Protein-protein interactions of the nicotinamide/nicotinate mononucleotide adenylyltransferase of *Leishmania braziliensis*

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BACKGROUND Nicotinamide adenine dinucleotide (NAD) plays a central role in energy metabolism and integrates cellular metabolism with signalling and gene expression. NAD biosynthesis depends on the enzyme nicotinamide/nicotinate mononucleotide adenylyltransferase (NMNAT; EC: 2.7.7.1/18), in which converge the de novo and salvage pathways.

OBJECTIVE The purpose of this study was to analyse the protein-protein interactions (PPI) of NMNAT of *Leishmania braziliensis* (LbNMNAT) in promastigotes.

METHODS Transgenic lines of *L. braziliensis* promastigotes were established by transfection with the pSP72αneoLbNMNAT-GFP vector. Soluble protein extracts were prepared, co-immunoprecipitation assays were performed, and the co-immunoprecipitates were analysed by mass spectrometry. Furthermore, bioinformatics tools such as network analysis were applied to generate a PPI network.

FINDINGS Proteins involved in protein folding, redox homeostasis, and translation were found to interact with the LbNMNAT protein. The PPI network indicated enzymes of the nicotinate and nicotinamide metabolic routes, as well as RNA-binding proteins, the latter being the point of convergence between our experimental and computational results.

MAIN CONCLUSION We constructed a model of PPI of LbNMNAT and showed its association with proteins involved in various functions such as protein folding, redox homeostasis, translation, and NAD synthesis.

Key words: *Leishmania braziliensis* - protein-protein interaction - Co-IP-MS/MS - NAD - NMNAT

The cellular functions of proteins occur via concerted interactions with other proteins. Such protein-protein interactions (PPI) are the foundation of all biological processes occurring in different physiological and pathological conditions. They regulate several cellular functions, including cell cycle progression, signal transduction, metabolism, gene expression, vesicular transport, nuclear transport, and cell migration. Furthermore, members of the PPI network are potential therapeutic targets for the development of new drugs. Therefore, it is important to study and understand the specific nature of these interactions.

One such protein is the nicotinamide/nicotinate mononucleotide adenylyltransferase of *Leishmania braziliensis* (LbNMNAT), which is the key member in the biosynthesis of nicotinamide adenine dinucleotide (NAD) and has been functionally identified. PPI analysis by interaction networks has enabled the study of several biological aspects of human pathogens, such as the protozoan parasites *Leishmania* spp., *Plasmodium* spp., and *Trypanosoma* spp., which require the continuous discovery of pharmacological targets owing to their resistance mechanisms against existing drugs. In the case of *Leishmania*, PPI networks based on structural information have opened possibilities for further studies aimed at the development of novel drugs. These studies are crucial in the context of neglected diseases such as leishmaniasis, which is endemic to the tropical and subtropical regions of the world and causes approximately 30,000 deaths annually. Currently, there is no effective leishmaniasis vaccine available for humans. Therefore, it is necessary to explore the pathogenesis mechanisms of this parasite further, with focus on the identification of the PPI of key enzymes such as NMNAT. This offers a potential strategy for identifying and characterising new pharmacological targets. As proteins generally do not function in an isolated manner but are a part of a dynamic network, identifying the interaction partners of LbNMNAT can provide considerable information on their physiological function in the parasite.

Co-immunoprecipitation in combination with mass spectrometry (Co-IP-MS/MS) has been widely used for the sensitive and reliable identification of PPI and the characterisation of protein complex members. The interactions identified through the Co-IP-MS/MS strategy have been validated by other technologies, which reinforce the efficacy of MS to identify protein-protein associations.

In this study, we aimed to investigate for the first time, the PPI of LbNMNAT by using experimental and bioinformatics tools such as Co-IP-MS/MS and PPI network construction, respectively. The identification of the members of the interaction network around LbNMNAT...
will expand the comprehension on its regulation in *L. braziliensis*. Furthermore, these regulatory members represent potential pharmacological targets to modulate NAD biosynthesis in the parasite.

**MATERIALS AND METHODS**

**Culture of *L. braziliensis* promastigotes -** *L. braziliensis* (M2904 MHOM/BR/75M2904) promastigotes were cultivated in Schneider’s medium (pH 7.4; Sigma), supplemented with 10% (v/v) foetal bovine serum (FBS) at 26°C, in 25-cm² (T25) flasks. When the culture reached the stationary growth phase, the parasites were diluted to an initial concentration of 1 × 10⁶ cells/mL.

