Prions are infectious agents which cause rapidly lethal neurodegenerative diseases in humans and animals following long, clinically silent incubation periods. They are composed of multichain assembles of misfolded cellular prion protein. While it has long been assumed that prions are themselves neurotoxic, recent development of methods to obtain exceptionally pure prions from mouse brain with maintained strain characteristics, and in which defined structures—paired rod-like double helical fibers—can be definitively correlated with infectivity, allowed a direct test of this assertion. Here we report that while brain homogenates from symptomatic prion-infected mice are highly toxic to cultured neurons, exceptionally pure intact high-titer infectious prions are not directly neurotoxic. We further show that treatment of brain homogenates from prion-infected mice with sodium lauroylsarcosine destroys toxicity without diminishing infectivity. This is consistent with models in which prion propagation and toxicity can be mechanistically uncoupled.

Highly infectious prions are not directly neurotoxic

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Prions are infectious agents composed of polymers of misfolded prion protein which cause fatal brain diseases such as Creutzfeldt-Jakob disease. These diseases involve progressive loss of neuronal cells, and it has been long assumed that prions are directly toxic to cells as they propagate. However, recent studies have suggested that prion infectivity and neurotoxicity can be uncoupled and involve distinct mechanisms. Using highly purified infectious prions we demonstrate that prions are not directly neurotoxic and that toxicity present in infected brain tissue can be distinguished from infectious prions. This has fundamental implications for understanding of prion diseases and how to treat them and may have wide relevance in other neurodegenerative diseases which involve propagation and spread of proteopathic seeds.

Significance

Prions are infectious agents composed of polymers of misfolded prion protein which cause fatal brain diseases such as Creutzfeldt-Jakob disease. These diseases involve progressive loss of neuronal cells, and it has been long assumed that prions are directly toxic to cells as they propagate. However, recent studies have suggested that prion infectivity and neurotoxicity can be uncoupled and involve distinct mechanisms. Using highly purified infectious prions we demonstrate that prions are not directly neurotoxic and that toxicity present in infected brain tissue can be distinguished from infectious prions. This has fundamental implications for understanding of prion diseases and how to treat them and may have wide relevance in other neurodegenerative diseases which involve propagation and spread of proteopathic seeds.

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Competing interest statement: J.C. is a Director and J.C. and J.D.F.W. are shareholders of D-Gen Limited, which supplied one antibody used in the study. This article is a PNAS Direct Submission. This open access article is distributed under Creative Commons Attribution License 4.0 (CC BY).

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We assessed neurotoxicity of pRML in a high-throughput, multiparametric assay consisting of time-course studies and end-point analyses of neuronal morphology. Neurite retraction was measured in phase-contrast mode in a cortico-hippocampal mouse neuronal culture on an IncuCyteS3 live-cell imaging platform followed by immunocytochemistry and high-content image analysis of neurons on an Opera Phenix platform. For immunostaining we used the neuron-specific markers NeuN, MAP2, and spinophilin that detected neuronal nuclei, neurites, and dendritic spines, respectively. We first examined toxicity of RML prion-infected and normal, uninfected, brain homogenate in these assays. Brain homogenate dilutions from clinically sick RML prion-infected wild type (Prnp+/+) mice containing from 104.2 to 105.7 IU/mL showed marked neurotoxicity (Figs. 1–4 and Movies S1 and S2) and, although normal brain homogenate at higher concentrations also showed nonspecific toxic effects, this was never comparable to the extent of toxicity elicited by RML-infected brain homogenate. Indeed, when applied over a range of equivalent concentrations, RML-infected brain homogenate was always significantly more toxic than normal brain homogenate (Figs. 1–4 and SI Appendix, Fig. S2, and Movies S1 and S2). Importantly, no significant prion propagation was detected by ASCA in primary mouse neurons treated with infectious RML brain homogenates for 72 h (SI Appendix, Fig. S1), indicating that prion propagation was not the basis for the observed specific neurotoxicity. In sharp contrast to findings with RML-infected brain homogenate, pRML prions, diluted in tissue culture medium, did not induce neurite retraction even when as much as 105.3 IU/mL was added, equivalent to a 10-fold higher prion titer than that present in the brain homogenate eliciting maximal toxicity (Fig. 2 A and B). We also evaluated neurite fragmentation (Fig. 2C) and dendritic spine density (Fig. 2 D and E) and found that none of the pRML preparations induced toxicity in the cultured neurons at any stage of treatment, again in sharp contrast to RML prion-infected brain homogenate (Fig. 2 and Movie S3).

