Prion strain-dependent tropism is maintained between spleen and granuloma and relies on lymphofollicular structures

Iman Al-Dybiat1, Mohammed Moudjou1, Davy Martin1, Fabienne Reine1, Laetitia Herzog1, Sandrine Truchet1, Patricia Berthon1, Hubert Laude1, Human Rezaei1, Olivier Andréoletti3, Vincent Béringue1 & Pierre Sibille1

In peripherally acquired prion diseases, prions move through several tissues of the infected host, notably in the lymphoid tissue, long before the occurrence of neuroinvasion. Accumulation can even be restricted to the lymphoid tissue without neuroinvasion and clinical disease. Several experimental observations indicated that the presence of differentiated follicular dendritic cells (FDCs) in the lymphoid structures and the strain type are critical determinants of prion extraneural replication. In this context, the report that granulomatous structures apparently devoid of FDCs could support prion replication raised the question of the requirements for prion lymphotropism. The report also raised the possibility that nonlymphoid tissue-tropic prions could actually target these inflammatory structures. To investigate these issues, we examined the capacity of closely related prions, albeit with opposite lymphotropism (or FDC dependency), for establishment in experimentally-induced granuloma in ovine PrP transgenic mice. We found a positive correlation between the prion capacity to accumulate in the lymphoid tissue and granuloma, regardless of the prion detection method used. Surprisingly, we also revealed that the accumulation of prions in granulomas involved lymphoid-like structures associated with the granulomas and containing cells that stain positive for PrP, Mfge-8 but not CD45 that strongly suggest FDCs. These results suggest that the FDC requirement for prion replication in lymphoid/inflammatory tissues may be strain-dependent.

Prions cause neurodegenerative and fatal disorders resulting from the conversion of the host-encoded cellular prion protein PrPc into a pathological misfolded protein termed PrPsc. Human prion diseases can be of genetic, sporadic or infectious origin1. In farm animals, prions have been responsible for the bovine spongiform encephalopathy (BSE) epidemic that arose mainly in Great Britain and western Europe in the 1990s and ultimately adapted to humans as the variant Creutzfeldt-Jacob disease (vCJD)2. Scrapie in sheep and goats and chronic wasting disease (CWD) in cervids, which has recently reached the European countries3, are other forms of prion diseases in animals. Multiple conformers of PrPsc or strains of prions are recognized in the same host species, exhibiting specific incubation periods, a stereotyped neuropathology, and specific tropism for the lymphoid tissue (for a review4). Colonization of secondary lymphoid organs such as spleen, lymph nodes, gut-associated lymphoid tissue and tonsils occurs long before neuroinvasion following peripheral infection by so-called lymphoid tissue tropic (LTT) prions5–7. Such extraneural prion replication has been observed in natural cases of sheep scrapie, CWD, humans infected with vCJD and in many experimental rodent models of prion disease. Lymphoid cell participation in this process has been extensively studied using mice with constitutive or induced deficits in specific cell populations8. A large body of experimental evidence indicates that follicular dendritic cells (FDCs) play a key role in prion replication in lymphoid tissues. FDCs have been identified as the first site of PrPsc accumulation in lymphoid tissues9,10, and the restriction of PrPc expression to FDCs is sufficient to sustain prion...

1VIM, INRA, Université Paris-Saclay, 78350, Jouy-en-Josas, France. 2UMR Infectiologie et Santé Publique, 37380, Nouzilly, France. 3UMR INRA ENVT 1225, Interactions Hôtés Agents Pathogènes, Ecole Nationale Vétérinaire de Toulouse, 23 Chemin des Capelles, 31076, Toulouse, France. Correspondence and requests for materials should be addressed to V.B. (email: vincent.beringue@inra.fr) or P.S. (email: pierre.sibille@inra.fr)
replication in the spleen\(^{11-13}\). The absence or dedifferentiation of FDCs or ablation of PrP\(^C\) expression in FDCs is mostly associated with the absence of prion replication in the spleen\(^{11,12,14,15}\), but not in the lymph nodes\(^{16}\). These contrasted findings suggested that despite their resemblance, there were features that drive striking different behaviors between both lymphoid structures. In addition, tertiary lymphoid organs may also develop under chronic inflammation stimuli and have been described to replicate prion ectopically in structures containing FDC-positive lymphoid follicles\(^{17,18}\).

The recent finding that the prion species barrier is lower in lymphoid tissues than in the central nervous system (CNS)\(^{19}\) and that BSE prions responsible for vCJD can circulate unnoticed in the lymphoid system of exposed humans with a higher prevalence than normally expected\(^{20}\) has raised the need to identify factors that can affect prion replication in lymphoid or inflammatory organs. In this context, the observations of Heikenwalder and colleagues that granulomatous structures devoid of FDCs were able to replicate prions in experimental mouse models of prion diseases was puzzling\(^{21}\), raising the possibility that such structures may exhibit some permissiveness to LTT toward non-LTT prions. To address this issue, we studied the capacity of prions with opposite lymphotropism to accumulate in the spleen and granuloma. We further showed that prions mainly accumulated in lymphofollicular structures and were associated with both FDCs (identified as PrP\(^{+}\), mfge-8\(^+\)CD45\(^−\)stromal cells) and CD68\(^+\)macrophages in the lymphofollicular structures surrounding the granuloma.

Results

Experimental setup and description of the prion strains used. For granuloma induction, ovine PrP transgenic mice (tg338 line\(^{22}\)) were treated with two successive subcutaneous injections of Complete Freund’s Adjuvant (CFA) emulsified in phosphate-buffered saline, approximately 40 days and 20 days before prion challenge (Fig. 1)\(^{21}\). The granulomas were clearly detectable by eye or palpation on the back of the mice from the time of prion inoculation until the terminal stage of disease.

To calibrate the experiment, granulomatous tg338 mice were inoculated with 127 S scrapie prions via the intraperitoneal (ip) route. The 127 S is an LTT prion that induces short, prototypal disease in tg338 mice\(^{23}\). Half of the inoculated mice were euthanized at day 30 post-infection, at a time when PrP\(^{Res}\) is easily detected in the spleen\(^{23}\). The other mice were euthanized at near the terminal disease stage, at day 95.

