Asparagine and glutamine ladders promote cross-species prion conversion

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Prion transmission between species is governed in part by primary sequence similarity between the infectious prion aggregate, PrPSc, and the cellular prion protein of the host, PrPC. A puzzling feature of prion formation is that certain PrPC sequences, such as that of bank vole, can be converted by a remarkably broad array of different mammalian prions, whereas others, such as rabbit, show robust resistance to cross-species prion conversion. To examine the structural determinants that confer susceptibility or resistance to prion conversion, we systematically tested over 40 PrPC variants of susceptible and resistant PrPSc sequences in a prion conversion assay. Five key residue positions markedly impacted prion conversion, four of which were in steric zipper segments where side chains from amino acids tightly interdigitate in a dry interface. Strikingly, all five residue substitutions modulating prion conversion involved the gain or loss of an asparagine or glutamine residue. For two of the four positions, Asn and Gln residues were not interchangeable, revealing a strict requirement for either an Asn or Gln residue. Bank voles have a high number of Asn and Gln residues and a high Asn:Gln ratio. These findings suggest that a high number of Asn and Gln residues at specific positions may stabilize β-sheets and lower the energy barrier for cross-species prion transmission, potentially because of hydrogen bond networks from side chain amides forming extended Asn/Gln ladders. These data also suggest that multiple PrPSc segments containing Asn/Gln residues may act in concert along a replicative interface to promote prion conversion.

Prion diseases are fatal neurodegenerative disorders of humans and animals caused by prion protein aggregates accumulating in the brain and spinal cord (1). β-Sheet–rich prion aggregates, known as PrPSc, template the misfolding of the cellular prion protein monomer, PrPC, similar to seeding mechanisms that occur with other amyloidogenic proteins such as amyloid-β, α-synuclein, and islet amyloid polypeptide (2, 3). PrPSc–templated conversion of PrPC monomers typically requires a high degree of sequence similarity (4–6); however, conversion of dissimilar PrP sequences can occur and induce prion disease in other species; for example, bovine spongiform encephalopathy prions have infected humans, cats, and zoo bovids (7, 8).

Mammalian PrPC is highly conserved in both sequence and structure, consisting of ~210 amino acids with a disordered N terminus (residues 23–120) and a globular, C-terminal domain arranged as three α-helices and a short anti-parallel β-sheet (9, 10). We found that conversion of mouse or human PrPC by elk chronic wasting disease (CWD)3 prions occurs efficiently when the PrPSc sequence within a loop segment rich in polar and aromatic residues, the β2–α2 loop, is mutated to match the elk sequence (VDQYNNQNTF) (11, 12). Further studies of the β2–α2 loop sequence revealed that substituting Tyr169 with glycine, leucine, or glutamine inhibited prion conversion, whereas substitutions of bulky aromatic residues (Y169W and Y169F) enabled conversion, indicating a requirement for highly specific amino acid side chain properties, in this case, an aromatic side chain (13). Collectively, these findings suggest that prion conversion between different PrP sequences does not require an exact match in the side chain between PrPC and PrPSc; however, side chain complementarity in amyloid-prone segments is essential. The β2–α2 loop has been identified as a steric zipper segment of PrP, in which side chains emerging from two β-sheets tightly interdigitate, forming a dry interface (14). Although residue mismatches between species may diminish steric zipper formation, polar, hydrophobic, and aromatic residues in steric zipper segments may also stabilize early aggregates via aromatic residue stacks, serine stacks, and asparagine ladders and thus may promote conversion to a β-sheet–rich isoform (15, 16).

The bank vole is a rodent that has proven remarkably susceptible to a diverse array of prions from humans and animals, and bank vole PrPC has recently been termed the “universal accep-

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The abbreviations used are: CWD, chronic wasting disease; PMCA, protein misfolding cyclic amplification; clPMCA, cell-lysate PMCA; PK, proteinase K; Hu, human; Bv, bank vole; Rb, rabbit; ANOVA, analysis of variance; RML, Rocky Mountain Laboratory.
Asn/Gln ladders promote cross-species prion conversion

The amino acid sequences of bank vole, rabbit, and human PrPC differ at 27 positions in fully processed PrP (Fig. 1A), yet show highly similar secondary and tertiary structure by NMR spectroscopy (27–29). To identify the residues that impact prion cross-seeding, we first compared the conversion of three mammalian PrPC sequences with single or multiple substitutions. A variety of cross-seeded conversion assay, cell-lysate protein misfolding cyclic amplification (clPMCA) (12, 13, 26), to identify the key residue positions that promote or inhibit prion conversion. We systematically substituted single residues in bank vole, rabbit, and human PrPC and seed with dissimilar elk or mouse prions to examine the mechanism of prion conversion. Conversion of three mammalian PrPC sequences with single or multiple substitutions has revealed Asn/Gln residues in PrP, often within known steric zippers, as powerful promoters of prion conversion among dissimilar sequences, suggesting that within a β-sheet, the linear chains of hydrogen bonds among the amides in Asn/Gln side chains stabilize and enhance prion assembly.

