Modulation of the extent of structural heterogeneity in α-synuclein fibrils by the small molecule thioflavin T

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The transition of intrinsically disordered, monomeric α-synuclein into β-sheet–rich oligomers and fibrils is associated with multiple neurodegenerative diseases. Fibrillar aggregates possessing distinct structures that differ in toxicity have been observed in different pathological phenotypes. Understanding the mechanism of the formation of various fibril polymorphs with differing cytotoxic effects is essential for determining how the aggregation reaction could be modulated to favor nontoxic fibrils over toxic fibrils. In this study, two morphologically different α-synuclein fibrils, one helical and the other ribbon-like, are shown to form together. Surprisingly, a widely used small molecule for probing aggregation reactions, thioflavin T (ThT), was found to tune the structural heterogeneity found in the fibrils. The ribbon-like fibrils formed in the presence of ThT were found to have a longer structural core than the helical fibrils formed in the absence of ThT. The ribbon-like fibrils are also more toxic to cells. By facilitating the formation of ribbon-like fibrils over helical fibrils, ThT reduced the extent of fibril polymorphism. This study highlights the role of a small molecule such as ThT in selectively favoring the formation of a specific type of fibril by binding to aggregates formed early on one of multiple pathways, thereby altering the structural core and external morphology of the fibrils formed.

The conversion of soluble, functionally active proteins into insoluble, β-sheet–rich, aggregated structures is associated with a variety of neurodegenerative diseases such as Parkinson’s disease (PD) and Alzheimer’s disease (AD). Detailed structural analyses of the characteristics of cross-β-sheet architecture (5–7) present in these aggregates reveal their polymorphism (8). It is now known that fibrils formed by a single protein can show multiple distinct conformations under different growth conditions (9–11) as well as under identical growth conditions (12–15). The importance of studying fibrillar heterogeneity originated from the prion strain phenomenon, in which a single prion protein is known to cause multiple different pathologies by adopting amyloid-like conformations that differ mainly in their external morphologies and molecular structures (16–18). A prion strain propagates a specific pathology faithfully by presenting a specific amyloid template for existing monomer to add on to. Proteins other than the prion protein can also acquire different fibrillar morphologies, showing different levels of toxicity, and propagate faithfully (5, 10). Importantly, fibrils of different morphologies, which differ in their cytotoxicity levels, have been shown to exist in the brains of different AD patients (8, 19). The differences in the toxicity potentials suggest that various fibrils may have different levels of stability, packing, and hydrophobicity (20–22). It is now clear that fibril polymorphism is responsible for different pathological phenotypes (23). It is therefore important to understand the origin of fibril polymorphism.

In PD, the central molecular species is the protein α-synuclein, which is an intrinsically disordered protein of 140 residues. It is expressed mainly in the neurons of the central nervous system (CNS). It acquires helical and β-hairpin structure upon binding to membranes and β-wrap proteins, respectively (27, 28). The function of this protein has not yet been ascertained conclusively, although some studies suggest that it is involved in the process of vesicle release and trafficking (29). α-Synuclein is known to aggregate and form Lewy bodies in dopaminergic neurons of the brain. Lewy bodies are made of cross-β-sheet–rich structured aggregates called amyloid fibrils. The process of α-synuclein fibrillation is associated with a variety of neurodegenerative diseases besides PD, including dementia with Lewy bodies (1), AD, and multiple system atrophy (2, 3). α-Synuclein has been shown to form differently structured amyloid aggregates (10, 12, 13, 15), but there is little understanding of how this happens.

Structural heterogeneity in α-synuclein fibrils originates mainly during the multi-step process of fibrillation in which monomers self-assemble into various on- and off-pathway oligomers that vary in size, shape, structure, stability, and packing (30, 31). Structurally distinct oligomers appear to grow into different types of fibrillar aggregates with structural cores that resemble the oligomers (31). The structural core of α-synuclein fibrils have been characterized by solid-state NMR (10, 12, 32–34), hydrogen–deuterium exchange mass spectrometry (HDX-MS) (31, 35), HDX-NMR (13, 33), electron paramag-
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Thioflavin T (ThT) modulates fibrillar heterogeneity by its binding to cross-β-sheet structures that stack on top of each other to form a protofilament (39). The protofilaments twist together to form fibrils (39). Cryo-electron microscopy and scanning transmission electron microscopy studies have also been used to characterize the structure of α-synuclein protofibrils. Two protofibrils, each made of three β-strands from the same subunit, form the protofibrils in which sequence segment 8–94 forms the structural core (15). Two protofibrils were found to associate with each other asymmetrically to form fibrils (15). It was hypothesized that the number of protofibrils and their nature of association in the fibrils may determine the final morphology of α-synuclein fibrils and, hence, their structural heterogeneity (15).

The process of α-synuclein fibrillation has been described by the nucleation-dependent polymerization model, which involves a lag phase followed by an exponential phase (40). During the lag phase, monomers undergo structural rearrangements to form transient nuclei, which further grow by the addition of monomers to form fibrils. The molecular structure of the nucleus is likely to determine the molecular structure of the fibrils (41), and modulation of the nucleation and elongation rates by varying the growth conditions is expected to modulate the amount of various fibrillar polymorphs (42).

