Quantification of bovine viral diarrhoea virus ribonucleic acid in serum of infected animals by one-step reverse transcriptase quantitative real-time polymerase chain reaction

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Abstract

Bovine viral diarrhoea virus (BVDV) can cause either acute transient or persistent infection. Identification and removal of persistently infected animals from infected herds is a crucial component to control BVDV infection. Only limited data on serum virus concentration in infected animals are available to date. Using one-step reverse transcriptase quantitative real-time polymerase chain reaction, we quantified the serum viral load in 40 BVDV infected animals. To control nucleic acid extraction, complementary DNA synthesis and polymerase chain reaction amplification, each serum sample was spiked with a known small amount of reference canine coronavirus. Detected ribonucleic acid copy number ranged from $2.2 \times 10^6$ to $7.4 \times 10^8$ per 1 ml of serum of persistently infected animals and from $6.6 \times 10^4$ to $3.3 \times 10^7$ of transiently infected animals. These findings support the idea that it is impossible to accurately distinguish between transiently and persistently infected animals just from a single blood sample. To use this testing as a means of declining costs of BVDV control programmes cannot be recommended and paired serum samples have to be investigated to confirm persistent infection.

Reverse transcription quantitative PCR, ruminant

Bovine viral diarrhoea virus (BVDV) is a member of the Pestivirus genus within the Flaviviridae family. Bovine viral diarrhoea viruses form two separate species, BVDV 1 and BVDV 2 which are further divided into several genetic subtypes. Pestiviruses are small enveloped viruses that contain single stranded positive sense RNA of approximately 12.5 Kb in length. The viral genome consists of a single open reading frame, flanked at both ends with short untranslated regions (5´UTR, 3´UTR). The ORF encodes one polyprotein which is subsequently cleaved into 12 individual viral proteins. According to the presence or absence of the cytopathic effect on cultured epithelial cells, BVDVs are divided into two different biotypes, cytopathic (cp) and noncytopathic (ncp).

Bovine viral diarrhoea is one of the most economically important diseases of cattle worldwide. Bovine viral diarrhoea virus infection can produce a broad range of clinical signs from mild to lethal mucosal disease or can infect animals without any obvious signs of disease. Bovine viral diarrhoea virus causes either acute transient (TI) or persistent (PI) infection (Baker 1995). The virus first replicates in epithelial cells of the affected mucosa and surrounding lymphoid tissue, which is followed by the spread to regional lymph nodes, viraemia and dissemination of the virus throughout the body (Howard 1990; Marshall et al. 1996; Bruschke et al. 1998). Acute infection results in short-term transient viraemia. Thereafter, the virus is cleared by the host immune system and specific antiviral antibodies are detected. In pregnant viraemic animals, the virus can cross the placenta and infect the foetus. Persistent infection develops when foetuses are exposed to ncp biotype of BVDV before they are immuno-competent on day 120 of gestation. This leads to a lack of recognition of viral antigens by the host immune system, persistent viraemia and replication of the virus to high titres in all tissues and organs (Ohmann 1988). Persistently infected
animals shed large amounts of virus in most secretions and excretions into the environment and are a major source of the infectious virus for their herd mates.

Identification and removal of PI animals from infected herds is a crucial component to control BVDV infection (Lindberg and Alenius 1999; Letellier et al. 2005; Lindberg and Houe 2005; Houe et al. 2006). Serum samples are widely used to identify PI animals (Saliki and Dubovi 2004; Sandvik 2005). Nowadays, viral antigens and/or viral RNA are detected in serum samples by antigen ELISA (enzyme-linked immunosorbent assay) and RT-PCR (reverse transcription polymerase chain reaction) or quantitative RT-qPCR. The RT-PCR has more advantages as it is not affected by the presence of antiviral antibodies and is also more sensitive than antigen ELISA. Pooled serum samples are investigated by PCR to reduce the costs of identification of PI individuals (Munoz-Zanzi et al. 2000; Mars and van Maanen 2005; Laureyns et al. 2010).

The aim of our study was to quantify and evaluate serum viral load in BVDV infected animals by the RT-qPCR targeting the 5’ UTR genome region and employing hydrolysis probes.

