Abstract

Different transmissible spongiform encephalopathy (TSE)-associated forms of prion protein (e.g. PrPSc) can vary markedly in ultrastructure and biochemical characteristics, but each is propagated in the host. PrPSc propagation involves conversion from its normal isoform, PrPC, by a seeded or templated polymerization mechanism. Such a mechanism is also the basis of the RT-QuIC and eQuIC prion assays which use recombinant PrP (rPrPSc) as a substrate. These ultra-sensitive detection assays have been developed for TSE prions of several host species and sample tissues, but not for murine models which are central to TSE pathogenesis research. Here we have adapted RT-QuIC and eQuIC to various murine prions and evaluated how seeding activity depends on glycosylphosphatidylinositol (GPI) anchoring and the abundance of amyloid plaques and protease-resistant PrPSc (PrPRes). Scrapie brain dilutions up to $10^{-8}$ and $10^{-13}$ were detected by RT-QuIC and eQuIC, respectively. Comparisons of scrapie-affected wild-type mice and transgenic mice expressing GPI anchorless PrP showed that, although similar concentrations of seeding activity accumulated in brain, the heavily amyloid-laden anchorless mouse tissue seeded more rapid reactions. Next we compared seeding activities in the brains of mice with similar infectivity titers, but widely divergent PrPRes levels. For this purpose we compared the 263K and 139A scrapie strains in transgenic mice expressing P101L PrPSc. Although the brains of 263K-affected mice had little immunoblot-detectable PrPRes, RT-QuIC indicated that seeding activity was comparable to that associated with a high-PrPRes strain, 139A. Thus, in this comparison, RT-QuIC seeding activity correlated more closely with infectivity than with PrPRes levels. We also found that eQuIC, which incorporates a PrPSc immunoprecipitation step, detected seeding activity in plasma from wild-type and anchorless PrP transgenic mice inoculated with 22L, 79A and/or RML scrapie strains. Overall, we conclude that these new mouse-adapted prion seeding assays detect diverse types of PrPSc.

Introduction

Misfolding of cellular prion protein (PrPC) into the scrapie prion protein (PrPSc) isoform is a key event in the pathogenesis of prion disorders [1,2]. PrPSc is the main component of the TSE infectious agent [3–8] and is able to propagate itself by seeding and templating a conformational change in PrPC, a glycosylphosphatidylinositol (GPI)-anchored glycoprotein [2,4,9,10]. Unlike PrPSc, PrPRes tends to be aggregated [11–15], partially resistant to proteases [3,14], rich in beta sheet [16–20], and lacking in native alpha helices [16,18,19].

In the brain PrPSc can accumulate in deposits ranging from large fibrillar amyloid plaques [21–24] to smaller diffuse non-amyloid oligomers [25,26]. Diffuse forms are predominant in many human and animal TSEs. However, PrPSc amyloid is a prominent feature of some genetic human prion diseases such as Gerstmann-Straussler-Scheinker syndrome (GSS) [27] and prion protein cerebral amyloid angiopathy (PrP-CAA) [28]. In numerous TSE types, both amyloid and non-amyloid deposits can be found in the same tissue. However, in scrapie-infected transgenic mice expressing prion protein lacking the glycosylphosphatidylinositol anchor (GPI), PrPSc appears to be exclusively contained in amyloid plaques [29,30]. Both large amyloid fibrils and non-amyloid aggregates of PrPSc are associated with high levels of infectivity [13,29], but smaller non-fibrillar oligomers have been found to have the highest specific infectivity per unit protein with several scrapie strains [13,31]. Nonetheless, the relative contributions of different PrPSc aggregates to prion propagation and TSE pathogenesis in vivo remains unclear.

