Prions in Milk from Ewes Incubating Natural Scrapie

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Abstract
Since prion infectivity had never been reported in milk, dairy products originating from transmissible spongiform encephalopathy (TSE)-affected ruminant flocks currently enter unrestricted into the animal and human food chain. However, a recently published study brought the first evidence of the presence of prions in mammary secretions from scrapie-affected ewes. Here we report the detection of consistent levels of infectivity in colostrum and milk from sheep incubating natural scrapie, several months prior to clinical onset. Additionally, abnormal PrP was detected, by immunohistochemistry and PET blot, in lacteal ducts and mammary acini. This PrPSc accumulation was detected only in ewes harbouring mammary ectopic lymphoid follicles that developed consequent to Maedi lentivirus infection. However, bioassay revealed that prion infectivity was present in milk and colostrum, not only from ewes with such lymphoproliferative chronic mastitis, but also from those displaying lesion-free mammary glands. In milk and colostrum, infectivity could be recovered in the cellular, cream, and casein-whey fractions. In our samples, using a Tg 338 mouse model, the highest per ml infectious titre measured was found to be equivalent to that contained in 6 mg of a posterior brain stem from a terminally scrapie-affected ewe. These findings indicate that both colostrum and milk from small ruminants incubating TSE could contribute to the animal TSE transmission process, either directly or through the presence of milk-derived material in animal feedstuffs. It also raises some concern with regard to the risk to humans of TSE exposure associated with milk products from ovine and other TSE-susceptible dairy species.

Results
In this study, we first investigated material collected between 2003 and 2006 from a sheep flock with a high incidence of natural scrapie (Langlade Flock) [12] and in which Maedi lentivirus has
A decade ago, a new variant form of Creutzfeldt-Jakob disease was identified. The emergence of this prion disease in humans was the consequence of the zoonotic transmission of bovine spongiform encephalopathy through dietary exposure. Since then, the control of human exposure to prions has become a priority, and a policy based on the exclusion of known infectious materials from the food chain has been implemented. Because all investigations carried out failed to reveal evidence of infectivity in milk from affected ruminants, this product has continuously been considered as safe. In this study, we demonstrate the presence of prions in colostrum and milk from sheep incubating natural scrapie and displaying apparently healthy mammary glands. This finding indicates that milk from small ruminants could contribute to the transmission of prion disease between animals. It also raises some concern with regard to the risk to humans associated with milk products from ovine and other dairy species.

been endemic for more than 10 years. Sheep from this flock were investigated for the presence of PrPSc in (i) mammary glands, (ii) lymphoreticular system (LRS) and (iii) central nervous system (CNS) (Table 1). The sheep carried various PRP polymorphisms at codons 136, 154 and 171 that are associated either with high susceptibility (ARQ/VRQ, VRQ/VRQ, ARQ/ARQ) or resistance (homozygote and heterozygote ARR) to TSEs [12].

The majority of susceptible genotype animals were clinically suspect of scrapie at the time of culling. The majority of susceptible genotype sheep were clinical suspect for scrapie at culling. All investigated animals (n = 175) were PCR positive for Maedi lentivirus. Abnormal PrP (PrPSc) detection was carried out using Western Blotting (SHa31 anti-PrP antibody), ELISA (TeSeE Sheep and Goat, Bio-Rad) and immunohistochemistry (8G8 anti-PrP antibody). The majority of susceptible genotype sheep (ARQ/VRQ, VRQ/VRQ) were clinical suspect for scrapie at the time of culling (Figure 1A and 1B). Double labelling indicated that these PrPSc positive cells were also positive with CD68, a marker of phagocyte cells which could encompass both macrophages and dendritic cell subsets [16,19].

Table 1. PrPSc in central nervous system, lympho-reticular system, mammary gland and milk duct lumen of natural scrapie exposed ewes bearing various genotypes at codons 136, 154 and 171 of the PRP gene.