**Purified RML Prions in Prnp<sup>0/0</sup> Mouse Brain Homogenate Are Not Neurotoxic in Cell Culture.** It is possible that, in order to cause neurotoxicity, infectious prions require cofactors present in mouse brain. Therefore we resuspended pRML prions in brain homogenate from uninfected Prnp<sup>0/0</sup> mice (which are devoid of PrPC and unable to propagate prions) (24, 25). We tested four concentrations of infectious pRML containing 10<sup>4.2</sup> to 10<sup>5.7</sup> IU/mL using the above-described multiparametric assay of neurotoxicity. As with pRML prions in tissue culture media, pRML prions diluted in Prnp<sup>0/0</sup> brain homogenate prior to application to neurons showed no evidence of neurotoxicity as measured by neurite retraction (Fig. 3 A and B, and Movie S4), neurite fragmentation (Fig. 3C), or reduction in spine density (Fig. 3 D and E). Prnp<sup>0/0</sup> brain homogenate itself at a concentration of 10<sup>-4</sup> did not cause toxicity in primary neurons, whereas RML prion-infected brain at the same concentration was toxic (Fig. 3 B–E, and Movies S1 and S5).

**Purified RML Prions in Prnp<sup>+/+</sup> Mouse Brain Homogenate Are Not Neurotoxic in Cell Culture.** We also added pRML to brain homogenate from uninfected wild-type Prnp<sup>+/+</sup> mice to levels of 10<sup>4.2</sup> to 10<sup>5.7</sup> IU/mL and applied it to primary neurons for 3 d as before. As incubation of pRML prions with Prnp<sup>+/+</sup> brain homogenate could conceivably result in generation of prions or putative toxic PrP species in vitro prior to addition to the neuronal culture, we kept incubation time to a minimum and did not exceed

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**Fig. 1.** Development of the IncuCyte-based neurite retraction assay. (A and B) Neurite outgrowth in primary neurons plated at 15,000 cells/well plated after 10 d in vitro (DIV); n = 6. Dark blue traces: neurites; yellow: cell features excluded from neurite analysis. (Scale bar, 200 μm.) (C) Neurite retraction in response to the treatment with RML-infected and uninfected Prnp<sup>+/+</sup> brain diluted to a concentration of 10<sup>-3</sup> (RML, 10<sup>5.2</sup> IU/mL) for 12 h occurred in a cell-density–dependent manner (n = 3, *P < 0.03, unpaired two-tailed t test). (D) Dose-response curve for the neurite retraction assay; n = 3 to 12 independent cell cultures were analyzed in triplicate after 12 h of treatment; mean ± SD. Data were fitted using a nonlinear regression model [Y = 100/(1+10<sup>(LogIC50-X)/HillSlope)</sup>]. ****P < 0.0001, one-way ANOVA with Tukey’s multiple comparison test.
Four concentrations of pRML diluted in nate were tested. We did not observe neurite retraction even at nonspecific toxicity from normal brain at 10⁻³ concentration of normal brain homogenate as vehicle for pRML. Collectively, these data demonstrate apparent, no additional toxicity was elicited by inclusion of high concentrations of pRML. Collectively, these data demonstrate that authentic ex vivo prions are not directly neurotoxic, either when directly added to cell cultures or when first mixed with uninfected brain homogenate to produce a prion titer at or far exceeding the prion titers of brain homogenate dilutions prepared from clinically sick RML prion-infected mice which are highly neurotoxic.

