Characterization of epidemic diarrhea outbreaks associated with bovine torovirus in adult cows

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Abstract Bovine torovirus (BToV) is recognized as an enteric pathogen of calves, but its etiological role in diarrhea and epidemiological characterization in adult cows remain unclear. In 2007-2008, three outbreaks of epidemic diarrhea occurred in adult cows at three dairy farms in Niigata Prefecture, Japan. BToV was the only enteric pathogen detected in these outbreaks, as determined by electron microscopy, reverse transcription-PCR, bacteria and parasite tests of fecal samples, and antibody tests with paired sera. The epidemiological features of the three outbreaks were similar to those of bovine coronavirus infection, except for the absence of bloody diarrhea, with diarrhea spreading among most adult cows, but not in calves, within several days and diarrhea lasting for 3-5 days with anorexia. Decreased milk production and mild respiratory symptoms were also observed in two of the outbreaks. Nucleotide sequence analysis of the BToV nucleocapsid, spike, and hemagglutinin-esterase (HE) genes revealed a close relatedness among the detected BToV strains from each outbreak and those of Japanese BToV strain Aichi/2004. Furthermore, we isolated a BToV strain, designated Niigata (TC), from a fecal sample using a human rectal tumor cell line. Sequence analysis of this isolate and Aichi/2004 indicated that both strains have truncated HE genes with deletions in the 3′ region that occurred through cell culture-adaptation. The short projections that are believed to be formed by the HE protein on virus particles were not observed in these cultured strains by electron microscopy. Taken together, these results suggest that BToV causes epidemic diarrhea in adult cows and should be included in the differential diagnosis of diarrhea in adult cows. In addition, our findings indicate that the HE protein of BToV may not be necessary for viral replication.

Introduction

Epidemic diarrhea in adult cattle is frequently observed worldwide, particularly among dairy cattle, and results in large economic losses from marked reductions in milk production. A major causative agent of adult cattle diarrhea is bovine coronavirus (BCV) [19, 24], which leads to the disease winter dysentery (WD). However, other viruses, including bovine torovirus (BToV), are also suspected to be etiologic agents of diarrhea [14, 15, 21], although the epidemiological situation of infections caused by these viruses in adult cattle remains unclear.

BToV, a member of the genus Torovirus within the family Coronaviridae, is a spherical, oval, elongated, or kidney-shaped enveloped virus and displays a double fringe of peplomers on its surface [3]. The virus has a single-stranded RNA genome of positive polarity encoding RNA polymerase and four structural proteins: the spike (S),
membrane (M), hemagglutinin-esterase (HE), and nucleocapsid (N) proteins [5]. Since the first isolation of BToV in 1982 from a case of neonatal calf diarrhea in the United States [34], several reports have confirmed the association of BToV with calf diarrhea in experimentally infected gnotobiotic calves and under field conditions [6, 8, 9, 13, 18]. Serological surveys have also revealed a high prevalence of antibodies to BToV in cattle populations, ranging between 55% and 95% [2, 14, 16, 33, 35]. Recently, Kuwabara et al. [16] succeeded in isolating BToV using a human rectal tumor cell line (HRT-18), which is expected to facilitate the diagnosis of BToV infection by virus neutralization tests and virus isolation.

BToV has been detected in fecal samples from adult cattle with diarrhea [10, 11, 21]. In addition, a few studies have reported that adult cows with WD-like diarrhea exhibited various degrees of seroconversion to BToV [14, 15]. These results suggest that BToV may be a causative agent of epidemic diarrhea of adult cattle. However, the epidemiological data for BToV infection in adult cattle under field conditions are limited, and the etiologic role of BToV in epidemic diarrhea of adult cattle remains unclear.

In the present study, we report the epidemiological features of three outbreaks of epidemic diarrhea in adult cows between May 2007 and February 2008 at three dairy farms in Niigata Prefecture, Japan, in which BToV was detected as the only pathogen. Further, we describe the isolation of a cytopathogenic BToV strain from diarrheic feces using HRT-18 cells, as well as the molecular characterization of the detected BToVs.

