Shared species of crocodilian trypanosomes carried by tabanid flies in Africa and South America, including the description of a new species from caimans, *Trypanosoma kaiowa* n. sp.

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**Abstract**

**Background:** The genus *Trypanosoma* Gruby, 1843 is constituted by terrestrial and aquatic phylogenetic lineages both harboring understudied trypanosomes from reptiles including an increasing diversity of crocodilian trypanosomes. *Trypanosoma clandestinus* Teixeira & Camargo, 2016 of the aquatic lineage is transmitted by leeches to caimans. *Trypanosoma grayi* Novy, 1906 of the terrestrial lineage is transmitted by tsetse flies to crocodiles in Africa, but the vectors of Neotropical caiman trypanosomes nested in this lineage remain unknown.

**Results:** Our phylogenetic analyses uncovered crocodilian trypanosomes in tabanids from South America and Africa, and trypanosomes other than *T. grayi* in tsetse flies. All trypanosomes found in tabanids clustered in the crocodilian clade (terrestrial lineage) forming six clades: Grayi (African trypanosomes from crocodiles and tsetse flies); Ralphi (trypanosomes from caimans, African and Brazilian tabanids and tsetse flies); Terena (caimans); Cay03 (caimans and Brazilian tabanids); and two new clades, Tab01 (Brazilian tabanid and tsetse flies) and Kaiowa. The clade Kaiowa comprises *Trypanosoma kaiowa* n. sp. and trypanosomes from African and Brazilian tabanids, caimans, tsetse flies and the African dwarf crocodile. *Trypanosoma kaiowa* n. sp. heavily colonises tabanid guts and differs remarkably in morphology from other caiman trypanosomes. This species multiplied predominantly as promastigotes on log-phase cultures showing scarce epimastigotes and exhibited very long flagellates in old cultures. Analyses of growth behavior revealed that insect cells allow the intracellular development of *Trypanosoma kaiowa* n. sp.

**Conclusions:** Prior to this description of *Trypanosoma kaiowa* n. sp., no crocodilian trypanosome parasitic in tabanid flies had been cultured, morphologically examined by light, scanning and transmission microscopy, and phylogenetically compared with other crocodilian trypanosomes. Additionally, trypanosomes thought to be restricted to caimans were identified in Brazilian and African tabanids, tsetse flies and the dwarf crocodile. Similar repertoires of trypanosomes found in South American caimans, African crocodiles and tabanids from both continents support the recent diversification of these transcontinental trypanosomes. Our findings are consistent with trypanosome host-switching likely mediated by tabanid flies between caimans and transoceanic migrant crocodiles co-inhabiting South American wetlands at the Miocene.

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Keywords: Tabanids, Tsetse flies, Crocodile, Caiman, Evolution, Taxonomy, Morphology, Transoceanic dispersal

Background

Trypanosomes are ubiquitous parasites of all classes of vertebrates, from fishes to mammals, including a few species parasitic in humans [1–4]. More than 70 trypanosome species have been described in reptiles (lizards, snakes, turtles and crocodilians) based on their morphology, and host of origin [2]. Until recently, Trypanosoma grayi Novy, 1906 [5, 6] and Trypanosoma cecili Lainson, 1977 [7] were the only trypanosomes described in African crocodiles and Brazilian caimans, respectively. Previous phylogenies, including T. grayi from tsetse flies and crocodiles [3, 8, 9] and trypanosomes from South American caimans [10–12] enabled the description of three additional species: Trypanosoma terena Teixeira & Camargo, 2013, Trypanosoma ralphi Teixeira & Camargo, 2013, and Trypanosoma clandestinus Teixeira & Camargo, 2016.

The genus Trypanosoma is composed of two major phylogenetic lineages, in line with the main habitats of their vertebrate hosts, the terrestrial and the aquatic lineage [3, 4]. Trypanosomes of fishes, anurans, cheloniens, platypus and some lizards [3, 4, 10, 11, 13–19] plus T. clandestinus of caimans [12] cluster together in the aquatic lineage. Trypanosomes of land mammals, bats, birds, snakes and lizards, nest in the terrestrial lineage, together with the crocodilian trypanosomes T. grayi, T. terena and T. ralphi [4, 9–12, 20–23].

In Africa, trypanosomes of the terrestrial lineage occur in species of the family Crocodylidae: Crocodylus niloticus Laurenti, Mecistops cataphractus Cuvier, and Osteolaemus tetraspis Cope [5, 6, 12, 24]. In South America, trypanosomes are hosted by species of Alligatoridae (subfamily Caimaninae): Caiman yacare Daudin, Caiman crocodilus Linnaeus, Melanosuchus niger Spix, and Paleosuchus trigonatus Schneider [7, 10–12]. The extant species of the Crocodylinae and Caimaninae appeared on earth after an episode of mass extinction of crocodylians during the global cooling that extended from the mid-Miocene to the Pliocene [25–28]. The genus Crocodylus Laurenti originated in Australasia during the mid-Miocene (8–13 mya) and through transoceanic dispersal reached South America and Africa, although the exact dispersion route remains uncertain [26]. Fossil records from the mid- to late Miocene [27, 28] indicate that ancestors of caimans and crocodiles lived in sympatry in northern South America, a food-rich waterlogged ensemble of lowlands that gave origin to the contemporary Amazonia hydrographic system [29–31]. We previously hypothesised that the encounter of crocodiles with caimans created the opportunity for host-switching of trypanosomes, likely mediated by hematophagous invertebrates [10, 11]. The vectors of the African T. grayi are tsetse flies [5, 6, 32]. Leeches, that are common ectoparasites of crocodilians, are known vectors of T. clandestinus of caimans [12]. However, these aquatic leeches have never been reported to carry trypanosomes of the crocodilian clade. Thus, unknown hematophagous invertebrates other than leeches should be vectors of caiman trypanosomes of the terrestrial lineage. There is a range of contemporary hematophagous invertebrates that thrived on earth during the Cretaceous sharing habitats with crocodilians: leeches, ticks, mosquitoes, and flies of many families such as the Glossinidae, Psychodidae, Ceratopogonidae, and Tabanidae [33–36].

