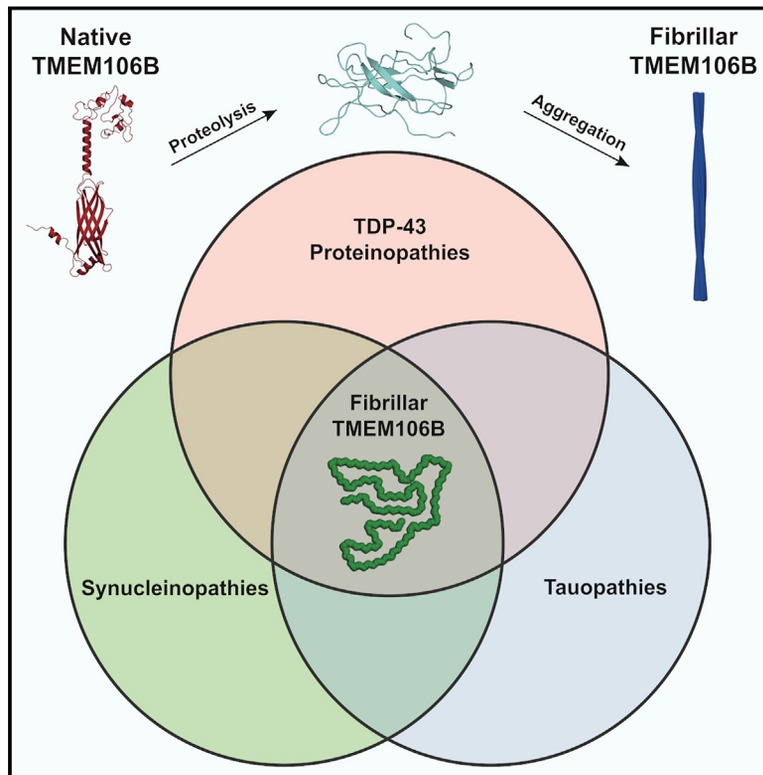


Homotypic fibrillization of TMEM106B across diverse neurodegenerative diseases

Graphical abstract



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In brief

Cryo-EM and mass spectrometry-based proteomics of insoluble amyloid fibrils derived from postmortem human brains afflicted with diverse neurodegenerative diseases reveals widespread fibrillization of an endolysosomal membrane protein, TMEM106B, pointing toward a potentially pathogenic commonality between distinct proteinopathies.

Highlights

- Cryo-EM structures of brain-derived TMEM106B fibrils from neurodegenerative diseases
- Endolysosomal membrane protein TMEM106B C-terminal fragment forms amyloid fibrils
- TMEM106B fibrillization is widespread among diverse neurodegenerative proteinopathies
- Identification of fibrillization pathway potentially implicated in neurodegeneration

Article

Homotypic fibrillization of TMEM106B across diverse neurodegenerative diseases

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SUMMARY

Misfolding and aggregation of disease-specific proteins, resulting in the formation of filamentous cellular inclusions, is a hallmark of neurodegenerative disease with characteristic filament structures, or conformers, defining each proteinopathy. Here we show that a previously unsolved amyloid fibril composed of a 135 amino acid C-terminal fragment of TMEM106B is a common finding in distinct human neurodegenerative diseases, including cases characterized by abnormal aggregation of TDP-43, tau, or α -synuclein protein. A combination of cryoelectron microscopy and mass spectrometry was used to solve the structures of TMEM106B fibrils at a resolution of 2.7 Å from postmortem human brain tissue afflicted with frontotemporal lobar degeneration with TDP-43 pathology (FTLD-TDP, $n = 8$), progressive supranuclear palsy (PSP, $n = 2$), or dementia with Lewy bodies (DLB, $n = 1$). The commonality of abundant amyloid fibrils composed of TMEM106B, a lysosomal/endosomal protein, to a broad range of debilitating human disorders indicates a shared fibrillization pathway that may initiate or accelerate neurodegeneration.

INTRODUCTION

Neurodegenerative proteinopathies are characterized by the deposition of filamentous protein aggregates in neurons and/or glia (Jucker and Walker, 2013). The transactive response DNA-binding protein-43 (TDP-43), the microtubule-associated phosphoprotein tau, and α -synuclein protein can each misfold and accrue intracellularly into tangled filamentous inclusions manifesting in neurodegenerative diseases known collectively as TDP-43 proteinopathies, tauopathies, and synucleinopathies, respectively (Lee et al., 2001). The structures of TDP-43 and tau filaments in frontotemporal lobar degeneration (FTLD) (Arakhamia et al.,

2020; Arseni et al., 2022; Shi et al., 2021; Zhang et al., 2020) and α -synuclein fibrils in multiple system atrophy (MSA) (Schweighauser et al., 2020) have recently been reported. What has emerged from these studies is that each disease has a homotypic molecular fold, or conformer, characteristic of the underlying neuropathology (Arakhamia et al., 2020; Falcon et al., 2018, 2019; Fitzpatrick et al., 2017; Zhang et al., 2020). The evidence for the “one conformer per disease” paradigm is growing, with the presence of unknown buried cofactors (Arakhamia et al., 2020; Falcon et al., 2019; Zhang et al., 2020) and diverse patterns of posttranslational modification (Arakhamia et al., 2020; Kametani et al., 2020) mediating the structural diversity of fibrillar polymorphs.

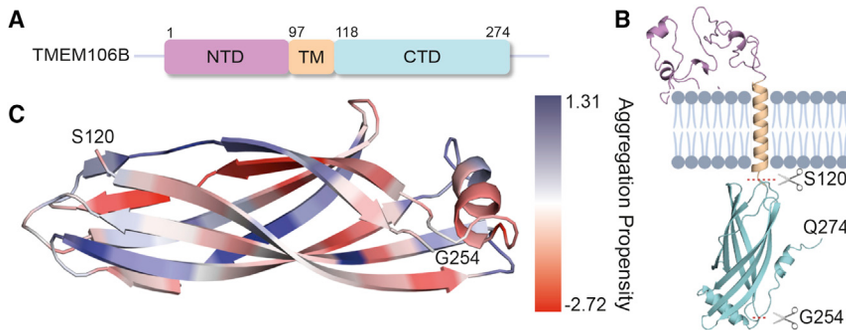


Figure 1. Overview of native TMEM106B

(A) Schematic view of TMEM106B with its N-terminal domain (NTD), transmembrane domain (TM), and C-terminal domain (CTD).

(B) A predicted structure of TMEM106B from ROBETTA (Kim et al., 2004) colored with the same scheme as (A), indicating sites at which the C-terminal domain may get cleaved.

(C) Aggregation propensity mapped onto a predicted structure of TMEM106B(120–254) from AlphaFold (Jumper et al., 2021) (positive values in blue indicate soluble regions and negative values in red correspond to aggregation-prone regions).

Here, using a combination of cryoelectron microscopy (cryo-EM) and mass spectrometry (MS), we show that a previously unsolved amyloid fibril found in cases representing a variety of neurodegenerative conditions is composed of a 135 amino acid C-terminal fragment of TMEM106B, a known risk gene for FTLN-TDP and aging (Feng et al., 2021; Van Deerlin et al., 2010). The fibrillization of TMEM106B into identical fibrillar structures in a wide range of sporadic or genetic TDP-43 proteinopathies (FTLN-TDP), a tauopathy (PSP), and a synucleinopathy (DLB) points toward a commonality between these diverse neurodegenerative diseases. This suggests that the formation of amyloid fibrils composed of TMEM106B, a lysosomal/endosomal protein, may contribute to pathogenicity via a loss or gain of function.

RESULTS

Structures of TMEM106B amyloid fibrils in diverse neurodegenerative diseases

TMEM106B is a 274 amino acid type II transmembrane protein (Figure 1A) that under physiological conditions spans the lysosomal and endosomal membranes of neurons, glia, endothelial cells, and pericytes (Figure 1B) (Busch et al., 2013; Lang et al., 2012). Current knowledge points to a physiological role for TMEM106B in the proper sorting and transport of endosomes and lysosomes in these various cell types (Feng et al., 2020; Lang et al., 2012; Lüningschrör et al., 2020; Rademakers et al., 2021; Schwenk et al., 2014; Stroobants et al., 2021; Zhou et al., 2020). TMEM106B consists of a C-terminal domain that projects in the lysosomal lumen, a single-pass transmembrane domain, and an N-terminal domain that extends into the cytosol (Figure 1B) (Lang et al., 2012).

The N terminus, TMEM106B(1–96) (Figures 1A and 1B), is involved in lysosomal and endosomal trafficking via interactions with MAP6 and can interact with itself as well as with TMEM106C to form homo- and hetero-multimers at the surface of the lysosome (Schwenk et al., 2014; Stagi et al., 2014). The transmembrane domain, TMEM106B(97–117), is a single-pass α -helix (Figures 1A and 1B). Although there is *in vitro* evidence that the luminal C terminus, TMEM106B(118–274), may undergo proteolytic cleavage to release a C-terminal fragment (Figure 1B) (Brady et al., 2014), this has not been confirmed *in vivo* under physiological or disease conditions. Importantly, the 135 amino acid C-terminal fragment that we identified in fibrils from FTLN-TDP, PSP, and DLB cases, TMEM106B(120–254), is predicted

to have a high β strand content and to be prone to aggregation (Figure 1C) (Jumper et al., 2021). TMEM106B has many known posttranslational modifications, with a heavily glycosylated C terminus (Lang et al., 2012). Here, we show that a truncated C-terminal fragment, TMEM106B(120–254), aggregates into amyloid fibrils in diverse genetic and sporadic proteinopathies and present experimentally determined atomic models of the fibrils extracted from 11 postmortem human brains.

Protein filaments were isolated from insoluble protein fractions derived from human postmortem brain tissue representing a number of neurodegenerative disorders. Remarkably, cryo-EM 3D reconstructions of fibrils extracted from five FTLN-TDP type A cases (four with different *GRN* mutations and one sporadic case), two FTLN-TDP type B cases, one FTLN-TDP type C case, two cases of PSP (a 4R tauopathy), and one case of DLB (Table S1; Figures S1 and S2), found two common fibril subtypes (Figures 2 and S3). In each of these cases, there were wide, 2-fold symmetric, twisted ribbons with a helical pitch of $\sim 2,100$ Å, a maximum width of ~ 260 Å, and a minimum width of ~ 120 Å, which we designate “doublet” fibrils (Figures 2 and S3A). There were also narrow, twisted rods displaying a regular helical pitch of $\sim 2,100$ Å and an approximately uniform width of ~ 125 Å, which we refer to as “singlet” fibrils (Figures 2 and S3B). A structural variation of the singlet fibril was found in one case (Figure 2, FTLN-TDP case 7/type B case 2), with a shorter helical pitch of $\sim 1,250$ Å (Figure S3C). The ratio of singlet-to-doublet fibril populations were found to vary between 1:2 and 2:1 for the majority of cases. All density maps show clear inter- β sheet separation perpendicular to the long axis of the fibrils, with separation of β strands discernible at resolutions better than 4.5 Å (Figure 2) (Fitzpatrick et al., 2017). The backbone trace of all cross-sections is clear (Falcon et al., 2018) and shows a generic fibrillar fold common to TDP-43 proteinopathies (FTLN-TDP), a tauopathy (PSP), and a synucleinopathy (DLB) (Figure 2).

Density maps of the unknown singlet and doublet fibril subtypes were determined to resolutions of 3.0 Å and 2.7 Å, respectively (Figures 2–5, and S4; Tables S2 and S3). To identify these unknown fibrils, MS-based proteomics of fractionated tissue from FTLN-TDP case 1 (Table S5) was used to identify 600 potential protein candidates. The well-resolved sidechain density of the doublet fibril was mapped to this reduced proteome using cryoID (Ho et al., 2020) and *findMySequence* (Chojnowski et al., 2021) software, which unambiguously identified an amino acid sequence corresponding to residues 120–254 of TMEM106B (see STAR Methods). Fibrils formed by

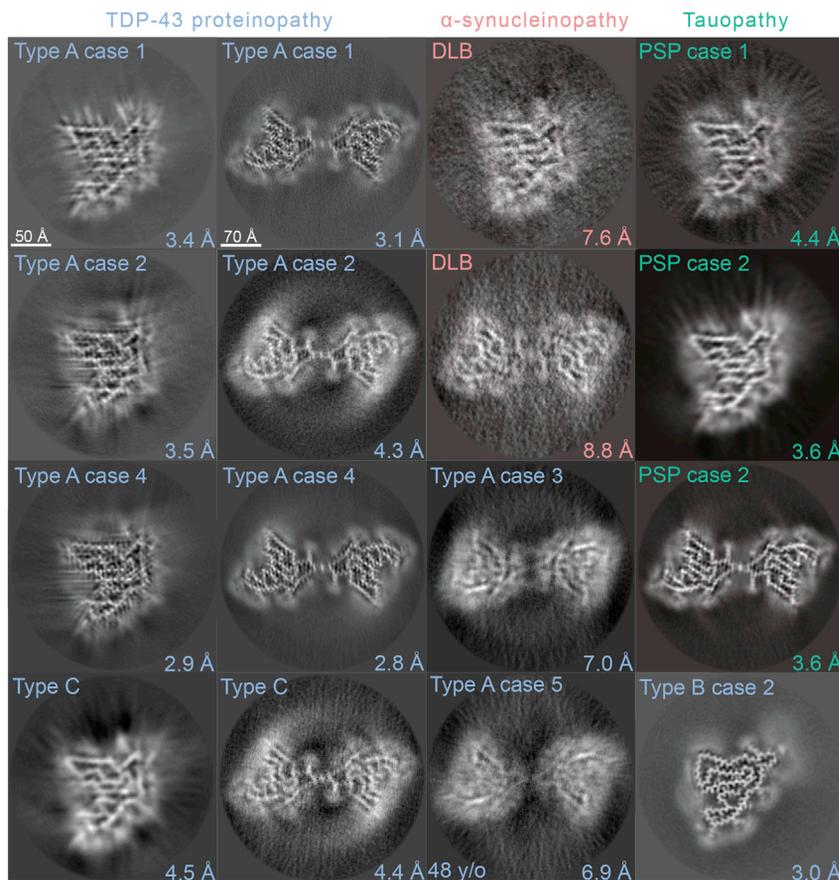


Figure 2. Cryo-EM reconstructions of TMEM106B fibrils

Cross-sections (10 z-slice average) of the TMEM106B(120–254) singlet and doublet fibril unsharpened density maps from eight cases of FTLD-TDP, two cases of PSP, and one case of DLB. See also [Figures S1–S3](#) and [S6](#) and [Tables S2](#) and [S3](#).

