Field and experimental evidence of a new caiman trypanosome species closely phylogenetically related to fish trypanosomes and transmitted by leeches

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A R T I C L E   I N F O

Article history:
Received 28 July 2015
Received in revised form 25 September 2015
Accepted 16 October 2015

Keywords:
Crocodilian
Trypanosoma
Reptilian parasites
Phylogeny
Phylogeography
Leeches
Co-infection
South America

A B S T R A C T

Trypanosoma terena and Trypanosoma ralphi are known species of the South American crocodilians Caiman crocodilus, Caiman yacare and Melanosuchus niger and are phylogenetically related to the tsetse-transmitted Trypanosoma grayi of the African Crocodylus niloticus. These trypanosomes form the Crocodilian clade of the terrestrial clade of the genus Trypanosoma. A PCR-survey for trypanosomes in caiman blood samples and in leeches taken from caimans revealed unknown trypanosome diversity and frequent mixed infections. Phylogenies based on SSU (small subunit) of rRNA and gGAPDH (glycosomal Glyceraldehyde Phosphate Dehydrogenase) gene sequences revealed a new trypanosome species clustering with T. terena and T. ralphi in the crocodilian clade and an additional new species nesting in the distant Aquatic clade of trypanosomes, which is herein named Trypanosoma clandestinus n. sp. This new species was found in Caiman yacare, Caiman crocodilus and M. niger from the Pantanal and Amazonian biomes in Brazil. Large numbers of dividing epimastigotes and unique thin and long trypomastigotes were found in the guts of leeches (Haementeria sp.) removed from the mouths of caimans. The trypanosomes recovered from the leeches had sequences identical to those of T. clandestinus of caiman blood samples. Experimental infestation of young caimans (Caiman yacare) with infected leeches resulted in long-lasting T. clandestinus infections that permitted us to delineate its life cycle. In contrast to T. terena, T. ralphi and T. grayi, which are detectable by hemoculturing, microscopy and standard PCR of caiman blood, T. clandestinus passes undetected by these methods due to very low parasitemia and could be detected solely by the more sensitive nested PCR method. T. clandestinus n. sp. is the first crocodilian trypanosome known to be transmitted by leeches and positioned in the aquatic clade closest to fish trypanosomes. Our data show that caimans can host trypanosomes of the aquatic or terrestrial clade, sometimes simultaneously.

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1. Introduction

Flagellates of the genus Trypanosoma (Euglenozoa, Kinetoplastea, Trypanosomatidae) are obligate parasites of all vertebrate classes and are distributed into two major phylogenetic lineages: the Terrestrial clade composed of trypanosomes of mammals, snakes, lizards, crocodilians and birds and the Aquatic clade containing trypanosomes of aquatic leeches and aquatic (fishes) or semi-aquatic (chelonians, anurans and platypus) hosts and, oddly, a trypanosomone of chameleon (Stevens et al., 2001; Hamilton et al., 2005, 2007).

Trypanosomes infect Neotropical and Afro tropical crocodilians (Telford, 1995, 2009; Viola et al., 2008a; Fermino et al., 2013). Crocodilians of the family Alligatoridae predominate in the New World and are distributed in the genera Caiman, Paleosuchus and Melanosuchus, generically called caimans. The Afrotropical Crocodylidae originated in Australasia and after transoceanic dispersal...
American caimans. Their close relative to detect tissue-dwelling or scarce blood examination of peripheral blood. However, these methods may fail date were detected by culturing, standard PCR and microscopic (Hoare, 1929, 1931), and species from crocodilians were molecularly characterized: (Oaks, 2011) as vectors of trypanosomes of reptiles, including tsetse and semi-aquatic animals including (2008b). Leeches are known to transmit trypanosomes of aquatic (Martin and Desser, 1991; Siddall and Desser, 1992), snakes (Siddall and Desser, 1992) and platypus (Paparini et al., 2014). Revealed a number of trypanosomes from wildlife displaying blood (2007, 2008; Viola et al., 2008a,b; 2009; Cavazzana et al., 2010; Lemos et al., 2015). In addition, molecular studies the alligatorid trypanosomes present varied cultivation. The alligatorid trypanosomes present varied morphology in the blood and tissues of their hosts that could be the result of the inherent polymorphism of trypanosomes or concomitant infections with more than one species (Viola et al., 2008a; Fermino et al., 2013). The alleged polymorphisms of blood trypanosomes may also result from the occurrence of mixed infections that have been disclosed by molecular analyses of parasites directly from blood samples of anurans, fishes and mammals (Ferreira et al., 2007; Gu et al., 2007; Grybchuk-leremenko et al., 2014; Thompson et al., 2013; Lemos et al., 2015). In addition, molecular studies revealed a number of trypanosomes from wildlife displaying blood forms that are morphologically indistinguishable (Ferreira et al., 2007, 2008; Viola et al., 2008a,b; 2009; Cavazzana et al., 2010; Fermino et al., 2013; Lemos et al., 2015). These facts have prevented the correct appraisal of the whole assemblage of trypanosomes and the extent of their diversity and host and geographical ranges.