Tabanids cyclically transmit Trypanosoma theileri Laveran, 1902 of domestic and wild ruminants, carry other trypanosomes of ungulates, and possibly transmit trypanosomes to marsupials in Australia [1, 22, 37–41]. Many species of tabanids are generalist bloodsuckers, and some species feed preferentially on crocodiles, monitor lizards, snakes, chelonians and birds [42–46]. Given that some tabanid species prefer caiman blood as a food source [44], we searched for trypanosomes in tabanid flies captured near rivers and lakes inhabited by caimans or crocodiles. In this study, we report results from field surveys, morphological and behavioral features, and phylogenetic analyses of trypanosomes from tabanids, tsetse flies, caimans and crocodiles. Data obtained enabled the description of a new trypanosome species parasitic in tabanid flies and Ca. yacare.

Methods

Collection of tabanids and tsetse flies

We collected tabanids while they were feeding on caimans (Ca. yacare) captured in the Miranda River, Mato Grosso do Sul state, in the Pantanal biome, which is a vast wetland in central Brazil and home to a considerable population of caimans (Fig. 1). Two adult caimans were captured [11, 12] and immobilized for about 3 h to serve as baits for tabanids. Tabanids were dissected and had their gut contents microscopically examined for trypanosomes according to procedures adopted for tsetse fly dissection detailed in a previous study [47]. Positive gut contents were inoculated into culture media and smeared on glass slides, which were fixed with methanol and stained with Giemsa.

We also collected tabanids near rivers and lakes inhabited by caimans or crocodiles. Tabanid flies were
collected manually in the Amazonia biome, Tabajara Forest Reserve, Rondonia State, Brazil (8°56′S, 62°03′W), on the margins of the Ji-Paraná River (Fig. 1). In addition, tabanids and tsetse flies were caught on borders of the Zambezi River (Chupanga, district of Sofala) and Lake Jurema in the Gorongosa National Park in Mozambique, Lake Chamo in the Nech Sar National Park in Ethiopia, and Lake Edward in the Queen Elizabeth National Park in Uganda (Fig. 1). Flies from Amazonia and Africa could not be examined by microscopy and were preserved intact in ethanol. We confirmed the identification of selected specimens as representatives of the morphological diversity of tabanid, and tsetse flies by \( \text{cox1} \) barcoding [48].

**Morphology and behavior of trypanosomes in vitro**

Guts of tabanids caught in the Pantanal biome that were found to be microscopically positive for trypanosomes were seeded in cultures of Hi-5 insect feeder cells (\( \text{Trichoplusia} \) sp.) overlaid with TC100 (= Grace’s) medium containing 10% FBS, and incubated at 25 °C [11, 12]. Flagellates were subsequently cultured in LIT (Liver Infusion Tryptose) medium supplemented by 10% FBS at 25 °C. The growth behavior of trypanosomes co-cultivated with Hi-5 cells was investigated in the supernatant and in cells attached to coverslips as we described previously [49]. Hi-5 cells in TC100 (~10^4 cells/well) were seeded onto 24-well plates containing glass coverslips, incubated for 24–48 h, and then seeded with 10² flagellates/well. After ~8 h, cultures were washed twice with PBS (phosphate buffered saline) and incubated with fresh TC100 medium. After 1, 3, 5, 7 and 10 days, cells were washed three times with PBS, and the cells fixed with methanol and stained with Giemsa.

Hi-5 cells seeded with trypanosomes were examined daily by inverted phase-microscopy. Examination of fixed flagellates stained with Giemsa in coverslips and mounted onto glass slides were performed by light microscopy. Both isolates TCC1611 and TCC2918 were evaluated regarding behavior in Hi-5 cultures, and experiments with TCC2918 were repeated three times. Flagellates from the supernatant of Hi-5 cultures were fixed with methanol on glass slides and stained with Giemsa.

Transmission (TEM) and scanning (SEM) electron microscopical analyses

For SEM, trypanosomes from log-phase Hi-5 cultures were washed three times with PBS, fixed in 2.5% glutaraldehyde plus 1% formaldehyde in 0.1 M cacodylate buffer (pH 7.2) at room temperature overnight, and processed
using the protocols usually employed for trypanosomes [11, 12, 17, 18, 20, 23, 49, 50]. Briefly, fixed flagellates were adhered to 0.1% poly-L-lysine-coated coverslips, post-fixed in the dark in 1% osmium tetroxide in 0.1 M cacodylate buffer, pH 7.2, for 30 min, dehydrated in an ascending ethanol series, dried in critical point drier equipment CPD 030 (Leica Microsystems, Wetzlar, Germany) and covered with gold in sputtering equipment SCD 050 (Leica Microsystems). A FEI Quanta 250 (FEI Company, Hillsboro, United States) was used for SEM. For TEM, trypanosomes were fixed and dehydrated as above for SEM, embedded overnight in epoxy resin, trimmed, and cut in thin sections (80–100 nm). Sections were stained with aqueous 2% uranyl acetate and lead citrate before being examined using a LEO 906E at 80 kV (Zeiss, Jena, Germany). Images were captured by a CCD camera MegaView III.

