High occurrence of *Mycoplasma suis* infection in swine herds from non-technified farms in Mossoró, state of Rio Grande do Norte, Northeastern Brazil

Alta ocorrência de infecção por *Mycoplasma suis* em suínos de criações não tecnificadas de Mossoró, estado do Rio Grande do Norte, Nordeste brasileiro

Mariana Aparecida Toledo¹; Alexandro Iris Leite²; Luiz Ricardo Gonçalves¹; Keyla Carstens Marques de Sousa¹; Renan Bressianini do Amaral¹; Glaucenyra Cecília Pinheiro da Silva¹; Rosangela Zacarias Machado¹; Marcos Rogério André*¹

¹ Faculdade de Ciências Agrárias e Veterinárias, Universidade Estadual Paulista “Júlio de Mesquita Filho” – UNESP, Jaboticabal, SP, Brasil
² Departamento de Ciência Animal, Universidade Federal Rural do Semi-Árido – UFERSA, Mossoró, RN, Brasil

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Abstract

*Mycoplasma suis*, the etiological agent of swine hemoplasmosis, has been neglected in swine herds around the world. Swine hemoplasmosis is frequently associated with hemolytic anemia, disgalacty, infertility and immunosuppression, and it results in significant economic losses. This study investigates the occurrence of *M. suis* in non-technified swine herds in the northeastern region of Brazil using quantitative PCR (qPCR) based on the 16S rRNA gene. Between March and August 2013, blood samples from 147 swine were collected during slaughter in the city of Mossoró, state of Rio Grande do Norte, northeastern Brazil. One hundred and twelve samples (76.19%) were positive for *M. suis* by qPCR assays. The range of Cqs and quantification (copies of a *M. suis*-16S rRNA gene fragment/µL) was 20.86–37.89 and 1.64×10⁻¹–6.64×10⁷, respectively. One can conclude that *M. suis* infection have high occurrence (76.19%) in non-technified swine-rearing systems in Mossoró in the state of Rio Grande do Norte, Brazil.

Keywords: Swine hemoplasmosis, swines, hemotropic mycoplasmas, Brazil.

Resumo

*Mycoplasma suis*, agente etiológico da hemoplasmose suína, tem sido negligenciado nas criações de suínos ao redor do mundo. A hemoplasmose suína é frequentemente associada à anemia hemolítica, disgalactia, infertilidade e imunossupressão, acarretando em perdas econômicas. O objetivo do presente trabalho foi investigar, por meio da PCR quantitativa (qPCR) baseada no gene rRNA 16S, a ocorrência de *M. suis* em amostras de sangue de suínos de criações não tecnificadas na cidade de Mossoró, Estado do Rio Grande do Norte. Entre março a agosto de 2013, foram colhidas amostras de sangue de 147 suínos de criações não tecnificadas da referida região. Cento e doze amostras (76,19%) foram positivas para *M. suis* por qPCR. A média dos Cqs e da quantificação (número de cópias do gene 16S rRNA de *M. suis* por microlitro) foi de 20,86 – 37,89 e 1,64 x 10⁻¹–6,64 x 10⁷, respectivamente. Conclui-se que a infecção por *M. suis* apresenta alta ocorrência (76,19%) em criações de suínos não tecnificadas na cidade de Mossoró, estado do Rio Grande do Norte.

Palavras-chave: Hemoplasmose suína, suínos, micoplasmas hemotrópicos, Brasil.

Introduction

Knowledge about pathogens that can potentially affect animal welfare and cause economic losses is critical in swine production. Considering that Brazil is the fourth largest producer and exporter of pork meat, constant surveillance of infectious diseases is mandatory in order to maintain high levels of production (BRASIL, 2016). *Mycoplasma suis*, a bacterium belonging to the Mollicutes Order, is one of the pathogens that affects swine herds and may result in economic losses (HOELZLE, 2008). These bacteria (also known as hemoplasmas) are uncultivable wall-less organisms that parasitize the surface of erythrocytes, leading to structural deformations (HOELZLE, 2008). As a result, hemolytic anemia
might be noted in the acute phase of the disease (ZACHARY & SMITH, 1985).

