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The 3D structure of lipidic fibrils of α-synuclein

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Abstract:

 α -synuclein (α Syn) is abundant in neurons, but its misfolding and abnormal fibrillization are associated with severe neurodegenerative diseases. Although interactions between α Syn and phospholipid membranes are relevant during α Syn fibril assembly, insights into the interactions of α Syn fibrils with phospholipids have remained elusive. Here, we present six novel polymorphic atomic structures of α Syn fibrils aggregated in the presence of phospholipids. The structures reveal that phospholipids favor a novel protofilament fold, mediate an unusual arrangement of protofilaments, and fill the central cavities between the protofilaments. These findings provide a structural rationale for fibril-induced lipid extraction, a mechanism likely to be involved in the development of α -synucleinopathies.

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15 Main Text:

The pathological causes for α -synucleinopathies, including Parkinson's disease (PD), multiple system atrophy, and dementia with Lewy bodies (DLB), are largely unknown, but accumulating evidence suggests that these diseases are related to the misfolding and aggregation of α -synuclein (α Syn) into amyloid fibrils (*1-3*). A pathological hallmark for PD and DLB is the presence of large neuronal inclusions called Lewy bodies (LB) and α Syn fibrils have been identified as one constituent of LB (*4-6*). Several studies on the composition of LB extracted postmortem from brain tissue of PD-confirmed patients revealed that lipids and membranous organelles are a significant component of LB (*7-9*). Although molecular interactions between α Syn and lipids have been discussed as relevant for PD pathology for decades (*10*), detailed structural characterization is desirable. Here, we report cryo-electron microscopy (cryo-EM) structures of α Syn fibrils grown in the presence of phospholipid vesicles and used molecular dynamics (MD) simulations together with published Nuclear Magnetic Resonance (NMR) data confirming specific lipid-fibril interactions.

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De novo aggregation in a 1:1 mixture of 1-palmitoyl-2-oleoyl-*sn*-glycero-3-phosphate (POPA) and 1-palmitoyl-2-oleoyl-*sn*-glycero-3-phosphocholine (POPC) with a 5:1 lipid to protein ratio was induced by sonication under protein misfolding cyclic amplification conditions and

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completed under gentle orbital shaking to elongate the fibrils (11) (Fig. S1). The lipids were selected as a simplification of negatively charged synaptic vesicles (12) to recapitulate the established binding of monomeric α Syn to lipids via its N-terminus (13, 14). We confirmed the presence of α Syn fibrils by cryo-EM screening and selected three preparations of α Syn fibrils for which independent image data sets were collected. Extensive classification during 3D reconstruction revealed three dominant protofilament folds (*L*1, *L*2, and *L*3) that form in total six different fibrils. Short sonication periods favor fibrils of the *L*1 fold (Fig. 1A-E), while extensive sonication is needed to yield larger populations of *L*2 and *L*3 fibrils (Fig. 2A-F, S1, S2).

The *L1* fibrils were determined to a resolution of 3.2 Å for *L1A* (Fig. S3A) and 3.0 Å for *L1B* and *L1C* (Fig. S3B, C), allowing to accurately model residues M1-Q99 (Fig. S4-S6). Each monomer in the *L1* protofilament comprises ten β -strands (β 1 to β 10) with nine connecting loops. Strands β 2 and β 8 form the tightly packed core, stabilized by a predominant hydrophobic steric zipper (*15*).

While the *L1A* fibril consists of a single *L1* protofilament (Fig. 1B), *L1B* and *L1C* fibrils are
composed of two identical and intertwined *L1* protofilaments (Fig. 1C, D). The *L1B* and *L1C* fibrils differ in their protofilament interfaces. In the *L1B* fibril, both protofilaments are related by an approximate 2₁ screw symmetry and the protofilaments are tilted by ~37° to each other (Fig. S7). Protofilament dimerization, mediated by hydrophobic interactions between residues M1 and V40 as well as G41, accommodates a wide cavity in the protofilament interface. The protofilaments in the *L1C* fibril, on the other hand, are related by C₂ symmetry and ionic interactions between residues K45 and E46 form the inter-protofilament interface.