Materials and methods

Clinical specimens

Three epidemic outbreaks of adult cattle diarrhea occurred between May 2007 and February 2008 in Niigata Prefecture, Japan. A total of 16 fecal samples were collected from 16 diarrheic adult cows aged 24 to 71 months: 5 samples (Niigata1-1 to Niigata1-5) from outbreak 1 (Farm 1), 7 samples (Niigata2-1 to Niigata2-7) from outbreak 2 (Farm 2), and 4 samples (Niigata3-1 to Niigata3-4) from outbreak 3 (Farm 3). Nasal swabs were also collected at the time of fecal sampling from the 7 cows involved in outbreak 2. Paired serum samples were collected in the acute phase of diarrhea and 14 to 19 days later. Fecal samples were diluted 1:10 in 0.01 M phosphate-buffered saline (PBS; pH 7.4), and nasal swabs were suspended in 2 ml PBS. The resulting suspensions were clarified by low-speed centrifugation at 3,000×g for 10 min and used for virus isolation and reverse transcription-polymerase chain reaction (RT-PCR). Fecal samples were also tested for Salmonella species using a standard technique, while Coccidium species and Cryptosporidium species were detected by a sucrose floatation method. In addition, fecal samples were examined for rotavirus double-stranded RNA by polyacrylamide gel electrophoresis [17].

Virus-neutralization and hemagglutination inhibition tests

Virus neutralizing antibody titers against BToV in paired sera from affected cows were measured using HRT-18 cells and the BToV strain Aichi/2004 [16]. Virus neutralization (VN) tests were also conducted for bovine viral diarrhea (BVDV) type 1, bovine adenovirus type 3 (BAdV-3), bovine herpesvirus 1 (BHV-1), and bovine respiratory syncytial virus (BRSV) as described previously [36]. Antibody titers against BCV, adenovirus type 7 (BAdV-7), and bovine parainfluenza virus type 3 (BPIV-3) were determined by hemagglutination inhibition (HI) tests [20]. Seroconversion was defined as a fourfold or greater increase in paired serum antibody titers to the examined virus.

Electron microscopy

Fecal suspensions and the supernatants of infected cell cultures were partially purified by ultracentrifugation through a 30% (wt/wt) sucrose cushion, negatively stained with 2% sodium phosphotungstic acid (pH 7.0), and examined using an electron microscope (JEM-100S; JEOL, Ltd., Tokyo, Japan).

Virus isolation

Two lines of human rectal tumor (HRT-18) and Madin-Darby bovine kidney (MDBK) cells were used for virus isolation. One line of HRT-18 cells (designated HRT-18 Aichi cells), which had been maintained in Aichi Prefecture, Japan, was used for the first isolation of BToV in cell culture [16], and the second HRT-18 cell line (designated HRT-18 Niigata cells) had been maintained in our laboratory. Confluent monolayers of these cells in 24-well plates were washed with Eagle’s MEM (EMEM) and then inoculated with 0.1 ml of the fecal suspensions diluted from 1:10 to 1:1000 in EMEM with and without 10 μg/ml trypsin (Type I; Sigma-Aldrich, St. Louis, MO, USA). After adsorption for 60 min at 37°C, the cells were washed once with EMEM. 0.5 ml EMEM was added, and they were further incubated for 4 or 5 days at 37°C and examined for cytopathic effects (CPE). After incubation, the cells and culture supernatant were frozen and thawed once to harvest cell lysates, and subsequent passages were carried out in the same manner with 0.1 ml of cell lysate. After
two passages, the viral isolate was cloned three times in HRT-18 cells by limiting dilution.

The isolate was identified by indirect immunofluorescence and VN tests with gnotobiotic calf antisera against BToV (kindly supplied by Dr. Linda Saif, The Ohio State University) [9] and BCV as described previously [16, 28]. For VN tests, BToV strain Aichi/2004 [16] and BCV strain Kakegawa [1] were used as reference viruses. Electron microscopy and RT-PCR were also performed for the identification of the isolate.

