The Genome Sequence of *Leishmania* (*Leishmania*) *amazonensis*: Functional Annotation and Extended Analysis of Gene Models

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Abstract

We present the sequencing and annotation of the *Leishmania* (*Leishmania*) *amazonensis* genome, an etiological agent of human cutaneous leishmaniasis in the Amazon region of Brazil. *L. (L.) amazonensis* shares features with *Leishmania* (*L.) *mexicana* but also exhibits unique characteristics regarding geographical distribution and clinical manifestations of cutaneous lesions (e.g. borderline disseminated cutaneous leishmaniasis). Predicted genes were scored for orthologous gene families and conserved domains in comparison with other human pathogenic *Leishmania* spp. Carboxypeptidase, aminotransferase, and 3'-nucleotidase genes and ATPase, thioredoxin, and chaperone-related domains were represented more abundantly in *L. (L.) amazonensis* and *L. (L.) mexicana* species. Phylogenetic analysis revealed that these two species share groups of amastin surface proteins unique to the genus that could be related to specific features of disease outcomes and host cell interactions. Additionally, we describe a hypothetical hybrid interactome of potentially secreted *L. (L.) amazonensis* proteins and host proteins under the assumption that parasite factors mimic their mammalian counterparts. The model predicts an interaction between an *L. (L.) amazonensis* heat-shock protein and mammalian Toll-like receptor 9, which is implicated in important immune responses such as cytokine and nitric oxide production. The analysis presented here represents valuable information for future studies of leishmaniasis pathogenicity and treatment.

Key words: genome; *Leishmania* *amazonensis*; interactome; amastin; heat-shock protein

1. Introduction

Leishmaniases are neglected infectious diseases caused by parasites belonging to the Trypanosomatidae family and the *Leishmania* genus. Leishmaniases are prevalent in tropical countries; ~12 million people are affected by these diseases worldwide with 350 million people at risk of infection and an estimated yearly incidence of 2 million cases.¹ *Leishmania* spp. are digenetic parasites that develop as promastigotes in the gut of phlebotomine sandflies and as intracellular amastigotes in...
the macrophages of vertebrate hosts. The Lainson and Shaw classification\(^2\) subdivides the *Leishmania* genus into two subgenera based on the localization of promastigotes in the insect alimentary tract. The subgenus *Leishmania* comprises species limited to the midgut and foregut of the sand fly, whereas the subgenus *Viannia* includes species that develop a prolonged phase in the hindgut with later migration of flagellates to the midgut and foregut of the vector's alimentary tract. More recently, a third subgenus has been included in *Leishmania* classification, the subgenus *Sauroleishmania*, which comprises species that exclusively parasitize lizards.\(^3\) A brief classification of *Leishmania* subgenera and species associated with the diverse array of leishmaniasis clinical manifestations (cutaneous, mucocutaneous, and visceral forms) is provided in Fig. 1A.

Leishmaniasis are primarily zoonotic diseases, and a variety of mammals acts as reservoirs of *Leishmania* species. Specifically, rodents, edentates, and marsupials typically harbor cutaneous leishmaniasis, whereas wild canines and domestic dogs are the main reservoirs of zoonotic visceral leishmaniasis. In human hosts, disease outcomes are determined by a combination of parasitic properties (dermotropic versus viscerotropic species) and host factors, such as genetic variability and immune responses to infection.\(^4\) Among the causative species of cutaneous leishmaniasis in Brazil, recent data indicate that 8% are attributed to *Leishmania (L.) amazonensis*.\(^6\) This species can cause simple and diffuse forms of cutaneous leishmaniasis (DCL) and was implicated recently in borderline disseminated cutaneous leishmaniasis, an intermediate form of disease.\(^7\)

Infections with *Leishmania* species belonging to the *Leishmania (L.) mexicana* complex involve the dermal infiltration of macrophages that harbor parasites in large parasitophorous vacuoles (PVs). Most *Leishmania* species including *Leishmania (L.) major*, *Leishmania (L.) donovani*, and *Leishmania (V.) braziliensis* lodge intracellularly within small membrane-bound PVs that typically contain a single parasite and undergo fusion as the amastigotes divide.\(^9\) In contrast, amastigotes of *L. (L.) amazonensis* and *L. (L.) mexicana* are housed in large numbers within spacious PVs (Fig. 1B) that fuse together.\(^9\) These enlarged PVs may subvert host cell defenses by facilitating conditions of relatively diluted hydrolytic enzymes.\(^10,11\) The mechanistic basis of spacious PV development remains unknown and likely is triggered by unidentified parasitic factors produced by *L. (L.) amazonensis* and other species from the *L. (L.) mexicana* complex, such as *L. (L.) mexicana* and *Leishmania (L.) pifanoi*.\(^12\)

During the past decade, several reports have attempted to elucidate the factors used by *Leishmania* to interact with its vertebrate host and establish an infection. Like other kinetoplastids, gene expression in *Leishmania* is regulated mainly at the post-transcriptional level by RNA stability, rather than by promoters.\(^13\) Genes are organized into polycistronic transcriptional units, and protein-encoding genes are co-transcribed by RNA polymerase II. Precursor mRNAs subsequently are trans-spliced and polyadenylated.\(^14\)–\(^16\)

The number of chromosomes has been established for several *Leishmania* species.\(^17\)–\(^21\) The molecular karyotypes of Old World *Leishmania* species [ *L. (L.) infantum*, *L. (L.) donovani*, *L. (L.) major*, and *L. (S.) tarentolae*] each comprise 36 chromosomes,\(^17\) whereas the New World species, *L. (V.) braziliensis*, and *L. (L.) mexicana* have 35 and 34 chromosomes, respectively, due to fusion events involving 2–4 chromosomes.\(^18,19\)

