

Integrative determination of the atomic structure of mutant huntingtin exon 1 fibrils from Huntington's disease

Mahdi Bagherpoor Helabad^{1,2}, Irina Matlahov³, Jan O. Daldrop⁴, Greeshma Jain³, Patrick C.A. van der Wel^{3,*} and Markus S. Miettinen^{1,4,5,6,*}

¹*Department of Theory and Bio-Systems, Max Planck Institute of Colloids and Interfaces, 14424 Potsdam, Germany*

²*Institute for Drug Discovery, Leipzig University Medical Center, 04103 Leipzig, Germany*

³*Zernike Institute for Advanced Materials, University of Groningen, Groningen, The Netherlands*

⁴*Fachbereich Physik, Freie Universität Berlin, 14195 Berlin, Germany*

⁵*Department of Chemistry, University of Bergen, 5007 Bergen, Norway*

⁶*Computational Biology Unit, Department of Informatics, University of Bergen, 5008 Bergen, Norway*

*Corresponding authors:
p.c.a.van.der.wel@rug.nl; markus.miettinen@iki.fi

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Abstract. Neurodegeneration in Huntington's disease (HD) is accompanied by the aggregation of fragments of the mutant huntingtin protein, a biomarker of disease progression. A particular pathogenic role has been attributed to the aggregation-prone huntingtin exon 1 (HttEx1) fragment, whose polyglutamine (polyQ) segment is expanded. Unlike amyloid fibrils from Parkinson's and Alzheimer's diseases, the atomic-level structure of HttEx1 fibrils has remained unknown, limiting diagnostic and treatment efforts. We present and analyze the structure of fibrils formed by polyQ peptides and polyQ-expanded HttEx1. Atomic-resolution perspectives are enabled by an integrative analysis and unrestrained all-atom molecular dynamics (MD) simulations incorporating experimental data from electron microscopy (EM), solid-state NMR, and other techniques. Visualizing the HttEx1 subdomains in atomic detail helps explaining the biological properties of these protein aggregates, as well as paves the way for targeting them for detection and degradation.

1 Introduction

Huntington's disease (HD) is one of a family of incurable neurological genetic disorders resulting from an aberrant expansion of a CAG trinucleotide repeat in a disease-specific gene.^{1,2} In HD, the mutation impacts the huntingtin (Htt) protein, which ends up with an expanded polyglutamine (polyQ) tract in its first exon (HttEx1, Fig. 1A).³ In HD patients and model animals, mutant Htt fragments matching HttEx1 form inclusions in brain areas affected by neurodegeneration. In cells, μm -sized inclusions are seen to contain fibrillar protein aggregates (Fig. 1B-C). The fibrillar HttEx1 adopts a morphology with all the common features of amyloid fibrils found in diseases such as Parkinson's and Alzheimer's disorders: highly stable protein deposits built around extended intermolecular β -sheets forming a characteristic cross- β architecture.⁴⁻⁶ The protein fibrils are of interest due to their potential involvement in pathogenic and neurotoxic processes—but also as biomarkers of disease onset and progression. Consequently, there is a

need to resolve their atomic structures to facilitate the design of inhibitors, modulators, and diagnostic tools; e.g., a better understanding of HttEx1 fibril structure is essential to the development of ligands used to detect fibril formation *in vivo*.⁷ Recent years have seen important progress in amyloid structure determination through application of cryogenic electron microscopy (cryo-EM) and solid-state NMR spectroscopy (ssNMR):^{4,8,9} Breakthrough studies have produced structures of amyloid fibrils formed by tau, amyloid- β , and α -synuclein, associated with Alzheimer's, Parkinson's, and other amyloid disorders.⁹

30 The polyQ-based amyloid fibrils from HD and other CAG repeat disorders, however, have proved more challenging, and still lack truly atomic-level structures or even structural models.² The aggregation propensity of polyQ proteins correlates strongly with their polyQ tract length, which inversely correlates with the age of onset.^{2,10} In their native state, polyQ segments are typically considered to lack a stable secondary structure, forming an intrinsically disordered region. This is also true for the polyQ-containing segment of Htt (HttEx1), which is invisible in cryo-EM of the full-length protein.^{11,12} However, in HD, N-terminal fragments of Htt can result from protease activity and missplicing, enabling subsequent aggregation into cellular inclusions.^{13,14} Cryogenic electron tomography in cellular context has revealed that μm -sized expanded inclusions contain nm-wide HttEx1 fibrils (Fig. 1B, C), which resemble those formed by purified HttEx1 *in vitro* (Fig. 1D).^{15,16}

40 Prior structural studies have shown that HttEx1 fibrils comprise a highly ordered polyQ amyloid core surrounded by flexible non-amyloid flanking regions.^{17–20} Fiber X-ray diffraction studies showed the core itself to comprise anti-parallel β -sheets in a cross- β pattern (Fig. 1E);^{21–24} these experiments spurred proposals of various possible fibril models,^{21–24} which however lacked atomic-level detail. Moreover, all early X-ray-based models proved inconsistent with later work^{5,25–27} that revealed important new structural data via combinations of cryo-EM, ssNMR, and other techniques.^{5,17–20,27–32} A few recent studies are of particular interest. Firstly, an integration of ssNMR, X-ray, and EM measurements permitted the manual assembly of a schematic fibril architecture (Fig. 1F-G).^{5,18,25,26} Subsequently, a cryo-EM study of HttEx1 fibrils³³ produced a medium-resolution density map, which—although it lacked the detail for a de-novo atomistic structure—was interpretable in terms of the said³⁴ polyQ amyloid core architecture (Fig. 1F, G).

50 The apparent success of such a unified (but as-yet qualitative) model underlines the need for a modern integrative-structural-biology approach³⁵ to fully describe these fibrils that have much in common with other amyloids, but also a number of truly unique features.

Here we employ rigorous physics-based molecular modeling to integrate the rich set of collective experimental knowledge into the first atomic-resolution polyQ and HttEx1 fibril structures that are fully consistent with all the key experimental data, including sheet-to-sheet distances from fiber X-ray diffraction, fiber dimensions from EM, and structural constraints from ssNMR. The obtained structures are analyzed in the context of known structural features of HttEx1 amyloid fibrils, allowing a detailed explanation of their conformational and spectroscopic characteristics. Moreover, based on the disposition and accessibility of key residues and segments, the molecular architecture of these HD-related fibrils explains biological properties, such as the (in)ability for post-aggregation polyubiquitination and degradation.

2 Results

2.1 Architecture of the internal polyQ fibril core

We started the integrative structure determination of HttEx1 fibril from its polyQ amyloid core. Most experimental studies, in particular recent ssNMR studies, argue for a molecular architecture common

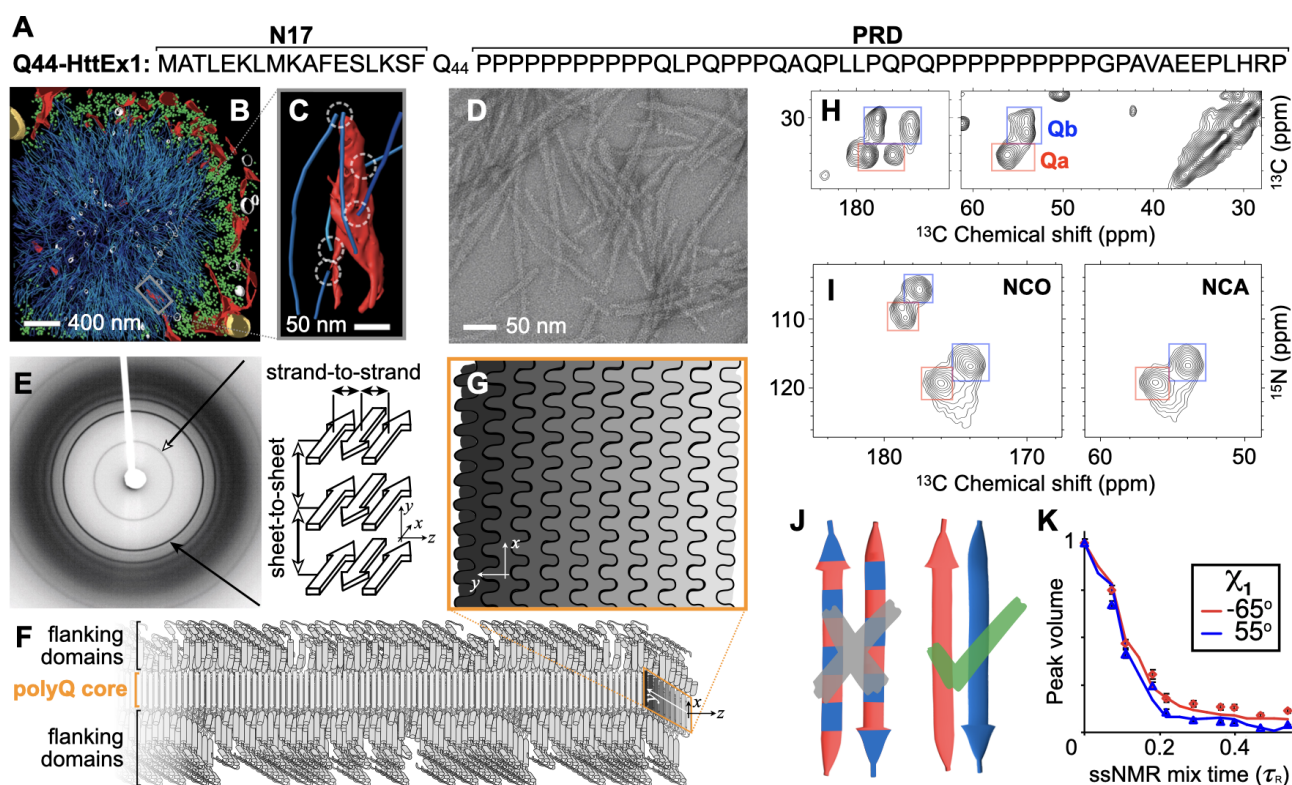


Figure 1. Huntingtin exon 1 sequence and aggregates. (A) Sequence of Q44-HttEx1. N- and C-terminal flanking domains marked as N17 and PRD, respectively. (B) Colorized EM tomogram of a HttEx1 inclusion in mouse primary neuronal cells, showing protein fibrils (cyan) and mitochondria (gold). (C) Magnified region showing individual filaments interacting with ER membranes (red). (D) Negatively-stained TEM of Q44-HttEx1 aggregates formed *in vitro*; average fiber width ~ 6.5 nm. (E) X-ray fiber diffraction of K₂Q₃₁K₂ fibrils, detecting the cross- β reflections of the amyloid core that represent the inter-strand and inter-sheet distances shown on the right. (F) Schematic model of a Q44-HttEx1 fibril, showing the N17 and PRD flanking domains outside the polyQ core. (G) Cross-section of the (6–7-nm wide) core showing the layering and interdigitation of the (differently shaded) β -sheets. (H) 2D ¹³C–¹³C ssNMR spectrum of ¹³C, ¹⁵N-labeled Q44-HttEx1 fibrils showing backbone and sidechain cross-peaks. Peaks for C α –CX correlations of two distinct “a” and “b” Gln conformers are marked with red and blue boxes, respectively. (I) 2D NCO and NCA ssNMR spectra of Q44-HttEx1 fibrils. (J) Pairs of antiparallel β -strands, schematically showing two previously proposed arrangements of the “a” (red) and “b” (blue) types of Gln. Given the ssNMR data (panels H–I), only the right one fits the experimental data. (K) ssNMR measurements sensitive of the side-chain dihedral angle χ_1 , with simulated curves for “a” (red) and “b” (blue) types of Gln. Panels B–C adapted with permission from Ref. 15, D–E and K from 5, F–G from 34, and I from Ref. 36.

65 to polyQ-containing peptide and protein fibrils.³⁷ Various techniques show that polyQ amyloid features antiparallel β -sheets (Fig. 1E),^{23,24,27,30,38} in contrast to the parallel in-register fiber architecture of many other amyloid fibers.^{8,39} Another distinct feature of polyQ amyloid is that it forms long β -strands with few turns (Fig. 1F),^{5,24,26,33,34} unlike the majority of other amyloid structures that have short β -strands connected by turns or loops.⁸ The polyQ fiber core is multiple nanometers wide, devoid of water molecules, and forms a block-like structure featuring a stacking of less than ten β -sheets.^{24–26,33} Its cross-section must contain polyQ segments from multiple protein monomers (schematically illustrated in Fig. 1F-G).³⁴ A large proportion of the Gln residues is buried, far away from the solvent, in a repetitive and pseudo-symmetric context. This ordered and repetitive nature is demonstrated by high-quality ssNMR spectra, with few and relatively narrow peaks (Fig. 1H).^{5,18,20,25,26,29,34} These ssNMR studies consistently show a characteristic

75 spectroscopic fingerprint that identifies two dominant Gln residue conformations—see the two sets of peaks from so-called “a” and “b” conformers (red and blue boxes) present at roughly equal intensities in the 2D ssNMR spectra of Q44-HttEx1 fibrils (Fig. 1H-I). The two Gln conformers reflect two types of β -strand structures that co-assemble into a single antiparallel β -sheet (Fig. 1J).^{5,20,26}

Integrating these experimental data, we concluded that the internal polyQ core structure can be captured
80 by an infinitely-repeating eight-Gln unit that features antiparallel β -sheets (Fig. 2A). This construction has translational symmetry and lacks the twisting seen in many amyloid fibrils, since EM analyses of our and others’ HttEx1 fibrils lack strong signs of twisting.³³ The repeating ‘unit cell’ (Fig. 2A) fulfills several *a priori* requirements: First, its peptide chain segments are two residues long, the minimum needed to capture the ‘odd/even’ side-chain alternation in a β -strand. Given that ssNMR shows only two ssNMR-detected
85 conformers, in long strands, two residues should be sufficient to capture the structural variation within this amyloid core. Second, the unit cell contains two β -strands (that can differ in conformation), the minimum needed to describe an antiparallel β -sheet. This contrasts with parallel in-register β -sheets that could be represented with a single repeating β -strand.⁴⁰ Third, the unit cell contains a stacking of two β -sheets, to permit the probing of distinct sheet–sheet dispositions. Fourth, certain ranges of side-chain dihedral angles
90 (χ_1, χ_2, χ_3) are required in order to fulfill a crucial feature of the interdigitating polyQ amyloid:^{5,23} the side-chain–side-chain hydrogen bonding that allows the polar Gln residues to be stowed in the waterless core environment (see Fig. 2A and Supplementary Fig. S1). Thus, e.g., the χ_2 dihedral must be close to 180°, as also known from ssNMR and Raman experiments.^{5,41} This minimal model (Fig. 2A) fulfills the above-noted experimental constraints, and still yields a library of roughly 1 000 distinct architectures (see
95 Methods for details).