Starting from the N terminus, strands β 1– β 3 forms a Z shape, with the tight junction Y132 forming a putative hydrogen bond with N154 that effectively hold neighboring strands β 5 and β 6 in place. A closely packed β helix is formed by strands β 3, β 4, and β 5. Following a gentle turn at N154, strand β 6 is interrupted by E161, which forms a salt bridge with H239 that contributes to the positioning of strand β 18. A short β 7 strand leads into a long β 8 strand that forms a short antiparallel sheet-sheet interface with strand β 17. Strands β 8– β 10 form a large, enclosed hydrophobic cavity, involving residues I176, A179, L181, I184, and I186. The cavity is likely to contain small cofactors as detached densities are observed frequently in sharpened maps. Strand β 11 marks the beginning of the C-terminal three-layer fold forming a tight convergence point with strand β 16 and loops V230–V232 through

TMEM106B(120–154) exhibit ultrastructural polymorphism with a common protofilament ([Figure 4A](#)) existing as either a singlet fibril ([Figure 5A](#) and [5C](#)) or in a 2-fold symmetrical juxtaposition as a doublet fibril ([Figure 3A](#)). The inter-protofilament interface of the doublet fibrils is mediated by an apparently non-proteinaceous anionic cofactor that binds to the sidechains of residues K178 and R180 of each protofilament with near-stoichiometric occupancy ([Figure 3B](#)).

The protofilament core consists of residues 120–254, derived from the luminal topological domain ([Figures 4A](#) and [S5A](#)), which is distinct from the stable N-terminally cleaved fragments from the cytoplasmic topological domain that have previously been reported ([Brady et al., 2014](#)). The 135 residues adopt a C-shaped protofilament with the N terminus embedded in the center, surrounded by a long connecting strand that meanders back on itself to form a closely packed, three-layered C-terminal fold ([Figures 4A](#) and [S5A](#)). In total, 19 β strands were identified (β 1– β 19, [Figures 5D](#), [S5A](#), and [S5C](#)) and found to be linked with either short loops (Q138–R140, T166–A167, G177–A179, G188–P189, V199–P200, E206–S208, C214–T215, K220–N223, Q229–V232, F237–G238, Q245–V251), a gentle turn (N154), or tight single residue turns (G127, Y132, N145, E161, N183, Q194). Diverse sidechain-sidechain interactions facilitate the packing or positioning of β strands ([Figures 4A](#), [S5A](#), and [S5C](#)).

the hydrophobic residues I186, L190, M227, and V230. Hydrophobic clustering also locks strands β 12, β 13 in place with strand β 15 nestled in between through small, apolar residues V199, I217, and I219. A bulky, aromatic β 14 strand may be stabilized by π - π interactions between Y209 and Y211, and between Y211 and F213, as the density indicates that two rotamers are accessible to Y211. Strand β 15 packs against the extreme C terminus through the formation of a disulfide bond between C214 and C253, connecting to strand β 16 via the loop K220–N223 (a potential cation- π interaction is formed by K220 and H222). N223 also plays an important role in facilitating the interdigitation of sidechains between strands β 16 and β 18 by forming a hydrogen bond with E246. The potential strain is relieved by forming an L-shape pin turn with the short β 17 strand. A multi-sheet, cross- β interface between strands β 16 \rightarrow β 18 \rightarrow β 1 cements the center of the fold with a short β 19 strand closing the protofilament structure at the potential cleavage site G254. The disulfide bond between C214 and C253 ([Figure 4B](#)) stabilizes the fibril structure. Ultrastructural polymorphism of the singlet and doublet fibrils is mediated by a non-proteinaceous density nestled between K178 and R180, connecting the two protofilaments of the doublet fibril ([Figures 3](#) and [S5C](#)). In cross-section, this measures approximately $9 \text{ \AA} \times 4 \text{ \AA}$ and is presumably anionic, given that it is bound to positively charged sidechains ([Figures 3B](#) and [S5C](#)).

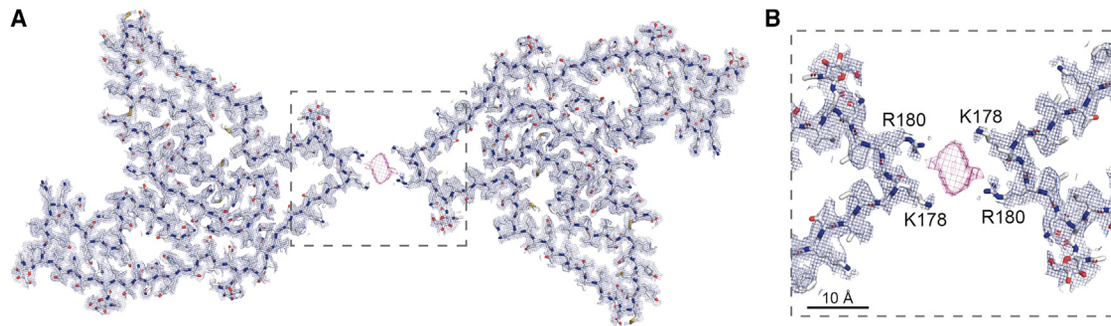


Figure 3. Cryo-EM structure of a TMEM106B doublet fibril

Cryo-EM density (mesh) and atomic model (sticks) of (A) a TMEM106B doublet fibril and (B) the interface between the two protofilaments in the TMEM106B doublet fibril mediated by a non-proteinaceous, anionic cofactor (purple mesh) that binds to the sidechains of residues K178 and R180 of each protofilament. See also [Figure S5](#) and [Table S2](#).

Structural role of posttranslational modifications in TMEM106B amyloid fibrils

Post-translational modifications were preserved for the filament sample and allowed assignment of glycans in the atomic model that match with previous studies of the native protein. Native TMEM106B has six potential N-glycosylation sites all located on the luminal domain, with the first three on residues N145, N151, and N164 (N1–N3) being non-complex glycosylation and the last three on N182, N183, and N256 (N4–N6) being complex N-glycosylation ([Lang et al., 2012](#)). In the protofilament structure, we found that these N-glycosylations were largely retained as densities consistent with NAG were identified at residues N183 ([Figure 4C](#)), N164 ([Figure 4D](#)), N151 ([Figure 4E](#)), and N145 ([Figure 4F](#)). All prominent non-proteinaceous densities are shown schematically in [Figures S5A–S5C](#).

The structure of TMEM106B(182–185) is significant as this region has a disease-risk associated polymorphism (T185S, [Table S1](#)). It includes a potentially phosphorylated T185, a heavily glycosylated N183 ([Figures 4A and 4C](#)), and a N182 that also shows proximal, non-proteinaceous density in some maps. The density near T185 shows a putative phosphate density. The phosphorylation of T185 is thought to determine the glycosylation state of N183 ([Nicholson et al., 2013](#)), which shows a large, strong density connected to the N δ 2 atom corresponding to multiple glycans ([Figure 4C](#)). Interestingly, TMEM106B(120–254) fibrils were observed irrespective of the presence of the risk (T185) or protective (S185) haplotype ([Table S1](#)). All TMEM106B(120–254) fibril structures have been built using a threonine at residue 185, as it is not possible to distinguish between a serine and threonine sidechain (van der Waals volume of 73 Å³ and 93 Å³, respectively) or an average of both sidechains at a resolution of 2.7 Å.

Singlet TMEM106B(120–254) amyloid fibrils display molecular polymorphism

The protofilament structure (singlet fibril) formed by TMEM106B(120–154) is identical across many cases with a uniform inter-strand twist angle of -0.4° ([Figure S3B](#); [Tables S2 and S3](#)). However, distinct misfolding minima exist with a similar, but structurally distinct, conformer displaying an inter-strand twist

angle of -0.67° . The more twisted singlet fibril is also comprised of residues S120 to G254, adopting a C shape similar to the less twisted singlet fibril and involving an embedded N terminus and a three-layered C terminus ([Figures 5A and S5B](#)). However, the C-terminal fold is less kinked and more compact, showing an ordered rectangular arrangement. Notably, in sharp contrast to the outward-pointing corner turn observed in the TMEM106B protofilament ([Figures 4A, S5A, and S5C](#)), in this instance the chain curves inward and forms an L shape pointing directly toward the extreme C terminus ([Figures 5A and S5B](#)). In total, 19 β strands were identified (β 1– β 19), matching the number of β strands we found in the less twisted singlet fibril. Although residues S120–G254 form both low- and high-twist singlet fibrils, comparison of the secondary structure motifs adopted by TMEM106B(120–254) in each molecular polymorph ([Fitzpatrick and Saibil, 2019](#); [Petkova et al., 2005](#)) show differences with 21 in 93 residues having a β strand/loop mismatch ([Figure 5D](#)).

In general, β strands (β 1– β 6) at the N terminus are identical in terms of amino acid composition and highly similar in spatial arrangements. However, from β 7 onward, the more twisted singlet fibril shows significant variation from its less twisted counterpart ([Figures 5C, S5A, and S5B](#)). Strand β 8 is shorter, packed against β 17 by forming a short, hydrophobic antiparallel sheet-sheet interface. However, this interface is not tightly packed due to the lack of interaction between F237 and Q170. Strand β 8 then leads into β 9 with a flexible short loop consisting of S172–I176. There is a strong, large density visibly attached to K178 ([Figures 5B and S5B](#)) that could be an anionic cofactor or posttranslational modification ([Arakhamia et al., 2020](#)). Strands β 9– β 13 organize into a continuous long strand, with β 10 marking the beginning of the C-terminal three-layered fold. These five strands (β 9– β 13) are arranged horizontally in a straight line, connected with short loops (G188–P189, V199–P200) and tight single residue turns (N183, Q194). Strand β 12 is packed closely against β 16, possibly by hydrophobic clustering with the small hydrophobic residues I195, I224, and L226. Strand β 13 continues into β 14 with a near 90-degree turn, with A204 at the corner. We hypothesize that a more complex interaction occurs in this aromatic region as the density suggests two rotamers for Y209, which is able to form either a methionine-aromatic

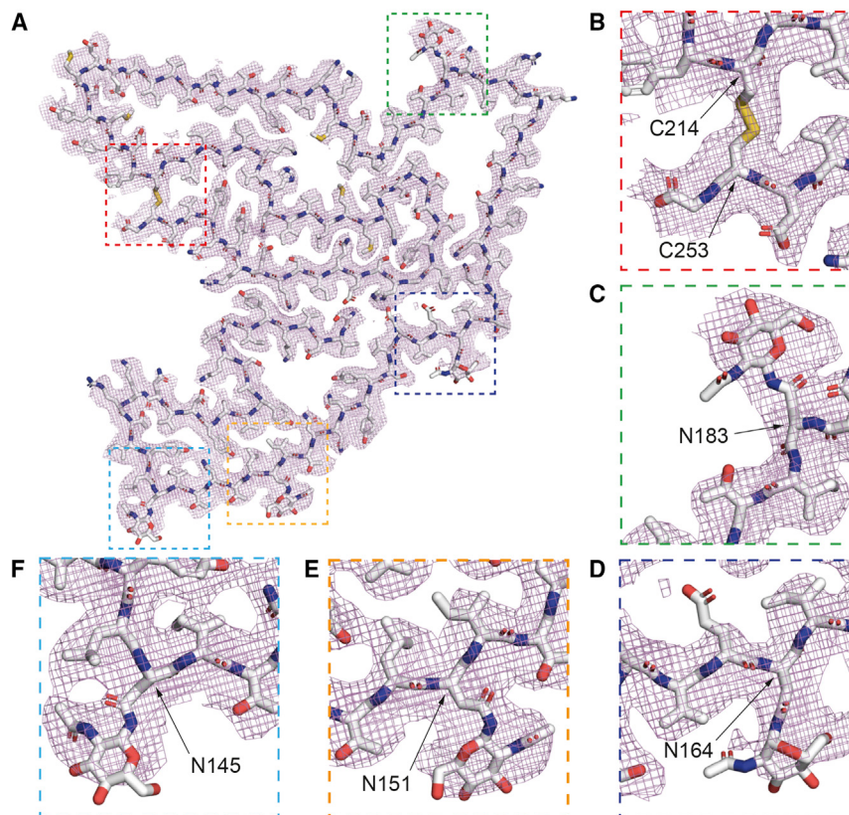


Figure 4. Cryo-EM structure of a TMEM106B protofilament highlighting the key structural features

(A) Cryo-EM density (mesh) and atomic model (sticks) of a TMEM106B protofilament. (B) A disulfide bond between C214 and C253. (C) Polymorphic site T185S and glycosylated asparagine at N183. (D–F) Glycosylated N164 (D), N151 (E), and N145 (F). See also [Figures S4 and S5](#) and [Table S2](#).

by cryo-EM, [Figure 2](#) and [Table S2](#)), B, and C along with five α -synucleinopathies ([Figure S6](#)). To probe for TMEM106B, we used a polyclonal antibody that binds to residues (204–253). As this antibody is not specific to a fibrillar epitope, all TMEM106B species are potentially detected. We observe bands consistent with monomeric TMEM106B(120–254), molecular weight of 15.49 kDa, at \sim 16 kDa ([Figure S6](#)) and multiple bands \sim 25 kDa ([Figure S6](#)) that may correspond to glycosylated TMEM106B(120–254) monomers. Notably, there are many higher molecular weight bands at 110 kDa, 135 kDa, and 190 kDa that are consistent with TMEM106B oligomeriza-

interaction with M207 or π - π interaction with Y211. At the C-terminal end of β 14, C214 again forms a disulfide bond with C253, the same as observed in the less twisted singlet fibril. This disulfide bond results in a sharp tip that steers into a conserved region consisting of β 15– β 17 and positions β 19 into forming a flattened and smooth strand following β 18.