Nothing is known about the vectors of trypanosomes of South American caimans. Their close relative T. grayi is transmitted to African crocodiles by tsetse flies. A range of hematophagous flies serve as vectors of trypanosomes of reptiles, including tsetse flies for crocodiles (Hoare, 1929, 1931) and sand flies for lizards (Ayala, 1971; Ayala and McKay, 1971; Cristensen and Telford, 1972; Viola et al., 2008b). Leeches are known to transmit trypanosomes of aquatic and semi-aquatic animals including fishes (Letch, 1977; Karlsbaek et al., 2005; Khan, 1976; Hayes et al., 2014; Lemos et al., 2015), anurans (Martin and Desser, 1991; Siddall and Desser, 1992), snakes (Pessoa and Fleury, 1969; Chia and Miller, 1984), turtles (Woo, 1969; Siddall and Desser, 1992) and platypus (Paparini et al., 2014). Trypanosomes of anurans (Ayala, 1971; Martin and Desser, 1991; Siddall and Desser, 1992; Ferreira et al., 2008) and serpents (Chia and Miller, 1984; Viola et al., 2008b) are transmitted by leeches in aquatic environments and by insects such as sand flies and culicids in more terrestrial niches (Desser et al., 1975). Moreover, a survey performed using nested PCR suggested that terrestrial leeches could transmit trypanosomes of toads and marsupials in Australia (Hamilton et al., 2005). Therefore, flies and leeches could transmit the trypanosomes of the semiaquatic caimans, although no vectors have been identified to date.

In the present study, we surveyed for trypanosomes in blood and tissues samples from 122 specimens of South American caimans and the guts of 208 leeches collected from the mouths of caimans. Field surveys, morphological and experimental infections of caimans and phylogenetic analyses were employed to investigate life cycles, species diversity and phylogenetic relationships of crocodilians trypanosomes.

2. Materials and methods

2.1. Collection sites, caiman handling and blood sampling

Caimans were captured in the Amazonian (AM), Araguaia-Tocantins (TO), Paraguay-Paraná (PP) and Orinoco (OR) river basins. The collection sites are shown in Fig. 1 and Table 1. The capture of animals and all ensuing procedures were conducted as described previously (Fermino et al., 2013) according to the recommendations of IBAMA (the Brazilian Institute for the Environment and Renewable Natural Resources), permit Number 10080-2 and the protocol (108/2013) approved by the Committee on the Ethics of Animal Experimentation of the Institute of Biomedical Sciences, University of São Paulo.

Caiman blood samples and liver tissue preserved in ethanol (v/v) were stored in the Blood Sample Collection (BSC) and Tissue Sample Collection (TSC) of the Trypanosomatid Culture Collection of the University of São Paulo (TCC – USP). Total DNA was extracted from both blood and tissue samples as previously described using the traditional phenol-chloroform method. DNA samples from all trypanosome-positive crocodilian samples were also preserved in the TCC.

2.2. Collection and identification of leeches and survey for trypanosomes

Leeches were collected from the mouths of the caimans, and identified as Haementeria sp. by COI barcodes (Folmer et al., 1994). Leeches were immediately preserved in ethanol (v/v) or brought to the laboratory and kept for 15–20 days in plain water until they completely digested the ingested blood meal and became almost transparent. Then, the guts of the leeches were examined by microscopy. Leeches positive for trypanosomes were used for smears on glass slides, inoculation into culture tubes and DNA preparations using the phenol-chloroform method.

2.3. PCR amplification and phylogenetic analyses of SSU rRNA and gGAPDH gene sequences

The nested-PCR of SSU rRNA sequences (~987 bp) of trypanosomes from crocodilian blood samples was performed as described previously (Noyes et al., 1999). PCR amplification of the V7V8 region of the SSU rRNA and gGAPDH sequences from leech trypanosomes was performed as previously described (Borghesan et al., 2013). The nested PCR amplification of gGAPDH sequences was performed using the first round primers (GAPDH SF) 5′ GTG CGG GTK GTY GAC ATG AAC A3′ and (GAPDH SR) 5′ TTG GAG TCR TAG ATG CAG CT3′ and the second round internal primers (GAP SF) 5′ GTG AAG GCC CAC GGC AAC A3′ and (GAP SR) 5′ CGG AGG ATG YCC TTC ATG 3′. The two rounds of amplification (30 cycles each) were performed using PCR reaction mixtures and conditions described previously (Borghesan et al., 2013). Amplified DNA were cloned, and 10–20 clones were sequenced for each gene from each trypanosome sample. Sequencing of cloned amplified DNA allowed for the detection of mixed
trypanosome sequences in a single blood sample (Fermino et al.,
2013). The SSU rRNA and gGAPDH sequences determined in this
study were deposited in GenBank under the accession numbers
shown in Table 1.

For phylogenetic inferences using Parsimony (P), Maximum
Likelihood (ML) and Bayesian inference (BI) analyses, sequences
were aligned using CLUSTALX (Thompson et al., 1997) and
manually refined. Three alignments were created: A1, consisting of
the V7V8 region of the SSU rRNA gene (687 characters); A2, con-
taining the gGAPDH sequences (608 characters); and A3, consisting
of concatenated SSU rRNA and gGAPDH sequences (1,295 charac-
ters) using previous alignments including a large set of taxa for
guidance (Hamilton et al., 2005). The parsimony and bootstrap
analyses were performed using PAUP* version 4.0b10 software.

**Fig. 1.** Geographical origin of crocodilian trypanosomes included in the V7V8 SSU rRNA dendrogram inferred to compare the barcode sequences between the new and known trypanosomes from crocodilians and other species of aquatic and semi-aquatic hosts. The clade comprising *T. clandestinus* n. sp. nested into the Aquatic clade closely related to fish trypanosomes whereas sequences of the other new species formed the clade Cay03, which clustered with *T. terena*, *T. grayi* and *T. ralphi* in the Crocodilian Terrestrial clade. The host species and geographic origin and Genbank accession numbers of sequences from the crocodilian trypanosomes are shown in Table 1. Numbers at nodes are bootstrap support values >50% (P/ML) derived from 500 replicates.
for the gene data sets. The GTR model was used in individual
previously (Ferreira et al., 2008; Borghesan et al., 2013). The general
Table 1

