First isolation and genetic characterization of pseudocowpox virus from cattle in Japan

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Abstract

Background: Pseudocowpox virus (PCPV) infects cattle worldwide with zoonotic potential but has not been isolated in Japan. Thus, the epidemiological status of PCPV infection in cattle is undetermined.

Results: In May 2016, a cattle in a farm in Yamaguchi Prefecture showed white vesicles and hyperemia in the mucosa under the tongue surface, but not on the teats and coronary cushions. A parapoxvirus was isolated from the oral lesion swab and was genetically characterized based on the full-length sequence of B2L gene encoding viral envelope. Phylogenetic analysis showed that the isolated virus was classified into PCPV.

Conclusion: This case indicates its potential spread in Japan. This is the first report of isolation of PCPV in Japan.

Keywords: Pseudocowpox virus, Cattle, Oral lesions, Isolation, B2L gene

Background

Pseudocowpox virus (PCPV) is a member of the genus Parapoxvirus in the family Poxviridae, which includes bovine papular stomatitis virus (BPSV) and orf virus (ORFV) [1]. Parapoxviruses are commonly known as causative agents of dermal diseases in ruminants worldwide, leading to papular stomatitis and contagious pustular dermatitis, especially in the regions of the lips, nostrils, oral mucosa, and teats. The importance of PCPV is increasingly recognized, primarily because of economic losses to farmers in connection with disease outbreaks and because of their zoonotic potential [2].

In Japan, although serological surveys have revealed that seroprevalence of parapoxvirus is very high in cattle and sheep [3, 4] and multiple BPSVs have been isolated [5], no PCPV has yet been isolated; thus, the epidemiological status of PCPV infection in cattle is undetermined.

We here report the first case of the isolation of PCPV in Japan. We determined the full-length sequence of the B2L gene encoding viral envelope of this isolate, and evaluated its phylogenetic relation to known members of this virus group.

Methods

Clinical and epidemiological investigations

In May 2016, a breeding cow (Japanese Black, female, 13-month old) in a farm in Yamaguchi Prefecture, in the western part of Japan showed anorexia, mild fever, frothy salivation, and hyperemia in the mucosa under the tongue surface. No lesions were observed on the teats or coronary cushions. A few days after the onset of clinical signs, the cattle showed white vesicles on the tongue surface (Fig. 1). These signs were convalescent in about 1 week. No other cattle in the herd showed clinical signs.

Sample collection

An oral swab sample was collected from the mucosal lesions of the affected cattle and was homogenized with Eagle's minimum essential medium (MEM). The sample was centrifuged at 800×g for 10 min at 4 °C. The supernatant was filtered through a 450-nm membrane (Merck Millipore, Cork, Ireland) and used for virus isolation and DNA extraction.

Virus isolation

Virus isolation was performed through inoculation in primary bovine testis (BT) cells; two cell lines, hamster lung (HmLu-1) and Madin-Darby bovine kidney (MDBK) cells were also tested for comparison. The cells were grown in Eagle’s MEM with kanamycin (Nissui Pharmaceutical, Tokyo, Japan) supplemented with 0.295%
tryptose phosphate broth (TPB), 0.015% sodium bicarbonate, 0.03% L-glutamine, and 5–10% fetal bovine serum at 37 °C. The cells were cultured in rolling tubes and 24-well plates, washed with maintenance medium (MEM containing 0.295% TPB, 0.015% sodium bicarbonate, 0.03% L-glutamine, and 0.1% bovine serum albumin), and inoculated with 0.1 ml of the processed samples. After adsorption for 60 min at 37 °C, the inocula were replaced with 0.5 ml of maintenance medium. The rolling tube-cultured and 24-well plates-cultured cells were incubated by rotary cultures and stationary cultures, respectively, and observed daily for cytopathic effects (CPE) for at least 7 days. Cultures without CPE were passaged twice in a blinded manner.