The *L2A* fibril was determined to a resolution of 2.7 Å (**Fig. 2B, S3D, S8**), 3.1 Å for *L2B* (**Fig. 2C, S3E, S9**), and 2.8 Å for *L3A* (**Fig. 2D, S3F, S10**). The *L2* fold is similar but not identical to α Syn "polymorph 2" (PDB-ID: 6SST), firstly reported by Guerrero-Ferreira *et al.* (*16*) (**Fig. 2E, S11A, B**). The main structural difference between the folds is a shift of strand β 5 by about 10 Å with respect to strands β 8 and β 1. On the other hand, the *L3* fold is similar to the fold determined for the E46K variant (PDB-ID: 6UFR) (*17*) (**Fig. 2F, S11C**). Here, the main structural differences are shifts of strands β 4- β 7 relative to their position in 6UFR and the presence of the N-terminal 14GVVAAA₁₉, which forms another β -strand neighboring β 7.

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The *L2A* fibril is composed of three identical *L2* protofilaments related by a C_3 symmetry. Interestingly, the three protofilaments are separated by ~20 Å and thus show no direct protein-

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protein contacts (Fig. 2B). In the L2B fibril, two identical but asymmetrically arranged L2 protofilaments form the mature fibril and the L3A fibril reveals a similar protofilament orientation (Fig. 2C-F). In both L2B and L3A fibrils, the helical axes of the two protofilaments point in opposite directions, which leads to an identical structure and elongation kinetics of both fibril ends. These structures belong to a symmetry class of amyloid fibrils which has been postulated (18) but, so far, not been observed experimentally.

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For all fibril structures, the cross-section cryo-EM maps reveal additional ring- and rodshaped densities at the fibril surface (Fig. 1F-H, 2G-I, 3A), which together are reminiscent of the cross-section of phospholipid micelles. In addition, previous ssNMR experiments identified residues of α Syn that bind to phospholipids (11), and most of those are neighboring these nonproteinaceous densities (Fig. 3A, B, D, F). We, therefore, assign the micelle-like cross-section densities to phospholipids bound to the fibril surfaces and refer to these protein-lipid aggregates as lipidic fibrils.

To validate this interpretation, we performed MD simulations of free lipid diffusion in the 15 presence of the aSyn fibril structures determined here. Comparable simulations have successfully identified binding sites for biomarkers on other amyloid fibrils (19, 20). The initially randomly distributed lipid molecules associate towards micelle-like aggregates (Movie S1) and subsequently bind to predominantly hydrophobic areas on the fibril surface (Fig. S12-S18). The conversion of the SUVs used for the preparation of the lipidic fibrils to such small lipid aggregates upon fibril formation was confirmed by ³¹P ssNMR (Fig. S19). We calculated average density grids, showing 20 the probability distribution of the lipids relative to the α Syn fibrils, averaged over multiple independent MD trajectories. Comparison to the cryo-EM cross-sections shows that the average lipid density from MD simulations almost perfectly matches the micelle-like densities in the cryo-EM cross-sections (Fig. 3C, E, G). Additionally, the averaged lipid density from MD simulations 25 also reflects the periodical arrangement of the rod-shaped densities along the helical axis seen in the cryo-EM maps. Consequently, the MD data corroborates our assumption that the nonproteinaceous densities in the cryo-EM cross-sections likely are phospholipids bound to the fibril surface. In particular, the ring-shaped densities in the cryo-EM cross-sections likely constitute the lipid head groups, while the rod-shaped densities can be attributed to the lipid acyl chains. Fig. 4 30 shows a POPC molecule modeled into the most well-defined of these densities.

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The central cavity in the *L1B* fibril is occupied by lipids, with their head groups pointing towards charged residues K6, K21, and E20 and their acyl chains pointing towards the center. The cavity also harbors two areas likely occupied by chloride ions (Cl⁻), bridging interactions of positively charged residues K6 and K21 with the choline moiety of the lipids (Fig. 3D, E). For the L1C fibril, MD simulations revealed a high probability for Cl⁻ ions in the hydrophilic interface involving residues K43, K45, and E57 (Fig. 3F, G).

A striking feature of the L2A fibril is the bridging of lipid molecules that span the ~ 20 Å gap between the protofilaments. Furthermore, head group densities of these lipids partially overlap with densities for Cl⁻ (Fig. 3A, H). Hence, the negatively charged phosphate groups, the positively charged choline moiety, and the Cl⁻ ions together form the bridge between K34 and K43 in the individual protofilaments by forming a well-ordered interaction network. This bridging is also seen for L2B and L3A fibrils, where Cl⁻ ions colocalize with head groups at the interface between K43 and K45 (Fig. 3A, I, J).

