Potential Pathogenetic Role of Bovine Herpesvirus 4 in Two Dairy Cows with Dermatitis-Pyrexia-Hemorrhagic Syndrome

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Dermatitis, pyrexia, and hemorrhagic syndrome (DPHS) is a rare bovine syndrome of unclear etiology. We describe two DPHS cases, the first to occur in Italy, with clinicopathological findings suggesting a potential pathogenetic role of bovine herpesvirus-4 (BoHV-4).

CASE REPORT

Two Italian Holstein-Friesian cows, 5 and 6 years old, from the same herd were referred to the Teaching Hospital of the Department of Veterinary Science of Turin University because of acute onset of pyrexia, pruritus, and skin lesions. One animal (cow A) was referred 1 week after the onset of clinical signs and displayed forced decubitus; the other animal (cow B) was referred 36 h after the onset of clinical signs. Both animals were anorectic, played forced decubitus; the other animal (cow B) was referred 36 A) was referred 1 week after the onset of clinical signs and dis-

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region of the herpesvirus DNA polymerase gene was obtained by pan-herpesvirus-nested PCR assay using degenerate consensus primers (1). Direct immunofluorescence was negative for BVD/MD, BRSV, ovine herpesvirus 2 (OHV-2), and blue tongue virus (BTV) but showed a signal for BoHV-4. Nested PCR gave amplicons of expected sizes (from 215 to 315 bp) that were directly sequenced on an ABI 3130 genetic analyzer (Life Technologies). BLAST analysis revealed maximum identity with published BoHV-4 sequences. In addition, electron microscopy showed herpesvirus-like particles in all the cell cultures inoculated with the lung and spleen samples. The numbers of virus particles varied depending on the sample, with a higher number found in the spleen samples.

Phylogenetic analysis was performed on the glycoprotein B (gB) and thymidine kinase (TK) genes of BoHV-4. Amplification was carried out on samples that were positive by pan-herpesvirus diagnostic PCR employing the nested-PCR protocols described by Wellenberg et al. (2) for gB and Verna et al. (3) for TK. PCR products of the expected sizes were obtained during the first amplification round for both gene targets and directly sequenced on an ABI 3130 genetic analyzer (Life Technologies). The sequences retrieved from all samples were identical and used to create a representative consensus sequence. The newly generated consensus and BoHV-4 sequences available in GenBank were aligned using BioEdit. Nucleotide substitution models were evaluated using jModelTest2 (4), and the best model was selected according to Akaike information criterion analysis. Molecular Evolutionary Genetics Analysis (MEGA; version 6) was used for calculating P distance matrices, and phylogeny inference was used according to the maximum-likelihood criterion. The nucleotide substitution model was set according to jModelTest2 output, and evolutionary distances for both data sets were calculated using the Jukes-Cantor model. The robustness of the hypothesis was tested in 1,000 non-parametric bootstrap analyses. Determination of sequence simi-
larity based on nucleotide homology revealed that the analyzed gene regions of the detected BoHV-4 isolate shared a high degree of nucleotide sequence identity with known BoHV-4 sequences, 97.9 to 100% (gB) and 96.3 to 100% (TK), except with four gB sequences detected in Turkey in 2009 to 2011, which showed much lower values (71.3%).

We detected no unique amino acid changes in our isolate for both the predicted partial gB and TK proteins in a comparison with sequences in GenBank. Phylogenetic trees based on the aligned nucleotide sequences of BoHV-4 gB and TK genes are shown in Fig. 3.

To the best of our knowledge, this is the first report of dermatitis, pyrexia, and hemorrhagic syndrome (DPHS) in Italy. The clinical signs and histopathological lesions in both cows resembled those previously described in dairy cows affected by the syndrome (5–10). Although its etiology is still unclear, citrinin mycotoxicosis (5, 11) or di-ureido isobutane (DUIB) poisoning (6) has been suggested as a potential cause. Toxic causes were excluded in the present cases. The clinicopathological findings suggested a potential pathogenetic role of BoHV-4. Indeed, published data, the complete blood cell count of cow B, electron microscopy, and viral isolation results from the lungs and spleens of both animals all suggested an active viral infection (12–15).

Moreover, vasculitis has been reported as a consequence of systemic herpesvirus infections caused by malignant catarrhal fever (MCF) (16) and BoHV-4 (17), and bovine endothelial cell cultures were found to be susceptible to BoHV-4 infection (18). Although the pathogenesis of vasculitis remains unsolved, it is
FIG 3 Phylogeny inferred by maximum-likelihood analysis obtained by alignment of 335 nucleotides covering the partial glycoprotein B (gB) gene region of BoHV-4 (A) and 200 nucleotides covering the partial thymidine kinase (TK) gene region of BoHV-4 (B). The phylogenetic tree includes the novel BoHV-4 isolate (TO-1, red rhombus) and sequences available in GenBank. Country of origin and collection date, when available, are indicated together with the accession number for each sequence. Bootstrap (1,000 replicates) values of >50 are shown at the internal nodes. The length of each pair of branches represents the distance between sequence pairs. The scale bar represents the percentage of nucleotide differences.
usually assumed that a small fraction of infected lymphocytes induces the proliferation and deregulation of uninfected cells with a nonspecific cytotoxic effect [19]. In the present cases, both electron microscopy and viral culture of skin samples were negative, consistent with previous observations in cases of cutaneous MCF, in which viral particle detection and viral antigen expression in skin samples may be limited [19]. Furthermore, BoHV-4 has been isolated from cattle with a variety of inflammatory lesions, such as mammary pustular dermatitis, metritis, pneumonia, diarrhea, respiratory infections, interdigital dermatitis, and vaginitis [20–23].

We carried out phylogenetic analyses on partial gB and TK nucleotide sequences of BoHV-4. The gB analysis (Fig. 3A) showed that the BoHV-4 sequence locates within a major cluster with other GenBank sequences from Belgium, Germany, Italy, Ireland, Brazil, and Turkey, confirming that this region of gB is quite conserved among BoHV-4 strains [24] and, therefore, not helpful in providing clues about their geographical origin. Of note, four Turkish isolates published in GenBank diverge from the major cluster into a separate branch, though they do not carry differences at the amino acid level. Phylogeny based on TK sequences revealed that the BoHV-4 sequence identified in this study clustered within genotype 1, without significant divergence from other known strains. Figure 3B illustrates the main clusters corresponding to the proposed genotypes, 1, 2, and 3, that include European (Mover-like, genotype 1) and American strains isolated from bison and cat, the African strain isolated from buffalo (DN599-like, genotype 2), and Argentinean and Brazilian strains (genotype 3) from bovine (3, 25).

The clinicopathological findings were suggestive of an active BoHV-4 infection; however, molecular characterization of other field isolates and broadening of the spectrum of BoHV-4 phylogenetic targets are needed to clarify BoHV-4’s role in the pathogenesis of DPHS syndrome.

Nucleotide sequence accession numbers. Sequences reported in this study were submitted to GenBank with accession numbers KT166423 and KT166424.

REFERENCES


