

1 **Exploring *Leishmania*-Host Interaction with Reactome, a Database of Biological  
2 Pathways and Processes**

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14 Running title: *Leishmania* infection in Reactome

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16 Keywords: *Leishmania*, Reactome, pathway, ORA, enrichment

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20

21 **Abstract**

22 Leishmaniasis is a parasitic disease with a wide range of clinical manifestations. Multiple  
23 aspects of the *Leishmania*-host interaction, such as genetic factors and modulation of  
24 microbial functions in host cells, influence pathogenesis, disease severity and treatment  
25 outcome. How do scientists contend with this complexity? Here, we work towards  
26 representing detailed, contextual knowledge on *Leishmania*-host interactions in the  
27 Reactome pathway database to facilitate the extraction of novel mechanistic insights from  
28 existing datasets. The Reactome database uses a hierarchy of abstractions that allows for

29 the incorporation of detailed contextual knowledge on biological processes matched to  
30 differentially expressed genes. It also includes tools for enhanced over-representation  
31 analysis that exploits this extra information. We conducted a systematic curation of  
32 published studies documenting different aspects of the *Leishmania*-host interaction. The  
33 “*Leishmania* infection pathway” included four sub-pathways: phagocytosis, killing  
34 mechanisms, cell recruitment, and *Leishmania* parasite growth and survival. As proof-of-  
35 principle of the usefulness of the released pathway, we used it to analyze two previously  
36 released transcriptomic datasets of human and murine macrophages infected with  
37 *Leishmania*. Our results provide insights on the participation of ADORA2B signaling  
38 pathway in the modulation of IL10 and IL6 in infected macrophages. This work opens the  
39 way for other researchers to contribute to, and make use of, the Reactome database.

40 **Importance**

41 Leishmaniasis is a neglected disease infectious disease which affects more than 1.5 million  
42 people annually. Many researchers in the field apply -omic technologies to dissect the basis  
43 of clinical and therapeutic outcomes and access drug targetable features in the host-parasite  
44 interaction, among others. However, getting mechanistic insights from -omics data to such  
45 end is not an easy task. The most common approach is to use the -omics data to inquire  
46 pathways databases. The retrieved list of pathways often contains vague names that lack the  
47 biological context. In this study, we worked to create the *Leishmania* infection pathway in  
48 the Reactome database. With two practical examples from transcriptomics and microarray  
49 data, we demonstrated how this pathway facilitates the analysis of such data. In both  
50 datasets, we found a common mechanism of IL10 and IL6 production that the authors did  
51 not advert in their previous analysis, providing proof-of-principle of the tool’s enhanced  
52 potential for knowledge extraction. *Leishmania* infection pathway is in its first version, and  
53 must be expanded to cover the current knowledge base of the *Leishmania*-host interaction.  
54 We strongly encourage contributions from domain experts for the completion of  
55 *Leishmania* infection pathways.

56

57

58 **Introduction**

59 The interaction between a parasite and its host is a fight for dominance. Who wins? How  
60 does disease progress? A multitude of factors determine the course of infection. These  
61 depend upon both the status of the host and the parasite. Omics data shed light on the  
62 multitude of activated mechanisms within the host-parasite interactome, by providing long  
63 lists of differentially expressed (DE) molecules. However, such data is a means for  
64 generating knowledge, not an end. The ultimate goal is to interpret this data to build a  
65 mechanistic understanding of the interactions at hand. This exercise is made tractable by  
66 the existence of pathway databases that curate and organize current knowledge (1–3).  
67 Typically, the databases are used to find known biological processes that could underlie the  
68 data. A common methodology is Over-Representation Analysis (ORA). This takes the set  
69 of DE genes from the data, and iteratively compares them to the set of genes involved in  
70 each separate pathway in the database. It uses the overlap between these two sets to predict  
71 the statistical likelihood of the biological pathway being represented in the data (4).  
72 Mechanistic hypotheses on the processes underlying the data are then proposed by the  
73 researcher, based on the ORA results.

74 Biological process labels within a database often lack context (e.g. ‘immune system’). Does  
75 the process occur within one particular cell type, or more? Across species? In a diseased  
76 organism? In the context of a pathogen-host interaction? It is difficult to build detailed  
77 hypotheses from such labels using ORA, or indeed other analytical approaches.

78 The Reactome database builds a hierarchy of abstractions into which the observed features  
79 of any biological process can be incorporated. At the top level of the hierarchy, high level  
80 characteristics are represented: Is it a disease? Is it infectious or metabolic? At lower levels,  
81 features such as specific, temporally-ordered sequences of cellular processes are  
82 represented (3). Choices such as how many levels of abstraction to include, and what each  
83 should represent, depend to some extent upon the expertise of the curator. Therefore, it is  
84 critical that the curator has expert domain knowledge or that they collaborate with an  
85 appropriate expert in the field.

86 The purpose of this work was to add representative features and variability of the  
87 *Leishmania* spp.-host interaction into the Reactome database. In the terminology of the  
88 database, this representation is known as a ‘pathway’. Our pathway is sufficiently flexible

89 as to allow for expansion and revision as new results are published. It incorporates detailed  
90 information on biological processes known to be activated during the *Leishmania*-host  
91 interaction. In particular, we focused on those processes correlated with the outcome of  
92 infection.

93 We explicitly demonstrate the utility of our database curation. We took two existing  
94 datasets to which ORA or manual revision of the literature was previously applied. With  
95 our expanded database, we uncover new mechanistic insights. In the long term, we hope  
96 that the research community will be able to use our pathways as a source of primary  
97 consultation, and as a curated database for functional and mechanistic interpretation of new  
98 data derived from -omics technologies, functional tests, or *in-silico* experiments. We  
99 believe that this pathway will enable fast and curated access to the integrative mechanisms  
100 of importance in leishmaniasis.

101

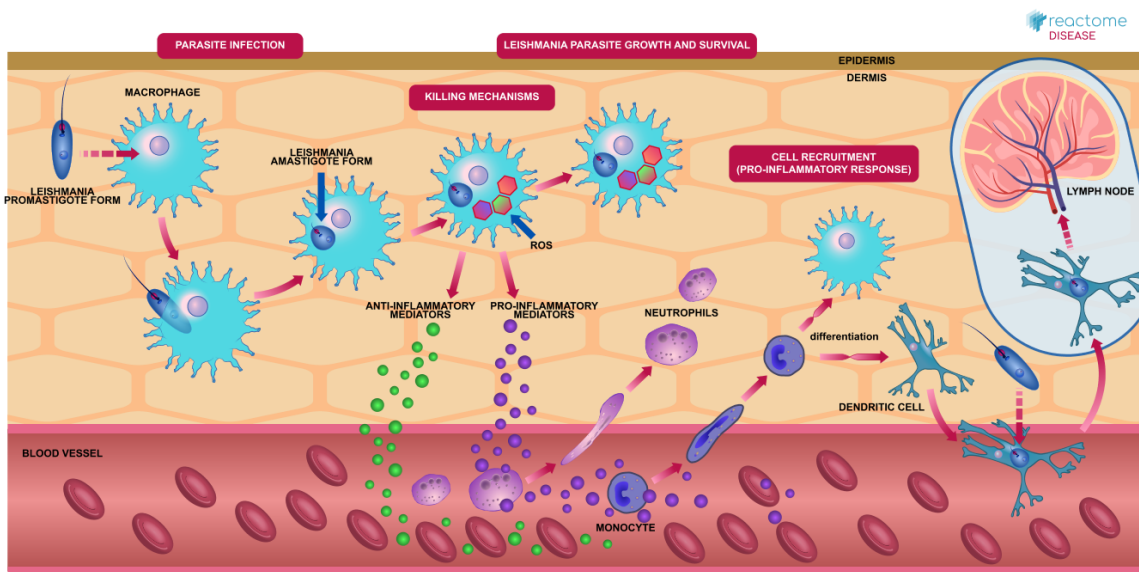
## 102 **Results**

### 103 **Abstracting the top-level pathway: phagocytosis, killing mechanisms, cell recruitment 104 and responses favoring *Leishmania* parasites.**

105 The curation of a pathway in Reactome starts by structuring pathways into hierarchical  
106 sections. The parent *Leishmania* infection pathway was structured into four subpathways  
107 describing the major processes involved: phagocytosis, killing mechanisms, cell  
108 recruitment and *Leishmania* parasite growth and survival. For each subpathway, there is  
109 extensive literature on the specific host responses to *Leishmania* infection, and their  
110 implication in the outcome of infection (5–7).

