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Cryo-EM structure of a novel α-synuclein filament subtype from multiple system atrophy

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Multiple system atrophy (MSA) is a progressive neurodegenerative disease characterized by accumulation of α -synuclein cross- β amyloid filaments in the brain. Previous structural studies of these filaments by cryo-electron microscopy (cryo-EM) revealed three discrete folds distinct from α -synuclein filaments associated with other neurodegenerative diseases. Here, we use cryo-EM to identify a novel, low-populated MSA filament subtype (designated Type I₂) in addition to a predominant class comprising MSA Type II₂ filaments. The 3.3-Å resolution structure of the Type I₂ filament reveals a fold consisting of two asymmetric protofilaments, one of which adopts a novel structure that is chimeric between two previously reported protofilaments. These results further define MSA-specific folds of α -synuclein filaments and have implications for designing MSA diagnostics and therapeutics.

Keywords: cryo-electron microscopy; multiple system atrophy; neurodegeneration; prion; protein aggregation; α -synuclein

 α -Synuclein is an intrinsically disordered, 140–amino acid protein [1] that localizes primarily to the axon terminals of presynaptic neurons, where it participates in membrane binding, vesicle trafficking, and neurotransmitter release [2,3]. The misfolding and aggregation of α -synuclein in the brain are hallmarks of a group of neurodegenerative diseases known as synucleinopathies [4,5], which includes Parkinson's disease and dementia with Lewy bodies. One such disease, multiple system atrophy (MSA), is clinically characterized by cerebellar dysfunction or parkinsonism and neuropathologically characterized by glial cytoplasmic inclusions (GCIs) in oligodendrocytes [6,7]. GCIs contain abundant fibrils that appear to be the result of soluble α -synuclein misfolding into self-templating prion conformations that ultimately misassemble as insoluble amyloid filaments [8–12]. Structural studies on filaments isolated from five MSA patients revealed three distinct filament types (denoted Type I, II₁, and II₂). These findings contrast with Lewy body pathologies, which appear to contain a single α -synuclein filament type, and juvenile-onset synucleinopathy, which contains one α synuclein fold in either a singlet or doublet association [13–15].

The previously determined MSA filament types each consist of two asymmetric protofilaments (PF-A and PF-B) that associate along an extended interface nearly spanning the width of the filament core (Fig. S1) [13].

Abbreviations

ADRC, Alzheimer's Disease Research Center; cryo-EM, cryo-electron microscopy; EMDB, Electron Microscopy Data Bank; FSC, Fourier shell correlation; GCI, glial cytoplasmic inclusions; MSA, multiple system atrophy; PDB, Protein Data Bank; PF, protofilament; RMSD, root-mean-square deviation.

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This is an open access article under the terms of the Creative Commons Attribution License, which permits use, distribution and reproduction in any medium, provided the original work is properly cited. Also common to each type is a central channel flanked by basic residues (Lys43, Lys45, and His50 of each protofilament) with unassigned density present within the channel. However, major conformational differences lie within the protofilaments that compose each mature fibril core. Type I MSA filaments contain two PFs, one PF-IA and one PF-IB. Residues Gly14 to Phe94 are resolved in PF-IA, and Lys21 to Gln99 are resolved in PF-IB. While both PF-IA and PF-IB contain a hairpin motif in the N-terminal region and a three-layered L-shaped motif in the C-terminal region, PF-IA also contains a single-layered L-shaped motif (residues 32–45) joining the N- and C-terminal motifs. Thus, the residues composing the terminal motifs differ in each protofilament, and the fold of the three-layered L-shaped motif also differs in the packing of the inner layer relative to the central layer.