A key additional consideration is that ssNMR has unambiguously shown sequential residues within each β -strand to have the same backbone conformations (based on identical chemical shifts). Thus the “a” and “b” conformers strictly occupy distinct strands, within which uniform backbone and χ_1 dihedral angles are found (Fig. 1J). Moreover, these two distinct β -strand types differ in their ψ and χ_1 dihedral angles
100 (Fig. 1K). After applying this additional filtering, 30 distinct unit cell architectures still remained possible. To investigate the structural stability of these 30 models, we carried out all-atom MD simulations in a fully periodic setting on them; see Methods for details. Only two of the candidate models proved stable throughout the simulations (Fig. 2B and Supplementary Fig. S2).

Notably, these two stable 3D lattices (called **M1** and **M2**) nicely capture known features of polyQ amyloid
105 structure. For instance, the experimental constraints offered by ssNMR dihedral angle measurements are open to more than one interpretation, as is typical for such experiments.^{5,42} Previously, this ambiguity and angular uncertainty yielded qualitative models useful for illustrative purposes, but not robust upon close inspection. Here, we have narrowed down the viable and physically plausible models to only two, with these models identifying specific narrow regions within the equivocal dihedral angle space (Fig. 2D).
110 In both models, the antiparallel β -sheet harbors strand-specific Gln structures occupying the side chain rotamers known as pt20° and mt-30°, according to Ref. 43. These conformations, as required, enable hydrogen bonding interactions between the side chains of the Gln conformers, in addition to typical strand-to-strand backbone hydrogen bonds (Fig. 2A,E). We used ssNMR to detect the protons involved in this hydrogen bonding ($N_\epsilon H$ protons), in the form of 2D 1H - ^{15}N HETCOR spectra (Fig. 2F). The 1H
115 chemical shifts are indistinguishable between the “a” and “b” Gln conformers. Structurally, this implies that the H-bonding distances should be very similar.^{44,45} This consideration was not imposed in the modelling, but is indeed observed in the stable structures (Supplementary Fig. S3), which demonstrate comparable side-chain–side-chain and side-chain–backbone distances in both the “a” and “b” conformers. In sum, we

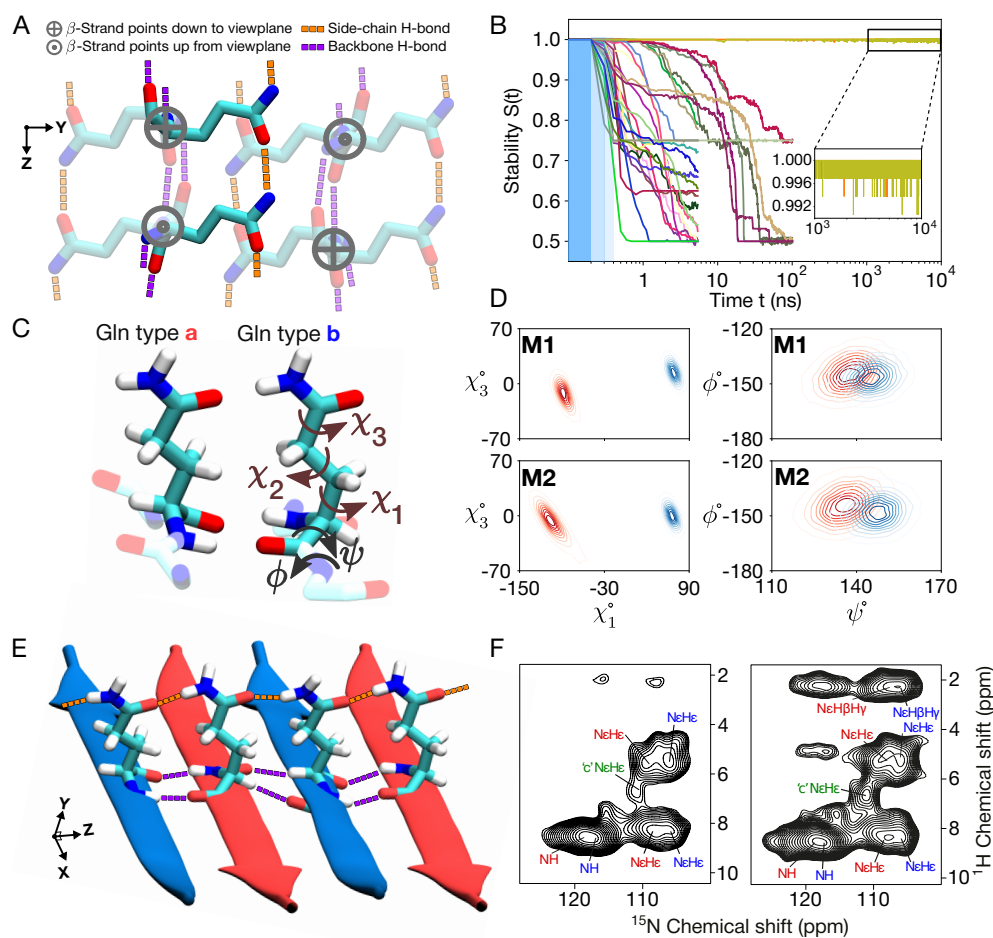


Figure 2. Structure of the internal polyQ amyloid core. (A) The eight-Gln building block used to generate polyQ core candidates. Six residues are shown faded out to bring the two non-faded in focus: These two are on adjacent β -strands of an antiparallel β -sheet that is stabilized not just by the backbone hydrogen bonds (purple), but also by the continuous chains of side-chain hydrogen bonds (orange). The latter are crucial for packing the polar glutamines into the waterless amyloid core. When generating the core candidates, all the χ_1 and χ_3 dihedral angles were independently rotated to explore all potential hydrogen bond networks. (B) Stabilities of the 30 experimentally-feasible polyQ core candidates (represented by color-coded lines) as a function of MD simulation time. The three blue-shaded sections indicate position restraints of 1 000, 500, and 100 kJ/mol over consecutive 100-ps periods. Notably, only two candidates maintain stability throughout the 10- μ s MD simulations; we denote these **M1** and **M2**. (C) Atomic-level structures of the type “a” and “b” Glns in **M2**. Gln dihedral angle names shown on “b”. (D) The side-chain χ_1 - χ_3 (left panels) and backbone ψ - ϕ (right) dihedral angle distributions of conformers “a” (red) and “b” (blue) for the final models **M1** (top) and **M2** (bottom). (E) Illustration of the inter-side-chain hydrogen-bond chain (orange) formed within β -sheet in **M2**. (F) ^1H - ^{15}N 2D ssNMR spectra of $\text{K}_2\text{Q}_{11}\text{pGQ}_{11}\text{K}_2$ peptide fibrils with a contact time of 100 μs (left) and of 350 μs (right) showing core and surface signals from the polyQ. Peaks from the core conformers “a” (red) and “b” (blue) are indicated. The side chain peaks that participate in hydrogen bonding ($\text{N}_\epsilon\text{H}_\epsilon$ and $\text{N}_\epsilon\text{H}_{\beta/\gamma}$) are marked. Note that the H_ϵ ^1H shifts are indistinguishable for the “a” and “b” conformers. Conformer “c” (green) refers to glutamines outside the core, on the surface of the fibril.

obtained valid atomic-resolution structures for the polyQ amyloid core, in two models that have many
 120 features in common. They formed the basis for further analysis and modeling, as will be discussed next.

2.2 Structure of polyQ₁₅ peptide fibrils

The above 3D models capture the internal core of polyQ protein fibrils, but lack the defined width and water-facing surface features of the real polypeptide fibrils. To model a proper experimental system, we modified the ‘infinite cores’ **M1** and **M2** into models of fibrils formed by the polyQ₁₅ peptide (Fig. 3A).
125 These widely studied^{5,21,24,46} peptides contain a 15-residue polyQ segment, flanked by charged residues to enhance solubility. Experimental analysis by ssNMR spectroscopy and X-ray diffraction has shown that atomic conformations in the polyQ₁₅ fibril core match those of the disease-relevant HttEx1 fibrils; e.g., the ssNMR signals from Gln within the D₂Q₁₅K₂ fibrils (Fig. 3A) and the Q44-HttEx1 fibrils (Fig. 1H) are indistinguishable. Using both the **M1** and **M2** models, we constructed D₂Q₁₅K₂ fibrils comprising
130 seven anti-parallel β -sheets, consistent with the 5.5- to 6.5-nm fibril width seen experimentally by TEM (Fig. 3A); Fig. 3B shows the **M2** D₂Q₁₅K₂ fibril in cartoon representation. We then subjected both polyQ₁₅ models to 1- μ s atomistic MD simulations in explicit water; for details, see Methods. The structure of the buried Gln residues within the core remained stable throughout the simulations (Supplementary Fig. S4).

There is a high degree of agreement between the χ_1 , χ_2 , and ψ dihedral angle distributions observed
135 in MD simulations of our polyQ₁₅ models and the ssNMR angle constraints obtained via HCCH (χ_1 , χ_2) and NCCN (ψ) experiments (Fig. 3C and Supplementary Fig. S5). As already noted, the “a” and “b” ssNMR signals reflect different χ_1 and ψ values in the two Gln conformers, but a similar χ_2 value. This is beautifully reproduced by the simulations of both polyQ₁₅ models (Fig. 3C). The largest deviation affects the **M2** model, in terms of its broader distribution of the conformer “a” χ_1 angle, which in part lies outside
140 the shaded region representing the (ambiguous) ssNMR constraint. Interestingly, experimental studies have shown⁵ that the conformer “a” is more dynamic than conformer “b”, reminiscent of the increased heterogeneity of this conformer in the MD simulations of both models (see also Supplementary Fig. S4). Also, the sheet-to-sheet and strand-to-strand distances within the models match well with repeat distances from X-ray diffraction (Supplementary Fig. S6). In sum, the MD simulations support the stability of these
145 polyQ amyloid core structures and recapitulate key data from multiple experimental techniques.

Our modeling reveals an important new feature of polyQ₁₅ fibrils, which, so far, has not been seen or studied: the solvent-facing external structure of the polyQ amyloid core. Fiber surface mediates interactions with cellular surroundings but also with any purposefully administered compounds, such as thioflavin-T or amyloid-specific tracers used in positron emission tomography (PET).⁷ The two outermost β -sheets
150 exposed to water (top and bottom sheets in Fig. 3B) allow us to compare the surface-exposed Gln residues versus the residues buried in the fibril core. Figure 4A highlights the water-exposed Gln side-chains (green) in the polyQ₁₅ model. These surface glutamines appeared as an averaged signal in ssNMR due to their increased dynamics (Fig. 2F green) and thus could not be resolved. Figure 4 shows the χ_1 - χ_2 and χ_3 - χ_2 dihedral angle distributions of the core Gln residues (panel B, gray) and compares them with the surface
155 residues (panel C, green) for model **M1** (see Supplementary Fig. S7 for **M2**). For the core residues, the rotamer states are similar to those discussed above (pt20° and mt-30°). For the surface residues, additional rotameric regions emerge, including Gln rotamers where χ_2 deviates from 180°, close to rotamer pm0°. ⁴³ Interestingly, surface residues do not show fully free side-chain motion and retain defined structural features that determine the molecular profile of the corrugated polyQ amyloid core surface.

