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# Phylogenetic relationships of *Leishmania* species based on trypanosomatid barcode (SSU rDNA) and gGAPDH genes: Taxonomic revision of *Leishmania* (*L*.) *infantum chagasi* in South America $\stackrel{\circ}{\sim}$



Arlei Marcili<sup>a,\*</sup>, Marcia Ap. Sperança<sup>b</sup>, Andrea P. da Costa<sup>a</sup>, Maria de F. Madeira<sup>c</sup>, Herbert S. Soares<sup>a</sup>, Camila de O.C.C. Sanches<sup>b</sup>, Igor da C.L. Acosta<sup>a</sup>, Aline Girotto<sup>a</sup>, Antonio H.H. Minervino<sup>a</sup>, Maurício C. Horta<sup>d</sup>, Jeffrey J. Shaw<sup>e</sup>, Solange M. Gennari<sup>a</sup>

<sup>a</sup> Departamento de Medicina Veterinária Preventiva e Saúde Animal, Faculdade de Medicina Veterinária, Universidade de São Paulo, SP, Brazil

<sup>b</sup> Centro de Ciências Naturais e Humanas, Universidade Federal do ABC, Santo André, SP, Brazil

<sup>c</sup> Instituto de Pesquisa Clínica Evandro Chagas, Fundação Oswaldo Cruz, Rio de Janeiro, RJ, Brazil

<sup>d</sup> Colegiado de Medicina Veterinária, Universidade Federal do Vale do São Franscisco, Petrolina, PE, Brazil

<sup>e</sup> Departamento de Parasitologia, Instituto de Ciências Biomédicas, Universidade de São Paulo, São Paulo, SP, Brazil

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# ABSTRACT

Phylogenetic studies on trypanosomatid barcode using V7V8 SSU rRNA and gGAPDH gene sequences have provided support for redefining some trypanosomatid species and positioning new isolates. The genus *Leishmania* is a slow evolving monophyletic group and including important human pathogens. The phylogenetic relationships of this genus have been determined by the natural history of its vertebrate hosts, vector specificity, clinical manifestations, geographical distribution and molecular approaches using different markers. Thus, in an attempt to better understand the phylogenetic relationships of *Leishmania* species, we performed phylogenetic analysis on trypanosomatid barcode using V7V8 SSU rRNA and gGAPDH gene sequences among a large number of *Leishmania* species and also several Brazilian visceral *Leishmania infantum chagasi* isolates obtained from dogs and humans. Our phylogenetic analysis strongly suggested that *Leishmania hertigi* and *Leishmania equatoriensis* should be taxonomically revised so as to include them in the genus *Endotrypanum*; and supported ancient divergence of *Leishmania enriettii*. This, together with recent data in the literature, throws light on the discussion about the evolutionary southern supercontinent hypothesis for the origin of *Leishmania* spe. and validates *L. infantum chagasi* from Brazil, thus clearly differentiating it from *L. infantum*, for the first time.

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#### 1. Introduction

The genus *Leishmania* is an important group of parasites that is the causative agent of a complex disease called leishmaniasis, whose clinical manifestations exhibit different degrees of severity, ranging from asymptomatic, cutaneous/mucocutaneous lesions, to a more severe life-threatening visceral disease (Colmenares et al., 2002). *Leishmania* protozoans are transmitted by sandflies (Diptera; Psychodidae; Phlebotominae) (Schlein, 1993) to different species of mammals and lizards (Desjeux, 2004; Orlando et al.,

\* Corresponding author. Address: Av. Prof. Dr. Orlando Marques de Paiva, 87, São Paulo, SP 05508-270, Brazil. Tel.: +55 11 30911446; fax: +55 11 30917928.

E-mail address: amarcili@usp.br (A. Marcili).

2002; Rioux et al., 1992). The mammalian *Leishmania* species are transmitted in zoonotic and anthroponotic cycles, and are divided into two subgenera according to the developmental stage of the parasite in the gut of the sandfly vector (Lainson et al., 1987a). This division is also associated with its geographical distribution: *L.* (*Viannia*) is composed of species restricted to the Neotropics and *L.* (*Leishmania*) has species distributed in both the New and the Old World (Shaw, 1994).

Visceral leishmaniasis is the most severe form of the disease with a worldwide estimate of 500,000 human cases with 59,000 deaths every year, and it is thus a serious public health problem (Postigo, 2010). The species causing visceral leishmaniasis belong to the *L* complex of the subgenus *L*. (*Leishmania*) including: *L. eishmania donovani* (*L*.) *donovani*, which causes anthroponotic visceral leishmaniasis in India, Bangladesh, Nepal and Pakistan; Leishmania (*L*.) *infantum*, responsible for zoonotic leishmaniasis in the

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 $<sup>\,^{*}\,</sup>$  Note: Nucleotide sequences reported in this paper are available in the GenBank database. Accession numbers in Table 1.

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Mediterranean region (Europe and Africa); and *Leishmania* (*L.*) *chagasi*, which shows high genetic similarity with *L.* (*L.*) *infantum*, and has been correlated with zoonotic leishmaniasis in different countries of the Americas (New World) (Lukes et al., 2007). Dogs are the main reservoir of all of these visceral *Leishmania* zoonotic species.

The phylogenetic relationships among *Leishmania* species have been determined through differences in the natural history of their vertebrate hosts, vector specificity, clinical manifestations, geographical distribution and, more recently, using molecular approaches with different markers (Asato et al., 2009; Boite et al., 2012; Fraga et al., 2010; Lukes et al., 2007; Mauricio et al., 2004, 2007; Orlando et al., 2002; Shaw, 1997). However, these studies have included few species and/or natural isolates and controversy still surrounds the geographical origins and spread of *Leishmania* species. Also, some species included in the genus *Leishmania* have questionable taxonomic status, such as *Leishmania hertigi, Leishmania equatoriensis* and *Leishmania enriettii* (Momen and Cupolillo, 2000).

In the New World, Brazil is the country that accounts for the highest number (90%) of visceral leishmaniasis cases (Romero and Boelaert, 2010). The origin of these cases is unknown, since recent studies have been unable to confirm that the human disease was in this country prior to the European invasion of South America (Momen and Cupolillo, 2000), thus suggesting that L. (L.) chagasi arrived in Latin America from Europe. This hypothesis is corroborated by structural and biochemical findings, including studies using various molecular markers, such as the internal transcribed spacer of the ribosomal gene (Kuhls et al., 2005; Mauricio et al., 2004), gp63 (Mauricio et al., 2001; Quispe Tintaya et al., 2004), mini-exons (Mauricio et al., 2004), cysteine proteases (Hide et al., 2007; Quispe Tintaya et al., 2004), cytochrome oxidase II (Ibrahim and Barker, 2001) and microsatellites (Kuhls et al., 2011; Lukes et al., 2007). Based on these data, the taxonomy of the species causing visceral leishmaniasis in the Americas was recently revised and it was suggested that the species name L. (L.) infantum chagasi should be used (Kuhls et al., 2005, 2011; Lainson and Shaw, 1988; Mauricio et al., 1999, 2001, 2004). Controversially, studies based on vertebrate hosts and the biogeography of mammals have demonstrated a scenario in which there was ancient introduction of L. infantum chagasi into the Americas (Lainson et al., 1987b). Independent of the initial origin of *L. infantum chagasi*, molecular markers capable of distinguishing L. infantum from L. infantum chagasi would be very useful for epidemiological investigations.