The Type II filament subtypes (II₁ and II₂) each have two protofilaments [13] and contain identical PF-IIA, which span residues 14-94. Comparing PF-IA and PF-IIA, the N-terminal regions spanning residues 14-42 are conformationally identical. However, these protofilaments differ in the C-terminal region, including the conformation of the three-layered L-shaped motif. PF-IIA contains a small channel within the filament core, surrounded by Val52, Thr54, Ala56, Thr59, Glu61, Thr72, Gly73, and Val74, that is not present in PF-IIB because these residues form closer contacts with each other. PF-IIB₁ and PF-IIB₂ include residues 36-99 and contain a three-layered L-shaped motif (residues 47-99). In contrast to PF-IB, these protofilaments lack the N-terminal hairpin and contain a shorter single-layered L-shaped motif (residues 36-46) instead. PF-IIB₁ and PF-IIB₂ are distinguished from each other by a change in a surface loop at the C-terminal end of the ordered core (residues 81–90). The surface loop conformation of PF-IIB₁ is similar to the loop in PF-IB, while PF-IIB₂ has a distinct loop conformation that is shifted approximately 4 A toward the C terminus of the protofilament core. The overall helical symmetry of each MSA filament type is similar, although Type I filaments have a slightly greater twist (-1.42°) compared to Type II₁/II₂ filaments (-1.34°) .

MSA filament types are heterogeneously distributed between different patients [13]. For example, in the original report from Schweighauser et al., two patients had almost exclusively Type I filaments, one patient had almost exclusively Type II filaments, and the remaining two patients had both Type I and II filaments in varying ratios. This heterogeneity suggests that there may be a greater variety of folds than previously reported. In this study, we use cryo-EM to obtain high-resolution (3.2–3.3 Å) structures of filaments purified from tissue of another patient with MSA. While the majority (66%) of the data consists of Type II₂ filaments, our results reveal a novel proto-filament subtype in a subset (5%) of filaments that are related to Type I filaments.

Materials and methods

MSA filament extraction

Deidentified human tissue was obtained from the Massachusetts Alzheimer's Disease Research Center (ADRC). All donors provided consent to donate brains for research purposes in accordance with the standards of the ADRC. This study was exempt from institutional review board approval in accordance with the institutional review board policy of the University of California San Francisco. Filament extraction was performed using a modified protocol from Schweighauser et al. [13]. In summary, 1–2 g of freshly frozen cerebellum tissue from a patient pathologically diagnosed with MSA was homogenized at 20 mL \cdot g⁻¹ of tissue in 10 mM Tris-HCl (pH 7.4), 800 mM NaCl, 1 mM EGTA, and 10% sucrose. N-lauroylsarcosinate (final concentration of 1% w/v) was added to the homogenate and incubated at 37 °C for 30 min. Afterward, homogenates were centrifuged at 10 000 g for 10 min. The supernatant was kept and centrifuged at 100 000 g for 20 min. The pellet was resuspended at 500 μ L·g⁻¹ of frozen tissue in 10 mM Tris-HCl (pH 7.4), 800 mM NaCl, 1 mM EGTA, and 10% sucrose. The suspension was then centrifuged at 3000 g for 5 min. The supernatant was kept and diluted threefold with 50 mM Tris-HCl (pH 7.4), 150 mM NaCl, 10% sucrose, and 2% sarkosyl. This suspension was then centrifuged at 150 000 g for 1 h. The filament-enriched pellet was resuspended in 30 mM Tris-HCl (pH 7.4) using 100 µL·g⁻¹ of frozen tissue.

Cryo-EM sample preparation and data collection

Purified filaments (3 µL) were added to a 200 mesh 1.2/1.3R Au Quantifoil grid coated with a 2-nm thick carbon layer, which was not glow discharged. After 30 s, grids were blotted for 7.5 s at room temperature and 100% humidity using a FEI Vitrobot Mark IV, followed by plunge freezing in liquid ethane. A total of 42 224 super-resolution movies were collected at a nominal magnification of $105\ 000 \times$ (physical pixel size: 0.417 A/pixel) on a Titan Krios (Thermo Fisher Scientific, Waltham, MA) operated at 300 kV and equipped with a K3 direct electron detector and BioQuantum energy filter (Gatan, Inc., Pleasanton, CA) set to a slit width of 20 eV. A defocus range of -0.8 to $-1.8 \ \mu m$ was used with a total exposure time of 2.024 s fractionated into 0.025-s subframes. The total dose for each movie was 46 electrons/Å². Movies were motion corrected using MotionCor2 [16] in Scipion [17] and were

Fourier cropped by a factor of 2 to a final pixel size of 0.834 Å/pixel. Motion-corrected and dose-weighted micrographs were manually curated in Scipion to remove micrographs lacking filaments, those at low resolution, or those with significant ice contamination, resulting in 4392 remaining micrographs.

