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Gallagher-Jones, Marcus Glynn, Calina Boyer, David R et al.

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# Sub-ångstrom cryo-EM structure of a prion protofibril reveals a polar clasp

Marcus Gallagher-Jones<sup>#1</sup>, Calina Glynn<sup>#1</sup>, David R. Boyer<sup>2</sup>, Michael W. Martynowycz<sup>3</sup>, Evelyn Hernandez<sup>1</sup>, Jennifer Miao<sup>1</sup>, Chih-Te Zee<sup>1</sup>, Irina V. Novikova<sup>5</sup>, Lukasz Goldschmidt<sup>2</sup>, Heather T. McFarlane<sup>2</sup>, Gustavo F. Helguera<sup>4</sup>, James E. Evans<sup>5</sup>, Michael R. Sawaya<sup>2</sup>, Duilio Cascio<sup>2</sup>, David Eisenberg<sup>2</sup>, Tamir Gonen<sup>3</sup>, and Jose A. Rodriguez<sup>1,\*</sup>

<sup>1</sup>Department of Chemistry and Biochemistry, UCLA-DOE Institute for Genomics and Proteomics, University of California Los Angeles, Los Angeles, CA 90095, USA.

<sup>2</sup>Department of Biological Chemistry and Department of Chemistry and Biochemistry, University of California Los Angeles, Howard Hughes Medical Institute, UCLA-DOE Institute for Genomics and Proteomics, Los Angeles, CA 90095, USA.

<sup>3</sup>Janelia Research Campus, Howard Hughes Medical Institute, 19700 Helix Drive, Ashburn, VA 20147, USA.

<sup>4</sup>Laboratory of Pharmaceutical Biotechnology, Institute of Biology and Experimental Medicine, Buenos Aires, AR.

<sup>5</sup>Environmental Molecular Sciences Laboratory, Pacific Northwest National Laboratory, Richland, WA 99354.

#### **Abstract**

The atomic structure of the infectious, protease-resistant,  $\beta$ -sheet-rich and fibrillar mammalian prion remains unknown. Through the cryo-EM method, MicroED, we reveal the sub-1Å resolution structure of a protofibril formed by a wild-type segment from the  $\beta 2$ - $\alpha 2$  loop of the bank vole prion protein. The structure of this protofibril reveals a stabilizing network of hydrogen bonds that link polar zippers within a sheet, producing motifs we name 'polar clasps'.

## **Summary sentence:**

Ultrahigh-resolution cryo-EM structure reveals a prion protofibril stabilized by a dense three-dimensional network of hydrogen bonds.

<sup>#</sup> These authors contributed equally to this work.

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<sup>\*</sup>Correspondence to Jose Rodriguez, Department of Chemistry and Biochemistry, UCLA-DOE Institute for Genomics and Proteomics, University of California Los Angeles, Los Angeles, CA 90095, USA, jrodriguez@mbi.ucla.edu. Jose A. Rodriguez; jrodriguez@mbi.ucla.edu.

Author contributions: J.A.R. directed the work. J.A.R., J.M., E.H., M.W.M. and C.G. grew, evaluated and optimized crystals. J.A.R., D.R.B., C.G., M.R.S., H.T.M., M.G.J., C.Z., I.V.N, J.E.E. and D.C. collected data. J.A.R., C.G., J.M., M.G.J., M.R.S., D.C., M.W.M., G.H., E.H., L.G., D.S.E., and T.G. analyzed the data. C.G., M.G.J. and J.A.R. wrote the article, with input from all authors.

Competing interests: The authors declare no competing interests.

## **Keywords**

cryo-EM; electron diffraction; prion; amyloid; protofibril

## Main text

Micro electron diffraction (MicroED) is a cryo-electron microscopy (cryo-EM) method that facilitates the determination of atomic structures from sub-micrometer thin protein nanocrystals and fragmented crystallites  $^{1-3}$ . Several high-resolution amyloid structures have been determined by MicroED. These include: two structures from the toxic core of the Parkinson's associated protein,  $\alpha$ -synuclein<sup>4</sup>, two from the type-2 diabetes associated protein, IAPP<sup>5</sup>, and five *ab initio* structures  $^{2,6}$ . These add to structures of the fibril cores of  $\alpha$ -synuclein<sup>7</sup>, amyloid-beta<sup>8</sup>, tau<sup>9</sup>, and a fungal prion  $^{10}$  recently determined by complementary methods. However, an atomic structure of the infectious, scrapie form of the mammalian prion protein (PrPSc) remains unknown  $^{11}$ . PrPSc shares some structural hallmarks of amyloid – appearing as ropelike filaments or rods made of tightly mating beta sheets  $^{12,13}$ . However, PrPSc also differs from other amyloids: it resists proteolysis and denaturation, is infectious, and can spread within and between species to cause disease  $^{11,14,15}$ .

