Leishmania donovani is the only cause of visceral leishmaniasis in East Africa; previous descriptions of L. infantum and "L. archibaldi" from this region are a consequence of convergent evolution in the isoenzyme data

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SUMMARY

Isoenzyme-based studies have identified 3 taxa/species/'phylogenetic complexes' as agents of visceral leishmaniasis in Sudan: L. donovani, L. infantum and "L. archibaldi". However, these observations remain controversial. A new chitinase gene phylogeny was constructed in which stocks of all 3 putative species isolated in Sudan formed a monophyletic clade. In order to construct a more robust classification of the L. donovani complex, a panel of 16 microsatellite markers was used to describe 39 stocks of these 3 species. All "L. donovani complex" stocks from Sudan were again found to form a single monophyletic clade. L. donovani ss stocks from India and Kenya were found to form 2 region-specific clades. The partial sequence of the glutamate oxaloacetate transaminase (GOT) gene of 17 L. donovani complex stocks was obtained. A single nucleotide polymorphism in the GOT gene appeared to underlie the isoenzyme classification. It was concluded that isoenzyme-based identification is unsafe for stocks isolated in L. donovani endernic areas and identified as L. infantum. It was also concluded that the name L. archibaldi is invalid and that only a single visceralizing species, Leishmania donovani, is found in East Africa.

Key words: aspartate aminotransferase, microsatellites, MLEE, kala azar, Ethiopia, chagasi.

INTRODUCTION

Visceral leishmaniasis (VL) is caused by members of the "Leishmania donovani complex" (Lainson & Shaw, 1987). The 2 main members are L. donovani (Laveran & Mesnil, 1903) in the Old World and L. infantum Nicolle, 1908 in the Old World and New World (L. chagasi is a junior synonym of L. infantum (Mauricio, Stothard & Miles, 2000)). "L. archibaldi" is a possible third member of the group that is found in Sudan (Pratlong et al. 2001). These 3 taxa are morphologically indistinguishable but are associated with different epidemiology, ecology and pathology. L. infantum and L. donovani have largely discrete geographical distributions; L. infantum is present around the Mediterranean basin and also in the New World, where it was probably introduced by

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European colonists (Courtenay et al. 2002; Mauricio et al. 2000). L. donovani is found in the Indian subcontinent and East Africa where it is associated with epidemics in adults as well as children. L. donovani is believed to be largely anthroponotic, although a species of mongoose has been found infected in Sudan (Elnaiem et al. 2001) and Dereure et al. (2003) reported a high (>70%) seroprevalence in dogs during 1998 and 1999 in one village in eastern Sudan. Dogs are the principal reservoir host for L. infantum but visceral disease in immunocompetent humans occurs only sporadically and almost exclusively in children (Ashford, 2000).

Multilocus enzyme electrophoresis-based studies (MLEE) have suggested that there may be some overlap in the range of *L. donovani* and *L. infantum*, particularly in East Africa where *L. infantum* is occasionally identified despite the predominance of *L. donovani* (Pratlong et al. 2001).

The classification of parasites from East Africa is further complicated by the supposed presence of "L. archibaldi". The name "Leishmania donovani varietas archibaldi" was originally applied to Leishmania from Sudan on dubious grounds by Castellani & Chalmers (1919). Brumpt (1936) later

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included L. donovani var. archibaldi as a synonym of L. donovani. Nicoli (1963) subsequently revised the genus Leishmania and treated 'var. archibaldi' as the subspecies L. donovani archibaldi. He applied the name to parasites from throughout East Africa and suggested, on the basis of its occurrence in rodents, that it should probably have specific status, even calling it L. archibaldi in the same article. The name "L. archibaldi" was more formally, but still provisionally, used by Lainson & Shaw (1987), who applied it to VL parasites from Sudan, suggesting that the parasite in Kenya and Somalia might be different.

With the advent of MLEE the possibility arose for a more objective taxonomic study of these parasites. Moreno et al. (1986) found that stocks that corresponded biologically with L. donovani could be distinguished from those corresponding with L. infantum on the basis of numerical analysis of isoenzyme data. The recent taxonomic re-descriptions of the L. donovani complex are, however, essentially based upon differences at a single 'diagnostic' enzyme locus since an inspection of the raw data shows that only glutamate oxaloacetate transaminase (GOT), segregates between these phenotypes. GOT₁₀₀ was assigned to L. infantum, which was considered to be invariant for this allele (Moreno et al. 1986; Rioux et al. 1990). The L. donovani GOT enzyme variant had a mobility 13% greater than the L. infantum allele and consequently was designated GOT_{113} . All L. donovani stocks were believed to have the GOT113 allele. Thus the mobility of alleles of the GOT enzyme was considered sufficient to distinguish between these 2 species.

When a series of stocks became available from Sudan, it was found that both the L. donovani and L. infantum GOT character-states were represented, as well as a third state-GOT₁₁₀. Rioux et al. (1990) used the name L. archibaldi for the zymodeme showing this new character-state, and produced a classification of Leishmania, which included L. archibaldi as a 'phylogenetic complex', defined by its GOT mobility. A total of 3 L. archibaldi zymodemes have now been described, all with the GOT₁₁₀ allele but with variations at other enzyme loci. Other Sudanese stocks were identified as L. donovani and L. infantum according to their respective GOT mobilities (Pratlong et al. 2001).

