Prion diseases are caused by the pathological deposition of the native α-helical prion protein PrP\textsuperscript{C} into its pathological β-sheet-rich isoform PrP\textsuperscript{Sc}. In current models of PrP\textsuperscript{Sc}, helix1 is assumed to be preferentially converted into β-sheet during aggregation of PrP\textsuperscript{C}. This was supported by the NMR structure of PrP\textsuperscript{C} since, in contrast to the isolated helix1, helix2 and helix3 are connected by a small loop and are additionally stabilized by an interhelical disulfide bond. However, helix1 is extremely hydrophilic and has a high helix propensity. This prompted us to investigate the role of helix1 in prion aggregation using native PrP\textsuperscript{C}.

In conclusion, we provide strong direct evidence that helix1 is not converted to β-sheet. This is confirmed by Fourier transformation infrared spectroscopy since there is almost no difference in β-sheet content of fibrillar PrP\textsuperscript{23–159} fibrils compared with PrP\textsuperscript{23–144} fibrils. In contrast to earlier assumptions helix1 is not involved in the aggregation process. However, helix1 is not resistant to digestion with proteinase K in fibrillar PrP\textsuperscript{23–159}, suggesting that helix1 is not converted to β-sheet. This is demonstrated by Fourier transformation infrared spectroscopy since there is almost no difference in β-sheet content of fibrillar PrP\textsuperscript{23–159} fibrils compared with PrP\textsuperscript{23–144}. In conclusion, we provide strong direct evidence that in contrast to earlier assumptions helix1 is not converted to β-sheet during aggregation of PrP\textsuperscript{C} to PrP\textsuperscript{Sc}.

Prion diseases are caused by the pathological deposition of the prion protein in its aggregated form. In sporadic Creutzfeldt-Jakob disease (CJD)\textsuperscript{3} the endogenous prion protein aggregates spontaneously. In contrast, mutations within the prion protein gene cause familial CJDs and Gerstmann-Sträussler-Scheinker syndrome (GSS) including two stop mutations Y145Stop and Q160Stop with a GSS-like phenotype. The most important aspect is the transmission of prion protein aggregates from one individual or species to another, causing prion diseases such as scrapie in sheep, bovine spongiform encephalopathy (BSE) in cattle, chronic wasting disease in deer, or variant CJD in humans. So far all experimental data strongly support the hypothesis that only one protein, the prion protein, is responsible for this virus-like infectivity (1).

Native prion protein (PrP\textsuperscript{C}) is attached to the extracellular plasma membrane surface by a glycosylphosphatidylinositol lipid anchor and undergoes endocytosis. In vivo endocytosis was shown to be essential for conversion of PrP\textsuperscript{C} into the pathological and infectious PrP\textsuperscript{Sc}. This suggested that changes in pH are involved in the conversion process. Indeed, a β-sheet-rich oligomeric intermediate (β-PrP) was identified at acidic pH under denaturing conditions (2, 3). Longer incubation resulted in fibril formation (4, 5).

Many efforts have been made to elucidate the molecular mechanisms of prion protein aggregation. Investigation of known mutations in prion protein associated with GSS and CJD showed only partially a thermodynamic destabilization of the ground states (6, 7). Therefore, a thermodynamic destabilization of PrP\textsuperscript{C} was excluded as a general mechanism for PrP\textsuperscript{Sc} formation. The oligomeric β-sheet-rich β-PrP was even more stable than PrP\textsuperscript{C} (8). However, there was a high energetic barrier between the PrP\textsuperscript{C} and the β-PrP form, which clearly demonstrates a kinetic control of the conversion process (8). This barrier was shown to be significantly dependent on pH (9).

The N-terminal segment of PrP\textsuperscript{C} from amino acid 23 to 126 is largely unstructured (10). Two pathological GSS-like mutations, Y145Stop and Q160Stop, result in C-terminal truncated isoforms. The truncation occurs just after the central region from amino acid 90 to 145, which was shown to be converted into β-sheet (11, 12). In contrast to the full-length prion protein, the Y145Stop mutant was shown to aggregate under native non-denaturing rather neutral pH conditions (13). Working under non-denaturing conditions appears promising since for the first time an in vitro model for the species barrier was established solely by recombinant PrP\textsuperscript{23–144} (12, 14). The present investigation aimed at clarifying the role of helix1 in the conversion process of the prion protein under native conditions.