Results

Replacing asparagine and glutamine residues in bank vole PrP<sup>C</sup> inhibits its conversion

The amino acid sequences of bank vole, rabbit, and human PrP<sup>C</sup> differ at 27 positions in fully processed PrP (Fig. 1A), yet show highly similar secondary and tertiary structure by NMR spectroscopy (27–29). To identify the residues that impact prion cross-seeding, we first compared the conversion of bank vole, human, and rabbit PrP<sup>C</sup> seeded by elk and mouse prions. The two mouse prions used, RML and 87V, differ in amino acid sequence in two positions (supplemental Fig. S1) and in the PrP<sup>Sc</sup> biochemical properties and disease phenotype in mice (30). Each PrP<sup>C</sup> sequence was expressed in PrP-deficient RK13 cells, and cell lysates were seeded with prions or were unseeded (supplemental Fig. S2) and subjected to clPMCA. Samples were then analyzed for proteinase K (PK)–resistant PrP<sup>Sc</sup> by Western blotting, using the anti-PrP 3F4 antibody epitope for detection of newly converted PrP<sup>Sc</sup> (31). Bank vole PrP<sup>C</sup> (BvPrP<sup>C</sup>) was readily seeded by elk CWD as well as mouse 87V and RML prions, whereas human (HuPrP<sup>C</sup>) and rabbit PrP<sup>C</sup> (RbPrP<sup>C</sup>) were not converted by any of the prions (Fig. 1B), confirming that the conversion efficiency reported for BvPrP<sup>C</sup>, HuPrP<sup>C</sup>, and RbPrP<sup>C</sup> (32–35) could be reproduced in the clPMCA assay.

To identify the HuPrP<sup>C</sup> residues that obstruct prion conversion, bank vole PrP<sup>C</sup> with human residue substitutions was used as a substrate in clPMCA (Fig. 1C). Fifteen residue substitutions were tested singly or grouped (positions 227–229) (Fig. 1A), five of which had a major effect on conversion. BvPrP<sup>C</sup> with the V166M or Q168E substitutions from HuPrP<sup>C</sup> (β2–α2 loop) markedly reduced conversion when seeded with mouse RML, mouse 87V, or elk CWD (∼10–50% conversion relative to BvPrP<sup>C</sup>) (Fig. 1, D and E). The N143S substitution (β1–α1 loop) was strongly inhibitory for RML and 87V mouse prions (14 and 7% conversion, respectively), but not CWD (>100% conversion) (Fig. 1, D and E). Additionally, N155H was inhibitory for 87V-induced conversion (47% conversion), and Q219E was inhibitory for RML-induced conversion (20% conversion) (Fig. 1, D and E). Given that all three PrP<sup>Sc</sup> sequences include Asn<sup>143</sup> and Gln<sup>219</sup> (supplemental Fig. S1), successful conversion is not likely due simply to a primary sequence match with PrP<sup>C</sup> but instead was influenced by the PrP<sup>Sc</sup> conformation. BvPrP<sup>C</sup> with 10 other human residue substitutions was converted to high PrP<sup>Sc</sup> levels by all three prions (Fig. 1, D and E), indicating that most human residues do not inhibit cross-seeding by CWD or certain mouse prions. Thus, substitution of two human PrP<sup>C</sup> residues (Met<sup>166</sup> and Glu<sup>168</sup>) inhibited conversion of BvPrP<sup>C</sup> by all prions tested, and three other substitutions (Ser<sup>143</sup>, His<sup>155</sup>, Glu<sup>219</sup>) inhibited prion conversion to 50% or less in a PrP<sup>Sc</sup> sequence- and/or conformation-specific manner.