| Genotype | number | PrPSc in obex | PrPSc in tonsil, prescapular lymph node, spleen and mammary lymph node | PrPSc in mammary gland | Ectopic mammary lymphoid follicles | PrPSc in lacteal ducts |
|----------|--------|---------------|---------------------------------------------------------------------|------------------------|-----------------------------------|------------------------|
| VRQ/VRQ  | n = 110 | pos           | pos                                                                 | pos n = 45             | n = 45                            | n = 27                  |
|          |        |               |                                                                     | neg n = 65             | n = 0                             | n = 0                   |
| ARQ/VRQ  | n = 11  | pos           | pos                                                                 | pos n = 2              | n = 2                             | n = 1                   |
|          |        |               |                                                                     | neg n = 9              | n = 0                             | n = 0                   |
| ARQ/ARQ  | n = 13  | pos           | pos n = 6                                                           | pos n = 3              | n = 3                             | n = 1                   |
|          |        |               |                                                                     | neg n = 3              | n = 0                             | n = 0                   |
|          |        |               |                                                                     |                        | neg n = 7                          |                        |
|          |        |               |                                                                     |                        | neg n = 7                          |                        |
|           |         | neg           | pos                                                                 | neg                    | n = 3                             | n = 0                   |
| ARR/ARR  | n = 32  | neg           | pos                                                                 | neg                    | n = 3                             | n = 0                   |
|          |        |               |                                                                     |                        | n = 16                            |                        |
|          |        |               |                                                                     |                        |                                   |                        |
|          |        |               |                                                                     |                        |                                   |                        |

All investigated animals (n = 175) were PCR positive for Maedi lentivirus. Abnormal PrP (PrPSc) detection was carried out using Western Blotting (SHa31 anti-PrP antibody), ELISA (TeSeE Sheep and Goat, Bio-Rad) and immunohistochemistry (8G8 anti-PrP antibody). The majority of susceptible genotype sheep (ARQ/VRQ, VRQ/VRQ and ARQ/ARQ) were clinical suspect for scrapie at culling.

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Author Summary

A decade ago, a new variant form of Creutzfeldt-Jakob disease was identified. The emergence of this prion disease in humans was the consequence of the zoonotic transmission of bovine spongiform encephalopathy through dietary exposure. Since then, the control of human exposure to prions has become a priority, and a policy based on the exclusion of known infectious materials from the food chain has been implemented. Because all investigations carried out failed to reveal evidence of infectivity in milk from affected ruminants, this product has continuously been considered as safe. In this study, we demonstrate the presence of prions in colostrum and milk from sheep incubating natural scrapie and displaying apparently healthy mammary glands. This finding indicates that milk from small ruminants could contribute to the transmission of prion disease between animals. It also raises some concern with regard to the risk to humans associated with milk products from ovine and other dairy species.
Figure 1. PrPSc detection in mammary gland from scrapie-incubating sheep. (A) PrPSc immunolabelling (8G8 monoclonal antibody - DAB brown deposit - bar: 80 μm) in mammary gland from a ewe incubating scrapie (preclinical phase - 15 months old - ARR/ARQ genotype) and harbou- ring lympho-proliferative mastitis with ectopic lymphoid follicles (Foill). In the milk duct lumen (arrow heads), several PrPSc positive cells are identifiable. (B) In mammary gland acini (Aci), positive PrPSc staining can be observed; either associated with cells or distributed as free granules. (C) Double labelling for PrPSc (R521 polyclonal serum - black deposits) and CD68 (KiM6 clone - red deposits) indicates that intracellular PrPSc in milk ducts and acini lumen is associated with phagocytic cells. (D) PrPSc immunolabelling (8G8 anti-PrP antibody - DAB brown deposit- bar: 200 μm) and (E) PET blot (Sha31 antibody - NB7/BCIP black deposits - bar: 200 μm) of two successive mammary gland sections confirmed that material in milk ducts is proteinkase K resistant (arrow heads indicate lining).

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Discussion

Failure to transmit TSE using milk from scrapie affected or incubating ewes in conventional rodents models, has been previously reported [7]. In our study, the use of a transgenic mouse model expressing the ovine PRP gene reduced or abolished the species barrier phenomenon [25] and could explain, at least partially, the positive results we obtained. However the low infectivity level we measured (100.1 to 101.6 ID50/ml) indicates that, even in this transgenic animal model, direct inoculation with 20 μl per mouse of whole milk or colostrum would be unlikely to transmit disease. It also means that, despite no PK resistant PrP being detected in milk and colostrum fractions, the approach we used (i.e immunoprecipitation of all detectable PrP on beads) was sufficiently efficient to concentrate infectivity. Measurements of the concentrative efficacy of this method are currently under investigation and will be reported elsewhere.