The genomes of two Old World *Leishmania* species, *L. (L.) major* Friedlin, and *L. (L.) infantum* JPCM5, and one New World species, *L. (V.) braziliensis* M2904, have been sequenced and annotated.\(^19,22\) Recently, the genomes of *L. (L.) mexicana*, 16 clinical isolates of *L. (L.) donovani*, and the lizard parasite, *L. (S.) tarentolae*, were sequenced and assembled using high-throughput DNA sequencing technologies.\(^20,23,24\) Despite evolutionary divergence within the *Leishmania* genus, *Leishmania* comparative genomics suggests a high degree of synteny.\(^19,20,23,24\) *Leishmania* spp. from the *Leishmania* and *Viannia* subgenera exhibit highly conserved gene sequences with remarkably few genes or paralog groups that are unique to any given species. However, *L. (S.) tarentolae* lacks genes associated with the intracellular life stages of human pathogenic *Leishmania* spp.\(^20\) On the other hand, the *L. (V.) braziliensis* genome includes features that are lacking in the genomes of Old World *Leishmania* spp., such as transposable elements and RNA interference (RNAi) machinery.\(^19,25,26\)

To obtain a broader understanding of the pathogenesis of leishmaniasis, we sequenced the genome of the New World species, *L. (L.) amazonensis*. Using a comparative bioinformatics approach with other available *Leishmania* genomes, we searched for conserved domains and orthologous gene families among predicted proteins of *L. (L.) amazonensis*. In addition, we inferred the phylogeny of the surface glycoprotein, amastin, and generated a hybrid protein interactome to identify potential interactions between *L. (L.) amazonensis* secreted proteins and mammalian host factors.

## 2. Materials and methods

All the procedures employed in this study, except for phylogenetic analyses, are summarized in the workflow presented in Fig. 2. The workflow was divided into genome assembly and annotation steps, and functional and extended analyses of gene models.
2.1. Genomic DNA extraction

*Leishmania* (L.) *amazonensis* (MHOM/BR/71973/M2269 strain) was extracted from the cutaneous lesions of a patient from Cafezal city, in the state of Pará, Brazil, in 1973. Since then, this strain has been maintained in the laboratory by inoculating hamsters and mice and by axenic culture. Parasites were cultivated in M199 culture medium supplemented with 10% fetal bovine serum. Genomic DNA was extracted from \(10^9\) promastigotes after the parasites were incubated in lysis buffer [50 mM Tris–HCl (pH 8.0), 62.5 mM EDTA (pH 9.0), 2.5 M LiCl, 4% Triton X-100] at 37°C for 5 min. DNA was purified using phenol–chloroform extraction (1:1 v/v) and ethanol precipitation. The resulting pellets were resuspended in 50 \(\mu\)l of 10 mM TE [Tris–HCl (pH 8.0), 1 mM EDTA] containing 0.6 \(\mu\)g/\(\mu\)l of RNase A (Life Technologies Corporation, USA), and were incubated at 37°C for 30 min. Genomic DNA was precipitated with 2.5 \(\mu\)l of 100% ethanol and 0.3 M sodium acetate, centrifuged at 15 700g at 4°C for 15 min, and resuspended in DNase-free water.

2.2. Genome sequencing and assembly

*Leishmania* (L.) *amazonensis* DNA sequences were obtained using the whole-genome shotgun strategy on a combination of 454 GS-FLX Titanium (Roche) and Solexa (Illumina, Inc.) instruments from the University of North Carolina (UNC, USA) sequencing facility. The GS-FLX sequencer generated single-end fragment reads (454 reads) with a mean length of 315 bp. The Illumina Genome Analyzer generated 76-bp paired-end fragment reads (Solexa reads) with an average insert size of 400 bp. Using an Perl script developed in-house, Illumina reads were filtered out if the average Phred quality score was lower than 20. For 454 sequences, reads comprising fewer than 100 bp or more than 500 bp (5% of each side of the normal distribution of read sizes) and reads with more than 1 unknown nucleotide (N) also were filtered out. The Solexa reads were assembled into longer scaffolds using the Velvet 0.7.56 *de novo* assembler with a k-mer parameter of 43. This value was calculated using the Velvet Optimizer script (Victorian Bioinformatics Consortium, Monash University, Australia),...
which tested a range of k-mers from 31 to 69. Newbler software\textsuperscript{29} was then used to assemble the 454 reads. The N\textsubscript{50} scaffold and contig lengths of the *L. (L.) amazonensis* assembly were 22,275 and 17,272 bp, respectively. Solexa scaffolds and 454 contigs were combined by the Zorro assembler (Laboratório de Genômica e Expressão, UNICAMP, Brasil, http://lge.ibi.unicamp.br/zorro/downloads/Zorro2.2/DOC) to create the final hybrid genome assembly. The hybrid assembly was performed by combining the 4441 Illumina scaffolds and the 27,856 contigs from the 454 reads. Briefly, the Zorro pipeline consisted of (i) masking repeat regions in the contigs, (ii) detecting overlaps, (iii) unmasking repeat regions, and (iv) assembling hybrid contigs. In Phase 1, the repeat regions were determined by counting the occurrences of k-mers in the 454 reads. The assembler then masked k-mers in the contigs that occurred at high frequency. The absence of repeats enabled accurate overlap detection in Phase 2. The contigs were then unmasked in Phase 3, and the correct hybrid assembly was obtained in Phase 4 by merging all overlapping contigs into hybrid contigs. Overlap detection and consensus generation were performed using the Minimus package.\textsuperscript{30} The hybrid contigs were ordered and oriented with the Bambus program\textsuperscript{31} using paired-end information and manual verification, which yielded 2627 scaffolds.

