Morphological and molecular characterization of *Haemoproteus coatneyi* and *Haemoproteus erythrogravidus* (Haemosporida: Haemoproteidae) in Passeriformes in Brazil's Atlantic Forest

Caracterização morfológica e molecular de *Haemoproteus coatneyi* e *Haemoproteus erythrogravidus* (Haemosporida: Haemoproteidae) em Passeriformes da Mata Atlântica, Brasil

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Abstract

*Haemoproteus* spp. are protozoan parasites found in birds around the world. These parasites are identified through the morphology of gametocytes, phylogenetic analysis based on the mitochondrial *cytb* gene, and the parasite’s geographic distribution. The absence of erythrocytic merogony, high intraspecific genetic variation and low parasitemia in wild birds makes it essential to use integrative approaches that assist in the identification of these parasites. Thus, microscopic and molecular analyses, combined with spatial distribution, were carried out to verify the presence of *Haemoproteus* spp. in wild birds in Brazil. Light microscopy revealed one *Tangara sayaca* bird was parasitized by *Haemoproteus coatneyi* and, two specimens of *Zonotrichia capensis* presented *Haemoproteus erythrogravidus*. The morphology of the gametocytes of these two parasitic species showed high similarity. The molecular analysis revealed the presence of one lineage of *H. coatneyi* and two lineages of *H. erythrogravidus*, one of which is considered a new lineage. These lineages were grouped phylogenetically in separate clades, with low genetic divergence, and the *H. erythrogravidus* lineage emerged as an internal group of the lineages of *H. coatneyi*. The geographic distribution demonstrated that the two species occur in the American continent. This is the first report of *H. erythrogravidus* in Brazil.

Keywords: birds, haemosporidians, *Tangara sayaca*, *Zonotrichia capensis*, geographic distribution.

Resumo

*Haemoproteus* spp. são protozoários parasitos encontrados em aves de todo o mundo. A identificação desses parasitos é realizada por meio da morfologia dos gametócitos, da análise filogenética, baseada no gene mitocôndrial *cytb* e na distribuição geográfica do parasito. A ausência de merogonia eritrocítica, a alta variação genética intraespecífica e a baixa parasitemia em aves silvestres, tornam essencial a utilização de abordagens integrativas que auxiliem na identificação desses parasitos. Assim, análises microscópicas e moleculares, aliadas à distribuição espacial, foram realizadas para verificar a presença de *Haemoproteus* spp. em aves silvestres no Brasil. A microscopia óptica demonstrou que uma ave *Tangara sayaca* estava parasitada por *Haemoproteus coatneyi*, e dois espécimes de *Zonotrichia capensis* apresentavam *Haemoproteus erythrogravidus*, cujas morfologias dos gametócitos apresentaram alta similaridade. A análise molecular recuperou uma linhagem de *H. coatneyi* e duas linhagens de *H. erythrogravidus*, sendo uma dessas considerada nova linhagem. Essas linhagens se agruparam...
The traditional taxonomic classification of avian haemosporidians is based on life history and morphology. The main form of identification of haemosporidian species is still based on the morphological characterization of blood stages found in the vertebrate host, combined with the molecular characterization of species (Valkiūnas & Iezhova, 2018). However, several factors can influence or hinder the correct identification of these hemoparasites. In the case of the genus *Haemoproteus*, for example, the morphological description is restricted to the characteristics of the gametocytes, given that the species of this group do not perform erythrocytic merogony and that they have few characters for differentiation at the species level. Moreover, some of these characters are extremely similar, making their identification difficult, especially in cases of cryptic species (Garnham, 1966; Dimitrov et al., 2016). In addition, wild birds generally have lower parasitemia than captive birds or those used in experimental infections to assess the susceptibility of the host to haemosporidians. This low parasitemia can make it impossible to differentiate species, given the absence of all erythrocytic stages or mature gametocytes (Dimitrov et al., 2016).

Two species of the genus *Haemoproteus* that infect wild birds are *Haemoproteus coatneyi* and *Haemoproteus erythrogravidus*. *Haemoproteus coatneyi* has been described only through morphological analysis (Burry-Caines & Bennett, 1992), whereas *H. erythrogravidus* has been described by morphological and molecular data based on a cytb fragment, and also through complete mitochondrial genome sequencing (Mantilla et al., 2016). According to these authors, *H. erythrogravidus* is closely related to *H. coatneyi*, distinguished morphologically by the formation of a marked protrusion on the envelope of infected erythrocytes by the majority of developing gametocytes and by the extremely attenuated width of the growing dumbbell-shaped macro and microgametocytes.