To evaluate the source of  $PrP^{Sc}$  stability, we investigated segments in mammalian prion proteins (PrP) that might form the core of  $PrP^{Sc}$  fibrils. Informed by structure-based prediction of amyloid-prone sequences  $^{16}$  we identified  $^{168}QYNNQNNFV^{176}$ , a segment of the  $\beta 2$ - $\alpha 2$  loop of bank vole ( $Myodes\ glareolus$ ) PrP (Figure 1), a universal prion acceptor  $^{17,18}$ . This segment lies within the predicted cross-beta core of PrP fibrils (Supplementary Fig. 1), shows high conservation in rodents and other mammals (Supplementary Fig. 1), and is rich in asparagines  $^{19,20}$ , which may stabilize prion fibrils  $^{21}$ . At sub-millimolar concentrations this segment produces highly ordered aggregates (Supplementary Fig. 2) that, when aligned and illuminated by x-rays, produce cross- $\beta$  diffraction (Supplementary Fig. 2). Like  $PrP^{Sc}$ , aggregates formed by this segment are resistant to high concentrations of urea, guanidine, and a range of pH, but are sensitive to sodium hydroxide (Supplementary Fig. 3). Given its shared biophysical properties with  $PrP^{Sc}$ , we labeled our segment proto- $PrP^{Sc}$  and set out to uncover the structural basis for its stability.

From optimized microcrystals (Supplementary Fig. 4), we determined a micro-focus x-ray diffraction structure of proto-PrPSc to 1.1Å resolution by molecular replacement (Supplementary Fig. 4 and Supplementary Table 1). Conditions in which we observed microcrystals of proto-PrPSc also produced showers of nanocrystals, evident in electron micrographs (Figure 1). These nanocrystals diffracted to 0.72Å by MicroED (Figure 1). Merging diffraction from multiple crystals we achieved a high completeness, 0.75Å resolution dataset (Supplementary Table 1). From these data, we obtained an *ab initio* solution that was overall similar to our microfocal x-ray diffraction structure of proto-PrPSc (Supplementary Fig. 4) and suitable for atomic refinement (Supplementary Fig. 5). This

ultrahigh resolution MicroED structure of proto-PrPSc shows features invisible in the x-ray structure and critical to our understanding of its stability.

The structure of proto-PrPSc reveals a prion protofibril with amyloid like features: beta strands arranged parallel and in register as a class two steric zipper 12 in which sheets pair front to back (Figure 2). Two tightly mating curved sheets make up the proto-PrPSc fibril (Figure 2), although side chains in these sheets interdigitate less in proto-PrPSc than is observed in conventional amyloid structures 12. Although sheets stack in a parallel, face to back configuration, the convex face of one sheet nestles against the concave face of its neighbor, approximately 10.3Å away, with a high degree of surface complementarity (Sc, 0.807) (Figure 2). The interface between these sheets is large, concealing 204.5Å2 per strand, and is entirely devoid of waters at its core (Figure 1). Atoms in the MicroED structure of proto-PrPSc are extremely well ordered, with an average B-factor of 6.0Å2 overall and 2.8Å2 within its core (Supplementary Fig. 6 and Supplementary Table 1). These values are less than half the overall B-factor in our x-ray diffraction structure of proto-PrPSc (Supplementary Table 1), confirming a greater degree of order in our nanocrystallites compared to larger microcrystals of the same segment.

A three-dimensional network of hydrogen bonds stabilizes proto-PrPSc (Supplementary Table 2): hydrogens at its core participate in intra-residue C5 bonds<sup>22</sup> (Figure 3), and asparagine and glutamine residues stack along its fibril axis (Figure 2). Glutamines and asparagines in proto-PrPSc form networks of hydrogen bonds reminiscent of proton wires<sup>23</sup>, polar ladders<sup>24</sup>, and the polar zippers first proposed by Max Perutz<sup>25,26</sup> (Supplementary Fig. 7). Neighboring polar ladders in proto-PrPSc are additionally linked by hydrogen bonds within a strand (Figure 3), a motif we refer to as a 'polar clasp' (Supplementary Fig. 7). Stacks of phenylalanine and tyrosine residues shield clasps at the core of proto-PrPSc (Figures 1 and 2). The importance of this aromatic embrace is underscored by a lack of clasps in structures of shorter segments from this region of PrP that lack tyrosine 169<sup>12,27</sup>. On this evidence, we hypothesize that polar clasps and stacked aromatic residues act in concert to stabilize proto-PrPSc, as they might for PrPSc.