2.3. Gene identification and annotation

2.3.1. Alignment of *Leishmania* spp. proteins with the *L. (L.) amazonensis* genome Predicted proteins for *L. (V.) braziliensis* (8153 proteins), *L. (L.) infantum* (8154 proteins), *L. (L.) major* (8298 proteins), and *L. (L.) mexicana* (8007 proteins) were mapped onto the *L. (L.) amazonensis* genome sequence using the Exonerate program (v. 2.2.0),\textsuperscript{32} which performs genomic searches and spliced alignments in a single run.

2.3.2. Ab initio prediction of gene models Glimmer, v. 3.02\textsuperscript{33} and Genemark.hmm, v. 3.3\textsuperscript{34} programs were used to perform *ab initio* gene predictions. The Glimmer long-orfs program was trained on non-overlapping open reading frames (ORFs) exceeding 200 bp in length. Genemark.hmm was executed in self-training mode by considering ORFs of at least 200 bp in length.

2.3.3. Combined gene models The above resources were used to automatically create *L. (L.) amazonensis* gene models using EVidenceModeler (EVM, v. r03062010) software.\textsuperscript{35} For individual outputs generated by Exonerate, Glimmer and Genemark.hmm were given values reflecting our data confidence to define gene structure. We considered the following confidence values: Glimmer = 3, Genemark = 3,
nucleotide-to-protein match = 6. The gene model was considered reliable only if it was predicted by all ab initio software or if it had similarity with one of the compared species. A final set of predicted gene models was then selected and manually inspected. Incomplete genes or genes overlapping gap regions were inspected with input from blastx alignments against protein databases and scaffold edges.

2.3.4. **Automatic annotation and analysis of protein domains** The automatic annotation software, AutoFACT (v. 3.4), was used for functional annotation of gene models. AutoFACT enables transitive annotation based on sequence similarity searches in several databases. We used the blastp algorithm (e-value 1 x 10^-7) to model gene alignments against the following protein databases: non-redundant proteins (NCBI RefSeq database, downloaded 8/6/2010), Swiss-Prot (only manually curated proteins, downloaded 4/15/2010), UniRef90, and UniRef100 (UniProt databases of clustered protein families, downloaded 4/15/2010), CDD (conserved protein domains, downloaded 4/15/2010), PFAM (protein domains, downloaded 4/15/2010), KEGG (metabolic pathways, downloaded 4/15/2010), and KEGG domains, downloaded 4/15/2010). We set AutoFACT to consider the following order of importance for annotation: UniRef100, UniRef90, KEGG, non-redundant proteins, and CDD. Data from CDD–PFAM analyses were extracted from AutoFACT and were used for comparative analyses of CDD–PFAM protein domains among L. (L.) amazonensis, L. (V.) braziliensis, L. (L.) infantum, L. (L.) major, and L. (L.) mexicana. These data also were evaluated using reverse PSI-BLAST (RPS-BLAST, e-value cutoff of 1 x 10^-5).

2.3.5. **Orthologous gene analysis** A data set composed of all the *Leishmania* spp. gene models was created and compared all-against-all using blastp (e-value cutoff of 1 x 10^-5). The results were submitted to OrthoMCL (v. 1.4) software, which clustered the proteins into orthologous and paralogous families. We applied the default software parameters, including an inflation index of 1.5. The inflation index regulates the cluster tightness (granularity) associated with sensitivity and selectivity. Clusters of proteins that presented bidirectional similarities between at least two *Leishmania* species were considered orthologs.

2.3.6. **Pseudogene identification** *Leishmania* and *Trypanosoma* spp. proteins were aligned against the *L. (L.) amazonensis* genome using blastx with an e-value threshold of 1 x 10^-5. The coordinates of the first hit alignment for all proteins were converted to GFF file format using an in-house Perl script. The BEDTools package was used to identify *L. (L.) amazonensis* regions exceeding 200 bp that showed similarity with *Leishmania* or *Trypanosoma* spp. proteins without overlapping with the gene model predictions. These regions were compared against the NCBI non-redundant database using the blastx program and manual annotation.

2.3.7. **Calculation of the codon adaptation index** The CodonW v. 1.4.4 software (http://bioweb.pasteur.fr/seqanal/interfaces/codonw.html) was employed to calculate the codon usage indices of each *L. (L.) amazonensis* predicted gene. The codon adaptation index (CAI) estimates the extent of bias toward codons known to be preferred in highly expressed genes. This index ranges from 0 to 1.0 with higher values indicating stronger codon usage bias and a higher expression level. The frequency of codon usage in highly expressed genes defines the relative fitness values for each synonymous codon. These values were calculated from the relative synonymous codon usage rather than from the raw codon usage and therefore were essentially independent of amino acid composition. Because fitness values are highly species specific, we first identified a set of highly expressed genes in *L. (L.) amazonensis*. This set was input into the calculation of the CAI. The effective number of codons (N_c) also was used to quantify the codon usage bias of each gene. The N_c ranges from 20 for a gene with extreme bias (using only 1 codon per amino acid) to 61 for a gene with no bias (using synonymous codons equally). Sequences for which N_c values are less than 30 were considered highly expressed, whereas sequences with N_c values exceeding 55 were considered poorly expressed genes.

