DOI: 10.1021/bi101449f



Diverse Effects on the Native β -Sheet of the Human Prion Protein Due to Disease-Associated Mutations[†]

Wei Chen, Marc W. van der Kamp, and Valerie Daggett*

Department of Bioengineering, University of Washington, Seattle, Washington 98195-5013, United States

Received September 8, 2010; Revised Manuscript Received October 14, 2010

ABSTRACT: Prion diseases are fatal neurodegenerative disorders that involve the conversion of the normal cellular form of the prion protein (PrP^C) to a misfolded pathogenic form (PrP^{Sc}) . There are many genetic mutations of PrP associated with human prion diseases. Three of these point mutations are located at the first strand of the native β -sheet in human PrP: G131V, S132I, and A133V. To understand the underlying structural and dynamic effects of these disease-causing mutations on the human PrP, we performed molecular dynamics of wild-type and mutated human PrP. The results indicate that the mutations induced different effects but they were all related to misfolding of the native β -sheet: G131V caused the elongation of the native β -sheet, A133V disrupted the native β -sheet, and S132I converted the native β -sheet to an α -sheet. The observed changes were due to the reorientation of side chain—side chain interactions upon introducing the mutations. In addition, all mutations impaired a structurally conserved water site at the native β -sheet. Our work suggests various misfolding pathways for human PrP in response to mutation.

Prion diseases, or transmissible spongiform encephalopathies (TSEs), are fatal neurodegenerative disorders, including Creutzfeldt-Jakob disease (CJD), Gerstmann-Straüssler-Scheinker syndrome (GSS), fatal familial insomnia (FFI), and kuru in humans, scrapie in sheep, bovine spongiform encephalopathy (BSE) in cattle, and chronic wasting disease (CWD) in cervids (1, 2). Prion diseases are attributed to the conformational conversion of the normal cellular prion protein (PrP^C) to its misfolded "scrapie" form, PrP^{Sc} (2, 3). PrP^C is a glycoprotein anchored to the cell membrane via a glycophosphatidylinositol (GPI) moiety (2, 4). PrP^C contains a flexible and largely unstructured N-terminal region and a C-terminal globular domain (Figure 1). The C-terminal domain consists of three α -helices (HA, HB, and HC) and a short two-stranded antiparallel β -sheet (S1 and S2). While PrP^{C} is mainly α -helical and is sensitive to protease digestion, PrP^{Sc} is enriched in β -sheet structure and is proteinase K-resistant (5, 6). The precise structure of PrP^{Sc}, however, is still unknown, and in fact it appears to be quite heterogeneous.

Human prion diseases can be infectious, sporadic, or genetic in origin (1, 2). Many single nucleotide mutations in the human prion protein gene (PRNP) have been linked with human prion diseases: familial CJD, GSS, and FFI (7, 8). Three mutations that are related to GSS, G131V (9), S132I (10), and A133V (11), are located at the first strand of the native β -sheet, S1 (Figure 1). The pathological mechanism as to how these mutations cause GSS is

unknown. To reveal the underlying structural ramifications upon mutation, one crucial step is to study how these mutations affect and convert PrP^C to PrP^{Sc} .

To this end, we have performed molecular dynamics (MD) simulations of human PrP^{C} structures containing residues 90-230 with the three mutations mentioned above at neutral pH and physiological temperature (310 K). Our simulations suggest that these mutations induced different conformational changes in human PrP^{C} , although they are all located at S1. With G131V, two additional hydrogen bonds formed between the main chains of strands S1 and S2, resulting in the extension of the native β -sheet. In contrast, A133V caused the loss of hydrogen bonds and disrupted the native β -sheet. By introducing S132I, hydrogen bonds of the native β -sheet were reordered. As a result, the native β -sheet was converted to an α -sheet (12). Reorganization

FIGURE 1: Structure of wt human PrP^C. The starting structure used for MD simulations, residues 90–230. Secondary structure elements in the C-terminal globular domain are labeled. The three residues involved in the point mutations studied in this work are shown as

 $^{^{\}dagger}$ We are grateful for support provided by the National Institutes of Health (GM81407 to V.D.).

^{*}To whom correspondence should be addressed. Phone: (206) 685-7420. Fax: (206) 685-3300. E-mail: daggett@u.washington.edu.

Abbreviations: PrP, prion protein; PrPC, cellular prion protein;

¹Abbreviations: PrP, prion protein; PrP^C, cellular prion protein; PrP^{Sc}, "scrapie" prion protein; TSE, transmissible spongiform encephalopathy; CJD, Creutzfeldt—Jakob disease; GSS, Gerstmann—Straüssler—Scheinker syndrome; FFI, fatal familial insomnia; BSE, bovine spongiform encephalopathy; CWD, chronic wasting disease; GPI, glycophosphatidylinositol; MD, molecular dynamics; wt, wild type; *il*mm, *in lucem* molecular mechanics; rmsd, root-mean-square deviation; NMR, nuclear magnetic resonance.