Rabbits have resisted intracerebral challenge with Creutzfeldt–Jakob disease, kuru, sheep scrapie, and mouse-adapted scrapie prions (strain ME7) (32, 33), and are considered one of the most highly prion-resistant species. We next measured the conversion of BvPrP<sup>C</sup> having nine single or grouped rabbit-specific residue substitutions (Fig. 1C). The N100G substitution showed the most dramatic inhibition, with <10% conversion, whereas K220Q resulted in only 37–57% conversion by any prion (Fig. 1, F and G). The M138L, I184V, and M205I rabbit substitutions reduced CWD-seeded conversion to ~30, 40, and 25% conversion, respectively (Fig. 1, F and G). Additionally, N174S completely blocked mouse 87V-seeded conversion of BvPrP<sup>C</sup> but only reduced RML-seeded conversion to ~40% (Fig. 1, F and G). Thus, six of nine rabbit substitutions diminished conversion of bank vole PrP<sup>C</sup> by more than 50%. Collectively, these results demonstrate that human and rabbit residues impede prion conversion to varying levels, depending on the PrP<sup>Sc</sup> sequence and/or conformation. Remarkably, the five most inhibitory substitutions, N100G, N143S, Q168E, N174S, and Q219E, involved the loss of an asparagine or glutamine residue from the bank vole sequence, suggesting that Asn/Gln residues are important for the conversion of bank vole PrP.

Bank vole asparagine and glutamine residues enable prion conversion of human and rabbit PrP<sup>C</sup>

Because the N100G, N143S, Q168E, N174S, and Q219E substitutions inhibited conversion of BvPrP<sup>C</sup>, we reasoned that the converse bank vole amino acid substitutions may enable prion conversion of otherwise resistant HuPrP<sup>C</sup> and RbPrP<sup>C</sup> (Fig. 2A). We measured seeded conversion of HuPrP<sup>C</sup> having the S143N, E168Q, or E219Q substitutions and RbPrP<sup>C</sup> containing the G100N or S174N substitutions. HuPrP<sup>C</sup> with the S143N or E168Q substitutions seeded with 87V or CWD prions, respectively, led to low PrP<sup>Sc</sup> levels (Fig. 2, B and C), whereas E219Q had no effect on conversion (Fig. 2, B and C). In contrast, mouse RML prions did not convert any of the HuPrP<sup>Sc</sup> sequences with single substitutions (Fig. 2, B and C). To identify any other residues that impact cross-seeding of HuPrP<sup>C</sup>, we assessed 12 other Hu-to-Bv amino acid substitutions including the adjacent residues 219–220 and 227–229. Only the Q168E and S170N substitutions in HuPrP<sup>C</sup> resulted in modest conversion when seeded by CWD prions (Fig. 2, B and C), as previously reported (12), whereas no other substitution enabled conversion by mouse RML or 87V prions.
We next measured conversion of RbPrPC with the bank vole substitutions G100N or S174N. Remarkably, the single G100N substitution resulted in efficient cross-seeding by CWD, 87V, and RML prions (100, 84, and 83%, respectively) (Fig. 2, D and E), indicating that the Asn100 residue is critical for conversion by all three prions. The single S174N substitution had minimal impact on conversion by RML, 87V, and CWD prions (4, 41, and 43%, of BvPrPC, respectively) (Fig. 2, D and E), whereas RbPrPC containing both G100N and S174N was converted to levels similar to RbPrPC containing the single G100N substitution (Fig. 2, D and E). These results suggest that certain asparagine and glutamine residues in BvPrPC (Asn100, Asn143, Gln168, Asn170) promote cross-species prion conversion, even in the context of a different PrP sequence.

**Bank vole substitutions variably promote conversion of human PrPC in a prion-dependent manner**

Because no single BvPrP substitution enabled conversion of HuPrPC by more than 50%, we next assessed conversion of HuPrPC having multiple substitutions. HuPrPC with amino acid substitutions from HuPrPC with amino acid substitutions from HuPrPC having multiple substitutions. HuPrPC with amino acid substitutions from HuPrPC having multiple substitutions. HuPrPC with amino acid substitutions from HuPrPC having multiple substitutions. HuPrPC with amino acid substitutions from HuPrPC having multiple substitutions. HuPrPC with amino acid substitutions from HuPrPC having multiple substitutions. HuPrPC with amino acid substitutions from HuPrPC having multiple substitutions. HuPrPC with amino acid substitutions from HuPrPC having multiple substitutions. HuPrPC with amino acid substitutions from HuPrPC having multiple substitutions. HuPrPC with amino acid substitutions from HuPrPC having multiple substitutions. HuPrPC with amino acid substitutions from HuPrPC having multiple substitutions. HuPrPC with amino acid substitutions from HuPrPC having multiple substitutions. HuPrPC with amino acid substitutions from HuPrPC having multiple substitutions.

