Allelic Origin of Protease-Sensitive and Protease-Resistant Prion Protein Isoforms in Gerstmann-Sträussler-Scheinker Disease with the P102L Mutation

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Abstract

Gerstmann-Sträussler-Scheinker (GSS) disease is a dominantly inherited prion disease associated with point mutations in the Prion Protein gene. The most frequent mutation is associated with GSS involves a proline-to-leucine substitution at residue 102 of the prion protein, and is characterized by marked variability at clinical, pathological and molecular levels. Previous investigations of GSS P102L have shown that disease-associated pathological prion protein, or PrP Sc, consists of two main conformers, which under exogenous proteolysis generates a core fragment of 21 kDa and an internal fragment of 8 kDa. Both conformers are detected in subjects with spongiform degeneration, whereas only the 8 kDa fragment is recovered in cases lacking spongiosis. Several studies have reported an exclusive derivation of protease-resistant PrP Sc isoforms from the mutated allele; however, more recently, the propagation of protease-resistant wild-type PrP Sc has been described. Here we analyze the molecular and pathological phenotype of six GSS P102L cases characterized by the presence of 21 and 8 kDa PrP fragments and two subjects with only the 8 kDa PrP fragment. Using sensitive protein separation techniques and Western blots with antibodies differentially recognizing wild-type and mutant PrP we observed a range of PrP Sc allelic conformers, either resistant or sensitive to protease treatment in all investigated subjects. Additionally, tissue deposition of protease-sensitive wild-type PrP Sc molecules was seen by conventional PrP immunohistochemistry and paraffin-embedded tissue blot. Our findings enlarge the spectrum of conformational allelic PrP Sc quasispecies propagating in GSS P102L thus providing a molecular support to the spectrum of disease phenotypes, and, in addition, impact the diagnostic role of PrP immunohistochemistry in prion diseases.

Citation: Monaco S, Fiorini M, Farinazzo A, Ferrari S, Gelati M, et al. (2012) Allelic Origin of Protease-Sensitive and Protease-Resistant Prion Protein Isoforms in Gerstmann-Sträussler-Scheinker Disease with the P102L Mutation. PLoS ONE 7(2): e32382. doi:10.1371/journal.pone.0032382

Editor: Noriyuki Nishida, Nagasaki University Graduate School of Biomedical Sciences, Japan

Received November 19, 2011; Accepted January 29, 2012; Published February 23, 2012

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Funding: This work was supported by grants R01 NS029822 to BG from National Institute of Neurological Disease and Stroke (NINDS), P30 AG10133 from National Institute on Aging (NIA) to BG (both Institutes are part of the National Institutes of Health) and by the grant “Disabilità cognitiva e comportamentale nelle demenze e nelle psicosi” from Fondazione Cariverona to SM. The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

Competing Interests: The authors have declared that no competing interests exist.

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Introduction

Prion diseases are fatal neurodegenerative disorders of humans and animals, occurring as idiopathic, genetic, and iatrogenic conditions [1]. Human prion disorders include Creutzfeldt-Jakob disease (CJD), variant Creutzfeldt-Jakob disease, kuru, familial insomnia, sporadic fatal insomnia, and Gerstmann-Sträussler-Scheinker (GSS) disease [2]. In these disorders, the cellular prion protein, or PrP C, is converted to a protease-resistant conformer with abnormal sedentation properties, or PrP Sc [3]. In some prion diseases variant PrP Sc discloses distinct proteolytic sites generating a core fragment named PrP 27–30 [4], reflecting the tertiary structure of PrP Sc, in addition to variable C-terminal truncated fragments (CTFs) and internal fragments [5,6,7,8,9]. PrP Sc conformers are classified on the basis of the glycosylation status and the molecular mass of their protease-resistant unglycosylated backbone, or PrP 27–30 core fragment, migrating at 21 kDa in type 1 PrP Sc and at 19 kDa in type 2 PrP Sc [10], although controversial results have been obtained by different authors who have proposed the classification of PrP 27–30 into four types [11]. However, several prion strains, either in naturally occurring disorders or in experimentally-induced conditions, harbour protease-sensitive PrP Sc (sPrP Sc ) quasispecies, in association or not with protease-resistant PrP Sc (rPrP Sc ) [12,13,14]; importantly, sPrP Sc molecules cause neurological dysfunction [15,16]. The neuropathological hallmarks of prion diseases are prion protein (PrP) deposition, spongiosis, astroglisis, and neuronal death [17]; however, it is still unknown which molecular forms of PrP Sc are detected by immunohistochemistry (IHC).

GSS is an autosomal dominant prion disease associated with point mutations in the Prion Protein gene (PRNP). The most frequent mutation associated with GSS involves a proline-to-leucine substitution within the unstructured N-terminal region of the PrP at codon 102 (P102L), segregating with methionine at codon 129 on the mutant allele [18]. Other mutations have been described either at the N-terminal or at the C-terminal PrP region.
The genetic heterogeneity of GSS partly explains the variability in age of onset, disease duration, and the ample number of clinical variants. However, phenotypic heterogeneity is also observed both within and between pedigrees with the same mutation, especially in P102L-GSS cases [19,20,21]. Accordingly, while in some P102L patients progressive gait disturbances, truncal ataxia, lower limb hyporeflexia and dysarthria are the presenting clinical features, followed over years by dementia and pyramidal signs, in other cases a rapid evolving dementia, indistinguishable from sporadic CJD (sCJD), is observed. As yet, the molecular basis for such clinical variability remains to be fully explained. Neuropathological examination in GSS P102L discloses marked variation in the degree and brain regional distribution of spongiform changes, which largely correlates with the biochemical detection of proteinase K (PK)-resistant PrPSc with a core fragment of 21 kDa [8,9,22].