111 *Leishmania* parasites are transmitted to the host through the bite of a sand-fly that injects  
112 the motile promastigote form into the dermis of humans and other warm-blooded animals.  
113 Therein, the parasite interacts with the host cell(s) to establish the intracellular niche, where  
114 it will adopt the amastigote form (8). Paradoxically, macrophages, professional  
115 phagocytes of the innate immune system, are the main host cells for *Leishmania*. The first  
116 interaction of the parasite with the host cell is crucial to the outcome of the infection (5).  
117 The type of phagocytic receptor or pattern recognition receptor stimulated might influence

118 the signaling cascade(s) that will trigger or inhibit cellular mechanisms involved in parasite  
119 killing or permissiveness for infection. Deregulated immune responses contribute to  
120 pathology (9). Pro and anti-inflammatory mediators must be expressed at the “right” time  
121 and in the “appropriate” magnitude, in order to have a healing response (9). This discussion  
122 motivates our abstraction of existing knowledge into the categories of phagocytosis, killing  
123 mechanisms, cell recruitment (pro-inflammatory response), and *Leishmania* parasite growth  
124 and survival (figure 1).



126 **Figure 1: Textbook-style diagram representing the top-level pathway “*Leishmania***  
127 **infection”**. The major steps occurring in the dermis were compartmentalized into four  
128 categories: phagocytosis, killing mechanisms, cell recruitment, and *Leishmania* parasite  
129 growth and survival. On the webpage (<https://reactome.org/PathwayBrowser/#/R-HSA-9658195>), the magenta rectangular labels are interactive and take the user to the content of  
130 each subpathway.  
131

### 132 ***Leishmania* infection pathways: From a sketch on paper to Reactome database.**

133 Reactome’s curation tool is a graphical user interface (GUI), that connects to its central  
134 database with which new information can be added to existing or new pathways. We started  
135 with a sketch of the overall pathways we wanted to curate for the first version of the  
136 “*Leishmania* infection pathway”. Then, we identified what molecules/entities and reactions  
137 already existed in the database and which ones needed to be added. Similarly, we accounted

138 for the molecular interactions that were already described in existing Reactome pathways.  
139 If new reactions were required, they were created from scratch, supported by literature  
140 references. Table 1 summarizes these metrics.

141 **Table 1: Curation and annotation metrics for the creation of *Leishmania* infection**  
142 **pathways in Reactome.**

Pathway	Entities		Reactions/Catalysis/Regulation		New literature references
	New	Re-used	New	Re-used	
Phagocytosis	9	141	12	14	7
Killing mechanisms	5	18	9	0	8
Cell recruitment	3	28	3	24	9
<i>Leishmania</i> parasite growth and survival	15	51	19	12	17
Total	32	238	42	50	41

143

144 **Structuring the lowest-level pathways: signaling cascades traceable from a membrane**  
145 **protein to the production of effector molecules.**

146 As part of the host response against *Leishmania*, many signaling cascades are modulated  
147 (activated or inactivated). Once a cellular/membrane receptor is stimulated, the downstream  
148 signal transduction can result in the activation of many molecules with different effects in  
149 the system (e.g. interleukins inducing the polarization of several types of T-cells, or  
150 chemokines mediating the recruitment of immune cells to different tissues, among others).  
151 That is why general pathway labelling such as “TNF-signaling”, might not be informative  
152 enough, and can allow for erroneous biological interpretations if the gene lists contained  
153 within enriched pathways are not carefully analysed. There could be many regulatory  
154 processes that favor one direction rather than another in a specific signaling pathway (as it  
155 can be noticed in TNF-pathway in Reactome, R-HSA-75893  
156 <https://reactome.org/PathwayBrowser/#/R-HSA-75893> and KEGG, hsa04668-  
157 [https://www.genome.jp/kegg-bin/show\\_pathway?hsa04668](https://www.genome.jp/kegg-bin/show_pathway?hsa04668)). Therefore, we structured the  
158 lowest-level pathways in each of four subpathways, starting off from an activated  
159 membrane protein (e.g., receptors, ion channels or enzymes). This was followed by  
160 inclusion of signalling and accessory molecules and finished with synthesis of effector

161 molecules that are consistent with the overall biological processes underlying the  
162 subpathway (e.g., reactive oxygen species –ROS- for “killing mechanisms”).

163 For this first version of the *Leishmania* infection pathway, we chose the membrane proteins  
164 FCGR3A, FZD7, P2RX4, P2RX7, ADORA2B, and CD163 and their downstream signaling  
165 cascades. Although additional membrane molecules are known to participate in the initial  
166 macrophage-*Leishmania* interaction, such as complement receptors, toll-like receptors -  
167 TLRs-, or chemokine receptors among others (10–12), we selected less “classical”  
168 membrane molecules to increase the breath of mechanistic interpretation of host-  
169 *Leishmania* –omic datasets. Shown in figure 2 are the structures of the four subpathways,  
170 each of which contains a set of reactions. Supporting references evidencing their relevance  
171 in the context of leishmaniasis will be discussed below.

-  + Leishmania infection
-  + Parasite infection
-  + Leishmania phagocytosis
  - ⊕  + FCGR3A-mediated phagocytosis
-  + Killing mechanisms
  - ⊕  + WNT5:FZD7-mediated leishmania damping
  - ⊕  + Cell recruitment (pro-inflammatory response)
  - ⊕  + Purinergic signaling in leishmaniasis infection
-  + Leishmania parasite growth and survival
  - ⊕  + Anti-inflammatory response favouring Leishmania parasite infection
    - ⊕  + CD163 mediating an anti-inflammatory response
    - ⊕  + FCGR3A-mediated IL10 synthesis
    - ⊕  + ADORA2B mediated anti-inflammatory cytokines production
    - ⊕  + LTC4-CYSLTR mediated IL4 production

172

173 **Figure 2:** Hierarchical structure of the *Leishmania* infection pathway showing the four sub-  
174 pathways and their contents. Each indentation introduces a new subpathway; as an  
175 example, “parasite infection” is the parent pathway for the subpathway “*Leishmania*  
176 phagocytosis”, which, to date only contains *FCGR3A*-mediated phagocytosis.

177

178 **I. Parasite Infection/Leishmania phagocytosis:** The phagocytosis subpathway  
179 was built to account for the different types of phagocytic receptors that  
180 *Leishmania* parasites can utilize for their entry into host cells. We started by

181 adding FCGR3A-mediated phagocytosis. To build it, we referenced the pre-  
182 existing Reactome pathway “FcG receptor (FCGR) dependent phagocytosis”  
183 (<https://reactome.org/content/detail/R-HSA-2029480>). Overall, 14 reactions  
184 were re-used from this pathway while 12 new reactions were created to  
185 represent phagocytosis in the context of *Leishmania* infection (table 1). The  
186 starting point was the binding of immunoglobulin G antibodies (IgG) to either  
187 an unknown *Leishmania* amastigote (abbreviate as: Lma) surface molecule or  
188 the glycoinositol phospholipid - GIPL (13, 14), as shown in figure 3a. These  
189 interactions correspond to binding reaction types in Reactome, with the product  
190 of binding being a complex comprising the inputs. The complexes “IgG:Lma  
191 surface” and “IgG:GIPL” represent the opsonization of the *Leishmania*  
192 amastigote by the antibody IgG, during a “second round” of host-parasite  
193 contact where the proliferative form and infective form in the host is the  
194 amastigote. We assumed the same course when opsonization occurs via these  
195 known, or other unknown molecules. Therefore, we collated the two complexes  
196 into one entity, which in Reactome is represented by the defined set “IgG:Lma  
197 antigens”. From here, downstream reactions continue towards the activation of  
198 actin filaments that then continue to the formation of the phagocytic cup. The  
199 overall diagram depicting each step of the pathway can be accessed through this  
200 link <https://reactome.org/PathwayBrowser/#/R-HSA-9664422&PATH=R-HSA-1643685,R-HSA-5663205,R-HSA-9658195,R-HSA-9664407,R-HSA-9664417>. A  
201 close-up depicting a portion of the pathway is found in figure 3A.  
202

203  
204 The route of entry into the macrophage can affect the fate of *Leishmania*  
205 parasites (5). We expect to incorporate the internalization processes that are  
206 mediated by other receptors into the *Leishmania* phagocytosis subpathway. This  
207 includes complement receptors (CR3 and CR1), mannose receptor-MR, and  
208 fibronectin receptors-FNRs (5). Similarly, the “Parasite infection” subpathway  
209 will be populated with the steps that describe the maturation of the phagocytic  
210 cup, and so on.