In the second set of experiments, two closely related prions, derived from the adaptation of classical BSE (C-BSE) and atypical L-type BSE (L-BSE) to ovinized tg338 mice\(^{24}\), were inoculated to granulomatous tg338 mice. Once terminally adapted to the tg338 host, C-BSE and L-BSE prions display similar biochemical and biological features\(^{24}\), except opposite tropism for the lymphoid tissue, as shown in Fig. 2. Western blot analysis for the presence of PrP\(^{Res}\) indicated that the capacity of C-BSE prions to establish in the spleen was lost on serial passage, whereas it was preserved for L-BSE prions (Fig. 2A). At the 4\(^{th}\) passage, all the spleens from L-BSE infected animals were positive for PrP\(^{Res}\) (Fig. 2B). Conversely, the spleens collected from C-BSE infected animals were hardly positive. At the 4\(^{th}\) passage, quantification of infectivity by an incubation time bioassay indicated that the spleens of mice infected with C-BSE prions that were PrP\(^{Res}\) negative were over 10\(^4\)-fold less infectious than the spleens of mice infected with L-BSE prions (Table 1). Worthy of note, spleen and brain material from L-BSE infected mice induced equivalent strain phenotype in reporter tg338 mice with regard to incubation time (Table 1), PrP\(^{Res}\) electrophoretic pattern (Fig. 2C), PrP\(^{Res}\) distribution in the hippocampus (Fig. 2D) and vacuolation distribution in the brain (Fig. 2E). The patterns of PrP\(^{Res}\) and vacuolation distribution were superimposable to that of C-BSE in these mice\(^{24}\). Therefore, there was no divergent evolution of L-BSE prions on passage through spleen compared to brain.

Figure 1. Experimental setup. At 42 and 21 days before prion inoculation, the mice were injected subcutaneously with a PBS/CFA emulsion. At day 0, the mice were inoculated with brain homogenate from terminally ill mice infected with either 127 S scrapie (intraperitoneal inoculation), L- or C-BSE (both by intracerebral inoculation, see results section). For each prion inoculum, brain, granuloma and spleen were collected at various days post-inoculation, and samples were processed for either histology, immunofluorescence, western blot or PMCA/RT-QuIC in vitro prion amplification.
For the inoculation of granulomatous tg338 mice, brain extracts of both L-BSE and C-BSE prions from the 5th passage in tg338 mice (i.e., stabilized L- and C-BSE strains) were used. Intracerebral inoculation was performed because the time to disease after intraperitoneal inoculation would have exceeded the expected resolution time of the granuloma. In addition, at the dose used, intracerebral infection can be considered as both intracerebral and intravenous infection, due to inoculum spillover in the bloodstream. Consequently, this route always targets the spleen in addition to the CNS as efficiently and within the same time window as an intraperitoneal route. A proportion of mice were euthanized at 2 intermediate time points at 1/3 (day 43) and 2/3 (day 85) of the incubation period.
with a previous report conducted in a wild-type mouse model. We next addressed whether prion strain-specific tropism toward the spleen was also maintained in granulomatous structures by inoculating granulomatous tg338 mice with either C-BSE or L-BSE prions. By western blot analysis (Fig. 4A), L-BSE PrPres was detected in spleens from day 43 onwards, and the levels of accumulation increased until the last stage of disease. As expected, C-BSE PrPres was not detected in the spleens of tg338 mice throughout the time course of disease. A parallel situation was found in the granulomas, which remained constantly PrPres-negative, regardless of the sampling time point. In L-BSE-inoculated animals, some granulomas were PrPres-positive, yet their percentages and levels of PrPres accumulation overall were lower than in the spleen, as observed for the 127 S-infected samples: at day 43 dpi, PrPres was found in the granuloma of one L-BSE-inoculated animal (1/4); one sample was also PrPres-positive at 85 dpi (1/4), and only 2/7 showed clear detectable amounts of PrPres. One out of the 4 animals tested showed very weak but positive signals by WB in the brain, and one provided inconsistently positive signals in the granuloma. The PrPres was also detected in both spleen and granuloma of the 127 S-infected animals (Fig. 3B).

Brains, spleens and granulomas were collected and processed for PrPSc detection using either histological or biochemical techniques (detailed in the methods section).

### Time course accumulation of 127S scrapie prions in the granuloma and spleen.

As shown in Fig. 3A, 127 S PrPres was detected by western blot analysis in the spleens of all granulomatous tg338 mice at day 30 post-infection (although one of them showed only a barely detectable signal). At this time point, PrPres was detected neither in the brain nor in the granuloma of the infected animals. At day 95, the three tissues contained detectable amounts of PrPres. One out of the 4 animals tested showed very weak but positive signals by WB in the brain, and one provided inconsistently positive signals in the granuloma. The PrPres was also detected in both spleen and granuloma of the 127 S-infected animals (Fig. 3B).

To improve the detection and titrate the prion-templating activity in the collected tissues, we used the mb-PMCA (miniaturized bead protein misfolding-cyclic amplification) method, which is highly efficient for amplifying 127 S prions. Healthy tg338 brain lysates were seeded at serial 10-fold dilutions of the tissue of interest and subjected to one 48-h single round of mb-PMCA. The amplified products were tested for the presence of PrPres by western blotting. The results are summarized in Fig. 3C. For the brain, PMCA-positive reactions were observed in only one brain at day 30. At day 95, all the brains analyzed showed positive reactions. Based on the limiting dilution values, there was an overall increase in the templating activity by 10^3–10^4-fold (depending on the brains analyzed) during this period. For the spleen, all the samples analyzed were PMCA-positive and at their maximum seeding activity from day 30 onwards, suggesting a plateau of activity, as previously described. For the granulomas, 3 out of the seven tested samples were PMCA-positive at day 30, with limiting dilution values approaching that found for the spleen, thus suggesting a substantial replicative activity in this tissue. At day 95, all the granulomas showed a positive score with limiting dilution values ranging from 10^-4 to 10^-8 (see supplementary table for percentages of PMCA-positive samples).

Collectively, these data indicated that intraperitoneally inoculated 127 S prions could be detected in a proportion of the granulomas, as early as in the spleen. The positive samples exhibited titers comparable to those observed in the spleen. This difference in the number of affected granuloma versus spleen was fully consistent with a previous report conducted in a wild-type mouse model.

| tissue   | Dilution | L-BSE | C-BSE |
|----------|----------|-------|-------|
|          | n/n0     | Mean survival time* (days ± sem) | n/n0 | Mean survival time* (days ± sem) |
| spleen   | 1/1      | 6/6   | 144 ± 2^a | 4/4 | 229 ± 4^a |
|          | 1/1      | 6/6   | 142 ± 2^a | 6/6 | 138 ± 1   |
|          | 10-4     | 6/6   | na        | 6/6 | 176 ± 3   |
| brain    | 10-5     | 6/6   | 261 ± 16  | 6/6 | 217 ± 9^a |
|          | 10-6     | 6/6   | 281,307   | 2/6 | 286,534   |
|          | 10-7     | 0/6   | >660      | 0/6 | >660      |