Host specificity and geographic distribution are currently not considered valid taxonomic characters for identifying haemosporidian species. However, these aspects have been used successfully in the taxonomy of Haemoproteidae (Valkiūnas & Ashford, 2002) and can be used as additional information in molecular and morphological methods. *Haemoproteus coatneyi* does not appear to be host-specific, and has been recorded in more than 20 avian species (Valkiūnas, 2005). However, only two hosts (*Zonotrichia capensis* and *Anisognathus somptuosus*) have been recorded for *H. erythrogravidus* (MalAvi, 2020; GenBank, 2020), although there are several studies of this species. In addition, *H. coatneyi* has a greater geographic distribution, and has been recorded in Canada, the United States, Mexico, Colombia and Peru, while *H. erythrogravidus* has been identified only in Colombia, Peru and Brazil (MalAvi, 2020; GenBank, 2020).

Thus, integrated methodologies that perform morphological and molecular analysis of parasite lineages, specificity of the host-parasite relationship, and geographic distribution may be essential for the identification of very similar species of the genus *Haemoproteus*. In order to improve the understanding of infection by *Haemoproteus* spp. in wild birds, this study focused on the morphological and molecular characterization of *Haemoproteus* spp. and compared the data obtained with previous records in different hosts and geographic locations.
**Material and methods**

**Area of study and blood collection**

The birds were caught between the years 2013 and 2015 in fragments of Atlantic Forest in the Zona da Mata of Minas Gerais, Brazil, as follows: Sítio Paraíso da Barra - Chácara, 21.68° S, 43.22° W, 796 m altitude; Sítio Vista Alegre - Juiz de Fora, 21.48° S, 43.15° W, 494 m altitude; Granja Passarada, Juiz de Fora, 21.43° S, 43.22° W, 715 m altitude and Fazenda Volta Grande - Santa Bárbara do Monte Verde, 21.58° S, 43.41° W, 798 m altitude.

The birds were trapped using mist nets and were marked with rings in order to avoid recapturing them. The bird species were identified according to Ridgely & Tudor (2009) and Sigrist (2014), based on morphometric analysis and phenotypic characteristics such as size, plumage color, beak and legs morphology. Blood samples were collected by brachial venipuncture from the species: *Arremon semitorquatus*, *Zonotrichia capensis* and *Ammodramus humeralis*, belonging to the family Passerellidae, and *Tangara sayaca*, belonging to the family Thraupidae. Immediately after preparing blood smears on slides, they were air dried and then fixed in absolute methanol for 3 min at the same collection site. For the molecular procedures, approximately 30 μl of blood were placed in microtubes without anticoagulants and stored in a freezer at -20°C.

**Morphological and morphometric analyses**

Blood smears were stained with Giemsa solution (SIGMA) diluted in distilled water at a ratio of 1:9. The slides were examined under Olympus BX-51 light microscope coupled to an Olympus Evolt E-330 digital camera to record the evolutionary forms of parasites. The slides were then examined under 600X magnification for 20 min to detect the presence of hemoparasites. Then, 100 microscopic fields were observed under 1000x magnification to calculate the parasitemia. Image-Pro Plus 6.0 imaging software was used to perform the morphometric analysis of gametocytes, following parameters set by Valkiūnas (2005).

