

# 1 LeishMANIAdb: a comparative 2 resource for *Leishmania* proteins

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14  
15 **Abstract** Leishmaniasis is a detrimental disease causing serious changes in quality of life and  
16 some forms lead to death. The disease is spread by the parasite *Leishmania* transmitted by  
17 sandfly vectors and their primary hosts are vertebrates including humans. The pathogen  
18 penetrates host cells and secretes proteins (the secretome) to repurpose cells for pathogen  
19 growth and to alter cell signaling via host-pathogen Protein-Protein Interactions (PPIs). Here we  
20 present LeishMANIAdb, a database specifically designed to investigate how *Leishmania* virulence  
21 factors may interfere with host proteins. Since the secretomes of different *Leishmania* species are  
22 only partially characterized, we collected various experimental evidence and used computational  
23 predictions to identify *Leishmania* secreted proteins to generate a user-friendly unified web  
24 resource allowing users to access all information available on experimental and predicted  
25 secretomes. In addition, we manually annotated host-pathogen interactions of 211 proteins, and  
26 the localization/function of 3764 transmembrane (TM) proteins of different *Leishmania* species.  
27 We also enriched all proteins with automatic structural and functional predictions that can  
28 provide new insights in the molecular mechanisms of infection. Our database, available at  
29 <https://leishmaniadb.ttk.hu> may provide novel insights into *Leishmania* host-pathogen interactions  
30 and help to identify new therapeutic targets for this neglected disease.

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## 32 Introduction

33 Leishmaniasis is a neglected tropical disease causing severe symptoms, affecting around 1 million  
34 new people yearly, with annual deaths estimated to be around 60,000 *Torres et al. (2017)*. Although  
35 over 90% of cases occur in poor regions south of the Equator, due to climatic changes it also ap-  
36 pears in new areas, and it has already shown up in Mediterranean European countries *Gianchec-*  
37 *chi and Montomoli (2020)* and Texas, USA *McIlwee et al. (2018)*. To this date no approved human  
38 vaccine is available and treatment is most effective at an early stage of the infection. *Leishmania*  
39 parasites are unicellular, flagellated trypanosomatids, belonging to the class Kinetoplastea. Upon  
40 infection, the amastigote stage pathogen (with reduced flagella) is engulfed by phagocytes, where it  
41 ends up in a stable parasitophorous vacuole that protects it *Arango Duque et al. (2019)*. *Leishmania*

42 cells then proliferate unhindered within host cells until egress and spreading to nearby phagocytes  
43 *Real et al. (2014)*. The parasite secretes proteins that enter various parts of the cell *Atayde et al.*  
44 *(2015)*. The secreted virulence factors can then interfere with cell signaling by interacting with the  
45 host proteins: they increase glycolytic metabolism *Ohms et al. (2021)*, perturb microbicidal path-  
46 ways *Matheoud et al. (2013)*, escape the innate immune response, and repurpose macrophages  
47 for parasite replication *Atayde et al. (2016)* by disturbing with cellular protein-protein interactions  
48 (PPIs). Interestingly, these mechanisms are somewhat unique to *Leishmania* among trypanosomes,  
49 which are usually extracellular pathogens and do not enter host cells. In contrast, *Leishmania* se-  
50 cretes proteins which are critical for host cell subjugation, but how they enter the cytoplasm of  
51 host cells is still poorly understood.

52 The host targeted interactions are often mediated via Short Linear Motifs (SLiMs) in many dis-  
53 tant, unrelated intracellular pathogens, ranging from viruses and bacteria to unicellular eukary-  
54 otes *Davey et al. (2011)*. SLiMs are flexible protein segments composed of a restricted number of  
55 residues (between 3-10), that usually bind to structured protein domains. Their short length and  
56 structural flexibility enable them to bind to a wide range of domains. Cellular SLiMs typically bind  
57 their targets with low micromolar affinity. These weak and transient interactions enable SLiMs to  
58 work in cooperative regulatory systems *Van Roey et al. (2014)*. Pathogens mimic host SLiMs to in-  
59 teract with host cell proteins *Davey et al. (2011)*. Pathogen SLiMs often bind with higher affinities  
60 than the cellular ones, outcompeting the native interactions, permanently re-wiring the host reg-  
61 ulation network. A few modulatory SLiMs have already been discovered in eukaryotic pathogens,  
62 such as the *Toxoplasma gondii* MapK docking motif *Pellegrini et al. (2017)* and the stage-specific  
63 (promastigote-amastigote) phosphorylation motifs from *Leishmania* *Tsigankov et al. (2013)*. In ad-  
64 dition, several putative SLiMs were recently detected in *Leishmania* such as heparin-binding se-  
65 quences or RGD integrin-binding motifs but their function has not been confirmed yet *Peysse-  
66 lon et al. (2013)*.

67 Numerous studies investigated *Leishmania* secretomes. Most of them expose promastigotes to  
68 a heat shock and pH change (attempting to emulate the conditions that promote promastigote-to-  
69 amastigote stage transition) and then analyze the *Leishmania* conditioned medium by proteomics  
70 to identify secreted proteins *Cuervo et al. (2009)*, and measure their protein abundance or by  
71 transcriptomics to detect mRNA levels *Lahav et al. (2011)*. While high-throughput experiments  
72 inherently suffer from a certain level of noise, experiments on individual proteins may be more  
73 reliable - in the case of *Leishmania* the vast majority focuses on leishmanolysin (GP63), a surface-  
74 anchored protease important for pathogenesis *Gregory et al. (2008)*; *Guay-Vincent et al. (2022)*.  
75 Furthermore, data were collected on different *Leishmania* species/strains identified via names and  
76 identifiers varying from one source to another, making a unified overview challenging. Another  
77 key step towards understanding the infection mechanism would be the identification of *Leishma-  
78 nia* surface proteins that can mediate the attachment of the pathogen to the host cell. Some sur-  
79 faceome experiments were carried out on *Leishmania*-related species, and human host proteins  
80 binding to the surface of 24 strains of intact *Leishmania* have been identified *Fatoux-Ardore et al.*  
81 *(2014)*. Beside the characterization of *Leishmania* secretomes, the identification of host-*Leishmania*  
82 PPIs is needed to narrow down virulence factors perturbing the host cell regulation to modules  
83 interfering with host proteins. SLiMs have low information content and simply scanning them in  
84 *Leishmania* secretomes may yield many false positives. Their structural and functional context,  
85 such as accessibility, conservation and localization, are all key elements to successfully identify  
86 those that may have a role in rewiring the host cell regulation. Notably, SLiMs also play a key role  
87 in maintaining housekeeping processes in *Leishmania*, therefore to find candidate SLiMs that may  
88 alter the host regulation, we need to discriminate SLiMs of proteins that reach the host cytoplasm  
89 or nucleus but limited information about these proteins are available. Currently the only publicly  
90 available database dealing with *Leishmania* proteins is TriTrypDB *Shanmugasundram et al. (2023)*,  
91 which is part of the VEuPathDB *Amos et al. (2022)*. TriTrypDB is a functional genomic resource for  
92 Trypanosomatidae, offering proteomic datasets, however it does not focus on protein structure,

93 protein motif search and interactions.