Table 1. Mean survival time of tg338 reporter mice inoculated with spleen material from tg338 mice infected with L-BSE or C-BSE. *Mean survival time of tg338 reporter mice inoculated with spleen material from tg338 infected with L-BSE or C-BSE. For pair comparison: this indicates that spleens of L-BSE infected animal are 1000 to 10000-fold more infectious than those of C-BSE infected animals. The symbols indicate inoculum dilution factors for which the incubation times (i.e. the inoculum concentration) are similar.
Figure 3. Detection of PrPSc in the brain, spleen and granuloma of 127S-infected mice. At various times post-infection, groups of animals were sacrificed and brain, spleen and granuloma were removed for tissue processing and analysis. (A) Proteinase-K digestion followed by western blot analysis using biotinylated Sha31 anti-PrP antibody. Note that the exposure times for brain and spleen WB membranes ranged from 3 to 5 minutes, while the granuloma samples required longer exposure times (over 3 hours). (B) Immunofluorescence (IF) detection of PrPSc in the spleen or granuloma of tg338 mice inoculated with 127S prions. Labeling of PrPSc in spleens (B-1, B-3) and granulomas lymphofollicular structures (B-2, B-4) from control (B-1, B-2) or infected animals (B-3, B-4). Ten-micrometer-thick slices were cut on a cryostat, fixed in 5% paraformaldehyde, permeabilized in 0.5% X-100 Triton, and treated with 3.5 N guanidinium thiocyanate before the avidin-biotin blocking reaction followed by biotinylated Sha31 incubation and streptavidin-Alexa-488 labeling, slide mounting using fluoromount and acquisition under a mono CCD camera. Images in false colors: blue, DAPI counterstaining; green; biotinylated Sha31, labeling guanidinium-resistant PrPSc. Bars: 20 μm. (C)
The collected tissues were then analyzed by mb-PMCA to further demonstrate the distinct tropism of L-BSE and C-BSE prions for granulomatous structures. Lowly-diluted (10⁻⁴ dilutions representing the lowest dilution achievable) spleen and granuloma extracts from C-BSE-infected mice did not produce PrPSc-positive amplicons by PMCA amplification (Fig. 4C). In contrast, PMCA-amplified products seeded with spleen and granuloma extracts from L-BSE-infected mice were mostly PrPSc-positive from day 43 onwards. However, the efficacy of the mb-PMCA assay to amplify C-BSE vs L-BSE prions appeared discrepant compared to the bioassay. The limiting dilution values achieved with C-BSE brain material were constantly >100-fold lower than with L-BSE brain material, despite similar values by bioassay in indicator tg338 mice (Table 1). Thus, we refined the mb-PMCA analysis with the data obtained using the RT-QuIC (Real Time Quaking-Induced Conversion) assay to accurately and quantitatively detect the replicative activity of L-BSE and C-BSE prions in central and peripheral tissues. Indeed, using this assay, the limiting dilution values from C-BSE and L-BSE brain materials were much closer throughout the time course of the disease (Fig. 4C). By RT-QuIC, C-BSE granulomas remained negative. A few spleens appeared to be positive above the detection threshold (10⁻⁴ dilution), in particular at 43 dpi (Fig. 4C). The number of positive samples and the limiting dilution values decreased over time. Such low values were found in PrP knock-out mice inoculated with C-BSE and L-BSE prions (Fig. 4C, open symbols), which suggested that we were detecting the residual fraction of the inoculum that had spilled over into the blood and become trapped in the spleen.

RT-QuIC, like PMCA, confirmed that a proportion of L-BSE granulomas were positive from day 43 onwards (Fig. 4C). The proportion of PMCA (or RT-QuIC)-positive granulomas increased with time, ranging from 66% of positive animals at day 43 to 75% (50% with RT-QuIC) at 85 dpi and 100% (57%) at 132 dpi (see supplementary table for percentages of PMCA- and RT-QuIC-positive samples). The templating activity of the positive granulomas was highly variable, with limiting dilution values ranging from 10⁻² to 10⁻⁵. The number of positive granulomas and the templating activity were significantly higher than those found in PrP0/0 mice, suggesting the active replication of L-BSE prions in this structure.

Collectively, our results indicated that prion strain-dependent tropism for the lymphoid tissue was preserved in granulomatous structures, as observed in the spleen.

**PrPSc deposits mainly co-localize with CD68⁺ macrophages and C4⁺ or mfi-g-8⁺ cells in the granuloma core and the surrounding lymphofollicular structures.** While evaluating the structural integrity of the granulomas isolated from the infected mice, we frequently noticed the presence of ovoid adjacent structures filled with small cells and a high nuclear density, observable by hematoxylin/eosin staining of a granuloma sampled from a 127 S-inoculated mouse (see Fig. 5 and Supplementary Fig. 1 for an overview of the main immunofluorescence staining pattern using B, T lymphocytes, macrophages and FDC markers, B220, CD68 and mfi-g-8 or C2/C4 antigens staining, respectively). Overall, the granulomas lymphofollicular structures were highly reminiscent of the inflammatory tertiary follicles that develop in the surroundings of chronic inflammatory stimuli (see38 and39 for a review). Interestingly, the strongest specific PrPSc-positive signal was located in these lymphofollicular structures (Fig. 5C, white arrowhead), while no specific staining could be observed in uninfected control animals (Fig. 5D). This particular staining was much more intense than that within the regular granuloma structure, which was particularly difficult to discriminate from the uninfected animals due to the high background levels.

Since the FDC counts in the lymphofollicular structures surrounding the granuloma were not significantly different from those in the spleen (see Supplementary Fig. 2), we next aimed to identify the cell types supporting prion accumulation in these lymphofollicular structures. Spleens were used as controls. Using consecutive slides, we could observe that FDC-M1 antibody also stained cells in close vicinity to the PrPSc-positive cells (data not shown). The antigen recognized by this antibody was seriously damaged by guanidinium treatment, thus precluding the use of FDC-M1 to establish a formal association of PrPSc and FDCs with a double-staining procedure. Therefore, histological slices from granulomas and spleens collected from mice at various time points post-infection were stained with biotinylated anti-PrP Sha31 and antibodies against the following cellular markers: B cells (B220), macrophages (CD68) and FDCs (C4 complement marker, recognized by FDC-M2 antibody). As anticipated based on a previous report40, PrPSc accumulation in the spleen was mainly located within the germinal center light zone. B220 labeling did not colocalize with the PrPSc signal in the spleen (not shown), and co-staining was also unnoticeable in the granuloma (Fig. 6, panels A, 127 S infected samples; panels E, L-BSE-infected samples) while C4 as well as CD68 staining were superimposed on the 127 S- (Fig. 6, CD68, panels B; C4, panels C) or L-BSE-PrPSc signals (Fig. 6, C4, panels D; C4, panels G). Z-stacks were generated for the CD68, B220 and C4-labeled slides (Fig. 6, panels A-B, B2- and C-2) and confirmed that CD68⁺ and C2/C4⁺ cells were close to the 127 S-PrPSc signals, while B220⁺ cells were clearly not associated with Sha31 positive staining in the granuloma of 127 S-infected animals. Similarly, multiple labeling studies revealed that L-BSE PrPSc staining in the lymphofollicular structures surrounding the granuloma was associated with CD68⁺ or C2/C4⁺ cells, but not with B220⁺ cells (Fig. 6, panels E-1 to G-2).

