Increased Susceptibility of Human-PrP Transgenic Mice to Bovine Spongiform Encephalopathy Infection following Passage in Sheep

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Increased susceptibility of human-PrP transgenic mice to bovine spongiform encephalopathy following passage in sheep.

Running title: Sheep BSE transmission to human-PrP transgenic mice

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Abstract

The risk of transmission of ruminant transmissible spongiform encephalopathy (TSE) to humans was thought to be low due to the lack of association between sheep scrapie and incidence of human TSE. However a single TSE agent strain has been shown to cause both bovine spongiform encephalopathy (BSE) and human vCJD, indicating that some ruminant TSEs may be transmissible to humans. While the transmission of cattle BSE to humans in transgenic mouse models has been inefficient, indicating the presence of a significant transmission barrier between cattle and humans, BSE has been transmitted to a number of other species. Here we aimed to further investigate the human transmission barrier following passage of BSE in a sheep. Following inoculation with cattle BSE, gene targeted transgenic mice expressing human PrP showed no clinical or pathological signs of TSE disease. However following inoculation with an isolate of BSE that had been passaged through a sheep, TSE associated vacuolation and proteinase-K resistant PrP deposition were observed in mice homozygous for the codon 129-methionine PRNP gene. This observation may be due to higher titres of the BSE agent in sheep, or an increased susceptibility of humans to BSE prions following passage through a sheep. However these data confirm that, contrary to previous predictions, it is possible that a sheep prion may be transmissible to humans and that BSE from other species may be a public health risk.
Introduction.

The transmissible spongiform encephalopathies (TSEs) are a group of fatal infectious neurodegenerative diseases that include scrapie in sheep, bovine spongiform encephalopathy (BSE) in cattle, and Creutzfeldt-Jakob disease (CJD) in humans. TSEs are characterised by the accumulation in the brain of PrP$^{\text{Sc}}$, which is a conformational variant of the normal cellular host prion protein (PrP$^{\text{C}}$). The abnormal form of the protein is protease resistant, detergent insoluble and aggregates in diffuse or amyloid deposits in the central nervous system (CNS) and lymphoreticular system of infected animals. TSEs are infectious diseases, and can be transmitted between animals of the same and different species by a number of routes including oral, environmental, or iatrogenic exposure. The host range is a specific characteristic of each strain, but TSE agents usually transmit more readily within rather than between species. Low transmission rates are often observed upon transmission to a new species, but on further passage in the new species increased transmission rates and shorter incubation times are usually observed. This effect is referred to as the species barrier. The ability of individual TSE agents to cross a species barrier can be examined experimentally by direct inoculation of different species, or modelled in transgenic mice expressing PrP sequences from these species. Modelling species barriers in mice is particularly important when assessing the risks of infection in humans. Such experiments can assess risk posed by different TSE agents, and also the potential for mutation and adaptation of the agent to the new species. These experiments can highlight changes that may result in the emergence of a new agent strain with a much wider or unknown host range.
One TSE agent which has shown the ability to transmit to several different species is BSE, where infection has been observed in captive and domestic feline species, and exotic ungulates (kudu & nyala) probably due to ingestion of contaminated feed (13). In 1996 a new variant form of CJD (vCJD) was reported in humans which presented with unusual pathology and an extremely young age range compared to other human TSEs (30). Strain typing experiments demonstrated that vCJD was caused by the same agent strain as BSE (9, 23), indicating that exposure to contaminated foods may have also resulted in transmission of BSE to humans. Humans had previously been thought to be at low risk from contracting ruminant TSEs, as sheep scrapie has been endemic in many countries for hundreds of years without any related foci of human TSE. However the link between BSE and vCJD proved that ruminant TSEs are a public health risk, and that new ruminant TSEs may potentially be transmissible to humans.

During the BSE epidemic, sheep were undoubtedly exposed to the BSE agent, however no cases of BSE in sheep have been documented in the field. Sheep can however be infected with BSE via oral, intravenous, and intracerebral routes (17, 18), producing clinical TSE with incubation periods ranging from months to years (depending on the PrP genotype of the sheep), proving that this species is susceptible to infection with the BSE agent. It is possible that low level BSE infection did exist in sheep during the height of the BSE epidemic, however it is unknown whether this could have been masked by coinfection with scrapie. BSE has however been documented in goats (14, 25), providing evidence that infection of small ruminants in the field has occurred. Whether the presence of BSE infection in sheep and goats would represent a risk to humans is
currently unclear. No previous association between sheep scrapie and human TSE has
been documented. This may be due to incompatibility between sheep and human PrP, or
the inability of natural scrapie strains to replicate efficiently in a human host. If the agent
strain is ultimately responsible for this lack of transmission rather than a sheep/human
barrier, it is possible that BSE in other ruminant species may pose a risk to humans.