160 2.3 Htt exon 1 fibril model

As already noted, the protein inclusions seen in HD patients and various HD model animals prominently feature the mutant HttEx1 protein fragment. Until now, no atomistic model of HttEx1 fibrils has been

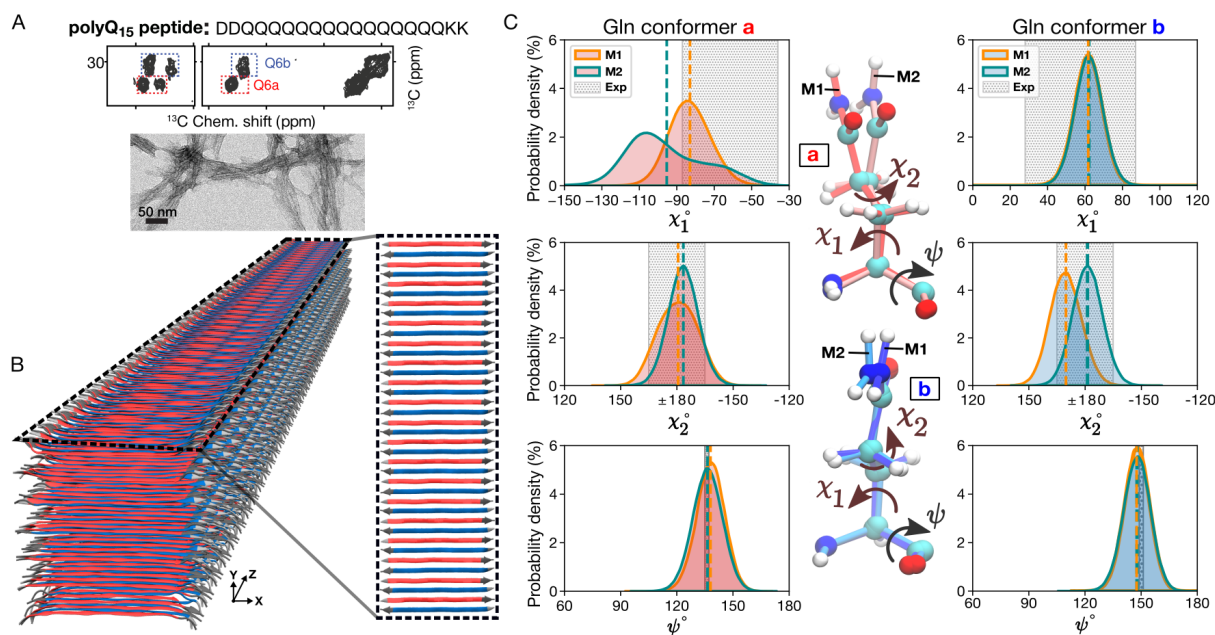


Figure 3. PolyQ peptide fibril structure. (A) Amino acid sequence of $D_2Q_{15}K_2$ peptide (top), $2D^{13}C-^{13}C$ ssNMR spectrum (middle) of $D_2Q_{15}K_2$ fibrils with single ^{13}C -labeled Gln (Q6), and negatively-stained TEM of the peptide aggregates (bottom). The ssNMR data adapted from Ref. 47. Signals from type "a" and "b" Gln conformers are shown in red and blue boxes, respectively. (B) 3D cartoon representation of the structural model of the $D_2Q_{15}K_2$ fibril. The alternation of β -strands of "a" (red) and "b" (blue) Gln conformers within a single sheet is shown on the right. (C) The χ_1 , χ_2 , and ψ dihedral angle distributions of "a" (left) and "b" (right) conformers in the context of the $D_2Q_{15}K_2$ fibril for the M1 (orange) and M2 (green) models; dashed vertical lines represent mean values. The data were obtained from 1- μ s MD simulations. Gray shading depicts the ssNMR constraints for the dihedral angles. The structures at the center show representative "a" (top) and "b" (bottom) conformers of the M1 and M2 models.

reported, although a diversity of schematic or cartoon-style models has been published over the years. Here, we build on our above-presented polyQ amyloid core model structures to construct an experiment-based
 165 molecular structure of Q44-HttEx1 fibrils. The Q44-HttEx1 construct is used to model the disease-relevant HD protein in a variety of experimental studies,^{18,19,34,36,48} as HD patients commonly have CAG repeat expansions that yield Htt proteins with approximately forty residues in their polyQ domain.² Experimentally, similar to our analysis of the polyQ₁₅ peptide fibrils above, the large majority of the polyQ residues are observed to be buried in the HttEx1 fibril core, in a multi-nm-size block-like core architecture.³⁴ Different
 170 from fibrils made from short polyQ segments, the polyQ-expanded protein fibrils have protofilament architectures^{19,34} where the polyQ segment adopts a β -hairpin structure;⁵ the Q44-HttEx1 fibrils feature a β -hairpin with a single turn and two ~ 20 residue-long β -strands. Solid-state NMR studies with site-specific amino acid labels have indicated that the ordered β -sheet core extends from the final N17-domain residue (F17) to the penultimate glutamine in the polyQ segment.^{18,25} Unlike the polyQ segment, the two polyQ-flanking segments of HttEx1 (see Fig. 1A) are solvent exposed, lack β -structure, and display increased motion and disorder.^{17,19,20,33} The short N-terminal N17 segment has been reported to adopt a random coil or α -helical structure, depending on context;^{25,49} in fibrils, Electron Paramagnetic Resonance (EPR) and ssNMR have shown the N17 segment to display partial order, and partly α -helical structure.^{17,19,25} The longer C-terminal proline-rich domain (PRD) is disordered, except for the polyproline-II (PPII) helices at
 180 the locations of the two oligoproline segments.

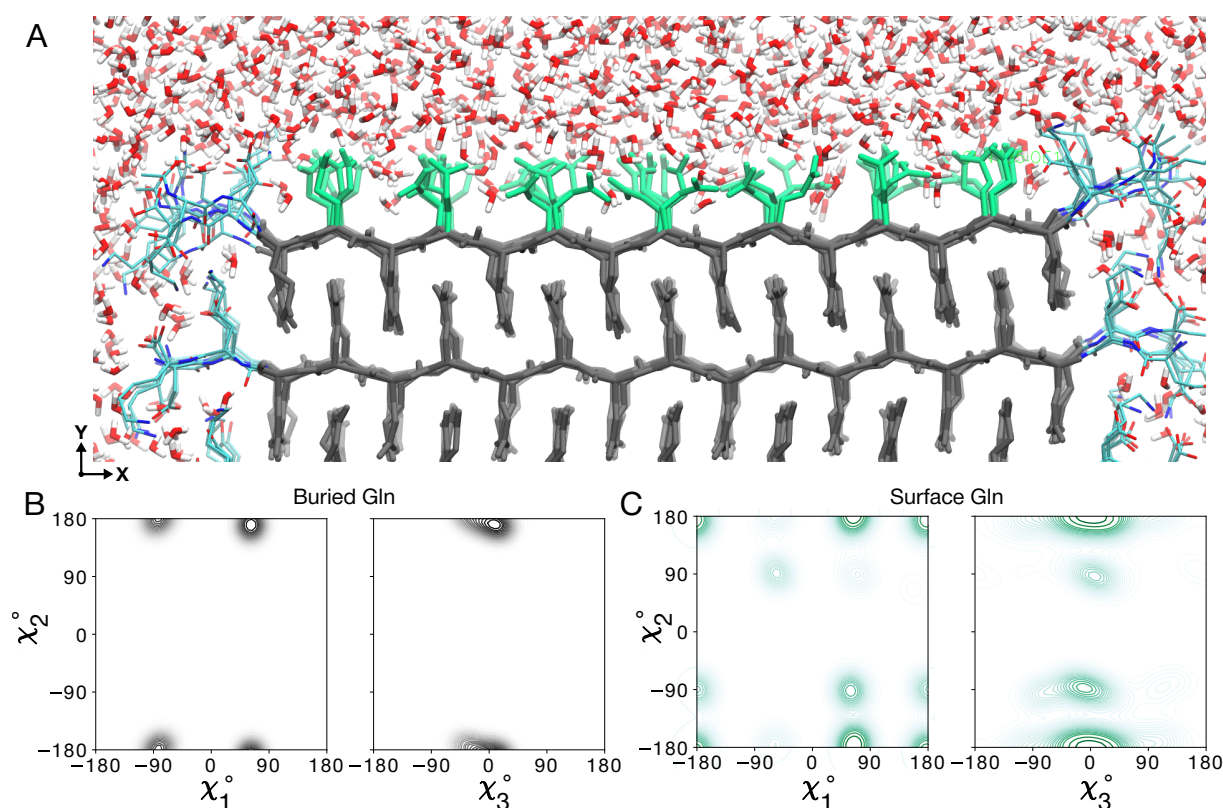


Figure 4. Water-facing surface of polyQ amyloid. (A) Atomistic MD snapshot of the $D_2Q_{15}K_2$ peptide fibril's polyQ surface in contact with water. Exposed and buried Gln residues are colored green and gray, respectively. Note how the Gln side-chains internal to the (model **M1**) amyloid core are well-ordered, while the water-facing side-chains display more mobility. (B) Side-chain dihedral angle distributions for the buried Gln residues, and (C) for the Gln residues on the fibril surface. The surface-facing residues show more disorder, but are nonetheless constrained to eight varyingly prominent specific rotamer states.

Using these experimental data as input, we constructed the Q44-HttEx1 core base-architecture after the schematic model described earlier.^{5,18,19,34} The 44-residue polyQ segment has a β -hairpin conformation with a tight β -turn, containing an even number of residues, and two long β -strands extending from F17 to the penultimate glutamine. A notable feature of polyQ protein aggregates is that the polyQ domains can come together in many different orientations and alignments, yielding a large propensity for heterogeneity and stochastic modes of assembly.^{18,33,34} Here, a configuration was selected to construct a model that mimics our previous schematic model,³⁴ with flanking domains evenly distributed on both sides of the fibril core. To this end, we attached the appropriate HttEx1 flanking domains to the terminal regions of the polyQ section, thereby achieving the formation of monomer building blocks as illustrated in the top panel of Fig. 5A. The thus constructed HttEx1 fibril was subjected to unrestrained all-atom MD simulations to assess its stability and monitor its structural dynamics. The fibril conformation obtained after a 5- μ s simulation is shown in Fig. 5A.

2.4 Structural analysis of HttEx1 fibril

The resulting structure (Fig. 5A) reveals interesting features that permit comparison to experimental studies. Firstly, consistent with the polyQ₁₅ fibril model, the HttEx1 polyQ core structure is found to be highly stable, with the bulk of the glutamine residues residing within the rigid and dehydrated core. A

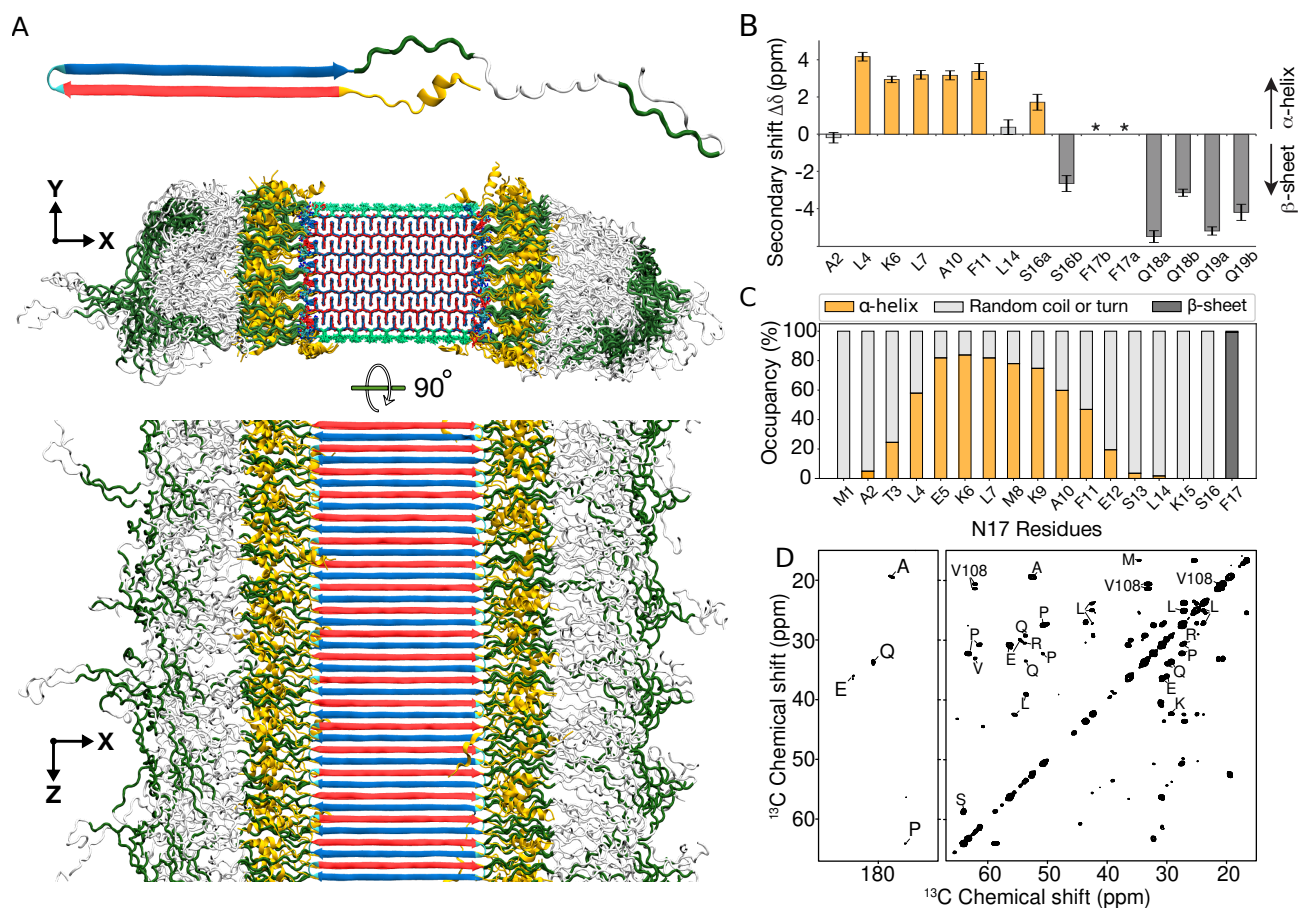


Figure 5. Structure of Q44-HttEx1 amyloid fibril. (A) Atomic-level structural model for mutant HttEx1 fibril. The top image shows a representative monomer within the fibril, with its β -hairpin polyQ segment to the left and the largely disordered flanking segments to the right. The middle image shows a cross-section, and the bottom image a side view of the fibril. The polyQ core is shown with the conformer-identifying red (for “a”) and blue (for “b”) β -strands, the N-terminal flanking segments yellow, and the C-terminal polyproline II helices dark green. Surface residues of the polyQ amyloid core are light green. (B) Secondary chemical shift values for residues in the N-terminal end, indicating local α -helix (positive values) or β -sheet (negative) conformations. Doubled peaks indicating multiple co-existing conformations are marked with letters a and b. The asterisks mark F17, for which a peak ($^{13}\text{C}_\beta$) was not detected, but other resonances indicate β -sheet structure. (C) Secondary structure distribution of the 17 residues in the N-terminal flanking domain during the last 500 ns of 5- μs MD simulation. (D) A 2D TOBSY ssNMR spectrum of Q44-HttEx1 fibrils, in which observed cross-peaks correspond to highly flexible residues outside the fibril core. Most, but not all, peaks originate from the C-terminal tail of the PRD. Spectrum was acquired at 600 MHz at 8.33 kHz MAS.