**Construction of the pSP72RαneoαLbNMNAT-GFP recombinant vector -** The fragment *Lbmnat* was polymerase chain reaction (PCR)-amplified using the plasmid pQE30-LbNMNAT as a template and the following oligonucleotides: forward 5’-GGCTCTAGAATGTTATCCTACTGCT-3’ and reverse 5’-GGCTCTAGGAGGCAAGCCTC-3’. The PCR product was cloned in the pGEMT-Easy (Promega) vector and was released from this vector employing the endonuclease XbaI (Fermentas), using the T4 ligase. The digested fragment was ligated with the expression vector pSP72RαneoαGFP (kindly donated by Dr Marcela Camacho), using the T4 ligase. The vector was previously digested with XbaI and dephosphorylated with alkaline phosphatase from calf intestine (Promega) for 2 h at 37°C. The digested vector was verified by DNA sequencing.

**Transfection of *L. braziliensis* promastigotes -** The recombinant vectors pSP72RαneoαLbNMNAT-GFP and pSP72RαneoαGFP (empty) were used to transfect *L. braziliensis* promastigotes, following a standardised electroporation protocol. Briefly, parasites from 10 mL of a confluent culture were washed with 3 mL of cold Cytomix (25 mM HEPES pH 7.6, 120 mM KCl, 10 mM K₂HPO₄, 5 mM MgCl₂, 2 mM EDTA, 0.15 mM CaCl₂), centrifuged at 1000 g for 10 min at 4°C. The precipitate of parasites was re-suspended in 1 mL of Cytomix and 450 μL of this suspension was transferred to a 4-mm electroporation cell containing 25 μF of the corresponding plasmids. The cell was incubated on ice for 10 min and 3 pulses of 1600 V, 25 μF capacitance and infinite resistance were applied, with intervals of 20 s between the pulses. The electroporated cells were incubated 24 h at 26°C in a final volume of 5 mL of Schneider’s medium supplemented with 20% (v/v) FBS. Then, 5 mL of this medium supplemented with 120 μg/mL of genetin G418 (Gibco-BRL) was added. The cultures were monitored daily by microscopic observations. Fifteen days later, live parasites were observed, and the established transgenic lines were cultured in the presence of 60 μg/mL G418. The transgenic parasites were used in immunodetection assays and fluorescence microscopy. GFP emission in paraformaldehyde-fixed cells was observed using the Nikon Eclipse C1 Plus microscope and the EZ-C1 software.

**Soluble protein extract preparation -** Transgenic parasites of each cell line were collected by centrifugation at 6000 g for 10 min at 4°C, using 100 mL of the culture. The parasites were washed two times with 10 mL of PBS (pH 7.4), re-suspended in 500 μL of lysis buffer (0.1% PBS, cocktail of protease inhibitors (1:200; Sigma), 0.1% (v/v) Triton X-100) and incubated for 30 min at 4°C with constant agitation. The suspension was centrifuged at 12000 g for 5 min at 4°C and the supernatant was stored at -20°C.

**Co-IP assays -** Soluble protein extracts (900 μL) were incubated with 100 μL of protein A-Sepharose (Sigma) for 1 h at 4°C with constant agitation and centrifuged at 3000 g for 3 min at 4°C. The anti-GFP antibody (Molecular Probes) was added to the clarified supernatant, to a final concentration of 0.2 μg/mL. The reaction was incubated for 16 h at 4°C with agitation. Then, 100 μL of protein A-Sepharose was added and incubated for 2 h with agitation at 4°C. The immunoprecipitate was obtained by centrifugation at 3000 g for 5 min at 4°C. The precipitate was washed three times with washing buffer (50 mM Tris-HCl, pH 7.5, 150 mM NaCl, 5 mM MgCl₂, 5% (v/v) glycerol). The immunoprecipitate was re-suspended in 100 μL of 1X sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE) loading buffer (50 mM Tris-HCl, pH 6.8, 10% (v/v) glycerol, 2% (w/v) SDS, 0.1% (w/v) bromophenol blue, 100 mM β-mercaptoethanol) and heated at 92°C for 10 min. The process was monitored by SDS-PAGE and western blot. For immunodetection, anti-GFP [1:500 in tris-buffered saline (TBS)] was used as the primary antibody, and peroxidase-conjugated anti-rabbit immunoglobulin G (IgG) (1:5000 in TBS, Sigma) was used as the secondary antibody. Membranes were revealed with 4-chloronaphthol (Promega) and hydrogen peroxide.