Protease-resistant PrPSc (PrPRes) is often used as a definitive biological marker for TSE infections, but several studies have shown that infectivity is not always well-correlated with PrPRes level [32–35]. Indeed, infectivity can sometimes be associated with forms of PrPSc that are largely proteinase K (PK)-sensitive (sPrPSc)
RT-QuIC and eQuIC with Mouse Scrapie Strains

RT-QuIC analysis of anchorless PrP (GPI-) brain tissue

To evaluate the seeding activity associated with predominantly amyloid forms of PrPSc, we analyzed the same scrapie strains in transgenic mice that express only GPI-anchorless PrP (GPI- mice) [29,30]. As mentioned above, these mice accumulate PrPSc that, in contrast to the largely non-amyloid diffuse and amorphous accumulations in wild-type mice, appears to be exclusively contained in amyloid fibrils and plaques. By immunoblotting of PK-treated brain homogenates, the levels of PrPRes present in the brains of the GPI- mice that we tested appeared to be less than or comparable to the levels accumulating in WT mice (Figure 3).

However, quantitative immunoblot comparisons of heavily glycosylated, GPI-anchored WT PrPSc with largely unglycosylated, anchorless PrPRes can be difficult due to apparent differences in the binding efficiency and/or immune detection of these types of molecules on blotting membranes [30,48,49] (data not shown). Furthermore, PrPRes levels in individual brains can vary markedly during the prolonged and subtle clinical phase of disease in the hemizygous GPI- mice used in this study. Further complicating matters, a recent study reported that analyses by capture ELISA indicated that GPI- mice can accumulate up to 25–50 fold more PrPSc than wild-type mice when inoculated with the RML or ME7 strains of scrapie [50], a conclusion that has differed markedly from at least some immunoblot-based determinations. In any case, our measurements of seeding activity by end-point dilution RT-QuIC [41] using the moPrPSc 23–231 substrate revealed that hemizygous GPI- mice infected with each scrapie strain had SD50 concentrations that were indistinguishable from their WT counterparts (Figure 2B, dark purple bars). Interestingly, the same dilutions of brain homogenates from the GPI- mice gave much shorter lag phases than those from WT mice (Figures 2A–C). Despite these differences in reaction kinetics, we could not detect any difference between the GPI- and WT-seeded (RML) RT-QuIC products with respect to PK-resistant fragments on SDS-PAGE (Figure 5, lanes 4 & 8). Overall, these data indicate that predominantly amyloid forms of PrPSc have abundant seeding activity and that samples of a given scrapie strain with similar end-point dilutions (i.e. SD50/ml) can seed strikingly different RT-QuIC reaction kinetics (i.e. lag phases) depending on whether the host mouse expresses wild-type or GPI- PrPSc.

Seeding activity in mice with little PrPRes

To determine if prion seeding activity can be detected in hosts with clinical TSE disease but little or no detectable PrPSc, we compared two scrapie strains in knock-in transgenic mice homozygous for P101L PrP (101LL mice) [51]. Inoculation of
the 263K scrapie strain causes TSE disease and high infectivity titers in the brain but little or no PrPRes in these mice as detected by immunoblotting and other assays [35]. In contrast, when these animals are inoculated with the 139A scrapie strain, they accumulate readily detectable amounts of PrPRes [37]. Indeed, our immunoblot-based comparisons indicated that brain synaptosome preparations from the 263K-inoculated mice contained 81-fold less PrPRes than those from 139A-inoculated mice in the clinical phase of disease (Figure 6). Previous work has shown that a majority of the infectivity fractionates with synaptosomes, and that similar titers are found with these two strains ([35]; unpublished data). Despite the large difference in PrPRes levels with these strains, we measured similar levels of seeding activity by end-point dilution RT-QuIC (Figure 7 A, B, E). Moreover, the profile of PK-resistant bands in the RT-QuIC reaction products was also similar between the two strains (Figure 8, lanes 4 & 6). Altogether, the data indicated abundant seeding activity associated with both high-PrPRes and very low-PrPRes TSE strains.