Under different solution conditions, α-synuclein has been shown to form two types of fibrils, which differ in their morphology, structural core, toxicity, and infectivity (10). Even under the same aggregation conditions, α-synuclein can form morphologically and structurally different fibrils (12, 13, 15). Understanding the molecular mechanism by which different types of aggregates arise and how fibril formation can be modulated is crucial for the development of therapeutics for PD and other protein aggregation diseases.

Small molecules have been used extensively to modulate and inhibit the process of aggregation for several disease-linked proteins (43–45) including α-synuclein (46, 47). The binding of small molecules to fibrils can reduce the amount of toxic oligomers by blocking fibril dissociation (48), by modifying the fibril surface, which acts as an efficient catalyst to generate toxic oligomers (49), or by driving the equilibrium toward fibril formation (45). The small molecule thioflavin T (ThT) is used widely to monitor the aggregation process for many proteins because of its ability to bind to cross-β-sheet structures found in amyloid fibrils, which modulates the fluorescence properties of ThT (50). ThT binds to different types of fibrils with different affinities (51), and it can accelerate protein aggregation by binding to monomer or fibrils (52, 53). It is, however, not known whether small molecules such as ThT that bind fibrils can modulate a fibril formation reaction, such that one type of fibrils is preferred over another, or whether the presence of such small molecules during fibril formation affects the internal structure as well as the external morphology of the fibrils.

In this study, the process of α-synuclein fibrillation in the absence and presence of ThT was studied. In the absence of ThT, two types of coexisting fibrils were observed. 70% of the fibrils were helical in external morphology and had a shorter structural core, whereas the remaining 30% of the fibrils had a flat, ribbon-like morphology and an extended structural core. The addition of ThT during aggregation enhances the rate constant of α-synuclein fibril formation and reduces structural heterogeneity with only ribbon-like fibrils being formed.

Results

Effects of ThT on the fibrillation of α-synuclein

To study the effects of ThT on the fibrillation of α-synuclein, 100 μM protein was incubated at pH 7.0 and 37 °C in the absence and presence of 1 mM ThT. The fibrillation process was monitored by measuring the ThT fluorescence emission at 482 nm (Fig. 1). In the absence of ThT, fibril formation by α-synuclein followed a nucleation-dependent polymerization mechanism with a lag phase of ~30 h duration. In contrast, the presence of ThT accelerated the fibrillation of the protein by reducing the lag phase and accelerating the elongation phase. The ThT fluorescence emission signal obtained at saturation for the reaction in the presence of ThT was 2-fold higher than for the reaction in the absence of ThT, indicating that either the amount of fibrils was greater or the fibrils differed in their binding ability to ThT. For fibrils formed in the absence and presence of 1 mM ThT, similar amounts of monomer were found to have converted into fibrils at saturation (supplemental Fig. S1). Hence, it was likely that the fibrils formed in the absence and presence of 1 mM ThT differed in their ability to bind ThT.
Effect of ThT on the size and morphology of the fibrils

Atomic force microscopy (AFM) images were obtained for fibrils formed by 100 μM M protein both in the absence and presence of ThT (Fig. 2, a and b). Two types of fibrils were observed to have formed in the absence of ThT (Fig. 2c). One type was helical, with a periodicity of 74 ± 6 nm (Fig. 2e), and the other was flat and ribbon-like with no periodicity. The mean heights of the helical and ribbon-like fibrils were 5.7 ± 0.7 and 6.3 ± 1.5 nm, respectively. Interestingly, only one type of fibril was obtained in the presence of ThT; these were flat and ribbon-like with a mean height of 7.4 ± 0.8 nm (Fig. 2d). Comparable results were obtained when the fibrils were formed from 50 μM protein (data not shown). The flat, ribbon-like fibrils formed in the presence of ThT had a larger diameter and less heterogeneity than the flat, ribbon-like fibrils formed in the absence of ThT (Fig. 2, c and d); the standard deviation of the fibril height distribution was 0.8 nm in the former case and 1.5 nm in the latter case. The difference in the heights of the ribbon-like fibrils formed in the absence and presence of ThT could conceivably result from intercalation of the ThT molecules in the fibrils formed in the presence of ThT.

To check whether the two types of fibrils behaved like prion strains, seeds were formed by sonicating the fibrils formed in the absence and presence of ThT, and seeding assays were carried out both in the absence and presence of ThT. It was found that both types of seeds (at 3% concentration) abolished the lag phase regardless of whether ThT was present during the aggregation of 100 μM α-synuclein (supplemental Fig. S2). Further-

Figure 2. Two different morphologies of α-synuclein fibrils are observed in the absence of ThT. a and b, AFM images of fibrils formed in the absence of ThT (a) and in the presence of 1 mM ThT (b) at a time corresponding to three times the t₅₀ of ThT fluorescence-monitored kinetics. c and d, show the fibril height distribution from a and b, respectively. e, the height profile of a single helical fibril from a, where an arrow points to the area from which the height profile was taken. f, the height profile of a single, flat, ribbon-like fibril from b, where an arrow points to the area from which the height profile was taken.
in the absence and presence of 1 mM ThT differed in their secondary structures. This peak near 1630 cm$^{-1}$ appeared at a lower wave number for the fibrils made in the presence of ThT compared with the fibrils made in the absence of ThT, which suggested an increase in the number of $\beta$-strands in the fibrils or a change in the twist angle of the $\alpha$-helix.