noteworthy observation lies in the β -turn conformation embedded within the β -hairpin structure: The simulation data underscores predominance of the type II turn over a type I' conformation (Supplementary Fig. S8), a finding that concurs with the experimental evidence gathered from ssNMR study of such compact turns.⁵ As for polyQ₁₅ fibrils, a minority population of glutamines (the light-green side-chains of the middle image in Fig. 5A) is exposed to the solvent; these surface side-chains, as analyzed in Fig. 4 above, show a semi-rigid behaviour, retaining a relatively high degree of structural order. The constrained dynamics of these solvent-exposed glutamines stand in large contrast to the dynamic disorder of both non-polyQ flanking domains—whose disposition and structure are of substantial interest, given that the flanking domains are known to govern key structural and pathogenic properties of the protein and its

aggregates.^{17,19,34,47,50–52} Figure 5A illustrates the high level of disorder that appears in the MD ensemble of the flanking domains, manifested in both the N17 and the PRD. Although such pronounced dynamics interfere with the detailed experimental study of the flanking domains, their appearance in simulations fits with the dynamic disorder of the flanking domains in HttEx1 fibrils observed in many experimental studies, whether based on EPR, EM, or ssNMR.^{15,17,19,20,33,47}

The secondary-structure preferences of N17, Q44, and PRD are summarized in Supplementary Fig. S9. There has been significant interest especially in the N17 segment, as it drives HttEx1 aggregation, but also harbors post-translational modifications that regulate HttEx1 (dis)aggregation and degradation.^{53–55} The fate of N17 in the fibrils has remained somewhat opaque, with seemingly conflicting reports of the presence and absence of (partial) α -helicity. The obtained HttEx1 fibril model may provide some insights, as its N17 segment displays a mixed secondary structure content, with much disorder (Fig. 5C and Supplementary Figs. S10–13). Virtually all N17 residues are seen to show some propensity for disorder, such that a subset of the proteins in the modelled fibril has an N17 devoid of α -helical structure (Fig. 6D). Yet, in close to half of the protein monomers, an α -helix is observed within N17. These findings match well to ssNMR analysis of the structure of N17 in fibrillar samples:^{19,25} On the one hand, signals from an α -helical N17 were detected, with helicity constrained to only part of the segment (Fig. 5B). The helical residues seen experimentally coincide remarkably well with the residues found to favour helicity in the MD fibril (Fig. 5C), providing further support for the validity of this structural ensemble. The observation that a significant part of N17 is not α -helical finds experimental support as well. Indeed, in some types of HttEx1 fibrils, no evidence of a structured or helical N17 has been found, indicating a propensity for substantial disorder.^{20,56} In other studies, N17 residues are observed to display both ordered and dynamic (ssNMR) signals, showing that different protein monomers have different degrees of (dis)order in the N17 segment.³⁴ This same finding is clearly recapitulated in the obtained MD fibril model: It displays an innate heterogeneity in structure and dynamics of the N17 segment, even among monomers in the same fibril.

C-terminal to the polyQ segment, the PRD is known to display a gradient of dynamics.^{17–20,34,56} Experimentally, the PRD is relatively rigid proximal to the polyQ core, whilst its very tail end is highly flexible. The latter is evidenced by ssNMR measurements of NMR relaxation parameters and dynamics-selective spectra (Fig. 5D).^{19,20,28} The MD simulations indicate that the mobility of the PRD is not only constrained by its attachment to the rigid polyQ core, but also by pronounced PRD–PRD intermolecular interactions. Prior work has inferred a propensity for such interactions, especially in context of filament–filament interactions.^{19,29,34} The current model suggest that such interactions are also prominent in structuring the flanking domains of isolated protofilaments.

A notable feature of the HttEx1 fibril structural ensemble that was not a priori expected is that the flanking domains are not showing much interaction with the polyQ amyloid core surface. Throughout the simulation, the flanking domains display substantial dynamics, but preferentially cluster together. This behaviour seems reminiscent of the type of liquid phase separation seen for, e.g., the nuclear pore complex, mediated by the proline-rich segments. This leaves the outer polyQ core surface easily accessible not only to solvent, but also amyloid-binding molecules such as thioflavin-T (ThT) as well as PET ligands.⁷ An in-depth and comprehensive visualisation of the described HttEx1 fibril structure can be further enhanced by viewing the accompanying movie in Extended Data 1.

2.5 Implications for huntingtin biology

Cryogenic electron tomography analysis of cellular httEx1 aggregates have indicated that the fibrils may be structurally similar to those formed by purified proteins,¹⁵ which would suggest that our model also

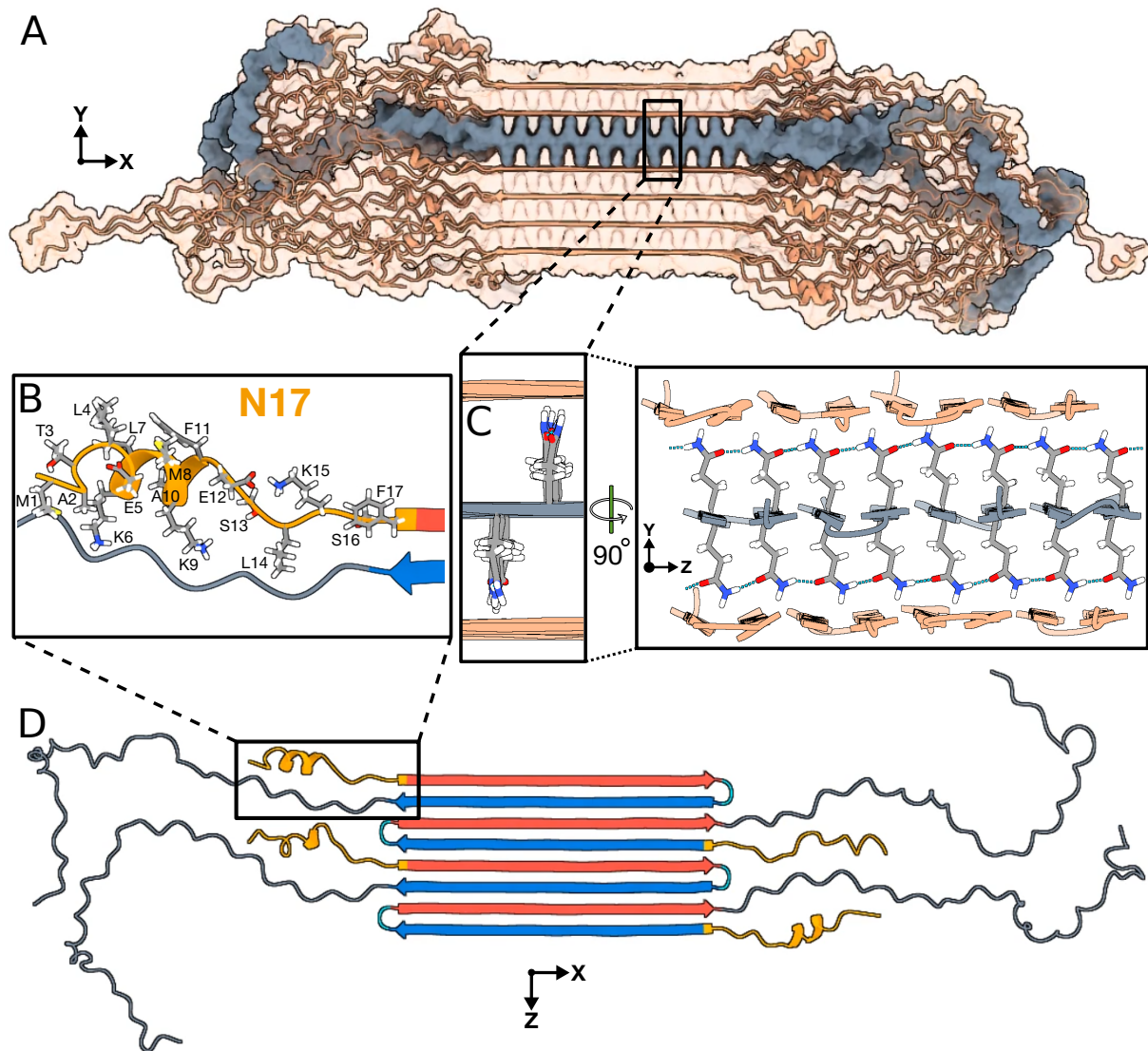


Figure 6. Structural analysis of the Q44-HttEx1 amyloid fibril. For an illustrative 3D exploration of Q44-HttEx1 fibril structure, see the associated video content available in Extended Data 1. (A) A graphical depiction of the HttEx1 fibril. The region shaded in gray denotes a single sheet within the fibril's architecture. (B) An atomic view of the N17 domain within the fibril, naming the specific amino acids. (C) An atomic depiction of the glutamine side-chains within the fibril. The high stability of the fibril structure is primarily attributed to the extensive hydrogen bonding interactions among the glutamines, as depicted in the right panel. (D) Side view representation of the β -sheet highlighted in panel A. A quartet of HttEx1 monomers is visible. The polyQ is color-coded for the type "a" (red) and "b" (blue) strands; the tight β -turn is cyan. The N17 and PRD domains are orange and gray, respectively. Note the structural variation between different monomers in the same fibril sheet, including in particular the range of helical content in the N17.

applies to cellular inclusions. Notably, structural features of our model appear to match, and thus rationalize, reported biological properties of HttEx1 fibrils. In particular, the surface-accessible features of amyloid fibrils are crucial for their biological, possibly pathological, properties. In our fibril architecture, the N17 segment is found to reside outside the fibril core, where it displays conformational and dynamic disorder. Despite these dynamics it is, however, tightly enclosed by the C-terminal PRD domains. Consequently, residues in N17 may be solvent accessible, but are nonetheless largely inaccessible to larger macromolecules such as chaperones, kinases, ubiquitinases, and other potential N17 interaction partners. Although in Fig. 6B

the known phosphorylation sites T3, S13, and S16 may appear accessible, Fig. 6A shows this monomer in context, underlining that the N17 segment is fully surrounded by the longer PRD segments. Similarly, the only HttEx1 ubiquitination sites involve lysines K6 and K9 in N17, rendering them inaccessible. The buried nature of N17 also rationalizes the low level of engagement by the TRiC chaperone:^{31,57} Although
260 known to bind this part of HttEx1, prior EM studies have shown TRiC to be unable engage HttEx1 fibrils, except near the fibril ends. N17 is also implicated in membrane interactions, such that its preferential exposure at fibril ends may explain the latter to be engaged with the ER membranes in Fig. 1C.^{52,58,59}

Naturally—although we expect the obtained fibril structure to be a good representation of the fibrils present in our samples and also to have strong predictive and descriptive qualities for cellular HttEx1
265 aggregates—it cannot be excluded that cellular factors (ranging from chaperones to membrane interactions) may modulate the aggregation mechanism to the extent that the mature fibril architecture differs from the one obtained *in vitro*. Yet, unlike for other amyloid proteins, a remarkable feature of the HttEx1 ssNMR studies is that multiple groups have studied a variety of HttEx1 fibrils and always found the same signature spectra that are connected to the structural parameters used to construct our model. Indeed, there is little
270 evidence for a qualitatively different fibril architecture. Thus, we are inclined to expect that while cellular conditions may change certain details in the fibril structure (for example, the fibril polymorph formed in presence of membranes may feature more exposed N17 segments⁵²), they would not fundamentally change the protofilament architecture.

Yet, we do need to note that polyQ-based protein aggregates display a persistent and characteristic
275 structural heterogeneity that is impossible to fully capture in practical MD simulations. Prior studies have discussed the predictable propensity for polyQ segments to self-assemble in a somewhat disordered or stochastic fashion, due to the lack of sequence variation.^{5,18,60} For instance, a polyQ chain extending an existing fibril can be added in different orientations. This variability is to some extent already displayed in our model: The alternating monomers in the single sheet in Fig. 6D have differently organized β -hairpins
280 in their polyQ segment and their N17 segments on opposite sides of the filament. What is explicitly missing from our model is that incoming polyQ segments can add to the fibril with register mis-alignments, resulting in shorter or longer β -strands in slightly different sequence positions, without a major energetic cost. This would be expected to yield fibril cores with varying fibril widths, local defects, and protein-to-protein structural variations^{18,33,60} not accessible to more canonical amyloid fibrils formed by other proteins. This
285 means that polyQ protein fibrils are expected (and observed^{33,34}) to display inherent structural variations between fibrils in a single sample, and even within single fibrils. Thus it appears that, like snowflakes, each HttEx1 fibril is unique (which prohibits the canonization of a single definitive protofilament structure) but still characterizable by well-defined structural features. This indistinguishability in local (atomic) structure of differing fibril architectures manifests for instance in the near-identity of ssNMR peak positions reported
290 for polymorphic HttEx1 fibrils.

Indeed, like other amyloids, polyQ proteins form different fibril polymorphs depending on experimental conditions.^{19,29,61} This structural variation imbues the fibrils with different degrees of cytotoxicity.^{29,61,62} Notably, the HttEx1 polymorphs often reflect a type of 'supramolecular' polymorphism, with different
295 supramolecular assemblies formed from similarly structured protofilaments.³⁴ The current model is expected to illuminate the atomic level conformation of one such protofilament, as it is based on experimental constraints that define its structure. While we do not explicitly probe the conformations of supramolecular polymorphs, the clustering of the dynamic flanking domains, seen for our single filaments, likely has some relevance to the inter-fibril interactions—thought to be mediated by the PRD segments as well.

300 Finally, let us briefly discuss the implications of our models for HttEx1 fibrils formed by proteins
with longer (and shorter) polyQ segments. Our HttEx1 fibril modelling focused on the HD-relevant Q44
that falls into the regime of common expansion lengths seen among patients. Yet, famously, mutant
proteins can differ widely in their polyQ lengths. As illustrated in the polyQ₁₅ peptide fibril, and discussed
elsewhere,^{37,56,62} proteins with shorter polyQ lengths can still form fibrils (at least *in vitro*). However, in
305 such fibrils the polyQ segment may not form a β -hairpin, but instead occupy a single extended β -strand.
Naturally, this would modulate the disposition of flanking segments on the fibril surface. Nonetheless, the
qualitative architecture would remain unchanged. Conversely, HttEx1 with longer polyQ lengths, such
as those associated with juvenile HD, would be expected to form amyloid cores featuring multiple turns,
unlike the single-turn structures analyzed here for Q44-HttEx1.