| Sample* | Host     | River basin | GPS         | GenBank Accession numbers |
|---------|----------|-------------|-------------|---------------------------|
| T. clavata (BSC88) | C. yacare | PP | 19°57'57"01'W | KF768260 KF768285 |
| T. clavata (BSC64) | Haemaptera | PP | 19°57'57"01'W | KF768270 KR107951 |
| BSC91* | C. yacare | PP | 19°57'57"01'W | KF768252 KF768284 |
| BSC34k1*, BSC35c1*, BSC39, BSC46, BSC49c1*, BSC51c1* | M. niger | AM | 2’19’5’65’70’W | KF768253-8 – |
| BSC34k1* | C. crocodilus | AT | 7’32’5’49’22’W | KF768259 – |
| BSC387 | C. yacare | PP | 19°57'57"01'W | KF768261 – |
| BSC388 | C. crocodilus | AT | 7’66’5’49’29’W | KF768262 KF768286 |
| TSC03 | Haemaptera | PP | 19°57'57"01'W | – KF768287 |
| TSC05 | Haemaptera | PP | 19°57'57"01'W | KF768263 KF768288 |
| TSC06, TSC07, TSC08, TSC09, TSC10 | Haemaptera | PP | 19°57'57"01'W | KF768264-8 – |
| TSC02 | Haemaptera | AT | 7’66’5’49’29’W | KF768269 KF768289 |
| TSC65 | Haemaptera | PP | 19°57'57"01'W | KF768271 – |
| Clade terena | T. yacare (TCC621) | C. yacare | PP | 19°57’57’01’W | EU596252 EU596256 |
| BSC27 | O. tetraspis* | GB | 11°40’15’04’W | KF546505 – |
| BSC28 | C. crocodilus | OR | 6’83’3’67’69’W | KF546504 KF546518 |
| BSC33k1*, BSC51c2*, BSC57c1* | M. niger | AM | 2’19’5’65’70’W | – KF768290-2 |
| BSC50*, BSC53 | M. niger | AM | 2’19’5’65’70’W | – KF546519-20 |
| TCC1611 | C. yacare | PP | 19°57’57’01’W | KF546517 KF546503 |
| BSC56 | C. crocodilus | AM | 7’27’5’64’80’W | – KF768293 |
| TSC40 | P. trigonatus | AM | 3’80’3’61’73’W | KF768272 – |
| TSC43 | C. crocodilus | AM | 3’20’3’51’86’W | KF768273 – |
| Clade raphid | T. raphid (TCC1838) | M. niger | AM | 7’27’5’64’80’W | KF546512 KF546527 |
| BSC20* | C. yacare | PP | 19°57’57’01’W | KF546515 – |
| BSC37 | – | – | – | – KF768294 |
| BSC38 | M. niger | AM | 2’19’5’65’70’W | – KF768274 – |
| BSC40 | M. niger | AM | 2’19’5’65’70’W | KF768275 KF768295 |
| BSC42, BSC43c1* | M. niger | AM | 2’19’5’65’70’W | KF768276-7 – |
| BSC42k2* | M. niger | AM | 2’19’5’65’70’W | – KF768296 |
| BSC51* | M. niger | AM | 2’19’5’65’70’W | – KF456524 |
| BSC53 | M. niger | AM | 2’19’5’65’70’W | KF768278 – |
| BSC56 | M. niger | AM | 2’19’5’65’70’W | – KF546525 |
| BSC57c2* | M. niger | AM | 2’19’5’65’70’W | – KF768297 |
| BSC64k2* | C. crocodilus | AT | 7’32’5’49’22’W | KF546516 – |
| TCC624 | C. yacare | PP | 19°57’57’01’W | EU596253 EU596257 |
| TCC625, TCC1100 | C. yacare | PP | 19°57’57’01’W | KF546506 EU596259 |
| TCC1092 | C. yacare | PP | 19°57’57’01’W | EU596258 EU596254 |
| TCC1101, TCC1102, TCC1119 | C. yacare | PP | 19°57’57’01’W | KF546507-9 EU596261-3 |
| TCC1220 | C. yacare | PP | 19°57’57’01’W | KF546510 EU596255 |
| TCC1829 | C. crocodilus | AM | 7’27’5’64’80’W | KF546511 KF546521 |
| TCC1974 | C. yacare | PP | 19°57’57’01’W | KF546513 KF546522 |
| TCC2218 | C. crocodilus | AT | 7’32’5’49’22’W | KF546514 KF546523 |
| Clade Cay03 | – | – | – | – |
| BSC32, BSC35c2*, BSC35c2*, BSC39 | M. niger | AM | 2’19’5’65’70’W | – KF768298-301 |
| BSC43k2*, BSC44, | M. niger | AM | 2’19’5’65’70’W | KF768279-80 KF768302-3 |
| BSC46 | M. niger | AM | 2’19’5’65’70’W | – KF768304 |
| BSC49k2* | M. niger | AM | 2’19’5’65’70’W | KF768281 – |
| BSC50c3*, BSC51c4*, BSC52 | M. niger | AM | 2’19’5’65’70’W | – KF768305-7 |
| BSC53 | M. niger | AM | 2’19’5’65’70’W | KF768282 KF768308 |
| BSC63 | C. crocodilus | AM | 8’80’3’63’95’W | KF768283 – |
| BSC452 | C. crocodilus | OR | 6’83’3’67’69’W | – KR107954 |

*Identification code of the samples deposited at the Trypanosomatid Culture Collection of the University of São Paulo (TCC-USP): TCC, culture number; BSC, blood sample number; TSC, tissue sample number. "Mixed infections, “c” designates the clone number. South American river basins: AM, Amazonas; AT, Araguaia-Tocantins; PP, Paraná-Paraguay; OR, Orinoco. # GB, Guinea Bissau.

(Swofford, 2002) with 500 random sequence addition replicates followed by branch swapping (RAS-TBR). The ML analyses were performed using RAxML v2.2.3 (Stamatakis, 2006) as described previously (Ferreira et al., 2008; Borghesan et al., 2013). The general time reversible (GTR) model of nucleotide substitution with a proportion of invariable sites and gamma distribution was selected for the gene data sets. The GTR model was used in individual analysis of each gene as well as in the combined Bayesian analysis. The analysis was run for 1,000,000 generations with trees sampled every 100 generations using four chains and 25% of the early sample trees discarded as “burn-in.”