Although the L2 and L3 folds appear reminiscent of reported structures (16, 17), fibril-lipid 15 interactions favor novel quaternary protofilament arrangements. In the L2A fibril, lipid-mediated interactions seem essential by connecting the neighboring protofilaments. Lipid-mediated interactions might also be responsible for the protofilaments pointing in opposite directions in the L2B and L3A fibrils, as in this configuration, the segment 34KEGVLYVGSK45 in both protofilaments is in contact with the same phospholipid micelle.

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While micelle-like lipid arrangements at the fibril surface can potentially also result from diffusion of lipid molecules after fibril formation, lipids in the central cavity of L2A seem less likely to originate from this process, as lipids mediate the interaction between the protofilaments, suggesting the presence of lipids already during fibril assembly. It is curious to note that the segment 35EGVLYV40 in the lipidic L1 fibrils or 34KEGVLYVGSK45 in lipidic L2 and L3 fibrils are in contact with lipids, which suggests that this stretch of residues could play an important role throughout fibril formation. Indeed, residues within this sequence, such as Y39, have previously been identified to play a crucial role in lipid binding, aggregation kinetics, and function (21, 22).

Although the mechanisms of α Syn aggregation and toxicity *in vivo* are still under debate (23, 24), disruption of intracellular vesicles is one potential mechanism for the cellular toxicity mediated by α Syn fibrils (25) as well as oligomers (26). Reynolds *et al.* proposed a mechanism in which the abnormal aggregation of aSyn is linked to continuous lipid extraction mediated by the

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growing aggregates, eventually leading to membrane disruption (27). The finding of direct fibrillipids interactions due to their lipid-associated aggregation may provide the structural basis for the proposed lipid extraction mechanism (27). In addition, the lipid-coated fibrils reported here give a structural rationale to the previously suggested lipid co-aggregation with α Syn fibrils (28).

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In recent years, a growing number of *ex vivo* cryo-EM fibril structures have been discovered that are characteristic of different diseases (29-31). That none of these so far are lipidic fibrils might be explained by the use of detergent during the isolation of fibrils from patient tissue.

In conclusion, we report six fibril structures of α Syn, revealing how lipid molecules bind directly to the fibril surface. Insights obtained from these lipidic fibrils emphasize that studying α Syn aggregates in the presence of lipids is relevant for understanding the molecular basis of α synucleinopathies. Furthermore, modulation of fibril-lipid interactions may also provide a promising strategy in searching for novel therapeutic interventions. bioRxiv preprint doi: https://doi.org/10.1101/2022.03.02.481946; this version posted March 7, 2022. The copyright holder for this preprint (which was not certified by peer review) is the author/funder, who has granted bioRxiv a license to display the preprint in perpetuity. It is made available under a Granus Control of the preprint in perpetuity. It is made

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Author contributions:

15	Conceptualization: CG, GFS
	Protein expression and purification: SB
	Preparation of fibril samples: LeA
	ssNMR data acquisition and interpretation: LeA, EEN, LBA
	Cryo-EM grid preparation and data collection: CD
20	Image processing, reconstruction and model building: BF
	Visualization: BF, LeA
	Project administration: BF, LeA, CG, GFS
	Supervision: CG, GFS
	Writing – original draft: BF, LeA, CG, GFS

25 Writing – review & editing: All authors

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Competing interests:

Authors declare that they have no competing interests.

Data and materials availability:

Cryo-EM maps and motion-corrected micrographs have been deposited in the Electron Microscopy Data bank (EMDB) under the accession numbers EMD-XXX (*L1A*), EMD-XXX (*L1B*), EMD-XXX (*L1C*), EMD-XXX (*L2A*), EMD-XXX (*L2B*), and EMD-XXX (*L3A*). The corresponding atomic models have been deposited in the Protein Data Bank (PDB) under the accession numbers: XXXX (*L1A*), XXXX (*L1B*), XXXX (*L1C*), XXXX (*L2A*), XXXX (*L2B*), and XXXX (*L3A*). Additional data will be available from the corresponding authors upon reasonable request.