Figure 1. Human and rabbit amino acid substitutions inhibit conversion of bank vole PrPC. A, alignment of the PrPC sequences from bank vole, human, and rabbit reveals amino acid sequence differences between species at 27 positions (labeled in red and blue), 21 of which were investigated here (in red). Locations of the β-strands and α-helices are shown. B, mouse RML, mouse 87V, and elk CWD prions convert bank vole BvPrPC, but not HuPrPC or RbPrPC. C, schematic for experimental approach. BvPrPC with HuPrPC or RbPrPC substitutions was seeded with mouse or elk prions, and the newly converted PrPSc was measured after 24 h. D, representative immunobots show conversion of BvPrPC with amino acid substitutions from HuPrPC. E, quantitative analysis shows that substitutions N143S, Q168E, and Q219E strongly inhibit BvPrPC conversion, depending on the PrPSc seed. F, representative immunobots show conversion of BvPrPC with amino acid substitutions from RbPrPC. G, quantitative analysis shows that the N100G and N174S substitutions strongly inhibit conversion, depending on the PrPSc seed. E' and G' show the quantified data with a segmented y axis. For A, the GenBank accession numbers for bank vole, human, and rabbit PrPC are AF367624, DQ408531, and U28334, respectively. Quantified data are from three to five independent experiments (E and G). The error bars indicate the observed variance. *, p < 0.05; **, p < 0.01; ***, p < 0.001, one-sample t test. One-way ANOVA with Tukey post hoc test revealed statistically significant differences for CWD between residue substitutions at positions 100 and 203, positions 138 and 225, and positions 205 and 225 (*, p < 0.05), as well as between residues 100 and 225 (**, p < 0.01).
plete bank vole β2–α2 loop sequence (M166V, E168Q, and S170N substitutions) was converted by CWD prions as efficiently as bank vole PrPC; no additional bank vole substitutions further enhanced conversion (Fig. 3, A and B). These results indicate that the bank vole β2–α2 loop sequence is necessary and sufficient for elk CWD cross-seeding of HuPrPC.

In contrast to elk CWD, mouse prions RML and 87V minimally converted HuPrPC containing the bank vole β2–α2 loop sequence (<15%; Fig. 3, A and B). Because the N143S substitution in the β1–α1 loop had inhibited BvPrP conversion (Fig. 1E), we next tested whether HuPrPC with the bank vole β1–α1 and β2–α2 loop segments (S143N, Y145W, M166V, E168Q, and S170N) would be converted by mouse 87V prions. These five-residue exchanges, which add three more Asn/Gln residues to the human sequence and remove a charged residue, led to substantially higher levels of HuPrPC conversion by 87V prions (101%), but not by RML prions (8%; Fig. 3, C and D). These results indicate that in two critical segments known to form steric zippers, the β1–α1 loop (positions 143–145) (36) and the β2–α2 loop (positions 166–170) (14), bank vole residues enable efficient conversion of HuPrPC seeded by mouse 87V prions. Because RML was not converted, these data also suggest that...
mouse RML and 87V prions differ in the number and location of PrP^C segments required for efficient conversion.

To identify the residues controlling conversion of HuPrP^C by RML prions, 10 additional BvPrP^C substitutions (in groups of 1–3) were incorporated into the HuPrP^C–bank vole PrP^C chimera and tested for conversion. Of the 10 residues tested, RML prions only converted to 20% PrPC having the additional I215V, E219Q, or E219Q, R220K substitutions (Fig. 3, C and D; note for Glu219, four of five measurements resulted in 28–35% conversion). These experiments indicate that four Asn/Gln substitutions, which were in the loop segments (Asn143, Gln168, and Asn170), and the C terminus of a-helix 3 (residue 219), enable limited conversion of human PrP^C by RML. The Val215 substitution also enhanced conversion by RML. Interestingly, HuPrP^C with the bank vole loop, together with the individual substitutions S143N, I215V, or E219Q, was not converted to high levels by RML prions (Fig. 3, A and B), indicating that exchanges in all three segments were essential for conversion of HuPrP^C by RML prions.

