Since January 2020 Elsevier has created a COVID-19 resource centre with free information in English and Mandarin on the novel coronavirus COVID-19. The COVID-19 resource centre is hosted on Elsevier Connect, the company’s public news and information website.

Elsevier hereby grants permission to make all its COVID-19-related research that is available on the COVID-19 resource centre - including this research content - immediately available in PubMed Central and other publicly funded repositories, such as the WHO COVID database with rights for unrestricted research re-use and analyses in any form or by any means with acknowledgement of the original source. These permissions are granted for free by Elsevier for as long as the COVID-19 resource centre remains active.
Detection of bovine coronavirus using a TaqMan-based real-time RT-PCR assay

Nicola Decaro *, Gabriella Elia, Marco Campolo, Costantina Desario, Viviana Mari, Arianna Radogna, Maria Loredana Colaianni, Francesco Cirone, Maria Tempesta, Canio Buonavoglia

Department of Public Health and Animal Sciences, Faculty of Veterinary Medicine of Bari, Strada per Casamassima Km 3, 70010 Valenzano, Bari, Italy

A R T I C L E   I N F O

Article history:
Received 10 March 2008
Received in revised form 8 May 2008
Accepted 14 May 2008
Available online 24 June 2008

Keywords:
Cattle
Bovine coronavirus
Diagnosis
Real-time RT-PCR

A B S T R A C T

A real-time reverse transcriptase-polymerase chain reaction (RT-PCR) for the detection of bovine coronavirus (BCoV) RNA in clinical samples is described. The assay is based on TaqMan technology, consisting of two primers and one probe labeled with the reporter dye 6-carboxyfluorescin that binds selectively to the transmembrane-protein gene of BCoV. The BCoV real-time RT-PCR assay was able to detect the tested BCoV and BCoV-like viruses (canine respiratory coronavirus and bubaline coronavirus), whereas other common viral pathogens of cattle were not recognised by the established oligonucleotide set, thus showing that the test was specific for bovine-like CoVs. The detection limit of the assay was 20 BCoV RNA copies (1-log higher with respect to traditional gel-based RT-PCR) and the reproducibility was satisfactory, thus allowing for a sensitive and accurate measurement of the viral RNA load in clinical samples. Two hundred and twenty clinical specimens (92 rectal, 82 nasal and 46 ocular swabs) were subjected to gel-based and real-time RT-PCR. By conventional amplification, 43 rectal, 54 nasal and 34 ocular samples tested positive, whereas the TaqMan assay was able to detect the BCoV nucleic acid in 49 rectal, 60 nasal and 37 ocular swabs. The rapidity and high throughput of the BCoV TaqMan assay makes this method a powerful tool for a sensitive and specific diagnosis of BCoV infection in cattle.

© 2008 Elsevier B.V. All rights reserved.

1. Introduction

Bovine coronavirus (BCoV) is a group 2 coronavirus (CoV) that is responsible for different clinical forms of disease in cattle, including enteritis in newborn calves, “winter dysentery” in adult cows, and respiratory signs in calves and cows. The same virus strain could also be responsible for the simultaneous appearance of enteric and respiratory disease in the same animals as well as in calves and cows (Decaro et al., 2008a,b). It is thought that BCoV is able to cross the species barrier very frequently, giving rise to bovine-like CoVs which can infect different mammals. The bovine origin of some group 2a CoVs has been suggested for viruses such as human coronavirus (HCoV) OC43 (Vijgen et al., 2005), porcine hemagglutinating encephalomyelitis virus (PHEV) (Vijgen et al., 2006), canine respiratory coronavirus (CRCoV) (Erles et al., 2003) and other ruminant CoVs (Hasoksuz et al., 2007; Jin et al., 2007; Decaro et al., 2008c).