Our hypothesis of proto-PrP<sup>Sc</sup> stability relies on the locations of hydrogen atoms throughout the structure. Hydrogens in the MicroED structure of proto-PrP<sup>Sc</sup> are unambiguously assigned, informed by pronounced difference density in ultrahigh resolution maps (Supplementary Figs. 5, 8, and 9 and Supplementary Table 4). Hydrogens at the core of proto-PrP<sup>Sc</sup> are as evident as those seen in structures of small organic compounds determined by electron diffraction<sup>28</sup>, including our own structure of carbamazepine (Supplementary Fig. 10 and Supplementary Table 3). Density in ultrahigh resolution maps of proto-PrP<sup>Sc</sup> suggests hydrogen may occupy positions that deviate from idealized geometry (Supplementary Fig. 5). Improved hydrogen positions indicated by ultrahigh resolution maps in MicroED could bolster the accuracy of calculations based on observed hydrogen bond networks.

Ultrahigh resolution maps of proto-PrPSc mirror features that appear in electron density maps of the highest resolution x-ray structures and show similar residual density, which in the latter is assigned to valence electrons<sup>29</sup> (Supplementary Fig. 9). In addition, in polar

clasps at the core of proto-PrP<sup>Sc</sup>, density is seen surrounding donor hydrogens and oxygen acceptors (Figure 3 and Supplementary Figs. 5 and 9). This density is not explicitly accounted for by our atomic model during refinement, and is not evident in maps of other small molecules determined by MicroED or in lower resolution maps of proto-PrP<sup>Sc</sup> (Supplementary Figs. 8 and 10, and Supplementary Table 4). However, this density is preserved across maps produced by refinement programs with different scattering factor libraries (Supplementary Fig. 11). A full account of the complex features present in ultrahigh resolution cryo-EM maps may require advances in refinement that incorporate molecular vibrations, crystal bending, non-kinematic scattering and chemical bonding<sup>29–33</sup>.

Through MicroED, cryo-EM now surpasses the one ångstrom barrier and yields the structure of proto-PrPSc, a denaturant-resistant prion protofibril. Cryo-EM maps of proto-PrPSc at 0.75Å resolution reveal a three-dimensional network of stabilizing hydrogen bonds that link residues between and within its beta strands through polar clasps.

## **Materials and Methods**

## Phylogenetic and Sequence Analysis of bank vole PrP 27-30

The top 130 most similar wild type sequences to bank vole PrP 90–231 were generated using NCBI BLAST. Of these 130 sequences, all sequences containing the proto-PrPsc peptide were selected and aligned.

## Characterization of proto-PrPSc peptide

The synthetic peptide representing sequence <sup>168</sup>QYNNQNNFV<sup>176</sup> of the bank vole prion protein was purchased from GenScript (Piscataway, NJ). The peptide used for our experiments was received at greater than 98% purity achieved by reverse phase HPLC. The peptide was qualified on a Bruker (ultrafleXtreme) MALDI-TOF/TOF mass spectrometer, reported as m/z (intensity, arbitrary units). The spectrum has a mass list that includes a [M+H]+ peak at 1141.3 Da (expected 1140.5 Da), a [M+Na]+ peak at 1163.4 Da (expected 1162.49 Da), and a [M+K]+ peak at 1179.3 Da (1178.46 Da).

## Characterization of carbamazepine

Lyophilized powder of carbamazepine (5H-dibenzo[b,f]azepine-5-carboxamide,  $C_{15}H_{12}N_2O$ ) with purity > 99% was purchased from Sigma-Aldrich (St Louis, MO) and crystallized without further purification.

