Full atomistic model of prion structure and conversion

Giovanni Spagnolli¹*, Marta Rigoli¹,², Simone Orioli²,³, Alejandro M. Sevillano⁴, Pietro Faccioli²,³, Holger Wille⁵, Emiliano Biasini¹*, Jesús R. Requena⁶*

1 Department of Cellular, Computational and Integrative Biology (CIBIO)—University of Trento, Povo TN, ITALY, 2 Department of Physics, Povo, Trento TN, ITALY, 3 INFN-TIFPA, Povo (Trento), ITALY, 4 Department of Pathology—University of California—San Diego, San Diego, California, United States of America, 5 Department of Biochemistry and Centre for Prions and Protein Folding Diseases—University of Alberta, Edmonton (AB), CANADA, 6 CIMUS Biomedical Research Institute & Department of Medical Sciences, University of Santiago de Compostela-IDIS, Santiago, SPAIN

* giovanni.spagnolli@unitn.it (GS); emiliano.biasini@unitn.it (EB); jesus.requena@usc.es (JRR)

Abstract

Prions are unusual protein assemblies that propagate their conformationally-encoded information in absence of nucleic acids. The first prion identified, the scrapie isoform (PrP⁰) of the cellular prion protein (PrP⁵), caused epidemic and epizootic episodes [1]. Most aggregates of other misfolding-prone proteins are amyloids, often arranged in a Parallel-In-Register-β-Sheet (PIRIBS) [2] or β-solenoid conformations [3]. Similar folding models have also been proposed for PrP⁰, although none of these have been confirmed experimentally. Recent cryo-electron microscopy (cryo-EM) and X-ray fiber-diffraction studies provided evidence that PrP⁰ is structured as a 4-rung β-solenoid (4RβS) [4, 5]. Here, we combined different experimental data and computational techniques to build the first physically-plausible, atomic resolution model of mouse PrP⁰, based on the 4RβS architecture. The stability of this new PrP⁰ model, as assessed by Molecular Dynamics (MD) simulations, was found to be comparable to that of the prion forming domain of Het-s, a naturally-occurring β-solenoid. Importantly, the 4RβS arrangement allowed the first simulation of the sequence of events underlying PrP⁰ conversion into PrP⁰. This study provides the most updated, experimentally-driven and physically-coherent model of PrP⁰, together with an unprecedented reconstruction of the mechanism underlying the self-catalytic propagation of prions.

Author summary

Prions are unusual infectious pathogens that do not contain any nucleic acid. They consist of assemblies of misfolded proteins. The scrapie isoform of the mammalian prion protein, PrP⁰, is the most notorious prion, and is responsible for deadly neurodegenerative diseases affecting humans, like Creutzfeldt-Jakob disease, and animals, such as bovine spongiform encephalopathy (“mad cow disease”) and chronic wasting disease affecting elk and deer in North America and, more recently, Europe. Understanding the structure (“shape”) of the PrP⁰ prion is critical to understand how it propagates. We have created a very detailed model of PrP⁰, which includes all its atoms, using computational
techniques. The model resembles a 4-rung cork-screw. The model is stable and agrees
with all available experimental data on PrPSc. This structure allowed us to model for the
first time the process of prion propagation at high resolution. These data inspire several
new hypotheses to elucidate prion biology and design therapeutics for prion diseases

Introduction

Prion diseases are infectious neurodegenerative disorders characterized by an invariably lethal
outcome caused by a proteinaceous infectious agent named “prion” [1]. The central event in
these pathologies is the conversion of PrPC, a GPI-anchored protein of unknown function,
into a misfolded isoform (PrPSc) which accumulates in the central nervous system of affected
individuals [6]. While PrPC structure has been widely characterized, and consists of a N-termi-
nal disordered tail and a C-terminal globular domain [7], no high-resolution information is
available for PrPSc due to technical challenges posed by its high insolubility and aggregation
propensity [8]. In order to fill this gap, different atomistic models based on low-resolution
experimental data have been proposed, including a Left-handed-β-Helix (LβH) structure span-
ing residues 89 to 170 while retaining the two C-terminal α-helices of PrPC [9], and a Parallel
In-Register Beta-Sheet (PIRIBS) architecture, characterized by intermolecular stacking of
aligned PrP monomers [10]. Recent cryo-EM data obtained using infectious, anchorless PrPSc
fibrils [4] provided strong evidence indicating that PrPSc fibrils consist of two independent
protofilaments, and of the existence of 2 nm structural units repeating along each protofila-
ment axis, suggestive of a 4-rung β-solenoid (4RβS). This would be fully compatible with the
LβH model, and much less so with the PIRIBS model, which posits that PrPSc fibrils are not
made up by two protofilaments, but rather, by a single wider filament that features two subdo-
 mains or lobes separated by a cleft (Goveman et al., 2014). Moreover, the PIRIBS model fails to
accommodate glycosylated residues in PrPSc, which would result in the introduction of exces-
sive steric clashes [11]. It should be pointed out, however, that the low resolution of available
experimental data still does not allow to definitively discard any option (a detailed comparison
of PIRIBS and solenoid-based models can be found in [12]). Of note, while consistent with the
mentioned experimental constraints, the proposed LβH model is incoherent with a recent re-
evaluation of previous FTIR data suggesting that PrPSc does not contain α-helices [8].