During conversion to PrP\textsuperscript{Sc} a significant part of the prion protein adopts β-sheet structure (15). However, the role of helix1 (16) in this process is still unclear. Originally, helix1 was thought to be
Plasmid Construction and Protein Purification—The open reading frame coding for humPrP (23–231) was amplified from human tissue and cloned into the plasmid pBluescript II SK(+) (Stratagene). The open reading frames for humPrP (23–144) and humPrP (23–159) were amplified from the plasmid pBluescript II SK(+)//humPrP (23–231) using the oligonucleotide primers 23MSup (5'-CGACGTAAGCTTACATGAGCAAGAAGC CCCCAAG-3') and 145do (5'-ACGATTATGAGCTGAC CATATTAGTC ACTGCCGAAAATG-3') or 160do (5'-GGGCCCTGTAGTCGACTTATGTGGTAGA CGGATG-3'). The resulting open reading frames were cloned into the plasmid pET27b (Novagen) via the Ndel and SalI restriction sites. Full-length prion protein and stop mutants were expressed in BL21(DE3) and purified as described previously (18).

CD Spectroscopy—Far UV circular dichroism spectra were recorded on a CD6 (Jobin Yvon) in 25 mM sodium acetate, pH 6.0, at 25 °C using a quartz cell with a path length of 0.1 mm. Protein concentrations were 80 μM. All spectra were corrected for absorbance of buffer.

Thioflavin T Fluorescence Assay—Mutant proteins were directly dissolved in water, and the pH was adjusted to 6.5. The reaction was started by transferring protein solutions to a buffer (25 mM acetate and 25 mM phosphate at pH 6.5) and incubated at 25 °C. In seeding experiments 1% (w/w) sonicated seed of human PrP (23–231) was added to 200 μM protein to proteinase K (PK) ratios of 104:1 (50 μg/ml PK) and 52:1 ratio (100 μg/ml PK), respectively. The digestion was stopped with 1 mM phenylmethylsulfonyl fluoride, boiled for 10 min in SDS sample buffer, and analyzed on a 17.5% SDS–polyacrylamide gel as described (19). For Western blot analysis 6H4 (1:10,000) and 1E4 (1:10,000) were used, and immunoreactivity was detected by chemiluminescence. Edman sequencing was performed using a Procise cLC protein sequencer (Applied Biosystems) according to the manufacturer’s standard protocols.

Fourier Transformation InfraRed Spectroscopy (FTIR)—FTIR measurements were performed on a Nicolet 6700 (Thermo Electron Corp.) constantly purged with dried air. All buffers and mutant proteins were exchanged three times with D2O by lyophilization. Aggregation was performed at a pD of 6.5 using 25 mM phosphate and 25 mM acetic acid as buffer. For measurements, CaF2 windows with a 50-μm Teflon spacer were used. In total 256 spectra were averaged with a resolution of 2 cm⁻¹. Spectra were corrected for buffer absorbance.

Peak fitting was done with PeakFit v. 4 (SPSS Inc.) using the option of automatic peak detection by the second derivative method. For fitting the implemented real Voigt function was used considering both Gaussian and Lorentzian peak broadening.

Electron Microscope Analysis of Aggregates—After termination of the aggregation aliquots of 10 μl were placed on a carbon-coated copper grid and stained with 2% aqueous (w/v) uranyl acetate for 5 min. After washing and drying, protein aggregates on the grids were visualized using a Zeiss 10B electron microscope operating at 60 kV.

