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Isolation of avian bornaviruses from psittacine birds using QT6 quail cells in Japan

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ISOLATION OF AVIAN BORNAVIRUSES
ABSTRACT

Avian bornaviruses (ABVs) were recently discovered as the causative agents of proventricular dilatation disease (PDD). Although molecular epidemiological studies revealed that ABVs exist in Japan, no Japanese isolate has been reported thus far. In this study, we isolated four strains of *Psittaciform 1 bornavirus* from psittacine birds affected by PDD using QT6 quail cells. To our knowledge, this is the first report to isolate ABVs in Japan and to show that QT6 cells are available for ABV isolation. These isolates and QT6 cells would be powerful tools for elucidating the fundamental biology and pathogenicity of ABVs.

KEY WORDS

avian bornavirus, isolation, parrot bornavirus, *Psittaciform 1 bornavirus*, QT6 cells
In 2008, two independent groups discovered novel bornaviruses, named avian bornaviruses (ABVs), from psittacine birds affected by proventricular dilatation disease (PDD) [3, 6], which is often fatal and accompanied with gastrointestinal dysfunction and/or neurologic symptoms. Recent extensive epidemiological studies revealed that 15 types of ABV were detected from many species of birds all over the world [7, 13].

Virus isolation is important for diagnosis of ABV infection as well as fundamental studies of ABVs. For example, Gray et al. used a parrot bornavirus (PaBV) isolate for infection experiments, which fulfilled the Koch’s postulates and demonstrated that ABVs are indeed the causative agent of PDD [2]. In addition, bornaviruses are known to be unique RNA viruses in that they establish non-cytolytic, persistent infections in the host cell nucleus. Therefore, studies of ABVs may also provide interesting insights into interactions between RNA viruses and their hosts as shown by Borna disease virus researches [4, 9]. In Japan, although we and others detected ABV nucleic acids from several birds [5, 12, 16, 17], no one has isolated ABV so far. Ogawa et al. tried to isolate an ABV from 10% cerebrum homogenate of an infected bird using Japanese quail fibroblast cells (QE-1 cells) [12], and we also inoculated a fecal sample, which was positive for PaBV-5 nucleic acid, to QT6 Japanese quail fibroblast-like cells [11], but the attempts failed [5]. Although the above trials failed and there is no report showing that Japanese quail is infected by ABVs, quail-derived cell cultures were reported to be available for isolation of ABVs [14]. Thus, QT6 may be also susceptible to ABV infection.

In the course of molecular epidemiological studies of ABVs, we got four brain samples from parrots affected with PDD (Table 1). We performed RT-PCR using random hexamer and primers MH175 and MH170 which amplify 221 bp segment of
ABV N gene [5], and sequencing analyses, revealing that the parrots were infected with *Psittaciform 1 bornavirus*: two were PaBV-2, and the other two were PaBV-4 (Table 1) (full methods and primer sequences are available in the supplementary information).

Therefore, we used these brain samples for isolation of ABVs. Inocula were prepared as described by Rubbenstroth *et al.* [15] with slight modifications (supplementary information), which were added to semi-confluent QT6 cells. The inoculated cells were passaged as described in the supplementary information.

After 3 to 12 weeks post inoculation, we performed indirect immunofluorescent assay (IFA) using anti-Borna disease virus 1 (BoDV-1) P antibodies (supplementary information). Positive signals were found in the inoculated QT6 cells, but not control cells, indicating that the QT6 cells were infected with bornaviruses (Fig. 1). We named these isolated strains as IH-1, KOKO, 7I6 and 7I10, respectively (Table 1).

