Outbreak of anaplasmosis associated with the presence of different *Anaplasma marginale* strains in dairy cattle in the states of São Paulo and Goiás, Brazil

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**Abstract**

The present study reports the genetic diversity of *Anaplasma marginale* during anaplasmosis outbreaks in rural properties of the states of São Paulo and Goiás, Brazil. Mortality rates of 3.5% (37/1,050) in calves, 4.7% (45/954) in heifers and 1.1% (25/2,200) in lactating cows were observed in a cattle herd of the municipality of Mambai, state of Goiás, central-western Brazil. In a cattle herd from the municipality of Lins, state of São Paulo, in southeastern Brazil, none of the animals died, despite presenting clinical signs suggestive of bovine anaplasmosis and exhibiting a drastic decrease in milk production. Thus, blood samples were collected from 100 animals with clinical signs suggestive of bovine anaplasmosis in the municipalities of Mambaí and Lins. Based on the microsatellite structure of the MSP1a of *A. marginale*, the genotypes E and H were observed in Lins, and the C, D and E genotypes were found in Mambaí. The analysis of the tandem repeat structures of the MSP1a showed nine different strains (α-10-15, α-β-3, α-β-13, α-β, 192, t-β-100, α-β-100, 193-β-100, 191-13-18, 191-13-18) in Lins and two (α-β-3-1-β and E-F-φ-3-4) in Mambaí. Three new tandem repeats of MSP1a (191, 192 and 193) were described. The α-10-15 and α-β-1-1-3 strains were predominantly associated with the occurrence of clinical anaplasmosis and mortality in calves, heifers and lactating cows.

**Keywords:** Anaplasmosis, Brazil, dairy herd, genotypes, MSP1a, outbreaks.

**Resumo**

O presente estudo relata a diversidade genética de *Anaplasma marginale* durante surtos de anaplasmose bovina no Brasil em propriedades localizadas nos Estados de Goiás e São Paulo. No rebanho bovino de Mambai, Estado de Goiás, Centro-oeste do Brasil, observaram-se taxas de mortalidade de 3,5% (37/1050) nos bezerros; 4,7% (45/954) nas novilhas e 1,1% (25/2200) nas vacas em lactação. No rebanho bovino de Lins, Estado de São Paulo, Sudeste do Brasil, embora os animais tenham apresentado sinais clínicos sugestivos de anaplasmose bovina, culminando em redução drástica da produção leiteira, nenhum animal veio a óbito. Assim, amostras de sangue de 100 animais com sinais clínicos sugestivos de bovina anaplasmosis foram coletadas em Mambaí-GO e Lins-SP. Baseando-se na estrutura do microssatélite da MSP1a de *A. marginale*, observou-se a presença dos genótipos E e H em Lins e C, D e E em Mambai. A análise da estrutura de repetição em tandem das MSP1a mostrou nove diferentes estirpes (α-10-15, α-β-3, α-β-13, α-β, 192, t-β-100, α-β-100, 193-β-100, 191-13-18) e duas (α-β-1-1-3 e E-F-φ-3-4) em Mambai. Três novos “tandem repeats” da MSP1a (191, 192 e 193) foram descritos. Foi observado predomínio dos estirpes α-10-15 e α-β-1-1-3 associado à ocorrência de anaplasmose clínica e mortalidade em bezerros, novilhas e vacas em lactação.

**Palavras-chave:** Anaplasmose, Brasil, rebanho leiteiro, genótipos, MSP1a, surtos.

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Introduction

Anaplasma marginale is a pathogen transmitted by ticks that belongs to the Anaplasmataceae family, order Rickettsiales (DUMLER et al., 2001). This rickettsia is a gram negative, obligate intracellular bacterium that can be biologically transmitted by ticks, can be mechanically transmitted by bloodsucking flies and fomites contaminated with infected blood, and, less commonly, can be transmitted via the placenta (KOCAN et al., 2010; AUBRY & GEALE, 2011).

Brazil is considered an endemic area for anaplasmosis in cattle (VIDOTTO et al., 2006; POHL et al., 2013), and the infection of cattle by multiple strains of A. marginale has recently been reported in the country (POHL et al., 2013; SILVA et al., 2015a, b). The global distribution and variation in pathogenicity of A. marginale are associated to the genetic diversity and variability of this bacterium (FUENTE et al., 2007). The genetic diversity of A. marginale strains is high in cattle in endemic regions worldwide (PALMER et al., 2001; FUENTE et al., 2007; CABEZAS-CRUZ et al., 2013; SILVA et al., 2015a).