NMR Analysis of humPrP23–159—humPrP23–159 was 13C- and 15N-labeled in M9 minimal medium according to standard procedures. The NMR sample contained 1 mM 13C,15N-labeled humPrP23–159 in 93% H2O, 7% D2O, 10 mM sodium acetate buffer, pH 4.5 and 6.5. All spectra were acquired at 293 K on a
Bruker Avance 900 MHz spectrometer equipped with a cryo probe head. NMR data were processed and analyzed using nmrPipe (20) and Sparky. Three-dimensional HACANNH, HNCO, and HNN experiments (21) were collected to obtain sequence-specific backbone assignment. Secondary chemical shifts were calculated as differences between the measured Cα/C′ shifts and empirical random coil values at pH 3.0 (22). Random coil values for prolines and aspartates were taken from Wishart and Sykes (23).

RESULTS

Structure of the Native Q160Stop Mutant—Before starting the aggregation experiments, both mutant proteins were subjected to capillary electrophoresis, which was shown previously to detect oligomers early in the aggregation process (24). The electropherograms showed only one single peak each, which clearly demonstrated their monomeric state (data not shown).

Because human PrP23–159 includes the helix1, we examined whether it was detectable using CD spectroscopy (Fig. 1). However, the CD spectrum of human PrP23–159 is almost identical to that of human PrP23–144, which corresponds to a spectrum of random coil, as shown previously for human PrP23–144 (25). It appeared that the fraction of amino acids in human PrP23–159 corresponding to helix1 is too low to be detected by any analytical technique that determines the sequence-independent average conformation, such as CD spectroscopy.

Therefore, the structure of the helix1 segment in human PrP23–159 was analyzed by NMR spectroscopy. NMR chemical shifts, especially of Cα and C′ atoms, are a very sensitive indicator for sequence-dependent secondary structure both in globular and unfolded proteins (26). These shifts show small but distinct deviations from random coil values. The deviations, called secondary chemical shifts, are shown in Fig. 2. A continuous stretch of positive secondary chemical shifts was
Prion Helix1 Is Not Converted to $\beta$-Sheet

FIGURE 3. Extension of humPrP$_{23-144}$ by helix1 promotes aggregation in humPrP$_{23-159}$. Independent of the used protein concentrations of 340 $\mu$M (A), 200 $\mu$M (B), and 80 $\mu$M (C), humPrP$_{23-159}$ (circles) aggregated significantly faster compared with humPrP$_{23-144}$ (triangles). Unseeded aggregation experiments were performed at pH 6.5, and fibril formation was monitored by thioflavin T (THT) fluorescence. Solid lines represent the result for fitting of the data according to Equation 1.

found for residues 144–154, indicative of helical structure. Most other residues show slightly negative secondary chemical shifts, reporting on a propensity to form $\beta$-structure. Interestingly, the highest propensity to form $\beta$-structure is present for residues 98–143, a region known to be converted into $\beta$-sheet on PrP$_{Sc}$ formation (11). In particular Ile-139, one of two residues that are critical for the conformational properties of humPrP$_{23-144}$ fibrils (14), has the most negative secondary chemical shift, i.e. the strongest propensity for $\beta$-structure.

NMR secondary chemical shifts clearly demonstrate the presence of a significantly populated helix 1 in humPrP$_{23-159}$. Comparison of average shift magnitudes for the helix1 region observed in humPrP$_{23-159}$ and humPrP$_{23-230}$ (Biological Magnetic Resonance Data Bank entry 4402) revealed an $\alpha$-helical content of 33% in humPrP$_{23-159}$ with respect to the full-length protein (data not shown). As expected from the data on humPrP$_{140-158}$ (27), helix1 is present in humPrP$_{23-159}$ at pH 4.5 as well as pH 6.5 (Fig. 2B).

humPrP$_{23-159}$ Aggregates Faster than humPrP$_{23-144}$—To obtain direct insight into the role of the helix1 during prion protein aggregation, we investigated the aggregation kinetics of humPrP$_{23-159}$ compared with humPrP$_{23-144}$. We demonstrate here for the first time that humPrP$_{23-159}$ readily aggregates under physiological non-denaturing conditions (Fig. 3). However, in contrast to the Y145Stop mutant, humPrP$_{23-159}$ is extended C-terminally by the helix1 segment. Apart from the presence of helix in the helix1 segment of humPrP$_{23-159}$, this segment is still rather polar, including 50% charged residues. Therefore, we expected this to significantly increase the lag phase with respect to humPrP$_{23-144}$. In addition, full-length prion protein has not yet been shown to aggregate under these conditions. Most unexpectedly, we observed the opposite: independent from the protein concentration, humPrP$_{23-159}$ aggregated significantly faster compared with humPrP$_{23-144}$, which clearly demonstrates an aggregation-promoting effect of helix1. From triple experiments, the average aggregation halftime of humPrP$_{23-159}$ at 340 $\mu$M was 39 h compared with 67 h for the Y145Stop mutant. At 200 and 80 $\mu$M, the aggregation halftimes were 68 and 171 h for humPrP$_{23-159}$ or 138 h and 242 h for humPrP$_{23-144}$.