Sarkosyl Abolishes the Toxicity of RML-Infected Brain Homogenate without Diminishing Infectious Prion Titer. In a further series of experiments, and as a prerequisite for undertaking fractionation of the toxic species from RML-infected brain homogenate, we sought to identify a detergent that is compatible with preservation of toxic activity. Based on the knowledge that sarkosyl has no detrimental effects on prion infectivity (20) we examined this detergent first. Remarkably, we found that treatment of 10% (wt/vol) RML brain homogenate with final concentrations of ≥2% (wt/vol)
sarkosyl completely abolished toxicity in primary cultured neurons (Fig. 5 A–C and SI Appendix, Fig. S3) without diminishing the prion infectivity titer in the same samples as measured by the ASCA (Fig. 5D). Indeed, in accordance with previous findings (20), prion infectivity levels in the RML brain homogenates treated with the highest concentrations of sarkosyl were slightly higher than that of untreated RML brain homogenate which might reflect changes in normal brain homogenate (SI Appendix, Fig. S3). Thus, while the specific toxicity of RML brain homogenate is sarkosyl-sensitive, the nonspecific toxicity of normal brain homogenate is not. Indeed, after sarkosyl treatment the residual toxicity of RML brain homogenate becomes equivalent to that produced by the same concentration of normal brain homogenate (SI Appendix, Fig. S3). Collectively, these data with sarkosyl clearly suggest that the structure of the specific neurotoxic active species generated in RML-infected brain is distinct from that of infectious prions.

**Discussion**

The molecular basis of prion neurotoxicity and neurodegeneration remains unknown. Here we report a multiparametric assay of prion neurotoxicity in primary neurons that readily differentiates prion-infected and noninfected mouse brain homogenates. This assay is a combination of kinetic and end-point high-content imaging of neuronal morphology, and it measures neurite retraction, neurite fragmentation, and changes in dendritic spine density upon semi-acute treatment of 10-d-old cortico-hippocampal neuronal cultures in a multiwell plate. The 2 log10 dynamic range of the assay enabled us to undertake a systematic and quantitative investigation of prion toxicity in diseased mouse brains.
Using this multiparametric assay, we examined exceptionally pure preparations of highly infectious, ex vivo prion rods [100 to 200 nm in length (21, 22)] devoid of contaminating proteins (20) or small oligomeric PrP assemblies (21, 22) [akin to those described by Prusiner and colleagues (1, 26)] in cell culture alongside RML prion-infected mouse brain homogenates. We found that prion-infected brain homogenates were toxic as measured by neurite retraction and fragmentation as well as by reduction in dendritic spine density. In sharp contrast, highly purified infectious RML rods correspond to PK-resistant material, RML-infected brain homogenate (BH) comprises both PK-resistant and PK-sensitive prion species (12–14, 20). We found that a 30-min treatment of RML BH with increasing concentrations of sarkosyl abolished neurotoxicity while infectivity was unaffected or even slightly increased. Therefore, neurotoxicity does not seem attributable to infectious prion assemblies regardless of their aggregate size or relative protease sensitivity.

We do not consider these findings as contradictory to earlier publications which have reported neurotoxicity in partially purified prion preparations or crude prion-infected brain homogenate (7, 8), which contains multiple PrP assemblies of different sizes and relative protease resistance, making it impossible to attribute toxicity to a defined PrP species, infectious or otherwise. The lack of detectable direct toxicity of highly purified prion preparations or sarkosyl-treated infected brain homogenate is consistent with models of prion neurotoxicity being mediated by PrPSc, which are distinct from fibrillar infectious prion assemblies and which may be generated by a distinct mechanistic process (3, 4, 14, 15) although further studies are required to confirm the identity of neurotoxic species. In future work, these neurotoxicity assays will be applied to study the kinetics of the generation of neurotoxic species during the incubation period of prion infection and the biochemical properties of neurotoxic species, which should facilitate fractionation of
proprion-infected brain homogenate to isolate the neurotoxic species and allow their identification and structural characterization.

