First molecular detection of *Mycoplasma ovis* (Hemotropic mycoplasmas) from Sheep in Brazil

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Abstract

*Mycoplasma ovis* is an emerging zoonotic pathogen with a worldwide distribution and can cause mild to severe hemolytic anemia, icterus, and poor weight gain in animals. Although *M. ovis* has been described in small ruminants worldwide, data on *M. ovis* in sheep in Brazil is unknown. The objective of the present study was to present the first report of hemotropic mycoplasma (HM) in sheep from Brazil. We evaluated factors associated with this infection, such as age group, tick presence, and anemia. Blood samples were collected from 33 sheep from a farm in southern Brazil and screened for hemoplasmas using PCR. Out of 33 samples, 26 (78.8%) tested positive for *M. ovis*. The sequencing of positive samples showed 100% identity with multiple *M. ovis* 16S rDNA sequences. No association was observed between the presence of *M. ovis* and the FAMACHA© score (*p = 0.620*). *Rhipicephalus (Boophilus) microplus* (15/33, 45.4%) was the tick species found on the animals. No significant association between *M. ovis* infection and presence of ticks (*p = 0.4134*) and age group (*p = 0.4221*) was observed. This is the first report of *M. ovis* infection in sheep from Brazil and only the second report of this pathogen in sheep in Latin America.

Keywords: Hemoplasmas, small ruminants, 16S rRNA gene, Rio Grande do Sul.

Resumo

*Mycoplasma ovis* é um patógeno zoonótico emergente com distribuição mundial e pode causar anemia hemolítica de leve a grave, icterícia e baixo ganho de peso em animais. Embora *M. ovis* tenha sido descrito em pequenos ruminantes em todo o mundo, os dados sobre *M. ovis* em ovinos no Brasil são desconhecidos. O objetivo deste estudo foi apresentar o primeiro relato de micoplasmas hemotrópicos em ovinos no Brasil. Avaliamos os fatores associados a essa infecção, como faixa etária, presença de carrapatos e anemia. Amostras de sangue foram coletadas de 33 ovelhas de uma fazenda no sul do Brasil e testadas para hemoplasmas usando a PCR. Das 33 amostras, 26 (78,8%) apresentaram resultado positivo. O sequenciamento das amostras positivas mostrou 100% de identidade com múltiplas sequências de *M. ovis* 16S rDNA. Não foi observada associação entre a presença de *M. ovis* e o escore FAMACHA© (*p = 0,620*). *Rhipicephalus (Boophilus) microplus* (15/33, 45,4%) foi a espécie de carrapato encontrada nos animais. Não houve associação significativa entre infecção por *M. ovis* e presença de carrapatos (*p = 0,4134*) e faixa etária (*p = 0,4221*). Este é o primeiro relato de infecção por *M. ovis* em ovinos no Brasil e o segundo relato deste patógeno em ovinos na América Latina.


Introduction

Hemotropic mycoplasmas (HM; formerly classified as *Eperythrozoon* & *Haemobartonella*) are small pleomorphic, uncultivable bacteria, which attach themselves to the surface of erythrocytes. They can cause hemolytic anemia in a wide variety of mammals including pigs, cattle, cats, dogs, deer, and humans (MESSICK, 2004; HOELZLE, 2008; GROEBEL et al., 2009; ZHOU et al., 2009; GRAZZIOTIN et al., 2011a; GIROTTO et al., 2012; MACHADO et al., 2017; SYKES et al., 2010; MAGGI et al., 2013).

The description of HM in small ruminants began with *Mycoplasma ovis* (formerly *Eperythrozoon ovis*) and “*Candidatus Mycoplasma haemovis*” (HORNOK et al., 2012). At present, it is not clear whether these hemoplasmas are single species or separate
strains/species remains, mainly because the complete genome sequence of \textit{M. ovis} strain Michigan contains two copies of the 16S rDNA genes, corresponding to the previously reported sequences for \textit{M. ovis} and \textit{C. M. haemovis} \cite{Deshuillers2014}.

Animal infections with HM are characterized by asymptomatic to subtle chronic anemia, and occasionally by an overt life-threatening hemolytic anemia. \cite{Messick2004, Neimark2004, Hoelzle2008}. \textit{Mycoplasma ovis} (formerly \textit{Eperythrozoon ovis}) transmission by hematophagous ticks, such as \textit{Rhipicephalus bursa}, has experimentally been demonstrated in Russia \cite{Nilkowski1969}. Since \textit{R. bursa} has never been found in the New World, it is still unknown which vector participates in the transmission of \textit{M. ovis} in our geographic location.