**Analysis of co-IP proteins by MS -** Here, 60 μL of the immunoprecipitates was analysed using T12 SDS-PAGE at 200 V for 5 min. Staining of the gel was completed with colloidal Coomassie G250 (10% (w/v) (NH₄)₂SO₄, 0.1% (w/v) Coomassie G250, 3% (v/v) H₂PO₄, 20% (v/v) ethanol) for 16 h. The immunoprecipitated protein bands were extracted from the gel and identified by commercial nano-LC-MS/MS (https://www.alphalyse.com/). Samples were reduced with 150 μL 10 mM DTT in 50 mM NH₄HCO₃ and incubated for 30 min at 50°C. Then, the samples were alkylated with 150 μL of 55 mM iodoacetamide in 50 mM NH₄HCO₃ and incubated for 30 min at room temperature, and digested with trypsin. The peptides were concentrated by lyophilisation and redissolved prior to being injected in a UPLC Dionex nano-LC system. The peptides were separated using a trap column (Acclaim PepMap 100, 75 μm × 2 cm) coupled to an analytic column (Acclaim PepMap RSLC, 75 μm × 15 cm) at 35°C using eluent A (0.1% formic acid) and B (0.1% formic acid in 90% acetonitrile) with the following gradient: 10 min 5% B, 30 min 45% B, 5 min 98% B, 10 min 5% B; flow: 0.3 μL/min. MS/MS analysis was conducted in the Bruker Maxis Impact QTOF instrument using a CaptiveSpray with nanoBoosteR source (0.2 Bar), positive polarity mode, 50-2200 m/z mass range, 1500V capillary voltage, 3 L/min of dry gas, and 150°C of dry temperature.
**Protein identification** - Proteomics data analysis was performed using the PatternLab for Proteomics software 4.0. Initially, the protein database of *L. braziliensis* (MHOM/BR/75/M2904) was loaded in the program from UniProt. As search parameters, we allowed two missed cleavages after trypsin digestion, precursor mass tolerance of 40 ppm, and carbamidomethylation (c) and methionine oxidation as fixed and variable modifications, respectively. Label-free quantification between common identified proteins in the biological replicates and the control sample was completed based on spectrum counting by means of the ratio: Spectrum counts in a biological replicate/Spectrum counts in the control sample. (9) Fold change ≥ 1.5 was selected as the parameter.

**Identification of interactions of the LbNMNAT protein by network analysis** - The PPI network was created with the platform STRING V.10.5 (Search Tool for the Retrieval of Interacting Genes/Proteins).(10) Resources used were neighbourhood, gene-fusion, text mining, co-expression, co-occurrence, and databases as well as experimental results, establishing a value of 0.4 as a minimum interaction score. Functional enhancement of the proteins was performed using functional classification systems in GO (Gene Ontology), and the functional and conserved modules were predicted using the databases InterPRO (https://www.ebi.ac.uk/interpro/), PFAM (https://pfam.xfam.org/), and KEGG (http://www.genome.jp/kegg/). Finally, the hypothetical proteins and interesting interactions were analysed. To contrast the results with experimentally confirmed physical PPI, a search was made for NMNAT physical interactions in *Homo sapiens*, *Escherichia coli*, *Saccharomyces cerevisiae*, and Arabidopsis *thaliana* in the online databases DIP (http://dip.doe-mbi.ucla.edu/dip/Main.cgi), BioGRID (https://thebiogrid.org/), and IntAct (https://www.ebi.ac.uk/intact/).

**RESULTS**

LbNMNAT-GFP co-immunoprecipitates with diverse interaction partners - Two transgenic cellular lines of *L. braziliensis* were established using the pSP72neoLbNMNAT-GFP and pSP72neoGFP vectors. Soluble protein extracts were obtained and analysed by western blot, whereas the parasites were observed by fluorescence microscopy to confirm LbNMNAT-GFP expression in the promastigotes. Parasites transfected with the empty vector expressed GFP (27 kDa), whereas those transfected with the recombinant pSP72neoLbNMNAT-GFP vector expressed the LbNMNAT enzyme fused to GFP (60 kDa) (Fig. 1).

