Identification of novel proteins and mRNAs differentially bound to the *Leishmania* Poly(A) Binding Proteins reveals a direct association between PABP1, the RNA-binding protein RBP23 and mRNAs encoding ribosomal proteins

Ludmila A. Assis¹,², Moezio V. C. Santos Filho¹,², João R. da Cruz Silva¹, Maria J. R. Bezerra¹, Irassandra R. P. U. C. de Aquino¹, Klesion C. Merlo³, Fabiola B. Holetz³, Christian M. Probst³, Antonio M. Rezende¹, Barbara Papadopoulou⁴, Tamara D. C. da Costa Lima⁵, Osvaldo P. de Melo Neto¹*¹

¹ Department of Microbiology, Aggeu Magalhães Institute, Oswaldo Cruz Foundation, Recife, Pernambuco, Brazil, ² Department of Genetics, Federal University of Pernambuco, Recife, Pernambuco, Brazil, ³ Laboratory of Gene Expression Regulation, Carlos Chagas Institute, Oswaldo Cruz Foundation, Curitiba, Paraná, Brazil, ⁴ CHU de Quebec Research Center and Department of Microbiology-Infectious Disease and Immunology, Laval University, Quebec, Quebec, Canada, ⁵ University Center Tabosa de Almeida, Caruaru, Pernambuco, Brazil

* osvaldo.pompilio@cpqam.fiocruz.br

Abstract

Poly(A) Binding Proteins (PABPs) are major eukaryotic RNA-binding proteins (RBPs) with multiple roles associated with mRNA stability and translation and characterized mainly from multicellular organisms and yeasts. A variable number of PABP homologues are seen in different organisms however the biological reasons for multiple PABPs are generally not well understood. In the unicellular *Leishmania*, dependent on post-transcriptional mechanisms for the control of its gene expression, three distinct PABPs are found, with yet undefined functional distinctions. Here, using RNA-immunoprecipitation sequencing analysis we show that the *Leishmania* PABP1 preferentially associates with mRNAs encoding ribosomal proteins, while PABP2 and PABP3 bind to an overlapping set of mRNAs distinct to those enriched in PABP1. Immunoprecipitation studies combined to mass-spectrometry analysis identified RBPs differentially associated with PABP1 or PABP2, including RBP23 and DRBD2, respectively, that were investigated further. Both RBP23 and DRBD2 bind directly to the three PABPs in vitro, but reciprocal experiments confirmed preferential co-immunoprecipitation of PABP1, as well as the EIF4E4/EIF4G3 based translation initiation complex, with RBP23. Other RBP23 binding partners also imply a direct role in translation. DRBD2, in contrast, co-immunoprecipitated with PABP2, PABP3 and with RBPs unrelated to translation. Over 90% of the RBP23-bound mRNAs code for ribosomal proteins, mainly absent from the transcripts co-precipitated with DRBD2. These experiments suggest a novel and specific route for translation of the ribosomal protein mRNAs, mediated by RBP23, PABP1.
and the associated EIF4E/EIF4G3 complex. They also highlight the unique roles that different PABP homologues may have in eukaryotic cells associated with mRNA translation.

Author summary
Flagellated protozoa belonging to the Leishmania genus are the causative agents of the Leishmaniasis, neglected tropical diseases with a worldwide distribution. These parasites are well known for unique molecular mechanisms which include a lack of regulation of mRNA synthesis, generally the major stage for the control of gene expression in eukaryotic cells. Regulation of mRNA translation is therefore critical for survival and adaptation during a life cycle involving multiple life forms. Such regulation is likely dependent on RNA binding proteins (RBPs), such as the Poly-A Binding Protein (PABP) that binds to the 3' ends of the mRNAs. Three PABPs are found in Leishmania, more than in other unicellular organisms, with presumably distinct but undefined roles. Here we identify different mRNAs targets and specific RBPs with whom they interact. We define a specific association between one of the Leishmania PABPs, a single RBP and mRNAs encoding ribosomal proteins, an important subset of cellular mRNAs. Our results shed new light on the interactions required for the translation of these mRNAs, highlighting the relevance of regulating their translation in different organisms and the convergence of regulatory mechanisms acting through their 5' and 3' ends. The data also expands on the known PABP roles.

Introduction
The trypanosomatid protozoa constitute a group of parasitic microorganisms which include several species pathogenic to humans, all belonging to the Leishmania and Trypanosoma genera [1]. These are early divergent eukaryotes characterized by a lack of well-defined RNA polymerase II promoters and constitutive polycistrionic transcription. Expression of most of the trypanosomatid genes is regulated post-transcriptionally and it is assumed that this regulation mainly targets mechanisms associated with the processing, transport, stability and translation of mature mRNAs [2–5]. Thus, trypanosomatids emerge as relevant models for the understanding of eukaryotic mechanisms mediating post-transcriptional regulation.

In eukaryotes, a large number of RNA binding proteins (RBPs) recognize and bind specifically to sequences in target mRNAs and are required for their processing, stability, subcellular transport, storage, and translation, with many of these RBPs being specifically associated with regulatory motifs within the untranslated regions (UTRs) of the mRNAs [6–13]. Trypanosomatids have an unusually large number of RBPs belonging to distinct functional families and with different mRNA binding domains. These include those with the RRM (RNA Recognition Motif), ZF (Zinc Finger), PUF (Pumilio) and ALBA (Acetylation Lowers Binding Affinity) domains [14–18]. Many reports have identified differentially regulated RBPs acting as post-transcriptional regulators of gene expression in trypanosomatids, usually binding to sequence elements located within the 3' UTRs of mature transcripts [5,17–21]. Different mRNAs containing the same motifs and coding for functionally related proteins appear to be similarly regulated [22,23], but detailed mechanisms are generally not well defined.

The cytoplasmic poly(A)-binding proteins, PABPCs or simply PABPs (distinct from the NPABPs, nuclear and more functionally restricted), are the best known and the most abundant
of the eukaryotic RBPs. They participate in most, if not all, mRNA associated events, a consequence of their high affinity to the 3' end adenosine tract, the poly(A) tail, found in almost all eukaryotic mRNAs [24–27]. The PABPs interact with many other proteins, thus participating in various cellular events, with one of the best defined being translation. Within the cytoplasm, PABPs bound to the poly(A) tail interact with the translation initiation complex eIF4F bound to the cap structure of mature mRNAs, thus contributing to the formation of a closed loop structure that might enhance translation or promote translation re-initiation. eIF4F is a heterotrimeric complex of three subunits (eIF4E, eIF4G and the RNA helicase eIF4A) that facilitates the recruitment of the 40S ribosomal subunit to the mRNA. Its eIF4E subunit allows binding to the capped 5' end of the mRNAs, while a direct interaction between eIF4G and PABP is likely to mediate many of the PABP roles in translation [25,28]. A variable number of PABP homologues can be found in different organisms, some of which may be simultaneously expressed [25,27,29], however the mechanisms by which different homologues can be specifically recruited to distinct sets of mRNAs, and their functional implications, still needs to be better defined.

Three PABP homologues (PABP1, PABP2 and PABP3) were identified in *Leishmania* and other trypanosomatid species with PABP3 found to be lost from the *Trypanosoma* lineage. *Leishmania* PABP2 and PABP3 co-precipitate together and both can migrate to the nucleus after transcription inhibition, while PABP1 remains predominantly in the cytoplasm [30,31]. Similarly, the *Trypanosoma brucei* PABP2, but not PABP1, can also accumulate in the nucleus upon stress conditions [31], reinforcing a nuclear role for PABP2, and PABP2/PABP3, in *Trypanosoma* and *Leishmania* species, respectively. Recent proteomic analysis showed that PABP2 interacts with a wider range of protein partners, while PABP1 co-precipitates with a smaller and distinct set of polypeptides [32,33]. In trypanosomatids, multiple eIF4F complexes have been identified based on several distinct homologues of their eIF4E (six) and eIF4G (five) subunits. The EIF4G3/EIF4E4 based complex, which requires EIF4AI, is so far the only one whose role in translation is better understood. This complex specifically co-immunoprecipitates with PABP1, with a novel and direct interaction between PABP1 and EIF4E4 having been identified [3,34–36]. The PABP1/EIF4E4 interaction is likely to be associated with a common mechanism of regulation, since both proteins were seen to be simultaneously phosphorylated in multiple serine/threonine residues, an event which in *T. brucei* was found to be mediated by at least one cell-cycle regulated protein kinase [33,36–38].

The evidence presented so far implies that the *Leishmania/Trypanosoma* PABP1 binds to mRNA populations that are distinct than those bound by PABP2 (or PABP2/PABP3 in *Leishmania*), even though both PABPs are abundant proteins that are simultaneously expressed [30–32]. Although there is still no clear explanation for this differential mRNA selection, it is likely to be mediated by RBPs interacting with specific protein partners and mRNA targets. Here, we carried out a search for RBPs specifically associated with each of the *Leishmania infantum* PABP homologues and we selected RBP23 and DRBD2, which co-precipitated with PABP1 or PABP2/PABP3, respectively, for further characterization. Reciprocal co-immunoprecipitation and mass-spectrometry analysis confirmed a strong association between RBP23 and PABP1, as well as with the EIF4E4/EIF4G3 complex, while DRBD2 was found to be more associated with PABP2 and a larger number of different protein partners. RNA immunoprecipitation of individual proteins followed by large-scale RNA-sequencing revealed specific mRNA populations co-precipitating with each PABP homologue as well as RBP23 and DRBD2. Noteworthy is the largely specific co-precipitation of mRNAs encoding ribosomal proteins with both PABP1 and RBP23, with these mRNAs mostly absent from those found with DRBD2 or the two other PABPs. The data presented in this study suggest a new mechanism for the selection of mRNAs encoding ribosomal proteins during translation initiation in
trypanosomatids, with those mRNAs being targeted by both PABP1, as part of a larger repertoire of mRNAs, and more specifically by RBP23.

Results

Native Leishmania major PABPs co-immunoprecipitated distinct sets of mRNAs

Immunoprecipitation (IP) assays have previously shown that Leishmania PABP1 does not co-precipitate with either PABP2 or PABP3. In contrast, reciprocal assays have confirmed that PABP2 and PABP3 co-precipitate together [30]. Since multiple PABP molecules are expected to bind to a single mRNA molecule, lack of co-precipitation of PABP1 with the PABP2/PABP3 pair might reflect binding to different mRNA targets. In T. brucei, lacking PABP3, the PABP1 and PABP2 orthologues differ in their pattern of migration to starvation-stress granules, known to be associated with the bulk of cellular mRNAs. PABP1 and known protein partners are mainly absent from these granules which nevertheless are enriched with PABP2 and associated proteins and this has been interpreted as indicating that while PABP2 binds to most parasite transcripts, PABP1 is most likely associated with a subpopulation of mRNAs [32]. Here, to better define the functional distinctions between the Leishmania PABPs, we first investigated their mRNAs targets in Leishmania major using purified polyclonal sera generated against the three native paralogues. Following IP assays, co-immunoprecipitated mRNAs were extracted and sequenced by SOLiD. Transcripts with the highest number of reads and log2 > 2 enriched, when compared with the negative control (beads plus extracts but with the sera omitted) (S1 Table), were grouped according to a list of GO functional terms. These were modified as deemed fit to illustrate relevant differences between the proteins. mRNA targets co-precipitated with the three PABPs were generally restricted to transcripts encoding soluble proteins either known to be abundant or encoded by multiple genes. A total of 141 mRNAs were found to be enriched with PABP1, while PABP2 and PABP3 co-precipitated with 142 and 125 enriched transcripts, respectively. Although several putative targets were shared between the three L. major PABPs, the analysis revealed relevant differences in the percentages of mRNAs assigned to different GO categories (S1 Fig). A greater percentage of mRNAs encoding ribosomal proteins, grouped within the term "structural constituent—ribosome", were found associated with PABP1 (~60% vs. ~32% and ~31% for PABP2 and PABP3, respectively). In contrast, both PABP2 and PABP3 have a higher proportion of mRNAs encoding proteins with either binding function (~28% and ~30% of the mRNAs for PABP2 and PABP3 vs. ~18% for PABP1) or catalytic (~17% and ~13% of the mRNAs vs. ~6% for PABP1) and transporter (~6% and 8% of the mRNAs vs. ~4% for PABP1) activities. The differences in bound mRNAs between PABP1 and PABP2/PABP3 is even more noticeable when only the top-most, log2 > 4 enriched, messages are considered (S1 Fig). Although 60% of the top-most enriched mRNAs with PABP1 encode ribosomal proteins, these transcripts represent only 22% and 10% of those co-precipitated with PABP2 and PABP3. As a comparison, histone mRNAs represent 15% of the top-most mRNAs with PABP1 while these same messages comprise 39% and 52% of the most enriched transcripts found with PABP2 and PABP3. These results are in agreement with PABP1 binding to distinct mRNAs from those associated with PABP2 or PABP3 and having a marked preference for mRNAs encoding ribosomal proteins, with the latter two proteins having much more similar mRNA association patterns.
mRNAs co-immunoprecipitated with *L. infantum* HA-tagged PABPs