Results

Construction of an experimentally-driven atomistic model of infectious
mouse prion

To satisfy current experimental evidence and theoretical structural constraints, we built a new
atomistic model of mouse PrPSc based on the 4RβS conformation, and tested its stability by
means of all-atom MD simulations. The construction of the model took into account an array of
experimental data, including: (i) cryo-EM [4] and X-ray fiber-diffraction studies [5], which
showed that the fold of a mouse glycosylphosphatidylinositol (GPI)-anchorless, infectious
PrPSc is compatible with a 4RβS architecture with L- or T- shaped cross-section [4]; (ii) circu-
lar dichroism (CD) and FTIR spectroscopy, which ruled out the presence of α-helices, and sug-
gest that PrPSc contains approximately 40–50% β-sheet and 50–60% coil/turns [8]; (iii) Mass
Spectrometry (MS) analyses indicating the presence of an intact disulphide bond between resi-
dues C178 and C213 (mouse sequence) [13], as well as mapping Proteinase K (PK)-sensitive
residues, which reflect amino acids likely excluded from the resistant core of the protein [14]
and (iv) the possibility of accommodating complex glycans at positions N180 and N196 [11]. All these constraints were comprehensively included into a 2D threading scheme spanning mouse PrP (moPrP) residues 89–230 (S1 Fig), also considering the structural propensities of different residues: polyglycine tracts and prolines were positioned in loops due to their destabilizing effects on β-strands; charged sidechains were excluded from the inner core of the protein or counterbalanced by salt bridges. This scheme was then modelled onto the 3D arrangement of a naturally-occurring β-solenoid protein (Dickeya dadantii Pectate Lyase; PDB 1AIR). The resulting structure (depicted in Fig 1) features an inner core containing mainly hydrophobic or mildly polar side-chains (T94, T106, L108, V111, Y127, M128, W144, Y149, V165, Y168, I181, I183, V188, F197, T198 and T200), few polar side-chains involved in hydrogen bonding (N142-HB-Y168, H168-HB-T198, Q216-HB-T200 and Q218-HB-S221), and a salt bridge (R147-SB-D166). Conversely, the majority of the highly-polar residues (N and Q) including the glycosylation sites (N180 and N196) and charged side-chains (E, D, K and R) are exposed to the solvent. The structure also encompasses identified PK cleavage sites localized in loops/turns, or at the edge of the β-strands, and the intact disulphide bond between C178 and C213. Importantly, the final model fitted with a previously described, low-resolution cryo-EM map of infectious PrPSc (S2 Fig).

Molecular dynamics simulations of the new 4RβS model

To test the physical consistency of the 4RβS model, we challenged its stability by all-atom MD simulations in explicit solvent. First, three independent, 20 ns simulations were performed in the isothermal-isobaric ensemble (T = 300 K, P = 1 Bar) restraining hydrogen bonds distances between atoms involved in β-sheets. This process allowed relaxation of protein loops and side chains of the core (S3 Fig). Next, the imposed restraints were released, and three plain-MD trajectories of 100 ns each were simulated. The stability of the model was then assessed by both Root Mean Squared Deviation (RMSD) of atomic positions relative to the initial frame (t = 0 ns) and secondary structures content. Interestingly, we obtained values in the same range of fluctuation for our 4RβS model and the prion-forming domain of the fungal protein Het-s, a naturally occurring β-solenoid whose structure has been solved by solid-state NMR (ssNMR; PDBs 2KJ3 and 2RNM; Fig 2, and S4 Fig). Conversely, by applying an identical workflow to the previously proposed model of laterally-stacked LβH trimer of PrPSc, we observed a profound instability of the β-helical domain, which was already evident after few tens of ns of dynamics (Fig 2 and S4 and S5 Figs).

Next, we tried to adapt alternative threading schemes to the 4RβS architecture. We obtained two new schemes carrying small deviations (i.e. one or two residue shifts) from the original model but still fully coherent with the experimental constraints (S6 Fig). Every other attempt to model threading schemes with a larger deviation (>3 residue shifts) resulted in violation of experimental constraints. Interestingly, MD simulations revealed that the two alternative models are also significantly stable (S6 Fig). These data suggest that the new threading schemes may represent putative alternative arrangements of the PrP polypeptide into an RML prion fibril. This conclusion implies an intrinsic flexibility of 4RβS architecture to accommodate slightly different arrangements, which could be related to the ability of prions to occur in different conformations specifying prion strains.

To mimic the fibrillary behavior of PrPSc, we built a tetrameric 4RβS model by stacking monomers in a head-to-tail fashion. As expected, this assembly showed a comparable MD stability to the monomer (Fig 3). Moreover, the 4RβS tetramer fits with the two main signals obtained by Fourier transform single particle analysis of cryo-EM data (19.1 Å and the ~40 Å) [4], which reflect distances between the same residues on two contiguous or alternate
monomers, respectively, as well as with the observed volume of the protofilaments (Fig 3). Finally, by introducing complex sugar precursors (GlcNAc2Man3Fuc) at positions N180 and N196 of each monomer, we observed the absence of steric clashes, confirming that the 4RβS model accommodates the presence of glycans (S6 Fig). Collectively, these findings indicate that, in contrast to the LβH model, the 4RβS is a solid arrangement for PrPSc, built on the most

Fig 1. View of the 4RβS PrPSc Model. The structure of PrPSc modelled as a 4RβS (β-strands represented as red arrows) is depicted in the center of the figure. Residues are displayed in each individual rung (1–4) with different colors. PK cleavage sites identified by mass spectrometry in two different previous reports are colored in red [14] [15]. Glycosylation sites are labelled in green. Prolines are colored in purple. Cystine is indicated in cyan.

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recent experimental data, and showing a conformational stability comparable to that of a natural prion.

