Characterization of goat prions demonstrates geographical variation of scrapie strains in Europe and reveals the composite nature of prion strains

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Bovine Spongiform Encephalopathy (BSE) is the only animal prion which has been recognized as a zoonotic agent so far. The identification of BSE in two goats raised the need to reliably identify BSE in small ruminants. However, our understanding of scrapie strain diversity in small ruminants remains ill-defined, thus limiting the accuracy of BSE surveillance and spreading fear that BSE might lurk unrecognized in goats. We investigated prion strain diversity in a large panel of European goats by a novel experimental approach that, instead of assessing the neuropathological profile after serial transmissions in a single animal model, was based on the direct interaction of prion isolates with several recipient rodent models expressing small ruminants or heterologous prion proteins. The findings show that the biological properties of scrapie isolates display different patterns of geographical distribution in Europe and suggest that goat BSE could be reliably discriminated from a wide range of biologically and geographically diverse goat prion isolates. Finally, most field prion isolates showed composite strain features, with discrete strain components or sub-strains being present in different proportions in individual goats or tissues. This has important implications for understanding the nature and evolution of scrapie strains and their transmissibility to other species, including humans.

Transmissible spongiform encephalopathies (TSEs) or prion diseases include fatal neurological diseases of humans and animals which can have different origin and epidemiology, i.e. acquired, sporadic or familial. Prion diseases are caused by a conformational switch of soluble cellular prion protein monomers (PrPC) into misfolded, partially protease-resistant and insoluble prion protein aggregates (PrPSc), which then propagate through an autocatalytic conformational conversion.

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TSEs have been reported in several livestock species, such as small ruminants, bovines, cervids and, more recently, camels. Thus, humans may be exposed to TSE infected tissues of animal origin through the food chain. Some of these TSEs in animals, such as chronic wasting disease (CWD) in cervids and classical scrapie in sheep and goats, efficiently spread by natural transmission and are thus infectious under field conditions. So far, there is no proof of zoonotic transmission of scrapie or CWD, although their zoonotic potential remains incompletely characterized: BSE is the only known animal TSE agent with proven transmissibility to man. The chance that BSE may circulate in sheep and goats led to the need to reliably identify BSE outbreaks in small ruminants. In particular, discriminatory methods able to differentiate between scrapie and BSE have been introduced, leading to the identification of BSE in two goats.

Prions occur as different strains, which are identified by bioassay experiments in rodents. However, the strain properties are thought to depend on the specific conformation of PrPSc and the molecular or immunohistochemical phenotypes in the original host species (often referred to as "types") allow some differentiation. As a notable example, BSE in small ruminants can be discriminated from scrapie by Western blot (WB) analysis of proteinase-K treated PrPSc (PrP<sup>res</sup>)<sup>21–23</sup> and by immunohistochemistry (IHC)<sup>24,25</sup>. Classical scrapie, Nor98 (also referred to as atypical scrapie), CH1641 scrapie and BSE are the TSE types which can be recognized in sheep based on WB PrP<sup>res</sup> banding patterns.

Classical scrapie is the prototypic animal TSE and it is known to be caused by more than one TSE strain<sup>13–19</sup>. CH1641 is a peculiar scrapie strain which has been propagated by experimental transmissions in sheep<sup>20</sup>. CH1641 is characterized by N-terminal PrP<sup>res</sup> features resembling those of BSE<sup>21–23</sup>, but is distinguishable from BSE by discriminatory immunohistochemistry<sup>26</sup> and by improved molecular typing approaches<sup>27–29</sup>. Intensified surveillance in the EU allowed to identify molecular and pathological properties reminiscent of experimental CH1641 in naturally affected sheep and goats<sup>13,15,24,25,27–28</sup>, these cases have been often referred to as CH1641-like scrapie. Finally, Nor98 was first detected in 1998 in Norway as a new form of scrapie characterized by distinctive clinical and pathological features<sup>29</sup>, and has been later recognized as a sporadic, putatively spontaneous or poorly transmissible, form of scrapie affecting small ruminants worldwide<sup>30</sup>.

The PrP<sup>res</sup>-based discrimination scheme is of practical relevance, as it provided the basis for BSE surveillance in small ruminants, but is far from representing the whole variability of scrapie strains. Contrary to Nor98 which represents a single prion strain<sup>31</sup>, several strains of classical scrapie in sheep have been documented in UK and France<sup>16–19</sup>, and one in Italy<sup>32</sup>. Unfortunately, goat scrapie strains have been less investigated and very little knowledge has been gathered on scrapie strain diversity in goats, thus limiting the accuracy and reliability of BSE surveillance in goats. However, this knowledge is necessary both for the ability to properly identify BSE in small ruminants and for the understanding of the zoonotic potential of scrapie.

The overall aim of this project was to characterize goat TSEs in order to i) allow a clear-cut discrimination of BSE in goat and ii) investigate the diversity of goat TSE strains in the field in the EU. We selected a panel of EU goat TSE isolates and characterised their molecular and biological properties in different EU labs. While the detailed molecular features of PrP<sup>res</sup> in the chosen goat TSE isolates panel will be reported elsewhere in parallel<sup>33</sup>, we here report the results of the bioassay experiments. Importantly, all experiments were done with aliquots of the same brain macerates, so as to reduce any sampling bias and to allow proper comparison of different animal models. Thus, we compared the transmission properties of the isolates after intracerebral inoculation in a panel of rodent models able to provide the most broad and sensitive approach for strain typing.

**Results and Discussion**

**Rodent models display differential susceptibility to goat TSEs.** Wild type mice (RIII) and bank voles (Bv109M), as well as several lines of transgenic mice expressing PrP from goat (tg501, hereafter referred to as tg-gtARQ), ARQ-sheep (tg-shpIX, referred to as tg-shARQ), VRQ-sheep (tg338, referred to as tg-shVRQ), bovine (tg110, referred to as tg-bov) or mouse (tg20) were inoculated with the panel of isolates displayed in Table 1. Most of the isolates had the classical scrapie PrP<sup>res</sup> type, hereafter referred to as “21 K”, which can be discriminated from experimental BSE in goat by the lower MM (referred to as “19 K”) and the reduced binding with mAb P4<sup>34</sup>. Two of the isolates from the panel were an Italian goat with atypical scrapie (I15) and an unusual goat case with 19 K PrP<sup>res</sup> originating from a large scrapie outbreak in an English herd<sup>35,36</sup>, later shown to have strain properties distinct from BSE<sup>35</sup> and clearcut CH1641-like biochemical features (UKB2)<sup>35</sup>. The details of rodent models and their overall ability to propagate goat prions are presented in Table 2; the individual attack rates and survival times of all inoculated rodent lines for 33 isolates are reported in Supplementary Table S1. Overall, the transmission rate varied across animal models, as determined by the number of isolates that induced disease in more than 20% of recipient animals (Table 2). Tg-gtARQ and tg-shARQ, which were permissive to all isolates, showed the highest ability to propagate different goat prions, followed by tg-shVRQ. Rodent models expressing bank vole, mouse and particularly bovine PrP were more selective, allowing to propagate only a subset of goat isolates (77%, 72% and 60%, respectively). The overall transmission rate was strongly related to the degree of PrP sequence homology between donor and recipient species, being higher in rodent models expressing small ruminant PrPs than in those with heterologous PrPs. In contrast, the PrP<sup>res</sup> expression level did not give per se a major contribution to the transmission rate, as previously shown by titration experiments<sup>37</sup>. This is well exemplified in tga20 mice, which were as permissive to infection as wild type mice although expressing ten-fold levels mouse PrP<sup>res</sup>. The survival time induced by the different isolates was highly variable, so that in all models either short, around 200 days post inoculation (dpi), or long, >500 dpi, survival times were observed (Table 2).