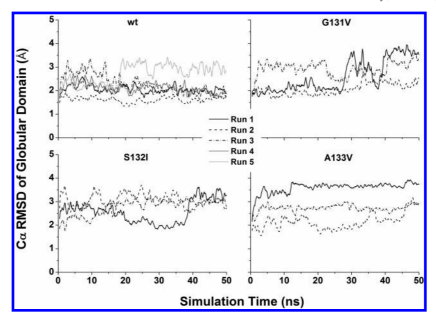


FIGURE 2: $C\alpha$ rmsd of the globular domain in MD simulations. Structures were aligned on the $C\alpha$ atoms of the globular domain (residues 128-228) before rmsd measurements. The results of both wt and mutants are shown. For clarity, a running average over 500 ps is shown.

of side chain—side chain interactions at the native β -sheet and the neighboring part of the N-terminal region was responsible for the observed conformational changes. Furthermore, the three mutations impaired a structurally conserved water site connecting S1, S2, and HC. In particular, the G131V mutation fully abolished the water site. These results suggest that there are various pathways for the early misfolding of PrP^C due to mutations.

METHODS

Structure Preparation. The starting structure of wild-type (wt) human PrP^{C} for MD simulations was modeled based on a NMR structure of human PrP^{C} (PDB code 1QLX) (13), which contains residues 125–228. Because the major proteinase K-resistant fragment of PrP^{Sc} (i.e., PrP 27–30) consists of residues $\sim 90-230$ (14), we modeled the N-terminal residues 90-127 and the C-terminal residues 229-230 as described previously (15). Consequently, the starting structure for the MD simulations of wt human PrP^{C} contained residues 90-230. As before, His140 was protonated on $N_{\delta 1}$, and other histidines were protonated on $N_{\epsilon 2}$ (15).

To model the starting structures of mutants G131V, S132I, and A133V, for which experimental structures do not exist, we mutated the corresponding residues *in silico*. First, the side chain atoms of the mutated residue, except for C_{β} , were deleted. Then, different rotamers of the target residue type were obtained from the Dynameomics rotamer library (16, 17) and added to generate multiple structures; after 100 steps of steepest descent minimization on all side chains, the structure with the smallest interaction energy between the side chain of the mutated residue and the surrounding residues was selected as the starting structure for MD simulations.

MD Simulations. Explicit solvent all-atom MD simulations were performed with in lucem molecular mechanics (ilmm) (18). The Levitt et al. force field (19) and the F3C model (20) were used for protein and water, respectively. Electrostatic interactions between atoms separated by three bonds or less were scaled by 0.4. The force-shifted nonbonded cutoff (21) was set to 10 Å, and the nonbonded list was updated every two steps. A time step of 2 fs was used.

The starting structures were first energy-minimized for 1000 steps with the steepest descent method in vacuo. Then, they were solvated in preequilibrated water boxes at 310 K. The final water density of the system was adjusted to the experimentally measured density of 0.993 g/mL (22). The minimal distance between the protein and the walls of the box was 10 A. To preclude bad interactions during solvation, water molecule positions were adjusted by 1000 steps of steepest descent minimization, 500 steps of MD, and another 500 steps of minimization. After that, 500 steps of steepest descent minimization were performed for the proteins. The production MD simulation of the whole system was then performed in the microcanonical (NVE, constant number of particles, volume, and energy) ensemble for 50 ns. System coordinates were saved every 1 ps. Five and three independent MD simulations were performed for wt PrP^C and each mutant, respectively, using different random number seeds to generate independent trajectories, for a total of 700 ns, or $0.7 \mu s$.

Analysis. Analysis of MD trajectories was performed with *il*mm (18) and VMD (23). Root-mean-square deviation (rmsd) of the C α atoms of the C-terminal globular domain (residues 128–228) and HA (residues 144–156) were calculated by aligning structures on the C α atoms of the globular domain and a stable core region (residues 174–186 and 200–219) (15, 24), respectively. A hydrogen bond was defined if the hydrogen—acceptor distance was \leq 2.6 Å and the donor—hydrogen—acceptor angle was \geq 135°. Hydrophobic contacts were identified when the C–C distance between CH_n groups was \leq 5.4 Å. The DSSP algorithm (25) with additional definitions (26) was used to assign secondary structures. Molecular graphics were generated using VMD (23).

RESULTS

HA Shifts Position in both wt and Mutant Simulations. The three disease-related mutations (G131V, S132I, and A133V) are located at the S1 strand (Figure 1). Three independent simulations at neutral pH and 310 K were performed for each mutant. For comparison, five simulations at neutral pH and 310 K were performed for the wt PrP structure. In some simulations, the rmsd of all $C\alpha$ atoms of the globular domain (residues 128–228) relative to the starting structure was \sim 2 Å through the

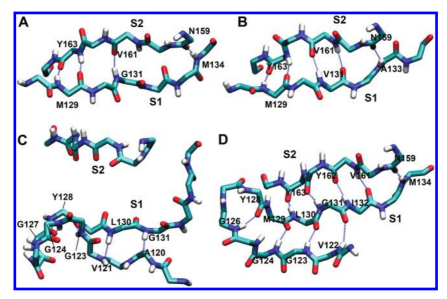


FIGURE 3: Structures of the native β -sheet. (A) The starting native sheet. (B) The elongated native sheet at 49 ns of run 1 of PrP mutant G131V. (C) The disrupted native sheet at 50 ns of run 2 of PrP mutant A133V. (D) The converted native sheet to an α -sheet at 50 ns of run 2 of PrP mutant S132I. Only backbone atoms are shown. Dashed lines represent hydrogen bonds. Residues participating in hydrogen bonds are labeled.