## Aggregation of proto-PrPSc

Proto-PrP<sup>Sc</sup> peptide was solubilized in ultrapure water at 0.22–1.75 mM. 50 µL of each sample was added to a 96-well clear flat bottom plate in triplicate and evaluated for aggregate formation by reading absorbance at 350 nm on an Infinite M1000 Pro plate reader (Tecan). Readings were measured immediately after solubilization, after three hours, and after six hours of shaking at 900 rpm at 37° C. Wells were imaged after six hours using a Leica M205 C light microscope (Leica Microsystems) and after three and six hours by electron microscopy as described below. Electron microscope images are representative of more than five images taken at each concentration and time point.

## Fibril diffraction from proto-PrPSc aggregates

Solutions containing aggregates of *proto-PrPSc* were clarified by centrifugation. Pelleted aggregates were resuspended in a concentrated volume in water, applied between two pulled capillaries and left to dry overnight. Oriented aggregates formed between the capillary ends, were resupplied with additional solution containing aggregates, and left to dry again. This was repeated several times to grow the bulk of aligned aggregates on a capillary. Aligned aggregates were then diffracted using 5 minute exposures to a FRE+ rotating anode generator with VARIMAX HR confocal optics producing Cu K- $\alpha$  radiation (Rigaku, Tokyo, Japan) and detected using a RIGAKU R-AXIS HTC imaging plate detector at a distance of 156 mm from the source.

## Chemical denaturation of proto-PrPSc aggregates

Proto-PrP<sup>Sc</sup> was solubilized at approximately 3.5 mM and allowed to form aggregates. This solution was diluted to approximately 1:4 before treatment with either 0.5 – 6.0 M urea, 0.5–4.5 M guanidinium-HCl, 0.75 M HCl, 0.1 M citrate pH 2 or 4, 0.1M Sodium/potassium phosphate pH 6 or 8, 0.1 M CAPS pH 10, or 0.75 M NaOH. Aggregate content was measured by absorbance compared to a control solution consisting of 0.01% (w/v) 1um latex spheres in water. Spectra across the visible range (250–700 nm) were collected using a Nanodrop One (Thermo). A single point of reference, OD at 600 nm, was used for comparison across sampled conditions.

## Growth of proto-PrPSc crystals

Peptide was weighed out as powder and dissolved in ultrapure water at near maximum solubility, 3.5 mM. Crystals were grown at room temperature by the hanging-drop method in a 96-well Wizard screen. Crystals appeared in various conditions, and were further optimized in 24-well hanging-drop vapor diffusion experiments. The best crystals of proto-PrPSc grew within 24 hours at a peptide concentration of 1.75 mM in the presence of 0.1 M 2-(*N*-morpholino) ethanesulfonic acid (MES), pH 6.0 and either 10% ethanol or 10% 2-Methyl-2,4-pentanediol (MPD).

#### Transmission electron microscopy

Approximately two microliters of aggregated proto-PrPSc were applied to 300-mesh formvar-carbon coated grids (Ted Pella Inc., Redding, Ca.) for two minutes before excess liquid was removed and grids left to dry. Grids were imaged either on a Tecnai T12 or F20 electron microscope (Thermo-Fisher; formerly FEI). Samples were imaged at a magnification of  $2100\times$  with a dose rate of less than  $30~\text{e}^-/\text{Å}^2$ .

#### Microfocus x-ray data collection

Crystals grown in 0.1M MES pH 6.0 and 10% ethanol and mixed with 100% glycerol as cryoprotectant were harvested from 24-well hanging drops using MiTeGen loops and flash frozen in liquid nitrogen. Seventy-three diffraction images were collected, each spanning an  $3^{\rm o}$  wedge, from a single crystal at a temperature of 100K at the advanced photon source (APS) beamline 24-ID-E, equipped with an ADSC Q315 CCD detector, using a 10  $\mu m$  beam with a 0.98 Å wavelength.

## Microfocus x-ray data processing and structure determination

Diffraction images collected from a single crystal of proto-PrPSc were indexed and integrated in Denzo, yielding a dataset with 80.75% overall completeness at 1.1 Å resolution in space group P1. A suitable molecular replacement solution was obtained from this data using the PHASER program and an idealized beta strand nonapeptide alanine model as a probe. The model was refined using REFMAC, against the measured data to a final  $R_{\rm work}/R_{\rm free}$  of 0.14/0.16.