**DNA extraction, PCR and sequencing**

Only the samples that were positive for the presence of gametocytes in blood smears were subjected to molecular analysis. DNA was extracted from a 20 μl blood sample from each bird, using Wizard® Genomic DNA purification kit (Promega®, Sao Paulo, Brazil), according to the manufacturer's recommendations. DNA was quantified by spectrophotometry on a Nanodrop ND-2000® spectrophotometer (ThermoScientific, Wilmington, DE, USA). The total DNA concentration was standardized to 100 ng/μL and stored at -20°C for further amplification by PCR. A 618 bp fragment of the mitochondrial cyt b gene of the haemosporidian parasites was amplified using nested PCR. The primers used in the first reaction, HaemFNI (5′-CATATATTAAGAGAAITATGGAG-3′) and HaemNRI (5′-ATAGAAAGATAAGAAATACCATTC-3′), served to amplify species of the genera *Plasmodium*, *Haemoproteus* and *Leucocytozoon* (Hellgren et al., 2004). The reaction was carried out in a final volume of 25 μL, containing: 12.5 μL of 1X of Go Taq Green Master Mix (Promega, MA, USA); 2.5 μL of ultra-pure water; 1.0 μM of each primer; and 5 μL of genomic DNA (20–100 ng). In the second reaction, the primers HaemF (5′-ATGGTGCTTTCGATATATGCATG-3′) and HaemR2 (5′-GCATTATCTGGATGTGATAATGGT-3′) were used to amplify a fragment of approximately 479 bp of the *Plasmodium* and *Haemoproteus cyt b* genes (Bensch et al., 2000). The reaction was carried out in a final volume of 25 μL, containing: 12.5 μL of 1X of Go Taq Green Master Mix; 0.5 μM of each primer; and 2 μL of the product of the first reaction. Nuclease-free water was used in the amplification reactions for the negative control and DNA extracted from a sample of *Plasmodium (Haemamoeba) gallinaceum* obtained from an experimental infection study conducted by the Federal University of Minas Gerais (Brazil) was used as a positive control (Rodrigues et al., 2020).

The following amplification conditions were established for the two reactions: 94°C for 3 minutes; 35 cycles of 94°C for 30 seconds; 52°C for 30 seconds; 72°C for 45 seconds; and a final extension at 72°C for 10 minutes. The PCR products were separated by electrophoresis on 2% agarose gel, stained with Blue Green Loading Dye I (LGC Biotecnologia®, Cotia, São Paulo, Brazil) and examined under ultraviolet light using a GE Healthcare® UV transilluminator. The products amplified in the second PCR reaction were purified with an QIAquick® Purification
Kit (Qiagen®, São Paulo, Brazil), and subjected to bidirectional sequencing in a 3130xL Genetic Analyzer (Applied Biosystems®, Carlsbad, California), by Sanger sequencing approach (Sanger et al., 1977), following the manufacturer’s instructions.

Phylogenetic analyses

Phylogenetic reconstructions were performed using a dataset containing 129 sequences attributed to *Haemoproteus* morphospecies, including all *H. coatneyi* (*n* = 44) and *H. erythrogravidus* (*n* = 6) sequences available in the MalAvi and GenBank databases (accessed on March 26, 2020), as well as the three sequences obtained in this study. The sequences of other haemosporidians amplified during the molecular analysis were automatically excluded. Three sequences of *Plasmodium* sp. morphotypes and one sequence of *Leucocytozoon buteonis* were used as outgroups.

Using the standard configuration, the sequences were aligned in the MAFFT software (Katoh et al., 2019) and then visually inspected. After removing poorly aligned sites with Gblocks server (Talavera & Castresana, 2007), a matrix with 479 bp was obtained. The phylogeny of *Haemoproteus* was analyzed by Bayesian inference using MrBayes software in XSEDE v. 3.2.6 (Ronquist et al., 2012), available on the CIPRES Portal. A trillion generations were carried out in two simultaneous and independent Markov Chain Monte Carlo simulations, excluding 25% of the trees generated (“Burn-in”), using the GTR + GAMMA + I model, with 4 gamma categories (Tavaré, 1986; Yang, 1994). The best model of nucleotide substitution was evaluated using the Jmodel Test software implemented in MEGA X (Kumar et al., 2018). The resulting trees were viewed and edited using the FigTree 1.4.4 software. To calculate the genetic divergence between the lineages obtained in this study and others already deposited in databases, the MEGA X software (Kumar et al., 2018) was used, following the program’s standard configuration.

**Results**

Blood smears from 65 birds were examined, 37 of which were *Tangara sayaca* (Thraupidae), 28 Passerellidae including 1 *Ammodramus humeralis*, 12 *Arremon semitorquatus* and 15 *Zonotrichia capensis*. The average prevalence found in the family Thraupidae was 2.70% (*n* = 1) with 0.04% parasitemia, and that in the family Passerellidae was 7.14% (*n* = 2), with parasitemia ranging from 0.15 to 0.36%.