2.4. Experimental infection of caimans

Five farm-born Caiman yacare 8–10 months old and weighing ~260 g were kept indoors at 27 °C in large glass tanks of water with a land platform. Blood samples were collected upon the arrival of the animals and for 2 months at ~15 days intervals to evaluate by nested PCR whether the animals were free of trypanosomes prior to experimental infection. All procedures during the experimental period were performed in accordance with the protocol (#36781-1) approved by the IBAMA and the “Sistema de Autorização e Informação em Biodiversidade” (SISBIO).

Approximately 80 leeches were collected from wild Caiman...
The mouths of trypanosomes in a sample of 208 leeches randomly collected from et al., 1979; Khan et al., 1980; Leslie et al., 2011). We surveyed for trypanosomes, although they are very common and abundant ects (Table 1, Fig. 1).

We examined the number of leeches (TSC codes) from which trypanosomes could be detected using SSU rRNA and gGAPDH sequences. The amplification products were cloned and 10 clones from each sample were sequenced.

3. Results

3.1. New species disclosed by barcoding of caiman and leech trypanosomes using SSU rRNA and gGAPDH sequences

Preliminary tests showed that the nested PCR methods targeting both SSU rRNA and gGAPDH sequences could detect trypanosomes in the blood of caimans that tested positive or negative by blood microscopic examination and hemoculturing. Then, we examined 122 caiman blood samples and found trypanosome sequences in 40 caimans, yielding a prevalence of ~33%.

The analyses of V7V8 SSU rRNA and gGAPDH trypanosome sequences obtained from blood or tissue samples from 34 caimans revealed that 23 (68%) of the caimans presented sequences from a single species, whereas 11 (32%) presented mixed infections with two to four trypanosome species: T. terena and T. ralphi were detected in ~50% of the mixed infections. Sequences corresponding to two new trypanosome species were detected exclusively using the nested-PCR methods. Samples sharing a new sequence formed a clade (Cay03) that nested together with T. terena and T. ralphi in the previously described Crocodilian Clade (Fermino et al., 2013); the other new sequences formed the new Clandestinus clade, which were positioned within the Aquatic clade of Trypanosoma (Table 1, Fig. 1).

Leeches have not been reported to be vectors of crocodilian trypanosomes, although they are very common and abundant ectoparasites and vectors of other crocodilian parasites (Glassman et al., 1979; Khan et al., 1980; Leslie et al., 2011). We surveyed for trypanosomes in a sample of 208 leeches randomly collected from the mouths of Caiman yacare and Caiman crocodilus; a total of 120 leeches (~58%) were found to be positive for trypanosomes by microscopy. The number of leeches (TSC codes) from which trypanosomes were sequenced are listed in Table 1.

The leeches taken from Caiman yacare from The Pantanal were identified as Haementeria sp. by BLAST analyses of COI sequences (Genbank accession number: leech TSC404Le – KP972452). However, although a single sequence was obtained from 10 randomly selected leeches, the sequences obtained did not completely match any sequence in GenBank, suggesting that they may represent a leech species still lacking a DNA barcode. The leech sequence determined in this study diverged 17% from the closest relative COI sequences from Haementeria officinalis (Genbank accession number: JN850907).

Fresh gut samples of leeches removed from Caiman yacare exhibited intense flagellate proliferation. These flagellates always failed to grow when inoculated into the different culture media used to culture a range of trypanosomes including those from the clade Crocodilian (Viola et al., 2008a; Fermino et al., 2013) and anurans (Ferreira et al., 2007). Sequences of V7V8 SSU rRNA and gGAPDH identical to those of the Clandestinus clade from caiman blood samples were the only ones detected in all leeches examined (Table 1, Figs. 1 and 2).

3.2. Phylogenetic positioning of caiman and leech trypanosomes within the aquatic clade

In the dendrogram inferred using V7V8 SSU rRNA sequences from trypanosomes of caimans and leeches aligned with sequences of other trypanosome species, the crocodilian trypanosomes of the terrestrial clades Terena, Ralphi and Gayl (Fermino et al., 2013) plus the new clade Cay03 were separated from the Clandestinus clade, which included sequences from trypanosomes allied to fish trypanosomes (Fig. 1). The trypanosomes of the Crocodilian and Clandestinus clades were separated by ~5.0% V7V8 SSU rRNA divergence.

Inferred phylogenetic trees using gGAPDH sequences (Fig. 2) and concatenated V7V8 SSU rRNA and gGAPDH sequences (Supplementary Material) strongly supported the distribution of the caiman trypanosomes in the Terrestrial and Aquatic clades of Trypanosoma. The trypanosomes of the terrestrial Crocodilian clade, comprising T. terena, T. grayi and T. ralphi plus the new species provisionally referred to as Cay03, clustered with the trypanosomes from lizards, snakes, birds and mammals. The trypanosomes identified in the leeches always clustered with the caiman trypanosomes of clade Clandestinus, comprising highly homogeneous sequences that were strongly supported within the Aquatic clade (Fig. 2). These results enabled the description of the new species Trypanosoma clandestinus n. sp. in caimans and leeches.

The divergence of gGAPDH sequences between the five crocodilean trypanosome species ranged from 6.0% to 14%. The largest divergence separated T. grayi from T. clandestinus, and the shortest divergence separated T. ralphi from Cay03. T. clandestinus diverged 11.7%, 12.5% and 11% from T. terena, T. ralphi and Cay03, respectively. The divergence of gGAPDH sequences within each species varied from 0.5% in T. clandestinus to 2.1% in the more heterogeneous clade T. grayi. All gGAPDH sequences obtained from leeches (TSC 4–10 and TSC 62–65) were identical to the sequence of T. clandestinus (Table 1; Figs. 1 and 2).