PCR amplification, sequencing and phylogenetic analyses of SSU rDNA and gGAPDH sequences
DNA from caiman blood and gut contents of tabanid and tsetse flies preserved in ethanol as previously described [12, 47]. Briefly, blood and gut samples were centrifuged to remove ethanol, dried at 37 °C, resuspended in 200 µl of DIGSOL solution (containing proteinase K and RNase), and incubated at 37 °C overnight. DNA was then precipitated with 400 µl of ammonium acetate (4 M), washed in ethanol, dried, and resuspended in 50–100 µl of TE.

DNA from cainan blood samples and the guts of tabanid and tsetse flies were submitted to nested-PCRs for amplification of V7V8 SSU rDNA (~900 bp) and gGAPDH (~680 bp) sequences, as previously described [12]. Sequences from 5–7 clones of each amplified sample were determined, screened for chimera by the RDP4 package, and those that were representatives of the whole genetic diversity were deposited in GenBank (Additional file 1: Table S1). For phylogenetic inferences, newly generated and published sequences were aligned with Clustal W and MUSCLE programs. We created three alignments: (i) A, consisting of V7V8 rDNA sequences of all trypanosomes found in crocodilians, including *T. clandestinus* and trypanosomes of the aquatic clade; (ii) B, including gGAPDH sequences, determined in the present and previous studies, restricted to trypanosomes of the clade crocodilian (several isolates from each clade), plus sequences of closely related trypanosomes from tapisrs (Brazil) [21] and rhinoceros (Sumatra) available on GenBank; (iii) C, consisting of concatenated V7V8 SSU rDNA and gGAPDH sequences from isolates representative of each species/clade of the major crocodilian clade, and trypanosomes representative of other main clades within the terrestrial and aquatic lineages. Phylogenetic analyses were inferred by maximum likelihood (ML) and Bayesian inferences (BI) as described previously [4, 9, 11, 12, 18]. The ML analysis was performed using RAxML v8.2 [51]. Tree searches employed GTR + GAMMA and proportion of invariable sites, model parameters were estimated in RAxML over the duration of tree search, and bootstrap supports estimated with 500 replicates. The BI analysis was conducted using MrBayes [52] with GTR + GAMMA. The first 25% of the trees was discarded as ‘burn in’.

Results
Tabanids from Brazil and Africa and tsetse flies harbour trypanosomes from the crocodilian clade
Two species of tabanids, *Phaeotabanus fervens* Linnaeus (48 flies) and *Tabanus occidentalis* Linnaeus (3 flies), herein identified by cox1 barcoding, were caught while feeding on *Ca. yacare* captured in the Miranda River, in the Pantanal biome, Brazil (Fig. 1). Nineteen flies, all females, were positive for trypanosomes by microscopy of their gut contents, yielding an overall trypanosome prevalence of ~37%. We obtained gGAPDH and/or V7V8 SSU rDNA sequences of trypanosomes in 17 out of these 19 positive guts: 15 from *P. fervens* and two from *Ta. occidentalis* (Additional file 1: Table S1, Figs. 2, 3, 4). Most sequences were highly similar to those of known caiman trypanosomes, namely *T. terena, T. ralphi* and *Trypanosoma* sp. Cay03. However, sequences not identical to any previously known caiman trypanosomes were obtained from gut DNA samples from 10 *P. fervens* and one *Ta. occidentalis*. These sequences were distributed into two new clades: Kaiowa and Tab01 (Figs. 2, 3).

Barcode analysis revealed that mixed infections by more than one trypanosome were highly prevalent in the Brazilian tabanids (Fig. 2). In the Pantanal biome, 65% of tabanids held mixed infections. Single infections were detected in just six flies: four harbouring caiman trypanosomes, one harbouring a bird trypanosome, and another harbouring *T. theileri*. We never detected *T. clandestinus* in tabanid flies.

Tabanids (*Atylotus* spp.; 83 flies) from Amazonia showed a 40% infection rate, and gGAPDH sequences obtained from these flies revealed that they were infected with *T. ralphi* (Additional file 1: Table S1, Fig. 3). Sequences from *T. theileri* and bird trypanosomes (currently being characterized) were also obtained from these flies. Of the Ethiopian tabanids (*Tabanus taeniola* Pali-sot de Beauvois and *Ancala africana* Griffith & Pidgeon) 11 out of 33 flies yielded sequences of trypanosomes, five of which were highly similar to those of *T. kaiowa* n. sp. (Additional file 1: Table S1, Fig. 4). DNA from guts of 37 tabanids from Mozambique submitted to PCR-sequencing of gGAPDH yielded no crocodilian trypanosomes. Regarding tsetse flies, gGAPDH sequences herein
analysed revealed the presence of *T. ralphi* in previously caught *Glossina morsitans morsitans* Wiedemann from Mozambique [47]. Also, in the present study, we detected *T. kaiowa* n. sp. and *Trypanosoma* sp. Tab01 in *Glossina pallidipes* Austen from Uganda (Additional file 1: Table S1, Figs. 3, 4).