Markers based on sequences of the SSU ribosomal gene have been used to study the diversity and phylogenetic relationships of trypanosomatids (Borghesan et al., 2013; Noyes et al., 1999; Stevens et al., 1999; Teixeira et al., 2011, 1994). V7V8 SSU rDNA has been named trypanosomatid barcode and has been used in several trypanosomatid phylogenetic studies and for describing inter and intraspecific relationships (Cortez et al., 2006; Da Silva et al., 2004; Ferreira et al., 2007; Lima et al., 2012; Marcili et al., 2009a,b; Rodrigues et al., 2006; Viola et al., 2009; Teixeira et al., 2011). In addition to the ribosomal gene, the gGAPDH sequence has made it possible to position trypanosomatid species (Hamilton et al., 2004, 2007).

Thus, in an attempt to better understand the phylogenetic relationships of *Leishmania* species, we performed phylogenetic analysis on *Trypanosomatidae* barcode using V7V8 SSU rRNA and gGAPDH gene sequences among a large number of *Leishmania* species and also several Brazilian visceral *L. i. chagasi* isolates obtained from dogs and humans. The evolutionary insights from the results relating to the *L. hertigi, L. equatoriensis* and *L. enriettii* reference strains, and from *L. i. chagasi* isolates from humans and dogs from Brazil, strongly suggest that taxonomic revision of these species is required.

# 2. Material and methods

# 2.1. Isolation and culturing of Leishmania species

L. i. chagasi was isolated through culturing popliteal lymph node aspirates obtained from fine-needle puncture performed on dogs in several Brazilian regions (Table 1). The culturing was done in biphasic blood agar base and liver infusion tryptose (LIT) medium (Camargo, 1964). Promastigote forms from positive cultures were used to infect monolayers of J774 peritoneal macrophage lineage cells in a 25 cm<sup>2</sup> flask that was kept in RPMI medium containing 10% bovine calf serum at 28 °C. Infected J774 cells were monitored until cell lysis and release of promastigote forms to the RPMI medium occurred. The promastigote forms of the Leishmania species were transferred to and expanded in LIT medium containing 10% bovine calf serum and 2% male human urine for DNA preparation and cryopreservation. All isolates were included in the Brazilian Tripanosomatid Collection of the School of Veterinary Medicine of the University of São Paulo, Brazil. The study was approved by the Bioethical Committee in Animal Research of the Faculty of Veterinary Medicine of the University of São Paulo.

# 2.2. PCR amplification of 18S rDNA and glycosomal glyceraldehyde 3phosphate dehydrogenase (gGAPDH) coding sequences

DNA from Leishmania isolates was extracted from culture supernatants using the phenol-chloroform method. The DNA samples were subjected to conventional polymerase chain reaction (PCR) with high fidelity Taq DNA polymerase for amplification of the barcode, which comprised a fragment of around 900 base pairs (bp) corresponding to the trypanosome V7V8 SSU rDNA (Ferreira et al., 2008; Marcili et al., 2009a,b; Viola et al., 2008), and of the approximately 850 bp fragment corresponding to the gGAPDH trypanosome coding sequence described previously (Hamilton et al., 2007). The obtained PCR products were purified using Spinx-X columns (Costar) and cloned into the pCR<sup>™</sup>2.1 TA vector (Invitrogen). Sequences of three to five clones of each PCR product were determined to minimize sequence mistakes generated by Tag DNA polymerase and occurrence of different haplotypes in Leishmania isolates. The sequences trace files were analyzed and compiled using the Segman program of DNAStar software. The nucleotide sequences generated were deposited in GenBank (Table 1).

#### 2.3. Phylogenetic analysis

The sequences of trypanosomatid barcode obtained were aligned with sequences previously determined for other trypanosomatid species available in GenBank (Table 1) using ClustalX (Thompson et al., 1997) and secondary structure comparative analysis and were adjusted manually using GeneDoc (Nicholas and Nicholas, 1997). The gGAPDH sequences of the visceral leishmaniasis species obtained were aligned to evaluate intraspecific variability. The barcode and gGAPDH sequences were used to construct a phylogenetic tree using maximum parsimony, as implemented in PAUP version 4.0b10 (Wilgenbusch and Swofford, 2003) with 500 bootstrap replicates, random stepwise addition starting trees (with random addition sequences) and TBR branch swapping. Sequences of V7V8 SSU rDNA and gGAPDH genes of L. donovani complex were concatenated to perform intraspecific sequence variation analysis. Bayesian analysis was performed using MrBayes v3.1.2 (Huelsenbeck and Ronquist, 2001) with 1,000,000 generations. Trees were sampled every 100 generations using chains, and 25% of the early sample trees were discarded as "burn-in". The GTR+I+G substitution model was used.

# Table 1

Trypanosomatid isolates, host, geographical origin and sequences of SSU rDNA and gGAPDH genes employed in the phylogenetic analyses performed in this study.