**All-atom reconstruction of prion conversion**

The 4RβS model allowed us to develop an original scheme to perform for the first time a simulation of the conformational transition from PrP\(^{C}\) to PrP\(^{Sc}\). In order to bridge the gap between the computationally-accessible and the biologically-relevant time scales, we employed a
specific kind of biased dynamics called ratchet-and-pawl MD (rMD) [16] in the framework of an all-atom realistic force field [17]. rMD-based methods have been successfully applied to simulate protein folding and other conformational transitions [18, 19]. However, this scheme provides a sampling of the transition path ensemble only if the biasing force is applied along a reliable reaction coordinate [20]. Therefore, the first step towards developing our rMD simulation was to build a statistical model to identify the reaction coordinate of the process. Using the Markov State Mode formalism, we demonstrate that among all the possible misfolding...
patterns from PrP\textsuperscript{C} to PrP\textsuperscript{Sc} the prominent reaction mechanism is the sequential formation of rungs. A biasing coordinate was then built by coupling this dynamical information with the all atom 4R\betaS structure. The associated rMD simulations yielded a transition pathway with full atomistic resolution in which the C-terminal rung of the solenoid acts as a primary conversion surface for PrP\textsuperscript{C} unstructured N-terminus (residues 89–124). This first event initiates a cascade of conformational transitions in which each newly formed rung acts as a template for the formation of the following one, ultimately leading to the complete conversion of PrP\textsuperscript{C} into PrP\textsuperscript{Sc} (Fig 4 and S1 Movie). This analysis provides a rigorous description of the active role of protofilament ends in the templated propagation of prions, and it is compatible with previous observations suggesting the presence of a structured intermediate conformer of PrP\textsuperscript{C} in its transition to PrP\textsuperscript{Sc} [21].
Discussion

The elucidation of the structure of PrP\textsuperscript{Sc} at atomic resolution has proven to be a phenomenal experimental challenge, mainly due to its high insolubility and aggregation propensity. Previously generated computational models of PrP\textsuperscript{Sc} have the virtue of providing a plausible 3D structure, but fail to comprehensively accommodate most recent experimental data [8]. Here, we filled this gap by exploiting the information arising from cryo-EM and X-ray fiber-diffraction studies [4, 5], which clearly defined the general architecture of PrP\textsuperscript{Sc} as a 4R\boldsymbol{\beta}\boldsymbol{S} and refined the structure by including experimental constraints obtained by mass spectrometry. The model we have created allowed us to perform the first reconstruction of how the information encoded into the conformation of a protein could be propagated in a directional fashion, a concept directly underlying the infectious nature of prions.

Virtues and limits of the 4R\boldsymbol{\beta}\boldsymbol{S} model

The search for effective therapies against prion diseases has so far been mostly unsuccessful [22], partly due to the lack of detailed information regarding the structure of PrP\textsuperscript{Sc}. In fact, defining the structure of infectious prion particles at high-resolution would allow the rational design of molecules that directly target PrP\textsuperscript{Sc} and inhibit its replication and accumulation into the nervous system. Unfortunately, this task is complicated by the wide heterogeneity and high propensity to aggregate of prions, aspects that make canonical high-resolution techniques such as NMR and X-ray crystallography inapplicable [8]. In an attempt to fill this gap, low-resolution methods have provided important experimental constraints illuminating the macroscopic architecture of prion fibrils and laying the groundwork for the application of computational techniques such as those used in this study. While these approaches still suffer from the lack of direct high-resolution information, they are undoubtedly valuable for interpreting the available information and inspire new hypotheses and approaches. Indeed, the new model described in this study possesses several important features. First, it displays an intrinsic coherence with state-of-the-art knowledge about infectious PrP\textsuperscript{Sc} fibrils and it appears to be as stable as the naturally-occurring prion Het-s [3]. Importantly, it also represents a unique workbench for interpreting future structural data or available biological evidence. In fact, it will be possible to test the effect of variations in the PRNP gene on the stability of the model by inferring mutations or polymorphisms known to favor or inhibit prion propagation and analyze the resulting structures by MD. As an example, it is known that the presence of an isoleucine at the polymorphic site 109 (M/I) in Bank vole PrP results in a substantially higher susceptibility to prion disease [23]. Assuming that the structure of prions in mouse and Bank vole are similar, as suggested by the difference in just 9 residues between murine and Bank vole PrP, and in light of the known higher propensity of isoleucine to participate in \boldsymbol{\beta}-strands as compared to methionine [24], our model may explain the effect of the 109 polymorphism on prion replication in Bank vole. Our 4R\boldsymbol{\beta}\boldsymbol{S} structure could in principle be adapted to model prions from Bank vole, human, deer, ovine and other species, as well as be employed to test how the effect of known polymorphisms and mutations affect prion stability and/or propagation. Similarly, it will be possible to rationally predict novel amino acid substitutions that could influence the stability of the 4R\boldsymbol{\beta}\boldsymbol{S} architecture and test their effects in surrogate in vitro (e.g. PMCA) or cell-based assays for prion replication. As a result, these strategies will produce new experimental constraints, which in turn will help to further improve the accuracy of the model, leading to an iterative cycle of prediction, testing and refinement. While waiting for a technological innovation that will definitively solve the structure of infectious prions, this strategy appears as the most feasible and promising among the available options.
Using the 4RβS model to address fundamental questions in prion biology

