RESEARCH ARTICLE

Generation of a Persistently Infected MDBK Cell Line with Natural Bovine Spongiform Encephalopathy (BSE)

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Abstract

Bovine spongiform encephalopathy (BSE) is a zoonotic transmissible spongiform encephalopathy (TSE) thought to be caused by the same prion strain as variant Creutzfeldt-Jakob disease (vCJD). Unlike scrapie and chronic wasting disease there is no cell culture model allowing the replication of proteinase K resistant BSE (PrPBSE) and the further in vitro study of this disease. We have generated a cell line based on the Madin-Darby Bovine Kidney (MDBK) cell line over-expressing the bovine prion protein. After exposure to naturally BSE-infected bovine brain homogenate this cell line has shown to replicate and accumulate PrPBSE and maintain infection up to passage 83 after initial challenge. Collectively, we demonstrate, for the first time, that the BSE agent can infect cell lines over-expressing the bovine prion protein similar to other prion diseases. These BSE infected cells will provide a useful tool to facilitate the study of potential therapeutic agents and the diagnosis of BSE.

Introduction

Transmissible spongiform encephalopathies (TSEs) are progressive neurodegenerative disorders causing degeneration of neurons and include Creutzfeldt-Jakob disease (CJD), bovine spongiform encephalopathy (BSE), scrapie, transmissible mink encephalopathy (TME), and chronic wasting disease (CWD). In 1996 a new disease, variant CJD (vCJD) was identified with evidence suggesting that vCJD and BSE are caused by the same prion strain. vCJD is most likely caused from consumption of contaminated beef or beef by-products [1]. The causative agent of TSEs most likely is an infectious protein (PrPSc) which unlike bacteria and viruses, does not contain any nucleic acid to propagate itself. PrPSc is generated from a normal host-encoded cellular prion protein (PrPC) during disease and is conformational different to the normal cellular protein [2]. These conformational differences result in an increased resistance to degradation allowing for detection of the disease associated PrPSc.
The development of cell lines for a specific prion disease may be advantageous for a variety of studies, for example, screening of anti-prion substances, formation and inhibition of pathogenic prions \([3–6]\). However, the availability of cells susceptible for TSE infection is still very limited. The majority of susceptible cells are mouse-derived \([7,8]\). In addition, propagation of chronic wasting disease (CWD) has been successfully achieved in a mule deer-derived fibroblast-like cell line \([9]\) and rabbit RK13 cells expressing elk PrP and the HIV-1 GAG precursor protein (RKE-Gag) \([10]\). However, no cells susceptible to infection with natural BSE from cattle have been established. To date, BSE related research relies heavily on the use of mice or transgenic mice expressing animal species-specific PrP\(^C\) \([11,12]\), or on large animal studies \([13]\). There is a strong requirement for replacing the animal models with in vitro systems using cell lines susceptible to BSE infection, to reduce the time and cost of such studies. Such systems will significantly facilitate the diagnosis of BSE as well as the study of potential therapeutic agents and disease pathogenesis.

In this study, we report for the first time a cell line which is persistently infected with BSE utilizing Madin-Darby Bovine Kidney (MDBK) cells over-expressing bovine PrP established using a lentiviral expression system. These results provide evidence that PrP\(^{BSE}\) is able to replicate persistently in an in vitro cell culture.

**Materials and Methods**

**Prion protein genes (PRNPs) and cloning**

Primer sequences were designed against the bovine PRNP gene (GenBank: AJ298878) and, for cloning, NotI and XhoI restriction enzyme sites were added to a forward primer (5'-AGCCGCGGCACCATGTGAAAAGCCACATAGG-3') and a reverse primer (5'-CGGCTCGAGCTATCCTACTATGAGAAAATGA-3'), respectively. Using the above primer pair, the complete PRNP coding region from brain tissues of cattle (Korean native cattle; Hanwoo) which were collected from abattoir (Hyup Sin Food Co. Ltd., Korea) was amplified by PCR as follows: initial denaturing at 94°C for 5 min, followed by 30 cycles of 94°C for 30 seconds, 60°C for 30 seconds and 72°C for 90 seconds, and a final elongation stage at 72°C for 15 min. The amplicons were cloned into pGEM-T Easy vector (Promega) and DNA sequencing of the target genes were performed (Macrogen Inc., Korea).