Thus the name archibaldi has been used by Rioux and co-workers' for stocks with the GOT_{In0} allele, while Castellani and Chalmers applied the name to all Sudanese parasites, Nicoli used it for all east African parasites, and Lainson and Shaw applied it to all parasites from Sudan with possibly another taxon being responsible for leishmaniasis in Kenya and Somalia. The World Health Organization (Anon, 1990) included L. archibaldi in their list of Leishmania species, but did not indicate whether they used it in the sense of Lainson and Shaw, or

that of Rioux et al., or indeed of any of the other workers.

While isoenzyme-based studies have indicated the presence of L. infantum and "L. archibaldi" in Sudan (Pratlong et al. 2001), this contrasts with DNAbased studies using Gp63, Internal Transcribed Spacer (ITS) and anonymous genomic sequences, that have found only a single clade of parasites in Sudan (El-Tai et al. 2001; Mauricio et al. 2001), and with sequences of anonymous genomic DNA, which failed to segregate with isoenzyme-based species (Lewin et al. 2002). Phylogeographical studies are usually conducted using neutral markers such as the housekeeping genes used for MLEE, however functional genes can also help reveal the effects of selective pressures. A PCR-RFLP study of the Gp63 gene (which is involved in invasion of the macrophage) of the L. donovani complex indicated geographical clustering that did not correlate with MLEE data (Guerbouj et al. 2001). In an attempt to clarify the classification of Leishmania from East Africa we have used the phylogeny of the chitinase gene which is involved in the interaction between the sandfly and the parasite and we have also developed a panel of 20 evolutionarily neutral microsatellite markers for L. infantum and L. donovani (Jamjoom et al. 2002b). Microsatellites are the mainstay of modern genetic studies and have largely replaced isoenzymes for many applications from individualbased studies through to shallow phylogenies (Sunnucks, 2000). Because they are PCR-based and allelic variants vary in length, microsatellite loci are well adapted for analysing large numbers of samples from small amounts of DNA and the data are, in principle, easily exchanged between laboratories. Most microsatellite loci are believed to be selectively neutral and are typically highly polymorphic. This makes it generally possible to discover more genetic diversity using microsatellites than with MLEE. We have now applied 16 of these markers to a collection of 41 stocks from Sudan, Kenya and India together with 5 L. infantum controls from Brazil, Spain, Algeria, and the UK (introduced). We have also sequenced parts of the chitinase and GOT genes in order to confirm our findings.

MATERIALS AND METHODS

Parasites

The stocks used in this study are listed in Table 1.

Sequencing the chitinase gene

The sequences of the L. donovani (AF009354) and L. mexicana chitinase genes were used to identify conserved regions of this gene and to design 2 pairs of primers, ChitA forward and reverse and ChitB forward and reverse (Table 2). PCR was carried out in

25 µl volumes in 0.2 ml PCR tubes with each reaction containing 2.0 mm MgCl₂, 75 mm Tris-HCl, 20 mm $(NH)_4SO_4$, 0.01% (v/v) Tween 20, 0.4 mM each dNTP, 25 ng genomic DNA, 40 pmol ChitA or ChitB primers and 2.5 units of Taq polymerase (ABgene). Thermal cycling conditions were: 94 °C for 5 min, then 30 times (94 °C for 60 sec, 58 °C for 60 sec and 72 °C for 60 sec) and finally 72 °C for 10 min. PCR products were cloned into pCR[®]-2.1-TOPO® (Invitrogen). The chitinase sequences were determined by cycle sequencing using CEQ DTCS chemistry (Beckman Coulter Inc.) and electrophoresis on a CEQ8000XL capillary sequencer (Beckman Coulter Inc). Sequences were aligned using MegAlign in the DNASTAR package and trees were compiled using DNAPARS and DNA-DIST in PHYLIP.

Microsatellite primers

A full description of the microsatellite loci and their PCR conditions is given in detail by Jamjoom et al. (2002b) except that for this work the forward primer for each locus was 5'-labelled with either D3 or D4 fluorescent dyes (PrOligo or Research Genetics) for visualization (Table 2). PCR products were pooled, according to their variation in allelic size and the fluorescent dye, along with a 400 base-pair (D1) size standard (Beckman Coulter Inc.) and separated by capillary electrophoresis through a denaturing acrylamide gel on a Ceq8000XL automated sequencer (Beckman Coulter Inc.). Microsatellite alleles were sized using the cubic model in the Ceq8000XL fragment analysis software. Alleles were converted to presence/absence data and the resultant table was used to construct phenetic trees using distance and Parsimony methods in PAUP 4.10b (Swofford, 1998). Deviation from genotypic linkage equilibrium among the entire set of locus-pair combinations was analysed using Fisher's exact test as implemented by GENEPOP v.3.3 (Raymond & Rousset, 1995).