Cryo-EM image processing

A graphical overview of the data processing workflow is provided in Fig. S2. All image processing was done in RELION 4 [18,19]. Dose-weighted summed micrographs were imported into RELION 4. The contrast transfer function was estimated using CTFFIND-4.1 [20]. Filaments were manually picked, and segments were extracted with a box size of 900 pixels downscaled to 300 pixels, resulting in 257 982 segments. Reference-free 2D classification was used to remove contaminants and segments contributing to straight filaments, resulting in 255 032 remaining segments. These were re-extracted with a box size of 288 pixels without downscaling, followed by another round of reference-free 2D classification that did not filter out more contaminant segments. One round of 3D classification with image alignment was performed on the segments using a reference map consisting of the existing MSA Type I filaments (PDB code: 6XYO), a regularization parameter (T) of 20, and fixing helical parameters to -1.42° twist and 4.76 Å rise. We were unable to resolve Type I₂ filaments by allowing the helical parameters to vary. One class (12 802 segments) corresponded to Type I₂ filaments. One round of 3D auto-refinement was run using this map low-pass filtered to 10 Å, allowing rise and twist parameters to vary. The map was sharpened using the standard postprocessing procedures in RELION. Full statistics are shown in Table S1.

Two out of the 12 classes (57 809 segments) corresponded to Type II₂ filaments with no breaks in the polypeptide density. Segments corresponding to these classes were pooled and subjected to one round of 3D classification without image alignment using a reference map consisting of one of these classes (A3; Fig. S3), a regularization parameter (T) of 20, and fixing helical parameters to -1.34° twist and 4.76 Å rise. The highest resolution class (31 115 segments) was subjected to 3D auto-refinement and postprocessing to give a refined Type II₂ filament map.

Of the 12 classes, 4 classes (110 255 segments) displayed clear separation between protofilaments in the z-direction but modest resolution in the x- and y-directions, suggesting suboptimal alignment. Segments corresponding to these classes were pooled and subjected to one round of 3D classification with image alignment using a reference map consisting of the existing MSA Type II₂ filaments (PDB code: 6XYQ), a regularization parameter (T) of 20, and fixing helical parameters to -1.34° twist and 4.76 Å rise. Most of these segments (110 071) converged into one class

corresponding to Type II_2 filaments, which was not further refined. The remaining classes (74 166 segments) from the initial round of 3D classification were low resolution and excluded from further processing.

To eliminate bias in 3D classification and further verify the distribution of filaments, we generated a 3D initial model ab initio from 2D class averages using RELION's relion_helix_inimodel2d feature. This initial model was used as a reference map for 3D classification using the otherwise same inputs as the first round of 3D classification described above, resulting in classes corresponding to Type II₂ filaments (206 869 segments, 81% of the data) and Type I₂ filaments (15 369 segments, 6% of the data).

Model building, refinement, and analysis

A single strand of the previously solved MSA Type II₂ (PDB code: 6XYQ) or Type I (PDB code: 6XYQ) filament structures was placed in the density using Chimera [21]. Subsequent model building was performed using COOT [22] and ISOLDE [23] followed by refinement in PHENIX [24]. The model was then translated to produce a stack occupying all regions with continuous density on the map. Refinement statistics are shown in Table S1. Data were deposited into the Protein Data Bank (PDB) under accession codes 9CD9 (Type II₂) and 9CDA (Type I₂) and into the Electron Microscopy Data Bank (EMDB) under accession codes EMD-45464 (Type II₂) and EMD-45465 (Type I₂).