Considering that the conditions were not optimal for the rigorous characterization of supposed FDCs in the granulomas (guanidinium treatment, non-specific FDC-M2 antibody, etc), we needed to get further convinced that the PrPSc/mfi-g-8⁺ or PrPSc/C2/C4⁺ cells were indeed follicular dendritic cells (ie differentiated stromal...
Figure 4. Detection of PrPSc in the brain, spleen and granuloma of L-BSE or C-BSE-infected mice. At various times post-infection, groups of animals were sacrificed and brain, spleen and granuloma were removed for tissue processing and analysis. (A) Proteinase-K digestion followed by western blot analysis using biotinylated Sha31 anti-PrP antibody. Note that the exposure times for brain and spleen WB membranes ranged from 3 to 5 minutes, while the granuloma samples required longer exposure times (over 3 hours). (B) Immunofluorescence (IF) detection of PrPSc in the spleen or granuloma of tg338 mice inoculated with L- and C-BSE prions. Labeling of PrPSc in spleens (B-1, B-3, B-5) and granulomas lymphofollicular structures (B-2, B-4, B-6) from control (B-1, B-2), L-BSE-infected animals (B-3, B-4) or C-BSE-infected animals (B-5, B-6); (C) PMCA and RT-QuIC templating activity of PrPSc present in individual brains, black ■; spleens, red ●; and granulomas, blue ▲, from C-BSE or L-BSE-inoculated mice. Gray squares represent the inoculum titer. Open symbols represent PrP0/0 animals. See Supplementary Table 1 for a percentage of positive samples.
cells), not just macrophages (ie lymphoid cells) that could have loaded prion particles and mfge-8 antigens as well. For that purpose, a general lymphoid marker should discriminate between stromal and lymphoid cells. Therefore, spleen and granuloma sections sampled from a 127 S-infected animal were triple-labeled with antibodies recognizing CD45, mfge-8 and PrP. Preservation of mfge-8 antigen precluded the use of guanidinium isothiocyanate, so PrP labeling mainly identified the cellular antigen of the prion protein, not the misfolded conformer. The resulting tri-labeled slides were observed under a confocal inverted microscope (Fig. 7). As expected from the literature and our double-label study, we were able to detect PrPC positive cells decorated with mfge-8 in the germinal centers of the spleen. These cells did not stain for CD45, as for true conventional splenic FDCs. When granuloma sections were studied, some cells positive for PrP and mfge-8 were also stained by CD45, establishing that they belonged to the lymphoid lineage. But we also could detect PrPC+/mfge-8+ cells that did not stain positive for CD45 (Fig. 7 panel 2). This finding strengthened the odds that there were indeed true FDCs in the granulomatous environment. Both types of cells could be observed in the same microscope field (see Supplementary Fig. 3A). Spatial fluorophore distribution within the tissue was further investigated (see Supplementary Fig. 3B). These observations confirmed that both PrP and mfge-8 labeling could be observed in the absence of neighboring CD45 (see individual and merged channels in yellow-boxed of Supplementary Fig. 3b showing slicing along the yellow line in both spleen and granuloma). Surprisingly, triple-labeled cells could hardly be detected in the spleen of the infected mice: at least, the mfge-8+/CD45+ cells did not stain for PrP.

In conclusion, both the spleen and granuloma contained macrophages and FDCs, with which misfolded PrP could be associated when mice were inoculated with lymphoid tissue tropic prions, either 127 S scrapie or L-BSE adapted to ovinized mice. Furthermore, we observed that the cellular organization surrounding the granuloma was reminiscent of tertiary lymphoid organs. These structures displayed higher PrPSc signals than the granuloma core.

**Discussion**

Here we provide evidence that the strain-dependent prion LTT is mostly preserved between the spleen and inflammatory, granulomatous structures from infected mice. Both LTT prions studied herein replicated in the spleen and in the granuloma of the infected animals, while the non-LTT prion did not. We further showed that prion accumulation in the granuloma might preferentially proceed through adjacent lymphofollicular structures.

![Figure 5](https://www.nature.com/scientificreports/structure.png)
suggesting that the lymph node environment is much more adapted to the prion accumulation/replication than the regular inflammatory granuloma structure.

Certain prions can actively replicate in the lymphoid tissue of infected animals. Although this step is not mandatory, it allows their efficient routing to the CNS after extraneural infection. By contrast, other prions are not detected within the lymphoreticular system, although we can reasonably suspect that they nevertheless circulate at very low levels through these structures. The determinants of prion tropism toward specific brain regions and lymphoid tissue are mostly unknown but are strain-specific. The adaptation of atypical L-BSE prions to ovine PrPtg338 mice led to the individualization of a prion closely resembling classical C-BSE prions in terms of incubation time, PrPres electrophoretic profile and resistance to proteolytic digestion, neuroanatomical deposition of PrPres and vacuolation, and infectious titer. Despite these phenotypic similarities, the two agents showed marked dissimilar tropisms for the lymphoid tissue, with the C-BSE agent losing its ability to propagate efficiently in the spleen upon iterative passage. Beyond suggesting that minimal conformational differences between PrPSc conformers (which were not possible to observe by any available conventional strain typing method, such as guanidinium denaturation or proteolytic digestion) could result in strikingly different lymphotropism, these two agents represented an ideal experimental paradigm to further study host requirements for lymphoid replication.

Within lymphoid tissues, FDCs are believed to be the main site of prion replication. When a former publication found that prions could accumulate in granuloma supposedly devoid of FDCs, the question arose concerning whether those structures would share the same kind of prion strain-dependent selectiveness with the spleen despite an apparently distinct prion-permissive cell type. We induced subcutaneous granulomas in tg338 mice and challenged them with prion strains differing in their lymphoid tropism. The time course of spleen and granuloma contamination was superimposable for the LTT 127S scrapie prion and the other LTT agent L-BSE. However, both strains replicated at lower and more variable levels in the granuloma than in the spleen. In marked contrast, the non-LTT C-BSE prion was not detected in any of these structures, even after in vitro prion amplification by mb-PMCA. Using RT-QuIC as another cell-free assay with superior sensitivity for C-BSE prions, a few granulomas and spleens provided positive scores, yet the proportion and limiting values were consistent with the inoculum remanence, as evidenced in PrP0/0 mice. There was therefore a 3-4 orders of magnitude difference in splenic or granuloma prion titers (obtained using either PMCA or RT-QuIC) between L- and C-BSE despite a similar titer in the brain. Thus, the L-BSE that was adapted to ovineized mice was a true LTT prion, while the classical C-BSE was not.