Sequencing the GOT gene

Glutamate oxaloacetate transaminase is also known as aspartate aminotransferase. There are 2 aspartate aminotransferases in *T. brucei*, a nuclear encoded cytoplasmic enzyme and a nuclear encoded mitochondrial enzyme (Berger et al. 2001). The 2 genes have no significant homology as judged by BLAST search of one against the other or attempts to align them (HAN, unpublished observations). However when used as probes in a BLAST search of kinetoplastid sequences in GeneDB (http://www.genedb.org/), they both hit the same sequence in clone AC078900 from *L. major* chromosome 35, which contains the aspartate aminotransferase gene annotated as LmjF35.0820 in GeneDB. This

sequence was used to design the following primers using the Primer3 program (Rozen & Skaletsky, 2000): GOT_1F 10-29, GOT_5R 1259-1277, and GOT_7R 1190-1211 (Table 2). The numbers after each primer name indicate the position of the primer in the predicted GOT gene. The primer pairs GOT_1F and GOT_5R and GOT_1F and GOT_7R both gave PCR products which were cloned into the pGEM EASY vector (Promega), cycle sequenced with the BigDyeV3.1 kit (ABI) and sequences were obtained on an ABI3100 capillary sequencer. Sequences were aligned in the SeqMan package of DNASTAR and scanned for polymorphisms by eye. A single nucleotide T→G polymorphism at position 823 was typed by direct sequencing of the PCR product of the primers GOT_SNP_F GOT_SNP_R (Table 2).

RESULTS

Comparison of the chitinase gene sequence of the New World species L. mexicana with that of L. donovani revealed significant differences (93% identity). Therefore, this gene was tested for suitability for an intraspecific classification of the "L. donovani complex". Parsimony informative sites (199) were identified within the 1028 bp alignment, a classification was produced of 27 stocks (Fig. 1) and the species complexes could be identified with confidence. Both parsimony and distance methods produced trees with identical topology and only small differences (<8) in bootstrap values. There were 8 informative sites within the L. donovani complex and relationships within this group were less well supported. All stocks from Kenya and Sudan fell into 2 clear geographically defined groups with 56% and 58% bootstrap support, respectively. Stocks from Ethiopia and the Indian subcontinent clustered with the Sudanese stocks with 91% bootstrap support. L. infantum from Europe and the Mediterranean formed a clade with 85% support. "L. infantum" from Sudan clustered tightly with other Sudanese isolates and not with L. infantum from Europe and Mediterranean. Stocks labelled "L. archibaldi" were indistinguishable from other stocks from Sudan and did not form a distinct clade. The 8 informative sites in the alignment formed 4 haplotypes (Table 3). These haplotypes showed that a Kenyan L. donovani and Mediterranean L. infantum clade was supported by 5 positions, the L. infantum clade was supported by 2 positions and the Sudanese clade combining L. donovani, "L. infantum" and "L. archibaldi" stocks was supported by just a single position.

Although the chitinase data described above gave a classification that correlated with geography and biology better than classifications based on MLEE, the amount of the data underlying this conclusion was limited. Therefore, data from presence/absence of alleles at 16 microsatellite loci for 41 stocks of

Table 1. Leishmania stocks

(Leishmania stocks used in the study. The first column shows the putative species of the studied stocks, species names based on MLEE but reassigned in this study are shown in inverted commas. The second column presents the WHO code. This indicates host: MHOM Humans; IDUB Phlebotomus duboscqi; IMRT Phlebotomus martini; MCAN Canis familiaris, MNYC Nyctomys sp.; MRHO Rhombomys sp./Country; BD Bangladesh; BR Brazil; BZ Belize; DZ Algeria; ES Spain; ET Ethiopia; GB Great Britain; IL Lebanon; IN India; IQ Iraq; IR Iran, IT Italy; KE Kenya; NP Nepal; SD Sudan; SN Senegal; SU Soviet Union; XX Unknown/Year of isolation/Laboratory code, if 2 or more laboratories have given codes then these are separated by a semi colon. The third column is the zymodeme where known. Zymodemes are indicated as typed by the Laboratoire d'Ecologie Medicale et Parhotogie Parasitaire (LEM), Montpellier (MON), n.d. indicates zymodeme is not determined. The fourth column is the clinical disease indicated by CL for cutaneous leishmaniasis and VL for visceral leishmaniasis. The fifth column shows the GOT⁶²³ genotype as determined by direct sequencing of PCR products, the traces from these genes can be seen at www.genomics.liv.ac.uk/animal/supplementary_data/jamjoom/traces.pdf. The sixth column shows the studies completed for each stock: Ch, Chitinase gene sequence and GenBank Accession Number; *indicates a previous WHO reference strain (Chance & Walton, 1982).)