2.4. **Phylogeny of amastin surface proteins** A phylogeny of *Leishmania* amastins was built from a set of *L. (L.) amazonensis* amastins predicted by our assembled genome and from all predicted amastin proteins in the *T. cruzi*, *T. brucei*, *L. (L.) major*, *L. (L.) infantum*, *L. (V.) braziliensis*, *L. (S.) tarentolae*, and *L. (S.) mexicana* genomes. These amastin sequences were extracted from TriTrypDB (accessed 6/28/2012) by searching for ‘amastin’ or ‘amastin-like’ entries. The 181 amastin/amastin-like protein sequences encoded by *L. (L.) braziliensis*, *L. (L.) infantum*, *L. (L.) major*, *L. (L.) mexicana*, and *L. (S.) tarentolae* and 24 amastin/amastin-like protein sequences from *L. (L.) amazonensis* were aligned using Geneious software (v. 5.6.3) in which an embedded MUSCLE software was applied. The alignment was performed using default parameters. The phylogenetic tree was inferred by Bayesian methods using MrBayes v. 3.1.2 with tree parameter optimization during the generations. A Bayesian tree was inferred based on 1 x 10^7 generations with a burn-in value of 75 000. Data were saved every 100 generations and were run in 4 chains.
2.5. Interactome of mammalian proteins and Leishmania secreted proteins

From the CDD and OrthoMCL analyses, protein families that were found to be expanded in L. (L.) amazonensis [i.e. more gene/domain copies when compared with L. (V.) braziliensis, L. (L.) infantum, and L. (L.) major] were also screened for orthologous in human protein databases. To identify human proteins that are orthologous to L. (L.) amazonensis proteins, we performed a blastp search (e-value cutoff of $1 \times 10^{-10}$) against the human protein databases using the parasite’s expanded proteins as query. These selected gene models were then analyzed using TargetP 1.1 and SecretomeP 2.0 prediction tools. Our objective was to classify proteins that could be secreted to the extracellular compartment or exposed for interaction with host cell components. We considered as putative secreted proteins those gene products for which TargetP returned a signal peptide value exceeding 0.8 and for which other values, such as the mitochondrial targeting peptide and the chloroplast transit peptide, were below 0.2. L. (L.) amazonensis proteins implicated in non-classical secretion or ectodomain shedding were predicted using SecretomeP v. 2.0 with the recommended threshold of 0.5.

Considering that amastigote is the developmental form which maintains a durable relationship with the mammalian host cell, an additional step was included in the screening. Using the screened gene models above mentioned, we performed a blastp analysis (e-value cutoff of $1 \times 10^{-5}$) against L. (L.) mexicana proteins expressed by amastigotes and identified in proteomic databases. L. (L.) amazonensis gene models that attend to these four criteria were considered for hybrid interactomes: (i) expanded or exclusive in L. (L.) amazonensis; (ii) orthologous to human proteins; (iii) possibly secreted (start codon was manually verified); and (iv) possibly expressed by amastigotes.

Some of the screened gene products could allow for interactions and interferences with native mammalian interactomes. Interactome networks for parasite-secreted proteins and for human and mouse proteins were created using IPA software (Ingenuity Systems, Redwood City, CA, USA) which was configured to build interactomes considering only those proteins expressed by cells of the immune system.

3. Results and discussion

3.1. Annotation of Leishmania (L) amazonensis gene models

Approximately 37 million 76-bp paired-end reads (average insert size, 400 bp) were assembled into 4411 scaffolds (coverage, ~94×) using Velvet de novo assembler software. The 454 reads (179 112 reads; average read length, 312 bp) were assembled using Newbler software into 27 856 contigs (coverage, ~2×). The final assembly was performed using Zorro, which combined scaffolds from Solexa and contigs from 454 sequencing to generate 2627 scaffolds (≥1000 bp in length) that specified an (L.) amazonensis genome size of 29.6 Mb.

Ab initio gene prediction was conducted using Glimmer and Genemark.hmm programs, which gave 8032 and 11 641 gene models (gene lengths ≥150 nt), respectively. Comparative gene predictions consisted of proteins from L. (V.) braziliensis, L. (L.) infantum, L. (L.) mexicana, and L. (L.) major aligned against the final L. (L.) amazonensis genome assembly. The final set of 8168 gene models was created using a combination of ab initio and comparative gene model analyses, EVM software to identify consensus gene structures supported by these two approaches, and manual annotation. Incomplete genes and genes overlapping gap regions were discarded from further analyses. Following manual annotation, 8100 gene models remained with an average length of 1793 bp; this is consistent with other sequenced Leishmania species. These data are summarized and compared with other Leishmania genomes in Table 1.

The final set of gene models were annotated automatically by comparing them against protein databases (blastp) and summarizing with AutoFACT software. The results indicated that 55% of the gene models showed high sequence similarity (e-value $1 \times 10^{-10}$) to functionally annotated proteins, whereas 42% of the models were similar to unassigned proteins (i.e. proteins with no functional annotation). The remaining 3% yielded no hits with any databases and were considered unclassified (Fig. 2).

In agreement with previous reports, our genome sequence analysis indicated that more than 90% of the 8100 L. (L.) amazonensis genes are shared with other human pathogenic Leishmania spp. with little variation in orthologous gene content. Despite varying clinical manifestations and features of lesions, Leishmania spp. harbor a conserved genomic core encoding functions ranging from fundamental biological processes to complex host–parasite interaction networks.

We performed a comparative annotation of tRNAs from the L. (L.) amazonensis, L. (L.) infantum, L. (V.) braziliensis, and L. (L.) major genomes. Using tRNAscan-SE software, the annotation yielded very similar numbers among the studied Leishmania species (Supplementary data, Table S1). Given the number of tRNAs for each codon in L. (L.) amazonensis, we calculated the CAI for all annotated gene models (Supplementary data, Table S2). The index measures...
the codon usage bias presented by highly expressed genes and can be comparatively employed to score native genes (higher CAI, more adapted to the pattern of codon usage) and possibly foreign or transferred genes (lower CAI, less adapted to the pattern of codon usage).56 Ranging from 0 (non-expressing gene, likely pseudogene) to 1 (highly expressed gene), we found that L. (L.) amazonensis predicted genes have a mean CAI value of 0.49 and median of 0.48 (Supplementary data, Fig. S1A). These data will be useful for future analyses of recent events of horizontal gene transfer in Leishmania.