Electron microscopy revealed that humPrP$_{23-159}$, like humPrP$_{23-144}$, indeed forms fibrils (Fig. 4). However, no significant differences were observed for both mutants. Both form a fiber network composed of highly bundled fibrils with average diameters ranging from 50 to 70 nm but maximal up to 350 nm. These bundles extend over several micrometers and are formed by fibrils with diameters from 8–12 nm made up of two protofibrils. This is identical to the results shown for humPrP$_{23-144}$ (13).

Seeded Aggregation of humPrP$_{23-159}$ Is Faster Compared with humPrP$_{23-144}$—In contrast to the unseeded reaction, in a seeded aggregation the speed of aggregation is additionally determined by the nature of the seed. Therefore, we investigated whether the increased aggregation propensity of humPrP$_{23-159}$ is an intrinsic property of the protein by comparing the seeding reactions of both mutants by seeding with humPrP$_{23-144}$ and humPrP$_{106-126}$ fibrils.

Interestingly, the seeding of humPrP$_{23-159}$ with either humPrP$_{23-159}$ or humPrP$_{23-144}$ fibrils was much faster compared with seeding of humPrP$_{23-144}$ (Fig. 5). The average aggregation halftimes for seeding of humPrP$_{23-159}$ were almost identical with 4.3 h for seeding with humPrP$_{23-159}$ fibrils and 4.7 h for seeding with of humPrP$_{23-144}$. In contrast, seeding of humPrP$_{23-144}$ with both mutant fibrils was significantly different since this resulted in a pronounced biphasic aggregation kinetic (Fig. 5, D and E). Furthermore, fibrils of humPrP$_{23-144}$ were significantly more efficient in seeding comparisons with humPrP$_{23-159}$ fibrils. The average prephase aggregation halftimes were 2.7 and 10.9 h for seeding with humPrP$_{23-144}$ and humPrP$_{23-159}$ whereas the average aggregation halftimes of the main phase were 10.4 and 47.8 h, respectively.
The difference between both mutants could further be demonstrated by seeding with \textit{humPrP}_{106–126} fibrils (Fig. 5, C and F). To our surprise we observed a significant seeding of \textit{humPrP}_{23–144} which is in contrast to a previous report (13). At present the reason for this difference remains unclear, although the lower concentration of \textit{humPrP}_{23–144} in our experiments is more favorable for detecting any seeding effect. However, again seeding of \textit{humPrP}_{23–159} was faster compared with \textit{humPrP}_{23–144}, resulting in aggregation half-times of 42 and 71 h. In conclusion, the results of the seeded and unseeded reactions clearly demonstrate that the higher aggregation propensity of \textit{humPrP}_{23–159} compared with \textit{humPrP}_{23–144} is an intrinsic property of the protein.

