A Rapid Bioassay for Classical and L-Type Bovine Spongiform Encephalopathies

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ABSTRACT

The rapid detection of infectivity of several agents that cause Creutzfeldt-Jakob disease has previously been achieved by assaying for deposits of abnormal prion protein (PrPSc) in follicular dendritic cells in the spleens of transgenic mice carrying the human prion protein gene. In this study, transgenic mice expressing the bovine prion protein were inoculated intraperitoneally with classical (C-type) or atypical L-type bovine spongiform encephalopathies (BSE). Proteinase-resistant PrPSc were detected in the spleens of all transgenic mice at 75 days after inoculation with both types of BSE. Infectivity in PrPSc-positive spleens of the transgenic mice revealed that prions of C- and L-type BSE replicated. These results suggest that bioassay system by the transgenic mice could be useful for the rapid detection of BSE infectivity with discriminating between C- and L-type BSEs.

Keywords: Bovine Spongiform Encephalopathy; Follicular Dendritic Cell; Transgenic Mice

1. Introduction

Transmissible spongiform encephalopathies (TSEs) are fatal, infectious neurodegenerative diseases; TSEs include bovine spongiform encephalopathy (BSE) in cattle, scrapie in sheep and goats, chronic wasting disease mainly in deer and elk, and Creutzfeldt-Jakob disease (CJD) in humans. BSE was first defined in 1986 in the United Kingdom [1]. There was little variation in the distribution of histopathological changes and the deposition of disease-associated prion protein (PrPSc), which is converted from the host-encoded normal form (PrPc), and also in the glycoform pattern and molecular mass of proteinase K (PK) resistant fragments of PrPSc, named as PrPcore, among BSE cases [2-5]. In addition, transmissibility of BSE agents to inbred mice has revealed defined incubation period from inoculation to development of clinical signs and distinct histopathological characteristics in the brain, as compared to the other TSE agents [6,7].

It was thought that all BSE cases shared the same origin and a single set of characteristics. However, atypical BSEs possessing different prion characteristics have been reported since 2003 in Japan, France, Italy, the United States, Germany, Poland, the Netherlands, Sweden, and the United Kingdom [8-17]. As determined by Western blotting, atypical BSEs are classified into the two types, which show higher (H-type) or lower (L-type) molecular mass of PrPcore on the basis of that in classical (C-type) BSE [8,9,12]. In 2006, a Japanese black cow was diagnosed as an atypical BSE [11]. Transmission of this agent to cattle or transgenic (Tg) mice over-expressing the bovine PrP gene realized that characteristics of histopathological changes and PrPcore were similar to L-type BSE agents found in Italy, Germany, France and Canada [18-21]. Incubation period of the Tg mice affected with L-type BSE was approximately 180 days post intracerebral (IC) inoculation and was shorter period than that with C-type BSE which was approximately 230 days [20, 22-24].

Transmission of human CJD agents to humanized knock-in (Ki) mice demonstrated that PrPSc could be detected in the follicular dendritic cells (FDCs) in the lymphoreticular tissues within 14 days post intraperitoneal (IP) inoculation [25]. Accordingly, the assessment of PrPSc accumulation in splenic FDCs can be used as a rapid method (FDC assay) to ascertain the infectivity of human CJD at 75 days post IP inoculation [25].

We report here that the FDC assay can be used to
demonstrate C- and L-type BSE infection in a Ki+Tg#40 mouse that we generated by crossing Ki mouse with Tg mouse. Transmission of BSE into the Ki+Tg mouse was shown rapidly by detection of PrP\textsuperscript{Sc} in splenic FDC after IP inoculation compared to the period when the mice develop clinical signs after IC.

2. Materials and Methods

2.1. Transgenic Mouse with a Bovine PrP Gene

We generated Ki+Tg#40 mice by crossing a Ki mouse with a Tg mouse, which carries the same gene construct with the open reading frame of the bovine PrP gene with 6 octarepeats [26]. The Ki+Tg#40 mice carry 12 copies of the bovine PrP gene on the hemizygote of Tg insertion and express about 6-fold more PrP\textsuperscript{C} in the brain than found in Ki mouse brains (Figure S1). When inoculated via IC route with a BSE agent, the Ki+Tg#40 mice showed shorter incubation period than Ki mice and PrP\textsuperscript{Sc} deposition in the brain and spleen, differently from the Tg#40 mice backcrossed into PrP-knockout background that showed in the brain but not in the spleen (Table S1).