For the subsequent analyses a subset of 20 isolates was selected (Table 3), excluding those isolates which were known to harbour low levels of PrP<sup>res</sup> or which were not inoculated in all rodent models. This allowed us to reduce confounding factors due to different infectious titres among the isolates and to compare more easily the transmission properties of the different isolates across rodent models. Additionally, tga20 have been excluded from...
Table 1. Geographic origin, PrP genotype and PrP<sup>Sc</sup> type of goat TSE isolates included in the study. *Only amino acid variations are reported, indicated as the codon number followed by the letter codes of the two amino acids at that position. As the P-to-S PrP polymorphism at codon 240 is exceedingly frequent in EU goats, the amino acids at this position are reported for all goats. § The PrP<sup>Sc</sup> type was determined by analyzing the same brain macerates here used for transmission studies<sup>33</sup>. §§ This case was found to have unusual 19 K PrP<sup>Sc</sup> type in the brain<sup>28</sup>, but strain properties distinct from BSE;<sup>35</sup> we later identified CH1641-like PrP<sup>Sc</sup> properties in this isolate<sup>33</sup>. ^ and ^^ respectively isolates deriving from the same farm. ** Experimentally infected with sheep scrapie by the intracerebral route. ° Experimentally infected with goat scrapie by the oral route. # Experimentally infected with cattle BSE by the intracerebral route.

| Country of origin | Original ID | Code | PrP genotype* | PrP<sup>Sc</sup> type§ |
|------------------|-------------|------|---------------|-----------------|
| Italy            | 114921/1/1  | I2   | 240PP         | Classical scrapie |
| Italy            | 121429/1/1  | I3   | 240PP         | Classical scrapie |
| Italy            | 128710/1/1  | I4   | 211QR, 240PS  | Classical scrapie |
| Italy            | 17666/1/1   | I5   | 240PP         | Classical scrapie |
| Italy            | 87016/1/1   | I15  | 154RH; 240PS  | Nor98           |
| Italy            | 85788/1/1   | I7   | 240PP         | Classical scrapie |
| Italy            | 85792/1/1   | I9   | 143HR, 240PS  | Classical scrapie |
| Italy            | 117463/1/1  | I11  | 240PS         | Classical scrapie |
| Italy            | 144508/1/1  | I12  | 240PS         | Classical scrapie |
| Netherlands      | 577227      | N1^  | 143HR, 240PS  | Classical scrapie |
| Netherlands      | 586632-32   | N2^  | 143HR, 240PS  | Classical scrapie |
| Netherlands      | 586632-33   | N3^  | 240PP         | Classical scrapie |
| France           | CP40        | F2** | 240PS         | Classical scrapie |
| France           | 1028        | F3   | 240PP         | Classical scrapie |
| France           | 2119        | F6   | 240PS         | Classical scrapie |
| France           | 2143        | F10  | 240PS         | Classical scrapie |
| France           | 2154        | F11° | 142IM, 240PP  | Classical scrapie |
| France           | 9041        | F14  | 142IM, 240PS  | Classical scrapie |
| France           | 9135        | F16  | 240PS         | Classical scrapie |
| Spain            | C-163P      | S2   | 240PS         | Classical scrapie |
| Spain            | C-645P      | S3   | 240PP         | Classical scrapie |
| Greece           | 1658        | G1   | 240PP         | Classical scrapie |
| Greece           | 1663        | G2   | 240PP         | Classical scrapie |
| Greece           | 1676        | G3   | 143HR, 240PP  | Classical scrapie |
| Greece           | 6989        | G4   | 240PP         | Classical scrapie |
| Cyprus           | Zyp13       | C1   | 240PP         | Classical scrapie |
| Cyprus           | Zyp21       | C2   | 240PP         | Classical scrapie |
| Cyprus           | Zyp27       | C3   | 240PP         | Classical scrapie |
| UK               | BR, G08-1469| UKB2^^| 127GS, 240PP | CH1641-like<sup>35</sup> |
| UK               | BR, G08-1475| UKA2^^| 240PS        | Classical scrapie |
| UK               | BR, G08-1446| UKD2^^| 127GS, 240PP | Classical scrapie |
| UK               | BR, G08-1460| UKC2^^| 240SS        | Classical scrapie |
| France           | 1075        | gbBSE#| 211RQ, 240PS | BSE            |

Table 2. Main features of rodent models and overall outcome of transmission experiments (data are derived from Supplementary Table S1). *number of inocula giving attack rate &gt;20%/number of inocula tested in that particular model.

| Species | Line | Name in the paper | PrP sequence | PrP expression level | % positive transmissions* | Mean Inc. time (range) |
|---------|------|-------------------|--------------|----------------------|---------------------------|------------------------|
| Mouse   | tg501| goat/sheep-ARQ    | 2×           | 100% (26/26)         | 221–>650                  |
| Mouse   | tghshPIX| goat/sheep-ARQ  | 4×           | 100% (32/32)         | 169–531                   |
| Mouse   | tg338| sheep-VRQ         | 8×           | 93% (27/29)          | 173–745                   |
| Mouse   | tg110| cattle            | 8×           | 60% (15/25)          | 195–649                   |
| Mouse   | tga20| mouse PrP<sup>+</sup>| 10×         | 72% (13/18)          | 177–447                   |
| Mouse   | RIIL| mouse PrP<sup>+</sup>| 72% (18/25) | 363–971               |
| Bank vole | Bv109M | vole 109M         | wild type    | 77% (20/26)          | 173–584                   |
Table 3. Transmission features of 20 selected goat TSE isolates in the different rodent models. *Previously reported in;40 in that paper the 154 H/Q genotype of the atypical/Nor98 case I15 was erroneously assigned as wild type.