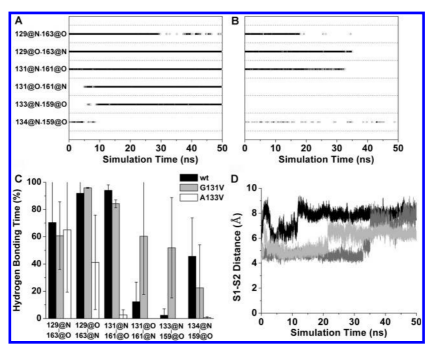


FIGURE 4: Changes in the native sheet upon G131V and A133V mutations. Main chain hydrogen bonds of the native sheet are plotted vs simulation time for (A) run 1 of PrP mutant G131V and (B) run 2 of PrP mutant A133V. A symbol represents that the hydrogen bond indicated by a y label exists at the time indicated by an x value. (C) Percentage of hydrogen-bonding times in the period of 25–50 ns averaged over all runs. x labels indicate the hydrogen bonds. Error bars show the standard deviation. (D) The distances between the centers of masses of the backbone atoms of the S1 and S2 strands during MD simulations of PrP mutant A133V. The results of run 1 (black), run 2 (dark gray), and run 3 (light gray) are shown.

simulations (Figure 2), indicating that the globular domain was stable. However, in other cases the Cα rmsd of the globular domain jumped to 3–4 Å, manifesting conformational changes. This is attributed to the movements of HA as indicated by the similar behavior of the rmsd of HA alone when structures were aligned on the stable core region (residues 174–186 and 200–219) (24) (Supporting Information Figure S1). Different types of HA movements were observed (Supporting Information Figure S2). HA moved away from the HB/HC plane (e.g., G131V run 1), moved downward (e.g., wt run 5), or moved toward the center (e.g., A133V run 1). By measuring the moving vectors of the HA N-terminus (residues 144–147) relative to its starting

position as described previously (27), we quantified the HA movements along different directions (Supporting Information Figure S3). The biggest outward movements of HA occurred in G131V run 1 and S132I run 1 while the largest downward movements were observed in wt run 5 and G131V run 2. Because HA movements occurred in both wt and mutant simulations, this structural flexibility might be an intrinsic property of human PrP at 310 K.

The G131V Mutation Induces the Elongation of the Native β -Sheet. In the starting PrP structure, the short antiparallel native β -sheet contains a β -bulge (28) and consists of four main chain hydrogen bonds: M129:N-Y163:O, M129:O-Y163:N,

G131:N-V161O, and M134:N-N159:O (Figure 3A). Upon mutation to G131V, two new hydrogen bonds (V131:O-V161:N and A133:N-N159:O) formed while one of the original hydrogen bonds (M134:N-N159:O) broke, which resulted in the elongation of the native β -sheet and the elimination of the β -bulge (Figures 3B) and 4A). In two out of three simulations of G131V, we observed the stable elongation of the native β -sheet (Supporting Information Figure S4). By comparison, in the wt simulations, the two new hydrogen bonds formed sporadically, but they were unstable (Supporting Information Figure S4). The hydrogen bond occupancies over the last 25 ns of the simulations were calculated for the six hydrogen bonds in the native β -sheet (Figure 4C). The two new hydrogen bonds were intact significantly longer in G131V

The A133V Mutation Disrupts the Native β -Sheet. In contrast to the G131V mutation, introduction of A133V resulted in partial or full disruption of the native β -sheet hydrogen bonding and a shift with S1 interacting with the neighboring part of the N-terminal region (Figure 3C). In all three simulations at least two of the original hydrogen bonds broke (Supporting Information Figure S4), and all of them broke in run 2 (Figure 4B). Except for the M129:N-Y163:O hydrogen bond, the hydrogen-bonding times of the other three original hydrogen bonds were much shorter for A133V than wt, indicating the disruptions of these hydrogen bonds due to the A133V mutation (Figure 4C). In conjunction with the loss of hydrogen bonds, S1 and S2 separated (Figure 4D). In different simulations, the separation of S1 and S2 occurred at different times and to different degrees. In run 1, the separation occurred at the start of the simulation, and the distance between the centers of mass of the backbone atoms of S1 and S2 increased to \sim 8 Å. In run 2, the separation occurred at \sim 33 ns. The distance between S1 and S2 first increased to \sim 5 Å for a short period of time and then to \sim 8 Å. In run 3, S1 was separated from S2 at \sim 22 ns and stayed at a position ~ 6.5 Å away from S2.