Species	WHO code	Zymodeme	Discase	GOT ⁶²³ Genotype	Analysis performed
The state of the s	NATIONA/II /80/Eco.dlin. St. 1	£ 6	1.)		(% (AV\$18223) Me
to, major	MED ELO/07/11/50/1	MON.4	3 5		Ch (AV518225), We
t. mado	NATIONALIZATION TORK TOOL TO		3 5	2	CE (11. (12.2.2.), 11.5.
L. major	MILCHAISO/00/LV330; LKC-L30	11.d.	3 5	5	CII. (A I 316220), IMS.
L. major	1DUB/SN/XX/LV399;DK3/	n.d.	5		Cn. (A1518227), ivis.
L. major	MHOM/SN/XX/LV622;DK72	n.d.	CF		Ch. (AY518228), Ms.
L.major	MRHO/IR/75/ER	n.d.	$C\Gamma$		Ch. (AY518229), Ms.
L, tropica	MHOM/SU/60/LV357;LRC-L39*	n.d.	CT	99	Ch. (AY518230), Ms.
L. tropica	MHOM/IQ/66/LV556*	n.d.	CL		Ch.(AY518258), Ms.
L. donovani	MHOM/BD/97/LDON;BG1	MON-2	VL		Ch. (AY518231), Ms., GOT (AY529106)
L. donovani	MHOM/ET/67/HU3;LV9*	MON-18	VL,	99	Ch. (AY518233), Ms., GOT (AY529111)
L. donovani	MHOM/SD/90/D83;LEM2131	MON-18	VL	99	Ch. (AY518234), Ms., GOT (AY529105)
L. donovani	MHOM/SD/90/D92;LEM2132	MON-18	VL	99	Ch. (AY518235), Ms., GOT (AY529107)
L. donovani	MHOM/SD/90/2828;LEM2139	MON-18	٧Ľ	GG	Ch. (AY518236), Ms., GOT (AY529107)
L. donovani	MHOM/SD/90/D100;LEM2140	MON-18	VĽ	99	Ch. (AY518237), Ms., GOT
L. donovani	MHOM/SD/92/18	MON-18	VL		Ms.
L. donovani	MHOM/IN/00/Devi;LEM138	MON-2	VI.		Ms.
L. donovani	MHOM/KE/75/H9;LEM496	MON-32	VL		Ch. (AY518238), Ms.
L. donovani	MHOM/KE/55/LRC-L53;LEM707	MON-36	VL		Ms.
L. donovani	IMRT/KE/62/LRC-L57;LEM719	MON-37	VIL		Ch. (AY518239), Ms.
L. donovani	MHOM/ET/84/ADDIS164;LEM980	MON-83	VL		Ch. (AY518240), Ms.
L. donovani	MHOM/KE/73/MRC74	LON-51	VL		Ms,
L. donovani	MHOM/ET/00/Hussen	LON-42	VL		Ch. (AY518241), Ms.
І., допочані	MHOM/NP/02/NEP2	n,d,	VL		Ch. (AY518242), Ms., GOT (AY529112)
L. donovani	MHOM/NP/02/NEP3	n.d.	VI.		Ch. (AY518243), Ms., GOT (AY529112)
L. donovani	MHOM/NP/02/NEP5	n,d.	VL		Ch. (AY518244), Ms., GOT (AY529114)
І., допочані	MHOM/1N/75/K13;LV613	n,d.	VL		Ms.
L. donovani	MHOM/IN/77/Munni;LV614	n.d.	VL		Ms,
L. donovani	MHOM/IN/77/Agindra;LV615	n.d.	۸F		Ms.
L. donotsani	MHOM/IN/77/Pandey;LV616	n,d.	VL		Ms.
L. donovani	MHOM/IN/77/Shibchandra;LV617	n.d.	VL		Ms.
L. donovani I Amazani	MHOM/IN/77/Devi;LV619 MHOM/IN/77/Mand-iba-1 V621	n.d.	AL VI		Ms.
L. aonoemn	1911 1C/191/ 118/ 1 / 1910 19 19 19 19 19 19 19 19 19 19 19 19 19	1.0.	7 >		IVIS.

donovani.	MHOM/IN/77/Rai;LV636	n,d.	VL		Ms.
donovani "L. archibaldi"	MHOM/SD/90/D75;LEM2134	MON-82	VL		Ch. (AY518245), Ms.
donovani "L. archibaldi"	MHOM/SD/90/2655;LEM2135	MON-82	VL	$_{ m GT}$	Ch. (AY518246), Ms., GOT (AY529109)
donovani "L. archibaldi"	MHOM/SD/90/D99;LEM2137	MON-82	VL	GT	Ch. (AY518247), Ms.
., donovani "L. archibaldi"	MFIOM/SD/91/D1783;LEM2211	MON-82	VL	GT	Ch. (AY518248), Ms.
donovani "L. archibaldi"	MHOM/SD/92/82	MON-82	VL		Ch. (AY518232), Ms.
. infantum	MCAN/GB/96/LV755	MON-1	VL	LL	Ch. (AY518249), Ms.
, infantum	MHOM/BR/76/150406;M4192	MON-1	VL		Ch. (AY518250), Ms.
infantum	MCAN/ES/98/LEM-935;JPC;M5	MON-1	VL	$_{ m LL}$	Ch. (AY518251), Ms., GOT (AY529110)
donovani "L. infantum"	MHOM/SD/91/D1809;LEM2213	MON-30	VL	TT	Ch. (AY518252), Ms., GOT (AY529115)
donovani "L. infantum"	MHOM/SD/97/LEM3435	MON-30	VL	LL	Ch. (AY518253), Ms., GOT (AY529116)
donovani "L. infantum"	MHOM/SD/97/LEM3431	MON-30	VL	TT	Ch. (AY518254), Ms., GOT
donovam "L. infantum"	MHOM/SD/97/LEM3441	MON-30	VL	$_{ m LL}$	Ch. (AY518255), Ms., GOT
donovani "L. infantum"	MHOM/SD/92/30	MON-30	VL		Ch. (AY518256), Ms.
infantum	MHOM/ES/81/BCN1;LEM307	MON-29	VL		Ms.
. infantum	MHOM/DZ/83/;LEM425	MON-80	VL		Ms,
infantum	MHOM/IT/85/ISS175;LEM1733	MON-111	VL		Ms.
., amazonensis	MHOM/BR/97/M2269	n,d.	$C\Gamma$	99	Ch. (AY518257), GOT
допоглат	MHOM/SD/62/1-S		VL		Ch., AF009354
mexicana	MNYC/BZ/62/M379		$_{\rm CF}$		Ch., AY572789
The state of the s	And the second s				

