Histological and virological findings in severe meningoencephalitis associated with border disease virus in Alpine chamois (Rupicapra rupicapra rupicapra) in Aosta Valley, Italy

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Abstract
In 2015, a young female Alpine chamois (Rupicapra rupicapra rupicapra), originated from the Aosta Valley Region, Northernwestern Italy, was conferred to the National Reference Centre for Wild Animal Diseases for pathologic examinations. Histological analysis revealed a severe meningoencephalitis characterized by lymphocytic and plasmacellular infiltration, gliosis, perivascular cuffs, and leptomeningitis at the level of brain and brain stem. Laboratory investigations included polymerase chain reaction, sequencing and characterization by phylogenetic analysis, and evaluation of the internal ribosome entry site secondary structure in the 5’ untranslated region. These tests identified the pathological agent as border disease virus, a known health risk in domestic small ruminants. Genetic characteristics of the isolated strains, closely related to ovine and caprine strain sequences from neighboring regions of Piedmont, France, and Switzerland, suggested geographic segregation and micro-evolutive steps within the species.

Keywords: Alpine chamois, Border disease virus, Meningoencephalitis, Pestivirus, Rupicapra rupicapra rupicapra.

Introduction
In the northwestern Italian region of Aosta Valley, a regular and wide monitoring program is operational under the guidance and supervision of the National Reference Centre for Wild Animal Diseases (CeRMAS). In the framework of this activity, carcasses of wild ruminants are regularly conferred to CeRMAS for gross pathologic, histologic, bacteriologic, and virologic examinations, as part of the health surveillance of wild animals.

On July 8, 2015, a young Alpine chamois (Rupicapra rupicapra rupicapra) was submitted to CeRMAS. The animal was found near a road in the vicinity of the municipality of Antey Saint André in Valtournenche, an alpine area with an altitude of about 1,000 m above sea level, characterized by the concomitant presence of cattle, sheep, goats farms, and wild ruminant ungulates (ibex, chamois, and roe deer). Collected by personnel of the Forest Rangers and transported to the wild fauna rescue centre of Quart, the animal, showing depression, difficulty of movement, and loss of fear for humans, was admitted for suspected heat exhaustion and subsequently euthanized due to its critical condition. Considering that the Alpine chamois population mating season occurs between the beginning of November and mid-December with subsequent births from mid-May until mid-June, the age of the animal was estimated between 1 and 2 months.

In line with the overall presentation of the case, a new border disease virus (BDV) was detected in the spleen and lungs of the animal and proposed as BDV genotype 8 (Caruso et al., 2017). Despite the presence of a severe meningoencephalitis, testing was inconclusive at the brain level. In order to confirm the presence of a new cluster in the species and verify a link between BDV and the central nervous system (CNS) lesions, a further investigation was carried out in collaboration with the Office International des Epizooties reference centre for classical swine fever (CSF) and bovine viral diarrhea viruses (BVDV) at the Animal and Plant Health Agency (APHA), UK. The results not only allow us to corroborate the previous results but also allow us to conclude that the meningoencephalitis was likely associated with the BDV infection.

Case Details
Post-mortem examination
After recording morpho-biometric data (age, sex, and weight), gross pathology lesions were determined at post-mortem examination according to a standard protocol. The organs with the most significant lesions were subjected to histological, bacteriological, parasitological, and virological procedures to complement the patho-morphologic investigations.

Brain, brain stem, and lungs were fixed in 10% neutral buffered formalin, embedded in paraffin wax, and cut into 4 µm thickness sections using a microtome. The sections were placed on slides, deparaffinized in Bioclear (Gene Linx International, Inc., Dublin, OH), rehydrated in decreasing gradations of alcohol, washed,
and stained with hematoxylin and eosin followed by dehydration in increasing concentrations of alcohol, cleared in Bioclear (Gene Linx International), mounted with a coverslip using Eukitt mounting medium (Kindler GmbH, Freiburg, Germany), and analyzed under an Olympus BX 60 light microscope (Olympus Europa GmbH, Hamburg, Germany).

**Bacteriological and parasitological tests**

Bacteriological examination was performed by Blood and MacConkey Agar over 48 h at 37°C in an aerobic atmosphere. Colonies of interest, isolated from the lung, were characterized by morphology, catalase and oxidase test, and Gram staining and typing by APINE Biomerieux biochemical tests. A parasitological examination has been carried out from the intestine by fecal floatation by 100% zinc sulphate solution.