A recently published study reported the successful transmission of scrapie to lambs through consumption of colostrum/milk collected from ewes at the late incubation or clinical stage of the disease [11]. This study was the first to identify Prion presence in mammary secretions from Scrapie affected ewes. However, because of its design, this study did not elicit information as to which fraction, colostrum or milk, induced disease transmission. Moreover, since lateral contamination occurred between lambs and in some cases several ewes were used to feed a lamb, it was not possible in this experiment to clearly determine which of the donor ewes were shedding infectivity in colostrum/milk. The data we provided here brings definitive and unambiguous evidence of the presence infectivity in both milk and colostrum from naturally incubating scrapie ewes. In our model, infectivity was detected up to 20 months before clinical disease onset and a majority of ewes (10 out of 13 at the moment of writing) were demonstrated to have shed prion infectivity in their milk.

Taken together, the results reported by Konold et al. [11] and those obtained in our study, raise the issue of the use of sheep milk or milk by-products for animal feeding. Currently ruminants’ milk represents a major source of protein in milk-replacer and feedstuffs used in a variety of farm animal species. The use of TSE incubating ewe milk in such products could give rise to dietary exposure of animals both intra and interspecies. Given the low level of infectivity apparent in milk and the species barrier phenomenon, the interspecies transmission risk associated with ewe milk certainly remains limited. However, in the current stage of knowledge, the possibility of such transmission cannot be ruled out.

As all scrapie samples in our study were collected from a single flock, it is likely that investigated ewes were exposed to only a limited range of TSE agents and possibly to a single one. Consequently, caution should be taken before inferring those observations to other situations. Interactions between host genotype and TSE agent are known to impact on the kinetics of...
| Organ/Age | 4 Months | 7 Months | 10 Months | 13 Months | 20 Months | 32 Months |
|-----------|----------|----------|-----------|-----------|-----------|-----------|
|           | VRQ/VRQ  | ARQ/VRQ  | VRQ/VRQ   | ARQ/VRQ   | VRQ/VRQ   | ARQ/VRQ   |
| Obex      | —        | —        | —         | —         | —         | —         |
| Tonsil    | 4        | +++      | —         | —         | —         | —         |
| Spleen    | 4        | —        | 4         | 4++       | 4         | 4++       |
| Duodenal PP | 4      | +++      | 1(a)      | 4         | +++       | 4         |
| Jejunum PP | 4       | +++      | 1(c)      | 4         | +++       | 4         |
| Ileum PP  | 4        | +++      | 3(a-b-c)  | 4++       | 4         | 4++       |
| Caecum PP | 4        | +++      | 3(a-b-c)  | 4++       | 4         | 4++       |
| NLM-jejunal | 4    | +++      | 4         | 4++       | 4         | 4++       |
| ileal MLN | 4        | +++      | 2(a-b)    | 4         | +++       | 4         |
| Médiastinal LN | 4   | +++      | 3(a-b-c)  | 4++       | 4         | 4++       |
| Prescapular LN | 4   | +++      | 3(a-b-c)  | 4++       | 4         | 4++       |
| Retro hepatic LN | 4 | +++      | 2(a-b)    | 4++       | 4         | 4++       |
| AbomasumENS | —      | —        | —         | —         | 1(c)      | 1(d)      |
| DuodenumENS | —     | —        | 1(a)      | 0/+       | 4         | 4++       |
| Ileum ENS | —        | —        | 4         | +++       | 4         | 4++       |
| Caecum ENS | —       | —        | 3(a-b-c)  | 4        | 2(a-c)    | 4         |
| Colon ENS | —        | —        | —         | 4         | 2(a-c)    | 4         |

Groups of 4 ARQ/VRQ and 4 VRQ/VRQ sheep (named a-b-c-d) were killed at different time of the incubation period. Clinical signs occurred at 20 months in VRQ/VRQ animals and at 32 months in ARQ/VRQ. A systematic PrPSc detection was realized using immunohistochemistry (8G8 antibody) in a large panel of the collected sheep tissues. PrPSc accumulation level was scored according to a semi-quantitative scale: (−) no PrPSc, (+) minimal PrPSc deposits, (++) moderate PrPSc deposits and (+++) strong PrPSc deposits.

LN: Lymph Node; MLN: Mesenteric Lymph Node; PP: Peyer's Patches; ENS: Enteric Nervous System.