94 We developed LeishMANIAdb to expedite *Leishmania* research by unifying scattered informa-  
95 tion from the literature in a user-friendly way and to extend available resources by adding protein  
96 level information. We collected high-throughput experiments and interaction studies on individual  
97 proteins, and used various prediction methods to enrich proteins with structural information.

## 98 Results

### 99 Selection of *Leishmania* proteomes and homology mapping of various kinetoplastid 100 proteins

101 We selected 5 *Leishmania* species (reference proteomes: *L. braziliensis*, *L. donovani*, *L. infantum*, *L.*  
102 *major*, *L. mexicana*), 13 *Leishmania* strains (*Lbraziliensis MHOMBR75M2903*, *Lbraziliensis MHOMBR75M2904*,  
103 *Lbraziliensis MHOMBR75M29042019*, *Ldonovani BPK282A1*, *LdonovaniCL-SL*, *Ldonovani HU3*, *Ldono-*  
104 *vani LV*, *Linfantum JPCM5*, *LmajorFriedlin*, *Lmajor Friedlin2021*, *Lmajor LV39c5*, *Lmajor SD75.1*, *Lmexi-*  
105 *cana MHOMGT2001U1103*), and 6 related species (reference proteomes: *Bodo saltans*, *Leptomonas*  
106 *seymouri*, *Trypanosoma brucei*, *Trypanosoma cruzi*, *Trypanosoma rangeli*, *Trypanosoma theileri*) as an out-  
107 group. The 5 *Leishmania* proteomes and the 6 related kinetoplastid proteomes were selected  
108 based on their quality (i.e. number of fragments and missing proteins) and were downloaded  
109 from UniProtKB **UniProt Consortium (2023)**. *Leishmania* proteins were also cross-referenced to  
110 TriTrypDB **Shanmugasundram et al. (2023)**. Around 30% of the cross-referenced proteins have dif-  
111 ferent sequences deposited into these resources, and in most cases the difference is due to the  
112 position of the initiator methionine. For data compatibility we always use the UniProt sequence  
113 version but the conflicts are highlighted in LeishMANIAdb. We also performed a similarity search  
114 between these proteins and linked close homologs (see Methods) so annotations and predictions  
115 can be easily compared between them. All manual annotations and experimental data from dif-  
116 ferent sources were mapped to these proteins. The 13 *Leishmania* strain proteomes were down-  
117 loaded from TriTrypDB. Altogether LeishMANIAdb contains 40 537 searchable *Leishmania* proteins  
118 from reference proteomes, 108 766 proteins from different strains and 68 924 other kinetoplastid  
119 proteins to strengthen predictions.

### 120 Manual annotation of host-pathogen PPIs and TM protein localization

121 We manually curated hundreds of proteins, using two strategies. The first type of annotation was  
122 the collection of host-pathogen PPI experiments on individual proteins, with the majority of them  
123 involving leishmanolysin (GP63). We collected 29 papers reporting 82 *Leishmania* PPIs with dif-  
124 ferent hosts. Although experiments were mapped back to specific proteins, the results are also  
125 displayed on close homologs (with a note that the experimental data is derived from a different  
126 protein) resulting in 211 proteins that contain PPI data. Interactions were reported using the Mini-  
127 mum Information required for reporting a Molecular Interaction eXperiment MIMIx **Orchard et al.**  
128 **(2007)** community standard description. The second type of manually curated data was the local-  
129 ization and functional annotation of TM proteins. The aim was to find surface proteins that may  
130 facilitate the infection, but we annotated hundreds of other TM proteins with their localizations  
131 too. For this task, we used all close homologous proteins defined in the previous step. Altogether  
132 342 protein families were annotated and these annotations were shared between 3764 proteins  
133 (which is 45.11% percent of the predicted TM proteomes, and 9.28% of all proteins of the 5 species  
134 combined).

### 135 The definition of *Leishmania* secretome and protein localization is still incomplete

136 *Leishmania* not only exploits host-secretory pathways to distribute effectors but also utilizes an  
137 unusual mechanism to deliver proteins to the cytosol of infected cells by releasing exosomes into  
138 the parasitophorous vesicle, which might fuse with the vesicular membrane to release their pro-  
139 tein content **Silverman et al. (2010)**. Therefore computational methods based on signal peptides