The polyclonal nature of the antibodies used for the set of IPs carried out for the native proteins might be associated with some degree of cross-reactivity and this might be one of the reasons for the overlap seen in mRNAs bound by PABP1 and the PABP2/PABP3 pair. We thus opted to carry out a more refined and independent experiment for the analysis of PABP-bound mRNAs using a previously described *L. infantum* cell line expressing PABP1 [33] and newly generated cell lines expressing either PABP2 or PABP3, all having an identical C-terminal HA epitope tag. When compared with the ectopic PABP1-HA, represented by isoforms indicative of phosphorylation events, both HA-tagged PABP2 and PABP3 were visualized as single bands in whole cellular extracts derived from the transfected cell lines (S2 Fig). Cytoplasmic extracts of these cell lines were then prepared after lysis through cavitation, in the absence of any detergent, and then used in IPs performed with monoclonal antibodies immobilized on magnetic beads and directed to the HA tag, therefore avoiding cross-reacting events. As negative controls, parallel IPs were performed using cytoplasmic extracts made from cells having no HA-tagged protein. Co-purified mRNAs were extracted from the IPs and used for Illumina sequencing. The new approach uses different lysis method, antibody and immunoprecipitation/sequencing strategies.

The full set of mRNAs identified with the new IP approach are listed in the S2 Table. A first analysis using a Volcano plot confirms a much-reduced overlap in mRNAs bound by PABP2 or PABP3 vs. PABP1. In contrast, no significant differences between the mRNAs associated with PABP2 or PABP3 can be detected (S3 Fig). A total of 108 mRNAs were found to co-precipitate with PABP1, fulfilling the criteria of log₂ fold enrichment greater than two (Fig 1). Forty-five of the enriched transcripts (~42%) are mRNAs encoding ribosomal proteins, 10 (~9%) encode enzymes with “catalytic activity” and 26 (24%) encode proteins classified with the term “binding function”. PABP2 and PABP3 co-precipitated a total of 135 and 118 transcripts, respectively. For both PABPs, mRNAs encoding ribosomal proteins represented only ~2% of the co-precipitated transcripts, while 24% to 26% of those encode proteins with “catalytic activity” and roughly 30% encode proteins classified with “binding function”. Other GO functional categories were generally poorly represented among the mRNAs found associated with the three proteins. Noteworthy however, is the increased presence of histone mRNAs associated with PABP3 (six different transcripts—~5%) in comparison with both PABP2 (two transcripts—1.5%) and PABP1 (one transcript—<1%), although the mRNAs co-precipitated with the HA-tagged *L. infantum* PABPs are not as enriched with histone transcripts as those co-precipitated with their native *L. major* orthologues (see S1 Table).

To better define the mRNAs specifically bound by individual HA-tagged PABPs and also to identify those commonly bound by distinct PABP pairs, we did a Venn diagram analysis of the transcripts co-precipitated with the three HA-tagged *L. infantum* PABPs (S4 Fig). A total of 66 mRNAs bound exclusively to PABP1, with 43 of those encoding ribosomal proteins. Meanwhile, 59 mRNAs bound exclusively to PABP2 and 37 to PABP3. PABP2 was associated with a larger number of mRNAs encoding hypothetical or uncharacterized proteins while the histone mRNAs were preferentially associated to PABP3. As expected, PABP2 and PABP3 shared more mRNAs (45 total) than PABP1 and PABP2 (eight) or PABP1 and PABP3 (eleven). Twenty-three mRNAs, however, co-precipitated with all three proteins. Overall, the experiment with the HA-tagged proteins proved to be more specific than the one with the native proteins, with PABP1 having a clear preference for mRNAs encoding ribosomal proteins and those being noticeably absent from the transcripts associated with PABP2 or PABP3.
Leishmania PABPs co-immunoprecipitate and interact with distinct RNA-binding proteins

Aside from their strong specificity for poly(A) sequences, PABPs have no known sequence specific affinity which could justify their selective binding to different sets of mRNAs. Binding to poly(A) has been shown to be mediated by specific residues within the PABPs’ RRMs 1 and 2, which are generally conserved in Leishmania PABP1 and PABP3, whereas substitutions in these residues have been identified for PABP2 which might lead to changes in sequence binding specificity [30]. For Leishmania PABP1 at least, and most likely for the PABP2/PABP3 pair as well, their recognition of specific mRNAs targets might require the assistance of partner RBPs with distinct RNA binding specificities. Here, putative RNA binding proteins that might be involved in specific mRNA recognition were initially identified in a pilot mass spectrometry analysis of the proteins co-immunoprecipitated with the three HA-tagged L. infantum PABPs (S5 Fig). These experiments led to the identification of RNA-binding proteins specifically

---

**Fig 1.** Analysis of mRNA populations associated with the three HA-tagged *Leishmania infantum* PABPs. mRNA groups associated with the recombinant HA-tagged PABP1, PABP2 and PABP3 from *L. infantum* (Illumina sequencing). Enriched genes in at least two of three available RNA-seq datasets were manually classified and grouped using the gene ontology (GO) terms according to their molecular function. ‘Enriched’ means at least 2-fold more abundant than in the negative control.

https://doi.org/10.1371/journal.pntd.0009899.g001

*Leishmania* PABPs co-immunoprecipitate and interact with distinct RNA-binding proteins

Aside from their strong specificity for poly(A) sequences, PABPs have no known sequence specific affinity which could justify their selective binding to different sets of mRNAs. Binding to poly(A) has been shown to be mediated by specific residues within the PABPs’ RRMs 1 and 2, which are generally conserved in *Leishmania* PABP1 and PABP3, whereas substitutions in these residues have been identified for PABP2 which might lead to changes in sequence binding specificity [30]. For *Leishmania* PABP1 at least, and most likely for the PABP2/PABP3 pair as well, their recognition of specific mRNAs targets might require the assistance of partner RBPs with distinct RNA binding specificities. Here, putative RNA binding proteins that might be involved in specific mRNA recognition were initially identified in a pilot mass spectrometry analysis of the proteins co-immunoprecipitated with the three HA-tagged *L. infantum* PABPs (S5 Fig). These experiments led to the identification of RNA-binding proteins specifically
associated with PABP1 (RBP23 and the uncharacterized LINF_180008000), PABP2 (DRBD2) or the three proteins (ZC3H41). Here we opted to investigate further both RBP23, confirmed to be strictly associated with PABP1 in both Leishmania and T. brucei [32,33], and DRBD2. In T. brucei, RBP23 is a cytoplasmic protein with a reticulated distribution, while DRBD2 was also localized within the cytoplasm, but distributed in the perinuclear region [39]. Both are small proteins characterized by the presence of RRM domains at their extremities, with the C-terminal RBP23 RRM and the two DRBD2 domains already mapped when these two proteins were originally described [40]. A second atypical RRM-like domain is also identifiable for RBP23 using current secondary structure and domain prediction tools, mapped to its N-terminus. To first confirm if direct interactions occur between RBP23 or DRBD2 and the Leishmania PABP homologues, both RBP23 and DRBD2 genes were cloned and the corresponding proteins used for in vitro pull-down assays. The three PABPs were first expressed in *Escherichia coli* with an N-terminal Glutathione S-transferase (GST) tag. After immobilization in Glutathione Sepharose these were then incubated with 35S-labeled RBP23 or DRBD2, produced by in vitro transcription/translation. The two 35S-labeled proteins bound to all three GST-tagged PABPs, with no binding seen for the negative GST control (Fig 2). However, a stronger signal was observed for both labeled proteins with PABP1, presumably indicating a stronger interaction. These results confirm the ability of both RBP23 and DRBD2 to directly interact with the Leishmania PABPs, with their binding motifs likely being conserved within different PABP sequences. Nevertheless, the lack of specificity seen in vivo by the co-immunoprecipitation assays, not only between RBP23 and PABP1 but also between DRBD2 and PABP2, indicates that this specificity might require binding to their mRNA targets or other protein partners.

**RBP23 binding partners**

To fully investigate the RBP23 and DRBD2 interactions in vivo, both proteins were ectopically expressed in *L. infantum* with a C-terminal HA-tag. As for the PABPs, expression was confirmed with the anti-HA antibody in exponentially grown promastigotes, with both proteins migrating as single bands in agreement with predicted sizes (Fig 3A). No isoforms suggestive of post-translational modifications were seen. Cytoplasmic extracts were then generated, and preliminary IPs performed. The amount of RBP23 detected in the cytoplasmic extract and/or IPs was generally much lower than that observed for DRBD2 (Fig 3B), contrasting with equivalent levels of expression for both proteins in whole cell extracts. RBP23 is thus likely to be much more susceptible to degradation than DRBD2.

To confirm the interaction with PABP1 and to identify additional protein partners of functional relevance, proteins co-immunoprecipitated with the HA-tagged RBP23 were next submitted to mass-spectrometry. For these assays we used extracts that were not RNase treated so that both directly bound partners as well as proteins bound to the same mRNAs as RBP23 would be precipitated. Normalized intensities from two sets of replicates, and with a minimum of two peptides found for each replicate, were first used to generate a full list of proteins enriched 1.5-fold or more with RBP23 in comparison to the negative control (S3 Table). For a more detailed analysis, we first considered only co-precipitated proteins that were enriched by four-fold or more in comparison with the control. These were categorized, based on both the intensity values and the ratio of enrichment, according to the “strength” of association with RBP23 (Tables 1 and S4). Proteins classified within the top two categories, possibly reflecting more robust associations, are summarized in Fig 3C. PABP1 was found to be foremost among those, being the top-most protein in intensity and with a greater than 65-fold enrichment. In contrast, although PABP2 can be considered high ranking among the co-precipitated proteins listed in the S3 Table, due to its overall intensity value, it is only enriched between three-
four-fold, similarly to PABP3. EIF4G3 and EIF4E4 are also among the top-most proteins enriched with RBP23, as well as three other proteins previously shown to specifically co-precipitate with the \textit{L. infantum} HA-tagged PABP1 \cite{33}: the zinc-finger ZC3H41; the uncharacterized LINF\_180008000; and LINF\_050009500, another uncharacterized protein structurally similar to Skp1 (S-phase kinase associated protein 1).

