Identification of the recently described new type of bovine papillomavirus (BPV-8) in a Brazilian beef cattle herd

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Bovine papillomavirus type 8 (BPV-8) was first detected and described in teat warts as well as in healthy teat skin from cattle raised in Japan. The entire viral genome was sequenced in 2007. Additionally, a variant of BPV-8, BPV-8-EB, was also identified from papillomatous lesions of a European bison in Slovakia. In Brazil, despite the relatively common occurrence of BPV infections, the identification and determination of viral types present in cattle is still sporadic. The aim of this study is to report the occurrence of the recently described BPV-8 in Brazil. The virus was identified in a skin warts obtained from a beef cattle herd located in Parana state, southern Brazil. The papilloma had a macular, non-verrucous gross aspect and was located on the dorsal thorax of a cow. Polymerase chain reaction (PCR) was performed using generic primers for partial amplification of L1 gene. The obtained amplicon (480bp) was cloned and two selected clones were sequenced. The nucleotide sequence was compared to existing papillomaviral genomic sequences, identifying the virus as BPV type 8. This study represents the first report of BPV-8 occurrence in Brazil, what suggests its presence among Brazilian cattle.

INDEX TERMS: Bovine, bovine papillomavirus type 8, BPV-8, cutaneous papillomatosis, molecular analysis.
INTRODUCTION

In Brazil, bovine papillomavirus (BPV) infections are endemic in beef and dairy cattle throughout the country. Despite the high frequency of BPV infection, identification of BPV types in Brazilian cattle herds is still sporadic. By using specific primers, BPV type 1 was described in skin warts, peripheral blood and plasma from cattle with cutaneous papillomatosis, while BPV type 2 was detected in whole blood and urinary bladder tumors from cattle with chronic enzootic haematuria and cutaneous papillomatosis (Santos et al. 1998, Freitas et al. 2003, Wosiacki et al. 2005, 2006).

The partial analysis of L1 gene, the most conserved gene within the papillomaviral genome, by polymerase chain reaction (PCR) assay with degenerate primers and sequencing of amplicons has allowed a rapid identification of unreported viral types in human and animal hosts. A new PV type can be established once its complete genome has been cloned and its L1 gene displays less than 90% identity with the closest known PV type. A subtype is defined when the L1 sequences show identities between 90 and 98% (Manos et al. 1989, Forslund et al. 1999, Antonsson & Hansson 2002, Ogawa et al. 2004).

While hundreds of human papillomavirus (HPV) types have been identified, only six BPV types had been characterized until the early 1980s (Jarret et al. 1984, Bernard 2005). However, recent studies employing PCR with generic primers FAP59/FAP64 in combination with cloning and sequencing, have described 15 putative new BPV types (Forslund et al. 1999, Antonsson & Hansson 2002, Ogawa et al. 2004). After characterization of their complete genome sequences, four of these Japanese isolates were recently recognized as new viral types (BPV-7, -8, -9, and -10) (Ogawa et al. 2007, Tomita et al. 2007, Hatama et al. 2008). In addition, four putative new BPV types have been identified in cutaneous lesions from cattle herds in Parana state, Brazil (Claus et al. 2008).

In 2007, the complete genome sequence of BPV type 8 (GenBank accession number: DQ098913) was determined. Based on the FAP59/FAP64 amplicon sequence, this virus was previously designated as a putative new BPV type, named BAPV2, by the same research group (Ogawa et al. 2004). BPV-8 was first detected from papilomas as well as healthy teat skin from cattle in Japan. In addition, the BPV-8-EB, a variant of BPV-8, was detected in papillomatous lesions of a European bison from Slovakia, demonstrating that this new BPV type was present, simultaneously, in Asia and Europe (Literák et al. 2006, Tomita et al. 2007).

The aim of the current study is to report the first identification of BPV-8 in Brazil, what suggests its occurrence throughout Brazilian cattle herds.

MATERIALS AND METHODS

Papilloma specimen

The papilloma specimen was collected from a cow in a beef cattle herd in the Parana state, southern Brazil. The macular, non-verrucous, cutaneous papilloma was located on the dorsal thorax of a cow. The skin wart was collected and the fragment of papilloma specimen was triturated in phosphate buffered saline solution (PBS pH 7.2). The suspension (10% w/v) was centrifuged for 15min at 3000 x g at 4°C. An aliquot (250µl) of the supernatant was treated with lysis buffer (10mM Tris; 1mM ethylenediamine tetra-acetic acid [EDTA]; 0.5% Nonidet P40; 1% SDS; 0.2mg/ml proteinase K). After homogenization, the sample was incubated at 56°C for 30min.

DNA extraction

A combination of phenol/chloroform/isoamyl alcohol and silica/guanidine isothiocyanate methods was performed for extraction of total DNA (Alfieri et al. 2006). Briefly, the fraction of supernatant was treated with an equal volume of phenol/chloroform/isoamyl alcohol (25:24:1), homogenized and heated at 56°C for 15min (Sambrook & Russell 2001). After centrifugation at 10,000 x g for 10min, the aqueous phase was mixed with silica/guanidine isothiocyanate (Boom et al. 1990). The DNA was eluted in 50 µl of ultrapure sterile water and kept at -20°C until use. An aliquot of ultrapure sterile water was included as negative control in the DNA extraction procedure.