| TSE isolates | tg-gtARQ | tg-shARQ | tg-shVRQ | tg-bov | RIII | Bv109M |
|--------------|----------|----------|----------|--------|------|--------|
| **Country of origin** | **ID** | **AR** | **ST±SD** | **PrP* type** | **AR** | **ST±SD** | **PrP* type** | **AR** | **ST±SD** | **PrP* type** | **AR** | **ST±SD** | **PrP* type** |
| France | gBSE | 6/6 | 346±16 | 19K | 12/14 | 230±59 | 19/21K | 6/6 | 657±47 | 19K | 6/6 | 260±14 | 19K | 15/15 | 363±27 | 19K |
| Italy | I2 | 6/6 | 526±123 | 21K | 4/11 | 410±28 | 21K | 6/6 | 745±26 | 21K | 1/6 | 453 | 21K | 0/13 | 11/11 | 226±48 | 21K |
| Italy | I3 | 4/4 | 644±14 | 21K | 6/6 | 471±74 | 21K | 6/6 | 641±103 | 21K | 1/7 | 464 | 21K | 0/13 | 11/11 | 222±27 | 21K |
| Italy | I15 | 6/6 | 552±76* | 7K | 4/4 | 354±64 | 7K | 5/5 | 208±39 | 7K | 0/6 | 0/13 | 1/10 | 0/10 |
| Italy | I12 | 4/4 | 591±42 | 21K | 10/12 | 420±99 | 21K | 3/3 | 595±173 | 21K | 0/6 | 0/14 | 10/10 | 173±23 | 21K |
| N-lands | N1 | 4/4 | 437±44 | 21K | 12/13 | 249±13 | 21K | 6/6 | 540±78 | 21K | 3/3 | 649±9 | 19K | 10/10 | 439±15 | 21K | 6/12 | 423±134 | 21K |
| N-lands | N3 | 6/6 | 463±21 | 21K | 11/14 | 270±14 | 21K | 5/5 | 266±79 | 21K | 6/6 | 324/703 | 19K | 15/15 | 460±21 | 21K | 2/9 | 476±894 | 21K |
| France | F2 | 5/5 | 239±21 | 21K | 11/11 | 177±25 | 21K | 5/5 | 173±16 | NA | 4/5 | 342±163 | 19K | 2/13 | 900±900 | NA | 2/2 | 216±624 | 21K |
| France | F3 | 6/6 | 287±14 | 21K | 13/14 | 208±26 | 21K | 6/6 | 224±37 | 19K | 3/6 | 290±48 | 19K | 15/15 | 300±36 | 21K | 5/11 | 526±276 | 21K |
| France | F6 | 4/4 | 468±15 | 21K | 13/14 | 235±37 | 19/21K | 6/6 | 285±124 | 19K | 6/6 | 523±166 | 19K | 13/13 | 471±20 | 21K | 6/11 | 436±252 | 19/21K |
| France | F10 | 6/6 | 449±19 | 21K | 11/11 | 251±65 | 21K | 6/6 | 284±40 | 21K | 4/4 | 556±68 | 19K | 15/15 | 430±11 | 21K | 4/9 | 276±93 | 21K |
| France | F11 | 6/6 | >650 | 21K | 7/9 | 266±36 | 21K | 5/5 | 351±30 | 21K | 0/6 | 0/11 | 417±39 | 21K | 2/11 | 348±674 | 21K |
| France | F16 | 3/3 | 451±22 | 21K | 5/9 | 238±78 | 21K | 6/6 | 413±68 | 21K | 3/6 | 348±58 | 21K | 6/8 | 971±66 | 21K | 15/15 | 193±20 | 21K |
| Spain | S2 | 6/6 | 228±15 | 21K | 14/15 | 222±44 | 19/21K | 6/6 | 198±26 | 19K | 6/6 | 384±149 | 19/21K | 11/11 | 475±31 | 21K | 6/13 | 459±181 | 19/21K |
| Spain | S3 | 6/6 | 221±16 | 21K | 13/15 | 210±32 | 21K | 6/6 | 195±4 | 19K | 5/5 | 271±19 | 19K | 14/14 | 451±11 | 21K | 5/9 | 384±210 | 19/21K |
| Greece | G3 | 4/4 | 466±35 | 21K | 6/6 | 292±53 | 21K | 5/5 | 540±57 | 19K | 0/6 | 0/13 | 477±21 | 21K | 0/4 |
| Cyprus | C1 | 4/4 | 483±15 | 21K | 13/13 | 295±15 | 21K | 5/5 | 453±65 | 19K | 2/2 | >650 | 19K | 15/15 | 437±13 | 21K | 9/14 | 292±86 | 21K |
| Cyprus | C2 | 5/5 | 475±31 | 21K | 13/15 | 279±54 | 21K | 6/6 | 459±70 | 19K | 2/5 | >650 | 19K | 14/14 | 435±8 | 21K | 8/11 | 277±47 | 21K |
| UK | UKB2 | 7/7 | 345±19 | 19/21K | 6/7 | 169±34 | 19/21K | 6/6 | 174±6 | 19K | 7/7 | 205±12 | 19K | 13/13 | 429±33 | 21K | 9/11 | 481±108 | 19/21K |
| UK | UKA2 | 5/5 | 245±36 | 19/21K | 10/10 | 231±59 | 21K | 5/5 | 300±168 | 19K | 5/5 | 255±69 | 19K | 15/15 | 418±22 | 21K | 17/18 | 255±86 | 19/21K |

Categorization of goat TSE isolates shows geographical patterns. The above data clearly suggest that the transmission rate in a panel of different rodent models reflects the biological properties of goat TSE isolates. Furthermore, we found that the survival times varied too, with isolates showing either long or short survival time depending on their specific interaction with the recipient rodents (Table 3). Since both the survival time and attack rate yield information on the transmissibility, we sought to incorporate them in a single parameter as a measure of the transmission efficiency (TE) (see methods).

By drawing graphs with the TE’s of the isolates in different models, it was possible to represent the overall outcome of a given isolate as a transmission profile, which allowed a comparison of the different isolates across all models (Fig. 1). It showed that goat BSE and Nor98 had distinctive transmission profiles, different from each other and from all scrapie isolates (Fig. 1A). Goat BSE was characterized by being the only TSE isolate with lower TE in tg-shVRQ and Bv109M than in any other rodent model. Most importantly, we found that a combination of 3 rodent models, tg-gtARQ or tg-shARQ, tg-shVRQ and tg-bov mice, allowed a clear-cut discrimination of experimental goat BSE from all field scrapie isolates (Fig. 2).
Classical scrapie isolates showed a strong variability of transmission profiles, suggesting the presence of different scrapie strains. Grouping scrapie isolates by country of origin (Fig. 1B) showed that their variability correlates, at least in part, with their geographical origin. In several instances, isolates from the same country had transmission profiles very similar among them and distinct from other countries (see for example graphs for Italy, Cyprus and Spain in Fig. 1B). Notably, the 3 isolates from Italy and the 2 from Spain were sampled in geographically far distant regions, indicating that specific biological properties could be relatively uniform in some countries. In contrast, different transmission profiles were evident among scrapie isolates from France (Fig. 1B). Strikingly, 2 isolates from the same goat herd in the Netherlands and sharing the same PrP\textsubscript{Sc} type had different behaviour when comparing their TE in tg-shVRQ mice (Fig. 1B).

Building on the strong similarities of the transmission profiles of some isolates, we then grouped all natural scrapie isolates by matching their transmission profiles. This resulted into 4 distinct categories of isolates (Fig. 3). TE variation among categories of isolates and rodent models was analysed by ordinary two-way ANOVA, showing significant variation of TEs depending on (i) the rodent model (18.60% of total variation; \( p < 0.001 \)), (ii) the category (22.97% of total variation; \( p < 0.001 \)) and (iii) their interaction (44.75% of total variation; \( p < 0.001 \)).

Category 1 included 3 isolates from Italy and 1 from France which were mainly characterized by a higher transmission efficiency in Bv109M than in any other rodent model. Category 2 comprised 3 isolates from France and 1 from the Netherlands, characterized by having the highest TEs in mice which express small ruminant PrP. Category 3, with 2 isolates from Spain and 2 from UK, was similar to category 2 but had higher TEs in tg-bov; interestingly, this category also included the CH1641-like UKB2 isolate. Finally, category 4 included isolates from Greece, Cyprus and 1 from the Netherlands. This latter category was similar to category 2 but had lower TEs in tg-shVRQ. Category 1 was the most distinct, while more overlap was evident among the other three categories, particularly for TEs in tg-gtARQ, tg-shARQ and RIII mice. Though the transmission profiles of isolates belonging to the same categories were remarkably similar, residual TE variation was evident in some rodent model, such as in Bv109M for categories 3 and 4. This might depend on the relatively coarse analysis that we used to derive the TE values, or might imply further strain variation within some categories.
Figure 2. Discrimination of goat BSE from natural goat TSE isolates based on the transmission efficiency in three rodent models. (A) Graph showing the TE profiles of goat TSE isolates in tg-gtARQ, tg-shVRQ and tg-bov. The TSE isolates are represented with different colours according to their PrPSc type (legend on the right). Note that goat BSE could be discriminated from all scrapie isolates, including the CH1641-like, because of the lower transmission efficiency in tg-shVRQ mice compared with both, tg-gtARQ and tg-bov mice. (B) The same features are shown in B, where tg-shARQ mice are included instead of tg-gtARQ mice.