The major surface proteins (MSPs) MSP5, MSP4 and MSP1a have been widely used for molecular characterization of A. marginale (AUBRY & GEALE, 2011). MSP1 is an adhesin important to bovine erythrocytes and tick cells. The protein contains tandem repeats that are involved in pathogen-host interaction, which provides relevant information about the A. marginale phenotypes transmitted by ticks (FUENTE et al., 2003a). Furthermore, analysis of the repeated sequences of MSP1 alpha has allowed the identification of A. marginale strains worldwide (CABEZAS-CRUZ et al., 2013). Despite the genetic diversity of MSP1a, this gene is considered a stable genetic marker, which is conserved in infections in cattle and ticks (PALMER et al., 2001; BOWIE et al., 2002; FUENTE et al., 2003b).

Among the different strains of A. marginale identified worldwide, some have been associated with the occurrence of anaplasmosis outbreaks. The α-β-τ and -10-15 strains have been previously described in outbreaks of bovine anaplasmosis in Mexico (ALMAZÁN et al., 2008) and Argentina (RUYBAL et al., 2009). In addition, the 72-62-61 strain has been reported as the most common in a cattle herd from the state of Minas Gerais, Brazil (POHL et al., 2013). In Brazil, significant variations in the MSP1a tandem repeats of strains of A. marginale have been found in the states of Paraná (VIDOTTO et al., 2006), Minas Gerais (FUENTE et al., 2004; POHL et al., 2013), Rio de Janeiro (SILVA et al., 2015a) and São Paulo (SILVA et al., 2015b). In MSP1a tandem repeats of strains of A. marginale described in Brazil, the most frequent sequences are 16, α, t in Paraná (VIDOTTO et al., 2006), 72 and α in Minas Gerais (POHL et al., 2013), 4 and t in Rio de Janeiro (SILVA et al., 2015a) and, α and β in São Paulo (SILVA et al., 2015b). The present study aimed to understand the genetic diversity of A. marginale during an outbreak of bovine anaplasmosis in rural properties located in the municipalities of Lins, state of São Paulo (southeastern), and Mambai, state of Goiás (central-western), Brazil.

Materials and Methods

Area and animals

A cross-sectional study was performed in rural properties of the municipalities of Lins (latitude: 21º 40’ 43” S, longitude: 49º 44’ 33” W and altitude: 437 m), state of São Paulo (southeastern) in October 2012, and Mambai (latitude: 14º 29’ 16” S, longitude: 46º 06’ 47” W and altitude: 709 m), state of Goiás (central-western), in December 2012, Brazil. The properties located in the municipality of Lins and Mambai had a cattle herd of 1,010 Holstein animals and 4,204 Holstein, Jersey and crossbred animals, respectively. Fifty animals with clinical signs suggestive of bovine anaplasmosis were evaluated on each property. The animals from the property located in Mambai were treated during the anaplasmosis outbreak with the following acaricides: Combo (cypermethrin + chlorpyrifos + piperonyl butoxide, Hertape, Brazil), Acatak (fluazuron, Novartis, Brazil) and Fluatac (fluazuron + abamectin, Ouro Fino, Brazil), according to the manufacturer’s recommendation.

Blood samples

Ethylenediamine tetraacetic acid (EDTA)-blood samples of 50 animals of each property were collected from the caudal or jugular veins of individual cattle. Serum samples were prepared from blood samples that were collected without EDTA, incubated at room temperature for 1 h and then centrifuged at 1000 x g for 15 minutes. Additionally, Giemsa-stained blood smears were prepared for microscopic examination. EDTA-blood and serum samples were stored at –20 °C. DNA was extracted from 200 µL of each of the 100 whole-blood samples using a QIAamp DNA Blood Mini kit (Qiagen, Madison, WI, USA) according to the manufacturer’s instructions.

Enzyme-linked immunosorbent assay (ELISA) and indirect fluorescent antibody test (IFAT)

ELISA and IFAT were performed as previously described by Machado et al. (1997) and Andrade et al. (2004). An A. marginale isolate from a calf in Jaboticabal, state of São Paulo, Brazil, was used to infect a calf for crude ELISA and IFAT antigen production (ANDRADE et al., 2004). For this purpose, a 3-month-old splenectomized calf was inoculated with 200 mL of A. marginale-infected blood (1.0 x 10^7 infected erythrocytes/mL). The rickettsia peak (1.0 x 10^7 A. marginale-infected erythrocytes/mL) was observed 7 days after the experimental infection. After the blood had been collected and processed for crude ELISA/IFAT antigen production (MACHADO et al., 1997), the experimentally infected animal was treated with oxytetracycline administered intramuscularly (20 mg/kg).