To determine whether the fibrils made in the presence of ThT had ThT incorporated into their structure, 100 $\mu$M fibrils made in the presence and absence of ThT were incubated with the same concentration (1 mM) of ThT. Free and loosely bound ThT was then removed by washing the fibril pellet with buffer following centrifugation. ThT fluorescence was measured for equal concentrations of the fibrils to compare the extent of ThT bound to fibrils made in the absence and presence of ThT (Fig. 3a). The ThT fluorescence of fibrils made in the presence of ThT was about 5-fold higher than that of the fibrils made in the absence of ThT but to which ThT was subsequently added. Comparable results were obtained when the concentration of associated ThT was determined by measuring the absorbance at 412 nm (supplemental Fig. S3a) after first dissolving the fibrils in 8 M GdnHCl. Hence, ThT remained tightly bound to the fibrils made in its presence, presumably by intercalation inside the fibrils. These results suggested that the fibrils made in the absence and presence of 1 mM ThT differed in their structures.

To determine whether the fibrils made in the presence and absence of ThT differed in their secondary structures, infrared spectra were acquired (Fig. 3b and supplemental Fig. S3, b and c). The presence of the peak near 1630 cm$^{-1}$ for both types of fibrils suggested that the fibrils had typical parallel $\beta$-sheet structures. This peak near 1630 cm$^{-1}$ appeared at a lower wave number for the fibrils made in the presence of ThT compared with the fibrils made in the absence of ThT, which suggested an increase in the number of $\beta$-strands in the fibrils or a change in the twist angle of the $\beta$-sheet in fibrils made in the presence of ThT (54, 55).

Structural characterization of fibrils by hydrogen–deuterium exchange mass spectrometry

HDX-MS was used to further characterize the difference in the internal structures of the fibrils made in the absence and presence of 1 mM ThT. In HDX-MS studies, the amide hydrogen sites that are protected against HDX can be localized to specific segments of the protein sequence by proteolytic fragmentation at low pH after the HDX reaction is complete. A peptide map of $\alpha$-synuclein, covering 100% of the sequence, was first generated by controlled proteolysis using pepsin at low pH (supplemental Fig. S4). The measured mass of each peptide was found to be the same as its calculated mass (supplemental Table S1), except for peptide 95–109 with a mass 18 daltons less than its calculated mass, probably due to water loss. A 5-min labeling pulse was given by incubating the fibrils as well as monomeric protein in deuterated buffer at pH 7.0, 25 °C. Control samples having no deuteration (0% D) and complete deuteration (95% D) were also run to calculate the amount of deuterium incorporation in different samples (supplemental Fig. S5).

Interestingly, sequence segments spanning residues 1 to 38 showed bimodal mass distributions for the fibrils formed in the absence of ThT but highly protected unimodal mass distributions for the fibrils formed in the presence of 1 mM ThT (Fig. 4). The bimodal mass distributions for the sequence segments spanning residues 1 to 38 indicated the existence of two different conformations differing in structure in this region. One of the conformations showed protection against HDX and hence was structured, whereas the other conformation showed no protection against HDX and hence was not structured in the sequence segment 1–38. Taken together, the data presented in Fig. 4 show that fibrils formed in the absence of ThT existed in at least two conformations. One of the conformations had a structured core from residues 1 to 94, whereas the other conformation had a structured core only from residues 39 to 94 (Fig. 5, a and b). On the other hand, fibrils formed in the presence of 1 mM ThT had only a single conformation with a structured core from residues 1 to 94 (Fig. 5c).

Differences in molecular structures of fibrils with different morphologies

To correlate the structural core with external morphology for fibrils formed in the absence of ThT, the relative amounts of the two conformations were quantified by fitting the bimodal mass distributions obtained for sequence segments 1–17 and 18–38 to the sum of two Gaussian distributions (Fig. 6a). In addition, the fractions of helical and flat fibrils were quantified by counting the numbers of the two types of fibrils from AFM images (Fig. 6a). About 70% of the fibrils were found to be helical, and the remaining ~30% fibrils had a flat morphology. Interestingly, about 70% of the fibrillar protein molecules had a...
structural core region extending from residues 39 to 94, whereas the remaining ~30% of the fibrillar protein molecules had a fibril core that extended from residues 1 to 94. These observations suggested that the helical fibrils are likely to have a core region extending from residues 39 to 94, whereas the flat, ribbon-like fibrils are likely to have a core region extending from residues 1 to 94. To further establish whether the flat, ribbon-like and helical fibrils had core regions extending from residues 1 to 94 and residues 39 to 94, respectively, the structural cores and morphologies of the fibrils were studied at 0.3 mM ThT instead of in the absence of ThT. Importantly, the AFM and HDX-MS studies showed that about 30% of the fibrils were helical in morphology and had a core region extending from residues 39 to 94, whereas the flat fibril region was found to be part of the structural core. This suggested that the N-terminal region plays an important role in determining the morphology of the fibrils. Helical fibrils formed in the absence of ThT did not convert into ribbon-like fibrils after incubating the fibrils in 1 mM ThT for 24 h at 25 °C (data not shown), suggesting that the fibrils have a stable structure and do not interconvert.