Soluble protein extracts from parasites expressing LbNMNAT-GFP were used to perform co-immunoprecipitation experiments. To verify the dynamic nature of the PPI, two independent and complementary biological replicates were performed using asynchronous cellular cultures. To avoid unreliable positive results due to the co-immunoprecipitation of unspecified proteins, a parallel Co-IP assay was performed using soluble protein extracts from parasites that expressed GFP exclusively (control experiment). The obtained samples were analysed by SDS-PAGE and western blot. LbNMNAT-GFP was detected in each case, revealing that this protein can be immunoprecipitated adequately and specifically from soluble extracts (Fig. 2).

The co-immunoprecipitated proteins were analysed by nano-LC-MS/MS. Those common among the biological replicates 1 and 2 along with those common with the control sample but showing an upregulation of 1.5 fold or higher in the replicates are listed in Table I. We identified 103 possible interacting proteins, and the complete protein sets for all Co-IP-MS/MS experiments are listed in Supplementary data (Table). The putative functions of these proteins were obtained from the databases UniProt, GeneDB, and TriTrypDB.

As mentioned, two independent Co-IP-MS/MS assays, using the soluble protein extracts from LbNMNAT-GFP-expressing parasites were performed. The proteins identified in each experiment were grouped and classified into six biochemical categories: proteins involved in translation, protein folding, redox homeostasis, biosynthetic processes, and other biological processes, and proteins with uncharacterised functions (Table II). The observed variation in the number of members per category could be attributed to the samples being obtained from asynchronous cellular cultures in which the promastigotes can exhibit relevant parasite development-related biochemical changes, which in some cases can modulate biological functions. Fig. 3 shows the functional category-based distribution of proteins identified using Co-IP-MS/MS in biological replicates 1 and 2 together.

**Generation of a PPI network based on the LbNMNAT protein** - Several in silico methods have been developed to confirm the interactions detected experimentally. The computational methods for PPI prediction include approaches based on sequences, structures, chromosomal proximity, genetic fusion, in silico double hybrid, phylogenetic trees, gene ontology, and gene expression. Several web servers are available for the development of such analyses. (11) In this study, we used an integrative approach for obtaining the physical and functional associations to create a PPI network; accordingly, we used the STRING web resource, which manages diverse sources of information facilitating a comparative analysis. (12)

A PPI network for the LbNMNAT protein was generated (Fig. 4). The red node XP_001563913.1 (A4H990) represents the protein of interest and the remaining nodes represent the proteins interacting with LbNMNAT (Table III). The score for each node represents the confidence for each association and is derived by comparing the predictions and a reference set. (13) During the creation of this network, it was arbitrarily established that the score of the minimal required interaction should be 0.4, employing all the sources of information except textmining. To validate the selected score, a PPI network based on the HsNMNAT1 protein was constructed, establishing two scores of minimal interaction: 0.9 (high) and 0.4 (medium). Identical results, in terms of number and identity of protein nodes, were obtained (data not shown). The physical interaction of HsNMNAT1 with RNA-binding proteins and sirtuins has been demonstrated, (14,15) thus validating our bioinformatics approach.
For LbNMNAT, we tested the same scores of minimal interaction (0.9 and 0.4) and detected 5 and 11 protein nodes, respectively. Considering that the obtained confidence scores were above 0.6 in the last case (0.4), the PPI network was constructed using this score of minimal interaction to present a more expanded network, which contains RNA-binding protein interactors, in accordance with our data obtained from Co-IP experiments. Among the parameters that describe the topology of the network, 11 nodes with a total of 16 connections, a 2.91 grade per node and a clustering coefficient of 0.885 were found.

The totality of the network proteins is predicted computationally from its nucleic acid sequence and has not been demonstrated experimentally. This can be attributed to the presence of high number of hypothetical proteins (~60%) in trypanosomatid genomes. Next, functional enhancement of the network was performed considering the information from the KEGG, PFAM, and INTERPRO databases, from which data on the biological routes and the predominant domains in the interacting proteins were collected. Five of the proteins, including NMNAT, are related to the metabolic route of nicotinate and nicotinamide, while four of the members exhibit RNA-binding motifs (Fig. 4), in concordance with our Co-IP-MS/MS experimental analysis.