In order to address the transmissibility of cattle and sheep derived BSE to humans we
performed inoculations of BSE infected sheep brain into a panel of gene targeted
transgenic mice expressing human PrP under the same spatial and temporal controls as
wild type PrP (4). Three lines of transgenic mice were used representing the genetic
diversity in the human population due to the PrP codon 129 methionine/valine
polymorphism; HuMM (40%), HuMV (50%) and HuVV (10%). This polymorphism is
known to affect human susceptibility to TSE, and to date all confirmed clinical cases of
vCJD have occurred in individuals who are methionine homozygous at PrP codon 129.
Although previous experiments showed no disease transmission from cattle BSE in any
of these human transgenic mouse lines (4), we show here that experimental sheep BSE
produced pathological evidence of disease transmission in ~70% of HuMM transgenic
mice, suggesting that sheep BSE may be a greater risk to humans than cattle BSE.
Materials and Methods

Transgenic Mice

Inbred gene targeted human PrP transgenic mice with the codon-129 methionine/valine polymorphism have been described previously (4). Transgenic lines homozygous for the polymorphism were crossed to produce all three genotypes represented in the UK population (designated HuMM, HuMV, and HuVV, respectively) as previously described (4). Brain tissue from a group of uninoculated HuMM and HuVV mice that had previously been allowed to age to over 700 days were utilised as controls in immunohistochemical analyses. In addition a gene targeted bovine transgenic line expressing bovine PrP with the 6-octapeptide repeat region (Bov6), and wild-type 129/Ola mice were used as control lines (4). The Bov6 gene targeted transgenic line is the same line described as “BovTg” in Bishop et al, 2006 (4).

Preparation of Inocula

Brain tissue from the cortex of a female, Cheviot sheep (NPU J2501; ARQ/ARQ) experimentally infected via the oral route with cattle BSE (19) was used to prepare 2 separate inocula (inoculum-1 and inoculum-2). The sheep was culled with confirmed clinical and pathological BSE at 596 days post inoculation. Natural scrapie isolates (VRQ/VRQ) from the NPU flock were used as controls. Inocula were prepared from cortex tissue in sterile saline at a concentration of 10% (w/v). The cattle BSE brainstem pool used in comparative experiments was supplied by the Veterinary Laboratories Agency, Weybridge UK (infectivity titre 10^{3.3} ID_{50} units/g tissue, measured in RIII mice; M Simmons & R Lockey, personal communication).
Inoculation of transgenic lines.

Experimental sheep BSE inoculum-1 was used to infect groups of gene targeted transgenic mice expressing human PrP with the codon 129 methionine/valine polymorphism (HuMM, HuVV, HuMV), control 129/Ola mice, and gene targeted Bov6 mice. In a later experiment, the same transgenic panel was inoculated with experimental sheep BSE inoculum-2 to confirm data obtained with inoculum-1. Data shown (for comparison) from inoculation of HuMM, HuMV, HuVV and Bov6 transgenic mice with a cattle BSE brainstem pool (provided by the Veterinary Laboratories Agency, Weybridge, UK) was generated and published previously. The 129/Ola wild type mice inoculated with cattle BSE described here were inoculated in the same manner with the same BSE brainstem pool as part of this work, but were not included in the original publication.

All mice were intracerebrally (i.c.) inoculated with 0.02ml of inoculum per mouse into the right mid-hemisphere. Following the inoculation, mice were monitored daily and scored once a week for signs of clinical disease. Mice were culled at a pre-defined clinical endpoint, or due to welfare reasons, and brain tissue recovered at post mortem. One half of the brain was fixed in formal saline, further trimmed to expose a number of different regions of the brain (frontal cortex, cortex, hippocampus, thalamus, cerebellum and brain stem) then wax embedded to allow 6µm sections to be cut for use in pathological analysis of the tissue. The second half of each brain and the spleen (where available) were snap frozen in liquid nitrogen for biochemical analysis. Each mouse was genotyped post-mortem to confirm PrP genotype. All mouse experiments were reviewed
and approved by the Local Ethical Review Committee, and performed under license from
the UK Home Office in accordance with the UK Animals (Scientific Procedures) Act
1986.

**Analysis of vacuolar pathology**

Sections were cut (6µm) from each mouse brain and stained using haematoxylin and
eosin (H&E). Nine regions of the grey matter (dorsal medulla, cerebellar cortex, superior
colliculus, hypothalamus, thalamus, hippocampus, septum, cerebral cortex, and forebrain
cerebral cortex) and three regions of white matter (cerebellar white matter, midbrain
white matter and cerebral peduncle) were examined and scored on a scale of 0 (no
vacuolation) to 5 (severe vacuolation) for the presence and severity of vacuolation. Mean
vacuolation scores for each mouse group in each experiment were calculated and plotted
with standard error of mean (SEM) against scoring areas to produce a lesion profile (21).