**Infectious recombinant lentiviruses**

Infectious recombinant lentiviruses were produced following published methods \([14,15]\) by Macrogen Inc.(Seoul, Korea). Briefly, the cloned PRNPs were inserted into pLEX MCS transfer vector (Thermo Scientific Open Biosystems) using restriction enzymes NotI and XhoI (New England Biolabs). This cloned transfer vector was then mixed with a VSV-G expression vector and a gag-pol expression vector in a relative molar ratio of 1:1:1, and co-transfected into 293T cells using lipofectamine Plus (Invitrogen, USA). The cell culture supernatant containing recombinant virus was recovered 48 hours after transfection, and filtered using a membrane filter with a pore size of 0.45 μm (Nalgene, USA), and stored immediately at −70°C. A titer value of the infectious recombinant virus was indirectly measured in HeLa cell, using fluorescent microscopy, to detect GFP expression in the transduced cell which has transfected with only pLEX vector containing GFP gene instead of bovine PRNP.

**Cell and transduced cell lines**

MDBK cell was obtained from the American Type Cell Collection (ATCC). Cells were grown in completed medium (Dulbecco’s modified Eagle’s medium/F12 supplemented with 10% fetal...
bovine serum, antibiotics (penicillin and streptomycin), non-essential amino acid, and L-glutamine). To determine puromycin concentration for selection of transduced cells, cell lines were treated with 0 to 10 μg/ml of puromycin and cultured for 3 ~ 4 days and observed for cell death. The optimal concentration was determined to be in the range of 1.5 to 2.5 μg/mL and used for selecting the transduced cell.

The transduced cell lines were produced as follows: Day 1 before recombinant lentiviral infection, cell lines were plated on a 12-well plate for culture and allowed to grow to 60 to 70% confluence on the inoculation day. After removing the culture supernatant, 0.5 mL of infectious recombinant lentivirus was inoculated and 8 μg/mL of Polybrene (Hexodimethrine bromide, Sigma H9268) was added, followed by gently shaking the mixture. After incubating overnight (for 15 to 16 hours) at 37°C under 5% CO₂ atmosphere, the inoculum was replaced by fresh completed medium and, on the following day, puromycin was added to the culture medium and transduced cells were selected while continuously culturing for 7 to 14 days.

**Immunofluorescence assay**

The transduced cells or the normal cells were cultured in chamber slide (Nunc, Lab_TeK chamber slide) and fixed with 4% paraformaldehyde (PFA, Sigma P6148) for 30 minutes and dried in order to determine whether or not prion protein was expressed on the cell membranes. A primary antibody, 6H4 (Prionics, 1 mg/mL, 1:100) was added to the fixed cells, and incubated for 1 hour at room temperature. After washing the cells 3 times using phosphate buffered saline (PBS), anti-mouse IgG FITC conjugate (KPL; Kirkegaard & Perry Laboratories, 0.5 mg/mL, 1:200) was added and incubated for 1 hour at room temperature. After washing the cells 6 times using PBS they were mounted and observed on a fluorescent microscope (Olympus, Tokyo, Japan).

**Immunoblotting**

Cultured cells were washed once using PBS and adherent cells collected from the culture flask in PBS using a scraper and transferred to a centrifuge tube. After centrifugation at 1,200 rpm for 5 minutes to pellet the cells, the supernatant was discarded and the pellet was dispersed by gently tapping the tube. The cells were lysed by adding lysis buffer (0.5% Triton X-100, 0.5% sodium deoxycholate, 10 mM Tris-HCl [pH7.5], 150 mM NaCl, 5 mM EDTA) to the suspension, and incubated on ice for 30 minutes. After this incubation, the product was subjected to refrigerated centrifugation at 1,200 rpm for 5 minutes and the supernatant was transferred into a new tube.

For PrP<sup>C</sup> analysis, using Dynabeads (Dynabeads M-280 Sheep anti-mouse IgG) coupled to 6H4 (1.5 μg of 6H4 per 30 μL beads suspension), the supernatant was subjected to a semi-purification according to the manufacture instructions. For PrP<sup>Sc</sup> analysis, the supernatant containing ~ 0.5 ~ 1 mg of total protein was incubated with 20 μg/mL proteinase K for 20 min at 37°C prior to addition of 1 mM 4-(2-Aminoethyl)-benzenesulfonyl fluoreide (Roche) and incubation for 5 minutes in ice. The proteinase K treated proteins were centrifuged at 20,000 x g for 2 hours and the pelleted protein was resuspended in 25 μL 1 x sample loading buffer (Invitrogen). For removing N-linked glycosylation from PrP<sup>Sc</sup> in cells before PK digestion, PNGase F was used according to the manufacturer’s recommendation (New England Biolabs). US/FluorchemQ analysis (Alpha Innotech, USA) was used for estimating migration of unglycosylated PrP after treatment of PNGase F.