\textit{Helix1 in humPrP}_{23–159}—\textbf{Fibrils Is Not PK-resistant}—The comparison of the aggregation kinetics clearly demonstrated that helix1 is involved in the aggregation process, but the mechanism as to how it promotes aggregation remained unclear. It would be most obvious if helix1 is directly converted into a protease-resistant \textit{β}-sheet on aggregation. To examine this, prion aggregates of both mutants were subjected to digestion with PK and analyzed by SDS-PAGE. Due to significant smearing with classical SDS-PAGE we used an alternative SDS-PAGE system, which was demonstrated to separate the also aggregation-prone A\textit{β}-peptides differing by only one amino acid in size (28). Fibris of both mutants were quite resistant to concentrations as high as 100 μg/ml PK for 1 h (Fig. 6A). The apparent size of 7 kDa for the PK-resistant fragment of the Y145Stop mutant fibrils is identical to that reported earlier (29). The increased apparent size of 8 kDa for the PK-resistant fragment of \textit{humPrP}_{23–159} fibrils (Fig. 6A) suggested either N- or C-terminal extension of the 160Stop fragment compared with the Y145Stop mutant. This difference may result from different known N-terminal cleavage sites of human PrP\textsuperscript{Sc} at residue 82 (type 1) and 97 (type 2) (30). Western blot analysis using antibody 3B5, which recognizes the type 1 cleavage pattern, was negative for the PK-resistant fragments (data not shown), whereas the 1E4 antibody directed against amino acids 98–109 (Fig. 6E) recognized PK-resistant fragments of both mutants (Fig. 6C), suggesting, rather, a type 2 cleavage pattern. Indeed, Edman sequencing of both PK-resistant cores resulted in cleavage sites at amino acids 97 and 98 (Fig. 6B), which corresponds exactly to the cleavage pattern type 2 (30). This suggested that the PK-resistant fragment of \textit{humPrP}_{23–159} is C-terminal-extended by the helix1. However, in contrast to the soluble isoform, none of the helix1 antibodies 6H4 (Fig. 6D) and 12F10 (data not shown) recognized the PK-resistant fragments of \textit{humPrP}_{23–159} fibrils.

Thus, helix1 appears not to be converted into PK-resistant \textit{β}-sheet in \textit{humPrP}_{23–159} fibrils. The observed size difference of the PK-resistant fragments is probably due to C-terminal extension of the \textit{humPrP}_{23–159} fragment by a few amino acids since, as mentioned above, the used SDS-PAGE system is able to separate A\textit{β}-peptides differing only by one amino acid in size (28).

\textit{Helix1 Is Not Converted to β-Sheet}—To further confirm that helix1 is not converted into \textit{β}-sheet, we analyzed the aggregates by FTIR. If helix1 from amino acid 144–156 (31) is completely converted, there should be an increase of 9.5% in the \textit{β}-sheet fraction in aggregates from \textit{humPrP}_{23–159} compared with \textit{humPrP}_{23–144}.

Curve fitting analysis (Fig. 7) resulted in average peak wave numbers of 1627, 1640, 1652, and 1664 cm\textsuperscript{-1}, almost identical to that observed previously for \textit{humPrP}_{23–144} (13). It was...
recently demonstrated that amide I peaks in the range 1611–1630 cm\(^{-1}\) are characteristic for \(\beta\)-sheet formed by amyloid fibrils, whereas native \(\beta\)-sheet clustered from 1630 to 1643 cm\(^{-1}\) (32). Therefore, the peaks at 1615 and 1627 cm\(^{-1}\) were assigned to amyloid \(\beta\)-sheet and \(\beta\)-sheet, respectively. The observed maximum at 1640 cm\(^{-1}\) is very close to the classical random coil amide I at 1643 cm\(^{-1}\). The amide I peak at 1652 cm\(^{-1}\) corresponded to classical \(\alpha\)-helix at 1650 cm\(^{-1}\) in D\(_2\)O. However, the formation of classical \(\alpha\)-helix on aggregation appeared rather unlikely. Because the non-protease-resistant N-terminal segment from amino acids 23–98 comprises more than 50% of both stop mutant molecules, this helix-like peak was attributed to the N-terminal segment. The sum of both peak areas at 1640 and 1652 cm\(^{-1}\) is very close to the classical random coil amide I at 1643 cm\(^{-1}\). The amide I peak at 1652 cm\(^{-1}\) corresponded to classical \(\alpha\)-helix at 1650 cm\(^{-1}\) in D\(_2\)O. However, the formation of classical \(\alpha\)-helix on aggregation appeared rather unlikely. Because the non-protease-resistant N-terminal segment from amino acids 23–98 comprises more than 50% of both stop mutant molecules, this helix-like peak was attributed to the N-terminal segment. The sum of both peak areas at 1640 and 1652 cm\(^{-1}\) is very close to the classical random coil amide I peak at 1643 cm\(^{-1}\). The amide I peak at 1652 cm\(^{-1}\) corresponded to classical \(\alpha\)-helix at 1650 cm\(^{-1}\) in D\(_2\)O. However, the formation of classical \(\alpha\)-helix on aggregation appeared rather unlikely. Because the non-protease-resistant N-terminal segment from amino acids 23–98 comprises more than 50% of both stop mutant molecules, this helix-like peak was attributed to the N-terminal segment. The sum of both peak areas at 1640 and 1652 cm\(^{-1}\) is very close to the classical random coil amide I wave number observed in PrP\(_{23–144}\) aggregates (13). Thus, in contrast to seeding of humPrP\(_{23–159}\) fibrils with humPrP\(_{106–126}\) resulted in only one amyloid \(\beta\)-sheet peak located at 1620 cm\(^{-1}\) (Fig. 7F). This corresponds exactly to the amyloid \(\beta\)-sheet wave number observed in PrP\(_{106–126}\) aggregates (13). Thus, in contrast to seeding of humPrP\(_{23–159}\) fibrils with humPrP\(_{106–126}\) fibrils are able to transmit their structural properties to humPrP\(_{23–144}\) on seeding.