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L. infantum, L. donovani and "L. archibaldi" were generated and analysed by parsimony (Fig. 2) and minimum evolution using mean character differences. There were only 3 minor differences between the parsimony and minimum evolution trees, all within the large Sudanese clade, these were: (1) LEM980 from western Ethiopia clustered inside the Sudanese clade; (2) the polytomy at the root of the Sudanese clade in the parsimony tree was resolved into 2 clades in the minimum evolution tree but the grouping of taxa was consistent between the 2 trees; (3) LEM2211 could be resolved from the large polytomy that is adjacent to the parsimony tree but not in the mean distance tree. A table of all the microsatellite data is available at www.genomics.liv.ac.uk/ animal/supplementary_data/jamjoom/ms_data.xls. All stocks from the Indian subcontinent, Kenya and Sudan fell into 3 clear geographically defined groups with 85-100% bootstrap support by parsimony and 98-100% support by the distance method. Classical L. infantum stocks from Europe, Brazil and Algeria formed a fourth discrete clade with 96-100% bootstrap support. Once again all stocks from Sudan clustered in a single clade. The "L. infantum" from Sudan clustered tightly with other Sudanese isolates and not with other L. infantum from Europe or Brazil. Stocks labelled "L. archibaldi" were indistinguishable from other stocks from Sudan and did not form a distinct clade. Two wellsupported clades were identifiable within the 17 Sudanese stocks. Members of all 3 putative species were found in each of the clades.

Since the tree was not rooted, it is only possible to draw limited conclusions about the relationships between the geographical clades. The Indian and Kenyan clades clustered together with 77% and 89% bootstrap support. However, no rooting point is possible that would create the clade that was observed in the chitinase phylogeny composed of the Kenyan L. donovani and the classical L. infantum stocks. Consequently the two phylogenies are inconsistent in this respect.

The microsatellite dataset was scanned for loci that might be in linkage disequilibrium and hence on the same chromosome. Table 4 shows the locus pairs for which there was evidence of disequilibrium but after applying a Bonferoni correction for the large number of tests involved only LIST7-037 × LIST7-035 remained significant.

Although 16 microsatellite loci were used in this classification it is possible to identify members of the different clades with fewer microsatellite markers. Table 5 shows the microsatellite alleles from 3 microsatellite loci that segregated with specific geographical regions.

As noted in the Introduction, the MLEE-based assignment of stocks to named species within the "L. donovani complex" is based exclusively on the mobility of the enzyme GOT. Therefore, to examine

Table 2. Sequences of 16 primer pairs for polymorphic, dinucleotide microsatellite loci (Jamjoom et al. 2002b) and primers used to amplify and sequence regions of the chitinase and GOT genes in the Leishmania donovani 'complex'

(Also shown are single nucleotide polymorphism primers GOT_SNP. Dye refers to the fluorescent label used for allelic visualization on a CEQ8000XL (Beckman Coulter Inc.).)

Locus	5'Dye	Forward	Reverse
LIST7-021	D4.	CCGAATACACAAGCCTCCTC	TCAGGCTTCGTCGTTTCTTT
LIST7-022	D3	GTCGCTCTGTCTCTGTGTGC	TCCGCATTTTCCTCTCCTT
LIST7-023	D3	CTTTGCGTTGCGCACTAA	GCTTGTGTTCCGTGTGTGTT
LIST7-024	D4	TAAACTGCATGGTCCCCTCT	ACAAGCACCATCATCCACAT
LIST7-025	D3	GGAGTCGTCTCTCTGTTACGC	ATCGCGTGCATGGGTATT
LIST7-027	D4	CTCTCTCGTCACCACAGCAC	AGGGGACAAGACACAGATGG
LIST7-028	D3	CACTCCACTGCGTTGGATA	CTTTGACCGCCGTTCTTT
LIST7-029	D4	GCAGAGCTTCTGCTTGGATT	GCATTGCTGTTCTCATCCAC
LIST7-031	D3	CACTGGTGGAAATAGAAAGACT	GGAGAACTAAAACGAGCAGCA
LIST7-032	D3	CTAGAGGCGTGCGGATGTA .	TCGCAGTTTTCGGTCCA
L1ST7-033	D4	CATTGCTGAGTGCTGCTAGTG	ATGAGCGTACTGGGCACAC
LIST7-035	D3	AAAGGTATGATACGCCTGTGG	ACCGCAAAGAACGGACAT
LIST7-036	D4	CTCTCTCGTCACCACAGCAC	TCCCTCTCGTTGGTGAAGTT
LIST7-037	D4	ATGCTGAGCCCATCAAGACT	GATGTCCCCGTTTACTCCAA
LIST7-039	D3	CTCGCACTCTTTCGCTCTTT	AGACGAGAGGAACGGAAAA
LIST7-040	D4	GCAGAGCGAGACACACAGAC	GTGCACGTTGATGTGCTTCT
ChitA		TTCTGCAGCTGGCGTGTCTTGTA	TGAGCTTGCGGCGGTGGTCCTG
ChitB		TCCTTCCCCGCTCTTTACTGTCTT	TACCCGTCATACTCGCTCAAT
GOT_1F 10-29		CCCATCACTCACGATTCACA	
GOT_5R 1259			AGCACAGATGTCCACGCA
GOT_7R 1190-1211		•	GAAGATTCAGGCACAAGCTCCC
GOT_SNP		CGAATCCATCACGCTCTTTAC	TGTTCTTCGACTCCGCCTAC