Materials and Methods

All experimental procedures involving RML prions were carried out in microbiological containment level 2 or level 3 facilities with strict adherence to safety protocols and guidelines. Work with animals was performed in accordance with licenses approved and granted by the UK Home Office (Project Licenses 70/6454, 70/7274, and 70/9022) and confirmed to University College London institutional and Animal Research: Reporting of In Vivo Experiments guidelines.

Prion Inoculation and Brain Homogenate Preparation. Mouse-adapted RML prions used in this study were originally supplied by Charles Weissmann, Zurich Institute for Molecular Biology. Two batches of 200 brains from terminally affected intracerebrally inoculated CD1 mice (Hsd:ICR [CD1]; Harlan) were homogenized in D-PBS lacking CaCl2 or MgCl2 ions (Invitrogen) using tissue grinders as described previously (20) to produce two 1-L pools of 10% (wt/vol) RML brain homogenate (designated I13100 and I17700). Control 10% (wt/vol) BH was prepared in D-PBS from the brains of uninfected CD1 (Prnp0/0) mice. All brain homogenates were stored as aliquots at −70°C.

Purification of Prion Rods from Brain Homogenate. Ex vivo infectious RML prion rods were purified from 200-μL aliquots of 10% (wt/vol) of the RML brain homogenates I13100 and I17700 without the use of Proteinase K as described previously (20). The method produces a recovery of ~10% of the prions present in the starting 10% (wt/vol) brain homogenate so that resuspension of the purified P4 pellet fraction in buffer at one-tenth of the volume of the 10% (wt/vol) brain homogenate from which it was derived produces a recovery of the infectivity titer of which is closely similar to that of the starting 10% (wt/vol) brain homogenate (20). Concentration of purified prions or buffer exchange was achieved by centrifugation at 16,000 × g for 30 min and resuspension of the pellet fraction into the desired volume and buffer of choice. Typically, we prepared stock preparations of the P4 pellet fraction resuspended in D-PBS containing 0.1% (wt/vol) sarkosyl in one-fiftieth of the volume of 10% (wt/vol) brain homogenate from which it was derived.

ASCA of RML Infectivity. The infectivity of purified RML prions and RML brain homogenate was determined by ASCA in RML prion-susceptible PK1/2 cells (27) as described (23). In brief, PK1/2 cells were dispensed into 96-well plates at 18,000 cells per well. The next day the cells were infected with prion-containing or mock samples using a Biomek FX liquid handling robot. After 3 d of infection, cells were split 1:8 by Biomek FX and grown to confluence for the next 3 d. After two further 1:8 splits, ~25,000 cells were transferred into each well of activated ELISPOT plates, which were then treated with Proteinase K and developed using anti-PrP monoclonal antibody ICSM18 (D-Gen) and an alkaline-phosphatase-conjugated anti-IgG1 polyclonal secondary antibody (Southern Biotechnology Associates) as described (23). Spot counts reporting PK-resistant PrP-positive cells were determined with a Bioroeder 5000-EF system (BioSys).