The presence of ThT modulated the fibrillar conformation, and it was important to determine how this occurred. To this end, α-synuclein aggregation reactions were carried out at different concentrations of ThT ranging from 0 to 1 mM, and AFM imaging was used to determine the relative fraction of helical fibrils formed at each concentration of ThT (Fig. 6c). We found that the relative fraction of helical fibrils decreased monotonically with an increase in the concentration of ThT present and that no helical fibrils had formed at 1 mM ThT concentration. The structure of the final aggregates was also determined using HDX-MS (supplemental Fig. S7). The relative amounts of the two fibrillar conformations were quantified by fitting the bimodal mass distributions obtained for the sequence segment 18–38 to the sum of multiple Gaussian distributions (supplemental Fig. S7). Interestingly, the relative amount of fibrillar
protein with a core extending from residues 39 to 94 decreased with increasing concentration of ThT (Fig. 6a and supplemental Fig. S7). Thus, the relative amount of helical fibrils decreased from 70% when the fibrillation was carried out in the absence of ThT to 0% when fibrillation was carried out in the presence of 1 mM ThT. These results showed that ThT reduced structural heterogeneity in the fibrils formed by α-synuclein.

**Monitoring the formation of α-synuclein oligomers during fibrillation**

To determine whether oligomer formation occurred during the fibril formation reaction carried out in either the absence or presence of ThT, α-synuclein was incubated at two different concentrations (100 and 690 μM (10 mg/ml)) under the fibrillation conditions (pH 7.0 and 37 °C) for 5 h. Size-exclusion chromatography was used to detect whether oligomers had formed. Oligomers were not observed to have formed in either the absence or presence of ThT (supplemental Fig. S8, a and b). Oligomers could only be observed (supplemental Fig. S8c) when aggregation was carried out at high a protein concentration (10 mg/ml) in a different (PBS) buffer at pH 7.4 and 37 °C for 5 h, as described previously (31).

**Characterization of the toxicity levels of the fibrils**

To determine whether the fibrils with different structures were differentially toxic to cells, the toxicities of the fibrils formed in the absence and presence of 1 mM ThT were measured using HEK-293T cells (human embryonic kidney cells). Equal amounts of fibrils (2.5 μM) were incubated with HEK-293T cells for 24 h, and toxicity was measured using the Wst-1 assay (Fig. 7). To eliminate the effect of any free ThT, the fibrils were first washed multiple times with Milli-Q water to remove any free and loosely bound ThT. For toxicity assays, 1 μM ThT was used as a buffer control, and cells without the addition of any buffer served as the control. It was found that about 113 ± 22% of the cells were viable after incubating them with fibrils formed in the absence of ThT, whereas about 76 ± 8% of the cells were viable in the case of fibrils formed in the presence of ThT. It could be concluded that the flat, ribbon-like fibrils formed in the presence of ThT were significantly more toxic to the cells than the fibrils formed in the absence of ThT. It would therefore appear that fibrils with a flat, ribbon-like morphology are more toxic than fibrils with a helical morphology. Control experiments were carried out to ensure that the observed toxicity was due to fibrils and not to any free (unbound) ThT. To this end, toxicity assays were carried out at various free ThT concentrations (supplemental Fig. S9). Free ThT was found to be toxic to cells at concentrations greater than 1 μM (supplemental Fig. S9) but not at lower concentrations.

**Discussion**

The current study was focused on understanding the structural and mechanistic basis for heterogeneity in α-synuclein fibrils and its modulation by the small molecule ThT. In the case of several neurodegenerative diseases, the aggregating protein concerned has been found to adopt distinct fibrillar conformations (5, 8, 15, 19). Thus, understanding the structural and physical basis for fibril heterogeneity may shed light on the different pathological behaviors of distinct fibrillar conformations.

**Structural heterogeneity in α-synuclein fibrils and their toxicity levels**

In this study, both helical and flat, ribbon-like fibrils were found to form under the same aggregation conditions (Fig. 2, a and c). In previous studies as well, α-synuclein was found to form two types of fibrils, under particular aggregation conditions, which differed in their secondary structures, morphologies, folds, and the extent and distribution of β-sheets but had the same structural core encompassing residues 38–95 (12, 13). In contrast, the two types of fibrils observed to form together in the current study had different structural cores (Fig. 5, a and b). The structural core of the helical fibrils comprises residues 39–94, and that of the flat, ribbon-like fibrils comprises residues 1–94 (Fig. 5, a and b). Multiple studies using solid-state NMR (10, 12, 32–34), HDX-MS (31, 35), HDX-NMR (13, 33), EPR spectroscopy (36, 37), and proteinase K treatment (38) have previously identified the structural core of the α-synuclein fibrils. Most of the studies have determined that residues 31–109 form the structural core of the α-synuclein fibrils (10, 31–94).
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Figure 6. Quantification of the two types of fibrils formed by α-synuclein. a, shows data for fibrils formed in the absence of ThT. b, shows data for fibrils formed in the presence of 0.3 mM ThT. The brown bars represent the percentages of fibrils that were helical and ribbon-like, as determined from counting the fibrils seen in the AFM images. The percentages of protein molecules in fibrils that had and did not have a structural core extending down to residue 1 were determined from the bimodal mass distributions seen for sequence segments 1–17 (orange) and 18–38 (green) as described in the legend for Fig. 4. The percentage of fibrillar protein molecules with a core extending down to residue 1 matches the percentage of flat, ribbon-like fibrils formed in both the absence (a) and presence (b) of 0.3 mM ThT. c, shows the percentages of fibrillar protein molecules with a core extending from residues 39 to 94, formed at various ThT concentrations, as quantified from the higher mass peak of the mass distributions of sequence segments 18–38 (supplemental Fig. S7). Also shown are the percentages of helical fibrils formed at various ThT concentrations, determined by counting the two types of fibrils seen in AFM images. The red line through the plot of % helical fibril data is described as 63 exp(−3.8 [ThT]). The error bars represent S.D. from three independent experiments.