Several proteins among those shortlisted in Table 1 and Fig 3C have specific functions associated with RNA metabolism. Noteworthy, are the CNOT10, CNOT11 and CAF1 subunits of the CCR4-NOT complex. Two proteins linked with mitochondrial mRNA metabolism, PPR and MRB1, were also found, as well as STRAP, an RNA-binding protein whose \textit{T. brucei} orthologue localizes to stress granules under starvation conditions \cite{41}. Uncharacterized co-precipitated proteins whose \textit{T. brucei} orthologues have both been shown to be associated with mRNA include LINF\_200021500 and LINF\_050009500 \cite{42}. The LINF\_200021500 orthologue was also found to enhance expression in a tethering assay, and seen to bind to MKT1, a known translational regulator \cite{43,44}. Another uncharacterized protein whose orthologue was also found to stimulate expression is LINF\_110013800. Considering that in \textit{T. brucei} the PABP1, EIF4E4, EIF4G3, EIF4A1 and LINF\_18008000 orthologues all also enhance expression, with some at least through translation stimulation, the strong association seen with these proteins reinforce a possible role for RBP23 during translation initiation.
DRBD2 binding partners

Results derived from a mass-spectrometry analysis of DRBD2-HA samples were analysed generally as described for RBP23, with the full list of enriched proteins shown in the S5 Table. Proteins enriched 4-fold or more and categorized as to the “strength” of association with DRBD2.
Table 1. Proteins specifically co-immunoprecipitating with *L. infantum* RBP23.

| TRITrypDB ID | Name | Description | Intensity | Ratio | Strength |
|--------------|------|-------------|-----------|-------|----------|
| Bait | | | | | |
| —LINF_170011800 | RBP23 | RNA-binding protein 23 | 7.6x10^8 | 291.1 | +++ |
| Translation initiation factors and partners | | | | | |
| —LINF_350055900 | PABP1 | Poly(A)-binding protein 1 | 1.4x10^9 | 66 | +++ |
| —LINF_160022100 | EIF4G3 | Eukaryotic translation initiation factor 4G3 | 8.3x10^7 | 25.4 | +++ |
| —LINF_300009600 | EIF4E | Eukaryotic translation initiation factor 4E | 7.5x10^7 | 129.8 | +++ |
| —LINF_340033000 | EIF3G | Eukaryotic translation initiation factor 3 G | 2.9x10^7 | 4.3 | ++ |
| —LINF_330010200 | 4E1-IP2 | EIF4E1-interacting protein 2 | 2.1x10^6 | 6.5 | - |
| —LINF_270023100 | EIF4E1 | Eukaryotic translation initiation factor 4E1 | 1.7x10^6 | 4.8 | - |
| RNA-binding proteins and related | | | | | |
| —LINF_270019700 | ZC3H41 | CCCH-Zinc finger 41 | 6.5x10^8 | 10.1 | +++ |
| LINF_270018900 | STRAP | Serine-threonine kinase receptor-associated protein | 5.2x10^7 | 6.1 | +++ |
| LINF_110015300 | MRB1AP | Mitochondrial-associated protein | 2x10^7 | 36.8 | +++ |
| LINF_300016700 | TRRM2 | Three RRMs containing protein 2 | 6.8x10^6 | 7.8 | - |
| LINF_310020800 | PuREBP1 | Purine-Responsive Element Binding Protein 1 | 1.8x10^7 | 13.7 | +++ |
| Polyadenylation and Deadenylation | | | | | |
| LINF_360070900 | CNOT10 | CCR4-NOT complex subunit 10 | 1.4x10^8 | 7.9 | +++ |
| LINF_230005100 | CNOT11 | CCR4-NOT complex subunit 11 | 6.5x10^7 | 7.3 | +++ |
| LINF_220022600 | CAF1 | CCR4-associated factor | 4x10^7 | 10.7 | +++ |
| LINF_240059000 | PAP | Poly A polymerase, regulatory subunit | 3.3x10^6 | 11.8 | - |
| LINF_100018400 | PBP1 | PAB1-binding protein | 2.3x10^6 | 5.6 | - |
| Enzymes | | | | | |
| LINF_180022300 | GSH1 | Gamma-glutamylcysteine synthetase | 5x10^6 | 11 | ++ |
| LINF_060014300 | DHFR-TS | Dihydrofolate reductase-thymidylate synthase | 3x10^6 | 5.9 | - |
| LINF_300036300 | tRNA (Guanine-1)-methyltransferase | 7.6x10^5 | 5.2 | - |
| Intracellular transport and cell motility | | | | | |
| LINF_290029700 | DNM1 | Dynamin-1-like protein | 3.8x10^7 | 5 | ++ |
| Uncharacterized proteins with identifiable domains | | | | | |
| —LINF_180008000 | Protein with NFT2 and RRM domains | 7.4x10^7 | 23.9 | +++ |
| —LINF_210020800 | PPR domain protein | 4.2x10^7 | 16.4 | +++ |
| —LINF_050009500 | Skp1-like | Skp1 (S-phase kinase-associated protein 1)—like | 3.5x10^7 | 10.6 | +++ |
| LINF_080013100 | BAR domain protein | 1.4x10^7 | 11.7 | ++ |
| LINF_260021000 | RING-zinc finger protein | 4.1x10^6 | 12.2 | ++ |
| Other selected uncharacterized proteins | | | | | |
| LINF_040006400 | | | 9x10^7 | 15.6 | +++ |
| LINF_110013800 | | | 3.4x10^7 | 4.1 | ++ |
| LINF_200021500 | | | 3.4x10^7 | 81.4 | +++ |
| LINF_190005100 | | | 2.9x10^7 | 10.3 | +++ |

The polypeptides listed are also detailed in the S3 Table and includes only identified proteins which fit into the following parameters: two or more peptides found for each RBP23 replicate; a minimum of 4-fold (4x) enrichment over the negative control (“Ratio”); a minimum average intensity (“Intensity”) 1000-fold (1000x) less than that seen for the RBP23 bait. The “Strength” column defines the strength of association with RBP23 taken into account both the average intensity and the enrichment in comparison with the negative control, according with criteria defined in the bottom of the S3 Table. Proteins co-immunoprecipitated with both *L. infantum* HA-tagged PABP1 [33] and RBP23 are indicated by arrows. Proteins also found co-precipitated with DRBD2 are underlined.

https://doi.org/10.1371/journal.pntd.0009899.t001
are listed in Table 2 and S6 Table and the top-most are also represented in Fig 4A. The data confirm that DRBD2 preferentially co-precipitates not only with PABP2 but also with PABP3. It furthers highlights the absence of PABP1, or most other proteins associated with RBP23, from the topmost co-precipitated proteins. Although RBP23 is found in the list from Table 2, it is one of the lowest ranking proteins included there. Only two proteins were found among the top candidates co-precipitated with both RBP23 and DRBD2: ZC3H41 and the uncharacterized LINF_190005100. Several RNA-binding proteins, not generally seen with RBP23, were found in the DRBD2 pull-down, contrasting with a reduced number of uncharacterized proteins. Considering only those from the top two categories, and omitting mitochondrial proteins, a total of four proteins with a zinc-finger domain (ZC3H31, ZCH39, ZCH40 and ZCH41), one RRM containing RBP (RBP43), a pumilio domain protein (PUF6), a putative RBP with NTF2 domain (LINF_210009700) and one DEAD-box RNA helicase (HEL67/DDX3/Ded1p) were found in our analyses.

A relevant observation from the DRBD2 data is the co-precipitation of 18 proteins with defined functions associated with mitochondrial RNA (Table 2), most of which showing substantial enrichments (greater than 15-fold). The mitochondrial-related proteins represent 38% of those found associated with DRBD2, as opposed to the putative RBP23 partners, of which only 2% belong to this category (Fig 4B). Orthologues to several of the proteins co-precipitated with the *Leishmania* DRBD2, but none of the mitochondrial proteins, were also found in a recent analysis of DRBD2 binding partners from *T. cruzi* [45]. Overall, the set of proteins found associated with DRBD2 highlights a markedly distinct profile in comparison with RBP23. It also suggests possible involvement in many different processes associated with the metabolism of targeted mRNAs but with minor, if any, role in translation.

### Comparative analysis of selected proteins co-immunoprecipitated with RBP23 and DRBD2

To better define the association of RBP23 or DRBD2 with different co-immunoprecipitated putative partners, we directly compared the enrichment ratios seen with either RBP23 or DRBD2 for several proteins known to be specifically associated with major RNA processes. First, as shown in Fig 5A, the comparison clearly highlights the association of RBP23 with PABP1, while DRBD2 preferentially interacts with PABP2 and also co-precipitates with PABP3. A differential association was also seen with a number of proteins and enzymes involved with different aspects of RNA metabolism (Fig 5B), such as various splicing and polyadenylation factors, some enriched only with RBP23 and others with DRBD2. In contrast, enrichments for over 60 ribosomal protein subunits and all 11 experimentally validated eIF3 subunits [47] were found with RBP23, but not with DRBD2. These results are consistent with RBP23 being associated with polysomal mRNAs, in contrast to DRBD2, and also in agreement with published tethered results showing that RBP23 enhances expression while DRBD2 has been shown to repress expression [42,43]. As previously, highlighted, another contrasting difference between the two proteins is the association of RBP23 with six subunits of the CCR4-Not complex, some with a very substantial enrichment, not seen for DRBD2.

Here we also sought to investigate in more detail the association of both RBP23 and DRBD2 with eIF4F subunits and related proteins. As well as the eIF4E4/eIF4G3 complex, eIF4A1, its known helicase partner in trypanosomatids, was differentially co-precipitated with RBP23 only (Fig 5C). eIF4E1, another eIF4E homologue, and one of its known protein partners, eIF4E1-IP2, was also found enriched with RBP23. In contrast, two eIF4G homologues, eIF4G1, with partners eIF4G1-IP and eIF4G1-IP2 (also called RBP43), and eIF4G2, were more enriched with DRBD2, with eIF4G3 being enriched not only with RBP23 but also with
Table 2. Proteins specifically co-immunoprecipitating with *L. infantum* DRBD2.

| TriTrypDB ID | Name       | Description                          | Intensity   | Ratio | Strength |
|--------------|------------|--------------------------------------|-------------|-------|----------|
| —LINF_350027200 | DRBD2     | dsRNA-binding domain 2               | 3.7x10^8    | 110.3 | +++      |
| —LINF_350047100 | PABP2     | Poly(A)-binding protein 2            | 7.4x10^8    | 12.4  | +++      |
| LINF_250005800 | PABP3     | Poly(A)-binding protein 3            | 1.3x10^8    | 13.9  | +++      |
| LINF_160022100 | EIF4G3    | Eukaryotic translation initiation factor 4G3 | 1.9x10^7    | 4.4   | -        |

### Translation initiation factors

| TriTrypDB ID | Name       | Description                          | Intensity   | Ratio | Strength |
|--------------|------------|--------------------------------------|-------------|-------|----------|
| —LINF_350027200 | DRBD2     | dsRNA-binding domain 2               | 3.7x10^8    | 110.3 | +++      |
| LINF_250005800 | PABP3     | Poly(A)-binding protein 3            | 1.3x10^8    | 13.9  | +++      |
| LINF_160022100 | EIF4G3    | Eukaryotic translation initiation factor 4G3 | 1.9x10^7    | 4.4   | -        |

### RNA-binding proteins

| TriTrypDB ID | Name       | Description                          | Intensity   | Ratio | Strength |
|--------------|------------|--------------------------------------|-------------|-------|----------|
| LINF_270019700 | ZC3H41    | CCCH-Zinc finger 41                  | 2.5x10^8    | 4.7   | +++      |
| LINF_210009700 | ZC3H39    | CCCH-Zinc finger 39                  | 1x10^6      | 63.3  | +++      |
| —LINF_190007900 | ZC3H40    | CCCH-Zinc finger 40                  | 8.8x10^7    | 44.4  | +++      |
| LINF_330019100 | PUF6      | Pumilio protein 6                    | 4.9x10^7    | 15.6  | ++       |
| LINF_360013000 | ZC3H31    | CCCH-Zinc finger 31                  | 4.2x10^7    | 11    | +        |
| LINF_250008000 | RBP43/G1-IP2 | RNA-binding protein 43               | 3.1x10^7    | 13.6  | ++       |
| LINF_180019090 | RBP12     | RNA-binding protein 12               | 2.7x10^7    | 7.1   | -        |
| LINF_270028100 | TRRM1     | Three RRM containing protein 1       | 1.5x10^7    | 4.8   | -        |
| LINF_300016700 | TRRM2     | Three RRM containing protein 2       | 1.3x10^7    | 4.7   | -        |
| LINF_320012700 | PAN3      | Poly(A) Specific Ribonuclease Subunit | 7.1x10^6    | 6     | -        |
| LINF_170011800 | RBP23     | RNA-binding protein 23               | 6.5x10^6    | 4.8   | -        |