Figure 6. Double immunofluorescent labeling of lymphofollicular structures in granuloma from 127 S- (panels A-D) or L-BSE- (panels E-H) -infected animals. Ten-micrometer-thick slices were cut on a cryostat, fixed in 5% PFA, permeabilized in 0.5% X-100 Triton, and treated with 3.5 M guanidinium thiocyanate, followed by the avidin-biotin blocking reaction. Incubation was performed with biotinylated Sha31 (PrPSc, green) + anti-B220 (panels A and E), anti-CD68 (panels B and F) or anti-C4 (FDC-M2, panels C and G) (red) antibodies, followed by anti-rat-Cy3 streptavidin-Alexa-488 conjugates, slide mounting using fluoromount and acquisition under a mono CCD camera. Panels D and H correspond to spleen and granuloma sections stained using IgG2a isotype control antibody. Images in false colors: blue, DAPI counterstaining. Bars: 10 µm. Longitudinal panels (A-2 – G-2) represent Z reslicing alongside yellow lines drawn in panels A-1 to G-1, respectively.
Figure 7. Triple immunofluorescence labeling of Spleen (panels 1) and Granuloma sections (panels 2) from 127S-infected animals. Nine-micrometer-thick slices were cut on a cryostat, fixed in 5% PFA, permeabilized in 0.5% X-100 Triton, blocked with 5% Bovine serum albumin and 2% rat and mouse sera, followed by the avidin-biotin blocking reaction. Incubation was performed with antibodies against CD45 (rabbit anti CD45 serum, gray, panels A) PrP (biotinylated Sha31, green, panels B), and mfge-8 (FDC-M1 rat monoclonal antibody, red, panels C). Incubation was followed with, anti-rat-Cy3 + streptavidin-Alexa-488 + anti-rabbit-Alexa 645 labeling, slide mounting using fluoromount and acquisition under a mono CCD camera. Panels D1 and D2 show the merged image built with the 4 channels (images in false colors: blue, DAPI counterstaining). Panels E1 and E2 correspond to spleen and granuloma sections stained using IgG2c isotype control and secondary antibodies as staining control. Bars: 10µm. Yellow arrows show PrP+/mfge-8+ cells that do not stain positive for CD45.
fully suitable environments for the detection of prions. qRT-PCR experiments supported the view that the local cellular granulomatous environment did not provide the appropriate cytokines (lymphotoxins-α, -β, TNF, CXCL13, CCL19, CCL21, CxCR4 and CxCR5) required for lymph node development and maintenance of FDCs. Macrophages have been described to play a particularly important role in the spleen of infected animals, where they can capture and sequester a significant amount of PrPSc. Here, CD68+ cells were found in the general core of the granuloma, and these cells were also positive for PrPSc. This observation indicated that the main granuloma PrPSc positive signal could be due to macrophage scavenging activity. Both LTT prions used in this study (127 S and L-BSE) could be detected in the CD68+ macrophages of the granuloma core, but that was not the case with C-BSE prion. This observation suggested that the scavenging activity of the macrophage is somehow associated to the local production or accumulation of PrPSc in the granuloma.

Surprisingly, our data established that lympho-follicular structures could develop adjacent to the canonical granulomatous structure that we induced in mice following Freund’s adjuvant injection. These structures contained small cells with a highly reduced cytoplasmic compartment. In addition, immunofluorescence labeling identified B220+, CD68+ and mfge-8+ and C2/C4+ cells. We were also able to observe PrPSc deposition, which could additionally co-localize with CD68+ cells, but not with B220 antigen. Though not specifically complete for FDCs, mfge-8 and C2/C4 antigens are good markers and commonly used for the identification of these stromal cells in the lympho-follicular environment. Although a combination of these markers with several other markers of stromal origin (CD54, CD106, podoplanin, clusterin and PrP) and see (for a review), they are the most specific markers for the identification of FDCs in the context of IF or IHC staining, especially if they are combined with the use of CD45 as an exclusion marker for the Stromal/Lymphoid cells discrimination. A definitive element in favor of the presence of true FDCs would however require the experimental presentation of immune complexes following peripheral or iv administration of fluorescent antigens.

Granulomas induced with complete Freund’s adjuvant seemed to induce the ectopic development of nearby lymphofollicular structures. However, these structures were not observed in every microscopic section studied, which might be. This is to be related to the fact that PK-resistant PrPSc signals and prion seeding activity were absent in a number of these structures throughout the disease pathogenesis of 127 S or L-BSE prions. These negative granulomas could correspond to dissected areas that were sampled without the surrounding lympho-follicular structures. Unfortunately, we could not formally link the prion-positive status of the granuloma with these ectopic structures, since each sample could not be used for both immunohistochemistry and western blot/cell-free assays. In addition, IF prion detection in the case of L-BSE was difficult to confirm in the granuloma core. Therefore, it would be interesting to focus on the distinct cellular organization of the granuloma structure in relation to the prion strains: infections with RML and 127 S scrapie strains (respectively used in and the present study) resulted in massive granuloma staining, while L-BSE prion detection was apparently more restricted to lymphoid-like structures. Consequently, lymphocompetence should be regarded as a continuous criterion, particularly in an inflammatory context. This phenomenon is of public health importance when considering vCJD as a lymphocompetent prion that can also replicate in various organs and benefit from favorable conditions in the context of concurrent chronic inflammatory diseases.

Materials and Methods
Ethics statement. All animal experiments were performed in strict accordance with EU directive 2010/206/EC and were approved by the local ethics committees of the authors’ institutions (Comité d’éthique appliqué à l’expérience animal, Comethea, permit number 12/034 and 12/127).

Transgenic mouse lines and prion strains. The ovine PrP tg338 mouse line (Val136Arg154Gln171 allele) is homozygous and overexpresses approximately eight-fold the heterologous PrP on a mouse PrP null background. PrP0/0 mice were kindly provided by C. Weissmann (Zürich I line). Mice were maintained under specific pathogen-free (SPF) conditions until the infection or inoculation procedures. Then, the mice were housed in a biosafety level 3 laboratory.