Trypanosomatid Species	CBT <sup>a</sup>	Isolate code <sup>b</sup>	Host	Geographic origin <sup>c</sup>		Acession number <sup>d</sup> Sequences	
						SSUrRNA	gGAPDH
<b>Phytomonas</b> P. serpens P. sp						AF016320 AF016322	
Hernetomonas							
H. megaseliae H. muscarum H. samuelpessoai			Megaselia scalaris Zelus leucogrammus			U01014 L18872 U01016	
<b>Crithidia</b> C. fasciculata			-			Y00055	
Wallacoina							
W. brevicula W. inconstans			Nabis brevis Calocoris sexguttatus	Russia Russia		AF153045 AF153044	
Leptomonas							
L. podlipaevi L. seymouri L. costaricensis			Dysdercus suturellus Ricolla simílima	Costa Rica		DQ383649 AF153040 DO383648	
Endotrumanum						2 23030 10	
E. monterogei Endotrypanum sp.		TCC 889	Nyssomyia trapidoi Psathyromyia dendrophyla	Ecuador Rondonia	EC BR	JQ863389 EU021240	
Endotrypanum sp.		TCC 890	Lutzomyia gomezi	Rondonia	BR	EU021238	
<b>Leishmania</b> L. adleri L. tarentolae						M80291 M84225	
L. hoogstraali						KF041810	
L. tropica	70	MHOM/SU/1958/STRAIN OD	Homo sapiens	Azerbaijão	AZ	KF041809	
L. amazonensis			Homo sapiens	Brazil	BR	GQ332354	
L. major L. major			Homo sapiens	l urcomenistao Israel	I M II	GQ332361 AF244350	
L. hertigi	74	MCOE/PA/1965/C8	Coendou sp.	Panamá	PA	KF041804	
L. lainsoni	75	MHOM/BR/1981/M6426	Homo sapiens	Brazil	BR	KF041805	
L. donovani	77	MHOM/ET/1967/L82;HV3;LV9	Homo sapiens	Ethiopia	ET	KF041801	FR799617
L. equatoriensis	79	MCHO/EC/1982/LSP1	Choloepus hoffmanni	Ecuador	EC	KF041802	
L. mexicana	80	MHOM/BZ/1982/BEL21	Homo sapiens	Belize	BZ	KF041806	
L. shawi	87	MCEB/BR/1984/M8408	Cebus apella	Brazil	BR	KF041808	
L. donovani	89	MHOM/ET/1967/HU3	Homo sapiens	Ethiopia	ET	KF041800	XM_003862962
L. guyanensis	90	MHOM/BR/1975/M4147	Homo sapiens	Brazil	BR	KF041803	
L. naiffi	91	MDAS/BR/1979/M5533	Dasypus sp.	Brazil	BR	KF041807	
L. braziliensis		MHOM/BR/1975/M2903	Homo sapiens	Brazil	BR	<u>KF041799</u>	
L. enriettii	47	1001.00	Cavia porcellus	Paraná	BR	KF041798	
L. infantum		JCPM5		Europe		M81430	FR/96462/XM_001467109
L. infantum	88	MHOM/BR/2002/LPC-RPV	Homo sapiens	Brazil	BR	KF041793	
L. infantum chagasi	72	MHOM/BR/1974/PP75	Homo sapiens	Brazil	BR	KF041792	
L. infantum chagasi	13	C808	Canis lupus familiaris	Rio de Janeiro	BR	KF041776	KF041811
L. infantum chagasi	15	C1148	Canis lupus familiaris	Rio de Janeiro	BR	KF041777	KF041812
L. infantum chagasi	16	C1194	Canis lupus familiaris	Rio de Janeiro	BR	KF041778	KF041813
L. infantum chagasi	22	MCAN/BR/1984/CCC17.482	Canis lupus familiaris	Ceará	BR	KF041779	
L. infantum chagasi	24	MCAN/BR/1995/CHULINHA	Canis lupus familiaris	Bahia	BR	KF041780	
L. infantum chagasi	25	MCAN/BR/2002/LVV-129	Canis lupus familiaris	Mato Grosso do Sul	BR	KF041781	KF041814
L. infantum chagasi	26	MCAN/BR/2005/NMT-DOTM	Canis lupus familiaris	Distrito Federal	BR	KF041782	<u>KF041815</u>
L. infantum chagasi	28	MCAN/BR/2004/LIBPI-18	Canis lupus familiaris	Piauí	BR	KF041783	
L. infantum chagasi	29	MCAN/BR/2007/LIBPI-51	Canis lupus familiaris	Piauí	BR	KF041784	<u>KF041816</u>
L. infantum chagasi	30	MCAN/BR/2010/BURRINHO II	Canis lupus familiaris	Rio Grande do Sul	BR	KF041785	
L. infantum chagasi	31 27	INICAN/BR/2010/LAIKA	Canis lupus familiaris	KIO Grande do Sul	BK	<u>KF041786</u>	<u>KF041817</u>
L. IIIJantum chagasi	/د مد		Canis lupus familiaris	rernambuco Dará	BK	KF041787	
L. IIIJantum chagasi	39		Canis lupus familiaris	r'dl'a Dará	BK	KF041788	KF041818
L. IIIJUIIIUIII Chagasi	40 42		Canis lupus familiaris	r di d Dará	ык рр	KF041789	<u>KF041819</u>
L. infantum chagasi	43		Canis iupus juitillaris	r al a Dará	DK RD	KF041790	<u>KF041820</u>
L. infantum chagasi	-1-1	1.447	Carità iupus juininunis	i ai a São Daulo	חם חם	<u>KF041791</u>	
L. mjumum chagasi		LVA4/	nomo supiens	SaU l'auto	DK	<u>KF041794</u>	

Trypanosomatid Species	CBT <sup>a</sup>	Isolate code <sup>b</sup>	Host	Geographic origin <sup>c</sup>		Acession number <sup>d</sup> Sequences		
						SSUrRNA	gGAPDH	
L. infantum chagasi		C1549	Canis lupus familiaris	Mato Grosso	BR	KF041797		
L. infantum chagasi		C1149	Canis lupus familiaris	Rio de Janeiro	BR	KF041795		
L. infantum chagasi		C1548	Canis lúpus familiaris	Mato Grosso	BR	KF041796		

<sup>a</sup> CBT, Code number of the isolates/strains cryopreserved in the Coleção Brasileira de Tripanossomatídeos (CBT).

<sup>b</sup> Original codes of isolates.

<sup>c</sup> Bahia, Ceará, Distrito Federal, Mato Grosso, Mato Grosso do Sul, Pará, Pernambuco, Piauí, Rio de Janeiro, Rondonia and São Paulo are Brazilian states.

<sup>d</sup> Sequences determined in this study and deposited in the Genbank are underlined.

# 3. Results

To evaluate phylogenetic relationships and polymorphisms within Leishmania species from different hosts and geographical origins, as well as to analyze the degree of genetic relatedness among these isolates and those of other trypanosomatid genera, we compared sequences of V7-V8 SSU rDNA genes aligned with those from reference strains from GenBank (Table 1). Most Leishmania species were aggregated in a monophyletic group, except for L. equatoriensis and L. hertigi, which were segregated with an isolate from the genus Endotrypanum (0.38% divergence of sequences and 95% bootstrap and 100% posterior probability) (Fig. 1). A single isolate of L. enriettii obtained from the host Cavia porcellus in the Brazilian state of Paraná (CBT 47) segregated at the basal branch in all phylogenetic results obtained and was placed together with Leishmania sp. (MAR 1), a human isolate from Martinique (100% bootstrap and 100% posterior probability). The divergence of sequences of L. enriettii and other Leishmania species ranged from 3.84% to 5.48% (Fig. 1).

The different species of the genus Leishmania were segregated into three groups: I. parasitic Leishmania species of Old World lizards including Leishmania adleri, Leishmania tarentolae and Leishmania hoogstraali (0.51% divergence of sequences and 92% bootstrap and 95% posterior probability); II. several human-infecting Leishmania species of different geographical origins that presented a broad spectrum of clinical manifestations (1.28% divergence of sequences and 83% bootstrap and 100% posterior probability); III. New and Old World species of visceral leishmaniasis (0.23% divergence of sequences and 80% bootstrap and 100% posterior probability), separated according to the geographical origin of the isolates. Phylogenetic analyses on the SSU rDNA V7V8 region grouped L. infantum and L. donovani distributed in the Old World separately from the Brazilian isolates of the New World L. infantum chagasi, with 100% similarity and 72/100% bootstrap and posterior probability, respectively (Fig. 1).

In all analyses using the trypanosomatid barcode region, L. i. chagasi isolates segregated into two branches according to their American (L. infantum chagasi) or Old World (L. infantum) origin. L. i. chagasi isolates from humans and dogs from South America (Brazil) were tightly clustered together (100% sequence similarities) (Fig. 1). All L. i. chagasi SSU rDNA V7V8 region and gGAPDH sequences were identical between them. The phylogenetic analysis based only on the gGAPDH gene separated L. infantum and L. donovani but was unable to separate the isolates of L. infantum chagasi from L. infantum. Phylogenetic analysis based on concatenated V7V8 SSU rDNA and gGAPDH genes enabled separation of the L. donovani complex species (Fig. 2). All L. i. chagasi isolates were clustered together (100% bootstrap/100% posterior probability and 100% of similarity) and were segregated from L. infantum and L. donovani, with 90% and 74% bootstrap and 100% posterior probability, respectively (Fig. 2).

#### 4. Discussion

With the aim of gaining better understanding of the phylogenetic relationships of *Leishmania* species, we performed phylogenetic analysis on *Trypanosomatidae* barcode using V7V8 SSU rRNA and gGAPDH gene sequences among a large number of *Leishmania* species and also several Brazilian visceral *L. infantum chagasi* isolates obtained from dogs and humans.