From the gut contents of *P. fervens* (ISC203), we obtained one culture, which was cryopreserved as TCC2918. *SSU* rRNA and gGAPDH gene sequences from the isolate TCC2918 were identical to those of a previous isolate TCC1611 from *Ca. yacare* [11], which is being designed as the type-material of *Trypanosoma kaiowa* n. sp. (Figs. 3, 4).

**Phylogenetic inferences of trypanosomes from caimans, crocodiles, tabanid and tsetse flies**

The comparison of *V7V8 SSU* rDNA barcode sequences (Fig. 3) of trypanosomes from caimans and crocodiles, and from tabanids and leeches collected over caimans, confirmed their separation in the major terrestrial and aquatic lineages [12]. *Trypanosoma* sp. Cay03, *Trypanosoma* sp. Tab01, *T. grayi*, *T. terena*, *T. ralphi* and *T. kaiowa* n. sp. clustered into a single and strongly supported phylogenetic assemblage: the crocodilian clade of the terrestrial lineage of trypanosomes. *Trypanosoma* sp. Seputh from Sumatran rhinoceros, inhabitants of marshy areas, and *Trypanosoma terrestris*
Fig. 3 Phylogenetic analysis based on V7V8 SSU rRNA gene sequences of trypanosomes from caimans, crocodiles, tabanids and tsetse flies. Maximum Likelihood inference (903 characters, Ln = −4371.770725) supported the clades Ralphi, Kaiowa, Terena, Cay03 and Grayi in the terrestrial lineage. Trypanosomes of the aquatic lineages, including T. clandestinus, were used as outgroups. Numbers at the nodes (Bayesian Inference/Maximum Likelihood) represent the posterior probability > 0.8 and bootstrap support > 60%, respectively, derived from 500 replicates.

Fig. 4 Phylogenetic tree (ML) inferred using concatenated SSU rRNA and gGAPDH gene sequences of trypanosomes from caimans, crocodiles, tabanids, and tsetse flies. The analyses were inferred by Maximum Likelihood (ML, 1720 characters, Ln = −17593.067085) and Bayesian Inference (BI). Trypanosomes of the crocodilian clade were distributed in the clades Ralphi, Terena, Cay03, Tab01, Kaiowa, and Grayi. Trypanosomes of both terrestrial and aquatic lineages were used as outgroups. Numbers at nodes (ML/BI) are bootstrap support > 50%, and Bayesian posterior probability > 0.8 derived from 500 replicates. Insert: phylogram (ML) using gGAPDH sequences. Numbers at nodes are bootstrap values derived from 500 replicates.
restricted to African crocodiles. Interestingly, *Trypanosoma shawi* Nunes, 1987 [53] from capybaras, semi-aquatic Neotropical rodents that often share aquatic habitats with caimans and tapirs, was phylogenetically close to these trypanosomes (unpublished results). Nevertheless, corroborating the results of our previous study [12], *T. clandestinus* from caimans transmitted by aquatic leeches clustered with fish trypanosomes in the aquatic lineage.

Phylogenetic relationships inferred (ML and BI analyses) using either gGAPDH sequences or concatenated SSU rDNA and gGAPDH sequences strongly supported the partition of trypanosomes within the clade crocodilian into two main groups (Fig. 4). One phylogenetic group consists of the closely-related clades Kaiowa, Terena and Tabo1 while the other consists of the clades Ralphi, Grayi (well-supported), and Cay03 (weakly supported) (Fig. 4). Each clade was led by one trypanosome species (separated by 3–6% gGAPDH sequence divergence), and each species showed small intraspecies divergence: 0.5% in *T. terena*; 0.6% in *T. ralphi*; 0.4% in *T. kaiowa*; and 2.2% in *T. grayi* from The Gambia and Cameroon. The relevant genetic divergence among the isolates from tsetse flies and *Crocodylus niloticus* isolates from *T. kaiowa* n. sp. and *T. grayi* from semi-aquatic *neo*-leeches clustered with fish trypanosomes in the aquatic lineage.

The separation of *T. kaiowa* n. sp. from the remaining species of crocodilian trypanosomes was confirmed by all analyses of the present study (Figs. 3, 4). It is worth noting that in our previous study [11], some isolates from caimans were included in the clade Terena despite their relevant divergences. The most divergent isolate included in this clade was TCC1611 from caiman, whose gGAPDH and SSU rDNA sequences showed to be identical to those of the isolate TCC2918 from a tabanid fly reported in the present study. These two isolates clustered with sequences obtained directly from guts of tabanid and tsetse flies, forming the just now well-resolved clade Kaiowa (Figs. 3, 4). The degrees of intra-clade sequence divergence and the phylogenetic positioning strongly support *T. kaiowa* n. sp. as the single species in the clade Kaiowa. The trypanosome from the African dwarf crocodile (BSC27) differs from *T. kaiowa* n. sp. by only 0.5% in gGAPDH sequences and, hence, should be regarded as a genotype of *T. kaiowa* n. sp. In contrast, large divergences (~8%) separated *T. kaiowa* n. sp. from *T. grayi*, so far restricted to African crocodiles.