Materials and Methods

Serum samples

The serum samples were collected during a voluntary BVDV control/eradication program conducted in the Czech Republic in 2009–2011. To identify PI animals, animals older than 6 months were screened for antibodies using a commercially available ELISA kit (BVD-MD IgG ELISA, Test-line, Brno, Czech Republic). Thereafter, the antibody-negative animals were tested for virus by antigen ELISA (BVDV Antigen Test Kit/ Serum Plus, IDEXX, Bern, Switzerland). The antibody-negative virus-positive animals were retested at least 3 weeks apart. Repeated detection of viral antigen in antibody-negative cattle confirmed PI. On the contrary, seroconversion proved transient BVDV infection. A total of 77 serum samples from 40 infected animals were tested to quantify viral RNA (Table 1). Serum samples were stored at −80 °C until processed and used in RT-qPCR.

RNA extraction

Viral RNA was extracted from 140 µl of serum samples using QIAamp Viral RNA kit (Qiagen, Hilden, Germany). Immediately before RNA extraction, the serum samples were spiked with 10 µl of Canine Coronavirus (CCoV) strain 1-71 (ATCC VR-809, Collection of Animal Pathogenic Microorganisms, VUVeL, Brno, Czech Republic) representing 9.5 median tissue culture infective dose (TCID50). The amounts of reagents added to the sample before loading onto QIAamp spin columns were then proportionally increased as recommended by the supplier. RNA was eluted in a volume of 60 µl and extracted RNA samples were stored until used in RT-qPCR at −80 °C.

Primers and probe

Sequences of primers and probe were selected from 5’ UTR of the BVDV genome and were based on primers and probe designed previously by Mahlum et al. (2002) and Young et al. (2006), respectively. After alignment of 226 sequences deposited in the GenBank, degeneracy was incorporated into specific nucleotide sites of forward (BVDV 183-F) and reverse (BVDV 343-R) primer. Both primers and probe (P-BVDV-224) were further lengthened to increase their melting temperatures. Details of primers and probe are shown in Table 2. The primers and probes were synthesized by Generi Biotech s. r. o., Hradec Králové, Czech Republic.

Synthesis of RNA standard for RT-qPCR

A 160 bp fragment of 5’UTR of BVDV 1 (CZ3192) amplified by RT-PCR using primers 183-F and 343-R was separated on 1.5% agarose gel and purified using a Gel Extraction kit (Qiagen, Hilden, Germany). Purified cDNA was cloned into pGEM (Promega, Madison, USA). Plasmid was linearized and purified using QIAquick PCR purification kit (Qiagen, Hilden, Germany). Purified linearized plasmid served as a template for in vitro transcription by T7 RNA polymerase (Promega, Madison, USA). RNA transcript was treated with RQ1 RNase-Free DNase (Promega, Madison, USA) to remove plasmid DNA and purified by RNeasy kit (Qiagen, Hilden, Germany). The RNA transcript was dissolved in nuclease-free water and quantified using NanoDrop 2000 spectrophotometer and converted into copy number. RT-qPCR was performed with 10-fold serial dilution (10^1–10^8 RNA copies per reaction).

RT-qPCR for BVDV quantification

The reverse transcription and PCR was performed in a single tube using QuantiTect Probe RT-PCR kit (Qiagen, Hilden, Germany). Reaction mix recipes were prepared according to the manufacturer’s instructions with a final reaction volume of 10 µl. The reaction mixture consisted of: 5 µl of 2 × QuantiTect Probe RT-PCR Master Mix,
1 µM forward primer 183-F, 1 µM reverse primer 343-R, 0.2 µM probe P-BVDV-224, 0.1 µl of QuantiTect RT Mix and 2 µl of template. The assay was carried out in white LightCycler 480 Multiwell Plates 96 (Roche, Mannheim, Germany) and was run in LightCycler 480 Instrument (Roche, Mannheim, Germany). The thermocycling consisted of reverse transcription at 50 °C for 20 min, PCR initial activation step at 95 °C for 15 min, followed by 45 two-step cycles of denaturation at 95 °C for 30 s and combined annealing/extension at 60 °C for 60 s. Fluorescence signals were measured during the elongation step.

**CCoV reference and CCoV RT-qPCR**

The specific amplification and detection of CCoV RNA was performed in a separate RT-qPCR reaction. The RT-qPCR mastermix and thermocycling conditions were the same as for BVDV RT-qPCR. The primers (forward CCoV N724-F and reverse CCoV N871-R) and probe (P-CCoV N816) were selected from the nucleoprotein gene of CCoV 1-71 sequence AB105373 obtained from the GenBank. Details of primers and probes are listed in Table 2.