310 **3 Conclusions**

In this report we have described our construction of experimentally informed and validated structural
models of polyQ₁₅ and Q44-HttEx1 amyloid fibrils. These models provide the best atomistic views of these
disease-associated protein inclusions to date, derived from a multi-technique structural analysis through
integrative modelling. The obtained HttEx1 structure rationalizes a variety of experimental findings with
315 notable biological and biomedical implications. The polyQ segment is mostly buried within the fibril,
but the model reveals the structural and dynamical features of the minority of solvent-facing residues.
These surface residues have proved challenging for experimental study, necessitating tailored and targeted
approaches in, e.g., ssNMR.³⁶ A better structural understanding of this special polyQ surface will be
useful in efforts to design polyQ-amyloid-specific binders, e.g., for PET imaging.⁷ The visualization
320 of the dynamically and structurally heterogeneous flanking domains enhance our understanding of their
accessibility in the fibrils and pave the way for more in-depth analyses of their role in intracellular
interactions with proteins and organelles.

4 Methods

Protein production and fibrillation

325 Mutant huntingtin exon 1 with a 44-residue polyQ core was expressed as part of a maltose binding protein
(MBP) fusion protein as previously reported.^{18,19} The fusion protein MBP-Q44-HttEx1 was expressed
in *Escherichia coli* BL21 (DE3) pLysS cells (Invitrogen, Grand Island, NY). Uniformly ¹³C and ¹⁵N
labeled MBP-Q44-HttEx1 protein was expressed with ¹³C D-glucose and ¹⁵N ammonium chloride for
MAS ssNMR studies. Then, cells were pelleted at 7 000 g, resuspended in phosphate buffered saline (PBS),
330 pH 7.4 and lysed in presence of 1 mM phenylmethanesulfonyl fluoride (PMSF) by a HPL 6 maximator
(Benelux BV, The Netherlands). After that, cells were centrifuged at 125 000 g for 1 h using an Optima
LE-80K ultra-centrifuge (Beckmann Coulter). The supernatant was filtered over Millex-GP syringe-driven
0.22 μ m PES membranes (Millipore Sigma, Burlington, MD). The MBP-Q44-HttEx1 protein was purified
by fast protein liquid chromatography (FPLC) using a 5 ml HisTrap HP nickel column (GE Healthcare,
335 Uppsala, Sweden) with 0.5 M imidazole gradient (SKU I5513-100G, Sigma, St. Louis, MO) on an AKTA
system (GE Healthcare, Chicago, IL). The imidazole was removed from the purified protein using an
Amicon Ultra centrifugal filter with a regenerated cellulose membrane (Millipore Sigma, Burlington, MA).
At least 3 washes with imidazole-free PBS buffer were done. Protein concentration was calculated from the
absorbance at 280 nm. According to ProtParam tool by ExPasy⁶³ the extinction coefficient of the fusion

340 protein is $66\ 350\ \text{M}^{-1}\text{cm}^{-1}$. Protein aggregation was initiated by addition of Factor Xa protease (SKU PR-V5581, Promega, Madison, WI) at 22°C , in order to cleave off the MBP fusion tag^{19,34} and release Q44-HttEx1. After 3 days, the obtained mature fibrils were washed with PBS to remove the cleaved MBP.

Transmission Electron Microscopy (TEM)

345 Transmission electron microscopy (TEM) was performed on mature $\text{D}_2\text{Q}_{15}\text{K}_2$ peptide fibrils. The fibrils were re-suspended in MiliQ and then $5\ \mu\text{l}$ of the sample was deposited on the plain carbon support film on 200 mesh copper grid (SKU FCF200-Cu-50, Electron Microscopy Sciences, Hatfield, PA). The grid was glow discharged for 0.5–1 min before adding the sample. After 30 s of the sample deposition, the excess MiliQ was removed by blotting, and immediately the negative staining agent 1% (w/v) uranyl acetate was applied. After 0.5–1 min, the excess stain was removed and the grid was air dried. The images were
350 recorded on a Tecnai T12 electron microscope.

NMR experiments

The hydrated $\text{U-}^{13}\text{C}$, ^{15}N -labeled Q44-HttEx1 fibrils were packed into a 3.2 mm ssNMR rotor (Bruker Biospin) using a previously described packing tool.⁶⁴ The fibrils were packed in the rotor by centrifugation at $\approx 130\ 000\ \text{g}$ in a Beckman Coulter Optima LE-80K ultracentrifuge equipped with an SW-32 Ti rotor
355 for 1 hour. Experiments were performed on a wide-bore Bruker Avance-I 600 MHz (14.1 T) spectrometer or Bruker Avance Neo 600 MHz (14.1 T) spectrometer, using triple-channel (HCN) 3.2 mm MAS EFree probes. All experiments were acquired using two-pulse phase modulated (TPPM) proton decoupling of 83 kHz during acquisition⁶⁵. The 2D ^{13}C – ^{13}C DARR experiments⁶⁶ were performed using a $3\text{-}\mu\text{s}$ 90° pulse on ^1H , $4\text{-}\mu\text{s}$ 90° pulses on ^{13}C , a ^1H – ^{13}C CP contact time of 1 ms at 275 K, a DARR mixing time
360 of 25 ms, and a recycle delay of 2.8 s. 2D ^{13}C – ^{15}N NCO experiments were done at 277 K using a $3\text{-}\mu\text{s}$ 90° pulse on ^1H , $8\text{-}\mu\text{s}$ 180° pulse on ^{13}C , ^1H – ^{15}N contact time of 1.5 ms, ^{15}N – ^{13}C contact time of 4 ms and recycle delay of 2.8 s. 2D ^{13}C – ^{15}N NCA experiments were done at 277 K using a recycle delay of 2.8 s, a $3\text{-}\mu\text{s}$ 90° pulse on ^1H , $8\text{-}\mu\text{s}$ 180° pulse on ^{13}C , 1.5 ms and 4 ms ^1H – ^{15}N and ^{15}N – ^{13}C contact times, respectively. In NCA and NCO experiments, the power levels for ^{15}N and ^{13}C during N–C transfer steps
365 were 50 kHz and 62.5 kHz, respectively. The 2D refocused-INEPT ^{13}C – ^{13}C 2D spectrum was obtained with total through-bond correlation spectroscopy (TOBSY; $\text{P}9\frac{1}{3}$) recoupling, measured at MAS rate of 8.3 kHz, using a 6 ms of TOBSY mixing time, a $3\text{-}\mu\text{s}$ 90° pulse on ^1H , $4\text{-}\mu\text{s}$ 90° on ^{13}C , at a temperature of 275 K.⁶⁷ The ^1H – ^{15}N heteronuclear correlation (HETCOR) experiments were done using a MAS rate of 13 kHz, 100 and 350 μs CP contact times, 4 s recycle delay, a $3\text{-}\mu\text{s}$ 90° ^1H pulse, and at 275 K temperature.
370 100 kHz homonuclear FSLG ^1H decoupling was applied during the t_1 evolution time.

MD simulations

Calculations were carried out on the fast, free, and flexible Gromacs engine.⁶⁸ All systems were first energy-minimized using steepest descent with one conjugate gradient step every 100 steps, then equilibrated through a 200-ps MD simulation in the NVT ensemble with positional restraints (1000 kJ/mol) on heavy
375 atoms, followed by three 100-ps NPT runs with positional restraints (1000, 500, and 100 kJ/mol) on heavy atoms. The production MD simulations were done in the NPT ensemble, obtained through the Bussi–Donadio–Parrinello⁶⁹ thermostat ($T = 300\ \text{K}$, $\tau_T = 0.2\ \text{ps}$) and the isotropic (for the polyQ lattice systems) or semi-isotropic (Q_{15} and HttEx1 systems, xy and z coupled separately) Parrinello–Rahman⁷⁰ ($P = 1\ \text{bar}$, $\tau_P = 2\ \text{ps}$, $\kappa_P = 4.5 \times 10^{-5}\ \text{bar}^{-1}$) barostat. The van der Waals interactions were switched off between

380 1.0 and 1.2 nm; long-range electrostatics were treated via Particle Mesh Ewald^{71,72} with fourth-order interpolation, a real-space cut-off at 1.2 nm, and size-optimized fast Fourier transform parameters (grid spacing of roughly 0.16 nm). Covalent bonds involving hydrogens were constrained to their equilibrium lengths by (fourth-order double-iteration) parallel linear constraint solver (P-LINCS)⁷³. Timestep was 2 fs, Verlet neighbour lists updated every 20 fs with the neighbour list radius automatically determined.

385 PolyQ amyloid core lattices

To build atomistic-resolution models of the internal structure of the polyQ amyloid core, we assumed it to comprise a lattice of antiparallel β -sheets stacked such that the Gln side-chains interdigitate. The pairs of side-chain dihedral angles (χ_1, χ_3) were set to satisfy the known characteristic of the interdigitating polyQ amyloid core:^{5,23} the existence of side-chain–side-chain hydrogen bond interactions. We took the fibril-axis
390 direction to align with the Cartesian coordinate z , and to be perpendicular to the β -strands (aligned with x). To construct the 3D lattice, we considered a minimal “unit cell” consisting of eight Gln residues arranged in a $2 \times 2 \times 2$ pattern (Fig. 2A): Along x , the unit cell contains a minimal peptide chain segment of two amino acids ($\underline{2} \times 2 \times 2$), representing the alternating (‘odd/even’, i.e., pointing ‘above’ and ‘below’ the β -sheet plane) residues of the β -strand. Along z , to describe an antiparallel β -sheet, minimum two
395 β -strands are needed ($2 \times 2 \times \underline{2}$); in contrast to parallel in-register sheet structures that could be represented with a single repeating β -strand. Along y , the unit cell contains two distinct neighbouring β -sheets to permit and probe distinct sheet–sheet interfaces ($2 \times \underline{2} \times 2$). Figure 2A illustrates how each eight-Gln unit cell contains four Gln–Gln pairs, which establish backbone hydrogen bonds (shown in purple) as well as side-chain hydrogen bonds (in orange). All the hydrogen bonds are aligned roughly along the fibril axis, z .
400 For each Gln–Gln pair, there are 8 distinct classes of (χ_1, χ_3) orientations that permit side-chain–side-chain hydrogen bond chains along z (see Supplementary Fig. S1). As there are 4 Gln–Gln pairs in the unit cell, there are $8^4 = 4096$ possible plausible atomistic structures of the 8-Gln unit cell. Accounting for rotational and translational symmetry of the 3D lattice reveals, however, that at most 1280 of these structures are unique. An important further consideration is that ssNMR experiments conclusively demonstrate that
405 consecutive residues within each β -strand must adopt the same backbone conformations, as indicated by identical chemical shifts. As a result, each distinct strand exclusively contains either type “a” or type “b” Gln residues (Fig. 1I,J). After applying this filter, the number of possible distinct unit cells is reduced to 30. For each of these 30 unit cells candidates, we performed all-atom MD simulations. The fully periodic MD simulation box contained 40 identical unit cells, that is, a total of $8 \times 40 = 320$ Gln residues, organized as a
410 stack (along y) of 4 antiparallel β -sheets, with each sheet comprising 8 β -strands of periodic (along x) Q₁₀ peptides.

Acknowledging the limitations of a classical mechanics approximation (force field) of an inherently quantum system, we employed three state-of-the-art MD force fields in parallel: AMBER14SB,⁷⁴ OPLSAA/M,⁷⁵ and CHARMM36m.⁷⁶ Gromacs version 2018.3 was used. During the 10- μ s production
415 runs, the stability of the 30 structural candidates was evaluated at 5 ns, 100 ns, 200 ns, and 1 μ s. At each of these time points, only the candidates that maintained stability above 0.9 (see Eq. (1)) were further continued. The stability $S(t)$ of the given structural candidate at a simulation time t was defined based on the $\chi_1(t)$ and $\chi_3(t)$ dihedral angles compared to the initial energy-minimized MD structure:

$$S(t) = \frac{N_1(t) + N_3(t)}{2N}, \quad (1)$$

where $N = 320$ is the total number of residues, and $N_i(t)$, $i = \{1, 3\}$ is the number of residues whose
420 $\chi_i(t)$ is within 90° of its initial value at time t .

Solvated polyQ₁₅ fibrils

Q₁₅ peptide fibrils were constructed from each of the two stable ssNMR-verified core models, **M1** and **M2**. Two successive aspartic acid (DD) residues were added to the N-terminus and two lysines (KK) to the C-terminus of the Q₁₅ peptide to mimic the D₂Q₁₅K₂ peptide widely studied by experiments.^{5,21,24,46}
425 The N-terminus was further capped with an acetyl (Ace) group, and the C-terminus was set uncharged (with -COOH capping) to match the peptides used in our experiments. A 7-sheet fibril structure was constructed, with each sheet comprising eight Ace-D₂Q₁₅K₂ peptides. The simulation box was set up to form a quasi-infinite fibril (along z) under periodic boundary conditions and solvated with $\sim 9\,800$ water molecules in a cuboid box of $\sim 11 \times 11 \times 3.8$ nm³.

430 State-of-the-art MD force fields AMBER14SB,⁷⁴ OPLSAA/M,⁷⁵ and the TIP3P⁷⁷ water model were used for calculations carried out on the Gromacs version 2021.3. Production run length was 1 μ s.

