Transovarial transmission and finding of *Trypanosoma rhipicephalis* in the hemolymph of *Rhipicephalus sanguineus sensu lato*¹

Transmissão transovariana e achado de *Trypanosoma rhipicephalis* na hemolinfa de *Rhipicephalus sanguineus sensu lato*

Yasmim Kaulich¹, Carolina Marotta Ribeiro², Jaqueline Rodrigues de Almeida Valim³, Juliana Ferreira dos Santos⁴, Thays Figueiroa dos Santos⁴, Claudia Bezerra da Silva⁴ & Adivaldo Henrique da Fonseca⁵

¹Veterinary Student, Scientific Initiation Scholarship FAPERJ, Awarded the Tokarnia Research Prize - SEMEV, IV. Departamento de Epidemiologia e Saúde Pública, Universidade Federal Rural do Rio de Janeiro – UFRRJ, Seropédica, RJ, Brasil
²Post Doctorate. Programa de Pós-graduação em Ciências Veterinárias, Universidade Federal Rural do Rio de Janeiro – UFRRJ, Seropédica, RJ, Brasil
³Veterinary Student, Scholarship. Departamento de Epidemiologia e Saúde Pública, Universidade Federal Rural do Rio de Janeiro – UFRRJ, Seropédica, RJ, Brasil
⁴Veterinary Student, Scientific Initiation Scholarship PIBIC. Departamento de Epidemiologia e Saúde Pública, Universidade Federal Rural do Rio de Janeiro – UFRRJ, Seropédica, RJ, Brasil
⁵Veterinary, Dr, Post Doctorate. Programa de Pós-graduação em Ciências Veterinárias, Universidade Federal Rural do Rio de Janeiro – UFRRJ, Seropédica, RJ, Brasil

Abstract

*Trypanosoma rhipicephalis* is a species isolated from *Rhipicephalus microplus* ticks collected from native bovine Seropédica, in the state of Rio de Janeiro. This study aimed to investigate the interaction of the tick *Rhipicephalus sanguineus s. l.* with *T. rhipicephalis* by in vitro artificial feeding. Eight females of *R. sanguineus s. l.* partially fed on rabbits. Tick infection was performed by an artificial feeding system using plastic tips for 12 hours. Canine blood used for feeding the ticks test group was previously infected with *T. rhipicephalis* 10³/mL. The hemolymph smear test was performed in all females after experimental infection. The daily posture was collected and organized in pools of each female per posture day. The eggs were divided into three groups, the first group for eggs PCR, the second for hatching and larval PCR, and the third group of eggs for isolation in cell culture. The evaluation of the presence of DNA in the macerated eggs of experimentally infected ticks showed two positive PCR samples. The evaluation of the presence of DNA in experimentally infected females showed all samples tested positive. For the hemolymph test, tick number 7 presented epimastigote developmental forms and amastigotes of *T. rhipicephalis*. Experimental infection by artificial feeding proved to be a suitable tool to study the interaction of *T. rhipicephalis* in *R. sanguineus s. l.* Ticks. The results show the transovarial transmission of *T. rhipicephalis* by *R. sanguineus s. l.*, as well as the interaction of the protozoan in the organism of this tick species.

Keywords: ticks, artificial feeding, trypanosomatids.

Resumo

*Trypanosoma rhipicephalis* é uma espécie isolada do carrapato *Rhipicephalus microplus* coletado de bovino nativo de Seropédica, estado do Rio de Janeiro. O presente estudo teve como objetivo investigar a interação do carrapato *Rhipicephalus sanguineus s. l.* com *T. rhipicephalis* por meio da alimentação artificial *in vitro*. Foram utilizadas 8 fêmeas de *R. sanguineus s. l.* parcialmente alimentadas em coelhos. A infecção dos carrapatos foi realizada por meio de um sistema de alimentação artificial utilizando ponteiras plásticas por um período de 12 horas. O sangue canino utilizado para alimentar os carrapatos do grupo teste foi previamente infectado com 10³ de *T. rhipicephalis/mL*. O teste de esfregaço de hemolinfa foi realizado em todas as fêmeas após a infecção experimental. A postura diária foi coletada e organizada em *pools* de cada fêmea por dia de postura. Os ovos foram separados em 3 grupos, sendo o primeiro grupo para a PCR dos ovos, o segundo para sua eclosão e PCR das larvas e o terceiro grupo para isolamento dos ovos em cultivo celular. A avaliação da presença de DNA no macerado de ovos dos carrapatos infectados...
experimentalmente mostrou duas amostras positivas na PCR. A avaliação da presença de DNA nas fêmeas infectadas experimentalmente mostrou todas as amostras do grupo teste positivas. Pelo teste de hemolinfa, o carrapato número 7 apresentou formas evolutivas epimastigota e amastigotas de *T. rhipicephalis*. A infecção experimental por meio da alimentação artificial mostrou-se uma ferramenta adequada para estudar a interação de *T. rhipicephalis* em *R. sanguineus s. l.* Os resultados comprovam a transmissão transovariana do *T. rhipicephalis* por *R. sanguineus s. l.*, assim como a interação do protozoário no organismo desta espécie de carrapato.