### Mitochondrial RNA-binding proteins

| TriTrypDB ID | Name       | Description                          | Intensity   | Ratio | Strength |
|--------------|------------|--------------------------------------|-------------|-------|----------|
| —LINF_090018500 | KMRP2/MRP2 | Mitochondrial RNA binding protein 2  | 8.2x10^7    | 16.5  | +++      |
| LINF_180005000 | KPAF1     | Kinetoplast polynucleation factor 1  | 7.7x10^7    | 38.8  | +++      |
| LINF_270017500 | KMRP1/MRP1 | Mitochondrial RNA binding protein 1  | 3.8x10^7    | 11.7  | ++       |
| LINF_280005300 | MERS2     | Mitochondrial edited mRNA stability factor 2 | 3.5x10^7    | 34.5  | ++       |
| LINF_220011900 | RESC2/GAP2 | Guide RNA associated protein 2       | 3.2x10^7    | 44.3  | ++       |
| LINF_310012100 | RESC12/REMC5 | RNA editing mediator complex protein 5 | 2.1x10^7    | 14.2  | ++       |
| LINF_330036600 | RESC1/GAP1 | Guide RNA associated protein 1       | 1.9x10^7    | 22.4  | ++       |
| LINF_320031100 | MERS1     | Mitochondrial edited mRNA stability factor 1 | 1.7x10^7    | 100.8 | +        |
| LINF_080016600 | RESC6/GRBC6 | gRNA binding complex subunit 6       | 1.5x10^7    | 8.3   | -        |
| LINF_330007800 | RESC13/RGG2 | RGG-containing protein 2             | 1.4x10^7    | 26.1  | +        |
| LINF_310012000 | RESC11B/MRB4150 | Mitochondrial RNA binding complex 1 subunit | 1.4x10^7    | 12.9  | +        |
| LINF_200014400 | RESC15/PAMC1 | Polyadenylation mediator complex 1    | 9.9x10^6    | 11.8  | +        |
| LINF_360057200 | RESC8/REMC1 | RNA editing mediator complex protein 1 | 9.18x10^6   | 16.8  | +        |
| LINF_330020200 | RESC5/GRBC5 | gRNA binding complex subunit 5       | 8.8x10^6    | 15.5  | -        |
| LINF_300013200 | KRGG1/RGG1 | RGG-containing protein 1             | 6.2x10^6    | 11.3  | -        |
| LINF_210011600 | RESC17/PAMC3 | Polyadenylation mediator complex 3    | 5.8x10^6    | 8.3   | -        |
| LINF_250021900 | KRBP72/MRB1590 | Mitochondrial RNA binding complex 1 subunit | 4.6x10^6    | 4.2   | -        |
| LINF_360078000 | RESC19/MERS3 | Mitochondrial edited mRNA stability factor 3 | 3.8x10^6    | 32.7  | -        |

### RNA helicases

| TriTrypDB ID | Name       | Description                          | Intensity   | Ratio | Strength |
|--------------|------------|--------------------------------------|-------------|-------|----------|
| —LINF_320009100 | HEL67     | ATP-dependent RNA helicase            | 1.4x10^8    | 7.7   | +++      |
| LINF_070008800 | DBP2B     | ATP-dependent DEAD/H RNA helicase     | 3.6x10^7    | 4.9   | +        |
| LINF_050006300 | DDX21     | Nucleolar RNA helicase II             | 2.2x10^7    | 7     | -        |
| LINF_220021200 | ATP-dependent DEAD/H RNA helicase     | 4.6x10^6    | 4.2   | -        |

### RNA methyltransferases

| TriTrypDB ID | Name       | Description                          | Intensity   | Ratio | Strength |
|--------------|------------|--------------------------------------|-------------|-------|----------|
| LINF_110010500 | G1-IP     | mRNA cap guanine-N7 methyltransferase | 3.4x10^7    | 6.2   | -        |
| LINF_200021600 | tRNA (Uracil-5')-methyltransferase | 4.3x10^6    | 7.4   | -        |

### Other enzymes

(Continued)
DRBD2. Little or no enrichment was seen with either of the RBPs for EIF4E3, the most abundant of the trypanosomatid eIF4Es, and its EIF4G4 partner. Once again, these differences highlight the very distinct profile of protein partners co-precipitating with both RBPs.

Distinct mRNA populations co-immunoprecipitated with RBP23 and DRBD2

To define the putative mRNA targets bound by the either RBP23 or DRBD2 and to investigate overlaps and differences in comparison to their PABP partners, sequencing of RNA populations co-immunoprecipitated with HA-tagged RBP23 and DRBD2 was performed using the same procedures carried out with the HA-tagged PABPs. The full list of mRNAs specifically enriched with either RBP23 or DRBD2, according to the same criteria defined for the analysis with the mRNAs bound to the three PABPs, is found in the S7 Table. These mRNAs are also functionally grouped in Fig 6A. A total of 114 enriched mRNAs were co-precipitated with RBP23, of which the vast majority (>80%) encode known ribosomal proteins. RBP23 thus show a much greater specificity than that seen for either the native or HA-tagged PABP1. Among the remaining enriched transcripts, at least three are directed linked to ribosomal function: the translation factor EIF5A, the nuclear RNA binding protein NRBP and the nascent polypeptide associated complex subunit. Comparison of the mRNAs co-precipitated with the L. infantum PABP1 to those found associated with RBP23 revealed that ~90% of the ribosomal protein mRNAs found with PABP1 also co-precipitated with RBP23. In contrast, none of the mRNAs encoding uncharacterized proteins found with PABP1, the second most abundant category of mRNAs found with this protein, were among those co-precipitated with RBP23. A very interesting observation, however, is that the top-most mRNA enriched with RBP23 was its own transcript, not found among those associated with PABP1.

A remarkably different profile was observed for the mRNAs co-precipitated with DRBD2 (Fig 6B). From a total of 300 enriched transcripts, only a minor fraction of those (~2%) encode proteins with a structural activity role. Among the 14 transcripts that encode ribosomal protein mRNAs, half encode subunits of mitochondrial ribosomes, not found associated with uncharacterized proteins with other identifiable domains

Uncharacterized proteins

| TriTrypDB ID | Name   | Description                                      | Intensity      | Ratio | Strength |
|--------------|--------|--------------------------------------------------|----------------|-------|----------|
| LINF_050009500 | Skp1-like | 5'-phase kinase-associated protein 1—like        | 1.4x10⁷         | 4.4   | -        |
| LINF_190005100 |         |                                                  | 3.7x10⁷         | 31.4  | ++       |
| LINF_200005800 |         |                                                  | 6.2x10⁶         | 7.1   | -        |

The polypeptides listed here are detailed in S5 Table and include only identified proteins which fit into the following parameters: two or more peptides found for each DRBD2 replicate; a minimum of 4-fold (4x) enrichment over the negative control (“Ratio”); a minimum average intensity (“Intensity”) 1000-fold (1000x) less than that seen for the DRBD2 bait. The “Strength” column defines the strength of association with DRBD2 taking into account both the average intensity and the enrichment in comparison to the negative control, according to the criteria defined in the bottom of the S5 Table. Proteins co-purifying with both T. cruzi DRBD2 [45] and L. infantum HA-tagged DRBD2 are indicated by arrows. Proteins in common with RBP23 are underlined.

*include recently assigned names [46].

https://doi.org/10.1371/journal.pntd.0009899.t002
Fig 4. Top proteins co-purifying with HA-tagged DRBD2 and their subcellular compartmental localization compared to HA-tagged RBP23. (A) Bar chart with proteins co-precipitated with DRBD2 with the enrichment strength defined as ‘+++’ (orange), ‘++’ (green) and ‘+’ (blue), according to Table 2. Proteins co-purifying with both *T. cruzi* DRBD2 [45] and its *L. infantum* HA-tagged orthologue are indicated by arrows. Proteins in common with RBP23 are underlined. (B) Pie charts with subcellular localization of DRBD2 (left chart) or RBP23 (right chart) partners. This was defined based on cellular component classes: cytoskeleton; cytoplasm; mitochondrion; nuclear; organelles/granules (except mitochondria); n/d or not defined. For uncharacterized proteins, their localization was defined based on their respective orthologues from *T. brucei* and according to TrypTag.org.

https://doi.org/10.1371/journal.pntd.0009899.g004
RBP23 (classified under mitochondrial in Fig 6B). For the remaining transcripts, most can be classified into three categories: ~31% encode binding proteins, ~26% encode proteins with uncharacterized functions and ~21% encode enzymes with catalytic activity. When compared with the PABP2 or PABP3 co-precipitated transcripts, a very limited overlap was seen (<3%) in specific mRNAs co-precipitated with DRBD2. It is possible that the reduced overlap might reflect the much greater diversity of mRNAs, both in number and functional categories, bound by these proteins, with only a limited fraction of those being identifiable by the approach used. Nevertheless, a possible difference seen between DRBD2 and PABP2/PABP3 is a greater association with DRBD2 of mRNAs encoding mitochondrial proteins.
**In silico 3’UTR motifs within target mRNAs associated with the RBP23 or DRBD2 proteins**

In order to define 3’UTR sequence elements which could mediate any specific association between RBP23/DRBD2 and target mRNAs, we first used transcript length data available from *L. donovani* orthologues [48] to compare the length of coding sequence and untranslated regions of the top-most 20 transcripts enriched with either RBP23 or DRBD2 (Fig 7A). This comparison highlights noticeable differences in length between these mRNAs, with the RBP23-associated mRNAs having much shorter 5’UTRs and coding regions than the DRBD2-associated transcripts, as well as smaller 3’UTRs. With the sole exception of the RBP23 transcript, the other top 19 transcripts co-precipitated with RBP23 encode ribosomal proteins and a remarkable feature of these mRNAs in trypanosomatids is their small 5’UTRs, as reported from *T. brucei* [49]. Based on the *L. donovani* transcriptome data, and after excluding the 39 nucleotides spliced leader (mini-exon) sequence, these can be as small as eight nucleotides in length [48]. As highlighted in Fig 7A, the native RBP23 transcript differs from the top-most messages bound by the HA-tagged RBP23 in having a larger 5’UTR and coding...
region. mRNAs encoding both the native RBP23 and the ectopically expressed HA-tagged protein were specifically co-precipitated. This observation excludes any binding by RBP23 to the 5' or 3'UTRs of its native mRNA, absent from the ectopically encoded protein. It implies either an association with the protein coding sequence or, alternatively, a co-translational dimerization between the tagged protein and the nascent polypeptide, leading to its co-precipitation with the attached mRNA. It also reinforces that the nature of the association between ribosomal protein mRNAs and RBP23 is most-likely distinct from that between RBP23 and its own transcript.

Next, we searched for any sequence elements within the same set of RBP23 or DRBD2 co-precipitated transcripts that would allow them to be specifically selected by either protein. Considering the much-reduced size for the 5'UTRs of the RBP23-associated mRNAs, the search focused on elements localized to their 3'UTRs using 300 nucleotides immediately downstream of the *L. infantum* coding sequences and, independently, the mapped 3'UTRs of ribosomal protein mRNAs and RBP23 (blue) and DRBD2 (orange) 5' (▼) and 3' (▲) UTRs, CDS (■) and total transcript lengths (●) according to *L. donovani* orthologous sequences [48]. The mean values are represented by black lines and the RBP23 transcript is indicated by the same symbols coloured in black. (B) Putative motifs found within 3'UTRs from the top 20 mRNAs associated to either HA-tagged RBP23 or DRBD2 in *L. infantum* and *L. donovani*. The motifs found in searches with mRNAs enriched with the HA-tagged RBP23, including the RBP23 transcript as well as the top 19 mRNAs encoding ribosomal proteins, are shown on the left. Those motifs found for the top 20 mRNAs associated to HA-tagged DRBD2 are shown on the right. All logos were generated by MEME (Multiple Em for Motif Elicitation) and the relative height of each nucleotide represents the conservation at that specific position. The e-values are in the right corner of each logo. 

https://doi.org/10.1371/journal.pntd.0009899.g007
the equivalent *L. donovani* transcripts. Motifs rich in uridines were found to be a common feature within the 3'UTRs of the RBP23 bound transcripts in both *L. infantum* and *L. donovani* sequences (Fig 7B). These motifs were found in one to four copies in most 3'UTRs of the ribosomal protein mRNAs tested that co-precipitated with RBP23 but were noticeably absent from the 3'UTR of the RBP23 transcript (S6 Fig). Comparatively, for the 3'-UTRs of the mRNAs associated with DRBD2, a motif with UG repeats was found in both *Leishmania* species in nearly all mRNAs investigated (Figs 7B and S6). The markedly distinct motifs identified within the transcripts found associated with each protein indicate that, at least for the functionally related RBP23-bound messages, they might have a role in defining the association with these RBPs. It remains to be seen whether they would be directly recognized by these proteins or their recognition would require extra RNA-motifs and/or protein partners that might allow a more precise mRNA selection and binding.

**Discussion**

Although multiple PABP homologues have been previously reported from different organisms, mainly metazoans and plants [25,27,29], a selective association of those with specific mRNAs targets has not been generally defined. This study identifies, for the first time in *Leishmania*, the association between one PABP homologue and a specific group of cytoplasmic mRNAs, those encoding ribosomal proteins. These constitute an important subset of mRNAs, encoding a large group of abundant proteins whose expression is in general tightly regulated [50]. Our data agrees with the known importance, seen in metazoans and other organisms, for the regulation of translation of ribosomal protein mRNAs. It also highlights the fact that the regulatory mechanisms associated with these mRNAs converge on interactions between their 5' and 3' ends. The description of how a specific PABP homologue can be involved in such mechanisms, through direct interactions with another RBP, further expands on the known roles seen for different PABPs in model organisms.