**Immunohistochemical analysis of PrP deposition in brain**

To identify PrP deposits in the brain, two methods of immunohistochemical (IHC)
analysis were employed. (i) ABC kit (Vectastain); brain sections (6µm) were pre-treated
using hydrated autoclaving at 121°C for 15 min and exposure to formic acid (95%) for
5min prior to incubation with 0.44µg/ml anti-PrP monoclonal antibody 6H4 (Prionics) at
room temperature overnight. Biotinylated secondary anti-mouse antibody (Jackson
Immuno Research Laboratories, UK) was added at 2.6µg/ml and incubated at room
temperature for 1h. PrP\text{Sc} was visualised by a reaction with hydrogen peroxidase-
activated diaminobenzidine (DAB). (ii) When levels of PrP\text{Sc} detected using the ABC kit
were low or zero, the DAKO Catalysed Signal Amplification kit (CSA II (Vectastain)) was used. IHC was performed using the same principals as in (i), but with an additional streptavidin-biotin-peroxidase amplification step (see manufacturer’s information). Sections were pre-treated as above prior to incubation with 0.44µg/ml monoclonal antibody 6H4 at room temperature overnight. Anti-mouse immunoglobulins, supplied with the CSA II kit, were added and incubated for 60min at room temperature. PrP<sub>Sc</sub> detection by a reaction with hydrogen peroxidase-activated DAB.

**Immunohistochemical detection of glial activation**

To detect astrocyte activation, brain sections (6µm) were incubated with 1.45µg/ml anti-glial fibrillary acidic protein (GFAP, Dako UK Ltd) antibody at room temperature for 1h. To detect microglia activation, brain sections (6µm) were pre-treated using hydrated microwaving for 10min prior to incubation with 0.05µg/ml anti-Iba1 antibody (Wako Chemicals GmbH) at room temperature for 1h. For both primary antibodies, a biotinylated secondary anti-rabbit antibody (Jackson Immuno Research Laboratories, UK) was added at 2.6µg/ml and incubated at room temperature for 1h. Astrocytes and microglia were visualised by a reaction with hydrogen peroxidase-activated DAB.

**Detection of amyloid plaques by thioflavin fluorescence**

Sections (6µm) were processed and exposed to 1% thioflavin-S (Sigma, UK) solution as described previously (28). Viewed under a fluorescence microscope amyloid deposits fluoresce bright green.
Identification of PrP\textsuperscript{Sc} in spleen tissue

Spleen tissues that were available from HuMM transgenic mice inoculated with experimental sheep BSE inoculum-2 were screened by the IDEXX HerdChek assay following the manufacturers’ guidelines. Buffer volumes for homogenisation were adjusted to ensure that all homogenates were 30\% (w/v) for consistency.

Identification of PrP\textsuperscript{Sc} by immunoblot

Brain tissues from HuMM transgenic mice, Bov6 transgenic mice and wild type 129/Ola mice inoculated with experimental sheep BSE inoculum-2 were prepared for analysis by immunoblot. Samples from the experimental sheep BSE source brain and from isolates of natural scrapie were also prepared as controls. Due to the focus of PrP deposition and the amyloid nature of much of the disease associated PrP in HuMM transgenic mice, a centrifugal concentration extraction procedure (including proteinase K digestion) was performed as described previously (24) to maximise the possibility of identifying PrP-res.

Samples were loaded and run on 16\% Tris-Glycine acrylamide gels (Novex, Invitrogen) at varying concentrations (0.6mg – 64mg brain equivalent) to allow comparison between lanes, and immunoblotted onto PVDF membrane. To achieve a detectable signal, approximately 64mg brain equivalent was loaded from HuMM brain tissue, compared to 0.88mg brain equivalent from experimental sheep BSE infected Bov6 controls.

Monoclonal antibodies 6H4 (0.1\(\mu\)g/ml) and 12B2 (0.2\(\mu\)g/ml) were used to detect PrP, and bands were visualised using HRP labelled anti-mouse secondary antibody (Jackson Immuno Research Laboratories, UK) and a chemiluminescence substrate (Roche)
Results.

BSE strain characteristics are retained following transmission in sheep.

For both experimental sheep BSE inoculum-1 and inoculum-2, 100% transmission rates were observed in 129/Ola and Bov6 control mice. Experimental sheep BSE inoculum-1 produced incubation times in 129/Ola and Bov6 mice of 474 ±22 days and 564 ±8 days respectively (Table 1), similar to those observed in a previous experiment following inoculation of these lines with a cattle BSE brain pool (4). Inoculum-2, prepared from the same BSE infected sheep, gave incubation times of 403 ±17 days and 487 ±3 days in 129/Ola and Bov6 mice respectively (Table 1). In each line of mice, the lesion profiles of cattle and sheep BSE were similar indicating no change in the targeting properties of BSE following passage through sheep (Figure 1). Lesion profiles of sheep BSE inoculum-1 and inoculum-2 were also similar, although the degree of vacuolation was slightly reduced for inoculum-2. This may represent the shortened incubation times observed in these mice. Although these shortened incubation times may reflect a higher level of agent replication in the tissue sample used to prepare inoculum-2, overall the incubation time ratio and targeting of cattle BSE and sheep BSE in control mice indicates no major change in agent characteristics following passage in sheep.

