Association of Maedi Visna virus with Brucella ovis infection in rams

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Maedi Visna Virus (MVV) is a lentivirus which affects sheep in several countries of Europe, Africa and Asia and in the USA. It is responsible for progressive inflammatory diseases of the lungs, mammary glands, joints and central nervous system (Brodie et al., 1998, Pepin et al., 1998). The major site of viral replication is represented by the monocyte-macrophage lineage, but specific tropisms for other cell types have been reported (Brodie et al., 1995). The virus is usually transmitted via respiratory aerosol and milk, but other excretion routes, including the venereal, cannot be excluded.

The shedding of lentiviruses in semen has been well documented in primates. In humans, it is considered one of the main routes of human immunodeficiency virus transmission (Krieger et al., 1991; Kurth et al., 1991). In rhesus macaques, Miller et al. (1994) detected the simian immunodeficiency virus (SIV) in epididymal macrophages and T-cells by means of immunohistochemistry (IHC) and in situ hybridisation (ISH).

With regard to MVV, its presence in the semen has been assessed in rams concurrently affected by Brucella ovis (De la Concha-Bermejillo et al., 1996). These data, obtained by virus isolation and liquid-phase procedures, have led to the speculation that the venereal transmission of MVV plays an important role in the epidemiology of this infection.

In this study, we have investigated the localisation of MVV in the epididymis of rams experimentally infected with B. ovis, in order to identify the involved cells in a tissue context, and to confirm the relevance of this concurrent infection in the transmission of the retroviral disease.

Materials and Methods

Experimental infection

Ten Massese breed rams, aged 4-5 months, seronegative for B. ovis and B. melitensis, were
experimentally infected by conjunctival instillation of a solution containing \textit{B. ovis} (strain BG1/94) at a titre of $2.4 \times 10^9$ UFC/mL (Farina et al., 1995). The animals received 0.25 mL of the inoculum in each eye. Blood samples were taken weekly from all the rams. Serological tests were performed according to the protocol reported by Cerri et al. (1999). ELISA tests for MVV were also carried out on all the subjects as previously described (Saman et al., 1999).

**Post-mortem exams**

The seropositive rams for \textit{B. ovis} were sacrificed at intervals of fifteen days starting from nine weeks after inoculation, and complete necropsies were carried out. Epididymal samples were subjected to bacteriological tests as previously described (Cerri et al., 1999). Tissue samples from epididymis and lungs from all the animals were fixed in 10% formalin and paraffin wax-embedded.

**Histopathology and Immunohistochemistry (IHC)**

Sections (5 \textmu m) prepared by routine methods, were examined after staining with haematoxylin and eosin. IHC was performed on the epididymal tissues to reveal the presence of \textit{B. ovis}. Selected sections were labelled immunohistochemically with a commercial avidin-biotin alkaline phosphatase complex kit (Vectastain ABC Kit; Vector Laboratories, Burlingame, CA, USA) after microwave pre-treatment (10 min at 650 W). Selected sections were incubated overnight at 4°C with a primary rabbit antiserum to \textit{B. ovis}, used at the predetermined optimal dilution of 1:500, raised at the Department of Animal Pathology, Pisa, Italy (Cerri et al., 1999). The specific sera were replaced with Tris-buffered saline (TBS) or pre-immunization sera as a control for non-specific reactions. As negative control, tissues from non-infected rams were used. The reactions were developed with alkaline-phosphatase. Harris’ hematoxylin was used as a counterstain.

Epididymal and lung samples from all the subjects were also subjected to IHC tests specific for the detection of MVV. A monoclonal antibody to the p28 core protein (VMRD, Pullman, WA, USA) was used at the pre-determined optimal dilution of 1:500. The reactions were carried out accordingly with a protocol previously reported (Renzoni et al., 1995), and diaminobenzidine (DAB) was used to develop the reactions.

**Liquid-phase PCR**

DNA was extracted from paraffin wax-embedded samples of lungs and epididymis by a lysis buffer method, as previously described (Wright and Manos, 1990). A nested PCR was performed to amplify a portion of the MVV proviral DNA Pol region.