**RT-PCR and DNA sequencing**

Viral RNA was extracted from fixed brain tissue samples using the RNeasy FFPE Qiagen kit according to the manufacturer’s protocol. All samples were tested using a short target real-time RT-PCR capable of differentiating between BDV and BVDV (La Rocca and Sandvik, 2009). The 5'-UTR was amplified by reverse transcription using the Superscript® III reverse transcription kit (Invitrogen, UK) as per the manufacturer’s protocol, followed by PCR with primers V324 and V326 (Vilcek et al., 1994). Sequencing of the purified amplicons on both strands was performed applying the Big Dye cycle sequencing technology and using the automated ABI3730 DNA sequencer.

**Genetic typing**

A phylogenetic tree based on the 5'-UTR was constructed with Clustal X (Chenna et al., 2003) using the neighbor-joining method (Saitou and Nei, 1987) and visualization with Newick tree format option in Mega (version 7.0.26) (Kumar et al., 2016). Qualitative and quantitative evaluation of genomic sequence divergence, in terms of palindromic nucleotide base pairings variations, has been applied through the consideration of relevant secondary structure regions in the 5'-UTR of the viral RNA, the three variable regions, V1, V2, and V3 genomic sequences, according to the genotyping based on the palindromic nucleotide substitution (PNS) method (Giangaspero and Harasawa, 2007). Internal ribosome entry site (IRES) types were identified according to base-pair (bp) combinations at the level of low-variable positions and ranked with alphabetic nomenclature according to increasing divergence in the species. The classification among BDV strains according to the PNS analysis based on changes in the secondary structure was compared with those based on the primary structure of the 5'-UTR performed through sequence alignment and construction of phylogenetic trees.

**Histopathology**

Microscopy revealed a severe and diffuse non-suppurative lymphohistiocytic meningoencephalitis (Fig. 1A–D). Associated histological lesions observed at the level of CNS tissues were focal cortical hemorrhage (red cells in brain parenchyma), mild perivascular mononuclear cuffs, gliosis, neutrophilic aggregates, and leptomeningitis.

Lungs presented suppurative broncho-pneumonia characterized by bronco-alveolar spaces filled with neutrophils, foamy macrophages, fluid, and occasionally fibrin aggregates, interlobular edema, and hyperemia. No lesions were evident at the level of kidneys and liver.

Specimens of formalin-fixed and paraffin-embedded tissues submitted to IHC staining were all negative for pestiviruses (data not shown).

**Bacteriological and parasitological examination**

Additional tests for the screening of pathogens, other than pestiviruses, showed the presence of bacteria in the lungs. Translucent colonies, grown on Blood...
Agar, formed by capsulated Gram-negative cocco-bacilli, positive at catalase and oxidase tests, were characterized by biochemical identification as Mannheimia haemolytica. No pathogenic bacteria have been found in kidneys, liver, and intestine. The parasitological examination revealed a high level of infestation by gastro-intestinal strongili and coccidia.

**Molecular detection and analysis of BDV in brain sample**

Applying the BDV specific PCR (Strong et al., 2010), two strains of BDV were detected in the two areas of the Alpine chamois brain (mesencephalon and occipital cortex), as well as in the spinal cord and lungs. The 5’-UTR sequences of isolates Chamois-VdA-2 and Chamois-VdA-3 have been deposited in the DDBL/EMBL/GenBank DNA database under accession numbers MG725337 and MG725338, respectively. Strains Chamois-VdA-2 and Chamois-VdA-3 belonged to BDV species, clustered as genotype BDV-c, subgenotype 3. The 5’-UTR sequences were very similar to the isolate Italy-58987 (Caruso et al., 2017), detected in the spleen of the same animal. Based on secondary structure characterization, the main difference was evident at the level of the IRES Domain D (Deng and Brock, 1993), showing a longer V3 locus when compared to the secondary sequence of isolate Italy-58987. According to the sequencing chromatogram of the 5’-UTR from the brain sample of strains Chamois-VdA-2 and Chamois-VdA-3, thymine and guanine nucleotides have been obtained in position 169, respectively (Fig. 2), suggesting a quasispecies phenomenon. A phylogenetic tree based on the 5’-UTR is presented in Figure 3.