The S132I Mutation Causes α-Sheet Formation. In the first simulation of mutation S132I, the M129/L130 and Y162/ Y163 peptide planes "flipped" at ~41 ns, allowing for formation of the L130:N-Y162:O hydrogen bond while the original M129: O-Y163:N hydrogen bond was lost (Supporting Information Figure S4). The resulting sheet is now in an α -sheet conformation (12, 29). In run 2, even more significant changes were observed (Figure 3D). Besides the flipping of the peptide planes, hydrogen bonding between S1 and S2 shifted by one residue: G131:N now formed an hydrogen bond with Y162:O (instead of V161:O) (Figure 5A). Two new hydrogen bonds, I132:N-V161: O and I132:O-V161:N, also formed. Together, these changes resulted in a mixed α/β -sheet, with the α -sheet on one side (residues 130–132 and 161–163) and the β -sheet on the other (residues 132–134 and 159–161). This conformational change occurred at the very beginning of the simulation (Figure 5). Around 10 ns, an additional α-strand (residues 122–126) formed in the N-terminal region (Figures 3D and 5), yielding a threestranded mixed α/β -sheet. However, in run 3 such changes were not observed (Supporting Information Figure S4).

Reorientation of Side Chain-Side Chain Interactions Is Responsible for Changes in the Native β -Sheet. To understand why the mutations had different effects on the native β -sheet, we investigated the side chain—side chain interactions in detail. With the G131V mutation, the bulky side chain of V131 tended to push the side chain of S132 away to the other side of the β -sheet (Figure 6A). This repositioning allowed the side chain

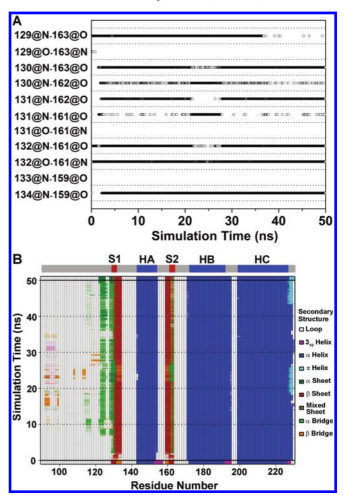


FIGURE 5: Changes in the native sheet upon S132I mutation. (A) Time course of main chain hydrogen bonds of the native sheet in run 2 of PrP mutant S132I. (B) Secondary structure analysis of PrP in run 2 of PrP mutant S132I. The secondary structure elements present in the starting structure are shown on the top.

oxygen (O_y) of S132 to form an hydrogen bond with the side chain nitrogen ($N_{\epsilon 2}$) of Q160 (Figure 6B). As a result, the original hydrogen bond between Q160:N_{E2} and V131:O was disrupted, which in turn allowed V131:O to hydrogen bond with V161:N. The local conformational changes further propagated down to A133 to allow the formation of the A133:N-N159:O hydrogen bond. Ultimately, the native β -sheet was elongated by two more hydrogen bonds. In run 1 of G131V, we even saw the reorientation of the side chains of A133 and M134 (Figure 6A). Initially, the A133 side chain pointed up while the M134 side chain pointed down. During the simulation, the side chains changed their orientations so that the A133 side chain pointed down while the M134 side chain pointed up. This is reasonable because the favored configuration of a β -sheet is that side chains are alternating above and below the sheet.

Changes in side chain-side chain interactions were also observed in the S132I mutant. Because the I132 side chain is bulky and hydrophobic, it interacted with the hydrophobic side chain of Y163, bringing them closer together. This movement caused a shift of hydrogen bonding between S1 and S2 by one residue, as mentioned above. This shift brought the L130 side chain closer to Y162. As a result, the L130 side chain turned downward (Figure 6C). These changes are illustrated by the increase in distance between the L130 and Y162 side chains and the decrease in distance between the L130 and Y163 side chains

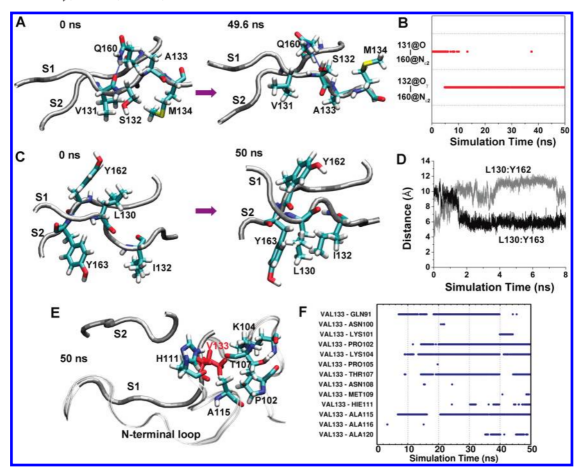


FIGURE 6: Reconfiguration of side chain interactions of the native sheet due to mutations. (A) Conformations of strands S1 and S2 at the indicated times in run 1 of PrP mutant G131V. The backbones of the strands are shown as tubes, and relevant residues are represented as sticks and labeled, as is the case in panels C and E. (B) Time course of hydrogen bonds between S131:O and Q160:N $_{e2}$ and between S132:O $_{\gamma}$ and Q160:N $_{e2}$ in run 1 of PrP mutant G131V. (C) Conformations of strands S1 and S2 at indicated times in run 2 of PrP mutant S132I. (D) The distance between the centers of mass of the L130 side chain and the Y162 (or Y163) side chain in run 2 of PrP mutant S132I. (E) The structure of the S1 and S2 strands and N-terminal region at 50 ns in run 2 of PrP mutant A133V. V133 is in red. The backbone of the N-terminal region is shown as a ribbon. (F) Time course of hydrophobic side chain—side chain contacts between V133 and residues of the N-terminal region in run 2 of PrP mutant A133V.