Among the routes of transmission of diverse HM species, we can comment the involvement of other arthropods, such as fleas, \textit{Ctenocephalides felis}, as a probable vector for \textit{M. haemofelis} and \textit{“Candidatus M. haemominutum”} \cite{Shaw2004, Woods2005, Lappin2006}, or the louse \textit{Polyplax serrata}, transmitting \textit{Mycoplasma coccoidees} to rats \cite{Berkenkamp1988}. The canine tick \textit{Rhipicephalus sanguineus} is the vector of \textit{Haemobartonella canis}, currently known as \textit{Mycoplasma haemocanis} \cite{Seneviratna1973}. There are also evidences of vertical transmission in cattle \cite{Girotto-Soares2015}. This scale serves as a tool in the selection of animals that are sensitive to anemias. The FAMACHA system is a tool developed to identify anemic sheep and goats by evaluating the ocular mucosa when compared to a standard chart \cite{VanWyk2002, Kenyon2012}. The FAMACHA score is directly related to anemia and indirectly with the packed cell volume (PCV). The scale ranges from 1 to 5, with 1 being red mucous and the animal being considered non-anemic (PCV > 27), and 5 being white mucous, suggesting severe anemia (PCV < 13) \cite{VanWyk2002, Kenyon2012, Maia2015}. This scale serves as a tool in the selection of animals that are sensitive to anemias. The FAMACHA system is in line with the recommendations of the Integrated Parasite (pest) Management (IPM) \cite{Nari2003, Kahn2012}.

To date, there are no studies in Brazil describing \textit{M. ovis} infections in sheep. The publications, so far, refer to \textit{M. ovis} infections in captive and free-living deer \cite{Grazziotin2011a, Grazziotin2011b}, one \textit{M. ovis} infection in goats \cite{Machado2017}, and there is a study in sheep from Argentina \cite{Aguirre2009}.

Although hemoplasmosis has been reported causing significant economic losses in sheep farming around the world, data for Brazilian herds remain scarce. In this context, the main aim of this study is to describe the first report of hemotrophic mycoplasmosis in sheep from Brazil. In addition, we evaluated factors associated with this infection in beef sheep on a farm in Rio Grande do Sul State in southern Brazil.

**Material and Methods**

**Study area and samples**

The samples for this study were obtained from a sheep beef production farm located in Cachoeira do Sul (29°53′54″S, 53°00′20″W), state of Rio Grande do Sul, southern Brazil, in April 2016. All the sheep in the farm, 33 animals, were collected for whole blood samples in EDTA-K$_3$, by jugular venipuncture. These sheep are routinely collected for sanitary analysis and the samples were given for research proposes. After the collection, the EDTA-K$_3$ tubes were refrigerated, transported to the laboratory, and immediately stored at -20 °C prior to DNA extraction and further processing and analysis. At the time of sampling, the animals had no obvious clinical signs compatible with hemoplasmosis. On the farm, the animals were kept in a pasture with a daily food supplementation; salt was offered \textit{ad libitum}. Cattle and horses used the same grazing area. The flock was composed of mixed breed sheep, with a predominance of Texel breed.

Epidemiological data collected from the farm included addressing age, presence of ticks, and FAMACHA$^0$ scoring, since it was not possible to evaluate the PCV from the samples. Sheep age was determined and later stratified into groups of < 1.5 years and > 1.5 years. Ticks were removed directly from the animals and kept in tubes with 70% ethanol for further classification according to morphological taxonomic keys \cite{Barros-Battesti2006}. The results for the FAMACHA$^0$ score were stratified for statistical purposes into groups of non-anemic (score ≤ 2) and anemic (score > 2) animals (Table 1).

**Table 1. Infection rate of hemoplasmas (Mycoplasma ovis) and result of the proportion test for the variable ticks, age group, Famacha$^0$ score in sheep in South Brazil.**

<table>
<thead>
<tr>
<th>Variables</th>
<th>Age group</th>
<th>Ticks</th>
<th>Stratified famacha score</th>
<th>Score famacha$^0$</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>&lt; 1.5 year</td>
<td>≥ 1.5 year</td>
<td></td>
<td></td>
</tr>
<tr>
<td>$GP_{pcr}$</td>
<td>10</td>
<td>16</td>
<td>13</td>
<td>23</td>
</tr>
<tr>
<td>$GN_{pcr}$</td>
<td>4</td>
<td>3</td>
<td>2</td>
<td>6</td>
</tr>
<tr>
<td>total</td>
<td>14</td>
<td>19</td>
<td>15</td>
<td>29</td>
</tr>
<tr>
<td>Infection rate (%)</td>
<td>71.4</td>
<td>84.2</td>
<td>72.2</td>
<td>86.6</td>
</tr>
<tr>
<td>P value</td>
<td>0.4221</td>
<td>0.4134</td>
<td>0.4221</td>
<td>0.620</td>
</tr>
</tbody>
</table>