![Figure S1. Western blotting of normal brains of Ki+Tg#40 mice and Ki mice. All samples without PK treatment were subjected to Western blotting with 6H4. Five samples of Ki+Tg#40 mice were loaded at 0.1 mg of the brain equivalent to wet weight. For estimating PrP\textsuperscript{C} expression level in the brain of Ki+Tg#40 mice, a sample of Ki mice was loaded at 1-fold, 0.1 mg; 4-fold, 0.4 mg; 6-fold, 0.6 mg; 8-fold, 0.8 mg of the brain equivalent to wet weight. The right bars indicate molecular mass markers, 38 kDa, 28 kDa and 19 kDa.](image)

Table S1. Transmissibility of BSE/UK1 cattle brain to three lines of transgenic mice. Ki mice were generated by genetargeting procedure with the bovine PrP gene. Tg#40 mice were generated by random insertion procedure and by backcrossed into Prnp knockout background. We generated Ki+Tg#40 mice by backcrossing Tg#40 into Ki mouse background.

<table>
<thead>
<tr>
<th>mouse line</th>
<th>incubation period (mean days ± SD)</th>
<th>PrP\textsuperscript{Sc} by immunohistochemical staining</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ki</td>
<td>607 ± 32</td>
<td>5/5</td>
</tr>
<tr>
<td>Tg#40</td>
<td>366 ± 5</td>
<td>3/3</td>
</tr>
<tr>
<td>Ki+Tg#40</td>
<td>350 ± 29</td>
<td>4/4</td>
</tr>
</tbody>
</table>

\[^1\text{PrP}\textsuperscript{Sc} positive number/examined number of mouse; }^1\text{The same data were shown in Table 1.}

2.2. BSE Inoculum

Two cattle brains infected with C-type BSE were obtained from the TSE Archive, Veterinary Laboratories Agency, Weybridge, the United Kingdom (reference number: PG2458/98; designated as BSE/UK1) and from the National Institute of Infectious Disease, Tokyo, Japan (the sixth Japanese case of BSE; designated as BSE/JP6). A cattle brain infected with L-type BSE (the 24th Japanese case of BSE; designated as BSE/JP24) was also obtained from the National Institute of Infectious Disease. These 3 cattle brains were found to be positive for BSE PrP\textsuperscript{Sc} by Western blotting. As a negative control, a normal brain sample from a healthy cow was obtained from a slaughterhouse and was found to be negative for BSE PrP\textsuperscript{Sc} by Western blotting. To prepare the inoculum for transmission and Western blotting, each cattle brain sample was homogenized at 10% (w/v) with phosphate-buffered saline (PBS) using Multi-beads shocker (Yasui Kikai Corp., Osaka, Japan), and debris was removed by centrifuging at 900 \times g for 10 min.

2.3. Transmission Studies

This animal experiment was approved by the Animal Experiment Committee at the National Institute of Animal Health. For the FDC assay, Ki+Tg#40 mice were inoculated via IP route with 50 \textmu l of 10% cattle brain homogenate and were euthanized 75 days after inoculation. The spleens of mice were harvested and divided in two pieces; one was fixed with 4% buffered formalin for immunohistochemical staining and the other was frozen at -80\degree C for Western blotting and subsequent transmission. Mice were defined as positive for BSE transmission if PrP\textsuperscript{Sc} deposition in splenic germinal center was observed by immunohistochemical staining and/or if the presence of PrPcore in the spleen was detected by Western blotting.

To confirm the BSE infectivity in cattle brain or mouse spleen, Ki+Tg#40 mice were inoculated via IC route with 20 ul of 10% homogenate from cattle brain or PrP\textsuperscript{Sc}-positive spleens of three Ki+Tg#40 mice at 75 days after IP inoculation. Incubation periods were defined as the day from inoculation to euthanasia with clinical signs of mouse BSE, including tremors, a high-arched back, incoercible food intake, and significant loss of body weight. BSE transmission to mice was confirmed by histopathological analysis with observation of vacuolation and PrP\textsuperscript{Sc} deposition in the brain.

2.4. Immunohistochemical Staining

Hydrolitic autoclaving pretreatment was used to enhance antigen retrieval. An anti-PrP rabbit serum previously raised against the N-terminal of PrP was used as the primary antibody [27]. Universal immunoperoxidase poly-
(Histofine MAX PO kit; Nichirei Biosciences Inc., Tokyo, Japan) was used as the secondary antibody for the detection of PrPSc accumulation.