As opposed to the variable topographical distribution of spongiform degeneration, typical multi-centric plaques are widely distributed throughout cerebral and cerebellar cortical and subcortical regions, and appear to be constituted by an abnormal PrP isoform that upon PK-digestion generates an internal 8 kDa fragment (PrP8), overall spanning from residues 74–90 to 146–153; this fragment is natively present in brain extract of GSS P102L, although is enriched after PK treatment. Experimental transmission of GSS P102L brain tissues with only PrP8 induces PrP-amyloid deposition, but not spongiform encephalopathy [23]. Earlier studies have shown that abnormal PrP isoforms detected in P102L mutation derive from mutated PrP (PrPM) [8]. Recently, this view has been challenged and proof has been obtained of wild-type PrP (PrPWt) involvement in three GSS P102L cases with the 21 kDa fragment [24]. Additionally, evidence has been provided that in transgenic (Tg) mice overexpressing the P101L mutation [Tg(PrP,P101L)], corresponding to GSS P102L mutation, spontaneous formation of mutant sPrPSc molecules occurs in brain tissues [15,16]. Moreover, in Tg(PrP,P101L) expressing high levels of mutant PrP, spontaneous prion disease is accompanied by generation of sPrPSc [14]. Endogenous generation of sPrPSc was also observed in Tg196 mice expressing lower levels of P101L mutated protein (about 2-fold higher than wild type), following inoculation with a synthetic peptide bearing the P101L mutation [25]. While the occurrence of rPrPSc and sPrPSc has been extensively investigated in experimental models of GSS P102L, no such studies have been systematically performed in the human disorder. The present study was designed to investigate in eight subjects with GSS P102L the extent of the contribution of PrPWt and PrPM in the generation of PrPSc conformers, including sPrPSc and rPrPSc molecules. For this purpose, we selected brain tissues from six cases with the 21 kDa and the 8 kDa fragments, either methionine homozygous or heterozygous at codon 129 of the PrP gene, and two cases harbouring only PrP8, both methionine homozygous. Frozen brain samples were investigated by conventional biochemical techniques, in addition to sensitive protein separation techniques and Western blot with antibodies differentially recognizing the two allelic forms of PrP and PrPWt. In addition, fixed tissues from the foregoing subjects were investigated by IHC and PET-blot to assess the extent of the contribution of distinct allelic PrP molecules to tissue deposition under different patterns. The results reported here demonstrate (i) a major contribution of PrPWt to the generation of rPrPSc species; (ii) the propagation of considerable amounts of mutated and wild-type sPrPSc molecules; (iii) the extensive tissue deposition and high PET blot signals of both allelic PrPSc species in all cases, including subjects lacking PrP27-30. Our findings strongly support experimental studies showing a major role of sPrPSc in causing neurodegeneration in a genetic human prion disease.

Results

Immunoblot analyses of protease-resistant PrPSc in P102L GSS

Demographic, genetic, and clinical data of all investigated subjects are summarized in Table 1. Following treatment with PK and immunoblot with 3F4, frontal cortex homogenates from six GSS P102L subjects showed the presence of three major PrP27-30 bands migrating at 30, 25, and 20 kDa, and a single 8 kDa band; two cases showed only trace amounts of the 8 kDa fragment in the absence of PrP27-30 (Fig. 1A). The migration of the unglycosylated PrP27-30 band was intermediate as compared to type 1 and type 2 sCJD-associated PrPSc, whereas the glycosylation profile showed a relative abundance of the diglycosylated form, similar to sCJD E200K, but at variance with the “monoglycosylated

Table 1. Demographic, molecular, and clinical features of investigated P102L GSS cases.

| CASE (code) | Codon 129 | Gender | Age at onset | Disease duration | Clinical signs at onset | Symptoms at evolution |
|-------------|-----------|--------|--------------|------------------|------------------------|----------------------|
| 1 (91-466)  | M/V       | M      | 56           | 60               | Ataxia, dysarthria, depression | Dementia |
| 2 (92-331)  | M/M       | M      | 31           | 36               | Ataxia, dysarthria | Cognitive decline, ataxia, dysarthria, abnormal eye movements |
| 3 (96-422)  | M/M       | M      | 65           | 60               | Ataxia, dysarthria and dysphasia | Dementia, agitation, sleep disturbance |
| 4 (98-196)  | M/M       | M      | 57           | 9                | Ataxia, mild cognitive impairment | Cerebellar signs and dementia |
| 5 (98-197)  | M/V       | M      | 31           | 11               | Confusion | Mental deterioration, apraxia, ataxia |
| 6 (2004-59) | M/F       | F      | 41           | 5                | Mild cognitive impairment, behavioral changes, dysarthria, mild ataxia | Abnormal sleep, agitation, dysphasia |
| 7 (2005-51) | M/V       | M      | 46           | 72               | Weakness, gait ataxia | Dementia |
| 8 (2005-52) | M/M       | M      | 50           | 36               | Cognitive decline | Behavioral disturbances, dementia |

doi:10.1371/journal.pone.0032382.t001
confirming the exclusive origin of this internal fragment from the mutated allele. Differently from 3F4-positive PrP27-30, including the total pool of mutated and wild type rPrPSc, ICSM 35-positive PrP27-30 showed a relative abundance of the monogydosylated forms, and a core fragment fragment migrating at 20–27 kDa. Semi-quantitative densitometric analyses showed that wild-type PK-resistant PrPSc accounted for 5–10% of the total 3F4-reactive PK-resistant PrPSc. Within each case, immunoblots with ICSM 35 and 3F4 showed similar results in brain homogenates from at least three different samples of the frontal cortex.