140 and localization predictions are not sufficient to predict *Leishmania* secretomes. To overcome this  
141 limitation we also used high-throughput experiments *Silverman et al. (2008)*; *Cuervo et al. (2009)*;  
142 *Hassani et al. (2011)*; *Forrest et al. (2020)*; *Pissarra et al. (2022)* to increase the coverage of *Leishma-*  
143 *nia* secretomes. Strikingly, the number of proteins in these secretomes varies to a large extent, and  
144 some proteins cannot be identified by mass spectrometry. Other datasets include proteins found  
145 in glycosomes *Jamdhade et al. (2015)*, stage-dependent (promastigote/amastigote) phosphopro-  
146 teomics *Tsigankov et al. (2013)*, housekeeping gene localizations *Jardim et al. (2018)*, exosome con-  
147 tent *Silverman et al. (2010)*, protein and mRNA abundance data *Lahav et al. (2011)*; *Pescher et al.*  
148 *(2011)*. When we mapped back all secretome and abundance experiments to *Leishmania infantum*  
149 (from orthologous proteins of other *Leishmania* species), the number of identified proteins ranges  
150 from 10 to 2,000 (Figure 1/A), and even when experimental conditions were similar they yielded  
151 highly different amounts of proteins. For example, pioneer secretome studies only provided a few  
152 hundred hits, while the latest ones are more inclusive with thousands of hits. Gene duplication  
153 is often acting on protein families responsible for host-pathogen PPIs, therefore we also collected  
154 proteins that are highly expanded. Notably, as all kinetoplastids have a polycistronic transcrip-  
155 tion system, the main way to amplify expression of critical proteins is through gene duplication.  
156 Thereby highly expanded gene families can be directly mapped to functions critical for these para-  
157 sites *Jackson et al. (2016)*. In this case we could discriminate between proteins with already many  
158 paralogs within kinetoplastids and *Leishmania*-exclusive amplified proteins. When we searched  
159 for homologs of *Leishmania infantum* proteins, we found distinct amino acid transporter and co-  
160 factor families already expanded in all kinetoplastids including *Leishmania*. In contrast, amastins,  
161 leishmanolysin, 3'A2-related proteins, kinase-containing putative receptor proteins (and several  
162 uncharacterized proteins) seemed to be highly abundant in *Leishmania* proteomes compared to  
163 all kinetoplastids (Figure 1/B). Comparing complete proteomic datasets yielded only a small over-  
164 lap. We defined 1) *Leishmania*\_novelty proteins, which are proteins without close homologs in  
165 SwissProt, without characterized Pfam domains, and expanded in *Leishmania infantum* (compared  
166 to other kinetoplastids); 2) abundant proteins, which are proteins showing increased abundance  
167 upon infection; 3) secreted proteins experimentally identified in at least two secretome experi-  
168 ments. These definitions provided markedly different protein sets, with some overlap between  
169 secreted and abundant proteins (611 proteins) and with only 22 proteins contained in all datasets  
170 (Figure 1/C).

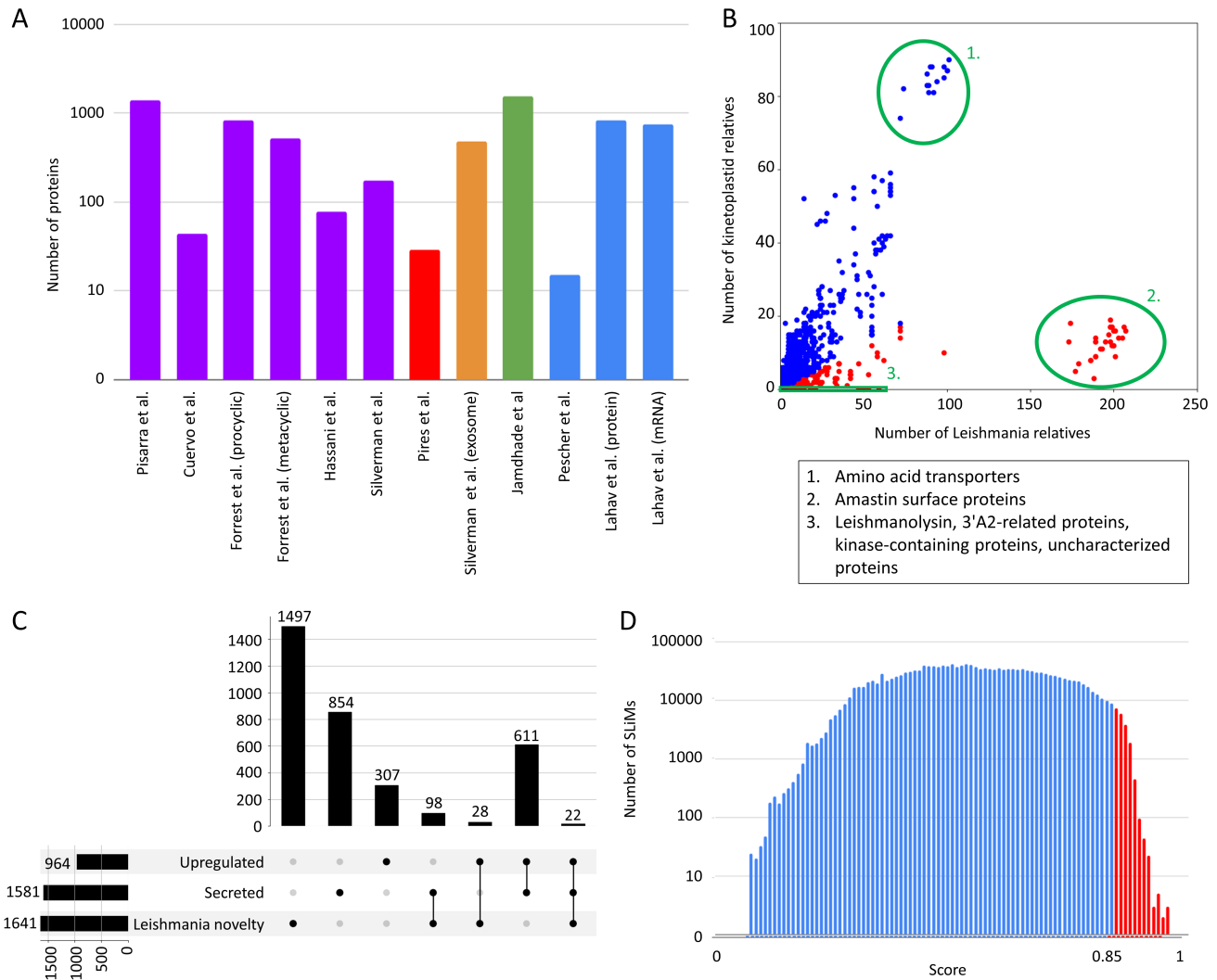
### 171 **AlphaFold2 provides an alternative way estimate structural features**

172 We used different methods to predict the structural features of proteins. Classical sequence-based  
173 methods can detect globular domains *Paysan-Lafosse et al. (2023)*, TM regions *Dobson et al. (2015)*  
174 and intrinsically disordered regions (IDRs) *Erdős et al. (2021)*. However, the use of AlphaFold2 (AF2)  
175 *Jumper et al. (2021)* provides alternative ways to obtain structural information. In LeishMANIAdb  
176 we used structures available in the AlphaFold database *Varadi et al. (2022b)* (however we could  
177 not find 3192 proteins ( 6% of all *Leishmania* proteins in LeishMANIAdb)). We not only displayed  
178 the predicted 3D structure of the proteins, but also information derived from the AF2 models such  
179 as the secondary structures and the position of the lipid bilayer for membrane proteins using the  
180 method introduced in the TMAAlphaFold database *Dobson et al. (2023)*. Although AF2 was originally  
181 built to predict protein structure, the scientific community quickly realized it is as much (if not more)  
182 efficient at predicting protein disorder *Akdel et al. (2022)*. To analyze IDRs we displayed predicted  
183 local distance difference test (pLDDT) values and relative surface accessibility from AF2. For IDR  
184 prediction in TM proteins, we tailored MemDis *Dobson and Tusnady (2021)* to incorporate features  
185 from AF2 instead of sequence-based predictors (see Methods).