The quality of blood smears and precise microscopic examination permitted confirmation of a single infection by *Haemoproteus coatneyi* in one *T. sayaca* (Figure 1) and of *Haemoproteus erythrogravidus* in two *Z. capensis* (Figures 2 and 3). The main similarities between the two parasite species were young and mature gametocytes with

*Figure 1. Haemoproteus coatneyi* in the blood of *Tangara sayaca* (lineage TSJB01). Young and growing gametocytes (a – d); macrogametocytes (e – k); microgametocytes (l – p). Arrowheads indicate the pigment granules. Arrows indicate the nucleus of the parasites. Bar = 12 µm.
longitudinal growth, adhered to the nucleus and in a dumbbell-shaped, nucleus subterminal to terminal position in the macrogametocytes, which were indistinguishable in the microgametocytes, and pigment granules randomly distributed in the macrogametocyte cytoplasm and generally grouped at the microgametocytes poles. Some mature gametocytes of the two parasites also showed a slight constriction in the central part of the parasite and a tendency to form a protrusion similar to a “balloon” on the non-parasitized side of the erythrocyte. The vacuoles in the cytoplasm was verified only in *H. erythrogravidus*.

**Figure 2.** *Haemoproteus erythrogravidus* in the blood of *Zonotrichia capensis* (lineage ZCSB01). Young gametocytes (a – d); macrogametocytes (e– j); microgametocytes (k – p). Black arrowheads indicate pigment granules. Black arrow indicates the parasite's nucleus. White arrowheads indicate attenuation in the central part of the gametocytes. Black arrows with open heads indicate vacuoles. White arrows indicate protrusions formed in the infected erythrocytes. Bar = 11 µm.

**Figure 3.** *Haemoproteus erythrogravidus* in the blood of *Zonotrichia capensis* (lineage ZCSB02). Young gametocytes (a – d); macrogametocytes (e – j); microgametocytes (k – p). White arrowheads indicate attenuation in the central part of the gametocytes. Black arrow indicates the parasite's nucleus. Black arrows with open heads indicate vacuoles. Black arrow heads indicate pigment granules. White arrows indicate protrusions formed in the infected erythrocytes. Bar = 11 µm.
The parasite morphometry measured during the microscopic analysis was compared to the original descriptions of *H. coatneyi* and *H. erythrogravidus*, and is shown in Table 1.

In the molecular analysis, three *Haemoproteus* spp. lineages were observed. The TSJB01 lineage corresponds to the morphospecies described in *T. sayaca* (Figure 1). The ZCSB01 and ZCSB02 lineages correspond to the morphospecies described in *Z. capensis* (Figures 2 and 3). The sequences obtained were deposited in the GenBank database under access numbers MT656006 (TSJB01), MT656007 (ZCSB01) and MT656008 (ZCSB02). The chromatograms analysis confirmed the presence of a single infection in the absence of double peaks in both directions, forward and reverse. The ZCSB02 lineage did not show 100% similarity with any other one previously included in the genetic databases (MalAvi and GenBank), as is therefore considered a new lineage.

In the phylogenetic reconstruction (Figure 4), the lineages obtained from the databases, together with our lineages of *H. coatneyi* and *H. erythrogravidus*, were grouped into four clades (Figure 4; Clades I – IV). Clade I, in which the new TSJB01 lineage was inserted, was composed of 20 *H. coatneyi* lineages. Six lineages of *H. erythrogravidus* were grouped in clade II, including our ZCSB01 and ZCSB02 (II-a) lineages, in addition to 8 lineages of *H. coatneyi* (II-b). Clade III was composed of 14 *H. coatneyi* lineages. Clade IV was composed of two *H. coatneyi* lineages and one *Haemoproteus cyanomitrae* lineage. It should be kept in mind that the clade that groups the *H. erythrogravidus* (II-a) lineage emerged as an internal group of a larger clade that also contains of *H. coatneyi* lineages (clade II). Furthermore, in clades III and IV, both *H. coatneyi* lineages showed a lineage of the species *Haemoproteus tartakovskyi*.