211       II. **Killing mechanisms:** this subpathway was designed to contain the signaling  
212       cascades that converge in the production of antimicrobial molecules in the  
213       context of leishmaniasis. We started by curating the activation of the receptor  
214       Frizzled-7 (FZD7) by the ligand WNT5 and its downstream cascade. To build  
215       this pathway we reviewed publications that contain the original experimental  
216       data used to determine the reactions details (15–19). WNT5 is known for being  
217       a highly specific regulated gene in response to microbial infection (20–22)  
218       including leishmaniasis (23), where it seems to be involved in mechanisms that  
219       dampen the parasite load within the macrophage. Complementary, FZD7 acts as  
220       a receptor of WNT5 which, upon binding, is implicated in the initiation of the  
221       non-canonical WNT pathway that leads to re-organization of the cytoskeleton to  
222       allow a process called planar cell polarity (PCP) (22). The activation of the  
223       WNT5:FZD7 non-canonical signaling cascade that drives PCP is being studied  
224       for its involvement in inflammatory responses (24). Treatment of RAW264.7  
225       macrophages with recombinant WNT5 induced NADPH oxidase-mediated ROS  
226       production, which has been suggested to contribute to the macrophage control of  
227       *L. donovani*. Consequently, detailed understanding of how the WNT signaling  
228       network defines host responses to infection could be important to identify new  
229       potential therapeutic targets (22).

230

231       We represented in 9 reactions, the activation of FZD7 by the WNT5 ligand,  
232       resulting in the production of ROS (table 1). Unlike the phagocytosis pathway,  
233       these reactions correspond to a host's response to the infection, even if no  
234       parasite components are depicted in the diagram (found at  
235       <https://reactome.org/PathwayBrowser/#/R-HSA-9673324&PATH=R-HSA-1643685,R-HSA-5663205,R-HSA-9658195,R-HSA-9664420>). In future versions we will  
236       incorporate the cross talk with signaling cascades, like TLR-signaling, that  
237       activate antimicrobial functions and synthesis of antimicrobial molecules.  
238

239

240       III. **Cell recruitment:** this subpathway was aimed at bringing together signaling  
241       pathways that converge in the induction of gene expression and synthesis of

242 chemokines and pro-inflammatory cytokines. It is known that a  
243 proinflammatory response early in the infection enhances host cell microbicidal  
244 mechanisms (25). However, the recruitment of inflammatory cells to the site of  
245 infection, once the parasite load has been controlled, transforms the course of  
246 infection and can lead to immunopathology (9). Therefore, it is important to  
247 curate and represent specific pathways that have shown to be activated upon  
248 *Leishmania* infection, resulting in the production of pro-inflammatory  
249 mediators.

250

251 The first specific mechanism we curated was the activation of the purinergic  
252 receptors P2RX4 and P2RX7. The liberation of ATP normally occurs in tissues  
253 facing stressful stimuli such as infection (26). Binding of ATP to purinergic  
254 receptor activates the inflammasome leading to subsequent activation of  
255 interleukin 1 beta-IL1 $\beta$ , which promotes the recruitment and activation of  
256 macrophages (27). We represented this process in 27 reactions (table 1), that  
257 included a regulatory step mediated by NTPDase1 and NTPDase5, which  
258 reduces ATP to adenosine (28). The molecular diagram can be found at  
259 <https://reactome.org/PathwayBrowser/#/R-HSA-9664424&SEL=R-HSA-9660826&PATH=R-HSA-1643685,R-HSA-5663205,R-HSA-9658195>.

260

261  
262 There are many other pathways promoting cell recruitment as a response to  
263 *Leishmania* infection with different consequences for the parasite and the host  
264 (29). In future expansions of this subpathway, it would be possible to highlight  
265 cross talk between different cascades that target the same effector molecules.

266

267 **IV. *Leishmania* parasite growth and survival:** this subpathway covers the host  
268 responses that favor intracellular parasite survival, and the mechanisms used by  
269 the parasite to hijack host cell functions. To survive as an intracellular parasite,  
270 *Leishmania* evades the activation of host cell microbicidal machineries. Many  
271 mechanisms facilitate this purpose. On the host side, the production of anti-  
272 inflammatory mediators often occurs alongside the repression of expression of

273 antimicrobial molecules, together with the recruitment of regulatory immune  
274 cells (e.g., regulatory T-cells). From the parasite side, inactivation of host  
275 molecules through mechanisms such as cleavage or activation of phosphatases  
276 are part of its repertoire (9). Induction of anti-inflammatory molecules was the  
277 first mechanism that we curated, compiling the steps that describe the cleavage  
278 of the membrane protein CD163, the activation of the receptors FCGR3A and  
279 ADORA2B, and ending with the corresponding production of the known anti-  
280 inflammatory molecules sCD163, IL4, and IL10, as well as the dual functioning  
281 IL6.

282

283 Macrophages infected with *L. amazonensis* or *L. donovani* strongly express the  
284 membrane protein CD163 (30–32), and soluble CD163 (sCD163) has been  
285 proposed as biomarker of visceral leishmaniasis. The hypothesis of the  
286 association between sCD63 and an anti-inflammation status is that it interferes  
287 with the proliferation of T-cells (33, 34). sCD163 is formed from the increased  
288 shedding of CD163 mediated by the metalloprotease ADAM17 (35, 36).  
289 Posteriorly, it might translocate to the cytoplasm of T-cells (through an  
290 unknown mechanism) where it binds with a protein involved in the proliferation  
291 process (33, 34). In “CD163 mediated anti-inflammatory responses” we  
292 represented, in 9 reactions, the production of sCD163 including the steps that  
293 precede the activation of ADAM17. Additionally, we included the positive  
294 regulation of glucocorticoids, IL6 and IL10 on CD163 gene expression (37–40).  
295 In figure 3B we show the molecular diagram depicting the pathway.