(Figure 6D). The shift between S1 and S2 also eliminated the β -bulge at S132 and A133, eventually resulting in the elongation of the native β -sheet.

A different mechanism was observed for the A133V mutation. V133 formed hydrophobic interactions with residues 91–122 in the flexible N-terminal region for significant periods during the simulations (Figure 6E,F and Supporting Information Figure S5), presumably due to the solvent-exposed starting position of its hydrophobic side chain. In comparison, A133 in the wt simulations seldom interacted with the residues of the N-terminal region. The new hydrophobic interactions brought the neighboring N-terminal sequence (residues 117–125) closer to S1, which led to formation of both hydrogen bonds and hydrophobic interactions between S1 and neighboring N-terminal residues (Figure 3C). Altogether, these interactions disrupted the hydrogen bonds between the backbones of S1 and S2, ultimately leading to their separation.

Mutations Affect a Conserved Water Site in PrP^{C} . A tightly bound, structurally conserved water in the native β-sheet of PrP^{C} was identified by the crystal structures of the sheep PrP^{C} (PDB 1UW3) (30) and the human PrP^{C} (PDB 1I4M and 3HAK) (31, 32) and MD simulations (33). The water forms hydrogen bonds with the carbonyl oxygen of S132, the amide nitrogen of V161, and the side chain oxygen of Q217 (O_{ε1}), which connects three PrP^{C} regions (S1, S2, and HC) (Figure 7A). In our

simulations of wt human PrP^{C} , we also observed that water molecules entered the water site and stayed for long or short periods of time (Supporting Information Figure S6). Besides the hydrogen-bonding pattern predicted from the crystal structures, a different pattern also occurred in the simulations: the water molecule interacted with the carbonyl oxygen of S132, the amide nitrogen of V161, the carbonyl oxygen of N159, and the side chain nitrogen of Q217 ($N_{\rm f2}$) (Figure 7B).

To investigate how the mutations would affect the conserved water site, we analyzed water occupancy in the site over the simulations. We defined a water molecule as a coordinating water if it simultaneously connected S1, S2, and HC; i.e., it formed hydrogen bonds with S132:O, N159:O or V161:N, and Q217:O_{E1} or Q217:N_{e2} at the same time. In all simulations of wt PrP, coordinating waters were observed (Supporting Information Figure S6). Particularly in run 4, a water molecule was locked in the site throughout the simulation. However, waters were in exchange with bulk water in runs 1 and 3. In contrast to the wt simulations, no coordinating waters were observed in the site in the G131V simulations, indicating that the water site was abolished due to the elongation of the native β -sheet. For the S132I mutant, several water molecules moved in and out of the site sequentially in run 1. But no coordinating waters were found in run 2 because of the shift and elongation of the native β -sheet, as described above. In run 3, only at the very beginning was

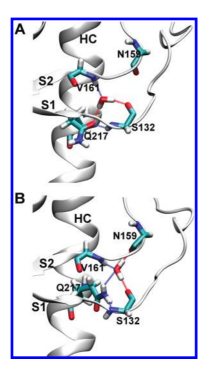


FIGURE 7: The conserved water site at the native β -sheet. (A) One mode of water coordination at the water site as observed in both the crystal structures of PrP and MD simulations of wt human PrP. (B) Another mode of water coordination at the water site as observed in MD simulations of wt human PrP. Coordinating residues and water are shown as sticks. Dashed lines indicate hydrogen bonds.

coordination water present because the bulky side chain of I132 pushed the side chain of Q217 aside. For the A133V mutant, coordinating waters were present in the first halves of runs 2 and 3 but were displaced half-way through the simulation, and almost no coordinating waters existed in run 1. The disappearance of the coordinating waters was well correlated with the separation of S1 and S2 (Figure 4D). In run 1, the two strands separated from the beginning, so no coordinating waters were possible. In runs 2 and 3, the separation occurred at \sim 22 and \sim 33 ns, respectively. As a result, no coordinating waters were found in the second halves of the simulations. Altogether, the three mutations at S1 impaired the water site although to different degrees.

DISCUSSION

Many single nucleotide mutations in *PRNP* have been identified to be associated with human prion diseases (7, 8), but the underlying mechanism of pathogenesis is still unclear. Here we describe MD simulations of human PrP with point mutations located on or near the S1 strand that are implicated in the development of GSS: G131V (9), S132I (10), and A133V (11). The simulations showed that these three mutations induced different effects on the native β -sheet: G131V elongated the native β -sheet from three to five hydrogen bonds, A133V disrupted the native β -sheet, and S132I converted the native β -sheet to an α -sheet. The various effects caused by these mutations suggest that PrP mutants can undergo diverse misfolding pathways and eventually may form different structures of PrP^{Sc} that manifest as different prion strains.

Previously, Santini et al. also simulated the G131V mutant although their simulations are only 4 ns long, much shorter than ours (34). However, Santini et al. did not observe the elongation of the native β -sheet from three to five hydrogen bonds for the G131V mutant. The elongation of the native β -sheet eliminated

the β -bulge at S132 and A133 (Figure 3B), which may facilitate interactions of S1 with the N-terminal region and thus the growth of the native β -sheet from two strands to three or more strands, as observed in simulations under acidic conditions (35, 36).