## MicroED sample preparation

Nano-scale needle crystals of proto-PrPSc were grown in batch in 0.1 M MES, pH 6.0 and 10% ethanol. Crystals were diluted in mother liquor and fragmented by force of pipetting to create an approximately monodisperse solution of crystals. Carbamazepine was crystallized in batch by dilution into neat isopropanol at 100  $\mu$ g/mL. Two microliters of either solution were placed on a holey carbon grid (1/4, 2/2, 2/4, #300 copper; Ted Pella Inc.) prior to plunge freezing into liquid ethane and transferring into liquid nitrogen for storage. Grids were held by a liquid nitrogen cooled Gatan 626 cryo-holder for transfer into and manipulation within the electron microscope.

#### MicroED data collection

MicroED data collection from nine sub-micron thick needle crystals and a single sub-micron carbamazepine crystal was performed as previously described  $^{34}$ . Briefly, crystals of either proto-PrPSc or carbamazepine lying in a frozen-hydrated state on holey carbon grids were inspected visually in overfocused diffraction mode on either a cryo-cooled FEI Tecnai F20 microscope operated at 200 kV (Janelia Research Campus), or a Titan environmental TEM operated at 300 kV (Environmental Molecular Sciences Lab, PNNL). Diffraction patterns used for structure determination were collected on a TVIPS TemCam-F416 CMOS detector in rolling-shutter mode with three second exposures while proto-PrPSc crystals were unidirectionally rotated at a constant rate of  $0.27^{\rm o}$  s $^{-1}$  over angular wedges ranging from  $-55^{\rm o}$  to  $+72^{\rm o}$ . A single carbamazepine crystal was rotated at a speed of  $0.2^{\rm o}$  s $^{-1}$  over an angular wedge ranging between  $-45^{\rm o}$  to  $+45^{\rm o}$  with 5 second exposures. Beam intensity was held constant, with an average dose rate of 0.003-0.005 e $^{-}$ Å $^{-1}$  sec $^{-1}$ , corresponding to a total dose of  $\sim 1-3$  e $^{-}$ Å $^{-2}$  per data set. We used a camera length of 520 mm, the equivalent of a sample to detector distance of 950 mm in a corresponding lensless system. All diffraction was performed using a circular selected area aperture of  $\sim 1 \, \mu m^2$  in projection.

#### MicroED data processing

Diffraction movies were converted to the SMV file format using TVIPS tools as previously described<sup>35</sup>. Indexing and integration were performed in XDS<sup>36</sup>. Integrated diffraction intensities from partial data sets of nine different proto-PrPSc crystals were sorted and merged in XSCALE<sup>36</sup>. Merged intensities were converted to amplitudes at various resolution cutoffs to produce separate 1.1Å, 1.0Å, 0.9Å, 0.8Å, and 0.75Å data sets. *Ab initio* structure determination was performed on each of these data sets using SHELXD<sup>37</sup>. Phases obtained from the atomic assembly generated by direct methods were used to produce maps of sufficient quality for subsequent model building in Coot<sup>38</sup> and refinement in Phenix<sup>39</sup>

using electron scattering form factors to produce a final structure in space group P 1 with a final  $R_{work}/R_{free}$  of 24/25. Refinement in REFMAC was carried out in parallel to a final  $R_{work}/R_{free}$  of 23/25. The structure refined in Phenix (PDB ID 6AXZ) was used in all subsequent analysis and is shown in figures. *Ab initio* structure determination for carbamazepine was performed in SHELXT<sup>37</sup> in which a solution was found in space group P 21<sub>m</sub> with no errors in chemical assignment or atom positions for all carbon, nitrogen, and oxygen atoms. This solution was refined in SHELXL<sup>37</sup> using electron form factors to an R value of 21.8%. A structure with hydrogen positions refined to best match difference density in the map lowered the R value to 19.8%.

## Analysis of buried surface area (Sa) and surface complementarity (Sc) for proto-PrPSc

The structure of proto-PrPSc was used to calculate both  $S_a$  and  $S_c$  from an assembly consisting of two sheets generated by translational symmetry, each consisting of ten stacked beta strands.  $S_a$  was computed as an average of the buried surface area per chain in our assembly, calculated as the difference between the sum of the solvent accessible surface area of the two sheets and the solvent accessible surface area of the entire complex, divided by the total number of strands in both sheets.  $S_c$  was calculated using the CCP4 suite for all points at the interface between the two aforementioned sheets.

## Comparison of x-ray and MicroED structures

Sequence alignment and structural superposition of both an individual chain and an assembly of two sheets from x-ray and MicroED structures of proto-PrPSc were performed in PyMOL $^{40}$ . This alignment produced an all atom R.M.S.D. of 0.162Å on which structural similarity was assessed.