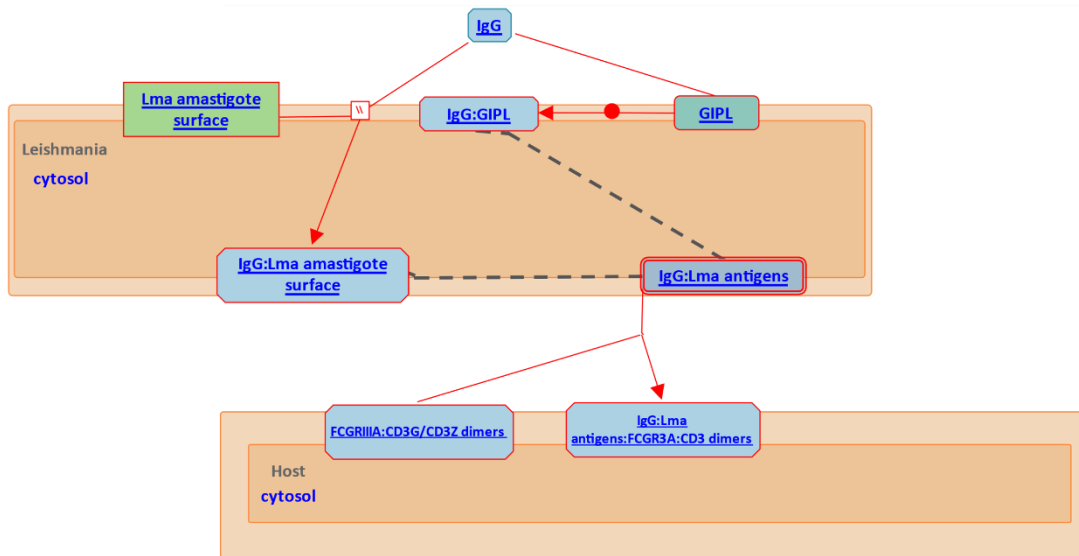
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297 IL-10 is an important immunoregulatory cytokine produced by many cell  
298 populations; in macrophages it is induced after the stimulation of TLRs, FCG  
299 receptors or by TLR-FCGR crosstalk (41). Classically, its function is considered  
300 to be the limitation and termination of inflammatory responses and the  
301 regulation of differentiation of several immune cells (42). In the context of  
302 leishmaniasis, IgG-opsonized amastigotes have been shown to induce IL10  
303 production through FCGRs, which in turn suppresses the killing mechanisms in

304 phagocytic cells (14). We represented, in 21 steps, the activation of FCGR3A  
305 that leads to the activation of the transcription factor CREB1, ending with the  
306 production of IL10.

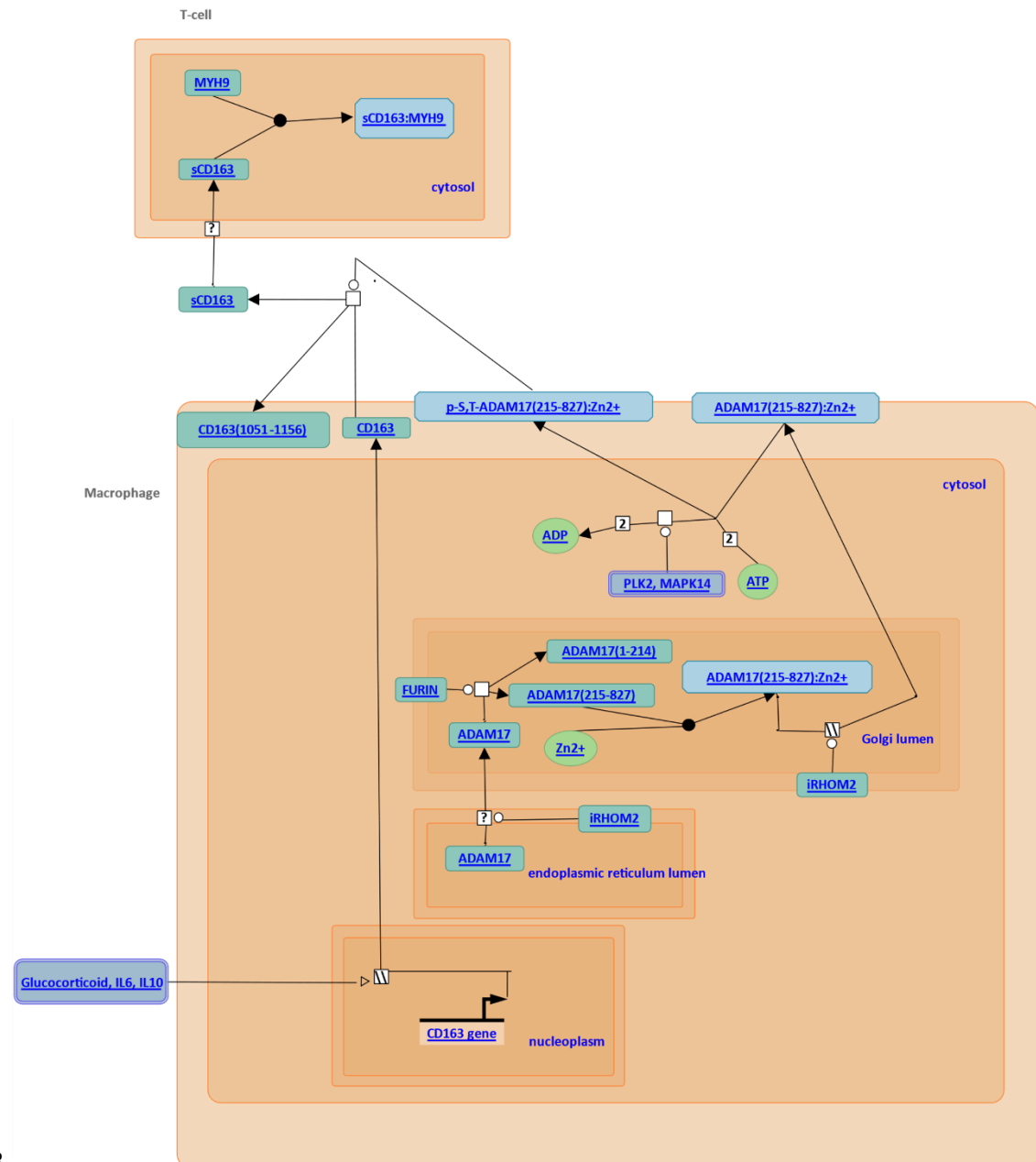
307

308 Finally, we curated ADORA2B-mediated anti-inflammatory responses.  
309 ADORA2B is a receptor for the ribonucleoside adenosine. Its activation leads to  
310 the production of anti-inflammatory cytokines which have been shown to favor  
311 *Leishmania* infection and survival (43–45). Apparently, this pathway exerts an  
312 opposing/regulatory response to the purinergic signaling pathway. The blockade  
313 in the production of pro-inflammatory cytokines may come with the inhibition  
314 of killing mechanisms (46). We represented this pathway with 6 reactions,  
315 starting with the binding of adenosine to ADORA2B, and ending with synthesis  
316 of IL6. Both FCGR3A and ADORA2B signaling pathways activate  
317 transcription factors, generating a positive feedback loop for transcription of  
318 more anti-inflammatory cytokines (47, 48). In future versions, these reactions  
319 might be incorporated, as well as other pathways leading to the synthesis of  
320 other anti-inflammatory mediators known to be induced during the *Leishmania*-  
321 host interactions. Moreover, other mechanisms that favor the persistence of  
322 *Leishmania* parasites must be added into new subpathways (e.g., Polyamine  
323 synthesis). The representation of these pathways in Reactome is available in the  
324 following link: <https://reactome.org/PathwayBrowser/#/R-HSA-9662851&PATH=R-HSA-1643685,R-HSA-5663205,R-HSA-9658195,R-HSA-9664433>.  
325



326

A.



327 B.

328 **Figure 3:** Standard graphical representation of pathways in Reactome. **A.** Fragment of the  
329 diagram for FCGR3A-mediated phagocytosis. Shown are the reactions corresponding to the  
330 parasite opsonization process by IgG. Parasitic components are highlighted in red. **B.**  
331 CD163 mediating an anti-inflammatory response. Each diagram shows the participating  
332 entities in the granularity of chemical compounds (green ovals), proteins (green rectangles),  
333 complexes (blue rectangles) and sets (blue rectangles with a double border). The

334 arrangement of the entities in the reactions can be easily followed on the web page, by  
335 clicking on the arrows that connect adjacent steps.