The electrophoresis was conducted at 200 V for 35 minutes using a pre-casted NuPAGE 12% Bis-Tris gel (Invitrogen) and proteins were transferred to a polyvinylidene difluoride (PVDF) membrane by an electrical device at 150 V for 1 hour. After transfer the PVDF
membrane was blocked using a blocking solution comprising 0.02% I-block (Tropix) dissolved in a tris-buffered saline (TBS), the membrane was incubated with the 6H4 monoclonal antibody (1 mg/mL, 1:3,000) or rabbit anti-PrP polyAb S1 (produced by the Animal and Plant Quarantine Agency, Korea) at room temperature for 1 hour. The membrane was washed 3 times using TBST (which comprises 0.05% (v/v) Tween 20 in TBS) and reacted with goat anti-mouse IgG (or anti-rabbit IgG) alkaline phosphatase-conjugated secondary antibody (KPL, 1 mg/mL, 1:3,000) at room temperature for 1 hour. Finally, the membrane was washed 3 times using TBST for 5 minutes and expression of prion protein was visualized using CDP star (Applied Biosystem). Chemiluminescence signals were analyzed using the LAS 4000 imaging system (Fuji, Japan).

ELISA and scrapie cell assay
To determine whether PrPSc infection was established in the transduced cell, a bovine brain homogenate from classical BSE infected cattle (supplied by TSE Archive, Animal Health and Veterinary Laboratories Agency UK) was prepared and used to challenge these cells according to the following procedure. First, the BSE brain homogenate was diluted in DMEM supplemented with 100 ng/mL of phorbol 12-myristrate 13-acetate (PMA, Sigma P8139), 10% fetal bovine serum (Gibco-BRL 16000–044), antibiotics (Gibco-BRL 15140–155), non-essential amino acids (Gibco-BRL 11140–050) to a final concentration of 0.25% and 100 μL of the diluted homogenate was inoculated onto a 96-well tissue culture plate containing 60 to 70% confluent cells. Once confluent, cells were serially transferred to a 24-well tissue culture plate, a 25 cm² flask and a 75 cm² flask at 4 to 6 day intervals.

BSE infection of the cultured cells in the 75 cm² flask was analyzed by ELISA (IDEXX HerdCheck) according to the manufacturer’s instructions. The transduced cells were subjected to limiting dilution using a 96-well plate after confirmation of infection by ELISA. The limiting dilution was performed by plating out an average of 0.3 cells per well of a 96-well plate. The individual clones were screened using the published protocol of the standard scrapie cell assay (SCA) [16] modified and performed as below. The single cloned cells were propagated until confluent. About 25,000 cells per well were suspended in DMEM containing 10% fetal bovine serum and transferred into each well of 96-well enzyme linked immune-spot assay (ELISPOT) plates (MultiScreen-IP filter plate; Millipore). Plates were dried and proteinase K (PK) treated (1 μg/mL in lysis buffer), followed by protease inhibition with phenylmethylsulfonyl fluoride (PMSF; sigma) and denaturation with Tris-guanidinium thiocyanate (GSCN). The denaturated PrPSc proteins were detected with the primary antibody, 6H4 (1 mg/mL, 1:3,000) and secondary antibody, alkaline phosphatase-conjugated anti-mouse IgG (1 mg/mL, 1:3,000, KPL) and detected with alkaline phosphatase conjugate substrate, BCIP/NBT (KPL). The immunostained spots were visualized with CTL ELISPOT equipment (Cellular Technology, Ltd, Shaker Heights, OH)

Results
Production of infectious recombinant lentiviruses and transduced MDBK
In order to produce infectious recombinant lentivirus for transducing the MDBK cell line and expressing recombinant bovine prion protein, bovine prion protein gene (PRNP) was amplified by PCR. The amplified bovine PRNP was compared with accession numbers AJ298878 (100%), and EU224471 (99.87%) registered in GenBank, it contained 6 octarepeats. The infectious recombinant lentivirus was produced using the amplified bovine PRNP as described in materials and methods. The measured titer of the lentivirus was more than 10⁶ transduction unit (TU) as determined by fluorescent antibody (FA) test (data not shown).
The transduced MDBK cell line was produced using recombinant lentivirus containing bovine PRNP and puromycin resistance to select transduced cell clones. In Fig. 1, we compared to the non-transduced MDBK control (A), the transduced MDBK with only pLEX vector (B) and MDBK C1–2F with bovine PRNP inserted into pLEX vector (C) for the expression of prion protein on their cell surface. MDBK C1–2F cells showed that PrPC was highly expressed on the cell surface compared with the controls (MDBK cells express moderate levels of intrinsic PrPC). By Western blot (WB), the prion protein expression levels between the three types of cells was determined by comparison with the house-keeping protein GAPDH. The transduced MDBK cell, MDBK C1–2F cell, exhibits the highest expression level of prion protein (Fig. 1D, lane 3).