Since the original hypothesis of prions as protein particles responsible for transmissible brain pathologies, later evolved into a much wider concept of protein-based inheritance, tens of laboratories have intensively investigated the molecular features of these infectious agents. However, despite enormous efforts worldwide, many important questions of prion biology remain unanswered. For example, we still don’t fully understand the nature of the biological barriers that disfavor or fully impede the propagation of prions among different species. The reconstruction of prion conversion described in this manuscript could provide a unique opportunity to try to address this issue. For example, it will be possible to test the extent by which our templated conversion model tolerates sequence mismatches between the 4RβS template and the PrP substrate. These analyses could be used to interpret available biological evidence, for example the unusual ability of Bank vole PrP molecules to escape species barriers. Interestingly, our model may also help to improve current understanding of the prion strain phenomenon, which until now has been generally conceptualized as the same PrP polypeptide adopting different conformations. We show that the 4RβS scaffold can accommodate slightly different sequence threading schemes while remaining coherent with low-resolution experimental constraints and maintaining structural stability (S6 Fig). This observation suggests surprising flexibility of the 4RβS architecture to the alignment of residues in the β-strands, a feature that may introduce small conformational variability in each PrPSc monomer and result in large structural effects at the level of fibrils. This conclusion implies that the different biochemical and biological features of prion strains can be encoded in the small differences of PrP sequence alignment along the 4RβS scaffold.

The 4RβS model inspiring therapy

Computer-aided technologies are among the most promising strategies for the discovery of novel therapeutics against a wide variety of human illnesses, including prion diseases. In most cases, these approaches require the availability of high-resolution information about the target protein, but in absence of experimentally solved structures, as is the case for PrPSc, may also be applied to structural models. From this standpoint, our 4RβS may represent a unique opportunity to rationally design new anti-prion compounds. These molecules may be directed against two main structural elements of a putative 4RβS fibril: the “sticky” end, which exposes unpaired β-strands to the solvent, or the lateral surface. In the former case, the compound could block the insertion of a new PrP polypeptide into the growing fibril, while in the latter it would act as a stabilizer and prevent breakage into smaller, more diffusible elements. Interestingly, a similar mechanism has already been proposed to explain the anti-prion effects of luminescent conjugated polythiophenes (LCPs), molecules presenting consecutive thiophene rings exposing regularly spaced carboxyl side groups which could promote the interaction with lateral surfaces of prion fibrils [25]. It will undoubtedly be interesting to use our 4RβS structure to model the binding architecture of LCPs, as such information could then be exploited to optimize these compounds and/or rationally design immunological mimics and immunogens giving rise to PrPSc-specific antibodies or prophylactic vaccines against prion diseases.

Materials and methods

Template selection & model building

In order to accommodate the moPrP 89–230 sequence, we selected a β-solenoid architecture capable of satisfying the following requirements: (i) Approximate number of residues in extended conformation higher than 12 per rung. Requirement needed to fit the secondary
structure content of approximately 40% of β-sheets[8]; (ii) Capacity of accommodating two extended loops per rung. Namely, the possibility of introducing arbitrarily long sequences between consecutive β-strands in order to allow the exclusion of PK cleavage sites, glycine tracts and prolines from the core of the solenoid; (iii) L or T cross-section solenoidal-shape. Architecture coherent with cryo-EM maps; (iv) Capacity of accommodating bulky aromatic residues in the hydrophobic core (e.g. moPrP 89–230 contains 11 Tyr, 3 Phe and 1 Trp). All these requirements are satisfied by the right-handed, L-shaped β-solenoid architecture. In contrast, left-handed-solenoid structures typically display smaller rungs[26], impairing the modelling of the desired number of residues in extended conformation as well as the accommodation of bulky residues in the core. The left-handed structure of the prion forming domain of Het-s (PDB 2KJ3 and 2RN) is an exception in terms of β-sheet content, however its hydrophobic core is mainly composed of small side-chains and it allows the insertion of only one arbitrarily long loop connecting two consecutive rungs[3]. The threading scheme for the PrP 89–230 sequence (S1 Fig) was obtained by following the L-shaped β-solenoid architecture. In the threading process, PK cleavage sites, prolines and glycine tracts were positioned in the loops (when this was not possible, at the edges of beta strands). Charged side-chains were excluded from the inner core of the solenoid or paired with residues forming salt bridges. The presence of an intact disulphide bond between C178 and C213 and solvent exposure of N180 and N196 sidechains (in order to accommodate glycans) were also considered.

The 3D structure of the monomeric model of PrPSc was constructed following these steps: (i) 4 Rungs of Dickeya dadantii Pectate Lyase (PDB 1AIR; two repetitions of residues 168–235) were used to obtain the scaffold for the β-solenoid core; (ii) The original loops of the protein were removed; (iii) The residues in the hydrophobic core of the original PDB structure were replaced with PrP residues (as indicated in S1 Fig) using Chimera [27]; (iv) New loops were generated using the MODELLER tool[28] in Chimera. Each loop was selected from a set of 20 proposed conformations. Structures resulting in extended atomic clashes were discarded, and finally the best performing model in term of DOPE score was selected; (v) Side-chain geometry was optimized using the Dunbrack’s rotamer library. In particular, for highly polar and charged side-chains in the hydrophobic core, geometries capable of forming hydrogen bonds or salt bridges with nearby residues were selected; (vi) The energy of the system was minimized in vacuum with the steepest descent algorithm in Gromacs 4.6.5[29]. System topology was generated using Amber99SB-ILDN force field[17]. A restraining potential during energy minimization between the H and O atoms involved in hydrogen bond formation between backbone residues was added, as in Eq.1:

\[
\begin{align*}
\left( r_{ij} \right) = & \begin{cases} 
0 & \text{for } r_0 < r_0 \\
\frac{1}{2} k_d (r_{ij} - r_0)^2 & \text{for } r_0 \leq r_{ij} < r_1 \\
\frac{1}{2} k_d (r_1 - r_0) (2r_{ij} - r_1 - r_0) & \text{for } r_{ij} > r_1
\end{cases}
\end{align*}
\]

Where \( r_{ij} \) is the distance between the H and the O atoms involved in hydrogen bond formation; \( r_0 \) is the original hydrogen bond distance (2 < \( r_0 < 2.5 \)) Å and \( r_1 \) is an upper limit equal to 4 Å, while \( k_d \) is set to \( 2 \cdot 10^7 \) kJ/(mol·nm²).