Additionally, we scored regions in the L. (L.) amazonensis genome that showed similarities with predicted genes in other Leishmania and trypanosomatid species but were not identified as ORFs due to premature stop codons or frame shifts. This approach yielded 36 genomic regions corresponding to potential pseudogenes (Supplementary data, Table S3). One L. (L.) amazonensis pseudogene identified in our analysis was a fragment of argonaute 1 (AGO1), which is involved in the RNAi machinery. In the genome of L. (L.) amazonensis, we did not detect the known trypanosomatid argonaute and dicer variants, AGO1, DCL1 or DCL2. In addition, proteins containing two RNAse III domains (characteristic of dicer) or PAZ and Piwi domains (characteristic of argonaute) were not detected in this analysis. Our results suggest that RNAi pathways are absent in L. (L.) amazonensis, corroborating the hypothesis that RNAi via dicer and argonaute has been lost from the Leishmania subgenus following its divergence from the Viannia subgenus.26

Leishmania generally is considered to be a diploid organism because it carries two copies of most of its homologous chromosomes.19,22,24,57,58 However, there is increasing evidence suggesting that aneuploidy can occur in Leishmania species.23,24,58,59 Chromosome copy numbers can vary considerably among strains and species from different geographic regions, even among recent isolates.23,24 We mapped the L. (L.) amazonensis reads generated by Illumina against the L. (L.) mexicana chromosomes using the software SSAHA2.60 The median of coverage along each chromosome indicated a probable extra copy of chromosomes 7 and 26 and 3 extra copies of chromosome 30 in L. (L.) amazonensis (Supplementary data, Fig. S1B). In L. (L.) amazonensis, the exact number of chromosomes has not been defined. Preliminary studies using pulsed field gel electrophoresis have reported that the L. (L.) amazonensis karyotype consists of 25 chromosomal bands ranging in size from 0.2 to 2.2 Mb.61 Some bands exhibited variable ethidium bromide staining intensities possibly due to co-migration of chromosomes of similar sizes. Further studies will be needed to define the number of chromosomes and ploidy in L. (L.) amazonensis.

### 3.2. Functional analysis of gene models

In our functional analysis of gene models, we focused on the common factors, rather than the species-specific factors, predicted in L. (L.) amazonensis and L. (L.) mexicana genomes and not predicted in the genomes of the other species. We chose one genome data set for each disease outcome to compare with L. (L.) amazonensis and L. (L.) mexicana. Specifically, we chose leishmaniasis causative agents representative of cutaneous [L. (L.) major], mucocutaneous [L. (V.) braziliensis], and visceral [L. (L.) infantum] infections.

We searched for L. (L.) amazonensis genes that could be expanded or contracted in terms of gene or domain copies compared with the other species, particularly L. (L.) mexicana. We present a discussion of
some of these expanded genes/domains that could participate in: (i) parasite tropism in host organisms via adhesion molecules or amastin surface proteins; (ii) the development of large PVs by lipid synthesis; and (iii) intracellular establishment by enzymes related to nutritional acquisition and resistance to host intracellular defenses, such as oxidative burst.

3.2.1. Expanded and contracted orthologous gene families

The OrthoMCL software was applied to genome data sets from L. (L.) amazonensis, L. (L.) mexicana, L. (L.) infantum, L. (V.) braziliensis, and L. (L.) major to identify Leishmania orthologous gene families. We identified 7826 orthologous gene families [7488 orthologous gene families containing \( \geq 1 \) L. (L.) amazonensis protein] with 468 families comprising at least 7 members. Most (6784) of the orthologous gene families were shared by other Leishmania spp. Eight families were found only in L. (L.) amazonensis and L. (L.) mexicana and 23 families were unique to L. (L.) amazonensis (Fig. 3A, Supplementary data, Table S4). The five largest families identified across human pathogenic Leishmania spp. using OrthoMCL also were identified in L. (L.) amazonensis: dyneins (Family 1), glycopolypeptide GP63 (Family 3, leishmanolysins), histone H4 (Family 4), ABC transporters (Family 8), and amastins (proteins (Family 14; Supplementary data, Table S4). We identified several families associated with 2-fold or more gene copies in L. (L.) amazonensis than in any of the other human pathogenic species. These families include aminotransferases (family 256), 60S ribosomal protein L37 (family 216), and hypothetical proteins (families 323, 5508, and 7732). Among the families with at least 2-fold fewer gene copies in L. (L.) amazonensis we highlighted the GP63 gene family. The list of contracted gene families requires further investigation because some of them could be present in unassembled regions.

Comparative genome analyses of L. (L.) amazonensis and L. (L.) mexicana indicated an expansion in the gene family encoding a class-IV branched-chain amino acid aminotransferase (OrthoMCL family 256) that consisted of 270–415 amino acid residues and shared few regions of sequence similarity. Branched-chain aminotransferases catalyze the synthesis of leucine, isoleucine, and valine, and may be used to fulfill the parasite's nutritional requirements. They also may be involved in parasite sterol and prenol lipid synthesis because leucine is the main isoprenoid precursor for L. (L.) mexicana promastigotes and amastigotes.

A 3'-nucleotidase/nuclease (OrthoMCL family 7761) gene was predicted as being exclusive to the L. (L.) amazonensis and L. (L.) mexicana genomes. This gene encodes an enzyme responsible for nucleic acid hydrolysis that was found to be dramatically up-regulated on the cell surface of the trypanosomatid, *Chritidia luciliae*, under purine starvation conditions. An ecto-3'-nucleotidase/nuclease was detected experimentally in L. (L.) amazonensis; this component has important implications for parasite nutrition, adhesion to host cells, and infectivity. Substantial differences in the gene copy numbers between certain *Leishmania* spp. may account for the observed phenotypic variability in terms of pathogenesis and virulence. Several genome features could contribute to quantitative variation in gene copies among *Leishmania* spp. The expansion and contraction of genes in tandem arrays could result in up- or down-regulation of gene expression associated with copy-number variation. In addition, extensive variation in aneuploidy frequencies within parasite populations has been reported for several *Leishmania* spp. and for different *Leishmania* isolates within the same species.