Results

Partially purified, sarkosyl-insoluble filaments were isolated from the postmortem cerebellum of a 67-year-old male neuropathologically diagnosed with MSA. Negative-stain transmission electron microscopy on this sample revealed filaments that were morphologically similar to those in previous reports (Fig. S4) [13,25]. After cryo-EM imaging using standard collection methods (Fig. 1A; see 'Materials and methods'), reference-free 2D classification resulted in 2D class averages that were similar to those previously reported [13], exhibiting a crossover distance of approximately 600 Å (Fig. S5). However, it was unclear whether these 2D classes corresponded to a known filament type. After 3D classification and helical reconstruction, we were able to obtain high-resolution structures of two MSA filament types (Fig. 1B). The primary type (Fig. 1C), refined to 3.2 Å and representing 66% of the data (Table S1), was nearly identical to Type II₂ filaments (PDB: 6XYO), with an all-atom root-meansquare deviation (RMSD) of 0.5 Å (Fig. S6). We observed previously reported regions of unknown density, such as non-proteinaceous density within the



Fig. 1. Cryo-EM analysis of *ex vivo* MSA filaments reveals a novel filament subtype: Type I_2 filaments. (A) Representative micrograph. The red arrow indicates an MSA filament. (B) Cross section of two major conformations of MSA filaments, Type I_2 and Type I_2 , identified after 3D classification of filament segments. The resolutions of the final reconstructions and abundance in the data are indicated. The remaining 29% of data consisted of low-resolution classes. (C) Density map and model of Type II_2 filaments. The two constituent protofilaments PF-IIA and PF-IIB₂, along with residues at the termini of the ordered core, are labeled and color coded in pink and purple, respectively. Regions of unmodeled density are also shown. (D) Density map and model of Type I_2 filaments. The two constituents PF-IA₂ and PF-IB, along with residues at the termini of the ordered core, are labeled and color coded in yellow and red, respectively. Regions of unmodeled density are also shown.

central channel flanked by basic residues and a peptide-like density contacting residues Lys80-Val82 of PF-IIA [13]. We were able to resolve an additional residue (Leu100) at the C-terminal region of the PF-IIB₂ ordered core.

In addition to the successful reproduction of the Type II₂ MSA filament, we also resolved a filament exhibiting a conformation of α -synuclein that has not been reported previously. We refer to this low-populated filament subtype as a Type I₂ filament, which represents 5% of the data (Fig. 1B, Fig. S7). The structure of Type I₂ filaments, refined to 3.3-Å resolution, resembles Type I (hereafter called Type I₁) and comprises two protofilaments, PF-IA₂ and PF-IB (Fig. 1D). The ordered core of these protofilaments includes residues Gly14-Gly93 for PF-IA₂ and Lys21-Leu100 for PF-1B. The PF-IB protofilament is common between Type I₁ and Type I₂ filaments, with an RMSD of 0.8 Å to the existing Type I₁ filament model (PDB: 6XYO) [13].

The differences between Type I_1 and Type I_2 filaments exist within the second protofilament: PF-IA for Type I_1 and PF-IA₂ for Type I_2 (Fig. 2A). While the

N-terminal regions comprising residues 14-56 are nearly identical between the two protofilaments, the Cterminal regions of the protofilaments (residues 57-93) adopt distinct conformations. Specifically, this region in PF-IA2 is shifted outwards and away from the interprotofilament interface. This conformational change shifts the register of the hydrophobic intraprotofilament interface containing residues Val74 and Ala76 by two residues and creates a channel within the filament core, resulting in a net loss of hydrophobic interactions, including those made by the Val52 and Thr54 sidechains (Fig. 2B). Additionally, potential interstrand polar contacts between the Glu61 sidechain and Gly73 backbone amide are lost. Interestingly, the C-terminal region of PF-IA₂ is virtually identical to the analogous region in PF-IIA found within Type II MSA filaments, although PF-IIA contains the C-terminal Phe94 residue that was not well resolved in PF-IA₂ (Fig. 2C). Comparing PF-IA and PF-IA₂, the average Ca displacement of residues 57-93 is 4.8 Å but is 0.3 Å between PF-IIA and PF-IA2. Just as in PF-IIA, no additional, non-proteinaceous densities are observed in the C-terminal channel of PF-IA₂.