**Prion conversion enhanced by Asn/Gln substitutions is highly position- and side chain-dependent**

Strikingly, five of five single substitutions that inhibit conversion of BvPrP^C to less than 20% or that promote conversion of HuPrP^C and RbPrP^C to greater than 20% involved the loss or gain of Asn/Gln residues, respectively. Mammalian PrP^C from 24 species revealed 26–31 Asn/Gln residues scattered throughout the protein; bank vole PrP^C has 31 Asn/Gln residues, an unusually high number, and a notably high Asn:Gln ratio (0.94) (Tables 1 and 2). To determine whether the position of the Asn or Gln substitution within a segment impacts conversion, Asn/Gln substitutions were placed at sites flanking positions 143 and 219 in HuPrP^C. For RML and 87V prions, there was little to no conversion when the Asn143 substitution was transposed to
Interestingly, although leucine is somewhat structurally similar to glutamine in volume, the Leu143 and Leu219 substitutions in Hu-Bv PrPC were not converted by 87V or RML, respectively (0–1%) (Fig. 4, A–D), suggesting that the side chain hydrogen bonding of the amide in the Asn/Gln is critical to prion conversion. These results indicate that the strong promoting effect of the Asn100 and Asn143 residues on prion conversion is highly position- and side chain-dependent, requiring specifically an asparagine residue.

**Discussion**

Prion transmission can be exquisitely sensitive to primary sequence differences between PrPSc and PrPSc, because even one mismatched residue can obstruct prion conversion (37–39). Bank vole PrPSc, however, is extraordinarily susceptible to conversion seeded by prions from other species, despite sequence differences (19, 21, 24, 25, 35, 40–42). Here we have investigated the residues that govern cross-species prion conversion by systematically testing over 40 mutated bank vole, human, and rabbit PrPSc sequences seeded by three dissimilar PrPSc seed sequences. Our studies revealed five amino acid substitutions that profoundly promote or inhibit prion conversion, all of which involve the gain or loss of an asparagine or glutamine residue (positions 100, 143, 168, 170, and 174). Remarkably, substituting Asn/Gln residues into PrP conferred susceptibility to sequences that otherwise resisted conversion. The relevant positions of the Asn/Gln residues varied with the incoming prion seed, consistent with PrPSc conformation determining the critical PrPSc–PrPSc interaction domains. Additionally, we have also established that certain positions tolerate either an Asn or a Gln, whereas others strictly require an Asn or a Gln for prion conversion.

The addition of Asn/Gln residues resulted in a sequence match with the PrPSc seed (supplemental Fig. S1), which could explain why conversion was enhanced. However, six other substitutions led to a sequence match with the seed but did not promote conversion. Additionally, for at least two positions, 170 and 219, a mismatched polar residue also promoted conversion. Additionally, for at least two positions, 170 and 219, a mismatched polar residue also promoted conversion, indicating that the side chain structure did not require a precise match with the PrPSc seed but did require a structurally similar residue. Notably, the position of the Asn/Gln residue was also important, because there were Asn/Gln residues in positions that did not significantly impact prion conversion (S97N or H155N), even in combination with the bank vole

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**Table 1**

| Species                  | Asn | Gln | Asn/Gln total | Asn/Gln ratio |
|--------------------------|-----|-----|---------------|---------------|
| Rodents                  |     |     |               |               |
| Bank vole (Myodes glareolus) | 15  | 16  | 31            | 0.94          |
| Meadow vole (Microtus pennsylvanicus) | 15  | 16  | 31            | 0.94          |
| Deer mouse (Peromyscus maniculatus bairdii) | 15  | 16  | 31            | 0.94          |
| Syrian golden hamster (Mesocricetus auratus) | 15  | 16  | 31            | 0.94          |
| Chinese hamster (Cricetulus griseus) | 15  | 16  | 31            | 0.94          |
| Armenian hamster (Cricetulus migratorius) | 14  | 16  | 30            | 0.88          |
| Mouse (Mus musculus)     | 13  | 16  | 29            | 0.81          |
| **Primates**             |     |     |               |               |
| Black-handled spider monkey (Ateles geoffroyi) | 14  | 15  | 29            | 0.93          |
| Common marmoset (Callithrix jacchus) | 13  | 16  | 29            | 0.81          |
| Squirrel monkey (Saimiri sciureus) | 12  | 17  | 29            | 0.71          |
| Tufted capuchin (Sapajus apella) | 12  | 16  | 28            | 0.75          |
| Macaque (Macaca mulatta, Macaca fascicularis) | 11  | 16  | 27            | 0.69          |
| Human (Homosapiens)      | 11  | 16  | 27            | 0.73          |
| Chimpanzee (Pan troglodytes) | 10  | 16  | 26            | 0.63          |
| **Ruminants**            |     |     |               |               |
| Deer (Odocoileus sp.)    | 12  | 17  | 29            | 0.71          |
| Cow (Bos taurus)         | 12  | 17  | 29            | 0.71          |
| Sheep (Ovis aries) ARR/ARQ | 12  | 16/17 | 28/29 | 0.71–0.75 |
| Elk (Cervus canadensis)  | 12  | 16  | 28            | 0.75          |
| **Carnivores**           |     |     |               |               |
| Cat (Felis catus)        | 12  | 16  | 28            | 0.75          |
| Ferret (Mustela putorius furo) | 11  | 17  | 28            | 0.62          |
| Mink (Neovison vison)    | 11  | 16  | 27            | 0.69          |
| Raccoon (Procyon lotor) | 11  | 16  | 27            | 0.69          |
| Dog (Canis familiaris)   | 11  | 16  | 27            | 0.69          |
| **Leporids**             |     |     |               |               |
| Rabbit (Oryctolagus cuniculus) | 10  | 18  | 28            | 0.56          |
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β1–α1 and β2–α2 residues. Among the five key Asn/Gln residues, four residues (at positions 100, 168, 174, and 219) were located within steric zipper segments identified by the ZipperDB 3D profiling method using the seven-residue zipper (Rosetta energies below an energetic threshold of -27 kcal/mol) (Table 3), and two segments have been crystallized (positions 143 and 174) (14, 36). Taken together, these findings suggest that PrPCβ1–α1 and PrPCβ2–α2 loop interactions in short segments regulate prion conversion.