Several tools are available for the traditional diagnosis of BCoV infection, including electron microscopy (EM) (Bulgin et al., 1989), virus isolation (VI) on human rectal tumor (HRT-18) and Madin Darby bovine kidney (MDBK) cells (Dea et al., 1980), hemagglutination (HA) (Sato et al., 1977), and enzyme-linked immunosorbent assay (ELISA) (Sato and Akashi, 1993; Silva et al., 1999). However, these techniques have been proven to be poorly sensitive and often inconclusive due to the presence of pleomorphic CoV-like particles in clinical samples, virus instability in the environment, time-consuming procedures, or presence of maternally derived antibodies in the faeces (Sato et al., 1977; Dea et al., 1980; Saif, 1990; Sato and Akashi, 1993). Because of its higher sensitivity and versatility, the detection of viral nucleic acid by polymerase chain reaction (PCR) has been established as the diagnostic gold standard for a number of infectious diseases. Several reverse transcriptase (RT)-PCR-based methods have been developed for the detection and identification of BCoV and BCoV-like RNA in faecal and respiratory specimens of cattle (Tsunemitsu et al., 1999; Cho et al., 2001; Erles et al., 2003; Takiuchi et al., 2006). However, none of those RT-PCR assays were designed to be quantitative. In addition, PCR assays are frequently exposed to risks of carryover contamination, especially when a high sample throughput is required.

In this paper, the development of a real-time RT-PCR assay based on TaqMan technology is reported for the rapid and sensitive diag-
nosis of BCoV infection and accurate quantification of BCoV nucleic acid in diagnostic samples.

2. Materials and methods

2.1. Oligonucleotide design and synthesis

The transmembrane (M) gene sequences of BCoV strains were retrieved from the GenBank database (http://www.ncbi.nlm.nih.gov/Genbank/index.html) and aligned using the BioEdit software package (Hall, 1999). The primers and TaqMan probe were designed using the Beacon Designer Software, Version 2.0 (Premier Biosoft International, Palo Alto, CA, USA) to amplify a conserved 85-bp fragment within the aligned M genes. The primers and probe were synthesized by MWG Biotech AG (Ebersberg, Germany). The TaqMan probe was labeled with 6-carboxyfluorescein (FAM) at the 5′ end and with a nonfluorescent quencher 1 (NFQ1) at the 3′ end. The position and sequence of the primers and probe used for the assay are reported in Table 1.

2.2. Standard RNA for absolute quantification

To obtain a standard for the TaqMan assay, a 919-bp RT-PCR product containing the full-length M gene of BCoV strain 339/06 (Decaro et al., 2008b) was amplified using primer pair 28380F/29298R (Decaro et al., 2008c) and the RT-PCR product was cloned into pCR®/T/T/NT-TOPO vector (Invitrogen Srl, Milan, Italy) and transcribed with RibomAX™ Large Scale RNA Production System-T7 (Promega Corporation, Madison, WI) from the T7 promoter, according to the manufacturer's guidelines. After DNase treatment to remove residues of plasmid DNA, the transcripts were purified using a commercial column (QiAamp® RNA Easy kit, Qiagen S.p.A., Milan, Italy) and quantified by spectrophotometric analysis. Ten-fold dilutions of the RNA transcript, representing 10⁶ to 10² copies RNA μl⁻¹ of template, was carried out in a mixed faecal/nasal swab suspension from a calf which tested negative for BCoV by HA and gel-based RT-PCR. Aliquots of each dilution were frozen at −70 °C prior to use.

2.3. Field samples collection and preparation

A total of 220 samples, including 92 rectal, 82 nasal and 46 ocular swabs collected from cattle with enteric and/or respiratory signs, were analysed. The faecal and nasal swabs were homogenised (10%, w/v) in Dulbecco’s minimal essential medium (D-MEM). Sample suspensions were clarified by a brief centrifugation at high speed in a micro-centrifuge and aliquots of 140 μl of the supernatants were used for RNA extraction.

Viral RNA was extracted from each sample suspension with QIAamp® Viral RNA Mini Kit (Qiagen S.p.A.) in accordance with the manufacturer’s protocol. Template RNAs were eluted in 50 μl of elution buffer water and stored at −70 °C prior to use.

2.4. Real-time RT-PCR

Duplicates of the standard dilutions and RNA templates were subjected simultaneously to reverse transcription with the GeneAmp® RNA PCR kit (Applied Biosystems, Applera Italia, Monza, Italy). One microlitre of each duplicate of the standard dilutions or template RNA was reverse transcribed in a 20-μl reaction volume containing PCR buffer 1× (KCI 50 mM, Tris–HCl 10 mM, pH 8.3), MgCl₂ 5 mM, 1 mM of each deoxynucleotide (dATP, dCTP, dGTP, dTTP), RNase Inhibitor 1 U, MulV reverse transcriptase 2.5 U, random hexamers 2.5 U. Synthesis of c-DNA was carried out at 42 °C for 30 min, followed by a denaturation step at 99 °C for 5 min.