### 2.5. Western Blotting

The PrPcore was extracted from animal tissues by collagenase and PK treatments, with some modifications to a method described previously [16]. Samples were concentrated after the extractions using the phosphotungstic acid method [28]. PNGase F (New England Biolabs, Beverly, MA) was used for the deglycosylation of PrPcore. Prior to electrophoresis, the samples were resolved in NuPAGE® LDS Sample Buffer (Invitrogen Corp., Carlsbad, CA) containing 1 mM dithiothreitol after treatment at 100°C for 10 min on a heat block. The samples were subsequently separated by a NuPAGE® 12% Bis-Tris gel (Invitrogen Corp.) in NuPAGE® MOPS SDS Running Buffer (Invitrogen Corp.) and were transferred electrically onto an Immobilon-P transfer membrane (Millipore, Bedford, MA). The antiPrP monoclonal antibody 6H4 (Prionics AG, Zurich, Switzerland) was used as the primary antibody. The PrP signal was generated using Envision (Dako Japan Inc., Tokyo, Japan) and SuperSignal® West Dura Extended Duration Substrate (Pierce Biotechnology Inc., Rockford, IL) and was imaged directly with the Fluochem™ imaging system (Alpha Innotech, San Leandro, CA).

The relative amounts of di-, mono-, and un-glycosylated forms of PrPcore were calculated as the mean percentages of the total signal of the 3 glycoforms from 5 repetitions of the Western blot analysis of the cattle and mouse brains. These calculations were performed using AlphaEaseFC™ software (Alpha Innotech).

### 3. Results

The spleens of all the Ki+Tg#40 mice at 75 days after IP inoculation with BSE cattle brains were positive for PrPSc deposition (Table 1). Immunohistochemical staining showed that PrPSc accumulations were finely stained in the germinal center of the spleen with no distinction between BSE/UK1 and BSE/JP24 (Figure 1(a)). Western blotting showed PrPcore in the spleen of Ki+Tg#40 mice at 75 days post IP inoculation with BSE cattle brains, but not in the brain (Figure 1(b)). None of the mice inoculated with a normal cattle brain showed any positive results for PrPSc (data not shown).

All Ki+Tg#40 mice inoculated via IC route developed clinical signs. The mean incubation periods were 350 days with BSE/UK1, 431 days with BSE/JP6, and 313 days with BSE/JP24 (Table 1). Sequential passage with the PrPSc-positive mouse spleens that were harvested at 75 days post IP inoculation showed that all Ki+Tg#40 mice developed clinical signs after IC inoculation (Table 1). The mean incubation periods were 365 days with BSE/UK1 mouse spleen and 338 days with BSE/JP24 mouse spleen.

Histopathological analysis using haematoxylin-eosin staining showed severe vacuolation in the hippocampus of the Ki+Tg#40 mice inoculated with the BSE/JP24 cattle brain or mouse spleen (Figures 2(a) and (b)). Moreover, mild vacuolation was observed in the brain cortex and thalamus of these mice. In contrast, less vacuolation

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**Table 1. Transmission of BSE to Ki+Tg#40 mice.**

<table>
<thead>
<tr>
<th>Inoculum</th>
<th>IP route</th>
<th>IC route</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>PrPSc-positive in spleen (n/nt)</td>
<td>Incubation period (mean days ± SD, n/nt)</td>
</tr>
<tr>
<td>BSE/UK1 cattle brain</td>
<td>4/4</td>
<td>350 ± 29, 4/4</td>
</tr>
<tr>
<td>BSE/JP6 cattle brain</td>
<td>4/4</td>
<td>431 ± 23, 7/7</td>
</tr>
<tr>
<td>BSE/JP24 cattle brain</td>
<td>5/5</td>
<td>313 ± 10, 6/6</td>
</tr>
<tr>
<td>BSE/UK1 mouse spleen</td>
<td>nt</td>
<td>365 ± 35, 6/6</td>
</tr>
<tr>
<td>BSE/JP24 mouse spleen</td>
<td>nt</td>
<td>338 ± 18, 5/5</td>
</tr>
</tbody>
</table>

1) PrPSc-positive mouse spleen at 75 days post IP inoculation with BSE cattle brains. 2) 75 days after inoculation. 3) n/nt: number of positive mice per number of inoculated mice, nt: not tested.
was observed in the hippocampus of mice inoculated with the BSE/UK1 cattle brain or mouse spleen (Figures 2(c) and (d)). Immunohistochemical staining revealed a fine granular deposition of PrPSc in the thalamus of mice inoculated with the BSE/JP24 cattle brain or mouse spleen (Figures 2(e) and (f)), while a coarse granular deposition of PrPSc was found in the thalamus of mice inoculated with BSE/UK1 cattle brain or mouse spleen (Figures 2(g) and (h)). A similar pattern of coarse granular deposition was observed in the mice inoculated with BSE/JP6 cattle brain (data not shown).