Polymerase chain reaction (Semi-Nested-PCR and quantitative real time PCR)

A total of 100 DNA samples were analyzed by quantitative real-time PCR (qPCR) reactions according to Carelli et al. (2007) for the msp1a gene. qPCR positive samples were additionally
submitted to a semi-Nested PCR (nPCR) targeting *msp1a* sequence (LEW et al., 2002). The nPCR reactions were performed using the primers 1733F (5'-TGTCCTGAGAAGAACCACCCTTGCTTACC-3') and 2957R (5'-AAACCTTGTAGCCCAACACCCTTGCTTACC-3'). Thus, the primers used were 1733F and 3134R in the first reaction and, 1733F and 2957R in the second reaction.

**Sequence of A. marginale msp1a microsatellite**

A microsatellite is located at the 5'-untranslated region (UTR) of the *msp1a* gene between the putative Shine-Dalgarno (GTAGG) sequence and the translation initiation codon (ATG) (Fuente et al., 2001). The microsatellite structure is GTAGG (G/ATTT)m (GT)n T ATG (ESTRADA-PEÑA et al., 2009). An analysis of the repeated sequences was performed according to the nomenclature proposed byFuente et al. (2007). The SD-ATG distance was calculated according to the formula:

\[
(4 \times m) + (2 \times n) + 1
\]

(1)

**Phylogenetic analysis**

For *msp1a* phylogenetic analysis, nucleotide sequences were aligned with MUSCLE (v3.7) configured for high precision (EDGAR, 2004) followed by removal of the ambiguous regions with Gblocks (v0.91b) (CASTRESANA, 2000). After alignment, regions with gaps were removed. Phylogenetic trees were reconstructed using the maximum likelihood (ML) and neighbor joining (NJ) methods as implemented in PhyML (v3.0 aLRT) (ANISIMOVA & GASCUEL, 2006) and PHYLIP (v3.66) (FELSENSTEIN, 1989), respectively. The reliability of the internal branches of the ML and NJ trees was assessed using the bootstrapping method (1,000 bootstrap replicates). Graphical representation and editing of the phylogenetic trees were performed with TreeDyn (v198.3) (CHEVENET et al., 2006). Detection of selection pressure on individual codons was calculated using two methods: single likelihood ancestor counting (SLAC) and fixed effects likelihood (FEL) implemented in the Datamon-key webserver (DELPORTE et al., 2010).

**Statistical analysis**

The frequencies of positive animals for *A. marginale* were compared via the Fisher's exact test, with 95% confidence level. Associations between the prevalence and its possible influencing determinants were measured by means of odds ratios (ORs). The operational procedures were performed with the aid of the statistical software R Foundation for Statistical Computing, version 2.12.2 (2011).

### Results

**Cattle mortality rate during the anaplasmosis outbreak**

An overall mortality of 2.5% (107/4,204) was observed in the cattle herd from the municipality of Mambai (state of Goiás) during the anaplasmosis outbreak (Figure 1), with 3.5% (37/1,050) of the calves, 4.7% (45/954) of the heifers and 1.1% (25/2,200) of the lactating cows dying. Animals presented fever, anemia and reduction in milk production. In contrast, in the property located in the municipality of Lins (state of São Paulo), none of the animals died, although they presented apparent clinical signs of bovine anaplasmosis (fever, anemia and reduction in milk production) and positive results in the blood smear examination and the qPCR. In the property located in the municipality of Mambai, the heifers and calves were four and three times, respectively, more likely to die from anaplasmosis than lactating cows (Figure 1).

**Anaplasma marginale detection and rickettsemia (qPCR) in dairy cattle**

The prevalence of *A. marginale* in the samples from the municipalities of Lins and Mambai was 84% and 34% by blood smear, 52% and 50% by IFAT, 58% and 54% by ELISA and 94% and 38% by qPCR, respectively (Table 1). The absolute quantification of *A. marginale* assessed by qPCR ranged from $7.77 \times 10^6$ to $1.97 \times 10^7$ in animals sampled in Lins and Mambai.

![Figure 1](image_url)  
**Figure 1.** Animal mortality rate during the anaplasmosis outbreak in a rural property located in the municipality of Mambai, state of Goiás, Central-West Region, Brazil. A total of 4,204 animals were evaluated including 1,050 calves, 954 heifers and 2,200 cows. The death cause was confirmed as anaplasmosis only for animals with apparent clinical signs and positive results in the blood smear and/or PCR assays.