Real-time PCR for the simultaneous detection and quantification of BCoV RNA was performed on a 7500 Real-time PCR System (Applied Biosystems) with iTaq™ Supermix added with ROX (Bio-Rad Laboratories Srl, Milan, Italy). The quantitative assay targeting the M gene was conducted in a 50 μl reaction mixture containing 25 μl of master mix, 600 nM of primers BCoV-F and BCoV-R, 200 nM of probe BCoV-Pb and 20 μl of c-DNA. The thermal profile consisted of activation of iTaq DNA polymerase at 95 °C for 10 min, followed by 45 cycles of denaturation at 95 °C for 15 s annealing for 30 s and extension at 60 °C for 1 min.

The increase in the fluorescent signal was registered during the extension step of the reaction and the data was analysed with the appropriate sequence detector software (7500 System Software v.1.3.1, Applied Biosystems).

2.5. Internal control

In order to verify the absence of RNA losses during the extraction step and the presence of RT-PCR inhibitors in the RNA templates, an internal control (IC), consisting of an RNA synthetic transcript containing the M gene of canine coronavirus (CCoV) type II (Decaro et al., 2005), was added to the lysis buffer (AVL Buffer, Qiagen S.p.A.) at a concentration of 10,000 RNA copies μl⁻¹ of buffer prior to nucleic acid extraction. The fixed amount of the IC added to each sample had been calculated to give a mean Ct value in a genotype-specific real-time RT-PCR assay (Decaro et al., 2005) of 34.18 with a S.D. of 0.65 as calculated by 50 separate runs. Samples in which the Ct value for the IC was >35.48 (average plus 2S.D.) were excluded from the analysis.

2.6. Evaluation of real-time RT-PCR performances

To exclude cross-reactivities between BCoV and other viral pathogens responsible for enteric and/or respiratory diseases of ruminants, the specificity of the assay was evaluated by testing
isolates of the following viruses: bovine rotaviruses, bovine viral diarrhea virus, bovine respiratory syncytial virus, bovine parainfluenza virus, and bovine herpesvirus types 1 and 4. The ability of the assay to detect the nucleic acid of bovine-like CoVs was assessed by testing the strictly related CRCoV (Decaro et al., 2007) and bubaline coronavirus (BuCoV) (Decaro et al., 2008c).

Faecal, nasal and ocular samples collected from 10 uninfected calves as well as sterile water were also included in the analysis as negative controls and blanks, respectively.

To evaluate the detection limit of the real-time PCR assay, 10-fold dilutions of a faecal sample containing 2 × 10^2 copies BCoV RNA μL^−1 were made in a mixed faecal/nasal sample homogenate from a BCoV-negative calf and tested subsequently.

Serial 10-fold dilutions of standard RNA which contained from 10^1 to 10^9 copies RNA transcripts and the corresponding Ct values were used to plot the standard curve for BCoV RNA absolute quantification.

Reproducibility of the assay was evaluated by testing repeatedly clinical samples containing BCoV RNA titres spanning the whole range covered by real-time RT-PCR, as described previously (Decaro et al., 2004, 2005; Elia et al., 2006). The intra-assay reproducibility was evaluated by testing the same samples 10 times in the same experiment, whereas the inter-assay reproducibility was assessed by testing the same samples in 10 independent experiments. CVs were calculated by dividing the standard deviation of each tested sample by its mean and multiplying that result by 100.

2.7. Gel-based RT-PCR

The detection of BCoV RNA in clinical samples and M-gene RNA transcript dilutions was carried out using SuperScript™ One-Step RT-PCR for Long Templates (Life Technologies, Invitrogen, Milan, Italy) and primers specific for the spike-protein gene that are able to detect bovine-like CoVs (Erles et al., 2003; Decaro et al., 2007, 2008c). The following thermal protocol was used: reverse transcription at 50 °C for 30 min, inactivation of Superscript II RT at 94 °C for 2 min, 45 cycles of 94 °C for 30 s, 55 °C for 30 s, 68 °C for 30 s, with a final extension at 68 °C for 10 min. The PCR products were detected by electrophoresis in 1.5% agarose gels and visualised under UV light after ethidium bromide staining. The position and sequence of the primers used for conventional amplification are reported in Table 1.

