERS depends on size, shape, metal of the nanoparticle at the tip apex. If the size of the metallic tip would predominantly determine the lateral resolution - a (sub-) nanometer resolution would be out of reach. Recent ultra-high vacuum low temperature TERS results on tetraphenylporphyrins have convincingly demonstrated a spatial resolution of 0.5 nm that cannot be explained by the classical concepts. Several recent quantum mechanical approaches also revealed that the high spatial resolution does not solely correlate with the tip apex diameter. Important are direct “chemical” interactions between the front-most tip features and the sample and also the particular atomistic composition of the metal nanoparticle. Both effects confine the local response area drastically.

Several TERS studies on insulin fibrils, collagen fibrils and other amyloid fibrils have been published so far and particularly for insulin it was demonstrated that the protein secondary structure of fibril surfaces clearly differs from the core: conventional and UV resonance Raman spectroscopy show that the latter consists of a pure β-sheet, whereas surface-sensitive TERS reveals that the former is also composed of α-helix and unordered protein secondary structures as well as their mixtures. In this context we conservatively assume that the surface area extends to a maximum of 2–3 nm because of the quickly decaying fields.

In our previous TERS studies of different insulin fibril types, spectra were collected for arbitrary areas to gain statistically averaged information regarding the secondary structure and amino acid abundance on fibril surfaces.

The present work pursues a systematic characterization of pre-defined areas on insulin fibrils grown at pH 1.5. These fibrils feature an almost flat topology and are presumed to grow by side-by-side with the association of intertwined protofilaments. Initially we demonstrate the reproducibility and stability of the AFM-based TERS setup as being fundamental for analysis at nanometer resolution. In all ensuing experiments, TERS spectra were continuously recorded on consecutive equidistant points along the fibril main axes. For a detailed data evaluation, spectra are analyzed using a band-fitting procedure, such that even amino acids with non-characteristic bands can be identified, enabling the distinction of mainly hydrophilic and hydrophobic domains. A two-dimensional amino acid distribution is visualized by generating maps from the fitted data. Finally, we show that data assessment by multivariate data analysis (N-FINDR) yields similar results.

Results and Discussion

Prior to the actual TERS experiments discussed we specifically addressed the aspect of the instrumental reliability. Details on this first experiment can be found in Supplementary Figs S1 and S2. The repeated detection of transitions from one spectral pattern to the other indicates the reliability of the experiment where tip contamination and thermal drift could be excluded.

In the following sections it will be demonstrated that several amino acids can be directly identified in the TERS spectra with one or two marker bands, assuming that no other bands interfere (e.g. tyr, asn/gln). Other peptide features like cys-cys are known to show several bands due the presence of conformers. In this case up to six bands can be detected, simultaneously. The classification of amino acids with several overlapping modes requires a correlation of additional bands (e.g. lys, his, gly) to ensure the assignment. The band assignment in Table 1 agrees with that in ref. 23 and the literature cited therein and is further supported by additional references where necessary. Overlap of bands is a particular issue when it comes to the assignment of the secondary structure from amide III bands. Since amide III bands are sensitive to the conformation assignment of modes between 1230–1300 cm$^{-1}$ to amide III should be possible by correlating them with amide I bands. Since the absence of amide I bands in plasmon-enhanced spectroscopy is common and known such a correlation can be difficult.

Utilizing the amide III band region alone for a conformation assignment is challenging because other modes can interfere and must be explicitly excluded.

Systematic surface characterization of insulin fibrils. In a second TERS experiment, information about structural organization was obtained along a 24.5-nm line on a fibril surface. In this case, 50 spectra along the fibril main axis (step-size: 0.5 nm) were collected, as indicated by the gray line in the AFM topography image in Fig. 1a.