Two-dimensional analysis of protease-resistant PrPSc in P102L brains

After 2-D separation and immunoblot with 3F4, PrP27-30 resolved as two sets of glycosylated charge isomers of 30 and 27 kDa, with pIs ranging from 4.0 to 8.0, and three unglycosylated 20 kDa spots, migrating between 8.1-6.8; in addition, three basic spots were seen in the 8 kDa zone (Fig. 2A). After deglycosylation, two additional 3F4-reactive 20 kDa spots, pIs of 6.5 and 6.1, were seen (Fig. 2B), whereas ICSM 35 detected three major spots at 8.1, 7.7, 7.3, and a minor spot at 6.8 (Fig. 2C). Taken together, the PrPSc core fragment in GSS P102L is composed as follows: (i) PrP35 accounts for two major 6.5 and 6.8 spots, and minor 6.1, 7.3, 7.7, and 8.1 charge isomers; (ii) PrP27-30 contributes a major 7.7 spot, and two 7.3 and 8.1 charge isomers (Fig. 2F). Immunoblot with 3E2 showed additional spots in the acidic zone of the gel, with an orthogonal migration between 20 and 14 kDa (Fig. 2D); after deglycosylation, the PrP27-30 core fragment resolved as six spots with pIs ranging from 6.1 to 6.8 (Fig. 2E), whereas CTFs of 16–17 kDa and 12–14 kDa were confined to a 4.5–6 range (Fig. 2E). The heterogeneous composition of the core fragment in P102L GSS was not dependent on the glycosylation pattern, since the PrPSc core fragment in GSS P102L is characterized by a “diglycosylated dominant” pattern, was composed by a single 6.8 spot (Fig. 2F). Collectively, these results show that divergent allelic PrPSc conformers and glycotyes propagate in GSS P102L. Immunoblots of the detergent-insoluble fraction probed with 3F4 (Fig. 2G) and 3E2 (Fig. 2H), revealed PrPSc species migrating at 20–30 kDa and 8 kDa, in addition to CTFs of 16–17 and 12–14 kDa, thus demonstrating the occurrence of endoproteolytic products, matching those obtained after PK digestion. Graphic representation of the allelic contribution of PrP35 and PrP27-30 to PK-resistant PrPSc is illustrated in the cartoon (Fig. 2I).

Sedimentation velocity in sucrose gradients of mutated and wild-type PrP in P102L GSS

As expected, in control human brains, small aggregates of 3F4- and ICSM 35-reactive PrPSc species were recovered only in the top fractions. On the contrary, in the six GSS cases with PK-resistant PrPSc, 3F4-reactive species were collected in all gradient fractions, being more abundant in fractions 9 to 11; the latter fractions contained also higher Mr PrP aggregates, and, in addition, fraction 11 contained aberrant PrP species of 8–16 kDa (Fig. 3A); fractions stained with mAb ICSM 35 disclosed the presence of small PrP aggregates mainly in the top fractions, in addition to variable amounts of large assemblies in the fractions 9–11, and to barely detectable PrP forms in the intermediate fractions (Fig. 3B). The distribution and association of aberrant PrP species in the two cases lacking PK-resistant PrPSc, namely cases # 3 and # 8, provided divergent results, as to the partitioning of 3F4- and ICSM-positive PrP molecules throughout the fractions. Indeed, 3F4-positive PrP molecules were detectable throughout all fractions, being more abundant in fractions 1–3 and 10–11, in

Figure 1. Molecular characterization of mutant and wild-type protease-resistant PrPSc in GSS P102L. (A) Western blot with the 3F4 antibody of proteinase K-resistant PrPSc in GSS P102L subjects shows the presence of three bands migrating at 29, 25 and 20 kDa (cases 1–2 and 4–7), in addition to variable amounts of an 8 kDa fragment (cases 1–8). (B) Comparative analysis of PrP27-30, before and after glycans removal (Fig. 1B, lanes 5–8), therefore suggesting a common proteolytic cleavage site of glycosylated and unglycosylated PrPSc species. Cerebellar homogenates were available in four P102L subjects and showed the presence of barely detectable amounts of the 8 kDa fragment in three cases, and of PrP27-30 in one.

Immunoblots probed with ICSM 35 antibody, which fails to recognize human PrPM, while detecting PrP35 [24], showed the presence of PK-resistant PrPSc in brains from subjects with PrP27-30 (Fig. 1C). In all cases, ICSM 35 failed to detect PrP6, thus
the absence of higher molecular and 8–16 kDa species (Fig. 3C); on the contrary, ICSM 35-positive PrP aggregates provided a decreasing signal through fractions 1 to 5 and a weak positivity in the bottom fraction (Fig. 3D). PK-treatment in the six cases with PK-resistant PrPSc showed the presence of PrP27-30 in all fractions, more abundantly in those reaching the bottom of gradient; the last two fractions also contained high amount of the 8 kDa fragment (Fig. 3E). In these samples, ICSM 35-positive PrP27-30 was seen in fractions 9–11 (Fig. 3F). In cases lacking PrP27-30, PK-resistant PrP species of 8 kDa were decorated by mAb 3F4 in the bottom fractions (Fig. 3G), whereas ICSM 35-positive species were completely degraded by the protease treatment (Fig. 3H).