Prion seeding specificity is widely recognized to be controlled by one or two key residues (22, 23), for example, PrP residues 138/139 in the β1–α1 loop control seeding specificity between mouse and hamsters (38, 39) and residues 168 and 170 in the β2–α2 loop impact deer CWD–human transmission barriers (12). The mechanism underlying how residues in the β1–α1 and β2–α2 loop control conversion may be explained by microcrystal structures of the 138–143 and the 170–175 segments determined to less than 3 Å, where amino acid side chain

Table 3
Rosetta energy calculations for five steric zipper segments identified in the prion protein

| Asn/Gln position | Zipper segments identified by ZipperDB | Rosetta energy kcal/mol |
|------------------|----------------------------------------|-------------------------|
| 100              | QWKPSK                                 | -29.2                   |
| 143              | MTHGWNN                               | -26                     |
| 168              | QYNQQNN                                | -34.6                   |
| 174              | QYQNSQ                                 | -34.6                   |
| 219              | QYQRESQ                                | -31.2                   |

Figure 4. The impact of Asn/Gln residues on prion conversion is highly position-dependent. A and B, HuPrPβ1 with the BvPrPβ2 loop and the 143Q or 143L substitution is not converted by RML or 87V prions. Additionally, transposing the asparagine substitution from position 143 to flanking positions in HuPrPβ1 no longer promotes conversion by RML or 87V prions. C and D, HuPrPβ1 with the E219N, but not the E219L, substitution together with the BvPrPβ1–α1 and β2–α2 loop sequences is efficiently converted by RML prions. Shifting the glutamine substitution from position 219 to flanking positions does not promote conversion by RML prions. E and F, RbPrPβ1 with the 100Q substitution was minimally converted by RML, 87V, or elk CWD prions. G and H, HuPrPβ1 with the Gln170 substitution was converted by elk CWD prions. I, PrPβ1 in lysates used for A–H show similar PrP β levels. The blots in A, E, and I show single Western blots at the same exposure with intervening lanes removed for clarity. PrP β sequence changes are shown in red below immunoblots. Quantified data are from three to five (B), three or four (F), or three (H) independent experiments. Standard error bars indicate the observed variance. *, p < 0.05; **, p < 0.01, one-sample t test. One-way ANOVA with Tukey post hoc test revealed statistically significant differences for RML between Asn219 and residue Asn218, Asn220, Asn221, and Leu219 (*, p < 0.05).
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Figure 5. Proposed mechanism for prion conversion involving asparagine/glutamine residues in discrete segments of bank vole PrPC. A, a ribbon diagram of the NMR BvPrPC structure (Protein Data Bank code 2K56; Ref. 27) with critical Asn/Gln residues shown in magenta. The N-terminal domain of PrPC (residues 97–118, gray) is modeled using Rosetta software (63). Residues 23–96 are not shown. B, another view of the NMR BvPrPC structure rotated horizontally by 90°. C, zipper structure from PrPSc peptide (residues 170–175; Protein Data Bank code 3VFA; Ref. 36) fibril demonstrating that alignment of Asn/Gln residues along the length of the fibril axis promotes side chain hydrogen bonds in a motif known as an asparagine/glutamine-ladder. β-Strands are represented by gray arrows, Asn174 residues are shown in magenta, Gln219 residues are in cyan, and side chain hydrogen bonds in the asparagine/glutamine-ladder are indicated by black dotted lines.