Comparison of TMEM106B in pathological and non-pathological human tissue

As the discovery of TMEM106B fibrils is a recent and surprising finding, antibodies that are specific to TMEM106B fibrillar signatures, such as PHF1 (pS396/404) or pS129 available for tau and α -synuclein filaments, respectively, have not yet been generated and validated. Antibodies that recognize fibrillar phospho-epitopes are necessary to unambiguously image TMEM106B fibrils in patient tissue using immunofluorescence and to quantitatively measure the amount of TMEM106B fibrils in insoluble protein fractions.

Despite the current lack of reagents that specifically bind to an exclusively fibrillar epitope, we performed western blot analysis on fractionated tissue pellets from five neurologically normal individuals aged between 68 and 73 years of age using a commercially available TMEM106B C-terminal antibody ([Figure S6](#)). For comparison, we also included a 48-year-old FTL-TDP type A homogenate (case 5; TMEM106B fibrils solved by cryo-EM, [Figure 2](#) and [Table S2](#)) in an adjacent lane and performed separate western blot analysis on fractionated tissue pellets from FTL-TDP types A (case 4; TMEM106B fibrils solved

([Jiang et al., 2022](#)) ([Figure S6](#)). The general trend we observe is that these high molecular weight species are more abundant in pathological cases compared to non-pathological controls, although there are exceptions (e.g., FTL-TDP case 4 is a sporadic case that shows a high concentration of TMEM106B fibrils by cryo-EM but not by western blot).

Importantly, we and others ([Jiang et al., 2022](#)) contend that western blots do not reliably detect TMEM106B aggregation. An antibody has recently been generated that detects non-fibrillar TMEM106B by binding to residues (239–250), which are buried in the fibril core ([Schweighauser et al., 2021](#)). Such an antibody can only measure accumulation of monomeric TMEM106B in the soluble or insoluble fractions and is therefore not suitable for detecting fibrils.

DISCUSSION

Our study has identified previously undetected amyloid fibrils composed of aggregated TMEM106B(120–254) protein fragments in postmortem brain tissue from multiple TDP-43 proteinopathies, a tauopathy, and a synucleinopathy. These observations were both unexpected and surprising. We further note that TMEM106B(120–254) fibrils were not observed in our previous studies of Alzheimer's disease ([Arakhamia et al., 2020](#); [Fitzpatrick et al., 2017](#)) or CBD tau filaments ([Arakhamia et al., 2020](#)). Although this suggests that TMEM106B(120–254) fibrils may not be a consistent feature of all tauopathies, our absence of evidence could simply be due to the limited n-number and

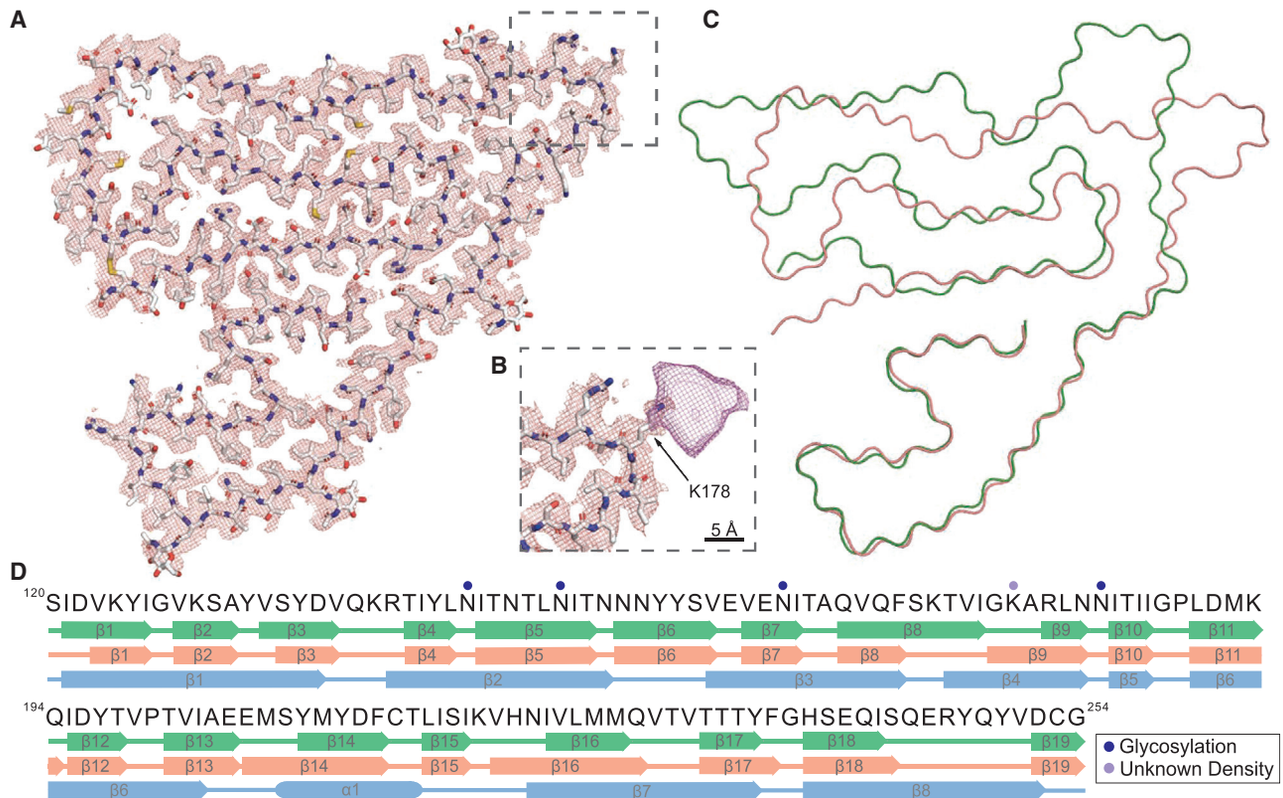


Figure 5. Cryo-EM structure of a highly twisted TMEM106B singlet fibril and comparison of singlet fibril molecular polymorphs

(A) Cryo-EM density (mesh) and atomic model (sticks) of a highly twisted TMEM106B singlet fibril.

(B) Magnified view of an unknown density bound to K178 in a highly twisted TMEM106B singlet fibril.

(C) Overlay of the atomic models (C α chain shown) of the low-twist (green) and high-twist (pink) singlet fibrils.

(D) Comparison of secondary structure motifs formed by the low-twist (green) singlet fibril, high-twist (pink) singlet fibril, and native protein predicted by AlphaFold (blue) with experimentally determined post-translational modifications of the highly twisted TMEM106B singlet fibril.

See also [Figure S5](#) and [Table S2](#).

varying brain regions in our cryo-EM studies of other tauopathies. It has also recently been suggested that variations in tissue fractionation protocols may significantly affect TMEM106B fibril yield ([Schweighauser et al., 2021](#)).

Notably, the abundance of TMEM106B(120–254) fibrils in two of our PSP cases is very high. We find that in PSP case 1, which also contains the PSP tau fold ([Shi et al., 2021](#)) ([Figure S3D](#); [Table S3](#)), TMEM106B(120–254) singlet fibrils are equally abundant as tau filaments. In PSP case 2, TMEM106B(120–254) singlet and doublet fibrils are approximately equal in number, with no PSP tau filaments observed in this sample ([Figure 2](#); [Table S3](#)). This is perhaps not surprising as PSP tau filaments are extremely sparse in the majority of PSP tissue samples we have screened (See [STAR Methods](#)). In contrast, PSP case 3 only contained PSP tau filaments ([Figure S3E](#)), and TMEM106B(120–254) fibrils were not observed. Recently, two cases with FTLD-TDP type B were shown to contain narrow, highly twisted filaments formed by TDP-43 (cross section ~ 40 Å \times 70 Å, helical pitch ~ 600 Å), although these fibrils were not found in all cortical regions and were not confirmed in other FTLD-TDP subtypes ([Arseni et al., 2022](#)). We observe TMEM106B(120–254) fibrils to be the predominant fibrillar spe-

cies in all FTLD-TDP cases we studied ([Figure 2](#); [Table S2](#)), irrespective of fractionation protocol (See [STAR Methods](#)). Finally, DLB is traditionally characterized by untwisted α -synuclein filaments with widths less than 100 Å ([Schweighauser et al., 2020](#)). In the present DLB case, we observe copious amounts of TMEM106B(120–254) singlet and doublet fibrils ([Figure 2](#); [Table S3](#)), but narrow, untwisted α -synuclein filaments were not observed (presumably due to insufficient fibril yield, see [STAR Methods](#)).

Interestingly, TMEM106B fibril structures similar to those identified in our study in patients across a wide range of proteinopathies as well as neurologically normal aged individuals was recently reported ([Schweighauser et al., 2021](#)). Although the findings are similar, our study is distinct in focusing primarily on genetic and sporadic forms of FTLD-TDP ($n = 8$, [Table S1](#)), whereas the work described in the recent report draws mainly from various tauopathies and synucleinopathies. The consensus finding that TMEM106B fibrils occur across diverse proteinopathies makes it difficult to determine whether TMEM106B fibrils are disease specific or are a general feature of neurodegeneration and aging. Importantly, the observation of increased TMEM106B fibrils in neurologically normal aged individuals

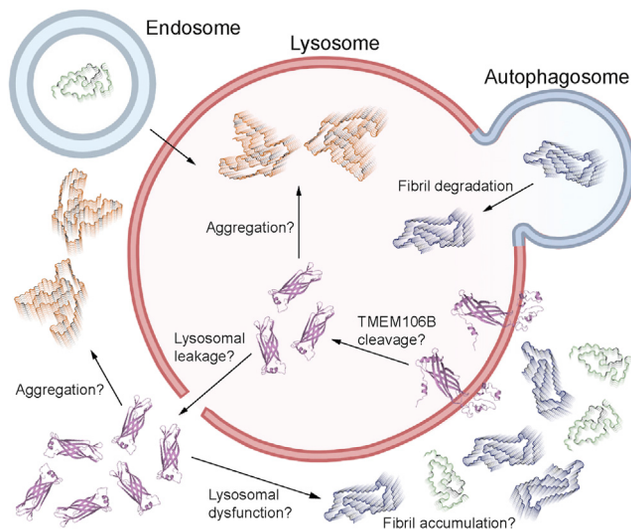


Figure 6. Structure-based model of a possible interrelationship between TMEM106B, aggregated filaments, and lysosomes in neurodegenerative diseases

Under physiological conditions, TMEM106B spans the lysosomal/endosomal membrane. Upon cleavage of the C-terminal fragment, TMEM106B(120–254) fibrils may form. It is currently unknown whether TMEM106B(120–254) fibrillizes in the lumen or whether lysosomal leakage occurs and TMEM106B(120–254) fragments aggregate into fibrils in the cytosol. Based on our set of TMEM106B fibril structures, we speculate that the aggregation of TMEM106B(120–254) into fibrils leads to lysosomal dysfunction, which promotes the accumulation of aberrantly aggregated amyloid fibrils such as those formed by TDP-43 (PDB:7PY2, green sticks), tau (PDB:7P65, blue sticks), or α -synuclein.

(Schweighauser et al., 2021) does not preclude a role in disease etiology as a similar observation is seen for tau (Brier et al., 2016; Chatterjee et al., 2021) and α -synuclein filaments (Hatton et al., 2022). Additionally, it is unclear whether the formation of TMEM106B amyloid fibrils represents a primary pathogenic process or a non-specific secondary phenomenon (e.g., general downstream marker of lysosomal stress). The observation of TMEM106B fibrils in some, but not all, cases of PSP (Figures 2, S3D, and S3E) (Shi et al., 2021) and DLB (Figure 2) (Schweighauser et al., 2021) suggests that the degree of fibrillization is highly variable in neurodegenerative diseases. Nevertheless, the combined trend across the two studies is that in the majority of cases with known neurodegeneration, fibrillization of TMEM106B is substantially greater when compared to age-matched controls (Figure S6) (Schweighauser et al., 2021).

