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Parkinson's protein α -synuclein : membrane interactions and fibril structure

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**4 Characterization of Fibrils Obtained by Seeded
Fibrillization of a Series of Spin-Labeled α -
Synuclein Variants by Transmission Electron
Microscopy**

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4.1 Introduction

The protein α -Synuclein (α S) is 140 amino-acids long and is present mainly in the human brain. The protein monomer is intrinsically disordered (1), which under certain conditions forms amyloid fibrils (2–9). Polymorphism of fibrils was described previously, for example, for Amyloid β (10), Immunoglobulin (Ig) light chains (11), ovalbumin (12), lysozyme (13), and also α S (14–17).

In an accompanying pulsed-EPR study (chapter 5), we sought to determine the inner intrinsic fold of α S in fibrils using α S with spin labels at different positions. For that, independent information about the fibril morphology was desired. The main question was whether the fibrils of all spin-label constructs have the same morphology. To do so, we prepared the fibrils of nine doubly spin-labeled variants (α S42/69, α S42/75, α S42/85, α S56/69, α S56/75, α S56/90, α S69/85, α S69/90, and α S75/85) and characterized the morphology by transmission electron microscopy (TEM). All the proteins were fibrillized as described in the Materials and methods section. We used the same fibrils for the successive EPR study in chapter 5.

In this study, we found that the morphology of the fibrils of all protein constructs grown under the conditions described in the present study is similar.

4.2 Materials and methods

4.2.1 Preparation of fibrillar α S

The expression, purification and MTSL-labelling of the protein α S has been described in chapter 5 (18–20).

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4.2.1.1 Preparation of α S-fibril seeds

For seeds, we first prepared wild type (wt) α S fibrils following the protocol from Sidhu *et al.* (17). Briefly, the wt- α S protein solution (concentration = 100 μ M, in 10 mM Tris- Cl buffer pH 7.4 containing 10 mM NaCl and 0.1 mM EDTA) was aliquoted into 15 Eppendorf tubes (Eppendorf LoProtein Bind 2 ml), 500 μ l each. All tubes were incubated at a temperature of 37 $^{\circ}$ C shaking continuously at 500 rpm in a Thermo mixer (Eppendorf). The time evolution of α S fibrillization was monitored by the standard Thioflavin T (ThioT) fluorescence assay. For each tube, fibrillization was stopped when ThioT fluorescence intensity had reached a plateau. The fibrillization was completed in 6-7 days. The content of each Eppendorf tube was divided into two samples of 250 μ l each, which were frozen quickly in liquid nitrogen and stored at - 20 $^{\circ}$ C. To start the seeded fibrillization experiment, one aliquot was thawed and sonicated in a bath sonicator (Branson 2510) for one minute to break the fibrils into seeds, which were then added to the samples to be fibrillized.

4.2.1.2 Seeded fibrillization of spin-labeled α S

We prepared the fibrils by mixing the monomeric α S (spin-labeled and wild type α S) with the wt- α S seeds. The total monomer concentration used for making fibrils was 100 μ M. To this mixture 2 % monomer equivalent seeds were added. Diamagnetic dilution was employed to diminish the effect of intermolecular interaction. We used 1:20 (SL α S:wt) diamagnetic dilution for all doubly labeled α S. A typical sample for doubly labeled α S consisted of 5 μ M spin-labeled α S, 95 μ M wt- α S and 2 μ M α S-monomer equivalent seeds. The total volume for each sample was 3.0 ml, which was aliquoted into five Eppendorf tubes and put on the thermomixer. The seeded fibrillization was performed under the same conditions

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as used for wt- α S fibrils (17). The time evolution of seeded fibrillization was monitored by the standard Thioflavin T (ThioT) fluorescence assay. Most of the mutants completed the aggregation, i.e., fluorescence intensity reached the plateau, in 24 hours except for α S69/90, which took 9 days to complete the aggregation. These samples, after harvesting, were used for the pulsed EPR experiments, described in chapter 5. A small amount of fibril solution from each sample was used to prepare TEM-grids to visualize the morphology.