$^*≤ 2$ not anemic; $^+> 2$ anemic; $^{**}GP_{pcr}$ group positive in PCR to \textit{M. ovis}; $^{**}GN_{pcr}$ group negative in PCR to \textit{M. ovis}; $^*${Score Famacha; 1 = red, non-anemic (>27 PCV); 2 = red-pink, non-anemic (23 to 27 PCV); 3 = pink, mildly-anemic (18 to 22 PCV); 4 = pink-white, anemic (13 to 17 PCV); 5 = white, severely anemic (<13 PCV).
DNA samples

Total Genomic DNA from 500 μL of blood was extracted using a standard phenol-chloroform protocol (SAMBROOK & RUSSEL, 2006) and quantified in a NanoDrop Lite Spectrophotometer (Thermo Scientific®); subsequently, the samples were stored at -20°C until PCR testing.

PCR amplification

All samples were tested by a conventional PCR protocol using the forward primer HBT F 5’ ATACGGCCCATATTCTACG 3’ and the reverse primer HBT R 5’TGCTCCACCACTTGTTCA 3’, targeting a 595- to 620-bp fragment of the hemotrophic mycoplasma 16S rRNA gene, as previously described by Criado-Fornelio et al. (2003).

Sequencing and phylogenetic analysis

The PCR products were submitted to electrophoresis through a 1.5% agarose gel and examined by a UV transilluminator. Amplicons of the expected size of three positive samples were purified with the PureLink kit (Invitrogen®) and sequenced in an automatic sequencer (Sanger) according to the manufacturer’s protocol. Generated sequences were submitted to BLAST® analysis (ALTSCHUL et al., 1990) to determine the closest similarities in GenBank®. Partial sequences of the 16S rRNA gene of hemotrophic mycoplasmas derived from sheep were aligned with corresponding 16S rRNA sequences of 20 Mycoplasma species retrieved from GenBank®, using Clustal/W v.1.8.1 (THOMPSON et al., 1994). A maximum likelihood phylogenetic tree using the GTR + G + I substitution model was generated using the Mega 7 software (KUMAR et al., 2016) with 100 bootstrap replicates. The substitution model was selected using the Mega 7 software (KUMAR et al., 2016), according to the lowest Bayesian Information Criterion (BIC) score. Sequence NR 113659 of Mycoplasma pneumoniae, a non-hemotrophic mycoplasma, was used as outgroup. An identity matrix calculated with the BioEdit software used sequences of M. ovis deposited in GenBank® and the ones found in this study to evaluate the similarity (Table 2).

Statistical Analysis

The variables age group, tick presence, and FAMACHA® score (considering anemic animals with a score greater than 2) were analyzed for the groups GPpcr (positive group in PCR) and GNpcr (negative group in PCR) by Fisher’s exact test. Differences in the distribution of the scores obtained in the FAMACHA® method for the GPpcr and GNpcr groups were analyzed through the Mann Whitney U Test of independent samples. The level of significance used as criteria value for rejection of the null hypothesis was 5% (p ≤ 0.05). Statistical analyses were conducted using IBM SPSS Statistics for Windows, version 22.0 (IBM Corp. Armonk, NY, 2013).

Results

Of the 33 animals, 26 (78.8%) were positive to hemoplasmas by PCR amplification of the 16S rRNA gene (Table 1). The 16S rRNA gene sequences of three M. ovis-like isolates had a 100% identity to M. ovis in sheep (KU983741 and KU983740) taken from GenBank®. Three isolates having 16S rRNA gene sequences were included in the phylogenetic tree (Figure 1). The DNA sequences generated in the present study have been deposited in GenBank® under the accession numbers MH379798 (616 pb), MH379799 (616 pb), and MH379800 (613 pb). The phylogenetic tree (Figure 1) depicts the three sequences of M. ovis generated in this study forming a cluster with previous M. ovis 16S rRNA sequences found at GenBank®.

We collected 68 ticks from 15 out of 33 sheep (45.4%). All ticks identified as R. (B.) microplus. Of these 15 animals, 13 were in the GPpcr (group positive PCR M. ovis). The characteristics of the ticks were as follows 11 males, 33 females, 1 nymphs, 1 larva; in the two remaining animals, part of GNpcr (group negative PCR M. ovis), 3 nymphs and 1 larva were collected (Table 1). Since the sheep in GPpcr were positive in blood when the ticks were removed, we did not perform PCR assays of the ectoparasites.