Susceptibility of human PrP transgenic mice to experimental sheep BSE.

Following inoculation with experimental sheep BSE inoculum-1, three human transgenic mice (1 x HuMM, 1 x HuVV, 1 x HuMV) were scored as showing clinical signs of TSE disease at 449, 609 and 707 days post inoculation respectively, but had no confirmatory vacuolar pathology in the brain. All other human transgenic mice showed no clinical
signs of TSE disease and no TSE associated vacuolar pathology (Table 1). This agreed
with previous data generated following inoculation of these human transgenic mouse
lines with cattle BSE (4), indicating the presence of a significant transmission barrier to
the BSE agent in humans. However, in order to rule out the presence of subclinical
disease, the oldest mice were screened for abnormal PrP deposition by
immunohistochemistry (IHC) using the anti-PrP antibody 6H4. Of all animals screened,
one HuMM mouse which was culled due to old age (706 days post inoculation) showed a
small focus of PrP deposition in the thalamus (Figure 2). No other deposition was
observed in this animal, or any of the other animals examined.

In order to confirm this observation, a second separate sample was taken from the same
BSE infected sheep brain to prepare inoculum-2, which was used to infect a repeat panel
of human transgenic mice. In this experiment, 4 mice (2 x HuVV and 2 x HuMV) were
scored as showing clinical signs of TSE disease (between 494-539 days post inoculation),
but showed no confirmatory TSE associated pathology in the CNS (vacuolation or PrP
deposition). All other human transgenic mice were negative for clinical signs of TSE
disease (Table 1). On pathological examination, all HuVV and HuMV mice were
negative for vacuolar pathology and PrP deposition. However, 16/23 HuMM mice
contained abnormal PrP deposition in the brain, and 3 of the 16 scored positive for
vacuolar pathology (Tables 1 and 2). PrP deposition was present in 50% of HuMM mice
culled (due to welfare reasons) between 377 and 589 days post inoculation (n=14), and in
all of the remaining HuMM mice culled between 600 - 750 days (n=9). Variable patterns
and levels of PrP deposition were observed in HuMM mice (Table 2, Figure 3 and Figure
Staining was evident in the cochlear nucleus (when identifiable in the tissue section) as punctate-like deposits, in a pattern similar to targeting of BSE in wild type mice (Figure 3a) (9). Punctate perineuronal and intraneuronal staining (Figure 3b & 3c) was also evident in the midbrain and areas of the thalamus. Most strikingly, thioflavin-S fluorescent PrP amyloid plaques were present in the hippocampus, corpus callosum, dorsal lateral geniculate nucleus (dLGN) and thalamus of 8/16 mice positive for TSE pathology (Figure 4h and Supplemental Information: Figure S1). The 8 mice showing thioflavin-S fluorescence were all over 589 days post inoculation (Table 2), indicating that plaque formation was occurring later in the disease process, and that PrP deposition in animals culled prior to 589 days was not amyloid. The PrP plaques were shown to be bi-lateral in a number of tissues where whole brains, or significant portions of the contralateral half had been analysed (Supplemental Information: Figure S2). Florid plaques (associated with vCJD disease in humans) were also evident in the hippocampus (Figure 3d). The vacuolar pathology observed in 3 of the PrP-positive HuMM mice was extremely limited, with the main focus of vacuolation occurring in the hippocampus and thalamus, closely linked with the presence of PrP amyloid plaques (Figure 4d). No PrP deposition was observed in aged (750 day old) uninoculated HuMM and HuVV mice that were stained with 6H4 to control for possible age related PrP deposition due to the transgene (Figure 4c).

As PrP deposition in HuMM transgenic mice was focused to specific brain areas, PrP\textsuperscript{Sc} had to be concentrated by centrifugal purification (SAF prep) to visualise by immunoblot. PrP\textsuperscript{Sc} was extracted from a HuMM transgenic mouse (with PrP deposition; same animal...
as shown in Figure 4b, d, f, h), a Bov6 transgenic mouse and a wild type mouse
inoculated with experimental sheep BSE inoculum-2 and analysed by immunoblot
alongside PrP\textsuperscript{Sc} extracted from the source BSE inoculated sheep, and four scrapie
controls. Blots probed with MAb 6H4 revealed a low level of Proteinase K-resistant PrP
(PrP-res) in the HuMM mouse compared to Bov6 and 129/Ola controls (Figure 5a). The
HuMM sample represented approximately 1/6\textsuperscript{th} brain equivalent, reflecting the foci of
deposition in the original tissue (Figure 4d) and the amyloid nature of much of the
depositd PrP, which may not have been resolved in the polyacrylamide gel. PrP-res
levels were too low in both the HuMM mouse and the original source BSE infected sheep
brain to determine the size of the low molecular weight PrP-res band. However when the
gel was re-probed with the N-terminal MAb 12B2, reduced levels of staining were
observed in lanes containing the source BSE infected sheep brain, and brain homogenate
from the HuMM mouse, 129/Ola mouse and Bov6 mouse infected with experimental
sheep BSE (Figure 5b). Such reduced staining with MAb 12B2 is characteristic of BSE
infection (27, 31)