The results from the genus *Trypanosoma* using gGAPDH and SSU rDNA have shown congruent phylogenetic relationships (Hamilton et al., 2004, 2007), while the same gene sequences used with the genus *Leishmania* revealed unrelated results (Fig. 2). The divergence values among the sequences of the *Leishmania* species ranged from 2.29% to 0.90%, thereby separating the Sauroleishmania species group from the *Vianna* and *Leishmania* subgenera. These values are low in comparison with the divergence rates of *Trypanosoma cruzi* intraspecific parasites (Marcili et al., 2009a,b,c,), but are similar to divergence differences among *Trypanosoma rangeli* and *Trypanosoma theileri* lineages (da Silva et al., 2004; Rodrigues et al., 2006). These results are also compatible with the low evolution rates of *Leishmania* parasites, in comparison with other trypanosomatids (Stevens and Rambaut, 2001).

Our data from phylogenetic analysis on concatenated and independent V7V8 SSU rRNA and gGAPDH sequences corroborated the monophyletic origin of the genus *Leishmania* and also the data obtained using different SSU rDNA gene sequences (Briones et al., 1992; Orlando et al., 2002; Uliana et al., 1991; van Eys et al., 1992) and the gGAPDH gene (Hannaert et al., 1998), thus positioning the Sauroleishmania group as ancestral.

L. hertigi and L. equatoriensis were grouped within the genus Endotrypanum and were excluded from Leishmania. The biological cycles of L. hertigi and L. equatoriensis exhibit amastigote forms resembling Leishmania parasites (Lainson and Shaw, 1969), and species of the genera Leishmania and Endotrypanum are transmitted by sandflies that are genetically closely related (Cupolillo et al., 2000), which may explain the controversy regarding the taxonomic position of these organisms. However, differing from Leishmania species, which are able to parasitize different vertebrates such as sloths, anteaters, armadillos, opossums, rodents, canids, etc., and are distributed worldwide, the genus Endotrypanum is composed of parasites unique to sloth species that are found in Brazil, Colombia, Guyana and Central America (Herrer, 1971; Silva et al., 2013; Zeledon and Ponce, 1977). L. hertigi and L. equatoriensis were included in the genus Leishmania by molecular studies based on methodologies such as isoenzyme patterns or nucleic acid polymorfism sizes determined by enzymes restriction patterns (Cupolillo et al., 1998a,b, 2000). Such methods are useful to evaluate intraspecific polymorphisms, but have no evolutionary and/or taxonomic information. These studies, demonstrated that L. hertigi and L. equatoriensis presented polymorphisms more similar to the *Endotrypanum* than to the Leishmania genus species, indicating possible taxonomic prob-



**Fig. 1.** Maximum parsimony and Bayesian tree inferred from SSU rDNA gene sequences on 57 trypanosomatids with *Phytomonas* as the outgroup (841 characters; 65 parsimony-informative sites). Numbers at nodes are the support values for the major branches (bootstrap/posterior probability; 500 replicates). The sequences obtained in this study are underlined.

lems. Also, the SSUrDNA gene sequences were not employed for phylogenetic positioning and taxonomic resolution of these parasites. Moreover, molecular analysis using PCR-RFLP on SSU rDNA and cytochrome b gene sequences has demonstrated that *L. hertigi* is more closely related to *Endotrypanum* than to *Leishmania* (Asato

et al., 2009; Noyes et al., 1997). Thus, we suggest that the taxonomic position of *L. hertigi* and *L. equatoriensis* should be revised so as to include these species in the genus *Endotrypanum*.

Phylogenetic analysis on the SSU rDNA gene showed that *Leishmania* (*L*) *enriettii*, which is a parasite found in naturally



**Fig. 2.** Dendogram based on concatenated SSU rDNA and gGAPDH sequences on 14 *Leishmania* species of the *L. donovani* complex, which was used for maximum parsimony and Bayesian inferences with 1469 characters. Numbers at nodes are the support values for the major branches (bootstrap/posterior probability; 500 replicates). The sequences obtained in this study are underlined.

infected domestic guinea pigs (C. porcellus) in the Brazilian states of Paraná and São Paulo (Machado et al., 1994; Muniz and Medina, 1948; Thomaz-Soccol et al., 1996) and probably in other sylvatic rodents (Lainson, 1997), has an ancient origin and was the first species to diverge within the genus Leishmania. This result is corroborated by molecular and distance analysis on the Leishmania cytochrome b and DNA and RNA polymerase gene sequences (Asato et al., 2009; Noves et al., 2002). More recently, molecular characterization and distance analysis on the RNA polymerase II gene of Leishmania sp. isolated from skin lesions of red kangaroos in a rural area of Australia, and also from captive macropods, including several Northern wallaroos, a black wallaroo and agile wallabies, showed that these parasites were also grouped within the L. enriettii complex (Dougall et al., 2009; Rose et al., 2004). Considering that the phylogenetically closely related parasite L. (L.) enriettii is present in Australia, the early divergence and ancient origin of these species can be supported by the southern supercontinent hypothesis for the evolutionary history of other Trypanosomatidae parasites that have been isolated from bats. In this hypothesis, T. cruzi and related parasites evolved in isolation in the mammals of South America, Antarctica and Australia (Hamilton et al., 2012). In this scenario, to understand the evolutionary history of Leishmania, it would be important to characterize the biological cycle of these L. (L). enriettii parasites, which is still poorly known. In the Brazilian state of Paraná, the sandfly Lutzomyia monticola was incriminated as a possible vector for L. enriettii (Luz et al., 1967), while in Australia, the potential vectors were found to be midges (Diptera: Ceratopogonidae) (Dougall et al., 2011). The evolutionary implication of different dipteran vectors in the life cycle of the L. (L.) enriettii group needs further investigation. In addition, it is important to note that human infection caused by parasites molecularly similar to L. (L.) enriettii has been described in HIV patients in Martinique, in the French West Indies (Noyes et al., 2002), suggesting that this group of ancient Leishmania parasites is also present in Central America.

The evolutionary history of *L. i. chagasi*, the etiological agent of visceral leishmaniasis in South America has been widely debated. Some hypotheses suggest that *L. i. chagasi* was introduced along with the expansion of the canids in the New World about 2–3 million years ago (Lainson et al., 1987b). However, due to the failure to

distinguish *L. chagasi* from *L. infantum* through several biochemical and molecular studies performed so far (Hide et al., 2007; Ibrahim and Barker, 2001; Kuhls et al., 2005, 2011; Lukes et al., 2007; Mauricio et al., 2001, 2004; Quispe Tintaya et al., 2004), the taxonomy of the species causing visceral leishmaniasis in the Americas was recently revised and it was suggested that the name *L. (L.) infantum chagasi* should be used (Kuhls et al., 2005, 2011; Lainson and Shaw, 1988; Mauricio et al., 1999, 2001, 2004). However, our phylogenetic analysis using Trypanosomatidae barcode resulted in separation of *L. chagasi* and *L. infantum*, thus revitalizing the discussion on the origin of *L. chagasi*.