Fig. 2. Comparison of MSA Type I₂ filaments to previously solved MSA filament structures. (A) Overlay of Type I₁ (PDB: 6XYO) and Type I₂ filaments aligned using PF-IB. The black arrow indicates the direction of the conformational change in the C-terminal region. The average C α displacement of residues 57–93 is 4.8 Å. (B) Comparison of intraprotofilament contacts in the C-terminal regions of PF-IA (Type I₁) and PF-IA₂ (Type I₂). (C) Overlay of Type I₂ and Type II₂ (PDB: 6XYQ) filaments aligned using residues 47–93 of PF-IIA. For clarity, PF-1B and PF-IIB₂ are not shown.

The morphological and helical parameter similarities between Type II₂ and Type I₂ MSA filaments prompted us to inquire whether both types could coexist within the same filament or if the types are instead segregated to individual filaments. Previous cryo-EM analysis of amyloid filaments suggested that multiple morphologies and folds could coexist within the same filaments [14,26]. To address this question, we mapped segments classifying Type I₂ or Type II₂ conformations to the original micrographs (Fig. 3A-C). Due to the 13-fold greater abundance of Type II₂ compared to Type I₂ segments in the dataset, we only analyzed micrographs containing at least one Type I₂ segment, corresponding to 437 micrographs out of 4392 (10%). We found that 71% of the 556 filaments in these micrographs contained segments that classified exclusively or predominantly (90% or higher) to either Type I_2 or Type II_2 (Fig. 3D). The remaining minority of filaments contained a more mixed distribution of types, although Type I₂ tended to predominate (e.g., Fig. 3A). These results suggest that each

MSA type preferentially segregates into separate filaments, though it is possible that a small number of filaments contain mixed types. A precise quantification of this dataset is complicated by the possibility of suboptimal classification, the extent of which is unknown.

Discussion

Cryo-EM studies have been instrumental in determining the structures of disease-associated amyloid fibrils from synucleinopathies and other neurodegenerative diseases [13,14,27–30]. Proteins associated with these diseases adopt numerous polymorphic folds that are associated with specific neurodegenerative diseases [31]. In some diseases, such as MSA, α -synuclein can adopt multiple distinct folds that are heterogeneously distributed between patients [13]. While Type I₁ and II₂ filaments appear to predominate in MSA, the lower-population type II₁ also exists within some patients. Here, we have expanded the spectrum of known MSA filament types to include a fourth subtype, Type I₂ filaments.

Type I₂ filaments share one protofilament fold (PF-IB) with Type I_1 filaments but differ in the PF-IA₂ protofilament, which instead bears similarities to the C-terminal region of the PF-IIA protofilament found in Type II filaments. The chimeric nature of PF-IA₂ suggests that common intermediates may exist in the misfolding and aggregation pathways of α -synuclein in MSA but then diverge, leading to multiple folds. The exact nature of these pathways and the relative contribution of each filament type to disease remains unknown. While MSA has two phenotypes-MSA-C (cerebellar) and MSA-P (parkinsonian) [32]-whether the identity and distribution of filament types correlates to phenotype remains undetermined. Schweighauser et al. previously analyzed 10 tissue samples across three brain regions (putamen, cerebellum, and frontal cortex) and five MSA patients, revealing a heterogeneous distribution of MSA filament types in different samples (Table S2). Even when considering our results, a possible relationship between brain region and MSA filament type cannot be determined due to the small number of samples; this remains an important open question. It also remains unclear whether different filament conformations in MSA may arise independently.