Table 1. Comparative morphometry of the gametocytes of *Haemoproteus coatneyi* in *Tangara sayaca* and *Haemoproteus erythrogravidus* in *Zonotrichia capensis*.

| Characteristic          | *H. coatneyi* *a*          | *H. coatneyi* Burry-Caines & Bennett (1992) | *H. erythrogravidus* *a*          | *H. erythrogravidus* Mantilla et al. (2016) |
|-------------------------|---------------------------|---------------------------------------------|----------------------------------|--------------------------------------------|
| Uninfected erythrocytes |                           |                                             |                                  |                                            |
| Length                  | 10.9–13 (11.9 ± 0.5)      | 11.9 (0.8)                                  | 10.1–11.9 (11 ± 0.5)            | 10.3-11.7 (10.9 ± 0.4)                     |
| Width                   | 6–7.1 (6.3 ± 0.3)         | 6.2 (0.6)                                   | 5.6–6.7 (6.2 ± 0.3)            | 5.8-6.5 (6.1 ± 0.2)                       |
| Area                    | 53.6–71.6 (60.9 ± 3.8)    | 59.2 (7.4)                                  | 46.7–60.6 (55.2 ± 3.7)         | 41.8-56.3 (47.9 ± 3.7)                     |
| Nucleus of uninfected erythrocytes |                    |                                             |                                  |                                            |
| Length                  | 5.3–6.4 (5.8 ± 0.3)       | 5.5 (0.5)                                   | 5.1–6.1 (5.6 ± 0.3)            | 4.7-5.6 (5.1 ± 0.1)                       |
| Width                   | 2.3–3.1 (2.6 ± 0.2)       | 2.4 (0.3)                                   | 2.4–3.1 (2.7 ± 0.2)            | 1.9-2.4 (2.1 ± 0.1)                       |
| Area                    | 11.6–15.3 (13.4 ± 1)      | 11.0 (1.8)                                  | 11.6–14.2 (13.1 ± 0.8)         | 7.5-9.1 (8.2 ± 0.5)                       |
| Macrogametocytes        |                           |                                             |                                  |                                            |
| n = 21                  |                           |                                             |                                  |                                            |
| Infected erythrocytes   |                           |                                             |                                  |                                            |
| Length                  | 11.2–13.3 (12.4 ± 0.5)    | 12.3 (0.9)                                  | 11–13 (11.8 ± 0.5)             | 10.3-11.7 (11.1 ± 0.5)                     |
| Width                   | 4.7–7.1 (5.9 ± 0.6)       | 6.2 (0.7)                                   | 5–7.4 (6.4 ± 0.7)              | 5.4-7.2 (6.4 ± 0.7)                       |
| Area                    | 54.1–74.1 (63.4 ± 6.1)    | 63.4 (8.7)                                  | 52.7–72.1 (64.2 ± 5.2)         | 49.6-56.7 (52.7 ± 2.1)                     |
| Nucleus of infected erythrocytes |                    |                                             |                                  |                                            |
| Length                  | 4.4–5.7 (5.1 ± 0.3)       | 5.4 (0.5)                                   | 5.1–6.1 (5.6 ± 0.3)            | 4.1-5 (4.6 ± 0.2)                         |
| Width                   | 1.7–2.7 (2.3 ± 0.2)       | 2.3 (0.4)                                   | 2.3–3.6 (2.7 ± 0.3)            | 2-2.2 (2.1 ± 0.05)                        |
| Area                    | 9.1–12.8 (10.4 ± 1.1)     | 10.6 (2.1)                                  | 10.5–17 (12.8 ± 1.3)           | 6.4-8 (7.4 ± 0.5)                         |

*aAll measurements are provided in minimum and maximum values. The averages are in parentheses together with the standard deviation; *bRange of variation and the average value of the nuclear displacement ratio (NDR) of infected erythrocytes which is calculated according to the formula: NDR = 2x/(x+y) Bennett & Campbell (1972). n = Number of gametocytes, macrogametocytes and microgametocytes measured.
Table 1. Continued...

| Characteristic | H. coatneyi * (This study) | H. coatneyi Burry-Caines & Bennett (1992) | H. erythrogravidus * (This study) | H. erythrogravidus Mantilla et al. (2016) |
|----------------|---------------------------|------------------------------------------|-----------------------------------|-----------------------------------------|
| Gametocytes    | n = 21                    | n = 150                                  | n = 21                            | n = 25                                  |
| Length         | 11.4–15 (13.1 ± 1.1)       | 16.5 (3.4)                               | 11.4–14.1 (12.9 ± 0.7)            | 10.3–12 (11.1 ± 0.6)                    |
| Width          | 1.5–3.2 (2.1 ± 0.4)        | 2.0 (0.9)                                | 0.9–2.1 (1.4 ± 0.3)               | 2.3–7.2 (4.1 ± 1.7)                     |
| Area           | 30.8–50.6 (36.8 ± 5.9)     | 39.4 (7.9)                               | 26.3–37.8 (31.7 ± 3.8)            | 34.3–40 (36.9 ± 1.8)                    |