Protease-sensitive PrPSc conformers

Using the “mild PK” protocol, or limited digestion with PK at 4°C, in the six cases with PK-resistant PrPSc we detected levels of 3F4-positive sPrPSc higher than amounts of rPrPSc obtained with the classical “harsh-PK” protocol. The signal of the cold-PK resistant isoforms, including a high Mr smear, was greatly enhanced after PTA precipitation, and reduced to a core fragment migrating in a 20–22 kDa zone, after PNGase treatment (Fig. 4A). Immunoblot with ICSM 35 showed that also the wtPrP contributed to the pool of sPrPSc, although at lower levels than the mutated PrP isoform (Fig. 4B). Intriguingly, moderate to high amounts of 3F4-positive sPrPSc species were recovered in the two cases lacking the 21 kDa fragment (Fig. 4C), in addition to ICSM 35-positive high Mr sPrPSc species (Fig. 4D, arrow). In all investigated subjects, sPrPSc species included the 8 kDa fragment, which was detected by the 3F4 antibody, but not the ICSM 35 (Table 2). Collectively, these findings are at variance with results obtained with GSS P101L prions in transgenic mice, which lack PrP8 and have a sPrPSc core fragment of 22–24 kDa [14]. Brain homogenates from frontal cortex of GSS P102L subjects with and without PrP27-30, digested with increasing concentrations of PK, ranging from 5 to 50 μg/ml, and probed with 1E4, did not show the ladder-like pattern observed in subjects with “variably protease-sensitive prionopathy” (VPSP) and GSS A117V [26].

PrP IHC

In all investigated subjects, tissue deposition of PrP occurred as different patterns, including synaptic-like, granular, diffuse, unicentric and multicentric plaques (Fig. 5A–F, Table 3). We also noted a fine granular type of PrP deposition in the white matter, suggestive
of an axonal location. Plaques were labeled by 3F4 (Fig. 5A, 5B, 5C) and ICSM 35 (Fig. 5D, 5E, 5F) antibodies, and, as far as individual plaques could be identified in consecutive paraffin- or plastic-embedded sections (data not shown), we were unable to observe ICSM 35-negative plaques. On the contrary, diffuse and synaptic-like PrP deposits were exclusively labeled by 3F4, but not...
as granular and plaque deposits, or in a wavy motif. This further observed in the white matter with both antibodies. Taken together, glial cell membranes. Additionally, punctuate PrP deposition was revealed the association of fibrillary prion protein aggregates with at codon 129 (H187R-129V haplotype) [27]; IHC with ICSM 35 recently described by Colucci et al. in GSS H187R coupled with V IHC with 3F4 showed synaptic-type positivity and "filamentous" deposition was seen within all cortical layers of the frontal cortex. doi:10.1371/journal.pone.0032382.g004
decreased by PNFase treatment (lane 10).
in the upper part of the gel in a 50–70 kDa zone (lane 9, arrow), partially and 5) or treated (lanes 4 and 6) preparations. (D) In case of the 8 kDa fragment are seen (lane 3–6) in PNGase untreated (lanes 3 NaPTA precipitation (lanes 5 and 6) consistent amounts of PrP27-30 and band (lanes 1,2, arrow); in contrast, after cPK digestion (lanes 3 to 6) and # deglycosylation. (C) Immunoblot with 3F4 of brain extracts from case # 8, after conventional PK treatment, shows a barely detectable 8 kDa band (lanes 1,2, arrow); in contrast, after cPK digestion (lanes 3 to 6) and NaPTA precipitation (lanes 5 and 6) consistent amounts of PrP27-30 and of the 8 kDa fragment are seen (lane 3–6) in PNGase untreated (lanes 3 and 5) or treated (lanes 4 and 6) preparations. (D) In case # 8, immunoblot with ICSM 35 did not reveal PrPSc after cPK treatment (lanes 7 and 8), whereas NaPTA precipitation showed a smear migrating in the upper part of the gel in a 50–70 kDa zone (lane 9, arrow), partially decreased by PNase treatment (lane 10).
doi:10.1371/journal.pone.0032382.g004
ICSM 35 antibody. Noteworthy, the use of antibody directed against the N- and C-terminus disclosed a peripheral staining of plaques (data not shown). PrP immunostaining in case # 2 digested with PK before (lane 1) and after PNGase treatment (lane 2); lanes 3 to 6 show products of "cold" PK (cPK)-treated homogenates, before (lanes 3 and 5) and after (lanes 4 and 6) PNGase treatment; in lanes 5 and 6 samples were NaPTA-precipitated; (B) cPK-treated (lanes 7–10) and NaPTA precipitated products (lanes 9 and 10) were revealed with ICSM 35, prior (lanes 7 and 9) or after deglycosylation. (C) Immunoblot with 3F4 of brain extracts from case # 8, after conventional PK treatment, shows a barely detectable 8 kDa band (lanes 1,2, arrow); in contrast, after cPK digestion (lanes 3 to 6) and NaPTA precipitation (lanes 5 and 6) consistent amounts of PrP27-30 and of the 8 kDa fragment are seen (lane 3–6) in PNGase untreated (lanes 3 and 5) or treated (lanes 4 and 6) preparations. (D) In case # 8, immunoblot with ICSM 35 did not reveal PrPSc after cPK treatment (lanes 7 and 8), whereas NaPTA precipitation showed a smear migrating in the upper part of the gel in a 50–70 kDa zone (lane 9, arrow), partially decreased by PNase treatment (lane 10).