Despite the small number of cases, this trend is most pronounced in the FTL-D TDP spectrum of neurodegenerative disease. All eight cases of FTL-D TDP in our study, representing all major pathological and genetic subtypes (Table S1; Figure S1), contained large amounts of TMEM106B fibrils per gram of frontal cortex (Figures 2 and S6). One relatively young patient (5th decade at death) with FTL-D TDP type A was replete with TMEM106B fibrils (Figures 2 and S6), arguing against TMEM106B fibrillization representing a non-specific feature of aging (Schweighauser et al., 2021). Furthermore, the observation that genetic variation in TMEM106B is a strong modifier of FTL-D TDP disease risk suggests that TMEM106B fibrillization may play

a more central role in FTL-D TDP pathogenesis (Finch et al., 2011; Van Deerlin et al., 2010; van der Zee et al., 2011).

If TMEM106B(120–254) is a potential driver of neurodegeneration, it will be important to determine what triggers the formation of these amyloid fibrils. Cellular models have revealed that overexpression of TMEM106B is associated with proteolytic cleavage that generates one or more N-terminal fragments and a presumed (though not yet observed) C-terminal fragment (Brady et al., 2014). We speculate that cleavage of the native TMEM106B fold by protease(s) generates an aggregation-prone fragment TMEM106B(120–254). As the isoelectric point of this luminal C-terminal fragment is pH 5.4, the acidic environment of the lysosome (pH 4.5–5.5) would favor charge-neutralization and possibly fibrillization (Smirnovas et al., 2005). The misfolding and aggregation of TMEM106B(120–254) may be influenced by posttranslational modifications (Arakhamia et al., 2020), such as glycosylation (N145, N151, N164, N183), and binding of cofactors (K178, R180) increasing the likelihood of molecular (Petkova et al., 2005; Zhang et al., 2019), and ultrastructural (Arakhamia et al., 2020; Fitzpatrick et al., 2013, 2017), polymorphism which manifest as amyloid fibril subtypes.

Based on our TMEM106B(120–254) fibril structures, we predict that cleavage at S120 may be due to thrombin, granzyme A, kallikrein-related peptidase 4, or cathepsin P (Table S4; Figure S5D). Granzyme A is particularly intriguing due to its role in the inflammatory response and the biogenic link between lytic granules and lysosomes (van Daalen et al., 2020). The degree to which proteolytic cleavage of the C-terminal fragment occurs *in vivo* is uncertain; however, it raises the possibility that factors that increase lysosomal stress, such as aging or reduced levels of progranulin (PGRN), combined with the presence of the higher expressing TMEM106B risk haplotype may drive more TMEM106B into the proteolytic pathway, resulting in the generation of TMEM106B C-terminal fragments in excess of the critical aggregation concentration favoring fibril formation (Hortschansky et al., 2005).

Given the presence of these previously undetected amyloid fibrils in myriad proteinopathies, it is pertinent to ask what possible effect the cleavage and aggregation of TMEM106B(120–254) could have on the trafficking and degradation of TDP-43, tau, and α -synuclein filaments. For now, we can only speculate, but given our current knowledge, we outline various possibilities based upon our structural observations (Figure 6). As TMEM106B is a lysosomal/endosomal protein, its loss of function via fibrillization could disrupt lysosomal trafficking on microtubules, impairing cellular clearance mechanisms (Brady et al., 2014; Lüningschrör et al., 2020; Schwenk et al., 2014). If TMEM106B(120–254) forms fibrils in the lumen, degradation of these protein aggregates places an extra burden on the lysosome. This will impede disposal of aberrantly aggregated filaments delivered to the lysosomes by autophagosomes or endosomes (Jiang and Bhaskar, 2020). If the lysosomal burden is increased, there may also be lysosomal leakage resulting in the release of TMEM106B(120–254) fragments and aggregates into the cytosol where they can propagate. Generally, we hypothesize that the fibrillization of TMEM106B(120–254) inside or outside the lysosome will trigger lysosomal dysfunction promoting the accumulation of TDP-43,

tau, and α -synuclein filaments in susceptible individuals (Figure 6). If TMEM106B(120–254) fibrils were to accumulate intracellularly, the loss/gain of function arising from the fibrillization of TMEM106B(120–254) would lead to markedly less efficient removal of TDP-43, tau, and α -synuclein filaments through neuronal or glial lysosomal degradation. Ultimately, our conjecture is that TMEM106B(120–254) fibrils exacerbate lysosomal stress and amplify or accelerate the accumulation of pathological proteins (Monaco and Fraldi, 2020).

In conclusion, we have determined the structures of TMEM106B(120–254) fibrils identified in a range of phenotypically distinct proteinopathies using cryo-EM. This unknown fibrillization pathway in all major TDP-43 proteinopathies, a tauopathy, and a synucleinopathy raises many intriguing questions. While future experiments will determine whether TMEM106B(120–254) fibril formation represents a loss or gain of function, our findings identify a common fibrillization pathway and widespread structural target that may be implicated in neurodegeneration.

Limitations of the study

At the time of publication, an antibody that binds exclusively to TMEM106B(120–254) fibrils does not exist. This has prevented immunohistochemistry experiments to directly visualize TMEM106B(120–254) fibrils in postmortem human tissue using immunofluorescence or immunogold electron microscopy.

STAR★METHODS

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SUPPLEMENTAL INFORMATION

Supplemental information can be found online at <https://doi.org/10.1016/j.cell.2022.02.026>.

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AUTHOR CONTRIBUTIONS

A.W.P.F., I.R.A.M., and M.H.B.S. conceived the experiments. I.R.A.M. identified patients and performed neuropathology on FTLD-TDP cases and PSP cases 2 and 3. D.W.D. and M.D. identified patients and performed neuropathology on PSP case 1. A.W.P.F. and C.L. purified amyloid fibrils from FTLD-TDP cases, PSP cases 2 and 3, and DLB. A.W.P.F. and C.L. performed western blots on sarkosyl-insoluble pellets fractionated from FTLD-TDP, DLB, and non-pathological tissue samples. Y.C. purified amyloid fibrils from PSP case 1. A.W.P.F., J.W., and C.W. performed cryo-EM experiments. A.W.P.F., A.C., X.X., J.W., T.A., and M.S. processed cryo-EM data. A.W.P.F., C.L., and G.Z. performed and analyzed mass spectrometry experiments. Y.C., D.W.D., and L.P. provided a PSP fibril sample. J.P., B.H., L.M.F., M.L.D., and R.R. made intellectual contributions. R.R. performed genetic analyses on all FTLD-TDP cases. A.W.P.F., X.X., J.W., and M.H.B.S. built and refined atomic models. A.W.P.F., A.C., X.X., J.W., T.A., M.S., S.D., M.T., C.N.C., R.R., D.W.D., L.P.,

M.H.B.S., and I.R.A.M. prepared the manuscript. M.H.B.S., I.R.A.M., and A.W.P.F. supervised the project.

DECLARATION OF INTERESTS

The authors declare no competing interests.

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STAR★METHODS

KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Antibodies		
Rabbit polyclonal anti-TMEM106B (204-253)	Novus Biologicals	Cat#NBP1-91311; RRID: AB_11019681; QC18333-42825
IRDye 800CW Donkey anti-Rabbit IgG Secondary Antibody	LI-COR Biosciences	Cat#926-32213; RRID: AB_621848; D11005-09
Mouse monoclonal anti-TMEM106B (1-46)	Proteintech	Cat#60333-1-Ig; RRID: AB_2881442
Rabbit polyclonal anti-TMEM106B (101-200)	Bioss	Cat#bs-11694R; RRID: AB_2905622
Rabbit polyclonal anti-TMEM106B (111-190)	Biorbyt	Cat#orb158617; RRID: AB_2905623
Rabbit polyclonal anti-TMEM106B (150-275)	Proteintech	Cat#20995-1-AP; RRID: AB_10694293
Rabbit polyclonal anti-TMEM106B (204-253)	Sigma Aldrich	Cat#SAB2106773; RRID: AB_2905624
Rabbit polyclonal anti-TMEM106B (218-252)	LSBio	Cat#LS-C757550; RRID: AB_2905625
Mouse monoclonal anti-pTau (phosphorylated at Ser262 and Ser356)	P. Seubert, Elan Pharmaceuticals; San Francisco, CA; USA	12E8
Mouse monoclonal anti-pTau (phosphorylated at Ser396 and Ser404)	P. Davies, Albert Einstein College of Medicine; New York, NY; USA	PHF1; RRID: AB_2315150
Mouse monoclonal anti-pTau (phosphorylated at Ser202)	P. Davies, Albert Einstein College of Medicine; New York, NY; USA	CP13; RRID: AB_2314223
Rabbit polyclonal anti-Tau (Human-specific)	L. Petrucelli, Mayo Clinic; Jacksonville, FL; USA	E1; RRID: 2819185
Rabbit polyclonal anti-TDP-43	Proteintech	Cat#10782-2-AP; RRID: AB_615042
Mouse monoclonal anti-pTau (phosphorylated at Ser202 and Thr205)	Thermo Fisher	Cat#MN1020; RRID: AB_223647
Mouse monoclonal anti- α -synuclein	Invitrogen	Cat# 180215; RRID: AB_86714
Mouse monoclonal anti-beta amyloid	Dako	Cat#M0872; RRID: AB_2056966
Rabbit polyclonal anti-ubiquitin	Dako	Cat#Z0458; RRID: AB_2315524
Biological Samples		
FTLD-TDP frontal cortex tissue	University of British Columbia (UBC)	https://neurology.med.ubc.ca/programs/alzheimer-disease-and-related-disorders-program/
PSP Case 1 caudate tissue	Mayo Clinic Brain Bank	https://www.mayo.edu/research/departments-divisions/department-neuroscience-florida/brain-bank
PSP Cases 2 and 3 frontal cortex tissue	University of British Columbia (UBC)	https://neurology.med.ubc.ca/programs/alzheimer-disease-and-related-disorders-program/
DLB frontal cortex tissue	Mayo Clinic Brain Bank	https://www.mayo.edu/research/departments-divisions/department-neuroscience-florida/brain-bank
Neurologically normal frontal cortex tissue	University of British Columbia (UBC)	https://neurology.med.ubc.ca/programs/alzheimer-disease-and-related-disorders-program/
Chemicals, Peptides, and Recombinant Proteins		
Intercept Blocking Buffer	LI-COR Biosciences	Cat#927-70001; 210816
iBlot 2 PVDF Mini Stacks	Invitrogen/Thermo Fisher Scientific	Cat#IB24002; 2PM131221-01
Novex WedgeWell 4-20% Tris-Glycine Gel	Invitrogen/Thermo Fisher Scientific	Cat#XP-4205BOX; 21030390
SDS-PAGE Running Buffer [10X]	GBiosciences	Cat#786-029G; 181405
LI-COR Odyssey CLx	LI-COR Biosciences	CLX-0707

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REAGENT or RESOURCE	SOURCE	IDENTIFIER
iBlot 2 Gel Transfer Device	Invitrogen/Thermo Fisher Scientific	Cat#IB21001; 0302180117
BLUEstain 2 Protein ladder 5-245 kDa	Gold Biotechnology	Cat#P008-500; 051301P008
Paraformaldehyde 32% Solution	Electron Microscopy Sciences	Cat#15714-S; 190107-10
XCell Sure Lock Mini-Cell	Invitrogen/Thermo Fisher Scientific	Cat#EI0001
cOMplete Mini, EDTA-free protease inhibitor cocktail tablets	Roche Diagnostics	Cat#11836170001; 35440400
Phenylmethylsulfonyl fluoride (PMSF)	Gold Biotechnology	Cat#P-470-10; 0302.091819A
Polytron Homogenizer	Kinematica	Cat#11030012

Deposited Data

Raw mass spectrometry data	This paper	MassIVE: MSV000088875
Cryo-EM density map of FTLN-TDP Type A (all cases combined) singlet	This paper	EMDB: EMD-26296
Cryo-EM density map of FTLN-TDP Type A (all cases combined) protofilament	This paper	EMDB: EMD-26279
Cryo-EM density map of FTLN-TDP Type A (case 1) singlet	This paper	EMDB: EMD-26295
Cryo-EM density map of FTLN-TDP Type A (case 1) doublet	This paper	EMDB: EMD-26294
Cryo-EM density map of FTLN-TDP Type A (case 1) protofilament	This paper	EMDB: EMD-26274
Cryo-EM density map of FTLN-TDP Type A (case 2) singlet	This paper	EMDB: EMD-26275
Cryo-EM density map of FTLN-TDP Type A (case 2) doublet	This paper	EMDB: EMD-26293
Cryo-EM density map of FTLN-TDP Type A (case 3) singlet	This paper	EMDB: EMD-26292
Cryo-EM density map of FTLN-TDP Type A (case 3) doublet	This paper	EMDB: EMD-26291
Cryo-EM density map of FTLN-TDP Type A (case 4) singlet	This paper	EMDB: EMD-26276
Cryo-EM density map of FTLN-TDP Type A (case 4) doublet	This paper	EMDB: EMD-26290
Cryo-EM density map of FTLN-TDP Type A (case 5) doublet	This paper	EMDB: EMD-26289
Cryo-EM density map of FTLN-TDP Type B case 1 (case 6) singlet	This paper	EMDB: EMD-26288
Cryo-EM density map of FTLN-TDP Type B case 2 (case 7) high-twist singlet	This paper	EMDB: EMD-26278
Cryo-EM density map of FTLN-TDP Type C (case 8) singlet	This paper	EMDB: EMD-26277
Cryo-EM density map of FTLN-TDP Type C (case 8) doublet	This paper	EMDB: EMD-26287
Cryo-EM density map of PSP case 1 singlet	This paper	EMDB: EMD-26284
Cryo-EM density map of PSP case 2 singlet	This paper	EMDB: EMD-26285
Cryo-EM density map of PSP case 2 doublet	This paper	EMDB: EMD-26286
Cryo-EM density map of PSP case 2 protofilament	This paper	EMDB: EMD-26273
Cryo-EM density map of PSP case 1 tau filament	This paper	EMDB: EMD-26268
Cryo-EM density map of PSP case 3 tau filament	This paper	EMDB: EMD-26283
Cryo-EM density map of DLB singlet	This paper	EMDB: EMD-26281
Cryo-EM density map of DLB doublet	This paper	EMDB: EMD-26282
Atomic model of FTLN-TDP Type A (all cases combined) protofilament	This paper	PDB: 7U16

