RT-QuIC detection of pathological prion protein in subclinical goats following experimental oral transmission of L-type BSE

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Abstract

Objective: The spread of bovine spongiform encephalopathy (BSE) agent to small ruminants is still a major issue in the surveillance of transmissible spongiform encephalopathies (TSEs). L-type bovine spongiform encephalopathy (L-BSE) is an atypical form of BSE with an unknown zoonotic potential that is transmissible to cattle and small ruminants. Our current knowledge of bovine atypical prion strains in sheep and goat relies only on experimental transmission studies by intracranial inoculation. To assess oral susceptibility of goats to L-BSE, we orally inoculated five goats with cattle L-BSE brain homogenates and investigated pathogenic prion protein (PrPsc) distribution by an ultrasensitive in vitro conversion assay known as Real-Time Quaking Induced Conversion (RT-QuIC).

Results: Despite a prolonged observation period of 80 months, all these animals and the uninfected controls did not develop clinical signs referable to TSEs and tested negative by standard diagnostics. Otherwise, RT-QuIC analysis showed seeding activity in five out of five examined brain samples. PrPsc accumulation was also detected in spinal cord and lymphoreticular system. These results indicate that caprine species are susceptible to L-BSE by oral transmission and that ultrasensitive prion tests deserve consideration to improve the potential of current surveillance systems against otherwise undetectable forms of animal prion infections.

Keywords: Prion, L-BSE, RT-QuIC, Goat, Oral transmission, PrPsc, Ultrasensitive detection

Introduction

L-type bovine spongiform encephalopathy (L-BSE) is an atypical form of BSE that is naturally occurring in cattle and experimentally transmissible to small ruminants. Early identification and characterization of TSEs prion strains different from scrapie in sheep and goat is a primary goal of active surveillance as species barrier breakthrough may boost the zoonotic potential of the relevant prion strain. Experimental transmission by the intracerebral route to sheep and goats showed that both species are susceptible to L-BSE with an efficiency of infection as high as 100% and preferential accumulation of PrPsc in the central nervous system (CNS) and to a lesser extent in the
peripheral nervous system, in the cerebrospinal fluid and in the lymphoreticular system (LRS) [1–5].

The intracerebral route offers the possibility to evaluate the species barrier between old and new host, the neuropathogenesis of the disease and the biochemical signature of the prion strain, yet it may not recapitulate the natural history of the disease after oral exposure, by far the main route of possible infection in field conditions.

We here investigate L-BSE transmissibility and PrPSc distribution in orally infected goats.

**Main text**

**Methods**

**Animals**

Seven, six-month-old goats were purchased from herds with no record of scrapie cases. Five animals were Saanen breed and two were crossbred. Genetic analysis to determine PRNP gene polymorphisms and to exclude subjects carrying mutations that could confer resistance against BSE: L168 and M142 [6, 7] or Scrapie (I142M, N146S/D, R154H, R211Q, Q222K) was performed as previously described [4, 8].

**Inoculation of goats**

PrPSc-positive frontal cortex (5 g tissue) from a L-BSE naturally affected cow (IT-141387/02) was ground in a mechanical grinder and homogenized at 30% weight/volume (wt/vol) in sterile buffered saline (PBS). Prior to inoculation, the animals were kept in the new environment for one month for adaptation and were clinically examined to rule out clinical abnormalities. L-BSE inoculum was checked for microbiological sterility and administered to four Saanen and one mixed breed goats (Table 1) by a tube passed through the esophagus down to rumen or abomasum. Two goats were challenged with 15 ml each of PBS and served as controls. The inoculated animals were housed in a bio-safety level 3 containment facility.

**Clinical evaluation**

Daily clinical evaluation was carried out by the animal husbandry staff and the veterinarian. Neurologic examination was performed monthly by a board-certified neurologist. For this purpose, a clinical examination protocol previously used to diagnose scrapie in sheep was applied [9].

**Tissue sample collection**

At various times after exposure, namely 45, 60, 65, 74 and 78 months the animals were anaesthetized with propofol (PropoVet®, Abbott Animal Health) administered intravenously (i.v.), were euthanized with i.v. enbutramide/mebezonium iodide/tetracaine hydrochloride (Tanax®, Intervet Inc. Merck) and were subjected to necropsy. The whole brain, the cerebrospinal fluid (CSF), the entire spinal cord, the LRS (tonsils, submandibular, retropharyngeal, mesenteric and mediastinal lymph nodes), were sampled. Each sample was divided equally: one half was fixed in 10% buffered formalin and the other one was frozen at −80 °C.