To confirm the above observation, we extracted RNA from the cells and carried out RT-PCR using random hexamer and the primers MH175 and MH170 [5] for reverse-transcription and PCR, respectively (supplementary information). Specific bands were detected in all RNA samples extracted from the inoculated cells (Fig. 2a). We determined the nucleotide sequences of amplicons, showing the sequences were almost identical to those in brains (data not shown). We further performed western blotting using the rabbit antibodies against BoDV-1 P protein (supplementary information). The western blotting analysis detected specific bands of expected molecular weight in the inoculated cells (Fig. 2b). The difference of band intensities among the samples may be due to different infection rates, protein expression levels and/or affinities to the antibodies. These data demonstrated that we indeed isolated ABVs from the brain samples.
In this study, we isolated four strains of *Psittaciform 1 bornavirus* using QT6 quail cells. To our knowledge, this is the first report showing that QT6 cells are susceptible to ABVs, at least PaBV-2 and PaBV-4. So far, several avian cell lines were reported to be susceptible to ABVs [14, 15]. However, due to the national regulation for infectious diseases of poultry in Japan, it is time-consuming and sometimes difficult to import chicken and quail cells from abroad. Duck embryonic fibroblast (DEF) was also reported to be available for ABV isolation [2, 15], but it is also laborious to prepare cultures from fertilized eggs. In addition, for propagation of ABVs and preparation of virus stocks, it is usually necessary to passage inoculated cells for long term, since ABVs slowly propagate in cell culture. Because of the limited life span of DEF, low-passage DEF should be regularly provided to propagate ABVs as described in [2]. Therefore, our finding is useful for further isolation of ABVs at least in Japan and maybe also in other countries that have similar regulations. Although it is not sure that QT6 cells are susceptible to other types of ABVs, the cell line is probably available to isolate other types of ABVs because the QT6 cell line was reported to be susceptible to other avian viruses, such as avian influenza viruses [8]. Since it was reported that susceptibilities to ABVs differ among avian cell lines [15], it is interesting to compare the susceptibilities of QT6 cells and the previously reported cell lines. And, more, QT6 cells were also reported to show high transfection efficiency [1]. Therefore, the cells are helpful to study the molecular biology of ABVs as well as to establish reverse genetics system for ABVs.

As described above, we tried to isolate PaBV-5 using QT-6 cells, but it failed. This may be explained by the difference of materials we used to isolate the viruses. When trying to isolate PaBV-5, we inoculated a feces-derived sample to QT6 cells, which
showed much higher cytotoxicity than the brain-derived samples. In addition, in infected birds, brains usually contain higher amount of ABVs than feces [14]. Alternatively, QT6 cells might not be susceptible to PaBV-5. Further studies are needed to assess the possibilities.

Our isolates belong to PaBV-2 and PaBV-4, which are dominant types of ABV in the world [2] and were formally shown to cause PDD [2, 10]. Although many PaBV-2 and PaBV-4 have been isolated thus far, there is no report, which systematically investigates differences of viral genome sequences, replication efficiencies and pathogenicity among the strains. Although we have not yet determined the full genome sequences of isolated viruses, it may be interesting to compare their sequences and their properties in vitro and in vivo.

Taken together, we isolated four Japanese strains of ABVs from psittacine birds using QT6 cells. Our isolates and findings are useful for further investigation of the fundamental biology and pathology of ABVs.

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**FIGURE LEGENDS**

**Fig. 1. Visualization of bornaviral antigens by indirect immunofluorescence assay.** Immunofluorescence assays using anti-BoDV-1 P rabbit polyclonal antibodies. Bright fields and fluorescence images are shown. The cells and inocula were indicated on the left side of each panel. Inoculum 1, 12 weeks post inoculation (w. p. i.); inoculum 2, 3 w. p. i.; inoculum 3, 3 w. p. i.; and inoculum 4, 4 w. p. i. 144:

**Fig. 2. Detection of bornaviral nucleic acid and phosphoproteins by RT-PCR and western blotting.** (a) RT-PCR with primers MH175 and MH170. Cells and virus strains
were indicated above the figure. (b) Western blotting using anti-BoDV-1 P rabbit polyclonal antibodies. Cells and infected virus strains were shown above the figure.
Figure 1

Bright field Anti-BoDV-1-P

QT6/ control

QT6/ inoculum 1

QT6/ inoculum 2

QT6/ inoculum 3

QT6/ inoculum 4
Figure 2
| Inoculum number | Host               | Virus species                        | Virus | Strain | Sampling date  |
|----------------|--------------------|--------------------------------------|-------|--------|----------------|
| 1              | *Ara militaris*    | *Psittaciform 1 bornavirus*          | PaBV-2| IH-1   | August 2011    |
| 2              | *Aratinga jandaya* | *Psittaciform 1 bornavirus*          | PaBV-2| KOKO   | January 2013   |
| 3              | *Diopsittaca nobilis* | *Psittaciform 1 bornavirus*       | PaBV-4| 7I6    | February 2014  |
| 4              | *Ara ararauna*     | *Psittaciform 1 bornavirus*          | PaBV-4| 7110   | February 2014  |
Supplementary information