We found that MSA Type I_2 and II_2 filaments tended to segregate to individual filaments, but it is possible that a small number of filaments contain a mixed population of both types. These results contrast with previous analyses of morphologically similar filament folds. Lewy body α -synuclein filaments were found to contain a roughly even distribution of both



Fig. 3. Analysis of MSA Type I_{2^-} and II_{2^-} classified segments within individual filaments. (A–C) After one round of 3D classification, segments corresponding to either type were color coded and mapped to the original micrographs. (D) Histogram of the fraction of segments classified as Type I_2 per filament in micrographs containing at least one I_2 segment.

twisted and untwisted segments, although that analysis is limited by the lack of a structure of the untwisted morphology [14]. Another study investigating ex vivo antibody light chain filaments from a patient with systemic AL amyloidosis found two folds that unambiguously coexist in most filaments and differ only slightly in a surface-exposed region of the filament core [26]. The overall high homology allows for the formation of mixed filaments that maintain the favorable interstrand interactions needed for filament growth and stability. In the case of MSA Types I_2 and II_2 , the folds are mostly dissimilar from each other except for residues 57-93 of PF-A, likely preventing the formation of a stable interface between both types. We attempted to model one Type I₂ filament rung directly adjacent to a Type II₂ filament rung, which were aligned using the homologous residues 57-93 of PF-IIA (Fig. S3). However, the model revealed severe steric clashes in the nonhomologous regions of the filaments. We do note the presence of a small number of filaments (11/556 filaments) containing numerous segments (≥ 10) classified into each type, although the inter-rung steric clashes lend credence to the argument that this results from classification stochasticity instead of the presence of truly mixed filaments.

Overall, these results improve our understanding of the α -synuclein structures underlying MSA and provide a framework for developing therapeutics or diagnostics that bind to MSA filaments with high affinity and selectivity. The existence of multiple MSA α -synuclein folds suggests that these putative molecules should target a conserved region of the filament, as even low-population filament types can predominate if other folds are inhibited. Furthermore, it is possible that Type I_2 filaments may be the major species in certain patients with MSA. Structural analysis on tissue samples from different patients and brain regions will be required to elucidate how MSA filament types are distributed. However, conformation-specific molecules would also be useful for rapidly quantifying each filament type in a tissue sample without needing to resort to cryo-EM [15,33,34]. These molecules should help us continue investigating whether multiple folds may exist in single filaments.

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Conflict of interest

SBP is the founder of Prio Pharma, which did not contribute support to this study. The remaining authors have no conflicts of interest to declare.

Author contributions

NY, DRS, NAP, and GEM designed the research; NY, FC, ET, AM, DAM, and GEM performed experiments; NY, SBP, DRS, NAP, and GEM analyzed the data; NY, DRS, NAP, and GEM wrote the manuscript. All authors read and approved the final manuscript.

Peer review

The peer review history for this article is available at https://www.webofscience.com/api/gateway/wos/peer-review/10.1002/1873-3468.15048.

Data accessibility

Refined atomic models have been deposited in the Protein Data Bank (PDB) under accession numbers 9CD9 (Type II₂) and 9CDA (Type I₂). Corresponding cryo-EM maps have been deposited in the Electron Microscopy Data Bank (EMDB) with accession numbers EMD-45464 (Type II₂) and EMD-45465 (Type I₂). Please address requests for materials to the corresponding author.

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Supporting information

Additional supporting information may be found online in the Supporting Information section at the end of the article.

Fig. S1. Structures of MSA filaments reported by Schweighauser et al.

Fig. S2. Flowchart of MSA filament data processing.

Fig. S3. Model of MSA Type II_2 - I_2 mixed cofilament wherein adjacent Type II_2 and Type I_2 rungs were aligned using residues 57–93 of PF-IIA.

Fig. S4. Representative negative-stain transmission electron micrographs of *ex vivo* MSA filaments.

Fig. S5. Two-dimensional class averages of MSA filaments extracted at two different box sizes.

Fig. S6. Characterization of MSA Type II_2 filaments in the dataset.

Fig. S7. Characterization of MSA Type I_2 filaments in the dataset.

 Table S1. Cryo-EM data collection and model refinement statistics.

 Table S2. MSA filament type distribution across different brain regions in different cases.