<table>
<thead>
<tr>
<th>Local</th>
<th>Frequency of positivity (%)</th>
<th>Rickettsemia qPCR*</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Blood smear</td>
<td>ELISA</td>
</tr>
<tr>
<td>Lins SP</td>
<td>84%</td>
<td>58%</td>
</tr>
<tr>
<td>Mambai GO</td>
<td>34%</td>
<td>54%</td>
</tr>
</tbody>
</table>

*msp1a* copies per mL of blood in all positive animals.

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**Table 1.** Frequency (%) of animals positive for *Anaplasma marginale*. *Anaplasma marginale* was detected by direct examination (blood smears), serological tests (ELISA/IFAT) and molecular analysis (qPCR) of samples from naturally infected dairy cattle in São Paulo and Goiás states, Brazil.
1.27 × 10^1 to 1.49 × 10^4 in animals sampled in Mambai. All animals with positive results by blood smear were also positive by qPCR. The ELISA showed 89.6% and 92.6% agreement with the IFAT in the samples from the municipalities of Lins and Mambai, respectively. Variations in the serological and molecular prevalence of *A. marginale* were observed in calves, heifers, and pregnant and lactating cows (Table 2). In Mambai, the ELISA and qPCR prevalence of *A. marginale* was significantly higher in calves (p < 0.04) and heifers (p < 0.05) than in pregnant cows. Additionally, lactating cows showed a lower molecular prevalence of *A. marginale* than calves and heifers (p < 0.03).

**Analysis of *A. marginale* MSP1a sequences in cattle**

According to the structure of the MSP1a microsatellite of *A. marginale*, the genotypes E and H were observed in the samples from Lins and the genotypes C, D and E were found in cattle from Mambai (Table 3). The analysis of the MSP1a tandem repeats structure showed nine different *A. marginale* strains (t-10 -15, α-β^3, α-β^1-13, α-β^3-192, τ-β-100, α-β^3-Γ, 193-β-100, 191-13-Γ and 191-13-β) in cattle from Lins and two strains (α-β^1-Γ and E-F-φ^2-F^2) in cattle from Mambai (Table 3). This is the first report of the H genotype and the E-F-φ^2-F^2 strain in Brazil. Three new tandem repeats, 192, 191 and 193, were described in the present study (Figure 2). The tandem repeats 191 and τ only differed by the amino acid located at position 28: leucine (L) was observed at the tandem repeat τ, whereas serine (S) was found at the tandem repeat 191. The tandem repeats 192 and Γ differed only by the amino acid located at position 13, serine (S) and arginine (R) were found at tandem repeats Γ and 192, respectively. The tandem repeats 193 and 4 differed by two amino acids located at the positions 5 and 18, serine (S) and glutamine (Q) were observed at the tandem repeat 4, and threonine (T) and proline (P) were observed at the tandem repeat 193.

**Phylogenetic analysis**

The phylogenetic analysis identified four distinct groups among the studied samples (Figure 3). The analysis (NJ and ML) produced similar topologies and the same relationships for all major clusters that were identified in the present study and represented in the NJ tree. The phylogenetic tree was constructed based on the sequence of the MSP1a of *A. marginale* identified in cattle from the property located in Lins (t-10-15, α-β^3, α-β^1-13, α-β^3-192, α-β-100, 193-β-100, 191-13-Γ and 191-13-β) and Mambai (α-β^3-Γ and E-F-φ^2-F^2). In addition to the strains identified in the present study, *A. marginale* strains isolated in Brazil (states of Minas Gerais, Paraná, São Paulo and Rio de Janeiro), South Africa (SA), Argentina (Ar), the United States (USA), the Philippines (Ph),

### Table 2. Frequency (%) of calves, heifer, pregnant and lactation cows positive for *Anaplasma marginale*. *Anaplasma marginale* was detected by direct examination (blood smears), serological tests (ELISA/IFAT) and molecular analysis (PCR) of samples from naturally infected dairy cattle in Goiás State, Brazil.