Next we tested the PK-resistance of the seeding activities associated with the 263K and 139A strains in the 101LL transgenic mice (Figure 7 C, D). Synaptosomes were permeabilized with Triton X-100 and treated with 100 μg/mL PK prior to end-point dilution RT-QuIC. Following PK treatment, little PrPRes was present in the 263K synaptosomes, and the expected size shift in banding pattern was observed in 139A synaptosomes (Figure 6). The PK treatment appeared to cause a modest (~4-fold) decrease in the mean SD50/mg brain value for 263K synaptosomes from three separate experiments (Figure 7E, red bars), but this was of minimal statistical significance (p = 0.056). No effect of PK treatment on the mean SD50/mg brain was seen with 139A synaptosomes (Figure 7E, orange bars). Overall, the results suggested that the 263K seeding activity may be somewhat sensitive to PK digestion, but less so than the total synaptosomal PrP content.

eQuIC detection of prion seeding activity in mouse plasma

Because blood plasma contains strong inhibitors of RT-QuIC reactions, we used the eQuIC [44] assay to analyze plasma samples from scrapie-infected mice. For this assay, beads coupled with antibody 15B3 [52] were used to capture prion seeding activity from plasma prior to detection by RT-QuIC [44]. Unexpectedly, in contrast to previous results obtained with RT-QuIC alone, the use of moPrPSen 23–231 substrate with antibody coated beads in the reaction didn’t support efficient PrPRes detection. More optimal reaction conditions were observed using moPrPSen 90–231 as substrate, 300 mM NaCl and 48°C (data not shown). In contrast to the use of this substrate in RT-QuIC as described above, we saw only rare spontaneous ThT-positive responses in negative control reactions under these conditions with beads present in the reaction well (see below). We tested the reaction sensitivity by spiking uninfected mouse plasma with dilutions of brain homogenates from RML-infected mice (Figure 9). We observed positive reactions with dilutions as extreme as 5×10⁻¹⁵ in 0.2 mL of plasma, which contained ~2 a fingers of PrPRes. These results showed that capture of mouse PrPRes with 15B3 antibody allowed the detection of highly diluted mouse seeding activity in plasma and enhanced RT-QuIC sensitivity by ~10⁵.

In an attempt to improve the reaction speed and sensitivity, we also tried adding fresh substrate to the reaction. This step has been helpful in previously described eQuIC assays for hamster and human prions in plasma samples [44]. However, with the murine-adapted eQuIC system [44], we observed only decreased sensitivity following substrate replacement (data not shown). Thus we abandoned the substrate replacement step in subsequent eQuIC assays for murine prions.

We also tested whether eQuIC (without substrate replacement) can detect PrPRes naturally present in the plasma of scrapie-affected mice. Samples were collected in the clinical phase of disease from 9 scrapie-affected WT mice inoculated with, RML or 79A scrapie strains. eQuIC analysis showed that seven of these infected samples gave multiple positive replicate reactions (three with 4/4
Figure 2. RT-QuIC comparison of multiple mouse-adapted scrapie strains. (A), Brain tissues dilutions ($5 \times 10^{-7}$ and $5 \times 10^{-8}$) from WT mice infected with 22L, ME7 and RML scrapie strains were used to seed RT-QuIC reactions containing moPrPSen23–231 substrate. A final concentration of 130 mM NaCl was used for the reaction. The average ThT fluorescence from a set of quadruplicate wells is reported on the vertical axis. (B), RT-QuIC end-point dilution analysis of brain homogenates from WT (light purple bars) and GPI– (dark purple bars) mice infected with 22L, ME7 and RML. Four replicate wells were used for each brain homogenates dilution. The means ± SD of Spearman-Kärber estimates of the SD50/mg brain tissue from three different experiments are shown.

doi:10.1371/journal.pone.0048969.g002

Figure 3. PrPRes levels in brains of GPI– and WT mice infected with multiple mouse-adapted scrapie strains. Normal brain homogenate as well as 22L, RML and ME7-infected brain homogenates were compared by immunoblotting. The sample brain equivalents were loaded into each lane. Lanes 1–2: WT and GPI– NBH undiluted, respectively; Lanes 3–8: WT and GPI– 22L BH undiluted and serially diluted 2-fold and 4-fold; Lanes 9–14: WT and GPI– RML BH undiluted and serially diluted 2-fold and 4-fold; Lanes 15–20: WT and GPI– ME7 BH undiluted and serially diluted 2-fold and 4-fold. A final concentration of 20 μg/mL PK was used to digest brain homogenates. Bands were detected with monoclonal antibody 6D11 as described in materials and methods.