**Nucleus of Gametocytes**

| Length         | 1.3–3.3 (2.2 ± 0.5)        | 2.4 (0.6)                                | 1.3–3 (2.2 ± 0.4)                 | 1.4–3.1 (2.2 ± 0.4)                     |
| Width          | 0.7–2.0 (1.2 ± 0.3)        | 2.1 (0.5)                                | 0.8–1.9 (1.4 ± 0.3)               | 1–2.2 (1.6 ± 0.39)                      |
| Area           | 1.4–4.9 (2.8 ± 0.9)        | 4.4 (1.6)                                | 1.5–4.1 (2.8 ± 0.8)               | 2.3–4.5 (3.6 ± 0.7)                     |

**No. Pigment granules**

| 9–17 (13 ± 2.7) | 11.6 (2.9)                | 8–14 (11 ± 1.6)                         | 12–17 (13.5 ± 1.6)                |

**NDR**

| 0.4–1 (0.8 ± 0.2) | 0.8 (0.2)                | 0.8–1.3 (1 ± 0.1)                       | 0.5–0.8 (0.6 ± 0.1)                |

**Microgametocytes**

| n = 21 | — | n = 21 | n = 11 |

**Infected erythrocytes**

| Length         | 11.7–13.4 (12.6 ± 0.5)   | — | 11.1–12.9 (12 ± 0.5) | 13.8–16.9 (15.3 ± 1)               |
| Width          | 5.1–7.7 (6.4 ± 0.8)      | — | 6–7.7 (6.8 ± 0.4)    | 6.8–8.4 (7.6 ± 0.6)                |
| Area           | 52.5–78.9 (67.4 ± 7.5)   | — | 56.6–74 (65.8 ± 4.2) | 36.7–45.9 (40.6 ± 2.9)             |

**Nucleus of infected erythrocytes**

| Length         | 5.1–6 (5.5 ± 0.3)        | — | 5.2–6.1 (5.7 ± 0.3)  | 6.2–8.2 (6.9 ± 0.6)                |
| Width          | 2.1–3 (2.5 ± 0.2)        | — | 2.5–3.7 (2.9 ± 0.3)  | 2.2–3.1 (2.6 ± 0.3)                |
| Area           | 10.9–13.7 (12.3 ± 0.7)   | — | 11.1–16.9 (13.7 ± 1.5) | 15.4–21.7 (18 ± 2.1)              |

**Gametocytes**

| n = 21 | — | n = 21 | n = 11 |

| Length         | 12.3–14.9 (13.4 ± 0.8)   | — | 12.9–15.4 (14.2 ± 0.9) | 8.4–12.1 (10.4 ± 1.2)               |
| Width          | 1.7–4.3 (2.9 ± 0.8)      | — | 2.5–3.7 (2.9 ± 0.3)    | 0.7–3.3 (2.1 ± 0.7)                |
| Area           | 31.1–54 (43.9 ± 6.6)     | — | 30–44.1 (36.5 ± 3.9)   | 21.3–30.6 (27.4 ± 2.7)             |