### Trypanosoma kaiowa n. sp.: morphology and predicted life-cycle

Flagellates were not detected in blood smears of any caiman assuredly infected exclusively with *T. kaiowa* n. sp. Therefore, blood trypomastigotes of this species still required morphological characterization. However, SSU rDNA and gGAPDH sequences (15 cloned sequences from each isolate) from the isolates of caiman blood (TCC1611) and tabanid gut (TCC2918) were identical, indicating that we have obtained pure cultures of *T. kaiowa* n. sp. from both the vertebrate host, and its putative vector.

Early hemocultures of the isolate TCC1611 (Fig. 5a) from *Ca. yacare* (Fig. 5b) captured in the Pantanal, Brazil, exhibited long and pointed trypomastigotes and epimastigotes with perceptible undulating membranes. These morphological features differed from the large roll-shaped trypomastigotes with exuberant undulating membranes of *T. terena* and *T. ralphi* both in early hemocultures and in blood of caimans [10–12]. The small flagellates presented in TCC1611 hemocultures in general resembled promastigote forms (Fig. 5a).

We also analyzed the morphology of flagellates in gut content from the tabanid fly (*Phaeotabanus fervens*) (Fig. 5b, c) infected with the isolate TCC2918 of *T. kaiowa* n. sp. Microscopic examination of gut contents revealed large clumps of small dividing parasites attached to the gut walls (Fig. 5c). The masses of flagellates are sometimes so dense that it is impossible to distinguish separate individuals, except for small flagellates detached from the clumps of parasites (Fig. 5c). Flagellates found free in the lumen of tabanid guts were mostly slender promastigotes, but a few epimastigotes and forms with kinetoplast at the posterior end resembling promastigotes were also detected (Fig. 5c). Trypanosomes were restricted to the digestive tract of the examined tabanids. However, salivary gland infection in later stages of the tabanid infection still needs to be investigated.

### Development of Trypanosoma kaiowa n. sp. co-cultured with Hi-5 insect cells (25 °C)

Light microscopy analyses of in Giemsa-stained preparations were conducted to evaluate the development of *T. kaiowa* n. sp. co-cultivated with a monolayer of Hi-5 insect cells in TC100 medium at 25 °C. In early cultures (24 h), inoculated promastigotes adhere to the surface of the insect cells by their flagella forming clusters of flagellates, and the flagellates likely invade the cells via the flagellum (Fig. 6a, b). After ~3 days, a few cells exhibited scarce rounded forms within vacuoles (Fig. 6c, d). However, 5–7 days cultures suggested that the parasites freely multiply in the cytoplasm of Hi-5 cells, and after ~7 days many cells show many rounded
Fig. 5 Trypanosoma kaiowa n. sp. predicted life-cycle. a Light microscopy of Giemsa-stained preparations of early hemocultures (BALB/LIT medium) from Caiman yacare naturally infected with T. kaiowa n. sp. showing large trypomastigote (T) and epimastigote (E) forms with noticeable undulating membrane, and small forms resembling promastigotes (P). b Caiman yacare and the tabanid Phaeotabanus fervens in the Pantanal, Brazil. c Clumps of flagellates adhered to a fragment of tabanid wall gut (yellow arrow), slim promastigotes (P), epimastigotes (E), and trypomastigotes (T) from gut contents of tabanids. Abbreviations: n, nucleus; k, kinetoplast; f, flagellum. Scale-bars: 10 µm
parasites (Fig. 6e, f). After 7–10 days, both rounded and short promastigote forms were, apparently, released by cells to the supernatant (Fig. 6g). The development of *T. kaiowa* n. sp. co-cultivated in Hi-5 cells was followed during ~20 days by observations of living (inverted phase microscopy), and fixed (Giemsa stained) cells. Growth behavior of *T. kaiowa* n. sp. in the supernatant of Hi-5 cultures was similar to cultures in LIT medium described below. Promastigotes multiply in the supernatant in general forming rosettes (Fig. 6h).

In early log-phase (2–3 days) of LIT cultures at 25 °C, *T. kaiowa* n. sp. forms small rosettes that quickly evolved to compact masses of numerous flagellates attached by their flagella (Fig. 7a). The flagellates gradually detach from the rosettes yielding abundant flagellates in the supernatant (~7 days). Log-phase trypanosomes lack perceptible undulating membranes, overall resembling small promastigotes (Fig. 7b, c). A few forms with lateral kinetoplast and an inconspicuous undulating membrane (characteristic of epimastigotes) were detected (Fig. 7d–f). Unexpectedly for stationary cultures, *T. kaiowa* n. sp. multiplied forming large rosettes of flagellates greatly varying in size (Fig. 7e) that gradually became very large flagellates with a pointed posterior extremity (Fig. 7h, i). A number of very long, often dividing flagellates, were observed in *T. kaiowa* n. sp. in old (> 15 days) cultures.

**Ultrastructural characterisation of *T. kaiowa* n. sp. by scanning and transmission electron microscopy**

Scanning electron microscopy of *T. kaiowa* n. sp. flagellates co-cultivated with monolayers of Hi-5 cultures corroborated previous evidence from light microscopy of flagellates adhering to cell surface by their flagella, and apparently (Fig. 8a) penetrating Hi-5 cells (Fig. 8b). Flagellates multiplied intensively (log-phase) either in supernatants (Fig. 7c) or adhered to coverslips forming large rosettes of flagellates attached by their flagella (Fig. 8d–e). Also confirming light microscopy, most flagellates resembled promastigote forms lacking a perceptible undulating membrane (Fig. 8c, f–h). A few pointed epimastigotes with an inconspicuous undulating membrane were detected in log-phase cultures of *T. kaiowa* n. sp. (Fig. 8f, h).