Both the strains resulted related to strains CH-BD3 and CH-BD4 (Peterhans et al., 2010; Casaubon et al., 2012), Italy-103761 (Peletto et al., 2016), reported from Swiss and Italian small ruminants, and clustered as subgenotypes BDV-c1 and BDV-c3. They were also related to the French ovine strains, reported from the region of Provence Alpes Côte d’Azur (PACA) (Dubois et al., 2008) and to the strain Rupi-05 (Martin et al., 2015), isolated in an Alpine chamois in the French Alps, all included in the subgenotype BDV-c2 (Table 1). Among BDV-c2 strains, Rupi-05 was the closest to BDV-c1 and BDV-c3.

Comparison of the classification among BDV strains according to the PNS analysis based on changes in the secondary structure with those based on the primary structure of the 5’-UTR performed through sequence alignment and construction of phylogenetic trees showed some differences. Apart from different numerical and alphabetic nomenclature applied for linear sequence analysis and for PNS method, respectively, discrepancies have been observed in relation to the allocation of strains into genotypes and subgenotypes. In particular, strains reported by PNS as
Table 1. BDV species genotype c strains (n 17) evaluated according to the palindromic nucleotide substitution (PNS) method at the 5‘ untranslated region of RNA. Genotype nomenclature is indicated according to PNS and primary sequence analysis (PSA).

| Genotype | PNS | PSA | Strain | Origin | Country | Collection date | Accession | Reference |
|----------|-----|-----|--------|--------|---------|-----------------|-----------|-----------|
| c1.1     | Switzerland | CH-BD3 | Sheep | Switzerland | 2006 | JQ994199 | Stalder et al. (unpublished) |
| c1.1     | Switzerland | CH-BD4 | Sheep | Switzerland | 2006 | JQ994200 | Stalder et al., unpublished |
| c1.2     | 8 | Italy-103761 | Goat | Italy | 2014 | KT072634 | Peletto et al. (2016) |
| c2       | 6 | 06-F-0299/357 | Sheep | France: PACA | 2006 | EF694000 | Dubois et al. (2008) |
| c2       | 6 | 06-F-0299/369 | Sheep | France: PACA | 2006 | EF694001 | Dubois et al. (2008) |
| c2       | 6 | 06-F-0299/420 | Sheep | France: PACA | 2006 | EF694002 | Dubois et al. (2008) |
| c2       | 6 | 06-F-0299/477 | Sheep | France: PACA | 2006 | EF694003 | Dubois et al. (2008) |
| c2       | 6 | 10F03356 | Sheep | France | 2010 | KC859384 | Martin et al. (2015) |
| c2       | 6 | 90-F-6335 | Sheep | France: PACA | 1990 | EF693990 | Dubois et al. (2008) |
| c2       | 6 | 91-F-7014 | Sheep | France: PACA | 1991 | EF693993 | Dubois et al. (2008) |
| c2       | 6 | 92-F-7119 | Sheep | France: PACA | 1992 | EF693994 | Dubois et al. (2008) |
| c2       | 6 | 94-F-7446/1 | Sheep | France: PACA | 1994 | EF693996 | Dubois et al. (2008) |
| c2       | 6 | 94-F-7446/2 | Sheep | France: PACA | 1994 | EF693997 | Dubois et al. (2008) |
| c2       | 6 | RUPI-05 | Alpine chamois | France: PACA | 2011 | KC859383 | Martin et al. (2015) |
| c3       | 8 | Italy-58987 | Alpine chamois | Italy | 2015 | KX573913 | Caruso et al. (2017) |
| c3       | Chamois-VdA-2 | Alpine chamois | Italy | 2015 | MG725337 | Present work |
| c3       | Chamois-VdA-3 | Alpine chamois | Italy | 2015 | MG725338 | Present work |

PACA: Provence Alpes Côte d’Azur; Host: Alpine chamois (Rupicapra rupicapra rupicapra); Goat (Capra hircus); Sheep (Ovis aries).

Discussion

The anamnestic report, gross pathology, and histological lesions suggested that the animal was exposed to a severe trauma (head wounds and brain hemorrhage), possibly a car accident, given the location where it was rescued, and in line with the reduced sensorial capacity caused by CNS lesions. However, car accidents occur easily when wild animals are affected by neurological disorders, including infections which change the natural behaviour and can cause a loss of shyness.
The features of meningoencephalitis, with a prevalent non-suppurative response, strongly suggested a viral infection (Storts, 1995). However, the presence of a virus in the brain had not been confirmed before (Caruso et al., 2017). BDV was detected here and may thus be associated with severe meningoencephalitis, considering that some of the microscopical lesions (such as cortical hemorrhages) may otherwise be the result of the trauma.