**Sequencing**

Primers BD1 and BD3 (Vilcek et al. 2001) were used to partially sequence the Npro protein gene of BVD viruses from infected animals 34, 35, 36, 37 and 38. Viral RNA was transcribed and amplified by a Transcriptor One-Step RT-PCR kit (Roche, Mannheim, Germany) according to the manufacturer’s instructions. Fifty µl of reaction mixture consisted of 10 µl of 5 × Reaction Buffer, 0.4 µM of both forward and reverse primers, 1µl of Transcriptor Enzyme mix and 5 µl of template. The resulting PCR products were gel purified using a Gel Extraction kit (Qiagen, Hilden, Germany) and afterwards sent to Eurofins Genomics (Vienna, Austria) for sequencing.

**Results**

**Linearity and limit of detection of BVDV RT-qPCR**

A standard curve was established using standard BVDV RNA that was 10-fold serially diluted from $1 \times 10^{10}$ to $1 \times 10^{2}$ copies per reaction and amplified in duplicate. The standard curve showed an efficiency of 1.91, a correlation coefficient of 0.999, a slope of -3.56, and an intercept of 42.23. The assay was linear over a range of $1 \times 10^{2}$ to $1 \times 10^{10}$ RNA copies and detected at least 100 RNA copies.

**RT-qPCR normalisation**

Tenfold serial dilutions of CCoV viral stock ($9.5 \times 10^{5}$ TCID50) were extracted and tested in duplicate in CCoV RT-qPCR. The concentration of 9.5 TCID50 (quantification cycle (Cq) value above 30) was chosen to normalize BVDV RT-qPCR. To evaluate precision and reproducibility of the assay, 80 spiked serum samples were amplified and the mean Cq and SD obtained were 31.67 and 0.4, respectively.

### Table 2. Nucleotide sequences and location of the polymerase chain reaction primers and hydrolysis probes used in this study.

<table>
<thead>
<tr>
<th>Primer name</th>
<th>Sequence (5’-3’)</th>
<th>Genome position</th>
<th>Amplicon length</th>
</tr>
</thead>
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<tr>
<td>BVDV 183-F</td>
<td>ACAGGGNAGTCGTCARTGGTTC</td>
<td>183-204&lt;sup&gt;a&lt;/sup&gt;</td>
<td>160 bp</td>
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<tr>
<td>BVDV 343-R</td>
<td>CTCTGCWRCACCCTATCAGGCTGT</td>
<td>320-343&lt;sup&gt;a&lt;/sup&gt;</td>
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<tr>
<td>CCoV N724-F</td>
<td>CACACCTGGAAGAGAACTGCAG</td>
<td>724-745&lt;sup&gt;b&lt;/sup&gt;</td>
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<tr>
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<td>849-871&lt;sup&gt;b&lt;/sup&gt;</td>
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<td>P-BVDV 224</td>
<td>FAM-CTCGAGATGCCAYGTGGAGGAGGTG-5’-BHQ1</td>
<td>224-254&lt;sup&gt;a&lt;/sup&gt;</td>
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<tr>
<td>P-CCoV N816</td>
<td>FAM-TGCCAATGGKARCAGTGCGATCCATACCAA-BHQ1</td>
<td>816-846&lt;sup&gt;b&lt;/sup&gt;</td>
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</tbody>
</table>

<sup>a</sup> position relative to BVDV strain NADL (GenBank Accession number AJ133739); <sup>b</sup> position relative to CCoV isolate 1-71 (GenBank Accession number AB105373).
BVDV quantification

The concentration of BVDV RNA copies was determined only for samples in which Cq values for CCoV reference ranged between 31.67 ± 0.4. Samples in which CCoV Cq values were outside this range were retested and repeated; detection of Cq value above or below the threshold ranges excluded samples from quantification (n = 10). A standard curve was established for each RT-qPCR run and individual standard curves showed efficiency between 1.8 and 1.9 and regression coefficient above 1.99. RNA copy numbers from $2.2 \times 10^6$ to $7.4 \times 10^8$ per ml were detected in serum samples of 37 PI animals and from $6.6 \times 10^4$ to $3.3 \times 10^7$ in 3 acutely infected animals (Table 1).

Sequencing

Sequences obtained during the study were deposited in the GenBank under their accession numbers (MK410569-73). The 381 bp nucleotide long partial sequences of the Npro protein gene were identical except one synonymous substitution detected in the sequence of BVDV from PI animal 34 (data not shown).