**Rapid test**

Diagnosis of TSE was performed on the medulla oblongata firstly by an enzyme-linked immunosorbent assay (ELISA)-based method (IDEXX HerdChek BSE-scrapie antigen test kit EIA rapid test—IDEXX Laboratories).

| ID | Inoculum | Route of infection | Breed | Sex | Genotype | Age* (mo) | SV (mo) | BH PrPSc | PrPSc peripheral distribution |
|----|----------|-------------------|-------|-----|----------|-----------|---------|----------|-------------------------------|
| G1 | L-BSE    | OS                | Mixed Breed | F   | 240 P/P  | 6         | 74      | +        | LNs: retropharyngeal, submandibular, mesenteric; tonsil; thoracic medulla |
| G2 | L-BSE    | OS                | Saanen | F   | 240 P/P  | 6         | 45      | +        | LNs: mesenteric, mediastinal, retropharyngeal, submandibular; tonsil; thoracic medulla |
| G3 | L-BSE    | OS                | Saanen | F   | 240 P/P  | 6         | 65      | +        | LNs: submandibular, mesenteric; tonsil; lombar and sacral medulla |
| G4 | L-BSE    | OS                | Saanen | F   | 240 P/P  | 6         | 60      | +        | LNs: submandibular, mesenteric, mediastinal, retropharyngeal; tonsil; thoracic medulla |
| G5 | L-BSE    | OS                | Saanen | F   | 240 P/P  | 6         | 78      | +        | LNs: mesenteric, mediastinal; tonsil; |
| CTR1 | L-BSE    | IC                | Saanen | F   | 240 P/P  | 6         | 41      | +        | CSF |
| CTR2 | None     | -                 | Mixed Breed | F   | 240 P/P  | 6         | 41      | –        | Negative |
| CTR2 | None     | -                 | Saanen | F   | 240 P/P  | 6         | 74      | –        | Negative |

OS oral administration, IC intracranial, F Female, SV survival time, BH brainstem homogenate, LNs Lympho nodes; mo months, n. a. not available. *Age at inoculation time
followed by a confirmatory western blot method. The CEA internal western blot technique has been described elsewhere [10]. These methods represent the gold standard techniques for making the diagnosis of TSE in CNS samples [11, 12].

Pathology and immunohistochemistry
After fixation, the brain of each goat was coronally cut at the level of obex, medulla, pons, cerebellum, midbrain, diencephalon, and telencephalon. The sections were processed and embedded in paraffin and stained with haematoxylin and eosin (H&E). Immunohistochemical (IHC) analysis and evaluation of PrPsc deposition and PrPsc distribution patterns were performed as previously described [4] at the level of obex, rostral medulla, cerebellar vermis, midbrain, thalamus and frontal area.

Tissues samples preparation for RT-QuIC analysis
Pools of brain areas was obtained mixing same amount (10 mg) of each area separately sampled (brainstem, hypothalamus, cerebellum, basal nuclei, frontal, temporal and occipital cortex). Goat brain homogenates (BHs) (10% wt/vol) were prepared as previously described [5]. Peripheral tissues samples were digested with 0.25% (10% wt/vol) collagenase A (Roche) in PBS/2 mM CaCl2 at 37 °C with 800 rpm overnight continuous shaking, homogenized in Beads Beater and finally clarified at 2000×g for 5 min. The supernatant was collected and frozen at −80 °C.

RT-QuIC analysis
Goat BHs or peripheral tissues homogenates (THs) were serially diluted in 0.1% sodium dodecyl sulfate (SDS, Sigma)–N2 (Gibco)–PBS and subjected to RT-QuIC by using chimeric hamster-sheep (Ha-S) 23-231 and Hamster 90–231 recombinant PrPSEN (rPrPSEN) produced as previously described [13]. The RT-QuIC reaction mix was composed of 10 mM phosphate buffer (pH 7.4), 300 mM NaCl, 10 μM Thioflavin T (ThT), 1 mM Ethylenediaminetetraacetic acid (EDTA), and 0.1 mg/ml of rPrPSEN. Aliquots of this mix (98 μl) were loaded into each well of a 96-well plate with a clear bottom (Nunc) and seeded with 2 μl of 10-3 to 10-4 BH or TH dilutions. Normal goat TH dilutions were used as negative controls, and 10−4 BH dilutions of goat with clinical L-BSE were included as positive controls. The plate was then sealed with a plate-sealer film (Nunc) and incubated at 42 °C or 55 °C in a BMG FLUOstar plate reader with cycles of 1 min of shaking (700 rpm double orbital) and 1 min of rest throughout the incubation. ThT fluorescence measurements (excitation, 450±10 nm; emission, 480±10 nm; bottom read) were recorded every 45 min. RT-QuIC reactions were deemed acceptable when the negative controls remained negative for at least 120 h. RT-QuIC CSF assay was performed as reported previously [5].