The results reported here show that the heterogeneity of PrPSc species associated with GSS P102L mutation is wider than previously reported and is characterized by different degree of involvement of PrPWM and PrPWI (Fig. 7). Our findings also provide important clues as to the relative contribution of allelic PrP molecules to neuropathological changes and tissue accumulation of PrP and, in turn, on the sensitivity of IHC and PET blot in detecting pathological forms of prion protein.

We show that the composition of rPrPSc forms in GSS P102L is more heterogeneous than previously recognized, since, in addition to PrP27-30 and PrP6, discrete amounts of CTFs with an intermediate migration at 16–17 kDa and 12–14 kDa were found. Using conventional and sensitive protein separation techniques, we confirm the different glycotype between rPrPM and rPrPWt, and, moreover, we show that rPrPM and rPrPWt have distinct conformations. These conclusions are highlighted by the heterogeneous composition of PrP27-30 core fragment, including distinct 3F4- and ICSM 35-positive PrPSc charge isomers, whose characterisation was not possible under conventional Western blot [24]. This hybrid pattern, reflecting the distinct contribution of rPrPM and rPrPWt, was independent from polymorphic codon 129 since it was observed in MM and MV subjects. In addition, the 3F4-positive core fragment was at variance with fCJD E200K, sharing a diglycosylated dominant pattern of PrP27-30 with GSS P102L, therefore ruling out any influence of the glycosylation status. The combined contribution of PrPWM and PrPWI to PrP27-30 species is likely to account for the molecular complexity of pathological prion protein in GSS P102L cases and for its influence into the phenotypic variability throughout propagation of distinct and multiple molecules.

The detection of CTF16-17 in GSS P102L subjects with PrP27-30 speaks in favor of an aberrant endogenous proteolytic pathway of PrPSc, as suggested by their detection in the native insoluble fraction, and provides molecular support to similarities with sCJD with type 1 PrPSc and MM2 subtype [6]. On the contrary, the significance of CTF12-14 is less clear, since these species have been detected in all molecular types of sCJD; their presence in GSS P102L cases is intriguing, given the strict contiguity of the N-terminus of CTF12-14 at amino acid 154–167 with the C terminus of PrP8 at 146–153. Zou et al. previously observed that, in sCJD, CTF12-14 likely originate from a subpopulation of PrPSc distinct

Figure 4. Protease-sensitive PrPSc conformers in GSS P102L brain tissues. (A) Immunoblot with 3F4 of brain homogenate from case # 2 digested with PK before (lane 1) and after PNGase treatment (lane 2); lanes 3 to 6 show products of “cold” PK (cPK)-treated homogenates, before (lanes 3 and 5) and after (lanes 4 and 6) PNGase treatment; in lanes 5 and 6 samples were NaPTA-precipitated; (B) cPK-treated (lanes 7–10) and NaPTA precipitated products (lanes 9 and 10) were revealed with ICSM 35, prior (lanes 7 and 9) or after deglycosylation. (C) Immunoblot with 3F4 of brain extracts from case # 8, after conventional PK treatment, shows a barely detectable 8 kDa band (lanes 1,2, arrow); in contrast, after cPK digestion (lanes 3 to 6) and NaPTA precipitation (lanes 5 and 6) consistent amounts of PrP27-30 and of the 8 kDa fragment are seen (lane 3–6) in PNGase untreated (lanes 3 and 5) or treated (lanes 4 and 6) preparations. (D) In case # 8, immunoblot with ICSM 35 did not reveal PrPSc after cPK treatment (lanes 7 and 8), whereas NaPTA precipitation showed a smear migrating in the upper part of the gel in a 50–70 kDa zone (lane 9, arrow), partially decreased by PNase treatment (lane 10).

ICSM 35 antibody. Noteworthy, the use of antibody directed against the N- and C-terminus disclosed a peripheral staining of plaques (data not shown). PrP immunostaining in case # 3 was indistinguishable from that observed in GSS cases with PrP27-30 (Fig. 5G,5H). Conversely, in case # 8 an atypical pattern of PrP deposition was seen within all cortical layers of the frontal cortex. IHC with 3F4 showed synaptic-type positivity and “filamentous” PrP aggregates in a wavy pattern of deposition (Fig. 5G,5H). In this case, packed PrP deposits were similar to the “curly” pattern recently described by Colucci et al. in GSS H187R coupled with V at codon 129 (H187R-129V haplotype) [27]; IHC with ICSM 35 revealed the association of fibrillary prion protein aggregates with glial cell membranes. Additionally, punctuate PrP deposition was observed in the white matter with both antibodies. Taken together, in the two cases lacking PrP27-30 at immunoblot, deposition of mutated and wt PrP was observed under diverging patterns, either as granular and plaque deposits, or in a wavy motif. This further confirms that sPrPSc is involved into mechanisms of aggregation and tissue deposition.

Discussion

The results reported here show that the heterogeneity of PrPSc species associated with GSS P102L mutation is wider than previously reported and is characterized by different degree of involvement of PrPWM and PrPWI (Fig. 7). Our findings also provide important clues as to the relative contribution of allelic PrP molecules to neuropathological changes and tissue accumulation of PrP and, in turn, on the sensitivity of IHC and PET blot in detecting pathological forms of prion protein.