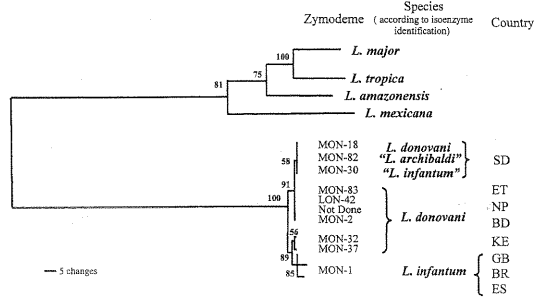


Fig. 1. Phylogeny based on partial sequencing (1028 nucleotides) of the chitinase gene. The phylogeny was compiled using the parsimony option in PAUP and bootstrap values were compiled using 100 bootstrap replicates. "Leishmania infantum" and "Leishmania archibaldi" from Sudan clearly cluster with Leishmania donovani and not with classical Leishmania infantum zymodeme MON1. The taxa included are indicated in Table 1, together with the GenBank Accession Numbers of the sequences. Country abbreviations are as in the legend to Table 1.

the validity of this distinction, the sequence of 1127 of the 1260 base pairs of this enzyme was obtained for 16 of the stocks under investigation (Table 1). Within this sequence only a single informative polymorph-

ism was identified that give rise to an amino acid change. This $T \rightarrow G$ polymorphism at position 823 would cause a change in amino acid 275 from tyrosine in the GOT_{100} ("L. infantum") stocks to aspartate in

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Table 3. Summary of the chitinase haplotypes within the *Leishmania donovani* 'complex' (The number of stocks sequenced in each group is indicated in parentheses.)

	Positio	on of the p	oolymorphi	sm in the cl	nitinase gene			
Stocks	129	158	212	315	435	663	847	1034
Sudanese (14) and Ethiopian (1)	Т	С	G	G	T	С	С	A
Kenya (2)	C	C	A	\mathbf{T}	C	G	T	A
L. infantum (MON-1) (3)	C	\mathbf{T}	\mathbf{A}	\mathbf{T}	C	Ġ	${f ilde{T}}$	Ĝ
Other stocks (Ethiopia, Nepal, Bangladesh) (6)	С	C.	G	G	Т	С	C	Ā
	Sudanese	L. infantum		Kenya	and L . $infar$	ntum		L. infantum

the GOT113 ("L. donovani") stocks. The additional negative charge provided by the aspartate residue in such L. donovani stocks would be expected to cause the GOT of these stocks to migrate faster towards the anode and consequently give rise to the faster observed mobility of GOT. All stocks that had been identified as L. infantum by MLEE had the 275tyr allele including 2 "L. infantum" stocks from Sudan (LEM2213 and LEM3435). Three stocks that had been identified as "L. archibaldi" by MLEE had the 275^{tyr} allele whilst one had the 275^{asp}. Eighteen stocks were sequenced directly from PCR products (3 "L. archibaldi", 6 L. infantum, 6 L. donovani, 1 L. tropica L. major and 1 L. amazonensis). All 6 L. infantum as identified by MLEE had the "T" allele, all 6 L. donovani had the "G" allele and all 3 "L. archibaldi" were heterozygotes (Table 1). The other 3 species typed had the "G" allele suggesting that this is the primitive condition. Electropherograms of these data are available from www.genomics.liv.ac.uk/animal/supplementary_data /jamjoom/traces.pdf.

DISCUSSION

Validity of "L. archibaldi" and L. infantum in Sudan

The chitinase sequence-based classification and the microsatellite-based classification both show that Leishmania from Sudan form a single clade. This is inconsistent with the present, isoenzyme-based classification which indicates that parasites from Sudan belong to 3 distinct clades, but consistent with previous DNA-based studies (El-Tai et al. 2001; Mauricio et al. 2001). Our GOT data show that the isoenzyme-based identification of L. donovani and L. infantum is dependent on a single nucleotide polymorphism and confirms previous observations made by Mauricio and Miles at the London School of Hygiene and Tropical Medicine (Mauricio personal

communication). Since both the $275^{\rm tyr}$ and the $275^{\rm asp}$ alleles of GOT are clearly viable, it is quite conceivable that the $T \rightarrow G$ transition that underlies the amino acid polymorphism could have occurred independently on more than one occasion. Such an independent mutation could have given rise to the "L. infantum" stocks found in Sudan quite separately from the same mutation that gave rise to the L. infantum around the Mediterranean. Consequently although GOT_{100} may be an adequate marker for identifying L. infantum and L. donovani in the Mediterranean, India and the New World, it is not suitable for this purpose in Africa.