Data sets were normalized to a percentage of the maximal fluorescence response of the instrument, and the obtained values were plotted against the reaction times. Data are displayed as the average of four technical replicates.

Results
No animal developed clinical signs of prion encephalopathy prior to euthanasia. All inoculated animals (n=5) and the uninfected controls (n=2) tested negative by standard diagnostics performed on brainstem. Also, histological and immunohistochemical analyses performed on the selected brain macro-areas and on all the peripheral tissues sampled resulted negative.

For ultrasensitive PrPsc detection, brainstem and a pool of different brain areas from each animal were homogenized, serially diluted in logarithmic steps in 0.1% SDS–N2–PBS and subjected to RT-QuIC by using chimeric Ha-S 23-231 and Ha 90-231 rPrPSEN. Five out of five examined brainstem samples showed positive RT-QuIC reactions (Fig. 1A) with stronger fluorescence increase for G4 goat and an average lag phase of 56 h. Furthermore, PrPsc accumulation was also detected in pooled brain samples of four (G1, G2, G3, G4) out of five examined goats. Normal control brain homogenates (NBH) showed no response (Fig. 1B).

To identify brain areas associated to PrPsc deposition, dilutions of multiple brain regions from G1 and G2 goats were tested: we detected a significant increase in fluorescence and shorter lag phase in cerebral cortex and cerebellum samples but not in basal nuclei and hypothalamus (Fig. 1C, D).

By using a modified version of RT-QuIC sample homogenization procedure, we also tested PrPsc distribution in the LRS (tonsils, lymph nodes) and in spinal cord of all L-BSE orally administered goats. The submandibular lymph nodes (LNs) demonstrated seeding reactions in four out of four L-BSE goats tested with an average lag phase of 36 h, whereas mesenteric, mediasinal and retropharyngeal lymph nodes showed seeding activity in 5/5, 3/4, 3/4, respectively, with longer lag phases (Fig. 2). Lower seeding activity was also detected in tonsils of all examined animals (5/5) (data not showed). RT-QuIC analysis of spinal cord also revealed PrPsc deposition in the thoracic segment of 3/5 animals with stronger fluorescence increase for G2 goat. Goat G3 (see Table 1) produced a weakly positive signal in lumbar and sacral gray matter samples. No seeding activity was revealed in cervical gray matter. Further, CSF analyses by second generation of RT-QuIC, performed as previously described [5], did not detect PrPsc in five out of five goats months after oral L-BSE administration. CSF from goat
affected with L-BSE was used as positive control and showed strong fluorescence increase in the first hours of reaction.

Discussion and conclusions
Data here presented indicate that caprine species are susceptible to L-BSE after oral administration and are able to produce very low levels of prions in both lymphatic and central nervous tissues as demonstrated by optimized, high-sensitive, RT-QuIC assay.

At variance with goats intracerebrally infected with L-BSE [4], in this study, no animal developed clinical signs of disease despite prolonged periods of observation, suggesting a comparatively low efficiency of the oral route versus the intracerebral one in L-BSE, a feature that further distinguish this strain from classical BSE [14, 15].

Interestingly, all goats tested negative by standard diagnostics for PrPsc performed on brainstem. This finding, associated with the low amount of PrPsc detected in different brain areas, suggests a partial strain-specific transmission barrier. Indeed, inoculation of a prion into a new host species can produce prolonged incubation periods and/or subclinical infection [16, 17]. In addition, the lack of clinical signs suggests that naturally L-BSE-infected goats may be asymptomatic similarly to what proposed by Okada et al. for oral L-BSE in cattle [17].