CNS tropism and the adverse impact of BDV during fetal development are well known in domestic small ruminants. BDV is responsible for congenital viral infection affecting sheep (Ovis aries), with abortions, barren ewes, stillbirths, and small weak lambs, which can show tremors, abnormal body conformation, and abnormal birth coat. The disease in goats (Capra aegagrus hircus) is rare and characterized by abortions (Nettleton et al., 1998). It is not known whether these conditions also occur in free-ranging chamois (Rupicapra rupicapra), considering the difficulty to study abortions and post-natal diseases in wildlife. However, despite CNS lesions are observable in chamois, the suspected etiopathological role of Pestivirus is not confirmed. In our knowledge, only one strain (AND-3, accession number HE615085) has been reported from the host brain of a Pyrenean chamois (Rupicapra pyrenaica) in Andorra in 2009. Clinical signs, gross and microscopical lesions observed in the Italian chamois here, were similar to those described in natural and experimental infections in Pyrenean chamois (Marco et al., 2007; Martin et al., 2013).

Broncho-pneumonia is frequently found in chamois (Citterio et al., 2003), but in our case, it may have been promoted as secondary infection by immunosuppressive effects conferred by and attributed to BDV infection: we observed the typical features of suppurative bronchopneumonia and isolated Mannhemia haemolytica.

Characteristic genetic cluster of BDV species was restricted in Italy related to other isolates from France and Switzerland. Secondary structure alignment and computing of divergence values by comparison with other Pestivirus sequences suggested relatedness with different strains defined as genotypes BDV-6 (Dubois et al., 2008; Martin et al., 2015), BDV-8 (Caruso et al., 2017), and BDV Switzerland (Peterhans et al., 2010; Casaubon et al., 2012). In this genetic group, PNS BDV-c clustering was related to geographic origin (Switzerland C1.1, France C.2, and Italy C1.2/C.3) and host (domestic C1.1/C1.2/C.2 or wild C.2/C.3). Strains defined as genotypes 6, 8, or BDV Switzerland have not been compared to each other for taxonomical definition by the authors. Therefore, it was not possible to consider sequence similarities that in the present study resulted according to secondary structure analysis and construction of a phylogenetic tree. Thus, this might be a further case of taxonomical clustering without complete consideration of deposited sequences and published different interpretations and nomenclatures provided by other authors.

Genetic relation among strains isolated in France, Italy, and Switzerland, belonging to genotype BDV-c, suggested geographic segregation and evolutionary dynamic of a specific subgroup in the BDV species, related also to exchanges between domestic and wild ruminants (Fig. 3). All these strains resulted in circulating in three restricted neighboring regions. In each region, strains showed distinctive genomic characteristic. Most of the sequences originated from sheep, chronologically, first reported between 1990 and 1994, all from the French southeastern region PACA. About a decade later, other closely related sequences have been reported from ovine in Switzerland, showing also recent circulation in cattle (Frei et al., 2014). Then, in 2014, the virus was detected in small ruminants from the Italian northwestern region of Piedmont. In the following year, Alpine chamois from Aosta valley has been shown to harbor a similar virus. Alpine chamois from France was also affected by a related virus earlier in 2011. Since virologic monitoring has been regularly performed, it is likely that these virus types were not circulating previously. A common geographic trait was the Alpine mountain chain, a natural barrier.
sus scrofa

Rupicapra rupicapra

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dominant Mycobacterium bovis or Corynebacterium pseudotuberculosis in Alpine chamois (Domenis et al., 2018), roe deer affected by Mycobacterium bovis (Balseiro et al., 2009; Domenis et al., 2010), or ibex affected by Mycoplasma mycoides subsp. capri LC serovar (MmeLC), causative agent of infectious agalactia (Giangaspero et al., 2010). Such health risks are different from those frequent in wild fauna as M. conjunctivae agent of infectious keratoconjunctivitis, a highly contagious ocular disease, reported in wild animals in the Alpine regions of Italy (Grattarola et al., 1999), less related to spillover from domestic animals. Monitoring and improved management of wild fauna are important to protect the unique and precious Alpine ecosystem in Europe. In the present study, according to secondary structure analysis, the BDV species resulted heterogeneous. Sequence characteristics of the genotype BDV-c genomic cluster of the BDV species, circulating in regions of France, Italy, and Switzerland, suggested geographic segregation. Determination of BDV species heterogeneity is important for diagnostic efficiency and prophylactic purposes, taking into account the adverse health and economic impact on small ruminant farming and potential negative impact on wild fauna.

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