Figure 3. Categories of natural scrapie isolates according to their TE profiles. All natural TSE isolates except the atypical scrapie I15 were grouped according to the similarity of their TE profiles, resulting into 4 categories. Each graph depicts the TE profiles of the isolates belonging to the same category, as reported on the top of each graph. Individual TSE isolates are referred to as in Table 1.
Overall, these findings show that goat BSE has distinctive biological properties reflecting in a TE profile which can be discriminated from all other goat TSE isolates, including the UKB2 isolate characterised by 19 K PrPSc. On the other hand, the biological properties of natural scrapie isolates appear much variable and consistent with geographical patterns of scrapie strain variation in Europe.

Molecular characterization of PrPSc in rodents reveals the composite nature of scrapie isolates. The biochemical features of the PrPSc conformers propagated in the brain of recipient rodents were investigated by WB analysis. Brain tissues from mice and voles were analysed in different laboratories, each using its own WB method. The details of WB results in the different rodent models are described in Supplementary Results. Overall, rodent models propagated three main PrPSc types, herein referred to as 21 K, 19 K and 8 K based on the apparent MM of their unglycosylated fragment (see Methods paragraph “Definitions and nomenclature”). The most frequent PrPSc type was the scrapie-like pattern 21 K, which was similar to PrPSc detected in goats with classical scrapie (Fig. 4), while 19 K had a BSE-like MM (Fig. 4) and was similarly discriminated by decreased binding with N-terminal antibodies such as 12B2 and P4 (Supplementary Figs S1–S3). The third PrPSc type, 8 K, was detected only in rodent models able to propagate the Nor98 isolate I15, i.e. in tg-gtARQ, tg-shARQ and...
tg-shVRQ mice. They faithfully reproduced the Nor98 molecular signature (Fig. 4), as previously shown\(^{31,39}\). The 8 K PrP\(^{Sc}\) type in tg-gtARQ inoculated with the Nor98 115 isolate had been already reported\(^{40}\).

No obvious variation of PrP\(^{Sc}\) features, such as the degree of glycosylation or the presence of additional PrP\(^{res}\) fragments, has been observed among individual rodents showing 21 K PrP\(^{Sc}\). A thorough examination of PrP\(^{Sc}\) in RIII mice by triplex WB\(^{41}\) (see Supplementary Results) didn’t reveal any variation in the 21 K molecular profile observed after inoculation of isolates with different geographical origin. Furthermore, it showed that all scrapie isolates, including the 19 K UKB2, resulted in 21 K PrP\(^{Sc}\) in RIII mice (Supplementary Fig. S2).

In contrast to RIII mice, the propagation of 19 K PrP\(^{Sc}\) in all other rodent models was not restricted to experiments with goat BSE, as a similar 19 K PrP\(^{Sc}\) type was also derived from the 19 K UKB2 isolate and from a subset of classical scrapie isolates. In most of these cases the scrape-derived BSE-like PrP\(^{Sc}\) type could still be discriminated from BSE PrP\(^{Sc}\). The 19 K PrP\(^{Sc}\) type propagated upon inoculation of scrapie isolates in tga20 (not shown) and tg-bov mice (Fig. 4) was much less glycosylated than 19 K derived from goat BSE, in agreement with findings in the same transgenic mouse line inoculated with sheep scrapie and BSE\(^{42}\). In tg-gtARQ mice, some scrapie isolates induced PrP\(^{Sc}\) with double 19/21 kDa unglycosylated bands, instead of pure 19 K as observed in goat BSE (Fig. 4). In Bv109M, 19 K derived from scrapie cases was frequently, but not invariably, accompanied by an additional 13 kDa C-terminal PrP\(^{res}\) fragment (Supplementary Fig. S4), reminiscent of that observed in sheep with CH1641\(^{25}\). Despite the above-mentioned differences, in Supplementary Table S1, Tables 3 and 4 all PrP\(^{Sc}\) types with a MM of the unglycosylated band BSE-like are indicated as 19 K, whereas 19/21 kDa double PrP\(^{Sc}\) bands and experiments replicating alternatively 21 K or 19 K in individual recipients are indicated as 19 K/21 K.

The PrP\(^{Sc}\) types observed in the different rodent models are summarized in Table 4. Goat BSE induced faithful 19 K PrP\(^{Sc}\) type in all recipient rodents; however, a subset of recipient tg-shARQ and Bv109M also propagated 21 K (2/12 and 4/8, respectively). Concerning Bv109M, this finding is in line with previous data showing that 19 K or 21 K could be selectively propagated in individual voles inoculated with two other BSE-related prions, sheep BSE and variant CJD\(^{43}\). Similar findings have been observed in transgenic mice overexpressing human PrP\(^{Sc}\), in which might suggest that PrP overexpression and/or the mouse genetic background could have putatively driven the emergence of 21 K upon inoculation of BSE in tg-shARQ mice. There was general consensus among the different WB methods and rodent models that PrP\(^{Sc}\) with BSE-like properties can be propagated after inoculation with some goat scrapie isolates. However, the probability for this to occur strongly depended on the rodent model considered, with as much as 76% and 44% of the scrapie isolates propagating 19 K in tg-bov and tg-shVRQ, respectively. In contrast, 13% and 0–8% of the scrapie isolates did so in recipient mice with ARQ or mouse PrP, respectively (Table 4). Furthermore, rodents less able to propagate 19 K from classical scrapie isolates, such as tg-gtARQ and Bv109M, mainly showed mixed 19 K/21 K features. These data strongly suggest that the ability to propagate 19 K from classical scrapie isolates might depend on the PrP sequence of the recipient rodent model. However, other factors such as the PrP expression level\(^{46}\) or background genetic factors\(^{41}\) might have played a role, too.

In accordance with our findings in tg-bov, the emergence of the 19 K PrP\(^{Sc}\) signature has been observed upon transmission of classical scrapie into cattle\(^{47–49}\) and in bovine PrP tg mice\(^{50,51}\). Transmission studies in bank voles suggested that the emergence of 19 K in cattle experimentally infected with scrapie could have been linked to the emergence of CH1641 biological properties\(^{46}\). Thus, bovine PrP might selectively propagate the 19 K component even when present in minimal amounts in the starting inoculum. It is more difficult to explain why 19 K was also propagated in transgenic mouse lines expressing small ruminant PrP sequences, also considering that scrapie cases with 19 K PrP\(^{Sc}\) are rather uncommon in small ruminants. In the case of tg-shVRQ mice, a recent seminal paper showed that the overexpression of PrP determines the selection of the 19 K strain component in these mice\(^{46}\). Furthermore, the 19 K scrapie-derived strain showed strain features indistinguishable from CH1641 in tg-shVRQ mice\(^{46,50}\). Finally, experimental transmission per se could have favoured the propagation of the 19 K component, as it has been shown that experimental challenge of sheep with classical scrapie can lead to the emergence of a 19 K CH1641-like component, which was hidden in the homotypic inoculum\(^{31}\). We then co-analysed the PrP\(^{Sc}\) type and the transmission efficiency in recipient rodent models (Fig. 5). Among all isolates, Goat-BSE only was able to induce 19 K PrP\(^{Sc}\) in all recipient rodent models, while the Nor98 isolate 115 invariably induced the 8 K signature. Among scrapie isolates, only those in category 1 were associated with the propagation solely of the 21 K in all rodent models, while all the others induced also the propagation of 19 K in one or more rodent models. Isolates from categories 2-to-4 invariably propagated 19 K in tg-bov, while their ability to induce 19 K as a minor component in tg-shARQ, tg-gtARQ or Bv109M varied depending on the category.

| Rodent model | PrP\(^{Sc}\) types induced by field TSE cases | % 19 K* | PrP\(^{Sc}\) types induced by goat BSE |
|--------------|------------------------------------------|--------|----------------------------------|
| tg-shARQ     | 21 K/19 K/8 K                            | 13%    | 19 K                            |
| tg-gtARQ     | 21 K/19 K/8 K                            | 13%    | 19 K/21 K                        |
| tg-shVRQ     | 21 K/19 K/8 K                            | 44%    | 19 K                            |
| tg-bov       | 21 K/19 K/8 K                            | 76%    | 19 K                            |
| tga20        | 21 K/19 K/8 K                            | 8%     | 19 K                            |
| RIII         | 21 K                                    | 0%     | 19 K                            |
| Bv109M       | 21 K/19 K/8 K                            | 30%    | 19 K/21 K                        |

Table 4. Summary of the PrP\(^{Sc}\) types reproduced by rodent models (data are derived from Supplementary Table S1). *percentage of transmitted classical scrapie isolates inducing 19 K PrP\(^{Sc}\).
being highest in category 3 and absent in category 4. Thus, the ability to induce 19 K might well be correlated with the biological properties of the isolates.