**DISCUSSION**

Proteins carry out their cellular functions via concerted interactions with other proteins, being PPI the basis of all biological processes. In this study, we found that proteins with diverse cellular functions interacted with LbNMNAT, and this can be explained by the cellular complexity, particularly the regulation of interactions in response to a particular signal, stimulus, or specific cellular state. Furthermore, they depend on many factors such as the stage of cellular development, the cell cycle phase, external conditions, and the presence of other proteins.

The LbNMNAT protein and its potential association with the ribosome - In the functional classification of each set of co-immunoprecipitated samples, proteins mostly related to translation were found; 31 sequences are related to this process. Some of these are RNA-binding proteins or structural constituents of the ribosome. To validate their potential connection with LbNMNAT, interactions already characterised in others NMNATs of widely studied organisms were reviewed.

In *S. cerevisiae*, for example, NMNAT1 interacts physically with the RNA-binding proteins, MPT5 and HEK2 proteins. Human HsNMNAT1 has been found to interact with RBM4B, another RNA-binding protein which is also related to circadian regulation, and with RPL30 (ribosomal protein L30), a structural component of the ribosome. Furthermore, HsNMNAT1 interacts with STAUI, which is associated with double-stranded RNA and can play a role in the specific positioning of the mRNAs in specific areas of the cell. Additionally, through the affinity capture-MS technique, the interaction of HsNMNAT1 with the ribosomal proteins L11 (RPL11) and L22 (RPL22) was identified. These proteins are necessary for rRNA formation, maturation, and processing.

Currently, the significance of the interaction of NMNAT with the ribosome is unknown. Further research focusing on the stability of the complex and the interactions among their diverse components is essential because ribosome assembly and protein translation are finely coordinated with cellular growth, proliferation, differentiation, and development and because NAD is involved in all these processes.
The LbNMNAT protein may participate in the folding of other proteins - Among the co-immunoprecipitated proteins, 14 candidates were functionally related to the folding process. Cytoplasmic chaperones belonging to the T-complex and heat shock proteins (Hsp70, Hsp83, and Hsp60 chaperonin) were found. The Leishmania genome contains four genes encoding mitochondrial chaperonins: two of them, CPN60.2 and CPN10, are orthologs of the bacterial GroEL/GroES system. Pull down experiments for NMNATs in other organisms revealed that the homologous NadD of E. coli interacts with GroEL (Hsp60). 

Given the presence of T-complex proteins, a network of theoretical interactions was created using the software STRING. A network of high reliability (p-value: 5.67e-05) was obtained, with 10 nodes and 25 interactions (data not shown). This network predicts coordinated interactions among the different chaperonins of L. braziliensis, including the immunoprecipitated candidates. Potential interaction of LbNMNAT can be related to its protein-folding capacity, as has already been demonstrated in other NMNATs with chaperone functions independent of its NAD synthesizing activity. This function can be considered as a response to stressful conditions, specifically drastic temperature changes, which is important for the survival of the parasite throughout its biological cycle. The NMNAT-chaperone interaction has been demonstrated previously. In H. sapiens, HsNMNAT2 not only acts as a chaperone to reduce protein aggregates but also interacts with Hsp90 to retract them. Studies in Drosophila melanogaster have demonstrated that its NMNAT is necessary for thermal tolerance and responses against oxidative stress. Additionally, regulation of DmNMNAT and Hsp70 under stressful conditions has been shown to be considerably different at the transcriptional level, suggesting that the NMNAT, which exhibits constitutive expression, may represent a different class of stress-response proteins. Because it is a housekeeping protein, NMNAT is available under normal conditions and can provide a primary response to a stress condition, reducing the resulting proteotoxicity.
**Table I**