4.2.1.3 Preparation of unseeded fibrils of α S69/90

The unseeded fibril of α S69/90 (referred to as **α S69/90** in the text) was prepared following the method described by Hashemi Shabestari *et al.* (21), described briefly. The total monomer concentration used for making fibrils was kept 100 μ M. We used a diamagnetic dilution of 1:20 (SL α S:wt) similar to the seeded fibrillization case. The fibrillization mixture was prepared by mixing a 5 μ M spin-labeled α S69/90 with 95 μ M wt- α S protein in 10 mM Tris-Cl buffer, pH 7.4, containing 50 mM NaCl. The total volume of the mixture was 2.8 ml, which was aliquoted into four Eppendorf tubes (Eppendorf LoProtein Bind 2ml), 700 μ l each. All tubes were incubated at a temperature of 37 °C with constant shaking at 1000 rpm in a Thermo mixer (Eppendorf). The time evolution of fibrillization was monitored by the ThioT assay. For each tube, fibrillization was stopped when the ThioT-fluorescence intensity reached a plateau. The fibrillization was completed in 6-7 days. A small amount of fibril solution was used for TEM and the remaining fibril solution was harvested as described above and double electron-electron resonance (DEER) experiment (see appendix B) was performed.

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4.2.2 Transmission electron microscopy

Negative staining of α S fibril samples was done by placing a fresh carbon-coated grid (200 meshes) on top of a drop (10 μ l) of the α S-fibril solution for 2 minutes. The grid was then washed 3 times on a drop of distilled water. Subsequently, the grids were placed directly on top of a small drop of 3.5 % uranyl acetate for 1.5 minutes and the excess uranyl acetate was blotted away by touching the grids to a filter paper at an angle of 45°. Afterwards the grids were placed in a Petri dish with filter paper to let them dry. The grids were examined with a FEI Technai-12 G2 Spirit Biotwin transmission electron microscope (FEI, Eindhoven, Netherlands) and micrographs were taken with a Veleta side-mounted TEM camera using Radius acquisition software (both Olympus Soft Imaging Solutions, Münster, Germany). Images were measured using the image processing feature within the Radius software package.

4.3 Results

Figures 4.1 and 4.2 show the TEM images of α S fibrils prepared under seeded conditions. These Figures show that all fibrils used in this study have a similar appearance, which we describe in the following. Single fibrils have a width of 5.6 ± 1 nm (Figure 4.1a, bottom, black arrows) and are often found to twin with another fibril that runs parallel to generate a fibril with a width of 9-10 nm marked by black arrows (see Figure 4.1a & Figure 4.2g, middle, and Table 4.1). Many of these twinned fibrils are twisted. We further characterize twinned fibrils with the periodicity of the twist, measured between the points marked by white arrows in Figure 4.1c and 4.2f, middle. With the exception of α S56/75, α S75/85 and α S69/90, all fibrils show families of periodicities around 170 nm, 240 nm, and 290 nm. Fibrils of α S56/75 show periodicity around 53 nm, 103 nm, and 152 nm,

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while fibrils of α S75/85 and α S69/90 show just one family of periodicity around 177 nm and 190 nm respectively.

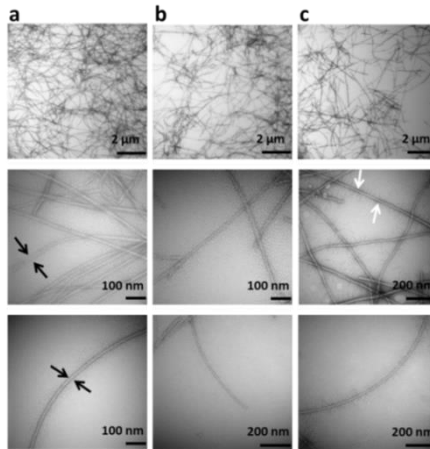


Figure 4.1. Morphological characterization of seeded fibrils of α S by TEM. a. α S42/69, b. α S42/75, c. α S42/85. Black arrows depict the width of a single fibril as shown for α S42/69 fibril. White arrows depict the points of cross-over of a twist in the fibril and the periodicity was measured between the points as shown for α S42/85 fibril.

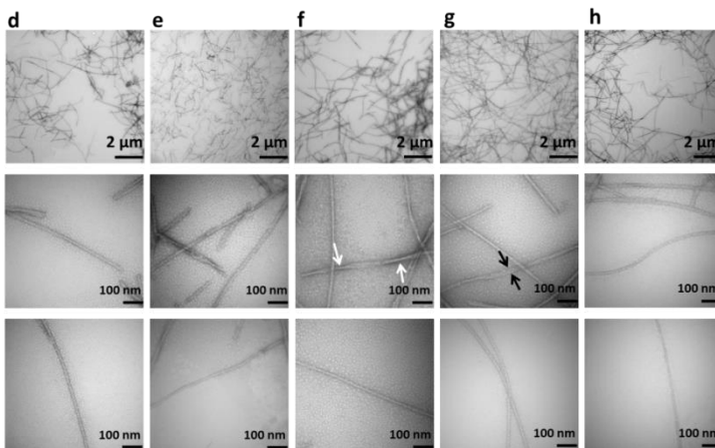


Figure 4.2. Morphological characterization of seeded fibrils of α S by TEM. d. α S56/69, e. α S56/75, f. α S56/90, g. α S69/85, and h. α S75/85. White arrows depict the points of cross-over of a twist in the fibril as shown for α S56/90. Black arrows depict the width of a twin fibril running together.