## Analysis of aromatic residues in proto-PrPSc

Distances were measured between stacked aromatic residues that flank the core of proto- $PrP^{Sc}$ . Aromatics stacked between strands along planes separated by 3.7 or 3.4Å for  $Y^{169}$  or  $F^{175}$ , respectively, in a parallel-displaced configuration  $^{41}$ . We measured the inter-sheet hydrogen bond created by  $Y^{169}$  to the backbone carbonyl of  $N^{171}$  on an opposing strand. Both  $Y^{169}$  and  $F^{175}$  formed aromatic ladders that channel polar residues into a 12.8Å long region at the core.

#### Analysis of hydrogen bond networks

Informed by the locations of hydrogens, visible in our maps at a  $0.7-2.5~\sigma$  range, we evaluated hydrogen bond networks determined by the program HBplus  $(v.3.06)^{42}$  and by manual inspection using PyMOL. We measured asparagine ladders between residues  $N^{170}$ ,  $N^{171}$ ,  $N^{173}$ , and  $N^{174}$  and the corresponding asparagine residues on the strands above and below. We also measured glutamine ladders between residues  $Q^{168}$  and  $Q^{172}$  and their corresponding residues on strands above and below. We analyzed distances associated with intra-strand hydrogen bonds formed by three pairs of residues:  $Q^{168}$ - $N^{170}$ ,  $N^{171}$ - $N^{173}$  and  $Q^{172}$ - $N^{174}$ . Polar contacts were also measured between  $Q^{168}$ - $N^{170}$ , with a 3.1 Å donor-acceptor (D-A) distance and 2.3Å H-O distance,  $Q^{172}$ - $N^{174}$ , with a 3.0 Å D-A distance and 2.1Å H-O distance and  $Q^{171}$ - $Q^$ 

each case, the linked residues faced the same side of the beta strand and bridged residues at positions  $[N/Q]^i$  and  $[N]^{i+2}$  within the strand. While the HBplus program does not identify C5 hydrogen bonds in our structure, we measured these bonds in residues  $N^{171}$ ,  $Q^{172}$  and  $N^{173}$ , based on the criteria that carbonyl oxygens bond with intra-residue amide protons if their geometry permits, with H-O distances shorter than  $2.5 \text{\AA}^{22}$ .

## Amylome profiling and informatic analysis of polar clasps

A subset of the predicted amylome was analyzed <sup>16</sup>, consisting of six residue segments found to score favorably when threaded onto a template based on the structure of the yeast prion NNQQNY. We chose all segments that scored two standard deviations better than the mean score for all peptides (Z-score >2). This subset of six residue segments represents 95,381 out of 7,900,599 total segments, or 1.2% of all possible segments of this size in the human proteome. For each segment, we searched for a [Y/F] X [N/Q] X [N/Q] X [Y/F] motif across the region of the protein to which the segment belonged; a 10-residue window including two residues upstream and downstream of a profiled segment. Protein identifiers were obtained from the RefSeq database for each of the 35 catalogued segments. Where a segment appeared in more than one protein, each was counted independently.

## Calculation of contour maps

 $2F_o$ - $F_c$  and  $F_o$ - $F_c$  density maps were calculated from the final refined MTZ file using the FFT tool in CCP4. Maps were converted to MRC format in Chimera and imported into MATLAB. Contour plots were calculated such that the number of contours spanned the minimum to the maximum values of the maps with intervals of one standard deviation  $(\sigma)$  between contour levels.

#### **Data Availability**

Atomic coordinates and structure factors for proto-PrPSc have been deposited in the Electron Microscopy and Protein Data Banks as EMD-7017, 6AXZ respectively. Source data for all figures and files is available from the authors upon reasonable request, please see author contributions for specific datasets. A life sciences reporting summary for this article is available.

## **Supplementary Material**

Refer to Web version on PubMed Central for supplementary material.