Since canids are natural reservoirs for L. infantum and L. chagasi, and the evolutionary histories of vertebrate hosts are important tools in reconstructing the evolutionary history of parasites of the family Trypanosomatidae (Hamilton, 2012; Hamilton et al., 2012), a discussion about the origin and radiation of canids could correlate with the introduction of these parasites in South America. as believed previously (Lainson et al., 1987b). The first records of the order Carnivora are from the Paleocene and Eocene, with radiation in North America and subsequently in Eurasia (Azzaroli et al., 1988; Sotnikova and Rook, 2010). The members of this order were originally absent in South America and Australia, where their niches were occupied by marsupials (Eisenberg and Redford, 1999). Procyonids were the first carnivores to migrate to South America during the Miocene, and other groups arrived only after the switching on the Isthmus of Panama during the Pliocene (Linares, 1981) Furthermore, several studies based on fossil records have shown that vast diversification of canids occurred in North America (Spassov and Rook, 2006; Sotnikova, 2006; Wang and Telford, 2007; Garrido and Arribas, 2008). Thus, if North American canids were responsible for the introduction of L. chagasi into South America, they should harbor L. donovani complex parasites. However, cases of visceral leishmaniasis in North America are rare, and are related to transplacental transmission, blood transfusion and exchange of dogs from endemic regions, without occurrences of vector transmission (Boggiatto et al., 2011; Duprey et al., 2006; Gibson-Corley et al., 2008; Giger et al., 2002; Owens et al., 2001). Furthermore, serological studies on Grav and Red Fox populations in North America have shown low positivity and possibly even cross-reactivity with T. cruzi, which is endemic in the region studied (Rosypal et al., 2010a,b). These pieces of evidence go against an ancient origin for L. i. chagasi in South America.

Therefore, the more likely origin of *L. i. chagasi* corresponds to the introduction of *L. infantum* around five hundred years ago, during the colonization of South America by Europeans, through infected domestic dogs. The low divergence found between *L. infantum* and *L. chagasi* (0.26%) based on V7V8 SSU rRNA does not explain and/or distinguish between the two hypotheses regarding the evolutionary origin of these species. However, the biogeography of canid hosts, *L. donovani* complex parasite occurrence and distribution in North America and the low divergence of *L. chagasi/L. infantum* V7V8 SSU rRNA values strongly support the notion that *L. chagasi* originated due to introduction of parasites through human actions. This hypothesis is also corroborated by the bottleneck signature of *L. chagasi* in the New World, accompanied by a thousand-fold decrease in population diversity in comparison with *L. infantum* in Europe (Kuhls et al., 2011).

The populational variability shown by microsatellites demonstrated gene flow between New and Old World *L. infantum* and corroborated the notion of recent introduction of *L. infantum* species into the Americas (Kuhls et al., 2008, 2011). However, with this approach, it was not possible to distinguish between the New and Old World parasite populations, probably due to the characteristic gene flow of microsatellites.

Nevertheless, our result validates the existence of the subspecies L. i. chagasi as a valid taxonomic unit, and its specific identification by the V7V8 SSU rRNA sequence enables epidemiological studies. This is of particular importance to Brazil and other Latin American countries, where visceral leishmaniasis caused by *L. i. chagasi* is expanding, following the dispersion of its major vector, *Lutzomyia longipalpis*, which has become adapted to periurban environments (Dias et al., 2011; Nascimento et al., 2008) and has spread to new areas in Argentina and Uruguay, thereby expanding the endemic regions affected by human-threatening visceral leishmaniasis (Cordoba-Lanus et al., 2006; Gould et al., 2013; Salomon et al., 2011).

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#### References

- Asato, Y., Oshiro, M., Myint, C.K., Yamamoto, Y., Kato, H., Marco, J.D., Mimori, T., Gomez, E.A., Hashiguchi, Y., Uezato, H., 2009. Phylogenic analysis of the genus *Leishmania* by cytochrome b gene sequencing. Exp. Parasitol. 121, 352–361.
- Azzaroli, A., De Giuli, C., Ficcarelli, G., Torre, D., 1988. Late Pliocene to Early Mid-Pleistocene mammals in Eurasia: faunal succession and dispersal events. Palaeogeography, Palaeoclimatology, Palaeoecology 66, 77–100.
- Boggiatto, P.M., Gibson-Corley, K.N., Metz, K., Gallup, J.M., Hostetter, J.M., Mullin, K., Petersen, C.A., 2011. Transplacental transmission of *Leishmania infantum* as a means for continued disease incidence in North America. PLoS Negl. Trop. Dis. 5, e1019.
- Boite, M.C., Mauricio, I.L., Miles, M.A., Cupolillo, E., 2012. New insights on taxonomy, phylogeny and population genetics of *Leishmania* (*Viannia*) parasites based on multilocus sequence analysis. PLoS Negl. Trop. Dis. 6, e1888.
- Borghesan, T.C., Ferreira, R.C., Takata, C.S., Campaner, M., Borda, C.C., Paiva, F., Milder, R.V., Teixeira, M.M., Camargo, E.P., 2013. Molecular phylogenetic redefinition of *Herpetomonas* (Kinetoplastea, Trypanosomatidae), a genus of insect parasites associated with flies. Protist 164, 129–152.
- Briones, M.R., Nelson, K., Beverley, S.M., Affonso, H.T., Camargo, E.P., Floeter-Winter, L.M., 1992. *Leishmania tarentolae* taxonomic relatedness inferred from phylogenetic analysis of the small subunit ribosomal RNA gene. Mol. Biochem. Parasitol. 53, 121–127.
- Camargo, E.P., 1964. Growth and differentiation in *Trypanosoma cruzi*. I. Origin of metacyclic trypanosomes in liquid media. Rev. Inst. Med. Trop. Sao Paulo 6, 93– 100.
- Colmenares, M., Kar, S., Goldsmith-Pestana, K., McMahon-Pratt, D., 2002. Mechanisms of pathogenesis: differences amongst *Leishmania* species. Trans. R. Soc. Trop. Med. Hyg. 96 (Suppl. 1), S3–S7.
- Cordoba-Lanus, E., De Grosso, M.L., Pinero, J.E., Valladares, B., Salomon, O.D., 2006. Natural infection of *Lutzomyia neivai* with *Leishmania* spp. in northwestern Argentina. Acta Trop. 98, 1–5.
- Cortez, A.P., Ventura, R.M., Rodrigues, A.C., Batista, J.S., Paiva, F., Anez, N., Machado, R.Z., Gibson, W.C., Teixeira, M.M., 2006. The taxonomic and phylogenetic relationships of *Trypanosoma vivax* from South America and Africa. Parasitology 133, 159–169.
- Cupolillo, E., Pereira, L.O., Fernandes, O., Catanho, M.P., Pereira, J.C., Medina-Acosta, E., Grimaldi Jr., G., 1998a. Genetic data showing evolutionary links between Leishmania and Endotrypanum. Mem. Inst. Oswaldo Cruz. 935, 677–683.
- Cupolillo, E., Momen, H., Grimaldi Jr., G., 1998b. Genetic diversity in natural populations of New World Leishmania. Mem. Inst. Oswaldo Cruz. 935, 663–668.
- Cupolillo, E., Medina-Acosta, E., Noyes, H., Momen, H., Grimaldi Jr., G., 2000. A revised classification for *Leishmania* and *Endotrypanum*. Parasitol. Today 16, 142–144.
- Da Silva, F.M., Noyes, H., Campaner, M., Junqueira, A.C., Coura, J.R., Anez, N., Shaw, J.J., Stevens, J.R., Teixeira, M.M., 2004. Phylogeny, taxonomy and grouping of *Trypanosoma rangeli* isolates from man, triatomines and sylvatic mammals from widespread geographical origin based on SSU and ITS ribosomal sequences. Parasitology 129, 549–561.

Desjeux, P., 2004. Leishmaniasis. Nat. Rev. Microbiol. 2, 692.