RNAi-mediated depletion of RBP23 has been previously shown to affect growth of the *T. brucei* bloodstream forms [51], with these cells displaying a gain-of-fitness phenotype during differentiation, but depletion does not seem to affect procyclic cells [52]. In *L. infantum*, studies with isobaric tagging methodology (iTRAQ) showed that RBP23 is significantly downregulated during differentiation and in the mature amastigote forms [53], suggesting that this protein is needed during exponential growth phases. Our data confirm the association of RBP23 with PABP1 and the EIF4E4/EIF4G3 complex, all three proteins known to enhance expression in tethering assays and also present in polysomes [33,42,43,54]. The mass-spectrometry results also indicate an association of RBP23 with ribosomes or polysomes in *Leishmania*, although in *T. brucei* its orthologue was not found associated with the polysomal fraction [54]. The RBP23 association with a translationally active complex based on EIF4E4/EIF4G3 is reinforced by the presence of EIF4AI in the immunoprecipitated fractions, with the lower EIF4AI enrichment possibly being either a consequence of greater abundance or a loose association with the complex.

Some of the additional proteins that co-precipitated with RBP23 highlight other likely functions, apart from translation. An early association with the mRNA is reinforced by the RBP23 co-precipitation with mRNA polyadenylation and nucleo-cytoplasmic transport factors. Both RBP23 and DRBD2, together with the two *T. brucei* PABP homologues, were also co-purified with TSR1, involved in splicing regulation, mRNA stability, and rRNA processing [55]. The CCR4-NOT deadenylase complex is an important mRNA regulator, interacting with transcription and translation activators and repressors and promoting mRNA decay [56]. The
association with CCR4-NOT components may then indicate that RBP23 can remain bound to mRNA targets after their translation.

DRBD2 and two other proteins found here as co-precipitating partners, RBP12 and the uncharacterized LIN_F_20005800, all have T. brucei orthologues shown to reduce expression in tethering assays, while the orthologues of two other putative DRBD2 partners, PABP2 and PUF6, enhance expression [42,43]. T. brucei ZC3H31/34 and the LIN_210009700 orthologue (Tb927.10.2240) interact with the translational regulator MKT1 [44] while the ZC3H39/40 pair were identified as regulators of the mitochondrial respiratome [57]. The LIN_210009700 orthologue was further shown to interact with DRBD3, a cytoplasmic, stabilizing protein, which under oxidative stress remains bound to mRNA and concentrates in the nucleus, a typical behaviour of a protein involved in mRNA transport [58]. Our analysis also show a very relevant DRBD2 co-precipitation with proteins associated with the metabolism of mitochondrial RNAs. Although this is not supported by the current information regarding the DRBD2 localization [39], it also suggests a possible mitochondrial function. In all the evidence so far indicates that DRBD2 acts in different complexes with different contrasting functions.

In mammals, ribosomal proteins, as well as some translation factors, are encoded by mRNAs containing the 5′-terminal oligopyrimidine (TOP) sequence and which is recognized by the La-related protein 1 (LARP1) [50]. Leishmania RBP23, in contrast, is unlikely to specifically recognize its mRNA targets through 5′UTR TOP motifs, since all trypanosomatid mRNAs have the 5′ spliced leader sequence starting with two consecutive adenines [59]. Other sequence motifs within the 5′UTRs of mRNAs encoding ribosomal proteins and which could mediate specific recognition are unlikely, due to their small sizes [49]. These small 5′UTRs, however, raise another issue regarding the need for an elf4F complex, with an associated elf4A helicase. Once thought to be mainly required for the translation of mRNAs having structured 5′UTRs, elf4A has been recently shown to be required for the translation of most mRNAs, in both yeast and mammals [60,61]. The protozoan Giardia lamblia, which lacks the elf4G subunit of elf4F [62], is also characterized by mRNAs having very short 5′UTRs [63] which nevertheless seem to require elf4A, plus one of two Giardia elf4Es, to mediate their interaction with the translation pre-initiation complex [64]. These reports are consistent with a strict requirement for elf4A, and the trypanosomatid elf4AI, for the translation of mRNAs regardless of the size of their 5′UTRs.

Generally, mRNAs encoding ribosomal proteins have their translation differentially regulated through diverse mechanisms. In yeasts, the phosphorylation of an activator (Ifh1) and a repressor (Crf1) controls the transcription of genes encoding these mRNAs [65]. RNA sequencing analysis demonstrated that they are also especially enriched with the closed loop translation initiation components, elf4E and both isoforms of elf4G [66]. In mammals, the LARP1 protein, when associated with the TOP motif and the 5′ cap, acts as a translational repressor that binds to PABP and consequently form a translation-inactive mRNA loop. The mammalian target of rapamycin complex 1 (mTORC1), a kinase complex, phosphorylates LARP1 and releases it from 5′TOP mRNAs. This leads to the elf4F assembly, since mTORC1 also controls phosphorylation of 4E-BP1, thereby allowing elf4G to bind to elf4E and recruit the 43S complex to start translation [50,67,68].

Both Leishmania PABP1 and EIF4E are simultaneously phosphorylated during exponential growth [33,36], possibly associated with activation of translation of their target mRNAs. RBP23 may help recruit PABP1 and the EIF4E4/EIF4G3 based complex to selected mRNAs, mainly mRNAs encoding ribosomal proteins, to promote their translation. Here, we propose a model where the trypanosomatid RBP23 binds to common 3′UTRs motifs shared by these mRNAs and mediates their association with PABP1 (Fig 8). The RBP23 specificity for the
ribosomal protein mRNAs, in comparison to a broader range of mRNAs associated with PABP1, and the specificity of the later to poly(A) sequences, strongly implicates RBP23 as the protein responsible for mRNA recognition, on its own or dependent on additional protein partners. Thus, RBP23 may mark the ribosomal protein mRNAs as targets for translation mediated by the complex PABP1/EIF4E4/EIF4G3/EIF4A1. Preferential binding of this eIF4F complex to the mRNAs encoding ribosomal proteins would direct them to a translation pathway distinct from most other cellular mRNAs, perhaps avoiding competition and in agreement with these mRNAs behaving distinctively than most other messages during translation [69]. It would also allow their translation to be specifically regulated, presumably involving the phosphorylation of EIF4E4 and PABP1, an event already shown to be cell-cycle regulated [33,36,38]. Thus, according to the proposed model, translation regulation of the mRNAs encoding ribosomal proteins in trypanosomatids would also require interactions involving proteins bound to both the 5' (EIF4E4) and 3' (PABP1) ends of the mRNAs, as seen for the translation regulation of these mRNAs in mammals. It remains to be seen whether the EIF4E4/PABP1 interaction may also lead to a translation-inactive loop that may be enhanced or abolished by their simultaneous phosphorylation.

Materials and methods

Plasmid constructs and DNA manipulations

*L. infantum* genomic DNA from the MHOM/MA/67/ITMAP-263 strain was isolated using DNAzol (Life Technologies) following the manufacturer's instructions. Full length *RBP23, DRBD2, PABP2* and *PABP3* genes were amplified using primers flanked by BamHI and HindIII restriction sites (listed in S8 Table) and cloned into the BamHI-HindIII sites of a modified version of the *Leishmania* expression vector pSPBT1YNEn [70]. The modification consisted of the insertion of a 27 nucleotide element encoding the HA epitope (YPYDVPDYA) immediately after the coding sequence and prior to the translation stop codon. The *RBP23* and *DRBD2* genes were also subcloned into the BamHI-HindIII sites of the pET-21a vector (Novagen), while the *PABP2* and *PABP3* genes were further subcloned into the same sites of a modified pGEX4T3 expression vector (GE Healthcare) having an added HindIII site.

Parasite growth, transfections and expression analysis

*Leishmania major* MHOM/IL/81/Friedlin and *Leishmania infantum* MHOM/MA/67/ITMAP-263 promastigotes were cultured in Schneider's insect medium supplemented with 1% Penicillin-Streptomycin, 10% heat-inactivated Fetal Bovine Serum and 2% hemin at pH 7.2, 25˚C. Transfection of the circular plasmids was carried out by electroporation, with exponentially grown cells resuspended in HEPES-NaCl buffer (21 mM HEPES pH 7.05, 137 mM sodium chloride, 5 mM potassium chloride, 0.7 mM disodium phosphate, 6 mM glucose), incubated with the plasmid DNA and submitted to a pulse of 450 V, 500 μF on the Gene Pulser Xcell electroporation system (Bio-Rad). Transfected cells were selected with G418 (20 μg/ml, Sigma). For the expression analysis of cells expressing the HA-tagged proteins, late exponentially grown *L. infantum* cultures were harvested and resuspended directly into denaturing SDS-PAGE sample buffer, submitted to 15% SDS-PAGE and blotted with an anti-HA mouse monoclonal antibody (100 ng/ml, Applied Biological Materials).
Sequence analyses

Secondary structure and domain predictions for RBP23 were performed using the Phyre2 automatic fold recognition server [71]. Domains were also identified using the InterPro tool available at http://www.ebi.ac.uk/interpro/search/sequence/. The 3’ UTR analyses were
performed through MEME [72] using 300 nucleotides immediately after the stop codon from
the top 20 mRNA sequences bound to the L. infantum RBP23-HA and DRBD2-HA [48]. The
already defined 3’ UTRs of L. donovani orthologues were also retrieved and analysed. Default
parameters were used to detect the motifs, using any number of sites per sequence and a width
ranging between six and 50 nucleotides.

**Cytoplasmic extract preparation**

For the RNA sequencing analysis, whole *Leishmania major* (MHOM/IL/81/Friedlin) cyto-
plasmic extracts were produced from late exponentially grown promastigote cultures harvested
and washed once in ice cold PBS, prior to resuspension in IPM1 buffer [100 mM KCl, 5 mM
MgCl$_2$, 10 mM HEPES, protease inhibitors (Roche), RNaseOUT and 0.5% IGEPAL CA-630
(Sigma)] to a concentration of 1 to 2x10$^9$ cells/ml. The resuspended cells were left for 10 min-
utes on ice and then centrifuged for 10 more minutes at 17.000 g, 4˚C, with the sediment dis-
carded and the supernatants, the cytoplasmic extracts, aliquoted and stored at -80˚C. Whole
cytoplasmic extracts from L. infantum wild-type and strains expressing various HA-tagged
proteins were generated after cell lysis using nitrogen cavitation [73]. Briefly, late exponentially
grown L. infantum promastigotes were harvested and washed once in ice cold PBS, followed
by resuspension in HEPES-lysis buffer (20 mM HEPES-KOH pH7.4, 75 mM potassium ace-
tate, 4 mM magnesium acetate, 2 mM DTT, supplemented with EDTA-free protease inhibitors
from Roche) to a concentration of 1 to 2x10$^9$ cells/ml. The resuspended cells were trans-
ferred into the cavitation chamber of the cell disruption vessel (Parr Instruments) and incubated
at 4˚C under 70 bar pressure for 40 minutes, followed by rapid decompression/lysis. The lysates
were submitted to centrifugation, as described above, with the supernatants aliquoted and
stored likewise.

For mass-spectrometry analysis, total cytoplasmic extracts from L. infantum expressing
HA-tagged PABP1, PABP2, PABP3, as well as a phosphorylation PABP1 mutant (TP-SP),
were generated after lysis using acid-washed glass beads, 425–600 μm (Sigma), as described
previously [33]. For parasites expressing HA-tagged RBP23 and DRBD2, and wild-type con-
trols, total cytoplasmic extracts were obtained after cell lysis through nitrogen cavitation, as
described above. None of the cytoplasmic extracts used for the immunoprecipitation assays
and subsequent mass spectrometry analysis were treated with RNase.