Figure 7. Cytotoxicity of α-synuclein fibrils. The cytotoxicity of α-synuclein fibrils (2.5 μM) to HEK-293T cells was measured using water-soluble tetrazolium (Wst-1 assay kit). Fibrils were made by incubating 100 μM α-synuclein at 37 °C in the absence and presence of 1 mM ThT. Fibrils were added to a final concentration of 2.5 μM to the cells in each well. The concentration of ThT associated with 2.5 μM fibrils was ~0.25 μM. After incubating the cells for 24 h, Wst-1 reagent was added to the wells. After incubating them for 2.5 h, the OD at 477 nm was measured. Absorbance data were normalized to that of the untreated cells (Control) having 100% cell viability (ns = non-significant; ****, p < 0.0001 using an unpaired t test). Buffer represents cells treated with only 1 μM ThT as described under “Experimental procedures.” The error bars represent S.E. from five independent experiments, each with three replicates.

12, 13, 15, 31–38). In a previous study, ribbon-like fibrils were found to form under one aggregation condition and cylindrical fibrils under another aggregation condition, with the two types also differing in their morphology, structural core, toxicity, and infectivity (10). The cylindrical fibrils observed in that study resembled the helical fibrils observed in the current study in having a structural core composed of residues 39–94; and the ribbon-like fibrils observed in that study were similar to the ribbon-like fibrils observed in the current study in having a structural core formed by residues 1–94 (10). Although the structurally distinct fibrils were shown to differ in their toxicity potentials (10), the properties of the fibrils that determine their toxicity potentials are not well-known. The factors that are likely to affect the toxicity of fibrils include their stability, their ability to interact with membranes, and their surface hydrophobicity (20–22).

The flat, ribbon-like fibrils, which possess the extended structural core, are somewhat more toxic than the helical fibrils (Fig. 7). Interestingly, in a previous study, fibrils formed by peptides of two different lengths, which were derived from the α-synuclein NACore (segment 68–78) and subNAC (segment 69–77) regions, which differed in their structural core, were also found to differ in their cytotoxicity (39). Fibrils made of the longer peptide (NACore), which had a longer structural core, were more cytotoxic than those made by the shorter peptide (39). Further studies are required to delineate the biological importance of the length of the amyloid core.

It is interesting to note that although the fibrils formed in the absence of ThT include flat ribbon-like fibrils similar to those formed in the presence of ThT, which are toxic, little if any toxicity was observed for them. It seems that this might be due to the relatively low proportion of ribbon-like fibrils formed in the absence of ThT.

It should be noted that in this study, toxicity was measured after the addition of the fibrils to cells for 24 h. It is possible that during this incubation with the cells, the fibrils break down into smaller aggregates and that it is these smaller aggregates that are toxic. In this context, it should also be noted that oligomers and prefibrillar aggregates formed by α-synuclein are reported to be toxic (56–60) as are fibrils (10, 61, 62).
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Different internal structures lead to different external fibril morphologies

Different fibril morphologies can originate from distinct molecular structures or from different arrangements of the same molecular structure (6). In the current study, α-synuclein fibrils possessing different morphologies were found to comprise internal structural cores of different lengths. It is likely that the fibrils differing in the length of the structural core also differ in extent and organization of β-sheets. The observation that the flat, ribbon-like fibrils with an extended structural core had a larger diameter/height than the helical fibrils (Fig. 2, c and d) suggests that they are likely to have a higher fold symmetry than the helical fibrils, as observed previously for Aβ fibrils (6). In the case of the small protein barstar as well, fibrils of very different morphology and diameter, which were, however, formed under different solution conditions, were found to have inner structural cores of different lengths (63, 64). At present, the link between a short structural core and helical fibrillar morphology, and between a longer structural core and flat, ribbon-like fibrillar morphology is not understood in the case of α-synuclein.

Origin of α-synuclein fibril heterogeneity and its reduction by ThT

Structural heterogeneity in α-synuclein fibrils could arise from heterogeneity at the monomer level, which would lead to the formation of distinct nuclei. Because of its intrinsically disordered nature, α-synuclein can adopt various conformations in a given solution condition as characterized by electrospray ionization mass spectrometry (65), single-molecule AFM (66), and NMR (67). Modifications in the amino acid sequence and solution conditions have been shown to modulate the conformational composition of the ensemble of α-synuclein molecules (66). The conformational diversity could determine the relative proportion of different aggregates that can be formed, as suggested previously for other proteins (23). It can then be expected that the stabilization of a specific conformational state upon binding to a small molecule might lead to the formation of specific aggregates.