Western blot analysis showed that the PrPcore characteristics of BSE/JP24 in the cattle brain, the spleen and brain of the transmitted Ki+Tg#40 mice were different from those of BSE/UK1 and BSE/JP6 in the corresponding tissues (Figure 3(a)). The relative amounts of the di-, mono-, and un-glycosylated forms of PrPcore in the cattle brain were present in the approximate ratio of 37:40:23 in BSE/JP24, 71:21:7 in BSE/UK1, and 81:17:2 in BSE/JP6 (Figure 3(b)). Similarly, the different forms of PrPcore in the mouse brains were present in an approximate ratio of 43:35:22 in BSE/JP24, 77:17:7 in BSE/UK1, and 82:14:3 in BSE/JP6. In the mouse spleens, the different forms of PrPcore were present in an approximate ratio of 54:34:12 in BSE/JP24, 77:18:5 in BSE/UK1, and 79:18:3 in BSE/JP6. The relative amount of the di-glycosylated form in the brains and the spleens of mice transmitted
with BSE/JP24 was slightly higher than that in the original cattle brain, but was not predominant (Figure 3(b)).

Western blot analysis also revealed that the unglycosylated molecular masses of PrPcore in the BSE/JP24 cattle brain, mouse spleen and mouse brain were lower than those in BSE/UK1 and BSE/JP6, which was confirmed by deglycosylation with PNGase F (Figure 3(c)).

4. Discussion

The FDC assay in the Ki+Tg#40 mice realized that PrPSc was generated in the mouse spleens within 75 days post IP inoculation with BSE cattle brains (Figure 1(a)). The PrPSc-positive mouse spleen had a well-established infectivity to Ki+Tg#40 mice (Table 1). In the brain of the Ki+Tg#40 mice, no PrPSc was not detected at 75 days post IP inoculation with C-type or L-type BSE (Figure 1(b)) although the PrP Sc in the brain was detected when the Ki+Tg#40 mice developed overt clinical signs post IC (Figures 2 and 3) and IP inoculation (data not shown). It is reported in other TSEs following peripheral routes of infection that PrPSc accumulation occurs in FDC of lymphoid tissues prior to in the neural tissues [29,30]. These results support that Ki+Tg#40 mice showing PrPSc accumulation in FDC should develop BSE at preclinical stage after IP inoculation.

Immunohistochemical staining of the mouse spleen at 75 days post IP inoculation showed appearance of PrPSc accumulation in L-type BSE was indistinguishable from that in C-type (Figure 1(a)). Subsequent passage of the PrPSc-positive mouse spleen, however, realized in the affected brain that histopathological properties of L-type and C-type BSEs were distinct from each other and similar to those after transmitted with the original cattle brain (Figure 2). These results suggest that prions of L-type and C-type BSEs should replicate in the mouse spleen at 75 days post IP inoculation.

It was easy to discriminate L-type BSE from C-type by Western blotting of the mouse spleen at 75 days post IP inoculation. The molecular mass of deglycosylated PrPcore in L-type BSE was apparently lower than that from C-type BSE (Figure 3(c)). The relative amount of the diglycosylated PrPcore in L-type BSE was less than that in C-type BSE, but slightly increased comparing to that in the original cattle brain (Figure 3). It has reported in other TSEs that the glycoform pattern and molecular mass of PrPcore was controlled not only by the prion-specific conformation but also by the PrPSc distribution in the tissue and brain regions [31]. The slight increasing of di-glycosylated PrPcore in the L-type BSE was shown in the mouse brain as like as in the spleen (Figure 3), suggesting that glycoform pattern of PrPcore could be controlled slightly by unknown factors in mouse during the replication of prion.

In conclusion, the FDC assay in Ki+Tg#40 mouse enables to verify the infectivity of C- and L-type BSEs rapidly and could also discriminate L-type from C-type BSE. For establishing the FDC assay as diagnosis tool of BSE, we have preceded the further studies of sensitivity and specification. This FDC assay system may archive a more rapid risk-evaluation of BSE infection to humans and cattle.

5. Acknowledgements

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