In the first step, a couple of primers (sense 4231: 5’ ATAGTAAGTGCAATCAAGATGC3’; antisense 4448 5’ TCCCGATTGTCTTACTC3’) was used.
to amplify a 218 bp sequence. The PCR mixture contained 1 × PCR buffer 5 µL, 3.5 mM MgCl₂, 200 µM dNTP mix, 0.5 µM sense primer, 0.5 µM antisense primer, Taq 2 units, DNA template 1700 ng, and H₂O (up to a total volume of 50 µL each sample). The amplification process consisted of one cycle at 95°C for 5 min, followed by 30 cycles at 95°C for 1 min, 59°C for 1 min, 72°C for 1 min, with a final extension at 72°C for 2 min.

In the second step, a couple of primers (Sense 4257 5’ - TCATTGCATCTAGAA TTT G - 3’; antisense 4398 5’ - TGTAATCCACTTGCCAAT - 3’) was designed to target a 142 bp sequence internal to the previously amplified portion. The composition of PCR mixture was the same as used in the first step, and, as a template, 1.5 µL of the first PCR products were employed. The amplification was performed at the following conditions: 95°C for 5 min, followed by 35 cycles at 95°C for 1 min, 48°C for 1 min, 72°C for 1 min, with a final extension at 72°C for 2 min. The PCR products were then analysed by agarose gel electrophoresis. Reagent and negative (unaffected) controls were used to check the specificity of the reaction.

**In situ PCR (IS-PCR)**

All stages of the in situ protocol were performed with a capillary gap system (Microprobe; Biomeda, Foster City, CA, USA) with the exception of amplification and hybridization. Sections (5 mm) of epididymis and lung samples from all the subjects were mounted on special capillary gap slides, and pre-treatments, hybridization, and post-hybridization procedures were performed as previously reported (Sanna et al., 1999). The PCR was designed to amplify a 218 bp sequence of MVV proviral DNA Pol region; we used the same primers employed in the first step of liquid phase PCR, and the PCR mixture contained 3 mM MgCl₂, 200 mM dNTP mix, 10 × PCR buffer 5 ml, 0.2 mM sense primer, 0.2 mM antisense primer, Taq 1.25 units, and H₂O up to a total volume of 50 ml per slide. Coverslips were added and sealed with Hybaid Easy Seal Starter Kit (Hybaid Ltd., Waldegrave, Teddington, Middlesex, UK). The amplification process consisted of one cycle at 95°C for 5 min, followed by 30 cycles at 95°C for 1 min, 59°C for 1 min, with a final extension at 72°C for 2 min, and it was performed in an in situ thermal cycler (Hybaid Omnislide 2000). After completion of the PCR, the slides were re-transferred into the capillary slide-holder, washed three times (10 min each step) with phosphate-buffered saline (PBS), post-fixed for 2 min in 4% paraformaldehyde and re-washed with PBS for 1 min. Slides were then hybridized with an internal digoxygenin-labelled probe (5’ CATTGACGATGATGATT 3’), diluted in hybridization solution IV (Fluka Chemika-Biochemika, Buchs, Switzerland). Reagent and unaffected controls (lung samples negative for MVV, and rabbit kidneys) were used.
Results

Experimental infection and post-mortem findings
Between 2 and 5 weeks after infection, all the ten rams became seropositive for Brucella ovis, which was isolated from epididymis of two rams, and from the blood of all the animals with the exception of one subject. ELISA tests for MVV constantly gave negative results. At necropsy, no gross finding was detected.

Histopathology
The histopathological examination revealed B. ovis-related lesions only in the two rams in which mature spermatozoa were present in the lumen of the epididymis (Figure 1). These subjects showed hyperplasia of the epididymal epithelium, microabscesses in the ducts, and oedema, perivascular cuffing and infiltration of lymphocytes, plasmacells and neutrophils in the interstitium (Figure 1).
Six animals (including those presenting the epididymal changes), showed lung lesions which were indicative of a MVV infection. They consisted of a thickening of the alveolar walls and an interstitial chronic flogosis with typical lymphatic clusters scattered in the parenchyma.