Structural heterogeneity in α-synuclein fibrils could also arise from the utilization of multiple pathways for fibril formation. For several other proteins, including barstar (64, 68) and the mouse prion protein (70), structurally distinct amyloid fibrils have been shown to form on different aggregation pathways under different aggregation conditions. In the case of α-synuclein, morphologically and structurally different aggregates have been shown to form from structurally distinct oligomers (31). One oligomer was found to be on pathway to fibril formation and had a structural core similar to that of fibrils, and the other oligomer with a different structural core grew into amorphous aggregates (31). In the current study, the inability to detect the formation of oligomers during fibril formation under the aggregation conditions used, either in the absence or presence of ThT (see “Results” and supplemental Fig. S8), did not permit us to determine whether the ribbon-like and helical fibrils arose from structurally distinct oligomers formed on different pathways of fibril formation.

The formation of distinct α-synuclein fibrils suggests that multiple nucleation events can take place under the same aggregation condition. Changes in the aggregation condition may affect the nucleation rate and elongation rate differently for distinct nuclei, which would play an important role in modulating structural heterogeneity. In the current study, the observation that the presence of ThT accelerates nucleation and elongation (Fig. 1) suggests that ThT binds to early species formed during the fibrillation of α-synuclein. Binding of ThT to early aggregates on one pathway, when multiple pathways are operative, will stabilize those aggregates and result in a reduction in fibril heterogeneity.

Role of ThT in modulating the fibril formation reaction of α-synuclein

ThT is known to bind to cross-β-sheet structures found in amyloid fibrils and has been used widely to monitor the amyloid fibril formation reaction of many proteins (50). The binding of ThT to fibrils can occur in different ways. ThT can bind perpendicular to the long fibril axis in the cavities formed by the side chains of aromatic/hydrophobic residues across consecutive β-strands on the surface of the β-sheet in the fibrils (51, 71–73). ThT is also known to bind in a parallel orientation to the peptide strand (74). It also appears that ThT can bind to the peptide backbone via π–π interactions (75). Hence, it is not surprising that ThT can bind diverse types of amyloid fibrils with different affinities (51). If fibrillar structures (nuclei), which form very early, differ in their binding affinity for ThT, then the early structure (nucleus) that binds most tightly to ThT will be stabilized and, hence, populated the most. The relative amounts of different fibrils formed will reflect the relative amounts of the initial aggregates (nuclei). Hence, the relative amounts of different fibrils that form will depend on the concentration and binding affinity of the ThT present. Thus, ThT must bind strongly to the early fibrillar structures (nuclei) that lead to the formation of the flat, ribbon-like fibrils. In fact, ThT is seen to remain bound to the final, mature, flat, ribbon-like fibrils (Fig. 3a).

The observation that ThT accelerates the fibrillation of α-synuclein by increasing the nucleation rate (as indicated in the reduction of the lag phase) (Fig. 1) suggests that ThT does indeed bind to early structures (nuclei) formed during the fibrillation process, resulting in an increase in the nucleus concentration and the elongation rate. In the case of Aβ, a previous study shows that the nucleation rate is affected more than the elongation rate in the presence of ThT, which also suggests that ThT binds to the early structures on the fibrillation pathway (52, 76).

It is also possible that ThT accelerates the amyloid fibril formation reaction by binding to monomer. In the case of α-synuclein, the electrostatic interaction of ThT with monomeric α-synuclein might favor nucleation or the conformation conversion of monomer to aggregation-prone structures that ultimately form the nucleus. Indeed, ThT is known to bind to the negatively charged C terminus of monomeric α-synuclein (53). Nevertheless, fibrils made from α-synuclein truncated at its C terminus can still bind ThT (77), indicating that ThT binds differently to monomer and to fibril.
The C terminus is known to be the most solvent-exposed region in both oligomers (31, 78, 79) and fibrils (10, 12, 13, 15, 31–38). The C terminus is also known to stabilize the disordered conformation of the protein by interacting with the N terminus or NAC domain (80–83). It is likely that the interaction of the C terminus with the N-terminal region suppresses the participation of the latter in forming the structural core of amyloid fibrils (53). Not surprisingly, then, C-terminal truncation significantly accelerates the fibrillation of α-synuclein (77, 84). Binding of the C-terminal domain to the NAC region protects the NAC region from participating in aggregation. Binding of the C terminus to other proteins (85), polyamines (86, 87), and metal ions (9, 88) increases the rate of fibril formation, suggesting that neutralization of the negatively charged C terminus prevents its binding to the NAC region and the N terminus of the protein. Thus, the interaction of ThT with the C terminus of α-synuclein may play an important role in modulating the fibrillation reaction.

It should be noted that, in this study, a range of ThT concentrations was used to investigate the effect of ThT in modulating structural heterogeneity in α-synuclein fibrils (Fig. 6). It was found that fibril formation by α-synuclein was not modulated at ThT concentrations below 30 μM, a concentration known to be the critical micelle concentration for ThT (89). It seems therefore that structural modulation by ThT during fibril formation, as seen in this study, is effected by micellar ThT. ThT is used widely to monitor fibril formation reactions, and the present study suggests that it is safe to do so at concentrations below its critical micelle concentration.