The constant SNR of consecutively recorded raw spectra shown in Fig. 1b allow for straightforward data analysis. Due to similarities in functional groups, many amino acid residues cannot be immediately distinguished by their spectra. This applies, for example, to alanine, leucine, isoleucine and valine, which contain terminal CH$_3$ groups. Nevertheless, heteroatom-containing functional groups or aromatic ring systems can clearly be instantly identified in the TERS spectra since those bands do not overlap with other modes. For example, in Fig. 1b, several modes at 650–800 cm$^{-1}$ (orange spots at 0–6.5 nm and again at 10–17.5 nm) could be assigned to cystine assemblies (cys-cys). Evidently, those modes appear at varying spectral positions and therefore must originate from different cys-cys residues with different molecular environments. This is not surprising because the insulin sequence comprises six cysteine residues (pair-wise linked via a disulfide (S-S) bridge) with different conformations. It must be noted that in all presented experiments, S-S modes were obscured by the silicon signal of the tip and
Table 1. Marker bands in the TERS spectra of different insulin fibrils used for the identification of specific amino acids.

| Amino Acid | Wavenumber/cm$^{-1}$ | Assignment |
|------------|------------------------|------------|
| asn, gln (NH$_3^+$) | 1070–1080 | lys$_{34–37}$ (NH$_3^+$, add. to 1150 cm$^{-1}$) |
| arg$_{35,36}$ (NH$_3^+$, add. to 1170 cm$^{-1}$) | 1083–1090 |
| asn, gln (NH$_3^+$) | 1139–1148 |
| arg$_{36,43}$ (NH$_3^+$, add. to 1075 cm$^{-1}$) | 118–119, 1196–1204 |
| CH, tyr$_{34}$ (CH, add. to 825/855 cm$^{-1}$) | 1198–1216 |
| CH, tyr$_{34}$ (CH, add. to 825/855 cm$^{-1}$) | 1210–1216 |
| CH, tyr$_{34}$ (CH, add. to 825/855 cm$^{-1}$) | 1226–1230 |
| CH, tyr$_{34}$ (CH, add. to 825/855 cm$^{-1}$) | 1237–1250 |
| CH, tyr$_{34}$ (CH, add. to 825/855 cm$^{-1}$) | 1251–1255 |
| CH, tyr$_{34}$ (CH, add. to 825/855 cm$^{-1}$) | 1256–1267 |
| CH, tyr$_{34}$ (CH, add. to 825/855 cm$^{-1}$) | 1268–1271 |
| CH, tyr$_{34}$ (CH, add. to 825/855 cm$^{-1}$) | 1271–1277 |
| CH, tyr$_{34}$ (CH, add. to 825/855 cm$^{-1}$) | 1278–1283 |

Consequently, were excluded from further analysis. This also applies to C-C modes at approximately 950 cm$^{-1}$ overlapping with the silicon overtone (from the TERS tip). The consecutively detected bands at approximately 1700 cm$^{-1}$ could be assigned to the carboxylic residues of glutamic acid (glu) and the C-termini. The characteristic amino modes of asn/glcn can be identified at 1060 cm$^{-1}$ and 1135 cm$^{-1}$ (see Table 1 and the complete dataset in Supplementary Table S3).
In the third experiment investigating a different fibril with a different tip, the probed area was expanded to 49.5 nm (line with a point-to-point distance of 1 nm). Based on the first two experiments, it was concluded that although a larger step-size compromises the spatial resolution slightly, larger areas can be probed without the loss of spectral information. The acquired spectra (see Fig. 2a) were processed as above, and the respective band parameters plotted in Fig. 2b.