Production of PrP\textsuperscript{BSE} infected cell line

In order to determine whether PrP\textsuperscript{Sc} infection could occur in the bovine PrP transduced cells, a 0.25% (w/v) bovine brain homogenate from naturally BSE infected cattle was used to challenge the transduced cells. BSE infection was firstly identified by ELISA performed on the challenged cultured cells from a 75 cm\textsuperscript{2} flask (data not shown). The infected cells were subjected to limiting dilution using 96-well plates to isolate pure infected clones. Isolated clones were analysed using the scrapie cell assay [16]. Six wells (ID5, IIIG1, IVB2, IVF12, VD7 and VG3) out of 5 plates (480 wells plated) showed strong positive responses and two showed weak positive responses (IVC4 and IIID9), see Fig. 2. In addition, the optical densities (mean ± standard deviation, n = 7) of IVF12 and VD7 were found to be 3.118 ± 0.026 and 3.151 ± 0.007 respectively. These values when compared with the value of 0.05 ± 0.004 of MDBK C1–2F control cells confirmed the results of the SCA and indicated that the cells are infected with naturally occurring BSE.

Confirmation of persistent PrP\textsuperscript{BSE} infected cell (M2B cell)

To confirm whether these cells were persistently infected, one clone, VD7 (see Fig. 2) was passaged further and cloned by limiting dilution adjusting the number of cells to approximately...
0.3 cells per well in 96-well plate. As shown in Fig. 3, proteinase K resistant PrPSc in the BSE infected uncloned cells disappeared before undergoing 20 passages (lane 3 to 6 in Fig. 3). However, the cloned infected cells, M2B (stand by M DBK with 2-times cloning after challenge of BSE infected bovine brain homogenate) exhibited persistent infection, proteinase K resistant PrPSc even after 26 passages by continuous subculture (lane 7 to 9 in Fig. 3). In contrast the unchallenged control cells have no proteinase K resistant prion protein band (lane 2 of Fig. 3), since normal prion protein in these cells was completely degraded by proteinase K treatment. The infected cells maintained their infection as evidenced by a proteinase K resistant PrPSc signal and is consistently seen throughout subsequent passages up to passage 83 (Fig. 4 lanes 4–7). In comparison to the nonglycosylated PrPSc of the original BSE inoculum (Fig. 4, lane 2 and Fig. 5 lane 2 after PNGase F deglycosylation), those of the cell line derived PrPSc have a lower molecular weight (about 1.3 kDa) as detected by WB (Figs. 4 and 5).
Discussion

Until now, BSE susceptible cell lines have not been developed. Although Heining et al. [17] recently have shown that PrPres could be detected in the PrP0/0 Chimp1 cell clone after incubation with BSE infected cow brain homogenate, this cell line has not been established as a persistently infected cell line against natural BSE infected bovine materials. Various cell lines have been developed which are susceptible to infection mainly with murine-adapted or ovine scrapie-infected materials and also recently two cell lines have been produced allowing infection with chronic wasting disease (CWD)[10,18–21]. No reliable cell model exists for human prion diseases including variant Creutzfeldt-Jakob disease (vCJD) although the human neuroblastoma cell line SH-SY5Y was shown to be permissive to human prion infection [22]. In this study, we have produced a persistently BSE-infected cell line, M2B, which overexpresses bovine prion protein in MDBK cells using a lentiviral expression system [14,15], and infected these cells with bovine brain homogenate from cattle infected with classical BSE. Prior to establishing this cell line we attempted to use primary bovine brain cells, especially neuronal cells however these cells were not easily immortalized using simian virus 40 large T antigen [23] or successfully passaged after infection (data not shown). Non-neuronal cells have been reported to be permissive to infection with chronic wasting disease (CWD)[10,18–21]. Vorberg et al. [25] demonstrated that the mouse fibroblast cell lines NIH/3T3 and L929 are permissive to infection with mouse-adopted scrapie in spite of expressing low level of cellular mouse prion protein. Here we have used MDBK to generate a BSE susceptible cell line, although PrPSc expression level of kidney is lower than any other tissue [26] and, to our knowledge there are no reports of PrPSc detection from bovine kidney. Ryder et al. and Hadlow et al. [27,28] found that PrPSc has been detected in the kidney of cats with feline spongiform encephalopathy (FSE) and experimental mink with transmissible mink encephalopathy (TME). Additionally cell lines based on the rabbit kidney cell line RK13 have been shown to be susceptible to a range of prion disease [29,30]. On the basis of above reports, we over-expressed bovine prion protein in MDBK using a lentiviral expression system to increase expression level of prion protein to potentially