Not a single trypanosome sequence of the Crocodilian clade was recovered from leeches. Moreover, T. clandestinus was phylogenetically more related to trypanosomes of fishes and placed closest to trypanosomes from Brazilian freshwater fishes than to any other reptilian trypanosome from the aquatic (Trypanosoma chelodinae) or terrestrial clade (Trypanosoma serpentinis, Trypanosoma cascavelli, Trypanosoma varani and T. sp. Gecko) (Fig. 1). The terrestrial crocodilian clade was sister to the main clade comprising trypanosomes from snakes, birds, lizards and mammals (Fig. 2).

3.3. Leeches as vectors of Trypanosoma clandestinus n. sp. and long-term infection of caimans

Although leeches are known to parasitize crocodilians, they had not been shown to transmit their trypanosomes. In this study, we achieved the experimental infection of caimans with trypanosomes from naturally-infected leeches recovered from the mouths of one
Caiman yacare captured in the Pantanal. SSU rRNA and gGAPDH sequences of the leech trypanosomes corresponded to the sequence previously determined for the leech TSC64 sample (Table 1).

We employed 30 randomly selected leeches for the experimental infestation of two young caimans. After exposure to the infected leeches, we collected blood samples from the caimans at ~15 day intervals for 5 months. Microhematocrit buffy coats show scarce trypanosomes 45 days after infection. Blood samples tested by PCR confirmed the infection of caimans (BSCs 386 and 387) with T. clandestinus (Table 1). The gGAPDH sequences from the infected caiman trypanosome were 99.8–100% similar to those from the leech TSC64. The control caiman and the caimans fed with macerated leeches remained negative by PCR throughout the experiment. Similarly, the ingestion of leeches taken from turtles naturally infected by trypanosomes did not infect turtles (Woo, 1969), suggesting that active leech inoculation of trypanosomes into the bloodstream is required for transmission.

![Phylogenetic tree (ML) based on gGAPDH sequences showing the Terrestrial and Aquatic clades of Trypanosoma and the positioning of T. clandestinus. The Crocodilian clade, which is formed by T. terena, T. ralphi, T. gray and Cay03 nests in the Terrestrial Clade whereas the Clandestinus clade comprising T. clandestinus nests in Aquatic clade. Typanosomatid genera other than Trypanosoma were used as outgroups in the phylogenetic trees (608 characters, Ln = −7611.8717). Numbers at nodes are bootstrap support (P/ML) >50% and Bayesian posterior probability >0.25 derived from 500 replicates.](image-url)
3.4. Diversity of crocodilian trypanosomes, mixed infections and lack of host-species restriction

The surveys using nested-PCR methods for the detection and barcoding of trypanosomes in the blood of four species of caimans disclosed trypanosome diversity and host-species ranges that are greater than those previously reported by hemoculturing. Two new trypanosome species were uncovered increasing to five the number of species (3) previously known (Fermino et al., 2013). Four species placed into the Terrestrial clades (Terena, Ralphi, Gray and Cay03) and one within the Aquatic clade (Clandestinus) of Trypanosoma, each one comprising one trypanosome species and its genotypes (Figs. 1 and 3).

In addition, results corroborated that diverse host species can successfully be used as hosts by different trypanosome species. The T. clandestinus clade comprises species found in Caiman yacare, Caiman crocodilus and M. niger. The new clade Cay03 was detected in M. niger and Caiman crocodilus. The hosts of T. terena include four of the six species of Brazilian alligatorids: M. niger, Caiman crocodilus, Caiman yacare and P. trigonatus. The list of species hosts for T. ralphi is similar, except for the absence of P. trigonatus (Table 1, Figs. 1 and 2). Thus, among all South American alligatorids, trypanosomes remain unreported only in Caiman latirostris and P. palpebrosus, which have not been thoroughly examined yet.

Altogether, results from this and previous phylogenetic studies revealed that the species of crocodilian trypanosomes lack host-restriction at species, genus and even family levels. In addition, a crocodilian species may host more than one trypanosome species.

Fig. 3. Proposed life cycle of T. clandestinus and its developmental and morphological features in caiman blood and leeches. Giemsa-stained blood smears showed blood trypanomastigotes of experimentally-infected Caiman yacare, and epi- and trypomastigotes found in the gut of one leech of the genus Haementeria sp. collected in the mouth of a wild Caiman yacare captured in the Pantanal wetland of Brazil. The caiman and the leech trypanosomes were molecularly identified as T. clandestinus. (a–c) epimastigotes; (b) epimastigote dividing by binary fission; (d, g) short trypomastigote; (e, f) long and thin trypomastigotes. Arrow points to the long and thin posterior extremity of very long and slender trypomastigotes. K, kinetoplast; N, nucleus; F, flagellum.
Indeed, one of the infected caimans harbored two or more trypanosomes. For instance, the *Caiman yacare* blood sample BSC29 yielded trypanosome sequences of the Ralphi and Clandestinus clades. An extreme example is offered by one *M. niger* (BSC51) that carried trypanosome sequences from the four Crocodilian subclades (Table 1, Figs. 1 and 2).

### 3.5. The distribution of trypanosome species in caimans of the South American river basins

This study shows that *T. clandestinus* is widespread in the Amazonian, Araguaia/Tocantins and Parana/Paraguay basins, but was not found in the Orinoco basin (Table 1, Fig. 1). Trypanosomes of clad CayO3 occurred in the Amazonian and Orinoco basins, but were not detected in the other basins including the Parana/Paraguay basin, even though large number of caimans were examined (Table 1, Fig. 1). We have previously shown that *T. terena* and *T. ralphii* were widespread throughout all of the river basins studied (Fermino et al., 2013). Therefore, the analysis of phylogeographical patterns did not support a strong spatial structure for the crocodilian trypanosomes, because most species occurred throughout the geographic distribution range of their hosts (Fig. 1).