This strategy was applied to favor the movement of the side-chains and accommodation of loops, while impairing the backbone deviation of the residues involved in the β-solenoid core formation; (vi) Additional optimization of the backbone (in order to remove Ramachandran outliers) and side-chain geometry was performed using Coot[30]; (vii) Absence of steric clashes was verified using Chimera, setting the VDW-overlap threshold as 0.6 Å, subtracting...
0.4 Å to account H-Bonding pairs and ignoring contacts of pairs <4 bonds apart. The 3D structure of the tetrameric model of PrP\textsuperscript{Sc} was assembled by stacking four monomers in a head-to-tail fashion using Chimera, maintaining the proposed threading. The strands 198-TETD and 215-TQYQKESQAYY were stacked on the top of 94-THNQ and 105-KTNLKHVAGAA of the forthcoming monomer, respectively. The structure was then energy minimized following the same protocol used for the monomer. Glycans (DManp$\alpha_1$-6DManp$\alpha_1$-3DManp$\beta_1$-4DGlcpNAc$\beta_1$-4[LFucp$\alpha_1$-6]DGlcpNAc$\alpha_1$-Asn) attached to residues N180 and N196 were added using the doGlycans tool to the 4R\textbeta S tetramer, and the structure was energy minimized in vacuum and then explicit solvent using Gromacs 5.1.2.

**Molecular dynamics simulations**

Molecular Dynamics simulations were performed on the supercomputers Finis Terrae II (CESGA, Santiago de Compostela, Spain) and Marconi (CINECA, Bologna, Italy) using Gromacs 5.1.2. The following protocol was applied for the whole set of simulations performed in this work. Protein topologies were generated using Amber 99SB-ILDN force field. The structures were accommodated in a dodecahedral box with periodic boundary conditions. The minimum wall distance from the protein was set to 12 Å. The box was filled with TIP3P water molecules. The charge of the system was neutralized with the addition of Na$^+$ or Cl$^-$ ions. The system was energy minimized in explicit solvent using a steepest descent algorithm, retaining the restraining potential (Eq 1). From the minimized structure, three independent equilibrations with position restraints on heavy atoms were launched: in the NVT ensemble (300 K) for 500 ps using the Nose-Hoover thermostat, and then in the NPT ensemble (300 K, 1 Bar) for 500 ps using the Nose-Hoover thermostat and the Parrinello-Rahman barostat. For each equilibrated system, a 20 ns MD simulation with restraining potential (Eq 1) was launched. Finally, restraints were released and each trajectory was extended with 100 ns of plain-MD. This protocol yielded three independent, unbiased 100 ns MD trajectories for each structure of interest. MD simulations were performed using a leap-frog integrator with step equal to 2 fs. LINCS algorithm was applied to handle bonds constraints. Cut-off for short-range Coulomb and Van der Waals interactions was set to 12 Å, while long range electrostatics was treated using Particle Mesh Ewald.

**Generation of the propagation model**

A key step to perform reliable rMD simulations is the identification of a reasonable reaction coordinate. To this end, we first developed a statistical coarse-grained model which enables us to identify the dominant reaction pathway underlying the conversion of PrP\textsuperscript{C} into a 4R\textbeta S PrP\textsuperscript{Sc}. In order to describe the reaction kinetics we developed a simple stochastic model in which the key rate-limiting processes are assumed to be the irreversible formation and docking of the four rungs of the 4R\textbeta S. We indicated R\textsubscript{0} as the C-terminal rung of the pre-formed 4R\textbeta S, and with R\textsubscript{1}, R\textsubscript{2}, R\textsubscript{3}, R\textsubscript{4} the consecutive rungs of the converting monomer. The instantaneous state of the system can be represented as a 4-dimensional vector of binary entries $S = [n_0, n_1, n_2, n_3]$, where $n_k = 1$ in the presence of docking between rung $R_k$ and rung $R_{k+1}$, and $n_k = 0$ otherwise. We emphasize that this model excludes the presence standalone rungs, which would correspond to an entropically-unfavorable single extended conformation, not stabilized by hydrogen bonds of nearby $\beta$-strands. On the other hand, misfolded rungs can be stabilized either upon docking to a pre-existing misfolded region (template mechanism) or through a process in which two rungs simultaneously form and dock. We modeled the transition of an initial state $S_0 = [0,0,0,0]$ (in which the PrP\textsuperscript{C} monomer is in the native state and none of the rungs are formed) to the fully misfolded state $S_P = [1,1,1,1]$ (where the monomer is completely...
misfolded and incorporated into PrP\textsuperscript{Sc}). The resulting network is represented in S8A Fig which contains all the possible combinations of docking events leading S\textsubscript{R} to S\textsubscript{P} through a sequence of irreversible transitions. The model can be simplified considering that rate \( k_2 \) is expected to be negligible as compared to \( k_0 \). Indeed, while the event associated with \( k_0 \) only requires the structuring of a disordered PrP region, the events associated with \( k_2 \) require the loss of native content (breakage of hydrogen bonds in the helical regions) together with the simultaneous formation of two rungs (two-fold entropic cost). Thus, it is possible to disregard all the reaction pathways in which the first step of the reaction is a transition occurring at a rate \( k_2 \) and consider only events starting from \([1,0,0,0]\). The network of the resulting simplified model is depicted in S8B Fig. Furthermore, we can set \( k_3/k_1 \gg 1 \) and \( k_3/k_2 \gg 1 \), since the docking of two pre-formed rungs occurs at a rate much faster than all processes involving misfolding. The reaction kinetics in this stochastic model was simulated through a Kinetic Monte Carlo algorithm (arbitrarily setting \( k_3/k_1 = 10^6 \)) and the resulting reaction mechanisms were studied as a function of the \( k_2/k_1 \) ratio. Namely, we enumerated all the reaction pathways and computed the corresponding probability:

| Step 1   | Step 2   | Step 3   | Step 4   |
|----------|----------|----------|----------|
| Path 1 [1,0,0,0] | [1,1,0,0] | [1,1,1,0] | [1,1,1,1] |
| Path 2 [1,0,0,0] | [1,0,1,0] | [1,1,1,0] | [1,1,1,1] |
| Path 3 [1,0,0,0] | [1,0,0,1] | [1,0,1,1] | [1,1,1,1] |
| Path 4 [1,0,0,0] | [1,0,0,1] | [1,1,0,1] | [1,1,1,1] |
| Path 5 [1,0,0,0] | [1,1,0,0] | [1,1,0,1] | [1,1,1,1] |
| Path 6 [1,0,0,0] | [1,0,1,0] | [1,0,1,1] | [1,1,1,1] |

We find that Path 1 (which consists in the consecutive formation of all rungs by templating on previously misfolded structures) dominates over all others as soon as \( k_1/k_2 \geq 4 \). Using Kramers' theory and assuming comparable pre-factors, we find that the templating mechanism is the most prominent reaction pathway when the activation energies for single and double rung formations obey the relationship:

\[
\ln \frac{k_1}{k_2} = \Delta G_2^i - \Delta G_1^i \gg 1.5 k_B T \quad \text{(Eq 2)}
\]

We expect this condition to be always satisfied. Indeed, processes 1 and 2 are characterized by the formation of the same number of hydrogen bonds, leading approximately to the same enthalpic gain. However, the process described by rate \( k_2 \) requires the breaking of the double amount of native contacts, together with a two-fold entropy loss (compared to the single rung formation) when forming two de novo rungs. We can therefore conclude that the propagation reaction proceeds through a subsequent formation of individual rugs, using as template pre-existing 4RβS free end. We emphasize that our approach is likely to underestimate the dominance of the sequential misfolding mechanism. Indeed, it does not account for the direct cooperativity of hydrogen bonds and the long-range electrostatics favoring \( \beta \)-strand formation in presence of pre-formed \( \beta \)-sheets, as directly supported by previous computational and experimental evidence [31, 32].

The coarse-grained information about the reaction mechanism obtained by means of our theoretical model can be exploited to set up a fully atomistic rMD simulation of the PrP\textsuperscript{C} \( \rightarrow \) PrP\textsuperscript{Sc} transition, using the 4R\textbeta\textsuperscript{S} as a target structure (Fig 4 and S1 Movie). The 3D structure of moPrP\textsuperscript{C} (residues 105–230) was obtained by linking moPrP 121–231 (PDB 1XYX) to the
adapted N-terminal fragment (residues 105–120) of huPrP^C (PDB 5yj5) which was mutated to the moPrP sequence. The initial state for the conversion simulation was generated by modifying a central dimer extracted from the tetrameric 4RβS at the end of 20 ns of restrained molecular dynamics simulation. The initial contact point between PrP^Sc and PrP^C was generated by leaving residues 89–104 of the C-terminal monomer anchored to the β-solenoid, which was then replaced by moPrP^C retaining the original disulfide bond. The rationale behind this modelling scheme derives from multiple previous reports indicating that the same region is a primary contact point between PrP^C and PrP^Sc [33]. The complex was energy minimized in vacuum and then in explicit solvent in a dodecahedral box (with periodic boundary conditions and minimum wall distance of 20 Å) also containing six Cl^- ions to neutralize the charge of the system, using a steepest descent algorithm (protein topology was generated by Amber99SB-ILDN force field). The system was then equilibrated in the NVT ensemble (350 K) for 200 ps using the Nose-Hoover thermostat, and then in the NPT ensemble (350 K, 1 Bar) for 200 ps using the Nose-Hoover thermostat and the Parrinello-Rahman barostat. The model was then subjected to a modified protocol of rMD simulations, adapting the method to a sequential biasing. This scheme resulted in a progressive rMD simulation in which the structure of PrP^C was targeted to the growing solenoid in a rung-by-rung fashion. Target structures included an entire 4RβS solenoid and additional rungs at the growing C-terminus: Step 1, residues 89–115; Step 2, residues 89–151; Step 3, residues 89–190; Step 4, residues 89–230. In this rMD scheme, the equation of motion is modified by an history-dependent biasing force \( F_i^{\text{rMD}}(X,t) \), defined as:

\[
F_i^{\text{rMD}}(X,t) \equiv -k_R \nabla z(X)(z[X(t)] - z_m(t))\theta(z[X(t)] - z_m(t))
\]

where \( k_R \) is the ratchet spring constant determining the strength of the biasing force applied on the system, in this case \( 10^{-2} \) kJ/mol. \( \theta \) function is equal to 1 if its argument is positive, its value its 0 otherwise. \( Z[X(t)] \) is a collective coordinate defined as:

\[
z[X(t)] = \sum_{i<j \geq 35} \left( C_{ij}[X(t)] - C_{ij}^0 \right)^2
\]

where \( C_{ij}[X(t)] \) is the instantaneous contact map and \( C_{ij}^0 \) is the contact map of the target state. Their entries are chosen to smoothly interpolate between 0 and 1, depending on the relative distance of the atoms \( i \) and \( j \). The \( z_m \) function indicates the smallest value assumed by the reaction coordinate \( z[X(t)] \) up to time \( t \) and.