Genetic analysis

DNA was extracted both from the oral swab sample and from the BT cells showing a CPE using magLEAD 12gC (Precision System Science, Chiba, Japan). Polymerase chain reaction (PCR) amplifications were carried out with TaKaRa Ex Taq Hot Start Version (TaKaRa Bio, Shiga, Japan) using a SimpliAmp Thermal Cycler (Thermo Fisher Scientific, Kanagawa, Japan) for detection of the full-length (1137 bp) and partial-length (554 bp) B2L gene encoding envelope of parapoxvirus with the primer sets OVB2LF1/OVB2LR1 [6] and PPV1/PPV4 [5], respectively. PCR products were purified using NucleoSpin Gel and PCR Clean-up (Macherey-Nagel, Düren, Germany), and the nucleotide sequence was determined by direct sequencing using a BigDye Terminator Cycle Sequencing Kit v3.1 (Applied Biosystems, Austin, TX, U.S.A.). Sequence data were aligned using the ClustalW method [7]. Phylogenetic analysis was performed using MEGA 6 software [8]. Phylogenetic trees were constructed using maximum-likelihood methods, and the reliability of the branches was evaluated by bootstrapping with 1000 replicates. Nucleotide and deduced amino acid sequences were compared with those of available corresponding parapoxviruses (Table 1). Bovine viral diarrhea virus [9], epizootic hemorrhagic disease virus [10], bluetongue virus [10], ovine herpesvirus 2 [11], and bovine herpesvirus 1 [12], were not detected by PCR using specific primers for detection of each viruses (data not shown).

Results

At the third passage, a distinct CPE was observed from 2 to 3 days after inoculation in the rolling tube-cultured BT cells, characterized by a rounded morphology and cell detachment (Fig. 2). We designated the isolate as strain YG2828. However, at the third passage, no CPE appeared in the other cell lines or in stationary cultures. For histological observations, confluent monolayers of BT cells in the chamber slide system were inoculated with the isolate. Twenty-four hours after inoculation, the BT cells were fixed with acetone and stained with hematoxylin-eosin, which revealed pyknosis, and eosinophilic and basophilic cytoplasmic inclusion bodies (data not shown).

Fragments of expected size were amplified by PCR using both primer sets. Neither deletions nor insertions in the nucleotide sequence of the YG2828 strain were found (Additional file 1: Figure S1). Based on the nucleotide/amino acid identities and phylogenetic analysis of the full-length B2L gene, the YG2828 strain was classified as PCPV (Fig. 3). The nucleotide identities against published parapoxviruses ranged from 85.8 to 98.6% (Table 1), and showed the highest identity (98.6%) to three PCPV strains: F05.990C and F10.3081C isolated from cattle in 2005 and 2010 in Finland, respectively, and VR634 isolated from a human in the USA in 1963 with "milker’s nodules" on the hands. The deduced amino acid identities ranged from 84.4 to 99.2%, and showed the highest identity (99.2%) to the VR634 strain (Table 1), even though these strains were isolated independently, chronologically, and geographically.

Discussion

As noted above, seroprevalence of parapoxvirus is very high in cattle in Japan and multiple BPSVs have been isolated [5], but no PCPV has yet been isolated. In this study, a PCPV was firstly isolated in Japan by rotary cultures, but not stationary cultures. Similarly, Mavromoustakis et al. [13] reported that significantly (P < 0.01) less herpes simplex virus was produced in stationary than in rotary cultures. Although the procedures of rotary cultures are more burdensome than those of static cultures, we suggest that rolling of inoculated cultures should be conventionally applied in clinical virology laboratories to aid in the isolation of PCPV. In the affected cattle in this study, there was no evidence of infection on the teats and udder, which are the more common lesion sites of pseudocowpox infection. The classification of parapoxviruses was
formerly based on the natural host range, clinical signs, and serology [14], however this does not always reflect the classification revealed by molecular analysis [5] as evidenced by the present study.

Previously, there has only been one report describing the PCR detection of PCPV DNA in Japan, in which PCPV DNA was detected from oral lesions in a calf in Iwate Prefecture, in the northern part of Japan, but virus isolation was unsuccessful [15]. Notably, the partial-length sequence of the B2L gene determined from the PCR product (accession no. AB921003) was identical to that of the present strain YG2828 (data not shown). Thus, our results confirmed that PCPV can be isolated from atypical sites besides the teats and udder, and suggest that YG2828-like PCPV may cause oral lesions in cattle. Moreover, since parapoxviruses cross-react antigenically and two similar strains infected cattle in different locations in separate years, YG2828-like PCPV might be spreading among the cattle population in Japan. It is known that cattle are frequently infected with parapoxvirus subclinically [15] and PCPV has zoonotic potential [2]. Therefore, we recommend to wear gloves for people with regular exposure to cattle mucosa.