**Immunoprecipitation studies**

For immunoprecipitations (IPs) with the HA-tagged proteins, whole L. infantum cytoplasmic
extracts from wild type or recombinant HA-tagged strains were mixed with Pierce Anti-HA
magnetic beads as per manufacturer’s protocol. Briefly, 0.2 mg of the beads were washed three
times with PBS followed by the incubation with 0.5 ml of cytoplasmic extracts (equivalent to
~5x10$^8$ to 10$^9$ cells) for 1 h at 4˚C. After removal of the depleted supernatant, the beads were
washed three times with PBS and the bound antigen-antibody complexes eluted in SDS-PAGE
sample buffer. Roughly 20% of the IPs were then analysed through SDS-PAGE and western-
blotting using antibodies against the HA-tag to confirm the efficiency of the precipitation reac-
tion. Immunoprecipitations were performed in triplicates and duplicates for RNA-sequencing
and mass spectrometry, respectively.

For IP assays against native PABPs, exclusively for RNA-sequencing, we used approxi-
ately 0.1 mg of protein A sepharose beads, whole L. major cytoplasmic extracts and previ-
ously described affinity purified antibodies raised against each of the three *Leishmania* PABPs
[30]. After a first wash with PBS, the beads were incubated with the antibodies (50 μL) in PBS
overnight at 4˚C, washed again with PBS and then incubated with the whole L. major extracts
(equivalent to 1×10⁶ cells) for 1 h, at 4˚C, in the presence of RNaseOUT. After spinning for 2
minutes at 600 g, 4˚C, the supernatant was removed and the beads washed three times sequential-
ly with IPM1 buffer containing 1% IGEPAL CA-630 and RNaseOUT.

RNA extraction and cDNA library construction

For the RNA sequencing of ligands bound to the HA-tagged proteins, RNA was extracted with
the RNAeasy Mini Kit (QIAGEN) from three independent immunoprecipitation experiments. These
were carried out with different batches of cytoplasmic extracts derived from wild-type L.
infantum and transfected cells expressing HA-RBP23, HA-DRBD2 and the three HA-tagged
PABPs. The RNA samples were quantified by Qubit RNA HS Assay Kit (Thermo Fisher) using
to read the concentration the Qubit 2.0 Fluorometer. Between 0.1 to 4 μg of total RNA was
used to construct the cDNA libraries with the TrueSeq Stranded mRNA Library Prep Kit (Illu-
mina). The libraries were validated quantitatively through qPCR using the KAPA Library
Quantification Kit and qualitatively by visualization in agarose gel. Finally, the libraries were
normalized and pooled prior to sequencing using the MiSeq Reagent Kit v3, 150 cycle (Illu-
mina). Similar RNA extractions were performed for the L. major IPs carried out with the
native anti-PABP affinity purified antibodies, and total RNA was used to prepare cDNA librar-
ies using the SOLiD Whole Transcriptome Analysis Kit followed by evaluation with an Agilent
Bioanalyzer (Agilent). The cDNA libraries were used for clonal amplification according to the
SOLiD Full-Scale Template Bead preparation protocol and sequenced with the SOLiD4 System
(Applied Biosystems).

RNA sequencing analysis and data normalization

RNA-Seq reads obtained from SOLiD data were mapped against the Leishmania major Fri-
edlin genome assembly version 8.1 available at the TriTrypDB database by the SHRiMP soft-
ware version 2.2.3 [74], with default parameters. All mappings whose score was higher than
350 were considered for further analyses, where all samples were normalized and the differen-
tial expression was assessed by the EdgeR package [75], included in the Bioconductor package
version 2.6 [76]. mRNAs co-precipitated with each HA-tagged protein were analysed with the
following bioinformatic tools: (1) FastQC, to evaluate the quality of the sequences (https://
www.bioinformatics.babraham.ac.uk/projects/fastqc/); (2) Trimmomatic [77], version 0.36, to
remove the adapters and the low-quality sequences; (3) STAR [78], to map the reads on the L.
infantum genome and count reads associated with each gene; (4) DEseq2 Galaxy version [79],
to statistically compare the samples. Genes were considered enriched when a four-fold
increase (log₂ ratio > 2) was observed when compared to the negative control, with FDR of
0.01 or 0.05 considered, respectively, for the IPs with the native proteins or HA-tagged ones.
mRNAs enriched with the different proteins were then classified according to a modified list
of gene ontology (GO) functional terms. For the RBP23 mRNA analysis, the transcript
sequence containing the 3’ UTR and the transcript sequence with HA, but without the 3’ UTR,
was used as reference. Bowtie2, a tool for aligning sequencing read [80] was used to generate
the bam files, which were viewed in IGV, integrative genomics viewer, a visualization tool
[81,82].

Mass-spectrometry analysis

For protein digestion and mass-spectrometry, IPs of HA-tagged PABP1, TP-SP, PABP2 and
PABP3 were analysed by the Proteomics platform of the Quebec Genomics Center at the CHU
de Quebec Research Center-Université Laval (http://www.crchudequebec.ulaval.ca/en/
services/proteomics/about-us/), as described [83]. The results of two independent experiments
were analysed by the Scaffold Proteome software used to validate the protein identification based on the \textit{L. infantum} genome. Only proteins identified with \(>2\) peptide and a probability of \(>80.0\%\) were considered. Eluted proteins from IP of HA-tagged RBP23 and DRBD2, also from two independent experiments, each with two sets of negative controls, were submitted to the mass spectrometry facility P02-004 at the Carlos Chagas Institute–Fiocruz-PR. The samples were loaded into 15\% SDS-PAGE gels and allowed to migrate into the resolving gel, when the electrophoresis was interrupted prior to protein fractionation. Gel slices containing the whole IP products were then excised and submitted to an in-gel trypptic digestion and mass spectrometry analysis and validation was performed as previously described [47]. Spectra were searched against the \textit{L. infantum} protein sequence database (\textit{L. infantum} JPCM5, version from March 29, 2016, available at TriTrypDB; https://tritrypdb.org/). To normalize the data from the IPs with the HA-tagged RBP23 and DRBD2, a first normalization was performed based on the sum of the intensities for each replicate, using the highest sum to normalize the remaining samples. For each co-precipitated polypeptide, for the negative control and both RBPs, the averages derived from the normalized intensities were calculated. These averages were then used to calculate the ratio between the values from the IPs with each HA-tagged protein and those control IPs using extracts from non-transfected cells (enrichment ratio).

\textbf{In vitro pull-down assays}

Pull-down assays were performed using Glutathione Sepharose 4B beads (GE Healthcare) as well as GST-tagged and \(^{35}\text{S}\)-labeled recombinant proteins, as described previously [36,84]. GST-tagged PABPs and the GST control, whose genes were cloned into the pGEX4T3 expression vector, were expressed in \textit{Escherichia coli}, affinity purified, immobilized on the beads and incubated with \(^{35}\text{S}\)-labeled RBP23 and DRBD2. The labeled proteins were produced after linearization with HindIII of the corresponding constructs in the pET21a plasmid, followed by transcription with T7 RNA polymerase in the presence of the cap analogue and translation in the rabbit reticulocyte lysate (Promega or Ambion) supplemented with \(^{35}\text{S}\)-methionine (Perkin Elmer). The labeled signals were visualized by autoradiographic films exposed unto 15\% SDS-PAGE gels.

\textbf{Supporting information}

\textbf{S1 Fig. Analysis of mRNA populations associated with the three native PABPs in \textit{Leishmania major}.} Upregulated transcripts in at least two of three available RNA-seq datasets (SOLiD sequencing) were manually classified and grouped using the gene ontology (GO) terms according to their molecular function. A) mRNA groups associated with PABP1, PABP2 and PABP3 from \textit{L. major}. All mRNAs enriched at least 2-fold more than the negative control are represented; B) Bar chart representing the enrichment values only of mRNAs co-immunoprecipitated with the three \textit{L. major} PABPs that were enriched at least 4-fold. The mRNAs with the same names indicate different transcripts encoding proteins with the same name but whose genes localize to different chromosomes.

\textbf{(TIF)}

\textbf{S2 Fig. Expression analysis of the HA-tagged \textit{L. infantum} PABPs.} Western-blot analysis of exponentially grown \textit{L. infantum} cell lines expressing the HA-tagged \textit{L. infantum} PABPs. Approximate molecular weights of 63, 65 and 61 kDa were seen for PABP1-HA, PABP2-HA and PABP3-HA, respectively.

\textbf{(TIF)}
S3 Fig. Volcano Plot comparing mRNA content of *L. infantum* PABPs RNA-seq datasets. Axis Y represents the statistical test significance 0.05 (>0.05 red and <0.05 black) and the axis X represents the genes that are at least 4-fold up- or down-regulated (log₂ fold change >2); Δ means a very high value to be ranked. (A) PABP2xPABP1; (B) PABP3xPABP1; and (C) PABP3xPABP2.

(TIF)

S4 Fig. Comparison of RNA-seq datasets from RNA-immunoprecipitation studies using the three *Leishmania infantum* PABPs. The mRNAs associated to each PABP homologue are indicated in a Venn diagram from their respective RNAseq datasets.

(TIF)

S5 Fig. Summary of RBPs co-purifying with the *L. infantum* HA-tagged PABPs. RBPs shown here are differentially co-precipitated with PABP1 and PABP2/3 in two independent immunoprecipitation (IP) assays. The values represent the number of peptide hits from proteins found with each PABP. Only proteins identified with two or more peptides in a minimum of two replicates and with a probability >80.0% were considered. TP-SP represents a PABP1 phosphorylation mutant (described in [33]), used as a second PABP1 sample, since it was presumed that the mutations would not impact on the interactions with major binding partners.

(TIF)

S6 Fig. Localization of the most-relevant motifs found on the topmost 10 mRNAs enriched with the *L. infantum* HA-tagged RBP23 or DRBD2. A) U-rich motifs found within the 3’ UTRs of the RBP23 associated mRNAs; B) Motifs having UG-repeats found within the 3’ UTRs of the DRBD2 associated mRNAs.

(TIF)

S1 Table. mRNAs specifically co-precipitated with the three *L. major* PABP homologues.

(XLSX)

S2 Table. mRNAs specifically co-precipitated with the three, HA-tagged, *L. infantum* PABP homologues.

(XLSX)

S3 Table. Full set of protein partners specifically co-precipitated with the HA-tagged RBP23.

(XLSX)

S4 Table. Top-most protein partners specifically co-precipitated with the HA-tagged RBP23.

(XLSX)

S5 Table. Full set of protein partners specifically co-precipitated with the HA-tagged DRBD2.

(XLSX)

S6 Table. Top-most protein partners specifically co-precipitated with the HA-tagged DRBD2.

(XLSX)

S7 Table. mRNAs specifically co-precipitated with the HA-tagged *L. infantum* RBP23 and DRBD2.

(XLSX)
S8 Table. Oligonucleotides used as primers for the amplification and cloning of the PABP2, PABP3, RBP23 and DRBD2 genes.

(XLSX)

Acknowledgments

We thank all the members of the de Melo Neto’s laboratory for helpful discussions and technical assistance. The authors thank FIOCRUZ for using the Technological Platforms Network for the funding and use of the automatic sequencing facility at the Aggeu Magalhães Institute and the Proteomics facility at the Carlos Chagas Institute.

Author Contributions

Conceptualization: Ludmila A. Assis, Fabiola B. Holetz, Tamara D. C. da Costa Lima, Osvaldo P. de Melo Neto.

Data curation: Ludmila A. Assis, Moezio V. C. Santos Filho, Maria J. R. Bezerra, Fabiola B. Holetz, Christian M. Probst, Osvaldo P. de Melo Neto.

Formal analysis: Ludmila A. Assis, Moezio V. C. Santos Filho, Maria J. R. Bezerra, Fabiola B. Holetz, Christian M. Probst, Osvaldo P. de Melo Neto.

Funding acquisition: Fabiola B. Holetz, Osvaldo P. de Melo Neto.

Investigation: Ludmila A. Assis, Moezio V. C. Santos Filho, Joao R. da Cruz Silva, Maria J. R. Bezerra, Irassandra R. P. U. C. de Aquino, Kleison C. Merlo, Fabiola B. Holetz, Tamara D. C. da Costa Lima.

Methodology: Ludmila A. Assis, Moezio V. C. Santos Filho, Joao R. da Cruz Silva, Maria J. R. Bezerra, Irassandra R. P. U. C. de Aquino, Kleison C. Merlo, Fabiola B. Holetz, Tamara D. C. da Costa Lima, Osvaldo P. de Melo Neto.

Project administration: Osvaldo P. de Melo Neto.

Resources: Barbara Papadopoulou.

Software: Christian M. Probst, Antonio M. Rezende.

Supervision: Barbara Papadopoulou, Tamara D. C. da Costa Lima, Osvaldo P. de Melo Neto.