PCR assay

The PCR assay was performed using the primers FAP59 (forward; 5’-TAACTGTGCGTCCCTATT-3’) and FAP64 (reverse; 5’-CCWATATCWVHCATITCICCATC-3’) (Forslund et al. 1999) with slight modifications (Claus et al. 2007). Reaction was carried out in a solution containing 5 µl of the extracted DNA and 45µl of PCR-mix consisting of 1µl (20 pmol) of each primer; 200 µM of each deoxyribonucleotide triphosphate (dNTP); 2.5 units of Platinum Taq DNA polymerase (Invitrogen Life Technologies, USA); 1x PCR buffer (20mM Tris-HCl pH 8.4, 50mM KCl); 1.5mM MgCl₂, and ultrapure sterile water, to a final volume of 50µl. Amplification was performed in a thermocycler, using the following amplification conditions: an initial step of 10min at 94°C followed by 40 cycles of 1min at 94°C; 1min at 50°C, and 1min at 72°C, and a final extension step of 10min at 72°C. The amplified product was analyzed by electrophoresis in a 2% agarose gel in TBE buffer pH 8.4, stained with ethidium bromide (0.5µg/ml), and visualized under UV light.

Cloning and identity analysis

The PCR amplicon was purified (PureLink Quick Gel Extraction Kit, Invitrogen Life Technologies, Carlsbad, CA, USA) from agarose gel and submitted to cloning (TOPO TA Cloning Kit for Sequencing, Invitrogen Life Technologies, Carlsbad, CA, USA). The insert from two selected clones was sequenced according to the manufacturer’s instructions (MegaBACE 1000/ Automated 96 Capillary DNA Sequencer, GE Healthcare, Little Chalfont, Bucks, UK). Sequencing was performed in both directions by using M13 forward and reverse primers (DYEnamic ET Terminator Cycle Sequencing Kit, Amersham Biosciences, Little Chalfont Buckinghamshire, UK).

Sequences were examined with the PHRED software for quality analysis of chromatogram readings. The sequences were accepted if their base quality was equal to or higher than 20. The consensus sequences were determined using the CAP3 software and sequence identity was verified with all sequences deposited in the GenBank using BLAST software (http://
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www.ncbi.nlm.nih.gov/BLAST). The alignment was obtained with the aid of the BioEdit software (Hall 1999).

Phylogenetic tree was obtained by the Neighbour-joining method with the Kimura two-parameter distance estimate (Kimura, 1980), using MEGA version 3.1 program (Kumar et al. 2004). Statistical analysis of phylogenetic tree was determined by bootstrap method on 1000 replicates.

RESULTS AND DISCUSSION

A product of approximately 480 bp was amplified from the cutaneous papilloma. The negative control for PCR amplification yielded no amplified product. The product was sequenced and a 429 bp nucleotide sequence (BPV-8: nt. 5.739-6.172) was deposited in GenBank (accession number EF635385 as BPV/BR-UEL1). In addition, by nucleotide sequence comparison, it was demonstrated that sequences from both clones were identical.

The sequence analysis of the BPV L1 fragment identified from the cutaneous wart (BPV/BR-UEL1) revealed a 100% identity with BPV-8 sequences from Asia and Europe strains (Fig.1). While BPV-8 was originally detected in both teat papillomas and healthy teat skin swabs in Japan (Ogawa et al. 2004), the BPV-8 Brazilian strain was identified in a different anatomical location (dorsal thorax). In agreement with the current findings, the variant of BPV-8, designated BPV-8-EB, was also identified in a non-teat papilloma in European bison (Literák et al. 2006, Tomita et al. 2007). These findings suggest that BPV-8 does not have a predilection for a specific anatomical location.

Degenerate PCR primers with a high degree of nucleotide identity with conserved L1 gene have enabled the detection of a broad range of HPV types in a single round PCR assay (Manos et al. 1989, Forslund et al. 1999). These primers have also been applied to papillomas from animal species and have allowed the identification of putative new PV types. The use of FAP59 and FAP64 primers has also demonstrated the existence of a great diversity of BPV types (Antonsson & Hansson 2002, Ogawa et al. 2004). Recently, by using the FAP primer pair, BPV types 1, 2, and 6 were identified in skin warts from cattle from Parana state (Claus et al. 2007). Besides, the detection of four putative new BPV types (BPV-BR/UELI to -5), in Brazil, consisted the first report of the presence of different BPV types in the American continent (Claus et al. 2008).

Papillomavirus genomes seem to be very stable genetically and the occurrence of genetic changes, such as mutation or recombination, is an uncommon event. In addition, it is recognized that the HPV genome evolved with similar frequency to their host genome (De Villiers et al. 2004, Bernard 2005). The identification of BPV-8 in Asia (Japan), Europe (Slovakia), and South America (Brazil), within a short period of time, suggests the previous existence of this BPV type in these regions. It is possible that the presence of BPV-8 in Brazil had been ignored to date in the face of the scarce studies concerning the characterization of BPVs in the country. The result of the current study suggests that the low diversity observed in BPV (BPV-1 to BPV-10) to date may be due to the low number of viruses examined rather than due to a low diversity in the BPV genome. In addition, the detection of this viral type in Brazil points to its possible distribution throughout Brazilian cattle herds.

The recent researches in BPV suggest that this virus can be as diverse as HPV, and emphasize the importance of new investigations involving the molecular epidemiology of BPV infections in beef and dairy cattle herds around the world.

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