As in the previous experiments, cys-cys and asn/gln can be frequently found. This observation agrees well with the recently calculated propensity values for selected amino acids on insulin fibril surfaces. The examined area in Fig. 1 is depleted of tyrosine, whereas Fig. 2 shows abundant tyr (blue spots in Fig. 2b). At first glance the repeated detection of this residue between ca. 10–20 nm of the measured line might give the impression of a much poorer spatial resolution compared to the fist two experiments. Due to the TERS tip preparation process it is only natural that not all tips show exactly the same resolution. With respect to the data in Fig. 1 and Supplementary Fig. S1 denoting a spatial resolution around 1 nm we assume that the resolution in Fig. 2 was within the same range. Most likely a cluster of tyr was present in this specific area of the fibril. Glycine (gly, lilac) and histidine (his, turquoise) can be identified at the end of the line: coexisting bands at 1020 cm\(^{-1}\) and 1320 cm\(^{-1}\) for glycine and three bands (1180 cm\(^{-1}\), 1330 cm\(^{-1}\), 1495 cm\(^{-1}\)) for histidine (assignment see Table 1). It should be noted that a combination of bands was used for reliable amino acid assignment because a single band is not specific enough in most cases. The identification of polar and non-polar amino acids along the measured line in Fig. 2b enables again the classification of regions with mainly hydrophilic character (checked) and domains with a mixture of hydrophilic and hydrophobic residues (dotted) in Fig. 2c. Similar to the previous experiment, domains with coexisting non-polar (cys-cys, gly) and polar (asn/gln, tyr, his) amino acids prevail in the measured region. Slight compositional differences compared to the previous fibril are not surprising and can be explained by the confined examined region and the often-reported polymorphism of insulin fibrils.
Regarding secondary structure elements, amide I bands are only very rarely detected within a short range (14–18 nm of the measured line in Fig. 2). According to our previous work on homopeptides, all spectra of the gly containing peptide show amid I bands. Consequently, one might expect amide I bands at the end of the measured line in Fig. 2a. However, in the case of insulin gly is embedded between amino acids with bulky side chains (e.g. phe). These bulky residues potentially lead to the suppression of amide I bands in our spectra. In contrast, bands in the amide III band region above 1260 cm$^{-1}$ are ubiquitous, and the positions remain nearly unaffected despite changes in chemical composition. Consequently, as in the previous experiment, those bands could be assigned to α-helix/unordered structures; conversely, β-sheet structures in the amide III band region around 1230 cm$^{-1}$ could not be definitively identified due to a likely overlap with the CH$_2$ modes of cys-cys and tyr$^{34–38}$.

**Figure 2.** (a) Raw TERS spectra acquired along a 49.5-nm line (step-size: 1 nm) on an insulin fibril (λ = 532 nm, t$_{acq}$ = 1 s). (b) Band fitting parameters plotted versus position on fibril; spot size corresponds to the band intensity and is normalized to each band; color code: cys-cys-orange; tyr-blue; asn, gln-magenta; amide I, III-red; his-turquoise; gly-lilac, CH, CH$_2$-green; unspecified bands-black. (c) Schematic sketch of the surface hydrophilicity, mainly hydrophilic domain-checked; domains with a mixture of hydrophilic and hydrophobic residues-dotted.
In previous studies, both residues were mainly found in β-sheet-dominated areas\textsuperscript{21,22}, and it is very likely that the probed area, which is rich in cys-cys and tyr, has a large β-sheet contribution.

Finally, the same tip was used to measure along 4 parallel lines (lateral offset: 1 nm, each with 50 equidistant measuring points). In this way, the end of a fibril was probed, as indicated by the rectangle in the topography of Fig. 3a (all 200 raw spectra are available in the Supplementary Fig. S4; all bands are given in the Supplementary Table S3).

The complete dataset was analyzed as before. Additionally, the N-FINDR algorithm\textsuperscript{46}, which has already successfully been applied to TERS spectra to distinguish cell membrane components (protein and lipids), was also implemented\textsuperscript{47}. Regarding the graph in Fig. 3b, marker bands of selected amino acids and the peptide bond (amide bands) were fitted, and the obtained intensity values were plotted to obtain TERS maps of the fibril surface.

In order to suppress outliers with very high signal intensities in the 1450 cm\textsuperscript{-1}, 1270 cm\textsuperscript{-1} and 1300 cm\textsuperscript{-1} maps the color bars were set to lower intensity values than the maximum value. The maximum values are provided separately above the color bar. Sharp transitions denoting structural differences within each line and between lines are clearly perceptible. For instance, cys-cys was detected along the 1st and 4th lines, albeit at different positions (796 and 766 cm\textsuperscript{-1}, respectively, orange map). This observation again indicates cys-cys moieties with different chemical environments. In contrast, asn/gln (magenta maps) is absent on the 1st line but almost continuously present along the 2nd and 3rd lines. Regarding the 4th line, the tip was too far from asn/gln-rich regions. It should be mentioned that tyr was exclusively detected along the first line (see Supplementary Fig. S4) and was therefore omitted from mapping. Amino acids with methyl groups (ala, val, ile and leu, gray maps in Fig. 3b) are omnipresent on lines 2–4, with varying intensities. In comparison to the previous measurements, band fluctuations in the 1320–1470 cm\textsuperscript{-1} region are less prominent, and signals below 1300 cm\textsuperscript{-1} show an overall weaker enhancement along