Validation: Ludmila A. Assis, Moezio V. C. Santos Filho, Joao R. da Cruz Silva, Maria J. R. Bezerra, Irassandra R. P. U. C. de Aquino, Kleison C. Merlo, Fabiola B. Holetz, Tamara D. C. da Costa Lima.

Visualization: Ludmila A. Assis, Moezio V. C. Santos Filho, Osvaldo P. de Melo Neto.

Writing – original draft: Ludmila A. Assis, Osvaldo P. de Melo Neto.

Writing – review & editing: Ludmila A. Assis, Osvaldo P. de Melo Neto.

References

1. Jackson AP. Genome evolution in trypanosomatid parasites. Parasitology. 2015; 142: S40–S56. https://doi.org/10.1017/S0031182014000894 PMID: 25068268

2. Clayton CE. Networks of gene expression regulation in Trypanosoma brucei. Mol Biochem Parasitol. 2014; 195: 96–106. https://doi.org/10.1016/j.molbiopara.2014.06.005 PMID: 24995711

3. Clayton C. Regulation of gene expression in trypanosomatids: living with polycistronic transcription. Open Biol. 2019; 9. https://doi.org/10.1098/rsob.190072 PMID: 31164049
4. Martinez-Calvillo S, Vizuete-De-Rueda JC, Florencio-Martinez LE, Manning-Cela RG, Figueroa-Angulo EE. Gene expression in trypanosomatid parasites. J Biomed Biotechnol. 2010; 2010. https://doi.org/10.1155/2010/525241 PMID: 20691133

5. De Gaudenzi JG, Noe G, Campo VA, Frasch AC, Cassola A. Gene expression regulation in trypanosomatids. Essays Biochem. 2011; 51: 31–46. https://doi.org/10.1042/ebse0510031 PMID: 2203440

6. Yamashita A, Takeuchi O. Translational control of mRNAs by 3'-untranslated region binding proteins. BMB Rep. 2017; 50: 194–200. https://doi.org/10.5483/bmbrep.2017.50.4.040 PMID: 28287067

7. Khong A, Parker R. The landscape of eukaryotic mRNPs. RNA. 2020; 26: 229–239. https://doi.org/10.1016/j.molcel.2020.00.001 PMID: 32083019

8. Singh G, Pratt G, Yeo GW, Moore MJ. The clothes make the mRNA: past and present trends in mRNP fashion. Annu Rev Biochem. 2015; 84: 325–354. https://doi.org/10.1146/annurev-biochem-080111-092106 PMID: 25784054

9. Harvey RF, Smith TS, Muloney T, Queiroz RML, Pizzinga M, Dezi V, et al. Trans-acting translational regulatory RNA binding proteins. Wiley Interdiscip Rev RNA. 2018; 9: 1–19. https://doi.org/10.1002/wrna.1465 PMID: 29341429

10. Müller-McNicol M, Neugebauer KM. How cells get the message: dynamic assembly and function of mRNA–protein complexes. Nat Rev Genet. 2013; 14: 275–287. https://doi.org/10.1038/nrg3434 PMID: 23478349

11. Wurth L, Gebauer F. RNA-binding proteins, multifaceted translational regulators in cancer. Biochim Biophys Acta—Gene Regul Mech. 2015; 1849: 881–886. https://doi.org/10.1016/j.bbagrm.2014.10.001 PMID: 25316157

12. Cho H, Cho HS, Hwang I. Emerging roles of RNA-binding proteins in plant development. Curr Opin Plant Biol. 2019; 51: 51–57. https://doi.org/10.1016/j.pbi.2019.03.016 PMID: 31071564

13. Alonso CR. A complex “mRNA degradation code” controls gene expression during animal development. Trends Genet. 2012; 28: 78–88. https://doi.org/10.1016/j.tig.2011.10.005 PMID: 22257633

14. Romagnoli BAA, Holetz FB, Alves LR, Goldenberg S. RNA Binding Proteins and gene expression regulation. Front Cell Infect Microbiol. 2020; 10: 1–17. https://doi.org/10.3389/fcimb.2020.00001 PMID: 32083019

15. Kramer S, Carrington M. Trans-acting proteins regulating mRNA maturation, stability and translation in trypanosomatids. Trends Parasitol. 2011; 27: 23–30. https://doi.org/10.1016/j.pt.2010.06.011 PMID: 20609625

16. Kolev NG, Ullu E, Tschudi C. The emerging role of RNA-binding proteins in the life cycle of Trypanosoma cruzi. Cell Microbiol. 2014; 16: 482–489. https://doi.org/10.1111/cmi.12268 PMID: 24438230

17. Dupé A, Dumas C, Papadopoulou B. An Alba-domain protein contributes to the stage-regulated stability of amastin transcripts in Leishmania. Mol Microbiol. 2014; 91: 546–561. https://doi.org/10.1111/mmi.12478 PMID: 24620725

18. Azizi H, Dumas C, Papadopoulou B. The Pumilio-domain protein PUF6 contributes to SIDER2 retrotranson-mediated mRNA decay in Leishmania. RNA. 2017; 23: 1874–1885. https://doi.org/10.1261/roma.062950.117 PMID: 28877997

19. Romanuik MA, Cervini G, Cassola A. Regulation of RNA binding proteins in trypanosomatid protozoan parasites. World J Biol Chem. 2016; 7: 146–157. https://doi.org/10.4331/wjbc.v7.i1.146 PMID: 26981203

20. Clayton C. The regulation of trypanosome gene expression by RNA-binding proteins. Knoll LJ, editor. PLoS Pathog. 2013; 9: e1003680. https://doi.org/10.1371/journal.ppat.1003680 PMID: 24244152

21. Karamysheva ZN, Guarinzo ŠAG, Karamyshev AL. Regulation of translation in the protozoan parasite leishmania. Int J Mol Sci. 2020; 21: 1–15. https://doi.org/10.3390/ijms21082981 PMID: 32340274

22. Najafabadi HS, Lu Z, MacPherson C, Mehta V, Adoue V, Pastinen T, et al. Global identification of conserved post-transcriptional regulatory programs in trypanosomatids. Nucleic Acids Res. 2013; 41: 8591–600. https://doi.org/10.1093/nar/gkt647 PMID: 23877242

23. Ouellette M, Papadopoulou B. Coordinated gene expression by post-transcriptional regulations in African trypanosomes. J Biol. 2009; 8: 100. https://doi.org/10.1186/jbionet203 PMID: 20017896

24. Wigington CP, Williams KR, Meers MP, Bassell GJ, Corbett AH. Poly(A) RNA-binding proteins and polyadenosine RNA: new members and novel functions. Wiley Interdiscip Rev RNA. 2014; 13: 601–622. https://doi.org/10.1002/wrna.1233 PMID: 24789627

25. Eliseeva IA, Lyabin DN, Ovchinnikov LP. Poly(A)-binding proteins: structure, domain organization, and activity regulation. Biochemistry (Moscow). 2013; 78: 1377–91. https://doi.org/10.1134/S0006297913100147 PMID: 24490729

26. Goss DJ, Kleinman FE. Poly(A) binding proteins: Are they all created equal? Wiley Interdiscip Rev RNA. 2013; 4: 167–179. https://doi.org/10.1002/wrna.1151 PMID: 23424172
27. Smith RWP, Blee TKP, Gray NK. Poly(A)-binding proteins are required for diverse biological processes in metazoans. Biochem Soc Trans. 2014; 42: 1229–1237. https://doi.org/10.1042/BST20140111 PMID: 25110030

28. Chen Z, Jolley B, Caldwell C, Galley DR. Eukaryotic translation initiation factor eIFiso4G is required to regulate violaxanthin De-epoxidase expression in Arabidopsis. J Biol Chem. 2014; 289: 13926–36. https://doi.org/10.1074/jbc.M114.555151 PMID: 24706761

29. Galley DR, Liu R. Phylogenetic analysis reveals dynamic evolution of the poly(A)-binding protein gene family in plants. BMC Evol Biol. 2014; 14: 238. https://doi.org/10.1186/s12862-014-0238-4 PMID: 25421536

30. da Costa Lima TD, Moura DNM, Reis CRS, Vasconcelos JRC, Ellis L, Carrington M, et al. Functional characterization of three leishmania poly(a) binding protein homologues with distinct binding properties to RNA and protein partners. Eukaryot Cell. 2010; 9: 1484–1494. https://doi.org/10.1128/EC.00148-10 PMID: 20675580

31. Kramer S, Bannerman-Chukualim B, Ellis L, Boulden EA, Kelly S, Field MC, et al. Differential localization of the two l Poly(A) Binding Proteins to the nucleolus and RNP granules suggests binding to distinct mRNA pools. Preiss T, editor. PLoS One. 2013; 8: e54004. https://doi.org/10.1371/journal.pone.0054004 PMID: 23382864

32. Zoltnar M, Krienitz N, Field MC, Kramer S. Comparative proteomics of the two T. brucei/PABPs suggests that PABP2 controls bulk mRNA. Acosta-Serrano A, editor. PLoS Negl Trop Dis. 2018; 12: e006879. https://doi.org/10.1371/journal.pntd.006879 PMID: 30040867

33. de Melo Neto OP, da Costa Lime TDC, Merlo KC, Romão TP, Rocha PO, Assis LA, et al. Phosphorylation and interactions associated with the control of the Leishmania Poly-A Binding Protein 1 (PABP1) function during translation initiation. RNA Biol. 2018; 6286: 1–17. https://doi.org/10.1080/15476286.2018.1449958 PMID: 29569995

34. Freire E, Sturm N, Campbell D, de Melo Neto O. The role of cytoplasmic mRNA cap-binding protein complexes in Trypanosoma brucei and other trypanosomatids. Pathogens. 2017; 6: 55. https://doi.org/10.3390/pathogens6040055 PMID: 29077018

35. Zinoviev A, Léger M, Wagner G, Shapira M. A novel 4E-interacting protein in Leishmania is involved in stage-specific translation pathways. Nucleic Acids Res. 2011; 39: 8404–15. https://doi.org/10.1093/nar/gkr555 PMID: 21764780

36. de Melo Neto OP, da Costa Lime TDC, Xavier CC, Nascimento LM, Romão TP, Assis LA, et al. The unique Leishmania EIF4E N-terminus is a target for multiple phosphorylation events and participates in critical interactions required for translation initiation. RNA Biol. 2015; 12: 1–13. https://doi.org/10.1080/15476286.2015.1086865 PMID: 26331814

37. Hu H, Gourguechon S, Wang CC, Li Z. The G1 cyclin-dependent kinase CRK1 in Trypanosoma brucei regulates anterograde protein transport by phosphorylating the COPII subunit Sec31. J Biol Chem. 2016; 291: 15527–15539. https://doi.org/10.1074/jbc.M116.715185 PMID: 27252375

38. An T, Liu Y, Gourguechon S, Wang CC, Li Z. CDK phosphorylation of translation initiation factors couples protein translation with cell-cycle transition. Cell Rep. 2018; 25: 3204–3214.e5. https://doi.org/10.1016/j.celrep.2018.11.063 PMID: 30540951

39. Dean S, Sunter JD, Wheeler RJ. TrypTag: a Trypanosoma genome-wide protein localisation resource. Trends Parasitol. 2017; 33: 80–82. https://doi.org/10.1016/j.pt.2016.10.009 PMID: 27863903

40. De Gaudenzi J, Frasch AC, Clayton C. RNA-binding domain proteins in kinetoplastids: a comparative analysis. Eukaryot Cell. 2005; 4: 2106–14. https://doi.org/10.1128/EC.4.12.2106-2114.2005 PMID: 16339728

41. Fritz M, Vanselow J, Sauer N, Lamer S, Goos C, Siegel TN, et al. Novel insights into RNP granules by employing the trypanosome’s microtubule skeleton as a molecular sieve. Nucleic Acids Res. 2015; 43: 8013–8032. https://doi.org/10.1093/nar/gkj731 PMID: 26187993

42. Lueong S, Merce C, Fischer B, Hoheisel JD, Erben ED. Gene expression regulatory networks in Trypanosoma brucei insights into the role of the mRNA-binding proteome. Mol Microbiol. 2016; 100: 457–471. https://doi.org/10.1111/mmi.13328 PMID: 26784394

43. Erben ED, Fadda A, Lueong S, Hoheisel JD, Clayton C. A genome-wide tethering screen reveals novel potential post-transcriptional regulators in Trypanosoma brucei. PLOS Pathog. 2014; 10: e1004178. https://doi.org/10.1371/journal.ppat.1004178 PMID: 24945722