In contrast to the G131V mutation, A133V caused separation of the S1 and S2 strands. We have not seen any reports of the effects of this mutation on the PrP structures. However, another mutation, T183A, has been shown by coarse-grained MD simulations to disrupt the native β -sheet (37). Therefore, other than the growth of the native sheet, the disruption of the native β -sheet is probably another early misfolding event.

Interestingly, the S132I mutation converted the native β -sheet to an α-sheet and further extended to three strands with participation of the N-terminal region (Figure 3D). In the α sheet, the main chain amide groups are on one side and carbonyl groups are on the other side, forming a polar sheet. This alignment has been suggested to give rise to attractive forces for self-assembly and thus facilitate formation of amyloid protofibrils (29). In the early stage of the unfolding of bovine PrP^C and simulations of human and hamster PrP^C, an α-sheet was also observed (29, 36, 38). The fact that the formation of α -sheets occurs in the simulations of numerous amyloidogenic proteins under amyloidogenic conditions (low pH or upon mutation, typically) and also in our simulations of S132I suggests that it is a common mechanism for amyloidogenesis (12, 29, 36, 38–43).

A structurally conserved water site was identified in the β -bugle of the native β -sheet in several crystal structures (Figure 7) (30-33). Water in this site connects S1, S2, and HC and has been suggested to stabilize the structure (33). Indeed, in the simulations of sheep PrP by De Simone et al., a water molecule was locked in the site once it entered (33). By comparison, in our wt simulations, which are much longer than the simulations of De Simone et al. (50 vs 10 ns), we observed exchange of waters in the site with bulk water, except that in run 4 a water was locked in the site throughout the simulation. The three mutations in the native β -sheet all impaired the water site, either by eliminating the β -bulge at S132 and A133 or by separation of the S1 and S2 strands.

The predictions of the structural changes of the native β -sheet based on the MD simulations could be tested by experiments. One possible way is to measure hydrogen bond scalar couplings $^{(h3}J_{\rm NC'})$ for the native β -sheet by nuclear magnetic resonance (NMR) (44). $^{h3}J_{\rm NC'}$ depends strongly on the geometry of hydrogen bonds (45, 46). We calculated $^{h3}J_{\rm NC'}$ based on structures from the last 25 ns of the MD trajectories. As expected, the changes of h3 J_{NC} due to mutations were observed corresponding to the formation or breakage of hydrogen bonds in the native β sheet (Supporting Information Figure S7). Compared with wt, the G131V mutant increases the magnitude of $^{h3}J_{NC'}$ of the V131: O-V161:N and A133:N-N159:O hydrogen bonds while the A133V mutant reduces $^{h3}J_{NC'}$ for several hydrogen bonds: M129: O-Y163:N, G131:N-V161:O, and M134:N-N159:O. On the other hand, the S132I mutant should yield new couplings. In addition, mutagenesis experiments could be used to test our predictions. A hydrogen bond between Q160:N_{ϵ 2} and G131:O exists in the wt PrP structure. In G131V this hydrogen bond was broken when the V131:O formed a hydrogen bond with V161:N (Figure 6A), suggesting that these two hydrogen bonds compete with each other. Therefore, if Q160 is mutated to another residue such as alanine, G131:O should be able to form a hydrogen bond with V161:N, resulting in the elongation of the native β -sheet.

The simulations described here suggest that multiple early misfolding pathways for PrP exist in response to mutation: one is the growth of a β -sheet via the elongation of the native β -sheet; the second is the disruption of the native β -sheet; the third is the growth of an α -sheet via the conversion of the native β -sheet. In contrast, we have found more consensus among misfolding events in the wt PrP at low pH even among different species (15, 26, 35, 36, 38). The simulations presented here shed further light on the mechanisms of the conversion of PrP^C to PrP^{SC}, particularly variations due to mutation.

ACKNOWLEDGMENT

We are grateful for computational resources provided by Microsoft. We thank Alex D. Scouras for technical assistance.

SUPPORTING INFORMATION AVAILABLE

Cα rmsd of HA in MD simulations (Figure S1), HA movements in MD simulations (Figure S2), movements of the HA N-terminus (Figure S3), time course of main chain hydrogen bonds between the S1 and S2 strands in MD simulations (Figure S4), time course of hydrophobic side chain—side chain contacts between V133 and residues of the N-terminal region (Figure S5), time course of water molecules residing in the conserved water site at the native β -sheet (Figure S6), and predicted hydrogen bond scalar couplings ($^{h3}J_{NC}$) for hydrogen bonds between S1 and S2 (Figure S7). This material is available free of charge via the Internet at http://pubs.acs.org.