Figure 4. Immunoblotting detection of the persistent PrP^{BSE}-infected cell according to serial passages. The lysates of brain homogenates (lane 1; BSE-noninfected bovine brain, lane 2; BSE-infected bovine brain-original BSE source) and MDBK cells expressing normal bovine prion protein by using infectious recombinant lentivirus (lane 3: transduced MDBK–MDBK C1–2F–passage(p) 7; lane 4: BSE-infected transduced MDBK (M2B)–p17; lane 5: M2B-p48; lane 6: M2B-p70 and lane 7: M2B-p83) were treated with proteinase K (PK) for detection of PrP^{BSE} in transduced MDBK which was sequentially passaged after inoculating BSE-infected bovine brain homogenate. Molecular mass marker (M) in kilodaltons (kDa) is shown on the left. The result shown is representative of multiple independent experiments.

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enhance susceptibility to prion infection [14,15]. As shown in Fig. 1, the recombinant prion protein is well expressed on the cell surface of bovine PRNP transfected cell compared to control cells but all cells express prion protein in cell lysates as shown by immunoblot. In our infection test, the control cells, non-transduced and empty vector transfected only, did not propagate prion infection, whereas the bovine PRNP transduced cell lines were able to be infected.

Recently studies were reported that the plasma membrane is the initial or potential site of prion conversion [31,32], so this cell surface-expressed pattern may be considered as an important factor in establishing the persistently BSE-infected M2B cell line in this study.

Because of the low infection rate in infected cell clones, we added PMA to the culture media which activates protein kinase C (PKC) and increases expression levels of PrP\textsuperscript{C} in cells [33,34] potentially increasing the PrP\textsuperscript{Sc} detection rates. Furthermore, the highly sensitive cell-based infectivity assay, the scrapie cell assay (SCA), was employed and this assay can potentially detect single infected cells [16]. In this study, we obtained 6 strongly infected cell clones from five 96-well plates using SCA. On the basis of our result, the number of infected cells is very limited as previously reported [9,35], especially in uncloned cells where infection was lost upon continuous passaging compared with cloned cells (Fig. 3). Therefore in order to obtain persistently infected cells it is required to clone the cells in order to achieve cell lines with a high proportion of infected cells. These results could be explained (1) by the lack of stability of cells during prolonged propagation [16] or (2) by a kinetic argument, i.e. persistent infection of cell is only achieved when the rate of prion synthesis exceeds degradation [36].

When PrP\textsuperscript{Res} from M2B cells was compared to that of the original BSE inoculum, nonglycosylated PrP\textsuperscript{Res} derived from cells had a slightly smaller molecular weight (Fig. 5). This result is similar to the pattern of unglycosylated PrP\textsuperscript{Res} from SMB cell infected with the Chandler scrapie [37] and from Rov and MovS cells infected with the 127S scrapie [38]. The former reports suggested that the difference of migration pattern in WB might be resulted from conformational differences in between cell and host brain [37,39] or through differences in the composition of the GPI anchor[40,41]. More recently Dron et al. shows that PrP\textsuperscript{Sc} endogenous truncation may be greater in cell lines than in brain[38].
Additionally, we examined whether PrP\textsuperscript{BSE} from M2B cell transmits to conventional VM mice. PrP\textsuperscript{BSE} from M2B was successfully infected to VM mouse and its phenotypes were consistent with those of PrP\textsuperscript{BSE} classical BSE (data not shown). Furthermore, we have tried to apply the lentiviral system to produce other TSE agent-infected cells. A persistently CWD-infected cell line which was inoculated with naturally CWD-infected elk brain homogenate has been successfully developed in elk PrP expressing cells that were established using this lentiviral expression system [39].

In conclusion, we have demonstrated the generation of a persistently BSE-infected cell line which over-expresses bovine PrP based on the MDBK cell line. This successful achievement of generating BSE infected cells provides a methodology to potentially produce human prion infected cell models, including for vCJD, using a lentiviral expression system and also provides a research tool for research on the diagnosis of BSE and other related prion diseases as well as the study of potential therapeutic agents.

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Author Contributions
Conceived and designed the experiments: DT OW IC YJ. Performed the experiments: DT HK MHN MK YL HS. Analyzed the data: DT HK MHN OW. Contributed reagents/materials/analysis tools: DT HK. Wrote the paper: DT MHN OW.

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