Furthermore, the emergence of 19 K strongly correlated with TE in tg-bov, i.e. isolates with the highest TE in tg-bov were also those inducing 19 K in rodent models less prone to replicate 19 K, such as tg-shARQ, tg-gtARQ or Bv109M. This suggests that the 19 K component isolated in the different models might all represent the same 19 K strain, supposedly CH1641, which is present as a hidden component in different proportions in isolates from different categories, being highest in category 3, lowest in category 4 and absent in category 1. This is in agreement with the finding that the UKB2 isolate, which is CH1641-like in origin33, was the most able to induce 19 K in recipient rodents, while other isolates in which the CH1641-like component was hidden, i.e. not detected by WB in the isolate, also induced less 19 K in the recipient models.

Although the transmission efficiency allowed to group isolates by their biological properties, the extent to which these categories equate to specific scrapie strains is difficult to assess. Indeed, the operational definition of prion strains relies on the pathological characterization of serially passaged, rodent-adapted isolates. However, natural scrapie isolates can be frequently mixtures of different strain components or sub-strains instead of being uniform strains, such as those which can be selectively propagated after serial experimental passages46,52–54. In this scenario, TE-based categories may not represent distinct strains, but they could rather reflect the presence of different proportions of the same sub-strains, whose effect on TE is amplified by the selective propagation of specific sub-strains in the different recipient rodent models. It can be hypothesized that isolates from categories 2-to-4 contained mixtures of different proportions of 19 K, supposedly CH1641, and one or more 21K-associated strains. Accordingly, the tg-shVRQ and tg-bov datasets explained most of the variability among categories 2-to-4, as their exclusion from the TE matrix strongly weakened the differences among categories 2-to-4 (Supplementary Fig. S5). Thus,tg-shVRQ and tg-bov would have always propagated 19 K because of their preferential selectivity for this strain component, either due to low susceptibility to 21 K or to high susceptibility to 19 K; but they did so with different TE according to the quantity of 19 K in the original isolates. Conversely, rodent models less prone to select 19 K, such as tg-shARQ, tg-gtARQ and Bv109M, mainly propagated 21 K, accompanied by 19 K only when present in the isolates at high levels (i.e. isolates with high TE in tg-bov). In agreement with this interpretation, which implies that 21 K and 19 K are distinct sub-strains pre-existing as a mixture in the original isolates, rodent models less prone to select 19 K propagated 21 K even when inoculated with the 19 K UKB2 isolate, implying that this CH1641-like isolate also contained a 21 K component. Furthermore, individual Bv109M and tg-shARQ mice inoculated with the same scrapie isolate, alternatively propagated 19 K or 21 K, and the UKA2 and UKB2 isolates even induced the propagation of 19 K and 21 K a mixture in tg-gtARQ. Again, this implies that 19 K and 21 K should be considered discrete components, further corroborating our interpretation.

Figure 5. Comparison between PrPSc types and transmission efficiency of goat TSE isolates in different rodent models. Parallel matrices showing the efficiency of transmission (left graph depicting TE values) and the corresponding PrPSc types (right graph) propagated by the goat TSE isolates in different rodent models, as indicated on the top of the graphs. In the left panel, TE values are represented by a colour gradient (legend on the right of the graph). In the right panel, PrPSc types are represented by different colours, according to the legend on the right of the graph. Classical scrapie isolates are ordered according to the TE categories, showing that there is some association between TE-based categories and the emergence of 19 K in one or more recipient models.
isolates from sheep53,56. It has been shown that 19 K and 21 K accumulate at variable levels and proportions in different goats, in different tissues, i.e. the brain and the lymph nodes (LN)23–25. We investigated scrapie outbreak in a UK goat herd, from which the CH1641-like UKB2 and the classical scrapie isolate UKA2 were sampled. In this herd, UKB2 was the only goat out of 71 tested which showed 19 K PrPSc in the brain; interestingly, all goats had a classical scrapie pattern in lymph nodes (LN), including the UKB2 isolate.

Investigation of a UK goat herd suggests CH1641 as a hidden strain component in classical scrapie. Notably, the above-mentioned hypothesis would also explain the reason for the partial variability in TE profiles observed between goats deriving from the same herd (see N1 and N3 in Fig. 1), as it postulates that sub-strain components might coexist in different proportions in individual isolates. To get further insight into this phenomenon in goats exposed to a shared source of infectivity, we extended our studies towards a thoroughly investigated scrapie outbreak in a UK goat herd, from which the CH1641-like UKB2 and the classical scrapie isolate UKA2 were sampled. In this herd, UKB2 was the only goat out of 71 tested which showed 19 K PrPSc in the brain; interestingly, all goats had a classical scrapie pattern in lymph nodes (LN), including the UKB2 isolate.

As shown before, the 19 K UKB2 and the 21 K UKA2 brain isolates were both able to induce either 19 K or 21 K PrPSc in individual Bv109M (Fig. 5). Interestingly, 19 K was detected as the main PrPSc type in 7 out of 9 voles infected with UKB2, but only in 1 out of 15 of those infected with UKA2. Similarly, 7 out of 7 tg-gtARQ mice infected with UKB2 propagated a 19 K component, while only 3 out of 5 of those infected with UKA2 did so (see Supplementary Results). These findings suggest that the proportion of individual rodents propagating 19 K might reflect the presence of different proportions of 19 K in the original isolates. We thus inoculated Bv109M with the brain isolates from 2 additional scrapie infected goats from this herd, and with LN isolates from all 4 goats.

Overall, 3 brain (UKA2, UKB2, UKC2) and 2 LN (UKA2, UKC2) isolates gave positive transmissions in Bv109M, possibly reflecting low PrPSc levels in the pre-clinical donor goats UKB2 and UKD2 (Table 5). The only brain isolate which did not transmit in Bv109M was UKD2 that gave poor or negative transmissions also in the other rodent models, consistent with a very low infectivity titer (Supplementary Table S1). The other 3 brain isolates from 2 additional scrapie infected goats from this herd, and with LN isolates from all 4 goats.

Finally, it is worth to highlight that even after the exclusion of the tg-shVRQ and tg-bov datasets from the TE matrix, the TE profile of category 1 remains distinguishable from all the others (Supplementary Fig. S5). This finding strongly suggests that isolates in category 1, besides being apparently devoid of the the 19 K sub-strain, have distinct biological properties and might even contain one single strain component. Thus, the simplest explanation for our data implies the presence of at least 2 different scrapie strains with different geographical distribution, one of which endows variable proportions of a supposedly CH1641 19 K component.