**Immunohistochemistry**

IHC for *B. ovis* gave positive results in the two rams which had shown appreciable histological epididymal lesions. A clear labelling was observed in mononuclear and epithelial ductal cells (Figure 2). IHC for MVV revealed the presence of the p28 viral core protein in the lungs of the rams presenting MVV-related microscopical lesions. MVV was mainly associated with alveolar and interstitial macrophages, but it was also present inside the lymphatic clusters. In the epididymis context, positive results were obtained only in the two animals which had shown immunohistochemical positivity for *B. ovis*. The signals were associated with interstitial macrophages, and epithelial ductal elements (Figure 3). No result was obtained from negative and reagent controls.
**Liquid phase-PCR**

DNA extracted from paraffin wax blocks was subjected to PCR amplification of MVV proviral DNA. A product of the expected size (142 bp) was observed in the samples which had shown immunohistochemical positivity for p28 MVV protein. No product was obtained from lung and epididymal samples from the other animals and from a healthy sheep lung kept as a negative control throughout the procedure (Figure 4).

**In situ-PCR**

The *in situ* PCR disclosed in the two subjects immunohistochemically positive for *B. ovis*, a number of cells that contained a dark blue pigment, the signal for amplified MVV proviral DNA. The positive areas corresponded to interstitial macrophages located near the ducts, and elements scattered in the epithelium (Figure 5). The unaffected and reagent control samples constantly gave negative results (Figure 6).

**Discussion**

This work was initially conducted to study the lesions caused by *B. ovis* and the pathogen distribution following experimental infection. As we were carrying out extensive studies on MVV infection, lung samples were also taken from the rams subjected to the experimental infection, and IHC, specific for MVV p28 protein was performed. Surprisingly, we were able to assess that some animals were immunohistochemically positive for MVV. The presence of the lentivirus was then confirmed by liquid phase PCR tests. Such observations appear to be in contrast with the results of the ELISA tests for MVV, but this could be compatible with previous findings of seronegativity that usually follows the colostral immunity received by the lambs (Brodie et al., 1995).

In the epididymis positive for *B. ovis*, prominent lesions were observed, with an involvement of both interstitial and ductal structures that substantially corresponded to those described in the literature (Ladds, 1993). IHC clearly showed that the bacterium was associated with the reported lesions. Interestingly, at the epididymal level, IHC for MVV gave good results only in the two rams which were positive for *B. ovis* by immunohistochemistry. The tissue distribution of the two pathogens matched well, as MVV p28 protein was found in interstitial macrophages at the inflamed sites and inside the affected epithelium. These observations were confirmed by IS-PCR, which disclosed positive signals arising from the same areas. It is remarkable that not all the subjects presenting lymphoproliferative lesions in the lung, whose IHC and liquid-phase PCR tests for MVV were positive, showed a similar labelling at the epididymal level.

Evaluated together, these data demonstrate that MVV may affect ovine epididymis, apparently taking advantage of the concurrent infection by *B. ovis*. As the ovine epididymis is not a usual target organ for MVV, its involvement may be facilitated in consequence of monocyte/macrophages recruitment and activation that occurs during the bacterial infection.

The presence of the lentivirus in an epithelial context, as demonstrated by both IHC and IS-PCR, is very interesting. MVV proviral DNA in cells other than macrophages had been described by Staskus et al. (1991), but Brodie et al. (1995) assessed that the virus could enter a variety of cell types, but productive infection was restricted to cells of macrophage lineage in affected tissues. Our IHC results show that the presence of MVV in the epididymal ductal cells is productive. Similar results were obtained by some of us on mammary glands subjected to experimental infection, and, also on that occasion, IHC and IS-PCR data matched very well (Vitali et al., 1997); on the other hand, we had demonstrated that caprine arthritis encephalitis virus, a lentivirus strictly related to MVV, shows a tropism for a broad cell spectrum; ependymal epithelia, for example, seemed to play a crucial role in the diffusion of the disease to different CNS districts via cerebrospinal fluid (Sanna et al., 1999). Recently, MVV proteins were found in the cytoplasm of ovine bronchiolo-alveolar elements (Gelmetti et al., 2000), dendritic cells (Ryan et al., 2000), and microglial cells (Ebrahimi et al., 2000). Thus, the infection of the epithelial cells in the epididymal context may be considered as not surprising at all.

A question to be answered is how the epithelial layers could become infected. Probably, the macrophages in the inflamed areas may act as a reservoir for viral infection, and facilitate the spreading of the virus towards the ductal structures. In any case, the presence of MVV-infected cells in the epididymal epithelia highlights the possibility that sperm of infected rams may contain
viral particles as a consequence of cell turnover. This possibility may per se increase the risk of disease transmission via semen.

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References