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Table 2. Allelic derivation and semiquantitative analysis of protease-resistant and protease-sensitive pathological prion protein species in P102L GSS cases.

| Case (code) | Conventional PK treatment (100 µg/ml 37°C) | Cold PK treatment (250 µg/ml 4°C) |
|------------|------------------------------------------|----------------------------------|
|            | PrP27-30 8 kDa | PrP27-30 8 kDa | PrP27-30 8 kDa | PrP27-30 8 kDa |
| 1 (91-466) | +++ | +++ | +/- | - | +++ | +++ | - |
| 2 (92-331) | ++ | ++ | +/- | - | ++ | ++ | +++ | - |
| 3 (96-422) | - | ++ | - | - | ++ | + | - | - |
| 4 (98-196) | ++ | + | +/- | - | ++ | + | +++ | - |
| 5 (98-197) | +++ | + | +/- | - | +++ | + | +++ | - |
| 6 (2004-59) | +++ | + | +/- | - | +++ | + | +++ | - |
| 7 (2005-51) | +++ | ++ | +/- | - | ++ | ++ | -(*) | - |
| 8 (2005-52) | - | ++ | + | - | - | ++ | +/- | - |

(*) in the upper part of the gel a smear is observed (see figure 5D). Scores: – (negative); +/- (faint); + (weak); ++ (moderate); +++ (intense).

from the population which generates PrP27-30 [29]; in addition, due to their inability to detect PrP8 in preparations containing CTF12-14, these authors considered unlikely the possibility that two separate cleavages of the full-length PrPSc could result in the generation of PrP-CTF12-14 and PrP8. The hypothesis that PrP8 and CTF12-14 are generated from different populations of PrPSc is further strengthened by our findings showing the failure to detect CTF12-14 in cases expressing only PrP8. Taken together, our present results support the view that in GSS P102L PrP-CTF12-14, PrP8, CTF16-17, and CTF12-14 species are generated by multiple and distinct conformational pools of protease-resistant PrPSc, likely originating from diverging subsets of protease-resistant molecules. Although the allelic origin of N-terminally truncated fragments in GSS could not be assessed, we argue that CTFs may originate from both PrPM and PrPWT.

Over the past few years, it has been shown that several prion strains harbor protease-resistant and protease-sensitive PrPSc molecules, and also that some human genetic and sporadic conditions are characterized by the exclusive presence of protease-sensitive PrPSc conformers. Further, a number of observations have suggested that molecular species other than protease-resistant PrP may be involved in the pathogenesis of GSS [12,13,14,15]. Hedge et al. first provided evidence that brain tissues from two GSS subjects with the A117V mutation and from Tg mice inoculated with brain homogenates from GSS P102L patients expressing valine or methionine at codon 129, and in the brain of a P102L M129 GSS case [14]. Collectively, the foregoing experiments have provided evidence that in the P102L mutation alternative misfolding pathways take place, inducing the formation of abnormal conformers other than PK-resistant PrPSc, which can revealed by the “cold PK” assay. However, one biochemical feature of these experimental studies is the striking absence of PrP8, either among protease-resistant PrP species or sPrP forms, and the lack of information regarding the allelic derivation of distinct abnormal PrP species.

In this study, the issue of sPrPSc and rPrPSc in P102L was investigated, after the evidence of different pools of PrPSc with variable distribution in sedimentation profile, and diverging resistance to conventional PK treatment. We introduced PK digestion in “cold condition” to preferentially detect sPrPSc species, in parallel to the canonical PK digestion at 37°C which retrieves rPrPSc molecules. Obtained results confirm earlier experimental findings in Tg mice on the presence of significant levels of “cold PK-resistant” PrPSc in all cases. Under the foregoing experimental conditions we confirm that sPrPSc from the mutated and wt alleles generate fragments with molecular weight similar to those of rPrPSc, an issue not investigated by Wadsworth and collaborators [24]. Noteworthy, cold PK sensitive PrP27-30 e PrP8 were found also in cases lacking rPrPSc, and they were highly retrieved by PTA precipitation. The biochemical properties of P102L sPrPSc species were distinct from molecular PrP forms encountered in GSS A117V and in VPSPr [26]. These findings indicate that different human prion diseases are characterized by the generation of multiple conformational quasispecies of sPrPSc [31].

Despite the wide use of PrP IHC in the diagnosis of prion diseases, it is still unclear which molecular forms of PrPSc are detected by IHC after formic acid treatment and hydrolytic
autoclaving, or even after PK treatment; this represents a major limitation in using IHC as a diagnostic tool in prion diseases. It has been argued that autoclaving of tissue sections, treated with formic acid, might destroy only PrPC and sPrPSc (also lost after limited proteolysis with PK), but not rPrPSc that is only denatured. However, this explanation is in contrast with our present IHC and PET blot results, showing PrP deposition in cases lacking rPrPSc species. Therefore, the diagnostic sensitivity of IHC and PET blot as compared with immunoassay, needs to be reassessed in prion diseases, also taking account of the fixation protocol. According to our present results, PrPM contributes to synaptic-type staining, whereas both PrPSc and PrPSc concurred to granular, axonal, and amyloid PrP deposition. Further, PET blot analysis with both 3F4 and ICSM35 showed a comparable staining of PrP plaques, in sections that were stained following PK digestion. To explain the present results we suggest that fixation with formalin may induce protein-protein covalent cross links, owing to reactivity of formaldehyde with side chain moieties of lysyl, arginyl, tyrosyl, aspartyl, histidyl, and seryl residues. These changes might induce sPrPSc to acquire resistance to chemical and physical treatments, including protease digestion. This line of thinking is readily evident when one considers that cases bearing only the internal PrP8 fragment, in the absence of 3F4- and ICSM 35-positive PrP27-30, showed marked deposition of PrPSc. The present findings lend support to results obtained in GSS H187R-129V, showing intense PrP deposition under “curly” plaques in spite of minimal amount of rPrPSc [27], and expand the range of GSS mutations associated with curly-like sPrPSc deposition.