3.2.2. Expanded and contracted conserved domains

Another comparison between *Leishmania* genomes was based on the identification of predicted protein domains in CDD–PFAM databanks. The result of CDD–PFAM analysis, included in the AutoFACT annotation, was retrieved and manually evaluated. In total, 2509 protein domains were identified; 2186 of these domains were detected in at least one L. (L.) amazonensis protein (Supplementary data, Table S5). Most (1881) of the identified domains were shared by other *Leishmania* spp; 20 domains were found exclusive to L. (L.) mexicana complex [L. (L.) amazonensis, and L. (L.) mexicana], and 26 domains were unique to L. (L.) amazonensis (Fig. 3B, Supplementary data, Table S5). The expanded and contracted protein domains in L. (L.) amazonensis were evaluated by the same criteria applied in OrthoMCL (i.e. variations in gene copies when compared with other *Leishmania* genomes). Among the most prevalent domains were the heat-shock protein (HSP)70 chaperone (CDD: 143803) and the vacuolar protein sorting-associated protein MRS6 (CDD: 34648). The following protein domains were detected only in L. (L.) amazonensis and L. (L.) mexicana: thioredoxin domain (CDD: 32932), tat-binding protein 1 (CDD: 148614), sterile alpha motif (SAM) domain of bicaudal C homolog protein 1 (BCC1, CDD: 188919), hydroxylase (CDD: 188206), and ATPase (CDD: 190944).

Examples of contracted protein domains identified in L. (L.) amazonensis, we highlight: amastins (CDD: 140228), UDP-GlcNAc-dependent glycosyltransferase (CDD: 140237), leishmanolysin peptidase M8 (CDD: 189994), cathepsin (CDD: 185513), tryptaredoxin peroxidase (CDD: 140280), non-long terminal repeat reverse transcriptases (RTs-nLTR, CDD: 73156), rim ABC transporters (CDD: 185513), adenylyl/guanylyl cyclase (CDD: 128359), and paraglaxar nod rod protein (CDD: 140353).
The present study detected a thioredoxin domain unique to *L. (L.) amazonensis* and *L. (L.) mexicana* (NCBI accession COG3118). Thioredoxin functions as a hydrogen donor or disulfide reductase and is involved in the response to oxidative stress and in protein folding. Reactive oxygen species can be scavenged directly by thioredoxin or by thioredoxin-related dehydrogenases. Parasitic thiol and dithiol proteins may buffer the redox environment of PVs; this could account for the resistance of *L. (L.) amazonensis* to nitric oxide (NO) production in interferon (IFN)-γ-activated macrophages.

3.3. Extended gene model analysis

3.3.1. Amastin phylogeny suggests specialized amastins in the *Leishmania* (L.) mexicana complex

Previous *Leishmania* phylogenetic analyses, based on comparisons of isoenzymes, DNA sequences, and HSP profiles among species agreed with the adopted Linnean classification and with complexes proposed by Lainson and Shaw in 1987. The phylogenies from these studies indicated that *L. (L.) amazonensis* has an evolutionary proximity to *L. (L.) mexicana*, a finding that was interpreted as these species comprising a monophyletic clade. These studies also indicate that parasites responsible for cutaneous/mucocutaneous lesions could be as different and divergent from one another as they are from the parasites that cause visceral leishmaniasis. For instance, the phylogenetic distance between the cutaneous-associated species, *L. (L.) amazonensis* and *L. (L.) major*, is similar to the distance between *L. (L.) amazonensis*/*L. (L.) major* and *L. (L.) donovani*, which causes visceral leishmaniasis. Therefore, the same disease outcomes in *Leishmania* mammalian hosts can result from a variety of evasive strategies and factors distinctively featured by *Leishmania* spp.

Amastin belongs to a multi-gene family in *Leishmania* that encodes small surface proteins of ≏ 200 amino acids. Several members of the amastin gene family are dispersed throughout the genomes of all *Leishmania* species and exhibit various expression patterns. Phylogenetic analysis of trypanosomatid amastins defined four subfamilies of amastin (α, β, γ, and δ) with distinct genomic organizations and expression patterns during the cell cycles of *T. cruzi* and *Leishmania* spp. δ-amastins comprise the largest and most diverse amastin subfamily. In *T. cruzi*, δ-amastin expression was associated with parasite infectivity to host cells. In *Leishmania*, the amastin N-terminal signature peptides are among the most immunogenic of all leishmanial surface antigens in mice and generate strong immune responses in humans with visceral leishmaniasis. DNA microarray analyses have implicated amastin in the intracellular survival of the parasite. Amastin gene expression was detected predominantly in amastigotes of several *L. (L.) donovani* strains isolated from patients with visceral and post-kala-azar dermal leishmaniasis. The roles of amastin genes in parasite homeostasis and growth inside acidic PVs also have been addressed. As transmembrane proteins, amastins could contribute to proton or ion trafficking across the membrane to adjust cytoplasmic pH under the harsh conditions of

![Figure 3](http://data.oxfordjournals.org/)

**Figure 3.** Diagrammatic representation of (A) species-specific orthologous gene families (OrthoMCL analysis) and (B) conserved domains (CDD–PFAM analysis). A core of 6784 orthologous families and 1881 domains was conserved in all studied *Leishmania* species [*L. (L.) amazonensis*, *L. (L.) mexicana*, *L. (L.) major*, *L. (L.) infantum*, and *L. (V.) braziliensis*]. We detected 8 orthologous families and 20 conserved domains that were exclusive to *L. (L.) mexicana* complex. A complete list of orthologous families and conserved domains is presented in Supplementary data, Table S4 and S5, respectively.
the phagolysosome. As a surface epitope, amastin may be recognized by opsonizing host IgG antibodies and could promote parasite uptake by host macrophages (via Fc receptors) and subsequent release of interleukin (IL)-10.\(^8\)\(^1\) We speculate that amastins could be involved in certain peculiar characteristics of \(L.\) (\(L.\)) \textit{amazonensis}, such as its propensity to induce DCL, its development inside spacious PVs, and its resistance to the highly oxidative phagolysosomal environment in host cells.