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Table 4.1. Morphology parameters of Seeded fibrils of α S based on TEM

αS fibril sample	Single fibril width (nm) \pm stdv (nm) [no of fibrils checked]	Twinned fibril width (nm) \pm stdv (nm) [no of fibrils checked]	Periodicity (nm) \pm stdv (nm) [no of fibrils checked]
αS42/69	5.5 \pm 0.9 [17]	9.45 \pm 1.0 [7]	182.5 \pm 9.5 [10], 237.8 \pm 7.9 [2], 285.7 \pm 14.7 [2]
αS42/75	5.4 \pm 1.0 [16]	10.6 \pm 1.2 [5]	165.8 \pm 8.6 [11], 235 [1]
αS42/85	6.0 \pm 0.6 [36]	9.5 \pm 1.6 [13]	164.3 \pm 9.9 [11], 227.8 \pm 15 [13], 293.4 \pm 9.7 [4]
αS56/69	6.1 \pm 1.1 [7]	9.0 \pm 1.3 [9]	171.8 \pm 7.7 [16], 242.2 \pm 6.0 [4]
αS56/75	5.7 \pm 1.0 [7]	10.6 \pm 2.2 [4]	53 \pm 6.0 [2], 103 \pm 12.4 [2], 152 [1]
αS56/90	5.2 \pm 1.1 [9]	9.2 \pm 1.0 [6]	165 \pm 6.0 [4], 206 \pm 13.5 [12], 266.8 \pm 11.4 [6]
αS69/85	6.3 \pm 1.0 [9]	11.1 \pm 2.3 [8]	181.2 \pm 6.0 [2], 308.3 \pm 6.0 [3]
αS69/90	5.7 \pm 1.0 [22]	8.7 \pm 0.9 [4]	190.4 \pm 6.0 [2]
αS75/85	5.0 \pm 1.4 [5]	9.2 \pm 0.1 [3]	177.4 \pm 9.9 [5]

To illustrate the comparison of fibril morphology, we show the width and the periodicity of fibrils (given in Table 4.1) as a scatter plot in Figures 4a and b. Figure 4.3a shows the plot of the width of fibrils as a function of the type of fibrils (single and twinned fibril), and Figure 4.3b shows the plot of the periodicities of the fibrils. Figure 4.3a illustrates that the width of the fibrils are the same within the error margins of the measurement. Figure 4.3b shows that there are three distinct clusters of periodicities for most of the fibrils. With the exception of α S56/75, the

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periodicity of the fibrils within the clusters agrees within experimental error. The fibrils of α S75/85 and α S69/90 show just one class of periodicity of 177 nm and 190 nm respectively, take into account the smallest number of observations compared to the other fibrils. For all fibril samples, the difference between the clusters of periodicities is around 50 nm.

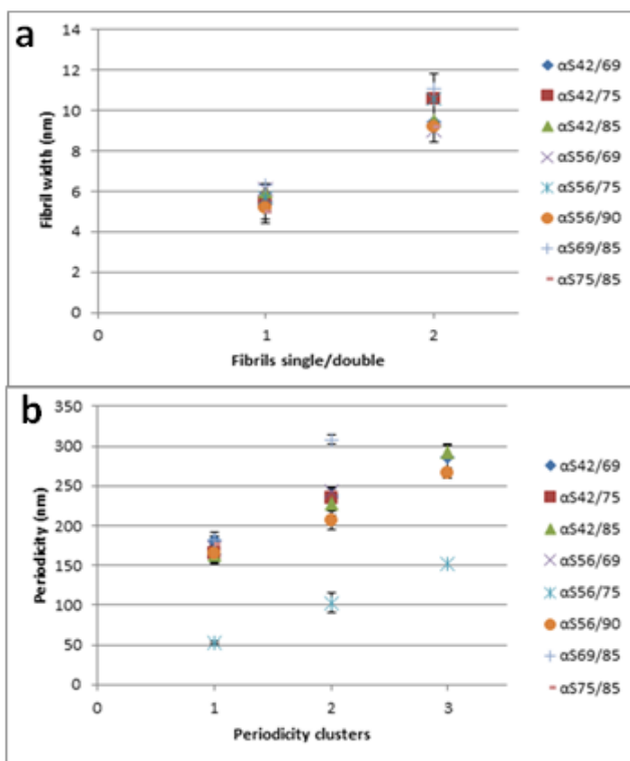


Figure 4.3. Scatter plots representing quantitative morphological features (listed in Table 4.1) of α S fibrils prepared under seeded condition (for details see Materials and methods): a. comparison of width of fibrils (given in Table 4.1); b. comparison of periodicities of fibrils (given in Table 4.1). Cluster 1: shortest periodicities; cluster 2: medium periodicities; cluster 3: longest periodicities.