Transmission electron microscopy of log-phase *T. kaiowa* n. sp. co-cultivated with Hi-5 cultures revealed flagellates adhering to and, apparently, invading the insect cells (Fig. 9a, b). Transverse sections of a Hi-5 cell showed multiple sections of one long internalized flagellum (Fig. 9a), and small flagellate within vacuoles in the cytoplasm of Hi-5 cells (Fig. 9b). The ultrastructural organization of *T. kaiowa* n. sp. flagellates from the supernatants of Hi-5 cultures revealed all common organelles of trypanosomatids. However, a set of ultrastructural features of this new species merit to be highlighted:
a cytostome (Fig. 9c, d) that deeply invaginates into the cytoplasm (Fig. 8d); large multivesicular bodies possibly containing virus particles (Fig. 9d); many acidocalcisomes (Fig. 9e); short kinetoplast (Fig. 9d, f); free flagella (portion outside the flagellar pocket) exhibiting well-developed parflagellar structure (Fig. 9c, g), and noticeable lamella (Fig. 9g); multiple glycosomes near the nucleus (Fig. 9h), and well-developed tubules of the spongiome near the flagellar pocket that contains many vesicles (Fig. 9i).

**New species description**

**Family Trypanosomatidae Doflein, 1951**

**Genus Trypanosoma Gruby, 1843**

*Trypanosoma kaiowa* Teixeira & Camargo n. sp.
**Type-host**: Caiman yacare Daudin (Crocodylia: Alligatoridae).

**Other hosts**: Caiman crocodilus Linnaeus; Paleosuchus trigonatus Schneider; Osteolaemus tetraspis Cope.

**Type-locality**: Miranda River (19°57′ S, 57°01′ W) in the Pantanal, Mato Grosso do Sul, Brazil.

**Other localities**: Tabajara Reserve, Rondonia, Brazilian Amazonia; Lake Chamo, Ethiopia; Lake Edward, Uganda; Lagos de Cufada Natural Reserve, Guinea Bissau.

**Type-material**: Hapantotype: the culture of the isolate TCC1611 from Caiman yacare. Paratypes: culture TCC2918 from the tabanid Phaeotabanus fervens, and caiman blood and tissue (liver) samples (BSC28; BSC60; TSC40, TSC43) and tabanid guts (ISC204, ISC208, ISC211) infected with *T. kaiowa* n. sp. The cultures of *T. kaiowa* n. sp are cryopreserved at the Trypanosomatid Culture Collection of the University of São Paulo, TCC-USP, registered in the World Culture Collection, Japan. Blood/liver samples of crocodilians were preserved in ethanol at BSC (blood sample collection) and TSC (tissue sample collection), and insect samples were preserved in ethanol at ISC (Insect Sample Collection). DNA from *T. kaiowa* n. sp. cultures, caiman blood samples and tabanid guts are all conserved at TCC-USP together with Giemsa-stained smears of cultures and tabanid guts infected with *T. kaiowa* n. sp.

**Invertebrate host (putative vector)**: Phaeotabanus fervens Linnaeus (Diptera, Tabanidae). Additional vectors: Tabanus occidentalis Linnaeus, Ancala africana Griffith & Pidgeon, Tabanus taeniola Palisot de Beauvois, and Glossina pallidipes Austen.

**Site in host**: Blood and liver of crocodilians; the digestive tract of tabanids and tsetse flies.

**Representative DNA sequences**: DNA sequences were deposited in the GenBank database as follows:

![Fig. 8](image-url)
KF546503 and MG680219 (V7V8 SSU rRNA gene); KF546517 and MG680244 (gGAPDH gene).

ZooBank registration: To comply with the regulations set out in article 8.5 of the amended 2012 version of the International Code of Zoological Nomenclature (ICZN) [55], details of the new species have been submitted to ZooBank. The Life Science Identifier (LSID) of the article is urn:lsid:zoobank.org:pub:5849C465-F9F8-4C4C-A331-95B745F4021C.
**Etymology:** “Kaiowá” refers to a formerly large Indian Nation occupying a vast area of Mato Grosso do Sul, Paraguay, and Argentina. The Kaiowá and Terena are closely related indigenous people, as closely related to each other are *T. terena* and *T. kaiowa* found in caimans.

**Description**
Tabanid flies exhibited clumps of small oval-shaped flagellates attached to the gut walls, and free-swimming slender flagellates lacking perceptible undulating membranes resembling promastigotes, epimastigotes with inconspicuous undulating membranes, and trypomastigote forms. Early hemocultures exhibited long, wide and pointed trypomastigotes that evolved to rosettes of slender promastigotes. Measurements of log-phase promastigotes (LIT cultures) were 8.3–13 (10.3 ± 1.2) µm in length; 1.5–2.5 (2.0 ± 0.32) µm in width, and 12–25 (18.3 ± 3.6) µm of free flagellum. A few epimastigotes and very large and pointed flagellates were observed in old cultures. Light microscopy and ultrastructural features of all stages of *T. kaiowa* n. sp. are shown in Figs. 5, 6, 7 and 8.