Materials and methods

**Virus isolation**

The inocula were prepared as described by Rubbenstroth *et al.* [2] with slight modifications. The brains were homogenized in DMEM containing 2% fetal calf serum (FCS), and then sonicated on ice. The homogenates were centrifuged at 10,000 × g for 10 min at 4°C. The supernatants were filtrated with 0.45 μm filter, which was used as the inocula. The inocula were added to semi-confluent QT6 cells, and the media were changed within 12 hours after the inoculation. The cells were first passaged after 24 hours, and then the cells were maintained in DMEM containing 10% FCS. The cells were passaged every three or four days.

**RT-PCR and sequencing analysis**

The RNAs were isolated from the brain samples and the inoculated cells using QIAamp Viral RNA Mini Kit (QIAGEN), High pure viral nucleic acid kit (Roche), or RNeasy Plus Mini Kit according to the manufacturer’s instructions. The extracted RNA samples were reverse-transcribed with random hexamer using Verso cDNA Synthesis Kit (Life Technologies) or PrimeScript RTase (TaKaRa) in accordance with the manufacturer’s protocol, which was used as the templates for the PCR analyses.

PCR was carried out using Ex Taq Hot Start Version (TaKaRa) and primers MH175 (5’-AARGARTAYTIAAYGARTGYATGGAYGC-3’) and MH170 (5’-GGRTTYTCTYTTTICTCCARTAAAAANGC-3’) [1] with the following cycles: initial denaturation at 94°C for 2 min, 40 cycles of 94°C for 30 sec, 55°C for 30 sec and 72°C for 20 sec, and final extension at 72°C for 3 min. The PCR products were analyzed by
agarose gel electrophoresis. For sequencing, the PCR products were purified with NucleoSpin Gel and PCR Clean-up (MACHEREY-NAGEL) or by ethanol precipitation. The purified PCR product of sample 1 (Table 1) was cloned with TOPO TA Cloning Kit for sequencing (Life Technologies) and sequenced with the M13 forward primer (5’-GTAAA ACACCGCCAGT-3’). The other PCR products were directly sequenced with primers MH175 or MH170.

*Immunofluorescence assay (IFA)*

For IFA, the inoculated cells were fixed with 4% paraformaldehyde for 10 min at room temperature (RT). The reaction was quenched with 0.1M glycine in PBS for 10 min at RT. The cells were permeabilized with 0.1% saponin in PBS for 10 min, and then blocked with 5% FCS in PBS for 30 min. The cells were incubated with rabbit anti-Borna disease virus 1 (BoDV-1) P antibodies diluted at 1:500 with 1% FCS in PBS for 1 h. After four times washing with PBS, the cells were reacted with anti-rabbit IgG-Alexa Fluor 488 (Life technologies) diluted at 1:1000 with 1% FCS in PBS for 1 h. The cells were washed four times with PBS, and stained with DAPI for 10 min. After five times washing, the images were obtained using LSM-710 confocal microscope (Zeiss).

*Western blotting*

The inoculated cells were once washed with PBS, and directly lysed with SDS sample buffer. The cell lysates were electrophoresed with 12% acrylamide gel, and then transferred to a PVDF membrane. The membrane was blocked with 3% skim-milk in Tris-buffered saline containing 0.1% Tween-20 (TBS-T). The blot was incubated with
rabbit anti-BoDV-1 P antibodies (HB03) diluted at 1:1000 with 3% skim-milk in TBS-T for 1 h at RT. After three times washing with TBS-T, the membrane was incubated with anti-rabbit-Ig conjugated with horseradish peroxidase (Dako) for 1 h at RT. After four times washing with TBS-T, the membrane was incubated with ECL Prime Western Blotting Detection System (GE healthcare), and analyzed by Fusion Solo S (Vilber Lourmat).

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