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REAGENT or RESOURCE	SOURCE	IDENTIFIER
Atomic model of FTLD-TDP Type A (T185S) (all cases combined) protofilament	This paper	PDB: 7U18
Atomic model of FTLD-TDP Type A (case 1) protofilament	This paper	PDB: 7U11
Atomic model of FTLD-TDP Type A (case 2) singlet	This paper	PDB: 7U12
Atomic model of FTLD-TDP Type A (case 4) singlet	This paper	PDB: 7U13
Atomic model of FTLD-TDP Type B case 2 (case 7) high-twist singlet	This paper	PDB: 7U15
Atomic model of FTLD-TDP Type B case 2 (case 7) (T185S) high-twist singlet	This paper	PDB: 7U17
Atomic model of FTLD-TDP Type C (case 8) singlet	This paper	PDB: 7U14
Atomic model of PSP case 1 tau filament	This paper	PDB: 7U0Z
Atomic model of PSP case 2 protofilament	This paper	PDB: 7U10

Software and Algorithms

MotionCor2	(Zheng et al., 2017)	https://msg.ucsf.edu/software
Gctf v1.06	(Zhang, 2016)	https://www2.mrc-lmb.cam.ac.uk/research/locally-developed-software/zhang-software/
RELION	(He and Scheres, 2017)	https://www2.mrc-lmb.cam.ac.uk/relion
Chimera	(Pettersen et al., 2004)	https://www.cgl.ucsf.edu/chimera/
Coot	(Emsley et al., 2010)	https://www2.mrc-lmb.cam.ac.uk/personal/pemsley/coot/
Phenix.real_space_refine (v1.17.1-3660)	(Afonine et al., 2018)	https://phenix-online.org/
MolProbity	(Chen et al., 2010)	http://molprobity.biochem.duke.edu/
CamSol method	(Sormanni et al., 2015)	http://www.vendruscolo.ch.cam.ac.uk/camsolmethod.html
PyMOL		https://www.pymol.org
AlphaFold	(Jumper et al., 2021)	https://alphafold.ebi.ac.uk/
TrRosetta	(Yang et al., 2020)	https://yanglab.nankai.edu.cn/trRosetta/
Robetta	(Kim et al., 2004)	http://rosetta.bakerlab.org/
DeepEMHancer	(Sanchez-Garcia et al., 2021)	https://cosmic-cryoem.org/tools/deepemhancer/
Deep Tracer	(Pfab et al., 2021)	https://deepttracer.uw.edu/home
Cryo-ID	(Ho et al., 2020)	https://github.com/EICN-UCLA/cryoID
findMySequence	(Chojnowski et al., 2021)	https://gitlab.com/gchojnowski/findmysequence
ProCleave server	(Li et al., 2020)	http://procleave.erc.monash.edu/
COSMIC	(Cianfrocco et al., 2017)	https://github.com/cianfrocco-lab/COSMIC-CryoEM-Gateway
MEROPS database	(Rawlings et al., 2018)	https://www.ebi.ac.uk/merops/
MaxQuant	(Cox and Mann, 2008)	https://www.maxquant.org/

Other

Holey carbon grids (Au R1.2/1.3, 300 mesh)	Quantifoil	N/A
BioRender		https://biorender.com/
atom2svg		https://gist.github.com/biochem-fan/026ec2f191fee9285424d12fc2b84ce7

RESOURCE AVAILABILITY

Lead contact

Further information and requests for resources and reagents should be directed to and will be fulfilled by the lead contact, Anthony W. P. Fitzpatrick (anthony.fitzpatrick@columbia.edu).

Materials availability

This study did not generate new unique reagents.

Data and code availability

- All cryo-EM maps have been deposited to the Electron Microscopy Data Bank (EMDB; <https://www.ebi.ac.uk/pdbe/emdb/>). All refined atomic models have been deposited to the Protein Data Bank (PDB; <https://www.rcsb.org/>). All EMDB and PDB accession numbers are listed in the [key resources table](#). The mass spectrometry raw data has been deposited to MassIVE (<https://massive.ucsd.edu>) under the identifier MSV000088875.
- The paper does not report original code.
- Any additional information required to reanalyze the data reported in this paper is available from the lead contact upon request.

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Acquisition of human tissues

The tissues used for this study were from the frontal cortices of brains with FTLD-TDP (n = 11, 4 males, 7 females), PSP (n = 3, 2 male, 1 female), DLB (n = 6, 3 males, 3 females), and neurologically normal controls (n = 5, 3 males, 2 females), with the exception of PSP case 1 tissue being taken from the caudate region. Patients had been followed at the University of British Columbia Clinic for Dementia and Related Disorders. Informed consent for autopsy and the use of tissue for research was obtained from the legal next-of-kin in accordance with institutional ethics guidelines. Human brain samples were collected under a Mayo Clinic IRB-approved autopsy protocol. Tissues were de-identified prior to use, with no identifiers to patient names linked to brain samples. Patient confidentiality is strictly kept. As such, the utilization of autopsy derived, de-identified human brain tissue in the current study is exempt from IRB regulations. All tissue was handled with universal precautions. For human postmortem brain tissue requests from the Mayo Clinic Brain Bank, see <https://www.mayo.edu/research/departments-divisions/department-neuroscience-florida/brain-bank>.

Overview of FTLD-TDP cases

Filaments for cryo-EM analysis were extracted from fresh-frozen post-mortem brain tissue samples from eight cases with FTLD-TDP, including four with FTLD-TDP type A caused by different *GRN* mutations (one of which also carried the *C9orf72* repeat expansion, Case 5) and four sporadic FTLD-TDP (one type A, two type B and one type C), in which *GRN* and *C9orf72* mutations were excluded (Table S1). The *TMEM106B* haplotypes were determined for each case.

Clinical history of FTLD-TDP Type A case 1

The patient was a 60-year-old man with a five-year history of behavioral variant FTD (bvFTD) and mild parkinsonism. There was a family history of early-onset dementia and genetic sequencing of the patient identified a *GRN* mutation (c.1072C>T).

Clinical history of FTLD-TDP Type A case 2

The patient was a 55-year-old woman with a seven-year history of non-fluent variant primary progressive aphasia (nfvPPA) and subsequent behavioral abnormalities. Detailed family history was not available; however, genetic analysis identified a *GRN* mutation (c.1180-1G>A).

Clinical history of FTLD-TDP Type A case 3

The patient was a 60-year-old woman with a history of psychosis who developed progressive aphasia and behavioral changes at age 54. There was a family history of dementia and parkinsonism and genetic analysis identified a *GRN* mutation in the patient (c.102delC).

Clinical history of FTLD-TDP Type A case 4

The patient was an 89-year-old man with a 10-year history of sporadic nfvPPA.

Clinical history of FTLD-TDP Type A case 5

The patient was a 48-year-old woman who developed progressive non-fluent aphasia at age 44 and had evidence of mild bulbar weakness the year prior to her death. There was a strong family history of FTD and genetic analysis demonstrated a *GRN* mutation (c.87_90dup) as well as the abnormal *C9orf72* repeat expansion in the patient and other affected family members.

Pathology of FTLD-TDP Type A cases

All five cases with FTLD-TDP type A pathology had abundant compact neuronal cytoplasmic inclusions (NCI), short dystrophic neurites (DN) and occasional lentiform neuronal intranuclear inclusions (NII) concentrated in the upper neocortical laminae (Figure S1). Case 1 and Case 3 also had tau-immunoreactive neurofibrillary tangles (NFT) restricted to the entorhinal cortex and/or hippocampus

(Braak stage I and III, respectively). In addition to FTLD-TDP, Case 5 also had mild loss of lower motor neurons in the hypoglossal nucleus with rare TDP-43-positive inclusions and abundant dipeptide repeat immunoreactive neuronal inclusions, characteristic of the C9orf72 repeat expansion.

Clinical history and pathology of FTLD-TDP Type B case 1 (case 6)

The patient was a 62-year-old man with a four-year history of behavioral and cognitive changes followed by progressive weakness, consistent with amyotrophic lateral sclerosis (ALS). TDP-43 immunohistochemistry demonstrated abundant NCI throughout all layers of the neocortex, with most having a diffuse, granular morphology, typical of FTLD-TDP type B. There was also significant loss of spinal cord lower motor neurons with many containing TDP-43-immunoreactive inclusions. NFT were restricted to the hippocampus (Braak stage III).

Clinical history and pathology of FTLD-TDP Type B case 2 (case 7)

The patient was a 74-year-old Asian woman who developed progressive limb and bulbar weakness at age 72. Electrophysiological studies revealed denervation in multiple muscle groups, consistent with a diagnosis of ALS. Cognitive assessment at age 73 demonstrated significant problems with short-term memory, attention and abstract thought. Postmortem examination confirmed ALS with degeneration of the corticospinal tracts and loss of lower motor neurons with many of the remaining cells containing TDP-43 immunoreactive cytoplasmic inclusions. TDP-43 pathology in the primary motor cortex and prefrontal cortex consisted of abundant diffuse, granular neuronal cytoplasmic inclusions and small neuropil dots in all cortical laminae, consistent with FTLD-TDP type B. Neurofibrillary tangles were restricted to the hippocampus (Braak stage III/VI).

Clinical history and pathology of FTLD-TDP Type C case 1 (case 8)

The patient was a 69-year-old man with a 14-year history of progressive language problems with prominent word finding difficulties, characteristic of semantic variant PPA (svPPA), and mild behavioral problems. TDP-43 immunohistochemistry demonstrated numerous DN in the neocortex, many of which were long and tortuous, typical of FTLD-TDP type C. He also had a number of small chronic cerebral infarcts.

Clinical history and pathology of FTLD-TDP Type A case 6 (case 9)

The patient was a woman who developed bvFTD at age 53 and died at age 60. There was a strong family history of FTD and genetic analysis identified a *GRN* mutation (c.87_90dup). Post-mortem evaluation demonstrated FTLD-TDP type A as the only significant pathology.

Clinical history and pathology of FTLD-TDP Type B case 3 (case 10)

The patient was a 58-year-old woman who presented with nvPPA followed by mild changes in personality and behavior. During the year prior to death, she developed swallowing difficulty and electrophysiological studies demonstrated lower motor dysfunction. There was no family history of neurological disease and no *GRN* or *C9orf72* mutations were detected in the patient. Post mortem evaluation demonstrated typical changes of FTLD-TDP type B and ALS with TDP-43-positive inclusions in lower motor neurons. There were rare diffuse senile plaques in the neocortex and neurofibrillary tangles were restricted to the entorhinal/transentorhinal cortex (Braak II/VI).

Clinical history and pathology of FTLD-TDP Type C case 2 (case 11)

The patient was a 72-year-old woman who developed language difficulty consistent with svPPA, personality and behavioral changes and poor memory in her early sixties. There was no family history of neurological disease and no *GRN* or *C9orf72* mutations were detected. Post mortem exam demonstrated FTLD-TDP type C pathology and neurofibrillary tangles restricted to the hippocampus (Braak IV/VI).

Clinical history and pathology of PSP case 1

The patient was a 68-year-old white man who developed double vision six years prior to death. He subsequently developed slurred speech, bradykinesia, difficulty walking and unexplained falls, clinically diagnosed as Parkinson-plus syndrome. Post-mortem evaluation revealed findings of cortical-predominant PSP with severe neuronal and glial tau pathology in the motor and premotor cortices, and marked white matter oligodendroglial pathology at the gray-white junction. There was no TDP-43 pathology.

Clinical history and pathology of PSP case 2

The patient was a 75-year-old man with a 10-year history of progressive speech apraxia, cognitive changes and ophthalmoplegia. Gallyas-positive, tau-immunoreactive NFT and glial inclusions, including tufted astrocytes, ramified astrocytes, threads and coil bodies were present throughout the neocortex and numerous subcortical regions, consistent with PSP. There was no TDP-43 or α -synuclein-immunoreactive pathology.

Clinical history and pathology of PSP case 3

The patient was a 59-year-old woman with eight years of progressive cognitive impairment, behavioral changes, supranuclear gaze palsy, gait problems, dysphagia and akinetic-rigidity, clinically diagnosed as PSP. Typical neuropathological features of PSP were present including numerous tau-positive NFT, tufted astrocytes and other glial inclusions throughout the neocortex and many other brain regions. There was no TDP-43 and α -synuclein pathology.

Clinical history and pathology of DLB case 1

This 68-year-old man had a 10-year history of parkinsonism, with subsequent cognitive impairment. His parkinsonism was characterized by tremors, stiffness, gait difficulties, falls and handwriting difficulties. His cognitive problems included change in personality, disorientation and visual hallucinations. Post mortem examination demonstrated degeneration of the substantia nigra and locus coeruleus. Histologic studies showed diffuse α -synuclein immunoreactive cortical Lewy bodies and minimal Alzheimer type pathology (Braak stage I, Thal phase 1, CERAD neuritic plaque score 0).