336

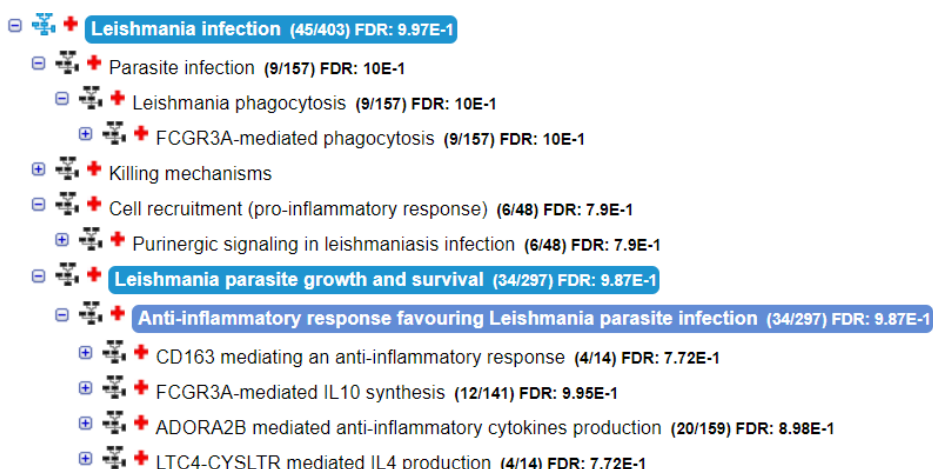
### 337 ***Leishmania* infection pathways enhancing transcriptome data analysis**

338 The first version of *Leishmania* infection pathways was released in March 2020. All  
339 released data can make use of Reactome pathway analysis tools. We used the ORA tool to  
340 test the impact of context-dependent labels on published datasets (prior march-2020) that  
341 explored the *Leishmania*-host interaction.

342 In 2015, Dillon *et al.* (49) explored the early response of macrophages to infection with *L.*  
343 *major*, using RNA-seq. With the differentially expressed (DE) gene list (>|2|-fold,  
344 uninfected macrophages versus infected macrophages at 4 hours post-infection), they  
345 performed ORA against the curated pathways in the KEGG database. For the up-regulated  
346 genes, the results included cytokine-cytokine receptor interactions, TNF-signaling pathway  
347 and NFkappa B-signaling pathway, among other pathways. An overall interpretation of  
348 these enriched pathways may suggest that the early infection of macrophages by *L. major*  
349 leads to induction of pro-inflammatory responses; as an example, TNF-signalling as  
350 represented in KEGG (PATHWAY: hsa04668), classically leads to the recruitment of  
351 inflammatory cells. However, transduction of the signal may induce the activation of  
352 factors that contribute to opposite responses, such as tissue regeneration (with VEGF and  
353 EDN1) (50, 51). Interestingly, among their gene-specific analyses, Dillon *et al.* identified a  
354 set of genes involved in anti-inflammatory responses (*Csf1*, *Csf3*, *Il10*, *Il11r*, *Il1rn*, *Socs3*,  
355 *Hmox1*, *Egfr* and, *Vegf*). However, the mechanisms that lead to production of these effector  
356 molecules, or how they contribute to achieve or maintain the underlying immune status  
357 couldn't be inferred from the KEGG pathway analysis. To overcome this gap, we  
358 implemented ORA from the same DE gene list (supplementary table S1) in Reactome.  
359 Among enriched pathways (table S1), were some of the *Leishmania* infection subpathways.  
360 As reported by Dillon and colleagues, FCGR3A subpathway (figure 4A) was found  
361 downregulated. Interestingly, the most over-represented pathway was the ADORA2B  
362 mediated anti-inflammatory cytokine production (Fold change-FC = 3.25). This pathway  
363 contributes to the expression of IL6 (FC = 7.68) and IL10 (FC = 17.05), through the

364 activation of the transcription factor CREB (FC = 2.75) (figure 4B). This pathway,  
365 simultaneously, leads to the activation of killing mechanisms, as well as to the production  
366 of IL10. Neither authors nor we found pathways involved in the former. Therefore, this  
367 suggests an alternative mechanism mediated by ADORA2B, that leads to production of  
368 IL10, and does not induce parallel pro-inflammatory consequences or the activation of  
369 antimicrobial molecules.

370 Complementary, we analyzed the microarray data from Gregory *et al.* (52) which  
371 represents the transcriptomic response of murine macrophages to infection with *L. major*  
372 and *L. donovani*. The authors stated that there were few differences in the number of genes  
373 and the magnitude of the expression with both species. Like Dillon's dataset, the  
374 ADORA2B subpathway was enriched among upregulated genes (figure 4C and  
375 supplementary table S2). These results highlight the identification of ADORA2B-pathway  
376 driving anti-inflammatory cytokine production, consistently activated in macrophages  
377 infected with different *Leishmania* species. These results substantiate the usefulness of the  
378 *Leishmania* infection pathways for gaining mechanistic insights from new and previously  
379 published data. In expanded versions of this pathway, secondary analyses like the one we  
380 performed, would shed light on response patterns against the infection with different  
381 *Leishmania* species, as well as species-dependent responses.

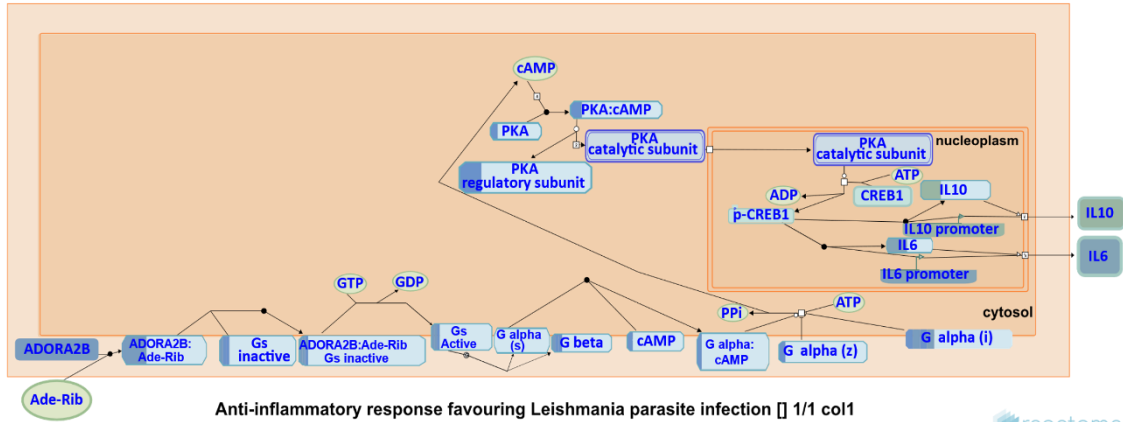


382 A. **CD163 mediating an anti-inflammatory response (4/14) FDR: 7.72E-1**

383

384

385



386 B.  [Anti-inflammatory response favouring Leishmania parasite infection](#) [1/1 col1]

- + Leishmania infection (18/403) FDR: 8.97E-1
- + Parasite infection (6/157) FDR: 8.93E-1
- + Leishmania phagocytosis (6/157) FDR: 8.93E-1
- + + FCGR3A-mediated phagocytosis (6/157) FDR: 8.93E-1
- + + Killing mechanisms (1/20) FDR: 6.88E-1
- + Cell recruitment (pro-inflammatory response) (3/48) FDR: 6.34E-1
- + + Purinergic signaling in leishmaniasis infection (3/48) FDR: 6.34E-1
- + Leishmania parasite growth and survival (8/297) FDR: 9.96E-1
- + Anti-inflammatory response favouring Leishmania parasite infection (8/297) FDR: 9.96E-1
- + + CD163 mediating an anti-inflammatory response
- + + FCGR3A-mediated IL10 synthesis (4/141) FDR: 9.64E-1
- + + ADORA2B mediated anti-inflammatory cytokines production (6/159) FDR: 9E-1
- + + LTC4-CYSLTR mediated IL4 production

387 C.

388 Figure 4: Results of applying ORA in Reactome, on the datasets generated by Dillon *et al.*  
389 and Gregory *et al.* A. Number of genes found in each subpathway from Dillon's dataset,  
390 with the associated false discovery rate (FDR) B. ADORA2B pathway enriched in Dillon's  
391 dataset shows upregulated genes involved in ADORA2B signaling cascade leading to the  
392 production of IL6 and IL10. C. Number of genes found in each subpathway from Gregory's  
393 dataset, and associated FDR.