**Figure 3.** (a) AFM topography of a fibril for which 200 spectra on four 1 nm laterally shifted lines were recorded consecutively; each line: 49 nm with 50 equidistant measuring points; (λ = 532 nm, t\textsubscript{acq} = 5 s. (b) Maps generated from the fitted band intensities; pixel brightness corresponds to band intensity. (c) Maps obtained from N-FINDR algorithm analysis; pixels represent the contribution of so-called “endmember” spectra (linear combination of spectra). All maps show the distribution of assigned amino acids and amide bands within the investigated area.
all four lines. Hence, it can be concluded that this specific fibril area has a widespread hydrophobic character with a more homogeneous surface composition compared to the previous species.

Information on the secondary structure is again provided by the position of amide I bands (see Table 1). Along the first two lines, α-helix/unordered structures obviously dominate, and bands in the amide III band region above 1260 cm⁻¹ are present, even when amide I bands are not. This observation validates the general assignment of these bands to α-helix/unordered structures in all the presented experiments. The maps generated from the N-FINDR analysis⁶⁵,⁶⁷ (Fig. 3c) agree very nicely and provide comparable chemical information. Note that it is not possible to compare intensities of the maps in Fig. 3b, since they were obtained from completely different data treatments. A considerable advantage of this method is that not all bands of the dataset have to be assigned, but the so-called endmember spectra, which are obtained from data analysis. Each endmember spectrum matches a raw spectrum at one specific point of the measured grid, whereas all other points of the dataset are a linear combination of all endmember spectra. In other words, N-FINDR considers the entire spectrum for the analysis, whereas in the band-fitting procedure, the user decides which bands are relevant. The endmember spectra and corresponding raw spectra for the present experiment are provided in Supplementary Fig. S5. The contribution of each endmember spectrum to the dataset can be imaged in terms of a map; adding up those images yields maps such as that shown in Fig. 3c, which illustrates the distribution of the selected amino acids and amide bands. Please see refs 46–48 for details regarding the N-FINDR method and its application on Raman and TERS spectra. Such multivariate analysis can be more straightforward and timesaving compared to the extensive assignment of single features necessary in the band-fitting procedure, particularly with large datasets. Nonetheless, an unequivocal advantage of the band-fitting procedure is the partitioning of bands and the visualization of correlating bands. In this way, non-specific modes can be assigned, possibly leading to the identification of further sample components. Clearly, the respective maps in Fig. 3b,c match nicely. Note deviations in the amide III-fitted map and N-FINDR map are due to outliers (white pixels in Fig. 3b), which cause all other pixels to appear dark-red and are thus hardly distinguishable from amide III-free areas (black pixels). In particular, the sharp transitions in the orange cys-cys maps highlight the confined volume that is probed with the TERS tip and the high spatial resolution of the measurement.

Conclusion
We present a systematic TERS investigation on spatially confined areas for four individual insulin fibrils. Clearly the experiments reveal the consistency of the acquired data and the stability of the setup as a basis for all of the presented data and the respective discussion. TER spectra, without any further enhancement of the so-called gap mode or any further molecular resonance enhancement, were collected at equidistant points along up to 50 nm-long lines on insulin fibrils with a point-to-point distance between 0.5 and 1 nm. This way the surface composition of the probed regions could be determined with extreme lateral resolution. Due to the intrinsically high resolution even amino acids without specific marker bands could be identified by exclusion arguments and spectra could consequently be assigned to an exact portion of the insulin sequence.