doi:10.1371/journal.pone.0048969.g003
positive replicates, two with 3/4 and two with 2/4) while the two remaining scrapie-affected mice gave 1/4 positive replicates (Figure 10A). In contrast, tests of 4 negative control mice gave 0/4 positive replicates, while 1 negative control specimen gave 1/4 positives, with the latter being an apparent false positive occurring late in the reaction (over 55 h). We also got similarly positive reactions (all 4/4 positive replicates) from plasma samples from clinically affected WT and GPI

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mice inoculated with 22L scrapie (Figure 10B). Collectively, our data showed the ability of the 15B3-based eQuIC to detect a variety of different mouse-adapted scrapie strains endogenous to plasma in the clinical phase of disease.

**Discussion**

Here we demonstrate the in vitro amplified detection of mouse-adapted scrapie strains by RT-QuIC and e-QuIC assay. In general, the use of full-length moPrPSen23–231 and low NaCl concentrations allowed rapid and sensitive mouse seed amplification with a very low incidence of false positive reactions in the RT-QuIC. The truncated moPrPSen90–231 substrate tended to undergo spontaneous (prion seed-independent) conversion in RT-QuIC reactions, but, curiously, did not show this tendency in eQuIC reactions. We speculate that the presence of antibody coated beads and/or the altered kinetics of the eQuIC might...
diminish spontaneous nucleation of moPrPSc<sub>90–231</sub>. By the same token, interactions of moPrP<sub>23–231</sub> with the beads might have slowed the eQuIC reaction rate relative to that observed in the absence of the beads. In any case, the versatility of RT-QuIC and eQuIC is indicated by the sensitive detection of several mouse-adapted scrapie strains with divergent PrP<sup>Sc</sup> characteristics.

Another highly sensitive assay, protein misfolding cyclic amplification (PMCA) [53], has been shown to be capable of amplifying detection of mouse prion strains [54–60], but with extended overall reaction time for optimal sensitivity. For instance, Murayama and colleagues were able to detect Chandler (RML) PrP<sup>Sc</sup> in 10<sup>−11</sup> brain dilution after three rounds of amplification taking >120 hours total [57]. In comparison, we have found that RT-QuIC can detect comparable RML brain dilutions in <40 hours.

Our detection of mouse PrP<sup>Sc</sup> in plasma extends the use of the e-QuIC, which was shown previously to detect prion seeding activity endogenous to hamster plasma or spiked into human plasma [44]. As with the latter studies, our mouse brain homogenate spiking experiments showed that eQuIC was much more sensitive (~100,000 fold) than RT-QuIC alone, allowing detection up to 10<sup>−12</sup>-fold dilutions of TSE brain homogenate spiked into plasma.

In contrast to previous eQuIC studies with vCJD and hamster-adapted scrapie [44], and other studies with PMCA [61], the substrate replacement step in the eQuIC protocol was not helpful. The reason for this difference is not clear. However, one possibility is that the murine seed particles are more fragile or less adherent to the beads or surface of the well than are the analogous particles of other host species. If so, then removing reaction fluid to refresh the substrate may deplete the seeds and nascent seeded products and compromise, rather than enhance, the reaction rate and sensitivity. Another possibility is that, relative to other substrates such as hamster and human rPrP<sup>Sc</sup> [44], mouse

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**Figure 7. PK-sensitivity of seeding activity in synaptosomes from 1011L mice infected with 263K and 139A.**

5 × 10<sup>−6</sup> (10<sup>−5</sup>) to 5 × 10<sup>−11</sup> (10<sup>−10</sup>) dilutions of PK-treated (100 μg/mL) (C, D) or control (-PK) (A, B) detergent permeabilized synaptosomal fractions. 263K (A, C) and 139A (B, D) synaptosomal fractions were seeded into quadruplicate reactions. (E) End-point dilution RT-QuIC analysis of 263K (red bars) and 139A (orange bars) strains. The mean ± SD of Spearman-Kärber estimates of the SD<sub>50</sub>/mg brain tissue from three different experiments are shown.

doi:10.1371/journal.pone.0048969.g007
rPrPsen may more readily adopt, and remain in, a state that is readily susceptible to seeded conversion to amyloid; in that case, its replacement would not accelerate the RT-QuIC reaction rate.