### 186 **Short linear motif candidates that may hijack host cell regulation**

187 We scanned *Leishmania* proteins for SLiMs using the regular expressions stored in the Eukaryotic  
188 Linear Motif (ELM) resource *Kumar et al. (2022)*. Scanning SLiMs alone would mostly yield false pos-





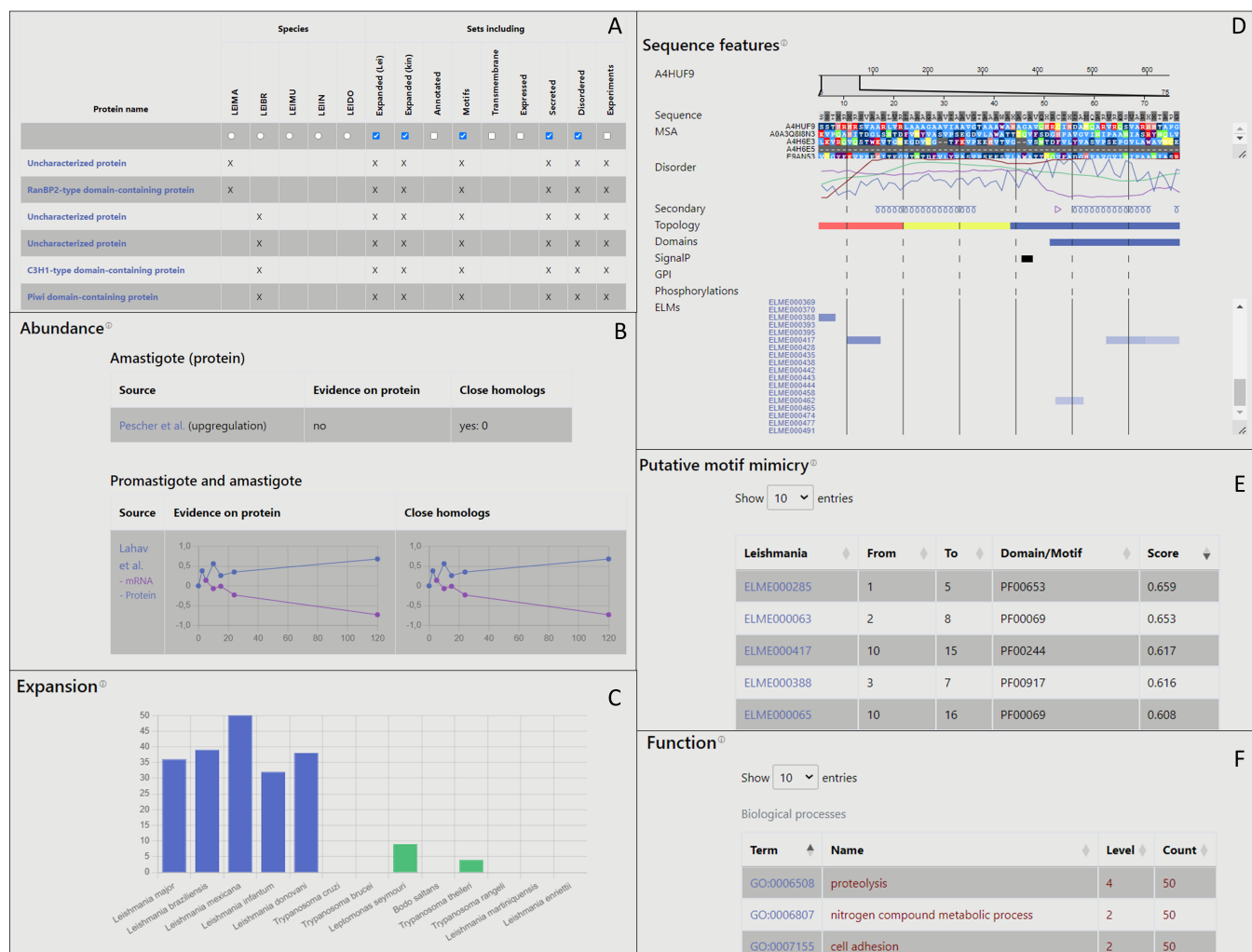
**Figure 1.** LeishMANIAdb content. All data were calculated on *Leishmania infantum*. A: Number of proteins in different proteomic datasets (purple: promastigote secretome, red: amastigote secretome, orange: exosome, green: housekeeping genes, blue: higher protein abundance level upon infection). B: Number of kinetoplastid and *Leishmania* close homologs. Each dot represents a protein (red: at least 80% of close homologs are in *Leishmania*, blue: other proteins). Green circles represent distinctive groups. C: Overlap between abundant, secreted and *Leishmania* novelty proteins (for more detail see text). D: Distribution of all predicted SLiMs with different scores. Red marks candidate motifs above 0.85 cutoff (for more details see text).

189 itive hits, so we developed a scoring system that ranges from 0 to 1, and that takes into account  
190 most information we collected. We aimed to develop a scoring system where conservation and  
191 accessibility/disorder has a reasonably high weight, while keeping in mind that proteomic experi-  
192 ments and localization information are a good way to narrow down the potentially large number of  
193 false positive hits. Unfortunately, due to the lack of data, in the case of *Leishmania* it is not possible  
194 to construct a benchmark set to evaluate motif scores. We can still assume that a good starting  
195 point can be when most predictions and proteomic data agree. Considering *Leishmania infantum*  
196 alone, we detected over a million putative motifs, from which 1.21% had a score above 0.85, on  
197 343 proteins (Figure 1/D).