**Glial Activation in infected HuMM transgenic mice**

Brian sections from selected HuMM, HuVV and HuMV transgenic mice inoculated with
experimental sheep BSE inoculum-2 were screened for activation of astrocytes (anti
GFAP) and microglia (anti-Iba1). Aged uninoculated control HuMM and HuVV mice
(\approx 750 days) showed modest GFAP and Iba1 immunoreactivity in isolated cells with
slender processes (Fig. 4a,e). PrP amyloid deposits were not seen in these animals (Fig.
4c, g). In contrast, astro- and micro-gliosis were observed in HuMM mice inoculated with
sheep BSE inoculum-2 in brain areas with abundant PrP amyloid deposition e.g., the lateral geniculate nucleus (LGN) (Figure 4b, d, f, h). Gliosis was not detected in experimental sheep BSE inoculated HuMV and HuVV mice (data not shown).

Peripheral accumulation of PrPSc in spleen

The IDEXX HerdChek assay was used to analyse spleen tissue available from 17 of the 23 HuMM mice inoculated with experimental sheep BSE inoculum-2. Positive assay readouts were obtained for 10 of the 17 spleens analysed (Table 2). The detection of abnormal PrP in spleen tissue by the IDEXX HerdChek assay did not correlate with PrP deposition observed in the brain of the same animal. Of the 10 IDEXX positive samples, three were from mice in which the brain was scored negative by IHC with anti PrP antibody 6H4 (culled at 422 and 658 days post inoculation). In contrast, the three oldest spleen samples available (623, 687 and 750 days post inoculation) were all negative, despite moderate PrP deposition in the brain (Table 2).

Susceptibility of human transgenic mice to natural sheep scrapie.

Brain homogenate from two VRQ/VRQ sheep with clinically and pathologically confirmed scrapie was used to inoculate the transgenic mouse panel. For both isolates, the presence of clinical and pathological signs of TSE disease were observed only in some control 129/Ola wild type mice with long incubation times (Table 1), as has been observed previously in C57 and RIII mice (8). Clinical signs were recorded in several other mice (2 x Bov6 mice, 8 x HuMV and 1 x HuVV mouse), but none of these animals showed signs of TSE associated pathology. No clinical signs of disease were observed in
any remaining transgenic mice, and animals were either culled for intercurrent disease or
due to old age. No evidence of TSE associated vacuolar pathology or PrP deposition was
observed in the human or bovine PrP gene targeted transgenic lines following
pathological examination of brain tissue (data not shown).
Discussion

While transmission of BSE from cattle to humans via oral exposure has been proposed as the origin of vCJD, the risk posed to humans from BSE infection in other species is currently unknown. Sheep can be experimentally infected with BSE, and it has long been a concern that sheep may have become infected during the BSE epidemic. Although no evidence of such infection has been identified in the field, cases of BSE in goats have been reported (14, 25). Here we have shown that inoculation of experimental sheep BSE into gene targeted HuMM transgenic mice resulted in the identification of TSE related pathology (PrP deposition, vacuolation, and gliosis) in ~70% of the animals overall, and 100% of HuMM mice surviving over 600 days. PrP^Sc was detected in brain tissue by immunoblot, and in spleen tissue of several mice using the IDEXX HerdChek assay. Some of the oldest HuMM mice (623-750 days post inoculation) that showed PrP deposition in the brain did not have detectable PrP^Sc in spleen. This variability is not unusual, as extremely old mice that show PrP^Sc in the brain often have no corresponding deposition in the spleen. This is likely due to loss of germinal centres in the spleen caused by aging (7). No evidence of disease transmission was observed in HuMV or HuVV mice, mirroring the prevalence of vCJD disease observed to date in the UK population. Although the presence of disease pathology indicates agent replication and early phase disease, we cannot predict whether these mice would have developed clinical disease if lifespan had been extended, or if this represents a persistent sub-clinical state. Subpassage of brain material from HuMM mice will be performed to confirm agent replication, and assess adaptation and host range of the agent.
Our results cast new light on the existing data concerning BSE transmission to humans. Since the identification of the link between BSE and vCJD, many studies have been performed to demonstrate or model the transmission of BSE to humans using \textit{in vitro} conversion techniques or by inoculation of transgenic mice expressing human PrP. In these transmission studies, cattle BSE has shown limited transmissibility to human PrP transgenic mice (~0-30%), and considerable variation in susceptibility exists between different transgenic lines with varying constructs and protein expression levels (1, 3, 4, 11). Highest levels of susceptibility to BSE in mice expressing human PrP with codon 129-methionine (~30%) were reported by Asante et al (1) in the 2x over-expression Tg35 model (Hu-129M), which included identification of both limited clinical disease and subclinical disease. However lower attack rates of approximately 20% have been reported in Tg650 mice, which have a higher expression level of 129-Met human PrP, of around 5-8 fold (3, 26). The gene targeted transgenic mice utilised in our studies, which express wild type levels of human PrP from the endogenous mouse \textit{Prnp} locus, previously showed no incidence of disease following cattle BSE inoculation (4). These observations have lead to the assumption that over-expression of PrP is essential to model human disease susceptibility in mice, and that rodent models with wild type physiological levels of PrP expression do not live long enough to display signs of disease as would be seen in the longer lifespan human species. It is therefore significant that the inoculation of experimental sheep BSE described here has resulted in the identification of TSE related pathology in the gene targeted human PrP transgenic mice. Additionally, previously published data has shown that short incubation times can be achieved in HuMM and HuVV gene targeted mice (5). Our data show clearly that gene targeted transgenic lines
are useful in the study of cross species susceptibility, and that such susceptibility depends on the agent/host combination, rather than the lifespan of a mouse. The inclusion of data obtained in both overexpressing and gene targeted transgenic mice may therefore inform more accurately on the assessment of the true zoonotic potential of a particular TSE isolate.