## 394 Discussion

395 Over-representation analysis (ORA) has become one of the standard methods for extracting  
396 mechanistic information from -omics datasets. As part of the workflow, ORA matches the  
397 omics data with molecular data curated in pathway databases. As a result, it gives a list of  
398 pathway names in which the genes are known to be involved. Mechanistic insight must  
399 then be built from a list of labels, a problem made much harder when labels lack biological

400 context. There are several specific cases in the literature, relating to *Leishmania*-host  
401 interaction, where this has proven to be an issue (49, 53, 54). In this work, we addressed  
402 this by creating leishmaniasis-context pathways in the Reactome database. We created four  
403 subpathways, labelled as: *Leishmania* phagocytosis, killing mechanisms, cell recruitment  
404 (pro-inflammatory response), and *Leishmania* parasite growth and survival. Inside each, we  
405 labeled the pathways according to the receptor or membrane protein directing the signaling  
406 cascade. This structure facilitates the generation of high-level mechanistic insights from  
407 low-level processes highlighted by ORA, which are sensitive to particular experimental  
408 contexts (e.g. the source of the biological sample the data was derived from).

409 Manual literature search is often required for in depth interpretation of the outcome of ORA  
410 for extracting biological/functional insights. However, this strategy is time consuming,  
411 prone to omissions, and certainly incompatible with unbiased exploration of novel  
412 mechanisms/functions in the data, which is the goal of –omic approaches. This is because  
413 the search is necessarily constrained to cover a small number of topics within the  
414 researcher's area of expertise, and it is unachievable to manually trawl the entirety of  
415 biological literature relating to a particular gene, microorganism or disease.

416 We have re-analyzed two previously published transcriptomic datasets (from *Leishmania*-  
417 infected macrophages), to provide proof-of-concept of the usefulness of context-dependent  
418 databases such as the one described in this study. Results from this secondary analysis  
419 revealed the putative participation of a signaling pathway (ADORA2B mediated anti-  
420 inflammatory cytokines production) in the parasite-mediated induction of anti-  
421 inflammatory molecules.

422 Although *Leishmania* infection pathways in its first version is far from representing the full  
423 current knowledge about the interaction between the parasite and the host, we have shown  
424 that our database curation has led to new mechanistic insights from existing datasets.  
425 Moreover, these findings are generated from the use of a workflow that skips particular  
426 time-consuming, problematic manual curation steps, aligning the available data  
427 interpretation tools to the nature of unbiased hypothesis generation from –omics datasets.

428 **Materials and Methods**

429 ***Leishmania* Infection Curation Process**

430 The abstraction of *Leishmania* infection into a structure of pathways and reactions fulfilling  
431 the Reactome paradigm was accomplished by the Reactome working group. This consisted  
432 of a consortium of biocurators, software developers and leishmaniasis researchers. The  
433 latter selected the mechanisms of interest. The selection was based on biological pathways  
434 known to be associated with the infectious outcome, directly or indirectly. From here,  
435 domain experts and Reactome curators worked side by side to translate the selected  
436 pathways into the Reactome data structure using the curator tool version 3.3.

437 Reactome represents, categorizes, and annotates all known entities/molecules in each  
438 reaction. Different components can interact with each other only in ways prescribed by the  
439 Reactome data-model. For instance, the representation of an individual protein in the  
440 database requires several steps. In an example, for the protein ADAM17, the UniProt ID  
441 (P78536) is retrieved, as well as the GO cellular compartment it functions, and which  
442 species this protein belongs to (e.g., *Homo sapiens*). Note that if the represented process  
443 implies the transition of the same protein from one compartment to another, different  
444 instances of this protein must be created (e.g ADAM17 [endoplasmic reticulum], ADAM17  
445 [golgi apparatus] and ADAM17 [plasma membrane]). However, all these instances still  
446 point to the same UniProt ID. On the other hand, if several proteins individually fulfil the  
447 same role in a reaction (e.g., phosphorylation), these are grouped together into a single  
448 entity called “*defined set*” (PKL2 [cytosol] and MAPK14 [cytosol] in the *defined set* PKL2,  
449 MAP14 [cytosol] that phosphorylates ADAM17 [plasma membrane]). The *defined set*  
450 entity is also specific to each cellular compartment. Therefore, each individual molecule  
451 within the *defined set* must have a compartment-specific representation. Otherwise, in the  
452 quality assurance procedure, this will come out as an error. Molecules of the same or  
453 different types can form complexes, for example ADAM17:Zn<sup>2+</sup> which represents the  
454 functional form of ADAM17. Once these entities are created, the reactions in which they  
455 take part, are created. Binding is a common type of reaction in Reactome. Here, the output  
456 is a complex type entity (e.g., the interaction between sCD163 and MYH9 is represented as  
457 a binding that ends in the formation of the sCD163:MYH9 complex). If we are not at the  
458 final point of the pathway or dealing with a secondary product, the complex can participate

459 in subsequent reactions. In that case, the reaction from which this complex came from is  
460 indicated as the “preceding event” of the subsequent reaction.

461 Overall, we curated *Leishmania* infection pathways following the general structure of a  
462 signaling pathway. Namely, a ligand binding to a receptor, then the stimulated receptor  
463 effecting a downstream signaling cascade, up to the activation of effector molecules (eg.  
464 cytokines, nitric oxide, etc.). However, other pathways were conceived with different  
465 starting points. For instance, the first step of the CD163 example pathway consisted of  
466 ADAM17 activation. This begins in the endoplasmic reticulum, and its maturation process  
467 follows through the Golgi apparatus until its translocation to the plasma membrane, where  
468 its phosphorylation by kinases (either PLK2 or MAPK14), activates the cleavage reaction  
469 of CD163. It is the soluble portion of CD163, sCD163, that has been found as a regulator of  
470 the inflammatory responses in *Leishmania* infection, through the inhibition of the  
471 proliferation of T lymphocytes.

472 Reactome’s criteria for the acceptance of a particular molecular interaction and its  
473 supporting reference have been previously explained (3, 55). Once the reactions were  
474 successfully integrated into Reactome’s central database, we reached out to an experienced  
475 researcher in leishmaniasis, Dr. David Gregory (ORCID: 0000-0001-6534-7150) (52, 56,  
476 57) to review the curated material on the basis of his expertise in the field (52, 57–59).  
477 Only material reviewed by an independent domain expert is allowed to be published in a  
478 Reactome release. The contributions of authors and reviewers can be directly accessed  
479 through Reactome’s search interface, for example  
480 <https://reactome.org/content/query?q=David+Gregory> shows a detailed description of credit  
481 attribution in Reactome (60). We are highly interested in further contributions from domain  
482 experts for the completion of *Leishmania* infection pathways.

483 Once an orderly list of reactions was created, these were transformed according to the  
484 Reactome paradigm by using the curator tool. Some reactions were created from scratch to  
485 populate a low-level pathway, while others that were present within other pathways were  
486 duplicated (or reused) in the context of *Leishmania* infection (table 1). Within a reaction,  
487 each component was re-used if it already existed as part of a different process in the

488 database, otherwise it was created. Reuse of existing components and reactions ensures no  
489 redundancy in the database.

490 Reaction details, such as input/catalyst/output molecules, reaction type, preceding reactions  
491 and experimental species, were captured during the curation process. Reactions were linked  
492 together based on preceding-following relationships to generate pathways.

493

#### 494 **Funding**

495 This work was supported by the UK Global Challenges Research Fund CABANA grant  
496 BB/P027849/1, Wellcome Trust grant 107595/Z/15/Z to MAG, and US National Institutes  
497 of Health grant U41HG003751 (Reactome). JM was also funded by the Colombian  
498 National “Departamento Administrativo de Ciencia, Tecnología e Innovación”  
499 (COLCIENCIAS), Ph.D program, number 756-2016.