The TriTrypDB provides a set of annotated amastin and amastin-like surface proteins in the \textit{Leishmania} and \textit{Trypanosoma} genomes. Our search of the TriTrypDB yielded 181 annotated genes encoding amastin or amastin-like proteins in \(L.\) (\(L.\)) \textit{brazilienis}, \(L.\) (\(L.\)) \textit{infantum}, \(L.\) (\(L.\)) \textit{major}, \(L.\) (\(L.\)) \textit{mexicana}, and \(L.\) (\(S.\)) \textit{tarentolae}. The \(L.\) (\(L.\)) \textit{amazonensis} genome presented in this study identified 12 orthologous groups annotated as amastin or amastin-like proteins (families 14, 19, 20, 3539, 3935, 5852, 6119, 6120, 6543, 7556, 7771, and 7778). All families corresponded to 1 representative gene model, except family 19 (3 gene models were associated with this orthologous group). Thus, our OrthoMCL analysis predicted 14 gene models were associated with this orthologous representative gene model, except family 19 (3756, 7771, and 7778). All families corresponded to 14, 19, 20, 3539, 3935, 5852, 6119, 6120, 6543, 7556, 7771, and 7778). All families corresponded to 1 representative gene model, except family 19 (3 gene models were associated with this orthologous group).

Thus, our OrthoMCL analysis predicted 14 gene models were associated with this orthologous representative gene model, except family 19 (3 gene models were associated with this orthologous group).

The CDD–PFAM analysis also identified gene models containing amastin domains that were not identified in the OrthoMCL analysis. We gathered all genes identified as amastin/amastin-like proteins (OrthoMCL) or containing amastin domains (CDD–PFAM) and built a list of 24 \(L.\) (\(L.\)) \textit{amazonensis} predicted amastin/amastin-like surface proteins to perform our phylogenetic analysis.

Alignment of the \textit{Leishmania}-annotated amastins with 24 amastin/amastin-like proteins identified in the \(L.\) (\(L.\)) \textit{amazonensis} genome (OrthoMCL and CDD–PFAM combined scoring) allowed us to build an amastin phylogenetic tree (Fig. 4). By placing the phylogenetic root halfway between the two most divergent/distant amastins (midpoint rooting) we identified clades composed of species-related amastins. Early branching clades could represent a class of amastin surface proteins conserved in \textit{Leishmania} prior to its radiation (Fig. 4, blue branches). These \textit{Leishmania} pre-speciation amastins are gathered in \(\alpha\), \(\beta\), and \(\gamma\) subfamily clades.\(^7\)\(^4\)\(^7\)\(^4\) The presence of species-specific clades of \(\delta\)-amastins at terminal tree branches (Fig. 4, red, green, yellow, and purple branches) suggests that several amastin surface proteins appeared because of environmental selective pressures or pathogen specification. At least in part, this could be associated with the diverse leishmaniasis outcomes of different \textit{Leishmania} species. It is possible to identify amastin subfamilies unique to \(L.\) (\(L.\)) \textit{major}, \(L.\) (\(V.\)) \textit{brazilienis}, and \(L.\) (\(L.\)) \textit{infantum}, and three groups in which \(L.\) (\(L.\)) \textit{amazonensis} and \(L.\) (\(L.\)) \textit{mexicana} amastins represent a distinct clade of amastins (Fig. 4, red branches). These amastins could play a role in the unusual housing of these parasites within spacious PVs of infected macrophages.

3.3.2. Secreted \textit{Leishmania} HSPs could interfere with native host interactomes One of the most striking features of the \(L.\) (\(L.\)) \textit{mexicana} complex is the development of giant PVs in infected macrophages that harbor amastigotes. We speculate that the formation of large PVs may be related to factors secreted by the parasite, the subversion of host native vesicular trafficking, and potentially the production/incorporation of parasitic components into PV membranes. \(L.\) (\(L.\)) \textit{amazonensis} amastigotes interact with the internal membranes of PVs via their posterior poles.\(^8\) The posterior pole behaves like an adhesion site between the parasite and PV membranes, although no adhesion factors or junction components have been identified to date. De Souza Leao et al.\(^8\)\(^2\) suggested that in \(L.\) (\(L.\)) \textit{amazonensis}-infected macrophages, the internalization and degradation of major histocompatibility complex (MHC) class II molecules by amastigotes occur through their posterior poles. This degradation could be performed by secreted components inserted into the PV membrane. The posterior pole also may be interpreted as part of a parasitic secretory pathway in which secreted proteins directly encounter the host cell cytosol, bypassing the acidic milieu of the PV. Once in the host cell cytosol, secreted factors may be transferred to the host cell nucleus and/or plasma membrane, affecting gene expression, cellular functions and metabolic processes. However, the classically described site for parasite exocytosis and endocytosis is the anterior pole where the flagellar pocket is located. The flagellar pocket faces the lumen of the PV, and most secreted \textit{Leishmania} proteins, regardless of their association with exosomes, are expected to reach the acidic (pH 4.5–5.0) PV milieu from there.\(^8\)\(^3\)