We therefore propose that all stocks that have been identified by MLEE as L. infantum and that were isolated within the L. donovani endemic area of East Africa should be renamed, as L. donovani. We further suggest that the species "L. archibaldi" should be considered invalid, since it did not form a monophyletic clade; even if L. archibaldi does form a monophyletic clade within the Sudanese L. donovani, this would render L. donovani itself polyphyletic and hence invalid.

Further, if genuine L. infantum does exist in Sudan then it might be expected that parasites identified as L. infantum would have a similar epidemiology to those around the Mediterranean. A recent study of zymodemes of parasites isolated from 52 humans and 20 dogs in Sudan found no epidemiological difference in the diseases caused by the 3 putative species (Dereure et al. 2003). However, reanalysis of the published data shows that a significantly larger proportion of dogs were infected with L. infantum zymodeme parasites than L. donovani zymodeme parasites (P<0.001, Chi Squared test). Although this would appear to support the MLEE-based hypothesis that there is L. infantum in East Africa, we do not believe that this observation is sufficient evidence in itself to warrant the retention of species status for L. infantum zymodeme parasites in East Africa.

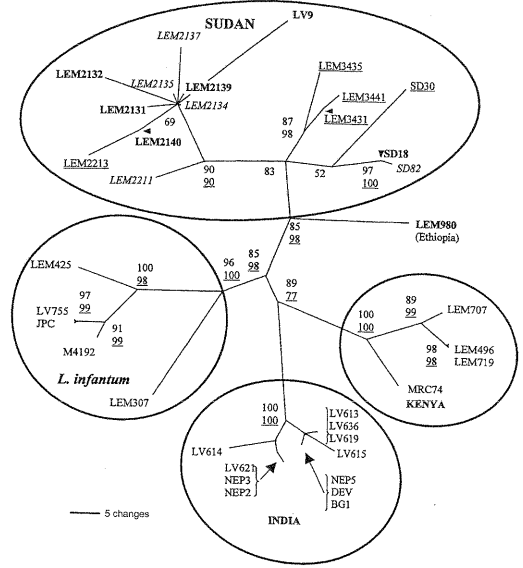


Fig. 2. Dendrogram of microsatellite data compiled using the parsimony option in PAUP. Bootstrap values were calculated using 100 bootstrap replicates. Bootstrap values from the distance tree are shown underlined underneath the bootstrap values for parsimony where the same clade was found in both trees. The MLEE-based species identification of Sudanese stocks is indicated by the typeface. Leishmania donovani in bold; Leishmania infantum underlined; Leishmania archibaldi in italics. These 3 putative taxa do not form distinct clades within the Sudanese clade nor does "Leishmania infantum" from Sudan cluster with Leishmania infantum from around the Mediterranean. Two distinct groups of parasites are found in Sudan but these groups do not have any correlation with MLEE identifications.

Inconsistencies with isoenzyme data

The data presented here are inconsistent with isoenzyme data in 2 respects. Firstly, we have presented molecular evidence that shows that "L. archibaldi" is a heterozygote for an amino acid that would cause a change in charge of the enzyme and would account for the difference in mobility of L. donovani and L. infantum. Heterozygotes at GOT have not been reported on MLEE gels by the Montpellier groups although Mauricio et al. (2001) refer to MON82 as a putative ASAT (=GOT) hybrid. The GOT enzyme is a homodimer and consequently the heterozygote would be expected to appear as a triplet with a strong middle band in MLEE gels. Given the small difference in relative mobility of the L. donovani GOT $_{113}$ enzyme compared with the L. infantum GOT $_{100}$ it is possible that a heterozygote with a strong intermediate band could be scored as a homozygote GOT $_{110}$. Secondly, 2 GOT loci are reported for L. donovani complex stocks in MLEE studies but our data only account for one of these loci. There is

Table 4. Microsatellite loci that may be in linkage disequilibrium

(Probability that 6 locus pairs are in linkage disequilibrium and hence presumed to be on the same chromosome. Note that LIST7-021, LIST7-027, LIST7-036 and possibly LIST7-022 appear to be on the same chromosome. * After applying a Bonferoni correction for the large number of tests that were made on this dataset only the LIST7-037 × LIST7-035 pair remains significant.)

Locus 1	Locus 2	Probability
LIST7-021	LIST7-027	0-975
LIST7-024	LIST7-029	0-970
LIST7-021	LIST7-036	0-964
LIST7-027	LIST7-036	0-990
LIST7-036	LIST7-022	0.980
LIST7-037	LIST7-035	1.00*

Table 5. Microsatellite loci with alleles that are associated with particular geographical regions

(The definition used to include Leishmania infantum stocks was parasites with a GOT¹⁰⁰ allele and that had been isolated outside the Leishmania donovani endemic areas. The Sudan group includes samples with a GOT¹⁰⁰ allele but since they were isolated within the Leishmania donovani endemic areas were excluded from the Leishmania infantum group. Further studies may show that some or all of these alleles are not restricted to particular geographical regions. These loci all appear to be independent since there was no evidence of linkage among this set of locus-pair combinations using Fisher's exact test as implemented by GENE-POP v.3.3 (Raymond & Rousset, 1995).)