### 3.6. Morphology and life cycle of *T. clandestinus* in caimans and leeches

Trypanosomes on blood smears of wild caimans could not be used for description of *T. clandestinus* trypanostigotes due to the common occurrence of mixed infections with two or more trypanosome species. Our source of blood trypanostigotes were the caimans experimentally infected with *T. clandestinus* from leeches harboring exclusively this species of trypanosome. Unfortunately, the caimans presented very low parasitemias rarely detected by microhematocrit, although the infection persisted for at least eight months. The few blood trypanostigotes examined (*N* = 5) in smears from microhematocrit buffy coats exhibited a serpentine body with a pointed posterior extremity, a central rounded nucleus and a posteriorly located small kinetoplast. The flagellum was short and the undulating membrane was convoluted (Fig. 3). The blood forms of *T. clandestinus* were smaller and thinner than the wider trypanostigotes of *T. terena* and *T. ralphii* (Fermino et al., 2013).

Leeches of the genus *Haementeria* are common ecto-parasites of alligatorids. Because the leeches were rather small the examination of the segments of their digestive tract was not attempted and gut contents were wholly mixed for examination. The leeches removed from the caimans were examined after the engorged blood was completely digested, thus ensuring that the flagellates observed ~20 days after leeches collection developed in their guts.

As expected for a cyclically transmitted parasite, most leeches were heavily infected with epimastigotes that showed many dividing forms, while some leeches examined after the 20th day presented large numbers of trypanostigotes (Fig. 3). The morphology of the flagellates found in the gut contents of one *Haementeria* leech (TSC09) removed from a *Caiman yacare* that had only sequences (14 clones) identical to that of *T. clandestinus* was selected for the illustration of the trypanosomes found in leeches (defined as type material for species description). The epimastigotes (*N* = 25) were generally slender, with body lengths varying from 30 to 50 μm (average 42 μm) and widths varying from 2.7 to 4.6 μm (average 3.9 μm). The spherical nuclei and kinetoplasts were close together and located near the middle of the body (Fig. 3a–c) and the flagellum length varied from 2.3 to 6.8 μm (average 5.5 μm). The undulating membrane can be narrow or markedly frilled in the widest forms (Fig. 3b). The polymorphism of the trypanostigotes was much more accentuated than that of the epimastigotes. Short trypanostigotes (*N* = 15) measuring 23–42 μm in length (average 38 μm) and 2.2 to 2.5 in width (average 2.1 μm) displayed nuclei and kinetoplasts in general displaced towards the anterior end (Fig. 3d, g). Long trypanostigotes, extremely slim at both extremities were abundant. These forms (*N* = 25) measured 60–107 μm long (average 77 μm) and 0.7–2.5 μm wide (average 1.25 μm) and contained small and elongated nuclei and punctual kinetoplasts located at the center of the body, but not in close proximity to one another. In these thin trypanostigotes the free-flagellum ranged 11–20 μm (average 13.3 μm) (Fig. 3e, f).

Although available data of *T. clandestinus* in leeches and experimentally infected caimans are not sufficient to explain the entire development of this species in its vertebrate host and vector, overall results allowed us to deduce the following life cycle for *T. clandestinus*: *Haementeria* sp. leeches ingest blood containing trypanostigotes upon piercing the caiman buccal mucosa. In the gut of leeches, blood trypanostigotes differentiate into epimastigotes that proliferate and progressively transform into short and long trypanostigotes. Probably, caimans become infected through the bite of leeches carrying trypanostigotes in their proboscis. Although few trypanostigotes were found in the peripheral blood of infected caimans, each leech ingested a considerable volume of caiman blood yielding the high prevalence of *T. clandestinus* in leeches.

### 4. Taxonomic summary

#### 4.1. New species description

Phylum Euglenozoa Cavalier-Smith 1981: Class Kinetoplastea Honigberg 1963: Order Trypanosomatida (Kent 1880) Hollande 1982. Genus *Trypanosoma* Gruby 1843. *Trypanosoma clandestinus* Teixeira and Camargo n. sp.

**Type Material:** Hapantotype: trypanostigotes on glass slides of blood smears of blood sample BSC386 from *Caiman yacare*, and epi- and trypanostigotes on smears of gut contents from the leech *Haementeria* sp. (sample TSC 64). The slides are deposited at the Trypanosomatid Culture Collection of the University of Sao Paulo (TCC~USP). Brazilian leeches infected with *T. clandestinus* are cryopreserved in liquid nitrogen. Paratypes: trypanostigotes on glass-slide blood smears of *Caiman yacare* blood sample BSC387 stored at TCC-USP. **Vertebrate type host:** *Caiman yacare* (Crocodylia, Alligatoridae), Vertebrate additional hosts: *Caiman crocodilus* and *M. niger*. **Invertebrate type host:** *Haementeria* sp. (Hirudinea, Rhynchobdellida), identified by COI sequences deposited in GenBank (KP972452). **Habitats:** blood of caimans and digestive tube of leeches. **Type locality:** Miranda River (520° 14’ W 56° 22’), Mato Grosso do Sul, the Pantanal, Brazil. **Additional localities:** AM and AT river basins (Table 1). **Morphology:** Caiman blood trypanostigotes and leech epi- and trypanostigotes shown in Fig. 3. **Molecular diagnosis:** DNA sequences deposited in GenBank of trypanosomes from *Caiman yacare* (BSC386) of the genes V7V8 SSU rRNA (KP768285) and gGAPDH (KP768260) and from trypanosomes of the leech *Haementeria* sp. (TSC64), V7V8 SSU rRNA (KR107951) and gGAPDH (KP768270). **Etymology:** clandestinus, Lat. n. m.: hidden, concealed, occult.