These additional hydrogen bonds, more typically between amides of identical side chains in adjacent strands, greatly increase the stability of the β-sheet, because the parallel arrays of aligned bonds are hyperpolarized (47) and are even stronger than those in ice (48, 49). We suggest a model in which intermolecular interactions between complementary PrP segments that have a strong, hyperpolarized hydrogen bond network, such as the β2–α2 loop, stabilize newly incorporated PrP monomers onto β-sheet-rich fibrils, driving aggregation of dissimilar PrP sequences. These intermolecular contact sites may also serve as critical templating sites, directing PrP into a particular PrPSc fold. This model would be consistent with experimental findings from the β1–α1 and β2–α2 loop, where sequence differences can switch the prion conformation (11, 50, 51).

Several yeast prions, such as Ure2, Sup35, and Rnq1, contain Asn/Gln-rich domains that are required for assembly into functional prion fibrils (52, 53) and may mediate prion cross-seeding (54). The Asn/Gln residues at the N terminus of Ure2p and Sup35p play a particularly critical role in stabilizing the amyloid state (55). This Asn/Gln-rich domain is also modular, causing aggregation when introduced into other proteins (56). By comparison, mammalian prions contain one short Asn/Gln-rich domain (β2–α2 loop), with additional Asn/Gln residues interspersed throughout the protein. Thus, the Asn/Gln residues within PrP are not constrained to a single Asn/Gln-rich domain but seem to collectively promote conversion from multiple segments. Stacking of Asn/Gln residues into ladders along the length of a prion fibril would maximize interstrand hydrogen bonding and would be compatible with a parallel, in-register β-sheet structure for PrPSc.

Our studies afford insight into the mechanism of prion conversion. Asn/Gln substitutions at positions flanking Asn143 or Gln219 (positions 142, 144, 145, 218, 220, and 221) had little differences result in microcrystals having different symmetry (14, 36). Thus, it is not entirely surprising that residue differences in these particular zipper segments had such a remarkable impact on prion transmission barriers. The surprising element here was that the residues that most profoundly impacted cross-species prion conversion were largely dominated by Asn/Gln residues.

The critical Asn/Gln residue positions for cross-species conversion differed depending on the PrPSc seed (Fig. 5), consistent with the recognized role of PrPSc conformation in prion conversion. The results using mutant PrP have enabled a partial mapping of interacting residues critical for certain prions, suggesting segments that may be exposed in PrPSc. PrPSc–PrPSc interaction at residues 215 and 219 was important for RML only, indicating that the distal C terminus is a key intermolecular contact segment for RML, but not 87V or CWD prions. Additionally, the β2–α2 loop was key for CWD-seeded conversion but insufficient for RML or 87V. Findings from our lab and others that the key interaction domains in PrPSc vary with the PrPSc conformation are consistent with those from Sup35 yeast prions, showing that short segments control prion nucleation, conformation, and species barriers (43).

How do asparagine and glutamine residues lower the energy barrier for prion cross-seeding? Of relevance to Huntington’s disease, fibril assembly of the huntingtin protein is caused by expansion of poly(Q) tracts (44, 45). Perutz et al. (46) have proposed that Asn/Gln residues mediate protein-protein interactions in huntingtin and other Asn/Gln-rich aggregating proteins through the formation of hydrogen bond networks that stabilize β-strands, termed “polar zippers.” Within a β-sheet, asparagine and glutamine residues were proposed to markedly enhance β-strand stability through intrasheet hydrogen bonds formed between both side chain and main chain amides (Fig. 5).
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Effect on conversion of hHuPrP<sup>C</sup>. This finding indicates that the precise Asn/Gln position within the segment was critical for prion conversion, which is consistent with steric zipper formation at segment 138–143. For instance, Gly<sup>142</sup> is located at a tight turn in the crystal structure of the 138–143 segment (36). Asn/Gln substitutions at this position would be predicted to clash with zipper formation and thus hinder prion conversion. Additionally, other substitutions in hHuPrP<sup>C</sup> that increase total Asn/Gln content, such as S97N or H155N, did not strongly promote conversion, supporting the importance of the specific position of the Asn/Gln residue in facilitating prion conversion.