The present findings widen the spectrum of propagating PrPSc isoforms in P102L GSS, thus providing further molecular support to explain the ample phenotypic heterogeneity observed in this inherited condition.

Materials and Methods

Ethics statement
All studies and procedures were done under protocols approved by the Institutional Review Board at Indiana University. The brains of all subjects included in the study were obtained after the families had expressed a strong wish that we study them neuropathologically and carry out research. The brains were harvested with full cooperation of the pathologists involved and following the family’s written informed consent.

Patients and tissue samples
The eight subjects analyzed in the present study were from four unrelated kindred (Table 1). Pedigrees, clinical features, neuropathological and molecular data of 5 cases (91-466, 92-331, 96-422, 98-196, 98-197) have been previously reported. Frozen and formalin-fixed brain tissues from all patients were used for the present study. Control brains were obtained from subjects not affected by prion diseases. The use of stored...
brain tissues in this study was approved by the ethical committee.

Genetic analysis
Analysis of PRNP was performed by using genomic DNA extracted from frozen brain tissue. Amplification of the PRNP coding region was obtained by polymerase chain reaction; the codon 129 genotype and the codon 102 mutation were determined by restriction enzyme analysis, as previously described [9]. PRNP sequence analyses showed the presence of allelic P102L point mutation coupled with methionine at codon 129 in all cases. Five patients were methionine homozygous at PRNP codon 129, whereas three patients were heterozygous.

Immunoblot analysis
Brain samples were homogenized in 9 volumes of lysis buffer (100 mM sodium chloride, 10 mM EDTA, 0.5% Nonidet P-40, 0.5% sodium deoxycholate, 10 mM Tris, pH 7.4) and clarified by centrifugation at 1,000 g for 10 min. The supernatant (S1) was stored at −280°C until use and the pellet discarded. S1 was centrifuged at 100,000 g for 90 min at 4°C. The supernatant (S2) was saved and the pellet was resuspended in lysis buffer and centrifuged as above. The final pellet (P3) was designated as detergent-insoluble, whereas the detergent-soluble fraction was referred to the final combination of S2 and S3. Protease resistance was assayed by incubating sample aliquots containing 40 μg of total protein with 100 μg of proteinase K (Boehringer Mannheim, Germany) per milliliter at 37°C for 60 minutes. For “cold PK” digestion, clarified 10% brain homogenates containing 1% NP-40 were incubated with 250 μg of PK/ml for 1 h on ice and thereafter treated according to the protocol of Tremblay et al.

Table 3. Immunohistochemical analysis of allelic prion protein species in P102L GSS.

| Case (code) | Frontal Cortex | Cerebellum |
|-------------|----------------|------------|
|             | 3F4            | ICSM 35    | 3F4        | ICSM 35 |
| 1 (91-466)  | +++ Multicentric cortical and subcortical plaques; granular cortical staining. Axonal staining (Fig. 3A, 3C, 3G) | +++ As with 3F4, but less intense granular and axonal deposition (Fig. 3B, 3D, 3H) | +++ Amyloid plaques; granular deposition in the granular layer (Fig. 3E) | +++ Granular and axonal staining (Fig. 3F) |
| 2 (92-331)  | +++ Diffuse staining and multicentric plaques | ++ Weaker Reactivity of plaque core, as compared to 3F4 | +++ Diffuse staining and multicentric amyloid plaques | ++ Weaker staining of plaque core |
| 3 (96-422)  | +++ Granular deposition, multicentric amyloid plaques (Fig. 3I) | ++ As with 3F4, but less intense granular and axonal deposition (Fig. 3J) | +++ Amyloid plaques in molecular and granular layers; axonal deposition | ++ Weaker staining of amyloid plaques |
| 4 (98-196)  | N.A.           | N.A.       | N.A.       | N.A.     |
| 5 (98-197)  | N.A.           | N.A.       | N.A.       | N.A.     |
| 6 (2004-59) | +++ Diffuse plaques, granular deposition on the whole cortex. Plaques and axonal staining in the white matter | ++ As with 3F4, except for reduced granular and axonal staining | +++ Numerous multicentric amyloid plaques involving the whole cerebellum. Axonal staining | ++ Granular and axonal deposition |
| 7 (2005-51) | +++ Granular staining and amyloid plaques | ++ As with 3F4, except for the absence of granular staining | N.A.       | N.A.     |
| 8 (2005-52) | +++ “Curly-like” pattern in the cerebral cortex. Intense axonal staining in the white matter (Fig. 3K and 3M) | ++ Diffuse staining (Fig. 3L and 3N) | + Synaptic-type deposition in the molecular layer | – |

N.A. Not available. Scores: – (negative); + (light); ++ (mild); +++ (moderate); ++++ (intense).