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692

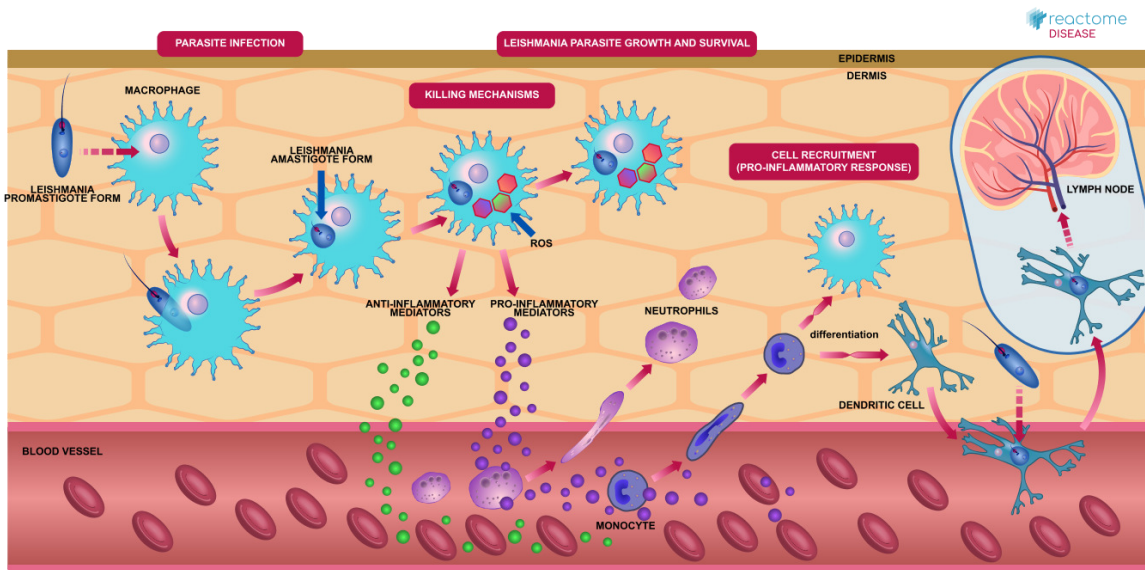
693 **Supplementary material description**

694 S1: It is an excel file with two tabs. The first one contains the gene input from the paper by  
695 Dillon *et al.* (49). The second one contains the output of applying ORA to that gene list.

696 S2: It is an excel file with two tabs. The first one contains the gene input from the paper by  
697 Gregory *et al.* (52). The second one contains the output of applying ORA to that gene list.

698

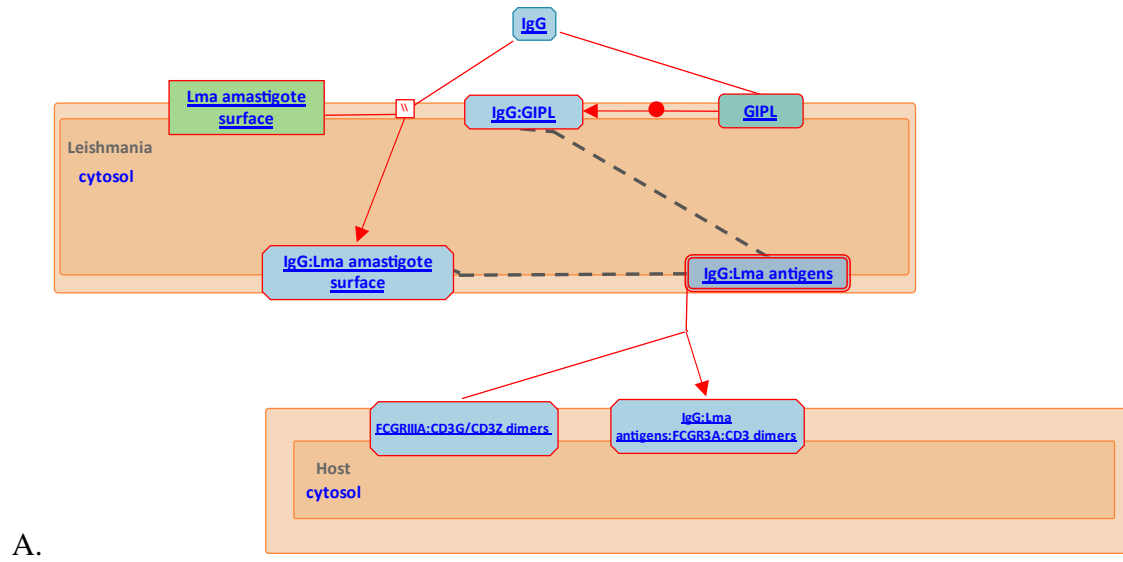
699



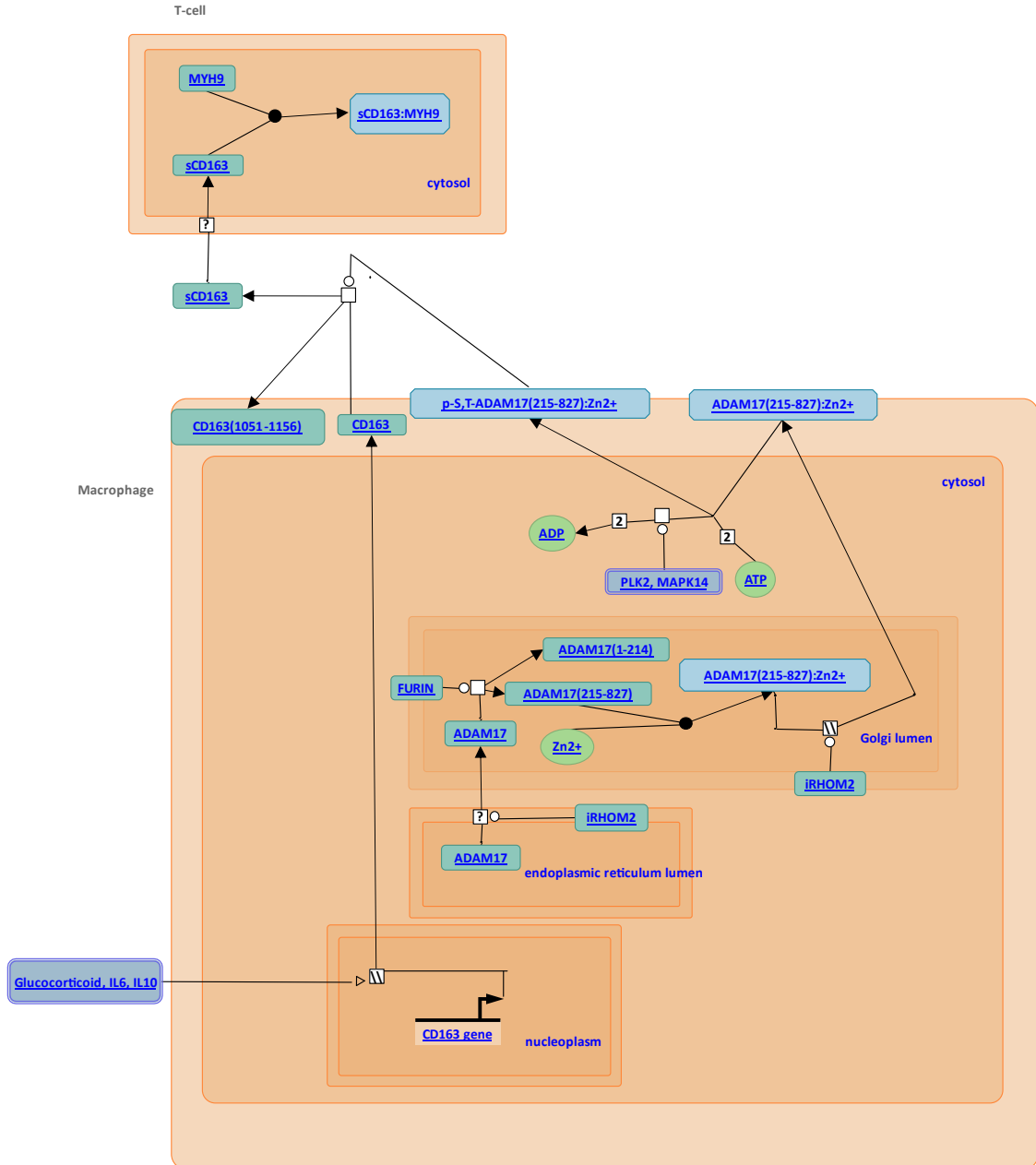
**Figure 1: Textbook-style diagram representing the top-level pathway “*Leishmania* infection”.** The major steps occurring in the dermis were compartmentalized into four categories: phagocytosis, killing mechanisms, cell recruitment, and *Leishmania* parasite growth and survival. On the webpage (<https://reactome.org/PathwayBrowser/#/R-HSA-9658195>), the magenta rectangular labels are interactive and take the user to the content of each subpathway.