<table>
<thead>
<tr>
<th>Class</th>
<th>Frequency of positivity (%)</th>
<th>Blood smear</th>
<th>IFAT</th>
<th>ELISA</th>
<th>qPCR</th>
</tr>
</thead>
<tbody>
<tr>
<td>Calves (n=10)</td>
<td>80</td>
<td>60</td>
<td>60</td>
<td>50</td>
<td></td>
</tr>
<tr>
<td>Heifer (n=20)</td>
<td>85</td>
<td>60</td>
<td>65</td>
<td>50</td>
<td></td>
</tr>
<tr>
<td>Pregnant cow (n=10)</td>
<td>70</td>
<td>20</td>
<td>20</td>
<td>20</td>
<td></td>
</tr>
<tr>
<td>Lactation cow (n=10)</td>
<td>50</td>
<td>50</td>
<td>60</td>
<td>20</td>
<td></td>
</tr>
</tbody>
</table>

### Table 3. Organization of the MSP1a tandem repeats and *Anaplasma marginale* strains isolated in naturally infected dairy cattle on farms located in the municipalities of Lins, state of São Paulo, and Mambai, state of Goiás, Brazil. *Anaplasma marginale* strain identification is based on *MSP1a*, including locality/microsatellite genotype – (tandem repeats structure); SD-ATG distance; and rickettsemia in the strains identified in cattle.

<table>
<thead>
<tr>
<th>Structure of MSP1a tandem repeats</th>
<th>Genotype</th>
<th>m</th>
<th>n</th>
<th>SD-ATG distance (nucleotide)</th>
<th>Rickettsemia* (msp1a copies/mL)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lins SP/7 (t-10-15)</td>
<td>H</td>
<td>3</td>
<td>6</td>
<td>25</td>
<td>7.75 × 10^4</td>
</tr>
<tr>
<td>Lins SP/10 (t-10-15)</td>
<td>H</td>
<td>3</td>
<td>6</td>
<td>25</td>
<td>2.02 × 10^4</td>
</tr>
<tr>
<td>Lins SP/16 (t-10-15)</td>
<td>H</td>
<td>3</td>
<td>6</td>
<td>25</td>
<td>5.95 × 10^4</td>
</tr>
<tr>
<td>Lins SP/11 (191-13-18)</td>
<td>E</td>
<td>2</td>
<td>7</td>
<td>23</td>
<td>6.34 × 10^4</td>
</tr>
<tr>
<td>Lins SP/12 (α-β^3)</td>
<td>E</td>
<td>2</td>
<td>7</td>
<td>23</td>
<td>7.29 × 10^4</td>
</tr>
<tr>
<td>Lins SP/110 (α-β^1-13)</td>
<td>E</td>
<td>2</td>
<td>7</td>
<td>23</td>
<td>6.13 × 10^4</td>
</tr>
<tr>
<td>Lins SP/703 (α-β^2-192)</td>
<td>E</td>
<td>2</td>
<td>7</td>
<td>23</td>
<td>9.24 × 10^4</td>
</tr>
<tr>
<td>Lins SP/1136 (τ-β-100)</td>
<td>E</td>
<td>2</td>
<td>7</td>
<td>23</td>
<td>1.42 × 10^4</td>
</tr>
<tr>
<td>Lins SP/1228 (α-β^3-Γ)</td>
<td>E</td>
<td>2</td>
<td>7</td>
<td>23</td>
<td>1.21 × 10^4</td>
</tr>
<tr>
<td>Lins SP/1450 (193-β-100)</td>
<td>E</td>
<td>2</td>
<td>7</td>
<td>23</td>
<td>6.42 × 10^4</td>
</tr>
<tr>
<td>Lins SP/1453 (191-13-Γ)</td>
<td>E</td>
<td>2</td>
<td>7</td>
<td>23</td>
<td>2.53 × 10^4</td>
</tr>
<tr>
<td>Mambai GO/1017B (α-β^3-Γ)</td>
<td>E</td>
<td>2</td>
<td>7</td>
<td>23</td>
<td>4.74 × 10^4</td>
</tr>
<tr>
<td>Mambai GO/1568B (α-β^3-Γ)</td>
<td>D</td>
<td>2</td>
<td>6</td>
<td>21</td>
<td>1.49 × 10^4</td>
</tr>
<tr>
<td>Mambai GO/1806B (E-F-φ^2-F^2)</td>
<td>C</td>
<td>2</td>
<td>5</td>
<td>19</td>
<td>3.57 × 10^4</td>
</tr>
</tbody>
</table>

*msp1a copies per mL of blood in all positive animals. The microsatellite (sequence in bold) was located between the Shine-Dalgarno (SD; sequence in brackets) and the translation initiation codon (ATG) with the structure: GTAGG (G/ATTT)m (GT)nT ATG. The SD-ATG distance was calculated in nucleotides as (4 × m) + (2 × n) + 1.*