44. Singh A, Minia I, Droll D, Fadda A, Clayton C, Erben E. Trypanosomone MKT1 and the RNA-binding protein ZC3H11: interactions and potential roles in post-transcriptional regulatory networks. Nucleic Acids Res. 2014; 42: 4652–68. https://doi.org/10.1093/nar/gkt1416 PMID: 24470144

45. Wippel HH, Malargin JS, Inoue AH, Leprevost FDV, Carvalho PC, Goldenberg S, et al. Unveiling the partners of the DRBD2-mRNP complex, an RBP in Trypanosoma cruzi and ortholog to the yeast SR-
46. Aphasizheva I, Alfonzo J, Cames J, Cestari I, Cruz-Reyes J, Göringer HU, et al. Lexis and grammar of mitochondrial RNA processing in trypanosomes. Trends Parasitol. 2020; 36: 337–355. https://doi.org/10.1016/pt.2020.01.006 PMID: 32191849

47. Rezende AM, Assis LA, Nunes ECC, da Costa Lima TDD, Marchini FKK, Freire ERR, et al. The translation initiation complex elf3 in trypanosomatids and other pathogenic excavates—identification of conserved and divergent features based on orthologue analysis. BMC Genomics. 2014; 15: 1175. https://doi.org/10.1186/1471-2164-15-1175 PMID: 25539953

48. Camacho E, González-de la Fuente S, Rastrojo A, Peiró-Pastor R, Solana JC, Tabera L, et al. Complete assembly of the Leishmania donovani (H3U strain) genome and transcriptome annotation. Sci Rep. 2019; 9: 1–15. https://doi.org/10.1038/s41598-018-37168-2 PMID: 30626917

49. Jensen BC, Ramasamy G, Vasconcelos EJR, Myler PJ, Parsons M. Extensive stage-regulation of translation revealed by ribosome profiling of Trypanosoma brucei. BMC Genomics. 2014; 15: 911. https://doi.org/10.1186/1471-2164-15-911 PMID: 25331479

50. Fonseca BD, Lahr RM, Damgaard CK, Alain T, Parsons M. LARP1 on TOP of ribosome production. Wiley Interdiscip Rev RNA. 2018; 9: 1–16. https://doi.org/10.1002/wrna.1480 PMID: 29722158

51. Wurst M, Robles A, Po J, Luu VD, Brems S, Marentije M, et al. An RNAi screen of the RRM-domain proteins of Trypanosoma brucei. Mol Biochem Parasitol. 2009; 163: 61–65. https://doi.org/10.1016/j.molbiopara.2008.09.001 PMID: 18840477

52. Alsford S, Turner DJ, Obado SO, Sanchez-Flores A, Glover L, Berriman M, et al. High-throughput phenotyping using parallel sequencing of RNA interference targets in the African trypanosome. Genome Res. 2011; 21: 915–924. https://doi.org/10.1101/gr.115089.110 PMID: 21363968

53. Rosenzweig D, Smith D, Opperdoes F, Stern S, Olafson RW, Zilberstein D. Retooling Leishmania metabolism: from sand fly gut to human macrophage. FASEB J. 2008; 22: 590–602. https://doi.org/10.1096/fj.07-9254com PMID: 17884972

54. Klein C, Terrao M, Inchastegui Gil D, Clayton C. Polysomes of Trypanosoma brucei. Association with Initiation Factors and RNA-Binding Proteins. PLoS One. 2015; 10: e0135973. https://doi.org/10.1371/journal.pone.0135973 PMID: 26287607

55. Gupta SK, Chikne V, Eliaz D, Tkacz ID, Naboishchikov I, Carmi S, et al. Two splicing factors carrying serine-arginine motifs, TSR1 and TSR1IP, regulate splicing, mRNA stability, and rRNA processing in Trypanosoma brucei. RNA Biol. 2014; 11: 715–731. https://doi.org/10.4161/rna.29143 PMID: 24922194

56. Collart MA. The Ccr4-Not complex is a key regulator of eukaryotic gene expression. Wiley Interdiscip Rev RNA. 2016; 7: 438–454. https://doi.org/10.1002/wrna.1332 PMID: 26821858

57. Treneman A, Glover L, Hutchinson S, Horn D. A post-transcriptional respiratorie regulon in trypanosomes. Nucleic Acids Res. 2019; 47: 7063–7077. https://doi.org/10.1093/nar/gkz455 PMID: 31127277

58. Fernandez-Moya SM, Garcia-Perez A, Kramer S, Carrington M, Estevez AM, Fernandez-Moya SM, et al. Alterations in DRBD3 ribonucleoprotein complexes in response to stress in Trypanosoma brucei. Mol Biochem Parasitol. 2017; 214: 306–312. https://doi.org/10.1016/j.molbiopara.2017.05.012 PMID: 28559306

59. Michaeli S. Trans-splicing in trypanosomes: machinery and its impact on the parasite transcriptome. Future Microbiol. 2011; 6: 459–74. https://doi.org/10.2217/fmb.11.20 PMID: 21526946

60. Sokabe M, Fraser CS. A helicase-independent activity of elf4A in promoting mRNA recruitment to the human ribosome. Proc Natl Acad Sci U S A. 2017; 114: 6304–6309. https://doi.org/10.1073/pnas.1620426114 PMID: 28559306

61. Yourik P, Aitken CE, Zhou F, Gupta N, Hinnebusch AG, Lorsch JR. Yeast elf4A enhances recruitment of mRNAs regardless of their structural complexity. Elife. 2017; 6: 1–27. https://doi.org/10.7554/eLife.31476 PMID: 29192585

62. Li L, Wang CC. Capped mRNA with a single nucleotide leader is optimally translated in a primitive eukaryote, Giardia lamblia. J Biol Chem. 2004; 279: 14656–14664. https://doi.org/10.1074/jbc.M309879200 PMID: 14722094

63. Adam RD. Biology of Giardia lamblia. Clin Microbiol Rev. 2001; 14: 447–475. https://doi.org/10.1128/CMR.14.3.447-475.2001 PMID: 11432808

64. Adedoja AN, McMahan T, Neal JP, Hamal Dhakal S, Jois S, Romo D, et al. Translation initiation factors GleIF4E2 and GleIF4A can interact directly with the components of the pre-initiation complex to facilitate translation initiation in Giardia lamblia. Mol Biochem Parasitol. 2020; 236: 111258. https://doi.org/10.1016/j.molbiopara.2020.111258 PMID: 31968220
65. Kim MS, Hahn JS. Role of CK2-dependent phosphorylation of Ifh1 and Crf1 in transcriptional regulation of ribosomal protein genes in Saccharomyces cerevisiae. Biochim Biophys Acta—Gene Regul Mech. 2016; 1859: 1004–1013. https://doi.org/10.1016/j.bbagrm.2016.06.003 PMID: 27321754

66. Costello J, Castelli LM, Rowe W, Kershaw CJ, Talavera D, Mohammad-Qureshi SS, et al. Global mRNA selection mechanisms for translation initiation. Genome Biol. 2015; 16: 10. https://doi.org/10.1186/s13059-014-0559-z PMID: 25650959

67. Fonseca BD, Zakaria C, Jia JJ, Graber TE, Svitkin Y, Tahmasebi S, et al. La-related protein 1 (LARP1) represses terminal oligopyrimidine (TOP) mRNA translation downstream of mTOR complex 1 (mTORC1). J Biol Chem. 2015; 290: 15996–16020. https://doi.org/10.1074/jbc.M114.621730 PMID: 25940091

68. Philippe L, Vasseur JJ, Debart F, Thoreen CC. La-related protein 1 (LARP1) repression of TOP mRNA translation is mediated through its cap-binding domain and controlled by an adjacent regulatory region. Nucleic Acids Res. 2018; 46: 1457–1469. https://doi.org/10.1093/nar/gkj1237 PMID: 29244122

69. Antwi EB, Haanstra JR, Ramsamy G, Jensen B, Droll D, Rojas F, et al. Integrative analysis of the Trypanosoma brucei gene expression cascade predicts differential regulation of mRNA processing and unusual control of ribosomal protein expression. BMC Genomics. 2016; 17: 306. https://doi.org/10.1186/s12866-016-2624-3 PMID: 27118143

70. Padmanabhan PK, Dumas C, Samant M, Rochette A, Simard MJ, Papadopoulos B. Novel features of a PIWI-like protein homolog in the parasitic protozoan Leishmania. PLoS One. 2012; 7: e52612. https://doi.org/10.1371/journal.pone.0025612 PMID: 22365111

71. Kelley L, Mezulis S, Yates C, Wass M, Sternberg M. The Phyre2 web portal for protein modelling, prediction, and analysis. Nat Protoc. 2015; 10: 845–858. https://doi.org/10.1038/nprot.2015.053 PMID: 25950237

72. Bailey, Timothy L.; Elkan C. Fitting a mixture model by expectation maximization to discover motifs in biopolymers. Proceedings of the Second International Conference on Intelligent Systems for Molecular Biology. Menlo Park, California: AAAI press; 1994. pp. 28–36.

73. Mureev S, Kovtun O, Nguyen UTT, Alexandrov K. Species-independent translational leaders facilitate cell-free expression. Nat Biotechnol. 2009; 27: 747–752. https://doi.org/10.1038/nbt.1556 PMID: 19649099

74. David M, Dzamba M, Lister D, Ilie L, Brudno M. SHRiMP 2: Sensitive yet practical short read mapping. Bioinformatics. 2014; 30: 2114–20. https://doi.org/10.1093/bioinformatics/btu170

75. Robinson MD, McCarthy DJ, Smyth GK. edgeR: A Bioconductor package for differential expression analysis of digital gene expression data. Bioinformatics. 2009; 26: 139–140. https://doi.org/10.1093/bioinformatics/btp616 PMID: 19910308

76. Gentleman RC CV, Bates DM, Bolstad B, Dettling M, Dudoit S, Ellis B, et al. Bioconductor: open-source development for computational biology and bioinformatics. Genome Biol. 2004; 5. https://doi.org/10.1186/gb-2004-5-10-r80 PMID: 15461798

77. Bolger AM, Lohse M, Usadel B. Trimmomatic: a flexible trimmer for Illumina sequence data. Bioinformatics. 2014; 30: 2114–20. https://doi.org/10.1093/bioinformatics/btu170 PMID: 24695404

78. Dobin A, Davis CA, Schlesinger F, Drenkow J, Zaleski C, Jha S, et al. STAR: Ultrafast universal RNA-seq aligner. Bioinformatics. 2013; 29: 15–21. https://doi.org/10.1093/bioinformatics/bts635 PMID: 23104886

79. Afgan E, Baker D, Batut B, Van Den Beek M, Bouvier D, Ech M, et al. The Galaxy platform for accessible, reproducible and collaborative biomedical analyses: 2018 update. Nucleic Acids Res. 2018; 46: W537–W544. https://doi.org/10.1093/nar/gky379 PMID: 29790989

80. Langmead B, Salzberg SL. Fast gapped-read alignment with Bowtie 2. Nat Methods. 2012; 9: 357–9. https://doi.org/10.1038/nmeth.1923 PMID: 22386286

81. Robinson JT, Thorvaldsdottir H, Winckler W, Guttman M, Lander ES, Getz G, et al. Integrative genomics viewer. Nat Biotechnol. 2011; 29: 24–26. https://doi.org/10.1038/nbt.1754 PMID: 21221095

82. Robinson JT, Thorvaldsdottir H, Wenger AM, Zehir A, Mesirov JP. Variant review with the integrative genomics viewer. Cancer Res. 2017; 77: e31–e34. https://doi.org/10.1158/0008-5472.CAN-17-0337 PMID: 29092934

83. Dupe A, Dumas C, Papadopoulos B. Differential Subcellular Localization of Leishmania Alba-Domain Proteins throughout the Parasite Development. PLoS One. 2015; 10: e0137243. https://doi.org/10.1371/journal.pone.0137243 PMID: 26334886

84. Dhala R, Reis CR, Freire ERR, Rocha POO, Katz R, Muniz JRCR, et al. Translation initiation in Leishmania major characterisation of multiple eIF4F subunit homologues. MolBiochemParasitol. 2005; 140: 23–41. https://doi.org/10.1016/j.molbiopara.2004.12.001 PMID: 15694484