Supplementary Materials:

Materials and Methods

15 Figs. S1 to S14 Tables S1 to S2 Movies S1

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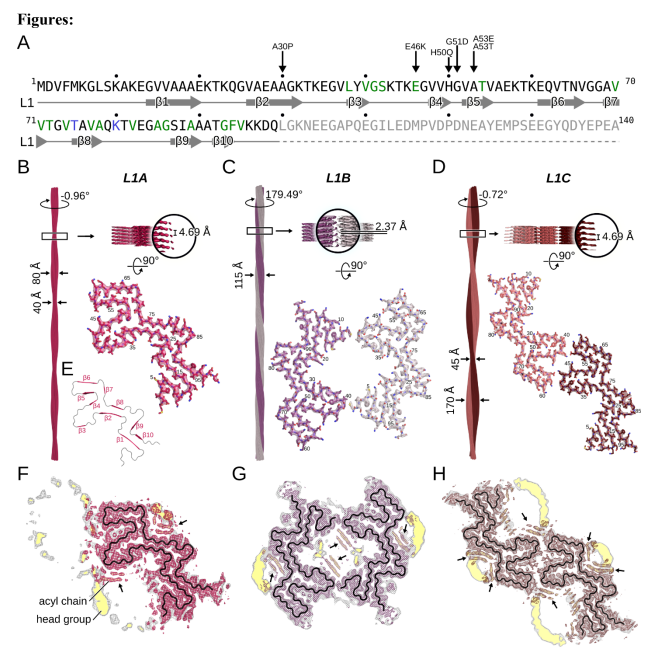


Fig. 1. Cryo-EM structures of the *L1* aSyn fibrils.

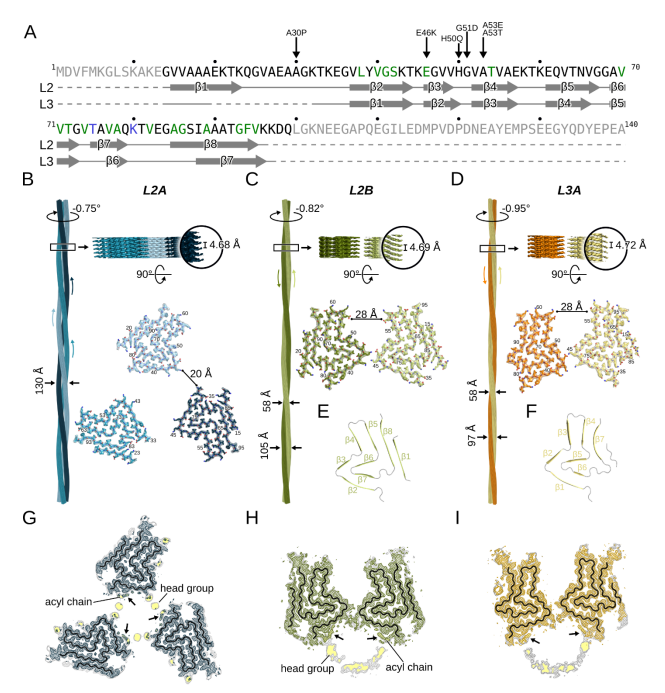
A: Sequence and secondary structure of human aSyn. Familial PD mutation sites (black arrow) localized within the lipid binding N-terminal region (residues 1–60) (32). Green-colored residues bind to the lipid acyl chain, blue to the choline moiety (11), and gray were not resolved. **B-D:** Cryo-EM structures of L1A (**B**), L1B (**C**), and L1C (**D**) fibrils (protofilaments colored differently). The atomic models are shown as sticks. Labels denote the fibril width, the helical twist and rise, and residue numbers. The density maps in the lower panels are displayed using the carve feature in PyMOL at a distance of 2 Å. **E**: Backbone of the L1 fibrils with the $\beta 1 - \beta 10$ colored magenta and loops in gray. **F-H:** Overlay of a sharpened high-resolution map shown in magenta (**F**), purple (**G**), and brown (**H**) and an unsharpened, 4.5 Å low-pass filtered density in gray. The backbone is

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shown as a black ribbon. Densities highlighted with a yellow background are reminiscent of lipid micelles.