## 198 **The LeishMANIAdb web resource**

199 To visualize all the collected and calculated information we developed an open-access resource. In  
200 LeishMANIAdb users can search for proteins using their UniProt Accession (AC), Entry name (for-  
201 merly ID), gene name and protein name. We also provide several protein sets as examples to help  
202 users browsing the database. Currently proteins are sorted based on 1) species: *L. braziliensis*, *L.*  
203 *donovani*, *L. infantum*, *L. major*, and *L. mexicana*; 2) manual curation data; 3) experimental data: se-  
204 creted proteins, protein abundance/mRNA level data, proteins with any kind of experimental data  
205 listed above; 4) computationally predicted information: proteins expanded in *Leishmania* (score  
206  $\geq 0.8$  - see Methods, Supplementary Material), transmembrane proteins, proteins with high dis-  
207 ordered content (at least 70% predicted disorder), proteins with high-scoring linear motifs (score  
208  $\geq 0.85$ ) and novel kinetoplastid proteins (proteins without SwissProt close homologs or Pfam do-  
209 mains). After searching (or selecting a protein set) users can further narrow their selection by  
210 choosing any other criterion (Figure 2/A).

211 The entry page for proteins consists of 10 sections, which are only visible if they contain data.  
212 The “Quick info” displays the protein name, species, cross-references and its number of amino acid  
213 residues. Data curation appears under the “Annotations” section. PPI (curated at the MIMix level),  
214 localization and function annotations are mirrored from close homologous proteins. We also dis-  
215 play functional annotations by Jardim et al. *Jardim et al. (2018)*. The “Localization” section contains  
216 high-throughput experiment data - promastigote and amastigote secretion, an exosome experi-  
217 ment and the glycosome. Protein localization, signal peptide and glycosylphosphatidyl (GPI) an-  
218 chor predictions are also displayed here. Since the reliability of both predictions and experiments  
219 may vary, we also display all this data for close homologous proteins, so users can quickly check  
220 the robustness of information. Furthermore, we also collected Gene\_Ontology (GO) *The Gene On-*  
221 *tology Consortium (2019)* annotations for cellular compartments. In this case, the specificity of  
222 the term (how deep it is on the tree) is shown in the level column. GO annotations are collected  
223 for all close homologous proteins too, and the number of occurrences of each term is displayed.  
224 We highlighted terms that are associated with the inspected protein itself that is displayed on the  
225 page. The “Abundance” module can display the mRNA and protein level experiments: static/single  
226 point (upregulated or not) or time-course experiments (e.g., mRNA and protein levels available for  
227 7 timesteps across 120 hours, Figure 2/B). In the “Expansion” section the number of close homologs  
228 are displayed by species with a color-code to identify *Leishmania* intracellular and extracellular/free-  
229 living relatives (Figure 2/C). The “Sequence features” displays various information (Figure 2/D). On  
230 the top, the gapless multiple sequence alignment (MSA) of proteins from the reference proteomes  
231 is visible. In this alignment gaps from the entry protein were removed (the original alignment with  
232 all strains can be downloaded) so other protein features could be visualized. Protein disorder, sec-  
233 ondary structures, transmembrane topology prediction, domains, signal peptides and GPI anchors  
234 and stage-dependent phosphorylation are also displayed. Predicted SLiMs are shown with a color-  
235 coded score (see Methods, Supplementary Material). In the “Structure” section the AF2 predicted  
236 structure is available (with the position of the membrane domain for TM proteins). The “Putative  
237 motif mimicry” section is the table format version of SLiMs from the “Sequence features” module  
238 (Figure 2/E). The “Function” section contains GO Molecular Function and Biological Process terms.



**Figure 2.** Figure 2: Layout of LeishMANIAdb: A: the search/browse result menu. B: Expression section of the entry page. C: Expansion section. D: Sequence features section. E: Putative motif mimicry section. F: Function section.

239 As done for the Cellular Component, terms are mirrored from close homologous proteins, and they  
 240 can be sorted based on their specificity (how deep they are located in the tree) and occurrences  
 241 considering homologs (Figure 2/F). Finally, each result from a BLAST search against SwissProt and  
 242 kinetoplastids relatives are listed in the “Homologs” table.

243 For each protein the full MSA, high-throughput experiments, annotation and predicted sequen-  
 244 tial features can be downloaded from the bottom of the page. Batch download is also available to  
 245 download the full database or different protein sets.

## 246 Discussion

### 247 Reliability of data

248 In LeishMANIAdb we aimed to collect high-throughput experimental data, PPI data on individual  
 249 proteins, predictions and localization information based on distant homologs. We noticed that the  
 250 amount of data from MS experiments differs highly, and therefore likely the quality also varies.  
 251 Not unexpected from high-throughput techniques applied to less-studied organisms, this can be  
 252 attributed to the quality of sample preparation, MS experimental techniques and most likely to  
 253 the sequences in the background databases. One striking finding was that the secretory datasets  
 254 contain a large number of proteins that are likely to take part in the housekeeping processes of

255 *Leishmania* cells, such as cytoskeletal proteins, nuclear histones and metabolic enzymes. Exosomes  
256 are known to contain a relatively high amount of „background“ proteins leaking from the cytosol  
257 of cells. Another explanation is that several housekeeping genes (such as intracellular chaperones  
258 and enzymes) are moonlighting proteins, they are generally constitutively expressed and have high  
259 levels of expression, while they are fulfilling other functions outside the cells *Jeffery (2018)*. Due  
260 to the lack of comparative studies, we cannot assess the enrichment ratios of secreted molecules,  
261 to see if there is selective exosomal packaging of a well-defined subset of *Leishmania* proteins.  
262 However, *Leishmania* exosomal-like secretion also differs from the typical exosomal sorting seen  
263 in other eukaryotic organisms because budding primarily initiates at the cell membrane, and not  
264 inside multivesicular bodies (endosomes). Therefore it is equally possible that in Leishmaniids, the  
265 budding is non-selective for its cytoplasmic cargos. Instead it would be initiated by cell surface  
266 receptors and primarily serve as a defense mechanism against membrane-attached host comple-  
267 ment and other immune complexes, removing them before they could damage the parasite mem-  
268 brane. Currently, testing of the latter hypothesis is impossible, since only soluble components but  
269 not the integral membrane proteins of *Leishmania* exosomes have been studied in depth in the  
270 above cited studies.