**Palavras-chave:** carrapato, alimentação artificial, tripanosomatídeos.

**Introduction**

Parasites of the genus *Trypanosoma* are unicellular and flagellated microorganisms belonging to the Trypanosomatidae family with wide geographical distribution. Hematophagous arthropods act as biological and mechanical vectors for different species of this family, infecting a wide variety of hosts, ranging from plants to animals, vertebrates and invertebrates (Hoare, 1972); (Morzaria et al., 1986).

Ticks, as blood feeders, are predisposed to the ingestion of a variety of parasites during feeding and have been proposed as vectors for *Trypanosoma* species (Morzaria et al., 1986); (Burgdorfer et al., 1973); and (Krige et al., 2019). Artificial tick feeding is an important tool for studying the transmission of pathogens in the absence of vertebrate hosts (Chabaud, 1950); (Valim et al., 2017). Cell culture of tick embryonic cells is an important tool for studying the interaction between cells of these arthropods and pathogens transmitted by them because it may help define the complex nature of the host–vector–pathogen relationship (Bell-Sakyi et al., 2007).

*Trypanosoma rhipicephalis* Marotta et al. (2018) is a species isolated from *R. microplus* ticks from native Seropédica cattle (Marotta et al., 2018). Little is known about its biological cycle; aspects related to possible pathogenesis and interaction with other species are unknown (Marotta et al., 2018).

The present study aims to investigate the interaction of *Rhipicephalus sanguineus sensu lato* and *T. rhipicephalis* by means of artificial feeding of ticks.

**Material and methods**

The experiments were performed at the Parasitic Diseases Laboratory of the Department of Epidemiology and Public Health of the Veterinary Institute (IV) of the Federal Rural University of Rio de Janeiro (UFRRJ), located in building 1 of the Veterinary Institute, in the municipality of Seropédica, Rio de Janeiro.

Eight females of *R. sanguineus s. l.* from the tick colony maintained at the Parasitic Disease Laboratory of the UFRRJ were partially fed on rabbits, following the methodology described by Valim et al. (2017). Two groups (G) were formed, one specimen was fed uninfected blood (G1 control), and seven were fed infected blood (G2 test). Tick infection was performed by an artificial feeding system using plastic tips (Figure 1) for a period of 12 hours following the procedures described by Valim et al. (2017). Blood used for artificial tick feeding was collected aseptically from the dog’s cephalic vein by the Vacutainer® system coupled to a 5 mL tube containing citrate anticoagulant. The material was labelled and stored under refrigeration at 4 °C for up to 24 hours.

To perform *Trypanosoma* blood infection, the dog’s blood serum complement system was previously inactivated so as not to cause reactions with the inoculated agent. For inactivation of the complement system, collected blood was centrifuged at 5000 rpm for 10 minutes. Then, the supernatant was kept for 40 minutes in a 56 °C water bath. The pellet was resuspended with PBS in a volume equivalent to the initial blood volume. Then two washes with PBS were performed. After the serum reached room temperature (26 °C), it was homogenized with the red blood cells. Canine blood used to feed G2 test ticks was previously infected with $10^8$ *T. rhipicephalis/mL*.

The hemolymph smear test was performed in all females after the experimental infection, being evaluated from the first DPI (the day after infection) until the posture, i.e., 13th DPI. The females used were kept frozen for later DNA extraction and PCR analysis. Hemolymph samples collected from the distal (tarsal and/or tibial) section of one or more telogin legs were deposited on a glass slide, identified, fixed for three minutes in methanol and stained by the Giemsa 10% method for 25 minutes.
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The daily posture was collected and organized in pools of each female per day of laying. Eggs were separated into three groups, the first for egg PCR, the second for hatching and larval PCR, and the third group for isolation of eggs in cell culture.

The DNA from the pool of macerated eggs, larvae, and infected females used in the experiment were extracted using the phenol–phenol/chloroform method previously described (Santolin et al., 2013). During each extraction battery, a blank sample was extracted along with the test samples as procedural control. A conventional Polymerase Chain Reaction (PCR) was performed to detect the presence of the 24S rDNA Trypanosomatid gene, following the protocol of the original article (Souto et al., 1999). Two positive samples (*Trypanosoma amblyomii*) and two negative samples (ultrapure water) were used during the PCR reaction.