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prion dissemination in ewes. Such interactions could also influence, not only how early the shedding of infectivity via colostrum and milk takes place but also the levels of infectivity.

Prion infectivity was detected here in sheep with healthy mammary glands as well as those with chronic lymphoproliferative mastitis. However, our preliminary results suggest that the presence of ectopic lymphoid follicles in ewes with lymphoproliferative mastitis could increase prion shedding in milk. Acute and subacute bacterial mastitis are extremely common in dairy animals and can impact on milk composition [26]. The effect of these conditions on milk prion shedding was not addressed in this study and remains to be evaluated.

Our study was carried out in a flock affected by Maedi-Visna Lentivirus. This non oncogenic retrovirus is largely spread in sheep and remains to be evaluated.

A potential enhancement of the prion infectivity shedding in milk from persistently Maedi Virus infected ewes (with no pathological manifestations) cannot be ruled out [30]. However, because of the relative high prevalence of such infection in dairy ewes production areas, such hypothetical effect would not impact, in our opinion, on the significance of our observations.

There are major differences in terms of peripheral pathogenesis between BSE in cattle and TSE in other ruminants. In cattle BSE, peripheral tissues PrPSc accumulation and infectivity is marginal and this is particularly true of lymphoid tissues [31]. Such differences prevent the observations reported here in sheep being directly extrapolated to BSE in cattle. Nevertheless, these results clearly call for the re-examination of milk from BSE affected cattle for the presence of prions.

Finally, the consequences for humans of the presence of prions in sheep milk should certainly be given consideration. However, it is our opinion that its relative impact on global TSE dietary exposure is of lower magnitude than other prion sources, such as lymphoid tissues from small ruminants incubating TSE [32,33].

Methods
Scrapie affected animals and Maedi PCR diagnosis
Scrapie positive ewes included in this experiment were all Romanov sheep born and bred in the Langlade flock. In this flock a natural scrapie epidemic has been occurring at a high incidence since 1993 [12].

Since 1997, all animals belonging to this flock are:

(a) genotyped at two months of age for codon 136 (A/V), 154 (R/H) and 171 (Q/H/R) of the PRP gene by the SNP taqman probe method (Labogena, Jouy en Josas).

(b) necropsied with collection of central nervous system, lymphoid tissues and several other tissues (including mammary gland). Samples are both formalin fixed/paraffin embedded and frozen stored. The retrospective study involved a set of samples collected between 2003 and 2006. Susceptible genotype sheep included in this retrospective study were, in the majority, clinically suspect for scrapie at the time of culling.

For the prospective study a group (n=13) of Langlade ewes, having susceptible genotypes ARQ/VRQ and VRQ/VRQ, was constituted. In the first 12 hours post lambing, 5 to 20 ml of colostrum was collected in TSE free conditions, the lambs having been separated for 4 hours from the ewes. Similarly, at 20 days post lambing, individual samples of milk (10 to 50 ml) were collected.

Milk from two VRQ/VRQ cheviot TSE free sheep (Arthur Rickwood, UK) was collected and included in the study as a negative control. This flock is the only source in Europe of sheep free of classical scrapie. The TSE free status of the dams was confirmed by post-mortem laboratory examination.

PCR detection of Maedi virus was carried out on DNA extracted from mammary tissue. Primers (Forward:CCACGTGCGGCGC-CAGCTGCGAGA-Reverse:TGAGACTGAAATTGTAACCGCG-AAG) and PCR conditions (40 cycles – annealing 58°C) were those published by Sonigo et al [34]. Reference positive case and a negative controls were included in each PCR run. PCR products (291 bp) were migrated on a 2% agarose TBE gel. Positive samples

Table 3. End-point titration of a brain homogenate (posterior brainstem- 12.5% weight/volume homogenate) in Tg338 mice.

| Dilution | Number of positive mice | Incubation period in days (mean+/− SD)* |
|----------|-------------------------|----------------------------------------|
| neat     | 6/6                     | 221+/− 20                              |
| 10⁻¹     | 6/6                     | 348+/− 16                              |
| 10⁻²     | 12/12                   | 481+/− 32                              |
| 10⁻³     | 10/12                   | 594+/− 34                              |
| 10⁻⁴     | 7/12                    | 713+/− 43                              |
| 10⁻⁵     | 3/12                    | 805, 824, 852*                         |
| 10⁻⁶     | 0/12                    | >900                                   |

The donor ewe was born and bred in the Langlade Flock. This ewe was at the terminal stage of Scrapie at the moment of culling. Each mouse was intracerebrally inoculated with 20 μl of homogenate. Mice were considered positive when abnormal PrP deposition was detected in brain. Incubation periods are presented as mean+/− SD except for that dilution with which less than 20% of mice were found positive. In that case (*) incubation times of the positive mice are individually presented.