RT-PCR

Viral RNA was extracted from fecal suspensions, nasal swabs, and infected cell culture supernatants using TRIzol LS Regent (Invitrogen Corp., Carlsbad, CA, USA) according to the manufacturer’s instructions. RT-PCR assays were performed using a OneStep RT-PCR kit (QIAGEN, Valencia, CA, USA) and primer pairs targeting fragments of the N-, S-, and HE-specific genes of torovirus. The primers, which were identical to those described previously by Hoet et al. [10] and Smits et al. [23], were as follows: 1344 (5'-GAGAAAGAGCAAGATGAATT-3'; positions 27761 to 27781) and 294 (5'-CTTACATGGACACTCAAAC-3'; positions 28403 to 28424) for N genes, primers S5 (5'-GTGTTAAGTTGTGCAATAAT-3'; positions 20956 to 20976) and S3 (5'-TGACAACTCTATATGCT-3'; positions 21677 to 21697) for S genes, and primers 1433 (5'-AAGTTTGAGTAGCCATTTTAC-3'; positions 26448 to 26468) and 1383 (5'-TAGCATTATTGATAGCATAGA-3'; positions 27781 to 27801) for HE genes. The positions were based on the genome of BToV strain Breda1 (AY427798). RT-PCR was also performed for detection of the specific genes of BCV [30], group A rotavirus (GAR) [7], group B rotavirus (GBR) [4], group C rotavirus (GCR) [29], and BVDV [32].

Sequence analysis

PCR products of the N, S, and HE genes of BToV were sequenced directly by cycle sequencing with an automatic sequencer (ABI PRISM 3100; Applied Biosystems, Tokyo, Japan). Sequencing was also performed for the HE gene of strain Aichi/2004 [16], which was passaged nine times in HRT-18 Aichi cells, and for the N, S, and HE genes from the large-intestinal content of the original specimen from which Aichi/2004 was isolated, designated as Aichi/2004(LIC). Sequence data were analyzed by the Clustal W method [26] using the Megalign 7.2 program of Lasergene software (DNASTAR, Madison, WI, USA). Phylogenetic trees were constructed using the MEGA 4 program [25].

Nucleotide sequence accession numbers

The newly determined sequences have been deposited in the DDBJ nucleotide sequence database and assigned the following accession numbers: Niigata1 (N gene, AB661450; S gene, AB661453; HE gene, AB661456), Niigata2 (N gene, AB661451; S gene, AB661454; HE gene, AB661457), Niigata3 (N gene, AB661452; S gene, AB661455; HE gene, AB661458), Niigata3(TC) (HE gene, AB661459), Aichi/2004(LIC) (HE gene, AB661461), and Aichi/2004 (HE gene, AB661460).

Results

Epidemiological situation of diarrhea outbreaks

Outbreak 1

On May 16, 2007, diarrhea and anorexia were observed in a few adult lactating cows on a dairy farm (Farm 1), with 12 of 24 adult cows on the farm displaying similar symptoms within several days. The diarrheal feces were watery and brownish. All affected cows recovered from diarrhea within 4-5 days, but milk production had decreased by up to 7% on the second day following the first finding of diarrhea and continued for one week.

Outbreak 2

On November 24, 2007, a few adult lactating cows on a dairy farm (Farm 2) showed pasty or watery diarrhea that was accompanied by nasal discharge. Within 4 days, 15 of 47 adult cows showed diarrhea and most of them had nasal discharge. Milk production decreased by approximately 20% on the fourth day after the onset of diarrhea. All affected cows recovered from diarrhea within 4-5 days. The farm was located approximately 80 km from Farm 1. No epidemiological relationship was observed between Farms 1 and 2.

Outbreak 3

On February 15, 2008, symptoms of pasty or watery diarrhea, anorexia, and mild cough were observed in a few adult lactating cows on a dairy farm (Farm 3), located 2 km away from Farm 1. Within one week, all 22 adult cows on the farm showed similar clinical signs. All cows recovered from diarrhea after 3-4 days, and no clinical signs were observed on the tenth day after the first finding of diarrhea. Milk production did not change during the outbreak.