The 127 S scrapie prion strain was obtained through serial transmission and subsequent biological cloning by limiting dilutions of the PG127 field scrapie isolate from Tg338 mice. The 127 S infectious titer was 10⁵⁰ LD₅₀/g of tg338 brain pools of 127 S sick tg338 mouse brains were prepared as 20% (wt/vol) homogenates in 5% glucose using a tissue homogenizer (Precellys 24 Ribolyzer, Ozyme, Bertin technologies, France).

L-BSE and C-BSE prions were obtained through serial transmission (5 passages) of cattle classical and atypical L-type BSE in tg338 mice (see Table 1).

Induction of granulomatous structures in tg338 mice and prion infection. Tg338 mice were inoculated subcutaneously with 100 µL of 50% Complete Freund’s adjuvant CFA according to a previously described protocol. The injection was performed either on the abdominal or on the dorsal side of mice and was repeated twice in 21 days between each injection. Visible swelling was observed or detected by eye or by local palpation of the injection site.

The granulomatous tg338 mice were inoculated intraperitoneally with 100 µL of a 2% (w/v) 127 S brain homogenate or intracerebrally with 20 µL of a 10% (w/v) C-BSE or L-BSE brain homogenate. PrP0/0 mice were inoculated with the same dose. Uninfected control mice were intracerebrally inoculated with the same dose of uninfected brain homogenate. Brains, spleens and granulomas were harvested at selected time-points post-infection, at 30 and 95 dpi for 127 S and at 43, 85 and 132 dpi for L-BSE and C-BSE prions, for which the incubation times were longer than for the former scrapie strain. For PrPSc detection by immunoblot, PMCA amplification and RT-QuIC analysis, the collected tissues were homogenized with a tissue homogenizer (Precellys 24 Ribolyzer, Ozyme, Bertin technologies, France) and adjusted to 20% (w/v) in 5% glucose (alternatively, 10% homogenized suspensions were realized for spleen samples or when the granulomas were too small). The samples...
were stored at −80 °C. For immunofluorescence studies, spleens and granulomas were immediately embedded in O.C.T medium (Fontenay-sous-Bois, France) and snap frozen on dry ice before further storage at −80 °C. For histological studies, spleen, granuloma and brain samples were fixed in 4% formalin solution before embedding in paraffin.

**PK treatments for PrP<sup>res</sup> detection.** Brain, spleen and granuloma homogenized samples were treated with PK (final concentration of 115 μg/mL) for 90 min at 37 °C. Loading Buffer (2 × 50 mM Tris, pH 6.8, 2% SDS, 0.01 bromophenol blue, 2 mM dithiothreitol 50% glycerol) was added, and the samples were denatured at 100 °C for 5 minutes.

**Miniaturized bead-Protein Misfolding Cyclic Amplification (mb-PMCA).** The mb-PMCA procedure has been described in a previous work<sup>35</sup>. In brief, mouse brain lysate from uninfected tg338 mice (10% brain lysate in cold PMCA buffer: 50 mM Tris-HCl pH 7.4, 5 mM EDTA, 300 mM NaCl, 1% Triton-X-100) were used as substrate for 127S scrapie, L-BSE and C-BSE prion strains. PMCA was performed in a final volume of 36 μL of lysate per well in 96-well PCR microplates (Axygen, Union City, CA, USA). Each well was first filled with one 2.381-mm-diameter Teflon bead (Marteau et Lemarié, Pantin, France). A 4-μL aliquot was serially diluted in the 36 μL-containing wells. The microplate was sealed and placed on a plexiglass rack designated for the cup horn of the Q700 sonicator (Misonix, Farmingdale USA) and subjected to 96 cycles of 30 sec of sonication at 37 °C. After PMCA, the microplates were removed, and 10-μL aliquots of each sample were PK-digested as previously described<sup>35</sup>. The samples were then stored at −20 °C and subjected to dot blot and western blot analyses.

**RT-QulC.** RT-QulC amplifications were performed as previously described<sup>38,49</sup>. Briefly, the inocula were serially diluted in phosphate-buffered saline (PBS), pH 7.4 containing 0.01% SDS and 1/100 N2 supplement (Thermo Fisher cat# 17502048). Then, 2.04 μL of each dilution was loaded in triplicate in individual wells of a black 96-well optical bottom plate containing 100 μL of PBS mix (10μM thioflavin T, 1 mM EDTA and 100 μg/mL of recombinant ovine PrP, V136-R154-Q171 allele)<sup>44</sup>. The plate was sealed using Nunc Amplification Tape (Nalgene Nunc International Cat# 232702), placed in a Xenius XM spectrofluorometer (Safas, Monaco) and incubated for 48–60 h at 47 °C. One minute of orbital shaking (600 rpm) was applied every minute, and the fluorescence was recorded every 30 minutes. Data were treated using the 2<sup>nd</sup> derivate for detection of the inflection timepoint. The sample titer was estimated as the last dilution before the inflection timepoint get across the background level (determined as the inflection timepoint calculated based on the unseeded control).

**SDS-PAGE and western blot analysis.** PMCA samples were run on Criterion XT 12% Bis-Tris precast gels (Bio-Rad, Hercules, CA, USA), electrotransferred onto nitrocellulose membranes (Bio-Rad) and probed with biotinylated Sha31 anti-PrP monoclonal antibody<sup>69</sup>. Streptavidin-peroxidase incubation and ECL detection were achieved as described for the dot blot assay. When necessary, the PrP<sup>res</sup> content of PMCA samples was determined with ImageLab software after acquisition of the signals with a Chemidoc digital imager (Bio-Rad, USA).

**Histochemistry and immunofluorescence.** For histochemistry, brain spleen and granuloma samples of C-BSE and L-BSE-infected mice and uninfected control mice with granulomas were fixed with freshly prepared 4% formalin before being embedded in molten paraffin and stored at room temperature until ready for sectioning. Using a microtome, embedded spleen and granuloma tissues were cut into 5-μm-thick sections and mounted on Superfrost Plus slides. The sections were then stained with hematoxylin/eosin.