REFERENCES

- 1. Aguzzi, A., Baumann, F., and Bremer, J. (2008) The prion's elusive reason for being. *Annu. Rev. Neurosci.* 31, 439–477.
- Aguzzi, A., and Calella, A. M. (2009) Prions: protein aggregation and infectious diseases. *Physiol. Rev.* 89, 1105–1152.
- Caughey, B., Baron, G. S., Chesebro, B., and Jeffrey, M. (2009) Getting a grip on prions: oligomers, amyloids, and pathological membrane interactions. *Annu. Rev. Biochem.* 78, 177–204.
- Linden, R., Martins, V. R., Prado, M. A., Cammarota, M., Izquierdo, I., and Brentani, R. R. (2008) Physiology of the prion protein. *Physiol. Rev.* 88, 673–728.
- 5. Caughey, B. W., Dong, A., Bhat, K. S., Ernst, D., Hayes, S. F., and Caughey, W. S. (1991) Secondary structure analysis of the scrapic-associated protein PrP 27–30 in water by infrared spectroscopy. *Biochemistry* 30, 7672–7680.
- Pan, K. M., Baldwin, M., Nguyen, J., Gasset, M., Serban, A., Groth, D., Mehlhorn, I., Huang, Z., Fletterick, R. J., and Cohen, F. E.; et al. (1993) Conversion of α-helices into β-sheets features in the formation of the scrapie prion proteins. *Proc. Natl. Acad. Sci. U.S.A.* 90, 10962–10966.
- Mead, S. (2006) Prion disease genetics. Eur. J. Hum. Genet. 14, 273–281.
- 8. Van der Kamp, M. W., and Daggett, V. (2009) The consequences of pathogenic mutations to the human prion protein. *Protein Eng.*, *Des. Sel. 22*, 461–468.
- Panegyres, P. K., Toufexis, K., Kakulas, B. A., Cernevakova, L., Brown, P., Ghetti, B., Piccardo, P., and Dlouhy, S. R. (2001) A new PRNP mutation (G131V) associated with Gerstmann-Straussler-Scheinker disease. *Arch. Neurol.* 58, 1899–1902.
- Hilton, D. A., Head, M. W., Singh, V. K., Bishop, M., and Ironside, J. W. (2009) Familial prion disease with a novel serine to isoleucine mutation at codon 132 of prion protein gene (PRNP). *Neuropathol. Appl. Neurobiol.* 35, 111–115.
- Rowe, D. B., Lewis, V., Needham, M., Rodriguez, M., Boyd, A., McLean, C., Roberts, H., Masters, C. L., and Collins, S. J. (2007) Novel prion protein gene mutation presenting with subacute PSP-like syndrome. *Neurology* 68, 868–870.
- Daggett, V. (2006) α-sheet: the toxic conformer in amyloid diseases?
 Acc. Chem. Res. 39, 594–602.
- 13. Zahn, R., Liu, A., Luhrs, T., Riek, R., von Schroetter, C., Lopez Garcia, F., Billeter, M., Calzolai, L., Wider, G., and Wuthrich, K.