Amino acid differences between tandem repeats A and 191, 192 and 193. The one-letter code is used for the different amino acids of the tandem repeats. Conserved amino acid positions are highlighted with asterisks. The amino acid at position 10 (-), which is evolving under negative selection (p < 0.25 using both FEL and SLAC methods), and residues of the immunodominant B-cell epitope is highlighted with asterisks. The amino acid at position 10 (-), which

** Figure 2.** Amino acid differences between tandem repeats A and 191, 192 and 193. The one-letter code is used for the different amino acids of the tandem repeats. Conserved amino acid positions are highlighted with asterisks. The amino acid at position 10 (-), which is evolving under negative selection (p < 0.25 using both FEL and SLAC methods), and residues of the immunodominant B-cell epitope (GARCIA-GARCIA et al., 2004) (box) are also shown.

The Mambaí sample 1806 was grouped with strains described in Curitiba (state of Paraná) as strains reported in Tamaulipas (Mexico), with 91% bootstrap support. The Lins' sample number 703 was positioned in a third group along with strains identified in Belo Horizonte (state of Minas Gerais), Táiacu (state of São Paulo), Jaboticabal (state of São Paulo) and Seropédica (state of Rio de Janeiro), Brazil, as well as strains reported in Tamaulipas (Mexico), with 91% bootstrap support. The Lins' sample number 703 was positioned in a third group along with strains identified in Curitiba (state of Paraná) and Táiacu (state of São Paulo), as well as with strains described in Argentina, Philippines and Taiwan, with 75% bootstrap support. The Mambai sample 1806 was grouped with strains described in Seropédica (state of Rio de Janeiro) with 84% of bootstrap support.

**Discussion**

*Anaplasma marginale* is endemic in Brazil, where anaplasmosis outbreaks cause economic losses to the livestock industry (VIDOTTO et al., 1998; KOCAN et al., 2010). However, studies are lacking on the *A. marginale* strains that circulate among cattle herds, causing clinical cases of bovine anaplasmosis in the world. The present study reports two bovine anaplasmosis outbreaks. One outbreak occurred in the municipality of Mambai, state of Goiás, Brazil, causing the death of 107 animals (37 calves, 45 heifers and 25 lactating cows). The other anaplasmosis outbreak occurred in a rural property located in the municipality of Lins, state of São Paulo, Brazil; no animals died, but lactating cows showed a drastic reduction in milk production.

Prior to the present study, the animals of the property located in the municipality of Mambai were kept in an alternate grazing system with center pivot irrigation. In this system, the animals had virtually no contact with ticks. However, due to a prolonged food shortage during the month of February 2012, some animals were transferred to an area of silvopasture, and when returned to the original area, they were infested with ticks. The bovine anaplasmosis outbreak began 14 to 20 days after the animals were transferred back to the center pivot irrigation system. Furthermore, due to the amount of feces accumulated during mechanical milking, a high number of flies were present at the fluid treatment station. Thus, two explanations exist for the occurrence of a bovine anaplasmosis outbreak on the property located in Mambai: (1) mechanical transmission of *A. marginale* by the high number of *Stomoxys calcitrans* flies originated from larvae that developed in faeces, accumulated in the environment and/or (2) the biological transmission of *A. marginale* by *Rhipicephalus* (Boophilus) *microplus* after the animals were placed in an area infested with cattle tick. Thus, in both scenarios, clinical cases of anaplasmosis and animal mortality occurred because animals were kept for a long period in a production system with no contact with *A. marginale* and, then, were exposed to the pathogen through tick and/or fly infestations. In contrast, in the rural property located in the municipality of Lins, the animals studied were from Uruguay and had been newly introduced into the herd. The introduction of infected animals into the herd has been proposed as a source of genetic diversity of *A. marginale* worldwide (FUENTE et al., 2007).

*Anaplasma marginale* prevalence was 94% (47/50) and 38% (19/50) according to the qPCR in bovines from the properties located in Lins and Mambai, respectively. However, of these samples, only 23% (11/47) of the animals from Lins and 16% (3/19) of the animals from Mambai were positive according to nPCR. In the present study, only samples with absolute quantification greater than 2.53 × 10^4 in the qPCR were positive by nPCR and subsequently sequenced. Thus, the different results observed in the nPCR and qPCR assays may have occurred due to differences in the level of rickettsiae in the samples and differences in the sensitivity of the PCR assays (FUENTE et al., 2001). In the present study, the samples negative by nPCR showed variation of 1.27 × 10^3 to 3.83 × 10^2 in absolute quantification by qPCR.