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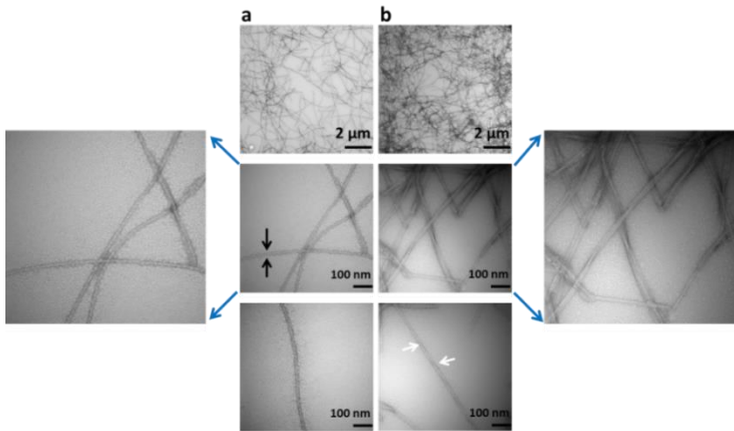


Figure 4.4. Morphological characterization of fibrils of α S by TEM. Comparison of morphology of the fibril prepared from the same α S69/90 protein a. under seeded conditions (for details, see Materials and methods); b. under the conditions described in Hashemi Shabestari *et al.* (21); Far left: middle panel of (a) scaled up (1.5x); Far right: middle panel of (b) scaled up (1.5x) for better view. Black arrows depict the width of the fibril, white arrows depict the points of cross-over.

Table 4.2. Comparison of fibril morphology of α S grown under seeded and unseeded conditions (21) based on TEM.

α S fibril sample	Single fibril width (nm) \pm stdv (nm) [no of fibrils checked]	Twinned fibril width (nm) \pm stdv (nm) [no of fibrils checked]	Periodicity (nm) \pm stdv (nm) [no of fibrils checked]
α S69/90	5.7 ± 1.0 [22]	8.7 ± 0.9 [4]	190.4 ± 6.0 [2]
αS69/90	5.6 ± 1.0 [9]	10.6 ± 1.2 [5]	116.6 ± 6.0 [4], 233 ± 8 [6], 278.7 ± 6.0 [3], 321.5 ± 9 [3], 352 [1]

Figure 4.4 shows the TEM images of fibrils prepared from the same α S69/90 protein, but under two different fibrillization conditions. Figure 4.4a shows the fibril morphology for the seeded-fibrillization conditions used in the present study (see Materials and methods), and Figure 4.4b for the fibrillization conditions

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described in Hashemi Shabestari *et al.* (21,22). For clarity, we use the abbreviation **α S69/90** for the latter fibrils (21). Comparing the images in Figure 4.4a and b, in Figure 4.4b fibrils appear somewhat wider (far right, enlarged view) than fibrils in Figure 4.4a (far left, enlarged view). Single fibrils, in Figure 4.4a, have a width of 5.7 ± 1 nm (shown with black arrows and Table 4.2), similar to the width of the fibrils shown in Figure 4.4b (Table 4.2). Fibrils in seeded conditions show twinned fibrils of width 8.7 ± 0.9 and a periodicity of 190 ± 6 nm (as shown in Figure 4.4a and Table 4.2). On the other hand, twinned fibrils of **α S69/90** show twists with several different periodicity lengths (shown in Table 4.2 and Figure 4.4b; twists are indicated by white arrows). The separation between the periodicity lengths varies from 45 nm to 100 nm. The differences in morphology of fibrils, grown under seeded conditions and conditions described by Hashemi Shabestari *et al.* (21), are larger than those seen between fibrils grown from different spin-label constructs under the same conditions i.e., under seeded conditions.