- (2000) NMR solution structure of the human prion protein. *Proc. Natl. Acad. Sci. U.S.A.* 97, 145–150.
- Prusiner, S. B., Groth, D. F., Bolton, D. C., Kent, S. B., and Hood, L. E. (1984) Purification and structural studies of a major scrapie prion protein. *Cell* 38, 127–134.
- 15. Van der Kamp, M. W., and Daggett, V. (2010) The influence of pH on the human prion protein: Insights into the early steps of misfolding. *Biophys. J.* 99, 2289–2298.
- Van der Kamp, M. W., Schaeffer, R. D., Jonsson, A. L., Scouras, A. D., Simms, A. M., Toofanny, R. D., Benson, N. C., Anderson, P. C., Merkley, E. D., Rysavy, S., Bromley, D., Beck, D. A., and Daggett, V. (2010) Dynameomics: a comprehensive database of protein dynamics. Structure 18, 423–435.
- Scouras, A. D., and Daggett, V. (2010) The dynameomics rotamer library: amino acid side chain conformations and dynamics from comprehensive molecular dynamics simulations in water, *Protein Sci.*, submitted.
- Beck, D. A. C., Alonso, D. O. V., and Daggett, V. (2000–2010) in Lucem Molecular Mechanics (ilmm), University of Washington, Seattle, WA.
- Levitt, M., Hirshberg, M., Sharon, R., and Daggett, V. (1995) Potential-energy function and parameters for simulations of the molecular-dynamics of proteins and nucleic-acids in solution. *Comput. Phys. Commun.* 91, 215–231.
- Levitt, M., Hirshberg, M., Sharon, R., Laidig, K. E., and Daggett, V. (1997) Calibration and testing of a water model for simulation of the molecular dynamics of proteins and nucleic acids in solution. *J. Phys. Chem. B* 101, 5051–5061.
- Beck, D. A. C., and Daggett, V. (2004) Methods for molecular dynamics simulations of protein folding/unfolding in solution. *Methods* 34, 112–120.
- 22. Kell, G. S. (1967) Precise representation of volume properties of water at 1 atm. *J. Chem. Eng. Data* 12, 66–69.
- 23. Humphrey, W., Dalke, A., and Schulten, K. (1996) VMD: visual molecular dynamics. *J. Mol. Graphics* 14 (33–38), 27–38.
- Viles, J. H., Donne, D., Kroon, G., Prusiner, S. B., Cohen, F. E., Dyson, H. J., and Wright, P. E. (2001) Local structural plasticity of the prion protein. Analysis of NMR relaxation dynamics. *Biochemistry* 40, 2743–2753.
- Kabsch, W., and Sander, C. (1983) Dictionary of protein secondary structure: pattern recognition of hydrogen-bonded and geometrical features. *Biopolymers* 22, 2577–2637.
- Scouras, A. D., and Daggett, V. (2008) Species variation in PrPSc protofibril models. J. Mater. Sci. 43, 3625–3637.
- Van der Kamp, M. W., and Daggett, V. (2010) Pathogenic mutations in the hydrophobic core of the human prion protein can promote structural instability and misfolding, *J. Mol. Biol.* (DOI: 10.1016/j. jmb.2010.09.060).
- Richardson, J. S., Getzoff, E. D., and Richardson, D. C. (1978) The β bulge: a common small unit of nonrepetitive protein structure. *Proc. Natl. Acad. Sci. U.S.A.* 75, 2574–2578.
- Armen, R. S., DeMarco, M. L., Alonso, D. O., and Daggett, V. (2004)
 Pauling and Corey's α-pleated sheet structure may define the prefibrillar amyloidogenic intermediate in amyloid disease. *Proc. Natl. Acad. Sci. U.S.A. 101*, 11622–11627.
- 30. Haire, L. F., Whyte, S. M., Vasisht, N., Gill, A. C., Verma, C., Dodson, E. J., Dodson, G. G., and Bayley, P. M. (2004) The crystal structure of the globular domain of sheep prion protein. *J. Mol. Biol.* 336, 1175–1183.
- Knaus, K. J., Morillas, M., Swietnicki, W., Malone, M., Surewicz, W. K., and Yee, V. C. (2001) Crystal structure of the human prion protein reveals a mechanism for oligomerization. *Nat. Struct. Biol.* 8, 770–774.
- 32. Lee, S., Antony, L., Hartmann, R., Knaus, K. J., Surewicz, K., Surewicz, W. K., and Yee, V. C. (2010) Conformational diversity in prion protein variants influences intermolecular β -sheet formation. *EMBO J.* 29, 251–262.
- De Simone, A., Dodson, G. G., Verma, C. S., Zagari, A., and Fraternali, F. (2005) Prion and water: tight and dynamical hydration sites have a key role in structural stability. *Proc. Natl. Acad. Sci. U.S.* A. 102, 7535–7540.
- 34. Santini, S., Claude, J. B., Audic, S., and Derreumaux, P. (2003) Impact of the tail and mutations G131V and M129V on prion protein flexibility. *Proteins* 51, 258–265.
- Alonso, D. O. V., DeArmond, S. J., Cohen, F. E., and Daggett, V. (2001) Mapping the early steps in the pH-induced conformational conversion of the prion protein. *Proc. Natl. Acad. Sci. U.S.A.* 98, 2985–2989.
- DeMarco, M. L., and Daggett, V. (2004) From conversion to aggregation: protofibril formation of the prion protein. *Proc. Natl.* Acad. Sci. U.S.A. 101, 2293–2298.

- 37. Chebaro, Y., and Derreumaux, P. (2009) The conversion of helix H2 to β -sheet is accelerated in the monomer and dimer of the prion protein upon T183A mutation. *J. Phys. Chem. B* 113, 6942–6948.
- 38. DeMarco, M. L., and Daggett, V. (2007) Molecular mechanism for low pH triggered misfolding of the human prion protein. *Biochemistry* 46, 3045–3054.
- 39. Armen, R. S., Alonso, D. O. V., and Daggett, V. (2004) Anatomy of an amyloidogenic intermediate: conversion of β -sheet to α -sheet structure in transthyretin at acidic pH. *Structure* 12, 1847–1863.
- Armen, R. S., Bernard, B. M., Day, R., Alonso, D. O. V., and Daggett, V. (2005) Characterization of a possible amyloidogenic precursor in glutamine-repeat neurodegenerative diseases. *Proc. Natl. Acad. Sci. U.S.A.* 102, 13433–13438.
- 41. Armen, R. S., and Daggett, V. (2005) Characterization of two distinct β_2 -microglobulin unfolding intermediates that may lead to amyloid fibrils of different morphology. *Biochemistry* 44, 16098–16107.

- Schmidlin, T., Kennedy, B. K., and Daggett, V. (2009) Structural changes to monomeric CuZn superoxide dismutase caused by the familial amyotrophic lateral sclerosis-associated mutation A4V. *Bio*phys. J. 97, 1709–1718.
- 43. Steward, R. E., Armen, R. S., and Daggett, V. (2008) Different disease-causing mutations in transthyretin trigger the same conformational conversion. *Protein Eng.*, *Des. Sel.* 21, 187–195.
- 44. Cordier, F., Nisius, L., Dingley, A. J., and Grzesiek, S. (2008) Direct detection of N−H···O=C hydrogen bonds in biomolecules by NMR spectroscopy. *Nat. Protoc.* 3, 235–241.
- 45. Barfield, M. (2002) Structural dependencies of interresidue scalar coupling ^{h3} J_{NC} and donor ¹H chemical shifts in the hydrogen bonding regions of proteins. *J. Am. Chem. Soc.* 124, 4158–4168.
- Sass, H. J., Schmid, F. F. F., and Grzesiek, S. (2007) Correlation of protein structure and dynamics to scalar couplings across hydrogen bonds. J. Am. Chem. Soc. 129, 5898–5903.