Solvated Q44-HttEx1 fibrils

The ssNMR data suggest that the HttEx1 aggregates display the same spectral patterns observed in polypeptide polyQ fibrils.^{5,37} This indicates the presence of a common atomic polyQ core structure among
435 them. Consequently, we constructed an atomistic-resolution structure of the mutant HttEx1 fibril utilizing the polyQ core model described in the preceding section. In contrast to fibrils composed of short polyQ segments, the Q44-HttEx1 fibrils exhibited a distinct structural characteristic, consisting of a β -hairpin structure with a single turn. Hence, the constructed core domain of our Q44-HttEx1 encompasses a stack of seven antiparallel β -sheets composed of 44-residue polyQ hairpins (Q44). The hairpin arms
440 consisted of “a” and “b” β -strands connected by a two-residue β -turn modeled based on a type I’ tight turn, known for its strong preference towards adopting a β -hairpin structure.⁷⁸ The N-terminus of HttEx1, comprising 17 residues, was modeled utilizing the crystal structure of a single Htt(1–17) peptide in complex with the C4 single-chain Fv antibody (PDB ID: 4RAV).⁴⁹ The residue F17 was included within the β -sheet core, pairing with the penultimate glutamine residue, as supported by the findings of ssNMR
445 investigations.²⁵ The PRD domain of the HttEx1, including 50 residues (P62–P111), was modelled as an end-to-end-distance-maximized random coil, with the two oligoproline (P62–P72 and P90–P99) segments in a polyproline-II-helix conformation. The termini were set to NH₃⁺ and COO⁻ for the N- and C-terminal, respectively. The turns of neighboring Q44 hairpins were positioned on opposite sides of the fibril (Fig. 5A). The system was solvated with $\sim 840\,000$ water molecules, resulting in a total of $\sim 2\,800\,000$ atoms, in a
450 cuboid box of $\sim 37 \times 37 \times 19$ nm³.

The production MD simulations were conducted for a duration of 5 μ s on Gromacs version 2021.4 with AMBER14SB⁷⁴ protein force field and TIP3P⁷⁷ water.

5 Data Availability

The data that support the findings of this study are openly available at Zenodo (DOI:
455 10.5281/zenodo.8143752).

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7 Competing Interests

The authors declare no competing interests.

Supplementary Material

1 SUPPLEMENTARY FIGURES

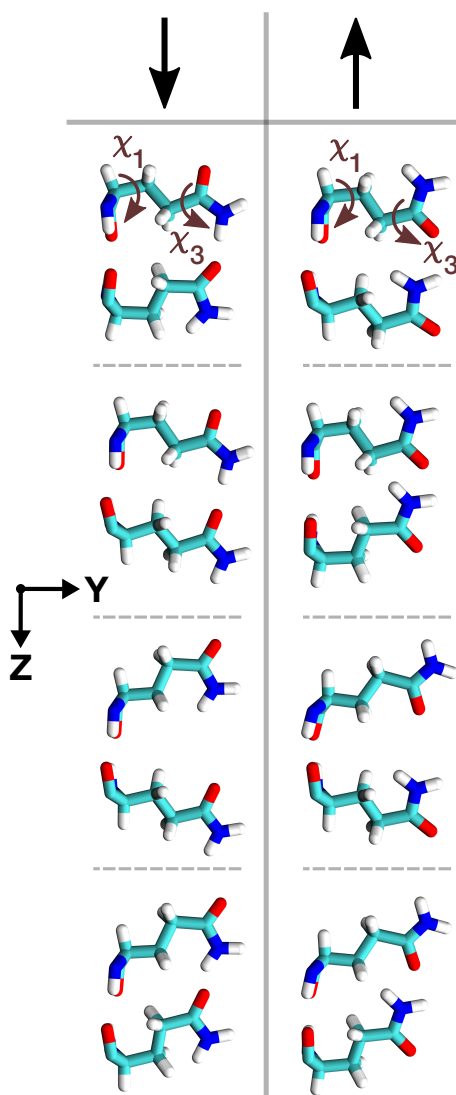


Figure S1. Schematic visualization of the eight (χ_1, χ_3) rotamer classes of a Gln-Gln pair, in which a side-chain-side-chain hydrogen bonding interaction could take place. The side-chains of the pair are located on neighbouring β -strands of an antiparallel β -sheet inside the polyQ amyloid core. Depending on the H-bond direction, the eight distinct structural classes are categorized into two forms: ↓ and ↑.

Supplementary Material

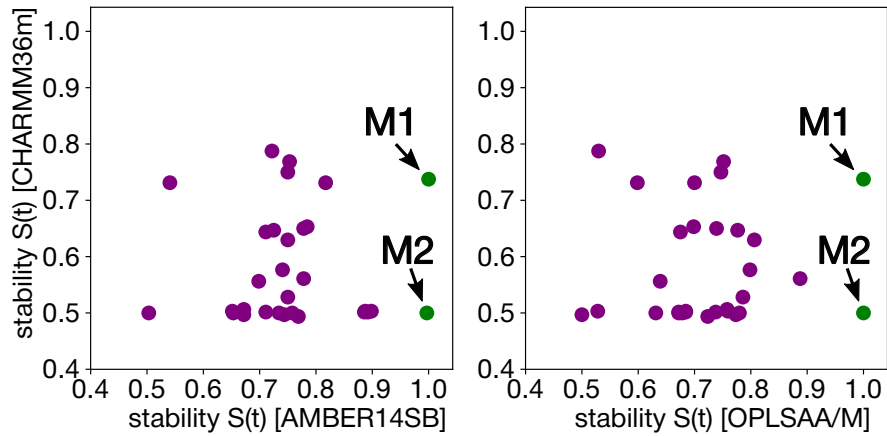


Figure S2. The stabilities $S(t)$ (see Eq. (1) in Methods of the main text) of the 30 candidate structures for the polyQ amyloid core examined with up-to-1- μ s molecular dynamics simulations using three distinct force fields: CHARMM36m (Huang et al. (2017)), AMBER14SB (Maier et al. (2015)) and OPLSAA/M (Robertson et al. (2015)). Strikingly, both **M1** and **M2** models demonstrated complete stability when simulated with AMBER14SB and OPLSAA/M, while in CHARMM36m none of the 30 candidates did.

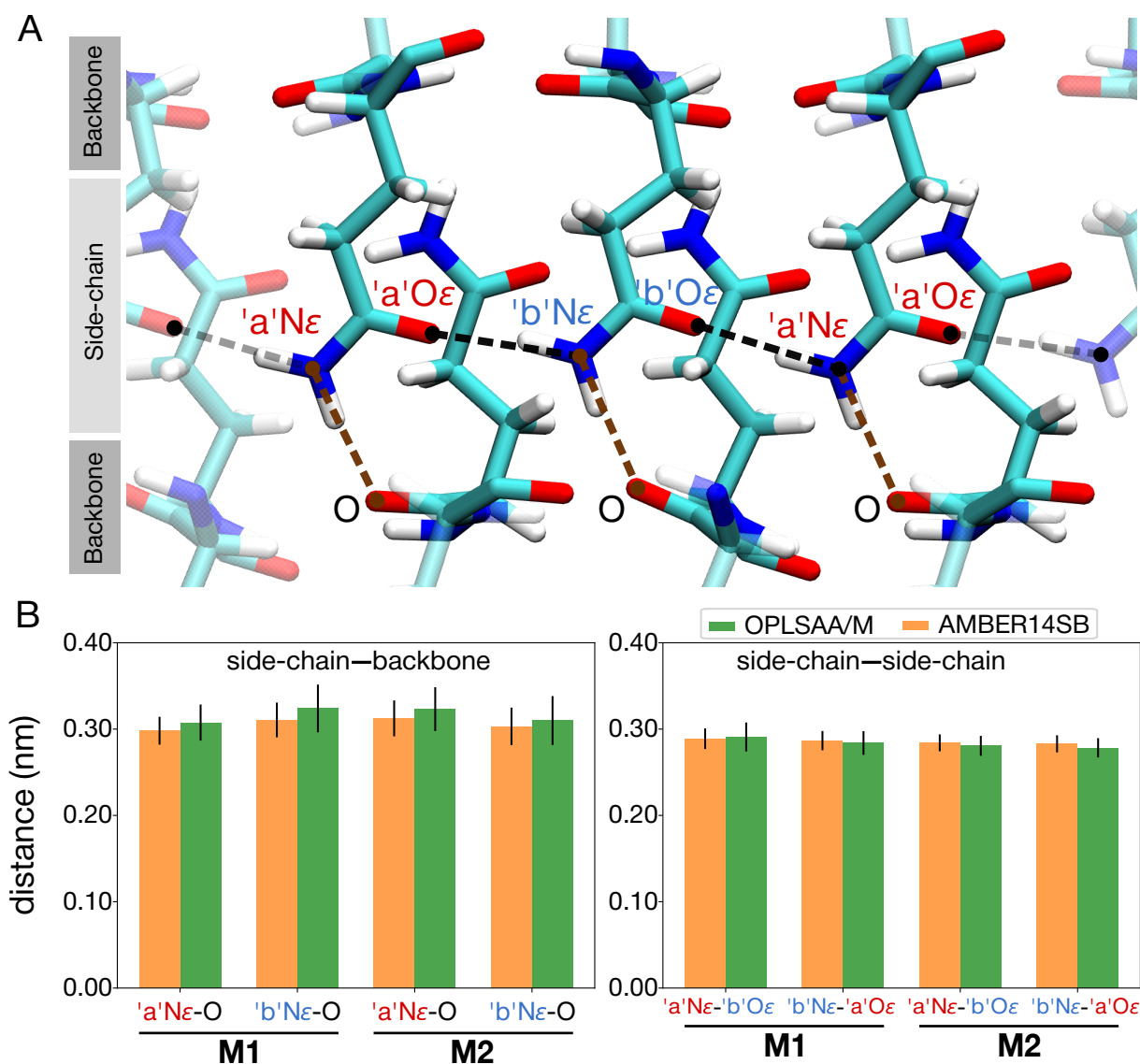


Figure S3. (A) Spatial arrangement of the side-chain and backbone atoms in the polyQ amyloid core model **M2**. Dashed lines represent the distances between the N_{ϵ} atoms of the “a” and “b” types of Gln residues and their nearest backbone O atoms (brown), as well as the N_{ϵ} – O_{ϵ} distances between side-chains (black). (B) Side-chain–backbone distances (left panel), i.e., the distances between the N_{ϵ} atoms of the “a” and “b” types of Gln residues and their closest backbone O atoms. Side-chain–side-chain distances (right panel), i.e., the N_{ϵ} – O_{ϵ} distances between the side chains, either from “a” to “b” or from “b” to “a”. The calculations are presented for both the **M1** and **M2** models, using two force fields: OPLSAA/M (green) and AMBER14SB (orange); error bars show standard deviation.

Supplementary Material

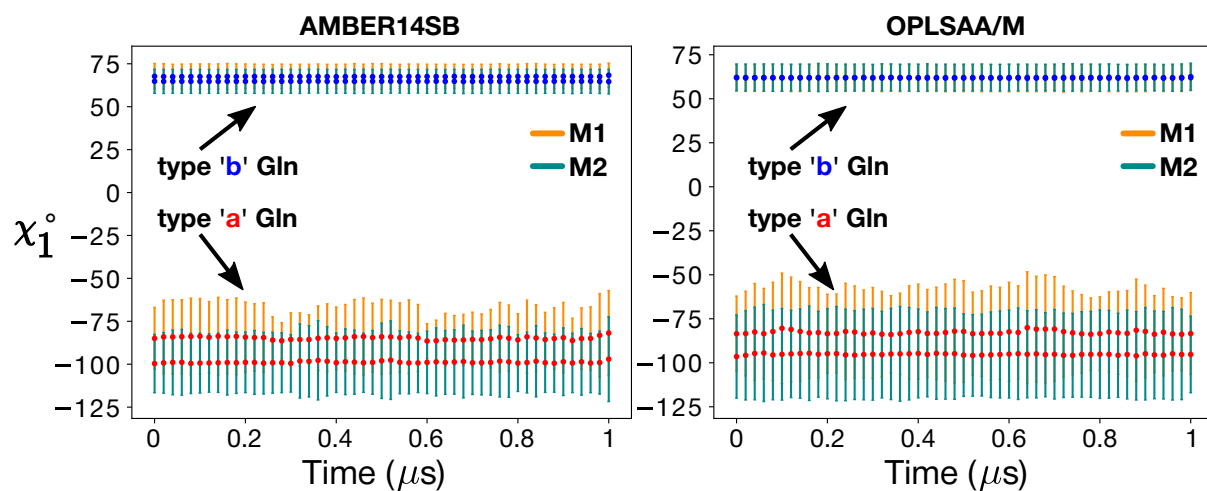


Figure S4. Both polyQ₁₅ models **M1** (orange) and **M2** (green) remained stable throughout 1- μs MD simulations in both the AMBER14SB (left) and OPLSAA/M (right) force fields. Shown are the mean χ_1 dihedral angles in type “a” (red) and “b” (blue) Gln side-chains, calculated over consecutive 20-ns windows; error bars show standard deviation.