**Taxonomical Comments:** The lack of host specificity of caiman trypanosomes and the existence of similar blood trypanostigotes shared by different species, intra-species polymorphisms and mixed infections do not permit the assumption that the trypanosomes observed in blood smears belong to a single or multiple species. Therefore, blood trypanostigotes from naturally infected caimans could not be the type material, unless they are proven to belong to a single species. In the absence of derived cultures, it is
necessary to verify whether only one trypanosome sequence is present through sensitive PCR and sequencing of a large number of cloned sequences from each blood sample. Alternatively, flagellates from vectors (epi- and trypomastigotes) exhibiting sequences of a single trypanosome species and blood trypomastigotes from animals experimentally infected with molecularly authenticated trypanosomes from the vectors can serve as type material. In this paper, we used the last two procedures to validate the new species because cultures of *T. clandestinus* are not available.

In addition, sequence divergences separating *T. terena*, *T. ralphi* and Cay03 were large enough to permit considering Cay03 as a new species within the Terrestrial clade. However, we still do not have a proper type material to taxonomically validate Cay03 as a new species.

5. Discussion

Recent molecular studies have uncovered a vast diversity of wildlife parasites within the genus *Trypanosoma*. However, the molecular phylogenetic studies of trypanosomes of reptiles are still limited to a few species of turtles (Jacques et al., 2001), lizards and snakes (Hamilton et al., 2007; Viola et al., 2008b, 2009) and crocodilians (Viola et al., 2008a; Fermino et al., 2013). In this study, surveys of trypanosomes in blood samples from South American caimans using sensitive nested-PCR methods revealed an unexpected high prevalence (33%) of infection and common mixed infections by two or more trypanosome species (45%). The phylogenetic positioning of caiman trypanosomes using SSU rRNA and gGAPDH sequences increased our knowledge about the species richness of crocodilian trypanosomes and raised the number of molecularly characterized species from three to five. Two phylogenetically-supported new trypanosome species were identified. One species was designated *T. clandestinus* n. sp. and molecularly and morphologically characterized in the present study. This species was shown to be transmitted by leeches and to cluster within the Aquatic clade of *Trypanosoma*. Altogether, the information concerning the development and morphological features in leeches and in blood of caimans experimentally infected with this trypanosome species through the bite of leeches, permitted us to outline the probable life cycle of *T. clandestinus*.

Due to the frequent occurrence of mixed infections and the large range of host species, it is impossible to identify a new species of crocodilian trypanosome based on the morphology of blood trypanosomes and host of origin. Phylogenetic analyses are obligatory and, preferably, should be preceded by culturing and morphological characterization of the candidate new species. Before this study, the use of these combined approaches prompted the description of two species of crocodilian trypanosomes (*T. terena* and *T. ralphi*) in South American alligatorids (Fermino et al., 2013). Similarly, cultures derived from tsetse flies and *C. niloticus* allowed the molecular characterization of *T. grayi* (Stevens et al., 2001; Hamilton et al., 2009; Fermino et al., 2013).

In contrast with *T. terena*, *T. ralphi* and *T. grayi*, which are commonly detected by microscopy, hemocultures and conventional PCR, *T. clandestinus* remains hidden in its crocodilian host and is exclusively detected by highly sensitive nested PCR methods. Contrasting with the very low parasitemia of caiman hosts, *T. clandestinus* can be easily detected by microscopy and conventional PCR in leeches of the genus *Haementeria*, which are ecto-parasites often found in large clumps in the buccal cavity of crocodilians. The abundant trypanosomes multiplying in the digestive tract of *Haementeria* sp. taken from caimans did not grow in culture under any of the conditions we attempted but offered sufficient material for morphological and molecular analyses, and for experimental infections of caimans. *T. clandestinus*-carrying leeches were used to infect farm-raised *Caiman yacare*. Blood trypanosomes from the infected caimans yielded V7V8 SSU rRNA and gGAPDH sequences identical to those determined for the trypanosomes multiplying in leeches. Thus, results demonstrated that leeches were capable of transmitting *T. clandestinus* to *Caiman yacare*.

*T. clandestinus* is the first crocodilian trypanosome known to be transmitted by leeches and placed into the Aquatic clade. This finding is in agreement with the role of aquatic leeches as vectors of trypanosomes of fishes, anurans, and the platypus (Martin and Desser, 1991; Siddall and Desser, 1992; Hayes et al., 2014; Paparini et al., 2014). *T. clandestinus* was the sole trypanosome identified in the gut of *Haementeria* sp., even in leeches taken from caimans infected with other two or more trypanosome species. This finding support our hypothesis that the crocodilian trypanosomes of the Terrestrial clade are transmitted by hematophagous flies, similarly to some trypanosomes of anurans, lizards and snakes (Ayala, 1971; Ayala and McKay, 1971; Martin and Desser, 1991; Siddall and Desser, 1992; Ferreira et al., 2008; Viola et al., 2008b, 2009) and *T. grayi* of African crocodiles (Hoare, 1929, 1931).

In this study, phylogenetic analyses evidenced, for the first time, that crocodilians of distinct species harbor phylogenetically distant trypanosomes of the two major *Trypanosoma* clades. The Aquatic clade harbors *T. clandestinus* transmitted by leeches and highly phylogenetically related to fish trypanosomes. The Terrestrial clade harbors *T. grayi* transmitted by tsetse flies plus *T. terena*, *T. ralphi* and Cay03 of unknown vectors. In contrast, all anuran trypanosomes nested in the Aquatic clade even though they are transmitted by leeches or by flies (Ferreira et al., 2007, 2008). It will be interesting to see whether trypanosomes from aquatic snakes transmitted by aquatic leeches (Pessoa and Fleury, 1969; Chia and Miller, 1984) cluster in the Aquatic clade or in the Terrestrial lizard-snake clade of trypanosomes transmitted by sand flies (Viola et al., 2009). To date, a chameleon trypanosome of unknown vector is the only trypanosome from lizard placed in the Aquatic clade (Stevens et al., 2001; Hamilton et al., 2007).