Swine hemoplasmosis has worldwide geographic distribution (MESSICK, 2004) and is frequently associated with severe or moderate chronic hemolytic anemia, disgalacty, infertility and immunosuppression, and it results in economic losses (HOELZLE, 2008).

*M. suis* can be experimentally transmitted by several routes (HENRY, 1979; HENDERSON et al., 1997), and, possibly, by the louse *Haematopinus suis* (MESSICK, 2004), the stable fly *Stomoxys calcitrans*, and the mosquito *Aedes aegypti* (PRULAGE et al., 1993). Additionally, other routes of transmission of *M. suis* have been speculated since the agent has been detected in urine samples, in water used for drinking, vaginal and nasal secretions, and environmental dust samples (DIETZ et al., 2014).

Although laboratory diagnoses of *M. suis* infections are frequently based on observations of stained-blood smears, this technique lacks sensitivity and specificity. Molecular techniques, mainly qPCR, have been shown to be accurate and quick tests for the diagnosis of *M. suis*, which enable its detection and quantification (GUIMARÃES et al., 2011). Despite the apparent importance of *M. suis* to the swine industry worldwide, reports of this hemoplasma species in swine herds around the world have been rare. The pathogen has only been detected in pigs from China (YUAN et al., 2009; SONG et al., 2014), Germany (RITZMANN et al., 2009; HOELZLE et al., 2010) and Brazil (GUIMARÃES et al., 2007). The zoonotic potential of *M. suis* still needs further investigation since the agent has been molecularly detected in veterinarians and individuals who come into close contact with pig herds in Shanghai, China (YUAN et al., 2009). The goal of this study was to investigate the occurrence of *M. suis* in non-technified swine herds in the northeastern region of Brazil using molecular techniques.

**Materials and Methods**

**Sampling area, swine blood collection and DNA extraction**

Between March and August 2013, blood samples were collected by jugular vein puncture from 147 pigs, during slaughter. The pigs used in the present study were originally from non-technified swine herds in Mossoró, state of Rio Grande do Norte, Northeastern Brazil. All of the swine farmers have their rearing systems classified as subsistence without technification; the work involved family labor. The sampled herds did not contain infirmaries or quarantine rooms. The DNA was stored at -20°C until the time of cPCR and qPCR testing.

**Mycoplasma suis qPCR based on the 16S rRNA gene**

DNA samples exhibiting positive results in conventional PCR for the GAPDH gene underwent qPCR for *M. suis* based on 16S rRNA gene, using the primers F (5’-CCCTCATTTGACCTCAACTCAT-3’) and R (5’-CCAAGAGTTGTCACTGATGACC-3’), which flank a fragment of 437 base pairs (bp) of the GAPDH gene (BIRKENHEUER et al., 2003). Polymerase Chain Reaction was performed as previously described by Birkenheuer et al. (2003), with some modifications. The amplification reaction was performed using a final volume of 25 µL containing a mixture of 5 µL of the DNA sample, 0.2 mM of each deoxynucleotide, 1 µM of each primer, 0.75 mM MgCl₂, 0.25 µL Taq DNA polymerase, 10X PCR buffer and sterile ultra-pure water q.s.p 25 µL. The cycling conditions were: initial denaturation at 94°C for 3 minutes, 35 cycles consisting of denaturation at 94°C for 1 minute, annealing at 50°C for 2 minutes and extension at 72°C for 2 minutes, followed by final extension at 72°C for 7 minutes. The amplified products then underwent horizontal agarose gel electrophoresis stained with 1.5% ethidium bromide (0.5 µL/mL) in TEB (pH 8.0; Tris base 44.58 M; 0.44 M boric acid; 12.49 mM EDTA). The electrophoresis was performed at 90V/50mA for 90 minutes. A molecular weight marker of 100 bp (Thermo Fisher Scientific, Waltham, MA USA) was used to determine the length of amplified products. Amplicons were visualized and analyzed using an ultraviolet transilluminator (Chemidoc, Bio-Rad, Hercules, California, USA) coupled to a computational program of image analysis (Image Lab, Bio-Rad, Hercules, California, USA).