271 From a computational point of view, predicting any features on *Leishmania* proteins might be  
272 highly challenging, as methods established were mostly trained on sequences that show little or  
273 no similarity to *Leishmania* proteins. The 5 *Leishmania* reference proteomes contain 10 267 unchar-  
274 acterized proteins combined, which is 25% of LeishMANIAdb. TMAAlphaFold provides an objective  
275 quality measurement option for alpha-helical membrane proteins. When we compared the TM  
276 proteome of Homo sapiens and *Leishmania infantum* we noticed that the ratio of good and excel-  
277 lent quality structures was much lower in *Leishmania*, probably caused by the different coverage  
278 of kinetoplastid and human structures deposited into the PDB (Figure 3/A).

## 279 Case studies

280 LeishMANIAdb can be utilized for different purposes, and can be a good starting point for various  
281 analyses. We selected three examples that highlight some use cases of the resource.

282 Using the Browse menu, after selecting a category users can further narrow down their search  
283 for proteins selecting additional categories to refine the results. For instance, if users are looking  
284 for *Leishmania* SLiMs that may alter or rewire host cell regulation network, they can look for pro-  
285 teins that were experimentally proven to be secreted, and then select proteins with disordered re-  
286 gions because SLiMs are mostly located in IDRs. “Kinetoplastid novelty” selection ensures that the  
287 protein and its domains are not present in other organisms, while *Leishmania* novelty/expansions  
288 select proteins that are new or highly expanded in *Leishmania* species. Last, by selecting high-  
289 scoring motifs users get a list of proteins where the motif is most likely to be functional (Figure 3/B  
290 shows the Venn diagram of the selection). These proteins may be an interesting starting point for  
291 further analyses.

292 When performing systematic searches to identify possible parasite hits of integrin ligand motifs  
293 (that only functions in the host, as kinetoplastids have no integrins), we identified a striking set of  
294 examples in a family of poorly-known *Leishmania* genes called 3'A2 related ORFs. This kinetoplastid-  
295 specific family of genes is actually expanded in *Leishmania* species together with the canonically se-  
296 creted A2 proteins, which are known pathogenicity factors *Zhang and Matlashewski (2001)*. While  
297 the actual sequences of these proteins are poorly conserved and very little is known about their  
298 subcellular location, the *Leishmania* versions have at least one transmembrane region and a C-  
299 terminal cytoplasmic tail, with an N-terminal signal peptide (or possibly another TM segment).  
300 Nevertheless, in the predicted, largely disordered extracellular segment we observed multiple,  
301 short, conserved stretches that may have amyloidogenic properties (high Val, Ala and Gly content,  
302 upon visual inspection), presumably capable of oligomerization and amphiphilic interaction with  
303 membranes (Figure 3/C). A highly conserved cysteine residue preceding the first amyloidogenic  
304 sequence might help the homodimerization by forming a disulphide bridge with neighboring 3'A2

305 related protein. Strikingly, in *Leishmania infantum* and *Leishmania donovani* (both species capable  
306 of causing visceral leishmaniasis), the N-terminus of these proteins carries canonical RGD (Arg-Gly-  
307 Asp) sequences, immediately after the putative signal peptide cleavage site. In addition, *Leishmania*  
308 *donovani* and *Leishmania infantum* proteins contain an NGR motif where asparagine deamidation  
309 might yield an isoDGR motif. If these proteins are expressed on the cell surface, they might bind  
310 to host integrins in an oligomeric state, and might even attack the host membrane as if it were a  
311 beta-barrel pore-forming toxin. However, much more experiments are needed to test any of these  
312 hypotheses.

313 Amastins are a large family of kinetoplastid-specific membrane proteins that belong to the  
314 broader claudin-like superfamily, implicated in the maintenance of parasitophorous vacuoles *de Paiva*  
315 *et al. (2015)*. Accordingly, the majority of amastins have 4 tightly packed TM segments, with cy-  
316 tosolitic tail regions. Similarly to their vertebrate counterparts that form tight cell-cell junctions by  
317 complex oligomerization processes, amastins might also engage in a variety of interactions with  
318 internal as well as external, host proteins. Although their exact function is not known, among the  
319 221 identified amastins with 4 TM regions we looked for SLiMs that occur in multiple proteins.  
320 Predicted SLiMs (within disordered regions) were packed in their cytoplasmic tail regions (Figure  
321 3/D). Since these regions face inward the parasite, we further narrowed hits based on their bind-  
322 ing domain to be present in *Leishmania*. We identified multiple potential phosphorylation sites  
323 and protein-protein interaction motifs, such as SH3 ligands (*Leishmania* species do encode SH3 do-  
324 main proteins) as well as vesicular trafficking signals. The tail region of amastins seem to be highly  
325 variable, likely acting as a hotspot in the pathogen-host arms race.