For immunofluorescence, frozen spleens and granulomas were cut on a cryostat machine (Thermo Fisher scientific). The 7–10-μm-thick sections were then adhered on Superfrost glass slides (Thermo Fisher scientific) and dried under air before storage at −80 °C for subsequent treatments. The tissue sections were fixed with 5% paraformaldehyde (PFA) for 45 minutes and then rinsed with 0.01 M phosphate-buffered saline (PBS), saturated and permeabilized with 5% bovine serum albumin (BSA) and 0.5% Triton-X100 for approximately 1 hour, followed by a 10-minute incubation with each component of the Avidin-Biotin blocking kit (Zymed, Invitrogen) according to the manufacturer's instructions. The sections were denatured with 3–3.5 M guanidinium thiocyanate and sufficiently washed with 1X PBS followed by double staining by incubation with two primary antibodies: 1/2000 dilution of biotinylated Sha31 mouse monoclonal anti-PrP (Sha31b) and one of the various cellular antigens (rat monoclonal antibodies): FDC-M1 (recognizing mfge-8 antigen, BD Bioscience-Pharmingen, 1/30) and FDC-M2 (recognizing C2/C4 antigen, AMS biotechnology, UK, 1/100) for follicular dendritic Cells, B220 (Invitrogen, 1/200) for B cells, CD3 (AbD Serotec, 1/200) for T cells, and CD68 (Hycult, 1/500) for macrophages, for 1 h at room temperature under saturated humidity. The sections were then washed and incubated with the secondary conjugates streptavidin-Alexa-633 or Alexa-488 (Invitrogen, 1/500) and CY3-donkey anti-rat (Jackson Immunoresearch, 1/500) or goat anti-mouse IgG (Jackson Immunoresearch, 1/50) for B cells, CD3 (AbD Serotec, 1/200) for T cells, and CD68 (Hycult, 1/500) for macrophages, for 1 h at room temperature under saturated humidity. The sections were then washed and incubated with the secondary conjugates streptavidin-Alexa-633 or Alexa-488 (Invitrogen, 1/500) and CY3-donkey anti-rat (Jackson Immunoresearch, 1/50) for B cells, CD3 (AbD Serotec, 1/200) for T cells, and CD68 (Hycult, 1/500) for macrophages, for 1 h at room temperature under saturated humidity. The sections were then washed and incubated with the secondary conjugates streptavidin-Alexa-633 or Alexa-488 (Invitrogen, 1/500) and CY3-donkey anti-rat (Jackson Immunoresearch, 1/50) for B cells, CD3 (AbD Serotec, 1/200) for T cells, and CD68 (Hycult, 1/500) for macrophages.