The results of the present study showed that the genetic diversity of the MSP1a of *A. marginale* was low among the cattle sampled in the property located in Mambai, and only two strains were identified (α-β-γ-δ-ε-F) and E-F-φ^2-.F). In the microsatellite analysis, the genotypes C, D and E were observed. Three possibilities can be considered to explain the genetic diversity of *A. marginale* from cattle on the property located in Mambai. (1) In an endemic region for bovine anaplasmosis, the low genetic diversity of *A. marginale* has been related to a lack of vectors (RUYBAL et al., 2009).
Figure 3. Characterization of *A. marginale* MSP1a sequences. Neighbor-joining phylogenetic tree of *Anaplasma marginale* MSP1a. The tree was constructed using the neighbor-joining method with *A. marginale* MSP1a sequences from strains identified in dairy cattle in the states of São Paulo and Goiás, Brazil. Bootstrap values are represented as percent on internal branches (1,000 replicates). The strains t-10-15, α-β2, α-β3-13, α-β2-192, α-β-100, 193-β-100, 191-13-Γ, 191-13-18, α-β-Γ and E-F-φ2-F2 are shown, as well as the *A. marginale* strains from cattle isolated in Minas Gerais (MG), Paraná (PR), São Paulo (SP), Rio de Janeiro (RJ), South Africa (SA), Argentina (Ar), the United States (USA), the Philippines (Ph), Israel (Is), Mexico (Me), Puerto Rico (Pr) and Taiwan (Ta). The strains isolated from cattle are identified as in the legend of the figure. The MSP1a GenBank accession numbers of the respective sequences used in the phylogenetic tree are shown.
The animals from Mambai had been subjected to an intensive tick control program, and they were free of ticks prior to the anaplasmosis outbreak. Thus, genetic diversity is likely low in cattle herds with a low level of infestation by tick vectors and where transmission occurs through bloodsucking flies. (2) Furthermore, cattle transfers between different geographic regions are an important source of dispersion of different *A. marginale* strains (FUENTE et al., 2007). However, the herd from Mambai was a closed herd; that is, there were no new animal introductions into the herd, and the replacement heifers were chosen from the nine infected calves born on the property. (3) Furthermore, herds infected by *A. marginale* for long periods exhibit high genetic diversity in the strains of this rickettsia (PALMER et al., 2001). Thus, the first and third possibilities seem to be the more applicable to animals on the property located in Mambai. The transmission of *A. marginale* by *R. (B.) microplus* into this herd was a recent event, and the transmission was maintained by bloodsucking flies, which resulted in low genetic diversity.

Our results showed that the genetic diversity of *A. marginale* was high in the cattle from the property located in the municipality of Lins, and nine different strains and two genotypes were identified. The results found in Lins are similar to those reported by Pohl et al. (2013) from cattle sampled in Belo Horizonte, state of Minas Gerais, southeastern Brazil, in which eight different strains of *A. marginale* and four different genotypes (B, D, E and G) were observed in the 13 samples sequenced. In our study, the high diversity of *A. marginale* in cattle from Lins, state of São Paulo, is most likely a consequence of different transmission mechanisms (biological and mechanical), each contributing to the genetic diversity of *A. marginale* in the herd. On both properties, despite acaricides were routinely used for controlling ticks and flies, infestation of the herd by these arthropods were reported even after treatment. Therefore, the occurrence of two independent transmission mechanisms (mechanical and biological) reflects a population with high genetic diversity in endemic areas (FUENTE et al., 2001). This same hypothesis was investigated in endemic regions by Silva et al. (2015a) in Brazil, where the animals were infected with multiple *A. marginale* strains. In the southeast and central-western regions of Brazil, where the present study was conducted, the tick *R. (B.) microplus* completes three to five generations per year (KASAI et al., 2000) and may, over time and during the feeding process, transmit new *A. marginale* strains to the cattle.

The prevalence of the C, D, E and H genotypes of *A. marginale* observed in the present study may indicate better adaptation by these genotypes, which has allowed for more efficient infection of the host. The genotypes identified in the present study have a dinucleotide tandem repeats between the Shine-Dalgarno sequence and the starting amino acid (SD-ATG) of 19, 21, 23 and 25 nucleotides. Estrada-Peña et al. (2009) evaluated the distribution of nine different genotypes in four distinct ecosystems worldwide and observed that the genotype E is the most common in South America, especially in Brazil and Argentina. However, the genotypes B, C, D and G have also been described previously in Brazil and Argentina (FUENTE et al., 2004; VIDOTTO et al., 2006; POHL et al., 2013; SILVA et al., 2015a, b). The present study describes for the first time the presence of the H genotype in Brazil.