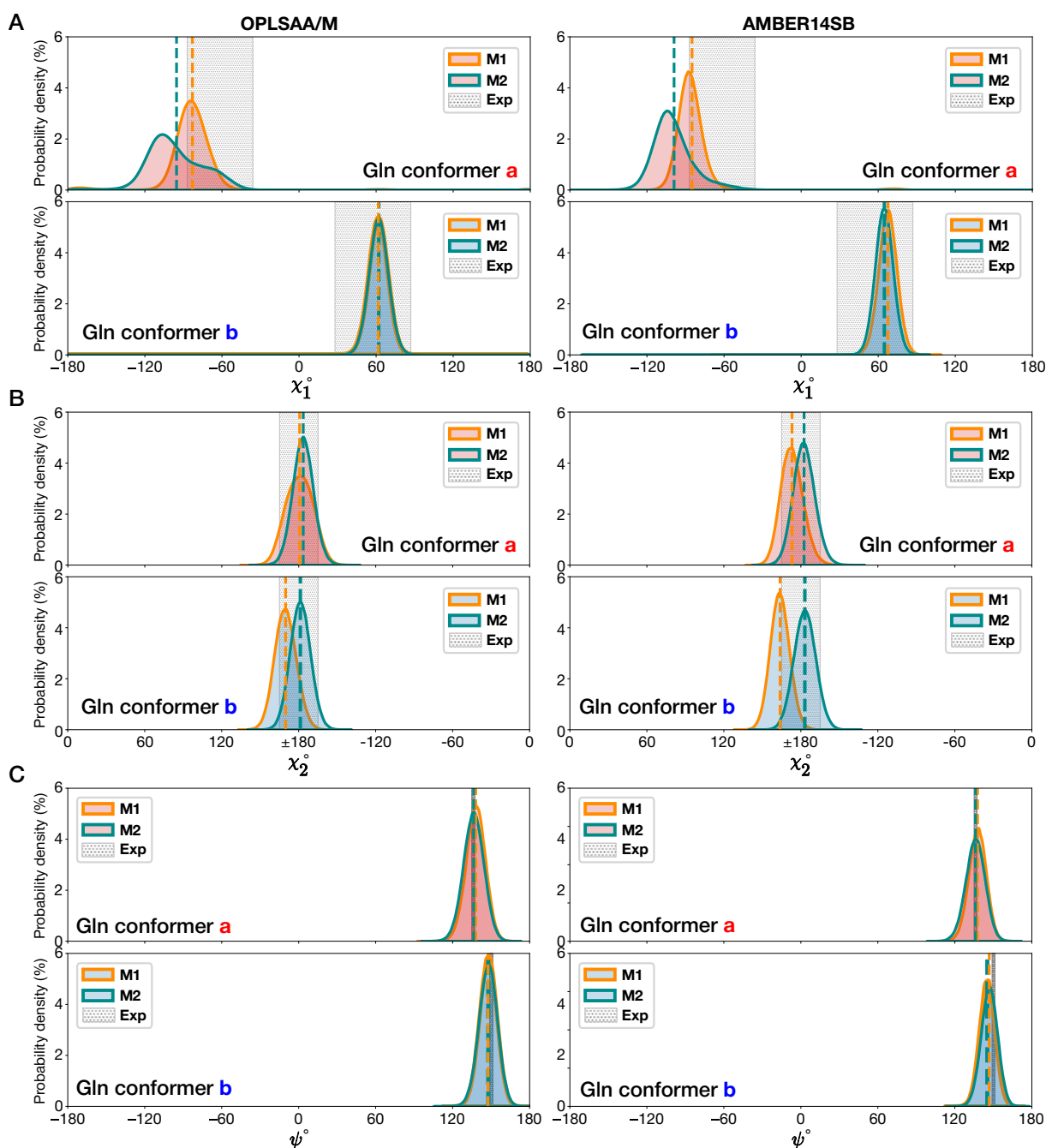


Figure S5. Distributions of the (A) χ_1 , (B) χ_2 , and (C) ψ dihedral angles in the M1 (orange lines) and M2 (green lines) models of the D₂Q₁₅K₂ fibril obtained from 1- μ s MD simulations using the OPLSAA/M (left) and AMBER14SB (right) force fields. The upper panels show the type “a” conformers (red-shaded distributions) and the lower panels the type “b” (blue-shaded). The gray-shaded regions represent the ssNMR-informed constraints. The dashed vertical lines depict the mean values of the corresponding dihedral angles.

Supplementary Material

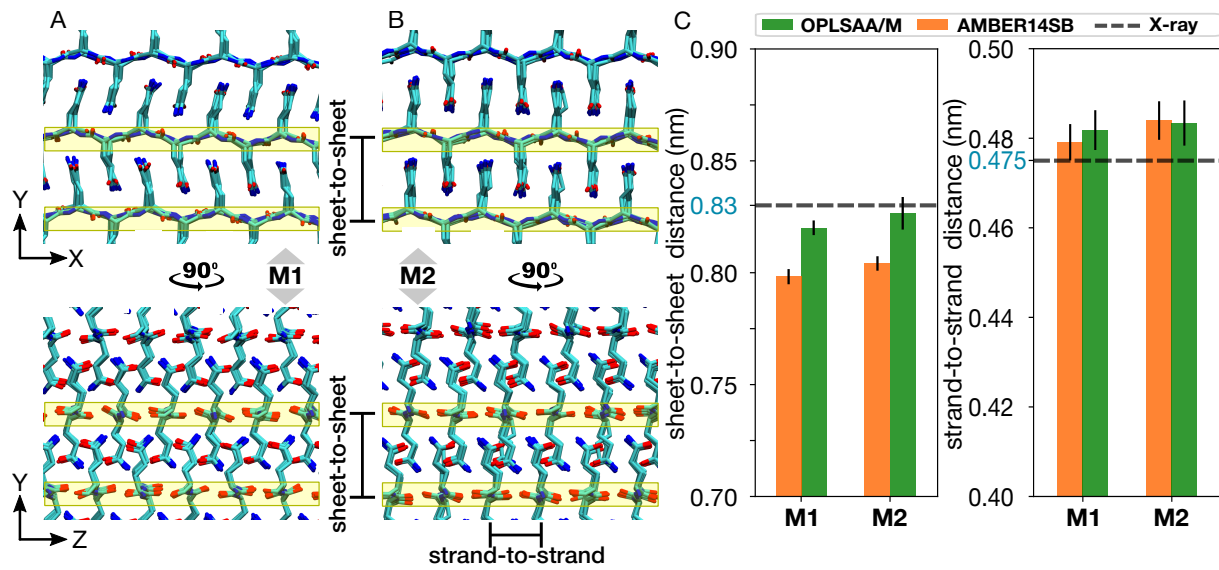


Figure S6. Illustration of the structural differences between the (A) **M1** and (B) **M2** models, visualized through views along the fibril axis z (top panels) and along the β -strand direction x (bottom panels). The yellow boxes highlight backbone atoms. (C) Sheet-to-sheet (left panel) and strand-to-strand (right panel) distances calculated for both the **M1** and **M2** models of polyQ₁₅ using the AMBER14SB (orange) and OPLSAA/M (green) force fields. The dashed black horizontal lines corresponds to the data obtained from X-ray experiments Perutz et al. (2002); Sikorski and Atkins (2005). Error bars represent standard deviation.

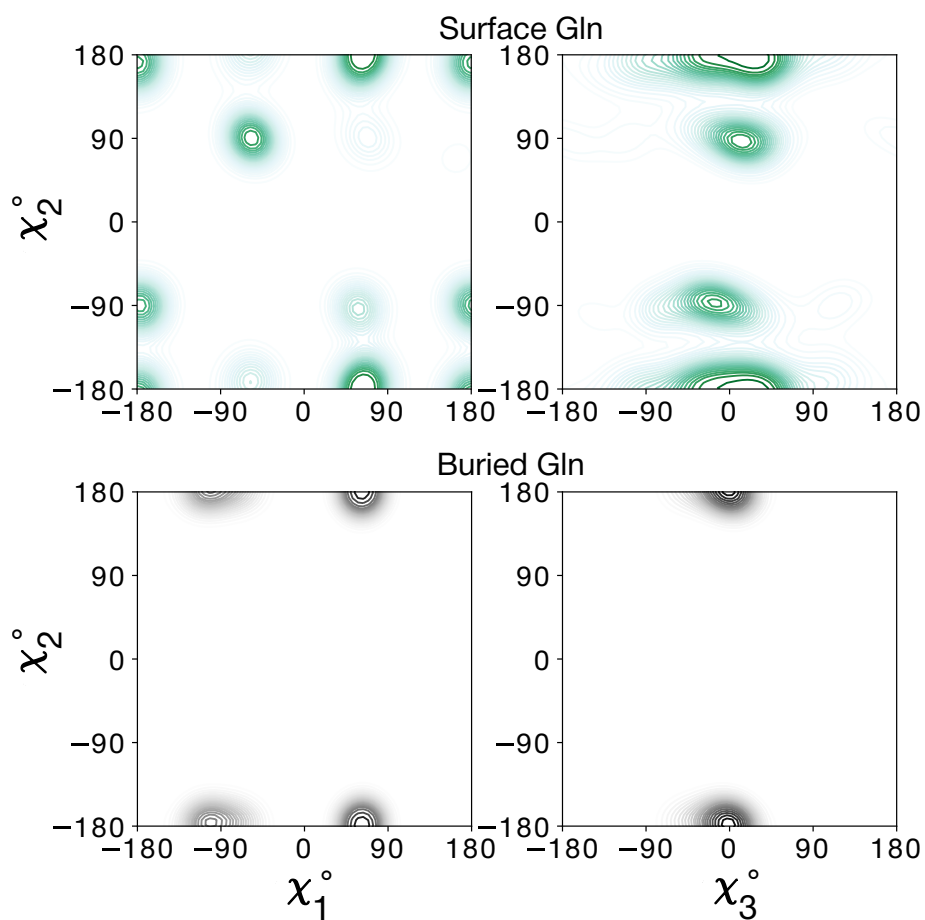


Figure S7. Distributions of the side-chain dihedral angles for Gln residues in the **M2** model of the polyQ₁₅ fibril. The water-facing side-chains (top) show more disorder than those internal to the polyQ amyloid core (bottom), but are nonetheless constrained to eight varyingly prominent specific rotamer states.

Supplementary Material

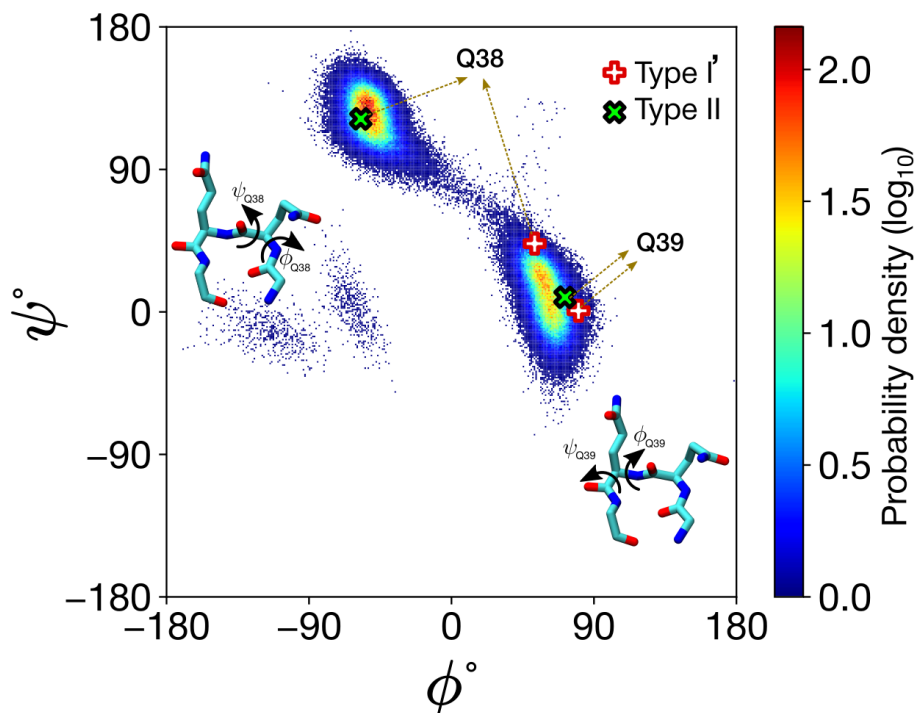


Figure S8. Ramachandran plot for the β -turn residues (Q38 and Q39) in the Q44-HttEx1 polyQ amyloid core. The distribution was obtained over the last 1 μ s of the 5- μ s MD simulation. The β -turn was initially prepared as a type I' conformer (whose canonical dihedrals are indicated with the red/white plus-signs); however, during the simulation it transitioned to the type II (black/green crosses).

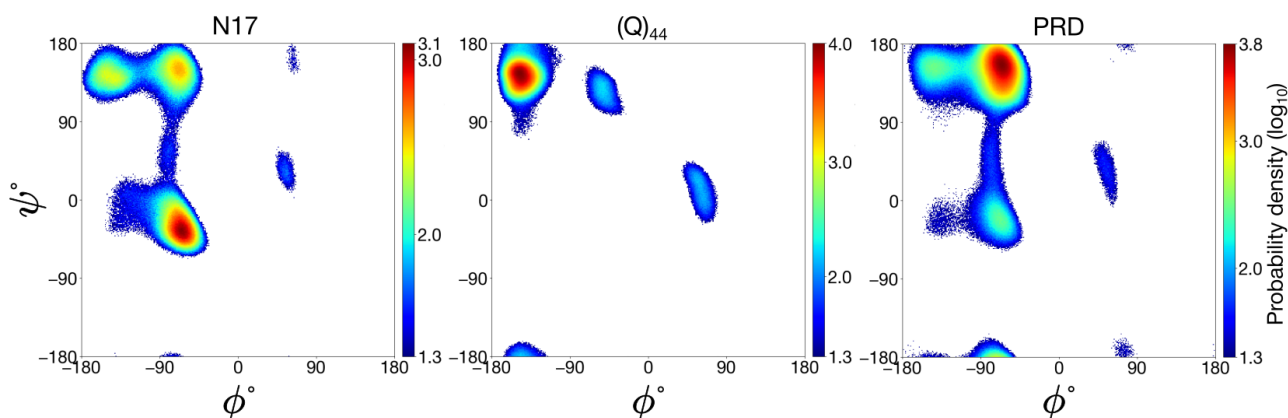


Figure S9. Ramachandran plots of the N17, Q₄₄, and PRD domains elucidate the characteristic secondary structures present in these three disparate domains of the HttEx1 protein: α -helical, β -sheet, and PPII-helical, respectively. The distributions were obtained over the last 1 μ s of the 5- μ s MD simulation. For clarity, \log_{10} (probability density) values below 1.3, characterized primarily by noise, have been excluded.