A key goal in TSE diagnostics is detection of prion seeding activity in blood. Here we found that 92% of plasma samples from mice clinically affected with multiple scrapie strains gave clear positive reactions. However, in the remainder of the mice the plasma seeding activity levels appeared to be near the detection limit. This could be due to naturally low plasma PrPSc concentrations, or to the presence of eQuIC inhibitors in plasma. Nevertheless, negative control samples gave no spontaneous conversion of the substrate within 55 h. Further work will be needed to determine if additional gains in sensitivity can be made without increasing the occurrence of false positive reactions.

Our comparison of the WT and GPI− PrP seeds revealed a curious discordance between seed concentration and reaction speed. The reason for the markedly shorter lag phases of RT-QuIC reactions seeded with infected brain from GPI− mice is unclear. Previous work has shown that for a given type of prion seed, lag phases tend to be inversely correlated with seed concentration in RT-QuIC reactions [41,43]. However, end-point dilution QuIC indicated that the seed, or SD50, concentrations in the brains of the GPI− and WT mice that we examined were indistinguishable for a given strain. End-point dilution RT-QuIC should measure primarily the concentration, rather than the relative seeding capacity, of individual seed particles. Clearly, however, PrPSc seed particles can vary widely in size [13,31] and presumably other characteristics such as seeding activity per particle [13]. For example larger particles, such as plaques or bundles of fibrils, could have many more seeding surfaces than individual fibrils, protofilaments, or small oligomeric seeds. Given that PrPSc in GPI− PrP transgenic mice accumulates exclusively in the form of large amyloid fibrils and plaques, we suspect that the average seed particle is larger, with more seeding surfaces, than those in WT brain homogenates. This higher per-particle seeding activity could support faster RT-QuIC kinetics for a given overall seed particle concentration. Alternatively, or additionally, the lack of GPI anchors and/or glycans on the GPI− PrPSc may allow better access of rPrPsen substrate molecules to seeding sites on PrPSc particles, thus improving the rate of conversion per unit seed in the reaction.

Figure 8. Total protein staining of RT-QuIC conversion products from reactions seeded with synaptosomes from 263K- and 139A-infected 101LL mice. RT-QuIC reactions were seeded with 5 × 10⁻⁶ dilutions of synaptosomal fractions. Products were PK digested (+) or not (−) at final concentration of 10 μg/mL and analyzed by SDS-PAGE. The gel was stained with Deep Purple protein stain. Lanes 1,3: no PK and PK treated uninfected products; Lanes 2,4: no PK and PK-treated 263K products. Lanes 5,6: no PK and PK-treated 139A products. The oval indicates the 18 and 19 kDa bands while the bracket represents the 12, 13 and 14 kDa bands in the PK-digested products of the scrapie-seeded reaction.

doi:10.1371/journal.pone.0048969.g008

Figure 9. eQuIC detection of RML PrPSc spiked into mouse plasma without substrate replacement. A 5 × 10⁻⁶ dilution of NBH or 5 × 10⁻⁴ to 5 × 10⁻¹³ dilutions of RML infected brain tissue containing from ~20 pg to 2 ag of PrPRes, respectively, were spiked into 0.2 mL of mouse plasma. PrPSc was immunoprecipitated using 15B3-coated beads and a portion of the beads was used to seed quadruplicate eQuIC reactions. moPrPsen 90–231 was used as a substrate in all reactions. The mean ThT fluorescence of the four replicates is shown.

doi:10.1371/journal.pone.0048969.g009
The use of the 101.LL PrP knock-in transgenic mice allowed us to directly compare, in a single host model, the seeding activities associated with scrapie strains giving high versus unusually low brain levels of PrPRes in the clinical phase of disease. Our observation of similar seed concentrations with the two strains provided evidence that RT-QuIC seeding activity correlates more closely with infectivity levels, which were equivalent, than with PrPRes levels. The seeding activity of the low PrPRes 263K strain, appeared to be marginally more sensitive to PK than that of 139A but neither strain of seed was as sensitive to PK as the vast majority of PrP in the infected brain tissue. Previous work has also failed to identify levels of PK-sensitive PrPSc in this model that could account for the discrepancy between PrPRes and TSE infectivity [35].