Figure 6. Paraffin-embedded tissue-blot analysis of mutant and wild-type PrP. Detection of proteinase K-resistant forms of mutant (A, B) and wild-type (C, D) prion protein in the cerebellum of case #1, as decorated by 3F4 and ICSM 35 antibodies; g denotes the granular layer, and m denotes the molecular layer. doi:10.1371/journal.pone.0032382.g006
Figure 7. Diagram of PK-resistant and PK-sensitive PrPSc molecules detected in GSS with the P102L mutation. PrPSc species originating from the mutant allele are depicted in orange, whereas wild-type PrPSc molecules are depicted in green. PrP8 is an internal PK-resistant fragment spanning residues 74/90 to 146/153. PrP27-30 is the main C-terminal PK-resistant PrP fragment generated by PK cleavage between PrPSc residues 74 and 103. PrP-CTF 16/17 and PrP-CTF 12/13 are smaller PK-resistant fragments whose allelic origin remains undetermined (depicted in blue). doi:10.1371/journal.pone.0032382.g007

Pathologic PrP Isoforms in P102L GSS

PK-resistant PrPSc

PK-sensitive PrPSc

[14]. The reaction was stopped by the addition of protease inhibitor (5 mM phenylmethylsulfonyl fluoride). Phosphotungstate (PTA) precipitation, after cold PK digestion, was done as previously described [32]. For N-deglycosylation, samples were treated with N-glycosidase F (PNGase F) according to the manufacturer’s instructions (Boehringer, Mannheim) for 8 hours at 37°C. Proteins were dissolved in sample buffer (3% SDS, 3% β-mercaptoethanol, 2 mM EDTA, 10% glycerol, 62.5 mM Tris, pH 6.8) and boiled for 5 minutes. An equivalent of 0.4 mg of wet tissue was loaded on 12% SDS-PAGE gels and proteins transferred onto PVDF membrane (Immobilon P, Millipore) for 2 hours at 60 V. Membranes were blocked with 1% non-fat dry milk in TBST (10 mM Tris, 150 mM NaCl, 0.1% Tween-20, pH 7.5) for 1 hour at 37°C and incubated overnight at 4°C with the following anti-human PrP monoclonal antibodies: 3F4 recognizing human PrP residues 106–110 (1:10.000), ICSM 35 whose epitope spans residues 93–105 and specifically detects wild-type but not P102L human PrP, 6H4 directed to residues 144–152 (Pronics, 1:5000), 3E2 directed to human PrP residues 214 to 231 (1:500) (kindly donated by Dr. Capucci), and mouse monoclonal antibody 1E4 against human PrP residues 97–108 (Cell Sciences, Inc., Canton, MA). Blots were developed with an enhanced chemiluminescence system (ECL, Amersham) and PrP was revealed with 3F4 (1:10.000, Dako), 3E2 (1:500) and ICSM 35 (1:1000, D-Gen).

Sedimentation velocity in sucrose gradient

The S1 fractions prepared by centrifugation of 20% brain homogenates at 8000 g for 5 min at 4°C were incubated with an equal volume of 2% Sarkosyl for 30 min on ice. The samples were loaded atop a 10–60% step sucrose gradient and centrifuged at 200,000 g for 1 h at 4°C, as described by Tzaban et al. [33]. Eleven fractions were collected from the top of the tube for Western blot analysis of PrP.

PrP immunohistochemistry

Paraffin sections that were 8 µm thick were deparaffinized, rehydrated in descending graded alcohols, treated with 98 percent formic acid for 20 minutes at room temperature, and autoclaved at 121°C for 10 minutes in 1.5 mM hydrochloric acid. Sections were rinsed and then incubated overnight at 4°C with monoclonal antibody 3F4 (1:500 dilution) and ICSM 35 (1:250). Subsequent antibody detection involved incubation with a biotinylated goat anti-mouse secondary antibody for one hour (1:500 dilution, Vector Laboratories) at room temperature, followed by incubation with the avidin–biotin–peroxidase complex (Vectastain ABC-Elite kit, Vector Laboratories) according to the manufacturer’s instructions. The samples were then stained with 0.06 percent 3,3′-diaminobenzidine as the chromogen and 0.006 percent hydrogen peroxide in 50 mM Tris buffer, pH 7.6.

Paraffin-embedded tissue blot (PET-blot)

Paraffin sections, 7 mm thick, were collected on precut 0.45 µm-pore nitrocellulose membrane and dried for 30 minutes. The sections were thereafter deparaffinized, rehydrated, and treated with 250 µg/ml of proteinase K for 4 h at 37°C. Sections were incubated with monoclonal antibodies 3F4 and ICSM 35. After washing in TBST, incubation with an alkaline phosphatase-coupled rabbit anti-mouse antibody (Dako) was done. The membranes were then adjusted to alkaline pH by exposure to NTM (100 mmol/L Tris-HCl, pH 9.5; 100 mmol/L NaCl, 2% SDS and a trace of bromophenol blue and loaded on a 16% SDS-PAGE. Immunoblotting was performed as described above and PrP was revealed with 3F4 (1:10.000, Dako), 3E2 (1:500) and ICSM 35 (1:1000, D-Gen).
50 mmol/L MgCl₂. The visualization of the antibody reaction was provided by formazan reaction using NBT/BCIP. Blots were evaluated with an Olympus microscope.

Acknowledgments

We would like to thank Dr. Lorenzo Capucci for provision of the 3E2 monoclonal antibody.

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Author Contributions

Conceived and designed the experiments: SM GZ BG. Performed the experiments: MF AF SF MG. Analyzed the data: SM PP GZ BG. Contributed reagents/materials/analysis tools: SM BG. Wrote the paper: SM GZ BG.