In line with previous results [18], RT-QuIC detected lower levels of prions than traditional diagnostic tools. Rapid and confirmatory tests failed to identify any PrPsc in the subclinical animals, while RT-QuIC allowed us to detect misfolded prion protein in multiple brain regions, spinal cord and lymphoreticular system. Studies have established that the rate of fluorescence increase in RT-QuIC, while not measuring infectivity, is directly related to the concentration of prions in the examined tissues and may reassure about the possibility of goat to play as silent L-BSE spreaders in natural conditions. However, we believe that prudence must be always adopted when dealing with the risk of prion spread in field conditions as also suggested by recent data by Denkers and colleagues, who showed that the oral route of infection for chronic wasting disease in deer, may be much more efficient than previously thought [21]. Furthermore, although the mere presence of PrPsc is not indicative of a possible infectivity of the tissue, the finding of positivity in the lymphoreticular muscle must alert
to the potential distribution of PrPsc in peripheral body regions which may increase the risks for humans. Bioassay of infectivity by inoculation of susceptible animals with brains of these goats may help to clarify this issue.

Based on the results achieved with this prion form and also other animal strains, it would be useful to consider the possibility to enlarge current diagnostic criteria to include, in defined conditions (e.g. very limited amounts of source tissue, or preclinical testing), the application of ultrasensitive diagnostic methods. This will not only improve the sensitivity of our surveillance systems but will also help to protect food chain from accidental spillovers of the agent of L-BSE.

Limitations

The primary limitation of this work is that infectivity was not demonstrated by bioassay and the infectious titre was not determined. Therefore, we cannot comment the degree of risk for human.

Despite these limitations, this work specifically demonstrates prion-seeding activity in tissues of goats orally exposed to L-BSE and provide RT-QuIC as useful method to enhance surveillance of TSEs.

Abbreviations

BSE: Bovine Spongiform Encephalopathy; TSEs: Transmissible Spongiform Encephalopathies; L-BSE: L-type Bovine Spongiform Encephalopathy; PrPsc: Pathogenic Prion Protein; RT-QuIC: Real-Time Quaking Induced Conversion; CNS: Central Nervous System; LRS: Lymphoreticular System; PBS: Sterile Buffered Saline; CSF: Cerebrospinal Fluid; H&E: Haematoxylin and Eosin; ELISA: Enzyme-linked immunosorbent assay; IHC: Immunohistochemical; mAb: Monoclonal antibody; Brs: Brain Homogenates; THs: Tissues Homogenates; Ha-S: Hamster-Sheep; rPrPSen: Recombinant prion protein; LNs: Lymph Nodes; wt/vol: Weight/volume; SDS: Sodium Dodecyl Sulfate; ThT: Thioflavin T; EDTA: Ethylenediaminetetraacetic acid; CEA: National reference Center for Animal Encephalopathies.

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Authors’ contributions

AF carried out most of the RT-QuIC experiments, analyzed results, prepared the figures, and wrote the manuscript. MM participated in designing the study, analyzed the data and supervised the execution of some experiments. Sample collection, acquisition, analysis and interpretation of data: AD, GL, CP, LD, GC, EB, MG, TA, EM, LM, PLA, DM, FC, MC, CC, GC was responsible for coordinating research activity, experimental design, data analysis, funding and writing the manuscript. All authors read and approved the final manuscript.

Fig. 2 RT-QuIC detection of PrPsc in different lympho nodes (A submandibular, B mesenteric, C mediastinal and D retropharyngeal) of L-BSE infected goats. Normal control tissue homogenates (black) showed no response. Goat ID are indicated next to the curve. RT-QuIC reactions were seeded with 2 μl of 10–4 LNs homogenate dilution and rHaPrPSen 29-231 substrate was used at 42 °C to detect PrP L-BSE. Each ThT reading is indicated as the percentage of the maximum value achievable by the plate readers as a function of reaction time. The positive threshold was calculated as 10% fluorescence increases of tissue homogenate from normal controls.
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Availability of data and materials
The datasets used in the current study are available from the corresponding author by request.

Declarations

Ethics approval and consent to participate
All procedures involving animals and their care were conducted in conformity with national and international laws and policies (EEC Council Directive 116/92 and 26/2014). The study was approved by the Italian Ministry of Health with authorization number 694/2015-PR of 17th of July 2015.

Consent for publication
Not Applicable.

Competing interests
There is no conflict of interest with any of the authors.

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