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Figure 3. Infectivity testing in a reference brain sample and colostrum/milk fractions from scrapie incubating ewes. (A,B) Survival curve in Tg338 mice (transgenic mice over-expressing ovine VRQ PRP allele) intracerebrally inoculated with colostrum (A) and milk (B), collected from ewes incubating scrapie. Samples were first fractionated into cellular pellet (△), cream (▼), and casein whey (○). An immunoprecipitation of PrP on magnetic beads coated with anti-PrP antibodies was then carried out. Beads from each fraction were inoculated into five or six Tg338 mice. (A) Colostrum fractions from a ewe harbouring mammary ectopic lymphoid follicles associated with Maedi lesions (white symbols) and from a ewe with a healthy mammary gland (black symbols). (B) Milk fractions from the same ewes as in A (black symbols and white symbols) and of the cellular fraction.
from a second scrapie incubating ewe with a healthy mammary gland (grey symbols). The experiment was terminated after 900 days (normal Tg338 mouse lifespan). Incubation periods have to be compared to those of successive 1/10 dilutions of brain (obex- vertical dotted lines) material from a sheep clinically affected with scrapie. The start point (mean) corresponds to the inoculation of 2.5 μg of brain tissue per mice. (C) Western-blotting (anti-PrP SHa31 antibody) of without (lane 1) and with (lane 2) PK treatment of brain material from a Tg338 mouse inoculated with scrapie positive brain (10^{-3} diluted); (lanes 2–6) PK digested brain material from mice inoculated with milk and colostrum cellular fraction – (lane 3) milk from a ewe with a healthy mammary gland – (lane 4) colostrum from a ewe with a healthy mammary gland – (lane 5) milk from TSE free control – (lane 6) colostrum from a Maedi affected (ectopic lymphoid follicle) ewe. (D) Intracerebral end point titration of a 12.5% obex homogenate, prepared from a terminally scrapie affected sheep (Langlade isolate), in a Tg338 mouse model. This titration allowed the determination of the infectious dose 50 (ID50) of the brain sample (10^{9.1} ID50/g), see the text. (E) Variation of the incubation period as a function of the infectious dose inoculated intracerebrally in Tg338 mice (obex – Langlade isolate), see the text.

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were identified on the basis of PCR product size (by comparison with a positive control plasmid);

PrPSc distribution in ARQ/VRQ and VRQ/VRQ sheep organism

An ARQ/VRQ (n = 72 – age 18 months) ewe cohort was inseminated with semen from a single VRQ/VRQ ram. Natural mating or contact between ewes and rams was strictly avoided. The resulting Lambs (birth cohort September 2003) were PRP genotyped at the age of 2 months. Groups of 4 VRQ/VRQ and 4 ARQ/VRQ animals were euthanized at 32 months, when first clinical signs were observed. A final group of 4 ARQ/VRQ animals was euthanized at 20 months of age. Scrapie clinical signs occurred in homozygote VRQ animals intravenously pentobarbital (DOLETHALND, 10 mg/kg) injection at 20 months of age. Scrapie clinical signs occurred in homozygote VRQ animals genotyped at the age of 2 months. Groups of 4 VRQ/VRQ and 4

Tissue processing and immunohistochemistry (IHC) detection

This method was performed as previously described [16]. PrPSc IHC detection was first performed using 8G8 antibody raised against human recombinant PrP protein and specifically recognizing the 95–108 amino acid sequence (SQWKNKP) of the PrP protein.

For each sample a negative serum control was included, in which the primary antibody was either omitted or replaced by purified mouse Ig2a serum. In addition, anti-PrP monoclonal antibodies were replaced by isotype-matched monoclonal antibodies irrelevant to the protein under investigation. PrPSc/CD68 double labelling was performed as previously described [16], using KiM6 monoclonal mouse anti human CD68 (Serotec, London, UK) and a rabbit anti-PrP serum (R521- diluted 1/1000– CIV, the Netherlands). For double-labelling, cross-reactivity controls were performed, in order to verify the absence of inter-species reactivity of secondary antibodies toward primary antibodies. The absence of affinity between the two secondary antibodies was also checked.