The reasons for the increased susceptibility of HuMM mice to experimental sheep BSE in respect to cattle BSE are currently unknown, and are the subject of further investigation in our laboratory. One possible explanation is that our BSE infected sheep brain contained a significantly higher titre of BSE than found in the cattle BSE brainstem pool, resulting in the shortened incubation times in control 129/Ola and Bov6 mice with experimental sheep BSE inoculum-2 compared to cattle BSE, and the pathological features observed in HuMM mice. Incubation times in control mice were also shorter with experimental sheep BSE inoculum-2 compared to inoculum-1, although the ratio between 129/Ola and Bov6 mice was similar for each inoculum. Such variation in incubation time on primary passage of experimental sheep BSE is however common, and has been observed in previous experiments (Supplemental Information: Table S1). The observed difference in incubation times between inoculum-1 and inoculum-2 is therefore not unexpected. Previous studies by Gonzalez et al (22) have shown relatively high infectivity titres of sheep passaged BSE in RIII mice, which were equivalent to those obtained in Romney sheep. The infectivity titre of the cattle BSE brainstem pool used in our transmissions was $10^{3.3}$ ID$_{50}$ units/g in RIII mice (personal communication; R. Lockey & M. Simmons, Veterinary Laboratories Agency, UK). Those reported by
Gonzalez et al (22) for sheep passaged BSE were $10^5$ ID$_{50}$ units/g in RIII mice, suggesting higher titres may indeed be attained in sheep brain. However reported infectivity titres for BSE in cattle have been variable (6, 20, 22). We are therefore performing titration analyses of experimental sheep BSE brainstem in Bov6 mice to provide a direct comparison with titration data already available for the cattle BSE brainstem pool.

An alternative hypothesis is that passage of BSE through a sheep has altered the strain characteristics of the agent, producing a variant with increased virulence and/or host range. This possibility is supported by recent data describing enhanced virulence of experimental sheep BSE in bovine PrP transgenic mice (BoPrP-Tg110) and porcine PrP transgenic mice (PoPrP-Tg001) compared to cattle BSE (15, 16). BoPrP-Tg110 mice and PoPrP-Tg001 mice (which over-express PrP 8x and 4x respectively) produced significantly shorter incubation times following inoculation with an experimental sheep BSE brainstem pool than with cattle BSE isolates. The differences in incubation time observed in BoPrP-Tg110 mice were maintained on subpassage (15), indicating that the original variation was probably not due to infectivity titre discrepancies between the two BSE sources. However full titration of these tissue homogenates in mice would be required to confirm this was indeed the case. In PoPrP-Tg001 mice, incubation times shortened significantly on subpassage, and were maintained on further subpassage, indicating adaptation to the “new” host (16). In the study described here, lesion profiles obtained in control 129/Ola mice and Bov6 transgenic mice were similar for both cattle and experimental sheep BSE. Although we were unable to resolve the size of the PrP-res
low molecular weight band in both the experimental sheep BSE brain homogenate and
the experimental sheep BSE infected HuMM mouse, both showed reduced staining with
MAb 12B2 which is characteristic of BSE infection (27). There were therefore no
differences in strain characteristics between experimental sheep BSE and cattle BSE,
with the exception of the transmissibility to HuMM mice (which could be due to
increased infectivity titre).