There were no statistical differences between the groups GPpcr and GNpcr in terms of the presence of ticks (p = 0.4134), age group (p = 0.4221), qualitative results of the stratified FAMACHA® score groups: non-anemic (score ≤ 2) and anemic (score > 2) (p = 0.4221), and the distribution of the FAMACHA® score (1 to 5) (p = 0.620) (Table 1).

Table 2. Identity matrix of Mycoplasma ovis sequence of the present study (1, 2 and 3) and isolates of Mycoplasma deposited in Genbank®.

<table>
<thead>
<tr>
<th>Sequences</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
<th>6</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. [MH379798] <em>Mycoplasma ovis</em> 1 Sheep</td>
<td>ID</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2. [MH379799] <em>M. ovis</em> 2 Sheep</td>
<td>1</td>
<td>ID</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3. [MH379800] <em>M. ovis</em> 3 Sheep</td>
<td>1</td>
<td>1</td>
<td>ID</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>4. [KU983740] <em>M. ovis</em> sheep</td>
<td>0.998</td>
<td>0.998</td>
<td>0.998</td>
<td>ID</td>
<td></td>
<td></td>
</tr>
<tr>
<td>5. [KF313922] <em>M. ovis</em> Human</td>
<td>0.998</td>
<td>0.998</td>
<td>0.998</td>
<td>1</td>
<td>ID</td>
<td></td>
</tr>
<tr>
<td>6. [HQ634377] <em>Mycoplasma spp. Ozotoceros bezoarticus</em></td>
<td>0.983</td>
<td>0.983</td>
<td>0.983</td>
<td>0.985</td>
<td>0.985</td>
<td>ID</td>
</tr>
</tbody>
</table>

ID: Identical.
Discussion

This is the first molecular detection of *M. ovis* in sheep in Brazil and only the second report of *M. ovis* in sheep from Latin America (AGUIRRE et al., 2009). Considering that *Haemonchus contortus* is an important cause of anemia in sheep (due to the consumption of whole blood) and severely impacts ovine breeding, any agent that causes hemolysis, such as *M. ovis*, gains special importance. There are distinct pathogenic processes that both contribute to the development of anemia in animals; however, hemoplasmosis remains largely neglected in small ruminant in Latin America. Given this, it is crucial to report the occurrence of this pathogen in Brazil.

Studies with experimental *M. ovis* (mentioned as *Eperythrozoon ovis*) infection in sheep have shown that the control group (no infection) had a mean of 10.35 g/dL Hb, with a PCV of 30.50%, while the infected group showed 6.45 g/dL Hb, with an average PCV of 18.17%. This suggests that *M. ovis* infection causes a large hematological alteration in sheep (DADDOW, 1979). Another study has evaluated the prevalence, diagnosis, and hematological changes that HM causes in sheep. Subclinical infection was associated with changes in mean corpuscular volume (MCV), mean corpuscular hemoglobin concentration (MCHC), red cell distribution width (RDW) and absolute monocyte count (HAMPEL et al., 2014). However, these changes were not evident in our study due to the evaluation method used, the FAMACHA© score, which measures PVC indirectly (see Table 1). Therefore, although there was no significant association between of the stratified FAMACHA© score groups: non-anemic (score ≤ 2) and anemic (score > 2) and the distribution of the FAMACHA© score (1 to 5) (Table 1) and presence *M. ovis* infection hematological alterations can be present in the studied herd, but were not detected.

The occurrence of *M. ovis* observed in this study (78.8%) was higher than that found for sheep in Hungary (51.5%) (HORNOK et al., 2009), goats in China (41%) (SONG et al., 2014), and goats in Brazil (39.30%) (MACHADO et al., 2017). In Argentina, a study described an outbreak of mycoplasmosis resulting in sheep death. Necropsy showed signs of anemia, mild subcutaneous jaundice, marked hepatic jaundice, splenomegaly, friable kidneys, congestion and pulmonary edema. The sheep were positive in the PCR for *M. ovis* (AGUIRRE et al., 2009).

We could not relate the involvement of a vector in the studied population, and therefore, there might be other transmission routes. Given the lack of information about the vectors in Brazil, other routes of transmission may be involved, such as other vectors, or iatrogenic and vertical transmission.