Table 4. Estimation of infectious titre in colostrum and milk from scrapie incubating ewes with apparently healthy mammary glands or lymphoproliferative mastitis (consecutive to Maedi infection).

| Ewe | Ecto. Lymph. follicles | Fraction | Quantity of material submitted to IP | Starting whole milk volume | Pos mice | Incubation period in days (mean/- SD) | Estimated infectious titre (ID50 IC in Tg 338) | Global infectious titre/ml |
|-----|----------------------|----------|-------------------------------------|--------------------------|----------|--------------------------------------|-----------------------------------------------|--------------------------|
| 0942 | pos | colostrum | Cell pellet | 10^5 cells | 10 ml | 6/6 | 524+/−45 | 85 | 10^{1.2} /ml |
| | | | | 3.6 ml | 3.6 ml | 6/6 | 609+/−81 | 15 | |
| | | | | 1.3 ml | 20 ml | 6/6 | 612+/−62 | 15 | |
| | | | | | | | | |
| | | Milk | Cell pellet | 1.3 10^6 cells | 30 ml | 5/5 | 355+/−58 | 1250 | 10^{1.6} /ml* |
| | | | | 3.6 ml | 3.6 ml | N.A | N.A (>400) | |
| | | | | 1.3 ml | 38 ml | N.A | N.A (>400) | |
| 0248 | neg | colostrum | Cell pellet | 6 10^6 cells | 8 ml | 6/6 | 685+/−39 | 5 | 10^{3.3} /ml |
| | | | | 3.6 ml | 3.6 ml | 3/6 | <1 | |
| | | | | 1.3 ml | 17 ml | 1/6 | <1 | |
| | | Milk | Cell pellet | 10^6 cells | 10 ml | 6/6 | 717+/−45 | 2 | 10^{0.1} /ml |
| | | | | 3.6 ml | 3.6 ml | 3/6 | <1 | |
| | | | | 1.3 ml | 35 ml | 1/6 | <1 | |
| 0370 | neg | Milk | Cell pellet | 510^5 cells | 7 ml | 6/6 | 509+/−34 | 90 | 10^{1.1} /mlb |
| | | | | 3.6 ml | 3.6 ml | N.A | N.A | |
| | | | | N.A | N.A | N.A | N.A | |

For each fraction (cell pellet, casein whey, cream) the quantity of the material submitted to immunoprecipitation process is detailed and linked to the initial volume of colostrum or milk from which it was prepared. In samples for which a 100% attack rate was observed, mean incubation period were used to estimate the infectious titre (Figure 3E). For each considered fraction the infectious titre per ml of starting material was calculated. The global infectious titre per ml of colostrum and milk was finally obtained by adding the value corresponding to each fraction.

N.A: not available at the moment of writing. *Infectivity was estimated from the only those fractions for which results are available. Consequently the calculated infectious titre/ml of milk is certainly underestimated.

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Paraffin embedded tissue blot (PET blot)

PET blot was performed using a method previously described [35]. Immunodetection was carried out using SHa31 monoclonal antibody (4 μg/ml), which recognizes the 145–152 sequence of PrP (YEDRYRE), followed by application of an alkaline phosphatase labelled secondary antibody (Dako ref D0314 – 1/500 diluted). Enzymatic activity was revealed using NBT/BCIP substrate chromogen. For each tissue sample, serial sections of 4 μm thickness for PET blot and 2 μm for IHC were collected onto membranes or glass slides respectively. Both methods were then washed and the resulting preparations were subject to comparison. This experimental design allowed the use of shape and localization of labeling on the IHC sections to identify the nature of PET blot PrP<sup>Sc</sup> positive structures.

Milk and colostrum fraction preparation

Milk and colostrum samples were all collected under TSE sterile conditions. An aliquot of each collected sample was submitted to a standard somatic cell count (SCC) (by flow cytometry) by an accredited laboratory (LIAL, Auch, France).