The altered agent properties of sheep BSE observed by Espinosa and colleagues (15, 16)
may suggest that passage through a sheep causes BSE to transmit in a manner more
similar to natural scrapie than cattle BSE. To investigate this we inoculated our
transgenic panel with two isolates of natural sheep scrapie. No disease pathology was
observed in any transgenic mice following inoculation with either isolate of natural
scrapie. Hence, in the experiments described here, the susceptibility of the HuMM mice
to experimental sheep BSE does not appear to be due to a general susceptibility to ovine
prions, but is instead linked specifically to the replication of the BSE strain of agent in
sheep brain. Although agent strain characteristics of BSE are not altered when assayed in
Bov6 or 129/Ola mice following passage in sheep, both samples of experimental sheep
BSE did show positive TSE pathology in HuMM transgenic mice, which has not been
seen previously with cattle BSE inoculations in these mice. Whether this is simply due to
agent infectivity titre or a more subtle change in agent characteristics is the subject of
further analysis in our laboratory.
Although sheep can be experimentally infected with BSE by oral, intravenous or intracerebral exposure (18), no cases of sheep BSE have been reported in the field. The possible increased risk of disease transmission identified in these studies is thus not of major concern to the public at present. Natural BSE infection has however been identified in goats (14, 25), indicating that small ruminants have been exposed to sources of contamination. We cannot rule out the possibility that sheep may have been infected with BSE during the height of the BSE epidemic, as these animals were undoubtedly exposed to similar feed sources (although with different levels of exposure compared to cattle). Such infection may have been limited and/or localised, and resolved very quickly. BSE in small ruminants may however represent an increased risk to humans due the wider distribution of BSE infectivity identified in peripheral sheep tissues (2, 17, 19, 29) compared to BSE in cattle which is mainly restricted to the CNS (10). While TSEs remain in the environment and continue to infect animals (even at low prevalence) there remains the potential for cross-species transmission and the emergence of TSE isolates with altered strain properties, or host ranges. Our data therefore emphasise the need for continued surveillance to identify, monitor and characterise any new emerging TSE agents that are identified in ruminants, and assess the potential risks posed to other species.
Acknowledgements.

The authors would like to acknowledge I. McConnell, V. Thomson, S. Cumming, S. Shillinglaw, R. Greenan and K. Hogan for experimental setup, care and scoring of the animals; A. Coghill, A. Boyle, S. Mack and G. McGregor for histology processing and scoring; Dr M. Jeffrey (Veterinary Laboratories Agency, Lasswade, UK) for pathology advice; and the Veterinary Laboratories Agency, Weybridge, UK for providing the cattle BSE brain pool. This work was funded by contracts SE1439 and SE1441 from the UK Department for Environment, Food and Rural Affairs (Defra), and NIH-NIAID Agreement No.Y1-A1-4893-02 and FDA Agreement No. 224-05-1307. The findings and conclusions in this article have not been formally disseminated by the Food and Drug Administration and should not be construed to represent any Agency determination or policy.
1 References

2


    Alperovitch, S. Poser, M. Pocchiari, A. Hofman, and P. G. Smith. 1996. A
    New Variant Of Creutzfeldt-Jakob-Disease In the UK. Lancet 347:921-925.
31. Yull, H. M., D. L. Ritchie, J. P. Langeveld, F. G. van Zijderveld, M. E. Bruce,
    J. W. Ironside, and M. W. Head. 2006. Detection of type 1 prion protein in
Table 1. Transmission of cattle BSE, experimental sheep BSE and natural scrapie to gene targeted human and bovine transgenic mice

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<tr>
<th>TSE Isolate</th>
<th>Mouse Line</th>
<th>Incubation time*</th>
<th>Number affected†</th>
<th>Incubation time*</th>
<th>Number affected†</th>
<th>Incubation time*</th>
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<th>Number affected†</th>
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<th>Number affected†</th>
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<td>Cattle BSE (brain pool)</td>
<td>129/Ola</td>
<td>447 ±27</td>
<td>8/8</td>
<td>551 ±12</td>
<td>22/22</td>
<td>&gt;765</td>
<td>0/18</td>
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<td>Sheep BSE inoculum-1</td>
<td>129/Ola</td>
<td>474 ±22</td>
<td>11/11</td>
<td>564 ±8</td>
<td>17/17</td>
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* Days ± SEM, calculated from mice showing both clinical and pathological signs of TSE. >n represents the survival in days of the oldest mouse in groups where both clinical and pathological signs of disease were not observed in any animals

† number of mice showing TSE pathology (vacuolation and/or PrP deposition)/number of mice inoculated

‡ data from Bishop et al 2006 (4)
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<th>Clinical signs</th>
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<td>+++</td>
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<td>23</td>
<td>750</td>
<td>-ve</td>
<td>+</td>
<td>+</td>
<td>+++</td>
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</table>

*Animals died or culled for welfare reasons.

† PrP deposition scored as; -, no deposition; +, very low deposition; ++, low deposition; ++++, moderate deposition.

‡ an OD reading above the positive threshold value for the IDEXX assay is represented as +.

n/a, no spleen tissue available.
Figure Legends.

**Figure 1.** Pattern of vacuolation observed in brains of 129/Ola wild type mice (a), and Bov6 mice (b) following inoculation with the cattle BSE brainstem pool, and experimental sheep BSE inoculum-1 and inoculum-2. Profile produced from 9 grey matter areas (1, dorsal medulla; 2, cerebellar cortex; 3, superior colliculus; 4, hypothalamus; 5, thalamus; 6, hippocampus; 7, septum; 8, cerebral cortex; 9, forebrain cerebral cortex) and 3 white matter areas (1*, cerebellar white matter; 2*, midbrain white matter; 3*, cerebral peduncle). Average scores taken from a minimum of six mice per group and plotted against brain area ±SEM.