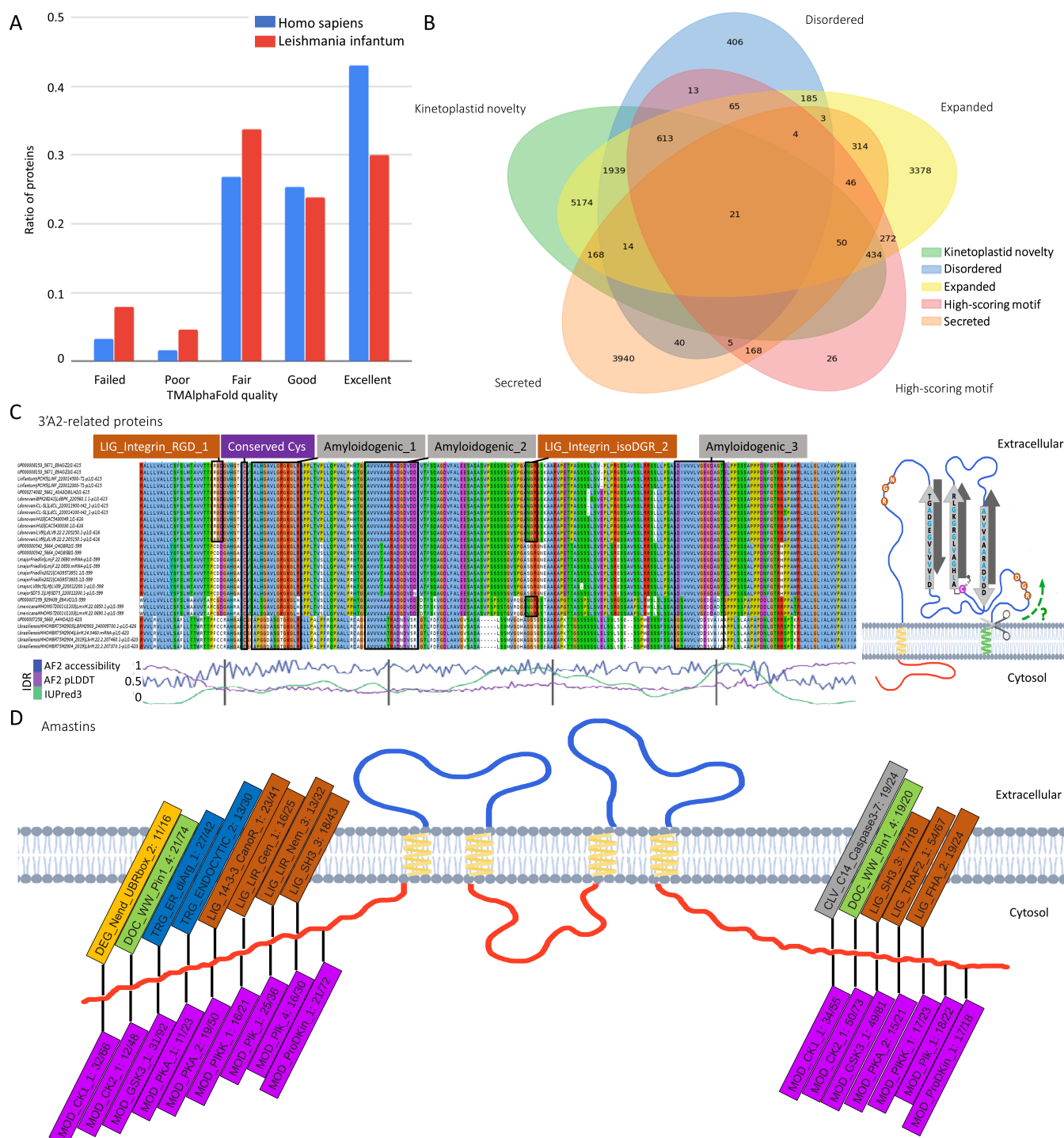
### 326 **Comparison with other resources**

327 In the past decades several databases were built to investigate *Leishmania*, however they are un-  
328 fortunately often offline and no longer updated by now. LeishCyc *Saunders et al. (2012)* focused  
329 on biochemical pathways. LeishDB *Torres et al. (2017)* included coding genes and non-RNAs and  
330 provided new annotation to them. The cysteine protease database in *Leishmania* species *Rana*  
331 *et al. (2012)* was designed to find data related to cysteine protease and LeishBase was a struc-  
332 tural database. There are a few active databases: Leish-EXP ([http://www.hpppi.iicb.res.in/Leish-](http://www.hpppi.iicb.res.in/Leish-ex)  
333 [ex](http://www.hpppi.iicb.res.in/Leish-ex)) (which has not so far been published in a peer-reviewed journal) contains proteins exclusively  
334 present in *Leishmania*. Leish-EXP incorporates localization tools, includes GO annotations and cal-  
335 culates physico-chemical properties of proteins. LmSmdB *Patel et al. (2016)* focuses on metabolic  
336 and biosynthetic pathways. TriTrypDB *Shanmugasundram et al. (2023)* is a kinetoplastid database  
337 that is part of the VEuPathDB resource *Amos et al. (2022)*. These databases contain a lot of ex-  
338 perimental data and various tools to analyze eukaryotic pathogens, but they are mostly focused  
339 on genomic data - although proteomic datasets, and some protein prediction algorithms are also  
340 incorporated.

341 There are also a handful of databases that include information on host-pathogen interactions:  
342 HPIDB *Ammari et al. (2016)*, PHIDIAS *Xiang et al. (2007)* and PHI-base *Urban et al. (2022)* contain  
343 information about PPIs between the host and pathogen, while ImitateDB *Tayal et al. (2022)* specif-  
344 ically focuses on motif mimicry. These resources contain no or very little data about *Leishmania*.

345 In LeishMANIAdb our main goal was to include protein information relevant to the infection and  
346 to complement previously established and still available resources. We included several proteomic  
347 datasets, and enriched experimental information with state-of-the-art prediction tools. Still, the  
348 most powerful way to explore uncharted proteomes is to inspect MSAs and check for conserved  
349 residues and regions - LeishMANIAdb contains precalculated alignments for all proteins. We also  
350 added hundreds of annotations to thousands of proteins, including localization and interaction  
351 information. While several databases seem to be shut down after a couple of years, our laboratory  
352 hosts several resources and we routinely update them. We plan to do so with LeishMANIAdb as  
353 well as to expand its repertoire to host-*Leishmania* interactions involving glycans and glycolipids,  
354 which play major roles in the infection.





**Figure 3.** Figure 3: A: Distribution of membrane protein quality levels of AlphaFold structure in Homo sapiens and *Leishmania infantum*. B: Venn diagram of proteins that are 1) secreted 2) novel kinoplastid 3) expanded (or new) in *Leishmania* 4) disordered 5) contain candidate SLIMs. C: left: Multiple Sequence Alignment of 3'A2 related proteins (alignment is available under UniProt AC: E9AGZ3). Amyloidogenic regions, conserved cysteine and Integrin-binding motifs are highlighted; right: proposed topology of 3'A2 related proteins D: Frequent SLIMs in the cytoplasmic tail regions of amastins (the numbers denote the unique/total occurrences).