The most interesting question was whether the C-terminal extension, including helix1, is converted into \(\beta\)-sheet. Theoretically, complete conversion of this extension (amino acids 145–159) into \(\beta\)-sheet should result in an increase of up to 11% in \(\beta\)-sheet compared with the Y145Stop mutant. Considering only the unseeded reactions, we obtained 24.1% pure random coil, 21.8% helix-like structure, and 30.7% \(\beta\)-sheet for the Q160Stop mutant (Table 1). For Y145Stop, we obtained 21.4% pure random coil, 20.5% helix-like structure, and 34.7% \(\beta\)-sheet. However, we had to take into consideration that Q160Stop mutant is longer by 15 amino acids. Thus, the percentages of Y145Stop were corrected by multiplying with 0.89.

![FIGURE 6. Helix 1 segment is not resistant to proteinase K in humPrP\(_{23–159}\) fibrils. Fibrils of humPrP\(_{23–159}\) and humPrP\(_{23–144}\) were subjected to digestion with proteinase K. A, SDS-PAGE analysis of digested fibrils according to Wiltfang et al. (28). This SDS-PAGE system was demonstrated to separate \(\beta\) peptides differing only by one amino acid (28). B, Edman sequencing of the PK-resistant fragments revealed identical N-terminal sequences. C, Western blot analysis using antibody 1E4 confirms the proteinase K cleavage pattern 2 of both fibrils. D, Western blot analysis demonstrate that the helix1 is not resistant to digestion with proteinase K since neither antibody 6H4 nor 12F10 (data not shown) recognize digested humPrP\(_{23–159}\) fibrils (F), in contrast to the undigested monomeric isoform (M). E, in contrast to the antibody 1E4, the 6H4 antibody is only able to recognize humPrP\(_{23–159}\) due to the location of epitopes schematically depicted for both mutants.](image)
resulting in 19.1% pure random coil, 18.2% helix-like structure, and 30.9% β-sheet. There is almost no difference in total β-sheet since β-sheet of humPrP23–159 is only 0.2% higher compared with humPrP23–159 fibrils. However, there is an increase of 3.6% in α-helix like structure in humPrP23–159 fibrils, which suggests that some α-helix could be present. Finally, random coil was increased by 5.0%. Taken together, there is no significant difference in β-sheet but 8.6% of the possible 11% secondary structure, corresponding to the fraction of the C-terminal extension in humPrP23–159 compared with humPrP23–144, is due to random coil or helix. This together with the data from proteinase K digestion clearly indicates that helix1 is not converted into β-sheet.