Altogether, we have shown that RT-QuIC: 1) allows highly rapid and sensitive detection of murine prion seeds; 2) works with multiple mouse-adapted scrapie strains and types of tissues (e.g. brain, brain fractions, plasma); and 3) detects diverse types of PrPSc with different ultrastructures and protease sensitivities, with seeding activity correlating more closely with infectivity than with PrPRes levels. Given the extensive use of mouse TSE models to elucidate the underlying biological principles of prion transmission and pathogenesis, we predict that there will be many interesting applications of the RT-QuIC and eQuIC assays for mouse-adapted TSE strains.

Materials and Methods

Recombinant prion protein purification

Genes encoding mouse PrP (residues 23 to 231 and 90–231 accession no. M13685) were amplified and ligated into the pET24 and pET41 vector (Novagen), respectively. Hamster-sheep chimeric PrP (Syrian hamster residues 23 to 137 followed by sheep residues 141 to 234 of the R154 Q171 polymorph [accession no. AY907689]) was amplified and ligated into the pET41 vector (EMD Biosciences), and sequences verified. Protein expression and
purification were performed as previously described [41]. Purity of rPrPsen proteins was ~99% as estimated by SDS-PAGE, immunoblotting, and mass spectrometry (data not shown).

Brain tissues homogenate preparation

Wild type C57BL/10 (Prnp+/+) mice and transgenic mice (tg44) expressing only anchorless mouse PrP (GPI mice) were infected with 22L, ME7 and RML (Chandler) scrapie strains and euthanized at a clinical stage of disease by deep isoflurane anesthesia. In the case of the GPI mice, ME7 inoculations were done with homozygous for the transgene (Tg44+/+), while the RML and 22L scrapie inoculations were done in mice hemizygous for the transgene (Tg44+/−). Brain tissues were collected and 10% (w/v) brain homogenates (BH) were prepared as previously described [30]. Unless otherwise indicated, brain tissues were homogenized using glass Dounce homogenizer in nine volumes (10% w/v) of 1X PBS, pH 7.4, with 0.1 mM phenylmethanesulfonylfluoride (PMSF), 1 µg/mL aprotinin and 0.7 µg/mL pepstatin A protease inhibitors (Sigma). Following a 2 min 2000 g clarification spin, the supernatant was collected, aliquotted and stored at −80°C for the transgene (Tg44+/

Production of synaptosomal preparations from 101LL infected mice

Brain tissue was harvested from 101LL mice infected with 139A or hamster 263K scrapie following cull by cervical dislocation at a pre-defined clinical endpoint. Brain tissue from animals with confirmed clinical and pathological disease was homogenized in 0.32 M sucrose at 100 mg/mL (w/v) and clarified by centrifugation at 2000 xg for 10 min at 4°C. Supernatants were transferred to clean centrifuge tubes, and centrifuged at 12,000 xg for 15 min at 4°C. Pellets were washed twice in 0.32 M sucrose before being resuspended in 0.32 M sucrose at 100 mg/mL, wet weight basis equivalent.

Plasma sample préparation

For plasma collections normal and clinical mice were anesthetized with isoflurane and exsanguinated via heart stick. Blood was immediately transferred to a BD Vacutainer (sodium citrate; Becton-Dickinson) tube and mixed gently. Samples were centrifuged at 3000 rpm in a Eppendorf 5415R centrifuge for 15 min. The plasma fraction was transferred to a new tube and stored at −20°C.