The contact map \( C_{ij}(X) \) is defined as:

\[
C_{ij}(X) = \begin{cases} 
1 - \left( \frac{r_{ij}}{r_0} \right)^6 & \text{if } r_{ij} < r_c \\
1 - \left( \frac{r_{ij}}{r_0} \right)^{10} & \text{if } r_{ij} > r_c \\
0 & \text{if } r_{ij} = r_c \\
6/10 & \text{if } r_{ij} = r_c
\end{cases}
\]

Where \( r_{ij} \) is the distance between the \( i \)-th and the \( j \)-th atom; \( r_0 \) is a typical contact distance set to 7.5 Å; \( r_c \) is an upper limit to improve computational efficiency by excluding excessively distant atoms from the calculation (and it is set to 12.3 Å). In this way, no bias force acts on the system when the protein spontaneously proceeds towards the target state, while the external biasing force is only applied when the polypeptide tends to backtrack toward the initial state. We terminated each rMD simulation when the RMSD of the protein relative to the final state is stably lower than 0.8 Å. rMD simulations were performed in explicit solvent using Gromacs.
4.6.5 with the plugin Plumed 2.0.2 [34]. Integration of motion was performed using a leap-frog algorithm with step equal to 2 fs. Temperature was maintained at 350 K (approximately PrP melting temperature) and pressure at 1 Bar using Nose-Hoover thermostat and the Parrinello-Rahman barostat. LINCS algorithm was applied to handle bonds constraints. Cut-off for short-range Coulomb and Van der Waals interactions was set to 10 Å, while long range electrostatics was treated using Particle Mesh Ewald.

**Data analysis**

Root Mean Squared Deviation of atomic positions (RMSD) was calculated “all-atoms” using Gromacs 5.1.2. The specific calculation of the β-solenoid/helix core RMSD involves all the following residues for the different systems:

4RβS Model: T94, H95, N96, Q97, K105, T106, N107, L108, K109, H110, V111, A112, G113, A114, A115, M128, L129, G130, S131, G141, N142, D143, W144, E145, D146, R147, Y148, Y149, R150, E151, D166, Y168, S169, N180, I181, T182, I183, K184, Q185, H186, T187, V188, T189, T190, T198, E199, T200, D201, T215, Q216, Y217, Q218, K219, E220, S221, Q222, A223, Y224, Y225.

Het-s Dimer: R225, N226, S227, A228, K229, D230, I231, R232, T233, E234, E235, R236, A237, R238, V239, Q240, L241, G242, V244, T261, N262, S263, V264, E265, T266, V267, V268, G269, K270, G271, E272, S273, R274, V275, L276, I277, G278, N279, E280.

LβH Model: G89, Q90, G91, G92, G93, T94, H95, N96, Q97, W98, N99, K100, N107, L108, K109, H110, V111, A112, G113, A114, A115, A116, G118, A119, V120, V121, G122, G123, L124, G125, G126, T127, M128, L129, G130, S131, A132, M133, S134, R135, P136, M137, I138, H139, F140, G141, N142, D143, W144, E145, D146, D166, Q167, Y168, S169.

Secondary structure content and fraction of native/misfolded contacts were obtained using the Timeline and the Trajectory tools in VMD 1.9.2. Fraction of native and misfolded contacts are defined in the following equation

\[
S(X) \equiv \frac{\sum_{i,j>4} \theta(r_0 - r_i(X))\theta(r_0 - r_j(X))}{\sum_{i,j>4} \theta(r_0 - r_i(X))} \tag{Eq 6}
\]

where \(r_{ij}\) is the distance between the \(i\)-th and \(j\)-th \(C_\alpha\) atoms, \(r_0\) is the typical distance defining a contact between two residues and it is set at 7.5 Å, \(X\) is the instantaneous configuration of the protein during the simulation, \(X_\text{S}\) is the atomic configuration of the protein in the reference state, that is the native PrP\textsuperscript{Sc} structure for the native contacts calculation, and the PrP\textsuperscript{Sc} structure for the misfolded contacts calculations. This quantity is evaluated only for the converting monomer.

Initial and final values of RMSD and secondary structure content were calculated by averaging the values over the last 5ns (of restrained or unrestrained MD) and then calculating the mean and standard deviation for the three trajectories. Inter-monomer distances in the tetramers are calculating using Chimera as an average of the distances between the mid-residue of each β-strand and the same residue in the first or the second next monomer. In particular, for β-strand-1: N96, G130, Y168, T200; for β-strand-2: N107, D143, T192, Y217; for β-strand-3: G113, Y149, V188, A223. Tetramer volumes were calculated using Chimera. Molecular graphics images were produced using the UCSF Chimera package from the Computer Graphics Laboratory, while graphs are plotted using Gnuplot.
Supporting information

S1 Fig. 2D threading scheme of moPrP residues 89–230. 2D arrangement based on the general architecture of right-handed β-solenoid proteins with L-shaped cross section. Blue thick arrows indicate β-strands, thin cyan and thin orange arrows indicate the sidechain orientation (toward the solvent or toward the hydrophobic core, respectively). The scheme is used to thread the moPrP 89–230 sequence by considering different constraints. PK cleavage sites identified by mass spectrometry in two different previous reports are colored in red. Glycosylation sites are labelled in green. Prolines are colored in purple. Cystine is indicated in cyan. (TIF)