- Dias, E.S., Regina-Silva, S., Franca-Silva, J.C., Paz, G.F., Michalsky, E.M., Araujo, S.C., Valadao, J.L., de Oliveira Lara-Silva, F., de Oliveira, F.S., Pacheco, R.S., Fortes-Dias, C.L., 2011. Eco-epidemiology of visceral leishmaniasis in the urban area of Paracatu, state of Minas Gerais, Brazil. Vet. Parasitol. 176, 101–111.
- Dougall, A., Shilton, C., Low Choy, J., Alexander, B., Walton, S., 2009. New reports of Australian cutaneous leishmaniasis in Northern Australian macropods. Epidemiol. Infect. 137, 1516–1520.
- Dougall, A.M., Alexander, B., Holt, D.C., Harris, T., Sultan, A.H., Bates, P.A., Rose, K., Walton, S.F., 2011. Evidence incriminating midges (Diptera: Ceratopogonidae) as potential vectors of *Leishmania* in Australia. Int. J. Parasitol. 41, 571–579.
- Duprey, Z.H., Steurer, F.J., Rooney, J.A., Kirchhoff, L.V., Jackson, J.E., Rowton, E.D., Schantz, P.M., 2006. Canine visceral Leishmaniasis, United States and Canada, 2000–2003. Emerg. Infect. Dis. 12, 440–446.
- Eisenberg, J.F., Redford, K.H., 1999. Mammals of the neotropics the central neotropics, 3. University of Chicago Press, Chicago.
- Ferreira, R.C., Campaner, M., Viola, L.B., Takata, C.S., Takeda, G.F., Teixeira, M.M., 2007. Morphological and molecular diversity and phylogenetic relationships among anuran trypanosomes from the Amazonia, Atlantic Forest and Pantanal biomes in Brazil. Parasitology 134, 1623–1638.
- Ferreira, R.C., De Souza, A.A., Freitas, R.A., Campaner, M., Takata, C.S., Barrett, T.V., Shaw, J.J., Teixeira, M.M., 2008. A phylogenetic lineage of closely related trypanosomes (Trypanosomatidae, Kinetoplastida) of anurans and sand flies (Psychodidae, Diptera) sharing the same ecotopes in Brazilian Amazonia. J. Eukaryot. Microbiol. 55, 427–435.
- Fraga, J., Montalvo, A.M., De Doncker, S., Dujardin, J.C., Van der Auwera, G., 2010. Phylogeny of *Leishmania* species based on the heat-shock protein 70 gene. Infect. Genet. Evol. 10, 238–245.
- Garrido, G., Arribas, A., 2008. Canis accitanus nov. sp., a new small dog (Canidae, Carnivora, Mammalia) from the Fonelas P-1 Plio–Pleistocene site (Guadix basin, Granada, Spain). Gé obios 41, 751–761.
- Gibson-Corley, K.N., Hostetter, J.M., Hostetter, S.J., Mullin, K., Ramer-Tait, A.E., Boggiatto, P.M., Petersen, C.A., 2008. Disseminated *Leishmania infantum* infection in two sibling foxhounds due to possible vertical transmission. Can. Vet. J. 49, 1005–1008.
- Giger, U., Oakley, D.A., Owens, S.D., Schantz, P., 2002. Leishmania donovani transmission by packed RBC transfusion to anemic dogs in the United States. Transfusion 42, 381–383.
- Gould, I.T., Perner, M.S., Santini, M.S., Saavedra, S.B., Bezzi, G., Maglianese, M.I., Antman, J.G., Gutierrez, J.A., Salomon, O.D., 2013. Visceral leishmaniasis in Argentina. Cases notification and distribution of vectors (2006–2012). Medicina (B Aires) 73, 104–110.
- Hamilton, P.B., 2012. Is *Trypanosoma vivax* genetically diverse? Trends Parasitol. 28, 173.
- Hamilton, P.B., Stevens, J.R., Gaunt, M.W., Gidley, J., Gibson, W.C., 2004. Trypanosomes are monophyletic: evidence from genes for glyceraldehyde phosphate dehydrogenase and small subunit ribosomal RNA. Int. J. Parasitol. 34, 1393–1404.
- Hamilton, P.B., Gibson, W.C., Stevens, J.R., 2007. Patterns of co-evolution between trypanosomes and their hosts deduced from ribosomal RNA and protein-coding gene phylogenies. Mol. Phylogenet. Evol. 44, 15–25.
- Hamilton, P.B., Teixeira, M.M., Stevens, J.R., 2012. The evolution of *Trypanosoma cruzi*: the 'bat seeding' hypothesis. Trends Parasitol. 28, 136–141.
- Hannaert, V., Opperdoes, F.R., Michels, P.A., 1998. Comparison and evolutionary analysis of the glycosomal glyceraldehyde-3-phosphate dehydrogenase from different Kinetoplastida. J. Mol. Evol. 47, 728–738.
- Herrer, A., 1971. Leishmania hertigi sp. n., from the tropical porcupine, Coendou rothschildi Thomas. J. Parasitol. 57, 626–629.
- Hide, M., Bras-Goncalves, R., Banuls, A.L., 2007. Specific cpb copies within the *Leishmania donovani* complex: evolutionary interpretations and potential clinical implications in humans. Parasitology 134, 379–389.
- Huelsenbeck, J.P., Ronquist, F., 2001. MRBAYES: bayesian inference of phylogenetic trees. Bioinformatics 17, 754–755.
  Ibrahim, M.E., Barker, D.C., 2001. The origin and evolution of the *Leishmania*
- Ibrahim, M.E., Barker, D.C., 2001. The origin and evolution of the *Leishmania donovani* complex as inferred from a mitochondrial cytochrome oxidase II gene sequence. Infect. Genet. Evol. 1, 61–68.
- Kuhls, K., Mauricio, I.L., Pratlong, F., Presber, W., Schonian, G., 2005. Analysis of ribosomal DNA internal transcribed spacer sequences of the *Leishmania donovani* complex. Microbes Infect. 7, 1224–1234.
- Kuhls, K., Chicharro, C., Canavate, C., Cortes, S., Campino, L., Haralambous, C., Soteriadou, K., Pratlong, F., Dedet, J.P., Mauricio, I., Miles, M., Schaar, M., Ochsenreither, S., Radtke, O.A., Schonian, G., 2008. Differentiation and gene flow among European populations of *Leishmania infantum* MON-1. PLoS Negl. Trop. Dis. 2, e261.
- Kuhls, K., Alam, M.Z., Cupolillo, E., Ferreira, G.E., Mauricio, I.L., Oddone, R., Feliciangeli, M.D., Wirth, T., Miles, M.A., Schonian, G., 2011. Comparative microsatellite typing of new world *Leishmania infantum* reveals low heterogeneity among populations and its recent old world origin. PLoS Negl. Trop. Dis. 5, e1155.
- Lainson, R., 1997. On Leishmania enriettii and other enigmatic Leishmania species of the Neotropics. Mem. Inst. Oswaldo Cruz 92, 377–387.
- Lainson, R., Shaw, J.J., 1969. Some reservoir-hosts of *Leishmania* in wild animals of Mato Grosso State, Brazil. Two distinct strains of parasites isolated from man and rodents. Trans. R. Soc. Trop. Med. Hyg. 63, 408–409.
- Lainson, R., Shaw, J.J., 1988. Observations on the development of *Leishmania* (*L.*) *chagasi* Cunha and Chagas in the midgut of the sandfly vector *Lutzomyia longipalpis* (Lutz and Neiva). Ann. Parasitol. Hum. Comp. 63, 134–145.