Clinical history and pathology of DLB cases used for western blot analysis

DLB cases used for western blot analysis had a pathological diagnosis of DLB or variants thereof. Namely, case 2 (male, 57 years old, DLB A53T), case 3 (male, 66 years old, familial DLB), case 4 (female, 62 years old, DLB), case 5 (female, 81 years old, severe DLB), case 6 (female, 71 years old DLB A53T).

Clinical history and pathology of neurologically normal cases (NN) used in western blot analysis

Neurologically normal cases NN1 (male, 70 years old), NN2 (male, 73 years old), NN3 (female, 73 years old), NN4 (male, 68 years old), and NN5 (female, 70 years old) had no significant neurodegenerative pathology apart from age-related senile plaques (sparse in NN5, moderate in NN3).

METHOD DETAILS

Histology, immunohistochemistry, immunoblotting

At the time of publication, an antibody that binds exclusively to TMEM106B(120–254) fibrils does not exist. This has prevented immunohistochemistry experiments to directly visualize TMEM106B(120–254) fibrils in postmortem human tissue using immunofluorescence or immunogold electron microscopy.

FTLD-TDP

The post-mortem evaluation followed consensus recommendations for the neuropathological assessment of dementia (Montine et al., 2012) and FTLD (Cairns et al., 2007). FTLD-TDP subtypes were assigned in accordance with recent consensus criteria (Mackenzie et al., 2011). Five μ m thick sections of formalin fixed, paraffin embedded tissue were stained using hematoxylin and eosin and Bielschowsky silver methods. Routine diagnostic Immunohistochemistry (IHC) was performed using the Ventana BenchMark XT automated immunostainer with primary antibodies against TDP-43, hyperphosphorylated tau (pS202, pT205), α -synuclein, beta-amyloid, and ubiquitin.

In an attempt to demonstrate TMEM106B in post-mortem tissue, we tested a number of commercially available antibodies raised against different regions of the protein, using a wide range of IHC protocols. A mouse monoclonal antibody against amino acids 1–46 of the N-terminal region showed a pattern of reactivity consistent with the expected lysosomal localization of physiological TMEM106B, consisting of small cytoplasmic dots in neuronal perikarya and oligodendrocytes. However, this antibody did not label any structures that resembled the pathological inclusions seen with TDP-43 IHC. Despite employing a number of different pretreatment and amplification steps, none of the antibodies against C-terminal epitopes showed any staining that was interpreted as being specific for either physiological or pathological forms of TMEM106B. These included rabbit polyclonal antibodies against TMEM106B epitopes (101–200), (111–190), (150–275), (204–253), and (218–252).

PSP

For Case 1 paraffin sections were immunostained for tau (CP13, 1:1,000) using a DAKO Autostainer. For Cases 2 and 3 sections of formalin fixed, paraffin embedded tissue were stained using hematoxylin and eosin, Bielschowsky and Gallyas silver methods. Immunohistochemistry was performed using the Ventana BenchMark XT automated immunostainer with primary antibodies against TDP-43, hyperphosphorylated tau (pS202, pT205), α -synuclein, beta-amyloid, and ubiquitin.

DLB

Immunohistochemistry for α -synuclein (NACP) was performed after formic acid with the DAKO Autostainer.

Genetic sequencing

DNA was extracted from frozen brain tissue using standard techniques. Sanger sequencing analysis of all exons of the *GRN* gene and expansion analysis of the GGGGCC repeat in *C9orf72* using the 2-step protocol was performed in all FTLD-TDP cases as described previously (Baker et al., 2006; DeJesus-Hernandez et al., 2011). *TMEM106B* SNP rs3173615 (encoding T185S) was genotyped in all cases using an inventoried Taqman SNP genotyping assay C__27465458_10, as described (Nicholson et al., 2013).

Human brain tissue fractionation

All cases except PSP case 1

Added 10 volumes (w/v) of cold buffer I (10 mM Tris, 800 mM NaCl, 1 mM EGTA, 5mM EDTA, 10% sucrose, 1mM PMSF, Roche protease and phosphate inhibitor, pH 7.4) to frozen brain tissue from frontal cortex and homogenized by polytron (PT2500E, Kinematica, AG). The homogenized samples were centrifuged at 20,100 g for 20 min at 4°C and the supernatant was collected. The pellet was resuspended with 1 volume of cold buffer I, homogenized again and centrifuged at 21,100 g for 20 min at 4°C. The supernatant was then collected and combined with saved supernatant after the 1st centrifugation step. The supernatant was incubated with sarkosyl at a final concentration of 1% with gently shaking for 1 h at room temperature. Subsequently, the samples were centrifuged at 100,000 g for 1 h at 4°C. The final pellet was resuspended in 20 μ l cold buffer II (5 mM Tris pH 7.4) and saved for cryo-EM grid preparation. With regard to α -synuclein fibrils in our DLB case, we note that the tissue fractionation protocol used in the present study is similar to a previous cryo-EM paper (Schweighauser et al., 2020). However, we did not observe thin (< 100 Å wide), untwisted fibrils characteristic of α -synuclein filaments in our images. Similarly, despite tissue fractionating type B case 2 (case 7) using our protocol or a recently published protocol for obtaining TDP-43 fibrils (Arseni et al., 2022), we do not observe a significant number of TDP-43 fibrils in our images. Although we performed identical purification methods to isolate filaments from neurologically normal tissue samples (n = 5), these cases all had markedly smaller sarkosyl-insoluble pellets than those derived from diseased tissue (2-3 times smaller per gram of frontal cortex).

PSP case 1

To obtain a sufficient yield of tau filaments from PSP tissue, we evaluated the amount of tau present within the sarkosyl-insoluble (P3) fraction from four different brain regions in three different PSP cases, without pronase-treatment to preserve posttranslational modifications (Arakhamia et al., 2020; Carlomagno et al., 2021). As tau levels were consistently highest in the P3 fractions from caudate and frontal cortex, we screened these two regions in an additional 12 PSP cases. Using EM, we observed a tractable filament yield (~0.006 mg/mL) with low background in the P3 fraction from the caudate of case #1 (Figure S3D). To further characterize the PSP case used for cryo-EM, we evaluated total and phosphorylated tau levels in the P3 fraction of case #1 and 3 additional PSP cases by western blot (Figure S2C). The similarities in positivity for common disease-associated phosphoepitopes indicates case #1 is a typical PSP case (Figure S2C), suggesting that the morphology of tau filaments in this case will be representative of PSP.

Mass spectrometry

A sample imaged by cryo-EM, FTLD-TDP type A case 1, was incubated in 4M guanidinium hydrochloride to disaggregate the amyloid fibrils. In solution, digestion was performed using LysC and trypsin, followed by desalting and LC-MS/MS for protein identification and quantification. The data were processed by MaxQuant (Cox and Mann, 2008). Mass spectrometry data was searched against Uniprot human protein database. The TMEM106B fragment that was identified by mass spectrometry in FTLD-TDP type A case 1 was TMEM106B(130-139) (Table S5A). Mass spectrometry was performed in an identical manner on a second sample, FTLD-TDP type A case 6 (case 9), to identify proteins and phosphorylation sites (Table S5). To identify phosphorylation sites, the digest was enriched with TiO₂ for phosphopeptides before being subjected to LC-MS/MS. The TMEM106B fragments that were identified by mass spectrometry in FTLD-TDP type A case 6 (case 9) were TMEM106B(130-139) and TMEM106B(248-255) (Table S5B), with no phosphorylation sites detected on these peptides (Table S5C). Raw data was deposited on MassIVE (Project Accession # MSV000088875).

Western blots of sarkosyl-insoluble fractions from donor brains

Sarkosyl-insoluble pellets from fractionated tissue samples were resolved on Novex WedgeWell 4%–20% Tris-Glycine gels (Invitrogen) and electrophoresis performed at 25 Milliamps, 2 h. Proteins were dry transferred onto 0.2 μ m PVDF membranes using iBlot 2 Mini Stacks in an iBlot 2 Gel Transfer Device (Invitrogen) at 23 Volts for 6 min. Before blocking in Intercept Blocking Buffer (LI-COR), membranes were fixed in 1% paraformaldehyde for 30 min at room temperature. Blots were incubated with primary antibody TMEM106B against epitope (204-253) overnight at 4 degrees. Membranes were washed three times (10 min each) in PBS-T, then incubated with secondary antibody IRDye 800CW Donkey anti-Rabbit at 1:10,000 for 1 h at room temperature. Membranes were washed three times (10 min each) in PBS-T and imaged on LI-COR Odyssey CLx.

Predicted native fold of TMEM106B and aggregation propensity

The native structure of TMEM106B was predicted using AlphaFold (Jumper et al., 2021), TrRosetta (Yang et al., 2020), and Robetta (Kim et al., 2004). All three methods converged on highly similar structures of the full length-protein with the most variability predicted to be in the disordered N terminus. Predicted structures of TMEM106B(120-254) using these tools have a backbone root mean square deviation < 2 Å. The aggregation propensity of each TMEM106B(120-254) amino acid at pH 5 were predicted using CamSol (Sormanni et al., 2015) with an average insolubility of -1.39. Predictions of TMEM106B(120-254) aggregation propensity at lysosomal pH 5 and cytosolic pH 7 using the CAMSOL method are similar (integer pH values only possible).

Cryo-electron microscopy

All cases except PSP case 1

For cryoEM, 3 μ l aliquots of purified TMEM106B filaments from human brain tissue was applied to glow-discharged holey carbon Quantifoil Au R1.2/1.3 grids (Electron Microscopy Science, Hatfield, PA), blotted with filter paper to create a thin film by removing

excess sample using a FEI Vitrobot Mark III at 25°C and 100% humidity (blot force setting 0 and blotting time between 3.5–5.5 s) and plunge-frozen into liquid ethane. High-resolution data was collected primarily at the Zuckerman Institute, Columbia University with additional movies collected at the Columbia University Cryo-EM facility and New York Structural Biology Center (NYSBC), on Titan Krios microscopes equipped with BioQuantum-K3 imaging systems (see [Tables S2](#) and [S3](#) for details). All movies were collected in counting mode with a nominal pixel size of 1.07 Å/pixel, dose rate of 1.5 e/Å²/frame, over 40 frames. The underfocus range was set between 1.4 and 2.8 μm ([Tables S2](#) and [S3](#)).

PSP Case 1

3 μL aliquots of purified tau filaments from PSP human brain tissue (~0.006 mg/mL) were applied to glow-discharged holey carbon grids (Quantifoil Au R1.2/1.3, 300 mesh), blotted with filter paper and plunge-frozen into liquid ethane using an FEI Vitrobot Mark IV. Images of the frozen grids were collected on three FEI Titan Krios microscopes on multiple occasions due to the paucity of the tau filaments. Imaging conditions were always identical ([Table S3](#)), but minute differences in Å/pixel had to be corrected for during image processing. All images were acquired at 300 kV and energy filtered (slit width of 20 eV) using a Gatan K2/K3 Summit detector in counting mode. All movies were collected at a nominal pixel size of 1.0605 Å, using a dose rate of 1.5 electrons per Å² per frame and an underfocus range of 1 to 3 μm ([Table S3](#)).

Helical reconstruction

Movie frames were gain-corrected, aligned, dose-weighted, and summed using MotionCor ([Zheng et al., 2017](#)) implemented in RELION ([Zivanov et al., 2018](#)). Since amyloid fibrils extend over a few microns, spanning entire micrographs, during motion correction no patching was employed. Parameters for the contrast transfer function were estimated from each motion-corrected micrograph using Gctf ([Zhang, 2016](#)). All subsequent image processing was mainly performed using helical reconstruction in RELION ([He and Scheres, 2017](#)), as previously described ([Fitzpatrick et al., 2017](#)). Singlet and doublet fibrils were manually picked and separated with initial segmentation at 10.7 Å/pixel and then 3.21 Å/pixel to determine helical parameters ([Table S2](#)). Initial models of the singlet and doublet fibrils were generated from 2D class averages of helical pitch-views of the fibrils using a cylinder as an initial model and helically reconstructed in C1 using 3D classification with a T regularization value of 20. The doublet initial model quickly converged using FTLD-TDP particles; the singlet fibril initial model was generated from a PSP dataset in the following manner.

Helical processing of TMEM106B(120-254) singlet fibrils from PSP case 1 began with segments extracted at 3.1815 Å/pixel and 2D classified to determine the helical parameters. The low twist angles hampered initial model generation and so a cylinder was used as a starting model in 3D classification of 8 classes with a T regularization value of 20. One class consistently showed a C-shaped protofilament, irrespective of starting model (which was low-pass filtered to 40 Å). Particles from this class were re-extracted at 1.0605 Å/pixel and 3D classification with local optimization of the helical twist and rise was performed with T = 100, decreasing the angular and translational sampling as the rotational and translational accuracy of the helical 3D reconstruction improved. A final subset of particles was 3D auto-refined with T = 100 and postprocessed in RELION ([Zivanov et al., 2018](#)). Per-particle estimates of ctf values were used to improve resolution during a final refinement with an overall resolution of 4.4 Å being attained. This map had a fairly even distribution of views and was used as an initial model for refinement of TMEM106B(120-254) singlet fibrils from all other cases.