### Table 5. Transmission in Bv109M of brain and LN TSE isolates from scrapie-positive goats deriving from the same UK goat herd.

| ID | Clinical scrapie | Tissue | PrPSc in the inoculum | Attack rate | Survival time (days ± SD) | 21 K PrPSc (n° voles) | 19 K PrPSc (n° voles) |
|----|------------------|--------|-----------------------|------------|---------------------------|----------------------|----------------------|
| UKA2 | yes | Brain | 21 K | 17/18 | 255 ± 86 | 14 | 1 |
| | | LN | 21 K | 2/5 | 302–638 | 2 |
| UKC2 | yes | Brain | 21 K | 2/9 | 760; 760 | 1 | 1 |
| | | LN | 21 K | 11/12 | 421 ± 197 | 10 | 1 |
| UKB2 | no | Brain | 19 K | 9/11 | 481 ± 108 | 2 | 7 |
| | | LN | 21 K | 0/7 |
| UKD2 | no | Brain | 21 K | 0/11 |
| | | LN | 21 K | 0/6 |

As shown before, the 19 K UKB2 and the 21 K UKA2 brain isolates were both able to induce either 19 K or 21 K PrPSc in individual Bv109M (Fig. 5). Interestingly, 19 K was detected as the main PrPSc type in 7 out of 9 voles infected with UKB2, but only in 1 out of 15 of those infected with UKA2. Similarly, 7 out of 7 tg-gtARQ mice infected with UKB2 propagated a 19 K component, while only 3 out of 5 of those infected with UKA2 did so (see Supplementary Results). These findings suggest that the proportion of individual rodents propagating 19 K might reflect the presence of different proportions of 19 K in the original isolates. We thus inoculated Bv109M with the brain isolates from 2 additional scrapie infected goats from this herd, and with LN isolates from all 4 goats.

Overall, 3 brain (UKA2, UKB2, UKC2) and 2 LN (UKA2, UKC2) isolates gave positive transmissions in Bv109M, possibly reflecting low PrPSc levels in the pre-clinical donor goats UKB2 and UKD2 (Table 5). The only brain isolate which did not transmit in Bv109M was UKD2 that gave poor or negative transmissions also in the other rodent models, consistent with a very low infectivity titer (Supplementary Table S1). The other 3 brain isolates from 2 additional scrapie infected goats from this herd, and with LN isolates from all 4 goats.

Overall, 3 brain (UKA2, UKB2, UKC2) and 2 LN (UKA2, UKC2) isolates gave positive transmissions in Bv109M, possibly reflecting low PrPSc levels in the pre-clinical donor goats UKB2 and UKD2 (Table 5). The only brain isolate which did not transmit in Bv109M was UKD2 that gave poor or negative transmissions also in the other rodent models, consistent with a very low infectivity titer (Supplementary Table S1). The other 3 brain isolates from 2 additional scrapie infected goats from this herd, and with LN isolates from all 4 goats.

Brain and LN isolates from 3 of these goats were also inoculated in wild type mouse lines at APHA, in order to discriminate the unusual CH1641-like case from BSE. In these experiments, strains have been identified by their PrPSc deposition patterns in recipient mice, a method previously shown to allow the detection of BSE in goats and its discrimination from classical scrapie after a single passage in mice55. All isolates transmitted in all wild type mouse lines inoculated, with the exception of LN isolates from UKD2 and UKB2 (Table 6), thus confirming their low infectious titre. Transmissions to mice invariably resulted in the propagation of classical scrapie, which is in line with findings in wild type mice inoculated with isolates from this same herd55 (Table 3) and with CH1641-like isolates from sheep53,56.

This finding can be explained by the results obtained by discriminatory IHC of the donor goat brain. In doing so it should be recalled that an intraneuronal PrPSc accumulation detectable not only by C-terminal but also by N-terminal antibodies is indicative for classical scrapie55. This pattern was indeed detected in the UKB2 brain and was mainly seen in the obex region, the brain area from which the original UKB2 isolate was obtained, while the neurons of the frontal cortex showed a strong CH1641-like deposition (Supplementary Fig. S6). These differences are reflected in the wild-type mice bioassay. Indeed, although both regions produced classical scrapie, the sample from the frontal cortex was much less efficient in all experiments (Table 6). As wild type mice appear unable to replicate the 19 K CH1641-like component, these findings suggest that the lower transmission efficiency of the frontal cortex isolate in mice reflects the lower titre of the 21 K component in the frontal cortex compared to the obex, in agreement with discriminatory IHC findings in the same brain areas.

Overall, the findings obtained with goat isolates from this herd are in agreement and support our previous conclusion that 19 K and 21 K represent sub-strain components pre-existing in the prion isolates. We have also shown that 19 K and 21 K accumulate at variable levels and proportions in different goats, in different tissues, i.e.
CH1641 isolates. Thus, experimental CH1641 would represent an artificially selected extreme phenotype of a few 21 K as visible in rodent bioassays, would represent the most deviant phenotypes in nature. Circulating classical scrapie strain, while the rare CH1641-like natural cases, which still endow significant levels

* Strain identification was based on IHC as described elsewhere.

Table 6. Transmission in wild type mouse lines of brain and lymphoid TSE isolates from scrapie-positive goats deriving from the same UK goat herd. *Strain identification was based on IHC as described elsewhere.

| ID   | Tissue                  | Mouse line | Attack rate | Survival time (days ± SD) | Strain* |
|------|-------------------------|------------|-------------|---------------------------|---------|
| UKA2 | Brain (Obex)            | C57        | 14/14       | 438 ± 20                  | Clas. Scrapie |
|      | VM                      | 15/15      | 542 ± 59    | Clas. Scrapie              |
|      | LN                      | C57        | 20/20       | 514 ± 62                  | Clas. Scrapie |
|      | VM                      | 18/18      | 539 ± 65    | Clas. Scrapie              |
| UKB2 | Brain (Obex)            | C57        | 10/16       | 744 ± 144                 | Clas. Scrapie |
|      | RI III                  | 10/15      | 636 ± 171   | Clas. Scrapie              |
|      | VM                      | 5/15       | 685 ± 137   | Clas. Scrapie              |
|      | LN                      | C57        | 19/20       | 516 ± 49                  | Clas. Scrapie |
|      | RI III                  | 10/11      | 436 ± 110   | Clas. Scrapie              |
|      | VM                      | 18/18      | 566 ± 61    | Clas. Scrapie              |
|      | LN                      | C57        | 5/10        | 660 ± 51                  | Clas. Scrapie |
|      | RI III                  | 3/8        | 485 ± 75    | Clas. Scrapie              |
|      | VM                      | 0/16       |             | Clas. Scrapie              |
| UKD2 | Brain (Obex)            | C57        | 6/11        | 655 ± 42                  | Clas. Scrapie |
|      | VM                      | 4/13       | 820 ± 77    | Clas. Scrapie              |
|      | LN                      | C57        | 0/14        |             | Clas. Scrapie |
|      | VM                      | 0/15       |             | Clas. Scrapie              |

LN vs brain, and even in different brain areas of goats exposed to a common scrapie source. In principle, these observations might be explained by postulating a supposedly rare case of co-infection with 2 different scrapie strains, 21 K and 19 K. However, this seems unlikely as the 19 K/21 K divergence was observed in 16 out of 31 classical scrapie cases deriving from 5 different EU countries (Supplementary Table S1). This situation is similar to that observed in sheep scrapie, where the 19 K/21 K phenotype divergence has been observed with 18 out of 20 geographically unrelated isolates, thus suggesting an ontogenic link, rather than a simple strain mixing.