In summary, the small molecule ThT has been shown to modulate the fibril formation reaction of α-synuclein and to thereby modulate the structural heterogeneity of the fibrils that form. The presence of ThT during the fibril formation favors the formation of flat, ribbon-like fibrils over the formation of helical fibrils. The flat, ribbon-like fibrils have inner structural core extending from residues 1 to 94, whereas the helical fibrils have an inner structural core extending from residues 39 to 94. The current study highlights the potential use of small molecules in modulating the fibril formation reaction of proteins so that the less toxic aggregates are favored over the more toxic aggregates.

Experimental procedures

Protein expression and purification

The plasmid pRK172 containing the human α-synuclein gene was a kind gift from Prof. A. L. Fink. The protein was expressed and purified as described previously (90) with some modification to the procedure. *Escherichia coli* BL21(DE3) codon plus (Stratagene) cells transformed with pRK172 were grown overnight at 37 °C in LB medium containing 100 μg/ml ampicillin and then subcultured into 1000 ml of LB containing 100 μg/ml ampicillin. At an OD_{600} of 0.8–1.0, the cells were induced by adding IPTG at a final concentration of 10 μg/ml and pelleted down after 5 h. They were resuspended in osmotic shock buffer (30 mM Tris-HCl, 40% w/v sucrose, 2 mM EDTA, pH 7.2), incubated at 25 °C for 15 min, and centrifuged to remove supernatant. The pellet was resuspended in Milli-Q water containing about 1 mM MgCl₂, and the supernatant was obtained after centrifugation. This supernatant was dialyzed twice against 20 mM Tris-HCl buffer, pH 8.0, at 4 °C. The protein was denatured by adding 8 M urea solution and loaded onto a DEAE FF ion-exchange column (5 ml, HiTrap, GE Healthcare). The loaded protein was washed and then eluted out using a gradient of 100–300 mM NaCl. The eluted protein was concentrated by ultrafiltration (Millipore), flash-frozen in liquid N₂, and stored at −80 °C.

Chemicals, buffers, and aggregation condition

All of the reagents used were of the highest purity grade available from Sigma-Aldrich unless specified otherwise. GdnHCl was procured from USB Corp. The stored protein was concentrated by filtration using a YM3 filter (Millipore) and denatured in 6 M GdnHCl before injecting it into a size-exclusion column (Superdex 200 10/300 GL). The protein was eluted out in 20 mM sodium phosphate, 0.01% sodium azide, and 0.1 mM EDTA, pH 7.0 (aggregation buffer) and passed through a YM100 filter to remove any aggregated protein. For aggregation reactions, 500 μl of 100 μM protein was agitated at 750 rpm and 37 °C (Eppendorf ThermoMixer) in a 1.5-ml centrifuge tube. For aggregation reactions in the presence of ThT, 500 μl of 100 μM protein solution containing 0.01 to 1 mM ThT was agitated at 750 rpm and 37 °C.

Aggregation studies

Aliquots were withdrawn from the aggregation reaction at various time points, and the amount of fibrils was monitored by measuring the ThT fluorescence at pH 8.0 in 20 mM Tris-HCl buffer. In the assay, 10 μM ThT and 1 μM protein were used. For aggregation reactions carried out in the absence of ThT, 5-μl aliquots were withdrawn and mixed with 495 μl of 20 mM Tris-HCl buffer containing 10 μM ThT. For aggregation reactions carried out in the presence of 1 mM ThT, 5-μl aliquots were withdrawn and mixed with 495 μl of 20 mM Tris-HCl buffer. Then fluorescence was monitored using a Fluoromax-4 spectrofluorometer with the excitation and emission wavelengths set at 440 and 482 nm, respectively.

Quantification of ThT bound to α-synuclein fibrils

To 45 μl of 100 μM fibrils prepared in the absence of ThT, 4.5 μl of 10 mM ThT was added. To 45 μl of 100 μM fibrils prepared in the presence of 1 mM ThT, 4.5 μl of Milli-Q water was added. In both cases, the fibrils were then incubated at 25 °C for 90 min and then spun down at 20,000 × g for 10 min at 9 °C. The pellet was resuspended in 50 μl of aggregation buffer (without ThT). This was repeated four times to remove the free/loosely bound ThT. Fibril concentration was determined using the BCA assay kit (Thermo Scientific), and equal amounts of fibrils formed in the absence and presence of ThT were used to monitor the ThT fluorescence and compare the amount of ThT retained in each fibril sample. The amount of ThT bound to the fibrils was also measured by dissolving the fibrils in 8 M GdnHCl and monitoring the absorbance at 412 nm.

Atomic force microscopy

50 μl of fibrils applied onto freshly cleaved mica were incubated for 3 min. The mica surface was rinsed twice with filtered
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Milli-Q water and dried under vacuum for 45–60 min. The AFM images were acquired using a FastScan Bio (Bruker) instrument and analyzed using WSxM software. The height distribution of the fibrils was obtained by measuring the heights of about 200 fibrils from 10 images of fibrils formed in the presence of ThT and about 1000 fibrils from 36 images of fibrils formed in the absence of ThT. The nature of the periodicity seen in the helical fibrils was determined by measuring 32 fibrils from nine images.