Quantification of Prion Titer. Enzyme-linked immunosorbent assay detection of protease-resistant PrP on prion-infected cells on ELISPOT plates results in the appearance of focal PrP “spots” distinct from any nonspecific background immunoreactivity seen on the plate (23, 27). The dose–response is dynamic between ~50 and 1,000 spots per well; however, because the assay is nonlinear, every experiment must include serial dilution of RML prions of known prion titer (intracerebral LD50 units/mL determined from rodent bioassay) to produce a standard curve against which unknown samples can be calibrated. This method reports prion titer in tissue culture infectious units per milliliter, which in this paper for simplicity is abbreviated as “IU/mL.” A stock 10% (wt/vol) RML brain homogenate I8700 used as a reference preparation reported a prion titer of 10−2.2 intracerebral LD50 units/mL when end-point–titrated once in Tg20 mice (20). The infectious titer of the 10% (wt/vol) RML BH stock (I13100) used in toxicity assays and for prion purification was measured in ASCA as described above and was 10−2.9 IU/mL, consistent with previously determined titers of RML BH (14, 20). The infectious titer of four independently purified batches of prion rods (P4 pellets resuspended in D-PBS containing 0.1% [wt/vol] sarkosyl in one-fiftieth of the total volume of 10% [wt/vol] brain homogenate from which the pellets were derived) was evaluated in an ASCA in PK1/2 cells (as...
Working Concentrations of Control and Prion-Infected Brain. RML and uninfected brain dilutions were prepared from 10% (wt/vol) stock homogenates. As an example, a 1:1000 concentration of brain was made by adding 10 μL of 10% (wt/vol) BH to 990-μL cell culture medium (a 1:1000 concentration of brain in tissue culture medium therefore corresponds to 100-fold dilution of 10% [wt/vol] BH). As described above, the 10% (wt/vol) RML BH stock (113100) was added to 1:1,000 dilution of the prion purifications for an infectivity titer of 10^{-2} IU/mL or 10^{-2} I5u/mL brain. To facilitate easy correlation, the values of RML brain concentration and corresponding infectivity titer (IU/mL) are provided here: 10^{-1} (10^{-2}), 10^{-2} (10^{-3}), 3 x 10^{-3} (10^{-4}), 10^{-3} (10^{-5}), 10^{-4} (10^{-6}), and 10^{-5} (10^{-7}).

Detergent Treatment of Brain Homogenates. Sarkosyl (20% [wt/vol], pH 7.7) was prepared from N-lauroylsarcosine (sarkosyl) sodium salt (Sigma catalog no. 61743) in sterile high-purity water. RML BH (10% [wt/vol]) or normal BH (10% [wt/vol]) were adjusted to final concentrations of 0.1 to 4% (wt/vol) sarkosyl in the sample and incubated at 37 °C with shaking at 300 × g in a thermomixer for 30 min. Subsequently, samples were diluted in cell culture medium and immediately analyzed for infectious titer in the ASCA in PK1/2 cells and for toxicity in primary neurons. The highest concentration of detergent in brain homogenate applied to cultured neurons was 0.04% (wt/vol) sarkosyl which had no intrinsic toxic effect on neuronal cells.

Primary neuronal culture. All of the reagents for primary neuronal cultures were from Thermo Fisher Scientific unless indicated otherwise. Primary cortico-hippocampal cultures for prior toxicity studies were prepared from embryonic day 17 FVB mouse brains. Cortices and hippocampi were dissected in chilled dissection medium consisting of calcium- and magnesium-free Hanks Balanced Salt Solution, 10 mM 4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid, 2 mM L-glutamine and 100 μM of penicillin-streptomycin. Brain tissue was next treated with 0.25% (wt/vol) trypsin and 1,000 U of benzonase (VWR) for 15 min at 37 °C and washed twice with dissection medium. Cells were then resuspended in the plating medium consisting of Dulbecco's Modified Eagle's Medium supplemented with 2% (vol/vol) B27, 0.25% (vol/vol) GlutaMAX (Thermo Fisher Scientific) in D-PBS (70 μL/well) and immediately analyzed for infectious titer in the ASCA in PK1/2 cells and for toxicity in primary neurons. The highest concentration of detergent in brain homogenate applied to cultured neurons was 0.04% (wt/vol) sarkosyl which had no intrinsic toxic effect on neuronal cells.