**Figure 2.** Limited PrP<sup>Sc</sup> accumulation in the thalamus of one HuMM mouse, 706 days post inoculation with experimental sheep BSE (inoculum-1). Panel (b) is higher magnification of the boxed area in panel (a). PrP<sup>Sc</sup> deposition appears to be restricted to the thalamus (b). Images obtained after staining with anti-PrP antibody 6H4 and counterstained with haemotoxylin. Magnification as shown.

**Figure 3.** Variation in pattern and location of PrP<sup>Sc</sup> accumulation in the brain of HuMM mice infected with experimental sheep BSE (inoculum-2). Punctate deposition in the cochlear nucleus (a) similar to BSE targeting in wild type mice. Peri-neuronal (b) and intra-neuronal (c) PrP deposition seen in the midbrain and areas of the thalamus. H&E stain of a large mature florid plaque located in the hippocampus (d). Punctate (e) and linear (f) PrP deposition in the thalamic region. Images a-c and e-f obtained after staining with anti-PrP antibody 6H4 and counterstained with haemotoxylin. Magnification as shown.
Figure 4. Comparative analysis of serial sections through the lateral geniculate nucleus (thalamus) of an uninoculated aged (750 days) HuMM mouse and a HuMM mouse infected with experimental sheep BSE (inoculum-2). HuMM mouse infected with sheep BSE (inoculum-2) shows astro- and microgliosis (b & f) visible when stained with anti-GFAP and anti-Iba1 (respectively). Several amyloid plaques are clearly visible fluorescing green with thioflavin-S (h) and stained with anti-PrP antibody 6H4 (d). Sections from a control aged HuMM mouse show mild astro and microgliosis, absence of PrP deposits or amyloid plaques (a, c, e, g). Sections used for immunohistochemical analysis were counterstained with haemotoxylin. x20 Magnification.

Figure 5. Comparative western blot (WB) analysis of the proteinase-K resistant fragment (PrPSc) of the prion protein. Discrimination between BSE and natural scrapie is achieved using two monoclonal antibodies 6H4 (a) and 12B2 (b). Lane 1 – 4.2mg equivalent brain material (mgE) of natural scrapie isolate from the NPU flock. Lane 2 – 20mgE of inocula NPU J2501, Cheviot sheep experimentally infected via the oral route with cattle BSE. Lanes 3 & 5 – 1.2 & 1.5mgE of 129/Ola mice infected with a natural scrapie isolate. Lane 4 – 64mgE of HuMM transgenic mouse infected with experimental sheep BSE (inoculum-2). Lanes 6-7 – 2.8 & 0.88mgE of 129/Ola and Bov6 (respectively) infected with experimental sheep BSE (inoculum-2). Lane 8 – 0.6mgE ME7/SV control. Molecular markers (M) of the standards are indicated on either side of the panels (kDa).
Uninoculated Aged HuMM Mouse | Sheep BSE Infected HuMM Mouse

(a) anti-GFAP

(c) 6H4

(e) anti-lba1

(g) Thioflavin-S

(b)

(d)

(f)

(h)
Figure S1. PrPSc deposition in the thalamus of a HuMM mouse inoculated with experimental sheep BSE inoculum-2 (a). Plaques and punctate staining seen in the thalamus and dLGN regions. Highlighted thalamic region (b) with plaques as well as punctate deposition. Stained with anti-PrP antibody 6H4 and counterstained with haemotoxylin. Magnification as shown.
Figure S2. Whole brain section of a HuMM mouse inoculated with experimental sheep BSE inoculum-2, illustrating bilateral PrP\textsuperscript{Sc} immunopositive fine punctate, coarse and plaque-like deposits in the thalamus and hippocampus. Stained with anti-PrP antibody 6H4 and counterstained with haemotoxylin. x4 magnification.
Table S1. Transmission of experimental sheep BSE to RIII wild type mice.

<table>
<thead>
<tr>
<th>Sheep</th>
<th>Tissue</th>
<th>Clinical +ve</th>
<th>Incubation time range (days)</th>
<th>Incubation time (days ±SE)</th>
<th>Clinical -ve</th>
<th>Survivors, path negative &gt;660dpi (number)</th>
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<tbody>
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<td>551± 19</td>
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<tr>
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<td>527± 18</td>
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<td>10</td>
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</table>

Brain homogenates prepared from four individual sheep infected with BSE were inoculated separately into groups of RIII mice. Animals were monitored for signs of TSE disease, and culled at a defined clinical endpoint or due to intercurrent illness. The data show the variability that can be observed on primary inoculation of mice with experimental sheep BSE. Incidence of disease ranged from 78% to 100%, with incubation times ranging from 350 days to 607 days. These observations are not uncommon in primary transmission studies.

N. Hunter and M. Bruce. Data from Defra funded project SE1428.