Each sample type was first diluted (1/2 for milk and 1/5 for colostrum) in PBS containing 10% acid-citrate-dextrose (Sigma-Aldrich ref C3821) and 10 mM EDTA-2K and homogenized by inversion. After standing for 30 min at 4°C, the cream was collected with a single use spatula. A Pasteur pipette was then passed through the layer of residual cream and the liquid was aspirated; this was passed through a 200 μm filter and collected in 50 ml tubes. The liquid sample was then centrifuged at 2000 rpm at 4°C for 5 minutes. The supernatant (casein-whey) was collected and stored frozen while cell pellets were transferred to a new 50 ml tube. The cells were washed three times by successive centrifugation/resuspension phases in PBS and counted in a Malassez cell before frozen storage.

PrP immunoprecipitation in colostrum and milk fractions

Cells and whey were extracted for 10 min at 37°C in NP40/DOC buffer (NP40 0.5% (V/V), deoxycholate 1% (W/V), EDTA 10 mmol.L<sup>-1</sup>, NaCl 150 mmol.L<sup>-1</sup>, Tris 10 mmol.L<sup>-1</sup> pH 7.4). The cream was extracted for 10 min at 37°C in sarkosyl buffer (N-lauroyl sarcosine 10% (W/V), EDTA 10 mmol.L<sup>-1</sup>, NaCl 150 mmol.L<sup>-1</sup>, Tris 20 mmol.L<sup>-1</sup> pH 7.4). Three different monoclonal antibodies (SHa31, SAF-34 and βS-36) [20,36] were immobilized by covalent coupling to magnetic beads (Dynal Biotech) and used to perform immunoprecipitation. Two successive immunoprecipitations (overnight 4°C), using a mixture of the three different antibody coated beads, were performed and any non-immunoprecipitated residual PrP in the supernatant was measured using a two site sandwich immunoassay (capture antibody 11C6 [20], tracer antibody Bar-224 [20]- CEA Saclay). Whenever the concentration of residual PrP was in excess of 5% of that of the unimmunoprecipitated control a third immunoprecipitation (2 h/RT) was performed. Beads were washed three times (two washings in PBS/Tween 1% and one in PBS) before re-suspension in the appropriate volume of 5% glucose. The concentration factor was 15 times for cream (equivalent to 300 μL inoculated per mouse) and 30 for whey (equivalent to 600 μL inoculated per mouse).

Colostrum and milk bioassay

Bioassay experiments were carried out in ovine VRQ PrP transgenic mice (Tg338), which are considered to be highly efficient for the detection of sheep scrapie infectivity [22]. Immunoprecipitated cream, casein whey and cell pellet fractions were re-suspended in 130 μL of sterile 5% glucose. Six mice were intracerebrally inoculated with each sample (20 μL). Colostrum inoculations were carried out in UMR INRA ENVT 1225 (Toulouse, France) facilities while milk fractions were tested at both the NVI (Oslo, Norway) and at INRA IASP (Tours, France). Samples from each animal were inoculated on different days so as to avoid any risk of cross contamination. Mice were then clinically monitored until the occurrence of TSE clinical signs, at which time they were culled. CNS and spleen samples were individually collected and Western blot (WB) tested. Mice inoculated with control TSE free sheep tissue and milk fractions were culled 950 days post inoculation. The majority of the bioassays are still in progress.

Estimation of infectious titre

A sample of obex from a VRQ/VRQ Langlade sheep, clinically affected with scrapie, was homogenized (12.5% weight/volume) before intracerebral inoculation (20 μL) of successive 1/10 dilutions in groups of Tg338 mice (6 or 12 mice). The Infectious Dose 50 of this brain homogenate was determined using a four parameter logistic regression approach, excluding the last point of end titration (no positive animals).

Incubation periods in mice were then plotted on a graph, the different Infectious Dose parameters being calculated for each dilution. A linear regression function was computed using this dataset and then used to estimate the infectious titre (number of Infectious Dose 50) contained in the colostrum and milk samples.

PrP<sup>Sc</sup> Western-blot detection (WB)

A Western blot kit (TeSeE Western Blot, Bio-Rad) was used following the manufacturer’s recommendations.

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Author Contributions

Conceived and designed the experiments: CL, JG OA. Performed the experiments: CL SS SLB JM SM SL FC HC PC DB TM FL CFT NM FS OA. Analyzed the data: CL SS SLB FS JG OA. Contributed reagents/materials/analysis tools: JLW HS. Wrote the paper: CL SS SLB JG OA.

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