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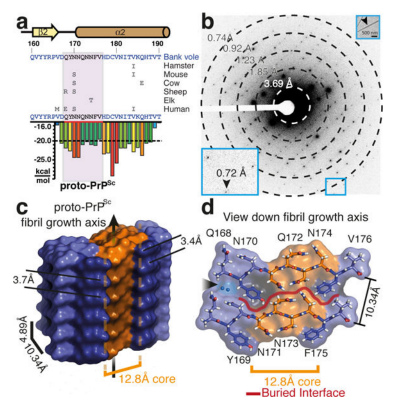


Figure 1. Structure of proto-PrPSc from sub-Å diffraction of nano-scale crystals by MicroED.  $\bf a$ , Sequence of the bank vole PrP  $\beta 2$ - $\alpha 2$  loop, highlighting local secondary structure and variation in sequence compared to hamster, mouse, cow, sheep, elk, and human. The sequence of proto-PrPSc is highlighted in a plot that indicates propensity for amyloid formation (scored in kcal/mol). The region encoding for proto-PrPSc is shown in magenta.  $\bf b$ , Single frame of MicroED data showing strong diffraction to 0.72Å (inset). A second inset shows a proto-PrPSc crystal diffracted by MicroED (arrow). Scale bar 500nm.  $\bf c$ , Side and  $\bf d$ , top view of an isosurface model of proto-PrPSc. Stacked aromatics are highlighted in dark purple, a highly constrained core region in orange, two waters are noted with a black arrow, and a red line outlines the interface between sheets in panel  $\bf d$ .

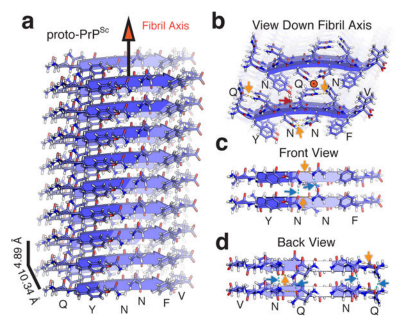


Figure 2.

Filament structure of proto-PrPSc and its stabilizing network of hydrogen bonds. a, Structure of proto-PrPSc determined by MicroED shown as an assembly of ten chains. Distances between strands and sheets, and the fibril growth axis are labeled. b, A view down the fibril axis highlighting the interface between adjacent sheets. An orange circle denotes the direction of the fibril axis. c, Front and d, back views of the fibril with two strands representing a sheet. Polar contacts are labeled as orange arrows for intra-chain and blue arrows inter-chain.

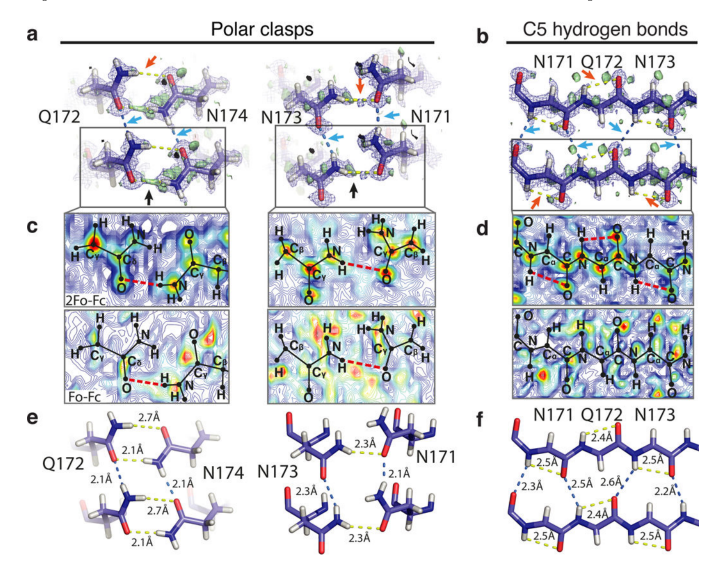


Figure 3. Sub-Å cryo-EM maps show stabilizing hydrogen bonds in proto-PrPSc. **a**, Polar clasps formed by hydrogen bonds that link Q<sup>172</sup> to N<sup>174</sup>, and N<sup>171</sup> to N<sup>173</sup>, and **b**, C5 hydrogen bonds (orange arrows) in the context of backbone hydrogen bonds at the core of proto-PrPSc. Bridging density can be seen in both the  $2F_0$ - $F_c$  and  $F_0$ - $F_c$  maps (blue mesh and green isosurface respectively) for a subset of the hydrogen bonds (black arrows). Contour maps show  $2F_0$ - $F_c$  or  $F_0$ - $F_c$  density for **c**, polar clasps or **d**, C5 bonds. Contours are averaged over a ~1.5Å thick volume with high contour values are shown in red, low values in blue, and hydrogen bonds as dashed red lines. Distances for both intra (yellow lines) and inter (blue lines) chain hydrogen bonds are highlighted in **e**, polar clasps or **f**, C5 bonds.