We found no significant association between the presence of HM and tick infestation, as observed in previous studies (RJEIBI et al., 2015). In a study conducted in northeastern Brazil no association was found between tick infestation and *M. ovis* infection in goats. The authors discuss that the lack of association may be linked to the control of tick used in the property and also to local factors of the region studied (MACHADO et al., 2017). Many of sheep were infested with ticks, which is not common for wool-covered sheep. There is a probability that this infestation was influenced by the unusually warm weather, favoring tick activity and reproduction. In addition, the cattle from the same property received strategical treatment for tick infestation, using acaricides with significant residue; we therefore speculate that this might have favored the infestation of *R. (B.) microplus* in the sampled sheep. In Mexico, *R. (B.) microplus* has been found infected with *M. ovis* present in sick sheep (MARTÍNEZ-HERNÁNDEZ et al., 2018). In the Old World, *R. bursa* is a known vector for *M. ovis* (NIKOL’SKII & SLIPCHENKO, 1969), and the lack of this species in the New World raises suspicion that *R. (B.) microplus* could be the vector, since it is the most common species infesting sheep in this locality and belongs to the same genus as the one from Russia.
It should be noted that the sampled animals were apparently healthy, and there was no statistically significant difference in distribution of FAMACHA® scores between GPpcr and GNpcr. This could be explained by the selection of resistant animals, eliminating from the herd those that presented signs of anemia, which has been practiced for more than 10 years on this property. Also, nutritional supplementation has a positive effect on the resistance of parasitized sheep (HAILE et al., 2004; LOUVANDINI et al., 2006; MELO et al., 2017).

The high occurrence of M. ovis in the studied sheep can be explained by the potential of hemotropic mycoplasmas to spread easily among animals due to the variety of transmission routes previously described, such as ticks, fleas, mosquitoes, and lice (DADDOW, 1981; SENEVIRATNA et al., 1973; BERKENKAMP & WESCOTT, 1988; SHAW et al., 2004; WOODS et al., 2005; LAPPIN et al., 2006; AKTAS & OZUBEK, 2017). Contamination via surgical procedures, such as vaccination, injectable anti-parasitic treatment, ear tagging, castration, tail docking, and mulesing, increases the risk of bloodborne transmission (CAMPBELL et al., 1971; PHILIBYE et al., 2006). Transplacental transmission also occurs, thus allowing a wide circulation of the pathogen not only in sheep, but also in other domestic and wild mammals, as already demonstrated in Brazil for free-living and captive deer (GRAZZIOTIN et al., 2011a; GRAZZIOTIN et al., 2011b). Transplacental transmission of “Candidatus Mycoplasma haemobos” between cows and their fetuses, both collected from an abattoir, has been reported in southern Brazil (GIROTTO-SOARES et al., 2016).

It is possible that mycoplasmas can act in synergy with other agents contributing to the worsening of infections and increasing lethality, especially in herds infected concomitantly with Haemonchus contortus. A synergic interaction between hemoplasmas and Anaplasma sp. was demonstrated in outbreaks that resulted in mortality of sheep (HORNOK et al., 2009; MARTÍNEZ-HERNÁNDEZ et al., 2018) and cattle (HORNOK et al., 2012). Hornok et al. (2009) have shown that 73% of the animals in their study had coinfections. Besides this synergic relation, hemotropic mycoplasmas can cause severe anemia (GENOVA et al., 2011) and even death in cattle (GLADDEN et al., 2016) as an isolated agent. Close and frequent human-animal contact, associated with the presence of arthropod vectors, may represent a risk for Public Health (MACHADO et al., 2017), considering that M. ovis has already been found infecting humans associated with other pathogens (SYKES et al., 2010; MAGGI et al., 2013).

Conclusions

In the New World, further studies evaluating the pathogenicity and transmission of sheep hemoplasmas are important. Large-scale studies are necessary to verify the occurrence in other regions of Brazil. Phylogenetic analyses showed that all three sequences of M. ovis generated at the present study formed a cluster with previous 16S rRNA of M. ovis sequences found at GenBank®.

Few data is available about M. ovis infections in Brazil, and the clinical and economic significance of hemoplasmas. The fact that we did not find statistical differences between groups GPpcr and GNpcr and the presence of ticks, age group, qualitative results of the FAMACHA® score may not show the reality of other herds.

We provide molecular evidence for the presence of hemotropic mycoplasma in sheep, representing the first report of M. ovis in sheep in Brazil, with a high occurrence of this organism in sheep from a farm in the south of Brazil. For sheep farmers, this should be a warning sign, given that this pathogen is an important cause of anemia and economic losses, whether it is the only infectious agent or simultaneously occurs with other diseases. In this sense, it is relevant to add M. ovis as a differential diagnosis for hemolytic disorders in Brazilian sheep.

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Detection of hemoplasmas in sheep from Brazil