**Results**

The authors were successful in infecting *R. sanguineus* s. l. with *T. rhipicephalis*, a species recently described in *R. microplus* by Marotta et al. (2018). The ticks *R. sanguineus* s. l. were infected by artificial feeding, and epimastigote and amastigote forms (Figure 2) of *T. rhipicephalis* were recovered from the hemolymph. The morphological forms observed were pleomorphic, such as those observed in in vitro cultures isolated from Rio de Janeiro native ticks, previously described by Marotta et al. (2018).

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**Figure 1.** Artificial feeding system of *Rhipicephalus sanguineus* s. l. using plastic tips with dog's blood infected with *Trypanosoma rhipicephalis*.

**Figure 2.** Epimastigote evolutionary forms of *Trypanosoma rhipicephalis* on tick hemolymph smear *Rhipicephalus sanguineus* s. l., Giemsa stained 10%, 100X magnification.
Evaluation of the presence of DNA in experimentally infected females showed all positive test group samples. By hemolymph test, tick number 7 showed epimastigote evolutionary forms of *T. rhipicephalis*.

The evaluation of DNA presence in the macerated eggs of experimentally infected ticks showed two positive PCR samples as well as the evaluation of the presence of DNA in macerated larvae, which presented one positive sample. The macerated eggs of the control G1 did not present positive bands by PCR. The PCR positive egg pool samples were from two G2 test females, one from the 4th to 6th DPI of tick number 4 and the other from the 4th to 6th DPI of tick number 6. The PCR positive larval pool sample also belonged to tick number 6 on the 6th to 8th DPI.

**Discussion**

This study describes for the first time how the tick *Rhipicephalus sanguineus s. l.* can be infected with *T. rhipicephalis* sing artificial feeding. The original description of *T. rhipicephalis* and preliminary studies conducted by Marotta et al. (2018) indicate its phylogenetic proximity to the KG1 species described by Thekisoe et al. (2007). *Trypanosoma KG1* was isolated from naturally infected *Haemaphysalis hystricis* ticks in Japan (Thekisoe et al., 2007). The morphological and phylogenetic proximity between *T. rhipicephalis* and *Trypanosoma KG1* seems to be related to the fact that it is also isolated from naturally infected ticks.

Another species with phylogenetic proximity to *T. rhipicephalis* is the species *T. caninum* isolated from an axenic culture of the intact skin of a domestic dog captured in different states of Brazil, including Rio de Janeiro (Madeira et al., 2014). Although the *T. caninum* vector is not yet known, it is possible that it is transmitted by ticks which then justifies the phylogenetic proximity between *T. rhipicephalis* and *T. caninum*. *T. rhipicephalis* has not been found to infect or cause any evidence of clinical signs in animals; there is a possibility that *T. rhipicephalis* has bovines as a mammalian host because tick *R. microplus* infests this species preferentially.

Although most Trypanosoma species are transmitted by hematophagous insects, ticks also appear to be likely vectors of some species of this genus (Madeira et al., 2014), (Kaufer et al., 2017), (Krige et al., 2019). In Brazil, a trypanosomatid with morphological characteristics similar to *Trypanosoma theileri* is naturally found in *R. microplus* and reported as nonpathogenic for cattle and usually transmitted by tabanids (Martins et al., 2008), (Ribeiro et al., 1988; Brum et al., 2012).

The *T. rhipicephalis* species in the present study showed biological characteristics compatible with other protozoa of the genus *Trypanosoma*, and evolutionary forms without visible free flagella in axenic cultures were seen. Recently, Barros et al. (2015) described aflagellar epimastigote forms in *T. caninum* confirmed by electron microscopy.

These results suggest that *R. sanguineus s. l.* may interact with the agent, considering its presence in the hemolymph. The present study also suggests the possible transovarial transmission of the protozoan by the tick *R. sanguineus s. l.* Transovarian transmission is a very significant mechanism for maintaining *T. rhipicephalis* in the environment and says a lot about its propagation and maintenance in the next generation of ticks.

*R. sanguineus* is a three-host life cycle species and is a competent disseminator of disease-causing microorganisms between infected and uninfected animal hosts (Dantas-Torres et al., 2013); (Sonenshine & Roe, 2014). The one-host tick *R. microplus* also takes three blood meals; however, feeding is restricted to the same host, making parasites ingested by one-host tick dependent on transovarial transmission for successful circulation (Kahl, 2018).

This study contributes a research tool for future studies on the possible elucidation of the *T. rhipicephalis* biological cycle and the involvement of the tick *R. sanguineus s. l.*

**Conclusion**

Experimental infection through artificial feeding proved to be an adequate tool to study the interaction of *T. rhipicephalis* in *R. sanguineus s. l.* The results indicate a possible transovarial transmission of *T. rhipicephalis* by *R. sanguineus s. l.*, as well as its interaction with the protozoan.
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