We hypothesized that \textit{Leishmania} secreted factors could mimic mammalian factors, thus perturbing native host protein interactions. To identify possible interactions between parasitic and mammalian host factors, we constructed hybrid protein interaction networks in which human and mouse databases were compared against the list of proteins that are potentially secreted by \(L.\) (\(L.\)) \textit{amazonensis} amastigotes. Our CDD–PFAM and OrthoMCL analyses identified nine conserved domains and three orthologous gene families that were exclusive to or expanded in \(L.\) (\(L.\)) \textit{amazonensis} and/or \(L.\) (\(L.\)) \textit{mexicana} and are also orthologous to human proteins (Supplementary data, Table S6). The CDD 143803, an HSP 70 domain, exists as six copies in \(L.\) (\(L.\)) \textit{amazonensis}, four copies in \(L.\) (\(L.\)) \textit{mexicana}, and three copies in the other species’ genomes. We considered this as an expanded number of HSP70 domains.
in the *L. (L.) mexicana* complex. The six gene models in which these domains were identified (A42670, A6630, A68920, A73510, A30200, and A45910) were submitted to the TargetP and SecretomeP servers and gene models A30200 and A45910 were predicted for secretion. Additionally, A30200 and A45910 were similar to two *L. (L.) mexicana* proteins identified in a proteomic data analysis of *L. (L.) mexicana* amastigotes\(^5\) (LmxM.28.2770 and LmxM.34.4710, respectively; Supplementary data, Table S6). Although these genes are not exclusive to the *L. (L.) mexicana* complex, their similarity to two products from amastigote proteomic data is suggestive that, at least in *L. (L.) amazonensis* and *L. (L.) mexicana*, these products are expressed by the intracellular form of the parasite and could be secreted within host cells.

**Figure 4.** Bayesian consensus phylogeny of amastin surface proteins. The phylogram is represented by a consensus of 214 amastin sequences. The root was inferred using midpoint rooting. WAG was used as the substitution matrix for the protein alignment. Posterior probabilities exceeding 0.5 are shown in the branches. The tree topology suggests early branching of similar amastins shared by different species (blue). These branches were classified as *Leishmania* pre-speciation amastins, composed by α, β, and γ subfamily clades. We highlighted the terminal taxa (late branching or apomorphic) of species-specific δ-amastin clades of *L. (L.) major* (yellow), *L. (V.) braziliensis* (green), and *L. (L.) infantum* (purple). Complex-specific clades of *L. (L.) amazonensis* and *L. (L.) mexicana* amastin surface proteins are in red. The scale of the generated tree (see 0.4 bar) represents the number of substitutions per sequence position. The classification of amastin clades in subfamilies α, β, γ, and δ was based on the amastin phylogeny performed by Jackson et al. (2010).
The A30200 and A45910 gene models are candidates for the construction of hybrid interactomes, given that they present a protein domain expanded in both *L. (L.) amazonensis* and *L. (L.) mexicana* (CDD 143803), are predicted to be secreted and are similar to proteins identified in the proteome of *L. (L.) mexicana* amastigotes. The A30200 and A45910 gene models present a considerable similarity with the mammalian hypoxia up-regulated protein 1 (HYOU1) and the HSP 70 kDa protein 5 (HSPA5), respectively. We created HYOU1 and HSPA5 interactome subsets to identify host components that could be affected by the secretion of both A30200 and A45910 gene products. This analysis led us to identify toll-like receptor (TLR) 9 and IL-6 as putative targets of the *L. (L.) amazonensis* A30200 and A45910 gene products (Fig. 5A and B). The HYOU1 and HSPA5 both directly interact with TLR9, a receptor implicated in the recognition of CpG DNA motifs and present in endolysosomal compartments where it is activated by proteolytic cleavage. Thus, TLR9 is preferentially expressed in the granulomas of human cutaneous leishmaniasis caused by *L. (V.) braziliensis*, and TLR9-deficient mice are more susceptible to *L. (L.) major* infection. Thus, TLR9 is implicated in the immune response against *Leishmania*. The interaction between host TLR9 and the putative secreted *L. (L.) amazonensis* HYOU1/HSPA5-mimic could block TLR9 function and favor intracellular establishment of the parasite. TLR9 also is implicated in the production of NO via NO synthase 2, tumor necrosis factor, IL-6, and IL-12B. The production of IL-6 is inhibited in dendritic cells differentiated from monocytes in the presence of *L. (L.) amazonensis* and is present at low levels in the sera of Chiclero’s ulcer patients infected for 3–8 months. Linares *et al.* reported that in vitro infection with *L. (L.) amazonensis* amastigotes decreases NO production by macrophages stimulated with IFN-γ plus lipopolysaccharide. Thus, although hypothetical and genome based, our proposed interactome can be used to identify components implicated in the establishment of *Leishmania* infection of mammalian host cells. Moreover, the interactome provides a model for studying *Leishmania*-secreted proteins and their influence on important effectors of the host cell immune response.

### 3.4. Conclusions

We present the genome of the protozoan *L. (L.) amazonensis* together with functional annotations and extended analyses focused on host–parasite interactions. We examined the genome sequences of *L. (L.) amazonensis* and *L. (L.) mexicana* for potentially expressed genes at expanded copy numbers. Confirming that a few *Leishmania* species-specific genes may exist despite striking conservation at the gene level, we report conserved domains, orthologous gene families, and amastin surface proteins unique to *L. (L.) amazonensis* and *L. (L.) mexicana*. Additionally, we propose an innovative approach to interactome analysis that emphasizes the role of parasite secreted proteins in host interaction networks.

### 4. Availability

The *Leishmania* (Leishmania) *amazonensis* Genome Database is available at the URL http://www.lge.ibi.unicamp.br/leishmania. This Whole Genome Shotgun...
The Genome Sequence of \textit{L. (L.) amazonensis}

project was deposited at DDBJ/EMBL/GenBank under the accession APNT00000000 (SUBID SUB120161, BioProject PRJNA173202). The version described in this paper is the first version, APNT01000000. While revising this manuscript, we realized that another Brazilian group from Instituto Oswaldo Cruz (IOC)—Fiocruz, Rio de Janeiro, is sequencing the genome of \textit{L. (L.) amazonensis}.

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