The *t*-10-15 and *α*-β*-Γ*- strains were dominant in cattle sampled on the properties located in Lins and Mambai, respectively. The *t*-10-15 strain previously had been described in cattle from Brazil (VIDOTTO et al., 2006; SILVA et al., 2015a, b), and the strain *α*-β*-Γ*- had been detected in cattle in Argentina (RUYBAL et al., 2009), Mexico (ALMAZÁN et al., 2008), Taiwan (CABEZAS-CRUZ et al., 2013) and Brazil (SILVA et al., 2015b), countries where bovine anaplasmosis is an endemic disease. In the present study, the most commonly observed tandem repeats of the MSP1a were *α*, *β* and *τ*. These results support previous studies on the *A. marginale* strains that circulate in South America, which show the most common tandem repeats are 4, 8, 16, 56, 60, 64, 67, *α*, *β*, *γ*, *π* and *τ* (ESTRADA-Peña et al., 2009; SILVA et al., 2015a, b). In addition, for the isolates described from South America, the most common tandem repeats in the first repetition are 4, 16, 72, *α* and *τ* in Brazil (VIDOTTO et al., 2006; POHL et al., 2013; SILVA et al., 2015a, b) and *α*, *τ* and *γ* in Argentina (RUYBAL et al., 2009). Previous studies have associated the presence of the strain *α*-β*-Γ*- with the occurrence of clinical anaplasmosis in Mexico (ALMAZÁN et al., 2008) and Brazil (SILVA et al., 2015b); however, the present study is the first that associates the occurrence of the *t*-10-15 strain with animal mortality. Thus, we believe that the *A. marginale* strains *t*-10-15 and *α*-β*-Γ*- may be involved in the occurrence of bovine anaplasmosis outbreaks in endemic areas.

When we compared the repeated sequences found in the *A. marginale* samples evaluated in the present study with sequences already known (CABEZAS-CRUZ et al., 2013), we identified three new repeated sequences. The tandem repeats 191 and *τ* differed only by the amino acid located in position 28. In contrast, the tandem repeats 192 and *Γ* differed only by the amino acid located at position 13, and the tandem repeats 193 and 4 differed by two amino acids located at positions 5 and 18. These findings suggest that the tandem repeats 191, 192 and 193 may have evolved recently from the *τ*, *Γ* and 4 tandem repeats, providing evidence for increased genetic diversity of *A. marginale* in cattle. Following this hypothesis, the phylogenetic analysis based on the MSP1a sequences indicated that the strains Lins 11 (191-13-18), Lins 1453 (191-13-Γ) and Lins 1450 (193-Γ-100) possibly evolved from the strain *t*-10-15, whereas Lins 703 (α-β*-2-192) evolved from the strain α-β*-Γ*. Considering that in certain regions ticks might not play an important role in transmitting *A. marginale* between cattle, the strains most commonly found in these regions might be transmitted mechanically. The negatively charged amino acids, aspartate (D) and glutamic acid (E), at position 20 are critical for the interaction of the cell of the invertebrate host with the MSP1a of *A. marginale* (FUENTE et al., 2003a). These amino acids affect the secondary structure of MSP1a, which seems to affect the transmission of *A. marginale* by ticks (CABEZAS-CRUZ et al., 2013). In agreement with these results, 50% of the tandem repeats of the MSP1a of *A. marginale* obtained in the present study had the amino acid glycine (G) at position 20. Thus, in the two studied herds,
the circulating *A. marginale* strains can be transmitted by both mechanisms: biologically and mechanically.

**Conclusion**

The τ-10-15 and α-β3-Γ strains were predominantly associated with the occurrence of clinical anaplasmosis in calves from southeastern Brazil, whereas the α-β3-Γ strain was predominantly associated with mortality of calves, heifers and lactating cows in central-western Brazil. In addition to the occurrence of clinical anaplasmosis, we also observed high rickettsemia and low serological prevalence of *A. marginale*. The genetic diversity of MSP1a was low in cattle from the municipality of Mambai, where animals died due to anaplasmosis, and in high animals from the municipality of Lins, where the animals did not die but had drastic reduction in milk production. One of the factors that likely contributed to the occurrence of clinical anaplasmosis in the studied herds may be the low population of ticks and bloodsucking flies, which may have reduced the levels of antibodies and the responsiveness of the cattle to *A. marginale*.

**References**