- Lainson, R., Ryan, L., Shaw, J.J., 1987a. Infective stages of *Leishmania* in the sandfly vector and some observations on the mechanism of transmission. Mem. Inst. Oswaldo Cruz 82, 421–424.
- Lainson, R., Shaw, J.J., Silveira, F.T., Braga, R.R., 1987b. American visceral leishmaniasis: on the origin of *Leishmania* (*Leishmania*) chagasi. Trans. R. Soc. Trop. Med. Hyg. 81, 517.
- Lima, L., Silva, F.M., Neves, L., Attias, M., Takata, C.S., Campaner, M., de Souza, W., Hamilton, P.B., Teixeira, M.M., 2012. Evolutionary insights from bat trypanosomes: morphological, developmental and phylogenetic evidence of a new species, *Trypanosoma (Schizotrypanum) erneyi* sp. nov., in African bats closely related to *Trypanosoma (Schizotrypanum) cruzi* and allied species. Protist 163, 856–872.
- Linares, O.J., 1981. Tres nuevos carnívoros procionidos fosiles de Mioceno de Norte y Sudamérica. Ameginiana 18, 113–121.
- Lukes, J., Mauricio, I.L., Schonian, G., Dujardin, J.C., Soteriadou, K., Dedet, J.P., Kuhls, K., Tintaya, K.W., Jirku, M., Chocholova, E., Haralambous, C., Pratlong, F., Obornik, M., Horak, A., Ayala, F.J., Miles, M.A., 2007. Evolutionary and geographical history of the *Leishmania donovani* complex with a revision of current taxonomy. Proc. Natl. Acad. Sci. U.S.A. 104, 9375–9380.
- Luz, E., Giovannoni, M., Borba, A.M., 1967. Infecção de Lutzomyia monticola por Leishmania enriettii. Anais da Faculdade de Medicina da Universidade Federal do Paraná 9, 121–128.
- Machado, M.I., Milder, R.V., Pacheco, R.S., Silva, M., Braga, R.R., Lainson, R., 1994. Naturally acquired infections with *Leishmania enriettii* Muniz and Medina 1948 in guinea-pigs from Sao Paulo, Brazil. Parasitology 109 (Pt. 2), 135–138.
- Marcili, A., Lima, L., Cavazzana, M., Junqueira, A.C., Veludo, H.H., Maia Da Silva, F., Campaner, M., Paiva, F., Nunes, V.L., Teixeira, M.M., 2009a. A new genotype of *Trypanosoma cruzi* associated with bats evidenced by phylogenetic analyses using SSU rDNA, cytochrome b and Histone H2B genes and genotyping based on ITS1 rDNA. Parasitology 136, 641–655.
- Marcili, A., Lima, L., Valente, V.C., Valente, S.A., Batista, J.S., Junqueira, A.C., Souza, A.I., da Rosa, J.A., Campaner, M., Lewis, M.D., Llewellyn, M.S., Miles, M.A., Teixeira, M.M., 2009b. Comparative phylogeography of *Trypanosoma cruzi* TCIIc: new hosts, association with terrestrial ecotopes, and spatial clustering. Infect. Genet. Evol. 9, 1265–1274.
- Marcili, A., Valente, V.C., Valente, S.A., Junqueira, A.C., da Silva, F.M., Pinto, A.Y., Naiff, R.D., Campaner, M., Coura, J.R., Camargo, E.P., Miles, M.A., Teixeira, M.M., 2009c. Trypanosoma cruzi in Brazilian Amazonia: Lineages TCI and TCIIa in wild primates, Rhodnius spp. and in humans with Chagas disease associated with oral transmission. Int. J. Parasitol. 39, 615–623.
- Mauricio, I.L., Howard, M.K., Stothard, J.R., Miles, M.A., 1999. Genomic diversity in the *Leishmania donovani* complex. Parasitology 119 (Pt. 3), 237–246.
- Mauricio, I.L., Gaunt, M.W., Stothard, J.R., Miles, M.A., 2001. Genetic typing and phylogeny of the *Leishmania donovani* complex by restriction analysis of PCR amplified gp63 intergenic regions. Parasitology 122, 393–403.
- Mauricio, I.L., Stothard, J.R., Miles, M.A., 2004. Leishmania donovani complex: genotyping with the ribosomal internal transcribed spacer and the mini-exon. Parasitology 128, 263–267.
- Mauricio, I.L., Gaunt, M.W., Stothard, J.R., Miles, M.A., 2007. Glycoprotein 63 (gp63) genes show gene conversion and reveal the evolution of Old World Leishmania. Int. J. Parasitol. 37, 565–576.
- Momen, H., Cupolillo, E., 2000. Speculations on the origin and evolution of the genus Leishmania. Mem. Inst. Oswaldo Cruz 95, 583–588.
- Muniz, J., Medina, H., 1948. Hospital (Rio J) 33, 7-25.
- Nascimento, E.L., Martins, D.R., Monteiro, G.R., Barbosa, J.D., Ximenes, M.F., Maciel, B.L., Duarte, I., Jeronimo, S.M., 2008. Forum: geographic spread and urbanization of visceral leishmaniasis in Brazil. Postscript: new challenges in the epidemiology of *Leishmania chagasi* infection. Cad. Saude Publica 24, 2964–2967.
- Nicholas, K.B., Nicholas Jr., H.B., Deerfield, D.W., 1997. GeneDoc: Analysis and Visualization of Genetic Variation. EMBNEW NEWS 4, 14.
- Noyes, H.A., Arana, B.A., Chance, M.L., Maingon, R., 1997. The Leishmania hertigi (Kinetoplastida; Trypanosomatidae) complex and the lizard Leishmania: their classification and evidence for a neotropical origin of the Leishmania– Endotrypanum clade. J. Eukaryot. Microbiol. 44, 511–517.
- Noyes, H.A., Stevens, J.R., Teixeira, M., Phelan, J., Holz, P., 1999. A nested PCR for the ssrRNA gene detects *Trypanosoma binneyi* in the platypus and *Trypanosoma* sp. in wombats and kangaroos in Australia. Int. J. Parasitol. 29, 331–339.
- Noyes, H., Pratlong, F., Chance, M., Ellis, J., Lanotte, G., Dedet, J.P., 2002. A previously unclassified trypanosomatid responsible for human cutaneous lesions in Martinique (French West Indies) is the most divergent member of the genus *Leishmania* ssp. Parasitology 124, 17–24.
- Orlando, T.C., Rubio, M.A., Sturm, N.R., Campbell, D.A., Floeter-Winter, L.M., 2002. Intergenic and external transcribed spacers of ribosomal RNA genes in lizardinfecting Leishmania: molecular structure and phylogenetic relationship to mammal-infecting Leishmania in the subgenus *Leishmania (Leishmania)*. Mem. Inst. Oswaldo Cruz 97, 695–701.
- Owens, S.D., Oakley, D.A., Marryott, K., Hatchett, W., Walton, R., Nolan, T.J., Newton, A., Steurer, F., Schantz, P., Giger, U., 2001. Transmission of visceral leishmaniasis through blood transfusions from infected English foxhounds to anemic dogs. J. Am. Vet. Med. Assoc. 219, 1076–1083.
- Postigo, J.A., 2010. Leishmaniasis in the world health organization eastern mediterranean region. Int. J. Antimicrob. Agents 36 (Suppl. 1), S62–S65.
- Quispe Tintaya, K.W., Ying, X., Dedet, J.P., Rijal, S., De Bolle, X., Dujardin, J.C., 2004. Antigen genes for molecular epidemiology of leishmaniasis: polymorphism of cysteine proteinase B and surface metalloprotease glycoprotein 63 in the *Leishmania donovani* complex. J. Infect. Dis. 189, 1035–1043.