- ☒ **Leishmania infection**
- ☒ **Parasite infection**
  - ☒ **Leishmania phagocytosis**
    - ⊕ **FCGR3A-mediated phagocytosis**
- ☒ **Killing mechanisms**
  - ⊕ **WNT5:FZD7-mediated leishmania damping**
  - ⊕ **Cell recruitment (pro-inflammatory response)**
    - ⊕ **Purinergic signaling in leishmaniasis infection**
- ☒ **Leishmania parasite growth and survival**
  - ⊕ **Anti-inflammatory response favouring Leishmania parasite infection**
    - ⊕ **CD163 mediating an anti-inflammatory response**
    - ⊕ **FCGR3A-mediated IL10 synthesis**
    - ⊕ **ADORA2B mediated anti-inflammatory cytokines production**
    - ⊕ **LTC4-CYSLTR mediated IL4 production**

**Figure 2:** Hierarchical structure of the *Leishmania* infection pathway showing the four sub-pathways and their contents. Each indentation introduces a new subpathway; as an example, “parasite infection” is the parent pathway for the subpathway “*Leishmania* phagocytosis”, which, to date only contains FCGR3A-mediated phagocytosis.



A.

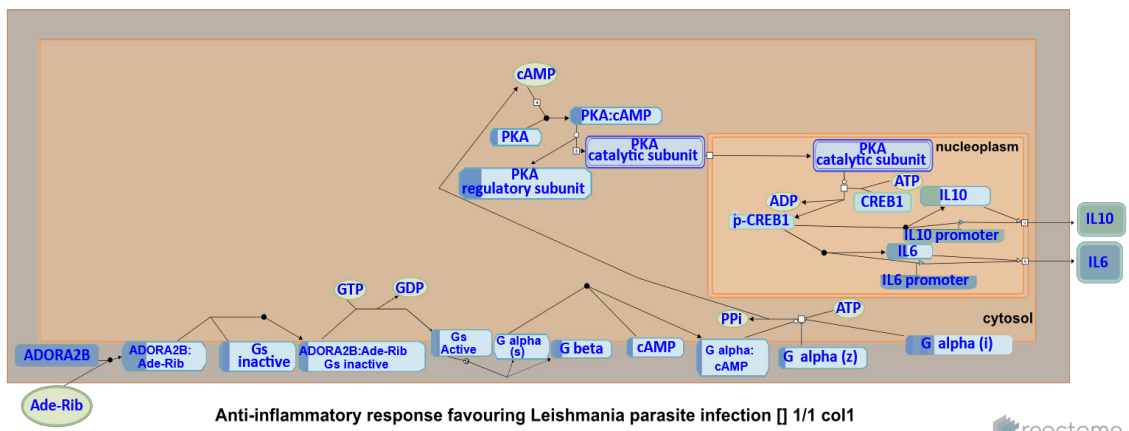


**Figure 3:** Standard graphical representation of pathways in Reactome. **A.** Fragment of the diagram for FCGR3A-mediated phagocytosis. It shows the reactions corresponding to the parasite opsonization process by the IgG. Parasitic components are highlighted in red. **B.** CD163 mediating an anti-inflammatory response. Each diagram shows the participating entities in the granularity of chemical compounds (green ovals), proteins (green rectangles), complexes (blue rectangles) and sets (blue rectangles with a double border). The arrangement

of the entities in the reactions can be easily followed on the web page, by clicking on the arrows that connect adjacent steps.

-   **Leishmania infection (45/403) FDR: 9.97E-1**
  -   **Parasite infection (9/157) FDR: 10E-1**
    -   **Leishmania phagocytosis (9/157) FDR: 10E-1**
      - ⊕   **FCGR3A-mediated phagocytosis (9/157) FDR: 10E-1**
  - ⊕   **Killing mechanisms**
    -   **Cell recruitment (pro-inflammatory response) (6/48) FDR: 7.9E-1**
      - ⊕   **Purinergic signaling in leishmaniasis infection (6/48) FDR: 7.9E-1**
  -   **Leishmania parasite growth and survival (34/297) FDR: 9.87E-1**
    - ⊕   **Anti-inflammatory response favouring Leishmania parasite infection (34/297) FDR: 9.87E-1**
      - ⊕   **CD163 mediating an anti-inflammatory response (4/14) FDR: 7.72E-1**
      - ⊕   **FCGR3A-mediated IL10 synthesis (12/141) FDR: 9.95E-1**
      - ⊕   **ADORA2B mediated anti-inflammatory cytokines production (20/159) FDR: 8.98E-1**
      - ⊕   **LTC4-CYSLTR mediated IL4 production (4/14) FDR: 7.72E-1**

A.



B.

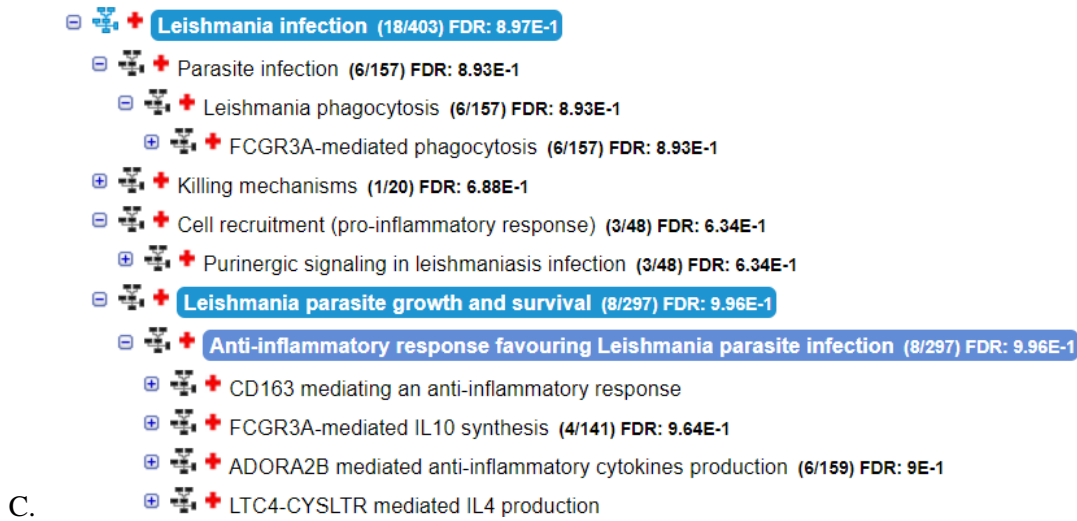


Figure 4: Results of applying ORA in Reactome, on the datasets generated by Dillon *et al.* and Gregory *et al.* A. Number of genes found in each subpathway from Dillon's dataset, with the associated false discovery rate (FDR) B. ADORA2B pathway enriched in Dillon's dataset shows upregulated genes involved in ADORA2B signaling cascade leading to the production of IL6 and IL10. C. Number of genes found in each subpathway from Gregory's dataset, and associated FDR.