We were asparagine and glutamine residues in the key positions interchangeable? PrP positions 100 and 143 required an asparagine and not a glutamine for conversion, indicating that a specific side chain length was crucial in certain positions (Fig. 5). On the other hand, positions 170 and 219 tolerated either an asparagine or glutamine; however, leucine substitutions at position 143 or 219 blocked conversion. Together, these findings underscore the importance of the hydrogen bond stabilizing ladder provided by the asparagine or glutamine side chain. In Sup35 yeast prions, asparagine residues enhance prion formation and may promote conversion, supporting the importance of the specific Asn/Gln content, such as S97N or H155N, did not strongly stabilize nascent PrP<sup>Sc</sup>.

The number of key interspersed Asn/Gln residues (Table 2) may be involved early in PrP nucleation and is favored by asparagine residues (59, 60). Our studies suggest that the high number of key interspersed Asn/Gln residues (Table 2) may stabilize nascent β-strands through a hydrogen bond ladder and may explain the elevated promiscuity of bank vole to cross-species prion conversion, as well as the spontaneous assembly of BvPrP<sup>C</sup> in transgenic mice (61).

These findings argue that certain interspersed asparagine and glutamine residues may facilitate the anchoring of PrP<sup>C</sup> to PrP<sup>Sc</sup> through strengthening a replicative interface, driving prion conversion between dissimilar sequences and lowering the barrier for aggregation. Identifying the key prion segments and specific residue interactions in early stages of conversion may facilitate predictions of cross-species prion transmission. We expect that the importance of interspersed asparagine and glutamine residues in protein aggregation may be a more general phenomena applicable to other amyloidogenic proteins and may present target segments for rational therapeutic design.

Experimental procedures

Cell-lysate protein misfolding cyclic amplification

The pcDNA3.1 vector (Invitrogen) with the mouse, human, bank vole, or rabbit Prnp encoding the 3F4 epitope (Met<sup>109</sup> and Met<sup>112</sup> human numbering) was used as a template for site-directed mutagenesis (QuikChange site-directed mutagenesis kit; Agilent). PrP-deficient RK13 cells (ATCC) were transfected with 5–10 μg of plasmid DNA using Lipofectamine 3000 (Invitrogen). At 24 h post-transfection, the cells were washed twice in PBS, harvested in 1 ml of PBS, and centrifuged for 1 min at 1,000 × g. The pellet was resuspended in PMCA buffer (PBS containing 1% Triton X-100, 150 mM NaCl, and 5 mM EDTA plus Complete<sup>TM</sup> protease inhibitors), passed repeatedly through a 27-gauge needle, and clarified by centrifuging at 2,000 × g for 1 min.

RML prions from C57BL/6 mice (Prnp encoding Leu<sup>109</sup>, Thr<sup>189</sup>, human numbering), 87V prions from VM/DK mice (Prnp encoding Phe<sup>109</sup> and Val<sup>189</sup>), and CWD prions from Elk (Met<sup>129</sup>) were used to seed human, bank vole, and rabbit PrP<sup>C</sup> (Prnp sequences shown in Fig. 1 and supplemental Fig. S1). The PrP<sup>C</sup> was newly prepared for each independent experiment. The prion seeds were derived from brain homogenate that was pooled from mice inoculated with the same prion strain or from naturally infected elk. The brain homogenate samples pooled to generate the seeds were consistent between the experiments. Prion-infected brain homogenate (10% w/v) was added into PrP<sup>C</sup>-expressing RK13 cell lysate (1:10, PrP<sup>Sc</sup>:PrP<sup>C</sup> by volume) and subjected to repeated 5-s sonications (S4000, Qsonica) with 10 min of incubation between each pulse over a total period of 24 h. Sonication power was maintained at 50–60%, and samples were continuously rotated in a water bath at 37 °C. Samples were then digested with 200 μg/ml PK for 30 min at 37 °C and analyzed by Western blot using the anti-PrP monoclonal antibody 3F4 (62). PrP<sup>C</sup> levels were measured by blotting 1–2 μl from unseeded lysates. Signals were quantified using a Fujifilm LAS-4000 imager and multi-gauge kit; Agilent).

Statistical analysis

One-way ANOVA with Tukey post hoc test and a one-sample t test were used to analyze the conversion data from each mutant PrP. For the one-sample t test, the conversion of human or rabbit substitutions in bank vole PrP<sup>C</sup> was compared against a mean of 100 (null hypothesis) to assess whether the mutation(s) inhibited PrP conversion. The conversion of bank vole substitutions in human or rabbit PrP<sup>C</sup> was compared against a mean of 0 to assess whether the mutation(s) promoted PrP conversion.

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