**Nucleus of Gametocytes**

| — | — | — | — |

**Length**

| — | — | — | — |

**Width**

| — | — | — | — |

**Area**

| — | — | — | — |

**Number of Pigment granules**

| 9–16 (12.2 ± 2.1) | — | 7–13 (9.7 ± 1.8) | 8–15 (10 ± 1.2) |

**NDR**

| 0–0.9 (0.4 ± 0.2) | — | 0.7–1.2 (0.9 ± 0.1) | 0.7–1.2 (0.9 ± 0.1) |

*All measurements are provided in minimum and maximum values. The averages are in parentheses together with the standard deviation; *Range of variation and the average value of the nuclear displacement ratio (NDR) of infected erythrocytes which is calculated according to the formula: NDR = 2x/(x+y) Bennett & Campbell (1972). n = Number of gametocytes, macrogametocytes and microgametocytes measured.
The average genetic divergence between lineages within clades are: clade I - 0.20% (0.00-1.91); clade II - 1.72% (0.00-2.97); clade III - 1.86% (0.00-2.89); clade IV - 2.82% (0.00-4.14). The average intraspecific genetic divergence between lineages within clade II, where *H. coatneyi* and *H. erythrogravidus* lineages were found, was 1.92% (clade II-a) and 1.68% (clade II-b), respectively. The maximum intraspecific genetic divergence between the *H. erythrogravidus* lineage available in the databases and our lineages (ZCSB01 and ZCSB02) was 2.89%, while the maximum intraspecific genetic divergence between the *H. coatneyi* lineages available in the databases and our lineage (TSJB01) was 3.10%. The intraspecific genetic divergence between *H. coatneyi* lineages was often greater than the interspecific genetic divergence. For example, the difference between the DENCOR01 and KF537297 lineages, both
from \( H. \) coatneyi, was 4.14\%, while the difference between the DENCOR01 lineage (\( H. \) coatneyi) and the CYAOLI03 lineage (\( H. cyanomitrae \)) was 2.28\%. The genetic divergences between the lineages used in this study to perform phylogenetic analyses are listed in Figure 5.

The lineages of the present study (ZCSB01, ZCSB02 and TSJB01) and the other \( H. \) coatneyi and \( H. erythrogravidus \) lineages were recorded in the American continent, in hosts of the order Passeriformes (Figure 6). The species \( H. \) coatneyi has been found in North, Central and South America, with the largest number of records in Colombia and Peru. This is the first report of this species in Brazil. \( H. \) erythrogravidus has been recorded only in South America, in the countries Colombia, Peru and Brazil, and in the latter country we recorded a new lineage infecting wild birds.

**Discussion**

In this study, morphological and molecular evidence was presented of hemoparasites of the genus *Haemoproteus* in wild birds of the Atlantic Forest. The phylogenetic reconstruction showed that the ZCSB01 and ZCSB02 lineages of *Z. capensis* and TSJB01 of *T. sayaca* are closely related to the *H. erythrogravidus* and *H. coatneyi* lineages, respectively. The TSJB01 lineage showed 100\% similarity with the NEOFAS02 strain, which was recorded in *Neothraupis fasciata*, but was not attributed to any morphospecies. This species of bird belonging to the family Thraupidae is recorded in Brazil’s Cerrado biome (MalAvi, 2020; Fecchio et al., 2017). The ZCSB01 lineage showed 100\% similarity with ZOCAP01, a lineage previously associated with the *H. erythrogravidus* morphospecies (Mantilla et al., 2016), while the ZCSB02 lineage showed 99\% similarity with ZCSB01 and ZOCAP01, thus corresponding to a new lineage, which will therefore be added to the databases in order to facilitate new studies on these species.

Based on published sequences, the distribution of haemosporidians and their hosts showed that \( H. \) coatneyi is the species with the greatest geographical distribution, occurring in South, Central and North America, whereas *H. erythrogravidus* occurs only in South America. It should be noted that this paper offers the first record of \( H. \) coatneyi in Brazil, thus expanding the body of knowledge about the geographic distribution of this parasite. There are areas of overlap in the occurrence of the two species, which may even be using the same vectors for dispersion. These patterns of spatial distribution of the diversity of parasitic species may be related to ecologically distinct habitats, environmental variables and the abundance of vectors (Sol et al., 2000; Sehgal et al., 2011; Loiseau et al., 2012; Clark et al., 2014). It is known that parasites have a series of effects on their hosts, influencing the temporal and spatial dynamics of natural populations and bird communities (Leite et al., 2013). In addition, knowledge about the spatial distribution of parasites, when correlated with the diversity and abundance of the vector, can contribute to protect public health, since these vectors can also transmit other diseases, including those that occur in humans (Grillo, 2009).