Importantly, we have also shown that the presence of 19 K depends on the biological properties of the isolates and relates to their geographical origin. Thus, we postulate that 19 K is a standard component of one or more, but not all, 21 K classical scrapie strains circulating in European goats. As in the vast majority of cases the dominant phenotype in the natural host is 21 K, 19 K is mostly present as a minor “hidden” component, which can be only recognized after selective propagation in susceptible hosts, such as tg-bov, tg-shVRQ and, to a lesser extent, Bv109M and tg-mice expressing wild type goat/sheep PrP. In this sense, 19 K might be continuously generated and selected against during scrapie replication in goat tissues, but positively selected during replication in specific rodent models. However, in rare cases 19 K can be positively selected even in its natural host, as shown by the variable presence in individual goats. The observed neurotropism of 19 K relative to 21 K is in line with data showing that the positive selection of 19 K in transgenic mice depends on high expression levels of PrP,

Our conclusions that 19 K is a “hidden” component of a classical scrapie strain allow to reconcile some conflicting data that have been obtained in the past, by postulating that experimental CH1641 and 21 K classical scrapie are actually two sides of the same coin. CH1641 is an experimental scrapie strain that was originally isolated from a British scrapie case and propagated by serial transmission in sheep. Our results may suggest that serial intracerebral transmission in sheep led to positive selection of 19 K over 21 K in CH1641, given the neurotropism of 19 K and evidences that experimental transmission per se could mediate the emergence of CH1641-like properties in recipient sheep. Interestingly, it has been shown that the PrP<sup>ΔC230</sup> type in natural CH1641-like isolates does not fully match experimental CH1641, but shows intermediate properties between classical scrapie and experimental CH1641 suggesting the co-existence of 19 K and 21 K in a mixture. In keeping with these data, 21 K has been selectively propagated in wild type mice inoculated with natural CH1641-like isolates, as shown here and in unrelated experiments, while it has been historically difficult to infect wild type mice with experimental CH1641 isolates. Thus, experimental CH1641 would represent an artificially selected extreme phenotype of a circulating classical scrapie strain, while the rare CH1641-like natural cases, which still endow significant levels of 21 K as visible in rodent bioassays, would represent the most deviant phenotypes in nature.

Conclusions

The conclusions of this work are both, of practical relevance for improving EU surveillance toward a better identification of zoonotic prion strains, and of general relevance for understanding the biological properties of natural TSE isolates. Our study provides a range of rodent models for goat TSEs and BSE discrimination, as well as baseline data from a wide range of goat isolates, both available to scientists for future studies. Most importantly, our findings show that the goat BSE sample here studied was reliably discriminated from a wide range of biologically and geographically diverse goat TSE isolates. The identification of BSE could already be attained after a single passage in rodents, based on the relative transmission efficiency in different rodent models and on the PrP<sup>ΔC230</sup> types associated. Previous studies showed overlapping disease phenotypes in goats with different PrP genotypes and
genetic backgrounds challenged with BSE,33,58 suggesting a low strain variation for BSE in goats and implying that our findings with a single BSE source are well representative of BSE in goats. Thus, our dataset provides the basis for building a reliable and practical approach in terms of discriminatory power and timing of the outcome for discriminating BSE cases from non-zoonotic TSEs in goat. To this purpose, a combination of 3 rodent models, tg-STARQ or tg-shARQ, tg-shVRQ and tg-bov mice, seems to best fulfill the above criteria (Fig. 2).

In this work, we investigated the biological properties of natural TSE isolates by introducing the relative transmission efficiency of TSE isolates across several models, i.e. the TE profiles, as a biological signature of the isolates. This, coupled to PrPSc typing in recipient hosts, allowed to investigate the phenomenon of scrapie strains from a new perspective, i.e. the biological properties of TSE isolates were derived by their specific interaction with different recipient PrP species rather than by isolating rodent-adapted strains by serial passage in a single model. Rodent-adapted strains may stem from the selective amplification of some original strain component in the new host,44 resulting in an incomplete representation of the original biological properties. In contrast, the approach used here may better represent the biological variability within each isolate, as it “reads” its composition through sieves with different mesh sizes, i.e. the various rodent models representing different selective environments. Indeed, thanks to the 19 K molecular signature which allowed to recognize the suggested CH1641 component, we unexpectedly found that a considerable number of classical scrapie isolates from European goats contains a mixture of at least two scrapie strain components or sub-strains, 19 K and 21 K, which co-exist in variable proportions in different individual goats and tissues and can be selectively propagated as discrete components in rodent models. It is conceivable that this phenomenon extends beyond the 19K/21K dichotomy currently detectable by WB, which would suggest that even isolates from which 19 K cannot be derived, such as those in category 1, could be actually composed of a mixture of PrPSc conformers. Detailed histopathological and immunohistochemical strain specific profiles of the ongoing subpassages in rodents may shed more light into it. We conclude that scrapie isolates have strain properties both, composite (i.e. they are mixtures of sub-strains which can be found in different proportions) and discrete (i.e. sub-strains can be isolated as distinct components in rodents), where it could be envisaged that phenomena such as competition or cooperation among sub-strains in different replication environments might play important roles in defining the biological properties of scrapie isolates and their evolution. Such a scenario is in line with the natural history of scrapie being a contagious disease, able to infect individuals with polymorphic PrP sequences and to colonize different tissues and cell types. All these factors represent changes in the replication environment and are likely to drive the emergence of PrPSc conformational variants which are under constant selective pressure as a mixture.46,59,60 Our observation that the variation of TE profiles is partially explained by geographical proximity is in agreement with this scenario.

Finally, this conclusion has obvious impact on the investigation of the zoonotic potential of scrapie, as it is conceivable that the presence and proportion of specific sub-strains could drive the ability of scrapie isolates to cross species barriers. Interestingly, a subset of scrapie isolates studied here was able to adapt to different PrP sequences similarly or even better than BSE.

Materials and Methods
Ethics statement. All experimental protocols were approved by the authors' respective institutions and national competent authorities and all procedures were carried out following ethical review in the authors' respective institutions and in accordance with European Council directives 86/609 and 2010/63, as well as in compliance with the respective national legislations. For the experiments at WBVR, the Netherlands: Dutch Central Authority for Scientific Procedures on Animals, permit number AVD01002016522. At Friedrich-Loeffler-Institute, Germany: the competent authority of the Federal State of Mecklenburg-Western Pomerania, reference number LALLF 7221.3-2.1-012/03 and 7221.3-2.1-027/02. At APHA, United Kingdom: Animal & Scientific Procedures Act, 1986, Home Office License Number 70/6310. At INRA, France: supervision by Institut national de la recherche agronome ethical committee (agreement numbers 02-032-02 for animal care facilities, 92–189 for animal experimentation). At INIA, Spain: Committee on the Ethics of Animal Experiments of the Instituto Nacional de Investigación y Tecnología Agraria y Alimentaria and the General Directorate of Madrid Community Government (Permit Numbers: CEEA 2009/003, CEEA 2011/050, PROEX 263/15). At ISS, Italy: Italian Legislative Decree 116/92 and then 26/2014; protocol approval and supervision by the Service for Biotechnology and Animal Welfare of the Istituto Superiore di Sanità, authorisation by the Italian Ministry of Health (decree number 84/12.B).