Co-immunoprecipitation in combination with mass spectrometry (Co-IP-MS/MS) identified proteins in biological replicates 1 and 2

| Protein name | Uniprot access | Score | Biological replicate | Fold change |
|--------------|----------------|-------|----------------------|-------------|
| Nicotinamide mononucleotide adenylyltransferase | A4H990 | 16.83 | 1 and 2 | - |
| Putative 40S ribosomal protein S15 | A4HC91 | 19.01 | 1 and 2 | - |
| Alpha tubulin | A4HHS1 | 7.16 | 1 and 2 | - |
| Putative heat-shock protein hsp70 (Fragment) | A4HGX9 | 6.71 | 1 and 2 | - |
| Putative cystathione gamma lyase | A4HMZ0 | 7.99 | 1, 2 and control | 1.5; 5.5 |
| Tubulin alpha chain | A4H727 | 19.83 | 1, 2 and control | 7; 8 |
| Putative 40S ribosomal protein S3 | A4H868 | 5.93 | 1, 2 and control | 2; 3 |
| Putative 40S ribosomal protein S3 | A4HLE6 | 5.93 | 1, 2 and control | 2; 3 |
| Putative heat shock 70-related protein 1, mitochondrial | A4HIH7 | 20.50 | 1 and control | 1.5 |
| Putative heat shock 70-related protein 1, mitochondrial | A4HH9 | 20.50 | 1 and control | 1.5 |
| T-complex protein 1 subunit | A4HNS7 | 13.56 | 2 and control | 1.7 |
| Contig, possible fusion of chromosomes 20 and 34 | E9AIH1 | 7.11 | 2 and control | 1.5 |
| Putative 3-ketoacyl-CoA thiolase | A4HCU5 | 5.27 | 2 and control | 3 |
| T-complex protein 1 subunit gamma | A4HCZ3 | 36.55 | 2 and control | 6.5 |
| Putative T-complex protein 1, theta subunit | A4HQL2 | 12.16 | 2 and control | 3 |
| T-complex protein 1 subunit epsilon | A4HKK2 | 21.02 | 2 and control | 2 |
| Putative translation elongation factor 1-beta | A4HP03 | 13.35 | 2 and control | 1.5 |
| Tubulin beta chain | A4HLC9 | 23.32 | 2 and control | 2.2 |
| Beta-tubulin | A4HLD6 | 8.47 | 2 and control | 2.5 |
| Beta-tubulin | A4HLD1 | 14.89 | 2 and control | 1.6 |
| Tubulin beta chain | A4HC48 | 23.32 | 2 and control | 2.2 |
| Tubulin beta chain | A4HLC8 | 23.32 | 2 and control | 2.2 |
| EF2-1 protein | A4HNM6 | 8.84 | 2 and control | 1.5 |
| Adenosylhomocysteinase | A4HPQ8 | 11.21 | 2 and control | 3 |
| Adenosylhomocysteinase | A4HPQ9 | 11.21 | 2 and control | 6 |
| ENOL protein (Fragment) | A4HTT5 | 5.72 | 2 and control | 2 |
| ENOL protein | A4HTT6 | 21.78 | 2 and control | 2.3 |

*: identification value assigned by PatternLab for Proteomics 4.0; **: label free quantification based on spectral counting. Proteins 5-8 present two values corresponding to fold changes between biological replicates 1 or 2 vs control sample, respectively.

LbNMNAT may be involved in redox homeostasis - The enzymes, gamma-glutamylcysteine synthetase 1 (GSH1) and tryparedoxin peroxidase (TXNPx), which are involved in redox homeostasis, were also identified. Of these, TXNPx presents a greater potential for interaction with LbNMNAT.

The parasite can survive in the host owing to its redox metabolism, which counter arrests the radicals produced by infected macrophages. To avoid cellular damage by reactive oxygen species (ROS) and reactive nitrogen intermediates (RNI), trypanosomatids possess peroxidases that reveal a unique characteristic in the use of reducing equivalents derived from trypanothione. The combined action of TXNPx with antioxidant enzymes is crucial for the maintenance of a low concentration of $\text{H}_2\text{O}_2$, regulating the oxidative and nitrosative stress via the displacement of reducing equivalents from NADPH to hydroperoxides and peroxyrenitrites. Functional analysis has revealed that TXNPx is involved in the antimony resistance phenotype in *L. braziliensis*, suggesting that an increase in the expression levels of this enzyme plays a role in the clinical resistance to compounds derived from this element.

Leishmania TXNPx has been studied previously,[24,25,26,27] and it would be interesting to investigate whether a physical association exists among them. Co-IP assays have shown that the corresponding human homologues interact with each other.[28] However, a direct interaction between these enzymes and the NMNAT protein has not been observed, and the interaction in homologues of other species has not been reported. Therefore, validation of a potential association through other experimental strategies is necessary, allowing the generation of new information in relation to metabolic routes and redox regulation under stressful conditions in the parasite.