Several common epidemiological features were observed among the three outbreaks. Affected cows did not show
bloody diarrhea or fever, and each cow recovered within 3-5 days after the onset of diarrhea without clinical treatment. Notably, no clinical signs, including diarrhea, were observed in calves during these outbreaks of adult cow diarrhea.

Examination of fecal samples

All 16 fecal samples were RT-PCR negative for BCV, BVDV, GAR, GBR, and GCR, and were also negative for species of *Salmonella*, *Coccidium*, and *Cryptosporidium*. In addition, none of the samples were positive for rotavirus RNAs by polyacrylamide gel electrophoresis. In contrast, all fecal samples except for Niigata2-2 were positive for N, S, and HE genes of BToV by RT-PCR. One nasal swab sample from outbreak 2 was also positive for the BToV N gene by RT-PCR, while we could not detect the S and HE genes of BToV in any nasal samples.

Virus antibody tests

Examination of paired sera from 14 of 16 affected cows showed seroconversion (≥4-fold increase) to BToV, and serum samples from the remaining two cows had antibody titers of ≥512 at the acute phase (Table 1). No significant change in BCV antibody titers was observed in any of the outbreaks (Table 1). Additionally, antibody titers of the other viruses examined (BVDV type 1, BAdV-3, BAdV-7, BHV-1, BRSV, and BPIV-3) did not markedly change in any of the paired sera.

Virus isolation in HRT-18 Aichi cells

After the second passage, CPE appeared in HRT-18 Aichi cells inoculated with one fecal sample, Niigata3-4, diluted 1:100 with EMEM containing trypsin. CPE, which was characterized by the enlargement of cells, was observed 2 to 3 days after inoculation, with the cells eventually detaching from the plastic surface. The isolate was cloned by limiting dilution three times, and the infective titers reached $10^{5.5}$ TCID$_{50}$/ml by passage 7. Indirect immunofluorescence microscopy of HRT-18 Aichi cells that had been inoculated with the isolate and reacted with gnotobiotic calf antiserum against BToV revealed cytoplasmic fluorescence, indicating the presence of BToV antigens [16]. In contrast, no fluorescent cells were observed when the infected cells were reacted with anti-BCV antiserum.

In VN tests, antiserum to BToV neutralized strain Aichi/2004 and the isolate at titers of 25,600 and 12,800, respectively. In contrast, antiserum to BCV failed to neutralize either strain (titer, <50). In addition, the isolate was positive in RT-PCR targeting the N, S, and HE genes of BToV. Electron microscopy and nucleotide sequencing of

| Table 1 | BToV neutralizing antibody and BCV hemagglutination inhibition antibody titers in adult cows with epidemic diarrhea at three outbreaks |
|---------|---------------------------------------------------------------------------------|
| Outbreak no. | Sample name | Age (months) | Sampling dates$^a$ | Antibody titer$^b$ |
|           |              |              |                      | BToV                     |
|           |              |              |                      | Acute phase | Convalescent phase | BCV                  |
|           |              |              |                      | Acute phase | Convalescent phase |                      |
| 1         | Niigata1-1   | 31           | 2007/5/18, 6/6/2007  | ≥2,048      | ≥2,048              | 40                  |
| 1         | Niigata1-2   | 29           | 2                    | <2          | 2,048              | 20                  |
| 1         | Niigata1-3   | 29           | 2                    | 2           | ≥2,048              | <10                 |
| 1         | Niigata1-4   | 29           | 2                    | 2           | 2,024               | <10                 |
| 1         | Niigata1-5   | 29           | 2                    | 2           | ≥2,048              | <10                 |
| 1         | Niigata2-1   | 29           | 2007/11/28, 12/12/2007 | 2           | 128                 | 40                  |
| 2         | Niigata2-2   | 59           | 2007/11/28, 12/12/2007 | 512         | 2,048               | 5,120               |
| 2         | Niigata2-3   | 26           | 2                    | 2           | 64                  | 40                  |
| 2         | Niigata2-4   | 71           | 2                    | 64          | 1,024               | 2,560               |
| 2         | Niigata2-5   | 66           | 2                    | 32          | 2,048               | 640                 |
| 2         | Niigata2-6   | 24           | 2                    | 32          | 256                 | 20                  |
| 2         | Niigata2-7   | 45           | 2                    | 64          | 2,048               | 5,120               |
| 3         | Niigata3-1   | 41           | 2008/2/18, 3/5/2008  | 64          | 1,024               | 640                 |
| 3         | Niigata3-2   | 50           | 2                    | 8           | 256                 | 160                 |
| 3         | Niigata3-3   | 43           | 2                    | 512         | 256                 | 160                 |
| 3         | Niigata3-4   | 24           | 2                    | <2          | 64                  | 80                  |