## 355 Methods

### 356 Resources

357 Protein sequences were retrieved from UniProtKB (release 2022\_05) *UniProt Consortium (2023)*  
358 and from TriTrypDB *Shanmugasundram et al. (2023)* based on the UniProt cross-references (*L.*  
359 *braziliensis*, *L. donovani*, *L. infantum*, *L. major*, *L. mexicana*, *Bodo saltans*, *Leptomonas seymouri*, *Try-*  
360 *pansoma brucei*, *Trypansoma cruzi*, *Trypansoma rangeli*, *Trypanosoma theileri*). Homologs in other  
361 kinetoplastids and in SwissProt were searched with BLAST using e-value: 10<sup>-5</sup>; sequence iden-  
362 tity>20%; coverage>50%. In the “Homologs” section all results are displayed, however for most  
363 other sections (and calculation) we only used homologous proteins until the first non-kinetoplastid  
364 SwissProt hit considering sequence identity (termed as “close-homologs”); Further similar kine-  
365 toplastid proteins were therefore considered as a different homology group. Furthermore, we down-  
366 loaded strains belonging to the 5 selected *Leishmania* species from TriTrypDB. In this case a more  
367 stringent condition was used in BLAST, by setting E-value: 10<sup>-5</sup>; sequence identity>80%; cover-  
368 age>80%. All kinetoplastid species and strains were used to calculate motif conservation.

369 We prepared three different type of MSAs using ClustalOmega *Sievers et al. (2011)*: 1) “non-  
370 redundant” MSA using homologous proteins from kinetoplastid reference proteomes; 2) the same  
371 MSA but with gaps removed from the “reference” protein that is currently displayed on the web-  
372 page; 3) a more redundant MSA using homologous kinetoplastid proteins in all species and strains  
373 (used to calculate motif conservation).

374 High-throughput experiments were first mapped to the corresponding protein using the iden-  
375 tifier provided in the original paper, then mirrored to close *Leishmania* homologs if their sequence  
376 were identical.

377 IDRs were predicted using IUPred3 *Erdős et al. (2021)* and using the AF2 models’ pLDDT and  
378 accessibility values - the latter was calculated by DSSP *Joosten et al. (2011)*, normalized using max-  
379 imum values calculated as in Tien et al. *Tien et al. (2013)*, the exposed value threshold defined  
380 as suggested by Rost et al. *Rost and Sander (1994)*. In the case of TM proteins, IDRs were also  
381 predicted by MemDis *Dobson and Tusnády (2021)*. In this in-house modified version, the Position-  
382 Specific Scoring Matrices (PSSMs) were generated using kinetoplastid sequence library and sec-  
383 ondary structure and accessibility were calculated using AlphaFold2. Topology was predicted by  
384 CCTOP *Dobson et al. (2015)*, however to minimise sporadic erroneous predictions, after an initial  
385 prediction we performed a constrained iteration where the topologies of homologous proteins  
386 were used as a constraint. Using this approach, closely related proteins will likely have the same  
387 topology. Secondary structure elements derived from AF2 structures are also displayed. Pfam  
388 domains were identified using InterPro *Paysan-Lafosse et al. (2023)*. Protein localization was as-  
389 signed by GO *The Gene Ontology Consortium (2019)*, predicted by DeepLoc *Almagro Armenteros*  
390 *et al. (2017)* and SignalP6.0 *Teufel et al. (2022)*. NetGPI *Gíslason et al. (2021)* was used to predict  
391 GPI-anchors (all prediction results are visible, therefore in case of a contradiction it is up to the  
392 user to judge the results).

393 To detect SLiMs that may alter or rewire host cell regulation, we used the regular expressions  
394 from ELM *Kumar et al. (2022)* on all *Leishmania* sequences. We defined different context filters  
395 and merged them into a single score to rank motifs (for more details see Supplementary Material):  
396 1) Disordered: The score is the average of the IUPred3, AF2-based pLDDT and accessibility val-  
397 ues. These disordered scores were first transformed so they range from 0 to 1, with 0.5 being the  
398 threshold, before calculating their mean; 2) Conservation of the motif was checked among close  
399 homologs with some permission for slight misalignment, and penalizing motifs that are present  
400 across all kinetoplastids - notably, in this case proteins from different *Leishmania* strains were also  
401 considered; 3) Localization: we used a simplified (intracellular/extracellular) distinction. Motif local-  
402 ization was determined using ELM GO annotations, secretion information and CCTOP prediction,  
403 while the domain localization was determined from TOPDOM (Varga et al., 2016). We looked for  
404 motif-domain pairs where they both have the same simplified (in/out) localization; 4) mRNA level:

405 using transcriptomic experiments about expression data; 5) protein level: from experiments about  
406 protein abundance; 6) Secretion score based on secretome experiments; 7) Expansion score: re-  
407 flecting how much the protein is expanded in *Leishmania* species (strains not included) compared  
408 to all kinetoplastids; 8) Outgroups score favoring proteins without homologs in SwissProt. Struc-  
409 ture data reflects structure data deposited in the PDB *Varadi et al. (2022a)* before 26.03.2023 and  
410 the AlphaFold database (v3). All other data was downloaded in October, 2022 from the source  
411 databases.

## 412 **Manual curation**

413 We manually curated hundreds of proteins, using two strategies. First, we searched PubMed and  
414 Google scholar for “*Leishmania* host-pathogen protein interaction” and manually processed the re-  
415 sults. Each protein in the experiments was mapped to the corresponding UniProt entry. Then we  
416 mapped interaction data to the 5 *Leishmania* proteomes. When the experiment was performed on  
417 a protein from different species, we mirrored it to the closest homology group in LeishMANIAdb,  
418 and we also indicated on the webpage that the experiment is from a different protein. All interac-  
419 tions were reported according to the community standard MIMix level *Orchard et al. (2007)*.

420 Next we searched for possible surface proteins. For this task we considered all homologs. We  
421 collected topology prediction, GPI-anchor prediction, GO terms and also searched for homologous  
422 proteins that were measured *Bausch-Fluck et al. (2018)*; *Langó et al. (2017)* or predicted to be on  
423 the surface. We manually processed the entries using this approach, taking distant homologues,  
424 domain architectures and conservation patterns into consideration.

## 425 **Website design**

426 The LeishMANIAdb website is written in PHP (v8.0) using the Laravel (v9.19) framework. All down-  
427 loaded, predicted or calculated data are stored in a local MySQL (v8.0) database. To visualize se-  
428 quence features over amino acid sequences, we developed a javascript package using React (18.2),  
429 while 3D structures are visualized using the original (for non-TM proteins) or a locally modified ver-  
430 sion of Mol\* *Sehnal et al. (2021)* for TM proteins. The modified version can display the membrane  
431 as two planes around the investigated TM protein using the results of TMDet *Tusnády et al. (2005)*.

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