LbNMNAT PPI network - Considering the PPI network, the proteins linked to the nicotinate/nicotinamide metabolism show a functional association. Among these proteins, putative nicotinate phosphoribosyltransferase (NaPR) (XP_001567885.1) and putative NAD synthase (NADS) (XP_001567539.1) showed the highest scores. These enzymes belong to the salvage pathway of NAD synthesis. The substrate for NaPR is nicotinic acid produced by nicotinamidase, which has been experimentally characterised.[29] The product of the NaPR-catalysed
reaction is nicotinic acid mononucleotide, the subsequent substrate of LbNMNAT. NADS catalyses the conversion of the LbNMNAT product, NAAD, into NAD, thus completing the salvage pathway. The three proteins (NaPRT, LbNMNAT, and NADS) act in series, which explains their high scores.

SIR2 in other Leishmania spp. has been studied previously, and the parasite is known to express three sirtuins: SIR2RP1, SIR2RP2, and SIR2RP3. After using the databases BioGRID and IntAct and identifying the homologues that interact in other species, it was found that human NMNAT1 physically interacts with SIRT1 and NMNAT2 interacts with SIR4 in S. cerevisiae. Some authors have hypothesised that the close physical proximity of NMNAT1 and SIRT1 facilitate the most efficient use of NAD, probably through a substrate-channelling mechanism. Some mention that the enzymes involved in NAD biosynthesis can form a complex between this molecule and SIRT1, creating a micro-domain of high concentration of NAD that allows the regulation of sirtuin activity. Through direct validation, it would be interesting to study whether the physical interaction of NMNAT-SIR2 occurs in Leishmania.

We also found four RNA-binding proteins in the network. Their linkage was the product of relating homologous neighbour genes in other genomes (neighbourhood) and of the interaction of putative homologues in other species (Campylobacter curvus). The common characteristic of these proteins is that they all exhibit the RRM motif, found in several RNA-binding proteins. Our experimental results, in which 31 co-immunoprecipitated proteins (identified by MS) correspond to RNA-binding proteins,
In conclusion - Using experimental and computational methods, we developed an integrated model of protein-protein interactions of the LbNMNAT enzyme. The model indicates the interaction of proteins related to a wide range of cellular activities, including protein folding, redox homeostasis, translation, and NAD synthesis. Our results expand the knowledge of NAD metabolism in L. braziliensis and offer new directions for scientific exploration, from an interactome point of view.

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AUTHORS’ CONTRIBUTION
LOJ designed and performed the experimental and bioinformatics procedures, namely protein soluble extract preparation, Co-IP assays, MS/MS analysis, and identification of protein interactors by STRING predictions; and prepared the manuscript. LECR designed and performed recombinant vector construction, in vitro cell culture and promastigote transfection, and MS/MS analysis; and prepared the manuscript. MHRH provided a general direction of the research. All the authors have read and approved the manuscript.

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List of predicted LbNMNAT interacting proteins according to STRING

| Access*   | Identifier (ID)** | Predicted functional partners | Score |
|-----------|-------------------|-------------------------------|-------|
| A4H990    | XP_001563913.1    | LbNMNAT                       | -     |
| A4HLF1    | XP_001567885.1    | Putative nicotinate phosphoribosyltransferase (413 aa) | 0.964 |
| A4HKF4    | XP_001567539.1    | Putative NAD synthase (293 aa) | 0.964 |
| A4HL2     | XP_00156981.2     | ATP-NAD kinase-like protein (1257 aa) | 0.911 |
| E9AIR9    | XP_003723185.1    | Putative NAD-dependent SIR2 (284 aa) | 0.909 |
| A4HFG6    | XP_00156810.1     | Putative ribokinase (329 aa)   | 0.856 |
| A4HFG7    | XP_001562450.1    | Putative nitrate (279 aa)      | 0.658 |
| A4HMS6    | XP_001568356.1    | Putative RNA binding protein (458 aa) | 0.629 |
| A4HLK8    | XP_001567942.1    | Putative RNA-binding protein (129 aa) | 0.629 |
| A4HUH0    | XP_001566631.1    | Hypothetical protein (210 aa)  | 0.629 |
| A4HHR3    | XP_001566494.1    | Putative RNA binding protein (648 aa) | 0.629 |

*: access number in UniProt; **: access number in NCBI.
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