3. Results

3.1. Analytical performance of the BCoV real-time RT-PCR assay

The template controls and BCoV-negative specimens did not produce any detectable fluorescence signal. The other tested viral pathogens were not detected with the exception of the bovine-like CoVs CRCoV and BuCoV, confirming that the real-time RT-PCR assay is specific for BCoV and BCoV-like viruses.

The detection limit of the assay was assessed as 2 × 10^1 RNA copies μL^−1 of template, whereas gel-based PCR was able to detect up to 2 × 10^2 copies μL^−1 of template.

Ten-fold dilutions of standard RNA were used to construct a standard curve covering a linear range of nine orders of magnitude (from 10^1 to 10^9 copies of standard RNA) and linearity was observed over the entire quantification range (slope = −3.366).

To determine the reproducibility of the assay, intra-assay and inter-assay coefficients of variation (CVs) were calculated (Fig. 1). Intra-assay CVs ranged from 18.42% (samples containing 7 × 10^7 DNA copies) to 41.03% (2 × 10^2 DNA copies), while the inter-assay CVs ranged from 28.93% (3 × 10^3 DNA copies) to 48.57% (2 × 10^2 DNA copies).

3.2. Internal control detection

The IC was detected in all the examined samples, with Ct values below the threshold value of 35.48, thus ruling out relevant RNA losses during nucleic acid extraction or DNA polymerase inhibition during real-time PCR amplification.

3.3. Analysis of clinical samples

The results of the analysis of 220 clinical samples collected from cattle with enteric and/or respiratory disease are summarised in Fig. 2. By using gel-based RT-PCR, 43/92 rectal, 54/82 nasal and 34/46 ocular swabs were found to contain BCoV RNA. Conversely, 49 rectal, 60 nasal and 37 ocular swabs tested positive for BCoV by real-time RT-PCR. Only one faecal sample tested positive by conventional amplification and negative by real-time analysis, whereas seven rectal, six nasal and three ocular swabs were found to be positive by the TaqMan assay and negative by conventional amplification. The number of samples in agreement between the two tests were 84 for the faecal swabs (42 positive and 42 negative samples), 76 for the respiratory specimens (54 positive and 22 negative samples) and 43 for the ocular swabs (34 positive and 9 negative samples). Totally, 203 samples were in agreement by both tests (130 positive and 73 negative samples). The analysed samples contained a wide range of BCoV RNA amounts per μL of template, from 2.20 × 10^1 to 5.53 × 10^9 (rectal swabs), from 3.15 × 10^1 to 8.09 × 10^8 (nasal swabs), and from 3.87 × 10^1 to 8.04 × 10^6 (ocular swabs).

4. Discussion

A real-time RT-PCR assay for the rapid and sensitive detection of the BCoV RNA in clinical samples of cattle was established. The assay was shown to be reproducible and linear over a range of 9 orders of magnitude, from 10^1 to 10^9 RNA copies, thus ensuring an accurate measurement of BCoV RNA loads in clinical samples. When compared with a classical RT-PCR protocol, the processing time required by TaqMan RT-PCR is shorter, the contamination risks are lower because of the lack of post-PCR steps and the specificity is increased by the probe hybridisation. The assay was also able...
to detect the nucleic acid of bovine-like CoVs (CRCoV and BuCoV). Other BCoV-like CoVs, such as hCoV-OC43 and PHEV, that were not included in our analyses, should be tested in order to assess whether they can be identified by the established assay. Although those viruses are not available in the laboratory where the BCoV TaqMan assay was developed, analysis of the M-gene sequences of strains from different geographical areas led us to predict a probable detection of these bovine-like CoVs.

The BCoV TaqMan assay was found to be 1-log more sensitive than gel-based RT-PCR. In fact, when the clinical samples collected from affected cattle were processed, real-time RT-PCR was able to detect BCoV RNA in seven rectal, six nasal and three ocular swabs that tested negative by conventional amplification. In contrast, only one faecal sample gave a positive result by gel-based RT-PCR and tested negative by the TaqMan assay. Sequencing of the M gene of this BCoV revealed the presence of two mismatches in the probe-binding region that prevented the correct annealing of the TaqMan probe (data not shown). CoVs are thought to mutate several base substitutions per round of replication (Jarvis et al., 1989). The high strain variation, TaqMan oligonucleotides were designed in a very conserved region of the viral replicase gene. The SYBR Green assay was able to detect as few as 10 CoV RNA copies. However, unlike the BCoV assay, that assay was unable to detect specifically BCoV and BCoV-like viruses, so it would appear more useful for the detection of new, uncharacterised CoVs than for specific identification of bovine-like CoVs in clinical samples.