**DISCUSSION**

The aggregation-promoting effect of the highly charged helix1 in humPrP23–159 relative to humPrP23–144 is surprising since, in general, truncation of non-amyloidogenic regions in 

| Secondary structure assignment | Wave number | Percentage of secondary structure element |
|-------------------------------|-------------|------------------------------------------|
|                               | 1615        | 11.6                                     |
| Amyloid β-sheet               | 1627        | 11.8                                     |
| β-Sheet                       | 1640        | 10.8                                     |
| Random                        | 1652        | 21.1                                     |
| α-Helix                       | 1664        | 15.6                                     |
| Turn/loop                     | 1675        | 9.0                                      |
| Amyloid β-sheet               | 1684        | 3.1                                      |

| Secondary structure assignment | Wave number | Percentage of secondary structure element |
|--------------------------------|-------------|------------------------------------------|
|                               | 1615        | 12.4                                     |
| Amyloid β-sheet               | 1627        | 13.1                                     |
| β-Sheet                       | 1640        | 18.1                                     |
| Random                        | 1652        | 17.9                                     |
| α-Helix                       | 1664        | 16.7                                     |
| Turn/loop                     | 1675        | 9.5                                      |
| Amyloid β-sheet               | 1684        | 4.5                                      |

**TABLE 1**

Quantitative analysis for FTIR spectra of prion aggregates

![FTIR spectroscopic analysis of fibrillar humPrP23–159 (A–C) and humPrP23–144 (D–F) to determine their secondary structure content by peak fitting (Table 1).](image)
proteins prone to aggregation promotes their aggregation. Calculation of the aggregation tendency using Zygregator (34) resulted in a significant lower aggregation propensity for PrP\(_{95–159}\) (Zagg = -0.80) compared with PrP95–144 (Zagg = -0.18). Normally charged amino acids are even added to peptides to increase their solubility, thereby preventing their aggregation. Interestingly, the increased aggregation propensity of \textit{humPrP}\(_{23–159}\) compared with \textit{humPrP}\(_{23–144}\) correlates also \textit{in vivo} with the 6-year earlier onset of clinical symptoms in one patient with a prion Q160Stop mutation (35) compared with the Y145Stop mutation (29).

Our results clearly demonstrate that helix1 is involved in the aggregation of \textit{humPrP}\(_{23–159}\). However, at present the unusual nature of this aggregation-promoting effect is unclear, although it was speculated that helix1 might form intermolecular salt bridges on aggregation of the prion protein (36).

Although \textit{humPrP}\(_{23–159}\) aggregated significantly faster compared with \textit{humPrP}\(_{23–144}\), both PK-digestion and FTIR analysis clearly demonstrate that helix1 is not converted into \textit{β-sheets}. Thus, we provide here for the first time strong direct evidence that helix1 in PrP\(_{\text{Sc}}\) is not converted into \textit{β-sheet}. This is confirmed by the fact that both PrP\(_{\text{Sc}}\) as well as PrP\(_{\text{Sc}}\) can be precipitated with the antibody 6H4, which recognizes an epitope in helix1 (37). Antibodies directed against helix1 are even able to block the conversion from PrP\(_{\text{Sc}}\) to PrP\(_{\text{Sc}}\) (38, 39). Furthermore, the unusual high propensity for α-helix formation of the helix1 segment (27) and its unusual low aggregation propensity (40) are in line with our results. Helix1 was completely dispensable for the generation of infectious PrP\(_{\text{Sc}}\) (41, 42), which further supports our results.

At present neither NMR nor x-ray crystallography is able to completely resolve the three-dimensional structure of PrP\(_{\text{Sc}}\). Therefore, evidence in the puzzle of the PrP\(_{\text{Sc}}\) structure can only be obtained by different biochemical approaches. We provide here strong direct evidence that in contrast to earlier assumptions (43, 17), helix1 is not converted to \textit{β-sheet} in PrP\(_{\text{Sc}}\). However, our results leave the possibility that helix1 is shortened in PrP\(_{\text{Sc}}\) since there is an increase in apparent helix content of 4% compared with maximal possible 9.5% in fibrils of \textit{humPrP}\(_{23–159}\) relative to \textit{humPrP}\(_{23–144}\). Interestingly, our data agree with results of molecular dynamic simulations for pH-induce conformational conversion. Although helix1 experienced some loss of helix, none of the lost amino acids was involved in the formation of \textit{β-sheet} (44).

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