**Conventional PCR for the endogenous gene GAPDH**

In order to rule out the presence of inhibitors in extracted DNA samples and thereby avoid false-negative results in qPCR for *M. suis*, all of the DNA samples were submitted to conventional PCR (cPCR) using the primers GAPDH-F (5’-CCCTCATTTGACCTCAACTCAT-3’) and GAPDH-R (5’-CCAAGAGTTGTCACTGATGACC-3’), which flank a fragment of 437 base pairs (bp) of the GAPDH gene (BIRKENHEUER et al., 2003). Polymerase Chain Reaction was performed as previously described by Birkenheuer et al. (2003), with some modifications. The amplification reaction was performed using a final volume of 25 µL containing a mixture of 5 µL of the DNA sample, 0.2 mM of each deoxynucleotide, 1 µM of each primer, 0.75 mM MgCl₂, 0.25 µL Taq DNA polymerase, 10X PCR buffer and sterile ultra-pure water q.s.p 25 µL. The cycling conditions were: initial denaturation at 94°C for 10 minutes followed by 40 cycles at 95°C for 15 seconds, 58.5°C for 45 seconds and 72°C for 2 minutes, followed by final extension at 72°C for 10 minutes. The amplified products then underwent horizontal agarose gel electrophoresis stained with 1.5% ethidium bromide (0.5 µL/mL) in TEB (pH 8.0; Tris base 44.58 M; 0.44 M boric acid; 12.49 mM EDTA). The electrophoresis was performed at 90V/50mA for 90 minutes. A molecular weight marker of 100 bp (Thermo Fisher Scientific, Waltham, MA USA) was used to determine the length of amplified products. Amplicons were visualized and analyzed using an ultraviolet transilluminator (Chemidoc, Bio-Rad, Hercules, California, USA) coupled to a computational program of image analysis (Image Lab, Bio-Rad, Hercules, California, USA).
72°C for 30 seconds. The qPCR assays followed the Minimum Information for Publication of Quantitative Real-Time PCR Experiments (MIQE) (Bustin et al., 2009).

The sensitivity of the qPCR was determined using 10-fold serial dilutions of plasmid DNA. In the present study, the assay sensitivity was 10 copies/µL. In the number of plasmid copies was determined according to the formula: E = 10^−1/slope. To determine the limit of detection and quantification of the TaqMan assay, standard curves were generated by serial dilutions from 10^−5 to 10^0 copies of plasmids.

The amplification efficiency (E) was calculated from the slope of the standard curve in each run using the following formula: E = 10^(−1/slope). To determine the limit of detection and quantification of the TaqMan assay, standard curves were generated by serial dilutions from 10^−5 to 10^0 copies of plasmids.

### Results

Out of 147 GAPDH-positive samples, 112 (76.19%) were positive for *M. suis*. The samples were processed in five different plates with reactions efficiencies (E) of 95.1%, 91.9%, 93%, 98.1% and 97.5%, respectively (Table 1). The range of Cqs and quantification were 20.79-39.82 and 1.02 x 10^−5 - 5.62 x 10^2, respectively. Three samples demonstrated inconsistent quantification results and larger variation in the numerical value of Cqs and, consequently, in the number of *M. suis* 16S rRNA DNA copies. This finding was most likely due to the Monte Carlo effect (Bustin et al., 2009), which represents an inherent limitation of the technique, particularly in samples with low number of *M. suis* DNA. In the present study, the assay sensitivity was 10 copies/µL.

### Discussion

There are few studies reporting the occurrence of *M. suis* in swine herds around the world. The few reports that exist on this subject are from China for conventional PCR (Yuan et al., 2009), Brazil (Guimarães et al., 2007) and Germany (Ritzmann et al., 2009; Hoelzle et al., 2010) for qPCR. The occurrence of *M. suis* noted in this study was similar to that found in China (Yuan et al., 2009) but lower than that found in studies conducted in Germany (Ritzmann et al., 2009; Hoelzle et al., 2010). The pigs sampled in this study were from non-technified swine herds in northeastern Brazil. These herds lacked nutritional programs, adequate facilities, defined genetics, and standardized management techniques. Similarly, the animals sampled in Shanghai, China (Yuan et al., 2009) were from commercial swine farms that exhibited poor sanitary conditions and provided a favorable environment for the development of arthropod vectors, which may play an important role in *M. suis* transmission between pigs (Yuan et al., 2009). Besides reporting a *M. suis* occurrence rate of 86% in pigs, 32 out of 65 veterinarians and individuals who worked with the swine herds in Shanghai tested positive for this hemoplasma species.