TSE isolates from goats. With the aim to characterize the highest biological variability, natural TSE cases were selected in order to encompass the geographical, pathological and genetic variability observed in the field; details on the country of origin, the PrP genotype and the PrPSc type of the selected TSE isolates are reported in Table 1. In doing so, the panel of isolates included 30 brain tissues from naturally TSE-affected goats, along with one isolate from a goat experimentally infected with BSE and two isolates from goats experimentally infected with sheep or goat scrapie (Table 1). Only the three Dutch and the four UK brain samples were derived from animals from single holdings, all the others from different farms. The goat brain samples had initially been found TSE-positive in the EU active surveillance plan with approved screening tests for the detection of TSEs and subsequent scrapie-BSE discriminatory analyses. A detailed analysis by Western blotting (WB) of PrPSc features of the inocula will be reported elsewhere.49 One Italian goat (I15, Table 1) with Nor98/atypical scrapie has already been published before.60 Fifty percent brain macerates in water were prepared under sterile conditions in a microbial safety cabinet with sterile plastic and glass materials. Details, e.g. paper tissues and gloves were changed and the cabinet cleaned between each individual brain sample treatment. Samples were weighed, immersed in an equal part of water, minced with a scalpel blade, and left for 18 hr at 4 °C. Material was ground in a Pyrex glass Dounce and disrupted up to a homogenous paste. Complete homogenisation was obtained by forcing the suspension several times in a ten mL syringe through a 19 G needle. The 50% macerate was aliquoted in 1.5 mL tubes and stored at −80 °C.
Rodent models. The rodent models used in this study are reported in Table 2; most of these models have been previously published and employed for characterizing sheep and goat scrapie. All transgenic mouse lines are on PrP null background and are homozygous for the transgene. In this study, different transgenic mouse lines have been used in order to model small ruminant PrP, expressing either wild type goat PrP (tg501), wild type sheep PrP (tgShpIX) and the most susceptible sheep PrP polymorphic variant (tg338) with the valine to alanine substitution at codon 136 (VRQ, as opposed to the ARQ wild type sequence). Tg501 mouse line expressing the goat wild type PrPSc in homozygosis was obtained by interbreeding the previously described hemizygous tg501 mouse line expressing one fold the PrPSc levels in the goat brain. This lead to homozygosis for the goat PrP transgene within a murine PrP background (PrP mu+/−go+/+). As expected, PrPSc expression levels determined in brain from homozygous tg501 animals was about 2-fold higher than PrPSc levels in goat brain as determined by dilution experiments in WB (data not shown).

Of note, although sheep and goat have different PRNP nucleotide sequences, they share the same PrP amino acid sequence. Hence, the transgenic mouse line used here to model wild type (ARQ/ARQ) sheep PrP, tgShp IX, share the same PrP amino acid sequence with tg501. However, the genetic background is different (B6CBAx129Ola in tgShp IX and FVB in tg501) as is the PrP expression level with 4-8x in tgShp IX and 2x in tg501 as compared to sheep/goat brain respectively. These two lines will be referred to as tg-shARQ and tg-gtARQ in order to emphasize their similarity in terms of PrP sequence.

Bioassay. Aliquots of the TSE isolates were dispatched to all participating laboratories as 50% macerates. Brain macerates were adjusted to 10% (w/v) in phosphate buffered saline (PBS) or 5% glucose. Groups of six-to-eight weeks old rodents were anesthetized and inoculated intracerebrally with 20 μl of brain homogenate. Animals were examined twice a week until neurological signs appeared, after which they were examined daily. Animals were culled with carbon dioxide before neurological impairment was such as to compromise their welfare, in particular their ability to drink and feed adequately, or at the set end point of the experiment. At post-mortem, brains from inoculated mice were removed and divided sagittally. Half brain was frozen for WB analysis of PrPSc and half fixed in formol-saline for histological analysis. The attack rate was calculated as the number of animals scoring positive at post-mortem/number inoculated. Animals found dead or culled for intercurrent disease before 100 days post-inoculation (dpi) and scoring negative at post-mortem were excluded from analyses. The survival time for animals scoring positive at post-mortem was calculated as the time from inoculation to culling or death.

In order to analyse and compare the transmission efficiency of isolates among the various rodent models, we introduced a new parameter, called transmission efficiency (TE), which takes into consideration both the attack rate and the survival time. To do so, mean survival times were categorized into 6 categories according to the observed overall variation (see SI Table): those with the shortest mean survival time (<200 dpi) were assigned the highest TE value (ST = 1); all others experiments having mean survival times >200 dpi were divided into 5 categories by increasing steps of 100 dpi. Then, equally spaced ST values were assigned to these categories by decreasing TE of 0,17 for each step (which represent 1 divided by the number of categories). In summary, each category was assigned to a survival time value (ST) between 0 and 1, as follow: 1,00 for survival times <200 dpi; 0,83 for survival times between 200 and 299 dpi; 0,67 for survival times between 300 and 399 dpi; 0,50 for survival times between 400 and 499 dpi; 0,33 for survival times between 500 and 599 dpi; 0,17 for survival times >600 dpi. TE was then calculated by multiplying the attack rate and ST values.

Molecular analysis of PrPSc in rodent models by western blotting. In all rodent models the detection and typing of PrPSc was performed by WB analysis of protease K-treated PrPSc in brain homogenates. Each laboratory used its own WB method and typing strategy, which are described in detail in Supplementary Methods.

Pathology and Immunohistochemistry. At APHA, wild type mice (C57/BL6 and VM) were challenged with brain tissue or mesenteric lymph node from three of the four UK isolates (G08-1469, G08-1475, G08-1446). In these experiments, TSE diagnosis was based on the detection of PrPSc using IHC.

Fixed brains were cut at 4 different coronal levels to reveal frontal cortex, hippocampus with thalamus, midbrain, and brain stem with cerebellum. Histological sections from the above levels were prepared and stained either with H&E to assess spongiosis or were subjected to immunohistochemistry using the polyclonal antibody Rb486 as described.

Discriminatory IHC in goat tissues was performed according to published protocols.

Definitions and nomenclature. We will use PrPSc as a general term for indicating pathological aggregates of misfolded PrP; the term PrPres will be used specifically to refer to the protease-resistant core of PrPSc.

Different types of PrPSc are discriminable by molecular methods, yielding different PrPres cores. In WB, PrPres appears usually as three protein bands with apparent molecular masses between 17 and 30 kDa; the top, middle and lower one are respectively diglycosylated, monoglycosylated and non-glycosylated forms of the same amino acid sequence. An exception is atypical/Nor98 scrapie which has variably processed PrPres bands, of which a low apparent molecular mass (MM) of the unglycosylated PrPres fragment, i.e. 21 K or 19 K.

The apparent MM of the unglycosylated PrPres fragments varied with the different WB methods used here, owing to the different gels and molecular markers used. For example, classical scrapie and BSE were measured as 21 kDa and 20 kDa, respectively, in RIII mice; as 19 kDa and 18 kDa in tg-shARQ, tg-goat and tg-bov; and as 18 kDa and 17 kDa in Bv109M. To address this issue, and considering that scrapie and BSE have been historically mostly referred as 21 kDa and 20 kDa, respectively, we introduced a common nomenclature to refer to the different PrPSc types as follows: 19 K for BSE-like PrPSc; 21 K for scrapie-like PrPSc and 8 K for Nor98-like PrPSc.
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Author contributions
R.N., U.A., J.M.T. and J.P.M.L. conceived and designed the study. A.M.M., J.C.E., C.F., L.V.K., J.S., I.L., O.A., L.P., M.A.D.B., P.A.C., J.G.J., G.V., C.D.A., B.C., F.L., J.P.M.L., J.M.T., M.H.G. and R.N. conducted the experiments. A.M.M., J.C.E., C.F., R.N., J.S. and J.P.M.L. prepared the figures. T.S., P.P.S., J.P.M.L., P.L.A., C.A. and O.A. selected and provided samples. R.N. drafted the manuscript. All authors discussed the results and critically reviewed the manuscript prior to submission.

Competing interests
The authors declare no competing interests.