In conclusion, a real-time RT-PCR assay for the rapid and sensitive detection of BCoV RNA was developed. This assay can be used as a powerful tool for the simultaneous analysis of several samples (up to 96 samples in the same plate) in a short time. Considering the difficulties of BCoV isolation and titration on cell cultures, the established assay can be helpful in BCoV-pathogenesis studies and vaccine trials.

Acknowledgements

This work was supported by grants from University of Bari, Italy: project ex 60% 2007 “Messa a punto di un sistema real-time RT-PCR per la identificazione e la quantificazione dell’RNA del coronavirus bovino”. The authors are grateful to P.J. Collins (CIT Department of Biology, Cork, Ireland) for the English revision of the manuscript.

References

Bulgin, M.S., Ward, A.C., Barrett, D.P., Lane, V.M., 1989. Detection of rotavirus and coronavirus shedding in two beef cow herds in Idaho. Can. Vet. J. 30, 235–238.

Cho, K.O., Hasoksuz, M., Nielsen, P.R., Chang, K.O., Lathrop, S., Saif, L.J., 2001. Cross-protection studies between respiratory and calf diarrhea and winter dysentery coronavirus strains in calves and RT-PCR and nested PCR for their detection. Arch. Virol. 146, 2401–2419.

Dea, S., Roy, R.S., Begin, M.E., 1980. Bovine coronavirus isolation and cultivation in continuous cell lines. Am. J. Vet. Res. 41, 30–38.

Decaro, N., Pratelli, A., Campolo, M., Elia, G., Martella, V., Tempesta, M., Buonavoglia, C., 2004. Quantitation of canine coronavirus RNA in the faeces of dogs by TaqMan RT-PCR. J. Virol. Methods 119, 145–150.

Decaro, N., Martella, V., Ricci, D., Elia, G., Desario, C., Campolo, M., Cavaliere, N., Di Trani, L., Tempesta, M., Buonavoglia, C., 2005. Genotype-specific fluorescent RT-PCR assays for the detection and quantitation of canine coronavirus type I and type II RNA in faecal samples of dogs. J. Virol. Methods 130, 72–78.

Decaro, N., Desario, C., Elia, G., Mari, V., Lucente, M.S., Cordioli, P., Colaianni, M.L., Martella, V., Buonavoglia, C., 2007. Serological and molecular evidence that canine respiratory coronavirus is circulating in Italy. Vet. Microbiol. 121, 225–230.

Decaro, N., Campolo, M., Desario, C., Cirone, F., D’abramo, M., Lorusso, E., Greco, G., Mari, V., Colaianni, M.L., Elia, G., Martella, V., Buonavoglia, C., 2008a. Respiratory disease associated with bovine coronavirus infection in cattle herds in Southern Italy. J. Vet. Diagn. Invest. 20, 28–32.

Decaro, N., Mari, V., Desario, C., Campolo, M., Elia, G., Martella, V., Greco, G., Cirone, F., Colaianni, M.L., Cordioli, P., Buonavoglia, C., 2008b. Severe outbreak of bovine coronavirus infection in dairy cattle during the warmer season. Vet. Microbiol. 126, 30–39.

Decaro, N., Martella, V., Elia, G., Campolo, M., Mari, V., Desario, C., Lucente, M.S., Lorusso, A., Greco, G., Corrente, M., Tempesta, M., Buonavoglia, C., 2008c. Biological and genetic analysis of a bovine-like coronavirus isolated from water buffalo (Bubalus bubalis) calves. Virology 370, 213–222.

Elia, G., Decaro, N., Martella, V., Cirone, F., Lucente, M.S., Lorusso, E., Di Trani, L., Buonavoglia, C., 2006. Detection of canine distemper virus in dogs by real-time RT-PCR. J. Virol. Methods 136, 171–176.