Discussion

We used one-step RT-qPCR utilizing a hydrolysis probe to quantify BVDV RNA in serum samples. For monitoring sample processing and amplification efficiency, a known amount of reference virus was added to each serum sample at the beginning of the isolation procedure (Cleland et al. 1999; Niesters 2001; Castelain et al. 2004; Young et al. 2006; Ward et al. 2009). Whenever the efficiency of RT-qPCR is inhibited and/or RNA is lost during sample processing, the Cq of reference virus will be higher. Thus the presence of amplifiable RNA as well as of inhibitory substances in extracted material is simultaneously monitored by this method.

In our study, sera of young animals aged between 6–12 months were collected, each animal was sampled twice during the time period ranging from 21 days up to 2 months to confirm PI. A relatively stable value of BVDV concentration was detected among sera from first and second sampling. The viral load ranging from $2.2 \times 10^6$ to $7.4 \times 10^8$ RNA copy number per ml was detected in serum of PI animals by RT-qPCR. Serum load levels of $3.3 \times 10^7$, $1.8 \times 10^7$, and $6.6 \times 10^4$ RNA copy number of BVDV were detected in acutely infected animals 37, 38, and 39, respectively (Table 1). As the virus replicates to high titres in PI animals, it is generally supposed that PI animals have a higher concentration of BVDV in their sera than acutely infected animals have. Only the serum viral load of three TI animals was quantified in our study. These animals tested positive for viral RNA and antigen in the first sampling and were antibody positive three weeks apart, which confirmed transient infection. The PI animals (34, 35, and 36) housed together with TI animals in the same herd G had serum viral loads ranging from $6.6 \times 10^6$ to $1.3 \times 10^7$ and these values are comparable with the viral load of $3.3 \times 10^7$ and $1.8 \times 10^7$ of TI animals 37 and 38, respectively. This is in accordance with the results obtained by antigen ELISA where corrected optical density (OD) of TI and PI reached similar values (data not shown). On the contrary, in TI animal 39, the viral serum load of $6.6 \times 10^4$ was significantly lower and the corrected OD value in antigen ELISA was also lower, which together with no detection of antiviral antibodies suggests the beginning of infection. Hanon et al. (2012) tested whole blood samples with commercial real-time RT-PCR test and found that the mean Cq value for PI animals is much lower than for TI. Nevertheless, they also detected TI animals with the same or even lower Cq value than the Cq value of some PI animals was. They also detected TI animals which tested positive by antigen ELISA. These results agree with our findings that the serum viral load in acutely infected animals can reach the same value as in PI animals.
Table 1. Quantification of bovine viral diarrhoea virus ribonucleic acid copies in 1 millilitre of serum by reverse transcriptase quantitative polymerase chain reaction.

<table>
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<th>Herd</th>
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<th>1</th>
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<th>3</th>
<th>4</th>
<th>5</th>
<th>6</th>
<th>7</th>
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<th>9</th>
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<td>S2</td>
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<td>3.3 × 10^8</td>
<td>x</td>
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<td>1.4 × 10^7</td>
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<td>6.5 × 10^7</td>
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Animal No. = animal number; S1 = first sampling; S2 = second sampling taken at least 3 weeks apart from the first sampling; x = quantification cycle of canine coronavirus out of range of 31.67 ± 0.4; bovine viral diarrhoea virus ribonucleic acid copy number not quantified; ND = not done
To characterize BVD viruses in the infected PI and TI animals from herd G (animals 34, 35, 36, 37 and 38), we partially sequenced their Npro protein gene. The obtained 381 bp long nucleotide sequences were identical except for one synonymous substitution detected in the sequence of BVDV from PI animal 34, suggesting that TI and PI animals were infected with the same virus and our results are not affected by viral strain variations.

Our results further revealed that the variability of virus concentration among different herds was more considerable than among animals in one herd. There are herds with relatively higher (herd E, mean viral load level $1.5 \times 10^8$ RNA/ml) or lower (herd C, mean viral load $6.3 \times 10^6$ RNA/ml) serum viral loads. The variability of virus concentration in serum between individual PI animals in one herd mostly does not exceed one order of magnitude. To answer the question whether the variability of BVDV serum concentration revealed among PI animals is due to different viral strains infecting them, sequence analysis should be performed.

In summary, we detected that serum viral load in PI animals ($2.2 \times 10^6$ to $7.4 \times 10^8$ RNA copy number per ml) can reach the same value as in TI animals ($6.6 \times 10^4$; $1.8 \times 10^7$, and $3.3 \times 10^7$). Data of the present study revealed that it is not possible to use virus quantification in serum samples to distinguish between persistently and transiently infected animals and paired serum samples should always be investigated to confirm persistent infection.

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