	Microsatellite loci						
Region	LIST7-027	LIST7-036	LIST7-023				
Sudan	187; 189; 191; 193; 195	242; 244; 247; 249	171; 173; 175				
Kenya	183	238	153				
India	185; 184	241	151				
L. infantum	177; 179	232; 234	151				

conflicting evidence from MLEE and genome sequencing projects for the number of GOT loci that generate products visible on MLEE gels. Both GOT loci are scored as having enzymes with the same relative mobility in all zymodemes in the L. donovani complex and in 10 out of 11 L. major zymodemes (Maazoun et al. 1986; Pratlong et al. 2001; Rioux et al. 1990). Two aspartate aminotransferases (=GOT) are annotated in GeneDB for L. major Friedlin strain (LmjF24.0370 and LmjF35.0820), however, these 2 genes have no sequence homology when one is used in a BLAST search against the other, and they have different predicted isoelectric points (pH 7.0 and pH 7.9) and hence would be expected to have different mobilities by MLEE. Since these 2 known GOT loci are on different chromosomes it is unlikely

that their relative mobilities are so tightly linked that they change from GOT₁₀₀ in L. infantum to GOT₁₁₃ in L. donovani together. Consequently one of these genes is probably not being expressed at a level detectable by MLEE in promastigotes (the parasite stage usually studied by MLEE). In contrast in L. tropica, L. gerbili, L. aethiopica and Namibian stocks all zymodemes have 2 GOT enzymes visible on MLEE gels with substantial differences in relative mobility (15-87%) and which vary independently (Lanotte, Rioux & Serres, 1986). For these species 2 loci are clearly being detected on the MLEE gels and this is consistent with the L. major genome sequence data for 2 GOT loci. Gene expression studies might help to resolve these apparent anomalies. Although sequence data are presented here that would account for the MLEE data on L. infantum and L. donovani. further work will be required to confirm that the "L. archibaldi" heterozygotes that are predicted by the sequence data would account for the intermediate mobility of the "L. archibaldi" GOT isoenzyme allele. If the MLEE phenotype of "L. archibaldi" is shown to be a heterozygote then this could be interpreted as indicating that "L. archibaldi" is a hybrid between L. infantum and L. donovani. However it has been shown here that the "L. infantum" in Sudan is in fact L. donovani and probably the consequence of an independent and relatively recent mutation in the GOT gene in Sudan from local L. donovaní stocks, an example of convergence. The microsatellite data (www.genomics.liv.ac.uk/animal/ supplementary_data/jamjoom/ms_data.xls) was scanned for evidence that "L. archibaldi" stocks as a group may be heterozygotes of other Sudanese stocks but none was found in the 16 loci tested. Since the "L. archibaldi" stocks were scattered in 2 distinct clades in the Sudanese group with high bootstrap support it would appear that "L. archibaldi" is not a genetic entity at all and if individual stocks are hybrids then they are different hybrids that have arisen on more than one occasion. Further work will also be required to confirm that the gene sequenced here is expressed in promastigotes at a level that could be detected by isoenzymes and to discover the expression pattern of the other aspartate aminotransferase.

Geographical grouping of clades

The microsatellite classification identified strongly supported geographically based groups. However, additional stocks will have to be typed to confirm these observations since a previous study on this scale also identified regional groups but also found that some strains from Portugal, Lebanon, Iran and Italy clustered with a clade that was otherwise similar to the Sudanese clade identified here (Mauricio et al. 2001). Although there were clear regional groups, the relationships between the groups could not all be

resolved with confidence. This is partly because the tree was unrooted. Only 2 of the microsatellite primers isolated from L. donovani are informative within the potential outgroup L. major, and conversely only 2 microsatellites isolated from L. major are polymorphic in L. donovani (Jamjoom et al. 2002a; Jamjoom et al. 2002b). It is therefore not possible to root the microsatellite tree using the currently available loci. However, the Kenyan and Indian clades appeared to be most closely related to each other, consistent with an African rather than Asian origin for the Indian group as previously suggested by Ashford (1986).

Identification of L. infantum

New methods are now required to identify L. infantum. A total of 5 alleles at 3 loci were found exclusively in L. infantum but studies that include larger numbers of L. infantum stocks will be required to confirm that these alleles are diagnostic for this species. Unfortunately this means that currently there is no single marker for identifying L. infantum, although it is possible to classify stocks as L. infantum by using the microsatellites described here or Gp63 intergenic region PCR-RFLP (Mauricio et al. 2001).

A new diagnostic test is not required in Sudan

The confirmation that visceral leishmaniasis in Sudan is only caused by a single clade of *L. donovani* makes it possible to apply a single serological test to patients from Sudan with greater confidence. The possibility that the most widely used existing test, which is based on an *L. donovani* antigen, might be missing infections caused by other species can now be disregarded.

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