To achieve a high-resolution doublet fibril map, particles from selected 3D classes were re-extracted at 1.07 Å/pixel and re-aligned and averaged to generate 2D classes. Optimal 2D class averages showing clear β strand separation were selected and used for helical reconstruction in C1 using 3D classification with a T regularization value of 100. A persistent 2-fold symmetry was apparent in the doublet fibril reconstructions and this was imposed in subsequent 3D auto-refinements. Other symmetries, such as a 2-start helix or a helix with opposite hand, were tested but failed to attain high-resolution. Helical twist and rise were optimized during 3D auto-refinements ([Table S2](#)) and once the resolution of the 3D reconstruction was better than 4 Å, particles were corrected using Bayesian polishing and local ctf parameter refinement in RELION ([Zivanov et al., 2018](#)) to improve resolution further. Due to the low average inter-strand twist of approximately –0.4°, many reconstructions appeared to have anisotropic cross-sections. The two-fold symmetry of the doublet fibril was exploited to correct for this anisotropy by invoking `relion_particle_symmetry_expand` and refining the two masked, subtracted, protofilaments of the doublet cross-section separately, then combining the aligned particles in a final 3D auto-refinement. This improved our reconstructions significantly and a combined, sharpened map using the best particles from each FTLD-TDP case was generated and used for model building. Postprocessing was performed in RELION ([Zivanov et al., 2018](#)) and DeepEMHancer ([Sanchez-Garcia et al., 2021](#)) via COSMIC ([Cianfrocco et al., 2017](#)). Local resolution estimates of the combined map were determined using RELION ([Zivanov et al., 2018](#)) and displayed using Chimera ([Pettersen et al., 2004](#)) (Figure S4). The final overall resolution estimates were calculated from Fourier shell correlations at 0.143 between the two independently refined half-maps in RELION ([Zivanov et al., 2018](#)) (Figure S4).

Helical processing of tau filaments from PSP case 1 was performed using RELION ([He and Scheres, 2017](#)), as previously described ([Fitzpatrick et al., 2017](#)). Twisted tau filaments were manually selected and segments were extracted at various Å/pixel (1 × , 2 × , 3 × , 5 × binned) and 2D classified to determine the helical parameters and ascertain whether - or not - the filament had C2 symmetry. Initial models were generated from 2D class averages of helical pitch views and used as a reference for helical reconstruction in C1 using 3D classification with a T regularization value of 100. When a clear chain trace emerged in the helical cross-section at 1.0605 Å/pixel, a final subset of particles was 3D auto-refined with T = 20.

During the preparation of this manuscript, the structure of tau filaments from a PSP case was solved to a resolution of 2.7 Å. The public release of this high-resolution map (EMD-13218) and raw micrographs (EMPIAR-10765) helped in resolving continuous

main-chain density in 3D reconstructions of tau filaments from PSP case 1 using our final subset of particles. Helical twist and rise were optimized in the range -0.7° to -0.9° and 4.7 Å to 4.8 Å, respectively. Once the helical parameters were optimized (Table S3), helical symmetry remained fixed in all subsequent refinements. Postprocessing was performed in RELION (Zivanov et al., 2018), and DeepEMHancer (Sanchez-Garcia et al., 2021) via COSMIC (Cianfrocco et al., 2017) also aiding interpretation of the maps. The final overall resolution of 4.2 Å was calculated from Fourier shell correlations at 0.143 between the two independently refined half-maps in RELION (Zivanov et al., 2018) (Table S3). Tau filaments from PSP case 3 were extremely sparse, with only 3,942 particles at 3.22 Å/pixel being used in a final 3D auto-refinement with T = 4 resulting in a map consistent in size and shape with EMD-13218 low-pass filtered to 6.5 Å (Figure S3E).

Model building

A continuous C α chain was generated from the combined map using DeepTracer (Pfab et al., 2021). Tryptic digest-mass spectrometry of a fibril sample extracted from FTLN-TDP type A case 1 identified 600 proteins which we used to help identify the fibril's constituent protein. By cross-referencing the proteins in the sample identified by mass spectrometry to the 2.7 Å density map using cryo-ID (Ho et al., 2020) and *findMySequence* (Chojnowski et al., 2021), we unambiguously determined that the fibril was composed of TMEM106B(120-254). TMEM106B(120-254) sidechains were manually added to the 135 amino acid C α chain and iteratively improved using real-space refinement in COOT (Emsley et al., 2010). A 3-chain stack of identical rungs was generated in Chimera and multiple rounds of backbone and sidechain geometry refinement were performed manually followed by local real-space refinement in COOT (Emsley et al., 2010). For refinement of the atomic model built into the combined map (Figures 4 and S4), we used phenix.real_space_refine applying secondary structure, non-crystallographic symmetry, and hydrogen bond restraints (Afonine et al., 2018). Once we had a good atomic model of the TMEM106B(120-254) chain, additional glycans were added to match the density map in COOT (Emsley et al., 2010). A final refinement of the atomic model, including glycans was then performed and validated using MolProbity (Chen et al., 2010) (Table S2). All other structures were generated using this atomic model as an initial reference and iteratively refined in an identical manner. Figures were made using Pymol (<https://www.pymol.org>), Chimera (Pettersen et al., 2004), atom2svg (biochem-fan) and BioRender (biorender.com).

Predicted enzymatic cleavage

To investigate potential proteases responsible for TMEM106B cleavage at position 120 we utilized the ProCleave server (Li et al., 2020) and the MEROPS database (Rawlings et al., 2018). The MEROPS database of proteolytic enzymes, their substrates and inhibitors in 2017 and a comparison with peptidases in the PANTHER database. Nucleic Acids Res 46, D624-D632.) ProCleave results with scores above 0.5 at position TMEM106B(120) having the highest score are listed in Table S4: Granzyme A (single site), Thrombin (3 sites) and Kallikrein-related peptidase 4 (3 sites). Cathepsin P is not currently in the ProCleave suite but does have a predicted cleavage motif in MEROPS consistent with selective cleavage at position 120 of TMEM106B.

QUANTIFICATION AND STATISTICAL ANALYSIS

The 0.143 Fourier Shell Correlation (FSC) criterion was used to estimate resolution of cryo-EM density maps.

Supplemental figures

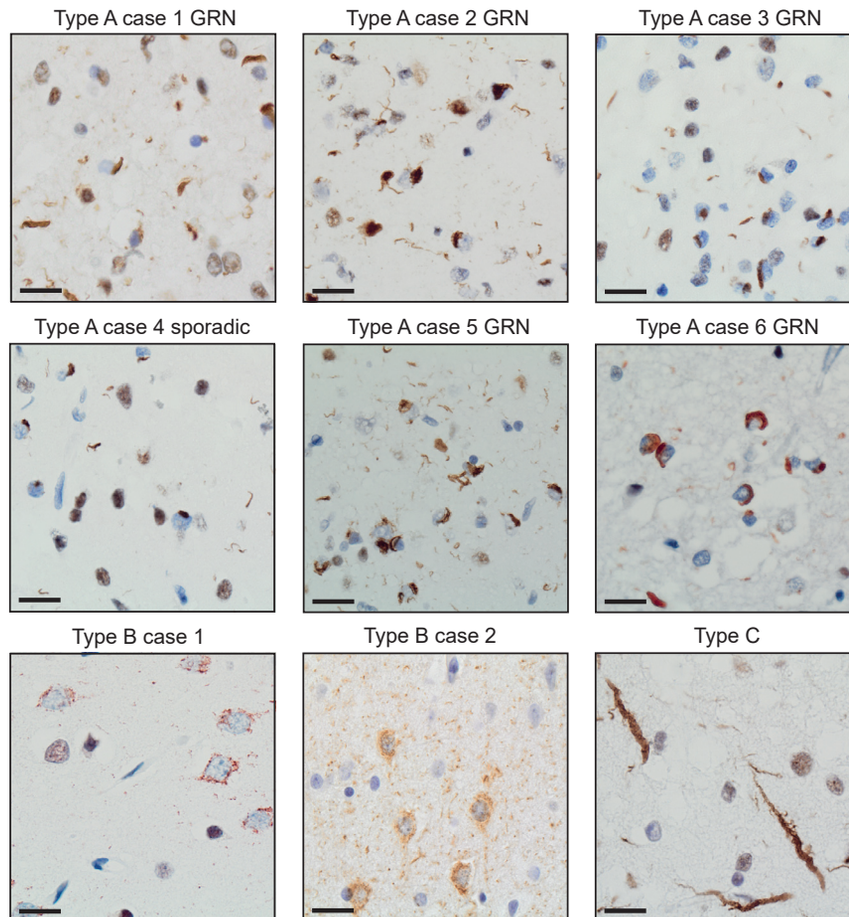


Figure S1. Immunohistochemistry of FTL-D-TDP cases used for cryo-EM and mass spectrometry, related to Figure 2 and STAR Methods
TDP-43 immunoreactive pathologic findings in frontal cortex of cases used for cryo-EM and mass spectrometry. Cases 1-5 and 9 (Type A case 6) showed changes typical of FTL-D-TDP type A, with abundant compact neuronal cytoplasmic inclusions, short dystrophic neurites and occasional lentiform neuronal intranuclear inclusions in the upper cortical laminae. Cases 6 (Type B case 1) and 7 (Type B case 2) had FTL-D-TDP type B pathology, characterized by diffuse, granular neuronal cytoplasmic inclusions in all cortical layers. Case 8 (Type C) had long, tortuous dystrophic neurites, diagnostic of FTL-D-TDP type C. Scale bar, 20 μ m.

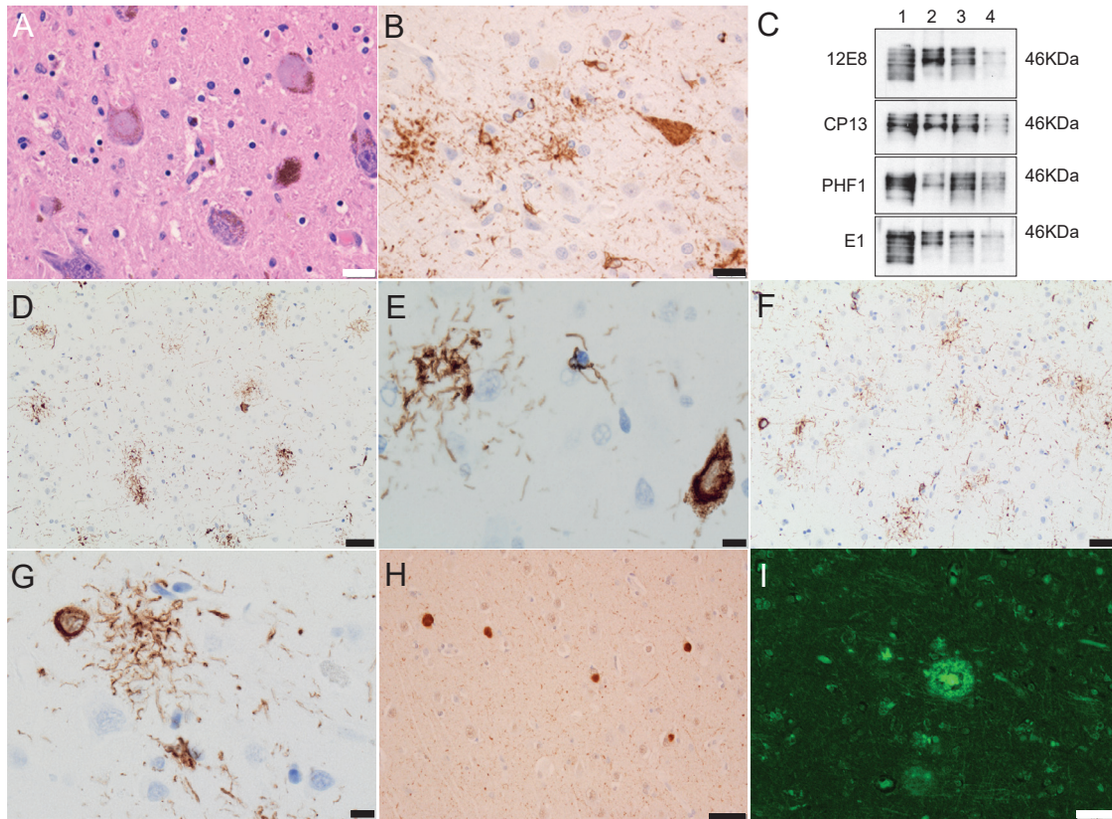


Figure S2. Neuropathological and biochemical analysis of three PSP cases used for cryo-EM, related to Figure 2 and STAR Methods

(A) and (B) The pigment-containing neurons of the substantia nigra have globus neurofibrillary tangles (hematoxylin and eosin), and tau immunohistochemistry shows neuronal and glial lesions (pretangles, tufted astrocytes, and oligodendroglial coiled bodies). Scale bar, 20 μm . (C) Sarkosyl-insoluble fraction extracted from the caudate/putamen of four different PSP cases as evaluated by immunoblotting for the tau antibodies 12E8 (pS262/356), CP13 (pS202), PHF1 (pS396/404), and E1 (human tau-specific). Tau filaments from (Mayo Clinic, Florida) PSP case #1 were subsequently used for cryo-EM analysis. (D-G) Pathologic findings in frontal cortex of PSP case 2 (D, E) and PSP case 3 (F, G) (both from UBC) had abundant neuronal and glial inclusions, including neurofibrillary tangles and tufted astrocytes, that were immunoreactive for hyperphosphorylated tau. (H, I) show immunohistochemistry of the DLB case used for cryo-EM. Frontal cortex with synuclein and also with thioflavin S fluorescent microscopy. Diffuse cortical LBs and minimal Alzheimer-type plaques are illustrated. (D, F) scale bar, 50 μm . (E, G) scale bar, 10 μm . (H, I) scale bar, 50 μm .