Erles, K., Toomey, C., Brooks, H.W., Brownlie, J., 2003. Detection of a group 2 coronavirus in dogs with canine infectious respiratory disease. Virology 310, 216–223.

Escutenaire, S., Mohamed, N., Isaksen, M., Thorén, P., Klineborn, B., Belák, S., Berg, M., Blomberg, J., 2007. SYBR Green real-time reverse transcription-polymerase chain reaction assay for the generic detection of coronaviruses. Arch. Virol. 152, 41–58.

Hall, T.A., 1999. BioEdit: a user-friendly biological sequence alignment and analysis program for Windows 95/98/NT. Nucl. Acids Symp. Ser. 41, 95–98.

Hasoksuz, M., Alekseev, K., Vlasova, A., Zhang, X., Spira, D., Halpin, R., Wang, S., Ghebremedhin, E., Saif, L.J., 2007. Biological, antigenic, and full-length genomic characterization of a bovine-like coronavirus isolated from a giraffe. J. Virol. 81, 4981–4990.

Jarvis, T.C., Kirkegaard, K., 1991. The polymerase in its labyrinth: mechanisms and implications of RNA recombination. Trends Genet. 7, 186–191.

Jin, L., Cebra, C.K., Baker, R.J., Mattson, D.E., Cohen, S.A., Alvarado, D.E., Rohrmann, F., 2007. Analysis of the genome sequence of an alpaca coronavirus. Virology 365, 198–203.

**Fig. 2.** Comparison between gel-based and real-time RT-PCR assays carried out on bovine clinical samples. Numbers indicate the samples positive (+) or negative (−) for BCoV. Results according to both techniques are shown in bold.
Lai, M.M.C., Holmes, K.V., 2001. Coronaviridae: the viruses and their replication. In: Knipe, D.M., Howley, P.M. (Eds.), Fields Virology, 4th edition. Lippincott Williams and Wilkins, Philadelphia, PA, pp. 1163–1183.

Marsilio, F., Di Martino, B., Decaro, N., Buonavoglia, C., 2005. Nested PCR for the diagnosis of calcivirus infections in the cat. Vet. Microbiol. 105, 1–7.

Saif, L.J., 1990. A review of evidence implicating bovine coronavirus in the etiology of winter dysentery cows: an enigma resolved? Cornell Vet. 80, 303–311.

Sato, K., Inaba, Y., Kurogi, H., Takahashi, E., Satoda, K., Onori, T., Matsumoto, M., 1977. Hemagglutination by calf diarrhea coronavirus. Vet. Microbiol. 2, 83–87.

Sato, M., Akashi, H., 1993. Detection of bovine coronavirus by enzyme-linked immunosorbent assay using monoclonal antibodies. J. Vet. Med. Sci. 55, 771–774.

Silva, M.R., O’Reilly, K.L., Lin, X., Stine, L., Storz, J., 1999. Sensitivity comparison for detection of respiratory bovine coronaviruses in nasal samples from feedlot cattle by ELISA and isolation with the G clone of HRT-18 cells. J. Vet. Diagn. Invest. 11, 15–19.

Takiuchi, E., Stipp, D.T., Alfieri, A.F., Alfieri, A.A., 2006. Improved detection of bovine coronavirus N gene in faeces of calves infected naturally by a semi-nested PCR assay and an internal control. J. Virol. Methods 131, 148–154.

Tsunemitsu, H., Smith, D.R., Saif, L.J., 1999. Experimental inoculation of adult dairy cows with bovine coronavirus and detection of coronavirus in feces by RT-PCR. Arch. Virol. 144, 167–175.

Vijgen, L., Keyaerts, E., Moes, E., Theelen, I., Wollants, E., Lemey, P., Vandamme, A.M., Van Ranst, M., 2005. Complete genomic sequence of human coronavirus OC43: molecular clock analysis suggests a relatively recent zoonotic coronavirus transmission event. J. Virol. 79, 1595–1604.

Vijgen, L., Keyaerts, E., Lemey, P., Maes, P., Van Reeth, K., Nauwynck, H., Pensaat, M., Van Ranst, M., 2006. Evolutionary history of the closely related group 2 coronaviruses: porcine hemagglutinating encephalomyelitis virus, bovine coronavirus, and human coronavirus OC43. J. Virol. 80, 7270–7274.