RT-QuIC and eQuIC with Mouse Scrapie Strains

RT-QuIC was performed as previously described [41] except for a few modifications. Briefly, 98 µL of fresh RT-QuIC buffer (10 mM phosphate buffer pH 7.4; 130–400 mM NaCl; 0.1 mg/mL rPrPsen; 10 µM Thioflavin T and 10 µM EDTA) were loaded into wells of a black 96-well plate with a clear bottom (Nunc). Reactions were seeded with 2 µL of the BH or synaptosomal fraction dilutions in a final volume of 100 µL (1:50 dilution). All reactions contained 0.002% final concentration of SDS. Plates were sealed (Nalgene Nunc International sealer) and incubated in a BMG Fluostar plate reader at 42°C for the designated period with cycles of 1 min shaking (700 rpm double orbital) and 1 min rest throughout the incubation. ThT fluorescence measurements (450+/−10 nm excitation and 480+/−10 nm emission; bottom read) were taken every 45 minutes.

SD50 calculations

SD50’s were determined by end point dilution RT-QuIC. In brief, for Spearman-Karber analysis [62] a dilution series with at least one dilution giving 100% ThT positive replicates and at least one dilution giving 0% ThT positive replicates was chosen. The dilution giving 50% positive replicates was calculated as described [41].

RT-QuIC products analysis

At the end of the reaction seeded conversion products were recovered from the wells with 0.5% sulphobetaine, treated with 10 µg/mL of PK for 60 min at 37°C, and analyzed by SDS-PAGE. The gel was stained with a total protein stain (Deep Purple, GE Healthcare).

eQuIC: 15B3 coating of magnetic beads

Rat anti-mouse IgM Dynabeads (Invitrogen) were briefly vortexed and 250 µL of beads (1 X 10^8 total beads) were transferred to new tubes for coating. Following incubation on a magnet, bead storage buffer was discarded and the beads washed twice with 5 original suspended bead volumes of coating buffer (0.1% bovine serum albumin in PBS). A final concentration of 0.38 mg/mL of 15B3 antibody (Prionics AG) was used to coat beads in 1 mL of coating buffer. Tubes were incubated with “end-over-end” rotation at room temperature for 2 h. Following three more washes with coating buffer the beads were resuspended in 250 µL coating buffer and stored at 4°C.

eQuIC of plasma samples

e-QuIC was performed as previously described [44], except for a few modifications. Frozen plasma samples were thawed at 37°C and centrifuged at 16000 xg for 1 min. The supernatant was used for 15B3 immunoprecipitation. Pooled normal mouse plasma (InnovoResearch) was used as a scrapie-negative control in all experiment. For spiking experiments, centrifuged pooled normal plasma was combined with dilutions of brain homogenates (the latter totaling ≤4% of the plasma volume) before 15B3 immunoprecipitation step. Forty µL of 15B3 coated beads were used per 0.2 mL of plasma. 15B3-coated beads were first captured from the coating buffer with a magnet, the coating buffer was discarded, and 0.2 mL of Immunoprecipitation buffer (IP, Prionics AG) was added. An equal volume of plasma was added and tubes were incubated with “end-over-end” rotation for 24 h at 37°C. The beads were incubated on the magnet for 2 minutes and plasma-IP buffer mixture was discarded. Beads were washed twice with 500 µL of Wash Buffer (WB, Prionics AG) and beads were resuspended in 10 µL of 1XPBS (pH 7.4). The beads were then combined with 0.05% SDS in PBS (1:1 v/v ratio) and, following incubation at room temperature for 20 min, 5 µL of beads (1:20 dilution in the plate) were added to 95 µL of eQuIC reaction buffer (10 mM PBS pH 7.4, 300 mM NaCl, 0.1 mg/mL rPrPsen, 100 µM ThT, and 10 µM EDTA) in a black 96-well plate with a clear bottom (Nunc). The reaction was incubated in a BMG Fluostar plate reader at 40°C using the same cycles of shake and rest previously described for the RT-QuIC [41].

Western blotting analysis

rPrPsen was detected by immunoblotting. In brief, 10% brain homogenates were digested with 20 µg/mL of protease K for...
References