5 Fig. 2. Cryo-EM structures of *L2* and *L3* αSyn fibrils.

A: See Fig. 1A for details. B-D: Cryo-EM structures of L2A (B), L2B (C), and L3A (D) fibrils (protofilaments colored differently). The atomic models are shown as sticks. Labels denote the fibril width, the helical twist and rise, and residue numbers. The density maps in the lower panels are displayed using the carve feature in PyMOL at a distance of 2 Å. E, F: Backbone trace of the L2 (E) and L3 (F) fibrils with the $\beta 1 - \beta 10$ colored green or yellow and loops in gray. G-I: Overlay of a sharpened high-resolution map shown in blue (G), green (H), and orange (I) and an

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unsharpened, 4.5 Å low-pass filtered density is shown in gray. The backbone of the model is shown as black ribbon. Unsharpened densities highlighted with a yellow background are reminiscent of lipid micelles.

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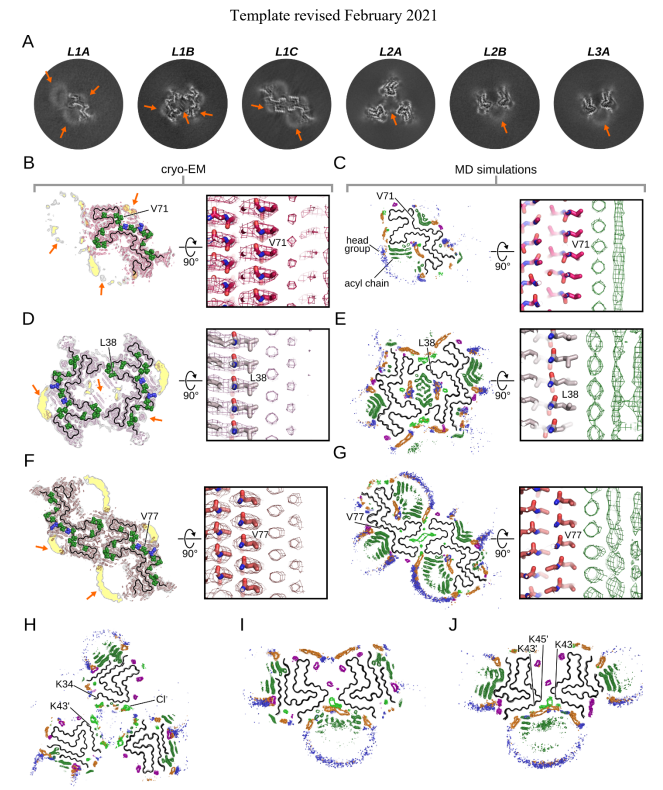


Fig. 3. Fibril-lipid interactions.

A: Central slice of the unsharpened refined cryo-EM maps. **B**, **D**, **F**: Superimposition of the reconstructed cryo-EM maps and the atomic model. Sharpened, high-resolution maps are shown in magenta (**B**, L1A), violet (**D**, L1B), and red (**F**, L1C). Unsharpened, 4.5 Å low-pass filtered density is shown in gray. The backbone of the model is shown as black ribbon, with residues binding to the acyl chain (green) or choline moiety (blue) of phospholipids shown as spheres (11).

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In A+D, the arrows highlight non-fibrillar densities. C, E, G: The grids indicate the probability density of the lipid acyl chain (dark green), phosphate (orange), and the choline nitrogen (blue), and sodium (purple), and chloride (light green) ions throughout MD simulations of the L1A (C), L1B, (E), and L1C (G) fibril. In B-G, the right panels show a close-up view visualizing the ordered packing of the lipid molecules along the helical axis. H-J: Probability density of the lipids throughout MD simulations of lipid diffusion for the L2A (H), L2B, (I), and L3A (J) fibril.

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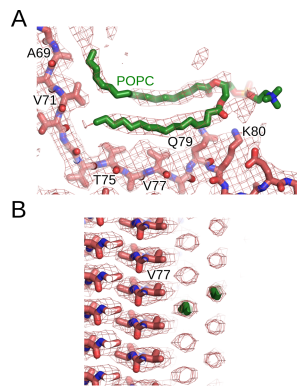


Fig. 4. A fibril-lipid binding-mode model.

One POPC molecule (green sticks) was modeled into the sharpened map of L1C and is shown in a view along the fibril axis (A) and perpendicular to the fibril axis (B).