$^a$ Acute phase, convalescent phase

$^b$ Paired sera showing seroconversion (≥4-fold increase) are in bold numbers

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the RT-PCR products were also conducted for identification of the isolate. From these results, we concluded that the isolate from fecal sample Niigata3-4 was BToV and designated the new isolate as strain Niigata3(TC). In contrast, no cytopathogenic viruses were isolated from any samples using HRT-18 Niigata or MDBK cells.

Electron microscopy

Coronavirus-like particles were observed by electron microscopy in 5 of 16 fecal samples (Niigata1-2, -3, -4; Niigata2-3, and Niigata3-2) from the three outbreaks. The particles were spherical, oval, elongated, or kidney-shaped, approximately 100-140 nm in diameter, and surrounded by club-shaped projections of 15-20 nm in length. The particles also had short projections of approximately 5 nm in length (Fig. 1a, b). No other virus-like particles were observed in any of the fecal samples.

In the culture supernatants of infected HTR-18 Aichi cells, coronavirus-like particles with long club-shaped projections similar to those detected in the fecal specimens were observed. However, short projections were not seen on these particles (Fig. 1c, d).

Analysis of BToV genes

The nucleotide sequences of the N (485 nucleotides [nt]), S (673 nt), and HE genes (1260 nt) of BToV from nine fecal samples obtained from the three farms and BToV isolate Niigata3(TC) were compared with those of published toroviruses. The nine selected fecal samples were as follows: Niigata1-1, -2, and -3 from outbreak 1, Niigata2-3 and -4 from outbreak 2, and Niigata3-1, -2, -3, and -4 from outbreak 3. In addition, the nucleotide sequences of the viral genes of Aichi/2004 and Aichi/2004(LIC) were compared (Fig. 2).

As the nucleotide sequences of BToV genes from the same outbreak were identical, BToV detected from the feces in each respective outbreak was designated as strains Niigata1, Niigata2, and Niigata3. For the N gene, the nucleotide sequences of Niigata1 and Niigata3 were identical to that of the B145 strain detected in the Netherlands and showed 99% nucleotide identity to Niigata2. For the S gene, Niigata2 and Niigata3 showed 99% nucleotide identity, and they displayed 96%-97% nucleotide identity to Niigata1 and Aichi/2004. For the HE gene, Niigata1, Niigata2, and Niigata3 showed 98%-99% nucleotide identity to each other and 98%-99% nucleotide identity to Aichi/2004. In addition, the nucleotide sequences of both the N and S genes between Niigata3 and Niigata3(TC) and between Aichi/2004(LIC) and Aichi/2004 showed 100% identity.

Phylogenetic analysis of the N, S, and HE genes indicated that strains Niigata1, Niigata2, and Niigata3 were grouped within the same cluster as other BToV strains, including Aichi/2004 and B145 (Fig. 2). In the HE gene, the detected open reading frames of strains Niigata1, Niigata2, Niigata3, and Aichi/2004(LIC) encoded 419 amino acids, which is identical to the number reported previously for other strains of BToV. In contrast, both cell-culture-adapted Aichi/2004 and Niigata3(TC) strains of BToV had incomplete HE open reading frames that only encoded 160 and 104 amino acids, respectively (Fig. 3).