Immunofluorescence. Primary neuronal cultures grown in 96-well plates were fixed in 3.6% (wt/vol) phosphate-buffered formaldehyde (100 μL/well) for 15 min, rinsed with 70 μL of phosphate-buffered saline (PBS) three times, and fixed with 4% (wt/vol) paraformaldehyde for 15 min. Cells were then incubated with 5% (vol/vol) BSA, 1% (vol/vol) PBS, 1% (vol/vol) Triton X100, 3% (wt/vol) bovine serum albumin (BSA) (Sigma), and 5% (vol/vol) horse serum (HS) (Thermo Fisher Scientific) in D-PBS (70 μL/well) and incubated with blocking buffer (3% [wt/vol] BSA and 5% [vol/vol] HS in D-PBS, 70 μL/well) for 1 h at room temperature. Anti-MA2 mouse monoclonal antibody (1:400, Thermo Fisher Scientific, catalog no. M13-1500) and anti-spinophilin/neurabin II rabbit polyclonal antibody (1:500, Millipore, catalog no. 06–852) were added to the cells to visualize neurites and dendritic spines, respectively. These antibodies were applied to cells in antibody buffer (3% [wt/vol] BSA, 1% [vol/vol] HS and 0.1% [vol/vol] Triton X100; 50 μL/well) and left overnight at 4 °C. Cells were then washed (2 × 100 μL D-PBS) and blocked (50 μL/well) with 5% (vol/vol) normal goat serum (Invitrogen) in D-PBS for 45 min. Next, the cells were incubated with secondary antibodies conjugated to Alexa Fluor dyes: goat anti-mouse Alexa 647 (Thermo Fisher Scientific, catalog no. A21235) and goat anti-rabbit Alexa 488 (Thermo Fisher Scientific, cat. No A11010) at a dilution of 1:1,000 in antibody buffer for 1.5 h at room temperature. Cells were then washed (2 × 100 μL D-PBS) and incubated with Alexa Fluor 555–conjugated anti–NeuN mouse antibody to visualize neuronal nuclei (1:500, Millipore, catalog no. MA3773AS, applied to cells in antibody buffer [50 μL/well] for 1 h at room temperature). The cells were washed again (2 × 100 μL D-PBS), and 150 μL of D-PBS was added to each well after the last wash. Then the plates were sealed and stored at 4 °C in the dark. Plates were imaged within 1 wk of staining.

Image Acquisition. Images were acquired either with an IncuCyte S3 (Sartorius) or Opera Phenix (Perkin-Elmer). For IncuCyte imaging, the imaging platform was kept inside a tissue culture incubator at 37 °C and 5% CO2. A total of four to nine images were taken per well with a 20× objective lens in phase contrast every 4 h from the same well coordinates. For Opera Phenix imaging, images of fixed, fluorescently stained cells were acquired with a 40× water immersion objective, and 15 views were taken per well. The laser power and exposure time was optimized for each channel. Flat-field profiles were calculated at imaging.

Statistical Analysis. The average neurite length value from the four to nine images per well acquired by IncuCyte, was found. There were at least three technical replicates per treatment tested in three independent experiments comprising independent cell cultures and independent prion preparations. Neurite length was normalized to the baseline neurite length recorded for ~20 h prior to the treatment (100%) on a per-plate basis. For Columbus image analysis, analysis outputs of each image were averaged between 15 views of each technical replicate. Then, an average was found between all of the technical replicates from each independent test. For neurite fragmentation analysis, raw averages were normalized on a per-plate basis between 0 and 100% where 0% was the medium (negative control) and 100% was the 10% (wt/vol) RML BH diluted 100-fold to a concentration of 10^{-3}. For dendritic spine density analysis, raw averages were normalized on a per-plate basis to obtain a percentage difference from the negative control. The mean and error of normalized values from all independent tests were plotted. Error bar type, statistical test, and P value are stated in figure legends. All statistical analyses were performed in GraphPad Prism 7.

Data Availability. The data that support the findings of this study are available in the article and SI Appendix.

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