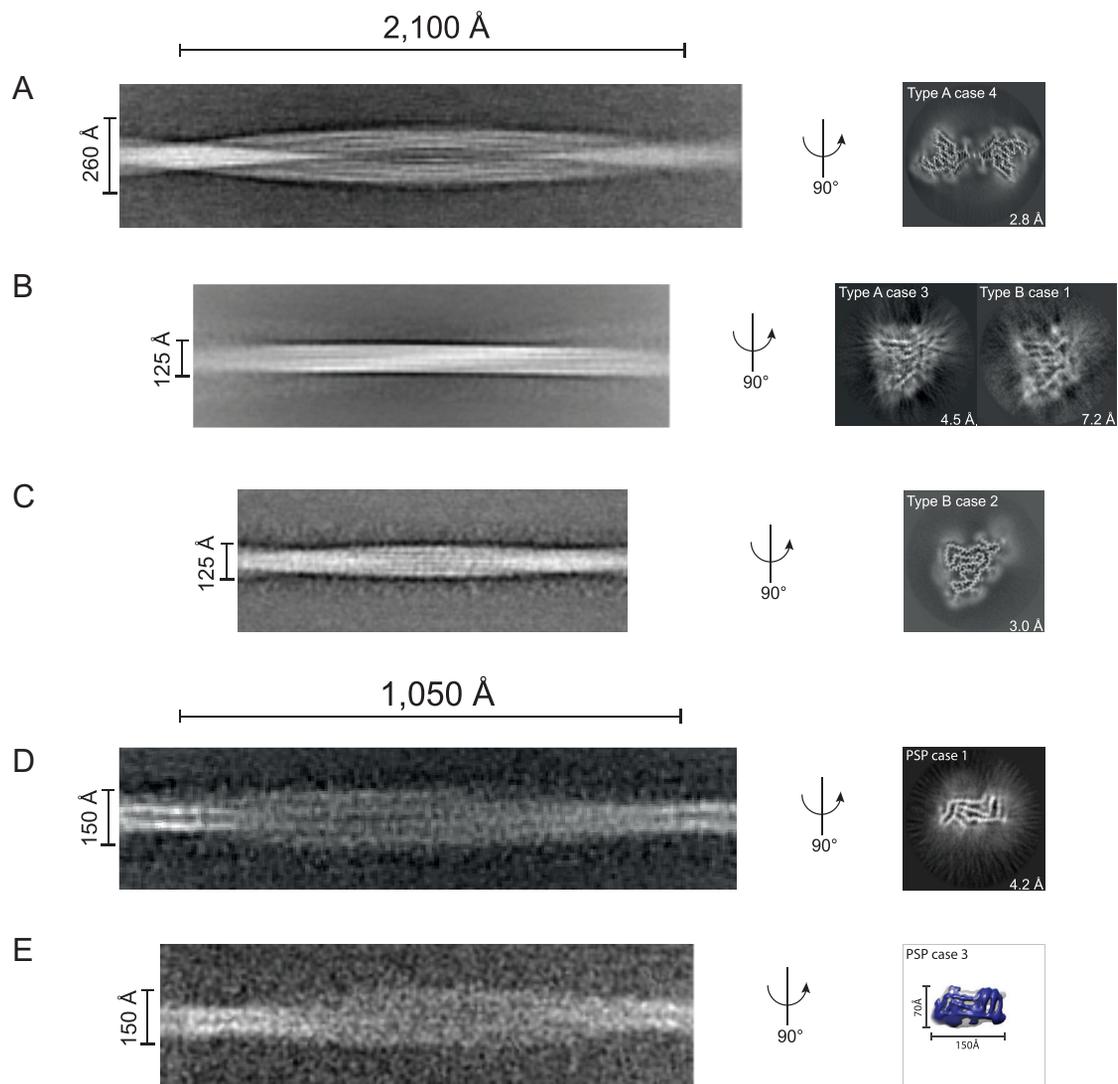


Figure S3. Comparison of the observed fibril types, related to Figure 2 and STAR Methods

(A-C) Pitch views of the (A) TMEM106B doublet, (B) singlet, and (C) highly twisted singlet fibrils and their respective cross-sections. (A-C) are on the same scale. (D, E) Pitch views of PSP tau filaments from PSP (D) case 1 and (E) case 3 and their respective cross-sections. (D, E) are on the same scale.

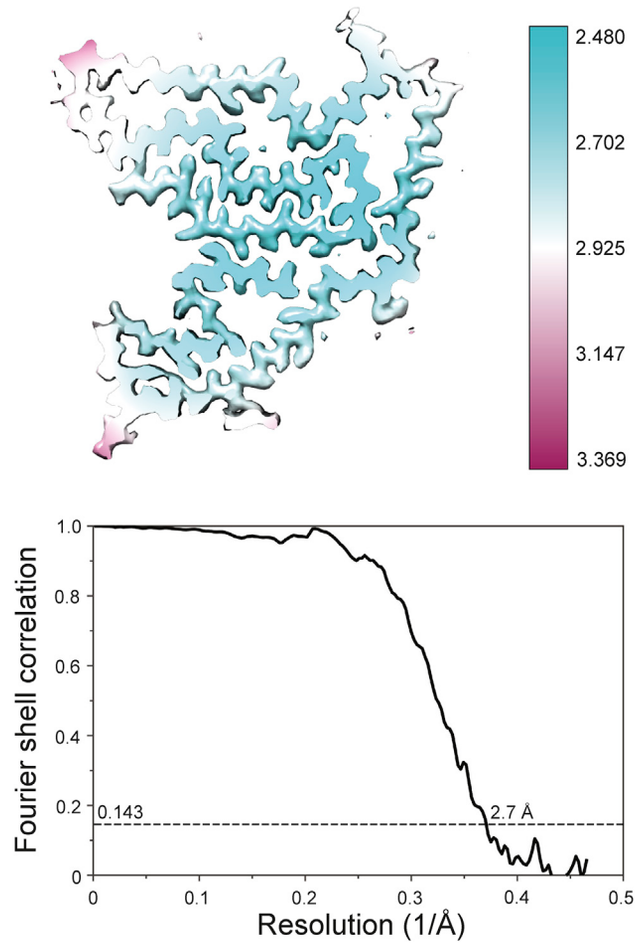


Figure S4. Cryo-EM resolution estimates of a combined cryo-EM protofilament map from multiple FTLT-TDP cases, related to Figure 4
Local resolution estimates (top) and Fourier Shell Correlation (FSC) curve (bottom) between two independently refined half-maps for the TMEM106B(120-254) protofilament.

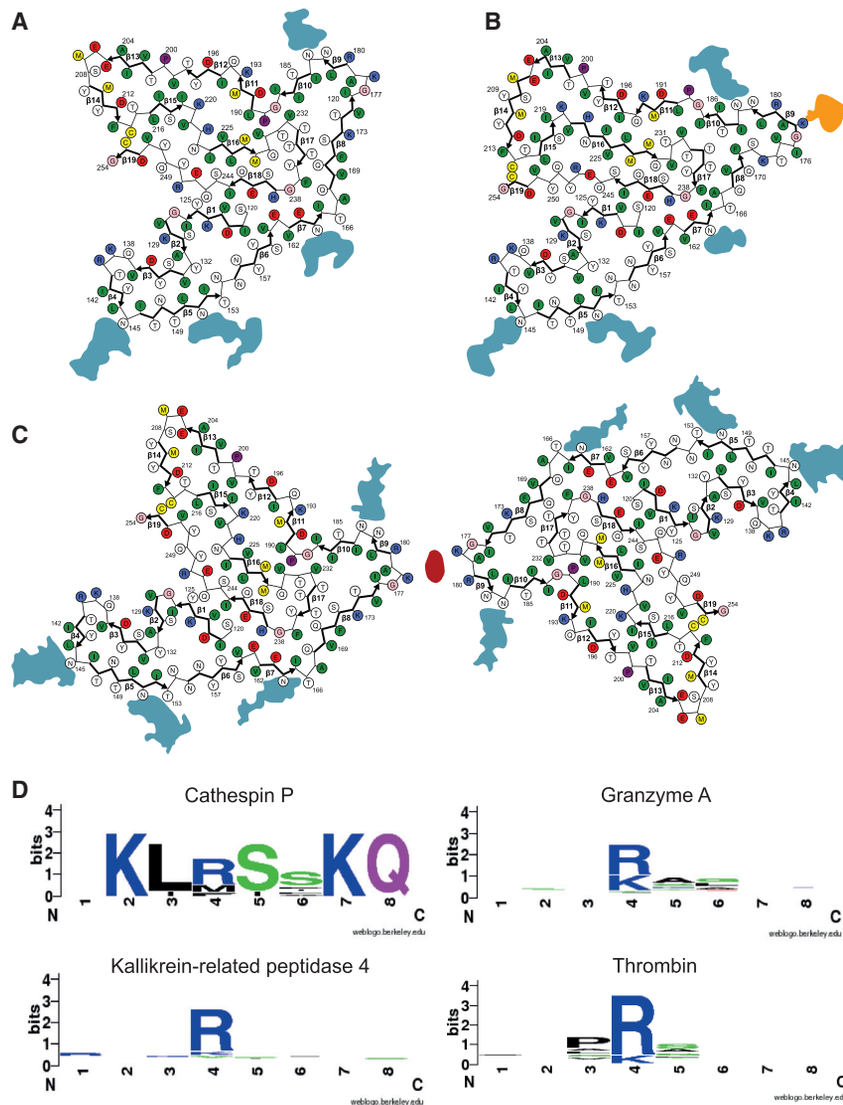


Figure S5. Schematic views of polymorphic TMEM106B(120-254) amyloid fibrils and fibril structure-based protease predictions, related to Figures 3-5 and Table S4

Singlet fibrils with (A) low inter-strand twist and (B) high inter-strand twist are shown. Bold line denotes β strand secondary structure with experimentally detected glycosylations shown in cyan. There is an unknown density (orange) attached to K178 in the highly twisted singlet fibril. (C) Doublet fibril with the amino acids K178 and R180 binding to an anionic cofactor highlighted in red. (D) Enzymes predicted to cleave TMEM106B at site S120.

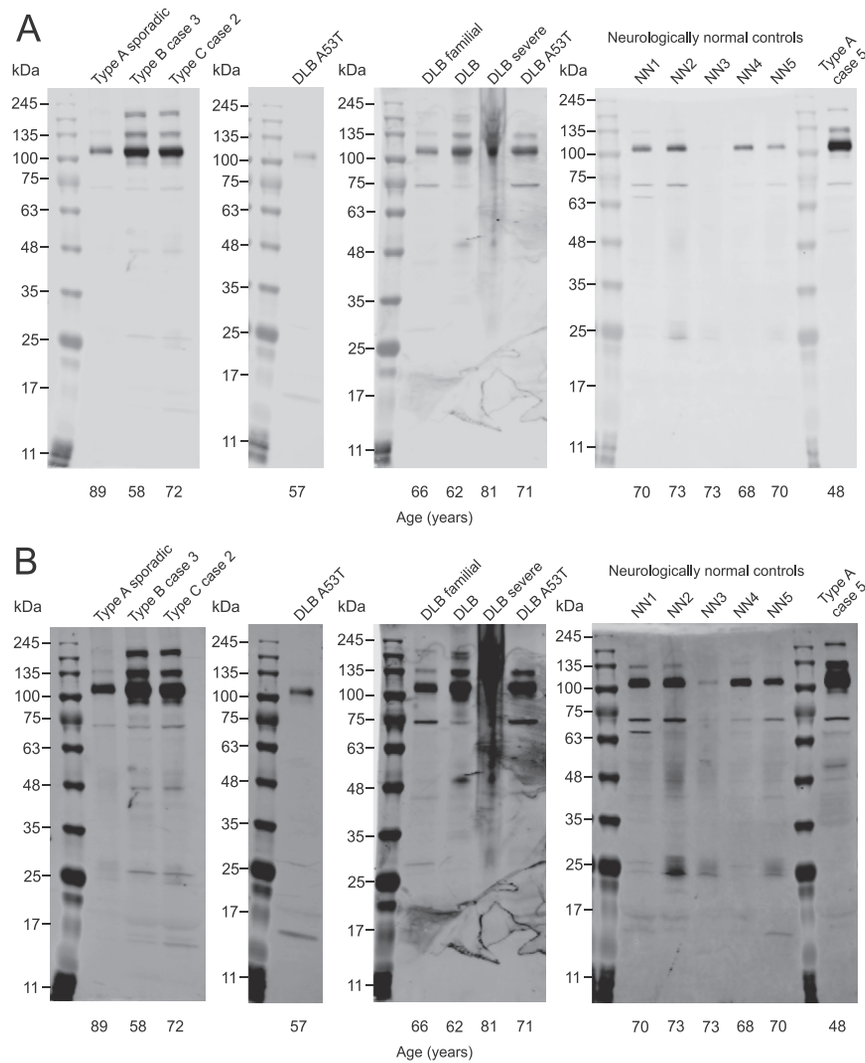


Figure S6. Western blots of sarkosyl-insoluble fractions from donor brains probed with a TMEM106B polyclonal antibody that binds to residues (204–253), related to Figure 2 and STAR Methods

(A) Equally-exposed western blots of sarkosyl-insoluble fractions from pathological and non-pathological tissues. (B) High contrast images of western blots in (A). Type A, Type B, Type C: FTLD-TDP subtypes. DLB: Dementia with Lewy Bodies. NN: Neurologically Normal.