Here, we demonstrated that different trypanosome species co-infected different species of crocodilians across South American river basins. The identification of new hosts and new trypanosomes permits us to anticipate that the species richness of crocodilian trypanosomes is much larger than presently known. However, the task of effectively cataloguing the species richness will have to face not only the conceptual issue of species delimitation (Adams, 2001; De Queiroz, 2007), but also the operational problems of new species descriptions. Collecting blood samples and leeches from wild crocodilians in their natural habitats can be very difficult. Moreover, the existence of species such as *T. terena* and *T. ralphi* that share morphologically indistinguishable blood and culture forms (Viola et al., 2008a; Fermino et al., 2013) or non-culturable species as *T. clandestinus* adds difficulties to the appreciation of the trypanosome diversity. *T. clandestinus* does not belong to the cryptic category of parasites (i.e., two or more species morphologically perceived as a single species), but to a category of parasites that go unnoticed in host blood samples by common methods of detection/identification of trypanosomes. Any inventory of parasite biodiversity and the proposition of well-supported hypotheses about their evolutionary histories depends on the sensitive detection and correct identification of species. The enormous realm of wildlife parasites remains largely underexplored and the occurrence of cryptic, clandestine and misclassified parasite species further complicates the prospect of such an inventory (Criscione et al., 2005; Bickford et al., 2007; De León and Nadler, 2010; Detwiler et al., 2010; Nadler and De León, 2011; Poulin et al., 2011b; Bray and Cribb, 2015). Further studies are required to assess
whether the specificity of the crocodilian trypanosomes is restricted to closely related crocodilian species or whether they can exploit distantly related hosts of other taxa across their broad geographic range (Poulin et al., 2011a). Caimans are known to be generalist predators that feed on diverse prey including fish, turtles, anurans and other aquatic and terrestrial animals. Animals sharing ecological niches and serving as food are known to share parasites with crocodilians (Junker and Boomker, 2006; Tellez and Nifong, 2014). In this context, it is worth emphasizing that T. clandestinus was placed closest to fish trypanosomes within a major assemblage of parasites transmitted by aquatic leeches. It is known that fishes and anurans play a role of paratenic hosts of Hepatozoon caimani, a highly prevalent caiman hemoparasite transmitted by culicids (Viana et al., 2012; Pereira et al., 2014). Therefore, it is possible that fish and other aquatic vertebrates share trypanosomes with caimans, including T. clandestinus, due to host-switching mediated by aquatic leeches and predation by crocodilians. We are currently examining trypanosomes of fishes and anurans that serve as food for the caimans in the Pantanal to investigate the role of ecological fitting favoring host-switching of trypanosomes sharing ecological niches. So far, we have not found trypanosomes identical to T. clandestinus in anurans (Ferreira et al., 2017) sharing ponds with Caiman variegatus. However, preliminary phylogenetic analysis revealed partial SSU rRNA sequences closely related to T. clandestinus, but not identical, in blood samples of fishes (BSCs 97, 98, 100) from The Pantanal (Fig. 1) and other Brazilian hidrographic basin (Lemos et al., 2015).

In addition, misclassified trypanosome species may obscure the etiology of infections of wild and farmed caimans. Understanding vectors and parasite species-specific associations have implications for conservation, management and production purposes. Molecular approaches have only recently been used to investigate helminthiasis of endangered species or farmed crocodilians (Zhao et al., 2015; Lott et al., 2015). There are no studies addressing the pathological effects of trypanosome infections in caimans. We did not find any obvious sign of illness in the caimans naturally and chronically infected (8 months) with T. clandestinus. However, the demonstration of trypanosomes in blood and tissue implants of kidney and lung of wild crocodilians (Lainson, 1977; Viola et al., 2008a) deserve further investigations. Trypanosomes in general have been considered non-pathogenic in wildlife, although studies are scarce and in general superficial. Recently, studies about trypanosomes infecting endangered Neotropical primates and Australian marsupials started producing relevant data concerning wildlife health and conservation (Maia da Silva et al., 2008; Thompson et al., 2013; Botero et al., 2013; Austen et al., 2015).

Results from the present study disclosed an increasing species richness, enlarged host/parasite geographical range, aquatic and terrestrial cycles of transmission and new vectors (leeches) of crocodilian trypanosomes. The diversity and phylogenetic relationships of these trypanosomes likely have been shaped by transoceanic dispersion and ecological fitting with multiple events of host-switching during the long shared evolutionary histories of trypanosomes, crocodilians of different families and leeches.

Acknowledgments

We thank Dr. Miguel U. Trefault Rodrigues for the caiman identifications and for providing some archived blood samples of caimans. We also thanks several students and anonymous field workers who helped us in the capture of caimans, fishes and anurans. This work was supported by grants from the Brazilian Agencies CNPq (PROTAX, PROAFRICA) and CAPES (PNPBP, PNPD). We acknowledge the Brazilian Ministry for Science, Technology and Innovation (MCTI) for field support through the Mamirauá Institute for Sustainable Development (IDSM). Bruno R. Fermino and Priscila Soares are PhD students supported by CAPES and CNPq, respectively. Robson C. Ferreira is a Post-doctoral student sponsored by PNPD-CAPES.

Conflict of interest

We confirm that there are no financial or personal interests that might lead to a conflict of interest.

Source of funding

All sources of funding were led in the Ms. There are none extra funding received for this work.

Appendix A. Supplementary data

Supplementary data related to this article can be found at http://dx.doi.org/10.1016/j.ijppaw.2015.10.005.

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