Yuan et al. (2009) also pointed out that ingesting food contaminated with blood of infected pigs and the reuse of needles were risk factors associated with swine hemoplasmosis in China. Herein, the poor sanitary conditions of swine herds in Mossoró might be quite similar to those found in swine-rearing establishments in Shanghai, China. On the other hand, studies carried out in Germany revealed a lower rate of occurrence of *M. suis* (10.0-13.9%) (Ritzmann et al., 2009; Hoelzle et al., 2010) than that found in this study (76.19%). This difference may be mainly due to the technified systems of swine herds in Germany.

According to Song et al. (2014), poor sanitary conditions of swine herds and environmental conditions may favor the maintenance and spread of the pathogen. The garbage that had accumulated might have increased the presence of vectors such as flies and mosquitoes. Therefore, impaired management techniques lead to a lack of environment cleanliness and, consequently, *M. suis* transmission by fomites, such as needles and other materials. The conditions of feeders and drinkers may favor the contamination of food and water with blood, saliva, nasal secretions, dust, and the urine of infected animals, which favors the transmission of the agent (Dietz et al., 2014).

Our study was conducted in Mossoró, which has an average temperature of 27.4°C and a temperature range of 21-36°C during a year (Mossoró, 2008), which may help explain the high occurrence rate of *M. suis*. On the other hand, an occurrence rate of 33.1% and 18.2% for *M. suis* assessed via Southern blotting and cPCR, respectively, was found among sows sampled in herds of the state of Santa Catarina, Southern Brazil, where lower temperatures are reported (Guimarães et al., 2007).

Additional studies are necessary to verify the occurrence of *M. suis* in non-technified herds in other Brazilian regions. Furthermore, it will be important to analyze the risk factors associated with infection, positivity and susceptibility according to age groups. Investigations of the real impact of swine hemoplasmosis on swine herd productivity around the world are critically needed. Finally, studies should be conducted in order to estimate the correlation

### Table 1. Parameters of *M. suis*-qPCR assays.

<table>
<thead>
<tr>
<th>Plate</th>
<th>(E)</th>
<th>slope</th>
<th>r²</th>
<th>Cq values</th>
<th>Quantification of <em>M. suis</em>-16S rRNA gene (number of copies/µL)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>95.1%</td>
<td>−3.446</td>
<td>0.993</td>
<td>30.42 (24.34-38.52)</td>
<td>1.09 x 10⁴ (1.02 x 10⁴-2.07 x 10⁴)</td>
</tr>
<tr>
<td>2</td>
<td>91.9%</td>
<td>−3.531</td>
<td>0.996</td>
<td>31.02 (24.61-37.83)</td>
<td>1.20 x 10⁴ (1.18 x 10⁴-3.35 x 10⁴)</td>
</tr>
<tr>
<td>3</td>
<td>93%</td>
<td>−3.503</td>
<td>0.967</td>
<td>28.99 (24.85-39.07)</td>
<td>1.99 x 10⁴ (1.55 x 10⁴-3.85 x 10⁴)</td>
</tr>
<tr>
<td>4</td>
<td>98.1%</td>
<td>−3.368</td>
<td>0.983</td>
<td>30.41 (20.79-32.86)</td>
<td>2.79 x 10⁴ (1.46 x 10⁴-5.62 x 10⁴)</td>
</tr>
<tr>
<td>5</td>
<td>97.5%</td>
<td>−3.397</td>
<td>0.984</td>
<td>31.35 (25.21-39.82)</td>
<td>7.94 x 10⁴ (1.03 x 10⁴-1.90 x 10⁵)</td>
</tr>
</tbody>
</table>

(ranging from − to); r²: correlation coefficient.
between the quantification of parasitaemia and the occurrence of clinical signs and decreases in productivity indices.

Conclusion

*M. suis* has a high rate of occurrence (76.19%) in non-technified swine-rearing establishments in Mossoró in the state of Rio Grande do Norte, Brazil.

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