Confocal microscopy was performed with a Zeiss LSM 700 microscope (Confocal facilities, MIMA2 Platform, INRA Jouy-en-Josas, France, [https://www6.jouy.inra.fr/mima2](https://www6.jouy.inra.fr/mima2)) equipped with CLSM 700 confocal laser scanning software, using a Zeiss confocal microscope (images in false colors: blue, DAPI counterstaining) and a Plan Neofluar x40 (NA 1.3(fig A) or a Plan- Apochromat (NA 1.4) × 63 oil-immersion objectives and the 488-, 546- or 645-nm excitation wavelengths of the laser. All images were analyzed using ImageJ 1.51j8 software (Schneider et al., 2012) ([http://rsb.info.nih.gov/ij/](http://rsb.info.nih.gov/ij/)).
References
1. Prusiner, S. B. Prions. Proc. Natl. Acad. Sci. USA 95, 13363–13383 (1998).
2. Collinge, J., Sidle, K. C., Meads, J., Ironside, J. & Hill, A. F. Molecular analysis of prion strain variation and the aetiology of ‘new variant’ CJD. Nature 383, 685–690 (1996).
3. Benestad, S. L., Mitchell, G., Simmons, M., Ytrehus, B. & Viker, T. First case of chronic wasting disease in Europe in a Norwegian free-ranging reindeer. Vet. Res. 47 (2016).
4. Igel-Egalon, A., Béringer, V. & Rezaei, H. & Sible, P. Prion Strains and Transmission Barrier Phenomena. Pathogens 7, 5 (2018).
5. Mabbott, N. A., Farquhar, C. F., Brown, K. L. & Bruce, M. E. Involvement of the immune system in TSE pathogenesis. Immunol. Today 19, 201–203 (1998).
6. Jeffrey, M., McGovern, G., Martin, S., Goodsir, C. M. & Brown, K. L. Cellular and sub-cellular localisation of PrP in the lymphoreticular system of mice and sheep. Arch. Virol. Suppl. 23–38 (2000).
7. Andreotti, O. et al. Early accumulation of PrP(Sc) in gut-associated lymphoid and nervous tissues of susceptible sheep from a Romanov flock with natural scrapie. J. Gen. Virol. 81, 3115–3126 (2000).
8. Forloni, G. The contribution of the immune system to prion diseases. Drug Discov. Today Dis. Mech. 1, 351–356 (2004).
9. Kitamoto, T., Muramoto, T., Mohri, S., Doh-Ura, K. & Tateishi, J. Abnormal isoform of prion protein accumulates in follicular dendritic cells in mice with Creutzfeldt-Jakob disease. J. Virol. 65, 6292–6295 (1991).
10. McBride, P. A., Eikelenboom, P., Kraal, G., Fraser, H. & Bruce, M. E. PrP protein is associated with follicular dendritic cells of spleens and lymph nodes in uninfected and scrapie-infected mice. J. Pathol. 168, 413–418 (1992).
11. Mabbott, N. A., Mackay, F., Minns, F. & Bruce, M. E. Temporary inactivation of follicular dendritic cells delays neuroinvasion of scrapie. Nat. Med. 6, 719–720 (2000).
12. Mabbott, N. A., Young, J., McConnell, I. & Bruce, M. E. Follicular Dendritic Cell Dedifferentiation by Treatment with an Inhibitor of the Lymphoendothelial Pathway Dramatically Reduces Scrapie Susceptibility. J. Virol. 77, 6845–6854 (2003).
13. McCulloch, L. et al. Follicular dendritic cell-specific prion protein (PrP) expression alone is sufficient to sustain prion infection in the spleen. PLoS Pathog. 7, e1002402 (2011).
14. Brown, K. L. et al. Scapie replication in lymphoid tissues depends on prion protein-expressing follicular dendritic cells. Nat Med 5, 1308–1312 (1999).
15. Mohan, J., Bruce, M. E. & Mabbott, N. A. Follicular dendritic cell dedifferentiation reduces scrapie susceptibility following inoculation via the skin. Immunology 114, 225–234 (2005).
16. Prinz, M. et al. Lymph nodal prion replication and neuroinvasion in mice devoid of follicular dendritic cells. Proc. Natl. Acad. Sci. USA 99, 919–924 (2002).
17. Maestrale, R. et al. A Lympho-Follicular Microenvironment Is Required for Pathological Prion Protein Deposition in Chronically Infected Tissues from Scrapie-Affected Sheep. PLoS ONE 6, e26830 (2013).
18. Heikenwalder, M. et al. Chronic lymphocytic inflammation specifies the organ tropism of prions. Science 307, 1107–1110 (2005).
19. Béringer, V. et al. Facilitated Cross-Species Transmission of Prions in Extraneural Tissue. Science 335, 472–475 (2012).
20. Gill, O. N. et al. Prevalent abnormal prion protein in human appendixes after bovine spongiform encephalopathy epizootic: large scale survey. BMJ 347, f6675–f6675 (2013).
21. Heikenwalder, M. et al. Lymphoendothelial prion replication in inflammatory stromal cells of granulomas. Immunity 29, 998–1008 (2008).
22. Vilote, J. L. et al. Markedly Increased Susceptibility to Natural Sheep Scapie of Transgenic Mice Expressing Ovine PrP. J. Virol. 75, 5977–5984 (2001).
23. Langevin, C., Andréoletti, O., Le Dur, A., Laude, H. & Béringer, V. Marked influence of the route of infection on prion strain apparent phenotype in a scrapie transgenic mouse model. Neurobiol. Dis. 41, 219–225 (2011).
24. Béringer, V. et al. A bovine prion acquires an epidemic bovine spongiform encephalopathy strain-like phenotype on interspecies transmission. J. Neurosci. Off. J. Soc. Neurosci. 27, 6965–6971 (2007).
25. Moudjou, M. et al. Highly Infectious Prions Generated by a Single Round of Microplate-Based Protein Misfolding Cyclic Amplification. mBio 5, e00829–13 (2014).
26. Béringer, V. et al. Opposite Effects of Dextran Sulfate 500, the Polyene Antibiotic MS-8209, and Congo Red on Accumulation of the Protease-Resistant Isoform of PrP in the Spleens of Mice Inoculated Intraperitoneally with the Scrape Agent. J. Virol. 74, 5432–5440 (2000).
27. Peretz, D. et al. Strain-specified relative conformational stability of the scrapie prion protein. Protein Sci. Publ. Protein Soc. 10, 854–863 (2001).
28. Williams, J. M. et al. Rapid End-Point Quantitation of Prion Seeding Activity with Sensitivity Comparable to Bioassays. PLoS Pathog. 6, 1 (2010).
29. Neyt, K., Perros, F., GeurtvensKessel, C. H., Hammad, H. & Lambrecht, B. N. Tertiary lymphoid organs in infection and autoimmunity. Trends Immunol. 33, 297–305 (2012).
30. Taylor, P. R. et al. The follicular dendritic cell restricted epitope, FDC-M2, is complement C4; localization of immune complexes in mouse tissues. Eur. J. Immunol. 32, 1883–1896 (2002).
31. Kimberlin, R. H. & Walker, C. A. The role of the spleen in the neuroinvasion of scrapie in mice. Virus Res. 12, 201–211 (1989).
32. Bartz, J. C., Dejioa, C., Tucker, T., Kincaid, A. E. & Bessen, R. A. Extraneuronal prion neuroinvasion without lymphoreticular system infection. J. Virol. 79, 11858–11863 (2005).
33. Shiikiya, R. A. et al. PrPSc formation and clearance as determinants of prion tropism. PLOS Pathog. 13, e1006298 (2017).
34. Montrasio, F. et al. Impaired Prion Replication in Spleens of Mice Lacking Functional Follicular Dendritic Cells. Science 288, 1257–1259 (2000).
35. Béringer, V. et al. Role of spleen macrophages in the clearance of scrapie agent early in pathogenesis. J. Pathol. 190, 495–502 (2000).
36. McCulloch, L., Brown, K. L. & Mabbott, N. A. Ablation of the cellular prion protein, PrP(C), specifically on follicular dendritic cells has no effect on their maturation or function. Immunology, https://doi.org/10.1111/imn.12031 (2012).
37. Mabbott, N. A. et al. Expression of mesenchyme-specific gene signatures by follicular dendritic cells: insights from the meta-analysis of miRNome data from multiple mouse cell populations. Immunology 133, 482–498 (2011).
38. Muller, S. N. & Germain, R. N. Normal development and function of B cells contributions to the homeostasis and functionality of the immune system. Nat. Rev. Immunol. 9, 618–629 (2009).
39. Heinen, E. et al. Retention of immune complexes by murine lymph node or spleen follicular dendritic cells. Role of antibody isotype. Scand. J. Immunol. 24, 327–334 (1986).
40. Pedersen, A. H., Ritchie, D. L., Head, M. W. & Ironside, J. W. Detection and Localization of PrPSc in the Skeletal Muscle of Patients with Variant, Iatrogenic, and Sporadic Forms of Creutzfeldt-Jakob Disease. Am. J. Pathol. 168, 927–935 (2006).
41. Bieler, H. et al. Normal development and behaviour of mice lacking the neuronal cell-surface PrP protein. Nature 356, 577–582 (1992).
42. Tixador, P. et al. The Physical Relationship between Infectivity and Prion Protein Aggregates is Strain-Dependent. PLoS Pathog 6, e1000859 (2010).
43. Orrá, C. D. et al. Bank Vole Prion Protein As an Apparently Universal Substrate for RT-QuIC-Based Detection and Discrimination of Prion Strains. PLoS Pathog 11, e1004983 (2015).
44. Rezaei, H. et al. High yield purification and physico-chemical properties of full-length recombinant allelic variants of sheep prion protein linked to scrapie susceptibility. *Eur. J. Biochem.* **267**, 2833–2839 (2000).

45. Féraudet, C. et al. Screening of 145 Anti-PrP Monoclonal Antibodies for Their Capacity to Inhibit PrPSc Replication in Infected Cells. *J. Biol. Chem.* **280**, 11247–11258 (2005).

**Acknowledgements**

We would like to thank Lucy De Guilhem de Lataillade for kind technical assistance in the IF studies and Christelle Rossignol for performing the paraffin sectioning and hematoxylin/eosin staining. The authors are grateful to the INRA MIMA2 imaging core facility (P. Adenot) for access to the virtual slide scanner (Pannoramic SCAN, 3DHISTECH) and to the confocal microscope. This work was funded by Alliance Biosecure (BoHuToHu) and by Fondation pour la Recherche Médicale (FRM) (Equipe FRM DEQ. 20150331689). Iman Al Dybiat’s PhD fellowship was funded by the Syrian Ministry of Higher Education as part of the Franco-Syrian cooperation program.

**Author Contributions**

I.A.D., P.S., M.M., D.M., H.R., F.R., L.H., S.T. participated substantially to all 580 experiments. P.B. performed the histological studies. P.S. and V.B. wrote and corrected the 581 manuscript. P.S., V.B., O.A. and H.L. contributed to design or fund the study.

**Additional Information**

**Supplementary information** accompanies this paper at https://doi.org/10.1038/s41598-019-51084-1.

**Competing Interests:** The authors declare no competing interests.

**Publisher’s note** Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

© The Author(s) 2019