S2 Fig. 4RβS model of PrPSc superimposed to the Cryo-EM map. The pictures illustrate that the size of 4RβS fits with the protofilament cross section (50 Å x 30 Å) obtained by cryo-EM. (A) Level of contouring was set to 3.05 to match the measured protofilament diameter as determined from the raw electron micrograph [4]. (B) Level of contouring was tightened to 3.60 to highlight the superimposition of the hydrophobic core of the 4RβS with the core electron density region of the protofilament. (C) Reconstruction of a PrPSc protofilament superimposed to the cryo-EM map. (D) Cross-sectional view of two PrPSc protofilaments compared with the orientation reported in [4]. (TIF)

S3 Fig. Summary of restrained MD simulations of the Het-s dimer, 4RβS and LβH PrPSc. Upper graphs report the RMSD deviation from initial configurations for the entire structure (blue lines) or the β-solenoid core (orange lines) of the different proteins during the 20 ns of restrained MD simulations. Filled curves represents standard error of the mean. The graphs indicate a minor rearrangement of the β-solenoid core that is almost identical for the Het-s dimer and the 4RβS model, characterized by a RMSD of 2.3 ± 0.2 Å and 2.4 ± 0.2 Å respectively (calculated as the average of the three trajectories over the last 5 ns, ± standard deviation). In contrast, the hydrophobic core of the LβH model displays a higher RMSD deviation (3.3 ± 0.3 Å). Lower graphs indicate the α-helical (blue lines) or β-sheet (red lines) content of each protein. The latter is stable for all the three structures, likely due to the presence of the restraining potential for the whole length of the three simulations. (TIF)

S4 Fig. Probability distribution of β-sheet content for the Het-s dimer, 4RβS and LβH PrPSc. Graphs show the probability of each residue to be in an extended conformation at the beginning of the simulation (green) or at the end of the simulation (purple) for the Het-s dimer, 4RβS and LβH PrPSc models. Initial or final probabilities were calculated from the last 5 ns of the restrained or unrestrained simulations, respectively. (TIF)

S5 Fig. Initial and final conformations of LβH PrPSc after MD simulations. Pictures show the energy minimized structure (1), the same structure at end of the restrained (2) or unrestrained (3) MD simulations of the LβH model. (TIF)

S6 Fig. Alternative threading schemes for the 4RβS. Panels A and B represent two possible alternative threading schemes not violating available experimental constraints. Only small deviations from the main model are allowed to agree with experimental data. Images (i) represents the 2D schemes of two putative alternative schemes. Images (ii) highlight the structure of the rung that differs from the main threading scheme, blue residues represent the shifted amino acids in the threading. Graphs (iii) report RMSD and secondary structure analysis of three 100
ns unrestrained MD simulations. Images (iv) report the initial and final frame of the respective MD simulations.

(TIF)

S7 Fig. Illustration of the energy minimized 4RβS tetramer carrying complex glycans. A model of energy minimized tetrameric 4RβS structure carrying glycan residues is depicted in two different orientations (A, B). Top view of a laterally stacked trimer of glycosylated 4RβS (C), compatible with the 2D crystals diffraction data. A scheme of the complex glycan precursor added is depicted in (D), blue squares indicate N-acetylglucosamine, green circles indicate mannos and the red triangle indicate fucose.

(TIF)

S8 Fig. Statistical model for prion propagation mechanism. (A) Schematic representation of the network of transitions leading to the incorporation of PrP\textsuperscript{C} into PrP\textsuperscript{Sc}. The purple arrow describes the formation of a single rung by templating the unstructured region of PrP\textsuperscript{C} (89–115) on PrP\textsuperscript{Sc} (rate \(k_0\)). The blue arrows describe the formation of a single by means of a templating process which involves the breaking the native contacts and the docking onto a preformed rung (rate \(k_1\)). The orange arrows indicate the spontaneous formation and docking of two de-novo rungs (rate \(k_2\)). Finally, the green arrows indicate the docking of preformed rungs belonging to two adjacent misfolded regions of the same chain (rate \(k_3\)). (B) Simplified version of the network assuming as a priming reaction step the formation of the first rung. (C) Relative probability of the 6 reaction pathways as a function of the \(k_1/k_2\) rate ratio.

(TIF)

S1 Movie. Full-atomistic model of prion propagation. Visualization of the rMD simulation reconstructing the entire refolding events of PrP\textsuperscript{C} onto the 4RβS PrP\textsuperscript{Sc} model.

(MP4)

S1 File. Atomic coordinates of the model.

(PDB)

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Author Contributions

Conceptualization: Giovanni Spagnolli, Pietro Faccioli, Holger Wille, Emiliano Biasini, Jesús R. Requena.

Data curation: Giovanni Spagnolli.

Formal analysis: Giovanni Spagnolli, Simone Orioli, Pietro Faccioli, Emiliano Biasini.

Funding acquisition: Emiliano Biasini, Jesús R. Requena.

Investigation: Giovanni Spagnolli, Marta Rigoli, Simone Orioli, Alejandro M. Sevillano, Pietro Faccioli, Jesús R. Requena.

Methodology: Giovanni Spagnolli, Pietro Faccioli, Emiliano Biasini.

Project administration: Emiliano Biasini.

Software: Giovanni Spagnolli, Pietro Faccioli.
Supervision: Pietro Faccioli, Emiliano Biasini, Jesús R. Requena.

Visualization: Giovanni Spagnolli, Marta Rigoli.

Writing – original draft: Giovanni Spagnolli, Emiliano Biasini.

Writing – review & editing: Pietro Faccioli, Holger Wille, Jesús R. Requena.

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