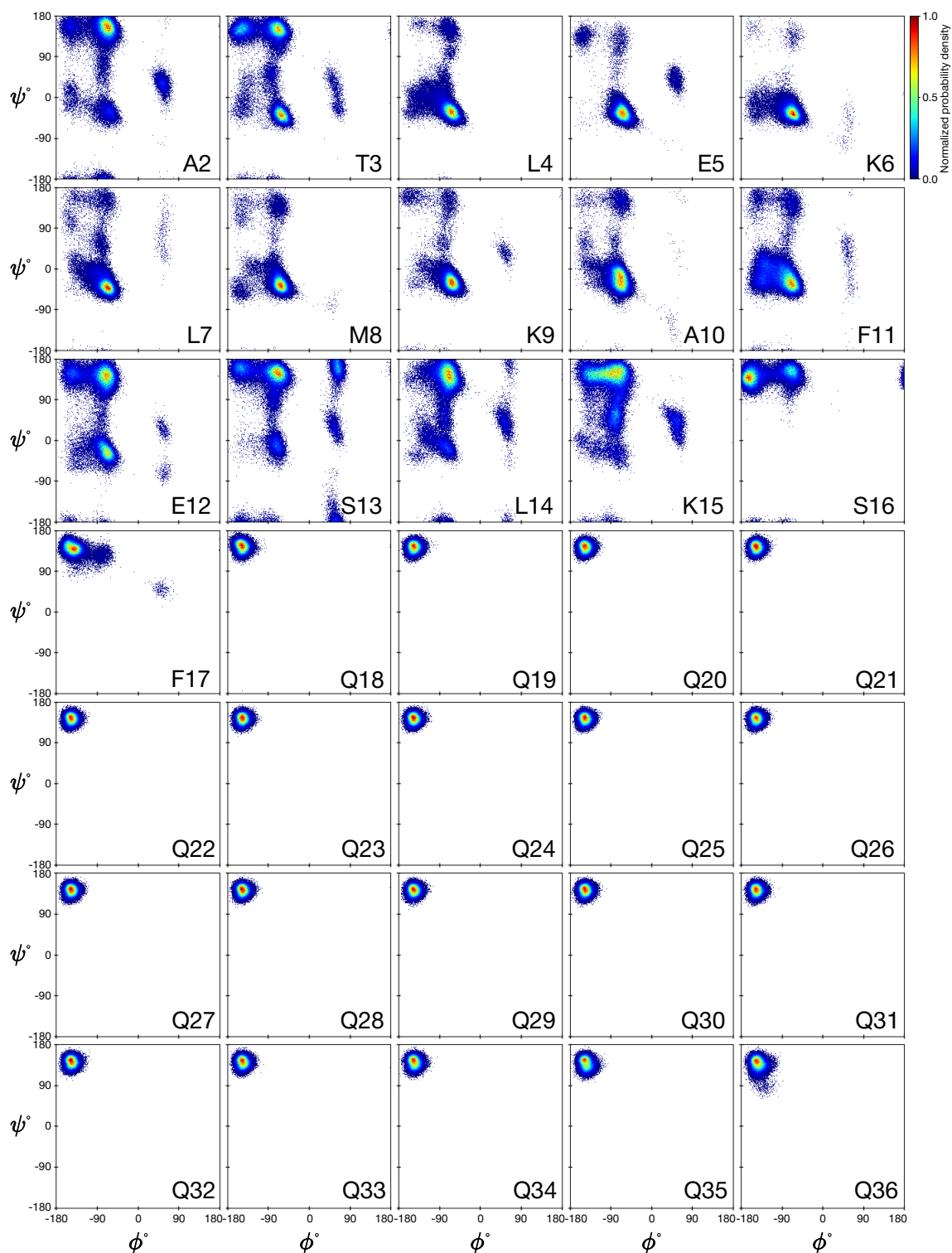


Figure S10. Ramachandran plots for residues 2–36 in the Q44-HttEx1 fibril. The plots showcase the conformational space explored by each individual residue, revealing variations in backbone torsion angles across the protein structure. The colorbar is calibrated such that a normalized value of 1 corresponds to the bin with the highest frequency among all protein residues. Each bin spans one degree in both ϕ and ψ . The distributions were obtained over the last 1 μ s of the 5- μ s MD simulation.

Supplementary Material

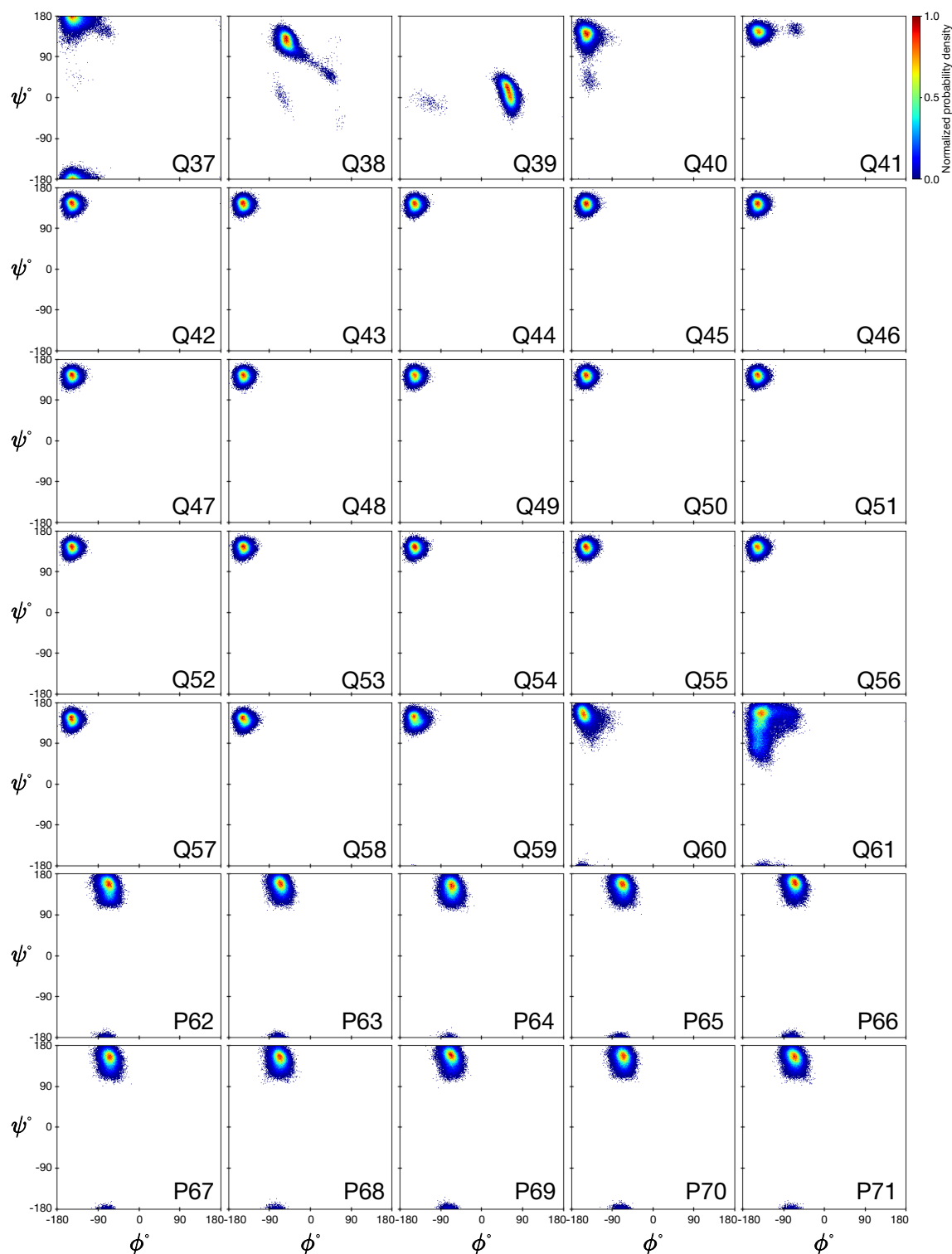


Figure S11. Ramachandran plots for residues 37–71 in the Q44-HttEx1 fibril. The plots showcase the conformational space explored by each individual residue, revealing variations in backbone torsion angles across the protein structure. The colorbar is calibrated such that a normalized value of 1 corresponds to the bin with the highest frequency among all protein residues. Each bin spans one degree in both ϕ and ψ . The distributions were obtained over the last 1 μ s of the 5- μ s MD simulation.

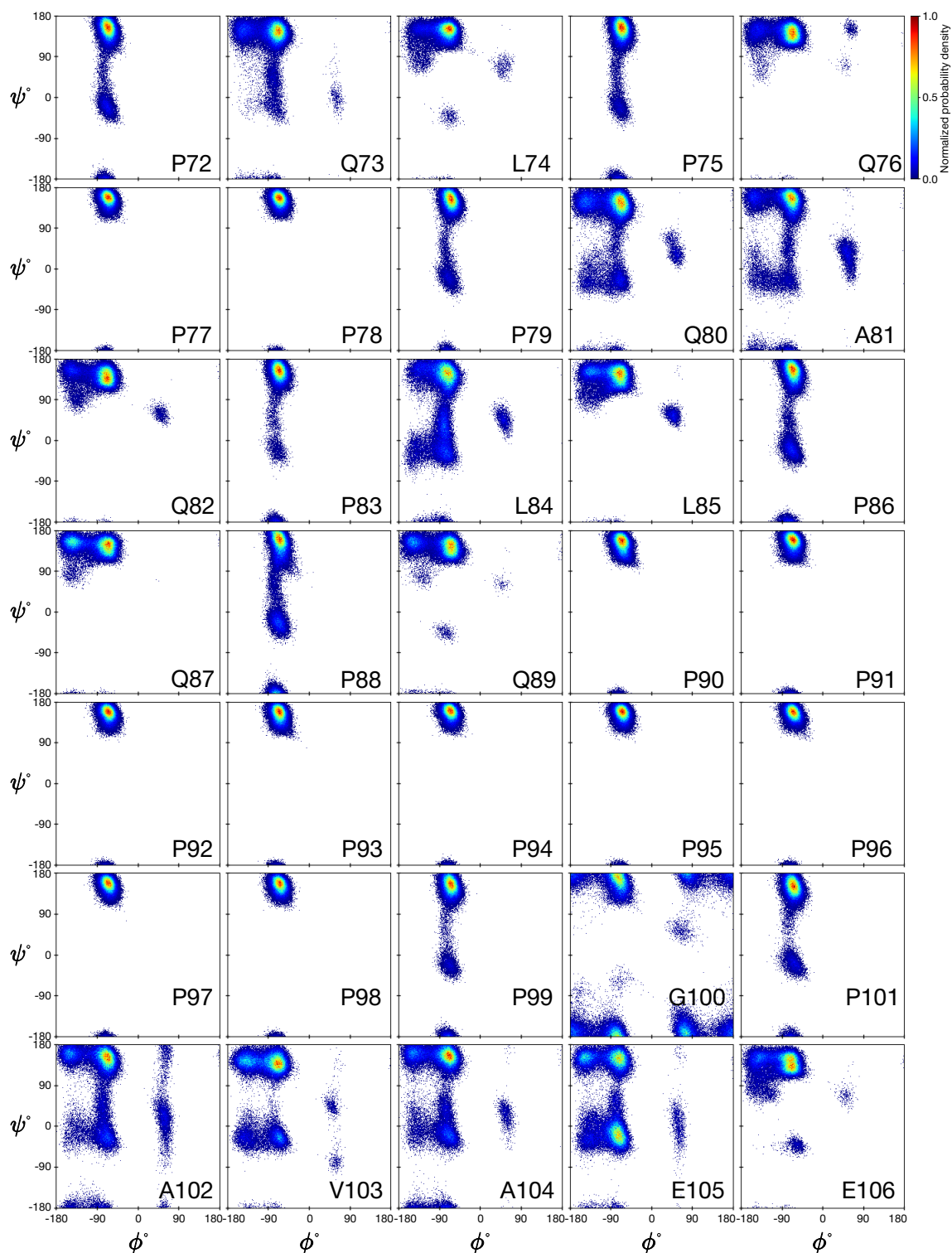


Figure S12. Ramachandran plots for residues 72–106 in the Q44-HttEx1 fibril. The plots showcase the conformational space explored by each individual residue, revealing variations in backbone torsion angles across the protein structure. The colorbar is calibrated such that a normalized value of 1 corresponds to the bin with the highest frequency among all protein residues. Each bin spans one degree in both ϕ and ψ . The distributions were obtained over the last 1 μ s of the 5- μ s MD simulation.

Supplementary Material

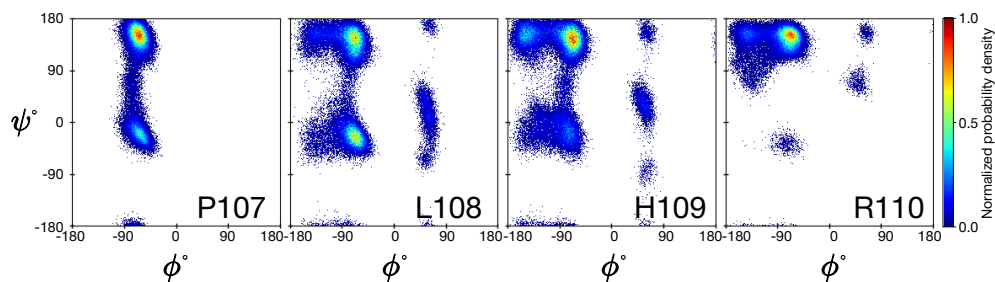


Figure S13. Ramachandran plots for residues 107–110 in the Q44-HttEx1 fibril. The plots showcase the conformational space explored by each individual residue, revealing variations in backbone torsion angles across the protein structure. The colorbar is calibrated such that a normalized value of 1 corresponds to the bin with the highest frequency among all protein residues. Each bin spans one degree in both ϕ and ψ . The distributions were obtained over the last 1 μ s of the 5- μ s MD simulation.

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