**Remarks**
In our previous phylogenetic study, the isolate TCC1611 from *Ca. yacare* was placed in the clade Terena. However, this isolate was not considered a paratype of *T. terena* “because of relevant genetic divergences” (see [12]). In the present study, phylogenetic and morphological data confirmed that *T. terena* and the isolate TCC1611 (*T. kaiowa* n. sp.) are indeed distinct species. In addition, long and pointed trypomastigotes of *T. kaiowa* n. sp. differed from the roll-shaped trypomastigotes of *T. terena* and *T. ralphi* present in early hemocultures [10–12], and the predominance of promastigotes differentiates *T. kaiowa* n. sp. from these two species, both exhibiting epimastigotes with noticeable undulating membranes [11]. In addition, phylogenetic analyses also supported *Trypanosoma* sp. Cay03 and *Trypanosoma* sp. Tab01 as candidate new species. Unfortunately, they cannot yet be designated as new species because there are no valid holotypes for them: no cultures are available, and trypanosomes were not detected either in smears of caiman blood/tissue or tabanids single-infected with these trypanosomes.

**Discussion**
The improvement of molecular approaches for the straightforward barcoding of Trypanosomatidae, including next-generation sequencing, has unveiling an increasing number of previously unidentified genera and species of trypanosomes in a wide range of vertebrate and invertebrate hosts [4, 9, 11–17, 19–23, 50, 56–62]. Recent descriptions of new species and genotypes has largely enriched the genetic diversity of the genus *Trypanosoma*, and drastically improved our understanding of their evolutionary histories, but are still far from counterbalancing the bias of studies towards pathogenic trypanosomatids in humans and livestock. In addition to many fieldwork difficulties and the permits required for capturing wild animals and collecting blood/tissue samples, additional challenges include the proper identification of vectors, the assessment of both host species and geographical ranges, investigation of life-cycles and developmental stages, and potential pathogenicity of any newly discovered species.

This study provides ecological, biological, behavioral and molecular evidence supporting tabanids as vectors of *T. kaiowa* n. sp. Our examination of tabanid guts revealed masses of small flagellates attached to the gut epithelium indicating that *T. kaiowa* n. sp. heavily colonizes tabanid guts. Similarly, during its development in naturally and experimentally infected tsetse flies, *T. grayi* also exhibited small flagellates attached by their flagella to the gut wall [6]. The development of *T. kaiowa* n. sp. in tabanids also appears to be like that reported for *Trypanosoma melophagium* Flu, 1908 and *Trypanosoma lewisi* Kent, 1880 in the gut epithelium of their vectors hippoboscid flies and fleas (exhibiting intracellular forms), respectively [1]. It is unknown whether, in natural infections, *T. kaiowa* n. sp. invades cells of tabanid guts similarly to what occurred herein within cultured Hi5 insect cells. All evidence indicates that tabanids are the vector of caiman trypanosomes, but confirmation still requires experimental infection of laboratory-raised caimans by tabanids, as has been done for *T. grayi* with tsetse flies and crocodiles [5, 6], and *T. clandestinus* with aquatic leeches and caimans [12].

A remarkable peculiarity of *T. kaiowa* n. sp. is the predominance of promastigote forms in tabanid guts and cultures, whereas typical epimastigotes are scarce in both cultures and tabanid flies. The predominance of promastigotes differentiates *T. kaiowa* n. sp. from *T. terena* and *T. ralphi* both exhibiting epimastigotes with noticeable undulating membranes [11]. In fact, predominance of promastigotes is rarely observed in cultures of mammalian trypanosomes [1], and whether this could be common in other trypanosomes remains to be investigated.

Our preliminary analyses demonstrated that *T. kaiowa* n. sp. can be co-cultivated at 34 °C with LLCMK2 mammalian cells, proliferates in the supernatants of cultures and, apparently, flagellates could enter the cells (data not shown). However, further investigations are necessary to verify a possible intracellular development of *T. kaiowa* n. sp. The ability of crocodilian trypanosomes to survive in mammals is an interesting question recently raised by the detection of *T. grayi* DNA in blood of cattle from...
Cameroon (Africa), where tsetse flies showed large prevalence rates of both T. grayi and trypanosomes of ungulates [60]. Moreover, trypanosomes sharing mammalian and reptilian hosts have been reported from naturally infected rodents, marsupials, snakes and lizards [61, 62].

Our molecular survey unveiled two species of crocodilian trypanosomes in tabanids, T. kaiowa n. sp. and T. ralphi, plus the still unnamed Trypanosoma sp. Cay03, and Trypanosoma sp. Tab01. All trypanosomes from caimans clustered with T. grayi forming the crocodilian clade of the terrestrial lineage. The only exception was T. clandestinus that remains the sole trypanosome species of caimans nested in the aquatic lineage of Trypanosoma [12]. So far, known vertebrate hosts of caiman trypanosomes are Ca. yacare, Ca. crocodilus, M. niger and P. trigonatus, which harbour single or mixed trypanosome infections throughout South American hydrographical basins [10–12]. Here, we demonstrated that different species of tabanids (P. fervens, Ta. occidentalis and Atylotus sp.) can harbour different trypanosomes of the crocodilian clade: T. ralphi, T. kaiowa n. sp., and the putative new species Trypanosoma sp. Cay03 and Trypanosoma sp. Tab01.