4.4 Discussion

In this study, we have used a seeded fibrillization procedure employing the conditions described in A. Sidhu *et al.* (17) (see also Materials and methods) to prepare fibrils of homogeneous morphology. Fibrils of nine different α S spin-labelled variants were prepared and their morphology checked by negative stain Transmission Electron Microscopy (TEM). The overall appearance of the fibrils of all constructs is similar. The width of single fibrils and twinned fibrils in all constructs agree within the error margin of the measurement. From this, we conclude that the fibrils of all α S constructs have the same morphology. The origin of the differences in the periods of the twisted fibrils are not clear, but since different periods occur within one sample, it is likely that different periods do not affect the internal structure of the fibril, i.e., the fold of the protein within the

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fibril. We also observe that fibrils grown under different conditions, i.e, α S69/90 and α S69/90, have a difference in their TEM appearance (shown in Figure 4.4).

Fibrils isolated from the brain of patients of Parkinson's disease show a width of ~ 5 nm for a single untwisted α S-fibril (23), which is consistent with our finding in the present study. Vilar *et al.* (14) have reported α S-fibrils that were described as straight fibrils with a width of 5.5 ± 0.5 nm. They also reported that two such straight fibrils often run together with a width of 13 ± 1.0 nm. The single-fibril width in this case is in agreement with the fibril width for a single fibril found in our study. The difference between 10 nm and 13 nm for the width of twinned fibrils may be related to the arrangement of the two filaments. If the two filaments are associated sidewise, this will give a larger width, i.e., in the range of 11-13 nm, than if the two filaments are partly on top of each other, which could explain the lower value of the width in the range of 9-10 nm observed in the present study. Bousset *et al.* (15) have reported cylindrical fibrils with a width of 13 ± 2 nm, which may be caused by the lateral association of two to three filaments as discussed above.

Comparing our fibril morphology with that of Sidhu *et al.*, (17) we note that a. the height measured by AFM is comparable to the width we observe, and b. the lengths of periods differ. In Sidhu *et al.*, (17) at least 100 fibrils were measured and the periodicity given is averaged over the entire length of fibrils, selecting fibrils of minimally 1 μ m long each. Such details could not be obtained in the present study, where we compare a large number of samples. Therefore, the periodicity information from the present study is less reliable than the one in Sidhu *et al.* (17). In addition, the differences in imaging methods, AFM and TEM, may lead to systematic variations in the parameters observed here and in Sidhu *et al.* (17).

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In conclusion, we observed that the fibrils of different protein constructs grown under seeded conditions are similar in morphology. We also found that the difference in morphology for the fibrils, grown from the same protein under the seeded conditions and the conditions described in (21), i.e., unseeded conditions, is larger compared to those observed between fibrils grown under the seeded conditions. For direct comparison with previously published fibril morphologies (14,15,17,24,25), observation of a larger number of fibrils, higher resolution techniques like cryo-electron microscopy (cryo-EM) and mass-per-length-ratio measurements by scanning transmission electron microscopy (STEM) (26) would be needed. These were beyond the scope of the present investigation.

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Appendix B to chapter 4

To check whether the difference in fibrillization conditions for α S69/90 and α S69/90 affects not only the morphology (see above), but also the internal structure, i.e., the fold of the protein inside fibrils, we performed DEER measurements on fibrils of α S69/90 and α S69/90. Figure B1 shows the comparison of the DEER traces of α S69/90 and α S69/90. The two traces look different. The modulation depth of α S69/90 is larger than of α S69/90 (see Table 5.2 in chapter 5). The modulation depth of α S69/90 is small, therefore, the distance distribution of this sample is not meaningful, because only a small fraction of the spin population contributes to it. As both fibrillizations were made with the same protein batch, the difference in modulation depth shows that for α S69/90, less spin-pairs are in the sensitive distance range of DEER (2 nm – 5 nm), which we interpret as a difference in the internal α S-fold between the two types of fibrils, α S69/90 and α S69/90.

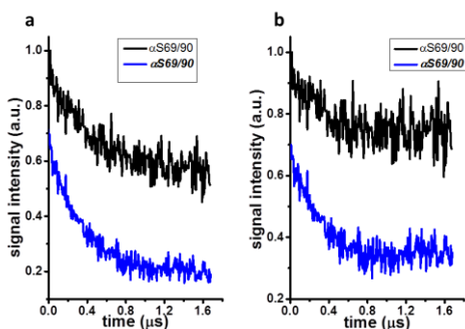


Figure B1. Comparison of DEER time traces of α S 69/90 fibril samples prepared under different fibrillization conditions: a. DEER time traces of α S69/90 (black lines) and α S69/90 (blue lines) before background corrections, b. DEER time traces of both fibril samples after background correction (black line: α S 69/90, blue lines: α S 69/90) (for details see text). Both the DEER traces are normalized to one by dividing the traces by the maximum of their intensity. DEER traces are shifted vertically with respect to each other for better visibility.