Discussion

BToV has been suspected to be a causative agent of adult cattle diarrhea based on its detection in diarrheic feces and because various degrees of seroconversion to BToV have been observed in outbreaks of WD-like diarrhea in adult cows [10, 11, 14, 15, 31]. However, prior to the present study, no conclusive evidence for a link between BToV and epidemic diarrhea in adult cattle had been reported. Furthermore, epidemiological characterization of these diarrheal outbreaks remained unclear. Here, we examined the etiological agents and conducted detailed serological tests with samples from three outbreaks of adult cattle diarrhea in Niigata, Japan, and found that BToV was the only causative pathogen.
In the three outbreaks examined, most affected cows developed watery diarrhea, anorexia, and decreased milk production, and additionally, some cows also showed slight respiratory signs. The observed clinical signs were similar to those encountered in WD and BCV infections. Notably, none of the calves developed diarrhea during the outbreaks, a diagnostic characteristic that is also observed in WD and BCV infections in adult cows [27, 31]. Taken together, these results suggest that adult cattle diarrhea caused by BToV may have been misdiagnosed as WD or BCV infection by clinical investigations.

We isolated the cytopathogenic BToV strain Niigata3(TC) using HRT-18 cells. Of note, a clear difference in sensitivity to BToV was observed between the two HRT-18 cell lines used in this study. Namely, the Niigata3(TC) strain could be isolated and propagated in HRT-18 Aichi cells, but not in HRT-18 Niigata cells. Similarly, strain Aichi2004 could also not be propagated in HRT-18 Niigata cells, although both HRT-18 cell lines displayed high sensitivity to BCV (data not shown). These results indicate that the use of HRT-18 Aichi cells is critical for the in vitro propagation of BToV. Although the reason for
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Fig. 3  Sequences of the HE genes of BToV detected in this study. Aichi/2004 and Niigata3(TC) are cell-culture-adapted BToV strains isolated using HRT-18 Aichi cells. All other strains were detected in fecal specimens. Open boxes indicate AUG initiation codons, and gray boxes indicate termination codons of ORFs. Accession numbers of the nucleotide sequences are indicated in the legend of Fig. 2.
the different sensitivity between the two HRT-18 cell lines to BToV is unknown, it is possible that the properties of the HRT-18 Aichi cells may have changed during passaging to increase BToV tropism.

BToVs have been classified into two serotypes on the basis of reactivity in the hemagglutination/HI test [35]. Recently, however, BToVs have been proposed to consist of a single serotype that includes at least two subtypes [12]. In the present study, rabbit antisera to strain Aichi/2004 showed no significant differences in virus neutralizing antibody titers between the Aichi/2004 and Niigata3(TC) strains, suggesting that these two strains belong to the same serotype. Nucleotide sequence analysis indicated that the BToV strains detected in the fecal samples from three outbreaks in Niigata prefecture were closely related to each other and to strain Aichi/2004. This finding, together with the serological results, suggests that these Japanese BToV strains may have evolved from a recent common ancestor.

Our sequencing analyses revealed that both cell-culture-adapted BToV strains, Niigata3(TC) and Aichi/2004, possessed truncated HE genes with deletions in the 3′ region that occurred through cell culture-adaptation. Electron microscopy photographs indicated that virus particles in fecal samples had long and short projections, whereas short projections were not seen on those in the culture supernatants of infected HTR-18 cells. Interestingly, in many laboratories strains of mouse hepatitis virus, the HE genes are inactivated, which apparently represents an in vitro artifact resulting from adaptation to propagation in cultured cells [37]. It is noteworthy that cell-culture-adapted equine torovirus Berne virus also possesses a truncated HE gene with deletions in the 5′ region [22]. Taken together with the data presented in this study, the HE protein in BToV may be unnecessary for viral replication. Further examination of the biological properties of the HE protein in fecal- and cell-culture-adapted strains of BToV is needed to confirm this speculation.

In conclusion, we presented the epidemiological characteristics of BToV diarrhea in adult cattle based on three outbreaks in Niigata, Japan. We propose that examination for BToV should be included in the routine diagnosis of adult cattle diarrhea.

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