Findings of ‘South American’ trypanosomes in African tabanid and tsetse flies suggest an underestimated diversity of crocodilian trypanosomes on the African continent, before the present study thought to be restricted to T. grayi, and exclusively transmitted by tsetse flies [5, 6]. In a recent molecular epidemiological study of Trypanosoma spp. in the guts of tsetse flies from Cameroon, T. grayi was a highly prevalent species [60].

Here, tsetse flies were found infected with three trypanosomes: T. ralphi in Mozambique, and T. kaiowa n. sp. and Trypanosoma sp. Tab01 in Uganda. Heterogeneous sequences of T. grayi were obtained from tsetse flies from The Gambia, and crocodiles from Cameroon [63, 64]. The wide host diversity and broad geographical distribution of crocodilian trypanosomes are indicative of their lack of host restriction and spatial structure. Regarding the vectors of these trypanosomes, the scenario is different: leeches transmit trypanosomes of the aquatic lineage (T. clandestinus), while tabanid and tsetse flies transmit trypanosomes of the terrestrial lineage (crocodilian clade). The evolutionary history of trypanosomes relied on their phylogenetic relationships and eco-geographical and paleontological data of their vertebrate and invertebrate hosts. Although all data suggested an ancient and common origin for the trypanosomes of the crocodilian clade, the very close phylogenetic relationships among Neotropical and Afrotropical trypanosomes of crocodilians underline a recent diversification of all trypanosomes so far identified in this clade. There are no caimans in Africa and no African dwarf crocodiles in South America, but these animals share highly similar genotypes of the same species of trypanosomes. This current disjunct distribution of crocodilian trypanosome species is consistent with the transcontinental dispersal of crocodilians harboring trypanosomes [10–12, 26–28].

Conclusions
To the best of our knowledge, data gathered in the present study provide the first evidence that tabanids can cyclically transmit trypanosomes to caimans and crocodiles in the Neotropic and Afrotropic. The description of T. kaiowa n. sp. in the present study relies on its phylogenetic positioning compared with other crocodilian trypanosomes. The description of T. kaiowa n. sp. is complemented by morphological and behavioural features of the flagellates in different growth conditions, and a predicted life-cycle in tabanid flies and caimans. The diversity and geographical dispersion of vertebrate and invertebrate hosts are also included in T. kaiowa n. sp. description. The close phylogenetic relationships of the crocodilian trypanosomes are in accordance with the transoceanic migration of crocodiles during the Miocene, and with the host-switching of trypanosomes between crocodiles and caimans. Our findings revealed that tabanid and tsetse flies are vectors of trypanosomes to crocodilians. Given that Glossinidae flies are absent from all continents other than Africa, adaptation to tsetse transmission would be secondary in the evolution of crocodilian trypanosomes, and tabanids, which are thought to have emerged during the Mesozoic (~140 mya), would be their ancestor vectors. However, this scenario relies on the contemporary geographical range of tsetse flies while records of Glossina-like fossils in the Nearctic and Palearctic at the Oligocene (30–40 mya) suggest that the ancestors of tsetse flies once flourished outside Africa, and possibly worldwide [36]. Data on crocodilian and tabanid trypanosomes from the Nearctic, Indomalayan and Australasian ecozones are still necessary for a better understanding of the evolutionary history and tracking the dispersion route of crocodilian trypanosomes.

Additional file

Additional file 1: Table S1. Host species, geographical origin and gGAPDH and V7V8 SSU rRNA gene sequences of trypanosomes from crocodilians, tabanids and tsetse flies.

Abbreviations
TCC: Trypanosomatid Culture Collection of the University of São Paulo, SP; Brazil; SSU: small subunit of ribosomal RNA; gGAPDH: glycosomal glyceraldehyde-3-phosphate dehydrogenase; PCR: polymerase chain reaction; Hi-5: insect (Trichoplusia ni) cell.

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Availability of data and materials
Cultures, DNA and glass-slides of the trypanosomes described in this study are deposited at the Trypanosomatid Collection of the University of São Paulo, Brazil. The newly generated DNA sequences were deposited in the GenBank database under the accession numbers MG680212–MG680225 (V7V8 SSU rRNA gene), MG680231–MG680259 (ggAPOH gene).

Authors’ contributions
EPC, MMGT, FP and BRF conceived, designed and coordinated field and laboratory studies. BRF performed part of the fieldwork in Brazil and Africa, participated in culture and morphological analysis, and was responsible for the phylogenetic analyses. LBV, CMFR, HAG participated in fieldworks and contributed with molecular characterization. FP was responsible for surveys in caimans and tabanids at the Pantanal. MC carried out culturing, morphological and biochemical analyses. CSAT performed electron microscopy. DS, JJK and AK supported fly collection in Africa. CAS) contribute with expertise and laboratory facility for electron microscopy, EPC, MMGT and BRF wrote the manuscript. All authors contributed to the revisions of the manuscript. All authors read and approved the final manuscript.

Ethics approval and consent to participate
All field procedures and caiman manipulations were performed according to the recommendations and authorisation of IBAMA (the Brazilian Institute for the Environment and Renewable Natural Resources), permit Number 10080-2, and approved by the Committees on the Ethics of Animal Experimentation of the Federal University of Mato Grosso do Sul, and Institute of Biomedical Science, University of São Paulo (Approved protocol 108/2-2013).

Consent for publication
Not applicable.

Competing interests
The authors declare that they have no competing interests.

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