The morphological characterization demonstrated that the gametocytes of *Haemoproteus* spp. identified in *T. sayaca* and in the two specimens of *Z. capensis* showed morphological similarities with the original descriptions of the gametocytes of *H. (Parahaemoproteus) coatneyi* and *H. (Parahaemoproteus) erythrogravidus*, respectively (Burry-Caines & Bennett, 1992; Mantilla et al., 2016). These gametocytes also showed a slight constriction in the central part of the parasite, a rather uncommon characteristic, but which is also mentioned in the original description of *H. coatneyi* (Burry-Caines & Bennett, 1992). In addition, some mature gametocytes from the two parasites showed a tendency to form a balloon-like protrusion on the non-parasitized side of the erythrocyte, a characteristic associated only with *H. erythrogravidus* (Mantilla et al., 2016). It is known that variations may occur in the shape and number of gametocytes in *Haemoproteus* species when smears are produced at different times in the infection process, i.e., in acute, chronic or recurrent infections, or when the preparation of blood smears is slow and air humidity is high (Valkiūnas, 2005). However, as the smears of these birds were of good quality and they were prepared correctly, the changes observed may be ascribed to the normal progressive development of the parasites, since different forms of the same *Haemoproteus* species have already been observed in smears prepared at different times from the same bird (Burry-Caines & Bennett, 1992). Thus, these two unusual characteristics are not enough to conclude that the species evaluated in this study are different from *H. coatneyi* and *H. erythrogravidus*.

Mantilla et al. (2016) used two morphological features to differentiate *H. coatneyi* from *H. erythrogravidus* that infected *Z. capensis* in Ecuador and Colombia. These features are similar to the unusual ones listed above, which were found in *T. sayaca* and *Z. capensis* (slight constriction in the central part of the parasite and balloon-like protrusion on the non-parasitic side of the erythrocyte). These characteristics have already been seen in other *Haemoproteus* species, such as: *Haemoproteus fringillae*, which had slightly visible protrusions in their envelopes and
Figure 5. Genetic divergence based on 479 bp fragments of the mitochondrial cytb gene between *Haemoproteus coatneyi* and *Haemoproteus erythrogravidus* lineages used in phylogenetic analyses. The lineages of the present study are given in bold font.
Haemoproteus attenuatus gametocytes, which have a markedly attenuated width in their central region, showing the dumbbell shape (Valkiūnas, 2005). Mantilla et al. (2016) stated that such protrusions were visible only in the final stages of development of gametocytes, and were less prominent; besides, the obtained H. fringillae and H. erythrogravidus sequences were clearly genetically dissimilar, with a 4.3% to 4.5% genetic divergence based on the 479 bp fragment of the cytb gene. The authors also stated that the feature of attenuated width was found only in H. attenuatus microgametocytes and that the genetic divergence of the two obtained H. erythrogravidus sequences was between 4.3% and 4.6% in cytb gene, which is assumed to differentiate these species.

In the present study, 50 strains of the two parasites were used to perform the phylogenetic analysis, a far larger number than that used by Mantilla et al. (2016) (n = 11). Thus, an average genetic distance of 0.83% was observed between the clade formed by the H. erythrogravidus lineages (clade II-a) and that formed by the H. coatneyi lineages (clade II-b), which is similar to the distance used by those authors to separate the species. According to Hellgren et al. (2007), species with a genetic distance above 5% are generally morphologically distinct. However, these species may present a lower genetic distance, provided they are clearly differentiated morphologically or that they present a high divergence in another gene that has phylogenetic resolution, in addition to cytb gene. Thus, there are examples of haemosporidian species whose cytb gene differs by less than 1%, as is the case of Haemoproteus jenniae and Haemoproteus siwa, whose cytb gene presents a genetic divergence of 0.6% and of 4%
in a segment of the apicoplast gene, in addition to being morphologically well differentiated (Levin et al., 2012). Another example is *Haemoproteus pallidus* and *Haemoproteus minutus*, which differ by only 0.7% in the complete sequence of cytb gene, but are clearly morphologically different (Hellgren et al., 2007). Considering the low genetic divergence and the fact that the morphological characteristics used for differentiation have been observed in several other species, it's possible that *H. erythrogravidus* is not a case of speciation.

According to the molecular, morphological and spatial distribution data, it was found that both *H. coatneyi* and *H. erythrogravidus* are parasitizing the order Passeriformes in Brazil. These two species have very similar gametocyte morphology and low genetic divergence, and may even be intraspecific variations of these two haemosporidian species. However, future phylogeny studies using mitochondrial genes and methods of computational demarcation of species may assist in the taxonomic delimitation of these two parasites.

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