Fourier transform infrared (FTIR) spectroscopy

For FTIR measurements, samples were spun down at 20,000 × g for 10 min at 9 °C, and fibrils were washed three times with aggregation buffer (without EDTA) made in D2O and then resuspended in the same buffer. A thin film was prepared on the diamond crystal by drying 3 μl of sample using N2 gas. FTIR spectra were acquired using a Thermo Nicolet 6700 FT-IR spectrometer. The FTIR spectra of the fibrils were analyzed after background subtraction of the spectrum of the appropriate aggregation buffer.

Peptide mapping

To generate a peptide map of α-synuclein, the protein, dissolved in water, was subjected to online pepsin digestion in 0.05% formic acid using an immobilized pepsin cartridge (Applied Biosystems) at a flow rate of 50 μl/min on a nanoAcquity UPLC (Waters). The eluted peptides were collected using a peptide trap column (C18 reversed-phase chromatography column), and eluted into an analytical C18 reversed-phase chromatography column using a gradient of 3–40% acetonitrile (0.1% formic acid) at a flow rate of 45 μl/min. The peptides were directed to the coupled Synapt G2 HD mass spectrometer (Waters). The peptides were sequenced using the MS/tandem MS (MS²) method followed by analysis with ProteinLynx Global Server software (Waters) and manual inspection.

HDX-MS measurements

Fibrils were prepared as described above. 80 μl of 100 μM fibrils were spun down at 20,000 × g for 20 min at 9 °C. The pellet was resuspended in 20 μl of aggregation buffer. To initiate deuterium labeling, 10 μl of the above sample was diluted into 190 μl of aggregation buffer (without EDTA) made in D2O and incubated at 25 °C for 5 min. After a 5-min pulse, 200 μl of the above sample was mixed with 400 μl of ice-cold quench buffer (0.1 M glycine-HCl, 8.4 M GdnHCl, pH 2.5) and incubated for 1 min on ice to dissolve the fibrils. The samples were desalted using a Sephade G-25 HiTrap desalting column equilibrated with water at pH 2.5 and Akta Basic HPLC. The desalted samples were injected into the HDX module coupled with a nanoACQUITY UPLC system (Waters) for online pepsin digestion using an immobilized pepsin cartridge (Applied Biosystems). Further processing of the sample for mass determination using a Waters Synapt G2 mass spectrometer was carried out as described previously (91).

Peptide masses were calculated from the centroid of the isotopic envelope using MassLynx software, and the shift in the mass of labeled peptide relative to the unlabeled peptide was used to determine the extent of deuterium incorporation. As the sample was in 95% D2O during labeling and was exposed to H2O after dissolution in GdnHCl, control experiments were carried out to correct for back-exchange and forward-exchange. To this end, monomeric α-synuclein was completely deuterated by incubating it in 20 mM sodium phosphate buffer, pH 7.0 (95% D2O) at 25 °C for 5 min. The fully deuterated α-synuclein sample was then processed in exactly the same way as the aggregates. The extent of deuterium incorporation in each peptide, % D, was determined using the following equation,

\[
\%D = \frac{(m(t) - m(0%))/(m(95%) - m(0%)) \times 100}
\]

where \( m(t) \) is the measured centroid mass of the peptide at time point \( t \), \( m(0%) \) is the measured mass of an undeuterated reference sample, and \( m(95%) \) is the measured mass of a fully deuterated reference sample (in 95% D2O) (69).

The percent deuterium incorporation for peptides showing a bimodal distribution was calculated as described previously (14, 91). The centroid mass for each peak was obtained by fitting the bimodal mass distributions to the sum of two Gaussian distributions using OriginPro 8. The % D for each peak was determined using Equation 1.

Cytotoxicity assay

Equal concentrations (1 mM) of ThT were added to the fibrils made under the two different conditions, and the fibrils were incubated at 25 °C for 15 min. The fibrils were spun down at 20,000 × g for 20 min at 9 °C, the supernatant was removed, and the fibrils were washed with an equal volume of autoclaved Milli-Q water. The washing process was repeated three times, and then the fibrils were resuspended in autoclaved Milli-Q water. The concentration of the fibrils was determined using the BCA assay after dissolving the fibrils in 4 M GdnHCl. Equal concentrations (2.5 μM) of fibrils made under the two different conditions were used for the cytotoxicity experiments. Autoclaved Milli-Q water containing 1 μM ThT was used as a buffer control in the cytotoxicity assay. For the assay, HEK-293T cells (ATCC) were cultured in DMEM (Gibco) supplemented with 10% FBS at 37 °C in a 5% CO2-humidified environment. Cells were plated in a 96-well plate at a density of 5000 cells/well to a final volume of 100 μl. After incubation for 24 h, 10 μl of fibrils (25 μM) or ThT only (0.1–5 μM) was added to each well, and the cells were further incubated for 24 h at 37 °C in 5% CO2. The Wst-1 assay kit (Roche) was used to measure the viability of the cells. 9 μl of Wst-1 reagent was added into each well, and the cells were incubated for 2.5 h at 37 °C in 5% CO2. The optical density at 477 nm was measured with a microplate reader (SpectraMax M5). Data were normalized with respect to the data obtained with untreated (control) cells.

Author contributions—H. K., J. S., J. B. U., and P. K. designed the experiments. H. K. performed the experiments. H. K., J. S., and J. B. U. analyzed the results and wrote the manuscript. The final version of the manuscript was approved by all of the authors.
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