To the best of our knowledge, these experiments for the first time allow the identification and full spectral mapping of alternating hydrophobic, hydrophilic (or a mixture of both) nanometer-sized domains on insulin fibrils. These properties play the most significant role in amyloid fibril biological activity and are associated with toxicity as well as with the mechanism of fibrillation. It is noteworthy that this information is not readily accessible by any other technique. The subsequent data evaluation furthermore enabled the assignment of secondary structures based on amide III bands, which is important if the amide I spectral region provides insufficient information. Finally, to investigate the possibility of an automated method for analyzing large TERS datasets, spectra were assessed with the N-FINDR algorithm. The agreement of the maps generated from band fitting with those from N-FINDR analyses indicates that N-FINDR provides a convenient and automated method to obtain detailed information from a TERS dataset of a peptide. In particular, such a method is promising for a first assessment when considering much larger datasets.

The results are encouraging for future projects aimed at the identification and localization of interaction sites of amyloid fibrils with membranes and drugs. The presented combination of surface characterization and data evaluation is not restricted to protein assemblies but can be applied to a wide range of samples for (bio-)chemical surface analysis at the nanometer scale.

Methods
Sample Preparation. Bovine insulin (Sigma-Aldrich, St. Louis, MO) was dissolved (60 mg/mL) in HCl (pH 1.5) and heated at 70°C for 2.5 hours without stirring. After cooling to room temperature, an aliquot of the fibrillar gel was re-suspended in HCl (pH 1.5) solution with a 1:100 dilution factor (V/V), and 10 μL was dropped onto a pre-cleaned cover slip. After incubation for 2–3 min, the suspension was gently sucked off from the glass surface with a pipette and washed twice with pH 1.5 HCl. The samples were dried under an argon flow prior to TERS measurements.

TERS measurements. TERS measurements were performed using an AFM mounted on an inverted microscope, as previously described in detail in ref. 21. Commercial non-contact AFM cantilever tips (NSG10, NT-MDT) were evaporated with 25 nm silver and stored under argon until use. Closed loop feedback in the x and y directions was used to maintain the relative positioning uncertainty below 1 nm. The absolute drift of the system was detected by measuring a fibril sample and was between 15–40 nm/hour with a local precision of 1–2 nm for fibril features measured on a 1 x 1 μm scan area. Experimental conditions were selected such that in these limits the maximum drift was 4–10 nm for the longest experiment. The nominal absolute positioning precision of the AFM’s closed loop system was 0.3 nm so the drift between two positions in an experiment is negligible ensuring a reliable basis for the relative location analysis. For excitation, a laser at 532 nm with a power...
of 770 μW was used on the sample. Depending on the activity of the tips, the acquisition time was set to 10 s (protofilament), 1 s (single lines) and 5 s (four lines) to achieve a signal-to-noise ratio allowing the assignment of at least 10 bands per spectrum. After each measurement, a reference spectrum was recorded for a substrate spot next to the fibril to exclude tip contamination. All shown spectra are raw spectra and were neither baseline nor background corrected.

**Band-fitting procedure.** Non-linear band fitting (Lorentzian profile) was accomplished using Igor Pro 6.12 (Wavemetrics, USA). Only bands with a signal-to-noise ratio above two were considered. Depending on the spectral areas of interest, up to ten bands were fitted simultaneously. For the graphs in Supplementary Fig. S2 and Figs 1 and 2, intensities were plotted as spots versus the measured position on the fibril. The spot size corresponds to the band intensity and is normalized to each band separately. The maps in Fig. 3b were generated from the fitted intensities of selected bands by plotting them versus a position of the fibril.

**N-FINDR analysis.** Data were imported into MatLab v2013a (The Mathworks, Natick, MA) and analyzed with in-house written scripts. The spectral region was limited to 600–1800 cm\(^{-1}\) and was vector normalized before being analyzed with the N-FINDR spectral unmixing algorithm using a non-negativity constrained fitting of endmembers. A detailed protocol of the method can be found in ref. 48.

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**Author Contributions**

T.D.-G., D.K., I.K.L. and V.D. conceived and designed the experiments. D.K. prepared the samples. T.D.-G. performed the measurements. T.D.-G., D.K., M.A.B.H. and P.S. analyzed the data, T.D.-G. wrote the manuscript.

All authors reviewed the manuscript.

**Additional Information**

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