**Conclusion**

A PCPV was firstly isolated in Japan from the oral lesion swab of cattle showing white vesicles and hyperemia in

| Virus* | Strain | Host | Nucleotide | Amino acid | Accession number |
|--------|--------|------|------------|------------|-----------------|
| PCPV   | YG2828 | Cattle | –          | –          | LC230119        |
|        | VR634  | Cattle | 98.6       | 99.2       | GQ329670        |
|        | TQ     | Cattle | 98.2       | 98.7       | AY424972        |
|        | F07.798R | Reindeer | 98.4     | 98.9       | JF773692        |
|        | F07.801R | Reindeer | 98.4     | 98.9       | JF773693        |
|        | F05.990C | Cattle | 98.6       | 98.7       | JF773694        |
|        | F10.3081C | Cattle | 98.6       | 98.7       | JF773695        |
|        | F00.120R | Reindeer | 98.4     | 98.9       | GQ329669        |
|        | Arero/05/2013 | Camel | 97.2       | 97.6       | KU645549        |
|        | Arero/04/2013 | Camel | 97.0       | 97.1       | KU645548        |
|        | Arero/02/2014 | Camel | 97.2       | 97.6       | KU645546        |
|        | Hordha/01/2011 | Camel | 97.0       | 97.1       | KU645563        |
| ORFV   | NZ2    | Sheep | 94.1       | 95.0       | DQ184476        |
|        | F07.808R | Reindeer | 93.7     | 93.9       | JF773698        |
|        | SD/DY  | Sheep | 93.9       | 93.7       | JQ904794        |
|        | Adet/O03/2012 | Sheep | 94.0       | 95.0       | KT438515        |
| BPSV   | RS     | Cattle | 85.8       | 84.4       | AY424973        |
|        | BV-TX09c1 | Cattle | 85.8       | 84.7       | KMB75472        |
|        | BV-TX09c5 | Cattle | 85.8       | 84.7       | KMB75471        |
|        | BV-AR02 | Cattle | 86.1       | 84.7       | AY386265        |

*PCPV, pseudocowpox virus; ORFV, orf virus; BPSV, bovine papular stomatitis virus

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**Fig. 2** Cytopathic effect observed in BT cells at passage 3. The cells were tested at day 3 after inoculation. Non-infected control (**a**) and infected (**b**) cells are shown.
the mucosa under the tongue surface, but not on the teats and coronary cushions, by rotary cultures. Genetic characterization based on the full-length sequence of B2L gene revealed that the isolated virus was genetically close to strains isolated from cattle in the USA and Finland. PCPV is responsible for significant economic losses in the cattle production. Further virological and epidemiological studies to characterize this strain and the possibility of its spread in Japan are highly required.

**Additional file**

**Additional file 1: Figure S1.** Alignment of the deduced amino acid sequences of the full-length B2L gene. Amino acids identical to the pseudocowpox virus strain YG2828 at given positions are represented by dots. (PPTX 73 kb)

**Abbreviations**

BPSV: Bovine papular stomatitis virus; BT: Bovine testis; CPE: Cytotoxic effects; HmLu-1: Hamster lung; MDBK: Madin-Darby bovine kidney; MEM: Minimum essential medium; ORFV: Orf virus; PCPV: Pseudocowpox virus; PCR: Polymerase chain reaction; TPB: Tryptose phosphate broth

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**Availability of data and materials**

Sequence data obtained in this study is available in the GenBank (accession no. LC230119).

**Authors’ contributions**

AO and YI analyzed all data in the experiment and were major contributors in writing the manuscript. AO and AY performed virus isolation and PCR experiments from samples from the affected cattle. AO and YI performed the nucleotide/amino acid identities and phylogenetic analysis. HN performed the histological experiment. All authors read and approved the final manuscript.

**Ethics approval**

This study was approved by the Gifu University Animal Care and Use Committee (Approval number 14094). Sampling of oral swab was performed with informed owner consent.

**Consent for publication**

Not applicable

**Competing interests**

All authors declare that they have no competing interests.

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