- Rioux, J.A., Ashford, R.W., Khiami, A., 1992. Ecoepidemiology of leishmaniases in Syria. 3. *Leishmania major* infection in *Psammomys obesus* provides clues to life history of the rodent and possible control measures. Ann. Parasitol. Hum. Comp. 67, 163–165.
- Rodrigues, A.C., Paiva, F., Campaner, M., Stevens, J.R., Noyes, H.A., Teixeira, M.M., 2006. Phylogeny of *Trypanosoma (Megatrypanum) theileri* and related trypanosomes reveals lineages of isolates associated with artiodactyl hosts diverging on SSU and ITS ribosomal sequences. Parasitology 132, 215–224.
- Romero, G.A., Boelaert, M., 2010. Control of visceral leishmaniasis in Latin America a systematic review. PLoS Negl. Trop. Dis. 4, e584.
- Rose, K., Curtis, J., Baldwin, T., Mathis, A., Kumar, B., Sakthianandeswaren, A., Spurck, T., Low Choy, J., Handman, E., 2004. Cutaneous leishmaniasis in red kangaroos: isolation and characterisation of the causative organisms. Int. J. Parasitol. 34, 655–664.
- Rosypal, A.C., Tripp, S., Kinlaw, C., Hailemariam, S., Tidwell, R.R., Lindsay, D.S., Rajapakse, R.P., Sreekumar, C., Dubey, J.P., 2010a. Surveillance for antibodies to *Leishmania* spp. in dogs from Sri Lanka. J. Parasitol. 96, 230–231.
- Rosypal, A.C., Tripp, S., Lewis, S., Francis, J., Stoskopf, M.K., Larsen, R.S., Lindsay, D.S., 2010b. Survey of antibodies to *Trypanosoma cruzi* and *Leishmania* spp. in gray and red fox populations from North Carolina and Virginia. J. Parasitol. 96, 1230– 1231.
- Salomon, O.D., Basmajdian, Y., Fernandez, M.S., Santini, M.S., 2011. Lutzomyia longipalpis in Uruguay: the first report and the potential of visceral leishmaniasis transmission. Mem. Inst. Oswaldo Cruz 106, 381–382.
- Schlein, Y., 1993. Leishmania and sandflies: interactions in the life cycle and transmission. Parasitol. Today 9, 255–258.
- Shaw, J.J., 1994. Taxonomy of the genus *Leishmania*: present and future trends and their implications. Mem. Inst. Oswaldo Cruz 89, 471–478.
- Shaw, J., 1997. RAPD for the differentiation of *Leishmania*. Trans. R. Soc. Trop. Med. Hyg. 91, 93–94.
- Silva, D.A., Madeira, M.D., Barbosa Filho, C.J., Schubach, E.Y., Barros, J.H., Figueiredo, F.B., 2013. *Leishmania (Leishmania) hertigi* in a porcupine (*Coendou* sp.) found in Brasilia, Federal District, Brazil. Rev. Bras. Parasitol. Vet. 22, 297–299.
- Sotnikova, M., 2006. A new canid Nurocyon chonokhariensis gen et sp. nov (Canini, Canidae, Mammalia) from the Pliocene of Mongolia. Courier Forschungsinstitut Senckenberg 256, 11–21.
- Sotnikova, M., Rook, L., 2010. Dispersal of the Canini (Mammalia, Canidae: Caninae) across Eurasia during the late Miocene to Early Pleistocene. Quaternary International. 212, 86–97.
- Spassov, N., Rook, L., 2006. Eucyon marinae sp. nov. (Mammalia, Carnivora), a new canid species from the Pliocene of Mongolia, with a review of forms referable to the genus. Rivista Italiana di Paleontologia e Stratigrafia 112, 123–133.
- Stevens, J., Rambaut, A., 2001. Evolutionary rate differences in trypanosomes. Infect. Genet. Evol. 1, 143–150.
- Stevens, J.R., Noyes, H.A., Dover, G.A., Gibson, W.C., 1999. The ancient and divergent origins of the human pathogenic trypanosomes, *Trypanosoma brucei* and *T. cruzi*. Parasitology 118 (Pt. 1), 107–116.
- Teixeira, M.M., Campaner, M., Camargo, E.P., 1994. Detection of trypanosomatid Phytomonas parasitic in plants by polymerase chain reaction amplification of small subunit ribosomal DNA. Parasitol. Res. 80, 512–516.
- Teixeira, M.M., Borghesan, T.C., Ferreira, R.C., Santos, M.A., Takata, C.S., Campaner, M., Nunes, V.L., Milder, R.V., de Souza, W., Camargo, E.P., 2011. Phylogenetic validation of the genera Angomonas and Strigomonas of trypanosomatids harboring bacterial endosymbionts with the description of new species of trypanosomatids and of proteobacterial symbionts. Protist 162, 503–524.
- Thomaz-Soccol, V., Pratlong, F., Langue, R., Castro, E., Luz, E., Dedet, J.P., 1996. New isolation of *Leishmania enriettii* Muniz and Medina, 1948 in Paranastate, Brazil, 50 years after the first description, and isoenzymatic polymorphism of the *L. enriettii* taxon. Ann. Trop. Med. Parasitol. 90, 491–495.
- Thompson, J.D., Gibson, T.J., Plewniak, F., Jeanmougin, F., Higgins, D.G., 1997. The CLUSTAL\_X windows interface: flexible strategies for multiple sequence alignment aided by quality analysis tools. Nucleic Acids Res. 25, 4876–4882.
- Uliana, S.R., Affonso, M.H., Camargo, E.P., Floeter-Winter, L.M., 1991. Leishmania: genus identification based on a specific sequence of the 18S ribosomal RNA sequence. Exp. Parasitol. 72, 157–163.
- van Eys, G.J., Schoone, G.J., Kroon, N.C., Ebeling, S.B., 1992. Sequence analysis of small subunit ribosomal RNA genes and its use for detection and identification of *Leishmania* parasites. Mol. Biochem. Parasitol. 51, 133–142.
- Viola, L.B., Campaner, M., Takata, C.S., Ferreira, R.C., Rodrigues, A.C., Freitas, R.A., Duarte, M.R., Grego, K.F., Barrett, T.V., Camargo, E.P., Teixeira, M.M., 2008. Phylogeny of snake trypanosomes inferred by SSU rDNA sequences, their possible transmission by phlebotomines, and taxonomic appraisal by molecular, cross-infection and morphological analysis. Parasitology 135, 595– 605.
- Viola, L.B., Attias, M., Takata, C.S., Campaner, M., De Souza, W., Camargo, E.P., Teixeira, M.M., 2009. Phylogenetic analyses based on small subunit rRNA and glycosomal glyceraldehyde-3-phosphate dehydrogenase genes and ultrastructural characterization of two snake Trypanosomes: *Trypanosoma* serpentis n. sp. from *Pseudoboa nigra* and *Trypanosoma* cascavelli from *Crotalus* durissus terrificus. J. Eukaryot. Microbiol. 56, 594–602.
- Wang, X., Tedford, R.H., 2007. The behavioral biology of dogs. In: Jensen, Per (Ed.), Evolutionary History of Canids. Cromwell Press, Trowbridge, UK, pp. 1–20.
- Wilgenbusch, J.C., Swofford, D., 2003. Inferring evolutionary trees with PAUP\*. Curr. Protoc. Bioinform. (Chapter 6, Unit 6 4).
- Zeledon, R., Ponce, C.E.D.E.P., 1977. Finding of *Leishmania hertigi* in the Costa Rican porcupine. J. Parasitol. 63, 924–925.